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Multiple Myeloma

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Multiple Myeloma What & Why

Multiple myeloma [MM] is a complex hematologic malignancy. It is crucial for any practitioner to be aware of the pathophysiology, presenting clinical manifestations, differential diagnoses, implications for care, and other disease specific details to avoid

misdiagnosis and provide proper treatment/referral. As a future nurse practitioner providing care in an orthopedic office in addition to the surgical environment

there is a high likelihood of caring for a patient with this disease. American Cancer Society (2018)

- Estimates "30,770 newly diagnosed cases of
- MM in the United States 12.770 deaths
- 1 out of 132 life time risk of developing MM" (p.1) **Presentation: Multiple**

Mveloma

Plasma cell malignancy, insidious onset Malignant plasma cells multiply in bone marrow and produce an overabundance of monoclonal protein (Michels & Peterson, 2017). Monoclonal protein produced: IgG, IgM, IgA (most commonly). IgE, IgD (rarely), Kappa or Lambda light chain proteins (Michels & Peterson, 2017). African Americans are twice as likely to develop MM compared to Caucasians. African Americans also present earlier in life (Michels & Peterson, 2017).

- 85% of patients diagnosed with MM are older than 65 years (Michels & Peterson, 2017).
- 65 years old is current median age of diagnosis (Brigle & Rogers, 2017). 46.6% 5 year survival rate (Brigle & Rogers, 2017).
- Findings on presentation for patients with multiple myeloma presented in Table 1 (Michels & Peterson, 2017). World Health Organization
- Classification, differentiation between: 1. Multiple myeloma [MM]
- 2. Monoclonal gammopathy of undetermined significance [MGUS]
- 3. Solitary plasmacytoma of the hone
- 4. Extraosseus plasmacytoma
- 5. Monoclonal immunoglobulin deposition diseases
- (Brigle & Rogers, 2017)

Underlying Pathophysiology

Demise of TP53 tumor suppressor

in disease stage).

2017, p. 226).

(epigenetic events):

respectively.

activity (can indicate poor prognosis)

Nuclear Factor kB (NF-kB) regulation is

inactivated by mutations (presents late

Adhesion molecules between MM cells

and bone marrow stroma cells

Downregulation results in MM cell

Changes not related to gene sequencing

DNA hypomethylation and

Target genes related to DNA

methylation include links to,

expression as result of histone

Disease progression related to genetic

Clonal evolution: clonal competition,

result of diversity and aggression.

Clones are more genetically complex

and produce a dominant cell line as a

survival of the fittest clone.

Rogers, 2017, p. 227).

demethylation.

complexity:

(Brigle & Rogers, 2017)

(Brigle & Rogers, 2017)

"dexamethasone resistance, cell

adhesion, and cell signaling" (Brigle &

Increase in oncogene transcription and

growth outside of bone marrow and

"development of stromal-independent

plasma cell Leukemia" (Brigle & Rogers

hypermethylation in early progression

of MGUS to MM and late stage MM

regulated by NF-kN cells

Plasma Cell Disease: Incurable, Heterogeneous

- Monoclonal plasma cell crowding in bone marrow due to overgrowth (Brigle & Rogers, 2017). Followed by increased production of
- immunoglobulins or immunoglobulin chains that also crowd other cells in bone marrow (Brigle & Rogers, 2017). Pathoaenesis of abnormal plasma cell production:
- Cyclin D protein dysregulation (early event) Almost always a progression from MGUS.
- (Brigle & Rogers, 2017). Two karyotype subclasses of MM
- 1.Hyperdiploid- chromosomes 3,5,7,9,11,19, and 21 contain extra copies or trisomies. 2. IaH Translocation- at 14q32 of IgH
- locus with the possibility of various individual partner genes that all result in an upregulation of cyclin D proteins. 40% of MM cases are hyperdiploid (better prognosis than patients with any of the various IgH translocation abnormalities).
- 30% of MM cases present with IgH translocations 15% of MM patients have both trisomies and IgH translocations. (Brigle & Rogers, 2017)
- Further genetic alterations that potentiate growth of abnormal plasma cells: Chromosome 13 loss MYC and RAS oncogene initiation
- Changes of chromosome 1 copies
- Table 1. Findings on Presentation for Patients with Multiple Myeloma (Michels & Peterson, 2017)
- Symptom or Laboratory Finding Percentage of Patients Anemia (hemoglobin < 12 g per dL 73 [120 g per L]) 58 Bone pain Elevated creatinine (> 1.3 mg per dL 48 [115 µmol per L]) Fatigue or generalized weakness 32 Hypercalcemia (calcium > 10.1 mg per 28 dL [2.52 mmol per L]) 24 Weight loss

Significance of Pathophysiology: Bone **Marrow Microenvironment**

Bone marrow: location of myeloma cell growth Bone marrow stromal cells: adhere to MM cells and expansion

- 2 compartments of bone marrow: 1) cellular 2)non-cellula Cellular compartment: hematopoietic and non-hematopoietic cells: Myeloid cells, T and • B lymphocytes, natural killer cells (NK), osteoclasts, bone marrow stromal cells, bone marrow-derived mesenchymal stromal cells, fibroblasts, osteoblasts, adipocytes, endothelial cells, blood vessels Non-cellular compartment: Extracellular
- matrix, cytokines, chemokines, growth factors, exosomes produced by cellular compartment
- Both compartments and their interactions crucial to MM progression (Brigle & Rogers, 2017)



Nalker, R. C., Brown, T. L., Jones-Jackson, L. B., Blanche, D. L. & Bartel, T. (2012). Imaging of multiple myeloma and related plasma cell dyscrasias. The Journal of Nuclear Medicine, 53, 1091-1101. doi:10.2967/jnumed.111.098830

Adipocytes

Bone marrow-derived mesenchymal stem cells

- "pluripotent potential" (can develop into fibroblasts, adipocytes, osteoblasts, (Rigle & Rogers, 2017, p. 228) Support growth and overproduction of MM cells through secretion of adhesion molecules, cytokines, and chemokines, Osteoblast and Osteoclast activity
- Osteoblast suppression Osteoclast activation due to MM cell secretion of cytokines. IL-6 secretion by
- osteoclast stimulates further MM cell proliferation Overall increase of osteolysis leading to
- bone lesions (Figure 1). Therapy targeting osteoclasts is key in maior reduction or delay in skeletal patholoav related to MM. (Brigle & Rogers, 2017)

and, "support tumor cell proliferation, migration, drug resistance, and expression of anti-apoptotic proteins" (Rigle & Rogers, 2017 p. 227)

Activate NF-kB to secrete cytokines which enhances the growth, adhesion, and overproduction of MM cells.

- Interleukin (IL)-6 a major cytokine involved in vascular endothelial growth factor (VEGF) secretin which increases vascularity inside the bone marrow. Enable microRNAs (miRNAs) to promote production of monoclonal light chains leads to end organ damage (Michels & Peterson,
- effect on drug resistance (Brigle & Rogers, 2017)

2017). symptomatic Patients may present without symptoms but MM is then discovered by laboratory analysis and findings of hypercalcemia, proteinuria, and anemia. (Michels & Peterson, 2017) Long Term Effects: Serum Hyperviscosity and dysfunctional plasma cell infiltration and

MM cell growth and overproduction through exosome release, also may have 2017).

Diagnostic Criteria According to Michels and Peterson, 2017: Both criteria must be met:

Signs & Symptoms

nausea, vomiting, weakness, fatigue,

anemia, bone pain, renal dysfunction

hypercalcemia, pathologic fractures.

Refer to Table 1 (Michels & Peterson,

weight loss, recurrent infections,

Nonspecific:

- Clonal bone marrow plasma cells ≥ 10% or biopsy-proven bony or extramedullary plasmacytoma
- (2) Any one or more of the following myeloma-defining events: Evidence of end-organ damage that can be attributed to the underlying plasma cell proliferative disorder, specifically:

Hypercalcemia: serum calcium > 1 mg per dL (0.25 mmol per L) higher than the upper limit of normal or > 11 mg per dL (> 2.75 mmol per L)

Renal insufficiency: creatinine clearance < 40 mL per minute per 1.73 m2 (0.67 mL per second per m2) or serum creatinine > 2 mg per dL (177 µmol per L) Anemia: hemoglobin > 2 g per dL (20 g per L) below the lower limit of normal, or a hemoglobin value < 10 g per dL (100 g per L) Bone lesions: one or more osteolytic lesions on skeletal radiography, CT, or positron emission tomography/CT Clonal bone marrow plasma cells ≥ 60% Involved: uninvolved serum free light chain ratio ≥ 100 (involved free light chain level must be ≥ 100

mg per L) More than one focal lesion on MRI studies (≥ 5 mm size) " (p. 376, Table 3). Implications for Care

International Staaina System usina serum beta2-microalobulin and albumin levels International Myeloma Working Group (IMWG recently presented new Revised International Staging System to include chromosomal alterations. Predicts, "progression free and

overall survival and has been recommended for use in future studies". (Michels & Peterson, 2017, p.378)

Implications for Care Treatmen

- Mveloablative chemotherapy using two or three drug combinations and autologous stem cell transplantation (ASCT)
- Common chemotherapy drugs used Corticosteroids: dexamethasone,
- methylprednisolone, prednisone Alkylating agents: Melphalan (Alkeran),
- cyclophosphamide Immunomodulatory drugs: thalidomide,
- lenalidomide(Revlimid) Proteasome inhibitors: bortezomid (Velcade), carfilzomib (kyprolis)
- (Michels & Peterson, 2017, p.380) Special Treatment Considerations
- Avoid nephrotoxic medications and studies using contrast media due to renal dysfunction
- Acute kidney injury with MM: at least 3L per day of intravenous normal saline in addition to dexamethasone to decrease serum light chain if elevated. (Michels & Peterson, 2017)
- Bone Modifying Agents
- Zoledronic acid (Reclast), pamidronate, denosumab
- Refer to the American Society of Clinical Oncology's 2017 Clinical Practice Guidelines update cited in the additional sources section for specific recommendations, selection, dosing, duration, and monitoring of BMAs with MM

(Anderson, Ismaila, & Kyle, 2017) High Risk of Thromboembolic Events

- Prevention with low-molecular-weightheparin or warfarin with target INR of 2 to 3 (American Society of Clinical Oncology).
- Stratify thromboembolic risk factors for treatment and consider aspirin alone for low-risk patients (IMWG). (Michels & Peterson, 2017, p.380) High Risk of Recurrent and/or Life Threatenina Infection
- Vigilance and speed in recognition and treatment
- May use prophylactic antibiotics: Trimethorprim/sulfamethoxazole. fluoroquinolone, penicillin
- Intravenous immune globulin Prophylactic antivirals if taking
- proteasome inhibitors due to risk of varicella-zoster virus
- Immunizations: pneumococcal pneumonia. Haemophilus influenza. influenza virus, especially for stem cell transplant patients.

Peterson, 2017).



Anemia Common with MM : Restrictive transfusion policy supported for hemoglobin levels less than 7 g per dL (Michels &



Conclusion

Providers should be aware of evaluation

and management of MM in relation to other

plasma cell dyscrasias for prompt diagnosis

and treatment. Refer to Figure 1 in the

March 2017 American Family Physician

Journal Article, Volume 95, Number 6,

Treatment for an excellent flowchart to

assist with management and treatment

Providers should also be aware of plasma

Overall diagnostic workup for suspicion

of MM includes, "lab studies, urine studies,

evaluation" (Brigle & Rogers, 2017, p.231).

n Oncology Nursing Volume 33, Number 3,

Myeloma (Brigle & Rogers, 2017, Table 7, p.

Populations that should be considered

regarding survival improvement: older

populations (75+ years) and minority

populations (Costa et al., 2016)

decade has allowed progression of study of

Significant expansion of molecular

pathophysiology of MM over the last

various MM pathoaenesis pathways

(Lawasut et al., 2013).

Pathobiology and Diggnosis of Multiple

Current findings confirm survival

improvements for patients with MM.

For specific diagnostics refer to Seminars

bone marrow biopsy, and radiographic

cell diseases that progress to MM (MGUS

and Smoldering MM) (Brigle & Rogers,

Multiple Myeloma: Diagnosis and

(Michels & Peterson, 2017).

2017).

231).



Endothelial Cells Promote neovascularization in bone marrow Contribute to overproduction of MM cells by supplying massive quantity of IL-6.

- Tregs Enhanced in bone marrow because of
- factor
- Increase in apoptotic ligands and receptors • Decrease, suppression, and alternation of
- function in antitumor mediated cells (CD8+ cytotoxic T cells and NK cells)

· Secrete leptin, adiponectin, IL-6 which are involved in MM cell growth. Large quantity of adipocytes in bone marrow

- Immune System Alterations • Increase in immunosuppressive cells (CD4+
- increased levels of IL-6 and a growth

(Brigle & Rogers, 2017)