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Bonding modes of species chemisorbed

on tungsten single crystal faces

Ъy

Leon William Anders

A Dissertation Submitted to the Graduate Faculty in Partial Fulfillment of The Requirements for the Degree of DOCTOR OF PHILOSOPHY

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Major: Physical Chemistry

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CHAPTER I. INTRODUCTION

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In recent years it has become experimentally possible to study the chemisorption of gases on single crystal planes of metal surfaces. This permits observation of chemisorption on metal surfaces of known structure, in contrast to earlier polycrystalline techniques. In the absence of direct spectroscopic observation, many indirect methods have been utilized in an attempt to assign molecular structures to various adsorbed states. Low energy electron diffraction (LEED) studies have uncovered the symmetry relationship between the substrate and the adsorbate. Flash desorption spectroscopy (FDS) has been used to determine the number of bonding states and kinetics and the energy of desorption from these states. From these techniques it has been clearly demonstrated in the last few years that gases chemisorbed on surfaces exhibit marked variations in structure and kinetics with crystal structure, but the investigators have not been able to assign positively bonding structures to the chemisorbed species.

This work was stimulated by the investigation by Tamm and Schmidt (1, 2) of hydrogen chemisorption on various tungsten single crystal surfaces. In the case of hydrogen on a W(100) surface, other investigators (3, 4, 5) have also reported results. From these findings, various conflicting models have been proposed. Tamm and Schmidt (1) proposed a mixture of atomic and molecular species; Estrup and

Anderson (3) as well as Madey and Yates (4) proposed that hydrogen atoms were bonded at a single site. Later Yonehara and Schmidt (5) proposed that hydrogen atoms were bonded at different sites.

The early part of this work was concerned with developing a qualitative description of chemisorbed hydrogen on W(100), W(110), and W(111). This was extended, following a suggestion by Dr. L. S. Bartell, by an Extended Hückel Molecular Orbital (EHMO) investigation to elucidate the bonding and structure of chemisorbed species on an array of atoms representing a truncated surface. The W(100) crystal face was chosen as the adsorbing surface, with hydrogen nitrogen, and oxygen as adsorbates. The hydrogen (1-5), nitrogen (6-9), and oxygen (10-12) on W(100) systems all have received experimental investigation, and it is generally concluded that the adsorbates are dissociated in their most strongly adsorbed state not only on a polycrystalline surface, but also a single crystal surface of tungsten. Although they are dissociatively adsorbed, molecular hydrogen and nitrogen desorb in FDS experiments. Molecular oxygen will not desorb in FDS experiments, but the dissociatively adsorbed oxygen desorbs as oxides of tungsten and even as atomic oxygen (13).

In developing an overall understanding of the bonding between a chemisorbed species and a surface, the literature was searched for experimental evidence which would yield insight into the problem.

Besides the above diatomic gases which have been used in chemisorption studies to probe the properties of a surface, NO and CO also have been used extensively. On polycrystalline tungsten, Yates and Madey (14) reported that NO chemisorbs nondissociatively at room temperature, but dissociation occurred at elevated temperatures ($\sim 500 - 600^{\circ}$ K). From the NO dissociation molecular nitrogen is desorbed and the majority of the oxygen is desorbed as oxides of tungsten. Weinberg and Merrill (15) report similar results on the W(100) surface and also found that when all of the nitrogen is desorbed, a typical (2 x 1) oxygen pattern remains (10).

In contrast to these common diatomic molecules, CO has been assumed to be associatively adsorbed on tungsten surfaces. Clavenna and Schmidt (16) recently reported CO was probably associatively adsorbed on a W(100) surface with the possible exception of the adsorbed state which desorbed at ~ 1425°K. More recent results (17-19) have been reported supporting a dissociatively adsorbed model for CO on polycrystalline tungsten. In Table I, a comparison is made between the desorption of associatively adsorbed states (molecular) and dissociatively adsorbed states (atomic) for the diatomic molecules of interest. The behavior of hydrogen, oxygen, nitrogen, and nitrous oxides has been well documented with regard to the separation of the two possible adsorbed states. As indicated by Goymour and King (17),

	D _e (Kcal/mole)	Molecular State		Atomic State	
Molecule		Desorption Temperature (°K)	Activation Energy for Desorption (Kcal/mole)	Desorption Temperature (°K)	Activation Energy for Desorption (Kcal/mole)
H ₂ (1)	104	100°		450° 600°	26 32
0 ₂ (11-13)	118			Desorbs as 1	netal oxides
NO (14, 15)	150	520°		1100°	47 (N ₂)
N ₂ (8)	225	1 80°	10	1250°	74
CO (16)	256	400°	21	(?) 1025° 1150° 1425°	(?) 57 62 74

Table I. A comparison of thermal desorption data of diatomic molecules from a W(100) surface.

it has been maintained in the literature that even the states of CO desorbing at a temperature from ~1000 - 1425°K have resulted from associatively adsorbed CO. In Table I the desorption activation energies and temperatures for these high temperature states of CO are similar to those for nitrogen states known to be dissociatively adsorbed.

It therefore seemed desirable to re-examine the question of CO dissociation on tungsten, and a study of mixed adsorption of $C^{16}O$ and $^{18}O_2$ on (100), (110) and (111) tungsten single crystal faces was under-taken for this purpose. Very recently and while the present study was in progress, Goymour and King (17-19) have presented evidence, based on polycrystalline FDS experiments, for dissociative adsorption of CO on tungsten and have presented a model for such adsorption.

CHAPTER II. MOLECULAR ORBITAL INVESTIGATION OF ATOMIC SPECIES CHEMISORBED ON A W(100) SURFACE

Introduction

Today there is no available experimental technique or any combination of techniques which can be used to assign unequivalently a bonding structure between an adsorbate species and the atoms of a surface. The advent of metal single crystal surfaces in chemisorption research has brought about increased speculation as to the preferred bonding structures for chemisorbed species. Unfortunately, no general overall understanding has emerged which can serve to predict sites and preferred structure for new systems based on previous knowledge. Certainly, there is no reason to suspect that chemisorbed atoms like hydrogen, nitrogen, oxygen, etc. would exhibit strange and unusual bonding properties when interacting with the atoms of a metal surface. The bonding properties of most atoms are generally known and their roles in a particular chemical compound are usually predictable. If a surface chemist were able to develop guidelines from which bonding between an adatom and surface atoms could be predicted, he would be able to understand and describe more concretely surface processes of current concern.

In this study, a molecular orbital approach was used in hopes that some guidelines could be established for predictions of chemisorption bonding. A simple molecular orbital technique developed by Hoffmann (20), which was sensitive to bonding geometries, was utilized in this study. Hoffmann extended the Hückel Molecular Orbital technique which depended only on the interactions of adjacent atoms to include all interactions. Therefore, it is referred to as the Extended Hückel Molecular Orbital (EHMO) technique. A listing of the computer program used for the molecular orbital calculations can be found in Appendix D.

Results and Discussion

This study was restricted to the tungsten (100) surface with adatoms of hydrogen, nitrogen, and oxygen. The application of the EHMO technique to the surface "molecules" formed by

$$W_n(100) + X \rightarrow W_n(100) X$$

is described in Appendix A. The tungsten surface was represented by a finite array of tungsten atoms (W_n) to which the adatom could bond forming the surface "molecule" $W_n X$.

The results and a full description of the molecular orbital calculations for hydrogen, nitrogen, and oxygen atoms chemisorbed to the various sites on a W(100) surface are found in Appendices A, B, and C respectively.

The tungsten (100) surface has three symmetric bonding sites as illustrated in Fig. 1a. An adatom at site A will be coordinated to





Fig. 1. Bonding structure of chemisorbed atoms on the W(100) surface (a - W(100) surface A - 5 CN site, B - 2 CN site, C - 1 CN site, b - hydrogen bonded at 1 CN sites in a (1 x 1) structure, c - nitrogen bonded at 5 CN sites in a C(2 x 2) structure and d - oxygen bonded at 2 CN sites in a (2 x 1) structure).

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five surface atoms (5 CN), at site B to two surface atoms (2 CN), and at site C to a single surface atom (1 CN). It was found that hydrogen (Appendix A) preferred the 1 CN site, the nitrogen (Appendix B) preferred the 5 CN site, and oxygen (Appendix C) preferred the 2 CN site. It appears that each adatom selected a site on the tungsten (100) surface determined by the bonding properties of the adsorbate atom.

The most direct tests of the above bonding models are the LEED patterns of stable structures of these adatoms. The adsorption of hydrogen on a W(100) surface leads to a (1×1) pattern (3, 5) or one hydrogen atom for every surface atom. Such a structure, as shown in Fig. 1b, is consistent with hydrogen bonded at every 1 CN site. After annealing a nitrogen adsorbed layer a $C(2 \times 2)$ structure (7, 8) is produced on a W(100) surface. From the knowledge (Appendix B) that nitrogen selected the 5 CN site because of the ability of the p_x and p_y orbitals to bond to the four atoms surrounding the 5 CN site, it is not unreasonable that nitrogen atoms would select a $C(2 \times 2)$ surface structure (Fig. 1c). In this structure, a nitrogen atom is surrounded by four empty adjacent 5 CN sites. The p_x and p_y orbitals of nitrogen can form strong bonds with the four atoms around this site. LEED has observed a stable (2×1) oxygen (10) surface structure on the W(100) surface. A (2x1) surface structure results if the oxygen atoms occupy the 2 CN sites on a W(100) surface as illustrated in Fig. 1d. Each

oxygen atom is bonded to two surface atoms, and each surface atom to one oxygen atom.

Summary and Conclusions

The bonding properties of chemisorbed species on metal surfaces appear to be very similar to those expected from their usual bonding characteristics. Where there is no activation for dissociation (usually the case for diatomic molecules) the atoms of an adsorbing molecule will select sites where they can best satisfy their valency requirements and maximize the bonding interactions with the surface atoms. These guidelines should be able to help understand additional chemisorbed systems. One example will be used to indicate the transferability of derived results. In Fig. 2, a W(110) surface is illustrated. There are three probable sites for a chemisorbed species to be located. If an oxygen atom were bonded to this surface, either site A or site B would satisfy its divalent property. At site B, the tungsten atoms are touching and probably would not form a stable bond with an oxygen atom. Site A appears attractive to an atom with a higher valency, because of its high coordination number. Although the four atoms are not equally spaced at this site, two atoms are separated by 3.16Å and the other two by 4.46Å. To the oxygen atom, this site is very similar to the 2 CN site on the W(100) surface. If the oxygen atoms were to produce a surface structure as illustrated in Fig. 2,



Fig. 2. Proposed surface structure of oxygen atoms at 2CN sites on a W(110) surface.

the bonding between the oxygen atoms and the surface atoms would be identical to that on the W(100) surface. Each oxygen is bonded to two tungsten atoms and these tungsten atoms are not bonded to any other oxygen atom. This surface structure is a $p(2 \ge 1)$ and a resulting LEED pattern should be observed. Tracy and Blakely (21) have reported such a structure for oxygen adsorption on a W(110) surface.

The usefulness of this basic understanding will be illustrated in the next chapter when selecting preferred sites for C and O adatoms and associatively adsorbed CO and O_2 on the single crystal surfaces of tungsten (100, 110, and 111) crystal faces.

CHAPTER III. CO AND ¹⁸O₂ CO-ADSORPTION ON SINGLE CRYSTAL SURFACES OF TUNGSTEN

Introduction

The interaction of CO with tungsten has been extensively studied and reviews of the literature have recently been published (22-24). On tungsten as on most metal surfaces, it has generally been accepted that CO is nondissociatively chemisorbed. All common diatomic gases including CO exhibit multiple peaks in the desorption spectra from Flash Desorption Spectrometry (FDS), but for CO, unlike the other diatomic gases, it has been assumed that even the high temperature desorption peaks result from undissociated adsorbed CO.

Very recently Goymour and King (17-19) have shown that a model based on dissociative chemisorption of CO can more satisfactorily explain their experimental results. They have presented the following facts in a brief history (17) of related experimental evidence which has been reported in favor of and against the hypothesis of dissociative adsorption:

(1) The argument of Ehrlich, Hickmott, and Hudda (25) that their failure to find traces of C or O residues on their surfaces following CO chemisorption implies associative chemisorption is not compelling. The major desorption product from a mixture of C and O adatoms appears always to be CO, and in this case it is scarcely surprising if little C and O residues remain after heating a CO adlayer. Thus Becker et al. (26) and Singleton (27) found that carbon impurities are removed as CO from a tungsten surface when that surface is heated to 2000°K in an oxygen ambient.

(2) The energy balance for dissociative adsorption is (24)

$$q_{co} = q_{o} + q_{c} - D_{co}$$

where q_x is the adsorption heat of species x and D_{CO} is the dissociation energy of gaseous CO (256 Kcal/mole). From Brennan and Graham (28) $q_0 = 163$ Kcal/mole, and from Redhead (29) $q_c \cong 170$ Kcal/mole; therefore q_{CO} is 77 Kcal/mole. If the activation energy for adsorption is zero, then q_{CO} can be compared with the activation energy for desorption of CO from the various surfaces of tungsten which are listed in Table II. These comparisons indicate that it is thermodynamically feasible for CO to be dissociated on tungsten surfaces.

(3) Madey et al. (30) observed isotopic mixing in the β states of CO from mixed adsorption of ${}^{12}C^{18}O$ and ${}^{13}C^{16}O$. Their explanation was based upon a four-centered complex formed by the interaction of two adsorbed CO molecules, although dissociative adsorption is also a very plausible explanation. Madey and Yates (31) later reported that isotopic exchange occurred on W at temperatures between 1300 and 2200°K where the steady state CO coverage is very

Tungsten Surface	State	E _{d0} (Kcal/mole)
Polycrystalline	<u>8</u> 1	53
W (4)	β ₂	100
W(100) (12)	$\beta_{3} \ (<0.1)$	93
	β_3 (>0.1)	74
	β ₂	62
	β_1	57
W(110) (13)	β_1	40 - 55
	β ₂	67

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Table II.	A comparison of activation energy of desorption for (CO on
	various tungsten surfaces.	

small and it is difficult to see why desorption should take place preferentially through a four-centered complex. It was also shown by Madey et al. (30) that isotopic exchange does not occur in the α states but only in the β states of CO.

(4) Propst and Piper (32), using the relatively low resolution energy-loss technique, observed a strong vibrational band attributed to the β -CO state on W(100) which they claimed could only be explained by the presence of atomically adsorbed C and O.

These facts along with the kinetic and surface stoichiometry data reported by Goymour and King (17) and also the mixed adsorption of CO and ${}^{18}O_2$ very recently reported by Goymour and King (19) strongly suggest that β -CO is dissociatively adsorbed on tungsten. The object of the present work was to use ${}^{18}O_2$ which will dissociatively adsorb on W surfaces as a chemical probe in mixed adsorption experiments on single crystal surfaces of tungsten (100, 110, 111) to test the hypothesis of dissociatively adsorbed CO. Isotopic mixing of oxygen in the desorbed CO molecules could serve as primary evidence for the hypothesis. It should also be possible to vary the ratio of C: ${}^{16}O:{}^{18}O$ adsorbed on the surface and if the CO is dissociatively adsorbed the ratio of C ${}^{16}O$ and C ${}^{18}O$ desorbed from the W surfaces should also vary. If more than one β (dissociated CO) state exist on a particular surface, then the ability of the adsorbed $^{18}O_2$ to displace any or all of these states, or the ability of the coadsorbed $^{18}O_2$ to alter the energy of CO desorption (e.g. lowering the temperature of the maximum desorption of an adsorbed state) are also of interest.

Experimental Procedure

The ultra-high vacuum Pyrex apparatus is schematically illustrated in Fig. 3. During bake-outs to ~350°C, the system was pumped by a mercury diffusion pump trapped with liquid nitrogen. At the end of the bake-out, with the oven still hot, the pumping was shifted to an Ultek 20 L/sec differential ion pump. After cooling of the apparatus, background pressures of ~ 10^{-10} Torr were routinely obtained. Pressures were measured with an Electron Technology Inc. ion gauge equipped with a thoriated iridium filament controlled with a Granville-Phillips ionization gauge controller. The reaction cell was also equipped with a quadrupole mass spectrometer (QMS) (Electronic Associates Inc.) enabling partial pressures of the various gaseous species to be monitored. The tungsten single crystals were cut from a 7 mm diameter rod of 99.998% purity. The rod was properly aligned by back reflectance x-ray patterns according to the procedure outlined by Wood (33). Sample wafers ~20 mil thick were spark cut from the rod to expose selected single crystal surfaces. These wafers were mechanically polished to a mirror finish. Back reflectance x-ray



Fig. 3. A schematic diagram of the experimental apparatus for flash desorption spectrometry.

patterns indicated that the three crystal surfaces (100), (110), and (111) were oriented to within 1.5° of the nominal orientations. For each of the crystals the edge area was about 10% of the total area. Each single crystal surface sample was mounted to a 15 mil tungsten main support rod which was attached to a pyrex dewar via a 45 mil tungsten rod. A thermocouple (W vs. Re) was also attached to each sample. Generally, the crystals were heated in vacuo to 2200° K by electron bombardment until a background pressure of less than 10^{-8} Torr could be obtained (about 24 hours); this was interspersed with flashes to at least 2600° K for several minutes each. During the cleaning period, the crystals were also flashed to 2000° K for several minutes in an O₂ atmosphere of ~ 10^{-6} Torr. At the end of this procedure the crystal could be flashed to above 2000°K without a measurable pressure increase.

The heating of the crystal by electron bombardment was accomplished by maintaining the crystal +400 volts above filament potential and by preselecting an emission current with which to bombard the sample. With a controller which monitored the emission current and signalled a programmable filament power supply, a filament temperature could be maintained to give the proper emission current. The calibrated thermocouple was used to obtain time-temperature curves for various emission currents for each sample. Emission currents could then be selected which gave linear heating rates over

the temperature ranges of interest. The flash desorption curves were all recorded as ion current (y axis) vs. time (x axis); the latter could be converted to a temperature scale from the above heating curves.

In the following experiments, the desorption species of principal concern were $C^{16}O$ and $C^{18}O$ molecules. Therefore, the QMS was set to sweep mass (m/e⁻) 28-30 at a rate of six sweeps per second. The ion current of each mass on each sweep was recorded by a high speed chart recorder enabling each species to be monitored simultaneously for each flash desorption spectrum.

The dosing gases were controlled by variable leak values. The ${}^{18}O_2$ was obtained from YEDA R&D Co. LTD, a Division of Miles Laboratories, and was 99.29% ${}^{18}O_2$. The C¹⁶O was C.P. Grade, 99.5% CO, obtained from Matheson Co. The CO was vacuum distilled prior to use as the dosing gas. The dosing pressure of each gas was monitored with the QMS which was calibrated with the ionization gauge.

Results and Discussion

<u>Mixed adsorption of CO and $\frac{18}{02}$ on a W(100) surface</u>

A series of CO desorption spectra obtained from a W(100)surface after varying exposures to CO at room temperature are shown in Fig. 4. These are very similar to those found in the literature (16, 34) for the interaction for CO with this single crystal surface. Yates and King (34) reported three bonding states: α which desorbed at ~400°K and two higher temperature states β_1 and β_2 . The desorption temperatures for these states were not reported. Clavenna and Schmidt (16) also reported an α state desorbing at ~ 400°K; with an ultra-high vacuum system which apparently had a higher pumping speed, they were able to resolve the next state which compares to the β_1 state of Fig. 4 into states which they labeled β_1 desorbing at ~1040°K and β_2 desorbing at ~1130°K. These two states combined desorbed at a temperature centered at ~1100°K and compares with the β_1 state in Fig. 4 which desorbs at ~1060°K and with the β_1 states reported by Yates and King (34). The higher temperature state β_2 in Fig. 4 desorbs at ~1450°K which compares with a state desorbing at ~1480°K and labeled β_3 by Clavenna and Schmidt. The spectra reported by Clavenna and Schmidt better resolve the $\beta_1 - \beta_2$ group and the higher temperature β_3 state than the spectra of Fig. 4. The β_2 state in Fig. 4 is only clearly observed in curves A and B and unfortunately because



Fig. 4. Flash desorption spectra of CO adsorbed on a W(100) surface (A - 2.4L, B - 4.5L, C - 9.0L, and D - 18.2L (1 Langmuir (L) = 10⁻⁶ Torr-sec.)).

TEMP. (°K)

14

4 6 8 10 12

x10²

of poor resolution at larger CO coverages (curves C and D of Fig. 4), the tail of the flash desorption spectra of the β_1 state covers up the β_2 state. The resolution of a flash desorption spectra can be improved primarily by controlling two experimental parameters: the pumping speed of the system and the heating rate. The design of the system and size of the pump determine the pumping speed, therefore, this can not be changed after construction of the system and in the experiments of this study, the heating rate was optimized for the best resolution possible. The only other observable difference between the spectra in Fig. 4 and those previously reported (16, 34) is the amount of CO desorbed in the α state. Yates (35) suggested that possibly the crystal was not cool enough during the dosing period to accommodate a larger amount of CO adsorbed in the α state. Yates and King (34) using ESD were able to resolve the α state into two substates α_1 (340°K) and α_2 (380°K). Yates (35) indicated that the α_1 state was very labile and could be desorbed by pumping on the system, whereas the α_2 state was more stable. Although cool water flowed through the dewar reservoir, Yates (35) thought that the temperature of the crystal might have been high enough during dosing to preclude formation of the α_1 state, so that only the α_2 state appeared in Fig. 4.

Using LEED and FDS, Anderson and Estrup (36) reported that two β states resulted from room temperature adsorption of CO on the

W(100) surface. The initial adsorption produced a (1×1) LEED pattern, and upon heating the surface to desorb the β_1 state at ~1150°K, the remaining CO, the β_2 state, produced a C(2 x 2) LEED pattern. This β_2 state desorbs at ~1380°K. These β states compare remarkably well with those found in this study and elsewhere (16, 34). They were able to readsorb CO upon the β_2 (C(2 x 2) structure) state to produce a structure indistinguishable from that resulting from adsorption on a clean surface.

The results of first dosing the W(100) surface with CO followed by a dose of ${}^{18}O_2$ are shown in Figs. 5-8. In these figures, the flash desorption curves of both $C^{16}O$ and $C^{18}O$ are shown (recorded simultaneously as previously explained). In Fig. 5, the surface was dosed with a 2.4L dose of $C^{16}O$ followed by increasing dosages of ${}^{18}O_2$ as indicated for curves B-F. It is observed that most of the $C^{18}O$ is desorbed in the β_1 state. It is difficult to observe any peak of $C^{18}O$ desorbing from the β_2 state, in fact at larger ${}^{18}O_2$ doses (curves E and F) even the $C^{16}O$ which desorbed from the β_2 state has been displaced. The amount of $C^{16}O$ desorbing in the α state increases slightly as the coverage of ${}^{18}O_2$ increases and seems to reach a limiting value as observed from curves E and F. The isotopic mixing of oxygen in the CO molecules desorbed from the β_1 state suggests CO has been dissociatively adsorbed. A suggestion that associatively



Fig. 5. Flash desorption spectra of CO from a W(100) surface predosed with 2.4L of C¹⁶O followed with increasing dosages of ¹⁸O₂ (A - 0.0L, B - 1.2L, C - 2.6L, D - 5.3L, E - 10.7L and F - 21.3L).


Fig. 6. Flash desorption spectra of CO from a W(100) surface predosed with 4.5L of C¹⁶O followed with increasing dosages of 18O₂ (A - 0.0L, B - 1.2L, C - 2.6L, D - 5.3L, E - 10.7L and F - 21.3L).



Fig. 7. Flash desorption spectra of CO from a W(100) surface predosed with 9.0L of C¹⁶O followed with increasing dosages of ¹⁸O₂ (A - 0.0L, B - 1.2L, C - 2.6L, D - 5.3L, E - 10.7L and F - 21.3L).



Fig. 8. Flash desorption spectra of CO from a W(100) surface predosed with 18.2L of CO followed with increasing dosage of ¹⁸O₂ (A - 0.0L, B - 1.2L, C - 2.6L, D - 5.3L, E - 10.7L and F - 21.3L).

adsorbed CO in an α state could be converted to dissociated CO in the β states as a result of heating during the flash desorption technique is not supported by the above facts. If CO were only associatively adsorbed at room temperature, then dosing with $^{18}O_2$ a surface with preadsorbed CO should block the sites needed for the conversion of α -CO to β -CO. This conflicts with results shown in Figs. 5-8; therefore, the CO must be adsorbing in the β states at room temperature. With larger ¹⁸O₂ doses (curves D-F), it also must be noted that some $C^{18}O$ is desorbed from the α state. Therefore, at very high coverages of $^{18}O_2$, the associatively adsorbed α -CO must be able to interact with the adsorbed $^{18}O_2$. In Fig. 6, where the surface was predosed with 4.5L dose of C¹⁶O followed with increasing $^{18}O_2$ doses, the observations are the same as those stated above. Figures 6 and 7 with predose 9.0L of $C^{16}O$, and Fig. 8 with predose of 18.2L of $C^{16}O$ all show that as the ${}^{18}O_2$ coverage increases, there is a decrease in the temperature of the desorption peak maximum for the β_1 state.

The flash desorption spectra of a set of experiments where a W(100) surface was predosed with varying amounts of ${}^{18}O_2$ always followed by the same dose of $C^{16}O$ are shown in Fig. 9. It is observed that the ${}^{18}O_2$ initially blocks sites for CO to adsorb in the β state, but creates sites for more CO to adsorb associatively in α states; as the



Fig. 9. Flash desorption spectra of CO from a W(100) surface predosed with increasing dosages of ${}^{18}O_2$ (A - 2.6L, B - 5.3L, C - 10.7L and D - 21.3L) followed by a constant dose of C¹⁶O (9.0L).

amount of preadsorbed ${}^{18}O_2$ increases, it eventually also blocks the CO from adsorbing in the α state. If the W(100) surface were saturated with ${}^{18}O_2$, it would prevent the surface from adsorbing any CO. From Fig. 9 it must also be observed that isotopic mixing of the oxygen in the desorbed CO molecules still occurs primarily in those desorbed in the β state.

It is possible to estimate the amount of CO desorbed from the surface by determining the area under each of the flash desorption curves. The total amounts of CO ($C^{16}O$ and $C^{18}O$) desorbed from all states in each of the four experiments (Figs. 5-8) are plotted in Fig. 10. It is evident that in a given experiment with a constant $C^{16}O$ predose, the amount of CO desorbed is independent of the ${}^{18}O_2$ coverage. This shows that ${}^{18}\text{O}_2$ is unable to displace any of the preadsorbed $C^{16}\text{O}$. Figure 11 plots the ratio of $C^{18}O:CO_{total}$ as a function of ${}^{18}O_2$ dose on the surface for each of the experiments in which C¹⁶O was preadsorbed. Each C¹⁶O predosed surface appears to reach a limiting ratio $(C^{18}O:CO_{total})$ as the amount of ${}^{18}O_2$ dosed increases to a saturated level. The limiting ratios for each predosed surface are plotted in Fig. 12 as functions of the $C^{16}O$ predose. The limiting $C^{18}O:CO_{total}$ ratio at low $C^{16}O$ dose appears to be ~ 0.75 and at high $C^{16}O$ dose appears to be ~ 0.24 .



Fig. 10. Total amount of CO (C¹⁶O + C¹⁸O) desorbed from a W(100) surface where the surface was predosed with C¹⁶O (A - 2.4L, B - 4.5L, C - 9.0L and D - 18.2L) with increasing dosages of ¹⁸O₂ (1.2L, 2.6L, 5.3L, 10.7L and 21.3L) in each set of C¹⁶O doses.



Fig. 11. The ratio of C¹⁸O/CO_{total} desorbed from a W(100) surface predosed with C¹⁶O (A - 2.4L, B - 4.5L, C - 9.0L and D -18.2L) with increasing dosages of ¹⁸O₂ (1.2L, 2.6L, 5.3L, 10.7L and 21.3L) in each set of C¹⁶O doses.



Fig. 12. The limiting ratio of C¹⁸O/CO_{total} desorbed from a W(100) surface predosed with C¹⁶O (2.4L, 4.5L, 9.0L, and 18.2L) followed by a saturated dose of ¹⁸O₂.

Before proposing a model which is consistent with the above results of mixed adsorption of $C^{16}O$ and $^{18}O_2$ on a W(100) surface, it would be appropriate to first summarize the experimental observations.

(1) The FDS spectra of CO desorbed from a W(100) surface exhibited three bonding states: α , β_1 , and β_2 . Because the amount of α desorbed was smaller than other reported results, it is assumed that only the more stable α_2 state was present on the W(100) surface.

(2) Preadsorbed CO was not displaced from the surface by dosing with $^{18}O_2$ (Fig. 10).

(3) The adsorption of ${}^{18}O_2$ on a CO covered surface eliminated the β_2 state of adsorbed CO. There was a small increase in CO desorbed in the α_2 state which appeared to reach a rather low limiting value. The fact the amount of CO desorbing in the α state does not continue to increase provides evidence that there is no conversion from an α state to β state. With increasing ${}^{18}O_2$ coverage, the temperature of maximum desorption for the β_1 state shifted to lower temperatures.

(4) From mixed adsorption of CO and ${}^{18}O_2$, isotopic exchange of oxygen occurred in the desorbed CO molecules primarily in the β_1 state and also to a very small extent in the α state at very high ${}^{18}O_2$ coverages.

(5) At very low preadsorbed coverages of CO followed by a

saturating dose of ${}^{18}O_2$, the ratio of $C^{18}O:CO_{total}$ desorbed appeared to reach a limiting value of ~.75. At a saturated coverage of CO followed by a saturating dose of ${}^{18}O_2$, the ratio of $C^{18}O:CO_{total}$ appeared to reach a lower limiting value of ~.24.

(6) With increasing coverages of preadsorbed ${}^{18}O_2$, it is possible to decrease the amount of CO which can be adsorbed on the surface. The β states are the first to be blocked by the preadsorbed ${}^{18}O_2$. As the sites for the β states are blocked, there is an increasing amount of CO desorbed in the α state, but at still higher ${}^{18}O_2$ coverages, the α state is also blocked for CO adsorption.

Goymour and King (17-19) have very recently published results for CO adsorption on polycrystalline tungsten from which they concluded CO was dissociatively adsorbed. They have proposed a model (17) based on the W(100) surface which was consistent with their kinetic and surface stoichiometric results from their polycrystalline work. The essence of the model proposed by Goymour and King is schematically illustrated in Fig. 13. They proposed that at room temperature a CO molecule would dissociate into C and O adatoms in the β state occupying the 5 CN sites marked A in Fig. 13. Nitrogen has been shown (Appendix B) to prefer the 5 CN site, and its 2p orbitals were influential in this preference; the carbon atom for the same reason should prefer this site. This is not, as shown in Appendix C, a



Fig. 13. Possible sites occupied by chemisorbed species resulting from co-adsorption of C¹⁶O and ¹⁸O₂ on a W(100) surface (A - 5 CN site, B - 2 CN site, C - 1 CN site, ● - C or O adatoms and ● associatively adsorbed oxygen).

preferred site for oxygen adsorbing alone, but presumably the carbon is able to influence the oxygen to select a 5 CN site when co-adsorbed (the LEED results indicate that C and O must select the same kind of site in CO adsorption). It has been shown (18) that, if the CO adsorption occurs by dissociation into these sites and requires the availability of 5 CN sites in adjacent pairs, then random adsorption will lead to a maximum coverage of CO in the β state in which only 85% of the available 5 CN sites are occupied. It was proposed (18) that the remaining 5 CN sites could adsorb an associated CO molecule in an α state or two associated CO molecules in an α state at the 2 CN sites (B in Fig. 13) adjacent to an empty 5 CN site. ESD (34, 37) has resolved two α states; the low temperature α_1 state is labile (35) and yields a CO^+ ion by ESD and the high temperature α_2 state yields an O^+ ion by ESD. These two α states might well correspond to the two α sites suggested by Goymour and King (18). Suppose the α_1 state is bonded at site B and the α_2 state at site A in Fig. 13. In the former case the carbon atom is presumably bonded to two tungsten atoms, and lies above their plane. Its desorption as CO^+ on ESD seems likely. In the latter case the carbon atom is coordinated to 5 tungsten atoms and is nestled down into the plane of surface atoms. The bonding of carbon to tungsten is presumed stronger, and to oxygen weaker, than at site B. Loss of O^+ from this configuration on ESD seems reasonable. If the C

and O atoms from the dissociated CO molecule in the β state are bonded at the 5 CN sites, it is very probable that the electrons would not be able to desorb any ionic species. Yates and King (34) have reported that they were unable to desorb any ions by the ESD technique when only the β states are present on a W(100) surface.

The observations which have been summarized above can readily be explained based upon the above model for dissociatively adsorbed CO. The initial interaction of a CO molecule with the W(100) surface is probably at either a 1 CN or 2 CN site (C or B of Fig. 13). If two adjacent 5 CN sites (A of Fig. 13) are available then the CO molecule will dissociate into C and O adatoms at these sites. If only one 5 CN site is empty, we will assume that the CO will bond associatively at this site. The bonding state of CO observed in the flash desorption spectra of Fig. 4 from a W(100) surface can be assigned bonding sites based on the above model for CO adsorption. The CO desorbed in the β_1 and β_2 states would be C and O adatoms bonded at the 5 CN sites. The CO desorbed in the α state (α_2) would be associatively adsorbed CO molecules occupying the 15% of 5 CN sites not occupied by C and O adatoms. The fact that CO desorbed as two states β_1 and β_2 from C and O adatoms in the same sites can be explained by lateral interactions as has been done in other systems (H on W(100) and N in W(100)). When the CO desorbs from the surface, as the β_1 state, it will be

assumed to be occurring by the recombination of adjacent C and O adatoms. The remaining C and O adatoms are undergoing rearrangement to form a more stable surface structure, a β_2 state, which desorbs at a higher temperature. This explanation is also consistent with the LEED observations of Anderson and Estrup (36).

The results summarized above where oxygen was co-adsorbed on the surface to serve as a chemical probe add support to the model of dissociatively adsorbed CO; the isotopic mixing of the oxygen in the CO molecules desorbed in the β states is particularly significant.

Goymour and King (19) have shown by ESD that oxygen adsorbed onto a CO saturated surface seemed to be identical in population and ESD cross section to the β_1 -O₂ state formed by oxygen adsorption on clean W with oxygen preadsorbed in the β_2 state. Although it does not desorb as a molecular oxygen, it has been described as an associated adsorbed O₂ species (38-40) on a clean polycrystalline tungsten surface. The primary ESD product of this species is an O⁺ ion. When heating the surface, this species undergoes a change which occurs between 600 and 900°K and the ESD product is no longer an O⁺ ion.

From the above discussion, it can be assumed that the oxygen adsorbed on W(100) with preadsorbed CO is molecular oxygen. From ESD results on polycrystalline tungsten, this molecularly adsorbed state begins to dissociate at about 600-900°K. Therefore, as the β_1

state of CO desorbs at ~1060°K, the carbon atoms have a random selection of the two isotopes of oxygen resulting in a mixture of $C^{16}O$ and $C^{18}O$ desorbed for the β states. The small amount of isotopic mixing in the α_2 state could result from the possibility that the associatively adsorbed CO in desorbing from a 5 CN site could interact with a molecularly adsorbed oxygen; the probability of such interaction should increase with increasing oxygen saturation and this is what is observed.

The β_2 state of dissociated CO has been proposed as resulting from C and O atoms in a C(2x2) structure formed after the β_1 state has desorbed and the C and O atoms have rearranged to seek a more stable bonding configuration. The presence of adsorbed oxygen could prevent this rearrangement and therefore all the CO would be desorbed at a temperature characteristic of the β_1 state. The presence of the adsorbed oxygen with which the surface tungsten atoms must also bond will cause those bonds between the tungsten atoms and those of the C and O adatoms to be weakened. Therefore, as there is an increase of oxygen adsorbed on a W(100) surface, preadsorbed CO (C and O adatoms) bonds should be weakened, and the CO should desorb at lower temperatures as observed.

Goymour and King (19) found that oxygen would displace part of the α -CO from their polycrystalline tungsten surface which was

saturated with CO. They found no displacement of the CO desorbed in the β state. In this study, oxygen did not displace any of the preadsorbed CO on a W(100) surface (Fig. 10). According to the model just presented, it would be difficult for the adsorbed oxygen which is assumed to be occupying sites above the surface atoms to displace any preadsorbed CO (which is assumed to be occupying 5 CN sites whether associatively (α) or dissociatively (β) adsorbed). On the other hand, \neg any CO preadsorbed in the more labile α_1 -CO state might be displaced. Goymour and King (19) reported the α -CO which was not displaced from their polycrystalline surfaces shifted to a higher desorption temperature. This supports the hypothesis of two bonding states of α -CO with the α_2 -CO being more stable and not displaced by the adsorbing oxygen.

In Fig. 9 it was observed that with an increasing preadsorbed oxygen dose the adsorption of CO in the β states was decreased while adsorption in the α_2 state was mutually increased. According to our model, the β states are dissociated and a pair of vacant adjacent 5 CN sites is needed to adsorb β -CO. Oxygen preadsorption will reduce adjacent pairs of empty sites much more than single sites (which suffice for associative α_2 -CO adsorption) and therefore as observed a decrease in CO desorbed as β and increased in α -CO. With increased coverages of oxygen, even these sites were blocked for CO adsorption and at a saturated coverage of oxygen, the surface should be blocked to any CO

adsorption.

By an assessment of sites available for adsorbed α -CO, β -CO and $18O_2$ it is possible to offer an explanation for the limiting ratio of $C^{18}O:CO_{total}$ which is desorbed from the surface. It has been assumed that the β states of adsorbed C and O atoms occupy 5 CN sites. With a low initial CO dose, the C atom will have one adjacent 5 CN site occupied by its initial oxygen partner and three empty adjacent sites. It can also be assumed that, as the ${}^{18}O_2$ adsorbs on this tungsten surface with low coverage of adsorbed CO, it dissociates and will be influenced by the C adatoms to occupy the surrounding three empty 5 CN sites. Then as the CO desorbed, the C atom will have a choice of one 160 and three 180atoms with which to combine. This should lead, at very low CO coverages, to a ratio of 3:4 (.75) for the $C^{18}O:CO_{total}$ desorbed from the surface, and this is what Fig. 11 shows. At saturated coverages of CO, Goymour and King (18) determined that about 15% of the 5 CN sites capable of adsorbing C and O atoms from dissociated CO were not occupied. These sites have adjacent 5 CN sites which are occupied, resulting from random adsorption.

On a tungsten surface which is saturated with preadsorbed CO, the β -CO is in the form of C and O adatoms at 5 CN sites in a (1×1) surface structure. A saturated dose of ${}^{18}O_2$ upon this C and O adatom layer results in co-adsorption and as reported above, oxygen is

molecularly bonded to the surface. The most reasonable structure for such a chemisorbed species is one bridging from site C to site C (Fig. 13). Each oxygen will be bonded to a single tungsten atom, and a single bond will exist between the oxygen atoms. The co-adsorbed oxygen molecules could easily occupy every other set of surface atoms in every row. This (2×1) structure of associatively adsorbed ${}^{18}O_2$ is formed on the surface (Fig. 13).

Therefore, from a surface with saturated preadsorbed CO followed with a dose of saturating ¹⁸O₂, 15% of the CO desorbed is α -C¹⁶O from associatively adsorbed CO. The remaining 85% of the 5 CN sites are occupied by C and O adatoms in a (1 x l) structure. This (1 x l) unit mesh defines a surface structure of one adatom per unit mesh and assuming LEED can not distinguish between C and O adatoms, then the (1 x l) structure contains either one carbon or one oxygen atom. It would require two unit meshes (2 x l) to contain a CO molecule. If the C adatom in this structure will recombine only with O adatoms, then the carbon has a choice of ¹⁶O and ¹⁸O if the associatively adsorbed ¹⁸O₂ has dissociated at the temperatures of β -CO desorption. If the co-adsorbed oxygen dissociates according to

then, as assumed, each (2×1) unit mesh should contain three oxygen

atoms $({}^{16}\text{O} + 2{}^{18}\text{O})$ and one carbon atom. It is very unlikely that the above dissociation is complete when β -CO desorbs. If one-half the oxygen molecules have dissociated, coupled with the probability that only one-half of these are dissociating next to a C adatom, then the carbon has a choice of oxygen isotopes in the ratio of ${}^{16}\text{O}$: ${}^{18}\text{O}$ which is 1:.5. Therefore, from the 85% of the sites occupied by C and O adatoms, 57.5 molecules of CO will be desorbed from every 100 sites occupied. Based upon the above assumptions, one-third of these CO molecules would be $C^{18}\text{O}$. From the arguments above, the limiting ratio of $C^{18}\text{O}$:CO_{total} desorbed should be ~.24 which compares well with the value of ~.23 determined from Fig. 12.

Mixed adsorption of CO and $\frac{18}{02}$ on a W(110) surface

The flash desorption spectra of CO from a W(110) surface are shown in Fig. 14. At low coverages only a β state is present which desorbs at ~1100°K. As the coverage increases, an α state desorbing at a much lower temperature (~400°K) appears and the β state increases and appears to split into 2 states, β_1 and β_2 , desorbing at ~920°K and ~1100°K respectively. In the present study, ${}^{16}O_2$ occasionally was found in the ambient when heating the crystal to higher temperatures for cleaning. It was also observed that the size of the β_1 state depended upon the amount of ${}^{16}O_2$ which appeared in the ambient. This dependence is depicted in Fig. 15 which indicates that the splitting



Fig. 14. Flash desorption spectra of CO adsorbed on a W(110) surface (A - 2.5L, B - 4.4L, C - 9.2L and D - 18.6L).



Fig. 15. Flash desorption spectra of CO from a W(110) where CO was dosed (18.6L) simultaneously with ¹⁸O₂ (A - ~.05L, B - .15L, C - .60L and D - 1.80L) from the ambient.

of the β state into β_1 and β_2 results from interactions of co-adsorbed species on the surface.

May and Germer (41) reported two peaks in their flash desorption spectra of CO from a W(110) surface, a β state at ~1100°K and an α state at ~450°K. Kohrt and Gomer (42) observed, in their step desorption spectra from CO adsorbed on a W(110) surface at 100°K, a virgin state (v) desorbing at ~390°K and two β states, β_1 desorbing at ~980°K and β_2 desorbing at ~1120°K. From their findings, they concluded that heating the surface to 400-600°K after it was saturated with virgin CO caused some of the CO to desorb, but that some of the virgin CO was converted irreversibly to the β states. They reported that the surface was able to accommodate approximately one-half the number of CO molecules in beta configurations as in virgin configuration. After formation of the beta layer, they report that additional CO could be adsorbed on the surface at 100°K. In their step desorption spectra this CO desorbed at ~150°K and at ~300°K. They labeled these states α_1 and α_2 respectively.

In the mixed adsorption experiments of CO followed by ${}^{18}O_2$ (Figs. 16-19), it was observed that as the amount of ${}^{18}O_2$ dosed on the surface increased, the β_2 state was gradually displaced until all the CO desorbed as the β_1 state. The temperature of maximum desorption for the β_1 state even shifted to lower temperature as the amount of



Fig. 16. Flash desorption spectra of CO from a W(110) surface predosed with 2.5L of $C^{16}O$ followed with increasing dosages of ${}^{16}O_2$ (A - 0.0L, B - 1.2L, C - 2.6L, D - 5.3L, E - 10.7L and F - 21.3L).



Fig. 17. Flash desorption spectra of CO from a W(110) surface predosed with 4.4L of C¹⁶O followed with increasing dosages of ¹⁸O₂ (A - 0.0L, B - 1.2L, C - 2.6L, D - 5.3L, E - 10.7L and F - 21.3L).



Fig. 18. Flash desorption spectra of CO from a W(110) surface predosed with 9.2L of C¹⁶O followed with increasing dosages of ¹⁸O₂ (A - 0.0L, B - 1.2L, C - 2.6L, D - 5.3L, E - 10.7L and F - 21.3L).



Fig. 19. Flash desorption spectra of CO from a W(110) surface predosed with 18.6L of C¹⁶O followed with increasing dosages of ¹⁸O₂ (A - 0.0L, B - 1.2L, C - 2.6L, D - 5.3L, E - 10.7L and F - 21.3L).

 ${}^{18}O_2$ on the surface increased. As the amount of ${}^{18}O_2$ on the surface increased following the predose of CO, there was no large increase in the amount of CO desorbed in the α state. Such an observation would have been expected if α -CO was being converted to β -CO by heating the surface during the flash desorption.

Figures 16-19 also show that isotopic mixing of the oxygen in the CO molecules desorbed from the β states is extensive, but that it is negligible in those desorbed from the α state. In a given experiment the CO molecules (C¹⁶O and C¹⁸O) desorb simultaneously as evidenced by their substantially identical FDS temperatures of peak maxima.

If the surface is predosed with ${}^{18}O_2$, the oxygen is able to block the adsorption of CO. In Fig. 20, a small dose of ${}^{18}O_2$ (curve A) displaces the β_2 state first and causes a slight increase in the amount of CO desorbed in the α state. With larger doses of ${}^{18}O_2$, the amount of CO desorbed from the surface decreases and the temperature of maximum desorption of the β_1 state is again shifted to lower temperatures until finally it is almost impossible to adsorb any CO on the oxygen covered surface. Figure 20 also illustrates the desorption of $C^{18}O$ in these experiments. It is observed that as the ${}^{18}O_2$ dosed increases the amount of $C^{18}O$ desorbed first increases (curves A - C of Fig. 20), then decreases (curves C-D) because the amount of CO on the surface decreases. There is almost no $C^{18}O$ desorbed from the



Fig. 20. Flash desorption spectra of CO form a W(110) surface predosed with increasing dosages of ¹⁸O₂ (A - 1.2L, B - 2.6L, C - 5.3L, D - 10.7L, and E - 21.3L) followed by a constant dose of C¹⁶O (9.2L).

surface as α -CO.

The total amounts of CO desorbed from the W(110) surface in each of the experiments as determined from the area under flash desorption curves are plotted in Fig. 21. A constant amount of CO is desorbed from the surface after being predosed with a constant amount of CO followed by increasing coverages of ${}^{18}O_2$; this proves that the oxygen is not able to displace any of the CO adsorbed on the surface at room temperature.

The ratios of $C^{18}O:CO_{total}$ desorbed from the surface in each of the experiments (Figs. 16-19) are plotted in Fig. 22. In each experiment, there appears to be a limiting ratio of $C^{18}O:CO_{total}$ desorbed. The limiting ratios for each of the four sets of experiments are plotted in Fig. 23. At low coverages of preadsorbed CO followed by a saturated dose of ${}^{18}O_2$, the limiting ratio is ~.5 and for a surface saturated with preadsorbed CO followed by a saturated dose of ${}^{18}O_2$, the limiting ratio is ~.25.

The above results will be summarized before proposing a consistent model.

(1) CO adsorbed on a W(110) surface at room temperature desorbs as an α -state at ~400°K and as two β states, β_1 at ~920°K and β_2 at ~1100°K. The separation into two states appears to result from interactions of co-adsorbed species on the surface.



Fig. 21. Total amount of CO (C¹⁶O + C¹⁸O) desorbed from a W(110) surface where the surface was predosed with C¹⁶O (A - 2.5L, B - 4.4L, C - 9.2L and D - 18.6L) with increasing dosages of ¹⁸O₂ (1.2L, 2.6L, 5.3L, 10.7L and 21.3L) in each set of C¹⁶O doses.



Fig. 22. The ratio of C¹⁸O/CO_{total} desorbed from a W(110) surface predosed with C¹⁶O (A - 2.5L, B - 4.4L, C - 9.2L and D - 18.6L) with increased dosages of ¹⁸O₂ (1.1L, 2.4L, 5.3L, 10.6L and 21.2L) in each set of C¹⁶O doses.

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Fig. 23. The limiting ratio of $C_{0,0}^{18}O/CO_{0,0}^{18}O$ desorbed from a W(110) surface predosed with $C_{0,0}^{16}O$ (2.5L, 4.4L, 9.2L and 18.6L) followed by a saturated dose of $1_{0,0}^{18}O_{2}$.

(2) Preadsorbed CO was not displaced from the surface by dosing with ${}^{18}O_2$ nor was there any extensive increase in the amount of CO desorbed in the α state. The lack of such an observation suggests no conversion is occurring between adsorbed states.

(3) Mixed adsorption of CO and ${}^{18}O_2$ results in isotopic mixing of the oxygen in the CO molecules desorbed from the β states and very little mixing in the CO molecules desorbed in the α state.

(4) From a low coverage of CO followed with a saturated coverage of ${}^{18}O_2$, the limiting ratio of $C^{18}O:CO_{total}$ desorbed from the surface is ~.5 and from a saturated coverage of CO and ${}^{18}O_2$, the limiting ratio is ~.25.

(5) A surface saturated with preadsorbed ${}^{18}O_2$ first preferentially blocks CO adsorption in the β state and slightly increases the amount of CO desorbed as α -CO (Fig. 20). As the amount of preadsorbed ${}^{18}O_2$ was increased, the amount of CO adsorbed in all states decreased.

The above results suggest a model for adsorption of CO on the W(110) surface similar to that for the W(100) surface. The CO which desorbs in the β states, where there is extensive isotopic mixing of the oxygen when it is co-adsorbed with $^{18}O_2$, will be assumed to be in the form of C and O adatoms. The CO desorbed as the α state is proposed to be associatively adsorbed CO. In the flash desorption of

the adsorbed CO, a point of concern is the conversion of α -CO to the β -CO by the heating process. If the CO were preadsorbed associatively on the surface, the co-adsorbed ¹⁸O₂ should be able to block the sites for the conversion of α -CO to β -CO. Since increasing doses of ¹⁸O₂ on a surface with preadsorbed CO leads to no increase in the amount of CO desorbing in the α state, it appears that there is no conversion from an associatively adsorbed CO in an α state to a β state.

The above model should also be consistent with other reported results for the adsorption of CO on the W(110) surface. Kohrt and Gomer (42) reported the formation of a virgin state, when dosing a W(110) surface at a temperature of 100°K or lower with CO. In a step desorption experiment, virgin CO desorbed at 200-450°K, followed by the desorption of two beta states, β_1 at ~970°K and β_2 at ~1120°K. Because of the complex kinetics observed from the desorption of the virgin state, they suggested there was competition between desorption and conversion to β state, with the latter predominating at low θ . If after forming the virgin state by dosing a surface at 100°K or less, they heated the surface to only 450-600°K where all the virgin state had desorbed, they were then able to readsorb CO forming a new state. This state was different from the original low temperature virgin state and was labeled α . It desorbed as α_1 at ~150°K and α_2 at ~300°K. This also supports the hypothesis that the CO adsorbed in

the virgin state is converted irreversibly to the β states.

In view of the model proposed for dissociative adsorption of CO, it must be reasoned that the W(110) surface at ~ 100°K is not able to dissociatively adsorb CO. Therefore, it adsorbs the CO associatively as virgin CO, but when the surface is heated to 450-600°K where half of the virgin CO desorbs, the remaining CO dissociates to C and O adatoms. Tungsten with this chemisorbed layer is able to adsorb additional CO associatively as the α states observed by Kohrt and Gomer (42). In this study, the CO was dosed on a W(110) surface at room temperature, and as observed the β states formed immediately and were not formed by the conversion of associative adsorbed CO molecules. With larger doses of CO, as the β states were saturated, a small amount of CO adsorbed associatively (no isotopic mixing) as an α -CO. This compares with the α_2 -CO state desorbed at ~ 300°K observed by Kohrt and Gomer.

On the W(110) surface there are three symmetric sites available for the chemisorbed species. These sites are indicated in Fig. 24. At site A, a chemisorbed species could possibly be coordinated to four surface atoms. Although two of the surface atoms are farther away, the two nearer surface atoms are separated by the same distance as those forming a 2 CN site on the W(100) surface. Although the two tungsten atoms are touching at site B, a chemisorbed


Fig. 24. Possible sites occupied by chemisorbed species from coadsorption of CO and ¹⁸O₂ on a W(110) surface (A - 2 CN site, B - 2 CN' site, C - 1 CN site, ● - C or O adatoms and ● - associatively adsorbed oxygen). species could be coordinated to these surface atoms. At site C, a chemisorbed atom would be coordinated to a single surface atom.

May and Germer (41) reported a C(9x5) pattern from a W(110) surface with CO adsorbed in the β state. Such patterns are of very little help in assigning possible sites and stoichiometry for the adsorbed species. In a LEED study of oxygen on the W(110) surface, Tracy and Blakely (21) reported that atomically chemisorbed oxygen produced a p(2x1) pattern. This structure corresponds to one-half a monolayer of oxygen atoms. The p(2x1) structure can be produced by the O adatoms occupying every other row of sites A as marked in Fig. 24. With the O adatoms in this surface structure, the bonding shows great similarity to the bonding of O adatoms at 2 CN sites on the W(100) surface where it produced a (2x1) LEED pattern. In the (2x1) structure on W(100), each O adatom was bonded to two surface tungsten atoms, and these tungsten atoms were not bonded to any other O adatoms. If the O adatoms on a W(110) surface occupied the proposed sites indicated in Fig. 24, then each oxygen atom would also be bonded to two of the surface tungsten atoms, and these two tungsten atoms would not be directly involved in the bonding of O adatoms at adjacent sites. At full oxygen coverage, each tungsten atom would be bonded to a single oxygen atom which is consistent with the bonding model also proposed for the W(100) surface. The carbon atom should

seek a site of the highest coordination number possible to satisfy its multi-valency. On the W(110), this would be site A (Fig. 24). Therefore, it is very likely that the C adatoms would occupy the same sites as the O adatoms.

In dissociative adsorption of CO on the W(110) surface, the C and O adatoms should occupy every other row of sites A (Fig. 24). In order to observe C¹⁶O and C¹⁸O as desorbed CO molecules, it must be possible to adsorb $^{18}O_2$ upon a W(110) surface with preadsorbed $C^{16}O_2$. And as indicated by the plots in Fig. 21, the adsorbed $C^{16}O$ is not displaced by the adsorption of ${}^{18}O_2$. Therefore, the ${}^{18}O_2$ must be co-adsorbed with the preadsorbed CO. As was concluded for the W(100) surface, it will be assumed that the ${}^{18}O_{2}$ will be associatively adsorbed. The adsorbed 18_{O_2} could be bridged across site A bonding to the two distant tungsten atoms. The adsorbed oxygen could be occupying every other site A between the rows of C and O adatoms. The sum of the amount of CO desorbed in the β_1 and β_2 states is a constant in a set of experiments where the same predose of $C^{16}O$ is followed by increased ${}^{18}O_2$ coverages. The shift in the desorption of the β -CO all to β_1 -CO (lower temperature state) is not likely to result from site changes of C and O adatoms, but rather from weakened bonds between surface atoms of tungsten and the C and O adatoms due to the co-adsorbed ${}^{18}O_2$. At the temperatures where

the β_1 -CO desorbs, the associatively co-adsorbed ${}^{18}O_2$ will be dissociating and the O adatoms will want to occupy sites A between the rows of C and O adatoms. Therefore, every tungsten atom would be coordinated to two adatoms, a structure certainly not as stable as the previous structure where every tungsten atom was coordinated to only one adatom.

In Fig. 15 the presences of two β states in flash desorption spectra were attributed to co-adsorbed ${}^{16}O_2$ which was noticed to be varying in the ambient. In fact it is very likely that the β_1 state exists only as a result of some co-adsorbed species (not necessarily always oxygen). The β_1 state could desorb from the surface where the C and O adatoms are co-adsorbed with ${}^{18}O_2$ according to the above bonding scheme where co-adsorbed species are trying to occupy all of the A sites available (Fig. 24). The β_2 state could be assumed to result from the surface were only C and O adatoms are bonded in the p(2x1) structure. The co-adsorption could also account for the C(9x5) LEED pattern reported by May and Germer (41), because May, Germer and Chang (43) later reported a C(11x5) LEED pattern resulting from experiments of known co-adsorption of oxygen and carbon monoxide on the W(110) surface.

When the surface was predosed with increasing amounts of $^{18}O_2$, followed with constant doses of $C^{16}O$, the amount of $C^{16}O$

adsorbed on the surface decreases. In Fig. 20, it is observed that a surface with preadsorbed oxygen has a larger amount of CO desorbing in the α state than in any experiment (Figs. 15-18) where the surface was predosed with $C^{16}O$. It should also be noted, even in the ${}^{18}O_2$ predosed surfaces followed with a dose of $C^{16}O$, that there is no $C^{18}O$ desorbed in the α state. The CO desorbed in the β states is desorbed primarily as β_1 with isotopic exchange of the oxygen occurring. The increase in the amount of α -CO desorbed from the surface where the CO was dosed on a surface with preadsorbed ${}^{18}O_2$ occurred as a result of the O adatoms blocking sites for dissociation by the depletion of adjacent empty A sites, but allowed single A sites to exist for associatively adsorbed CO.

At very small coverages of preadsorbed $C^{16}O$, it should be possible to complete the p(2x1) structure with ¹⁸O adatoms. In the limiting case of small coverages, the C and ¹⁶O adatoms from the dissociated CO molecule should occupy adjacent sites in the p(2x1) structure, and remaining sites in this structure should be occupied by ¹⁸O adatoms. As the carbon atoms desorb, it then has a choice of two O adatoms, one ¹⁸O and the other ¹⁶O. Therefore, the limiting ratio of $C^{18}O:CO_{total}$ desorbed from a surface in the case of a low coverage of CO and saturated with ¹⁸O₂ should be .5, which compares remarkably well with the value determined from the plot in Fig. 23.

When CO adsorbs on the surface and dissociates into C and O adatoms, it is necessary to have two adjacent sites vacant. On the W(110) surface, the C and O adatoms were assumed to occupy every other row of A sites (Fig. 24). Random dissociative adsorption of CO according to the above hypothesis results in $\sim 25\%$ of the A sites on the W(110) surface not having empty adjacent sites (44). The α -CO (associatively adsorbed CO) will be assumed to be bonded at these sites. A p(2x1) surface structure was assumed to result from C and O adatoms from dissociatively adsorbed CO. One-half a molecule would be adsorbed per unit cell or one molecule in a p(2x2) structure. If 18 O, were dosed on a surface of saturated preadsorbed CO, the co-adsorbed ${}^{18}O_2$ could occupy every other site in the rows between the C and O adatoms (Fig. 24); these oxygen molecules would also produce a $p(2x^2)$ structure or one molecule per unit cell of this structure. Upon heating the surface in a flash desorption experiment, the dissociation of adsorbed molecular $^{18}O_2$ occurs according to

$$\begin{array}{c} 0 - 0 = 20 \\ / \\ * \\ \end{array}$$

It will be assumed a C adatom can recombine only with an O adatom for CO desorption. According to the above bonding scheme, there is a C and ¹⁸O adatom per p(2x2) unit mesh and the associatively co-adsorbed ¹⁸O₂ also formed a p(2x2) unit mesh. Therefore, each

p(2x2) unit mesh has C and ¹⁶O adatoms and a molecule of ¹⁸O₂. The probability of an adsorbed ¹⁸O₂ is near the C adatom is .5 and coupled with the assumption only one-half of the ¹⁸O₂ molecules have dissociated at the temperature where CO desorbs results in the C adatom having a choice of oxygen isotopes in the ratio ¹⁶O:¹⁸O of 1:.5. The resulting ratio of $C^{16}O: C^{18}O$ desorbed from the β state should be 2: 1. Therefore, from a surface with a saturated predose of $C^{16}O$ followed by a saturating ¹⁸O₂ dose, 25% of the CO will desorb as associatively adsorbed $C^{16}O$. From 100 sites, the remaining 75 sites should desorb 37.5 molecules of CO as the β state. From the above ratio one-third should be $C^{18}O$ or the limiting ratio β of $C^{18}O:CO_{total}$ would be .20. This ratio estimated from Fig. 22 is ~.24.

The above discussion can be summarized very briefly. The isotopic mixing of the oxygen in the desorbing CO molecules can best be interpreted in terms of a model for dissociative adsorption of CO. If the adsorbed CO is in the form of C and O adatoms, it is very understandable why adsorption of oxygen does not displace any preadsorbed CO. The bonding of the C and O adatoms at sites A in Fig. 24 to produce a p(2x1) structure explains the limiting ratio of $C^{18}O: CO_{total}$ desorbed at low CO coverages and is consistent with bonding models for chemisorbed species presented in Chapter II.

Mixed adsorption of CO and $^{18}O_2$ on a W(111) surface

The flash desorption spectra of CO adsorbed on a W(111) surface are plotted in Fig. 25. The CO desorbs in two states, a β state at ~1050°K and an α state at ~450°K. These two states desorb as very sharp symmetric peaks, an indication of the absence of any other states present on the surface. The desorption of CO is unlike the hydrogen desorption from a W(111) surface in which Tamm and Schmidt (2) observed multiple desorption peaks. In the case of CO adsorption, it is not even possible to form any new desorption peaks by the coadsorption of oxygen on a surface with preadsorbed CO. The results of the mixed adsorption of $C^{16}O$ and ${}^{18}O_2$ on the W(111) surface are plotted in Figs. 26-29. In Fig. 26, it is observed, for the small predose of $C^{16}O$ followed by increasing coverages of $^{18}O_2$, that the amount of C¹⁸O desorbed increases proportionally. At high coverages of ${}^{18}O_2$, the amount of $C^{18}O$ desorbed is very large compared to $C^{16}O$ desorbed in the high temperature state, but the amount of $C^{18}O$ desorbed in this state decreases as the amount of CO predosed on the surface increases (Figs. 26-29). It is also interesting to note in this series of experiments that the temperature of maximum desorption of this high temperature state decreases as the coverage of oxygen increases. The isotopic mixing of the oxygen in the desorbing CO molecules from a surface with co-adsorbed $C^{16}O$ and $18O_2$ occurs



Fig. 25. Flash desorption spectra of CO adsorbed on a W(111) surface (A - 2.3L, B - 4.8L, C - 8.0L and D - 19.9L).



Fig. 26. Flash desorption spectra of CO from a W(111) surface predosed with 2.3L of C¹⁶O followed with increasing dosages of ¹⁸O₂ (A - 0.0L, B - 1.4L, C - 2.5L, D - 5.7L, E - 10.7L and F - 22.0L).



Fig. 27. Flash desorption spectra of CO from a W(111) surface predosed with 4.8L of C¹⁶O followed with increasing dosages of ¹⁸O₂ (A - 0.0L, B - 1.4L, C - 2.5L, D - 5.7L, E - 10.7L and F - 22.0L).

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Fig. 28. Flash desorption spectra of CO from a W(111) surface predosed with 8.0L of C¹⁶O followed with increasing dosages of ¹⁸O₂ (A - 0.0L, B - 2.5L, C - 5.7L, D - 10.7L and E - 22.0L).





Fig. 29. Flash desorption spectra of CO from a W(111) surface predosed with 19.9L of C¹⁶O followed with increasing dosages of ¹⁸O₂ (A - 0.0L, B - 2.5L, C - 5.7L, D - 10.7L and E - 22.0L).

almost exclusively in the CO desorbed in the β state. When oxygen is dosed on a surface with preadsorbed CO, there is no significant increase in the amount of α -CO desorbing. Such an observation suggests no conversion process of α -CO and β -CO is occurring on the surface. If β -CO was formed by conversion of the associatively adsorbed α -CO, the oxygen adsorption should prevent this conversion. Therefore, an increase in the amount of α -CO desorbed would result. No such observations were made (Figs. 26-29) providing no evidence for a conversion process.

In Fig. 30, it is observed that at low coverages of preadsorbed oxygen there is a decrease in the overall amount of CO adsorbed on the surface compared to a clean surface. The preadsorbed oxygen appears to initially block sites for CO adsorption in the β state and create more sites for CO to adsorb as α -CO (Fig. 30). But as the amount of preadsorbed oxygen increases, there is a decrease in the amount of CO desorbed in both states.

As observed on the other tungsten surfaces in this study, oxygen when dosed on a surface with preadsorbed CO is not able to displace any of the CO. Figure 31 shows, that with increasing doses of oxygen on surfaces with constant coverages of preadsorbed CO, the total amount of CO ($C^{16}O + C^{18}O$) desorbed is constant but the ratio of $C^{18}O:CO_{total}$ which is desorbed from the surface (Fig. 32) changes.



Fig. 30. Flash desorption spectra of CO from a W(111) surface predosed with increasing dosages of ¹⁸O₂ (A - 2.5L, B - 5.7L, C -10.7L and D - 22.0L) followed by a constant dose of C¹⁶O (8.0L).



Fig. 31. Total amount of CO (C¹⁶O + C¹⁸O) desorbed from a W(111) surface where the surface was predosed with C¹⁶O (A - 2.3L, B - 4.8L, C - 8.0L and D - 19.9L) with increasing dosages of 180₂ (1.4L, 2.5L, 5.7L, 10.7L and 22.0L) in each set of C¹⁶O doses.



Fig. 32. The ratio of C¹⁸O/CO_{total} desorbed from a W(111) surface predosed with C¹⁶O (A - 2.3L, B - 4.8L, C - 8.0L and D - 19.9L) with increased dosages of ¹⁸O₂ (1.4L, 2.5L, 5.7L, 10.7L and 22.0L) in each set of C¹⁶O doses.

This ratio appears to reach limiting values for a selected predose of CO followed by a saturating ${}^{18}O_2$ dose. These limiting ratios of $C^{18}O:CO_{total}$ which desorb from a surface with preadsorbed CO are (Fig. 33) 0.65 at low coverages of CO and a surface saturated with ${}^{18}O_2$, and 0.24 for a saturated coverage of CO followed by a saturating ${}^{18}O_2$ dose.

Following is a summary of the above results which must be consistent with any proposed model for CO adsorption and interaction with co-adsorbed $^{18}O_2$.

(1) CO adsorbed on a W(111) surface at room temperature desorbs as an α state at ~450°K and a β state at ~1050°K.

(2) Co-adsorption of CO and ${}^{18}O_2$ result in isotopic mixing of the oxygen in the CO molecules desorbed from the β state.

(3) Preadsorbed CO is not displaced from the surface by dosing with oxygen nor is the oxygen able to create new desorption states. The adsorbed oxygen on a surface with preadsorbed CO does not increase the amount of CO desorbing in the α state providing evidence to dispute the existence of a conversion process to the β state by heating the surface.

(4) A surface with preadsorbed ${}^{18}O_2$ initially blocks sites for adsorption in the β state and creates more sites for adsorption of α -CO. A surface with a saturated adlayer of oxygen appears to be



Fig. 33. The limiting ratio of C¹⁶O/CO_{total} desorbed from a W(111) surface predosed with C¹⁶O (2.3L, 4.8L, 8.0L, 19.9L) followed by a saturated dose of ¹⁸O₂.

able to block subsequent adsorption of CO in any state.

(5) There are limiting ratios of $C^{18}O:CO_{total}$ desorbed from a surface with preadsorbed CO followed by a saturating ${}^{18}O_2$ dose. At low coverage of preadsorbed CO, this ratio appears to be ~.65 and from a surface with a saturated coverage of preadsorbed CO, this ratio appears to be ~.24.

The above results are very similar to those found on the other tungsten surfaces studied and therefore, also imply a model for the dissociative adsorption of CO. The CO in the β state on the W(111) is assumed to be dissociatively adsorbed based on the isotopic mixing of the oxygen in the desorbed CO molecules from a surface of coadsorbed C¹⁶O and ¹⁸O₂. The CO molecules in the α state must be associatively adsorbed because of the lack of any significant isotopic mixing. The decrease in the temperature of maximum desorption of the CO in the β state as the coverage of adsorbed oxygen increases can either result from weakened bonds with the surface due to the coadsorbed oxygen or from the increased O adatom concentration enabling easier recombination of the C adatom with oxygen.

On the W(111) surface, three symmetric sites seem attractive for bonding chemisorbed species to the surface. In Fig. 34, these sites are illustrated as site A which has three-fold symmetry and is above



Fig. 34. Possible sites occupied by chemisorbed species from coadsorption of CO and ¹⁸O₂ on a W(111) surface (A - 4 CN site, B - 4 CN' site, C - 1 CN site, ● - Oadatoms, ⊗ -C adatoms and ● - associatively adsorbed oxygen).

an atom exposed to the surface and also surrounded by other surface atoms. Site B also has three-fold symmetry and is above a surface atom which is more exposed as a surface atom. The C sites are above the most exposed surface atoms on this surface. Based on the results and prediction in Chapter II, a carbon adatom would probably select site A to satisfy its multi-valent property. The proximity and the easy occupation of site B during dissociative adsorption of CO probably would result in O adatoms selecting these sites. If dissociative adsorption requires an empty site A and an empty adjacent site B, it is very probable, as was found and predicted on the other tungsten surfaces (18, 44), that in random dissociative adsorption $\sim 20\%$ of the A sites would not have available empty adjacent B sites at saturation coverage. These sites could then be occupied by associatively adsorbed CO which would be stable on the surface at room temperature. This is very similar to the model proposed on the other tungsten surfaces.

We therefore assume that adsorption of CO at room temperature results in C and O adatoms bonded at adjacent A and B sites (Fig. 34), which desorb as CO molecules in the β state, and CO adsorbed associatively at A sites where adjacent B sites are unavailable, which desorb as CO molecules in the α state. On a surface with preadsorbed CO, it is easy to understand why the co-adsorption of oxygen was not able to displace any of the preadsorbed CO from these bonding sites.

It was observed that preadsorbed oxygen initially blocked sites for β adsorption while creating more sites for associatively adsorbed CO (Fig. 30). If the preadsorbed oxygen is dissociatively adsorbed at the B sites, this increases the probability of A sites not having an empty adjacent B site for CO to dissociatively adsorb. Therefore, there is an increase in A sites for the associatively adsorbed CO (observed in Fig. 30).

At very low coverages of CO followed by a saturated dose of ${}^{18}O_2$, the C adatom bonded at A sites would find one ${}^{16}O$ adatom in an adjacent B site and two ${}^{18}O$ species adsorbed at the other adjacent B sites. Therefore, the limiting ratio of $C^{18}O:CO_{total}$ desorbed should be ~. 67 which compares favorably with the value of ~. 65 determined from the plot in Fig. 33. On a W(111) surface saturated with preadsorbed $C^{16}O$ and followed with a saturating ${}^{18}O_2$ dose, the only place for the co-adsorbed ${}^{18}O_2$ molecules to bond are at the remaining unoccupied C sites. It is very likely that the oxygen molecule would be bridged between two adjacent C sites. In this configuration, there is one-half an adsorbed ${}^{18}O_2$ molecule per C adatom. Again, it will be assumed that upon heating the surface during flash desorption, the oxygen dissociates according to

$$\frac{1}{2} \circ - \circ \neq \circ$$

Based upon previous evidence, ~20% of the CO desorbed will be α -C¹⁶O from associatively adsorbed CO at A sites. If the C adatom will recombine only with oxygen atoms, it has a choice of one ¹⁶O adatom from the dissociated CO molecule and assuming the ¹⁸O₂ dissociation is only one-half complete, and that only one-half the C adatoms are adjacent to the adsorbed ¹⁸O₂, then the C adatom has a choice of one-fourth ¹⁸O adatom for every ¹⁶O adatom. Therefore, the 80% of the CO desorbed from the β state will be one-fifth C¹⁸O. The limiting ratio of C¹⁸O:CO_{total} desorbed from the above surface saturated with preadsorbed CO followed by a saturating dose of ¹⁸O₂ is ~.18 and compares with the value estimated in Fig. 33 of ~.24.

Summary and Conclusions

A model for dissociative adsorption of CO is consistent with all the observations in this study where CO chemisorption was examined on three single crystal surfaces (100, 110, 111) of tungsten. The observations indicated the CO which desorbed at the high temperatures was dissociated on the surface at room temperature. On all the surfaces, a stable associatively adsorbed species was found to desorb at lower temperatures. All these chemisorbed species were not displaced by co-adsorbed oxygen. The CO desorbing at the higher temperature was a mixture of $C^{16}O$ and $C^{18}O$ which resulted from the isotopic mixing of the oxygen from the dissociative adsorbed CO and the co-adsorbed

 18 O₂ which was at least partially dissociated before the CO desorbed. On all three tungsten surfaces, it was assumed the associatively adsorbed CO was bonded at sites where vacant sites needed for the dissociation of the CO molecule were not present. On the W(100) and W(111) surfaces, it was proposed that adsorbed oxygen from an oxygen dose on a clean surface would not occupy the sites preferred for C adatoms from CO dissociation. Therefore, on the W(100) surface preadsorbed oxygen at 2 CN sites(Fig. 13) should decrease the availability of vacant adjacent 5 CN sites needed for CO to adsorb and dissociate. As observed (Fig. 9), this results in an increase of α -CO desorbed from the W(100) surface where CO was dosed on a surface with preadsorbed oxygen. On the W(111) surface, it was assumed the preadsorbed oxygen occupied sites B in Fig. 34. When the CO adsorbs at sites A, vacant adjacent sites B are needed for the CO to dissociate. If these sites are occupied by preadsorbed O adatoms, then they are not available for CO dissociation. Therefore, as observed in Fig. 30, there is an increase in the amount of α - CO desorbed from the W(111) surface where CO was dosed on a surface with preadsorbed oxygen. On the W(110) surface, the oxygen adatoms from adsorbed oxygen and the adatoms from dissociated CO were assumed to occupy the same sites. Therefore, the random adsorption of either CO or O_2 on a W(110) surface should result in approximately the same number of sites present on the surface with unavailable vacant adjacent sites. These

sites were assumed to be occupied by associatively adsorbed CO. Therefore, in contrast to the other surfaces (100, 111), the amount of α - CO desorbed from a W(110) surface where CO was dosed on a surface with preadsorbed oxygen (Fig. 20) is about the same as the amount of α - CO desorbed from a surface where only CO was adsorbed (Fig. 14).

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A MOLECULAR ORBITAL INVESTIGATION OF

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CHEMISORPTION. I. HYDROGEN ON TUNGSTEN (100) SURFACE

A Molecular Orbital Investigation of

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Chemisorption. I. Hydrogen on Tungsten (100) Surface

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Abstract

The relative bonding energies of hydrogen chemisorbed at three symmetric sites on a W(100) surface were obtained by means of the Extended Hückel Molecular Orbital Theory (EHMO). The preferred site for hydrogen chemisorption was found to be the single coordination number (1 CN) site or the site above a surface tungsten atom. The W(100) surface was represented by finite arrays of tungsten atoms which were shown to be adequate for obtaining semi-quantitative results. The basis set for the calculations contained the valence orbitals of tungsten and, initially, the 5p orbitals which were nonbonding but provided the necessary repulsion at small internuclear separation. The repulsive energy provided by these orbitals was replaced by an analytical exponential repulsive energy term. This allowed the 5p orbitals to be omitted from the basis set to simplify computation. Functionally, the energy change for the reaction $W_n + H \rightarrow W_n H$ was calculated for various assumed configurations of the $W_n H$ "molecule". The bonding between tungsten atoms was found to be changed as a result of $W_n H$ formation, and the change varied with hydrogen position. Energy barriers to surface diffusion were also calculated, and found to agree reasonably with experimental values.

I. INTRODUCTION

In recent years, it has become experimentally possible to study the chemisorption of atoms and molecules on single crystal surfaces. To account for the experimental observations from these well defined surfaces, various bonding models have been proposed, but the detailed description of the adsorbed species remains in the realm of speculation. Unfortunately, no available experimental techniques give conclusive information about the molecular structures or site distributions of the adsorbed species. In view of current experimental limitations and the importance of the implications in the field of catalysis, a theoretical attack seems justified. The prohibitive expense of

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ab initio quantum mechanical calculations for such systems make it attractive to explore nonrigorous, semiempirical methods which have been found in the past to account quite well for structural features of molecular systems.

In this paper, we apply the extended Huckel molecular orbital theory, a semiempirical molecular orbital technique, to investigate the preferred structure of the "surface molecule". EHMO has been used previously to study the chemisorption of hydrogen on graphite, ¹ and of organic molecules, 2 carbon monoxide³ and hydrogen⁴ on nickel. In the case of carbon monoxide on nickel,³ the dependence of binding energy on distance at the possible chemisorption sites was not analyzed. The preferred site was selected by comparison of binding energies at a fixed carbon-nickel distance. These investigations used only the valence orbitals in the basis set, and as a result only the hydrogengraphite interaction produced a minimum in the binding energy curve. The interaction of hydrogen as well as carbon, nitrogen, oxygen and fluorine on graphite⁵ was also investigated using CNDO technique in order to alleviate the problem of unreasonable charge transfer which. can occur in the EHMO calculations.

Although it is known that this technique has many limitations, it nevertheless has been successful in determining preferred molecular configurations. Therefore, it is a promising tool for

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investigating the structure of the elusive surface molecules and for obtaining a better understanding of the bonding between chemisorbed atoms and the surface atoms. We have chosen to investigate the chemisorption of hydrogen on a tungsten (100) surface because of the experimental data which are available for comparison. $^{6-11}$

II. PROCEDURE

Hoffmann¹² developed a semiempirical molecular orbital theory, including all overlap integrals, referred to as the "extended Hückel" Molecular Orbital theory (EHMO). It was originally utilized to solve
organic conformational problems, but subsequently has been used to yield insight on other molecular geometric problems, including the structural chemistry of crystals.¹³ Hoffmann stressed that the method's most reliable characteristic is its ability to predict molecular geometries and to assess the relative importance of sigma and pi orbitals. On the other hand, it is quite crude in predicting charge distributions and excitation energies in electronic transitions.

The molecular orbitals are formed from a linear combination of atomic orbitals (LCAO),

$$\Psi_j = \sum_q C_{jq} \chi_q$$
,

where the atomic orbitals χ_q are the single parameter, Slater-type atomic orbitals (STO). According to the standard variation principle, the LCAO function with q atomic orbitals leads to a set of q simultaneous linear homogeneous equations, each of which contains q coefficients C_{iq} ,

$$\sum_{\mathbf{q}} (\mathbf{H}_{\mathbf{pq}} - \epsilon_{j} S_{\mathbf{pq}}) C_{j\mathbf{q}} = 0 \quad .$$

For these equations to have nontrivial solutions, the determinant of the coefficients must vanish,

$$||H_{pq} - \epsilon_j S_{pq}|| = 0$$

The eigenvalues ϵ_j given by the solution of this determinant are the energies of the molecular orbitals. These energies ϵ_j are substituted back into the set of simultaneous linear homogeneous equations which are

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then solved to give the coefficients C_{jq} . These coefficients C_{jq} form the wave function of the molecular orbital at each energy ϵ_j . In the Hoffmann formalism, the diagonal elements H_{pp} are set equal to the valence orbital ionization potential (VOIP) of χ_p ; the non-diagonal elements H_{pq} were approximated by the Mulliken Wolfsberg-Helmholtz approximation¹⁴

$$H_{pq} = 0.5K(H_{pp} + H_{qq})S_{pq}$$

with a Helmholtz Factor K equal to 1.80.

The EHMO method does not explicitly account for either electronelectron or core-core interactions between atoms, but relies on extensive cancellations which hold approximately for nonpolar molecules. These cancellations will also hold for other molecules near the equilibrium internuclear separations where the chemical bond is formed. Excessive charge transfers, a common problem in the EHMO method, were partially alleviated by adjusting the VOIP as functions of atomic charge q and iterating to self-consistency. The atomic charge was computed by the Mulliken method¹⁵ and the VOIP of the atomic orbitals were determined as functions of charge q according to the following¹⁶

 $VOIP(q) = Aq^2 + Bq + C$.

These adjusted VOIP gave more realistic charge transfers than fixed VOIP. Bonding energies for different structures in the neighborhood of their energy minima were similarly shifted by shifts in VOIP so

that conclusions as to preferred sites were unaffected by variations in VOIP.

Table I lists the orbital exponents for the STO and the values for the constants, A, B and C, which are needed to determine the VOIP as a function of atomic charge. The values for the hydrogen 1s orbital were obtained from Basch, Viste and Gray.¹⁶ A value of 1.20 was used for the orbital exponent of the hydrogen 1s orbital. For the tungsten 5p orbital, the value of C was obtained from the tables of Herman and Skillman;¹⁷ the orbital exponent for the STO was obtained by the rules of Burns¹⁸ and assumed to be charge independent. The values of C and the orbital exponents for the STO of the tungsten 5d and 6s orbital were obtained from Lohr and Lipscomb;¹⁹ the values of C for these orbitals agreed with neutral atom ionization measurements of Lotz.²⁰ A linear charge dependence²¹ for the VOIP of the tungsten 5d and 6s orbitals was used with the value of B equal to 3.5 ev/e⁻.

The EHMO method was first applied to the diatomic molecules, W-W and W-H. These calculations showed that it was necessary to include the inner orbitals of the tungsten atoms as well as the valence orbitals in the basis set to obtain a characteristic potential energy curve; when only valence orbitals were used too few non-bonding orbitals were populated to provide the requisite repulsion at small internuclear separations. It was found that the addition of the inner

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orbitals overcame this deficiency, although they made little contribution to the formation of the chemical bond. Therefore, if the repulsive energy could be calculated by an empirical relationship, these orbitals could be omitted, thus reducing the size of the basis set and saving a substantial computation effort. The repulsive energy between atoms in a molecule can be represented approximately by a series of exponential terms of the form

$$E_{\text{Rep}} = \sum_{ij} A_{ij} e^{-r_{ij}/\alpha_{ij}}$$

where the parameters, A_{ij} and α_{ij} , are unique for a pair of atoms. These parameters for pairwise interactions were derived from the repulsive energy, E_{Rep} , of the diatomic molecules. The repulsive energy was determined as the difference in bonding energies obtained by a calculation with only valence orbitals and by a calculation which also included the inner orbitals. Comparison of E_{Rep} for WH and W_2 H indicated that repulsive energies in the tungsten hydrogen system could be assumed pairwise additive.

In modeling the tungsten (100) surface for the EHMO calculations, it was necessary to truncate the infinite surface to a limited number of metal atoms. These selected surface arrays must represent the possible sites for bonding of a chemisorbed atom. The tungsten (100) appears to offer three distinct symmetric sites which might be considered attractive adsorption sites. These are shown in Figure 1, which also

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shows the metal atom arrays used in calculations. The three sites can be distinguished by the numbers of surface tungsten atoms to which the hydrogen atom is coordinated (CN). Thus the 1 CN site is on top of a tungsten atom, the 2 CN site bridges two tungsten atoms, and the 5 CN site is in the four-fold hole. The size of the surface array representing the 1 CN and 5 CN sites was increased to 21 tungsten atoms with an insignificant change in bonding energies. The presence of edges on these truncated surface arrays caused abnormal charge distribution, but these effects were reduced by adjusting the VOIP of the edge atoms. After this initial adjustment, there was no further adjustment of the tungsten orbitals' VOIP.

III. RESULTS

A. Diatomic Molecules

Energy-distance calculations were made for diatomic molecules to determine the parameters needed for the exponential repulsive energy term. These calculations also provided a basis for comparison with the chemisorbed system, in which the hydrogen was bonded to larger arrays of tungsten atoms. In Fig. 2, the bonding energy is plotted as a function of inter-nuclear separation r along the molecular axis z for the W-W and W-H molecules. The bonding energy was calculated by subtracting from the total energy the energy of the ions

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of the W-H molecule or the energy of the atoms of the W-W molecule at infinite separation. These arbitrary conventions compensate in part for the total neglect of coulomb interactions in the EHMO method and have no influence on any conclusions drawn later.

Figures 2a and 2b display the clear need to include some of the inner orbitals of tungsten in the basis set in order to obtain a characteristic potential energy curve. The differences between curves a and b in Figs. 2a and 2b are the repulsive energies. In Fig. 3, $\ln \Delta \epsilon$ is plotted as a function of internuclear separation r for W-W and W-H molecules. From these plots, the values for the parameters A and α needed to calculate the repulsive energy by the analytical exponential term were derived. Best values were found to be: $A_{WW} =$ 4096.ev, $\alpha_{WW} = 0.289 \text{\AA}^{-1}$, $A_{WH} = 139.ev$, $\alpha_{WH} = 0.279 \text{\AA}^{-1}$.

As previously stated, the VOIP of the valence orbitals were adjusted as a function of atomic charge and iterated to self-consistency. For the diatomic molecule W_2 the atomic charge on each tungsten atom was of course zero, for WH the charge on hydrogen was -0.10 and on tungsten +0.10. Corresponding VOIP are implied by Table I. The molecular orbitals (LCAO coefficients) for the diatomic molecules at equilibrium separation are tabulated in Table II. Those for W-H will prove useful in comparing the bonding of hydrogen to tungsten in the diatomic molecule with that in "surface molecules" such as W_0H

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used to model chemisorption.

Although the diatomic molecules in Table II are not known to exist, their EHMO bond lengths can be compared with the sum of the covalent radii of the atoms in the molecule. The Pauling single bond covalent radii of hydrogen and tungsten are 0.32 and 1.30 angstroms, respectively. The sum of the Pauling covalent radii for the diatomic tungsten molecule gives a bond length of 2.60Å for a single bond. The calculated bond length of 2.14Å implies a multiple bond whose order can be estimated from Pauling's²² empirical rule

 $D(n) = D(1) - 0.71 \log n$.

Here D(1) is the single bond length (2.60Å), D(n) the observed bond length (2.14Å), and n the bond order. Pauling's rule leads to a value of 4.4 for the bond order in reasonable agreement with the implications of Table II (5 occupied bonding orbitals, 1 occupied antibonding orbital). Hence the calculated bond lengths appear to be reasonable and self consistent.

A molecule of a hydrogen atom and two tungsten atoms was used to determine whether the repulsive energies as calculated in the analytical exponential term were pairwise additive. In this molecule, the tungsten atoms were separated by 3.16 Å, a characteristic distance on a tungsten (100) surface, and the hydrogen was equidistant

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from these two tungsten atoms. The repulsive energies plotted as curve a in Fig. 4 are determined by the differences between the energies calculated by the EHMO technique where the basis set of atomic orbitals included the 5p as well as the valence (6s, 5d) orbitals of the tungsten atoms and those calculated where the 5p orbitals were omitted from the basis set. The repulsive energies, as calculated by the analytical exponential repulsive term for two tungsten-hydrogen interactions, are plotted as curve b in Fig. 4. From the comparison of curves a and b, it is confirmed that the repulsive energy is reasonably well represented by the sum of all pairwise interactions.

B. Hydrogen Chemisorbed on a W(100) Surface

Before calculations were performed with the hydrogen at the proposed chemisorbed sites, it was necessary to make adjustments on the VOIP of the edge tungsten atoms. Because the edge atoms of the surface arrays were missing nearest and next nearest neighbors, their VOIP were adjusted to prevent abnormal charge distribution. The VOIP of the 5d and 6s orbitals of the four corner atoms of the arrays in Figs. 1a and 1b were adjusted to -8.52 ev and -7.52 ev, respectively; these same orbitals on the other four edge atoms were adjusted to -8.76 ev and -7.76 ev, respectively. The orbitals of the eight edge atoms of the array in Fig. 1c were adjusted to -8.68 ev

for the 5d orbitals and -7.68 ev for the 6s orbital. The VOIP of the remaining atoms in these arrays were not adjusted.

Mattheiss' ²⁶ band theory calculation on tungsten provides two check points for our EHMO calculations on the W₉ molecule. First, he finds a d-band width of 10.47 eV. We take as the EHMO analog the difference in energies between highest and lowest molecular orbitals having d character on atom 1 in the surface array of Fig. 1a. The lowest energy of such an orbital is Ψ_2 (shown in Fig. 6a), the highest (not shown because it is unoccupied) is 9.45 eV higher. Second, Mattheiss finds that the bottom of the d band lies 1.36 eV above the bottom of the conduction band. Our analog is the difference between Ψ_1 in Fig. 6a and the first occupied orbital of d character (not totally symmetric and therefore not shown in Fig. 6a): this difference is 1.34 eV, remarkably close to Mattheiss' result.

Calculations were then carried out with hydrogen bonded at the three possible symmetric sites. The minimum charge transfer to the hydrogen occurred with hydrogen bonded at the 5 CN site and the maximum with the hydrogen bonded at the 1 CN site. In the initial calculations, the VOIP of the hydrogen 1s orbital was adjusted as a function of atomic charge q and iterated to self consistency holding the tungsten orbital VOIP constant at values previously given. The final iteration VOIP were -11.34 ev at the 5 CN site, -9.54 ev at the 2 CN site, and -8.84 ev at the 1 CN site. These differences raised

106 12a the possibility that energy differences between various absorption configurations might reflect too strongly the different parametrizations of the hydrogen 1s VOIP. Since these energy differences were of principle concern, the calculations were repeated using a VOIP of -10.30 for all sites. Comparing the two sets of calculations, it was found that differences in bond energy and equilibrium distances between configurations were almost unaffected, and most importantly that the order of bonding energies among the three configurations remained unchanged. Hence use of a fixed hydrogen VOIP of -10. ev was adopted for all subsequent calculations for computational simplicity.

107 12Ъ The dependence of bonding energy E on internuclear separation r is plotted in Fig. 5 for hydrogen bonded at each of the three symmetric sites. The internuclear separation was the distance of hydrogen to the nearest tungsten atom(s) in the surface atom arrays. In the case of the 5 CN site, it was the distance to the tungsten atom in the hole, for the 2 CN site, it was the distance to the two surface atoms, and for the 1 CN site, it was the distance to the single tungsten atom. The bond energy E was calculated according to the following relationship:

$$E = \sum_{W_n H} n_i \epsilon_i - \sum_{W_n} n_i \epsilon_i - \epsilon_H + \sum_{W_n H} Ae^{-r/\alpha}$$

where the first term is the summation over the occupied molecular orbitals of energy ϵ_i of the surface "molecule" W_nH , the next term, is the summation over the occupied molecular orbitals of energies ϵ_i of the W_n part of the "molecule" separated from the hydrogen and the next term is the energy of the singly occupied atomic orbital of a separated hydrogen atom. The last term, the repulsive energy term, represents the repulsion of the occupied inner non-bonding orbitals (omitted from initial calculations), and the summation extends over all hydrogen-tungsten pairs in the surface array. This term is not necessary to determine the bond energy if the inner orbitals are included in the basis set for the MO calculations. Table III presents bonding energies, equilibrium distances, and charges corresponding to the energy minima in Fig. 5.

A somewhat simpler description and a better understanding of the bonding between hydrogen and the surface arrays can be obtained by using group theory. First, the surface "molecules" must be assigned to a symmetry group. The 1 CN and 5 CN sites on the W(100) surface have fourfold symmetry and the surface arrays in Fig. 1 representing the surface for each of these symmetric sites belong to the C_{4v} symmetry group. The 2 CN site has twofold symmetry and the surface array representing this site belongs to the C_{2v} symmetry group. Because the hydrogen 1s orbital is totally symmetric and belongs to the a₁ irreducible representation in these symmetry groups, it will interact only with the molecular orbitals of the tungsten arrays belonging to this same totally symmetric irreducible representation a_1 . For the C_{4v} surface arrays with nine tungsten atoms (Figs. 1a and 1b), there are twelve totally symmetric molecular orbitals formed by the valence orbitals of which only five are occupied. The C_{2v} surface array (Fig. 1c) with twelve tungsten atoms has twenty totally symmetric molecular orbitals of which eleven are occupied. An additional totally symmetric molecular orbital is formed as the hydrogen 1s orbital interacts with molecular orbitals of the W_n arrays.

Molecular orbital energy diagrams are shown in Fig. 6 for the hydrogen interacting with each of the surface arrays at equilibrium separation. The totally symmetric molecular orbitals of the tungsten arrays are indicated on the left side of the panels in Fig. 6 with the hydrogen 1s orbital on the right side and in the middle are the molecular orbitals formed by the W_nH surface "molecule". The lines joining the molecular orbitals of the separated W_n and H atomic orbital with those of the combined W_nH show the change in energy of the W_n molecular orbitals as they interact with the hydrogen in qualitative form; corresponding quantitative information can be obtained by comparing Tables IV and V (which tabulate the occupied totally symmetric molecular orbitals of the W_9 and W_{12} surface arrays) with Tables VI, VII and VIII (which tabulate the totally symmetric orbitals of the W_nH surface "molecules"). From these tables, it is observed that hydrogen makes only a small contribution to some of the molecular orbitals of the surface "molecule". These are indicated in Fig. 6 as orbitals not joined by a line to the hydrogen ls orbital.

It is interesting to note that, for the 1 CN and 2 CN sites, interaction of the hydrogen atom with the k occupied molecular orbitals in the totally symmetric a₁ representation leads to k+1 molecular orbitals all of which are below the highest occupied molecular orbital

(HOMO), and which are correspondingly occupied and contribute to bonding in the W_n H molecule. In the 5 CN site one of the molecular orbitals in the W_n H molecule is above the HOMO (Ψ_6 in Fig. 6c) and so does not contribute to bonding.

IV. DISCUSSION

The observed^{6, 9} LEED patterns from a $C(2 \ge 2)$ surface structure of hydrogen on the W(100) surface indicate that hydrogen must be chemisorbed at a site with 4-fold symmetry, either the 5 CN or 1 CN site. Evidence for this comes from the diffraction spots of the $C(2 \ge 2)$ surface structure, where the spots at the half order position $\left[\left(\frac{1}{2}, \frac{1}{2}\right), \left(\frac{1}{2}, \frac{3}{2}\right)\right]$, etc.] are diffuse, resulting from $C(2 \ge 2)$ domains on the surface which are not in phase. The facts that the spots at the integral order positions $\left[(1,0), (1,1), \text{ etc.}\right]$ are sharp and only those at the half order positions are diffuse indicate that hydrogen must be chemisorbed at sites with 4fold symmetry in the $C(2 \ge 2)$ surface structure.²³ From the results of our calculations as shown in Table III, the 1 CN site can be chosen as the preferred site with 4-fold symmetry for the chemisorbed hydrogen atom.

Early experimental evidence led to conflicting models for hydrogen chemisorbed on a W(100) surface including: atoms bonded at a single site, ^{6,8} atoms bonded at different sites, ⁹ and a mixture of atomic and molecular species bonded to the surface. ⁷ Recent evidence^{10,11}

indicates that atomic species are bonded at one kind of surface site; the two peaks, β_1 and β_2 , in the flash desorption spectrum are a result of a density dependent interaction on the surface. The high temperature state, β_2 , has been associated with a C(2 x2) surface structure and as the remaining sites are populated, a (1 x 1) surface structure results. From the results of our calculating, these sites must be the 1 CN sites on the W(100) surface.

In the WH molecule, the hydrogen 1s orbital appears only in the single molecular orbital Ψ_1 (see Table III). In this case the tungstenhydrogen bond can be interpreted as a simple single bond involving a hydrogen 1s orbital and a tungsten sd hybrid orbital. It is tempting to interpret the 1 CN preference for hydrogen adsorption as reflecting a preference for simple single bonding to the tungsten atom below it. The present treatment is not well designed to provide a justification for such an interpretation. Table VI shows that the molecular orbitals Ψ'_1 , Ψ'_2 , Ψ'_4 and Ψ_6 all have substantial hydrogen 1s contributions; while the principal tungsten orbital contributions to Ψ'_1 and Ψ'_2 arise from the tungsten atom underlying the hydrogen atom this atom contributes almost negligibly to Ψ'_4 and Ψ_6 , whose tungsten contributions come largely from the d orbitals of the four tungsten atoms surrounding the adsorption site. Such molecular orbitals are

not well suited for interpretation in terms of localized bonding; for this purpose optimally localized orbitals are more appropriate as shown by Ruedenberg and Edmiston.²⁴ The tungsten atoms which are involved with the bonding of the hydrogen atom retain their bonds with the neighboring tungsten atoms, but their strengths change as a result of the chemisorbed hydrogen. The reduced overlap population calculated by the Mulliken¹⁹ method can assess the relative change in strengths of these tungsten-tungsten bonds. With hydrogen bonded at the 1 CN site, the reduced overlap population of the central tungsten atom with each of its four nearest neighboring tungsten atoms is reduced 15%.

The small difference in the bonding energy between a 1 CN site and a 2 CN site can account for the observed mobility of hydrogen on tungsten surfaces. Gomer, et. al.²⁵ observed the diffusion of hydrogen on a tungsten surface at a temperature of 180° K and reported the activation energy for diffusion E_d was generally 10 to 20% of the bonding energy E_a , that both E_a and E_d/E_a were lowest on most closelypacked faces, and that the lowest value of E_d observed was 5.9 kcal (0.26 ev). If the diffusion of hydrogen on the W(100) surface occurs from one 1 CN site to another 1 CN site via a 2 CN site, and the only energy barrier to diffusion is the difference between bonding energies at these two sites, then the activation energy for diffusion would be 0.25 ev.

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Table I. Atomic orbital parameters for Extended Hückel Molecular Orbital calculations. Orbital exponents are for Slater type orbitals. Valence orbital ionization potential (VOIP) parameters reflect dependence of VOIP on charge q according to VOIP = $Aq^2 + Bq + C$.

Atom	Orbital	Orbital Exponent	$ \begin{array}{c} A \\ (ev/ e ^2) \end{array} $	B (ev/[e])	C (ev)
Hydrogen	16	1.20	13.60	27.20	13.60
Tungsten	5p	4.91	0.00	0. 00	46.07
	5 d	2.20	0.00	3.50	9.00
	6s	1.40	0.00	3.50	8.00

W-W	Ψ_1	Ψ_{2}	Ψ_3	Ψ_4	Ψ_5	Ψ_6
Energy (ev)	(-10.63)	(-10.57)	(-10.57)	(-9.57)	(-9.57)	(-9.23)
W(1,2)						
5d _z 2	16					63
$5d_{xz}$		±. 62				
$5d_{x^2-y^2}$. 68		
5dvz			±. 62			
5d _{xy}					. 68	
68	. 52					16
W-H	 Ψ1	Ψ2	Ψ3	Ψ_4	Ψ ₅	
Energy (ev)	(-12.23)	(-9.42)	(-9.42)	(-9.42)	(-9.42)	
н						
ls	62					
w						
$5d_{z}^{2}$	47					
⁵ d _{xz}		1.00				
$5d_{x}^{2}y^{2}$			1.00			
$5d_{yz}$				1.00		
5d _{xy}					1.00	
,	24					

Table II. Occupied molecular orbitals for the diatomic molecules, W-W and W-H.

Bond Energy (ev)	r _e (Å)	Charge e
3.00	1.65	36
2.74	1.96	26
1.72	1.61	+. 04
	Bond Energy (ev) 3.00 2.74 1.72	Bond Energy (ev) re (Å) 3.00 1.65 2.74 1.96 1.72 1.61

Table III. Energies, equilibrium distances, and charges for hydrogen bonded to different W(100) surface sites.

Molecular Orbital	Ψ_1	Ψ_2	Ψ_3	Ψ_4	Ψ_5
Energy (ev)	(-12,43)	(-10.69)	(-10.38)	(-9 . 77)	(-8.47)
W(1)					
5d _z 2		54		10	19
6s	24				
W(2, 3, 4, 5)					
5dz2		. 22	.17	.16	
5d _{xz} (2,5)		±. 15		±. 31	±.07
$5d_{x^2-y^2}$		±. 12	±. 30	±. 19	±. 05
5d _{yz} (3,4)		±. 15		. ±. 31	±.07
65	18	·			
W(6, 7, 8, 9)				•	
5d _z 2		10	19		.26
5d _{xz}		·	±.11		±. 28
5dyz			±. 11		±. 28
5d _{xy}		±. 12		±.22	±. 10
6s	12				

Table IV. Totally symmetric molecular orbitals for the surface arrays representing the 1 CN and 5 CN sites.

	ψı	ψ ₂	ψ3	ψ_4	ψ5	ψ6	ΨŢ	ψ8	ψ9	ψ10	ψ_{11}
	(-12.78)	(-11, 31)	(-10.87)	(-10.82)	(-10.25)	(-10.03)	(~9.68)	(-9.62)	(-8.97)	(-8.72)	(-8.32)
W(1,2)											
5d-2			36	09		08	.09	1 5			.15
$5d_{y}2_{y}2_{z}$.33			19	06	05	08	30	14	.08
⁵ dyz 6s	. 18	±. 12	±.08	±. 32	±. 21		±.17	±. 18	±.10	±. 1 2	
W(3,4)											
5d_2			.36	09		08	09	15			15
5d _{xz}		±. 12	±. 08	±.32	±.21	• - •	±.17	±.18	±. 10	±. 12	
5d _x 2-y2 6s	. 18	33			19	.05	.05	+.08	30	.14	.08
W(5, 6, 7, 8)											
5d_2			. 12	. 05	17	.07	08	.06	. 08	12	.11
$5d_{wz}$			±.05	±.07	±,14	±, 23		±.09	•	-	±.26
$5d_{x}^{2}-v^{2}$		13	.05	11	.10	16	.05		20	05	14
5dvz						±.07	±.1 3	±. 19	±. 14	±. 24	±. 10
$5d_{xy}$				±. 12		±. 08	±. 24	±.1 8	±.09	±.1 8	±. 08
68	.10										
W(9, 10, 11, 12)											
5d_2			.12	±. 05	.17	.07	.08	.06	.08	12	11
$5d_{xz}^{2}$			• •			±. 07	±. 13	±. 19	±.1 4	±. 24	±.1 0
$5d_{x}2-y^{2}$.13	±. 05	.11	.10	.16	.05		±. 20	.05	14
5dyz			±. 05	±. 07	±. 14	±. 23		±.0 9			±. 26
5d _{xy} 6s	.10			±.12		±. 08	±. 24	±.1 8	±. 09	±. 18	±. 08

Table V. Totally symmetric molecular orbitals for the surface array representing the 2 CN site.

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Molecular Orbital	$\Psi_1^{'}$	Ψ ₂	 Ψ ₃	$\Psi_4^{'}$	Ψ_6	 Ψ ₅
Energy (ev)	(-12.74)	(-11.52)	(-10.39)	(-10.01)	(-9.66)	(-8.46)
Н						
ls	.36	49	06	.36	.26	05
W(1)						
5d _{z2}	.16	52		06		.12
6s	.24	.06		.05		
W(2,3,4,5)						
5d _z 2		.07	22	.11	.26	
$5d_{xz}$ (2, 5)	·	±. 05		±. 30	±. 19	±. 06
$5d_{x^2-v^2}$		±. 06	±.27	±.10	±.27	±. 08
$5d_{yz}$ (3, 4)		±.05		±.30	±.1 9	±. 06
6s	.13	.11		08	05	
W(6, 7, 8, 9)						
5d _z 2			.21		07	26
5d _{xz}			±. 11			±. 28
$5d_{vz}$			±.11			±. 28
5d _{xy}		±. 04		±. 22	±. 12	±.11
68	.10	.07				

Table VI. Totally symmetric molecular orbitals for the W₉H surface "molecule" where hydrogen is bonded at a 1 CN site.

a	ta 2 CN	site.										
Molecular Orbital Energy (ev)	Ψ ₁ ' (-13,15)	Ψ ['] 2 (-11, 82)	Ψ ₄ ' (-11, 25)	Ψ ['] 3 (-10.85)	Ψ_{5}'	Ψ ₆ (-10, 03)	Ψ ₇ (-9, 84)	Ψ ['] 8 (-9,65)	¥9 (-9.38)	Ψ ₁₀ (-8, 73)	[₩] 12 (-8,62)	Ψ ['] 11 (-8, 32)
•••												
H 18	.35	44	. 13	02	16	02	. 18	04	-,22	.06	-,21	.02
W(1,2)												
5d~2		. 05		36	09	. 08		. 16	09			. 15
5d.22	07	. 26	. 24	•	. 09			. 10	10	05	19	. 08
5dur	±.09	±.24	±.27	±. 13			±.07		±. 14	±.13	• • •	•
68	.18										-,05	
W(3,4)												
5d 2		09	. 11	. 25	. 22	. 09	16	. 06	07			15
5d			+ 14	+ 23	+ 31			+ 25	+ 04	+ 07	+ 16	
5d a a		08	- 32		51	- 07	11	05	01	01		12
5°x ² -y ²	16	00	-, 36		.15	07	• • • •		. 15	4	* •20	• 14
08	• 1 3											
W(5,6,7,8)												
5d_2				. 13	. 09	09	. 18			12	05	. 13
5d		±.05		•	•••	±.21	±.14		±.20	±.07	±.21	±. 23
5d22		08	~. 08		. 08	. 18	- 11	. 06	. 26			-, 16
5d			±.06	±.05		±.07	14		±. 21	±. 25		±. 10
5d			±.03	±.09	±.06	±.08	±.05	±. 30		±. 13	±.15	±.08
6s	.06	09		,				-150	.05	06	.08	
W(9, 10, 11, 12												
6d o				00	17	05	,,		07	0.0	10	0.9
54			τ 0¢	08	I/	05 04	11	09	107 1 AF	-,08	10	-,08
5d a a			±,05	00	12	≭ ,00	± , U5	±, 64	±.05	≭,10 10	4,17	
54			. 14	.09	-,13	-,15		.00	. 10	. 12	20	11
⁵ ayz				±.07	±.10	±.25	±.09	±,06			1 07	±. 28
⁵ ⁴ xy	00	04			±.11	±,06	±,20	±. 08	±,13	±, 61	±,U/	±. 10
40	.08	.00										

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Table VII. Totally symmetric molecular orbitals for the W12H surface "molecule" where hydrogen is bonded

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		, ,				
Molecular Orbital	Ψ_1	Ψ2	$\Psi_3^{'}$	$\Psi_4^{'}$	$\Psi_5^{'}$	Ψ ₆
Energy (ev)	(-13.22)	(-11.75)	(-10.42)	(-9.96)	(8.62)	(7.36)
н						
ls	. 44	.29	04	11	13	.17
w						
$5d_{z^2}$.19	.38	.15	.10	11	.54
6s	.16	17				
W(2, 3, 4, 5)						
$5d_{z^2}$	05	11	23	.09	08	.38
$5d_{xz}$ (2, 5)				±.35		
$5d_{x}2_{-y}2$	±. 08	±. 15	±. 25		±. 05	±. 16
5d _{yz} (3, 4)				±. 35		
65	. 12	11				
W(6, 7, 8, 9)						
$5d_{z^2}$.21		.25	.17
$5d_{xz}$			±.10		±. 28	

±.10

.06

- -.11

±.28

.08

±.12

-.05

-.25

 $5d_{yz}$

5d_{xy}

6s

Table VIII. Totally symmetric molecular orbitals for the W9H surface "molecule" where hydrogen is bonded at a 5 CN site.

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Figure Captions

- Figure 1. Surface arrays used to model adsorption on different W(100) surface sites.
- Figure 2. Dependence of bond energy on bond distance for W₂ (left figure) and WH (right figure) molecules with and without inclusion of W5p orbitals.
- Figure 3. Dependence of non-bonding (5p orbital) repulsive energy on distance for W_2 and WH molecules.
- Figure 4. Additivity of non-bonded orbital repulsive energy.
 - (a) Difference in E(r) for W₂H between calculations including and omitting W 5p orbitals.
 - (b) E(r) calculated by pairwise addition of analytic repulsive energy functions.
- Figure 5. Bond energies as functions of distances for hydrogen adsorbed on different W(100) surface sites.
- Figure 6. Molecular orbital-energy diagrams for the totally symmetric occupied orbitals for hydrogen adsorbed on different W(100) surface sites.



Figure 1. Surface arrays used to model adsorption on different W(100) surface sites.



Figure 2. Dependence of bond energy on bond distance for W_2 (left figure) and WH (right figure) molecules with and without inclusion of W 5p orbitals.



Figure 3. Dependence of non-bonding (5p orbital) repulsive energy on distance for W₂ and WH molecules.



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Figure 4. Additivity of non-bonded orbital repulsive energy.

- (a) Difference in E(r) for W_2H between calculations including and omitting W 5p orbitals.
- (b) E(r) calculated by pairwise addition of analytic repulsive energy functions.



Figure 5. Bond energies as functions of distances for hydrogen adsorbed on different W(100) surface sites.



Molecular orbital-energy diagrams for the totally Figure 6. symmetric occupied orbitals for hydrogen adsorbed on different W(100) surface sites.

c) 5CN SITE

APPENDİX B.

A MOLECULAR ORBITAL INVESTIGATION OF

CHEMISORPTION. II. NITROGEN ON TUNGSTEN (100) SURFACE

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A Molecular Orbital Investigation of Chemisorption. II. Nitrogen on Tungsten (100) Surface

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Abstract

The relative bonding energies of nitrogen chemisorbed at three symmetric sites on a W(100) surface, represented by finite arrays of tungsten atoms,¹ were obtained by means of the Extended Hückel Molecular Orbital Theory (EHMO). The preferred site for nitrogen chemisorption was found to be the five coordination number (5 CN) site or the fourfold site with a tungsten atom below four tungsten atoms surrounding the nitrogen atom. The 5p orbital repulsive energy, in the case of hydrogen chemisorption, could be adequately approximated by the sum over pairs of empirical exponential repulsive terms; in the case of nitrogen chemisorption this same method was approximately
10% in error at the equilibrium bond distance, and repulsive energies were therefore obtained from calculations including tungsten 5p orbitals but with smaller arrays.

I. INTRODUCTION

In a previous paper, ¹ the Extended Huckel Molecular Orbital (EHMO) technique was applied to the bonding of hydrogen to a finite cluster of tungsten atoms representing a truncated tungsten (100) surface. The relative stabilities of hydrogen bonded at the three symmetric sites of a tungsten (100) surface were compared. In addition, the contributions made by the various orbitals of the surface atoms were investigated. According to the calculations, hydrogen will preferentially chemisorb on top of a surface tungsten atom forming a bond closely resembling that of a WH diatomic molecule.

In this paper this same molecular orbital (EHMO) technique has been used to investigate the bonding of nitrogen on a tungsten (100) surface. The nitrogen atom differs from the hydrogen atom particularly in its ability to form multiple bonds with the surface. The present calculations should be able to assess the influence of the additional nitrogen orbitals on a preferred chemisorption site.

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II. PROCEDURE

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The EHMO theory as formulated by $Hoffmann^2$ and described in our previous paper¹ has been used to investigate the preferred structure of the W_nN surface "molecule" consisting of nitrogen chemisorbed on a truncated W(100) surface. The surface arrays which represent the three symmetric sites for chemisorption on this surface are shown in Fig. 1. These are the same sites previously described.¹

Table I lists the orbital exponents for the Slater type orbitals (STO) of the nitrogen 2s and 2p orbitals derived from Slater's rules.³ It also includes values, taken from the work of Bacch, Viste and Gray,⁴ for the parameters A, B, and C which are needed to calculate the valence orbital ionization potential (VOIP) as a function of charge,¹ i.e.,

$$VOIP = Aq^2 + Bq + C .$$
 (1)

The EHMO technique was first applied to the WN diatomic molecule. A comparison of curves a and b in Fig. 2 shows that it is necessary to include the 5p orbitals of the tungsten atom as well as the valence orbitals in the basis set to obtain a characteristic potential energy curve. The population of these non-bonding orbitals provides the repulsion needed at small internuclear separations. This repulsive energy can be approximated by

$$E_{\rm R} = A_{\rm e}^{-r/\alpha} \qquad (2)$$

As the nitrogen atom forms a surface molecule, hopefully the repulsive energy for nitrogen interacting with a larger group of tungsten atoms can be approximated by the pairwise summation of exponential terms according to Eq. (2).

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III. RESULTS

A. Diatomic Molecule

In Fig. 2, the bonding energy is plotted as a function of internuclear separation r along the molecular axis z for the WN molecule. The bonding energy was calculated by subtracting from the total energy (energy of the occupied molecular orbitals), the energy of the ions of the W-N molecule at infinite separation.¹ The difference between curves a and b in Fig. 2 is the repulsive energy resulting from the presence of the 5p orbitals of tungsten in the basis set. In Fig. 3 $\ln E_{\text{Rep}}$ is plotted as a function of internuclear separation r. Values for the parameters A and α needed to calculate the repulsive energy according to Eq. (2) were found from this plot to be A_{WN} = 2324.8 ev and $\alpha_{\text{WN}} = .234\text{\AA}^{-1}$. For the final iteration, the charge on the nitrogen atom in the WN molecule at equilibrium separation was -.38 and that on the tungsten atom +.38.

Single bond covalent radii (0.70Å for N and 1.30Å for W) predict a length of 2.00Å for a W-N single bond. Pauling's empirical relationship⁵ between bond length and bond order (see previous paper¹),

coupled with the 1.65 $^{\text{A}}$ equilibrium bond length indicated by Fig. 2, gives a value of 3.1 for the bond order of the WN molecule. Table II, which lists the LCAO for the occupied molecular orbitals of the WN molecule, shows that there are three molecular orbitals (Ψ_2 , Ψ_3 and Ψ_4) which make major contributions to the bonding: this suggests that the bond order of 3.1 is a plausible value.

Figure 4 compares the W₂N molecule repulsive energies determined (curve a) as the difference between the energies calculated by the EHMO technique with the inclusion and omission of the tungsten 5p orbitals and (curve b) by summing analytical exponential terms over pairwise interactions. There is about a 10% difference in the repulsive energies in the neighborhood where the chemical bond is formed. Because of this difference, larger molecules (W_4N and W_5N) which still lended themselves to easy computation were investigated; these molecules are reductions of the surface arrays in Fig. 1. The tungsten atoms omitted from the surface arrays in Fig. 1 for the above calculations are at a sufficient distance from the nitrogen atom at each of the symmetric sites that their 5p orbitals should not contribute to the repulsive energy when the larger surface arrays are used. In Fig. 5 the repulsive energies between the nitrogen atom at each of the bonding sites and the tungsten atoms in the above molecules are plotted.

Curves a in Fig. 5 are based on repulsive energies calculated

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from the W_5N or W_4N arrays as explained above, curves b by pairwise summation of analytical exponential terms. Repulsive energies at bonding distances obtained from curves a are 10% higher for 1 CN site, about the same for the 2 CN site, and about 10% lower for the 5 CN site than those obtained from curves b. Approximation of repulsive energy by pairwise summation of exponential terms (Eq. 2) was therefore judged insufficiently reliable, and it was approximated instead from direct calculations on the abbreviated arrays W_4N and W_5N .

B. Nitrogen Chemisorbed on a W(100) Surface

The adjustments of the VOIP of the edge atoms of the surface arrays in Fig. 1 to reduce charge transfer were the same as those previously adopted.¹ The VOIP of the orbitals of the nitrogen atom were iterated at equilibrium separation to self consistency for charge q at each of the three symmetric sites. On the final iteration, the VOIP for the 2s and 2p orbitals of nitrogen were: -18.93 ev and -10.01 ev at the 5 CN site, -17.17 ev and -8.93 ev at the 2 CN site and -17.31 ev and -9.02 ev at the 1 CN site. To avoid possible oversensitivity of results to VOIP parametrization, the calculations were repeated using -17.77 ev and -9.29 ev for the 2s and 2p orbitals of nitrogen respectively at all the sites. As in the case of hydrogen chemisorption¹ the same ordering in site preference was obtained by both methods

and constant nitrogen 2s and 2p VOIP's were used in further calculations. The bonding energies as a function of internuclear separation are plotted in Fig. 6. The internuclear separation was the distance of nitrogen to the nearest tungsten atom(s) in the surface atom arrays. In the case of the 5 CN site, it was the distance of the tungsten atom in the hole, for the 2 CN site, it was the distance to the two surface atoms, and for the 1 CN site, it was the distance to the single tungsten atom. The bonding energy E was calculated according to the following relationship:

$$\mathbf{E} = \sum_{\mathbf{W}_{n}N} \mathbf{n}_{i} \boldsymbol{\epsilon}_{i} - \sum_{\mathbf{W}_{n}} \mathbf{n}_{i} \boldsymbol{\epsilon}_{i} - \sum_{N} \mathbf{n}_{i} \boldsymbol{\epsilon}_{i} + \mathbf{E}_{Rep}$$
(3)

where the first term is the summation over occupied molecular orbitals of energy ϵ_i of the W_nN surface molecule, the next term is the summation over the occupied molecular orbitals of energies ϵ_i of the W_n part of the molecule at infinite separation from the nitrogen atom and the next term is the summation over the occupied atomic orbitals of energy ϵ_i of the nitrogen atom also at infinite separation. The last term is the repulsive energy E_{Rep} determined from the W_5N and W_4N molecules and plotted as curves a in Fig. 5. Table III summarizes the results of calculations plotted in Fig. 6.

The surface arrays in Fig. 1a and 1b belong to the C_{4v} symmetry groups and the surface array in Fig. 1c to the C_{2v} symmetry group.¹ In these groups, the 2s and $2p_z$ orbitals of nitrogen belong to the totally symmetric a₁ irreducible representation. Therefore, these orbitals

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of nitrogen can interact only with the totally symmetric molecular orbitals of the W_n surface arrays. In the C_{4v} symmetry group, the $2p_x$ and $2p_y$ orbitals of nitrogen belong to the e irreducible representation and will interact with molecular orbitals of the W₉ arrays which also belong to this representation. These same orbitals of nitrogen belong to the b_1 ($2p_x$) and b_2 ($2p_y$) irreducible representations in the C_{2v} symmetry group and interact with the molecular orbitals of the W₁₂ array belonging to these same irreducible representations.

Molecular orbital energy diagrams for the molecular orbitals belonging to the totally symmetric a_1 irreducible representation are shown in Fig. 7. On the left side of each panel in these figures are the molecular orbitals from the W_n arrays, on the right side are the atomic orbitals of nitrogen and in the middle are the molecular orbitals for the W_n H surface "molecules". The occupied totally symmetric (a1) molecular orbitals in Fig. 7 are tabulated in Tables IV and V for the W_g and W_{12} arrays respectively. These can be compared with the totally symmetric molecular orbitals of the W_n N surface "molecules" (Table VI - 1 CN, Table VII - 2 CN and Table VIII - 5 CN). Because of the totally symmetric orbitals (2s and $2p_z$) of nitrogen, there must be two additional totally symmetric molecular orbitals in the W_n N surface "molecule" than in the W_n "molecule". In Fig. 7a and 7c, these molecular orbitals are Ψ_6 and Ψ_7 . For nitrogen bonded at the 1 CN

site (Fig. 7a) these molecular orbitals are lower in energy than the highest occupied molecular orbital (HOMO) and therefore are occupied, whereas at the 5 CN site (Fig. 7c) Ψ_7 is above the HOMO. At the 2 CN site (Fig. 7b) the additional molecular orbitals (Ψ_{12} and Ψ_{13}) of the W_{12} N formed by the nitrogen 2s and $2p_z$ orbital interacting with the W_{12} surface array are below the HOMO but one of the totally symmetric orbitals (Ψ_{11}) has become unoccupied in the reordering of all the molecular orbitals in the W_{12} N surface "molecule".

In Fig. 8, molecular orbital energy diagrams are shown which involve the $2p_x$ and $2p_y$ orbitals of nitrogen. In the case of the W_9 arrays for the 1 CN and 5 CN sites (Fig. 8a and c) these molecular orbitals belong to the e irreducible representations and for the 2 CN site (Fig. 8b) they belong to the b_1 and b_2 irreducible representations. The molecular orbitals represented in Fig. 8 for the W_9 and W_{12} are tabulated in Tables IX and X respectively. These can be compared with the molecular orbitals of the W_nN surface "molecule" (Table XI -I CN, Table XII - 2 CN, Table XIII - 5 CN).

The additional molecular orbitals formed by the $2p_x$ and $2p_y$ atomic orbitals of nitrogen as they interact with the W_n surface arrays are Ψ_{15} and Ψ_{16} in the case of the W_9N and Ψ_{19} and Ψ_{20} in the $W_{12}N$ surface "molecules". The molecular orbitals Ψ_{19} and Ψ_{20} of the $W_{12}N$ "molecule" are below the highest occupied molecular

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(HOMO) and are therefore occupied. But in the case of the W₉N "molecules", Ψ_{15} and Ψ_{16} (having the same energy) are the HOMO; these molecular orbitals of the W₉N "molecule" with nitrogen at the 1 CN site (Fig. 8a) are occupied with a single electron, whereas at the 5 CN site (Fig. 8c), they are occupied with three electrons.

IV. DISCUSSION

Low-energy electron-diffraction (LEED)⁶ has indicated that nitrogen adsorbed on a W(100) surface produces a C(2x2) structure and flash desorption spectrometry (FDS)⁷ indicates it desorbs as a single β state. It has been proposed that this adsorbed structure has only half the surface sites occupied with atomic species. Estrup and Anderson⁶ originally proposed that the preferred chemisorption sites are alternate 5 CN sites. It has also been shown that a molecular bound state can be converted to another β state, presumably also atomically adsorbed nitrogen,^{7,8} and proposed that these atoms could be filling 5 CN sites unoccupied in the original C(2x2) structure.

Our calculations indicate that nitrogen prefers the 5 CN site. At this site, the σ bond formed by the 2s and $2p_z$ atomic orbitals of nitrogen with the atoms of the surface array is not as favorable as at the other sites (Fig. 7); hydrogen also formed a weaker σ -bond at this site¹. But in the case of nitrogen the $2p_z$ and $2p_y$ atomic orbitals

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interact more favorably with the tungsten atoms surrounding the 5 CN site (Fig. 8) than at the other sites, and this factor dominates the selection of the 5 CN site for chemisorption. As Table XIII shows, these atomic orbitals are able to form the lowest energy molecular orbitals of e symmetry at the 5 CN site because of their overlap with the appropriate d orbitals of the four surrounding tungsten atoms.

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Orbital	Orbital Exponent	A [ev/(e) ²]	B [ev/e]	C [ev]
2s	1.950	3.48	20.25	28.02
2p	1.950	3.72	14.14	16.05

Table I. Orbital exponents for STO of the nitrogen atom and parameters needed to calculate VOIP as a function of atomic charge.

					 \T/	
Energy (ev)	Ψ1 -21.61	Ψ2 -12.74	Ψ ₃ -12.74	Ψ ₄ -11.51	£5 -10.29	Ψ ₆ -10.29
N						
2s	. 86			. 13		
2p _z	.08	•		.72		
2p _x		-, 69				
2py			69			
w						
5d _z 2	. 21			48		
⁵ d _x 2		54				
$5d_{x^2-y^2}$					1.00	
5dyz			- .54			
5d xv						1.00
6s	. 13			.29		

Table II. Occupied molecular orbitals for the diatomic molecule WN.

	Site	Bond En er gy (Å)	ге (Å)	Charge
	5 CN	-3.25	2,05	04
•	2 CN	-3.15	2.13	-1.36
	1 CN	-2.85	1.88	-1.13

Table III. Energies, equilibrium distances, and charges for nitrogen bonded to different W(100) surface sites.

Molecular Orbital Energy (ev)	Ψ ₁ (-12.43)	¥2 (~10.69)	Ψ ₃ (-10.38)	Ψ ₄ (-9. 77)	Ψ ₅ (-8.47)
W(1)		<u> </u>			······································
5d _{z2}		54		10	19
6s	24				
W(2, 3 , 4, 5)					
$5d_{z2}$. 22	.17	.16	
5d _{xz} (2, 5)		±. 15		±. 31	±.07
5d, 2- y2		±. 12	±. 30	±. 19	±. 05
$5d_{vz}$ (3, 4)		±. 15		±. 31	±.07
6s	18				
W(6, 7, 8, 9)			·		
5d _z 2		10	19		.26
5d _{xz}			±.11		±. 28
5d _{vz}			±.11		±. 28
5d _{xy}		±. 12		±.22	±. 10
6s	12				

Table IV. Totally symmetric molecular orbitals for the surface arrays representing the 1 CN and 5 CN sites.

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	ψ1	ψ ₂	ψ3	ψ4	ψ5	ψ6	ψ7	ψ8	ψ9	ψ ₁₀	ψ ₁₁
	(-12.78)	(-11, 31)	(-10.87)	(-10.82)	(-10,25)	(-10.03)	(-9.68)	(-9, 62)	(-8,97)	(-8.72)	(-8.32)
W(1,2)											
5d~2			36	09		08	.09	1 5			.15
$5d_{y}2_{y}2_{z}$.33			19	06	05	08	30	14	.08
$5d_{vz}$		±. 12	±. 08	±. 32	±.21		±.1 7	±. 18	±. 10	±. 12	
68	.18										
W(3,4)											
5d, 2			.36	09		08	09	15			1 5
5d _{xz}		±. 12	±. 08	±.32	±,21		±,17	±. 18	±.10	±.12	
$5d_{x^2-y^2}$		33			19	.05	.05	+.08	30	.14	.08
6s ,	.18										
W(5,6,7,8)											
5d,2			.12	.05	17	.07	. 08	.06	.08	12	.11
$5d_{xz}^{2}$			±. 05	±. 07	±.1 4	±.23		±. 09			±. 26
$5d_{x}^{2}-v^{2}$		13	.05	11	.10	16	.05		20	05	14
5dvz						±.07	±.1 3	±.1 9	±.1 4	±. 24	±.1 0
$5d_{xv}$				±. 12		±. 08	±. 24	±.1 8	±. 09	±.1 8	±.0 8
68	.10							·			
W(9, 10, 11, 12)											
5d, 2			.12	±. 05	.17	.07	.08	.06	.08	12	11
5d _{xz}						±.07	±. 13	±. 19	±.1 4	±.24	±. 10
$5d_{x}2-y^{2}$.13	±. 05	.11	.10	.16	05		±.20	.05	14
5dyz			±. 05	• ±.07	±. 14	±. 23		±.0 9			±. 26
5d _{xv}				±. 12		±. 08	±. 24	±. 18	±.09	±. 18	±.0 8
6s ′	. 10										

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Table V. Totally symmetric molecular orbitals for the surface array representing the 2 CN site.

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Molecular Orbital	Ψ6	Ψ_1	Ψ2	Ψ'3	Ψ'_4	Ψ ₇	Ψ_5
Energy (ev)	(-18.33)	(-12.17)	(-10.57)	(-10.38)	(-9.82)	(-9.47)	(-8.43)
N	<u></u>						
2s	. 89	14	21	.04	.04	.03	
$2p_z$		03	.46	10	33	65	12
W(1)							
5d _z 2	.17	21	.38	05	07	06	.08
65	. 11	.19	07				
W(2, 3, 4, 5)						:	
5d _z 2			21	16	.07	25	08
5d _{xz} (2, 5)			±.1 2	±.06	±.3 3	· .	±. 05
$5d_{x}2_{-y}2$			±.08	±. 30	±.11	±. 22	±.10
5d _{yz} (3, 4)			±.1 2	±. 06	±. 33		±.0 5
6в		.18	.08				
W(6, 7, 8, 9)							
5d _z 2			.10	.19	·	.10	2 5
5d _{xz}				±. 11		±. 05	±. 28
⁵ dyz				±.11		±.0 5	±. 28
⁵ d _{xy}			±. 10		±.23		±.12
68		.12			•		

Table VI. Totally symmetric molecular orbital for W9N surface "molecule" where nitrogen is bonded at a 1 CN site.

Molecular Orbital	Ψ_{12}	Ψ_1^{+}	Ψ ₂	Ψ_3	Ψ_4	Ψ5	Ψ ₆	Ψ_7	Ψ [.] 8	Ψ9	Ψ ₁₀	Ψ ₁₃	Ψ_{11}
Energy (ev)	(-18.47)	(-12,42)	(-11,35)	(-10,98)	(-10,63)	(-10, 22)	(-10,03)	(-9.79)	(-9,62)	(-9.33)	(-8.72)	(-8,55)	(-8,26)
N													
25	86	-,17	.05	. 05	-,10	16	05	. 03	08		. 03	03	. 03
2p _z		.02	.18	. 26	26	47	05	. 34	16	. 32	01	. 28	. 17
W(1, 2)													
⁵ d _z 2			. 09	.36			.07	.07	. 11	07			09
5dx2_y2	.06		. 32		.06	14			.07	. 12	08	14	08
^{5d} yz	±.1 1	±.14	±.14	±,08	±. 15	±.17	±.07	±.11	±.13	±. 09	±. 12		±.06
68	±. 09	. 13				05							
W(3,4)		•											
5d ₂ 2				31	. 14	12	.09	. 19	05	05	09	- ,15	. 18
^{5d} xz			±.14	±. 08	±.3 7			±. 16	±.1 9	±.07	±.20	±.23	±.08
^{5d} x ² -y ²			31	. 12	.10	14	11	06		. 17			-,16
6 s		.18				05							
W(5, 6, 7, 8)													
5dz2				09		17	14	10			-,12		14
5d _{xz}				±. 05		±. 16	±,17	±. 10		±.19	±.05	±.29	±.13
⁵ d _x 2-y2			-, 12		.09		. 20	.08	.08	. 22			.17
5dyz			±. 05	±. 05				±. 15	±. 08	±. 20	±. 25		±, 10
5d _{xy}					±.1 1		±. 08	±,07	±,30	±, 11	±, 14	±.06	±.05
68		. 12				.05					05		
W(9, 10, 11, 12)													
5dz2				.10		.16			08	. 09	10	16	b
5d _{xz}			±.06				±. 06	±.1 7	±.1 7		±.21	±.14	±,15
^{5d} x ² -y ²			. 11	09		.06	13	.07		.11	. 10	22	.08
5dyz					•		±. 26	±. 08	±.17				±,31
- 5d _{xy}						±. 07		±,22		±.0 8	±.20	±.0	7 ±.12
68		. 10									•		

Table VII. Totally symmetric molecular orbitals for the $W_{12}N$ surface "molecule", where nitrogen is bonded at a 2 CN site.

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Molecular Orbital	Ψ_6	Ψ_1	Ψ'2	Ψ'_3	Ψ'_4	Ψ'_5	Ψ_7
Energy (ev)	(-18.71)	(-11.97)	(-10.70)	(-10.36)	(-9.39)	(-8,61)	(-7.49)
N							······································
2s 2p	82	.11	.10	25	.17	09	.06
² μ _Z		11	.50	. 25	• 40	.02	41
W(1)							
5d _z 2	12	.28	23	.09	31		.35
68	-,06	21					.08
W(2, 3, 4, 5)							
5d _z 2			.15	 17	13	07	. 22
5d _{xz} (2, 5)			±.21	±.1 8	±.1 6		±.21
5d _x 2_y2	±.06		±. 08	±. 24			±. 17
5d _{yz} (3, 4)			±. 21	±.1 8	±.1 6		±.21
6s	05	14					
W(6, 7, 8, 9)							
5d _z 2			11	.19	.08	.24	•
5d _{xz}			±. 05	±.07	±. 06	±. 30	
$5 d_{yz}$			±. 05	±.07	±. 06	±. 30	
⁵ d _{xy}			±.0 6	±. 10	±. 29		±. 35
6в		13			֥ 09		

Table VIII - Totally symmetric molecular orbitals for the W₉N surface molecule where nitrogen is bonded at a 5 CN site.

Molecular Orbital	$\Psi_1 \Psi_2$	¥3 ¥4	¥5 ¥6	Ψ 7 Ψ 8	Ψ ₉ Ψ ₁₀	$\Psi_{11} \Psi_{12}$	Ψ_{13} Ψ_{14}
Energy (ev)	(-10.68)	(-10,17)	(-9.71)	(-9.58)	(-9.24)	(-8.91)	(-8.80)
W(1)		~				· · · · · · · · · · · · · · · · · · ·	
5d _{xz}	50 .10		.21	. 19	.06 .08	19	0737
5dyz	.10 .50		21	. 19	08 .06	19	.3707
W(2, 5)			•				
5dz2	±.16	±.13 ±.15	±.26	±. 11	±.14 ±.18	±.10	±.07
5d _{xz}	23 .05	05	. 13		1722	29	. 10
5dx2_y2	±.12	±.17 ±.21	±.14	±.26	±.17 ±.22	±.07	
5dyz	17	.1613	16	. 35		08	. 05
5d _{xy}	±.22	±.19 ±.15	±.07			±.32	±.07
6.		±.07 ±.08	±.20	±.20	±.12 ±.16		±. 08
W(3,4)							
5dz2	±.16	±.15 ±.13	±. 26	±.11	±.18 ±.14	. ±.10	±.07
5d _{xz}	.17	.13 .16	.16	.35		08	05
^{5d} x ² -y ²	±,12	±.21 ±.17	±. 14	±. 26	±.22 ±.17	±.07	
5dyz	.05 .23	.05	13		.2217	29	10
^{5d} xy	±.22	±.15 ±.18	±.07			±. 32	' ±. 07
6.		±.08 ±.07	±. 20	±.20	±.16 ±.12		.08
W (6, 9)							
5dz2		±.21 .	±.14 ±.10			±.26 ±.32	±.08 ±.11
⁵ d _{xz}	.0809	08 .19	07	.18 .09	. 09	.0911	21 .11
^{5d} x ² -y ²	±.09 ±.06	±.11	±.17 ±.24		±.18	±.10 ±.08	±.17 ±.12
5dyz	.0611	0520	06 .05	.19	. 10	13 .07	. 24
^{5d} xy	±.08 ±.10	±.07	±.08 ±.05		±.26	±.09 ±.11	±.25 ±.37
6.		±.10	±.19 ±.14	±.15 ±.18	±.23	· ±.05	±.05 ±.08
W (7, 8)							
۶dz2		±.21	±.10 ±.14		±.05	±.32 ±.26	±.11 ±.08
^{5d} xz	.11 .06	.2005	.05 .06	. 19	.10	.07 .13	. 24
^{5d} x ² -y ²	±.06 ±.09	±.11	±.24 ±.17		±.18	±.08 ±.10	±.12 ±.17
^{5d} yz	0908	.19 .08	07	.09 .18		.11 .09 .	1121
^{5d} ×y	±.10 ±.07	±.07	±.05 ±.08		±.26	±.11 ±.09	±. 37
64		±.10	±.14 ±.19	±.18 ±.15	±.23	±.05	±.08 ±.05

Table IX. Molecular orbitals which belong to the e representation in the C_{4v} symmetry group for the W₉ surface arrays representing the 1 CN and 5 CN sites.

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Molecular Orbital	Ψ_1	Ψź	Ψ3	Ý4	Ψş	¥ 6	¥7	₩ 8	¥9	¥10	¥11	ŵ ₁₂	ч ₁₃	Ÿ ₁₄	¥15	¥16	¥17	¥18
Finargy (ev)	(-10, 92)	(-10, 92)	(-10, HH)	(-10, 88)	(-10.56)	(-10.56)	(-10.01)	(-10.01)	(-9.80)	(-9, 80)	(-9, 59)	(-9.59)	(-9.38)	(-9.38)	(-9.07)	(-9.07)	(-8, 62)	(-8. 62
w.1, 21													-					
54.2	e. 2H	+. 14			z. 10				±. 18	±, 19	\$.09	±. 07	±. 10	*. 08			±.14	×. 05
Nd _{an}	17	. 21	-, 06			Он	10		. 17	- 15			. 20	24	13			08
1.d. 2	1, 17	1.14	a, 05	±. 09	*.11			±.05					±. 12	±. 10			±.27	±. 10
`dy≖					25	05	.08	23			. 14	. 11	19	16			. 05	
···!=y	+. 12	4.14	1.14	1.08	1. OH	+. 37							±. 09	* . 11	±. 20			±.11
ti n				s, 18	#, OH													
W(1.4)																		
اليولەن [.]	4, 15	», 1H				±.10			±.19	±. 18	±.07	±. 09	*. 08	*. 10			±.05	s. 14
hd _{xr}					. 05	21	. 23	. 04			. 11	24	. 26	19				05
54_2-v2	1, 24	1.17	+. 09	±.05		a. 11	±.05						. *. 10	*. 12			*. 10	±. 27
٠d	21			06	#. ON			. 10	. 15	. 17			24	20		- , 13	.08	
5-1 x v	+. 14	+.12	1. UH	*. 14	#. 37	±.08							*.11	±.09		±. 20	±. 11	
he			+, 18	±. 09		*. OB												
W(5, #)																		
54 <u>.</u> 2	1,09					±.12	±.0 7		±. 27	±.16	a. 09	. 08			±. 09	±.14	*. 14	
5dgg					11			±.23		. 09		18	. 11	. 09	14	27	18	08
54 <u>22-</u> 2		1.17			±.11		±.08				* . 21	±.18	* . 18	* . 12	±.14	±.09		*. 13
Sdva.		. 15					. 09		. 08	±.16	. 18	. 15	10			13	24	18
hd av	#. ON				±.07			*. 30	±.07	≞. 09	±.08	±.08		#. 20·	±,07	±. 18		±. 15
6.			1.23	*. 16	a. 12													
W(6, 7)																		
54-2		1.09			±.07	# 10	*. 09		#. 14	z. 28	s . 06	±. 11			±.10	±. 13	±. 12	±. 07
ده					. 10	. 06	17	. 16	±.08		. 17	07	11	09	17	. 26	. 09	18
5d.2.2	*. 17				a. 10		±. 08				*. 23	±.15	*. 15	±.16	a. 13	±. 11	±.09	±. 10
5d	. 15						06	07	17	.06	. 19	. 13		*. 10	. 06	12	07	. 29
5dev		*. OH			*. 09		z. 19	*.23	±.09	±. 06	±. 05	±. 09	*. 19			±.19	±. 08	#. 13
fie .				±. 2×	±.10													
W(7, 12)								•••	. 10							+ 10		a 13
54z2	1.04				1 . 10	2.07		.04	z. 20	=. 14	*. 11	1.00	••		*. 15	E. 10	=.07	E. 16
""xz							.07	07	. 08		. 15	19	. 10	15	12	08	. 27	
⁷ a _x 2-y2		•. 17				4.10		¥.0A			1,15	1.23	#.10	=	*	x . 15	*.10	±,09
""YE						10	. 10	. 17		08	.07	.17	09	19	-, 20	11	. 10	.09
רי גע	3,08		+. 28		*. OH	*. 10 *. 10	±,23	±.14	4.06	±,04	±.09				8.17		2 . 13	2. 08
W(10, 11)																		
5dg2		1,04			*. 12			±.07	±. 16	±.27	±.08	±. 09			±.14	±. 09		±. 14
-d ₂₂	. 15							. 09	16	08	. 15	18		10	13		18	. 24
	±.17					±.11		*. 08			±.18	±.21	±. 12	±.18	±.09	a. 14	±.13	
hdys						. 11	23		09		. 18		. 09	11	. 27	14	. 08	18
∽d _{≍y}		*. OH				±.07	*. 30		±. 09	±.07	±, 08	±.08	±. 20		#. 18	07	±. 25	
65			*. 16	#. 23		*.12												

faile X. Molecular orbitale which belong to the b1 and b2 representations in the C2+ symmetry group for the W12 surface array representing the 2 CN site.

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Molecular Orbital	Ψ1 Ψ2	Ψ, Ψ4	Ψ ₇ Ψ8	Ψ'5 Ψ'6	Ψ13 Ψ14	¥9 ¥10	Ψ ₁₁ Ψ ₁₂	$\Psi_{15} \Psi_{16}$
Energy (ev)	(-10.92)	(-10,14)	(-9.87)	(-9.70)	(-9.35)	(-9.24)	(-8.90)	(-8.23)
N			·····					
Zp_	\$7	.0808	48		3025	. 05		46 .23
² Py	37	0808	. 48		25 .30	. 05		2346
w(1)								
5d xz	. 55		. 14	.10 .16		.0512	0506	19 .09
^{5d} yz	. 55		14	16 .10		1205	06 .05	0919
W(2,5)								
5dz2	±.12	±.16 ±.16		±.14 ±.23	±.13 ±.11	±.08 ±.19	±.08 ±.10	±.10 ±.05
5-d _{xz}	. 1.9		13	.07 .11	05	11 .27	1925	
5dx2-y2	±.12 ±.12	±.18 ±.18		±.06 ±.10	±.26 ±.21	±.08 ±.22		±.14 ±.07
` ^{hd} y≖	17	1111	37	11 .07	11 .13		05	0510
^{5d} xy		±.20 ±.20	±.07	±.06	±.11 ±.13		±.23 ±.17	
6 #		±.08 ±.08	±.11	±.12 ±.19	±.06 ±.05	±.08 ±.20	±. 05	±. 12 ±.06
W(3, 4)								
5elz2	±.12	±.16 ±.16		±.23 ±.14	±.11 ±.13	±.19 ±.08	±.10 ±.08	±.05 ±.10
hd xr	12	. 11 11	. 37	. 07 . 11	13 11		05	10 .05
⁵ ط x ² -y ²	±.12	±.18 ±.18		±.10 ±.06	±.21 ±.26	±.22 ±.08		±.07 ±.14
^{∽d} yz	. 19		. 13	11 .07	.05	···.27 .11	25 . 19	
"dxy	±.17	±.20 ±.20	±.07	±.06	±.13 ±.11		±.17 ±.23	
6 8		±.08 ±.08	±.11 .	±.19 ±.12	±.05 ±.06	±.20 '±.08	±.05	±.06 ±.12
W(6, 9)								
5dz2		±. 20	±.05 ±.05	±.17		±. 05	±.42 ±.05	
^{5d} xz	0706	.18 .09	. 16	.06	20	0506	11 .16	. 22
^{5d} x ² -v ²	±.05 ±.05	±.08	±.13 ±.11	±.28 ±.08	±.12	±.15 ±.07	±.06	±.10 ±.29
5dyz	0607	18 .09	16	07	20	07	0618	. 19 . 11
hdxy	+.05 ±.05	±. 10	±.07 ±.08	±.10	±.24	±.13 ±.29		±.30 ±.10
figs.		±. 12	±.09 ±.11	±.06 ±.25	±.09	±.10 ±.22	±.09	±.11
w(7, 8)								
hdz 2		±. 20	±.05 ±.05	±.17		±.05	±.05 ±.42	
^{≻d} x≖	07 .06	0918	• .16	.07	20	. 07	.1806	.1119
54x2_y2 -	±.05 ±.05	±.08	±.11 ±.13	±.07 ±.28	±. 12	±.07 ±.15	±. 06	±. 29 ±.10
''dyz	.0607	.0918	16	06	. 20	06 .05	.16 .11	10 .22
"dxy	±.05 ±.05	±.10	±.08 ±.07	±.10	±.24	±.29 ±.13		±. 30
fon.		±. 12	±.11 ±.09	±.25 ±.06	±.09	±.22 ±.10	±.09	±. 11

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Fable XI. Molecular orbitals which belong to the e-representation for the WgN surface "molecule" where nitrogen is bonded at a 1 CN site.

Molecular Orbital	Ψı	Ψz	Ψ3	¥4	₩5	Ψ6	¥7	Ψ8	¥9	¥10	Ψ11	Ψ12	Ψ ₁₃	Ψ ₁₄	Ψ15	Ψ ₁₆	Ψ19	420	Ψ ₁ -	¥15
Energy (ev)	(-11, 18)	(-11, 14)	(-10.90)	(-10.89)	(-10.70)	(-10, 63)	(-10.28)	(-10, 02)	(-9.86)	(-9.84)	(-9.67)	:- 2. 61)	(-9, 55)	1-9.46)	(- 2, 23)	1-9.071	⊬â.93)	(+ 8.72)	HB. 581	1. 5.50)
N																				
5 ⁵ ×	. 35		03			.10		-,08		22		+,10		31	. 34			-153	. 34	
Zpy		. 39		02	25		. 33		. 15		21		15			. 02	. 44			20
W(1, 2)																				
5d.22		±, 23			±,14		±,13		±.22 .	•	*.12		±. 13							±.11
5d _{xz}	.19		. 21			09		13		24				23	- , 10			. 14		
⁵ d _x 2_y2		±.17		±,16			±.16		±.05				±,05							±.23
^{5d} yz		. 21		-,05	. 27		14		.06		. 14		05				. 22			. 00
5d _{xy}	±.28					±,31		±.07		±,05		±.07		±.10	±.19			±.06	±.15	
ί s		±.0 6		±,20			±. 07													
W(3,4)												•								
5dz2	±.18		±.11					±.07		±.27		±.14		±. 06	±.12				±.23	
5d _{xz}			.07			22		. 23		06		.11		30	05				.06	
5dx2-y2	±.08		±,22			±.13		*. 05						±,16	±,05			±.16	±, 23	
5dyz		. 19		-,12	09		.19		. 11		. 22		. 21			-,11	. 27			
5dxy	•	±.07		±, 11	±.4 0		±. 16						±.10			±.21	±,15			±.07
6.	±,10		±.16			±,11														
W(5, 6, 7, 8)																				
5dg2		±. 07	±. 07			±.11		±. 08	±. 20	±.07	±,20	±.14	±. 06		±.05	±.12	±. 21	±.09	±.09	±.05
^{5d} xz	.05				±.10			10	±.23	.07	±,15	.14		06	. 10	±.27	±.08	.16	.08	±.13
^{5d} x ² -y ²	±. 05	±.10	±,12				*. 11	±,07					±.36	±,06	±.09	±,10		±,13	±.08	
5dyz	±. 08	±. 06		.06	.07		10	±. 08	08	±.13	. 13	±, 05	15	±.11		12	.10	±.28	±, 12	22
^{5d} xy		±. 08			±,07		±,19	±,11	±.13	±.08	±,07	±,11		±,12	±.0 8	±.20	±. 20	±.08	±, 23	±, 12
68	±. 05	±. 08	±.10	±,25		±.07	±. 10						·							
W(9, 10, 11, 12)																				
5dz2		±.05	±.05	±,05	*, 05	±.06	±,12	±,05	±.08	±.27	±.1 0	±. 09	±.08	±.11	±, 12	±.09	±.06	±, 11	±. 10	±,08
^{5d} xz	06	±. 06	.09		±. 10				±. 19			.17		-,21	. 10	±.06	±.07		18	±.22
5dx2_v2	±,05	±.07	±.08	±,09	±.05	±, 13	±.10			±.08		±, 30		±.12	±.07	±.14	±.05	±.08	±.10	±. 12
5dyz					.06	±. 10		±,19	10	±,14	.11	±. 09	11	±. 06	±.27	-,15			±.1 7	.16
5d _{xy}	±.07				±. 05			±.29			±, 23				±.20	±, 05	±.06	±.08		± , 19
68	±, 13		±.19	±,12		±.15								±,05						

Table XII. Molecular orbitals which belong to the b1 and b2 representations for the W12N surface molecule where nitrogen is bonded at a 2 CN site.

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Molecular Orbital	Ψ_1	Ψ	Ψĵ	Ψio	Ψ3	¥4	¥5	₽ 6	¥7	₩8	Ψ_{11}	Ψ ₁₂	Ψ ₁₃	Ψ_{14}	₩15	₽ 16
Energy (ev)	(-11.30)		(-10.31)		(-10, 16)		(-9. 68)		(-9.51)		(-8, 94)		(-8.81)		(-8.35)	
N																
۷. دور		-51	. 30		02	. 09	. 05		. 08			. 05	02		·11	. 12
2py	-151			30	09	02		. 05		.08	.05			. 02	12	22
W(1)															•	
^{5d} x7		40	24				13		19			22	37		. 11	13
^{5d} yz	40			. 24				13		19	22			.37	. 13	.11
W(2.5)																
Sdz2	±.	16			±.05	±.19	±. 30		±. 05			±.17	±. 08			
^{5d} xz			33				19		22			±. 24	.06		. 10	12
^{∽d} x²-y²	÷.	22	±.07		±. 08	±. 33			±. 11				±,07			
^{5d} yz				31	. 11				. 05	33	09					
• ^{• d} ×y	±. 20				±. 23	±.06		±. 05			±.28			±. 05	±.0 6	±.05
6 n	±.	05	±. 07			±.07	±, 23		±, 21			±. 10	±. 10	±.11	±.07	±.0 9
W(3, 4)																
5d _z 2	±. 16				±.19	±.05		±. 30		±.05	±.1 7			±. 08		
^{5d} x=			. 31			11			33	05		09	05		•	
⁵ d _x 2-y2	±.22			±.07	*. 33	±.08				±.11				±.07 ·		
^{5d} yz		•		. 33				19		22	24			06	. 12	. 10
^{5 d} xy	±.	20			±. 06	±.23	±. 05	±. 23	•			±. 28	±. 11		±. 05	±.06
6.	05			±.07	±.07					±. 21	±. 10			±. 10	±. 09	±.07
W(6, 9)																
54z2					±. 18	±. 11	±.14	±.11			*. 30	±. 27	±.06	±.06	±.19	
Sdxz			- 15		19	09			14	-,14	16		. 10	22	. 14	33
^{5d} x ² -y ²			±.10	±.11		±. 06	±.17	±. 22	±. 14	±. 14			±.17	±. 17		±. 32
^{5d} vz			. 06	15		. 21						16	. 22	10	. 18	.30
^{'ed} xy			±. 09	±.08	±.09	±.05	±.09	±.07	±. 11	±.11	±. 13	±. 11	±.30	±.30	±. 19	
b N			±.07	±.06	±.06		±.21	±.17	±. 14	±, 14	±.10	±.09	±.07	±.07	±. 11	
W(7, 8)																
5dz2					±. 11	±. 18	±.11	±. 14			±. 27	±. 30	±,06	±.06		±. 19
^{>d} xz			. 15	.06	. 21				15	15	.16		. 10	. 22	. 30	18
'ndx²-y2			±. 11	±.10	±.06		±.22	±.17	±. 10	±. 10			±. 17	±.17	±. 32	
'dyz				15	. 09	19			. 05	. 05		.16	22	10	. 33	. 14
^{Sd} xy			±.08	±.09	±. 05	±. 09	±.07	±.09	±. 16	±.16	±.11	±. 13	±. 30	±.30		±. 19
ta n			±.06	±.07		±.06	±. 17	±.21	±. 20	±. 20	±.09	±. 10	±.07	±.07		±. 11

Table XIII. Molecular orbitals which belong to the e-representation for the W9N surface "molecule" where nitrogen is bonded at a 5 CN site.

Figure Captions

- Figure 1. Surface arrays used to model adsorption on different W(100) surface sites.
- Figure 2. Dependence of bond energy on bond distance for WN molecule with and without inclusion of W 5p orbitals.
- Figure 3. Dependence of non-bonding (5p orbital) repulsive energy on distance for the WN molecules.
- Figure 4. Additivity of non-bonded orbital repulsive energy.
 - a) Difference in E(r) for W₂N between calculations including and omitting W 5p orbitals.
 - b) E(r) calculated by pairwise addition of analytic repulsive energy functions.
- Figure 5. Additivity of non-bonded orbital repulsive energy of abbreviated surface arrays.
 - a) Difference in E(r) for W_nN (n = 4, 5) between calculations including and omitting W 5p orbitals.
 - b) E(r) calculated by pairwise addition of analytic repulsive energy functions.
- Figure 6. Bond energies as functions of distances for nitrogen adsorbed on different W(100) surface sites.
- Figure 7. Molecular orbital energy diagrams for the totally symmetric occupied orbitals for nitrogen adsorbed on different W(100) surface sites.

Figure 8. Molecular orbital - energy diagrams for the occupied orbitals belonging to the representations $e(W_9)$, and b_1 and b_2 (W₁₂) for nitrogen adsorbed on different W(100) surface sites.







Figure 2. Dependence of bond energy on bond distance for WN molecule with and without inclusion of W 5p orbitals.



Figure 3. Dependence of non-bonding (5p orbital) repulsive energy on distance for the WN molecules.





- a) Difference in E(r) for W_2N between calculations including and omitting W 5p orbitals.
- b) E(r) calculated by pairwise addition of analytic repulsive energy functions.



Figure 5. Additivity of non-bonded orbital repulsive energy of abbreviated surface arrays.

- a) Difference in E(r) for W_nN (n = 4, 5) between calculations including and omitting W 5p orbitals.
- b) E(r) calculated by pairwise addition of analytic repulsive energy functions.



Figure 6. Bond energies as functions of distances for nitrogen adsorbed on different W(100) surface sites.

 $\overset{\omega}{3}$



Figure 7. Molecular orbital - energy diagrams for the totally symmetric occupied orbitals for nitrogen adsorbed on different W(100) surface sites.

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Figure 8. Molecular orbital - energy diagrams for the occupied orbitals belonging to the representations $e(W_9)$, and b_1 and b_2 (W_{12}) for nitrogen adsorbed on different W(100) surface sites.

APPENDIX C.

A MOLECULAR ORBITAL INVESTIGATION OF

CHEMISORPTION. III. OXYGEN ON TUNGSTEN (100) SURFACE

A Molecular Orbital Investigation of

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Chemisorption. III. Oxygen on Tungsten (100) Surface

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Abstract

The relative bonding energies of oxygen chemisorbed at three symmetric sites on a W(100) surface represented by finite arrays of tungsten atoms¹ were obtained by means of Extended Huckel Molecular Orbital Theory (EHMO). The preferred site for chemisorbed oxygen was the 2 CN site. The selection of this site by an O adatom is consistent with the usual divalent property of oxygen. Based upon this site selection, a surface structure for O adatoms on the W(100) can be proposed for observed (2 x 1) LEED patterns (8).
I. INTRODUCTION

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In applying the Extended Hückel Molecular Orbital (EHMO) technique to the chemisorption of hydrogen¹ and nitrogen² on a tungsten (100) surface, we have found that the preferred site, for these atoms are different. Hydrogen preferred a site where it could most effectively bond to a single surface atom and nitrogen a site where it could interact with five tungsten atoms satisfying its multiple valency. An atom with yet different bonding properties is the next obvious step in the study of trying to develop an overall understanding of chemisorption and sites selected by adsorbates on any surface. In this paper, we have selected oxygen because of its usual divalent properties. This should add an additional test in determining if a metal surface will accept an adsorbate at a site determined by the chemical properties of the adsorbate atom.

II. PROCEDURE

The EHMO procedure has been outlined in our two previous papers.^{1,2} In Table I are listed the orbital exponents for the Slater type orbitals (STO) of the oxygen 2s and 2p orbitals which have been derived from Slater's rules.³ The valence orbital ionization potential

$$VOIP(q) = Aq^2 + Bq + C$$
 (1)

was calculated as a function of atomic charge from the parameters A, B and C which are also listed in Table I. These values were taken from the work of Basch, Viste and Gray.⁴ As in the previous studies,^{1,2} we are reporting the results of the diatomic molecule, WO, and the surface "molecules", W_nO , where the oxygen atom is placed at each of the three symmetric sites of the surface arrays in Fig. 1.

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III. RESULTS

A. Diatomic Molecule

For the diatomic molecule, WO, the bonding energy E as a function of internuclear separation r is plotted in Fig. 2. The presence of the 5p orbitals of tungsten in the basis set produced a minimum in curve a, because the complete occupation of these non-bonding orbitals provide the repulsion at smaller internuclear separation. $ln E_{Rep}$ (difference between curves a and b) is plotted as a function of r in Fig. 3. An analytical exponential term of the form previously indicated¹

$$E = Ae^{-r/\alpha}$$
(2)

can be used to approximate this repulsive energy. The values for the parameters A and α needed to calculate the repulsive energy by this analytical exponential term can be determined from the plot of Fig. 3 $(A_{WO} = 4308.6 \text{ ev and } \alpha_{WO} = .209 \text{ Å}^{-1}).$

From the sum of the single bond covalent radii (0.66Å for O and 1.30Å for W) and the equilibrium bond length for the WO molecule of 1.68Å (Fig. 2), Pauling's empirical relationship⁵ between bond length and bond order gives a value of 2.4 for the bond order. From Table II, it is observed that Ψ_2 , Ψ_3 and Ψ_4 are the most significant bonding molecular orbitals, as was the case for the WN molecule which had a bond order of 3.1. The resulting smaller bond order for the WO molecule can be explained by a comparison of overlap population of the atomic orbitals forming the σ and π bonds of these diatomic molecules. The overlap population in the σ bond for the WN molecule was .69 compared to .48 for the WO molecule and in the π bond for WN molecule, the overlap population was .86 compared to .62 in the WO molecule. The total overlap population for the WN molecule is 1.55 compared to 1.10 for the WO molecule. This reduction in overlap population can account for the reduction in bond order.

It was originally proposed that the computation could be simplified if the non-bonding 5p orbitals which provide the repulsive energy could be omitted from the basis set. In the case of hydrogen¹ this was possible, but it was shown² that nitrogen bonded to larger groups of atoms would not allow the repulsive energy to be calculated by the summation of pairwise interactions. A similar comparison is made for the interaction of oxygen with larger arrays of atoms. The repulsive energies resulting from the presence of the 5p orbitals of tungsten in the basis set are plotted as curves a in Fig. 4. The repulsive energies calculated by the summation of pairwise repulsions

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determined by the analytical exponential term are plotted as curves b. As the case with nitrogen, it is also observed that the repulsive energies can not be properly approximated by the pairwise summation of exponential term when the oxygen is bonded to larger arrays of tungsten atoms. Therefore, it was approximated instead by direct calculation of the reduced arrays of Fig. 1 ($W_5 - 1$ CN and 5 CN sites and $W_4 - 2$ CN sites).

B. Oxygen Chemisorbed on a W(100) Surface

The previous¹ adjustments of the VOIP for the orbitals of the edge atoms of the surface atom arrays (Fig. 1) were used to reduce charge transfer resulting from edge effects. With no further adjustment of the VOIP of the tungsten atoms, the VOIP for the orbitals of the oxygen atom were iterated to charge self-consistency at equilibrium separation at each of the symmetric sites. On the final iteration, the VOIP for the 2s and 2p orbitals of oxygen were: -23.53 ev and -8.77 ev at the 1 CN site, -23.73 ev and -8.92 ev at the 2 CN site and -25.91 ev and -10.64 ev at the 5 CN site. To insure the

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results were not dependent upon VOIP parametrization, ^{1, 2} the calculations were repeated using -23.73 ev and -8.92 ev for the 2s and 2p orbitals of oxygen respectively at all the sites. From these calculations, the bonding energies for oxygen on each of the surface arrays were determined as a function of internuclear separation r and plotted in Fig. 5. The internuclear separation r was the distance to the nearest tungsten atom(s) for the oxygen at each of the sites. The bonding energy E was calculated according to the following relationship:

$$E = \sum_{W_n O}^{n_i} \epsilon_i - \sum_{W_n}^{n_i} \epsilon_i - \sum_{O}^{n_i} \epsilon_i + E_{Rep}$$
(3)

where the first term is the summation over the energies ϵ_i of the occupied molecular orbitals of the surface "molecule" W_nO , the next term is the summation over the energies ϵ_i of the occupied molecular orbitals of the W_n surface "molecule" representing the surface for each site, and the next term is the summation over the energies ϵ_i of the occupied atomic orbitals of the oxygen atom at infinite separation. The repulsive energy E_{Rep} is that determined in prior calculation using abbreviated surface arrays in which the 5p orbitals could be included in the basis set. A summary of the above results where oxygen is at equilibrium separation at each site is found in Table III.

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The only molecular orbitals of interest from the W_n surface arrays are those belonging to the same irreducible representation in the appropriate symmetry groups (W₉ - C_{4v} and W₁₂ - C_{2v}) as the atomic orbitals of oxygen. In Fig. 6, the molecular orbital energy diagrams are shown for the molecular orbitals belonging to the totally symmetric irreducible representation for surface "molecules" with oxygen bonded to each of the surface arrays in Fig. 1. On the sides of each panel in Fig. 6 are the occupied totally symmetric molecular orbitals of the W_n arrays and the atomic orbitals of oxygen (2s and $2p_{z}$). The occupied totally symmetric molecular orbitals of the $W_{n}O$ surface molecules are represented in the center of the panels. The totally symmetric orbitals for the W_q and W_{12} arrays are tabulated in Tables IV and V and can be compared with the tabulated totally symmetric orbitals for the W_nO surface "molecules" (Table VI - 1 CN, Table VII - 2 CN and Table VIII - 5 CN). From these tables, it is observed that the lowest molecular orbital in each surface "molecule" is extensively 2s in character. The other additional molecular orbital formed as a result of the $2p_z$ atomic orbital interacting with the totally symmetric orbitals for each of the W_n arrays (Ψ_7 - 1 CN and 5 CN and Ψ_{13} - 2 CN) increase in energy in going from the 1 CN to the 2 CN to the 5 CN site. The Ψ_7 molecular orbital at the 5 CN is even above the highest occupied molecular orbital (HOMO). Similar behavior was also observed for the interaction of the totally symmetric

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atomic orbital of hydrogen¹ and nitrogen² with the W_n arrays.

The $2p_x$ and $2p_y$ orbitals of oxygen, as in the case of nitrogen,² play a very important role in the selection of the site for preferred bonding. In Fig. 7 the molecular orbital energy diagrams are shown for these atomic orbitals interacting with the appropriate molecular orbitals of the W_n arrays forming the molecular orbitals of the W_nO surface "molecules". A comparison can be made between the tabulated molecular orbitals (Table XI - 1 CN, Table XII - 2 CN and Table XIII - 5 CN) for the W_nO surface 'molecules' with the tabulated molecular orbitals (Table IX - W_9 , Table X - W_{12}) of appropriate symmetry for the W_n arrays. With oxygen at the 5 CN site, the $2p_x$ and $2p_v$ orbitals interact primarily with three sets of degenerate orbitals of the W9 array. The resulting orbitals of the W9O "molecule" are not as energetically favorable as those similar molecular orbitals of the W_qN "molecule".² The influence of these p orbitals of oxygen in selecting the 2 CN site is consistent with the usual divalent property of oxygen.

IV. DISCUSSION

As reported by Madey,⁶ the adsorption oxygen on W(100) surface is a complex process. Oxygen will not desorb from the W(100) surface as molecular oxygen, but as oxides of tungsten and even as atomic

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oxygen.⁷ Estrup and Anderson⁸ reported an oxygen (2×1) LEED pattern formed by heating a streaked (4×1) structure formed by room temperature of oxygen.

Our calculations indicate that oxygen prefers the 2 CN site. The divalent property of oxygen has predominated in selecting the bonding site. In Fig. 5, the equilibrium internuclear W-O distance (2.05 Å) is comparable to the sum of the W metallic radius (1.37 Å) and the O covalent radius (.66 Å). The bonding of oxygen at the 2 CN sites can be effectively described as two single bonds to each tungsten atom at this site. If the oxygen atoms were bonded on the W(100) surface at 2 CN sites as depicted in Fig. 8, each oxygen atom would be bonded to two tungsten atoms and these two tungsten atoms would not be bonded to any other oxygen atom. Such a surface structure of adsorbed oxygen would account for the observed (2×1) LEED pattern.

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Orbital	Orbital Exponent	A [ev/(e) ²]	B [ev/e]	C [ev]
2s	2.275	3.46	22.89	32.33
2 p	2.275	3.46	18.57	15.80

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Table I. Orbital exponents for STO of the oxygen atom and parameters needed to calculate VOIP as a function of atomic charge.

W-0	Ψ_1	Ψ_2	Ψ_3	Ψ_{4}	Ψ_5	Ψ_6
Energy (ev)	(-27.21)	(-12.50)	(-12.50)	(-11.95)	(-9.80)	(-9.80)
0						
2s	.93			.10		
$2p_z$. 84		
² p _x			81			
2py		81				
w						
5d _z 2	.13			35		
^{5d}xz			44			
$5d_{x^2-y^2}$					1.00	
^{5d} xy						1.00
6s				.18		

Table II. Occupied Molecular Orbitals for the diatomic molecule WO.

Site	Bond Energy (ev)	re (Å)	Charge e
1 CN	-1.06	1.93	 45
2 CN	-1.30	2.05	40
5 CN	80	1.98	±. 90

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Table III.	Energies,	equilibrium	distances,	and charges for	oxygen
	bonded to	different W(100) surface	e sites.	

Molecular Orbital Energy (ev)	Ψ ₁ (-12.43)	¥2 (-10.69)	Ψ ₃ (-10.38)	Ψ ₄ (-9. 77)	Ψ ₅ (-8.47)	
W(1)						
$5d_{z^2}$		54		10	i9	-
6s	24					
W(2, 3, 4, 5)						
$5d_{z^2}$. 22	.17	.16		
$5d_{xz}$ (2,5)		±. 15		±. 31	±.07	
$5d_{x^2-y^2}$		±.12	±. 30	±. 19	±. 05	
5d _{yz} (3, 4)		±.15		±.31	±.07	
6s	18					
W(6, 7, 8, 9)						
5d _z 2		10	19		.26	
5d _{xz}			±.11		±. 28	
5d _{yz}			±.11		±. 28	
^{5d} xv		±.12		±.22	±. 10	
6s	12					

Table IV. Totally symmetric molecular orbitals for the surface arrays representing the 1 CN and 5 CN sites.

	ψ1	ψ ₂	\$ 3	¥'4	ψ5	ψ6	ψ7	ψ8	¥9	ψ ₁₀	ψ ₁₁
	(-12.78)	(-11.31)	(-10.87)	(-10.82)	(-10.25)	(-10.03)	(-9.68)	(-9.62)	(-8.97)	(-8.72)	(-8,32)
W(1,2)											
5d-2			36	09		08	.09	1 5			. 15
$5d_{y}2_{-y}2$.33			19	06	-,05	08	30	14	.08
5 d _{y 2}		±. 12	±. 08	±. 32	±.21		±. 17	±. 18	±.1 0	±.12	
6s 92	.18										
W(3,4)											
5d.2			.36	09		08	09	1 5			15
5d		±.12	±.08	±.32	±.21	• • •	±.1 7	±. 18	±. 10	±.12	
5d, 2, ,2		33			19	.05	.05	+.08	30	.14	.08
6s 6s	.18										
W(5,6,7,8)											
5d_2			. 12	.05	17	.07	.08	.06	.08	12	.11
5 d _w z			±.05	±. 07	±.14	±.23		±.09			±.26
$5d_{x}2 - v^{2}$		13	.05	11	.10	16	.05		20	05	14
5d _{vz}						±. 07	±.1 3	±.1 9	±.1 4	±. 24	±. 10
5dxv ·				±. 12		±. 08	±. 24	±. 18	±. 09	±.1 8	±. 08
6s	.10										
W(9, 10, 11, 12)											
5d2			.12	±. 05	.17	.07	.08	.06	.08	12	11
5dyz			• -	-		±.07	±. 13	±. 19	±. 14	±. 24	±. 10
$5d_{x}^{2} - v^{2}$.13	±. 05	.11	.10	.16	05		±. 20	.05	14
$5d_{vz}$			±. 05	• ±.07	±.14	±. 23		±. 09			±. 26
$5d_{xy}$				±. 12		±. 08	±.24	±. 18	±. 09	±. 18	±. 08
65 ⁷	.10										

Table V. Totally symmetric molecular orbitals for the surface array representing the 2 CN site.

Molecular Orbital	${}^{\Psi}6$	Ψ'_1	Ψ'_2	Ψ'_3	$\Psi_4^{'}$	Ψ_7	$\Psi_5^{'}$
Energy (ev)	(-23.98)	(-12.23)	(-10.49)	(-10.37)	(-9.78)	(-9.23)	(-8.42)
0							
2s	. 95	. 09	.14	05	.03	05	
2p _z		. 02	31	.11	18	. 78	16
W(1)							
$5d_z^2$.10	.17	37	.10	09	.15	.07
6s	.07	20	.06				
W(2, 3, 4, 5)							
$5d_{z2}$. 25	.12	.14	. 20	08
5d _{xz} (2,5)			±.14	±.09	±.32	±. 05	
$5d_{x^2-y^2}$			±.05	±.32	±.1 7	±.1 5	±. 10
5d _v 2 (3, 4)			±.14	±.09	±.32	±. 05	
6s		18	07				
W(6, 7, 8, 9)							
$5d_{z^2}$			13	16		09	25
⁵ d _{xz}				±.11		±. 06	±. 28
5d _{vz}				±. 11		±.06	±.2 7
5d _{xy}			±.11	±. 06	±. 22		±. 12
6s		12					

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Table VI. Totally symmetric molecular orbital for W₉O surface "molecule" where oxygen is bonded at a 1 CN site.

Molecular Orbital	Ψ12	Ψ_1'	¥2'	¥3	Ψ4	₩5	₩'6	₩'8	¥'7	¥ģ	Ψ10	¥ ₁₃	Ψ.1
Energy (ev)	(-24.07)	(-12.49)	(-11.32)	(-10.92)	(-10. 63)	(-10.08)	(-10.02)	(-9.71)	(-9.59)	(-9.27)	(-8.72)	(-8, 53)	(-8.26)
0		_											
2:	93	. 11	. 03	. 02	07	.10	06	~.03	09	. 03	02	. 02	. 02
2pg	01	02	. 11	17	03	.31	09	32	29	. 45		32	. 21
W(1, 2)													
5d_2			. 05	37			.07	14		05		06	08
5d_2_v2		08	.31		.06	.14		08		. 15	. 09	. 12	08
5dys	±.07	±. 12	±.16		±,14	±.11	±. 09	±.09	±. 21		±.13		±.06
60	*. 06	14											
W(3, 4)													
5d_2				.32	. 16		. 10	14	12				. 18
5d _{ma}			±. 14	±.11	±.36			±.24	*. 09	±.07	±.09	±.14	±.09
5d_2_v2			32	08	.11	. 11	14			. 20	20	21	16
68		18											
W(5, 6, 7, 8)													
5ds2				. 11		. 14	18				. 12		15
5d <u>**</u>						±.24	±. 11			±.17		±. 30	±. 12
5dx2_v2			12		. 10		. 21	•	. 11	. 20			.17
5dyz			±. 06	±. 05		±. 07		* . 12	±.17	±, 17	*. 25		±.10
5day				±. 05	±. 10		±.07	*. 19	±, 23	±.15	*, 15		
68		12									.05		
W(9, 10, 11, 12)													
5d ₂ 2				10	12	17			05	.09	. 10	.16	
5d xs			±.06				±. 05	±.23	±.08		±.21	±, 13	±.15
5dx2-y2			. 12	.08	14	11	11	07		. 12	10	. 22	. 07
5dyz				±.06	±. 10	±. 06	*. 26	±.07					±.31
5dxy					±.14	±, 15		±. 10	±.24		±.20	±.07	*. 12
6e .		10											

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Table VIL. Totally symmetric molecular orbitals for the W12O surface "molecule" where oxygen is bounded at a 2 CN site.

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Molecular Orbital	Ψ_6	$\Psi_1^{'}$	Ψ'_3	$\Psi_2^{'}$	$\Psi_{4}^{'}$	Ψ_5'	Ψ_{7}
Energy (ev)	(-24.15)	(-12.01)	(-10.49)	(-10.18)	(-9.38)	(-8.62)	(-7.59)
0							
2s	.91	09	.04		. 15	08	07
2p _z		. 07	. 25	45	.56	.04	. 23
W(1)							
5d _{z2}	.07	25	23	.04	33	05	27
6s	.04	.21			04		07
W(2, 3, 4, 5)							
5d _z 2		.06	.24		11	08	22
$5d_{xz}$ (2,5)			±.07	±. 31	±.11		±.21
$5d_{x^2-v^2}$	±. 04	±. 10	±. 20	±.1 5		±. 06	±. 17
5d _{yz} (3,4)			±.07	±.31	±.11		±.21
6s	. 03	.14					
W(6, 7, 8, 9)							
$5d_{z2}$			20	07	.07	.24	
$5d_{xz}$			±.09		±. 06	±. 29	
^{5d} yz			±. 09		±.06	±. 29	
⁵ d _{xy}				±.16	±. 25		±. 36
6s		. 13			09		.05

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Table VIII. Totally symmetric molecular orbitals for the W_9 O surface "molecule" where oxygen is bonded at a 5 CN site.

гер	resenting the 1 C	N and 5	CN site	s.				
Molecular Orbital	$\underline{\Psi_1}$ $\underline{\Psi_2}$	¥3	¥4	¥5 ¥6	Ψ ₇ Ψ ₈	$\Psi_9 \Psi_{10}$	$\Psi_{11} = \Psi_{12}$	Ψ_{13} Ψ_{14}
Energy (ev)	(-10.68)	(-	10.17)	(-9.71)	(-9.58)	(-9.24)	(-8.91)	(-8.80)
W(1)								
^{5d} xz	50 .10			.21	. 19	.06 .08	19	0737
^{5d} yz	.10 .50			21	. 19	08 .06	19	.3707
W(2.5)				•				
۶dz2	±.16	±.13	±.15	±.26	±.11	±.14 ±.18	±.10	±.07
5d _{xz}	23 .05		- 05	. 13		1722	29	. 10
5dx2-y2	±.12	±.17	±.21	±.]4	±.26	±.17 ±.22	±.07	
5dyz	17	.16	13	16	. 35		08	. 05
^{5d} xy	±.22	±.19	±.15	±.07			±.32	±.07
6.		±.07	±.08	±.20	±, 20	±.12 ±.16		±. 08
W(3, 4)								
5dz2	±.16	±. 15	±.13	±.26	±.11	±.18 ±.14	. ±. 10	±.07
5d _{x2}	. 17	. 13	.16	.16	.35		08	05
^{5d} x ² -y ²	±.12	±.21	±. 17	±.14	±. 26	±.22 ±.17	±,07	
^{5d} yz	.05 .23	.05		13		.2217	- 29	10
^{5d} xy	±.22	±. 15	±. 18	±.07			±.32	±.07
6.		±.08	±.07	±.20	±, 20	±.16 ±.12		. 08
W(6, 9)								
۶dz2		±.21		±.14 ±.10			±.26 ±.32	±.08 ±.11
^{5d} xz	.0809	08	.19	07	.18 .09	. 09	.0911	21 .11
^{5d} x ² -y ²	±.09 ±.06		±.11	±.17 ±.24		±.18	±.10 ±.08	±.17 ±.12
⁵ dyz	.0611	05	20	06 .05	.19	. 10	13 .07	. 24
^{5d} xy	±.08 ±.10	±.07		±.08 ±.05		±,26	±.09 ±.11	±.25 ±.37
t s		±.10		±.19 ±.14	±.15 ±.18	±.23	±.05	±.05 ±.08
W(7,8)								
4,2			±.21	±.10 ±.14		±.05	±.32 ±.26	±.11 ±.08
"dar	.11 .06	. 20	05	.05 .06	. 19	. 10	.07 .13	. 24
hdxd-yd	1,06 ±,09	*.11		±.24 ±.17		±.18	±.08 ±.10	±.12 ±.17
""Jyr	-,09 -,08	. 19	, Он	-,07	.09 .18		.11 .09	1121
'''' xy	+.10 ±.07		±.07	±.05 ±.08		±.26	±.11 ±.09	±.37 ±.25
6 m			*.10	±.14 ±.19	±.18 ±.15	±.23	±.05	±.08 ±.05

Table IX. Molecular orbitals which belong to the e representation in the C_{4v} symmetry group for the W_0 surface arrays representing the 1 CN and 5 CN sites.

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Istia X - Muletular urbitals which belong to the by and by representations in the Gey symmetry group for the Wild surface areay representing the 2 CN site.

 Misler slær Grissigi	¥1	¥.·	···	¥.		۴	· ¥7	¥,,	 ¥ŋ	Ψ ₁₀	¥11	Ψ ₁₂	¥13	¥14	¥ ₁₅	¥16	¥17	¥18
Findgy last	1.10, 12	1.640.921	(10. AN)	с. за, яну	1-10.54	1-10,56)	(-10, 01)	(-10 01)	(.9, 80)	(-9, 80)	(-9, 59)	(-9, 59)	(-9.38)	(-9. 38)	(-9.07)	(-9.07)	(-8. 62)	(-8.62)
50.2	e, 1n	•. 15			s. 10				s. 18	±. 19	+. 09	£.07	*. 10	*. 08			±. 14	±. 05
		21	-, 06			-,04	10		. 17	15			. 20	24	13			98
··· · · · · · · · · · · · · · · · · ·	1.17	1.14	1.05	1.04	+. 11			+. 05					* . 12	±.10			* . 27	±. 10
"Hy #					21	05	. O.H	21			. 14	. 11	19	16			. 05	
" day	• 12	+ 14	1.14	*, 0K	1, OM	•. 17							±. 09	±. 11	±. 20			#. 11
f				•.1#	4. OH													
W.1 A																		
74-2	4, 15	• 18				a 10			1.19	#. 18	*. 0 7	s. 09	£. 08	a. 10			±. 05	±. 14
					. 05	- 21	. 21	. 0.8			, 11	14	. 16	19				05
54. S	•. 14	+ 17	e. n.e	6,05		•. 11	4.05						. ±. 10	±. 12			±. 10	·±. 27
	23	17		"4	•, ан			. 10	. 15	. 17			24	20		13	. 08	
· · ·	+.14	•.12	1. IM	a. 14	8. 17	= 0N							±.11	±.09		±.20	±. 11	
			•. 1 *	±, 09		•.08												
W/rS. Ht																		
54.2	. 09					#. 12	. 07		s 27	# 16	8.09	. 08			±.09	±. 14	±. 14	
94					-, 11			1.25	-	. 04		28	. 11	. 09	14	27	18	08
N4-22		4. 17			1.11		1. ON				•. 21	#. 1H	±. 1 8	*. 12	1.14	±.04		#. 13
54		. 15					. 09		. 08	z. 16	. 18	. 15	10			13	24	18
7- 14-	#, GM				∎. 07			+, 10	s. 07	±.09	a. 08	±. 08		±.20·	±. 07	±.18		*. 1 5
fin.			4.21	0,16	s. 12													
W(h, 7)																	. 13	
54,2		÷ 04			a, 07	± 10	*.09	• •	#. 14	€ 28	# ()h	*. 11			*. 10	*. 13	±. 12	E . 07
***					. 10	08	17	. 16	z. 08			16	14	09	17	. 20	. 09	10
³⁴ ≝2-y2 5.1	•, 17				4.10		. 06	. 07	- 17	0.6	19	13	.	* 10	a. 15 06	- 12	- 07	29
¥π Sal					. 09		• 19	• 24	. 09	*.06	* 05	• 09	* . 19			*. 19	*. 08	* 13
(.a.		•,		4.28	1, 10								,					
-																		
w(9, 12)					. 10			00				- 04				A 10	a 07	• 13
·****	1.04	15					07	. 07		17		. 19	10	- 15	- 12	- 06	2.0.	
		. 12				• 10		• DB	. • •		* 15	• 23	*. 16	* 13	1.11	±. 13	*. 10	±. 09
∎ - y* Nel _						10	. 16	17		•. UH	. 07	.17	09		26	17	. 18	. 09
	•. d#					*. 04	4.21	. 19	. 06	a. D*/	. 09	• • •			#. 19		x . 13	a . 08
••• •••	-		•		8, 134	s, 10				-								
w.t. 11																		
دهه به برور به د ما		. 84			+. 12			1.07	#. 16	1.27	• 08	±.09			z , 14	±. 09		±, 14
	. 15							.09	16	OH	15	18		10	13		18	. 24
14.2	+.17					*. 22		#, DH			#, 3H	*, 21	±, 12	±. 18	a, 09	#, 14	±. 13	
he he he						. 11	23		09		. 18		. 09	11	.27	14	. 08	18
heley.		• 11×6				± 07	#. 10		*. 09	*. 07	±. 0X	#. 0R	. 20	-	*. 18	07	*. 15	
1			\$ 16	4.23		a.12												
	•																	

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Molecular Orbital	¥'ı	¥2	Ψ3	₩4	¥s	¥6	⊎ '7	₩8	\$9	¥10	¥,11	¥,12	¥13	¥14	₽ ₁₅	¥16
Energy (ev)	(-10	(-10.78)		(-10, 18)		(-9. 72)		(-9.69)		(-9.25)		(-9.22)		(-8. 90)		. 32)
0																
2p _x	. 22			06	19	17	-, 25	- 07		. 19	. 50	05	.06	03	54	34
2p		. 22	06	5	. 17	19	07	-, 25	. 19		05	50	~.03	06	.34	54
₩(1)																
5d	53				.21	. 19				. 05	19		.06		21	13
5dys		. 53			19	.21			. 05			. 19		06	. 13	. 21
W(2, 5)																
5d ₈ 2 '	*.15			±.20	A. 13	*.11	a. 21	a. 06		*. 25			*.11	s. 06	a .06	
54 ₂₃	22			. 05		-	12			Z3	. 18		. 28	15	07	
5dx2-y2	n. 12			*. 27	*, 15	±. 13				#.34	±. 12				±. 10	±. 06
5dys		15	18		24	. 27	. 05	.17			•	08		06	. 05	08
Sday		±. 20	±. 25	±. 05				a. 06	±.11			±. 16	±.13	±.24		
66				*. 10		•	*. 26	±. 07		±. 17	±. 12		*. 05		±. 09	±.06
W(3,4)																
5d_2		÷. 15	a. 20		±. 11	±. 13	a. 06	a. 21	±. 25				a. 06	±, 11		*. 06
5d ₂₂	15			18	. 27	.24	. 17	05			. 08		. 06		.08	05
5d_2_v2		±. 12	±.27	s. 05	a. 13	a.15			a. 34			±. 12			±. 06	±.10
Sdyg		. 22	. 05					12	23			18	15	28		07
54	a. 20		±.05	±. 25			6.06			±.11	±.16		±. 24	±.13		
60 ^{-'}			±. 10				±, 07	a. 26	±. 17			*. 12		±. 05	±. 06	*. 0 8
W(6, 9)																
5d _{g2}			±. 12	s. 17		*. 09	*, 14	±. 08					±.12	±.41		±. 05
5d ₂₈	. 08	07	. 20	05	.07	.15	. 10		. 11	. 11	. 08	09	-,14	14		. 18
^{5d} x2-y2	s. 07	a. 06	*. 08	±. 06	*. 29		a. 06	· #. 10	±.17	±. 13	±. 05		*, 05		*.22	*. 05
5dys	.06	09	11	. 17	05	. 15		. 10	.07	.13	.07	09	. 19		.07	. 17
5dxy	±.07	±.08	±.05	±. 07			±. 08		±.08	±,11	*. 26	±. 31			±.07	*.33
60			n , 06	*_ 09		±. 05	n. 28	a. 16	±. 12	a. 15	±. 08	*. 10		±. 09		*. 09
W(7, 8)																
5d _{a2}			±. 17	*. 12	a. 09		4. 08	n. 56					*.41	*. 1Z	±. 05	
5d ₂₈	. 09	.06	17	11	. 15	. 05	. 10		13	.07	. 09	. 07		.19	.17	±. 07
5dx2-y2	* . 06	±. 07	*. 06	*. 08		±. 29	±. 10	a. 06	a. 13	±. 17		±. 05		±. 05	±. 05	±. 22
5dys	07	08	05	20	15	. 07		.10	. 11	11	09	08	14	.14	18	
Sday	s. 08	±.07	±.07	±. 05				*. 08	±.11	*. 08	±. 31	±. 25			±. 33	±. 07
60			a. 09	a. 06	±. 05		a. 16	a. 28	a. 15	a. 12	±. 10	*. 08	±. 09		±. 09	

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Table XL Molecular orbitals which belong to the e representation for the WgO surface "molecule" where oxygen is bonded at a 1 CN site.

Molecular Orbital	¥1	₩2	¥3	₩4	¥۶	¥ 6	¥7	¥8	* 9	♥ ₁₀	¥11	₩ 12	¥ ₁₃	¥ ₁₄	¥15	¥16	¥ ₁₇	¥ ₁₈	♥ 19	* ₂0
Energy (ev)	(-11.04)	(-11.02)	(-10, 90)	(-10, 88)	(-10. 67)	(-10.59)	(-10,19)	(-10, 01)	(-9.83)	(-9. 83)	(-9.66)	(-9.60)	(-9, 52)	(-9. 42)	(-9.21)	(-9.07)	(-8.87)	(-8.68)	(-8.54)	(-8, 53
0																				
2pz		. 26	03			09		. 05		. 16		.08		-, 24	. 37			. 49	. 55	
2py	24			.01	-, 23		-,31		.14		23		. 20			02	. 50			28
W(1, 2)																				
5dg2		*. 23			A ,13		±. 11		±.24		±.1 0		±.14				±.07			±. 10
5d _{xs}	21		. 19			. 09		. 12		. 23				29				12		
5d_2_v2		±.21		±, 12		•	*, 13		*. 06				±.07				#. 24			±. 20
5dys		16		. 05	. 28		. 21				. 17			12			. 13			
5day	*. 28		a. 05			*. 33		*. 05				±. 07			* , 19				± , 15	•
6a				±. 20	±. 06		* ,05													
W(3, 4)														٠						
5d_2	* , 21		*. 09			*. 06		\$,05		*,2 7		*. 12		\$, 10	\$.0 9				±.1 9	
5das			.07			. 23		24				16		27					.06	
5d_2_2	#.11		* , 21			#, 13		± ,06						*. 16				±. 22	±.18	
5d		-, 24			•. 39		17		. 15		. 16		-, 24			. 12	. 25			
5day		*. 10		±.15			±,13									± ,21	±.17			±, 05
6.	±. 09		±. 17			±, 10														
W(5, 6, 7, 8)																				
5d_2		±.08	±. 07			±, 11		±.09	±. 24	±.08	±, 15	±, 13	*. 06		±.07	±, 12	#.20	±,05	±.10	
5d	05				#. 10		*.07	. 10	. 19	05	±,16	15	*. 07		. 11	* , 27	±.08	-,16	. 05	±, 12
5dx2.v2	±.06	±.12	*. 11		* , 05		*. 07	*. 08			±.07		±. 36		*, 11	±. 10		±.14		
5d	±.09	.08	±. 05		. 05		. 10	*. 08	09	±.14	. 18		. 10	±. 08		. 12	. 07	*.30		23
5d _{xy}		±. 08			*. 07		±,23	±, 11	±.08	±.10	±. 06	±.11		±,14			±.18		±,23	±.13
6.	*. 05	±. 05	*.11	±.25	±. 07	±.07	*. 07													
W(9, 10, 11, 12)	•																			•
5d_2	±. 05	*. 06		±,05	±.07	±.0 6	±, 12		±. 08	±.29	±.11	±.05	#. 0 7	*. 08	±. 13	±,09	\$.06	±, 12	±.08	±.09
5d _{ya}	. 07	±.08	08		*.0 9				4.19	-,05		21		-,15	.11	±.06	*, 11	. 07	16	±, 20
5d_2_v2	±.07	±,10	±.07	±. 07		±.1 3	±.10			±.05		±. 29		±.19	, ·	*, 14	±,06		¥.09	±, 11
Sd _{va}					~. 05	* , 10	05	*, 20	•.09	*, 12	. 14	±.09	.08	±,08	±.27	. 15	,		±.16	. 17
5d.v	±.08					±.06	±. 06	±.28			±.21		±.0	s ±.06	±.20	±.05		* ,07		±. 19
63	±.12		±. 20	±, 13		±,14														

Table XIL Molecular orbitals which belong to the b1 and b2 representations for the W12O surface "molecule" where oxygen is bonded at a 2 CN site.

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Molecular Orbital	¥1	¥₂	* 5	*6	¥'3	*	*,	*'8	¥,	¥10	\$'11	¥'12	Ψ'B	¥14	Ψ_{15}^{+}	¥ 16
Energy (ev)	(-10. 98)		(-20.19)		(-10.15)		(-9. 68)		(- 9	. 50)	(-8. 93)	(-	8.80}	(-8.34)	
0																
20-	39		23	. 12		31	- 06			11	04	. 05	02	02	16	. 14
2py		39	. 12	. 23	31			.06	.11		. 05	04	. 02	0Z	. 14	. 16
W(1)																
5d ₂₂₈	47		. 10	05			14			. 18	. 13	17	23	28	. 14	12
5dys		47	05	10				14	-, 18		17	13	. 28	23	12	14
W(2, 5)																
54 ₂ 2	±.17		±. 12	a. 06		s. 14	±. 30			±.06	a. 10	±. 13	s. 05	*. 06		
Sd _{XX}	10		. 21	11		18	19			. 23	. 15	19		. 05	. 12	10
5d2-y2	a. 21		*. 06			a. 36				a. 09				±.05		
5dym		. 08	.16	. 29	07				32			+. 05				
54xy		*. 22	±. 07	s. 14	s. 18			4,05			±.23	±.1 8	±. 09	±. 07	a. 05	*.06
6e			a. 10	a, 06		•	±, 23			4, 23	n , 06	±. 07	a. 06	*.08	#. 08	± , 07
₩(3, 4)																
5d _{z2}		a. 17	±. 06	e, 12	±.14			±. 30	a. 06		a. 13	a. 10	a. 06	±. 05		
5d <u>_2</u>	-08.		29	.16		- 07				. 32	. 05	07				
5dx2_72		±. 21		a , 06	±. 36				±. 09				#. 05			
5dys		10	11	21	.18			19	23		19	15	05		10	1Z
5dxy	e. 22		a. 14	a. 07		s. 18	*. 05				±.18	#. 23	*. 07	±.09	*. 06	#. 05
6e	•		±. 06	a , 10				±. 23	*. 23		±. 07	*. 06	#. 08	±. 06	±.07	≜. 08
W(6, 9)																
5d_2				±,14	±,12	*. 10	a. 13	a. 11		•	±. 05	±. 40		a. 09		±. 1 8
5dga	. 05	. 05	17		20					.14	14	07	11	. 21	. 32	14
5dx2_y2	n. 05	*. 05	s. 15				±. 18	±. 21	*. 11	a. 13	a. 05		±. 24		a. 32	
Sd ym	. 04	. 05	. 14	. 10	07	. 19		•	14		. 12	10	. 06	. 23	30	18
5day	*. 05	±. 05			±,11	a. 09	4. 09	±.07	s. 15	s. 13		*.16	A. 05	±. 42		*. 19
60				a. 12			±.21	s.1 7	* 19	a. 16		±. 12		a. 10		±. 10
W(7.8)																
5d_2			m. 14		±.10	a. 12	s. 11	±.1 3			a. 40	a. 05	±. 0 9		a. 18	
5d _{an}	. 05	04	10	.14	. 19	. 07				. 14	. 10	. 12	. 23	06	.18	30
5dx2_y2	a. 05	a. 05		4. 15			±.21	s. 18	=. 13	a , 11		a. 05		a. 24		±.32
5dys	05	. 05		.17	. 09	20			14		07	. 14	21	11	14	32
5d xy	*. 05	*. 05				±.11	±. 07	a. 09	±. 13	±. 15	±. 16		±. 42	a. 05	s. 19	
64			a. 12				s, 17	±.21	a. 16	±.19	±. 12		±. 10		±. 10	

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Table XIII. Molecular orbitals which belong to the e representation for the WgO explace "molecule" where oxygen is bonded at a 5 CN site.

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Figure Captions

- Figure 1. Surface arrays used to model adsorption on different W(100) surface sites.
- Figure 2. Dependence of bond energy on bond distance for WO molecule with and without inclusion of W 5p orbitals.
- Figure 3. Dependence of non-bonding (5p orbital) repulsive energy on distance for the WO molecules.
- Figure 4. Additivity of non-bonded orbital repulsive energy of abbreviated surface arrays.
 - a) Difference in E(r) for W_nO (n = 4, 5) between calculations including and omitting W 5p orbitals.
 - b) E(r) calculated by pairwise addition of analytic repulsive energy functions.
- Figure 5. Bond energies as functions of distances for oxygen adsorbed on different W(100) surface sites.
- Figure 6. Molecular orbital-energy diagrams for the totally symmetric occupied orbitals for oxygen adsorbed on different W(100) surface sites.
- Figure 7. Molecular orbital-energy diagrams for the occupied orbitals belonging to the representations $e(W_9)$, and b_1 and b_2 (W_{12}) for oxygen adsorbed on different W(100) surface sites.
- Figure 8. Suggested structure for oxygen adsorbed on the W(100) based upon an oxygen adatom preferring a 2 CN site.



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Figure 2. Dependence of bond energy on bond distance for WO molecule with and without inclusion of W 5p orbitals.



Figure 3. Dependence of non-bonding (5p orbital) repulsive energy on distance for the WO molecules.



Figure 4. Additivity of non-bonded orbital repulsive energy of abbreviated surface arrays.

- a) Difference in E(r) for W_nO (n = 4, 5) between calculations including and omitting W 5p orbitals.
- b) E(r) calculated by pairwise addition of analytic repulsive energy functions.



Figure 5. Bond energies as functions of distances for oxygen adsorbed on different W(100) surface sites.



Figure 6. Molecular orbital-energy diagrams for the totally symmetric occupied orbitals for oxygen adsorbed on different W(100) surface sites.



Figure 7. Molecular orbital-energy diagrams for the occupied orbitals belonging to the representations $e(W_9)$, and b_1 and b_2 (W_{12}) for oxygen adsorbed on different W(100) surface sites.



Figure 8. Suggested structure for oxygen adsorbed on the W(100) based upon an oxygen adatom preferring a 2 CN site.

APPENDIX D.

COMPUTER PROGRAM FOR EXTENDED

HUCKEL MOLECULAR ORBITAL (EHMO) CALCULATIONS

C MAIN PROGRAM TO CALCULATE THE GVERLAP MATRIX RGALD HOFFMANN CHEMISTRY DEPARTMENT HARVARD UNIVERSITY CAMBRIDGE C THIS PREGRAM CONSTRUCTS AN OVERLAP MATRIX FOR UP TO 140 ORBITALS O С UP TO 50 ATOMS. THE ORBITALS MAY BE OF ANY PRINCIPAL QUANTUM NUMB С ANGULAR MCMENTUM S P D OR F AND THERE IS NO LIMITATION ON THE C NUMBER OF ORBITALS OF ANY TYPE EXCEPT THAT THE TOTAL MUST BE LESS С C THAN 14C. THE PROGRAM USES SEVEN FILES 5= INPUT 6=OUTPUT 7=PUNCHOUT С 3,4,8,9 ARE INTERMEDIATE FILES С RESTRICTION - NO MORE THAN 5 DIFFERENT TYPE ORBITALS ON ANY ATOM С RESTRICTION DIFFERENT CRBITALS ON SAME CENTER MUST HAVE DIFFERENT С INFUT CONSISTS OF THE FOLLOWING С С FIRST CARD IN FORMAT (712, F14.0) С N, MS, MC, MMM, MCRB, MK, CON С N IS THE NUMBER OF DIFFERENT ATOMS THE OTHER INTEGERS ARE CONTROL PARAMETERS C MK=0 USES ARITHMETIC MEAN WOLFSBERG HELMHOLTZ FORMULA С С MK=1 USES DIRECT PROPORTIONALITY OF HAMILTONIAN TO OVERLAP С MK=2 USES GEOMETRIC MEAN WOLFSBERG-HELMHOLTZ FORMULA MCRE SPECIFIES AN ORBITAL NUMBER FOR WHICH A POPULATION ANALYSIS С IN ADDITION TO THE GENERAL ONE WILL BE CONE IF MORE=0 JNLY TOTAL POPULATION ANALYSIS WILL BE PERFORMED С С С MS=1 PUNCHES OUT S С MC=1 PUNCHES OUT C MMM=1 BYPASSES PCPULATION ANALYSIS С MS GREATER THAN 1 PRODUCES TRACE PRINTOUT C C MR GREATER THAN ZERD, AN EXPONENTIAL REPULSIVE ENERGY WILL BE ACCED TO THE TOTAL ENERGY С SECOND CARD IS A TITLE CARD IN FORMAT(72H) THEN THERE FOLLOW 2N CARDS, TWO FOR EACH ATOM C С FOR EACH ATOM THE FIRST CARD IS IN FORMAT (13, F7.3, 3F10.6, F12.3, С C F6.3) THE SECOND CARD IN FORMAT (5(211, F5.3, F6.2)) С CN THE FIRST CARD ONE READS IN NORB(1) = THE NUMBER OF DIFFERENT TY С С CRBITALS - ORBITALS ARE DIFFERENT IF THEY HAVE DIFFERENT N AND L THE NEXT ENETRY ON THIS CARD IS THE CHARGE ON THE ATCK CORE С THE NEXT THREE ENTRIES ARE CARTESIAN COORDINATES X, Y, Z OF THE ATOM ON THE SECOND CARD AS MANY 13 CHARACTER SPECIFICATIONS ARE READ IN С С С ARE PROVIDED BY NORB(I) ON THE FIRST CARD EACH SET SPECIFIES AN С FIRST ITS N AND L QUANTUM NUMBERS, THEN THE SLATER EXPONENT, THEN TH С CCULOMB INTEGRAL THE LAST CARD IN THE DATA IS IN FORMAT(7211) C С THIS READS IN THE OCCUPATION NUMBERS OF THE VARIOUS MOLECULAR OREI С STAFTING FROM THE MOST BONDING UP TO THE ANTIBONDING GNES С THE CREERING FOR P LEVELS IS Z X Y C THE ORDERING FOR Ð LEVELS IS Z******2 XZ X**2-Y**2 YZ XY C THE ORDERING FOR Ρ LEVELS IS ZX C THE ORDERING FOR D LEVELS IS Z**2 XZ X**2-Y**2 YZ XY CNE HAS NO CHOICE AT AL ABOUT THE DIRECTION IN WHICH THE BASIS С C ORBITALS ARE POINTING THEY ARE ALWAYS DEFINED WITH RESPECT TO TH C ABSOLUTE MOLECULAR COORDINATE SYSTEM С ALMOST ALL THE COMMUNICATION TO SUBPROGRAMS IS DONE THROUGH COMMON AS LSUAL THE NAMES OF THE SUBROUTINES HAVE LITTLE OR NO RELATION TO THEIR FUNCTION C C

ссс с с с с с с с с	ONE SET OF DATA MAY BE PLACED IMMEDIATELY FOLLOWING ANOTHER THERE IS A LIMITATION IN THE DIMENSIONING OF AFUNCT AND BFUNCT SC THAT JMAX=N1+N2+L1+L2+1 MUST BE LESS THAN 20 FOR THE HIGHEST PCSSIBLE COMBINATION OF THESE ARGUMENTS THIS CAN BE CHANGED BY EXPANDING THE DIMENSION STATEMENT IN AFUNCT AND BFUNCT FOR A ANC B BUT ONE MUST BE CAREFUL IN DCING THIS, SEE NOTE AT BEGINNING OF B FUNCT DIMENSION XX(15C),YY(15O),ZZ(15O),AR(15O),AL(15O) DIMENSION NOR(50,5),SLA(50,5),HII(50,5),NO(50,5),LO(50,5) CIMENSION ALPHA(15O) DIMENSION ALPHA(15O)
	DIMENSION ICOUNT(50,2)
	DIMENSION TOCC(15C)
	CIMENSION D1(7), C2(7), M1(7), M2(7), M1SIG(7), M2SIG(7)
	DIMENSION DITTO(84), DITTA(84)
	DIMENSION FACT(25)
	CIMENSION DIM(5)
	DIMENSION NIN(7)
	DIMENSION LQ(150),MQ(150) DIMENSION ENERGIADA DISCON XICEA VICEA DICEA DICEA
	CONNON /AAA/ S
	CEMMON /EAB/ ICCC, XX, YY, N, NDIM
	CCMMON /CAC/ ZZ, LQ
	CEMMON /GAD/ A, B
	COMMON /CAE/ ICCUNT, MC, ALPHA, CORE
	COMMON /CAF/ MS, MC, MMM, MORB, MK, CON
	CCMMON /CAG/ L1, L2, N1, N2, SK1, SK2
	CORMON /CAT/ SEPARN; RHEE; MAJ; MAN Cormon /cat/ Nin
	$\begin{array}{c} c \in \mathbb{R} \\ c \in $
	CEMMON /CAS/ XIY XZY IIY IZY ZZ
	CEMMON /CAL/ FACT
	COMMON /GAM/ AR, AL, MR
	CCMMGN /DAN/ ER, SUM
1	FCRMAT (712,F14.6,212)
6	FGRMAT(1H1)
5	FCFMAT (72H
1	
715	FURMAILDIZII970-0970-233 ECONAT/100 EGY 200 DISTANCE MATRIX ////
2	FORMAT (13, F7, 2, 3F10, 6, F12, 3, F6, 3)
8	FCRMAT (617, 3F10.6, F6.3, F6.2, F12.3, F6.3)
\$	FCRMAT(110H ORBITAL ATOM N L M SIGMA X
1	1 Y Z EXP ALPHA PRE-EXP EXPON)
814	FGRMAT(44HOH(I,J)=0.5*K*S(I,J)*(H(I,I)+H(J,J)) WITH K= F7.2)
815	FCFMAT(25HOH(I,J)=K*S(I,J) WITH K= F7.2)
616 571	$FLK = AI \left(\frac{1}{2} \right) = K + S \left(\frac{1}{2} \right) + S \left(\frac{1}{2} \right) + H \left(\frac{1}{2} \right) + H \left(\frac{1}{2} \right) = H \left(\frac{1}{2} \right) +
822	FORMATING THE UNDITALS ARE NUMBERED AS FULLUNS
\$77	FCRMAT(7211)
	NTRN=1
1111	READ (5,1)N,MS,FC,MMM,MCRB,MK,MR,CCN,MG,NPP
	IF(N) 3495,3495,3496

.

```
3496 CONTINUE
       WRITE (6,6)
       REAC (5,5)
       WRITE (6,5)
       CC 10 I=1,N
      READ ( 5 ,2)NOR(I,1),CORE(I),XX(I),YY(I),ZZ(I),AR(I),AL(I)
      NCPB=NCR(I,1)
       REAC ( 5 ,4)(NC(I,J),LC(I,J),SLA(I,J),HII(I,J),J=1,NORB)
   10 CONTINUE
      WRITE ( 6 ,82
WRITE ( 6 ,9)
                 ,821)
      K=1
      DC 100 I=1,N
      ICCUNT(I,1)=K
      NCFE=NOR(I,1)
       LC(I) = LC(I, I)
      CC S8 J=1,NORB
      IF(LQ(I)-LO(I,J)) 31,32,32
   31 LQ(I) = LG(I,J)
   32 CENTINUE
      KF=K+2*LC(I,J)
      KS=KF-K+1
С
С
      GECRGE IS A LITTLE SUBROUTINE WHICH ORDERS M AND SIGMA FOR A GIVEN
С
      CALL GECFGE(M1, M1SIG, KS)
      DC 20 M=1,KS
      ₩₩=M+K-1
      ALPHA(MM)=HII(I,J)
   20 WRITE ( 6 ,8)MM,I,ND(I,J),LD(I,J),MI(N),MISIG(M),XX(I),YY(I),
     1ZZ(I),SLA(I,J),ALFHA(MM),AR(I),AL(I)
   98 K=K+KS
  100 ICCUNT(I_{2})=K-1
      NDIM=ICCUNT(N,2)
С
     . ICCUNT IS AN ARRAY WHICH TELLS FOR ATOM I THE NUMBERS OF THE FIRST
C
      OREITALS BELONGING TO THAT ATCM
С
С
      THIS IS USEFUL IN THE POPULATION ANALYSIS WHEN TOTALS OVER ATOMS A
С
      WKK=WK+1
      GC TO (817,818,819),MKK
  817 WRITE ( 6,814)CCN
  GC TO 820
818 WRITE ( 6
               6 ,815)CON
      GE TO 820
  819 WRITE ( 6 ,816)CCN
  820 CONTINUE
      IF(NDIM-140) 823,823,824
  824 WRITE ( 6,822)
      STCP
  823 CONTINUE
С
С
      NOIM = TOTAL NUMBER OF CRBITALS
С
С
      NCW THE DISTANCE MATRIX IS COMPUTED AND PRINTED OUT
```

```
С
      CC 714 I=1,N
      DG 714 J=1,N
  714 S(I,J)=SQRT((XX(I)-XX(J))**2+(YY(I)-YY(J))**2+(ZZ(I)-ZZ(J))**2)
      WRITE ( 6,715)
      CALL PEGLEG(S,N)
¢
С
      CALL REPU
C
C
c
c
      WE NOW COMPUTE ALL THE FACTORIALS NECESSARY FOR THE GVERLAP COMPUT
      HIGHEST IS FACT(24)
                                AND FACT(I)=FACTORIAL(I-1)
С
      RESTRICTION N1 CR N2 MUST BE LESS THAN 13
      WHICH IS NOT MUCH OF A RESTRICTION AFTER ALL
С
Ċ
      FACT(1)=1.0
      DC 2293 I=2,25
      II = I - 1
2293 FACT(I)=FLOAT(II)*FACT(II)
      DC 94C I=1,NDIM
CC 940 J=1,NDIM
  94C S(I,J)=C.C
```
CITTC(1)=1.0 DITTA(1)=1.0 M1(1)=C M2(1)=0 MISIG(1)=C M2SIG(1)=CNIN(3)=1 NIN(5)=10 NIN(7) = 35NM1=N-1 THE EVALUATION OF THE OVERLAP IS DONE IN A SET OF EIGHT NESTED С С DC LCCPS THE OUTERMOST ONE DEFINES AN ATOM PAUR, J GREATER THAN С THE NEXT TWO LOOPS IN DEFINE AN ORBITAL PAIR II AND JJ IF THERE č MCRE THAN ONE CRBITAL ON A GIVEN ATOM THE NEXT TWO DO LOOPS RUN OVER ALL THE VALUES OF M AND SIGMA FOR BOTH ORBITALS THE INNERFOST PAIR OF LOOPS RUNS OVER M AND SIGMA GAIN THIS TIME С Ċ С THE RESOLVED SET THE ARRAY LQ CONTAINS THE MAXIMUM L QUANTUM NUMBER FOR A GIVEN AT THE ROTATION SUERGUTINE CERES (WHICH CALLS RTNMXN) THEN NEED BE С С С CALLED CNLY ONCE FOR EACH SET OF ATOMS 5555 DC 1000 I=1,NM1 IP=I+1IKIN= ICCUNT(I,1) x1 = xx(1)Y1=YY(I)Z1 = ZZ(I)NORBI=NOR(I,1) CC 1000 J=IP,N JKIN=ICCUNT(J,1) N1S=0 N2S=C NI=0NJ=C X2=XX(J) Y2=YY(J) $Z_2 = Z_2(J)$ NORBJ=NOR(J,1) SEPARN=SCRT((X1-X2)**2+(Y1-Y2)**2+(Z1-Z2)**2) LLCI=LC(I) LLCJ=LC(J) CALL CERESILLGI, DITTO, SEPARN) CALL CERES(LLQJ,DITTA,SEPARN) SEFARN=SEPARN/0.529171 DC 900 II=1,NORBI N1=NC(I,II) L1=LC(I,II)IN=IKIN+NI N1S=2*L1+1 NI=NI+N1S NJ=0 N2S=0SK1=SLA(I,II) IF(L1) 901,902,901 SO1 CALL GEORGE(M1, MISIG, N1S)

```
S02 CONTINUE
      DC S10 JJ=1,NORBJ
      N2=NG(J,JJ)
       SK2=SLA(J,JJ)
      L2=LC(J,JJ)
       JIN=JKIN+NJ
      N2S=2*L2+1
      NJ=NJ+N2S
      IF(L2) 903,904,903
  903 CALL GEORGE(M2, M2SIG, N2S)
  S04 CONTINUE
С
С
      MCLPAB ACQUIRES INFORMATION FROM AFUNCT AND BFUNCT THRU COMMON.
С
      JMAX=N1+N2+L1+L2+1
      CALL AFUNCT(SEPARN*(SK1+SK2)/2.0, JMAX)
      CALL BFUNCT(SEPARN*(SK1-SK2)/2.0,JMAX)
      CC 80C III=1,N1S
      CALL SEAN(III,N1S,DITTO,D1,NIN)
      CC 801 JJJ=1,N2S
      S(IN, JIN)=0.0
      CALL SEAN(JJJ,N2S,DITTA,D2,NIN)
С
      THE ORBITAL HAS BEEN EXPANDED WRT A LOCAL COGRDINATE SYSTEM.
С
С
      JAP AND KAM ARE INDICES RANGING OVER THE LOCAL COORDINATE SYSTEMS
С
      CF I AND J RESPECTIVELY. WHEE REPRESENTS THE OVERLAP INTEGRAL
      BETWEEN LOCAL ORBITALS JAM AND KAM.
С
С
      CC 118 JAM=1,N1S
      CC 118 KAM=1,N2S
      IF(M1(JAN)-M2(KAM)) 118,120,118
  12C IF(M1SIG(JAM)-M2SIG(KAM)) 118,121,118
  121 MAJ=MI(JAM)
      MAK=M2(KAM)
      CALL MCLPAB
      BECAUSE MCLPAB CALCULATES OVERLAPS BETWEEN A RT.-HANDED SYSTEM AND
С
C
      A LEFT-HANDED SYSTEM, WITH Z-AXES PCINTING TOWARDS EACH OTHER, WE
С
      MUST CHANGE SOME SIGNS.
С
      CC 124 KIX=1,7,2
      IF(L2-(KIX - 1)+MAK) 124,123,124
  124 CONTINUE
      WHEE =- WHEE
  123 S(IN, JIN)=S(IN, JIN)+D1(JAM)+WHEE+D2(KAM)
  118 CENTINUE
  801 JIN=JIN+1
      JIN=JIN-N2S
  EOG IN=IN+1
  91C IN=IN-N1S
  900 CONTINUE
 1000 CONTINUE
      CC 1201 I=1,NDIM
      S(I,I)=1.0
      DC 1201 J=I,NDIM
 1201 S(J,I)=S(I,J)
```

С С С THE CCCUPATION NUMBERS ARE READ IN FROM MOST BONDING TO ANTIBONDIN BUT THEN THE DIRECTION HAS TO BE REVERSED BECAUSE THE EIGENVALUES c c CCME GUT IN REVERSE ORDER READ (5 ,977) (MQ(I), I=1, NDIM) DC 850 I=1,NDIM J=NDIM-I+1 856 IDCC(J)=MQ(I) CALL IGCR IF(MG) 3495, 3495, 3001 3495 GC TC 1111 3001 ENER(NTRN)=SUM NTEN=NTRN+1 IF(NTRN.GT.NPP) GC TO 3002 GC TC 1111 3002 READ (5,3112) IS, MO, XS, YS, XSF, XM, YSF, YM 3112 FCFMAT(213,6F5.2) REAC (5,3113) EINF, (DIS(I), I=1, NPP) 3113 FCRMAT (F10.4,12F5.3) REAC(5,3114) XL,YL,GL,DL 3114 FCRMAT (2CA4) WRITE(6,3) FCFMAT(//12X, "DISTANCE", 4X, "ENERGY", /) 3 WRITE(6,2013) (DIS(I), ENER(I), I=1, NPP) WRITE (6,3) DC 3115 I=1,NPP ENER(I)=ENER(I)-EINF write(6,2013) DIS(I),ENER(I) 2013 FCRMAT(10X, 2F10.3) 3115 CENTINUE CALL GRAPH(NPP, DIS, ENER, IS, MO, XS, YS, XSF, XM, YSF, YM, XL, YL, GL, DL) STEP END

с		SUBROUTINE RTNMXN(O) THIS SUBROUTINE CALCULATES THE TRANSFORMATION COEFFICIENTS OF ANGU
С		MCMENTUM FUNCTIONS THESE ARE CONSTRUCTED BY RECURSION FORMULA
С		RESTRICTION- F ORBITALS ARE THE HIGHEST L THAT CAN BE TRANSFORMED
C		THE CROERING OF COEFFICIENTS IS SUCH THAT M PRIME VARIES FASTEST
C		FOLLOWED BY SIGMA PRIME FOLLOWED BY M FOLLOWED BY SIGMA
ć		SIGMA =0 IS THE REAL CUSINE COMBINATION, SIGMA=1 IS THE SINE
C C		THE SUBRUITINE ORIGINALLY WRITTEN BY R.M. PHIZER AND HUUTTED
٤ د		FUR THE SPECIAL LASE WHERE CALPHARI SALPHARO . ACTUALLY
Ċ		THE DEFINITION OF EVER ANGLES OF THE KUTATION WHICH TAKEN THE
c c		ALCUAR COORDINATE SISTEM INTO THE LUCAR ONE IS THE WITH
L		CINFNSION D(84)
		DIMENSION SA(2.4), SG(2.4), IEPS(4), DP(20), DN(10)
		COMMON /RTN/ LMAX, CBETA, SBETA, CGAMMA, SGAMMA
		SA(1,1)=1.0
		Se(1,1)=1.0
		SA(2,1)=C.O
		SG(2,1)=0.0
		IEFS(1)=1
		DO IC M=1,LMAX
		SA(1, M+1)=SA(1, M)
		SA(29M+1)=SA(29M) SC(1 H1)=SC(1 H)+CCAHHA_SC(2 H)+SCANMA
		SC(1, MT1)=SC(1, M)+CCAMMA=SC(2, M)+SCAMMA SC(2, V+1)=SC(2, M)+CCAMMA=SC(1, M)+SCAMMA
	10	TEPS(M+1)=2
	10	
		IP#2=0
		IM=0
		IMM1=C
		IMM2=C
		IP=1
		CP(1)=1.C
		CM(1)=0.0
		CL 20C L=1,LMAX
		LF1=L=1 1T2H1=L1H1
		E12010AT(#172W1)
		F2=F1CAT(1M)*1T2W1)
		F3=FLGAT(LM1*L)
		IL=L+1
		CC 190 MPP=1,IL
		NP=MPP-1
		CC 18C MM=1,NPP
		h=hh-1
		IF=IP+1
		IF(MP-LM1) 40,8C,120
	4 C	IPM1=IPM1+1 IPM2-IPM2-1
		[FFZ=1FFZ+1 CNST1~C1/SOPT/E/OAT///~WD)+//~WD)+//~WN+//
		UNGIT=(1) GRNINE LUMINNE METALEMETALIMETALEMETALEMETAL
		IF(M) 44,44,60
	44	DP(IP)=CNST1+(CBETA+DP(IPM1)-CNST2+DP(IPM2))
	-	GC TO 180

```
6C IM=IM+1
    IMNI=IMM1+1
    IMM2=IMM2+1
    CNST3=FLCAT(MP*M)/F3
    DP(IP)=CNST1*(CEETA*DP(IFM1)-CNST2*DP(IPM2)-CNST3*DM(IMM1))
    CF(IM)=CNST1*(CEETA+DM(IMM1)-CNST2+DM(IMM2)-CNST3+DP(IPM1))
    GC TC 18C
 EC IPM1=IPM1+1
    CNST1=F1/SQRT(FLOAT(LT2M1*(L-M)*(L+M)))
                                                         .
    IF(M) 84,84,100
 84 DP(IP)=CNST1*CBETA*DP(IPM1)
    GC TC 180
10C IM=IM+1
    IMN1=IMM1+1
    CNST3=FLCAT(MP*M)/F3
    CP(IP)=CNST1*(CBETA*DP(IPM1)-CNST3*DM(IMM1))
    CM(IM)=CNST1*(CEETA*DM(IMM1)-CNST3*DP(IPM1))
    GC TC 18C
120 IF(M-L) 130,160,130
130 CNST4=SQRT(F1/FLOAT((L-M)*(L+M)*IEPS(L)))
    IF(M) 145,140,145
140 IPM1=IPM1-LM1
    IMM1=IMM1-LM1
   GC TO 148
145 IM=IM+1
    IPM1=IPM1+1
    IMM1=IMM1+1
   CM(IM) =- CNST4*SEETA*DM(IMM1)
148 DP(IP)=-CNST4*SBETA*DP(IPM1)
   GC TO 18C
16C IN=IN+1
    IF(M-1) 164,164,168
164 CP(IP)=-CEETA*DP(IPM1)
   CM(IM)=-CP(IPM1)
   GC TO 180
168 CP(IP)=~.5*(CBETA*DP(IPM1)+DM(IMM1))
   DM(IM)=-.5*(CBETA*DM(IMM1)+DP(IPM1))
18C CENTINUE
1SC CONTINUE
200 CENTINUE
   F5IG=-1.0
   FSIGP=-1.0
   T = 1
   C(1)=1.C
   CC 400 L=1,LMAX
   IP=(L*(L+1)*(L+2))/6+1
   IM=((L-1)+L*(L+1))/6
   DC 390 ISIGG=1,2
   ISIG=ISIGG-1
   ICSIG=1-ISIG
   FSIG=-FSIG
   FM=-FSIG
   LL=L+1
   DO 38C MZ=ISIGG,LL
   M=MZ-1
```

NF1=MZ FN=-FM DC 37C ISIGPP=1.2 ISIGP=ISIGPP-1 ICSIGP=1-ISIGP FSIGP=-FSIGP CC 360 MPZ=ISIGPP,LL MP=MPZ-1 MPF1=MPZ I = I + IIF(MP-M) 300,330,330 30C I1F=IP+(M*(M+1))/2+MP I1M=IM+((M-1)*N)/2+MP GC TO 35C 33C IIF=IP+(MP*(MP+1))/2+M I1#=IN+((MP-1)*MP)/2+M 35C IF(I1M) 351,352,351 351 D(I)=FM*(FSIGP*DP(I1P)*SA(ISIGP+1, MPP1)*SG(ISIG+1, MP1)-FSIG* lcm(I1m)*SA(ICSIGP+1,MPP1)*SG(ICSIG+1,MP1)) GC TO 360 352 C(I)=FV*(FSIGP*CP(I1P)*SA(ISIGP+1, MPP1)*SG(ISIG+1, MP1)) 36C CONTINUE 376 CONTINUE 380 CONTINUE 39C CONTINUE 400 CENTINUE RETURN ENC

```
SLERGUTINE BFUNCT (RHC2, IX)
С
      THIS SUBROUTINE CALCULATES THE B-FUNCTIONS WITH INDICES OF O TO IX
С
      FOR THE GIVEN VALUE OF RHO2.
С
С
      CRIGINALLY WRITTEN BY R.M. STEVENS
      IF THE DIMENSION OF B IS CHANGED THAN ONE MUST ALSO MODIFY
С
С
      THE STATEMENT DEFINING IS SO THAT THE SECOND ENTRY IN THE ARGUMENT
      XMINOF IS THE DIMENSION CF B MINUS 1
С
C
      CIMENSION A(100), B(100)
      CCMMON /CAD/ A, B
      IA=IFIX(2.0*RH02)
      IR=IAES(IA)
      IS=MINC(IR+1,99)
      IF (RHC2) 25,35,25
   25 C = EXF(RHO2)
      + = EXP(-RHO2)
      R = C - H
С
      IF THE VALUE OF RHOZ IS TOO SMALL THE SINH MUST BE OBTAINED BY
С
              SUMMING THE INFINITE SERIES RATHER THAN BY ADDITION OF TWO
С
С
              EXPONENTIALS.
С
      IF (ABS(R)-0.1) 26,28,28
   26 RA=0.0
      T = RHC2
      CC 27 I=2,25
      T = T*RHC2**2/FLOAT((2*I-1)*(2*I-2))
   27 RA = RA+T
      F = (R#+RE02)*2.
   28 B(1) = R/RHO2
C
      AS MANY SUCCESSIVE B-FUNCTIONS ARE GENERATED FROM B-0 BY THE RE-
С
c
             CURRENCE FORMULAE AS ACCURACY WILL PERMIT.
С
      CC 51 I=2,IX,IS
      IF (IR) 32,40,32
   32 IL=IS-1
      CC 31 J=1,IL
      K = I + J - I
      IF ((-1)**K) 29,29,30
   29 E(K) = (R+FLOAT(K-1)*B(K-1))/RHO2
      GC TO 31
   3C B(K) = -(C+H-FLCAT(K-1)*B(K-1))/RHO2
   31 CENTINUE
   4C IN=I+IS-1
      IF (IN-IX) 39,39,38
С
      AFTER THE RECURRENCE FORMULAE HAVE BEEN APPLIED AN APPROPRIATE
С
С
             NUMBER OF TIMES THE NEXT B-FUNCTION IS OBTAINED BY SUM-
С
      MATICN OF THE INFINITE SERIES.
C
   35 IF((-1)**IN) 44,44,42
   42 \text{ TR} = \text{RHC2}
```

```
105 B(IN) = -2.*TR/FLCAT(IN+1)

EC 43 J = 1,500

TR = TR*RH02**2/FL0AT((2*J)*(2*J+1))
       IF (AES(TR/B(IN))-.0000001) 51,51,43
   43 E(IN) = E(IN)-2.*TR/FLOAT(IN+1+2*J)
   44 TR = 1.
  107 \text{ e(IN)} = 2.*TR/FLOAT(IN)
       DC 46 J = 1,500
       TR = TR*RH02**2/FL0AT((2*J)*(2*J-1))
       IF (ABS(TR/B(IN))-.0000001) 51,51,46
   46 B(IN) = B(IN)+2.*TR/FLOAT(IN+2*J)
   51 CENTINUE
       GC TC 38
С
       IF THE ARGUMENT IS ZERO A SEPARATE FORMULA MUST BE USED.
С
С
   35 JY = IX/2+2
       CC 36 I =1,JY
       E(2*I)=0.0
   36 3(2*I-1)=2./FLCAT(2*I-1)
   38 RETURN
      END
                               ٠
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SLERGUTINE BINCCE(I, J, B)
с
с
с
       THIS SUBROUTINE COMPUTES BINOMIAL COEFFICIENTS
      DIMENSION FACT(25)
      CCMMON /CAL/ FACT
      K=J-I
      B=FACT (J+1)/FACT(I+1)/FACT(K+1)
    1 RETURN
      END
      SUBROUTINE SEAN(I, N, DITTO, D, NIN)
THIS SUBROUTINE FINDS THE RIGHT ELEMENTS OF DITTO AND PUTS THEM IN
С
       SMALLER ARRAY D
С
      DIMENSION DITTO(84), D(7), NIN(7)
      IF(N-1) 1,2,1
    2 D(1)=1.0
      RETURN
    1 CONTINUE
      LIB=NIN(N)+N*(I-1)
      CC 85 JIE=1,N
      INGRID=JIB+LIB
   85 D(JIB)=CITTO(INGRID)
      RETURN
      ENC
```

SUBROUTINE SEAN(I, N, DITTO, D, NIN) THIS SUBROUTINE FINDS THE RIGHT ELEMENTS OF DITTO AND PUTS THEM IN С Ċ SMALLER ARRAY D DIMENSION DITTO(84), D(7), NIN(7) IF(N-1) 1,2,1 2 D(1) = 1.0RETURN 1 CONTINUE LIB=NIN(N)+N*(I-1) DC 85 JIB=1.N INGRID=JIB+LIB 85 D(JIB)=DITTO(INGRID) RETURN END

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SUBROUTINE AFUNCT (RHO1, J) C C THIS SUBROUTINE CALCULATES THE A-FUNCTIONS WITH INDICES OF 0 TO J FOR THE GIVEN VALUE OF RHO1. C ORIGINALLY WRITTEN BY R.M. STEVENS C DIMENSION A(100),B(100) CCMMCN /CAD/ A, B C = EXP(-RHO1) A(1) = C/RHO1 DC 15 I = 2,J 15 A(I) = (FLOAT(I-1)*A(I-1)+C)/RHO1 RETURN ENC

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SUBROUTINE MOLPAB
С
      THIS SUBROUTINE CALCULATES TWO-CENTER OVERLAP INTEGRALS.
C
ē
      THE INTEGRALS ARE CALCULATED BY TRANSFORMATION TO ELLIPSOIDAL
             COORDINATES AND THEREBY EXPRESSED IN TERMS OF C-FUNCTIONS.
C
C
      (SEE J.C.P., 24, 201.)
      ORIGINALLY WRITTEN BY R.M. STEVENS
С
С
      DIMENSION FACT(25)
      COMMON /OAG/ L1, L2, N1, N2, SK1, SK2
COMMON /OAH/ R, VEST, M1, M2
      COMMON /OAL/ FACT
      VEST = 0.0
      STRAD = 0.0
      RHOA = R*SK1
      RHOB = R * SK2
      F1=FACT(2*N1+1)
      F2=FACT(2*N2+1)
      IF=L1-M1+1
      F3=FACT(IF)
      IF=L2-N2+1
      F4=FACT(IF)
      IF=L1+M1+1
      F5=FACT(IF)
     IF=L2+M2+1
     F6=FACT(IF)
     TERM = 2.**(N1+N2-L1+L2)*SQRT(FLOAT((2*L1+1)*(2*L2+1))*F3*F4/F1/
    1F2/F5/F6*SK1**(2*N1+1)*SK2**(-2*N1-1))
      JEND = 1 + \{ (L1 - M1) / 2 \}
     KEND = 1 + ((L2 - M2)/2)
     DO 50 J=1, JEND
      JU=J-1
     IF=2*L1-2*JU+1
     F11=FACT(IF)
     IF=L1-M1-2*JU+1
     F13=FACT(IF)
     F15=FACT(JU+1)
      IF=L1-JU+1
     F17=FACT(IF)
     DC 50 K=1,KEND
     KU=K-1
     IF=2*L2-2*KU+1
     F12=FACT(IF)
     IF=L2-M2-2*KU+1
     F14=FACT(IF)
     F16=FACT(KU+1)
     IF=L2-KU+1
     F18=FACT(IF)
     CALL CFUNCT (N1-L1+2*JU,N2-L2+2*KU,L1-M1-2*JU,L2-M2-2*KU,M1,RHDA,
    1 RHOB, VALUE)
  50 STRAD = STRAD+VALUE*F11*F12/F13/F14/F15/F16/F17/F18*(-1.)**(JU+KU)
     VEST=TERM*STRAD
     RETURN
     END
```

```
SUBROUTINE GEORGE(M, MSIG, NSBSP)

DIMENSIGN M(7), MSIG(7)

M(1)=0

MSIG(1)=0

K=(NSBSP/2)+1

IF(K-1) 5,5,6

6 CCNTINLE

CC 10 J=2,K

M(J)=J-1

MSIG(J)=C

L=K+J-1

M(L)=J-1

10 MSIG(L)=1

5 RETURN

ENC
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C C C C C C C

С

THIS SUBROUTINE CALCULATES THE C-COEFFICIENTS OF ROOTHAAN FCR OVERLAP INTEGRALS ONLY THE C-FUNCTIONS WITH POSITIVE INDICES ARE NECESSARY AND THESE MAY BE CONVENIENTLY CALCULATED BY EXPANSION BY THE BINOMIAL THEOREM. CRIGINALLY WRITTEN BY R.M. STEVENS DIFENSION A(100), B(100) COMMON /GAD/ A, B 25 CCLNT = C.O VEST = TERM#STRAD RETURN END IAE = IA+1I88=I8+1 ICE=IC+1 ICE=ID+1 IEE=IE+1 DC 91 I6=1,IEB J6=I6-1 CALL BINCCE (J6, IE, B6) DC 91 I5=1,IEB J5=I5-1 CALL BINCCE (J5, IE, B5) DC 90 I4=1, IDB J4=I4-1 CALL BINCOE (J4, IC, B4) CC 90 I3=1,IC8 J3=I3-1 CALL BINCCE (J3, IC, B3) CC 90 I2=1,IBB J2=I2-1 CALL BINCCE (J2, I8, B2) CG 90 I1=1, IAB J1=I1-1 CALL BINCCE (J1, IA, B1) TERM = B1*B2*B3*B4*B5*B6*(-1.)**(J2+J5+J6+J4+IE+ID) IR = J1+J2+2*IE-2*J6+IC-J3+ID-J4+1 $IP = IA - J1 + IB - J2 + 2 \times IE - 2 \times J5 + IC - J3 + ID - J4 + 1$ CCUNT = CCUNT +A(IP)*B(IR)*TERM 90 CENTINUE 91 CENTINUE SNAG = CCUNT*(RHOB /2.)**(IA+IB+IC+ID+2*IE+1) 92 RETURN ENC

SLERCUTINE CFUNCT (IA, IB, IC, ID, IE, RHOA, RHOB, SNAG)

```
SLEROLTINE PEGLEG(A, N)
C MATRIX OUTPUT SLBROUTINE
CIMENSION A(140,140)
18 FCRMAT(14,2X,11F10.6)
19 FCFMAT(1H0/1H0,4X,11(6X,12,2X)//)
KITE=C
20 LCw=KITE+1
KITE=KITE+11
KITE=MINC(KITE,N)
wRITE ( 6,19)(I,I=LOW,KITE)
CC 32 I=1,N
32 wRITE ( 6,18)I,(A(I,J),J=LOW,KITE)
IF(N-KITE) 40,4C,20
4C RETURN
ENC
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SUPROUTINE CERES(LQ, DITTO, SEP) THIS SUBROUTINE CALCULATES THE EULER ANGLES OF ROTATION AND С PRECEEEDS TO CALL RINMAN FOR THE GENERATIONS OF THE TRANSFORMATIONS C C OF ANGULAR MOMENTUM FUNCTIONS **DIMENSION DITTO(84)** CCMMCN /CAJ/ X1, X2, Y1, Y2, Z1, Z2 CCMMON /RTN/ LMAX, CBETA, SBETA, CGAMMA, SGAMMA 20C FORMAT('ERROR, POSSIBLY ROUNDING OFF ERROR IN CALCULATION OF SEP') IF(LQ) 100,100,101 10C CITTC(1)=1.0 FETURN 1C1 CONTINUE LMAX=LC 22=22-21 IF(X2-X1) 20,5, 20 5 IF(Y2-Y1) 20, 10, 20 1C CGAMMA=1.C SGAMMA=0.0 CBETA=SIGN(1.0,ZZ) SBETA=C.C GC TC 30 2C ARGL=SEP*SEP-ZZ*ZZ IF (ARGU .GE. 0.) GO TO 25 IFIARGL .GE. -1.E-6) GG TO 24 WRITE (6,200) STEP 24 ARGL=C. 25 SXY=SCRT(ARGU) CBETA=ZZ/SEP SEETA=SXY/SEP CGAMMA=(X2-X1)/SXY SGAMMA = (Y2-Y1)/SXY 30 CALL RTNMXN(DITTO) RETLRN ENC

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SLEROUTINE IGOR
       CEMPUTES H MATRIX FROM S WHICH IS ASSUMED TO BE IN CORE
C
      H GOES OUT ON TAPE IN THEN DIAGONALIZATION AND POPULATION
С
      ANALYSIS ROUTINES ARE CALLED
С
С
                MS=1 PUNCHES OUT BCD S
                                              MC=1 PUNCHES OUT C
      CFTIGNS
С
      MM=1 BYPASSES POPULATION ANALYSIS
      MK=0 USES ADDITIVE WOLFSBERG-HELMHOLTZ FORMULA
С
      MK=1 USES DIRECT PROPORTIONALITY H(I,J)=K*S(I,J)
С
      MK=2 USES GEOMETRIC MEAN WOLFSBERG HELMHOLTZ FORMULA
C
      DIMENSION A(140,140)
      EIMENSION IX(15C)
      DIFENSION ALPHA(150), LATER(150)
      CIMENSION HDIAG(150),8(150)
      CIMENSION IDCC(150)
      CIMENSION CORE(50)
      CIMENSION ICOUNT(50,2)
      COMMON /CAA/ A
              /CAB/IX, ALPHA, B, N, NT
      CEMMON
              /CAE/ ICOUNT, LATER, HDIAG, CORE
/CAF/ MS, MC, MMM, MORB, MK, CON
      CEMMON
      CCMMON
      CCNP=0.5*CON
      ₩K=₩K+1
      PENIND 4
    1 FCFMAT(1H0,50X,23F TOTAL OVERLAP MATRIX
                                                        113
      WRITE ( 6,1)
CALL PEGLEG(A,NT)
    4 FCRMAT(8F5.6)
      IF(MS-1) 2,3,2
    3 kRITE ( 7 ,4)((4(I,J),J=1,NT),I=1,NT)
    2 GC TO (5,6,7),MK
    5 EE 8 I=1,NT
      DC S J=1,NT
    S E(J)=A(I,J)*CONP*(HDIAG(I)+HDIAG(J))
      E(I)=HEIAG(I)
    & WRITE ( 4)(B(J),J=1,NT)
      GC TC 15
   6 DC 10 I=1.NT
      CC 11 J=1,NT
  11 E(J)=A(I,J)*CON
     B(I)=HCIAG(I)
  10 WRITE ( 4)(B(J),J=1,NT)
   GC TC 15
7 CC 12 I=1,NT
     DC 13 J=1,NT
  13 E(J)=A(I,J)*CON*SCRT(HDIAG(I)*HDIAG(J))
     B(I)=HCIAG(I)
  12 WRITE ( 4)(B(J), J=1, NT)
  15 CONTINUE
     REWINC 8
     DC 19 I=1,NT
  19 WRITE ( 8)(A(I, J), J=1, NT)
     CALL SCREN
     IF(MC-1) 14,16,14
                7 ,4)((A(I,J),J=1,NT),I=1,NT)
  16 WRITE (
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14 IF(MMM-1) 17,18,17 17 CALL MELEA 18 RETURN ENC

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SUBROUTINE MELBA POPULATION ANALYSIS SUBROUTINE C IN CORE ASSUMES S ON IS BY ROWS CIMENSION A(140,140), IOCC(150), HDIAG(150), B(150), C(150) DIMENSION ICOUNT(50,2), LATER(150) **DIMENSION CORE(50)** CEPHEN /CAA/ A CCMMON /CAB/ ICCC, HDIAG, C, N, NT CCMMON /DAE/ ICOUNT, LATER, B, CCRE CEMMON /CAF/ MS, MC, MMN, MORB, MK, CON 3 FORMAT(//40X,35H MULLIKEN OVERLAP POPULATIONS FOR 13,10H ELECTR 1CNS//) ISLM=0 CC 5 I=1,NT 5 ISLM=ISUM+IOCC(I) WRITE (6,3)ISUM REWIND 3 DC 4 J=1,NT 4 WRITE (3)(A(I, J), I=1, NT) IF(MORE) 6,6,7 7 CC 8 I=1,NT 8 HDIAG(I)=A(I,MORB) 6 CONTINUE CALL CARUSO (IDCC, A, C, NT) CALL PEGLEG(A,NT) IF(MOR8) 9,9,10 10 REWIND 4 wRITE (4)((A(I,J),J=1,NT),I=1,NT) REWIND 8 DC 11 I=1,NT 11 READ (&)(A(I, J), J=1,NT) CC 12 I=1,NT DC 12 J=1,NT 12 A(I,J)=2.0*A(I,J)*HDIAG(I)*HDIAG(J)WRITE (6 ,13) FORB 13 FORMAT(1H0,20X,45H OVERLAP POPULATION FOR ELECTRON IN ORBITAL 113//) CALL PEGLEG(A,NT) CALL JENNY(ICOUNT,A, N, C) REWIND 4 READ (4)((A(1,J),J=1,NT),I=1,NT) 9 CONTINUE CALL IENNY(ICOUNT,A, N, C) CALL CALLAS(A, NT, C) CC 18 I=1,NT 18 B(I)=0.0 DE 15 M=1,NT RR=FLCAT(IOCC(M))/2.0 IF(RR) 15,15,17 17 DC 16 I=1,NT 16 B(I)=B(I)+RR#A(I,M) **15 CONTINUE** 78 FORMAT(1HC,50X,20H ORBITAL CHARGES 1/1 WRITE (6,78)

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F10.6//)

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SUBROLTINE CALLAS(A, NT, B)
COMPUTES TOTAL GROSS OVERLAP POPULATION MATRIX
С
       CIMENSIGN A(140,140), B(150)
       REWIND 3
    CC 3 J=1,NT
3 READ ( 3)(A(I,J),I=1,NT
                                       )
       CALL CCKI
С
       NEW A=SC
      REWINE 3
       CC 1 M=1,NT
       READ (3) (B(J), J=1, NT)
      CC 1 I=1,NT
    1 A(I,M) = A(I,M) * B(I) * 2.0
    2 FCRMAT(1H0,50X,50HCHARGE MATRIX FOR MO'S WITH TWO ELECTRONS IN EAC
     18
                        //}
      WRITE ( 6,2)
      CALL PEGLEG(A,NT)
      RETURN
      END
```

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С DIAGONALIZATION OF A REAL SYMMETRIC MATRIX BY THE JACOBI METHOD. С с с THERE IS A FACTOR QMULT INSERTED WHICH ESSENTIALLY SETS THE Ĉ CONVERGENCE CRITERION FOR THE DIAGONALIZATION THIS WAS ORIGINALLY EUT NOW HAS BEEN SET TO 100.0 WHIVH S TC BETTER THAN CNE PART ON 10 TO THE 8 C WHIVH SHOULD GIVE A TCTAL ENERGY С CIFENSION H(140,140), IA(150), HDIAG(150), X(150), IQ(150), ARRAY(252) CEMMEN /DAA/ H CCMMCN /CAB/IA, HDIAG, X, NDIM, N CCMMON /CUD/ ARRAY, IPIV, JPIV, CGSINE, SINE THE SUBROUTINE OPERATES ONLY ON THE ELEMENTS OF H THAT ARE TO THE С RIGHT OF THE MAIN DIAGONAL. THUS, ONLY A TRIANGULAR SECTION NEED С С EE STOREC. С NR IS THE NUMBER OF ROTATIONS С LAFITE , MOUTON AND YOUEM ARE SUBROUTINES USED TO PACK WRITE AND R С С THE VARIOUS QUANTITIES NEEDED TO COMPUTE THE EIGENVECTORS OUT ON TAFES - THE ROUTINE CAN CNLY HOLD ONE MATRIX IN CORE AT A TIME С С THE EIGENVALUES ARE PUT OUT IN THE VECTOR HOIAG С С С TAPE ITAPEU IS USED TO STORE IPIV, JPIV, COSINE, SINE FOR EACH FCTATION, IN ORDER THAT THE EIGENVECTORS MAY BE COMPUTED AFTER С С DIAGONALIZATION IS COMPLETE. С ¥€¥≈1 REWIND 3 999 FCFMAT(1+0,3HNR=,15) NR=0 IF (N-1) 1000,1000,17 С С SCAN FOR LARGEST OFF-DIAGONAL ELEMENT IN EACH ROW. X(I) CONTAINS С LARGEST ELEMENT IN THE ITH ROW, AND IQ(I) HOLDS SECOND SUBSCRIPT С **CEFINING POSITION OF ELEMENT.** С 17 N∦I1=N-1 DC 30 I=1,NMI1 X(I)=0. IPL1=I+1 CC 30 J=IPL1,N IF (X(I)-ABS(H(I,J))) 20,20,30 $2C \times (I) = ABS(H(I,J))$ IC(I)=J**3C CONTINUE** С С SET INDICATOR FCR SHUT-CFF. RAP=2**-27. С RAF=.745058E-08 CMULT=100.0 RAF=RAP+QMULT HCTEST=1.0E38 С

SLEROUTINE CUPID

FINC MAXIMUM OF X(I)'S FOR PIVOT ELEMENT, AND TEST FOR END OF С С PROBLEM. C 40 DC 70 I=1,NMI1 IF (I-1) 60,60,45 45 IF (XMAX-X(I)) 60,70,70 6C XMAX=X(I) IPIV=I JPIV=IC(I) 7C CONTINUE С IF XMAX LESS THAN HDTEST, REVISE HDTEST. С С IF (XMAX) 1000,1000,80 80 IF (HOTEST) 90,50,85 85 IF (XMAX-HDTEST) 90,90,148 90 HOIMIN=ABS(H(1,1)) CG 110 I=2,N IF (HCIMIN-ABS(H(I,I))) 110,110,100 10C HEIMIN=ABS(H(I,I)) 11C CONTINUE HOTEST=HDIMIN#RAP С С RETURN IF MAXIMUM H(I,J) LESS THAN (2++-27)+ABSF(H(K,K)-MIN) С IF (HDTEST-XMAX) 148,1000,1000 148 NR=NR+1 С С CCMPUTE TANGENT, SINE, AND COSINE, H(I,I), H(J,J) Ċ 150 TANG=SIGN(2.0,(H(IPIV,IPIV)-H(JPIV,JPIV)))*H(IPIV,JPIV)/(ABS(H(IPI 1V, IPIV)-H(JPIV, JPIV))+SQRT((H(IPIV, IPIV)-H(JPIV, JPIV))**2+4.0*H(IP 2IV, JPIV) ** 2) } CCSINE=1.0/SQRT(1.0+TANG**2) SINE=TANG*COSINE FII=H(IPIV, IPIV) +(IPIV,IPIV)=COSINE**2*(HII+TANG*(2.*H(IPIV,JPIV)+TANG* 1F(JPIV,JPIV))) H(JPIV, JPIV) =COSINE*+2*(H(JPIV, JPIV)-TANG+(2.+H(IPIV, JPIV)-TANG+ 1HII)) +(IPIV,JPIV)=0. С С PSEUDO-RANK THE EIGENVALUES. ADJUST SIN AND COS FOR COMPUTATION С H(I,K). С IF (H(IPIV, IPIV)-H(JPIV, JPIV)) 152,153,153 152 HTEMP = H(IPIV, IPIV) +(IPIV, IPIV)=H(JPIV, JPIV) +(JPIV, JPIV) =HTEMP С С С RECOMPUTE SINE AND COS HTEMP=SIGN(1.0,-SINE) +COSINE CCSINE=ABS(SINE) SINE=HTEMP

```
153 CONTINUE
с
с
      INSPECT THE IQ'S BETWEEN I+1 AND N-1 TO DETERMINE WHETHER A NEW
      MAXINUM VALUE SHOULD BE CONPUTED, SINCE THE PRESENT MAXIMUM IS IN
С
С
      ROW I OR J.
С
      CO 350 I=1,NMI1
      IF (I-IPIV) 210,350,200
  200 IF (I-JPIV) 210,350,210
  210 IF (IG(I)-IPIV) 230,240,230
  230 IF (IQ(I)-JPIV) 350,240,350
  24C K=IQ(I)
  25C HTEMP=H(I,K)
      H(I,K)=0.
      IFL1=I+1
      X(I)=0.
C
Ċ
      SEARCH IN DEPLETED RCW FCR NEW MAXIMUM.
С
      CC 320 J=IPL1,N
      IF (X(I)-ABS(H(I,J))) 300,300,320
  300 X(I)=ABS(H(I,J))
      IC(1)=J
  32C CONTINUE
      H(I,K)=HTEMP
  35C CENTINUE
      X(IPIV)=0.
      X(JPIV)=0.
С
C
      CHANGE THE OTHER ELEMENTS OF H.
С
      CC 53C I=1,N
      IF (I-IPIV) 370,530,420
  370 HTEMP=H(I.IPIV)
      H(I, IPIV)=COSINE+FTEMP+SINE+H(I, JPIV)
      IF (X(I)-ABS(H(I, IPIV))) 380,390,390
  380 X(I)=AES(H(I,IPIV))
      IC(I)=IPIV
  39C H(I, JPIV) =- SINE +HTEMP+CCSINE+H(I, JPIV)
      IF (X(I)-A8S(H(I, JPIV))) 400,530,530
  40C X(I)=ABS(H(I,JPIV))
      IQ(I)=JPIV
      GC TO 530
  42C IF (I-JPIV) 430,530,480
  43C HTEMP=H(IPIV,I)
      H(IPIV,I)=COSINE+HTEMP+SINE+H(I,JPIV)
      IF (X(IPIV)-ABS(H(IPIV,I))) 440,450,450
 440 X(IPIV)=ABS(H(IPIV,I))
      IG(IPIV)=I
 450 H(I,JPIV)=-SINE#HTEMP+COSINE#H(I,JPIV)
      IF (X(I)-ABS(H(I,JPIV))) 400,530,530
 48C HTEMP=+(IPIV,I)
      H(IPIV,I)=COSINE+HTEMP+SINE+H(JPIV,I)
      IF (X(IPIV)-ABS(H(IPIV,I))) 490,500,500
 490 X(IPIV)=ABS(H(IPIV,I))
```

IC(IPIV)=I 50C H(JPIV,I) =-SINE*HTEMP+CCSINE*H(JPIV,I) IF (X(JPIV)-ABS(H(JPIV,I))) 510,530,530 510 X(JPIV)=A85(H(JFIV,I)) IC(JPIV)=I 530 CONTINUE 540 CALL LAFITE(MOM) GG TO 4C 1000 WRITE (6,999)NR IF(MCM-1) 2014,2014,2015 2015 CALL MOUTON 2014 MEM=1 1002.00 1004 J=1.N 1004 HDIAG(J)=H(J,J) DC 14 I=1,N CC 12 J=1,N H(I,J)=0. 12 CONTINUE +(I,I)=1.14 CONTINUE IF (NR) 1005,1005,1006 1006 CONTINUE REWIND 3 NRV=1 1167 IF(NRV-NR) 1168,1168,1100 1168 CONTINUE CALL YOUEH(MOM) DC 1101 I=1.N HTEMP=H(I, IPIV) BC8=CCSINE*HTEMP SC8=-SINE#HTEMP CCE=SINE+H(I, JPIV) F(I, IPIV)=BOB+CCB TGB=COSINE*H(I, JPIV) H(I, JPIV) = SOB+TCB 1101 CONTINUE NRV=NRV+1 GC TO 1167 110C CONTINUE 1005 NM=N-1 CG 742 J=1,NM JP=J+1DC 895 K=1,J IF (HCIAG(K)-HDIAG(JP)) 841,895,895 841 HH=HDIAG(K) HDIAG(K)=HDIAG(JP) HCIAG(JP)=HH DC 1009 I=1,N $H \models H(I,K)$ H(I,K)=H(I,JP)H(I,JP)=HH 1009 CONTINUE 895 CONTINUE 742 CONTINUE RETURN

```
SLEROUTINE YQUEF(MCM)
READS IN THE PACKED INDICES NEEDED FOR EIGENVECTOR COMPUTATION
ANC PUT ON TAPE PREVIOUSLY BY LAFITE AND MOUTON
С
C
       CIMENSION ARRAY(252)
COMMON /CUC/ ARRAY, IPIV, JPIV, COSINE, SINE
     7 IF(MCM-1) 1,2,1
     2 REAC (3) (ARRAY(1),1=1,252)
     1 IF (NCM-252) 5,5,6
    6 MCM=1
GC TC 7
                                                                                       .
     5 IPIV=ARRAY(MOM)
       JPIV=ARRAY(MOM+1)
       CCSINE=ARRAY[MGP+2]
       SINF=ARRAY(MOM+3)
       *C*=*C*+4
       RETURN
       END
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```
SUBROUTINE LAFITE(M)
      PACKS QUANTITIES NECESSARY FOR COMPUTATION OF EIGENVECTORS INTO 
MAXIMUM SIZE RECORDS AND WRITES THEM OUT ON TAPE
С
C
       CIMENSICN ARRAY(252)
      CCMMON /CUC/ ARRAY, IPIV, JPIV, COSINE, SINE
      ARRAY(>)=IPIV
      ARRAY(M+1)=JPIV
      ARRAY(P+2)=COSINE
      ARRAY(M+3)=SINE
      ¥=¥+4
      IF(M-252) 2,2,3
    3 CENTINLE
                  3 ) (ARRAY(I), I=1,252)
      WRITE (
      ¥=1
    2 RETURN
      ENC
```

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SLBRCUTINE MOUTEN C WRITES DUT ON TAPE ANY UNFINISHED RECORD RELATING TO EIGENVECTORS EIMENSICN ARRAY(252) CCMMCN /CUD/ ARRAY, IPIV, JPIV, COSINE, SINE 1 WRITE (3) (ARRAY(I),I=1,252) RETURN ENC

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SUBREUTINE CARUSO(IOCC, U, A, NT)
COMPUTES TOTAL OVERLAP POPULATION MATRIX
С
       U IS IN CORE S ON TAPE IS
C
       CIMENSION U(140,140), A(150), IOCC(150)
       REWIND 8
       REWIND 3
       CC 2 I=1,NT
CC 2 J=1,NT
     2 L[[,J]=C.C
       CC 3 M=1,NT
REAC ( 3)(A(J),J=1,NT)
       RMM=2*IGCC(M)
       IF(RMM) 3,3,4
     4 CC 5 I=1,NT
CC 5 J=1,NT
     5 L(I,J)=U(I,J)+A(I)*A(J)*RMM
    3 CONTINUE
       CC 6 I=1,NT
READ ( 8)(A(J),J=1,NT)
CC 6 J=1,NT
    6 L(I,J)=L(I,J)*A(J)
       RETURN
       ENC
```

SUBROUTINE SOREN THIS IS THE MATRIX EQUATION SOLVER WHICH REPLACES OLD EIGEN AND С č FUNCTIONS WITH ONLY ONE MATRIX IN CORE ASSUMES S IS IN CORE AND H ON TAPE ITAPEH С DIFENSION A(140,140), IOCC(150), HOIAG(150), B(150), C(150) COMMON /DAA/ A /DAB/ IGCC, HDIAG, B, NDIM, N CCMMON /CAN/ ER, SUM COMMON REWIND 8 DG 222 I=1,N 222 WRITE (8) (A(I,J), J=1, N)REWIND 8 DG 1988 NN=1,N READ (8) (E(J), J=1,N) IF(NN-1)2,3,2 3 A(NN.NN)=1. GC TO 1988 2 NNN=NN-1 DC 5988 I=1,NNN A(NN, I)=0. CO 5988 K=1,I 5588 A(NN,I)=A(NN,I)+A(K,I)*B(K) DC 6988 L=1,NNN A(L,NN)=0. DC 6988 I=L,NNN 6988 A(L,NN)=A(L,NN)-A(NN,I)*A(L,I) A(NN,NN) = 1. XNORM=0. CC 4988 J=1.NN 4588 XNCRM=XNORM+A(J,NN)*B(J) XNORM=1./SQRT(XNORM) CC 7988 L=1,NN 7988 A(L,NN)=A(L,NN) *XNORM 1988 CONTINUE DG 3111 I=2,N ING=I-1 DG 3111 J=1,ING 3111 A(I,J)=0. A(2,1)=0. 3112 REWIND 9 CC 3211 I=1,N 3211 WRITE (9) $\{A\{J,I\}, J=I, N\}$ REWIND 4 CC 10 I=1,N 10 READ (4) (A(I,J),J=1,N) CALL BALDUR REMIND 4 CG 6 J=1,N 6 WRITE (4) (A(I,J),I=1,N)REWIND 9 CC 7 J=1,N 7 READ [9] (A(J,IJ,I=1,N) REWIND 9 CC 960 J=1,N

...

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960 WRITE (9)
                   (A(I,J),I=1,N)
     CALL BALL
     CALL CUPID
WRITE ( 6,S)(I,HDIAG(I),IOCC(I),I=1,N)
     SU#=0.0
     DC 1965 I=1,N
1965 SUM=SUM+HDIAG(I)*FLOAT(IOCC(I))
     CRLM=23.07*SUN
     WRITE ( 6,1972)SUM,CRUM
     ERLM=23.07*ER
     WRITE (6,1973) ER, ERUM
                                                                  .
     SLM=SUM+ER
     CRLM=23.07*SUM
     WRITE ( 6,1974JSUM,CRUM
1972 FORMAT(//50X,4HSUM,F16.8,3H EV,20X,2H= ,F16.8,5H KCAL//)
1973 FORMAT(41X,13HR EPUL ENERGY, F16.8,3H EV, 20X,2H= ,F16.8,5H KCAL//)
1974 FORMAT(41X,13HTCTAL ENERGY,F16.8,3H EV,20X,2H= ,F16.8,5H KCAL//)
   8 FORMAT(1H0,51X,2HE(,I3,3H) =,F12.6,6X,I3)
1234 CALL LCKI
     WRITE ( 6,9)
   $ FORMAT(1H0,55X,14HEIGENVECTORS ///)
     CALL PEGLEG(A,N)
     RETURN
     ENC
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SLEROUTINE LOKI
с
с
       CALCULATES A=BC WITH C IN CORE AND B ON TAPE ITAPEB STORED ROWWIS
      RESULTANT MATRIX IN CORE
      CIMENSION C(140,140), IX(150), A(150), B(150)
      CCMMCN /CAA/ C
CCMMON /CAB/IX, A, B, NDIM, N
REWINC 3
      REWIND S
      CC 2 I=1,N
      READ (5)
                       (E(J), J=1, N)
      EC 1 K=1,N
      A(K)=0.0
      CC 1 J=1,N
    1 \quad A(K) = A(K) + B(J) + C(J,K)
    2 WRITE (3)
                     (A(K), K=1, N)
      REWIND 3
    CC 3 I=1,N
3 REAC (3)
                       (C(I,J),J=1,N)
      FETURN
      END
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SUBREUTINE BALDUR CALCULATES A=BC WITH B IN CORE, C ON ITAPEC STORED BY COLUMNS С RESULTANT MATRIX IS IN CORE С DIMENSION B(140,140), IX(150), A(150), C(150) COMMON /CAA/ B COMMON /CAB/IX, A, C, NDIM, N REWIND 3 REWIND 9 CC 2 I=1,N REAC (5) (C(J), J=1, N)EC 1 K=1.N A(K)=0.0 CC 1 J=1,N 1 A(K) = A(K) + B(K, J) + C(J)2 WRITE (3) REWIND 3 (A(K),K=1,N) DC 3 I=1,N 3 REAC (3) (E(J,I),J=1,N) RETURN END

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SLERGUTINE JENNY(ICOUNT, P, N, A)
REELCTICN OF OVERLAP POPULATION MATRIX TO ATOM-ATOM
DIFENSION P(140,140), ICCUNT(50,2), A(150)
   REWIND 9
   CC 1 I=1,N
   IN=ICGUNT(I,1)
   IFF=ICCUNT(1,2)
  DC 2 J=1,N
  A(J)=0.0
   JIN=ICCLNT(J,1)
  JF=ICCUNT(J,2)
  CC 2 K=IN, IFF
  DC 2 L=JIN,JF
2 A(J) = A(J) + P(K,L)
1 WRITE ( S)(A(J), J=1, N)
  REWIND 9
DC 2 I=1,N
3 READ ( S)(P(I,J),J=1,N)
4 FCRMAT(1+0,40X,50F REDUCED OVERLAP POPULATION MATRIX ATOM BY ATOM
 1
                     1/}
  WRITE ( 6,4)
  CALL PEGLEG(P,N)
  RETURN
  ENC
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С

SUBROUTINE COKI CALCULATES A=BC WITH C IN CORE AND B ON TAPE ITAPEB STORED ROWWIS С RESULTANT MATRIX IN CORE С CIMENSION C(140,140), IX(150), A(150), B(150) CCMMON /GAA/ C CCMMCN /CAB/IX, A, B, NDIM, N REWIND 4 REWIND 8 EC 2 I=1,N READ (8) (B(J), J=1, N)CC 1 K=1,N A(K)=C.C CC 1 J=1,N $1 \land (K) = \land (K) + B(J) * C(J,K)$ 2 WRITE (4) (A(K),K=1,N) REWINE 4 CC 3 I=1,N 3 REAC (4) (C(I,J),J=1,N) RETURN END

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SLBRCUTINE BALL CALCULATES A=BC WITH B IN CORE, C CN ITAPEC STORED BY COLUMNS RESULTANT MATRIX IS IN CCRE DIMENSION B(140,140), IX(150), A(150), C(150) CCMMON /CAA/ B CCMMON /CAB/IX, A, C, NDIM, N REWINC 3 REWIND 4 CC 2 I=1,N (C(J), J=1,N) READ (4) DC 1 K=1, h A(K)=0.0 DC 1 J=1,N $1 \Delta(K) = \Delta(K) + B(K, J) + C(J)$ 2 WRITE (3) (A(K),K=1,N) REWIND 3 ED 3 1=1,N 3 READ (3) (E(J,I), J=1,N)RETURN

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END
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SUBROUTINE IENNY(ICOUNT, P, N, A)
Reduction of overlap population matrix to atom-atom
  DIFENSION P(140,140), ICCUNT(50,2), A(150)
  REWIND 4
  CC 1 I=1.N
  IN=ICCUNT(I,1)
  IFF=ICCUNT(I,2)
  DC 2 J=1,N
  A(J)=C-C
   JIN=ICCUNT(J,1)
  JF=ICCLNT(J,2)
  CC 2 K=IN, IFF
  DC 2 L=JIN,JF
2 A(J)=A(J)+P(K,L)
1 WRITE ( 4)(A(J),J=1,N)
  REWIND 4
  CC 3 I=1,N
3 READ ( 4)(P(I, J), J=1, N)
4 FCRMAT(1H0,40X,50H REDUCED DVERLAP POPULATION MATRIX ATOM BY ATOM
                  11)
 1
  WRITE ( 6 ,4)
  CALL PEGLEG(P,N)
  RETURN
  END
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SLERCUTINE REPU
С
      THIS SUBROUTINE CALCULATES AN EXPONENTIAL REPULSION ENERGY.
Ċ
      THIS ENERGY DEPENDS ON THO EMPIRICALLY DETERMINED PARAMETERS.
č
c
      CIFENSION S(140,140), IOCC(150), XX(150), YY(150), AR(150), AL(150)
      CCFFGN /CAA/ S
      CCFMON /CAB/ ICCC, XX, YY, N, NDIM
      CEMMON ZEAMZ AR, AL, MR
      CCMMCN /CAN/ ER
      ER=C
      IF (MR) 4,4,2
    2 K=N-1
      CC 3 I=1,K
      L=I+1
      CC_3_J=L,N
ER=ER+SQRT(AR(I)*AR(J))*EXP(-S(I,J)/(AL(I)+AL(J)))
    3 CONTINUE
    4 RETURN
      END
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