Variable Energy Photoelectron Spectroscopy: Periodic Trends in d-Orbital Energies for Organometallic Compounds of the Transition Metals

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Introduction

Molecular photoelectron spectroscopy (PE) came into prominence with the studies of Turner and his co-workers in the early 1960s.¹ It was shown that when a photon beam irradiates a molecule, and if the photon energy (hv) is larger than the binding energy (E_b or BE) of a molecular orbital (MO), the electrons in that MO will be ejected. On the basis of the relationship $hv = E_b + E_k$ a photoelectron spectrometer sorts the binding energies of molecular orbitals according to the different kinetic energies (E_k) of the outgoing electrons. Usually, in the outer valence levels of an organometallic molecule, there is a one-to-one correspondence between the binding energies and molecular orbital energies (Koopmans' theorem). Therefore, photoelectron spectroscopy is the most direct way to probe the orbital ordering of molecules.

What kind of information can we get from the photoelectron spectra of a molecule if these spectra are recorded at different photon energies? It was recognized early on by Price² that the binding energies provided by a photoelectron spectrum are independent of the photon energy employed, but band intensities (photoionization cross sections) vary with photon energy. The band intensity variations as a function of photon energy can disclose important information on the orbital origin of the ejected electrons. In his pioneering work, Price used He I (21.2 eV) and He II (40.8 eV) photons to explore the correlation between band intensity and photon energy.

Synchrotron radiation provides the intense tunable source of photons needed to study the continuous variation of atomic and molecular photoionization cross sections with photon energy. The increasing availability of this radiation, coupled with the advent of high-resolution electron spectrometers, has greatly enhanced our ability to make orbital assignments for PE spectra.³

Only in the last nine years (starting in 1987) has variable energy photoelectron spectroscopy been applied to organometallics, including $M(CO)_6$ (M = Cr, Mo, W),⁴ $M(\eta^5$ - $C_5H_5)_2$ (M = Fe, Ru, Os),⁵ U(η^8 - $C_8H_8)_2$,⁶ (η^5 - C_5H_5)PtMe₃,⁷ $(\eta^7 - C_7 H_7) M(\eta^5 - C_5 H_5)$ (M = Ti, Nb, Mo),⁸ $(\eta^7 - C_7 H_7) Ta(\eta^5 - C$ C_5H_4Me),⁸ $Cr(\eta^6-C_6H_6)_2$,⁹ $Mo(\eta^6-C_6H_5Me)_2$,⁹ $M(\eta^5-C_5H_5)_2$ (M = V, Cr, Co, Ni),¹⁰ and $[(\eta^5 - C_5 H_4^i Pr) MoS]_4$,¹¹ and it has been shown that the information content from a PE experiment is greatly enhanced in such studies. The merits of using synchrotron radiation to study the PE spectra of these molecules have been well documented by Green.³ However, prior to our work, variable energy photoelectron studies, along with related theoretical developments on photoionization cross sections, have not been reported for systematic studies of organometallic compounds of the late transition metals. For some important organometallic molecules of these metals, such as $M(\eta^3-C_3H_5)_2$ (M = Ni, Pd, Pt), the MO ordering assigned from photoelectron spectra and theoretical calculations has been very controversial.¹² The sharp difference of cross section behavior between the ligand orbital ionizations and metal d ioniza-

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Variable Energy Photoelectron Spectroscopy Li et al.



FIGURE 1. Theoretical photoionization cross sections of metal d and C 2p orbitals.¹³

tions made it possible for us to use synchrotron radiation to reassign the MO ordering of these molecules, and thus enabled us to reveal periodic trends in the molecular orbital energies of these molecules. These periodic trends, which have not been reviewed in the literature, will be summarized in this Account.

MO Assignments of Photoelectron Spectra Using Band Intensity Variations

With variable energy photoelectron spectroscopy, we monitor the photoelectron band intensities as a function of photon energy, usually between \sim 20 and \sim 150 eV. The relative band intensities vary because the atomic cross sections (and thus the molecular orbital cross sections) have very different trends with photon energy. For example, Figure 1 shows the theoretical cross sections as a function of photon energy (from 20 to 170 eV) for the carbon 2p and d orbitals of the late transition metals. These atomic orbitals form the major components of the outer valence region of organometallic molecules.¹³ Obviously, the cross section trends for the metal d and C 2p orbitals are very different. For example, the C 2p cross sections (along with other common ligand orbital cross sections such as N 2p and O 2p) show a rapid monotonic decay above the threshold. In contrast, the metal d cross sections all show an initial increase in cross section above the threshold (so-called delayed maxima), followed by a decrease. The rate of decrease in cross section is much smaller for the first-row transition metals compared to the second- and third-row metals, while the second-row metal 4d cross section goes through a minimum (Cooper minimum) at >100 eV.

A model (so-called Gelius model¹⁴) which is important for variable energy photoelectron spectroscopy states that the trend of photoionization cross sections as a function of photon energy for a molecular orbital is mainly determined by that of its predominant atomic orbital components.¹⁴ For organometallic compounds, the cross section variation of nonbonding metal d and ligand carbon 2p orbitals should behave like their atomic counterparts, and those of the bonding MOs behave in an intermediate manner. This statement is more reliable quantitatively when the photon energy is large (>50 eV), and when the magnitude of one atomic orbital cross section is dominant over all the other atomic orbital contributions.¹⁵ This rule has been successfully applied in our variable energy photoelectron spectroscopic studies of organometallic compounds of the late transition metals, such as $M(\eta^3-C_3H_5)_2$ (M = Ni, Pd, Pt).¹⁶ The above study was particularly successful because there are a few "nonbonding" ligand

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FIGURE 2. X α -SW MO diagrams for the formation of the outer valence of (η^{5} -C₅H₅)Pd(η^{3} -C₃H₅). The Pd 4d compositions are put beside the MOs of (η^{5} -C₅H₅)Pd(η^{3} -C₃H₅).

and metal d orbitals in the outer valence of these molecules, which makes the intensity changes of many bands in the low binding energy region behave like atomic metal d and carbon 2p orbitals. The sharp difference in the trends of cross section between metal d and ligand orbitals (Figure 1) allow us to assign these PE spectra with confidence. The above cross section rule is also supported by X α -SW cross section calculations for MOs for M(η^3 -C₃H₅)₂.¹⁶ The calculations indeed show that the nonbonding MOs behave like their atomic counterparts, and bonding MOs show intermediate behavior.

To illustrate the power of the variable energy approach, we look first at the PE spectrum of $(\eta^5-C_5H_5)Pd(\eta^3-C_3H_5)$.¹⁷ The reasons for us to use this molecule as an example are (1) the bands in the outer valence of its PE spectrum are well resolved, so the one-to-one relationship between photoelectron bands and molecular orbitals can be best seen; (2) there is a fairly broad distribution of metal d compositions among the MOs in the outer valence region, i.e., some MOs are Pd 4d based, some are mainly bonding MOs between Pd 4d and Cp, some are nonbonding Cp π_1 and ally π_1 , and one MO is from bonding between PdCp and allyl. The sequence of Pd percent compositions for these MOs is Pd 4d based > Pd-Cp bonding > Pd-Cpallyl bonding and π_1 MOs of allyl and Cp. This variation of metal d compositions has been disclosed very clearly by the different degrees of band intensity variations observed from the variable energy photoelectron spectra of this molecule.

Figure 2 shows a X α -SW MO diagram showing the formation of the MOs of (η^5 -C₅H₅)Pd(η^3 -C₃H₅) between PdCp and allyl fragments. The calculated Pd 4d compositions are shown beside each orbital. Among the nine occupied orbitals in the outer valence region, there are three orbitals (15a', 9a'', and 14a') whose Pd 4d compositions are >82%. They are designated the Pd 4d based MOs. These MOs are located in the middle part of the outer valence region. Among other MOs, there are three

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The above orbital ordering is confirmed by the variable energy photoelectron spectra of $(\eta^5-C_5H_5)Pd(\eta^3-C_3H_5)$ (Figure 3) recorded at 21.2, 40, 50, and 60 eV, respectively, covering the photon energy range in which the Pd 4d delayed maximum takes major effect. The spectra display eight bands corresponding to the first eight MOs in the outer valence region. The three bands (bands 3, 4, and 5) in the middle binding energy region increase in intensity with an increase in photon energy. They are assigned to the three Pd 4d based orbitals, as predicted from the MO diagram and the Gelius model. All other bands decrease in relative intensity, with bands 2 and 8 decreasing more than bands 1, 6, and 7. Therefore, it is reasonable to assign bands 2 and 8 to the MOs with the smallest Pd 4d character (band 2 to 10a" and band 8 to 12a'), and bands 1, 6, and 7 must be assigned to the three MOs with intermediate Pd 4d character (16a', 8a", and 13a', respectively).

In the $(\eta^5-C_5H_5)Pd(\eta^3-C_3H_5)$ case, the calculated orbital ordering is in agreement with that deduced from the variable energy spectra, and the ratios of intensity variation for each band are also consistent with the relative Pd 4d compositions calculated with the X α -SW method, strongly suggesting that the Gelius model has semiquantitative significance for MOs of variable Pd 4d composition. The variable energy technique, combined with MO calculations, is obviously a powerful method to confirm MO assignments.

Although the traditional He I (21.2 eV)/He II (40.8 eV) ratio of band intensity can help to assign PE spectra for some organometallic compounds, the low-energy and noncontinuous feature of helium light limits its usefulness. For example, in our studies for PE spectra of copper(I) β -diketonate complexes,¹⁸ we noticed that the mainly Cu 3d band cannot be assigned unambiguously by using the He I/He II intensity ratio. The Cu 3d band only shows the expected large increase in relative intensity above 40.8 eV (He II radiation). Only by taking spectra above 50 eV does the Cu 3d band assignment become very obvious.

It is also important to emphasise that, often, the energy ordering determined by the variable energy technique

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VOL. 30, NO. 5. 1997 / ACCOUNTS OF CHEMICAL RESEARCH 215



FIGURE 3. Representative variable energy photoelectron spectra of $(\eta^{5}$ -C₅H₅)Pd $(\eta^{3}$ -C₃H₅) recorded at 21.22, 40, 50, and 60 eV. The X α -SW Pd 4d compositions (%) are given at the top of each band. Band numbers are at the bottom with the Pd 4d based bands underlined.

does not agree with the ordering from MO calculations. For example, for Ni(η^3 -C₃H₅)₂, the orbital ordering did not agree with the ordering from any theoretical calculation—either our X α -SW calculation or many others!¹⁶ The variable energy method seems much more reliable than MO calculations for transition metal complexes.

Periodic Trends

To determine periodic trends in metal d orbital energies, it is highly desirable to have volatile gas phase analogues for a given transition metal group available, because the PE resolution in the gas phase is generally much better than for solid state spectra. Fortunately, stable volatile analogues are now available, and the cross section variations described above make it rather easy to identify the metal d levels.

General trends in metal d energies can be seen immediately by looking at a few spectra. For the early transition metal organometallic compounds, the PE spectra down a group are very similar. For example, the He I spectra of $M(CO)_6$ molecules (M = Cr, Mo, W)¹⁹ are shown in Figure 4. These spectra are qualitatively very similar, and even the metal d energies vary only by 0.16 eV from $Cr(CO)_6$ to $W(CO)_6$. If we now go to the next group (Figure

5), the spectra of $CpM(CO)_3$ analogues (M = Mn, Re)²⁰ are still qualitatively similar, but the metal d energies now differ by ~ 0.34 eV. In contrast, when we turn to the spectra of organometallic compounds of the later transition metals, some striking differences occur. Figure 6 shows the outer valence PE spectra of $M(\eta^3-C_3H_5)_2$ (M = Ni, Pd, Pt)¹⁶ and CpM(CO)₂ (M = Co, Rh, Ir)²¹ all recorded with He I photons. It is immediately apparent that the spectra of the first-row Ni and Co complexes differ greatly from the spectra of the second- and third-row analogues. For example, the Ni(η^3 -C₃H₅)₂ spectrum shows three resolvable peaks at 8 ± 0.4 eV BE, whereas the Pd and Pt analogues only show one peak in this region. But at higher binding energy (8.5-12 eV), there are many more peaks in the Pd and Pt spectra than in the Ni analogue. The variable energy spectra (Figure 7) show that peak 1 in the spectra of $Pd(\eta^3-C_3H_5)_2$ must be due to a mainly ligand orbital because the intensity of peak 1 decreases from 24 to 80 eV photon energy. In contrast, peaks 1 and 2 for Ni(η^3 -C₃H₅)₂ must be due mainly to Ni 3d orbitals because their intensity increases with photon energy. Thus, the Ni 3d orbitals must have $a \ge 1$ eV lower BE than the Pd 4d BE for the Pd analogue.

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FIGURE 4. He I spectra of M(CO)₆ (M = Cr, Mo, and W).^{19a} The metal nd (η = 3, 4, 5) bands and their binding energies are specified.



FIGURE 5. PE spectra of $(\eta^{5}-C_{5}H_{5})M(CO)_{3}$ (M = Mn and Re) recorded at 60 eV photon energy. The binding energies of metal 3d (Mn) and 5d (Re) based bands are specified.

It is also very important from Figure 6 that the secondand third-row spectra of analogues are very similar, with the spectra of $Pt(\eta^3-C_3H_5)_2$ and $CpIr(CO)_2$ showing one more resolved peak than $Pd(\eta^3-C_3H_5)_2$ and $CpRh(CO)_2$ respectively. Even the order of orbital assignments is the same, with the symmetry and character of orbitals in the second- and third-row complexes being identical. For example, assignments of bands, 1, 2, 3, 4, 5, and 7 for Pd- $(\eta^3-C_3H_5)_2$ are the same as these from bands 1, 2, 3, 4, 5, and 8 for Pt $(\eta^3-C_3H_5)_2$. Band 6 of Pd $(\eta^3-C_3H_5)_2$ is broad and contains two MOs corresponding to bands 6 and 7 of Pt $(\eta^3-C_3H_5)_2$.

From the above discussion, the qualitative differences in spectra are due mainly to the shift in Ni 3d and Co 3d binding energies to much lower values. Indeed, the ligand based orbitals, such as bands 5 and 6 for Ni(η^3 -C₃H₅)₂, have a BE very similar to those of the corresponding ligand orbitals in the Pd (bands 6 and 7) and Pt (bands 7 and 8) analogues.

The orbital sequences for the three compounds from the variable energy technique can now be summarized (with the band number in parentheses and the main character of the Mo-either metal (M) or ligand (L)following). Ni $(\eta^3$ -C₃H₅)₂: 13a_g (1, M) < 12a_g (1, M) < 6b_g $(2, M) < 7a_u (2, L) < 11a_g (3, M) < 5b_g (4, M) < 11b_u (5, L)$ < $10a_g$ (6, L). Pd(η^3 -C₃H₅)₂: $8a_u$ (1, L) < $17a_g$ (2, M) < $16a_g (3, M) < 8b_g (4, M) < 15a_g (5, M) < 7b_g (6, M) < 13b_u$ $(6, L) < 14a_g (7, L). Pt(\eta^3-C_3H_5)_2: 12a_u (1, L) < 21a_g (2, M)$ $< 20a_g$ (3, M) $< 10b_g$ (4, M) $< 19a_g$ (5, M) $< 9b_g$ (6, M) < $19b_u$ (7, L) < $18a_g$ (8, L). From the spectra in Figures 4–6, it is apparent that the average binding energy difference $(\Delta E_{\rm b})$ between metal 3d (first-row) and 4d (second-row) orbitals is much larger for the late transition metals than the early transition metals. The sequence of $\Delta E_{\rm b}$ is $M(CO)_6$ (M = Cr and Mo, 0.10 eV)¹⁹ < CpM(CO)₃ (M = Mn and Re, 0.34 eV)²⁰ < MCp₂ (M = Fe and Ru, 0.59 eV)²² < CpM(CO)₂ (M = Co and Rh, **0.71 eV**)²¹ < M(η^3 -C₃H₅)₂ $(M = Ni \text{ and } Pd, 1.17 \text{ eV})^{16} < (hfac)MPMe_3$ (M = Cu and Ag, hfac = $CF_3C(O)CHC(O)CF_3$, **1.9 eV**).¹⁸

Origin of the Difference in Metal d Ionizations

What is the reason for this trend, in which the energy difference between3d and 4d or 5d orbitals for analogous compounds appears to increase for the later transition metals? Lichtenberger et al.^{23,24} attributed the d binding energy separation in $CpM(CO)_2$ (M = Co and Rh) mainly to the larger relaxation energy associated with first-row complexes. The orbital relaxation effect is mainly due to the orbital contraction on ionization. This effect makes the ion state more stable relatively, and lowers the BE. On the other hand, Ziegler et al.25 proposed that trends in the thermal stability and kinetic lability of the metalcarbonyl bond in $M(CO)_6$ (M = Cr, Mo, and W), $M(CO)_5$ (M = Fe, Ru, and Os), and $M(CO)_4$ (M = Ni, Pd, and Pt)were due to the ground state energy differences between the first- and second- or third-row transition metals. They suggested that, in the ground state, the 4d and 5d metal orbitals are lower in energy (higher in BE) than the 3d orbitals, since d-d repulsions are smaller for the diffuse 4d and 5d orbitals than for the contracted 3d orbitals and that this difference should increase from the early to the

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FIGURE 6. He I spectra of M(η^3 -C₃H₅)₂ (M = Ni, Pd, and Pt) and (η^5 -C₅H₅)M(CO)₂ (M = Co, Rh, and Ir).

FIGURE 7. Band intensity increase with photon energy for Pd 4d and Ni 3d based bands in the spectra of $M(\eta^3-C_3H_5)_2$ (M = Ni and Pd). The photon energy range for the Pd compound is covering that for the Pd 4d delayed maximum.

late transition metals. Due to the "lanthanide contraction", the energies of 4d and 5d metal orbitals are relatively close. The Xa calculations carried out for CpM(CO)2 (M = Co, Rh, and Ir)²¹ and M(η^3 -C₃H₅)₂ (M = Ni, Pd, and Pt)¹⁶ in our work support Ziegler's interpretation in terms of the difference in ground state d orbital energies. It is also supported by the difference in slopes among experimental curves of relative band intensity variation (branching ratio) for the HOMOs of $CpM(CO)_2$ (M = Co, Rh, and Ir).²¹ The HOMOs of these molecules are the antibonding orbitals from the interaction of M(CO)₂ with Cp. The slope

difference suggests that the HOMO of the Co compound has more metal d character than those of the Rh and Ir analogues, thus supporting the ground-state d BE energy sequence 3d (Co) < 4d (Rh) \approx 5d (Ir).

Concluding Remarks

The large metal d binding energy difference between the first and second or third late transition rows has been confirmed by variable energy photoelectron spectroscopy. Its origin, most likely a ground-state d energy difference, may help us to understand some special features associated with the late transition metals, such as the relative σ and π bonding between metal and carbonyl ligands,²¹ and the very different reactions of olefins on Ni versus Pt surfaces (which may be largely due to the enhanced metal-olefin π bonding on Ni surfaces).^{26,27} The periodic trends of d energy separations for the late transition metals should be one of the essential factors for comprehending the different types of chemical reactions and catalytic processes associated with these metals.

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