## The Finite

# Element Method and Applications in Engineering Using ANSYS ${ }^{\circledR}$ 

Erdogan Madenci

Ibrahim Guven


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Springer

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## PREFACE

The finite element method (FEM) has become a staple for predicting and simulating the physical behavior of complex engineering systems. The commercial finite element analysis (FEA) programs have gained common acceptance among engineers in industry and researchers at universities and government laboratories. Therefore, academic engineering departments include graduate or undergraduate senior-level courses that cover not only the theory of FEM but also its applications using the commercially available FEA programs.

The goal of this book is to provide students with a theoretical and practical knowledge of the finite element method and the skills required to analyze engineering problems with ANSYS ${ }^{\circledR}$, a commercially available FEA program. This book, designed for seniors and first-year graduate students, as well as practicing engineers, is introductory and self-contained in order to minimize the need for additional reference material.

In addition to the fundamental topics in finite element methods, it presents advanced topics concerning modeling and analysis with ANSYS ${ }^{\circledR}$. These topics are introduced through extensive examples in a step-by-step fashion from various engineering disciplines. The book focuses on the use of ANSYS ${ }^{\circledR}$ through both the Graphics User Interface (GUI) and the ANSYS ${ }^{\oplus}$ Parametric Design Language (APDL). Furthermore, it includes a CD-ROM with the "input" files for the example problems so that the students can regenerate them on their own computers. Because of printing costs, the printed figures and screen shots are all in gray scale. However, color versions are provided on the accompanying CD-ROM.

Chapter 1 provides an introduction to the concept of FEM. In Chapter 2, the analysis capabilities and fundamentals of ANSYS ${ }^{\oplus}$, as well as practical modeling considerations, are presented. The fundamentals of discretization and approximation functions are presented in Chapter 3. The modeling techniques and details of mesh generation in ANSYS ${ }^{\circledR}$ are presented in Chapter 4. Steps for obtaining solutions and reviews of results are presented in Chapter 5. In Chapter 6, the derivation of finite element equations based on the method of weighted residuals and principle of minimum potential energy
is explained and demonstrated through example problems. The use of commands and APDL and the development of macro files are presented in Chapter 7. In Chapter 8, example problems on linear structural analysis are worked out in detail in a step-by-step fashion. The example problems related to heat transfer and moisture diffusion are demonstrated in Chapter 9. Nonlinear structural problems are presented in Chapter 10. Advanced topics concerning submodeling, substructuring, interaction with external files, and modification of ANSYS ${ }^{\circledR}$-GUI are presented in Chapter 11.

There are more than 40 example problems considered in this book; solutions to most of these problems using ANSYS ${ }^{\circledR}$ are demonstrated using GUI in a step-by-step fashion. The remaining problems are demonstrated using the APDL. However, the steps taken in either GUI- or APDL-based solutions may not be the optimum/shortest possible way. Considering the steps involved in obtaining solutions to engineering problems (e.g., model generation, meshing, solution options, etc.), there exist many different routes to achieve the same solution. Therefore, the authors strongly encourage the students/engineers to experiment with modifications to the analysis steps presented in this book.

We are greatly indebted to Connie Spencer for her invaluable efforts in typing, editing, and assisting with each detail associated with the completion of this book. Also, we appreciate the contributions made by Dr. Atila Barut, Mr. Erkan Oterkus, Ms. Abigail Agwai, Mr. Manabendra Das, and Mr. Bahattin Kilic in the solution of the example problems. The permission provided by ANSYS, Inc. to print the screen shots is also appreciated.

## TABLE OF CONTENTS

PREFACE ..... v
LIST OF PROBLEMS SOLVED ..... xiii
1 INTRODUCTION ..... 1
1.1 Concept ..... 1
1.2 Nodes ..... 3
1.3 Elements ..... 4
1.4 Direct Approach ..... 5
1.4.1 Linear Spring ..... 5
1.4.2 Heat Flow ..... 6
1.4.3 Assembly of the Global System of Equations ..... 8
1.4.4 Solution of the Global System of Equations ..... 12
1.4.5 Boundary Conditions ..... 13
2 FUNDAMENTALS OF ANSYS ..... 15
2.1 Useful Definitions ..... 15
2.2 Before an ANSYS Session ..... 16
2.2.1 Analysis Discipline ..... 16
2.2.2 Time Dependence ..... 18
2.2.3 Nonlinearity ..... 18
2.2.4 Practical Modeling Considerations ..... 19
2.3 Organization of ANSYS Software ..... 25
2.4 ANSYS Analysis Approach ..... 26
2.4.1 ANSYS Preprocessor ..... 27
2.4.2 ANSYS Solution Processor ..... 27
2.4.3 ANSYS General Postprocessor ..... 27
2.4.4 ANSYS Time History Postprocessor ..... 27
2.5 ANSYS File Structure ..... 27
2.5.1 Database File ..... 28
2.5.2 Log File ..... 28
2.5.3 Error File ..... 28
2.5.4 Results Files ..... 28
2.6 Description of ANSYS Menus and Windows ..... 29
2.6.1 Utility Menu ..... 30
2.6.2 Main Menu ..... 31
2.6.3 Toolbar ..... 32
2.6.4 Input Field ..... 32
2.6.5 Graphics Window ..... 32
2.6.6 Output Window ..... 32
2.7 Using the ANSYS Help System ..... 32
2.7.1 Help Contents ..... 33
2.7.2 Help Index ..... 33
2.7.3 Search in Help ..... 33
2.7.4 Verification Manual ..... 35
3 FUNDAMENTALS OF DISCRETIZATION ..... 37
3.1 Local and Global Numbering ..... 37
3.2 Approximation Functions ..... 37
3.3 Coordinate Systems ..... 43
3.3.1 Generalized Coordinates ..... 43
3.3.2 Global Coordinates ..... 43
3.3.3 Local Coordinates ..... 43
3.3.4 Natural Coordinates ..... 43
3.4 Shape Functions ..... 51
3.4.1 Linear Line Element with Two Nodes ..... 54
3.4.2 Quadratic Line Element with Three Nodes: Centroidal Coordinate ..... 56
3.4.3 Linear Triangular Element with Three Nodes: Global Coordinate ..... 58
3.4.4 Quadratic Triangular Element with Six Nodes ..... 59
3.4.5 Linear Quadrilateral Element with Four Nodes: Centroidal Coordinate ..... 62
3.5 Isoparametric Elements: Curved Boundaries ..... 64
3.6 Numerical Evaluation of Integrals ..... 68
3.6.1 Line Integrals ..... 68
3.6.2 Triangular Area Integrals ..... 72
3.6.3 Quadrilateral Area Integrals ..... 75
3.7 Problems ..... 78
4 ANSYS PREPROCESSOR ..... 83
4.1 Fundamentals of Modeling ..... 83
4.2 Modeling Operations ..... 83
4.2.1 Title ..... 84
4.2.2 Elements ..... 85
4.2.3 Real Constants ..... 89
4.2.4 Material Properties ..... 92
4.2.5 Element Attributes ..... 96
4.2.6 Interaction with the Graphics Window: Picking Entities ..... 96
4.2.7 Coordinate Systems ..... 99
4.2.8 Working Plane ..... 102
4.3 Solid Modeling ..... 105
4.3.1 Bottom-up Approach: Entities ..... 106
4.3.2 Top-down Approach: Primitives ..... 112
4.4 Boolean Operators ..... 118
4.4.1 Adding ..... 118
4.4.2 Subtracting ..... 120
4.4.3 Overlap ..... 120
4.4.4 Gluing ..... 121
4.4.5 Dividing ..... 121
4.5 Additional Operations ..... 124
4.5.1 Extrusion and Sweeping ..... 124
4.5.2 Moving and Copying ..... 128
4.5.3 Keeping/Deleting Original Entities ..... 128
4.5.4 Listing Entities ..... 130
4.5.5 Deleting Entities ..... 130
4.6 Viewing a Model ..... 131
4.6.1 Plotting: Pan, Zoom, and Rotate Functions ..... 131
4.6.2 Plotting/Listing Entities ..... 134
4.6.3 Numbers in the Graphics Window ..... 134
4.7 Meshing ..... 134
4.7.1 Automatic Meshing ..... 135
4.7.2 Manipulation of the Mesh ..... 141
4.8 Selecting and Components ..... 144
4.8.1 Selecting Operations ..... 144
4.8.2 Components ..... 148
5 ANSYS SOLUTION AND POSTPROCESSING ..... 149
5.1 Overview ..... 149
5.2 Solution ..... 150
5.2.1 Analysis Options/Solution Controls ..... 150
5.2.2 Boundary Conditions ..... 153
5.2.3 Initial Conditions ..... 154
5.2.4 Body Loads ..... 154
5.2.5 Solution in Single and Multiple Load Steps ..... 154
5.2.6 Failure to Obtain Solution ..... 158
5.3 Postprocessing ..... 160
5.3.1 General Postprocessor ..... 160
5.3.2 Time History Postprocessor ..... 160
5.3.3 Read Results ..... 161
5.3.4 Plot Results ..... 163
5.3.5 Element Tables ..... 167
5.3.6 List Results ..... 170
5.4 Example: One-dimensional Transient Heat Transfer ..... 170
6 FINITE ELEMENT EQUATIONS ..... 187
6.1 Method of Weighted Residuals ..... 187
6.1.1 Example: One-dimensional Differential Equation with Line Elements ..... 189
6.1.2 Example: Two-dimensional Differential Equation with Linear Triangular Elements ..... 197
6.1.3 Example: Two-dimensional Differential Equation with Linear Quadrilateral Elements ..... 216
6.2 Principle of Minimum Potential Energy ..... 235
6.2.1 Example: One-dimensional Analysis with Line Elements ..... 242
6.2.2 Two-dimensional Structural Analysis ..... 248
6.3 Problems ..... 289
7 USE OF COMMANDS IN ANSYS ..... 297
7.1 Basic ANSYS Commands ..... 297
7.1.1 Operators and Functions ..... 298
7.1.2 Defining Parameters ..... 304
7.2 A Typical Input File ..... 307
7.3 Selecting Operations ..... 309
7.4 Extracting Information from ANSYS ..... 314
7.5 Programming with ANSYS ..... 317
7.5.1 DO Loops ..... 317
7.5.2 IF Statements ..... 318
7.5.3 / OUTPUT and *VWRITE Commands ..... 321
7.6 Macro Files ..... 322
7.7 Useful Resources ..... 324
7.7.1 Using the Log File for Programming ..... 325
7.7.2 Using the Verification Problems for Programming ..... 326
8 LINEAR STRUCTURAL ANALYSIS ..... 329
8.1 Static Analysis ..... 329
8.1.1 Trusses ..... 329
8.1.2 Beams ..... 337
8.1.3 Three-dimensional Problems ..... 342
8.1.4 Two-dimensional Idealizations ..... 346
8.1.5 Plates and Shells ..... 373
8.2 Linear Buckling Analysis ..... 403
8.3 Thermomechanical Analysis ..... 412
8.4 Fracture Mechanics Analysis ..... 421
8.5 Dynamic Analysis ..... 433
8.5.1 Modal Analysis ..... 434
8.5.2 Harmonic Analysis ..... 444
8.5.3 Transient Analysis ..... 459
9 LINEAR ANALYSIS OF FIELD PROBLEMS ..... 477
9.1 Heat Transfer Problems ..... 477
9.1.1 Steady-state Analysis ..... 478
9.1.2 Transient Analysis ..... 520
9.1.3 Radiation Analysis ..... 543
9.2 Moisture Diffusion ..... 549
10 NONLINEAR STRUCTURAL ANALYSIS ..... 565
10.1 Geometric Nonlinearity ..... 569
10.1.1 Large Deformation Analysis of a Plate ..... 570
10.1.2 Post-buckling Analysis of a Plate with a Hole ..... 573
10.2 Material Nonlinearity ..... 578
10.2.1 Plastic Deformation of an Aluminum Sphere ..... 579
10.2.2 Plastic Deformation of an Aluminum Cylinder ..... 583
10.2.3 Stress Analysis of a Reinforced Viscoelastic Cylinder ..... 589
10.2.4 Viscoplasticity Analysis of a Eutectic Solder Cylinder ..... 592
10.2.5 Combined Plasticity and Creep ..... 599
10.3 Contact ..... 605
10.3.1 Contact Analysis of a Block Dropping on a Beam ..... 607
10.3.2 Simulation of a Nano-indentation Test ..... 613
11 ADVANCED TOPICS IN ANSYS ..... 621
11.1 Coupled Degrees of Freedom ..... 621
11.2 Constraint Equations ..... 624
11.3 Submodeling ..... 629
11.4 Substructuring: Superelements ..... 636
11.4.1 Generation Pass ..... 638
11.4.2 Use Pass ..... 644
11.4.3 Expansion Pass ..... 646
11.5 Interacting with External Files ..... 647
11.5.1 Reading an Input File ..... 647
11.5.2 Writing Data to External ASCII Files ..... 648
11.5.3 Executing an External File ..... 652
11.5.4 Modifying ANSYS Results ..... 654
11.6 Modifying the ANSYS GUI ..... 654
11.6.1 GUI Development Demonstration ..... 662
11.6.2 GUI Modification for Obtaining a Random Load Profile ..... 671
11.6.3 Function Block for Selecting Elements Using a Pick Menu ..... 675
REFERENCES ..... 679
INDEX ..... 681

## LIST OF PROBLEMS SOLVED

ANSYS Solution of a Two-dimensional Differential Equation with Linear Triangular Elements ..... 211
ANSYS Solution of a Two-dimensional Differential Equation with Linear Quadrilateral Elements ..... 230
Plane Stress Analysis with Linear Triangular Elements ..... 264
Plane Stress Analysis with Linear Quadrilateral Isoparametric Elements ..... 284
Elongation of a Bar Under Its Own Weight Using Truss Elements ..... 330
Analysis of a Truss Structure with Symmetry ..... 333
Analysis of a Slit Ring ..... 337
Elongation of a Bar Under Its Own Weight Using 3-D Elements ..... 342
Plane Stress Analysis of a Plate with a Circular Hole ..... 346
Plane Stress Analysis of a Composite Plate Under Axial Tension ..... 355
Plane Strain Analysis of a Bi-material Cylindrical Pressure Vessel Under Internal Pressure ..... 359
Deformation of a Bar Due to Its Own Weight Using 2-D Axisymmetric Elements ..... 366
Analysis of a Circular Plate Pushed Down by a Piston Head Using 2-D Axisymmetric Elements ..... 368
Static Analysis of a Bracket Using Shell Elements ..... 373
Analysis of a Circular Plate Pushed Down by a Piston Head Using Solid Brick and Shell Elements ..... 383
Analysis of an Axisymmetric Shell with Internal Pressure Using Shell Elements ..... 391
Analysis of a Layered Composite Plate Using Shell Elements ..... 397
Linear Buckling Analysis of a Plate ..... 403
Thermomechanical Analysis of an Electronic Package ..... 412
Fracture Mechanics Analysis of a Strip with an Inclined Edge Crack ..... 421
Modal Analysis of a Bracket ..... 434
Vibration Analysis of an Automobile Suspension System ..... 438
Harmonic Analysis of a Bracket ..... 444
Harmonic Analysis of a Guitar String ..... 453
Dynamic Analysis of a Bracket ..... 460
Impact Loading on a Beam ..... 465
Dynamic Analysis of a 4-bar Linkage ..... 471
Heat Transfer Analysis of a Tank/Pipe Assembly ..... 478
Heat Transfer Analysis of a Window Assembly ..... 499
Transient Thermomechanical Analysis of an Electronic Package ..... 522
Transient Thermomechanical Analysis of a Welded Joint ..... 532
Radiation Heat Transfer Analysis of a Conical Fin ..... 543
Moisture Diffusion Analysis of an Electronic Package ..... 549
Large Deformation Analysis of a Plate ..... 570
Postbuckling Analysis of a Plate with a Hole ..... 573
Plastic Deformation of an Aluminum Sphere ..... 579
LIST OF PROBLEMS SOLVED ..... XV
Plastic Deformation of an Aluminum Cylinder ..... 583
Stress Analysis of a Reinforced Viscoelastic Cylinder ..... 589
Viscoplasticity Analysis of a Eutectic Solder Cylinder ..... 592
Combined Plasticity and Creep Analysis of a Eutectic Solder Cylinder ..... 599
Contact Analysis of a Block Dropping on a Beam ..... 607
Simulation of a Nano-indentation Test ..... 613
Analysis of a Sandwich Panel Using Constraint Equations ..... 624
Submodeling Analysis of a Square Plate with a Circular Hole ..... 629
Substructuring Analysis of an Electronic Package ..... 636
GUI Development Demonstration ..... 662

## Chapter 1

## INTRODUCTION

### 1.1 Concept

The Finite Element Analysis (FEA) method, originally introduced by Turner et al. (1956), is a powerful computational technique for approximate solutions to a variety of "real-world" engineering problems having complex domains subjected to general boundary conditions. FEA has become an essential step in the design or modeling of a physical phenomenon in various engineering disciplines. A physical phenomenon usually occurs in a continuum of matter (solid, liquid, or gas) involving several field variables. The field variables vary from point to point, thus possessing an infinite number of solutions in the domain. Within the scope of this book, a continuum with a known boundary is called a domain.

The basis of FEA relies on the decomposition of the domain into a finite number of subdomains (elements) for which the systematic approximate solution is constructed by applying the variational or weighted residual methods. In effect, FEA reduces the problem to that of a finite number of unknowns by dividing the domain into elements and by expressing the unknown field variable in terms of the assumed approximating functions within each element. These functions (also called interpolation functions) are defined in terms of the values of the field variables at specific points, referred to as nodes. Nodes are usually located along the element boundaries, and they connect adjacent elements.

The ability to discretize the irregular domains with finite elements makes the method a valuable and practical analysis tool for the solution of boundary, initial, and eigenvalue problems arising in various engineering disciplines. Since its inception, many technical papers and books have appeared on the development and application of FEA. The books by Desai and Abel (1971), Oden (1972), Gallagher (1975), Huebner (1975), Bathe and Wilson (1976), Ziekiewicz (1977), Cook (1981), and Bathe (1996) have influenced the current state of FEA. Representative common engineering problems and their corresponding FEA discretizations are illustrated in Fig. 1.1.


Fig. 1.1 FEA representation of practical engineering problems.

The finite element analysis method requires the following major steps:

- Discretization of the domain into a finite number of subdomains (elements).
- Selection of interpolation functions.
- Development of the element matrix for the subdomain (element).
- Assembly of the element matrices for each subdomain to obtain the global matrix for the entire domain.
- Imposition of the boundary conditions.
- Solution of equations.
- Additional computations (if desired).

There are three main approaches to constructing an approximate solution based on the concept of FEA:

Direct Approach: This approach is used for relatively simple problems, and it usually serves as a means to explain the concept of FEA and its important steps (discussed in Sec. 1.4).

Weighted Residuals: This is a versatile method, allowing the application of FEA to problems whose functionals cannot be constructed. This approach directly utilizes the governing differential equations, such as those of heat transfer and fluid mechanics (discussed in Sec. 6.1).

Variational Approach: This approach relies on the calculus of variations, which involves extremizing a functional. This functional corresponds to the potential energy in structural mechanics (discussed in Sec. 6.2).

In matrix notation, the global system of equations can be cast into

$$
\begin{equation*}
\mathbf{K u}=\mathbf{F} \tag{1.1}
\end{equation*}
$$

where $\mathbf{K}$ is the system stiffness matrix, $\mathbf{u}$ is the vector of unknowns, and $\mathbf{F}$ is the force vector. Depending on the nature of the problem, $\mathbf{K}$ may be dependent on $\mathbf{u}$, i.e., $\mathbf{K}=\mathbf{K}(\mathbf{u})$ and $\mathbf{F}$ may be time dependent, i.e., $\mathbf{F}=\mathbf{F}(t)$.

### 1.2 Nodes

As shown in Fig. 1.2, the transformation of the practical engineering problem to a mathematical representation is achieved by discretizing the domain of interest into elements (subdomains). These elements are connected to each other by their "common" nodes. A node specifies the coordinate location in space where degrees of freedom and actions of the physical problem exist. The nodal unknown(s) in the matrix system of equations represents one (or more) of the primary field variables. Nodal variables assigned to an element are called the degrees of freedom of the element.

The common nodes shown in Fig. 1.2 provide continuity for the nodal variables (degrees of freedom). Degrees of freedom (DOF) of a node are dictated by the physical nature of the problem and the element type. Table 1.1 presents the DOF and corresponding "forces" used in FEA for different physical problems.


Fig. 1.2 Division of a domain into subdomains (elements).

Table 1.1 Degrees of freedom and force vectors in FEA for different engineering disciplines.

| Discipline | DOF | Force Vector |
| :---: | :---: | :---: |
| Structural/solids | Displacement | Mechanical forces |
| Heat conduction | Temperature | Heat flux |
| Acoustic fluid | Displacement potential | Particle velocity |
| Potential flow | Pressure | Particle velocity |
| General flows | Velocity | Fluxes |
| Electrostatics | Electric potential | Charge density |
| Magnetostatics | Magnetic potential | Magnetic intensity |

### 1.3 Elements

Depending on the geometry and the physical nature of the problem, the domain of interest can be discretized by employing line, area, or volume elements. Some of the common elements in FEA are shown in Fig. 1.3. Each element, identified by an element number, is defined by a specific sequence of global node numbers. The specific sequence (usually counterclockwise) is based on the node numbering at the element level. The node numbering sequence for the elements shown in Fig. 1.4 are presented in Table 1. 2.


Fig. 1.3 Description of line, area, and volume elements with node numbers at the element level.


Fig. 1.4 Discretization of a domain: element and node numbering.

Table 1.2 Description of numbering at the element level.

| Element <br> Number | Node 1 | Node 2 | Node 3 | Node 4 |
| :---: | :---: | :---: | :---: | :---: |
| 1 | 1 | 2 | 6 | 7 |
| 2 | 3 | 4 | 6 | 2 |
| 3 | 4 | 5 | 6 |  |

### 1.4 Direct Approach

Although the direct approach is suitable for simple problems, it involves each fundamental step of a typical finite element analysis. Therefore, this approach is demonstrated by considering a linear spring system and heat flow in a one-dimensional (1-D) domain.

### 1.4.1 Linear Spring

As shown in Fig. 1.5, a linear spring with stiffness $k$ has two nodes. Each node is subjected to axial loads of $f_{1}$ and $f_{2}$, resulting in displacements of $u_{1}$ and $u_{2}$ in their defined positive directions.

Subjected to these nodal forces, the resulting deformation of the spring becomes

$$
\begin{equation*}
u=u_{1}-u_{2} \tag{1.2}
\end{equation*}
$$



Fig. 1.5 Free-body diagram of a linear spring element.
which is related to the force acting on the spring by

$$
\begin{equation*}
f_{1}=k u=k\left(u_{1}-u_{2}\right) \tag{1.3}
\end{equation*}
$$

The equilibrium of forces requires that

$$
\begin{equation*}
f_{2}=-f_{1} \tag{1.4}
\end{equation*}
$$

which yields

$$
\begin{equation*}
f_{2}=k\left(u_{2}-u_{1}\right) \tag{1.5}
\end{equation*}
$$

Combining Eq. (1.3) and (1.5) and rewriting the resulting equations in matrix form yield

$$
\left[\begin{array}{cc}
k & -k  \tag{1.6}\\
-k & k
\end{array}\right]\left\{\begin{array}{l}
u_{1} \\
u_{2}
\end{array}\right\}=\left\{\begin{array}{l}
f_{1} \\
f_{2}
\end{array}\right\} \quad \text { or } \quad \mathbf{k}^{(e)} \mathbf{u}^{(e)}=\mathbf{f}^{(e)}
$$

in which $\mathbf{u}^{(e)}$ is the vector of nodal unknowns representing displacement and $\mathbf{k}^{(e)}$ and $\mathbf{f}^{(e)}$ are referred to as the element characteristic (stiffness) matrix and element right-hand-side (force) vector, respectively. The superscript ( $e$ ) denotes the element numbered as ' $e$ '.

The stiffness matrix can be expressed in indicial form as $k_{i j}^{(e)}$

$$
\begin{equation*}
\mathbf{k}^{(e)} \sim k_{i j}^{(e)} \tag{1.7}
\end{equation*}
$$

where the subscripts $i$ and $j(i, j=1,2)$ are the row and the column numbers. The coefficients, $k_{i j}^{(e)}$, may be interpreted as the force required at node $i$ to produce a unit displacement at node $j$ while all the other nodes are fixed.

### 1.4.2 Heat Flow

Uniform heat flow through the thickness of a domain whose in-plane dimensions are long in comparison to its thickness can be considered as a one-dimensional analysis. The cross section of such a domain is shown in Fig. 1.6. In accordance with Fourier's Law, the rate of heat flow per unit area in the $x$-direction can be written as

$$
\begin{equation*}
q=-k A \frac{d \theta}{d x} \tag{1.8}
\end{equation*}
$$



Fig. 1.6 One-dimensional heat flow.
where $A$ is the area normal to the heat flow, $\theta$ is the temperature, and $k$ is the coefficient of thermal conductivity. For constant $k$, Eq. (1.8) can be rewritten as

$$
\begin{equation*}
q=-k A \frac{\Delta \theta}{L} \tag{1.9}
\end{equation*}
$$

in which $\Delta \theta=\theta_{2}-\theta_{1}$ denotes the temperature drop across the thickness denoted by $L$ of the domain.

As illustrated in Fig. 1.6, the nodal flux (heat flow entering a node) at Node 1 becomes

$$
\begin{equation*}
q_{1}=\frac{k A}{L}\left(\theta_{1}-\theta_{2}\right) \tag{1.10}
\end{equation*}
$$

The balance of the heat flux requires that

$$
\begin{equation*}
q_{2}=-q_{1} \tag{1.11}
\end{equation*}
$$

which yields

$$
\begin{equation*}
q_{2}=-\frac{k A}{L}\left(\theta_{1}-\theta_{2}\right) \tag{1.12}
\end{equation*}
$$

Combining Eq. (1.10) and (1.12) and rewriting the resulting equations in matrix form yield

$$
\frac{k A}{L}\left[\begin{array}{cc}
1 & -1  \tag{1.13}\\
-1 & 1
\end{array}\right]\left\{\begin{array}{l}
\theta_{1} \\
\theta_{2}
\end{array}\right\}=\left\{\begin{array}{l}
q_{1} \\
q_{2}
\end{array}\right\} \quad \text { or } \quad \mathbf{k}^{(e)} \boldsymbol{\theta}^{(e)}=\mathbf{q}^{(e)}
$$

in which $\theta^{(e)}$ is the vector of nodal unknowns representing temperature and $\mathbf{k}^{(e)}$ and $\mathbf{q}^{(e)}$ are referred to as the element characteristic matrix and element right-hand-side vector, respectively.

### 1.4.3 Assembly of the Global System of Equations

Modeling an engineering problem with finite elements requires the assembly of element characteristic (stiffness) matrices and element right-hand-side (force) vectors, leading to the global system of equations

$$
\begin{equation*}
\mathbf{K u}=\mathbf{F} \tag{1.14}
\end{equation*}
$$

in which $\mathbf{K}$ is the assembly of element characteristic matrices, referred to as the global system matrix and $\mathbf{F}$ is the assembly of element right-handside vectors, referred to as the global right-hand-side (force) vector. The vector of nodal unknowns is represented by $\mathbf{u}$.

The global system matrix, $\mathbf{K}$, can be obtained from the "expanded" element coefficient matrices, $\mathbf{k}^{(e)}$, by summation in the form

$$
\begin{equation*}
\mathbf{K}=\sum_{e=1}^{E} \mathbf{k}^{(e)} \tag{1.15}
\end{equation*}
$$

in which the parameter $E$ denotes the total number of elements. The "expanded" element characteristic matrices are the same size as the global system matrix but have rows and columns of zeros corresponding to the nodes not associated with element (e). The size of the global system matrix is dictated by the highest number among the global node numbers.

Similarly, the global right-hand-side vector, $\mathbf{F}$, can be obtained from the "expanded" element coefficient vectors, $\mathbf{f}^{(e)}$, by summation in the form

$$
\begin{equation*}
\mathbf{F}=\sum_{e=1}^{E} \mathbf{f}^{(e)} \tag{1.16}
\end{equation*}
$$

The "expanded" element right-hand-side vectors are the same size as the global right-hand-side vector but have rows of zeros corresponding to the nodes not associated with element (e). The size of the global right-hand-side vector is also dictated by the highest number among the global node numbers.

The explicit steps in the construction of the global system matrix and the global right-hand-side-vector are explained by considering the system of linear springs shown in Fig. 1.7. Associated with element (e), the element equations for a spring given by Eq. (1.6) are rewritten as

$$
\left[\begin{array}{ll}
k_{11}^{(e)} & k_{12}^{(e)}  \tag{1.17}\\
k_{21}^{(e)} & k_{22}^{(e)}
\end{array}\right]\left\{\begin{array}{c}
u_{1}^{(e)} \\
u_{2}^{(e)}
\end{array}\right\}=\left\{\begin{array}{c}
f_{1}^{(e)} \\
f_{2}^{(e)}
\end{array}\right\}
$$

in which $k_{11}^{(e)}=k_{22}^{(e)}=k^{(e)}$ and $k_{12}^{(e)}=k_{21}^{(e)}=-k^{(e)}$. The subscripts used in Eq. (1.17) correspond to Node 1 and Node 2, the local node numbers of element (e). The global node numbers specifying the connectivity among the elements for this system of springs is shown in Fig. 1.7, and the connectivity information is tabulated in Table 1.3.


(3)

Fig. 1.7 System of linear springs (top) and corresponding FEA model (bottom).

Table 1.3 Table of connectivity.

| Element <br> Number | Local Node <br> Numbering | Global Node <br> Numbering |
| :---: | :---: | :---: |
| 1 | 1 | 1 |
| 2 | 2 | 2 |
| 2 | 1 | 2 |
| 3 | 2 | 3 |
|  | 1 | 2 |
| 4 | 2 | 3 |
|  | 1 | 3 |

In accordance with Eq. (1.15), the size of the global system matrix is $(4 \times 4)$ and the specific contribution from each element is captured as

Element 1: $\left[\begin{array}{cc}1 & 2 \\ k_{11}^{(1)} & k_{12}^{(1)} \\ k_{21}^{(1)} & k_{22}^{(1)}\end{array}\right]\left[\begin{array}{cccc|c}k_{11}^{(1)} & k_{12}^{(1)} & 0 & 0 \\ k_{21}^{(1)} & k_{22}^{(1)} & 0 & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 2 \\ 2\end{array} \Rightarrow \mathbf{k}^{(1)}\right.$


Element 3: \(\left[\begin{array}{cc}2 \& 3 <br>
k_{11}^{(3)} \& k_{12}^{(3)} <br>

k_{21}^{(3)} \& k_{22}^{(3)}\end{array}\right]\)| 2 |
| :---: |
| 3 |\(\Rightarrow\left[\begin{array}{cccc|c}0 \& 0 \& 0 \& 0 \& 1 <br>

0 \& k_{11}^{(3)} \& k_{12}^{(3)} \& 0 \& 2 <br>
0 \& k_{21}^{(3)} \& k_{22}^{(3)} \& 0 \& 3 <br>
0 \& 0 \& 0 \& 0\end{array}\right]=\mathbf{k}^{(3)}\)

Element 4: $\left.\left[\begin{array}{cc}3 & 4 \\ k_{11}^{(4)} & k_{12}^{(4)} \\ k_{21}^{(4)} & k_{22}^{(4)}\end{array}\right]\left[\begin{array}{l}3 \\ 4\end{array}\right]\left[\begin{array}{cccc}0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & k_{11}^{(4)} & k_{12}^{(4)} \\ 0 & 0 & k_{21}^{(4)} & k_{22}^{(4)}\end{array}\right] \begin{array}{ll}1 \\ 2 \\ 3\end{array}\right] \mathbf{k}^{(4)}$
Performing their assembly leads to

$$
\begin{equation*}
\mathbf{K}=\sum_{e=1}^{4} \mathbf{k}^{(e)}=\mathbf{k}^{(1)}+\mathbf{k}^{(2)}+\mathbf{k}^{(3)}+\mathbf{k}^{(4)} \tag{1.22}
\end{equation*}
$$

$$
\mathbf{K}=\left[\begin{array}{cccc}
k_{11}^{(1)} & k_{12}^{(1)} & 0 & 0  \tag{1.23}\\
k_{21}^{(1)} & \left(k_{22}^{(1)}+k_{11}^{(2)}+k_{11}^{(3)}\right) & \left(k_{12}^{(2)}+k_{12}^{(3)}\right) & 0 \\
0 & \left(k_{21}^{(2)}+k_{21}^{(3)}\right) & \left(k_{22}^{(2)}+k_{22}^{(3)}+k_{11}^{(4)}\right) & k_{12}^{(4)} \\
0 & 0 & k_{21}^{(4)} & k_{22}^{(4)}
\end{array}\right]
$$

In accordance with Eq. (1.16), the size of the global right-hand-side vector is $(4 \times 1)$ and the specific contribution from each element is captured as

Element 2: $\quad\left\{\begin{array}{c}f_{1}^{(2)} \\ f_{2}^{(2)}\end{array}\right] \begin{aligned} & 2 \\ & 3\end{aligned} \Rightarrow\left\{\begin{array}{c|c}0 & {[1} \\ f_{1}^{(2)} & 2 \\ f_{2}^{(2)} & 2 \\ 0 & \boxed{3} \\ \hline\end{array}\right] \mathbf{f}^{(2)}$

Similarly, performing their assembly leads to

$$
\begin{equation*}
\mathbf{F}=\sum_{e=1}^{4} \mathbf{f}^{(e)}=\mathbf{f}^{(1)}+\mathbf{f}^{(2)}+\mathbf{f}^{(3)}+\mathbf{f}^{(4)} \tag{1.28}
\end{equation*}
$$

$$
\mathbf{F}=\left\{\begin{array}{l}
f_{1}  \tag{1.29}\\
f_{2} \\
f_{3} \\
f_{4}
\end{array}\right\}=\left\{\begin{array}{c}
f_{1}^{(1)} \\
f_{2}^{(1)}+f_{1}^{(2)}+f_{1}^{(3)} \\
f_{2}^{(2)}+f_{2}^{(3)}+f_{1}^{(4)} \\
f_{2}^{(4)}
\end{array}\right\}
$$

Consistent with the assembly of the global system matrix and the global right-hand-side vector, the vector of unknowns, $\mathbf{u}$, becomes

$$
\mathbf{u}=\left\{\begin{array}{l}
u_{1}  \tag{1.30}\\
u_{2} \\
u_{3} \\
u_{4}
\end{array}\right\}=\left\{\begin{array}{c}
u_{1}^{(1)} \\
u_{2}^{(1)}=u_{1}^{(2)}=u_{1}^{(3)} \\
u_{2}^{(2)}=u_{2}^{(3)}=u_{1}^{(4)} \\
u_{2}^{(4)}
\end{array}\right\}
$$

### 1.4.4 Solution of the Global System of Equations

In order for the global system of equations to have a unique solution, the determinant of the global system matrix must be nonzero. However, an examination of the global system matrix reveals that one of its eigenvalues is zero, thus resulting in a zero determinant or singular matrix. Therefore, the solution is not unique. The eigenvector corresponding to the zero eigenvalue represents the translational mode, and the remaining nonzero eigenvalues represent all of the deformation modes.

For the specific values of $k_{11}^{(e)}=k_{22}^{(e)}=k^{(e)}$ and $k_{12}^{(e)}=k_{21}^{(e)}=-k^{(e)}$, the global system matrix becomes

$$
\mathbf{K}=k^{(e)}\left[\begin{array}{cccc}
1 & -1 & 0 & 0  \tag{1.31}\\
-1 & 3 & -2 & 0 \\
0 & -2 & 3 & -1 \\
0 & 0 & -1 & 1
\end{array}\right]
$$

with its eigenvalues $\lambda_{1}=0, \lambda_{2}=2, \lambda_{3}=3-\sqrt{5}$, and $\lambda_{4}=3+\sqrt{5}$. The corresponding eigenvectors are

$$
\mathbf{u}^{(1)}=\left\{\begin{array}{l}
1  \tag{1.32}\\
1 \\
1 \\
1
\end{array}\right\}, \quad \mathbf{u}^{(2)}=\left\{\begin{array}{c}
1 \\
-1 \\
-1 \\
1
\end{array}\right\}, \quad \mathbf{u}^{(3)}=\left\{\begin{array}{c}
-1 \\
2-\sqrt{5} \\
-2+\sqrt{5} \\
1
\end{array}\right\}, \quad \mathbf{u}^{(4)}=\left\{\begin{array}{c}
-1 \\
2+\sqrt{5} \\
-2-\sqrt{5} \\
1
\end{array}\right\}
$$

Each of these eigenvectors represents a possible solution mode. The contribution of each solution mode is illustrated in Fig. 1.8.

In order for the global system of equations to have a unique solution, the global system matrix is rendered nonsingular by eliminating the zero eigenvalue. This is achieved by introducing a boundary condition so as to suppress the translational mode of the solution corresponding to the zero eigenvalue.


Fig. 1.8 Possible solution modes for the system of linear springs.

### 1.4.5 Boundary Conditions

As shown in Fig. 1.7, Node 1 is restrained from displacement. This constraint is satisfied by imposing the boundary condition of $u_{1}=0$. Either the nodal displacements, $u_{i}$, or the nodal forces, $f_{i}$, can be specified at a given node. It is physically impossible to specify both of them as known or as unknown. Therefore, the nodal force $f_{1}$ remains as one of the unknowns. The nodal displacements, $u_{2}, u_{3}$, and $u_{4}$ are treated as unknowns, and the corresponding nodal forces have values of $f_{2}=0, f_{3}=0$, and $f_{4}=F$.

These specified values are invoked into the global system of equations as

$$
k^{(e)}\left[\begin{array}{cccc}
1 & -1 & 0 & 0  \tag{1.33}\\
-1 & 3 & -2 & 0 \\
0 & -2 & 3 & -1 \\
0 & 0 & -1 & 1
\end{array}\right]\left\{\begin{array}{c}
u_{1}=0 \\
u_{2} \\
u_{3} \\
u_{4}
\end{array}\right\}=\left\{\begin{array}{c}
f_{1} \\
f_{2}=0 \\
f_{3}=0 \\
f_{4}=F
\end{array}\right\}
$$

leading to the following equations:

$$
k^{(e)}\left[\begin{array}{ccc}
3 & -2 & 0  \tag{1.34}\\
-2 & 3 & -1 \\
0 & -1 & 1
\end{array}\right]\left\{\begin{array}{l}
u_{2} \\
u_{3} \\
u_{4}
\end{array}\right\}=\left\{\begin{array}{l}
0 \\
0 \\
F
\end{array}\right\}
$$

and

$$
\begin{equation*}
-k^{(e)} u_{2}=f_{1} \tag{1.35}
\end{equation*}
$$

The coefficient matrix in Eq. (1.34) is no longer singular, and the solutions to these equations are obtained as

$$
\begin{equation*}
u_{2}=\frac{F}{k^{(e)}}, \quad u_{3}=\frac{3}{2} \frac{F}{k^{(e)}}, \quad u_{4}=\frac{5}{2} \frac{F}{k^{(e)}} \tag{1.36}
\end{equation*}
$$

and the unknown nodal force $f_{1}$ is determined as $f_{1}=-F$. The final physically acceptable solution mode is shown in Fig. 1.9.

There exist systematic approaches to assemble the global coefficient matrix while invoking the specified nodal values (Bathe and Wilson 1976; Bathe 1996). The specified nodal variables are eliminated in advance from the global system of equations prior to the solution.


Fig. 1.9 Physically acceptable solution mode for the system of linear springs.

## Chapter 2

## FUNDAMENTALS OF ANSYS

### 2.1 Useful Definitions

Before delving into the details of the procedures related to the ANSYS program, we define the following terms:

Jobname: A specific name to be used for the files created during an ANSYS session. This name can be assigned either before or after starting the ANSYS program.

Working Directory: A specific folder (directory) for ANSYS to store all of the files created during a session. It is possible to specify the Working Directory before or after starting ANSYS.

Interactive Mode: This is the most common mode of interaction between the user and the ANSYS program. It involves activation of a platform called Graphical User Interface (GUI), which is composed of menus, dialog boxes, push-buttons, and different windows. Interactive Mode is the recommended mode for beginner ANSYS users as it provides an excellent platform for learning. It is also highly effective for postprocessing.

Batch Mode: This is a method to use the ANSYS program without activating the GUI. It involves an Input File written in ANSYS Parametric Design Language (APDL), which allows the use of parameters and common programming features such as DO loops and IF statements. These capabilities make the Batch Mode a very powerful analysis tool. Another distinct advantage of the Batch Mode is realized when there is an error/mistake in the model generation. This type of problem can be fixed by modifying a small portion of the Input File and reading it again, saving the user a great deal of time.

Combined Mode: This is a combination of the Interactive and Batch Modes in which the user activates the GUI and reads the Input File. Typically, this method allows the user to generate the model and obtain the solution using the Input File while reviewing the results using the

Postprocessor within the GUI. This method combines the salient advantages of the Interactive and Batch Modes.

### 2.2 Before an ANSYS Session

The construction of solutions to engineering problems using FEA requires either the development of a computer program based on the FEA formulation or the use of a commercially available general-purpose FEA program such as ANSYS. The ANSYS program is a powerful, multi-purpose analysis tool that can be used in a wide variety of engineering disciplines. Before using ANSYS to generate an FEA model of a physical system, the following questions should be answered based on engineering judgment and observations:

- What are the objectives of this analysis?
- Should the entire physical system be modeled, or just a portion?
- How much detail should be included in the model?
- How refined should the finite element mesh be?

In answering such questions, the computational expense should be balanced against the accuracy of the results. Therefore, the ANSYS finite element program can be employed in a correct and efficient way after considering the following:

- Type of problem.
- Time dependence.
- Nonlinearity.
- Modeling idealizations/simplifications.

Each of these topics is discussed in this section.

### 2.2.1 Analysis Discipline

The ANSYS program is capable of simulating problems in a wide range of engineering disciplines. However, this book focuses on the following disciplines:

Structural Analysis: Deformation, stress, and strain fields, as well as reaction forces in a solid body.

Thermal Analysis: Steady-state or time-dependent temperature field and heat flux in a solid body.

### 2.2.1.1 Structural Analysis

This analysis type addresses several different structural problems, for example:

Static Analysis: The applied loads and support conditions of the solid body do not change with time. Nonlinear material and geometrical properties such as plasticity, contact, creep, etc., are available.

Modal Analysis: This option concerns natural frequencies and modal shapes of a structure.

Harmonic Analysis: The response of a structure subjected to loads only exhibiting sinusoidal behavior in time.

Transient Dynamic: The response of a structure subjected to loads with arbitrary behavior in time.

Eigenvalue Buckling: This option concerns the buckling loads and buckling modes of a structure.

### 2.2.1.2 Thermal Analysis

This analysis type addresses several different thermal problems, for example:

Primary Heat Transfer: Steady-state or transient conduction, convection and radiation.

Phase Change: Melting or freezing.
Thermomechanical Analysis: Thermal analysis results are employed to compute displacement, stress, and strain fields due to differential thermal expansion.

### 2.2.1.3 Degrees of Freedom

The ANSYS solution for each of these analysis disciplines provides nodal values of the field variable. This primary unknown is called a degree of freedom (DOF). The degrees of freedom for these disciplines are presented in Table 2.1. The analysis discipline should be chosen based on the quantities of interest.

Table 2.1 Degrees of freedom for structural and thermal analysis disciplines.

| Discipline | Quantity | DOF |
| :--- | :--- | :---: |
| Structural | Displacement, stress, strain, <br> reaction forces | Displacement |
| Thermal | Temperature, flux | Temperature |

### 2.2.2 Time Dependence

The analysis with ANSYS should be time-dependent if:

- The solid body is subjected to time varying loads.
- The solid body has an initially specified temperature distribution.
- The body changes phase.


### 2.2.3 Nonlinearity

Most real-world physical phenomena exhibit nonlinear behavior. There are many situations in which assuming a linear behavior for the physical system might provide satisfactory results. On the other hand, there are circumstances or phenomena that might require a nonlinear solution. A nonlinear structural behavior may arise because of geometric and material nonlinearities, as well as a change in the boundary conditions and structural integrity. These nonlinearities are discussed briefly in the following subsections.

### 2.2.3.1 Geometric Nonlinearity

There are two main types of geometric nonlinearity:
Large deflection and rotation: If the structure undergoes large displacements compared to its smallest dimension and rotations to such an extent that its original dimensions and position, as well as the loading direction, change significantly, the large deflection and rotation analysis becomes necessary. For example, a fishing rod with a low lateral stiffness under a lateral load experiences large deflections and rotations.

Stress stiffening: When the stress in one direction affects the stiffness in another direction, stress stiffening occurs. Typically, a structure that has little or no stiffness in compression while having considerable stiffness in tension exhibits this behavior. Cables, membranes, or spinning structures exhibit stress stiffening.

### 2.2.3.2 Material Nonlinearity

A typical nonlinear stress-strain curve is given in Fig. 2.1. A linear material response is a good approximation if the material exhibits a nearly linear stress-strain curve up to a proportional limit and the loading is in a manner that does not create stresses higher than the yield stress anywhere in the body.


Fig. 2.1 Non-linear material response.

Nonlinear material behavior in ANSYS is characterized as:
Plasticity: Permanent, time-independent deformation.
Creep: Permanent, time-dependent deformation.
Nonlinear Elastic: Nonlinear stress-strain curve; upon unloading, the structure returns back to its original state-no permanent deformations.

Viscoelasticity: Time-dependent deformation under constant load. Full recovery upon unloading.

Hyperelasticity: Rubber-like materials.

### 2.2.3.3 Changing-status Nonlinearity

Many common structural features exhibit nonlinear behavior that is status dependent. When the status of the physical system changes, its stiffness shifts abruptly. The ANSYS program offers solutions to such phenomena through the use of nonlinear contact elements and birth and death options. This type of behavior is common in modeling manufacturing processes such as that of a shrink-fit (Fig. 2.2).

### 2.2.4 Practical Modeling Considerations

In order to reduce computational time, minor details that do not influence the results should not be included in the FE model. Minor details can also be ignored in order to render the geometry symmetric, which leads to a reduced FE model. However, in certain structures, "small" details such as fillets or holes may be the areas of maximum stress, which might prove to be extremely important in the analysis and design. Engineering judgment is essential to balance the possible gain in computational cost against the loss of accuracy.


Fig. 2.2 Element birth and death used in a manufacturing problem.

### 2.2.4.1 Symmetry Conditions

If the physical system under consideration exhibits symmetry in geometry, material properties, and loading, then it is computationally advantageous to model only a representative portion. If the symmetry observations are to be included in the model generation, the physical system must exhibit symmetry in all of the following:

- Geometry.
- Material properties.
- Loading.
- Degree of freedom constraints.

Different types of symmetry are:

- Axisymmetry.
- Rotational symmetry.
- Planar or reflective symmetry.
- Repetitive or translational symmetry.

Examples for each of the symmetry types are shown in Fig. 2.3. Each of these symmetry types is discussed below.

Axisymmetry: As illustrated in Fig. 2.4, axisymmetry is the symmetry about a central axis, as exhibited by structures such as light bulbs, straight pipes, cones, circular plates, and domes.

Rotational Symmetry: A structure possesses rotational symmetry when it is made up of repeated segments arranged about a central axis. An example is a turbine rotor (see Fig. 2.5).


Fig. 2.3 Types of symmetry conditions (from left to right): axisymmetry, rotational, reflective/planar, and repetitive/translational.


Fig. 2.4 Different views of a 3-D body with axisymmetry and its cross section (far right).


Fig. 2.5 Different views of a 3-D body with rotational symmetry.
Planar or Reflective Symmetry When one-half of a structure is a mirror image of the other half, planar or reflective symmetry exists, as shown in Fig. 2.6. In this case, the plane of symmetry is located on the surface of the mirror.

Repetitive or Translational Symmetry Repetitive or translational symmetry exists when a structure is made up of repeated segments lined up in a row, such as a long pipe with evenly spaced cooling fins, as shown in Fig. 2.7.

Symmetry in Material Properties, Loading, Displacements Once symmetry in geometry is observed, the same symmetry plane or axis should also be valid for the material properties, loading (forces, pressure, etc.), and


Fig. 2.6 Different views of a 3-D body with reflective/planar symmetry.


Fig. 2.7 A 3-D body with repetitive/translational symmetry.
constraints. For example, a homogeneous and isotropic square plate with a hole at the center under horizontal tensile loading (Fig. 2.8) has octant ( $1 / 8^{\text {dh }}$ ) symmetry in both geometry and material with respect to horizontal, vertical, and both diagonal axes. However, the loading is symmetric with respect to horizontal and vertical axes only. Therefore, a quarter of the structure is required in the construction of the solution.

If the applied loading varies in the vertical direction, as shown in Fig. 2.9, the loading becomes symmetric with respect to the vertical axis only. Although the geometry exhibits octant symmetry, half-symmetry is necessary in order to construct the solution.


Fig. 2.8 Example of quarter-symmetry.


Fig. 2.9 Example of half-symmetry with respect to vertical axis.
A similar plate, this time composed of two dissimilar materials is shown in Fig. 2.10. The loading condition allows for quarter-symmetry; however, the material properties are symmetric with respect to the horizontal axis only. Therefore, it is limited to half-symmetry. If this plate is subjected to a horizontal tensile load varying in the vertical direction, as shown in Fig. 2.11 , no symmetry condition is present.

Since a structure may exhibit symmetry in one or more of the aforementioned categories, one should try to find the smallest possible segment of the structure that would represent the entire structure. If the physical system exhibits symmetry in geometry, material properties, loading, and displacement constraints, it is computationally advantageous to use symmetry in the analysis. Typically, the use of symmetry produces better results as it leads to a finer, more detailed model than would otherwise be possible.


Fig. 2.10 Example of half-symmetry with respect to horizontal axis.


Fig. 2.11 Example of no symmetry.

A three-dimensional finite element mesh of the structure shown in Fig. 2.12 contains 18,739 tetrahedral elements with 5,014 nodes. However, the twodimensional mesh of the cross section necessary for the axisymmetric analysis has 372 quadrilateral elements and 447 nodes. The use of symmetry in this case reduces the CPU time required for the solution while delivering the same level of accuracy in the results.

### 2.2.4.2 Mesh Density

In general, a large number of elements provide a better approximation of the solution. However, in some cases, an excessive number of elements may increase the round-off error. Therefore, it is important that the mesh is adequately fine or coarse in the appropriate regions. How fine or coarse the mesh should be in such regions is another important question. Unfortunately, definitive answers to the questions about mesh refinement are not available since it is completely dependent on the specific physical system considered. However, there are some techniques that might be helpful in answering these questions:

Adaptive Meshing: The generated mesh is required to meet acceptable energy error estimate criteria. The user provides the "acceptable" error level information. This type of meshing is available only for linear static structural analysis and steady-state thermal analysis.


Fig. 2.12 Three-dimensional mesh of a structure (left) and 2-D mesh of the same structure (right) using axisymmetry.

Mesh Refinement Test Within ANSYS: An analysis with an initial mesh is performed first and then reanalyzed by using twice as many elements. The two solutions are compared. If the results are close to each other, the initial mesh configuration is considered to be adequate. If there are substantial differences between the two, the analysis should continue with a more-refined mesh and a subsequent comparison until convergence is established.

Submodeling: If the mesh refinement test yields nearly identical results for most regions and substantial differences in only a portion of the model, the built-in "submodeling" feature of ANSYS should be employed for localized mesh refinement. This feature is described in Chap. 11.

### 2.3 Organization of ANSYS Software

There are two primary levels in the ANSYS program, as shown in Fig. 2.13:
Begin Level: Gateway into and out of ANSYS and platform to utilize some global controls such as changing the jobname, etc.

Processor Level: This level contains the processors (preprocessor, solution, postprocessor, etc.) that are used to conduct finite element analyses.


Fig. 2.13 Schematic of ANSYS levels.
The user is in the Begin Level upon entering the ANSYS program. One can proceed to the Processor Level by clicking the mouse on one of the processor selections in the ANSYS Main Menu.

### 2.4 ANSYS Analysis Approach

There are three main steps in a typical ANSYS analysis:

- Model generation:
- Simplifications, idealizations.
- Define materials/material properties.
- Generate finite element model (mesh).
- Solution:
- Specify boundary conditions.
- Obtain the solution.
- Review results:
- Plot/list results.
- Check for validity.

Each of these steps corresponds to a specific processor or processors within the Processor Level in ANSYS. In particular, model generation is done in the Preprocessor and application of loads and the solution is performed in the Solution Processor. Finally, the results are viewed in the General Postprocessor and Time History Postprocessor for steady-state (static) and transient (time-dependent) problems, respectively. There are several other processors within the ANSYS program. These mostly concern optimizationand probabilistic-type problems. The most commonly used processors are described in the following subsections.

### 2.4.1 ANSYS Preprocessor

Model generation is conducted in this processor, which involves material definition, creation of a solid model, and, finally, meshing. Important tasks within this processor are:

- Specify element type.
- Define real constants (if required by the element type).
- Define material properties.
- Create the model geometry.
- Generate the mesh.

Although the boundary conditions can also be specified in this processor, it is usually done in the Solution Processor.

### 2.4.2 ANSYS Solution Processor

This processor is used for obtaining the solution for the finite element model that is generated within the Preprocessor. Important tasks within this processor are:

- Define analysis type and analysis options.
- Specify boundary conditions.
- Obtain solution.


### 2.4.3 ANSYS General Postprocessor

In this processor, the results at a specific time (if the analysis type is transient) over the entire or a portion of the model are reviewed. This includes the plotting of contours, vector displays, deformed shapes, and listings of the results in tabular format.

### 2.4.4 ANSYS Time History Postprocessor

This processor is used to review results at specific points in time (if the analysis type is transient). Similar to the General Postprocessor, it provides graphical variations and tabular listings of results data as functions of time.

### 2.5 ANSYS File Structure

Several files are created during a typical ANSYS analysis. Some of these files are in ASCII format while the others are binary. Brief descriptions of common file types are given below.

### 2.5.1 Database File

During a typical ANSYS analysis, input and output data reside in memory until they are saved in a Database File, which is saved in the Working Directory. The syntax for the name of the Database File is jobname.db. This binary file includes the element type, material properties, geometry (solid model), mesh (nodal coordinates and element connectivity), and the results if a solution is obtained. Once the Database File is saved, the user can resume from this file at any time. There are three distinct ways to save and resume the Database File:

- Use the Utility Menu.
- Click on SAVE_DB or RESUM_DB button on the ANSYS Toolbar.
- Issue the command SAVE or resume in the Input Field.


### 2.5.2 Log File

The Log File is an ASCII file, which is created (or resumed) immediately upon entering ANSYS. Every action taken by the user is stored sequentially in this file in command format (ANSYS Parametric Design Language (APDL)). The syntax for the name of the Log File, which is also saved in the Working Directory, is jobname.log. If jobname.log already exists in the Working Directory, ANSYS appends the newly executed actions instead of overwriting the file. The Log File can be utilized to:

- Understand how an analysis was performed by another user.
- Learn the command equivalents of the actions taken within ANSYS.


### 2.5.3 Error File

Similar to the Log File, the Error File is an ASCII file, which is created (or resumed) immediately upon entering ANSYS. This file captures all warning and error messages issued by ANSYS during a session. It is saved in the Working Directory with the following syntax for the name: jobname.err. If jobname.err already exists in the Working Directory, ANSYS appends the newly issued warning and error messages instead of overwriting the file. This file is particularly important when ANSYS issues several warning and error messages too quickly during an interactive session. The user can then consult the Error File to discover the exact cause(s) of each of the warnings or errors.

### 2.5.4 Results Files

The results of an ANSYS analysis are stored in a separate Results File. This file is a binary file and, depending upon the Analysis Type, the file's
extension takes a different form. The following syntax applies to the Results File name for the selected Analysis Type:

Structural analysis: jobname.rst
Thermal analysis: jobname.rth
Fluids analysis: jobname.rfl

### 2.6 Description of ANSYS Menus and Windows

When using the ANSYS program in Interactive Mode, the Graphical User Interface ( $G U I$ ) is activated. The $G U I$ has six distinct components:

Utility Menu: Contains functions that are available throughout the ANSYS session, such as file controls, selecting, graphic controls, and parameters. The ANSYS Help System is also accessible through this menu.

Main Menu: Contains the primary ANSYS functions organized by processors (Preprocessor, Solution, General Postprocessor, etc.).

Toolbar: Contains push-buttons for executing commonly used ANSYS commands and functions. Customized buttons can be created.

Input Field: Displays a text field for typing commands. All previously typed commands are stored in a pull-down menu for easy reference and access.

Graphics Window: Displays the graphical representation of the models/ meshes created within ANSYS. Also, the related results are reviewed in this window.

Output Window: Receives text output from the program. This window is usually positioned behind other windows and can be raised to the front when necessary.

Figure 2.14 shows a typical ANSYS GUI with each of the preceding components identified.


Fig. 2.14 Typical ANSYS GUI with separate components identified.

### 2.6.1 Utility Menu

The Utility Menu contains utility functions that are independent of ANSYS Levels (i.e., begin and processor levels), with some exceptions. The Utility Мепи contains ten items, each of which brings up a pull-down menu of subitems. Clicking the left mouse button on these subitems will result in one of the following:

- Bring up a submenu, indicated by the icon .
- Immediately execute a function.
- Bring up a dialog box, indicated by the icon ... .
- Bring up a picking menu, indicated by the icon + .

Brief descriptions of each of the menu items under the Utility Menu are given below.

File item under Utility Menu: Contains file- and database-related functions, such as clearing the database, reading an input file, saving the database to a file, or resuming a database from a file. This menu item can be used to exit the program.

Select item under Utility Menu: Includes functions that allow the user to select a subset of data and to create Components.

List item under Utility Menu：This menu item allows the user to list any data stored in the ANSYS database．Also，status information about different areas of the program and contents of files in the system are available．

Plot item under Utility Menu：This menu item allows the user to plot ANSYS entities such as keypoints，lines，areas，volumes，nodes，and elements．If a solution is obtained，results can also be plotted through this menu item．

PlotCtrls item under Utility Menu：Contains functions that control the view，style，and other characteristics of graphic displays．

WorkPlane item under Utility Menu：Use of WorkPlane offers great convenience for Solid Model generation．This menu item enables the user to toggle the Working Plane on and off，and to move，rotate，and maneuver it．Coordinate system operations are also performed under this menu item．

Parameters item under Utility Menu：Contains functions to define，edit， and delete scalar and array parameters．

Macro item under Utility Menu：This menu item allows the user to execute Macros and data blocks．Under this menu item，the user can also manipulate the push－buttons on the Toolbar．

MenuCtrls item under Utility Menu：Allows the user to format the menus，as well as manipulate the Toolbar．

Help item under Utility Menu：Brings up the ANSYS Help System．

## 2．6．2 Main Menu

The Main Menu contains main ANSYS functions and processors，such as the preprocessor，solution，and postprocessor．It has a tree structure，where menus and submenus can be expanded and collapsed．Similar to the Utility Menu，clicking the left mouse button on the Main Menu items results in one of the following：
－Expand or collapse the submenus attached to the menu item，indicated by icons $⿴ 囗 十 丌$ and $\boxminus$ ，respectively．
－Bring up a dialog box，indicated by the icon
－Bring up a picking menu，indicated by the icon $\gg$ ．

### 2.6.3 Toolbar

The Toolbar contains a set of push-buttons that execute frequently used ANSYS functions. When the user starts ANSYS, predefined push-buttons such as QUIT, SAVE_DB, and RESUM_DB appear in the toolbar. The user can create customized push-buttons and delete or edit the existing ones.

### 2.6.4 Input Field

This field allows the user to type in commands directly as opposed to the use of menu items. The Input Field consists of two main regions:

- Command entry box.
- History buffer.


### 2.6.5 Graphics Window

All ANSYS graphics are displayed in the Graphics Window. Also, the user performs all of the graphical "picking" in this window.

### 2.6.6 Output Window

All of the text output generated as a result of command responses, warnings, and errors appear in the Output Window. It is positioned behind the main ANSYS window, but can be raised to the front when necessary.

### 2.7 Using the ANSYS Help System

Information on ANSYS procedures, commands, and concepts can be found in the ANSYS Help System. The importance of knowing how to use the Help System cannot be overemphasized. It can be accessed within the Graphical User Interface (GUI) in three ways:

- By choosing the Help menu item under Utility Menu.
- By pressing the Help button within dialog boxes.
- By entering the help command directly in the Input Field.

The Help System is also available as a stand-alone program outside of ANSYS. The user can bring up the desired help topic by choosing it from the system's table of contents or index, through a word search, or by choosing a hypertext link. The Help System is built on the HTML platform in the form of web pages. As indicated in Fig. 2.15, there are three tabs on the left of the Help Window: Contents, Index, and Search. The help pages are displayed on the right side of the Help Window. Selected topics regarding the Help System are discussed in the following subsections.


Fig. 2.15 ANSYS Help System.

### 2.7.1 Help Contents

The first tab on the left side of the Help Window is the Contents Tab, as shown in Fig. 2.16. It is a collection of several different ANSYS Manuals containing thousands of pages. The Contents Tab is organized in a tree structure for easy navigation. It is recommended that beginner ANSYS users take the time to read the relevant chapters in each Manual. Throughout this book, the reader is referred to several specific chapters in these Manuals for a thorough understanding of the topics being discussed.

### 2.7.2 Help Index

The Index Tab (Fig. 2.17) is the second tab on the left side of the Help Window. Every single help page contained in the ANSYS Help System is exhaustively listed under this tab. It is useful for finding which help pages are available for a given topic. Upon typing the topic of interest, a list of help pages appears, giving the user a chance to browse for the most-relevant help page.

### 2.7.3 Search in Help

The user can perform a word search of the ANSYS Manuals through the Search Tab (Fig. 2.18), which is the third tab on the left side of the Help Window. As a result of an inquiry, a list of help pages containing the search word appears and the user can select which pages to display.


Fig. 2.16 ANSYS Help System with Contents tab activated.


Fig. 2.17 ANSYS Help System with Index tab activated.


Fig. 2.18 ANSYS Help System with Search tab activated.

### 2.7.4 Verification Manual

Although all of the ANSYS Manuals included under the Contents Tab are important sources of information, one particular Manual deserves special emphasis, the Verification Manual. The purpose of this Manual is to demonstrate the capabilities of ANSYS in solving fundamental engineering problems with analytical solutions. Another important feature of the Verification Manual is its suitability as an effective learning tool. There is a corresponding Input File for each of the verification problems included in this manual (in excess of 200). As mentioned earlier, the input files contain ANSYS commands to be executed sequentially when read from within ANSYS. Each of these commands corresponds to a specific action in the Interactive Mode. Once the verification problem that is the closest to the problem at hand is identified, the user can then study the corresponding Input File and learn the essential steps in solving the problem using ANSYS. The Verification Manual also serves as an excellent tool for learning to use ANSYS in Batch Mode.

## Chapter 3

## FUNDAMENTALS OF DISCRETIZATION

### 3.1 Local and Global Numbering

In solving an engineering problem with the finite element method (FEM), the domain is discretized by employing elements. The characteristics of the problem dictate the dimensionality of the problem, i.e., one, two, or three dimensional. A brief summary of the common element types utilized in a finite element analysis (FEA) is presented in Fig. 3.1. Once the domain of the problem is discretized by elements, a unique element number identifies each element and a unique node number identifies each node in the domain. As illustrated in Fig. 3.2, nodes are also numbered within each element, and are called local node numbers. The unique node numbering within the entire domain is called global node numbering. This is part of the computational procedure in FEA.

### 3.2 Approximation Functions

The variation of the field variable, $\phi^{(e)}$, over an element is approximated by an appropriate choice of functions, as illustrated in Fig. 3.3. The selection of these functions is the core of the finite element method. The approximation functions should be reliable in the sense that as the mesh becomes more refined, the approximate solution should converge to the exact solution monotonically. Oscillatory convergence is unreliable because it is possible to observe an increase in error with the refined mesh. Oscillatory and monotonic convergences are demonstrated in Fig. 3.4. Common approximation functions are usually polynomials since their differentiation and integration are rather straightforward compared to other functions.

In order to achieve a monotonically convergent solution, the polynomials chosen as approximation functions must satisfy four requirements:
$\longrightarrow \underset{\substack{\text { 2-node line } \\ \text { (linear) }}}{\stackrel{1}{0} \longrightarrow} \longrightarrow \underset{\substack{\text { 3-node line } \\ \text { (quadratic) }}}{\circ}$


3-node triangle
(linear)


4-node rectangle (linear)


6-node triangle (quadratic)


4-node quadrilateral (linear)



10-node triangle (cubic)


9 -node
rectangle (quadratic)


4-node tetrahedron (linear)


Fig. 3.1 Commonly used one-, two-, and three-dimensional finite elements.


Fig. 3.2 Element numbers, global node numbers, and local node numbers.


Fig. 3.3 Element approximation functions.


Fig. 3.4 Oscillatory and monotonic convergence of approximate solution.

Requirement 1. Continuous behavior of the approximation function within the element-no kinks or jumps.

Requirement 2. Compatibility along the common nodes, boundaries or surfaces between adjacent elements-no gaps between elements

The elements satisfying the continuity and compatibility requirements are called conformal elements (Fig. 3.5).

Requirement 3. Completeness, permitting rigid body motion of the element and ensuring (constant) variation of $\phi$ and its derivatives within the element.

The reason for this requirement is best illustrated by considering a cantilever beam under a concentrated load in the middle (Fig. 3.6). As a result of this loading, deformation occurs only to the left of the load. The section of the beam to the right of the load experiences only rigid-body translations and rotations (constant displacements and zero strain), i.e., no stresses and strains occur. Therefore, the element approximation functions must permit such behavior. Complete polynomials satisfy these requirements.

A complete polynomial of order $n$ in one dimension can be written in compact form as

$$
\begin{equation*}
P_{n}(x)=\sum_{k=1}^{n+1} \alpha_{k} x^{k-1} \tag{3.1}
\end{equation*}
$$

leading to complete polynomials of order 0,1 , and 2 (constant, linear, and quadratic) as

$$
\begin{align*}
& P_{0}(x)=\alpha_{1} \\
& P_{1}(x)=\alpha_{1}+\alpha_{2} x  \tag{3.2}\\
& P_{2}(x)=\alpha_{1}+\alpha_{2} x+\alpha_{3} x^{2}
\end{align*}
$$

In two dimensions, the compact form for a complete polynomial of order $n$ can be written as

$$
\begin{equation*}
P_{n}(x)=\sum_{k=1}^{\frac{(n+1)(n+2)}{2}} \alpha_{k} x^{i} y^{j} \quad i+j \leq n \tag{3.3}
\end{equation*}
$$



Fig. 3.5 Compatibility of approximation functions.



Fig. 3.6 A cantilever beam loaded at the middle and its FEA model.

Constant, linear, and quadratic complete polynomials in two dimensions can be written as

$$
\begin{align*}
& P_{0}(x, y)=\alpha_{1} \\
& P_{1}(x, y)=\alpha_{1}+\alpha_{2} x+\alpha_{3} y  \tag{3.4}\\
& P_{2}(x, y)=\alpha_{1}+\alpha_{2} x+\alpha_{3} y+\alpha_{4} x^{2}+\alpha_{5} x y+\alpha_{6} y^{2}
\end{align*}
$$

The Pascal triangle shown in Fig. 3.7 is useful for including the appropriate terms to obtain complete approximating functions in any order.


Fig. 3.7 Pascal's triangle for complete polynomials.
The order of the polynomial as an approximation function is dictated by the total number of nodes in an element, i.e., the number of coefficients, $\alpha_{i}$, in the approximation function must be the same as the number of nodes in the element.

Requirement 4. Geometric isotropy for the same behavior in each direction.

Using complete polynomials satisfies this requirement of translation and rotation of the coordinate system. If the required degree of completeness does not provide a number of terms equal to the number of nodes, then this requirement can be satisfied by disregarding the non-symmetrical terms. In the case of a 4 -noded rectangular element, the first-order complete polynomial has 3 coefficients, one less than the number of nodes. In order to circumvent this deficiency, the order of the polynomial can be increased to "complete" in the second degree, having 6 coefficients, two more than the number of nodes. As a result, two of the additional higher-order terms, which are $\alpha_{4} x^{2}, \alpha_{5} x y$, and $\alpha_{6} y^{2}$, must be removed from the approximation function.

In order to satisfy the condition of geometric isotropy, only the term $\alpha_{5} x y$ is retained in the approximation function, leading to

$$
\begin{equation*}
P_{2}(x, y)=\alpha_{1}+\alpha_{2} x+\alpha_{3} y+\alpha_{4} x y \tag{3.5}
\end{equation*}
$$

Approximation functions satisfying these four requirements ensure monotonic convergence of the solution as the element sizes decrease.

The element is referred to as $C^{0}$ continuous when only the field variable (none of its derivatives) maintains continuity along its boundary. If the field variable and its $r^{\text {th }}$ derivative maintain continuity, the element is $C^{r}$ continuous. A more extensive discussion is given by Huebner et al. (2001).

### 3.3 Coordinate Systems

### 3.3.1 Generalized Coordinates

The coefficients of the approximation functions, $\alpha_{i}$, are referred to as the generalized coordinates. They are not identified with particular nodes. The generalized coordinates are independent parameters that specify the magnitude of the prescribed distribution of the field variable. They have no direct physical interpretation, but rather are linear combinations of the physical nodal degrees of freedom.

### 3.3.2 Global Coordinates

Global coordinates are convenient for specifying the location of each node, the orientation of each element, and the boundary conditions and loads for the entire domain. Also, the solution to the field variable is generally represented with respect to the global coordinates. However, approximation functions described in terms of the global coordinates are not convenient to use in the evaluation of integrals necessary for the construction of the element matrix.

### 3.3.3 Local Coordinates

A local coordinate system whose origin is located within the element is introduced in order to simplify the algebraic manipulations in the derivation of the element matrix. The use of natural coordinates in expressing the approximation functions is particularly advantageous because special integration formulas can often be employed to evaluate the integrals in the element matrix. Natural coordinates also play a crucial role in the development of elements with curved boundaries (discussed under isoparametric elements, Sec. 6.2.2.5).

### 3.3.4 Natural Coordinates

A local coordinate system that permits the specification of a point within the element by a dimensionless parameter whose absolute magnitude never exceeds unity is referred to as a natural coordinate system. Natural coordinates are dimensionless. They are defined with respect to the element rather than with reference to the global coordinates. Also, the natural coordinates are functions of the global coordinates in which the element is defined. As illustrated in Fig. 3.8, the basic purpose of the natural coordinate system is
to describe the location of a point inside an element in terms of coordinates associated with the nodes of the element.

### 3.3.4.1 Natural Coordinates in One Dimension

As shown in Fig. 3.9, within a one-dimensional element (line segment), defined by two nodes (one at each end), the location of a point $P$ denoted by $x$ (global coordinate) on the element can be expressed in terms of length or centroidal coordinates.

### 3.3.4.1.1 Length Coordinates

The location of point $\mathrm{P}, x$, is expressed as a linear combination of the global nodal coordinates, $x_{1}$ and $x_{2}$, and the length coordinates, $\xi_{1}$ and $\xi_{2}$, as

$$
\begin{equation*}
x=\xi_{1} x_{1}+\xi_{2} x_{2} \tag{3.6}
\end{equation*}
$$

As shown in Fig. 3.9, $\xi_{1}$ and $\xi_{2}$ are defined as the ratios of lengths $\xi_{1}=L_{1} / L$ and $\xi_{2}=L_{2} / L$, with $L$ representing the length of the line segment, $L=x_{2}-x_{1}$. Since $L=L_{1}+L_{2}, \xi_{1}$, and $\xi_{2}$ are not independent of each other and must satisfy the constraint relation

$$
\begin{equation*}
\xi_{1}+\xi_{2}=1 \tag{3.7}
\end{equation*}
$$

Solving for $\xi_{1}$ and $\xi_{2}$ via these equations written in matrix form as

$$
\left\{\begin{array}{l}
1  \tag{3.8}\\
x
\end{array}\right\}=\left[\begin{array}{cc}
1 & 1 \\
x_{1} & x_{2}
\end{array}\right]\left\{\begin{array}{l}
\xi_{1} \\
\xi_{2}
\end{array}\right\}
$$

results in

$$
\begin{equation*}
\xi_{1}=\frac{x_{2}-x}{x_{2}-x_{1}} \quad \text { and } \quad \xi_{2}=\frac{x-x_{1}}{x_{2}-x_{1}} \tag{3.9}
\end{equation*}
$$

Such coordinates, whose behavior is shown in Fig. 3.10, have the property that one particular coordinate has a unit value at one node of the element and a zero value at the other node(s), i.e., $\xi_{1}\left(x_{1}\right)=1$ and $\xi_{1}\left(x_{2}\right)=0$, and $\xi_{2}\left(x_{1}\right)=0$ and $\xi_{2}\left(x_{2}\right)=1$.


Fig. 3.8 Local and global coordinates in two dimensions.


Fig. 3.9 Length coordinates in one dimension.


Fig. 3.10 Variation of length coordinates within the element.

### 3.3.4.1.2 Centroidal Coordinates

As shown in Fig. 3.11, $x$ (the location of point P ) with respect to a local coordinate system, $r$, located at the centroid of the line element becomes

$$
\begin{equation*}
x=r+x_{1}+\frac{L}{2} \tag{3.10}
\end{equation*}
$$

The local coordinate $r$ is normalized in the form $\xi=r /(L / 2)$ in order to achieve a dimensionless coordinate, $\xi$, and to ensure that its range never exceeds unity. Thus, the location of the point P becomes

$$
\begin{equation*}
x=\frac{L}{2} \xi+x_{1}+\frac{L}{2} \tag{3.11}
\end{equation*}
$$

Substituting for $L\left(L=x_{2}-x_{1}\right)$ and rearranging terms leads to

$$
\begin{equation*}
x=\frac{1}{2}(1-\xi) x_{1}+\frac{1}{2}(1+\xi) x_{2} \tag{3.12}
\end{equation*}
$$

or

$$
\begin{equation*}
x=\sum_{i=1}^{2} N_{i} x_{i} \tag{3.13}
\end{equation*}
$$

with $N_{1}=(1-\xi) / 2$ and $N_{2}=(1+\xi) / 2$. As shown in Fig. 3.12, $N_{1}(-1)=1$ and $N_{1}(1)=0$, and $N_{2}(-1)=0$ and $N_{2}(1)=1$.

### 3.3.4.2 Natural Coordinates in Two Dimensions

### 3.3.4.2.1 Area Coordinates

As shown in Fig. 3.13, within a two-dimensional element (triangular area) defined by three nodes, one at each apex, the location of a point $P$, denoted by ( $x, y$ ) (global coordinates), on the element can be expressed as linear combinations of the global nodal coordinates, $\left(x_{1}, y_{1}\right),\left(x_{2}, y_{2}\right)$, and $\left(x_{3}, y_{3}\right)$, and the area coordinates, $\xi_{1}, \xi_{2}$, and $\xi_{3}$, as

$$
\begin{align*}
& x=\xi_{1} x_{1}+\xi_{2} x_{2}+\xi_{3} x_{3} \\
& y=\xi_{1} y_{1}+\xi_{2} y_{2}+\xi_{3} y_{3} \tag{3.14}
\end{align*}
$$



Fig. 3.11 Centroidal coordinates in one dimension.


Fig. 3.12 Variation of centroidal coordinates within the element.


Fig. 3.13 Definition of area coordinates in a triangular element.

As illustrated in Fig. $3.13, \xi_{1}, \xi_{2}$, and $\xi_{3}$ are defined as the ratios of areas $\xi_{1}=A_{1} / A, \xi_{2}=A_{2} / A$, and $\xi_{3}=A_{3} / A$, with $A$ representing the area of the triangle. Since $A_{1}+A_{2}+A_{3}=1, \xi_{1}, \xi_{2}$, and $\xi_{3}$ are not independent of each other and must satisfy the constraint relation

$$
\begin{equation*}
\xi_{1}+\xi_{2}+\xi_{3}=1 \tag{3.15}
\end{equation*}
$$

Solving for $\xi_{1}, \xi_{2}$, and $\xi_{3}$ via Eq. (3.14) and (3.15) written in matrix form as

$$
\left\{\begin{array}{l}
1  \tag{3.16}\\
x \\
y
\end{array}\right\}=\left[\begin{array}{ccc}
1 & 1 & 1 \\
x_{1} & x_{2} & x_{3} \\
y_{1} & y_{2} & y_{3}
\end{array}\right]\left\{\begin{array}{l}
\xi_{1} \\
\xi_{2} \\
\xi_{3}
\end{array}\right\}
$$

results in

$$
\left\{\begin{array}{l}
\xi_{1}  \tag{3.17}\\
\xi_{2} \\
\xi_{3}
\end{array}\right\}=\frac{1}{2 A}\left[\begin{array}{ccc}
\left(x_{2} y_{3}-x_{3} y_{2}\right) & y_{23} & x_{32} \\
\left(x_{3} y_{1}-x_{1} y_{3}\right) & y_{31} & x_{13} \\
\left(x_{1} y_{2}-x_{2} y_{1}\right) & y_{12} & x_{21}
\end{array}\right]\left\{\begin{array}{l}
1 \\
x \\
y
\end{array}\right\}
$$

where $x_{m n}=x_{m}-x_{n}, y_{m n}=y_{m}-y_{n}$, and

$$
2 A=\left|\begin{array}{ccc}
1 & 1 & 1  \tag{3.18}\\
x_{1} & x_{2} & x_{3} \\
y_{1} & y_{2} & y_{3}
\end{array}\right|
$$

As shown in Fig. 3.14, one particular area coordinate has a unit value at one node of the element and a zero value at the other node(s); $\xi_{i}\left(x_{j}\right)=\delta_{i j}$, where $\delta_{i j}=1$ for $i=j$ and $\delta_{i j}=0$ for $i \neq j$.

The exact evaluation of the area integrals over a triangle can be obtained by employing the expression

$$
\begin{equation*}
I=\int_{A} \xi_{1}^{m} \xi_{2}^{n} \xi_{3}^{\ell} d x d y=\frac{m!n!\ell!}{(m+n+\ell+2)!} 2 A \tag{3.19}
\end{equation*}
$$

### 3.3.4.2.2 Centroidal Coordinates

In the case of a two-dimensional element with a quadrilateral shape defined by four nodes, one at each corner, the location of a point $P$, denoted by $(x, y)$, on the element can be expressed with respect to the centroidal coor-


Fig. 3.14 Area coordinates within a triangular element.
dinate system $(\xi, \eta)$ whose origin coincides with the centroid of the quadrilateral area, as shown in Fig. 3.15. The relationship between ( $x, y$ ) and $(\xi, \eta)$ can be expressed as

$$
\begin{align*}
& x=a_{x}+b_{x} \xi+c_{x} \eta+d_{x} \xi \eta \\
& y=a_{y}+b_{y} \xi+c_{y} \eta+d_{y} \xi \eta \tag{3.20}
\end{align*}
$$

Also, these relations map a quadrilateral shape in global coordinates to a unit square in natural (centroidal) coordinates. Evaluation of these equations along $\eta=-1$ leads to

$$
\begin{align*}
& x=a_{x}+b_{x} \xi-c_{x}-d_{x} \xi \\
& y=a_{y}+b_{y} \xi-c_{y}-d_{y} \xi \tag{3.21}
\end{align*}
$$

Eliminating the coordinate $\xi$ from the resulting equations yields the linear relationship between the global coordinates

$$
\begin{equation*}
y=A+B x \tag{3.22}
\end{equation*}
$$

in which $A$ and $B$ are known explicitly. Considering the remaining sides of the square in the centroidal coordinates defined by the lines $\eta=1, \xi=1$, and $\xi=-1$ results in a straight-sided quadrilateral.


Fig. 3.15 Centroidal coordinates within a quadrilateral element.

Evaluation of $x$ at $\xi= \pm 1$ and $\eta= \pm 1$ (four corners) leads to

$$
\begin{align*}
& x_{1}=a_{x}-b_{x}-c_{x}+d_{x} \\
& x_{2}=a_{x}+b_{x}-c_{x}-d_{x}  \tag{3.23}\\
& x_{3}=a_{x}+b_{x}+c_{x}+d_{x} \\
& x_{4}=a_{x}-b_{x}+c_{x}-d_{x}
\end{align*}
$$

Solving for the coefficients $a_{x}, b_{x}, c_{x}$, and $d_{x}$, substituting back into Eq. (3.20), and collecting the terms multiplying $x_{i}$ gives

$$
\begin{align*}
x & =\frac{1}{4}(1-\xi)(1-\eta) x_{1}+\frac{1}{4}(1+\xi)(1-\eta) x_{2} \\
& +\frac{1}{4}(1+\xi)(1+\eta) x_{3}+\frac{1}{4}(1-\xi)(1+\eta) x_{4} \tag{3.24}
\end{align*}
$$

A similar operation performed on $y$ in Eq. (3.20) yields

$$
\begin{align*}
y & =\frac{1}{4}(1-\xi)(1-\eta) y_{1}+\frac{1}{4}(1+\xi)(1-\eta) y_{2} \\
& +\frac{1}{4}(1+\xi)(1+\eta) y_{3}+\frac{1}{4}(1-\xi)(1+\eta) y_{4} \tag{3.25}
\end{align*}
$$

Defining

$$
\begin{array}{ll}
N_{1}=\frac{1}{4}(1-\xi)(1-\eta) & N_{2}=\frac{1}{4}(1+\xi)(1-\eta)  \tag{3.26}\\
N_{3}=\frac{1}{4}(1+\xi)(1+\eta) & N_{4}=\frac{1}{4}(1-\xi)(1+\eta)
\end{array}
$$

allows Eq. (3.24) and (3.25) to be rewritten as

$$
\begin{equation*}
x=\sum_{i=1}^{4} N_{i}(\xi, \eta) x_{i} \quad \text { and } \quad y=\sum_{i=1}^{4} N_{i}(\xi, \eta) y_{i} \tag{3.27}
\end{equation*}
$$

Note that $N_{i}$ can be written in compact form as

$$
\begin{equation*}
N_{i}=\frac{1}{4}\left(1+\xi \xi_{i}\right)\left(1+\eta \eta_{i}\right) \tag{3.28}
\end{equation*}
$$

with $\xi_{i}$ and $\eta_{i}$ representing the coordinates of the corner nodes in the natural coordinate system. It is worth noting that $N_{i}\left(\xi_{j}, \eta_{j}\right)=\delta_{i j}$, where $\delta_{i j}=1$ for $i=j$ and $\delta_{i j}=0$ for $i \neq j$. The variations of $N_{i}$ within a quadrilateral element are given schematically in Fig. 3.16.

### 3.4 Shape Functions

Shape functions constitute the subset of element approximation functions. They cannot be chosen arbitrarily. As discussed in the previous section, the element approximation functions are chosen to be complete polynomials with unknown generalized coordinates. For a one-dimensional element with $m$ nodes as shown in Fig. 3.17, the element approximation function for the field variable, $\phi(x)$, is assumed as a polynomial of order $(m-1)$

$$
\begin{equation*}
\phi^{(e)}(x)=\alpha_{1}+\alpha_{2} x+\alpha_{3} x^{2}+\alpha_{4} x^{3}+\cdots+\alpha_{m-1} x^{m-2}+\alpha_{m} x^{m-1} \tag{3.29}
\end{equation*}
$$

or

$$
\begin{equation*}
\phi^{(e)}(x)=\mathbf{g}^{T} \boldsymbol{\alpha} \tag{3.30}
\end{equation*}
$$

where

$$
\mathbf{g}^{T}=\left\{\begin{array}{lllll}
1 & x & x^{2} & \cdots & x^{m-1} \tag{3.31}
\end{array}\right\}
$$

and

$$
\boldsymbol{\alpha}^{T}=\left\{\begin{array}{lllll}
\alpha_{1} & \alpha_{2} & \alpha_{3} & \cdots & \alpha_{m} \tag{3.32}
\end{array}\right\}
$$

Note that the number of generalized coordinates $\left(\alpha_{i}, i=1,2, \ldots, m\right)$ is equal to the number of nodes within the element.

$$
N_{1}=(1-\xi)(1-\eta) / 4 \quad N_{2}=(1-\xi)(1-\eta) / 4
$$



Fig. 3.16 Variation of $N_{i}$ within a quadrilateral.


Fig. 3.17 A one-dimensional element with $m$ nodes.

The field variable, $\phi^{(e)}(x)$, can also be expressed within the element through the use of its nodal values, $\phi_{i}(i=1, m)$, in the form

$$
\begin{align*}
& \phi^{(e)}(x)=N_{1} \phi_{1}+N_{2} \phi_{2}+N_{3} \phi_{3}+N_{4} \phi_{4}+\cdots  \tag{3.33}\\
& \quad+N_{m-1} \phi_{m-1}+N_{m} \phi_{m}
\end{align*}
$$

or

$$
\begin{equation*}
\phi^{(e)}(x)=\mathbf{N}^{T} \varphi \tag{3.34}
\end{equation*}
$$

where

$$
\mathbf{N}^{T}=\left\{\begin{array}{lllll}
N_{1} & N_{2} & N_{3} & \cdots & N_{m} \tag{3.35}
\end{array}\right\}
$$

and

$$
\varphi^{T}=\left\{\begin{array}{lllll}
\phi_{1} & \phi_{2} & \phi_{3} & \cdots & \phi_{m} \tag{3.36}
\end{array}\right\}
$$

in which $N_{i}(i=1, m)$ are referred to as shape functions. These functions are associated with node $i$ and must have a unit value at node $i$ and a zero value at all other nodes. Furthermore, they must have the same degree of polynomial variation as in the element approximation function.

The explicit form of the shape functions can be determined by solving for the generalized coordinates, $\alpha_{i}$, in terms of the nodal coordinates, $x_{i}$, and nodal values, $\phi_{i}(i=1,2, \ldots, m)$, through Eq. (3.29), and rearranging the resulting expressions in the form of Eq. (3.34). At each node, the field variable $\phi^{(e)}(x)$ is evaluated as

$$
\begin{gather*}
\phi_{1}=\alpha_{1}+\alpha_{2} x_{1}+\alpha_{3} x_{1}^{2}+\alpha_{4} x_{1}^{3}+\cdots+\alpha_{m-1} x_{1}^{m-2}+\alpha_{m} x_{1}^{m-1} \\
\phi_{2}=\alpha_{1}+\alpha_{2} x_{2}+\alpha_{3} x_{2}^{2}+\alpha_{4} x_{2}^{3}+\cdots+\alpha_{m-1} x_{2}^{m-2}+\alpha_{m} x_{2}^{m-1}  \tag{3.37}\\
\vdots \\
\phi_{m}=\alpha_{1}+\alpha_{2} x_{m}+\alpha_{3} x_{m}^{2}+\alpha_{4} x_{m}^{3}+\cdots+\alpha_{m-1} x_{m}^{m-2}+\alpha_{m} x_{m}^{m-1}
\end{gather*}
$$

or in matrix form

$$
\left\{\begin{array}{c}
\phi_{1}  \tag{3.38}\\
\phi_{2} \\
\phi_{3} \\
\vdots \\
\phi_{m}
\end{array}\right\}=\left[\begin{array}{ccccc}
1 & x_{1} & x_{1}^{2} & \cdots & x_{1}^{m-1} \\
1 & x_{2} & x_{2}^{2} & \cdots & x_{2}^{m-1} \\
1 & x_{3} & x_{3}^{2} & \cdots & x_{3}^{m-1} \\
\vdots & \vdots & \vdots & \ddots & \vdots \\
1 & x_{m} & x_{m}^{2} & \cdots & x_{m}^{m-1}
\end{array}\right]\left\{\begin{array}{c}
\alpha_{1} \\
\alpha_{2} \\
\alpha_{3} \\
\vdots \\
\alpha_{m}
\end{array}\right\} \text { or } \varphi=\mathbf{A} \boldsymbol{\alpha}
$$

Solving for the generalized coordinates in terms of nodal coordinates and nodal values of the field variable yields

$$
\begin{equation*}
\alpha=\mathrm{A}^{-1} \boldsymbol{\varphi} \tag{3.39}
\end{equation*}
$$

Substituting for the generalized coordinates in Eq. (3.30) results in

$$
\begin{equation*}
\phi^{(e)}(x)=\mathbf{g}^{T} \mathbf{A}^{-1} \boldsymbol{\varphi} \tag{3.40}
\end{equation*}
$$

Comparison of Eq. (3.40) and (3.34) leads to the explicit form of the shape functions $N_{i}$ as

$$
\begin{equation*}
\mathbf{N}^{T}=\mathbf{g}^{T} \mathbf{A}^{-1} \tag{3.41}
\end{equation*}
$$

This formulation illustrates the determination of the shape functions for a one-dimensional element; its extension to two dimensions is straightforward. The properties of shape functions are:

1. $N_{i}=1$ at node $i$ and $N_{i}=0$ at all other nodes.
2. $\sum_{i=1}^{m} N_{i}=1$.

### 3.4.1 Linear Line Element with Two Nodes

### 3.4.1.1 Global Coordinate

For a line element with two nodes, the field variable, $\phi^{(e)}$, is approximated by a linear function (refer to Fig. 3.18) in terms of the global coordinate, $x$, as

$$
\begin{equation*}
\phi^{(e)}(x)=\alpha_{1}+\alpha_{2} x \tag{3.42}
\end{equation*}
$$

This element approximation function ensures the inter-element continuity of only the field variable. The nodal values of the function are identified by $\phi_{1}$ and $\phi_{2}$.

Evaluation of the function at each node with coordinates $x_{1}$ and $x_{2}$ leads to

$$
\begin{equation*}
\phi_{1}=\alpha_{1}+\alpha_{2} x_{1} \quad \text { and } \quad \phi_{2}=\alpha_{1}+\alpha_{2} x_{2} \tag{3.43}
\end{equation*}
$$

Solving for $\alpha_{1}$ and $\alpha_{2}$ and substituting for them in the element approximation function results in

$$
\begin{equation*}
\phi^{(e)}(x)=N_{1}(x) \phi_{1}+N_{2}(x) \phi_{2} \tag{3.44}
\end{equation*}
$$

where $N_{1}=\left(x_{2}-x\right) /\left(x_{2}-x_{1}\right)$ and $N_{2}=\left(x-x_{1}\right) /\left(x_{2}-x_{1}\right)$. These functions, referred to as interpolation or shape functions, are the same as the length coordinates, $\xi_{1}$ and $\xi_{2}$, and they also vary linearly with $x$ (Fig. 3.19), as does the element approximation function. Because $N_{i}\left(x_{j}\right)=\delta_{i j}$, where $\delta_{i j}=1$ for $i=j$ and $\delta_{i j}=0$ for $i \neq j$,

$$
\begin{equation*}
1=\sum_{i=1}^{2} N_{i} \tag{3.45}
\end{equation*}
$$



Fig. 3.18 Linear approximation for the field variable $\phi$ within a line element.



Fig. 3.19 Variation of linear shape functions within a 1-D line element.

### 3.4.1.2 Centroidal Coordinate

For a line element with two nodes, the field variable, $\phi^{(e)}$, is approximated by a linear function in terms of the natural (centroidal) coordinate, $\xi$, as

$$
\begin{equation*}
\phi^{(e)}(\xi)=\alpha_{1}+\alpha_{2} \xi \tag{3.46}
\end{equation*}
$$

This element approximation function ensures the inter-element continuity of the field variable. The nodal values of the function are identified by $\phi_{1}$ and $\phi_{2}$. Evaluation of the function at each node with coordinates $\xi=-1$ and $\xi=1$ leads to

$$
\begin{equation*}
\phi_{1}=\alpha_{1}-\alpha_{2} \quad \text { and } \quad \phi_{2}=\alpha_{1}+\alpha_{2} \tag{3.47}
\end{equation*}
$$

Solving for $\alpha_{1}$ and $\alpha_{2}$ and substituting for them in the element approximation function results in

$$
\begin{equation*}
\phi^{(e)}(\xi)=N_{1}(\xi) \phi_{1}+N_{2}(\xi) \phi_{2} \tag{3.48}
\end{equation*}
$$

where $N_{1}(\xi)=(1-\xi) / 2$ and $N_{2}(\xi)=(1+\xi) / 2$. These functions, referred to as interpolation or shape functions, vary linearly with $\xi$ (Fig. 3.20), as in the case of the element approximation function.

Also, they have the property

$$
\begin{equation*}
1=\sum_{i=1}^{2} N_{i} \tag{3.49}
\end{equation*}
$$

because $N_{i}\left(\xi_{j}\right)=\delta_{i j}$, where $\delta_{i j}=1$ for $i=j$ and $\delta_{i j}=0$ for $i \neq j$.

### 3.4.2 Quadratic Line Element with Three Nodes: Centroidal Coordinate

For a line element with three nodes, the field variable, $\phi^{(e)}$, is approximated by a quadratic function (schematic given in Fig. 3.21) in terms of the natural (centroidal) coordinate, $\xi$, as

$$
\begin{equation*}
\phi^{(e)}(\xi)=\alpha_{1}+\alpha_{2} \xi+\alpha_{3} \xi^{2} \tag{3.50}
\end{equation*}
$$

in order to ensure the inter-element continuity of the field variable. The element nodes are identified as 1,2 , and 3 , with their nodal values as $\phi_{1}, \phi_{2}$, and $\phi_{3}$. The middle node is located at the center of the line element. Evaluation of the function at each node with coordinates $\xi=-1, \xi=0$, and $\xi=1$ leads to

$$
\begin{equation*}
\phi_{1}=\alpha_{1}-\alpha_{2}+\alpha_{3}, \quad \phi_{3}=\alpha_{1}, \quad \phi_{2}=\alpha_{1}+\alpha_{2}+\alpha_{3} \tag{3.51}
\end{equation*}
$$

Solving for $\alpha_{1}, \alpha_{2}$, and $\alpha_{3}$ and substituting for them in the element approximation function results in

$$
\begin{equation*}
\phi^{(e)}(\xi)=N_{1}(\xi) \phi_{1}+N_{2}(\xi) \phi_{2}+N_{3}(\xi) \phi_{3} \tag{3.52}
\end{equation*}
$$

where $\quad N_{1}(\xi)=\xi /[2(\xi-1)], \quad N_{2}(\xi)=\xi /[2(\xi+1)], \quad$ and $\quad N_{3}(\xi)=$ $-(\xi+1)(\xi-1)$. These functions, referred to as interpolation or shape functions, vary quadratically with $\xi$ (Fig. 3.22), as in the case of element approximation function.


Fig. 3.20 Variation of linear shape functions within a 1-D line element.


Fig. 3.21 Quadratic approximation for the field variable $\phi$ within a line element.



Fig. 3.22 Variation of quadratic shape functions within a 1-D line element.

Also, they have the property

$$
\begin{equation*}
1=\sum_{i=1}^{3} N_{i} \tag{3.53}
\end{equation*}
$$

because $N_{i}\left(\xi_{j}\right)=\delta_{i j}$, where $\delta_{i j}=1$ for $i=j$ and $\delta_{i j}=0$ for $i \neq j$.

### 3.4.3 Linear Triangular Element with Three Nodes: Global Coordinate

Within a two-dimensional element (triangular area) defined by three nodes, one at each apex, the variation of the field variable, $\phi^{(e)}(x, y)$, can be approximated by a linear function (as illustrated in Fig. 3.23) of the form

$$
\begin{equation*}
\phi^{(e)}(x, y)=\alpha_{1}+\alpha_{2} x+\alpha_{3} y \tag{3.54}
\end{equation*}
$$

This function ensures the inter-element continuity of the field variable $\phi^{(e)}(x, y)$.

The element nodes are identified as 1,2 , and 3 in a counterclockwise orientation, with their nodal values as $\phi_{1}, \phi_{2}$, and $\phi_{3}$. The nodal coordinates are specified by $\left(x_{1}, y_{1}\right),\left(x_{2}, y_{2}\right)$, and $\left(x_{3}, y_{3}\right)$.

The nodal values of the field variable must be satisfied as

$$
\begin{align*}
& \phi_{1}=\alpha_{1}+\alpha_{2} x_{1}+\alpha_{3} y_{1} \\
& \phi_{2}=\alpha_{1}+\alpha_{2} x_{2}+\alpha_{3} y_{2}  \tag{3.55}\\
& \phi_{3}=\alpha_{1}+\alpha_{2} x_{3}+\alpha_{3} y_{3}
\end{align*}
$$

leading to the determination of the generalized coefficients in the form

$$
\left\{\begin{array}{l}
\alpha_{1}  \tag{3.56}\\
\alpha_{2} \\
\alpha_{3}
\end{array}\right\}=\frac{1}{2 A}\left[\begin{array}{ccc}
\left(x_{2} y_{3}-x_{3} y_{2}\right) & \left(x_{3} y_{1}-x_{1} y_{3}\right) & \left(x_{1} y_{2}-x_{2} y_{1}\right) \\
\left(y_{2}-y_{3}\right) & \left(y_{3}-y_{1}\right) & \left(y_{1}-y_{2}\right) \\
\left(x_{3}-x_{2}\right) & \left(x_{1}-x_{3}\right) & \left(x_{2}-x_{1}\right)
\end{array}\right]\left\{\begin{array}{l}
\phi_{1} \\
\phi_{2} \\
\phi_{3}
\end{array}\right\}
$$

where

$$
2 A=\left|\begin{array}{lll}
1 & x_{1} & y_{1}  \tag{3.57}\\
1 & x_{2} & y_{2} \\
1 & x_{3} & y_{3}
\end{array}\right|
$$



Fig. 3.23 Linear approximation for the field variable $\phi$ within a triangular element.

Substitution of $\alpha_{1}, \alpha_{2}$, and $\alpha_{3}$ into the expression for the element approximation function results in

$$
\begin{equation*}
\phi^{(e)}(x, y)=N_{1}(x, y) \phi_{1}+N_{2}(x, y) \phi_{2}+N_{3}(x, y) \phi_{3} \tag{3.58}
\end{equation*}
$$

where the shape functions $N_{1}=\xi_{1}, N_{2}=\xi_{2}$, and $N_{3}=\xi_{3}$ are the same as the area coordinates with properties $\xi_{i}\left(x_{j}, y_{j}\right)=\delta_{i j}$ and $\sum_{i=1}^{3} \xi_{i}=1$. Their variation within the element is given in Fig. 3.24.

### 3.4.4 Quadratic Triangular Element with Six Nodes

The field variable can be approximated by a complete quadratic function within a triangular element in the form

$$
\begin{equation*}
\phi^{(e)}(x, y)=\alpha_{1}+\alpha_{2} x+\alpha_{3} y+\alpha_{4} x^{2}+\alpha_{5} x y+\alpha_{6} y^{2} \tag{3.59a}
\end{equation*}
$$

or

$$
\begin{equation*}
\phi^{(e)}(x, y)=\mathbf{g}^{T} \boldsymbol{\alpha} \tag{3.59b}
\end{equation*}
$$

where the vectors $\mathbf{g}$ and $\boldsymbol{\alpha}$ are defined by

$$
\mathbf{g}^{T}=\left\{\begin{array}{llllll}
1 & x & y & x^{2} & x y & y^{2} \tag{3.60}
\end{array}\right\}
$$



Fig. 3.24 Variation of linear shape functions within a triangular element.
and

$$
\boldsymbol{\alpha}^{T}=\left\{\begin{array}{llllll}
\alpha_{1} & \alpha_{2} & \alpha_{3} & \alpha_{4} & \alpha_{5} & \alpha_{6} \tag{3.61}
\end{array}\right\}
$$

However, this representation requires a triangular element with six nodes, as shown in Fig. 3.25, in order to determine its six unknown coefficients, $\alpha_{i}$.

At each node, the field variable, $\phi^{(e)}\left(x_{i}, y_{i}\right)$, is evaluated as

$$
\left\{\begin{array}{l}
\phi_{1}  \tag{3.62}\\
\phi_{2} \\
\phi_{3} \\
\phi_{4} \\
\phi_{5} \\
\phi_{6}
\end{array}\right\}=\left[\begin{array}{llllll}
1 & x_{1} & y_{1} & x_{1}^{2} & x_{1} y_{1} & y_{1}^{2} \\
1 & x_{2} & y_{2} & x_{2}^{2} & x_{2} y_{2} & y_{2}^{2} \\
1 & x_{3} & y_{3} & x_{3}^{2} & x_{3} y_{3} & y_{3}^{2} \\
1 & x_{4} & y_{4} & x_{4}^{2} & x_{4} y_{4} & y_{4}^{2} \\
1 & x_{5} & y_{5} & x_{5}^{2} & x_{5} y_{5} & y_{5}^{2} \\
1 & x_{6} & y_{6} & x_{6}^{2} & x_{6} y_{6} & y_{6}^{2}
\end{array}\right]\left\{\begin{array}{l}
\alpha_{1} \\
\alpha_{2} \\
\alpha_{3} \\
\alpha_{4} \\
\alpha_{5} \\
\alpha_{6}
\end{array}\right\} \text { or } \varphi=\mathbf{A} \alpha
$$



Fig. 3.25 Variation of linear shape functions within a triangular element.

Solving for the generalized coordinates in terms of nodal coordinates and nodal values of the field variable yields

$$
\begin{equation*}
\alpha=\mathbf{A}^{-1} \varphi \tag{3.63}
\end{equation*}
$$

Substituting for the generalized coordinates in Eq. (3.59) results in

$$
\begin{equation*}
\phi^{(e)}(x, y)=\mathbf{g}^{T} \mathbf{A}^{-1} \varphi \tag{3.64}
\end{equation*}
$$

However, $\phi^{(e)}(x, y)$ can also be expressed within the element through the use of its nodal values $\phi_{i}$ as

$$
\begin{equation*}
\phi^{(e)}(x, y)=\sum_{i=1}^{6} N_{i}(x, y) \phi_{i} \quad \text { or } \quad \phi^{(e)}(x, y)=\mathbf{N}^{T} \boldsymbol{\varphi} \tag{3.65}
\end{equation*}
$$

where $\mathbf{N}$ is the vector of shape functions, $N_{i}(i=1,6)$. Comparison of the last two equations results in the explicit form of the shape functions $N_{i}$ as

$$
\begin{equation*}
\mathbf{N}^{T}=\mathbf{g}^{T} \mathbf{A}^{-1} \tag{3.66}
\end{equation*}
$$

In providing the explicit forms of the shape functions, lengthy expressions are avoided by utilizing the expressions for the area coordinates of $\xi_{1}, \xi_{2}$, and $\xi_{3}$, as derived in Eq. (3.17), thus leading to

$$
\begin{equation*}
\mathbf{N}^{T}=\left\{\left(2 \xi_{1}-1\right) \xi_{1} \quad\left(2 \xi_{2}-1\right) \xi_{2} \quad\left(2 \xi_{3}-1\right) \xi_{3} \quad 4 \xi_{1} \xi_{2} \quad 4 \xi_{2} \xi_{3} \quad 4 \xi_{3} \xi_{1}\right\} \tag{3.67}
\end{equation*}
$$

or

$$
\begin{array}{lll}
N_{1}=\left(2 \xi_{1}-1\right) \xi_{1}, & N_{2}=\left(2 \xi_{2}-1\right) \xi_{2}, & N_{3}=\left(2 \xi_{3}-1\right) \xi_{3}  \tag{3.68}\\
N_{4}=4 \xi_{1} \xi_{2}, & N_{5}=4 \xi_{2} \xi_{3}, & N_{6}=4 \xi_{3} \xi_{1}
\end{array}
$$

Variation of these shape functions within the element is shown in Fig. 3.26.

### 3.4.5 Linear Quadrilateral Element with Four Nodes: Centroidal Coordinate

For a quadrilateral element with four nodes, the field variable, $\phi^{(e)}(x, y)$, is approximated by a linear function (refer to Fig. 3.27) in terms of the natural (centroidal) coordinates, $-1 \leq \xi \leq 1$ and $-1 \leq \eta \leq 1$, as

$$
\begin{equation*}
\phi^{(e)}(\xi, \eta)=\alpha_{1}+\alpha_{2} \xi+\alpha_{3} \eta+\alpha_{4} \xi \eta \tag{3.69}
\end{equation*}
$$

This element approximation function ensures the inter-element continuity of only the field variable. The nodal values of the function are identified by $\phi_{1}$, $\phi_{2}, \phi_{3}$, and $\phi_{4}$. Evaluation of the function at each node with coordinates $\left(\xi_{1}=-1, \eta_{1}=-1\right), \quad\left(\xi_{2}=1, \eta_{2}=-1\right), \quad\left(\xi_{3}=1, \eta_{3}=1\right)$, and $\left(\xi_{4}=-1, \eta_{4}=1\right)$ leads to

$$
\left\{\begin{array}{l}
\phi_{1}  \tag{3.70}\\
\phi_{2} \\
\phi_{3} \\
\phi_{4}
\end{array}\right\}=\left[\begin{array}{cccc}
1 & -1 & -1 & 1 \\
1 & 1 & -1 & -1 \\
1 & 1 & 1 & 1 \\
1 & -1 & 1 & -1
\end{array}\right]\left\{\begin{array}{l}
\alpha_{1} \\
\alpha_{2} \\
\alpha_{3} \\
\alpha_{4}
\end{array}\right\}
$$

Solving for $\alpha_{1}, \alpha_{2}, \alpha_{3}$, and $\alpha_{4}$ results in

$$
\left\{\begin{array}{l}
\alpha_{1}  \tag{3.71}\\
\alpha_{2} \\
\alpha_{3} \\
\alpha_{4}
\end{array}\right\}=\frac{1}{4}\left[\begin{array}{cccc}
1 & 1 & 1 & 1 \\
-1 & 1 & 1 & -1 \\
-1 & -1 & 1 & 1 \\
1 & -1 & 1 & -1
\end{array}\right]\left\{\begin{array}{l}
\phi_{1} \\
\phi_{2} \\
\phi_{3} \\
\phi_{4}
\end{array}\right\}
$$

and their substitution in the element approximation function yields

$$
\begin{equation*}
\phi^{(e)}(\xi, \eta)=\sum_{i=1}^{4} N_{i}(\xi, \eta) \phi_{i} \tag{3.72}
\end{equation*}
$$



Fig. 3.26 Variation of quadratic shape functions within a triangular element.


Fig. 3.27 Bi-linear approximation for the field variable $\phi$ within a quadrilateral element.
in which

$$
\begin{equation*}
N_{i}=\frac{1}{4}\left(1+\xi \xi_{i}\right)\left(1+\eta \eta_{i}\right) \tag{3.73}
\end{equation*}
$$

with $\xi_{i}$ and $\eta_{i}$ representing the coordinates of the corner nodes in the natural coordinate system. The shape functions have the property $N_{i}\left(\xi_{j}, \eta_{j}\right)=\delta_{i j}$, where $\delta_{i j}=1$ for $i=j$ and $\delta_{i j}=0$ for $i \neq j$. They are graphically illustrated in Fig. 3.28.

### 3.5 Isoparametric Elements: Curved Boundaries

The modeling of domains involving curved boundaries by using straightsided elements may not provide satisfactory results. However, the family of elements known as "isoparametric elements" is suitable for such boundaries. The shape (or geometry) and the field variable of these elements are described by the same interpolation functions of the same order. The representation of geometry (element shape) in terms of linear (or nonlinear) shape functions can be considered as a mapping procedure that transforms a square in local coordinates to a regular quadrilateral (or distorted shape) in global coordinates (Fig. 3.29) (Ergatoudis et al. 1968).


Fig. 3.28 Variation of bi-linear shape functions within a quadrilateral element.


Fig. 3.29 Mapping from a unit square to an arbitrary straightsided quadrilateral.

The most widely used elements are triangular or quadrilateral because of their ability to approximate complex geometries. An arbitrary straight-sided quadrilateral in global coordinates, $(x, y)$, can be obtained by a point mapping from the "standard square" defined in natural coordinates, $(\xi, \eta)$. The mapping shown in Fig. 3.29 can be achieved by

$$
\begin{align*}
x & =\frac{1}{4}(1-\xi)(1-\eta) x_{1}+\frac{1}{4}(1+\xi)(1-\eta) x_{2} \\
& +\frac{1}{4}(1+\xi)(1+\eta) x_{3}+\frac{1}{4}(1-\xi)(1+\eta) x_{4}  \tag{3.74}\\
y & =\frac{1}{4}(1-\xi)(1-\eta) y_{1}+\frac{1}{4}(1+\xi)(1-\eta) y_{2} \\
& +\frac{1}{4}(1+\xi)(1+\eta) y_{3}+\frac{1}{4}(1-\xi)(1+\eta) y_{4}
\end{align*}
$$

or

$$
\begin{equation*}
x=\sum_{i=1}^{4} N_{i}(\xi, \eta) x_{i} \quad \text { and } \quad y=\sum_{i=1}^{4} N_{i}(\xi, \eta) y_{i} \tag{3.75}
\end{equation*}
$$

in which

$$
\begin{equation*}
N_{i}=\frac{1}{4}\left(1+\xi \xi_{i}\right)\left(1+\eta \eta_{i}\right) \tag{3.76}
\end{equation*}
$$

with $\left(\xi_{1}=-1, \eta_{1}=-1\right),\left(\xi_{2}=1, \eta_{2}=-1\right),\left(\xi_{3}=1, \eta_{3}=1\right)$, and $\left(\xi_{4}=-1\right.$, $\eta_{4}=1$ ).

In the case of an element with curved boundaries in global coordinates, quadratic shape functions can be used to map it on to a unit square in local coordinates, as shown in Fig. 3.30. The mapping can be achieved by

$$
\begin{equation*}
x=\sum_{i=1}^{8} N_{i}(\xi, \eta) x_{i} \quad \text { and } \quad y=\sum_{i=1}^{8} N_{i}(\xi, \eta) y_{i} \tag{3.77}
\end{equation*}
$$

in which


Fig. 3.30 Mapping from a unit square to a quadrilateral with curved sides.

$$
\begin{align*}
& N_{1}=\frac{1}{4}(1+\xi)(1+\eta)-\frac{1}{2}\left(N_{5}+N_{8}\right) \\
& N_{2}=\frac{1}{4}(1-\xi)(1+\eta)-\frac{1}{2}\left(N_{5}+N_{6}\right) \\
& N_{3}=\frac{1}{4}(1-\xi)(1-\eta)-\frac{1}{2}\left(N_{6}+N_{7}\right) \\
& N_{4}=\frac{1}{4}(1+\xi)(1-\eta)-\frac{1}{2}\left(N_{7}+N_{8}\right)  \tag{3.78}\\
& N_{5}=\frac{1}{2}\left(1-\xi^{2}\right)(1+\eta) \\
& N_{6}=\frac{1}{2}(1-\xi)\left(1-\eta^{2}\right) \\
& N_{7}=\frac{1}{2}\left(1-\xi^{2}\right)(1-\eta) \\
& N_{8}=\frac{1}{2}(1+\xi)\left(1-\eta^{2}\right)
\end{align*}
$$

When the elements have curved boundaries, or arbitrary nodal locations (such as the quadrilaterals), the integrals appearing in the expression for the element matrix are most easily evaluated by using a natural coordinate system. Since it is more advantageous to use natural coordinates, the variables of integration are changed so that the integrals can be evaluated using natural coordinates. In two dimensions, the integral over an arbitrary quadrilateral region of $d x d y$ becomes an integral over a square area of $d \xi d \eta$ in a natural coordinate system in the form

$$
\begin{equation*}
\int_{A} f(x, y) d x d y=\int_{-1-1}^{1} \int_{-1}^{1} g(\xi, \eta)|\mathbf{J}| d \xi d \eta \tag{3.79}
\end{equation*}
$$

where $|J|$ is the determinant of the Jacobian matrix relating the term $d x d y$ to $d \xi d \eta$ from advanced calculus as

$$
\begin{equation*}
d x d y=|\mathbf{J}| d \xi d \eta \tag{3.80}
\end{equation*}
$$

The Jacobian matrix, $\mathbf{J}$, is given by

$$
\mathbf{J}=\left[\begin{array}{ll}
\frac{\partial x}{\partial \xi} & \frac{\partial y}{\partial \xi}  \tag{3.81}\\
\frac{\partial x}{\partial \eta} & \frac{\partial y}{\partial \eta}
\end{array}\right]
$$

whose determinant is always positive, $|\mathbf{J}|>0$, for a one-to-one mapping.
It is not necessary to use interpolation or shape functions of the same order for describing both the geometry and field variable of an element. If the geometry is described by a lower-order model (in comparison to that for the field variable), the element is called a "subparametric element." On the other hand, if the geometry is described by a higher-order interpolation function, then the element is termed a "superparametric" element.

### 3.6 Numerical Evaluation of Integrals

The evaluation of line or area integrals appearing in the finite element equations can be performed numerically by employing the Gaussian integration method (Stroud and Secrest 1966). This method locates sampling points (also called Gaussian points) to achieve the greatest accuracy.

### 3.6.1 Line Integrals

The line integrals encountered commonly are of the form

$$
\begin{equation*}
I=\int_{a}^{b} f(x) d x \tag{3.82}
\end{equation*}
$$

The limits of this integral can be changed by introducing a new variable as

$$
\begin{equation*}
x=\frac{1}{2}[(b-a) \xi+(b+a)] \tag{3.83}
\end{equation*}
$$

Thus, the integral given by Eq. (3.82) can be rewritten as

$$
\begin{equation*}
I=\int_{-1}^{1} f(\xi) J d \xi \tag{3.84}
\end{equation*}
$$

in which the variables $\xi$ and $J$ are given by

$$
\begin{equation*}
\xi=\frac{2}{b-a}\left[x-\frac{(b+a)}{2}\right] \tag{3.85}
\end{equation*}
$$

and

$$
\begin{equation*}
J=\frac{d x}{d \xi}=\frac{b-a}{2} \tag{3.86}
\end{equation*}
$$

Integrals expressed in the form of Eq. (3.84) are almost always evaluated numerically. The most commonly used Gaussian integration technique approximates the integral in the form

$$
\begin{equation*}
I=\int_{-1}^{1} f(\xi) d \xi \approx \sum_{i=1}^{n} w_{i} f\left(\xi_{i}\right) \tag{3.87}
\end{equation*}
$$

The weights of the numerical integration are denoted by $w_{i}$, and the number of evaluation points, $\xi_{i}$ (referred to as the Gaussian points), depends on the order of the polynomial approximation of the integrand.

In general, the integrand $f(\xi)$ in Eq. (3.87) can be approximated as

$$
\begin{equation*}
f(\xi)=\alpha_{1}+\alpha_{2} \xi+\alpha_{3} \xi^{2}+\alpha_{4} \xi^{3}+\ldots+\alpha_{2 n} \xi^{2 n-1} \tag{3.88}
\end{equation*}
$$

resulting in

$$
\begin{equation*}
I=\int_{-1}^{1} f(\xi) d \xi=2 \alpha_{1}+\frac{2}{3} \alpha_{3}+\ldots+\frac{2}{2 n-1} \alpha_{2 n-1} \tag{3.89}
\end{equation*}
$$

and

$$
\begin{align*}
I=\sum_{i=1}^{n} w_{i} f\left(\xi_{i}\right) & =\alpha_{1} \sum_{i=1}^{n} w_{i}+\alpha_{2} \sum_{i=1}^{n} w_{i} \xi_{i}+\alpha_{3} \sum_{i=1}^{n} w_{i} \xi_{i}^{2}+\ldots  \tag{3.90}\\
& +\alpha_{2 n} \sum_{i=1}^{n} w_{i} \xi_{i}^{2 n-1}
\end{align*}
$$

Equating the coefficients of the $\alpha_{i}^{\prime}$ 's in Eq. (3.89) and (3.90) leads to

$$
\begin{align*}
& \sum_{i=1}^{n} w_{i}=2, \quad \sum_{i=1}^{n} w_{i} \xi_{i}=0 \\
& \sum_{i=1}^{n} w_{i} \xi_{i}^{2}=\frac{2}{3}, \quad \sum_{i=1}^{n} w_{i} \xi_{i}^{2 n-2}=\frac{2}{2 n-1}  \tag{3.91}\\
& \sum_{i=1}^{n} w_{i} \xi_{i}^{2 n-1}=0
\end{align*}
$$

providing $2 n$ equations in $n$ unknowns for positions $\xi_{i}$ and $n$ unknowns for weights $w_{i}$. Hence, for a polynomial of degree $p=2 n-1$, it is sufficient to use $n$ sampling points for exact integration, i.e., the exact integration is obtained if $n \geq(p+1) / 2$. This means that for " $n$ " sampling points, a polynomial of degree $(2 n-1)$ can be integrated exactly.

Rewriting Eq. (3.84) in its final form as

$$
\begin{equation*}
I=\int_{a}^{b} f(x) d x=\frac{b-a}{2} \int_{-1}^{1} f\left[\frac{b-a}{2} \xi+\frac{b+a}{2}\right] d \xi \tag{3.92}
\end{equation*}
$$

and assuming a third-order polynomial ( $p=3$ ) approximation for $f(\xi)$ in Eq. (3.92), this integral is approximated with two sampling points $(n=2)$ as

$$
\begin{equation*}
I \approx w_{1} f\left(\xi_{1}\right)+w_{2} f\left(\xi_{2}\right) \tag{3.93}
\end{equation*}
$$

where $-1 \leq \xi_{1}, \xi_{2} \leq 1$, and $w_{1}, w_{2}$ (Gaussian weights), $\xi_{1}$, and $\xi_{2}$ are to be determined. For each coefficient of the cubic representation of $f(\xi)$, Eq. (3.91) yields

$$
\begin{align*}
& \int_{-1}^{1} \xi^{3} d \xi=0=w_{1} \xi_{1}^{3}+w_{2} \xi_{2}^{3}  \tag{3.94a}\\
& \int_{-1}^{1} \xi^{2} d \xi=\frac{2}{3}=w_{1} \xi_{1}^{2}+w_{2} \xi_{2}^{2} \tag{3.94b}
\end{align*}
$$

$$
\begin{gather*}
\int_{-1}^{1} \xi d \xi=0=w_{1} \xi_{1}+w_{2} \xi_{2}  \tag{3.94c}\\
\int_{-1}^{1} d \xi=2=w_{1}+w_{2} \tag{3.94d}
\end{gather*}
$$

Multiplying Eq. (3.94c) by $\xi_{1}^{2}$ and subtracting it from Eq. (3.94a) gives

$$
\begin{equation*}
w_{2} \xi_{2}\left(\xi_{2}^{2}-\xi_{1}^{2}\right)=w_{2} \xi_{2}\left(\xi_{2}-\xi_{1}\right)\left(\xi_{2}+\xi_{1}\right)=0 \tag{3.95}
\end{equation*}
$$

For this equality to be valid, the possibilities are:

1. $w_{2}=0 \rightarrow \quad$ one-term formula-reject.
2. $\xi_{2}=0 \rightarrow w_{1}=0 \quad$ one-term formula-reject.
3. $\xi_{1}=\xi_{2} \rightarrow w_{1}=0 \quad$ one-term formula-reject.
4. $\xi_{2}=-\xi_{1} \rightarrow \quad$ ACCEPTED.

Thus, substituting for $\xi_{2}=-\xi_{1}$ in Eq. (3.94) leads to

$$
\begin{gather*}
w_{1}=w_{2}  \tag{3.96a}\\
2 \xi_{1}^{2}=\frac{2}{\sqrt{3}} \rightarrow \xi_{1}=\frac{1}{\sqrt{3}}, \xi_{2}=-\frac{1}{\sqrt{3}}  \tag{3.96b}\\
w_{1}=w_{2}=1 \tag{3.96c}
\end{gather*}
$$

The numerical integration, Eq. (3.93) becomes

$$
\begin{equation*}
I \approx f\left(-\frac{1}{\sqrt{3}}\right)+f\left(\frac{1}{\sqrt{3}}\right) \tag{3.97}
\end{equation*}
$$

The Gaussian points and weights for polynomials of order up to 5 are summarized in Table 3.1. The Gaussian points for higher order polynomial approximation are given by Abramowitz and Stegun (1972).

An example is considered that evaluates the line integral given by

$$
\begin{equation*}
I=\int_{-0.25}^{0.25} e^{x} d x \tag{3.98}
\end{equation*}
$$

Table 3.1 Positions and weights for Gauss integration.

| Gauss Points | $\xi_{i}$ | $w_{i}$ |
| :---: | :---: | :---: |
| $n=1$ | 0.00 | 2.00 |
| $n=2$ | $\pm \sqrt{1 / 3}$ | 1.00 |
| $n=3$ | 0.00 | $8 / 9$ |
|  | $\pm \sqrt{3 / 5}$ | $5 / 9$ |
| $n=4$ | $\pm 0.339981$ | 0.652145 |
|  | $\pm 0.861136$ | 0.347854 |
| $n=5$ | 0.00 | 0.568888 |
|  | $\pm 0.538469$ | 0.478628 |
|  | $\pm 0.906179$ | 0.236926 |

This integral can be rewritten as

$$
\begin{equation*}
I=\frac{1}{4} \int_{-1}^{1} e^{\xi / 4} d \xi \tag{3.99}
\end{equation*}
$$

Applying Gauss's formula with $n=2$ integration points, this integral is approximated as

$$
\begin{equation*}
I \approx \frac{1}{4}\left[e^{-1 / 4 \sqrt{3}}+e^{1 / 4 \sqrt{3}}\right]=0.505217 \tag{3.100}
\end{equation*}
$$

The exact solution is $I=2 \times \sinh (0.25)=0.505224$.

### 3.6.2 Triangular Area Integrals

The area integrals over a triangular region given in the form

$$
\begin{equation*}
I=\int_{A} f(x, y) d A \tag{3.101}
\end{equation*}
$$

can be rewritten as

$$
\begin{equation*}
I=\int_{0}^{1} \int_{0}^{1-\xi_{2}} f\left(\xi_{1}, \xi_{2}\right)|\mathbf{J}| d \xi_{1} d \xi_{2} \tag{3.102}
\end{equation*}
$$

in which $|\mathbf{J}|$ is the determinant of the Jacobian matrix expressed as

$$
\mathbf{J}=\left[\begin{array}{cc}
\frac{\partial x}{\partial \xi_{1}} & \frac{\partial y}{\partial \xi_{1}}  \tag{3.103}\\
\frac{\partial x}{\partial \xi_{2}} & \frac{\partial y}{\partial \xi_{2}}
\end{array}\right]=\left[\begin{array}{ll}
\left(x_{1}-x_{3}\right) & \left(y_{1}-y_{3}\right) \\
\left(x_{2}-x_{3}\right) & \left(y_{2}-y_{3}\right)
\end{array}\right]=2 A
$$

relating the area coordinates (discussed in Sec. 3.3.4.2.1) to Cartesian coordinates

$$
\left\{\begin{array}{c}
\frac{\partial}{\partial \xi_{1}}  \tag{3.104}\\
\frac{\partial}{\partial \xi_{2}}
\end{array}\right\}=[\mathbf{J}]\left\{\begin{array}{c}
\frac{\partial}{\partial x} \\
\frac{\partial}{\partial y}
\end{array}\right\}
$$

The extent of the triangular area of integration is defined by the coordinates $\left(x_{i}, y_{i}\right)$ (with $i=1,2,3$ ) of the vertices. The Gaussian approximation to the integration is expressed as

$$
\begin{equation*}
I=\int_{0}^{1} \int_{0}^{1-\xi_{2}} f\left(\xi_{1}, \xi_{2}\right)|\mathbf{J}| d \xi_{1} d \xi_{2} \approx 2 A \sum_{i=1}^{n} w_{i} f\left(\xi_{1 i}, \xi_{2 i}\right) \tag{3.105}
\end{equation*}
$$

in which the weights of the numerical integration are denoted by $w_{i}$. The number of evaluation points, $\xi_{1 i}$ and $\xi_{2 i}$, are referred to as the Gaussian integration points and they depend on the order of the polynomial approximation of the integrand. Depending on the degree of approximation, the weights and the evaluation points are given by Huebner et al. (2001).

An example is considered that evaluates the area integral given by

$$
\begin{equation*}
I=\int_{A} x y d A \tag{3.106}
\end{equation*}
$$

in which the area $A$ is defined by a triangle whose vertices are $(1,1),(3,2)$, and $(2,3)$, as shown in Fig. 3.31. This integral can also be evaluated exactly by using Eq. (3.19).


Fig. 3.31 A triangular element and its mapping.

The coordinates $(x, y)$ of a point within a triangular area can be expressed as linear combinations of the nodal coordinates $\left(x_{1}, y_{1}\right),\left(x_{2}, y_{2}\right)$, and $\left(x_{3}, y_{3}\right)$ and the area coordinates $\xi_{1}, \xi_{2}$, and $\xi_{3}$ as

$$
\begin{align*}
& x=\xi_{1} x_{1}+\xi_{2} x_{2}+\xi_{3} x_{3}=\xi_{1}+3 \xi_{2}+2 \xi_{3}=-\xi_{1}+\xi_{2}+2  \tag{3.107a}\\
& y=\xi_{1} y_{1}+\xi_{2} y_{2}+\xi_{3} y_{3}=\xi_{1}+2 \xi_{2}+3 \xi_{3}=-2 \xi_{1}-\xi_{2}+3 \tag{3.107b}
\end{align*}
$$

with

$$
\begin{equation*}
\xi_{1}+\xi_{2}+\xi_{3}=1 \tag{3.107c}
\end{equation*}
$$

Substituting for $x$ and $y$ in the integrand of Eq. (3.106) results in

$$
\begin{equation*}
I=2 A \int_{0}^{1} \int_{0}^{1-\xi_{2}}\left(2 \xi_{1}^{2}-\xi_{2}^{2}-\xi_{1} \xi_{2}-7 \xi_{1}+\xi_{2}+6\right) d \xi_{1} d \xi_{2} \tag{3.108}
\end{equation*}
$$

Utilizing $n=3$ Gaussian points as shown in Fig. 3.32, approximation to the integration by Eq. (3.105) becomes

$$
\begin{equation*}
I \approx 2 A\left[w_{1} f\left(\xi_{11}, \xi_{21}\right)+w_{2} f\left(\xi_{12}, \xi_{22}\right)+w_{3} f\left(\xi_{13}, \xi_{23}\right)\right] \tag{3.109}
\end{equation*}
$$

in which $w_{1}=w_{2}=w_{3}=1 / 6, \xi_{11}=1 / 2, \xi_{21}=0, \xi_{12}=1 / 2, \quad \xi_{22}=1 / 2$, $\xi_{13}=0$, and $\xi_{23}=1 / 2$. The area of the triangle is obtained from Eq. (3.18) as $2 A=3$. Thus, the Gaussian approximation leads to

$$
\begin{aligned}
I \approx & 2 A\left[w_{1}\left(2 \xi_{11}^{2}-\xi_{21}^{2}-\xi_{11} \xi_{21}-7 \xi_{11}+\xi_{21}+6\right)\right. \\
& +w_{2}\left(2 \xi_{12}^{2}-\xi_{22}^{2}-\xi_{12} \xi_{22}-7 \xi_{12}+\xi_{22}+6\right) \\
& \left.+w_{3}\left(2 \xi_{13}^{2}-\xi_{23}^{2}-\xi_{13} \xi_{23}-7 \xi_{13}+\xi_{23}+6\right)\right]
\end{aligned}
$$



Fig. 3.32 Three Gaussian points, located at mid-sides, for approximate integration.
and

$$
I \approx 3 \frac{1}{6}\left[\left(2 \frac{1}{4}-7 \frac{1}{2}+6\right)+\left(2 \frac{1}{4}-\frac{1}{4}-\frac{1}{4}-7 \frac{1}{2}+\frac{1}{2}+6\right)+\left(-\frac{1}{4}+\frac{1}{2}+6\right)\right]
$$

and

$$
\begin{equation*}
I \approx \frac{1}{2}\left[6+\frac{25}{4}\right]=6.125 \tag{3.110}
\end{equation*}
$$

For the exact evaluation, substituting for $x$ and $y$ in the integrand results in

$$
\begin{equation*}
I=\int_{A}\left(\xi_{1}^{2}+6 \xi_{2}^{2}+6 \xi_{3}^{2}+5 \xi_{1} \xi_{2}+13 \xi_{2} \xi_{3}+5 \xi_{1} \xi_{3}\right) d A \tag{3.111}
\end{equation*}
$$

Utilizing the formula of Eq. (3.19) for exact integration results in

$$
\begin{align*}
I & =\frac{2 A}{4!}(2!+6 \times 2!+6 \times 2!+5 \times 1!\times 1!+13 \times 1!\times 1!+5 \times 1!\times 1!) \\
& =\frac{3}{24}[13 \times 2+23]=\frac{49}{8}=6.125 \tag{3.112}
\end{align*}
$$

### 3.6.3 Quadrilateral Area Integrals

The quadrilateral area integrals appearing in the form

$$
\begin{equation*}
I=\int_{a}^{b} \int_{c}^{d} f(x, y) d x d y \tag{3.113}
\end{equation*}
$$

can be rewritten as

$$
\begin{equation*}
I=\int_{-1}^{1} \int_{-1}^{1} f(\xi, \eta)|\mathbf{J}| d \xi d \eta \tag{3.114}
\end{equation*}
$$

in which $|\mathbf{J}|$ is the determinant of the Jacobian matrix expressed as

$$
\mathbf{J}=\left[\begin{array}{ll}
\frac{\partial x}{\partial \xi} & \frac{\partial y}{\partial \xi}  \tag{3.115}\\
\frac{\partial x}{\partial \eta} & \frac{\partial y}{\partial \eta}
\end{array}\right] \text { relating }\left\{\begin{array}{c}
\frac{\partial}{\partial \xi} \\
\frac{\partial}{\partial \eta}
\end{array}\right\}=[\mathbf{J}]\left\{\begin{array}{c}
\frac{\partial}{\partial x} \\
\frac{\partial}{\partial y}
\end{array}\right\}
$$

These integrals can be evaluated first with respect to one variable and then with respect to the other leading to

$$
\begin{equation*}
I=\int_{-1}^{1} \int_{-1}^{1} f(\xi, \eta)|\mathbf{J}| d \xi d \eta \approx \sum_{i=1}^{n} \sum_{j=1}^{n} w_{i} w_{j} f\left(\xi_{i}, \eta_{j}\right)\left|\mathbf{J}\left(\xi_{i}, \eta_{j}\right)\right| \tag{3.116}
\end{equation*}
$$

in which $w_{i}$ represent the weights of the numerical integration, and $\xi_{i}$ and $\eta_{i}$ are the Gaussian integration points. They are given by Abramowitz and Stegun (1972) and depend on the order of the polynomial approximation of the integrand.

An example is considered that evaluates the area integral given by

$$
\begin{equation*}
I=\int_{A} x y d A \tag{3.117}
\end{equation*}
$$

in which the area $A$ is defined by a quadrilateral whose vertices are $(1,1)$, $(3,2),(4,4)$, and $(2,3)$ as shown in Fig. 3.33.

The coordinates $(x, y)$ of a point within a quadrilateral area can be expressed as linear combinations of the nodal coordinates $\left(x_{1}, y_{1}\right),\left(x_{2}, y_{2}\right),\left(x_{3}, y_{3}\right)$, and $\left(x_{4}, y_{4}\right)$ and the natural coordinates $\xi$ and $\eta$ as

$$
\begin{align*}
x & =\frac{1}{4}[(1-\xi)(1-\eta)+3(1+\xi)(1-\eta)  \tag{3.118}\\
& +4(1+\xi)(1+\eta)+2(1-\xi)(1+\eta)] \\
y & =\frac{1}{4}[(1-\xi)(1-\eta)+2(1+\xi)(1-\eta)  \tag{3.119}\\
& +4(1+\xi)(1+\eta)+3(1-\xi)(1+\eta)]
\end{align*}
$$



Fig. 3.33 A four-noded quadrilateral element and its mapping.

The Jacobian matrix is obtained as

$$
\mathbf{J}=\left[\begin{array}{cc}
1 & 1 / 2 \\
1 / 2 & 1
\end{array}\right]
$$

with its determinant $|J|=3 / 4$.
Utilizing two Gaussian points as shown in Fig. 3.34, the approximation to the integration becomes

$$
\begin{align*}
I & \approx \frac{3}{4}\left[w_{1} w_{1} f\left(\xi_{1}, \eta_{1}\right)+w_{1} w_{2} f\left(\xi_{1}, \eta_{2}\right)\right.  \tag{3.120}\\
& \left.+w_{2} w_{1} f\left(\xi_{2}, \eta_{1}\right)+w_{2} w_{2} f\left(\xi_{2}, \eta_{2}\right)\right]
\end{align*}
$$

in which $w_{1}=w_{2}=1, \quad \xi_{1}=-1 / \sqrt{3}, \xi_{2}=1 / \sqrt{3}, \quad \eta_{1}=-1 / \sqrt{3}$, and $\eta_{2}=$ $1 / \sqrt{3}$.

The function $f(\xi, \eta)$ is expressed as

$$
\begin{align*}
& f(\xi, \eta)=\frac{1}{16}[(1-\xi)(1-\eta)+3(1+\xi)(1-\eta) \\
&+4(1+\xi)(1+\eta)+2(1-\xi)(1+\eta)]  \tag{3.121}\\
& \times[(1-\xi)(1-\eta)+2(1+\xi)(1-\eta) \\
&+4(1+\xi)(1+\eta)+3(1-\xi)(1+\eta) 0]
\end{align*}
$$

Evaluation of the function $f(\xi, \eta)$ at Gaussian integration points results in their numerical evaluations as


Fig. 3.34 Two Gaussian points, in each direction, for approximate integration.

$$
\begin{aligned}
& f\left(\xi_{1}, \eta_{1}\right)=11.33012702 \\
& f\left(\xi_{1}, \eta_{2}\right)=6.16666667 \\
& f\left(\xi_{2}, \eta_{1}\right)=6.16666667 \\
& f\left(\xi_{2}, \eta_{2}\right)=2.66987298
\end{aligned}
$$

Finally, the approximation to the integral from Eq. (3. 120) is determined to be 19.75 .

By using Eq. (3.19), the exact evaluation of this integral can be obtained by integration over two triangular regions defined by the vertices $(1,1),(3,2)$, and $(2,3)$ and $(3,2),(4,4)$, and $(2,3)$. The exact integration over these two regions are obtained as 6.125 and 12.125 . Their summation provides the exact integration over a quadrilateral defined by vertices $(1,1),(3,2),(4,4)$, and $(2,3)$. Thus, the exact integration becomes 19.75 .

### 3.7 Problems

3.1. The completeness criterion for convergence of finite element solutions requires that the interpolating function must be able to reproduce exactly (that is, interpolate to the exact value at every point in the element). In particular, the approximation function $\phi(x, y)$ is specified as

$$
\phi(x, y)=a+b x+c y=\sum N_{i} \phi_{i}
$$

where $a, b$, and $c$ are arbitrary constants, $\phi_{i}$ are the nodal values, and $N_{i}(x, y)$ are the interpolating functions.
(a) Derive a set of three equations that the interpolating functions $N_{i}(x, y)$ must satisfy for completeness.
(b) Show that the standard and quadratic linear interpolation functions for a triangular domain satisfy these requirements.
3.2. Using the coordinate transformation equations given in Sec. 3.5 for an 8 -noded quadrilateral element, determine the isoparametric element shape whose nodal locations are

| Node No. | $x$ | $y$ |
| :---: | ---: | ---: |
| 1 | 6.0 | 3.0 |
| 2 | -4.0 | 3.0 |
| 3 | -5.0 | -3.0 |
| 4 | 4.0 | -3.0 |
| 5 | 1.0 | 4.0 |
| 6 | -3.0 | 0.5 |
| 7 | 0.0 | -2.0 |
| 8 | 5.0 | 0.0 |

3.3. The isoparametric formulation is useful for triangular, as well as for quadrilateral, elements. Also, the area coordinates $\left(\xi_{1}, \xi_{2}, \xi_{3}\right)$ are commonly employed for triangular elements instead of using the local coordinates $(r, s)$. However, because only two of these are independent coordinates, one of them, say $\xi_{3}$, can be eliminated in favor of $\xi_{1}$ and $\xi_{2}$. Thus, for a 3 -noded triangle, the interpolation functions are $N_{i}=\xi_{i}(i=1,2,3)$ and the coordinate transformations, using $\xi_{3}=1-\xi_{1}-\xi_{2}$, are

$$
\begin{aligned}
& x=\xi_{1} x_{1}+\xi_{2} x_{2}+\left(1-\xi_{1}-\xi_{2}\right) x_{3} \\
& y=\xi_{1} y_{1}+\xi_{2} y_{2}+\left(1-\xi_{1}-\xi_{2}\right) y_{3}
\end{aligned}
$$

As illustrated in Fig. 3.35, this clearly maps a triangle with vertices $(1,0),(0,1)$, and $(0,0)$ in the $\xi_{1}-\xi_{2}$ plane into a triangle with vertices $\left(x_{1}, y_{1}\right),\left(x_{2}, y_{2}\right)$, and $\left(x_{3}, y_{3}\right)$ in the $x-y$ plane. Also, the integrals in the $x-y$ plane may be related to integrals in the $\xi_{1}-\xi_{2}$ plane by

$$
\iint() d x d y=\int_{0}^{1} \int_{0}^{1-\xi_{1}}(\quad)|J| d \xi_{2} d \xi_{1}
$$



Fig. 3.35 A triangular element and its mapping.

Explicitly determine the coordinate transformations and the Jacobian matrix for the 6 -noded triangle having the side nodes located at the midpoint of each side. Explain how it is possible to obtain a triangular element in the $x-y$ plane with one or more curved sides. What is the form of the curve?
3.4. For a 4-noded element shown in Fig. 3.36, the mapping is achieved by

$$
x=\sum_{i=1}^{4} N_{i}(\xi, \eta) x_{i} \quad \text { and } \quad y=\sum_{i=1}^{4} N_{i}(\xi, \eta) y_{i}
$$

where

$$
\begin{array}{ll}
N_{1}=\frac{1}{4}(1-\xi)(1-\eta), & N_{2}=\frac{1}{4}(1+\xi)(1-\eta) \\
N_{3}=\frac{1}{4}(1+\xi)(1+\eta), & N_{4}=\frac{1}{4}(1-\xi)(1+\eta)
\end{array}
$$

(a) For this element explicitly determine the Jacobian determinant, and show that it is strictly linear in the local coordinates $\xi$ and $\eta$ and that the term proportional to the product $\xi \eta$ vanishes.
(b) Show that the Jacobian determinant becomes

$$
\begin{aligned}
|\mathbf{J}| & =\frac{1}{4}\left[\left(x_{4}-x_{3}\right)\left(y_{2}-y_{3}\right)-\left(x_{2}-x_{3}\right)\left(y_{4}-y_{3}\right)\right] \\
\text { for } \xi & =\eta=1
\end{aligned}
$$



Fig. 3.36 A four-noded quadrilateral element and its mapping.
(c) Using the definition of the cross-product of the vectors $\nu_{1}$ and $\nu_{2}$ shown in Fig. 3.36, show that at $\xi=\eta=1$

$$
|J|>0 \quad \text { if } \quad 0<\theta<\pi
$$

(d) Based on the results of parts (a) and (c), provide a short argument to show that $|J|>0$ throughout the element and, hence, the coordinate transformation $(\xi, \eta) \rightarrow(x, y)$ is unique and invertible if the interior angles at all nodes are less than $180^{\circ}$.

## Chapter 4

## ANSYS PREPROCESSOR

### 4.1 Fundamentals of Modeling

The fundamental concepts and the Begin and Processor Levels of the ANSYS finite element program are described in Chap. 2. Specifications of all the geometric and material properties, as well as the generation of solid and finite element models, are conducted at the preprocessor level.

There are two approaches for creating a finite element model: solid modeling and direct generation. The solid modeling approach utilizes Primitives (pre-defined geometric shapes) and operations similar to those of computeraided design (CAD) tools, and internally generates the nodes and the elements based on user specifications. Solid modeling is the most commonly used approach because it is much more versatile and powerful. However, the user must have a strong understanding of the concept of meshing in order to utilize the solid modeling approach successfully and efficiently.

Direct generation is entirely dependent on user input for the size, shape, and connectivity of each element and coordinates of each node before it creates the nodes and elements one at a time. It requires the user to keep track of the node and element numbering, which may become tedious-sometimes practically impossible-for complex problems requiring thousands of nodes. It is, however, extremely useful for simple problems as one has full control over the model.

A combination of the two approaches is not only possible, but also advantageous in many cases. A comprehensive list of some important advantages and disadvantages is given in Table 4.1.

### 4.2 Modeling Operations

Within the ANSYS Preprocessor, a finite element model is generated by utilizing various operations, which are explained in this section.

Table 4.1 Advantages and disadvantages of solid modeling and direct generation.

| Advantages | Disadvantages |
| :--- | :--- |
| Solid Modeling |  |
| $\begin{array}{l}\text { Powerful (sometimes the only feasible } \\ \text { way) in modeling three-dimensional } \\ \text { solid volumes with complex geometry. }\end{array}$ | $\begin{array}{l}\text { If the user does not have a good under- } \\ \text { standing of meshing, ANSYS may not } \\ \text { be able to generate the finite element } \\ \text { mesh. }\end{array}$ |
| User data input is rather low. | For simple problems, using solid |
| modeling may be ponderous |  |$]$| Common computer-aided design |
| :--- |
| (CAD)-type operations such as extru- |
| sions, dragging, and rotations are uti- |$\quad$| lized which are not possible when |
| :--- |
| working directly with the nodes and |
| elements. |
| With the basic (primitive) areas and |
| volumes (rectangular, circular etc. |
| areas; cubic, cylindrical, spherical etc. |
| volumes), the Boolean operations (add, |
| subtract, overlap etc.) can be used |
| easily to modify (or tailor) these basic |
| areas or volumes to obtain the desired |
| shape. |

### 4.2.1 Title

This operation defines the title for the ANSYS analysis. This is an optional but recommended step in a typical ANSYS session. It helps the user to keep track of the problems by appearing in the graphics display and output. It becomes extremely useful when the user conducts a case study that involves the same model with different boundary conditions, different material properties, etc. The following menu path is used to change (or specify) the title:

## Utility Menu > File > Change Title

which brings up the dialog box shown in Fig. 4.1. After entering the desired title in the text box, clicking on $O K$ completes the specification of the title.


Fig. 4.1 Dialog box for specifying the title.

### 4.2.2 Elements

The nodes and elements are the essential parts of a finite element model. Before starting meshing, the element type(s) to be used must be defined (otherwise ANSYS refuses to create the mesh). The ANSYS software contains more than 100 different element types in its element library. Each element type has a unique number and a prefix that identifies the element category, such as BEAM3, PLANE42, SOLID45, etc. The elements that are available in ANSYS can be classified according to many different criteria, such as dimensionality, analysis discipline, and material behavior. ANSYS classifies the elements in 21 different groups. In this section, the elements from four of these groups-specifically, structural, thermal, fluid, and FLOTRAN CFD-are considered for different analysis objectives.

1. Structural: For this group of elements, the degrees of freedom at the nodes are displacements. As shown in Fig. 4.2, the structural analysis employs plane, link, beam, pipe, solid, and shell elements. All of the above "subgroups" of elements include several element types with different degree-of-freedom (DOF) sets. Consider the entries Quad 4node 42, Quad 8node 82, and Brick 8node 45 from the Structural Solid subgroup. The first two elements types, quad42 and Quad82, are used for two-dimensional structural problems (plane stress, plane strain, or axisymmetric) whereas the third one is used for threedimensional structural problems. The difference between Quad42 and Quad82 elements is that they have a different number of nodes per element, which implies that they are employing different interpolation functions for the variation of the degrees of freedom along the edges of the element. In this particular case, the variation of displacements along the element edges is assumed to be linear for Quad 4node 42 and quadratic for Quad 8node 82, as shown in Fig. 4.3. The interpolation functions for the Brick 8node 45 element are linear.
2. Thermal: For this group of elements, the degrees of freedom at the nodes are temperatures. The thermal analysis employs mass, link, solid and shell subgroups. The element types in this group differ from each other with similar considerations as explained for structural discipline. Two commonly used thermal elements are shown in Fig. 4.4.


Fig. 4.2 Examples of structural elements in ANSYS.


Fig. 4.3 Linear and quadratic variations of displacements within a 2-D element.


Fig. 4.4 Examples of thermal elements in ANSYS.
3. Fluid: For this group of elements, depending on the type, the degrees of freedom appear as a pair, velocity-pressure or pressure-temperature, at the nodes. Included in this group are two- and three- dimensional acoustic, thermal-fluid coupled pipe, and contained-fluid types of elements.
4. FLOTRAN CFD: This group of elements is similar to the previous one, except it is based on the method of computational finite difference.

Each discipline requires the use of its own element types because the element type determines the degree-of-freedom set (displacements, temperatures, pressures, etc.) and the dimensionality of the problem (2-D or 3-D).

For example, the BEAMA element, shown in Fig. 4.5, has six structural degrees of freedom (displacements and rotations in and about the $x$-, $y$-, and $z$-directions) at each of the two nodes, is a line element, and can be modeled in 3-D space. The PLANE55 element, also shown in Fig. 4.5, which has a total of four thermal degrees of freedom (temperature at each node), is a 4noded quadrilateral element, and can be used only for two-dimensional problems.

In order to specify an element type, the user must be in the Preprocessor. The menu path for element specification is

## Main Menu > Preprocessor > Element Type > Add/Edit/Delete

When this action is taken, a dialog box, shown in Fig. 4.6, appears with the options of Add, Options, Delete, Close, and Help. Choosing Add brings up another dialog box with a list of all available elements, along with the Element type reference number. The element types that are defined in a particular ANSYS analysis are assigned reference numbers. This reference number is used when creating the mesh. If the analysis requires the use of more than one element type, switching from one type to another one is achieved by referring to this number (this point is further explained when discussing Element Attributes).


Fig. 4.5 BEAM4 element for 3-D problems and PLANE55 element for 2-D problems.


Fig. 4.6 Defining an element type in ANSYS.

If the user wants to delete an existing element type, it is achieved by using the same GUI path and choosing Delete, as shown in Fig. 4.7:

## Main Menu > Preprocessor > Element Type > Add/Edit/Delete

Many element types have additional options, known as keyoptions (KEYOPT), and are referred to as Keyopt(1), keyopt(2), etc. For example, as shown in Fig. 4.8, KEYOPT(3) for SOLID42 (4-noded quadrilateral 2-D structural element) allows the user to specify the type of two-dimensional idealization, i.e., plane stress, plane strain, axisymmetric, or plane stress with thickness.

Another example is shown in Fig. 4.9, in which Keyopt (7) for SOLID70 (8-noded thermal solid element for 3-D problems) permits the specification of a standard heat transfer or a nonlinear steady-state fluid flow through a porous medium.

Keyoptions are specified using the same GUI path and choosing Options from the Element Types dialog box.

### 4.2.3 Real Constants

As described in Chap. 1, the calculation of the element matrices requires material properties, nodal coordinates and geometrical parameters. Any data required for the calculation of the element matrix that cannot be determined from the nodal coordinates or material properties are called "real constants" in ANSYS. Typically, real constants are area, thickness, inner diameter, outer diameter, etc. Not all element types require real constants.

Real constants of a particular element type are briefly explained in the "Element Reference" of the ANSYS Help System. If the required real constants are not specified, ANSYS issues a warning. A good example for describing the real constants is the 3-D beam element (element type BEAM4). As shown in Fig. 4.10, the real constants for this type consist of the cross-sectional area (AREA), area moments of inertia (IZZ and IYY), thicknesses in the $z$ and $y$-directions ( $\boldsymbol{T K Z}$ and $\boldsymbol{T K Y}$ ), etc. In some cases, a complete set of real constants may not be required; in other cases, if the real constants are not specified, ANSYS may use a default value for that particular parameter. It is recommended that the "Element Reference" be consulted for the particular element type.

For each real constant set, ANSYS requires a reference number. If it is not assigned by the user, ANSYS automatically assigns a number, as shown in Fig. 4.10.


Fig. 4.7 Deleting an element type.


Fig. 4.8 Keyoptions for the PLANE42 element.


Fig. 4.9 Keyoptions for the SOLID70 element.


Fig. 4.10 Real constants for the BEAM4 element.

Real constants are specified using the following GUI path:

## Main Menu $>$ Preprocessor $>$ Real Constants $>$ Add/EdilVDelete

This brings up the Real Constants dialog box, where clicking on Add leads to another dialog box having a list of currently defined element types. Choosing the element type for which the real constants are specified (if there are no required real constants for the selected element type, a warning window pops up) and hitting $O K$ brings up a new dialog box. The real constants for that specific element type appear; after filling in the boxes, hitting $O K$ completes this operation.

For models having multiple element types, a distinct real constant set (that is, a different reference number) is assigned for each element type. ANSYS issues a warning message if multiple element types are referenced to the same real constant set. However, there are cases where it is necessary to specify several real constant sets for the same element type. This feature is explained further by considering a beam composed of three different sections, as shown in Fig. 4.11. Although the material properties are the same, each section has a different thickness, and thus different moment of inertia values. Modeling this beam with a beam type of element, BEAM3, requires the cross-sectional properties as the real constants. Since there are three different cross-sectional properties, a different real constant set is defined for each of these cross sections; the same element type (BEAM3) in the Real Constants dialog box is selected. Different parts of the beam are meshed one at a time, directing ANSYS to use the real constant set corresponding to the specific part of the beam. This concept is further clarified when discussing Element Attributes.

CAUTION: It is the user's responsibility to keep track of units that are used in the analysis. The user does not need to give ANSYS the system of units being used. The user should decide which system of units to use and be consistent throughout the analysis (i.e., dimensions of the input, real constants, material properties and loads). ANSYS WILL NOT CONVERT UNITS. Also, the solution quantities are given in terms of the units of the input.

### 4.2.4 Material Properties

For each element type, there are a minimum number of required material properties. This number depends on the type of analysis. The material properties may be:


Fig. 4.11 A beam with three different cross-sectional areas; three real constant sets are required.

- Linear or nonlinear.
- Isotropic, orthotropic, or anisotropic.
- Temperature dependent or independent.

All material properties can be input as functions of temperature. Some properties are called linear properties because typical solutions with these properties require only a single iteration. This means that the properties being used are neither time nor temperature dependent, and thus remain constant throughout the analysis.

In the presence of variable material properties, the nonlinear characteristics of the properties must be specified. For example, a material exhibiting plasticity, viscoplasticity, etc., requires the specification of a nonlinear stress-strain relation.

A complete list of linear material properties is given in Table 4.2 (properties related to electrical and magnetic analyses are not included).

Each material property set has a reference number, the same as the element types and real constants. In problems involving different materials, the user is required to specify multiple material property sets. ANSYS identifies each material by its unique reference number. The Help System should be consulted for the specification of nonlinear material properties.

The following menu path is used to specify constant isotropic or orthotropic material properties:

Main Menu $>$ Preprocessor $>$ Material Props $>$ Material Models
This brings up the Define Material Model Behavior dialog box, as shown in Fig. 4.12. On the left side of this window, material models are listed based on their material reference numbers. On the right side, available material models are organized based on the analysis type (e.g., structural, thermal,

Table 4.2 List of material properties for structural, thermal, and fluids disciplines.

| Label | Units | Description |
| :---: | :---: | :---: |
| EX | Force/Area | Elastic modulus, element $x$-direction |
| EY |  | Elastic modulus, element $y$-direction |
| EZ |  | Elastic modulus, element $z$-direction |
| ALPX | Strain/Temp | Coefficient of thermal expansion, element $x$ direction |
| ALPY |  | Coefficient of thermal expansion, element $y$ direction |
| ALPZ |  | Coefficient of thermal expansion, element $z$ direction |
| REFT | Temp | Reference temperature (as a property) |
| PRXY | None | Major Poisson's ratio, $x-y$ plane |
| PRYZ |  | Major Poisson's ratio, $y-z$ plane |
| PRXZ |  | Major Poisson's ratio, $x-z$ plane |
| NUXY |  | Minor Poisson's ratio, $x-y$ plane |
| NUYZ |  | Minor Poisson's ratio, $y$-z plane |
| NUXZ |  | Minor Poisson's ratio, $x-z$ plane |
| GXY | Force/Area | Shear modulus, $x-y$ plane |
| GYZ |  | Shear modulus, $y-z$ plane |
| GXZ |  | Shear modulus, $x-z$ plane |
| DAMP | Time | K matrix multiplier for damping |
| MU | None | Coefficient of friction (or, for FLUID29 and FLUID30 elements, boundary admittance) |
| DENS | Mass/Vol | Mass density |
| C | Heat/Mass $\times$ Temp | Specific heat |
| ENTH | Heat/Vol | Enthalpy |
| KXX | Heat $\times$ Length/(Time $\times$ Area $\times$ Temp) | Thermal conductivity, element $x$-direction |
| KYY |  | Thermal conductivity, element $y$-direction |
| KZZ |  | Thermal conductivity, element $z$-direction |
| HF | Heat/(Time $\times$ Area $\times$ Temp) | Convection (or film) coefficient |
| EMIS | None | Emissivity |
| QRATE | Heat/Time | Heat generation rate (MASS71 element only) |
| VISC | Force $\times$ Time/Length ${ }^{2}$ | Viscosity |
| SONC | Length/Time | Sonic velocity (FLUID29 and FLUID30 elements only) |



Fig. 4.12 Dialog box for defining material models.
etc.). Figure 4.13 shows an expanded view of the material models available under Structural analysis. As observed in the figure, if a linear material response is to be used, then the user double-clicks on the Linear option to expand. After double-clicking on the Elastic option under Linear, three options are available for the user: isotropic, orthotropic, and anisotropic. Upon double-clicking on any of these options, a new dialog box appears. Figure 4.14 (left) shows the dialog box corresponding to the isotropic option. If the material properties are temperature dependent, the Add Temperature button is used for adding columns for different temperatures, as shown in Fig. 4.14 (right).


Fig. 4.13 Expanded view of the material models under the Structural discipline.


Fig. 4.14 Dialog box for isotropic properties: not temperature dependent (left) and temperature dependent (right).

### 4.2.5 Element Attributes

Every element in ANSYS is identified by the element type, real constant set, material property set, and element coordinate system. These are called element attributes. In order to create a mesh, the element type(s) must be specified a priori and the material properties (and real constants, depending on the element type) must be specified in order to obtain a solution. The element coordinate system is defined internally.

### 4.2.6 Interaction with the Graphics Window: Picking Entities

When using ANSYS through the GUI, part of the interaction between the user and the software involves picking entities or locations in the Graphics Window. These interactions are performed using the Pick Menus. Figures 4.15 and 4.16 show two examples of such menus. Picking operations are performed using the left mouse button.

When picking entities through the Pick Menu, there are five distinct fields, as shown in Fig. 4.15:

1. Pick/Unpick Field: Using the radio-buttons, the user selects whether the entities are to be picked or unpicked. This feature is useful when the user picks entities other than the intended ones. Instead of using the radio-buttons, the user may use the right mouse button to toggle between the Pick and Unpick modes.
2. Picking Style Field: By default, the user picks entities one at a time (i.e., radio-button Single in the Pick Menu). However, if the number of entities to be picked is a large number, the Single picking mode may become tedious, and one of the other modes may be preferable in such situations. Available options include:


Fig. 4.15 Pick Мenu for picking entities.


Fig. 4.16 Pick Menu for picking locations.

Box: The user draws a rectangle in the Graphics Window by holding down the left mouse button; entities located inside the rectangular box are picked.

Polygon: The user draws a polygon in the Graphics Window. Vertices of the polygon are created by single clicks on the left mouse button. The polygon is finalized when the user clicks on the first vertex created. The entities located inside the polygon are picked.

Circle: When the entities follow a radial pattern, it may be more convenient to pick them through a circular region. This option permits the user to draw a circle in the Graphics Window by holding down the left mouse button.
3. Information Field: This field provides the user with useful information such as the number of currently picked entities, maximum number of entities that can be picked, and the last entity number picked.
4. Text Field: Using this option, the user may provide text input for the entities to be picked instead of picking them in the Graphics Window. This can be done in two different formats:

List of Items: When the radio-button next to List of Items is selected (default), the user may enter a list of the entity numbers to be picked, separated by commas, in the text field.

Min, Max, Inc: When the radio-button next to Min, Max, Inc is selected, the user may enter the entity numbers to be picked in the text field in the format Minimum, Maximum, Increment. For example, if the user enters 1,5,2, then ANSYS picks entities 1, 3 and 5.
5. Action Field: This field involves familiar actions, such as:

OK: Finishes the picking operation and closes the Pick Menu.
Apply: Applies the picking performed so far while keeping the Pick Мепи active.

Reset: The picking operations performed so far are ignored and the configuration is set to the one that existed when the Pick Menu appeared.

Cancel: Closes the Pick Menu without performing picking.
Pick All: All of the items under consideration are picked and the Pick Menu is closed.

Help: Displays the Help Page related to the current operation.
Picking locations is similar to picking entities, except for slight differences in the Pick Menu. This time the menu has four fields, as shown in Fig. 4.16:

1. Pick/Unpick Field: This field is the same as explained above.
2. Information Field: Similar to the previous case, this field provides the user with useful information such as the number of currently picked locations, maximum and minimum possible picking operations, and the Working Plane and Global Cartesian coordinates of the last location picked.
3. Text Field: Using this option, the user can provide the coordinates of the location to be picked instead of picking them in the Graphics Window. This can be done in two different formats: Working Plane or Global Cartesian Coordinates. In either case, the coordinates are separated by commas.
4. Action Field: This field is the same as explained above, with exception of the absence of the Pick All button.

### 4.2.7 Coordinate Systems

### 4.2.7.1 Global Coordinate Systems

When the user starts an ANSYS session, the coordinate system (CS) is Cartesian by default. However, there are many situations where using other coordinate systems (cylindrical or spherical) is more convenient. There are four predefined coordinate systems in ANSYS: Cartesian, cylindrical, spherical, and toroidal; the first three of them are shown in Fig. 4.17.

All of these coordinate systems have the same origin (global origin) and are called global coordinate systems. Although the session starts with the Cartesian CS, the user can switch to one of the other three coordinate systems at any time. The CS currently used is referred to as the active coordinate system (active CS); any action referring to the coordinates is performed in the active CS. For example, either a Cartesian or cylindrical CS can be used to create the nodes at the locations shown in Fig. 4.18. The nodes around the unit circle are equally spaced.

In reference to a Cartesian CS, Nodes 1, 2, and 6 can easily be created because the coordinates are explicitly given as $(\mathbf{0}, \mathbf{0}, \mathbf{0}),(\mathbf{1}, \mathbf{0}, \mathbf{0})$ and $(\mathbf{0}, \mathbf{1}$, 0 ), respectively. For nodes 3,4 , and 5 , trigonometric relations can be used to calculate the $x$-, $y$-, and $z$-coordinates with a desired precision or round-off.

An alternative to the calculation of these coordinates is to change the active CS from Cartesian to cylindrical. In the cylindrical coordinate system, any reference to $x$-, $y$-, and $z$-coordinates are treated as $r, \theta$, and $z$. The coordinates of the nodes 3,4 , and 5 in the cylindrical CS are specified as $(1,22.5,0),(1,45,0)$, and $(1,67.5,0)$, respectively.

By changing the active CS, unnecessary algebraic calculations and the potential loss of accuracy are avoided.

## CARTESIAN <br> CYLINDRICAL <br> SPIIERICAL



Fig. 4.17 Cartesian (left), cylindrical (middle), and spherical (right) coordinate systems.


Fig. 4.18 Six nodes, one at the origin; the remaining five lie in a quarter-circle pattern.

The coordinates of the nodes in the Cartesian and cylindrical coordinate systems are given in Table 4.3. The "loss of accuracy" can be observed by examining the possible $x$ - and $y$-coordinates of nodes 3,4 , and 5 .

The menu path to change the active CS is given as
Utility Menu > WorkPlane > Change Active CS to
Selection of one of the top three choices in the dialog box,, i.e., global Cartesian, global cylindrical, or global spherical, completes this operation. The CS that is chosen remains active and, in turn, all the coordinates are referenced to that CS, until the user changes it.

### 4.2.7.2 Local Coordinate Systems

The global coordinate systems all share the same origin (global origin) with a predefined orientation. There are situations where changing one type of global coordinate system to a different global coordinate system does not provide enough convenience or sometimes makes it even more complicated.

Table 4.3 Nodal coordinates in Cartesian and cylindrical coordinate systems.

| Node | Cartesian |  |  | Cylindrical |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | $x$ | $y$ | $z$ | $r$ | $\theta$ | $z$ |
| 1 | 0 | 0 | 0 | 0 | 0 | 0 |
| 2 | 1 | 0 | 0 | 1 | 0 | 0 |
| 3 | 0.924 | 0.383 | 0 | 1 | 22.5 | 0 |
|  | 0.9239 | 0.3827 |  |  |  |  |
|  | 0.92388 | 0.38268 |  |  |  |  |
| 4 | 0.707 | 0.707 | 0 | 1 | 45 | 0 |
|  | 0.7071 | 0.7071 |  |  |  |  |
|  | 0.70710 | 0.70710 |  |  |  |  |
| 5 | 6 | 6 |  | 1 | 67.5 | 0 |
|  | 0.383 | 0.924 | 0 |  |  |  |
|  | 0.3827 | 0.9239 |  |  |  |  |
|  | 0.38268 | 0.92388 |  | 1 | 0 | 1 |

It may turn out that what the user really needs is to change the orientation of the CS and/or location of the origin.

In such cases, the user needs to define a CS by offsetting the origin or changing the orientation, or both. Such a coordinate system is called a local coordinate system (local CS). A local CS can be created by specifying either a location for the origin or three keypoints or nodes. Only one CS can be active at a given time. ANSYS requires that local coordinate systems have reference numbers that are greater than or equal to 11 . The menu path for creating a local CS at specified location is given as

Utility Menu > WorkPlane > Local Coordinate Systems > Create Local CS > At Specified Loc +
This brings up a Pick Menu, requesting the user to enter the coordinates of the points in the text field inside the Pick Menu, or to pick the points by clicking the mouse pointer on the Graphics Window.

After picking the origin, clicking on $\boldsymbol{O K}$ brings up the dialog box shown in Fig. 4.19. There are several text boxes to fill out in this dialog box. First is the reference number (by default, it is 11). If a local CS was defined previously, as a default, ANSYS assigns the smallest available reference number that is greater than or equal to 11 . If this reference number is not desired, the user enters the new reference number for this CS. Below the


Fig. 4.19 Dialog box for creating a Local $C S$ at a specified location.
reference number box, there is a pull-down menu for the CS type: Cartesian, cylindrical, or spherical (toroidal is not discussed herein). The coordinates of the origin of the local $C S$ with respect to the global origin should already appear in the CS type menu. Finally, rotation angles with respect to the active CS (not necessarily global Cartesian) are entered.

### 4.2.8 Working Plane

Within the ANSYS environment, regardless of the dimensionality of the problem (2-D or 3-D), calculations are performed in a 3-D space. If the problem is 2-D, then ANSYS uses the $x-y$ plane, which is the $z=0$ plane.

The Working Plane (WP) is a 2-D plane with the origin of a 2-D coordinate system (Cartesian or polar) and a display grid. It is designed to facilitate solid model generation, where many solid model entities are created by referring to the origin of the WP.

In order to view the WP, the menu path is given as

## Utility Menu > WorkPlane > Display Working Plane

A checkmark appears on the left of this menu item. Similarly, one can turn the display WP off by using the exact same menu path, resulting in the disappearance of the checkmark. By default, only the triad that is attached to the WP is shown in the Graphics Window. Viewing the grid is achieved by the menu path:

Utility Menu $>$ WorkPlane $>$ WP Settings
This brings up the WP Settings Window. Clicking on the Grid and Triad radio-button turns on both the grid and the triad; clicking on the Grid Only radio-button turns on only the grid. Hitting the Apply or $\boldsymbol{O K}$ button activates the new setting.

Using the two radio-buttons at the top permits a switch between the Cartesian and cylindrical (polar) CS. The WP can be placed at any point in the 3D space with an arbitrary orientation. There can only be one working plane at a time. By default, the WP is the $x-y$ plane of the global CS. A working plane can be defined by specifying either three points or nodes or keypoints.

At this point, defining a WP by three points is explained. The menu path is given as

## Utility Menu > WorkPlane > Align WP with > XYZ Locations

A Pick Menu appears, prompting the user to enter the coordinates of the points in the text field or pick the points by clicking the mouse pointer on the Graphics Window.

The user needs three noncolinear points to define a plane. The first point is the origin of the WP. The second point defines the WP $x$-axis along the line defined between the first and itself. The third point defines the direction of the positive WP y-axis. Two examples of these operations are illustrated in Fig. 4.20. When all three points are entered, clicking on $O K$ in the pick menu completes the definition of the WP by three points.

As shown in Fig. 4.21, an existing WP can be moved to a new location by providing offset distances in the $x$-, $y$-, and $z$-directions, which yields a WP parallel to its previous orientation. Also, an existing WP can be rotated in all three directions, as shown in Fig. 4.22. If the user rotates the WP about the $z$-axis (which is the direction normal to the WP-not to the global CS), then the WP remains in the same plane but the WP $x$ - and $y$-axes rotate within the plane. These movements can be made by the following menu path:

## Utility Menu > WorkPlane > Offset WP by Increments

This brings up the Offset window. This window requires the offset values in $\mathbf{X}, \mathbf{Y}$, and $\mathbf{Z}$. In the Offset WP window, which is used for both translation and rotation, there are six push-buttons for translation and six push-buttons for rotation. These are used for incremental translation and rotation in and around, respectively, positive and negative $x$-, $y$-, and $z$-axes. The increment is given by a sliding button right below the buttons (one for translation and one for rotation). If the display WP is turned on, the resulting incremental translation or rotation can be observed immediately (without having to hit Apply).


Fig. 4.20 Four nodes in 3-D space (top left); WP defined on the plane defined by nodes 1,2 , and 3 (top right) and by nodes 1,2 , and 4 (bottom).


Fig. 4.21 WP first moved 3 units in the $x$-direction, then 3 units in the $y$-direction, and, finally, -3 units in the $z$-direction.


Fig. 4.22 WP rotated $-45^{\circ}$ about the $x$-axis (top left), $+45^{\circ}$ about the $z$-axis (top right), and $+45^{\circ}$ about $y$-axis (bottom).

CAUTION: The $\mathbf{X}$ and $\mathbf{Y}$ refer to the WP's $x$ - and $y$-axes (not global axes) and $z$ is the direction normal to the WP (not global CS); the positive direction is established by the right-hand rule.

### 4.3 Solid Modeling

The geometrical representation of the physical system is referred to as the solid model. In model generation with ANSYS, the ultimate goal is to create a finite element mesh of the physical system. There are two main paths in ANSYS to generate the nodes and elements of the mesh: (1) direct generation and (2) solid modeling and meshing.

In direct generation, every single node is generated by entering their coordinates followed by generation of the elements through the connectivity information. Since most real engineering problems require a high number of nodes and elements (i.e., hundreds or thousands), direct generation is not feasible. Solid modeling is a very powerful alternative to direct generation.

Solid modeling involves the creation of geometrical entities, such as lines, areas, or volumes, that represent the actual geometry of the problem. Once completed, they can be meshed by ANSYS automatically (user still has control over the meshing through user-specified preferences for mesh density, etc.). A solid model can be created by using either entities or primitives.

The entities refer to the keypoints, lines, areas, and volumes. The primitives are predefined geometrical shapes.

There is an ascending hierarchy among the entities from the keypoints to the volumes. Each entity (except keypoints) can be created by using the lower ones. When defined, each entity is automatically associated with its lower entities. If these entities are created by starting with keypoints and moving up, the approach is referred to as "bottom-up" solid modeling.

When primitives are used, lower-order entities (keypoints, lines, and areas) are automatically generated by ANSYS. Since the use of primitives involves the generation of entities without having to create lower entities, it is referred to as the "top-down" approach. Boolean or similar operations can be applied to the primitives to generate the complex geometries.

The bottom-up and top-down approaches can easily be combined since one may be more convenient at a certain stage and the other at another stage. It is not necessary to declare a preference between the two approaches throughout the analysis.

### 4.3.1 Bottom-up Approach: Entities

### 4.3.1.1 Keypoints

When the bottom-up solid model generation approach is used, the user starts by generating the keypoints. The higher entities (lines, areas, and volumes) can then be defined by using the keypoints. The keypoints necessary to create a higher-order entity for modeling different parts of the geometry should be generated a priori. When areas or volumes are generated using keypoints, the intermediate entities are generated automatically by ANSYS. The creation of keypoints on the WP and in the active CS is explained herein.

The following menu path is suggested to create a keypoint on WP:
Main Menu > Preprocessor > Modeling > Create > Keypoints > On Working Plane
This brings up a Pick Menu, where ANSYS expects the user to pick points on the WP. Once the points are picked by clicking on the left mouse key, hitting on the Apply or $\boldsymbol{O K}$ button completes this task ( $\mathbf{O K}$ closes the Pick Menu). When using this option, turning on the display WP with the grid visible is highly recommended.

IMPORTANT HINT: When using the Pick Menu, picking the exact location might become a real challenge and, in turn, result in the generation of unnecessary entities. These "extra" entities might cause confusion and possible errors in the course of the solid model generation. Whenever there is a

Pick Menu, the user can hold down (no release) the left mouse button and move the pointer on the Graphics Window. This action shows the mouse pointer coordinates on the Pick Menu. When the target coordinates are found, the user can release the button to finish the picking.

The following menu path is suggested to create keypoint(s) (KP) in the active CS:

## Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Keypoints $>\operatorname{In}$ Active CS

This brings up a dialog box with four input fields for the KP number and the $x$-, $y$-, and $z$-coordinates. Once this information is supplied, hitting $O K$ creates the KP and exits from this dialog box. Alternatively, the Apply button can be clicked on and more keypoints can be created.

The coordinates defining a KP can be modified as follows:
Main Menu > Preprocessor > Modeling > Move/Modify > Single KP
This brings up a Pick Menu. First, KP is picked from the Graphics Window, or its number is typed in the text field. Then, the new location is picked or the new coordinates are typed. If a KP is modified, any mesh that is attached to that KP is automatically cleared, and any higher-order entities that are associated with that KP also are modified accordingly.

### 4.3.1.2 Lines

Lines are used for either creating a mesh with line elements or creating areas and volumes. A straight line, an arc, and a cubic spline can be created, as shown in Fig. 4.23 and 4.24.

Creating a True Straight Line: By using the following menu path, a straight line can be created regardless of the active CS. The only input needed is two keypoints. The menu path is given as

```
Main Menu \(>\) Preprocessor \(>\) Modeling \(>\) Create \(>\) Lines \(>\) Straight Line
```

This brings up a Pick Menu, requesting keypoint numbers, which can be entered through the text field or picked from the Graphics Window. Multiple lines can be generated, one at a time, without closing the Pick Menu (by using Apply button). The straight line (L1) is generated by keypoints (KP1) and (KP2).

Creating a Straight Line in the Active CS: This method creates a straight line in the active CS. If the active CS is a Cartesian CS, then the line is a true straight line. If the active CS is a cylindrical CS , then the line is a helical spiral. The menu path is given as

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Lines $>\ln$ Active Coord


Fig. 4.23 A straight line.


Fig. 4.24 An arc (left) and a cubic spline (right).

This brings up a Pick Menu, requesting the keypoints. The keypoint numbers are supplied through either the text field or by picking them using the Graphics Window. Multiple lines can be generated, one at a time, without closing the Pick Menu. It works the same way as creating a true straight line.

Creating an Arc: Creating an arc requires three keypoints. The arc is circular, regardless of the active CS. It is generated between the first and the second keypoints. The third KP defines the plane of the arc, as well as the positive curvature side. It does not have to be at the center of the curvature. The menu path is given as

## Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Arcs $>$ By End KPs \& Rad

This brings up a Pick Menu, requesting the two end keypoints. These keypoint numbers are supplied through either the text field or by picking them using the Graphics Window. Upon hitting OK in the Pick Menu, ANSYS requests the third KP , which defines the positive curvature side. After entering it the same way and hitting $O K$ in the Pick Menu, a dialog box appears. The first field is the radius and the remaining 3 are the keypoints that have already been input. Entering the radius and hitting $O K$ completes this operation.

Creating a Spline: Several keypoints (minimum 2) are needed for creating a spline. The menu path is given as

## Main Menu > Preprocessor > Modeling > Create > Splines > Spline Thru KPs

This brings up a Pick Menu, requesting the keypoints to be picked. When finished, hitting $O K$ finishes the spline creation. Multiple splines can be generated, one at a time, without closing the Pick Menu by hitting the Apply button instead of $O K$.

Once the lines are defined, areas can be created by using them.

### 4.3.1.3 Areas

Areas are used to create a mesh with area elements and to create volumes. If the geometry involves a 2-D domain, the area(s) is (are) required to be flat, lying on the $x-y$ plane. If the geometry involves 3-D bodies, then the areas that define the faces of the volume(s) can be flat or curved. A mesh created from a flat area and volumes created from flat and curved areas are shown in Fig. 4.25 and 4.26.

In bottom-up approach, areas can be created by using either keypoints or lines.

Creating an Area Using Keypoints: A minimum of 3 keypoints is required, and the maximum number allowed is 18 . If more than 3 keypoints are used, they must lie in the same plane (co-planar), as shown in Fig. 4.27. The menu path is given as

Main Menu > Preprocessor > Modeling > Create >-Areas- Arbitrary > Through KPs
which brings up a Pick Menu, requesting the keypoints to be picked. When finished, clicking on $O K$ creates the area.

CAUTION: In the PC version, it is recommended that the input window be used.

Creating an Area Using Lines: In creating an area by lines, a minimum of 3 previously defined lines are required, and the maximum number of lines allowed is 10 . If more than 3 lines are used, they must be co-planar. Lines must be given in a clockwise or counterclockwise order, and they must form a simply connected closed curve. The menu path is given as

## Main Menu > Preprocessor > Modeling > Create > -Areas- Arbitrary > By Lines

This brings up a Pick Menu, requesting the lines to be picked. When finished, hitting $O K$ creates the area.

Another commonly used method to create areas is to use primitives as part of the top-down approach; this is discussed in Sec. 4.3.2.


Fig. 4.25 An area in the $x-y$ plane (left); meshed (right).


Fig. 4.26 Volume composed of flat areas (left) and flat and curved areas (right).

### 4.3.1.4 Volumes

Volumes are used to create a mesh with volume elements (Fig. 4.28). Volumes can be created by using either keypoints or areas. If keypoints are used, the areas and lines that are associated with the volume are automatically generated by ANSYS. Two basic methods are presented below.

Creating Volumes Using Keypoints: A maximum of 8 and a minimum of 4 keypoints are required to create a volume using keypoints. Keypoints must be specified in a continuous order. If the volume has 6 faces, two of the opposite faces are required to be specified by the user, and keypoints defining both of these faces should be given in either a clockwise or counterclockwise direction.

For example, a 6 -faced volume, shown in Fig. 4.29, requires 8 keypoints. The correct counterclockwise sequence of keypoints is 1-2-6-5-4-3-7-8. Incorrect sequences, such as 1-2-6-5-4-8-7-3 or 1-2-6-5-7-3-4-8 (Fig. 4.30), have neither a clockwise nor counterclockwise sense and fail to produce the


Fig. 4.27 Coplanarity of keypoints: 4 coplanar keypoints (left) and 4 noncoplanar keypoints (right).


Fig. 4.28 A meshed volume.


Fig. 4.29 Eight keypoints (left); volume created by picking keypoints in 1-2-6-5-4-3-7-8 order (right).


Fig. 4.30 Volumes created by picking keypoints in 1-2-6-5-4-8-7-3 order (left) and 1-2-6-5-7-3-4-8 order (right).

6-faced volume. Figure 4.31 illustrates volumes with 4 and 5 faces (tetrahedron and triangular prism). The menu path is given as

Main Menu > Preprocessor > Modeling > Create > -Volumes- Arbitrary > Through KPs
This brings up a Pick Menu, requesting the keypoints to be picked. When finished, clicking on $\boldsymbol{O K}$ creates the volume.

Creating Volumes Using Areas: At least four areas (maximum of ten) are required to create a volume through areas. Areas can be specified in any order. The surface defined by the area must be continuous. The menu path is given as

## Main Menu > Preprocessor > Modeling > Create > -Volumes- Arbitrary > By Areas

which brings up a Pick Menu, requesting the areas. When finished, hitting OK creates the volume.

### 4.3.2 Top-down Approach: Primitives

The primitives are predefined geometrical shapes that enable the user to create a solid model entity (area or volume) with the execution of a single menu item. The user is not required to create keypoints and lines prior to using primitives.

### 4.3.2.1 Area Primitives

Area primitives are available for the generation of rectangles, circles, and polygons. There are different ways to create each of these primitives. The basic methods are presented here.


Fig. 4.31 A tetrahedron, 4 faces (left), and a triangular prism, 5 faces (right).

Rectangle by Dimension: The menu path is given as
Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Rectangle $>$ By Dimensions
This brings up a dialog box asking for Working Plane $\boldsymbol{X}$ and $\boldsymbol{Y}$ coordinates of the two corners of the rectangle. After filling out the four fields in this box, clicking on $\boldsymbol{O K}$ creates the rectangle in the Graphics Window.

Rectangle by 2 Corners: The menu path is given as

## Main Menu > Preprocessor > Modeling > Create > Rectangle > By 2 Corners

This brings up a Pick Menu. There are two ways to finish this action. One way is to use the four fields in the pick menu to input WP coordinates of one corner and the dimensions of the rectangle. The other method is to use the left mouse button to click on the Graphics Window to define one corner. After this, as the mouse pointer is moved, ANSYS displays possible rectangles as outlines, with the dimensions quantitatively indicated. When the user finds the right dimensions, a left-click creates the rectangular area.

Solid Circular Area: The menu path is given as

## Main Menu > Preprocessor > Modeling > Create > Circle > Solid Circle

This brings up a Pick Menu, requesting Working Plane $\boldsymbol{X}$ and $\boldsymbol{Y}$ of the center of the circle and its radius. They can be supplied either by filling out the fields in the Pick Menu or using the mouse pointer. Picking the center of the circle, moving the pointer to find the desired radius (as the mouse pointer is moved, similar to creating rectangles by dimensions, ANSYS plots the
circle's outline with the radius identified), and clicking again finalizes the circle generation.

Circular Area by Dimensions: With this option, a solid circle, annulus, circular segment (wedge) or partial annulus can be generated, as shown in Fig. 4.32. The menu path is given as

```
Main Menu > Preprocessor > Modeling > Create > Circle > By Dimensions
```

This brings up a dialog box requesting the outer and inner (optional) radii and starting and ending angles of the circular sector. All four of the geometrical parameters are defined with respect to Working Plane. If the starting and ending angles are entered as 0 and 360, ANSYS creates a full circular solid area or annulus, depending on the radius information. Otherwise, a partial solid circle (wedge) or a partial annulus is created. If the "Optional inner radius" is left blank (or entered as 0), the area is a solid one; otherwise, it's an annulus.

Polygon: The menu path is given as

## Main Menu > Preprocessor > Modeling > Create > -Areas- Polygon > By Vertices

This brings up a Pick Menu, requesting the vertices. All the vertices are on the Working Plane. Naturally, the polygon must be closed; this is achieved by picking the first point one more time after picking the last point. If the user does not pick the first point to close the polygon and hits $\boldsymbol{O K}$ in the Pick Menu, ANSYS automatically closes the polygon by defining a line between the last and the first point.

### 4.3.2.2 Volume Primitives

Volume primitives are available for generation of blocks, cylinders, prisms, spheres, or cones. There are different ways to create each of these primitives. The basic methods are presented here.

Block: A block is a rectangular prism. It is created using the following menu path:

## Main Menu > Preprocessor > Modeling > Create > Volumes > Block > By Dimensions

This brings up a dialog box requesting six coordinates, the starting and ending $x$-, $y$-, and $z$-coordinates in the active coordinate system. Figure 4.33 shows the isometric view of a block created using $x_{1}=y_{1}=z_{1}=0, x_{2}=1, y_{2}=$ 2 , and $z_{2}=3$.


Fig. 4.32 Circular area primitives.


Fig. 4.33 Isometric view of a block created using $x_{1}=y_{1}=$ $z_{1}=0, x_{2}=1, y_{2}=2$, and $z_{2}=3$.

Cylinder: The user can create solid or hollow cylinders that encompass either the entire angular range or a part thereof. Cylinders are created using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Volumes > Cylinder > By Dimensions
Six parameters are requested in the dialog box:
RAD1 and RAD2: Outer and inner radii of the cylinder. If RAD2 is not specified (or specified as zero), then the cylinder is solid; otherwise, it's hollow.

Z1 and $Z 2$ : Starting and ending $z$-coordinates.
THETA1 and THETA2: Starting and ending angles, measured in degrees, with the active coordinate system $z$-axis defining the axis of rotation.

Figure 4.34 (left) shows an isometric view of a hollow cylinder created using RAD1 $=1$, RAD2 $=0.5, \mathbf{Z 1}=0, \mathbf{Z 2}=2, \boldsymbol{T H E T A 1}=0$ and $\boldsymbol{T H E T A} 2=$ 360. When the parameter THETA1 is changed to $135^{\circ}$ and the other parameters are kept the same, the partial hollow cylinder shown in Fig. 4.34 (right) is created.

Prism: Regular prisms are created using this option. A regular prism is a volume with a constant polygonal cross section in the Working Plane zdirection. The menu path for creating prisms is given as

## Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Volumes $>$ Prism $>$ By Side Length

This brings up a dialog box requesting the starting and ending $z$-coordinates $Z 1$ and $Z 2$, respectively; the number of sides (NSIDES); and the length of each side (LSIDE). The center of the polygonal area coincides with the Working Plane origin. Figure 4.35 (left) shows the isometric view of a prism created using $Z 1=0, Z 2=2$, NSIDES $=3$, and $L S I D E=1$. Fig. 4.35 (right) shows the prism when the parameter NSIDES is changed to 6 .

Sphere: The user can create solid or hollow spheres by using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Volumes $>$ Sphere $>$ By Dimensions
Four parameters are requested in the dialog box:
RAD1 and RAD2: Outer and inner radii of the cylinder. If RAD2 is not specified (or is specified as zero), then the sphere is solid; otherwise, it's hollow.

THETA1 and THETA2: Starting and ending angles, measured in degrees, with the Working Plane $z$-axis defining the axis of rotation.

Figure 4.36 (left) shows an isometric view of a solid sphere created using RAD1 $=1$, RAD2 $=0$, THETA1 $=0$, and THETA2 $=360$. The partial hollow sphere shown in Fig. 4.36 (right) is created when the parameters THETA1 and THETA2 are changed to $90^{\circ}$ and $270^{\circ}$, respectively, while the other parameters are kept same.

Cone: Complete or partial cones may be created using this option by following the menu path:

[^0]

Fig. 4.34 Isometric view of a hollow cylinder (left) created using RAD1 $=1$, RAD2 $=0.5, Z 1=0, Z 2=2$, THETA1 $=0$, and THETA2 $=360$ and a partial hollow cylinder (right) when the parameter THETA1 is changed to 135 .


Fig. 4.35 Isometric view of a prism (left) created using $\boldsymbol{Z 1}=0, \boldsymbol{Z} \mathbf{Z}$ $=2$, NSIDES $=3$, and $\operatorname{LSIDE}=1$ and the prism (right) when the parameter NSIDES is changed to 6 .


Fig. 4.36 Isometric view of a solid sphere (left) created using RAD1 $=1$, RAD2 $=0$, THETA1 $=0$, and THETA $2=360$ and the partial hollow sphere (right) when the parameters THETA1 and THETA2 are changed to $90^{\circ}$ and $270^{\circ}$, respectively.

In the dialog box, six parameters are requested:
RBOT and RTOP: Bottom and top radii of the cone. If RTOP is not specified (or is specified as zero), then a complete cone is generated. If a nonzero RTOP is specified, then the volume generated is a conical section with parallel top and bottom sides.
$\mathbf{Z 1}$ and $\mathbf{Z 2}$ : Starting and ending $z$-coordinates.
THETA1 and THETA2: Starting and ending angles, measured in degrees, with the Working Plane $z$-axis defining the axis of rotation. It is used for creating conical sections.

Figure 4.37 (left) shows an isometric view of a cone created using $R B O T=$ $1, R T O P=0, Z 1=0, Z 2=3, T H E T A 1=0$, and $T H E T A 2=360$. The partial conical section shown in Fig. 4.37 (right) is created when the parameters RTOP, Z2, and THETA1 are changed to $0.5,2$, and 135, respectively, while the other parameters are kept same.

### 4.4 Boolean Operators

Many engineering problems possess a complex geometry, making model generation a real challenge. However, the solid model entities can be subjected to certain operations that make model generation much easier. These operations, referred to as Boolean operations, utilize logical operators such as add, subtract, divide, etc. The Boolean operators are applied to generate more complex entities using simple entities (see Fig. 4.38).

### 4.4.1 Adding

The areas to be added must be co-planar (lie in the same plane). As shown in Fig. 4.39, the areas (or volumes) must have either a common boundary or an overlapping region. The original areas or volumes that are added will be deleted unless otherwise enforced by the user. The addition of areas or volumes results in a single (possibly complex geometry) entity, as shown in Fig. 4.40.

Adding entities can be performed by the following menu paths:

```
Main Menu > Preprocessor > Modeling > Operate > Add > Lines
Main Menu > Preprocessor > Modeling > Operate > Add > Areas
Main Menu > Preprocessor > Modeling > Operate > Add > Volumes
```



Fig. 4.37 Isometric view of a cone (left) created using RBOT $=1$, RTOP $=0, \mathbf{Z 1}=0, \mathbf{Z 2}=3$, THETA1 $=0$, and THETA $2=360$ and the partial conical section (right) when parameters RTOP, Z2, and THETA1 are changed to $0.5,2$, and 135 , respectively.


Fig. 4.38 Examples of entities that can co-exist in 3-D space.


Fig. 4.39 Two areas with a common boundary.


Fig. 4.40 Two areas added to produce one area.

### 4.4.2 Subtracting

Entities can be subtracted from each other to obtain new entities. Subtracting entities can be executed through the menu paths given below:

```
Main Menu > Preprocessor > Modeling > Operate > Subtract > Lines
Main Menu > Preprocessor > Modeling > Operate > Subtract > Areas
Main Menu > Preprocessor > Modeling > Operate > Subtract > Volumes
```

This brings up a Pick Menu, requesting the user to pick or enter the base entity from which to subtract. The user picks the entities to be subtracted and clicks on the $\boldsymbol{O K}$ button to complete the operation.

### 4.4.3 Overlap

This operation joins two or more solid model entities to generate three or more entities forming a union of the entire original group of entities, as shown in Fig. 4.41. It is similar to the $\boldsymbol{A d d}$ operation. The only difference between the two is that internal entities are generated in the overlapping operation.

This operation can be executed through the following menu paths:

```
Main Menu > Preprocessor > Modeling > Operate > Overlap > Lines
Main Menu > Preprocessor > Modeling > Operate > Overlap > Areas
Main Menu > Preprocessor > Modeling > Operate > Overlap > Volumes
```

This brings up a Pick Menu asking for the entities to be overlapped. Picking the entities followed by hitting $O K$ completes the operation.


Fig. 4.41 Two areas, A1 and A2 (left); the result of adding A1 and A2 (middle); and the result of overlapping A1 and A2 (right).

### 4.4.4 Gluing

This operation is used for connecting entities that are "touching" but not sharing any entities. If the entities are apart from or overlapping each other, gluing cannot be used. The glue operation does not produce additional entities of the same dimensionality but does create new entities that have one lower dimensionality. This operation can be executed through the following menu paths:

## Main Menu > Preprocessor > Modeling > Operate > Glue > Lines <br> Main Menu > Preprocessor > Modeling > Operate > Glue > Areas <br> Main Menu $>$ Preprocessor $>$ Modeling $>$ Operate $>$ Glue $>$ Volumes

Before gluing the two areas shown in Fig. 4.42, there are two lines at the interface between Area 1 (A1) and Area 2 (A2). One of these lines is attached to A1, defined by keypoints $\mathbf{2}$ and 3, and the other one is attached to A2, defined by keypoints $\mathbf{5}$ and $\mathbf{8}$. Before gluing, these two areas do not "know" of each other's existence because they do not "truly" share any entities. Gluing makes sure that they share entities. After gluing, there are two lines along the right vertical side of A1, and two lines along the left side of A2. The lines along the right vertical side of A1 are defined by keypoints 3 and $\mathbf{8}$ and keypoints $\mathbf{8}$ and $\mathbf{2}$ whereas the lines along the left vertical side of A2 are defined by keypoints $\mathbf{2}$ and $\mathbf{8}$ and keypoints $\mathbf{5}$ and $\mathbf{2}$. After gluing, the two areas share one line and two keypoints.

### 4.4.5 Dividing

A solid model entity can be divided into smaller parts by using other solid model entities. By default, a divided solid model entity is deleted after the operation. There is a wide range of choices for this operation. Some of the available options are presented in Fig. 4.43-4.46.


Fig. 4.42 Two areas with a common boundary plotted with line numbers (left); they do not share any lines as area 1 (A1) is defined by lines 1 through 4 and area 2 (A2) is defined by lines 5 through 8. After gluing (right), the areas share line 9 .


Fig. 4.43 A cylindrical volume is divided into two smaller cylindrical volumes by an area.


Fig. 4.44 Dividing A1 by A2 (left) produces A3 and A4 (right).


Fig. 4.45 Dividing an area by a line (requires the dividing line to be in the same plane as the area).


Fig. 4.46 Dividing L1 by L2 (left) results in L3 and L4 (right).

The menu paths for these operations are given as

## Volume by Area

Main Menu > Preprocessor > Modeling > Operate > Divide > Volume by Area

## Area by Volume

Main Menu > Preprocessor > Modeling > Operate > Divide > Area by Volume

```
Area by Area
Main Menu \(>\) Preprocessor \(>\) Modeling \(>\) Operate \(>\) Divide \(>\) Area by Area
Area by Line
Main Menu \(>\) Preprocessor \(>\) Modeling \(>\) Operate \(>\) Divide \(>\) Area by Line
```


## Line by Volume

Main Menu $>$ Preprocessor $>$ Modeling $>$ Operate $>$ Divide $>$ Line by Volume

## Line by Area

Main Menu $>$ Preprocessor $>$ Modeling $>$ Operate $>$ Divide $>$ Line by Area

## Line by Line

Main Menu > Preprocessor > Modeling > Operate > Divide > Line by Line

### 4.5 Additional Operations

### 4.5.1 Extrusion and Sweeping

In addition to Boolean operators, extrusion and sweeping of the existing entities can be used to generate higher entities. By extruding (dragging) an entity or by sweeping (rotating) it about an axis, one can create a new solid model entity, which is one order higher than the original one (e.g., lines from keypoints, volumes from areas). Extrusion is a subset of sweeping where areas are dragged along lines to create volumes.

The commonly used features of extrusion and sweeping operations are described in Fig. 4.47-4.54. Following are the menu paths used for these operations:

## Creating Lines by Rotating a Keypoint About an Axis

> Main Menu > Preprocessor > Modeling > Operate > Extrude/Sweep > -Keypoints- About Axis

## Creating Lines by Sweeping a Keypoint Along a Path

Main Menu > Preprocessor > Modeling > Operate > Extrude/Sweep >-Keypoints- Along Lines

## Creating Areas by Rotating Lines About an Axis

Main Menu > Preprocessor > Modeling > Operate > Extrude/Sweep > -Lines- About Axis
Creating Areas by Sweeping Lines Along a Path
Main Menu $>$ Preprocessor $>$ Modeling $>$ Operate $>$ Extrude/Sweep $>-$ Lines- Along Lines


Fig. 4.47 Keypoints 1 and 2 (left) are rotated $+60^{\circ}$ around the $z$-axis in two increments of $30^{\circ}$ to create the lines (right).


Fig. 4.48 Front view of an arc (left); the arc is rotated $+60^{\circ}$ about the $y$-axis in two increments of $30^{\circ}$ to create the curved areas (shown in oblique view on the right).


Fig. 4.49 Front view of a straight line (left); the line is rotated $+60^{\circ}$ about the $y$-axis in two increments of $30^{\circ}$ to create the curved areas (shown in oblique view on the right).


Fig. 4.50 Front view of a straight line (left); the line is rotated $+120^{\circ}$ about the $y$-axis in two increments of $60^{\circ}$ to create the curved areas (shown in oblique view on the right).


Fig. 4.51 Front view of an area (left); the area is rotated $+120^{\circ}$ about the $y$-axis in two increments of $60^{\circ}$ to create the volumes (shown in oblique view on the right).


Fig. 4.52 Oblique view of a path defined by lines and an area to be swept along the path (left) and the volume created by sweeping the area along the path (right).


Fig. 4.53 Oblique view of an area to be extruded along its normal (left) and oblique view of the volume created by extrusion of the area along its normal (right).


Fig. 4.54 Oblique view of an area to be offset in $x, y$, or $z$ (left) and oblique view of the volume created by offsetting the area in the $z$ direction (right).

## Creating Volumes by Rotating Areas About an Axis

Main Menu > Preprocessor > Modeling > Operate > Extrude/Sweep > -Areas- About Axis

## Creating Volumes by Sweeping Areas Along a Path

Main Menu > Preprocessor > Modeling $>$ Operate $>$ Extrude/Sweep > -Areas- Along Lines

## Creating Volumes by Extruding Areas

Main Menu > Preprocessor > Modeling > Operate > Extrude/Sweep > -Areas- Along Normal

## Creating Volumes by Offsetting Areas

Main Menu > Preprocessor > Modeling > Operate > Extrude/Sweep > -Areas- By XYZ Offset

### 4.5.2 Moving and Copying

Previously created entities can be moved or copied. Also, if a repeated symmetry or skew-symmetry exists in the geometry, the user can create a representative entity to create the geometry by copying it to a new location. Representative applications are described in Fig. 4.55-4.57.

The common menu paths for moving entities are specified as

```
Main Menu > Preprocessor > Modeling > Move/Modify > -Keypoints- Single KP
Main Menu > Preprocessor > Modeling > Move/Modify > Lines
Main Menu > Preprocessor > Modeling > Move/Modify > -Areas- Areas
Main Menu > Preprocessor > Modeling > Move/Modify > Volumes
```

The common menu paths for copying entities are specified as
Main Menu $>$ Preprocessor $>$ Modeling $>$ Copy $>$ Keypoints
Main Menu $>$ Preprocessor $>$ Modeling $>$ Copy $>$ Lines
Main Menu $>$ Preprocessor $>$ Modeling $>$ Copy $>$ Areas
Main Menu $>$ Preprocessor $>$ Modeling $>$ Copy $>$ Volumes

### 4.5.3 Keeping/Deleting Original Entities

During the performance of Boolean-type operations, there are input entities (e.g., original areas to be added or, when dividing a line with a volume, the original line and volume) and output entities. By default, ANSYS will delete the input entities, keeping only the output entity. However, the input entities can be "kept" through the menu path:

Main Menu > Preprocessor > Modeling > Operate > Settings


Fig. 4.55 Original line (left) and copies of the line created by offsets in $x$ (L2), in $y$ (L3), and in both $x$ and $y$ (L4)(right).


Fig. 4.56 Original area (left) and copies of the area created by offsets in $x$ (A2), in $y$ (A3), and in both $x$ and $y$ (A4) (right).


Fig. 4.57 The first volume is copied three times with 0.4 unit offset in the $x$-direction and then reflected with respect to the $x-z$ plane

This brings up a dialog box requesting the user to specify certain settings. The first setting option controls whether the input entities will be kept or deleted. Answering Yes instructs ANSYS to keep the input entities; otherwise, they are deleted.

### 4.5.4 Listing Entities

In most cases, plotting is an effective way to quickly examine the model. However, if there are unexpected errors or if the model is not what the user intended to create, it may be difficult to identify what went wrong. In such cases, the user can examine the model in a more accurate way by listing the entities. ANSYS provides options for listing solid model entities with detailed information. All of the lists are given in a new window (which can be saved to disk or printed) where the entities are sorted by their reference numbers in ascending order. Lists for the solid model entities can be obtained by following the menu paths:

Utility Menu $>$ List $>$ Keypoints $>$ Coordinates only
Utility Menu > List > Lines > Attribute format
Utility Menu > List > Areas
Utility Menu > List > Volumes

### 4.5.5 Deleting Entities

During solid modeling, it is very common for the user to create unintended solid modeling entities. These extra entities might make the modeling phase confusing and may potentially cause errors. In order to eliminate this possibility, the user should "clean up" the model by deleting these entities. The hierarchy of the solid model entities is important in that the entity (or entities) must not be used for the definition of any higher-order entities in order to be deleted. For example, the existence of an area automatically implies that lines and keypoints are attached to this area. None of the lines can be deleted as long as the area exists. The area must first be deleted, then the lower-order entities can be deleted. Similarly, a KP cannot be deleted as long as the line(s) to which the KP is attached exist(s). Only after the line(s) is(are) deleted, can the KP can be deleted. Solid model entities can be deleted in two different methods:

1. Delete the entity without deleting the lower-order entities that are attached to it.
2. Delete the entity and all the lower-order entities that are attached to it. In this case, if some of the lower-order entities are associated with other entities, they will not be deleted.

The following menu paths are used for these two methods:

## To Delete Entities Only

```
Main Menu > Preprocessor > Modeling > Delete > Keypoints
Main Menu \(>\) Preprocessor \(>\) Modeling \(>\) Delete \(>\) Lines Only
Main Menu > Preprocessor > Modeling > Delete > Areas Only
Main Menu \(>\) Preprocessor \(>\) Modeling \(>\) Delete \(>\) Volumes Only
```


## To Delete Entities and Below

Main Menu $>$ Preprocessor $>$ Modeling $>$ Delete $>$ Lines and Below
Main Menu $>$ Preprocessor $>$ Modeling $>$ Delete $>$ Areas and Below
Main Menu $>$ Preprocessor $>$ Modeling $>$ Delete $>$ Volumes and Below

### 4.6 Viewing a Model

ANSYS provides a very robust graphic utility to view solid model entities, nodes, elements, material properties, boundary constraints, loads, and results. The graphics-related utilities are accessible through the Plot and PlotCtrls (stands for Plot Controls) submenus under the Utility Menu. All of the entities can be viewed through the Plot submenu. However, the PlotCtrls submenu (as its name indicates) provides many options that enhance the use of the plot utility for many different purposes, such as plotting the numbers associated with entities, plotting the entities in different colors, ${ }^{1}$ and viewpoint and viewing angle adjustments.

For the sake of brevity, only the most frequently used items are explained in detail here. However, many more features are discussed in the examples.

### 4.6.1 Plotting: PanZoom, and Rotate Functions

The Pan-Zoom-Rotate tool is a very effective function of ANSYS for manipulating the view by panning, zooming, and rotating the model. The following menu path is used to activate this function:

Utility Menu > PlotCtrls > Pan, Zoom, Rotate
The Pan-Zoom-Rotate window appears, as shown in Fig. 4.58. There are eight different fields in this window:

1. Active Window Field: The Graphics Window can be divided into "sub"windows (up to 5) in ANSYS; only one of them can be the "active" window at any one time. This button identifies which window(s) are to be affected by the operations performed within the Pan-Zoom-Rotate window.

[^1]

Fig. 4.58 The eight fields in the Pan-Zoom-Rotate window.
2. Viewing Direction Field: This group of buttons changes the viewpoint. Clicking on the Top button will redraw the model (or entities) as seen from the top. In ANSYS, "top" corresponds to the positive global Ydirection. Similarly, "front" and "right" will redraw the model as seen from the positive global Z- and X-directions, respectively. The Iso and Obliq buttons redraw the model as seen from a point that lies on a line that passes through the origin and $(1,1,1)$ and $(1,2,3)$, respectively. Finally, the $\boldsymbol{W} \boldsymbol{P}$ button redraws the model by taking the positive Working Plane $z$-direction as the front of the model.
3. Zoom Field: Provides different zooming methods:

Zoom: Clicking on this button, followed by a single left-click, chooses the center of the region of interest. After the first click, moving the mouse toward and away from the center will display a moving square outline of the potential target region that the user would like to zoom in. Once decided, a second left-click will zoom in to the region indicated by the outline.

Box Zoom: This function works in a similar way. The user picks two corners of the zoom-in region. After picking the first corner by clicking the left mouse button, moving the mouse over the Graphics Window will show a moving outline of the potential zoom-in region. A second click will pick the second corner and ANSYS will redraw the zoom-in region.

Backup: Clicking on this button redraws the model in the previous viewing configuration.

Win Zoom: This button works like the Box Zoom button except that after picking the first point, ANSYS locks the aspect ratio of the potential zoom-in region at the same values as the aspect ratios of the active window (the redraw of the zoom-in region will fit perfectly in the window).
4. Pan/Zoom Field: The arrow buttons pan the model in the indicated directions and the dots zoom in and out. A small dot indicates zooming out and a large dot indicates zooming in. The Sliding Rate Control Bar, explained below, dictates the rate at which pan and zoom actions operate.
5. Rotate Field: These six buttons rotate the model about "Screen" $x$-, $y$-, and $z$-directions. The "screen origin" is the center of the active window. The positive "Screen" $x$-direction starts from the center of the window and extends to the right. Likewise, the positive "Screen" $y$ - and $z$ directions start at the center of the window and extend to, respectively, the top and out (of the monitor).
6. Rate Control Field: The Sliding Bar controls the rate of pan, zoom, and rotate that is performed in the active window. The range is from 1 to 100 (rate 1 pans/zooms at a smaller rate than rate 100 would).
7. Dynamic Mode Field: By clicking on this radio button, the user toggles on/off the option to pan and rotate dynamically. When the Dynamic Mode is active, the mouse pointer changes shape when it is over the Graphics Window. Pressing the left mouse button (without releasing) and moving around in the Graphics Window pans the model. Similarly, the right mouse button is used for rotating the model dynamically.
8. Action Field: Includes four action buttons:

Fit: Fits the whole model in the active window.
Reset: Restores the default orientation and size for viewing (front view).
Close: Closes the Pan-Zoom-Rotate window.
Help: Brings the help page for Pan-Zoom-Rotate window.

### 4.6.2 Plotting/Listing Entities

The following menu paths are used to plot and list the solid model (keypoints, lines, areas, and volumes) and mesh (nodes and elements) entities:

```
Utility Menu > Plot > Keypoints > Keypoints
Utility Menu > Plot > Lines
Utility Menu > Plot > Areas
Utility Menu > Plot > Volumes
Utility Menu > List > Volumes
Utility Menu > Plot > Nodes
Utility Menu > List > Nodes
Utility Menu > Plot > Elements
Utility Menu > List > Elements > Nodes + Attributes
```

The resulting plots are displayed in the Graphics Window and can be examined using the Pan-Zoom-Rotate window discussed in the previous section.

### 4.6.3 Numbers in the Graphics Window

Whenever an entity is being created, ANSYS either asks for a reference number or assigns the lowest available number for that type of entity. Therefore, every entity differs from the other entities of the same type by this reference number. When plotting these entities in the Graphics Window, by default, ANSYS will not show the entity numbers. Often times, it is important for the user to see the numbers printed when plotting entities. This can be done using the following menu path:

Utility Menu > Ploctrrls > Numbering
which brings up the Plot Numbering Controls dialog box, as shown in Fig. 4.59. The entity numbers for keypoints, lines, areas, volumes, and nodes can simply be turned on by placing a checkmark in the corresponding boxes. Element numbers can be turned on using the Elem/Attrib numbering pulldown menu. Instead of the element numbers, the user can display element attribute numbers (element type, real constant, and material) using the same option. Also, colors may be assigned to each entity number for more convenient viewing. The [/NUM] Numbering shown with pull-down menu in this dialog box allows the user to plot numbers with or without color assignments, as well as to plot using colors only (without numbers).

### 4.7 Meshing

As mentioned previously (Sec. 4.3), the mesh of the geometry under consideration may be generated directly, i.e., generation of nodes and elements,


Fig. 4.59 Plot Numbering Controls dialog box.
one at a time. However, this may prove to be a challenging task. Almost always Solid Modeling constitutes a part of the finite element analysis. Thus, the sole purpose of Solid Modeling is to create the mesh of the geometry, as conveniently and efficiently as possible. Once the Solid Model is completed, the user is ready to perform meshing. Regardless of whether a Solid Model is generated or not, the meshing can be performed only after the specification of element type(s). ANSYS offers several convenient options to assist in meshing. These include Automatic Meshing, Smart Sizing, and Mapped Meshing. In the following subsections, topics related to meshing are discussed in more detail.

### 4.7.1 Automatic Meshing

One of the most powerful features of ANSYS is automatic mesh generation. ANSYS meshes the solid model entities upon execution of an "appropriate" single command. With automatic meshing, the user can still provide specific preferences for mesh density and shape. If no preferences are specified by the user, ANSYS uses the default preferences. The following menu paths are used for automatic mesh generation after solid model generation:

Mesh Using Line Elements: This option is used for models utilizing onedimensional elements, such as trusses and beams. It requires existing lines. The following menu path is used to mesh lines:

## Main Menu > Preprocessor > Meshing > Mesh > Lines

This brings up a Pick Menu asking the user to either enter the line number(s) through the text field or pick line(s) from the Graphics Window. When all the lines are input (picked), hitting $\boldsymbol{O K}$ in the Pick Menu generates the mesh.

Mesh Using Area Elements: This option is used for models utilizing 2-D elements, and it requires existing areas. The following menu path is used to mesh areas:

## Main Menu > Preprocessor > Meshing > Mesh > Areas > Mapped $>3$ or 4 Sided Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Free

Meshing can be accomplished through either the Mapped or Free Meshing methods. If free meshing is chosen, the second menu path is used, bringing up a Pick Menu asking the user to either enter the area number(s) through the text field or pick area(s) from the Graphics Window. When all the areas are input (picked), hitting $O K$ in the Pick Menu generates the mesh. The Mapped meshing option is discussed in a later subsection.

Mesh Using Volume Elements: This option is used for models using 3-D elements, and it requires existing volumes.

```
Main Menu > Preprocessor > Meshing >Mesh > Volumes > Mapped > 4 to 6 Sided
Main Menu > Preprocessor > Meshing > Mesh > Volumes > Free
```

This brings up a Pick Menu asking the user to either enter the volume number(s) through the text field or pick volumes(s) from the Graphics Window. When all the volumes are input (picked), hitting $O K$ in the Pick Меnu will generate the mesh.

ANSYS allows the user to control the mesh density of the domains defined by solid model entities. The desired mesh density can be achieved by:

- Defining a target element edge size on the domain boundaries.
- Defining a default number of element edges on all lines.
- Defining the number of element edges on specific lines.
- Using smart sizing.
- Using mapped meshing.

These methods are discussed in detail in the following subsections.

### 4.7.1.1 Specifying Mesh Density Globally

There are two approaches for enforcing the mesh density globally. The first one involves specification of the element edge size; ANSYS attempts to generate a mesh with all elements having edge sizes as close as possible to the specified value. The second possibility is to specify a fixed number of elements along all the lines within the solid model. The following menu path is used for specifying the mesh density globally:

## Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Global > Size

This brings up the Global Element Sizes dialog box with two input parameters: SIZE and NDIV. SIZE denotes the target element edge length, and NDIV is the target number of elements along the lines. If SIZE is specified, NDIV is ignored. The following example explains these concepts. Consider a square area with sides 5 units long, as shown in Fig. 4.60 (left). If the global element size is specified as $1(S I Z E=1)$, the mesh shown in Fig. 4.60 (middle) is generated with each element having an edge size of 1 unit. If the user chooses to the specify the number of elements along lines instead of element sizes, then SIZE is left untouched (zero), and NDIV is set to a specific value, say 8. As a result of this operation, the mesh shown in Fig. 4.60 (right) is generated, with 8 elements along each line.

Specification of mesh density globally works well when the geometry of the problem is regular, with aspect ratio close to one. When domains of irregular shapes are considered, applying the same meshing targets to lines of different sizes results in meshes with high aspect ratios, leading to potentially erroneous results. Therefore, the techniques explained in the following subsections are more desirable.

### 4.7.1.2 Specifying Number of Element Edges on Specific Lines

When the geometry of the problem is irregular, i.e., not basic shapes such as triangles and rectangles, specifying the number of element edges along specific lines may be a good way to avert possible meshing problems. This strategy also helps to refine the mesh around regions where it may be crucial for accuracy. Similarly, certain regions in the geometry may not be critical, and keeping the mesh around these regions may help reduce the computational cost without losing accuracy. The number of element edges on specific lines can be specified using the following menu path:

## Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Lines > Picked Lines

which brings up the Pick Menu for line picking. After the user picks the lines and clicks on OK, the Element Sizes on Picked Lines dialog box appears. The second parameter, NDIV, dictates how many elements will be


Fig. 4.60 A square area (left) meshed with global element size (SIZE) specified (middle) and number of line divisions (NDIV) specified globally (right).
placed along the picked lines. The third parameter, SPACE, which stands for spacing ratio, is important when a mesh graded (biased) toward a direction is desired. The default value for SPACE is 1 (no bias, uniform spacing). If it is positive, the spacing is biased from one end of the line to the other end. If it is negative, then the bias is from the center toward the ends. Its magnitude defines the ratio of the largest division size to the smallest. These concepts are explained in the following examples. Consider the square area used for the example in the previous subsection, shown in Fig. 4.61 (left) with line numbers. Using the menu path above, NDIV is specified to be 5 for lines 2 and 4 , and 10 for lines 1 and 3 . After this operation, lines are plotted with the specified divisions clearly visible [Fig. 4.61 (middle)]. Meshing of the area produces the one shown in Fig. 4.61 (right). The same example is considered, this time with the specification of spacing ratios. The goal is to have a mesh graded from coarse at the center to fine at the edges in the $x$ direction, and from coarse at the top to fine at the bottom (in the $y$ direction). The same number of divisions is used for all the lines with a value of $8(N D I V=8)$. Using the menu path given above, spacing ratios (SPACE) for lines $1,2,3$, and 4 are specified as $-4,4,-4$, and 0.25 , respectively. The line plot after this operation is shown in Fig. 4.62 (left), and the corresponding mesh is given in Fig. 4.62 (middle). It is worth noting the reason why the parameter $\operatorname{SPACE}$ is different for lines 2 and 4 . Figure 4.62 (right) shows the line plot with the keypoint numbers. Line 2 is defined from keypoint 2 to keypoint 3 (from bottom to top) whereas line 4 is defined from keypoint 4 to keypoint 1 (from top to bottom). When SPACE is positive and greater than 1 , its value defines the ratio of the division length at the end of the line to the length of the division at the beginning of the line.


Fig. 4.61 A square area (left) with number of line divisions specified at specific lines (middle) and the resulting mesh (right).


Fig. 4.62 Biased line divisions (left), resulting mesh (middle), and lines with keypoint numbers plotted (right).

### 4.7.1.3 Smart Sizing

Instead of specifying the number of line divisions or element edge sizes, one can use the ANSYS "smart sizing" feature, where the mesh density is specified in a cumulative sense. In this method, the user specifies a level of refinement that ranges from $\mathbf{1}$ to $\mathbf{1 0}$; the smaller the number, the finer the mesh. To use smart sizing, follow the menu path given below:

## Main Menu $>$ Preprocessor $>$ Meshing $>$ Size Cntrls $>$ SmartSize $>$ Basic

This brings up a dialog box with a pull-down menu asking the user to choose a level of refinement. Selecting the level, followed by hitting $\boldsymbol{O K}$, activates the smart sizing. Now, the user is ready to mesh the solid model entities. As this option uses meshing options involving advanced geometry, there is no easy way to explain how it works. Therefore, it is suggested that the user experiment with it, and build a knowledge base that will be helpful later on.

### 4.7.1.4 Mapped Meshing

Another very commonly used (by experienced ANSYS users) meshing method is Mapped Meshing. The mapped meshing concept is valid only in
two- and three-dimensional problems (no line elements). The solid model entities (areas and volumes) meshed with this option use quadrilateral area elements or hexahedral (brick) volume elements.

The reason why mapped meshing is desirable is that it generates regular, thus computationally well-behaving, meshes. Not every area or volume can be mapped meshed. The areas or volumes to be mapped meshed must be "regular." This regularity is governed by two properties of the solid model entity: the number of sides (lines for areas and areas for volumes) and number of divisions on opposite sides (opposite sides must have an equal number of divisions). For areas, the acceptable number of sides is 3 or 4. If the area has 3 sides (defined by 3 lines), then the number of divisions in all 3 lines must be equal and even. If 4 lines define the area, as stated before, the lines on opposite sides must have the same number of divisions. These considerations are similar for mapped meshing of volumes. The number of areas that define the volume must be either 4 (tetrahedron), 5 (prism), or 6 (hexahedron). The number of divisions on opposite sides must be equal. If 4 or 5 areas define the volume, the number of divisions on the triangular areas must be equal and even.

To use mapped meshing, follow the menu path given below:

## Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 Sided Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Volumes $>$ Mapped $>4$ to 6 Sided

which brings up a Pick Menu for area picking. After the areas are picked and the $O K$ button is pressed, the mesh is generated. Figure 4.63 (left) shows a triangular area with corresponding free and mapped meshes given in Fig. 4.63 (middle) and 4.63 (right), respectively. It is clear from these figures that the mapped mesh delivers elements with controlled and desirable aspect ratios.

If the areas or volumes do not have the required number of sides that are given above, there might still be a way to "mapped mesh" these entities. For this, the user looks for sides that could be considered as a single side when combined. This way the number of sides can be reduced to the required numbers. This operation is performed through "concatenating" lines (for meshing of areas with more than 4 sides) and areas (for meshing of volumes with more than 6 sides). Line and area concatenations are performed using the following menu paths:

[^2]

Fig. 4.63 Triangular area (left) and corresponding free (middle) and mapped (right) meshes.
which brings up a Pick Menu for lines to be concatenated. The concatenation is explained through the following example. Consider the irregular area, shown in Fig. 4.64 (top), with line numbers plotted. Free meshing of this area produces the mesh given in Fig. 4.64 (middle), with elements having large aspect ratios. As observed from Fig. 4.64 (top), the area to be meshed is enclosed by 7 lines (sides). When mapped meshing areas, the maximum number of sides is 4 . Therefore, if the user wants to mapped mesh this area, line concatenations must be performed. For this purpose, lines 1,2 , and 3 are concatenated to produce a new line (line 8). Also, lines 4 and 5 are concatenated, producing line 9 . With these concatenations, the number of sides defining the area is reduced from 7 to 4 and mapped meshing is possible. After specifying the number of divisions on lines 6 and 7 as 3 , mapped meshing is performed by using the menu path given above, producing the mesh given in Fig. 4.64 (bottom), with elements having acceptable aspect ratios.

### 4.7.2 Manipulation of the Mesh

### 4.7.2.1 Changing Element Attributes

The user can change the attributes of the elements after the mesh is generated. This is achieved by using the following menu path:

```
Main Menu > Preprocessor > Modeling > Move/Modify > Elements > Modify Attrib
```

This will bring up a Pick Menu asking the user to pick element(s) from the Graphics Window. Once the element(s) are selected, clicking on $O K$ in the Pick Menu leads to a dialog box with two fields: (1) a pull-down menu containing the attributes, and (2) a new attribute reference number for the selected attribute. After selecting the attribute to be changed, followed by entering the new attribute reference number, clicking on $\boldsymbol{O K}$ finalizes the operation.

L2


Fig. 4.64 An irregular area (top) meshed using free meshing (middle) and meshed with mapped meshing after line concatenation (bottom).

### 4.7.2.2 Clearing and Deleting Mesh

After a mesh is generated, there are ways to re-mesh if the user is not satisfied with the result. If direct generation was used for meshing, the user can simply "delete" elements first, and then nodes. Note that deleting elements does not automatically delete the nodes. These tasks are performed as follows:

```
Main Menu > Preprocessor > Modeling > Delete > Elements
Main Menu > Preprocessor > Modeling > Delete > Nodes
```

If the solid model approach was used, the elements and nodes cannot be "deleted" since they are "attached" to the solid model entities. However, more conveniently, the solid models that elements and nodes are attached to can be "cleared." This deletes all elements and nodes attached to the solid model entity at once. The user can now re-mesh the solid model entities after certain changes are made in meshing controls. The menu paths for this operation are as follows:

```
Main Menu > Preprocessor > Meshing > Clear > Keypoints
Main Menu > Preprocessor > Meshing > Clear > Lines
Main Menu > Preprocessor > Meshing > Clear > Areas
Main Menu > Preprocessor > Meshing > Clear > Volumes
```

There are cases where it may be advantageous to remove the association between the solid model and the mesh. This is achieved by using the following menu path:

Main Menu > Preprocessor > Checking Ctrls > Model Checking
which brings up a dialog box with a pull-down menu. Selecting the item Detach in the pull-down menu and clicking on $O K$ removes the association between the solid model and the mesh.

### 4.7.2.3 Numbering Controls

When dealing with complicated geometries, the Boolean operations explained in Sec. 4.4 are used regularly. These operations often generate new entities while removing existing ones, which creates gaps in the entity numbering. For example, when an area (say, area 1) is subtracted from another one (area 2), the resulting area is given the smallest available area number (in this case, area 3). Immediately after the creation of this area (area 3), ANSYS internally deletes the input areas (areas 1 and 2). Similarly, all the keypoints and lines associated with the new area are given new numbers while the ones associated with the old areas are removed. Similar considerations apply to nodes and elements. ANSYS provides the user with
the option of "compressing" the entity numbers, which is performed as follows:

Main Menu > Preprocessor > Numbering Ctrls > Compress Numbers
which leads to a dialog box with a pull-down menu. After the entity label is selected from this menu, clicking $\boldsymbol{O K}$ finalizes this operation.

Another important concept in solid modeling and meshing is the possible existence of duplicate entities. This usually occurs when the user creates new entities by copying existing ones or by reflection about a plane. If the old and new entities occupy the same space and if the material is supposed to be continuous along the line (or plane) where duplicate items lie, then they must be merged. Although it may not be apparent in the Graphics Window, the existence of duplicate entities compromises the continuity of the mesh, and thus may lead to invalid solutions. The following menu path is used to merge entities:

## Main Menu $>$ Preprocessor $>$ Numbering Ctrls $>$ Merge ltems

This brings up a dialog box in which the first field is a pull-down menu for the label of entities to be merged.

### 4.8 Selecting and Components

In the case of a three-dimensional finite element model, graphical picking may become a tedious and, at times, frustrating experience. In such cases, the selection tool provided by ANSYS is highly useful. This tool is efficient as it utilizes concepts from logics. The selected entity can then be saved in the ANSYS database as a "component." Thus, the next time this group of entities needs to be selected, selection of the component is sufficient. Selecting operations are discussed first, and the components next.

### 4.8.1 Selecting Operations

In ANSYS, entities are stored in separate sets (e.g., a set of areas, a set of volumes, etc.). Initially all of the full sets are active, until a selection operation is performed. When individual entities in a full set are selected (subset), they become active. Entity sets are independent of each other, i.e., selecting a group of lines does not cause any change in the selection status of the sets of keypoints or areas. Selections can be made based on several criteria, as explained below.

Selections can be performed by using the following menu path:

```
Utility Menu > Select > Entities
```

which brings up the Select Entities dialog box, as shown in Fig. 4.65. This dialog box has five distinct fields:

1. Entity Field: The entity to be selected is chosen using this pull-down menu.
2. Criterion Field: The entity chosen in the Entity Field is selected based on the criterion chosen in this pull-down menu. The following criteria are possible:

By Num/Pick: Clicking $O K$ after choosing this criterion starts the Pick Menu and the entities are selected by picking.

Attached to: As discussed previously, the entities in ANSYS are associated with each other. For example, a line is composed of at least two keypoints; an area is made up of at least three lines; etc. Thus, keypoints and lines, and lines and areas, are mutually attached (the list may be extended). When this criterion is chosen, another field appears in the Select Entities dialog box, listing the possibilities for attachment. If Areas in the Entity Field and Attached to in the Criterion Field are chosen, the new field lists Lines and Volumes as possible attachments; choosing Volumes and clicking on $O K$ results in the selection of volumes that are attached to the currently selected areas.

By Location: Selects entities based on their location. Upon choosing this criterion, a new field appears in the Select Entities dialog box with radio-buttons for $x$-, $y$-, and $z$-coordinates and a text field for the minimum and maximum values for the coordinate (shown in Fig. 4.66). For example, in order to select the nodes located between $y=2$ and $y=$ 5 , the radio-button for the $y$-coordinate is activated and the expression " 2,5 " (without the quotation marks) is entered in the text field.

By Attributes: This criterion is used for selecting entities based on their attributes (element type, material, real constant, etc.).

Exterior: Using this criterion, entities along the outer boundaries of the model are selected.

By Results: If a solution is obtained, then entities (only nodes and elements) can be selected based on result values.
3. Domain Field: This field determines the domain of the entity set with which the criterion is applied, as explained below:


Fig. 4.65 Select Entities dialog box (by number and picking).


Fig. 4.66 Select Entities dialog box (by location).

From Full: Selection is made from the full set of entities regardless of the selection status of the particular entity set.

Reselect: This option is used to refine the selection. It is used to select entities from a previously selected subset. For example, if the goal is to select all the nodes having coordinates $x=2$ and $y=3$ (both at the same time), then the nodes with the coordinate $x=2$ are first selected From Full set and, then, using Reselect button, the nodes with coordinate $y=$ 3 are selected from the previously selected subset of nodes with coordinate $x=2$.

Also Select: This option is used to expand the selection. It is used to add entities to the currently selected subset, based on a different criterion.

Unselect: This option is used to deactivate (unselect) a group of entities from the selected subset.

## 4. Domain Action Field:

Sele All: Selects the full set of a specific entity.
Invert: Inverts the selected set; active entities become inactive and vice versa.

Sele None: Unselects the full set of a specific entity; the active set becomes empty.

Sele Belo: Following the hierarchy of entities (i.e., volume is highest and node is lowest), this option selects the lower entities attached to the selected set of entities chosen in the Entity field.

## 5. Action Field:

OK: Applies the selection operation and closes the Select Entities dialog box.

Apply: Applies the selection operation; the Select Entities dialog box remains open for further selections.

Plot: Plots the currently selected set of a specific entity.
Replot: Updates the plot.
Cancel: Closes the Select Entities dialog box without applying the selection operation.

Help: Displays the help pages related to selection operations.
In order to select "everything" (reset all entities to their full sets), the following menu path is used:

Utility Menu > Select > Everything

### 4.8.2 Components

Groups of selected entities can be saved in an ANSYS database for easy retrieval. These groups are called components, and they can only contain entities of the same kind. The main advantage of defining components is to avoid multiple selection operations every time the user needs to select the same group of entities. The following menu path is used for defining components:

Utility Menu > Select > Comp/Assembly > Create Component
which is followed by a dialog box requesting the name to be given to the component and the type of entity to include in the component. Upon clicking $O K$, the component is created using the currently selected subset of the entity type chosen. The following menu path is used when a component has to be selected:

Utility Menu > Select > Comp/Assembly > Select Comp/Assembly
Listing and deletion of components is performed by using the following menu paths:

Utility Menu > Select > Comp/Assembly > List Comp/Assembly
Utility Menu > Select > Comp/Assembly > Delete Comp/Assembly

## Chapter 5

## ANSYS SOLUTION AND POSTPROCESSING

### 5.1 Overview

A typical ANSYS session, regardless of the discipline, involves the following steps:

1. Model Generation

- Specify jobname (this step is optional but recommended).
- Enter Preprocessor.
- Define element types and options.
- Define real constant for the element types (if the element type(s) require real constants).
- Define material properties.
- Create the model:.
- Build solid model (using either top-down or bottom-up approach).
- Define meshing controls.
- Create the mesh.
- Exit the Preprocessor.

2. Boundary/Initial Conditions and Solution

- Enter Solution Processor.
- Define analysis type and analysis options.
- Specify boundary/initial conditions:
- Degree of freedom constraints.
- Nodal force loads.
- Surface loads.
- Body loads.
- Inertia loads.
- Initial conditions (if the analysis type is transient).
- Save database (this step is not required but is recommended).
- Initiate solution.
- Exit the Solution Processor.

3. Review Results

- Enter the appropriate Postprocessor (General Postprocessor or Time History Postprocessor).
- Display results.
- List results.

The first step involves operations concerning the ANSYS Preprocessor and was covered in detail in Chap. 4. The operations pertaining to the solution and postprocessing of the results are discussed in detail in this chapter. At the end, specific steps are demonstrated by considering a one-dimensional transient heat transfer problem.

### 5.2 Solution

After preprocessing, the model generation, including meshing, is complete. The user is ready to begin the solution phase of the ANSYS session. First, the analysis type is specified from among the three main types:

- Static.
- Transient (time-dependent).
- Submodeling and substructuring (discussed in Sec. 11.3 and 11.4).

If the problem under consideration falls into the Structural Analysis discipline, then there are additional analysis types, such as modal, harmonic, spectrum, and eigenvalue buckling. There are two main deciding factors in choosing the analysis type:

Loading conditions: If the boundary conditions change as a function of time or there are initial conditions, then the analysis type is Transient. However, if the analysis discipline is structural and if the loading is a sinusoidal function of time, then the analysis type is Harmonic. Similarly, if the loading is a seismic spectrum, the analysis type is Spectrum.

Results of Interest: If the analysis discipline is structural and if the results of interest are the natural structural frequencies, then the analysis type is Modal. Similarly, if the interest is in determining the load at which the structure looses stability (buckles), then the analysis type is eigenvalue buckling.

The analysis type is specified by using the following menu path:
Main Menu $>$ Solution > Analysis Type $>$ New Analysis
This brings up the dialog box shown in Fig. 5.1. The user selects a particular analysis type by clicking on the corresponding radio-button and clicks $\boldsymbol{O K}$. The common solution operations used in almost every ANSYS session are discussed in the following subsections.

### 5.2.1 Analysis Options/Solution Controls

ANSYS allows the user to select certain options during the solution phase. They are specified through either Analysis Options or Solution Controls. The Analysis Options, specific to the Analysis Type, permit the user to select


Fig. 5.1 Dialog boxes for selecting the type of analysis for structural (top) and thermal (bottom) disciplines.
the method of solution and related details; this step requires familiarity with the Analysis Type. Analysis Options can be specified by the following the menu path:

Main Menu > Solution > Analysis Type > Analysis Options
Because the Analysis Options are specific to the particular problems under consideration, related discussions are covered in various example problems throughout this book.

In addition to the Analysis Options, the user has the option of specifying preferences through the Solution Controls. The main difference between the Analysis Options and the Solution Controls is that the Solution Controls are not specific to the Analysis Type. The same set of options in the Solution Controls can be used in a structural or thermal analysis. The Solution Controls dialog box can be activated by using the following menu path:

Main Menu > Solution > Analysis Type >Sol'n Controls
As shown in Fig. 5.2, the Solution Controls dialog box has five different tabs:


Fig. 5.2 Solution Controls dialog box.

Basic: Involves selection of options specific to the analysis type, time-domain-related parameters, and results items to be written to the Results File.

Transient: This option provides control over the way the loading is applied (stepped or ramped over time), the damping coefficients, and the time integration parameters.

Sol'n Options: The equation solver is chosen under this option. Also, if the current analysis is a Restart from a previous analysis or is intended to be "restarted" later, this option controls the number of restart files to write and the frequency at which they are written.

Nonlinear: Involves nonlinear options, specification of the maximum number of equilibrium iterations, and the limits on physical values used to perform bisection when performing nonlinear structural analyses, such as plasticity deformation, creep, etc.

Advanced NL: This option is used to specify what the software should do when convergence is not achieved during a nonlinear analysis.

Within the Solution Controls, all of the options have default values that the user is not required to specify while performing the analysis. However, if the analysis fails to produce convergence, manual specification of these options may improve the chances of convergence. As part of the example
problems solved throughout this book, the Solution Controls items are manually specified for several problems in Chap. 8 and 9.

### 5.2.2 Boundary Conditions

In a well-posed mathematical problem, the conditions along the entire boundary must be known. These conditions are referred to as the boundary conditions, and they can be specified in three different ways:

Type I: Specification of the primary variable (degree of freedom).
Type II: Specification of variables related to the derivative of the primary variable.

Type III: Specification of a linear combination of the primary variable and its derivative.

In a Structural problem, the primary variables are the displacement components (see Sec. 2.2.1.3). When Type I boundary conditions are used, the displacement constraints are specified along a segment of the boundary. If tractions are specified along the boundary, the boundary conditions fall under Type II because the tractions are related to the derivatives of the displacement components. A special case of the traction boundary conditions is the point load (also called the Force/Moment load). When the structure is subjected to tractions over a rather small area of the boundary, it is reasonable to idealize this condition as a concentrated load applied at a point. While conducting an analysis with BEAM or SHELL element types, moment loads can also be applied. Displacements, pressures (normal tractions), forces, and moments can be specified using the following menu paths:

```
Main Menu > Solution > Define Loads > Apply > Structural > Displacement
Main Menu > Solution > Define Loads > Apply > Structural > Force/Moment
Main Menu > Solution > Define Loads > Apply > Structural > Pressure
```

In a Thermal problem, the temperature is the primary variable. Similar to Structural problems, Type I boundary conditions correspond to the specification of the primary variable, i.e., temperature over a portion of the boundary. Specified heat flux conditions fall under Type II boundary conditions. Finally, convective conditions correspond to Type III boundary conditions. Temperature, heat flux, and convective conditions along the boundaries can be specified using the following menu paths:

[^3]Each of the boundary conditions discussed in this section can be applied on Nodes or on appropriate solid model entities such as Keypoints, Lines, or Areas. If they are applied on the solid model entities, ANSYS transfers them to the nodes when the solution is initiated. Although the paths for specification of boundary conditions are shown under the Solution Processor, it is possible to apply them under the Preprocessor.

### 5.2.3 Initial Conditions

A well-posed transient problem requires the specification of initial conditions. For Structural problems, initial conditions may involve components of displacement, rotation, velocity, or acceleration. In a Thermal problem, initial conditions are typically the temperature distribution within the domain. Initial conditions can be specified only when the Analysis Type is selected as Transient; if the Analysis Type is selected as Static, the specification of initial conditions does not appear as an option in the menus. Initial conditions can be specified using the following menu paths:

Main Menu > Solution > Define Loads > Apply > Initial Condit'n

### 5.2.4 Body Loads

Body loads can be generated internally or externally as the result of a physical field acting on the body. They act within the domain expressed volumetrically. Gravity, inertia loads, and temperature change represent body loads in a Structural problem. They can be specified using one of the following menu paths:

```
Main Menu > Solution > Define Loads > Apply > Structural > Temperature
Main Menu > Solution > Define Loads > Apply > Structural > Inertia > Angular Velocity
Main Menu > Solution > Define Loads > Apply > Structural > Inertia > Angular Accel
Main Menu > Solution > Define Loads > Apply > Structural > Inertia > Coriolis Effects
Main Menu > Solution > Define Loads > Apply > Structural > Inertia > Gravity
```

Heat generation within the domain is also represented as a body load for Thermal problems and can be specified using the following menu path:

Main Menu > Solution > Define Loads > Apply > Thermal > Heat Generat

### 5.2.5 Solution in Single and Multiple Load Steps

After completing the finite element mesh and specifying the loading conditions (boundary, initial, and body loads), the solution can be initiated using the following menu path:

However, there are cases in which the loads are time-dependent, and the solution is achieved in multiple steps. Different load steps must be used if the loading on the structure changes abruptly. The use of load steps also becomes necessary if the response of the structure at specific points in time is desired. ANSYS accommodates the application of time-dependent loads through the use of multiple Load Steps. Time-dependent loading is commonly encountered in analyses involving the determination of dynamic response and viscoplastic- and creep-type material behaviors. Simulation of manufacturing processes also involves time-dependent thermal loading. Figure 5.3 illustrates different profiles of impact loading as a function of time. The solid lines designate the actual loading while the dashed lines denote the loading profiles as specified in ANSYS. The solid circles indicate the times at which a load step starts or ends. As observed in Fig. 5.3, ANSYS permits the user to specify either step or ramped loading. In all of the cases, the last load step is necessary in order to capture the response of the structure at times after the load is removed. The multiple load steps are also necessary in modeling a viscoplastic material subjected to thermal cycling, as shown in Fig. 5.4.

The following steps are used in order to use multiple load step solution method:

1. Apply the initial conditions as explained in Sec. 5.2.3.
2. Apply the boundary conditions appropriate for the first load step.
3. Specify time-related parameters: This is performed by using the following menu path:

Main Menu $>$ Solution $>$ Load Step Opts $>$ Time/Frequenc $>$ Time - Time Step
This brings up the Time and Time Step Options dialog box (Fig. 5.5) Enter the time at end of load step (TIME) and the time step size (DELTIM), which is optional. Choose between whether the loads are applied in a stepped or ramped manner (KBC). If the Automatic Time Stepping is $O F F$, then the user must specify the time step size. If the time step size is not specified and the Automatic Time Stepping is set as Prog Chosen (stands for program chosen), then ANSYS turns the Automatic Time Stepping $O N$. If the time step size is specified and the Automatic Time Stepping is ON, ANSYS starts the solution with the specified time step size and modifies it based on the convergence.
4. Write Load Step file: Load steps are written to Load Step Files by using the following menu path:


Fig. 5.3 Different profiles of impact loading as functions of time.


Fig. 5.4 Cyclic thermal loading.


Fig. 5.5 Time and Time Step Options dialog box.

This brings up the Write Load Step File dialog box (Fig. 5.6). Enter the load step file number (LSNUM) and hit $\boldsymbol{O K}$. This file is stored in the Working Directory and contains all the solution options, the time, and time-related parameters, as well as the boundary conditions.
5. Repeat steps $\mathbf{2 - 4}$ for the remainder of the load steps.
6. Initiate solution from Load Step files: Once all of the load step files are written, the solution is initiated by using the following menu path:

Main Menu > Solution > Solve > From LS Files
which brings up the Solve Load Step Files dialog box. Enter the starting and ending load step file numbers (LSMIN and LSMAX) and hit $\boldsymbol{O K}$.


Fig. 5.6 Write Load Step File dialog box.

### 5.2.6 Failure to Obtain Solution

There are two common reasons why ANSYS fails to provide a solution:
Singular coefficient matrix: As shown in detail in Chap. 1, every finite element solution involves the solution of a system of equations with a known coefficient matrix (stiffness), an unknown degree of freedom vector, and a known right-hand-side (force) vector. If the coefficient matrix is singular, the solution fails. The most common reasons why the coefficient matrix becomes singular are as follows:

1. Instability in the structure due to lack of constraints in static structural analyses. This leads to rigid-body translations and rotations, which makes the stiffness matrix singular. As an example of this phenomenon, Fig. 5.7 shows three distinct constraint configurations applied on the same 2-D square structure subjected to a distributed tensile load in the $x$-direction. The first configuration involves two constraints, both suppressing displacements in the $y$-direction, along the bottom surface of the structure. Because there are no constraints suppressing displacements in the $x$-direction, the structure is free to move in the $x$ direction under the applied load, thus leading to a singular stiffness matrix. In the second configuration, displacements in both the $x$ - and $y$-directions are suppressed at the same point (bottom left corner). Although this configuration prevents rigid-body translations, it fails to prevent rigid-body rotation around the corner node where displacement constraints are applied, causing the stiffness matrix to become singular. Finally, in the third configuration, two corners are constrained in the $y$-direction, with one of them also constrained in the $x$ direction. This is a stable configuration, preventing all possible rigidbody movements, leading to a nonsingular stiffness matrix and thus a successful unique solution.

insufficient
(rigid body translation in $x$-direction)

insufficient (rigid body rotation)

sufficient
(no rigid body movement)

Fig. 5.7 Instability due to lack of constraints (left and middle); stable configuration.

- Material properties that are physically impossible may make the coefficient matrix singular. Examples include zero or negative Young's modulus, thermal conductivity, density, or specific heat.
- There are structural elements within the ANSYS element library that carry loads only along their line of direction (SPAR elements simulating truss structures). Stability concerns of Statics apply to structures made up of these elements, and the user must make sure that the structure is stable.

2. Failed convergence: In finite element analyses, problems involving nonlinearity are solved through iterations. As described in Chap. 2, these nonlinearities arise through the material behavior (plasticity, creep, viscoelasticity, viscoplasticity, etc.) or geometric configuration (large deformations) of the structure. The "correct" solution is approached in small steps, referred to as convergence iterations. If the problem is time-dependent, then the small steps are taken in the time domain. If the problem is not dependent on time (e.g., plasticity), these small steps are taken in the application of the loads. At the end of each iteration, ANSYS checks whether the solution satisfies a convergence criterion "built-in" for different analysis types. If the criterion is not satisfied, the last step is repeated with a smaller step size. This is repeated until the convergence criterion is satisfied. However, there are limits on the number of convergence iterations and, if a converged solution is not achieved within those limits, ANSYS terminates the solution process. Because each nonlinear analysis type is different, there is no straightforward answer as to what to do to improve the chances of a successful convergence. However, several nonlinear problems are considered in Chap. 10 that may give the reader some ideas on convergence considerations.

### 5.3 Postprocessing

After a solution is obtained in an ANSYS session, the user can review the results in either the General Postprocessor or the Time History Postprocessor. If the problem is static (or steady state), then the General Postprocessor is the only postprocessor where the results can be reviewed. However, if the problem is dependent on time (transient), both processors are useful for distinctly different tasks. The postprocessors and common postprocessing operations are discussed briefly in the following (5.3.15.3.6).

### 5.3.1 General Postprocessor

In the General Postprocessor, the results of a solution at a specific time (if the problem is time dependent) are reviewed. Available options for review include graphical displays and a listing of results. It is also possible to perform sorting and mathematical operations on the results.

### 5.3.2 Time History Postprocessor

When the problem under consideration is time dependent, the time variation of the results at specific locations (nodes) are reviewed under the Time History Postprocessor. Upon entering this postprocessor, the Time History Variables dialog box appears (Fig. 5.8). This dialog box has three distinct areas: Toolbar, Variables, and Calculator. The first four buttons (from the left) in the Toolbar are the most commonly used ones:

Add Data Button: This button is used to define new variables, such as displacements, temperatures, etc., at specific nodes.

Delete Data Button: Used for deleting defined variables.
Graph Data: Using this button, the user can plot the time variation of variables.

List Data: Similar to plotting, this button is used for listing the results as functions of time.

As the new variables are defined, they appear in the Variables area. By default, TIME is the first variable and cannot be removed. In addition to the name of the variable, the Variables area includes useful information about the variable, such as its element or node number, what result item it corresponds to, and the range of its values.


Fig. 5.8 Time History Variables dialog box.

The last item (located at the right-most side) is the $X$-Axis button, which enables the user to select which variable to display on the $x$-axis in the graphical representations.

An example problem (time-dependent heat transfer) demonstrating the use of the Time History Postprocessor is given in Sec. 5.4.

### 5.3.3 Read Results

The results, obtained through the Solution Processor, are saved in results files (jobname.rst for structural, jobname.rth for thermal, and jobname.rfl for fluids problems), which are stored in the working directory. In order to review the results, the user needs to guide ANSYS so that the correct results file is selected. This is done by using the following menu path:

## Main Menu > General Postproc > Data \& File Opts

This brings up the Data and File Options dialog box (Fig. 5.9). The results file is selected by clicking on the browse button (button with three dots). After selecting the correct file, click on $\boldsymbol{O K}$.

If the solution does not involve multiple substeps and load steps, then there is only one results "set" the user can review. However, when the solution involves multiple substeps and load steps, there are many results sets and the user should select the correct (intended) one. The results sets can be selected using the following options:


Fig. 5.9 Data and File Options dialog box.

First Set: Results related to the first available set are read into the database using the following menu path:

Main Menu $>$ General Postproc $>$ Read Results $>$ First Set
Next Set: Results related to the set available immediately after the current set are read into the database using the following menu path:

Main Menu > General Postproc > Read Results > Next Set
Previous Set: Results related to the set available immediately before the current set are read into the database using the following menu path:

Main Menu > General Postproc > Read Results > Previous Set
Last Set: Results related to the last available set are read into the database using the following menu path:

Main Menu > General Postproc $>$ Read Results $>$ Last Set
Read By Picking: The following menu path is used for this option:
Main Menu > General Postproc > Read Results > By Pick
which brings up a dialog box (Fig. 5.10) listing the available results sets. The user selects the desired results set and clicks on Read and Close for the results to be read into the database.


Fig. 5.10 Results File dialog box.
Read By Load Step Number: Results related to a specific load step and substep are read into the database using the following menu path:

Main Menu > General Postproc > Read Results > By Load Step
which brings up the Read Results by Load Step Number dialog box (Fig. 5.11) in which the user specifies the load step number (LSTEP) and substep (SBSTEP) number within that load step and clicks on $\boldsymbol{O K}$ for the results to be read into the database.

Read By Time: Results related to a specific time (or frequency) value are read into the database using the following menu path:

Main Menu > General Postproc > Read Results > By Time/Freq
which brings up the Read Results by Time or Frequency dialog box (Fig. 5.12) in which the user specifies the value of time (or frequency) (TIME) and clicks on $\boldsymbol{O K}$ for the results to be read into the database.

### 5.3.4 Plot Results

After the desired results set is read into the database, the result quantities can be reviewed through graphics displays. The types of graphics displays include deformed shapes (structural analysis), contour plots, vector displays (thermal), and path plots.

In structural analyses, the deformed shape resulting from the applied loads and boundary conditions is displayed using the following menu path:
Main Menu > General Postproc > Plot Results > Deformed Shape
which brings up the Plot Deformed Shape dialog box (Fig. 5.13). The user is offered three distinct display modes:


Fig. 5.11 Read Results by Load Step Number dialog box.


Fig. 5.12 Read Results by Time or Frequency dialog box.


Fig. 5.13 Plot Deformed Shape dialog box.

- Display deformed shape only.
- Display deformed and undeformed shapes together.
- Display deformed shape with the outer boundary (edge) of the undeformed shape.

After the user makes a choice and clicks on $\boldsymbol{O K}$, the deformed shape appears in the Graphics Window.

Contour plots are obtained using one of the following menu paths:

```
Main Menu > General Postproc > Plot Results > Contour Plot > Nodal Solu
Main Menu > General Postproc > Plot Results > Contour Plot > Element Solu
```

which brings up the Contour Nodal (Element) Solution Data dialog box (Fig. 5.14). In this dialog box, the first two fields identify the quantity to be plotted. They include the degree of freedom (DOF) solution (displacements, temperatures, etc.), and derived quantities (stresses, strains, fluxes, etc.). Once the user makes the selection, upon clicking $\boldsymbol{O K}$, the contour plot appears in the Graphics Window.

Vector plots are obtained using the following menu path:
Main Menu > General Postproc > Plot Results > Vector Plot > Predefined
which brings up the Vector Plot of Predefined Vectors dialog box. Similar to the contour plots, this dialog box has two fields identifying the quantity to be plotted. Once the user makes a selection, upon clicking $\boldsymbol{O K}$, the vector plot appears in the Graphics Window.

In the ANSYS General Postprocessor, it is possible to obtain line plots along a path. Utilizing path plots involves:


Fig. 5.14 Contour Nodal Solution Data dialog box.
Defining Paths: This can be performed by different methods, one of which uses the following menu path:

Main Menu > General Postproc > Path Operations > Define Path > By Nodes
which brings up a Pick Menu for the nodes defining the path to be picked. Upon clicking on $O K$ in the Pick Menu, a dialog box appears asking for path specifications such as the user-defined name of the path and number of divisions between data points. Clicking $O K$ in this dialog box finishes the path definition.

Mapping Quantities onto Paths: Once the paths are defined, quantities of interest are mapped onto paths by using the following menu path:

## Main Menu > General Postproc $>$ Path Operations $>$ Map onto Path

which brings up the Map Result Items onto Path dialog box. The user specifies a unique label for the result item to be mapped and selects the result item; clicking $\boldsymbol{O K}$ completes the mapping operation.

Plotting Quantities on Graphs or on Geometry: The quantities mapped onto defined paths can be plotted using the following menu items:

[^4]which brings up the Plot of Path Items on Graph (Geometry) dialog box. The user selects the path item to be plotted from the list of defined path items and clicks on $\mathbf{O K}$; the plot appears in the Graphics Window.

The operations related to path plots are demonstrated through an example problem in Sec. 5.4.

### 5.3.5 Element Tables

In ANSYS, each element type possesses numerous output quantities available upon completion of the solution. Although several of these quantities are offered by their names under the postprocessors, some are not directly accessible, and the user needs to take additional steps in order to access them. One important purpose of using Element Tables is to access these result items. Another important role of Element Tables is that they enable the user to perform arithmetic operations involving several result items. Element tables are defined by using the following menu path:

## Main Menu > General Postproc > Element Table > Define Table

which brings up the Element Table Data dialog box (Fig. 5.15). In order to add new items to the element table, the user needs to click on the Add button, which brings up the Define Additional Element Table Items dialog box (Fig. 5.16). After selecting the result quantity and specifying a user label for it, clicking on $O K$ completes the element table item definition.

In order to explain the usage of element tables, an example based on the plane5s element type is considered. In Table 5.1, the output quantities provided by the element type plane55 are given. Table 5.2 lists the results quantities accessible through the element tables. For example, the heat flow rate per unit area across the element faces caused by input heat flux, denoted by HFLXAVG in Tables 5.1 and 5.2, is available for definition in the element tables. In Table 5.2, the matching item for this quantity is given as NMISC, along with numbers $6,12,18$, and 24 corresponding to different faces of the element. To store $H F L X A V G$ at the $4^{\text {th }}$ face of each element, in the Define Additional Element Table Items dialog box, By sequence num from the left list and NMISC from the right list must be selected. Entering 24 in the text field underneath the right list ensures that the HFLXAVG at the $4^{\text {th }}$ face of each element will be stored in the element table.

Element tables are plotted and listed using the following menu paths:

[^5]

Fig. 5.15 Element Table Data dialog box.


Fig. 5.16 Define Additional Element Table Items dialog box.

Table 5.1 Output quantities provided by the PLANE55 element type.

| Name | Definition |
| :---: | :--- |
| EL | Element Number |
| NODES | Nodes: I, J, K, L |
| MAT | Material number |
| VOLU | Volume |
| XC, YC | Location where results are reported |
| HGEN | Heat generations HG(I), HG(J), HG(K), HG(L) |
| TG:X, Y, SUM | Thermal gradient components and vector sum at |
| centroid |  |
| TF:X, Y, SUM | Thermal flux (heat flow rate/cross-sectional area) |
| components and vector sum at centroid |  |
| FACE | Face label |
| AREA | Face area |
| NODES | Face nodes |
| HFILM | Film coefficient at each node of face |
| TBULK | Bulk temperature at each node of face |
| TAVG | Average face temperature |
| HEAT RATE | Heat flow rate across face by convection |
| HFAVG | Average film coefficient of the face |
| TBAVG | Average face bulk temperature |
| HFLXAVG | Heat flow rate per unit area across face caused by |
| input heat flux |  |
| HEAT RATE/AREA | Heat flow rate per unit area across face by |
| convection |  |
| HFLUX | Heat flux at each node of face |

Table 5.2 Quantities obtained via the element table.

| Output Quantity <br> Name | Element Table Input |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | Item | FC1 | FC2 | FC3 | FC4 |
| AREA | NMISC | 1 | 7 | 13 | 19 |
| HFAVG | NMISC | 2 | 8 | 14 | 20 |
| TAVG | NMISC | 3 | 9 | 15 | 21 |
| TBAVG | NMISC | 4 | 10 | 16 | 22 |
| HEAT RATE | NMISC | 5 | 11 | 17 | 23 |
| HFLXAVG | NMISC | 6 | 12 | 18 | 24 |

It is also possible to perform arithmetic operations within each column or between the columns of the element table. Examples of such operations include: finding absolute values, finding the sum of each element table item, adding and multiplying element table items, etc.

### 5.3.6 List Results

Results of an ANSYS solution can be reviewed through lists. Although there are numerous different options for listing the results under postprocessors, only two of them are discussed in this section: nodal and element solutions. In order to list results computed at the nodes, the following menu path is used:

## Main Menu > General Postproc > List Results $>$ Nodal Solu

which brings up the List Nodal Solution dialog box. Once the user makes a selection as to what result quantities are to be reviewed and clicks on $O K$, the list appears in a separate window. Similar to nodal solution listings, the element results are listed by using the following menu path:
Main Menu > General Postproc $>$ List Results $>$ Element Solu
The usage of this option is similar to the nodal solution lists.

### 5.4 Example: One-dimensional Transient Heat Transfer

Consider the one-dimensional transient heat transfer problem shown in Fig. 5.17. The problem is time dependent; therefore, in addition to thermal conductivity, the specific heat and the density of the material are taken into account. The governing equation for this problem is written as

$$
\begin{equation*}
\rho c \frac{\partial T}{\partial t}=\kappa \frac{\partial^{2} T}{\partial x^{2}} \quad 0 \leq x \leq l \tag{5.1}
\end{equation*}
$$

with the boundary and initial conditions

$$
\begin{align*}
& T(x=0, t)=T_{a}=100 \\
& T(x=l, t)=T_{b}=0  \tag{5.2}\\
& T(x, t=0)=f(x)=0
\end{align*}
$$

$$
\kappa, \rho, c
$$

Fig. 5.17 One-dimensional transient heat transfer problem.

The analytical solution for this problem is given by (Carslaw and Jaeger, 1959, pp. 99-100):

$$
\begin{align*}
T(x, t)=T_{a}+\left(T_{b}-T_{a}\right) \frac{x}{l} & +\frac{2}{\pi} \sum_{n=1}^{\infty} \frac{T_{b} \cos n \pi-T_{a}}{n} \sin \frac{n \pi x}{l} e^{-\alpha n^{2} \pi^{2} t / l^{2}} \\
& +\frac{2}{l} \sum_{n=1}^{\infty} \sin \frac{n \pi x}{l} e^{-\alpha n^{2} \pi^{2} t / l^{2}} \int_{0}^{l} f\left(x^{\prime}\right) \sin \frac{n \pi x^{\prime}}{l} d x^{\prime} \tag{5.3}
\end{align*}
$$

where $\alpha=\kappa /(\rho c)$. Substituting $T_{a}=100, T_{b}=0, f(x)=0, l=2, \kappa=1$, $\rho=10$, and $c=3$, Eq. (5.3) yields

$$
\begin{equation*}
T(x, t)=100-100 \frac{x}{2}+\frac{2}{\pi} \sum_{n=1}^{\infty}\left(-\frac{100}{n}\right) \sin \frac{n \pi x}{2} e^{-n^{2} \pi^{2} t / 120} \tag{5.4}
\end{equation*}
$$

When computing the exact solution using this equation, the number of terms, $n$, is truncated at 40 for satisfactory convergence.

Subjected to the boundary conditions indicated in Fig. 5.18 in the time range $0 \leq t \leq 5$, this problem is solved by using two-dimensional PLANE55 elements in ANSYS.

The model is generated by using 4 element divisions along the vertical boundaries and 20 element divisions along the horizontal boundaries. The temperature variations along the midline at times $t=0.1,0.5$, and 5 are obtained by ANSYS and the exact solution is plotted.

## model generation

- Specify the jobname as Id_dif using the following menu path:

Utility Menu > File > Change Jobname

- In the dialog box, type 1d_dif in the [/FILNAM] Enter new jobname text field; click on the check box for New log and error files to show Yes; click on $O K$.
- Define the element type (ET command) using the following menu path:


## Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Thermal Solid on the left list and Quad 4node 55 on the right list; click on $\mathbf{O K}$.
- Click on Close.


Fig. 5.18 Two-dimensional representation of the 1-D transient heat transfer problem and corresponding boundary conditions.

- Specify material properties (MP command) using the following menu path:


## Main Menu $>$ Preprocessor $>$ Material Props $>$ Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Thermal, Conductivity, and, finally, Isotropic, which brings up another dialog box.
- Enter $\boldsymbol{I}$ for $\boldsymbol{K X X}$; click on $\boldsymbol{O K}$.
- In the Define Material Model Behavior dialog box, in the right window, double-click on Specific Heat, which brings up another dialog box.
- Enter 3 for $\boldsymbol{C}$; click on $\boldsymbol{O K}$.
- In the Define Material Model Behavior dialog box, in the right window, double-click on Density, which brings up another dialog box.
- Enter 10 for DENS; click on $O \boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:


## Material > Exit

- Create the solid model:
- Create a rectangular area using the following menu path: Main Menu > Preprocessor > Modeling > Create $>$ Areas $>$ Rectangle $>$ By Dimensions
- In the Create Rectangle by Dimensions dialog box, type 0 for $\mathbf{X 1}, 2$ for $X 2,0$ for $Y 1$, and 0.5 for $Y 2$; click on $O K$.

Create the mesh:

- Specify the number of elements along the vertical boundaries using the following menu path:

Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize $>$ Lines $>$ Picked Lines

- Pick the two vertical lines; click on $\boldsymbol{O K}$.
- Element Sizes on Lines dialog box appears; type 4 in the text field corresponding to NDIV (the second text field), and uncheck the first check box; click on $\boldsymbol{O K}$.
- Specify the number of elements along the horizontal boundaries using the following menu path:

```
Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Lines > Picked Lines
```

- Pick the two horizontal lines; click on $\boldsymbol{O K}$.
- Element Sizes on Lines dialog box reappears; type 20 in the text field corresponding to NDIV (the second text field), and uncheck the first check box; click on $\boldsymbol{O K}$.
- Create the mesh using the following menu path:

Main Menu > Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 sided

- In the Pick Menu, click on Pick All.
- Figure 5.19 shows the mesh.
- Save the model using the following menu path:

Utility Menu > File > Save as Jobname.db
The model is saved under the name $1 d \_$dif. $d b$ in the working directory.

## SOLUTION

- Specify the analysis type as transient using the following menu path:

Main Menu > Solution > Analysis Type > New Analysis

- Click on Transient; click on $\boldsymbol{O K}$.
- A new dialog box appears; click on $\boldsymbol{O K}$.


Fig. 5.19 Mesh used in the analysis.

- Specify temperature boundary conditions along the vertical boundaries using the following menu path:


## Main Menu > Solution $>$ Define Loads $>$ Apply $>$ Thermal $>$ Temperature $>$ On Nodes

- Pick Menu appears; click on the Box radio-button and draw a rectangle around the nodes along the left vertical boundary; click on $\boldsymbol{O K}$.
- Apply TEMP on Nodes dialog box appears; highlight TEMP, enter 100 for VALUE Load TEMP value; click on Apply.
- Pick Menu reappears; click on the Box radio-button and draw a rectangle around the nodes along the right vertical boundary; click on $\boldsymbol{O K}$.
- Apply TEMP on Nodes dialog box reappears; highlight TEMP, enter 0 for VALUE Load TEMP value; click on OK.
- Specify initial conditions within the domain using the following menu path:

Main Menu > Solution > Define Loads > Apply > Initial Condit'n > Define

- Pick Menu appears; click on Pick All.
- Define Initial Conditions dialog box appears; select TEMP on the Lab pull-down menu; enter $\boldsymbol{0}$ in the VALUE text field; click on $\boldsymbol{O K}$.
- Specify time parameters using the following menu path:

Main Menu > Solution > Load Step Opts > Time/Frequenc > Time - Time Step

- Time and Time Step Options dialog box appears.
- As shown in Fig. 5.20, enter 5 in the [TIME] Time at end of load step text field and $5 / 100$ in the [DELTIM] Time step size text field, and click on the Stepped radio-button for [KBC]; click on OK.
- Specify output controls using the following menu path:

Main Menu > Solution > Load Step Opts > Output Ctrls > DB/Results File

- Controls for Database and Results File Writing dialog box appears.
- As shown in Fig. 5.21, click on the Every substep radio-button for FREQ File write frequency; click on $O K$.
- Obtain solution using the following menu path:

Main Menu >Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window and click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


Fig. 5.20 Time and Time Step Options dialog box used in the analysis.


Fig. 5.21 Output Controls dialog box used in the analysis.

## GENERAL POSTPROCESSING

- Review results at the end of first substep using the following menu paths:

Main Menu $>$ General Postproc $>$ Read Results $>$ First Set
Main Menu > General Postproc > Plot Results > Contour Plot > Nodal Solu

- Contour Nodal Solution Data dialog box appears; select DOF solution on the left list and Temperature TEMP on the right list; click on $O K$.
- Figure 5.22 shows the contour plot of the temperature distribution at $t=0.05$ as it appears in the Graphics Window.
- Review results at the "next" substep using the following menu paths:

Main Menu > General Postproc $>$ Read Results $>$ Next Set
Main Menu > General Postproc > Plot Results > Contour Plot $>$ Nodal Solu

- Contour Nodal Solution Data dialog box appears; select DOF solution on the left list and Temperature TEMP on the right list; click on $\boldsymbol{O K}$.
- Figure 5.23 shows the contour plot of temperature distribution at $t=0.1$ as it appears in the Graphics Window.
- Review results at $t=0.5$ using the following menu path:

Main Menu > General Postproc > Read Results > By Time/Freq

- As shown in Fig. 5.24, Read Results by Time or Frequency dialog box appears; enter 0.5 for TIME Value of time or freq; click on OK.
- View the temperature contours, as shown in Fig. 5.25.
- Review results at the "last" substep using the following menu paths:

Main Menu > General Postproc > Read Results > Last Set
Main Menu $>$ General Postproc $>$ Plot Results $>$ Contour Plot $>$ Nodal Solu

- Contour Nodal Solution Data dialog box appears; select DOF solution on the left list and Temperature TEMP on the right list; click on $O K$.
- Figure 5.26 shows the contour plot of the temperature distribution at $t=5$ as it appears in the Graphics Window.
- Review thermal flux vector plot at $t=5$ using the following menu path:

Main Menu > General Postproc > Plot Results $>$ Vector Plot $>$ Predefined

- Vector Plot of Predefined Vectors dialog box appears; select Thermal flux TF on the right list and click on $O K$.
- Figure 5.27 shows the vector plot of the thermal flux as it appears in the Graphics Window.


Fig. 5.22 Temperature distribution contour plot at $t=0.05$.


Fig. 5.23 Temperature distribution contour plot at $t=0.1$.


Fig. 5.24 Read Results by Time or Frequency dialog box used in the analysis.


Fig. 5.25 Temperature distribution contour plot at $t=0.5$.


Fig. 5.26 Temperature distribution contour plot at $t=5$.


Fig. 5.27 Vector plot of thermal flux at $t=5$.

- Review results by path plots:
- Plot elements using the following menu path:

Utility Menu > Plot > Elements

- Turn node numbering on using the following menu path:

Utility Menu > PlotCtrls>Numbering

- Plot Numbering Controls dialog box appears; click on the check box for NODE Node numbers to show On; click on OK.
- Define the path using the following menu path:

Main Menu > General Postproc > Path Operations > Define Path > By Nodes

- Pick Menu appears; pick nodes $47(x=0, y=0.25)$ and $24(x=2, y=$ $0.25)$; click on $O K$.
- Define Path By Nodes dialog box appears, as shown in Fig. 5.28; enter a unique name (e.g., y025) identifying the path; click on $O K$.
- Close the Path Status Information Window.
- Turn node numbering off using the following menu path: Utility Menu >PlotCtrls > Numbering
- In the Plot Numbering Controls dialog box, click on the check box for NODE Node numbers to show Off; click on OK.
- Plot the path on geometry using the following menu path:

Main Menu > General Postproc > Path Operations > Plot Paths

- Figure 5.29 shows the result of this action.
- Map the temperature results onto the defined path using the following menu path:


## Main Menu > General Postproc > Path Operations > Map onto Path

- Map Result Items onto Path dialog box appears, as shown in Fig. 5.30; enter a unique name for the result item (e.g., t025, note that this is different than the name given for the path); select DOF solution from the left list and Temperature TEMP from the right list for the item; click on $O K$.
- Path plot the temperature results using the following menu path:

Main Menu > General Postproc $>$ Path Operations $>$ Plot Path Item $>$ On Graph

- Plot of Path Items on Graph dialog box appears; select T025 from the list; click on $O K$.
- Observe the temperature variation along the path $\mathbf{y 0 2 5}$ as it appears in the Graphics Window, as shown in Fig. 5.31.
- Map the flux results onto the defined path using the following menu path:

Main Menu $>$ General Postproc $>$ Path Operations $>$ Map onto Path

- Map Result Items onto Path dialog box appears, as shown in Fig. 5.32; enter a unique name for the result item (e.g., q025); select Flux \& gradient from the left list and Thermal flux TFX from the right list for the item; click on $\boldsymbol{O K}$.
- Path plot the flux results using the following menu path:

[^6]
## \By Nodes

[PATH] Define Path specifications

| Name Define Path Name : | y025 <br> nSets Number of data sets |
| :--- | :--- |
| nDiv Number of divisions | 30 |



Fig. 5.28 Defining path specifications.


Fig. 5.29 Geometry plot of path Y025.


Fig. 5.30 Map Result Items onto Path dialog box used for line plot of temperature along the path Y025.


Fig. 5.31 Temperature variation line plot along the path Y025 at $t=5$.


Fig. 5.32 Map Result Items onto Path dialog box used for line plot of flux along the path Y025.

- Plot of Path Items on Graph dialog box appears; unselect T025 and select Q025 from the list; click on $O K$.
- Observe the flux variation along the path $\mathbf{y} 025$ as it appears in the Graphics Window, as shown in Fig. 5.33.
- Finally, plot the flux on actual geometry using the following menu path:
Main Menu > General Postproc > Path Operations > Plot Path Item > On Geometry
- Plot of Path Items on Geometry dialog box appears; select Q025 from the list; click on $\boldsymbol{O K}$.
- Figure 5.34 shows the path plot of thermal flux as it appears in the Graphics Window.


## TIME HISTORY POSTPROCESSING

- Review time-dependent behavior of temperature at nodes located at $(0.1,0.25)$ and $(1.0,0.25)$ using the following menu path:


## Main Menu > TimeHist Postpro

- Time History Variables dialog box appears (Fig. 5.35).
- Click on the button with the green plus sign at the top-left to define a variable.
- Add Time History Variable dialog box appears.
- Successively double-click on the items Nodal Solution, DOF Solution, and Temperature; click on $\boldsymbol{O K}$.
- Pick Menu appears; pick the node located at $x=0.1, y=0.25$ (as indicated in Fig. 5.36); click on $O K$.
- Note the new variable TEMP_2 in the Time History Variables dialog box.
- Add a new variable for temperature at the center node by clicking on the button with the green plus sign and successively double-clicking on the items Nodal Solution, DOF Solution, and Temperature; click on OK.
- Pick the node located at $x=1, y=0.25$ (as indicated in Fig. 5.36); click on $\boldsymbol{O K}$.
- Note the new variable TEMP_3 in the Time History Variables dialog box.
- Highlight the rows TEMP_2 and TEMP_3 from the list (by pressing Ctrl on the keyboard and clicking on the rows with the left mouse button); click on the third from the left button to plot the time variation of these temperatures.
- The plot appears in the Graphics Window, as shown in Fig. 5.37.


Fig. 5.33 Flux variation line plot along the path Y025 at $t=5$.


Fig. 5.34 Path plot on geometry for the variation of flux.


Fig. 5.35 Time History Variables dialog box.


Fig. 5.36 Nodes to be picked to review the time-dependent behavior of temperature and flux.


Fig. 5.37 Temperature variation over time at two nodes.

- Review time-dependent behavior of thermal flux at nodes located at $(0.1,0.25)$ and $(1.0,0.25)$.
- In the Time History Variables dialog box, highlight the rows TEMP_2 and $\boldsymbol{T E M P} \_3$ from the list and click on the second from the left button (button with a red cross) to delete the temperature variables.
- In order to add thermal flux variables, click on the button with the green plus sign at the top-left to define a variable.
- Add Time History Variable dialog box appears.
- Successively double-click on the items Nodal Solution, Thermal Flux, and $\boldsymbol{X}$-Component of thermal flux; click on $\mathbf{O K}$.
- The Pick Menu appears; pick the node located at ( $x=0.1, y=0.25$ ) (as indicated in Fig. 5.36); and click on $\boldsymbol{O K}$.
- Note the new variable TFX_2 in the Time History Variables dialog box.
- Add a new variable for thermal flux at the center node by clicking on the button with the green plus sign and successively double-clicking on the items Nodal Solution, Thermal Flux, and X-Component of thermal flux; click on $O K$.
- Pick the node located at $(x=1, y=0.25)$ (as indicated in Fig. 5.36); click on $\boldsymbol{O K}$.
- Note the new variable TFX_3 in the Time History Variables dialog box.
- Highlight the rows $\boldsymbol{T F X} \_2$ and $\boldsymbol{T F X} \_3$ from the list (by pressing Ctrl on the keyboard and clicking on the rows with the left mouse button); click on the third from the left button to plot the time variation of these thermal fluxes.
- The plot appears in the Graphics Window, as shown in Fig. 5.38.
- Close Time History Variables dialog box.

Table 5.3 lists the temperature values along the midline ( $y=0.25$ ) obtained by ANSYS (columns 2-4) and the analytical solution given by Eq. (5.4) (columns 5-7) at times $t=0.1,0.5$, and 5 . The analytical solution is obtained by using $n=40$ in the series. Three separate ANSYS solutions are obtained with the final times $t=0.1,0.5$, and 5 , each utilizing 100 equal time steps. Figure 5.39 shows a graphical comparison of the analytical and ANSYS solutions.


Fig. 5.38 Flux variation over time at two nodes.


Fig. 5.39 Graphical comparison of ANSYS and analytical solutions.

Table 5.3 Temperature values along the midline $(y=0.25)$ obtained by ANSYS and Eq. (5.4) $(t=0.1,0.5,5.0)$.

| $x$ | ANSYS |  |  | EXACT $(n=40)$ |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | $t=0.1$ | $t=0.5$ | $t=5.0$ | $t=0.1$ | $t=0.5$ | $t=5.0$ |
| 0.00 | 100.0000 | 100.0000 | 100.0000 | 100.0000 | 100.0000 | 100.0000 |
| 0.10 | 24.2230 | 56.7890 | 88.1810 | 22.0671 | 58.3882 | 86.2490 |
| 0.20 | 3.7816 | 26.6910 | 72.7800 | 1.4306 | 27.3322 | 72.9034 |
| 0.30 | 0.4497 | 10.7250 | 60.1760 | 0.0239 | 10.0348 | 60.3332 |
| 0.40 | 0.0443 | 3.7954 | 48.6770 | $9.84 \mathrm{E}-05$ | 2.8460 | 48.8422 |
| 0.50 | 0.0038 | 1.2112 | 38.5010 | $2.09 \mathrm{E}-06$ | 0.6170 | 38.6475 |
| 0.60 | 0.0003 | 0.3550 | 29.7610 | $1.88 \mathrm{E}-06$ | 0.1015 | 29.8698 |
| 0.70 | $2.07 \mathrm{E}-05$ | 0.0969 | 22.4760 | $1.73 \mathrm{E}-06$ | 0.0126 | 22.5346 |
| 0.80 | $1.36 \mathrm{E}-06$ | 0.0249 | 16.5800 | $1.56 \mathrm{E}-06$ | 0.0012 | 16.5857 |
| 0.90 | $8.47 \mathrm{E}-08$ | 0.0061 | 11.9450 | $1.40 \mathrm{E}-05$ | $8.24 \mathrm{E}-05$ | 11.9033 |
| 1.00 | $5.00 \mathrm{E}-09$ | 0.0014 | 8.4047 | $1.24 \mathrm{E}-06$ | $4.32 \mathrm{E}-06$ | 8.3264 |
| 1.10 | $2.83 \mathrm{E}-10$ | 0.0003 | 5.7753 | $1.09 \mathrm{E}-06$ | $1.69 \mathrm{E}-07$ | 5.6746 |
| 1.20 | $1.54 \mathrm{E}-11$ | $6.94 \mathrm{E}-05$ | 3.8761 | $9.43 \mathrm{E}-07$ | $4.94 \mathrm{E}-09$ | 3.7666 |
| 1.30 | $8.14 \mathrm{E}-13$ | $1.46 \mathrm{E}-05$ | 2.5411 | $8.08 \mathrm{E}-07$ | $1.08 \mathrm{E}-10$ | 2.4340 |
| 1.40 | $4.17 \mathrm{E}-14$ | $3.00 \mathrm{E}-06$ | 1.6271 | $6.80 \mathrm{E}-07$ | $1.74 \mathrm{E}-12$ | 1.5307 |
| 1.50 | $2.08 \mathrm{E}-15$ | $6.00 \mathrm{E}-07$ | 1.0169 | $5.58 \mathrm{E}-07$ | $0.00 \mathrm{E}+00$ | 0.9360 |
| 1.60 | $1.03 \mathrm{E}-16$ | $1.17 \mathrm{E}-07$ | 0.6187 | $4.41 \mathrm{E}-07$ | $0.00 \mathrm{E}+00$ | 0.5551 |
| 1.70 | $4.85 \mathrm{E}-18$ | $2.25 \mathrm{E}-08$ | 0.3628 | $3.27 \mathrm{E}-07$ | $0.00 \mathrm{E}+00$ | 0.3167 |
| 1.80 | $2.27 \mathrm{E}-19$ | $4.24 \mathrm{E}-09$ | 0.1981 | $2.17 \mathrm{E}-07$ | $0.00 \mathrm{E}+00$ | 0.1684 |
| 1.90 | $1.04 \mathrm{E}-20$ | $7.61 \mathrm{E}-10$ | 0.0868 | $1.08 \mathrm{E}-07$ | $-1.69 \mathrm{E}-14$ | 0.0723 |
| 2.00 | 0 | 0 | 0 | $-8.39 \mathrm{E}-15$ | $-4.03 \mathrm{E}-15$ | $-3.85 \mathrm{E}-15$ |

## Chapter 6

## FINITE ELEMENT EQUATIONS

Finite element equations capture the characteristics of the field equations. Their derivation is based on either the governing differential equation or the global energy balance of the physical problem. The approach involving the governing differential equation is referred to as the method of weighted residuals or Galerkin's method. The approach utilizing the global energy balance is referred to as the variational method or Rayleigh-Ritz method.

### 6.1 Method of Weighted Residuals

The method of weighted residuals involves the approximation of the functional behavior of the dependent variable in the governing differential equation (Finlayson 1972). When substituted into the governing differential equation, the approximate form of the dependent variable leads to an error called the "residual." This residual error is required to vanish in a weighted average sense over the domain. If the weighting functions are chosen to be the same as the element shape (interpolation) functions used in the element approximation functions, the method of weighted residuals is referred to as Galerkin's method.

The governing differential equation for the physical problem in domain $D$ described in Fig. 6.1 can be expressed in the form

$$
\begin{equation*}
L(\phi)-f=0 \tag{6.1}
\end{equation*}
$$

where $\phi$ is a dependent variable and $f$ is a known forcing function. The ordinary or partial differential operator, $L$ whose order is specified by $p$, can be linear or nonlinear. The boundary conditions are given by

$$
\begin{equation*}
B_{j}(\phi)=g_{j} \text { on } C_{1} \tag{6.2}
\end{equation*}
$$

and

$$
\begin{equation*}
E_{j}(\phi)=h_{j} \text { on } C_{2} \tag{6.3}
\end{equation*}
$$

in which $B_{j}$ and $E_{j}$ are operators, with $j=1,2,3, \ldots, p$. The known functions $g_{j}$ and $h_{j}$ prescribe the boundary conditions on the dependent vari-


Fig. 6.1 Variation of the dependent (field) variable over a twodimensional domain under specified boundary conditions.
able and its derivatives, respectively. The conditions on the dependent variable over $C_{1}$ are referred to as essential or forced boundary conditions, and the ones involving the derivatives of the dependent variable over $C_{2}$ are referred to as natural boundary conditions.

The method of weighted residuals requires that

$$
\begin{equation*}
\int_{D}[L(\tilde{\phi})-f] W_{k} d D=0, \text { with } k=1,2,3, \ldots, n \tag{6.4}
\end{equation*}
$$

where $W_{k}$ are the weighting functions approximating the dependent variable as

$$
\begin{equation*}
\phi \approx \tilde{\phi}=\sum_{k=1}^{n} \alpha_{k} W_{k} \tag{6.5}
\end{equation*}
$$

while satisfying the essential boundary conditions on $C_{1}$. The unknown coefficients, $\alpha_{k}$, are determined by solving for the resulting system of algebraic equations.

Since the governing differential equation is valid for the entire domain, $D$, partitioning the domain into subdomains or elements, $D^{(e)}$, and applying

Galerkin's method with weighting functions $W_{k}=N_{k}^{(e)}$ over the element domain results in

$$
\begin{equation*}
\sum_{e=1}^{E} \int_{D^{(e)}} \mathbf{N}^{(e)}\left(L\left(\tilde{\phi}^{(e)}\right)-f\right) d D=0 \tag{6.6}
\end{equation*}
$$

in which $E$ is the number of elements and the superscript " $e$ " denotes a specific element whose domain is $D^{(e)}$. The approximation to the dependent variable within the element can be expressed as

$$
\begin{equation*}
\tilde{\phi}^{(e)}=\sum_{i=1}^{n} N_{i}^{(e)} \phi_{i}^{(e)} \tag{6.7}
\end{equation*}
$$

or

$$
\begin{equation*}
\tilde{\phi}^{(e)}=\mathbf{N}^{(e) T} \varphi^{(e)} \tag{6.8}
\end{equation*}
$$

where

$$
\mathbf{N}^{(e) T}=\left\{\begin{array}{lllll}
N_{1}^{(e)} & N_{2}^{(e)} & N_{3}^{(e)} & \ldots & N_{n}^{(e)} \tag{6.9}
\end{array}\right\}
$$

and

$$
\varphi^{(e) T}=\left\{\begin{array}{lllll}
\phi_{1}^{(e)} & \phi_{2}^{(e)} & \phi_{3}^{(e)} & \ldots & \phi_{n}^{(e)} \tag{6.10}
\end{array}\right\}
$$

with $n$ representing the number of nodes associated with element $e$. The nodal unknowns and shape functions are denoted by $\phi_{i}^{(e)}$ and $N_{i}^{(e)}$, with $i=1,2, . ., n$, respectively. The shape functions need not satisfy the boundary conditions; however, they satisfy the inter-element continuity conditions necessary for assembly of the element equations. The essential boundary conditions are imposed after assembling the global matrix. The natural boundary conditions are not imposed directly. However, their influence emerges in the derivation of the element equations.

The required order of the element continuity is equal to one less than the highest derivative of the dependent variable appearing in the integrand. This requirement is relaxed by applying integration by parts in the minimization procedure of the residual error in Galerkin's method.

### 6.1.1 Example: One-dimensional Differential Equation with Line Elements

The application of Galerkin's method is introduced by considering the ordinary differential equation given by

$$
\begin{equation*}
\frac{d^{2} \phi(x)}{d x^{2}}+\phi(x)-f(x)=0 \tag{6.11}
\end{equation*}
$$

in domain $D$ defined by $0 \leq x \leq 1$. The known forcing function is given by

$$
\begin{equation*}
f(x)=-x \tag{6.12}
\end{equation*}
$$

The boundary conditions, identified as the essential type, are $\phi(0)=0$ and $\phi(1)=0$. As shown in Fig. 6.2, the domain can be discretized with $E$ linear line elements, each having two nodes ( $n=2$ ). There are a total of $N$ nodes, and global coordinates of each node in domain $D$ are specified by $x_{i}$, with $i=1,2, \ldots, N$. Nodal values of the dependent variable associated with element $e$ are specified at its first and second nodes by $\phi_{1}^{(e)}$ and $\phi_{2}^{(e)}$, respectively.

The linear approximation function for the dependent variable in element $e$ can be expressed in the form

$$
\begin{equation*}
\tilde{\phi}^{(e)}=N_{1}^{(e)} \phi_{1}^{(e)}+N_{2}^{(e)} \phi_{2}^{(e)} \tag{6.13}
\end{equation*}
$$

or

$$
\begin{equation*}
\tilde{\phi}^{(e)}=\mathbf{N}^{(e) T} \varphi^{(e)} \tag{6.14}
\end{equation*}
$$

where

$$
\mathbf{N}^{(e) T}=\left\{\begin{array}{ll}
N_{1}^{(e)} & N_{2}^{(e)}
\end{array}\right\} \text { and } \varphi^{(e) T}=\left\{\begin{array}{ll}
\phi_{1}^{(e)} & \phi_{2}^{(e)} \tag{6.15}
\end{array}\right\}
$$

in which the shape functions are given by

$$
\begin{equation*}
N_{1}^{(e)}=\frac{x_{2}^{(e)}-x}{x_{2}^{(e)}-x_{1}^{(e)}} \text { and } N_{2}^{(e)}=\frac{x-x_{1}^{(e)}}{x_{2}^{(e)}-x_{1}^{(e)}} \tag{6.16}
\end{equation*}
$$



Fig. 6.2 Domain of the one-dimensional differential equation, discretized into $E$ elements.

They are the same as the length coordinates given by Eq. (3.9). Applying Galerkin's method by Eq. (6.6) leads to

$$
\begin{equation*}
\sum_{e=1}^{E} \int_{\substack{x_{1}^{(e)}}}^{x_{2}^{(e)}} \mathbf{N}^{(e)}\left(\frac{d^{2} \tilde{\phi}^{(e)}(x)}{d x^{2}}+\tilde{\phi}^{(e)}(x)-f(x)\right) d x=0 \tag{6.17}
\end{equation*}
$$

Integrating the first term in the integral by parts results in

$$
\begin{align*}
& \sum_{e=1}^{E}\left[\left.\mathbf{N}^{(e)} \frac{d \tilde{\phi}^{(e)}(x)}{d x}\right|_{x_{1}^{(e)}} ^{x_{2}^{(e)}}-\int_{x_{1}^{(e)}}^{x_{2}^{(e)}} \frac{d \mathbf{N}^{(e)}}{d x} \frac{d \tilde{\phi}^{(e)}}{d x} d x\right.  \tag{6.18}\\
& \left.\quad+\int_{x_{1}^{(e)}}^{x_{2}^{(e)}} \mathbf{N}^{(e)} \tilde{\phi}^{(e)}(x) d x-\int_{x_{1}^{(e)}}^{x_{2}^{(e)}} \mathbf{N}^{(e)} f(x) d x\right]=0
\end{align*}
$$

Substituting for the element approximation function $\left(\tilde{\phi}^{(e)}=\mathbf{N}^{(e) T} \boldsymbol{\varphi}^{(e)}\right)$ yields

$$
\begin{equation*}
\sum_{e=1}^{E} \mathbf{k}^{(e)} \boldsymbol{\varphi}^{(e)}=\sum_{e=1}^{E} \mathbf{f}^{(e)} \tag{6.19}
\end{equation*}
$$

where

$$
\begin{equation*}
\mathbf{k}^{(e)}=-\int_{x_{1}^{(e)}}^{x_{2}^{(e)}} \frac{d \mathbf{N}^{(e)}}{d x} \frac{d \mathbf{N}^{(e) T}}{d x} d x+\int_{x_{1}^{(e)}}^{x_{2}^{(e)}} \mathbf{N}^{(e)} \mathbf{N}^{(e) T} d x \tag{6.20}
\end{equation*}
$$

and

$$
\begin{equation*}
\mathbf{f}^{(e)}=\int_{x_{1}^{(e)}}^{x_{2}^{(e)}} \mathbf{N}^{(e)} f(x) d x-\left[\mathbf{N}^{(e)} \frac{d \tilde{\phi}^{(e)}(x)}{d x}\right]_{x_{1}^{(e)}}^{x_{2}^{(e)}} \tag{6.21}
\end{equation*}
$$

After substituting for the shape functions and their derivatives, as well as the forcing function, the expressions for the element characteristic matrix, $\mathbf{k}^{(e)}$, and the right-hand-side vector, $\mathbf{f}^{(e)}$, become

$$
\begin{align*}
\mathbf{k}^{(e)}= & -\frac{1}{\left(x_{2}^{(e)}-x_{1}^{(e)}\right)^{2}} \int_{x_{1}^{(e)}}^{x_{2}^{(e)}}\left[\begin{array}{cc}
1 & -1 \\
-1 & 1
\end{array}\right] d x \\
& +\int_{x_{1}^{(e)}}^{x_{2}^{(e)}}\left[\begin{array}{ll}
N_{1}^{(e)} N_{1}^{(e)} & N_{1}^{(e)} N_{2}^{(e)} \\
N_{2}^{(e)} N_{1}^{(e)} & N_{2}^{(e)} N_{2}^{(e)}
\end{array}\right] d x  \tag{6.22}\\
\mathbf{f}^{(e)}= & -\int_{x_{1}^{(e)}}^{x_{2}^{(e)}}\left\{\begin{array}{l}
N_{1}^{(e)} \\
N_{2}^{(e)}
\end{array}\right\} x d x-\left[\left\{\begin{array}{l}
N_{1}^{(e)} \\
N_{2}^{(e)}
\end{array}\right\} \frac{d \tilde{\phi}^{(e)}(x)}{d x}\right]_{x_{1}^{(e)}}^{x_{2}^{(e)}} \tag{6.23}
\end{align*}
$$

Evaluation of these integrals leads to the final form of the element characteristic matrix, $\mathbf{k}^{(e)}$, and the right-hand-side vector, $\mathbf{f}^{(e)}$

$$
\mathbf{k}^{(e)}=-\frac{1}{\left(x_{2}^{(e)}-x_{1}^{(e)}\right)}\left[\begin{array}{cc}
1 & -1  \tag{6.24}\\
-1 & 1
\end{array}\right]+\frac{\left(x_{2}^{(e)}-x_{1}^{(e)}\right)}{6}\left[\begin{array}{ll}
2 & 1 \\
1 & 2
\end{array}\right]
$$

and

$$
\begin{aligned}
\mathbf{f}^{(e)}= & -\frac{1}{6}\left(x_{2}^{(e)}-x_{1}^{(e)}\right)\left\{\begin{array}{l}
2 x_{1}^{(e)}+x_{2}^{(e)} \\
x_{1}^{(e)}+2 x_{2}^{(e)}
\end{array}\right\} \\
& -\left[\frac{d \tilde{\phi}^{(e)}\left(x_{2}^{(e)}\right)}{d x}\left\{\begin{array}{l}
N_{1}^{(e)}\left(x_{2}^{(e)}\right) \\
N_{2}^{(e)}\left(x_{2}^{(e)}\right)
\end{array}\right\}-\frac{d \tilde{\phi}^{(e)}\left(x_{1}^{(e)}\right)}{d x}\left\{\begin{array}{l}
N_{1}^{(e)}\left(x_{1}^{(e)}\right) \\
N_{2}^{(e)}\left(x_{1}^{(e)}\right)
\end{array}\right\}\right]
\end{aligned}
$$

or

$$
\mathbf{f}^{(e)}=-\frac{1}{6}\left(x_{2}^{(e)}-x_{1}^{(e)}\right)\left\{\begin{array}{c}
2 x_{1}^{(e)}+x_{2}^{(e)}  \tag{6.26}\\
x_{1}^{(e)}+2 x_{2}^{(e)}
\end{array}\right\}-\left\{\begin{array}{l}
-\frac{d \tilde{\phi}^{(e)}}{d x}\left(x_{1}^{(e)}\right) \\
\frac{d \tilde{\phi}^{(e)}}{d x}\left(x_{2}^{(e)}\right)
\end{array}\right\}
$$

The local and global nodes for the domain discretized with three elements, $E=3$, and four nodes, $N=4$, are numbered as shown in Table 6.1.

Table 6.1 Element connectivity and nodal coordinates.

| Element <br> Number <br> $(e)$ | Node 1 | Node 2 | $x_{1}^{(e)}$ | $x_{2}^{(e)}$ |
| :---: | :---: | :---: | :---: | :---: |
| 1 | 1 | 2 | 0 | $1 / 3$ |
| 2 | 2 | 3 | $1 / 3$ | $2 / 3$ |
| 3 | 3 | 4 | $2 / 3$ | 1 |

With the appropriate value of the nodal coordinates from Eq. (6.24) and (6.26), the element characteristic matrices and vectors are calculated as

$$
\begin{gather*}
\mathbf{k}^{(1)}=\frac{1}{18}\left[\begin{array}{cc}
52 & -55 \\
-55 & 52
\end{array}\right] \\
\mathbf{k}^{(2)}=\frac{1}{18}\left[\begin{array}{cc}
52 & -55 \\
-55 & 52
\end{array}\right]  \tag{6.27}\\
\mathbf{k}^{(3)}=\frac{1}{18}\left[\begin{array}{cc}
52 & -55 \\
-55 & 52
\end{array}\right] \\
\mathbf{f}^{(1)}=\frac{1}{54}\left\{\begin{array}{l}
3 \\
2
\end{array}\right\}+\left\{\begin{array}{l}
-\frac{d \tilde{\phi}^{(1)}(0)}{d x} \\
\frac{d \tilde{\phi}^{(1)}(1 / 3)}{d x}
\end{array}\right\}\left[\begin{array}{l}
2 \\
\mathbf{f}^{(2)}=
\end{array}\right.  \tag{6.28}\\
\frac{1}{54}\left\{\begin{array}{l}
4 \\
5
\end{array}\right\}+\left\{\begin{array}{l}
\frac{d \tilde{\phi}^{(2)}(1 / 3)}{d x} \\
\frac{d \tilde{\phi}^{(2)}(2 / 3)}{d x}
\end{array}\right\}
\end{gather*}
$$

$$
\mathbf{f}^{(3)}=\frac{1}{54}\left\{\begin{array}{l}
7  \tag{6.32}\\
8
\end{array}\right\}+\left\{\begin{array}{c}
-\frac{d \tilde{\phi}^{(3)}(2 / 3)}{d x} \\
\frac{d \tilde{\phi}^{(3)}(1)}{d x}
\end{array}\right\}
$$

As reflected by the element connectivity in Table 6.1 , the boxed numbers indicate the rows and columns of the global matrix, $\mathbf{K}$, and global right-hand-side vector, $\mathbf{F}$, to which the individual coefficients are added. The global coefficient matrix, $\mathbf{K}$, and the global right-hand-side vector, $\mathbf{F}$, are obtained from the "expanded" element coefficient matrices, $\mathbf{k}^{(e)}$, and the element right-hand-side vectors, $\mathbf{f}^{(e)}$, by summation in the form

$$
\begin{equation*}
\mathbf{K}=\sum_{e=1}^{E} \mathbf{k}^{(e)} \quad \text { and } \quad \mathbf{F}=\sum_{e=1}^{E} \mathbf{f}^{(e)} \tag{6.33}
\end{equation*}
$$

The "expanded" element matrices are the same size as the global matrix but have rows and columns of zeros corresponding to the nodes not associated with element (e). Specifically, the expanded form of the element stiffness and load vector becomes

$$
\begin{align*}
& \text { (1) 2 } 3 \text { 4 } \\
& \mathbf{k}^{(1)}=\frac{1}{18}\left[\begin{array}{cccc}
52 & -55 & 0 & 0 \\
-55 & 52 & 0 & 0 \\
0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0
\end{array}\right] \begin{array}{l}
1 \\
2 \\
3 \\
4
\end{array}  \tag{6.34}\\
& \mathbf{f}^{(1)}=\frac{1}{54}\left\{\begin{array}{l}
1 \\
2 \\
0 \\
0
\end{array}\right\}+\left\{\begin{array}{c}
-\frac{d \tilde{\phi}^{(1)}(0)}{d x} \\
\frac{d \tilde{\phi}^{(1)}(1 / 3)}{d x} \\
0 \\
0
\end{array}\right] \begin{array}{l}
1 \\
\frac{2}{3} \\
{[ }
\end{array} \tag{6.35}
\end{align*}
$$

$$
\begin{align*}
& \mathbf{k}^{(2)}=\frac{1}{18}\left[\begin{array}{cccc}
1 & 2 & 3 & 4 \\
0 & 0 & 0 & 0 \\
0 & 52 & -55 & 0 \\
0 & -55 & 52 & 0 \\
0 & 0 & 0 & 0
\end{array}\right]\left[\begin{array}{l}
1 \\
2
\end{array}\right]\left[\begin{array}{l}
2 \\
3
\end{array}\right.  \tag{6.36}\\
& \mathbf{f}^{(2)}=\frac{1}{54}\left\{\begin{array}{l}
0 \\
4 \\
5 \\
0
\end{array}\right\}+\left\{\begin{array}{c|c}
0 & 1 \\
-\frac{d \tilde{\phi}^{(2)}(1 / 3)}{d x} & 2 \\
\frac{d \tilde{\phi}^{(2)}(2 / 3)}{d x} & \begin{array}{l}
2 \\
0
\end{array} \\
\boxed{3}
\end{array}\right.  \tag{6.37}\\
& \text { (1) 2) } 3 \text { 4 } \\
& \mathbf{k}^{(3)}=\frac{1}{18}\left[\begin{array}{cccc}
0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 \\
0 & 0 & 52 & -55 \\
0 & 0 & -55 & 52
\end{array}\right] \frac{1}{2}  \tag{6.38}\\
& \mathbf{f}^{(3)}=\frac{1}{54}\left[\begin{array}{l}
0 \\
0 \\
7 \\
8
\end{array}\right\}+\left\{\begin{array}{c}
0 \\
0 \\
\frac{d \tilde{\phi}^{(3)}(2 / 3)}{d x} \\
\frac{d \tilde{\phi}^{(3)}(1)}{d x}
\end{array}\right\} \begin{array}{l}
1 \\
2 \\
3
\end{array} \tag{6.39}
\end{align*}
$$

In accordance with Eq. (6.33) and (6.19), the assembly of the element characteristic matrices and vectors results in the global equilibrium equations

$$
\begin{align*}
& \frac{1}{18}\left[\begin{array}{cccc}
52 & -55 & 0 & 0 \\
-55 & 52+52 & -55 & 0 \\
0 & -55 & 52+52 & -55 \\
0 & 0 & -55 & 52
\end{array}\right]\left\{\begin{array}{c}
\phi_{1}=\phi_{1}^{(1)} \\
\phi_{2}=\phi_{2}^{(1)}=\phi_{1}^{(2)} \\
\phi_{3}=\phi_{2}^{(2)}=\phi_{1}^{(3)} \\
\phi_{4}=\phi_{2}^{(3)}
\end{array}\right\}=\frac{1}{54}\left\{\begin{array}{c}
1 \\
2+4 \\
5+7 \\
8
\end{array}\right\} \\
& +\left\{\begin{array}{c}
-\frac{d \tilde{\phi}^{(1)}(0)}{d x} \\
\frac{\frac{d \tilde{\phi}^{(1)}(1 / 3)}{\frac{d \tilde{\phi}^{(2)}(2 / 3)}{d x} \frac{d \tilde{\phi}^{(2)}(1 / 3)}{d x}}}{\frac{d \tilde{\phi}^{(3)}(2 / 3)}{d x}} 0 \\
\frac{d \tilde{\phi}^{(3)}(1)}{d x}
\end{array}\right\} \tag{6.40}
\end{align*}
$$

or

$$
\frac{1}{18}\left[\begin{array}{cccc}
52 & -55 & 0 & 0  \tag{6.41}\\
-55 & 104 & -55 & 0 \\
0 & -55 & 104 & -55 \\
0 & 0 & -55 & 52
\end{array}\right]\left\{\begin{array}{c}
\phi_{1} \\
\phi_{2} \\
\phi_{3} \\
\phi_{4}
\end{array}\right\}=\frac{1}{54}\left\{\begin{array}{c}
1 \\
6 \\
12 \\
8
\end{array}\right\}+\left\{\begin{array}{c}
-\frac{d \tilde{\phi}^{(1)}(0)}{d x} \\
0 \\
0 \\
\frac{d \tilde{\phi}^{(3)}(1)}{d x}
\end{array}\right\}
$$

or

$$
\begin{equation*}
\mathbf{K} \Phi=\mathbf{F} \tag{6.42}
\end{equation*}
$$

After imposing the essential boundary conditions, $\phi_{1}=0$ and $\phi_{4}=0$, the global system of equations is reduced by deleting the row and column corresponding to $\phi_{1}$ and $\phi_{4}$, leading to

$$
\frac{1}{18}\left[\begin{array}{cc}
104 & -55  \tag{6.43}\\
-55 & 104
\end{array}\right]\left\{\begin{array}{l}
\phi_{2} \\
\phi_{3}
\end{array}\right\}=\frac{1}{54}\left\{\begin{array}{c}
6 \\
12
\end{array}\right\}
$$

Its solution yields

$$
\left\{\begin{array}{l}
\phi_{2}  \tag{6.44}\\
\phi_{3}
\end{array}\right\}=\left\{\begin{array}{l}
0.05493 \\
0.06751
\end{array}\right\}
$$

The exact solution to the differential equation given by

$$
\begin{equation*}
\phi(x)=\frac{\sin (x)}{\sin (1)}-x \tag{6.45}
\end{equation*}
$$

provides

$$
\left\{\begin{array}{l}
\phi_{2}  \tag{6.46}\\
\phi_{3}
\end{array}\right\}=\left\{\begin{array}{l}
0.0555 \\
0.0682
\end{array}\right\}
$$

The exact and FEM calculations of $\phi$ along the $x$-axis are shown in Fig. 6.3.

### 6.1.2 Example: Two-dimensional Differential Equation with Linear Triangular Elements

### 6.1.2.1 Galerkin's Method

The application of Galerkin's method in solving two-dimensional problems with linear triangular elements is demonstrated by considering the partial differential equation given by

$$
\begin{equation*}
\frac{\partial^{2} \phi(x, y)}{\partial x^{2}}+\frac{\partial^{2} \phi(x, y)}{\partial y^{2}}+A=0 \tag{6.47}
\end{equation*}
$$

in domain $D$, defined by the intersection of $y=0, y=2-\sqrt{3} x$, and $y=\sqrt{3} x$ (as shown in Fig. 6.4), where $A=1$.

The boundary conditions are specified as

$$
\begin{align*}
& -n_{y} \frac{\partial \phi(x, y=0)}{\partial y}=[\phi(x, y=0)-(B=1)] \quad \text { for } 0 \leq x \leq 2 / \sqrt{3}  \tag{6.48}\\
& \phi(x, y=\sqrt{3} x)=0 \quad \text { for } 0 \leq x \leq 1 / \sqrt{3}  \tag{6.49}\\
& \phi(x, y=2-\sqrt{3} x)=0 \quad \text { for } 1 / \sqrt{3} \leq x \leq 2 / \sqrt{3} \tag{6.50}
\end{align*}
$$

When independent of time, these equations provide the temperature field, $\phi(x, y)$, due to heat conduction in a domain having a heat generation of $A$ with one of its boundaries subjected to a convective heat transfer. The thermal conductivity and the film (surface) heat transfer coefficient are equal to unity, and the temperature of the surrounding medium is $B$.


Fig. 6.3 Comparison of the exact and FEA (approximate) solutions to the 1 D differential equation.


Fig. 6.4 The equilateral triangular domain.

The triangular domain can be discretized into four linear triangular elements, each having three nodes identified as 1,2 , and 3 (local node numbering), as illustrated in Fig. 6.5.

As shown in Fig. 6.6, the global coordinates of each node in domain $D$ are specified by $\left(x_{i}, y_{i}\right)$, with $i=1,2,3,4$, and 5 . These coordinates are presented in Table 6.2.


Fig. 6.5 Local node numbering for the linear triangular element.


Fig. 6.6 Finite element discretization of the domain.

Table 6.2 Nodal coordinates.

| Global Node <br> Number | Nodal <br> Coordinates | Nodal <br> Unknowns |
| :---: | :---: | :---: |
| 1 | $(0,0)$ | $\phi_{1}$ |
| 2 | $(2 / \sqrt{3}, 0)$ | $\phi_{2}$ |
| 3 | $(1 / \sqrt{3}, 1)$ | $\phi_{3}$ |
| 4 | $(1 / \sqrt{3}, 1 / 3)$ | $\phi_{4}$ |
| 5 | $(1 / \sqrt{3}, 0)$ | $\phi_{5}$ |

The nodal values of the dependent variable associated with the global coordinates are denoted by $\phi_{i}(i=1,2,3,4$, and 5). As shown in Fig. 6.5, the nodal values of the dependent variable associated with element $e$ are specified at its first, second, and third nodes by $\phi_{1}^{(e)}, \phi_{2}^{(e)}$, and $\phi_{3}^{(e)}$, respectively.

The linear element approximation function for the dependent field variable in a triangular element " $e$ " is written as

$$
\begin{equation*}
\tilde{\phi}^{(e)}=N_{1}^{(e)} \phi_{1}^{(e)}+N_{2}^{(e)} \phi_{2}^{(e)}+N_{3}^{(e)} \phi_{3}^{(e)} \tag{6.51}
\end{equation*}
$$

or

$$
\begin{equation*}
\tilde{\phi}^{(e)}=\mathbf{N}^{(e) T} \boldsymbol{\varphi}^{(e)} \tag{6.52}
\end{equation*}
$$

As derived in Chap. 3, the element shape functions in Eq. (3.17) are taken as

$$
\left\{\begin{array}{l}
N_{1}^{(e)}  \tag{6.53}\\
N_{2}^{(e)} \\
N_{3}^{(e)}
\end{array}\right\}=\frac{1}{2 \Delta^{(e)}}\left[\begin{array}{ccc}
\left(x_{2} y_{3}-x_{3} y_{2}\right) & y_{23} & x_{32} \\
\left(x_{3} y_{1}-x_{1} y_{3}\right) & y_{31} & x_{13} \\
\left(x_{1} y_{2}-x_{2} y_{1}\right) & y_{12} & x_{21}
\end{array}\right]\left\{\begin{array}{c}
1 \\
x \\
y
\end{array}\right\}
$$

where $x_{m n}=x_{m}-x_{n}, y_{m n}=y_{m}-y_{n}$, and $\Delta^{(e)}$ is the area of the element computed by

$$
2 \Delta^{(e)}=\left|\begin{array}{ccc}
1 & 1 & 1  \tag{6.54}\\
x_{1} & x_{2} & x_{3} \\
y_{1} & y_{2} & y_{3}
\end{array}\right|
$$

Applying Eq. (6.6), Galerkin's method, leads to

$$
\begin{equation*}
\sum_{e=1}^{E} \int_{D^{(e)}} \mathbf{N}^{(e)}\left(\frac{\partial^{2} \tilde{\phi}^{(e)}(x, y)}{\partial x^{2}}+\frac{\partial^{2} \tilde{\phi}^{(e)}(x, y)}{\partial y^{2}}+A\right) d x d y=0 \tag{6.55}
\end{equation*}
$$

Since the element approximation function is $C^{0}$ continuous, the secondorder derivatives in the integrand must be reduced by one so that the interelement continuity is achieved during the assembly of the global matrix. This reduction is achieved by observing that

$$
\begin{equation*}
\mathbf{N}^{(e)} \frac{\partial^{2} \tilde{\phi}^{(e)}}{\partial x^{2}}(x, y)=\frac{\partial}{\partial x}\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}(x, y)\right)-\frac{\partial \mathbf{N}^{(e)}}{\partial x} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}(x, y) \tag{6.56}
\end{equation*}
$$

and

$$
\begin{equation*}
\mathbf{N}^{(e)} \frac{\partial^{2} \tilde{\phi}^{(e)}}{\partial y^{2}}(x, y)=\frac{\partial}{\partial y}\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}(x, y)\right)-\frac{\partial \mathbf{N}^{(e)}}{\partial y} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}(x, y) \tag{6.5}
\end{equation*}
$$

Their substitution into the integrand in Eq. (6.55) and rearrangement of the terms result in

$$
\begin{align*}
\sum_{e=1}^{E}\left\{\int_{D^{(e)}}[ \right. & \left.\frac{\partial}{\partial x}\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}\right)+\frac{\partial}{\partial y}\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}\right)\right] d x d y \\
& \left.+\int_{D^{(e)}}\left[-\frac{\partial \mathbf{N}^{(e)}}{\partial x} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}-\frac{\partial \mathbf{N}^{(e)}}{\partial y} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}+\mathbf{N}^{(e)} A\right] d x d y\right\}=0 \tag{6.58}
\end{align*}
$$

Applying the divergence theorem to the first integral renders the domain integral to the boundary integral, and it yields

$$
\begin{align*}
\sum_{e=1}^{E}\left\{\oint_{C^{(e)}}\right. & {\left[\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}\right) n_{x}^{(e)}+\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}\right) n_{y}^{(e)}\right] d s }  \tag{6.59}\\
& \left.+\int_{D^{(e)}}\left[-\frac{\partial \mathbf{N}^{(e)}}{\partial x} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}-\frac{\partial \mathbf{N}^{(e)}}{\partial y} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}+\mathbf{N}^{(e)} A\right] d x d y\right\}=0
\end{align*}
$$

where $n_{x}^{(e)}$ and $n_{y}^{(e)}$ are, respectively, the $x$ - and $y$-components of the outward normal vector along the closed boundary defining the area of the element, $C^{(e)}$.

Substituting for the element approximation function yields

$$
\begin{align*}
& \sum_{e=1}^{E}\left\{\oint_{C^{(e)}} \mathbf{N}^{(e)}\left[\frac{\partial \tilde{\phi}^{(e)}}{\partial x} n_{x}^{(e)}+\frac{\partial \tilde{\phi}^{(e)}}{\partial y} n_{y}^{(e)}\right] d s\right. \\
&-\int_{D^{(e)}}\left[\frac{\partial \mathbf{N}^{(e)}}{\partial x} \frac{\partial \mathbf{N}^{(e) T}}{\partial x}+\frac{\partial \mathbf{N}^{(e)}}{\partial y} \frac{\partial \mathbf{N}^{(e) T}}{\partial y}\right] d x d y \varphi^{(e)}  \tag{6.60}\\
&\left.+\int_{D^{(e)}} \mathbf{N}^{(e)} A d x d y\right\}=0
\end{align*}
$$

This equation can be recast in matrix form as

$$
\begin{equation*}
\sum_{e=1}^{E}\left(-\mathbf{k}^{(e)} \varphi^{(e)}+\mathbf{f}^{(e)}+\mathbf{Q}^{(e)}\right)=0 \tag{6.61}
\end{equation*}
$$

where

$$
\begin{gather*}
\mathbf{k}^{(e)}=\int_{D^{(e)}}\left[\frac{\partial \mathbf{N}^{(e)}}{\partial x} \frac{\partial \mathbf{N}^{(e) T}}{\partial x}+\frac{\partial \mathbf{N}^{(e)}}{\partial y} \frac{\partial \mathbf{N}^{(e) T}}{\partial y}\right] d x d y  \tag{6.62}\\
\mathbf{f}^{(e)}=\int_{D^{(e)}} \mathbf{N}^{(e)} A d x d y  \tag{6.63}\\
\mathbf{Q}^{(e)}=\oint_{C^{(e)}} \mathbf{N}^{(e)}\left[\frac{\partial \tilde{\phi}^{(e)}}{\partial x} n_{x}^{(e)}+\frac{\partial \tilde{\phi}^{(e)}}{\partial y} n_{y}^{(e)}\right] d s \tag{6.64}
\end{gather*}
$$

in which $\mathbf{k}^{(e)}$ is the element characteristic matrix, $\mathbf{f}^{(e)}$ is the element right-hand-side vector, and $\mathbf{Q}^{(e)}$ is often referred to as the inter-element vector that includes the derivative terms along the boundary of the element. The boundary integral around each element is evaluated in a counterclockwise direction, i.e., this boundary integral is the sum of three integrals taken along each side of the element.

Depending on whether the element has an exterior boundary or not, the inter-element vector is divided into two parts

$$
\begin{equation*}
\mathbf{Q}^{(e)}=\mathbf{Q}_{e}^{(e)}+\mathbf{Q}_{i}^{(e)} \tag{6.65}
\end{equation*}
$$

in which $\mathbf{Q}_{e}^{(e)}$ represents the contribution of the derivative terms specified along the external boundary of the element $C_{e}^{(e)}$, and $\mathbf{Q}_{i}^{(e)}$ represents the contribution from the internal boundaries of the element shared with other adjacent elements. Because each of the boundary integrals is evaluated in a counterclockwise direction, the contributions coming from the vector $\mathbf{Q}_{i}^{(e)}$ vanish when the global system of equations are assembled, thus no further discussion is necessary. However, in the case of specified derivative boundary conditions, the contribution coming from $\mathbf{Q}_{e}^{(e)}$ must be included.

In view of the boundary conditions given by Eq. (6.48) and the discretization of the domain, the $1-5$ side of element 1 and the 5-2 side of element 2 are subjected to derivative boundary conditions.

With $n_{x}^{(1)}=n_{x}^{(2)}=0$ and $n_{y}^{(1)}=n_{y}^{(2)}=-1$, the contribution of the derivative boundary conditions appearing in Eq. (6.64) leads to the inter-element vectors as

$$
\begin{equation*}
\mathbf{Q}_{e}^{(1)}=\oint_{C_{1-5}^{(1)}} \mathbf{N}^{(1)}\left[B-\phi_{C}\right] d s \quad \text { and } \quad \mathbf{Q}_{e}^{(2)}=\oint_{C_{S-2}^{(2)}} \mathbf{N}^{(2)}\left[B-\phi_{C}\right] d s \tag{6.66}
\end{equation*}
$$

where $\phi_{C}$ is the unknown value of the field variable on the external boundary of the element $C_{e}$, along which the derivative boundary condition is specified.

Approximating the unknown field variable, $\phi_{C}$, by $\tilde{\phi}^{(e)}=\mathbf{N}^{(e) T} \boldsymbol{\varphi}^{(e)}$ in these equations leads to

$$
\begin{equation*}
\mathbf{Q}_{e}^{(1)}=\oint_{C_{1-5}^{(1)}} \mathbf{N}^{(1)}\left[B-\mathbf{N}^{(1) T} \boldsymbol{\varphi}^{(1)}\right] d s \tag{6.67a}
\end{equation*}
$$

and

$$
\begin{equation*}
\mathbf{Q}_{e}^{(2)}=\oint_{C_{s-2}^{(2)}} \mathbf{N}^{(2)}\left[B-\mathbf{N}^{(2) T} \boldsymbol{\varphi}^{(2)}\right] d s \tag{6.67b}
\end{equation*}
$$

which can be rewritten as

$$
\begin{equation*}
\mathbf{Q}_{e}^{(1)}=\oint_{C_{1-5}^{(1)}} \mathbf{N}^{(1)} B d s-\left\{\oint_{C_{1-5}^{(1)}} \mathbf{N}^{(1)} \mathbf{N}^{(1) T} d s\right\} \boldsymbol{\varphi}^{(1)} \text { or } \quad \mathbf{Q}_{e}^{(1)}=\mathbf{g}^{(1)}-\mathbf{h}^{(1)} \boldsymbol{\varphi}^{(1)} \tag{6.68}
\end{equation*}
$$

and

$$
\begin{equation*}
\mathbf{Q}_{e}^{(2)}=\oint_{C_{5-2}^{(2)}} \mathbf{N}^{(2)} B d s-\left\{\oint_{C_{5-2}^{(2)}} \mathbf{N}^{(2)} \mathbf{N}^{(2) T} d s\right\} \boldsymbol{\varphi}^{(2)} \tag{6.69a}
\end{equation*}
$$

or

$$
\begin{equation*}
\mathbf{Q}_{e}^{(2)}=\mathbf{g}^{(2)}-\mathbf{h}^{(2)} \boldsymbol{\varphi}^{(2)} \tag{6.69b}
\end{equation*}
$$

where

$$
\begin{equation*}
\mathbf{h}^{(1)}=\oint_{C_{1-5}^{(1)}} \mathbf{N}^{(1)} \mathbf{N}^{(1) T} d s \quad \text { and } \quad \mathbf{h}^{(2)}=\oint_{C_{5-2}^{(2)}} \mathbf{N}^{(2)} \mathbf{N}^{(2)^{T}} d s \tag{6.70}
\end{equation*}
$$

and

$$
\begin{equation*}
\mathbf{g}^{(1)}=\oint_{C_{1-5}^{(1)}} \mathbf{N}^{(1)} B d s \quad \text { and } \quad \mathbf{g}^{(2)}=\oint_{C_{5-2}^{(2)}} \mathbf{N}^{(2)} B d s \tag{6.71}
\end{equation*}
$$

With this representation of the inter-element vector, the element equilibrium equations given by Eq. (6.61) can be rewritten in their final form as

$$
\begin{align*}
& \left(\mathbf{k}^{(1)}+\mathbf{h}^{(1)}\right) \varphi^{(1)}=\mathbf{f}^{(1)}+\mathbf{g}(1) \\
& \left(\mathbf{k}^{(2)}+\mathbf{h}^{(2)}\right) \varphi^{(2)}=\mathbf{f}^{(2)}+\mathbf{g}^{(2)}  \tag{6.72}\\
& \mathbf{k}^{(3)} \varphi^{(3)}=\mathbf{f}^{(3)} \\
& \mathbf{k}^{(4)} \varphi^{(4)}=\mathbf{f}^{(4)}
\end{align*}
$$

With the derivatives of the shape functions obtained as

$$
\left\{\begin{array}{l}
\frac{\partial N_{1}^{(e)}}{\partial x}  \tag{6.73}\\
\frac{\partial N_{2}^{(e)}}{\partial x} \\
\frac{\partial N_{3}^{(e)}}{\partial x}
\end{array}\right\}=\frac{1}{2 \Delta^{(e)}}\left\{\begin{array}{l}
y_{23} \\
y_{31} \\
y_{12}
\end{array}\right\} \text { and }\left\{\begin{array}{l}
\frac{\partial N_{1}^{(e)}}{\partial y} \\
\frac{\partial N_{2}^{(e)}}{\partial y} \\
\frac{\partial N_{3}^{(e)}}{\partial y}
\end{array}\right\}=\frac{1}{2 \Delta^{(e)}}\left\{\begin{array}{l}
x_{32} \\
x_{13} \\
x_{21}
\end{array}\right\}
$$

the evaluation of the area integrals in Eq. (6.62) and (6.63) by using Eq. (3.19) leads to the final form of the element coefficient matrix, $\mathbf{k}^{(e)}$, and right-hand-side vector, $\mathbf{f}^{(e)}$

$$
\mathbf{k}^{(e)}=\frac{1}{4 \Delta^{(e)}}\left[\begin{array}{ccc}
x_{32}^{2}+y_{23}^{2} & x_{32} x_{13}+y_{23} y_{31} & x_{32} x_{21}+y_{23} y_{12}  \tag{6.74}\\
x_{32} x_{13}+y_{23} y_{31} & x_{13}^{2}+y_{31}^{2} & x_{13} x_{21}+y_{31} y_{12} \\
x_{32} x_{21}+y_{23} y_{12} & x_{13} x_{21}+y_{31} y_{12} & x_{21}^{2}+y_{12}^{2}
\end{array}\right]
$$

and

$$
\mathbf{f}^{(e)}=\frac{A \Delta^{(e)}}{3}\left\{\begin{array}{l}
1  \tag{6.75}\\
1 \\
1
\end{array}\right\}
$$

Their numerical evaluation results in

$$
\begin{align*}
& \mathbf{k}^{(1)}=\frac{\sqrt{3}}{6}\left[\begin{array}{ccc}
1 & -1 & 0 \\
-1 & 4 & -3 \\
0 & -3 & 3
\end{array}\right] \text { and } \mathbf{f}^{(1)}=\frac{1}{18 \sqrt{3}}\left\{\begin{array}{l}
1 \\
1 \\
1
\end{array}\right\}  \tag{6.76}\\
& \mathbf{k}^{(2)}=\frac{\sqrt{3}}{6}\left[\begin{array}{ccc}
4 & -1 & -3 \\
-1 & 1 & 0 \\
-3 & 0 & 3
\end{array}\right] \text { and } \mathbf{f}^{(2)}=\frac{1}{18 \sqrt{3}}\left\{\begin{array}{l}
1 \\
1 \\
1
\end{array}\right\}  \tag{6.77}\\
& \mathbf{k}^{(3)}=\frac{\sqrt{3}}{12}\left[\begin{array}{ccc}
4 & 2 & -6 \\
2 & 4 & -6 \\
-6 & -6 & 12
\end{array}\right] \text { and } \mathbf{f}^{(3)}=\frac{1}{9 \sqrt{3}}\left\{\begin{array}{l}
1 \\
1 \\
1
\end{array}\right\}  \tag{6.78}\\
& \mathbf{k}^{(4)}=\frac{\sqrt{3}}{12}\left[\begin{array}{ccc}
4 & 2 & -6 \\
2 & 4 & -6 \\
-6 & -6 & 12
\end{array}\right] \text { and } \mathbf{f}^{(4)}=\frac{1}{9 \sqrt{3}}\left\{\begin{array}{l}
1 \\
1 \\
1
\end{array}\right\} \tag{6.79}
\end{align*}
$$

in which the area of each element is computed as

$$
\begin{gather*}
\Delta^{(1)}=\frac{1}{2}\left|\begin{array}{ccc}
1 & 1 & 1 \\
0 & 1 / \sqrt{3} & 1 / \sqrt{3} \\
0 & 0 & 1 / 3
\end{array}\right|=\frac{1}{6 \sqrt{3}}  \tag{6.80}\\
\Delta^{(2)}=\frac{1}{2}\left|\begin{array}{ccc}
1 & 1 & 1 \\
1 / \sqrt{3} & 2 / \sqrt{3} & 1 / \sqrt{3} \\
0 & 0 & 1 / 3
\end{array}\right|=\frac{1}{6 \sqrt{3}} \tag{6.81}
\end{gather*}
$$

$$
\begin{gather*}
\Delta^{(3)}=\frac{1}{2}\left|\begin{array}{ccc}
1 & 1 & 1 \\
2 / \sqrt{3} & 1 / \sqrt{3} & 1 / \sqrt{3} \\
0 & 1 & 1 / 3
\end{array}\right|=\frac{1}{3 \sqrt{3}}  \tag{6.82}\\
\Delta^{(4)}=\frac{1}{2}\left|\begin{array}{ccc}
1 & 1 & 1 \\
1 / \sqrt{3} & 0 & 1 / \sqrt{3} \\
1 & 0 & 1 / 3
\end{array}\right|=\frac{1}{3 \sqrt{3}} \tag{6.83}
\end{gather*}
$$

Associated with the inter-element vector, the boundary integrals in Eq. (6.70) and (6.71) are rewritten as

$$
\begin{gather*}
\mathbf{h}^{(1)}=\int_{C_{1-5}^{(1)}}\left[\begin{array}{lll}
N_{1}^{(1)} N_{1}^{(1)} & N_{1}^{(1)} N_{2}^{(1)} & N_{1}^{(1)} N_{3}^{(1)} \\
N_{2}^{(1)} N_{1}^{(1)} & N_{2}^{(1)} N_{2}^{(1)} & N_{2}^{(1)} N_{3}^{(1)} \\
N_{3}^{(1)} N_{1}^{(1)} & N_{3}^{(1)} N_{2}^{(1)} & N_{3}^{(1)} N_{3}^{(1)}
\end{array}\right] d s  \tag{6.84a}\\
\mathbf{g}^{(1)}=\oint\left\{\begin{array}{l}
N_{1}^{(1)} \\
N_{2}^{(1)} \\
N_{3}^{(1)}
\end{array}\right\} d s \tag{6.84b}
\end{gather*}
$$

and

$$
\begin{gather*}
\mathbf{h}^{(2)}=\int_{C_{5-2}^{(2)}}\left[\begin{array}{lll}
N_{1}^{(2)} N_{1}^{(2)} & N_{1}^{(2)} N_{2}^{(2)} & N_{1}^{(2)} N_{3}^{(2)} \\
N_{2}^{(2)} N_{1}^{(2)} & N_{2}^{(2)} N_{2}^{(2)} & N_{2}^{(2)} N_{3}^{(2)} \\
N_{3}^{(2)} N_{1}^{(2)} & N_{3}^{(2)} N_{2}^{(2)} & N_{3}^{(2)} N_{3}^{(2)}
\end{array}\right] d s  \tag{6.85a}\\
\mathbf{g}^{(2)}=\oint\left\{\begin{array}{l}
N_{1}^{(2)} \\
N_{2}^{(2)} \\
N_{3}^{(2)}
\end{array}\right\} d s \tag{6.85b}
\end{gather*}
$$

in which $N_{3}^{(1)}$ and $N_{3}^{(2)}$ are zero along side 1-5 (with length $L_{1-5}$ ) and along side 5-2 (with length $L_{2-5}$ ), respectively. The remaining shape functions $N_{1}^{(1)}, N_{2}^{(1)}, N_{1}^{(2)}$, and $N_{2}^{(2)}$ reduce to a one-dimensional form as

$$
\begin{align*}
& N_{1}^{(1)}=\frac{L_{1-5}-s}{L_{1-5}} \quad \text { and } \quad N_{2}^{(1)}=\frac{s}{L_{1-5}}  \tag{6.86}\\
& N_{1}^{(2)}=\frac{L_{5-2}-s}{L_{5-2}} \quad \text { and } \quad N_{2}^{(2)}=\frac{s}{L_{5-2}} \tag{6.87}
\end{align*}
$$

in which $s$ is the local coordinate in the range of ( $0 \leq s \leq L_{1-5}$ ) along side 15 and ( $0 \leq s \leq L_{5-2}$ ) along side $5-2, L_{1-5}=1 / \sqrt{3}$, and $L_{5-2}=1 / \sqrt{3}$. With these shape functions, the evaluation of $\mathbf{h}^{(1)}, \mathbf{g}^{(1)}, \mathbf{h}^{(2)}$, and $\mathbf{g}^{(2)}$ leads to

$$
\mathbf{h}^{(1)}=\frac{1}{6 \sqrt{3}}\left[\begin{array}{lll}
2 & 1 & 0  \tag{6.88}\\
1 & 2 & 0 \\
0 & 0 & 0
\end{array}\right] \text { and } \mathbf{g}^{(1)}=\frac{1}{2 \sqrt{3}}\left\{\begin{array}{l}
1 \\
1 \\
0
\end{array}\right\}
$$

and

$$
\mathbf{h}^{(2)}=\frac{1}{6 \sqrt{3}}\left[\begin{array}{lll}
2 & 1 & 0  \tag{6.89}\\
1 & 2 & 0 \\
0 & 0 & 0
\end{array}\right] \quad \text { and } \quad \mathbf{g}^{(2)}=\frac{1}{2 \sqrt{3}}\left\{\begin{array}{l}
1 \\
1 \\
0
\end{array}\right\}
$$

Considering the correspondence between the local and global node numbering presented in Table 6.3, the element characteristic matrices and vectors can be rewritten as

Element 1 :
[1]
5
4

$$
\left[\begin{array}{ccc}
k_{11}^{(1)}+h_{11}^{(1)} & k_{12}^{(1)}+h_{12}^{(1)} & k_{13}^{(1)}+h_{13}^{(1)}  \tag{6.90}\\
k_{21}^{(1)}+h_{21}^{(1)} & k_{22}^{(1)}+h_{22}^{(1)} & k_{23}^{(1)}+h_{23}^{(1)} \\
k_{31}^{(1)}+h_{31}^{(1)} & k_{32}^{(1)}+h_{32}^{(1)} & k_{33}^{(1)}+h_{33}^{(1)}
\end{array}\right]\left\{\begin{array}{c}
\phi_{1}^{(1)} \\
\phi_{2}^{(1)} \\
\phi_{3}^{(1)}
\end{array}\right\}=\left\{\begin{array}{c}
f_{1}^{(1)}+g_{1}^{(1)} \\
f_{2}^{(1)}+g_{2}^{(1)} \\
f_{3}^{(1)}+g_{3}^{(1)}
\end{array}\right\}\left[\begin{array}{l}
1 \\
\hline
\end{array}\right.
$$

Element 2:
[5 $\quad 2$ [

$$
\left[\begin{array}{lll}
k_{11}^{(2)}+h_{11}^{(2)} & k_{12}^{(2)}+h_{12}^{(2)} & k_{13}^{(2)}+h_{13}^{(2)}  \tag{6.91}\\
k_{21}^{(2)}+h_{21}^{(2)} & k_{22}^{(2)}+h_{22}^{(2)} & k_{23}^{(2)}+h_{23}^{(2)} \\
k_{31}^{(2)}+h_{31}^{(2)} & k_{32}^{(2)}+h_{32}^{(2)} & k_{33}^{(2)}+h_{33}^{(2)}
\end{array}\right]\left\{\begin{array}{l}
\phi_{1}^{(2)} \\
\phi_{2}^{(2)} \\
\phi_{3}^{(2)}
\end{array}\right\}=\left\{\begin{array}{l}
f_{1}^{(2)}+g_{1}^{(2)} \\
f_{2}^{(2)}+g_{2}^{(2)} \\
f_{3}^{(2)}+g_{3}^{(2)}
\end{array}\right\}
$$

Table 6.3 Element connectivity.

| Element <br> Number <br> $(e)$ | Node 1 | Node 2 | Node 3 |
| :---: | :---: | :---: | :---: |
| 1 | 1 | 5 | 4 |
| 2 | 5 | 2 | 4 |
| 3 | 2 | 3 | 4 |
| 4 | 3 | 1 | 4 |

Element 3:

$$
\begin{gather*}
2 \\
{\left[\begin{array}{lll}
k_{11}^{(3)} & k_{12}^{(3)} & k_{13}^{(3)} \\
k_{21}^{(3)} & k_{22}^{(3)} & k_{23}^{(3)} \\
k_{31}^{(3)} & k_{32}^{(3)} & k_{33}^{(3)}
\end{array}\right]\left\{\begin{array}{l}
\phi_{1}^{(3)} \\
\phi_{2}^{(3)} \\
\phi_{3}^{(3)}
\end{array}\right\}=\left\{\begin{array}{l}
f_{1}^{(3)} \\
f_{2}^{(3)} \\
f_{3}^{(3)}
\end{array}\right\} \begin{array}{l}
2 \\
3 \\
4
\end{array}} \tag{6.92}
\end{gather*}
$$

Element 4:

$$
\begin{gather*}
3 \\
{\left[\begin{array}{lll}
k_{11}^{(4)} & k_{12}^{(4)} & k_{13}^{(4)} \\
k_{21}^{(4)} & k_{22}^{(4)} & k_{23}^{(4)} \\
k_{31}^{(4)} & k_{32}^{(4)} & k_{33}^{(4)}
\end{array}\right]\left\{\begin{array}{l}
\phi_{1}^{(4)} \\
\phi_{2}^{(4)} \\
\phi_{3}^{(4)}
\end{array}\right\}=\left\{\begin{array}{l}
f_{1}^{(4)} \\
f_{2}^{(4)} \\
f_{3}^{(4)}
\end{array}\right\}\left[\begin{array}{|c}
3 \\
1
\end{array}\right]} \tag{6.93}
\end{gather*}
$$

In the assembly of the element characteristic matrices and vectors, the boxed numbers indicate the rows and columns of the global matrix, $\mathbf{K}$, and global right-hand-side vector, $\mathbf{F}$, to which the individual coefficients are added, resulting in

$$
\begin{equation*}
\mathbf{K \Phi}=\mathbf{F} \tag{6.94}
\end{equation*}
$$

where

$$
\begin{align*}
& \mathbf{K}=\left[\begin{array}{ccc}
k_{11}^{(1)}+h_{11}^{(1)}+k_{22}^{(4)} & 0 & k_{21}^{(4)} \\
0 & k_{22}^{(2)}+h_{22}^{(2)}+k_{11}^{(3)} & k_{12}^{(3)} \\
k_{12}^{(4)} & k_{21}^{(3)} & k_{22}^{(3)}+k_{11}^{(4)} \\
k_{31}^{(1)}+h_{31}^{(1)}+k_{32}^{(4)} & k_{32}^{(2)}+h_{32}^{(2)}+k_{31}^{(3)} & k_{32}^{(3)}+k_{31}^{(4)} \\
k_{21}^{(1)}+h_{21}^{(1)} & k_{12}^{(2)}+h_{12}^{(2)} & 0
\end{array}\right.  \tag{6.95a}\\
& \begin{array}{l}
k_{13}^{(1)}+h_{13}^{(1)}+k_{23}^{(4)} \\
k_{23}^{(2)}+h_{23}^{(2)}+k_{13}^{(3)}
\end{array} \\
& k_{23}^{(2)}+h_{23}^{(2)}+k_{13}^{(3)} \\
& k_{23}^{(3)}+k_{13}^{(4)} \\
& \left.\begin{array}{cl}
k_{33}^{(1)}+h_{33}^{(1)}+k_{33}^{(2)}+h_{33}^{(2)}+k_{33}^{(3)}+k_{33}^{(4)} & k_{32}^{(1)}+h_{32}^{(1)}+k_{31}^{(2)}+h_{31}^{(2)} \\
k_{23}^{(1)}+h_{23}^{(1)}+k_{13}^{(2)}+h_{13}^{(2)} & k_{22}^{(1)}+h_{22}^{(1)}+k_{11}^{(2)}+h_{11}^{(2)}
\end{array}\right] \\
& \mathbf{F}=\left\{\begin{array}{c}
f_{1}^{(1)}+g_{1}^{(1)}+f_{2}^{(4)} \\
f_{2}^{(2)}+g_{2}^{(2)}+f_{1}^{(3)} \\
f_{2}^{(3)}+f_{1}^{(4)} \\
f_{3}^{(1)}+g_{3}^{(1)}+f_{3}^{(2)}+g_{3}^{(2)}+f_{3}^{(3)}+f_{3}^{(4)} \\
f_{2}^{(1)}+g_{2}^{(1)}+f_{1}^{(2)}+g_{1}^{(2)}
\end{array}\right\}  \tag{6.95b}\\
& \Phi=\left\{\begin{array}{l}
\phi_{1} \\
\phi_{2} \\
\phi_{3} \\
\phi_{4} \\
\phi_{5}
\end{array}\right\}  \tag{6.95c}\\
& k_{12}^{(1)}+h_{12}^{(1)} \\
& k_{21}^{(2)}+h_{21}^{(2)} \\
& 0
\end{align*}
$$

After imposing the essential boundary conditions, the global system of equations are reduced by deleting the rows and columns corresponding to $\phi_{1}, \phi_{2}$, and $\phi_{3}$, leading to

$$
\begin{align*}
& {\left[\begin{array}{c}
k_{33}^{(1)}+h_{33}^{(1)}+k_{33}^{(2)}+h_{33}^{(2)}+k_{33}^{(3)}+k_{33}^{(4)} \\
k_{32}^{(1)}+h_{32}^{(1)}+k_{31}^{(2)}+h_{31}^{(2)} \\
k_{23}^{(1)}+h_{23}^{(1)}+k_{13}^{(2)}-h_{13}^{(2)} \\
=\left\{\begin{array}{c}
(1) \\
2\left(h_{22}^{(1)}+k_{11}^{(2)}+h_{11}^{(2)}\right.
\end{array}\right]\left\{\begin{array}{c}
\phi_{4} \\
\phi_{5}
\end{array}\right\} \\
f_{2}^{(1)}+g_{2}^{(1)}+f_{1}^{(2)}+g_{1}^{(2)}
\end{array}\right\}}
\end{align*}
$$

With the explicit values of the coefficients, the nodal unknowns, $\phi_{4}$ and $\phi_{5}$, are determined as

$$
\begin{gather*}
\phi_{4}=\frac{4}{27}=0.14815  \tag{6.97a}\\
\phi_{5}=\frac{1}{3}=0.33333 \tag{6.97b}
\end{gather*}
$$

The expressions for $\mathbf{h}^{(1)}$ and $\mathbf{h}^{(2)}$ in Eq. (6.70) are derived based on a formulation consistent with the derivation of the element coefficient matrices, $\mathbf{k}^{(e)}$. An alternative to the consistent formulation is the use of lumped diagonal matrices and expressing $\mathbf{h}^{(1)}$ and $\mathbf{h}^{(2)}$ in the form

$$
\mathbf{h}^{(1)}=\oint_{C_{1-5}^{(1)}}\left[\begin{array}{ccc}
N_{1}^{(1)} & 0 & 0  \tag{6.98}\\
0 & N_{2}^{(1)} & 0 \\
0 & 0 & N_{3}^{(1)}
\end{array}\right] d s=\frac{1}{6 \sqrt{3}}\left[\begin{array}{lll}
3 & 0 & 0 \\
0 & 3 & 0 \\
0 & 0 & 0
\end{array}\right]
$$

and

$$
\mathbf{h}^{(2)}=\oint_{C_{j-2}^{(2)}}\left[\begin{array}{ccc}
N_{1}^{(2)} & 0 & 0  \tag{6.99}\\
0 & N_{2}^{(2)} & 0 \\
0 & 0 & N_{3}^{(2)}
\end{array}\right] d s=\frac{1}{6 \sqrt{3}}\left[\begin{array}{lll}
3 & 0 & 0 \\
0 & 3 & 0 \\
0 & 0 & 0
\end{array}\right]
$$

Replacing the components of $\mathbf{h}^{(1)}$ and $\mathbf{h}^{(2)}$ in Eq. (6.96) with the values obtained in Eq. (6.98) and (6.99), the nodal unknowns $\phi_{4}$ and $\phi_{5}$ are determined as

$$
\begin{align*}
& \phi_{4}=\frac{5}{36}=0.13889  \tag{6.100a}\\
& \phi_{5}=\frac{11}{36}=0.30556 \tag{6.100b}
\end{align*}
$$

Note that the discrepancy in the value of $\phi_{4}$ and $\phi_{5}$ obtained from the two methods is due to the small number of elements in the discretization of the domain.

### 6.1.2.2 ANSYS Solution

The governing equations for a steady-state heat transfer, described by Eq. (6.47) through (6.50), also can be solved using ANSYS. The solution procedure is outlined as follows:

## MODEL GENERATION

- Specify the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on $A d d$.
- Select Thermal Solid from the left list and Quad 4node 55 from the right list; click on $\boldsymbol{O K}$.
- Click on Close .

■ Specify material properties (MP command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Material Props $>$ Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Thermal, Conductivity, and, finally, Isotropic, which brings up another dialog box.
- Enter $\boldsymbol{I}$ for $\boldsymbol{K X X}$, and click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create nodes ( $\mathbf{N}$ command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Nodes $>\operatorname{In}$ Active CS

- A total of 5 nodes are created (Table 6.2).
- Referring to Table 6.2, enter $x$ - and $y$-coordinates of node 1 , and click on Apply. This action keeps the Create Nodes in Active Coordinate System dialog box open. If the Node number field is left blank, then ANSYS assigns the lowest available node number to the node that is being created.
- Repeat the same procedure for the nodes 2 through 5 .
- After entering the $x$ - and $y$-coordinates of node 5 , click on $O K$ (instead of Apply).
- The nodes should appear in the Graphics Window, as shown in Fig. 6.7.


Fig. 6.7 Generation of nodes.

- Create elements ( $\mathbf{E}$ command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Elements > Auto Numbered > Thru Nodes

- Pick Menu appears; refer to Fig. 6.8 to create elements by picking three nodes at a time and clicking on Apply in between.
- Observe the elements created after clicking on Apply in the Pick Menu.
- Repeat until the last element is created.
- Click on $O K$ when the last element is created.
- Review elements:
- Turn on element numbering using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Select Element numbers from the first pull-down menu; click on $O K$.
- Plot elements (EPLOT command) using the following menu path:

Utility Menu > Plot > Elements

- Figure 6.8 shows the outcome of this action as it appears in the Graphics Window.
- Turn off element numbering and turn on node numbering using the following menu path:

Utility Menu $>$ PlotCtrls $>$ Numbering

- Place a checkmark by clicking on the empty box next to NODE Node numbers.


Fig. 6.8 Generation of elements.

- Select No numbering from the first pull-down menu.
- Click on $\boldsymbol{O K}$.
- Plot nodes (NPLOT command) using the following menu path: Utility Menu > Plot > Nodes
- Figure 6.7 shows the outcome of this action as it appears in the Graphics Window.


## SOLUTION

- Apply temperature boundary conditions (D command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Thermal > Temperature > On Nodes

- Pick Menu appears; pick nodes 1, 2, and 3 (Fig. 6.7); click on $\boldsymbol{O K}$ on Pick Menu.
- Highlight TEMP and enter $\mathbf{0}$ for VALUE; click on $\boldsymbol{O K}$ (Fig. 6.9).
- Apply convection boundary conditions (SF command) using the following menu path:


## Main Menu > Solution > Define Loads > Apply > Thermal > Convection > On Nodes

- Pick Menu appears; pick nodes 1, 2 and 5 along the boundary (Fig. 6.7); click on OK on Pick Menu.
- Enter 1 for both VALI Film coefficient and VAL2I Bulk temperature; click on $O K$ (Fig. 6.10).
- Apply body load on elements (BFE command) using the following menu path:

[^7]

Fig. 6.9 Application of temperature boundary conditions on nodes.

| A Apply CONV on nodes |  |  |
| :--- | :--- | :--- |
| [5F] Apply Film Coef on nodes | Constant value |  |
| If Constant value then: |  |  |
| vali Film coefficient |  |  |

Fig. 6.10 Application of convection boundary conditions on nodes.

- Pick Menu appears; click on Pick All.
- Enter $\mathbf{l}$ for $\boldsymbol{V A L 1}$ leave other fields untouched, as shown in Fig. 6.11.
- Click on $\boldsymbol{O K}$.
- Obtain solution (SOLVE command) using the following menu path:

Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status. If OK, close the Status Report Window and click on OK in Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review temperature values (PRNSOL command) using the following menu path:

Main Menu > General Postproc > List Results > Nodal Solution

- Highlight DOF solution on the left list and Temperature TEMP on the right list; click on $\boldsymbol{O K}$.
- The list appears in a new window, as shown in Fig. 6.12.

| A. Apply HGEN on elems |  |
| :--- | :--- |
| [BFE] Apply HGEN on elems as a | Constant value |
| If Constant value then: |  |
| STLOC Starting location N |  |
| VAL1 Load HGEN at loc N |  |
| VAL2 Load HGEN at loc $\mathrm{N}+1$ |  |
| VAL3 Load HGEN at loc $\mathrm{N}+2$ |  |
| VAL4 Load HGEN at loc $\mathrm{N}+3$ |  |

Fig. 6.11 Application of heat generation condition on elements.


Fig. 6.12 Nodal solution for temperature.

### 6.1.3 Example: Two-dimensional Differential Equation with Linear Quadrilateral Elements

### 6.1.3.1 Galerkin's Method

In solving two-dimensional problems with quadrilateral isoparametric elements, Galerkin's method is demonstrated by considering the partial differential equation given by

$$
\begin{equation*}
\frac{\partial^{2} \phi(x, y)}{\partial x^{2}}+\frac{\partial^{2} \phi(x, y)}{\partial y^{2}}-A=0 \tag{6.101}
\end{equation*}
$$

in domain D defined by the intersection of $y=-3, x=-4, y=3$, and $y=3 x-15$. The constant, $A$, is known. As shown in Fig. 6.13, the flux vanishes along the boundary of the domain specified by $y=-3$ and $x=-4$, and along the remaining part of the boundary specified by $y=3$ and $y=3 x-15$, the dependent variable, $\phi(x, y)$, has a value of unity. These boundary conditions are expressed as

$$
\begin{gather*}
\phi(x, y)=1 \text { for } 4 \leq x \leq 6, y=3 x-15  \tag{6.102}\\
\frac{\partial}{\partial x} \phi(x, y=-3)=0 \text { for }-4 \leq x \leq 4  \tag{6.103}\\
\frac{\partial}{\partial x} \phi(x=-4, y)=0 \text { for }-3 \leq y \leq 3  \tag{6.104}\\
\phi(x, y=3)=1 \text { for }-4 \leq x \leq 6 \tag{6.105}
\end{gather*}
$$



Fig. 6.13 Description of domain, and boundary conditions.
The domain is discretized with four linear quadrilateral isoparametric elements, each having four nodes identified as 1, 2, 3, and 4, shown in Fig. 6.14. The nodal values of the dependent variable associated with element $e$ are specified at its first, second, third, and fourth nodes by $\phi_{1}^{(e)}, \phi_{2}^{(e)}, \phi_{3}^{(e)}$, and $\phi_{4}^{(e)}$, respectively. The discretization of the domain with global node numbering is shown in Fig. 6.14. The global coordinates of the nodal values of the dependent variable denoted by $\phi_{i}(i=1,2, \ldots, 9)$ are presented in Table 6.4.

The linear element approximation function for the dependent field variable in a quadrilateral isoparametric element " $e$ " is written as

$$
\begin{equation*}
\tilde{\phi}^{(e)}=N_{1}^{(e)} \phi_{1}^{(e)}+N_{2}^{(e)} \phi_{2}^{(e)}+N_{3}^{(e)} \phi_{3}^{(e)}+N_{4}^{(e)} \phi_{4}^{(e)} \tag{6.106}
\end{equation*}
$$

or


Fig. 6.14 FEM discretization of the domain into four quadrilaterals.

Table 6.4 Nodal coordinates.

| Global <br> Node Number | Nodal <br> Coordinates | Nodal <br> Variables |
| :---: | :---: | :---: |
| 1 | $(-4,-3)$ | $\phi_{1}$ |
| 2 | $(0,-3)$ | $\phi_{2}$ |
| 3 | $(4,-3)$ | $\phi_{3}$ |
| 4 | $(5,0)$ | $\phi_{4}$ |
| 5 | $(6,3)$ | $\phi_{5}$ |
| 6 | $(0,3)$ | $\phi_{6}$ |
| 7 | $(-4,3)$ | $\phi_{7}$ |
| 8 | $(-4,0)$ | $\phi_{8}$ |
| 9 | $(0,0)$ | $\phi_{9}$ |

$$
\begin{equation*}
\tilde{\phi}^{(e)}=\mathbf{N}^{(e) T} \boldsymbol{\varphi}^{(e)} \tag{6.107}
\end{equation*}
$$

where

$$
\mathbf{N}^{(e)}=\left\{\begin{array}{l}
N_{1}^{(e)}  \tag{6.108}\\
N_{2}^{(e)} \\
N_{3}^{(e)} \\
N_{4}^{(e)}
\end{array}\right\} \quad \text { and } \quad \varphi^{(e)}=\left\{\begin{array}{c}
\phi_{1}^{(e)} \\
\phi_{2}^{(e)} \\
\phi_{3}^{(e)} \\
\phi_{4}^{(e)}
\end{array}\right\}
$$

in which the shape functions $N_{1}^{(e)}, N_{2}^{(e)}, N_{3}^{(e)}$, and $N_{4}^{(e)}$ are expressed in terms of the centroidal or natural coordinates, $(\xi, \eta)$, shown in Fig. 6.15. For a linear (straight-sided) quadrilateral illustrated in Fig. 6.15, they are of the form

$$
\begin{equation*}
N_{i}^{(e)}=\frac{1}{4}\left(1+\xi \xi_{i}\right)\left(1+\eta \eta_{i}\right) \quad \text { with } \quad i=1,2,3,4 \tag{6.109}
\end{equation*}
$$

where $\xi_{i}$ and $\eta_{i}$ represent the coordinates of the corner nodes in the natural coordinate system, $\left(\xi_{1}=-1, \eta_{1}=-1\right)$, $\left(\xi_{2}=1, \eta_{2}=-1\right),\left(\xi_{3}=1, \eta_{3}=1\right)$, and $\left(\xi_{4}=-1, \eta_{4}=1\right)$.

Applying Eq. (6.6), Galerkin's method, leads to

$$
\begin{equation*}
\sum_{e=1}^{E} \int_{D^{(e)}} \mathbf{N}^{(e)}\left(\frac{\partial^{2} \tilde{\phi}^{(e)}(x, y)}{\partial x^{2}}+\frac{\partial^{2} \tilde{\phi}^{(e)}(x, y)}{\partial y^{2}}-A\right) d x d y=0 \tag{6.110}
\end{equation*}
$$



Fig. 6.15 Local node numbering for a linear isoparametric quadrilateral element.

Since the element approximation function is $C^{0}$ continuous, the secondorder derivatives in the integrand must be reduced by one so that interelement continuity is achieved during the assembly of the global matrix. This reduction is achieved by observing that

$$
\begin{equation*}
\mathbf{N}^{(e)} \frac{\partial^{2} \tilde{\phi}^{(e)}}{\partial x^{2}}(x, y)=\frac{\partial}{\partial x}\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}(x, y)\right)-\frac{\partial \mathbf{N}^{(e)}}{\partial x} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}(x, y) \tag{6.111}
\end{equation*}
$$

and

$$
\begin{equation*}
\mathbf{N}^{(e)} \frac{\partial^{2} \tilde{\phi}^{(e)}}{\partial y^{2}}(x, y)=\frac{\partial}{\partial y}\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}(x, y)\right)-\frac{\partial \mathbf{N}^{(e)}}{\partial y} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}(x, y) \tag{6.112}
\end{equation*}
$$

Their substitution into the integrand in Eq. (6.110) and rearrangement of the terms result in

$$
\begin{align*}
& \sum_{e=1}^{E}\left\{\int_{D^{(e)}}\left[\frac{\partial}{\partial x}\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}\right)+\frac{\partial}{\partial y}\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}\right)\right] d x d y\right. \\
& \left.\quad+\int_{D^{(e)}}\left[-\frac{\partial \mathbf{N}^{(e)}}{\partial x} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}-\frac{\partial \mathbf{N}^{(e)}}{\partial y} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}-\mathbf{N}^{(e)} A\right] d x d y\right\}=0 \tag{6.113}
\end{align*}
$$

Applying the divergence theorem to the first integral renders the domain integral to the boundary integral, and it yields

$$
\begin{align*}
& \sum_{e=1}^{E}\left\{\oint_{C^{(e)}}\left[\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}\right) n_{x}^{(e)}+\left(\mathbf{N}^{(e)} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}\right) n_{y}^{(e)}\right] d s\right. \\
&\left.+\int_{D^{(e)}}\left[-\frac{\partial \mathbf{N}^{(e)}}{\partial x} \frac{\partial \tilde{\phi}^{(e)}}{\partial x}-\frac{\partial \mathbf{N}^{(e)}}{\partial y} \frac{\partial \tilde{\phi}^{(e)}}{\partial y}-\mathbf{N}^{(e)} A\right] d x d y\right\}=0 \tag{6.114}
\end{align*}
$$

where $n_{x}^{(e)}$ and $n_{y}^{(e)}$ are, respectively, the $x$ - and $y$-components of the outward normal vector along the closed boundary defining the area of the element $C^{(e)}$.

Substituting for the element approximation function yields

$$
\begin{align*}
& \sum_{e=1}^{E}\left\{\oint_{C^{(e)}} \mathbf{N}^{(e)}\left[\frac{\partial \tilde{\phi}^{(e)}}{\partial x} n_{x}^{(e)}+\frac{\partial \tilde{\phi}^{(e)}}{\partial y} n_{y}^{(e)}\right] d s\right. \\
&+\int_{D^{(e)}}\left[-\frac{\partial \mathbf{N}^{(e)}}{\partial x} \frac{\partial \mathbf{N}^{(e) T}}{\partial x}-\frac{\partial \mathbf{N}^{(e)}}{\partial y} \frac{\partial \mathbf{N}^{(e) T}}{\partial y}\right] d x d y \varphi^{(e)}  \tag{6.115}\\
&\left.-\int_{D^{(e)}} \mathbf{N}^{(e)} A d x d y\right\}=0
\end{align*}
$$

This equation can be recast in matrix form as

$$
\begin{equation*}
\sum_{e=1}^{E}\left(\mathbf{k}^{(e)} \boldsymbol{\varphi}^{(e)}-\mathbf{f}^{(e)}+\mathbf{Q}^{(e)}\right)=0 \tag{6.116}
\end{equation*}
$$

where

$$
\begin{gather*}
\mathbf{k}^{(e)}=-\int_{D^{(e)}}\left[\frac{\partial \mathbf{N}^{(e)}}{\partial x} \frac{\partial \mathbf{N}^{(e) T}}{\partial x}+\frac{\partial \mathbf{N}^{(e)}}{\partial y} \frac{\partial \mathbf{N}^{(e) T}}{\partial y}\right] d x d y  \tag{6.117}\\
\mathbf{f}^{(e)}=\int_{D^{(e)}} A \mathbf{N}^{(e)} d x d y \tag{6.118}
\end{gather*}
$$

$$
\begin{equation*}
\mathbf{Q}^{(e)}=\oint_{C^{(e)}} \mathbf{N}^{(e)}\left[\frac{\partial \tilde{\phi}^{(e)}}{\partial x} n_{x}^{(e)}+\frac{\partial \tilde{\phi}^{(e)}}{\partial y} n_{y}^{(e)}\right] d s \tag{6.119}
\end{equation*}
$$

in which $\mathbf{k}^{(e)}$ is the element characteristic matrix, $\mathbf{f}^{(e)}$ is the element right-hand-side vector, and $\mathbf{Q}^{(e)}$ is often referred to as the inter-element vector that includes the derivative terms along the boundary of the element. The boundary integral around each element is evaluated in a counterclockwise direction, i.e., this boundary integral is the sum of four integrals taken along each side of the element.

Because the specified derivatives have zero values along the element boundaries, the inter-element vector, $\mathbf{Q}^{(e)}$ vanishes, i.e., $\mathbf{Q}^{(e)}=0$, thus reducing the element equilibrium equations to

$$
\begin{equation*}
\sum_{e=1}^{E}\left(\mathbf{k}^{(e)} \boldsymbol{\varphi}^{(e)}-\mathbf{f}^{(e)}\right)=0 \tag{6.120}
\end{equation*}
$$

The integrals contributing to the characteristic element matrix, $\mathbf{k}^{(e)}$, and the right-hand-side vector, $\mathbf{f}^{(e)}$, are evaluated over a square region in the natural coordinate system after an appropriate coordinate transformation given by

$$
\begin{equation*}
x=\sum_{i=1}^{4} N_{i}^{(e)}(\xi, \eta) x_{i}^{(e)} \quad \text { and } \quad y=\sum_{i=1}^{4} N_{i}^{(e)}(\xi, \eta) y_{i}^{(e)} \tag{6.121}
\end{equation*}
$$

Application of the chain rule of differentiation yields

$$
\left\{\begin{array}{c}
\frac{\partial N_{i}^{(e)}}{\partial \xi}  \tag{6.122}\\
\frac{\partial N_{i}^{(e)}}{\partial \eta}
\end{array}\right\}=\left[\begin{array}{ll}
\frac{\partial x}{\partial \xi} & \frac{\partial y}{\partial \xi} \\
\frac{\partial x}{\partial \eta} & \frac{\partial y}{\partial \eta}
\end{array}\right]\left\{\begin{array}{c}
\frac{\partial N_{i}^{(e)}}{\partial x} \\
\frac{\partial N_{i}^{(e)}}{\partial y}
\end{array}\right\} \text { with } i=1,2,3,4
$$

or

$$
\left\{\begin{array}{c}
\frac{\partial}{\partial \xi}  \tag{6.123}\\
\frac{\partial}{\partial \eta}
\end{array}\right\} N_{i}^{(e)}=\mathbf{J}\left\{\begin{array}{c}
\frac{\partial}{\partial x} \\
\frac{\partial}{\partial y}
\end{array}\right\} N_{i}^{(e)}
$$

where $\mathbf{J}$ is called the Jacobian matrix. It can be expressed as

$$
\mathbf{J}=\left[\begin{array}{ll}
J_{11} & J_{12}  \tag{6.124}\\
J_{21} & J_{22}
\end{array}\right]
$$

in which

$$
\begin{align*}
& J_{11}=\frac{\partial x}{\partial \xi}=\frac{1}{4}\left\{-(1-\eta) x_{1}^{(e)}+(1-\eta) x_{2}^{(e)}+(1+\eta) x_{3}^{(e)}-(1+\eta) x_{4}^{(e)}\right\}  \tag{6.125}\\
& J_{12}=\frac{\partial y}{\partial \xi}=\frac{1}{4}\left\{-(1-\eta) y_{1}^{(e)}+(1-\eta) y_{2}^{(e)}+(1+\eta) y_{3}^{(e)}-(1+\eta) y_{4}^{(e)}\right\}  \tag{6.126}\\
& J_{21}=\frac{\partial x}{\partial \eta}=\frac{1}{4}\left\{-(1-\xi) x_{1}^{(e)}-(1+\xi) x_{2}^{(e)}+(1+\xi) x_{3}^{(e)}+(1-\xi) x_{4}^{(e)}\right\}  \tag{6.127}\\
& J_{22}=\frac{\partial y}{\partial \eta}=\frac{1}{4}\left\{-(1-\xi) y_{1}^{(e)}-(1+\xi) y_{2}^{(e)}+(1+\xi) y_{3}^{(e)}+(1-\xi) y_{4}^{(e)}\right\} \tag{6.128}
\end{align*}
$$

Also, the Jacobian can be rewritten in the form

$$
\mathbf{J}=\left[\begin{array}{cccc}
\frac{\partial N_{1}^{(e)}}{\partial \xi} & \frac{\partial N_{2}^{(e)}}{\partial \xi} & \frac{\partial N_{3}^{(e)}}{\partial \xi} & \frac{\partial N_{4}^{(e)}}{\partial \xi}  \tag{6.129}\\
\frac{\partial N_{1}^{(e)}}{\partial \eta} & \frac{\partial N_{2}^{(e)}}{\partial \eta} & \frac{\partial N_{3}^{(e)}}{\partial \eta} & \frac{\partial N_{4}^{(e)}}{\partial \eta}
\end{array}\right]\left[\begin{array}{cc}
x_{1}^{(e)} & y_{1}^{(e)} \\
x_{2}^{(e)} & y_{2}^{(e)} \\
x_{3}^{(e)} & y_{3}^{(e)} \\
x_{4}^{(e)} & y_{4}^{(e)}
\end{array}\right]
$$

or

$$
\mathbf{J}=\frac{1}{4}\left[\begin{array}{cccc}
-(1-\eta) & (1-\eta) & (1+\eta) & -(1+\eta)  \tag{6.130}\\
-(1-\xi) & -(1+\xi) & (1+\xi) & (1-\xi)
\end{array}\right]\left[\begin{array}{cc}
x_{1}^{(e)} & y_{1}^{(e)} \\
x_{2}^{(e)} & y_{2}^{(e)} \\
x_{3}^{(e)} & y_{3}^{(e)} \\
x_{4}^{(e)} & y_{4}^{(e)}
\end{array}\right]
$$

Because the transformation between the natural and global coordinates has a one-to-one correspondence, the inverse of the Jacobian exists, and it can be expressed as

$$
\mathbf{J}^{-1}=\frac{1}{|\mathbf{J}|}\left[\begin{array}{cc}
J_{22} & -J_{12}  \tag{6.131}\\
-J_{21} & J_{11}
\end{array}\right]
$$

When the element is degenerated into a triangle by increasing an internal angle to $180^{\circ}, \mathbf{J}$ is singular at that corner. The inverse of the Jacobian matrix permits the expression for the derivatives in terms of global coordinates

$$
\left\{\begin{array}{l}
\frac{\partial N_{i}^{(e)}}{\partial x}  \tag{6.132}\\
\frac{\partial N_{i}^{(e)}}{\partial y}
\end{array}\right\}=\mathbf{J}^{-1}\left\{\begin{array}{c}
\frac{\partial N_{i}^{(e)}}{\partial \xi} \\
\frac{\partial N_{i}^{(e)}}{\partial \eta}
\end{array}\right\}
$$

Defining the element shape matrix $\mathbf{B}^{(e)}$ as

$$
\mathbf{B}^{(e)}=\left[\begin{array}{llll}
\frac{\partial N_{1}^{(e)}}{\partial x} & \frac{\partial N_{2}^{(e)}}{\partial x} & \frac{\partial N_{3}^{(e)}}{\partial x} & \frac{\partial N_{4}^{(e)}}{\partial x}  \tag{6.133}\\
\frac{\partial N_{1}^{(e)}}{\partial y} & \frac{\partial N_{2}^{(e)}}{\partial y} & \frac{\partial N_{3}^{(e)}}{\partial y} & \frac{\partial N_{4}^{(e)}}{\partial y}
\end{array}\right]=\left\{\begin{array}{c}
\frac{\partial}{\partial x} \\
\frac{\partial}{\partial y}
\end{array}\right\} \mathbf{N}^{(e)^{r}}
$$

permits the element matrix $\mathbf{k}^{(e)}$ be written in the form

$$
\begin{equation*}
\mathbf{k}^{(e)}=-\int_{D^{(e)}} \mathbf{B}^{(e) T} \mathbf{B}^{(e)} d x d y=-\int_{-1}^{1} \int_{-1}^{1} \mathbf{B}^{(e) T} \mathbf{B}^{(e)}|\mathbf{J}| d \xi d \eta \tag{6.134}
\end{equation*}
$$

A similar operation is performed for evaluation of $\mathbf{f}^{(e)}$

$$
\begin{equation*}
\mathbf{f}^{(e)}=A \int_{D^{(e)}} \mathbf{N}^{(e)} d x d y=A \int_{-1}^{1} \int_{-1}^{1} \mathbf{N}^{(e)}|\mathbf{J}| d \xi d \eta \tag{6.135}
\end{equation*}
$$

Due to the difficulty of obtaining an analytical expression for the determinant and inverse of the Jacobian matrix, these integrals are evaluated numerically by the Gaussian integration technique described in detail in Sec. 3.6.

Prior to the calculation of the element characteristic matrices, their Jacobian matrices are obtained for each element using Eq. (6.130) as

$$
\begin{align*}
\mathbf{J}^{(1)} & =\frac{1}{4}\left[\begin{array}{cccc}
-(1-\eta) & (1-\eta) & (1+\eta) & -(1+\eta) \\
-(1-\xi) & -(1+\xi) & (1+\xi) & (1-\xi)
\end{array}\right]\left[\begin{array}{cc}
-4 & -3 \\
0 & -3 \\
0 & 0 \\
-4 & 0
\end{array}\right]  \tag{6.136}\\
& =\frac{1}{4}\left[\begin{array}{ll}
8 & 0 \\
0 & 6
\end{array}\right] \quad \text { with }\left|\mathbf{J}^{(1)}\right|=3
\end{align*}
$$

$$
\begin{align*}
& \mathbf{J}^{(2)}=\frac{1}{4}\left[\begin{array}{cccc}
-(1-\eta) & (1-\eta) & (1+\eta) & -(1+\eta) \\
-(1-\xi) & -(1+\xi) & (1+\xi) & (1-\xi)
\end{array}\right]\left[\begin{array}{cc}
0 & -3 \\
4 & -3 \\
5 & 0 \\
0 & 0
\end{array}\right]  \tag{6.137}\\
&=\frac{1}{4}\left[\begin{array}{cc}
9+\eta & 0 \\
1+\xi & 6
\end{array}\right] \quad \text { with }\left|\mathbf{J}^{(2)}\right|=\frac{3}{8}(9+\eta) \\
& \begin{aligned}
\mathbf{J}^{(3)} & =\frac{1}{4}\left[\begin{array}{ccc}
-(1-\eta) & (1-\eta) & (1+\eta) \\
-(1-\xi) & -(1+\xi) & (1+\xi) \\
(1-\xi)
\end{array}\right]\left[\begin{array}{cc}
0 & 0 \\
5 & 0 \\
6 & 3 \\
0 & 3
\end{array}\right] \\
& =\frac{1}{4}\left[\begin{array}{ccc}
11+\eta & 0 \\
1+\xi & 6
\end{array}\right] \quad \text { with }\left|\mathbf{J}^{(3)}\right|=\frac{3}{8}(11+\eta) \\
\mathbf{J}^{(4)} & =\frac{1}{4}\left[\begin{array}{ccc}
-(1-\eta) & (1-\eta) & (1+\eta) \\
-(1-\xi) & -(1+\xi) & (1+\xi) \\
(1-\xi)
\end{array}\right]\left[\begin{array}{cc}
-4 & 0 \\
0 & 0 \\
0 & 3 \\
-4 & 3
\end{array}\right] \\
& =\frac{1}{4}\left[\begin{array}{ll}
8 & 0 \\
0 & 6
\end{array}\right] \quad \text { with }\left|\mathbf{J}^{(4)}\right|=3
\end{aligned} \tag{6.138}
\end{align*}
$$

The inverse of the Jacobian matrices are obtained as

$$
\begin{gather*}
{\left[\mathbf{J}^{(1)}\right]^{-1}=4\left[\begin{array}{ll}
\frac{1}{8} & 0 \\
0 & \frac{1}{6}
\end{array}\right]}  \tag{6.140}\\
{\left[\mathbf{J}^{(2)}\right]^{-1}=\frac{4}{6(9+\eta)}\left[\begin{array}{cc}
6 & 0 \\
-(1+\xi) & 9+\eta
\end{array}\right]}  \tag{6.141}\\
{\left[\mathbf{J}^{(3)}\right]^{-1}=\frac{4}{6(11+\eta)}\left[\begin{array}{cc}
6 & 0 \\
-(1+\xi) & 11+\eta
\end{array}\right]}  \tag{6.142}\\
{\left[\mathbf{J}^{(4)}\right]^{-1}=4\left[\begin{array}{cc}
\frac{1}{8} & 0 \\
0 & \frac{1}{6}
\end{array}\right]} \tag{6.143}
\end{gather*}
$$

The element shape matrices $\mathbf{B}^{(e)}$ are obtained as

$$
\begin{gather*}
\mathbf{B}^{(1)}=\left[\begin{array}{cccc}
-\frac{1}{8}(1-\eta) & \frac{1}{8}(1-\eta) & \frac{1}{8}(1+\eta) & -\frac{1}{8}(1+\eta) \\
-\frac{1}{6}(1-\xi) & -\frac{1}{6}(1+\xi) & \frac{1}{6}(1+\xi) & \frac{1}{6}(1-\xi)
\end{array}\right]  \tag{6.144}\\
\mathbf{B}^{(2)}=\frac{1}{9+\eta}\left[\begin{array}{cccc}
-(1-\eta) & (1-\eta) & (1+\eta) & -(1+\eta) \\
-\frac{1}{3}(4-5 \xi+\eta) & -\frac{5}{3}(1+\xi) & \frac{4}{3}(1+\xi) & \frac{1}{3}(5-4 \xi+\eta)
\end{array}\right]  \tag{6.145}\\
\mathbf{B}^{(3)}=\frac{1}{11+\eta}\left[\begin{array}{cccc}
-(1-\eta) & (1-\eta) & (1+\eta) & -(1+\eta) \\
-\frac{1}{3}(5-6 \xi+\eta) & -2(1+\xi) & \frac{5}{3}(1+\xi) & \frac{1}{3}(6-5 \xi+\eta)
\end{array}\right]  \tag{6.146}\\
\mathbf{B}^{(4)}=\left[\begin{array}{cccc}
-\frac{1}{8}(1-\eta) & \frac{1}{8}(1-\eta) & \frac{1}{8}(1+\eta) & -\frac{1}{8}(1+\eta) \\
-\frac{1}{6}(1-\xi) & -\frac{1}{6}(1+\xi) & \frac{1}{6}(1+\xi) & \frac{1}{6}(1-\xi)
\end{array}\right] \tag{6.147}
\end{gather*}
$$

Numerical evaluation of the element characteristic matrices results in

$$
\begin{align*}
& \mathbf{k}^{(1)}=\left[\begin{array}{cccc}
\frac{25}{36} & -\frac{1}{36} & -\frac{25}{72} & -\frac{23}{72} \\
-\frac{1}{36} & \frac{25}{36} & -\frac{23}{72} & -\frac{25}{72} \\
-\frac{25}{72} & -\frac{23}{72} & \frac{25}{36} & -\frac{1}{36} \\
-\frac{23}{72} & -\frac{25}{72} & -\frac{1}{36} & \frac{25}{36}
\end{array}\right]  \tag{6.148}\\
& \mathbf{k}^{(2)}=\left[\begin{array}{cccc}
0.688943 & -0.0222762 & -0.282179 & -0.384488 \\
-0.0222762 & 0.85561 & -0.384488 & -0.448846 \\
-0.282179 & -0.384488 & 0.60759 & 0.0590766 \\
-0.384488 & -0.448846 & 0.0590766 & 0.774257
\end{array}\right]  \tag{6.149}\\
& \mathbf{k}^{(3)}=\left[\begin{array}{cccc}
0.753348 & 0.0799856 & -0.316655 & -0.516679 \\
0.0799856 & 0.920014 & -0.516679 & -0.483321 \\
-0.316655 & -0.516679 & 0.680566 & 0.152768 \\
-0.516679 & -0.483321 & 0.152768 & 0.847232
\end{array}\right] \tag{6.150}
\end{align*}
$$

$$
\mathbf{k}^{(4)}=\left[\begin{array}{cccc}
\frac{25}{36} & -\frac{1}{36} & -\frac{25}{72} & -\frac{23}{72}  \tag{6.151}\\
-\frac{1}{36} & \frac{25}{36} & -\frac{23}{72} & -\frac{25}{72} \\
-\frac{25}{72} & -\frac{23}{72} & \frac{25}{36} & -\frac{1}{36} \\
-\frac{23}{72} & -\frac{25}{72} & -\frac{1}{36} & \frac{25}{36}
\end{array}\right]
$$

Similarly, the right-hand-side vectors are calculated as

$$
\mathbf{f}^{(1)}=A\left\{\begin{array}{l}
3  \tag{6.152}\\
3 \\
3 \\
3
\end{array}\right\}, \mathbf{f}^{(2)}=A\left\{\begin{array}{c}
3.25 \\
3.25 \\
3.5 \\
3.5
\end{array}\right\}, \mathbf{f}^{(3)}=A\left\{\begin{array}{c}
4 \\
4 \\
4.25 \\
4.25
\end{array}\right\}, \mathbf{f}^{(4)}=A\left\{\begin{array}{l}
3 \\
3 \\
3 \\
3
\end{array}\right\}
$$

The element definitions (or connectivity of elements), as shown in Fig. 6.14, are presented in Table 6.5.

Considering the correspondence between the local and global node numbering as shown in Table 6.5, the element equations can be rewritten as

$$
\begin{align*}
& \begin{array}{llll}
1 & 2 & 9 & 8
\end{array} \\
& \text { Element 1: }\left[\begin{array}{llll}
k_{11}^{(1)} & k_{12}^{(1)} & k_{13}^{(1)} & k_{14}^{(1)} \\
k_{21}^{(1)} & k_{22}^{(1)} & k_{23}^{(1)} & k_{24}^{(1)} \\
k_{31}^{(1)} & k_{32}^{(1)} & k_{33}^{(1)} & k_{34}^{(1)} \\
k_{11}^{(1)} & k_{42}^{(1)} & k_{43}^{(1)} & k_{44}^{(1)}
\end{array}\right]\left\{\begin{array}{l}
\phi_{1}^{(1)} \\
\phi_{2}^{(1)} \\
\phi_{3}^{(1)} \\
\phi_{4}^{(1)}
\end{array}\right\}=\left\{\begin{array}{c}
f_{1}^{(1)} \\
f_{2}^{(1)} \\
f_{3}^{(1)} \\
f_{4}^{(1)}
\end{array}\right] \tag{6.153}
\end{align*}
$$

Table 6.5 Element connectivity.

| Element <br> Number <br> $(e)$ | Node 1 | Node 2 | Node 3 | Node 4 |
| :---: | :---: | :---: | :---: | :---: |
| 1 | 1 | 2 | 9 | 8 |
| 2 | 2 | 3 | 4 | 9 |
| 3 | 9 | 4 | 5 | 6 |
| 4 | 8 | 9 | 6 | 7 |

Element 2: $\left[\begin{array}{cccc}2 & 3 & 4 & 0 \\ k_{11}^{(2)} & k_{12}^{(2)} & k_{13}^{(2)} & k_{14}^{(2)} \\ k_{21}^{(2)} & k_{22}^{(2)} & k_{23}^{(2)} & k_{24}^{(2)} \\ k_{31}^{(2)} & k_{32}^{(2)} & k_{33}^{(2)} & k_{34}^{(2)} \\ k_{41}^{(2)} & k_{42}^{(2)} & k_{43}^{(2)} & k_{44}^{(2)}\end{array}\right]\left\{\begin{array}{l}\phi_{1}^{(2)} \\ \phi_{2}^{(2)} \\ \phi_{3}^{(2)} \\ \phi_{4}^{(2)}\end{array}\right\}=\left\{\begin{array}{l}f_{1}^{(2)} \\ f_{2}^{(2)} \\ f_{3}^{(2)} \\ f_{4}^{(2)}\end{array}\right] \begin{aligned} & 2 \\ & 3 \\ & 4 \\ & \square\end{aligned}$

Element 3: $\left[\begin{array}{lllll}k_{11}^{(3)} & k_{12}^{(3)} & k_{13}^{(3)} & k_{14}^{(3)} \\ k_{21}^{(3)} & k_{22}^{(3)} & k_{23}^{(3)} & k_{24}^{(3)} \\ k_{31}^{(3)} & k_{32}^{(3)} & k_{33}^{(3)} & k_{34}^{(3)} \\ k_{41}^{(3)} & k_{42}^{(3)} & k_{43}^{(3)} & k_{44}^{(3)}\end{array}\right]\left[\begin{array}{l}\phi_{1}^{(3)} \\ \phi_{2}^{(3)} \\ \phi_{3}^{(3)} \\ \phi_{4}^{(3)}\end{array}\right\}=\left\{\begin{array}{l}f_{1}^{(3)} \\ f_{2}^{(3)}\end{array}\right\} \begin{aligned} & 9 \\ & f_{3}^{(3)} \\ & f_{4}^{(3)}\end{aligned} \begin{aligned} & {[ } \\ & 5\end{aligned}$
Element 4: $\left[\begin{array}{cccc}8 & 0 & 6 & 7 \\ k_{11}^{(4)} & k_{12}^{(4)} & k_{13}^{(4)} & k_{14}^{(4)} \\ k_{21}^{(4)} & k_{22}^{(4)} & k_{23}^{(4)} & k_{24}^{(4)} \\ k_{31}^{(4)} & k_{32}^{(4)} & k_{33}^{(4)} & k_{34}^{(4)} \\ k_{41}^{(4)} & k_{42}^{(4)} & k_{43}^{(4)} & k_{44}^{(4)}\end{array}\right]\left\{\begin{array}{l}\phi_{1}^{(4)} \\ \phi_{2}^{(4)} \\ \phi_{3}^{(4)} \\ \phi_{4}^{(4)}\end{array}\right\}=\left\{\begin{array}{l}f_{1}^{(4)} \\ f_{2}^{(4)} \\ f_{3}^{(4)} \\ f_{4}^{(4)}\end{array}\right] \begin{aligned} & 8 \\ & 6 \\ & 6\end{aligned}$
In the assembly of the element characteristic matrices and vectors, the boxed numbers indicate the rows and columns of the global matrix, $\mathbf{K}$, and global right-hand-side vector, $\mathbf{F}$, to which the individual coefficients are added, resulting in
$\left[\begin{array}{ccccccc}k_{11}^{(1)} & k_{12}^{(1)} & 0 & 0 & 0 & 0 & 0 \\ k_{21}^{(1)} & k_{22}^{(1)}+k_{11}^{(2)} & k_{12}^{(2)} & k_{13}^{(2)} & 0 & 0 & 0 \\ 0 & k_{21}^{(2)} & k_{22}^{(2)} & k_{23}^{(2)} & 0 & 0 & 0 \\ 0 & k_{31}^{(2)} & k_{32}^{(2)} & k_{33}^{(2)}+k_{22}^{(3)} & k_{23}^{(3)} & k_{24}^{(3)} & 0 \\ 0 & 0 & 0 & k_{32}^{(3)} & k_{33}^{(3)} & k_{34}^{(3)} & 0 \\ 0 & 0 & 0 & k_{42}^{(3)} & k_{43}^{(3)} & k_{44}^{(3)}+k_{33}^{(4)} & k_{34}^{(4)} \\ 0 & 0 & 0 & 0 & 0 & k_{43}^{(4)} & k_{44}^{(4)} \\ k_{41}^{(1)} & k_{42}^{(1)} & 0 & 0 & 0 & k_{13}^{(4)} & k_{14}^{(4)} \\ k_{31}^{(1)} & k_{32}^{(1)}+k_{41}^{(2)} & k_{42}^{(2)} & k_{43}^{(2)}+k_{12}^{(3)} & k_{13}^{(3)} & k_{14}^{(3)}+k_{23}^{(4)} & k_{24}^{(4)}\end{array}\right.$
$\left.\begin{array}{cc}k_{14}^{(1)} & k_{13}^{(1)} \\ k_{24}^{(1)} & k_{23}^{(1)}+k_{14}^{(2)} \\ 0 & k_{24}^{(2)} \\ 0 & k_{34}^{(2)}+k_{21}^{(3)} \\ 0 & k_{31}^{(3)} \\ k_{31}^{(4)} & k_{41}^{(3)}+k_{32}^{(4)} \\ k_{41}^{(4)} & k_{42}^{(4)} \\ k_{44}^{(1)}+k_{11}^{(4)} & k_{43}^{(1)}+k_{12}^{(4)} \\ k_{34}^{(1)}+k_{21}^{(4)} & k_{33}^{(1)}+k_{44}^{(2)}+k_{11}^{(3)}+k_{22}^{(4)}\end{array}\right]\left\{\begin{array}{c}\phi_{1} \\ \phi_{2} \\ \phi_{3} \\ \phi_{4} \\ \phi_{5} \\ \phi_{6} \\ \phi_{7} \\ \phi_{8} \\ \phi_{9}\end{array}\right\}=\left\{\begin{array}{c}f_{1}^{(1)} \\ f_{2}^{(1)}+f_{1}^{(2)} \\ f_{2}^{(2)} \\ f_{3}^{(2)}+f_{2}^{(3)} \\ f_{3}^{(3)} \\ f_{4}^{(3)}+f_{3}^{(4)} \\ f_{4}^{(4)} \\ f_{4}^{(1)}+f_{1}^{(4)} \\ f_{3}^{(1)}+f_{4}^{(2)}+f_{1}^{(3)}+f_{2}^{(4)}\end{array}\right\}$
or

$$
\begin{equation*}
\mathbf{K} \boldsymbol{\Phi}=\mathbf{F} \tag{6.158}
\end{equation*}
$$

the global stiffness matrix and right-hand-side vector are numerically evaluated as
$\mathbf{K}=\left[\begin{array}{ccccc}0.694444 & -0.0277778 & 0 & 0 & 0 \\ -0.0277778 & 1.38339 & -0.0222762 & -0.282179 & 0 \\ 0 & -0.0222762 & 0.85561 & -0.384488 & 0 \\ 0 & -0.282179 & -0.384488 & 1.5276 & -0.516679 \\ 0 & 0 & 0 & -0.516679 & 0.680566 \\ 0 & 0 & 0 & -0.483321 & 0.152768 \\ 0 & 0 & 0 & 0 & 0 \\ -0.319444 & -0.347222 & 0 & 0 & 0 \\ -0.347222 & -0.703932 & -0.448846 & 0.139062 & -0.316655\end{array}\right.$
(6.159)
$\left.\begin{array}{cccc}0 & 0 & -0.319444 & -0.347222 \\ 0 & 0 & -0.347222 & -0.703932 \\ 0 & 0 & 0 & -0.448846 \\ -0.483321 & 0 & 0 & 0.139062 \\ 0.152768 & 0 & 0 & -0.316655 \\ 1.54168 & -0.0277778 & -0.347222 & -0.836123 \\ -0.0277778 & 0.694444 & -0.319444 & -0.347222 \\ -0.347222 & -0.319444 & 1.38889 & -0.0555556 \\ -0.836123 & -0.347222 & -0.0555556 & 2.91649\end{array}\right]$
and

$$
\mathbf{F}=\left\{\begin{array}{c}
3  \tag{6.160}\\
6.25 \\
3.25 \\
7.5 \\
4.25 \\
7.25 \\
3 \\
6 \\
13.5
\end{array}\right\}
$$

After imposing the essential boundary conditions, i.e., $\phi_{3}=\phi_{4}=\phi_{5}=\phi_{6}=$ $\phi_{7}=1$, the global system of equations is reduced by deleting the rows and columns corresponding to $\phi_{3}, \phi_{4}, \phi_{5}, \phi_{6}$, and $\phi_{7}$, leading to

$$
\begin{align*}
& {\left[\begin{array}{cccc}
k_{11}^{(1)} & k_{12}^{(1)} & k_{14}^{(1)} & k_{13}^{(1)} \\
k_{21}^{(1)} & k_{22}^{(1)}+k_{11}^{(2)} & k_{24}^{(1)} & k_{23}^{(1)}+k_{14}^{(2)} \\
k_{41}^{(1)} & k_{42}^{(1)} & k_{44}^{(1)}+k_{11}^{(4)} & k_{43}^{(1)}+k_{12}^{(4)} \\
k_{31}^{(1)} & k_{32}^{(1)}+k_{41}^{(2)} & k_{34}^{(1)}+k_{21}^{(4)} & k_{33}^{(1)}+k_{44}^{(2)}+k_{11}^{(3)}+k_{22}^{(4)}
\end{array}\right]\left\{\begin{array}{l}
\phi_{1} \\
\phi_{2} \\
\phi_{8} \\
\phi_{9}
\end{array}\right\}} \\
& =\left\{\begin{array}{c}
f_{1}^{(1)} \\
\quad f_{2}^{(1)}+f_{1}^{(2)}-k_{12}^{(2)}-k_{13}^{(2)} \\
f_{4}^{(1)}+f_{1}^{(4)}-k_{13}^{(4)}-k_{14}^{(4)} \\
f_{3}^{(1)}+f_{4}^{(2)}+f_{1}^{(3)}+f_{2}^{(4)}-k_{42}^{(2)}-\left(k_{43}^{(2)}+k_{12}^{(3)}\right) \\
-k_{13}^{(3)}-\left(k_{14}^{(3)}+k_{23}^{(4)}\right)-k_{24}^{(4)}
\end{array}\right\} \tag{6.161}
\end{align*}
$$

which is numerically evaluated as

$$
\mathbf{K}=\left[\begin{array}{cccc}
0.694444 & -0.0277778 & -0.319444 & -0.347222  \tag{6.162}\\
-0.0277778 & 1.38339 & -0.347222 & -0.703932 \\
-0.319444 & -0.347222 & 1.38889 & -0.0555556 \\
-0.347222 & -0.703932 & -0.0555556 & 2.91649
\end{array}\right]
$$

and

$$
\mathbf{F}=\left\{\begin{array}{c}
3  \tag{6.163}\\
6.55446 \\
6.66667 \\
15.3098
\end{array}\right\}
$$

Finally, the solution of the reduced global system yields

$$
\left\{\begin{array}{l}
\phi_{1}  \tag{6.164}\\
\phi_{2} \\
\phi_{8} \\
\phi_{9}
\end{array}\right\}=\left\{\begin{array}{l}
15.8119 \\
13.5401 \\
12.2471 \\
10.6332
\end{array}\right\}
$$

### 6.1.3.2 ANSYS Solution

The governing equations for a steady-state heat transfer, described by Eq. ( 6.101 ) through (6.105), also can be solved using ANSYS. The solution procedure is outlined as follows:

## MODEL GENERATION

- Specify the element type (ET command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Element Type $>$ Add/Edit/Delete

- Click on Add.
- Select Thermal Solid from the left list and Quad 4node 55 from the right list; click on OK.
- Click on Close.
- Specify material properties (MP command) using the following menu path:

Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Thermal, Conductivity, and, finally, Isotropic, which brings up another dialog box.
- Enter $\mathbf{1}$ for $\boldsymbol{K X X}$, and click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create nodes ( $\mathbf{N}$ command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Nodes $>$ In Active CS

- A total of 9 nodes will be created (Table 6.4).
- Referring to Table 6.4 , enter $x$ - and $y$-coordinates of node 1 , and Click on Apply. This action will keep the Create Nodes in Active Coordinate System dialog box open. If the Node number field is left blank, then ANSYS will assign the lowest available node number to the node that is being created.
- Repeat the same procedure for the nodes 2 through 9 .
- After entering the $x$ - and $y$-coordinates of node 9 , click on $O K$ (instead of Apply).
- The nodes should appear in the Graphics Window, as shown in Fig. 6.16.
- Create elements ( $\mathbf{E}$ command) using the following menu path:


## Main Menu > Preprocessor > Modeling > Create > Elements > Auto Numbered > Thru Nodes

- Pick Menu appears; refer to Fig. 6.17 to create elements by picking four nodes at a time and clicking on Apply in between.
- Observe the elements created after clicking on Apply in the Pick Menu.
- Repeat until the last element is created.
- Click on $O K$ when the last element is created.


Fig. 6.16 Generation of nodes.


Fig. 6.17 Generation of elements.

- Review elements:
- Turn on element numbering using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Select Element numbers from the first pull-down menu; click on $\boldsymbol{O K}$.
- Plot elements (EPLOT command) using the following menu path:

Utility Menu > Plot > Elements

- Figure 6.17 shows the outcome of this action as it appears in the Graphics Window.
- Turn off element numbering and turn on node numbering using the following menu path:
- Place a checkmark by clicking on the empty box next to NODE Node numbers.
- Select No numbering from the first pull-down menu.
- Click on $\boldsymbol{O K}$.
- Plot nodes (NPLOT command) using the following menu path: Utility Menu > Plot $>$ Nodes
- Figure 6.16 shows the outcome of this action as it appears in the Graphics Window.


## SOLUTION

- Apply temperature boundary conditions (D command) using the following menu path:

Main Menu > Solution > Define Loads $>$ Apply $>$ Thermal $>$ Temperature $>$ On Nodes

- Pick Menu appears; pick nodes 3 through 7 along the boundary (Fig. 6.16) and click on OK on Pick Menu.
- Highlight TEMP and enter $\mathbf{1}$ for VALUE; click on OK (Fig. 6.18).
- Apply body load on elements (BFE command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Thermal > Heat Generat > On Elements

- Pick Menu appears; click on Pick All.
- Enter 1 for VAL1 (leave other fields untouched, as shown in Fig. 6.19).
- Click on OK.
- Obtain solution (SOLVE command) using the following menu path: Main Menu $>$ Solution $>$ Solve $>$ Current LS
- Confirmation Window appears along with Status Report Window.
- Review status/ If OK, close the Status Report Window and click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review temperature values (PRNSOL command) using the following menu path:

Main Menu > General Postproc > List Results > Nodal Solution

- Highlight DOF solution on the left list and Temperature TEMP on the right list; click on $O K$.
- The list will appear in a new window, as shown in Fig. 6.20.


Fig. 6.18 Application of temperature boundary conditions on nodes.


Fig. 6.19 Application of heat generation condition on elements.


Fig. 6.20 Nodal solution for temperature.

### 6.2 Principle of Minimum Potential Energy

Galerkin's method is not always suitable for all structural problems because of difficulties in mathematically describing the structural geometry and/or the boundary conditions. An alternative to Galerkin's method is the principle of minimum potential energy (Washizu 1982; Dym and Shames 1973).

The energy method involves determination of the stationary values of the global energy. This requires the approximation of the functional behavior of the dependent variable so that the global energy becomes stationary. The stationary value can be a maximum, a minimum or a neutral point. With an understanding of variational calculus, the minimum stationary value leading to stable equilibrium (Fig. 6.21) is obtained by requiring the first variation of the global energy to vanish.

Avoiding the details of variational calculus, the concepts of differential calculus can be used to perform the minimization of the global energy. In solid mechanics, this is known as the principle of minimum potential energy, which states that among all compatible displacement fields satisfying the boundary conditions (kinematically admissible), the correct displacement field satisfying the equilibrium equations is the one that ren-


Fig. 6.21 Schematics of stable, neutral, and unstable equilibrium points of the global energy.
ders the potential energy an absolute minimum. A solution satisfying both equilibrium equations and boundary conditions is, of course, "exact"; however, such solutions are difficult, if not impossible, to construct for complex problems. Therefore, approximate solutions are obtained by assuming kinematically admissible displacement fields with unknown coefficients. The values of these coefficients are determined in such a way that the total potential energy of the system is a minimum.

The principle of virtual work is applicable for any material behavior, whereas the principle of minimum potential energy is applicable only for elastic materials. However, both principles yield the same element equations for elastic materials.

The total potential energy of the structural system shown in Fig. 6.22 is defined as

$$
\begin{equation*}
\pi_{p}=W+\Omega \tag{6.165}
\end{equation*}
$$

in which $W$ is the strain energy and $\Omega$ is the potential energy arising from the presence of body forces, surface tractions, and the initial residual stresses. Strain energy is the capacity of the internal forces (or stresses) to do work through strains in the structure.

For a linear elastic material, the strain energy of the deformed structure is given by

$$
\begin{equation*}
W=\frac{1}{2} \int_{V}\left(\varepsilon-\varepsilon^{*}\right)^{T} \sigma d V \tag{6.166}
\end{equation*}
$$



Fig. 6.22 A 3D body with displacement constraints, body and concentrated forces, and surface tractions.
where $\boldsymbol{\sigma}$ is the vector of stress components arising from the difference between the total strains, $\varepsilon$, and initial strains, $\varepsilon^{*}$. It can be expressed as

$$
\begin{equation*}
\sigma=\mathbf{D}\left(\varepsilon-\varepsilon^{*}\right) \tag{6.167}
\end{equation*}
$$

in which

$$
\boldsymbol{\sigma}^{T}=\left\{\begin{array}{llllll}
\sigma_{x x} & \sigma_{y y} & \sigma_{z z} & \sigma_{x y} & \sigma_{y z} & \sigma_{z x} \tag{6.168}
\end{array}\right\}
$$

and

$$
\varepsilon^{T}=\left\{\begin{array}{llllll}
\varepsilon_{x x} & \varepsilon_{y y} & \varepsilon_{z z} & \gamma_{x y} & \gamma_{y z} & \gamma_{z x} \tag{6.169}
\end{array}\right\}
$$

and the material property matrix

$$
\mathbf{D}=\frac{E}{(1+v)(1-2 v)}\left[\begin{array}{cccccc}
1-v & v & v & 0 & 0 & 0  \tag{6.170}\\
v & 1-v & v & 0 & 0 & 0 \\
v & v & 1-v & 0 & 0 & 0 \\
0 & 0 & 0 & \frac{(1-2 v)}{2} & 0 & 0 \\
0 & 0 & 0 & 0 & \frac{(1-2 v)}{2} & 0 \\
0 & 0 & 0 & 0 & 0 & \frac{(1-2 v)}{2}
\end{array}\right]
$$

where $\sigma_{i j}$ and $\varepsilon_{i j}$ represent the stress and strain components, with $i, j=x, y, z$ being the Cartesian coordinates. The elastic modulus and

Poisson's ratio are denoted by $E$ and $v$, respectively. In the presence of temperature change, the initial strains can be expressed as

$$
\varepsilon^{* T}=\left\{\begin{array}{llllll}
\alpha \Delta T & \alpha \Delta T & \alpha \Delta T & 0 & 0 & 0 \tag{6.171}
\end{array}\right\}
$$

where $\alpha$ is the coefficient of thermal expansion and $\Delta T$ is the temperature change with respect to a reference state.

The potential energy arising from the presence of body forces, $\mathbf{b}$, surface tractions, $\mathbf{T}$, and the initial residual stresses, $\boldsymbol{\sigma}^{*}$, is given by

$$
\begin{equation*}
\Omega=-\int_{V} \mathbf{u}^{T} \mathbf{b} d V-\int_{S_{\sigma}} \mathbf{u}^{T} \mathbf{T} d S+\int_{V}^{T} \varepsilon^{*} \boldsymbol{\sigma}^{*} d V \tag{6.172}
\end{equation*}
$$

with

$$
\begin{align*}
\mathbf{b}^{T} & =\left\{\begin{array}{lll}
b_{x} & b_{y} & b_{z}
\end{array}\right\}  \tag{6.173}\\
\mathbf{T}^{T} & =\left\{\begin{array}{lll}
T_{x} & T_{y} & T_{z}
\end{array}\right\}  \tag{6.174}\\
\mathbf{u}^{T} & =\left\{\begin{array}{lll}
u_{x} & u_{y} & u_{z}
\end{array}\right\} \tag{6.175}
\end{align*}
$$

in which $b_{x}, b_{y}$, and $b_{z}$ are the components of body force (in units of force per unit volume), and $T_{x}, T_{y}$, and $T_{z}$ represent the components of the applied traction vector (in units of force per unit area) over the surface defined by $S_{\sigma}$. The entire surface of the body having a volume of $V$ is defined by $S$, with segments $S_{u}$ and $S_{\sigma}$ subjected to displacement and traction conditions, respectively. The displacement components are given by $u_{x}, u_{y}$, and $u_{z}$ in the $x-y$-, and $z$-directions, respectively. Also, included in the expression for the total potential is the initial residual stresses denoted by $\sigma^{*}$. The initial stresses could be measured, but their prediction without full knowledge of the material's history is impossible.

After partitioning the entire domain occupied by volume $V$ into $E$ number of elements with volume $V^{e}$, the total potential energy of the system can be rewritten as

$$
\begin{equation*}
\pi_{p}\left(u_{x}, u_{y}, u_{z}\right)=\sum_{e=1}^{E} \pi_{p}^{(e)}\left(u_{x}, u_{y}, u_{z}\right) \tag{6.176}
\end{equation*}
$$

in which

$$
\begin{align*}
\pi_{p}^{(e)}=\frac{1}{2} \int_{V^{(e)}} \varepsilon^{T} \mathbf{D} \varepsilon d V & -\int_{V^{(e)}} \varepsilon^{T} \mathbf{D} \varepsilon^{*} d V+\frac{1}{2} \int_{V^{(e)}} \varepsilon^{* T} \mathbf{D} \varepsilon^{*} d V  \tag{6.177}\\
& -\int_{V^{(e)}} \mathbf{u}^{T} \mathbf{b} d V-\int_{S_{\sigma}^{(e)}} \mathbf{u}^{T} \mathbf{T} d S+\int_{V^{(e)}} \varepsilon^{T} \boldsymbol{\sigma}^{*} d V
\end{align*}
$$

where the superscript $e$ denotes a specific element.
Based on kinematical considerations, the components of the total strain vector, $\boldsymbol{\varepsilon}$, in terms of the displacement components are expressed as

$$
\left\{\begin{array}{l}
\varepsilon_{x x}  \tag{6.178}\\
\varepsilon_{y y} \\
\varepsilon_{z z} \\
\gamma_{x y} \\
\gamma_{y z} \\
\gamma_{z x}
\end{array}\right\}=\left[\begin{array}{ccc}
\frac{\partial}{\partial x} & 0 & 0 \\
0 & \frac{\partial}{\partial y} & 0 \\
0 & 0 & \frac{\partial}{\partial z} \\
\frac{\partial}{\partial y} & \frac{\partial}{\partial x} & 0 \\
0 & \frac{\partial}{\partial z} & \frac{\partial}{\partial y} \\
\frac{\partial}{\partial z} & 0 & \frac{\partial}{\partial x}
\end{array}\right]\left\{\begin{array}{l}
u_{x} \\
u_{y} \\
u_{z}
\end{array}\right\} \text { or } \quad \varepsilon=\mathbf{L} \mathbf{u}
$$

in which $\mathbf{L}$ is the differential operator matrix.
The finite element process seeks a minimum in the potential energy based on the approximate form of the dependent variables (displacement components) within each element. The greater the number of degrees of freedom associated with the element (usually means increasing the number of nodes), the more closely the solution will approximate the true equilibrium position. Within each element, the approximation to the displacement components can be expressed as

$$
\begin{align*}
& u_{x}^{(e)} \approx \tilde{u}_{x}^{(e)}=\sum_{r=1}^{n} N_{r}^{(e)} u_{x_{r}}^{(e)} \\
& u_{y}^{(e)} \approx \tilde{u}_{y}^{(e)}=\sum_{r=1}^{n} N_{r}^{(e)} u_{y_{r}}^{(e)}  \tag{6.179}\\
& u_{z}^{(e)} \approx \tilde{u}_{z}^{(e)}=\sum_{r=1}^{n} N_{r}^{(e)} u_{z_{r}}^{(e)}
\end{align*}
$$

with $n$ representing the number of nodes associated with element $e$. The nodal unknowns and shape functions are denoted by $u_{x_{r}}^{(e)}, u_{y_{r}}^{(e)}, u_{z_{r}}^{(e)}$, and $N_{r}^{(e)}$, respectively. In matrix form, the approximate displacement components can be expressed as

$$
\begin{equation*}
\tilde{\mathbf{u}}^{(e)}=\mathbf{N}^{(e) T} \mathbf{U}^{(e)} \tag{6.180}
\end{equation*}
$$

in which

$$
\begin{align*}
& \tilde{\mathbf{u}}^{(e) T}=\left\{\begin{array}{lll}
\tilde{u}_{x}^{(e)} & \tilde{u}_{y}^{(e)} & \tilde{u}_{z}^{(e)}
\end{array}\right\}  \tag{6.181}\\
& \mathbf{N}^{(e) T}=\left[\begin{array}{ccccccccccc}
N_{1} & 0 & 0 & N_{2} & 0 & 0 & \ldots & N_{n} & 0 & 0 \\
0 & N_{1} & 0 & 0 & N_{2} & 0 & \ldots & . & 0 & N_{n} & 0 \\
0 & 0 & N_{1} & 0 & 0 & N_{2} & \ldots & . & 0 & 0 & N_{n}
\end{array}\right]_{3 \times 3 n}  \tag{6.182}\\
& \mathbf{U}^{(e) T}=\left\{\begin{array}{llllllllll}
u_{x_{1}}^{(e)} & u_{y_{1}}^{(e)} & u_{z_{1}}^{(e)} & u_{x_{2}}^{(e)} & u_{y_{2}}^{(e)} & u_{z_{2}}^{(e)} & \cdots & u_{x_{n}}^{(e)} & u_{y_{n}}^{(e)} & u_{z_{n}}^{(e)}
\end{array}\right\} \tag{6.183}
\end{align*}
$$

With the approximate form of the displacement components, the strain components within each element can be expressed as

$$
\begin{equation*}
\boldsymbol{\varepsilon} \approx \mathbf{B}^{(e)} \mathbf{U}^{(e)} \tag{6.184}
\end{equation*}
$$

where

$$
\begin{equation*}
\mathbf{B}^{(e)}=\mathbf{L} \mathbf{N}^{(e) T} \tag{6.185}
\end{equation*}
$$

leading to the expression for the total potential in terms of element nodal displacements, $\mathbf{U}^{(e)}$

$$
\begin{equation*}
\pi_{p}^{(e)}=\frac{1}{2} \mathbf{U}^{(e) T} \mathbf{k}^{(e)} \mathbf{U}^{(e)}-\mathbf{U}^{(e) T} \mathbf{p}^{(e)}+\frac{1}{2} \int_{V^{(e)}} \varepsilon^{* T} \mathbf{D} \varepsilon^{*} d V \tag{6.186}
\end{equation*}
$$

in which the element stiffness matrix, $\mathbf{k}^{(e)}$, and the element force vector, $\mathbf{p}^{(e)}$, are defined as

$$
\begin{equation*}
\mathbf{k}^{(e)}=\int_{V^{(e)}} \mathbf{B}^{(e) T} \mathbf{D} \mathbf{B}^{(e)} d V \tag{6.187}
\end{equation*}
$$

and

$$
\begin{equation*}
\mathbf{p}^{(e)}=\mathbf{p}_{\mathrm{b}}^{(e)}+\mathbf{p}_{\mathrm{T}}^{(e)}+\mathbf{p}_{\varepsilon^{*}}^{(e)}-\mathbf{p}_{\sigma^{*}}^{(e)} \tag{6.188}
\end{equation*}
$$

with $\mathbf{p}_{\mathrm{b}}^{(e)}, \mathbf{p}_{\mathrm{T}}^{(e)}, \mathbf{p}_{\varepsilon^{*}}^{(e)}$, and $\mathbf{p}_{\sigma^{*}}^{(e)}$ representing the element load vectors due to body forces, surface tractions (forces), initial strains, and initial stresses, respectively, defined by

$$
\begin{align*}
& \mathbf{p}_{\mathbf{b}}^{(e)}=\int_{V^{(e)}} \mathbf{N}^{(e)} \mathbf{b} d V \\
& \mathbf{p}_{\mathbf{T}}^{(e)}=\int_{S_{\sigma}^{(e)}} \mathbf{N}^{(e)} \mathbf{T} d S  \tag{6.189}\\
& \mathbf{p}_{\varepsilon^{*}}^{(e)}=\int_{V^{(e)}} \mathbf{B}^{(e) T} \mathbf{D} \varepsilon^{*} d V \\
& \mathbf{p}_{\sigma^{*}}^{(e)}=\int_{V^{(e)}} \mathbf{B}^{(e) T} \boldsymbol{\sigma}^{*} d V
\end{align*}
$$

Evaluation of these integrals results in the statically equivalent nodal forces in the elements affected by the body force, the surface tractions, and the initial strains and initial stresses. In the presence of external concentrated forces acting on various nodes, the potential energy is modified as

$$
\begin{align*}
\pi_{p} & =\frac{1}{2} \mathbf{U}^{T}\left\{\sum_{e=1}^{E} \mathbf{k}^{(e)}\right\} \mathbf{U}-\mathbf{U}^{T}\left\{\sum_{e=1}^{E}\left(\mathbf{p}_{\mathbf{b}}^{(e)}+\mathbf{p}_{\mathbf{T}}^{(e)}+\mathbf{p}_{\varepsilon^{*}}^{(e)}-\mathbf{p}_{\sigma^{*}}^{(e)}\right)-\mathbf{P}_{\mathbf{c}}\right\} \\
& +\frac{1}{2} \sum_{e=1}^{E} \int_{V^{(e)}} \varepsilon^{* T} \mathbf{D} \boldsymbol{\varepsilon}^{*} d V \tag{6.190}
\end{align*}
$$

where $\mathbf{P}_{\mathbf{c}}$ is the vector of nodal forces and $\mathbf{U}$ represents the vector of nodal displacements for the entire structure. Note that each component of the element nodal displacement vector, $\mathbf{U}^{(e)}$, appears in the global (system) nodal displacement vector, $\mathbf{U}$. Therefore, the element nodal displacement vector $\mathbf{U}^{(e)}$ can be replaced by $\mathbf{U}$ with the appropriate enlargement of the element matrices and vectors in the expression for the potential energy by adding the required number of zero elements and rearranging. The summation in the expression for the potential energy implies the expansion of the element matrices to the size of the global (system) matrix while collecting the overlapping terms.

Minimization of the total potential energy requires that

$$
\begin{equation*}
\left\{\frac{\partial \pi_{p}}{\partial \mathbf{U}}\right\}=0 \tag{6.191}
\end{equation*}
$$

leading to the system (global) equilibrium equations in the form

$$
\begin{equation*}
\mathbf{K U}=\mathbf{P} \tag{6.192}
\end{equation*}
$$

in which $\mathbf{K}$ and $\mathbf{P}$ are the assembled (global) stiffness matrix and the assembled (global) nodal load vector, respectively, defined by

$$
\begin{equation*}
\mathbf{K}=\sum_{e=1}^{E} \mathbf{k}^{(e)} \tag{6.193}
\end{equation*}
$$

and

$$
\begin{equation*}
\mathbf{P}=\sum_{e=1}^{E}\left(\mathbf{p}_{\mathbf{b}}^{(e)}+\mathbf{p}_{\mathbf{T}}^{(e)}+\mathbf{p}_{\varepsilon^{*}}^{(e)}-\mathbf{p}_{\sigma^{*}}^{(e)}\right)-\mathbf{P}_{\mathbf{c}} \tag{6.194}
\end{equation*}
$$

This global equilibrium equation cannot be solved unless boundary constraints are imposed to suppress the rigid-body motion. Otherwise, the global stiffness matrix becomes singular.

After obtaining the solution to the nodal displacements of the system equilibrium equations, the stresses within the element can be determined from

$$
\begin{equation*}
\boldsymbol{\sigma}=\mathbf{D B}^{(e)} \mathbf{U}^{(e)}-\mathbf{D}^{*}+\boldsymbol{\sigma}^{*} \tag{6.195}
\end{equation*}
$$

The global stiffness matrix and the load vector require the evaluation of the integrals associated with the element stiffness matrix and the element nodal load vector.

### 6.2.1 Example: One-dimensional Analysis with Line Elements

The application of this approach is demonstrated by computing the displacements and strains in a rod constructed of three concentric sections of different materials. As shown in Fig. 6.23, the rod has a uniform cross section and is subjected to a concentrated horizontal load, P , at the second joint, and the boundary conditions are specified as $u_{x}(x=0)=0$ and $u_{x}(x=L)=0$.

The domain is discretized with 3 linear line elements having two nodes, as shown in Fig. 6.24. The global coordinates of each node in domain $D$ are specified by $x_{i}$, with $i=1,2,3,4$. The nodal values of the dependent variable associated with element $e$ are specified at its first and second nodes by $u_{x_{i}}^{(e)}$ and $u_{x_{j}}^{(e)}$, respectively.
For the domain discretized with three elements and four nodes, the local and global nodes are numbered as shown in Table 6.6.


Fig. 6.23 A rod constrained at both ends, subjected to a concentrated force.


Fig. 6.24 Finite element discretization of the rod with three elements.

Table 6.6 Local and global node numbers.

| Element <br> Number <br> $(e)$ | Node 1 | Node 2 |
| :---: | :---: | :---: |
| 1 | 1 | 2 |
| 2 | 2 | 3 |
| 3 | 3 | 4 |

Within each element, the approximation to the displacement component can be expressed as

$$
\begin{equation*}
u_{x}^{(e)} \approx \tilde{u}_{x}^{(e)}=\sum_{r=1}^{2} N_{r}^{(e)} u_{x_{r}}^{(e)} \tag{6.196}
\end{equation*}
$$

The nodal unknowns and shape functions are denoted by $u_{x_{r}}^{(e)}$ and $N_{r}^{(e)}$, respectively. In matrix form, the approximate displacement components can be expressed as

$$
\begin{equation*}
\tilde{\mathbf{u}}^{(e)}=\mathbf{N}^{(e) T} \mathbf{U}^{(e)} \tag{6.197}
\end{equation*}
$$

with

$$
\mathbf{N}^{(e) T}=\left\{\begin{array}{lll}
N_{1}^{(e)} & N_{2}^{(e)}
\end{array}\right\} \quad \text { and } \quad \mathbf{U}^{(e)}=\left\{\begin{array}{l}
u_{x_{1}}^{(e)}  \tag{6.198}\\
u_{x_{2}}^{(e)}
\end{array}\right\}
$$

in which the shape functions are

$$
\begin{equation*}
N_{1}^{(e)}=\frac{x_{2}^{(e)}-x}{x_{2}^{(e)}-x_{1}^{(e)}} \quad \text { and } \quad N_{2}^{(e)}=\frac{x-x_{1}^{(e)}}{x_{2}^{(e)}-x_{1}^{(e)}} \tag{6.199}
\end{equation*}
$$

With the approximate form of the displacement components and $\mathrm{L}=\partial / \partial x$, the shape matrix can be obtained from

$$
\mathbf{B}^{(e)}=\frac{\partial}{\partial x}\left[\begin{array}{ll}
N_{1}^{(e)} & N_{2}^{(e)} \tag{6.200}
\end{array}\right]
$$

For a constant cross section, $A^{(e)}$, and elastic modulus, $E^{(e)}$, in each element, the element stiffness matrix is

$$
\begin{align*}
\mathbf{k}^{(e)} & =\int_{V^{(e)}} \mathbf{B}^{(e) T} \mathbf{D} \mathbf{B}^{(e)} d V \\
& =A^{(e)} \int_{x_{1}^{(e)}}^{x_{2}^{(e)}} \frac{\partial}{\partial x}\left[\begin{array}{l}
N_{1}^{(e)} \\
N_{2}^{(e)}
\end{array}\right] E^{(e)} \frac{\partial}{\partial x}\left[\begin{array}{ll}
N_{1}^{(e)} & N_{2}^{(e)}
\end{array}\right] d x \tag{6.201}
\end{align*}
$$

Substituting for the shape functions, the element stiffness matrix becomes

$$
\mathbf{k}^{(e)}=-\frac{A^{(e)} E^{(e)}}{\left(x_{2}^{(e)}-x_{1}^{(e)}\right)^{2}} \int_{x_{1}^{(e)}}^{x_{2}^{(e)}}\left[\begin{array}{cc}
1 & -1  \tag{6.202}\\
-1 & 1
\end{array}\right] d x
$$

Integration along the element length results in

$$
\begin{align*}
\mathbf{k}^{(e)} & =-\frac{A^{(e)} E^{(e)}}{\left(x_{2}^{(e)}-x_{1}^{(e)}\right)}\left[\begin{array}{cc}
1 & -1 \\
-1 & 1
\end{array}\right]=-\frac{A^{(e)} E^{(e)}}{L^{(e)}}\left[\begin{array}{cc}
1 & -1 \\
-1 & 1
\end{array}\right]  \tag{6.203}\\
& =-\alpha^{(e)}\left[\begin{array}{cc}
1 & -1 \\
-1 & 1
\end{array}\right]
\end{align*}
$$

in which $L^{(e)}=\left(x_{2}^{(e)}-x_{1}^{(e)}\right)$ and $\alpha^{(e)}=A^{(e)} E^{(e)} / L^{(e)}$. The element stiffness matrices are computed as

$$
\mathbf{k}^{(1)}=\left[\begin{array}{cc}
1 & 2 \\
\alpha^{(1)} & -\alpha^{(1)}  \tag{6.204}\\
-\alpha^{(1)} & \alpha^{(1)}
\end{array}\right]\left[\begin{array}{l}
1 \\
2
\end{array}\right.
$$

$$
\begin{gather*}
2 \\
\mathbf{k}^{(2)}=\left[\begin{array}{cc}
\alpha^{(2)} & -\alpha^{(2)} \\
-\alpha^{(2)} & \alpha^{(2)}
\end{array}\right] \begin{array}{|c}
2 \\
3
\end{array}  \tag{6.205}\\
\mathbf{k}^{(3)}=\left[\begin{array}{cc}
\alpha^{(3)} & -\alpha^{(3)} \\
-\alpha^{(3)} & \alpha^{(3)}
\end{array}\right] \begin{array}{|c}
4 \\
\hline
\end{array}
\end{gather*}
$$

The element load vector, $\mathbf{p}_{\mathbf{T}}^{(e)}$, due to the unknown nodal forces, $T_{x_{i}}$ and $T_{x_{j}}$ at nodes $i$ and $j$, respectively (Fig. 6.25), can be obtained from

$$
\begin{align*}
\mathbf{p}_{\mathbf{T}}^{(e)} & =\int_{S_{\sigma}^{(e)}} \mathbf{N}^{(e)} \mathbf{T} d S=\left\{\begin{array}{l}
N_{1}^{(e)}\left(x=x_{1}^{(e)}\right) \\
N_{2}^{(e)}\left(x=x_{1}^{(e)}\right)
\end{array}\right\} T_{x_{1}} \\
& +\left\{\begin{array}{l}
N_{1}^{(e)}\left(x=x_{2}^{(e)}\right) \\
N_{2}^{(e)}\left(x=x_{2}^{(e)}\right)
\end{array}\right\} T_{x_{2}} \tag{6.207}
\end{align*}
$$

Evaluating the shape functions results in a load vector of the form

$$
\begin{align*}
\mathbf{p}_{\mathbf{T}}^{(e)} & =\left\{\begin{array}{c}
-1 \\
0
\end{array}\right\} T_{x_{1}}+\left\{\begin{array}{l}
0 \\
1
\end{array}\right\} T_{x_{2}} \\
& =\left\{\begin{array}{c}
-T_{x_{1}} \\
0
\end{array}\right\}+\left\{\begin{array}{c}
0 \\
T_{x_{2}}
\end{array}\right\}=\left\{\begin{array}{c}
-T_{x_{1}} \\
T_{x_{2}}
\end{array}\right\} \tag{6.208}
\end{align*}
$$

Associated with each element, the load vectors become

$$
\mathbf{p}_{\mathbf{T}}^{(1)}=\left\{\begin{array}{c}
-T_{x_{1}}  \tag{6.209}\\
T_{x_{2}}
\end{array}\right\}, \quad \mathbf{p}_{\mathbf{T}}^{(2)}=\left\{\begin{array}{c}
-T_{x_{2}} \\
T_{x_{3}}+P
\end{array}\right\}, \quad \mathbf{p}_{\mathbf{T}}^{(3)}=\left\{\begin{array}{c}
-T_{x_{3}} \\
T_{x_{4}}
\end{array}\right\}
$$

The global coefficient matrix, $\mathbf{K}$, and the load vector, $\mathbf{P}_{\mathbf{T}}$, are obtained from the "expanded" element coefficient matrices, $\mathbf{k}^{(e)}$, and the element load vectors, $\mathbf{p}_{\mathbf{T}}^{(e)}$, by summation in the form


Fig. 6.25 A typical linear line element with two nodes.

$$
\begin{equation*}
\mathbf{K}=\sum_{e=1}^{E} \mathbf{k}^{(e)} \quad \text { and } \quad \mathbf{P}_{\mathbf{T}}=\sum_{e=1}^{E} \mathbf{p}_{\mathbf{T}}^{(e)} \tag{6.210}
\end{equation*}
$$

The "expanded" element matrices are the same size as the global matrix but have rows and columns of zeros corresponding to the nodes not associated with element (e). Specifically, the expanded form of the element stiffness and load vector becomes

$$
\begin{align*}
& \begin{array}{llll}
1 & 2 & 3 & 4 \\
1
\end{array} \\
& \mathbf{k}^{(1)}=\left[\begin{array}{cccc|c}
\alpha^{(1)} & -\alpha^{(1)} & 0 & 0 & 1 \\
-\alpha^{(1)} & \alpha^{(1)} & 0 & 0 & 2 \\
0 & 0 & 0 & 0 & 3 \\
0 & 0 & 0 & 0
\end{array}\right] \mathbf{p}_{\mathbf{T}}^{(1)}=\left[\begin{array}{cc}
-T_{x_{1}} & {[1]} \\
T_{x_{2}} & 2 \\
0 & 2 \\
0 & 3
\end{array}\right.  \tag{6.211}\\
& \begin{array}{lllll}
1 & 2 & 3 & 4 & 1
\end{array}
\end{align*}
$$

$$
\begin{align*}
& \text { (1) } 2 \text { 3 } 3 \text { (1) } \\
& \mathbf{k}^{(3)}=\left[\begin{array}{cccc|c}
0 & 0 & 0 & 0 & 1 \\
0 & 0 & 0 & 0 & 2 \\
0 & 0 & \alpha^{(3)} & -\alpha^{(3)} & 3 \\
0 & 0 & -\alpha^{(3)} & \alpha^{(3)} & \mathbf{p}_{\mathrm{T}}^{(3)} \\
\hline
\end{array}\right] \begin{array}{c|c}
0 & 1 \\
0 & 2 \\
-T_{x_{3}} & 3 \\
T_{x_{4}} & 4
\end{array} \tag{6.213}
\end{align*}
$$

In accordance with Eq. (6.210) and (6.192), the global equilibrium equations can be written as

$$
\left[\begin{array}{cccc}
\alpha^{(1)} & -\alpha^{(1)} & 0 & 0  \tag{6.214}\\
-\alpha^{(1)} & \left(\alpha^{(1)}+\alpha^{(2)}\right) & -\alpha^{(2)} & 0 \\
0 & -\alpha^{(2)} & \left(\alpha^{(2)}+\alpha^{(3)}\right) & -\alpha^{(3)} \\
0 & 0 & -\alpha^{(3)} & \alpha^{((3))}
\end{array}\right]\left\{\begin{array}{l}
u_{x_{1}} \\
u_{x_{2}} \\
u_{x_{3}} \\
u_{x_{4}}
\end{array}\right\}=\left\{\begin{array}{c}
-T_{x_{1}} \\
T_{x_{2}}-T_{x_{2}} \\
T_{x_{3}}+P-T_{x_{3}} \\
T_{x_{4}}
\end{array}\right\}(
$$

Enforcing the boundary conditions of $u_{x_{1}}=u_{x_{4}}=0$ leads to

$$
\left[\begin{array}{cccc}
\alpha^{(1)} & -\alpha^{(1)} & 0 & 0  \tag{6.215}\\
-\alpha^{(1)} & \left(\alpha^{(1)}+\alpha^{(2)}\right) & -\alpha^{(2)} & 0 \\
0 & -\alpha^{(2)} & \left(\alpha^{(2)}+\alpha^{(3)}\right) & -\alpha^{(3)} \\
0 & 0 & -\alpha^{(3)} & \alpha^{(3)}
\end{array}\right]\left\{\begin{array}{c}
0 \\
u_{x_{2}} \\
u_{x_{3}} \\
0
\end{array}\right\}=\left\{\begin{array}{c}
-T_{x_{1}} \\
0 \\
P \\
T_{x_{4}}
\end{array}\right\}
$$

This system of equations can be partitioned in the form

$$
\left[\begin{array}{cccc}
-\alpha^{(1)} & \left(\alpha^{(1)}+\alpha^{(2)}\right) & -\alpha^{(2)} & 0  \tag{6.216a}\\
0 & -\alpha^{(2)} & \left(\alpha^{(2)}+\alpha^{(3)}\right) & 0
\end{array}\right]\left\{\begin{array}{c}
0 \\
u_{x 2} \\
u_{x 3} \\
0
\end{array}\right\}=\left\{\begin{array}{l}
0 \\
P
\end{array}\right\}
$$

or

$$
\left[\begin{array}{cc}
\left(\alpha^{(1)}+\alpha^{(2)}\right) & -\alpha^{(2)}  \tag{6.216b}\\
-\alpha^{(2)} & \left(\alpha^{(2)}+\alpha^{(3)}\right.
\end{array}\right]\left\{\begin{array}{l}
u_{x 2} \\
u_{x 3}
\end{array}\right\}=\left\{\begin{array}{l}
0 \\
P
\end{array}\right\}
$$

and

$$
\left[\begin{array}{cccc}
\alpha^{(1)} & -\alpha^{(1)} & 0 & 0  \tag{6.217a}\\
0 & 0 & -\alpha^{(3)} & \alpha^{(3)}
\end{array}\right]\left\{\begin{array}{c}
0 \\
u_{x_{2}} \\
u_{x_{3}} \\
0
\end{array}\right\}=\left\{\begin{array}{c}
-T_{x_{1}} \\
T_{x_{4}}
\end{array}\right\}
$$

or

$$
\begin{equation*}
T_{x_{1}}=\alpha^{(1)} u_{x_{2}} \quad \text { and } \quad T_{x_{4}}=-\alpha^{(3)} u_{x_{3}} \tag{6.217b}
\end{equation*}
$$

Solution to nodal displacements results in

$$
\begin{align*}
& u_{x_{2}}=\frac{\alpha^{(2)}}{\left(\alpha^{(1)} \alpha^{(2)}+\alpha^{(1)} \alpha^{(3)}+\alpha^{(2)} \alpha^{(3)}\right)} P  \tag{6.218}\\
& u_{x_{3}}=\frac{\alpha^{(1)}+\alpha^{(2)}}{\left(\alpha^{(1)} \alpha^{(2)}+\alpha^{(1)} \alpha^{(3)}+\alpha^{(2)} \alpha^{(3)}\right)} P \tag{6.219}
\end{align*}
$$

With these nodal displacements, the reaction forces are computed as

$$
\begin{align*}
& T_{x_{1}}=\frac{\alpha^{(1)} \alpha^{(2)}}{\left(\alpha^{(1)} \alpha^{(2)}+\alpha^{(1)} \alpha^{(3)}+\alpha^{(2)} \alpha^{(3)}\right)} P  \tag{6.220}\\
& T_{x_{4}}=-\frac{\alpha^{(3)}\left(\alpha^{(1)}+\alpha^{(2)}\right)}{\left(\alpha^{(1)} \alpha^{(2)}+\alpha^{(1)} \alpha^{(3)}+\alpha^{(2)} \alpha^{(3)}\right)} P \tag{6.221}
\end{align*}
$$

Finally, the strains are computed as

$$
\begin{align*}
& \varepsilon_{x x}^{(1)}=\frac{1}{L^{(1)}}\left(u_{x_{2}}-u_{x_{1}}\right)=\frac{\alpha^{(2)}}{\left(\alpha^{(1)} \alpha^{(2)}+\alpha^{(1)} \alpha^{(3)}+\alpha^{(2)} \alpha^{(3)}\right) L^{(1)}} P  \tag{6.222}\\
& \varepsilon_{x x}^{(2)}=\frac{1}{L^{(2)}}\left(u_{x_{3}}-u_{x_{2}}\right)=\frac{\alpha^{(1)}}{\left(\alpha^{(1)} \alpha^{(2)}+\alpha^{(1)} \alpha^{(3)}+\alpha^{(2)} \alpha^{(3)}\right) L^{(2)}} P  \tag{6.223}\\
& \varepsilon_{x x}^{(3)}=\frac{1}{L^{(3)}}\left(u_{x_{4}}-u_{x_{3}}\right)=-\frac{\left(\alpha^{(1)}+\alpha^{(2)}\right)}{\left(\alpha^{(1)} \alpha^{(2)}+\alpha^{(1)} \alpha^{(3)}+\alpha^{(2)} \alpha^{(3)}\right) L^{(3)}} P \tag{6.224}
\end{align*}
$$

### 6.2.2 Two-dimensional Structural Analysis

The three-dimensional analysis of either "thin" or "long" components subjected to in-plane external loading conditions can be reduced to a twodimensional analysis under certain assumptions referred to as "plane stress" and "plane strain" conditions.

### 6.2.2.1 Plane Stress Conditions

A state of plane stress exists for thin components subjected only to in-plane external loading, i.e., no lateral loads (Fig. 6.26). Due to a small thickness-to-characteristic length ratio and in-plane external loading only, there is no


Fig. 6.26 Thin body with in-plane loading; suitable for plane stress idealization.
out-of-plane displacement component, $u_{z}$, and the shear strain components associated with the thickness direction, $\gamma_{x z}$ and $\gamma_{y z}$, are very small and assumed to be zero. Therefore, the stress components, $\sigma_{z z}, \sigma_{x z}$, and $\sigma_{y z}$, associated with the thickness direction vanish. Under these assumptions, the displacement, $\mathbf{u}$, stress, $\boldsymbol{\sigma}$, strain, $\boldsymbol{\varepsilon}$, and traction, $\mathbf{T}$, vectors, and material property matrix, $\mathbf{D}$, reduce to

$$
\begin{align*}
\mathbf{u}^{T} & =\left\{\begin{array}{ll}
u_{x} & u_{y}
\end{array}\right\} \\
\boldsymbol{\sigma}^{T} & =\left\{\begin{array}{lll}
\sigma_{x x} & \sigma_{y y} & \sigma_{x y}
\end{array}\right\} \\
\boldsymbol{\varepsilon}^{T} & =\left\{\begin{array}{lll}
\varepsilon_{x x} & \varepsilon_{y y} & \gamma_{x y}
\end{array}\right\}  \tag{6.225}\\
\mathbf{T}^{T} & =\left\{\begin{array}{ll}
T_{x} & T_{y}
\end{array}\right\}
\end{align*}
$$

and

$$
\mathbf{D}=\frac{E}{1-v^{2}}\left[\begin{array}{ccc}
1 & v & 0  \tag{6.226}\\
v & 1 & 0 \\
0 & 0 & \frac{(1-v)}{2}
\end{array}\right]
$$

with

$$
\begin{equation*}
\varepsilon_{z z}=-\frac{v}{E}\left(\sigma_{x x}+\sigma_{y y}\right) \tag{6.227}
\end{equation*}
$$

The initial strains arising from $\Delta T$, the temperature change with respect to the reference state, can be expressed as

$$
\varepsilon^{* T}=\left[\begin{array}{lll}
\alpha \Delta T & \alpha \Delta T & 0 \tag{6.228}
\end{array}\right]
$$

### 6.2.2.2 Plane Strain Conditions

A state of plane strain exists for a cylindrical component that is either "long" or fully constrained in the length direction under the action of only uniform lateral external loads (two examples are shown in Fig. 6.27). Because the ends of the cylindrical component are prevented from deforming in the thickness direction, it is assumed that the displacement component $u_{z}$ vanishes at every cross section of the body. The uniform loading and crosssectional geometry eliminates any variation in the length direction, leading to $\partial() / \partial z=0$. Also, planes perpendicular to the thickness direction before


Fig. 6.27 Long bodies with in-plane loading; suitable for plane strain idealization.
deformation remain perpendicular to the thickness direction after deformation. These assumptions result in zero transverse shear strains, $\gamma_{x z}=\gamma_{y z}=0$. Under these assumptions, the displacement, u, stress, $\boldsymbol{\sigma}$, strain, $\boldsymbol{\varepsilon}$, and traction, $\mathbf{T}$, vectors, and material property matrix, $\mathbf{D}$, reduce to

$$
\begin{align*}
& \mathbf{u}^{T}=\left\{\begin{array}{ll}
u_{x} & u_{y}
\end{array}\right\} \\
& \boldsymbol{\sigma}^{T}=\left\{\begin{array}{lll}
\sigma_{x x} & \sigma_{y y} & \sigma_{x y}
\end{array}\right\}  \tag{6.229}\\
& \boldsymbol{\varepsilon}^{T}=\left\{\begin{array}{lll}
\varepsilon_{x x} & \varepsilon_{y y} & \gamma_{x y}
\end{array}\right\} \\
& \mathbf{T}^{T}=\left\{\begin{array}{ll}
T_{x} & T_{y}
\end{array}\right\}
\end{align*}
$$

and

$$
\mathbf{D}=\frac{E}{(1+v)(1-2 v)}\left[\begin{array}{ccc}
1-v & v & 0  \tag{6.230}\\
v & 1-v & 0 \\
0 & 0 & \frac{(1-2 v)}{2}
\end{array}\right]
$$

The initial strain vector due to this temperature change can be expressed as

$$
\varepsilon^{* T}=\left[\begin{array}{lll}
(1+v) \alpha \Delta T & (1+v) \alpha \Delta T & 0 \tag{6.231}
\end{array}\right]
$$

where $\Delta T$ is the temperature change with respect to a reference state.
The material property matrices for both plane stress and strain conditions have the same form, and it is convenient to present it in the form

$$
\mathbf{D}=\left[\begin{array}{ccc}
D_{1} & D_{1} D_{2} & 0  \tag{6.232}\\
D_{1} D_{2} & D_{1} & 0 \\
0 & 0 & D_{12}
\end{array}\right]
$$

where

$$
\begin{equation*}
D_{12}=\frac{D_{1}\left(1-D_{2}\right)}{2} \tag{6.233}
\end{equation*}
$$

with $D_{1}=E /(1-v)^{2}$ and $D_{2}=v$ for plane stress, and $D_{1}=E(1-v) /$ $(1+v)(1-2 v)$ and $D_{2}=v /(1-v)$ for plane strain.

### 6.2.2.3 Finite Element Equations with Linear Triangular Elements

The displacement components $u_{x}$ and $u_{y}$ within a triangular element can be approximated as

$$
\begin{align*}
& u_{x}^{(e)}=\tilde{u}_{x}^{(e)}=N_{1}^{(e)} u_{x_{1}}^{(e)}+N_{2}^{(e)} u_{x_{2}}^{(e)}+N_{3}^{(e)} u_{x_{3}}^{(e)} \\
& u_{y}^{(e)}=\tilde{u}_{y}^{(e)}=N_{1}^{(e)} u_{y_{1}}^{(e)}+N_{2}^{(e)} u_{y_{2}}^{(e)}+N_{3}^{(e)} u_{y_{3}}^{(e)} \tag{6.234}
\end{align*}
$$

in which $N_{1}^{(e)}, N_{2}^{(e)}$, and $N_{3}^{(e)}$ are the linear shape functions and $\left(u_{x_{1}}^{(e)}, u_{y_{1}}^{(e)}\right),\left(u_{x_{2}}^{(e)}, u_{y_{2}}^{(e)}\right)$, and ( $\left.u_{x_{3}}^{(e)}, u_{y_{3}}^{(e)}\right)$ are the nodal unknowns (degrees of freedom) associated with first, second, and third nodes, respectively. An example of a triangular element with its nodal degrees of freedom and local nodal numbering is shown in Fig. 6.28. In matrix form, the approximate displacement components become

$$
\begin{equation*}
\tilde{\mathbf{u}}^{(e)}=\mathbf{N}^{(e) T} \mathbf{U}^{(e)} \tag{6.235}
\end{equation*}
$$



Fig. 6.28 Typical linear triangular element with nodal degrees of freedom.
in which

$$
\tilde{\mathbf{u}}^{(e) T}=\left\{\begin{array}{ll}
\tilde{u}_{x}^{(e)} & \tilde{u}_{y}^{(e)} \tag{6.236}
\end{array}\right\}
$$

and

$$
\mathbf{N}^{(e) T}=\left[\begin{array}{cccccc}
N_{1} & 0 & N_{2} & 0 & N_{3} & 0  \tag{6.237}\\
0 & N_{1} & 0 & N_{2} & 0 & N_{3}
\end{array}\right]
$$

and

$$
\mathbf{U}^{(e) T}=\left\{\begin{array}{llllll}
u_{x_{1}}^{(e)} & u_{y_{1}}^{(e)} & u_{x_{2}}^{(e)} & u_{y_{2}}^{(e)} & u_{x_{3}}^{(e)} & u_{y_{3}}^{(e)} \tag{6.238}
\end{array}\right\}
$$

The element shape matrix, $\mathbf{B}^{(e)}$, becomes

$$
\mathbf{B}^{(e)}=\left[\begin{array}{cccccc}
\frac{\partial N_{1}}{\partial x} & 0 & \frac{\partial N_{2}}{\partial x} & 0 & \frac{\partial N_{3}}{\partial x} & 0  \tag{6.239}\\
0 & \frac{\partial N_{1}}{\partial y} & 0 & \frac{\partial N_{2}}{\partial y} & 0 & \frac{\partial N_{3}}{\partial y} \\
\frac{\partial N_{1}}{\partial y} & \frac{\partial N_{1}}{\partial x} & \frac{\partial N_{2}}{\partial y} & \frac{\partial N_{2}}{\partial x} & \frac{\partial N_{3}}{\partial y} & \frac{\partial N_{3}}{\partial x}
\end{array}\right]
$$

Substituting for the derivatives of the shape functions, this matrix simplifies to

$$
\mathbf{B}^{(e)}=\frac{1}{2 \Delta^{(e)}}\left[\begin{array}{cccccc}
y_{23}^{(e)} & 0 & y_{31}^{(e)} & 0 & y_{12}^{(e)} & 0  \tag{6.240}\\
0 & x_{32}^{(e)} & 0 & x_{13}^{(e)} & 0 & x_{21}^{(e)} \\
x_{32}^{(e)} & y_{23}^{(e)} & x_{13}^{(e)} & y_{31}^{(e)} & x_{21}^{(e)} & y_{12}^{(e)}
\end{array}\right]
$$

Both the element shape and material property matrices are independent of the spatial coordinates, $x$ and $y$, thus leading to the evaluation of the element stiffness matrix, $\mathbf{k}^{(e)}$, as

$$
\begin{equation*}
\mathbf{k}^{(e)}=\mathbf{B}^{(e) T} \mathbf{D} \mathbf{B}^{(e)} V^{(e)} \tag{6.241}
\end{equation*}
$$

where $V^{(e)}=t \Delta^{(e)}$, with element area $\Delta^{(e)}$ and constant thickness $t$. The evaluation of the load vectors, $\mathbf{p}_{\mathbf{b}}^{(e)}$ and $\mathbf{p}_{\mathrm{T}}^{(e)}$, arising from the body forces and surface tractions (forces), respectively, involve integrals of the form

$$
\begin{equation*}
\int d x d y, \quad \int x d x d y, \quad \int y d x d y \tag{6.242}
\end{equation*}
$$

By choosing the centroid of the triangle as the origin of the $(x, y)$ coordinate system, the integrals involving either $x$ or $y$ in the integrand vanish. The load vector arising from the body forces can be obtained from

$$
\left.\mathbf{p}_{\mathrm{b}}^{(e)}=\int_{V^{(e)}}\left[\begin{array}{cc}
N_{1} & 0  \tag{6.243}\\
0 & N_{1} \\
N_{2} & 0 \\
0 & N_{2} \\
N_{3} & 0 \\
0 & N_{3}
\end{array}\right]\left\{b_{x}\right\} b_{y}\right\} d V=\left\{\begin{array}{l}
N_{1} b_{x} \\
N_{1} b_{y} \\
N_{2} b_{x} \\
N_{2} b_{y} \\
N_{3} b_{x} \\
N_{3} b_{y}
\end{array}\right\} d V
$$

reducing to

$$
\mathbf{p}_{\mathrm{b}}^{(e) T}=\frac{t \Delta^{(e)}}{3}\left[\begin{array}{llllll}
b_{x} & b_{y} & b_{x} & b_{y} & b_{x} & b_{y} \tag{6.244}
\end{array}\right]
$$

in which $b_{x}$ and $b_{y}$ are the components of the body force vector.
The evaluation of the element load vector due to the applied traction forces (distributed loads as shown in Fig. 6.29) requires their explicit variation along the edges of the element. For an element of constant thickness subjected to uniform load of $T_{x}$ in the $x$-direction along its 1-2 edge, the vector $\mathbf{p}_{\mathbf{T}}^{(e)}$ can be written as


Fig. 6.29 Surface force along side 1-2 of the triangular element.

$$
\mathbf{p}_{\mathbf{T}}^{(e)}=t \int\left[\begin{array}{cc}
N_{1} & 0  \tag{6.245}\\
0 & N_{1} \\
N_{2} & 0 \\
0 & N_{2} \\
0 & 0 \\
0 & 0
\end{array}\right]\left\{\begin{array}{c}
T_{x} \\
0
\end{array}\right\} d l=t \iint_{L_{1-2}}\left\{\begin{array}{c}
N_{1} T_{x} \\
0 \\
N_{2} T_{x} \\
0 \\
0 \\
0
\end{array}\right\} d l
$$

in which $N_{3}=0$ along the 1-2 edge and $L_{1-2}$ is the length of the 1-2 edge . Since $N_{1}$ and $N_{2}$ vary linearly along the 1-2 edge, they can be expressed in terms of the natural coordinates, $\xi_{1}$ and $\xi_{2}$, as derived in Chap. 3

$$
\begin{equation*}
N_{1}=\xi_{1}=\frac{x_{2}^{(e)}-x}{x_{2}^{(e)}-x_{1}^{(e)}} \quad \text { and } \quad N_{2}=\xi_{2}=\frac{x-x_{1}^{(e)}}{x_{2}^{(e)}-x_{1}^{(e)}} \tag{6.246}
\end{equation*}
$$

The integrals in the expression for $\mathbf{p}_{\mathrm{T}}^{(e)}$ are evaluated as

$$
\begin{align*}
& \int_{L_{1-2}} N_{1} T_{x} d l=\int_{0}^{1} \xi_{1} T_{x} L_{1-2} d \xi_{1}=\frac{T_{x} L_{1-2}}{2}  \tag{6.247}\\
& \int_{L_{1-2}} N_{2} T_{x} d l=\int_{0}^{1} \xi_{2} T_{x} L_{1-2} d \xi_{2}=\frac{T_{x} L_{1-2}}{2}
\end{align*}
$$

Thus, the load vector, $\mathbf{p}_{\mathbf{T}}^{(e)}$, takes the form

$$
\mathbf{p}_{\mathrm{T}}^{(e) T}=t \frac{T_{x} L_{1-2}}{2}\left[\begin{array}{llllll}
1 & 0 & 1 & 0 & 0 & 0 \tag{6.248}
\end{array}\right]
$$

as illustrated in Fig. 6.30.
Note that this result corresponds to equivalent point forces acting at the first and second nodes. The element load vectors arising from the initial strains and stresses can be written as

$$
\begin{align*}
& \mathbf{p}_{\varepsilon^{*}}^{(e)}=\mathbf{B}^{(e) T} \mathbf{D} \boldsymbol{\varepsilon}^{*} V^{(e)}  \tag{6.249}\\
& \mathbf{p}_{\sigma^{*}}^{(e)}=\mathbf{B}^{(e) T} \boldsymbol{\sigma}^{*} V^{(e)}
\end{align*}
$$

### 6.2.2.4 Example of a Plane Stress Analysis with Linear Triangular Elements

### 6.2.2.4.1 Derivation of a System of Equations and Its Solution

Using linear triangular elements, determine the nodal displacements and the element stresses in a thin plate subjected to displacement constraints and surface tractions as shown in Fig. 6.31. Also, the plate is exposed to a temperature change of $10{ }^{\circ} \mathrm{C}$ from the reference temperature. The plate thickness is 0.5 cm and the Young's modulus, $E$, and the Poisson's ratio, $v$, are $15 \times 10^{6} \mathrm{~N} / \mathrm{cm}^{2}$ and 0.25 , respectively. The coefficient of thermal expansion is $6 \times 10^{-6} /{ }^{\circ} \mathrm{C}$.

In order to illustrate the solution method, the plate is discretized into two triangular elements, as shown in Fig. 6.32.

The global coordinates of each node are specified by ( $x_{p}, y_{p}$ ), with $p=1,2,3,4$, and are presented in Table 6.7.


Fig. 6.30 Equivalent nodal forces for the surface force along side 1-2 of the triangular element.


Fig. 6.31 Geometry and loading of the problem.



Fig. 6.32 Global and local numbering of nodes and elements.

Table 6.7 Global nodal coordinates.

| Global Node <br> Number | Nodal <br> Coordinates | Nodal <br> Unknowns |
| :---: | :---: | :---: |
| 1 | $(0,0)$ | $u_{x_{1}}, u_{y_{1}}$ |
| 2 | $(2,0)$ | $u_{x_{2}}, u_{y_{2}}$ |
| 3 | $(2,3)$ | $u_{x_{3}}, u_{y_{3}}$ |
| 4 | $(1,3)$ | $u_{x_{4}}, u_{y_{4}}$ |

The global unknown nodal displacement vector is given by

$$
\mathbf{U}^{T}=\left\{\begin{array}{llllllll}
u_{x_{1}} & u_{y_{1}} & u_{x_{2}} & u_{y_{2}} & u_{x_{3}} & u_{y_{3}} & u_{x_{4}} & u_{y_{4}} \tag{6.250}
\end{array}\right\}
$$

Considering the correspondence between the local and global node numbering schemes, the elements are defined (connected) as shown in Table 6.8.

The areas of each element are calculated to be

$$
\begin{equation*}
\Delta^{(1)}=3 \mathrm{~cm}^{2} \quad \text { and } \quad \Delta^{(2)}=3 / 2 \mathrm{~cm}^{2} \tag{6.251}
\end{equation*}
$$

Under plane stress assumptions, the material property matrix becomes

$$
\mathbf{D}=10^{6}\left[\begin{array}{ccc}
16 & 4 & 0  \tag{6.252}\\
4 & 16 & 0 \\
0 & 0 & 6
\end{array}\right] \mathrm{N} / \mathrm{cm}^{2}
$$

The initial strains arising from the temperature change is written as

$$
\varepsilon^{* T}=10^{-6}\left[\begin{array}{lll}
60 & 60 & 0 \tag{6.253}
\end{array}\right]
$$

The element load vectors arising from the applied tractions are

$$
\begin{align*}
& \mathbf{p}_{\mathbf{T}}^{(1) T}=t \frac{T_{x} L_{1-4}}{2}\left[\begin{array}{llllll}
1 & 0 & 0 & 0 & 1 & 0
\end{array}\right]  \tag{6.254}\\
& \mathbf{p}_{\mathbf{T}}^{(2) T}=t \frac{T_{y} L_{3-4}}{2}\left[\begin{array}{llllll}
0 & 0 & 0 & 1 & 0 & 1
\end{array}\right] \tag{6.255}
\end{align*}
$$

With the specified values of the thickness and the distributed loads, these element load vectors become

$$
\mathbf{p}_{\mathbf{T}}^{(1) T}=300 \sqrt{10}\left[\begin{array}{llllll}
1 & 0 & 0 & 0 & 1 & 0 \tag{6.256}
\end{array}\right] N
$$

and

$$
\mathbf{p}_{\mathbf{T}}^{(2) T}=-150\left[\begin{array}{llllll}
0 & 0 & 0 & 1 & 0 & 1 \tag{6.257}
\end{array}\right] N
$$

Table 6.8 Element connectivity.

| Element <br> Number <br> $(e)$ | Node 1 | Node 2 | Node 3 |
| :---: | :---: | :---: | :---: |
| 1 | 1 | 2 | 4 |
| 2 | 2 | 3 | 4 |

For the first element, $e=1$, the components of the element shape matrix $\mathbf{B}^{(1)}$ are computed as

$$
\begin{array}{ll}
y_{23}^{(1)}=y_{2}^{(1)}-y_{3}^{(1)}=y_{2}-y_{4}=-3, & x_{32}^{(1)}=x_{3}^{(1)}-x_{2}^{(1)}=x_{4}-x_{2}=-1 \\
y_{31}^{(1)}=y_{3}^{(1)}-y_{1}^{(1)}=y_{4}-y_{1}=3, & x_{13}^{(1)}=x_{1}^{(1)}-x_{3}^{(1)}=x_{1}-x_{4}=-1  \tag{6.258}\\
y_{12}^{(1)}=y_{1}^{(1)}-y_{2}^{(1)}=y_{1}-y_{2}=0, & x_{21}^{(1)}=x_{2}^{(1)}-x_{1}^{(1)}=x_{2}-x_{1}=2
\end{array}
$$

leading to

$$
\mathbf{B}^{(1)}=\frac{1}{6}\left[\begin{array}{cccccc}
-3 & 0 & 3 & 0 & 0 & 0  \tag{6.259}\\
0 & -1 & 0 & -1 & 0 & 2 \\
-1 & -3 & -1 & 3 & 2 & 0
\end{array}\right]
$$

For the second element, $e=2$, the components of the element shape matrix $\mathbf{B}^{(2)}$ are computed as

$$
\begin{array}{ll}
y_{23}^{(2)}=y_{2}^{(2)}-y_{3}^{(2)}=y_{3}-y_{4}=0, & x_{32}^{(2)}=x_{3}^{(2)}-x_{2}^{(2)}=x_{4}-x_{3}=-1 \\
y_{31}^{(2)}=y_{3}^{(2)}-y_{1}^{(2)}=y_{4}-y_{2}=3, & x_{13}^{(2)}=x_{1}^{(2)}-x_{3}^{(2)}=x_{2}-x_{4}=1  \tag{6.260}\\
y_{12}^{(2)}=y_{1}^{(2)}-y_{2}^{(2)}=y_{2}-y_{3}=-3, & x_{21}^{(2)}=x_{2}^{(2)}-x_{1}^{(2)}=x_{3}-x_{2}=0
\end{array}
$$

leading to

$$
\mathbf{B}^{(2)}=\frac{1}{3}\left[\begin{array}{cccccc}
0 & 0 & 3 & 0 & -3 & 0  \tag{6.261}\\
0 & -1 & 0 & 1 & 0 & 0 \\
-1 & 0 & 1 & 3 & 0 & -3
\end{array}\right]
$$

The evaluation of the stiffness matrices, $\mathbf{k}^{(1)}$ and $\mathbf{k}^{(2)}$, requires the products of $\mathbf{B}^{(1) T} \mathbf{D}$ and $\mathbf{B}^{(2) T} \mathbf{D}$. Also, these products appear in the evaluation of the element load vectors arising from the temperature change. Therefore,

$$
\mathbf{B}^{(1) T} \mathbf{D}=\frac{10^{6}}{6}\left[\begin{array}{ccc}
-48 & -12 & -6  \tag{6.262}\\
-4 & -16 & -18 \\
48 & 12 & -6 \\
-4 & -16 & 18 \\
0 & 0 & 12 \\
8 & 32 & 0
\end{array}\right]
$$

$$
\mathbf{B}^{(2) T} \mathbf{D}=\frac{10^{6}}{3}\left[\begin{array}{ccc}
0 & 0 & -6  \tag{6.263}\\
-4 & -16 & 0 \\
48 & 12 & 6 \\
4 & 16 & 18 \\
-48 & -12 & 0 \\
0 & 0 & -18
\end{array}\right]
$$

The element stiffness matrices become

$$
\mathbf{k}^{(1)}=\frac{10^{6}}{12}\left[\right]
$$

and

$$
\left.\mathbf{k}^{(2)}=\frac{10^{6}}{12}\left[\begin{array}{cccccc}
i=2 & j=3 & k=4  \tag{6.265}\\
0 & 0 & -6 & -18 & 0 & 18 \\
0 & 16 & -12 & -16 & 12 & 0 \\
-6 & -12 & 150 & 30 & -144 & -18 \\
-18 & -16 & 30 & 70 & -12 & -54 \\
0 & 12 & -144 & -12 & 144 & 0 \\
18 & 0 & -18 & -54 & 0 & 54
\end{array}\right], ~ \begin{array}{cc}
6
\end{array}\right]
$$

The boxed numbers above each column pair indicate the nodal order of degrees of freedom in each element stiffness matrix.

The thermal load vectors associated with each element are obtained as

$$
\mathbf{p}_{\varepsilon^{*}}^{(1)}=\left\{\begin{array}{c}
-900  \tag{6.266}\\
-300 \\
900 \\
-300 \\
0 \\
600
\end{array}\right\} N \quad \text { and } \quad \mathbf{p}_{\varepsilon^{*}}^{(2)}=\left\{\begin{array}{c}
0 \\
-300 \\
900 \\
300 \\
-900 \\
0
\end{array}\right\} N
$$

Rewriting the element stiffness matrices and the load vectors, in the expanded order and rearranged form according to the increasing nodal degrees of freedom of the global stiffness matrix, $\mathbf{K}$ yields

Associated with the first element:

$$
\begin{align*}
& \begin{array}{lll}
1 & 2 & 3 \\
\hline
\end{array} \\
& \mathbf{k}^{(1)}=\frac{10^{6}}{12}\left[\begin{array}{cccccccc}
75 & 15 & -69 & -3 & 0 & 0 & -6 & -12 \\
15 & 35 & 3 & -19 & 0 & 0 & -18 & -16 \\
-69 & 3 & 75 & -15 & 0 & 0 & -6 & 12 \\
-3 & -19 & -15 & 35 & 0 & 0 & 18 & -16 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
-6 & -18 & -6 & 18 & 0 & 0 & 12 & 0 \\
-12 & -16 & 12 & -16 & 0 & 0 & 0 & 32
\end{array}\right]  \tag{6.267}\\
& \mathbf{p}_{\mathbf{T}}^{(1)}=300 \sqrt{10}\left\{\begin{array}{l}
1 \\
0 \\
0 \\
0 \\
0 \\
0 \\
1 \\
0
\end{array}\right\} N \text { and } \quad \mathbf{p}_{\varepsilon^{*}}^{(1)}=\left\{\begin{array}{c}
-900 \\
-300 \\
900 \\
-300 \\
0 \\
0 \\
0 \\
600
\end{array}\right\} N \tag{6.268}
\end{align*}
$$

Associated with the second element:

$$
\begin{equation*}
\mathbf{k}^{(2)}=\frac{10^{6}}{12}\left[\right] \tag{6.269}
\end{equation*}
$$

$$
\mathbf{p}_{\mathbf{T}}^{(2)}=-150\left\{\begin{array}{l}
0  \tag{6.270}\\
0 \\
0 \\
0 \\
0 \\
1 \\
0 \\
1
\end{array}\right\} N \quad \text { and } \quad \mathbf{p}_{\varepsilon^{*}}^{(2)}=\left\{\begin{array}{c}
0 \\
0 \\
0 \\
-300 \\
900 \\
300 \\
-900 \\
0
\end{array}\right\} N
$$

Summation of the element stiffness matrices

$$
\begin{equation*}
\mathbf{K}=\sum_{e=1}^{E} \mathbf{k}^{(e)} \tag{6.271}
\end{equation*}
$$

and load vectors

$$
\begin{equation*}
\mathbf{P}=\sum_{e=1}^{E=2}\left(\mathbf{p}_{\mathbf{T}}^{(e)}+\mathbf{p}_{\varepsilon^{*}}^{(e)}\right) \tag{6.272}
\end{equation*}
$$

results in the global stiffness matrix and the global load vector as

$$
\mathbf{K}=\frac{10^{6}}{12}\left[\begin{array}{cccc}
75 & 15 & -69 & -3 \\
15 & 35 & 3 & -19 \\
-69 & 3 & (75+6) & -15 \\
-3 & -19 & -15 & (35+6) \\
0 & 0 & -6 & -12 \\
0 & 0 & -18 & -16 \\
-6 & -18 & -6 & (18+12) \\
-12 & -16 & (12+18) & -16 \\
0 & 0 & -6 & -12 \\
0 & 0 & -18 & -16 \\
-6 & -18 & -6 & (12+18) \\
-12 & -16 & (18+12) & -16 \\
150 & 30 & -144 & -18 \\
30 & 64 & -12 & -48 \\
-144 & -12 & (12+144) & 0 \\
-18 & -48 & 0 & (32+48)
\end{array}\right]
$$

and

$$
\mathbf{P}=\left\{\begin{array}{c}
(300 \sqrt{10}-900)  \tag{6.274}\\
-300 \\
900 \\
(-300-300) \\
900 \\
(-150+300) \\
(300 \sqrt{10}-900) \\
-150+600
\end{array}\right\} N
$$

The final form of the global system of equations becomes

$$
\begin{align*}
& \frac{10^{6}}{12}\left[\begin{array}{cccccccc}
75 & 15 & -69 & -3 & 0 & 0 & -6 & -12 \\
15 & 35 & 3 & -19 & 0 & 0 & -18 & -16 \\
-69 & 3 & (75+6) & -15 & -6 & -18 & -6 & (12+18) \\
-3 & -19 & -15 & (35+6) & -12 & -16 & (18+12) & -16 \\
0 & 0 & -6 & -12 & 150 & 30 & -144 & -18 \\
0 & 0 & -18 & -16 & 30 & 70 & -12 & -54 \\
-6 & -18 & -6 & (18+12) & -144 & -12 & (12+144) & 0 \\
-12 & -16 & (12+18) & -16 & -18 & -54 & 0 & (32+54)
\end{array}\right] \\
& \times\left\{\begin{array}{l}
u_{x_{1}} \\
u_{y_{1}} \\
u_{x_{2}} \\
u_{y_{2}} \\
u_{x_{3}} \\
u_{y_{3}} \\
u_{x_{4}} \\
u_{y_{4}}
\end{array}\right\}=\left\{\begin{array}{c}
(300 \sqrt{10}-900) \\
-300 \\
900 \\
(-300-300) \\
900 \\
(-150+300) \\
(300 \sqrt{10}-900) \\
-150+600
\end{array}\right\} \tag{6.275}
\end{align*}
$$

Applying the prescribed values of the displacement components leads to

$$
\begin{align*}
\frac{10^{6}}{12}\left[\begin{array}{cccccccc}
75 & 15 & -69 & -3 & 0 & 0 & -6 & -12 \\
15 & 35 & 3 & -19 & 0 & 0 & -18 & -16 \\
-69 & 3 & (75+6) & -15 & -6 & -18 & -6 & (12+18) \\
-3 & -19 & -15 & (35+6) & -12 & -16 & (18+12) & -16 \\
0 & 0 & -6 & -12 & 150 & 30 & -144 & -18 \\
0 & 0 & -18 & -16 & 30 & 70 & -12 & -54 \\
-6 & -18 & -6 & (18+12) & -144 & -12 & (12+144) & 0 \\
-12 & -16 & (12+18) & -16 & -18 & -54 & 0 & (32+54)
\end{array}\right] \\
\times\left\{\begin{array}{c}
u_{x_{1}} \\
0 \\
0 \\
0 \\
0 \\
u_{y_{3}} \\
u_{x_{4}} \\
u_{y_{4}}
\end{array}\right\}=\left\{\begin{array}{c}
(300 \sqrt{10}-900) \\
-300 \\
900 \\
(-300-300) \\
900 \\
(-150+300) \\
(300 \sqrt{10}-900) \\
-150+600
\end{array}\right\} \tag{6.276}
\end{align*}
$$

Eliminating the rows and columns corresponding to zero displacement components simplifies the global system of equations to

$$
\begin{align*}
& \frac{10^{6}}{12}\left[\begin{array}{cccc}
75 & 0 & -6 & -12 \\
0 & 70 & -12 & -54 \\
-6 & -12 & (12+144) & 0 \\
-12 & -54 & 0 & (32+54)
\end{array}\right]\left\{\begin{array}{l}
u_{x_{1}} \\
u_{y_{3}} \\
u_{x_{4}} \\
u_{y_{4}}
\end{array}\right\}  \tag{6.277}\\
& =\left\{\begin{array}{c}
(300 \sqrt{10}-900) \\
(-150+300) \\
(300 \sqrt{10}-900) \\
(-150+600)
\end{array}\right\}
\end{align*}
$$

The solution to this system of equations results in the values for the unknown displacement components as

$$
\left\{\begin{array}{l}
u_{x_{1}}  \tag{6.278}\\
u_{y_{3}} \\
u_{x_{4}} \\
u_{y_{4}}
\end{array}\right\}=\left\{\begin{array}{c}
0.0000357839 \\
0.000157003 \\
0.0000171983 \\
0.000166367
\end{array}\right\} \mathrm{cm}
$$

### 6.2.2.4.2 ANSYS Solution

The nodal displacements of the plate subjected to uniform temperature can also be obtained using ANSYS. The solution procedure is outlined as follows:

## MODEL GENERATION

- Specify the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Structural Solid from the left list and Quad 4node 42 from the right list; click on $\boldsymbol{O K}$.
- Click on Options.
- In order to specify the 2-D idealization as plane stress with thickness, in the newly appeared dialog box pull down the menu for Element behavior K3 and select Plane strs w/thk; click on OK (Fig. 6.33).
- Click on Close.
- Specify real constants ( R command) using the following menu path:

Main Menu > Preprocessor > Real Constants > Add/Edit/Delete

- Click on Add.
- Click on $\boldsymbol{O K}$.
- Enter 5 e-3 for Thickness THK; click on $\boldsymbol{O K}$.
- Click on Close.

Specify material properties (MP command) using the following menu path:

## Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring another dialog box.
- Enter $\mathbf{1 5 0 e 9}$ for $\boldsymbol{E X}$, and $\mathbf{0 . 2 5}$ for PRXY; click on OK.
- In the Define Material Model Behavior dialog box, in the right window, under Structural find Thermal Expansion, Secant Coefficient, and Isotropic, which will bring another dialog box (Fig. 6.34).
- Enter $\boldsymbol{6 e}-\mathbf{6}$ for $\boldsymbol{A P L X}$; click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:


Fig. 6.33 Specification of element options.


Fig. 6.34 Specification of material behavior.

- Create nodes ( $\mathbf{N}$ command) using the following menu path:

Main Menu > Preprocessor $>$ Modeling $>$ Create $>$ Nodes $>\operatorname{In}$ Active CS

- A total of 4 nodes will be created (Table 6.7).
- Referring to Table 6.7, enter $x$ - and $y$-coordinates of node 1 (be sure to convert the coordinates to meters), and Click on Apply. This action will keep the Create Nodes in Active Coordinate System dialog box open. If the Node number field is left blank, then ANSYS will assign the lowest available node number to the node that is being created.
- Repeat the same procedure for the nodes 2 through 4.
- After entering the $x$ - and $y$-coordinates of node 4 , click on $O K$ (instead of Apply).
- The nodes should appear in the Graphics Window, as shown in Fig. 6.35 .
- Create elements ( $\mathbf{E}$ command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Elements > Auto Numbered > Thru Nodes

- Pick Menu appears; refer to Fig. 6.36 to create elements by picking three nodes at a time and clicking on Apply in between.
- Observe the elements created after clicking on Apply in the Pick Menu.
- Repeat until the last element is created.
- Click on $O K$ when the last element is created.
- Review elements:
- Turn on element numbering using the following menu path: Utility Menu > PlotCtris > Numbering
- Select Element numbers from the first pull-down menu; click on $O K$.
- Plot elements (EPLOT command) using the following menu path:

Utility Menu > Plot > Elements

- Figure 6.36 shows the outcome of this action as it appears in the Graphics Window.
- Turn off element numbering and turn on node numbering using the following menu path:

Utility Menu > PlotCtris $>$ Numbering

- Place a checkmark by clicking on the empty box next to NODE Node numbers.
- Select No numbering from the first pull-down menu.
- Click on $O K$.
- Plot nodes (NPLOT command) using the following menu path:

Utility Menu > Plot > Nodes

- Figure 6.35 shows the outcome of this action as it appears in the Graphics Window.


Fig. 6.35 Generation of nodes.


Fig. 6.36 Generation of elements.

## SOLUTION

- Apply displacement boundary conditions ( $\mathbf{D}$ command) using the following menu path:


## Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; pick nodes 1 and 2 along the bottom horizontal boundary (Fig. 6.35) and click on $\boldsymbol{O K}$ on Pick Menu.
- Highlight $\boldsymbol{U Y}$ and enter $\boldsymbol{0}$ for VALUE; click on Apply.
- Pick Menu reappears; pick nodes 2 and 3 along the right vertical boundary (Fig. 6.35) and click on $O K$ on Pick Menu.
- Highlight $\boldsymbol{U X}$ and remove the highlight on $\boldsymbol{U Y}$; enter $\mathbf{0}$ for VALUE; click on $O K$.
- Apply force boundary conditions on nodes ( $\mathbf{F}$ command) using the following menu path:

Main Menu > Solution $>$ Define Loads $>$ Apply $>$ Structural $>$ Force/Moment $>$ On Nodes

- Pick Menu appears; pick nodes 1 and 4 along the slanted boundary; click on $\mathbf{O K}$.
- Enter $3 e 3{ }^{*}$ sqrt(0.1) for VALUE (Fig. 6.37).
- Click on Apply.
- Pick Menu reappears; pick nodes 4 and 3 along the top horizontal boundary; click on $\boldsymbol{O K}$.
- Pull down the menu for Direction of force/mom and select $\boldsymbol{F Y}$; Enter -150 for VALUE; click on $O K$.
- Apply thermal load (TUNIF command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Temperature > Uniform Temp

- Uniform Temperature dialog box appears; Enter 10 for Uniform temperature.
- Click on $O K$.


Fig. 6.37 Application of external loads.

- Obtain solution (SOLVE command) using the following menu path:


## Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status. If OK, close the Status Report Window and click on $\boldsymbol{O K}$ in Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

■ Review deformed shape (PLDISP command) using the following menu path:

Main Menu > General Postproc > Plot Results > Deformed Shape

- In the Plot Deformed Shape dialog box, choose the radio-button for Def + undef edge; click on $O K$.
- The deformed shape will appear in the Graphics Window, as shown in Fig. 6.38.
- Review displacement values (PRNSOL command) using the following menu path:

Main Menu > General Postproc > List Results > Nodal Solution

- Highlight DOF solution on the left list and All DOFs DOF on the right list; click on $\boldsymbol{O K}$.
- The list will appear in a new window, as shown in Fig. 6.39.


Fig. 6.38 Deformed configuration.


Fig. 6.39 List of nodal displacements.

### 6.2.2.5 Finite Element Equations with Linear Quadrilateral Isoparametric Elements

The displacement components $u_{x}$ and $u_{y}$ within a quadrilateral element can be approximated as

$$
\begin{align*}
& u_{x}^{(e)}=\tilde{u}_{x}^{(e)}=N_{1}^{(e)} u_{x_{1}}^{(e)}+N_{2}^{(e)} u_{x_{2}}^{(e)}+N_{3}^{(e)} u_{x_{3}}^{(e)}+N_{4}^{(e)} u_{x_{4}}^{(e)} \\
& u_{y}^{(e)}=\tilde{u}_{y}^{(e)}=N_{1}^{(e)} u_{y_{1}}^{(e)}+N_{2}^{(e)} u_{y_{2}}^{(e)}+N_{3}^{(e)} u_{y_{3}}^{(e)}+N_{4}^{(e)} u_{y_{4}}^{(e)} \tag{6.279}
\end{align*}
$$

in which $N_{1}^{(e)}, N_{2}^{(e)}, N_{3}^{(e)}$, and $N_{4}^{(e)}$ are the linear shape functions and $\left(u_{x_{1}}^{(e)}, u_{y_{1}}^{(e)}\right), \quad\left(u_{x_{2}}^{(e)}, u_{y_{2}}^{(e)}\right), \quad\left(u_{x_{3}}^{(e)}, u_{y_{3}}^{(e)}\right)$, and $\left(u_{x_{4}}^{(e)}, u_{y_{4}}^{(e)}\right)$ are the nodal unknowns (degrees of freedom) associated with first, second, third, and fourth nodes, respectively. The shape functions for the linear (straight-sided) quadrilateral shown in Fig. 6.40 are defined in terms of the centroidal or natural coordinates, $(\xi, \eta)$, as


Fig. 6.40 Variation of the natural coordinates in a typical quadrilateral element.

$$
\begin{equation*}
N_{p}=\frac{1}{4}\left(1+\xi \xi_{p}\right)\left(1+\eta \eta_{p}\right) \quad \text { with } \quad p=1,2,3,4 \tag{6.280}
\end{equation*}
$$

where $\xi_{p}$ and $\eta_{p}$ represent the coordinates of the corner nodes in the natural coordinate system, $\quad\left(\xi_{1}=-1, \eta_{1}=-1\right), \quad\left(\xi_{2}=1, \eta_{2}=-1\right)$, $\left(\xi_{3}=1, \eta_{3}=1\right)$, and ( $\xi_{4}=-1, \eta_{4}=1$ ).

In matrix form, the approximate displacement components become

$$
\begin{equation*}
\tilde{\mathbf{u}}^{(e)}=\mathbf{N}^{(e) T} \mathbf{U}^{(e)} \tag{6.281}
\end{equation*}
$$

in which

$$
\tilde{\mathbf{u}}^{(e) T}=\left\{\begin{array}{cc}
\tilde{u}_{x}^{(e)} & \tilde{u}_{y}^{(e)} \tag{6.282}
\end{array}\right\}
$$

and

$$
\mathbf{N}^{(e) T}=\left[\begin{array}{cccccccc}
N_{1} & 0 & N_{2} & 0 & N_{3} & 0 & N_{4} & 0  \tag{6.283}\\
0 & N_{1} & 0 & N_{2} & 0 & N_{3} & 0 & N_{4}
\end{array}\right]
$$

and

$$
\mathbf{U}^{(e) T}=\left\{\begin{array}{llllllll}
u_{x_{1}}^{(e)} & u_{y_{1}}^{(e)} & u_{x_{2}}^{(e)} & u_{y_{2}}^{(e)} & u_{x_{3}}^{(e)} & u_{y_{3}}^{(e)} & u_{x_{4}}^{(e)} & u_{y_{4}}^{(e)} \tag{6.284}
\end{array}\right\}
$$

The element shape matrix $\mathbf{B}^{(e)}$ can be expressed as

$$
\begin{equation*}
\mathbf{B}^{(e)}=\mathbf{L} \mathbf{N}^{(e) T} \tag{6.285}
\end{equation*}
$$

in which the differential operator matrix is

$$
\mathbf{L}=\left[\begin{array}{cc}
\frac{\partial}{\partial x} & 0  \tag{6.286}\\
0 & \frac{\partial}{\partial y} \\
\frac{\partial}{\partial x} & \frac{\partial}{\partial y}
\end{array}\right]
$$

The element shape matrix can be rewritten as

$$
\mathbf{B}^{(e)}=\left[\begin{array}{cccccccc}
\frac{\partial N_{1}}{\partial x} & 0 & \frac{\partial N_{2}}{\partial x} & 0 & \frac{\partial N_{3}}{\partial x} & 0 & \frac{\partial N_{4}}{\partial x} & 0  \tag{6.287}\\
0 & \frac{\partial N_{1}}{\partial y} & 0 & \frac{\partial N_{2}}{\partial y} & 0 & \frac{\partial N_{3}}{\partial y} & 0 & \frac{\partial N_{4}}{\partial y} \\
\frac{\partial N_{1}}{\partial y} & \frac{\partial N_{1}}{\partial x} & \frac{\partial N_{2}}{\partial y} & \frac{\partial N_{2}}{\partial x} & \frac{\partial N_{3}}{\partial y} & \frac{\partial N_{3}}{\partial x} & \frac{\partial N_{4}}{\partial y} & \frac{\partial N_{4}}{\partial x}
\end{array}\right]
$$

However, the shape functions are defined in terms of the centroidal or natural coordinates, $(\xi, \eta)$. Therefore, they cannot be differentiated directly with respect to the $x$ - and $y$-coordinates. In order to overcome this difficulty, the global coordinates are expressed in terms of the shape functions in the form

$$
\begin{equation*}
x=\sum_{p=1}^{4} N_{p}(\xi, \eta) x_{p} \quad \text { and } \quad y=\sum_{p=1}^{4} N_{p}(\xi, \eta) y_{p} \tag{6.288}
\end{equation*}
$$

With this transformation utilizing the same shape functions as those used for the displacement components, the concept of isoparametric element emerges, and the element is referred to as an isoparametric element.

The derivatives of the shape functions can be obtained as

$$
\begin{align*}
& \frac{\partial N_{p}}{\partial x}=\frac{\partial N_{p}}{\partial \xi} \frac{\partial \xi}{\partial x}+\frac{\partial N_{p}}{\partial \eta} \frac{\partial \eta}{\partial x} \\
& \frac{\partial N_{p}}{\partial y}=\frac{\partial N_{p}}{\partial \xi} \frac{\partial \xi}{\partial y}+\frac{\partial N_{p}}{\partial \eta} \frac{\partial \eta}{\partial y} \tag{6.289}
\end{align*} \quad \text { with } \quad p=1,2,3,4
$$

Application of the chain rule of differentiation yields

$$
\begin{align*}
& \frac{\partial N_{p}}{\partial \xi}=\frac{\partial N_{p}}{\partial x} \frac{\partial x}{\partial \xi}+\frac{\partial N_{p}}{\partial y} \frac{\partial y}{\partial \xi} \\
& \frac{\partial N_{p}}{\partial \eta}=\frac{\partial N_{p}}{\partial x} \frac{\partial x}{\partial \eta}+\frac{\partial N_{p}}{\partial y} \frac{\partial y}{\partial \eta} \tag{6.290}
\end{align*}
$$

In matrix form, it can be expressed as

$$
\left\{\begin{array}{c}
\frac{\partial}{\partial \xi}  \tag{6.291}\\
\frac{\partial}{\partial \eta}
\end{array}\right\}=\left[\begin{array}{ll}
\frac{\partial x}{\partial \xi} & \frac{\partial y}{\partial \xi} \\
\frac{\partial x}{\partial \eta} & \frac{\partial y}{\partial \eta}
\end{array}\right]\left\{\begin{array}{c}
\frac{\partial}{\partial x} \\
\frac{\partial}{\partial y}
\end{array}\right\} \quad \text { or }\left\{\begin{array}{c}
\frac{\partial}{\partial \xi} \\
\frac{\partial}{\partial \eta}
\end{array}\right\}=\mathbf{J}\left\{\begin{array}{c}
\frac{\partial}{\partial x} \\
\frac{\partial}{\partial y}
\end{array}\right\}
$$

where $\mathbf{J}$ is called the Jacobian matrix, whose inverse does not exist if there is excessive distortion of the element leading to the intersection of lines of constant $\xi$ and $\eta$ inside or on the element boundaries, as illustrated in Fig. 6.41. If the quadrilateral element is degenerated into a triangle by increasing an internal angle to $180^{\circ}$, then $\mathbf{J}$ is singular at that corner. It is possible to obtain the element stiffness because $\mathbf{J}$ is still unique at the Gaussian integration points. However, the stresses at that corner are indeterminate. A similar situation occurs when two adjacent corner nodes are made coincident to produce a triangular element. Therefore, any internal angle of each corner node should be less than $180^{\circ}$, and there is a loss of accuracy as the internal angle approaches $180^{\circ}$.

In the absence of excessive distortion, the transformation between the natural and global coordinates has a one-to-one correspondence and $\mathbf{J}^{-1}$ inverse exists. It can be expressed as

$$
\mathbf{J}^{-1}=\frac{1}{|\mathbf{J}|}\left[\begin{array}{cc}
\frac{\partial y}{\partial \eta} & -\frac{\partial y}{\partial \xi}  \tag{6.292}\\
-\frac{\partial x}{\partial \eta} & \frac{\partial x}{\partial \xi}
\end{array}\right]
$$

where the determinant of the Jacobian matrix is

$$
\begin{equation*}
|\mathbf{J}|=\frac{\partial x}{\partial \xi} \frac{\partial y}{\partial \eta}-\frac{\partial x}{\partial \eta} \frac{\partial y}{\partial \xi} \tag{6.293}
\end{equation*}
$$

in which


Fig. 6.41 Internal angle exceeding $180^{\circ}$.

$$
\begin{align*}
& \frac{\partial x}{\partial \xi}=\sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \xi} x_{p}=\frac{1}{4}\left\{-(1-\eta) x_{1}+(1-\eta) x_{2}+(1+\eta) x_{3}-(1+\eta) x_{4}\right\} \\
& \frac{\partial y}{\partial \xi}=\sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \xi} y_{p}=\frac{1}{4}\left\{-(1-\eta) y_{1}+(1-\eta) y_{2}+(1+\eta) y_{3}-(1+\eta) y_{4}\right\}  \tag{6.294}\\
& \frac{\partial x}{\partial \eta}=\sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \eta} x_{p}=\frac{1}{4}\left\{-(1-\xi) x_{1}-(1+\xi) x_{2}+(1+\xi) x_{3}+(1-\xi) x_{4}\right\} \\
& \frac{\partial y}{\partial \eta}=\sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \eta} y_{p}=\frac{1}{4}\left\{-(1-\xi) y_{1}-(1+\xi) y_{2}+(1+\xi) y_{3}+(1-\xi) y_{4}\right\}
\end{align*}
$$

Substituting for the derivatives and rearranging the terms permit the Jacobian to be rewritten in the form

$$
\mathbf{J}=\left[\begin{array}{cccc}
\frac{\partial N_{1}}{\partial \xi} & \frac{\partial N_{2}}{\partial \xi} & \frac{\partial N_{3}}{\partial \xi} & \frac{\partial N_{4}}{\partial \xi}  \tag{6.295}\\
\frac{\partial N_{1}}{\partial \eta} & \frac{\partial N_{2}}{\partial \eta} & \frac{\partial N_{3}}{\partial \eta} & \frac{\partial N_{4}}{\partial \eta}
\end{array}\right]\left[\begin{array}{ll}
x_{1} & y_{1} \\
x_{2} & y_{2} \\
x_{3} & y_{3} \\
x_{4} & y_{4}
\end{array}\right]
$$

or

$$
\mathbf{J}=\frac{1}{4}\left[\begin{array}{cccc}
-(1-\eta) & (1-\eta) & (1+\eta) & -(1+\eta)  \tag{6.296}\\
-(1-\xi) & -(1+\xi) & (1+\xi) & (1-\xi)
\end{array}\right]\left[\begin{array}{cc}
x_{1} & y_{1} \\
x_{2} & y_{2} \\
x_{3} & y_{3} \\
x_{4} & y_{4}
\end{array}\right]
$$

Its determinant can be expressed in the form

$$
|\mathbf{J}|=\frac{1}{8}\left[\begin{array}{llll}
x_{1} & x_{2} & x_{3} & x_{4}
\end{array}\right]\left[\begin{array}{cccc}
0 & 1-\eta & -\xi+\eta & -1+\xi  \tag{6.297}\\
-1+\eta & 0 & 1+\xi & -\xi-\eta \\
\xi-\eta & -1-\xi & 0 & 1+\eta \\
1-\xi & \xi+\eta & -1-\eta & 0
\end{array}\right]\left\{\begin{array}{l}
y_{1} \\
y_{2} \\
y_{3} \\
y_{4}
\end{array}\right\}
$$

In a concise form, the determinant can be also rewritten as

$$
\begin{equation*}
|\mathbf{J}|=\frac{1}{8}\left[\left(x_{31} y_{42}-x_{42} y_{31}\right)+\xi\left(x_{12} y_{23}-x_{23} y_{12}\right)+\eta\left(x_{41} y_{32}-x_{32} y_{41}\right)\right] \tag{6.298}
\end{equation*}
$$

where

$$
\begin{equation*}
x_{i j}=x_{i}-x_{j} \quad \text { and } \quad y_{i j}=y_{i}-y_{j} \tag{6.299}
\end{equation*}
$$

Determination of the inverse of the Jacobian matrix permits the expression for the derivatives of the natural coordinates in terms of the global coordinates, $x$ and $y$

$$
\left\{\begin{array}{l}
\frac{\partial \xi}{\partial x}  \tag{6.300a}\\
\frac{\partial \xi}{\partial y}
\end{array}\right\}=\frac{1}{|\mathbf{J}|}\left[\begin{array}{cc}
\frac{\partial y}{\partial \eta} & -\frac{\partial y}{\partial \xi} \\
-\frac{\partial x}{\partial \eta} & \frac{\partial x}{\partial \xi}
\end{array}\right]\left\{\begin{array}{l}
\frac{\partial \xi}{\partial \xi} \\
\frac{\partial \xi}{\partial \eta}
\end{array}\right\}
$$

and

$$
\left\{\begin{array}{l}
\frac{\partial \eta}{\partial x}  \tag{6.300b}\\
\frac{\partial \eta}{\partial y}
\end{array}\right\}=\frac{1}{|\mathbf{J}|}\left[\begin{array}{cc}
\frac{\partial y}{\partial \eta} & -\frac{\partial y}{\partial \xi} \\
-\frac{\partial x}{\partial \eta} & \frac{\partial x}{\partial \xi}
\end{array}\right]\left\{\begin{array}{l}
\frac{\partial \eta}{\partial \xi} \\
\frac{\partial \eta}{\partial \eta}
\end{array}\right\}
$$

By substituting for the derivatives of the global coordinates in terms of the natural coordinates, these expressions can be rewritten as

$$
\begin{array}{lll}
\frac{\partial \xi}{\partial x}=\frac{1}{|\mathbf{J}|} \sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \eta} y_{p} & \text { and } \quad \frac{\partial \xi}{\partial y}=-\frac{1}{|\mathbf{J}|} \sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \eta} x_{p} \\
\frac{\partial \eta}{\partial x}=-\frac{1}{|\mathbf{J}|} \sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \xi} y_{p} & \text { and } & \frac{\partial \eta}{\partial y}=\frac{1}{|\mathbf{J}|} \sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \xi} x_{p} \tag{6.301}
\end{array}
$$

Finally, the derivatives in the shape matrix becomes

$$
\begin{align*}
& \frac{\partial N_{p}}{\partial x}=\frac{1}{|\mathbf{J}|}\left\{\frac{\partial N_{p}}{\partial \xi} \sum_{q=1}^{4} \frac{\partial N_{q}}{\partial \eta} y_{q}-\frac{\partial N_{p}}{\partial \eta} \sum_{q=1}^{4} \frac{\partial N_{q}}{\partial \xi} y_{q}\right\} \\
& \frac{\partial N_{p}}{\partial y}=\frac{1}{|\mathbf{J}|}\left\{-\frac{\partial N_{p}}{\partial \xi} \sum_{q=1}^{4} \frac{\partial N_{q}}{\partial \eta} x_{q}+\frac{\partial N_{p}}{\partial \eta} \sum_{q=1}^{4} \frac{\partial N_{q}}{\partial \xi} x_{q}\right\} \tag{6.302}
\end{align*}
$$

These explicit expressions for the derivatives appearing in the element shape matrix permit the determination of the element stiffness matrix, $\mathbf{k}^{(e)}$, defined as

$$
\begin{equation*}
\mathbf{k}^{(e)}=\int_{V^{(e)}} \mathbf{B}^{(e) T} \mathbf{D} \mathbf{B}^{(e)} d V \tag{6.303}
\end{equation*}
$$

in which $V^{(e)}=t \Delta^{(e)}$, with $\Delta^{(e)}$ and $t$ representing the element area and constant element thickness. It can be rewritten in the form

$$
\begin{equation*}
\mathbf{k}^{(e)}=t \int_{\Delta^{(e)}} \mathbf{B}^{(e) T} \mathbf{D} \mathbf{B}^{(e)} d A \tag{6.304}
\end{equation*}
$$

The material property matrix $\mathbf{D}$ is usually independent of the spatial coordinates, $x$ and $y$, while the element shape matrix $\mathbf{B}^{(e)}$ requires differentiation of the shape functions with respect to $x$ and $y$. In order to overcome this difficulty, the integrals are evaluated over a square region in the natural coordinate system, with the transformation of coordinates given by

$$
\begin{equation*}
x=\sum_{p=1}^{4} N_{p}(\xi, \eta) x_{p} \quad \text { and } \quad y=\sum_{p=1}^{4} N_{p}(\xi, \eta) y_{p} \tag{6.305}
\end{equation*}
$$

With this transformation and utilizing the following relation

$$
\begin{equation*}
\iint_{A} d x d y=\int_{-1}^{1} \int_{-1}^{1}|\mathbf{J}| d \xi d \eta \tag{6.306}
\end{equation*}
$$

the element stiffness matrix, $\mathbf{k}^{(e)}$, can be rewritten as

$$
\begin{equation*}
\mathbf{k}^{(e)}=t \int_{-1-1}^{1} \int^{1} \mathbf{B}^{(e) T} \mathbf{D B}^{(e)}|\mathbf{J}| d \xi d \eta \tag{6.307}
\end{equation*}
$$

Due to the difficulty of obtaining analytical expression for the determinant and inverse of the Jacobian matrix, these integrals are evaluated numerically by the Gaussian integration technique. The element stiffness matrix can be evaluated numerically as

$$
\begin{equation*}
\mathbf{k}^{(e)}=t \sum_{p=1}^{P} \sum_{q=1}^{Q} w_{p} w_{q} \mathbf{B}^{(e)}\left(\xi_{p}, \eta_{q}\right)^{T} \mathbf{D} \mathbf{B}^{(e)}\left(\xi_{p}, \eta_{q}\right)\left|\mathbf{J}\left(\xi_{p}, \eta_{q}\right)\right| \tag{6.308}
\end{equation*}
$$

in which $w_{p}$ and $w_{q}$ are the weights and $\xi_{p}$ and $\eta_{q}$ are the integration points of the Gaussian integration technique explained in Sec. 3.6. For this quadrilateral isoparametric element, $P=2$ and $Q=2$ are sufficient for accurate integration.

For an element of constant thickness subjected to a uniform load of $T_{x}$ and $T_{y}$ in the $x$ - and $y$-directions, respectively, along its 1-2 edge, the vector $\mathbf{p}_{\mathbf{T}}^{(e)}$, arising from tractions can be written as

$$
\left.\mathbf{p}_{\mathbf{T}}^{(e)}=t \int\left[\begin{array}{cc}
N_{1} & 0  \tag{6.309}\\
0 & N_{1} \\
N_{2} & 0 \\
0 & N_{2} \\
N_{3} & 0 \\
0 & N_{3} \\
N_{4} & 0 \\
0 & N_{4}
\end{array}\right]\left\{\begin{array}{l}
T_{x} \\
T_{y}
\end{array}\right\} d l=t \int \begin{array}{c}
N_{1} T_{x} \\
N_{1} T_{y} \\
N_{2} T_{x} \\
N_{2} T_{y} \\
N_{3} T_{x} \\
N_{3} T_{y} \\
N_{4} T_{x} \\
N_{4} T_{y}
\end{array}\right\} d l
$$

Referring to Fig. 6.40, along the $1-2$ edge whose length is $L_{1-2}$, the coordinate $\eta$ has a constant value of -1 and $\xi$ varies between -1 and 1 , leading to

$$
\mathbf{p}_{\mathrm{T}}^{(e)}=t \frac{L_{1-2}}{2} \int_{-1}^{1}\left\{\begin{array}{l}
N_{1} T_{x}  \tag{6.310}\\
N_{1} T_{y} \\
N_{2} T_{x} \\
N_{2} T_{y} \\
N_{3} T_{x} \\
N_{3} T_{y} \\
N_{4} T_{x} \\
N_{4} T_{y}
\end{array}\right\} d \xi
$$

Along $\xi=-1$ to 1 and $\eta=-1$,

$$
\begin{align*}
& N_{1}=\frac{1}{4}(1-\xi)(1-\eta)=\frac{1}{2}(1-\xi) \\
& N_{2}=\frac{1}{4}(1+\xi)(1-\eta)=\frac{1}{2}(1+\xi)  \tag{6.311}\\
& N_{3}=\frac{1}{4}(1+\xi)(1+\eta)=0 \\
& N_{4}=\frac{1}{4}(1-\xi)(1+\eta)=0
\end{align*}
$$

The integrals in the expression for $\mathbf{p}_{\mathrm{T}}^{(e)}$ are evaluated as

$$
\begin{equation*}
t \frac{L_{1-2}}{2} \int_{-1}^{1} N_{1} T_{x} d \xi=t \frac{L_{1-2}}{4} \int_{-1}^{1}(1-\xi) T_{x} d \xi=t \frac{L_{1-2}}{2} T_{x} \tag{6.312}
\end{equation*}
$$

and

$$
\begin{equation*}
t \frac{L_{1-2}}{2} \int_{-1}^{1} N_{2} T_{y} d \xi=t \frac{L_{1-2}}{4} \int_{-1}^{1}(1+\xi) T_{y} d \xi=t \frac{L_{4-2}}{2} T_{y} \tag{6.313}
\end{equation*}
$$

Thus, the load vector, $\mathbf{p}_{\mathbf{T}}^{(e)}$, takes the form

$$
\mathbf{p}_{\mathrm{T}}^{(e) T}=t \frac{L_{1-2}}{2}\left[\begin{array}{llllllll}
T_{x} & T_{y} & T_{x} & T_{y} & 0 & 0 & 0 & 0 \tag{6.314}
\end{array}\right]
$$

Note that this result implies that the applied load is distributed equally at the first and second nodes of the 1-2 edge. This is a result of the linear variation of the shape function along the edges.

As carried out in the derivation of the element stiffness matrix, the load vectors due to body forces, initial strains, and initial stresses can be rewritten as

$$
\begin{gather*}
\mathbf{p}_{\mathbf{b}}^{(e)}=t \int_{-1}^{1} \int_{-1}^{1} \mathbf{N}^{(e)} \mathbf{b}|\mathbf{J}| d \xi d \eta  \tag{6.315}\\
\mathbf{p}_{\varepsilon^{*}}^{(e)}=t \int_{-1}^{1} \int_{-1}^{1} \mathbf{B}^{(e) T} \mathbf{D} \varepsilon^{*}|\mathbf{J}| d \xi d \eta  \tag{6.316}\\
\mathbf{p}_{\sigma^{*}}^{(e)}=t \int_{-1}^{1} \int_{-1}^{1} \mathbf{B}^{(e) T} \boldsymbol{\sigma}^{*}|\mathbf{J}| d \xi d \eta \tag{6.317}
\end{gather*}
$$

Application of the Gaussian integration technique leads to the evaluation of these load vectors in the form

$$
\begin{gather*}
\mathbf{p}_{\mathrm{b}}^{(e)}=t \sum_{p=1}^{P} \sum_{q=1}^{Q} w_{p} w_{q} \mathbf{N}^{(e)}\left(\xi_{p}, \eta_{q}\right) \mathbf{b}\left|\mathbf{J}\left(\xi_{p}, \eta_{q}\right)\right|  \tag{6.318}\\
\mathbf{p}_{\varepsilon^{*}}^{(e)}=t \sum_{p=1}^{P} \sum_{q=1}^{Q} w_{p} w_{q} \mathbf{B}^{(e)}\left(\xi_{p}, \eta_{q}\right)^{T} \mathbf{D} \varepsilon^{*}\left|\mathbf{J}\left(\xi_{p}, \eta_{q}\right)\right|  \tag{6.319}\\
\mathbf{p}_{\sigma^{*}}^{(e)}=t \sum_{p=1}^{P} \sum_{q=1}^{Q} w_{p} w_{q} \mathbf{B}^{(e)}\left(\xi_{p}, \eta_{q}\right)^{T} \mathbf{\sigma}^{*}\left|\mathbf{J}\left(\xi_{p}, \eta_{q}\right)\right| \tag{6.320}
\end{gather*}
$$

in which $w_{p}$ and $w_{q}$ are the weights and $\xi_{p}$ and $\eta_{q}$ are the integration points of the Gaussian integration technique.

### 6.2.2.6 Example of a Plane Stress Analysis with Linear Quadrilateral Isoparametric Elements

### 6.2.2.6.1 Derivation of a System of Equations and Its Solution

The previous example discussed in Sec. 6.2.2.4 is reconsidered to compute the nodal displacements and the element stresses. In order to illustrate the finite element solution method, the plate is discretized into one quadrilateral isoparametric element, as shown in Fig. 6.42.

The global coordinates of each node are specified by $\left(x_{p}, y_{p}\right)$, with $p=1,2,3,4$, and are tabulated in Table 6.9.

The global unknown nodal displacement vector is given by

$$
\mathbf{U}^{T}=\left\{\begin{array}{llllllll}
u_{x_{1}} & u_{y_{1}} & u_{x_{2}} & u_{y_{2}} & u_{x_{3}} & u_{y_{3}} & u_{x_{4}} & u_{y_{4}} \tag{6.321}
\end{array}\right\}
$$

Considering the correspondence between the local and global node numbering schemes the elements are defined in Table 6.10.


Fig. 6.42 Local numbering scheme of the FEM discretization with a quadrilateral element.

Table 6.9 Global nodal coordinates.

| Global <br> Node <br> Number | Nodal <br> Coordinates | Nodal <br> Unknowns |
| :---: | :---: | :---: |
| 1 | $\left(x_{1}=0, y_{1}=0\right)$ | $u_{x_{1}}, u_{y_{1}}$ |
| 2 | $\left(x_{2}=2, y_{2}=0\right)$ | $u_{x_{2}}, u_{y_{2}}$ |
| 3 | $\left(x_{3}=2, y_{3}=3\right)$ | $u_{x_{3}}, u_{y_{3}}$ |
| 4 | $\left(x_{4}=1, y_{4}=3\right)$ | $u_{x_{4}}, u_{y_{4}}$ |

Table 6.10 Element connectivity.

| Element <br> Number <br> $(e)$ | Node 1 | Node 2 | Node 3 | Node 4 |
| :---: | :---: | :---: | :---: | :---: |
| 1 | 1 | 2 | 3 | 4 |

For this element, $e=1$, the coefficients of the Jacobian matrix are determined from

$$
\begin{align*}
& \frac{\partial x}{\partial \xi}=\frac{1}{4}\left\{-(1-\eta)\left(x_{1}=0\right)+(1-\eta)\left(x_{2}=2\right)+(1+\eta)\left(x_{3}=2\right)\right.  \tag{6.322a}\\
&\left.-(1+\eta)\left(x_{4}=1\right)\right\}=\frac{1}{4}(3-\eta) \\
& \frac{\partial y}{\partial \xi}=\frac{1}{4}\left\{-(1-\eta)\left(y_{1}=0\right)+(1-\eta)\left(y_{2}=0\right)+(1+\eta)\left(y_{3}=3\right)\right.  \tag{6.322b}\\
&\left.-(1+\eta)\left(y_{4}=3\right)\right\}=0 \\
& \frac{\partial x}{\partial \eta}=\frac{1}{4}\left\{-(1-\xi)\left(x_{1}=0\right)-(1+\xi)\left(x_{2}=2\right)+(1+\xi)\left(x_{3}=2\right)\right.  \tag{6.322c}\\
&\left.+(1-\xi)\left(x_{4}=1\right)\right\}=\frac{1}{4}(1-\xi) \\
& \frac{\partial y}{\partial \eta}=\frac{1}{4}\left\{-(1-\xi)\left(y_{1}=0\right)-(1+\xi)\left(y_{2}=0\right)+(1+\xi)\left(y_{3}=3\right)\right.  \tag{6.322d}\\
&+(1-\xi)\left(y_{4}=3\right)=\frac{6}{4}
\end{align*}
$$

leading to the Jacobian matrix given by

$$
\mathbf{J}=\left[\begin{array}{ll}
\frac{1}{4}(3-\eta) & 0  \tag{6.323}\\
\frac{1}{4}(1-\xi) & \frac{6}{4}
\end{array}\right]
$$

with its determinant

$$
\begin{equation*}
|\mathbf{J}|=\frac{3}{8}(3-\eta) \tag{6.324}
\end{equation*}
$$

The inverse of the Jacobian matrix becomes

$$
\mathbf{J}^{-1}=\left[\begin{array}{cc}
\frac{4}{(3-\eta)} & 0  \tag{6.325}\\
-\frac{2(1-\xi)}{3(3-\eta)} & \frac{2}{3}
\end{array}\right]
$$

The determinant of the Jacobian matrix can be also determined from

$$
\begin{equation*}
|\mathbf{J}|=\frac{1}{8}\left[\left(x_{31} y_{42}-x_{42} y_{31}\right)+\xi\left(x_{12} y_{23}-x_{23} y_{12}\right)+\eta\left(x_{41} y_{32}-x_{32} y_{41}\right)\right] \tag{6.326}
\end{equation*}
$$

in which

$$
\begin{array}{lll}
x_{31}=x_{3}-x_{1}=2 & x_{43}=x_{4}-x_{3}=-1 & x_{32}=x_{3}-x_{2}=0 \\
y_{31}=y_{3}-y_{1}=3 & y_{43}=y_{4}-y_{3}=0 & y_{32}=y_{3}-y_{2}=3 \\
y_{42}=y_{4}-y_{2}=3 & x_{21}=x_{2}-x_{1}=2 & x_{41}=x_{4}-x_{1}=1 \\
x_{42}=x_{4}-x_{2}=-1 & y_{21}=y_{2}-y_{1}=0 & y_{41}=y_{4}-y_{1}=3 \tag{6.327}
\end{array}
$$

Substituting for the following derivatives

$$
\begin{align*}
& \sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \xi} x_{p}=\frac{\partial x}{\partial \xi}=\frac{1}{4}(3-\eta) \\
& \sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \xi} y_{p}=\frac{\partial y}{\partial \xi}=0  \tag{6.328}\\
& \sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \eta} x_{p}=\frac{\partial x}{\partial \eta}=\frac{1}{4}(1-\xi) \\
& \sum_{p=1}^{4} \frac{\partial N_{p}}{\partial \eta} y_{p}=\frac{\partial y}{\partial \eta}=\frac{3}{2}
\end{align*}
$$

permits the derivatives of the shape functions as

$$
\begin{gather*}
\frac{\partial N_{p}}{\partial x}=\frac{8}{3(3+\eta)}\left\{-\frac{6}{4} \frac{\partial N_{p}}{\partial \xi}\right\}=-\frac{4}{(3+\eta)} \frac{\partial N_{p}}{\partial \xi}  \tag{6.329}\\
\frac{\partial N_{p}}{\partial y}=\frac{2(1+\xi)}{3(3+\eta)} \frac{\partial N_{p}}{\partial \xi}-\frac{2}{3} \frac{\partial N_{p}}{\partial \eta}
\end{gather*}
$$

Thus, the components of the element shape matrix, $\mathbf{B}^{(1)}$ are computed as

$$
\begin{align*}
& \frac{\partial N_{1}}{\partial x}=-\frac{(1-\eta)}{(3-\eta)}, \frac{\partial N_{2}}{\partial x}=\frac{(1-\eta)}{(3-\eta)}, \\
& \frac{\partial N_{3}}{\partial x}=\frac{(1+\eta)}{(3-\eta)}, \quad \frac{\partial N_{4}}{\partial x}=-\frac{(1+\eta)}{(3-\eta)} \\
& \frac{\partial N_{1}}{\partial y}=-\frac{1}{3} \frac{(1-\xi)}{(3-\eta)}, \quad \frac{\partial N_{2}}{\partial y}=-\frac{(2+\xi-\eta)}{3(3-\eta)},  \tag{6.330}\\
& \frac{\partial N_{3}}{\partial y}=\frac{(1+2 \xi-\eta)}{3(3-\eta)}, \quad \frac{\partial N_{4}}{\partial y}=\frac{2(1-\xi)}{3(3-\eta)} \\
& \mathbf{B}^{(1)}=\left[\begin{array}{cccc}
-\frac{(1-\eta)}{(3-\eta)} & 0 & -\frac{(1-\eta)}{(3-\eta)} & 0 \\
0 & -\frac{1}{3} \frac{(1-\xi)}{(3-\eta)} & 0 & -\frac{(2+\xi-\eta)}{3(3-\eta)} \\
-\frac{1}{3} \frac{(1-\xi)}{(3-\eta)} & -\frac{(1-\eta)}{(3-\eta)} & -\frac{(2+\xi-\eta)}{3(3-\eta)} & -\frac{(1-\eta)}{(3-\eta)}
\end{array}\right.  \tag{6.331}\\
& \frac{(1+\eta)}{(3-\eta)} \quad 0 \quad-\frac{(1+\eta)}{(3-\eta)} \quad 0 \\
& 0 \quad \frac{(1+2 \xi-\eta)}{3(3-\eta)} \quad 0 \quad \frac{2(1-\xi)}{3(3-\eta)} \\
& \left.\frac{(1+2 \xi-\eta)}{3(3-\eta)} \quad \frac{(1+\eta)}{(3-\eta)} \quad \frac{2(1-\xi)}{3(3-\eta)} \quad-\frac{(1+\eta)}{(3-\eta)}\right]
\end{align*}
$$

Under plane stress assumptions, the material property matrix, D becomes

$$
\mathbf{D}=10^{6}\left[\begin{array}{ccc}
16 & 4 & 0  \tag{6.332}\\
4 & 16 & 0 \\
0 & 0 & 6
\end{array}\right] \mathrm{N} / \mathrm{cm}^{2}
$$

The element stiffness matrix, $\mathbf{k}^{(1)}$, is computed as

$$
\mathbf{k}^{(1)}=10^{6}\left[\begin{array}{cccc}
4.8666 & 0.76713 & -4.3666 & 0.23287  \tag{6.333}\\
0.76713 & 2.3545 & 0.73287 & -1.0211 \\
-4.3666 & 0.73287 & 5.3666 & -1.7329 \\
0.23287 & -1.0211 & -1.7329 & 3.6878 \\
-2.7668 & -0.96574 & 2.2668 & -0.034264 \\
-0.96574 & -1.1244 & -0.53426 & -0.20891 \\
2.2668 & -0.53426 & -3.2668 & 1.5343 \\
-0.034264 & -0.20891 & 1.5343 & -2.4578 \\
-2.7668 & -0.96574 & 2.2668 & -0.034264 \\
-0.96574 & -1.1244 & -0.53426 & -0.20891 \\
2.2668 & -0.53426 & -3.2668 & 1.5343 \\
-0.034264 & -0.20891 & 1.5343 & -2.4578 \\
6.9663 & 0.56853 & -6.4663 & 0.43147 \\
0.56853 & 3.5845 & 0.93147 & -2.2512 \\
-6.4663 & 0.93147 & 7.4663 & -1.9315 \\
0.43147 & -2.2512 & -1.9315 & 4.9178
\end{array}\right]
$$

The initial strains arising from the temperature change are included in the vector $\varepsilon^{*}$ as

$$
\varepsilon^{* T}=10^{-6}\left[\begin{array}{lll}
60 & 60 & 0 \tag{6.334}
\end{array}\right]
$$

The element load vectors, $\mathbf{p}_{\mathrm{T} 1-4}^{(1)}$ and $\mathbf{p}_{\mathrm{T} 3-4}^{(1)}$, arising from the applied tractions are

$$
\begin{align*}
& \mathbf{p}_{\mathrm{T} 1-4}^{(1) T}=t \frac{T_{x} L_{1-4}}{2}\left[\begin{array}{llllllll}
1 & 0 & 0 & 0 & 0 & 0 & 1 & 0
\end{array}\right]  \tag{6.335}\\
& \mathbf{p}_{\mathrm{T} 3-4}^{(1) T}=t \frac{T_{y} L_{3-4}}{2}\left[\begin{array}{llllllll}
0 & 0 & 0 & 0 & 0 & 1 & 0 & 1
\end{array}\right] \tag{6.336}
\end{align*}
$$

With the specified values of the thickness and the distributed loads, these element load vectors become

$$
\begin{align*}
\mathbf{p}_{\mathbf{T} 1-4}^{(1) T} & =300 \sqrt{10}\left[\begin{array}{llllllll}
1 & 0 & 0 & 0 & 0 & 0 & 1 & 0
\end{array}\right] \mathrm{N}  \tag{6.337}\\
\mathbf{p}_{\mathbf{T} 3-4}^{(1) T} & =-150\left[\begin{array}{llllllll}
0 & 0 & 0 & 0 & 0 & 1 & 0 & 1
\end{array}\right] \mathrm{N} \tag{6.338}
\end{align*}
$$

The element load vector from all the applied tractions is

$$
\mathbf{p}_{\mathrm{T} 1-4}^{(1)}+\mathbf{p}_{\mathrm{T} 3-4}^{(1)}=\left\{\begin{array}{c}
300 \sqrt{10}  \tag{6.339}\\
0 \\
0 \\
0 \\
0 \\
-150 \\
300 \sqrt{10} \\
-150
\end{array}\right\} \mathrm{N}
$$

The thermal load vector of the element, $\mathbf{p}_{\varepsilon^{*}}^{(1)}$, is obtained as

$$
\mathbf{p}_{\varepsilon^{*}}^{(1)}=\left\{\begin{array}{c}
-900  \tag{6.340}\\
-300 \\
900 \\
-600 \\
900 \\
300 \\
-900 \\
600
\end{array}\right\} \mathrm{N}
$$

Thus, the total element load vector, $\mathbf{P}$ is

$$
\mathbf{P}=\left\{\begin{array}{c}
(300 \sqrt{10}-900)  \tag{6.341}\\
-300 \\
900 \\
(-300-300) \\
900 \\
(-150+300) \\
(300 \sqrt{10}-900) \\
-150+600
\end{array}\right\} \mathbf{N}
$$

After applying the boundary conditions, the global stiffness matrix is reduced to

$$
\mathbf{K}=10^{6}\left[\begin{array}{cccc}
4.8666 & -0.96574 & 2.2668 & -0.034264  \tag{6.342}\\
-0.96574 & 3.5845 & 0.93147 & -2.2512 \\
2.2668 & 0.93147 & 7.4663 & -1.9315 \\
-0.034264 & -2.2512 & -1.9315 & 4.9178
\end{array}\right]
$$

and the reduced load vector is

$$
\mathbf{P}=\left\{\begin{array}{c}
(300 \sqrt{10}-900)  \tag{6.343}\\
150 \\
(300 \sqrt{10}-900) \\
450
\end{array}\right\} \mathrm{N}
$$

The solution is given by

$$
\left\{\begin{array}{l}
u_{x_{1}}  \tag{6.344}\\
u_{y_{3}} \\
u_{x_{4}} \\
u_{y_{4}}
\end{array}\right\}=\left\{\begin{array}{c}
0.0000307806 \\
0.000150801 \\
0.0000222016 \\
0.000169468
\end{array}\right\} \mathrm{cm}
$$

### 6.2.2.6.2 ANSYS Solution

The nodal displacements of the plate subjected to uniform temperature can also be obtained using ANSYS. The solution procedure is outlined as follows:

## MODEL GENERATION

- Specify the element type (ET command) using the following menu path:


## Main Menu $>$ Preprocessor $>$ Element Type $>$ Add/EdivDelete

- Click on Add.
- Select Structural Solid from the left list and Quad 4node 42 from the right list; click on $O K$.
- Click on Options.
- In order to specify the 2-D idealization as plane stress with thickness, in the newly appeared dialog box, pull down the menu for Element behavior K3 and select Plane strs w/thk; click on OK (Fig. 6.43).
- Click on Close .


Fig. 6.43 Specification of element options.

- Specify real constants ( $\mathbf{R}$ command) using the following menu path:

Main Menu > Preprocessor > Real Constants > Add/Edit/Delete

- Click on Add.
- Click on $\boldsymbol{O K}$.
- Enter 5e-3 for Thickness THK; click on OK.
- Click on Close.

■ Specify material properties (MP command) using the following menu path:

## Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring another dialog box.
- Enter $150 e 9$ for $\boldsymbol{E X}$, and $\mathbf{0 . 2 5}$ for PRXY; click on $\boldsymbol{O K}$.
- In the Define Material Model Behavior dialog box, in the right window, under Structural, find Thermal Expansion, Secant Coefficient, and Isotropic, which will bring another dialog box (Fig. 6.44).
- Enter $\boldsymbol{6} \boldsymbol{e}-\mathbf{6}$ for $\boldsymbol{A P L X}$; click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

```
Material > Exit
```



Fig. 6.44 Specification of material behavior.

- Create nodes ( $\mathbf{N}$ command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Nodes > In Active CS

- A total of 4 nodes will be created (Table 6.7).
- Referring to Table 6.7, enter $x$ - and $y$-coordinates of node 1 (be sure to convert the coordinates to meters), and Click on Apply. This action will keep the Create Nodes in Active Coordinate System dialog box open. If the Node number field is left blank, then ANSYS will assign the lowest available node number to the node that is being created.
- Repeat the same procedure for the nodes 2 through 4 .
- After entering the $x$ - and $y$-coordinates of node 4 , click on $\boldsymbol{O K}$ (instead of Apply).
- The nodes should appear in the Graphics Window, as shown in Fig. 6.45 .
- Create one element ( $\mathbf{E}$ command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Elements > Auto Numbered > Thru Nodes

- Pick Мenu appears; pick four nodes in a clockwise (or counterclockwise) order.
- Click on $\boldsymbol{O K}$.


## SOLUTION

- Apply displacement boundary conditions (D command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes


Fig. 6.45 Generation of nodes.

- Pick Menu appears; pick nodes 1 and 2 along the bottom horizontal boundary (Fig. 6.45) and click on $O K$ on Pick Menu.
- Highlight $\boldsymbol{U Y}$ and enter $\boldsymbol{0}$ for VALUE; click on Apply.
- Pick Menu reappears; pick nodes 2 and 3 along the right vertical boundary (Fig. 6.45); click on OK on Pick Menu.
- Highlight $\boldsymbol{U X}$ and enter $\boldsymbol{O}$ for $\boldsymbol{V A L U E}$; click on $\boldsymbol{O K}$.
- Apply force boundary conditions on nodes ( $\mathbf{F}$ command) using the following menu path:
Main Menu >Solution > Define Loads > Apply > Structural > Force/Moment > On Nodes
- Pick Menu appears; pick nodes 1 and 4 along the slanted boundary; click on $\mathbf{O K}$.
- Enter $3 e 3{ }^{*}$ sqrt(0.1) for VALUE (Fig. 6.46).
- Click on Apply.
- Pick Menu reappears; pick nodes 4 and 3 along the top horizontal boundary; click on $\boldsymbol{O K}$.
- Pull down the menu for Direction of force/mom and select $\boldsymbol{F Y}$; Enter -150 for VALUE; click on OK.
- Apply thermal load (TUNIF command) using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Structural $>$ Temperature $>$ Uniform Temp

- Uniform Temperature dialog box appears; Enter 10 for Uniform temperature.
- Click on $\boldsymbol{O K}$.


Fig. 6.46 Application of external loads.

- Obtain solution (SOLVE command) using the following menu path:

Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status. If OK, close the Status Report Window and click on $\boldsymbol{O K}$ in Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

■ Review deformed shape (PLDISP command) using the following menu path:

Main Menu > General Postproc $>$ Plot Results $>$ Deformed Shape

- In the Plot Deformed Shape dialog box, choose the radio-button for Def + undef edge; click on OK.
- The deformed shape will appear in the Graphics Window, as shown in Fig. 6.47.
- Review displacement values (PRNSOL command) using the following menu path:

Main Menu > General Postproc > List Results > Nodal Solution

- Highlight DOF solution on the left list and All DOFs DOF on the right list; click on $\boldsymbol{O K}$.
- The list will appear in a new window, as shown in Fig. 6.48.


Fig. 6.47 Deformed configuration.


Fig. 6.48 List of nodal displacements.

### 6.3 Problems

6.1. Construct the finite element equations for the solution of the linear second-order ordinary differential equation given in the form

$$
p(x) \frac{d^{2} u(x)}{d x^{2}}+\frac{d p(x)}{d x} \frac{d u(x)}{d x}+q(x) u(x)=f(x)
$$

subject to the conditions given as

$$
u\left(x_{0}\right)=A, \quad u\left(x_{n}\right)=B
$$

by using the Galerkin technique within the realm of finite element method with linear interpolation functions.
6.2. By using a one-dimensional (line) $C^{l}$ continuous cubic element, derive the element coefficient matrix for the solution of the differential equation given as

$$
\frac{d^{2} u(x)}{d x^{2}}=f(x)
$$

Assume equally spaced nodal points.
6.3. By using quadratic interpolation functions, derive the element coefficient matrix for the solution of the differential equation given as

$$
\frac{d^{2} u}{d x^{2}}=e^{x}
$$

subject to the conditions

$$
u(0)=1 \text { and } \frac{d u}{d x}(4)=0
$$

Also, explicitly assemble both the global coefficient matrix and the right-hand vector for equally spaced nodal points located at $x=0,1$, 2,3 , and 4 .
6.4. Without giving any consideration to the boundary conditions, write down the contribution from the four elements, shown in Fig. 6.49, in the finite element formulation for the Poisson equation $\nabla^{2} \phi=C$. Denote all entries in the element coefficient matrices symbolically and write your answer in the form $[\mathbf{K}]\{\varphi\}+\{\mathbf{F}\}=\{0\}$.


Fig. 6.49 Four linear triangular elements forming a quadrilateral element.
6.5. In Problem 6.4, note that the interaction of the internal node 5 with all the adjacent elements is included in forming the equation arising from the field variable $\phi_{5}$ associated with the $5^{\text {th }}$ node. In the absence of external loads, the last row of the vector-matrix expression in the previous problem may be set directly equal to zero. Using the resulting equation, eliminate $\phi_{5}$ from the remaining four rows of the vector-matrix expression to obtain the element coefficient matrix and the contribution to the right-hand-side vector of a quadrilateral element made up of four simpler triangular elements.
6.6. Suppose a collection of elements (part of some larger collection) has a total of $n$ interior nodes and $m$ exterior (or boundary) nodes. The contribution from this collection to the global finite element equations can be written as

$$
[\mathbf{K}]^{e}\{\boldsymbol{\varphi}\}^{e}+\{\mathbf{f}\}^{e}
$$

The contributions from the exterior nodes, $\phi_{i}^{E}(i=1,2, \ldots, m)$, and the interior nodes, $\phi_{i}^{l}(i=m+1, \ldots, n+m)$, may be partitioned as

$$
\left[\begin{array}{cc}
\mathbf{K}^{E} & \mathbf{K}^{*} \\
\mathbf{K}^{* T} & \mathbf{K}^{I}
\end{array}\right]\left\{\begin{array}{c}
\boldsymbol{\varphi}^{E} \\
\boldsymbol{\varphi}^{I}
\end{array}\right\}+\left\{\begin{array}{c}
\mathbf{f}^{E} \\
\mathbf{f}^{I}
\end{array}\right\}
$$

where $\left[\mathbf{K}^{E}\right.$ ] is an $m \times m$ submatrix, $\left[\mathbf{K}^{I}\right]$ is an $n \times n$ submatrix, etc. Consideration of all of the contributions to the interior nodes results in

$$
\left[\mathbf{K}^{*}\right]^{T}\left\{\boldsymbol{\varphi}^{E}\right\}+\left[\mathbf{K}^{I}\right]\left\{\boldsymbol{\varphi}^{I}\right\}+\left\{\mathbf{f}^{I}\right\}=\{\mathbf{0}\}
$$

Proceeding from this point, eliminate the quantities $\varphi_{i}^{l}$ from the remaining equations to express the contribution from this collection of elements in the form

$$
\left[\mathbf{K}^{R}\right]^{e}\left\{\boldsymbol{\varphi}^{E}\right\}+\left\{\mathbf{f}^{R}\right\}
$$

where $\left[\mathbf{K}^{R}\right]$ is an $m \times m$ matrix. This technique is called substructuring.
6.7. For two-dimensional heat transfer in an isotropic body, the governing equation is

$$
\frac{\partial}{\partial x}\left(K \frac{\partial T}{\partial x}\right)+\frac{\partial}{\partial y}\left(K \frac{\partial T}{\partial y}\right)+q(x, y)=0
$$

where $T$ is temperature, $K$ is thermal conductivity, and $q(x, y)$ is the heat generation rate over the domain. Suppose the heat flux out of some portion, $S_{f}$, of the boundary is specified to have a constant value, $Q$, as shown in Fig. 6.50. Then, the boundary condition over $S_{f}$ becomes

$$
K\left(\frac{\partial T}{\partial v}\right)+Q=K\left[\left(\frac{\partial T}{\partial x}\right) n_{x}+\left(\frac{\partial T}{\partial y}\right) n_{y}\right]+Q=0
$$

where $\mathbf{n}=<n_{x}, n_{y}>$ is the unit normal vector to the boundary. Using the Galerkin technique, show in a general way how this boundary condition enters the right-hand-side vector.
6.8. Suppose that the heat flux is specified to be $Q$ over the side $4-5$ of the domain as shown in Fig. 6.51. Find explicitly the contribution of the interpolating function associated with node 4 to the right-hand-side vector in the system of equations derived in Problem 6.7:
(a.) for the case where element 3 is a linear triangular element.
(b) for the case where element 3 is a quadratic triangular element with a mid-side node between nodes 4 and 5 .

Hint: Use a local coordinate, $s$, directed along the side of the triangle from node 4 to node 5 . Note that the interpolating function associated with node 4 is linear in $s$ for linear interpolation and quadratic for quadratic interpolation.


Fig. 6.50 Heat generation within the body and flux boundary condition along $S_{f}$.


Fig. 6.51 Domain discretized with three triangular elements.
6.9. Explicitly evaluate the element coefficient matrix for the problem

$$
\frac{\partial^{2} \psi}{\partial x^{2}}+\frac{\partial^{2} \psi}{\partial y^{2}}=G
$$

using $2 \times 2$ Gaussian integration for a 4 -noded quadrilateral element whose nodal point locations are given by

| Node No. | $x$ | $y$ |
| :---: | :---: | :---: |
| 1 | 6.0 | 3.0 |
| 2 | -4.0 | 3.0 |
| 3 | -5.0 | -3.0 |
| 4 | 4.0 | -3.0 |

6.10. Using quadratic interpolation over a 6 -noded triangle (shown in Fig. 6.52), derive explicit expressions for the entries $K_{11}, K_{44}$, and $K_{15}$ in the element coefficient matrix for the Poisson equation

$$
\frac{\partial^{2} \phi}{\partial x^{2}}+\frac{\partial^{2} \phi}{\partial y^{2}}=f(x, y)
$$

6.11. Consider the 3 -noded triangular element subjected to traction boundary conditions along the $2-3$ side as shown in Fig. 6.53. Assuming plane stress idealization with thickness $t=0.01 \mathrm{~m}, E=200 \mathrm{GPa}$, and $v=0.25$, construct:
(a) the stiffness matrix.
(b) the equivalent nodal force vector.


Fig. 6.52 Three-noded triangular element under uniform traction.


Fig. 6.53 A six-noded triangular element.
6.12. Assume that the nodal displacement components of the triangular element considered in Problem 6.11 are as follows:

$$
\begin{array}{cc}
u_{1}=0 & v_{1}=0 \\
u_{2}=3.30078 \times 10^{-4} \mathrm{~m} & v_{2}=0 \\
u_{3}=1.85937 \times 10^{-4} \mathrm{~m} & v_{3}=4.6875 \times 10^{-6} \mathrm{~m}
\end{array}
$$

Find the stress components ( $\sigma_{x x}, \sigma_{y y}$ and $\sigma_{x y}$ ).
6.13. Assuming that the triangular element considered in Problem 6.11 is subjected to gravitational acceleration in the negative $y$-direction with mass density $\rho=7850 \mathrm{~kg} / \mathrm{m}^{3}$, find the equivalent nodal force vector.
6.14. Derive the equivalent nodal force vector for a 3-noded triangular element when it is subjected to a uniform temperature change of $\Delta T$. The coefficient of thermal expansion of the material is $\alpha$.
6.15. The equations governing the time-dependent motion of an elastic body are

$$
\frac{\partial}{\partial x_{j}}\left[\sigma_{i j}\right]-\rho \frac{\partial^{2} u_{i}}{\partial t^{2}}=0
$$

where $\rho$ is the mass density of the body. The term $\rho \partial^{2} u_{i} / \partial t^{2}$ may be interpreted as an "inertia" force, which is a special type of body force.
(a) Identifying the inertia force as a body force with $F_{i}=-\rho \partial^{2} u_{i} / \partial t^{2}$, derive the contribution from a single element to the global finite element formulation for the case of plane strain.
(b) If no tractions are specified over the surface of the body, write down the general form of the global finite element equations. Assuming

$$
\{\mathbf{u}\}=\{\tilde{\mathbf{u}}\} e^{i \omega t}
$$

write down an equation for $\omega$, the natural frequencies of vibration.
6.16. A two-dimensional situation that is often of theoretical interest (although less seldom of practical interest) is that of antiplane strain, in which $u_{1}=u_{2}=0$ and $u_{3}=u_{3}\left(x_{1}, x_{2}\right)$. Hence, the only non-zero components of strain are $\varepsilon_{13}$ and $\varepsilon_{23}$ and those of stress are $\sigma_{13}$ and $\sigma_{23}$, which are related by Hooke's law:

$$
\sigma_{13}=\frac{E \varepsilon_{13}}{(1+v)}, \quad \sigma_{23}=\frac{E \varepsilon_{23}}{(1+v)}
$$

Find the element coefficient matrix for this problem for the linear triangle (3-noded) using the integration formulas for area coordinates given previously.
6.17. Newton's method is a familiar recursive technique for finding the roots of a transcendental equation. Suppose the roots of $n$ transcendental equations, $\left\{g_{i}\left(a_{j}\right)\right\}=0$, in $n$ unknowns are to be found. Then, Newton's method can be generalized to

$$
\left\{x_{i}\right\}^{(m+1)}=\left\{x_{i}\right\}^{(m)}-\left(\left[\frac{\partial g_{i}}{\partial x_{j}}\right]^{-1}\right)^{(m)}\left\{g_{i}\right\}^{(m)}
$$

where

$$
\left[\frac{\partial g_{i}}{\partial x_{j}}\right]^{(m)}=\left[\begin{array}{cccc}
\frac{\partial g_{1}}{\partial a_{1}} & \frac{\partial g_{1}}{\partial a_{2}} & \cdots & \frac{\partial g_{1}}{\partial a_{n}} \\
\frac{\partial g_{2}}{\partial a_{1}} & \frac{\partial g_{2}}{\partial a_{2}} & \cdots & \frac{\partial g_{2}}{\partial a_{n}} \\
\vdots & \vdots & \ddots & \vdots \\
\frac{\partial g_{n}}{\partial a_{1}} & \frac{\partial g_{n}}{\partial a_{2}} & \cdots & \frac{\partial g_{n}}{\partial a_{n}}
\end{array}\right]^{(m)}
$$

and $\left\{g_{i}\right\}^{(m)}$ and $\left[\partial g_{i} / \partial a_{j}\right]$ are evaluated at $\left\{a_{i}\right\}^{(m)}$.
The finite element equations resulting from the nonlinear two-point boundary value problem

$$
\frac{d^{2} u}{d x^{2}}+g(u, x)=0
$$

have the form

$$
\left[K_{i j}\right]\left\{a_{i}\right\}+\left\{f_{i}\left(a_{j}\right)\right\}=\{0\} \quad(i=1,2, \ldots, n)
$$

where $\left\{a_{i}\right\}$ are the nodal values and $\left\{f_{i}\left(a_{j}\right)\right\}$ is some nonlinear function of the nodal values. Apply Newton's method to this problem to obtain a recursive formula for the nodal values. What is the major drawback of this approach?

## Chapter 7

## USE OF COMMANDS IN ANSYS

The distinct differences between the two modes of ANSYS usage, i.e., the Graphical User Interface (GUI) and Batch Mode, are covered briefly in Chap. 2, and the most common operations within the Preprocessor, Solution, and Postprocessors, mainly using the GUI, are covered in Chap. 4 and 5. This chapter is devoted to using the Batch Mode of ANSYS, which is the method preferred by advanced ANSYS users.

As mentioned in Chap. 2, every action taken by the user within the ANSYS GUI platform has an equivalent ANSYS command. Using ANSYS through the Batch Mode involves text (ASCII) files with specific ANSYS commands. These commands, along with specific rules, form a special programming language, ANSYS Parametric Design Language, or APDL, which utilizes concepts and structures very similar to common scientific programming languages such as BASIC, FORTRAN, etc. Using the APDL, the user can create (a) an Input File to solve a specific problem and (b) Macro File(s) that act as special functions, accepting several arguments as input. In either case, each line consists of a single command, and the lines are executed sequentially.

The basic ANSYS commands, operators, and functions are discussed in the following sections. After solving a simple problem by using the Batch Mode, more advanced APDL features are covered. The Batch Mode command files for each example problem included in this book are given on the accompanying CD-ROM.

### 7.1 Basic ANSYS Commands

There are around 1,500 ANSYS commands, each with a specific syntax and function. It is impractical (and perhaps impossible) for the user to learn the use of all of the commands. This apparent obstacle is overcome by using the ANSYS Help System, accessible from within the program, which is covered in Sec. 2.7. However, the solution of a typical problem often involves a limited number of commonly used commands. A selection of these common
commands is presented in tabular form in this section. Within the context of this book, they are grouped into the following six categories:

- Session and Database Commands (Table 7.1).
- APDL Commands (Table 7.2).
- Preprocessor Solid Model Generation Commands (Table 7.3).
- Preprocessor Meshing Commands (Table 7.4).
- Solution Commands (Table 7.5).
- General Postprocessor Commands (Table 7.6).

In Tables 7.1-7.6, the first column gives the command and the corresponding description is given in the second column. With the exception of some APDL commands, the commands can also be issued as a command line input in the Input Field in the ANSYS GUI. It is worth noting that some ANSYS commands are valid only in a specific processor or BEGIN level while the remaining ones are valid at all times. Most of the ANSYS commands require arguments separated by commas. For example, the syntax for the K command (to create keypoints) given in Table 7.3 is

$$
\mathrm{K}, \mathrm{NPT}, \mathrm{X}, \mathrm{Y}, \mathrm{Z}
$$

where NPT is the keypoint number and $\mathbf{X}, \mathbf{Y}$, and $\mathbf{z}$ are the $x$-, $y$-, and $z$ coordinates of the node,

As explained in Sec. 2.7, the help page related to the use of this command can be retrieved by issuing the following command line input in the Input Field in ANSYS:

HELP, K
This command brings up detailed information about the arguments.
Tables 7.1-7.6 serve as an introduction to the ANSYS commands. However, it is highly recommended that the user read the help pages before usage.

### 7.1.1 Operators and Functions

In the ANSYS Parametric Design Language (APDL), several fundamental mathematical operations can be utilized through the use of common operators and functions. A complete list of operators is given in Table 7.7. Table 7.8 lists selected mathematical functions available within APDL. Section 7.1.2 provides several examples demonstrating the definition and use of parameters in APDL. These examples are also useful in understanding the way mathematical operators and functions are used in ANSYS.

Table 7.1 Session and database commands

| Command | Description |
| :--- | :--- |
| /CLEAR | Clear the database (and memory) |
| /PREP7 | Enter the Preprocessor |
| /SOLU | Enter the Solution |
| /POST1 | Enter the General Postprocessor |
| /POST26 | Enter the Time History Postprocessor |
| FINISH | Exit the current processor; go to Begin |
| /EOF | level |
| /FILNAME | Marks the end of file (stop reading) |
| HELP | Specify jobname |
| SAVE | Display help pages related to the command |
| RESUME | Save the database |
| KSEL, LSEL, ASEL, VSEL, | Resume from an existing database |
| NSEL, ESEL, CMSEL | Select keypoints, lines, areas, volumes, |
| ALLSEL | nodes, elements, and components |
| CLOCAL, LOCAL | Select all entities |
| CSYS | Define local coordinate systems |
| ! | Switch between coordinate systems |
|  | Start comment-ANSYS ignores the |
|  | characters to the right of the exclama- |

Table 7.2 APDL commands.

| Command | Description |
| :--- | :--- |
| *AFUN | Switch between degrees and radians to be <br> used for angles |
| *GET | Store model or result information into <br> parameters |
| *VWRITE | Write formatted output to external files |
| *DO, *ENDDO | Beginning and ending of do loops |
| *IF, *ELSE, *ELSEIF, | Commands related to IF-THEN-ELSE |
| *ENDIF | blocks |
| *SET | Define parameters |

Table 7.3 Preprocessor solid model generation commands.

| Command | Description |
| :---: | :---: |
| BLC4 | Create rectangular area or prism volume |
| CYL 4 | Create circular area or cylindrical volume |
| $K, L, A, A L, V, V A$ | Create keypoints, lines, areas, and volumes |
| LARC | Create circular arc |
| SPLINE, BSPLIN | Create line through spline fit to keypoints |
| ADRAG | Create an area by dragging a line along a path |
| VRAG | Create a volume by dragging an area along a path |
| VEXT | Create a volume by extruding an area |
| AADD, VADD | Add areas and volumes |
| LGLUE, AGLUE, VGLUE | Glue lines, areas, and volumes |
| LOVLAP, AOVLAP, VOVLAP | Overlap lines, areas, and volumes |
| CM | Create components |
| KDELE, LDELE, ADELE, VDELE, CMDELE | Delete keypoints, lines, areas, volumes, and components |
| KPLOT, LPLOT, APLOT, VPLOT | Plot keypoints, lines, areas, and volumes in the Graphics Window |
| KLIST, LLIST, ALIST, VLIST, CMLIST | List keypoints, lines, areas, volumes, and components |

Table 7.4 Preprocessor meshing commands.

| Command | Description |
| :--- | :--- |
| ET | Specify element type |
| MP | Specify real constants |
| N | Specify material properties |
| E | Create nodes |
| TYPE | Create elements |
| Specify default element type attribute |  |
| REAL | number <br> Specify default real constant set attribute <br> number <br> MAT |
| LMESH, AMESH, VMESH | Specify default material property set <br> attribute number <br> Mesh the lines, areas, and volumes <br> Llear the mesh from lines, areas, and |
| LCLEAR, ACLEAR, | Cles <br> volumes (deletes the nodes and elements <br> attached to those entities) |
| LESIZE | Specify number of elements or element <br> sizes along selected lines |
| MSHKEY | Specify whether to use mapped or free <br> meshing |
| NDELE, EDELE | Delete nodes and elements |
| NPLOT, EPLOT | Plot nodes and elements in the Graphics <br> Window |

Table 7.5 Solution commands.

| Command | Description |
| :--- | :--- |
| SOLVE | Start solution for the current load step |
| LSSOLVE | Start solution from multiple load step files |
| D | Specify DOF constraints on nodes |
| F | Specify concentrated load boundary conditions on <br> nodes |
| SF, SFE, SFL, SFA | Specify surface (distributed) loads on nodes, <br> elements, lines, and areas |
| BF, BFE | Specify body loads on nodes and elements <br> TUNIF |
| Specify uniform thermal load on all nodes |  |
| IC | Specify initial conditions |
| LSREAD, LSWRITE | Read from and write to load step files |

Table 7.6 General postprocessor commands.

| Command | Description |
| :--- | :--- |
| FILE | Specify the results file for the results to be read from |
| SET | Specify the load step and substep numbers to be loaded |
| PLDISP | Plot deformed shape |
| PLNSOL | Plot contours of nodal solution |
| PLESOL | Plot contours of element solution |
| PRNSOL | List nodal solution items |
| PRESOL | List element solution items |

Table 7.7 List of operators within ANSYS.

| Operator | Description |
| :---: | :---: |
| + | Addition |
| - | Subtraction |
| $*$ | Multiplication |
| $/$ | Division |
| $* *$ | Exponentiation |
| $<$ | Less-than comparison |
| $>$ | Greater-than comparison |
| $=$ | Equal to (used in defining parameters) |

Table 7.8 Selected ANSYS functions.

| Function | Description |
| :---: | :---: |
| ABS (X) | Absolute value of $X$ |
| EXP (X) | Exponential of $X$ |
| LOG (X) | Natural logarithm of $X$ |
| LOG10 (X) | Base 10 logarithm of $X$ |
| SQRT (X) | Square root of $X$ |
| NINT (X) | Nearest integer to $X$ |
| RAND ( $\mathrm{X}, \mathrm{Y}$ ) | Random number within the range $X$ to $Y$ |
| $\operatorname{SIN}(\mathrm{X}), \operatorname{COS}(\mathrm{X}), \operatorname{TAN}(\mathrm{X})$ | Sine, cosine, and tangent of $X$ |
| $\begin{aligned} & \operatorname{SINH}(X), \operatorname{COSH}(X), \\ & \text { TANH (X) } \end{aligned}$ | Hyperbolic sine, hyperbolic cosine, and hyperbolic tangent of $X$ |
| $\begin{aligned} & \operatorname{ASIN}(X), \operatorname{ACOS}(X), \\ & \operatorname{ATAN}(X) \end{aligned}$ | Inverse sine, inverse cosine, and inverse tangent of $X$ |

### 7.1.2 Defining Parameters

Parameters in APDL can be defined by using either the *SET command or the "equal to" sign (=). For example the parameter "USRPRM" can be defined to have the value 22 by either

```
*SET,USRPRM,22
```

or
USRPRM $=22$
The rules for naming of parameters are:

- The first character of a parameter name must be a letter.
- Within the parameter name, only letters, numbers, and the underscore character ( _ ) are allowed.
- The maximum number of characters within a parameter name is 32 .

The use of common mathematical operations and functions (Tables 7.7-7.8) in parameter definitions is illustrated in the example below. Similar input files for various examples considered in this book are also provided on the CD-ROM.

A rectangular area consisting of two dissimilar materials, shown in Fig. 7.1, has a width and height of $w$ and $h$, respectively. The material interface starts on the left edge at point $(0, a)$, with an inclination angle $\theta$. Assuming the numerical values of $w=2, h=4, a=1$, and $\theta=30^{\circ}$, the following APDL block creates the solid model shown in Fig. 7.2:

```
/PREP7 ! ENTER PREPROCESSOR
*AFUN,DEG !SWITCH TO DEGREES
W=2 ! WIDTH
H=4 !HEIGHT
A=1 !Y-COORDINATE OF MATERIAL
                                    !INTERFACE AT LEFT EDGE
                            ! INCLINATION ANGLE
THETA=30
B= W*TAN(THETA)
! CREATE KEYPOINTS
K,1,0,0
K,2,W,0
K,3,0,A
K,4,W,A+B
K,5,0,H
K,6,W,H
```



Fig. 7.1 A rectangular area consisting of two dissimilar materials.


Fig. 7.2 ANSYS model created using numerical values of $w=2$, $h=4, a=1$, and $\theta=30^{\circ}$.
! CREATE LINES
L, 1, 2
L, 2, 4
L, 4, 3
L, 1, 3
L, 4, 6
L, 6, 5
L, 3, 5
! CREATE AREAS
AL, 1, 2, 3, 4
AL, 3, 5, 6, 7
Note that the distance $b$ is calculated using a mathematical operator (*) and the function taN to create keypoint 4 . The input block can be saved as a text file, "example.txt," in the Working Directory and read from within ANSYS using the following menu path:

## Utility Menu > File > Read Input from

It is also possible to read input files by issuing the /INPUT command in the Input Field in ANSYS GUI as follows:

```
/INPUT, EXAMPLE, TXT
```

Convenience in using the Batch Mode is demonstrated by modifying the length and angle parameters defined in the previous example. Figure 7.3 shows the solid model generated using $w=5, h=5, a=2$, and $\theta=15^{\circ}$ and Fig. 7.4 shows the one using $w=1, h=5, a=2$, and $\theta=60^{\circ}$.


Fig. 7.3 ANSYS model created using numerical values of $w=5, h=5, a=2$, and $\theta=15^{\circ}$.


Fig. 7.4 ANSYS model created using numerical values of $w=1, h=5, a=2$, and $\theta=60^{\circ}$.

It is worth noting that if a parameter is redefined in the input file, the new value is not reflected in the entities or parameters defined previously. For example, keypoint 4 is created using parameters $w, a$, and $b$. If the parameter $w$ is redefined (from 2 to 5 ) after the creation of keypoint 4 as shown below, the new value of $w$ is not reflected in the definition of keypoint 4 and the $x$ coordinate of keypoint 4 remains as 2.

```
W=2
A=1
THETA=30
B= W*TAN (THETA)
K,4,W,A+B ! CREATE KEYPOINT 4
W=5
```


### 7.2 A Typical Input File

Typical steps involved in solving an engineering problem are listed in Sec. 5.1. A similar data structure is observed in the Input Files used in the Batch Mode. In order to demonstrate the use of the Batch Mode for a complete analysis, a thin, square structure with a centric circular hole subjected to tensile loading in the $y$-direction, as shown in Fig. 7.5, is considered.

The length of the square and the radius of the circular hole are $w=4 \mathrm{in}$. and $r=1 \mathrm{in}$., respectively, and the thickness of the structure is $t=0.1 \mathrm{in}$. The geometry and material possess quarter-symmetry, therefore only one-fourth of the domain is modeled. Because the thickness is significantly smaller than the in-plane dimensions, a plane stress assumption is used. The elastic modulus and Poisson's ratio are $E=30 \times 10^{6}$ psi and $v=0.30$, respectively. The distributed tensile load is specified as $q=10 \mathrm{lbs} / \mathrm{in}$, and it is applied in the form of pressure loading with $q=-10$. The corresponding pressure is input as $q=-10$. The analysis is demonstrated by utilizing two separate solid modeling approaches: Bottom-up and Top-down.

The Input File below uses the Bottom-up approach, which starts building the model with keypoints, then line from keypoints, and, finally, areas using lines (explanations are given along with the commands; the commands between the dashed lines correspond to the Bottom-up approach in solid modeling).

```
/FILNAM, BOT-UP !SPECIFY JOBNAME
/PREP7 ! ENTER PREPROCESSOR
ET,1,42 !SELECT ELEMENT TYPE AS
! PLANE42
!SPECIFY PLANE STRESS WITH
! THICKNESS
! SPECIFY REAL CONSTANT
! THICKNESS)
SPECIFY ELASTIC MODULUS
SPECIFY POISSON'S RATIO
SIDE LENGTH OF SQUARE
HOLE RADIUS
!APPLIED SURFACE LOAD
```



Fig. 7.5 A thin, square structure with a centric circular hole subjected to tensile loading in the $y$-direction (left); due to symmetry, only one-fourth of the structure is modeled (right).

| K, 1, 0, 0 | ! CREATE KEYPOINTS |
| :---: | :---: |
| K, 2, R, 0 |  |
| K, 3, W/2, 0 |  |
| $\mathrm{K}, 4,0, \mathrm{R}$ |  |
| K, 5, 0, W/ 2 |  |
| K, 6,W/2,W/2 |  |
| L, 2, 3 | ! CREATE LINES |
| L, 3, 6 |  |
| L, 6, 5 |  |
| L, 5, 4 |  |
| LARC, 4, 2, 1, R | ! CREATE ARC |
| LESIZE, 1, , 10 | ! SPECIFY NUMBER OF |
| LESIZE, 4, , 10 | ! ELEMENTS ALONG LINES |
| LESIZE, 2, , 15 |  |
| LESIZE, 3, , 15 |  |
| LESIZE, 5, , 30 |  |
| AL, 1, 2, 3, 4, 5 | ! CREATE AREA |
| LCCAT, 2, 3 | ! CONCATENATE LINES FOR MAPPED <br> ! MESHING |
| MSHKEY, 1 | ! USE MAPPED MESHING |
| AMESH, ALL | ! MESH AREA |
| / SOLU | ! ENTER SOLUTION |
| NSEL, S, LOC , X, 0 | ! SELECT NODES AT X $=0$ |
| D, ALL, UX | ! SUPPRESS X-DISPLACEMENTS |
|  | ! AT SELECTED NODES |
| NSEL, S, LOC , Y, 0 | ! SELECT NODES AT Y $=0$ |
| D, ALL, UY | ! SUPPRESS Y-DISPLACEMENTS |
|  | $!$ AT SELECTED NODES |
| NSEL, S, LOC, Y, W/2 | ! SELECT NODES AT Y $=\mathrm{W} / 2$ |
| SF, ALL, PRES, -P | ! APPLY SURFACE LOADS |
| ALLSEL | ! SELECT EVERYTHING |
| SOLVE | ! SOLVE |
| / POST1 | ! ENTER POSTPROCESSOR |
| PLDISP, 2 | ! PLOT DEFORMED SHAPE |
| PLNSOL, S, Y | ! PLOT STRESS IN Y-DIR |
| /EOF | ! MARK THE END OF FILE |

Solid modeling using the Top-down approach to accomplish the same task is given below; the methods are interchangeable and the results are the same.

```
RECTNG,0,W/2,0,W/2 ! CREATE RECTANGLE
PCIRC,1 ! CREATE CIRCLE
ASBA,1,2 ! SUBTRACT CIRCLE FROM
! RECTANGLE
! SELECT LINES AT X = 0
! ADD LINES AT Y = 0 TO THE
! SELECTED SET
! SPECIFY NUMBER OF ELEMENTS
! ALONG LINES
! SELECT LINES AT X = W/2
! ADD LINES AT Y = W/2 TO
! THE SELECTED SET
! SPECIFY NUMBER OF ELEMENTS
! ALONG LINES
! CONCATENATE SELECTED LINES
! SWITCH TO GLOBAL CYLINDRICAL
! COORDINATE SYSTEM
! SELECT LINES AT r = R
! SPECIFY NUMBER OF ELEMENTS
! ALONG LINES
! SWITCH TO GLOBAL CARTESIAN
! COORDINATE SYSTEM
! SELECT EVERYTHING
```

The deformed shape of the structure and the contour variation of stresses in the $y$-direction after the solution are shown in Fig. 7.6 and 7.7, respectively.

### 7.3 Selecting Operations

Selecting operations play a key role when programming with APDL. The most commonly used ANSYS commands for selecting operations are given in Table 7.9.


Fig. 7.6 Deformed shape of the structure.


Fig. 7.7 Contour plot of normal stress in the $y$-direction.

Table 7.9 Commonly used ANSYS commands for selecting operations.

| Command | Description |
| :--- | :--- |
| ALLSEL | Select all the entities |
| KSEL, LSEL, ASEL, VSEL, | Select subsets of keypoints, lines, areas, <br> volumes, nodes, and elements <br> NSEL, ESEL |
| NSLE | Select nodes attached to the selected <br> elements <br> Select elements containing the selected <br> nodes |
| ESLN | Select nodes associated with the selected <br> lines, areas, and volumes <br> Select elements associated with the selected <br> lines, areas, and volumes |
| ESLL, ESLA, ESLV | Select keypoints contained in the selected <br> lines |
| KSLL | Select lines containing the selected key- <br> points |
| LSLK | Select lines contained in the selected areas <br> Select areas containing the selected lines |
| LSLA | Select areas contained in the selected <br> volumes |
| ASLL | Select volumes containing the selected <br> areas |
| ASLV |  |
| VSLA |  |

The basic group of selection commands involves the ones that allow the user to select a subset of entities, i.e., KSEL, LSEL, ASEL, VSEL, NSEL, and ESEL. The syntax for these commands is as follows:

```
KSEL, Type, Item, Comp, VMIN, VMAX, VINC, KABS
LSEL, Type, Item, Comp, VMIN, VMAX, VINC, KSWP
ASEL, Type, Item, Comp, VMIN, VMAX, VINC, KSWP
VSEL, Type, Item, Comp, VMIN, VMAX, VINC, KSWP
NSEL, Type, Item, Comp, VMIN, VMAX, VINC, KABS
ESEL, TYpe, Item, Comp, VMIN, VMAX, VINC, KABS
```

The first argument, "Type," determines the specific type of selection with the following possible values:

| S | Select a subset from the full set. |
| :--- | :--- |
| R | Select a subset from the current selected set. |
| A | Select a subset from the full set and add it to the current |
| selected set. |  |

Figure 7.8 graphically illustrates the concepts behind the argument Type. The following examples demonstrate the use of Type, along with the remaining arguments.

The argument Item, depending on the entity, may have several different meanings. The third through sixth arguments (Comp, VMIN, VMAX, and VINC) refer to the argument Item. The most commonly used Item arguments are:

- Entity name: KP for keypoints, LINE for lines, AREA for areas, VOLU for volumes, NODE for nodes, and ELEM for elements. In this case, the Comp field (stands for component) is left blank, and VMIN, VMAX, and VINC refer to the minimum and maximum values of the item range and value increment in range (if VINC is not specified, its default value is 1 ), respectively. For example, in order to select keypoints 21 through 30, the following statement is used:


Fig. 7.8 Graphical representation of the argument Type in selection logic.

- mat, read, type: Selects the entities based on their association with material, real constant, and element type attributes, with the exception of nodes. Similar to the entity name, the Comp field is left blank. The use of this item is demonstrated in the following example in which the elements with material property attribute number 2 are unselected:

```
ESEL,U,MAT, , 2
```

- LOC: This item allows the user to perform the selection operations based on the location of the entities, with the exception of elements. The Comp field in this case corresponds to the direction ( $x, y, z$ for a Cartesian coordinate system; $r, \theta, z$ for a cylindrical coordinate system; etc.). For example, the nodes located within the range $2.5 \leq z \leq 4$ can be added to the currently selected set of nodes using the following statement:

```
NSEL, A, LOC, Z, 2. 5,4
```

Several examples demonstrating the concepts used in selection operations are given below.

- Select nodes along planes $x=1$ and $x=1.5$ (but not the ones in between):

```
NSEL, S,LOC, X, 1
NSEL, A, LOC, X, 1.5
```

or
NSEL, S, LOC , X, 1, 1. 5, 0. 5

- Select nodes along planes $x=1$ and $x=1.5$ (but not the ones in between) and within the range $0 \leq y \leq 4$ :

```
NSEL,S,LOC,X,1
NSEL, A, LOC,X, 1.5
NSEL, R, LOC,Y, 0,4
```

- Select keypoints within the range $10<x<15$ (note that $x=10$ and $x=15$ are excluded):

```
TINY=1E-6 ! DEFINE SMALL NUMBER
KSEL, S,LOC,X,10+TINY,15-TINY
```

- Select keypoints with $x \leq 10$ and $x>15$ :

```
TINY=1E-6
KSEL,S,LOC,X,10+TINY,15
KSEL,INVE ! INVERT SELECTION
```

- Select elements 1 through 101 with the increment 2 :

ESEL, S, ELEM, 1, 101, 2

- Select elements with material attribute number 5 but without those whose real constant attribute number is 3 :

```
ESEL,S,MAT, , 5
ESEL,U,REAL, , 3
```

The remaining commands included in Table 7.9 perform more specific tasks, mostly utilizing the association between the entities. For example, the NSLE command is used for performing selection operations on nodes associated with the currently selected set of elements. The command line input given in the following example unselects the nodes that are attached to the selected set of elements:

```
NSLE,U
```

Command ESLN selects elements attached to the currently selected set of nodes. Analogous to the other selection commands, the first argument is Type, which determines the type of selection. The value of the second argument, EKEY, determines which elements are to be selected:

If EKEY $=0$, elements are selected if any of their nodes are in the selected node set.

If EKEY = 1, elements are selected only if all of their nodes are in the selected node set.

The following lines demonstrate the use of this command:

```
ESLN,S,0 ! CASE 1
ESLN,S,1 ! CASE 2
```


### 7.4 Extracting Information from ANSYS

Programming with the ANSYS Parametric Design Language often requires the extraction of data such as entity numbers and locations, geometric information, results, etc. Considering the fact that a typical FEA mesh consists of thousands of nodes and elements, the user does not usually have control over the entity numbering. Thus, the information that nodes exist at a specific location may be known without knowledge of their numbers. So, if the user is interested in extracting the variation of a certain solution item along a specific path, the aforementioned data extraction tasks must be performed. These tasks are achieved using the *GET command. The syntax of the *GET command is as follows:

[^8]The *GET command retrieves and subsequently stores data into parameters. The first argument, Par, is the parameter name given by the user. The help page for the *GET command provides a complete list of possible argument combinations, and it is highly recommended that the user refer to it. In order to explain the use of the *GET command, we consider the examples given below.

- Store the maximum and minimum node numbers in the currently selected node set in parameters maxnod and minnod:
*GET, maxnod, NODE, 0, NUM, MAX
*GET, minnod, NODE, 0, NUM, MIN
- Store the maximum and minimum element numbers in the currently selected element set in parameters maxel and minel:

```
*GET, maxel, ELEM, 0, NUM, MAX
*GET,minel, ELEM, O,NUM,MIN
```

- Store the number of nodes and elements in the currently selected node and element sets in parameters numnod and numel:
*GET, numnod, NODE, 0, COUNT
*GET, numel, ELEM, 0, COUNT
- Store the $x$ -,$y$, and $z$-coordinates of the node numbered maxnod in parameters $x l, y l$, and $z l$ :

```
*GET, x1,NODE,maxnod, LOC,X
*GET, y1, NODE, maxnod, LOC, Y
*GET, z1,NODE, maxnod, LOC, z
```

- Store the $x$-, $y$-, and $z$-displacements of the node numbered minnod in parameters $u 2, v 2$, and $w 2$ :

```
*GET,u2,NODE,minnod,U,X
*GET,v2,NODE,minnod,U,Y
*GET,w2,NODE,minnod,U,Z
```

- Store the rotations of the node numbered minnod about the $x$-, $y$-, and $z$ axes in parameters $r_{-} x, r_{-} y$, and $r_{-} z$ :

```
*GET,r_x,NODE,minnod,ROT,X
*GET, r_y,NODE, minnod,ROT, Y
*GET,r_z,NODE,minnod, ROT, z
```

- Store the shear stresses $\sigma_{x y}, \sigma_{y z}$, and $\sigma_{x z}$ and von Mises stress $\sigma_{e q v}$ at the node numbered maxnod in parameters $s_{-} x y, s_{-} y z, s_{-} x z$, and $s_{-} e q v$ :

```
*GET, s_xy,NODE,maxnod, S, XY
*GET, s_Yz,NODE, maxnod, S,YZ
```

*GET, s_xz, NODE, maxnod, $\mathrm{S}, \mathrm{Xz}$
*GET, s_eqv, NODE, maxnod, S, EQV

- Store the normal strains $\varepsilon_{x x}, \varepsilon_{y y}$, and $\varepsilon_{z z}$ at the node numbered minnod in parameters $e p s_{-} x x, e p s_{-} y y$, and $e p s_{-} z z:$

```
*GET, eps_xx, NODE,minnod, EPEL, X
*GET, eps_YY, NODE, minnod, EPEL, Y
*GET, eps_zz,NODE,minnod, EPEL, Z
```

- Store the $x$-, $y$-, and $z$-coordinates of the centroid of the element numbered maxel in parameters $c e_{-} x, c e_{-} y$, and $c e_{-} z$ :

```
*GET, ce__x, ELEM, maxel, CENT,X
*GET, ce_y, ELEM, maxel, CENT, Y
*GET, ce_z, ELEM, maxel, CENT, Z
```

- Store the area of the element numbered minel in parameters e_area:

```
*GET, e_area, ELEM,minel,AREA
```

As an alternative to the syntax given above, one can use readily available *GET functions that are predefined in compact form. A few of these functions are listed in Table 7.10. For example, the $x$-, $y$-, and $z$ displacements of the node numbered minnod can be stored in parameters $u 2$, $v 2$, and $w 2$ by using the following:

```
u2=UX(minnod)
v2=UY (minnod)
w2=UZ (minnod)
```

Table 7.10 Selected compact *GET functions.

| Command | Description |
| :---: | :---: |
| $\begin{aligned} & \text { NX(n), } N Y(n), \\ & N Z(n) \end{aligned}$ | Retrieve $x$-, $y$-, and $z$-coordinates of node numbered $\mathbf{n}$ |
| NDNEXT ( n ) | Retrieve node number of the next selected node having a node number greater than node $\boldsymbol{n}$ |
| ELNEXT (e) | Retrieve element number of the next selected element having an element number greater than element $e$ |
| $\begin{aligned} & \operatorname{UX}(\mathbf{n}), \quad \operatorname{UY}(\mathbf{n}), \\ & \operatorname{UZ}(\mathbf{n}) \end{aligned}$ | Retrieve $x$-, $y$-, and $z$-displacements of node numbered n |
| $\begin{aligned} & \operatorname{ROTX}(n), \quad \operatorname{ROTY}(n), \\ & \operatorname{ROTZ}(n) \end{aligned}$ | Retrieve rotations about $x-, y$, and $z$-axes of node numbered $n$ |
| TEMP ( $\mathbf{n}$ ) | Retrieve temperature at node numbered $n$ |
| PRES ( n ) | Retrieve pressure at node numbered n |

### 7.5 Programming with ANSYS

The ANSYS Parametric Design Language contains features that are common to other scientific programming languages. These include looping (DO loops) and conditional branching (IF statements), as well as writing formatted output to text files (/OUTPUT and *VWRITE commands). These concepts are discussed in the following subsections.

### 7.5.1 DO Loops

Do loops are program blocks containing a series of commands executed repeatedly, once for each value of the loop index. The APDL commands *DO and *ENDDO define the beginning and ending of a do loop, respectively. The syntax for the *DO command is

```
*DO,Par,IVAL,FVAL,INC
```

in which Par is the loop index and IVAL and FVAL designate the initial and final values of Par to be incremented by INC. For example, the following input block is used to find the arithmetic average of $x$-displacements along the boundary defined by $x=-2.5$ :

| /POST1 ! | ! ENTER POSTPROCESSOR |
| :---: | :---: |
| NSEL, S,LOC, X, -2.5 ! | ! SELECT NODES ALONG X $=-2.5$ |
| *GET, numnod, NODE, 0, COUNT ! | ! RETRIEVE NUMBER OF NODES |
| *GET, minnod, NODE, 0,NUM, MIN ! | ! RETRIEVE MINIMUM NODE NUMBER |
| sum=0 ! | ! INITIALIZE SUM OF |
|  | : DISPLACEMENTS |
| curnod=minnod ! | ! INITIALIZE CURRENT NODE |
|  | ! NUMBER |
| *DO, ii, 1, numnod ! | ! BEGIN DO LOOP |
| *GET, cux, NODE, curnod, U, X ! | ! RETRIEVE X- DISPLACEMENT OF |
|  | ! THE CURRENT NODE |
| sum=sum+cux ! | ! UPDATE SUMMATION |
| *GET, nextnod, NODE, curnod, NXT | NXTH ! STORE NEXT HIGHER NODE |
|  | ! NUMBER IN nextnod |
| curnod=nextnod ! | ! UPDATE CURRENT NODE |
| *ENDDO | ! END DO LOOP |
| avgdisp=sum/numnod ! | ! CALCULATE ARITHMETIC AVERAGE |

In the example above, ii is the loop index with the initial and final values of 1 and numnod, respectively. Before the do loop begins, the necessary information is obtained by using the *GET command (numnod for number of nodes and minnod for the minimum node number in the selected set of nodes). Also, two new parameters are defined:

- sum for the summation of displacements, which is updated within the do loop and finally divided by the number of nodes (numnod) to find the arithmetic average.
- curnod designating the node number of the "current node" within the do loop. Its initial value is set as the minimum node number in the selected set of nodes (minnod), and it is updated within the loop.

Additional do loops may be used within do loops. For example, the following input block creates 216 nodes, starting from the origin, with increments of $0.25,0.1$, and 0.5 in the $x$-, $y$-, and $z$-directions, respectively.

```
/PREP7 ! ENTER PREPROCESSOR
dx=0.25 ! DEFINE PARAMETER FOR INCREMENT IN X
dy=0.1 ! DEFINE PARAMETER FOR INCREMENT IN Y
dz=0.5 ! DEFINE PARAMETER FOR INCREMENT IN Z
    *DO,i,1,6 ! BEGIN DO LOOP IN i
    *DO,j,1,6 ! BEGIN DO LOOP IN j
*DO,k,1,6 ! BEGIN DO LOOP IN k
N,,(i-1)*dx,(j-1)*dy,(k-1)*dz ! CREATE NODE
*ENDDO ! END DO LOOP IN k
*ENDDO ! END DO LOOP IN j
*ENDDO ! END DO LOOP IN i
```


### 7.5.2 IF Statements

Conditional branching, which is the execution of an input block based on a condition, is accomplished by using the *IF command. The syntax for the *IF command is
*IF, VAL1, Oper1, VAL2, Base1, VAL3, Oper2, VAL4, Base2
in which VAL1, VAL2, VAL3, and VAL4 are numerical values or parameters that are compared (VAL1 is compared to VAL2, and VAL3 is compared to VAL4). The types of these comparisons are dictated by operator arguments Oper1 and Oper2. Finally, the arguments Base1 and Base2 specify the action to be taken based on the comparisons. Operator arguments Oper1 and Oper 2 may take the following selected logical values:

```
EQ Equal to (VAL1 = VAL2).
NE Not equal (VAL1 = VAL2).
LT Less than (VAL1<VAL2).
GT Greater than (VAL1 > VAL2).
LE Less than or equal to (VAL1\leqVAL2).
GE Greater than or equal to (VAL1\geqVAL2).
```

Common logical values for action arguments Base1 and Base 2 are:

| AND | True if both comparisons dictated by Oper1 and Oper2 are <br> true. |
| :--- | :--- |
| OR | True if either one of the comparisons dictated by Oper1 and <br> Oper2 is true. |
| XOR | True if either one but not both of the comparisons dictated by |
| OHENOper1 and Oper2 is true. <br> If the preceding logical comparison is true, continue to the <br> next line, otherwise skip to one of the following commands <br> (whichever appears first): *ELSE, *ELSEIF, or *ENDIF. |  |
| This point is explained in further detail below. |  |

In the event that the first action argument Base1 has the logical value THEN, which is often the case, then the conditional branching has the form

```
*IF,VAL1,Oper1,VAL2,THEN
```

and implies that this is an IF-THEN-ELSE block and that it must be ended by a *ENDIF command. Between the *IF (marking the beginning of the block) and *ENDIF (marking the end of the block) commands, the user may use *elseif and *else commands. A typical tf-Then-else block has the following form:

```
*IF,VAL1,Oper1,VAL2,THEN ! COMPARISON-1
... ! APDL-1 (ANY NUMBER OF APDL
! COMMANDS)
*ELSEIF,VAL1,Oper1,VAL2 ! COMPARISON-2 (OPTIONAL)
... ! APDL-2 (ANY NUMBER OF APDL
! ! COMMANDS)
*ELSE ! COMPARISON-3 (OPTIONAL)
... ! APDL-3 ANY NUMBER OF APDL
! COMMANDS
* ENDIF
```

There may be several *elseif commands. *elserf command usage is the same as for the *IF command (as far as the arguments are concerned) whereas the *ELSE command does not have any arguments. There can only be one *ELSE command, and it is the last IF-THEN-ELSE block command before the *ENDIF command. In the example above, note that:

If COMPARISON-1 is true, then the input block APDL-1 is executed and the input blocks APDL-2 and APDL-3 are ignored. If COMPARISON-1 is false and if COMPARISON-2 is true, then the input block APDL-2 is executed and the input blocks APDL-1 and APDL-3 are ignored. Finally, if neither COMPARISON-1 nor

COMPARISON-2 is true, then the input block APDL-3 is executed and the input blocks APDL-1 and APDL-2 are ignored.

The arithmetic average of $x$-displacements along the boundary defined by $x=-2.5$ was evaluated in the example considered in Sec. 7.5.1. In order to demonstrate the use of IF-THEN-ELSE blocks, the example is modified by computing the arithmetic averages of positive and negative $x$-displacements separately, and the number of nodes with zero $x$-displacement along the boundary specified as $x=-2.5$. This task can be performed by the following input block:

```
/POST1 ! ENTER POSTPROCESSOR
NSEL,S,LOC,X,-2.5 ! SELECT NODES ALONG
! X = -2.5
*GET, numnod, NODE,0, COUNT ! RETRIEVE NUMBER OF NODES
*GET,minnod,NODE,0,NUM,MIN ! RETRIEVE MINIMUM NODE
                                    ! NUMBER
sum_p=0 : INITIALIZE SUM OF POSITIVE DISPLACEMENTS
sum_n=0 ! INITIALIZE SUM OF NEGATIVE DISPLACEMENTS
cnt_p=0 ! INITIALIZE # OF NODES WITH POSITIVE
    ! DISPLACEMENTS
cnt_n=0 ! INITIALIZE # OF NODES WITH NEGATIVE
    ! DISPLACEMENTS
cnt_z=0 ! INITIALIZE # OF NODES WITH ZERO DISPLACEMENTS
curnod=minnod ! INITIALIZE CURRENT NODE
    NUMBER
*DO,ii,1,numnod !BEGIN DO LOOP
*GET,cux,NODE, curnod,U,X ! RETRIEVE X-DISPLACEMENT
                                    ! OF THE CURRENT NODE
*IF,cux,GT,0,THEN ! BEGIN IF-THEN-ELSE BLOCK
sum_p=sum_p+cux ! UPDATE SUM FOR POSITIVE
    ! DISPLACEMENTS
cnt_p=cnt_p+1 ! UPDATE NUMBER OF NODES
*ELSEIF,cUX,LT,0 ! cux IS NEGATIVE
sum_n=sum_n+cux !UPDATE SUM FOR NEGATIVE
! DISPLACEMENTS
cnt_n=cnt_n+1 ! UPDATE NUMBER OF NODES
*ELSE ! CUX IS ZERO
cnt_z=cnt_z+1 !UPDATE NUMBER OF NODES
*ENDIF ! END IF-THEN-ELSE BLOCK
\begin{tabular}{ll} 
*GET, nextnod,NODE, curnod, NXTH & ! STORE NEXT HIGHER NODE \\
& ! NUMBER IN nextnod \\
curnod=nextnod & ! UPDATE CURRENT NODE \\
*ENDDO & ! END DO LOOP \\
ave_d_p=sum_p/cnt_p & ! CALCULATE AVERAGE \\
ave_d_n=sum_n/cnt_n & ! POSITIVE \\
& ! CALCULATE AVERAGE
\end{tabular}
```

The arithmetic averages of positive and negative $x$-displacements are stored in parameters ave_d_p and ave_d_n, respectively. Also, the number of nodes with zero $x$-displacement is stored in parameter ent_z.

### 7.5.3 /OUTPUT and *VWRITE Commands

APDL offers the option of writing formatted output to text files through use of /OUTPUT and *VWRITE commands. The /OUTPUT command redirects the text output, normally written in the Output Window, to a text (ASCII) file whereas the *VWRITE command allows desired parameters to be written with FORTRAN (or C) formatting.

The syntax for the /OUTPUT command is

```
/OUTPUT, Fname, Ext, ,Loc
```

in which Fname and Ext are the file name and extension, respectively, and Loc decides whether to start writing from the top of this file or to append to it. If the field for Loc is left blank, then the output is written from the top of the file. If the value of LOC is specified as APPEND, then the output is appended. Once the desired data are written to the text file, the output should be redirected back to the Output Window using the same command with no arguments, i.e.,

```
/OUTPUT
```

The syntax for the *VWRITE command is

```
*VWRITE,Par1,Par2,...,Par19
```

in which Par1 through Par19 are the parameters to be written with formatting. As observed, up to 19 parameters can be written at a time. A FORTRAN or C format can be used and must be supplied in the next line. The FORTRAN format must be enclosed in parentheses and only real or alphanumeric formatting is allowed.

When appended to the input block given in Sec. 7.5.2, the commands in the following input block write the arithmetic averages of positive and negative $x$-displacements, and the number of nodes with zero $x$-displacement along the boundary of $x=-2.5$ to three parameters (ave_d_p, ave_d_n, and cnt_z) in a text file named data.out:

```
/OUTPUT,data,out ! REDIRECT OUTPUT TO FILE
*VWRITE,ave_d_p,ave_d_n ! WRITE PARAMETERS ON THE
! SAME LINE
```

```
(E15.8,2X, E15.8)
/OUTPUT
/OUTPUT,data,out, , APPEND
*VWRITE,cnt_z
(F8.0)
/OUTPUT
```

```
! FORMAT STATEMENT
! REDIRECT OUTPUT BACK TO
! OUTPUT WINDOW
! APPEND TO EXISTING FILE
! WRITE THE PARAMETER
! FORMAT STATEMENT
! REDIRECT OUTPUT BACK TO
! OUTPUT WINDOW
```

In this particular example, E15.8 in the format statement allocates 15 spaces for the parameter, 8 of which are used for the numbers after the decimal point. The 2 x enforces 2 blank spaces between the parameters. The parameters ave_d_p and ave_d_n are written in the following format:

```
\pm0.12345678E\pm00 士0.12345678E\pm00
```

Similarly, the format statement F8.0 allocates a total of 8 spaces for the parameter with no space for the numbers after the decimal point, and the parameter cnt_z must be an integer.

### 7.6 Macro Files

A more advanced level of APDL use is the Macro Files, which are similar to subroutines in the FORTRAN programming language. Macro Files are saved in separate text files with the file extension mac (e.g. macroI.mac) and written using the APDL. If they are saved in the Working Directory, they are automatically recognized by the ANSYS program. Otherwise, the user must declare their location using the /PSEARCH command. They are particularly useful for tasks that are repeated many times with different values of model variables such as geometry, material properties, boundary conditions, etc. A simple example on how Macro Files are used is given below.

The example under consideration involves the modeling of a spring that has a helix shape. The user needs to generate several models with different geometric properties as part of a design requirement. The coordinates of a point on the helix are given by the following set of parametric equations:

$$
\begin{align*}
& x=a \cos (t) \\
& y=a \sin (t)  \tag{7.1}\\
& z=b t
\end{align*}
$$

in which $a$ is the radius of the helix as it is projected onto the $x-y$ plane, $2 \pi b$ is the distance in the $z$-direction of one full turn, and $t$ is the independent parameter. For this purpose, two Macro Files are written, with names HELIX1.MAC and HELIX2.MAC, as given below:

```
! HELIX1.MAC
! MACRO FOR HELIX GENERATION
! ARG1 : COEFFICIENT A IN EQ. 7.1
! ARG2 : COEFFICIENT B IN EQ. 7.1
! ARG3 : NUMBER OF SEGMENTS TO BE USED FOR QUARTER CIRCLE
! ARG4 : NUMBER OF HELIX STEPS
/PREP7 ! ENTER PREPROCESSOR
PI=4*ATAN(I) ! DEFINE PI
T=0 ! INITIAL VALUE OF T
DT=2*PI/(4*ARG3-1) ! INCREMENT OF T
*DO,I,1,4*ARG3
K, ,ARG1*COS (T),ARG1*SIN(T),ARG2*T
T=T+DT
* ENDDO
HELIX2,ARG3 ! CALL MACRO HELIX2
LGEN,ARG4,ALL, , , , , 2*PI*ARG2
/EOF ! MARK END OF FILE
! HELIX2.MAC
! CALLED BY HELIX1.MAC
! ARG1 : NUMBER OF SEGMENTS TO BE USED FOR QUARTER CIRCLE
KSEL,S,KP, 1,ARG1+1
BSPLIN,ALL
KSEL, S, KP, , ARG1+1, 2*ARG1+1
BSPLIN,ALL
KSEL, S,KP, , 2*ARG1+1,3*ARG1 +1
BSPLIN,ALL
KSEL, S,KP, , 3*ARG1+1,4*ARG1 +1
BSPLIN,ALL
ALLSEL
LGLUE,ALL
/EOF
```

As long as these files are located in the Working Directory, issuing the following command produces the helix shown in Fig. 7.9 (oblique view):

```
HELIX1,1,0.1,4,4
```

Note that the arguments are specified as $a=1$ and $b=0.1$, and four segments are used in creating a quarter circle. Finally, the geometry possesses a total of 4 helix steps. When the radius is modified to be $a=0.5$, and the number of helix steps is increased to 10 using

HELIX1, 0.5, 0.1, 4, 10
the geometry shown in Fig. 7.10 is obtained.


Fig. 7.9 Helix created upon execution of user-defined macro HELIX1 with arguments $1,0.1,4$, and 4 .


Fig. 7.10 Helix created upon execution of user-defined macro HELIX1 with arguments $\mathbf{0 . 5 , 0 . 1 , 4}$, and 10 .

### 7.7 Useful Resources

There are three main resources for help in enhancing and accelerating knowledge of and skill in programming with APDL:

ANSYS Help System
Log File
ANSYS Verification Manual

The first topic is discussed in sufficient detail in Sec. 2.7. The following subsections briefly discuss the second and third topics.

### 7.7.1 Using the Log File for Programming

Every time ANSYS is used interactively (using GUI), a Log File is created in the Working Directory with the name jobname.log. If Jobname is not specified, the default for the Jobname is file and the Log File is named file.log. This file records every single action the user takes when using ANSYS, including the ones that are not directly related to the finite element method, such as graphics (zoom in/out, turning on/off entity numbering in the Graphics Window, etc.). Although it may appear to be a little "messy," it is extremely useful in learning which commands are used for certain actions when using GUI.

The following example illustrates how the user can utilize the Log File for learning certain commands. Suppose the domain is the same as the one considered in Sec. 7.2 and the user is required to write an input file to create the solid model. The model generation includes:

Creation of a square with side length of 2 .
Creation of a circle with radius 1 .
Subtraction of the circle from the square.
The details on how to perform these simple operations in GUI are left out as they should be clear based on the coverage in Chap. 4 and several examples throughout this book. Once the user performs these operations using the GUI, the Log File can be viewed from within ANSYS using

Utility Menu > List > Files > Log File
which should appear in a separate Output Window with contents as given below:

```
/ BATCH
/COM, ANSYS RELEASE 8.0 UP20030930 14:49:33 06/16/2004
/ PREP7
RECTNG, 2, ,2,
PCIRC,1, ,0,360,
ASBA, 1, 2
```

Creation of the square and the circle is achieved by using the commands RECTNG and PCIRC, respectively. Finally, area subtraction is performed using the ASBA command. Once the command names are identified, Help Pages for these commands must be read to learn the correct and most efficient usage. The Log File may not always be as clear-cut as the one shown above, especially when the GUI action involves graphical picking. However, as the user becomes more accustomed to the use of ANSYS, both in Batch and Interactive Modes, the Log File becomes easier to follow.

### 7.7.2 Using the Verification Problems for Programming

As mentioned in Sec. 2.7, the Verification Manual under the ANSYS Help System provides an excellent platform for programming in APDL. In order to demonstrate this point, Verification Problem 2 (one of the 238 problems) is selected. In this problem, a simply supported I-beam with known properties is subjected to a uniformly distributed transverse loading, as shown in Fig. 7.11. The help page for this problem can be viewed by using the following menu path within the ANSYS Help System, which appears on the left side of the Help Window (heading shown in Fig. 7.12):

## ANSYS Verification Manual > Verification Test Case Descriptions > VM2

The problem description, a sketch of the problem and corresponding FEA representation, the reference from which the problem is taken, and analysis assumptions are found at the top of the page. Included further down is a table (see Fig. 7.13) showing the results obtained by both analytical methods and the ANSYS software. As observed in Fig. 7.12, there is a hyperlink to the text file vm2. dat, which includes the input commands for the solution of this problem using the Batch Mode. Upon clicking on this hyperlink, the file appears as partially shown in Fig. 7.14. The user can go through this file line by line, referring frequently to the help pages of unfamiliar commands in order to learn the correct usage of commands.

Another important benefit from the Verification Manual is that one can learn how to solve problems with certain properties. The Verification Test Case Descriptions help page, accessed through the menu path

ANSYS Verification Manual > Verification Test Case Descriptions
is a good place to start. For example, if the problem at hand involves materials with viscoelastic behavior, it would be a good idea to scan the test case descriptions to find a solved problem with such materials. A quick glance at the list of test case descriptions reveals that Verification Problem 200 involves a viscoelastic material and that the user may benefit from examining this file in order to see how the problem is treated before moving on to the problem at hand, which is likely to be more complicated.


Fig. 7.11 Graphical description of Verification Problem 2 as given in ANSYS Verification Manual.

VM2
Beam Stresses and Deflections

## Overview

| Reference: | Timoshenko, [3.], pg. 98, problem 4 |
| :--- | :--- |
| Analysis Type(s): | Static Analysis (ANTYPE = 0) |
| Element Type(s): | 2-D Elastic Beam Elements (BEAM3) |
| Input Listing: | vm2. dat |

Fig. 7.12 Heading of the help page for Verification Problem 2 as given in ANSYS Verification Manual.

## Results Comparison

|  | Target | ANSYS | Ratio |
| :--- | :---: | :---: | :---: |
| Stress, psi | -11.400 | -11.404 | 1.000 |
| Deflection, in | 0.182 | 0.182 | 1.003 |

Fig. 7.13 Comparison of results as given in ANSYS Verification Manual.

## VM2 Input Listing

```
/COK, NNSYS MEDIL REL. 8.0 (9-17-2003) RET. VERIF, MANULL: REL. 8.0
/VERIFY,VE2
JPGPRT,500,100,1 : HACRO TO SET PRETS FOR JPEG PLOTS
/SHOW,JPEG
/PREP7
HP,PRXY, ,0.3
/TITLE, VH2, BEAK STRESSES NND DEFLECTIONS
C*** STR. OF KATL., TIMOSHENKO, PART 1, 3RD ED., PAGE 98, PROB. 4
antype,static
ET, 1, BEAM3
KEYOPT,1,9,9 I OUTPUT AT 9 intermEdIATE LOCations
R,1,50.65,7892,30
KP,EX, 1,3OE6
H,1
N,5,480
TILL
I,1,2
IGEN, 4, 1,1
D,2,UX,\ldots..UY
D,4,UY
STBEAK, 1, 1, PRES, (10000/12)
SFBEAK, 4, 1, PRES, (1E4/12)
FINISH
/SOLU
```

Fig. 7.14 VM2 input listing as given in ANSYS Verification Manual.

## Chapter 8

## LINEAR STRUCTURAL ANALYSIS

A linear analysis is conducted if a structure is expected to exhibit linear behavior. The deformation and load-carrying capability can be determined by employing one of the analysis types available in ANSYS, static or dynamic, depending on the nature of the applied loading. If the applied loading is determined as part of the solution for structural stability, a buckling analysis is conducted. If the structure is subjected to thermal loading, the analysis is referred to as thermomechanical.

### 8.1 Static Analysis

The behavior of structures under static loading can be analyzed by employing different types of elements within ANSYS. The nature of the structure dictates the type of elements utilized in the analysis. Discrete or framed structures are suitable for modeling with rod- and beam-type elements. However, the modeling of continuous structures usually requires a threedimensional model with solid elements.

Under certain types of loading and geometric conditions, the three-dimensional type of analysis can be idealized as a two-dimensional analysis. If the component is subjected to in-plane loading only and its thickness is small with respect to the other length dimensions, it is idealized as a plane stress condition. If the component with a uniform cross section is long in the depth direction and is subjected to a uniform loading along the depth direction, it is idealized as a plane strain condition. If the component has a circular cross section and is subjected to uniform and concentric loading, it possesses axisymmetry. If thin structural components are subjected to lateral loading, the plate and shell elements are suitable for analysis.

### 8.1.1 Trusses

A truss is a structure that is made of straight structural members capable of carrying loads only in their own direction, i.e., no shear forces, no moments. Thus, each member is under either axial tension or axial compression. These members are connected to each other by means of joints. It is assumed that
loads can only be applied at the joints. ANSYS provides several element types for modeling truss structures. They include LINK1 and LINK160 for two- and three-dimensional analyses, respectively. The degrees of freedom at each node for truss elements are the displacement components ( $u_{x}$ and $u_{y}$ for 2-D; $u_{x}, u_{y}$, and $u_{z}$ for 3-D). However, the vector sum of the deformations (elongation or contraction, not the displacements) is aligned with the direction of the element. Two example problems are given to demonstrate the usage of truss elements within ANSYS.

### 8.1.1.1 Elongation of a Bar under its Own Weight Using Truss Elements

Consider a steel bar of uniform cross section whose upper end is supported such that it is fixed from translational motion. The mass density, elastic modulus, and Poisson's ratio of steel are $\rho=0.284 \mathrm{lb} / \mathrm{in}^{3}, E=30 \times 10^{6} \mathrm{psi}$, and $v=0.3$, respectively. The radius and length of the bar are assumed to be $r=2 \mathrm{in}$ and $l=20 \mathrm{in}$, respectively, and the gravitational acceleration is $g=386.2205 \mathrm{in} / \mathrm{sec}^{2}$. The goal is to find the elongation of the bar at the lower end due to its own weight. The positive $y$-direction is the opposite direction of the gravitational acceleration, as shown in Fig. 8.1.

This problem can be solved using two-dimensional truss, two-dimensional axisymmetric plane, or three-dimensional elements. Since, we are interested in the elongation only, two-dimensional truss elements (LINK1) are used to obtain the solution.

## MODEL GENERATION

- Specify the element type (ET command) using the following menu path:


## Main Menu > Preprocessor > Element Type > Add/EditDelete

- Click on Add.
- Select Link from the left list and 2D Spar 1 from the right list; click on OK.
- Click on Close.
- Specify real constants ( $R$ command) using the following menu path:

Main Menu > Preprocessor > Real Constants > Add/Edit/Delete

- Click on $A d d$.
- Highlight Type 1 Link 1; click on OK.
- Enter 12.5664 (calculated based on radius, $r=2$ in ) for $A R E A$; click on $O K$.
- Click on Close.


Fig. 8.1 Schematic of a bar deformed due to its own weight.

- Specify material properties for the bar (MP command) using the following menu path:


## Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural and Density, which will bring up another dialog box.
- Enter 0.284 for $\boldsymbol{D E N S}$; click on $\boldsymbol{O K}$.
- In order to specify the elastic modulus and Poisson's ratio, in the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring up another dialog box.
- Enter 30e6 for $\boldsymbol{E X}$ and $\mathbf{0 . 3}$ for PRXY; click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create keypoints ( K command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Keypoints $>\operatorname{In}$ Active CS

- A total of 2 keypoints will be created.
- Enter $(x, y)$ coordinates of keypoint 1 as $(0,0)$; click on Apply. This action will keep the Create Keypoints in Active Coordinate System dialog box open. If the NPT Keypoint number field is left blank, then

ANSYS assigns the lowest available keypoint number to the keypoint that is being created.

- Repeat the same procedure for keypoint 2 using $(0,-20)$ for the $(x, y)$ coordinates.
- Click on $\boldsymbol{O K}$ (instead of Apply).
- Create a line ( $\mathbf{L}$ command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Lines > Lines > Straight Line

- Pick Menu appears; first pick keypoint 1 , then keypoint 2; click on $\boldsymbol{O K}$.
- Specify the number of divisions on the line (LESIZE command) using the following menu path:

Main Menu > Preprocessor > Meshing > Size Cntris > ManualSize > Lines > Picked Lines

- Pick Menu appears; pick the line; click on $O K$.
- Element Sizes on Picked Lines dialog box appears; enter 20 for NDIV; click on $\mathbf{O K}$.
- Create the mesh (LMESH command) using the following menu path:

Main Menu > Preprocessor > Meshing > Mesh > Lines

- Pick Menu appears; pick the line; click on $\boldsymbol{O K}$.
- Review elements.
- Turn on element numbering using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Select Element numbers from the first pull-down menu.
- Plot elements (EPLOT command) using the following menu path:

Utility Menu > Plot > Elements

- Turn off element numbering and turn on node numbering using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Place a checkmark by clicking on the empty box next to NODE Node numbers.
- Select No numbering from the first pull-down menu.
- Click on OK.
- Plot nodes (NPLOT command) using the following menu path: Utility Menu > Plot > Nodes


## SOLUTION

- Apply displacement constraints (D command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; pick node 1 (upper end); click on $\boldsymbol{O K}$ in the Pick Мепи.
- Highlight All DOF; click on OK.
- Apply gravitational acceleration (ACEL command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Inertia > Gravity

- Apply (Gravitational) Acceleration dialog box appears.
- Enter 386.2205 for $\boldsymbol{A C E L Y}$; click on $\boldsymbol{O K}$.

Obtain the solution (SOLVE command) using the following menu path:
Main Menu $>$ Solution $>$ Solve $>$ Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on $\boldsymbol{O K}$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review displacement values (PRNSOL command) using the following menu path:


## Main Menu > General Postproc > List Results > Nodal Solution

- Highlight DOF solution in the left list and Translation $\boldsymbol{U Y}$ in the right list; click on $O K$.
- The list appears. Note that the value for the $y$-displacement at node 2 (lower end) is listed as $\mathbf{- 0 . 7 3 1 1 4 E - 0 3}$ (in inches).


### 8.1.1.2 Analysis of a Truss Structure with Symmetry

Consider the steel truss structure shown in Fig. 8.2, which possesses symmetry with respect to the ordinate. Node and element numbers are also shown in this figure. Element 3 has a cross-sectional area of $A=20 \mathrm{in}^{2}$, while the other elements have $A=10 \mathrm{in}^{2}$. The elastic modulus for all of the elements is $E=30 \times 10^{6} \mathrm{psi}$. The goal is to find the displacements at the nodes and the stresses in the elements. Due to the symmetry condition, only


Fig. 8.2 Schematic of the truss structure with symmetry.
half the geometry is modeled with appropriate boundary conditions, i.e., the $x$-displacement at nodes 2 and 3 is zero and the applied force at node 2 is halved. Also, for the element located along the symmetry line, one half of the cross-sectional area is used.

The solution obtained using ANSYS is as follows:

## MODEL GENERATION

- Specify the element type (ET command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Element Type $>$ Add/Edit/Delete

- Click on Add.
- Select Link from the left list and 2D Spar 1 from the right list; click on OK.
- Click on Close.
- Specify real constants ( R command) using the following menu path:

Main Menu > Preprocessor > Real Constants > Add/Edit/Delete

- Click on Add.
- Highlight Type 1 Link 1; click on OK.
- Enter 10 for $\boldsymbol{A R E A}$; click on $\mathbf{O K}$.
- Click on Close.
- Specify material properties for the bar (MP command) using the following menu path:


## Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring up another dialog box.
- Enter 30e6 for $\boldsymbol{E X}$; click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create nodes ( $\mathbf{N}$ command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Nodes > In Active CS

- A total of 3 nodes will be created.
- Enter the $(x, y)$ coordinates of node 1 as $(0,0)$; click on Apply.
- Repeat the same procedure for nodes 2 and 3 using $(96,0)$ and $(96,72)$, respectively, for the $(x, y)$ coordinates.
- After entering the coordinates for node 3, click on $\boldsymbol{O K}$ (instead of Apply).
- A total of 3 elements will be created. Element 1 is defined by nodes 1 and 3 [1-3]. Similarly, elements 2 and 3 are defined by nodes [1-2] and [2-3], respectively. Create elements ( $\mathbf{E}$ command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Elements > Auto Numbered > Thru Nodes

- Pick Menu appears; create elements by picking two nodes at a time and clicking on Apply in between.
- Observe the elements created after clicking on Apply in the Pick Menu.
- Repeat until element 3 is created; click on $O K$.


## SOLUTION

- Apply displacement constraints (D command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Мепи appears; pick node 1 ; click on $O K$ in the Pick Menu.
- Highlight $\boldsymbol{U Y}$; click on Apply.
- Pick Menu reappears; pick nodes 2 and 3; click on $O K$ in the Pick Menu.
- Click on $\boldsymbol{U Y}$ to remove the highlight then click on $\boldsymbol{U} \boldsymbol{X}$ to highlight.
- Click on $\boldsymbol{O K}$.

Apply force boundary conditions ( $\mathbf{F}$ command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Force/Moment > On Nodes

- Pick Menu appears; pick node 2; click on $O K$ in the Pick Menu.
- Select $\boldsymbol{F Y}$ from pull-down menu and enter -10000 for Force/moment value; click on OK.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status, if OK, close the Status Report Window; click on $O K$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review displacement values (PRNSOL command) using the following menu path:

Main Menu > General Postproc $>$ List Results $>$ Nodal Solution

- List Nodal Solution dialog box appears. Highlight DOF solution in the left list and Translation $U Y$ in the right list; click on $O K$.
- The list appears with the values for the $y$-displacement at nodes 2 and 3 as $-0.19200 \mathrm{E}-01$ and $-0.16800 \mathrm{E}-01$, respectively.
- Review element stress values (etable command) using the following menu path:


## Main Menu > General Postproc > Element Table > Define Table

- Element Table Data dialog box appears. Click on Add, which brings up the Define Additional Element Table Items dialog box. Enter a label (Lab) for element stresses, say ELSTRS. Scroll down in the left list; click on By Sequence num; click on $\boldsymbol{L S}$ in the right list. Finally, enter $L S, 1$ in the last text field, as shown in Fig. 8.3; click on $\boldsymbol{O K}$.
- Note that the element table ELSTRS is now listed in the Element Table Data dialog box; click on Close.
- List the element table (pretab command) using the following menu path:

Main Menu > General Postproc > Element Table > List Elem Table

- In the List Element Table Data dialog box, highlight ELSTRS; click on OK.
- The list appears with stresses in elements 1,2 , and 3 as -1666.7 , 1333.3 , and 1000, respectively.


Fig. 8.3 Dialog box for retrieving element results based on sequence numbers.

### 8.1.2 Beams

A beam is a structural member capable of carrying axial, shear, and moment loads. Unlike truss members, loads can be applied anywhere along the beam geometry. ANSYS provides several element types for modeling beams. The most commonly used ones are beam3 and beam4 for two- and threedimensional analyses, respectively. At each node, both displacements and rotations are the degrees of freedoms for structural beam elements ( $u_{x}, u_{y}$, and $\theta_{z}$ for 2-D; $u_{x}, u_{y}, u_{z}, \theta_{x}, \theta_{y}$, and $\theta_{z}$ for 3-D). Two example problems are considered in this section for the demonstration of the usage of beam elements within ANSYS.

### 8.1.2.1 Analysis of a Slit Ring

A circular steel ring with a slit, as shown in Fig. 8.4, is subjected to a $50-\mathrm{lb}$ vertical force acting in the negative $y$-direction at the termination point while translations and rotations are constrained in every direction. The ring


Fig. 8.4 Schematic of a circular steel ring and the corresponding solid model.
has a solid circular cross section with radius 1 in . The structure is modeled using beam elements with cross-sectional area $A=\pi$, elastic modulus $E=30 \times 10^{6}$, Poisson's ratio $v=0.3$, and moment of inertia $I_{z z}=\pi / 4$. The goal is to find the displacements at the nodes and the moment diagram. The solid model used in the ANSYS solution is also shown in Fig. 8.4, with the keypoint and line numbers indicated.

## model generation

- Specify the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Beam from the left list and 3D Elastic 4 from the right list; click on $O K$.
- Click on Close.
- Specify real constants ( R command) using the following menu path:

Main Menu > Preprocessor > Real Constants > Add/Edit/Delete

- Real Constants dialog box appears; click on Add.
- Highlight Type 1 Beam 4; click on OK.
- Enter 3.14159265 for AREA and 0.785398 for IZZ and IYY; click on OK.
- Exit from the Real Constants dialog box by clicking on Close.
- Specify material properties for the beam (MP command) using the following menu path:


## Main Menu > Preprocessor > Material Props $>$ Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring up another dialog box.
- Enter $\mathbf{3 0 e} \boldsymbol{6}$ for $\boldsymbol{E X}$ and $\mathbf{0 . 3}$ for $\boldsymbol{P R X Y}$; click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create keypoints ( K command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Keypoints > In Active CS

- A total of 6 keypoints will be created.
- Enter the $(x, y)$ coordinates of keypoint 1 as (100, 0); click on Apply. This action will keep the Create Keypoints in Active Coordinate System dialog box open. If the NPT Keypoint number field is left blank, then ANSYS assigns the lowest available keypoint number to the keypoint that is being created.
- Referring to Fig. 8.4, repeat the same procedure for keypoints 2, 3, 4, 5, and 6 using ( $0,-100$ ), $(-100,0),(0,100),(100,0)$, and (0,0), respectively, for the $(x, y)$ coordinates.
- After generating keypoint 6 , click on $O K$ (instead of Apply).
- Note that keypoints 1 and 5 are coincident. This is intentional, so the slit can be modeled properly.
- Create arcs (LARC command) using the following menu path:


## Main Menu > Preprocessor > Modeling > Create > Lines > Arcs > By End KPs \& Rad

- A total of 4 lines (arcs) will be created.
- Pick Menu appears; pick keypoints 1 and 2 (end points of the arc); click on $O K$ in the Pick Menu.
- Pick keypoint 6 (center of the arc); click on $\boldsymbol{O K}$ in the Pick Menu.
- Arc by End KPs \& Radius dialog box appears; enter 100 for RAD Radius of the arc.
- Click on Apply; line 1 is created.
- Repeat this procedure for lines 2,3 , and 4 using keypoint pairs ( 2,3 ), $(3,4)$, and $(4,5)$, respectively. All lines use keypoint 6 as the center and 100 as the radius.
- Specify the number of divisions on all lines (LESIZE command) using the following menu path:

Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Lines > All Lines

- Element Sizes on All Selected Lines dialog box appears; enter 10 for NDIV; click on OK.
- Create the mesh (LMESH command) using the following menu path:

Main Menu > Preprocessor > Meshing > Mesh > Lines

- Pick Menu appears; click on Pick All.


## SOLUTION

- Apply displacement constraints (D command) using the following menu path:
- Pick Menu appears; pick one of the nodes at $x=100$ and $y=0$. There are two nodes at this location: nodes 1 and 32 . When picking, ANSYS asks the user which one of the nodes is to be picked. Click on the Next button in this Warning Window so that it shows Node 32; click on OK in the Pick Menu.
- Apply U, Rot on Nodes dialog box appears; highlight All DOF; click on OK.
- Apply force boundary conditions (F command) using the following menu path:

Main Menu > Solution $>$ Define Loads $>$ Apply $>$ Structural $>$ Force/Moment $>$ On Nodes

- Pick Menu appears; this time pick node 1 (instead of node 32); click on OK in the Pick Menu.
- Select $\boldsymbol{F Y}$ from the pull-down menu and enter $\mathbf{- 5 0}$ for Force/moment value; click on $O K$.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu > Solution > Solve >Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the deformed shape (PLDISP command) using the following menu path:

Main Menu $>$ General PostProc $>$ Plot Results $>$ Deformed Shape

- Select Def + undeformed; click on OK.
- The deformed shape is shown in Fig. 8.5 as it appears in the Graphics Window.
- Store bending moment values in the element table (ETABLE command) using the following menu path:


## Main Menu > General Postproc > Element Table > Define Table

- Element Table Data dialog box appears; click on Add.
- Define Additional Element Table Items dialog box appears; in the left list, scroll down to select By sequence number and select SMISC in the right list. Finally, type SMISC,6 in the last text field; click on $O K$ (Fig. 8.6).


Fig. 8.5 Derormea snape or the stees rong unaer applied boundary conditions.


Fig. 8.6 Define Additional Element Table Items dialog box for extracting nodal moment values.

- Note that SMIS6 now appears in the list in the Element Table Data dialog box. Exit from the Element Table Data dialog box by clicking on Close.
- Plot the moment diagram (PLLS command) using the following menu path:
Main Menu > General Postproc > Plot Results > Contour Plot > Line Elem Res
- Plot Line-Element Results dialog box appears; click on OK.
- Figure 8.7 shows the resulting moment diagram as displayed in the Graphics Window.


Fig. 8.7 Moment diagram of the steel ring.

### 8.1.3 Three-dimensional Problems

Almost all engineering problems are three-dimensional (3-D) by nature. However, depending on the specific geometry, loading conditions, and quantities of interest, it is common to approach the problem with the idealization of a lower dimensionality. If a representative idealization cannot be utilized, then a three-dimensional model must be created. The most commonly used three-dimensional structural element is SOLID45, which is an 8 -noded brick element. The degrees of freedom at each node for 3-D problems are $u_{x}, u_{y}$, and $u_{z}$. Determining the deformation of a bar under its own weight using three-dimensional elements in ANSYS is demonstrated in the following.

The elongation of a bar due to its own weight was modeled in Sec. 8.1.1.1 using two-dimensional link elements. That solution provided the displacement of the bar in the longitudinal direction, and the same crosssectional area is assumed. Three-dimensional elements provide the change in the cross-sectional area, as well as the displacement components. The reference frame shown in Fig. 8.8 is used in the 3-D solution.

## MODEL GENERATION

- Specify the element type (ET command) using the following menu path:


## Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Structural Solid from the left list and Brick 8node 45 from the right list; click on $\boldsymbol{O K}$.
- Click on Close.


Fig. 8.8 Schematic of a bar deformed due to its own weight.

- Specify material properties for the bar (MP command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Material Props $>$ Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural and Density, which will bring up another dialog box.
- Enter 0.2839605 for DENS; click on OK.
- In order to specify the elastic modulus and Poisson's ratio, in the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring up another dialog box.
- Enter 30e6 for $\boldsymbol{E X}$ and $\mathbf{0 . 3}$ for $\boldsymbol{P R X Y}$; click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create a volume (CYLIND command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Volumes > Cylinder > By Dimensions

- Create Cylinder by Dimensions dialog box appears. Enter 2 for RAD2, 20 for Z2, and 90 for THETA2; click on OK.

■ Create additional volumes by reflection (VSYMM command) using the following menu path:

Main Menu > Preprocessor > Modeling > Reflect > Volumes

- Pick Меnu appears; click on Pick All button, which brings up the Reflect Volumes dialog box.
- Click on the $\boldsymbol{Y}$ - $\boldsymbol{Z}$ Plane $X$ radio-button; click on Apply.
- Pick Menu reappears; click on Pick All button and in the Reflect Volumes dialog box click on the $\boldsymbol{X}-\boldsymbol{Z}$ Plane $\boldsymbol{Y}$ radio-button; click on OK.
m Glue the volumes (VGLUE command) using the following menu path:
Main Menu > Preprocessor > Modeling > Operate > Booleans > Glue > Volumes
- Pick Menu appears; click on Pick All button.
- Specify the global element size (ESIZE command) using the following menu path:

Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Global > Size

- Global Element Sizes dialog box appears; enter 1 for SIZE; click on OK.
- Create the mesh (VMESH command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Volumes $>$ Mapped $>4$ to 6 sided

- Pick Menu appears; click on Pick All.


## SOLUTION

- Apply displacement constraints (D command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; pick the nodes at $z=0$ plane (use different viewpoints if necessary); click on $\boldsymbol{O K}$ in the Pick Menu.
- Highlight All DOF; click on OK.
- Apply gravitational acceleration (ACEL command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Inertia > Gravity

- Apply (Gravitational) Acceleration dialog box appears.
- Enter -386.2205 for $A C E L Z$; click on OK.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status, if OK, close the Status Report Window; click on $O K$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the deformed shape (PLDISP command) using the following menu path:

Main Menu > General PostProc > Plot Results > Deformed Shape

- Select Def + undef edge; click on $\boldsymbol{O K}$.
- The deformed shape is shown in Fig. 8.9 as it appears in the Graphics Window.
- Review $z$-displacement contours (PLNSOL command) using the following menu path:

Main Menu > General PostProc > Plot Results > Contour Plot > Nodal Solu

- Contour Nodal Solution Data dialog box appears. Select DOF solution from the left list and Translation UZ from the right list; click on $O K$.
- The contour plot is shown in Fig. 8.9 as it appears in the Graphics Window.


Fig. 8.9 Deformed shape (left) and contour plot of the $z$ displacement (right) of the bar due to its own weight.

- Review displacement values (PRNSOL command) using the following menu path:


## Main Menu > General Postproc > List Results > Nodal Solution

- Highlight DOF solution in the left list and Translation $U Z$ in the right list; click on $O K$.
- The list appears in a separate window. It is a long list of $z$ displacements.
- At the bottom of the window maximum displacement value is printed as $0.72274 E-03$.


### 8.1.4 Two-dimensional Idealizations

As mentioned in Sec. 6.2.2, the reduction of the dimensionality of a problem from three to two through an idealization may reduce the computational cost significantly. There are three distinct two-dimensional idealizations: plane stress, plane strain, and axisymmetry.

Plane stress and strain idealizations are discussed in Sec. 6.2.2.1 and 6.2.2.2, respectively. Therefore, the descriptions given in the following subsections are brief.

### 8.1.4.1 Plane Stress

In a structural problem, if one of the dimensions is much smaller than the inplane dimensions, and if the structure is subjected to only in-plane loads along the boundary, then the plane stress idealization is valid. It reduces the computational cost significantly without a loss of accuracy in the quantities of interest. Plane stress idealization is demonstrated by considering a plate with a circular hole and a composite plate under axial tension.

### 8.1.4.1.1 Analysis of a Plate with a Circular Hole

A square plate $\left(9 \times 9 \mathrm{in}^{2}\right)$ with a circular hole (radius $r=0.25 \mathrm{in}$ ) is subjected to uniformly distributed tensile loading ( 1000 psi ) in the vertical direction along its top surface while being fixed along the bottom surface (Fig. 8.10). The plate is stiffened by means of increased thickness, from 0.063 in to 0.12 in. Plane stress idealization is used in the ANSYS solution, as the plate is thin and there are no lateral loads. The material properties are given as elastic modulus $\mathrm{E}=10 \times 0^{6} \mathrm{psi}$ and Poisson's ratio $v=0.25$. The goal is to obtain the displacement and stress fields resulting from the applied boundary conditions.


Fig. 8.10 Geometry and loading of the plate with a circular hole.

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:


## Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Solid in the left list and Quad 4 Node 42 in the right list; click on $\boldsymbol{O K}$.
- Click on Options.
- PLANE42 element type options dialog box appears; select the Plane strs w/thk item from the pull-down menu corresponding to Element behavior K3.
- Click on OK; click on Close.
- Specify the thickness information using real constants ( R command) using the following menu path:


## Main Menu > Preprocessor > Real Constants > Add/Edit/Delete

- Real Constants dialog box appears; click on Add. Click on OK; Real Constants Set Number 1 for PLANE42 dialog box appears.
- Type 0.063 in the Thickness THK text field; click on Apply.
- Change the Real Constant Set No. from 1 to 2 and modify the Thickness THK text field to be 0.12 ; click on $O K$.
- Exit from the Real Constants dialog box by clicking on Close.
- Specify material properties (MP command) using the following menu path:


## Main Menu > Preprocessor > Material Props > Material Models

- Define Material Model Behavior dialog box appears. In the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which brings up another dialog box.
- Enter $\mathbf{1 0 e 6}$ for $\boldsymbol{E X}$ and $\mathbf{0 . 2 5}$ for PRXY; click on OK.
- Close the Define Material Model Behavior dialog box by using the following menu path:

```
Material > Exit
```

- Create a square area around the hole (RECTNG command) using the following menu path:
Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Rectangle $>$ By Dimensions
- In the Create Rectangle by Dimensions dialog box, enter 0 and 0.5 for $X 1$ and $X 2$ and 0 and 0.5 for Y1 and Y2; click on $O K$.
- Create a circular area for the hole geometry (PCIRC command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Circle $>$ By Dimensions

- In the Create Circle by Dimensions dialog box, type 0.25 for Outer radius; click on $\boldsymbol{O K}$.
- Subtract the circle from the rectangle (ASBA command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Operate $>$ Booleans $>$ Subtract $>$ Areas

- Pick Menu appears; pick the rectangle; click on $\boldsymbol{O K}$; pick the circle; click on $\boldsymbol{O K}$.
- Create additional rectangular areas (RECTNG command) using the following menu path:
Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Rectangle $>$ By Dimensions
- In the Create Rectangle by Dimensions dialog box, enter 0.5 and 4.5 for $X 1$ and X2 and 0 and 0.5 for Y1 and Y2; click on Apply.
- Now, enter 0.5 and 4.5 for $X 1$ and $X 2$ and 0.5 and 4.5 for $Y 1$ and $Y 2$; click on Apply.
- Finally, enter 0 and 0.5 for $X 1$ and $X 2$ and 0.5 and 4.5 for $Y 1$ and $Y 2$; click on $O K$.
- Glue the areas (AGLUE command) using the following menu path:

Main Menu > Preprocessor $>$ Modeling $>$ Operate $>$ Booleans $>$ Glue $>$ Areas

- Pick Menu appears; click on Pick All button.
- The areas appear in the Graphics Window, as shown in Fig. 8.11.
- Specify the global element size (ESIZE command) using the following menu path:
Main Menu $>$ Preprocessor $>$ Meshing $>$ Size Cntrls $>$ ManualSize $>$ Global $>$ Size
- Global Element Sizes dialog box appears; enter 0.1 for SIZE; click on OK.
- Specify the number of divisions on selected lines (LESIZE command) using the following menu path:
Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize $>$ Lines $>$ Picked Lines
- Pick Мепи appears; pick the two lines identified in Fig. 8.12; click on OK.
- Element Sizes on Picked Lines dialog box appears; enter 6 for NDIV and remove the checkmark next to KYNDIV SIZE,NDIV can be changed so that it shows No; click on $O K$.
- Concatenate lines (LCCAT command) using the following menu path:

Main Menu > Preprocessor > Meshing > Concatenate > Lines

- Pick Menu appears; pick the two lines identified in Fig. 8.12; click on OK.


Fig. 8.11 Areas after gluing operation.


Fig. 8.12 Number of divisions on identified lines.

- Create the mesh (AMESH command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 sided

- Pick Menu appears; click on Pick All.
- Modify the real constant set attribute of the elements corresponding to the thicker portion of the plate (EMODIF command) using the following menu path:
Main Menu > Preprocessor > Modeling > Move / Modify > Elements > Modify Attrib
- Pick Menu appears; pick the elements corresponding to the areas indicated in Fig. 8.13 (click on the Box radio-button in the Pick Menu and draw a rectangle in the Graphics Window to pick the elements). Clicking on $\boldsymbol{O K}$ brings up the Modify Elem Attributes dialog box.
- Select Real const REAL from the pull-down menu and enter $\mathbf{2}$ in the $\boldsymbol{I I}$ New attribute number field; click on $O K$.


Fig. 8.13 Element attributes on elements attached to identified areas.

■ Create two successive reflective symmetric meshes (ARSYM command) using the following menu path:

## Main Menu > Preprocessor > Modeling > Reflect > Areas

- Pick Menu appears; click on Pick All.
- Reflect Areas dialog box appears; click on the $\boldsymbol{Y}-Z$ plane $X$ radiobutton; click on Apply.
- Pick Menu reappears; click on Pick All.
- Plot elements (EPLOT command) using the following menu path:

Utility Menu > Plot > Elements

- Reflect Areas dialog box reappears; click on the $X-Z$ plane $Y$ radiobutton; click on $\boldsymbol{O K}$.
- Although it is not apparent through visual inspection, there are duplicate entities (keypoints, lines and nodes) along the symmetry lines, thus there is no continuity. Therefore, merge duplicate entities using the following menu path:

Main Menu $>$ Preprocessor $>$ Numbering Ctrls $>$ Merge Items

- In the dialog box, select $A l l$ from the first pull-down menu; click on OK.
- Plot elements with different colors based on their real constant numbers using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Plot Numbering Controls dialog box appears. Select Real const num from the first pull-down menu (corresponding to Elem / Attrib numbering) and select Colors only from the second pull-down menu (corresponding to [/NUM] Numbering shown with); click on OK.
- Figure 8.14 shows the corresponding element plot with different colors ${ }^{1}$ based on material numbers.


## SOLUTION

- Apply displacement constraints (D command) using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Structural $>$ Displacement $>$ On Nodes

- Pick Menu appears; pick the nodes along the bottom surface of the plate (click on the Box radio-button in the Pick Menu and draw a rectangle in the Graphics Window to pick the nodes); click on $\boldsymbol{O K}$ in the Pick Menu.
- Highlight both $\boldsymbol{U X}$ and $\boldsymbol{U Y}$; click on $\boldsymbol{O K}$.

[^9]

Fig. 8.14 Elements of the plate, as they appear in the Graphics Window.

- Apply surface force (pressure) boundary conditions (SF command) using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Structural $>$ Pressure $>$ On Nodes

- Pick Menu appears; pick the nodes along the top surface of the plate; click on $O K$ in the Pick Menu.
- Type -1000 (negative 1000) for VALUE Load PRES value; click on OK.
- Pressure, by definition, acts normal toward the body along the surface. The direction of action in reference to the global coordinate system does not affect whether it is positive or negative. The only factor that dictates the sign is whether it acts toward or away from the body. Therefore, in order to apply the tensile loading, it is necessary to apply negative pressure.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on $\boldsymbol{O K}$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the normal stress contour plots in the $x$ - and $y$-directions (PLNSOL command) using the following menu path:

Main Menu > General Postproc > Plot Results > Contour Plot $>$ Nodal Solu

- Highlight Stress in the left list (second item) and highlight $X$-direction $S X$ in the right list; click on $O K$.
- The contour plot of $\sigma_{x x}$ appears in the Graphics Window, as shown in Fig. 8.15.
- The Contour plot of $\sigma_{y y}$ is obtained similarly by highlighting $Y$ direction $S Y$ in the right list and clicking on $O K$ (shown in Fig. 8.16).
- Review the variation of stresses along a path by means of a line plot. This operation requires the path to be defined first, followed by mapping the solution items of interest onto the path and, finally, obtaining the plot. The path that is defined in this case lies along the positive $x$-axis, starting from the left boundary of the hole and ending at the left boundary of the plate. Define the path (PPATH command) using the following menu path:

Main Menu > General Postproc > Path Operations > Define Path > By Nodes

- Pick Menu appears; pick the nodes with $(x, y)$ coordinates $(0.25,0)$ and (4.5, 0); click on $\boldsymbol{O K}$.
- By Nodes dialog box appears; enter a name describing the path, say $h r z$, in the Define Path Name text field; click on OK.
- Close the PATH Command Status Window.
- Map results onto path (PDEF command) using the following menu path:

Main Menu > General Postproc > Path Operations > Map onto Path

- Map Result Items onto Path dialog box appears; select Stress from the left list and $Y$-direction $S Y$ from the right list; click on $O K$.
- Obtain line plot of $\sigma_{y y}$ along the path (PLPATH command) using the following menu path:

Main Menu > General Postproc $>$ Path Operations $>$ Plot Path Item $>$ On Graph

- Plot of Path Items on Graph dialog box appears; select SY; click on OK.
- Figure 8.17 shows the line plots of $\sigma_{x x}$ and $\sigma_{y y}$ along the defined path.

$-954.667{ }^{-801.426^{-648.185}}-494.944^{-341.703}-188.462^{-35.221} 118.02{ }^{271.261} \quad 424.502$
Fig. 8.15 Contour plot of $\sigma_{x x}$.


Fig. 8.16 Contour plot of $\sigma_{y y}$.


Fig. 8.17 Line plots of $\sigma_{x x}$ and $\sigma_{y y}$ along the defined path.

### 8.1.4.1.2 Composite Plate under Axial Tension

A fiber-reinforced square plate, shown in Fig. 8.18, is subjected to a uniform stress field of 20 ksi along the top and bottom boundaries. The sides of the plate are 10 in long, and the fibers are oriented at a $45^{\circ}$ angle to the global Cartesian coordinate system. Material properties are specified as $E_{1}=10 \times 10^{3} \mathrm{ksi}, E_{2}=30 \times 10^{3} \mathrm{ksi}, G_{12}=15 \times 10^{3} \mathrm{ksi}$, and $v_{12}=0.1$. The goal is to find the displaced shape.

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Solid in the left list and Quad 4 Node 42 in the right list; click on $\boldsymbol{O K}$.
- Click on Close.
- Specify material properties by typing the following four commands in the Input Field (at the end of each command, hit the Enter key to execute):

MP, EX, 1, 10E6
MP, EY, 1, 30E6
MP, PRXY, 1, 0.1
MP, GXY, 1, 15E6


Fig. 8.18 Schematic of the composite plate, fiber orientation, and loading.

- Create keypoints ( K command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create $>$ Keypoints $>\ln$ Active CS

- A total of 4 keypoints will be created.
- Enter $(x, y)$ coordinates of keypoint 1 as ( $-5,-5$ ); click on Apply. This action will keep the Create Keypoints in Active Coordinate System dialog box open. If the NPT Keypoint number field is left blank, then ANSYS assigns the lowest available keypoint number to the keypoint that is being created.
- Repeat the same procedure for the keypoints 2,3 , and 4 using (5, $\mathbf{- 5}$ ), $(5,5)$, and $(-5,5)$, respectively, for the $(x, y)$ coordinates.
- Once keypoint 4 is created, click on $O K$ (instead of Apply).
$\square$ Create the area through keypoints (A command) using the following menu path:
Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Arbitrary $>$ Through KPs
- Pick Menu appears; pick keypoints 1 through 4 (in sequence); click on OK.
- Material properties refer to the fiber directions. However, the global Cartesian coordinates and the fiber directions are at an angle of $45^{\circ}$. Therefore, the element coordinate system needs to be aligned with the fiber orientation. For this purpose, create a local coordinate system (CLOCAL command) using the following menu path:
Utility Menu > WorkPlane > Local Coordinate Systems > Create Local CS $>$ At Specified Loc
- Pick Menu appears; type 0,0,0 in the text field in the Pick Menu; click on $\mathbf{O K}$,
- A dialog box appears; type 45 in the THXY Rotation about local $Z$ text field; click on $\mathbf{O K}$; local coordinate system 11 is created.
- Align the element coordinate system with local coordinate system 11 (ESYS command) using the following menu path:


## Main Menu > Preprocessor > Meshing > Mesh Attributes > Default Attribs

- Meshing Attributes dialog box appears. Select 11 from the ESYS Element coordinate sys pull-down menu; click on OK.
- Switch the active coordinate system to global Cartesian using the following menu path:

Utility Menu > Change Active CS to > Global Cartesian

- Specify the number of divisions on all lines (LESIZE command) using the following menu path:

Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Lines > All Lines

- Element Sizes on All Selected Lines dialog box appears; enter 20 for NDIV; click on $O K$.
- Mesh the square (AMESH command) using the following menu path:

Main Menu > Preprocessor > Meshing > Mesh > Areas > Mapped > 3 or 4 sided

- Pick Menu appears; click on Pick All.


## SOLUTION

- Apply displacement constraints (D command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; pick the center node, i.e., $x=0$ and $y=0$; click on OK in the Pick Menu.
- Highlight both $\boldsymbol{U} \boldsymbol{X}$ and $\boldsymbol{U} \boldsymbol{Y}$; click on Apply.
- Pick Menu reappears; pick the right-side center node, i.e., $x=5$ and $y=$ 0 ; click on $O K$ in the Pick Menu.
- Remove the highlight on $\boldsymbol{U X}$, leaving $\boldsymbol{U Y}$ highlighted; click on $\boldsymbol{O K}$.
- Apply surface force (pressure) boundary conditions (SF command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Pressure > On Nodes

- Pick Menu appears; pick the nodes along the top and bottom surfaces of the plate; click on $O K$ in the Pick Menu.
- Type - 20000 (negative 20000) for VALUE Load PRES value; click on OK.
- Pressure, by definition, acts normal toward the body along the surface. The direction of action in reference to the global coordinate system does not affect whether it is positive or negative. The only factor that dictates the sign is whether it acts toward or away from the body. Therefore, in order to apply the tensile loading, it is necessary to apply negative pressure.
- Obtain the solution (SOLVE command) using the following menu path:


## Main Menu >Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status, if OK, close the Status Report Window; click on $O K$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the deformed shape (PLDISP command) using the following menu path:
Main Menu $>$ General PostProc $>$ Plot Results $>$ Deformed Shape
- Select Def + undef edge; click on OK.
- The deformed shape is shown in Fig. 8.19 as it appears in the Graphics Window.
- Review the $x$-displacement at the top-right and the $y$-displacement at the top-left nodes (PRNSOL command) using the following menu path:

Main Menu > General Postproc > List Results > Nodal Solution

- Highlight DOF solution in the left list and All DOFs DOF in the right list; click on OK.
- The list appears. The $x$-displacement at the top-right node (node 22 ) is given as $0.45 E-2$, and the $y$-displacement at the top-left node (node 42) is given as $0.45 E-2$.
- In ANSYS, results can also be listed (or displayed) in different coordinate systems. By default, the Results Coordinate System is aligned with the Global Cartesian. Align the Results Coordinate System with local coordinate system 11 defined earlier using the following menu path:

[^10]

Fig. 8.19 Deformed shape of the composite plate.

- Options for Output dialog box appears. Select Local system from the first pull-down menu and enter 11 for Local system reference no; click on $\boldsymbol{O K}$.
- Now, review the nodal displacements one more time using the following menu path:

Main Menu > General Postproc > List Results > Nodal Solution

- When the results are transformed to the local coordinate system, the $x$ displacement at the top-right node becomes $0.6364 E-2$, and the $y$ displacement at the top-left node becomes $0.16499 E-2$. Corresponding analytical solution values are $0.6364 E-2$ and $0.16495 E-2$, producing negligible error values.


### 8.1.4.2 Plane Strain

In a structural problem, if one of the dimensions is significantly longer than the other dimensions defining a uniform cross-sectional area, and if the structure is subjected to only uniform lateral loads, then plane strain idealization is valid. Similar to plane stress idealization, because the number of nodes and elements in the model is reduced drastically, utilization of plane strain idealization leads to significant savings in computational cost without loss of accuracy in the quantities of interest. Stresses in a bi-material cylindrical pressure vessel are used to demonstrate plane strain idealization.

A bi-material cylinder is subjected to internal pressure, $p_{a}$, as shown in Fig. 8.20. The radius of the hollow portion is $a$, and the thicknesses of the inner


Fig. 8.20 Plane strain representation of a bi-material cylindrical pressure vessel under internal pressure.
and outer cylinders are $(b-a)$ and $(c-a)$, respectively. Perfect contact with no slipping is assumed along the interface, implying displacement continuity. Elastic properties of the inner and outer cylinders are ( $E_{1}, v_{1}$ ) and ( $E_{2}, v_{2}$ ), respectively. The goal is to compute the stress field. The problem is solved with ANSYS using $E_{2} / E_{1}=0.5, v_{a}=v_{b}=0.33, b / a=2$, and $c / a=4$, with $a=1, p_{a}=1$, and $E_{1}=2$.

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Solid in the left list and Quad 4 Node 42 in the right list; click on $O K$.
- Click on Options.
- PLANE42 element type options dialog box appears; select Plane strain item from the pull-down menu corresponding to Element behavior K3.
- Click on $O K$; click on Close.
- Specify material properties (MP command) using the following menu path:


## Main Menu $>$ Preprocessor $>$ Material Props $>$ Material Models

- The inner and outer cylinder will have material reference number 1 and 2, respectively. Define Material Model Behavior dialog box appears. In
the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which brings up another dialog box.
- Enter 2 for $\boldsymbol{E X}$ and 0.33 for $\boldsymbol{P R X Y}$; click on $\mathbf{O K}$.
- Add new material model using the following menu path:

Material > New Model

- Click on $O K$ in the new dialog box.
- In the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic; Enter 1 for $E X$ and 0.33 for PRXY; click on $\boldsymbol{O K}$.
- When finished, close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create partial hollow circular areas (PCIRC command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Circle $>$ By Dimensions

- In the Create Circle by Dimensions dialog box, type 2 for Outer radius, 1 for Inner radius, 0 for Theta1, and 90 for Theta2; click on Apply.
- Now, type 4 for Outer radius, 2 for Inner radius, 0 for Theta1, and 90 for Theta2; click on $O K$.
- Glue the areas (AGLUE command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Operate $>$ Booleans $>$ Glue $>$ Areas

- Pick Menu appears; click on Pick All button.
- Create the mesh. Since the problem involves two dissimilar materials, the inner circle (material 1) will be meshed first. Then the default material attribute will be changed to material 2 for the outer circle. Specify global element size (ESIZE command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Size Cntrls $>$ ManualSize $>$ Global $>$ Size

- Global Element Sizes dialog box appears; enter 0.1 for SIZE; click on OK.
- Mesh the inner circle (AMESH command) using the following menu path:

$$
\text { Main Menu > Preprocessor }>\text { Meshing }>\text { Mesh }>\text { Areas }>\text { Mapped }>3 \text { or } 4 \text { sided }
$$

- Pick Menu appears; pick the inner circle; click on $O K$.
- Change the default material attribute to 2 (MAT command) using the following menu path:


## Main Menu > Preprocessor > Meshing > Mesh Attributes > Default Attribs

- Meshing Attributes dialog box appears. Select 2 from the second pulldown menu; click on $O K$.
- Mesh the outer circle (AMESH command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 sided

- Pick Мепи appears; pick the outer circle; click on $O K$.


## SOLUTION

- Apply displacement constraints (D command) using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Structural $>$ Displacement $>$ On Nodes

- Pick Menu appears; pick the nodes along $x=0$ (coincident with $y$-axis); click on $O K$ in the Pick Menu.
- Highlight $\boldsymbol{U X}$; click on Apply.
- Pick Menu reappears; pick the nodes along $y=0$ (coincident with $x$ axis); click on $O K$ in the Pick Menu.
- Highlight $\boldsymbol{U} \boldsymbol{Y}$ and remove the highlight on $\boldsymbol{U} X$; click on $O K$.

■ Apply surface force (pressure) boundary conditions along the inner circular boundary. Since the boundary is circular, it is convenient to first switch to Cylindrical Coordinates and then select the nodes.

- Switch to Cylindrical Coordinates (CSYS command) using the following menu path:

Utility Menu > WorkPlane > Change Active CS to > Global Cylindrical

- Select nodes along the circular boundary (NSEL command) by using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; choose By Location in the second pull-down menu and type 1 in the Min, Max text field; click on $O K$. Because the active coordinate system is cylindrical, any reference to the $x$-coordinate is treated as a reference to the $r$-coordinate by ANSYS.
- Now, apply pressure boundary conditions (SF command) by using the following menu path:

[^11]- Pick Menu appears; click on Pick All.
- Type $\mathbf{I}$ for VALUE Load PRES value; click on OK.
- Select everything (ALLSEL command) using the following menu path: Utility Menu > Select > Everything
- Switch back to Cartesian Coordinates (CSys command) using the following menu path:

```
Utility Menu > WorkPlane > Change Active CS to > Global Cartesian
```

- Obtain the solution (SOLVE command) using the following menu path:

Main Menu >Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Obtain contour plots for $\sigma_{x x}$ and $\sigma_{y y}$ (PLNSOL command) using the following menu path:

Main Menu > General Postproc > Plot Results > Contour Plot $>$ Nodal Solu

- Highlight Stress in the left list (second item) and highlight $X$-direction $S X$ in the right list; click on $O K$.
- The contour plot appears in the Graphics Window, as shown in Fig. 8.21 (left).
- Repeat the same procedure for $\sigma_{y y}$, which produces the contour plot given in Fig. 8.21 (right)
- Since the problem possesses a circular geometry, it is often more useful to examine the stresses in cylindrical coordinates. For this purpose, change the results coordinate system to the global cylindrical system (RSYS command) using the following menu path:

Main Menu > General Postproc > Options for Outp

- Options for Output dialog box appears. Select Global cylindric from the first pull-down; click on $O K$.
- Now, obtain contour plots for $\sigma_{r r}$ and $\sigma_{\theta \theta}$ (PLNSOL command) using the following menu path:

[^12]

Fig. 8.21 Normal stresses in Cartesian coordinates: in $x$-direction, $\sigma_{x x}$ (left), and in $y$-direction, $\sigma_{y y}$ (right).

- Highlight Stress in the left list (second item) and highlight $\boldsymbol{X}$-direction $\boldsymbol{S X}$ in the right list; click on $\boldsymbol{O K}$.
- The contour plot appears in the Graphics Window, as shown in Fig. 8.22 (left).
- Repeat the same procedure for $\sigma_{\theta \theta}$, which produces the contour plot given in Fig. 8.22 (right).
- Review the variation of stresses along a path by means of a line plot.

Define the path (PPATH command) using the following menu path:
Main Menu > General Postproc > Path Operations > Define Path $>$ By Nodes

- Pick Menu appears; pick the nodes with $(x, y)$ coordinates $(1,0)$ and $(4,0)$; click on $O K$. The path lies along the positive $x$-axis, starting from the boundary of the hole and ending at the left boundary of the structure.
- By Nodes dialog box appears; enter a name describing the path, say hrz, in the Define Path Name text field; click on OK.
- Close the PATH Command Status Window.
- Map results onto path (PDEF command) using the following menu path:
Main Menu > General Postproc > Path Operations > Map onto Path
- Map Result Items onto Path dialog box appears; select Stress from the left list and $\boldsymbol{Y}$-direction $S \boldsymbol{Y}$ from the right list; click on $\boldsymbol{O K}$.
- Obtain line plot of $\sigma_{\theta \theta}$ along the path (PLPATH command) using the following menu path:

Main Menu > General Postproc > Path Operations > Plot Path Item > On Graph

- Plot of Path Items on Graph dialog box appears; select SY; click on OK.
- Figure 8.23 shows the line plots of $\sigma_{r r}$ and $\sigma_{\theta \theta}$ along the defined path.


Fig. 8.22 Normal stresses in cylindrical coordinates: in $r$-direction, $\sigma_{r r}$ (left), and in $\theta$-direction, $\sigma_{\theta \theta}$ (right).


Fig. 8.23 Radial and hoop stresses along $y=0$.

### 8.1.4.3 Axisymmetric

In a solid of revolution, location of a point in the body can conveniently be identified by cylindrical coordinates, $r, \theta$ and $z$, with $z$ being the axis of rotation. When a solid of revolution is subjected to loading that can also be obtained by revolution about the $z$-axis, then the results become independent of $\theta$. This is called an Axisymmetric Condition. Two problems are considered here.

### 8.1.4.3.1 Deformation of a Bar Due to its Own Weight Using 2-D Axisymmetric Elements

Deformation of a bar under its own weight was modeled in Sec. 8.1.1.1 using two-dimensional link elements. The problem was solved using threedimensional brick elements in Sec. 8.1.3. The solution to this problem also can be obtained using axisymmetric elements as the geometry and the loading (gravity) exhibit conditions for axisymmetry. The following axisymmetric solution utilizes the reference coordinate frame shown in Fig. 8.1.

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Solid in the left list and Quad 4 Node 42 in the right list; click on $\boldsymbol{O K}$.
- Click on Options.
- PLANE42 element type options dialog box appears; select the Axisymmetric item from the pull-down menu corresponding to Element behavior K3.
- Click on $O K$; click on Close.
- Specify material properties for the bar (MP command) using the following menu path:


## Main Menu > Preprocessor $>$ Material Props $>$ Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural and Density, which will bring up another dialog box.
- Enter 0.2839605 for DENS; click on OK.
- In order to specify the elastic modulus and Poisson's ratio, in the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring up another dialog box.
- Enter 30e6 for $\boldsymbol{E X}$ and $\mathbf{0 . 3}$ for $\boldsymbol{P R X Y}$; click on $O K$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create the rectangle defining the axisymmetric cross section (RECTNG command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Rectangle $>$ By Dimensions

- In the Create Rectangle by Dimensions dialog box, enter 0 and 2 for X1 and $X 2$ and 0 and -20 for $Y 1$ and $Y 2$; click on $O K$.
- Specify the global element size (ESIZE command) using the following menu path:

Main Menu > Preprocessor > Meshing > Size Cntrls $>$ ManualSize $>$ Global > Size

- Global Element Sizes dialog box appears; enter 0.2 for SIZE; click on OK.
- Create the mesh (AMESH command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Free

- Pick Menu appears; click on Pick All.


## SOLUTION

■ Apply displacement constraints (D command) using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Structural $>$ Displacement $>$ On Nodes

- Pick Menu appears; pick the nodes at $y=0$; click on $O K$ in the Pick Menu.
- Highlight All DOF; click on $O K$.
- Apply gravitational acceleration (ACEL command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Inertia > Gravity

- Apply (Gravitational) Acceleration dialog box appears.
- Enter 386.2205 for ACELY; click on OK.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu >Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

■ Review $y$-displacement contours (PLNSOL command) using the following menu path:

## Main Menu > General PostProc > Plot Results > Contour Plot > Nodal Solu

- Contour Nodal Solution Data dialog box appears. Select DOF solution from the left list and Translation $U Y$ from the right list; click on $O K$.
- The contour plot is shown in Fig. 8.24 as it appears in the Graphics Window.
- Review displacement values (PRNSOL command) using the following menu path:


## Main Menu > General Postproc > List Results > Nodal Solution

- Highlight DOF solution in the left list and Translation $U Y$ in the right list; click on $\boldsymbol{O K}$.
- The list appears in a separate window. It is a long list of $z$ displacements. At the bottom of the window, the maximum displacement value is printed as $-0.72464 E-03$.


### 8.1.4.3.2 Analysis of a Circular Plate Pushed Down by a Piston Head

An aluminum circular plate with a diameter of 40 in is pushed down by a steel piston head, as shown in Fig. 8.25. The piston head has two sections with diameters 20 in and 2 in . The elastic modulus and Poisson's ratio for the aluminum plate are given as $E_{a l}=10 \times 10^{6} \mathrm{psi}$ and $v_{a l}=0.35$, respectively, whereas the corresponding properties for steel are $E_{s t}=30 \times 10^{6} \mathrm{psi}$ and $v_{s t}=0.3$. The aluminum plate is clamped along the boundary (all degrees of freedom constrained). The goal is to obtain the displacement and stress fields when the piston is pushed down (at the top) by an amount of 0.1 in. This problem possesses the conditions necessary for axisymmetry to be employed. Following is the solution utilizing axisymmetric elements in ANSYS.

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:


## Main Menu > Preprocessor > Element Type > Add/EditDelete

- Click on $\operatorname{Add}$.
- Select Solid in the left list and Quad 4 Node 42 in the right list; click on $\boldsymbol{O K}$.
- Click on Options.


Fig. 8.24 Contour plot of z-displacement of a bar elongated due to its own weight.


Fig. 8.25 Schematic of a circular plate pushed down by a piston head.

- PLANE42 element type options dialog box appears; select Axisymmetric item from the pull-down menu corresponding to Element behavior K3.
- Click on $\mathbf{O K}$; click on Close .

■ Specify material properties (MP command) using the following menu path:

## Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring up another dialog box.
- Enter $\mathbf{1 0 e 6}$ for $\boldsymbol{E X}$ and $\mathbf{0 . 3 5}$ for $\boldsymbol{P R X Y}$; click on $\boldsymbol{O K}$.
- Add new material model using the following menu path:


## Material > New Model

- Click on $O K$ in the new dialog box.
- In the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic; Enter $30 e 6$ for $E X$ and 0.3 for PRXY; click on $O K$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Three rectangles defining the geometry will be created and overlapped. Create the rectangles defining the axisymmetric cross section (RECTNG command) using the following menu path:


## Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Rectangle $>$ By Dimensions

- In the Create Rectangle by Dimensions dialog box, enter 0 and 20 for $X 1$ and $X 2$ and 0 and 0.5 for Y1 and Y2; click on Apply.
- Now, enter 0 and 10 for $X 1$ and $X 2$ and 0 and 1.5 for $Y 1$ and $Y 2$; click on Apply.
- Finally, enter 0 and 1 for $X 1$ and $X 2$ and 0 and 5.5 for $Y 1$ and $Y 2$; click on $O K$.
- Overlap the rectangles (AOVLAP command) using the following menu path:

Main Menu > Preprocessor > Modeling > Operate > Booleans > Overlap > Areas

- Pick Menu appears, click on Pick All.
- The overlapping operation produces six areas (started with three), sharing lines along the interfaces.
- Specify the global element size (ESIZE command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Size Cntrls $>$ ManualSize $>$ Global $>$ Size

- Global Element Sizes dialog box appears; enter 0.2 for SIZE; click on OK.
- Create the mesh for the aluminum plate (AMESH command) using the following menu path:


## Main Menu >Preprocessor > Meshing > Mesh > Areas > Mapped > 3 or 4 sided

- Pick Menu appears; pick the bottom row of rectangles (corresponding to the aluminum plate); click on $\boldsymbol{O K}$ in the Pick Menu.
- Plot the areas (APLOT command) using the following menu path:

Utility Menu > Plot > Areas

- Change default element attribute for material number from 1 to 2 (MAT command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh Attributes $>$ Default Attribs

- Meshing Attributes dialog box appears; select 2 from the [MAT] Material number pull-down menu; click on OK.
- Create mesh for the steel piston (AMESH command) using the following menu path:

Main Menu > Preprocessor > Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 sided

- Pick Menu appears; pick the rectangles corresponding to the steel piston; click on $O K$ in the Pick Menu.
- Plot elements with different colors based on their material numbers using the following menu path:

Utility Menu > PlotCtris > Numbering

- Plot Numbering Controls dialog box appears. Select Material numbers from the first pull-down menu (corresponding to Elem / Attrib numbering) and select Colors only from the second pull-down menu (corresponding to [/NUM] Numbering shown with); click on OK. Figure 8.26 shows the corresponding element plot with different colors based on material numbers.


## SOLUTION

- Apply displacement constraints along the periphery of the aluminum plate (D command) using the following menu path:

[^13]

Fig. 8.26 Element plot with different colors based on material numbers.

- Pick Menu appears; pick the nodes along the right boundary $(x=20)$; click on $\boldsymbol{O K}$ in the Pick Menu.
- Highlight All DOF ; click on $\boldsymbol{O K}$.
- Apply displacement constraints along the top surface of the steel piston (D command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; pick the nodes along the top boundary $(y=5.5)$; click on $\boldsymbol{O K}$ in the Pick Menu.
- Remove the highlight on All DOF and highlight $\boldsymbol{U Y}$.
- Enter $\mathbf{- 0 . 1}$ in the text box for VALUE Displacement value; click on OK.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status, if OK, close the Status Report Window; click on $\boldsymbol{O K}$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

■ Review the deformed shape (PLDISP command) using the following menu path:

Main Menu $>$ General PostProc $>$ Plot Results $>$ Deformed Shape

- Plot Deformed Shape dialog box appears; select the Def + undef edge radio button; click on $\boldsymbol{O K}$.
- Corresponding deformed shape is shown in Fig. 8.27.


Fig. 8.27 Deformed shape with undeformed edge.

- Review the equivalent stress (von Mises) contour plot (PLNSOL command) using the following menu path:

Main Menu $>$ General PostProc $>$ Plot Results $>$ Contour Plot $>$ Nodal Solu

- Contour Nodal Solution Data dialog box appears. Select Stress from the left list; in the right list, scroll down to select von Mises SEQV. Click on $\boldsymbol{O K}$.
- Figure 8.28 shows the corresponding contour plot


### 8.1.5 Plates and Shells

Many engineering structures involve plates and shells where one dimension is much smaller than the other two. When these thin members are flat and only in-plane loads are applied, the problem can be solved using Plane Stress idealization. However, if they are curved and/or subjected to both inplane and out-of-plane loads, it is necessary to solve the problem in 3-D using shell elements. At each node of the shell elements, both displacements and rotations are the degrees of freedom. Three problems are solved utilizing shell elements.

### 8.1.5.1 Static Analysis of a Bracket

The bracket shown in Fig. 8.29 is clamped at the two top holes and is subjected to static vertical loading at the bottom two holes. Due to the symmetry in geometry, only one quarter of the structure is modeled at first. Once the top-left quarter is modeled and meshed, two symmetric reflection operations are utilized to create the rest of the bracket. The goal is to create the finite element model and obtain the static solution.

## MODEL GENERATION

- Specify the jobname as bracket using the following menu path:


## Utility Menu > File > Change Jobname

- In the dialog box, type bracket in the [/FILNAM] Enter new jobname text field; click on the checkbox for New $\log$ and error files to show Yes; click on $\boldsymbol{O K}$.


Fig. 8.28 Equivalent stress contours.


Fig. 8.29 Geometry, material properties, and loading on the bracket.

- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on $\operatorname{Add}$.
- Select Shell in the left list and Elastic 4node 63 in the right list; click on $\boldsymbol{O K}$.
- Click on Close.
- Specify the thickness using real constants ( R command) using the following menu path:


## Main Menu > Preprocessor > Real Constants > Add/Edit/Delete

- Click on $\operatorname{Add}$.
- Click on $\boldsymbol{O K}$; dialog box appears.
- Type 0.1 in the Shell thickness at node I TK(I) text field; click on $\boldsymbol{O K}$.
- Click on Close.

Specify material properties (MP command) using the following menu path:

## Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring up another dialog box.
- Enter $\mathbf{3 0 e 6}$ for $\boldsymbol{E X}$ and $\mathbf{0 . 3}$ for PRXY; click on $\boldsymbol{O K}$.
- In the Define Material Model Behavior dialog box, in the right window, double-click on Density, which will bring up another dialog box.
- Enter $\mathbf{0 . 0 0 0 7 3}$ for DENS; click on OK.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create the solid model.
- Move Working Plane origin using the following menu path:

Utility Menu > WorkPlane > Offset WP by Increments

- Offset WP dialog box appears; type $0,3,-2$ in the $X, Y, Z$ Offsets text field; click on $\boldsymbol{O K}$.
- Create a rectangular area using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Rectangle $>$ By Dimensions

- In the Create Rectangle by Dimensions dialog box, type -2 for X1, 0 for $\boldsymbol{X 2}, \mathbf{0}$ for $\boldsymbol{Y} \mathbf{1}$, and $\mathbf{2}$ for $\boldsymbol{Y} \mathbf{2}$; click on $O K$.
- Create a circular area using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Areas > Circle > By Dimensions

- In the Create Circle by Dimensions dialog box, type 1 for Outer radius, 90 for Theta1, and 180 for Theta2; click on OK.
- Subtract the circle from the rectangle using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Operate $>$ Booleans $>$ Subtract $>$ Areas

- Pick Мепи appears; pick the rectangle; click on $\boldsymbol{O K}$; pick the circle; click on $O K$.
- Move the Working Plane origin to the top-left hole center using the following menu path:

Utility Menu > WorkPlane > Offiset WP by Increments

- Offset WP dialog box appears; type $-1.5,1.5$ in the $X, Y, Z$ Offsets text field (because only $x$ - and $y$-increments are entered, no move will be applied in $z$-direction); click on $O K$.
- Create a circular area for the top-left hole using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Circle $>$ By Dimensions

- In the Create Circle by Dimensions dialog box, type $0.25 / 2$ for Outer radius, 0 for Theta1, and 360 for Theta2; click on OK.
- Subtract the circle from the rest of the area using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Operate $>$ Booleans $>$ Subtract $>$ Areas

- Pick Menu appears; pick the large area and click $O K$; pick the circle; click on $O K$.
- Move the Working Plane in order to create the additional rectangular area using the following menu path:

Utility Menu > WorkPlane > Offset WP by Increments

- Offset $W P$ dialog box appears; type $-0.5,0.5$ in the $\boldsymbol{X}, \boldsymbol{Y}, \mathbf{Z}$ Offsets text field; click on $\boldsymbol{O K}$.
- Create additional rectangular area using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Reclangle $>$ By Dimensions

- In the Create Rectangle by Dimensions dialog box, type 0 for $\mathbf{X 1}, 1$ for $X 2,-2$ for $Y 1$, and -5 for $Y 2$; click on $O K$.
- In order to create a curved area, create keypoints that define the axis of rotation using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Keypoints $>$ In Active CS

- Create Keypoints in Active Coordinate System dialog box appears; type 51 for NPT Keypoint number and 0 in the $X, Y, Z$ Location in active CS text fields; click on Apply.
- In the same dialog box, type 52 for NPT Keypoint number and - 0.5 for $x$ and $\boldsymbol{0}$ for $y$ and $z$ in the $\boldsymbol{X}, \boldsymbol{Y}, \boldsymbol{Z}$ Location in active $\boldsymbol{C S}$ text fields; click on $\boldsymbol{O K}$.
- Plot areas using the following menu path:

Utility Menu > Plot > Areas

- Create the curved area by sweeping the line at the bottom around an axis defined by the last two keypoints created using the following menu path:
- Pick Menu appears; the user is first asked to pick the line to be swept, and then to pick the keypoints defining the axis that the line to be swept about.
- Pick the horizontal line at the bottom; click on $\mathbf{O K}$; type 51 in the text field in the Pick Menu and hit Enter on the keyboard; type 52 followed by hitting Enter on the keyboard; click on $\boldsymbol{O K}$.
- Sweep Lines about Axis dialog box appears; type 45 for ARC Arc length in degrees; click on $O K$.
- Click on the Isometric View button.
- Figure 8.30 shows the result of this action.
- Although the areas created appear to be connected, ANSYS treats them as independent of each other (not connected). Therefore, the areas must be glued to each other. This is achieved by using the following menu path:

Main Menu > Preprocessor > Modeling > Operate > Booleans > Glue > Areas

- Pick Menu appears; click on Pick All.
- Create the mesh.
- Specify the number of elements around the hole using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Size Cntrls $>$ ManualSize $>$ Lines $>$ Picked Lines


Fig. 8.30 Solid model of a quarter of the bracket.

- Pick the four circular segments defining the hole; click on $\boldsymbol{O K}$.
- Element Sizes on Lines dialog box appears; type 2 in the text field corresponding to NDIV (the second text field); uncheck the first checkbox; click on $\boldsymbol{O K}$.
- Specify mesh density in the vicinity of the top-left corner using the following menu path:
Main Menu > Preprocessor > Meshing $>$ Size Cntrls > ManualSize > Keypoints $>$ Picked KPs
- Pick Menu appears; pick the top-left keypoint; click on OK.
- Element Sizes at Picked Keypoints dialog box appears; type 0.3 for SIZE Element edge length text field; click on OK.
- Specify global mesh density using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Size Cntrls $>$ ManualSize $>$ Global $>$ Size

- Global Element Sizes dialog box appears; type 0.5 for SIZE Element edge length text field; click on OK.
- Create the mesh using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Free

- In the Pick Menu, click on Pick All.
- Save the model using the following menu path:

Utility Menu > File > Save as Jobname.db
The model will be saved in the Working Directory under the name bracket.db.

- Create a reflective symmetric mesh using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Reflect $>$ Areas

- Pick Menu appears; click on Pick All.
- Reflect Areas dialog box appears; click on the $Y$ - $Z$ plane $X$ radiobutton; click on $\mathbf{O K}$.
- Although it is not apparent through visual inspection, there are duplicate entities (keypoints, lines and nodes) along the symmetry line, thus there is no continuity. Therefore, merge duplicate entities using the following menu path:

Main Menu $>$ Preprocessor $>$ Numbering Ctrls $>$ Merge Items

- In the dialog box, select All from the first pull-down menu; click on OK.
- Create a second reflective symmetric mesh.
- For this purpose, create a local coordinate system using the following menu path:

[^14]- Pick Menu appears; type 0, 0, 0 in the text field in the Pick Menu; click on $\boldsymbol{O K}$.
- A dialog box appears; type $\mathbf{- 4 5}$ in the THYZ Rotation about local X text field; click on $\mathbf{O K}$.
- Create a reflective symmetric mesh using the following menu path:

Main Menu > Preprocessor > Modeling > Reflect > Areas

- Pick Menu appears; click on Pick All.
- Reflect Areas dialog box appears; click on the $X-Z$ plane $Y$ radiobutton; click on $\boldsymbol{O K}$.
- Plot elements using the following menu path:

Utility Menu > Plot > Elements

- Figure 8.31 shows the isometric view of the mesh after the reflection.
- Merge duplicate entities using the following menu path:

Main Menu > Preprocessor > Numbering Ctris > Merge Items

- In the dialog box, select $A l l$ from the first pull-down menu; click on OK.
- Define components for future use.
- For this purpose, create a local coordinate system at the center of the top-left hole using the following menu path:

Utility Menu > WorkPlane > Local Coordinate Systems > Create Local CS > At Specified Loc

- Pick Menu appears; type -1.5, 4.5, -2 in the text field in the Pick Menu; click on $\boldsymbol{O K}$.
- A dialog box appears; select Cylindrical 1 in the KCS Type of coordinate system pull-down menu.
- Delete -45 in the THYZ Rotation about local $X$ text field; click on OK.
- Select nodes along the top-left hole by using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; choose By Location in the second pull-down menu and type $0.25 / 2$ in the Min, Max text field; click on OK. Because the active coordinate system is cylindrical, any reference to the $x$-coordinate will be treated as a reference to the $r$-coordinate by ANSYS.
- Create the component by using the following menu path:


Fig. 8.31 Bracket after meshing and two reflection operations.

- Component Manager dialog box appears (Fig. 8.32); click on the first button on the left (Create Component button).
- Create Component dialog box appears; click on the Nodes radio-button and name the component by typing $\boldsymbol{T L} \boldsymbol{B O L T}$ (stands for top-left bolt) in the text field (Fig. 8.33); click on $\boldsymbol{O K}$.
- Close the Component Manager.
- Create components for top-right, bottom-left, and bottom-right bolts in the same manner. The origin of the local cylindrical coordinates for each of these are given as

TR_BOLT: 1.5, 4.5, -2
BL_BOLT: $-1.5,-2,4.5$ and use -90 for the THYZ Rotation about local $\boldsymbol{X}$
BR_BOLT: $1.5,-2,4.5$ and use $\mathbf{- 9 0}$ for the THYZ Rotation about local $X$

- Save the model using the following menu path:

Utility Menu > File > Save as Jobname.db


Fig. 8.32 Component Manager dialog box (left-most button is used for creating components).


Fig. 8.33 Dialog box for creating components.

## SOLUTION

- Constrain displacement and rotation degrees of freedom along the top-left and -right holes. For this purpose, first select the components created earlier for these holes ( $\boldsymbol{T L} \_\boldsymbol{B O L T}$ and $\boldsymbol{T R} \_\boldsymbol{B O L T}$ ) using the following menu path:

Utility Menu > Select > Comp/Assembly > Select Comp/Assembly

- A dialog box appears; click on the by component name radio-button; click on $\boldsymbol{O K}$.
- A new dialog box with the components listed appears; highlight $T L \_B O L T$; click on $O K$. This action selects the nodes along the top-left hole.
- Specify the displacement boundary conditions using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; click on Pick All.
- In the new dialog box, highlight All DOF; click on $O K$.
- Repeat the same procedure for the top-right hole $\left(\boldsymbol{T R} \_B O L T\right)$.
- Apply force boundary conditions.
- For this purpose, create a local coordinate system at the center of the bottom-left hole using the following menu path:

Utility Menu > WorkPlane > Local Coordinate Systems >Create Local CS $>$ At Specified Loc

- Pick Menu appears; type -1.5, -2, 4.5 in the text field in the Pick Menu; click on $\boldsymbol{O K}$.
- A dialog box appears; select Cylindrical 1 in the KCS Type of coordinate system pull-down menu and type -90 in the THYZ Rotation about local $X$ text field; click on $O K$.
- Select the keypoints along the bottom-left hole using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; choose Keypoints in first pull-down menu and By Location in the second pull-down menu; type $0.25 / 2$ in the Min, Max text field; click on $O K$.
- Apply forces on the keypoints using the menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Force/Moment > On Keypoints

- Pick Menu appears; click on Pick All.
- In the new dialog box, select $\boldsymbol{F Y}$ from pull-down menu and enter $6 / 4$ for Force/moment value; click on OK.
- A Warning Window appears, informing the user that boundary conditions applied to solid modeling entities overwrite those that may have already been applied to finite element entities (nodes and elements) directly.
- Close the Warning Window.
- Repeat the same procedure for the bottom-right hole (use 1.5, -2, 4.5 for the local coordinate system origin and -90 for the THYZ Rotation about local X).
- Select everything using the following menu path:

Utility Menu > Select > Everything

- Save the model using the following menu path:

Utility Menu > File > Save as Jobname.db

- Obtain the solution using the following menu path:

Main Menu > Solution > Solve >Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the deformed shape using the following menu path:

Main Menu > General PostProc > Plot Results > Deformed Shape

- Select Def + undef edge; click on OK.
- The deformed shape is shown in Fig. 8.34 (left) as it appears in the Graphics Window.
- Review the stress contours using the following menu path:

Main Menu > General PostProc > Plot Results > Contour Plot > Nodal Solu

- Select stress in the left list and vonMises $\operatorname{SEQV}$ in the right list; click on $\boldsymbol{O K}$.
The equivalent stress contour plot is shown in Fig. 8.34 (right) as it appears in the Graphics Window.


### 8.1.5.2 Analysis of a Circular Plate Pushed Down by a Piston Head

The circular aluminum plate pushed down by a steel piston was analyzed in Sec. 8.1.4.3.2 by employing only axisymmetric elements. The geometry of the problem is shown in Fig. 8.25. The same problem is solved in this section by using a combination of shell and 3-D solid elements.

## MODEL GENERATION

■ Define the element types (ET command) using the following menu path:
Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Element Types dialog box appears; click on Add.


Fig. 8.34 Deformed shape (left), and contour plot of equivalent (von Mises) stresses (right).

- Select Shell in the left list and scroll down to select Linear Layer 99 in the right list; click on Apply.
- Select Solid in the left list and scroll down to select 20node 95 in the right list; click on $O K$.
- Define keyoptions on the shell element by highlighting Type 1 SHELL99 in the Element Types dialog box and clicking on Options.
- SHELL99 element type options dialog box appears; select Stress \& strain from the fourth pull-down menu (corresponding to Strains or stresses output K5) and select All layers from the Storage of layer data K8 pull-down menu.
- Finally, select Nodes @ top face from the last pull-down menu (corresponding to Node offset option K11); click on OK.
- Click on Close to exit from the Element Types dialog box.
- Specify the thickness using real constants ( R command) using the following menu path:
Main Menu > Preprocessor > Real Constants > Add/Edit/Delete
- Real Constants dialog box appears; click on Add.
- Element Type for Real Constants dialog box appears; highlight Type 1 SHELL99; click on OK.
- A new dialog box appears prompting the user to enter the Real Constant Set No. (with 1 default); click on $O K$.
- Real Constant Set Number 1, for SHELL99 dialog box appears; click on OK.
- Another dialog box appears, prompting the user to enter material, orientation (if orthotropic), and thickness information. Enter 0.5 in the third text box (corresponding to $\boldsymbol{T K}$ ); click on $\boldsymbol{O K}$.
- Exit from the Real Constants dialog box by clicking on Close.
- Specify material properties (MP command) using the following menu path:

Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which brings up another dialog box.
- Enter $\mathbf{1 0 e 6}$ for $\boldsymbol{E X}$ and $\mathbf{0 . 3 5}$ for PRXY; click on $\boldsymbol{O K}$.
- Add new material model using the following menu path:

Material > New Model

- Click on OK.
- In the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic; Enter 30e6 for $E X$ and 0.3 for PRXY; click on $O K$.
- Close the Define Material Model Behavior dialog box by using the following menu path:


## Material > Exit

- Create quarter circular areas (PCIRC command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Areas > Circle > By Dimensions

- In the Create Circle by Dimensions dialog box, type 1 for Outer radius, 0 for Theta1, and 90 for Theta2; click on Apply.
- Modify Outer radius to be 10; click on Apply.
- Finally, modify Outer radius (one more time) to be 20 ; click on $O K$.
- Overlap the areas (AOVLAP command) using the following menu path:

```
Main Menu > Preprocessor > Modeling > Operate > Booleans > Overlap > Areas
```

- Pick Menu appears; click on Pick All.
- Create quarter cylindrical volumes (CYLIND command) using the following menu path:


## Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Volumes $>$ Cylinder $>$ By Dimensions

- In the Create Cylinder by Dimensions dialog box, type 10 for Outer radius, 0 for $\mathbf{Z 1}, 1$ for Z2, 0 for Theta1, and 90 for Theta2; click on Apply.
- Modify Outer radius to be 1 and $Z 2$ to be 5 ; click on $O K$.

■ Overlap the volumes (VOVLAP command) using the following menu path:

Main Menu > Preprocessor > Modeling > Operate > Booleans > Overlap > Volumes

- Pick Menu appears; click on Pick All.

Specify size controls for meshing. First, the number of divisions on specific lines will be specified, followed by specification of the global element size.

- Select lines at $0.25 \leq x \leq 0.75$ and $0.25 \leq y \leq 0.75$ (LSEL command) using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; choose Lines from the first pulldown menu and By Location in the second pull-down menu. Type $0.25,0.75$ in the Min, Max text field; click on Apply.
- Now, select Y-coordinates and the Also Select radio-buttons without changing the text $(0.25,0.75)$ in the Min, Max text field; click on $O K$.
- A total of 12 lines are selected. Specify the number of element divisions along the selected lines (LESIZE command) using the following menu path:

Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Lines > Picked Lines

- Pick Menu appears; click on Pick All.
- Element Sizes on Picked Lines dialog box appears; enter 4 for NDIV. Remove the checkmark next to KYNDIV SIZE,NDIV can be changed so that it shows No; click on $O K$.
- Specify global element size (ESIZE command) using the following menu path:

Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Global > Size

- Global Element Sizes dialog box appears; enter 0.5 for SIZE; click on OK.
- Select everything (ALLSEL command) using the following menu path:

Utility Menu > Select > Everything

- Generate the mesh in two stages: (i) the mesh associated with the aluminum plate is created using SHELL99 elements followed by (ii) the generation of the mesh for the steel piston head using SOLID95 elements.
- Select areas attached to the volumes (ASLV command) using the following menu path:
- Select Entities dialog box appears; choose Areas from the first pulldown menu and Attached to in the second pull-down menu. Select Volumes and From Full radio-buttons; click on Apply.
- Now, click on Invert button. This inverts the selection, i.e., selected areas are unselected and vice versa. At this point, areas associated with the aluminum plate are selected.
- Create the mesh for the aluminum plate (AMESH command) using the following menu path:

Main Menu > Preprocessor > Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 sided

- Pick Menu appears; click on Pick All.
- Select everything (ALLSEL command) using the following menu path:

Utility Menu > Select > Everything

- Change the default element type attribute to 2 (TYPE command) and default material attribute to 2 (MAT command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh Attributes $>$ Default Attribs

- Meshing Attributes dialog box appears. Select 2 SOLID95 from the first pull-down menu and select 2 from the second pull-down menu; click on $O K$.
- Create the mesh for the steel piston head (VMESH command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Volumes $>$ Mapped $>4$ to 6 sided

- Pick Menu appears; click on Pick All.
- Obtain an isometric view of the mesh using the following menu path:

Utility Menu > PlotCtrls > Pan Zoom Rotate

- Pan Zoom Rotate window appears; click on Iso button.
- Plot elements with different colors based on their material numbers using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Plot Numbering Controls dialog box appears. Select Material numbers from the first pull-down menu (corresponding to Elem/Attrib numbering) and select Colors only from the second pull-down menu (corresponding to [/NUM] Numbering shown with); click on OK. Figure 8.35 (left) shows the corresponding element plot with different colors based on material numbers.


Fig. 8.35 Elements plotted without (left) and with (right) thickness information from real constants and curved surfaces.

- Note in the element plot that the aluminum plate elements do not have a thickness. This is because the thickness information is stored in real constants and the SHELL99 elements are plane elements. However, for visualization purposes, it is possible to plot plane elements with their thickness (/ESHAPE command) using the following menu path:

Utility Menu > PlotCtrls > Style > Size and Shape

- Size and Shape dialog box appears. Place a checkmark next to [/ESHAPE] Display of element shapes based on real constant descriptions so that it shows On. In the same dialog box, select 2 facets/edge for [/EFACET] Facets/element edge so that the elements with curved edges and surfaces are shown correctly (/EFACET command). Click on $\boldsymbol{O K}$.
- Figure 8.35 (right) shows the corresponding element plot with elements having curved edges/surfaces and thickness.
- Finally, merge duplicate entities (NUMMRG command) using the following menu path:


## Main Menu > Preprocessor > Numbering Ctrls > Merge Items

- In the dialog box, select $\boldsymbol{A l l}$ from the first pull-down menu; click on OK.


## SOLUTION

- Apply degree of freedom (DOF) constraints along the outer boundary of the aluminum plate. Since the boundary is circular, it is convenient to first switch to Cylindrical Coordinates and then select the nodes.
- Switch to Cylindrical Coordinates (CSYS command) using the following menu path:

Utility Menu > WorkPlane > Change Active CS to $>$ Global Cylindrical

- Select nodes along the circular boundary (NSEL command) by using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; choose Nodes in the first pull-down menu and By Location in the second pull-down menu. Click on $X$ coordinate and From Full radio-buttons; type 20 in the Min, Max text field; click on OK. Because the active coordinate system is cylindrical, any reference to the $x$-coordinate is treated as a reference to the $r$ coordinate by ANSYS.
- Now, apply DOF constraints (D command) by using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; click on Pick All.
- In the new dialog box, highlight All DOF; click on $\boldsymbol{O K}$.
- Select everything (ALLSEL command) using the following menu path:


## Utility Menu > Select > Everything

- Switch back to Cartesian Coordinates (CSYS command) using the following menu path:

Utility Menu $>$ WorkPlane $>$ Change Active CS to $>$ Global Cartesian

- Apply degree of freedom (DOF) constraints along the top surface of the steel piston.
- Select nodes (NSEL command) by using the following menu path:


## Utility Menu > Select > Enitities

- Select Entities dialog box appears; choose By Location in the second pull-down menu; click on the $Z$ coordinate and From Full radiobuttons; type 5 in the Min, Max text field; click on OK.
- Apply DOF constraints (D command) by using the following menu path:
- Pick Menu appears; click on Pick All.
- In the new dialog box, remove the highlight on All DOF and highlight UZ. Enter $\mathbf{- 0 . 1}$ in the text box for VALUE Displacement value; click on $\boldsymbol{O K}$.
- Apply symmetry conditions along the $x=0$ and $y=0$ planes for the entire structure.
- Select nodes (NSEL command) by using the following menu path: Utility Menu > Select > Entities
- Select Entities dialog box appears; choose Nodes in the first pull-down menu and By Location in the second pull-down menu. Click on the $X$ coordinate and From Full radio-buttons; type 0 in the Min, Max text field; click on $O K$.
- Apply DOF constraints (D command) by using the following menu path:
Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes
- Pick Menu appears; click on Pick All.
- In the new dialog box, remove the highlight on $\boldsymbol{U Z}$ and highlight $\boldsymbol{U X}$. Enter 0 in the text box for VALUE Displacement value; click on $O K$.
- Select nodes (NSEL command) by using the following menu path: Utility Menu > Select > Entities
- Select Entities dialog box appears; choose Nodes in the first pull-down menu and By Location in the second pull-down menu. Click on the $\boldsymbol{Y}$ coordinate and From Full radio-buttons; type 0 in the Min, Max text field; click on $O K$.
- Apply DOF constraints (D command) by using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; click on Pick All.
- In the new dialog box, remove the highlight on $\boldsymbol{U X}$ and highlight $\boldsymbol{U Y}$; click on $\mathbf{O K}$.
- Select everything (ALLSEL command) using the following menu path: Utility Menu > Select > Everything
- Obtain the solution using the following menu path:

Main Menu $>$ Solution $>$ Solve $>$ Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on $\boldsymbol{O K}$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the deformed shape using the following menu path:

Main Menu $>$ General PostProc $>$ Plot Results $>$ Deformed Shape

- Select Def + undef edge; click on $O K$.
- The isometric view of the deformed shape is shown in Fig. 8.36 as it appears in the Graphics Window.
- Review the equivalent stress (von Mises) contour plot (PLNSOL command) using the following menu path:

Main Menu > General PostProc > Plot Results >Contour Plot $>$ Nodal Solu

- Contour Nodal Solution Data dialog box appears. Select Stress from the left list; in the right list, scroll down to select von Mises SEQV. Click on $O K$.
- Figure 8.37 shows the corresponding contour plot.


### 8.1.5.3 Analysis of an Axisymmetric Shell with Internal Pressure

Consider the pressure vessel shown in Fig. 8.38 with elastic properties $E=10 \times 10^{6}$ psi and $v=0.3$. Its radius changes, as shown in Fig. 8.38, while the thickness remains constant, $t=0.25 \mathrm{in}$. The internal pressure is 300 psi . The goal is to find the meridional and hoop stresses in the shell. Examination of the geometry and loading reveals that the problem is axisymmetric. Therefore, axisymmetric shell elements within ANSYS are utilized in this section.

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on $A d d$.
- Select Shell in the left list and 2D axisymmetr 51 in the right list; click on $O K$.
- Click on Close.
- Specify the thickness using real constants ( R command) using the following menu path:

[^15]

Fig. 8.36 Isometric view of the deformed shape.


Fig. 8.37 Equivalent stress contours.


Fig. 8.38 Schematic of the axisymmetric shell with internal pressure.

- Click on Add.
- Click on $\boldsymbol{O K}$; dialog box appears.
- Type 0.25 in the Shell thickness at node I TK(I) text field; click on OK.
- Click on Close.
- Specify material properties (MP command) using the following menu path:


## Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring up another dialog box.
- Enter $\mathbf{1 0 e 6}$ for $\boldsymbol{E X}$ and $\mathbf{0 . 3}$ for $\boldsymbol{P R X Y}$; click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

```
Material > Exit
```

- Create keypoints ( K command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Keypoints > In Active CS

- In the Create Keypoints in Active Coordinate System dialog box, type, 25 for $\boldsymbol{X}$ and $\mathbf{0}$ for $\boldsymbol{Y}$; click on Apply (keypoint 1 is created).
- Referring to the schematic of keypoints shown in Fig. 8.39 and tabulated in Table 8.1, repeat this procedure for keypoints 2 through 6.


Fig. 8.39 Schematic of the keypoints used in the solid model.

Table 8.1 Keypoint numbers and coordinates for the axisymmetric shell.

| Keypoint <br> Number | $x$ | $y$ |
| :---: | :---: | :---: |
| 1 | 25 | 0 |
| 2 | 25 | 10 |
| 3 | 15 | 27.32 |
| 4 | 15 | 37.32 |
| 5 | 0 | 52.32 |
| 6 | 0 | 37.32 |

- Create straight lines ( L command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Lines > Lines > Straight Line

- Pick Menu appears; pick keypoints 1 and 2 ; line 1 is created.
- Repeat this for lines 2 and 3 using keypoint pairs 2-3 and 3-4, respectively; click on $O K$ in the Pick Menu.
- Create an arc (LARC command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Lines > Arcs > By End KPs \& Rad

- Pick Menu appears; pick keypoints 4 and 5 (end points of the arc); click on $O K$ in the Pick Menu.
- Pick keypoint 6 (center of the arc); click on $O K$ in the Pick Menu.
- Arc by End KPs \& Radius dialog box appears; enter 15 for RAD Radius of the arc.
- Click on $O K$; line 4 is created.
- Specify the number of divisions on the lines (LESIZE command) using the following menu path:


## Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Lines > All Lines

- Element Sizes on Picked Lines dialog box appears; enter 20 for NDIV; click on $O K$.
- Create the mesh (LMESH command) using the following menu path:


## Main Menu > Preprocessor > Meshing > Mesh > Lines

- Pick Menu appears; click on Pick All.


## SOLUTION

- Apply degree of freedom (DOF) constraints at end points (D command) using the following menu path:


## Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; pick the bottom node, i.e., $(x, y)=(25,0)$; click on OK.
- In the dialog box, select $\boldsymbol{U Y}$; click on Apply.
- Pick Menu reappears; pick the top node, i.e., $(x, y)=(0,52.32)$; click on OK.
- In the dialog box, remove the highlight on $\boldsymbol{U Y}$ and highlight $\boldsymbol{U X}$; click on $\mathbf{O K}$.
- Apply pressure (SFE command) using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Structural $>$ Pressure $>$ On Elements

- Pick Menu appears; click on Pick All.
- In the new dialog box, enter 300 for Value Load PRES value; click on OK.

Obtain the solution (SOLVE command) using the following menu path:
Main Menu >Solution > Solve >Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on $O K$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the deformed shape (PLDISP command) using the following menu path:

Main Menu > General PostProc > Plot Results > Deformed Shape

- Select Def + undef edge; click on OK.
- The deformed shape is shown in Fig. 8.40 (left) as it appears in the Graphics Window. It is clear from the figure that the bottom end of the conical section exhibits unexpected displacements/rotations. Problems with real applications (using realistic material properties, geometry, and loads) seldom produce displacements that can be visually detected. Therefore, ANSYS scales the displacements when displaying the deformed shape.


Fig. 8.40 Deformed shape with automatic scaling (left; amplified $\sim 23$ times) and manual scaling (right; amplified 5 times).

- Change the displacement scaling (/DSCALE command) using the following menu path:

Utility Menu > PlotCtrls > Style > Displacement Scaling

- This brings up the Displacement Display Scaling dialog box. Note the number 22.5304726012 in the User specified factor field. This means that the displacements are amplified by a factor of approximately 22 , so they can be clearly viewed. In order to change this setting, click on the User specified radio-button; replace the existing scaling factor with the desired value. Figure 8.40 (right) shows the deformed shape amplified by a factor of 5 .
- Store element stresses in the element table (ETABLE command) using the following menu path:


## Main Menu > General PostProc > Element Table > Define Table

- Element Table Data dialog box appears; click on Add.
- Define Additional Element Table Items dialog box appears; assign a user-defined label for the plate mid-plane meridional stresses, say STMR, in the Lab User label for item field.
- In the left list, scroll down and select By sequence num; in the right list, select $L S$.
- Finally, in the last field, type $L S, 5$; click on Apply. There are several quantities that are stored in sequences, i.e., LS, NMISC, or SMISC. The information as to which quantity is stored under which sequence is given in element help pages. In this particular example, the help page
for SHELL51 contains tables explaining which quantities are stored under which sequence.
- Similarly, store plate mid-plane hoop stresses by assigning a user label, say $\boldsymbol{S T H P}$, and entering $\boldsymbol{L S}, 7$; click on $\boldsymbol{O K}$.
- Click on Close in the Element Table Data dialog box.

■ View element table quantities (PRETAB command) using the following menu path:

## Main Menu > General PostProc > Element Table > List Elem Table

- List Element Table dialog box appears; select STMR and STHP; click on $\boldsymbol{O K}$.
- Element numbers and requested element table items are displayed in columns in a separate window. Figures 8.41 and 8.42 show the meridional and hoop stresses, respectively (plotted outside ANSYS).


### 8.1.5.4 Analysis of a Layered Composite Plate

A 10 in $\times 10$ in square composite plate with a stacking sequence of $\left[45^{\circ} /\right.$ $0^{\circ} /-45^{\circ} / 90^{\circ}$ ] is subjected to tensile loading of 100 MPa in the $y$-direction, as shown in Fig. 8.43. Unidirectional ply properties are $E_{L}=161 \mathrm{GPa}$, $E_{T}=9 \mathrm{GPa}, v_{L T}=0.26$, and $G_{L T}=6.1 \mathrm{GPa}$. The subscripts $L$ and $T$ designate longitudinal (fiber direction) and transverse (perpendicular to fiber direction), respectively. Each ply has a thickness of 0.16 mm . The goal is to find the displacement and stress fields in the plate.

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:


## Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Shell in the left list and Linear Layer 99 in the right list; click on $\boldsymbol{O K}$.
- Click on Close.
- Specify layer information using real constants ( $R$ command) using the following menu path:


## Main Menu > Preprocessor > Real Constants $>$ Add/Edit/Delete

- Real Constants dialog box appears; click on Add.
- Element Type for Real Constants dialog box appears; click on OK.
- Real Constants Set Number 1, for SHELL99 dialog box appears; click on $\boldsymbol{O K}$.


Fig. 8.41 Meridional stresses in the axisymmetric shell.


Fig. 8.42 Hoop stresses in the axisymmetric shell.


Fig. 8.43 Schematic of the layered composite plate.

- Another dialog box appears; type 4 in the Number of layers ( 250 max ) $N L$ text field; click on $O K$.
- Dialog box for layer information appears; material number (MAT), orientation angle (THETA), and thickness (TK) information for each layer (ply) are entered in this dialog box. Enter the related quantities, as shown in Fig. 8.44; click on $\boldsymbol{O K}$.
- Exit from Real Constants dialog box by clicking on Close .

Specify material properties (MP command) using the following menu path:

## Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Orthotropic, which will bring up another dialog box.
- Enter 161e9 for $E X, 9 e 9$ for $E Y$ and $E Z, 0.26$ for $P R X Y$ and $P R X Z$, 0.01 for PRXY, 6.1e9 for GXY and GXZ, and $1 e 9$ for $G Y Z$; click on OK.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create keypoints ( K command) using the following menu path:

Main Menu > Preprocessor $>$ Modeling $>$ Create $>$ Keypoints $>\operatorname{In}$ Active CS

- In the Create Keypoints in Active Coordinate System dialog box, type, $\mathbf{- 5}$ for $\boldsymbol{X}$ and $\mathbf{- 5}$ for $\boldsymbol{Y}$; click on Apply (keypoint 1 is created).


Fig. 8.44 Dialog box for entering layer information.

- Repeat this procedure for keypoints 2 through 4 using the following data:

$$
\begin{array}{ll}
x_{2}=5 & y_{2}=-5 \\
x_{3}=5 & y_{3}=5 \\
x_{4}=-5 & y_{4}=5
\end{array}
$$

- After creating keypoint 4, click on $O K$ instead of Apply.
- Create the area ( $\mathbf{A}$ command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create $>$ Areas $>$ Arbitrary $>$ Through KPs

- Pick Menu appears; pick keypoints 1, 2, 3, and 4 in this order; click on OK in the Pick Menu.
- Specify the number of divisions on all lines (LESIZE command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Size Cntrls $>$ ManualSize $>$ Lines $>$ All Lines

- Element Sizes on All Selected Lines dialog box appears; enter 40 for NDIV; click on $O K$.
- Create the mesh (LMESH command) using the following menu path: Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 sided
- Pick Menu appears; click on Pick All.


## SOLUTION

- Apply degree of freedom (DOF) constraints at the center node and right mid-node ( $\mathbf{D}$ command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; pick the center node, i.e., $(x, y)=(0,0)$; click on OK.
- In the dialog box, select $\boldsymbol{U X}$ and $\boldsymbol{U Y}$; click on Apply.
- Pick Menu reappears; pick the right mid-node, i.e., $(x, y)=(5,0)$; click on $\boldsymbol{O K}$.
- In the dialog box, remove the highlight on $\boldsymbol{U X}$ (leaving only $\boldsymbol{U Y}$ highlighted); click on $O K$.
- Constrain $z$ degrees of freedom (DOF) in all nodes (D command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; click on Pick All.
- In the dialog box, remove the highlight on $\boldsymbol{U Y}$ and highlight $\boldsymbol{U Z}$; click on $O K$.
- Plot nodes for clarity (NPLOT command) using the following menu path: Utility Menu > Plot > Nodes
- Apply pressure (SFE command) using the following menu path:

Main Menu >Solution > Define Loads > Apply > Structural $>$ Pressure $>$ On Nodes

- Pick Menu appears; using the Box radio-button in the Pick Menu, pick the nodes along $y=5$ and $y=-5$; click on $O K$.
- In the new dialog box, enter $-100 E 6^{*} 4^{*} .16 E-3$ for Value Load PRES value; click on OK.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu >Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the deformed shape (PLDISP command) using the following menu path:
Main Menu > General PostProc > Plot Results > Deformed Shape
- Select Def + undef edge; click on $O K$.
- The deformed shape is shown in Fig. 8.45 as it appears in the Graphics Window.
- Obtain a contour plot of the $y$-displacement ( $u_{y}$ ) (PLNSOL command) using the following menu path:

Main Menu > General Postproc > Plot Results > Contour Plot > Nodal Solu

- Highlight DOF solution in the left list and highlight Translation UY in the right list; click on $O K$.
- The contour plot appears in the Graphics Window, as shown in Fig. 8.46 .


Fig. 8.45 Deformed shape of the composite plate under uniaxial tension in the $y$-direction.

$\begin{array}{llllllllll}-.008204 & -.006381 & -.004550 & -.002735 & & -.912 \mathrm{E}-03 & .912 \mathrm{E}-03 & .002735 & .004558 & .006361\end{array}$
Fig. 8.46 Contour plot of the $y$-displacement of the composite plate.

The same problem can be solved following the procedure given in Sec. 8.1.4.1.2 using the following equivalent orthotropic material properties for the laminate:

$$
\begin{aligned}
& E_{x}=E_{y}=61.7 \mathrm{GPa} \\
& v_{x y}=0.3 \\
& G_{x y}=23.8 \mathrm{GPa}
\end{aligned}
$$

The contour plot for the $y$-displacement ( $u_{y}$ ) obtained by using this twodimensional approximation is shown in Fig. 8.47.


Fig. 8.47 Contour plot of the $y$-displacement of the composite plate utilizing two-dimensional plane stress idealization with orthotropic properties.

### 8.2 Linear Buckling Analysis

If the component is expected to exhibit structural instability, the search for the load that causes structural bifurcation is referred to as a "buckling load" analysis. Because the buckling load is not known a priori, the finite element equilibrium equations for this type of analysis involve the solution of homogeneous algebraic equations whose lowest eigenvalue corresponds to the buckling load, and the eigenvector represents the primary buckling mode.

There are two approaches in the ANSYS program for buckling analysis: (i) eigenvalue buckling (linear) and (ii) non-linear buckling. The first is considered here.

Eigenvalue buckling is used for calculating the theoretical buckling load of a linear elastic structure. Since it assumes the structure exhibits linearly elastic behavior, the predicted buckling loads are overestimated (unconservative). Steps involved in a typical Eigenvalue Buckling analysis are:

Build the model.
Obtain the static solution.
Obtain the eigenvalue buckling solution.
Expand the solution.
Review the results.

A static solution is needed to establish the stiffening of the structure under the applied load (stress stiffening). There are several buckling modes (theoretically, infinitely many!) in a structure. The first buckling mode is the one requiring the smallest load. The user specifies the number of buckling modes to be extracted. An eigenvalue buckling solution simply calculates the buckling loads for each of these modes. The solution is then expanded to include the deformation patterns in the structure (mode shapes) corresponding to the buckling loads. Results are reviewed in the General Postprocessor.

As an example, a rectangular plate is subjected to uniform compressive loading along its top edge while the bottom edge is constrained to move in the direction of loading, as shown in Fig. 8.48. The plate is 0.45 m long, 0.3 m wide, and 0.003 m thick. It is made of steel with elastic modulus $E=200$ GPa and Poisson's ratio $v=0.32$. The goal is to find the first four buckling modes and their corresponding buckling loads under given constraints and loading configuration.

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Shell in the left list and Elastic $4 n o d e 63$ in the right list; click on $O K$.
- Click on Close.


Fig. 8.48 Schematic of the rectangular plate.

- Specify the thickness using real constants ( $\mathbf{R}$ command) using the following menu path:

Main Menu > Preprocessor > Real Constants > Add/Edit/Delete

- Click on $\operatorname{Add}$.
- Click on $O K$; dialog box appears.
- Type 0.003 in the Shell thickness at node I TK(I) text field; click on OK.
- Click on Close.

Specify material properties (MP command) using the following menu path:

## Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring up another dialog box.
- Enter 200e9 for $\boldsymbol{E X}$ and $\mathbf{0 . 3 2}$ for PRXY; click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create the solid model, a rectangular area in this case, using the following menu path:


## Main Menu > Preprocessor > Modeling > Create > Areas > Rectangle > By Dimensions

- In the Create Rectangle by Dimensions dialog box, type 0 for X1, 0.3 for $X 2,0$ for $Y 1$, and 0.45 for $Y 2$; click on $O K$.
- Turn line numbering on using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Place a checkmark in the square box next to LINE Line numbers; click on OK.
- Plot lines using the menu path:

Utility Menu > Plot > Lines

- Specify the number of elements on selected lines for mapped meshing. On lines 1 and 3, use 15 divisions; on lines 2 and 4, use 25 divisions. Use the following menu path for this action:

Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Lines > Picked Lines

- Pick lines 1 and 3; click on $\boldsymbol{O K}$.
- Element Sizes on Picked Lines dialog box appears; type 15 in the text field corresponding to NDIV (the second text field), and uncheck the first checkbox; click on Apply.
- Repeat this procedure for the next set of lines (2 and 4) with their corresponding number divisions as 25 . After specifying the number of divisions for lines 2 and 4, click on $O K$ in the Element Sizes on Picked Lines dialog box.
- Create the mesh using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 sided

- Pick the area; click on $O K$.
- The mesh should appear in the Graphics Window, as shown in Fig. 8.49.


## SOLUTION

- Constrain the out-of-plane displacements (z-displacements) of the nodes along the entire boundary using the following menu path:

```
Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes
```

- In the Pick Menu click on the radio-button next to Box. This enables the user to pick several nodes at a time by drawing an area in the Graphics Window. Move the mouse pointer to a location slightly left and above the top-left corner of the meshed area.
- Click on the left mouse button (without releasing it) and draw a rectangle that encloses only the nodes along the $x=0$ boundary; release the left button (Fig. 8.50). Observe that each selected node is identified by a small square.
- Similarly, select nodes along all boundaries $(x=0.3, y=0$, and $y=$ 0.45 ); click on $O K$ in the Pick Menu.
- Highlight $U Z$; click on $O K$.

■ Apply remaining displacement constraints in the same manner as in the previous step.

- Constrain displacements in the $y$-direction along the $y=0$ boundary.
- Constrain the displacement in the $x$-direction at the boundary point $(x, y)=(0,0)$.

■ Apply the uniform load along the $y=0.45$ boundary using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Pressure > On Nodes

- Pick the nodes along the $y=0.45$ boundary; click on $O K$.


Fig. 8.49 Mesh of the rectangular plate.


Fig. 8.50 Selecting nodes using the Box option.

- Type $\mathbf{1}$; click on $\boldsymbol{O K}$.
- The eigenvalue buckling analysis calculates a scaling factor for the existing loads; therefore, if a unit load is applied, the scaling factor yields the buckling load.
- Turn on pre-stress effects using the following menu path:

Main Menu > Solution > Analysis Type > Sol'n Controls

- A dialog box appears; place a checkmark in the box next to Calculate prestress effects; click on $O K$ (Fig. 8.51).
- Obtain the static solution (SOLVE command) using the following menu path:

Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on $O K$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!
- Exit Solution Processor using the following menu path:

Main Menu > Finish

- Re-enter the Solution Processor and change the analysis type to eigenvalue buckling using the following menu path:


## Main Menu > Solution > Analysis Type > New Analysis

- Click on Eigen Buckling; click on OK.


Fig. 8.51 Solution Controls dialog box (Basic tab shown).

Set analysis options using the following menu path:
Main Menu > Solution > Analysis Type > Analysis Options

- Type 4 in the text field next to NMODE No. of modes to extract; click on $\boldsymbol{O K}$.
- In the new dialog box, click on $\boldsymbol{O K}$, leaving the settings at their default values.
- Instruct ANSYS to expand modes using the following menu path:

Main Menu > Solution > Load Step Opts > ExpansionPass > Single Expand $>$ Expand Modes

- Type 4 in the text field next to NMODE No. of modes to extract; click on $\boldsymbol{O K}$.
- Obtain the eigenvalue buckling solution (SOLVE command) using the following menu path:


## Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on $\boldsymbol{O K}$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the buckling loads using the following menu path:

Main Menu > General PostProc > Results Summary

- The list will appear in a new window, as shown in Fig. 8.52. The critical load for the first mode is given as $0.23788 \mathrm{E}+06 \mathrm{~Pa}(0.23788$ MPa ) in this list. This means that when the applied load $p$ is increased to this value, the plate will buckle in the first mode.
- Review the buckling modes.
- Read the results for the first buckling load using the following menu path:

Main Menu > General PostProc > Read Results > First Set

- Obtain contour plot of the z-displacement using the following menu path:

Main Menu > General Postproc > Plot Results > Contour Plot > Nodal Solu

- Highlight DOF solution in the left list and $U Z$ in the right list; click on OK.
- The contour plot will appear in the Graphics Window, as shown in Fig. 8.53.
- Read the results for the second buckling load using the following menu path:

```
Main Menu > General PostProc > Read Results > Next Set
```

- Obtain contour plot of the $z$-displacement using the following menu path:

Main Menu > General Postproc $>$ Plot Results $>$ Contour Plot $>$ Nodal Solu

- Highlight DOF solution in the left list and $\boldsymbol{U Z}$ in the right list; click on OK.
- The contour plot will appear in the Graphics Window, as shown in Fig. 8.54.
- Repeat for modes 3 and 4 to obtain plots similar to those given in Fig. 8.55 and 8.56 .
- Review the buckling mode shapes.
- Read the results for the desired mode (as shown in previous step) and plot the deformed shape using the following menu path:


## Main Menu > General Postproc > Plot Results > Deformed Shape

- Click on the Def shape only radio-button; click on OK.


Fig. 8.52 List of buckling loads for different buckling modes.


Fig. 8.53 Contour plot of $u_{z}$ field (z-displacement) under first buckling mode.

```
FRBG=257694
UZ (AVG)
RgYg=0
DMX =.095028
9MX =.095028
```



Fig. 8.54 Contour plot of $u_{z}$ field ( $z$-displacement) under second buckling mode.


Fig. 8.55 Contour plot of $u_{z}$ field ( $z$-displacement) under third buckling mode.

```
FREQ=505725
```

UZ (AVG)
RSYS=0
DMX $=.035874$
SMN $=-.035874$
$\operatorname{sex}=.035867$


Fig. 8.56 Contour plot of $u_{z}$ field ( $z$-displacement) under fourth buckling mode.

- Change the viewpoint to isometric using the menu path:


## Utility Menu > PlotCtris > Pan Zoom Rotate

- In the Pan Zoom Rotate window, click on the Iso button.
- Figure 8.57 shows the first four mode shapes of the plate.


### 8.3 Thermomechanical Analysis

Thermal strains and stresses constitute an important part of the design considerations for many practical engineering problems. They become especially critical when materials with different coefficients of thermal expansion form interfaces.

As an example of a thermomechanical analysis with ANSYS, consider an electronic device containing a silicon die (chip), epoxy die-attach substrate, and a molding compound, as shown in Fig. 8.58. A common cause of failure in electronic devices is the thermal stresses at elevated temperatures caused by a coefficient of thermal expansion mismatch. In the ANSYS solution, plane strain idealization is utilized. The device is subjected to a uniform temperature increase of $30^{\circ} \mathrm{C}$. Material properties of the constituent materials are given in Table 8.2. The goal is to obtain displacement and stress fields.


Fig. 8.57 Buckling modes 1 through 4 (from left to right).


Fig. 8.58 Geometry of the electronic package.

Table 8.2 Properties of the constituent materials in the electronic package.

|  | $E$ <br> $(\mathrm{GPa})$ | $v$ | $\alpha$ <br> $\left(10^{-6} /{ }^{\circ} \mathrm{C}\right)$ | Material <br> Reference <br> Number |
| :--- | :---: | :---: | :---: | :---: |
| Substrate | 22 | 0.39 | 18 | 1 |
| Die-attach | 7.4 | 0.4 | 52 | 2 |
| Silicon | 163 | 0.278 | 2.6 | 3 |
| Molding compound | 15 | 0.25 | 16 | 4 |

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Solid in the left list and Quad 4 Node 42 in the right list; click on $\boldsymbol{O K}$.
- Click on Options.
- PLANE42 element type options dialog box appears; select Plane strain item from the pull-down menu corresponding to Element behavior K3.
- Click on $O K$; click on Close.

Specify material properties (MP command) using the following menu path:

Main Menu > Preprocessor > Material Props $>$ Material Models

- Define Material Model Behavior dialog box appears. In the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which brings up another dialog box.
- Referring to Table 8.2, enter 22E9 for EX and 0.39 for PRXY; click on OK.
- In the right list, successively double-click on Structural, Thermal Expansion, Secant Coefficient, and, finally, Isotropic, which brings up another dialog box.
- Enter $18 E-6$ for $\boldsymbol{A L P X}$; click on $O K$.
- Add new material model using the following menu path:

Material > New Model

- Repeat the procedure for the remaining materials (2 through 4) referring to Table 8.2.
- When finished, close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create rectangles as identified in Fig. 8.59 (rectivg command) using the following menu path:
Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Rectangle $>$ By Dimensions
- Create Rectangle by Dimensions dialog box appears. Referring to Table 8.3, enter $\mathbf{0}$ for $\boldsymbol{X 1}, \mathbf{5 E}-\mathbf{3}$ for $\boldsymbol{X 2}, \mathbf{0}$ for $\boldsymbol{Y 1}$, and $\mathbf{2 E} \mathbf{- 3}$ for $\boldsymbol{Y 2}$; click on Apply.
- Repeat the procedure for the remaining areas (2 through 9 ). When creating Area 9 , click on $O K$ after entering the coordinates.


Fig. 8.59 Solid model of the electronic package.

Table 8.3 Coordinates defining the areas and the corresponding material reference numbers.

| Area | X1 | X2 | Y1 | $Y 2$ | Material Reference |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Number | -------------- (mm) ---------------- |  |  |  | Number |
| 1 | 0 | 5 | 0 | 2 | 1 |
| 2 | 5 | 7.5 | 0 | 2 | 1 |
| 3 | 7.5 | 10 | 0 | 2 | 1 |
| 4 | 0 | 5 | 2 | 2.1 | 2 |
| 5 | 5 | 7.5 | 2 | 2.1 | 4 |
| 6 | 0 | 5 | 2.1 | 3.1 | 3 |
| 7 | 5 | 7.5 | 2.1 | 3.1 | 4 |
| 8 | 0 | 5 | 3.1 | 4.1 | 4 |
| 9 | 5 | 7.5 | 3.1 | 4.1 | 4 |

- Glue the areas (AGLUE command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Operate $>$ Booleans $>$ Glue $>$ Areas

- Pick Menu appears; click on Pick All button.
- Mesh the areas (AMESH command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 sided

- Pick Menu appears; click on Pick All.
- At this point, all the elements have Material Reference Number 1. Attributes can be changed after the elements are created. For this purpose, areas are selected first. Then the elements that are attached to the selected areas are selected. Finally, elements are modified so they have the correct attributes. The correspondence between the areas and
material numbers are given in Table 8.3. Select areas (ASEL command) using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; select Areas from the first pulldown menu; click on $\boldsymbol{O K}$.
- Pick Menu appears; pick areas 5, 7, 8, and 9; click on $O K$.
- Now, select the elements that are attached to the selected areas (ESLA command) using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; select Elements from the first pulldown menu; select Attached to from the second pull-down menu. Click on the Areas radio-button; click on $\boldsymbol{O K}$.
- Modify the attributes of the selected set of elements (EMODIF command) using the following menu path:

Main Menu > Preprocessor > Modeling > Move / Modify > Elements > Modify Attrib

- Pick Menu appears; click on Pick All, which brings up the Modify Elem Attributes dialog box.
- Select Material MAT from the pull-down menu and enter 4 in the 11 New attribute number field; click on $O K$.
- Repeat this procedure for area 4 (material reference number 2 ) and area 6 (material reference number 3).
- When finished, select everything (allsel command) using the following menu path:

Utility Menu > Select > Everything

## SOLUTION

- Apply displacement constraints (D command) using the following menu path:


## Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; pick the nodes along $x=0$ ( $y$-axis); click on $\boldsymbol{O K}$ in the Pick Menu.
- Highlight $\boldsymbol{U X}$; click on Apply.
- Pick Menu reappears; pick the bottom-left corner node ( $x=0, y=0$ ); click on $O K$ in the Pick Menu.
- Remove highlight $\boldsymbol{U Y}$ (leave the $\boldsymbol{U X}$ highlighted); click on $\boldsymbol{O K}$.
- Apply the thermal load (TUNIF command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Temperature > Uniform Temp

- Uniform Temperature dialog box appears; enter 30 for TUNIF; click on OK.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu >Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

■ Review the deformed shape (PLDISP command) using the following menu path:

Main Menu > General PostProc > Plot Results > Deformed Shape

- Select Def + undef edge; click on OK.
- The deformed shape is shown in Fig. 8.60 as it appears in the Graphics Window.

■ Obtain the normal stress in the $y$-direction and shearing stress contour plots (PLNSOL command) using the following menu path:

Main Menu > General Postproc > Plot Results > Contour Plot > Nodal Solu

- In order to obtain the view of the normal stresses in the $y$-direction, highlight Stress in the left list (second item) and scroll down to highlight $Y$-direction $S Y$ in the right list; click on $O K$.
- The resulting contour plot, along with a zoomed-in view of the critical junction, is shown in Fig. 8.61.
- Similarly, in order to view the shear stresses, highlight Stress in the left list (second item) and scroll down to highlight $X Y$-shear $S X Y$ in the right list; click on $O K$.
- The resulting contour plot, along with a zoomed-in view of the critical junction, is shown in Fig. 8.62.
- Plot elements (EPLOT command) using the following menu path:


Fig. 8.60 Deformed shape of the electronic package under thermal load.


Fig. 8.61 Contour plot of the normal stress ( $\sigma_{y y}$ ) in the $y$-direction: in the entire package (left) and in the vicinity of the die/die-attach interface (right).


Fig. 8.62 Contour plot of the shear stress $\left(\sigma_{x y}\right)$ : in the entire package (left) and in the vicinity of the die/die-attach interface (right).

- Review variation of stresses along paths by means of line plots. Two paths are defined, both of which are vertical. The first path passes through the vertical cross section where the die and the die-attach terminate and form an interface with the molding compound ( $x=5 \mathrm{~mm}$.). The second path is located approximately in the middle of the die and die-attach. Both paths are plotted in Fig. 8.63 and 8.64 (element edges are removed in Fig. 8.64 for clarity). Define the path (PPATH command) using the following menu path:


## Main Menu > General Postproc > Path Operations > Define Path > By Nodes

- Pick Menu appears; pick the two nodes indicated with small squares, as shown in Fig. 8.63 (corresponding to the path V1); click on OK.
- By Nodes dialog box appears; enter a name describing the path, say V1, in the Define Path Name text field; click on OK.
- Close the Path Command Status Window.
- Define a second path as indicated in Fig. 8.63 and 8.64 (corresponding to the path V2); enter the name as V2.
- When multiple paths are defined, only one path is active at a given time, and mapping of results is performed on the active path. Activate the path V1 (PATH command) using the following menu path:


## Main Menu > General Postproc > Path Operations > Recall Path

- Recall Path dialog box appears. Select V1; click on OK.
- Map results onto path (PDEF command) using the following menu path:


## Main Menu > General Postproc > Path Operations > Map onto Path

- Map Result Items onto Path dialog box appears; select Stress from the left list and $Y$-direction $S Y$ from the right list; click on Apply.
- Map Result Items onto Path dialog box remains active; select Stress from the left list and $\boldsymbol{X Y}$-shear $\boldsymbol{S X Y}$ from the right list; click on Apply.
- Now, select Stress from the left list and scroll down in the right list to select von Mises SEQV; click on OK.
- At this point, normal stress in the $y$-direction ( $\sigma_{y y}$ ), $x y$ shear stress ( $\sigma_{x y}$ ), and equivalent stress ( $\sigma_{e q v}$ ) values are mapped onto path V1. Obtain line plot of $\sigma_{y y}$ and $\sigma_{x y}$ along the path V1 (PLPATH command) using the following menu path:

Main Menu > General Postproc > Path Operations > Plot Path Item > On Graph

- Plot of Path Items on Graph dialog box appears; select $\boldsymbol{S Y}$ and $\boldsymbol{S X Y}$; click on $O K$.
- Figure 8.65 shows the line plots of $\sigma_{y y}$ and $\sigma_{x y}$ along the defined path.


Fig. 8.63 Element plot with paths V1 and V2 identified.


Fig. 8.64 Element plot (element edges removed) with paths V1 and V2 identified.


Fig. 8.65 Line plots of $\sigma_{y y}$ and $\sigma_{x y}$ along path V1.

- Now, obtain line plot of $\sigma_{e q v}$ in the same graph with $\sigma_{y y}$ and $\sigma_{x y}$ along path V1 (PLPATH command) using the following menu path:
Main Menu > General Postproc > Path Operations > Plot Path Item $>$ On Graph
- Plot of Path Items on Graph dialog box appears; add SEQV to the existing selection ( $\boldsymbol{S Y}$ and $\boldsymbol{S X Y}$ ); click on $\boldsymbol{O K}$.
- Figure 8.66 shows the resulting line plot.
- Similar plots can be obtained for stresses along path V2. For this purpose, the user needs to activate path V 2 , followed by the mapping of quantities. Figure 8.67 shows the variation of $\sigma_{y y}, \sigma_{x y}$, and $\sigma_{e q v}$ along the path.


### 8.4 Fracture Mechanics Analysis

Computation of fracture parameters, such as the stress intensity factors or energy release rate, using finite element analysis requires either a refined mesh around the crack tip or the use of "special elements" with embedded stress singularity near the crack tip. Although conceptually the stress intensity factors are obtained in a straightforward manner, finite element analyses with conventional elements near the crack tip always underestimate the sharply rising stress-displacement gradients.

Instead of trying to capture the well-known $1 / \sqrt{r}$ singular behavior with smaller and smaller elements, Henshell and Shaw (1975) and Barsoum $(1976,1977)$ introduced a direct method by shifting the mid-side node of an 8 -noded isoparametric quadrilateral element to the one-quarter point from the crack tip node. Relocating the mid-side nodes to the one-quarter point achieves the desired $1 / \sqrt{r}$ singular behavior. In the case of linear elastic deformation, the elements PLANE2 (2-D, 6-noded triangle), PLANE82 (2D, 8-noded quadrilateral), and SOLID95 (3-D, 20-noded brick) in ANSYS are used to obtain the well-established singular stress field by shifting the mid-side nodes one-quarter away from the crack tip.

Once an accurate stress field is obtained, fracture parameters (i.e., stress intensity factors, $J$-integral, and energy release rate) can be calculated within the ANSYS postprocessor.

As an extension of the node collapsing approach, Pu et al. (1978) showed that the stress intensity factors, $K_{I}$ and $K_{I I}$ for opening and sliding modes, respectively, can be computed directly from the nodal displacements on opposite sides of the crack plane as


Fig. 8.66 Line plots of $\sigma_{y y}, \sigma_{x y}$, and $\sigma_{e q v}$ along path V1.


Fig. 8.67 Line plots of $\sigma_{y y}, \sigma_{x y}$, and $\sigma_{e q v}$ along path V2.

$$
K_{I}=G \sqrt{\frac{2 \pi}{r_{0}}} \frac{u_{y}\left(r_{0}, \theta=\pi\right)-u_{y}\left(r_{0}, \theta=-\pi\right)}{(\kappa+1)}
$$

and

$$
K_{I I}=G \sqrt{\frac{2 \pi}{r_{0}}} \frac{u_{x}\left(r_{0}, \theta=\pi\right)-u_{x}\left(r_{0}, \theta=-\pi\right)}{(\kappa+1)}
$$

in which $r_{0}$, usually restricted to one or two percent of the crack length, is the distance from the crack tip to the first side-node behind the crack tip, as shown in Fig. 8.68. The shear modulus is $G$, and the parameters are $\kappa=3-v / 1+v$ and $\kappa=3-4 v$ for plane stress and strain idealizations, respectively.

Under plane strain assumptions, the computation of the stress intensity factors within ANSYS is demonstrated by considering a strip with an inclined edge crack, as shown in Fig. 8.69. The crack is 1 in long and has an inclination angle of $45^{\circ}$. The width and length of the strip are 5 in and 25 in , respectively. The bottom surface of the strip is constrained in both directions while the top surface is subjected to a tensile load of 100 psi. Elastic modulus and Poisson's ratio of the strip are $30 \times 10^{6} \mathrm{psi}$ and 0.3 , respectively.

In the ANSYS solution, special meshing around the crack tip is utilized. It relocates the mid-side nodes to one-quarter away from the crack tip. Coordinates of the keypoints are listed in Table 8.4. Note that keypoint 2 is located at the crack tip, and keypoints 1 and 3 are coincident, each belonging to the opposite crack faces. Line numbers with their corresponding keypoints are listed in Table 8.5. The goal is to obtain stress intensity factors, as well as the displacement and stress fields.


Fig. 8.68 Displacements at nodal points located behind the crack tip.


Fig. 8.69 Geometry of the strip with inclined edge crack.

Table 8.4 Coordinates of the keypoints.

| Keypoint No. | $x$ | $y$ |
| :---: | :---: | :---: |
| 1 | 0 | $12.5+\frac{\sin (45)}{2}$ |
| 2 | $\cos (45)$ | $12.5-\frac{\sin (45)}{2}$ |
| 3 | 0 | $12.5+\frac{\sin (45)}{2}$ |
| 4 | 0 | 0 |
| 5 | 5 | 0 |
| 6 | 5 | $12.5-\left\{[5-\cos (45)] \times \tan (45)+\frac{\sin (45)}{2}\right\}$ |
| 7 | 5 | 25 |
| 8 | 0 | 25 |

Table 8.5 Line-keypoint correspondence in the solid model.

| Line No. | Keypoint 1 | Keypoint 2 |
| :---: | :---: | :---: |
| 1 | 4 | 5 |
| 2 | 5 | 6 |
| 3 | 6 | 2 |
| 4 | 2 | 1 |
| 5 | 1 | 4 |
| 6 | 6 | 7 |
| 7 | 7 | 8 |
| 8 | 8 | 3 |
| 9 | 3 | 2 |

## MODEL GENERATION

- Define element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Solid on the left list and Triangle 6node 2 on the right list; click on $\boldsymbol{O K}$.
- Click on Options.
- PLANE42 element type options dialog box appears; select Plane strain item from the pull-down menu corresponding to Element behavior K3.
- Click on OK; click on Close .
- Specify material properties (MP command) using the following menu path:


## Main Menu > Preprocessor > Material Props > Material Models

- Define Material Model Behavior dialog box appears; in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which brings up another dialog box.
- In the new dialog box, enter 30E6 for EX and 0.3 for PRXY; click on OK.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

■ Change the default angular unit to degrees (*AFUN command) using the following menu path:

Utility Menu > Parameters > Angular Units

- Angular Units for Parametric Functions dialog box appears; select Degrees DEG from the pull-down menu; click on $O K$.
- Create keypoints ( K command) using the following menu path:


## Main Menu > Preprocessor > Modeling > Create $>$ Keypoints $>$ In Active CS

- Create Keypoints in Active Coordinate System dialog box appears. Referring to Table 8.4, enter 0 and $12.5+S I N(45) / 2$ for $X$ and $Y$, leaving the text fields for NPT and Z blank. Click on Apply.
- Repeat the procedure for the remaining keypoints (2 through 8 ). When creating keypoint 8 , click on $\boldsymbol{O K}$ after entering the coordinates.
- Create lines (L command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Lines $>$ Lines $>$ Straight Line

- Pick Menu appears, prompting the user to pick two keypoints forming the line. Referring to Table 8.5 , pick the correct keypoints. When picking keypoint 1 or 3 , ANSYS displays a warning message informing the user that there are two coincident keypoints at the particular location. By clicking on the Next button in this message, pick the correct keypoint.
- Turn line and keypoint numbering on (/PNUM command) using the following menu path:


## Utility Menu > PlotCtrls > Numbering

- Plot Numbering Controls dialog box appears. Click on the boxes next to KP Keypoint numbers and LINE Line numbers (this places checkmarks), and select Numbers only from the [/NUM] Numbering shown with the pull-down menu. Click on $O K$.
- Plot lines (LPLOT command) using the following menu path:

Utility Menu > Plot > Lines

- Figure 8.70 shows the line plot with both keypoint and line numbers printed. Observe that keypoints 1 and 3 and lines 4 and 9 are coincident.


Fig. 8.70 Line plot with both keypoint and line numbers printed.

- Two areas are created. The first area utilizes lines $1-5$ while the second area is formed by lines $6,7,8,9$, and 3 . Create areas using lines (AL command) using the following menu path:


## Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Arbitrary $>$ By Lines

- Pick Menu appears, prompting the user to pick lines forming the area. Pick lines 1 through 5 (in this order) and click on $O K$ in the Pick Menu. When picking line 4, ANSYS informs the user that there are two coincident lines at the picked location. Make sure to pick line 4.
- Repeat the same procedure for the second area by picking lines $6,7,8$, 9 , and 3 . Similar to the previous case, make sure to pick line 9 (instead of line 4).
- For the stress intensity factor calculations, a local coordinate system aligned with the crack faces is needed. Create a local coordinate system using 3 keypoints (CSKP command) using the following menu path:

Utility Menu > WorkPlane > Local Coordinate Systems > Create Local CS > By 3 Keypoints

- Pick Menu appears; pick keypoints 2,6, and 7 (in this order) and click on $O K$ in the Pick Menu.
- Create CS By 3 KPs dialog box appears; click on OK.
- The local coordinate system is now active. Activate the global Cartesian coordinate system (CSYS command) using the following menu path:

Utility Menu > WorkPlane > Change Active CS to $>$ Global Cartesian

- Specify keypoint 2 to be the crack tip so that the elements around it have the singular stress capability (KSCON command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Size Cntrls $>$ Concentrat KPs $>$ Create

- Pick Menu appears; pick keypoint 2; click on $\boldsymbol{O K}$ in the Pick Menu.
- Concentration Keypoint dialog box appears; enter 1/20 for DELR Radius of 1st row of elems and 6 for NTHET No of elems around circumf. Select Skewed 1/4pt from the KCTIP midside node position pull-down menu and click on $O K$.
- Specify mesh density around keypoints (KESIZE command) using the following menu path:


## Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Keypoints > All KPs

- Element Size at All Keypoints dialog box appears; enter $5 / 3$ for SIZE Element edge length; click on OK.
- Specify mesh density around specific keypoints (KESIZE command) using the following menu path:

Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize $>$ Keypoints $>$ Picked KPs

- Pick Menu appears; pick keypoints 1 and 3 (since these keypoints are coincident, click on the location twice to pick both of them); click on OK in the Pick Menu.
- Element Size at Picked Keypoints dialog box appears; enter $1 / 3$ for SIZE Element edge length; click on $O K$.
- Specify mesh density around crack tip (KESIZE command) using the following menu path:


## Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Keypoints > Picked KPs

- Pick Menu appears; pick keypoint 2; click on $O K$ in the Pick Menu.
- Element Size at Picked Keypoints dialog box appears; enter 1/30 for SIZE Element edge length; click on $O K$.
- Mesh the areas (AMESH command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Free

- Pick Menu appears; click on Pick All.
- Close the Warning Window.
- Zoom in around the crack tip and observe the mesh pattern around it (Fig. 8.71).


Fig. 8.71 Mesh pattern around the crack tip.

## SOLUTION

- Apply displacement constraints (D command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; pick the nodes along $y=0$ ( $x$-axis); click on $O K$ in Pick Menu.
- Apply U,ROT on Nodes dialog box appears; highlight UY; click on Apply.
- Pick Menu reappears; pick the bottom-left corner node ( $x=0, y=0$ ); click on $\mathbf{O K}$ in Pick Menu.
- Apply U,ROT on Nodes dialog box reappears; highlight $\boldsymbol{U X}$ (leave the $\boldsymbol{U Y}$ highlighted) and click on $\boldsymbol{O K}$.
- Apply surface load (SF command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Pressure > On Nodes

- Pick Menu appears; pick the nodes along $y=25$; click on $O K$ in Pick Мепи.
- Apply PRES on nodes dialog box appears; enter -100 for VALUE Load PRES value; click on OK.
- Obtain solution (SOLVE command) using the following menu path:

Main Menu >Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window and click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

■ Review deformed shape (PLDISP command) using the following menu path:

Main Menu > General PostProc > Plot Results > Deformed Shape

- Select Def shape only; click on OK.
- The deformed shape near the crack is shown in Fig. 8.72 as it appears in the Graphics Window.
- Activate local coordinate system 11 (CSYS command) using the following menu path:

Utility Menu > WorkPlane > Change Active CS to > Specified Coord Sys


Fig. 8.72 Deformed shape of the crack (left) and the deformed shape in the vicinity of the crack tip (right).

- Change Active CS to Specified CS dialog box appears; enter 11 for KCN Coordinate system number; click on $O K$.
- Enforce the use of the same coordinate system (11) for the results calculations and display (RSYS command) using the following menu path:

Main Menu > General Postproc > Options for Outp

- Options for Output dialog box appears; select Local system from the [RSYS] Results coord system pull-down menu and enter 11 for Local system reference no.; click on OK.
- In order to calculate stress intensity factors, a path along the crack faces in the vicinity of the crack tip is defined (PPATH command) using the following menu path:


## Main Menu > General Postproc > Path Operations > Define Path $>$ By Nodes

- Pick Menu appears; a total of 5 nodes are needed for this operation. Crack tip node needs to be picked first, followed by the two nodes closest to the crack tip along the top crack face. Finally, the two nodes closest to the crack tip along the bottom crack face are picked. The crack tip node number is 18 in this particular problem. Node numbers for the two nodes closest to the crack tip along the top and bottom faces are 47 and 48 , and 521 and 522 , respectively. Before picking the nodes, it is recommended that the user zoom in around the crack tip and plot elements as shown in Fig. 8.73. Pick nodes 18, 47, 48, 522, and 521 (in this order) and click on $\boldsymbol{O K}$. The nodal locations to be picked are also shown in Fig. 8.73 (denoted by small squares).


Fig. 8.73 Elements around the crack tip is zoomed in for picking operation.

- By Nodes dialog box appears; enter a path name (say crck); click on OK.
- Close the new information window.
- Calculate stress intensity factors (KCALC command) using the following menu path:


## Main Menu > General PostProc > Nodal Calcs > Stress Int Factr

- Stress Intensity Factor dialog box appears; select Plane strain from the KPLAN Disp extrapolat based on pull-down menu and select Fullcrack model from the KCSYM Model type pull-down menu. Click on OK.
- Stress intensity factors are reported in a separate window (KI $=139.48$, $\mathrm{KII}=70.699$ ), as shown in Fig. 8.74.
- Review normal and shear stresses ahead of the crack tip, in the direction of the crack. For this purpose, define a new path (PPATH command) using the following menu path:

Main Menu > General Postproc > Path Operations > Define Path > By Nodes

- Pick Menu appears; pick nodes 18 (crack tip) and 34, as shown in Fig. 8.75; click on $\boldsymbol{O K}$ in the Pick Menu.
- By Nodes dialog box appears; enter a path name (say strs); click on $\boldsymbol{O K}$.
- Close the new information window.
- Map stresses onto the path strs (PDEF command) using the following menu path:

[^16]

Fig. 8.74 Stress intensity factor results.


Fig. 8.75 Nodal plot showing the starting (crack tip, node 18 ) and ending (node 34) nodes for the path definition.

- Map Result Items onto Path dialog box appears; select Stress from the left list and $\boldsymbol{Y}$-direction $\boldsymbol{S Y}$ from the right list; click on Apply.
- Map Result Items onto Path dialog box reappears; select Stress from the left list and $\boldsymbol{X Y}$-shear $\boldsymbol{S X Y}$ from the right list and click on $\boldsymbol{O K}$.
- Plot stresses along the path on the graph (PLPATH command) using the following menu path:

Main Menu > General Postproc > Path Operations $>$ Plot Path Item $>$ On Graph

- Plot of Path Items on Graph dialog box appears; select $S \boldsymbol{Y}$ from the list; click on $O K$. The graph appears in the Graphics Window, as shown in Fig. 8.76. Similarly, shear stresses are plotted by selecting item $\boldsymbol{S X Y}$ in the Plot of Path Items on Graph dialog box (shown in Fig. 8.77).


Fig. 8.76 Normal stress $\sigma_{y}$ (in coordinate system 11) ahead of the crack tip.


Fig. 8.77 Shear stress $\sigma_{x y}$ (in coordinate system 11) ahead of the crack tip.

### 8.5 Dynamic Analysis

There are three commonly used dynamic analysis types in ANSYS: (i) modal analysis, (ii) harmonic analysis, and (iii) transient analysis.

The results related to these types of analyses can be reviewed in both postprocessors (General Postprocessor and Time History Postprocessor). The General Postprocessor is used to review results over the entire model at specific times or frequencies while the Time History Postprocessor allows the user to review results at specific nodes in the model over the entire time or frequency range.

### 8.5.1 Modal Analysis

If the structural vibration is of concern in the absence of time-dependent external loads, a modal analysis is performed. Because the structural frequencies are not known a priori, the finite element equilibrium equations for this type of analysis involve the solution of homogeneous algebraic equations whose eigenvalues correspond to the frequencies, and the eigenvectors represent the vibration modes. The following steps are used in a typical modal analysis in ANSYS:

Build the model.
Apply loads and obtain the solution.
Expand the modes.
Review the results.
In certain cases, especially if the model has of a large number of degrees of freedom, it is advantageous to define Master Degrees of Freedom (MDOF). This procedure condenses the full matrices describing the structure into a smaller size, thus reducing the computational cost significantly. The only boundary conditions that are permissible in modal analysis are zero displacements. Any constraints/loads that are non-zero are ignored in the analysis. Once the modal analysis is complete, the solution is expanded to find results related to the complete structure-not just the MDOF. The results include natural frequencies, mode shapes, and corresponding parametric (relative) stress fields.

### 8.5.1.1 Modal Analysis of a Bracket

The bracket shown in Fig. 8.29 is clamped at the two top holes. The reduced method of modal analysis is used here, which requires master degrees of freedom. Master degrees of freedom are automatically selected by ANSYS. The reduced modal analysis is then expanded for the number of modes desired (in this case, four). The goal is to obtain the modal frequencies and corresponding mode shapes.

## MODEL GENERATION

The finite element model of this bracket is created in a separate problem (Sec. 8.1.5.1). Therefore, the model will not be regenerated here; it will be generated interactively using an input file. It is worth noting that the components consisting of the nodes along the holes are defined in the input file.

- Create the model by reading the input file "bracketinp" from the CDROM location Vnput_Files\ch08Vbracket.inp using the following menu path:
- Browse for the bracket.inp file in the folder given above; click on $\boldsymbol{O K}$.
- Wait until the model is generated.


## SOLUTION

■ Specify the Analysis Type as Modal, using the following menu path (ANTYPE command), which will bring up the New Analysis dialog box:

Main Menu > Solution > Analysis Type > New Analysis

- Click on the Modal radio-button; click on $O K$.
- Print 4 reduced mode shapes using the following menu path:

Main Menu $>$ Solution > Analysis Type $>$ Analysis Options

- Click on the Reduced radio-button; click on $O K$, which will bring up another dialog box.
- Enter 4 for $P R M O D E$; click on $O K$.
- Specify the master degrees of freedom using the following menu path:

Main Menu $>$ Solution $>$ Master DOFs $>$ Program Selected

- In the dialog box, type 20 for NTOT Total no. of master DOF; click on the checkbox to show Yes for NRMDF Exclude rotational DOF; click on $O K$.

Expand the first four modes using the following menu path:
Main Menu > Solution > Load Step Opts > ExpansionPass > Single Expand > Expand Modes

- Enter 4 for $N M O D E$; click on $O K$.
- Constrain displacement and rotation degrees of freedom along the top-left and -right holes. For this purpose, first select the components created earlier for these holes ( $\boldsymbol{T L} \_\boldsymbol{B O L T}$ and $\boldsymbol{T} \boldsymbol{R}_{\_} \boldsymbol{B O L T}$ ) using the following menu path:

Utility Menu > Select > Comp/Assembly > Select Comp/Assembly

- A dialog box appears; click on the by component name radio-button; click on $O K$.
- A new dialog box with the components listed appears; highlight $T L \_B O L T$; click on $O K$. This action selects the nodes along the top-left hole.
- Specify the displacement boundary conditions using the following menu path:

[^17]- Pick Menu appears; click on Pick All.
- In the new dialog box, highlight All DOF; click on OK.
- Repeat the same procedure for the top-right hole (TR_BOLT).
- Select everything (ALLSEL command) using the following menu path: Utility Menu > Select > Everything
- Obtain the solution using the following menu path:

Main Menu > Solution > Solve > Current LS

- Close the /STATUS information window.
- Click on $O K$ to start the solution.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the results using the following menu path:

Main Menu > General Postproc > Results Summary

- The mode frequencies appear in a separate window, as shown in Fig. 8.78 .
- Plot the mode shapes.
- Set the solution to First Set using the following menu path:

Main Menu > General Postproc $>$ Read Results $>$ First Set

- Plot the mode shape using the following menu path:

Main Menu > General Postproc > Plot Results > Deformed Shape

- Click on Def + undef edge radio-button; click on OK.
- The mode shape is shown in Fig. 8.79 as it appears in the Graphics Window.
- Set the solution to Second Set using the following menu path:

Main Menu > General Postproc > Read Results > Next Set

- Plot the mode shape using the following menu path:

Main Menu > General Postproc > Plot Results > Deformed Shape

- Click on Def + undef edge radio-button; click on OK.
- The mode shape is shown in Fig. 8.79 as it appears in the Graphics Window.
- Repeat for the third and fourth modes.


Fig. 8.78 Listing of the mode frequencies.


Fig. 8.79 Mode shapes 1 through 4.

### 8.5.1.2 Vibration of an Automobile Suspension

An automobile suspension system is simplified to consider only two major motions of the system: (i) up-and-down linear motion of the body, and (ii) pitching angular motion of the body.

The body is idealized as a lumped mass with weight, $W$, and radius of gyration, $r$, as shown in Fig. 8.80. The equivalent finite element model is depicted in Fig. 8.81. The numerical values of the geometric parameters used in Fig. 8.80 are as follows: $l_{1}=4.5 \mathrm{ft}, l_{2}=5.5 \mathrm{ft}$, and $r=4 \mathrm{ft}$; the gravitational acceleration is $g=32.2 \mathrm{ft} / \mathrm{sec}^{2}$. The corresponding coupled frequencies, $f_{1}$ and $f_{2}$, are to be determined. The elastic modulus of the beam is $E=4 \times 10^{9} \mathrm{psf}$, the lumped weight is $W=3220 \mathrm{lb}$, the spring constants are $k_{1}=2400 \mathrm{lb} / \mathrm{ft}$ and $k_{2}=2600 \mathrm{lb} / \mathrm{ft}$. The nodal coordinates and element properties and connectivity are given in Tables 8.6 and 8.7, respectively.


Fig. 8.80 Schematic of the automobile suspension system.


Fig. 8.81 Finite element method model of the automobile suspension system.

Table 8.6 Node numbers and coordinates of the automobile suspension system model.

| Node Number | $x$ | $y$ |
| :---: | :---: | :---: |
| 1 | 0.0 | 0.0 |
| 2 | 0.0 | 1.0 |
| 3 | 4.5 | 1.0 |
| 4 | 10.0 | 1.0 |
| 5 | 10.0 | 0.0 |

Table 8.7 Finite element method model of the automobile suspension system.

| Element <br> Number | Attributes <br> Element <br> Type <br> Reference <br> Number |  |  | Real <br> Constant <br> Set <br> Number | Material <br> Reference <br> Number |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | 2 | 1 | 1 | 1 | 2 |
|  | 1 | 2 | 1 | 2 | 3 |
| 3 | 3 | 3 | 1 | 3 |  |
| 4 | 1 | 4 | 1 | 3 | 4 |
| 5 | 2 | 51 | 1 | 4 | 5 |

## model generation

Define the element types (ET command) using the following menu path, which brings up the Element Types dialog box:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Specify element type reference number 1 as BEAM3 (BEAM in the left list; 2D elastic 3 in the right list); click on $O K$.
- Click on $A d d$ for element type reference number 2.
- Specify element type reference number 2 as Combin14 (Combination in the left list, Spring-damper 14 in the right list); click on OK.
- Highlight COMBIN14 entry by clicking on it; click on Options.
- Select 2-D longitudinal for $\boldsymbol{K} 3$; click on $\boldsymbol{O K}$.
- Click on $\operatorname{Add}$ for element type reference number 3.
- Specify element type reference number 3 as MASS21 (Structural Mass in the left list, 3D mass 21 in the right list).
- Highlight MASS21 entry by clicking on it; click on Options.
- Select 2-D w rot inert for K3; click on $O K$.
- Click on Close.
- Specify the real constants ( R command) using the following menu path:


## Main Menu > Preprocessor > Real Constants > Add/EditDelete

- Click on Add .
- Specify the left spring constant $\left(k_{1}\right)$ as real constant set number 1 :
- Highlight Type 2 COMBIN14 entry by clicking on it; click on OK.
- Enter 2400 for $\boldsymbol{K}$; click on $O K$.
- Specify the beam geometric properties as real constant set number 2:
- Click on $\operatorname{Add}$.
- Highlight Type 1 BEAM3 entry by clicking on it; click on OK.
- Enter $\boldsymbol{I}$ for AREA, $\boldsymbol{1}$ for $\boldsymbol{I Z Z}$, and $\boldsymbol{I}$ for HEIGHT; click on OK.
- Specify the mass properties as real constant set number 3:
- Click on Add.
- Highlight Type 3 MASS21 entry by clicking on it; click on OK.
- Enter $\mathbf{1 0 0}$ for MASS and $\mathbf{1 6 0 0}$ for IZZ; click on OK.
- Specify the arbitrary beam properties as real constant set number 4:
- Click on Add.
- Highlight Type 1 BEAM3 entry by clicking on it; click on $\boldsymbol{O K}$.
- Enter $\mathbf{1}$ for AREA, 1 for IZZ, and $\mathbf{1}$ for HEIGHT; click on OK.
- Specify the right spring constant $\left(k_{2}\right)$ as real constant set number 5 :
- Highlight Type 2 COMBIN14 entry by clicking on it; click on OK.
- Enter 2600 for $K$; click on $O K$.
- Click on Close.
- Specify material properties for the beam (MP command) using the following menu path:


## Main Menu $>$ Preprocessor $>$ Material Props $>$ Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which brings up another dialog box.
- Enter $\mathbf{4 e} \boldsymbol{9}$ for $\boldsymbol{E X}$; click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit
$\square$ Create the nodes ( $\mathbf{N}$ command) using the following menu path, which brings up the Create Nodes in Active Coordinate System dialog box:

Main Menu > Preprocessor > Modeling > Create $>$ Nodes $>$ In Active CS

- A total of 5 nodes will be created.
- Referring to Table 8.6, enter the $\boldsymbol{x}$ - and $\boldsymbol{y}$-coordinates of node 1 ; click on Apply. This action will keep the Create Nodes in Active Coordinate System dialog box open. If the Node number field is left blank, then ANSYS will assign the lowest available node number to the node that is being created.
- Repeat the same procedure for the nodes 2,3 and 4.
- After creating node 5 , click on $O K$ (instead of Apply).

■ Create the elements (the element attributes and connectivity information are given in Table 8.7).

- Referring to Table 8.7 , element 1 is a spring element (type 2) with spring stiffness $2400 \mathrm{lb} / \mathrm{ft}$ (real constant set number 1), composed of
nodes 1 and 2. By default, the element attributes are set to 1 . Before element 1 is created, the element type attribute needs to be specified as 2. Perform this operation by using the following menu path, which will bring up the Element Attributes dialog box:
Main Menu > Preprocessor > Modeling > Create > Elements > Elem Attributes
- Select 2 COMBIN14 in the [TYPE] Element type number pull-down menu; click on $O K$. Any element that is created after this point will have these attributes (i.e., element type number 2, material number 1 , and real constant set number 1).
- Create element 1 (E command) using the following menu path, which brings up a Pick Menu:

Main Menu > Preprocessor > Modeling > Create > Elements > Auto Numbered > Thru Nodes

- Pick (by clicking in the Graphics Window) nodes 1 and 2; click on $\boldsymbol{O K}$ in the Pick Menu.
- Create elements 2 through 5 in the same manner, each time setting the correct attributes as explained above.
- The elements should appear in the Graphics Window, as shown in Fig. 8.82 .
- The commands equivalent to changing attributes are: TYPE, MAT, and REAL.
- Obtain and review the list of elements by using the following menu path:

Utility Menu > List > Elements > Nodes + Attributes
Observe attributes (element type, real constants, and material properties) and node numbers for each element (Fig. 8.83).

## SOLUTION

■ Specify the Analysis Type as Modal, using the following menu path (ANTYPE command), which will bring up the New Analysis dialog box:

Main Menu > Solution > Analysis Type $>$ New Analysis

- Click on the Modal radio-button; click on $\boldsymbol{O K}$.

■ Expand the first mode (MXPAND command) using the following menu path:

Main Menu > Solution > Load Step Opts > ExpansionPass > Single Expand > Expand Modes

- Enter 1 for $N M O D E$; click on $O K$.


Fig. 8.82 Elements of the automobile suspension system model.


Fig. 8.83 Listing of the elements.

- Print two reduced mode shapes (MODOPT command) using the following menu path:

Main Menu > Solution > Analysis Type > Analysis Options

- Click on the Reduced radio-button; click on $\boldsymbol{O K}$, which brings up another dialog box.
- Enter 2 for PRMODE; click on $\boldsymbol{O K}$.
- Specify the master degrees of freedom (M command) using the following menu path:
Main Menu >Solution > Master DOFs > User Selected > Define
- After picking node 3, click on $O K$ in the Pick Menu, which brings up the Define Master DOFs dialog box.
- Select $\boldsymbol{U Y}$ for $\boldsymbol{L a b 1}$ and $\boldsymbol{R O T Z}$ for $\boldsymbol{L a b 2 - 6}$; click on $\boldsymbol{O K}$.
- Specify boundary conditions (D command):
- Constrain $x$ - and $y$-displacements at nodes 1 and 5 using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick nodes 1 and 5; click on $\boldsymbol{O K}$ in the Pick Menu, which brings up the Apply U,ROT on Nodes dialog box.
- Highlight $\boldsymbol{U X}$ and $\boldsymbol{U Y}$; click on $\boldsymbol{O K}$.
- Constrain $x$-displacement at node 3 in the same manner.
- Instruct ANSYS to print the solution at every substep using the following menu path (OUTPR command):

Main Menu > Solution > Load Step Opts > Output Ctrls > Solu Printout

- Select Nodal DOF solu from the first pull-down menu.
- Click on the Every substep radio-button; click on $\boldsymbol{O K}$.
- Obtain the solution using the following menu path (SOLVE command):

Main Menu >Solution > Solve >Current LS

- Close the /STATUS information window.
- Click on $\boldsymbol{O K}$ to start the solution.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Retrieve the coupled frequencies and store them in user-defined parameters using the following menu path:


## Utility Menu > Parameters > Get Scalar Data

- Highlight Results data in the left list, Modal results in the right list; click on $\boldsymbol{O K}$, which brings up another dialog box.
- Enter freq1 for Name of parameter to be defined field, and enter 1 for Mode number $N$; click on $O K$.
- Repeat the procedure above to define parameter freq 2 corresponding to Mode 2.

List the parameters defined above using the following menu path:
Utility Menu > List > Other > Named Parameter

- Highlight freq1 or freq2 from the list of parameters; click on $O K$.
- Figure 8.84 shows the outcome of this action for freq1.


Fig. 8.84 First mode frequency for the automobile suspension system model.

Plot the first mode shape.

- Set the solution to First Set using the following menu path:

Main Menu > General Postproc > Read Results > First Set

- Plot the mode shape using the following menu path:

Main Menu > General Postproc > Plot Results > Deformed Shape

- Click on Def + undeformed radio-button; click on OK.
- The mode shape is shown in Fig. 8.85 as it appears in the Graphics Window.


### 8.5.2 Harmonic Analysis

When a structure is subjected to cyclic loading, the resulting response is expected to be cyclic as well. ANSYS provides the user with the capability to solve this class of problems through the Harmonic analysis option. The restrictions on a harmonic analysis are:

All loads must be sinusoidal functions of time.
All loads must have the same frequency.
The structure must exhibit linearly elastic behavior (no geometric and material nonlinearities).

Sinusoidal loads are specified through the parameters amplitude, phase angle, and forcing frequency range. Amplitude is the peak value of the load, and phase angle is the time lag between multiple loads that are out of phase with each other. On the complex plane, it is the angle measured from the real axis. Finally, forcing frequency range is the frequency range of the harmonic load (in cycles/time).

### 8.5.2.1 Harmonic Analysis of a Bracket

A bracket, shown in Fig. 8.29, experiences a harmonic (cyclic) loading at two points of application. The forcing frequency varies from 0 to 400 Hz ; however, the two loads are $120^{\circ}$ out of phase, i.e., $F_{1}=1.5 \sin \omega t$ and $F_{2}=1.5 \sin (\omega-120) t$. The amplitude of each load is 1.5 lb . The bracket is clamped at its two upper bolt holes. The structural damping ratio is 0.03 . The mode superposition method is used to calculate the harmonic response. This method requires a modal analysis to be performed first. A reduced modal analysis is used, so the harmonic analysis results in a reduced solution, which is then expanded using Expansion Pass. Note that the phase angle input is crucial for the expansion pass. Time History Postprocessing of the reduced analysis reveals the appropriate phase for expansion.


Fig. 8.85 First mode shape for the automobile suspension system model.

## MODEL GENERATION

The finite element model of this bracket was created in Sec. 8.1.5.1 (modeling of a bracket). Therefore, the model will not be regenerated here; it will be generated using the input file bracket.inp. It is worth noting that the components consisting of the nodes along the holes are defined in the input file.

- Specify the jobname as br_harm using the following menu path:

Utility Menu > File > Change Jobname

- In the dialog box, type br_harm in the [/FILNAM] Enter new jobname text field; click on the checkbox for New log and error files to show Yes; click on $\boldsymbol{O K}$.
- Create the model by reading the input file bracket.inp from the CD-ROM location Vnput_Files $\backslash c h 08 V$ bracket.inp using the following menu path:

Utility Menu > File > Read Input from

- Browse for the bracket.inp from the folder given above; click on $\boldsymbol{O K}$.
- Wait until the model is generated.


## SOLUTION

- Specify the Analysis Type as Modal, using the following menu path (ANTYPE command), which brings up the New Analysis dialog box:

Main Menu > Solution > Analysis Type > New Analysis

- Click on Modal radio-button; click on $\boldsymbol{O K}$.
- Specify the reduced modal analysis method using the following menu path:

Main Menu > Solution > Analysis Type > Analysis Options

- Click on the Reduced radio-button; click on $\mathbf{O K}$.
- A new dialog box appears; click on $\boldsymbol{O K}$.
- Constrain displacement and rotation degrees of freedom along the top-left and -right holes. For this purpose, first select the components created earlier for these holes ( $\boldsymbol{T L} \_\boldsymbol{B O L T}$ and $\boldsymbol{T} \boldsymbol{R}_{\_} \boldsymbol{B O L T}$ ) using the following menu path:

Utility Menu > Select > Comp/Assembly > Select Comp/Assembly

- A dialog box appears; click on the by component name radio-button; click on $\boldsymbol{O K}$.
- A new dialog box with the components listed appears; highlight $\boldsymbol{T L} \_\boldsymbol{B O L T}$; click on $\boldsymbol{O K}$. This action selects the nodes along the top-left hole.
- Specify the displacement boundary conditions using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; click on Pick All.
- In the new dialog box, highlight All DOF; click on $\boldsymbol{O K}$.
- Repeat the same procedure for the top-right hole (TR_BOLT).
- Specify the master degrees of freedom at the four nodes of interest (as shown in Fig. 8.86) and specify that an additional 40 degrees of freedom are to be selected automatically by ANSYS.
- Select the bottom corner nodes using the following menu path:

Utility Menu > Select > Entities


Fig. 8.86 Nodes at which master degrees of freedom are specified.

- Select Entities dialog box appears; choose Nodes from the first pulldown menu and choose By Location in the second pull-down menu; click on the radio-button for $Z$ coordinates and type 5 in the Min, Max text field; click on Apply.
- In the same dialog box (Select Entities dialog box), click on the radiobutton for $X$ coordinates and type -1.99, 1.99 in the Min, Max text field; click on the radio-button for Unselect; click on $O K$.
- Specify the master degrees of freedom using the following menu path:

Main Menu >Solution > Master DOFs > User Selected $>$ Define

- Click on Pick All in the Pick Menu, which brings up the Define Master DOFs dialog box.
- Select All DOF for LabI; click on $O K$.
- Select bottom inner nodes using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; choose Nodes from the first pulldown menu and choose By Location in the second pull-down menu; click on the radio-button for $X$ coordinates and type $-1.01,1.01$ in the Min, Max text field; click on Apply.
- In the same dialog box (Select Entities dialog box) click on the radiobutton for Z coordinates and type 3 in the Min, Max text field; click on the radio-button for Reselect; click on $\mathbf{O K}$ (the left node among the two is referred to as bottom inner left node in the postprocessing).
- Specify the master degrees of freedom using the following menu path:

Main Menu > Solution > Master DOFs > User Selected $>$ Define

- Click on Pick All in the Pick Menu, which brings up the Define Master DOFs dialog box.
- Select All DOF for Lab1; click on OK.
- Select everything using the following menu path:

Utility Menu > Select > Everything

- Specify additional 40 master degrees of freedom using the following menu path:

Main Menu $>$ Solution $>$ Master DOFs $>$ Program Selected

- In the dialog box, type 40 for NTOT Total no. of master DOF ; click on the checkbox to show Yes for NRMDF Exclude rotational DOF; click on $O K$.
- Save the model using the following menu path:

Utility Menu > File > Save as Jobname.db

- Obtain the modal solution using the following menu path:

Main Menu > Solution > Solve > Current LS

- Close the /STATUS information window.
- Click on $O K$ to start the solution.
- Wait until ANSYS responds with Solution is done!
- Exit the Solution Processor using the following menu path:

Main Menu > Finish

- Specify the Analysis Type as Harmonic, using the following menu path, which bring up the New Analysis dialog box:

Main Menu > Solution > Analysis Type > New Analysis

- Click on Harmonic radio-button; click on $O K$.
- Specify the Mode Superposition method using the following menu path:


## Main Menu > Solution > Analysis Type > Analysis Options

- Select Mode superpos'n from the Solution method pull-down menu; click on $O K$.
- Click on radio-button for Cluster at modes; click on OK.
- Apply force at bottom-left and -right corner nodes.
- Select bottom-left corner node using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; choose Nodes from the first pulldown menu and choose By Location in the second pull-down menu; click on the radio-button for $Z$ coordinates and type 5 in the Min, Max text field; click on the From Full radio-button; click on Apply.
- In the same dialog box (Select Entities dialog box) click on the radiobutton for $X$ coordinates and type -2 in the Min, Max text field; click on the Reselect radio-button; click on $O K$.
- Define scalar parameter using the following menu path:

Utility Menu > Parameters > Scalar Parameters

- In the text field within the dialog box, type $A M P L=1.5$; click on Accept. Close the Scalar Parameters dialog box by clicking on Close.
- Apply the force on previously selected node (bottom-left corner) using the following menu path:

[^18]- Click on Pick All in the Pick Menu.
- In the new dialog box, select $\boldsymbol{F Y}$ from the first pull-down menu and type AMPL in the VALUE Real part of force/mom text field; click on OK.
- Select bottom-right corner node using the following menu path:


## Utility Menu > Select > Entities

- Select Entities dialog box appears; choose Nodes from the first pulldown menu and choose By Location in the second pull-down menu; click on the radio-button for $Z$ coordinates and type 5 in the Min, Max text field; click on the From Full radio-button; click on Apply.
- In the same dialog box (Select Entities dialog box) click on the radiobutton for $X$ coordinates and type 2 in the Min, Max text field; click on the Reselect radio-button; click on $\mathbf{O K}$.
- Select degrees as the unit for angles using the following menu path:

Utility Menu > Parameters > Angular Units

- In the dialog box, select Degrees for [*AFUN] Units for angular; click on $\boldsymbol{O K}$.
- Define scalar parameter using the following menu path:

Utility Menu $>$ Parameters $>$ Scalar Parameters

- In the text field within the dialog box:
- Type PHASE=120; click on Accept.
- Type $\boldsymbol{F R}=A M P L^{*}$ COS(PHASE); click on Accept.
- Type FI=AMPL*SIN (PHASE); click on Accept.
- Click on Close.
- Apply the force on previously selected node (bottom-right corner) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural $>$ Force/Moment $>$ On Nodes

- Click on Pick All in the Pick Menu.
- In the new dialog box, select $\boldsymbol{F Y}$ from the pull-down menu and type $\boldsymbol{F R}$ in the VALUE Real part of force/mom text field and type FI in the VALUE2 Imag part of force/mom text field; click on OK.
- Select everything using the following menu path:

Utility Menu > Select > Everything

- Specify the harmonic frequencies using the following menu path:

Main Menu $>$ Solution $>$ Load Step Opts $>$ Time/Frequenc $>$ Freq \& Substps

- Type 0 in the first text field and 400 in the second text field.
- Type 40 for NSUBST text field; select the Stepped radio-button; click on $\boldsymbol{O K}$.
- Specify the damping using the following menu path:

Main Menu $>$ Solution $>$ Load Step Opts $>$ Time/Frequenc $>$ Damping

- Type 0.03 in the DMPRAT text field (third text field); click on $\boldsymbol{O K}$.
- Save the model using the following menu path:

Utility Menu > File > Save as Jobname.db

- Obtain the harmonic solution using the following menu path:

Main Menu >Solution >Solve >Current LS

- Close the /STATUS information window.
- Click on $O K$ to start the solution.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Enter the Time History Postprocessor using the following menu path:

Main Menu > TimeHist Postpro

- Time History Variables dialog box appears. Click on the button with the green plus sign at the top-left to define a variable.
- Add Time History Variable dialog box appears.
- Successively double-click on Nodal Solution, DOF Solution, and YComponent of displacement; in the text field, type UYRCNR; click on OK.
- Pick Menu appears; pick the bottom-right corner node; click on $O K$.
- Note the new variable UYRCNR in Time History Variables dialog box.
- Add new variable for displacement in $y$-direction at the bottom innerleft node by clicking on the button with the green plus sign and successively double-clicking on Nodal Solution, DOF Solution, and YComponent of displacement; in the text field, type UYINLEFT; click on OK.
- Pick the bottom inner-left node; click on $\boldsymbol{O K}$.
- Note the new variable UYINLEFT in Time History Variables dialog box.
- Highlight the rows UYRCNR and UYINLEFT from the list (by pressing $C t r l$ on the keyboard and clicking on the rows with left mouse button); click on the third from left button to plot the variation of these displacements.
- The plot appears in the Graphics Window, as shown in Fig. 8.87.
- Change the frequency axis ( $x$-axis) limits in order to take a closer look at the first peak using the following menu path:

Utility Menu > PlotCtrls > Style > Graphs > Modify Axes


Fig. 8.87 Amplitudes of the displacements at two locations as functions of frequency under harmonic loading.

- In the dialog box, under [/XRANGE], select the radio button for Specified range and type 16 and 37 in the XMIN, XMAX Specified $X$ range text fields; click on $O K$.
- Obtain the plot with the new range using the following menu path: Utility Menu > Plot > Replot
- The plot appears in the Graphics Window, as shown in Fig. 8.88.
- Reset the frequency axis ( $x$-axis) limits using the following menu path: Utility Menu > PlotCtrls > Style > Graphs > Modify Axes
- In the dialog box, under [/XRANGE], select the radio button for Auto calculated; click on $\boldsymbol{O K}$.
- In order to plot the phase angles, change the settings of the time history graphing using the following menu path:


## Main Menu > TimeHist Postpro > Settings > Graph

- Select Phase Angle from the pull-down menu for [PLCPLX] Complex variable; click on $\mathbf{O K}$.
- Click on the third from left button in Time History Variables dialog box to plot the variation of these phase angles.
- The plot appears in the Graphics Window, as shown in Fig. 8.89.


Fig. 8.88 Close-up view of amplitudes of the displacements at two locations as functions of frequency under harmonic loading.


Fig. 8.89 Phase angles of the displacements at two locations as functions of frequency under harmonic loading.

### 8.5.2.2 Harmonic Analysis of a Guitar String

A stainless-steel guitar string of length $l=710 \mathrm{~mm}$ and diameter $d=0.254$ mm is stretched between two rigid supports by a tensioning force $F_{1}=84 \mathrm{~N}$ applied at the right end (refer to Fig. 8.90). The string is struck at a location $c=165 \mathrm{~mm}$ from the left end with a force $F_{2}=1 \mathrm{~N}$. The elastic modulus and density of the string are $E=190 \mathrm{GPa}$ and $\rho=7920 \mathrm{~kg} / \mathrm{m}^{3}$, respectively. The fundamental frequency $f_{1}$ is determined using truss elements (LINK1), and a Mode Superposition Harmonic Response Analysis is performed. In order to perform the harmonic analysis, static and modal analyses are first performed.

## MODEL GENERATION

- Specify the jobname as guitar using the following menu path:

Utility Menu > File > Change Jobname

- In the dialog box, type guitar in the [/FILNAM] Enter new jobname text field; click on the checkbox for New log and error files to show Yes; click on $\boldsymbol{O K}$.

■ Define element types using the following menu path, which brings up the Element Types dialog box:

Main Menu $>$ Preprocessor $>$ Element Type $>$ Add/EditDelete

- Click on Add.
- Specify element type reference \# I as LINKI (select Link from the left list and 2D Spar 1 from the right list); click on $O K$.
- Click on Close .

■ Specify real constants using the following menu path:
Main Menu > Preprocessor > Real Constants > Add/Edit/Delete

- Click on Add.
- Specify the cross-sectional area as real constant set number 1:
- Highlight Type 1 LINK1 entry by clicking on it; click on OK.
- Enter 50671E-12 for Cross-sectional area AREA; click on OK.
- Click on Close .
- Specify material properties for the string using the following menu path:


Fig. 8.90 Finite element model of a guitar string.

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring up another dialog box.
- Enter $\mathbf{1 9 0 e 9}$ for $E X$; click on $O K$.
- In the Define Material Model Behavior dialog box, in the right window, double-click on Density, which will bring up another dialog box.
- Enter 7920 for DENS; click on OK.
- Close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit
m Create nodes using the following menu path, which brings up the Create Nodes in Active Coordinate System dialog box:

```
Main Menu \(>\) Preprocessor \(>\) Modeling \(>\) Create \(>\) Nodes \(>\) In Active CS
```

- Type 1 for NODE Node number; type 0 for $x$ - and $y$-coordinates; click on Apply.
- In the same dialog box, type 31 for NODE Node number; type 0.71 for $x$-coordinate and 0 for $y$-coordinate; click on $O K$.
- Fill between nodes using the following menu path:

Main Menu > Preprocessor > Modeling > Create $>$ Nodes $>$ Fill Between Nds

- Pick Menu appears; pick the two nodes; click on $\boldsymbol{O K}$.
- A new dialog box appears; click on $O K$.
- The nodes should appear in the Graphics Window, as shown in Fig. 8.91 .
- Create elements using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Elements > Auto Numbered > Thru Nodes

Fig. 8.91 Nodes of the guitar string finite element model.

- Pick Menu appears; pick the first two nodes (nodes 1 and 2); click on OK.
- Create the rest of the elements by copying element 1 using the following menu path:

Main Menu > Preprocessor > Modeling > Copy > Elements > Auto Numbered

- Pick Menu appears; pick the element; click on $\boldsymbol{O K}$.
- A new dialog box appears; type 30 for ITIME Total number of copies; click on $\mathbf{O K}$.
- The elements should appear in the Graphics Window, as shown in Fig. 8.92 .


## STATIC SOLUTION

- Specify the Analysis Type as Static, using the following menu path, which will bring up the New Analysis dialog box:

Main Menu > Solution > Analysis Type > New Analysis

- Click on Static radio-button; click on $\boldsymbol{O K}$.
- Specify boundary conditions.
- Constrain $x$ - and $y$-displacements at node 1 , and $y$-displacement at the remaining nodes using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick node 1; click on $\boldsymbol{O K}$.
- Highlight All DOF; click on Apply.
- Click on the radio-button for Box in the Pick Menu; select all the nodes except node 1 ; click on $\boldsymbol{O K}$.
- Highlight $\boldsymbol{U Y}$, remove the highlight on All DOF by clicking on it; click on $\boldsymbol{O K}$.
- Apply the axial force using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Force/Moment > On Nodes

- Pick the last node (node 31); click on $\boldsymbol{O K}$.
- In the new dialog box, select $\boldsymbol{F} \boldsymbol{X}$ from the pull-down menu and type 84 for VALUE Force/moment value; click on OK.

[^19]Fig. 8.92 Elements of the guitar string finite element model.

■ Specify this analysis to be a prestressed analysis using the following menu path:

Main Menu > Solution > Analysis Type > Sol'n Controls

- A dialog box with tabs appears. By default, the Basic tab is active.
- On the left side, under Analysis Options, put a checkmark for Calculate prestress effects.
- On the right side, under Write Items to Results File, click on the radiobutton for All solution items and select Write every substep from the Frequency pull-down menu.
- Click on OK.
- Save the model using the following menu path:

Utility Menu > File > Save as Jobname.db
The model will be saved in the Working Directory under the name guitar.db.

■ Obtain the solution using the following menu path:
Main Menu > Solution > Solve > Current LS

- Close the /STATUS information window.
- Click on $O K$ to start the solution.
- Wait until ANSYS responds with Solution is done!
- Exit the Solution Processor using the following menu path:

Main Menu $>$ Finish

## MODAL SOLUTION

- Specify the Analysis Type as Modal, using the following menu path, which will bring up the New Analysis dialog box:

Main Menu > Solution > Analysis Type > New Analysis

- Click on Modal radio-button; click on OK.
- Specify analysis options using the following menu path:

Main Menu > Solution > Analysis Type > Analysis Options

- Click on the Subspace radio-button; type 6 for No. of modes to extract (first text field).
- Place a checkmark for [PSTRES] Incl prestress effects; click on $O K$, which brings up another dialog box.
- Click on OK.
- Modify boundary conditions.
- Delete constraints in the $y$-direction at nodes 2 through 30 (leaving the displacement constraints at nodes 1 and 31 intact) using the following menu path:

Main Menu >Solution $>$ Define Loads $>$ Deiete $>$ Structural $>$ Displacement $>$ On Nodes

- Click on the radio-button for Box in the Pick Menu; select all the nodes except nodes 1 and 31; click on $\boldsymbol{O K}$.
- Select $\boldsymbol{U Y}$ from the pull-down menu; click on $\boldsymbol{O K}$.

Save the model using the following menu path:
Utility Menu > File > Save as Jobname.db
The model will be saved in the Working Directory under the name guitar.db.

- Obtain the solution using the following menu path:

Main Menu > Solution > Solve > Current LS

- Close the /STATUS information window.
- Click on $O K$ to start the solution.
- Wait until ANSYS responds with Solution is done!
- Exit the Solution Processor using the following menu path:

Main Menu > Finish

## hARMONIC SOLUTION

- Specify the Analysis Type as Harmonic, using the following menu path, which will bring up the New Analysis dialog box:

Main Menu > Solution > Analysis Type > New Analysis

- Click on Harmonic radio-button; click on OK.
- Specify the mode superposition method using the following menu path:


## Main Menu > Solution > Analysis Type > Analysis Options

- Select Mode superpos'n from the [HROPT] Solution method pulldown menu.
- Select Amplitud + phase from the [HROUT] DOF printout format pull-down menu; click on $O K$.
- Click on $O K$ in the new dialog box.
- Modify the boundary conditions.
- Delete the axial force using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Delete $>$ Structural $>$ Force/Moment $>$ On Nodes

- Pick the last node (node 31); click on $O K$.
- In the new dialog box, select $\boldsymbol{F X}$ from the pull-down menu; click on OK.
- Apply the vertical force using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Force/Moment > On Nodes

- Pick node 8 ; click on $\boldsymbol{O K}$.
- In the new dialog box, select $\boldsymbol{F Y}$ from the pull-down menu and type -1 for VALUE Real part of force/mom; click on $O K$.
- Specify the harmonic frequencies and enforce step loading using the following menu path:

Main Menu > Solution > Load Step Opts > Time/Frequenc $>$ Freq \& Substps

- Type 0 in the first text field and 2000 in the second text field for HARFRQ.
- Type 250 for $N S U B S T$ text field; select the Stepped radio-button; click on $O K$.
- Save the model using the following menu path:

Utility Menu > File > Save as Jobname.db
The model will be saved in the Working Directory under the name guitar.db.

- Obtain the solution using the following menu path:

Main Menu >Solution >Solve >Current LS

- Close the /STATUS information window.
- Click on $O K$ to start the solution.
- Wait until ANSYS responds with Solution is done!
- Exit the Solution Processor using the following menu path:

Main Menu $>$ Finish

## POSTPROCESSING

- Enter the Time History Postprocessor using the following menu path:
- Time History Variables dialog box appears. Click on the button with the green plus sign at the top-left to define a variable.
- Add Time History Variable dialog box appears.
- Successively double-click on the items Nodal Solution, DOF Solution, and $\boldsymbol{Y}$-Component of displacement; click on $O K$.
- Pick Menu appears; pick the midpoint node (node 16); click on $\boldsymbol{O K}$.
- Note the new variable UY_2 in Time History Variables dialog box.
- Highlight the row $\boldsymbol{U Y} Y_{-} 2$ from the list; click on the third from left button to plot the variation of these displacements.
- The plot appears in the Graphics Window, as shown in Fig. 8.93.


### 8.5.3 Transient Analysis

Practical engineering problems having a non-cyclic transient loading can be solved using the Transient analysis in ANSYS. Typically, the following steps are used for a Transient analysis:

Build the model.
Specify initial conditions.
Specify Solution Controls.
Apply loads.
Write Load Step File.
Apply/change loads and Solution Controls for the next Load Step(s) and write Load Step File(s).
Obtain solution from Load Step Files.
Review the Results.


Fig. 8.93 Amplitudes of the displacement at the midpoint node as a function of frequency under harmonic loading.

Solution Controls are specified using the Solution Controls dialog box, which is accessed through the following menu path:

Main Menu > Solution > Analysis Type >Sol'n Controls
There are five tabs in the Solution Controls dialog box. The first two tabs, Basic and Transient, are sufficient for most transient analyses. The Basic tab contains options to specify nonlinear geometry (large deformation), time at the end of the load step, load step and substep numbers and sizes, automatic time stepping, and the amount and frequency of results data to be saved. The Transient tab involves options related to time integration and damping specifications.

### 8.5.3.1 Dynamic Analysis of a Bracket

The bracket shown in Fig. 8.29 is clamped at the two top holes and subjected to an impact load at the bottom-left hole. The load is 12 lb , in the positive $z$-direction, and is applied for 0.01 sec . Three load steps with ramped loading are used; as shown in Fig. 8.94, the times at the end of the first, second, and third load steps are $0.005,0.01$, and 0.1 sec , respectively, with a time step size of $\mathrm{EE}-4 \mathrm{sec}$. The objective is to obtain the timedependent response of the bracket.

## MODEL GENERATION

The finite element model of this bracket was created in Sec. 8.1.5.1 (modeling of a bracket). Therefore, the model will not be regenerated here; it will be generated using an input file. It is worth noting that the components consisting of the nodes along the holes are defined in the input file.

- Create the model by reading the input file bracket.inp from the CD-ROM location Vnput_Files $\backslash c h 08 \backslash b r a c k e t . i n p ~ u s i n g ~ t h e ~ f o l l o w i n g ~ m e n u ~ p a t h: ~$

Utility Menu > File > Read Input from

- Browse for the bracket.inp from the folder given above; click on OK.
- Wait until the model is generated.


Fig. 8.94 Time-dependent loading on the bracket.

## SOLUTION

There are three distinct loading regiments: (1) from zero loading to maximum loading, (2) from maximum loading to complete removal of the load, and (3) time period after load removal during which the transient response of the bracket is sought. Each of these distinct periods will be defined as a Load Step. Each load step will be written to a load step file prior to solution. Once all the load step files are written, the solution will be obtained using these files. Displacement constraints do not change throughout the analysis; they will be specified only once at the beginning.

- Specify the Analysis Type as transient using the following menu path:

Main Menu > Solution > Analysis Type > New Analysis

- Click on Transient; click on OK.
- A new dialog box appears; click on $O K$.

■ Constrain displacement and rotation degrees of freedom along the top-left and -right holes. For this purpose, first select the components created earlier for these holes ( $\boldsymbol{T L} \__{-} B O L T$ and $\boldsymbol{T} R_{-} B O L T$ ) using the following menu path:

## Utility Menu > Select > Comp/Assembly > Select Comp/Assembly

- A dialog box appears; click on the by component name radio-button; click on $O K$.
- A new dialog box with the components listed appears; highlight $T L_{\_} B O L T$; click on $O K$. This action selects the nodes along the top-left hole.
- Specify the displacement boundary conditions using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Structural $>$ Displacement $>$ On Nodes

- Pick Menu appears; click on Pick All.
- In the new dialog box, highlight All DOF; click on $O K$.
- Repeat the same procedure for the top-right hole (TR_BOLT).

■ Apply the load at the bottom-left hole.

- For this purpose, create a local coordinate system at the center of the bottom-left hole using the following menu path:

Utility Menu > WorkPlane > Local Coordinate Systems > Create Local CS > At Specified Loc

- Pick Menu appears; type -1.5, -2, 4.5 in the text field in the Pick Menu; click on $\boldsymbol{O K}$.
- A dialog box appears; select Cylindrical 1 in the KCS Type of coordinate system pull-down menu and type -90 in the THYZ Rotation about local $X$ text field; click on OK.
- Select the keypoints along the bottom-left hole using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; choose Keypoints in first pull-down menu; choose By Location in the second pull-down menu; type $0.25 / 2$ in the Min, Max text field; click on $O K$.
- Apply forces on the keypoints using the menu path:

Main Menu >Solution > Define Loads $>$ Apply $>$ Structural $>$ Force/Moment $>$ On Keypoints

- Pick Menu appears; click on Pick All.
- In the new dialog box, select $\boldsymbol{F Z}$ from pull-down menu and enter $12 / 4$ for Force/moment value; click on OK.
- Select everything using the following menu path:

Utility Menu > Select > Everything

- Set solution options for the first load step using the following menu path:

Main Menu > Solution > Analysis Type >Sol'n Controls

- Solution Controls dialog box appeara. By default, the Basic tab is active.
- On the left side, under Time Control, type 0.005 for Time at end of loadstep; select On from Automatic time stepping pull-down menu.
- Click on the radio-button for Time increment and type 0.0001 in Time step size text field.
- On the right side, under Write Items, click on the radio-button for All solution items and select Write every substep from the Frequency pulldown menu.
- Activate the Transient tab (by clicking on it).
- Under Full Transient Options, click on the radio-button for Ramped loading; click on OK to close the Solution Controls dialog box.
- Write first load step file using the following menu path:

Main Menu $>$ Solution $>$ Load Step Opts $>$ Write LS File

- Type 1 ; click on $\boldsymbol{O K}$.
- The first load step is written to a file.
- Remove the load at the midpoint node using the following menu path:

[^20]
## - Click on Pick All.

- Select $\boldsymbol{F Z}$ from the pull-down menu; click on $\boldsymbol{O K}$.

Set solution options for the second load step using the following menu path:

Main Menu > Solution > Analysis Type $>$ Sol'n Controls

- Activate the Basic tab. On the left side, under Time Control, type 0.01 for Time at end of loadstep, leave the other options unchanged.
- Click on OK to close the Solution Controls dialog box.
- Write the second load step file using the following menu path:

Main Menu > Solution > Load Step Opts > Write LS File

- Type 2; click on $\boldsymbol{O K}$.
- The second load step is written to a file.
- Set solution options for the third load step using the following menu path: Main Menu > Solution > Analysis Type >Sol'n Controls
- On the left side, under Time Control, type 0.1 for Time at end of loadstep; leave the other options unchanged.
- Click on OK to close the Solution Controls dialog box.
- Write the third load step file using the following menu path:

Main Menu $>$ Solution $>$ Load Step Opts $>$ Write LS File

- Type $\mathbf{3}$; click on $\mathbf{O K}$.
- The third load step is written to a file.
- Obtain the solution using the following menu path:

Main Menu >Solution > Solve $>$ From LS Files

- Solve Load Step Files dialog box appears.
- Type 1 for LSMIN Starting LS file number and 3 for LSMAX Ending LS file number; click on OK.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review time-dependent behaviors of $y$-displacement at the bottom-left and -right corner nodes using the following menu path:

Main Menu > TimeHist Postpro

- Time History Variables dialog box appears.
- Click on the button with the green plus sign at the top-left to define a variable.
- Add Time History Variable dialog box appears.
- Successively double-click on the items Nodal Solution, DOF Solution, and $\boldsymbol{Y}$-Component of displacement; click on $\boldsymbol{O K}$.
- Pick Menu appears; pick the bottom-left corner node; click on $\boldsymbol{O K}$.
- Note the new variable UY_2 in Time History Variables dialog box.
- Add new variable for displacement in $y$-direction at the bottom-right corner node by clicking on the button with the green plus sign and successively double-clicking on the items Nodal Solution, DOF Solution, and $\boldsymbol{Y}$-Component of displacement; click on $\mathbf{O K}$.
- Pick the bottom-right corner node; click on $\boldsymbol{O K}$.
- Note the new variable UY_3 in Time History Variables dialog box.
- Highlight the rows $\boldsymbol{U} Y_{-} 2$ and $\boldsymbol{U} Y_{-} 3$ from the list (by pressing Ctrl on the keyboard and clicking on the rows with left mouse button); click on the third from left button to plot the time variation of these displacements.
- The plot appears in the Graphics Window, as shown in Fig. 8.95.
- Close Time History Variables dialog box.
( $\times 10=*-1$ )


Fig. 8.95 Time variation of displacements in the $y$-direction at two locations.

### 8.5.3.2 Impact Loading on a Beam

A 3-ft-long steel beam is clamped at both ends and subjected to an impact loading of 100 lbf that lasts 0.0002 sec at the midpoint (Fig. 8.96). It is assumed that the loading increases linearly from 0 lbf to 100 lbf in 0.0001 sec , and decreases in the same manner, as demonstrated in Fig. 8.97. The beam has a square cross-sectional area ( $1 \mathrm{in} \times 1 \mathrm{in}$ ) with material properties $E=29 \times 10^{6}$ psi and $v=0.32$. The objective is to solve for the timedependent response of the beam for 0.5 sec .

The problem is solved twice: (i) without structural damping, and (ii) with structural damping.

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Click on Add.
- Select Beam in the left list and 2D elastic 3 in the right list; click on OK.
- Click on Close.
- Specify the cross-sectional area, moment of inertia, and thickness using real constants ( R command) using the following menu path:
Main Menu > Preprocessor > Real Constants > Add/Edit/Delete
- Click on Add.
- Click on $\boldsymbol{O K}$; dialog box appears.
- Type 1 in the Cross-sectional area AREA text field.
- Type $1 / 12$ in the Area moment of inertia IZZ text field.
- Type 1 in the Total beam height HEIGHT text field.
- Click on $\boldsymbol{O K}$.
- Click on Close.


Fig. 8.96 Schematic of the beam under impact loading.


Fig. 8.97 Time-dependent loading.

- Specify material properties (MP command) using the following menu path:

Main Menu > Preprocessor > Material Props $>$ Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which will bring up another dialog box.
- Enter 29e6 for $\boldsymbol{E X}$ and 0.32 for $\boldsymbol{P R X Y}$; click on $\mathbf{O K}$.
- In the Define Material Model Behavior dialog box, in the right window, double-click on Density, which will bring up another dialog box.
- Enter 0.2836 for DENS; click on OK.
- Close the Define Material Model Behavior dialog box by using the following menu path:


## Material > Exit

- Create the solid model. In this problem, the solid model is a line, which is composed of two keypoints. Therefore, first create the keypoints, then the line.
- Create keypoints using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Keypoints $>\operatorname{In}$ Active CS

- In the Create Keypoints in Active Coordinate System dialog box, type -18 for $X$ and $\boldsymbol{O}$ for $\boldsymbol{Y}$; click on Apply (keypoint 1 is created).
- Type $\mathbf{1 8}$ for $\boldsymbol{X}$ and $\boldsymbol{O}$ for $\boldsymbol{Y}$; click on $\boldsymbol{O K}$ (keypoint 2 is created)
- Create the line using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Lines $>$ Lines $>$ Straight Line

- Pick keypoints 1 and 2; line is created; click on $\boldsymbol{O K}$ in the Pick Menu.
- Specify the number of elements on the line as $\mathbf{3 0}$ divisions using the following menu path:

Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize $>$ Lines $>$ Picked Lines

- Pick the line; click on $\boldsymbol{O K}$.
- Element Sizes on Lines dialog box appears; type 30 in the text field corresponding to NDIV (the second text field), and uncheck the first checkbox; click on Apply.
- Create the mesh using the following menu path:

Main Menu > Preprocessor > Meshing > Mesh > Lines

- Pick the line; click on $\boldsymbol{O K}$.


## SOLUTION WITHOUT STRUCTURAL DAMPING

There are three distinct loading regiments: (1) from zero loading to maximum loading, (2) from maximum loading to complete removal of the load, and (3) time period after load removal during which the transient response of the beam is sought. Each of these distinct periods will be defined as a Load Step. Each load step will be written to a load step file prior to solution. Once all the load step files are written, the solution will be obtained using these files. Displacement constraints do not change throughout the analysis; they will be specified only once at the beginning.

- Specify the Analysis Type as transient using the following menu path:

Main Menu > Solution > Analysis Type > New Analysis

- Click on Transient; click on OK.
- A new dialog box appears; click on $\boldsymbol{O K}$.
- Specify displacement constraints using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick the nodes on either end; click on $\boldsymbol{O K}$ in the Pick Menu.
- Highlight All DOF; click on OK.
- Apply the load at the midpoint node (located at the origin for picking conveniently) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Force/Moment > On Nodes

- Pick the midpoint node; click on $O K$.
- Select $F Y$ from the pull-down menu and type -100 in the VALUE Force/moment value text field; click on $\boldsymbol{O K}$.
- Set solution options for the first load step using the following menu path:

Main Menu >Solution > Analysis Type >Sol'n Controls

- Solution Controls dialog box appears. By default, the Basic tab is active.
- On the left side, under Time Control, type 1e-4 for Time at end of loadstep, type 20 for Number of substeps, and type 20 for Min no. of substeps.
- On the right side, under Write Items, click on the radio-button for All solution items and select Write every substep from the Frequency pulldown menu.
- Activate the Transient tab (by clicking on it).
- Under Full Transient Options, click on the radio-button for Ramped loading; click on $O K$ to close the Solution Controls dialog box.
- Write the first load step file using the following menu path:

Main Menu > Solution > Load Step Opts > Write LS File

- Type $\boldsymbol{1}$; click on $\boldsymbol{O K}$.
- The first load step is written to a file.
- Remove the load at the midpoint node using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Delete $>$ Structural $>$ Force/Moment $>$ On Nodes

- Pick the midpoint node; click on $O K$.
- Select $\boldsymbol{F Y}$ from the pull-down menu; click on $O K$.
- Set solution options for the second load step using the following menu path:
Main Menu > Solution > Analysis Type >Sol'n Controls
- Activate the Basic tab. On the left side, under Time Control, type $2 e-4$ for Time at end of loadstep; leave 20 unchanged for both Number of substeps and Min no. of substeps.
- On the right side, under Write Items, click on the radio-button for All solution items and select Write every substep from the Frequency pulldown menu.
- Click on $O K$ to close the Solution Controls dialog box.
- Write the second load step file using the following menu path:

Main Menu $>$ Solution $>$ Load Step Opts $>$ Write LS File

- Type 2 ; click on $O K$.
- The second load step is written to a file.
- Set solution options for the third load step using the following menu path:

Main Menu > Solution > Analysis Type > Sol'n Controls

- On the left side, under Time Control, type 0.5 for Time at end of loadstep and type 200 for both Number of substeps and Min no. of substeps.
- On the right side, under Write Items, click on the radio-button for All solution items and select Write every substep from the Frequency pulldown menu.
- Click on $O K$ to close the Solution Controls dialog box.
- Write the third load step file using the following menu path:

Main Menu > Solution > Load Step Opts > Write LS File

- Type 3; click on $O K$.
- The third load step is written to a file.
- Obtain the solution using the following menu path:

Main Menu > Solution > Solve > From LS Files

- Solve Load Step Files dialog box appears.
- Type 1 for LSMIN Starting LS file number and 3 for LSMAX Ending LS file number; click on $O K$.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

■ Review time-dependent behaviors of $y$-displacement at the midpoint, reaction force in $y$-direction at the left support, and reaction moment about $z$-axis at the left support using the following menu path:

## Main Menu > TimeHist Postpro

- Time History Variables dialog box appears.
- Click on the button with the green plus sign at the top-left to define a variable.
- Add Time History Variable dialog box appears.
- Successively double-click on the items Nodal Solution, DOF Solution, and Y-Component of displacement; click on $O K$.
- Pick Menu appears; pick the midpoint node; click on $O K$.
- Note the new variable UY_2 in Time History Variables dialog box.
- Highlight the row of $U Y_{\_} 2$ from the list; click on the third from left button to plot the time variation of this displacement.
- The plot appears in the Graphics Window, as shown in Fig. 8.98.


Fig. 8.98 Time variation of the $y$-displacement at the midpoint.

- Note that in this plot the displacement has a periodical behavior with no sign of decay. This is because no damping was specified in the solution. A solution with damping will be obtained in the next section.
- Click on the fourth from left button to list the time variation of this displacement in a separate window.


## SOLUTION WITH STRUCTURAL DAMPING

- Repeat the previous solution in its entirety with one difference: when specifying solution options, specify structural damping as 0.005 . This is accomplished using the following menu path:

Main Menu > Solution > Analysis Type > Sol'n Controls

- Activate the Transient tab in the Solution Controls dialog box.
- Under Damping Coefficients, type 0.005 for Stiffness matrix multiplier (BETA); click on OK to close the Solution Controls dialog box.
- Note that this needs to be done before each load step is written and that all the time and substep information needs to be re-entered.


## POSTPROCESSING

- Review the time-dependent behavior of the $y$-displacement at the midpoint in the same manner as was done for the case without damping. The corresponding response is given in Fig. 8.99 as it appears in the Graphics Window.


Fig. 8.99 Time variation of the $y$-displacement at the midpoint when damping is included in the analysis.

### 8.5.3.3 Dynamic Analysis of a 4-bar Linkage

Consider the 4-bar linkage shown in Fig. 8.100. The far-left node is rotated about the $z$-axis in a clockwise direction. The modeling part of the analysis utilizes direct nodal and elemental definition. Since this model is a flexible kinematics analysis, numerous beam elements are defined for each bar. To reduce the initial vibrations caused by loading, loads are applied in two time steps. A four-revolution displacement is imposed over a time period of 0.6 sec . This is equivalent to a $400 \mathrm{rev} / \mathrm{min}(\mathrm{rpm})$ displacement.

The following input listing is used for model generation and solution:

```
/PREP7
ET,1, BEAM4
ET,2,COMBIN7
R,1,0.063,5.25e-3,2.08373e-5,0.063,1 ! SPECIFY REAL
    ! CONSTANT SET 1
R,2,10e6,10e6,10e6 ! SPECIFY REAL CONSTANT SET 2
RMORE,1.5528e-4 ! ADDITIONAL DATA FOR
    ! REAL CONSTANT SET 2
MP,EX,1,10e6 ! ELASTIC MODULUS FOR MATERIAL 1
MP,PRXY,1,0.3 !POISSON'S RATIO FOR MATERIAL 1
MP,DENS,1,2.587799e-4 ! DENSITY FOR MATERIAL 1
MP,DAMP,1,0.000139 ! DAMPING MULTIPLIER FOR MATERIAL 
MP,EX,2,10e6 !ELASTIC MODULUS FOR MATERIAL 2
MP,PRXY,2,0.3 !POISSON'S RATIO FOR MATERIAL 2
MP,DENS,2,2.587799e-4 ! DENSITY FOR MATERIAL 2
MP,DAMP,2,0.000278 ! DAMPING MULTIPLIER FOR MATERIAL 2
```

| N, 1, 0, 0 | ! CREATE NODES |
| :---: | :---: |
| N, 2, 2.5,0 |  |
| N, 3, 2.5,0 |  |
| CLOCAL, 11,0,2.5,0,0,66.024 ! DEFINE LOCAL CS 11 |  |
| CSYS, 11 | : SWITCH TO LOCAL CS |
| N, 4, 2.5,0 | : CONTINUE CREATING NODES |
| N, 5, 5, 5, 0 |  |
| N, 6, 8, 5, 0 |  |
| N, 7, 11, 0 |  |
| N, 8, 11, 0 |  |
| CSYS,0 | ! SWITCH TO GLOBAL CARTESIAN CS |
| *AFUN, DEG | ! USE DEGREES FOR TRIG. FUNCTIONS |
| CLOCAL, 12, 0, 2. $5+11 * \operatorname{COS}(66.024), 11 * \operatorname{SIN}(66.024), 0,-73.21$ |  |
|  | ! DEFINE LOCAL |
|  | ! COORDINATE SYSTEM (CS) 12 |
| CSYS, 12 | ! SWITCH TO LOCAL CS 12 |
| N, 9, 2, 5, 0 | ! CONTINUE CREATING NODES |
| N, 10, 5. 25,0 |  |
| N, 11, 8, 0 |  |
| N,12,10.5,0 |  |
| E, 1, 2 | ! CREATE BEAM ELEMENTS FOR THE |
| E, 3, 4 | ! FIRST TWO BARS |
| E, 4, 5 |  |
| E, 5, 6 |  |
| E, 6, 7 |  |
| MAT, 2 | ! SWITCH TO MATERIAL 2 |
| E, 8,9 | ! CREATE BEAM ELEMENTS FOR THE |
| E, 9,10 | ! LAST BAR |
| E, 10, 11 |  |
| E, 11, 12 |  |
| REAL, 2 | ! SWITCH TO REAL CONSTANT SET 2 |
| TYPE, 2 | ! SWITCH TO ELEMENT TYPE 2 |
| E, 2, 3 | ! CREATE REVOLUTE JOINT ELEMENTS |
| E, 7, 8 |  |
| FINISH | ! FINISH |
| / SOL | ! ENTER SOLUTION PROCESSOR |
| *AFUN, RAD | ! USE RADIANS FOR TRIG. FUNCTIONS |
| ANTYPE, TRAN, NEW | ! DEFINE NEW ANALYSIS AS TRANSIENT |
| NLGEOM, 1 | ! INCLUDE LARGE DEFLECTION EFFECTS |
| D, 1, UX, , , 12, 11, UY | ! SUPPRESS X AND Y DISPLACEMENTS <br> ! AT NODES 1 AND 12 |
| D, ALL, UZ | ! SUPPRESS Z DISPL. AT ALL NODES |
| D, 1, ROTZ, -0.041888 | ! SPECIFY A SMALL ROTATION OF |
|  | ! -0.041888 RAD (2.4 DEG) AT NODE 1 |
| ACEL, 209.4, 324.7 | ! SPECIFY GRAVITATIONAL |
|  | ! ACCELERATION IN X AND Y |
|  | ! DIRECTIONS |
| TIME, 0.001 | ! SPECIFY THE TIME AT THE END OF |
|  | ! THIS LOAD STEP AS 0.001 SECONDS |
| NSUBST, 1 | ! USE ONLY ONE SUBSTEP |
| TIMINT, OFF | ! TURN OFF TRANSIENT EFFECTS |
| NEQIT, 1 | ! USE 1 EQUILIBRIUM ITERATION |
|  | ! FOR THE SUBSTEP |
| CNVTOL, F, 1,0.001 | ! SPECIFY CONVERGENCE TOLERANCE |
| CNVTOL, M, 1, 0.001 | ! FOR FORCE AND MOMENT |


| OUTRES, ALL, ALL | ! WRITE RESULTS FOR ALL SUBSTEPS ! TO THE DATABASE FILE |
| :---: | :---: |
| SOLVE | ! OBTAIN SOL'N FOR THIS LOAD STEP |
| D, 1, ROTZ, -25.13274 | ! SPECIFY A ROTATION OF -25.13274 |
|  | $!\mathrm{RAD}$ (1440 DEG) AT NODE 1 |
| TIME, 0.6 | ! SPECIFY THE TIME AT THE END OF |
|  | ! THIS LOAD STEP AS 0.6 SECONDS |
| NSUBST, 99 | ! USE 99 SUBSTEPS |
| TIMINT, ON | ! INCLUDE TRANSIENT EFFECTS |
| NEQIT, 30 | ! USE A MAXIMUM OF 30 EQUILIBRIUM |
|  | ! ITERATIONS FOR EACH SUBSTEP |
| KBC , 0 | ! APPLY THE LOAD (ROTATION) IN A |
|  | ! RAMPED FASHION |
| SOLVE | ! OBTAIN SOL'N FOR THIS LOAD STEP |

## POSTPROCESSING

- Review time-dependent behavior of the torque at node 1 using the following menu path:


## Main Menu > TimeHist Postpro

- Time History Variables dialog box appears.
- Click on the button with the green plus sign at the top-left to define a variable.
- Add Time History Variable dialog box appears.
- Successively double-click on Reaction Forces, Structural Moments, and Z-Component of moment; click on OK.
- Pick Menu appears; pick node 1; click on $O K$.
- Note the new variable $M Z \_2$ in the Time History Variables dialog box.
- Highlight the row $M Z \_2$ from the list; click on the third from left button to plot the time variation of this displacement.
- The plot appears in the Graphics Window, as shown in Fig. 8.101.
- Plot the nodes using the following menu path:

Utility Menu > Plot > Nodes

- Add new variable for displacement in the $y$-direction at node 2 by clicking on the button with the green plus sign and successively doubleclicking on Nodal Solution, DOF Solution, and Y-Component of displacement; click on OK.
- Pick Menu appears; pick node 2 . Since nodes 2 and 3 are coincident, ANSYS asks the user which one to pick. Select node 2, click on $\boldsymbol{O K}$. Click on $O K$ in the Pick Menu.
- Note the new variable UY_3 in the Time History Variables dialog box.
- Highlight the row $U Y \_3$ from the list; click on the third from left button to plot the time variation of this displacement.
- The plot appears in the Graphics Window, as shown in Fig. 8.102.
- Close the Time History Variables dialog box.


Fig. 8.100 Schematic (top), and finite element model (bottom) of the 4-bar linkage.


Fig. 8.101 Time variation of the moment at the left joint (node 1).


Fig. 8.102 Time variation of the $y$-displacement at node 2.

- Obtain an animation of the motion of the structure.
- Enter General Postprocessor and read the results using the following menu path:

Main Menu > General Postproc > Read Results > Last Set

- Plot deformed shape using the following menu path:

Main Menu > General Postproc > Plot Results > Deformed Shape

- Select Def + undeformed; click on OK.
- Obtain animation using the following menu path:

Utility Menu > PlotCtris > Animate > Over Time

- Animate Over Time dialog box appears. Enter 100 for Number of animation frames. Enter 0.05 for Animation time delay (sec); click on OK.


## Chapter 9

## LINEAR ANALYSIS OF FIELD PROBLEMS

The field equation,

$$
\begin{equation*}
D_{x} \frac{\partial^{2} \phi}{\partial x^{2}}+D_{y} \frac{\partial^{2} \phi}{\partial y^{2}}-A \phi+B=0 \tag{9.1}
\end{equation*}
$$

(where $\phi$ is the field variable) governs a wide variety of physical problems, referred to as field problems. Several engineering problems, e.g., torsion of noncircular section, ideal irrotational fluid flow, seepage, heat transfer, and electrostatic, are embedded in this equation. For each of these problems, the parameters $D_{x}, D_{y}, A$, and $B$ designate a different physical property. For example, if the problem under consideration involves two-dimensional steady-state heat transfer with no heat generation within the body, then Eq. (9.1) becomes

$$
\begin{equation*}
D_{x} \frac{\partial^{2} T}{\partial x^{2}}+D_{y} \frac{\partial^{2} T}{\partial y^{2}}=0 \tag{9.2}
\end{equation*}
$$

where $D_{x}$ and $D_{y}$ are the thermal conductivity of the material in the $x$ - and $y$-directions, respectively, and $T$ represents temperature.

Most of the field problems can be solved using ANSYS software. However, only two types of field problems are considered within the context of this book: (i) heat transfer problems and (ii) moisture diffusion problems.

### 9.1 Heat Transfer Problems

In certain cases, a thermal analysis is followed by a stress analysis in order to evaluate the structural integrity of the component under the given thermal conditions. In a typical heat transfer problem, the goal is to obtain certain thermal quantities within a body under a specific set of boundary conditions. These quantities include: temperatures, thermal fluxes and gradients, and the amount of heat dissipated. There are two main types of thermal analyses:

Steady-state heat transfer: Solution is time independent.

Transient heat transfer: Subjected to specific initial conditions, the solution exhibits a time-dependent behavior. If the transient solution is obtained for a sufficiently long time period, the solution is expected to converge to the steady-state solution.

ANSYS accommodates three main heat transfer types: conduction, convection, and radiation. Convection boundary conditions require knowledge of the film coefficient and ambient temperature. There are a few different ways to impose radiation conditions within ANSYS. However, only one of them, the Radiosity Solver method, is explained within the context of this book.

Several boundary conditions are available, including specification of temperatures, heat fluxes, convective heat loss/gain, and radiation. As for the body loads, the user can specify heat generation rates within the domain.

In both steady-state and transient analyses, the material properties may be defined as temperature dependent. In the presence of temperature-dependent material properties, the analysis becomes nonlinear, thus requiring an iterative solution.

The most commonly used heat transfer elements are plane55 (twodimensional plane) and SOLID70 (three-dimensional brick).

### 9.1.1 Steady-state Analysis

When the boundary conditions and body loads do not vary with time and there are no specified initial conditions, the solution quantities do not vary with time. In such cases, steady-state solutions are obtained. A steady-state analysis is demonstrated by considering two problems.

### 9.1.1.1 Analysis of a Tank/Pipe Assembly

A cylindrical tank and a small pipe form a junction, as shown in Fig. 9.1 (only $1 / 8^{\text {th }}$ of the geometry is shown due to octant-symmetry). Inside the tank, there is fluid at a temperature of $450^{\circ} \mathrm{F}$. A steady flow of a fluid at a temperature of $100^{\circ} \mathrm{F}$ is experienced inside the pipe. The film coefficient along the inner surface of the tank is $250 \mathrm{Btu} / \mathrm{hr}^{-} \mathrm{ft}^{2}-{ }^{\circ} \mathrm{F}$ whereas the film coefficient along the inner surface of the pipe depends on the surface temperature. The geometric parameters and boundary conditions are given in Table 9.1, and the material properties are summarized in Table 9.2. Note in Tables 9.1 and 9.2 that the length units are in inches and feet. However, in order to obtain a physically correct solution the units must be consistent. Inches are used in this problem, therefore any parameter with the length unit in feet must be converted to inches. The goal is to determine the temperature distribution in the tank.


Fig. 9.1 The geometry of the tank and the pipe from different viewpoints.

Table 9.1 Geometric parameters and boundary conditions used in the analysis.

| Parameter | Value |
| :---: | :---: |
| Inside diameter, pipe | 0.8 in |
| Outside diameter, pipe | 1.0 in |
| Inside diameter, tank | 2.6 in |
| Outside diameter, tank | 3.0 in |
| Inside bulk fluid temperature, tank | $450^{\circ} \mathrm{F}$ |
| Inside film coefficient, tank | $250 \mathrm{Btu} / \mathrm{hr}-\mathrm{ft}^{2}-{ }^{\circ} \mathrm{F}$ |
| Inside bulk fluid temperature, pipe | $100^{\circ} \mathrm{F}$ |

Table 9.2 Material properties used in the analysis.

| Temperature <br> $\left({ }^{\circ} \mathrm{F}\right)$ | Density <br> $\left(\mathrm{lb} / \mathrm{in}^{3}\right)$ | Conductivity <br> $(\mathrm{Btu} / \mathrm{hr}-\mathrm{ft}-$ <br> $\left.{ }^{\circ} \mathrm{F}\right)$ | Specific <br> Heat <br> $\left(\mathrm{Btu} / \mathrm{lb}-{ }^{\circ} \mathrm{F}\right)$ | Film Coefficient <br> $(\mathrm{pipe})$ <br> $\left(\mathrm{Btu} / \mathrm{hr}-\mathrm{ft}^{2}-{ }^{\circ} \mathrm{F}\right)$ |
| :---: | :---: | :---: | :---: | :---: |
| 70 | 0.285 | 8.35 | 0113 | 426 |
| 200 | 0.285 | 8.90 | 0.117 | 405 |
| 300 | 0.285 | 9.35 | 0.119 | 352 |
| 400 | 0.285 | 9.80 | 0.122 | 275 |
| 500 | 0.285 | 10.35 | 0.125 | 221 |

## MODEL GENERATION

- Declare Preferences for Graphics User Interface (GUI) filtering (Fig. 9.2) using the following menu path:


## Main Menu > Preferences

- Place a checkmark corresponding to Thermal; click on $\boldsymbol{O K}$.
- This action eliminates several menu entries that are not related to thermal analyses, thus resulting in an abridged menu.
- Define the element type (ET command) using the following menu path:


## Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Element Types dialog box appears; click on Add.
- Select Thermal Solid in the left list and Brick 8 Node 70 on the right list; click on OK.
- Click on Close.
- Specify the temperature-independent material property, density (MP command), using the following menu path:

Main Menu > Preprocessor $>$ Material Props $>$ Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Thermal and Density, which will bring up another dialog box.
- Enter 0.285 for DENS; click on OK.
- Specify temperature-dependent material properties.
- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Thermal, Conductivity, and Isotropic, which will bring up another dialog box.


Fig. 9.2 Preferences for GUI Filtering dialog box.

- Click on the Add Temperature button four times (so that there are five temperature slots) (Fig. 9.3).
- Enter temperature values (i.e., 70, ..., 500) in the top row, from left to right in ascending order (Table 9.2).
- Enter the corresponding thermal conductivity values referring to Table 9.2. However, the values given in Table 9.2 are in units of Btu/hr-ft- ${ }^{\circ} \mathrm{F}$, which must be converted to $\mathrm{Btu} / \mathrm{hr}-\mathrm{in}-{ }^{\circ} \mathrm{F}$. This can be achieved within the dialog box, as shown in Fig. 9.4. Therefore, enter 8.35/12, ..., 10.35/12.
- View the variation of thermal conductivity as a function of temperature by clicking on the Graph button. $\boldsymbol{K X X}$ versus TEMP appears in the Graphics Window (Fig. 9.5).
- Click on $O K$.
- Specify specific heat in the same manner as thermal conductivity.
- Specify the temperature-dependent film coefficient inside the pipe.
- Although the tank and the pipe are of the same material, their film coefficients are different. Therefore, the film coefficient inside the pipe will be defined as a new material.
- In the Define Material Model Behavior dialog box, define a new material reference number using the following menu path:

[^21]

Fig. 9.3 Adding temperatures for tempera-ture-dependent material properties.


Fig. 9.4 Conductivity values at five different temperatures.


Fig. 9.5 Variation of conductivity with temperature as plotted in ANSYS.

- A dialog box appears with the material reference number set to the next available number (in this case 2) (Fig. 9.6); click on $\boldsymbol{O K}$.
- Note the new entry for Material Model Number 2 in the left list in the Define Material Model Behavior dialog box.
- In the right list, double-click on Convection or Film Coef., which brings up a new dialog box.
- Click on the Add Temperature button four times (so that there are five temperature slots); enter temperature values (i.e., 70, ... , 500) in the top row, from left to right in ascending order.
- Enter the film coefficient values (division by 144 is required for unit conversion), as shown in Fig. 9.7; click on OK.
- Close the Define Material Model Behavior dialog box.
- The film coefficient for the inside surface of the tank will be specified later during the application of boundary conditions.
- Define parameters.
- In order to demonstrate the use of parameters in ANSYS the following parameters and their values are defined:

```
ril = 1.3
rol=1.5
zI=2.0
ri2 = 0.4
ro2 =0.5
z2=2.0
```

- Parameters can be defined in two different ways:
- Using the Input Field, write the user-defined parameter name followed by the equal sign $(=)$ and the value assigned to the parameter (Fig. 9.8). Finish by the hitting Enter key on the keyboard.
- Define the parameters using the following the menu path:


## Utility Menu > Parameters > Scalar Parameters

The Scalar Parameters dialog box appears with an input box toward the bottom (Fig. 9.9). Write the user-defined parameter name followed by the equal sign (=) and the value assigned to the parameter; click on Accept. Observe that the newly defined parameter appears in the list.

- Define all the parameters using either method.
- Create the volume for the tank-a hollow partial cylinder-(CYLIND command) using the following menu path:


Fig. 9.6 Definition of a new material.


Fig. 9.7 Temperature-dependent film coefficient.


Fig. 9.8 Defining a parameter through the Input Field.


Fig. 9.9 Scalar Parameters dialog box.

- Enter ro1 for RAD1 Outer radius, ri1 for RAD2 Optional inner radius, $z 1$ for the second text box in the Z1, Z2 Z-coordinates row, and 90 for THETA2 Ending angle (degrees); click on OK.
- Observe the hollow partial cylinder created in the Graphics Window.
- View the tank from different angles using the following menu path: Utility Menu > PlotCtrls > Pan Zoom Rotate
- Click on Obliq.
- Figure 9.10 shows the outcome of this action as it appears in the Graphics Window.
- Create the volume for the pipe-a second hollow partial cylinder.
- In order to use the GUI for this action, first the Working Plane needs to be rotated using the following menu path:

Utility Menu > WorkPlane > Offset WP by Increments

- Offset WP dialog box appears.
- Find the second input field with the title: $X Y, Y Z, Z X$ Angles.
- Enter $\mathbf{0}, \boldsymbol{- 9 0}, \mathbf{0}$ in the second input field; click on $\boldsymbol{O K}$ (Fig. 9.11).
- Create the second hollow partial cylinder using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Volumes > Cylinder > By Dimensions

- In the same manner used for the creation of the cylinder for the tank, enter ro2 for RAD1 Outer radius, ri2 for RAD2 Optional inner radius, $z 2$ for the second text box in the Z1, Z2 Z-coordinates row, and -90 (not 90) for THETA2 Ending angle (degrees); click on OK.


Fig. 9.10 Hollow partial cylinder as it appears in the Graphics Window.


Fig. 9.11 Offset WP dialog box.

- Observe the second hollow partial cylinder created in the Graphics Window (Fig. 9.12).
- Turn the numbering on for the volumes using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Place a checkmark by clicking on the square box next to VOLU Volume numbers; click on OK.
- In the Pan Zoom Rotate Window, click on Left button.
- Figure 9.13 shows the outcome of this action as it appears in the Graphics Window; observe the different colors ${ }^{1}$ for each volume and the volume numbers.
- Reset the Working Plane to its original configuration by using the following menu path:

Utility Menu > WorkPlane > Offset WP by Increments

- Offset WP dialog box appears.
- Enter 0, 90,0 in the second input field; click on $\boldsymbol{O K}$.
- Use Boolean Operator Overlap to define the intersection volume between the two partial cylinders using the following menu path:

Main Menu > Preprocessor > Modeling > Operate > Booleans > Overlap > Volumes

- Pick Menu appears; pick the two volumes (or click on Pick All in the Pick Menu); click on OK.
- In the Pan Zoom Rotate Window, click on Left button.
- Figure 9.14 shows the outcome of this action as it appears in the Graphics Window.
- Delete the excess volumes using the following menu path:


## Main Menu > Preprocessor > Modeling > Delete > Volume and Below

- Pick Menu appears; pick the volumes to be deleted (carefully!); the volumes that need to be deleted are \#3 and \# 4.
- While picking the volumes, observe the number next to Volu No. in the Pick Menu change as the volumes are picked. This number corresponds to the last volume that is picked. If the wrong volume is picked, rightclick inside the Graphics Window and observe that the mouse pointer changes from upward arrow (pick) to downward arrow (unpick); then unpick the wrong volume and pick the correct one.

[^22]

Fig. 9.12 Two hollow partial cylinders as they appear in the Graphics Window.


Fig. 9.13 Two hollow partial cylinders with volume numbers turned on (both colors and numbers).


Fig. 9.14 Volumes after overlapping.

- Once the volumes to be deleted (3 and 4) are picked, click on $O K$ to finalize the deletion.
- Figure 9.15 shows the left view of the volumes after deletion as they appear in the Graphics Window.
- Set the element shape to be used as hexahedron (brick shaped) using the following menu path:


## Main Menu > Preprocessor > Meshing > Mesher Opts

- In the Mesher Options dialog box, click on the Mapped radio-button (Fig. 9.16); click on $\boldsymbol{O K}$.
- Set Element Shape dialog box appears (Fig. 9.17), click on OK.
- In 3-D, mapped meshing can only be performed on volumes with 4 to 6 sides. Examination of the volumes in the current model reveals that volume 7 has 7 sides. In order to be able to perform mapped meshing on this volume, certain areas ( $2-5$ pair) and lines ( $7-12,5-10$ pairs) must be concatenated.
- In order to be able to pick the areas and lines with ease, they should be in plain view. Rotate the model using rotation buttons on the Pan Zoom Rotate Window as follows: click on the negative $X$-rotation arrow button twice and negative $Y$-rotation arrow button once.


Fig. 9.15 Volumes after deletion.


Fig. 9.16 Mesher Options dialog box.


Fig. 9.17 Set Element Shape dialog box.

- Figure 9.18 shows the outcome of this action as it appears in the Graphics Window.
- Turn off volume numbers and turn on area numbers using the following menu path:

Utility Menu > PlotCtris > Numbering

- Remove the checkmark by clicking on the square box next to $\boldsymbol{V O L U}$ Volume numbers (turning volume numbers off); place a checkmark on the square box next to AREA Area numbers (turning area numbers on); click on $\boldsymbol{O K}$.
- Plot areas using the following menu path:


## Utility Menu > Plot > Areas

- Identify areas 2 and 5 (as shown in Fig. 9.19).
- Concatenate areas 2 and 5 using the following menu path:


## Main Menu > Preprocessor > Meshing > Concatenate > Areas

- Pick areas 2 and 5; click on $\boldsymbol{O K}$.
- Turn off area numbers and turn on line numbers.
- Plot lines.
- Identify lines 5, 7, 10, and 12 (as shown in Fig. 9.20).
- Concatenate lines 5 and 10 using the following menu path:

Main Menu > Preprocessor $>$ Meshing $>$ Concatenate $>$ Lines

- Pick lines 5 and 10; click on $\boldsymbol{O K}$.
- Similarly, concatenate lines 7 and 12.
- Specify the number of elements on selected lines for mapped meshing. On lines 18 and 47, use 2 divisions; on line 7, use 6 divisions; on line 12, use 5 divisions; on line 39 , use 11 divisions. For this action, use the following menu path:


Fig. 9.18 Volumes viewed after rotations.


Fig. 9.19 Area plot with area numbers (colors and numbers) turned on.


Fig. 9.20 Line plot with line numbers (colors and numbers) turned on.

- In the text field of the Pick Menu, type 18 followed by hitting enter on the keyboard; type 47 and hit enter again.
- Click on Apply.
- Element Sizes on Lines dialog box appears; type 2 in the text field corresponding to NDIV (the second text field); uncheck the first check box; click on Apply.
- Repeat this procedure for lines 7, 12, and 39 with their corresponding number of divisions as $\mathbf{6}, \mathbf{5}$, and 11, respectively. After specifying number of divisions for line 39, click on $\boldsymbol{O K}$ in the Element Sizes on Lines dialog box.
- Create the mesh using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Volumes $>$ Mapped $>4$ to 6 sided

- In the Pick Menu, click on Pick All.
- Observe the mesh using the different options in the Pan Zoom Rotate Window.
- Figure 9.21 shows the oblique view of the mesh.


## SOLUTION

- Apply uniform temperature to the entire model using the following menu path:


## Main Menu > Solution > Define Loads > Apply > Thermal > Temperature > Uniform Temp

- Dialog box appears; type 450; click on $O K$.


Fig. 9.21 Oblique view of the mesh.

- Apply boundary conditions.
- Change the active coordinate system to the global cylindrical coordinate system using the following menu path:

```
Utility Menu > WorkPlane > Change Active CS to > Global Cylindrical
```

- Select nodes at the inner surface of the tank for convective boundary conditions using the following menu path:
Utility Menu > Select > Entities
- Select Entities dialog box appears.
- Choose Nodes from the first pull-down menu.
- Chose By Location from the second pull-down menu.
- Click on the radio-button $X$ coordinates. In cylindrical coordinates, $x$ and $y$ correspond to $r$ and $\theta$; therefore, the selection criterion will utilize the $r$-coordinate as long as the cylindrical coordinate system is active.
- Type ril in the text field; click on $\mathbf{O K}$ (Fig. 9.22).
- Activate the ANSYS Output Window and examine the lines on the bottom to confirm that several nodes are selected as a result of this action (Fig. 9.23).
- Apply convective boundary conditions on the selected nodes using the following menu path:

[^23]

Fig. 9.22 Select Entities dialog box.


Fig. 9.23 Selection of nodes reported in the Output Window.

- Click on Pick All.
- In the dialog box, type $250 / 144$ for film coefficient (first text field) and 450 for bulk temperature (second text field); click on $O K$ (Fig. 9.24).
- Select nodes at the far edge of the tank for the temperature boundary conditions using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears.
- Choose Nodes from the first pull-down menu.
- Chose By Location from the second pull-down menu.
- Click on the radio-button $Z$ coordinates.
- Type $z 1$ in the text field; click on $O K$.
- Activate the ANSYS Output Window and examine the lines on the bottom to confirm that several nodes are selected as a result of this action.


Fig. 9.24 Apply CONV on nodes dialog box.

- Apply temperature boundary conditions on the selected nodes using the following menu path:

Main Menu > Solution > Define Loads > Apply > Thermal > Temperature > On Nodes

- Click on Pick All.
- In the dialog box, highlight TEMP and type 450 for VALUE Load Temp value; click on $O K$.
- Rotate the working plane $-90^{\circ}$ (negative $90^{\circ}$ ) about the $x$-axis using the following menu path:
Utility Menu > WorkPlane > Offset WP by Increments
- Offset WP dialog box appears.
- Type $\mathbf{0 , ~} \mathbf{9 0 , 0}$ in the $\boldsymbol{X Y}, \boldsymbol{Y Z}, \mathbf{Z X}$ Angles text field (second text field); click on $O K$.
- Define a local cylindrical coordinate system at the working plane origin using the following menu path:

Utility Menu > WorkPlane > Local Coordinate Systems > Create Local CS > At WP Origin

- Specify 11 for the KCN Ref number of new coord sys field (number 11 appears by default-leave it unchanged).
- Choose Cylindrical 1 from the pull-down menu; click on $\boldsymbol{O K}$.
- Select nodes at the inner surface of the pipe for the convective boundary conditions using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears.
- Choose Nodes from the first pull-down menu.
- Choose By Location from the second pull-down menu.
- Click on the radio-button $X$ coordinates.
- Type ri2 in the text field; click on $O K$.
- Apply the convective boundary condition with a temperature-dependent film coefficient on the selected nodes using the following menu path:

Main Menu > Solution > Define Loads > Apply > Thermal > Convection > On Nodes

- Click on Pick All.
- In the dialog box, type -2 (negative 2) for film coefficient and $\mathbf{1 0 0}$ for bulk temperature; click on $\boldsymbol{O K}$.
- Select everything using the following menu path:

Utility Menu > Select > Everything

- Obtain the solution.
- Although steady-state (no time dependence), this problem is nonlinear due to the existence of the temperature-dependent film coefficient and thus requires an iterative solution. Therefore, time is introduced as an auxiliary variable, and related parameters are specified.
- Specify automatic time stepping using the following menu path:

Main Menu > Solution > Load Step Opts > Time/Frequenc > Time and Substps

- Type 50 in the second text field corresponding to NSUBST (Fig. 9.25).
- Click on the $O N$ radio-button for Automatic time stepping; click on OK.
- Save using the following menu path:

Utility Menu > File > Save as

- Start the solution using the following menu path:

Main Menu >Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window and click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

meview temperature contours using the following menu path:
Main Menu > General Postproc > Plot Results > Contour Plot > Nodal Solu

- Select DOF solution from the left list and Temperature TEMP from the right list; click on $O K$.
- The temperature contour plot is shown in Fig. 9.26 as it appears in the Graphics Window.


Fig. 9.25 Time and Substep Options dialog box.


Fig. 9.26 Contour plot of temperatures.

- Review thermal flux vectors using the following menu path:

Main Menu > General Postproc > Plot Results $>$ Vector Plot $>$ Predefined

- Select Flux and gradient from the left list and Thermal flux TF from the right list; click on $\boldsymbol{O K}$.
- Figures 9.27 and 9.28 show the flux vectors as they appear in the Graphics Window, viewed from top and left, respectively.


### 9.1.1.2 Analysis of a Window Assembly

Consider a window assembly composed of concrete, glass, and aluminum, as shown in Fig. 9.29. The inside and outside ambient temperatures are $70^{\circ} \mathrm{F}$ and $0^{\circ} \mathrm{F}$, respectively, as indicated in Fig. 9.30. The film coefficient along the boundaries facing inside is $0.0139 \mathrm{Btu} / \mathrm{hr}-\mathrm{in}^{2}-{ }^{\circ} \mathrm{F}$ whereas the film coefficient along the boundaries facing outside is $0.0347 \mathrm{Btu} / \mathrm{hr}^{-} \mathrm{in}^{2} \mathrm{I}^{\circ} \mathrm{F}$. The thermal conductivities of glass, concrete, and aluminum are $0.05,0.06$, and $11 \mathrm{Btu} / \mathrm{hr}-\mathrm{in}-{ }^{\circ} \mathrm{F}$, respectively. The goal is to determine the steady-state temperature distribution in the system.

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Element Types dialog box appears; click on Add.
- Select Thermal Solid in the left list and Quad 4node 55 on the right list; click on $O K$.
- Click on Close.
- Specify thermal conductivities for aluminum (material 1), glass (material 2), and concrete (material 3) (MP command) using the following menu path:

Main Menu > Preprocessor > Material Props > Material Models

- In the Define Material Model Behavior dialog box, in the right window, successively double-click on Thermal, Conductivity, and Isotropic, which brings up another dialog box.
- Enter $\mathbf{1 1}$ for $K X X$; click on $O K$.
- Add new material for glass. In the Define Material Model Behavior dialog box, define a new material reference number using the following menu path:

Material > New Model


Fig. 9.27 Top view of thermal flux vectors.


Fig. 9.28 Left view of thermal flux vectors.


Fig. 9.29 The geometry of the window assembly.


Fig. 9.30 Boundary conditions for the window assembly.

- A dialog box appears with the material reference number set to the next available number (in this case 2); click on $\boldsymbol{O K}$.
- Note the new entry for Material Model Number 2 in the left list in the Define Material Model Behavior dialog box.
- In the right window, successively double-click on Thermal, Conductivity, and Isotropic, which brings up another dialog box.
- Enter 0.05 for $\boldsymbol{K X X}$; click on $O K$.
- Add new material for concrete and specify the thermal conductivity as 0.06 following the same procedure used for glass above.
- Close the Define Material Model Behavior dialog box by using the following menu path:


## Material > Exit

- Create the areas (RECTNG command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Rectangle $>$ By Dimensions

- Create Rectangle by Dimensions dialog box appears; enter 0 and 1.5 for the $\mathbf{X}$-coordinates ( $\mathbf{X 1}, \mathrm{X} 2$ ) and 0 and 2.25 for the $\boldsymbol{Y}$-coordinates (Y1, Y2); click on Apply.
- Referring to Table 9.3, create the remaining six areas. After creating the sixth area, click on $O K$ (instead of Apply).
- Figure 9.31 shows the six areas as they appear in the Graphics Window.
- Define a local coordinate system (LOCAL command) using the following menu path:

Utility Menu > WorkPlane > Local Coordinate Systems > Create Local CS > At Specified Loc

- Pick Мепи appears; type $1.75,1$ in the text box and hit enter (as shown in Fig. 9.32); click on $\boldsymbol{O K}$.

Table 9.3 Coordinates defining the rectangles.

| Area <br> Number | X1 | X2 | Y1 | Y2 |
| :---: | :---: | :---: | :---: | :---: |
| 1 | 0 | 1.5 | 0 | 2.25 |
| 2 | 1.5 | 1.75 | 0.75 | 2.25 |
| 3 | 0 | 1.75 | 2.25 | 2.5 |
| 4 | 0 | $3 / 16$ | 2.5 | 3 |
| 5 | $3 / 16$ | $3 / 8$ | 2.5 | 4 |
| 6 | $3 / 8$ | $9 / 16$ | 2.5 | 3 |



Fig. 9.31 Six areas as they appear in the Graphics Window.


Fig. 9.32 Pick Menu for creating a local coordinate system at a location.

- Create Local CS at Specified Location dialog box appears; select Cylindrical 1 from the KCS Type of coordinate system pull-down menu; click on $O K$.
- Observe that the local coordinate system appears in the Graphics Window.
- Create a keypoint in the local coordinate system ( K command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Keypoints > In Active CS

- Create Keypoints in Active Coordinate System dialog box appears; leave the NPT Keypoint number field blank (ANSYS issues the lowest available keypoint number to the newly created keypoints) and enter 1.5 and 70 for $X, Y, Z$ Location in active $C S$ (leave the $Z$ field blank). Since the active coordinate system is cylindrical, $X$ corresponds to the radius from the local coordinate system origin and $Y$ corresponds to the angle measured from the $x$-axis in the counterclockwise direction.
- Upon clicking on $\boldsymbol{O K}$, ANSYS creates the new keypoint and plots all of the keypoints, as shown in Fig. 9.33. Note that the most recently created keypoint is 25 .
m The slanted rectangular area and the circular portion at the end for the aluminum component are created next. For this purpose, the Working Plane is moved to an existing keypoint and rotated for the rectangular area. It is then moved one more time for the creation of the circular area.
- Activate the global Cartesian coordinate system (CSYS command) using the following menu path:

Utility Menu $>$ WorkPlane $>$ Change Active CS to $>$ Global Cartesian

- Move the Working Plane origin to keypoint 25 (KWPAVE command) using the following menu path:

Utility Menu > WorkPlane > Offset WP to > Keypoints

- Pick Menu appears; pick keypoint 25; click on OK. Observe the Working Plane origin at keypoint 25 in the Graphics Window.
- Rotate the Working Plane by $-20^{\circ}$ about the $z$-axis by (WPROTA command) using the following menu path:

Utility Menu $>$ WorkPlane $>$ Offset WP by Increments

- Offset WP dialog box appears, which has two main action areas, i.e., translations in the $x$-, $y$-, and $z$-directions, and rotations about $x$-, $y$-, and $z$-axes. Each of these areas also has a slider that controls the rate at


Fig. 9.33 Keypoints as they appear in the Graphics Window.
which translations or rotations are applied. By default, the translation slider is set to 1 and the rotation slider is set to $\mathbf{3 0}$. By moving the rotation slider to the left (or using the arrow keys), set the rate to be $20^{\circ}$. Upon clicking on the negative Z-button, the Working Plane is rotated $-20^{\circ}$ about the $z$-axis.

- Create the rectangular area (recting command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Areas > Rectangle > By Dimensions

- Create Rectangle by Dimensions dialog box appears; enter 0 and 1 for the $X$-coordinates ( $X 1, X 2$ ) and -0.25 and 0 for the $\boldsymbol{Y}$-coordinates ( $\mathbf{Y 1}$, Y2); click on $O K$.
- Turn keypoint numbering on (/PNOM command) using the following menu path:


## Utility Menu > PlotCtrls > Numbering

- Plot Numbering Controls dialog box appears; click on the box next to KP Keypoint numbers, which places a checkmark in the box. Select Numbers only from the [/NUM] Numbering shown with pull-down menu; click on $\boldsymbol{O K}$.
- Plot keypoints (KPLOT command) using the following menu path: Utility Menu > Plot > Keypoints > Keypoints
- Move the Working Plane origin to keypoint 27 (KWPAVE command) using the following menu path:

Utility Menu > WorkPlane > Offset WP to > Keypoints

- Pick Menu appears; pick keypoint 27; click on OK. Observe the Working Plane origin at keypoint 27 in the Graphics Window.
- Create the hollow circular area (PCIRC command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Circle $>$ By Dimensions

- Circular Area by Dimensions dialog box appears; enter 0.25 for RAD1 Outer radius and 0.125 for RAD2 Optional inner radius. Click on $O K$.
- Turn keypoint numbering off and area numbering on (/PNUM command) using the following menu path:

Utility Menu $>$ PlotCtrls $>$ Numbering

- Plot Numbering Controls dialog box appears; click on the box next to KP Keypoint numbers to remove the checkmark (turning keypoint numbering off) and click on the box next to AREA Area numbers to place a checkmark (turning area numbering on). Click on $O K$.
- Plot areas (APLOT command) using the following menu path: Utility Menu > Plot > Areas
- Figure 9.34 shows the areas as they appear in the Graphics Window.
- Overlap areas 7 and 8 (AOVLAP command) using the following menu path:

Main Menu > Preprocessor > Modeling $>$ Operate $>$ Booleans $>$ Overlap $>$ Areas

- Pick Menu appears; pick areas 7 and 8; click on $O K$ in the Pick Menu.
- Figure 9.35 shows the areas before and after overlapping.
- Delete area 9 and keypoints and lines attached to it (ADELE command) using the following menu path:


## Main Menu $>$ Preprocessor $>$ Modeling $>$ Delete $>$ Area and Below

- Pick Menu appears; pick area 9; click on OK in the Pick Menu.
- Observe that area 9 disappears in the Graphics Window.


Fig. 9.34 Areas as they appear in the Graphics Window.


Fig. 9.35 Areas before (left) and after (right) overlapping.

- Reset the Working Plane to its default configuration (WPSTYL command) using the following menu path:

Utility Menu > WorkPlane > Align WP with > Global Cartesian

- Working Plane is aligned with global Cartesian coordinate system (both the origin and the orientation).
- Toggle the Working Plane display off (WPSTYL command) using the following menu path:

Utility Menu > WorkPlane > Display Working Plane

- If a checkmark appears next to the menu entry Display Working Plane, which means the display is on, click on the menu entry to toggle off the display.
- Create a keypoint ( $\mathbf{K}$ command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create $>$ Keypoints $>$ In Active CS

- Create Keypoints in Active Coordinate System dialog box appears; enter 50 in the NPT Keypoint number field and 1.75 and 2 for $X, Y, Z$ Location in active CS (leave the $Z$ field blank). Click on $O K$.
- Activate local coordinate system 11 (created earlier) (CSYS command) using the following menu path:


## Utility Menu > Change Active CS to >Specified Coord Sys

- Change Active CS to Specified CS dialog box appears; enter 11 for KCN Coordinate system number; click on OK.
- Turn keypoint numbering on and area numbering off. Plot keypoints.

■ Create a line between keypoints 11 and 29 ( $L$ command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Lines $>$ Lines $>$ In Active Coord

- Pick Menu appears; pick keypoints 11 and 29. Keypoint 29 is coincident with keypoint 25; therefore, when attempting to pick keypoint 29 , ANSYS issues a warning to the user that there are more than one keypoints at the picked location and that the currently picked keypoint is keypoint 25. Click on Next in this Multiple_Entities Warning Window, which picks the next higher numbered keypoint (keypoint 29 in this case) at the coincident location. Click on $O K$ in the Multiple_Entities Warning Window; click on OK in the Pick Menu.
- The line that is just created is a curved line. This is because it is created in the active coordinate system, which is a local cylindrical coordinate system.
- Reset the active coordinate system to be global Cartesian (CSYS command) using the following menu path:

Utility Menu > WorkPlane $>$ Change Active CS to $>$ Global Cartesian
■ Create an area using keypoints 50,26,29, and 11 (A command) using the following menu path:
Main Menu > Preprocessor > Modeling > Create > Areas > Arbitrary > Through KPs

- Pick Menu appears; pick the keypoints listed above. Note that when picking keypoint 29 , since it is coincident with keypoint 25 , ANSYS warns the user that there are two keypoints at the picked location. Similar to the previous case, click on Next to pick keypoint 29. When
finished with picking all four keypoints, click on $O K$ in the Pick Menu and observe that area 7 appears in the Graphics Window.
- Since a curved line was created between keypoints 11 and 29 previously, and both of these keypoints are used in the definition of area 7 , the boundary of the area between keypoints 11 and 29 is a curved one. If the line were not created, then this boundary would have been a straight line.
- The circular hole within the aluminum component near the top-right corner of the concrete block is created next. For this purpose, the Working Plane origin is moved to keypoint 10 . After creating the circle, areas around the circle are overlapped and the ones defining the hole are deleted.
- Plot keypoints (KPLOT command) using the following menu path:

Utility Menu > Plot > Keypoints > Keypoints

- Move the Working Plane origin to keypoint 10 (kwPAve command) using the following menu path:

Utility Menu > WorkPlane > Offset WP to > Keypoints

- Pick Menu appears; pick keypoint 10 (keypoint 10 is coincident with keypoint 7 and they are located between keypoints 11 and 50 ) and click on OK. Observe the Working Plane origin at keypoint 10 in the Graphics Window.
- Create a circular area (PCIRC command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Circle $>$ By Dimensions

- Circular Area by Dimensions dialog box appears; enter 0.125 for RAD1 Outer radius and remove any leftover numbers from the RAD2 Optional inner radius field. Click on $O K$.
- Reset the Working Plane to its default configuration (WPSTYL command) using the following menu path:

Utility Menu > WorkPlane > Align WP with > Global Cartesian

- Working Plane is aligned with global Cartesian coordinate system (both the origin and the orientation).
- Toggle the Working Plane display off (WPSTYL command) using the following menu path:

Utility Menu > WorkPlane > Display Working Plane

- If a checkmark appears next to the menu entry Display Working Plane, which means the display is on, click on the menu entry to toggle off the display.
- Turn area numbering on (/PNUM command) and keypoint numbering off using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Plot Numbering Controls dialog box appears; click on the box next to AREA Area numbers to place a checkmark (turning area numbering on). Remove the checkmark for keypoints. Click on $\boldsymbol{O K}$.
- Plot areas (APLOT command) using the following menu path:

Utility Menu > Plot > Areas

- Overlap areas 2, 3, 7, and 8 (AOVLAP command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Operate $>$ Booleans $>$ Overlap $>$ Areas

- Pick Menu appears; pick areas $2,3,7$, and 8 and click on $\boldsymbol{O K}$ in the Pick Menu.
- Figures 9.36 and 9.37 show the areas before and after overlapping, respectively.
- Delete areas 9,13 , and 14 and the keypoints and lines attached to them (ADELE command) using the following menu path:
Main Menu $>$ Preprocessor $>$ Modeling $>$ Delete $>$ Area and Below
- Pick Menu appears; pick areas 9, 13, and 14; click on $O K$ in the Pick Menu.
- Observe that the areas disappear in the Graphics Window.
- Two fillets are created next.
- Turn line numbering on (/PNUM command) using the following menu path:

Utility Menu > PlotCtris > Numbering

- Plot Numbering Controls dialog box appears; click on the box next to LINE Line numbers to place a checkmark (turning line numbering on). Make sure no other numbering (keypoint and/or area) is on. Click on OK.
- Plot lines (LPLOT command) using the following menu path: Utility Menu > Plot > Lines
- Zoom-in to the region around the hole inside the aluminum component near the top-right corner of the concrete, as shown in Fig. 9.38.


Fig. 9.36 Areas before overlapping.


Fig. 9.37 Areas after overlapping.


Fig. 9.38 Lines around the hole.

- Create line fillets between lines 48 and 26 and between lines 26 and 39, both of them having a radius of 0.25 in (LFILLT command), using the following menu path:


## Main Menu > Preprocessor> Modeling > Create > Lines >Line Fillet

- Pick Menu appears, prompting the user to pick two intersecting lines. Pick lines 48 and 26; click on $O K$ in the Pick Menu.
- Line Fillet dialog box appears; enter 0.25 for RAD Fillet radius and 55 for PCENT Number to assign to generated keypoint at fillet center. Click on Apply to create the next fillet.
- Pick Menu reappears. Repeat the same procedure by picking lines 26 and 39 , using the same radius (0.25), and using 60 for $P C E N T$. When finished, click on $\boldsymbol{O K}$ in the Line Fillet dialog box.
- Turn keypoint numbering on (/PNUM command) using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Plot Numbering Controls dialog box appears; click on the box next to KP Keypoint numbers to place a checkmark. Make sure no other numbering (line and/or area) is on. Click on $\boldsymbol{O K}$.
- Plot keypoints (KPLOT command) using the following menu path:

Utility Menu > Plot > Keypoints > Keypoints

- Create areas (for fillets) through keypoints (A command) using the following menu path:

Main Menu > Preprocessor > Modeling > Create > Areas > Arbitrary > Through KPs

- Pick Menu appears; pick keypoints 7, 10, and 50; click on Apply in the Pick Menu.
- Pick Menu remains active; pick keypoints 40, 41, and 26; click on $O K$ in the Pick Menu.
- Turn area numbering on (/PNUM command) using the following menu path:

Utility Menu > PlotCtrls $>$ Numbering

- Plot Numbering Controls dialog box appears; click on the box next to AREA Area numbers to place a checkmark (turning area numbering on). Make sure no other numbering (keypoint and/or line) is on. Click on $O K$.
- Plot areas (APLOT command) using the following menu path:

Utility Menu > Plot > Areas

- Areas appear in the Graphics Window, as shown in Fig. 9.39.
- Add areas $2,3,10,15$, and 17 (AADD command) using the following menu path:

Main Menu > Preprocessor > Modeling > Booleans > Add > Areas

- Pick Menu appears; pick areas 2,3,10,15, and 17; click on $O K$ in the Pick Menu.
- Glue all areas (AGLUE command) using the following menu path:

Main Menu > Preprocessor > Modeling > Booleans $>$ Glue $>$ Areas

- Pick Menu appears; click on Pick All in the Pick Menu.
- Figure 9.40 shows the areas after adding and gluing operations as they appear in the Graphics Window.
- Specify size controls for meshing, compress entity numbers, and create the mesh.
- Specify global element size (ESIZE command) using the following menu path:


Fig. 9.39 Areas after fillet creation.


Fig. 9.40 Areas after adding and gluing operations.

- Global Element Sizes dialog box appears; enter 1/16 for SIZE Element edge length; click on OK.
- Turn keypoint numbering on (/PNUM command) using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Plot Numbering Controls dialog box appears; click on the box next to KP Keypoint numbers to place a checkmark. Make sure no other numbering (line and/or area) is on. Click on $\boldsymbol{O K}$.
- Plot keypoints (KPLOT command) using the following menu path: Utility Menu > Plot > Keypoints > Keypoints
- Specify element size near keypoints 1 and 2 as 0.25 (KESIZE command) using the following menu path:

```
Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Keypoints > Picked KPs
```

- Pick Menu appears; pick keypoints 1 and 2 (these are the bottom corners of the concrete block-keypoint 1 is at the global origin); click on OK in the Pick Menu.
- Element Size at Picked Keypoints dialog box appears; enter 0.25 for SIZE Element edge length; click on OK.
- Compress entity numbers (NUMCMP command) using the following menu path:

Main Menu > Preprocessor > Numbering Ctris > Compress Numbers

- Compress Numbers dialog box appears; select All from the Label Item to be compressed pull-down menu; click on OK.
- Turn area numbering on (/PNUM command) using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Plot Numbering Controls dialog box appears; click on the box next to AREA Area numbers to place a checkmark (turning area numbering on). Make sure no other numbering (keypoint and/or line) is on. Click on $\boldsymbol{O K}$.
- Plot areas (APLOT command) using the following menu path:

Utility Menu > Plot > Areas

- Areas appear in the Graphics Window, as shown in Fig. 9.41.
- Mesh is generated next.
- Change default material attribute to 3 (MAT command) using the following menu path:


Fig. 9.41 Areas after entity number compression.

- Meshing Attributes dialog box appears; select 3 from the [MAT] Material number pull-down menu; click on $\boldsymbol{O K}$. Mesh generated after this point will have material reference number 3, until it is changed to another number.
- Create mesh for area 2 (concrete) (AMESH command) using the following menu path:
Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Free
- Pick Меnu appears; pick area 2; click on $\boldsymbol{O K}$.
- Plot areas (APLOT command) using the following menu path:

Utility Menu > Plot > Areas

- Change default material attribute to 2 (MAT command) using the following menu path:

Main Menu > Preprocessor > Meshing > Mesh Attributes > Default Attribs

- Meshing Attributes dialog box appears; select 2 from the [MAT] Material number pull-down menu; click on $\boldsymbol{O K}$.
- Create mesh for area 4 (glass) (AMESH command) using the following menu path:
- Pick Menu appears; pick area 4; click on $O K$.
- Plot areas (APLOT command) using the following menu path: Utility Menu > Plot > Areas
- Change default material attribute to 1 (MAT command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh Attributes $>$ Default Attribs

- Meshing Attributes dialog box appears; select 1 from the [MAT] Material number pull-down menu; click on $O K$.
- Create mesh for the remaining areas (aluminum) (AMESH command) using the following menu path:

Main Menu > Preprocessor > Meshing > Mesh $>$ Areas > Free

- Pick Menu appears; pick the remaining areas (1,3,5-8); click on $\boldsymbol{O K}$.
- A warning message may appear; if so, close it.
- Turn the element numbering on so that the elements are plotted with different colors based on their material attribute (/PNUM command) using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Plot Numbering Controls dialog box appears; select Material numbers from the Elem / Attrib numbering pull-down menu; select Colors only from the [/NUM] Numbering shown with pull-down menu. Click on OK.
- Plot elements (EPLOT command) using the following menu path:

Utility Menu > Plot $>$ Elements

- Elements appear in the Graphics Window, as shown in Fig. 9.42.


## SOLUTION

- Apply convective boundary conditions.
- Turn line numbering on (/PNUM command) using the following menu path:

Utility Menu > PlotCtrls > Numbering

- Plot Numbering Controls dialog box appears; click on the box next to LINE Line numbers to place a checkmark. Select No numbering from the Elem / Attrib numbering pull-down menu, and select Numbers only from the [/NUM] Numbering shown with pull-down menu. Click on $O K$.


Fig. 9.42 Elements after meshing plotted with different colors based on their material number attributes.

- Plot lines (LPLOT command) using the following menu path:

Utility Menu > Plot > Lines

- Apply convective boundary conditions on the lines along the side facing outside (as shown in Fig. 9.30) (SFL command) using the following menu path:


## Main Menu > Solution > Define Loads > Apply > Thermal > Convection > On Lines

- Pick Мепи appears; pick the lines shown in Fig. 9.43; click on $O K$ in the Pick Menu.
- Apply CONV on lines dialog box appears; type 0.0347 for VALI Film coefficient and 0 for VAL2I Bulk temperature; click on OK.
- Apply convective boundary conditions on the lines along the side facing inside (as shown in Fig. 9.30) (SFL command) using the following menu path:


## Main Menu > Solution > Define Loads > Apply > Thermal > Convection > On Lines

- Pick Menu appears; pick the lines shown in Fig. 9.44; click on $O K$ in the Pick Menu.
- Apply CONV on lines dialog box appears; type 0.0139 for VALI Film coefficient and 70 for VAL2I Bulk temperature; click on OK.


Fig. 9.43 Lines facing outside.


Fig. 9.44 Lines facing inside.

- Obtain the solution (SOLVE command) using the following menu path:

Main Menu > Solution > Solve >Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on $O K$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Turn colors on ( / PNUM command) using the following menu path: Utility Menu $>$ PlotCtrls $>$ Numbering
- Plot Numbering Controls dialog box appears; click on the box next to LINE Line numbers to remove the checkmark. Select Colors only from the [/NUM] Numbering shown with pull-down menu; click on OK.
$\square$ Review temperature contours (PLNSOL command) using the following menu path:

Main Menu > General Postproc > Plot Results > Contour Plot > Nodal Solu

- Contour Nodal Solution Data dialog box appears. Select DOF solution from the left list and Temperature TEMP from the right list; click on OK
- The temperature contour plot is shown in Fig. 9.45 as it appears in the Graphics Window
- Review thermal flux and gradient vectors using the following menu path: Main Menu > General Postproc > Plot Results > Vector Plot > Predefined
- Vector Plot of Predefined Vectors dialog box appears. Select Flux \& gradient from the left list and Thermal flux TF from the right list; click on $O K$.
- Figure 9.46 shows the flux vectors as they appear in the Graphics Window.
- Repeat the same procedure for the thermal gradient by selecting Thermal grad $T \boldsymbol{T}$ from the right list. Fig. 9.47 shows the thermal gradient vectors as they appear in the Graphics Window.


### 9.1.2 Transient Analysis

A transient analysis is utilized to simulate the heat transfer phenomenon in the presence of time-dependent boundary conditions, body loads, and/or initial conditions. In a transient analysis, in addition to the initial and boundary


Fig. 9.45 Contour plot of temperatures.


Fig. 9.46 Vector plot of thermal flux.


Fig. 9.47 Vector plot of thermal gradient.
conditions and the body loads, the user must specify time-related quantities, such as time step size, number of load steps, number of substeps, and the final time. Depending on the values of these quantities, solutions to the same problem may differ considerably. As a rule of thumb, the solution is expected to be more accurate as the time step size decreases (increased number of substeps). However, this may increase the cost of analysis significantly. Therefore, the user has to build a certain level of knowledge through numerical experimentation on the time-dependent parameters.

### 9.1.2.1 Transient Thermomechanical Analysis of an Electronic Package

A common cause of failure in electronic devices is the thermal stresses at elevated temperatures caused by a coefficient of thermal expansion mismatch. In most thermomechanical analyses, the thermal load is assumed to be uniform. However, a nonuniform temperature distribution may be required for specific cases. In order to obtain a transient thermal stress field, first, a transient heat conduction (diffusion) solution is obtained, followed by a stress analysis utilizing the nonuniform thermal field at a specific time as body loading. ANSYS provides a convenient method for achieving this task. In order to demonstrate this capability, a problem involving an
electronic device is considered. The device contains a silicon die (chip), epoxy die-attach, substrate, and a copper heat spreader, as shown in Fig. 9.48. Thermal and mechanical properties of the constituent materials are given in Table 9.4. The solid model is generated utilizing rectangles with the coordinates given in Table 9.5, along with their associations with material reference numbers.

In this transient thermal analysis, the surrounding air is at a temperature of $T_{\infty}=25^{\circ} \mathrm{C}$ (ambient temperature). All of the surfaces, except the symmetry line, are subjected to convective heat loss with a heat transfer coefficient of $h=5 \mathrm{~W} /\left(\mathrm{m}^{2} \cdot{ }^{\circ} \mathrm{C}\right)$. There is no heat transfer through the symmetry line (insulation). In ANSYS thermal analyses, when boundary conditions are not specified along a boundary, insulation is imposed automatically. Heat is generated at the bottom face of the die, which is expressed in terms of a constant heat flux of $q=1000 \mathrm{~W} / \mathrm{m}$. The initial temperature of the device is assumed to be uniform at $T_{0}=25^{\circ} \mathrm{C}$. Heat transfer is simulated for a period of $5 \mathrm{~min}(300 \mathrm{sec})$, after which the device reaches a steady state.

Once the time-dependent temperature field is obtained, the stress field is calculated based on the temperatures at desired time points.

### 9.1.2.1.1 Transient Thermal Analysis

## MODEL GENERATION

- Specify the jobname (/FILNAM command) using the following menu path:


## Utility Menu > File > Change Jobname

- Change Jobname dialog box appears. Type TH; click on $\boldsymbol{O K}$.


Fig. 9.48 Schematic of the electronic package.

Table 9.4 Properties of the constituent materials in the electronic package.

|  | Substrate | Die- <br> Attach | Silicon | Copper |
| :--- | :---: | :---: | :---: | :---: |
| Material reference no. | 1 | 2 | 3 | 4 |
| $E(\mathrm{GPa})$ | 22 | 7.4 | 163 | 129 |
| $\nu$ | 0.39 | 0.4 | 0.278 | 0.344 |
| $\alpha\left(10^{-6} /{ }^{\circ} \mathrm{C}\right)$ | 18 | 52 | 2.6 | 14.3 |
| $\kappa\left(\mathrm{~W} / \mathrm{m} \cdot{ }^{\circ} \mathrm{C}\right)$ | 2 | 100 | 150 | 396 |
| $c\left(\mathrm{~J} / \mathrm{kg} \cdot{ }^{\circ} \mathrm{C}\right)$ | 840 | 535 | 703 | 384 |
| $\rho\left(\mathrm{~kg} / \mathrm{m}^{3}\right)$ | 220 | 6450 | 2330 | 8940 |

Table 9.5 Coordinates defining the areas and the corresponding material reference numbers.

| Area <br> No. | X1 | X2 | Y1 | Y2 | Mat. Ref. <br> No. |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | 0 | 5 | 0 | 2 | 1 |
| 2 | 5 | 7.5 | 0 | 2 | 1 |
| 3 | 7.5 | 10 | 0 | 2 | 1 |
| 4 | 0 | 5 | 2 | 2.1 | 2 |
| 5 | 0 | 5 | 2.1 | 3.1 | 3 |
| 6 | 0 | 5 | 3.1 | 5.1 | 4 |

- Define the element type (ET command) using the following menu path:


## Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Element Types dialog box appears; click on Add.
- Select Thermal Solid in the left list and Quad 4 Node 55 on the right list; click on OK.
- Click on Close.
- Specify material properties (MP command) using the following menu path:

Main Menu > Preprocessor $>$ Material Props $>$ Material Models

- Define Material Model Behavior dialog box appears. Structural properties are first specified, followed by thermal properties.
- In the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which brings up another dialog box
- Referring to Table 9.4, enter 22E9 for $\boldsymbol{E X}$ and $\mathbf{0 . 3 9}$ for $\boldsymbol{P R X Y}$; click on OK
- In the right list, successively double-click on Structural, Thermal Expansion, Secant Coefficient, and, finally, Isotropic, which brings up another dialog box.
- Enter $\mathbf{1 8 E - 6}$ for $\mathbf{A L P X}$; click on $\mathbf{O K}$.
- Specify density by successively double-clicking on Structural and Density, which brings up another dialog box. Type 220, click on OK.
- In order to specify thermal properties, first successively double-click on Thermal, Conductivity, and Isotropic; then enter 2 in the newly appeared dialog box; click on $\boldsymbol{O K}$. Thermal conductivity is specified.
- Specify specific heat by successively double-clicking on Thermal and Specific Heat; enter 840 in the newly appeared dialog box; click on OK.
- Add new material model using the following menu path:


## Material > New Model

- Repeat the procedure for the remaining materials (2 through 4 ).
- When finished, close the Define Material Model Behavior dialog box by using the following menu path:

```
Material > Exit
```

- Create rectangles as identified in Fig. 9.49 (RECTNG command) using the following menu path:


## Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Rectangle $>$ By Dimensions

- Create Rectangle by Dimensions dialog box appears. Referring to Table 9.5, enter $\mathbf{0}$ and $\mathbf{5 E - 3}$ for $\boldsymbol{X 1}$ and $\boldsymbol{X 2}$ and $\mathbf{0}$ and $\mathbf{2 E - 3}$ for $\boldsymbol{Y 1}$ and Y2; click on Apply.
- Repeat the procedure for the remaining areas (2 through 6 ). When creating Area 6 , click on $\boldsymbol{O K}$ after entering the coordinates.


Fig. 9.49 Solid model of the electronic device with area numbers turned on.

Glue the areas (AGLUE command) using the following menu path:
Main Menu $>$ Preprocessor $>$ Modeling $>$ Operate $>$ Booleans $>$ Glue $>$ Areas

- Pick Menu appears; click on Pick All button.

Mesh the areas (AMESH command) using the following menu path:
Main Menu > Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 sided

- Pick Menu appears; click on Pick All.
- At this point, all the elements have Material Reference Number 1. Attributes can be changed after the elements are created. For this purpose, the areas are selected and then the elements that are attached to the selected areas are selected. Finally, elements are modified to have the correct attributes. The correspondence between the areas and material numbers is given in Table 9.5. Select areas (ASEL command) using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; select Areas from the first pulldown menu; click on $O K$.
- Pick Menu appears; pick area 4 as defined in Table 9.5; click on $O K$.
- Now, select the elements that are attached to the selected areas (ESLA command) using the following menu path:


## Utility Menu > Select > Entities

- Select Entities dialog box appears; select Elements from the first pulldown menu; select Attached to from the second pull-down menu. Click on the Areas radio-button; click on $\boldsymbol{O K}$.
- Modify the attributes of the selected set of elements (EMODIf command) using the following menu path:


## Main Menu > Preprocessor > Modeling > Move $/$ Modify > Elements > Modify Attrib

- Pick Menu appears; click on Pick All, which brings up the Modify Elem Attributes dialog box.
- Select Material MAT from the pull-down menu; enter 2 in the $\boldsymbol{I I}$ New attribute number field; click on $O K$.
- Repeat this procedure for area 5 (material reference number 3) and area 6 (material reference number 4).
- When finished, select everything (ALLSEL command) using the following menu path:
- Save the database (SAVE command) using the following menu path:

Utility Menu > Save as Jobname.db

- The database is saved in file TH.db.


## SOLUTION

- Declare the new analysis to be a transient analysis (ANTYPE command) using the following menu path:
Main Menu > Solution > Analysis Type > New Analysis
- New Analysis dialog box appears; click on the Transient radio-button; click on $\boldsymbol{O K}$.
- Transient Analysis dialog box appears; click on $\boldsymbol{O K}$.
- Apply convective boundary conditions on lines (SFL command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Thermal > Convection > On Lines

- Pick Menu appears; pick the exterior lines except the ones along $x=0$ (lines to be picked are shown in Fig. 9.50); click on $\boldsymbol{O K}$ in the Pick Мепи.
- Enter 5 for VALI Film coefficient and 25 for VAL2I Bulk temperature; click on $O K$.
- Apply the heat flux condition (SFL command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Thermal > Heat Flux > On Lines


Fig. 9.50 Lines to be selected for convective boundary conditions.

- Pick Menu appears; pick the line between the die and the die-attach; click on $O K$ in the Pick Menu.
- Apply HFLUX on lines dialog box appears; enter 1000 for VALI Heat flux; click on $O K$.
- Apply the initial condition on the nodes (IC command) using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Initial Condit' $\gg$ Define

- Pick Menu appears; click on Pick All in Pick Menu.
- Define Initial Conditions dialog box appears; select TEMP from the pull-down menu; enter 25 for VALUE Initial value of DOF; click on OK.
- Specify solution controls using the following menu path:

Main Menu > Solution > Analysis Type > Sol'n Controls

- Solution Controls dialog box appears, which has five tabs. In the Basic tab, within the Time Control field, enter 300 for Time at end of loadstep and select Off from the Automatic time stepping pull-down menu.
- Enter 100 for Number of substeps.
- In the Write Items to Results File field, select Write every substep from the Frequency pull-down menu; click on $O K$.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu >Solution > Solve >Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on $O K$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Results are stored in the TH.rth file in the Working Directory. Review the temperature distribution at different time steps using the following menu path:
Main Menu > General Postproc > Read Results > By Load Step
- Read Results by Load Step Number dialog box appears; enter 10 for SBSTEP Substep number; click on OK.
- Plot temperature contours (PLNSOL command) at the selected substep using the following menu path:

Main Menu > General Postproc > Plot Results > Contour Plot > Nodal Solu

- Contour Nodal Solution Data dialog box appears; click on OK. The contour plot appears, as shown in Fig. 9.51.
- Repeat this procedure for substeps 50 and 100. Contour plot corresponding to substep 100 (time $=300 \mathrm{sec}$ ) is shown in Fig. 9.52.
- Stress fields corresponding to substeps 10,50 , and 100 are obtained in the next subsection.


### 9.1.2.1.2 Thermomechanical Analysis

- Clear the database (/CLEAR command) using the following menu path: Utility Menu > File > Clear \& Start New
- Clear Database and Start New dialog box appears; click on OK.
- Verify dialog box appears; click on Yes.
- Resume from the TH. $d b$ database file (resume command) using the following menu path:

Utility Menu > File $>$ Resume from

- Resume Database dialog box appears. Browse for TH.db; click on OK.


## PREPROCESSOR

- Change the element type from SOLID55 to SOLID42 (ETCHG command) using the following menu path:

Main Menu > Preprocessor > Element Type > Switch Elem Type

- Switch Elem Type dialog box appears. Select Thermal to Struc from the pull-down menu; click on $\boldsymbol{O K}$.
- A Warning Window appears; click on Close.


Fig. 9.51 Contour plot of temperature at substep 10 (time $=30 \mathrm{sec}$ ).


Fig. 9.52 Contour plot of temperature at substep 100 (time $=300 \mathrm{sec})$.

- Modify the element type to be Plane Strain (KEYOPT command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Element Types dialog box appears; click on Options.
- PLANE42 element type options dialog box appears; select Plane strain from the Element behavior K3 pull-down menu; click on OK.
- Click on Close in Element Types dialog box.


## SOLUTION

- Declare the new analysis to be static (ANTYPE command) using the following menu path:
Main Menu > Solution > Analysis Type > New Analysis
- New Analysis dialog box appears; click on the Static radio-button; click on $\boldsymbol{O K}$.
- Apply displacement constraints (D command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

- Pick Menu appears; pick the nodes along $x=0$ ( $y$-axis); click on $\boldsymbol{O K}$.
- Apply U,ROT on Nodes dialog box appears; highlight $\boldsymbol{U X}$; and click on Apply.
- Pick Menu reappears; pick the node at $(x, y)=(0,0)$; click on $\boldsymbol{O K}$.
- Apply U,ROT on Nodes dialog box reappears; highlight $\boldsymbol{U X}$ and $\boldsymbol{U Y}$; click on $\boldsymbol{O K}$.
- Specify the stress-free temperature (TREF command) using the following menu path:


## Main Menu $>$ Solution $>$ Define Loads $>$ Settings $>$ Reference Temp

- Reference Temperature dialog box appears; enter 25 for Reference temperature; click on $\boldsymbol{O K}$.
- Read nonuniform temperature field as body load (LDREAD command) using the following menu path:

Main Menu > Solution > Define Loads > Apply $>$ Structural $>$ Temperature $>$ From Therm Analy

- Apply TEMP from Thermal Analysis dialog box appears; enter 1 and 10 for Load step and substep no; click on Browse.
- Fname Name of results file dialog box appears; browse for and select the TH.rth file; click on Open.
- Click on $O K$ in the Apply TEMP from Thermal Analysis dialog box.
- In real applications in the electronics industry, the copper heat spreader and the die are not in perfect contact; a thermal grease compound is dispensed in between these two components. Therefore, the copper heat spreader will be excluded from the analysis here in order to represent the real situation better. This will be achieved by selecting the nodes associated the substrate, die-attach, and die only (excluding those associated with the heat spreader). Select elements with material numbers 1,2 , and 3 (ESEL command) using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; select Elements from the first pulldown menu and By Attributes from the second pull-down menu. Click on the Material num radio-button and enter 1,3,1 in the Min,Max,Inc text field. Click on Apply.
- In the Select Entities dialog box, select Nodes from the pull-down menu and Attached to from the second pull-down menu. Click on the Elements radio-button; click on $O K$.
- Now, only the nodes associated with the substrate, die-attach, and die are selected (active) and only they will be included in the solution process.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu >Solution >Solve >Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the deformed shape resulting from the nonuniform temperature loading (PLDISP command) using the following menu path:


## Main Menu > General Postproc > Plot Results > Deformed Shape

- Plot Deformed Shape dialog box appears; click on the Def + undef edge radio-button; click on $O K$.
- The deformed shape appears, as shown in Fig. 9.53.

■ Review normal and shear stress contours (PLNSOL command) using the following menu path:

Main Menu $>$ General Postproc $>$ Plot Results $>$ Contour Plot $>$ Nodal Solu

- Contour Nodal Solution Data dialog box appears; select Stress from the left list and $\boldsymbol{Y}$-direction $S Y$ from the right list; click on $O K$.
- The contour plot of normal stress in the $y$-direction (peeling stress) appears, as shown in Fig. 9.54.
- Repeating this procedure for shear stresses, the contour plot appears, as shown in Fig. 9.55.
- The thermomechanical solution may be repeated for any substep of interest. Figures 9.56 and 9.57 show the contour plots of normal stress in the $y$-direction and shear stress for substep 100 (last substep, time $=$ 300 sec ), respectively.


### 9.1.2.2 Transient Thermomechanical Analysis of a Welded Joint

Consider the welded steel joint shown in Fig. 9.58 (due to symmetry with respect to the plane of the weld pool, only half of the geometry is shown). The plate is 12.5 cm long, 2 cm high, and 10 cm wide. The weld pool is assumed to be 0.5 cm long. In this heat transfer analysis, the surrounding air is at a temperature of $T_{\infty}=229.82^{\circ} \mathrm{K}$ (ambient temperature). The temperature-dependent film coefficients for the top, bottom, and side surfaces are given in Table 9.6, as well as the temperature-dependent thermal conductivity The density and specific heat are assumed to be constant at $\rho=7850 \mathrm{~kg} / \mathrm{m}^{3}$ and $c=500 \mathrm{~J} /\left(\mathrm{kg} \cdot{ }^{\circ} \mathrm{K}\right)$, respectively. The temperature is fixed at $299.82^{\circ} \mathrm{K}$ along the surface coinciding with the $x-y$ plane, and there is no heat transfer along the opposite surface (symmetry plane). The initial temperature for the weld pool is $1852.94^{\circ} \mathrm{K}$, and for the remaining volume, $299.82^{\circ} \mathrm{K}$. In this thermomechanical analysis, the plate is fixed in all directions along the side coinciding with the $x-y$


Fig. 9.53 Deformed shape of the electronic package due to thermal loading at load step 1 , substep 10 .


Fig. 9.54 Contour plot of the peeling stress due to thermal loading at load step 1, substep 10 .


Fig. 9.55 Contour plot of the shearing stress due to thermal loading at load step 1 , substep 10 .


Fig. 9.56 Contour plot of the peeling stress due to thermal loading at load step 1 , substep 100 .


Fig. 9.57 Contour plot of the shearing stress due to thermal loading at load step 1, substep 100 .


Fig. 9.58 Schematic of the welded joint.

Table 9.6 Temperature-dependent properties.

| Temperature <br> $\left({ }^{\circ} \mathrm{K}\right)$ | Thermal <br> Conductivity <br> $\left[\mathrm{W} /\left(\mathrm{m} \cdot{ }^{\circ} \mathrm{K}\right)\right]$ | Film Coefficient $\left[\mathrm{W} /\left(\mathrm{m}^{2} \cdot{ }^{\circ} \mathrm{K}\right)\right]$ |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | Top Surface | Bottom <br> Surface | Side <br> Surfaces |  |  |
| 433.15 | 30.158 | 9.08 | 4.99 | 8.18 |  |
| 593.15 | 30.385 | 11.69 | 6.05 | 9.89 |  |
| 753.15 | 30.611 | 13.57 | 6.79 | 10.96 |  |
| 913.15 | 30.838 | 14.96 | 7.28 | 11.86 |  |
| 1073.15 | 31.064 | 16.19 | 7.69 | 12.51 |  |
| 1233.15 | 31.290 | 17.25 | 8.09 | 13.16 |  |
| 1393.15 | 31.517 | 18.15 | 8.42 | 13.66 |  |
| 1553.15 | 31.743 | 18.97 | 8.67 | 14.15 |  |
| 1713.15 | 31.969 | 19.79 | 8.99 | 14.55 |  |
| 1873.15 | 25.338 | 20.52 | 9.24 | 14.96 |  |

plane, and the symmetry plane is constrained in the $z$-direction only. The elastic modulus, Poisson's ratio, and coefficient of thermal expansion are $E=200 \mathrm{GPa}, v=0.3$, and $\alpha=60 \times 10^{-6} \mathrm{ppm} /{ }^{\circ} \mathrm{K}$, respectively. The goal is to obtain a time-dependent thermal solution followed by a thermomechanical solution, which provides displacement and stress fields at different times.

### 9.1.2.2.1 Transient Thermal Analysis

## mOdEL GENERATION

- Specify the jobname (/FILNAM command) using the following menu path:

Utility Menu > File > Change Jobname

- Change Jobname dialog box appears; type WELD; click on OK.
- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Element Types dialog box appears; click on Add.
- Select Thermal Solid in the left list and Brick 8node 70 on the right list; click on $\boldsymbol{O K}$.
- Click on Close.
- Specify material properties (MP command) using the following menu path:
- Define Material Model Behavior dialog box appears. Structural properties are specified first, followed by thermal properties.
- In the right window, successively double-click on Structural, Linear, Elastic, and, finally, Isotropic, which brings up another dialog box.
- Enter 200E9 for $E X$ and 0.3 for PRXY; click on $O K$
- In the right list, successively double-click on Structural, Thermal Expansion, Secant Coefficient, and, finally, Isotropic, which brings up another dialog box.
- Enter 3.6E-5 for ALPX; click on $O K$.
- Specify density by successively double-clicking on Structural and Density, which brings up another dialog box. Type 7850 ; click on $O K$.
- Specify specific heat by successively double-clicking on Thermal and Specific Heat; enter 500 in the newly appeared dialog box; click on OK.
- Specify temperature-dependent thermal conductivity by successively double-clicking on Thermal, Conductivity, and Isotropic.
- Conductivity for Material Number 1 dialog box appears; click on the Add Temperature button 9 times (so that there are 10 temperature columns). Referring to Table 9.6, enter temperature values and corresponding thermal conductivity values. When finished, click on $O K$.
- Temperature-dependent film coefficient values for top, bottom, and side surfaces are input as having different material reference numbers. Add new material model using the following menu path:


## Material > New Model

- In the right list, successively double-click on Thermal and Convection or Film Coef., which brings up the Convection of Film Coefficient for Material 2 dialog box. Click on the Add Temperature button 9 times (so that there are 10 temperature columns). Referring to Table 9.6, enter temperature values and corresponding film coefficient values at the top surface. When finished, click on $O K$.
- Repeat the procedure for film coefficient values at the bottom (material 3) and side (material 4) surfaces.
- When finished, close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create rectangular blocks (BLOCK command) using the following menu path:
- Create Block by Dimensions dialog box appears. Referring to Table 9.7, enter 0 and $10 E-2$ for $X 1$ and $X 2,0$ and $2 E-2$ for $Y 1$ and $Y 2$, and 0 and $3 E-2$ for $\mathbf{Z 1}$ and $\mathbf{Z 2}$; click on Apply.
- Repeat the procedure for the remaining volumes (2 through 4). When creating volume 4 , click on $O K$ after entering the coordinates (instead of Apply).
- Glue the volumes (VGLUE command) using the following menu path:

Main Menu > Preprocessor > Modeling > Operate > Booleans > Glue > Volumes

- Pick Menu appears; click on Pick All button.

Mesh the volumes (VMESH command) using the following menu path:
Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Volumes $>$ Mapped $>4$ to 6 sided

- Pick Menu appears; click on Pick All.

Save the database (SAVE command) using the following menu path:
Utility Menu > Save as Jobname.db

- The database is saved in file WELD.db.


## SOLUTION

- Declare the new analysis to be a transient analysis (ANTYPE command) using the following menu path:
Main Menu > Solution > Analysis Type $>$ New Analysis
- New Analysis dialog box appears; click on the Transient radio-button; click on $\boldsymbol{O K}$.
- Transient Analysis dialog box appears; click on $O K$.
- Specify solution controls using the following menu path:

Main Menu > Solution > Analysis Type > Sol'n Controls

Table 9.7 Coordinates defining the volumes.

| Volume <br> Number | XI | X 2 | Y 1 | Y 2 | Z 1 | Z 2 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $----------(\mathrm{cm})$---------------- |  |  |  |  |  |  |
| 1 | 0 | 10 | 0 | 2 | 0 | 3 |
| 2 | 0 | 10 | 0 | 2 | 3 | 8 |
| 3 | 0 | 10 | 0 | 2 | 8 | 12 |
| 4 | 0 | 10 | 0 | 2 | 12 | 12.5 |

- Solution Controls dialog box appears, which has five tabs. In the Basic tab, within the Time Control field, enter 3600 for Time at end of loadstep.
- Click on the Time increment radio-button and enter 36, 3.6, and 500 for Time step size, Minimum time step, and Maximum time step, respectively.
- In the Write Items to Results File field, select Write every substep from the Frequency pull-down menu; click on $O K$.
- Initial conditions are specified next. For this purpose, volumes are selected first, followed by selection of the nodes attached to the selected volumes.
- Select volumes (VSEL command) using the following menu path:

Utility Menu > Select > Select Entities

- Select Entities dialog box appears; choose Volumes from the first pulldown menu; click on $\boldsymbol{O K}$.
- Pick Menu appears; pick the volumes indicated in Fig. 9.59; click on OK in the Pick Menu.
- Select nodes attached to the selected volumes (NSLV command) using the following menu path:

Utility Menu > Select > Select Entities

- Select Entities dialog box appears; choose Nodes from the first pulldown menu and Attached to from the second pull-down menu. Click on the Volumes, all radio-button; click on $O K$.
- Apply initial conditions on the selected nodes (IC command) using the following menu path:
Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Initial Condit'n $>$ Define
- Pick Menu appears; click on Pick All.
- Define Initial Conditions dialog box appears; select TEMP from the pull-down menu and enter 299.8167 for VALUE Initial value of DOF; click on $O K$.
- Select everything (ALLSEL command) using the following menu path: Utility Menu > Select > Everything
- Repeat the same procedure to apply initial conditions on nodes attached to the remaining volume corresponding to the welded region. This time, apply a temperature of $1852.594^{\circ} \mathrm{K}$.
- Select everything (ALLSEL command) using the following menu path:


Fig. 9.59 Volumes to be selected for initial temperatures.

- Apply temperature boundary conditions on the selected nodes (D command) using the following menu path:

Main Menu > Solution > Define Loads $>$ Apply $>$ Thermal $>$ Temperature $>$ On Nodes

- Pick Menu appears; pick the nodes along the $z=0$ surface and click on $\boldsymbol{O K}$ (in order to pick the nodes conveniently, plot the nodes and obtain a top view).
- Apply TEMP on Nodes dialog box appears; highlight TEMP and type 299.8167 for VALUE Load Temp value; click on OK.

■ Convective boundary conditions are applied next. Temperature-dependent film coefficients were specified previously for the top (material 2), bottom (material 3), and side (material 4) surfaces.

- Apply convective boundary conditions on the nodes along the side surfaces (SF command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Thermal > Convection >On Nodes

- Pick Menu appears; pick the nodes along the $x=0$ and $x=10 \times 10^{-2}$ surfaces (side surfaces); click on OK in the Pick Menu. For this case, Front View is the most convenient viewpoint.
- Apply CONV on nodes dialog box appears; enter -4 for film coefficient (first text field) and 299.8167 for bulk temperature (second text field); click on Apply.
- Pick Menu reappears; pick the nodes along the $y=2 \times 10^{-2}$ surface (top surface); click on $O K$ in the Pick Menu. Front View is the most convenient viewpoint for this case, also.
- Apply CONV on nodes dialog box appears; enter -2 for film coefficient and 299.8167 for bulk temperature; click on Apply.
- Pick Menu reappears; pick the nodes along the $y=0$ surface (bottom surface); click on OK in the Pick Menu. Right View and Left View are the most convenient viewpoints for this case.
- Apply CONV on nodes dialog box appears; enter -3 for film coefficient and 299.8167 for bulk temperature; click on $O K$.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu > Solùtion > Solve >Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window and click on OK in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Results are stored in the WELD.rth file in the Working Directory. Review the temperature distribution at different time points using the following menu path:

Main Menu > General Postproc > Read Results > By Pick

- Results File: WELD_TH.rth dialog box appears, in which available solution sets are tabulated. Highlight set 5; click on Read and click on Close.
- Plot temperature contours (PLNSOL command) at the selected substep using the following menu path:

Main Menu $>$ General Postproc $>$ Plot Results $>$ Contour Plot $>$ Nodal Solu

- Contour Nodal Solution Data dialog box appears; click on OK. The contour plot appears, as shown in Fig. 9.60.
- Repeat this procedure for substep 22 (last substep). Corresponding contour plot is shown in Fig. 9.61.
$\pm$ Stress fields corresponding to substeps 10,50 , and 100 are obtained in the next subsection.


Fig. 9.60 Contour plot of temperature at substep $5($ time $=69.409 \mathrm{sec})$.


Fig. 9.61 Contour plot of temperature at substep 22 (time $=3600 \mathrm{sec})$.

### 9.1.2.2.2 Thermomechanical Analysis

- Clear the database (/CLEAR command) using the following menu path: Utility Menu > File > Clear \& Start New
- Clear Database and Start New dialog box appears; click on OK.
- Verify dialog box appears; click on Yes.
- Resume from the WELD.db database file (RESUME command) using the following menu path:

Utility Menu > File > Resume from

- Resume Database dialog box appears. Browse for WELD.db; click on OK.


## PREPROCESSOR

■ Change the element type from SOLID70 to SOLID45 (ETCHG command) using the following menu path:

Main Menu > Preprocessor > Element Type > Switch Elem Type

- Switch Elem Type dialog box appears. Select Thermal to Struc from the pull-down menu; click on $\boldsymbol{O K}$.
- A Warning Window appears; click on Close .


## SOLUTION

- Declare the new analysis to be static (ANTYPE command) using the following menu path:

Main Menu >Solution > Analysis Type $>$ New Analysis

- New Analysis dialog box appears; click on the Static radio-button; click on $\boldsymbol{O K}$.
- Apply displacement constraints (D command) using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Structural $>$ Displacement $>$ On Nodes

- Pick Menu appears; pick the nodes along $z=0$; click on $\boldsymbol{O K}$.
- Apply U,ROT on Nodes dialog box appears; highlight All DOF; click on Apply.
- Pick Menu reappears; pick the nodes along $z=12.5 \times 10^{-2}$; click on OK.
- Apply U,ROT on Nodes dialog box reappears; highlight $U Z$ and remove the highlight from All DOF; click on OK.
- Specify the stress-free temperature (TREF command) using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Settings $>$ Reference Temp

- Reference Temperature dialog box appears; enter 299.8167 for Reference temperature; click on OK.
- Read the nonuniform temperature field as body load (LDREAD command) using the following menu path:
Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Structural $>$ Temperature $>$ From Therm Analy
- Apply TEMP from Thermal Analysis dialog box appears; enter 1 and 5 for Load step and substep no.; click on Browse.
- Fname Name of results file dialog box appears; search for and select WELD.rth file; click on $O K$.
- Click on OK in the Apply TEMP from Thermal Analysis dialog box.

Obtain the solution (SOLVE command) using the following menu path:
Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window
- Review status; if OK, close the Status Report Window; click on $O K$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review contours for normal stresses in the $z$-direction (PLNSOL command) using the following menu path:

Main Menu > General Postproc > Plot Results > Contour Plot > Nodal Solu

- Contour Nodal Solution Data dialog box appears; select Stress from the left list and $\mathbf{Z}$-direction $\mathbf{S Z}$ from the right list; click on $\mathbf{O K}$.
- The contour plot of normal stresses in the $z$-direction appears, as shown in Fig. 9.62.
- The thermomechanical solution may be repeated for any substep of interest. Figure 9.63 shows the contour plots of normal stress in the $z$ direction for substep 22 (last substep).


### 9.1.3 Radiation Analysis

Radiation is the transfer of thermal energy between two surfaces through electromagnetic waves. No medium is required for radiation heat transfer to take place. The transfer of thermal energy between two surfaces depends on the difference between the fourth powers of absolute temperatures along the surfaces. When radiation conditions are present, the problem is certain to be nonlinear. A radiation analysis requires specific knowledge of concepts that are not covered in this book. Therefore, it is highly recommended that the user have a good understanding of the radiation phenomenon before using ANSYS for radiation analyses.

Radiation analyses in ANSYS require the Stefan-Boltzmann constant, as well as the emissivity, view factor, and space temperature values for the surfaces involved. A radiation analysis is demonstrated by solving a simple problem using the Radiosity Solver method.


Fig. 9.62 Contour plot of the stresses in the $z$-direction due to thermal loading at load step 1 , substep 5.


Fig. 9.63 Contour plot of the stresses in the $z$-direction due to thermal loading at load step 1 , substep 22.

Two partial hollow circular regions are radiating to each other, as shown in Fig. 9.64. The emissivity of the outer surface of the inner region and inner surface of the outer region are 0.9 and 0.7 , respectively, while the inner surface of the inner region and outer surface of the outer region are maintained at temperatures of $1500^{\circ} \mathrm{F}$ and $100^{\circ} \mathrm{F}$, respectively. The space temperature is $70^{\circ} \mathrm{F}$. Both regions have a thermal conductivity of 0.1 . The goal is to obtain the steady-state temperature and heat flux variations.


Fig. 9.64 Schematic of the conical fin cross section.

## MODEL GENERATION

- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Element Types dialog box appears; click on Add.
- Select Thermal Solid in the left list and Quad 4 Node 55 on the right list; click on $\boldsymbol{O K}$.
- Exit from the Element Types dialog box by clicking on Close .
- Specify material properties (MP command) using the following menu path:


## Main Menu > Preprocessor > Material Props > Material Models

- Define Material Model Behavior dialog box appears. In the right window, successively double-click on Thermal, Conductivity, and Isotropic, which brings up another dialog box.
- Enter $\mathbf{0 . 1}$ for $\boldsymbol{K} \boldsymbol{X} \boldsymbol{X}$; click on $\boldsymbol{O K}$.
- Close the Define Material Model Behavior dialog box by using the following menu path:


## Material > Exit

- Create partial hollow circular areas (PCIRC command) using the following menu path:


## Main Menu > Preprocessor > Modeling > Create > Areas > Circle > By Dimensions

- Circular Area by Dimensions dialog box appears. Referring to Fig. 9.64 , enter 0.5 and 0.25 for RAD1 and RAD2, and 0 and 180 for THETA1 and THETA2; click on OK.
- Move the center of the Working Plane by 0.2 in the $x$-direction using the following menu path:

Utility Menu > WorkPlane > Offset WP by Increments

- Offset $W P$ menu appears; click on the $+X$ button four times and observe the Working Plane triad move in the Graphics Window. Also observe, toward the bottom of the Offset WP menu, that 0.2 appears next to Global $X=$.
- Exit from the Offset WP menu by clicking on $\boldsymbol{O K}$.
- Create the remaining partial hollow circular area (PCIRC command) using the following menu path:

```
Main Menu > Preprocessor > Modeling > Create > Areas > Circle > By Dimensions
```

- Circular Area by Dimensions dialog box appears. Referring to Fig. 9.64, enter 1 and 0.75 for RAD1 and RAD2 and 0 and 180 for THETA1 and THETA2; click on OK.
- The areas appear in the Graphics Window, as shown in Fig. 9.65.
- Turn line numbering on using the following menu path: Utility Menu > PlotCtrls > Numbering
- Plot Numbering Controls dialog box appears; click on the square box next to LINE Line numbers so that a checkmark appears. Click on OK. Line numbers appear, as shown in Fig. 9.66.
- Specify the number of elements along selected lines (LESIZE command) using the following menu path:


## Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Lines > Picked Lines

- Pick Menu appears; pick lines $1,3,5$, and 7 ; click on $\boldsymbol{O K}$.
- Element Sizes on Picked Lines dialog box appears; enter 40 in the text field for NDIV and remove the checkmark next to KYNDIV. Click on Apply.
- Pick Menu reappears; pick lines $2,4,6$, and 8 ; click on $O K$.
- In the Element Sizes on Picked Lines dialog box, enter 10 in the text field for NDIV; click on OK.
- Mesh the areas (AMESH command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 sided

- Pick Menu appears; click on Pick All.
- The mesh appears in the Graphics Window, as shown in Fig. 9.67.


Fig. 9.65 Area plot for the conical fin.


Fig. 9.66 Area plot with line numbers turned on.


Fig. 9.67 Mesh representing the conical fin.

## SOLUTION

- Apply radiation conditions on lines (SFL command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Thermal > Radiation > On Lines

- Pick Menu appears; pick line 1 (refer to Fig. 9.66); click on $O K$ in the Pick Menu.
- Apply RDSF on Lines dialog box appears; enter 0.9 for VALUE Emissivity and If for VALUE2 Enclosure number. Click on Apply.
- Pick Menu reappears; pick line 7; click on $\boldsymbol{O K}$ in the Pick Menu.
- In the Apply RDSF on Lines dialog box, enter 0.7 for VALUE Emissivity and $\boldsymbol{I}$ for VALUE2 Enclosure number. Click on OK.
- Specify the temperatures on lines (DL command) using the following menu path:

Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Thermal $>$ Temperature $>$ On Lines

- Pick Menu appears; pick line 3; click on $O K$ in the Pick Menu.
- Apply TEMP on Lines dialog box appears; highlight TEMP from the list and enter 1500 for VALUE Load TEMP value; click on Apply.
- Pick Menu reappears; pick line 5; and click on $O K$ in the Pick Menu.
- In the Apply TEMP on Lines dialog box enter $\mathbf{1 0 0}$ for VALUE Load TEMP value; click on OK.
- Define Radiosity Solver options (Stef, toffst, RADOPt, SPCTEMP commands) using the following menu path:


## Main Menu > Solution > Radiation Opts > Solution Opt

- Radiation Solution Options dialog box appears; enter 460 for [TOFFST] Temperature difference, 0.5 for Radiation flux relax. factor, 0.01 for Convergence tolerance, and 70 for Value (underneath Space option). Click on OK.
- Specify the time-related parameters (TIME, DELTIM commands) using the following menu path:


## Main Menu $>$ Solution > Load Step Opts $>$ Time/Frequenc $>$ Time - Time Step

- Time and Time Step Options dialog box appears. Enter 1 for [TIME] Time at end of load step, 0.5 for [DELTIM] Time step size, 0.1 for [DELTIM] Minimum time step size, and 1 for Maximum time step size. Click on $O K$.
- Specify the maximum number of equilibrium iterations (NEQIT command) using the following menu path:

Main Menu > Solution > Load Step Opts > Nonlinear > Equilibrium Iter

- Equilibrium Iterations dialog box appears. Enter 1000 for [NEQIT] No. of equilibrium iter; click on $O K$.
n Obtain the solution (SOLVE command) using the following menu path:
Main Menu >Solution >Solve >Current LS
- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window; click on $O K$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Review the temperature distribution (PLNSOL command) using the following menu path:

Main Menu > General Postproc $>$ Plot Results $>$ Contour Plot $>$ Nodal Solu

- Contour Nodal Solution Data dialog box appears; select DOF solution from the left list and Temperature TEMP from the right list; click on OK.
- Temperature contours appear in the Graphics Window, as shown in Fig. 9.68.
- Review flux vectors (PLVECT command) using the following menu path: Main Menu > General Postproc > Plot Results > Vector Plot > Predefined
- Vector Plot of Predefined Vectors dialog box appears; select Flux \& gradient from the left list and Thermal flux TF from the right list; click on $O K$.
- Flux vectors appear in the Graphics Window, as shown in Fig. 9.69.


### 9.2 Moisture Diffusion

The moisture diffusion phenomenon is an important issue because polymeric materials are being used more extensively in engineering applications. The moisture diffusion into a water permeable medium is governed by

$$
\begin{equation*}
\frac{\partial C}{\partial t}=D\left(\frac{\partial^{2} C}{\partial x^{2}}+\frac{\partial^{2} C}{\partial y^{2}}+\frac{\partial^{2} C}{\partial z^{2}}\right) \tag{9.3}
\end{equation*}
$$



Fig. 9.68 Contour plot of temperature.


Fig. 9.69 Vector plot of heat flux.
where $C$ is the moisture concentration, $D$ is the moisture diffusivity, and $t$ designates time. This equation is analogous to the heat diffusion (transient heat transfer) equation given by

$$
\begin{equation*}
\frac{\partial T}{\partial t}=\alpha\left(\frac{\partial^{2} T}{\partial x^{2}}+\frac{\partial^{2} T}{\partial y^{2}}+\frac{\partial^{2} T}{\partial z^{2}}\right) \tag{9.4}
\end{equation*}
$$

where $T$ is the temperature and $\alpha$ is the thermal diffusivity written in terms of thermal conductivity $\kappa$, specific heat $c$, and density $\rho$ as

$$
\begin{equation*}
\alpha=\frac{\kappa}{\rho c} \tag{9.5}
\end{equation*}
$$

Although Eq. (9.3) and (9.4) are analogous, the finite element formulation for heat diffusion cannot be directly used for solving moisture diffusion problems involving multiple dissimilar materials. This is because the moisture concentration, $C$, unlike the temperature, $T$, is not continuous along
material interfaces. However, when $C$ is normalized with respect to the saturated moisture concentration, $C_{s a t}$, this incompatibility is removed and the finite element formulation for heat diffusion can now be used for solving moisture diffusion problems. The normalized moisture concentration is called the wetness parameter and is written as

$$
\begin{equation*}
w=\frac{C}{C_{s a t}} \tag{9.6}
\end{equation*}
$$

As is clear from Eq. (9.4), thermal conductivity, $\kappa$, specific heat, $c$, and density, $\rho$, need to be known to solve a heat diffusion problem. After the introduction of the wetness parameter, in order to utilize the finite element formulation for heat diffusion for solving moisture diffusion problems, the user must use the correspondence table given in Table 9.8.

Once the solution is obtained, the weight of the moisture absorbed by an element, $W^{(e)}$, is computed by the product of average moisture concentration and the volume of the element in the form

$$
\begin{equation*}
W^{(e)}=\left(\frac{1}{N} \sum_{i=1}^{N} \omega_{i} \cdot C_{s a t}\right) \cdot V^{(e)} \tag{9.7}
\end{equation*}
$$

in which $N$ is the number of nodes per element and $V^{(e)}$ represents the volume of the element. The wetness at the $i^{\text {th }}$ node is indicated by $\omega_{i}$. The total weight of the absorbed moisture can then be found by simply summing the weight of the moisture absorbed by each element.

When simulating absorption conditions, the exposed surfaces of the moisture-absorbing materials are subjected to "wet" conditions, i.e., $T=1$. Similarly, in desorption simulations, those surfaces are subjected to "dry" conditions, $T=0$.

Once the solution is complete, the analyst can use Eq. (9.6) to obtain the moisture concentration, $C$, in each material.

Table 9.8 Correspondence table for thermal/moisture analogy.

| Property | Thermal | Moisture |
| :---: | :---: | :---: |
| Primary variable | Temperature, $T$ | Wetness, $\omega$ |
| Density | $\rho\left(\mathrm{kg} / \mathrm{m}^{3}\right)$ | 1 |
| Conductivity | $\kappa\left(\mathrm{W} / \mathrm{m} \cdot{ }^{\circ} \mathrm{C}\right)$ | $D \cdot C_{s a t}(\mathrm{~kg} / \mathrm{s} \cdot \mathrm{m})$ |
| Specific heat | $c\left(\mathrm{~J} / \mathrm{kg} \cdot{ }^{\circ} \mathrm{C}\right)$ | $C_{s a i}\left(\mathrm{~kg} / \mathrm{m}^{3}\right)$ |

The moisture diffusion simulation using ANSYS is demonstrated by considering a typical example from the electronics industry: an analysis of the moisture diffusion of an electronic package.

Consider an electronic package consisting of a silicon die, die-attach, copper, and molding compound, as shown in Fig. 9.70. The package is preconditioned at $85 \%$ relative humidity and a temperature of $85^{\circ} \mathrm{C}$ for one week ( 168 hr ) before being placed in a reflow oven at $225^{\circ} \mathrm{C}$ for 5 min . The moisture diffusivity during absorption and desorption and the saturated moisture concentration values of the molding compound and the die-attach are tabulated in Table 9.9. After applying the substitutions required for a thermal-moisture analogy, the values given in Table 9.10 are obtained. These are the material properties to be used in the ANSYS analysis. As noted in Table 9.11, the silicon die and copper do not absorb moisture and so appropriate material properties are used for these components. The geometry of the problem possesses half-symmetry; thus only half the package is simulated, with insulation along the symmetry axis.

The solution has two stages. First, absorption is simulated. During absorption, all the external surfaces of the moisture-absorbing materials are subjected to $T=1$. After the absorption simulation is complete, desorption is simulated, with boundary conditions $T=0$ along the same surfaces.


Fig. 9.70 Schematic of the electronic device undergoing moisture diffusion.

Table 9.9 Moisture diffusion related material properties of the constituent materials in the electronic package.

|  | Molding <br> Compound | Die-Attach |
| :--- | :---: | :---: |
| $D\left(\mathrm{~mm}^{2} / \mathrm{sec}\right)$ absorption | $7.43 \times 10^{-7}$ | $1.25 \times 10^{-5}$ |
| $D\left(\mathrm{~mm}^{2} / \mathrm{sec}\right)$ desorption | 0.18 | 0.35 |
| $C_{\text {sat }}\left(\mathrm{mg} / \mathrm{mm}^{3}\right)$ | $7.06 \times 10^{-3}$ | $6.20 \times 10^{-3}$ |

Table 9.10 Moisture properties converted to heat diffusion properties.

| a. Absorption |  |  |  |  |
| :--- | :---: | :---: | :---: | :---: |
|  | Molding <br> Compound | Die-Attach | Silicon Die | Copper |
| Material ref. no. | 1 | 2 | 3 | 4 |
| $\kappa(\mathrm{~kg} / \mathrm{hr} \cdot \mathrm{m})$ | $1.88 \times 10^{-5}$ | $2.79 \times 10^{-4}$ | $1 \times 10^{-11}$ | $1 \times 10^{-11}$ |
| $c\left(\mathrm{~kg} / \mathrm{m}^{3}\right)$ | 7060 | 6200 | 1 | 1 |
| $\rho$ | 1 | 1 | 1 | 1 |
| b. Desorption |  |  |  |  |
|  | Molding | Die-Attach |  |  |
| Material ref. no. | 5 | 6 |  |  |
| $\kappa(\mathrm{~kg} / \mathrm{hr} \cdot \mathrm{m})$ | $4.24 \times 10^{-3}$ | $9.32 \times 10^{-3}$ |  |  |
| $c\left(\mathrm{~kg} / \mathrm{m}^{3}\right)$ | 7060 | 6200 |  |  |
| $\rho$ | 1 | 1 |  |  |

Table 9.11 Coordinates defining the areas.

| Area <br> Number | X 1 | X 2 | Y 1 | Y 2 |
| :---: | :---: | :---: | :---: | :---: |
| 1 | 0 | 2.50 | 0 | 1.00 |
| 2 | 0 | 2.95 | 0 | 1.00 |
| 3 | 0 | 3.35 | 0 | 1.00 |
| 4 | 0 | 4.00 | 0 | 1.00 |
| 5 | 0 | 4.00 | 0 | 0.20 |
| 6 | 0 | 4.00 | 0 | 0.25 |
| 7 | 0 | 4.00 | 0 | 0.65 |

## model generation

- Define the element type (ET command) using the following menu path:

Main Menu > Preprocessor > Element Type > Add/Edit/Delete

- Element Types dialog box appears; click on Add.
- Select Thermal Solid in the left list and Quad 4 Node 55 on the right list; click on $\boldsymbol{O K}$.
- Click on Close.
- Specify material properties (MP command) using the following menu path:

Main Menu > Preprocessor > Material Props $>$ Material Models

- Define Material Model Behavior dialog box appears. Specify conductivity by successively double-clicking on Thermal, Conductivity, and Isotropic; enter $1.88 E-5$ in the newly appeared dialog box; click on OK.
- Specify density by successively double-clicking on Thermal and Specific Heat, which brings up another dialog box. Type 7060, click on OK.
- Specify specific heat by successively double-clicking on Thermal and Density; enter $\boldsymbol{I}$ in the newly appeared dialog box; click on $\mathbf{O K}$.
- Add new material model using the following menu path:

Material > New Model

- Repeat the procedure for the remaining materials ( 2 through 6 ) by referring to Table 9.10.
- When finished, close the Define Material Model Behavior dialog box by using the following menu path:

Material > Exit

- Create rectangular areas (RECTNG command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Modeling $>$ Create $>$ Areas $>$ Rectangle $>$ By Dimensions

- Create Rectangle by Dimensions dialog box appears. Referring to Table 9.11, enter $\boldsymbol{0}$ and $\mathbf{2 . 5 E - 3}$ for $\boldsymbol{X 1}$ and $\mathbf{X 2}$ and $\boldsymbol{0}$ and $\mathbf{1 E - 3}$ for $\mathbf{Y 1}$ and $\boldsymbol{Y}$; click on Apply. Note that the units used in the analysis are in meters while the values given in Table 9.11 are in millimeters.
- Repeat the procedure for the remaining areas ( 2 through 7). When creating Area 7 , click on $O K$ after entering the coordinates.
- Overlap the areas (AOVLAP command) using the following menu path:

Main Menu > Preprocessor > Modeling > Operate > Booleans > Overlap > Areas
Pick Menu appears; click on Pick All button.

- Compress entity numbers (NUMCMP command) using the following menu path:
Main Menu > Preprocessor > Numbering Ctrls > Compress Numbers
- Compress Numbers dialog box appears; select $A l l$ from the pull-down menu; click on $O K$.
- Specify the global element size (ESIZE command) using the following menu path:

Main Menu > Preprocessor > Meshing > Size Cntrls > ManualSize > Global > Size

- Global Element Sizes dialog box appears; enter 4E-5 for SIZE Element edge length; click on OK.
- Mesh the areas (AMESH command) using the following menu path:

Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Mapped $>3$ or 4 sided

- Pick Menu appears; click on Pick All.
- At this point, all the elements have Material Reference Number 1. Attributes can be changed after the elements are created. For this purpose, areas are selected and then the elements that are attached to the selected areas are selected. Finally, elements are modified to have the correct attributes. The correspondence between the areas and material numbers are given in Table 9.12. Figure 9.71 shows the area plot with area numbers printed. Plot areas (APLOT command) using the following menu path:

Utility Menu > Plot > Areas

- Turn area numbers on (/PNUM command) using the following menu path:
Utility Menu >PlotCtrls > Numbering
- Plot Numbering Controls dialog box appears; place a checkmark next to AREA Area numbers by clicking on the box; select Numbers only from the pull-down menu next to [/NUM] Numbering shown with. Click on $\boldsymbol{O K}$. Areas appear with their numbers printed, as shown in Fig. 9.71 .

Table 9.12 Areas corresponding to specific materials.

| Material | Area Numbers |
| :---: | :---: |
| Molding compound | $2,4,5,6,8,9,11,13,14,15,16$ |
| Die-attach | 10 |
| Silicon die | 12 |
| Copper | $1,3,7$ |


| A2 | A4 | A6 | A8 |
| :---: | :---: | :---: | :---: | :---: |
| A12 | A14 | A16 | A11 |
| A10 | A13 | A15 | A9 |
| A1 | A3 | A5 | A7 |

Fig. 9.71 Area plot with area numbers turned on.

- Select areas (ASEL command) using the following menu path: Utility Menu > Select > Entities
- Select Entities dialog box appears; select Areas from the first pulldown menu; click on $\boldsymbol{O K}$.
- Pick Menu appears; pick area 10 ; click on $\boldsymbol{O K}$.
- Now, select the elements that are attached to the selected areas (ESLA command) using the following menu path:

Utility Menu > Select > Entities

- Select Entities dialog box appears; select Elements from the first pulldown menu; select Attached to from the second pull-down menu. Click on the Areas radio-button; click on $\mathbf{O K}$.
- Modify the attributes of the selected set of elements (EMODIF command) using the following menu path:

Main Menu > Preprocessor > Modeling > Move / Modify > Elements > Modify Attrib

- Pick Menu appears; click on Pick All, which brings up the Modify Elem Attributes dialog box.
- Select Material MAT from the pull-down menu and enter 2 in the $\mathbf{I I}$ New attribute number field; click on $O K$.
- Repeat this procedure for area 12 (material reference number 3), and areas 1,3 , and 7 (material reference number 4).
- When finished, select everything (aLLSEL command) using the following menu path:

Utility Menu > Select > Everything

- Plot the elements (EPLOT command) using the following menu path: Utility Menu > Plot > Elements
- Plot the elements with different colors based on their material reference numbers (/PNUM command) using the following menu path:

Utility Menu $>$ PlotCtris $>$ Numbering

- Plot Numbering Controls dialog box appears; remove the checkmark next to AREA Area numbers by clicking on the box. Select Material Numbers for Elem / Attrib numbering pull-down menu; select Colors only from the pull-down menu next to [/NUM] Numbering shown with; click on $O K$. When elements are plotted, they appear in different colors based on their material reference numbers, as shown in Fig. 9.72.


## SOLUTION

- Declare the new analysis to be a transient analysis (ANTYPE command) using the following menu path:

Main Menu > Solution > Analysis Type > New Analysis

- New Analysis dialog box appears; click on the Transient radio-button; click on $\mathbf{O K}$.
- Transient Analysis dialog box appears; click on $\mathbf{O K}$.
- Apply the initial condition on the nodes (IC command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Initial Condit'n > Define

- Pick Menu appears; click on Pick All in Pick Menu.
- Define Initial Conditions dialog box appears; select TEMP from the pull-down menu; enter 0 for VALUE Initial value of DOF; click on OK.

Apply temperature (wetness) boundary conditions on the lines (SFL command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Thermal > Temperature >On Lines

- Pick Меnu appears; pick the exterior lines that are shown in bold in Fig. 9.73; click on OK in the Pick Menu.
- Apply TEMP on Lines dialog box appears; select TEMP from the list and enter 1 for VALUE Load TEMP value; click on OK.
- Specify solution controls using the following menu path:

Main Menu > Solution > Analysis Type >Sol'n Controls

- Solution Controls dialog box appears, which has five tabs. In the Basic tab, within the Time Control field, enter 168 for Time at end of loadstep.
- Click on the Time increment radio-button; enter I for Time step size.
- In the Write Items to Results File field, select Write every substep from the Frequency pull-down menu; click on OK.


Fig. 9.72 Mesh representing the electronic device.


Fig. 9.73 Bold lines represent the surfaces of moisture diffusion.

- Obtain the solution (SOLVE command) using the following menu path:

Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window and click on $\mathbf{O K}$ in the Confirmation Window.
- Wait until ANSYS responds with Solution is done!

The solution for pre-conditioning is complete. Next, the solution for the reflow process is obtained. The moisture diffusivity values for the molding compound and the die-attach are different for pre-conditioning (absorption) and reflow (desorption). For this purpose, the material attributes of the molding compound and die-attach need to be modified to be 5 and 6 , respectively.

- Select elements corresponding to the molding compound (ESEL command) using the following menu path:

Utility Menu > Select Entities

- Select Entities dialog box appears; select Elements from the first pulldown menu; select By Attributes from the second pull-down menu. Click on the Material num radio-button and type 1 in the test field; click on $\boldsymbol{O K}$.

■ Modify the attributes of the selected set of elements (EMODIF command) using the following menu path:

Main Menu > Solution > Other > Change Mat Props > Change Mat Num

- Change Material Number dialog box appears; type 5 in the Mat New material number text field and type $A L L$ in the ELEM Element no. to be modified text field; click on OK.
- Repeat the same procedure for die-attach elements; first select elements with material attribute number 2 ; then modify their material attribute number to be 6 .
- When finished, select everything (ALLSEL command) using the following menu path:

Utility Menu > Seiect > Everything

- Apply temperature (wetness) boundary conditions on lines (SFL command) using the following menu path:

Main Menu > Solution > Define Loads > Apply > Thermal > Temperature > On Lines

- Pick Menu appears; pick the exterior lines shown in Fig. 9.73; click on OK in the Pick Menu.
- Apply TEMP on Lines dialog box appears; select TEMP from the list and enter 0 for VALUE Load TEMP value; click on OK.
- Specify solution controls using the following menu path:

Main Menu > Solution > Analysis Type > Sol'n Controls

- Solution Controls dialog box appears, which has five tabs. In the Basic tab, within the Time Control field, enter 168+5/60 for Time at end of loadstep.
- Click on the Time increment radio-button; enter 5/60/20 for Time step size.
- In the Write Items to Results File field, select Write every substep from the Frequency pull-down menu; click on $\boldsymbol{O K}$.
- Obtain the solution (SOLVE command) using the following menu path:

Main Menu > Solution > Solve > Current LS

- Confirmation Window appears along with Status Report Window.
- Review status; if OK, close the Status Report Window and click on OK in Confirmation Window.
- Wait until ANSYS responds with Solution is done!


## POSTPROCESSING

- Read the results for the last time point using the following menu path:
- Plot temperature (wetness) contours (PLNSOL command) at the last time point using the following menu path:
Main Menu > General Postproc > Plot Results > Contour Plot > Nodal Solu
- Contour Nodal Solution Data dialog box appears; click on OK. The contour plot appears, as shown in Fig. 9.74.
- Plot flux vector contours (PLVECT command) at the last time point using the following menu path:

```
Main Menu > General Postproc > Plot Results > Vector Plot > Predefined
```

- Vector Plot of Predefined Vectors dialog box appears; click on $O K$. The vector plot appears, as shown in Fig. 9.75.
m Obtain an animation of the temperature (wetness) over time using the following menu path:

Utility Menu $>$ PlotCtrls $>$ Animate $>$ Over Time
■ Animate Over Time dialog box appears; enter 100 for Number of animation frames, select Load Step Range radio-button and enter 1 and 2 for Range Minimum, Maximum. Click on OK; wait until the Animation Controls Window appears.

- Obtain the moisture content as a function of time. This portion of the postprocessing is performed using the ANSYS Parametric Design Language (APDL) as discussed in substantial detail in Chap. 7. It is assumed that the results are stored in a file named WET.RTH. The input file (written using APDL) is given below. There are two DO loops: one over the substeps and the other over the elements. During each of the loops over the elements, each element's area, saturated moisture concentration, and wetness parameter are used to calculate the element's moisture content in milligrams. Since this is a two-dimensional idealization, the package is assumed to have a unit thickness in the third direction. The results (time and total moisture content in the package) are written to two different text files, i.e., ABS.OUT for absorption and DES.OUT for desorption. Figures 9.76 and 9.77 show the variation of total moisture content in the package against time during absorption and desorption, respectively.


Fig. 9.74 Contour plot of wetness.


Fig. 9.75 Vector plot of flux.


Fig. 9.76 Moisture content vs. time during absorption.


Fig. 9.77 Moisture content vs. time during desorption.
/POST1
FILE, 'WET', 'RTH', '.' CSAT5 $=7.06 \mathrm{E}-3$
CSAT6 $=6.20 \mathrm{E}-3$
CSAT3 $=0$
CSAT4 $=0$
*GET, NUMEL, ELEM, 0, COUNT
SET, FIRST
*DO, J, 1, 188
ETABLE, , TEMP
*GET, MINEL, ELEM, 0, NUM, MIN
*GET, TIM, ACTIVE, 0, SET, TIME
SUM=0
*DO, I, 1, NUMEL
*GET, EA, ELEM, MINEL, AREA
*GET, EM, ELEM, MINEL, ATTR, MAT
*GET, EW, ETAB, 1, ELEM, MINEL
$\mathrm{EC}=1 \mathrm{E} 6 * \mathrm{EA}^{*} \mathrm{EW}{ }^{*} \mathrm{CSAT} \% \mathrm{EM} \%$
SUM $=$ SUM + EC
MINEL=ELNEXT (MINEL)

* ENDDO
*IF, J, LE, 168, THEN
/ OUTPUT, ABS, OUT, , APPEND

```
! ENTER POSTPROCESSOR
! DECLARE RESULTS FILE
!C_sat FOR MOLDING COMPOUND
!C_sat FOR DIE-ATTACH
! C_sat FOR DIE
!C_sat FOR COPPER
! OBTAIN NUMBER OF ELEMENTS
! READ 1 }\mp@subsup{}{}{\mathrm{ ST }}\mathrm{ RESULTS SET
! (SUBSTEP 1)
! LOOP OVER SUBSTEPS (TOTAL
! 188)
! STORE WETNESS IN ELEMENT
! TABLE
! OBTAIN MINIMUM ELEMENT
! NUMBER
! OBTAIN CURRENT TIME
! INITIALIZE TOTAL MOISTURE
! MASS
! LOOP OVER ELEMENTS
! OBTAIN ELEMENT AREA
! OBTAIN ELEMENT MATERIAL
! NUMBER
! OBTAIN ELEMENT WETNESS
! CALCULATE MOISTURE MASS (mg)
! UPDATE TOTAL MOISTURE MASS
! MOVE ON TO NEXT ELEMENT
! END LOOP OVER ELEMENTS
! IF TRUE - ABSORPTION
! REDIRECT OUTPUT TO FILE
! ABS.OUT
```

| *VWRITE, TIM, SUM | ! WRITE TIME AND MOISTURE MASS ! TO FILE |
| :---: | :---: |
| (E16.8.5X, E16.8) | ! FORMAT FOR *VWRITE COMMAND |
| /OUTPUT | ! REDIRECT OUTPUT TO OUTPUT ! WINDOW |
| *ELSEIF, J, GT, 168, THEN | ! IF TRUE - DESORPTION |
| / OUTPUT, DES, OUT, , APPEND | ```! REDIRECT OUTPUT TO FILE ! DES.OUT``` |
| *VWRITE, TIM, SUM | ! WRITE TIME AND MOISTURE MASS ! TO FILE |
| (E16.8, 5X, E16.8) | ! FORMAT FOR *VWRITE COMMAND |
| / OUTPUT | ! REDIRECT OUTPUT TO OUTPUT ! WINDOW |
| * ENDIF | ! |
| *IF, J, LT, 188, THEN | $!$ |
| SET, , , , , , J + 1 | ! READ NEXT RESULTS SET |
| * ENDIF | ! |
| * ENDDO | ! END LOOP OVER SUBSTEPS |

## Chapter 10

## NONLINEAR STRUCTURAL ANALYSIS

The nonlinear load-displacement relationship-the stress-strain relationship with a nonlinear function of stress, strain, and/or time; changes in geometry due to large displacements; irreversible structural behavior upon removal of the external loads; change in boundary conditions such as a change in the contact area and the influence of loading sequence on the behavior of the structure-requires a nonlinear structural analysis. The structural nonlinearities can be classified as geometric nonlinearity, material nonlinearity, and contact or boundary nonlinearity.

The governing equations concerning large deformations are nonlinear with respect to displacements and velocities. The material behavior can be linear or nonlinear, and the boundary conditions can also exhibit nonlinearity. Geometrical nonlinearity arising from large deformations is associated with the necessity to distinguish between the coordinates of the initial and final states of deformation, and also with the necessity to use the complete expressions for the strain components. The material can exhibit either timedependent or time-independent nonlinear behavior. Nonlinearity due to boundary conditions emerges from a nonlinear relationship between the external forces and the boundary displacements. The presence of contact conditions also leads to a nonlinear structural analysis because the extent of the contact region and the contact stresses are not known a priori.

The solution to the nonlinear governing equations can be achieved through an incremental approach. The incremental form of the governing equations can be written as

$$
\begin{equation*}
\mathbf{K}(\mathbf{u}) \Delta \mathbf{u}=\Delta \mathbf{P} \tag{10.1}
\end{equation*}
$$

in which $\Delta \mathbf{u}$ and $\Delta \mathbf{P}$ represent the unknown incremental displacement vector and the known incremental applied load vector, respectively. The solution is constructed by taking a series of linear steps in the appropriate direction in order to closely approximate the exact solution. Depending on the nature of the nonlinearity, the magnitude of each step and its direction may involve several iterations. The computational algorithms and the
associated parameters must be chosen with extreme care. The solution to nonlinear problems may not be unique.

When solving nonlinear problems, ANSYS uses the Newton-Raphson (N-R) method, which involves an iterative procedure. This method starts with a trial (assumed) solution, $\mathbf{u}=\mathbf{u}_{i}$, to determine the magnitude of the next step (increment), $\Delta \mathbf{u}_{i}=\mathbf{K}^{-1}\left(\mathbf{u}_{i}\right) \Delta \mathbf{P}$, and the corresponding out-of-balance load vector, $\Delta \mathbf{R}_{i}=\Delta \mathbf{P}-\mathbf{K}\left(\mathbf{u}_{i}\right) \Delta \mathbf{u}_{i}$, which is the difference between the applied loads and the loads evaluated based on the assumed solution. In order to satisfy the equilibrium conditions exactly, the out-of-balance load vector must be zero. However, as the nonlinear equilibrium conditions are solved approximately, a tolerance is introduced for the out-of-balance load vector in order to terminate the solution procedure. In each iteration, the N-R method computes the out-of-balance load vector and checks for convergence based on the specified tolerance. If the convergence criterion is not satisfied, the trial solution is updated as $\mathbf{u}_{i+1}=\mathbf{u}_{i}+\Delta \mathbf{u}_{i}$ based on the calculated incremental displacements, and the next incremental solution vector is determined as $\Delta \mathbf{u}_{i+1}=\mathbf{K}^{-1}\left(\mathbf{u}_{i+1}\right) \Delta \mathbf{P}$ leading to the computation of the new out-of-balance load vector $\Delta \mathbf{R}_{i+1}=\Delta \mathbf{P}-\mathbf{K}\left(\mathbf{u}_{i+1}\right) \Delta \mathbf{u}_{i+1}$; this procedure is repeated until convergence is accomplished.

Several methods for improving the convergence (or convergence rate) are available in ANSYS. These include automatic time stepping, a bisection method, and line search algorithms. The user may choose to have full control or let ANSYS choose the options.

In a nonlinear solution in ANSYS, there are three distinct levels: (1) Load Steps, (2) Substeps, and (3) Equilibrium Iterations. The number of load steps is specified by the user. Different load steps must be used if the loading on the structure changes abruptly. The use of load steps also becomes necessary if the response of the structure at specific points in time is desired. A solution within each load step is obtained by applying the load incrementally in substeps. Within each substep, several equilibrium iterations are performed until convergence is accomplished, after which ANSYS proceeds to the next substep. As the number of substeps used increases, the accuracy of the solution improves. However, this also means that more computational time is being used. ANSYS offers the Automatic Time Stepping feature to optimize the task of obtaining a solution with acceptable accuracy in a reasonable amount of time. The automatic time stepping feature decides on the number and size of substeps within load steps. When using automatic time stepping, if a solution fails to converge within a
substep, the bisection method is activated, which restarts the solution from the last converged substep.

The ANSYS program has default values for all of the nonlinear solution controls, including the convergence options. The SOLCONTROL command is used to turn these defaults on or off. The help page for the SOLCONTROL command provides a comprehensive list of the default values of nonlinear analysis settings when solution controls are on (SOLCONTROL, ON), which is the default setting. It is also possible to modify specific controls while leaving the rest for ANSYS to assign. Some of the commonly used commands for modifying/specifying nonlinear analysis settings with brief descriptions are:

AUTOTS Command: Turns automatic time stepping on or off.
DELTIM Command: Specifies time step size and/or minimum and maximum time step sizes to be used within a load step.
nSUBST Command: Specifies number of substeps and/or minimum and maximum number of substeps to be used within a load step.

NEQIT Command: Specifies maximum number of equilibrium iterations within a substep. If this number is reached with no converged solution, and if automatic time stepping is on, then ANSYS employs the bisection method to achieve convergence. Otherwise, the solution is terminated.

KBC Command: Specifies whether the loads are interpolated (ramped) for each substep from the values of the previous load step to the values of the current load step.

EQSLV Command: Specifies the type of solver to be used to solve the matrix system of equations. By default, it is the Sparse Solver, however, there are several other solvers available that may be more efficient for the particular problem being solved.

CNVTOL Command: Specifies convergence tolerance values for the nonlinear analyses.

NROPT Command: Specifies which type of Newton-Raphson method is used in the solution.

LNSRCH Command: Specifies whether a line search is to be used with the Newton-Raphson method in the solution.

PRED Command: Specifies whether a predictor algorithm is to be used in the solution.

## ARCLEN Command: Toggles the arc-length method on or off.

SSTIF Command: The stiffness of certain materials increases with the increased stress levels within the structure (e.g., cables and membranes).
This command toggles the stress stiffening effects on or off.
TIMINT Command: Toggles the transient effects on or off.
OUTRES Command: Specifies the amount and frequency of the data saved in the results file. By default, results associated with the last substep of each load step are written in the results file.

The commands described above require special attention, which may be crucial to the success of the analysis. There are no "golden standard" values for time step sizes, the number of equilibrium iterations, or the number of substeps. The user accumulates knowledge on the use of these features with every new analysis. It is highly recommended that the user consult the ANSYS Help pages on nonlinear analysis and individual commands, which provide detailed guidelines. Some general suggestions on achieving success in nonlinear analyses are as follows:

Nonlinear analyses require more computational time. Therefore, when solving nonlinear static problems, it may be helpful to solve a preliminary version of the problem with no nonlinearities. The results from the linear solution may indicate mistakes in modeling, meshing, and application of boundary conditions in a shorter time frame. Also, the linear solution provides information about the regions where high stress gradients are expected, thus guiding the user to modify the mesh (make it more refined) in those regions.

In nonlinear analyses, it is important to utilize all possible simplifications in order to improve convergence and reduce the computational cost. For example, if the problem can be simplified as a plane stress or plane strain idealization, then the user should take advantage of this opportunity.

Reading the contents of the Output Window and the Error File (jobname. err) is crucial in finding the specific reason why the solution does not converge.

Another important consideration in dealing with nonlinear problems is the path dependency of the solution. When all the materials in a problem exhibit linear behavior, the order in which the loads are applied does not make any difference in the results. However, when a nonlinear material behavior is
present, results obtained by applying the same set of loads in different orders may differ from each other.

Detailed step-by-step instructions for numerous example problems are provided in Chap. 6, 8, and 9. The command line equivalents of each of these example problems, written in the ANSYS Parametric Design Language (APDL), are included in the accompanying CD-ROM. The use of the APDL is described in Chap. 7.

In this chapter, the nonlinear structural analyses arising from (i) geometric nonlinearity, (ii) material nonlinearity, and (iii) contact conditions are considered in order to demonstrate the nonlinear features of ANSYS. However, APDL is chosen to be the main method of interacting with ANSYS because of its versatility and efficiency. Explanations are included in each command line after an exclamation mark (!). Step-by-step instructions are sporadically given when they are considered to be beneficial to the user. It is highly recommended that the user have a good understanding of APDL before delving into this chapter.

### 10.1 Geometric Nonlinearity

Geometric nonlinearities arise from the presence of large strain, small strains but finite displacements and/or rotations, and loss of structural stability. Large strains, over $5 \%$ may occur in rubber structures and metal forming. Slender structures such as bars and thin plates may experience large displacements and rotations with small strains. Initially stressed structures with small strains and displacements may undergo a loss of stability by buckling.

Two problems are considered. The first problem involves a thin cantilever plate subjected to a point load at one of the free corners. Because the plate is thin, the resulting displacement components are in comparable order to its geometric dimensions, thus the geometry changes. This requires the stiffness matrix to be modified by accounting for the changes in the geometry. Results are compared to the solution of the same problem obtained by disregarding the nonlinear geometry effects.

The second problem involves a composite plate with a circular hole subjected to compression. As the applied loading increases, the plate is expected to buckle. Eigenvalue Buckling Analysis is one of the methods that could be used to solve this problem (as explained in Chap. 8). However, Eigenvalue Buckling Analysis evaluates only the buckling load; it does not solve for the events after buckling occurs. Alternatively, nonlinear geometry
effects are turned on and post-buckling behavior is evaluated along with the buckling load.

### 10.1.1 Large Deformation Analysis of a Plate

Consider the cantilever plate with a transverse force at one corner shown in Fig. 10.1. The plate has a length, width, and thickness of $40 \mathrm{~m}, 30 \mathrm{~m}$, and 0.4 m , respectively. Its elastic modulus is 120 MPa , and the Poisson's ratio is 0.3 . The maximum applied load of 40 kN is reached in five equal increments. The nonlinear geometry option is used in the ANSYS solution. This is achieved by writing Load Step Files for each increment, and obtaining the solution from these files (LSSOLVE command). The goal is to find the displacement components as the applied load increases, at points A and B shown in Fig. 10.1.

## MODEL GENERATION

Element type shelu181 is used in the analysis. The number of element divisions on all of the lines is specified as 20, and mapped meshing is utilized. The following command input is used for model generation:

```
/PREP7 ! ENTER PREPROCESSOR
ET,1,181
R,1,0.4
MP,EX,1,1.2E8
MP,NUXY,1,0.3
K,1,0
K,2,40
K,3,40,30
K,4,0,30
A,1,2,3,4
LESIZE,ALL, ,,20
MSHKEY,1
AMESH, ALL
FINISH
```

```
SPECIFY ELEMENT TYPE AS SHELL 181
```

SPECIFY ELEMENT TYPE AS SHELL 181
SPECIFY PLATE THICKNESS
SPECIFY PLATE THICKNESS
SPECIFY ELASTIC MODULUS
SPECIFY ELASTIC MODULUS
SPECIFY POISSON'S RATIO
SPECIFY POISSON'S RATIO
CREATE KEYPOINTS
CREATE KEYPOINTS
CREATE AREA THROUGH KEYPOINTS
CREATE AREA THROUGH KEYPOINTS
SPECIFY NO. OF DIVISIONS ON LINES
SPECIFY NO. OF DIVISIONS ON LINES
ENFORCE MAPPED MESHING
ENFORCE MAPPED MESHING
MESH THE AREA
MESH THE AREA
EXIT PREPROCESSOR

```
EXIT PREPROCESSOR
```



Fig. 10.1 Cantilever plate with a transverse force at one corner.

## SOLUTION

Nonlinear geometry effects are turned on using the NLGEOM command. The maximum number of equilibrium iterations is specified as 1000 using the NEQIT command. After the specification of displacement constraints, the transverse load is specified in increments of 8000 N . After application of each increment, a load step file is written to the Working Directory (with the naming convention: file.s01, file.s02, ...) using the LSWRITE command. When the LSWRITE command is issued, ANSYS writes all of the specified boundary conditions to the load step file. When finished with the last load step ( $40,000 \mathrm{~N}$ ), the LSSOLVE command is issued to start the solution by reading the boundary conditions from load step files 1 through 5 . The following command input is used for the solution:

| / SOLU | ENTER SOLUTION PROCESSOR |
| :---: | :---: |
| ANTYPE, STATIC | ! SPECIFY ANALYSIS TO BE STATIC |
| NLGEOM, ON | ! TURN NONLINEAR GEOMETRY |
|  | $!$ EFFECTS ON |
| NEQIT, 1000 | ! SPECIFY MAX. \# OF EQUILIBRIUM |
|  | ! ITERATIONS |
| OUTRES, ALL, ALL | ! WRITE ALL SOLUTION ITEMS FOR |
|  | $!$ EVERY SUBSTEP |
| NSEL, S, LOC , X, 0 | ! SELECT NODES AT $\mathrm{X}=0$ |
| D, ALL, ALL | $!$ CONSTRAIN ALL DOFS AT SELECTED |
|  | ! NODES |
| ALLSEL | ! SELECT EVERYTHING |
|  | ! START FIRST LOAD INCREMENT |
| NSEL, S, LOC, X, 40 | SELECT NODES AT $X=40$ |
| NSEL, R, LOC, Y, 0 | $!$ SELECT NODES AT Y $=0$ FROM THE |
|  | SELECTED SET |
| F, ALL, FZ, -8000 | ! SPECIFY FORCE IN NEG. Z-DIR AT |
|  | SELECTED NODES |
| ALLSEL | ! SELECT EVERYTHING |
| LSWRITE | ! WRITE LOAD STEP FILE 1 |
|  | ! SECOND LOAD INCREMENT |
| NSEL, S, LOC, X, 40 | SELECT NODES AT X $=40$ |
| NSEL, R, LOC, Y, 0 | ! SELECT NODES AT Y $=0$ FROM THE |
|  | ! SELECTED SET |
| F, ALL, FZ, -16000 | ! SPECIFY FORCE IN NEG. Z-DIR AT |
|  | ! SELECTED NODES |
| ALLSEL | SELECT EVERYTHING |
| LSWRITE | ! WRITE LOAD STEP FILE 2 |
|  | $!$ THIRD LOAD INCREMENT |
| NSEL, S, LOC, X, 40 | SELECT NODES AT X $=40$ |
| NSEL, R, LOC, Y, 0 | SELECT NODES AT $Y=0$ FROM THE |
|  | $!$ SELECTED SET |
| F, ALL , FZ, -24000 | SPECIFY FORCE IN NEG. Z-DIR AT |
|  | $!$ SELECTED NODES |
| ALLSEL | ! SELECT EVERYTHING |
| LSWRITE | ! WRITE LOAD STEP FILE 3 |
|  | ! FOURTH LOAD INCREMENT |
| NSEL, S, LOC , X, 40 | ! SELECT NODES AT X $=40$ |


| NSEL, R, LOC, Y, 0 | SELECT NODES AT $Y=0$ FROM THE SELECTED SET |
| :---: | :---: |
| F،ALL, FZ, -32000 | SPECIFY FORCE IN NEG. Z-DIR AT SELECTED NODES |
| ALLSEL | SELECT EVERYTHING |
| LSWRITE | WRITE LOAD STEP FILE 4 |
|  | FIFTH LOAD INCREMENT |
| NSEL, S, LOC, X, 40 | SELECT NODES AT X $=40$ |
| NSEL, R, LOC, Y, 0 | SELECT NODES AT Y $=0$ FROM THE |
|  | SELECTED SET |
| F, ALL, FZ, -40000 | SPECIFY FORCE IN NEG. Z-DIR AT |
|  | S SELECTED NODES |
| ALLSEL | ! SELECT EVERYTHING |
| LSWRITE | [ WRITE LOAD STEP FILE 5 |
| LSSOLVE, 1,5,1 | ! SOLVE FROM LOAD STEP FILES |
| FINISH | EXIT SOLUTION PROCESSOR |

## POSTPROCESSING

In the postprocessing, node numbers for the nodes at points A and B are stored in parameters NA and NB, respectively. A do loop is used for extraction of the results data at different results sets corresponding to the load steps. The commands /OUTPUT and *VWRITE are used for redirecting output and writing parameters to external files. The use of shortcuts for *GET functions $\mathbf{U X}(N), \mathbb{U Y}(N)$, and $\mathbf{U Z}(N)$, is also demonstrated. These functions retrieve the $x$-, $y$-, and $z$-displacements of node $\boldsymbol{N}$. The following command input is used for postprocessing:

```
/POST1
NSEL, S, LOC, X, 40
NSEL, R, LOC, Y, O
*GET , NA, NODE, 0,NUM, MIN
NSEL, S,LOC,X,40
NSEL, R,LOC,Y, 30
*GET, NB,NODE, 0, NUM, MIN
ALLSEL
*DO,I,1,5
SET,I
/OUTPUT,ADISP,OUT, ,APPEND
*VWRITE,UX(NA),UY(NA),UZ (NA)
(E16.8,5X, E16.8,5X,E16.8,5X)
/OUTPUT
/OUTPUT,BDISP,OUT, ,APPEND
*VWRITE,UX (NB),UY (NB) ,UZ (NB)
(E16.8,5X, E16.8,5X, E16.8,5X)
/OUTPUT
* ENDDO
```

! ENTER POSTPROCESSOR
! SELECT NODES AT X $=40$
! RESELECT NODE AT $Y=0$
! STORE NODE \# OF PT A
! INTO NA
: SELECT NODES AT X $=40$
! RESELECT NODE AT $Y=30$
! STORE NODE \# OF PT B
! INTO NB
! SELECT EVERYTHING
! START DO LOOP
! SET RESULTS TO LOAD STEP I
! REDIRECT OUTPUT TO FILE
! ADISP.OUT
! WRITE DISPLACEMENTS TO FILE
! FORMAT STATEMENT
! REDIRECT OUTPUT TO OUTPUT
! WINDOW
! REDIRECT OUTPUT TO FILE
! BDISP. OUT
! WRITE DISPLACEMENTS TO FILE
! FORMAT STATEMENT
! REDIRECT OUTPUT TO
! OUTPUT WINDOW
! END DO LOOP

After execution of the command input given above, the $x$-, $y$-, and $z$ displacements at points A and B are written to files ADISP.OUT and BDISP.OUT, respectively. Figure 10.2 shows the variation of displacement components as the load increases incrementally. Also plotted in Fig. 10.2 are the $z$-displacements of points A and B obtained by disregarding geometric nonlinearity (indicated in the legend as "linear"). Displacements in the $x$ - and $y$-directions are identically zero for this case.

### 10.1.2 Post-buckling Analysis of a Plate with a Hole

Consider a 9.5 -in-square composite plate with a circular hole of radius 1.5 in, as illustrated in Fig. 10.3. The laminate lay-up is $[ \pm 30]_{3 s}$, with a total of 12 layers, symmetric with respect to the mid-plane, and the orientation of the layers alternates between $30^{\circ}$ and $-30^{\circ}$. Each layer has moduli of $E_{L}=18.5 \times 10^{6} \mathrm{psi}, E_{T}=1.6 \times 10^{6} \mathrm{psi}$, and $G_{L T}=0.832 \times 10^{6} \mathrm{psi}$, and a Poisson's ratio of $v_{L T}=0.35$. Each layer is 0.01 in thick, resulting in a total laminate thickness of 0.12 in .

Along the right edge of the laminate, an axial concentrated load of $12,000 \mathrm{lb}$ is introduced through a rigid end. This type of load introduction, requiring the $x$-displacement to be uniform, is enforced by coupled degrees of freedom, as explained in Chap. 11. All degrees of freedom (displacements and rotations) are constrained along the left edge. Along the horizontal edges, inplane displacements and rotations about these edges are permitted. In order to trigger the nonlinear response, a sinusoidal imperfection with an amplitude of $1 \%$ of the total laminate thickness is used as follows:

$$
\begin{equation*}
z=0.0012 \sin \left(\frac{\pi x}{9.5}\right) \sin \left(\frac{\pi y}{9.5}\right) \tag{10.2}
\end{equation*}
$$

The goal is to obtain the variations of the $z$-displacement at points $\mathrm{A}(4.75$, $6.25,0)$ and $\mathrm{B}(6.25,4.75,0)$, and the $x$-displacement at point $\mathrm{C}(9.5,4.75$, 0 ) as the applied load increases (points $\mathrm{A}, \mathrm{B}$, and C are indicated in Fig. 10.3).

## model generation

The element type used in the analysis is SHell99, which is a layered element especially useful for modeling composite plates. By default, the element coordinate system is derived from the local geometry of each element. Although this may be convenient in certain cases, in this problem it is required that the element coordinate system for each element be aligned with the global Cartesian system. This is achieved by using the ESYS command. After specifying real constants (thickness and layer information)


Fig. 10.2 Variation of displacement components at points A and B as the load increases incrementally.


Fig. 10.3 Composite plate with a hole.
and orthotropic material properties, and creating the solid model, the mesh is generated. The mesh has all of its nodes and elements on the $x-y$ plane, i.e., $z=0$. However, in order to capture the buckling behavior, the flatness of the plate is slightly perturbed by means of a double-sinusoidal surface, as explained earlier. This is achieved by modifying the $z$-coordinates of each node through Eq. (10.2). When a mesh is generated on solid modeling entities (lines, areas and volumes), the resulting nodes and elements are attached to solid modeling entities and their modification is not allowed. Therefore, the mesh (nodes and elements) is detached from the solid model using the MODMSH command, after which the nodal coordinates are modified utilizing a do loop in accordance with Eq. (10.2). The following command input is used for model generation:

```
/ PREP7
ET,1,SHELL99
ESYS,0
R,1,12,1
RMORE
RMORE, 1, 30,.01, 1, -30,.01
RMORE, 1, 30,.01,1, -30,.01
RMORE, 1, 30,.01, 1, -30,.01
MP,EX,1,18.5E6
MP,EY,1,1.6E6
MP,GXY,1,0.832E6
MP, PRXY,1,0.35
K,1,0
K,2,9.5
K,3,9.5,9.5
K,4,0,9.5
A, 1, 2, 3, 4
CYL4,4.75,4.75,1.5
ASBA,1,2
NUMCMP,ALL
LSEL,S, , 1,4
LESIZE, ALL, , , 10
LSEL,S, , 5,8
LESIZE,ALL, , ,5
AMESH,ALL
MODMSH, DETACH
*GET, NNUMBER, NODE ، 0, NUM, MAX
PI=4*ATAN(1)
*DO,I, 1, NNUMBER
*GET, TMPX,NODE , I, LOC, X
*GET, TMPY,NODE, I, LOC , Y
```

```
```

! ENTER PREPROCESSOR

```
```

! ENTER PREPROCESSOR
! SPECIFY ELEMENT TYPE
! SPECIFY ELEMENT TYPE
! ALIGN ELEM CS WITH GLOBAL
! ALIGN ELEM CS WITH GLOBAL
! CARTESIAN
! CARTESIAN
! SPECIFY REAL CONSTANTS
! SPECIFY REAL CONSTANTS
!
!
!
!
!
!
! SPECIFY MATERIAL
! SPECIFY MATERIAL
! PROPERTIES
! PROPERTIES
!
!
!
!
!
!
! CREATE KEYPOINTS
! CREATE KEYPOINTS
!
!
!
!
!
!
! CREATE SQUARE AREA
! CREATE SQUARE AREA
! CREATE CIRCULAR AREA
! CREATE CIRCULAR AREA
! SUBTRACT AREAS
! SUBTRACT AREAS
! COMPRESS ENTITY NUMBERS
! COMPRESS ENTITY NUMBERS
! SELECT LINES
! SELECT LINES
! SPECIFY \# OF DIVISIONS
! SPECIFY \# OF DIVISIONS
! SELECT LINES
! SELECT LINES
! SPECIFY \# OF DIVISIONS
! SPECIFY \# OF DIVISIONS
! MESH AREA
! MESH AREA
! DETACH MESH FROM SOLID
! DETACH MESH FROM SOLID
! MODEL
! MODEL
! GET MAXIMUM NODE \# {ALSO \#
! GET MAXIMUM NODE \# {ALSO \#
! OF NODES AS NUMBERS ARE
! OF NODES AS NUMBERS ARE
! COMPRESSED)
! COMPRESSED)
! DEFINE PI (3.1415...)
! DEFINE PI (3.1415...)
! START DO LOOP ON NODES
! START DO LOOP ON NODES
! GET X-COORD OF CURRENT
! GET X-COORD OF CURRENT
! NODE
! NODE
! GET Y-COORD OF CURRENT
! GET Y-COORD OF CURRENT
! NODE

```
```

! NODE

```
```

```
TMPZ=SIN(PI*NX(I)/9.5)*SIN(PI*NY(I)/9.5)*0.0012
    ! EQ. (10.2)
N,I,NX(I),NY(I),TMPZ ! REDEFINE CURRENT NODE
*ENDDO ! END DO LOOP
FINISH ! EXIT FROM PREPROCESSOR
```


## SOLUTION

In the solution phase of this problem, first, nonlinear geometry effects are turned on (NLGEOM command). Results associated with each substep are written to the results file using the OUtres command. In order to improve accuracy and convergence, the number of substeps is specified as 100 using the NSUBST command. Displacement constraints are then applied using the D command. The condition that the $x$-displacement be uniform along the right boundary ( $x=9.5$ ) is imposed by selecting all of the nodes along that boundary and issuing the CP command. This command defines a set of coupled degrees of freedom (DOF) of the selected nodes (in this case the $x$ displacement) and enforces these DOF to be equal. The concept of coupled DOF is described as a separate topic in Sec. 11.1. The following command input is used for the solution:

| / SOLU | ! ENTER SOLUTION PROCESSOR |
| :---: | :---: |
| ANTYPE, STATIC | ! SPECIFY STATIC SOLUTION |
| NLGEOM, ON | ! TURN NONLINEAR GEOMETRY ON |
| OUTRES, ALI, ALL | ! WRITE RESULTS FOR EVERY SUBSTEP |
| NSUBST, 100 | ! SET NUMBER OF SUBSTEPS TO BE 100 |
| NSEL, S, LOC, X, 0 | ! SELECT NODES AT $\mathrm{X}=0$ |
| D, ALL, ALL | ! CONSTRAIN ALL DOFS |
| NSEL, S, LOC, Y, O | ! SELECT NODES AT Y $=0$ |
| NSEL, A, LOC, Y, 9.5 | ! ADD TO SELECTION NODES AT Y $=9.5$ |
| D, ALL, UZ, 0 | ! CONSTRAIN Z-DISPLACEMENTS |
| D, ALL, ROTY, 0 | ! CONSTRAIN ROTATIONS ABOUT Y-AXIS |
| NSEL, S, LOC, X, 9.5 | ! SELECT NODES AT X $=9.5$ |
| D, ALL, UZ, 0 | ! CONSTRAIN Z-DISPLACEMENTS |
| D, ALL, ROTX, 0 | ! CONSTRAIN ROTATIONS ABOUT X-AXIS |
| D, ALL, ROTY, 0 | ! CONSTRAIN ROTATIONS ABOUT Y-AXIS |
| ALLSEL | ! SELECT EVERYTHING |
| D, ALL, ROTZ, 0 | ! CONSTRAIN ROTATIONS ABOUT Z-AXIS |
| NSEL, S, LOC, X, 9.5 | ! SELECT NODES AT X $=9.5$ |
| CP, 1, UX, ALL | ! COUPLE X-DISPL. OF SELECTED NODES |
| NSEL,R,LOC, Y, 4.75 | ! RESELECT NODE AT Y $=4.75$ |
| F, ALL, FX, -12000 | $!$ APPLY TOTAL LOAD IN X-DIRECTION |
| ALLSEL | ! SELECT EVERYTHING |
| SOLVE | ! OBTAIN SOLUTION |
| FINISH | ! EXIT SOLUTION PROCESSOR |

## POSTPROCESSING

Because the loading is applied slow enough that there are no dynamic effects within the structure, referred to as quasi-static, the nonlinear nature of the problem requires that the load is incrementally increased.

Turning on the nonlinear geometry effects using the NLGEOM command leads to the problem being solved in small increments (substeps), even though the analysis type is declared as static. ANSYS automatically assigns the value 1 as the time at the end of the load step. In nonlinear static solutions, time is a measure of the fraction of the total load applied at the current substep. For example, if time has a value of 0.2 at a particular substep, this means that the load applied during solution at that substep is $20 \%$ of the total load. In the following command input, the node numbers of points A , $B$, and $C$ are stored in parameters $\mathrm{NA}, \mathrm{NB}$, and NC , respectively, followed by extraction of the total number of substeps (parameter SB). A do loop is set up so that the results associated with every substep can be retrieved and written to external files sequentially. Within the do loop, the time associated with the current substep is extracted (parameter $\boldsymbol{T T}$ ) for subsequent scaling of the applied load (parameter TF), and $z$-displacements at points $A$ and $B$ and the absolute value of the $x$-displacement at point C are written to files AZD.OUT, BZD.OUT, and CXD.OUT, respectively. Figure 10.4 shows the variation of these displacements with incremental load. In Fig. 10.4, zdisplacements at points $A$ and $B$ are denoted by $w(A)$ and $w(B)$, respectively, and $u(C)$ designates the absolute value of the $x$-displacement at point $C$.

```
/POST1
NSEL,S,LOC,X,4.75
NSEL,R,LOC,Y,6.25
*GET, NA, NODE, 0, NUM, MIN
NSEL,S,LOC,X,6.25
NSEL,R,LOC,Y,4.75
*GET , NB, NODE, O, NUM, MIN
NSEL,S,LOC,X,9.5
NSEL,R,LOC,Y,4.75
*GET , NC, NODE, 0, NUM, MIN
ALLSEL
SET,LAST
*GET,SB,ACTIVE,0,SET,SBST
*DO,I,1,SB
SET,1, I
*GET, TT, ACTIVE, 0, SET, TIME
TF=TT*12000
/OUTPUT,AZD,OUT, ,APPEND
*VWRITE,TF,UZ (NA)
(E16.8,5X, E16.8)
```

```
! ENTER POSTPROCESSOR
! SELECT NODES AT X = 4.75
! RESELECT NODE AT Y = 6.25
! STORE NODE # OF PT A INTO NA
! SELECT NODES AT X = 6.25
! RESELECT NODE AT Y = 4.75
! STORE NODE # OF PT B INTO NB
! SELECT NODES AT X = 9.5
! RESELECT NODE AT Y = 4.75
! STORE NODE # OF PT C INTO NC
! SELECT EVERYTHING
! SET RESULTS TO LAST SUBSTEP
! STORE # OF SUBSTEPS TO SB
! START DO LOOP
! SET RESULTS TO SUBSTEP I
! STORE CURRENT TIME TO TT
! FIND CURRENT FORCE
! REDIRECT OUTPUT TO FILE AZD.OUT
! WRITE TIME & DISP TO FILE
! FORMAT STATEMENT
```



Fig. 10.4 Variation of displacement components at points A, B, and C as the load increases incrementally.

```
/OUTPUT
/OUTPUT, BZD, OUT, , APPEND
*VWRITE,TF,UZ (NB)
(E16.8,5X, E16.8)
/OUTPUT
/OUTPUT,CXD,OUT, ,APPEND
*VWRITE,TF,ABS (UX(NC))
(E16.8,5X,E16.8
/OUTPUT
* ENDDO
```

```
! REDIRECT OUTPUT TO
```

! REDIRECT OUTPUT TO
! OUTPUT WINDOW
! OUTPUT WINDOW
! REDIRECT OUTPUT TO FILE BZD.OUT
! REDIRECT OUTPUT TO FILE BZD.OUT
! WRITE TIME \& DISP TO FILE
! WRITE TIME \& DISP TO FILE
! FORMAT STATEMENT
! FORMAT STATEMENT
! REDIRECT OUTPUT TO
! REDIRECT OUTPUT TO
! OUTPUT WINDOW
! OUTPUT WINDOW
! REDIRECT OUTPUT TO FILE CXD.OUT
! REDIRECT OUTPUT TO FILE CXD.OUT
! WRITE TIME \& DISP TO FILE
! WRITE TIME \& DISP TO FILE
! FORMAT STATEMENT
! FORMAT STATEMENT
! REDIRECT OUTPUT TO
! REDIRECT OUTPUT TO
! OUTPUT WINDOW
! OUTPUT WINDOW
! END DO LOOP

```
! END DO LOOP
```


### 10.2 Material Nonlinearity

Material nonlinearities arise from the presence of time-independent behavior, such as plasticity, time-dependent behavior such as creep, and viscoelastic/viscoplastic behavior where both plasticity and creep effects occur simultaneously. They may result in load sequence dependence and energy dissipation (irreversible structural behavior).

ANSYS offers a wide variety of nonlinear material behavior models, including nonlinear elasticity, hyperelasticity, viscoelasticity, plasticity, viscoplasticity, creep, swelling, and shape memory alloys. Several of these nonlinear material models can be specified in a combined fashion (an exhaustive list
of models that can be combined is given in the ANSYS Structural Analysis Guide). In the following subsections, four problems are considered that demonstrate the solution methods involving plasticity with isotropic hardening, viscoelasticity, viscoplasticity with Anand's model, and combined plasticity and creep.

### 10.2.1 Plastic Deformation of an Aluminum Sphere

Consider a thin-walled aluminum sphere with a radius of $r=10$ in and a thickness of $t=0.2$ in, as shown in Fig. 10.5. The sphere is subjected to an internal pressure of $p_{0}=1600$ psi . The elastic modulus and Poisson's ratio of the shell are $E=10^{7} \mathrm{psi}$ and $v=0.3$, respectively. The plastic behavior of aluminum is governed by

$$
\begin{equation*}
\sigma_{e}=30000+136000\left(\varepsilon_{p}\right)^{1 / 2} \tag{10.3}
\end{equation*}
$$

in which $\sigma_{e}$ is the effective stress and $\varepsilon_{p}$ designates the plastic strain. Figure 10.6 shows the stress vs. total strain ( $\varepsilon=\varepsilon_{e}+\varepsilon_{p}, \varepsilon_{e}$ : elastic strain) curve based on Eq. (10.3). This curve is input in ANSYS by means of a data table for nonlinear material behavior, which is given through data points (see Table 10.1). The goal is to obtain the radial displacements, as well as the strain field.

## MODEL GENERATION

In order to model the thin-walled aluminum sphere, a 2 -noded axisymmetric shell element (Shell51) is used. For the nonlinear material behavior, the multiple-point isotropic hardening rule is chosen (TB command with MISO option). Twenty data points for strain and stress values are entered using the TBPT command [the data point $(0,0)$ is implied]. The stress value of the first data point defines the yield stress, i.e., $\sigma_{y s}=30000$ psi . Due to the symmetry conditions, only a quarter circle is modeled.


Fig. 10.5 Aluminum sphere subjected to internal pressure, and corresponding finite element mesh using axisymmetric shell elements (SHELL51).


Fig. 10.6 Nonlinear stress-strain behavior of aluminum.

Table 10.1 Data points for nonlinear stress-strain behavior of aluminum.

| $\varepsilon$ | $\sigma_{e}(\mathrm{psi})$ |
| :---: | :---: |
| 0 | 0 |
| 0.00300 | 30000 |
| 0.00350 | 33041 |
| 0.00400 | 34300 |
| 0.00450 | 35267 |
| 0.00500 | 36082 |
| 0.00550 | 36800 |
| 0.00600 | 37449 |
| 0.00650 | 38045 |
| 0.00700 | 38601 |
| 0.00750 | 39123 |
| 0.00800 | 39616 |
| 0.00850 | 40086 |
| 0.00875 | 40312 |
| 0.00900 | 40534 |
| 0.00925 | 40751 |
| 0.00950 | 40964 |
| 0.00975 | 41173 |
| 0.02400 | 49708 |
| 0.04000 | 56160 |
| 0.06300 | 63313 |


| /PREP7 | ! ENTER PREPROCESSOR |
| :---: | :---: |
| ET, 1, 51 | ! USE SHELL51 ELEMENT |
| R, 1, 0.2 | ! SPECIFY THICKNESS |
| MP, EX, 1, 1E7 | ! SPECIFY ELASTIC MODULUS |
| MP, NUXY, 1, 0.3 | ! SPECIFY POISSON'S RATIO |
| TB, MISO, 1, 1, 20, | ! SPECIFY MULTILINEAR ISOTROPIC ! HARDENING |
| TBPT, $0.00300,30000$ | ! ENTER STRAIN VS STRESS DATA ! POINTS |
| TBPT, $0.00350,33041$ | ! |
| TBPT, 0.00400,34300 | ! |
| TBPT, 0.00450,35267 | ! |
| TBPT, 0.00500,36082 | ! |
| TBPT, , 0.00550,36800 | $!$ |
| TBPT, 0.00600,37449 | $!$ |
| TBPT, 0.00650,38045 | ! |
| TBPT, $0.00700,38601$ | ! |
| TBPT, 0.00750,39123 | ! |
| TBPT, , 0.00800,39616 | $!$ |
| TBPT, 0.00850,40086 | ! |
| TBPT, 0.00875,40312 | $!$ |
| TBPT, , 0.00900,40534 | ! |
| TBPT, , $0.00925,40751$ | ! |
| TBPT, , 0.00950,40964 | ! |
| TBPT, 0.00975,41173 | ! |
| TBPT, 0.02400,49708 | $!$ |
| TBPT, 0.04000,56160 | $!$ |
| TBPT, , $0.06300,63313$ | ! |
| K, 1, 0, 0 | ! CREATE KEYPOINTS |
| $\mathrm{K}, 2,10,0$ | ! |
| K, 3, 0, 10 | ! |
| LARC, 2, 3, 1, 10 | ! CREATE ARC |
| LESIZE, ALL, , 20 | ! SPECIFY \# OF ELEMENTS ON LINE |
| LMESH, ALL | ! MESH LINE |
| FINISH | ! EXIT PREPROCESSOR |

## SOLUTION

The solution phase of this problem is rather straightforward and involves application of boundary conditions and the internal pressure. No specific solution controls options are specified; ANSYS uses default settings for nonlinear solution.

```
/ SOLU
P0=1600
NSEL,S,LOC,Y,O
D,ALL,UY,0
NSEL,S,LOC, Z,0
SFE,ALL,1,PRES, , P0
SOLVE
FINISH
```

```
! ENTER SOLUTION PROCESSOR
! DEFINE PARAMETER FOR INTERNAL
! PRESSURE
! SELECT NODES AT Y = 0
! CONSTRAIN Y-DISPL AT SELECTED NODES
! SELECT NODES AT Z = 0
! SPECIFY INTERNAL PRESSURE ALONG ELEMS
! OBTAIN SOLUTION
! EXIT SOLUTION PROCESSOR
```


## POSTPROCESSING

The elastic and plastic strains along the inner surface of the sphere are stored in element tables using the etable command. Three strain components are considered, i.e., meridional, through-the-thickness, and hoop strains. Finally, total strains are obtained by adding elastic and plastic strains for each component using the SADD command, which adds columns to the element table. The element tables are listed in the Output Window using the pretab command. Figure 10.7 shows the listing of total strains at elements as they appear in the Output Window.

```
/POST1 ! ENTER POSTPROCESSOR
ETABLE,E_MER_TP,LEPEL,1
ETABLE, E_THK_TP,LEPEL,2
ETABLE,E_HOP_TP,LEPEL, 3
ETABLE,P_MER_TP,LEPEL,1
ETABLE, P_THK_TP,LEPEL, 2
ETABLE, P_HOP_TP,LEPEL, 3
```

```
! STORE ELASTIC
```

! STORE ELASTIC
! STRAINS IN ELEMENT
! STRAINS IN ELEMENT
! TABLE
! TABLE
!
!
! STORE PLASTIC
! STORE PLASTIC
! STRAINS IN ELEMENT
! STRAINS IN ELEMENT
! TABLE
! TABLE
!

```
!
```



Fig. 10.7 Listing of element table items as they appear in the Output Window.

```
SADD,T_MER_TP,E_MER_TP,P_MER_TP,1,1 ! ADD ELEMENT TABLE
SADD,T_THK_TP,E_THK_TP,P_THK_TP,1,1 ! ITEMS TO FIND TOTAL
SADD,T_HOP_TP, E_HOP_TP,P_HOP_TP,1,1 ! STRAINS
PRETAB,E_MER_TP,E_THK_TP,E_HOP_TP ! LIST ELASTIC
! STRAINS
PRETAB, P_MER_TP,P_THK_TP,P_HOP_TP ! LIST PLASTIC
! STRAINS
!LIST TOTAL STRAINS
```


### 10.2.2 Plastic Deformation of an Aluminum Cylinder

Consider a thin-walled aluminum cylinder with a radius of $r=20 \mathrm{in}$, height of $r=72 \mathrm{in}$, a thickness of $t=0.5 \mathrm{in}$, and an extremely stiff panel on one end, as shown in Fig. 10.8. The cylinder and the stiff panel are first subjected to an internal pressure of $p_{0}=1500 \mathrm{psi}$. With the internal pressure in place, the stiff panel is subjected to four tangential forces of $10^{5} \mathrm{lb}$, as illustrated in Fig. 10.8. All displacements and rotations at the cylinder's end opposite to the stiff panel are constrained. The elastic modulus and Poisson's ratio of the aluminum shell are $E_{a l}=10^{7} \mathrm{psi}$ and $v_{a l}=0.3$, respectively. The same set of properties corresponding to the stiff panel are $E_{s t}=10^{11} \mathrm{psi}$ and $v_{a l}=0$. The plastic behavior of aluminum is governed by


Fig. 10.8 Geometry (top) and the corresponding mesh (bottom) of the aluminum cylinder and the stiff panel.

Eq. (10.3). The data points for the stress vs. strain are plotted in Fig. 10.6 and tabulated in Table 10.1. The goal is to obtain the plastic strain field resulting from the internal pressure and added torsion.

## MODEL GENERATION

Both the aluminum cylinder and the stiff panel are modeled using an 8noded shell element, SHELL93. For the nonlinear material behavior, the multiple-point isotropic hardening rule is chosen ( $\mathbf{T B}$ command with MISO option). Although the geometry is axisymmetric, the loading is not. Therefore, the entire geometry is modeled.

```
/PREP7
R=20
H=72
T=R/40
NDIV1=10
NDIV2=15
P=1500
F=100E3
ET,1,93
KEYOPT,1,4,0
KEYOPT,1,5,2
KEYOPT,1,6,1
KEYOPT,1,8,2
R,1,T
MP,EX,1,1E7
MP,NUXY,1,0.3
TB,MISO, 1, 1, 20,
TBPT, ,0.00300,30000
TBPT,,0.00350,33041
TBPT,,0.00400,34300
TBPT,,0.00450,35267
TBPT,,0.00500,36082
TBPT,,0.00550,36800
TBPT,,0.00600,37449
TBPT,,0.00650,38045
TBPT,,0.00700,38601
TBPT,,0.00750,39123
TBPT,,0.00800,39616
TBPT,,0.00850,40086
TBPT,,0.00875,40312
TBPT,,0.00900,40534
TBPT,,0.00925,40751
TBPT,,0.00950,40964
```

```
ENTER PREPROCESSOR
```

ENTER PREPROCESSOR
! PARAMETER FOR RADIUS
! PARAMETER FOR RADIUS
! PARAMETER FOR HEIGHT
! PARAMETER FOR HEIGHT
! PARAMETER FOR THICKNESS
! PARAMETER FOR THICKNESS
! \# OF DIVISIONS IN RADIAL
! \# OF DIVISIONS IN RADIAL
!DIRECTION
!DIRECTION
! \# OF DIVISIONS IN HEIGHT
! \# OF DIVISIONS IN HEIGHT
!DIRECTION
!DIRECTION
! INTERNAL PRESSURE
! INTERNAL PRESSURE
! TANGENTIAL FORCE
! TANGENTIAL FORCE
! USE SHELL93 ELEMENT TYPE
! USE SHELL93 ELEMENT TYPE
! NO USER SUBROUTINE FOR ELEMENT CS
! NO USER SUBROUTINE FOR ELEMENT CS
! PROVIDE NODAL STRESS PRINTOUT
! PROVIDE NODAL STRESS PRINTOUT
! PROVIDE NONLINEAR INTEG. PT.
! PROVIDE NONLINEAR INTEG. PT.
! PRINTOUT
! PRINTOUT
! STORE DATA FOR TOP, BOTTOM \& MID
! STORE DATA FOR TOP, BOTTOM \& MID
! SURFACES
! SURFACES
! SPECIFY THICKNESS FOR ALUMINUM
! SPECIFY THICKNESS FOR ALUMINUM
! SPECIFY ELASTIC MODULUS FOR
! SPECIFY ELASTIC MODULUS FOR
! ALUMINUM
! ALUMINUM
! SPECIFY POISSON'S RATIO FOR
! SPECIFY POISSON'S RATIO FOR
! ALUMINUM
! ALUMINUM
! MULTILINEAR ISOTROPIC HARDENING
! MULTILINEAR ISOTROPIC HARDENING
! PLASTICITY
! PLASTICITY
! ENTER STRAIN VS STRESS DATA
! ENTER STRAIN VS STRESS DATA
! POINTS

```
! POINTS
```

```
TBPT,.0.00975,41173 !
TBPT,,0.02400,49708 !
TBPT,,0.04000,56160 !
TBPT,,0.06300,63313 !
R,2,T
MP,EX, 2,1E11
MP, NUXY, 2,0
CYLIND,R,0,0,H
VDEL,ALL
ASEL,S,LOC, Z,0
ASEL,A, LOC, Z,H
ADEL, ALL
ALLSEL
LSEL,S,LOC,Z,H/2
LESIZE,ALL, , NDIV2
ALLSEL
LESIZE,ALL,,,NDIV1
LSEL,S,LOC, Z,0
LSEL,R,LOC,Y,0,R
LCCAT,ALL
LSEL,S,LOC, Z, O
LSEL, R,LOC,Y,0,-R
LCCAT,ALL
LSEL,S,LOC,Z,H
LSEL, R, LOC, Y, O, R
LCCAT,ALL
LSEL,S,LOC,Z,H
LSEL, R,LOC,Y,O,-R
LCCAT, ALL
ALLSEL
MSHKEY,1
MAT,1
REAL, 1
AMESH, ALL
WPOFFS,0,0,H
PCIRC,R,0,0,90
PCIRC,1.5*R,R,0,90
ASEL,S,LOC,Z,H
AGLUE, ALL
CSYS,1
```

```
LSLA,S !SELECT LINES ATTACHED TO SELECTED
LSEL,U,IOC,X,1.25*R ! UNSELECT LINES AT X = 1.25*R
LESIZE,ALL,,,NDIV1 !SPECIFY # OF DIVS ON SELECTED
LSELL,R,LOC,X,1.25*R
LESIZE,ALL,,,NDIV1/2
MAT, 2
REAL, 2
AMESH, ALL
CSYS
ARSYM,X,ALL
ARSYM,Y,ALL
ALLSEL
NUMMRG,ALL
FINISH
```

```
! AREAS
```

! AREAS
! LINES
! LINES
! SELECT LINES ATTACHED TO SELECTED
! SELECT LINES ATTACHED TO SELECTED
! AREAS
! AREAS

```
! RESELECT LINES AT X = 1.25*R
```

! RESELECT LINES AT X = 1.25*R
! SPECIFY \# OF DIVS ON SELECTED
! SPECIFY \# OF DIVS ON SELECTED
LINES
LINES
SWITCH MATERIAL ATTRIBUTE TO MAT
SWITCH MATERIAL ATTRIBUTE TO MAT
| \# 2
| \# 2
!SWITCH REAL c ATTRIBUTE TO REAL
!SWITCH REAL c ATTRIBUTE TO REAL
! \# 2
! \# 2
!MESH SELECTED AREAS
!MESH SELECTED AREAS
! SWITCH TO GLOBAL CARTESIAN CS
! SWITCH TO GLOBAL CARTESIAN CS
! REFLECT SELECTED AREAS ABOUT Y-Z
! REFLECT SELECTED AREAS ABOUT Y-Z
! PLANE
! PLANE
! REFLECT SELECTED AREAS ABOUT X-Z
! REFLECT SELECTED AREAS ABOUT X-Z
! PLANE
! PLANE
! SELECT EVERYTHING
! SELECT EVERYTHING
! MERGE DUPIICATE ENTITIES
! MERGE DUPIICATE ENTITIES
! EXIT PREPROCESSOR

```
! EXIT PREPROCESSOR
```


## SOLUTION

The solution is obtained in two steps. First, the internal pressure is applied and the first load step file is generated (LSWRITE command). Then, the concentrated loads are applied while the internal pressure is still present, which constitutes the second load step file. The solution is obtained sequentially from these load step files using the LSSOLVE command.

```
/ SOLU
NSEL,S,LOC, Z,O
D,ALL,ALL
ALLSEL
ESEL,S,MAT, ,1
SFE,ALL, 1, PRES, , P
CSYS,1
ASEL,S,LOC,Z,H
ASEL,R,LOC,X,-R,R
CSYS
ESLA,S
SFE,ALL, 1, PRES, , P
ALLSEL
LSWRITE,1
N=NODE(1.5*R,0,H)
```

```
: ENTER SOLUTION PROCESSOR
! SELECT NODES AT Z = 0
! CONSTRAIN ALL DOFS (DISPL &
! ROTATIONS)
! SELECT EVERYTHING
! SELECT ELEMENTS WITH MAT # 1
! APPLY INTERNAL PRES. ON SELECTED
! ELEMS
! SWITCH TO GLOBAL CYLINDRICAL CS
! SELECT AREAS AT Z = H
! RESELECT AREAS BETWEEN X = -R &
: X = R
! SWITCH TO GLOBAL CARTESIAN CS
! SELECT ELEMENTS ATTACHED TO
! SELECTED AREAS
! APPLY INTERNAL PRES. ON SELECTED
! ELEMS
! SELECT EVERYTHING
!WRITE LOAD STEP FILE # 1
! STORE NODE # IN N FOR GIVEN
! COORDS .
```

```
F,N,FY,F
N=NODE (-1.5*R,0,H)
F,N,FY,-F
N=NODE (0,1.5*R,H)
F,N,FX,-F
N=NODE (0, -1. 5*R,H)
F,N,FX,F
ALLSEL
LSWRITE, 2
LSSOLVE, 1,2
FINISH !EXIT SOLUTION PROCESSOR
```


## POSTPROCESSING

Once the solution is complete, the quantities of interest, i.e., radial, circumferential, and longitudinal displacements and circumferential plastic strains, are retrieved at a specific node and written to an external text file for both load steps. The channeling of data to the external text file, referred to as the Command File, is achieved by using the *CFOPEN command (detailed description is given in Sec. 11.5.2.2). After issuing the *CFOPEN command, quantities of interest are written to the file PLASTIC.OUT using the *VWRITE command. The aforementioned quantities are reviewed at a single node, which is chosen along the middle of the cylinder surface in the $z$-direction (i.e., $z=H / 2$ ). Plastic strains are calculated for elements. Therefore, plastic strains in the circumferential direction are stored in an element table using the ETABLE command. There are two elements attached to the selected single node. Therefore, the plastic strains at these two elements are averaged to find its value at the shared node. The results associated with specific load steps are read using the SET command.

```
/ POST1
*CFOPEN, PLASTIC,OUT
CSYS,1
RSYS,1
SET,1
ETABLE,EPT, EPPL,Y
NSEL,S,LOC, Z,H/2
NSEL, R,LOC,Y,O
ESLN,S
*GET, E1, ELEM,0 ,NUM, MAX
*GET,E2,ELEM, 0,NUM,MIN
```

```
! ENTER GENERAL POSTPROCESSOR
! OPEN "COMMAND FILE"
! SWITCH TO GLOBAL CYLINDRICAL
!CS
! USE CYLINCRICAL CS FOR
! RESULTS
! READ RESULTS OF LOAD STEP 1
! STORE PLASTIC STRAINS IN
! THETA DIRECTION IN ELEMENT
! TABLE
! SELECT NODES AT Z = H/2
! RESELECT NODES AT THETA = 0
! SELECT ELEMENTS ATTACHED TO
! SELECTED NODE
! STORE MAX ELEM # IN PARAMETER
! E1
! STORE MIN ELEM # IN PARAMETER
! E2
```

| *GET, EPT1, ETAB, 1, ELEM, E1 | ! STORE ELEM TABLE ITEMS FOR E1 |
| :---: | :---: |
| *GET, EPT2, ETAB , 1, ELEM, E2 |  |
|  | ! EPT2 |
| EPTAV $=(E P T 1+E P T 2) / 2$ | ! FIND AVERAGE OF EPT1 \& EPT2 |
| *GET, NODE1, NODE, 0, NUM, MAX | ! RETRIEVE SELECTED NODE NUMBER |
| * GET, UR1, NODE, NODE1, U, X | ! RETRIEVE RADIAL DISPLACEMENT |
| *GET, UT1, NODE, NODE1, U, Y | ! RETRIEVE CIRCUMF. |
|  | ! DISPLACEMENT |
| *GET, UZ1, NODE, NODE1, U, Z | ! RETRIEVE Z-DISPLACEMENT |
| *VWRITE, 'TIME = 1' | ! START WRITING TO COMMAND FILE |
| (A8) | ! FORMAT STATEMENT |
| *VWRITE, 'UR = ', UR1 | ! WRITE RADIAL DISPLACEMENT |
| (A9, E14.5) | ! FORMAT STATEMENT |
| *VWRITE, 'UTHETA $=$ ', UT1 | ! WRITE CIRCUMF. DISPLACEMENT |
| (A9, E14.5) | ! FORMAT STATEMENT |
| *VWRITE, 'UZ = ', UZ1 | ! WRITE Z-DISPLACEMENT |
| (A9, E14.5) | ! FORMAT STATEMENT |
| *VWRITE, 'EPLTH $=$ ', EPTAV | ! WRITE AVERAGED PLASTIC STRAIN |
| (A9, E14.5) | ! FORMAT STATEMENT |
| SET, NEXT | ! READ RESULTS OF LOAD STEP 2 |
| ETABLE, REFL | ! UPDATE ELEMENT TABLE |
| *GET, EPT1, ETAB, 1, ELEM, E1 | ! STORE ELEM TABLE ITEMS FOR E1 |
| *GET, EPT2, ETAB, 1, ELEM, E2 |  |
|  | ! EPT2 |
| $\mathrm{EPTAV}=(\mathrm{EPT} 1+\mathrm{EPT} 2) / 2$ | ! FIND AVERAGE OF EPT1 \& EPT2 |
| *GET, UR1, NODE, NODE1, U, X | ! RETRIEVE RADIAL DISPLACEMENT |
| *GET, UT1, NODE, NODE1, U, Y | ! RETRIEVE CIRCUMF. |
|  | ! DISPLACEMENT |
| *GET, UZ1, NODE, NODE1, U, Z | ! RETRIEVE Z-DISPLACEMENT |
| *VWRITE, 'TIME = ${ }^{\prime}$ ' | ! RESUME WRITING TO COMMAND |
|  | ! FILE |
| (A8) | ! FORMAT STATEMENT |
| *VWRITE, 'UR = ',UR1 | ! WRITE RADIAL DISPLACEMENT |
| (A9, E14.5) | ! FORMAT STATEMENT |
| *VWRITE, 'UTHETA $=$ ',UT1 | ! WRITE CIRCUMF. DISPLACEMENT |
| (A9, E14.5) | ! FORMAT STATEMENT |
| *VWRITE, 'UZ $=$ ', UZ1 | ! WRITE Z-DISPLACEMENT |
| (A9, E14.5) | ! FORMAT STATEMENT |
| *VWRITE, 'EPLTH = ', EPTAV | ! WRITE AVERAGED PLASTIC STRAIN |
| (A9, E14.5) | ! FORMAT STATEMENT |
| * CFCLOS | ! CLOSE "COMMAND FILE" |
| FINISH | ! EXIT GENERAL POSTPROCESSOR |

After the command input segment given above is executed, the contents of the file PLASTIC. OUT is given as

```
TIME = 1
    UR = 0.52447E+00
    UTHETA = -0.20702E-11
    UZ = 0.59981E-01
    EPLTH = 0.20770E-01
TIME = 2
    UR = 0.62531E+00
    UTHETA = 0.19934E+00
    UZ = 0.64357E-01
    EPLTH = 0.25750E-01
```


### 10.2.3 Stress Analysis of a Reinforced Viscoelastic Cylinder

Consider a hollow viscoelastic cylinder reinforced by an elastic material along the outer periphery subjected to internal pressure, as shown in Fig. 10.9. The cylinder is long in the out-of-plane direction. The inner and outer radii of the cylinder are $a=2$ in and $b=4$ in, respectively; the thickness of the reinforcing layer is $h=4 / 33 \mathrm{in}$. The elastic modulus and Poisson's ratio of the reinforcement are $E_{r}=3 \times 10^{7} \mathrm{psi}$ and $v_{r}=1 / \sqrt{11}$.

The shear and bulk moduli of the viscoelastic cylinder behave as

$$
\begin{gather*}
G(t)=G_{0} e^{-t}  \tag{10.4a}\\
K(t)=K_{\infty} H(t) \tag{10.4b}
\end{gather*}
$$

in which $G_{0}$ and $K_{\infty}$ are defined as $G(0)=G_{0}=E_{c} /\left(1+2 \nu_{c}\right)$ and $K_{\infty}=E_{c} /\left[3\left(1-2 v_{c}\right)\right]$. The elastic modulus and Poisson's ratio of the cylinder are $E_{c}=10^{5} p s i$ and $v_{c}=1 / 3$.

Within ANSYS, the viscoelastic material behavior is specified as

$$
\begin{align*}
& G(t)=G_{0}\left[\frac{G_{\infty}}{G_{0}}+\frac{G_{1}}{G_{0}} e^{\left(-t / \tau^{G}\right)}\right]  \tag{10.5a}\\
& K(t)=K_{0}\left[\frac{K_{\infty}}{K_{0}}+\frac{K_{1}}{K_{0}} e^{\left(-t / \tau^{K}\right)}\right] \tag{10.5b}
\end{align*}
$$



Fig. 10.9 Hollow viscoelastic cylinder with external reinforcement.

In accordance with Eq. (10.4), the parameters in Eq. (10.5) take the following values: $G(0)=G_{0}=G_{1}, \lim _{t \rightarrow \infty} G(t)=G_{\infty}=0, K_{0}=K_{\infty}$, and $K_{1}=0$. Therefore, the relative shear and bulk moduli, $\left(G_{1} / G_{0}\right)$ and ( $K_{1} / K_{0}$ ), respectively, are assigned the values of 1.0 and 0.0. The parameters $\tau^{G}$ and $\tau^{K}$ represent relative times for shear and bulk behavior, respectively; both of them are assumed to be 1.0 . The goal is to find the time-dependent behavior of radial and hoop stresses.

## MODEL GENERATION

The problem is solved using two-dimensional 8-noded PLANE183 elements with the axisymmetric option. A Prony series representation of viscoelasticity is specified using TB and TBDATA commands. The model is a rectangular cross section of the cylinder and the reinforcement layer with a height of 1 in .

| /PREP7 | ! ENTER PREPROCESSOR |
| :---: | :---: |
| ET, 1, PLANE183, , 1 | ! USE PLANE183 ELEM WITH |
|  | ! AXISYMMETRY |
| MP, EX, 1, 1.0E5 | ! ELASTIC MODULUS OF MAT 1 |
| MP, NUXY, 1, 1/3 | ! POISSON'S RATIO OF MAT 1 |
| TB, PRONY, 1, 1, SHEAR | ! USE PRONY SERIES VISCOELASTICITY |
|  | ! FOR SHEAR |
| TBDATA, 1, 1.0,1.0 | ! RELATIVE MODULUS AND TIME |
| TB, PRONY, 1, 1, BULK | ! USE PRONY SERIES VISCOELASTICITY |
|  | ! FOR BULK |
| TBDATA, 1, 0.0,1.0 | ! RELATIVE MODULUS AND TIME |
| MP, EX, 2, 3,0E7 | ! ELASTIC MODULUS OF MAT 2 |
| MP, NUXY, 2,1/SQRT (11) | ! POISSON'S RATIO OF MAT 2 |
| $\mathrm{R} 1=2$ | ! PARAMETER FOR INNER RADIUS OF |
|  | ! CYLINDER |
| $\mathrm{R} 2=4$ | ! PARAMETER FOR OUTER RADIUS OF |
|  | ! CYLINDER |
| $R 3=R 2+4 / 33$ | ! PARAMETER FOR OUTER RADIUS OF |
|  | ! REINFORC. |
| K, 1, R1 | ! CREATE KEYPOINTS |
| K, 2, R2 | ! |
| K, 3, R3 | $!$ ! |
| KGEN, $2,1,3,1,1$ | ! GENERATE KPS FROM EXISTING |
|  | ! PATTERN |
| L, 1, 2, 5 | ! CREATE LINES |
| L, 2, 3, 1 | $!$ |
| A, 4, 1, 2, 5 | ! CREATE AREAS |
| A, 5, 2, 3, 6 | ! |
| LESIZE, 1, , 10 | ! SPECIFY \# OF LINE DIVISIONS |
| LESIZE, $2, \ldots, 2$ | ! |
| ESIZE, 0.5 | ! SPECIFY ELEMENT SIZE |
| MSHKEY, 1 | ! ENFORCE MAPPED MESHING |
| MSHAPE, 0, 2D | ! ENFORCE QUADRILATERAL ELEMENT |
|  | ! SHAPE |
| MAT, 1 | ! SWITCH TO MATERIAL 1 |

```
AMESH, 1
MAT, 2
AMESH,2
FINISH
```

```
! MESH AREA 1
```

! MESH AREA 1

```
SWITCH TO MATERIAL 2
```

SWITCH TO MATERIAL 2
MESH AREA 2
MESH AREA 2
! EXIT PREPROCESSOR

```
! EXIT PREPROCESSOR
```


## SOLUTION

The solution is obtained in two load steps. The first load step, which spans an extremely small duration of 0.00001 sec , is used to set up the initial conditions for the problem. As shown in Fig. 10.9, the nodes along the bottom row are constrained to move in the vertical direction ( $y$-direction). In order to capture the plane strain characteristic of the problem, displacements are enforced to be the same on the top row of nodes. This is accomplished by coupling the degree of freedom in the $y$-direction of those nodes using the CP command. The solution controls option is turned off in this analysis, and values of the specific nonlinear solution controls items are specified. In order to reduce the size of the results file, only a limited amount of data is written (OUTRES command). The accuracy of the solution is improved by specifying a small convergence tolerance value using the CNVTOL command. Finally, the time step size is specified to be $0.1 \mathrm{sec} u s i n g$ the Deltim command.

```
/ SOLU
ANTYPE,STATIC
NSEL, S,LOC,Y,O
D, ALL,UY
NSEL,S,LOC,Y,1
CP,1,UY,ALI,
ALLSEL
NSEL,S, LOC, X, R1
PIO = 1
SF, ,PRES, PIO
ALLSEL
SOLCONTROL,O
OUTRES,BASIC,ALL
CNVTOL, F, , , 1E-7
TIME,0.00001
SOLVE
TIME,10
DELTIM,0.1
SOLVE
```

```
! ENTER SOLUTION PROCESSOR
! SPECIFY ANALYSIS TYPE
! SELECT NODES AT Y = 0
! CONSTRAIN Y-DISP AT SELECTED
! NODES
! SELECT NODES AT Y = 1
! COUPLE Y-DISP OF SELECTED NODES
!SELECT EVERYTHING
! SELECT NODES AT X = R1
! PARAMETER FOR INNER PRESSURE
! APPLY PRESSURE BC AT SELECTED
! NODES
SELECT EVERYTHING
! TURN SOLUTION CONTROLS OFF
! SAVE BASIC OUTPUT AT EVERY
! SUBSTEP
! SMALL CONVERGENCE TOLER. ENFORCED
! TIME AT THE END OF 1ST LOAD STEP
! OBTAIN SOLUTION FOR 1ST LOAD STEP
! TIME AT THE END OF 2ND LOAD STEP
! SPECIFY TIME STEP SIZE
! OBTAIN SOLUTION FOR 2ND LOAD STEP
```


## POSTPROCESSING

Once the solution is obtained, a component of nodes (the center row of nodes) is created using the CM command. The radial and hoop stresses are written to an external file, which is opened using the *CFOPEN command
and closed using the *CFCLOSE command. Between these two commands, desired quantities are written to the external file using the *VWRITE command followed by a format statement (written in FORTRAN syntax). Figures 10.10 and 10.11 show the variation of radial $\left(\sigma_{r r}\right)$ and hoop ( $\sigma_{\theta \theta}$ ) stresses, respectively, along $y=0.5$ in for times $t=0.1,1,5$, and 10 .

```
/POST1 !ENTER POSTPROCESSOR
ESEL,S,MAT,,1 !SELECT ELEMENTS WITH MAT 1
NSLE,S,CORNER !SELECT CORNER NODES ATTACHED
! TO THE SELECTED ELEMENTS
! RESELECT NODES AT Y = 0.5
! CREATE A COMPONENT OF
! NODES NAMED "NLIST"
! OPEN DATA FILE "STRS.OUT"
! READ RESULTS FOR 1ST LOAD
! STEP
! LOOP OVER RESULTS SETS
SELECT EVERYTHING
! OBTAIN TIME AT CURRENT
! RESULTS SET
! WRITE A SEPARATOR ROW TO
! FILE
I FORMAT STATEMENT
!WRITE CURRENT TIME TO FILE
! FORMAT STATEMENT
! SELECT COMPONENT "NLIST"
! OBTAIN NUMBER OF NODES
! LOOP OVER SELECTED NODES
OBTAIN NODE # CLOSEST
TO THE CENTER
! OBTAIN X-COORD OF
!THE NODE "NODNUM"
```




```
! WRITE }\mp@subsup{\sigma}{rr}{}\mathrm{ AND }\mp@subsup{\sigma}{00}{}\mathrm{ TO FILE
! FORMAT STATEMENT
! UNSELECT THE NODE "NODNUM"
! END LOOP OVER NODES
! IF TIM = 10, END LOOP
I READ NEXT RESULTS SET
! END LOOP OVER RESULTS SETS
! CLOSE DATA FILE
! SELECT EVERYTHING
```


### 10.2.4 Viscoplasticity Analysis of a Eutectic Solder Cylinder

Consider a cylindrical eutectic solder with a radius of 10 mm and a height of 100 mm , as shown in Fig. 10.12. The bottom surface of the cylinder is constrained in all directions while the top surface is subjected to a prescribed displacement in the $y$-direction as a sinusoidal function of time. In


Fig. 10.10 Variation of radial stress ( $\sigma_{r r}$ ) along $y=0.5$ in at different times.


Distance from Center (in)
Fig. 10.11 Variation of hoop stresses $\left(\sigma_{\theta \theta}\right)$ along $y=0.5$ in at different times.


Fig. 10.12 Cylindrical eutectic solder.
addition, the eutectic solder is exposed to temperature, which exhibits the same time-dependent behavior as the prescribed displacement.

These loading conditions are given by

$$
\begin{align*}
& u_{y}(x, y=100, z, t)=u_{y}^{\max } \sin \left(\frac{\pi t}{t_{f i n}}\right) \\
& T(x, y, z, t)=T_{\min }+\left(T_{\max }-T_{\min }\right) \sin \left(\frac{\pi t}{t_{f i n}}\right) \tag{10.6}
\end{align*}
$$

in which $u_{y}^{\max }$ and $T_{\max }$ are the maximum values of the applied displacement and temperature, respectively; $T_{\min }$ is the stress-free temperature; $t$ designates time; and $t_{f i n}$ is the time at the end of the process. The numerical values of these parameters are given as $u_{y}^{\max }=0.5 \mathrm{~mm}, T_{\min }=0^{\circ} \mathrm{C}$, $T_{\max }=125^{\circ} \mathrm{C}$, and $t_{f i n}=60 \mathrm{sec}$. The material properties of the solder vary with temperature, as tabulated in Table 10.2. In addition to the elastic material properties, Anand's viscoplastic material behavior is assumed for the solder, with related parameters listed in Table 10.3. The goal is to find strain field (elastic, inelastic and total strains) at different times.

## MODEL GENERATION

The problem is solved using two-dimensional viscoplastic elements (VISCO106) with the axisymmetric option. In order to specify temperaturedependent elastic properties, a temperature table is constructed using the

Table 10.2 Temperature-dependent variation of material properties of the eutectic solder.

| Temperature <br> $\left({ }^{\circ} \mathrm{C}\right)$ | Elastic Modulus <br> $(\mathrm{MPa})$ | Poisson's <br> Ratio | Coefficient of <br> Thermal Expansion <br> $\left(\mathrm{ppm} /{ }^{\circ} \mathrm{C}\right)$ |
| :---: | :---: | :---: | :---: |
| -35 | 40781 | 0.3540 | 24.27 |
| -15 | 37825 | 0.3565 | 24.48 |
| 5 | 34884 | 0.3600 | 24.66 |
| 25 | 31910 | 0.3628 | 24.80 |
| 50 | 28149 | 0.3650 | 25.01 |
| 75 | 24425 | 0.3700 | 25.26 |
| 100 | 20710 | 0.3774 | 25.52 |
| 125 | 16942 | 0.3839 | 25.79 |

Table 10.3 Numerical values of parameters used in Anand's material model for the eutectic solder.

| Parameter | Description | Value |
| :---: | :---: | :---: |
| $S_{0}$ | Initial deformation resistance | $12.41(\mathrm{MPa})$ |
| $Q / R$ | Ratio of activation energy to <br> universal gas constant | $9400\left(1 /{ }^{\circ} \mathrm{K}\right)$ |
| $\xi$ | Pre-exponential factor | $4 \times 10^{6}(1 / \mathrm{sec})$ |
| $m$ | Stress multiplier | 1.5 |
| $h_{0}$ | Strain rate sensitivity of stress | 0.303 |
| $n$ | Hardening/softening constant <br> reefistance saturation value <br> Strain rate sensitivity of <br> saturation value | $1379(\mathrm{MPa})$ |
| $a$ | Strain rate sensitivity of <br> hardening or softening | $13.79(\mathrm{MPa})$ |

MPTEMP command, followed by the specification of properties using the mPDATA command. Note that temperatures are specified in Kelvin. Anand's viscoplastic properties are specified using the TB and TBDATA commands. In the analysis, millimeter ( mm ) is used for length dimensions and megapascal ( MPa ) is used for stresses (also elastic modulus). Thus, the resulting displacements and stresses are in millimeters and megapascals, respectively. However, this is a special case where an inconsistent unit system works correctly, and it is highly recommended that a unit analysis be performed before applying similar approaches to other mixed-unit systems.

```
/PREP7 ! ENTER PREPROCESSOR
ET,1,VISCO106 ! USE VISCO106 ELEMENT
KEYOPT,1,3,1 !SPECIFY AXISYMMETRY
KEYOPT,1,5,2 ! REQUEST EXTRA OUTPUT
    ! (PLASTIC WORK)
    ! INITIALIZE MATERIAL TABLE
    ! SPECIFY TEMPERATURE POINTS
MPTEMP,1,238.15,258.15,278.15
MPTEMP,4,298.15,323.15,348.15,373.15,398.15
                            ! SPECIFY ELASTIC MODULUS
MPDATA, EX, 1, 1,40781,37825,34884
MPDATA, EX,1,4,31910,28149,24425,20710,16942
                            ! SPECIFY POISSON'S RATIO
MPDATA,NUXY, 1,1,0.354,0.3565,0.36
MPDATA,NUXY, 1,4,0.3628,0.365,0.37,0.3774,0.3839
                            SPECIFY COEFFICIENT OF THERMAL
                            ! EXPANSION
MPDATA,ALPX, 1,1,2.427e-5,2.448e-5,2,466e-5
MPDATA, ALPX,1,4,2.48e-5,2.501e-5,2.526e-5,2.552e-5,
                                    *2.579e-5
TB,ANAND,1 !SPECIFY ANAND'S VISCOPLASTICITY
    ! SPECIFY ANAND'S PARAMETERS
TBDATA, 1,12.41,9400.,4E6,1.5,0.303,1379
TBDATA, 7,13.79,0.07,1.3
RECTNG,0,10,0,100 ! CREATE RECTANGLE
ESIZE,2 !SPECIFY ELEMENT SIZE
MSHKEY,1 ! ENFORCE MAPPED MESHING
AMESH,ALL !MESH AREA
FINISH ! EXIT PREPROCESSOR
```


## SOLUTION

The solution is obtained in twenty load steps. Nonlinear geometry effects must be turned on when using viscoio6 elements. Automatic time stepping is turned on (AUTOTS command) so that ANSYS can adjust the time step size values for the substeps within each load step. However, the starting time step size and the minimum and maximum time step size to be used in the analysis are specified using the DELTIM command. A do loop is utilized for creating load step files, in which the applied displacement and temperature conditions are calculated and written to load step files. Finally, the solution is obtained using the LSSOLVE command.

```
/ SOLU
TMIN=273.15 !PARAMETER FOR MIN
! TEMPERATURE
! PARAMETER FOR MAX
! TEMPERATURE
DELT=TMAX-TMIN !TEMPERATURE DIFFERENCE
VMAX=0.5 !MAXIMUM APPLIED
! DISPLACEMENT
NLS=20 !NUMBER OF LOAD STEPS
TMX=60 !FINAL TIME (IN SECONDS)
```

| $\mathrm{PI}=4 *$ ATAN (1) | ! PARAMETER FOR $\pi$ |
| :---: | :---: |
| ANTYPE, STATIC | ! DECLARE ANALYSIS TYPE |
| NLGEOM, ON | ! TURN NONLINEAR GEOMETRY ON |
| KBC, 0 | ! APPLY RAMPED LOADING |
| AUTOTS, ON | ! TURN AUTOMATIC TIME |
|  | ! STEPPING ON |
| TREF, 273.15 | ! STRESS-FREE TEMPERATURE |
| DELTIM, 1E-2,1E-3, 0.3 | ! SPECIFY TIME STEP SIZE |
| NSEL, S, LOC, Y, 0 | ! SELECT NODES AT Y = 0 |
| D, ALL, ALL | ! CONSTRAIN ALL DOFS |
| ALLSEL | ! SELECT EVERYTHING |
| TIM=TMX/NLS | ! FIND TIME AT CURRENT LOAD |
|  | ! STEP |
| *DO, I, 1, NLS | ! BEGIN DO LOOP ON LOAD STEPS |
| TIME, TIM | ! SPECIFY TIME AT END OF LOAD |
|  | ! STEP |
| $V C=V M A X * S I N(P I * T I M / T M X)$ | ! FIND CURRENT DISPLACEMENT |
|  | ! BC |
| TC=TMIN+DELT*SIN (PI*TIM/TMX) | ! FIND CURRENT TEMPERATURE |
| NSEL, S, LOC, Y, 100 | ! SELECT NODES AT Y = 100 |
| D, ALL, UY, VC | ! APPLY CURRENT DISPLACEMENT |
|  | ! BC |
| ALLSEL | ! SELECT EVERYTHING |
| BFUNIF, TEMP, TC | ! APPLY CURRENT TEMPERATURE |
|  | ! LOAD |
| LSWRITE, I | ! WRITE LOAD STEP FILE |
| TIM $=$ TIM + TMX / NLS | ! UPDATE TIME AT END OF LOAD |
|  | ! STEP |
| * ENDDO | ! END DO LOOP ON LOAD STEPS |
| LSSOLVE, 1, NLS | ! SOLVE FROM LOAD STEP FILES |
| FINISH | ! EXIT SOLUTION PROCESSOR |

## POSTPROCESSING

Once the solution is obtained, the user can review a multitude of results items, including elastic, plastic, and total strain components and stress components. Also available is the plastic work. The command input segment given below obtains contour plots for a few of the items mentioned above, in two different load steps. The results for different load steps are read using the SET command. For clarity in the contour plots, the triad and the symbols for minimum and maximum quantities are turned off using the /TRIAD and /PLOPTS commands. The two load steps considered here are the $10^{\text {th }}$ and the last load steps. The $10^{\text {th }}$ load step corresponds to the maximum displacement and temperature conditions while the last load step brings the structure to its initial configuration. Although initially stress free, after applying time-dependent displacement and temperature loads and returning to a no-load state, the structure experiences residual stresses and strains. Figures 10.13 and 10.14 show the contour plots of plastic work at load steps 10 and 20 , respectively.


Fig. 10.13 Plastic work contours at load step $10(t=30 \mathrm{sec})$.


Fig. 10.14 Plastic work contours at load step $20(t=60 \mathrm{sec})$.

```
/ POST1
/TRIAD,OFF
/PLOPTS,MINM,O
SET,10
PLNSOL,NL,PLWK
PLNSOL,S,EQV
PLNSOL,EPPL,Y
SET,20
PLNSOL,NL, PLWK
PLNSOL,S,EQV
PLNSOL, EPPL,Y
```

```
ENTER POSTPROCESSOR
! TURN THE TRIAD OFF
! TURN MIN & MAX SYMBOLS OFF
! READ RESULTS AT LOAD STEP 10
! PLOT PLASTIC WORK CONTOURS
! PLOT EQUIVALENT STRESS CONTOURS
! PLOT CONTOURS FOR PLASTIC STRAIN
! IN Y-DIR.
! READ RESULTS AT LOAD STEP 20
! PLOT PLASTIC WORK CONTOURS
PLOT EQUIVALENT STRESS CONTOURS
! PLOT CONTOURS FOR PLASTIC STRAIN
I IN Y-DIR.
```


### 10.2.5 Combined Plasticity and Creep

In this subsection, the cylindrical eutectic solder column considered in the previous subsection is reconsidered, this time with nonlinear material properties by means of rate-independent plasticity combined with creep (Fig. 10.12). Both top and bottom surfaces of the cylinder are constrained in all directions. Temperature within the cylinder varies as a sinusoidal function of time as given by

$$
\begin{equation*}
T(x, y, z, t)=T_{\min }+\left(T_{\max }-T_{\min }\right) \sin \left(\frac{\pi t}{t_{f i n}}\right) \tag{10.7}
\end{equation*}
$$

in which $T_{\min }$ and $T_{\max }$ are the minimum and maximum values of the applied temperature, $t$ designates time, and $t_{f i n}$ is the time at the end of the process. The numerical values of these parameters are given as: $T_{\text {min }}=-35^{\circ} \mathrm{C}, T_{\text {max }}=125^{\circ} \mathrm{C}$, and $t_{\text {fin }}=60 \mathrm{sec}$. The material properties of the solder vary with temperature, as tabulated in Table 10.2. The material exhibits both isotropic hardening plasticity and creep behavior. For the creep behavior, the following strain rate equation is utilized:

$$
\begin{equation*}
\dot{\varepsilon}_{c r}=C_{1}\left[\sinh \left(C_{2} \sigma\right)\right]^{C_{3}} e^{-C_{4} / T} \tag{10.8}
\end{equation*}
$$

in which $\dot{\varepsilon}_{c r}$ is the creep strain rate, $T$ is the temperature in Kelvin, $\sigma$ is the equivalent stress, and $C_{1}, C_{2}, C_{3}$, and $C_{4}$ are specific parameters for the generalized Garofalo creep model. In this problem, values of these parameters are taken as $C_{1}=23.3 \times 10^{6}(1 / \mathrm{sec}), C_{2}=6.699 \times 10^{-2}, C_{3}=3.3$, and $C_{4}=8.12 \times 10^{3}\left(1 /^{\circ} \mathrm{K}\right)$. The plastic behavior at different temperatures is shown in Fig. 10.15. The goal is to find strain fields, including plastic and creep strains, at different times.


Fig. 10.15 Nonlinear stress-strain behavior of solder at different temperatures.

## MODEL GENERATION

The problem is solved using two-dimensional 8-noded PLANE183 elements with the axisymmetric option. In order to specify temperature-dependent elastic properties, a temperature table is constructed using the MPTEMP command, followed by the specification of properties using the MPDATA command. Note that temperatures are specified in Kelvin. The stress-strain variations for the isotropic hardening model at different temperatures are specified using the TB, TBTEMP, and TBPT commands. In the analysis, millimeter ( mm ) is used for length dimensions and megapascal ( MPa ) is used for stresses (also elastic modulus). Thus, the resulting displacements and stresses are in millimeters and megapascals, respectively. Again, this is a special case where an inconsistent unit system works correctly, and it is highly recommended that a unit analysis be performed before applying similar approaches to other mixed-unit systems.

```
/PREP7 ! ENTER PREPROCESSOR
ET,1,PLANE183 !USE PLANE183 ELEMENT
KEYOPT,1,3,1 !USE AXISYMMETRY
MPTEMP ! INITIALIZE MAT TEMP TABLE
MPTEMP,1,238.15,258.15,278.15 ! CONSTRUCT MAT TEMP
    ! TABLE
MPTEMP,4,298.15,323.15,348.15,373.15,398.15
                                    !SPECIFY TEMPERATURE
                            ! DEPENDENT ELASTIC MODULUS
MPDATA, EX, 1,1,40781,37825,34884
MPDATA, EX, 1,4,31910,28149,24425,20710,16942

MPDATA, NUXY, 1, 1,0.354,0.3565,0.36
MPDATA, NUXY, \(1,4,0.3628,0.365,0.37,0.3774,0.3839\) ! SPECIFY TEMPERATURE ! DEPENDENT CTE
MPDATA, ALPX, 1, 1,2.427E-5,2.448E-5,2.466E-5
MPDATA, ALPX, 1, 4, 2.48E-5, 2.501E-5, 2.526E-5,
\(\Rightarrow 2.552 \mathrm{E}-5,2.579 \mathrm{E}-5\)
MPTEMP
! INITIALIZE MAT TEMP TABLE
TB,MISO, 1,8,7, !SPECIFY ISOTROPIC HARDENING
TBTEMP, \(-35+273.15\) ! SPECIFY TEMPERATURE AS -35 C
TBPT, DEFI, 9.313161E-04,37.980 ! SPECIFY STRAIN VS
TBPT,DEFI, 8.931316E-03,50.677 ! STRESS DATA POINTS AT
TBPT,DEFI, 1.693132E-02,56.307 ! THIS TEMPERATURE
TBPT, DEFI, 2.493132E-02, 60. 142
TBPT, DEFI, 4.893132E-02, 67.620
TBPT, DEFI, 8.093132E-02,73.889
TBPT, DEFI, 1.009313E-01, 76.836
TBTEMP, \(-15+273.15\) ! SPECIFY TEMPERATURE AS -15 C
TBPT, DEFI, 9.295968E-04,35.162 ! SPECIFY STRAIN VS
TBPT,DEFI, 8.929597E-03,46.917 ! STRESS DATA POINTS AT
TBPT,DEFI,1.692960E-02,52.129 ! THIS TEMPERATURE
TBPT, DEFI, 2.492960E-02,55.680
TBPT, DEFI, 4.892960E-02, 62.603
TBPT, DEFI, 8.092960E-02, 68.407
TBPT, DEFI, 1.009296E-01, 71.136
!
!
\(!\)

TBTEMP, 5+273.15 ! SPECIFY TEMPERATURE AS 5 C
TBPT,DEFI,9.271872E-04,32.344 ! SPECIFY STRAIN VS
TBPT, DEFI, \(8.927187 \mathrm{E}-03,43.158\) ! STRESS DATA POINTS AT
TBPT,DEFI,1.692719E-02.47.952 ! THIS TEMPERATURE
TBPT, DEFI, 2.492719E-02, 51.218
TBPT, DEFI, 4.892719E-02,57.586
TBPT,DEFI, 8.092719E-02, 62.925
TBPT, DEFI, 1.009272E-01, 65.435
TBTEMP, 25+273.15 ! SPECIFY TEMPERATURE AS 25 C
TBPT, DEFI, \(9.252899 \mathrm{E}-04,29.526\) ! SPECIFY STRAIN VS
TBPT, DEFI, \(8.925290 \mathrm{E}-03,39.398\) ! STRESS DATA POINTS AT
TBPT, DEFI, 1.692529E-02, 43.774 ! THIS TEMPERATURE
TBPT, DEFI, 2.492529E-02, 46.756
!
TBPT, DEFI, 4.892529E-02,52.569
TBPT, DEFI, \(8.092529 \mathrm{E}-02,57.443\)
TBPT, DEFI, 1.009253E-01,59.734
TBTEMP, 50+273.15 !SPECIFY TEMPERATURE AS 50 C TBPT, DEFI, \(9.237984 \mathrm{E}-04,26.004\)
TBPT, DEFI, \(8.923798 \mathrm{E}-03,34.698\)
TBPT, DEFI, 1.692380E-02, 38.552 ! STRESS DATA POINTS AT ! THIS TEMPERATURE
TBPT, DEFI, 2.492380E-02, 41.178
TBPT, DEFI, 4.892380E-02,46.298
TBPT, DEFI, \(8.092380 \mathrm{E}-02,50.590\)
TBPT, DEFI, 1.009238E-01,52.608
TBTEMP, 75+273.15 :SPECIFY TEMPERATURE AS 75 C
TBPT,DEFI,9.204504E-04,22.482 ! SPECIFY STRAIN VS
TBPT, DEFI, \(8.920450 \mathrm{E}-03,29.998\) ! STRESS DATA POINTS AT ! THIS TEMPERATURE
TBPT, DEFI, 1.692045E-02, 33.330
TBPT, DEFI, 2.492045E-02, 35.601
TBPT, DEFI، 4.892045E-02, 40.027
```

TBPT,DEFI, 8.092045E-02,43.738
TBPT,DEFI,1.009205E-01,45.482 !
TBTEMP, 100+273.15 !SPECIFY TEMPERATURE AS 100 C
TBPT,DEFI,9.154515E-04,18.959 ! SPECIFY STRAIN VS
TBPT,DEFI,8.915451E-03,25.298 !STRESS DATA POINTS AT
TBPT,DEFI,1.691545E-02,28.108 ! THIS TEMPERATURE
TBPT,DEFI,2.491545E-02,30.023 !
TBPT,DEFI,4.891545E-02,33.756 !
TBPT,DEFI,8.091545E-02,36.885 !
TBPT,DEFI,1.009155E-01,38.357 !
TBTEMP,125+273.15 !SPECIFY TEMPERATURE AS 125 C
TBPT,DEFI,9.111675E-04,15.437 !SPECIFY STRAIN VS
TBPT,DEFI,8.911168E-03,20.598 !STRESS DATA POINTS AT
TBPT,DEFI,1.691117E-02,22.886 ! THIS TEMPERATURE
TBPT,DEFI,2.491117E-02,24.445 !
TBPT,DEFI,4.891117E-02,27.485 !
TBPT,DEFI,8.091117E-02,30.033 !
TBPT,DEFI,1.009112E-01,31.231 !
C1 = 23.3E6 !DEFINE CREEP MODEL
C2 =6.6990E-2 !PARAMETERS
C3 = 3.300 !
C4 = 6.7515E4/8.314 !
TB,CREEP,1,,,8 !SPECIFY GENERALIZED
! GAROFALO CREEP MODEL
TBDATA,1,C1,C2,C3,C4 !SPECIFY CREEP MODEL PAR.
RATE,1 !USE CREEP STRAIN RATE EFFECT
RECTNG,0,10,0,100 !CREATE RECTANGLE
ESIZE,2 !SPECIFY ELEMENT SIZE
MSHKEY,1 ! ENFORCE MAPPED MESHING
AMESH,ALL !MESH AREA
FINISH ! EXIT PREPROCESSOR

```

\section*{SOLUTION}

The solution is obtained in twenty load steps. Nonlinear geometry effects are turned on using the nLgeom command. Automatic time stepping is turned on (AUTOTS command) so that ANSYS can adjust the time step size values for the substeps within each load step. The loads (temperature change) are interpolated for each substep from the values of the previous load step to the values of the current load step using the KBC command. A do loop is utilized for creating load step files, in which the temperature conditions are calculated and applied (BFUNIF command) before they are written to load step files (LSWRIte command). Finally, the solution is obtained using the LSSOLVE command.
```

/ SOLU
TMIN=238.15 ! DEFINE PARAMETER FOR MIN
! TEMPERATURE
! DEFINE PARAMETER FOR MAX
! TEMPERATURE
NLS=20 !DEFINE PARAMETER FOR \# OF LOAD
! STEPS

```
```

TMX=60 ! DEFINE PARAMETER FOR FINAL TIME
PI=4*ATAN(1) ! DEFINE PARAMETER FOR }
ANTYPE,STATIC !SPECIFY ANALYSIS TYPE AS STATIC
NLGEOM,ON !TURN NONLINEAR GEOMETRY EFFECTS
KBC,0 !ENFORCE RAMPED LOADING
AUTOTS,ON !TURN AUTOMATIC TIME STEPPING ON
TREF,273.15
DELTIM,1E-2,1E-3,3
NSEL,S,LOC,Y,0
D,ALL,ALL
ALLSEL
NSEL,S,LOC,Y,100
D,ALL,ALL
ALLSEL
TIM=TMX/NLS
*DO,I,1,NLS !LOOP OVER LOAD STEPS
TIME,TIM
! SPECIFY STRESS-FREE TEMPERATURE
! SPECIFY TIME STEP SIZE PARAMETERS
! SELECT NODES AT Y = 0
! CONSTRAIN ALI DOFS
! SELECT EVERYTHING
! SELECT NODES AT Y = 100
! CONSTRAIN ALL DOFS
! SELECT EVERYTHING
! FIND END TIME FOR CURRENT LOAD
STEP
! SPECIFY TIME FOR CURRENT LOAD
! STEP
TC=TMIN+(TMAX-TMIN)*SIN(PI*TIM/TMX) ! FIND TEMP FOR
! CURRENT LS
BFUNIF,TEMP,TC !SPECIFY TEMPERATURE FOR
! CURRENT LS
LSWRITE,I !WRITE LOAD STEP FILE
TIM=TIM+TMX/NLS ! FIND END TIME FOR NEXT LOAD STEP
*ENDDO !END LOOP OVER LOAD STEPS
LSSOLVE,1,NLS !SOLVE FROM LOAD STEPS
FINISH ! EXIT SOLUTION PROCESSOR

```

\section*{POSTPROCESSING}

Once the solution is obtained, the user can review a multitude of results items, including elastic, plastic, creep, and total strain components, as well as the stress components. The command input segment given below obtains contour plots for a few of the items mentioned above, in two different load steps. The results for different load steps are read using the SET command. For clarity in the contour plots, the triad and the symbols for minimum and maximum quantities are turned off. Using the /TRIAD and /PLOPTS commands. The two load steps considered here are the \(10^{\text {th }}\) and the last load steps. The \(10^{\text {th }}\) load step corresponds to the maximum temperature condition while the last load step brings the structure to its initial configuration. Although initially stress free, after applying a time-dependent temperature load and returning to the initial temperature, the structure experiences residual stresses and strains. Figure 10.16 shows the contour plot of equivalent stress at load step 20, which exhibits a significant amount of stress. Figures 10.17 and 10.18 show contour plots of plastic and creep strains at load step 20 , respectively.

\(\begin{array}{lllllll}19.296 & 22.222^{25.147} & 28.073^{30.998} & 33.923^{36.849} & 39.774 & 42.699 & 45.625\end{array}\)
Fig. 10.16 Contours of residual equivalent stress at load step \(20(t\) \(=60 \mathrm{sec}\) ).


Fig. 10.17 Contours of plastic strain in the \(y\)-direction at load step \(20(t=60 \mathrm{sec})\).


Fig. 10.18 Contours of creep strain in the \(y\)-direction at load step \(20(t=60 \mathrm{sec})\).
```

/ POST1
/TRIAD,OFF
/ PLOPTS,MINM, O
SET,10
PLNSOL,NL, PLWK
PLNSOL,S,EQV
PLNSOL, EPPL,Y
PLNSOL,EPCR,Y ! PLOT CONTOURS FOR CREEP STRAIN IN
! Y-DIR.
! READ RESULTS AT LOAD STEP }2
! PLOT PLASTIC WORK CONTOURS
! PLOT EQUIVALENT STRESS CONTOURS
! PLOT CONTOURS FOR PLASTIC STRAIN
! IN Y-DIR.
! PLOT CONTOURS FOR CREEP STRAIN IN
! Y-DIR.

```

\subsection*{10.3 Contact}

Nonlinearity due to contact conditions arises because the prescribed displacements on the boundary depend on the deformation of the structure. Furthermore, no-interpenetration conditions are enforced while the extent of the contact area is unknown.

Contact between two bodies with no bonding (such as glue, solder, or weld) is a challenging problem, mainly stemming from the lack of prior knowledge of the contact regions. Another complication is that, in most of the cases, there is friction between the contacting bodies. Both of these two factors make contact analysis highly nonlinear. In addition to these, if the materials involved exhibit nonlinear material behavior or transient effects, achieving convergence becomes even more difficult.

Before starting a contact analysis, the user needs to be aware of two main considerations: the difference in stiffness of the contacting bodies and the location of possible contact regions. If one of the contacting bodies is significantly stiffer than the other, then the rigid-to-flexible contact option can be used in ANSYS, resulting in a considerable reduction in computational time and, possibly, less difficulty in convergence. There are three contact models in ANSYS: node-to-node, node-to-surface, and surface-tosurface. Each of these models requires different types of contact elements. The node-to-node contact model is used when the contact region is accurately known a priori and when the nodes belonging to either contact surfaces are paired, thus requiring these nodes to have same the coordinates. If a large amount of sliding between the contact surfaces is expected, node-tonode contact is not suitable. The node-to-surface contact model is used when a specific point on one of the surfaces, e.g., a corner, is expected to make contact with a rather smooth surface. This model does not require accurate a priori knowledge of the contact region, and the mesh pattern on either surface does not need to be compatible. The surface-to-surface contact model is used when contact regions are not known accurately and a significant amount of sliding is expected. In this model, one of the surfaces is called the contact surface and the other, the target surface.

In a typical ANSYS contact analysis, the model is first meshed with conventional elements (beam, plane, or solid), and then the contact elements are created along the potential contact regions. Different element types are used for different contact models.

A contact analysis can be highly complicated, requiring the user to have a good understanding beforehand. It is recommended that the user read the section entitled "Contact" in the ANSYS Structural Analysis Guide as it provides a highly detailed description on how to perform a contact analysis in ANSYS, as well as several helpful hints to achieve converged solutions.

In the following subsections, two contact problems are used to demonstrate a contact analysis within ANSYS. In the first problem, the node-to-surface
contact model is used in the simulation of a block dropping on a beam. Thus, the contact analysis is combined with dynamics, leading to timedependent results. The second problem is the simulation of a nanoindentation test of a thin film deposited on a hard substrate. The problem is solved using surface-to-surface contact elements, and the thin film exhibits an elastic-perfectly plastic material behavior.

\subsection*{10.3.1 Contact Analysis of a Block Dropping on a Beam}

Consider a \(6-\mathrm{in} \times 6\)-in \(\times 1\)-in block free falling onto a 100 -in-long beam, as shown in Fig. 10.19. The block and the beam are made of the same material, with elastic modulus \(E=1 \times 10^{6} \mathrm{psi}\) and density \(\rho=0.001 \mathrm{lb} / \mathrm{in}^{3}\). The beam has a cross-sectional area of \(0.5 \mathrm{in}^{2}\), an area moment of inertia of \(I_{z z}=0.05 \mathrm{in}^{4}\), and a height of 1 in . The block is initially at 100 in above the beam, with its center point 9 in to the right of the mid-point of the beam. The goal is to obtain the time-dependent response of the beam and the block.

\section*{MODEL GENERATION}

The problem is solved using two-dimensional elements for both the beam (BEAM3) and the block (PLANE42). In modeling the block, plane stress with thickness idealization is used. Between the block (contact surface) and the beam (target surface), 40 contact elements are defined using element type CONTAC48. This is achieved by first defining two components, one containing the nodes of the target surface and the other containing the nodes of the contact surface. Finally, the GCGEN command is used to create the contact elements.


Fig. 10.19 Schematic of the block and beam (left), and the corresponding finite element mesh (right).
```

/ PREP7
ET,1,BEAM3 !ELEMENT TYPE 1 IS BEAM3
ET,2,PLANE42,.,3 !ELEMENT TYPE 2 IS PLANE42
ET,3,CONTAC48 !ELEMENT TYPE 3 IS CONTAC48
KEYOPT, 3,7,1
R,1,.5,.05,1
R,2,1
R,3,2000
MP,EX,1,1E6
MP,DENS,1,.001
K,1,0,0
K,2,100,0
L,1,2
ESIZE,,10
LMESH,1
RECTNG,56,62,100,106
ESIZE,,1
TYPE,2
REAL, 2
AMESH,ALI
TYPE, 3
REAL, 3
NSEL, S,LOC,Y,0
CM,TARGET,NODE
ESEL,S,TYPE, ,2
NSLE
CM, CONTACT, NODE
ALLSEL,ALL
GCGEN, CONTACT, TARGET
FINISH

```
```

! ENTER PREPROCESSOR

```
! ENTER PREPROCESSOR
! SPECIFY KEYOPTION 7 FOR ELEM
! SPECIFY KEYOPTION 7 FOR ELEM
! TYPE 3
! TYPE 3
! REAL CONSTANT SET 1 FOR BEAM3
! REAL CONSTANT SET 1 FOR BEAM3
! REAL CONSTANT SET 2 FOR PLANE42
! REAL CONSTANT SET 2 FOR PLANE42
! REAL CONSTANT SET 3 FOR CONTAC48
! REAL CONSTANT SET 3 FOR CONTAC48
! SPECIFY ELASTIC MODULUS
! SPECIFY ELASTIC MODULUS
! SPECIFY DENSITY
! SPECIFY DENSITY
! CREATE KEYPOINT 1 (LEFT END OF
! CREATE KEYPOINT 1 (LEFT END OF
! BEAM)
! BEAM)
! CREATE KEYPOINT 2 (RIGHT END OF
! CREATE KEYPOINT 2 (RIGHT END OF
! BEAM)
! BEAM)
! CREATE LINE FOR BEAM
! CREATE LINE FOR BEAM
! USE 10 ELEMENTS PER LINE
! USE 10 ELEMENTS PER LINE
! MESH THE LINE WITH BEAM3 ELEMENTS
! MESH THE LINE WITH BEAM3 ELEMENTS
! CREATE RECTANGULAR AREA
! CREATE RECTANGULAR AREA
! USE 1 ELEMENT PER LINE
! USE 1 ELEMENT PER LINE
! SWITCH TO ELEMENT TYPE 2
! SWITCH TO ELEMENT TYPE 2
! (PLANE42)
! (PLANE42)
! SWITCH TO REAL CONSTANT SET 2
! SWITCH TO REAL CONSTANT SET 2
! MESH THE RECTANGULAR AREA
! MESH THE RECTANGULAR AREA
! SWITCH TO ELEMENT TYPE 3
! SWITCH TO ELEMENT TYPE 3
! (CONTAC48)
! (CONTAC48)
! SWITCH TO REAL CONSTANT SET 3
! SWITCH TO REAL CONSTANT SET 3
! SELECT NODES AT Y = 0
! SELECT NODES AT Y = 0
!DEFINE COMPONENT NAMED "TARGET"
!DEFINE COMPONENT NAMED "TARGET"
! SELECT ELEMENTS WITH TYPE 2
! SELECT ELEMENTS WITH TYPE 2
! SELECT NODES ATTACHED TO SELECTED
! SELECT NODES ATTACHED TO SELECTED
! ELEMENTS
! ELEMENTS
!DEFINE COMPONENT NAMED "CONTACT"
!DEFINE COMPONENT NAMED "CONTACT"
! SELECT EVERYTHING
! SELECT EVERYTHING
! GENERATE CONTACT ELEMENTS BETWEEN
! GENERATE CONTACT ELEMENTS BETWEEN
! CONTACT AND TARGET SURFACES
! CONTACT AND TARGET SURFACES
! EXIT PREPROCESSOR
```

! EXIT PREPROCESSOR

```

\section*{SOLUTION}

The transient solution is obtained in two load steps. The first load step encompasses a very short duration ( 0.002 sec ) and is solved using 2 substeps without time integration. This is done in order to set up the initial conditions for the transient solution. In the second load step, time integration is turned on (TIMINT command) and so is the automatic time stepping option (AUTOTS command). Also, the predictor option is turned on before the solution is obtained for the second load step using the PRED command. This option allows ANSYS to make a prediction of the displacements at the beginning of each substep, thus improving convergence. Structural damping is specified in terms of viscous damping, which produces a damping matrix in the form \([\mathbf{C}]=\alpha[\mathbf{M}]+\beta[\mathbf{K}]\), in which \([\mathbf{C}],[\mathbf{M}]\), and \([\mathbf{K}]\) are the
damping, mass, and stiffness matrices, respectively. In this case, only the damping through stiffness matrix is enforced by specifying the stiffness matrix multiplier \(\beta\) using the BETAD command.
\begin{tabular}{|c|c|}
\hline / SOLU & ! ENTER SOLUTION PROCESSOR \\
\hline ANTYPE, TRANS & ! DECLARE ANALYSIS TYPE A ! TRANSIENT \\
\hline NLGEOM, ON & ! TURN NONLINEAR GEOMETRY EFFECTS ! ON \\
\hline LUMPM, ON & ! USE LUMPED MASS ASSUMPTION \\
\hline KSEL, S, KP, 1, 2 & ! SELECT KEYPOINTS 1 AND 2 \\
\hline NSLK, S & ! SELECT NODES ATTACHED TO ! KEYPOINTS \\
\hline D, ALL, ALL, 0 & ! CONSTRAIN ALL DOFS AT SELECTED ! NODES \\
\hline ALLSEL, ALL & ! SELECT EVERYTHING \\
\hline ESEL, S, ENAME, , 42 & ! SELECT ELEMENTS OF TYPE PLANE42 \\
\hline NSLE, S & ! SELECT NODES ATTACHED TO SELECTED ! ELEMENTS \\
\hline D, ALI, ALI, 0 & \begin{tabular}{l}
! CONSTRAIN ALL DOFS AT SELECTED \\
! NODES
\end{tabular} \\
\hline ALLSEL, ALL & ! SELECT EVERYTHING \\
\hline ACEL, , 386 & ! DEFINE GRAVITATIONAL ACCELERATION \\
\hline TIME, . 0002 & ! SPECIFY TIME AT THE END OF \(1^{\text {ST }}\) ! LOAD STEP \\
\hline DELTIM, . 0001 & ! SPECIFY TIME STEP SIZE \\
\hline KBC, 1 & ! ENFORCE STEPPED LOADING \\
\hline BETAD, . 000318 & \begin{tabular}{l}
! STIFFNESS MATRIX MULTIPLIER FOR \\
! DAMPING
\end{tabular} \\
\hline TIMINT, OFF & ! TURN OFF TIME INTEGRATION \\
\hline CNVTOL, F, , . 00001 & \begin{tabular}{l}
! SPECIFY FORCE CONVERGENCE \\
! TOLERANCE
\end{tabular} \\
\hline OUTRES, ALL, LAST & ! SAVE ONLY THE LAST RESULTS SET \\
\hline SOLVE & ! OBTAIN SOLUTION FOR LOAD STEP 1 \\
\hline ESEL, S, ENAME، , 42 & ! SELECT ELEMENTS OF TYPE PLANE42 \\
\hline NSLE, S & \begin{tabular}{l}
! SELECT NODES ATTACHED TO SELECTED \\
! ELEMENTS
\end{tabular} \\
\hline DDELE, ALL, ALL & ! DELETE DOF CONSTRAINTS ON ! SELECTED NODES \\
\hline ALLSEL, ALL & ! SELECT EVERYTHING \\
\hline TIME, 3 & ! TIME AT THE END OF 2ND LS AS ! 3 SEC \\
\hline DELTIM, . \(02, .0002, .02\) & ! USE INITIAL TIME STEP SIZE 0.02 ! WITH MINIMUM 0.0002 AND MAXIMUM \(!0.02\) \\
\hline AUTOTS, ON & ! TURN ON AUTOMATIC TIME STEPPING \\
\hline TIMINT, ON & ! TURN ON TIME INTEGRATION \\
\hline CNVTOL, F & ! RESET FORCE CONVERGENCE TOLERANCE \\
\hline PRED, ON & ! ACTIVATE PREDICTOR FOR INITIAL ! GUESS \\
\hline OUTRES, ALL , ALL & ! SAVE RESULTS FOR EVERY SUBSTEP \\
\hline SOLVE & ! OBTAIN SOLUTION \\
\hline FINISH & ! EXIT SOLUTION PROCESSOR \\
\hline
\end{tabular}

\section*{POSTPROCESSING}

After the solution is obtained, results are reviewed in both the General Postprocessor (results associated with the whole structure at a specific time) and Time History Postprocessor (results associated with a specific node in the structure along the entire time domain). First, in the Time History Postprocessor, the displacements in the \(y\)-direction at the mid-point of the beam (parameter MID) and at four corners of the block (parameters BL, BR, TR, and TL) are stored (NSOL command) in user-defined parameters, and they are plotted against time (PLVAR command), as shown in Fig. 10.20. Similarly, the reaction forces in the \(y\)-direction at both ends of the beam are stored (RFOR command) in user-defined parameters (parameters FL and FR), and they are plotted against time (Fig. 10.21). In both of these graphs, labels and ranges for the \(x\) - and \(y\)-axes are modified using the /AXLAB, /AUTO, /XRANGE, and /Yrange commands. After reviewing the responses of specific nodes as functions of time, the deformed shape and moment diagram at a specific time ( 1.742 sec ) are reviewed in the General Postprocessor. The results associated with time 1.742 sec are read using the SET command, and the deformed shape is obtained using the PLDISP command (Fig. 10.22). The moment results associated with nodes I and \(\boldsymbol{J}\) of each element are stored in element table items momzi and momzJ, respectively, using the ETABLE command. Finally, the PLLS command is used for plotting the moment diagram (Fig. 10.23). The command input block below includes a /WAIT command after each plot command (PLVAR and PLDISP commands), which causes ANSYS to suspend operations for a specified duration ( 2 sec in this case).
```

/POST26
NSOL, 2,6,U,Y,MID
NSOL, 3,12,U,Y,BL
NSOL, 4, 13, U,Y,BR
NSOL, 5,14,U,Y,TR
NSOL,6,15,U,Y,TL
/AXLAB,Y,DISPLACEMENT
/AXLAB,X,TIME
/AUTO,1
PLVAR, 2, 3, 4,5,6
/WAIT,2
RFOR,7,1,F,Y,FL

```
```

```
! ENTER TIME HISTORY
```

```
! ENTER TIME HISTORY
!POSTPROCESSOR
!POSTPROCESSOR
! STORE MID-PT BEAM Y-DISP TO
! STORE MID-PT BEAM Y-DISP TO
! MID
! MID
! STORE BOTTOM LEFT BLOCK Y-DISP
! STORE BOTTOM LEFT BLOCK Y-DISP
! TO BL
! TO BL
! STORE BOTTOM RIGHT BLOCK Y-
! STORE BOTTOM RIGHT BLOCK Y-
! DISP TO BR
! DISP TO BR
!STORE TOP RIGHT BLOCK Y-DISP
!STORE TOP RIGHT BLOCK Y-DISP
! TO TR
! TO TR
! STORE TOP LEFT BLOCK Y-DISP TO
! STORE TOP LEFT BLOCK Y-DISP TO
! TL
! TL
! SPECIFY Y AXIS LABEL FOR GRAPH
! SPECIFY Y AXIS LABEL FOR GRAPH
! SPECIFY X AXIS LABEL FOR GRAPH
! SPECIFY X AXIS LABEL FOR GRAPH
! AUTOMATIC FOCUS/DISTANCE FOR
! AUTOMATIC FOCUS/DISTANCE FOR
! GRAPH
! GRAPH
! PLOT VARIABLES 2-6 VS TIME
! PLOT VARIABLES 2-6 VS TIME
! WAIT 2 SECONDS BEFORE
! WAIT 2 SECONDS BEFORE
! PROCEEDING
! PROCEEDING
! STORE LEFT Y-REACTION FORCE TO
! STORE LEFT Y-REACTION FORCE TO
! FL
```

```
! FL
```

```


Fig. 10.20 Time variation of \(y\)-displacements at four corners of the block and at the midpoint of the beam.


Fig. 10.21 Time variation of reaction forces in the beam in the \(y\) direction.

\section*{Cl}

Fig. 10.22 Deformed configuration at time 1.742 sec .


Fig. 10.23 Moment diagram of the beam at time 1.742 sec .
\begin{tabular}{|c|c|}
\hline \multirow[t]{2}{*}{RFOR, 8, \(2, \mathrm{~F}, \mathrm{Y}, \mathrm{FR}\)} & ! STORE RIGHT Y-REACTION FORCE \\
\hline & ! TO FR \\
\hline /YRANGE & ! RESET Y AXIS RANGE \\
\hline /AXLAB, Y, REACTION FORCE & ! SPECIFY Y AXIS LABEL FOR GRAPH \\
\hline PLVAR, 7, 8 & ! PLOT VARIABLES 7 \& 8 VS TIME \\
\hline \multirow[t]{2}{*}{/WAIT, 2} & ! WAIT 2 SECONDS BEFORE \\
\hline & ! PROCEEDING \\
\hline \multirow[t]{2}{*}{FINISH} & ! EXIT TIME HISTORY \\
\hline & ! POSTPROCESSOR \\
\hline / POST1 & ! ENTER GENERAL POSTPROCESSOR \\
\hline SET, , , , 1.742 & ! SET RESULTS FOR TIME 1.742 SEC \\
\hline \multirow[t]{2}{*}{ETABLE, MOMZI, SMISC, 6} & ! ELEM TABLE FOR MOMENT ON NODE \\
\hline & ! I \\
\hline \multirow[t]{2}{*}{ETABLE, MOMZJ, SMISC, 12} & ! ELEM TABLE FOR MOMENT ON NODE \\
\hline & ! J \\
\hline /DSCALE, 1, 1 & ! DO NOT SCALE DISPLACEMENTS \\
\hline /TRIAD, OFF & ! TURN OFF THE TRIAD \\
\hline PLDISP, 1 & ! PLOT DEFORMED SHAPE \\
\hline \multirow[t]{2}{*}{/WAIT, 2} & ! WAIT 2 SECONDS BEFORE \\
\hline & ! PROCEEDING \\
\hline ESEL, S, TYPE, , 1 & ! SELECT BEAM ELEMENTS (TYPE 1) \\
\hline NSLE, S & ! SELECT NODES ATTACHED TO \\
\hline & ! ELEMENTS \\
\hline PLLS, MOMZI, MOMZJ & ! PLOT ELEM TABLE ITEMS ON MESH \\
\hline ALLSEL, ALL & ! SELECT EVERYTHING \\
\hline FINISH & ! EXIT GENERAL POSTPROCESSOR \\
\hline
\end{tabular}

In order to obtain an animation of the time-dependent displacement response of the system:
- Read the results associated with load step 2 (SET command) using the following menu path:

\footnotetext{
Main Menu > General Postproc > Read Results > Last Set
}
- Obtain the deformed shape (PLDISP command) using the following menu path:

Main Menu > General Positproc > Plot Results > Deformed Shape
- Plot Deformed Shape dialog box appears; select Def + undef edge radiobutton; click on \(O K\).
- Create animation using the following menu path:

Utility Menu > PlotCtrls > Animate \(>\) Over Time
- Animate Over Time dialog box appears; enter 100 for Number of animation frames. Click on \(O K\) and wait until Animation Controls Window appears.

\subsection*{10.3.2 Simulation of a Nano-indentation Test}

Nano-indentation tests are commonly used for evaluation of the response of thin films. Consider a sol-gel layer deposited on a glass substrate, which is indented by means of a conical diamond indenter as shown in Fig. 10.24. The thickness values for the sol-gel film and the glass substrate are \(2 \mu \mathrm{~m}\) and \(6 \mu \mathrm{~m}\), respectively. The indenter has an angle of \(68^{\circ}\) measured from the axis of rotation. In order to correctly simulate the indentation phenomenon, a contact analysis is utilized. For this purpose, contact elements (CONTA174) are placed along the top surface of the film and target elements (TARGE170) are used along the bottom surface of the indenter. The indentation is simulated by applying displacement boundary conditions in the \(y\)-direction to the nodes along the bottom surface of the substrate. Consequently, the top surface of the film is pressed against the bottom surface of the indenter, thus exerting the contact elements against the target elements. The contact is assumed to be frictionless. The indentation is performed using several displacement steps, each of which is written to a load step file. Both loading and unloading are simulated. Displacement step sizes for loading and unloading are \(0.04 \mu \mathrm{~m}\) and \(0.03 \mu \mathrm{~m}\), respectively. All three materials are modeled using SOLID45 elements. Since the problem possesses symmetry with respect to the \(y\)-axis, only one octant \(\left(1 / 8^{\text {th }}\right)\) of the geometry is modeled. Normal displacements are constrained along the symmetry planes, and \(x\) - and \(z\)-displacements are constrained along the axis of rotation. The top surface of the indenter is constrained in all directions. The sol-gel film exhibits elastic-perfectly plastic behavior with yield stress of 700 MPa while the diamond indenter and the glass substrate are both elastic. The elastic modulus and Poisson's ratio values for the constituent materials are given in Table 10.4. The goal is to obtain indentation vs. force response for the film.


Fig. 10.24 Sol-gel film on glass substrate, and diamond indenter.
Table 10.4 Material properties used in nano-indentation simulation.
\begin{tabular}{l|c|c|c}
\hline & \begin{tabular}{c} 
Elastic \\
Modulus \\
(GPa)
\end{tabular} & \begin{tabular}{c} 
Poisson's \\
Ratio
\end{tabular} & \begin{tabular}{c} 
Material \\
Reference \\
Number
\end{tabular} \\
\hline Glass substrate & 75 & 0.23 & 1 \\
Sol-gel film & 4.5 & 0.35 & 2 \\
Diamond indenter & 1141 & 0.07 & 3 \\
\hline
\end{tabular}

\section*{MODEL GENERATION}

In the solid modeling phase, a bottom-up approach is used. The solid model is created, starting with keypoints, then lines, areas, and volumes. The elas-tic-perfectly plastic behavior of the thin film is incorporated using the bilinear isotropic hardening rule (TB command with BISO option) with a tangent modulus of zero. The model is first generated in the \(x-y\) plane and meshed using two-dimensional elements (PLANE42), after which the VROTAT command is used to generate the three-dimensional mesh by rotating the meshed areas about the \(y\)-axis by \(45^{\circ}\). Before issuing the VROTAT command, the default element type attribute must be changed to
the one for three-dimensional elements (element type 2 for SOLID45 in this case). Figure 10.25 shows an isometric view of the mesh used in this analysis.
```

/FILNAM, NANO
/PREP7
ET,1,42
ET,2,45
ET,3,TARGE170
ET,4,CONTA174
MP, EX,1,75000
MP,NUXY,1,0.23
MP,EX,2,4500
MP,NUXY,2,0.35
TB,BISO,2,1
TBTEMP,0
TBDATA, 1,700,0
MP,EX, 3,1141000
MP,NUXY,3,0.07
K,1
K,2,4
K, 3,10
K,4, 6
K,5,4,6
K,6,10,6
K,7, 7.2
K,8,4,7.2
K,9,10,7.2
K,10, , 8
K,11,4,8
K,12,10,8
L,1,2
L,2,3
L,4,5
L,5,6
L,7,8
L,8,9
L,10,11
L,11,12
L,1,4
L,2,5
L, 3,6
L,4,7
L,5,8
L,6,9
L,7,10
L,8,11
L,9,12
AL, 1, 3,9,10
AL, 2, 4, 10,11
AL, 3,5,12,13
AL, 4, 6, 13,14
AL, 5, 7, 15,16
AL, 6, 8, 16,17

```
```

SPECIFY JOBNAME

```
SPECIFY JOBNAME
ENTER PREPROCESSOR
ENTER PREPROCESSOR
ELEMENT TYPE 1 IS PLANE42
ELEMENT TYPE 1 IS PLANE42
!ELEMENT TYPE 2 IS SOLID45
!ELEMENT TYPE 2 IS SOLID45
! ELEMENT TYPE 3 IS TARGE170
! ELEMENT TYPE 3 IS TARGE170
! ELEMENT TYPE 4 IS CONTA174
! ELEMENT TYPE 4 IS CONTA174
!GLASS SUBSTRATE MAT PROPS
!GLASS SUBSTRATE MAT PROPS
!
! SOL GEL FILM MAT PROPS
! SOL GEL FILM MAT PROPS
!
! BILINEAR ISOTROPIC HARDENING RULE
! BILINEAR ISOTROPIC HARDENING RULE
!
! YIELD STRENGTH
! YIELD STRENGTH
! DIAMOND INDENTER MAT PROPS
! DIAMOND INDENTER MAT PROPS
!
! CREATE KEYPOINTS
! CREATE KEYPOINTS
! CREATE LINES
! CREATE AREAS
```



Fig. 10.25 Isometric view of the finite element mesh used for the nano-indentation simulation.

```
LESIZE,7,.,15 !SPECIFY LINE DIVISIONS
LESIZE,5,,,15
LESIZE, 3,,,15
LESIZE,1,,,15
LESIZE, 8, , ,10,4
LESIZE,6, , 10,4
LESIZE, 4, , ,10,4
LESIZE, 2, , ,10,4
LESIZE, 9, , 10,1/4
LESIZE, 10, , ,10,1/4
LESIZE, 11, , ,10,1/4
LESIZE,12,, ,8
LESIZE,13,, , 8
LESIZE,14,, , 8
LESIZE, 15, , , 8
LESIZE,16, , 8
LESIZE,17, , 8
MSHKEY,1 ! ENFORCE MAPPED MESHING
TYPE,1 !SWITCH TO ET 1
MAT,1
AMESH,1,6
TYPE,2
ESIZE,,4 !SPECIFY # OF ELEMS FOR SWEEP
VROTAT, 1, 2, 3,4,5,6,1,10,45 ! SWEEP AREAS TO CREATE
    ! VOLUME
```

```
NSEL,S,LOC,Y,6,8 ISELECT NODES AT 6 mm < Y < 8 \mum
ESLN,S,1 ! SELECT ELEMENTS WITH SELECTED
! NODES
EMODIF,ALL,MAT,2 ! MODIFY ELEMS TO BE GLASS
TYPE,4 ! SWITCH TO ET 4
AMESH,22 ! CREATE CONTACT AREA MESH
ESEL,S,TYPE,,4 !SELECT ELEMENTS WITH ET 4
ESURF , REVE ! ADJUST OUTWARD NORMAL OF ELEMENTS
! GENERATE SOLID MODEL FOR INDENTER
! CREATE KEYPOINTS
K,26,,10.5
K,24,3.7,9.5
K,25,3.7,10.5
L,23,24 ! CREATE LINES
L,24,25
L,25,26
L,26,23
AL,40,41,42,43
LESIZE,41,,,5
LESIZE,43,,,5
LESIZE,40, , 15
LESIZE,42,, ,15
TYPE, 1
MAT, 3
AMESH,27
TYPE,2
ESIZE,,4 !SPECIFY # OF ELEMS FOR SWEEP
VROTAT,27,,,,,,23,26,45!SWEEP AREAS TO CREATE VOLUME
TYPE,3 !SWITCH TO ET 3
AMESH,28 ! CREATE TARGET AREA MESH
AILSEL,ALL !SELECT EVERYTHING
FINISH ! EXIT PREPROCESSOR
```


## SOLUTION

As mentioned previously, the top surface of the diamond indenter is constrained in all directions. In addition, the vertical planes are not allowed to move in their normal direction due to the symmetry boundary conditions imposed by using the DSYM command. The indentation is simulated by prescribing a uniform $y$-displacement along the bottom surface of the glass substrate, which causes the film to contact the indenter, starting the indentation process. This displacement is applied in increments of $0.04 \mu \mathrm{~m}$ for the loading phase, and its values are stored in the array parameter DIS. The array parameter is created/initialized using the *DIM command, with an array size of 57 . The first 37 entries of this array correspond to the loading phase displacements and the following 20 entries are for the unloading phase displacements. Once all the displacement values are stored in the array DIS, 57 load step files are written (LSWRITE command) utilizing a
do loop. Nonlinear geometry effects are turned on (NLGEOM command), as well as automatic time stepping (AUTOTS command). A full NewtonRaphson method is utilized without the adaptive descent option (NROPT command). The maximum number of equilibrium iterations in each substep is set to 100 using the NEQIT command. Prior to initiating the solution using the LSSOLVE command, the two-dimensional elements must be unselected, as the sole purpose of their existence is for meshing.

| /SOLU | ! ENTER SOLUTION PROCESSOR |
| :---: | :---: |
| SOLCONTROL, 0 | ! TURN OFF SOLUTION CONTROLS |
| ANTYPE, STATIC | ! SPECIFY ANALYSIS TYPE AS STATIC |
| NSEL, S, LOC, Z | ! SELECT NODES AT $\mathrm{Z}=0$ |
| DSYM, SYMM, Z, 0 | ! APPLY SYMMETRY BC ABOUT X-Y PLANE |
| CLOCAL, 11, , , , , , 45 | ! DEFINE LOCAL CS |
| NSEL, S, LOC, Z | ! SELECT NODES AT z = 0 (IN LOCAL ! CS) |
| DSYM, SYMM, 2,11 | ! APPLY SYMMETRY BC IN LOCAL CS |
| CSYS, 0 | ! SWITCH TO GLOBAL CARTESIAN CS |
| NSEL, S,LOC, Y, 10.5 | ! SEIECT NODES AT Y = 10.5 |
| D, ALL, UY, 0 | ! CONSTRAIN Y-DISP AT SELECTED ! NODES |
| NSEL, S, LOC, X | ! SELECT NODES AT $\mathrm{X}=0$ |
| NSEL, R, LOC, Z | ! RESELECT NODES AT $\mathrm{z}=0$ |
| D, ALL, UX | ! CONSTRAIN X-DISP AT SELECTED ! NODES |
| D, ALL , UZ | ! CONSTRAIN Z-DISP AT SELECTED ! NODES |
| *DIM, DIS, ARRAY,57 | ! INITIALIZE ARRAY PARAMETER DIS |
| $\mathrm{A}=0$ | ! INITIALIZE PARAMETER A |
| *DO, I, 1,36 | ! START DO LOOP FOR LOADING |
| $\operatorname{DIS}(\mathrm{I})=\mathrm{A}$ | ! STORE DISP VALUE IN DIS FOR ! CURRENT LS |
| $\mathrm{A}=\mathrm{A}+0.04$ | ! UPDATE PARAMETER A |
| *ENDDO | ! END DO LOOP ON LOADING |
| $\mathrm{B}=0.03$ | ! INITIALIZE PARAMETER B |
| *DO, I, 37,57 | ! START DO LOOP FOR UNLOADING |
| $\mathrm{DIS}(\mathrm{I})=\mathrm{A}-\mathrm{B}$ | ! STORE DISP VALUE IN DIS FOR ! CURRENT LS |
| $\mathrm{B}=\mathrm{B}+0.03$ | ! UPDATE PARAMETER B |
| *ENDDO | ! END DO LOOP ON UNLOADING |
| NLGEOM, ON | ! TURN ON NONLINEAR GEOMETRY ! EFFECTS |
| AUTOTS, ON | ! TURN ON AUTOMATIC TIME STEPPING |
| OUTRES, , 1 | ! SAVE RESULTS FOR LAST SUBSTEP OF <br> ! EACH LS |
| NROPT, FULL, , OFF | ! USE FULL NEWTON-RAPHSON WITH NO <br> ! ADAPTIVE DESCENT |
| NEQIT, 100 | ! USE MAXIMUM 100 EQUILIBRIUM ! ITERATIONS |
| ALLSEL | ! SELECT EVERYTHING |
| *DO, I, 1, 57 | ! START DO LOOP FOR WRITING LOAD |
| NSEL, S, LOC, Y | ! SELECT NODES AT Y = 0 |

```
D,ALL,UX
D,ALL,UZ
D,ALL,UY,DIS(I)
ALLSEL
LSWRITE,I
* ENDDO
ESEL,U,TYPE,,1
LSSOLVE,1,57
FINISH
```

```
! CONSTRAIN X-DISP AT SELECTED
```

! CONSTRAIN X-DISP AT SELECTED
! NODES
! NODES
!CONSTRAIN Z-DISP AT SELECTED
!CONSTRAIN Z-DISP AT SELECTED
! NODES
! NODES
!SPECIFY Y-DISP ALONG THE BOTTOM
!SPECIFY Y-DISP ALONG THE BOTTOM
! SURFACE OF THE SUBSTRATE
! SURFACE OF THE SUBSTRATE
SELECT EVERYTHING
SELECT EVERYTHING
!WRITE LOAD STEP FILE
!WRITE LOAD STEP FILE
! END DO LOOP ON WRITING LOAD STEPS
! END DO LOOP ON WRITING LOAD STEPS
!UNSELECT ELEMENT TYPE 1
!UNSELECT ELEMENT TYPE 1
SOLVE FROM LS FILES (1 TO 57)
SOLVE FROM LS FILES (1 TO 57)
! EXIT SOLUTION PROCESSOR

```
! EXIT SOLUTION PROCESSOR
```


## POSTPROCESSING

With the goal of obtaining the load vs. indentation depth response of the sol-gel thin film deposited on a glass substrate, the following command input segment is used. Load and indentation depth values for each load step are extracted within a do loop, in which the results associated with the current load step are read using the SET command. The parameter SUM is the total reaction force in the $y$-direction along the bottom surface of the glass substrate. This quantity must be identical to the force exerted by the indenter to the top surface of the film because of equilibrium of forces. The command *GET is used on numerous occasions to retrieve model and results information about the nodes. For each load step, the applied indentation depth from array parameter DIS, and the corresponding total reaction force stored in parameter SUM are written to the file NANO_RF_D.OUT using a combination of /OUTPUT and *VWRITE commands. Figure 10.26 shows the resulting load vs. indentation depth response.


Indentation Depth (micrometer)
Fig. 10.26 Load vs. indentation depth for sol-gel film on glass substrate.

| / POST1 | ! ENTER GENERAL POSTPROCESSOR |
| :---: | :---: |
| *DO, J, 1, 57 | ! LOOP OVER LOAD STEPS |
| SET, J | ! READ RESULTS SET |
| NSEL, S, LOC, Y | ! SELECT NODES AT Y = 0 |
| *GET, NUMNNOD, NODE, 0, COUNT | ! STORE \# OF NODES IN NUMNOD |
| *GET, CURNOD, NODE, 0, NUM, MIN | ! STORE MIN NODE \# TO CURNOD |
| SUM=0 | ! INITIALIZE TOTAL REACTION F |
| *DO, I, 1, NUMNOD | ! LOOP OVER SELECTED NODES |
| *GET, RFY, NODE, CURNOD , RF, FY | ! STORE Y REACT. FORCE IN RFY |
| SUM=SUM+RFY | ! UPDATE TOTAL REACTION FORCE |
| CURNOD=NDNEXT (CURNOD) | ! UPDATE CURRENT NODE NUMBER |
| * ENDDO | ! END LOOP OVER SELECTED NODES |
| DISJ $=$ DIS ( J ) | ! STORE CURRENT INDENTATION |
|  | ! DEPTH TO PARAMETER DISJ |
| SUM=SUM / 100 | ! CONVERT TO MILLINEWTONS |
| / OUTPUT, NANO_RF__D, OUT, , APPEND! REDIRECT OUTPUT TO FILE |  |
| *VWRITE, DISJ, SUM | ! WRITE DISJ AND SUM TO FILE |
| (E16.8,5X, E16.8) | : FORMAT STATEMENT |
| / OUTPUT | ! REDIRECT OUTPUT TO OUTPUT W |
| * ENDDO | ! END LOOP OVER LOAD STEPS |

## Chapter 11

## ADVANCED TOPICS IN ANSYS

Selected advanced topics are discussed in this chapter. First, coupled degrees of freedom and constraint equations are explained. Discussions on submodeling and superelements are included next, with one example problem for each. Finally, a brief section on how to interact with external files from within the ANSYS environment is followed by a section on modification of the ANSYS GUI.

### 11.1 Coupled Degrees of Freedom

In certain engineering problems, the behavior of some of the unknown degrees of freedoms may be known. For example, certain points (nodes) may be expected to have the same displacement in a certain direction. One can take advantage of this behavior and enforce it in order to achieve an accurate solution with minimum computational resources. If a particular degree of freedom at several nodes is expected to have the same unknown value, these degrees of freedoms can be coupled. The use of coupled degrees of freedom is clarified with the following example: a long, hollow cylinder subjected to internal pressure (see Fig. 11.1)

This problem can be solved using two different approaches: (i) plane strain idealization and (ii) axisymmetry. Use of plane strain idealization involves meshing on the $x-y$ plane and does not require imposition of additional constraints. Figure 11.2 shows the plane strain model with applied boundary conditions (quarter-symmetry is used).

In order to utilize axisymmetry, the mesh is generated on the $x$-z plane, as shown in Fig. 11.3. Along the left vertical boundary, the inner pressure is specified while the right vertical boundary is traction free. Based on the problem definition (long in the $z$-direction), it is known that the $z$-displacement on the $x-y$ plane is uniform but its value is unknown. This condition is enforced by constraining the $z$-displacements along the bottom boundary and coupling the $z$-displacements of the nodes along the top surface.


Fig. 11.1 A long, hollow cylinder subjected to internal pressure.


Fig. 11.2 Plane strain idealization with applied boundary conditions.


Fig. 11.3 Axisymmetric model with applied boundary conditions.

Both plane strain and axisymmetric models are valid. The solution obtained using plane strain idealization produces identical results in the circumferential direction (when viewed in the cylindrical coordinate system). Although both models may have a similar number of nodes, the accuracy of the results in the radial direction provided by the axisymmetric solution is much higher than that of the plane strain solution.

A set of coupled degrees of freedom is defined using the CP command, which has the following syntax:

```
CP, NSET, Lab, NODE1, NODE2, . . ., NODE17
```

in which NSET is the coupled degree of freedom set number, Lab is the degree of freedom label (e.g., UX, UY, etc.) and NODE1 through NODE17 are the nodes to be coupled. It is common to first select the nodes to be coupled and then assign ALL for the argument NODE1. For example, the $z$ displacements of all nodes along $y=1$ are coupled, using the following command input:

```
NSEL, S, LOC, Y, 1
CP,1,UZ,ALL
```

The coupled degrees of freedom set must not include any degrees of freedom that are specified as boundary conditions; they must be unknown.

### 11.2 Constraint Equations

The coupled degrees of freedom feature discussed in the previous section is a subset of the constraint equations. When coupling two degrees of freedom, say $x$-displacements at nodes 2 and 4, the following constraint equation is enforced:

$$
\begin{equation*}
u_{x}^{(2)}-u_{x}^{(4)}=0 \tag{11.1}
\end{equation*}
$$

In ANSYS, it is also possible to specify more general constraint equations. The general form of these equations is

$$
\begin{equation*}
c_{0}=\sum_{i=1}^{N} c_{i} u_{j}^{(i)} \tag{11.2}
\end{equation*}
$$

in which $c_{i}(i=0, \ldots, N)$ are constant coefficients, $u_{j}^{(i)}$ denotes a degree of freedom at node $i$ in the $j$-direction $(j=x, y, z)$, and $N$ is the number of degrees of freedom involved in the constraint equation. The constraint equations are specified using the CE command with the following syntax:

```
CE, NEQN, CONST,NODE1, Lab1, C1, NODE2, Lab2, C2,NODE3, Lab3,C3
```

in which NEQN is the constraint equation reference number; CONST, C1, C2, and C3 are the coefficients $c_{i}$ of Eq. (11.2); NODE1, NODE2, and NODE3 denote the node numbers involved in the constraint equation; and, finally, Lab1, Lab2 and Lab3 are the degree of freedom labels. The use of this command is explained through an example.

Consider five equally spaced nodes, as shown in Fig. 11.4. Assume that the $x$-displacements of these nodes are expected to exhibit anti-symmetric behavior. This implies two constraint equations, which are written as

$$
\begin{align*}
& 0=1 \times u_{x}^{(1)}+(-1) \times u_{x}^{(5)} \\
& 0=1 \times u_{x}^{(2)}+(-1) \times u_{x}^{(4)} \tag{11.3}
\end{align*}
$$

The equivalent command line input using the CE command is given by

$$
\begin{aligned}
& C E, 1,0,1, U X, 1,5, U X,-1 \\
& C E, 2,0,2, U X, 1,4, U X,-1
\end{aligned}
$$

Constraint equations are particularly useful when a rigid connection between two separate regions is needed. For example, the modeling of a composite sandwich panel where the core is enclosed by two thin facesheets on the top and bottom surfaces is ideal for using rigid constraints. In this case, the facesheets are modeled using shell elements while solid brick elements are used


Fig. 11.4 Five equally spaced nodes with expected anti-symmetric displacement.
for the core, as shown in Fig. 11.5. However, the shell elements, by default, place the nodes at the mid-plane of the plate. If the nodes of the shell elements were shared with the solid brick elements, half of the facesheet element in the thickness direction would penetrate into the solid brick element, thus creating a wrong representation of the actual structure. The use of shell elements whose mid-plane is offset by means of rigid constraints removes this problem.

Rigid constraints are specified using the CERIG command with the following syntax:

CERIG, MASTE, SLAVE, Ldof, Ldof2, Ldof3, Ldof4, Ldof5
in which MASTE and SLAVE are the node numbers for the master and slave nodes, respectively, and Ldof,..., Ldof5 are the degree of freedom labels. Once the constraint is specified, only one of these two nodes is retained in the matrix system of equations, which is called the master node, while the remaining node is called the slave node. Generally, only the argument Ldof is used as it provides three labels, ALL, UXYZ, and RXYZ, in addition to the usual degree of freedom labels (UX, UY, UZ, ROTX, ROTY, and ROTZ). The use of the CERIG command is demonstrated by an analysis of a sandwich composite panel under transverse pressure.

The construction of a sandwich panel with facesheets and a core is illustrated in Fig. 11.6. The mid-plane of the sandwich panel coincides with the $(x, y)$ plane of the reference coordinate system. As shown in Fig. 11.6, the panel has a uniform thickness of $2 h$. The thicknesses of the core and facesheets are denoted by $h_{c}$ and $h_{f}$, respectively. The panel has a square planar geometry described by $2 a=2 b=6 \mathrm{in}$. The facesheets at the bottom and top of the core are of equal thickness, $h_{f}=0.01 \mathrm{in}$, and the core thickness is $h_{c}=0.5 \mathrm{in}$. The facesheets are homogeneous, elastic, and orthotropic material layers. The elastic modulus and Poisson's ratio values are


Fig. 11.5 A sandwich panel and corresponding finite element model utilizing rigid constraints between the shell and solid elements.



Fig. 11.6 Sandwich composite panel under transverse pressure.
defined with respect to the reference frame as $E_{x x}=E_{y y}=9.58 \times 10^{6} \mathrm{psi}$, $E_{z z}=1.66 \times 10^{6} \mathrm{psi}, G_{x y}=709 \times 10^{3} \mathrm{psi}, G_{y z}=G_{x z}=731 \times 10^{3} \mathrm{psi}, v_{x y}=$ 0.033 , and $v_{y z}=v_{x z}=0.352$. The core is isotropic with Young's modulus of $E_{c}=18.5 \times 10^{3} \mathrm{psi}$ and Poisson's ratio of $\nu_{c}=0.3$.

The panel is on roller supports along all edges and is subjected to a square step load of 1000 psi in the positive $z$-direction, applied over a square area ( $2 \times 2 \mathrm{in}^{2}$ ) centered along the bottom surface of the panel. Due to the quarter-symmetry of the problem, symmetry boundary conditions are applied along the $x=0$ and $y=0$ planes.

```
/PREP7
ET,1,99
KEYOPT,1,5,2
KEYOPT, 1,8,1
KEYOPT,1,11,0
ET,2,95
R,1
RMODIF,1,1,1,0,0,0,0,0 !
RMODIF,1,13,1,0,.01, !SPECIFY THICKNESS
UIMP,1,EX,EY,EZ,9.582E6,9.582E6,1.66E6, ! SPECIFY MAT
                                    ! PROPS
UIMP, 1, PRXY, PRYZ, PRXZ,0.03192,0.35231,0.35231, ! FOR
UIMP,1,GXY,GYZ,GXZ,709E3,731E3,731E3, ! FACESHEETS
UIMP,2,EX, , ,18.51E3, !SPECIFY MAT PROPS FOR CORE
UIMP, 2,NUXY, , ,.3,
LOCAL,11,0 !DEFINE A LOCAL CS
ESYS,11 !ALIGN ELEMENT CS WITH CS 11
W=6 ! PARAMETER FOR X-LENGTH OF PANEL
H=6 ! PARAMETER FOR Y-LENGTH OF PANEL
D=1 ! PARAMETER FOR SUBDIVISIONS
TF1=0.01 !TOP FACESHEET THICKNESS
TF2=0.01 ! BOTTOM FACESHEET THICKNESS
TC=0.5 ! CORE THICKNESS
P=1000 !APPLIED PRESSURE PSI
DIV1=10 ! PARAMETERS FOR NUMBER OF
DIV2=7 !DIVISIONS AT VARIOUS LOCATIONS
DIV3=5
DIV4=DIV1/2
RECTNG,,W/2,,H/2, ! CREATE RECTANGLES
RECTNG, D, ,H/2 !
RECTNG,,W/2,,D !
RECTNG, , 2*D, ,H/2 !
RECTNG, ,W/2,,2*D !
AOVLAP,ALL !OVERLAP ALL AREAS
ALLSEL !SELECT EVERYTHING
LSEL,S,LOC,X,D/2 ! SELECT LINES AT X = D/2
LSEL,A,LOC,Y,D/2 ! ALSO SELECT LINES AT Y = D/2
LESIZE,ALL,,,DIV1 !SPECIFY # OF DIVISIONS
LSEL,S,LOC,X,3*D/2 ! SELECT LINES AT X = 3D/2
LSEL,A,LOC,Y,3*D/2 ! ALSO SELECT LINES AT Y = 3D/2
LESIZE,ALL,,,DIV2 !SPECIFY # OF DIVISIONS
LSEL,S,LOC,X,5*D/2 !SELECT LINES AT X = 5D/2
LSEL,A,LOC,Y,5*D/2 !ALSO SELECT LINES AT Y = 5D/2
```



| SLAVE=NODE (NX (CURNOD), NY (CURNOD), -TC-TF1) ! STORE NODE \# |  |
| :---: | :---: |
|  | ! OF CORRESPONDING CORE NODE |
| CERIG, CURNOD, SLAVE, UXYZ ! APPLY RIGID CONSTRAINT |  |
| NSEL, S,LOC, $\mathrm{Z},-\mathrm{TC}-T \mathrm{~T} 1 / 2$ ! SELECT BOTTOM FACESHEET NODES |  |
| CURNOD=NDNEXT (CURNOD) ! UPDATE CURRENT NO |  |
| * ENDDO | ! END LOOP ON BOTTOM FACESHEET |
|  | ! NODES |
| FINISH | ! EXIT PREPROCESSOR |
| /SOLU | ! ENTER SOLUTION PROCESSOR |
| NSEL, S, LOC, X, 0 | ! SELECT NODES AT $X=0$ |
| D, ALL, UX | ! CONSTRAIN UX AT SELECTED NODES |
| NSEL, S, LOC, Y, 0 | ! SELECT NODES AT Y = 0 |
| D, ALL, UY | ! CONSTRAIN UY AT SELECTED NODES |
| NSEL, S, LOC, Y, H/2-1E-2 | ! SELECT NODES AT Y $=\mathrm{H} / 2-1 \mathrm{E}-2$ |
| D, ALL, UX | ! CONSTRAIN UX AT SELECTED NODES |
| D, ALL, UZ | ! CONSTRAIN UZ AT SELECTED NODES |
| NSEL, S, LOC, X, W/ 2-1E-2 | ! SELECT NODES AT $\mathrm{X}=\mathrm{H} / 2-1 \mathrm{E}-2$ |
| D, ALL, UY | ! CONSTRAIN UY AT SELECTED NODES |
| D, ALI, UZ | ! CONSTRAIN UZ AT SELECTED NODES |
| ALLSEL | ! SELECT EVERYTHING |
| NSEL, S, LOC, Z, -TC-TF1 | ! SELECT NODES AT Z $=-\mathrm{TC}-\mathrm{TF} 1$ |
| NSEL, R, LOC, X, 0, 1 | ! RESELECT NODES AT $\mathrm{X}=0$ \& 1 |
| NSEL, R, LOC, Y, 0, 1 | $!$ RESELECT NODES AT $Y=0$ \& 1 |
| SF, ALL, PRES, P | ! APPLY PRESSURE ON SELECTED NODES |
| ALLSEL | ! SELECT EVERYTHING |
| /NERR, 5,100000 | ! INCREASE ALLOWED \# OF ERROR MSG |
| SOLVE | ! OBTAIN SOLUTION |
| SAVE | ! SAVE MODEL |
| FINISH | ! EXIT SOLUTION PROCESSOR |
| / POST1 | ! ENTER GENERAL POSTPROCESSOR |
| PLNSOL, S, EQV | ! OBTAIN EQUIVALENT STRESS CONTOURS |
| PLNSOL, S, Z | ! OBTAIN STRESS CONTOURS IN Z-DIR |

Figures 11.7 and 11.8 show the contours for equivalent stress and stress in the $z$-direction, respectively, obtained through the command input given above.

### 11.3 Submodeling

Submodeling is a method used for obtaining more accurate results in a specific region of the domain. It requires an existing model, which is called the global model, and the corresponding solution. The global model uses a coarse mesh to minimize computational cost while the submodel has a much finer mesh to enhance (capture) accuracy. The solution to the global model provides the boundary conditions for the submodel. Consider the gear shown in Fig. 11.9. If the primary interest is the level of stress intensity around the region indicated in Fig. 11.9, then a separate finite element


Fig. 11.7 Contours of equivalent stress.


Fig. 11.8 Contours of normal stress in the $z$-direction.


Fig. 11.9 Meshes for the global model and submodel.
model (submodel) of that region is generated with a refined mesh (also shown in Fig. 11.9). The boundaries of the submodel that are not external boundaries are called cut boundaries. The degree of freedom results from the global model are interpolated and applied as boundary conditions along the cut boundaries. The boundary conditions along the external boundaries of the submodel must be the same as those of the global model. Cut boundaries do not need to coincide with the global element boundaries.

The steps required in a typical submodeling analysis are given as follows:

- Create global model and obtain solution.
- Save global model and solution.
- Create submodel and save.
- Select nodes along cut boundaries and write to a file.
- Resume from the global model.
- Retrieve global solution.
- Perform cut boundary interpolation (writes boundary conditions to cut boundary condition file).
- Resume from the submodel.
- Read cut boundary condition file.
- Apply remaining loads (body loads and/or conditions along external boundaries).
- Obtain submodel solution.

The following example-submodeling analysis of a plate with a circular hole-demonstrates how the submodeling method is used in an analysis.

A thin, square plate ( $10 \times 10 \mathrm{in}^{2}$ ) with a circular hole (radius $r=2.5 \mathrm{in}$ ) is subjected to a uniformly distributed tensile loading ( 1000 psi ) in the vertical direction along its top edge while being fixed along the bottom edge (Fig. 11.10). Because the plate is thin and subjected to in-plane loads, plane stress idealization is valid. The material properties are elastic modulus, $E=10 \times 10^{6} \mathrm{psi}$, and Poisson's ratio $v=0.25$. The goal is to perform a submodeling analysis around the hole to obtain more accurate stress results.

## GLOBAL MODEL ANALYSIS

The global model analysis is straightforward. However, in order for the submodeling analysis to be successful, it is crucial that distinct jobnames for both the global model and the submodel be specified (/FILNAM command). The jobname for the global model is specified as GLO in this example. Once the global model solution is obtained, the SAVE command is issued, which saves both the model and the solution.

```
/FILNAM,GLO
```

! SPECIFY JOBNAME AS 'GLO'
! ENTER PREPROCESSOR
! USE PLANE42 ELEMENT TYPE
! SPECIFY ELASTIC MODULUS
! SPECIFY POISSON'S RATIO
! CREATE RECTANGLE
! CREATE CIRCLE
! SUBTRACT CIRCLE FROM RECTANGLE
! SWITCH TO GLOBAL CYLINDRICAL CS
! SELECT LINES AT RADIUS 2.5
! SPECIFY LINE DIVISIONS
! SWITCH TO GLOBAL CARTESIAN CS
! SELECT EVERYTHING
! SPECIFY LINE DIVISIONS
! CONCATENATE LINES 2 AND 3
! ENFORCE MAPPED MESHING
! MESH AREA
! REFLECT MESHED AREA WRT YZ-PLANE
! REFLECT MESHED AREAS WRT XZ-PLANE
! MERGE ALL DUPLICATE ENTITIES
! EXIT PREPROCESSOR
! ENTER SOLUTION PROCESSOR
! SELECT NODES AT Y = -5
! CONSTRAIN ALL DOFS AT SELECTED
! NODES
! SELECT NODES AT Y = 5
! APPLY PRESSURE AT SELECTED NODES
! SELECT EVERYTHING


Fig. 11.10 Plate with a circular hole subjected to uniform tensile loading.

```
SOLVE !OBTAIN GLOBAL MODEL SOLUTION
SAVE ! SAVE MODEL AND RESULTS
FINISH ! EXIT SOLUTION PROCESSOR
/POST1 ! ENTER GENERAL POSTPROCESSOR
PLNSOL,S,Y !OBTAIN STRESS CONTOURS IN Y-
| DIRECTION
FINISH ! EXIT GENERAL POSTPROCESSOR
```


## SUBMODEL ANALYSIS

As part of the submodeling analysis, the mesh for the area enclosed by dashed lines indicated in Fig. 11.10 is created; the submodel encompasses this area. Figure 11.11 shows the mesh for the global model and the region where the submodel is created. The refined mesh for the submodel is shown in Fig. 11.12. The jobname for the submodel is specified as SUB in this example. All of the lines except the circular hole boundary are the cut boundaries. After creating the mesh for the submodel, nodes along the cut boundaries are selected and written to an external ASCII file (SBS.NOD) using the NWRITE command, after which the submodel is saved (SAVE command). The cut boundary interpolation can be performed using the cut boundary nodes in the SBS.NOD file and the global model solution in the results file (GLO.RST). For this purpose, the global model is loaded (RESUME command), and results are read from the results file GLO.RST.


Fig. 11.11 Mesh for the global model (left), and the location of the submodel (right).


Fig. 11.12 Mesh for the submodel (overlaid on the global model mesh).

Using the CBDOF command, the cut boundary interpolation is performed, which writes the interpolated results in the cut boundary condition file (SBS.CBD in this example). The submodel is then loaded and the SBS.CBD file is read as input.

Stresses in the $y$-direction obtained from the global model and submodel are plotted along the $x$-axis, as shown in Fig. 11.13.

```
/CLEAR,START
/FILNAM,SUB
/PREP7
ET,1, PLANE42
MP, EX,1,10E6
MP,NUXY,1,0.25
RECTNG,0,3,0,3
PCIRC,2.5
```

```
! CLEAR DATABASE AND START NEW
```

! CLEAR DATABASE AND START NEW
! SPECIFY JOBNAME AS 'SUB'
! SPECIFY JOBNAME AS 'SUB'
! ENTER PREPROCESSOR
! ENTER PREPROCESSOR
! USE PLANE42 ELEMENT TYPE
! USE PLANE42 ELEMENT TYPE
! SPECIFY ELASTIC MODULUS
! SPECIFY ELASTIC MODULUS
! SPECIFY POISSON'S RATIO
! SPECIFY POISSON'S RATIO
! CREATE RECTANGLE
! CREATE RECTANGLE
! CREATE CIRCLE

```
! CREATE CIRCLE
```



Fig. 11.13 Normal stress in the $y$-direction obtained from the global model and submodel, plotted against $x$.

```
ASBA,1,2
LESIZE,9,,,10
LESIZE,10,,,10
LESIZE,5,,,40
LESIZE,2,,,20
LESIZE,3,,,20
LCCAT,2,3
MSHKEY,1
AMESH,ALL
NSEL, S, LOC,Y,O
NSEL,A, LOC, X,O
NSEL,A,LOC,Y, 3
NSEL, A, LOC, X, 3
NWRITE,SBS,NOD
ALLSEL
SAVE
FINISH
RESUME,GLO,DB !RESUME FROM `GLO.DB' FILE
/ POST1
FILE,GLO,RST
CBDOF,SBS,NOD, ,SBS,CBD
! INTERPOLATION
FINISH !EXIT GENERAL POSTPROCESSOR
RESUME,SUB,DB !RESUME FROM `SUB.DB' FILE
/SOLU !ENTER SOLUTION PROCESSOR
/INPUT,SBS,CBD ! READ CUT BOUNDARY CONDITIONS
SOLVE !OBTAIN SOLUTION
FINISH ! EXIT SOLUTION PROCESSOR
/POST1 ! ENTER GENERAL POSTPROCESSOR
```

```
! OBTAIN STRESS CONTOURS IN Y-
! DIRECTION
```


### 11.4 Substructuring: Superelements

Consider the aircraft shown in Fig. 11.14. A detailed finite element model of an aircraft may contain several hundred thousand degrees of freedom. In order to reduce computational cost without losing accuracy, the matrix equations of the tail and wing sections can be condensed in terms of the corresponding interface nodes (Master DOFs) identified in Fig. 11.14. Thus, during the analysis, the size of the matrix system of equations becomes the number of unknown degrees of freedom of the fuselage section (main structure). Once the solution is obtained, all degrees of freedom belonging to the nodes of the fuselage are known, including the Master DOFs. Finally, the solution for the tail and wing sections is obtained by using the known Master DOFs as the boundary conditions.

The linear matrix equations representing a group of finite elements (and nodes) can be condensed into a smaller matrix system corresponding to a number of selected degrees of freedom (DOF). This procedure is commonly known as Static Condensation or Substructuring. The DOF selected in the analysis is referred to as the Master DOF. The resulting matrix may be considered as the element equations of a special element, which is called a Substructure or Superelement. Superelements are attached to the main structure through the Master DOF. The main advantage of substructuring is that it allows for the solution of large complex structures at a reduced computational cost (both memory and CPU time). Another advantage is that if the problem involves a combination of materials with linear and nonlinear material behavior, the linear portion can be condensed into a superelement so that only the matrices corresponding to the nonlinear portion are recalculated during the iterations of the nonlinear solution.

A substructuring analysis involves three main steps:
Generation Pass: Superelements are generated during this step. It involves selection of the master degrees of freedom, which are located along the interface between the superelement and the main structure. At the end of this step, matrices corresponding to each superelement are written to separate files in the Working Directory.

Use Pass: The main structure is generated in this step. Also, the superelements are introduced to the main structure. At the end of this step, the


Fig. 11.14 An aircraft structure and its finite element model divided into four parts (three superelements).
solution for the main structure is obtained, which includes the solution for the master degrees of freedom of each superelement.

Expansion Pass: During this step, complete solutions for the superelements are obtained by expanding the master degree of freedom solution available from the use pass.

Substructuring is demonstrated by considering an analysis of an electronic package with linear and nonlinear material behaviors.

The electronic package shown in Fig. 11.15 is composed of an FR4 substrate, two solder balls, and a silicon die. As indicated in the figure, thicknesses of the substrate and die are $h_{s}$ and $h_{d}$, respectively, and the stand-off height is $h_{b}$. The lengths of the die and substrate are $l_{1}$ and $l_{2}$, respectively. The pitch for the solder balls is $p$, and the balls are assumed to have circular geometry with radius $r$. Material properties of the die and the substrate are tabulated in Table 11.1. The solder behavior is viscoplastic using Anand's model, having properties given in Tables 10.2 and 10.3. The following numerical values of the parameters are used in the analysis: $h_{s}=2 \mathrm{~mm}$, $h_{b}=0.7 \mathrm{~mm}, h_{d}=1 \mathrm{~mm}, p=2 \mathrm{~mm}$, and $r=0.5 \mathrm{~mm}$. The temperature profile as a function of time is shown in Fig. 11.16, which is implemented in the ANSYS solution using four load steps. The stress-free temperature is taken as $0^{\circ} \mathrm{C}$. The temperature of the entire structure is first increased to $60^{\circ} \mathrm{C}$ over a period of 120 sec , followed by an increase to $100^{\circ} \mathrm{C}$ over 120 sec. Finally, the package is cooled down, following the same path along the temperature vs. time curve. The symmetry plane is constrained in the $x$ direction. Also, the $y$-direction displacement is suppressed at the bottom-left corner in order to prevent rigid-body translations. The goal is to solve this problem utilizing the substructuring technique. Figure 11.17 shows the superelement and main structure, and a close-up of the master degrees of freedom is shown in Fig. 11.18.

### 11.4.1 Generation Pass

A Generation Pass is composed of two main parts: (i) model generation and (ii) superelement generation. In this analysis, the solder balls are the main model and the substrate and the die constitute the superelement.

### 11.4.1.1 Model Generation

The solder material is modeled using two-dimensional VISCO106 viscoplastic elements. In order to specify temperature-dependent elastic properties, a temperature table is constructed using the MPTEMP command,


Fig. 11.15 Electronic package with FR4 substrate, solder balls, and silicon die.

Table 11.1 Properties of the constituent materials.

|  | Elastic <br> Modulus <br> $(\mathrm{GPa})$ | Poisson's <br> Ratio | CTE <br> $\left(\mathrm{ppm} /{ }^{\circ} \mathrm{C}\right)$ | Material <br> Reference <br> Number |
| :--- | :---: | :---: | :---: | :---: |
| Silicon die | 163 | 0.278 | 2.6 | 1 |
| FR4 substrate | 22.0 | 03.90 | 18.0 | 3 |



Fig. 11.16 Temperature load as a function of time.


Fig. 11.17 Superelement and main structure.


Fig. 11.18 Close-up of master degrees of freedom.
followed by the specification of properties using the MPDATA command. Note that temperatures are specified in Kelvin. Anand's viscoplastic properties are specified using the TB and TBDAtA commands. In order for the main model and superelement interface nodes to have the same numbers (for continuity), the full model is generated (shown in Fig. 11.17) and saved in a file named sUB1.DB. After this operation, the elements (and attached nodes) corresponding to the solder are deleted (EDEL and NDEL commands), leaving the mesh for only the superelement.

```
/FILNAME,GEN !SPECIFY JOBNAME AS GEN
/PREP7 ! ENTER PREPROCESSOR
ET,1,PLANE42 ! ELEMENT TYPE 1 IS PLANE42
ET,2,VISCO106 !ELEMENT TYPE 2 IS VISCO106
KEYOPT,2,5,2 ! EXTRA ELEM OUTPUT PLASTIC WORK
MPTEMP !DEFINE TEMPERATURE TABLE
MPTEMP,1,238.15,258.15,278.15 ! SPECIFY TEMPERATURES
MPTEMP,4,298.15,323.15,348.15,373.15,398.15 !
MP,EX,1,163E3 !ELASTIC MODULUS OF SILICON DIE
MP,NUXY,1,0.278 !POISSON'S RATIO OF SILICON DIE
MP,ALPX,1,2.6E-6 !CTE OF SILICON DIE
MPDATA, EX, 2,1,40781,37825,34884 ! ELASTIC MOD. OF SOLDER
MPDATA, EX, 2,4,31910,28149,24425,20710,16942 !
MPDATA,NUXY,2,1,0.354,0.3565,0.36 ! POISSON'S RAT. OF
    ! SOLDER
MPDATA, NUXY, 2,4,0.3628,0.365,0.37,0.3774,0.3839!
MPDATA,ALPX,2,1,2.43E-5,2.45E-5,2.47E-5 ! CTE OF SOLDER
MPDATA,ALPX,2,4,2.48E-5,2.50E-5,2.53E-5,2.55E-5,2.58E-5 !
MP, EX,3,22E3 ! ELASTIC MODULUS OF FR4 SUBSTRATE
MP,NUXY, 3,0.39 ! POISSON'S RATIO OF FR4 SUBSTRATE
MP,ALPX,3,18E-6 !CTE OF FR4 SUBSTRATE
TB,ANAND,2 ! ANAND'S VISCOPLASTIC MODEL FOR
    ! SOLDER
TBDATA, 1,12.41,9400,4E6,1.5,0.303,1379 ! SPECIFY ANAND'S
TBDATA,7,13.79,0.07,1.3 !MATERIAL PARAMETERS
HS=2 !SUBSTRATE THICKNESS
HB=0.7 !STAND-OFF HEIGHT
HD=1 !THICKNESS OF DIE
R=0.5 ! RADIUS OF SOLDER BALLS
P=2 ! PITCH
L1=5 !LENGTH OF DIE
L2=10 !LENGTH OF SUBSTRATE
R1=SQRT(R**2-(HB/2)**2)!HALF-LENGTH OF SOLDER AT
    ! TRUNCATION PT
CYL4,P/2,HS+HB/2,R ! CREATE 1ST SOLDER BALL
CYL4,3*P/2,HS+HB/2,R !CREATE 2ND SOLDER BALL
BLC4,0,0,L2,HS ! CREATE SUBSTRATE
BLC4,0,HS+HB,L1,HS ! CREATE DIE
ASEL,S,,1,2 !SELECT AREAS FOR SOLDER BALLS
CM,A1,AREA !CREATE COMPONENT OF AREAS
ASEL,S,,,3,4 !SELECT AREAS FOR DIE AND
    ! SUBSTRATE
CM,A2,AREA ! CREATE COMPONENT OF AREAS
ALLSEL !SELECT EVERYTHING
```

| ASBA, A1, A2, , KEEP | ! SUBTRACT AREAS (KEEP SUBTRACTED <br> ! AREAS) |
| :---: | :---: |
| AGLUE, ALL | ! GLUE ALL AREAS |
| ESIZE, 0.1 | ! SPECIFY ELEMENT SIZE |
| AMESH, ALL | ! MESH AREAS |
| ASEL, S, LOC, Y, HS / 2 | ! SELECT AREAS AT Y $=\mathrm{HS} / 2$ |
| ESLA, S | ! SELECT ELEMENTS ATTACHED TO AREAS |
| EMODIF, ALL, MAT, 3 | ! MODIFY MATERIAL ATTRIBUTE |
| ASEL, S, LOC, Y, HS + HB/2 | ! SELECT AREAS AT Y $=\mathrm{HS}+\mathrm{HB} / 2$ |
| ESLA, S | ! SELECT ELEMENTS ATTACHED TO AREAS |
| EMODIF, ALL , MAT, 2 | ! MODIFY MATERIAL ATTRIBUTE |
| EMODIF, ALL, TYPE, 2 | ! MODIFY ELEMENT TYPE ATTRIBUTE |
| ALLSEL | ! SELECT EVERYTHING |
| MODMSH, DETACH | ! DETACH SOLID MODEL FROM MESH |
| SAVE, SUB1, DB | ! SAVE AS SUB1. DB |
| ESEL, S, MAT , 2 | ! SELECT SOLDER ELEMENTS |
| EDEL, ALL | ! DELETE SELECTED ELEMENTS |
| NDEL, ALL | ! DELETE FREE NODES |
| ETDELE, 2 | ! DELETE VISCO106 ELEMENT TYPE |
| ALLSEL | ! SELECT EVERYTHING |
| FINISH | ! EXIT PREPROCESSOR |

### 11.4.1.2 Superelement Generation

The analysis type is specified as Substructuring. This makes sure that, when the SOLVE command is issued, the superelement file (GEN.SUB) is generated in the Working Directory. The superelement name is specified using the SEOPT command. The interface nodes between the main model and the superelement are defined as the master degrees of freedom using the $\mathbf{M}$ command (Fig. 11.18). After applying the displacement constraints (D command) and thermal load (BFUNIF command), the substructure is saved in the GEN. DB file, which is necessary for expanding the solution for the superelement after the solution for the main model is obtained. Figure 11.19 shows the superelement with the interface nodes indicated.

Although the superelement contains only materials with linear behavior, time values consistent with the applied thermal load profile (shown in Fig. 11.16) must be given. This is performed using the TIME command. The stress-free temperature is specified using the TREF command. Load vectors corresponding to each load step are obtained by using the SOLVE command repeatedly while changing the time value and thermal load.


Fig. 11.19 Superelement mesh with interface nodes indicated.


BFUNIF, TEMP, 373.15
TIME, 240

SOLVE
BFUNIF, TEMP, 333.15
TIME, 360

SOLVE

BFUNIF, TEMP, 273.15

TIME, 480
SOLVE

FINISH

```
! UNIFORM TEMPERATURE
! INCREASE TO 373.15 K
! TIME AT THE END OF LOAD
! STEP = 240 SEC
! OBTAIN LOAD VECTOR FOR
! SECOND LOAD STEP
! UNIFORM TEMPERATURE
! DECREASE TO 333.15 K
! TIME AT THE END OF LOAD
! STEP = 360 SEC
! OBTAIN LOAD VECTOR FOR
! THIRD LOAD STEP
! UNIFORM TEMPERATURE
! DECREASE TO 273.15 K
! TIME AT THE END OF LOAD
! STEP = 480 SEC
! OBTAIN LOAD VECTOR FOR
! FOURTH LOAD STEP
! EXIT SOLUTION PROCESSOR
```


### 11.4.2 Use Pass

At this point, the Working Directory includes the superelement file GEN. SUB. The main model is created conveniently by resuming from the SUB1.DB file and eliminating the elements (and attached nodes) corresponding to the substrate and the die. Element type 3 is defined as MATRIX50, which is used for superelements. The superelement file GEN. SUB is read by using the SE command, after which an element plot is obtained (EPLOT command) to verify the location of the superelement. In the solution phase of the Use Pass, thermal loading for the main model is applied, followed by reading the superelement load vector corresponding to the first load step using the SFE command with the SELV label. Finally, the solution for the main model is obtained. This procedure is repeated for load steps 2 through 4 . Figure 11.20 shows the main model with the interface nodes indicated.


Fig. 11.20 Main model mesh with interface nodes indicated.
/CLEAR
/FILNAME, USE
RESUME, SUB1,DB
/PREP7
ET, 3, MATRIX50

ESEL, U, MAT , , 2
EDEL, ALL
NDEL, ALI
ALLSEL
TYPE, 3
SE, GEN
EPLOT

ALLSEL
FINISH
/SOLU
NLGEOM, ON
KBC, 0

AUTOTS, ON
ESEL, S, TYPE, , 3
*GET, SENUM, ELEM, O , NUM, MAX
ALLSEL
TREF, 273.15
BFUNIF, TEMP, 333.15
SFE, SENUM, 1, SELV, , 1

TIME, 120
DELTIM, 10, 1, 10
SAVE
SOLVE
BFUNIF, TEMP, 373.15
TIME, 240
SFE, SENUM, 2, SELV, , 1
SOLVE
BFUNIF, TEMP, 333.15

TIME, 360
SFE, SENUM, 3, SELV, , 1
SOLVE
BFUNIF, TEMP, 273.15
! CLEAR DATABASE
! SPECIFY JOBNAME AS USE
! RESUME FROM SUB1.DB
! ENTER PREPROCESSOR
! MATRIX50 IS ELEMENT TYPE 3
! (FOR SUPERELEMENT)
! UNSELECT SOLDER ELEMENTS
! DELETE ALL SELECTED ELEMENTS
! DELETE ALL FREE NODES
! SELECT EVERYTHING
! SWITCH TO ELEMENT TYPE 3
! READ IN SUPERELEMENT
! VERIFY LOCATION OF
! SUPERELEMENT
! SELECT EVERYTHING
! ENTER PREPROCESSOR
! ENTER SOLUTION PROCESSOR
! TURN NONLINEAR GEOMETRY
! EFFECTS ON
! INTERPOLATE LOADS BETWEEN
! LOAD STEPS
! TURN AUTOMATIC TIME
! STEPPING ON
! SELECT SUPERELEMENT
! STORE ELEM \# INTO
! PARAMETER SENUM
! SELECT EVERYTHING
! STRESS-FREE TEMP IS 273.15 K
! UNIFORM TEMP. INCREASE TO
! 333.15 K
! READ IN SUPERELEMENT LOAD
! VECTOR 1
! TIME AT THE END OF
$!\mathrm{LS}=120 \mathrm{SEC}$
! TIME STEP SIZE SETTINGS
! SAVE DATABASE TO USE.DB
! OBTAIN SOLUTION FOR
! LOAD STEP 1
! UNIFORM TEMP. INCREASE TO ! 373.15 K
! TIME AT THE END OF
! LS $=240 \mathrm{SEC}$
! READ IN SUPERELEMENT LOAD
! VECTOR 2
! OBTAIN SOLUTION FOR L LOAD STEP 2
! UNIFORM TEMP. DECREASE TO ! 333.15 K
! TIME AT THE END OF $!\mathrm{LS}=360 \mathrm{SEC}$
! READ IN SUPERELEMENT LOAD ! VECTOR 3
: OBTAIN SOLUTION FOR ! LOAD STEP 3
! UNIFORM TEMP. DECREASE TO ! 273.15 K
! TIME AT THE END OF

|  | !LS $=480$ SEC |
| :--- | :--- |
| SFE, SENUM, 4, SELV, , 1 | ! READ IN SUPERELEMENT LOAD |
|  | !VECTOR 4 |
| SOLVE | !OBTAIN SOLUTION FOR |
|  | !LOAD STEP 4 |
| FINISH | !EXIT SOLUTION PROCESSOR |
| /POST1 | !ENTER GENERAL POSTPROCESSOR |
| PLNSOL,NL, PLWK | !PLOT CONTOURS OF PLASTIC WORK |
| FINISH | !EXIT GENERAL POSTPROCESSOR |

The solution contains all results items for the main model, including the degree of freedom results for the nodes along the interface between the main model and the superelement. The superelement degree-of-freedom solution is also stored in the file USE.DSUB, which is used in the Expansion Pass. Figure 11.21 shows the contour plots of plastic work at the end of the fourth load step.

### 11.4.3 Expansion Pass

In order to obtain a complete a solution for the superelement, the database is first cleared, followed by resuming (RESUME command) from the GEN. DB file. Expansion Pass is turned on using the ExpASS command. Names of the superelement file (GEN.SUB) and the file containing the superelement solution (USE.DSUB) are declared using the SEEXP command. The command EXPSOL is used for obtaining a solution at different time values (load steps).

| /CLEAR | ! CLEAR DATABASE |
| :---: | :---: |
| /FILNAME, GEN | ! SPECIFY JOBNAME AS GEN |
| RESUME | ! RESUME FROM GEN. DB |
| / SOLU | ! ENTER SOLUTION PROCESSOR |
| EXPASS, ON | ! ACTIVATE EXPANSION PASS |
| SEEXP, GEN, USE | ! SUPPLY SUPERELEMENT NAME TO BE <br> ! EXPANDED |
| EXPSOL, , 120 | $!$ TIME OF INTEREST IS 120 SEC |
| SOLVE | ! OBTAIN EXPANSION PASS SOLUTION |
| EXPSOL, , , 240 | ! TIME OF INTEREST IS 240 SEC |
| SOLVE | ! OBTAIN EXPANSION PASS SOLUTION |
| EXPSOL, , 360 | ! TIME OF INTEREST IS 360 SEC |
| SOLVE | ! OBTAIN EXPANSION PASS SOLUTION |
| EXPSOL, , , 480 | ! TIME OF INTEREST IS 480 SEC |
| SOLVE | ! OBTAIN EXPANSION PASS SOLUTION |
| FINISH | ! EXIT SOLUTION PROCESSOR |
| /POSTI | ! ENTER GENERAL POSTPROCESSOR |
| SET, LAST | ! SET RESULTS TO LAST LOAD STEP |
| PLNSOL, S, Y, 2 | ! PLOT CONTOURS OF STRESS IN Y! DIRECTION |
| PLNSOL, S, XY, 2 | ! PLOT CONTOURS OF STRESS IN XY! DIRECTION |
| FINISH | ! EXIT GENERAL POSTPROCESSOR |



Fig. 11.21 Contour plot of plastic work at the end of the fourth load step.

Figures 11.22 and 11.23 show the contour plots of the normal and shear stresses, $\sigma_{y y}$ and $\sigma_{x y}$, respectively, in the superelement at the end of fourth load step.

### 11.5 Interacting with External Files

Interaction with external files makes ANSYS a very versatile software. The most common such interactions include:

Reading an input file.
Writing data to ASCII (plain text) files.
Reading data from ASCII (plain text) files.
Passing a command string to the operating system.
These topics are briefly discussed in this section. However, the descriptions of commands are not complete; only their most common usage is described. Therefore, it is recommended that the user consult the ANSYS Help System for complete instructions.

### 11.5.1 Reading an Input File

As described in Chap. 7, input files are ASCII files containing ANSYS commands (one command per line) to be read and executed sequentially. In Chap. 7 through 10, several example problems were demonstrated utilizing command line input. The input files are read from within ANSYS by issuing the / INPUT command from the Input Field. The commonly used syntax for this command is as follows:

```
/INPUT, Fname, Ext
```

in which Fname and Ext are the path with the file name and extension of the input file, respectively. If the path is not specified, i.e., the argument Fname includes only the file name, then the file is searched for in the Working Directory.


Fig. 11.22 Contour plot of normal stress, $\sigma_{y y}$, in the superelement at the end of the fourth load step.


Fig. 11.23 Contour plot of shear stress, $\sigma_{x y}$, in the superelement at the end of the fourth load step.

### 11.5.2 Writing Data to External ASCII Files

There are two main motivations in writing data to files:
ANSYS software may be used as a pre- and postprocessor while the actual computations are performed by an external executable file. In such cases, the executable file almost always requires input data. Such data can be written to ASCII files, with a specific format, to be read by the executable file as input.

Although ANSYS provides powerful contour and line plot options, it may not always offer the most convenient environment for specific plotting formats. In such cases, it may be more efficient to write the
required data to an external ASCII file; this file can then be processed with specialized graphics or spreadsheet software to produce the desired plots.

There are two main commands that enable writing data to external files: /OUTPUT and *CFOPEN. Once either of these commands is issued, data are written to an external file using the *VWRITE command. These commands are discussed in the following subsections.

### 11.5.2.1 The Ioutput Command

The /OUTPUT command redirects text output to a specified file. The syntax for the IOUTPUT command is as follows:
/ OUTPUT, Fname, Ext, , Loc
in which Fname and Ext are the path (file name and extension of the file, respectively) and Loc determines whether to write starting from the top of the file (effectively deleting the contents of the file) or appending at the bottom of the file. Note that the third argument is not used. If the Loc field is left blank, then ANSYS writes data starting from the top of the file. In order to append an existing file, the phrase APPEND is entered in the Loc field. After issuing the /OUTPUT command with its arguments, text that normally appears in the Output Window is written to the external file. Therefore, immediately after writing the data to the file, the text output needs to be redirected to the Output Window by issuing the /OUTPUT command without any arguments. An example is shown below that writes the parameters in an external file named DATA.OUT in the Working Directory:

```
/OUTPUT, DATA,OUT, ,APPEND
*VWRITE,CH1,NUM1,NUM2
(A8, 2X,F12.8,2X,E13.6)
/OUTPUT
```


### 11.5.2.2 The *CFOPEN and *CFCLOS Commands

A better way of writing data to an external file is to open a "command" file using the *CFOPEN command. It has a syntax similar to the /OUTPUT command, with the same argument definitions:

```
*CFOPEN, Fname, Ext, ,Loc
```

After issuing this command with a file name and extension, anytime the *VWRITE command is issued, the data are written to the file. Once finished with the data transfer, the file can be closed using the *CFCLOS command, which has no arguments.

### 11.5.2.3 The *VWRITE Command

The syntax for the *VWRITE command is as follows:
*VWRITE, Par1, Par2, . . . Par19
in which Par1, ..., Par19 are user-defined parameters. The *VWRITE command line must be followed by a format statement enclosed in parenthesis. The format statement follows FORTRAN programming language syntax. There are five distinct format descriptors addressing real numbers (three different descriptors), characters, and blank spaces. Real number format descriptors are $\mathbf{F}, \mathbf{E}$, and $\mathbf{D}$, which have the following syntax

FW.d
Ew.d
Dw. d
in which $w$ is the total number of digits allocated for the number in the file (total width) while $d$ denotes the number of decimal places (number of digits to the right of the decimal point). The information given herein is common to FORTRAN compilers used in MS Windows operating systems. For compilers used under different operating systems, small changes may be expected, but the general concept remains the same. The descriptor $\mathbf{F}$ writes the real number with no exponent while descriptors $\mathbf{E}$ and $\mathbf{D}$ write with E and $D$ exponents, respectively. The number $w$ must account for the sign of the number, decimal point and the exponent field. The exponent field, when using descriptors $\mathbf{E}$ and $\mathbf{D}$, takes up four digits: one digit for the exponent character $\mathbf{E}$ or $\mathbf{D}$, one digit for the sign of the exponent, and two digits for the exponent itself. Therefore, in format statements using $\mathbf{E}$ and $\mathbf{D}$ descriptors, the difference $(w-d)$ must be at least 7 .

In order to specify the format for characters, the descriptor $\mathbf{A}$ is used with the following syntax:

AW
where $w$ is the total width. The maximum width allowed by ANSYS is 8 digits.

Finally, blank spaces are inserted using the $\mathbf{x}$ descriptor with the following syntax:

## $w x$

in which $w$ is the total number of blank spaces.
The use of descriptors $\mathbf{F}, \mathbf{E}, \mathbf{D}, \mathbf{A}$, and $\mathbf{X}$ is demonstrated next.

Consider the numbers 152.67328199 and $-3.251667 \times 10^{6}$, which are stored in parameters NUM1 and NUM2, respectively, and the character string result 1 stored in parameter CH1 in ANSYS. They can be written to an external ASCII file using the following commands lines:

```
*VWRITE, CH1,NUM1,NUM2
(A8, 2X,F12.8, 2X, E13.6)
```

which produces the following output in the ASCII file


There are two observations to be made: (i) when the character string is given more digits than its length, ANSYS inserts as many blank spaces as needed to make the field the same width as that specified and justifies the string to the left, and (ii) when using E or $\mathbf{D}$ descriptors, ANSYS places a $\mathbf{0}$ immediately to the left of the decimal.

When the format statement above is replaced with

$$
(\mathrm{A} 2,1 \mathrm{X}, \mathrm{~F} 11.8,1 \mathrm{X}, \mathrm{E} 13.2)
$$

The following output is written in the ASCII file:


Observations:
When the number of digits allocated by the descriptor $\mathbf{A}$ is less than the actual length of the character string, ANSYS truncates the character string.

When the number of digits allocated by the descriptor $\mathbf{F}$ (or $\mathbf{E}$ or $\mathbf{D}$ ) is less than the actual length of the real number, ANSYS prints the character * as many times as the specified number digits.

When the format statement is replaced with

$$
(\mathrm{A} 7,1 \mathrm{X}, \mathrm{E} 11.4,1 \mathrm{X}, \mathrm{D} 14.7)
$$

the following output is written in the ASCII file:


### 11.5.3 Executing an External File

In certain cases, executing an external file may be required. This could be software performing an independent analysis. The command /SYS, which passes a command string to the operating system, is used in such cases. Its syntax is written as follows:

```
/SYS,String
```

in which the argument String is the command string. When used for the aforementioned purpose, the argument String is the name of the executable file. For example, if the name of the executable file is ROOTS. Exe, then the following command is issued to execute this file:
/SYS,ROOTS.EXE
Upon execution of this command, the roors.exe file runs, and ANSYS does not allow any action to be taken during this time.

As an example, we consider the generation of a time-dependent random thermal load. In this example, an external executable file, RANDOM.EXE, generating random numbers is created using the FORTRAN programming language. The code requires values for five input parameters:

| TOT | Total time the load is applied. |
| :--- | :--- |
| ND | Number of data points (random numbers). |
| MIN | Minimum value of load. |
| MAX | Maximum value of load. |
| SEED | Seeding number for random number generation. |

The values of these parameters are provided through an input file named RANDOM.DAT. After the code generates numbers within the range [MIN, max], they are written to an output file named random. out. The input is unformatted while the output is formatted. The FORTRAN code is given as follows:

| REAL (8) NUM, TOT, DT, T, MIN, MAX | ! REAL VARIABLES |
| :---: | :---: |
| INTEGER(4) ND, F, I, SEED | ! INTEGER VARIABLES |
| OPEN (1, FILE= RANDOM. DAT') | ! DECLARE INPUT FILE |
| OPEN ( $2, ~ \mathrm{FILE}=$ 'RANDOM. OUT ' ) | ! DECLARE OUTPUT FILE |
| READ (1,*) TOT, ND, MIN, MAX, SEED | ! READ INPUT |
| DT=TOT/ND | ! FIND TIME STEP SIZE |
| $\mathrm{T}=\mathrm{DT}$ | ! INITIALIZE TIME |
| $\mathrm{F}=0$ | ! INITIALIZE RANDOM |
|  | ! NUMBER |
|  | ! GENERATOR FLAG |
| $A M P=M A X-M I N$ | ! FIND LOAD RANGE |
| NUM=DRAND (SEED) | ! GENERATE RANDOM |
|  | ! NUMBER |
|  | ! SET |

```
    DO I=1,ND ! START LOOP OVER DATA
    NUM=MIN+AMP*DRAND (F) ! FIND LOAD VALUE
    WRITE(2,100) T,NUM
    T=T+DT
ENDDO
100 FORMAT (2(E16.8,2X))
END
```

```
POINTS
```

POINTS
WRITE TIME AND LOAD
WRITE TIME AND LOAD
VALUES
VALUES
UPDATE TIME
UPDATE TIME
END LOOP OVER DATA
END LOOP OVER DATA
POINTS
POINTS
FORMAT STATEMENT
FORMAT STATEMENT
END OF PROGRAM

```
END OF PROGRAM
```

In order to create the executable file, one must compile and link the code above using a FORTRAN compiler.

The following command input segment, executed from within ANSYS, writes the required input to file RANDOM.DAT, executes the file random.exe (resides in the Working Directory), reads the time vs. load values from file RANDOM.OUT, and, finally, plots the variation of load vs. time. After the file RANDOM.EXE is executed, three array parameters are defined (*SET command) and given dimensions (*DIM command). The time and load values are stored in one large array using the *VREAD command. Finally, axis labels are specified using the /AXLAB command and the plot is obtained using the *VPLOT command.

| /PREP7 | ! ENTER PREPROCESSOR |
| :---: | :---: |
| TOT=8 | ! DEFINE PARAMETERS |
| $\mathrm{ND}=160$ | $!$ |
| $\mathrm{MN}=0$ | ! |
| $\mathrm{MX}=100$ | ! |
| SEED=12 | ! |
| / OUTPUT, RANDOM, DAT | ! REDIRECT OUTPUT TO RANDOM.DAT |
| *VWRITE, TOT, ND, MN, MX, SEED | ! WRITE PARAMETERS TO FILE |
| (E16.8,2X,F8.0,2X, E10.2, 2X, E10.2,2X, F8.0) ! FORMAT |  |
|  | $!$ STATEMENT |
| /OUTPUT | ! REDIRECT OUTPUT TO OUTPUT ! WIN. |
| /SYS, RANDOM.EXE | ! EXECUTE RANDOM.EXE |
| *SET, ARY | ! INITIALIZE PARAMETER ARY |
| *SET, TIM | ! INITIALIZE PARAMETER TIM |
| *SET, LOAD | ! INITIALIZE PARAMETER LOAD |
| *DIM, ARY, ARRAY, 2 *ND | ! DIMENSION PARAMETER ARY |
| *DIM, TIM, TABLE, ND | ! DIMENSION PARAMETER TIM |
| *DIM, LOAD, TABLE, ND | ! DIMENSION PARAMETER LOAD |
| *VREAD, ARY (1) , RANDOM, OUT | ! FILL ARY ARRAY FROM |
|  | : RANDOM. OUT |
| (2 (E16.8,2X) ) | ! FORMAT STATEMENT |
| *DO, I, 1, ND | ! LOOP OVER DATA POINTS |
| $T I M(I)=A R Y(2 * I-1)$ | ! SEPARATE ARRAY ARY INTO TIM |
| $\operatorname{LOAD}(\mathrm{I})=\operatorname{ARY}(2 * I)$ | ! AND LOAD ARRAYS |
| * ENDDO | ! END LOOP OVER DATA POINTS |
| /AXLAB, X, TIME | ! DEFINE X-AXIS LABEL |
| /AXLAB, Y, LOAD | ! DEFINE Y-AXIS LABEL |
| *VPLOT, TIM (1), LOAD (1) | ! PLOT TWO ARRAYS AGAINST EACH |
|  | ! OTHER |

When the command input segment above, with the given values of parameters, is read as an input file from within ANSYS, the plot shown in Fig. 11.24 is produced in the Graphics Window.

### 11.5.4 Modifying ANSYS Results

ANSYS software can be used as a postprocessor for results calculated by an external program utilizing the DNSOL or DESOL commands, which impose those results on the ANSYS mesh. The syntax for DNSOL and DESOL commands are:

```
DNSOL, NODE, Item, Comp,V1,V2,V3,V4,V5,V6
DESOL, ELEM, NODE, Item, Comp,V1,V2,V3,V4,V5,V6
```

in which NODE and ELEM are the node and elements numbers, respectively, at which the results are specified, Item and Comp are labels for the specific degree of freedom, and V1 through V6 are the values of the results. If the $y$ displacement (Item is $U$ and Comp is $\mathbf{Y}$ ) at node 12 is modified to be 2.5 , the following command input is used:

```
DNSOL,12,U,Y,2.5
```

Similarly, when shear stress in the $y z$-direction (Item is $\mathbf{S}$ and Comp is YZ) at node 212 of element 23 is modified to be 1200 , the following command input is used:

```
DESOL,23,212, S,YZ,1200
```


### 11.6 Modifying the ANSYS GUI

ANSYS allows the user to modify/customize its Graphical User Interface (GUI). This capability is very advantageous, particularly when the user performs similar types of analyses over and over again. However, it is not a straightforward task. The main resource for learning how to customize the GUI is the ANSYS UIDL Programmer's Guide, in which UIDL stands for User Interface Design Language. This guide is not included in the ANSYS Help System; it must be purchased separately.

This section does not provide exhaustive information on UIDL. It is written with the intention of giving the reader a few examples on how the UIDL is modified. First, a brief summary of steps involved in modifying the UIDL are discussed, after which two simple examples are given for demonstration purposes. The first example involves customized dialog boxes, and the second involves a customized pick menu.


Fig. 11.24 Variation of load with time as read from the RANDOM. OUT file.

The following steps are taken to customize the ANSYS GUI:

- Locate the Control Files UIFUNC1.GRN, UIFUNC2.GRN, and UIMENU.GRN, and the file menulist**.ans. The characters ** correspond to the ANSYS version being used. For example if ANSYS Release 8.0 is being used, then the file name is menulist80.ans. In later ANSYS releases, these files are located in the same directory. In a PC using a Windows operating system, a possible location is:

```
\Program Files\Ansys Inc\v80\ANSYS\gui\en-us\UIDL
```

- Create a temporary directory, e.g.:
\user\temp
- Copy the control files and the file menulist 80 . ans into the temporary directory.
- Using parts of the existing control files located in the temporary directory, create new control files (this point is explained in detail later).
- Copy the new control files and the menulist80. ans file to a Working Directory and add the names of the new control files, including their path, in the menulist80, ans file.
- Start ANSYS with the directory containing the new control files and the menulist80.ans file as the Working Directory and verify that the modified GUI works properly.
- When finished, copy the menulist80.ans file to additional Working Directories; the modified GUI will be available when ANSYS is started using those directories as Working Directory.

There are three control files: UIFUNC1.GRN, UIFUNC2.GRN, and UIMENU. GRN. The first two contain function blocks and the last one is for the menu blocks. The general logic of menu and function blocks is explained considering existing blocks. First, menu blocks are discussed, followed by a discussion on function blocks.

At the beginning of the UIMENU.GRN file, the following Control File Header is placed:

```
:F UIMENU.GRN
:D Modified on %E%, Revision (SID) = %I%
:I 403, 100064, 110164
:!
```

in which the : $\boldsymbol{F}$ command defines the name of the control file, : $\boldsymbol{D}$ gives a description of the control file, and the :I command stores internally generated index numbers. At the beginning of every control file, these three commands must appear in the sequence given above. In a new control file, the :I command must have comma-separated zeros in columns 9,18 and 27. Once ANSYS runs and reads the file, it assigns new numbers and modifies the control files. The Control File Header must be separated from the menu blocks by a line containing : ! (the fourth line above).

The Main Menu in ANSYS is created by the following menu block, taken from UIMENU. GRN file:

```
:N MenuRoot
:S 488, 76, 405
:T Menu
:A Main Menu
:D ANSYS ROOT MENU
Men_UVBA_Main_T1
Men_UVBA_Main_T2
Men_UVBA_Main_T3
Fnc_Preferences
Sep_
Men_Preproc
Men_Solution
Men_GenlPost
Men_TimePost
K_LN(DROPTEST)
Sep_
K_LN(DROPTEST)
Men_DropTest
Sep
Men_Topo
```

```
Men_ROM
Men_DesXplorerVT
K_LN(alpha)
Men_DesOpt
K_LN(ALPHA)
Men_DesOpt_al
Men_ProbDesign
Men_Aux12
Men_RunStat
Sep_
Fnc_UNDO
Sep_
Fnc_FINISH
K_LN(UTILMENU)
Men_UtilMenu
Men_UVBA_Main_B1
Men_UVBA_Main_B2
Men_UVBA_Main_B3
:E END
:!
```

which is composed of following three distinct sections:
Block Header Section: Commands belonging to this section always start with a colon (:). The first command must be $: \mathbf{N}$, which defines a unique internal name given to the block. The next command must be the : $\mathbf{S}$ command, which stores internally generated index numbers. In a new control file, the : $\mathbf{S}$ command must have comma-separated zeros in columns 9, 16, and 23. Once ANSYS runs and reads the file, it assigns new numbers and modifies the control files. The command : $T$ identifies the block as a menu block (Menu) or a command block (Cmd). The command $: T$ defines the heading of the block as it appears in the menu system. There are optional commands that can be used in the Block Header Section, such as :D, which is the description of the block.

Data Controls Section: This section contains four distinct data control commands: Men_, Fnc_, Sep_, and K_LN. The control commands Men_ and Fnc_ are calls to existing menu and function blocks, respectively. For example, the line Men_Preproc is a call to the menu block for the ANSYS Preprocessor in the same control file (UIMENU.GRN), which has the internal name Men_Preproc defined by the $: \mathrm{N}$ command. The function blocks are stored in one of two files: UIFUNC1.GRN or UIFUNC2.GRN. Thus, the line Fnc_Preferences calls the function block that is in UIFUNC2 . GRN file. Figure 11.25 shows the ANSYS Release 8.0 Main Menu, with Preferences (Fnc_Preferences) and Preprocessor (Men Preproc). The command Sep_ is used for inserting a separator bar between menu items. However, in newer releases of ANSYS, the whole

```
ANSYS Main Menu
圆 Preferences
⿴囗 Preprocessor
田 Solution
田 General Postproc
\Psi TimeHist Postpro
⿴囗十
⿴囗⿱一一⿻儿口一隹 Tool
田 Design Opt
# Prob Design
田 Radiation Opt
⿴囗 Run-Time Stats
眳 Session Editor
Finish
```

Fig．11．25 ANSYS Main Menu with functions and submenus．
menu is in one window with a tree－like structure（as opposed to separate windows for menus）．Therefore，the separators in menus are no longer visible．The command K＿LN is related to ANSYS keyword logic，which decides to show or not show certain menu items；this topic is not discussed within the context of this book．

Ending Section：Every menu block must end with these two lines：

```
:E END
:!
```

Function blocks are contained in the control files UIFUNC1．GRN and UIFUNC2．GRN．They execute ANSYS commands．As an example for function blocks，the creation of rectangles by dimensions function block，as given below，is considered（dialog box shown in Fig．11．26）．This block executes the recting command．The function block is taken from the UIFUNC2．GRN file．


Fig．11．26 Dialog box produced by the Fnc＿Recting function block．

```
:N Fnc_RECTNG
:S 355, 145, 203
:T Command
:C )! Fnc_RECTNG
:A By Dimensions
:D Create Rectangle by Dimensions
:K #(PREP7)
:H Hlp_C__RECTNG
Cmd_RECTNG
    Fld_0
        Typ_Lab
        Prm_[RECTNG] Create Rectangle by Dimensions
    Fld_2
        Prm_X1,X2 X-coordinates
        TYp_REAL2
        Def_Blank,Blank
    Fld_4
        Prm_Y1,Y2 Y-coordinates
        TYp_REAL2
        Def_Blank,Blank
    :E END
:!
```

Function blocks are also composed of three distinct sections:
Block Header Section: Definitions of commands belonging to this section are the same as those for the menu blocks. In addition to the commands described under menu blocks, there is another important command: : $\mathbf{C}$. This command is used for executing ANSYS commands before the actual function is executed. An exclamation mark designates, as in the APDL, comments.

Data Controls Section: Many more commands can be used within this section under function blocks than can be used under menu blocks. For a complete list of available commands, the reader should consult the ANSYS UIDL Programmer's Guide. A partial list is given in Table 11.2.

In the example above, the Data Controls Section starts with the line containing Cmd_recting. The command Cmd_ initiates a command sequence, in this case the RECTNG command. The Fld command defines the field number, which is associated with the command being used. If no data are collected in a field (such as a text block in the dialog box), then 0 is assigned to the Fld_ command, i.e., F1d_0. Each field has a type defined by the TYp command. In this case, the text block [RECTNG] Create Rectangle by Dimensions, in the dialog box (Fig. 11.26) is created using the Typ_Lab (stands for type label) command. Finally the label is printed using the Prm_ command, which is immediately followed by the text block.

Table 11.2 Function block data control commands.

| General Data Control Commands: |  |
| :---: | :---: |
| Cal | Calls another function block. If used, must be the last item before the : E END command is issued. |
| Inp_P | Defines pick menus. |
| Inp_NoApply | When used, Apply button is not shown. |
| Typ_Lab | Defines a label field. |
| Command Control Commands: |  |
| Cmd | Initiates a command sequence. |
| General Field Control Commands: |  |
| Def | Sets default values for the data to be shown in dialog boxes. |
| Fld | Assigns numbers to fields. |
| Prm | Prints text in dialog boxes. |
| Picking Control Commands: |  |
| Max | Sets maximum number of items to be picked. |
| Min | Sets minimum number of items to be picked. |
| Sel | Defines type of selection allowed. |
| Typ_Entity | Retrieves the numbers of the specified entities (NODE, ELEM, KEYP, LINE, AREA, VOLU). |
| TYP_XYZ | Retrieves the coordinates of a point. |
| Listing Control Commands: |  |
| Typ_Idx | Creates scrolling lists from subsequent $I d x_{-}$ commands. |
| Typ Lis_Options | Creates option button lists. |
| TYp_Lis_Radiob | Creates radio-button lists. |
| TYp_Lis | Creates single scrolling lists. |
| TYp_MLis | Creates multiple scrolling lists. |
| Numerical Control Commands: |  |
| TYp_Int | Creates single input field for integers. |
| TYp_Int2 | Creates two input fields for integers. |
| TYp Int3 | Creates three input fields for integers. |
| TYp_Real | Creates single input field for real numbers. |
| TYP_Real2 | Creates two input fields for real numbers. |
| TYP_Real3 | Creates three input fields for real numbers. |
| Character Control Commands: |  |
| Typ_Char | Creates single input field for characters. |
| Logical Control Commands: |  |
| Typ_Logi | Creates toggle buttons. |

If data are collected in a field, then non-zero, sequential field numbers must be assigned. The field number for the first data item is 2, i.e., F1d_2. In this particular example, X1 and X2 coordinates are collected, which constitute field numbers 2 and 3 . Hence, the next two data items, $\mathbf{Y} 1$ and $\mathbf{Y} 2$, are assigned field numbers 4 and 5 . All of the data collected in this case are real numbers. Also, two data items per row are collected, which is achieved using the command Typ_Rea12. Single and three real numbers per row are collected using Typ_Real and Typ_Real3 commands. Similarly, integer data can be collected using commands Typ_Int, TYp_Int2, and Typ_Int3. Default values for the data can be supplied by using the Def_ command. In this case, blank data fields are enforced, i.e., Def_Blank is used. If, for example, the default values of X1 and X2 are 0.1 and 0.4 , respectively, then Def_0.1,0.4 would be used.

Ending Section: Every function block must end with the following two lines:

```
:E END
:!
```

ANSYS dialog boxes offer numerous modes of interaction with the user. A partial list of these interactions is:

- Single Selection List: Only one selection can be made.
- Text Option Button List: Creates a pull-down menu.
- Radio-button List: Creates several radio-buttons (only one can be selected).
- Multiple Selection List: Multiple items can be selected.
- Side-by-side Parent Child List: Creates two lists, one on the left (parent) and the other on the right (child). Every item in the parent list has at least one child in the child list.
- Data Field: Creates data fields for real and integer numbers, characters, and yes/no types of interactions.

Each of these interactions passes values corresponding to the selection made to ANSYS commands (or user macro files). In the case of Data Fields, the value that is passed is specified in the text boxes.

GUI modification using UIDL is demonstrated by considering three examples. The first example demonstrates how to create function blocks utilizing the different modes of interaction listed above. In the second example, data are collected through a dialog box used as input for an external code (RANDOM.EXE of Sec. 11.5.3), which is subsequently executed. A menu item that reads and plots the output generated by the external executable file
is also generated. Finally, a simple example utilizing a customized picking menu is included.

### 11.6.1 GUI Development Demonstration

In this example, a menu item under the ANSYS Preprocessor menu is generated. Its name appears as DEMO. There are three function blocks under the DEMO menu: F1, F2, and F3. The function block F1 involves a Single Selection List, a Text Option Button List and a Radio-button List. In function block F2, creation of a Multiple Selection List and a Side-by-side Parent Child List is demonstrated. Finally, function block F3 utilizes various Data Fields. The following is a complete step-by-step list of actions taken to achieve this task.

## STEP 1

Copy the original control files UIFUNC1.GRN, UIFUNC2.GRN, and UIMENU.GRN, and the file menulist80.ans file to the temporary directory \user $\backslash$ temp.

## STEP 2

Create two temporary files with names zero1.dat and zero2. dat in the temporary directory. Open both files using a text editor. In the zero1. dat file, enter the following phrase:

```
:I 0, 0, 0
```

making sure the zeros are placed in columns 9,18 , and 27 . In the zero2 dat file, enter the following phrase:

```
:S 0, 0, 0
```

This time, the zeros must be placed in columns 9,16 , and 23 . Save both files and leave them open.

## STEP 3

Create a new control file in the temporary directory with a user-defined name, say USERM.GRN. Open both the USERM.GRN and UIMENU.GRN files using a text editor. Copy the first four lines from the UIMENU. GRN file and paste them in the file USERM.GRN. Modify the first two lines in the USERM. GRN file to be:

```
:F USERM.GRN
:D User Defined Menu
```

Copy the single line in the zerol. dat file and paste it to replace the third line in the USERM. GRN file. At this point, the USERM. GRN file looks like:

```
:F USERM.GRN
:D User Defined Menu
:I 0, 0, 0
:!
```


## STEP 4

Because the new menu item is going to be under the ANSYS Preprocessor, find the menu block corresponding to the ANSYS Preprocessor. It was observed earlier that the internal name for the Preprocessor is Preproc. Perform a word search of the UIMENU.GRN file for the phrase Men_ Preproc, which produces several results. Find the location with the phrase:

```
:N Men_Preproc
```

Copy the menu block Preproc from file UIMENU.GRN and paste it in the file USERM.GRN, after the first four lines modified earlier. In the USERM.GRN file, find the line starting with the : $\mathbf{S}$ command (sixth line). Copy the single line in the zero2. dat file and paste it to replace the sixth line in the USERM.GRN file.

The DEMO menu is going to be the first item available under the ANSYS Preprocessor. So, in the USERM.GRN file, first find the line with the phrase

```
Men_ElemType
```

then insert the following line above it:

```
Men_DEMO
```

which declares that there is a menu block with an internal name DEMO.
Create the menu block for DEMO at the bottom of the USERM.GRN file as follows:

```
:N Men_DEmo
:S 0, 0, 0
:T Menu
:A Demo Menu
:D Demo Menu
Fnc_F1
Fnc_F2
Fnc_F3
:E END
:!
```

Make sure the menu block for the Preprocessor is separated from the Demo menu block by the following line:

Creation of the customized menu is complete. Save the USERM.GRN file.

## STEP 5

Three function blocks are created next. For this purpose, create a new control file in the temporary directory with a user-defined name, say USERF. GRN. Open the newly created USERF. GRN file using a text editor. Copy the first four lines from the USERM. GRN file and paste them in the file USERF. GRN. Modify the first two lines in the USERF. GRN file to be

```
: F USERF.GRN
```

:D User Defined Functions
The first function block utilizes a Single Selection List, a Text Option Button List, and a Radio-button List. Enter the following text in the USERF.GRN file, starting from line 5:

```
:N Fnc_F1
:S 0, 0, 0
:T Command
:A Function F1
:D Function Fl Heading
Cmd_F1
Fld_0
        Typ_Lab
        Prm_Type the description of function F1 here
Fld_0
        Typ_Sep
Fld_0
        Typ_Lab
        Prm_Typ_Lis command is used to generate the list
Fld_2
        Typ_Lis
            Lis_Item 1,1
            Lis_Item 2,2
            Lis_Item 3,3
            Lis_Item 4,4
            Lis_Item 5,5
        Prm_Selected item appears in the box
Fld_0
    Typ_Sep
Fld_0
    Typ_Lab
    Prm_Typ_Lis_OptionB command is for pull-down menus
Fld_3
        Prm_Choose one option
        Typ_Lis_OptionB
        Lis_Option 1,10
        Lis_Option 2,20
        Lis_Option 3,35
        Lis_Option 4,89
Fld_0
    Typ_Sep
Fld_0
        Typ_Lab
        Prm_Typ_Lis_RadioB command is for radio-buttons
```

```
Fld_4
    Prm_Choose one of the buttons
    Typ_Lis_RadioB
        Lis_Button 1,4
        Lis_Button 2,7
        Lis_Button 3,23
:E END
:!
```

The function block above produces the dialog box shown in Fig. 11.27. There are three data fields (fields 2,3 , and 4 ). After making the selections and clicking OK , ANSYS issues the command F1 with three arguments (which is not a standard ANSYS command). For example, if the user selects Item 4 from the single selection list, Option 3 from the pull-down menu, and Button 2 from the radio-buttons, then ANSYS issues the following command input:

```
F1,4,35,7
```



Fig. 11.27 Dialog box generated by function block F1.

The arguments for each list item are specified in the UIDL segment for $\boldsymbol{F 1}$ given above. For example, selecting Button 3 passes 23 as the third argument of the command $F 1$, enforced by the line

Lis_Button 3,23
The second function block demonstrates Multiple Selection and Side-by-side Parent Child Lists. Append the following text in the USERF. GRN file, after the function block for $\mathbf{F 1}$ :

```
:N Fnc__F2
:S 0, 0, 0
:T Command
:A Function F2
:D Function F2 Heading
Cmd_F2
    Fld_0
        Typ_Lab
        Prm_Type the description of function F2 here
Fld_0
        Typ_Sep
Fld_0
        Typ_Lab
        Prm_Typ_MLis command is used to generate this list
Fld_0
        Typ_Lab
        Prm_Bnd_ command is used to set limits
    Fld_2
        Prm_Select a maximum of 5 items
        Typ_MLis
Bnd_1.0,5.0
        Lis_Multi Item 1,1
        Lis_Multi Item 2,2
        Lis_Multi Item 3,3
        Lis_Multi Item 4,4
        Lis_Multi Item 5,5
        Lis_Multi Item 6,6
        Lis_Multi Item 7,7
        Lis_Multi Item 8,8
        Lis_Multi Item 9,9
        Lis_Multi Item 10,10
Fld_0
    Typ_Sep
Fld_0
        Typ_Lab
        Prm_Typ_Idx command is for parent-child lists
        Fld_3
            Prm_Select one item from each list
            Typ_Idx
                Idx_Parent 1,Child 1 of Parent 1,1
                Idx_Parent 1,Child 2 of Parent 1,2
                Idx Parent 1, Child 3 of Parent 1,3
                Idx Parent 1,Child 4 of Parent 1,4
                Idx_Parent 1,Child 5 of Parent 1,5
```

```
    Idx_Parent 2,Child 1 of Parent 2,10
    Idx_Parent 2,Child 2 of Parent 2,15
    Idx_Parent 3,Child 1 of Parent 3,17
    Idx_Parent 4,Child 1 of Parent 4,-1
    Idx_Parent 4,Child 2 of Parent 4,0
    Idx_Parent 4,Child 3 of Parent 4,-21
    Idx_Parent 4,Child 4 of Parent 4,212
    Idx_Parent 4,Child 5 of Parent 4,18
    Idx_Parent 4,Child 6 of Parent 4,19
    Idx_Parent 4,Child 7 of Parent 4,20
:E END
:!
```

This function block for $\mathbf{F 2}$ produces the dialog box shown in Fig. 11.28. Similar to function block F1, clicking on OK in the dialog box issues the command F2 with appropriate arguments. In this case, however, the number of arguments is not predetermined due to the presence of the Multiple Selection List. For example, if items 1, 3, and 5 from the Multiple Selection List and Parent 4/Child 3 from the Parent Child List are selected, then ANSYS issues the following command input:
F2, 1, 3, 5, -21

On the other hand, if only item 3 from the Multiple Selection List and Parent 4/Child 3 from the Parent Child List are selected, then ANSYS issues the following command input:

F2, 3,-21


Fig. 11.28 Dialog box generated by function block F2.

Note that there are a minimum and maximum number of items that can be selected from the Multiple Selection List. In this case, their values are set to be 1 and 5 , respectively, using the Bnd_command.

Finally, function block F3 is created. Append the following text in the USERF . GRN file, after the function block for $\mathbf{F 2}$ :

```
:N Fnc_F3
:S 0, 0, 0
:T Command
:A Function F3
:D Function F3 Heading
Cmd_F3
    Fld_0
        Typ_Lab
        Prm_Type the description of function F3 here
Fld_0
        Typ_Sep
Fld_0
        Typ_Lab
        Prm_Typ_Int command is used for integer input fields
Fld_2
        Typ_Int
        Prm_Typ_Int
Fld_3
        Typ_Int2
        Prm_Typ_Int2
Fld_5
        Typ_Int3
        Prm_Typ_Int3
    Fld_0
        Typ_Sep
        Fld_0
        Typ_Lab
        Prm_Typ_Real command is used for real input fields
        Fld_8
        Typ_Real
        Prm_Typ_Real
        Fld_9
        Typ_Real2
        Prm_Typ_Real2
    Fld_11
        Typ_Real3
        Prm_Typ_Real3
    Fld_0
        Typ_Sep
Fld_0
        Typ__Lab
        Prm_Typ_Char command is used for character input fields
        Fld_14
        Typ_Char
        Prm_Typ_Char
```

```
Fld_0
    Typ_Lab
    Prm_Typ_Logi command is used for logical input fields
    Fld_15
    Typ_Logi,No,Yes
    Prm_Typ_Logi
:E END
:!
```

This function block produces the dialog box shown in Fig. 11.29. When the logical field, created using the Typ_Logi command, is unchecked, the corresponding argument has the value 0 ; otherwise (when checked), the value 1 is assigned.

## STEP 6

Copy the new control files, USERM. GRN and USERF.GRN, located in the temporary directory and paste them in the Working Directory where they will be accessed and processed by ANSYS. In this example, the following directory is considered to be the Working Directory: C: \user \.

## STEP 7

Add the file names of the new control files, USERM. GRN and USERF .GRN, to the menulist80.ans file. Copy the menulist80.ans file located in the temporary directory, paste it in the Working Directory, and, finally, open it using a text editor. Its contents should look similar to the following:

```
C: \Program Files \(\backslash\) Ansys Inc \(\backslash v 80 \backslash A N S Y S \backslash g u i \backslash e n-u s \backslash u i d 1 \backslash U I M E N U . G R N\)
C: \Program Files \(\backslash A n s y s\) Inc \(\backslash 880 \backslash A N S Y S \backslash g u i \backslash e n-u s \backslash u i d l \backslash U I F U N C 1 . G R N\)
C: \Program Files \(\backslash A n s y s\) Inc \(\backslash v 80 \backslash A N S Y S \backslash g u i \backslash e n-u s \backslash u i d l \backslash U I F U N C 2\). GRN
\(C: \backslash\) Program Files \(\backslash\) Ansys Inc \(\backslash v 80 \backslash A N S Y S \backslash g u i \backslash e n-u s \backslash u i d l \backslash M E C H T O O L\). AUI
```

Add the following two lines at the bottom of the file:

```
C:\user\USERM.GRN
C:\user\USERF.GRN
```


## STEP 8

Start ANSYS with C: \user as the Working Directory and verify that the customized GUI is working properly. If there are problems, edit the files in the temporary directory, then copy and paste them in the C:luser directory and restart ANSYS for verification.

## STEP 9

Once completed, the menulist80.ans file can be copied to additional Working Directories; the customized GUI will be available when ANSYS is started using those directories as the Working Directory. Figure 11.30 shows a portion of the newly created menu structure.


Fig．11．29 Dialog box generated by function block F3．

## ANSYS Main Menu

Preferences
曰 Preprocessor
$\square$ Demo Menu
目 Function F1
Function F2
Function F3
田 Element Type
田 Real Constants
Fig．11．30 Customized menu structure．

## STEP 10

Although the menu structure and the dialog boxes are working properly，the data collected through them have not been processed，i．e．，F1，F2，and F3 are not commands．ANSYS Macros are used for processing the data collected from customized dialog boxes．These files are written in the ANSYS Parametric Design Language（APDL）．In this case，three macro files are necessary：F1．MAC，F2．MAC，and F3．mAC．Macro files must have their extensions as MAC．If they reside in the Working Directory，they are automatically recognized by ANSYS．Otherwise，the／PSEARCH command
is used to point to their location. Macro files corresponding to customized dialog boxes expect arguments in the sequence they are entered in the dialog boxes. The first number in the dialog box data set corresponds to ARG1 in the macro file. A maximum of 19 scalar parameters can be passed on to each macro file from dialog boxes, i.e., ARG1, ARG2, ..., ARG9, AR10, AR11, ..., AR19. In the following subsection, this point is clarified.

### 11.6.2 GUI Modification for Obtaining a Random Load Profile

In this subsection, ANSYS GUI is customized to have an additional menu item with two functions under the Preprocessor. The first function collects data to use the external file RANDOM.EXE of Sec. 11.5.3 for generation of random loading as a function of time. The second function reads the random loading from an external file and plots the variation. Following the steps outlined in Sec. 11.6.1, the following menu block is created:

```
:F UM.GRN
:D User Menu
:I 0, 0, 0
:!
:N Men_Preproc
:S 0, 0, 0
:T Menu
:A Preprocessor
:C )/nopr
:C ) !*get,_z1,active,,routin
:C )!*IF,_z1,NE,17,THEN
:C /PREP7
:C )!*ENDIF
:C //go
:D Preprocessor (PREP7)
Men_UVBA_P7_T1
Men_UVBA_P7_T2
Men_UVBA_P7_T3
Men_Random
Men_ElemType
Men_Real_copy
Fnc_R_too
Men_Material
Fnc_DYNA_Shell
Men_Section_Defs
Sep_
Men_Modeling
Sep_
Men_Meshing
K_LN(lsdyna)
Sep_
Men_Trefftz_Dom
K_LN(ELECSTAT)
Sep_
Men_CheckCtrl
Men_NumCtrl
```

```
Men_Archive
Sep_
Men_CoupCeqn
K_LN(lsdyna+flotran)
Sep_
Men_FLOTRAN
K_LN(LSDYNA)
Sep_
Men_DYNAPREP
K_LN(1sdyna)
Sep
K_LN(lsdyna)
Men_FSI
K_LN(1sdyna)
Men_MFSET
K_LN(1sdyna+ROMES)
Sep_
K_LN(1sdyna+ROMES)
Men_ROM_Tool
K_LN(1sdyna)
Sep_
Men_Radiosity
Sep_
K_LN(ALPHA)
Men_QA_Test
K_LN(1sdyna)
Men_Loads
K_LN(1sdyna)
Sep_
Men_MultiPhys
Men_CrePath
Men_UVBA_P7_B1
Men_UVBA_P7_B2
Men_UVBA_P7_B3
:E END
:!
:N Men_Random
:S 0, 0, 0
:T Menu
:A Random Loading
:D Random Loading
Fnc_Generate
Fnc_ReadPlot
:E END
:!
```

As indicated in the first line of the listing, the control file for menus is named UM. GRN and the internal name for the customized menu block is Random, which is included at the bottom of the listing. The control file for customized functions is expected to have two function blocks: one for Generate and the other for Readplot. The contents of the control file, UF . GRN, for customized functions are as follows:

```
ADVANCED TOPICS
```

```
:F UF.GRN
```

:F UF.GRN
:D User Defined Functions
:D User Defined Functions
:I 0, 0, 0
:I 0, 0, 0
:!
:!
:N Fnc_Generate
:N Fnc_Generate
:S 0, 0, 0
:S 0, 0, 0
:T Command
:T Command
:A Generate Load
:A Generate Load
:D Generate Random Loading
:D Generate Random Loading
Cmd_F3
Cmd_F3
Fld_0
Fld_0
Typ_Lab
Typ_Lab
Prm_[GENERATE] Generates random loading
Prm_[GENERATE] Generates random loading
Fld_0
Fld_0
Typ_Sep
Typ_Sep
Fld_2
Fld_2
Typ_Real
Typ_Real
Prm_Duration of loading
Prm_Duration of loading
Fld_3
Fld_3
Typ_Int
Typ_Int
Prm_Number of data points
Prm_Number of data points
F1d_4
F1d_4
Typ_Real2
Typ_Real2
Prm_Range of load (MIN \& MAX)
Prm_Range of load (MIN \& MAX)
Fld_0
Fld_0
Typ_Lab
Typ_Lab
Prm_seed numbex for random number generation
Prm_seed numbex for random number generation
Fld_6
Fld_6
Typ_Int
Typ_Int
Prm_Seed number
Prm_Seed number
: E END
: E END
:!
:!
:N Fnc_ReadPlot
:N Fnc_ReadPlot
:S 0, 0, 0
:S 0, 0, 0
:T Command
:T Command
:A Read/Plot
:A Read/Plot
:D Read and Plot Random Loading
:D Read and Plot Random Loading
Cmd_F3
Cmd_F3
Fld_0
Fld_0
Typ_Lab
Typ_Lab
Prm_[ReadPlot] Reads and plots random loading data
Prm_[ReadPlot] Reads and plots random loading data
Fld_0
Fld_0
Typ_Sep
Typ_Sep
Fld_3
Fld_3
Prm_Select one option
Prm_Select one option
Typ_Lis_OptionB
Typ_Lis_OptionB
Lis__Read,1
Lis__Read,1
Lis_Read and Plot,2
Lis_Read and Plot,2
:E END
:E END
:!

```
:!
```

The resulting menu structure appears in the ANSYS GUI, as shown in Fig. 11.31. The dialog boxes corresponding to function blocks Generate and ReadPlot are shown in Fig. 11.32 and 11.33, respectively.

ANSYS Main Menu
Preferences
$\square$ Preprocessor
$\square$ Random Loading
Generate Load
圜 Read／Plot
田 Element Type
田 Real Constants
田 Material Props
Fig．11．31 Customized menu structure for random loading data generation．


Fig．11．32 Dialog box corresponding to function Generate．


Fig．11．33 Dialog box corresponding to function ReadPlot．

Two macro files corresponding to the function blocks are created and stored in the Working Directory. The first macro file is Generate.mAC:

```
TOT=ARG1 !STORE ARGUMENTS IN
ND=ARG2 !
MN=ARG3 !
MX=ARG4 !
SEED=ARG5 !
/OUTPUT,RANDOM,DAT !REDIRECT OUTPUT TO
    ! RANDOM.DAT
*VWRITE,TOT,ND,MN,MX,SEED ! WRITE PARAMETERS TO FILE
(E16.8, 2X, F8.0,2X, E10.2, 2X, E10.2, 2X,F8.0)
/OUTPUT !REDIRECT OUTPUT TO OUTPUT
    ! WINDOW
/SYS,RANDOM.EXE !EXECUTE RANDOM.EXE
```

The macro file Readplot . MAC is given as

```
* SET,ARY
*SET,TIM ! INITIALIZE PARAMETER TIM
*SET,LOAD ! INITIALIZE PARAMETER LOAD
*DIM,ARY,ARRAY,2*ND ! DIMENSION PARAMETER ARY
*DIM,TIM,TABLE,ND !DIMENSION PARAMETER TIM
*DIM,LOAD,TABLE,ND !DIMENSION PARAMETER LOAD
*VREAD,ARY(1),RANDOM,OUT ! FILL ARY ARRAY FROM RANDOM.OUT
(2(E16.8,2X)) !FORMAT STATEMENT
*DO,I,1,ND !LOOP OVER DATA POINTS
TIM(I)=ARY(2*I-1) !SEPARATE ARRAY ARY INTO TIM
LOAD (I) =ARY (2*I) !AND LOAD ARRAYS
*ENDDO ! END LOOP OVER DATA POINTS
*IF,ARG1,EQ,2,THEN
/AXLAB,X,TIME !DEFINE X-AXIS LABEL
/AXLAB,Y,LOAD !DEFINE Y-AXIS LABEL
*VPLOT,TIM(1),LOAD(1) ! PLOT TWO ARRAYS AGAINST EACH
! OTHER
* ENDIF
```

Finally, a copy of the external executable file RANDOM. EXE is stored in the Working Directory. The input shown in Fig. 11.34 produces the random loading shown in Fig. 11.35, which is obtained using the Readplot function.

### 11.6.3 Function Block for Selecting Elements Using a Pick Menu

The final example for demonstrating the ANSYS GUI environment involves a simple selection function. The function block is called Selel and is given below.


Fig. 11.34 Input for random load generation through a dialog box.


Fig. 11.35 Variation of load vs. time generated using Readplot function block.

```
:N Fnc_Selel
:S 0, 0, 0
:T Cmd_P
:A Element select
:D Pick elements
Inp_P
Cmd_ESEL
    Fld_2
    Typ_Def_S
    Fld_5
Prm_Pick elements within the global element
    TYp_ELEM
    Min_1
    Cnt_s
    PCN_1
    PFM_1
Cal_Fnc_Store
:!
```

The expression in the third line, :TCmAp, declares that this function block is for creating a Pick Menu. Similarly, the line Inp_P is used for picking, and it must be the first line in the data controls block. The line TYp_ELEM states that elements are to be picked. A minimum of 1 element must be picked, based on the specification Min_1. The line Cnt_s specifies that the selected entities produce an unordered list and that there is no limit on how many entities can be picked. The command PCN_1 specifies that a set of picked entities should not be ordered for retrieval picking. The command PFM_1 forces ANSYS to use commands FITEM and FLST for processing of the picked items. Finally, a call to another function block, named Store, is made in the line Cal_Fnc_Store.

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## INDEX

Absorption, 551, 552
Action Field, 98, 133, 147
Active coordinate system, 99, 114, 357, 362
Active Window Field, 131
Adaptive descent, 618
Adaptive meshing, 24
Adding, 118
Ambient temperature, 478, 523, 532
Amplitude, 444
Analysis disciplines, 16, 17, 85, 150
Analysis options, 27, 149, 150, 408, 435, 445
Analysis type, 27, 150
Anand's model, 579
Anisotropic, 93
ANSYS
Basic commands, 297
Extracting information, 314
File structure, 27
General Postprocessor, 27
Help System, 31, 32, 89, 93,
297, 324, 326, 647, 654
Contents, 33
Index, 33
Search, 33
Main Menu, 26
Modifying results, 654
Modifying the GUI, 654
Operators/functions, 303
Parametric Design Language, 15, 297
Preprocessor, 26, 83
Programming with, 317

Time History Postprocessor, 27
Verification Manual, 35, 324, 326
Verification problems, 326
APDL commands, 298
Approximating functions, 1,41 ,
See also Interpolation
functions
Arc(s), 108, 339
Arc-length, 568
Area(s), 4, 106, 109
Elements, 136
Integrals, 48, 72, 75
Primitives, 112
Arithmetic operations, 170
Automatic meshing, 135
Automatic time stepping, 155, 566, 567
Axisymmetric, 365
Elements, 366, 383
Shell elements, 391
Axisymmetry, 20, 346
Batch Mode, 15, 297
Beams, 337
Begin Level, 25
Bisection, 152, 566, 567
Block Header Section, 657
Body forces, 236
Body loads, 154
Boolean operators, 118
Bottom-up approach, 106
Boundary, 2
Integral, 202
Nonlinearity, 565

Boundary conditions, 13, 27, 149, 153, 187
Forced, 188
Natural, 188
Box Zoom, 133
Brick element, 342
Changing element attributes, 141
Changing-status nonlinearity, 19
Clearing/deleting mesh, 143
Combined mode, 15
Combined plasticity/creep, 579, 599
Commands
*GET, 314
*VWRITE, 321
/OUTPUT, 321
ADPL, 299
General Postprocessor, 298, 302
Preprocessor meshing, 298, 301
Preprocessor solid model generation, 298, 300
Session/database, 298, 299
Solution, 298, 302
Compatibility, 40
Complete polynomial, 40
Completeness, 40
Components, 148
Composite(s), 355, 573, 624
Layered plate, 397
Concatenating, 140, 349
Conduction, 478
Conformal elements, 40
Constraint equations, 621,624
Contact
Analysis, 605, 606, 607
Area, 565
Nonlinearity, 565
Sliding, 606
Contour plots, 165
Control Files, 655
Convection/convective, 153, 478

Convergence, 37
Criterion, 159
Failure, 159
Monotonic, 42
Tolerance, 567
Coordinates
Area, 46
Cartesian, 99, 103
Centroidal, 46, 48
Cylindrical, 99
Cylindrical (polar), 103
Generalized, 43
Global Cartesian, 99
Global coordinate systems, 43, 99
Length, 44
Local coordinate systems, 43, 100
Natural, 43
Spherical, 99
Toroidal, 99
Coupled degrees of freedom, 621
Creep, 19, 578
Strain rate, 599
Curved boundaries, 66
Cut boundaries, 631
Data button, 160
Data Controls Section, 657
Data/file options, 161
Database File, 28
Defining parameters, 304
Degrees of freedom, 3, 17
coupled, 621
Delete entities, 130
Density, 550, 551
Dependent variable, 187
Detach, 143
Differential operator matrix, 239
Diffusion, 522
Direct approach, 2, 5
Direct generation, 83
Dividing, 121
DO loops, 317

Duplicate entities, 144
Dynamic analysis, 433
Dynamic Mode Field, 133
Effective stress, 579
Eigenvalue, 12
Eigenvalue buckling, 17, 150, 403
Eigenvector, 12
Element matrix, 2
Attributes, 96
Characteristic, 6, 8
Force vector, 240
Right-hand-side, 8
Stiffness, 240
Element table data, 167
Element type(s), 27, 85
Elements, 4, 85, 575
Connectivity, 9, 28, 83, 105, 194
Isoparametric, 64, 217, 269, 278, 421
Linear quadrilateral isoparametric, 269, 278
Linear triangular, 255
Link, 366
Select using Pick Menu, 675
Shell, 373
Volume, 136
Ending Section, 658
Entities, 105
Equilibrium iterations, 566
Error File, 28
Evaluation of integrals, 68
Executing external file, 652
Expansion Pass, 444, 638, 646
External files, 647
Extrusion/sweeping, 124
Fluid, 87
Forcing frequency range, 444
Forcing function, 187
Fracture mechanics analysis, 421
Galerkin's method, 187

Gaussian
Integration, 68, 275
Points, 68
General Postprocessor, 26, 160
Generation Pass, 636, 638
Geometric isotropy, 42
Geometric nonlinearity, 18, 569
Global coordinate systems, 43, 99
Global element sizes, 137
Global equilibrium equations, 241
Global node numbering, 4, 37
Gluing, 121
Gobal model, 629
Graph data, 160
Graphical User Interface, 15
Development, 662
Modification, 671
Graphics Window, 29, 96
Gravity, 154, 366
Grid and triad, 103
GUI. See Graphical User Interface
Harmonic analysis, 17, 150, 444
Heat
Conduction, 4, 17, 197, 522
Diffusion, 550
Flow, 5, 6, 7, 153, 167
Flux, 4, 7, 16, 153, 167, 478, 523, 527
Generation, 94, 154, 169, 197, 477, 478
Insulation, 552
Transfer, 477
Help System, 31, 32, 89, 93, 297, 324, 326, 647, 654
Contents, 33
Index, 33
Search, 33
Hyperelasticity, 19
IF statements, 318
Impact load(ing), 460, 465
Inertia loads, 154

Information Field, 98
Initial conditions, 149, 154
Initial residual stresses/strains, 236, 237
Input Field, 29
Input file(s), 15, 30, 35, 297, 304, 305, 307, 325, 647
Interactive mode, 15
Interpolation functions, $1,64,68$, 85
Shape functions, $51,53,54$, 56, 59, 61, 64, 68, 189, 190, 200, 204, 218, 240, 251, 269, 277, 280
Isoparametric elements, 64, 217, 269, 278, 421
Isotropic, 93
Isotropic hardening, 579, 599
Jacobian matrix, 68, 272
Jobname, 15
Keeping/deleting original entities, 128
Keyoptions, 89
Keypoints, 106
Large deflection/rotation, 18, 159, 460, 565, 570
Large strain, 569
Length coordinates, 44
Line(s), 4, 106, 107
Elements, 136
Integrals, 68
Search, 567
Linear, 93
Buckling analysis, 403
Line element, 54
Quadrilateral element, 62
Spring, 5
List, 31
Data, 160
Entities, 130
Nodal solution, 170
Results, 170
Load step(s), 154, 566
Files, 155

Loading conditions, 150
Loading sequence, 565
Local coordinate systems, 43, 100
Local node numbers, 37
Log File, 28, 325
Macro files, 297, 322
Main Menu, 29, 31
Mapped meshing, 135, 136, 139, 301, 489
Master DOFs, 434, 446, 636
Material nonlinearity, 18, 565, 578
Material properties, 27, 92
Matrices
Global stiffness, 242
Global system, 8
Global system of equations, 3 , 8

Jacobian, 68, 272
Lumped mass, 438
Material property, 237
Singular, 12
Singular coefficient, 158
Stiffness, 6
System stiffness, 3
MenuCtrls, 31
Merge entities, 144
Mesh(ing), 134
Density, 24, 137
Manipulation, 141
Refinement test, 25
Method of weighted residuals, 187
Modal analysis, 17, 150, 434
Frequencies, 434
Shapes, 434
Superposition, 453
Model generation, 149
Model geometry, 27
Modeling operations, 83
Moisture
Concentration, 550, 551
Diffusion, 477, 549
Diffusivity, 550

Moving/copying, 128
Nano-indentation test, 607, 613
Natural coordinates, 43
Newton-Raphson, 566, 567
Nodal unknowns, 189
Nodal values, 52
Nodes, 1, 3
Common, 3
Master, 625
Slave, 625
Nonlinear, 93
Elastic, 19
Geometry option, 570
Load-displacement relationship, 565
Numbering, 134
Compressing, 144
Controls, 143
Operators /functions, 298
Orthotropic material, 93
Out-of-balance load vector, 566
Output Window, 29
Overlap, 120
Pan/Zoom Field, 133
Parameters, 31
Pascal triangle, 41
Path plots, 165
Phase change, 17
Picking entities, 96
Pick/Unpick Field, 96
Picking Style Field, 96
Planar/reflective symmetry, 21
Plane strain, 346
Conditions, 249
Plane stress, 346
Conditions, 248
Plastic deformation, 583
Plasticity, 19, 578
Plates/shells, 373
Plot, 31, 131
Controls (Ctrls), 31, 131
List entities, 134
Results, 163
Post-buckling analysis, 573

Postprocessing, 160
Postprocessor, 16
Potential energy, 3, 236
Predictor algorithm, 567
Preprocessor, 26, 83
Preprocessor level, 83
Pressures, 153
Primary heat transfer, 17
Primitives, 83, 105, 112
Processor Level, 25, 26
Quadratic line element, 56
Quadratic triangular element, 59
Radiation analysis, 543
Radiosity solver, 478, 543
Rate Control Field, 133
Rate-independent plasticity, 599
Rayleigh-Ritz method, 187
Read results, 161
Reading Input File, 647
Real constants, 27, 89
Repetitive/translational
symmetry, 21
Results files, 28
Rigid constraints, 625
Rotate, 131
Rotate Field, 133
Rotational symmetry, 20
Rotations, 569
Round-off error, 24
Sandwich panel, 625
Select, 30
Components, 144
Operations, 144, 309
Smart sizing, 139
Solid modeling, 83, 105
Solution, 150
Commands, 298
Controls, 150, 459, 460, 462, 528,567,591
Processor, 26
Trial, 566
Solve load step files, 157
Sparse solver, 567
Spectrum, 150

Spline, 109
Static analysis, 17, 329
Static condensation, 636
Steady-state, 523
Analysis, 478
Heat transfer, 477
Step size, 567
Stiffness matrix multiplier, 609
Strain rate equation, 599
Stress intensity factors, 421
Stress stiffening, 18, 404, 568
Structural, 85
Analysis, 16, 150
Damping, 465, 608
Stability, 569
Structural system, 236
Submodeling, 25, 629
Substeps, 566, 567
Substructuring, 636, 642
Superelements, 621,636
Generation, 642
Subtracting, 120
Surface
Contact, 606
Target, 606
Tractions, 236
Symmetry conditions, 20
Temperature, 7, 153, 550
Change, 154, 238
Text Field, 98
Thermal, 85
Thermal analysis, 16
Thermomechanical analysis, 17 , 412, 541
Three-dimensional problems, 342
Time dependence, 18
Time History Postprocessor, 26, 160, 444
Time step, 155
Size, 155
Time-dependent loads, 155
Time-related parameters, 155
Title, 84

Toolbar, 29, 32
Top-down approach, 106, 112
Transient
Analysis, 459, 520
Dynamic, 17
Heat transfer, 170, 478
Trial solution, 566
Truss(es), 329
Elements, 330
Structure, 333
Units, 92
Use Pass, 636, 644
Utility Menu, 29
Variational method, 3, 187, 235
Vector(s), 202
Global nodal load, 242
Inter-element, 202
Nodal displacements, 241
Nodal forces, 241
Nodal unknowns, 6
Plots, 165
Right-hand-side, 6
Unknowns, 3
Verification
Manual, 35, 324, 326
Problems, 326
View model, 131
Viewing Direction Field, 132
Viscoelastic/viscoplastic, 578
Viscoelasticity, 19, 579
Viscoplasticity, 579
Analysis, 592
Volume elements, 136
Volume(s), 4, 106, 110
Weighted residuals, 3, 187
Wetness parameter, 551
Working Directory, 15, 28
Working plane, 31, 99, 102
Write Load Step File, 157
Writing data to external ASCII files, 648
Zoom, 131, 132


[^0]:    Main Menu > Preprocessor > Modeling > Create > Volumes > Cone > By Dimensions

[^1]:    ${ }^{\text {i }}$ Colors have not been used in the printed version of the figures. See the accompanying CD-ROM for color versions of the figures.

[^2]:    Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Areas $>$ Mapped $>$ Concatenate $>$ Lines Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Volumes $>$ Mapped $>$ Concatenate $>$ Areas Main Menu $>$ Preprocessor $>$ Meshing $>$ Mesh $>$ Volumes $>$ Mapped $>$ Concatenate $>$ Lines

[^3]:    Main Menu $>$ Solution $>$ Define Loads $>$ Apply $>$ Thermal $>$ Temperature
    Main Menu > Solution > Define Loads > Apply $>$ Thermal $>$ Heat Flux
    Main Menu >Solution > Define Loads > Apply $>$ Thermal $>$ Heat Flow
    Main Menu > Solution > Define Loads > Apply > Thermal > Convection

[^4]:    Main Menu > General Postproc > Path Operations > Plot Path Item > On Graph
    Main Menu > General Postproc > Path Operations > Plot Path Item > On Geometry

[^5]:    Main Menu > General Postproc > Element Table > Plot Elem Table
    Main Menu > General Postproc > Element Table > List Elem Table

[^6]:    Main Menu > General Postproc > Path Operations > Plot Path Item > On Graph

[^7]:    Main Menu > Solution > Define Loads > Apply > Thermal > Heat Generat > On Elements

[^8]:    *GET, Par, Entity, ENTNUM, Item1, IT1NUM, Item2, IT2NUM

[^9]:    ${ }^{1}$ Colors have not been used in the printed version of the figures. See the accompanying CD-ROM for color versions of the figures.

[^10]:    Main Menu > General Postproc > Options for Outp

[^11]:    Main Menu > Solution > Define Loads > Apply > Structural > Pressure > On Nodes

[^12]:    Main Menu > General Postproc > Plot Results > Contour Plot > Nodal Solu

[^13]:    Main Menu > Solution > Define Loads > Apply > Structural > Displacement > On Nodes

[^14]:    Utility Menu > WorkPlane > Local Coordinate Systems > Create Local CS > At Specified Loc

[^15]:    Main Menu > Preprocessor > Real Constants > Add/Edit/Delete

[^16]:    Main Menu > General Postproc > Path Operations > Map onto Path

[^17]:    Main Menu >Solution > Define Loads > Apply > Structural > Displacement > On Nodes

[^18]:    Main Menu > Solution > Define Loads > Apply > Structural > Force/Moment > On Nodes

[^19]:    $\begin{aligned} & Y \\ & Z\end{aligned} 2 \times 345 \begin{array}{llllll}10111213141516171819202122232425262728293031\end{array}$

[^20]:    Main Menu $>$ Solution $>$ Define Loads $>$ Delete $>$ Structural $>$ Force/Moment $>$ On Keypoints

[^21]:    Material > New Model

[^22]:    ${ }^{1}$ Colors have not been used in the printed version of the figures. See the accompanying CD-ROM for color versions of the figures.

[^23]:    Main Menu > Solution > Define Loads > Apply > Thermal > Convection > On Nodes

