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Theoretical Studies of the Electronic Structure of Small Metal Clusters

Final Report covering the Period October 1, 1979 - March 31, 1982

NASA Grant NSG - 2407*

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* The NASA technical officer for this grant is David M. Cooper, Materials Science Division, NASA Ames Research Center, Moffett Field, CA 94035. The main thrust of our research efforts during the period of the NASA grant involved theoretical studies of the electronic structure of metal clusters, in particular clusters of Group IIA and IIB atoms. Early in the project it became clear that electron correlation involving d orbitals plays a more important role in the binding of these clusters than had been previously anticipated. This necessitated that we replace our computer codes for calculating two electron integrals and for constructing the resulting CI Hamiltonions with newer, more efficient procedures. Hence, considerable time was spent on program modification, interfacing, and testing. (See Appendix A)

I. Group II A metal atom clusters

Perhaps the most significant finding resulting from our studies is that although Mg_4 is essentially unbound at an SCF level of treatment² the inclusion of electron correlation effects involving d functions gives rise to appreciable ($\sim 0.8 \text{ eV}$) binding.¹ This is the first example of strong binding originating from correlation effects alone. This work was performed in collaboration with Professor C. Dykstra and utilized the CEPA and SCEPA methods for treating electron correlation. These calculations on Mg_4 included explicitly all electrons and represents one of the largest (in terms of the number of electrons treated explicitly and number of basis functions employed) calculations on a metal cluster to date.

During the course of our study of Mg_4 , Pople and coworkers³ published a paper on the electronic structure of beryllium clusters. These authors also found that correlation effects involving d functions plays an important role in the cluster binding. However, a fundamental difference exists between the Be₄ and Mg₄ cases in that Be₄ displays appreciable binding even in the absence of correlation effects. The discoveries that

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d functions play a fundamental role in the binding of beryllium and magnesium clusters has led us to perform new calculations on $B\bar{e}_2$ and $B\bar{e}_3$. Here we have found that it is essential to include both d functions and correlation effects to obtain the correct ordering of the pi and sigma anion states.⁴

We are presently carrying out a new set of calculations on Mg_4 utilizing a local potential to model the $1s^2 2s^2 2p^6$ atomic cores. If these local potential calculations successfully reproduce the results of the all-electron calculations, then, we can confidently employ local potentials to study the larger clusters.

As part of our continuing effort to understand electron attachment to group IIA metal clusters, we have also calculated⁵ electron affinities of atomic Be, Mg, and Ca. For the atoms the anion states lie energetically above the ground states of the neutral atoms, requiring scattering calculations for their investigation. Using the Harris-Michels method, we have shown that all three atoms have low-lying ²P anions. For Mg a broad feature due to ²D Mg⁻ is found near 2.5 e.V in the scattering cross section. For Ca the ²D anion is found to lie only 0.7 ev above the ²P anion.

II. Cu, and CuH.

During the course of the grant we began calculations on Cu_2 and CuHand their anions⁶. In these studies we have utilized the effective potential of Topiol <u>et.al</u>.⁷ for the copper atoms. The first challenge was to find a compact Cu basis set suitable for the d¹⁰s and d⁹s² states of Cu as well as the d¹⁰s² state of Cu⁻. After much experimentation, we arrived at a [4s3p5d/4s3p2d] basis set. With this basis we are able to

-2-

obtain a SCF $d^{10}s \rightarrow d^9s^2$ excitation energy of 0.59 eV in fair agreement with the all-electron value of 0.34 eV and a SDCI (single and double excitation configuration interaction) excitation energy of 1.41 eV in excellent agreement with the experimental value of 1.49 eV. A SDCI with this basis yields a dissociation energy of Cu₂ of 1.87 eV compared with the 2.05 eV experimental value. The above results are noteworthy since other investigators have concluded that a 5d/3d contraction is required to describe the $d^9s^2 \rightarrow d^{10}s$ atomic excitation energy and presumably also to describe the bonding in the Cu₂ molecule. The calculations on the Cu₂ and the Cu₂ potential energy curves as well as on CuH and CuH are near completion. We expect to submit this work in the near future to the Journal of Chemical Physics for publication.

III. <u>Reactions of Metal Atoms with H₂O</u>.

The second major thrust of our research program has been the investigation of the reactions of metal atoms with small molecules, especially H_20 . In particular, we have characterized the M·OH₂ and HMOH species for M= Be, B, C, Mg, Al, Si.^{9,10} For each of these species the geometries, stabilities, and vibraticnal frequencies have been determined. Perhaps the most interesting finding of these calculations is that for the B, C, Al, and Si insertion products both cis and trans isomers exist. Experimental evidence for the existence of cis/trans isomers of HSiOH has recently been found.¹¹

IV. Other Work Carried Out Under the NASA Grant

Theoretical investigations of several other systems were also completed during the course of our NASA grant. Using 2^{nd} order many-body perturbation theory we have determined the polarizabilities of LiF and Be0,¹²BF,¹³ and LiC1.¹⁴ The results for LiF and Be0 ¹² and those for

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BF have been published.¹³ Those for LiCl are near completion.¹⁴ The calculations on LiCl are of particular interest since in this case we have determined several properties using different pseudopotentials for the chlorine atom. Finally, we have utilized¹⁵ both the configuration interaction and many-body perturbation theory (through fourth-order) to calculate the electron affinities of C_2 and C_3 . We plan to extend these studies to include larger carbon clusters.

Appendix A: Program Implementation and Development

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During the course of the NASA grant we implemented the Pitzer-Schaefer integral program, which is considerably faster for integrals over d functions than the MOLECULE program we had been using. A significant gain in speed resulted from a switch over to Shepard's Unitary-group multiconfigurational self-consistent field (MCSCF) and configuration interaction (CI) programs. These programs were modified to allow for the use of local potentials to model the core electrons and to use input vectors from Purvis' SCF program. 3

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