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CORE-ELECTRON BINDING ENERGIES FOR COMPOUNDS OF BORON, CARBON AND CHROMIUM

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## Authors

Henrickson, D.N. Hollander, J.M. Jolly, W.L.

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### CORE-ELECTRON BINDING ENERGIES FOR COMPOUNDS OF BORON, CARBON AND CHROMIUM

D. N. Henrickson, J. M. Hollander, and W. L. Jolly

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of Boron, Carbon and Chromium

D. N. Hendrickson, J. M. Hollander, and W. L. Jolly

Contribution from the Department of Chemistry of the University of California and the Inorganic Materials Research and Nuclear Chemistry Divisions of the Lawrence Radiation Laboratory, Berkeley, California 94720

#### ABSTRACT

Boron <u>ls</u> and chromium <u>3p</u> binding energies have been measured for 25 boron compounds and 17 chromium compounds. The data correlate linearly with calculated atomic charges. A linear correlation is also demonstrated for CNDO-calculated atomic charges and carbon <u>ls</u> binding energies determined by other workers.

#### INTRODUCTION

Core-electron binding energies for several elements in a wide variety of compounds have been determined by X-ray photoelectron spectroscopy (XPS).<sup>1</sup>

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A review of many aspects of XPS can be found in K. Siegbahn, C. Nordling,
 A. Fahlman, R. Nordberg, K. Hamrin, J. Hedman, G. Johansson, T. Bergmark,
 S.-E. Karlsson, J. Lindgren and B. Lindberg, "ESCA Atomic Molecular and
 Solid State Structure Studied by Means of Electron Spectroscopy," Almqvist
 and Wiksells AB, Stockholm (1967).

These energies have been correlated with atomic charges calculated from molecular orbital eigenfunctions.<sup>2-4</sup> Although it has been pointed<sup>5</sup> out that, from

(2) For nitrogen compounds see: J. M. Hollander, D. N. Hendrickson, and
W. L. Jolly, <u>J. Chem. Phys.</u>, <u>49</u>, 3315 (1968); D. N. Hendrickson, J. M.
Hollander, and W. L. Jolly, submitted to Inorg. Chem.

(3) For carbon compounds see: R. Nordberg, U. Gelius, P. F. Hedén,
J. Hedman, C. Nordling, K. Siegbahn, and B. J. Lindberg, submitted to
Arkiv. för Kemi.

(4) For phosphorus compounds see: M. Pelavin, D. N. Hendrickson, J. M. Hollander, and W. L. Jolly, submitted to <u>J. Phys. Chem</u>.

(5) W. L. Jolly and D. N. Hendrickson, to be published.

theoretical considerations, one cannot expect better than a rough correlation between core-electron binding energy and atomic charge, such rough correlations are fairly easy to make and can give useful information about the bonding and structure of compounds. In this paper we report boron <u>ls</u> and chromium <u>3p</u> binding energies and show that they can be roughly correlated with atomic charges calculated by various molecular orbital methods. A CNDO-calculated charge correlation is also demonstrated for some carbon <u>ls</u> binding energies that were determined in another work.<sup>3</sup>

### EXPERIMENTAL

The experimental aspects of the determination of inner-electron binding energies have been reviewed by Siegbahn <u>et al</u>.<sup>1</sup> MgK<sub>a</sub> X-radiation (1253.6 eV) was used in this work. All compounds were solid and were treated as before<sup>2</sup> except where noted below.

It was necessary to cool the samples of  $(CH_3)_3 NBF_3$ ,  $B_{10}H_{14}$  and  $Cr(CO)_6$  to prevent their sublimation in the spectrometer. These samples were loaded into the spectrometer source house under dry nitrogen and cooled to ~ -60° before evacuation.

Photoelectrons originate from sites near the surface,<sup>1</sup> thus it was necessary to continuously clean the surface of a piece of chromium foil in order to see a peak that could be assigned to the unoxidized metal. This was accomplished by heating the chromium foil to ~ 650° in hydrogen at  $10^{-2}$  torr. The disappearance of the oxide layer was monitored by the 0 ls line. Chromium hydride formation need not be considered because of the low solubility of hydrogen in chromium metal at 650°.<sup>6</sup>

(6) T. R. P. Gibb, Jr., Progress in Inorganic Chemistry, 315 (1962).

-3-

The work function for the spectrometer material (aluminum) was assigned the value 4.0 eV, as in previous work.<sup>2,4</sup>

Compounds were either purchased or were gifts. Samples of many boron compounds were kindly supplied by Professor M. F. Hawthorne and Dr. S. Trofimenko.

#### CALCULATIONS

CNDO molecular orbital eigenfunctions<sup>7</sup> were obtained for some of the carbon and boron compounds. Simple extended Hückel calculations<sup>8</sup> were also made for some of the boron compounds. In the extended

(7) J. A. Pople, D. P. Santry, and G. A. Segal, J. Chem. Phys., 43,
Sl29, Sl36 (1965). A modified CNDO/l version involving empirically
evaluated repulsion integrals was used, as per P. M. Kuznesof and
D. F. Shriver, J. Am. Chem. Soc., 90, 1683 (1968). Calculational details
can be found in this latter reference.

Hückel method, Coulomb integrals, H<sub>ii</sub>, were approximated by the valence state ionization potentials (vsip) calculated by Hinze and Jaffé.<sup>9</sup> The off-diagonal elements were assigned as

$$H_{ij} = 1.75 S_{ij}(H_{ii} + H_{jj})/2$$

where  $S_{ij}$  is the overlap integral between the ith and jth atomic orbitals. Slater's rules<sup>10</sup> were used to calculate orbital exponents for atomic orbitals. Boron nitride was treated in an approximate manner.<sup>2</sup>

- (8) R. Hoffmann, <u>J</u>. <u>Chem</u>. <u>Phys</u>., <u>39</u>, 1397 (1963).
- (9) J. Hinze and H. H. Jaffé, J. Am. Chem. Soc., 84, 540 (1962).
- (10) J. C. Slater, Phys. <u>Rev.</u>, <u>36</u>, 57 (1930).

Simple iterative extended Huckel molecular orbital calculations<sup>11</sup> were completed for a few chromium compounds. Coulomb integrals for ligand orbitals were set equal to the negative of the appropriate neutral atom vsip's<sup>9</sup> corrected in each cycle for net atomic charge  $q_i$ .

$$H_{ii} = H_{ii}^{\circ} - (2.0 \text{ eV/charge})q_i \qquad (1)$$

Method was essentially that used for phosphorus compounds by P: C.
 Van der Voorn and R. S. Drago, J. Am. Chem. Soc., 88, 3255 (1966).

Chromium coulomb integrals (except where noted) were calculated for a neutral atom in a  $d^5p$  configuration;<sup>12</sup> charge correction of these values was made using equation (1) on each iterative cycle. Both the metal

-4-

(12) H. Basch, A. Viste, and H. B. Gray, <u>Theoret</u>. <u>Chim</u>. <u>Acta</u> (Berl.) <u>3</u>, 458 (1965).

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and ligand orbital exponents were taken as neutral atom orbital values,  $\mu_i^{\circ}$ , (assigned by Slater's rules<sup>10</sup>) corrected for charge

$$\mu_{i} = \mu_{i}^{\circ} + 0.35 q_{i}/n^{*}$$

Here  $n^*$  is the effective principal quantum number. The off-diagonal Hamiltonian elements were assigned by Cusachs' approximation<sup>13</sup>

 $H_{ij} = S_{ij}(H_{ii} + H_{jj}) (2 - |S_{ij}|)/2$ 

(13) L. Cusachs, J. Chem. Phys., <u>44</u>, 835 (1966).

Atomic charges q were obtained in each cycle by an application of Mulliken's population analysis.<sup>14</sup> Iterations were made on the chromium

(14) R. S. Mulliken, J. Chem. Phys., 23, 1833, 1841, 2338, 2343 (1955).

compounds until the atomic charges were self-consistent to at least 0.05.

V

Program PROXYZ<sup>15</sup> was used to obtain cartesian coordinates when necessary; the molecular parameters were obtained from crystal structure determinations<sup>16</sup> or from estimates.

(15) P. M. Kuznesof, Quantum Chemistry Program Exchange (Indiana University) QCPE 94 (1966).

(16) "Table of Interatomic Distances," L. E. Sutton ed., Special
Publication No.11, The Chemical Society, Burlington House London, 1958
and No. 18, Supplement, 1965.

Core-electron binding energies (the differences in energy between the Fermi level and the atomic level for the solids) were calculated from the relation<sup>1</sup>

 $E_{B} = E_{h\nu} - E_{kin} - \phi_{sp}$ 

Here  $E_{h\nu}$  is the X-ray energy,  $E_{kin}$  is the kinetic energy of the photoelectron in the spectrometer and  $\phi_{sp}$  is the work function of the spectrometer material.

#### RESULTS AND DISCUSSION

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Boron Compounds. Boron 1s electron binding energies were measured for some 25 compounds and are listed in Tables I and II. The range of boron 1s shifts is 8.4 eV, which is comparable to the observed phosphorus 2p shift range (~ 8.5 eV).<sup>4</sup> Larger shift ranges have been observed for nitrogen (~ 10.8 eV),<sup>2</sup> carbon (~ 15.9 eV),<sup>3</sup> and sulfur (~ 15 eV).<sup>1</sup>

Simple extended Hückel molecular orbital (EHMO) calculations were completed for 12 of the boron compounds. A fair linear correlation between measured boron <u>ls</u> electron binding energy and boron atom charge from the EHMO eigenfunctions is shown in Figure 1. The tetramethylammonium salt of the octahydrotriborate ion  $(B_3H_8)$  with two geometrically-different borons gave a single broad boron <u>ls</u> peak with no structure, and therefore only a single point represents this ion in Figure 1. The boron atom charges (via the EHMO method) range from near neutrality in the borohydrides to almost +2.4 in  $BF_h$ .

In our nitrogen studies<sup>2</sup> we found that the EHMO method tended to overemphasize charge separations in molecules. The CNDO nitrogen atom charge range was 4 - 5 times smaller than that obtained by the EHMO method for a series of nitrogen compounds. CNDO calculations were made for some of the boron compounds treated by the EHMO method. The results are given in Table I. Again the CNDO-calculated atom charge range is about 4 times smaller than that obtained using EHMO eigenfunctions. The greater charge separations obtained using the simple EHMO method are probably the result of using charge-independent Coulomb integrals.

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A fairly good correlation exists between the measured boron lsbinding energies and the CNDO-calculated boron charges, as can be seen in Figure 2. Using this CNDO correlation line and the boron binding energy for BN, we infer a boron charge of ~ +0.29. The measured nitrogen <u>ls</u> binding energy for BN<sup>2</sup> correlates with a nitrogen charge of -0.25. This result is a good indication of internal consistency; the sum of the charges should equal zero. Unfortunately, the number of boron compounds was insufficient to permit the distinguishing of two lines (one characteristic of anions and the other of neutral molecules), as observed in our nitrogen work.<sup>2</sup> However it is perhaps significant that the points for the three neutral molecules (2,3, and 6) fall below the line in Figure 2.

For all of the compounds listed in Table II it was possible to detect only a single relatively sharp boron peak. The carboranes (compounds 16-20) have geometrically-different boron atoms, but in an unsubstituted carborane (as in compound 20 where two 1,2-dicarbollide ions are bound to a single cobalt atom) delocalization of charge would be expected to make the charges on the boron atoms similar. It can be seen that as we substitute one S atom for a H atom on the 1,2-dicarbollide ion (as in compound 17) the boron ls binding energy is increased appreciably. Substitution of three bromines for three hydrogens on this same cage increases the boron binding energy even more, but still only one sharp peak can  $b \ge$  seen. Presumably delocalization of charge occurs in this 1,2-dicarbollide cage.

Carbon Compounds. Nordberg et al<sup>3</sup> have recently reported carbon 1s binding energies. They found that the carbon binding energies roughly correlate with carbon fractional atomic charges calculated by a modification of Pauling's method and with carbon atom charges calculated from simple EHMO eigenfunctions. The results of our CNDO calculations for some of the compounds treated by Nordberg et al 3 are given in Table III. The carbon 1s binding energies for these compounds are plotted vs. the CNDO-calculated carbon charges in Figure 3. In the cases of  $CF_{\underline{h}}$  and  $CHF_2$ , the binding energies obtained by Professor D. Thomas (measured both vs. Ne as well as  $CH_{\mu}$ ) were found to fit the correlation better and were used instead of the values given by Nordberg et al. We draw two conclusions. First, the CNDO carbon charges seem more realistic (i.e. not as large) than those obtained by the EHMO method. Second, a somewhat improved straight-line correlation is obtained in the CNDO plot. Only two points fall off the line (i.e. not within experimental uncertainty), the points for KCN and Na2CO3.

<u>Chromium Compounds</u>. Chromium <u>3p</u> electron binding energies are given in Table IV. Qualitatively the chromium <u>3p</u> binding energies increase with increasing oxidation state, the exception being  $Cr(CO)_6$ . Chromium <u>3p</u> binding energies are somewhat more difficult to obtain than nitrogen, carbon, phosphorus or boron core-electron binding energies, because the lower energies associated with ionization of a chromium <u>3p</u> electron give a lower photoionization cross section when using the same X-radiation.<sup>1</sup> Equally troublesome is the asymmetric background due probably to bremstrahlung (inelastically scattered electrons).

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Some preliminary iterative extended Hückel calculations, (see calculation section) were completed for a few of the chromium compounds; the results are given in Table V. In all cases the configuration of the chromium atom was assumed, and the calculations were iterated until all atom charges in the molecule were self-consistent to within 0.05. Two modifications of the metal input parameters were considered--one using Slater orbital exponents<sup>10</sup> coupled with neutral atom chromium valence orbital ionization potentials for  $\frac{d^5p}{p}$  configuration<sup>12</sup>, and the other where essentially the metal orbital exponents were changed. In both cases (see Figure 4) only a very rough correlation between chromium 3p binding energy and calculated chromium atom charge was found.

The chromium 3<u>p</u> binding energies of  $K_3[Cr(CN)_6]$  and  $K_3[Cr(CN)_5NO]$ are identical and therefore one might conclude that the latter compound is a Cr(III) compound, contrary to its usual formulation as a Cr(I) compound.<sup>17</sup> The assignment of the +3 oxidation state to chromium

(17) This conclusion has also been reached by R. D. Feltham on the basis of bond lengths, etc. (private communication).

corresponds to a formal NO<sup>-</sup> group. This conclusion is consistent with our nitrogen ls work, where we found the charge on the NO-nitrogen in  $K_2[Cr(CN)_5NO]$  to be more negative than expected for an NO<sup>+</sup> group.<sup>2</sup>

Acknowledgment. This work was supported by the U. S. Atomic Energy Commission. Our gratitude is due Professor M. F. Hawthorne for gifts of many interesting boron compounds.

### Table I

## Boron 1s Binding Energies and Calculated Charges

Compound		Binding Energy,	Calculated Boron Atom Charge	
Number	Compound	eV.	CNDO	Extended Hückel
1	NaBF4	195.1	+0.754	+2.361
Ź	(CH <sub>3</sub> ) <sub>3</sub> NBF <sub>3</sub>	193.8	+0.756	+2.161
3	B(OH) <sub>3</sub>	193.2	+0.651	+2.122
<b>1</b> 4	NaBH(OCH3)3	192.3	+0.404	+1.724
5	Na <sub>3</sub> <sup>B</sup> 36	192.2	+0.321	+2.098
6	р-F-C6 <sup>H</sup> 4 <sup>B</sup> (OH)2	191.9	+0.498	+1.715
7	p-Cl-C6H4B(OH)2	191.9		+1.713
8	$[c_{2}^{H}_{5}B-(10)]_{N-N}$ ] $_{3}^{-Bc_{2}}B_{5}^{H}_{5}$ ] $PF_{6}$	191.9		+1.228
9	BN	190.2		+1.48
ĺO	K2 <sup>BH</sup> 3CO2	187.5	-0.024	+1.604
ĺÌ	(CH <sub>3</sub> ) <sub>4</sub> NB <sub>3</sub> H <sub>8</sub>	187.4	+0.165 <sup>a</sup> -0.007	+0.518 <sup>a</sup> +0.078
12	$\operatorname{NaBH}_{\mu}$	187.4	0.000	+0.138
13	B amor.	187.5	≡ 0	≡ 0

<sup>a</sup> There are two structurally-different B atoms in the  $B_3H_8$  ion; the first boror charge refers to the unique boron. For purposes of plotting a weighted average was used.

# Table II

# Boron 1s Binding Energies

Compound Number	Compound	Binding Energy, eV.
14	$Na_2B_40$ , $10H_20$	192.8
15	$\left( \sum_{N-N-B} \left( \begin{array}{c} O \\ N-N \end{array} \right)_{3} - W(CO)_{2}(NO) \right)_{3}$	191.9
16	(CH3)4N[(8,9,12-Br3-1,2-	189.5
	в <sub>9</sub> с <sub>2</sub> н <sub>8</sub> ) <sub>2</sub> со]	· · · · · ·
17	$Cs[Co(1,2-B_9C_2H_{10}-8-S)_2]$	188.9
18	Rb[Co(1,2-B,C2H1)/B8C2H0)]	188.4
19	C2 <sup>B</sup> 9 <sup>H</sup> 11 <sup>·C</sup> 5 <sup>H</sup> 5 <sup>N</sup>	188.4
20	Cs[(1,2-B <sub>9</sub> C <sub>2</sub> H <sub>11</sub> ) <sub>2</sub> Co]	188.2
21	<sup>B</sup> 10 <sup>H</sup> 1 <sup>4</sup>	188.0
22	Na2 <sup>B</sup> 12 <sup>H</sup> 12 <sup>•</sup> xH2 <sup>0</sup>	187.7
23	$NaB(C_{6}^{H}_{5})_{4}$	187.7
24	Cs2 <sup>B</sup> 10 <sup>H</sup> 10	187.6
25	в <sub>ц</sub> С	186.7
· .		

## Table III

# Carbon 1s Binding Energies and Calculated Charges

Compound Number	Compound	Binding Energy, eV. <sup>a</sup>	CNDO-Calculated Carbon Atom Charge
1	СF <sub>Ц</sub>	300.7 (296.1 <sup>b</sup> )	+0.831
2	τ CF <sub>2</sub> O	294.1	+0.720
3	CF <sub>3</sub> H	295.1 (293.5 <sup>b</sup> )	+0.656
4	CF <sub>3</sub> COCH <sub>3</sub>	292.8	+0.637
5	~ 3 3 CO <sub>2</sub>	292.5	+0.576
6	<sup>2</sup> ≈ <sup>3</sup> <sup>2</sup> <sup>2</sup> <sup>2</sup>	292.3	+0.569
7	~ 3 2 2 NaHCO <sub>3</sub>	290.2	+0.409
8	ئ Na <sub>2</sub> CO <sub>3</sub>	289.7	+0.226
9	$2^{-3}$ CH <sub>3</sub> $\sim^{2}$ <sub>2</sub> Na	289.1	+0.437
6	$3^{\sim 2}$ CF <sub>3</sub> $\sim^{\sim}$ <sub>2</sub> Na	289.1	+0.401
10	$3^{\sim} 2$ CH <sub>3</sub> CO <sub>2</sub> Na	288.8	+0.320
4	$3\sim 2$ CF <sub>3</sub> $\sim$ CH <sub>3</sub>	288.7	+0.363
11	3∼ 3 NH <sub>L</sub> HCO2	288.6	+0.318
12	CH <sub>3</sub> CO CH <sub>3</sub>	288.0	+0.302
13	5 5	287.9	+0.329
10	CH <sub>2</sub> O	285.0	+0.072
	$^{\text{CH}}_{\sim 3}^{\text{CO}}_{\sim 2}^{\text{Na}}$	285.0	
14	CH <sub>4</sub>		+0.098
15	KCN	285.0	-0.482
12	CH <sub>3</sub> CO CH <sub>3</sub>	284.9	+0.098
9	<sup>CH</sup> 3 <sup>CO</sup> <sup>H</sup> 2	284.8	+0.125

## Table III (continued)

<sup>a</sup> Binding energies taken from reference 3 except where noted.

b D. Thomas, unpublished work.

# Table IV

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# Chromium 3p Binding Energies

· · · · · · · · · · · · · · · · · · ·	Compound	Binding Energy, eV.	
	· · · · · · · · · · · · · · · · · · ·		
$K_2 Cr_2 O_7$		48.7	
Cr0 <sub>3</sub>		48.2	
Na2Cr04.	+H <sub>2</sub> 0	47.9	•
$Cr_2(SO_4)$	3 xH_0	46.9	
Cr(CO) <sub>6</sub>		45.8	
[Cr(H <sub>2</sub> 0)	5](NO3)3.3H <sup>2</sup> O	45.4	
CrCl <sub>3</sub> ·6H	20	45.4	
[Cr(H <sub>2</sub> 0)	c1]c12	45.2	
K <sub>3</sub> [Cr(CN	) <sub>6</sub> ]	44.8	
K <sub>3</sub> [Cr(CN	) <sub>5</sub> NO]	44.8	
Cr(NH <sub>3</sub> )6	, ,	44.7	
Cr203 xH	20	44.6	
-	-(H <sub>2</sub> 0) <sub>4</sub> Cl <sub>2</sub> ]Cl	44.5	•
Cr2(CH3C	о <sub>2</sub> ) <sub>4</sub> (н <sub>2</sub> о)	44.5	
Cr foil(1	unreduced)	43.9	
Cr203		43.5	• 4
CrN	· · ·	43.2	
Cr foil(	reduced)	43.2	•
		•	•

#### Table V

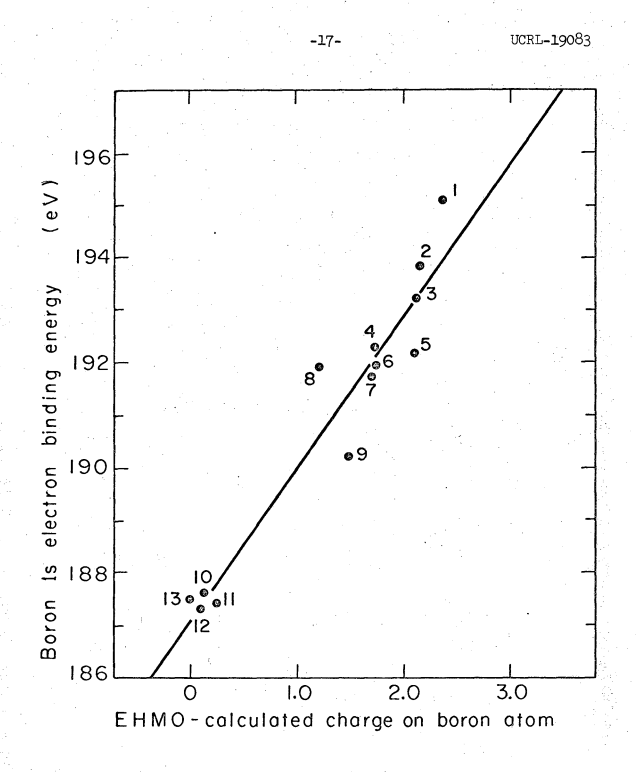
-16-

### Iterative Extended Hückel-Calculated

### Chromium Atom Charges

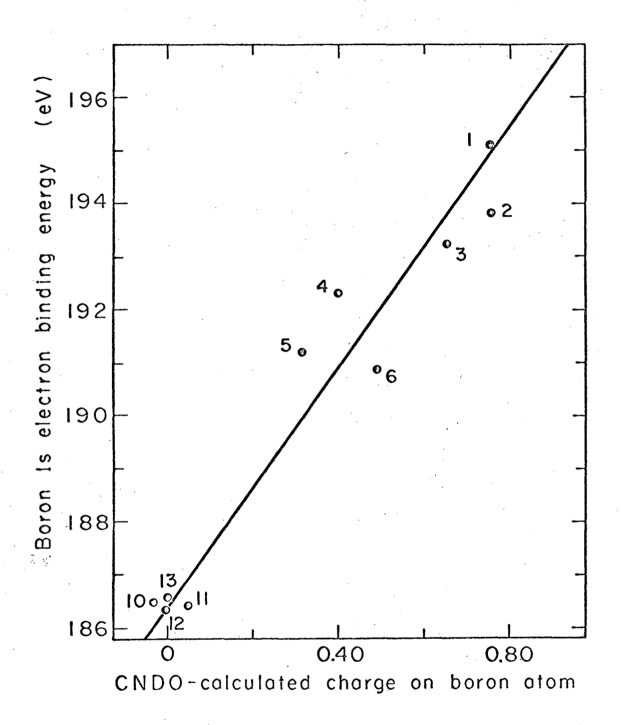
	Calculated Chromium Atom Charges		
Molecule	$d^{5}p$ Configuration <sup>a</sup>	modified $\frac{d^5p}{D}$ b Configuration	
Cr207 <sup>2-</sup>	<b>2.</b> 526	2.741	
Cr042-	2.430	2.618	
Cr(CN)6 <sup>3-</sup>	2.055	1.598	
<b>c</b> r(co) <sub>6</sub>	1.500	1.400	

a Slater exponents used,<sup>9</sup> H<sub>ii</sub> for metal: 3d(-8.40 eV), 4s(-8.01 eV), and 4p(-3.52 eV) as per reference 12 for neutral-atom d<sup>5</sup>p configuration.
b Metal orbital exponents: 3d(2.22), 4s and 4p(0.733). H<sub>ii</sub> for metal: 3d(-8.40 eV), 4s(-9.27 eV), and 4p(-3.52 eV).



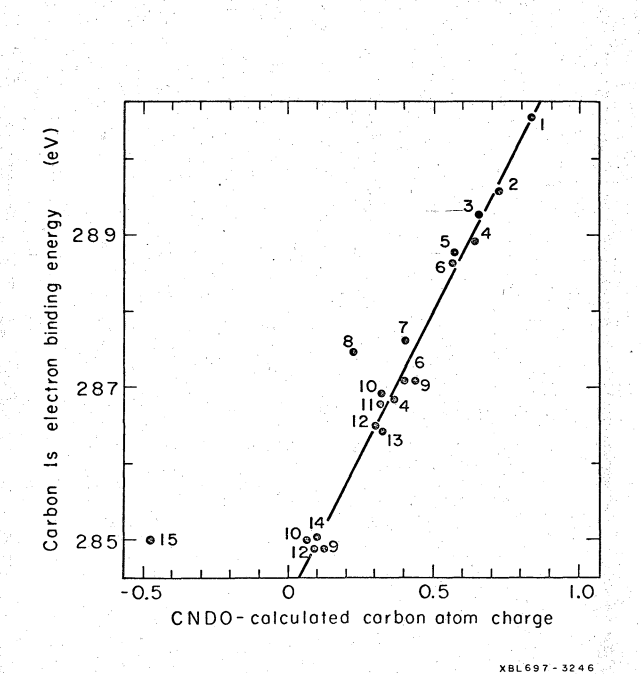
XBL 697-3245

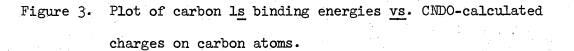
Figure 1. Plot of boron ls binding Energy vs. Extended Hückel-calculated charges on boron atoms.



XBL 697-3247 Figure 2. Plot of boron ls binding energy vs. CNDO-calculated charges on boron atoms.

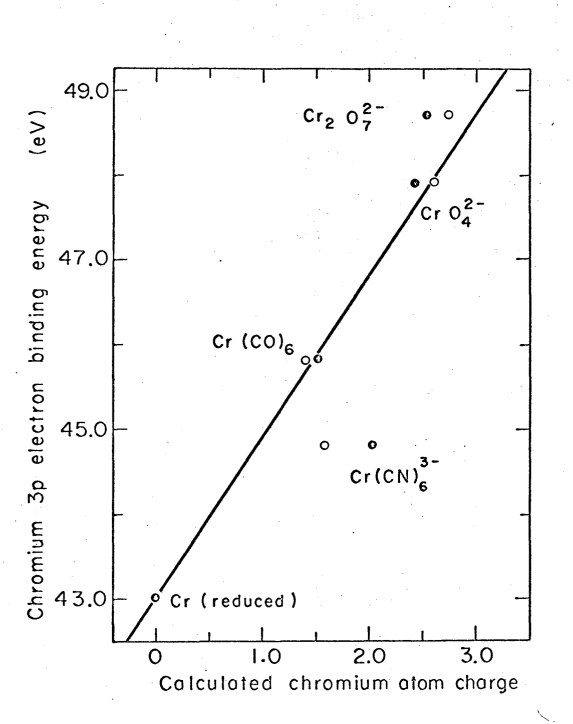
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-20-

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Figure 4. Plot of chromium 3p binding energies vs. iterative Extended
Hückel-calculated charges on chromium atoms. The symbols
and o refer to two different sets of input data as given
in footnotes a and b, respectively, in Table V.

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