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### ABSTRACT OF DISSERTATION

VAN DER WAALS CLUSTERS OF AROMATIC MOLECULES STUDIED USING

SUPERSONIC MOLECULAR JET SPECTROSCOPY

SAGE STRUCTURE MANAGER

van der Waals (vdW) clusters of aromatic solutes with various solvents are studied in the gas phase using supersonic molecular jet spectroscopy. Calculations involving ground state cluster binding energy, geometry, and intermolecular vibrational structure are also presented to complement the experiments. The analyses include: spectroscopic studies and theoretical modeling of the intermolecular vibronic structure of benzene solvated by argon, methane, water, and ammonia; spectroscopic studies and theoretical modeling of the intermolecular vibronic torsional structure in benzene solvated by methane, deuteromethane, and carbon tetrafluoride; solvation of pyrazine and pyrimidine by both small hydrocarbon and hydrogen bonding solvents; the study of pyrazine and pyrimidine dimers; and the solvation of macrocycles such as free base phthalocyanine  $(H_2^Pc)$  and magnesium phthalocyanine (MgPc) by small hydrocarbon solvents, hydrogen bonding solvents, and carbon dioxide.

The benzene/solvent studies lead to the elucidation of the detailed nature of the intermolecular vibrational structures and the geometries/symmetries present in the clusters. These studies reveal that the majority of the intermolecular vibronic transitions observed involve vdW bending and torsional motion parallel to the solute pi cloud. Furthermore, the clusters behave rigidly with regard to internal rotation of the cluster subunits and the clusters possess unique equilibrium geometries.

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The pyrazine and pyrimidine/solvent and dimer studies demonstrate the detailed effects of the solute ring nitrogens on cluster geometry and on the role of hydrogen bonding in the clusters.

The  $H_2Pc$  and MgPc/solvent cluster experiments and models suggest that stable solute solvation sites are located over the phthalocyano core and not over peripheral ring centers. Forbidden low frequency cluster chromophore out-of-plane vibronic transitions are also induced by solvation in both the  $H_2Pc$  and MgPc clusters.

Primary References:

1. J. A. Menapace and E. R. Bernstein, Journal of Physical Chemistry, 1987, <u>91</u>, 2843.

2. J. A. Menapace and E. R. Bernstein, Journal of Physical Chemistry, 1987, <u>91</u>, 2533.

3. J. A. Menapace and E. R. Bernstein, Journal of Chemical Physics, 1987, <u>86</u>, 0000.

4. J. Wanna, J. A. Menapace, and E. R. Bernstein, Journal of Chemical Physics, 1986, <u>85</u>, 777.

5. J. Wanna, J. A. Menapace, and E. R. Bernstein, Journal of Chemical Physics, 1986, <u>85</u>, 1795.

Secondary References:

1. M. Schauer and E. R. Bernstein, Journal of Chemical Physics, 1985, <u>82</u>, 726.

2. K. S. Law, M. Schauer, and E. R. Bernstein, Journal of Chemical Physics, 1984, <u>81</u>, 4871.

3. J. Wanna and E. R. Bernstein, Journal of Chemical Physics, 1986, 84, 927.

Joseph Arthur Menapace, Capt, USAF Doctor of Philosophy Department of Chemistry Colorado State University Fort Collins, CO 80523 Summer 1987 Dissertation: 339 Pages

DISSERTATION

VAN DER WAALS CLUSTERS OF AROMATIC MOLECULES STUDIED USING SUPERSONIC MOLECULAR JET SPECTROSCOPY

> Submitted by Joseph Arthur Menapace Department of Chemistry

In partial fulfillment of the requirements for the Degree of Doctor of Philosophy Colorado State University Fort Collins, Colorado

Summer 1987

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Committee on Graduate Work

Advisor

Department Head

ABSTRACT OF DISSERTATION

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van der Waals (vdW) clusters of aromatic solutes with various solvents are studied in the gas phase using supersonic molecular jet spectroscopy. Calculations involving ground state cluster binding energy, geometry, and intermolecular vibrational structure are also presented to complement the experiments. The analyses include: spectroscopic studies and theoretical modeling of the intermolecular vibronic structure of benzene solvated by argon, methane, water, and ammonia; spectroscopic studies and theoretical modeling of the intermolecular vibronic torsional structure in benzene solvated by methane, deuteromethane, and carbon tetrafluoride; solvation of pyrazine and pyrimidine by both small hydrocarbon and hydrogen bonding solvents; the study of pyrazine and pyrimidine dimers; and the solvation of macrocycles such as free base phthalocyanine  $(H_2Pc)$ and magnesium phthalocyanine (MgPc) by small hydrocarbon solvents. hydrogen bonding solvents, and carbon dioxide.

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The pyrazine and pyrimidine/solvent and dimer studies demonstrate the detailed effects of the solute ring nitrogens on cluster geometry and on the role of hydrogen bonding in the clusters.

The  $H_2Pc$  and MgPc/solvent cluster experiments and models suggest that stable solute solvation sites are located over the phthalocyano core and not over peripheral ring centers. The  $H_2Pc/hydrocarbon$  cluster results parallel those obtained for the benzene and N-heterocycle/ hydrocarbon clusters. The  $H_2Pc$  and MgPc/alcohol cluster spectra and calculated geometries suggest that the solvent OH groups are intimately involved in the intermolecular interactions. Forbidden low frequency cluster chromophore out-of-plane vibronic transitions are also induced by clustering. This low frequency motion is characterized using an outof-plane normal coordinate analysis on the  $H_2Pc$  moiety.

> Joseph Arthur Menapace Department of Chemistry Colorado State University Fort Collins, CO 80523 Summer 1987

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То

Ida M. Menapace,

Janet J. Juranek, and

John E. Juranek Sr.,

With Love... Joe

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# CHAPTER ONE

# INTRODUCTORY COMMENTS AND OVERVIEW

Introduction.

The material contained in this dissertation is largely presented in five publications. In the spirit of completeness and accuracy, these papers are incorporated directly into the dissertation. The first three papers appear as chapters in the body of the manuscript. The remaining two papers appear in reprint form as appendices. The rest of the dissertation entails the introductory comments and overview, unpublished results, future/proposed experiments, conclusions, and the computer programs utilized in this work.

The introductory comments and overview are meant to tie together the body of the dissertation since directly presenting the publications tends to fragment the discussion. They will also provide the reader with the general notions underlying the work presented in detail in each major section or chapter of the dissertation.

The unpublished results contain mostly those data which are not important enough to appear in papers, or data which need additional work to interpret. The data are presented in the dissertation to establish a record of the research conducted in specific areas and to provide future investigators with a basis for further study.

The future/proposed experiments involve both theoretical and experimental studies of solute/solvent cluster systems similar to those

presented in this dissertation. The primary motivation in these proposed experiments is to provide for continued research on cluster systems using the models, procedures, and experimental set-ups currently available in the Bernstein group laboratory.

The presentation of the computer programs used in the cluster studies is largely for the convenience of future investigators. The programs serve as easy references for those conducting research in the same or similar areas as well as provide documentation on the "how to's" of the models used in the studies. 

### Overview.

The five publications incorporated into this dissertation are divided into five chapters. The work covers: 1) the experimental study and theoretical modeling of the intermolecular vibronic structure of benzene solvated by argon, methane, water, and ammonia; 2) the experimental study and theoretical modeling of the intermolecular vibronic torsional structure in benzene solvated by methane, deuteromethane, and carbon tetrafluoride; 3) the detailed study of the solvation of pyrazine and pyrimidine by both small hydrocarbon and hydrogen bonding solvents: 4) the study of pyrazine and pyrimidine dimers; and 5) the solvation of macrocycles such as free base phthalocyanine and magnesium phthalocyanine by small hydrocarbon solvents, hydrogen bonding solvents, and carbon dioxide. This overview is intended to the together the major results of these studies and to create a sense of unity and purpose for the studies.

The Vibronic Structure of Solute/Solvent van der Waals Clusters A large body of data has been accumulated in this laboratory for small aromatic molecules such as benzene, pyrazine, and pyrimidine solvated

with small hydrocarbon solvents such as methane, ethane, and propane, and hydrogen bonding solvents such as water and ammonia. These spectroscopic studies yield observables such as spectral shift which can be related to the difference between the cluster binding energy between the ground and electronic excited states and to cluster geometry. In the majority of the cluster spectra, intermolecular vibronic transitions are observed which can be used to elucidate the cluster symmetry/geometry, electronic state mixing, and the intricacies of the intermolecular potential surface.

Modeling of the intermolecular interaction between the cluster constituents also complements the spectroscopic studies. These calculations allow one to elucidate some physical properties of the clusters such as ground state cluster binding energy and geometry. The calculations always yield results consistent with experimental observations in regard to the number of cluster configurations observed, their respective binding energies, and their qualitative geometries. Modeling of the intermolecular vibronic structure in these molecules adds to the understanding of the cluster systems as it allows one to obtain detailed information on cluster geometry/symmetry and to elucidate the intricacies of the potential established between the cluster constituents.

Presently, little information is available, either experimentally or theoretically, on the details of the intermolecular vibrational structure in the solute/solvent cluster systems. Furthermore, the van der Waals (vdW) modes are interesting as they play a key role in dynamical energy transfer processes in the clusters as well as represent precursors to a variety of motions occurring in liquids and solids.

The modeling of the intermolecular vibrational structure in this study is approached by extending the empirical models previously used to

calculate cluster ground state geometry and binding energy to include modeling of the ground state vdW motion. The vdW vibrational structure is modeled using a normal coordinate analysis in which the clusters are treated as "giant molecules" whose motion is governed by a force field describing both the intra- and intermolecular motion. The intermolecular force field is based upon an atom-atom Lennard-Jones potential function including general non-bonding (6-12), hydrogen bonding (10-12). and monopole charge (1) terms.

Using this model, the eigenvalues and eigenvector normal modes for the vdW motion occurring in benzene(Ar)<sub>1</sub>, /(CH<sub>4</sub>)<sub>1</sub>, /(H<sub>2</sub>O)<sub>1</sub>, and /(NH<sub>3</sub>)<sub>1</sub>, and s-tetrazine(Ar)<sub>1</sub> are calculated. The results of these calculations are then used to assign the vdW motions observed in vibronic spectra of the aforementioned cluster systems which are obtained using 1- and 2-color time-of-flight mass spectroscopy (TOFMS). Agreement between the calculations and experiments is excellent for cluster binding energies, symmetries, and vdW frequencies. Essentially, the S<sub>1</sub> + S<sub>0</sub> vibronic transition of the clusters are completely assigned based upon these calculations.

A number of approximate "diatomic molecule" models are also considered in this study to analyze the vdW structure in benzene(Ar)<sub>1</sub>. The cluster vdW modes are modeled using three methods: 1) a Taylor series expansion of the intermolecular potential along the vdW stretching and bending coordinates; 2) a Morse potential fit to the intermolecular potential along the vdW stretching and bending coordinates; and 3) a semi-classical energy level fit to the intermolecular potential using the JWKB method. The models essentially treat the clusters as simple two particle systems whose motions are restricted to

the principle axes of a Cartesian coordinate system and can, thereby, be analyzed using a diatomic molecule type approximation. These models are considered to show the consistency of the calculations between each diatomic molecule model and the intermolecular normal coordinate analysis.

The experimental and theoretical studies on the benzene/solvent systems give rise to several interesting notions which involve the detailed nature of the intermolecular interaction. First, the studies reveal that the weak vdW potential between the cluster solute and solvent is, for the most part, the same for the ground and excited electronic states. Second, the majority of the observed vdW vibronic transitions observed in the cluster spectra are those involving vdW bending and torsional motions parallel to the aromatic  $\pi$  system. Third, these modes are quite active in the Herzberg-Teller vibronic coupling mechanism. And fourth, vdW motions for which the cluster solvent penetrates the aromatic  $\pi$  system of the solute have high frequencies and are typically not observed.

The Intermolecular Vibronic Torsional Structure in Solute/Solvent vdW Clusters: Benzene/Methane, /Deuteromethane, and /Carbon Tetrafluoride - The modeling of the intermolecular motion in the vdW clusters discussed above is basically approached using, more or less, a rigid molecule bound energy well approximation. This approach seems quite reasonable for the vdW stretching and bending degrees of freedom which are essentially translations of the cluster constituents relative to one another. The vdW torsional modes, on the other hand, could in principle not be oscillatory. They could be free or hindered rotation of the cluster solvent relative to the cluster solute. In this regard the

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intermolecular normal coordinate analysis may not faithfully reproduce the vdW torsional motion present in the clusters.

To study this possibility, we chose to analyze benzene clustered with methane, deuteromethane, and carbon tetrafluoride. The motivation for studying this "isotopic" cluster solvent series centers upon the elucidation of the vdW torsional structure. Two limiting cases can be proposed in regard to the torsional structure in benzene( $CH_4$ )<sub>1</sub>, benzene $(CD_4)_1$ , and benzene $(CF_4)_1$ . In one case, the clusters can possess free internal rotation in which the cluster solvent freely rotates in three dimensions against the benzene frame and the system can be considered internally non-rigid. In the other case, the clusters possess torsional oscillations for which the cluster solvent librates against the benzene frame with a residence time long enough to give rise to "vibration like" motion in an internally rigid molecule regime. Understanding the vdW torsional motion is particularly interesting in these systems as the limiting cases pose questions regarding the physics governing the vdW torsional structure: 1) do the clusters posses free/ hindered internal rotation or do they possess torsional oscillations; and 2) in either case, what is the dependence of the intermolecular potential upon the relative orientation of the cluster constituents?

In the studies, the clusters are probed spectroscopically using supersonic molecular jet expansion and 2-color TOFMS techniques. The cluster  $S_1 + S_0$  intermolecular vibronic structures are then characterized by calculational modeling of the vdW motion. The calculations include: 1) an intermolecular normal coordinate analysis which treats all six vdW modes under a harmonic oscillator assumption; and 2) a three-dimensional hindered rigid rotor analysis which treats only the intermolecular torsional motion.

In the experimental and theoretical studies on the benzene $(CH_4)_1$ , henzene $(CD_4)_1$ , and benzene $(CF_4)_1$  systems, several interesting results are obtained. First, the cluster vibronic spectra show that the clusters are at least semi-rigid systems with regard to internal rotation of the cluster subunits and that the clusters possess unique equilibrium geometries. Second, the spectra demonstrate that the intermolecular motion present in the systems is oscillatory and, through the "isotopic" shifts observed, that the low-Jying eigenstates are nearly harmonic. They are not admixtures of vdW bends, stretches, and free internal rotation as would occur if the clusters were internally nonrigid. Third, both the intermolecular normal coordinate analysis and the three-dimensional hindered rigid rotor analysis indicate that the vdW torsional structure is oscillatory and that the motion is constrained by an orientationally dependent intermolecular potential whose barrier height is on the order of the cluster binding energy.

Hydrogen Bonded and Non-Hydrogen Bonded vdW Clusters: Comparison between Clusters of Pyrazine, Pyrimidine, and Benzene with Various Solvents - The study of the solvation of pyrazine, pyrimidine, and benzene by both small hydrocarbon and hydrogen bonding solvents has led to detailed information about the structure and energetics present in the solute/solvent systems. Our chief motivations for studying these clusters center upon the elucidation of the effects of the ring nitrogen atoms in pyrazine and pyrimidine solutes on cluster geometry and on the role of hydrogen bonding in the pyrazine, pyrimidine, and benzene cluster systems. These studies are of interest as hydrogen bonding interactions are known to play an important role in the intra- and intermolecular interactions responsible for secondary and tertiary

structure, molecular dynamics, and ionic and molecular solvation. Assignment of the spectra obtained in these studies is accomplished through the determination of the cluster spectral shifts, ionization energies, relative intensities, molecular forbidden cluster transitions, and intermolecular potential/cluster geometry calculations. Computer modeling of the cluster systems has proven essential to the understanding of the spectroscopic data. Through the interplay of the computer modeling of these clusters and the spectroscopic data, it is possible to assign probable geometries to many of the spectroscopic features. In many cases this allows one to find the spectral shift corresponding to a specific geometry of a cluster. The assignment of a geometry to the spectral shifts leads to some general conclusions about the types of interactions responsible for the spectral shifts.

In these studies we find that the clusters fall into two general categories: 1) a conventional set containing the aromatic solute/ hydrocarbon solvent clusters; and 2) henzene, pyrimidine, pyrazine ammonia and benzene water clusters. The aromatic solute/hydrocarbon cluster spectra are quite similar to one another in that they all possess bathychromic shifts with respect to the isolated chromophore transition. The cluster binding energies and calculated geometries are also similar. The presence of the nitrogen atoms in the aromatic ring of the pyrazine and pyrimidine solutes has a relatively small although discernible effect on the overall intermolecular interaction. In particular, the solvent hydrogen atoms preferentially orient towards the solute nitrogen atoms.

The solute/hydrogen bonding solvent cluster systems, on the other hand, possess vibronic spectra which are all unique and surprisingly

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erratic. In these clusters the spectral shifts range from -100 to  $500 \text{ cm}^{-1}$ , the vdW vibronic motion in the systems range from nonexistent to intense, the vdW modes are in some cases highly perturbing to the solute vibronic structure and energy, and the number of unique cluster geometries range from one to three in an apparently random fashion. In spite of these differences the models used to calculate the cluster geometries and binding energies corroborate the experiments as far as the comparison can be made in regard to cluster symmetry, number of configurations observed, and red and blue shifts in regard to hydrogen bonding. The hypsochromic shifts observed in the pyrazine and pyrimidine ammonia clusters suggest that hydrogen bonding between the ring nitrogen and the solvent hydrogens may be contributing to some extent to the total intermolecular interaction responsible for cluster formation/ stabilization. The benzene/ammonia and benzene/water spectra seem to suggest that some type of hydrogen bonding may be occurring in the benzene/water system between the aromatic  $\pi$  cloud and the solvent mojety as the benzene $(H_20)_1$  spectral shift is hypsochromic whereas the  $benzene(NH_3)_1$  spectral shift is bathychromic.

<u>Supersonic Molecular Jet Studies of the Pyrazine and Pyrimidine</u> <u>Dimers</u> - The understanding of the solute/solute interactions in the pyrazine and pyrimidine dimer systems is of interest for a number of reasons. First, the dimers serve as model systems for condensed phase structure, dynamics, and nucleation and growth of molecular aggregates. Second, the dimers provide insight into the understanding of the second ary and tertiary structures present in more complicated molecules. And third, studying these clusters in the gas phase yields information on the major interactions responsible for dimer formation and what types of

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dimer geometries are most probable in an isolated environment free from the extraneous perturbations present in liquid and solid phases.

فتنتعمهم

The pyrazine and pyrimidine dimers are analyzed using supersonic molecular jet expansion and 2-color TOFMS techniques for which the mass selected optical spectra of the first excited singlet  $n\pi^*$  transitions of the dimers are observed. On the basis of what we have learned from the study of the simple solute/solvent clusters, we are able to analyze the dimer systems using cluster ionization energy, vibronic structure, spectral shift, and modeling of the intermolecular potential between the cluster subunits. The interplay of the spectroscopic data and the calculations allows us to obtain a consistent set of geometries for the dimers present in the supersonic expansion.

In the pyrazine dimer system, the experiments and calculations suggest that the both parallel hydrogen bonded and perpendicular dimers are present in the supersonic expansion. The calculations also predict a parallel stacked/90° rotated dimer which is not observed. This species most likely forms an excimer in the excited state with a short lifetime and a large red shifted and broad spectrum. The major distinction between the two observed dimer species is determined to be the difference in the involvement of the  $\pi$  clouds in the overall dimer interaction as evidenced by differences in the cluster ionization energy between the two cluster species.

In the pyrimidine dimer, the calculations yield four planar hydrogen bonded species and a parallel stacked/displaced species. The observed dimer vibronic spectra are consistent with these calculated geometries. As in the pyrazine dimer, the ionization energy proves to be an important contribution in determining the number of different dimer geometries responsible for the observed spectra. Additionally, we find that the dimer spectral shifts depend upon the specific cluster geometry. A bathychromic shift is associated with the parallel stacked/ displaced dimer and hypsochromic shifts are associated with the planar hydrogen bonded dimers.

To explore further the agreement between the experiments and calculations on the dimer systems, we also conducted calculations on the tetrazine dimer. Three calculated geometries are obtained for the tetrazine dimer: a parallel stacked/90<sup>0</sup> rotated species, a planar hydrogen bonded species, and a perpendicular species. In this study we find that the calculated geometries are in agreement with geometries determined from rotational analysis.

Computer modeling is determined to be an essential component to the study of these dimer systems. The spectroscopic data are needed to validate the computer modeling techniques, and the calculations help to assign spectra which are otherwise quite puzzling. In this way the interplay between spectroscopy and computer modeling leads to a better understanding of the structure and energetics of the solute/solute systems.

Supersonic Molecular Jet Studies of Phthalocyanines and Their vdW Clusters with Small Molecules - In these studies, free base phthalocyanine ( $H_2Pc$ ) and magnesium phthalocyanine (MgPc) clustered with solvents such as small hydrocarbons ( $C_nH_{2n+2}$  (n=1, 2, 3)), hydrogen bonding solvents ( $H_2O$ , MeOH, EtOH), and  $CO_2$  are analyzed in an isolated ultracold molecular environment. The clusters are generated using a high temperature continuous supersonic molecular jet especially designed for efficient generation and study of the species in the gas phase. The

advantages of the supersonic molecular jet are exploited in these studies as the complicated phthalocyanine spectra are dramatically simplified and the solvent or environmental perturbations present are controlled in a set and reproducible manner.

Our basic motivation for studying these systems centers upon elucidation of the solvation properties of the macrocycles in a controlled and well-defined environment. In this regard, supersonic molecular jet investigations on the H<sub>2</sub>Pc and MgPc solute/solvent clusters can contribute to the resolution of a number of important concerns dealing with the behavior of the systems on the microscopic scale. Questions that we considered in these cluster studies are 1) what are the ground and excited state binding energies between the phthalocyanines and various solvents, 2) what are the preferential interaction sites on the phthalocyanine moiety, 3) what are the most favorable cluster geometries, 4) what types of interactions are important in the intermolecular interaction established between the solute and the solvent, and 5) are changes to the chromophore symmetry/geometry induced by clustering.

The  $H_2Pc$  and MgPc solute/solvent clusters are characterized by analysis of their gas phase fluorescence excitation spectra and modeling of the intermolecular potential between the cluster solutes and solvents. In these studies, we find that forbidden cluster chromophore out-of-plane motion is induced by clustering and that elucidation of the nature of this out-of-plane motion is essential to the understanding of the cluster spectra and in the identification of the number of different clusters of a specific composition observed. This prompted us to conduct an out-of-plane normal coordinate analysis on  $H_2Pc$  to

characterize the motion. The comparison between these calculations and the experiments makes possible the identification of specific species/ geometries responsible for the cluster vibronic transitions observed in the spectra.

Several interesting results are obtained from the cluster First, the cluster vibronic spectra and calculations suggest studies. that stable H<sub>2</sub>Pc and MgPc solvation sites are located over the phthalocyano core. Local minima over peripheral ring centers are either nonexistent or too shallow to accommodate minimum energy bound state geometries. Second, the H2Pc/hydrocarbon cluster experimental and theoretical results parallel those obtained for benzene and the Nheterocycle/hydrocarbon clusters. The spectral shifts observed in these solvent series are all bathychromic and the magnitudes of the spectral shifts increase with increasing solvent size and polarizability. Third, the H\_PC and MgPc/alcohol cluster spectra and calculated geometries suggest that the solvent OH group is intimately involved in the intermolecular interactions and contributes significantly to the observed spectral shifts. Fourth, MgPc clusters display weak vdW interactions between the cluster solute and solvent. Actual complexation in which the solvent donates electron density to the solute does not occur. Fifth, excited electronic state splitting may occur in the MgPc clusters due to the reduction in system symmetry upon cluster formation. The degenerate Q band in the isolated MgPc spectrum appears to split into its two components,  $Q_{\chi}$  and  $Q_{\gamma}$ , in the cluster spectra. Finally, forbidden low frequency cluster chromophore out-of-plane motion is induced by clustering in both the H<sub>2</sub>Pc and MgPc systems. Intensity of this motion arises from the reduction of the chromophore symmetry in the clusters.

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# CHAPTER TWO

THE VIBRONIC STRUCTURE OF SOLUTE/SOLVENT VAN DER WAALS CLUSTERS

Introduction.

The combination of laser spectroscopy and supersonic molecular jet expansions has made possible the study of a wide array of weakly bound van der Waals (vdW) molecules in the gas phase. These clusters, formed in the jet expansion, are stable in the post-expansion region and can be studied as isolated molecules. They are interesting both theoretically and experimentally because of their unique characteristics such as low binding energies, large intermolecular equilibrium distances, and low frequency intermolecular vibrational modes. Furthermore, the vdW clusters only slightly perturb the individual properties of their molecular constituents.<sup>1</sup> These characteristics set the vdW cluster apart as a distinct phase of matter to be explored and understood.

The electronic-vibrational spectroscopy of aromatic molecules like benzene<sup>2-4</sup> and s-tetrazine<sup>5</sup> clustered with various solvents reveals interesting information regarding unique cluster characteristics. Specifically, the studies show detailed information pertaining to the intermolecular energetics and dynamics of cluster interactions, espec ially in the area of the clusters' low frequency vdW vibrational modes. These modes are of considerable interest since they represent the pre cursors of a variety of condensed phase eigenstates such as phonons in liquids and solids. They are also key factors in cluster dynamics as they play a major role in the energy transfer processes of intramolecular vibrational redistribution (IVR) and vibrational predissociation (VP).

Presently, little information is available concerning the detailed intermolecular vibrational structure in molecule-molecule clusters either experimentally or theoretically. The majority of the theoretical work on cluster energetics and dynamics to date is focused upon the intermolecular modes in simple atom-molecule systems. The energetic studies range from quantitative treatment of the vdW stretch and qualitative discussion of the vdW bends $^{1,6,7}$  in the atom-molecule clusters to full quantitative treatment of both the vdW bends and stretch<sup>16</sup> in systems for which the intermolecular potential is easily modeled. In dynamical studies,<sup>6,7</sup> primary emphasis is placed upon the vdW stretching mode since it is presupposed that this motion is the major contributor to IVR and VP processes. The lusters are thereby treated quantitatively using a "dumbbell" approximation in which the intermolecular motion is restricted to a stretching mode form. Within this approximation, neglecting the quantitative contributions of the bending/torsional vdW modes in the dynamical scheme seriously limits the application of theoretical treatments to IVR and VP phenomena occurring in molecule-molecule clusters. The stretching mode "restriction" dictates that only certain energy transfer processes can be modeled. specifically those which involve the vdW stretch. Experimental evidence of the theory's limitation<sup>4</sup> is found in the observation of energy transfer from prepared states tangential to the vdW stretching motion. Part of the difficulty of incorporating the vdW bending and/or torsional modes into the theory is that no model has been demonstrated which

describes adequately either their energetics or mode nature in moleculemolecule clusters.

In previous publications,<sup>8</sup> we have used theoretical calculations to elucidate structure and binding energy in the vdW cluster ground state. Combining these calculations with experimental observables such as binding energies, ionization energies, spectral shifts, relative feature intensities, and the appearance of cluster constituent, symmetry forbidden, transitions has aided considerably in spectral assignment and understanding. The calculations always yield results consistent with experimental observations in regard to the number of cluster configurations observed, their respective binding energies, and their qualitative geometries. **WALLER** 

In the majority of the cluster spectra studied, vdW vibronic features are observed and sometimes assigned based upon overtone and combination band analysis. In other cases, however, these vibronic features are numerous and complex, making elucidation of the mode fundamentals difficult and sometimes impossible. The vdW vibronic feature assignments have been made based upon the assumptions that the vdW stretch occurs at higher frequency and with greater intensity than vdW bends and torsions. Assigning spectra using these assumptions is a difficult task without <u>a priori</u> knowledge of vibronic mode nature since one must consider that, in nonlinear polyatomic molecule-molecule clusters, six vdW modes exist of which only one is a stretching mode. (In a non-linear atom-polyatomic molecule cluster, three vdW modes exist).

The studies reported in this publication involve the calculation of a complete set of ground state vdW vibrational modes for

benzene(Ar)<sub>1</sub>, s-tetrazine(Ar)<sub>1</sub>, benzene(CH<sub>4</sub>)<sub>1</sub>, benzene(H<sub>2</sub>O)<sub>1</sub>, and benzene(NH<sub>3</sub>)<sub>1</sub>. The calculations are performed using a selfconsistent pairwise atom-atom intermolecular potential developed by Scheraga, et al.<sup>9</sup> containing general nonbonding (6-12), general hydrogen bonding (10-12), and monopole charge parameters. The calculated ground state vdW modes are compared with cluster vibronic spectra previously studied in this<sup>2,3</sup> and other laboratories.<sup>5</sup> Consequently, a number of spectral reassignments are suggested. Vibronic selection rules governing the vdW cluster S<sub>1</sub> + S<sub>0</sub> transitions are also derived based upon calculated ground state cluster mode symmetry, cluster geometry, and experimental observation. Within this framework, ramifications of Herzberg-Teller vibronic coupling are discussed as they pertain to experimental observations of Franck-Condon forbidden transitions.

The calculated vdW vibrations are presented for all systems considered as eigenvector normal modes and eigenvalue energies determined via normal coordinate analysis of the entire vdW cluster.<sup>10</sup> In performing the calculations, the high frequency intramolecular vibrations of the cluster constituents are assumed to be completely uncoupled from the low frequency vdW modes.

Simpler models are also considered in studying the vdW transitions of benzene(Ar)<sub>1</sub>. The system is studied using four methods: 1) a Taylor series expansion of the intermolecular vdW potential along the three Cartesian axes in which the term coefficients are related to the vibrational frequencies in these directions; 2) a Morse potential fit to the intermolecular vdW potential along each of the three Cartesian axes; 3) a Morse potential fit to the intermolecular vdW potential using  $\beta = \frac{6}{R_o}$ ;<sup>1,6,7</sup> and 4) a semi-classical energy level fit to the

intermolecular vdW potential using the JWKB method. These studies are presented to show the consistency of the calculations and to reveal the advantages and pitfalls of the models. The four models involve treating the vdW clusters as simple two particle systems whose motions are restricted to the principle axes of the Cartesian coordinate system. The intermolecular vdW potential surfaces are thereby reduced to onedimensional potential functions which can be analyzed in a diatomic molecule approximation.

The above outlined approaches are all based more or less on a rigid molecule, bound potential energy well approximation. This approach would seem quite reasonable for the stretching(s) and bending  $(b_x \text{ and } b_y)$  degrees of freedom which are essentially translations of the components of the vdW molecules with respect to one another. Torsional modes  $(t_x, t_y, t_z)$ , on the other hand, could in principle be modeled by a free/hindered rotor formalism.<sup>11,12</sup> A one-dimensional "free rotor" description has been applied to the symmetry axis (z) torsional motion of benzene and toluene  $(CH_4)_1$ ,  $(CD_4)_1$  and  $(CF_4)_1$  and compared to the experimental observations. Preliminary results suggest that this approach does not faithfully reproduce the experimentally observed spectra for this "isotopic" series in terms of line shapes, intensities, major features, and the number of observed transitions. A threedimensional "free rotor" model has also been applied to this problem in order to treat all torsional modes  $(t_x, t_y, t_z)$  simultaneously. Similar difficulties are experienced in fitting the experimentally observed spectra. An account of these studies will be submitted for publication in the near future.

The driving motivation in these studies is to answer the following questions: 1) if parametric calculations involving cluster

geometry and binding energy are consistent with experiment, can the same data set be utilized to calculate intermolecular vibrational modes; and 2) what are the advantages and pitfalls of the various models in regard to the complexity of calculation, the approximations made, and the nature of the results obtained?

# Experimental Procedures.

Experimental data pertaining to the benzene(Ar)<sub>1</sub> vdW vibronic spectrum are obtained employing the experimental apparatus and procedures similar to those used previously to study vdW clusters.<sup>8</sup> The benzene(Ar)<sub>1</sub> S<sub>1</sub> + S<sub>0</sub> spectrum is recorded using a pulsed supersonic molecular jet expansion in combination with 1-color time-of-flight mass spectroscopy (TOFMS). A single Nd<sup>+3</sup>/YAG pumped LDS 698 dye laser whose output is frequency doubled and then mixed with the Nd<sup>+3</sup>/YAG 1.064  $\mu$ m fundamental is used to probe the 6<sup>1</sup><sub>0</sub> region of the benzene(Ar)<sub>1</sub> cluster. A 5% Ar in He mixture is placed inline with liquid benzene in a trap maintained at room temperature. This three component gas mixture is then expanded using a pulsed nozzle maintained at 100 psig backing pressure. Apparatus chamber pressure is maintained at or below 4 x 10<sup>-6</sup> torr during the experiment.

# Theoretical Considerations.

The normal coordinate analyses of the vdW clusters are conducted employing the GF methods of Wilson.<sup>10</sup> These methods involve solving the characteristic equation of 3N-6 coupled harmonic oscillators for its 3N-6 non-zero eigenvalues and eigenvectors. The approach is to treat the vdW cluster as a "giant molecule" and treat both the intramolecular vibrational modes and the intermolecular vdW modes simultaneously. The

intermolecular vdW potential field used in the analyses is expressed in an intermolecular coordinate system. In this coordinate system, the intermolecular force field is diagonal in the 3N dimensional space.

The cluster constituent intramolecular vibrational frequencies are considerably higher than those of the vdW modes. A reasonable approximation in this context then is to assume that the intramolecular modes are completely uncoupled from the low frequency vdW modes. Thus, the intramolecular modes are taken to be those of the cluster con-The constituent force fields are generated using the central stituents. force approximation<sup>11</sup> including out-of-plane motion terms. Since the vdW modes arise from the restriction of cluster constituent translations and rotations and the central force approximation adequately reproduces these degrees of freedom, the necessary uncoupling of the intramolecular modes and the intermolecular vdW modes is maintained along with providing adequate vdW mode calculational results. Other, more sophisticated force field approximations are tested in the calculations; the central force approximation is determined to be adequate for calculations of the vdW modes. The only restriction which applies is that the intramolecular field yields mode eigenvalues in the proper frequency regions.

Within the central force approximation, the force field contains only diagonal terms in the internuclear coordinate system. This diagonal force field provides a simple and convenient means of using approximate force constants for intramolecular motion in the calculations. The intramolecular force constants chosen are those pertaining to general functional group stretches and bends.<sup>11</sup>

The most convenient choice for a coordinate system as the working basis for matrix diagonalization is the Cartesian system. This

coordinate system is chosen since it is the system used in the present cluster configuration calculations, and more importantly, it is the coordinate system in which the  $\underline{G}^{-1}$  matrix is diagonal and obvious.

In order to combine the intermolecular and intramolecular force fields algebraically, the force fields are transformed into the Cartesian coordinate system. The transformation yields two 3N dimensional F matrices. The intramolecular F matrix is designated as  $\underline{F}_{n}$  and the intermolecular <u>F</u> matrix is designated as <u>F'</u>. The <u>F</u> matrix consists of two diagonal blocks containing the coordinates of cluster constituent intramolecular motion. The F' matrix contains two off-diagonal blocks and diagonal entries corresponding to the "perturbations" yielding the vdW motion. Adding these two matrices results in the "giant molecule" F matrix of order 3N. This matrix is left-multiplied by the <u>G</u> matrix and numerically diagonalized with the eigenvalues and eigenvectors being determined in the usual fashion. Upon diagonalization, the eigenvalues and eigenvectors of intramolecular motion are identified along with those corresponding to cluster translation and rotation. These modes are discarded and the remaining modes are the eigenvalues and eigenvectors of the ground state vdW modes.

The intermolecular force constants used in the normal coordinate analysis are generated from the intermolecular vdW potential by making a harmonic oscillator approximation. Within this approximation, the force constant is simply the second derivative of the potential function,<sup>9</sup>

$$U(r_{ij}) = \sum_{i=1}^{n} \sum_{j=1}^{m} \left[ \left\{ \frac{A^{kl}}{r_{ij}^{12}} - \frac{C^{kl}}{r_{ij}^{6}} \right\} (1 - \delta_{HB}^{ij}) + \frac{332.0q_{i}q_{j}}{Dr_{ij}} + \frac{332.0q_{i}$$

The potential function contains a general nonbonding potential (NB), a monopole charge potential (MC), and a general hydrogen bonding potential (HB) ns a general nonbonding potential (NB), a monopole charge potential (MC), and a general hydrogen bonding potential (HB) in a Lennard-Jones (6-12-1-10-12) form. The total intermolecular interaction is taken as a sum of pairwise atom-atom interactions over all the atoms of each cluster constituent. The  $r_{ij}$ 's are the atom-atom distances between atom i on constituent k and atom j on constituent 1. The  $r_{ij}$ 's represent the coordinates in which the intermolecular force field is diagonal. The second derivative of Equation 2.1 with respect to  $r_{ij}$  gives the force constant of the atom-atom interaction as,

$$K_{jj}(r_{ij}) = \sum_{i=1}^{n} \sum_{j=1}^{m} \left[ \left\{ \frac{12 \cdot 13 \cdot A^{kl}}{r_{ij}^{14}} - \frac{6 \cdot 7 \cdot C^{kl}}{r_{ij}^{8}} \right\} \times (1 - \delta_{HB}^{ij}) + \frac{2 \cdot 332 \cdot 0q_{i}q_{j}}{Dr_{ij}^{3}} + \left\{ \frac{12 \cdot 13A^{'kl}}{r_{ij}^{14}} - \frac{10 \cdot 11 \cdot C^{'kl}}{r_{ij}^{14}} \right\} \delta_{HB}^{ij} \right]$$

in which the  $K_{ij}$ 's are elements of the <u>F</u>' matrix expressed in intermolecular coordinates. These terms are evaluated at the equilibrium configuration of the cluster assuming that the cluster constituents are frozen with regard to intramolecular motion. The potential term coefficients used in the configurational and intermolecular vibrational mode calculations (Equations 2.1 and 2.2) on the systems studies are derived from the theory and data set described by Scheraga et al.<sup>9</sup>

Three additional models are employed to study the benzene(Ar)<sub>1</sub> vdW cluster as a test case. In these models, the vdW cluster is assumed to be a "diatomic molecule" in the sense that the system is considered to be composed of two particles, the benzene molecule (solute) and the

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argon atom (solvent). The vdW modes are assumed to arise from restric ted one-dimensional motion of the benzene molecule relative to the argon atom. Only the atom-molecule benzene(Ar)<sub>1</sub> cluster will be considered using these models since its vdW modes are easily characterized by simple translations along any of the three Cartesian axes. The vdW stretching mode is taken as motion restricted to the one-dimensional translation moving the cluster constituents apart in opposite directions. The vdW bending modes are considered to be motions restricted to one-dimensional translations which move the cluster constituents parallel to one another in opposite directions.

In the first model considered, one-dimensional potential curves are mapped out by translating the solvent atom relative to the solute molecule in one of the three Cartesian directions. The intermolecular vdW potential is assumed to be represented by a Taylor series expansion about the equilibrium intermolecular distance.  $R_0$  in the form,

$$U(R) = U(R_0) + \left(\frac{dU}{dR}\right) \frac{R}{R=R_0} + \frac{1}{2} \left(\frac{d^2U}{dR^2}\right) \frac{R^2}{R=R_0}$$

2.3

$$\frac{1}{6} \left( \frac{d^3 U}{dR^3} \right)_{\substack{R=R_0}} R^3 + \frac{1}{24} \left( \frac{d^4 U}{dR^4} \right)_{\substack{R=R_0}} R^4 + \dots$$

The expansion coefficients are evaluated by a polynomial fit to the potential curves taking the displacement vector R as the independent variable and U(R) as the dependent variable. The second-order polynomial fit coefficient determines the effective harmonic force constant governing the frequency of bound state motion. The energy of this motion is given by<sup>12</sup>

$$w_{\rm e} = \frac{1}{2\pi c} \left(\frac{k_{\rm eff}}{\mu}\right)^{1/2} \qquad 2.4$$

2.5

with

$$k_{eff} = \left(\frac{d^2 U}{dR^2}\right) = 2(\text{second-order coefficient})$$

$$R = R_0$$

$$\frac{1}{\mu} = \frac{1}{m_1} + \frac{1}{m_2}$$

$$m_1 = \text{solute mass} \qquad m_2 = \text{solvent mass}$$

The third and fourth order polynomial fit coefficients represent an anharmonic correction term to the energy. From perturbation theory, the correction can be written to first order as<sup>12</sup>

$$\omega_{e} \chi_{e} = \frac{3h^{2}}{32\pi^{4} \omega_{e}^{2} \mu^{2} c^{2}} \left( \frac{5g^{2}h}{8\pi^{2} \omega_{e}^{2} \mu c} - j \right)$$

$$1 \ (d^{3}U)$$

with

g = (third-order coefficient) = 
$$\frac{1}{6} \left( \frac{d^2 U}{dR^3} \right)$$
  
j = (fourth-order coefficient) =  $\frac{1}{24} \left( \frac{d^4 U}{dR^4} \right)$   
R=R<sub>0</sub>

Higher order terms are neglected in the anharmonic corrections since  $w_e \gg w_e | \chi_e >>$  other corrections. A polynomial least-squares fit to tenth order in R is determined to be sufficient to faithfully reproduce the one-dimensional potential curves generated via translation. In passing we note that the Taylor series expansion could also be evaluated directly by taking successive derivatives of Equation 2.1. This may be the method of choice if one is only interested in the lowest order terms in evaluating Equation 2.4 or 2.5. or if one is interested in a more "exact" reproduction of the potential curves in the vicinity of the dissociation limit.
The second model considered involves fitting the one-dimensional potential energy curves, derived in the same manner as previously discussed, to a Morse function of the form.<sup>15</sup>

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$$U(R) = D_{e}(e^{-2\beta(R-R_{o})} - 2e^{-\beta(R-R_{o})}). \qquad 2.6$$

The energy levels and anharmonic corrections are evaluated using

$$\omega_{\rm e} = \beta \left( \frac{D_{\rm e}h}{2\pi^2 c \mu} \right) \qquad 2.7$$

$$\omega_{\rm e} \chi_{\rm e} = \frac{\beta h}{8\pi^2 c\mu}$$
 2.8

The last model used involves an energy level fit based upon the semi-classical JWKB method.<sup>13</sup> In this model, the energy levels are determined from the quantization of the action integral according to the Bohr-Sommerfeld restrictions. The governing equation is

$$I = (2\mu)^{-1/2} \oint [E - U_{eff}(R)]^{1/2} dR = h(v + 1/2)$$
 2.9

Energy level determination is accomplished by numerical integration of (2.9) taking  $U_{eff}(R)$  as the one-dimensional intermolecular vdW potential in a tenth order polynomial form. The path of integration is taken over one complete motion cycle with the boundary conditions established by the intermolecular potential at a specific energy E.

The three "diatomic molecule" models are similar to the linear oscillator model described by Leutwyler et al.<sup>16</sup> In all of the approximations, the atom-molecule systems are assumed to be composed of three uncoupled linear oscillators. The potential surface in these degrees of freedom can thereby be modeled by one-dimensional potential functions. The models presented here differ from that present in ref. 16 in the respect that the linear oscillator model<sup>16</sup> treats the system as strongly anharmonic; the Taylor series expansion, and Morse fit models treat anharmonicity from the perturbational standpoint. In this respect, the JWKB method is probably the most similar to the linear oscillator model since both approximations fit the intermolecular mode energies using the physical boundary conditions established by the intermolecular potential.

Results.

A. Benzene(Ar),

Figure 2.1 and Table 2.1 present the benzene(Ar)<sub>1</sub> vdW cluster  ${}^{1}B_{2u} + {}^{1}A_{1g}$  spectrum recorded in the region between 38561 cm<sup>-1</sup> and 38710 cm<sup>-1</sup> using 1-color TOFMS. The cluster  $6_0^1$  is "red shifted" by 21 cm<sup>-1</sup> with respect to the benzene  $6_0^1$ . The bathychromic shift is indicative of the greater binding energy in the cluster S<sub>1</sub> state relative to the cluster S<sub>0</sub> state. Three vibronic features are observed to the blue of the cluster  $6_0^1$ . No features are observed in the symmetry "forbidden" benzene  $0_0^0$  region; therefore, the cluster must have at least a three-fold axis of symmetry.

Figure 2.2 and Table 2.2 contain the calculational results of the ground state configuration and vdW modes of benzene(Ar)<sub>1</sub>. Configurational calculations yield a single geometry of minimum energy for the cluster possessing  $C_{6v}$  symmetry, Figure 2.2. In this geometry, the argon atom lies 3.44 Å above the benzene molecular plane along the z (six-fold) axis. The ground state cluster binding energy is calculated at 287 cm<sup>-1</sup> which makes the excited state binding energy 308 cm<sup>-1</sup>. The calculated intermolecular distance of 3.44 Å compares well with that of



Mass selective  $S_1 + S_0$  spectrum and calculated ground state vdW modes of benzene(Ar)<sub>1</sub>. Energy scale is relative to benzene(Ar)<sub>1</sub>  $6_0^1$  transition (38587.6 cm<sup>-1</sup>). Nozzle backing conditions:  $P_0 = 100$  psig,  $T_0 = 300$  K. Peak positions and assignments as per Table 2.1 and Figure 2.2.

## TABLE 2.1

vdW spectal features in benzene(Ar) $_1 6_0^1$  region and calculated ground state vdW modes (refer to Figure 2.1).

Energy Relative to Cluster $6_0^1$ (cm <sup>-1</sup> )	Calculated Ground <sup>a</sup> State Energy (cm <sup>-1</sup> )	Assignment <sup>a</sup>
0 (38587.6)		6 <mark>1</mark>
	11 (b <sub>xy</sub> )	
30.0		$6^1_0 b_{xyo}^2$
39.7	40 (s <sub>z</sub> )	$6^{1}_{0}$ s <sub>zo</sub>
61.8		$6^1_0 b_{xy0}^4$

a) vdW mode representations as per Figure 2.2.

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Calculated ground state minimum energy configuration (a) and eigenvalue/ eigenvector vdW modes (b)-(d) for benzene(Ar)<sub>1</sub>. Cluster symmetry is  $C_{6v}$ with an equilibrium intermolecular distance of 3.44 Å. Eigenvectors are normalized and displayed at 2x magnification (2 Å total displacement).

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Model	s <sub>z</sub> (cm <sup>-1</sup> ) a	b <sub>xy</sub> (cm <sup>-1</sup> ) a
Taylor Series	40.88 (1.51)	9.80 (.03)
Morse Fit	39.47 (1.36)	10.54 (.10)
JWKB	40.05 (1.43)	9.71 (.02)
Normal Coordinate Analysis	40.0	11.0

Calculated ground state vdW mode energies for benzene(Ar)<sub>1</sub>.

a) Energy presented is for a harmonic oscillator model. Values in parentheses are first order anharmonicity corrections calculated from "diatomic molecule" models. vdW mode representations as per Figure 2.2. 3.45  $\pm$  .2 Å obtained from rotational analysis,<sup>15</sup> adding independent proof to the adequacy of the calculations in predicting detailed information on cluster structure.

The eigenvalue energies from the normal coordinate analysis of  $benzene(Ar)_1$  are 40 cm<sup>-1</sup> and 11 cm<sup>-1</sup> for the vdW stretch and vdW bends, respectively. The eigenvector normal modes, Figure 2.2, reveal that the vdW stretch entails purely perpendicular motion of the argon atom relative to the benzene molecular plane. Furthermore, the calculations reveal that the two-fold degenerate vdW bending mode involves some combination of motion parallel to the benzene molecular plane. Both of these eigenvector results are consistent with group theoretical arguments as expected.

The "diatomic molecule" model calculations yield three sets of vdW mode energies. The average mode frequencies are 40 cm<sup>-1</sup> for the vdW stretch,  $s_z(a_1)$  and 10 cm<sup>-1</sup> for the vdW bends  $b_{xy}(e_1)$ . Figures 2.3 and 2.4 show the details of the one-dimensional potential curve mappings and the results of the model calculations. Note that the z-direction potential curve modeling the vdW stretch looks surprisingly similar in form to that of a typical diatomic molecule. All models yield both adequate potential curve fits and consistent vibrational energy level structures.

B. s-Tetrazine(Ar)<sub>1</sub>.

Figure 2.5 presents the results of the ground state configuration and vdW mode analysis of s-tetrazine(Ar)<sub>1</sub>. Only the normal coordinate analysis vibrational calculation is presented since this method yields the most informative results for our purposes and the consistency between the eigenvalue/eigenvector results and the "diatomic molecule" results has already been shown for the benzene(Ar)<sub>1</sub> case. Configurational calculations yield a single cluster geometry of minimum


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#### Figure 2.3

z-direction (vdW stretch) potential energy mapping of benzene(Ar)<sub>1</sub>. Coordinate system is as shown in Figure 2.2. Translation is along z the axis with x and y coordinates at equilibrium intermolecular distance values. Translation is displayed relative to equilibrium intermolecular distance, 3.44 Å. (6-12) potential energy mapping is represented by o: Taylor series expansion and energy levels are represented by ----; Morse fit potential energy curve is represented by ----. Vibrational mode constants as per Table 2.2.



x(y)-direction (two-fold degenerate vdW bend) potential energy mapping of benzene(Ar)<sub>1</sub>. Coordinate system is as shown in Figure 2.2. Translation is along x(y) axis with z and y(x) coordinates at equilibrium intermolecular distance values. Translation is displayed relative to equilibrium intermolecular distance 3.44 Å. (6-12) potential energy mapping is represented by o; Taylor series expansion and energy levels are represented by —; Morse fit potential energy curve is represented by ---. Vibrational mode constants as per Table 2.2.



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## Figure 2.5

Calculated ground state minimum energy configuration (a) and eigenvalue/ eigenvector vdW modes (b)-(d) for s-tetrazine(Ar)<sub>1</sub>. Cluster symmetry is  $C_{2V}$  with an equilibrium intermolecular distance of 3.45 Å. Eigenvectors are normalized and displayed at 2x magnification (2 Å total displacement).

energy possessing  $C_{2v}$  symmetry, Figure 2.5. In this geometry, the argon atom lies 3.45 Å above the s-tetrazine molecular plane along the z (twofold) axis. The calculated ground state cluster binding energy is 295 cm<sup>-1</sup>. The calculated intermolecular distance of 3.45 Å compares extremely well with the intermolecular distance of 3.45 Å obtained from rotational analysis. The calculated binding energy of 295 cm<sup>-1</sup> also lies within the experimental limits of 254 <  $D_0^{''}$  < 332 cm<sup>-1.5</sup> Again, the calculations and experiment are in exact agreement.

The normal coordinate analysis eigenvalues are 41.0 cm<sup>-1</sup> for the vdW stretching mode  $s_z(a_1)$ , 9 cm<sup>-1</sup> for the vdW bending mode  $b_y(b_1)$ , and 12 cm<sup>-1</sup> for the vdW bending mode  $b_x(b_2)$ . The eigenvector normal modes. Figure 2.5 show that the vdW stretch is restricted to motion perpendicular to the s-tetrazine molecular plane while the vdW bends are restricted to motion parallel to the molecular plane. As in the benzene(Ar)<sub>1</sub> analysis, these results are consistent with group theoretical arguments.

C. Benzene( $CH_4$ )<sub>1</sub>.

The ground state configuration and vdW eigenvalues/eigenvectors for benzene(CH<sub>4</sub>)<sub>1</sub> are shown in Figure 2.6. The results presented for the geometry and binding energy of benzene(CH<sub>4</sub>)<sub>1</sub> are in good agreement with previous reports from this laboratory using an exponential-six and Lennard-Jones potential form.<sup>2</sup> In this geometry. Figure 2.6, the methane center-of-mass lies at 3.47 Å above the benzene molecular plane on the principle z (three-fold) axis. The cluster ground state binding energy is 540 cm<sup>-1</sup>. The normal coordinate analysis reveals six vdW vibrations, two being two-fold degenerate. The ground state vibrational energies are 82 cm<sup>-1</sup> for the vdW stretch s<sub>z</sub>(a<sub>1</sub>). BENZENE-METHANE (a) B.E.=540cm<sup>-1</sup> (c) 82cm<sup>-1</sup>  $s_z$  stretch( $a_1$ ) 28 cm<sup>-1</sup> t\_torsion(a, 16 cm<sup>-1</sup> 16 cm<sup>-1</sup> b\_bend(e) b\_ bend(e) 89 cm<sup>-1</sup> 89 cm - l t\_torsion(e) torsion (e

## Figure 2.6

Calculated ground state minimum energy configuration (a) and eigenvalue/ eigenvector vdW modes (b)-(g) for benzene( $CH_4$ )<sub>1</sub>. Cluster symmetry is  $C_{3v}$  with an equilibrium intermolecular distance of 3.47 Å. Eigenvectors are normalized and displayed at 2x magnification (2 Å total displacement).

16 cm<sup>-1</sup> for the vdW bends  $b_{xy}(e)$ , and 28 cm<sup>-1</sup> and 89 cm<sup>-1</sup> for the vdW torsions  $t_z(a_2)$  and  $t_{xy}(e)$ , respectively. The eigenvector normal modes. Figure 2.6, transform as the translational and rotational representations of the  $C_{3v}$  point group, as indicated. The vdW stretching mode transforms as the translation of the cluster constituents away from one another along the z (three-fold) axis. The vdW bending modes transform as some combination of cluster constituent translations in opposite directions perpendicular to the three-fold axis in the xy plane. One vdW torsion mode transforms as a rotation of the cluster constituents about the z (three-fold) axis in opposite directions. The remaining two vdW torsions transform as rotations about orthogonal axes perpendicular to the three-fold axis.

D. Benzene $(H_20)_1$ .

The calculated benzene(H<sub>2</sub>O)<sub>1</sub> geometry used in the normal coordinate analysis is similar to that calculated previously.<sup>3</sup> Only one minimum energy configuration, which has a binding energy of 504 cm<sup>-1</sup>, is found. As shown in Figure 2.7, the cluster geometry possesses  $C_s$  symmetry with the H<sub>2</sub>O center-of-mass located 3.15 Å above the benzene molecular plane.

Six ground state vdW vibrations are calculated for the  $C_s$  cluster geometry. Their corresponding eigenvalues and eigenvectors are shown in Figure 2.7. The six vdW modes consist of a vdW stretch at 159 cm<sup>-1</sup>, two vdW bends at 14 cm<sup>-1</sup> and 18 cm<sup>-1</sup>, and three vdW torsions at 40 cm<sup>-1</sup>. 50 cm<sup>-1</sup>, and 156 cm<sup>-1</sup>. The eigenvector normal modes transform as the translational and rotational representations of the  $C_s$  point group: the vdW stretch transforms as a z translation; the vdW bends transform as x and y translations; and the vdW torsions transform as  $R_x$ ,  $R_y$ , and  $R_z$  rotations.



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## Figure 2.7

Calculated ground state minimum energy configuration (a) and eigenvalue/ eigenvector vdW modes (b)-(g) for benzene( $H_2^{(0)}$ )<sub>1</sub>. Cluster symmetry is C<sub>s</sub> with an equilibrium intermolecular distance of 3.15 Å. Eigenvectors are normalized and displayed at 2x magnification (2 Å total displacement).

E. Benzene(NH<sub>3</sub>)<sub>1</sub>.

Configurational calculations on the benzene(NH<sub>3</sub>)<sub>1</sub> cluster reveal two minimum energy geometries similar to those obtained previously.<sup>3</sup> One cluster geometry possesses  $C_{3v}$  symmetry with a binding energy of 711 cm<sup>-1</sup> while the other possesses  $C_s$  symmetry and a binding energy of 608 cm<sup>-1</sup>. In the  $C_{3v}$  cluster, Figure 2.8, the NH<sub>3</sub> center-ofmass is located 3.23 Å above the benzene molecular plane along the z (three-fold) axis. In the  $C_s$  cluster, Figure 2.9, the NH<sub>3</sub> center-ofmass is located 3.29 Å above the benzene molecular plane.

Using the potential surfaces generated from these two configurations, six ground state vdW vibrations are calculated for each geometry. Their corresponding eigenvalues and eigenvectors are shown in Figures 2.8 and 2.9. The  $C_{3v}$  cluster ground state normal modes transform in the same manner as those of benzene( $CH_4$ )<sub>1</sub>. In the  $C_s$  cluster, the ground state normal modes transform similar to those of benzene( $H_20$ )<sub>1</sub>.

Discussion.

To compare the calculated and experimentally observed vdW modes. we assume that the intermolecular potential surface of the cluster is identical in both  $S_0$  and  $S_1$  electronic states. This assumption is justifiable if one considers that cluster fluorescence excitation and dispersed emission spectra are similar for the vdW vibronic transitions.<sup>4</sup> Furthermore, the small spectral shifts of the chromophore and the weak intensity of the vdW modes signify only small changes in cluster binding energy. This is probably indicative of only slight variations of the potential surface between  $S_0$  and  $S_1$  electronic states

Comparisons between calculation and experiment are made using group theoretical arguments based on the selection rules governing the

BENZENE-AMMONIA  $C_{3v}$ 97 cm<sup>-1</sup> (b) y (c) y (c) y (c) (c)(c)

#### Figure 2.8

Calculated ground state minimum energy configuration (a) and eigenvalue/ eigenvector vdW modes (b)-(g) for benzene(NH<sub>3</sub>)<sub>1</sub>. Cluster symmetry is  $C_{3v}$  with an equilibrium intermolecular distance of 3.23 Å. Eigenvectors are normalized and displayed at 2x magnification (2 Å total displacement).



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Calculated ground state minimum energy configuration (a) and eigenvalue/ eigenvector vdW modes (b)-(g) for benzene( $NH_3$ )<sub>1</sub>. Cluster symmetry is C<sub>s</sub> with an equilibrium intermolecular distance of 3.29 Å. Eigenvectors are normalized and displayed at 2x magnification (2 Å total displacement). vibronic transitions. Specifically, transition moment matrix elements are qualitatively analyzed using the crude adiabatic approximation for which the vibrational mode dependence on the electronic wave function is explicit. In this case, a standard Herzberg-Teller (HT) expansion and adiabatic wave functions are used<sup>14</sup> with the electronic wave function vibrational mode dependence truncated at second order.

Using this expansion, two unique types of spectra can be generated. First, one could consider that the vdW modes do not participate in the vibronic coupling scheme (Case I) and that they merely enter into the expansion as an additional scalar product (overlap integral). This argument dictates that only totally symmetric Franck-Condon progressions and combination band, are spectroscopically observed. Furthermore, the intensities of these features are solely derived from the cluster chromophore vibronic mode with which they are in combination. Alternately, one could consider the vdW modes to be capable of vibronic coupling (Case II). In this case, they enter into the HT transition moment equation in the same manner as other vibronically active modes. The operator responsible for these transitions would be of the form

$$\left(\frac{\partial^2 U}{\partial Q_{\mathfrak{s}} \partial q_{\mathsf{v}} d\mathsf{W}}\right) \begin{array}{c} Q_{\mathfrak{s}} = Q^{\mathfrak{o}}_{\mathfrak{s}}, q_{\mathsf{v}} d\mathsf{W} = q^{\mathfrak{o}}_{\mathsf{v}} d\mathsf{W} \end{array} \qquad 2.10$$

in which  $Q_6$  is the cluster  $v_6$  vibrational mode and  $q_{vdW}$  is a specific cluster vdW mode. This argument allows the possibility of observing nontotally symmetric vdW fundamentals with "borrowed" intensity due to interelectronic state mixing.

In the individual cluster discussions, the above two cases are considered in order to assign and understand the observed cluster vibronic spectra.

#### A. Benzene(Ar)<sub>1</sub>.

Comparison of the calculated ground state vdW vibrations of benzene(Ar)<sub>1</sub> and the experimental  ${}^{1}B_{2u} + {}^{1}A_{1g}$  vibronic spectrum. Figure 2.1. Table 2.1. and Table 2.2. shows that vibronic assignments can be made based upon calculations for all models analyzed. Considering the transition moment matrix elements for the benzene(Ar)<sub>1</sub> S<sub>1</sub> + S<sub>0</sub> transition under C<sub>6v</sub> symmetry, one should expect to observe totally symmetric combination bands of the vdW stretch built upon the cluster 6<sup>1</sup><sub>0</sub>. The selection rule for these combination bands is  $\Delta v = 0$ ,  $\pm 1$ ,  $\pm 2$ , ... Also, one should expect to observe nontotally symmetric vdW bend combinations with the 6<sup>1</sup><sub>0</sub> with the selection rule being  $\Delta v = 0$ ,  $\pm 2$ ,  $\pm 4$  ... These selection rules hold for both Case I and Case II type spectra and imply that the vdW modes do not enter into the vibronic intensity borrowing mechanism.

The calculated vdW stretching mode at 40 cm<sup>-1</sup> compares quite well with the experimental vibronic feature at 39.7 cm<sup>-1</sup> to the blue of the benzene(Ar)<sub>1</sub>  $6_0^1$ . Thus, this feature is assigned to the benzene(Ar)<sub>1</sub> vdW stretch/cluster  $6_0^1$  combination band  $6_0^1 s_z(a_1)_0^1$  based upon the  $\Delta v = 0$ , ±1, ±2 ... selection rule.

For vdW bending modes, only odd overtones are expected to be observed as pointed out above. The experimentally observed features at  $30.9 \text{ cm}^{-1}$  and  $61.8 \text{ cm}^{-1}$  to the blue of the cluster  $6_0^1$  correspond to overtone features of the vdW bends; using the  $\Delta v = 0$ ,  $\pm 2$ ,  $\pm 4$  ... selection rule, the former feature is the first overtone of the vdW bends and the latter is the third. Considering the  $30.9 \text{ cm}^{-1}$  feature as the first overtone places the symmetry forbidden bend fundamental at about 15.5 cm<sup>-1</sup>. This energy lies close to the calculated two-fold degenerate ground state vdW bend at 11 cm<sup>-1</sup>. Thus, these two spectral features are assigned to the benzene(Ar)<sub>1</sub> vdW bends first and third overtone/cluster  $6_0^1$  combination bands,  $6_0^1 - b_{xy}(e_1)_0^2$  and  $6_0^1 - b_{xy}(e_1)_0^4$ .

Ramifications of Herzberg-Teller vibronic coupling in the benzene(Ar)<sub>1</sub> cluster are obvious. From the derived vibronic selection rules and experimental observation, the benzene(Ar)<sub>1</sub> cluster spectrum is best assigned based upon benzene Herzberg-Teller coupling (i.e., the  $6_0^1$  feature is allowed) with vdW totally symmetric modes and combinations forming short, weak Franck-Condon progressions built upon the intense benzene transition. In addition, the calculated and observed<sup>15</sup> C<sub>6V</sub> cluster symmetry is verified by the vdW vibronic structure.

B. s-Tetrazine(Ar)<sub>1</sub>.

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The selection rules governing the s-tetrazine(Ar)<sub>1</sub> vdW vibronic transitions under  $C_{2v}$  symmetry arise from Case I Franck-Condon arguments. The totally symmetric vdW stretch should be observed to the blue of the cluster  $0_0^0$  following a  $\Delta v = 0, \pm 1, \pm 2, ...$  selection rule. The vdW bends should only be observed in odd overtones ( $\Delta v = 0, \pm 2, \pm 4$  ...) built on the allowed s-tetrazine  $0_0^0$  transition. As in the benzene(Ar)<sub>1</sub> cluster, no Case II distinction can be made for s-tetrazine(Ar)<sub>1</sub> and thus no vdW Herzberg-Teller vibronic coupling is expected. Unfortunately, a complete experimental spectrum showing the details of the s-tetrazine(Ar)<sub>1</sub> vdW modes is not, as yet, available. The only information in this regard is the identification of the vdW stretching mode at 44 cm<sup>-1</sup> to the blue of the cluster  $0_0^0$  by Levy et al.<sup>5</sup> Other vdW features at 66 cm<sup>-1</sup> and 108 cm<sup>-1</sup> to the blue of the cluster  $0_0^0$  are observed, but they are neither assigned nor are their spectra published.

Figure 2.10 and Table 2.3 compare the calculated ground state vdW mode frequencies with those observed in the s-tetrazine(Ar) $_1^{-1}B_{3u} + {}^{1}A_{g}$ 



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Schematic  ${}^{1}B_{3u} + {}^{1}A_{g}$  spectrum (from ref. 5) and calculated ground state vdW modes of s-tetrazine(Ar)<sub>1</sub>. Energy scale is relative to s-tetrazine(Ar)<sub>1</sub> 0<sup>0</sup><sub>0</sub> transition (18104.9 cm<sup>-1</sup>). Relative feature intensities are not shown. Feature positions and assignments as per Table 2.3 and Figure 2.5. vdW bends are represented by b<sub>1</sub> in schematic spectrum (see text for explanation).



vdW spectral features in s-tetrazine(Ar)<sub>1</sub>  $0_0^0$  region (s-tetrazine  ${}^{1}B_{3u} + {}^{1}A_g$ ) and calculated ground state vdW modes (refer to Figure 2.10).

Energy Relative to Cluster $0^{\circ}_{o}$ (cm <sup>-1</sup> ) c	Calculated Ground State Energy (cm <sup>-1</sup> ) <sup>b</sup>	Assignment b		
0 (18104.9)		0 <mark>0</mark>		
	9 (b <sub>y</sub> )			
	12 (b <sub>x</sub> )			
44 <sup>a</sup>	41 (s <sub>z</sub> )	s <sup>1</sup> <sub>zo</sub>		
66		$s_{zo}^{1}b_{xo}^{2}$ or $s_{zo}^{1}b_{yo}^{2}$		
108		$s_{zo}^{1b} s_{xo}^{4}$ or $s_{zo}^{1b} s_{yo}^{4}$		

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a) Observed and assigned in Reference 5.

b) vdW mode representations as per Figure 2.5.

c) From Reference 5.

vdW vibronic spectrum. The experimentally assigned vdW stretch at 44 cm<sup>-1</sup> to the blue of the cluster  $0_0^0$  corresponds to the 41 cm<sup>-1</sup> calculated ground state stretch. This motion, like that in the benzene(Ar)<sub>1</sub> cluster, involves perpendicular motion of the argon atom relative to the s-tetrazine molecular plane. Based upon ground state calculations, the feature 66 cm<sup>-1</sup> to the blue of the cluster  $0_0^0$  probably corresponds to a vdW stretch/vdW bend overtone combination band,  $s_2(a_1)_0^1 b_x(b_2)_0^2$  or  $s(a_1)_0^1 b_y(b_1)_0^2$ ; the feature at 108 cm<sup>-1</sup> probably corresponds to the next allowed bend overtone/stretch combination band  $s(a_1)_0^1 b_x(b_2)_0^4$  or  $s(a_1)_0^1 b_y(b_1)_0^4$ . No speculation with regard to which band is responsible for the features observed will be made since no spectra of sufficient sensitivity are available for analysis.

C. Benzene( $CH_4$ )<sub>1</sub>.

In a previous analysis of the benzene( $CH_4$ )<sub>1</sub> cluster,<sup>2</sup> three major vdW vibronic features were reported in the cluster 6<sup>1</sup><sub>0</sub> region: these features were assigned to a bend fundamental (27.3 cm<sup>-1</sup>), a stretch fundamental (32.3 cm<sup>-1</sup>), and a stretch overtone (51.4 cm<sup>-1</sup>). The assignments were made based upon the assumptions described in the Introduction.

Considering the HT transition moment matrix elements and assuming a Case I type spectrum, the selection rules for the benzene( $CH_4$ )<sub>1</sub> vdW vibronic transitions are  $\Delta v = 0, \pm 1, \pm 2, \dots$  for the vdW stretch and  $\Delta v$ = 0,  $\pm 2, \pm 4, \dots$  for the vdW bends and torsions when in combination with the cluster  $6_0^1$ .

The selection rules involved in Case II can be viewed in two ways. The selection rules can be derived using the calculated cluster symmetry of  $C_{3v}$ , or they can be derived by considering the cluster symmetry as  $C_{6v}$ . The latter situation arises since the methane centerof-mass is calculated at 3.47 Å above the benzene molecular plane. At this distance, the methane could be viewed as a sphere above the benzene molecular plane and, hence, the use of the  $C_{6v}$  point group to represent the vdW mode symmetries could be warranted.

In  $C_{3v}$  symmetry Case II, the selection rules for vdW mode combinations with the cluster  $6_0^1$  are  $\Delta v = 0, \pm 1, \pm 2, \ldots$  for all six vdW modes. If  $C_{3v}$  is the correct cluster physical symmetry and HT vdW coupling exists, all modes can be observed in the cluster  $6_0^1$  region. In  $C_{6v}$  symmetry, Case II, the selection rules for the vdW mode combinations with the cluster  $6_0^1$  are  $\Delta v = 0, \pm 1, \pm 2, \ldots$  for the vdW stretch  $s_z(a_1)$  and torsion  $t_z(a_2)$  and  $\Delta v = 0, \pm 2, \pm 4, \ldots$  for the two-fold degenerate vdW bends  $b_{xy}(e_1)$  and torsions  $t_{xy}(e_1)$ . In this approximate high symmetry, only the cluster  $6_0^1$  transition.

Comparison of the experimental  $6_0^1$  vibronic spectrum of benzene(CH<sub>4</sub>)<sub>1</sub> and the calculated ground state vdW vibrations is shown in Figure 2.11 and Table 2.4. The observed feature at 27.3 cm<sup>-1</sup> to the blue of the cluster  $6_0^1$  corresponds to the t<sub>z</sub> torsion calculated at 28 cm<sup>-1</sup>. Thus, this feature and its observed overtones at 51.4 cm<sup>-1</sup> and 73.5 cm<sup>-1</sup> are reassigned to  $6_0^1$  vdW torsion combination bands  $6_0^1$  t<sub>z</sub>(a<sub>2</sub>)<sub>0</sub><sup>1</sup>,  $6_0^1$  t<sub>z</sub>(a<sub>2</sub>)<sub>0</sub><sup>2</sup>, and  $6_0^1$  t<sub>z</sub>(a<sub>2</sub>)<sub>0</sub><sup>3</sup>, using the C<sub>3V</sub> Case II or C<sub>6V</sub> Case II  $\Delta v = 0$ , ±1, ±2, ... selection rule. The  $\Delta v = 0$ , ±1, ±2, ... selection rule suggests that for the benzene(CH<sub>4</sub>)<sub>1</sub> system vdW mode Herzberg-Teller vibronic coupling is an important component of the overall intensity mechanism. In the present case, the occurrence of the nontotally symmetric t<sub>z</sub> torsion progression implies that the vdW modes are

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Mass selective  $S_1 + S_0$  spectrum (ref. 2) and calculated ground state vdW modes of benzene(CH<sub>4</sub>)<sub>1</sub>. Energy scale is relative to benzene(CH<sub>4</sub>)<sub>1</sub>  $6_0^1$ transition (38567.6 cm<sup>-1</sup>). Feature positions and assignments as per Table 2.4 and Figure 2.6. vdW spectral features in benzene(CH<sub>4</sub>)<sub>1</sub>  $6_0^1$  region and calculated ground state vdW modes (refer to Figure 2.11).

Energy Relative to Cluster 6 <mark>1</mark> (cm <sup>-1</sup> ) a	Calculated Ground State Energy (cm <sup>-1</sup> ) b	Assignment <sup>b</sup>
0 (38567.6)		6 <mark>1</mark>
16.1	16 (b <sub>xy</sub> )	$6^1_0h^{1}_{xyo}$
27.3	28 (t <sub>z</sub> )	$6^1_{\text{o}} t^{-1}_{\text{zo}}$
32.3		$6^1_0 b_{xyo}^2$
48.4		6 <sup>1</sup> <sub>0</sub> b <sub>xyo</sub> <sup>3</sup>
51.4		$6^1_{o}t^2_{zo}$
64.6		6 <sup>1</sup> <sub>0</sub> h <sub>xyo</sub>
73.5		$6^1_{o}t^3_{zo}$
	82 (s <sub>z</sub>	

89 (t<sub>xy</sub>)

a) From Reference 2 and unpublished spectra.

b) vdW mode representations as per Figure 2.6.

vibronically active and that assuming them to be nonparticipants in the coupling mechanism oversimplifies the physics necessary to explain the cluster's spectroscopy.

The intense feature at 32.3 cm<sup>-1</sup> to the blue of the cluster  $6_{0}^{1}$ corresponds to the first overtone of the two-fold degenerate vdW bending mode,  $6_0^1 b_{xy}(e)_0^2$ , calculated at 16 cm<sup>-1</sup>. This identification is based upon the observation of a feature at 64.6  $cm^{-1}$  which corresponds to the third overtone of the bends. Furthermore, weak intensity features are observed at about 16.1  $cm^{-1}$  and 48.4  $cm^{-1}$  which could correspond to the vdW bend fundamentals and second overtones. The observation of these features adds proof to the arguments suggesting that the vdW modes are, at least, minor participants in the Herzberg-Teller vibronic coupling scheme. The cluster symmetry and, hence, spectroscopy are thereby also best described using the calculated C3v point group in conjunction with Herzberg-Teller coupling rather than the approximate C<sub>6v</sub> point group. Based upon this, the features at 32.3  $cm^{-1}$  and 64.6  $cm^{-1}$  are reassigned to vdW bend overtone combinations with the cluster  $6_0^1$ ;  $6_0^1$  b<sub>xy</sub>(e) $_0^2$  and  $6_0^1 b_{XY}(e)_0^4$  using the  $C_{3V}$  selection rule  $\Delta v = 0, \pm 1, \pm 2, \dots$  The features are assigned in Table 2.4 and Figure 2.11. The weak features at 16.1 cm<sup>-1</sup> and 48.4 cm<sup>-1</sup> are assigned to  $6_0^1 b_{xy}(e)_0^1$  and  $6_0^1 b_{xy}(e)_0^3$ Based upon the relative intensities displayed in the spectrum, vdW vibronic coupling is an important factor in the intensity of this progression involving the e symmetry bending modes.

Neither the vdW stretch  $s_z(a_1)$  nor the two-fold degenerate torsions  $t_{xy}(e)$  are observed in the  $6_0^1$  spectrum. This could be due to poor Franck-Condon factors for these vibronic transitions since they are both calculated to be at relatively high energies (ca. 82 cm<sup>-1</sup> and

89 cm<sup>-1</sup>, respectively). Moreover, these modes could be participating in VP since the total energy  $6_0^1 s_2(a_1)_0^1$  or  $6_0^1 t_{XY}(e)_0^1$  is close to that of, if not above, the cluster's  $S_1$  binding energy. Both the vdW stretch and  $t_{XY}$  torsions involve motion perpendicular to the benzene molecular plane: this motion could couple well to the VP process.

D. Benzene $(H_20)_1$ .

The  $S_1 \leftarrow S_0$  vibronic spectrum of benzene(H<sub>2</sub>O)<sub>1</sub> previously observed in this laboratory<sup>3</sup> possesses two unique spectral regions located around the cluster  $0_0^0$  and  $6_0^1$  containing vdW vibronic features. No vdW vibronic assignments were made in either region and no correlation between the regions was suggested.

Examination of the HT transition moment matrix elements using  $C_s$  symmetry and Case I considerations leads to the selection rules  $\Delta v = 0, \pm 1, \pm 2, \dots$  for the vdW s<sub>z</sub> stretch, b<sub>y</sub> bend, and t<sub>x</sub> torsion and  $\Delta v = 0, \pm 2, \pm 4, \dots$  for the b<sub>x</sub> bend, t<sub>y</sub> and t<sub>z</sub> torsions. Under Case II arguments, the selection rule is  $\Delta v = 0, \pm 1, \pm 2, \dots$  for all six vdW modes: all modes are capable of vibronic coupling.

Vibronic spectra of benzene(H<sub>2</sub>O)<sub>1</sub> in both cluster  $0_0^{O}$  and  $6_0^{1}$ regions are reproduced in Figure 2.12 along with the calculated ground state vdW mode energies. The observed feature at 5.2 cm<sup>-1</sup> (4.8 cm<sup>-1</sup>) to the blue of the cluster  $0_0^{O}$  ( $6_0^{1}$ ) transition corresponds to the vdW b<sub>x</sub> bend fundamental calculated at 14 cm<sup>-1</sup>. The observed feature at 16.2 cm<sup>-1</sup> (15.8 cm<sup>-1</sup>) to the blue of the cluster  $0_0^{O}$  ( $6_0^{1}$ ) transition corresponds to the calculated totally symmetric vdW b<sub>y</sub> bend at 18 cm<sup>-1</sup>. Additionally, the observed features at 34.6 cm<sup>-1</sup> (34.6 cm<sup>-1</sup>) and 49 cm<sup>-1</sup> (48.4 cm<sup>-1</sup>) to the blue of the cluster  $0_0^{O}$  ( $6_0^{1}$ ) transition are associated with the t<sub>z</sub>(a'') and t<sub>x</sub>(a') torsion fundamentals calculated at 40 cm<sup>-1</sup>



Mass selective  $S_1 \leftarrow S_0$  spectra (ref. 3) and calculated ground state vdW modes of benzene(H<sub>2</sub>0)<sub>1</sub>. Energy scale is relative to benzene(H<sub>2</sub>0)<sub>1</sub>  $0_0^0$  and  $6_0^j$  transitions (38168.6 cm<sup>-1</sup> and 38655.4 cm<sup>-1</sup>, respectively). Feature positions and assignments as per Table 2.5 and Figure 2.7.  $s_z$  and  $t_y$  vdW modes are not shown.

and 50 cm<sup>-1</sup>, respectively. The occurrence of the nontotally symmetric fundamentals implies that the  $\Delta v = 0$ ,  $\pm 2$ ,  $\pm 4$ ,, ... selection rule for Case I in which the vdW modes are not vibronically coupled is violated. The violation suggests that the vdW modes participate in the vibronic coupling scheme and that the  $\Delta v = 0$ ,  $\pm 1$ ,  $\pm 2$ , ... selection rule should apply to all six vdW modes (Case II). Based upon this, the spectra are best assigned, Table 2.5 and Figure 2.12, using both nontotally and totally symmetric vdW progressions.

Assigning the benzene( $H_2O$ )<sub>1</sub> spectra using the C<sub>S</sub> point group representations corroborates the calculated cluster geometry. Treating the cluster in approximate high symmetries, such as C<sub>2v</sub>, leads to selection rules which are clearly violated when applied to spectral observation and assignment. Specifically, the bend and torsion fundamentals are forbidden under these higher symmetry approximations. Furthermore, the spectral assignments using C<sub>S</sub> symmetry arguments suggest that the water constituent is likely located above the benzene molecular plane.

Neither the vdW s<sub>z</sub> stretch nor the t<sub>y</sub> torsion are observed in either spectral region. This as in the benzene(CH<sub>4</sub>)<sub>1</sub> case, probably results from poor Franck-Condon factors for these particular modes; they are both calculated to be at relatively high energies (ca. 159 cm<sup>-1</sup> and 156 cm<sup>-1</sup>, respectively). Moreover, in the 6<sup>1</sup><sub>0</sub> region, these modes could be participating in VP since the total energy of the system at these levels (486.8 + 159 cm<sup>-1</sup> for the stretch; 486.8 + 156 cm<sup>-1</sup> for t<sub>y</sub>) is close to, if not above, the cluster binding energy (ca. = 500 cm<sup>-1</sup>). The decrease in the hypsochromic shift and the shift of intensity maximum in the vdW manifold in going from the cluster 0<sup>0</sup><sub>0</sub> to the 6<sup>1</sup><sub>0</sub> may also be indicative of the VP process.

TABLE 2.0
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vdW spectral features in benzene( $H_20$ )<sub>1</sub>  $0^0_0$  and  $6^1_0$  region and calculated ground state vdW modes (refer to Figure 2.12).

Energy Relative to <sup>a</sup> Cluster 0 <sup>0</sup> (cm <sup>-1</sup> ) or 6 <sup>1</sup> (cm <sup>-1</sup> )	Calculated Ground State Energy (cm <sup>-1</sup> ) b	Assignment <sup>b</sup>
0 (38168.6)		00
5.2	14 (b <sub>x</sub> )	b <sub>xo</sub> <sup>1</sup>
16.2	18 (b <sub>y</sub> )	b <sub>yo</sub> <sup>1</sup>
21.4		$b_{xo}^{1}b_{yo}^{1}$
25		$b_{x0}^{2}b_{y0}^{1}$
31.9		b 2 yo
34.6	40 (t <sub>z</sub> )	$t_{zo}^{1}$
39.4		$t_{zo}^{1}b_{xo}^{1}$
45.7		b 3 yo
49	50 (t <sub>x</sub> )	$t_{xo}^{1}$
50.8		$t_{zo}^{1}b_{yo}^{1}$
67.4		$t_{xo}^{1}b_{yo}^{1}$
70.3		$t_{zo}^2$
99.5		t <sup>2</sup> xo
103.5		$t_{zo}^{3}$
-	156 (t <sub>y</sub> )	
-	159 (s <sub>z</sub> )	
0 (38655.4)		6 <mark>1</mark>
4.8	14 (b <sub>x</sub> )	$6^{1}_{0}b^{1}_{x0}$
15.8	18 (b <sub>v</sub> )	$6^1_0 b_{VO}^{-1}$

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## TABLE 2.5 (Continued)

25.5		$6^1_0 b^2_{x0} b^1_{y0}$
30.3		$6^1_0 b_{y0}^2$
34.6	40 (t <sub>z</sub> )	$6^1_{O}t^{-1}_{ZO}$
39.6		$6^1_0 t_{z0} b_{x0}^1$
48.8	50 (t <sub>x</sub> )	$6^1_{o}t^{1}_{xo}$
60.5		$6^1_{0}b_{y0}^4$
97.8		$6^1_0 t^2_{X0}$
101.0		$6^1_0 t_{z0}^3$
	156 (t <sub>v</sub> )	

159 (s<sub>z</sub>)

a) From Reference 3.

b) vdW mode representations as per Figure 2.7.

#### E. Benzene $(NH_3)_1$ .

Benzene(NH<sub>3</sub>)<sub>1</sub> clusters recently observed in this laboratory<sup>3</sup> yield spectra in both the cluster  $0_0^0$  and  $6_0^1$  regions. Two cluster geometries are calculated for the system, one possessing  $C_s$  symmetry and the other possessing  $C_{3v}$  symmetry. From symmetry arguments the  $C_s$  symmetry cluster is the only contributor to the  $0_0^0$  spectrum while both cluster geometries contribute to the  $6_0^1$  spectrum. Neither of these spectra were analyzed nor assigned in the initial observation since they are so complicated. They were merely presented as an indication of the notion that cluster vibronic spectra can sometimes be very extensive and congested.

The benzene(NH<sub>3</sub>)<sub>1</sub> C<sub>s</sub> symmetry cluster follows the same vdW vibronic selection rules as derived for the benzene(H<sub>2</sub>O)<sub>1</sub> cluster. In this symmetry, the selection rules for Case I are  $\Delta v = 0, \pm 1, \pm 2, ...$  for the vdW stretch, b<sub>y</sub> bend, and t<sub>x</sub> torsion and  $\Delta v = 0, \pm 2, \pm 4, ...$  for the b<sub>x</sub> bend, t<sub>y</sub> and t<sub>z</sub> torsions. In Case II the selection rule is  $\Delta v = 0, \pm 1, \pm 2, ...$  for all six vdW modes. Furthermore, the calculated cluster geometries are qualitatively similar. Hence, their 0<sup>0</sup><sub>0</sub> spectra should be qualitatively similar: this is borne out in both experimental results and vibrational mode calculations.

The benzene(NH<sub>3</sub>)<sub>1</sub>  $C_{3v}$  symmetry cluster follows the same vibronic selection rules as presented for benzene(CH<sub>4</sub>)<sub>1</sub>. Here the selection rules are either  $\Delta v = 0, \pm 1, \pm 2, \dots$  for the vdW stretch and  $\Delta v = 0, \pm 2, \pm 4, \dots$  for the vdW bends and torsions (Case I) or  $\Delta v = 0, \pm 1, \pm 2, \dots$  for all six vdW modes (Case II).

The calculated ground state vibrational energies and the observed cluster  $0_0^0$  and  $6_0^1$  vibronic spectra are compared in Figure 2.13. Due to



Mass selective  $S_1 \leftarrow S_0$  spectra (ref. 3) and calculated ground state vdW modes of benzene(NH<sub>3</sub>)<sub>1</sub>. Energy scale is relative to benzene(NH<sub>3</sub>)<sub>1</sub>  $0_0^0$  and  $6_0^1$  transitions for  $C_s$  cluster (38021.1 cm<sup>-1</sup> and 38514.7 cm<sup>-1</sup>, respectively). Feature positions and assignments as per Table 2.6 and Figure 2.8 and 2.9.  $C_{3v}$  cluster  $s_z$  and  $t_{xy}$  modes are not shown.

the complex nature of the spectra which possibly results from hot bands, only tentative assignments of the vdW mode progressions are made. The tentative assignments are based upon both vibrational mode calculations and upon inference from the benzene( $H_2O$ )<sub>1</sub> and benzene( $CH_4$ )<sub>1</sub> cluster spectra.

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In the  $0_0^0$  spectrum, the most intense low energy feature is assigned to the origin of the cluster's  $S_1 + S_0$  transition. The smaller intensity features to the red of this feature are thus hot bands which yield the sequence structure in the  $0_0^0$  region. The observed feature at 8.8 cm<sup>-1</sup> to the blue of the cluster  $0_0^0$  corresponds to the nontotally symmetric vdW b<sub>x</sub> bend fundamental calculated at 15 cm<sup>-1</sup>. Additionally, the observed features at 15.0 cm<sup>-1</sup>, 45.1 cm<sup>-1</sup>, 54.2 cm<sup>-1</sup>, 99.6 cm<sup>-1</sup>, and 127.7 cm<sup>-1</sup> correspond to the calculated b<sub>y</sub> bend (21 cm<sup>-1</sup>), t<sub>z</sub> torsion (44 cm<sup>-1</sup>), t<sub>x</sub> torsion (48 cm<sup>-1</sup>), stretch (112 cm<sup>-1</sup>), and t<sub>y</sub> torsion (125 cm<sup>-1</sup>). The occurrence of the nontotally symmetric fundamentals suggests that the vdW modes participate in vibronic coupling (Case II) and that the  $\Delta v = 0$ , ±1, ±2, ... selection rule should apply to all six vdW modes. Using this selection rule, the spectrum is best assigned, Table 2.6, using both nontotally and totally symmetric vdW progressions.

In the  $6_0^1$  spectrum, the most intense low energy feature is assigned to the cluster  $6_0^1$  vibronic origin. This assignment bears a resemblance to the benzene(H<sub>2</sub>O)<sub>1</sub>  $6_0^1$  spectrum in the respect that the  $6_0^1$ features for both clusters are red shifted relative to that observed at the S<sub>1</sub> + S<sub>0</sub> origins (ca. 25 cm<sup>-1</sup> for benzene(NH<sub>3</sub>)<sub>1</sub> and 35 cm<sup>-1</sup> for benzene(H<sub>2</sub>O)<sub>1</sub>.

Table 2.6 presents the  $C_s$  symmetry geometry assignments in the  $6_0^1$  region. The assignments are made using the vdW fundamentals identified

	TA	BL	E	2		6
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Energy Relative to <sup>a</sup> Cluster 0 <sup>0</sup> (cm <sup>-1</sup> ) or 6 <sup>1</sup> (cm <sup>-1</sup> )	Calculated Ground State Energy (cm <sup>-1</sup> ) b	Assignment <sup>b,c</sup>
-39.2		by2
-36.7		
-33.4		$b_{x2}^{o}b_{y1}^{o}$
-31.3		by2bx0
-29.6		·
-26.7		b <sub>x1</sub> 0b <sub>v1</sub>
-19.2		b <sub>y1</sub>
- 6.7		b <sub>x1</sub>
- 3.8		
0 (38021.1)		00
8.8	15 (b <sub>x</sub> )	$b_{xo}^{1}$
15.0	21 (b <sub>v</sub> )	b l
17.9		b <sub>x0</sub> <sup>2</sup>
24.2		$b_{x0}^{1}b_{y0}^{1}$
26.7		b <sub>x0</sub> <sup>3</sup>
29.0		b <sup>2</sup> vo
32.1		$b_{x0}^{2}b_{y0}^{1}$
34.2		b 4
37.6		$b_{x0}^{1}b_{y0}^{2}$
41.7		x0 y0
45.1	44 (t <sub>2</sub> )	$t_{ro}^{-1}$

vdW spectral features in benzene(NH<sub>3</sub>)<sub>1</sub>  $0_0^0$  and  $6_0^1$  region and calculated ground state vdW modes (refer to Figure 2.13).
TABLE 2.6 (Continued)

50.9		
54.2	48 (t <sub>x</sub> )	t <mark>1</mark> xo
59.7		t j h j
64.7		$t_{xo}^{1}b_{xo}^{1}$
70.4		$t_{xo}^{1}b_{yo}^{1}$
75.5		t. 10 2
77.2		
80.1		t_10_10_1 t_x0b_x0b_y0
83.8		$t_{xo}^{1}h_{yo}^{2}$
99.6	112 (s <sub>z</sub> )	s <sub>zo</sub> <sup>1</sup>
101.8		$t_{xo}^{1}t_{zo}^{1}$
107.2		s <sup>1</sup> t <sup>1</sup> zotxo
111.8		$t_{x0} t_{z0} b_{x0} b_{x0}$
114.3		szobyo
119.3		$t_{x0}^{1}t_{z0}^{1}h_{x0}^{2}$
122.7		$s_{zo}^{1}b_{xo}^{1}b_{yo}^{1}$
127.7	125 (t <sub>y</sub> )	t 1 yo
- 9.0		$6^1_{o}h_{xo}^{1}$
- 4.9		
- 2.0		
0 (38514.7)		$6_0^1(C_s)$
2.0		
9.0	15 (b <sub>x</sub> )	$6^{1}_{0}b^{1}_{x0}$
13.1	21 (h <sub>y</sub> )	6 <sup>1</sup> <sub>0</sub> b <sub>y0</sub>
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TABLE 2.6 (Continued)

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26.2			$6^1_0 h_{XO}^3$
29.5			$6^1_0 h_{yo}^2$
32.0			$6^1_{0}b^2_{x0}b^1_{y0}$
34.4			6 <sup>1</sup> <sub>0</sub> b <sup>4</sup> <sub>x0</sub>
37.3			6 <sup>1</sup> <sub>0</sub> b <sup>2</sup> <sub>0</sub> b <sup>1</sup> <sub>x0</sub>
45.1		44 (t <sub>z</sub> )	$6^{1}_{0}t^{1}_{z0}$
48.7			
53.7			$6^{j}_{o}t^{1}_{zo}h^{1}_{xo}$
55.3		48 (t <sub>x</sub> )	$6^{1}_{0}t^{1}_{x0}$
58.6			$6^{1}_{0}t^{1}_{z0}h^{1}_{y0}$
61.0			$6^1_0 t_{z_0} b_{x_0}^2$
64.7			$6^{J}_{o}t^{1}_{xo}h^{1}_{xo}$
68.8			$6^{1}_{o}t^{1}_{xo}b^{1}_{yo}$
72.9	[0 (38587.6)]		$6_0^1(C_{3v})$
74.6			$6_0^1 t_{x0}^{1} b_{x0}^2$
79.5			$6^{j}_{o}t^{1}_{xo}h^{1}_{xo}h^{1}_{vo}$
83.3			$6_0^1 t_{x_0}^{1} b_{y_0}^2$
87.9		19 (b <sub>xy</sub> )	$6_0^1 h_{xy0}^{-1} (C_{3v})$
98.7		112 (s <sub>z</sub> )	$6^{1}_{0}s^{1}_{z0}$
100.4			$6_0^1 t_{x0} t_{z0}^{-1}$
103.9			$6_0^1 h_{xy0}^2 (C_{3v})$
107.1			$6_0^{\mathbf{j}} \mathbf{s}_{\mathbf{z}0}^{1} \mathbf{h}_{\mathbf{x}0}^{1}$
111.9			$6^{1}_{0}t^{-1}_{x0}t^{-1}_{z0}b^{-1}_{x0}$
114.0			$6^{1}_{0}$ s $\frac{1}{2}$ b $\frac{1}{2}$
123.4			$6^{1}_{0}s^{1}_{z0}b^{1}_{x0}b^{1}_{y0}$



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125.9	44 (t <sub>z</sub> )	$6_0^1 t_{z_0}^{-1} (C_{3v})$
-	97 (s <sub>z</sub> )	-
-	152 $(t_{xy})$	-

a) From Reference 3.

- b) vdW mode representations as per Figure 2.8 and 2.9.
- c)  $C_{3v}$  cluster  $6_0^1$  contributions tabulated relative to  $C_s$  cluster  $6_0^1$  origin as in Figure 2.13.

in the  $0^{0}_{0}$  spectrum and  $\Delta v = 0, \pm 1, \pm 2, \dots$  selection rule for all six vdW modes (Case II).

The contribution of the  $C_{3v}$  symmetry benzene(NH<sub>3</sub>)<sub>1</sub> cluster to the  $6_0^1$  spectrum is observed starting with the feature at 72.9 cm<sup>-1</sup> to the blue of the  $C_s$  cluster  $6_0^1$ . Assignment of this feature to the  $C_{3v}$  cluster  $6_0^1$  is based upon the observation that no intense features are seen in either the benzene(NH<sub>3</sub>)<sub>1</sub>  $0_0^0$  or the benzene(H<sub>2</sub>0)<sub>1</sub>  $6_0^1$  spectra at this energy. The observed feature at 15 cm<sup>-1</sup> to the blue of the  $C_{3v}$  cluster  $6_0^1$  corresponds to the two-fold degenerate vdW b<sub>xy</sub> bends at 19 cm<sup>-1</sup>. The observed feature at 53 cm<sup>-1</sup> to the blue of the  $C_{3v}$  cluster  $6_0^1$  is associated with the calculated  $t_z$  torsion at 44 cm<sup>-1</sup>. As in the benzene(CH<sub>4</sub>)<sub>1</sub> case, the observation of the b<sub>xy</sub> and  $t_z$  fundamentals implies that the vdW modes participate in the vibronic coupling mechanism (Case II) and that the  $\Delta v = 0, \pm 1, \pm 2, \ldots$  vibronic selection rule applies for all vdW modes. Based upon this, the best assignments for the  $C_{3v}$  cluster vdW vibronics are tabulated in Table 2.6.

F. Morse  $\frac{6}{R_0}$  Model.

Calculations of the ground state vdW modes of benzene(Ar)<sub>1</sub> are also conducted using a model described by Jortner et al.<sup>1.6,7</sup> This model contains a main feature which, at first glance, appears to make it generally applicable in predicting vdW stretching mode frequencies from calculated cluster binding energies and equilibrium intermolecular distances. However, the model turns out to be incorrect in this applica tion. The model employs a simple relationship between the cluster equilibrium intermolecular distance R<sub>0</sub> and the Morse potential parameter  $\beta$ . The relation is derived by equating the second derivative of Equation 2.6 with the second derivative of a single term (6-12) المتحد بحدون ومد بالمحديد

potential function. This derivation yields the relation  $\beta = \frac{6}{R_o}$  which can be used to predict the vdW stretching mode frequency by diatomic Morse fit methods.

Substituting  $\beta = \frac{6}{R_{a}}$  and the calculated ground state cluster binding energy into Equation 2.7 yields a vdW stretching mode energy of 47 cm<sup>-1</sup> for benzene(Ar), employing Scheraga's potential parameters. Even though the calculated vibrational energy is qualitatively correct, Figures 2.14 and 2.3 show that the  $\beta = \frac{6}{R_{a}}$  relation results in an inadequate reproduction of the potential curve, especially in the critical region near the equilibrium intermolecular distance. At  $R_0$ , the  $\beta = \frac{6}{R_1}$ fit overestimates the curvature of the potential by about 33% with respect to the curvature calculated from the one-dimensional potential energy mapping (1746 cm<sup>-1</sup>/Å<sup>2</sup> versus 1312 cm<sup>-1</sup>/Å<sup>2</sup>) and, therefore, overestimates the stretching mode energy. Additional calculations using the potential data described in ref. 1 and the above approximate model also result in an inadequate potential curve reproduction, Figure 2.15. The potential curvature at  $R_0$  using the  $\beta = \frac{6}{R_0}$  fit is overestimated by about 37% with respect to the corresponding curvature calculated via potential energy mapping (2322 cm<sup>-1</sup>/Å<sup>2</sup> versus 1695 cm<sup>-1</sup>/Å<sup>2</sup>). In this case, the vdW stretching mode is calculated at 55 cm $^{-1}$ .

The failure of the model under both data sets suggests that the model itself is inadequate in this application. The model fails in this application since the  $\beta = \frac{6}{R_0}$  relation holds exactly only in systems in which the molecule-atom Lennard Jones parameters have been determined <u>directly</u>. Applying this approximate model to a case in which the potential function is represented by pairwise <u>atom-atom</u> potentials does not take into account the differing contributions of each interaction to the



# Figure 2.14

z-direction (vdW stretch) potential energy mapping of benzene(Ar)<sub>j</sub> and  $\beta = \frac{6}{R_o}$  Morse fit. Translation is relative to equilibrium intermolecular distance, 3.44 Å. (6-12) potential energy mapping using data set from ref. 9 is represented by o; Taylor series expansion and energy levels are represented by ---;  $\beta = \frac{6}{R_o}$  Morse fit potential energy curve is represent by ---. Taylor series vibrational mode constant as per Table 2.2.  $\beta = \frac{6}{R_o}$  Morse fit vibrational constants are  $\omega_e = 47.22$  cm<sup>-1</sup>,  $\omega_e x_e =$ 1.94 cm<sup>-1</sup>.



# Figure 2.15

z-direction (vdW stretch) potential energy mapping of benzene(Ar)<sub>1</sub> and  $\beta = \frac{6}{R_o}$  Morse fit. Translation is relative to equilibrium intermolecular distance, 3.5 A. (6-12) potential energy mapping using data set from ref. 1 is represented by o; Taylor series expansion and energy levels are represented by ---- with  $\omega_e = 46.68 \text{ cm}^{-1}$  and  $\omega_e \chi_e = 1.81 \text{ cm}^{-1}$ ;  $\beta = \frac{6}{R_o}$ Morse fit potential energy curve is represented by ----  $\beta = \frac{6}{R_o}$  Morse fit vibrational mode constants are  $\omega_e = 54.84 \text{ cm}^{-1}$  and  $\omega_e \chi_e = 1.90 \text{ cm}^{-1}$ 

Exercise
 Exercise

potential energy and the equilibrium intermolecular distance. In this more complicated situation, no analytic relationship between  $\beta$  and R<sub>0</sub> exists.

Comparing the vdW stretching mode energies calculated using Jortner's and Scheraga's data sets suggests that a significant difference exists between the two data sets. Benzene(Ar)<sub>1</sub> configurational calculations using Jortner's data set yield a single cluster geometry of  $C_{By}$  symmetry with the argon atom located 3.5 Å above the benzene molecular plane. In this case, the ground state cluster binding energy is  $395 \text{ cm}^{-1}$ . This binding energy is 108 cm<sup>-1</sup> greater than that calculated using Scheraga's data set. The binding energy of 287  $cm^{-1}$  calculated using Scheraga's data set is probably more accurate since the benzene(Ar)<sub>1</sub> ground state binding energy should be very similar to that calculated for s-tetrazine $(Ar)_1$ . In the latter case, the calculated ground state binding energy of 295  $cm^{-1}$  compares well with that observed experimentally, 254 <  $D_0''$  < 332 cm<sup>-1</sup>.<sup>5</sup> Furthermore, generating Lennard-Jones parameters using the data set of ref. 1 yields a binding energy of 359  $cm^{-1}$  for s-tetrazine(Ar)<sub>1</sub> which is clearly not as accurate as the binding energy reported in this work. The difficulty is due to the consolidation of atom-atom parameters from different data sets. In this respect, great caution must be taken when consolidating parameters since each parameter set is, in general, only self-consistent and may have no meaning when combined with parameters from other sets.

## Conclusions

TOFMS studies have been employed to determine the general geo metry and symmetry of vdW clusters in the gas phase. Through computer modeling, a correlation between the details of the cluster geometry and spectral features has been demonstrated. Specifically, the parametric calculations yield useful information regarding cluster geometry, binding energy, and the vdW vibrations. These calculated results are consistent with experiment and serve as predictive and analytic tools which can be used to elucidate and understand the details of vdW cluster energetics.

Of the several models considered in studying the intermolecular vdW modes, simple diatomic approximations yield adequate results when applied to atom-molecule clusters. On the other hand, for moleculemolecule clusters a normal coordinate analysis is essential. The normal coordinate analysis is especially useful for analyzing systems which have little or no symmetry since no <u>a priori</u> knowledge of vdW mode nature is necessary to generate potential energy surface mappings.

Reassignments of and assignments to cluster vdW modes have been made based upon the knowledge gained from calculation. From comparison of calculation and experiment, several conclusions result. First, the actual excited state normal mode vdW frequencies are well fit by the calculated ground state cluster potential. This conclusion, though not surprising, gives independent proof of the invariance of the weak vdW potential between ground and excited electronic states as well as providing a means of using ground state vdW vibrational structure to predict vdW vibronic structure. Second, in the majority of the clusters analyzed, the observed vdW vibrations are those involving bending and torsional motions parallel to the aromatic  $\pi$  system. Furthermore, these modes are, in general, quite active in the Herzberg-Teller vibronic coupling mechanism and significant interelectronic state mixing results. Third, vdW motions which penetrate the aromatic  $\pi$  system have high

frequencies and are only observed in systems in which the Franck-Condon factors and binding energies are favorable. Finally, the observed vibronic structure supports the calculated cluster geometry in all cases.

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### REFERENCES

- M.J. Ondrechem, Z. Berkovitch-Yellin, and J. Jortner, J. Amer. Chem. Soc. <u>103</u>, 6586 (1981).
- 2. M. Schauer and E.R. Bernstein, J. Chem. Phys. 82, 726 (1985).
- J. Wanna, J.A. Menapace and E.R. Bernstein, J. Chem. Phys., <u>85</u>, 1795 (1986).
- 1. T.A. Stephenson and S.A. Rice, J. Chem. Phys. 81, 1083 (1984).
- D.V. Brumbaugh, J.E. Kenny, and D.H. Levy, J. Chem. Phys. <u>78</u>, 3415 72(1983).
- 6. M.L. Sage and J. Jortner, J. Chem. Phys. 82 5437 (1985).

- J.A. Beswick and J. Jortner, J. Chem. Phys. <u>68</u>, 2277 (1978); <u>69</u>, 512 (1978); <u>74</u>, 6725 (1981); <u>71</u> 4737 (1979).
- J. Wanna and E.R. Bernstein, J. Chem. Phys. <u>84</u>, 927 (1986), and ref. 2 and 3, for example.
- F.A. Momany, L.M. Carruthers, R.F. McGuire, and H.A. Scheraga, J. Phys. Chem. <u>78</u>, 1595 (1974); G. Nemethy, M.S. Pottle, and H.A. Scheraga, J. Phys. Chem. <u>87</u>, 1883 (1983).
- E.B. Wilson Jr., J.C. Decius, and P.C. Cross, "Molecular Vibrations, Theory of Infrared and Raman Vibrational Spectra," (McGraw-Hill Book Co., Inc., 1955).
- G. Herzberg, "Molecular Spectra and Molecular Structure: II Infrared and Raman Spectra of Polyatomic Molecules," (Van Nostrand Reinhold Co., 1945).
- 12. G. Herzberg, "Molecular Spectra and Molecular Structure: I Spectra of Diatomic Molecules." (Van Nostrand Reinhold Co., 1950).
- J.I. Steinfield, "Molecules and Radiation: An Introduction to Modern Molecular Spectroscopy," (MIT Press, 1978)
- 14. G. Fischer, "Vibronic Coupling: The Interaction Between Electron: and Nuclear Motions," (Academic Press, 1984)
- 15. K.H. Fung, H.L. Selzle and F.W. Schlag, Z. Naturforsch <u>36A</u>, 1978 (1981).

S. Leutwyler and A. Schmeizer, J. Chem. Phys. <u>79</u>, 4385 (1983); J.
 Boesiger and S. Leutwyler, Chem. Phys. <u>126</u>, 283 (1986)

### CHAPTER THREE

THE INTERMOLECULAR VIBRONIC TORSIONAL STRUCTURE IN SOLUTE/SOLVENT VAN DER WAALS CLUSTERS: BENZENE/METHANE, /DEUTEROMETHANE, AND /CARBON TETRAFLUORIDE

Introduction.

van der Waals (vdW) complexes of aromatic molecules with hydrocarbon solvents form a class of supramolecular systems whose intermolecular bonding has several interesting features. First, the intermolecular interaction is small and is dominated by long-range dispersive attractions and short-range exchange repulsions. These two features allow the interaction to be modeled using an intermolecular potential of known functional form. For example, the interaction can be modeled by additive atom-atom potentials set in a Lennard-Jones or an Exponential Six format.<sup>1</sup> Second, the interaction potential surface only changes slightly, if at all, upon electronic excitation of the cluster chromophore. Thus, small spectral shifts of the chromophore electronic transition and weak intensity intermolecular vdW mode vibronic transi tions are observed. $^{2,3}$  Third, the interaction results in the formation of specific minimum energy cluster configurations. These geometries are interesting since they give insight into the nucleation processes and solvation geometry occurring in both gas and condensed phase systems  $^{1}$ Fourth, the small binding energy of the complex and the low frequency odw vibrational modes are important since they play an essential role in

intramolecular vibrational redistribution (IVR) and vibrational predissociation (VP) cluster dynamic energy transfer processes.<sup>4</sup> And fifth, cluster structure, binding energy, vdW modes and dynamics (IVR and VP) are essentially dependent on the actual cluster structure and the intricacies of the intermolecular interaction.<sup>3,5</sup> In this paper, we report the spectroscopic results of the benzene/deuterated methane  $(ben(CD_4)_1)$  and benzene/carbon tetrafluoride  $(ben(CF_4)_1)$  clusters together with calculated modeling of selected cluster characteristics. The spectroscopic results include the  $\pi^* + \pi$  vibronic spectra of the clusters in their respective benzene constituent  $6\frac{1}{0}$  regions.

The calculated results include the geometry, the binding energy, and the full eigenvalue/eigenvector intermolecular vibrational structure for the electronic ground state of each cluster. The intermolecular ground state vibrational structure is modeled by two methods. (1) an intermolecular normal coordinate analysis (N(A) which determines all six intermolecular normal coordinate analysis (N(A) which determines all six intermolecular vdW mode fundamentals under a harmonic oscillator assumption.<sup>4</sup> and (2) a three dimensional hindered right before unalysis. (1) HRRA) for which an anisotropic perturbation the time upposed to the cluster solvents considered to be a perturbation to the constances potential well. The intermelecular targets as a decise to the constances obtained under a hordered to perturbation as

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ramifications of Herzberg-Teller (H-T) coupling for the observation of "forbidden" vibronic transitions.

The motivation for studying these systems concerns the elucida tion of the vdW torsional structure. Two limiting cases can be proposed in regard to the torsional structure of  $ben(CH_4)_1$ ,  $ben(CD_4)_1$ , and  $ben(CF_4)_1$ . In one case, the clusters possess free internal rotation between the cluster solute and solvent. The cluster solvent  $(CH_4, CD_4,$ or  $CF_4$ ) rotates freely in three dimensions against the benzene framework and the system is considered nonrigid. In the other case, the clusters possess torsional oscillations for which the cluster solvent librates against the benzene frame with a residence time long enough to give rise to "vibration like" motion in a rigid molecule regime.

Elucidation of the vdW torsions is of particular interest in these systems since the aforementioned limiting cases pose questions regarding the actual physics governing the torsional structure (1) do the clusters possess free internal rotation between the cluster solute and sourcent or do they possess forsional oscillations for which the we to constitute behave more or less barmonica of and (d) in erflet clusters of the dependence of the otermolecitar protection of the

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mixed with the Nd<sup>+3</sup>/YAG 1.064  $\mu$ m fundamental is used to probe the clusters' S<sub>1</sub> + S<sub>0</sub> transition in the isolated benzene 6<sup>1</sup><sub>0</sub> region (<sup>1</sup>B<sub>2u</sub> + <sup>1</sup>A<sub>1g</sub>;  $\pi^*$  +  $\pi$ ). Subsequent ionization of the clusters is accomplished using a R590 dye laser whose output is frequency doubled and mixed with the Nd<sup>+3</sup>/YAG 1.064  $\mu$ m fundamental. The ionization laser output is set at 45112 cm<sup>-1</sup>. A 5% mixture of deuterated methane or carbon tetrafluoride in helium is placed inline with liquid benzene in a trap at room temperature. The three component mixture is expanded using a pulsed molecular jet nozzle having a 500 micron orifice while maintaining 100 psig backing pressure. Apparatus chamber pressure is maintained a<sup>+</sup> or below 5 x 10<sup>-6</sup> torr during the experiments.

#### Theoretical Considerations.

The NCAs are conducted using the same methods as described in our previous publication on vdW cluster vibronic structure <sup>3</sup>. For  $ben(CD_1)_1$ , the calculated cluster ground state geometry, force field, and binding energy are taken as those of  $ben(CH_1)_1$ . Only the masses are changed for the deuterium hydrogen substitutions in the N(A). For  $ben(CH_4)_1$ , the cluster ground state geometry and binding energy are calculated via intermole rise geometry and binding energy are calculated via intermole rise energy minimization employing the methods previous device bet  $\frac{1}{2}$ . The N(A) is inducted via intermole rise energy minimization employing the methods previous device bet  $\frac{1}{2}$ .

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cluster solvent rotates freely in three dimensions against a fixed cluster solute framework. Application of a specified perturbation field results in the eventual restriction of this free rotation to torsional oscillation as the perturbation field magnitude is increased; that is, rotation ceases and vibrational oscillatory motion begins as the residence times of the solvent in the torsional potential well becomes longer.

In the 3D-HRRA, both cluster constituents are assumed internally rigid, their respective internal geometries remaining constant and at "equilibrium". The cluster solvents are taken as the rotating portions of the clusters since their zero field rotational constants are orders of magnitude greater than those of the cluster solute. The hindered solvent rotations are presupposed to contribute to the spectral features observed along with the intermolecular bending and stretching modes.

The 3D-HRRA involves setting up a molecule fixed coordinate system (x,y,z) and a space fixed coordinate system ( $\xi$ ,  $\eta$ ,  $\zeta$ ) on the cluster solvent as shown in Figure 3.1. Both systems have their origins at the nuclear center of mass of the cluster solvent. The molecule fixed coordinate system is chosen such that its principle axes lie alone the  $\xi_2$  rotational axes of the solvent tetrahedcon sl<sub>1</sub> period at the symmetry. The space fixed coordinate system is thereen to be define the tangles  $\theta = 1$  degrees to be degree and  $\eta = 0$  to  $\eta = 0$ .

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# Figure 3-1

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Stereoscopic projection of solvent tetrahedron showing relative orienta time of the molecule fixed is yzi and space fixed if it conditiate is stress or the displace the space force system. Successful the space f(x,y) = f(x,y)

of the rotational coordinates. Since a solvent rotation is periodic over 360 degrees, the potential can be expressed by an even function Fourier series written as $^7$ 

$$V(q) = V_0/2 + \sum_{p=1}^{\infty} A_p \cos(pq)$$
 3.1

with q a function of the rotational coordinates and  $V_0$  taken as the barrier height to internal rotation. The function q is determined by the potential dimensionality and the number of minima in the potential along each of the rotational coordinates. The first cosine term in the series describes the major form of the potential, and the rest of the terms in the series "fine tune" the potential shape. Since the coefficients  $A_p$  in these terms are assumed to be small, the series can be truncated at its first cosine term without sacrificing the general potential shape. Doing this yields the mathematically convenient potential form:

$$\frac{V}{V(q)} \sim (1 - \cos^2(q)) \qquad <\varepsilon_{c}$$

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potential magnitude depends upon the relative displacements of all three rotational coordinates as,

$$q = 2\theta + 2\chi + \phi \qquad 3.3$$

Inserting Equation 3.3 into Equation 3.2 yields the three dimensional potential function used in the calculations:

$$V(\theta, \phi, \chi) = \frac{V_o}{2} \left[1 - \cos(2\theta + 2\chi + \phi)\right] \qquad 3.4$$

in which  $V_0$  is the barrier height to internal rotation. The rotational wavefunctions chosen as the basis set for the calculation are the rigid rotor symmetric top wavefunctions which depend on the curvilinear coordinates and on the quantum numbers J, k and m. Under zero field conditions, these wavefunctions are solutions to the spherical top Schrodinger equation,<sup>8</sup>

<sup>.</sup> 

J'	-	J"‡	≤	2	3.7
k'		k"	=	2	3.8
m '	-	m"	=	1.	3.9

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Diagonal matrix elements contain only the zero field spherical top energies and a  $V_0/2$  potential term.

The resulting matrix is diagonalized for a selected value of  $V_0$  yielding eigenvalues corresponding to the solvent torsional eigenstates at the specified perturbation. The perturbation is varied until a reasonable fit with the experimental spectrum is obtained. The calculations are performed on a Cyber 205 computer using a basis set consisting of 680 wavefunctions to ensure convergence of the lowest eigenstates at their proper eigenvalues. Only the torsional structures of ben(CH<sub>4</sub>)<sub>1</sub> and ben(CD<sub>4</sub>)<sub>1</sub> are calculated since their respective rotational constants are large enough that the coupling of the rotational levels does not require an extremely large basis set to insure convergence. Matrix elements are determined via numerical integration using a nonadaptive integration routine. The matrix is prediagonalized into a tridiagonal form using orthogonal similarity transformations<sup>9</sup> and diagonalized using an implicit QL method <sup>10</sup>. The rotational constants used for methane and deuterated methane are 5.2 cm<sup>-1</sup> and 2.6 cm<sup>-1</sup> respectively.

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Two color TOFMS  $S_1 + S_0$  spectrum and calculated ground state vdW modes (NCA) of ben(CD<sub>1</sub>)<sub>1</sub>. Energy scale is relative to ben(CD<sub>1</sub>)<sub>1</sub>  $S_0^1$  transition (38567.1 m<sup>-1</sup>). Feature positions and assignments as per Table 3.1 and Figure ...

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TAB	

vdW spectral features in  $ben(CD_4)_1$  and  $ben(CH_4)_1$   $6_0^1$  regions

and usiculated ground state volw socies (refer to Figure 3.2). Ben(CH4) 1 d

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*	J.D HARRA C	Assignment b	Observed	NCA	3D-HRRA <sup>C</sup>	Assignment
	()(	6 <sup>-</sup>	0(38567.6)	0	0(4)	6.1 6
e •		6 1 h x vo	16.1	16(b <sub>xy</sub> )		60bxyo
:	21(4)	6111 60120	27.3	28(t <sub>z</sub> )	29(4)	6 <sup>1</sup> t 1 6 <sup>0</sup> t 20
-		6 0 x v 0	32.3			60 <sup>1</sup> bxy0
		•	48.4			6 <sup>1</sup> bxyo
	42(4)	6 <sup>1</sup> t 2 60 20	51.4		54(2)	6 <sup>1</sup> t 2 6 <sup>0</sup> t 20
.:	68(8)	6 1 t 1 6 0 t x y 0				•
		A <sup>1</sup> bzyo	84 . 8			6 <sup>1</sup> bxyo
	75(4)	61t 3 60t 20	73.5		73(4)	6 <sup>1</sup> t <sup>3</sup> 60 20
:		60°20		82(=,)		
	86(12)	81 t xyo 20				
				89(t <sub>xy</sub> )	84(8)	
		6 <sup>1</sup> s 20 b x y 0				
		6]#11]				
		61t 4 60t 20				

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.... , parentheses represent "structural" degeneracy calculated in 3D-HKRA.

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difference between  $S_1$  and  $S_0$  for  $ben(CD_4)_1$  is nearly identical to that of  $ben(CH_4)_1$ . Ten pronounced intermolecular vibronic features are observed to the "blue" of the cluster  $6_0^1$ . As in the case of the  $ben(CH_4)_1$  cluster, no features are observed in the symmetry forbidden benzene  $0_0^0$  region. Thus, the cluster geometry must possess at least a three-fold axis of symmetry.

The calculated  $ben(CD_4)_1$  geometry (Figure 3.3) is assumed to be the same as that calculated for  $ben(CH_4)_1$ . The geometry possesses  $C_{3v}$ point group symmetry. In this geometry, the  $CD_4$  center-of-mass lies 3.47 Å above the benzene molecular plane along the three-fold rotational axis. The ground state binding energy in this configuration is calculated at 540 cm<sup>-1</sup>. Using the cluster "red shift" of 41.2 cm<sup>-1</sup>, the excited state binding energy is calculated to be 581 cm<sup>-1</sup>.

The NCA reveals six vdW vibrations (Figure 3.3 and Table 3.1), two being two-fold degenerate. The ground state vibrational energies are 75 cm<sup>-1</sup> for the vdW stretch  $s_z(a_1)$ , 15 cm<sup>-1</sup> for the bends  $b_{xy}(e)$ . and 20 cm<sup>-1</sup> ( $t_z(a_2)$ ) and 64 cm<sup>-1</sup> ( $t_{xy}(e)$ ) for the vdW torsions. The eigenvector normal modes (Figure 3.3) transform in an identical fashion to those calculated for ben(CH<sub>4</sub>)<sub>1</sub>. The vdW stretching mode transforms as the translation of the cluster constituents away from one another along the z (three fold) axis. The vdW bends transform is some combiblation of cluster constituent translations is opposite fore to a perpendicipation to the three fold axis is the vdW bends transform is some combiblation of cluster constituent translations is opposite fore to a perpendicipation to the three fold axis is the vdW bends transform is some combiblation of cluster constituent translations is opposite fore to a perpendicipation to the three fold axis is a stream of the vdW bends transform is some combiblation of cluster constituent translations is opposite fore to a perpendicipation of the three fold axis is a stream of the vdW bends transform is some combiblation of cluster constituent translations is opposite fore to a perpendicipation of the three fold axis is a stream of the vdW bends transform is some combiblation of cluster constituent translations is opposite fore to a perpendicipation of the three fold axis is a stream of the vdW bends transform is a stream of the vdW bends is a stream of the three fold axis is a stream of the vdW bends transform is a stream of the vdW bends is a stream of the three fold axis is a stream of the vdW bends transform is a stream of the vdW bends is a stream of the vdW bends transform is a stream of the vdW bends transform is a stream of the vdW bends is a stream of the vdW bends transform is a stream of the vdW bends transform is a stream of the vdW bends is a stream of the vdW bends transform is a stream of the vdW bends transform is a stream of t



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The 3D HRRA results are also included in Table 3.1 for the lowest eigenstates in the torsional manifold. The torsional mode structure is calculated for  $V_0 = 300 \text{ cm}^{-1}$  and  $B = 2.6 \text{ cm}^{-1}$ . This perturbation results in a reasonable fit for the torsional features observed experimentally. The torsional "zero point energy" is 75 cm<sup>-1</sup>. Two distinct torsional manifolds result from the calculations: one manifold has eigenvalues grouped quartically (nearly four-fold degenerate located at approximately 21 cm<sup>-1</sup>, 46 cm<sup>-1</sup>, and 75 cm<sup>-1</sup> above the zero point energy) and the other manifold has eigenvalues grouped octally (nearly eight-fold degenerate located at approximately contained at approximately 20 cm<sup>-1</sup>.

For comparison, the observed vibronic features for  $ben(CH_4)_1$ are reproduced in Table 3.1 along with the results of the NCA and the 3D-HRRA. The NCA results for  $ben(CH_4)_1$  are those reported previously.<sup>3</sup> The 3D-HRRA torsional structure is calculated for  $V_0 = 300 \text{ cm}^{-1}$  and  $B = 5.2 \text{ cm}^{-1}$ . This perturbation is chosen since it is assumed that the barrier to internal rotation is nearly identical in the two systems as they only differ by isotopic substitution and have the same electronic structure. Additionally, using the same potential barrier of electronic origin for both  $CH_4$  and  $CD_4$  clusters provides a check on the calidity of the model in predicting the torsional mode structure of the dusters. For bench  $H_1^{+1}$  the torsional zero point energy is 96 cm<sup>-1</sup>.





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B Ben( $CF_4$ )<sub>1</sub>

Figure 3.4 and Table 3.2 present the  $ben(CF_4)_1$  cluster spectrum recorded using 2-color TOPMS in the region between 38578.6  ${
m cm}^{-1}$ and 38702.6 cm<sup>-1</sup>. Unlike the ben $(CD_4)_1$  and ben $(CH_4)_1$  cluster  $6_0^1$  transitions, the ben(CF<sub>4</sub>)<sub>1</sub> cluster  $6_0^1$  is blue shifted by 6.1 cm<sup>-1</sup> with respect to the benzene  $6_0^1$ . The small hypsochromic shift indicates that the binding energies in the  $S_0$  and  $S_1$  states are nearly identical with the ground state binding energy being slightly greater. The relative displacement between the two potential surfaces is also small since only 5 vdW transitions are observed and their intensities decrease abruptly at about 50 cm<sup>-1</sup> above the cluster  $6_0^1$  origin. No cluster spectrum is observed in the forbidden benzene  $0^0_0$  region indicating that the cluster possesses at least a three-fold rotation axis. This result is not unexpected since the same observation is made for the  $ben(CH_d)_1$  and ben $(CD_A)_1$  systems. The ground state configuration and vdW eigenvalues/ eigenvectors are shown in Figure 3.5 and Table 3.2. Only one minimum energy geometry is calculated for the cluster. The geometry has  $C_{3w}$ point group symmetry with the  $ext{CF}_4$  center-of-mass at 3.43 Å above the benzene molecular plane along the three-fold axis. The cluster ground state binding energy is 1064 cm<sup>-1</sup>.

The NCA reveals six vdW vibrations. Their ground state vibrational energies are 69 cm<sup>-1</sup> for the vdW stretch  $s_z(a_1)$ , 11 cm<sup>-1</sup> for the vdW bends  $b_{xy}(e)$ , and 13 cm<sup>-1</sup> ( $t_z(a_2)$ ) and 36 cm<sup>-1</sup> ( $t_{xy}(e)$ ) for the vdW torsions. The eigenvector normal modes transform as the translational and rotational representations of the  $C_{3v}$  point group as indicated and in the same manner as those for ben(CH<sub>4</sub>)<sub>1</sub> and ben(CD<sub>4</sub>)<sub>1</sub>.



# Figure 3.4

Two-color TOFMS  $S_1 \leftarrow S_0$  spectrum and calculated ground state vdW modes (NCA) of ben(CF<sub>4</sub>)<sub>1</sub>. Energy scale is relative to ben(CF<sub>4</sub>)<sub>1</sub>  $6_0^1$  transition (38614.7 cm<sup>-1</sup>). Feature positions and assignments as per Table 3.2 and Figure 3.5.

TABLE 🤇	3.2	2
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vdW spectral features in ben $(CF_4)_1 = 6_0^1$  region and calculated ground state vdW modes (refer to Figure 3.4).

Observed <sup>a</sup>	NCA a	Assignment b
0(38614.7)	0	6 <mark>1</mark>
15.7	11 (b <sub>xy</sub> )	$6^1_{o}b^{1}_{xyo}$
17.4	13 (t <sub>z</sub> )	$6^{1}_{o}t^{1}_{zo}$
32.3		6 <sup>1</sup> <sub>0</sub> b <sup>2</sup> <sub>xyo</sub>
35.4		$6^1_o t_{zo}^2$
39.5	36 (t <sub>xy</sub> )	$6^{1}_{0}t^{1}_{xy0}$
	69 (s <sub>z</sub> )	

a) Energies are reported in  $cm^{-1}$  relative to the  $6_0^1$  cluster origin.

b) vdW mode representations as per Figure 3.4.



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# Figure 3.5

Calculated ground state minimum energy configuration (a) and NCA eigenvalue/eigenvector normal modes (b) - (g) for  $ben(CF_4)_1$ . Cluster symmetry is  $C_{3v}$  with an equilibrium intermolecular distance of 3.43 Å. Eigenvectors are normalized and displayed at 2x magnification (2 Å total displacement). Discussion.

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In comparing the calculated ground state vdW vibrational structure and experimental vdW vibronic structure, we assume that the intermolecular potential surfaces of the clusters studied are identical inboth the  $S_1$  and  $S_0$  electronic states. Additionally, we utilize the group theoretical arguments developed in our previous publication<sup>3</sup> on vdW cluster vibronic structure to assign and understand the observed cluster spectra.

A.  $Ben(CD_4)_1$ .

The calculated ground state vdW vibrations (NCA) of  $hen(CD_4)_1$ and the experimental vibronic spectrum is shown in Figure 3.2 and Table 3.1. As previously, the assignments are made by direct comparison between the calculations and the experimental vibronic spectra. The intense feature at 22.5  $cm^{-1}$  to the blue of the cluster  $6^1_0$  corresponds to the  $t_z$  torsion calculated at 20 cm<sup>-1</sup>. Thus this feature and its observed overtones at 48.4 cm<sup>-1</sup>, 70.5 cm<sup>-1</sup>, and 100 cm<sup>-1</sup> are assigned to  $6_0^1$  vdW combination bands  $6_0^1 t_z(a_2)_0^1$ ,  $6_0^1 t_z(a_2)_0^2$ ,  $6_0^1 t_z(a_2)_0^3$ , and  $6_0^1 t_z(a_2)_0^4$ The features at 32 cm<sup>-1</sup> and 62.7 cm<sup>-1</sup> to the blue of the cluster  $6_0^1$ correspond to the first and third overtones of the two-fold degenerate vdW bending modes whose fundamental is calculated at 15 cm<sup>-1</sup>. They are assigned to the  $6_0^1$  combination bands  $6_0^1 b_{xy}(e)_0^2$ , and  $6_0^1 b_{xy}(e)_0^4$ , respec tively. The feature at 58.1 cm<sup>-1</sup> to the blue of the cluster  $6_0^1$  is assigned to a two-fold degenerate  $t_{xy}$  vdW torsions/cluster  $6_0^1$  combination band,  $6_0^1 t_{xy}(e)_0^1$ . The  $t_{xy}$  torsion fundamental is calculated at 64 cm $^{-1}$ . With this assignment and that of the  ${
m t}_{
m z}$  torsion, the feature at 84 cm<sup>-1</sup> is assigned to the  $6_0^1 t_{XY}(e)_0^1 t_Z(a_2)_0^1$  combination band. The vdW stretch is calculated at 75  ${
m cm}^{-1}$ . This mode is identified in the

cluster vibronic spectrum at 72.9 cm<sup>-1</sup> to the blue of the cluster  $6_0^1$  origin and is thus assigned as  $6_0^1 s_z(a_1)_0^1$ . Under this scheme, we assign the features at 93.3 cm<sup>-1</sup> and 97.5 cm<sup>-1</sup> as  $6_0^1 s_z(a_1)_0^1 b_{xy}(e)_0^1$  and  $6_0^1 s_z(a_1)_0^1 t_z(a_2)_0^1$  combination bands

The torsional structure resulting from the 3D-HRRA for  $ben(CD_4)_{1}$ . (Table 3.1), confirms the torsional assignments made using the NCA. The quartically grouped torsional levels at 21 cm<sup>-1</sup>, 46 cm<sup>-1</sup> and 75 cm<sup>-1</sup> correspond to those associated with the t<sub>z</sub> torsion and its overtones in the cluster spectrum at 22.5 cm<sup>-1</sup>, 48.4 cm<sup>-1</sup> and 70.5 cm<sup>-1</sup>. The octally grouped torsional levels at 68 cm<sup>-1</sup> correspond to the t<sub>xy</sub> torsion assigned at 58.1 cm<sup>-1</sup>.

The correspondence between these levels and those calculated using the NCA can be understood as follows. In the 3D-HRRA, four symmetrically equivalent minima exist in the torsional potential surface. These minima correspond to the four ways of placing the solvent tetrahedron upon the solute with a tetrahedral face toward the solute molecular plane. If the barrier between these minima is infinitely high, penetration of the local wavefunctions through the barrier separating the potential minima does not take place. In this case, each "well" contains eigenstates corresponding to intermolecular torsions which occur with small amplitude about each potential well minimum. Since the four "potential wells" are identical in shape and depth, a four-fold "structural" degeneracy exists in which all four potential wells contain identical torsional structure. Thus, for example, a non degenerate torsional eigenstate actually has a four fold structural Considering the actual situation in which the potential barrier is finite, tunneling occurs and the "structural" degeneracy is lifted via interaction of the local wavefunctions through the potential barrier. The eigenstate splitting due to this tunneling may, or may not, be observed depending on the experimental resolution and the relative difference between the eigenstate energy and the barrier height. Splitting of the "structural" degeneracy in the lower portion of the potential well is minimal unless the barrier is low. Based on the calculations, energy level splittings of the first few sets of eigenstates should not be observed unless the barrier is below 150 cm<sup>-1</sup>.

At moderate barriers (about 300 cm<sup>-1</sup>), the eigenstates in the lower portion of the well are nearly degenerate and behave more or less harmonically. For all practical purposes, we can assume these levels to be degenerate. The torsional level structure can then be determined from the eigenstates in one of the minima. Thus, the 3D-HRRA torsional structure calculation simplifies into the NCA. Physically, the cluster can be considered at least "semi-rigid" in the respect that it has a definable equilibrium configuration. The potential energy barrier separating one minimum from the others in the potential surface is large and may be of the order of the cluster binding energy.

The vibronic structure in both the  $ben(CH_4)_1$  and  $ben(CD_4)_1$ spectra suggests that the systems are more or less rigid. None of the observed vibronic state energies follow a free rotor formalism for which the energy level structure is described by Equation 3.2. If the systems behaved nonrigidly, the free rotor eigenstates would lie at approxi mately 2.6J(J+1) cm<sup>-1</sup> and 5.2J(J+1) cm<sup>-1</sup> for  $ben(CD_4)_1$  and  $ben(CH_4)_1$ . respectively, and a AJ - '1 selection rule would govern the transitions

In the ben(CH<sub>4</sub>)<sub>1</sub> and ben(CD<sub>4</sub>)<sub>1</sub> systems, the t<sub>z</sub> torsional mode shifts by 17.6 % upon deuterium substitution of the cluster solvent. Both the NCA and 3D-HRRA models predict a 28.6% frequency shift for t<sub>z</sub>, while the free rotor model would predict a 50% frequency shift for t<sub>z</sub> upon deuteration. Considering that only one mode is taken into account and the mode couplings may be different in the protonated and deuterated clusters, we conclude that torsional tunneling does not take place in the lower levels of the ben(CH<sub>4</sub>)<sub>1</sub> and ben(CD<sub>4</sub>)<sub>1</sub> cluster potential wells and that the clusters can be considered to be rigid.

The negligible isotopic red shift in the  $b_{xy}$  bends also suggests that the cluster is rigid and that the entire observed spectrum is not solely due to internal rotation. The same rationale holds for the vdW stretch: its observation also dispels the notion that only rotor modes occur in the spectrum. No experimental isotopic shift can be determined for the stretching mode, however, since it is not observed in the ben(CH<sub>4</sub>)<sub>1</sub> system. Theoretically, the mode should red shift by 8.5% upon deuteration of the cluster solvent. The ben(CH<sub>4</sub>)<sub>1</sub> stretch should then be at about 80 cm<sup>-1</sup> in the vibronic spectrum based upon the observed stretch in the ben(CD<sub>4</sub>)<sub>1</sub> system. (The ben(CH<sub>4</sub>)<sub>1</sub> stretch is calculated by the NCA to be at 82 cm<sup>-1</sup>).

The ben $(CD_4)_1$  cluster spectrum is also substantially richer than the ben $(CH_4)_1$  spectrum in the respect that both the vdW stretch and  $t_{xy}$ torsions are observed along with well developed  $t_z$  torsion and bend progressions. This is probably due to more favorable Franck-Condon factors in the ben $(CD_4)_1$  case resulting from the isotopic substitution.

As is the case for  $ben(CH_4)_1$ , H-T coupling influences the vibronic intensities (selection rules) in the  $ben(CD_4)_1$  spectrum. The

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nontotally symmetric modes  $t_z$ ,  $t_{xy}$  and  $b_{xy}$  all appear in the spectrum with  $\Delta v = 0, \pm 1, \pm 2, \pm 3, \dots$  selection rules as can be seen in Figure 3.2 and Table 3.1.

B.  $Ben(CF_1)_1$ .

Comparison of the calculated ground state vdW vibrations (NCA) of ben(CF<sub>4</sub>)<sub>1</sub> and the experimental cluster spectrum is presented in Figure 3.4 and Table 3.2. The intense feature at 17.4 cm<sup>-1</sup> to the blue of the ben(CF<sub>4</sub>)<sub>1</sub> 6<sup>1</sup><sub>0</sub> corresponds to the t<sub>z</sub> torsion calculated at 13 cm<sup>-1</sup>. Thus, this feature and its first overtone at 35.4 cm<sup>-1</sup> are assigned to the 6<sup>1</sup><sub>0</sub> vdW combination bands 6<sup>1</sup><sub>0</sub> t<sub>z</sub>(a<sub>2</sub>)<sup>1</sup><sub>0</sub> and 6<sup>1</sup><sub>0</sub>t<sub>z</sub>(a<sub>2</sub>)<sup>2</sup><sub>0</sub>. The shoulder feature at 15.7 cm<sup>-1</sup> to the blue of the cluster 6<sup>1</sup><sub>0</sub> corresponds to the vdW two-fold degenerate b<sub>xy</sub> bend fundamental calculated at 11 cm<sup>-1</sup>. Taking this feature as the combination band 6<sup>1</sup><sub>0</sub>b<sub>xy</sub>(e)<sup>2</sup><sub>0</sub> at 32.3 cm<sup>-1</sup>. Finally, the spectral feature at 39.5 cm<sup>-1</sup> in the ben(CF<sub>4</sub>)<sub>1</sub> spectrum corresponds to the t<sub>xy</sub> torsion calculated at 36 cm<sup>-1</sup>. Thus, the feature is assigned to the combination band 6<sup>1</sup><sub>0</sub>t<sub>xy</sub>(e)<sup>1</sup><sub>0</sub>.

The ben(CF<sub>4</sub>)<sub>1</sub> spectrum, like the ben(CD<sub>4</sub>)<sub>1</sub> and ben(CH<sub>4</sub>)<sub>1</sub> spectra, suggests that the system is rigid. Since the rotational constant for CF<sub>4</sub> is small (ca. 0.18 cm<sup>-1</sup>), the free rotor energy level structure should appear at about .37 cm<sup>-1</sup> intervals. This structure is not observed. Instead, the spectrum possesses oscillatory torsional structure commensurate with the NCA theoretical predictions. These theoretical and experimental results demonstrate the rigidity of the systems.

The  $S_1 \leftarrow S_0$  excitation of the ben $(CF_4)_1$  cluster involves very little change in cluster geometry. In this spectrum, the progression intensities decrease dramatically at energies greater than the  $6_0^1$  cluster origin. In the lowest two vdW mode progressions, the intensities decrease approximately 13% between the mode fundamentals and the first overtones. Furthermore, the high energy vdW stretch calculated at 69 cm<sup>-1</sup> is not observed. At these high energies, the Franck-Condon factors must be very small.

The observation of the nontotally symmetric vdW  $b_{xy}$  bends and the  $t_z$  torsion progressions with a  $\Delta v = 0, \pm 1, \pm 2$  ... selection rule suggests that interelectronic state mixing (H-T coupling) is an important contributor to the mode intensity mechanism. As in the ben(CH<sub>4</sub>)<sub>1</sub> and ben(CD<sub>4</sub>)<sub>1</sub> cases, H-T coupling becomes apparent in the low lying vdW modes of the ben(CF<sub>4</sub>)<sub>1</sub> system. In fact, the interelectronic state mixing is substantial in the ben(CF<sub>4</sub>)<sub>1</sub> system and can be demonstrated by the observation of the well defined bend fundamental at 15.7 cm<sup>-1</sup> and by the observation of the t<sub>xy</sub> torsion fundamental (39.5 cm<sup>-1</sup>). Both of these latter features should not be observed if H-T coupling is not present.

Summary and Conclusions.

Two-color TOFMS and supersonic molecular jet techniques have been employed to study the  $S_1 \leftarrow S_0$  vibronic spectra of ben $(CD_4)_1$  and ben $(CF_4)_1$  vdW clusters. These studies reveal detailed information regarding the geometries, the intermolecular energetics, and the physical nature of the vdW interactions present in the systems. The experimental observations demonstrate that the clusters are at least semi-rigid systems possessing unique equilibrium geometries and that the intermole cular motion present in the systems is oscillatory. Through comparison of these spectra with those of ben $(CH_4)_1$  previously studied in this laboratory: the "isotopic" shifts observed in the intermolecular vibronic structures demonstrate that all the low lying intermole claseigenstates are nearly harmonic. They are not admixtures of a culater intermolecular bends and stretches and free intermolecular rotation as would occur if the clusters behaved nonrigidly.

The rigidity of the clusters is further demonstrated by theoretical treatment of the intermolecular motion. Both the NCA and 3D HRRA models indicate that the intermolecular torsional motion is oscillators and that the motion is constrained by an orientationally dependent intermolecular potential. Detailed analysis of the intermolecular interaction reveals that the torsional motion is governed by a three dimensional potential possessing torsional barriers on the order of the cluster binding energy. At this barrier magnitude, torsional tunneling in the lower portions of the potential is minimal and the residence time of the cluster constituents in a particular well minimum is substan tially longer than the timescale of experimental observation.

The assignment of geometry and identification of the intermole cular modes using the theoretical models has also proven useful in understanding the physics governing the spectroscopic properties of the clusters. The observation of nontotally symmetric intermolecular mode fundamentals and combination bands in the vibronic spectra suggest that interelectronic state mixing (H-T coupling) is an important factor in the overall intensity mechanism governing the transitions. Addition ally, the interelectronic state mixing is most important for the lowlying intermolecular bending and torsional modes.

Both the NCA and the 3D-HRRA adequately model the intermolecular modes in the systems studied; however, the NCA is the more useful here te no julio te

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We wish to thank Professor W. Klemperer for many helpful discussions and suggestions concerning the rigidity of vdW molecules.

### REFERENCES

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- M. Schauer and E.R. Bernstein, J. Chem. Phys. <u>82</u>, 726 (1985); M. Schauer, K.S. Law and E.R. Bernstein, J. Chem. Phys. <u>82</u>, 736 (1985); J. Wanna and E.R. Bernstein, J. Chem. Phys. <u>84</u>, 927 (1986).
- 2. T.A. Stephenson and S.A. Rice, J. Chem. Phys. 81, 1083 (1984).
- 3. J.A. Menapace and E.R. Bernstein, J. Phys. Chem. <u>90</u>, 000 (1987) and references therein.
- J.A. Beswick and J. Jortner, J. Chem. Phys. <u>74</u>, 6725 (1981) and references therein; D.F. Kelley and E.R. Bernstein, J. Phys. Chem. (to be published).
- 5. D.F. Kelley and E.R. Bernstein, J. Phys. Chem. <u>90</u>, 5461 (1986).
- 6. T.M. Miller and B. Bederson, Adv. At. Mol. Phys. <u>13</u>, 1 (1977); "Handbook of Chemistry and Physics" 53rd Ed. (CRC Press, 1973); G. Nemethy, M.S. Pottle and H.A. Scheraga, J. Phys. Chem. <u>87</u>, 1883 (1983) and references therein.
- B.L. Crawford Jr., J. Chem. Phys. <u>8</u>, 273 (1940); K.S. Pitzer and W.D. Gwinn, J. Chem. Phys. <u>10</u>, 428 (1942); G. Herzberg, "Molecular Spectra and Molecular Structure: II. Infrared Raman Vibrational Spectra", (McGraw-Hill Book Co., Inc. 1955).
- 8. P.R. Bunker, "Molecular Symmetry and Spectroscopy", (Academic Press, 1979).
- R.S. Martin, C. Reinsch and J.H. Wilkinson, Num. Math. <u>11</u>, 181 (1968).
- R.S. Martin and J.H. Wilkinson. Num. Math. <u>12</u>, 377 (1968); A. Dubrille, Num. Math. <u>15</u>, 450 (1970).
- P.J. Breen, J.A. Warren, E.R. Bernstein and J.I. Seeman, J. Chem. Phys. (to be published).

### CHAPTER FOUR

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HYDROGEN BONDED AND NON-HYDROGEN BONDED VAN DER WAALS CLUSTERS: COMPARISON BETWEEN CLUSTERS OF PYRAZINE. PYRIMIDINE. AND BENZENE WITH VARIOUS SOLVENTS (Reprint contained in Appendix One) CHAPTER FIVE

SUPERSONIC MOLECULAR JET STUDIES OF THE PYRAZINE AND PYRIMIDINE DIMERS

(Reprint contained in Appendix Two)

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### CHAPTER SIX

### SUPERSONIC MOLECULAR JET STUDIES OF PHTHALOCYANINES AND THEIR VAN DER WAALS CLUSTERS WITH SMALL MOLECULES

Introduction.

Supersonic molecular jet spectroscopy can be utilized to study weakly bound solute/solvent van der Waals (vdW) clusters in the gas phase.<sup>1</sup> These investigations have increased our understanding of the intra- and intermolecular energetics and dynamics present in the solute/ solvent systems as well as the nucleation and growth of small clusters.<sup>2-11</sup> The vdW clusters studied in our laboratory thus far center around aromatic hydrocarbon and N-heterocycle solutes clustered with small hydrocarbon and hydrogen bonding solvents. We are currently expanding our studies of solute/solvent systems to include clusters of small molecules with macrocycles, such as free-base phthalocyanine (H<sub>2</sub>Pc) and magnesium phthalocyanine (MgPc).

Considerable interest exists in phthalocyanine (Pc) compounds as coloring agents, photosensensitizers, organic semiconductors, and as model systems for biologically important species, such as porphyrins. Pc's have exceptional color, chemical, temperature, and solubility characteristics which make them attractive for use as coloring agents in commercial dyeing and painting processes.<sup>12</sup> The unique electrical pro perties of Pc's serve as building blocks in a number of important technical applications involving electronic devices. For example, thin film sandwich photovoltaic cells containing Pc's have been fabricated which possess photocurrent quantum efficiencies of as much as 10%.<sup>13</sup> Pc films have also been shown to possess both n- and p- type semiconductor properties.<sup>14</sup>

Of particular interest are the interactions between Pc's and other molecular species. Pc films have been observed to respond electrically to the presence of vapors such as  $BF_3$ ,  $NO_2$ , and  $NH_3$ .<sup>15</sup> Interaction between the film surface and the vapors produces reversible and reproducible film conductivity changes which make the measuring of molecular concentrations of these species at the ppm to ppb levels possible. Pc's and porphyrins have also been studied as photocatalysts for the reduction of water to hydrogen.<sup>16</sup> Photochemical generation of hydrogen from water is a very attractive goal for photochemical storage of solar energy. Since water does not absorb at solar wavelengths present at the earth's surface, the photoreductive reactions must be sensitized by dyes. Pc's are good candidates as dyes in this respect since they absorb in the visible spectrum where peak solar radiation occurs.<sup>17</sup> The elucidation of the trapping and transduction mechanisms of solar energy into useful chemical energy is of particular importance in these systems as it may yield insight into the understanding of the basic process of photosynthesis in biologically active systems such as green plants and bacteria.<sup>18</sup> Studying the interactions between Pc's and small "solvents" may also prove useful in understanding the transport and storage of small molecules and the transfer of electrons in a variety of biologically active species such as hemoglobin, myoglobin, and cytochromes.<sup>19</sup>

Supersonic molecular jet investigations can contribute to the resolution of a number of important concerns dealing with the behavior of Pc/solvent systems on the microscopic scale. Questions that can be considered are (1) what are the ground and excited state interaction energies between Pc and various solvents, (2) what are the preferential intermolecular interaction sites on the Pc moiety, (3) what are the most favorable cluster geometries, (4) what types of interactions are important in the intermolecular potential established between the solute and solvent, and (5) are changes in the chromophore symmetry/geometry induced by clustering. The answers to these questions will certainly be helpful to those employing Pc's or related systems in reactions and other chemical processes.

In this paper, we report the spectroscopic results of  $H_2Pc$  and MgPc clustered with hydrocarbon solvents ( $CH_4$ ,  $C_2H_6$ , and  $C_3H_8$ ), hydrogen bonding solvents ( $H_2O$ , MeOH, EtOH), and  $CO_2$  along with the calculated modeling of selected cluster characteristics. The spectroscopic results include the  $\pi^* \leftarrow \pi$  fluorescence excitation (FE) spectra of the clusters near the Pc  $0_0^0$  transition. The calculated results include the ground state geometry and the binding energy for each of the cluster systems studied. The H\_Pc cluster spectra and calculated geometries and binding energies are compared with those obtained for the MgPc clusters. Differences and similarities between the H<sub>2</sub>Pc(MgPc) clusters and the aromatic hydrocarbon and N-heterocycle clusters are discussed. Specifi cally, trends in spectral shifts, relative solute/solvent orientations. and binding energies are noted. Finally, the results of an out-of-plane normal coordinate analysis for Pc are discussed and compared to both Pc and cluster spectra within the first few hundred wavenumbers of the origin transition.

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Experimental Procedures.

The majority of the apparatus used in these experiments has been previously described;  $2^{-7}$  therefore, only a description of the equipment and procedures unique to the present studies will be discussed. A high temperature continuous supersonic nozzle fabricated to generate and analyze the H<sub>2</sub>Pc and MgPc clusters in shown in figure 6.1. The expansion region is constructed by welding a stainless steel pinhole (Micro Engineering) A onto a 1/4 inch tube connector gland (Cajon VCR) B. The pinhole/gland is attached to the nozzle backing region C using a 1/4inch tube connector (Cajon VCR) D. Stainless steel gaskets are used to insure proper connector sealing throughout the nozzle operating temperature range. This setup allows different size pinhole assemblies (nozzle throats) to be attached to the nozzle to obtain various peak experimental backing pressure and gas throughput conditions. A 100 micron pinhole is used to perform the expansion in the H<sub>2</sub>Pc and MgPc cluster experiments. The nozzle backing region is constructed from 1/2 inch stainless steel tubing. The tubing extends through the vacuum chamber wall E passing through a vacuum quick disconnect (MDC Vacuum) F. Connection of the nozzle to the expansion gas line is accomplished using a stainless steel tube connector (Swagelok) G. Solid samples are placed in the nozzle backing region using a quartz "boat" H which can be inserted into the nozzle through the expansion gas line connection. The "boat" facilitates access to the nozzle backing region while the nozzle is in the vacuum chamber and is heated. It allows for easy inspection of the sample during and after an experiment without nozzle removal or nozzle cooling. The nozzle is heated using two heating coils (ARI Industries) I located around the expansion and backing regions. The



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High temperature continuous supersonic molecular jet nozzle (see text).

heaters are capable of maintaining the nozzle at temperature up to 650°C. The heaters are independently controlled using two transformer assemblies. Nozzle temperature is monitored using two iron-constantan thermocouples (Omega) J placed around the expansion and backing regions. The nozzle/heater assembly is enclosed in a stainless steel shield K to minimize heat loss and to maintain uniform nozzle heating.

FE spectra of the  $H_2Pc$ , MgPc, and their respective solvent clusters are obtained using a Nd<sup>+3</sup>/YAG pumped DCM (Exciton) dye laser. Dye laser output is 46-60 mJ/pulse in the vicinity of the  $H_2Pc$  and MgPc  $O_0^0$  transitions. Total excited state fluorescence is collected using a f/1.2 5 cm lens focused at 5-8 mm in front of the nozzle throat and detected by a water cooled RCA C31034 photomultiplier tube. The photomultiplier tube output is amplified 10x using an Ortec 9301 amplifier. The signal is then sent to a boxcar/computer for averaging and digital storage. Fluorescence wavelength calibration is provided by an optogalvanic cell with iron and neon lines as standards.

H<sub>2</sub>Pc (Aldrich) and MgPc (Eastman Kodak) are purified by vacuum sublimation before use. The solid samples are pelletized prior to insertion into the nozzle to minimize consumption. Granular samples placed in the nozzle are consumed at a rate of ca. 45 mg/hr. Pelletization reduces sample consumption by about a factor of thirty without significant loss in fluorescence intensity.

The  $H_2Pc$  and MgPc solid samples are heated to  $380-640^{\circ}$  C to provide sufficient vapor density to perform the spectroscopy. Methane, ethane, propane or carbon dioxide is doped into helium carrier gas at concentrations of up to 1% (partial pressure). The gas is then mixed with the  $H_2Pc$  or MgPc in the nozzle backing region and expanded using

pressures ranging from 100-200 psig. Water, methanol, or ethanol is seeded into the helium carrier gas by passing the carrier gas through an inline trap containing the liquid solvent.

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Calculations of the ground state cluster binding energies and geometries are conducted using an empirical intermolecular potential generated from additive atom-atom potentials set into a Lennard-Jones format. The potential includes general non-bonding (6-12), monopole charge (1), and hydrogen bonding (10-12) terms.<sup>20</sup> The H<sub>2</sub>Pc and MgPc structures used in the calculations are obtained from crystal structure data.<sup>21</sup> The atom-atom potential for magnesium is approximated using experimentally determined polarizabilities<sup>22</sup>, interatomic distance<sup>23</sup>, and the Slater-Kirkwood approximation<sup>24</sup>. Atomic partial charges employed to model the monopole charge interaction are taken from extended Huckel<sup>25</sup> calculations. The hydrocarbon and water structures are those previously used in studying benzene and N-heterocycle clusters.<sup>2,8-10</sup> The CO<sub>2</sub>, MeOH, and EtOH structures are taken from ref. 26, 27, and 28, respectively.

The out-of-plane normal coordinate analysis for  $H_2Pc$  is conducted using the FG matrix method of Wilson et. al.<sup>29</sup> The details of the analysis will be reported elsewhere.<sup>30</sup> Briefly, the nuclear motion is modeled using a set of 82 internal coordinates; 48 C-C(N) bond torsions, 18 C(N)-H bond wags, and 16 C-C(N) bond wags. The valence force field for the <u>P</u> matrix consists of the diagonal force constants describing the out-of-plane ground state motions in benzene.<sup>31</sup> In this gross approximation, all  $H_2Pc$  bond torsions force constants are assumed to be the same as the C-C torsions in benzene. All bond wag force constants are assumed to be the same as the benzene C-H wags. The secular equation

describing the nuclear motion is symmetry factored into four species sets ( $B_{1u}$ ,  $A_u$ ,  $B_{2g}$ , and  $B_{3g}$ ) under the  $D_{2h}$  point group. The factored equations are individually diagonalized to yield 15  $B_{1u}$ . 13  $A_u$ . 13  $B_{2g}$ , and 14  $B_{3g}$  out-of-plane frequency eigenvalues and eigenvector normal modes. (See Table 6.1).

Results.

A. Isolated Ultracold Molecular FE Spectra of H<sub>2</sub>Pc and MgPc.

Figure 6.2 presents the PE spectrum of the  $H_2PC S_1 + S_0$ transition ( $Q_x$  band) in the vicinity of the  $0_0^0$ . The spectrum is taken at 200 psig helium backing pressure ( $P_0$ ), a nozzle expansion region temperature ( $T_e$ ) of 570°C, and a nozzle backing region temperature ( $T_b$ ) of 460°C. The general nature of the spectrum has been previously described.<sup>32</sup> The purpose of its reproduction in this paper is 1) to present a detailed account of the low frequency vibronic transitions observed in the vicinity of the  $0_0^0$  which have not been previously reported in the literature Table 6.2, and 2) to provide an isolated chromophore spectrum which can be compared to the cluster spectra, all of which are taken under nearly identical experimental conditions.

The FE spectrum of the MgPc  $S_1 + S_0$  transition (Q band) in the vicinity of the  $0_0^0$  is shown in Figure 6.3. The spectrum is taken using  $P_0 = 120$  psig helium,  $T_e = 630^{\circ}$ C, and  $T_b = 397^{\circ}$ C. Table 6.3 lists the energies of the vibronic transitions observed in this portion of the spectrum.

B. Hydrocarbon Clusters:  $CH_4$ ,  $C_2H_6$ , and  $C_3H_8$ .

The  $H_2Pc/hydrocarbon$  cluster FE spectra observed in the vicinity of the  $H_2Pc$   $0_0^0$  are presented in Figure 6.4. The energies of

### TABLE 6.1

### Calculated out-of-plane mode frequencies for $\mathrm{H}_{2}\mathrm{Pc}.$

Calculated Mode Energy (cm <sup>-1</sup> )	D <sub>2h</sub> Symmetry Species	Observed Overtone <sup>a</sup> Energy (cm <sup>-1</sup> )	Mode Designation
14.8	B <sub>111</sub>	31.2 (15.6)	A
24.5	B <sub>1u</sub>	51.8 (25.9)	В
84.0	<sup>B</sup> 1u	163.7 (81.9)	с
33.1	A <sub>u</sub>	71.4 (35.7)	D
71.7	Au	141.7 (70.9)	E
38.6	<sup>8</sup> 2g	85.4 (42.7) or 100.6 (50.3)	F
72.1	B <sub>2g</sub>	176 (88) or 203.1 (101.6)	G
38.2	B <sub>3g</sub>	85.4 (42.7) or 100.6 (50.3)	н
71.4	<sup>B</sup> 3g	176 (88) or 203.1 (101.6)	J

a) Values in parentheses are forbidden fundamental mode energies inferred from overtone transitions.



### Figure 6.2

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FE spectrum of  $0_0^0$  region of  $H_2Pc$  taken at  $T_e = 570^{\circ}C$ ,  $T_b = 460^{\circ}C$ , and  $P_0 = 200$  psig He. Peak assignments are given in Table 6.2.

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### TABLE 6.2

Energy (vac. cm <sup>-1</sup> )	Wavelength (vac. Å)	Energy Relative to $0_0^o$ (cm <sup>-1</sup> )	Assignment <sup>a</sup>
15131.8	6608.6	0	000
15163.0	6595.0	31.2	A <sup>2</sup> <sub>o</sub>
15183.6	65 <b>86</b> .1	51.8	8 <mark>2</mark>
15193.9	6581.6	62.0	A <mark>4</mark>
15203.2	6577.6	71.4	$D_0^2$
15215.6	6572.2	83.8	$A_o^2 + B_o^2$
15217.2	6571.5	85.4	$F_0^2$ or $H_0^2$
15225.8	6567.8	93.9	(b)
15228.0	6566.9	96.2	$A_0^1 + C_0^1$
15232.4	6564.9	100.6	$F_0^2$ or $H_0^2$
15258.7	6553.6	126.9	(b)
15273.5	6547.3	141.7	E <sup>2</sup> o
15290.5	6540.0	158.7	$126.9 + A_0^2$
15295.5	6537.9	163.7	$c_o^2$
15307.8	6532.6	176	$G_0^2$ or $J_0^2$

## Observed vibronic transitions in the vicinity of the $\mathrm{H}_{2}\mathrm{Pc}~\mathrm{O}_{0}^{\mathrm{O}}.$

a) Assignments based on normal coordinate analysis results (see Table 6.1).

b) Modes possibly due to in-plane motion.

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Figure 6.3

FE spectrum of  $0_0^0$  region of MgPc taken at  $T_e = 630$  °C,  $T_b = 397$  °C, and  $P_0 = 120$  psig He. Peak assignments are given in Table 6.3.

Energy (vac. cm <sup>-1</sup> )	Wavelength (vac. A)	Energy Relative to $0_0^0$ (cm <sup>-1</sup> )	Assignment <sup>a</sup>
15612.1	6404.5	0	000
15643.0	6392.6	30.9	$A_o^2$
15666.7	6383.0	54.6	B <sub>o</sub> <sup>2</sup>
15673.2	6380.3	61.1	A <sup>4</sup> <sub>0</sub>
15704.4	6367.6	92.3	$F_0^2$ or $H_0^2$
15709.0	6365.8	<b>96</b> .9	(b)
15714.6	6363.5	102.5	$F_0^2$ or $H_0^2$
15742.1	6352.4	130.0	
15750.1	6349.2	138.0	Е <mark>2</mark>
15778.0	6337.9	165.9	
15783.5	6335.7	171.4	
15786 0	6334 7	173 9	(b)

### Observed vibronic transitions in the vicinity of the MgPc $0^0_{0}$

a) Assignments based on normal coordinate analysis results (see Table 6.1).

b) Modes are due to in-plane motion.



### Figure 6.4

PE spectra of  $H_2Pc/C_nH_{2n+2}$  clusters in the vicinity of the  $H_2Pc 0_0^0$ . The  $H_2Pc(CH_4)_1$  spectrum (A) is taken at  $T_e = 540^{\circ}C$ .  $T_h = 430^{\circ}C$ .  $P_0 = 200$  psig He, and .1%  $CH_4$ . The  $H_2Pc(C_2H_6)_1$  spectrum (B) is taken at  $T_e = 540^{\circ}C$ .  $T_b = 430^{\circ}C$ .  $P_0 = 160$  psig He, and .1%  $C_2H_6$ . The  $H_2Pc(C_3H_8)_1$  spectrum (C) is taken at  $T_e = 580^{\circ}C$ .  $T_b = 435^{\circ}C$ .  $P_0 = 160$  psig He. and .1%  $C_3H_8$ . Peak assignments are given in Table 6.4.

the observed cluster transitions are listed in Table 6.4. In these experiments, solvent concentration is varied between .07 and 1% and the backing pressure is varied between 0 and 300 psig. Below .07% solvent concentration, no cluster transitions are observed; above about .3%, the cluster spectra appear broad and featureless to the red of the  $H_2Pc 0_0^0$ . Below 50 psig backing pressure, the spectra become broad; at 100 psig to 300 psig, no new cluster transitions emerge in the region probed.

Sharp spectra of MgPc/hydrocarbon clusters could not be generated. Through the range of .06 to 3% solvent concentration, the spectra vary from no observable cluster transitions to broad cluster bands which extend some 60 cm<sup>-1</sup> to the red of the MgPc origin.

The ground state geometries calculated for the H2Pc/hydrocarbon clusters are shown in Figure 6.5. The cluster binding energies and solute/solvent center-of-mass coordinates are listed in Table 6.5. For  $H_2Pc(C_2H_6)_1$ , a geometry nearly isoenergetic with geometry II is calculated but not shown. The geometry is largely the same as II but has the  $C_2H_6$  long axis rotated by 90° about the H<sub>2</sub>Pc symmetry z axis with respect to geometry II. The geometry has a binding energy of 1564  $cm^{-1}$ . For  $H_2Pc(C_3H_8)_1$ , two additional cluster geometries are calculated and not shown. Geometry III is similar to geometry II in that the solvent  $C_2$  axis lies perpendicular to the H<sub>2</sub>Pc molecular plane; however, the  $C_3H_8$  is inverted in this geometry with respect to geometry I. The cluster binding energy is 1975  $cm^{-1}$ . Geometry IV is nearly isoenergetic with geometry I. The geometry is largely the same as II but has the  $C_{3}H_{8}$  rotated by 90° about the H<sub>2</sub>Pc symmetry z axis. This geometry has a binding energy of 2202  $cm^{-1}$ .

### TABLE 6.4

Species	Energy (vac. cm <sup>-1</sup> )	Cluster $0^{0}_{0}$ Rela- tive to H <sub>2</sub> Pc $0^{0}_{0}$ (cm <sup>-1</sup> )	Energy Relative to Cluster $0_0^0$ (cm <sup>-1</sup> )	Assignment
H <sub>2</sub> Pc(CH <sub>4</sub> ) <sub>1</sub>	15117.2	-14.6	0	000
	15148.5		31.3	$A_0^2$
	15163.5		51.3	$B_0^2$
<sup>н</sup> ₂ <sup>рс(С</sup> 2 <sup>н</sup> 6)1	15105.9	-25.9	0	I 0 <mark>0</mark>
	15121.5		15.6	$A_0^1$
	15138.6		32.7	$A_0^2$
	15113.0	-18.8	0	II 0 <mark>0</mark>
	15144.3		31.3	$A_o^2$
H <sub>2</sub> Pc(C <sub>3</sub> H <sub>6</sub> ) <sub>1</sub>	15098.4	-33.4	0	I 00
	15113.2		14.8	$A_0^1$
	15106.7	-25.1	0	11 0 <mark>0</mark>
	15122.4		15.7	$A_{0}^{1}$
	15138.1		31.4	$A_0^2$

 $\rm H_2Pc/Hydrocarbon$  cluster transitions in the vicinity of  $\rm H_2Pc~0_0^O$  .



solvent center-of-mass coordinates are given in Table 6.5.

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# Calculated cluster binding energies, solvent center-of-ass positions. and solvent orientation specifics.

Cluster	Pig. Ref.	Binding Energy	L. Solvent	ocation o Center-o	f f-Nass e	
		(- •••)	×	2 >	N	Solvent Orientation (Remarks)
H <sub>2</sub> Pc(CH <sub>4</sub> ) <sub>1</sub>	~	1176	.240	.208	3.005	Three solvent hydrogens point toward solute
H2Pc(C2H6)1	ŝ	1360	133	. 383	3.747	([) Lower solvent CH <sub>3</sub> is situated as CH <sub>4</sub> in H <sub>2</sub> Pc(CH <sub>4</sub> ) <sub>1</sub>
	ŝ	1574	.672	.022	3.206	(11). Centermost solvent CM <sub>3</sub> has two hydrogens toward solute.
H2Pc(C3Hg)1	ŝ	2024	203	- 929	3.458	(11). Terminal CM <sub>3</sub> groups on solvent point toward solute. Each CM <sub>3</sub> has two hydrogens pointing down.
	en	2225	.427	408	3.312	(1).
H2Pc(H20)1	60	1617	.042	030	2.550	Solvent hydrogens point toward solute
H <sub>2</sub> Pc(NeOH) <sub>1</sub>	æ	1884	- , 101	.200	2.985	<b>Two solvent</b> CM <sub>3</sub> hydrogens and hydroxyl hydrogens point tomard solute.
H <sub>2</sub> Pc(EtOH) <sub>1</sub>		2055	.592	209.	3.143	(1).
	80	2062	013	<b>900</b> -	3.488	(11). Solvent groups closest to solute are situsted as MeON in M <sub>2</sub> Pc(MeON) <sub>1</sub> .
MgPc(H <sub>2</sub> 0) <sub>1</sub>	0	1553	.492	1.478	3.011	Solvent hydrogens point toward solute.
NgPc (NeOH ) <sub>1</sub>	5	2013	.447	1.602	3.363	Solvent bydrozyl hydrogen points toward solute.
MgPc(EtOH) <sub>1</sub>	•	2954	076	. 937	3.351	
H2Pc(CO2)]	11	1992	.126	110	2.784	
Mg ( CO <sub>2</sub> ) <sub>1</sub>	11	2336	996	1.937	3.142	
a) The solut	e center o	of mass is located	at x - y		for all clu	sters.

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C. H<sub>2</sub>O, MeOH, and EtOH Clusters with H<sub>2</sub>Pc and MgPc.

Figures 6.6 and 6.7 and Table 6.6 present the  $H_2^0$ , MeOH, and EtOH, solute/solvent cluster spectra observed in the vicinity of their respective  $H_2Pc$  and MgPc  $0_0^0$  transitions. The  $H_2Pc$  cluster spectra are observed using  $P_0 = 150$  psig helium,  $T_e = 570^{\circ}C$ , and  $T_b = 435^{\circ}C$ . The MgPc cluster spectra are observed using  $P_0 = 135$  psig helium,  $T_e = 630^{\circ}C$ , and  $T_h = 480^{\circ}C$ .

The ground state geometries calculated for the above mentioned cluster series are shown in Figures 6.8 and 6.9. The cluster binding energies and geometry specifics are listed in Table 6.5. For  $H_2Pc(EtOH)_1$ , a geometry similar to geometry I is calculated but not shown. The cluster binding energy is 2053 cm<sup>-1</sup>. This geometry has the EtOH rotated by 90° about the  $H_2Pc$  symmetry z axis with respect to I.

D.  $CO_2$  Clusters of  $H_2Pc$  and MgPc.

The  $H_2Pc/CO_2$  and  $MgPc/CO_2$  FE spectra observed in the vicinity of the cluster chromophore  $O_0^0$  transitions are shown in Figure 6.10. The cluster transition energies are listed in Table 6.7. In these experiments, both backing pressure and  $CO_2$  concentration are varied. As in the  $H_2Pc/hydrocarbon$  cluster experiments, no new or additional sharp cluster transitions are observed under these conditions. The ground state cluster geometries for  $H_2Pc(CO_2)_1$  and  $MgPc(CO_2)_1$  are shown in Figure 6.11. The cluster binding energies are listed in Table 6.5.

Discussion.

Before analyzing the individual cluster systems in detail, we will discuss the low frequency out-of-plane vibrational motion of isolated H<sub>2</sub>Pc and MgPc. Elucidation of the out-of-plane motion in these molecules is essential to the understanding of the cluster spectra and

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### Figure 6.6

PE spectra of  $H_2Pc/ROH$  clusters in the vicinity of the  $H_2Pc$  0<sup>0</sup><sub>0</sub>. The spectra are obtained using  $T_e = 570$ °C,  $T_b = 435$ °C, and  $P_o = 150$  psig He. Traces (A), (B), and (C) correspond to  $H_2Pc(H_2O)_1$ ,  $H_2Pc(MeOH)_1$ , and  $H_2Pc(EtOH)_1$ , respectively. Peak assignments are given in Table 6.6.





FE spectra of MgPc/ROH clusters in the vicinity of the MgPc  $0_0^0$ . Traces (A), (B), and (C) correspond to MgPc(H<sub>2</sub>0)<sub>1</sub>, MgPc(MeOH)<sub>1</sub>, and MgPc(EtOH)<sub>1</sub>. respectively. The spectra are taken at T<sub>e</sub> = 630<sup>°</sup>C, T<sub>b</sub> = 480<sup>°</sup>C, and P<sub>0</sub> = 135 psig He. Peak assignments are given in Table 6.6. The arrows indicate cluster transitions.

### TABLE 6.6

Species	Energy (vac. cm <sup>-1</sup> )	Cluster $0_0^0$ Relative to to H <sub>2</sub> Pc/MgPc $0_0^0$ (cm <sup>-1</sup> )	Energy Relative to Cluster $0_0^0$ (cm <sup>-1</sup> )	Assignment
H <sub>2</sub> Pc(H <sub>2</sub> 0) <sub>1</sub>	15060.6	-71.2	0	00
	15076.3		15.7	$A_0^1$
	15091.8		31.2	$A_0^2$
	15108.2		47.6	$A_0^3$
	15115.7		55.1	$B_0^2$
H <sub>2</sub> Pc(MeOH) <sub>1</sub>	15049.8	-82.0	0	oo
	15066.1		16.3	$A_{O}^{1}$
	15081.2		31.4	$A_0^2$
H <sub>2</sub> Pc(EtOH) <sub>1</sub>	15040.9	-90.9	0	I 00
	15035.3 <sup>a</sup>		15.8	A <sup>1</sup> <sub>o</sub>
	15051.7 <sup>a</sup>		32.2	$A_o^2$
	15048.5	-83.3	0	00 II
	15064.6		16.1	$A_{O}^{1}$
	15080.2		31.7	$A_0^2$
MgPc(H <sub>2</sub> 0) <sub>1</sub>	15658.7	46.6	0	00
	15672.6		13.9	A <sub>0</sub> <sup>1</sup>
	15683.0			
	15687.8		29.1	$\Lambda_0^2$
	15696.4		37.7	$F_0^1$ or $H_0^1$
	15715.8		57.1	$B_0^2$

### $H_2Pc(H_2O)_1$ , $H_2Pc(MeOH)_1$ , $H_2Pc(EtOH)_1$ , $MgPc(H_2O)_1$ , $MgPc(MeOH)_1$ . and $MgPc(EtOH)_1$ cluster transitions in the vicinities of the chromophore $0_0^0$ .

Species	Energy (vac. cm <sup>-1</sup> )	Cluster $0_0^0$ Relative to to H <sub>2</sub> Pc/MgPc $0_0^0$ (cm <sup>-1</sup> )	Energy Relative to Cluster 0 <sup>0</sup> (cm <sup>-1</sup> )	Assignment
	15660.4	48.3	0	00
	15674.6		14.2	A <sup>1</sup> <sub>o</sub>
	15690.2		29.8	$A_0^2$
	15701.0		40.6	$F_0^1$ or $H_0^1$
	15717.3		56.9	B <sub>0</sub> <sup>2</sup>
MgPc(MeOH) <sub>1</sub>	15658.4	46.3	0	00
	15672.5		14.1	$A_0^1$
	15687.9		29.8	$A_0^2$
	15696.8		38.4	$F_{o}^{1}$ or $H_{o}^{1}$
	15713.7		55.3	$B_0^2$
	15660.4	48.3	0	0 <sup>0</sup> 0
	15674.8		14.4	$A_0^1$
	15690.3		29.9	$\Lambda_0^2$
	15701.2		40.8	$F_0^1$ or $H_0^1$
	15717.8		57.4	$B_0^2$
MgPc(EtOH) <sub>1</sub>	15661.7	49.6	0	0 <sup>0</sup> 0
	15676.3		14.6	$A_{O}^{1}$
	15684.7			
	15700.6		38.9	$\mathbf{F}_{\mathbf{O}}^{1}$ or $\mathbf{H}_{\mathbf{O}}^{1}$
	15714.6		52. <b>9</b>	$B_{0}^{2}$

a) Weak features (Figure 6.6) - assignment must be considered tentative



### Figure 6.8

Calculated minimum energy geometries for  $H_2Pc(H_2O)_1$  (A),  $H_2Pc(MeOH)_1$ (B), and  $H_2Pc(EtOH)_1$  (C). The cluster binding energies and solute/ solvent center-of~mass coordinates are given in Table 6.5.





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### Figure 6.9

Calculated minimum energy for  $MgPc(H_2O)_1$  (A),  $MgPc(MeOH)_1$  (B), and  $MgPc(EtOH)_1$  (C). The cluster binding energies and solute/solvent center-of-mass coordinates are given in Table 6.5.



### Figure 6.10

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FE spectra of  $H_2Pc(CO_2)_1$ , and  $MgPc(CO_2)_1$  in the vicinities of the cluster chromophore  $0_0^{\circ}$  transitions. The  $H_2Pc(CO_2)_1$  spectrum (A) is taken at  $T_e = 500 \circ C$ ,  $T_b = 485 \circ C$ ,  $P_o = 150$  psig He, and .2%  $CO_2$ . The  $MgPc(CO_2)_1$  spectrum (B) is taken at  $T_e = 443 \circ C$ ,  $T_b = 412 \circ C$ ,  $P_o = 150$  psig He and .2%  $CO_2$ . Peak assignments are given in Table 6.7. The arrows indicate cluster transitions.

### TABLE 6.7

Species	Energy (vac. cm <sup>-1</sup> )	Cluster $0_0^0$ Relative to to $H_2 Pc' MgPc$ $0_0^0 (cm^{-1})$	Energy Relative to Cluster 00 (cm <sup>-1</sup> )	Assignment
H <sub>2</sub> Pc(CO <sub>2</sub> ) <sub>1</sub>	15145.8	14.0	0	0 <mark>0</mark>
	15161.2		15.4	A <sup>1</sup> <sub>0</sub>
	15176.5		30.7	A <sub>0</sub> <sup>2</sup>
MgPc(CO <sub>2</sub> ) <sub>1</sub>	15630.0	17.9 <sup>a</sup>	0	00
	15649.9		19.9	A <mark>1</mark>
	15664.6		34.6	A <sup>2</sup> <sub>0</sub>
	15634.7	22.6 <sup>a</sup>	0	00
	15655.8		21.1	A <sup>1</sup> <sub>O</sub>
	15667.0		32 3	1 <mark>2</mark>

# $H_2Pc(CO_2)_1$ and $MgPc(CO_2)_1$ cluster transitions in the vicinity of the chromophore $0_0^0$ .

a) Transitions correspond to same cluster species (see text)





### Figure 6.11

Calculated minimum energy geometries for  $H_2Pc(CO_2)_1$  (A) and  $MgPc(CO_2)_1$ (B). The cluster binding energies and center-of-mass coordinates are given in Table 6.5. in the identification of the number of different clusters of a specific composition observed. The nature of molecular motion associated with each spectral feature is determined by using the results of an out-of-plane normal coordinate analysis for isolated  $H_2Pc$ . Table 6.1 lists the nine lowest energy out-of-plane modes calculated. Given the qualitative nature of the force field used in the analysis, the calculated vibrational energies compare quite well with the fundamental energies inferred from the observed overtone transitions presented in Figure 6.2 and Table 6.2.

Three  $B_{1u}$  fundamentals are calculated at 14.8, 24.5 and 84.0 cm<sup>-1</sup>. In the isolated  $H_2Pc$  spectrum, these vibrations are observed as symmetric overtones at 31.2 ( $A_0^2$ ), 51.8 ( $B_0^2$ ), 62.0 ( $A_0^4$ ), and 163.7 ( $C_0^2$ ) cm<sup>-1</sup>. The symmetry forbidden fundamentals are thus located at 15.6, 25.9 and 81.9 cm<sup>-1</sup> using a harmonic oscillator assumption. Two  $A_u$  modes calculated at 33.1 and 71.7 cm<sup>-1</sup> correspond to the forbidden out-of-plane fundamentals at 35.7 and 70.9 cm<sup>-1</sup>. The two  $B_{2g}$  ( $B_{3g}$ ) modes calculated at 38.6 (38.2) and 72.1 (71.4) cm<sup>-1</sup> correspond to the fundamentals of the observed overtone transitions at 85.4 and 100.6 ( $F_0^2$  or  $H_0^2$ ) cm<sup>-1</sup> and at 176 and 203.1 ( $G_0^2$  or  $J_0^2$ ) cm<sup>-1</sup>. The low energy out-of-plane vibronic transitions in the vicinity of the MgPc origin are similarly assigned (Figure 6.3 and Table 6.3).

Five out of the nine lowest energy out-of-plane vibrations are responsible for the vibronic transitions observed in the first 100 cm<sup>-1</sup> of the  $H_2Pc$  S<sub>1</sub> manifold. These five modes involve large amplitude displacements of the four isoindole groups comprising  $H_2Pc$ . The qualitative forms of the motion can be described by the two sets of
operations indicated in Figure 6.12: (a) the tilt of the isoindole groups around the line A through the pyrrole  $\alpha$  carbons; and (b) the rotation of the isoindole groups around the line B between the inner ring nitrogens and the midpoint between outer benzene carbons. The  $B_{1\mu}$ vibration calculated at 14.8  $cm^{-1}$  corresponds to (a) type motion in which a set of opposite isoindole groups tilts out of the molecular plane in one direction and other set of opposite isoindole groups tilts out of the plane in the opposite direction. The B<sub>11</sub> vibration calculated at 24.5  $\text{cm}^{-1}$  is similar to the 14.8  $\text{cm}^{-1}$  mode; however, all the isoindole groups tilt out of the plane about the A axis (Figure 6.12) in the same direction. The mode form looks similar to the forming of a "bowl" out of the phthalocyano skeleton. The  $A_u$  mode (33.1 cm<sup>-1</sup>) involves type (b) motion of adjacent isoindole groups in one direction and the other two adjacent isoindole groups in the opposite direction. The motion looks like "ruffling" of the molecular skeleton. The B<sub>20</sub> mode  $(38.6 \text{ cm}^{-1})$  corresponds to (a) and (b) type motions of opposite isoindole groups. The vibration form is such that one set of the opposite groups tilts out of the molecular plane about A in different directions. The other set of opposite groups rotates about B in the same direction. Overall, the motion looks similar to the forming of a "chair" out of the phthalocyano molety. The  $B_{3g}$  mode (38.2 cm<sup>-1</sup>) form is similar to that associated with the  $B_{2g}$  mode. The motion is the same in both cases; however, the (a) and (b) motion is exchanged between the two opposite groups in the  $B_{3g}$  mode with respect to the motion in the  $B_{2g}$  mode.

Chromophore out-of-plane fundamental and/or overtone transitions are observed in the vicinity of the cluster origins (vide infra) for all of the H<sub>2</sub>Pc and MgPc systems discussed below. The observation of the



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# Figure 6.12

Operations showing qualitative out-of-plane vibrational motion in

H<sub>2</sub>Pc/MgPc.

cluster chromophore out-of plane fundamentals in the cluster spectra can be rationalized using the results of the above normal coordinate analy sis and group theoretical arguments: clusters have reduced symmetry with respect to the isolated chromophore  $H_2Pc$  or MgPc but nearly identical chromophore molecular vibrations. For example, in the  $H_2Pc(H_20)_1$ , the symmetry is reduced from  $D_{2h}$  to, at most,  $C_{2v}$  (see Figure 6.8). Under this reduced symmetry, the forbidden  $B_{1u}$  fundamental vibrations in isolated  $H_2Pc$  correlate to  $A_1$  vibrations in  $C_{2v}$ . The modes are there fore fully allowed by symmetry and should, in principle, be observed if Frank-Condon factors are favorable.

vdW vibrational mode eigenvectors and eigenvalues have been calculated for a number of different solute/solvent systems, including benzene, pyrazine, etc. with many of the same solvents employed in this work.<sup>10,11</sup> Based on these previous studies, we can estimate that the lowest energy vdW modes for the H<sub>2</sub>Pc and MgPc/solvent systems are ca. 50 cm<sup>-1</sup>. Thus, features between cluster origins and ca. +50 cm<sup>-1</sup> are most likely not vdW vibronic modes of the clusters. Moreover, these same studies demonstrate that high energy vdW modes (> 50 cm<sup>-1</sup>) do not have large intensity due to poor Franck-Condon factors. A fuller discussion of these issues is presented below.

On the basis of the above notions, the cluster spectra are analyzed using four premises. First, the cluster transitions are identified as those features not associated with the isolated H<sub>2</sub>Pc or MgPc molety. Second, the cluster origins are assigned to be the lowest energy cluster transitions observed. Third, the cluster vibronic mani folds associated with each of the origins should, and do, exhibit chromophore out of plane fundamental and overtone transitions commensurate with those observed and/or calculated for isolated  $H_2Pc$  or MgPc. And fourth, due to the similarity between the vdW potential surfaces for the chromophore  $S_0$  and  $S_1$  states and the relatively large energies (see below) of the vdW modes in these systems, little if any vdW vibronic mode intensity should be observed.

A. Hydrocarbon Clusters of H<sub>2</sub>Pc and MgPc.

The H<sub>2</sub>Pc/methane spectrum, Pigure 6.4a and Table 6.4, exhibits a cluster transition at 14.6 cm<sup>-1</sup> to the red of the H<sub>2</sub>Pc 0<sup>0</sup><sub>0</sub>. This transition is assigned to the 0<sup>0</sup><sub>0</sub> of a single H<sub>2</sub>Pc/methane species. As discussed above, features 31.3 and 51.3 cm<sup>-1</sup> to the blue of the cluster 0<sup>0</sup><sub>0</sub> correspond to cluster chromophore out-of-plane symmetric overtones,  $A^2_0$  and  $B^2_0$ . Both transitions are shifted by -14.6 cm<sup>-1</sup> with respect to their corresponding transitions in the isolated H<sub>2</sub>Pc spectrum. The  $A^1_0$  transition is not observed as it may be weak and/or within the linewidth of the H<sub>2</sub>Pc isolated molecule 0<sup>0</sup><sub>0</sub>.

The cluster transitions are most likely due to  $H_2Pc(CH_4)_1$  since no new additional sharp transitions are observed when either the  $CH_4$ concentration or the backing pressure is increased. Increasing the  $CH_4$ concentration and/or backing pressure should yield higher order clusters. From prior experience with other solute/solvent cluster systems,<sup>2-9</sup> higher order clusters typically yield more than one set of cluster transitions. Additive shifts are also observed in most cases resulting from inhomogeneous nucleation processes in which solvent mole cules bind symmetrically to opposite sides of the chromophore molecular plane.

The predominance of 1:1 solute/solvent clusters in the expansion may be rationalized on the basis of large solute/solvent binding energy

versus small solvent dimer binding energy. In the systems presently studied, the solute/solvent binding energies are sufficiently large that when a solvent dimer collides with a solute molecule, the dimer dissipates some of the cluster binding energy via vibrational predissociation. This interactive collision leaves one solvent molecule bound to the solute while the other solvent carries off enough of the cluster energy to stabilize the vdW bond until further collisional cooling can take place. These notions would lead one to conclude that 1:1 solute/ solvent clustering predominates in the other clusters systems analyzed as well.

The existence of a single  $H_2Pc(CH_4)_1$  geometry responsible for the cluster origin in the spectrum is further corroborated by the ground state configuration calculation depicted in Figure 6.5a. In this geometry, the  $CH_4$  cluster subunit is situated above the  $H_2Pc$  plane and is coordinated to the  $\pi$ -cloud of the aromatic ring. The cluster geometry is interesting in the respect that the cluster solvent uniquely lies nearly over the H2Pc core: none of the several potential cluster sites located over each of the H<sub>2</sub>Pc closed ring subunits is apparently a true local minimum. One can envision three distinct cluster sites on the H<sub>2</sub>Pc molety: 1) above the H<sub>2</sub>Pc core (most stable); 2) above one of its four five-membered rings; or 3) above one of its four six-membered rings (least stable). If all three of these sites were physically accessible, three different cluster spectral shifts should be observed corresponding to the three distinct sites. Since the single cluster origin in the spectrum suggests that only one geometry is stable, two out of the three speculated minima are either nonexistent or not sufficiently deep to accommodate bound state geometries. The observed

cluster origin at -14.6 cm<sup>-1</sup> from the  $H_2Pc$   $0_0^0$  probably corresponds to a geometry very similar to the calculated one (Figure 6.5a).

 $H_2Pc$  clustered with ethane, Figure 6.4b and Table 6.4, reveals two cluster origins at 25.9 and 18.8 cm<sup>-1</sup> to the red of the  $H_2Pc$  0<sup>0</sup><sub>0</sub>. Two weak transitions to the blue of the  $H_2Pc$  0<sup>0</sup><sub>0</sub> correspond to cluster chromophore out-of-plane vibrations  $A_0^2$  built upon the two cluster origins. The transition at -10.3 corresponds to the fundamental of the out-of-plane cluster chromophore vibration  $A_0^1$  (15.6 cm<sup>-1</sup>) built upon the origin at -25.9 cm<sup>-1</sup>. The cluster chromophore fundamental built upon the cluster origin at -18.8 cm<sup>-1</sup> is not observed as it may be weak and/or within the linewidth of the  $H_2Pc$  0<sup>0</sup><sub>0</sub>.

These cluster manifolds most likely correspond to  $H_2Pc(C_2H_6)_1$ clusters since, as in the  $H_2Pc(CH_4)_1$  case, as both concentration and backing pressure are varied, the spectrum does not yield any additional sharp transitions. Furthermore, no additive spectral shifts are observed indicative of higher order clusters with ethane subunits situated above and below the  $H_2Pc$  plane. The same arguments used to rationalize the predominance of 1:1 clusters in the  $H_2Pc(CH_4)_1$  case are applicable to this system as well.

The two  $H_2Pc(C_2H_6)_1$  cluster geometries shown in Figure 6.5b support the assignment of two cluster geometries. Both geometries have the  $C_2H_6$  situated over the center of the  $H_2Pc$  core as found for  $H_2Pc(CH_4)_1$ . Geometry I should yield a larger spectral shift than geometry II based upon polarizability arguments previously discussed for single ring cluster systems.<sup>2,8</sup> Briefly, the species with the larger spectral shift is associated with the solute/solvent relative orientation for which the direction of the large solvent polarizability is perpendicular to the solute molecular plane. Using this argument, a geometry similar to I would be associated with the cluster origin 25.9 cm<sup>-1</sup> to the red of the  $H_2PC \ 0_0^0$ . As mentioned in the Results Section, two specific solvent orientations are consistent with the qualitative solute/solvent geometry of calculated  $H_2Pc(C_2H_6)_1$  configuration II (see fig 6.5b). One would not expect a spectroscopic difference between the two geometries as they differ by a 90<sup>0</sup> rotation about the symmetry z axis of  $H_2Pc$ . These two directions as far as the  $H_2Pc$  moiety is concerned should be roughly equivalent in terms of polarizabilities and  $\pi$ -cloud overlap.

The H<sub>2</sub>Pc/propane clusters, Figure 6.4c and Table 6.4, are assigned on the basis of similar arguments presented for the other two hydrocarbon clusters studied. In the spectrum, two H<sub>2</sub>Pc(C<sub>3</sub>H<sub>8</sub>)<sub>1</sub> cluster origins appear at 33.4 and 25.1 cm<sup>-1</sup> to the red of the H<sub>2</sub>Pc 0<sub>0</sub><sup>0</sup>. Two cluster vibronic manifolds to the blue of each origin are assigned to cluster chromophore vibronic fundamentals and first overtones. For the cluster manifold beginning at -33.4 cm<sup>-1</sup>, the A<sub>0</sub><sup>1</sup> occurs at 14.8 cm<sup>-1</sup> to the blue of the cluster 0<sub>0</sub><sup>0</sup>. For the cluster manifold beginning at -25.1 cm<sup>-1</sup>, the A<sub>0</sub><sup>1</sup> and A<sub>0</sub><sup>2</sup> occur at 15.7 and 31.4 cm<sup>-1</sup> to the blue of the cluster 0<sub>0</sub><sup>0</sup>.

Geometries similar to those shown in Figure 6.5c could be associated with the two observed cluster manifolds. Different spectral shifts for the two geometries most likely result from difference in  $\pi$ -cloud solvation. The propane solvent interacts less with the H<sub>2</sub>Pc  $\pi$ -cloud in geometry II than in geometry I since the C<sub>2</sub> axis of the cluster solvent (direction of small polarizability) is perpendicular to the H<sub>2</sub>Pc molecular plane in geometry II but parallel to the plane in

geometry I. A geometry similar to I could thus be responsible for the cluster manifold beginning at  $-33.4 \text{ cm}^{-1}$  and a geometry similar to II could be associated with the manifold beginning at  $-25.1 \text{ cm}^{-1}$ .

The  $H_2Pc(C_2H_6)_1$  and the  $H_2Pc(C_3H_8)_1$  spectra exhibit  $A_0^1$  transitions which are forbidden in isolated H<sub>2</sub>Pc. A similar change in selection rules upon clustering has been reported for the benzene/solvent systems.<sup>2,9-11</sup> In the latter systems, the observations suggest that the presence of the cluster solvent over the solute molecular plane is sufficient to induce the forbidden benzene  $0_0^0$  transition if the threefold rotation axis of the solute is destroyed. Furthermore, the selection rules governing the intermolecular vdW motion follow the reduced symmetry of the cluster systems. This minimum perturbation may also be the driving force which induces the vibronic transitions in the  $H_2Pc/$ hydrocarbon clusters. The perturbation in the present instance may even be large enough to cause the cluster chromophore to adjust its geometry in an attempt to wrap itself around the cluster solvents to establish optimal  $\pi$ -cloud overlap. The observation of the B<sub>10</sub> fundamental at 15  $cm^{-1}$  serves as evidence supporting this notion as molecular displacement along this coordinate appears to yield favorable Franck-Condon factors. The  ${\bf A}^1_0$  transition is also observed in all the other H<sub>2</sub>Pc and MgPc clusters studied as well. These observations would lead one to conclude that the clusters are not planar in the excited electronic state.

In general the  $H_2Pc/hydrocarbon$  cluster series is similar to the benzene and N-heterocycle/hydrocarbon cluster series previously studied.<sup>2,8,10,11</sup> The  $H_2Pc/hydrocarbon$  clusters  $S_1 + S_0$  transitions all exhibit bathychromic shifts with respect to the cluster chromophore

transition. The direction of the shifts is the same as that observed in the single ring cluster systems. These shifts indicate that the binding energies of the clusters become larger in the electronic excited state than in the ground state. Upon excitation, the cluster chromophore  $\pi$ -cloud expands and becomes more diffuse. The  $\pi$ -cloud can thus participate more effectively in intermolecular bonding. The net result is stabilization of the vdW bond and a larger binding energy in the cluster  $S_1$  state relative to the  $S_0$  state.

As in the single ring cluster systems, the H<sub>2</sub>Pc/hydrocarbon cluster spectral shifts and calculated binding energies increase with increasing solvent size and  $\pi$ -cloud overlap. The shifts also depend upon the relative orientation of the cluster solute and solvent. The spectral shifts for the H<sub>2</sub>Pc/hydrocarbon series are about a factor of two smaller than those observed in the single ring aromatic/hydrocarbon systems. The difference in the spectral shift magnitudes can be attributed to smaller changes in the intermolecular interaction resulting from smaller overall  $\pi$ -cloud overlap between the solute and the solvents at the H<sub>2</sub>Pc core site. The conjugated  $\pi$ -electron path in H<sub>2</sub>Pc circles around the core yielding a  $\pi$ -cloud "void" at the central core of the molecule. Thus, cluster solvents situated over the core do not overlap with the solute  $\pi$ -cloud as much as they do in single ring systems. The large size and extensive delocalization of the solute  $\pi$ -cloud may also be contributing factors since they minimize electron density changes at the binding site when an electron is promoted from a  $\pi$ - and a  $\pi^*$ -orbital.

The cluster geometries calculated for the H<sub>2</sub>Pc/hydrocarbon series compare well with the geometries calculated, and in some cases

experimentally verified, for the single ring cluster series as far as the cluster solvent orientations are concerned. The geometries, however, are unique in the respect that both the spectra and the calculations suggest that the favorable binding site is the H<sub>2</sub>Pc core and is not at peripheral ring centers.

The one major difference between the H\_PC/hydrocarbon cluster spectra and the benzene/hydrocarbon cluster spectra is the absence of observable intermolecular vdW mode intensity in the H<sub>2</sub>Pc cluster spectra. The lowest energy vdW motion (bending) is expected to be observed at about 50  $cm^{-1}$  to the blue of the cluster origins based upon calculational modeling of intermolecular mode energetics.<sup>10,11</sup> The vdW stretch is expected to occur at about 100  $cm^{-1}$  above the cluster origin. The absence of  $\Delta v \approx \pm 1$  vdW mode transitions in the H<sub>2</sub>Pc clusters may be due to poor Franck-Condon factors. The large size and extensive delocalization of the H<sub>2</sub>Pc  $\pi$ -cloud yields little change in the electron density at the cluster site when a single electron is promoted from a  $\pi-$  to a  $\pi^{\star}$ -orbital. Thus, one might expect that the intermolecular potential surfaces of the two states are nearly superimposable even though the binding energy of the excited state is slightly different (ca. 1%) than that of the ground state. The net result is the observation of vdW sequence structure ( $\Delta v = 0$ ) giving rise to cluster origins and cluster chromophore vibrations only. These arguments probably hold for all the cluster systems investigated in this study as well.

B. H<sub>2</sub>O, MeOH, and EtOH Clusters of H<sub>2</sub>Pc and MgPc.

 $\rm H_2Pc$  clustered with  $\rm H_2O$ . Figure 6.6a and Table 6.6, yields a spectrum exhibiting a single cluster vibronic manifold with an origin 71.2 cm<sup>-1</sup> to the red of the  $\rm H_2Pc$   $O_0^O$ . The single cluster (chromophore)

vibronic manifold suggests that one  $H_2Pc/H_2O$  species is responsible for the observed spectrum. The spectrum is assigned to  $H_2Pc(H_2O)_1$  based upon the same arguments developed for the  $H_2Pc/hydrocarbon$  clusters. The assignment is supported by the ground state configuration calculations for which a single geometry is obtained. Figure 8a. In this geometry, the  $H_2O$  is situated over the  $H_2Pc$  core. The transitions at 15.7, 31.2, and 47.6 cm<sup>-1</sup> to the blue of the cluster  $O_0^O$  correspond  $H_2Pc$ out-of-plane vibrations  $A_0^1$ ,  $A_0^2$  and  $A_0^3$ . The feature at 55.1 cm<sup>-1</sup> to the blue of the cluster  $O_0^O$  corresponds to the  $B_0^2$  cluster chromophore transition.

Clustering  $H_2Pc$  with MeOH yields a cluster spectrum, Figure 6.6b and Table 6.6, associated with  $H_2Pc(MeOH)_1$  which has an origin 82.0 cm<sup>-1</sup> to the red of the  $H_2Pc$  0<sup>0</sup><sub>0</sub>. The cluster chromophore out-of-plane vibrations  $A_0^1$  and  $A_0^2$  are observed at 16.3 and 31.4 cm<sup>-1</sup> to the blue of the cluster 0<sup>0</sup><sub>0</sub>. The assignment of a single cluster manifold is supported by the single calculated ground state geometry obtained (Figure 6.8b).

 $H_2Pc$  cluster with EtOH, Figure 6.6c and Table 6.6, yields a spectrum exhibiting two cluster (chromophore) vibronic manifolds which can be assigned as due to two  $H_2Pc(EtOH)_1$  species. The first cluster origin is at 90.9 cm<sup>-1</sup> to the red of the  $H_2Pc$  0<sup>0</sup><sub>0</sub> and the second cluster manifold begins at 83.3 cm<sup>-1</sup> to the red of the  $H_2Pc$  0<sup>0</sup><sub>0</sub>. The cluster chromophore vibrations  $A_0^1$  and  $A_0^2$  can be identified in these manifolds at 16.1 and 31.7 cm<sup>-1</sup>.

The two  $H_2Pc(EtOH)_1$  cluster species responsible for the observed spectrum can be associated with geometries similar to those shown in Figure 6.8c. In geometry I the EtOH solvent interacts more with the  $H_2Pc \pi$ -cloud than in geometry II and, therefore, geometry I should

exhibit a larger spectral shift. Thus a geometry similar to 1 could be associated with the cluster whose origin is at  $-90.9 \text{ cm}^{-1}$  and a geometry similar to II could be associated with the cluster whose origin is at  $-83.3 \text{ cm}^{-1}$  with respect to the H<sub>2</sub>Pc  $0_0^0$ .

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The  $H_2Pc/H_2O$ , /MeOH, and /EtOH cluster spectra suggest that the solvent OH group is intimately involved in the intermolecular interaction. The red shifts for all three clusters are similar and larger than those observed in the  $H_2Pc/hydrocarbon$  systems. The MeOH and EtOH cluster shifts are larger than that observed in the  $H_2O$  cluster sugges ting that the shifts are dependent upon the combined effects of the OH group and the hydrophobic portions of the cluster solvents. If, on the other hand, the alkyl groups were pointing toward the solute, one would expect the spectral shifts for MeOH/EtOH clusters to be similar to those observed in the hydrocarbon clusters. In these geometries, the alkyl groups would be the major contributors in the intermolecular interaction.

The large spectral shifts, the interaction of the OH group with  $H_2Pc$ , and the ground state configuration calculations lead one to postulate that hydrogen bonding may be occurring in these cluster systems. Hydrogen bonding can occur to some extent between the solute inner ring nitrogens and/or pyrrole hydrogens and the solvent OH group: the  $H_2Pc$  inner ring nitrogens have large electron density<sup>19,33</sup> which can enhance the hydrogen bonding of solvent OH groups. Since the observed chromophore transition is  $\pi^* \leftarrow \pi$ , a large red shift can be expected.

The MgPc(H<sub>2</sub>O)<sub>1</sub>, (MeOH)<sub>1</sub>, (EtOH)<sub>1</sub> cluster spectra, Figure 6.7 and Table 6.6, are very similar to one another. The MgPc(H<sub>2</sub>O)<sub>1</sub> spectrum  $\sim$  and be assigned as arising from two different cluster manifolds with origin

at 46.6 and 48.3 cm<sup>-1</sup> to the blue of the MgPe  $0_0^0$ . Cluster chromophore out of plane vibrations are observed for both manifolds. The  $A_0^1$  and the  $\Lambda_0^2$  are observed at 13.9 (14.2) cm<sup>-1</sup> and 29.1 (29.8) cm<sup>-1</sup> to the blue of the origins  $B_0^2$  transitions are observed at 57.1 (56.9) cm<sup>-1</sup>. The MgPe(MeOH)<sub>1</sub> and MgPe(EtOH)<sub>1</sub> spectra are similarly assigned.

The  $MgPc(ROH)_1$  series differs from the  $H_2Pc(ROH)_1$  series in three aspects. First, the MgPc(ROH), spectra exhibit hypsochromic shifts with respect to the isolated MgPc spectrum. The direction of the spectral shifts indicate that the solute and solvents interact more strongly in  $S_{0}$  than in  $S_{1}$ . The major difference between the two solute systems studied is the phthalocyano core environment; the pyrrole hydrogens in H<sub>2</sub>Pc are replaced by a magnesium in MgPc. The hypsochromic shift suggests that the core is responsible for the larger ground state interaction. The greater stabilization in the ground state can, in principle, be due to interactions ranging from weak nonbonded vdW interactions to actual complexation in which the solvent donates an electron pair to the solute via the central metal atom.<sup>19</sup> On the basis of the size of the spectral shifts observed, the interaction is most likely due to weak vdW interactions. If the solvent were to donate significant electron density to the central Mg atom in MgPc (a coordina tion bond formation) or if the Mg were to move significantly out of the Pc molecular plane,  $^{34}$  cluster formation would drastically perturb the chromophore electronic environment and yield relatively large spectral shifts. Observation of weak vdW interactions between MgPc and  $H_2O_1$ . EtOH, and phenol in the IR region leads to similar conclusions. $^{35}$ 

Second, the spectral shifts in the MgPc(ROH)<sub>1</sub> series are virtually identical. This observation suggests that the OH group is

intimately involved in the interaction and, furthermore, is largely responsible for the observed spectral shifts. The ground state configuration calculations, Figure 6.9, support the notion that the OH group is the major contributor to the interaction. All three calculated geometries depicted in Figure 6.9 have the solvents situated so that the OH groups point towards the MgPc core. Moreover, one would not expect to observe sharp cluster transitions if the hydrophobic portions of the solvents were the major contributors to the interaction.

Third, the MgPc(ROH), spectra show two nearly isoenergetic cluster origins and vibronic manifolds. These cluster features could be due to either two cluster species which are nearly spectroscopically identical or they could be due to the splitting of the two-fold degenerate chromophore  $S_1$  state via vdW cluster formation. The former situation probably does not occur since these doublets are present in all the MgPc clusters observed. One would not typically expect to see this type of coincidence in the different cluster systems especially when two solvent series are considered. In MgPc clusters, clustering reduces the system symmetry (see Figure 6.9 for example) and the fourfold symmetry axis of the MgPc chromophore is destroyed. The degenerate Q band (E<sub>u</sub> in D<sub>4h</sub>) may thus split into  $Q_x$  and  $Q_y$  bands (B<sub>2u</sub> and B<sub>3u</sub> in  $D_{2h}$ ); the splitting is small, however, due to the small perturbation caused by vdW clustering. Similar removal of degeneracies occurs for benzene clusters, 2.9,10 and for ground state vibrations of MgPc/H<sub>2</sub>0, /EtOH, and /phenol systems.<sup>35</sup>

C. CO<sub>2</sub> Clusters of H<sub>2</sub>Pc and MgPc.

 $H_2Pc$  clustered with  $CO_2$ , Figure 6.10a and Table 6.7, yields a spectrum with a single cluster origin blue shifted by 14 cm<sup>-1</sup> with

respect to the isolated  $H_2Pc \ 0_0^0$ . The cluster manifold is assigned to a  $H_2Pc(CO_2)_1$  species. Cluster chromophore out-of-plane vibrations  $A_0^1$  (15.4 cm<sup>-1</sup>) and  $A_0^2$  (30.7 cm<sup>-1</sup>) built upon the cluster origin are also observed. The calculated ground state geometry, Figure 6.11a, further supports the existence of a single cluster species responsible for the spectrum. In this geometry, the  $CO_2$  moiety is situated over the  $H_2Pc$  core.

The  $H_2Pc(CO_2)_1$  spectrum is different than the other  $H_2Pc$  clusters studied in that it is the only system exhibiting a hypsochromic shift. The exact nature of the interaction responsible for the spectral shift in this system is not known and more work on  $CO_2$  solute/solvent systems is necessary to establish a firm argument. On the basis of the interactions responsible for the spectral shifts in the hydrocarbon and hydrogen bonded solute/solvent systems, the major interaction responsible for the shift appears to involve solvent/solute  $\pi$ -cloud overlap. In the ground state, the solute and solvent most likely interact through  $\pi$ - $\pi$  interactions. Upon excitation the diffuse nature of the solute  $\pi^*$  state reduces the interaction between the  $\pi$  systems and results in a loss in S<sub>1</sub> binding energy relative to S<sub>0</sub>.

MgPc clustered with  $CO_2$ , Figure 6.10b and Table 6.7, reveals a cluster spectrum with two parallel vibronic progressions. The appearance of the two cluster manifolds is probably due to the same type of electronic state splitting that occurs in the MgPc(ROH)<sub>1</sub> systems. The cluster origins are blue shifted by 17.9 and 22.6 cm<sup>-1</sup> with respect to the MgPc  $O_0^0$ . As in H<sub>2</sub>Pc(CO<sub>2</sub>)<sub>1</sub>, well defined cluster chromophore vibronic transitions are observed. The  $A_0^1$  and  $A_0^2$  transitions built upon the origin at 17.9 cm<sup>-1</sup> occur at 19.9 and 31.1 cm<sup>-1</sup> to the blue of this origin. The  $A_0^1$  and  $A_0^2$  transitions built upon the 22.6 cm<sup>-1</sup> origin occur at 21.1 and 32.3 cm<sup>-1</sup> to the blue of the origin. The existence of a single MgPc(CO<sub>2</sub>)<sub>1</sub> species responsible for the cluster transitions is supported by the ground state configuration calculation shown in Figure 6.11b.

The MgPc(CO<sub>2</sub>)<sub>1</sub> cluster spectrum differs from that of the other MgPc clusters studied in that the magnitude of the hypsochromic shift is smaller for MgPc(CO<sub>2</sub>)<sub>1</sub>. This difference can possibly be attributed to the difference in the major mode of interaction between the solute and solvent. In MgPc(CO<sub>2</sub>)<sub>1</sub>, the major interaction may be due to the same type of  $\pi$ -cloud interaction as suggested for the H<sub>2</sub>Pc(CO<sub>2</sub>)<sub>1</sub> interaction as the spectral shifts are comparable for the two systems. If the solvent oxygen/solute magnesium interaction were the major contributor, one could expect a larger hypsochromic shift comparable to those found for the MgPc(ROH)<sub>1</sub> systems.

Summary and Conclusions.

FE spectroscopy is used to probe the optical spectra of vdW clusters of  $H_2Pc$  and MgPc in the vicinity of the cluster  $0_0^0$  transitions. A continuous supersonic molecular jet capable of operating at tempera tures up to  $650^0$  C is employed to generate the ultracold molecular beam. Spectroscopic observables, such as spectral shift and forbidden chromo phore vibronic transitions, combined with computer modeling of ground state cluster intermolecular interactions, allow for the elucidation of the nature of the intermolecular potential and qualitative geometry of the  $H_2Pc$  and MgPc clusters studied. The conclusions drawn from this work are as follows:

1) The cluster vibronic spectra and theoretical calculations suggest that stable H<sub>2</sub>Pc and MgPc clusters have solvents situated over the center of the phthalocyano core. Local minima over peripheral solute rings are either nonexistent or too shallow to accommodate minimum energy bound state geometries;

2) H<sub>2</sub>Pc/hydrocarbon clusters are similar to the benzene and N-heterocycle/hydrocarbon clusters previously studied. The spectral shifts in these solute/solvent series are all bathychromic. The magnitudes of the spectral shifts increase with increasing solvent size and polarizability. The qualitative cluster geometries responsible for the observed spectra are similar with respect to solute/solvent orientation is concerned;

3) In both  $H_2Pc$  and MgPc clusters, the  $H_2O$ , MeOH, and EtOH moieties are situated over the Pc core in such a manner that the OH groups are intimately involved in the intermolecular interactions and contribute significantly to the spectral shifts;

4) Hydrogen bonding may be occurring to some extent between  ${\rm H_2P_C}$  and solvent ROH moieties;

5) MgPc clusters display weak vdW interactions between the cluster solute and solvent. Actual complexation in which the solvent donates an electron pair to the solute does not occur:

6) Forbidden low frequency cluster chromophore out-of plane vibronic transitions are induced by clustering in both  $H_2Pc$  and MgPc systems. Intensity due to this motion arises from the reduction of the chromophore symmetry in the clusters. The perturbation may be large enough to cause the cluster chromophore geometry to change in an attempt to maximize  $\pi$ -cloud overlap with the solvent: and

7) Excited electronic state splitting occurs in the MgPc clusters due to the reduction in system symmetry upon cluster formation. The degenerate Q band appears to split into its two components,  $Q_x$  and  $Q_y$ .

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#### REFERENCES

- D.H. Levy, L. Wharton and R.E. Smalley, "Chemical and Biochemical Applications of Lasers," (Academic Press, New York, 1977) Vol. IJ.
- 2. M. Schauer and E.R. Bernstein, J. Chem. Phys. 2, 726 (1985).
- M. Schauer, K.S. Law and E.R. Bernstein, J. Chem. Phys. <u>2</u>, 736 (1985).
- 4. K.S. Law and E.R. Bernstein, J. Chem. Phys. <u>82</u>, 2856 (1985).
- K.S. Law, M. Schauer, and E.R. Bernstein, J. Chem. Phys. <u>81</u>, 4871 (1984).
- E.R. Bernstein, K. Law, and M. Schauer, J. Chem. Phys. <u>80</u>, 207 (1984).
- M. Schauer, K. Law, and E.R. Bernstein, J. Chem. Phys. <u>81</u>, 49 (1984).
- 8. J. Wanna and E.R. Bernstein, J. Chem. Phys. 84, 927 (1986).
- 9. J. Wanna, J.A. Menapace, and E.R. Bernstein, J. Chem. Phys. <u>85</u>, 1795 (1986).
- 10. J.A. Menapace and E.R. Bernstein, J. Phys. Chem. 91, 0000 (1987).
- 11. J.A. Menapace and E.R. Bernstein, J. Phys. Chem., to be published (1987).
- 12. F.H. Moser and A.H. Thomas, "The Phthalocyanines", (CRC Press Boca Raton, 1983) Vol. I and II.
- R.O. Loutfy, C.K. Hsiao, and R. Ho, Can. J. Phys. <u>61</u>, 1416 (1983);
   P.C. Rieke and N.R. Armstrong, J. Phys. Chem. <u>89</u>, 1121 (1985);
   T.J. Klofta, P.C. Riecke, C.A. Linkous, W.J. Buttner,
   A. Nanthakumar, T.D. Mewborn and N.R. Armstrong, J. Electrochem. Soc., <u>132</u>, 2134 (1985);
   M. Shimura and A. Toyoda, Jpn. J. App. Phys. Part 1 <u>23</u>, 1462 (1984).
- M. Maitrot, G. Guillaud, B. Bondjema, J.-J. Andre, H. Strzelecka, J. Simon and R. Even, Chem. Phys. Lett. <u>133</u>, 59 (1987).

- H. Wohltjen, W.R. Barger, A.W. Snow and N.L. Jarvis, IEEE Transactions on Electron Devices, <u>32</u>, 1170 (1985); S. Baker, G.G. Roberts and M.C. Petty, IEEE Proc. Part I, Solid State Electron Devices <u>130</u>, 260 (1983); R.L. van Ewyk, A.V. Chadwick and J.D. Wright, J. Chem. Soc. Far. I, <u>76</u>, 2194 (1980).
- J.R. Darwent, Chem. Commun., 805 (1980); T. Tanno, D. Wohrle, M. Kaneko, and A. Yamada. Ber. Bunsen. Physik. Chem. <u>84</u>, 1032 (1980); G. Melendon and D.S. Miller, Chem. Commun., 533 (1980); I. Okura, M. Takeuchi and N. Kimthuan, Chem. Lett. 765 (1980).

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17. J.R. Bolton, Solar Energy, <u>20</u>, 181 (1978).

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- J. Barber, "Primary Processes of Photosynthesis", (Elsevier, New York, 1977).
- D. Dolphin, "The Porphyrins" (Academic Press, New York, 1979), Vols. III, V, and VII; J.E. Falk, "Porphyrins and Metalloporphyrins", (Elsevier, New York, 1964).
- F.A. Momany, L.M. Carruthers, R.F. McGuire and H.A. Scheraga, J. Phys. Chem. <u>78</u>, 1595 (1974); G. Nemethy, M.S. Pottle, and H.A. Scheraga, J. Phys. Chem. <u>87</u>, 1883 (1983).
- 21. B.F. Hoskins and S.A. Mason, Chem. Commun., 554 (1969); The phthalocyano skeleton in MgPc is assumed to be the same as  $H_2Pc$  in the calculations. The  $H_2Pc$  imino hydrogens are replaced by a magnesium atom at the center-most position to model the MgPc structure.
- 22. T.M. Miller and B. Bederson, Adv. At. Mol. Phys. <u>13</u>, 1 (1977).
- 23. A. Bondi, J. Phys. Chem. <u>68</u>, 441 (1964).
- 24. R.A. Scott and H.A. Scheraga, J. Chem. Phys. <u>45</u>, 2091 (1965).
- A. Henriksson and M. Sundbom, Theor. Chim. Acta. (Berl.) <u>27</u>, 213 (1972); A.M. Schaffer, M. Gouterman, and E.R. Davidson, Theor. Chim. Acta. (Berl.) <u>30</u>, 9 (1973).
- 26. G. Herzberg, "Molecular Spectra and Molecular Structure III. Electronic Spectra and Electronic Structure of Polyatomic Molecules" (van Nostrand, New York, 1966).
- 27. R.F. McGuire, F.A. Momany, and H.A. Scheraga, J. Phys. Chem. <u>76</u>, 375 (1972).
- 28. P. Jonsson, Acta. Cryst. B32, 232 (1976).
- E.B. Wilson Jr., J.C. Decius, and P.C. Cross, "Molecular Vibrations: Theory of Infrared and Raman Vibrational Spectra" (McGraw-Hill Book Co., Inc., 1955).
- 30. J.A. Menapace, Ph.D. Thesis, Colorado State University, 1987.

- P.C. Painter and J.L. Koenig, Spectrochim. Acta. <u>33A</u>, 1019 (1977).
- 32. P.S.H. Fitch, C.A. Haynam, and D.H. Levy, J. Chem. Phys. <u>73</u>, 1064 (1980).
- H.C. Longuet-Higgins, C.W. Rector and J.R. Platt, J. Chem. Phys. <u>18</u>, 1174 (1950).
- 34. M.S. Fischer, D.H. Templeton, A. Zalkin and M. Calvin, J. Am. Chem. Soc. <u>93</u>, 2622 (1971).

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35. B. Stymne, F.X. Sauvage and G. Wettermark, Spectrochim. Acta. <u>35A</u>, 1195 (1979).

#### CHAPTER SEVEN

#### UNPUBLISHED RESULTS

This chapter discusses the results of an out-of-plane normal coordinate analysis on  $H_2Pc$  and the results of the 2-color TOFMS experiments on  $H_2Pc$  and MgPc. These results are presented to establish an up-to-date record of our work in these areas and to outline the basic approaches used in the analyses.

Out-of-Plane Normal Coordinate Analysis on Isolated H<sub>2</sub>Pc.

Out-of-plane fundamental and/or overtone vibronic transitions are observed in the vicinity of the  $0_0^0$  transitions in the H<sub>2</sub>Pc and MgPc spectra as well as in all the H<sub>2</sub>Pc and MgPc solute/solvent van der Waals clusters presented in Chapter 6. Elucidation of the out-of-plane motion in these molecules has proven useful in the understanding of the cluster spectra and in the identification of the number of different clusters of a specific composition observed. In this section, we report the details of the out-of-plane normal coordinate analysis on isolated H<sub>2</sub>Pc. This normal coordinate analysis is used to determine the nature of the molecular motion occurring in both the H<sub>2</sub>Pc(MgPc) molecule and H<sub>2</sub>Pc(MgPc) cluster spectra.

The out-of-plane normal coordinate analysis on H<sub>2</sub>Pc is conducted using the FG matrix methods described by Wilson et al.<sup>1</sup> These methods essentially involve expressing the secular equation of N-3 coupled harmonic oscillators describing the out-of-plane motion in matrix form and solving for its N-3 non-zero eigenvalues and eigenvectors. To simplify the calculations, the molecular symmetry of  $H_2Pc$  is utilized to symmetry (block) factor the potential energy (<u>F</u>) and kinetic energy (<u>G</u>) matrices to yield "submatrices" each of which describes a particular species of vibrational motion dictated by the vibrational symmetry. The analysis is set up by 1) determining the symmetry of the vibrational motion, 2) selecting a complete set of internal coordinates to describe the motion, 3) obtaining the elements of the <u>F</u> and <u>G</u> matrices in this coordinate scheme, 4) constructing normalized symmetry coordinates using the complete internal coordinate set as a basis, and 5) symmetry factoring the <u>F</u> and <u>G</u> matrices in

The H<sub>2</sub>Pc molecule is depicted in Figure 7.1a. This planar molecule contains 18 hydrogen atoms, 32 carbon atoms, and 8 nitrogen atoms, and has 55 out-of-plane normal vibrations that are distributed among the following irreducible representations under the D<sub>2h</sub> point group:

$$\Gamma_{vib} = 15 B_{1u} + 13 A_u + 13 B_{2g} + 14 B_{3g}$$

Two types of internal coordinates are used to describe the  $H_2Pc$ out-of-plane vibrational motion, bond torsion and out-of-plane bond wagging. The bond torsion coordinate, Figure 7.1b, is defined in the situation when the atoms (2 and 3) at each end of a bond are also bonded to additional atoms (1 and 4) by bonds not collinear with the bond connecting atoms 2 and 3. The terminal bonds taken together with the connecting bond define two planes. Nuclear deformation may thus be expressed by the angle z defined as the dihedral angle between the two

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### Figure 7.1

a) Out-of-plane internal coordinates used in  $H_2Pc$  NCA. Coordinates a-n are bond torsions and coordinates o-w are bond wags. b) Definition of bond torsion  $\tau$  in terms of bond lengths and bond angles. c) Definition of bond wag  $\gamma$  in terms of bond lengths and bond angles.

planes. The displacement of the atoms in this internal coordinate for atoms 1-4 bonded in sequence is expressed by the following  $\hat{s}$  vectors.<sup>1</sup>

7.1a 
$$\tilde{s}_{\tau_1} = \frac{-\hat{e}_{12} \times \hat{e}_{23}}{r_{12} \sin^2 \phi_2}$$

7.1b 
$$\tilde{s}_{\tau_2} = \frac{r_{23} - r_{12} \cos \phi_2}{r_{23} r_{12} \sin \phi_2} - \frac{\hat{e}_{12} \times \hat{e}_{23}}{\sin \phi_2} - \frac{\cos \phi_3}{r_{23} \sin \phi_3} - \frac{\hat{e}_{13} \times \hat{e}_{32}}{\sin \phi_3}$$

7.1c  $\tilde{s}_{\tau_3} = [(14)(23)] \bar{s}_{\tau_2}$ 

7.1d 
$$\bar{s}_{\tau_A} = [(14)(23)] \bar{s}_{\tau_A}$$

for which the terms in square brackets are permutation operators. Equations 7.1c and 7.1d thereby state that the  $\dot{s}_{\tau_3}$  and  $\ddot{s}_{\tau_3}$  vectors can be obtained by permutation of atoms 1 and 4, and 2 and 3 in the expressions for  $\bar{s}_{\tau_1}$  and  $\bar{s}_{\tau_2}$ .

The out-of-plane wagging coordinate. Figure 7.1c is defined at an atom for which three coplanar bonds are coincident. The deformation angle 7 is formed by atoms 2, 3, and 4. The spectors describing the atomic displacements in this internal coordinate are  $\frac{1}{2}$ .

7.2a 
$$\bar{s}_{\gamma_2} = \frac{1}{r_{41}}$$

7.2b 
$$\bar{s}_{\gamma_2} = \frac{\sin \phi_2}{r_{42} \sin \phi_1}$$

7 2c 
$$s_{7,3} = \frac{\sin \phi_3}{r_{43} \sin \phi_1}$$
  
7 2d  $s_{7,4} = \frac{1}{r_{41}} = \frac{\sin \phi_2}{r_{42} \sin \phi_1} = \frac{\sin \phi_3}{r_{44} \sin \phi_1}$ 

In order to obtain a kinematically complete set of internal vibrational coordinates bond torsions are chosen at each non terminal bond, and out of plane wagging coordinates are chosen at each atom for which three coplanar bonds meet  $\frac{2}{2}$ . This prescription coordinates and at out of plane wagging coordinates which torsion coordinates and at out of plane wagging coordinates which completely describe the out of plane motion in the H<sub>2</sub>Pr molety. Twenty seven redundancies are distributed among the sibrational symmetries  $R_{10}(1) = A_{0}(9) = B_{20}(7)$  and  $R_{30}(7)$  in this internal coordinate scheme. These redundancies are carried through the calculations so that symmetry ourdinates can be used to block factor the secular equation cost the four symmetry species sets.

The 18 bond forsion coordinates are constend to Di Symmetry equivalent sets designated as a n-pu-Figure T-ball. The atoms is electrocoordinate are number using a can excel intention of statements with atom 1 moving in the induced and its moving to the boost operation measured in a counterclockwise direction when looking down the bond between it mo 2 and 3 with atom 2 nearer to the observer. When two is more that in the molecule contain the bond forsion coordinates the coordinate is oriented so that its terminal atoms point towards the H<sub>1</sub>Bell order and its terminal bonds are situated in a <u>sym</u> conformation.

The 34 out of plane wagging coordinates are located in 9 symmetry equivalent sets designated as o w in Figure 7.1a. The atoms in each coordinate are numbered using a counterclockwise convention (Figure  $7(t_{c})$  = Positive displacement in  $\gamma$  is measured by out of plane motion of atom 1 in the z direction. Coordinates containing hydrogen (o s) are numbered so that the hydrogen atom is designated as atom 1. (coordinates t and v are oriented such that atom 1. circesponds to the benzene C ff carbon atom. Coordinates u and w are numbered so that the bridge nitrogen atom is designated as atom 1.

The elements of the <u>G</u> matrix are obtained using the internal oordinate + s vectors (Equations 7-1 and 7-2) and

the which  $z_{1}^{-1}$  the reciprocal of the mass of atom a. The non-zero  $\frac{m_{a}}{m_{a}}$  the summation are those corresponding to atoms common to the two internal coordinates t and t in question. The geometric parameters used to determine the internal coordinate s vectors are based upon the resolds. It is neutron diffraction analysis of  $R_{c}Pc^{-1}$ . These parameters are capitly modified however to reduce the total number of different 5 matrix elements that need to be calculated. In this approximation all the isoindole subunits comprising  $R_{c}Pc^{-1}$  are assumed to contain identical interatomic distances and hond angles. The interatomic distances and hond angles. The interatomic parameters determined from the crystallographic data. The geometric parameters used in the calculations are presented in Figure 7.2. The summatrix elements calculated from Equations 7.1.3 are given in Table 7.1.



### Figure 7.2

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Geometric parameters used in  $H_2Pc$  NCA. Bond lengths are expressed in angstroms and bond angles are in degrees. The parameters are averaged to  $D_{4h}$  symmetry to simplify the calculations.

### TABLE 7.1

## Kinetic Energy (G) Matrix Elements for $H_2Pc$ Out-of-Plane Motion <sup>a</sup>

the states

Matrix Element	Matrix Element	Matrix Element
(A-g/mol) <sup>-1</sup>	$(A-g/mol)^{-1}$	$(A-g/mol)^{-1}$
G(alal) .56689	G(b1b1) .56689	G(c1c1) .56689
G(a1b1)45351	G(b1b2) .22676	G(c1c2) .22676
G(a1b2)45351	G(b1c1)45351	G(c1d1)45351
G(a1c1) .22676	G(b1c2)11338	G(c1e1) - 21223
G(a1c2) .22676	G(b1d1) .22676	G(c1e2)18273
G(ald1)11338	G(b1e1) .08058	G(c1f1) .09890
G(a1o1) .47094	G(b1e2) .05107	G(c1f2) .04945
G(a1o2)47094	G(b1f1)04945	G(c1o1) .26002
G(alp1)26002	G(b1o1)47094	G(c1o2)04909
G(a1p2) .26002	G(b1o2) .26002	G(c1p1)47094
G(alt1) .04909	G(b1p1) .47094	G(c1p2) .04909
G(alt2)04909	G(b1p2)04909	G(c1t1) .39084
	G(b1t1)22615	G(c1t2)20143
	G(b1t2) .03674	G(c1u1)07794
	G(b1u1) .03897	G(c1u2) .03897
G(d1d1) .56689	G(e1e1) .32338	G(f1f1) .33332
G(d1e1) .26331	G(e1e2) .10262	G(f1f2)27330
G(d1e2) .26331	G(e1f1) - 26292	G(f1g1) .29100
G(d1f1)09890	G(e1f2) .10302	G(f1g2) - 21388
G(d1f2) - 09890	G(e1g1) - 25421	G(f1h1) - 09307
G(d1o1)04909	G(e1g2) .05108	G(f1h2) .05879
G(d1o2) .04909	G(e1h1) .09148	G(f1p1) - 04283
G(d1p1) .26002	G(e1p1) .06978	G(f1t1) .17404
G(d1p2) - 26002	$G(p_1p_2) = .04423$	G(f1t2) 00738
G(d1t1) 37849	G(e1t1) = 27633	G(f(u)) = 29377
G(d1t2) .37849	G(p1t2) = 16750	$G(f_{1}u_{2}) = 18581$
G(d1u1) 07794	G(=1u1) 28850	0(1102) 10001
G(d1u2)07794	G(e1u2) 00059	
G(g1g1) 78450	G(h1h1) 78480	0(1111) 58699
$G(g_1g_2) = 06129$	G(h1h4) = 06129	G(1111) - 48351
G(g(h)) = 59954	G(h1m2) 05108	
G(g1n4) 09148	G(h1m3) = 25421	
G(g1n3) 05879	G(1100) = 20021 G(1100) = 21400	Q(11K1) 22070 Q(11k2) 22676
G(g1nd) = 09307	G(b1=4) 20100	G(11R2) 220/8
G(g1a2) 04788	G(b1-2) 25000	
G(g1+1) 10384	U(1186) *.23707 G(b1+1) - 04887	
C(g1u1) _ A0843	0(N)LIJ - 04007 C(5141) - 25000	G(1192) 47094
G(#102) _ G4284	G(D1U1) 20829 G(D1U1) 20829	G(11r1) - 26002
C(g1u4) C4866	G(NIV4) - 10364 G(b)	G(11r2) 26002
0(8144) 06007	G(h1w3) 04284	G(11VI) 04909
0(81#4) - 52858	G( <b>h1w4) 48542</b>	G(11v2) 04909

### TABLE 7.1 (Continued)

Matrix Element	Matrix Element	Matrix Element (A-g/mol) <sup>-]</sup>		
(A-g/mol) -1	(A-g/mol) <sup>-1</sup>			
G(1111) 58690	C(6161) 80000			
G(1112) 22678	G(K1K1) .30089 G(k1k2) 33878			
G(j)j(2) = .22070 G(j)k(1) = .45351	G(K1K2) .22070 G(k1)1) _ AB981			
G(11k2) - 1133R	G(kint) = 21222	G(11mc) = 00000		
G(11m1) .08058	G(k1n1) 09890	G(11n2) = 00800		
G(11m2) .05107	G(k1n2) 04945	G(1161) = 0.4909		
G(j1n1)04945	G(k1g1) = 26002	G(11q2) 04909		
G(j1q) - 47094	G(k1g2) = .04909	G(11qL) 26002		
G(j1g2) 26002	G(k1r1) - 47094	G(11r1) = 26002		
G(j1r1) .47094	G(k1r2) .04909	G(11v1) = 37849		
G(j1r2) - 04909	G(k1v1) .39084	G(11v2) 37849		
G(j1v1) - 22615	G(k1v2) - 20143	G(11w1) 07794		
G(j1v2) .03674	G(k1w1)07794	G(11w2) - 07794		
G(j1w1) .03897	G(k1w2) .03897			
G(m1m1) .32338	G(nini) .33 <b>332</b>	G(0101) 1.40434		
G(m1m2) .10262	G(njn2) - 27330	G(0102) - 28030		
G(m1n1) ~ 26292	G(n1r1)04283	G(o1p1) - 28030		
G(m1n2) .10302	G(n1s1) .31472	G(01p2) 04252		
G(m1r1) .06978	G(nlv1) .17404	G(citi) 04252		
G(m1r2)04423	G(n1v2) .00738			
G(m1s1)19027	G(n1w1) - 29377			
G(m1v1) - 27663	G(n1w2) - 18581			
G(m1v2) .16750				
G(m1W1) 28850				
G(m1w2) .00059				
G(p1p1) 1.40434	G(q1q1) 1.40434	G(r1r1)   40434		
G(p1t1) - 25098	G(q1q2) - 28030	G(r1v1) 25096		
G(p1t2) 03182	G(q1r1) - 28030	G(r1v2) 03182		
G( <b>p1u1</b> ) 03375	G(q1r2) 04252	G(r1w1) 03375		
	G(q1v1) 04252			
G(111) 1 55480	G(t1t1) 38645	G(u101) 42178		
G(s1v1) 03210	G(t1t2) 16585	G(u1u2)		
G(\$1\$2) 03210	G(tlul) 18867	G(u)w4) 04094		
G(s1w1) 21009	G(t1u2) 02525			
U(\$1W2) 21009				
G(V1V1) 38645	G(w1w1) 42170			
(i(vIv2) 16585	G( <b>w1w2</b> ) 02994			
G(VIW1) 18867				
G(v1w2) 02525				

a) Only those elements necessary to generate the <u>U</u>matrix in terms of symmetry coordinates are shown

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The elements of the P matrix in terms of the bond torsion and out-of-plane wagging coordinates, Table 7.2, are simply expressed as valence force constants  $F_{tt}$  for which t and t' correspond to the two internal coordinates in question. For the purpose of this calculation. the F matrix elements are taken as the diagonal force constants describ ing the out of plane motion of benzene <sup>4</sup>. In this approximation, all  ${\sf H}_{0}{\sf Pc}$  bond forsion force constants are assumed to be the same as those corresponding to the C-C torsions in benzene. The out of plane wagging force constants are assumed to be the same as those corresponding to benzene out of plane C H wags . All off diagonal force constants are taken as zero. This force field scheme, even though not quantitatively incourate will allow for the elucidation of the general nature of the out of plane motion in HyPC as it will yield characteristic frequencies for the particular molecular segments participating in the various sibrational motions. No ittempt is made to refine the force field soft an the calculations since the experimental and constants on the set of plane shrata as so Hyperso lympted to these possessing estimate  $\sim$  where the constraint is a track  $5^{-12}$  , in the constraint of the track the track  $\sim$  . The the company of the second statements that shall be the the formation of the second statements of the second state and the rest of the report of the repair of the second second second second second second second second second

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Potential Energy (F) Matrix Elements for $H_2Pc$ Out-of-Plane Notion <sup>a</sup>						
Matrix Element (mdyn-A/rad <sup>2</sup> )		Matrix El (mdyn-A/r	ement ad <sup>2</sup> )	Matrix Element (mdyn-A/rad		
P(alal)	1190	P(b1b1)	. 1190	F(c1c1)	. 1190	
F(d1d1)	1190	P(ele1)	. 1190	F(f1f1)	. 1190	
F(g1g1)	1190	P(hihi)	. 1190	F(1111)	. 1190	
₽(j1j1)	1190	F(k1k1)	. 1190	F(1111)	. 1190	
F(=1=1)	1190	P(nJn1)	. 1190	F(0101)	. 3237	
F(pip1)	3237	₹(q1q1)	3237	F(r1r1)	3237	
F(=1=1)	3237	F(titi)	3237	₹(ulul)	3237	
P(v1v1)	3237	F(m1m1)	1317			

a) Only those elements necessary to generate the <u>P</u> matrix in terms of symmetry coordinates are shown

TABLE 7.2

B <sub>1u</sub> :	B=1/2(b1-b2+b3-b4)	B <sub>2g</sub> :	$B \approx 1/2(b1-b2-b3+b4)$
	U=1/2(C1-C2+C3-C4)		U=1/2(C1-C2-C3+C4)
	L-1/2(e1-e2+e3-e4)		E=1/2(e1-e2-e3+e4)
	F=1/2(F1-F2+F3-F4)		P=1/2(f1-f2-f3+f4)
	G=1/2(g1-g2+g3-g4)		G=1/2(g1-g2-g3+g4)
	H=1/2(h1-h2+h3-h4)		H=1/2(h1-h2-h3+h4)
	J=1/2(j1-j2+j3-j4)		$I = 1/\sqrt{2(11-12)}$
	K=1/2(k1-k2+k3-k4)		J=1/2(j1+j2-j3-j4)
	M=1/2(m1-m2+m3-m4)		K=1/2(k1+k2-k3-k4)
	N=1/2(n1-n2+n3-n4)		L=1//2(11-12)
	0=1/2(01+02+03+04)		M = 1/2(m1+m2-m3-m4)
	P=1/2(p1+p2+p3+4p)		N=1/2(n1+n2-n3-n4)
	Q=1/2(q1+q2+q3+q4)		0=1/2(01+02-03-04)
	R=1/2(r1+r2+r3+r4)		P=1/2(p1+p2-p3-p4)
	S=1//2(s1+s2)		Q=1/2(q1-q2-q3+q4)
	T=1/2(t1+t2+t3+t4)		R=1/2(r1-r2-r3+r4)
	U=1/2(u1+u2+u3+u4)		T=1/2(t1+t2-t3-t4)
	V=1/2(v1+v2+v3+v4)		U=1/2(u1+u2-u3-u4)
	W=1/2(W1+W2+W3+W4)		V=1/2(v1-v2-v3+v4)
			W = 1/2(W1 - W2 - W3 + W4)
A., :	A=1/J2(a1+a2)	Basi	A=1/J2(a1-a2)
u	B=1/2(b1+b2+b3+b4)	-3 <b>g</b>	B=1/2(b1+b2-b3-b4)
	C=1/2(c1+c2+c3+c4)		C=1/2(c1+c2+c3+c4)
	D=1/2(/d1+d2)		D=1//2(d1-d2)
	E=1/2(e1+e2+e3+e4)		E=1/2(e1+e2-e3-e4)
	F=1/2(f1+f2+f3+f4)		F=1/2(f1+f2-f3-f4)
	G=1/2(g1+g2+g3+g4)		G=1/2(g1+g2-g3-g4)
	H=1/2(h1+h2+h3+h4)		H=1/2(h1+h2-h3-h4)
	I = 1/J2(11+12)		I = 1/2(11 - 12 - 13 + 14)
	J = 1/2(11 + 12 + 13 + 14)		$K = 1/2/k(1 k_2 k_3 k_4)$
	$K = 1/2(k_1 + k_2 + k_3 + k_4)$		M = 1 / 2 (m1 - m2 - m3 + m4)
	L=1/J2(11+12)		N = 1/2 (n1 + n2 + n3 + n4)
	$M_{v1}/2(m1+m2+m3+m4)$		
	$N=1/2(n_1+n_2+n_3+n_4)$		$P_{-1}/2(p_1, p_2, p_3, p_4)$
	0 = 1 + 2 + 01 + 02 + 03 + 04		$(p_1)^2(p_1)^2(p_2)^2(p_3)^2(p_4)$
	P=1/2(n1, n2+n3, n4)		$P_{-1} = c(q_1 + q_2 - q_3 - q_4)$
	0-1 2(0) 02×03 04)		
	vrs ∠iųs ųstųotųe) Polojei eokeoloje		$\begin{array}{cccccccccccccccccccccccccccccccccccc$
	N-1 6111 16473 F4) T-1 3/41 43.43 441		$\frac{1}{2} + \frac{1}{2} + \frac{1}$
	ISI 2131 3243 333 341 11 1 3741 433 441		U 1 2(u1 u2 u3+u4)
			V = 1 - 2(v1 + v2 - v3 - v4)
	V=1 (2(v) v2+v3 v4)		₩~1 2(w1+w2 w3 wd)

2(#1 #2+#3 #4)

Normalized Symmetry Coordinates for H<sub>2</sub>Pc in Terms of Bond Torsion and Out-of-Plane Wagging Coordinates.

### TABLE 7.4

Mode	Calç	Obsą <sup>a</sup>	Mode	Calç	Obsd <sup>a</sup>
	(cm <sup>-1</sup> )	(cm <sup>~1</sup> )		(cm <sup>-1</sup> )	(cm <sup>~1</sup> )
B <sub>111</sub> 1	14.8	15.6	B <sub>2</sub> 29	38.6	42.7/50.3
2	24.5	25.9	<sup>2</sup> <b>g</b> 30	72.1	88/101.6
3	84.0	81.9/90	31	154.5	
4	145.3	127	32	185.2	
5	158.9	140	33	317.2	
6	230.5	282	34	343.1	
7	333.7	342	35	510.1	
8	362.3	342	36	686.8	
9	608.3		37	737.4	
10	735.3	725	38	791.1	
11	739.6	725	39	889.0	
12	777.5	775	40	968.1	
13	962.7	945	41	1146.3	
14	967.8	945			
15	1014.4				
A., 16	33.1	35.7	Bo., 42	38.2	42.7/50.3
<sup>u</sup> 17	71.7	70.9	-3g 43	71.4	88/101.6
18	136.4		44	147.2	
19	310.4		45	183.3	
50	321.0		46	315.8	
21	427.6		47	339.5	
22	593.2		48	498.1	
23	690.5		49	648.3	
24	747.9		50	728.4	
25	884.5		51	740.7	
26	891.8		52	887.2	
27	1146.1		53	962.7	
28	1146.8		54	1013 2	
			55	1046 3	

# Observed and calculated frequencies and tentative assignments of the out-of-plane vibrations in H<sub>2</sub>Pc.

a) Prom Chapter 6 and References 5-7.

frequencies obtained from the vibronic spectrum discussed in Chapter 6 and from infrared spectra.<sup>5-7</sup> The comparison between the experimental vibronic transitions and the calculated mode frequencies is made on the premise that the ground and excited state potential surfaces are nearly superimposable. This notion is demonstrated experimentally by FE and DE spectra of  $H_2Pc$  for which the vibrational energies in the  $S_1$  and  $S_0$ states differ by at most 5-10%.<sup>8</sup> Overall, the calculated frequencies compare quite well with those observed given the qualitative nature of the force field used in the analysis. Based upon this calculation, we propose tentative out-of-plane mode assignments for the experimentally observed out-of-plane modes.

The eigenvectors corresponding to the calculated vibrations are listed in Table 7.5. From the eigenvector normal mode forms, the outof-plane motion can be qualitatively categorized into 1) macrocycle ring deformation, 2) isoindole ring deformation, and 3) C(N)-H out-of-plane wagging motion types.

The 0-100 cm<sup>-1</sup> region is dominated by macrocycle ring deformation for which large amplitude out-of-plane motion of the four isoindole subunits occurs. This motion is located principally at the pyrrole  $\alpha$ carbon/bridge nitrogen bonds (internal coordinates g and h). Modes 1. 2. 29, and 42 possess potential energy distributions (PED's) ranging from 40-70% in these coordinates.<sup>9</sup> From the H<sub>2</sub>Pc vibronic spectrum presented in Chapter 6, these modes are observed as symmetric over tones at 31 2(1<sup>1</sup><sub>0</sub>), 51 8(2<sup>2</sup><sub>0</sub>), 85 4(100.6) (29<sup>2</sup><sub>0</sub> or 42<sup>2</sup><sub>0</sub>), and 100.6 cm<sup>-1</sup>(85.4 cm<sup>-1</sup>) (42<sup>2</sup><sub>0</sub> or 29<sup>2</sup><sub>0</sub>) which places their forbidden funda mental transitions at 15.6, 25.9 42.7(50.3), and 50.3 cm<sup>-1</sup>(42.7 cm<sup>-1</sup>) respectively. Mode 3 is also observed in the far infrared at about

### TABLE 7.5

# Out-of-Plane Eigenvector Normal Modes Calculated for $H_2^{Pc}$ .

Mode	Eigenve	ctor in 1	Cerms of S	Symmetry C	Coordinate	es a	
<sup>B</sup> 1u <sup>1</sup>	1925	E+.3323	F~.5829	G5838	H+.1920	M3315	]
2	+.1445 1601	E2212 T4253	F4510 U1593	G+.4565 V4301	H+.1508 W	M2315	ł
3	1407	B+.1407	C+.2406 K- 2630	E3954 M+ 4325	F2622 N- 1782	G2505 T- 3704	1
	+.1546	V+.3658	W . 2000	M. 1020		1.0104	
4	+.2500 +.2154	B2500 M3503	C+.1753 N+.2378	E2864 T+.3941	F+.4221 V	J4221	I
5	4540 +.2686	B+.4540 J2686	C~.1235 K~.1022	E+.1986 N3632	F+.3185 T+.2143	G+.2819 V	ł
6	2158 2686	B+.2158 J+.2686	C+.2602 K+.2611	E4307 M4326	F+.3443 N	G3574	ł
7	+.4007 +.4019	B4007 K1001	C+.1469 N~.1467	F+ . 1347 P+ . 1469	G+.2009 R3105	H4019 T+.3080	1
8	+.3543 3398	B3543 K1220	C1204 N1550	F+ 3123 P- 1486	G3064 R3523	H+ . 3398 T 3374	•
9	- 2060 + 3238	E+.3418 S+.1214	F+ . 3697 T 2491	H+.2649 U1993	M- 4400 V+ 4342	N- 1226 W	F
10	+.1423 +.1549	F+.2272 R1440	G <b>1518</b> U	H+.7619	()+ 4266	P+ 2763	(
11	► 2943	0+.1455	P8353	Q 4128	R		
12	<b>2613</b> + . 1401	<b>E+ 4334</b> T 4163	P+:5511 U 1263	G 3320 W	H 2233	0 2469	
13	+ 2919 1314	J 2919 V	K+ 1121	P 4025	Q+ 7437	R+ 2462	
14	· 30 <b>4</b> 0	B- 3040	C 4032	0+ 7618	P 1213	R 1672	
15	+ 1864 + 1115	G 3675 V 2899	H 2153 W	M+ 3567	N 1856	R - 1001	
TABLE 7.5 (Continued)

Mode	Eigenvector in Te	erms of Symmetry C	oordinates <sup>a</sup>
А <sub>и</sub> 16	+.1503 C1656 1503 K+.1656	D+.3721 E1410 L3721 M+.1410	<b>F3954</b> G+.3954 H N <b>3513</b> U+ 3513 W
17	+.1385 A2577 1825 H+.1385	C+.2891 D4917 I2577 K+.2891	E+.1849 P1825 G L4917 M+ 1849 N
18	+.1640 A2530 +.3436 H1640	C+.2932 D3954 [+.2530 K2932	E+.1482 F3436 G L+.3954 M1482 N
19	1435 U+.1435 4671 A+.4560 +.1435 O+.1435	W B1114 D4671 Q+.1147 U+.1147	I+.4560 J1114 L W
20	• 4538 A- 4675 1189 K- 1184	B+.1189 C+.1184 M1480 O+.1480	E4538 I+.4674 J Q
21	1059 A+ 2007 • 2660 H- 1059 1197 P- 1197	B- 3027 C+ 2501 I+ 2007 J- 3027 R 4232 U- 4232	D- 1274 E+ 2660 G K+ 2501 L- 1274 N W
22	2790 C+ 2637 2637 L 1224 • 2228 V+ 1726	D+ 1224 E+ 4457 M- 2217 P+ 2217 W	G 4457 H+ 2790 K R 2228 T 1726 U
23	2202 C+ 2154 • 3039 M 1141 2859 V+ 2103	D+ 3039 E 1141 N 3936 P 3936 W	F 2202 K+ 2154 1 R 2859 T+ 2103
24	2115 E+ 4645 1335 Q 3217	0 4645 H+ 2115 R+ 1434 T 2901	M-1835 S-2217 R C 1434 R-2401 m
	<ul> <li>2260 ( 2463)</li> <li>4200 0 2953</li> <li>2479 V 1000</li> </ul>	0 1918 F- 2260 F 4200 y 2953 W	Koljąki – urski Mi Rulją≑si tori
<u>'</u> •	2119 - 2389 2389 - 2072 2494 - 78 1172	(), (), (), (), (), (), (), (), (), (),	non over proper e e Galago (1997), King e
. •	2284 A 2922 - 2820 K 1733 - 1343 T+ 1343	8 - 1425 - 2005 19 - 19 20 - 19 44 19	α μ. <b>3.3</b> μ. α. Α. μ. α.
ν.	2 - 24 - X - 2 274 2 2 2 - 14 - 17 2 3 2 2 4 4 - 17 - 17 2 3	Не (1 <b>1</b>	<b>.</b> · • • • •

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# TABLE 7.5 (Continued)

Mode	Eigenvector in T	erms of Sy	ymmetry C	oordinate	s <sup>a</sup>	_
B <sub>2g</sub> 29	+.1179 B1179 2328 M+.2209	C+.7966 T+.3514 R+ 1762	G+.2358 U+.1308	H1072 W	K+.1191	L
30	+.1944 E3189 +.2386 N2474	U+.1803	W	K+.3000	L-,0340	1.1
31	+.4865 B4865 +.4174 T1334	C+.23 <b>56</b> U	E3831	F1991	G2474	H
32	2591 B+.2591 1695 I+.2071 +.2327 W	C+.2861 K2518	E4743 L+.2789	F+.3997 M1045	G3046 N1103	H T
33	+.1522 B1522 1268 K1341	C+.1101 M+.1997	F+.1459 Q+.1220	H−.6255 ₩	I+.6320	J
34	+.5224 B5224 2029 P4420	C+.2596 T+.1149	G+.1907 W	I2203	J+.1214	к
35	+.1951 E3237 +.3102 L1786	F+.1469 R1474	G5439 T+.1811	H+.1498 U1829	J3424 V4103	K W
36	+ 1769 E- 2934 + 1406 N+ 1104	F1897 Q+.4846	G+.2833 R+.2496	K 2768 U+ . 3592	L3744 V2422	M W
37	• 1404 G1394	H+.8457	0+.4460	P+.1139	R	
38	+ 2411 E - 3999 2127 R - 1237	P5561 T+ 3913	G+ 4052 U+ 1936	H+.1672 W	0- 1267	Ŷ
ξų.	1437 G+ 1652 1061 N+ 5692	H+ 1397 Q+ 3794	I 3067 R 3486	K+ 3357 V+ 1822	L+ 2825 W	м
<b>\$</b> ()	• 3071 B 3071	C 4067	0+ 7691	P 1705	T	
4 1	2168 1+4128 +4584 R 1910	1 3424 V	K+ 2172	1 - 1147	₩ ~4~)	
11 - <b>1</b> 2 - <b>1</b> 3	. С. Ал. — 1150 1954 — М. — 1264	D+ 2251 D+ 2174	F - 1402 VI - 1666		H+ 1154	
1	11 - Xeller M. 12 - North M.	2 € 5 7 2 € 2 €	151 - 21 C W	1 <u>1</u> 1 1 1	<u></u>	ч
• •	1 1 1 − 11146 − 1 mil 1114	स्तित्तः क्रांडिलय	5 17 <b>46</b> 5 - 11 - 15	ч 1.(8 <sup>т</sup> ) #	1. <b></b>	

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TABLE 7.5 (Continued)

Mode	Eigenve	ctor in T	erms of S	Symmetry (	Coordinate	s a	
45	+.1615 +.4128 1479	A~.2996 H~.3138 V	C+.2421 J+.3138	D2706 K+.2646	E+.1013 M4396	P 2789 N- 2199	- 6 U
46	+.6131 +.1985	A~.6148 J~.1985	B+.1105 K+.1148	C+.1002 N1941	D+ 1202 0- 1218	E+ 1396 T - 1240	6 V
47	+.2509 +.4985	A~.2843 K+.1890	B+.1431 R+.1240	C+.1198 U+.4040	E2690 V	H- 4985	J
48	1501 +.1623	B+.3268 P+.1237	C2935 S+.1631	D 4908 T+ . 4080	G+.2145 U2014	M-:3564 V+:2426	N W
49	+.2737 +.3052 3401	C~.2640 N+.3212	D2379 P+.1007	E2581 R3008	G+.4123 S+.2819	H- 1837 T+ 1333	M V
50	+.1284 1031 +.2291	C1271 N+.1451 T3401	D2979 O+.4155 U+.1628	E+.1118 P2352 W	F+.4455 Q1579	G3629 R+.2259	H S
51	+.1861 1236	G~.1684 U~.1071	H+.1407 V+.1140	P+.8176 W	Q+.3895	R+.1560	S
52	1411 5828	A+.3139 0~.3970	C3430 P+.3518	D2800 T1677	E+.1051 U	F+.1211	G
53	2953	J+.2953	K+.4070	Q7522	R2439	S+.1332	v
54	1645 +.1130	G+.3629 V~.2908	H2170 W	M+.3595	N1909	R+.7091	S
55	+.3167 4583	A4127 P+.1911	B+.3424 T	C2172	D1148	E+.5470	0

a) Symmetry coordinates contributing less than 1% to the potential energy distribution are not included.

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We make the end of the second terms for modes 3. 16, 17, 40, and 13 are been characteristic in the g and h internal coordinates is they are dominated by pyrrole a carbon bridge mitropen bond toronomic opend to coupled with pyrrole skeleton torsion coordinates (e. f. and m). These modes cover in the distribut spectrum of  $162.7(3\frac{2}{0}-71.4(16\frac{2}{0})-141.7(172))$  $176(20001) < 90\frac{2}{0}$  or 447, and 298.1 m  $\frac{1}{2}(176.4)$  m  $\frac{1}{2}(140\frac{2}{0})$  or  $10\frac{2}{0}$  which places their form the simulamental transitions at 81.9-35.7-70.984(101.6) and 101.0 m  $\frac{1}{2}(38.2)$  m  $\frac{1}{2}$  respectively.

Counded ring deformation occurs mainly in the region between 100 and 700 cm<sup>-1</sup>. The dominant out of plane motions in this region arise from the coupled deformation of the benzene and pyrrole moleties. as can be seen by the large PED contributions from the a-f and i n coordinates in Table 7.5. T. Kobayshi<sup>5</sup> has assigned far infrared bands at 127–140–322, and 342 cm<sup>-1</sup> to isoindole ring deformations on the basis of comparative analyses of free base and metallophthalo cyanines. These bands correspond to the calculated modes 4(145.3 cm<sup>-1</sup>). 5(158.9 cm<sup>-1</sup>): 6(230.5 cm<sup>-1</sup>), 7(333.7 cm<sup>-1</sup>), and 8(362.3 cm<sup>-1</sup>). respectively. The far infrared studies also show that the band observed at 282 cm<sup>-1</sup> contains contributions from both the isoindole ring and bridge nitrogen motions. The eigenvector form for mode 6 corroborates, this assignment as it possesses a PED of about 25% in the pyrrole  $\alpha$ carbon/bridge nitrogen torsion coordinates (g and h).

In the region between 700–1100 cm  $^{-1}$ , C(N)-H out of plane wagging dominates. From the eigenvector normal mode forms, the C-H vibrations are, for the most part, uncoupled from the H<sub>2</sub>Pc ring modes as the out of-plane wagging coordinates o-r account for between 71 and 97% of the PED's responsible for the motions. The infrared spectrum of H<sub>2</sub>Pc

possesses bands at 725, 775, and 945 cm  $^{1}$  which can be attributed to  $B_{111}$  type out-of-plane C-H wagging.<sup>7</sup> The normal coordinate analysis reveals C-H wagging mode frequencies in this vicinity at 735.3 (mode 10), 739.6 (mode 11), 777.5 (mode 12), 962 (mode 13), and 967.8  $cm^{-1}$  (mode 14). The only large discrepancy between the calculated frequencies and the experimental data involves out-of-plane  $(B_{1n})$  wagging motion of the imino hydrogens. This mode is calculated to occur at 1014.4  $cm^{-1}$  (mode 15) with a PED of 49% in the s internal coordinate, but it has been assigned to a band observed in the vicinity of 700  $cm^{-16,7}$ based upon isotopic substitution. Intramolecular hydrogen bonding between the imino hydrogens and adjacent core nitrogen atoms may account for the large difference between the calculated and observed/ assigned modes. However, mode 9 calculated at 608.3  $cm^{-1}$  is near the assigned N-H out of plane wagging frequency and exhibits and 11% PED contribution from the N-H wagging coordinate, s, and 20% from the pyrrole torsion coordinate, m. Performing the normal coordinate analysis using deuterons at the pyrrole nitrogens results in mode 9 red shifting to 536.3 cm<sup>-1</sup> (9H/9D = 1.13) and mode 15 red shifting to 916 cm  $^{1}$  (9H/9D = 1.11). These results suggest that the mode responsuble for the band observed at 711 cm  $^{1}$  could be due to highly coupled motion between the N-H group and the pyrrole ring and not to characteristic N-H motion which should occur in the 900-1000 cm<sup>-1</sup> region.

#### SUMMARY AND CONCLUSIONS:

An out-of-plane normal coordinate analysis is conducted on  $H_2Pc$ to characterize the large amplitude low frequency motion present in both the isolated  $H_2Pc$  and MgPc moieties as well as their respective vdW clusters with small solvents (see Chapter 6). The out-of-plane normal

coordinate analysis is conducted using the FG matrix method. The nuclear motion is modeled by a set of 82 internal coordinates; 48 C-C(N) bond torsions, 18 C(N)-H bond wags, and 16 C-C(N) bond wags. The force field used in the model consists of the diagonal force constants describing the out-of-plane motion in benzene. The resulting eigenvalues and eigenvector normal modes are compared with experiment and tentative assignments are proposed based upon the calculations. In the comparison, observed vibronic symmetric overtone transitions and infrared active  $B_{1U}$  vibrational transitions are utilized. Overall, agreement between the observed and calculated frequencies is excellent given the qualitative nature of the force field used in the analysis and the fact that the force field is not refined/fit to the experiments.

H<sub>2</sub>Pc AND MgPc 2-Color TOFMS Experiments:

Ultracold 2-color TOPMS spectra of  $H_2Pc$  and MgPc could not be observed with the current experimental apparatus configuration available in the laboratory. In initial attempts to obtain 2-color TOPMS spectra of the macrocycles, spectra were observed which were not tunable when the pump laser frequency was varied. Both helium and argon carrier gases were employed to expand the molecules into the vacuum apparatus; however, the spectra were significantly broadened. These results are somewhat surprising since ultracold 2-color TOPMS spectra of small aromatic molecules such as benzene, pyrazine, and aniline have been observed in the existing apparatus.<sup>10-12</sup> One possible explanation for the nontunability is that the molecules are vibrationally/rotationally hot when they reach the section of the apparatus where the spectra are taken. In this case, molecular heating may occur in the section of the apparatus between the supersonic nozzle and the TOFMS flight tube.

In the existing apparatus.<sup>12</sup> Figure 7.3, the supersonic expansion is sampled using a compound angle conical skimmer.<sup>13</sup> The sampled portion of the supersonic expansion then travels about 50 cm to the TOFMS ionization area where it is optically probed. Based upon the aforementioned experimental observations, the expansion could be disturbed to some extent by the presence of the skimmer in the supersonic flow path. In order to understand the interferences in the supersonic flow which result in molecular heating, it is instructive to look at the physical properties of the supersonic expansion before, at, and after the skimmer.

Assuming that the expansion proceeds in an adiabatic and isentropic manner and that the expanding gases can be considered ideal, the centerline translational temperature  $T_1$ , the centerline static pressure  $P_1$ , and the centerline static density  $\rho_1$  of the expansion before the skimmer can be described by<sup>14</sup>

7.4a 
$$T_1 = \left(1 + \frac{\gamma - 1}{2} M_1^2\right)^{-1} T_0$$

7.4b 
$$P_1 = \left(1 + \frac{\gamma - 1}{2} M_1^2\right)^{\frac{-\gamma}{\gamma - 1}} P_0$$

7.4c 
$$\rho_1 = \left(1 + \frac{\gamma - 1}{2} M_1^2\right)^{\frac{-1}{\gamma - 1}} \rho_c$$

as a function of centerline Mach number  $M_1$ , nozzle backing region stagnation temperature  $T_0$ , stagnation pressure  $P_0$ , stagnation density  $\rho_0$ , and  $\gamma = C_{\rm p}/C_{\rm v} = 5/3$  for monatomic carrier gases. The centerline Mach



#### Figure 7.3

Supersonic molecular jet apparatus. I - FE chamber. OVBP - oil vapor booster pump. N - nozzle. S - skimmer. PMT - photomultiplier tube. Ti -2 inch tubing. T2 - 3 1/2 inch tubing. GV - 3 inch gate valve. II -TOFMS chamber. FT - flight tube. IA - ionization area. DP - diffusion pump. number is a function of downstream distance from the nerror throat and can be expressed as 15

7.5 
$$M_1 = 3.26 \left(\frac{x - 075D}{D}\right)^{7-1} \frac{1}{2} \left(\frac{7+1}{7-1}\right)_{3-26} \left(\frac{x - 075D}{D}\right)^{7-1}$$

for which x is the centerline downstream distance and D is the super sonic nozzle throat diameter. Since the vacuum chamber pressure is finite, acceleration of the expansion will cease at a terminal Mach number which can be determined using Equation 7.4a and<sup>14</sup> 12.000

7.6 
$$\frac{T_1}{T_0} = \left(\frac{P_1}{P_0}\right)^{\frac{\gamma-1}{\gamma}}$$

for which  $P_i$  is the vacuum chamber static pressure. For the continuous supersonic nozzle used in the experiments, typical nozzle backing pressures are around 150 psig. The static pressure in the vacuum chamber at this backing pressure is about 2 x  $10^{-3}$  torr. The nozzle throat diameter is typically 100 microns. Using Equations 7.4a, 7.5 and 7.6, the centerline terminal Mach number for the expansion is calculated to be at  $M_t$  = 36 and is reached at about 4 mm downstream of the nozzle throat.

The supersonic flow properties are described by Equations 7.4-6 until the flow encounters the skimmer or until the flow passes through the Mach disk which is located at a downstream position determined by<sup>15</sup>

7.7 
$$X_{MD} = .67 D \left(\frac{P_0}{P_1}\right)^{-1/2}$$

At the entworpoint consistent open processes colling and contrade the heat the flow and can return it to subsonic speed. In the experiment conducted, the supersonic flow is sampled at nozzle skimmer distances between 20 and 40 mm which is considerably less than the position of the Mach disk calculated to lie at 132 mm downstream using Equation 7.7

Figure 7.4 presents the centerline temperature profiles for supersonic expansions beginning at  $T_0 = 773$  K and  $T_0 = 298$  K as a function of downstream distance x calculated using Equations 7.4 and 7.5 with the centerline terminal Mach number taken at 36. On the basis of the calculations, the temperature profiles for supersonic expansions beginning at elevated temperatures ( $T_0 = 773$  K) are largely the same as those beginning at room temperature ( $T_0 = 298$  K). Thus,  $H_2$ Pc and MgPc should, therefore, be nearly as internally cold before the skimmer as other species observed in the supersonic molecular jet generated by using room temperature expansions assuming that rotational and vibrational temperatures equilibrate near the translational temperature.<sup>15,16</sup> These notions are corroborated experimentally in FE for which ultracold spectra of  $H_2$ Pc, MgPc, and their respective solute/solvent clusters are observed. (The FE spectra are taken between 4 and 8 mm from the nozzle throat, well upstream of the Mach disk.)

Large deviation from isentropic flow may occur at the skimmer which can result in molecular heating. The interferences from the skimmer can be attributed to shock waves forming in the expansion as the flow passes through the skimmer. The physical properties of the supersonic flow after passing through the shock waves can be expressed as a function of upstream conditions and the angle of the shock waves with



1 . .

### Figure 7.4

Downstream distance versus centerline translational temperature for supersonic expansions beginning at 773 K and 298 K.

respect to the flow direction. The centerline downstream translational temperature T<sub>2</sub>, static pressure P<sub>2</sub>, and static density  $\rho_2$  are<sup>14</sup>

7.8a 
$$T_2 = \left(\frac{2\gamma}{\gamma+1} M_1^2 \sin^2 \varepsilon \frac{\gamma-1}{\gamma+1}\right) \left(\frac{\gamma-1}{\gamma+1} + \frac{2}{(\gamma+1)M_1^2 \sin^2 \varepsilon}\right) T_1$$

7.8b 
$$P_2 = \left(\frac{2\gamma}{\gamma+1} M_1^2 \sin^2 \varepsilon - \frac{\gamma-1}{\gamma+1}\right) P_1$$

7.8c 
$$\rho_2 = \left(\frac{(\gamma+1) \ M_1^2 \ \sin^2 \varepsilon}{2 + (\gamma-1) \ M_1^2 \ \sin^2 \varepsilon}\right) \ \rho_1$$

for which  $M_1$  is the Mach number just upstream of the shock wave and  $\varepsilon$  is the flow deflection angle between the shock wave and the flow direction. Downstream/upstream translational temperature ratio versus flow deflection angle for an upstream Mach number of 36 is plotted in Figure 7.5. The plot demonstrates the severe heating effects that can occur in the flow as it passes through the shock wave. The most severe case is at  $\varepsilon = 90^{\circ}$  which corresponds to a normal shock wave. Here, the downstream temperature is about 405 times the upstream temperature. Even at smaller flow deflection angles (oblique shock waves), the heating is significant. For example at  $\varepsilon = 10^{\circ}$ , the downstream temperature is a factor of 13 greater than the upstream temperature. Bier and Hagena have shown, by Schlieren photographs, the types of possible skimmer interferences present in the expansion.<sup>17</sup> At large nozzle/skimmer distances, the shock wave system in the expanding jet gas is similar to that depicted in Figure 7.6a, for which the supersonic flow is contained in oblique and normal shock (Mach disk) waves. In



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#### Figure 7.5

Flow deflection angle versus downstream/upstream centerline translational temperature ratio for flow across a shock wave at an upstream Mach number of 36.



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### Figure 7.6

(d)

F.

Shock wave system in a supersonic molecular jet expansion in the presence of a skimmer downstream of the nozzle throat.

(e)

(f)

this case the scimmer statise lies in the subscript. the normal shock wave other not mall shock vave to any their molecules penetrate the shock wave boundaries. This interference is most severe source of nozzle beam heating since all molecules entry by the skimmer are in subsonic flow and internally hot. As the nozzle is moved towards the skimmer, the shock system in the jet stream is deformed, and the shock waves move towards the nozzle. Figures 7.6b-7.6c. Here, the normal shock wave is still detached from the skimmer orifice and results in "shock beam" formation. On further approach of the nozzle to the skimmer, the normal shock wave attaches to the skimmer orifice, Figures 7.6d-7.6f. The normal shock wave becomes oblique shock waves on both the upstream and downstream sides of the skimmer. The oblique shock wave upstream of the skimmer should not affect the supersonic flow at this nozzle/skimmer distance unless the skimmer mount or chamber walls are in close proximity of the skimmer orifice. Molecules entering the skimmer are in supersonic flow and may be subsequently heated upon traversing the oblique shock waves downstream of the skimmer. The amount of heating which occurs in the downstream oblique shock system depends upon the static pressure behind the skimmer and upon the flow deflection angle.

Molecular heating which occurs in the shock waves can be thought of as a sequence of collisional energy transfer processes up the mole cular vibrational and rotational manifolds. Each elementary step involves the deposition of a small amount of vibrational and rotational energy at the expense of translational energy. Since single ring systems yield ultracold 2-color TOFMS spectra and the macrocycles do not, differences must exist between the collisional energy transfer

element of the shows at or near the skimmer for the momentian systems considered. Two major distinctions can be diffed between the single ring systems and  $H_0Pe$  MgPe which may play a signiti introle in accounting for the difference in molecular heating by the shock system. First, the macrocycles are about 3 times larger in diameter than the single ring systems. Approximating the molecules as hard spheres, one finds that H<sub>2</sub>Pc/MgPc will undergo between 6 and 9 times more collisions in the shock system than the single ring molecules depending upon whether they collide with carrier gas molecules or with other macrocycles. Second,  $H_2Pc$  and MgPc possess a large number of vibrational modes compared to the single ring systems (168 for  $H_2Pc$ versus 30 for benzene); in particular, many of these modes are of low frequency. H<sub>2</sub>Pc has nine ground state vibrational modes with energies below 100 cm<sup>-1</sup>. These low-lying vibrational levels may facilitate heating of the macrocycles. As the large molecules pass through the shock system, the interconversion of small amounts of translational energy into vibrational energy should be more efficient than in the single ring systems for which the lowest ground state vibrational levels lie at about 400 cm<sup>-1</sup>. Either or both of these two factors may contribute to the molecular heating of the macrocycles.

Since the skimmer used in the initial experiments did not perform satisfactorily in maintaining internally cold H<sub>2</sub>Pc MgPc beams, a new skimmer was designed to minimize the shock wave effects on the super sonic flow. Particular attention was focused upon the length of the skimmer, the diameter of the skimmer orifice and the interior and exterior skimming angles.





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EXERCIS DEFENSE

The length of the skimmer is important since the skimmer must be sufficiently long so that the oblique shock wave system does not cause flow perturbations about the skimmer orifice due to skimmer mount or vacuum chamber wall interactions. These interactions tend to degrade supersonic beam skimming by creating a boundary layer at the skimmer mount or chamber walls which rotates the attached shock wave towards a critical angle (more normal to the supersonic flow) which could yield shock wave detachment and the formation of a bow shock wave similar to that shown in Figure 7.6b-7.6c.<sup>17,18</sup>

The diameter of the skimmer orifice dictates the gas throughput of the skimmer.<sup>19</sup> Too large a skimmer diameter will cause TOFMS chamber pumping problems as well as an increase in the intensity of the oblique shock wave system downstream of the skimmer orifice.

The internal skimming angle must be as large as possible to insure that molecules entering the skimmer orifice and striking the skimmer walls can be pumped away fast enough to minimize scattering/ heating of the beam molecules.<sup>19,20</sup> Too small an internal skimming angle will cause skimmer choking and boundary layer formation which results in an increase in the oblique shock wave intensity downstream of the skimmer orifice, similar to that occurring if the skimmer orifice diameter is too large.

The exterior skimming angle determines the upstream flow deflection caused by the skimmer. The angle must be kept small enough so that the oblique shock wave will not detach from the skimmer orifice and destroy the supersonic flow before the molecules pass through the skimmer. The maximum flow deflection angle  $\epsilon_{max}$  which can be negotiated by a supersonic flow without shock wave detachment is given by<sup>14</sup>

7.9 
$$\sin^2 \varepsilon_{\max} = \frac{1}{\gamma M_1^2} \left( \frac{\gamma + 1}{4} M_1^2 - 1 + \left[ (\gamma + 1) \left( 1 + \frac{\gamma - 1}{2} M_1^2 + \frac{\gamma + 1}{16} M_1^4 \right) \right]^{1/2} \right)$$

as a function of Mach number  $M_1$  at the skimmer orifice. The maximum flow deflection angle is related to the maximum exterior skimming angle  $\delta_{\max}$  and skimmer Mach number by  $^{14}$ 

7.10 
$$\frac{1}{\tan \delta_{\max}} = \left(\frac{9(\gamma+1)}{20} \frac{M_1^2}{M_1^2 \sin^2 \varepsilon_{\max} - 1} - 1\right) \tan \varepsilon_{\max}$$

The maximum flow deflection angle and the maximum exterior skimming angle are plotted versus Mach number in Figure 7.7.

From Figure 7.7, the maximum flow deflection angle and the maximum exterior skimming angle approach limiting values at Mach numbers above 6. The high Mach number of the supersonic flow used in the experiments thus makes it possible to use a skimmer with exterior skimming angles of up to  $46^{\circ}$ .  $41^{\circ}$  was selected for the exterior skimming angle used in the new skimmer to provide a safety factor to avoid the occurrence of a normal detached shock upstream of the skimmer orifice. A 35<sup>0</sup> interior skimming angle was selected for the new skimmer as it was the largest feasible angle with respect to the exterior skimming angle that the skimmer could be ground without causing structural problems. The skimmer shape was chosen as conical. The length of the skimmer was selected at 47 mm using the exterior skimming angle and a skimmer base diameter of 35 mm. This length is about a factor of 2 larger than that of the skimmer utilized in the initial TOFMS experiments. This larger skimmer should minimize possible upstream boundary layer and chamber wall effects.<sup>14,19,20</sup> The maximum diameter of the skimmer orifice was selected at 2 mm to avoid TOPMS chamber pumping



## Figure 7.7

Maximum flow deflection angle ( $\epsilon_{max}$ ) maximum skimming angle ( $\delta_{max}$ ) versus skimmer Mach number.

problems. Three skimmers were fabricated possessing the above length, skimming angles, and orifice diameters of .5, 1, and 2 mm in order to determine the optimal skimmer configuration since no theoretical relationship exists between skimmer orifice diameter and supersonic flow characteristics. These skimmers should give an indication of the interference caused by the oblique shock system downstream of the skimmer. The intensities of the shock waves should decrease with decreasing skimmer orifice diameter on the basis of skimmer throughput, boundary layer, and skimmer choking arguments.

2-color TOFMS spectra of  $H_2Pc$  taken using the three skimmers are shown in Figure 7.8. The spectra are taken using  $428^{\circ}C$  nozzle backing region temperature,  $560^{\circ}C$  nozzle tip temperature, 25 mm nozzle/skimmer distance, and 2.5 psig argon nozzle backing pressure. On decreasing the skimmer orifice from 2 to .5 mm, the spectra evolve from being broad and featureless (trace A) to being "quasi" cold (trace C). In both traces B and C, the  $H_2Pc$   $0_0^{\circ}$  and  $A_0^2$  are observed; however, the spectra are smeared out by rotational broadening and vibrational sequence congestion. The sequence structure in trace C is about 40 cm<sup>-1</sup> wide and peaks at about 16 cm<sup>-1</sup> to the red of the  $H_2Pc$   $0_0^{\circ}$ . Similar sequence structure is also observed to the red of the  $A_0^2$ . Assuming a 10% decrease in vibrational frequency between the ground and excited states for  $H_2Pc$ , a vibrational temperature of about 150 K can be estimated from the observed sequence structure.

These experiments identify the skimmer as being the major source of interference in the supersonic expansion. In particular, the observations suggest that the downstream oblique shock system is responsible for the molecular heating since the interference decreases with



## Figure 7.8

2-color TOFMS of  $H_2Pc$  in the vicinity of the  $Q_X$  band  $0_0^0$ . Trace A taken using a 2 mm skimmer orifice. Trace B taken using a 1 mm skimmer orifice. Trace C taken using a .5 mm skimmer orifice. decreasing skimmer orifice diameter. The maintenance of at least some degree of cold internal temperatures in the spectrum taken using the .5 mm orifice skimmer (trace C) demonstrates that the downstream shock wave intensity is still large enough to cause observable skimmer interference in the supersonic flow. This notion is corroborated by experiments using higher backing pressures of argon carrier gas (up to 50 psig) which yield broad spectra that can be attributed to an increase in the downstream skimmer interferences similar to those observed using the 2 mm orifice skimmer.

As mentioned above, the intensity of the downstream oblique shock wave depends upon the static pressure in the vacuum chamber directly downstream of the skimmer. In order to minimize the interference caused by the downstream shock system, the static pressure in this area should be kept lower than the upstream static pressure so that skimmer choking and boundary layer formation does not occur. 14,19,20 In the existing apparatus (Figure 7.3), the region between the skimmer and the TOFMS chamber is constructed using two lengths of 2 inch diameter (4 1/3 inches long) and 3 1/2 inch diameter (4 1/3 inches long) tubing and a 3 inch gate value assembly  $(4 \ 1/3 \ inches \ long)$ . The region directly downstream of the skimmer is displaced by about 70 cm from the TOFMS chamber diffusion pump. The tubing between the two vacuum chambers essentially forms a "dead end" region with a leak provided by the skimmer which may not be adequately evacuated by the TOFMS chamber pumping system. A pressure gradient may thus be present in this region which may give rise to the shock system interference observed.

Other vacuum systems<sup>21</sup> which utilize skimmers to sample the supersonic expansion differ in design from our existing apparatus.

These systems possess two key design features that minimize the effects of the downstream shock waves. First, the vacuum systems are designed for the efficient removal of gas molecules from the post-skimmer region by pumping directly downstream of the skimmer with pumps capable of sustaining static pressures of about  $10^{-6}$  torr. Second, the systems allow for minimization of chamber wall/boundary layer effects in the post-skimmer region by making the chambers large and divergent with respect to the skimmer base. Neither of these features is incorporated in the existing apparatus design.

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A proposed modification to the existing apparatus which incorporates the aforementioned design features is shown in Figure 7.9. The modification involves replacing the 3 inch gate valve, the interchamber tubing, and the 6 inch chamber connecting ports with 12 inch connecting ports and a 16 inch port flange with a divergent hole bored in its center. The 12 inch connecting ports will provide for more than adequate divergence downstream of the skimmer as well as provide for more efficient pumping of the region by the existing TOPMS pumping system. The 16 inch flange serves as both a chamber divider and as a skimmer mounting surface. The skimmer can be mounted in the chamber by clamping its base over the hole in the 16 inch flange with a small flange assembly. For FE experiments, the skimmer can be removed a a sealing plate can be placed over the flange hole to isolate the FE chamber from the TOFMS chamber. The connecting ports can be constructed with different lengths so that the 16 inch flange/skimmer assembly can be located closer to the TOFMS flight tube. This will allow for an overall shorter nozzle/TOFMS ionization area distance which should increase the molecular density at the ionization area by a factor of 2500/X $_{
m 2}$  for which X is the nozzle/TOFMS ionization area distance in cm.



### Figure 7.9

Proposed supersonic molecular jet apparatus. I - FE chamber. OVBP oil vapor booster pump. N - nozzle. S - skimmer. PMT - photomultiplier tube. F - 16 inch Flange. II - TOFMS chamber. FT - flight tube. IA - ionization area. DP - diffusion pump. SUMMARY AND CONCLUSIONS:

Initial attempts to obtain ultracold 2-color TOFMS spectra of H<sub>2</sub>Pc and MgPc yielded negative results. Both helium and argon carrier gases were employed to expand the macrocycles into the vacuum apparatus; however the spectra were significantly broadened. The nontunability of the spectra with varying pump laser frequency was attributed to molecular heating of the H\_Pc and MgPc moleties by shock waves in the vicinity of the skimmer used to sample the supersonic expansion. New skimmers were designed to minimize the shock wave effects on the supersonic flow. "Quasi" cold 2-color TOFMS spectra of H<sub>2</sub>Pc were obtained using these skimmers. Experiments identified the skimmer as being the major source of interference in the molecular expansion. In particular, the observations suggest that a downstream (Chamber II) oblique shock system is responsible for the molecular heating. Through identification of key apparatus design factors which should minimize the downstream shock wave effects, proposed modifications to the existing laboratory apparatus are suggested. The proposed modifications involve enlarging the region downstream of the skimmer to allow for more effective pumping and to minimize skimmer choking and boundary layer effects in the postskimmer region.

#### REFERENCES

- E.B. Wilson Jr., J.C. Decius, and P.C. Cross, "Molecular Vibrations, Theory of Infrared and Raman Vibrational Spectra," (McGraw-Hill Book Co., Inc., 1955).
- 2. J.C. Decius, J. Chem. Phys. <u>17</u>, 1315 (1949).
- 3. B.F. Hoskins and S.A. Mason, Chem. Comm., 554 (1969).
- P.C. Painter and J.L. Koenig, Stectrochimica Acta. <u>33A</u>, 1019 (1977).
- 5. T. Kobayashi, Spectrochimica Acta. 26A, 1313 (1970).
- 6. T. Kobayashi, F. Kurokawa, N. Uyeda, and E. Suito, Spectrochimica Acta. <u>26A</u>, 1305 (1970).
- B. Stymne, F.X. Sauvage, and G. Wettermark, Spectochimica Acta. <u>35A</u>, 1195 (1979).
- P.S.H. Fitch, C.A. Hayman, and D.H. Levy, J. Chem. Phys. <u>73</u> 1064 (1980).
- 9. The PED's are calculated by using % PED =  $100 \times |C_i|^2$  for which  $C_i$  is the eigenvector coefficient for symmetry coordinate i in the eigenvector normal modes.
- J. Wanna and E.R. Bernstein, J. Chem. Phys. <u>85</u>, 1795 (1986); J.
   Wanna, J.A. Menapace and E.R. Bernstein, J. Chem. Phys. <u>85</u>, 1795 (1986).
- E.R. Bernstein, K. Law and M. Schauer, J. Chem. Phys. <u>80</u>, 634 (1984); M. Schauer and E.R. Bernstein, J. Chem. Phys. <u>82</u>, 726 (1985).
- 12. M. Schauer, Ph.D. Thesis, Colorado State University, 1984. E.R. Bernstein, K. Law and M. Schauer, J. Chem. Phys. 80, 207 (1984).
- The skimmer used in the experiments is a model 1 skimmer purchased from Beam Dynamics, 623 E. 57th Street, Minneapolis, Minnesota 55417.
- W.C. Reynolds and H.C. Perkins, "Engineering Thermodynamics," (McGraw-Hill Book Co., Inc., New York, 1977); M.J. Zucrow and J.D. Hoffman, "Gas Dynamics," Vol. I (J. Wiley and Sons, Inc., New York, 1976).

- P.P. Wegener, "Molecular Beams and Low Density Gas Dynamics," (Marcel Dekker, Inc., New York, 1974).
- 16. A. Amirav, U. Even, J. Jortner, Chem. Phys. 51, 31 (1980).
- 17. K. Bier and O. Hegena, "Rarefied Gas Dynamics," edited by J.A. Laurmann (Academic Press, New York) Vol. 1, p. 478-496, 1963.
- 18. U. Bossel, Entropic <u>30</u>, 11 (1969).

•

- 19. A. Kantowitz and J. Grey, Rev. of Scientific Instruments <u>22</u>, 328 (1957).
- 20. K. Bier and O. Hegena, "Rarefied Gas Dynamics," edited by J.H. el Leeuw (Academic Press, New York) Vol. II, p. 260-278, 1966.
- 21. M.G. Liverman, S.M. Beck, D.L. Monts, and R.E. Smalley, "Rarefied Gas Dynamics" edited by R. Campargue (Academic Press, New York) Vol. II, p. 1037-1048, 1979; D.H. Levy, L. Wharton, and R.E. Smalley, "Laser Spectroscopy in Supersonic Jets," in <u>Chemical and Biological Applications of Lasers</u>, Vol. II, edited by C. Bradley Moore (Academic Press, New York) 1977; J.B. Fenn and J. Deckers, "Rarefied Gas Dynamics," edited by J.A. Laurmann (Academic Press, New York) Vol. I, p. 497-515, 1963; J.G. SkoFronick, Review of Scientific Instruments, <u>38</u> 1628, 1967.

#### CHAPTER EIGHT

#### FUTURE EXPERIMENTS AND CONCLUSIONS

This chapter presents the basic ideas and motivations for future experiments which can be conducted using the techniques and experimental set-ups currently available in the Bernstein group laboratory. The proposed studies entail those experiments that will extend or complement the work presented in this dissertation. This chapter also summarizes the conclusions drawn from the work accomplished on the molecules and clusters presented in Chapters 2-7. The conclusions include the most important results obtained from the work. For specific conclusions pertaining to each of the systems studies, the appropriate chapters in the dissertation should be consulted.

Future Experiments.

Effects of Macrocycle Ring Substitution on Solvation - The high temperature supersonic molecular jet techniques utilized to study  $H_2Pc$ . MgPc, and their respective vdW clusters with small solvents (Chapter 6) also gives us the opportunity to study the effects of ring substitution in the macrocycles on their solvation properties in an ultracold iso lated environment. For example in the  $H_2Pc$  and MgPc cluster studies, major differences exist between the two cluster solutes in that the  $H_2Pc$ clusters exhibit enhanced stabilization in the  $S_1$  state relative to the  $S_0$  state whereas the MgPc clusters exhibit enhanced stabilization in the  $S_0$  state relative to the  $S_1$  state. This difference appears to be due to the magnesium substitution in the phthalocyano core. Furthermore, the spectra suggest that the intermolecular interactions between MgPc and solvents having lone electron pairs possess major contributions from the central magnesium atom and the solvent electron pairs.

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These observations pose several questions pertaining to the physics and chemistry responsible for the solvation properties of the phthalocyanines (Pc's). Questions that can be considered include: 1) are the solvation properties of Pc's dependent upon the specific metal atom present in the phthalocyano core; 2) does the relative stabilization of one electronic state over another depend upon the specific central metal atom; 3) do cluster geometries and solute solvation sites change significantly with different central metal atoms; and 4) are the central metal atoms in the Pc moieties responsible for a major portion of the intermolecular interaction.

To explore the effects of central metal atom substitution in Pc's, initial experiments can be conducted on zinc phthalocyanine (ZnPc). The fluorescence quantum yield for ZnPc is about .3 which is comparable to those for  $H_2Pc$  (.7) and MgPc (.6).<sup>1</sup> This quantum yield is large enough to allow for observation of ZnPc and its respective vdW clusters in fluorescence excitation experiments. The experiments could initially entail clustering ZnPc with solvents such as those used in the  $H_2Pc$  and MgPc studies. By comparing the ZnPc cluster spectra and calculated cluster geometries with those for the MgPc cluster systems, the effects of the central metal atom on cluster properties can be explored. Subsequent experiments in this area could include spectroscopic and modeling studies on Pc's containing cadmium, sodium, and lithium.

Another set of experiments which can be conducted to study the effects of ring substitution on the Pc solvation properties involves the analysis of a tetrabenzoporphyrin (TBP) solute series clustered with various solvents. Macrocycles which can be initially studied in this class include free base tetrabenzoporphyrin (H<sub>2</sub>TBP), magnesium tetrabenzoporphyrin (MgTBP), and zinc tetrabenzoporphyrin<sup>2</sup> (ZnTBP). This class of compounds offers the opportunity to study the effects of the solute bridge nitrogens on the solvation properties of the Pc class as H<sub>2</sub>TBP, MgTBP, and ZnTBP differ from their respective Pc analogues only by replacement of the bridge nitrogens in Pc by methine bridges in TBP. The TBP compounds proposed for study also possess fluorescence quantum yields (.46 for  $H_2$ TBP, .55 for MgTBP, and .35 for ZnTBP)<sup>1</sup> which are favorable for observation in a supersonic molecular jet fluorescence excitation experiment. Through the analysis of the cluster spectral shifts, chromophore vibronic structures, and calculational modeling of the cluster geometries and binding energies, the effects of the bridge nitrogens on the macrocycle solvation properties may be established. Cluster studies using the TBP solutes are interesting and important as they will provide the link between the Pc's and their biologically active analogues, the porphyrins, in regard to cluster geometry, binding energy, preferred solute solvation sites, and major interaction types responsible for solvation.

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<u>Aromatic Solute/Aromatic Solvent Clusters</u> - All the aforementioned Pc and TBP solutes can be clustered with a number of small aromatic solvents such as benzene, pyrazine, pyrimidine, and pyridine. Studying these clusters gives us the opportunity to explore cluster

energetics and dynamics from the point of view of both the cluster solute and solvent. Supersonic molecular jet studies can be conducted by probing both the Pc/TBP solute and the benzene/N-heterocycle solvent optical transitions. The studies will yield insight into the importance of the solvent aromatic  $\pi$  cloud and/or nonbonded electron pairs on solvation. For example, clustering MgPc with pyrazine should provide detailed information on the participation of the solvent nonbonded electron pairs with the magnesium central metal atom. If the solvent nonbonded electron pairs are major contributors in cluster formation/ stabilization, the  $n\pi^*$  transition in pyrazine should not exist or be highly perturbed (blue shifted). The cluster systems are also interesting in the respect that they may allow one to determine if changes in the cluster binding energy upon electronic excitation depend on the specific electronic excitation analyzed. In particular, one could ask the question: will the change in cluster binding energy be different in magnitude if the cluster solvent optical transition is probed rather than the cluster solute optical transition? Furthermore, these studies may yield information on differences in the relative stabilization of the solute and solvent electronic states. Do the spectral shifts observed when probing the solvent optical spectrum differ in sign than those observed when probing the cluster solute spectrum; that is, are the spectral shifts both bathychromic or hypsochromic or is the solvent spectral shift bathychromic and the solute shift hypsochromic (or vice versa)?

<u>Phthalocyanine and Porphyrin Vibronic Structure</u> - Up to this point, we have only considered analyzing the Pc and TBP macrocycles in cluster systems. The vibrational spectroscopy of these molecules is

also important in the understanding of the vibrational structure present in biologically significant systems. In particular, elucidation of the vibrational structure in the Pc and TBP systems may yield valuable information regarding which portions of the macrocycles participate in the various vibrational motions. Supersonic molecular jet studies on isolated Pc and TBP can contribute to the resolution of the excited state energetics present in the systems since the techniques greatly simplify the vibronic spectra and thus facilitate their interpretation. As discussed in Chapter 6, the low frequency motion present in the H\_Pc and MgPc vibronic spectra is characterized using the results of an outof-plane normal coordinate analysis; and in Chapter 7, the calculations on H<sub>2</sub>Pc are compared to infrared data. These studies make it possible for us to determine the detailed nature of the out-of-plane motion occurring in specific regions within the H<sub>2</sub>Pc vibrational manifold. In order to obtain a complete understanding of the macrocycles vibrational structure, the in-plane motion should be characterized. The vibronic spectra of the Pc and TBP molecules presented above can be analyzed using the results of both in-plane and out-of-plane normal coordinate analyses. In these studies, the in-plane motion can be modeled using a refined force field determined from normal coordinate analyses on the in-plane motion occurring in porphyrins. $^{3-7}$  The combination of calculational modeling and spectroscopic studies on these systems should increase our understanding of the structural changes (nuclear motion) occurring in the various electronic states as well as the extent of interelectronic state mixing (vibronic coupling) between the different electronic states in the macrocycles. These studies will also be useful in the basic understanding of the spectroscopy in more complex systems such as substituted Pc's and porphyrins.

 $\underline{z} = \underline{z}$  Solute/Solvent Interaction in Clusters - The MgPc(CO<sub>2</sub>)<sub>1</sub> and H<sub>2</sub>Pc(CO<sub>2</sub>)<sub>1</sub> clusters presented in Chapter 6 exhibit vibronic spectra with distinctive spectral shifts. The  $H_2Pc(CO_2)_1$  cluster is the only cluster observed with H<sub>2</sub>Pc which possesses a hypsochromic shift upon excitation to the  $S_1$  state. The MgPc(CO<sub>2</sub>)<sub>1</sub> cluster also exhibits a hypsochromic shift comparable to that observed in  $H_2Pc(CO_2)_1$ . The exact nature of the intermolecular interaction responsible for the spectral shifts is not known; however, based upon cluster experiments with alcohol and hydrocarbon solvents, it is possible to speculate that  $\pi$ - $\pi$ solute/solvent interactions may be responsible for the observed spectral shifts. To study this behavior, a series of experiments on single ring solutes such as benzene, pyrazine, and pyrimidine solvated by carbon dioxide, carbon monoxide, and oxygen can be conducted using 2-color TOFMS techniques. The benzene/solvent vibronic spectra will reveal whether the solvents orient perpendicular or parallel to the solute molecular plane through the observation of forbidden cluster chromophore transitions. In these studies, the spectral shifts may be related to particular cluster geometries which may yield information on the major interactions contributing to the intermolecular potentials between the solutes and solvents. The pyrazine and pyrimidine/solvent clusters may tell us about the effects of the ring nitrogens on the cluster geometry. binding energy, and solute solvation. Computer modeling of cluster geometry, binding energy, and intermolecular vibrational structure may also aid in the understanding of the intricacies of the intermolecular interactions present in these systems.

<u>vdW Vibrational Structure in N-Heterocycle Dimers and Solute</u> <u>Solvent Clusters</u> - Calculation of the intermolecular vibrational struc ture for the N-heterocycle dimers and solute/solvent clusters discussed

in Chapters 4 and 5 can also be conducted. The modeling studies of the N-heterocycle solute/solvent clusters are interesting since they may show the effects of the solute nitrogen atoms on the intermolecular vibrational structure when compared to the analogous benzene solute/solvent clusters. Furthermore, the studies are important as little to no intermolecular vibronic structure is observed in the cluster spectra. In this regard the studies will yield predictions of the frequencies of those vdW modes not observed in the spectra as well as aid in assigning the weak features present in some of the spectra. Intermolecular vibrational structure modeling of the N-heterocycle dimer systems could serve as a major contribution to the understanding of their spectroscopy. The studies may be useful in the determination/ confirmation of the dimer geometries responsible for the observed spectra as well as help to elucidate the nature of the intermolecular energetics and dynamics present in the systems. Of particular interest is the elucidation of the vdW torsional structure present in the planar dimers, especially for the vdW torsional coordinate in which the cluster subunits rotate/librate about an axis joining their centers-of-mass. This motion is similar to torsional motion occurring in polyring systems such as biphenyl and biphridyl. Studying the torsional motion in the clusters may yield information concerning the long-range ( $\pi$ -electron effects) and short-range (hydrogen-hydrogen repulsions) interactions present between the cluster subunits which may be considered precursors to the interactions responsible for the potential well shape in the polyring systems. The studies will also contribute to the understanding of hydrogen bonding interactions in the cluster systems and how it affects cluster geometry and intermolecular energetics.

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Conclusions.

Supersonic molecular jet expansion, TOFMS, and FE spectroscopic techniques have been employed to study the gas phase optical spectra of a wide array of weakly bound vdW clusters in an ultracold isolated molecular environment. The techniques have proved useful in studying the vdW clusters of aromatic molecules ranging from benzene and its N-heterocyclic analogues to macrocycles such as H\_PC and MgPc solvated by small hydrocarbons/fluorocarbons, small alcohols, water, ammonia, argon, and carbon dioxide. The studies reveal detailed information regarding the geometries/symmetries, the intermolecular energetics, and the physical nature of the vdW interactions present in the cluster Spectroscopic observables, such as cluster spectral shifts, systems. forbidden cluster chromophore vibronic transitions, vdW motions, and cluster ionization energies, combined with computer modeling of cluster ground state binding energies, geometries, and intermolecular energetics proves essential in the analysis and understanding of the physics involved in cluster formation, stabilization, and behavior.

The experimental and theoretical studies of the benzene/solvent clusters reveal detailed information regarding cluster geometries, symmetries, and the nature of the intermolecular vibronic motion present in the systems. In these studies, we found that the preferred benzene solvation site is located over the aromatic  $\pi$  cloud where solute'solvent vdW interaction is maximal. Through computer modeling, we have demonstrated the correlation between cluster geometry and spectral features. Specifically, we showed that the calculations yield useful information regarding cluster geometry, binding energy, and intermolecular vibrational structure. These calculated results are consistent with

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experiment and can be used to elucidate the details of vdW cluster energetics. By comparing the calculations and the experiments, we are able to assign cluster vibronic transitions to specific cluster geometries and to specific vdW vibrations.

Detailed experimental and theoretical studies of the intermolecular torsional structure in the benzene/solvent clusters show that the clusters are rigid systems with regard to internal rotation of the cluster subunits. In this regard, the clusters possess unique equilibrium grometries for which the intermolecular motion is oscillatory. The motion is not an admixture of vdW bends, stretches, and free internal rotations as would occur if the clusters were internally nonrigid.

The N-heterocycle/solvent cluster studies demonstrate that the solute ring nitrogen atoms influence, to some extent, both the cluster geometry and spectroscopy. In the N-heterocycle/hydrocarbon clusters, we find that the presence of the nitrogen atoms in the aromatic rings has a small but discernible effect on the intermolecular interaction. In particular, the solvent hydrogen atoms preferentially orient towards the solvent nitrogen atoms. The overall behavior of these clusters with regard to geometry, binding energy, and spectroscopy, however, is very similar to that found in the benzene/hydrocarbon clusters.

The N-heterocycle/water and ammonia clusters, on the other hand, clearly demonstrate major interactions between the ring nitrogens and the solvent hydrogens. These studies suggest that hydrogen bonding is present between the ring nitrogens and the solvent hydrogens as evidenced by the hypsochromic spectral shifts present in the cluster spectra.

The single ring/solvent clusters also exhibit spectra and calculated geometries consistent with the notion that the intermolecular potential is primarily composed of solute/solvent  $\pi$  cloud interactions. In this regard, the preferred solvation sites on the cluster solutes are located over the aromatic  $\pi$  clouds. Thus, the cluster geometries are such that the solvents lie over the center of the solutes unless other large intermolecular interactions are present which may alter this behavior. In particular, hydrogen bonding interactions present in the N-heterocycle and benzene/solvent systems appear to cause geometrical changes in the clusters. The changes range from simple reorientation of the solvents to minimize solute ring nitrogen/solvent hydrogen distance or solvent hydrogen/solute  $\pi$  cloud distance to large reorganization of the cluster geometry from a "sitting atop" configuration to planar hydrogen bonded geometries as suggested by pyrimidine/ammonia and N-heterocycle dimer studies.

The spectroscopic studies of phthalocyanines and their respective vdW clusters with small molecules take advantage of the characteristics of the supersonic molecular jet. This technique makes it possible to simplify the complicated phthalocyanine spectra and to control the solvent and environmental effects on the chromophores in a set and controlled manner. The interplay of spectroscopy and an out-of-plane normal coordinate analysis enables us to characterize the low-lying outof-plane motion present in  $H_2Pc$  and MgPc. The correlation between the spectra and the calculations is excellent which allows for the elucidation of the motion types observed in the first few hundred wavenumbers of the  $H_2Pc/MgPc$   $S_1$  manifolds. The correlation also makes possible the identification of specific species/geometries responsible for the

cluster vibronic spectra observed in the vicinities of the chromophore Q band origins.

In the cluster studies, we find that the  $H_2Pc/hydrocarbon$  cluster experimental and theoretical results parallel those observed for the benzene and N-heterocycle/hydrocarbon clusters. The cluster geometries favor forms for which solvent/solute  $\pi$  cloud overlap is maximal. The spectral shifts are all bathychromic and the magnitudes of the shifts increase with increasing solvent size and polarizability. The  $H_2Pc$  and MgPc/water and /alcohol clusters exhibit spectra and calculated geometries which demonstrate that the solvent OH groups are large contributors to the spectral shifts and intermolecular interactions. In all the cluster systems studies, the preferred  $H_2Pc$  and MgPc solvation sites are located over the phthalocyano core. As in the benzene/water and benzene/ammonia clusters, the  $H_2Pc$  and MgPc clusters exhibit reduced symmetry with respect to the isolated chromophores. This reduction in symmetry is demonstrated by the observation of forbidden chromophore transitions and  $S_1$  state splitting in the clusters.

Overall, computer modeling proved to be an essential component of the studies on both the isolated molecules and the cluster systems. The spectroscopic data obtained are necessary to validate and refine the models used, and the calculations help to assign and understand spectra which are otherwise difficult, if not impossible, to uniquely interpret. In this way, the combination of spectroscopy and computer modeling leads to an increased understanding of the behavior of the molecules and clusters over what could be obtained by either study alone.

## REFERENCES

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- Martin Gouterman, Optical Spectra and Electronic Structure of Porphyrins and Related Rings, in "The Porphyrins" edited by David Dolphin (Academic Press, New York) Vol. III, 1978.
- 2. Uzi Even, Jacob Magen, and Joshua Jortner, J. Chem. Phys. <u>77</u>, 4384 (1982).
- 3. S. Sunder and H.J. Bernstein, J. of Raman Spectros. 5, 351 (1976).
- 4. M. Abe, T. Kitagawa, and Y. Kyogoku, J. Chem. Phys. <u>69</u>, 4526 (1978).
- 5. C.A. Melandres and V.A. Maroni, J. of Raman Spectros. <u>15</u>, 319 (1984).
- L.L. Gladkov and K.N. Solovyov, Spectrochimica Acta. <u>41</u>, 1437 (1985); <u>41</u>, 1443 (1985); <u>42</u>, 1 (1986).
- 7. L.L. Gladkov, A.T. Gradyushko, A.M. Shulga, K.N. Solovyov, and A.S. Starukhin, J. of Molecular Structure <u>47</u>, 463 (1978).

## APPENDIX ONE

HYDROGEN BONDED AND NON-HYDROGEN BONDED VAN DER WAALS CLUSTERS: COMPARISON BETWEEN CLUSTERS OF PYRAZINE, PYRIMIDINE, AND BENZENE WITH VARIOUS SOLVENTS

# Hydrogen bonded and non-hydrogen bonded van der Waals clusters: Comparison between clusters of pyrazine, pyrimidine, and benzene with various solvents <sup>a)</sup>

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Solute-solvent clusters of pyrazine, pyrimidine, and benzene (solutes) and  $C_n H_{2n+2}$  (n = 1,2), NH<sub>3</sub>, and H<sub>2</sub>O (solvents) are studied by the techniques of supersonic molecular jet spectroscopy and two-color time-of-flight mass spectroscopy (two-color TOFMS). Spectral shifts, van der Waals (vdW) modes, dissociation energies, and vdW mode-solute mode vibronic couplings are characterized for most of the observed clusters. Based on these data and previous results for hydrocarbon systems, cluster geometries can be suggested. Lennard-Jones potential (6-12-1) calculations are also performed for these clusters and in all instances for which comparisons can be readily made, calculated and experimentally estimated geometries and binding energies agree completely. Clusters of N-heterocyclic solutes and H<sub>2</sub>O are not observed experimentally. Systematics and trends among the clusters reported herein and those previously reported are discussed and analyzed.

## I. INTRODUCTION

Supersonic molecular jet spectroscopy has made possible the study of a wide variety of weakly bound, solute-solvent van der Waals (vdW) clusters in the gas phase. Cluster investigations have enhanced our understanding of intraand intermolecular interactions and potentials, vibrational energy dynamics and chemical reactions, structural properties of small aggregates of solute and solvent molecules, and nucleation and growth of small clusters. Clusters can also be considered as model systems for condensed phase behavior. Moreover, these vdW systems can be thought of as an important new state of matter in which the static and dynamic properties of small aggregates of weakly coupled molecules can be studied.

vdW clusters, after being produced in a supersonic jet expansion, can be probed by three distinct techniques: fluorescence excitation (FE), dispersed emission (DE), and two-color time-of-flight mass spectroscopy (two-color TOFMS). The latter technique is employed most often in our studies of clusters because it gives unique cluster identification, brackets the cluster binding energies, and elucidates cluster vibrational energy dynamics and vibrational predissociation.

In the past few years, we have reported several studies of vdW clusters using the three spectroscopic techniques mentioned above.<sup>1-7</sup> Cluster geometry, binding energy, nucleation and growth dynamics, and limits on the vibrational energy dynamics and vibrational predissociation times have been determined. The vdW solute-solvent clusters investigated initially are for the most part restricted to aromatic hydrocarbon solutes (e.g., benzene and toluene) and small alkane solvents ( $CH_4$ , $C_2H_6$ , and  $C_3H_8$ ) in which only one type of interaction, that is one potential form, is found to be important for the solute-solvent coordination. Expanding on these previous studies we are now exploring solute-solvent clusters with N-heterocyclic solutes (e.g., pyrazine and pyrimidine) and alkane solvents and N-heterocyclic solutes with hydrogen bonding solvents (e.g., water and ammonia). The initial report of this effort for pyrazine and methane, ethane and propane clusters has already appeared.<sup>7</sup>

In this paper we discuss the two-color TOFMS study of pyrimidine clustered with CH4 and C2H6, pyrazine and pyrimidine clustered with NH<sub>3</sub>, and benzene clusters with H<sub>2</sub>O and NH<sub>3</sub>. The pyrimidine-alkane clusters are presented for comparison with the previously published' pyrazine-alkane data: The effect of the ring nitrogen atoms on the cluster geometry can thereby be evaluated. The pyrazine and pyrimidine ammonia clusters reveal the role of hydrogen bonding interactions in simple clusters. Benzene-water and -ammonia clusters serve as an example of clusters with these more complicated solvent systems (i.e., two possible interaction potentials) interacting with aromatic hydrocarbons. Theoretical and experimental studies of such a series of systems should eventually lead to a fuller understanding of solute-solvent coordination structure, dynamics, and the hydrogen bonding interaction.

Although extensive efforts were made and a wide variety of experimental conditions explored, pyrazine and pyrimidine clusters with water were not observed. Both FE and two-color TOFMS detection techniques were employed. A broad feature (roughly 50 cm<sup>-1</sup> FWHM) was observed in FE at 580 cm<sup>-1</sup> to the blue of pyrazine  $0_0^0$  transition for pyrazine and water expanded with helium. No signal, however, was observed at the pyrazine water mass channel for two-color TOFMS. We must thus conclude that the pyrazine and the pyrimidine water clusters have not been a bserved. Either the excited  $n\pi^0$  state of the cluster is dissociative or internal conversion or intersystem crossing is so rapid for these clusters that the lifetime of the  $n\pi^0$  state is greatly reduced (~0.1 ps). Pyrimidine-fluorinated alcohol clusters J Chem Phys. 85 (4), 15 August 1986

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also exhibit broad features and reduced lifetimes.8

Hydrogen bonding interactions are known to play an important role in intra- and intermolecular interactions important for secondary and tertiary molecular structure, molecular dynamics, and ionic and molecular solvation. Hydrogen bonding interactions have received attention in previous supersonic molecular jet spectroscopic studies: These systems include phenol clustered with various proton accepting molecules,<sup>9</sup> indole and 2-aminopyridine clustered with various solvents,<sup>10</sup> benzoic acid dimers,<sup>11</sup> s-tetrazine dimers,<sup>12</sup> 1,4-dihydroxyanthraquinone,<sup>13</sup> and methylsalicylate.<sup>14</sup> In most instances blue-shifted cluster spectra are found due to the cluster stabilization of the ground state and destabilization of the excited state.<sup>15</sup>

In this and our other cluster studies, the experimental data are supplemented by potential energy Lennard-Jones (LJ) atom-atom (6-12-1) calculations of cluster structure, binding energy, and internal motion. The calculations and the potential are discussed thoroughly in our previous studies.<sup>7</sup> The LJ calculations produce identical geometries but slightly lower binding energies than the exp-6 potential form used in earlier calculations.<sup>1-6</sup> The LJ hydrogen bonding (LJ-HB) potential form proves to be more versatile than the exp-6 form since many more constants for different types of atom-atom interactions have been independently reported for the LJ potential.<sup>16</sup> None of the potential parameters employed in this work is fit to the cluster data.

As is well known and widely accepted, geometries of isolated molecules and clusters are best obtained through spectroscopic observation of rotational structure. In fact, rotational structure of a number of simple clusters has been observed under molecular jet conditions: s-tetrazine and iodine with He and Ar<sup>17</sup> and aniline with Ne and Ar.<sup>18</sup> The resolution available to us at present is 0.08  $\text{cm}^{-1}$ ; at this resolution only rotational envelopes are observable which do not lend themselves readily to a unique interpretation of cluster structure. We calculate that 0.005 cm<sup>-1</sup> resolution would be required to resolve rotational structure for  $C_4H_4N_2(NH_3)_1$  etc., under the restriction of a rigid geometry. We are thus for the present forced to employ less direct methods to obtain cluster geometry. Assignment of the spectra is accomplished through the determination of ionization energies, spectral shifts, relative intensities, (molecular) symmetry forbidden cluster transitions, and potential calculations. The understanding of these more complex systems rests heavily on the previous data obtained for other clusters.<sup>1-7</sup> For all systems discussed in this paper, complete agreement between spectroscopic data, calculations, and results for previously analyzed solute-solvent clusters is found.

#### **II. EXPERIMENTAL PROCEDURES**

The experimental apparatus and procedures are similar to those used previously for the study of vdW clusters. The vacuum system consists of two chambers with a pulsed nozzle and mass detection system in the second chamber. A skimmer separates the pulsed nozzle and the time-of-flight mass spectrometer. The first chamber contains either a pulsed or cw nozzle, the molecular beam from which can be taken into the second chamber through a skimmer. FE and DE experiments are carried out in the first chamber.

The two independent lasers used in the two-color TOFMS experiments are Nd<sup>+1</sup>/YAG pulsed lasers the doubled output of which pumps two dye lasers. The dye laser output can be mixed with the 1.064 $\mu$ m Nd<sup>+1</sup>/YAG fundamental, frequency mixed and doubled, or just doubled using various nonlinear KDP crystals. The laser output can be extended from greater than 4.5 to  $\sim 0.215 \,\mu$ m. One laser is employed to excite the cluster to its first excited  $n\pi^{\bullet}$  or  $\pi\pi^{\bullet}$  electronic state and the second laser then ionizes this cluster starting from the  $S_1$  vibronic manifold. The maximum ionization energy achievable with this second laser is roughly 46 500 cm<sup>-3</sup>.

The solute or cluster chromophore is typically placed in an in-line trap or filter cup directly behind the pulsed valve. Water is placed in a trap before the valve and solute; the helium carrier gas passes over both materials and into the valve. Gaseous solvents are premixed with the helium carrier gas in a holding tank ( $\sim 2000$  psi) at concentrations varying from 2.0 to 0.1 mol/mol%.

The LJ potential function (6-12-1) with the additional HB form is described in detail in a previous publication.<sup>7</sup> Table I contains a list of the previously unreported constants employed with this potential form. Pyrazine and pyrimidine structures used in the calculations are obtained from Ref. 19.

TABLE I. Parameters for the energy expression in the computer modeling.

	$E_{q} = 1.16 \times 10^{5} q_{i}q_{j}/2r_{q} + A_{q}/r_{q}^{12} - C_{q}/r_{q}^{0} (LJ)$					
		$A\left(\frac{\mathrm{cm}^{-1}\mathbf{\hat{A}}^{12}}{\mathrm{mol}}\right)$	$C\left(\frac{\operatorname{cm}^{-\epsilon}\dot{\mathbf{A}}^{\bullet}}{\operatorname{mol}}\right)$			
Amine	aromatic					
	N-N	1.312×10	1.403×10 <sup>9</sup>			
	N-C	1.728×10*	1 575 × 10 <sup>3</sup>			
	N-H	$2.523 \times 10^{\prime}$	4 527 × 10°			
	H-C	2.749 × 10'	5 217 × 10"			
	н-н	3.872 × 10 <sup>e</sup>	1 590 × 10 <sup>4</sup>			
	H-N	1 990×10'	4 527 × 10 <sup>4</sup>			
Water-s	romatic					
	0-N	7 548 × 10'	1 021 × 10 <sup>3</sup>			
	<u>о-с</u>	9 868 × 10'	1 130 × 10 <sup>3</sup>			
	0-H	1.363 × 10'	3.162 × 10 <sup>4</sup>			
	H-C	$3.160 \times 10^{7}$	5 217 × 10 <sup>4</sup>			
	H-H	4 537 × 10 <sup>e</sup>	1 590 × 10 <sup>4</sup>			
	H-N	$2.297 \times 10^{7}$	4 527 < 10 <sup>4</sup>			
		$E_{y} = 1.16 \times 10^{3}  q$	$(q_{i}/2r_{y} + A^{2}/r_{y}^{12} - B/r_{u}^{10})$ (HB)			
		$\mathcal{A}\left(\frac{cm^{-1}\mathbf{\dot{A}}^{12}}{moi}\right)$	$\boldsymbol{B}\left(\frac{cm^{-1}\mathbf{\dot{A}}^{10}}{mol}\right)$			
H N		1.150×10 <sup>7</sup>	2 882 × 10 <sup>e</sup>			
		9				
NH,	N	- 0 4 38				
	н	+ 0 1 <b>46</b>				
H <sub>2</sub> O:	0	- 0 34				
	н	+017				
C,H,N,	<sub>1</sub> : N	- 0 18 <b>2</b>				
	С	+ 0 0 <b>91</b>				
	н	0				
C,H,	С	- 0 0074				
	н	+ 0.0074				

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	N-H	2.523 × 10'	4.527×10 <sup>4</sup>		
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	H-N	1.990 × 107	4.527×10 <sup>4</sup>		
Water-a	romatic				
	0-N	7.548×10 <sup>7</sup>	1 021 × 10 <sup>3</sup>		
	о-с	9.868 × 10 <sup>7</sup>	1.130×10 <sup>4</sup>		
	0-н	1.363 × 10'	3.162×10*		
	H-C	3.160×107	5.217×104		
	H-H	4.537×10 <sup>e</sup>	1.590×10 <sup>4</sup>		
	H-N	2.297 × 107	4.527 × 104		
		$E_q = 1.16 \times 10^3  q$	$r_{i}q_{j}/2r_{y} + A'/r_{y}^{12} - B/r_{y}^{10}$ (HB)		
		$\mathcal{A}\left(\frac{\operatorname{cm}^{-1}\hat{\mathbf{A}}^{12}}{\operatorname{mol}}\right)$	$B\left(\frac{\operatorname{cm}^{-1}\dot{\mathbf{A}}^{10}}{\operatorname{mol}}\right)$		
H N	·	1.150×107	2.882 × 10*		
		9			
NH <sub>3</sub> :	N	0.438			
	H	+ 0.140			
H <sub>2</sub> O:	0	- 0.34			
	H N	+ 0.17			
C'H'N'	, N	- 0.182			
	C	+ 0.091			
~	н	0			
C <sub>6</sub> H <sub>6</sub> :	C	- 0.0074			
	H	+ 0.0074			

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#### III. RESULTS

This section contains the experimental and calculational results for the various clusters investigated. We first present pyrimidine with methane and ethane for comparison with the previously reported pyrazine clusters. Pyrimidine and pyrazine ammonia clusters are then discussed and benzene water and ammonia clusters are presented for a comparison with the .V-heterocyclic systems. Based on previous experience with a number of different clusters,<sup>1-7</sup> we have not made an extensive experimental study of the binding energies of these clusters. We rely on the calculations which have always fallen within the range bracketed by the experimental two-color TOFMS data. Preliminary checks made on both pyrazine and pyrimidine clusters are in agreement with the calculations. Vibrational vdW modes observed in the spectra will not be assigned in this publication. Future publications will assign them as bends, stretches, torsions, and combinations utilizing a normal coordinate calculational analysis.20

## A. Pyrimidine-methane

Figure 1 and Table II present the data for the pyrimidine  $(CH_4)_1$  and  $(CH_4)_2$  complexes taken near the pyrimidine  $0_0^0$ transition (31 073.0 cm<sup>-1</sup>). The cluster of pyrimidine  $(CH_a)_1$  has a spectral shift of -56.6 cm<sup>-1</sup> and a low frequency mode at 4.5 cm<sup>-1</sup> from this origin. These spectra are obtained by two-color TOFMS. Hints of other vdW modes can also be seen in the trace in Fig. 1 but we are hesitant to report such weak transitions. Pyrimidine (CH<sub>4</sub>)<sub>2</sub> spectra clearly show two clusters, similar to previously reported clusters for other aromatic systems.<sup>1-7</sup> The feature at -112.1 cm<sup>-4</sup> in Fig. 1 is assigned as the 0<sup>6</sup><sub>6</sub> transition of the isotropic (symmetric), additive shift cluster and the feature at -47.2 cm<sup>-1</sup> is attributed to the  $0_0^0$  transition of the anisotropic (asymmetric) cluster with both methanes on the same side of the aromatic ring. Note that without both mass and energy resolution, the spectra of pyrimidine (CH<sub>4</sub>)<sub>1</sub> and (CH<sub>4</sub>)<sub>2</sub> would not be resolved and the clusters could not be separated and uniquely identified. A vdW mode at 5.1 cm ~ from the isotropic cluster origin is observed.

Potential energy calculations using LJ potentials for these clusters generate geometries and binding energies comparable to those previously reported for other aromatic-al-



FIG. 1. Two-color time-of-flight mass spectra (two-color TOFMS) of pyrimidine (CH<sub>4</sub>)<sub>1</sub> and pyrimidine (CH<sub>4</sub>)<sub>2</sub> in the region of the pyrimidine origin (31.073.0 cm<sup>-1</sup>).

kane systems and in complete accord with the above experimental findings (see Fig. 2). Calculations for pyrimidine (CH4) clusters yield only one geometry for which the methane is coordinated with the aromatic  $\pi$  system of the pyrimidine ring. The calculated binding energy for this cluster is 514 cm<sup>-1</sup>. The methane-carbon atom is above the ring at 3.5 Å and is shifted  $\sim 0.1$  Å from the ring center toward the nitrogen atoms. The three hydrogen atoms of methane that point down toward the ring are equidistant from the ring at 3.1 Å: two of these hydrogens point directly at the ring nitrogen atoms. Again in agreement with the main experimental observations for these clusters, the calculations for pyrimidine (CH<sub>4</sub>)<sub>2</sub> clusters yield two distinct geometrical arrangements. The isotropic cluster has a calculated binding energy of 1029 cm<sup>-1</sup> and the anistropic cluster has a calculated binding energy of 879 cm<sup>-1</sup>.

#### **B. Pyrimidine-ethane**

The pyrimidine  $(C_2H_6)_1$  spectrum is quite complicated, consisting of a number of low intensity features and an intense feature at -60.7 cm<sup>-1</sup> with respect to the pyrimidine

Species	Energy (vac. cm <sup>-1</sup> )	Energy relative to pyrimidine 0% (cm <sup>-+</sup> )	Energy relative to cluster 06 (cm <sup>-1</sup> )	Assignment <sup>e</sup>
C.H.N.(CH.),	31 016.4	- 56.6	0	a
	31 020.9	- 52.1	45	
$C_{H_{A}N_{A}}(CH_{A})_{A}$	30 960 9	- 112.1	0	150 C
	30 966 0	- 107 0	51	
	31 025 8	- 47 2	0	aniso 08

TABLE II. Observed peaks in the spectra of pyrimidine-methane clusters

\*The = 112.1 cm<sup>-1</sup> shift is associated with two methanes added symmetrically above and below the pyrimidine ring (isotropic), as shown in Fig. 2. The = 47.2 cm<sup>-1</sup> shift is associated with the anisotropic configuration, as shown in Fig. 2.

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FIG. 2. Minimum energy configurations and binding energies for pyrimidine  $(CH_4)_1$  and pyrimidine  $(CH_4)_2$  as obtained using the LJ potential calculation described in the text.

origin (see Fig. 3). We might expect that little vdW vibronic intensity would be observed, based on pyrazine and other pyrimidine clusters. In order to begin to interpret these features we must consult the vapor phase room temperature pyrimidine monomer spectrum.<sup>21</sup> In these reports, features at -156 and +22 cm<sup>-1</sup> are assigned as the  $16a_1^2$  and  $16b_1^2$ sequence bands, respectively. The feature that appears in Fig. 3 at -39.1 cm<sup>-1</sup> may be assigned as the 16b  $\frac{1}{2}$  sequence band of the pyrimidine  $(C_2H_4)_1$  cluster built on the intense 60.7 cm<sup>-1</sup> cluster origin. If this identification is correct, then the clusters of pyrimidine  $(C_2H_6)_1$  are hot  $(T_{mb} \approx 260)$ K) and the features in Fig. 3 and Table II at -153.5. -99.3, -86.9, and -75.4 cm<sup>-1</sup> may well be hot bands associated with the cluster origins at -71.4, -60.7, and - 52.7 cm<sup>-1</sup>. Apparently the cluster formation process for pyrimidine ethane tends to warm the cluster. Table III summarizes these results.

Supersonic expansion of pyrimidine apparently does not produce the expected cooling for vibrational modes  $16b_1$  and  $16a_1$ . Vibrational temperatures for the  $16b_1$  mode have been reported to be in excess of 200 K. Ito and co-workers<sup>21d</sup> have also observed the  $16b_1^+$  transition for pyrimidine clustered with argon and nitrogen. In the present work, changes in backing pressure from 10 to 120 psi do not change the rela-

FIG. 3. Two-color TOFMS of pyrimidine  $(C_2H_6)_1$  in the region of the pyrimidine origin.

tive band intensities: apparently these modes present a bottleneck for vibrational cooling.

As can be seen in Fig. 4, three different configurations are calculated to be stable for the pyrimidine  $(C_2H_6)_1$  cluster. Configuration I has the long axis of ethane perpendicular to the plane of the pyrimidine ring. Configurations II and III have the ethane molecule long axis more or less parallel to the plane of the ring; for configuration II, the axis of ethane lies between a nitrogen and a carbon and for configuration III this axis lies between two carbon atoms. In each instance a CH<sub>3</sub> group lies more or less over the ring center.

### C. Pyrimidine-ammonia

The pyrimidine  $(NH_3)_1$  two-color TOFMS spectrum in the range 300 to 500 cm<sup>-1</sup> to the blue of the pyrimidine origin is presented in Fig. 5. The three features are associated with the 0<sup>o</sup><sub>0</sub> transitions of the pyrimidine  $(NH_3)_1$  clusters. The lack of significant vdW vibrational mode intensity indicates that the ground and excited state vdW potentials are nearly identical. The large cluster blue shift implies a strong hydrogen bonding interaction between the pyrimidine and the ammonia. The spectrum of Fig. 5 is taken with an ioniza-

TABLE III.	Observed	peaks in	the spectra of	f pyrimidir	e-ethane clusters.
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Species	Energy	Energy relative to pyrimidine 06 (cm <sup>-1</sup> )	Energy relative to cluster 06 (cm <sup>-1</sup> )	Tenta assigni	itive ment
C.H.N.(C.H.)	30 919 5	- 153.5			
	30 973.7	- 99.3			
	30 986.1	- 86.9			
	30 997 6	- 75.4			
	31 001.6	- 71.4	0		02
	31 012.3	- 60.7	0	11	og
	31 020.3	- 52.7	0	111	œ
	31 033.9	- 39.1	21.6	11	160 }

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tion energy of 45 110 cm<sup>-1</sup>; lowering the ionization energy to ~42 000 cm<sup>-1</sup> the two nearly degenerate features at + 367 cm<sup>-1</sup> decrease in intensity much more rapidly than the feature at + 496 cm<sup>-1</sup>. Table IV gives the energies and assignments for the pyrimidine  $(NH_3)_1$  clusters. Based on the idea that the three features in this spectrum are associated with three different pyrimidine  $(NH_3)_1$  cluster geometries, the two nearly isoenergetic configurations must be quite similar. The feature at +496 cm<sup>-1</sup> must correspond to a different (more hydrogen bonded) geometry.

Three different configurations are calculated for the pyrimidine  $(NH_3)_1$  cluster using the LJ-HB potential form (see Table I and Ref. 7). Two of these configurations have the ammonia above the pyrimidine plane and the third configuration has the NH<sub>3</sub> molecule not above the ring. These clusters are presented in Fig. 6 along with the calculated cluster binding energies. Configuration I has the ammonia N-atom 3.2 Å above the pyrimidine ring with the three am-





monia hydrogen atoms pointing down toward the  $\pi$  cloud. The two closest hydrogen atoms of the ammonia in this configuration are 2.7 Å from the pyrimidine plane. In configuration II, the ammonia nitrogen atom is 3.2 Å above the ring plane and points in the general direction of the ring N atoms. The closest two ammonia hydrogen atoms to the ring are at 2.7 Å. This cluster has a somewhat lower binding energy than cluster I (667 vs 689 cm<sup>-1</sup>). In the third pyrimidine (NH<sub>3</sub>)<sub>1</sub> configuration, the ammonia molecule does not reside over the pyrimidine ring: the NH<sub>3</sub> has a N-H bond in

TABLE IV. Pyrimidine (NH1)1.

Energy (vac. cm <sup>- i</sup> )	Energy relative to corresponding pyrimidine feature (cm <sup>-1</sup> )	Energy relative to corresponding pyrimidine- ammonia feature (cm <sup>-1</sup> )	Assignment
31 378.2	305.2	- 60.9	
31 439.1	366.1	0	1 08
31 441.0	368.0	0	11 08
31 485.7	412.7	46.6	Ŷ
31 520.0	447.0	80.9	
31 535.2	462.2	96.1	
31 561.3	488.3	122.2	
31 568.9	495.9	0	111 08
31 579.3	506.3	10.4	
31 610.1	537 1	41.2	
31 685 1	612.1	116.2	
32 044 6	360.6	0	1 64
32 050.8	366.8	0	11 64
32 097 2	355.2	0	[6b <sup>2</sup>
32 108 2		5	1 00 1
32 121.5	379 5	0	11.663
32 162.6	·		,
32 176 1	492.1	0	111.6a
32 224 7	482 7	, 0	III 66 <sup>2</sup>

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PYRIMIDINE (NH3)





FIG. 6. Minimum energy configurations and binding energies for pyrimidine (NH<sub>3</sub>), as obtained using a LJ plus HB potential calculation.

FIG. 7. Two-color TOFMS of pyrazine  $(NH_1)_1$  at the origin,  $10a_1^2$  and  $6a_2^2$  regions are shown. The cluster origin  $(0 \text{ cm}^{-1})$  is  $+117 \text{ cm}^{-1}$  to the blue of the pyrazine origin.

the plane of the pyrimidine ring with the H atom pointing to one of the pyrimidine nitrogens, 2.23 Å from it. The ammonia nitrogen is displaced in the xy plane by 3.3 and 2.8 Å along the x axis and y axis, respectively, as shown in Fig. 6, configuration III. This configuration has a calculated binding energy of 537 cm<sup>-1</sup>.

#### D. Pyrazine-ammonia

The pyrazine  $(NH_3)_1$  spectra in the  $0_0^0$ ,  $10a_0^1$ , and  $6a_0^1$  regions are presented in Fig. 7. One first notices the significant difference between these data and those of pyrimidine  $(NH_3)_1$ . The spectra all strongly suggest that only one configuration is present for the pyrazine  $(NH_3)_1$  system. The intense feature in the  $0_0^0$  spectrum is the cluster origin at  $+ 117 \text{ cm}^{-1}$  from the pyrazine  $0_0^0$  transition. The remaining features of this spectrum are vdW vibrational modes of the cluster. They will be analyzed in a future publication.<sup>20</sup> Table V gives the energies and the features observed in the spectrum. In this case, the ground and excited state potential surfaces must be significantly different.

The cluster  $10a_0^1$  transition is quite different from the cluster  $0_0^0$  or  $6a_0^1$  transitions. From this one concludes that strong vdW-internal mode coupling exists for the  $10a^1$  out of plane ring mode.<sup>22</sup> The vdW overtones and combination bands extend to more than 180 cm<sup>-1</sup> from the  $10a_0^1$  cluster origin feature. This is particularly striking in comparison with the  $6a_0^1$  vibronic band.

Only one configuration is calculated for the pyrazine  $(NH_3)_1$  cluster using the LJ plus HB potential function, in agreement with expectations from the spectra. The ammonia molecule hydrogen bonds to the ring nitrogens,

through two ammonia hydrogens. The nutrogen atom of the ammonia molecule is 3.2 Å above the ring plane with all three hydrogens pointing toward the ring. This geometry is depicted in Fig. 8. The calculated binding energy of the cluster is  $677 \text{ cm}^{-1}$ .

#### E. Benzene-ammonia

The origin and  $6_0^1$  transitions of the  $C_0H_0(NH_3)_1$  cluster are observed in two-color TOFMS. One can immediately conclude that at least one configuration of this cluster does not retain the benzene threefold axis. The spectra are traced in Fig. 9. The spectra for both transitions are far more complicated than any other cluster previously reported from our laboratory. Regions with similar structure can be found in the  $0^0_0$  and  $6^1_0$  spectra; in particular, the regions around + 20, - 20 and - 60 cm<sup>-1</sup> in each spectra bear some resemblance to one another. Nonetheless, features appearing in one spectrum do not appear in the other, and therefore at least two clusters of different geometry are probably responsible for the  $6_0^1$  transition. The  $0_0^0$  transition could arise from one reduced symmetry cluster. Since very little a priori spectroscopic analysis seems possible in this situation at the present time, great stock must be placed in the calculations.

Two different configurations are calculated for the  $C_6H_6(NH_3)_1$  cluster and both of them have the NH<sub>3</sub> molecule placed over the ring. Configuration I in Fig. 10 preserves the benzene  $C_3$  axis and configuration II does not. The high symmetry cluster has the N atom 3.3 Å from the ring plane and the three H atoms of NH<sub>3</sub> are 2.9 Å above the ring plane. Configuration II has the N atom 3.3 Å above the plane and slightly shifted from the ring center. Two H atoms of the

Energy (vac. cm <sup>- 1</sup> )	Energy relative to corresponding pyrazine feature (om -')	Energy relative to corresponding pyrazine- ammonia cluster (cm <sup>-1</sup> )	Assignment
30 967 3	913	- 25.7	
30 993 0	1170	0	രം
31 000.8	124 8	78	
31 019.5	143.5	26.5	
31 036.5	160.5	43 5	
31 043.6	167.6	50.6	
31 045.4	169.4	52.4	
31 067.3	191.3	74.3	
1 070.0	194.0	77.0	
31 077.4	201.4	84.4	
31 089.9	213.9	96.9	
1 093.7	217.7	100.7	
31 099.2	223.2	106.2	
31 378.7	119.7	0	10 a <sub>0</sub> 1
1 384.9	125.9	6.2	•
1 400.5	141.5	21.8	
1 403.7	144.7	25.0	
1 424.3	165.3	45.6	
441.8	182.8	63.1	
1 461.3	202.3	(82.6)*	
1 467.5	208.5	(88.8)	
1 507.5	248.5	128.8	
1 513.7	254.7	135.0	
1.539.5	280.5	160.8	
1 559.7	300.7	181.0	
1 580.2	120.5	0	6 a.
1 590.8	131.1	10.6	-
1 624.8	165.1	44.6	

\* May be associated with other pyrazine features.

NH, point towards C-C bonds and one points away from the ring plane. The NH<sub>3</sub> H-atoms in this configuration are at 2.9 (two of them) and 4.3 Å from the ring. The calculated binding energies of these two configurations are given in Fig. 10. Configuration II alone must generate the  $0_0^0$  spectrum in Fig. 9.

## F. Benzene-water

The  $C_6H_6(H_2O)_1$  cluster  $0_0^6$  and  $6_0^1$  are presented in Fig. 11. They are strikingly different from those of  $C_6H_6(NH_1)_1$ (Fig. 9) but bear a strong resemblance to the pyrazine  $(NH_3)_1$  spectra displayed in Fig. 7, as well as spectra of other systems studied in our laboratory.<sup>1-7</sup> In particular, a very clear vibronic progression is present for the  $0_0^6$  spectrum and will be analyzed in a future publication.<sup>20</sup> Table VI gives

## PYRAZINE - AMMONIA





-677 cm-

FIG. 8. Minimum energy configuration and binding energy for pyrazine  $(NH_1)$ , as obtained using a LJ HB form potential calculation.



FIG-9 Two-color TOFMS of benzene (NH , ) , in the region of the benzene  $\theta_{1}^{n}$  and  $\theta_{2}^{i}$  transitions.

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#### BENZENE-AMMONIA



FIG. 10. Minimum energy configurations and binding energies for benzene (NH<sub>3</sub>), as obtained using a LJ potential calculation.

these features and their energies. The  $6_0^1$  spectrum of  $C_6H_6(H_2O)_1$  has a different shift  $(+50 \text{ vs} + 85 \text{ cm}^{-1})$  and a different general pattern than the  $0_0^0$ . In addition, the  $6_0^1$  origin is split by roughly 1 cm<sup>-1</sup>. Considerable vibronic coupling must occur between the in plane carbon-carbon stretch  $6^1$  and the vdW modes. The  $C_0H_6(H_2O)_1$  cluster two-color TOFMS spectra are observed only at a higher ionization energy than required to observe the  $6_0^1$  spectrum of bare  $C_0H_6$ . The  $C_0H_6(H_2O)_1$  spectra depicted in Fig. 11 are taken with an ionization laser energy of 44 480 cm<sup>-1</sup> compared to 36 100 cm<sup>-1</sup> for the benzene monomer. The ionization energy for the benzene water cluster is 3300 cm<sup>-1</sup> higher than that of the benzene monomer.

Only one geometry is calculated for the  $C_8H_6(H_2O)_1$ cluster employing the LJ potential. The configuration and binding energy are given in Fig. 12. The oxygen atom of water is roughly centered over the ring at 3.2 Å above it and



FIG. 11. Two-color TOFMS of benzene  $(H_1O)_1$  in the region of benzene origin  $0_0^0$  and  $6_0^1$ . An expanded scale insert is shown of the  $6_0^1$  origin of the cluster.

the water hydrogen atoms are at 3.0 Å above the ring. The calculated water benzene binding energy is  $505 \text{ cm}^{-1}$ .

## IV. DISCUSSION

Cluster geometry is determined through analysis of twocolor TOFMS data for individual clusters and through calculations of cluster geometry and binding energies using an augmented LJ potential. Calculated binding energies have

TABLE VI Benzene (H,O),

7

Energy (vac. cm <sup>-+</sup> )	Energy relative to corresponding benzene feature (cm <sup>-1</sup> )	Energy relative to corresponding benzene- water feature (cm <sup>-+</sup> )	Assignment
	0	ρ 9	i
38 168 6	84 6	0	
38 173 9	89 9	53	
38 185 5	101 5	16.9	
39 191 0	107.0	22.4	
38 205 3	121 3	36.7	
38 221 6	137 6	530	
38 243 5	159 5	74 9	
38 274 9	190.9	106 3	
	6	h.	
		a	
38 655 4	48.4	0	
38 656 8	49-8		
38 683 8	76.8	27 7	
38 688 6	81.6	12.5	
38 694 2	87.2	181	
38 709 1	102 1	510	
38 720 7	113.7	64.6	
38 762 1	155 1	106.0	

"Shift taken with 6, at 49.1

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#### BENZENE-WATER



#### -505 cm-1

FIG. 12. Minimum energy configuration and binding energy for benzene  $(H_2O)_1$  as obtained using a LF potential calculation.

always been roughly bracketed by the experimentally observable range defined by two solute molecule vibrations (e.g., 520 to  $850 \text{ cm}^{-1}$  for the 1:1 complex).<sup>1-7</sup>

Before discussing the details of the clusters observed, some general remarks are in order. First, one should take note of the great apparent differences between the spectra of the various clusters studied: pyrazine and pyrimidine hydrocarbon spectra show little vdW vibronic structure but intense origins; the pyrazine (NH<sub>3</sub>)<sub>1</sub> spectrum displays elaborate and well developed vibronic progressions with a + 117 cm<sup>-1</sup> cluster shift for a single cluster; the pyrimidine (NH<sub>3</sub>)<sub>1</sub> spectra consist only of intense origins for three large shift  $(+366, +368, +496 \text{ cm}^{-1})$  configurations with no vdW vibronic development; the benzene (NH3) i spectra are red shifted, and too complex to interpret without further calculations; and the benzene (H2O), spectra are blue shifted with extensive vdW structure. Second, and perhaps even more astonishing, the LJ-HB (where appropriate) potential calculations parallel and reinforce these differences in all cases. That is, for example, the calculations suggest one hydrogen-bonded configuration for pyrazine (NH<sub>3</sub>), but three hydrogen-bonded configurations for pyrimidine (NH<sub>3</sub>), in agreement with the straightforward interpretation of the spectra. Third, the binding energies of the clusters seem relatively insensitive to the detailed configuration of the cluster. Fourth, spectral shifts are found to be a sensitive function of the detailed geometry of the cluster. Proximity to the  $\pi$  system is important for cluster red shifts, while hydrogen bonding yields in general cluster blue shifts with respect to the solute monomer origin.

#### A. Pyrimidine-methane

In the pyrimidine  $(CH_4)_1$  cluster the  $CH_4$  molecule is situated above the pyrimidine ring coordinated to the  $\pi$ cloud of the aromatic ring. The cluster has a simple spectrum with little vdW vibronic intensity following the pyrimidine vibronic origins. The cluster spectrum is red shifted, indicating that the excited state cluster is more tightly bound than the ground state cluster by roughly 60 cm<sup>-1</sup>. The overall appearance of the spectrum is similar to that of pyrazine methane.

The additive shift features in the spectrum of pyrimidine  $(CH_4)_2$  are attributed to the isotropic (symmetrical) geometry with a methane molecule on either side of the pyrimidine ring. The feature at  $-47.2 \text{ cm}^{-1}$  in the pyrimidine  $(CH_4)_2$  spectrum is attributed to the anisotropic configuration with both  $CH_4$  molecules on the same side of the ring. In this asymmetric geometry, one methane is more or less above the ring, and responsible for most of the  $CH_4-\pi$ -cloud interactions, and the other methane molecule is off the ring interacting primarily with the first methane, contributing little to the cluster spectral shift.

The overall behavior of this cluster system with regard to geometry, binding energy, and cluster population in the beam is very similar to that found for benzene, toluene, and pyrazine methane species.<sup>1-7</sup>

#### **B. Pyrimidine-ethane**

The pyrimidine  $(C_2H_6)_1$  cluster is similar to the pyrazine  $(C_2H_6)_1$  cluster:<sup>7</sup> both clusters have three geometrical configurations and each configuration of the two clusters has a similar binding energy. These configurations are also similar to those of the benzene  $(C_2H_6)_1$  cluster with the exception that, in the N-heterocyclic systems, two parallel ornentations of the  $C_2H_6$  long axis with respect to the ring plane are now possible.

The identification of calculated geometries with the three distinct spectroscopic features is of course tentative but can be pursued in the spirit of the arguments and correlations employed with the benzene and pyrazine systems.<sup>1-7</sup> Referring to Figs. 3 and 4, the feature at  $-71.4 \text{ cm}^{-1}$  can be associated with configuration I, the intense feature at  $-60.7 \text{ cm}^{-1}$  can be associated with configuration II, and the  $-52.7 \text{ cm}^{-1}$  feature is associated with configuration II. The lack of vdW vibronic structure for these transitions must be due to the similarity between the ground and excited state potentials for the clusters.

#### C. Pyrimidine-ammonia

The pyrimidine  $(NH_3)_1$  clusters have a unique spectrum which can only be interpreted as due to three distinct configurations with no vdW mode progression intensity following the well defined origins. Calculations, as pointed out previously, give exactly these conclusions and identification of origins in the spectrum with configurations seems straightforward. Configurations I and II of Fig. 6 are associated with the features at 366 and 368 cm -1 in Fig. 5. These two configurations are quite similar and have less hydrogen bonding interaction than the more blue-shifted single feature at 496 cm<sup>-1</sup>. The large spectral blue shifts of  $\sim$  365 and  $\sim$  500 cm<sup>-1</sup> must arise from the strong hydrogen bonding interactions. While none of these observations seem particularly striking in and of itself, in comparison with the pyrazine  $(NH_1)_1$  and benzene  $(NH_1)_1$  results, they are surprising; these will be discussed below.

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## D. Pyrazine-ammonia

The pyrazine  $(NH_3)_1$  cluster spectrum is completely different from that of pyrimidine  $(NH_3)_1$ . The cluster  $0_0^0$  is shifted + 117 cm<sup>-1</sup> from the pyrazine  $0_0^0$ , only one cluster geometry is present, and a rather extensive vdW vibronic structure is built upon the origin. The  $6a_0^1$  spectrum is quite similar. Two intense vdW vibronic transitions are associated with these cluster transitions. The  $10a_0^1$  vibration region of the pyrazine  $(NH_3)_1$  cluster, on the other hand, looks quite different from these other features and vibronic interactions between the out of plane  $10a^1$  carbon-carbon mode and the vdW modes are quite evident in the overall vibronic intensity pattern in the  $10a_0^1$  region.

Calculations predict only one configuration for this cluster system. The fact that the potential energy calculations can accurately parallel the spectral data for pyrimidine and pyrazine ammonia clusters which are clearly so different, gives us a high degree of confidence in the calculational process, the binding energies, and the potential form accuracy.

#### E. Benzene-ammonia

Assignment of the benzene  $(NH_3)_1$  spectra has not been attempted as yet because the spectra are too complicated. We present them only as part of the general picture indicating what spectra of relatively simple clusters of such systems can be like. The benzene  $(NH_3)_1$  clusters yield much more complicated spectra than either pyrazine or pyrimidine  $(NH_3)_1$  clusters do. In the benzene  $(NH_3)_1$  cluster, strong vdW vibronic interactions must be important for the cluster transition intensity.

Both calculated configurations (Fig. 10) contribute to the  $6_0^1$  transition but only configuration II generates the  $0_0^0$ spectrum. Note too, that both configurations most likely generate a red shift.

#### F. Benzene-water

Any  $C_0 H_0 (H_2 O)_1$  cluster will in principle generate a  $0_0^0$ transition. The  $0_0^0$  spectrum of  $C_0H_0(H_2O)_1$  (Fig. 11) thus strongly suggests that only one cluster geometry is realized for this system. The  $C_6H_6(H_2O)_10_0^0$  transition is much like the pyrazine (NH<sub>3</sub>)<sub>1</sub> spectrum. Well developed vdW vibronic features are observed. The 6<sup>1</sup> spectrum implies strong vibronic mixing between the in plane carbon-carbon deformation 61 and the vdW bends and torsions: the cluster shift is different for 6<sup>1</sup>/<sub>0</sub>, as is the intensity pattern. The cluster shift at the  $0_0^0$  transition is + 85 cm<sup>-1</sup> which indicates that the cluster excited state is destablized with respect to the ground state. The blue shift may be related to the unique hydrogen bonding capabilities of the H<sub>2</sub>O molecule with the  $\pi$  system of benzene. The calculated binding energy is probably  $\sim 50$  $cm^{-1}$  low for this cluster because the  $6_0^1$  transition at  $0_0^0$  plus 520 cm<sup>-1</sup> is observed. We have previously noted that the LJ potential binding energy is roughly 50 cm<sup>-1</sup> low compared to exp-6 and experimental values.

An infrared study of the 1:1 benzene-water complex in an argon matrix has been reported.<sup>23</sup> The work suggests that the water molecule hydrogen bonds to the benzene  $\pi$  system in a manner nearly identical to that found in the calculations presented in Fig. 12.

### V. CONCLUSIONS

The clusters studied in this work fall into two broad categories: a conventional set containing pyrimidine hydrocarbon clusters, the spectra of which are quite similar to those of other aromatic and pyrazine hydrocarbon systems. and benzene, pyrimidine, and pyrazine ammonia and benzene water clusters, the spectra of which are all unique and surprisingly erratic. In the latter grouping, spectral cluster shifts range from - 100 to + 500 cm<sup>-1</sup>, vdW vibronic spectra range from nonexistent to intense, vdW modes can be highly perturbing to the solute vibronic structure and energy, and the number of cluster configurations varies from one to three in an apparently arbitrary fashion. These differences notwithstanding, the Lennard-Jones (6-12-1) potential, augmented appropriately with hydrogen bonding interactions (10-12) as required, always gives geometry and binding energy results that are in complete agreement with the spectra as far as the comparison can be made (i.e., symmetry, numbers of configurations, red and blue shifts with regard to hydrogen bonding, etc.). The atom-atom LJ potential form has been chosen for these calculations because a large number of parameters for different types of atoms are available in the literature.

The benzene  $(NH_3)_1$  and  $(H_2O)_1$  spectra are quite different from one another. The shifts for these two clusters and their geometries seem to emphasize the importance of hydrogen bonding in the benzene  $(H_2O)_1$  cluster.

Pyrazine and pyrimidine water clusters are not found in these studies although they have been extensively investigated. These clusters are not observed perhaps because their excited states are dissociative, but more likely because of rapid excited state intersystem crossing and/or internal conversion.

<sup>1</sup>M. Schauer and E. R. Bernstein, J. Chem. Phys. 82, 726 (1985)

- <sup>1</sup>M. Schauer, K. S. Law, and E. R. Bernstein, J. Chem. Phys. 82, <sup>73</sup>6 (1985)
- <sup>1</sup>K. S. Law and E. R. Bernstein, J. Chem. Phys. 82, 2856 (1985).
- <sup>4</sup>K. S. Law, M. Schauer, and E. R. Bernstein, J. Chem. Phys. 81, 4871 (1984).
- E. R. Bernstein, K. Law, and M. Schauer, J. Chem. Phys. 80, 207 (1984).
- M. Schauer, K. Law, and E. R. Bernstein, J. Chem. Phys. 81, 49 (1984)
- 'J Wanna and E. R. Bernstein, J. Chem. Phys. 84, 927 (1986)
- <sup>4</sup>M. M. Carrabba, J. E. Kenny, W. R. Moomaw, J. Cordes, and M. Denton, J. Phys. Chem. 89, 674 (1985).
- <sup>3</sup>(a) H. Abe, N. Mikami, and M. Ito, J. Phys. Chem. 86, 1768 (1982). (b) H. Abe, N. Mikami, M. Ito, and Y. Udagawa, *ibid.* 86, 2567 (1982). (c)
- K, Fuke, H. Yoshiuchi, K. Kaya, Y. Achiba, K. Sato, and K. Kimura, Chem. Phys. Lett. 100, 179 (1984). (d) N. Gonohe, H. Abe, N. Mikami, and M. Ito, J. Phys. Chem. 89, 3642 (1985).
- <sup>10</sup>(a) J Hager and S. C. Wallace, J. Phys. Chem. 88, 5513 (1984), (b) 89, 3833 (1985).
- "D. E. Poeltl and J. K. McVey, J. Chem. Phys. 78, 4349 (1983)
- <sup>15</sup>(a) C. A. Haynam, D. V. Brumhaugh, and D. H. Levy, J. Chem. Phys. 79, 1581 (1983), (b) L. Young, C. A. Haynam, and D. H. Levy, *ibid.* 79, 1592 (1983).
- <sup>11</sup>G Smulevich, A. Amirav, V. Even, and J. Jortner, Chem. Phys. <sup>43</sup> (1982)
- <sup>14</sup>(a) P. M. Felker, W. R. Lambert, and A. H. Zewail, J. Chem. Phys. 77 1603 (1982), (b) L. A. Heimbrook, J. E. Kenny, B. E. Kohler, and G. W. Scott, J. Phys. Chem. 87, 280 (1983).

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Λ.

<sup>15</sup>M. Kasha, Discuss. Faraday Soc. 9, 14 (1950).
<sup>19</sup>(a) F. A. Momany, L. M. Carruthers, R. F. McGuire, and H. A. Scher-aga, J. Phys. Chem. 78, 1595 (1974); (b) G. Nemethy, M. S. Pottle, and H A. Scheraga, ibid. 87, 1883 (1983).

H. A. Scheraga, *ibid.* 87, 1883 (1983). <sup>1</sup>(a) C. A. Haynam, D. V. Brumbaugh, and D. H. Levy, J. Chem. Phys. 80, 2256 (1984); (b) D. V. Brumbaugh, J. E. Kenny, and D. H. Levy, *ibid.* 78, 3415 (1983); (c) K. E. Johnson, W. Sharfin, and D. H. Levy, *ibid.* 74, 163 (1981); (d) K. E. Johnson, L. Wharton, and D. H. Levy, *ibid.* 69, 2719 (1978).

"K. Yamanouchi, H. Watanabe, S. Koda, S. Tsuchiya, and K. Kuchitsu,

Chem. Phys. Lett. 107, 290 (1984)

<sup>19</sup>(a) P. J. Wheatley, Acta Crystallogr. 10, 182 (1957), (b) 13, 30 (1960). <sup>20</sup>J. A. Menapace and E. R. Bernstein, J. Chem. Phys. (to be published) <sup>21</sup>(a) A. E. W. Knight, C. M. Lawburgh, and C. S. Parmenter, J. Chem. Phys. 63, 4336 (1975); (b) I. Knoth, H. J. Neusser, and E. W. Schlag, J. Phys. Chem. 86, 891 (1982); (c) A. K. Jameson, H. Salgusa, and E. C. Lim. ibid. 87, 3007 (1983); (d) H. Abe, Y. Ohyanagi, M. Ichijo, N. Mikami, and M. Ito, ibid. 89, 3512 (1985).

<sup>22</sup>M. L. Sage and J. Jortner, J. Chem. Phys. 82, 5437 (1985).

<sup>21</sup>A. Engdahl and B. Nelander, J. Phys. Chem. 89, 2860 (1985)

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APPENDIX TWO

SUPERSONIC MOLECULAR JET STUDIES OF THE PYRAZINE AND

# PYRIMIDINE DIMERS

# Supersonic molecular jet studies of the pyrazine and pyrimidine dimers<sup>a)</sup>

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Mass selected optical spectra for the first excited singlet  $n\pi^*$  states of the pyrazine and pyrimidine dimers are presented. The species are created in a pulsed supersonic jet expansion. The spectra are analyzed based on ionization energy, vibronic structure, and relative energy with respect to the isolated monomer (cluster spectroscopic shift). Calculations of binding energy and geometry for these dimers are carried out employing a Lennard-Jones (6-12-1) and hydrogen bonding (10-12-1) potential. In the case of pyrazine, calculations and experiments agree that both parallel planar hydrogen bonded and perpendicular dimers are present in the expansion. The calculations also predict a parallel stacked and 90° rotated pyrazine dimer which is not observed. This latter species most likely forms an excimer in the excited state with a short lifetime and a highly red shifted broad spectrum. In the case of pyrimidine, calculations yield four planar hydrogen bonded species and a parallel stacked and displaced species. The spectra for the pyrimidine dimer are consistent with these configurations, in agreement with the calculations. No perpendicular configuration is calculated for the pyrimidine dimer and no spectroscopic features require postulating the existence of such a configuration. To explore further the agreement between calculated and experimental results for aromatic dimers, calculations are also presented for the tetrazine dimer. Three calculated geometries are obtained for the tetrazine dimer: a parallel stacked and 90° rotated species, a planar hydrogen bonded species, and a perpendicular species. Experimental spectra and calculations are in basic agreement for all dimers studied and, in general, support one another.

#### I. INTRODUCTION

Molecular dimers are of interest for a number of reasons. They can serve as model systems for condensed phase structure, dynamics, and nucleation and growth. Vibrational dynamics and reactions can be studied in clusters through the observation of vibrational dephasing or intramolecular vibrational redistribution (IVR) and vibrational predissociation (VP). Dimers also provide a model for higher order (i.e., secondary, tertiary, etc.) structure of more complex, flexible molecules. Finally, these small clusters can be treated as a new, weakly coupled state of matter appropriate in its own right for investigation and focused attention.

Small clusters or dimers are best studied by molecular jet techniques,<sup>1-6</sup> as the species are thereby isolated and free of extraneous perturbations. Of the molecular supersonic jet spectroscopy techniques available, the most useful is twocolor time of flight mass spectroscopy<sup>1</sup> (2-color TOFMS) because many different clusters [e.g., dimers, trimers, dimers (He)<sub>n</sub>, etc.] are simultaneously produced in the expansion process. Two-color TOFMS selects a cluster of particular mass, does not allow fragmentation of clusters to take place, thus maintaining mass integrity of the clusters, and yields a plot of cluster ion intensity in the chosen mass channel as a function of the cluster absorption spectrum.

Dimers of benzene, toluene, and benzene-toluene have been studied in our laboratory using the 2-color TOFMS technique.<sup>1</sup> Coupled with the experimental findings, a potential energy calculation of the structure and binding energy of these dimers based on an exponential-six (exp-6) function has also been reported.<sup>2</sup> Spectroscopic data and potential energy calculations have been analyzed to arrive at a set of consistent geometries for these dimers. The benzene dimer is suggested to have a parallel displaced structure and the toluene and toluene-benzene dimers are suggested to have both parallel displaced and perpendicular geometries. The benzene dimer characterization rests on isotopic substitution, absence of resolved splittings at the cluster 0%, observation of only one feature, respectively, for  $(C_6H_6)_{2}$ ,  $(C_6D_6)_2$ , and  $C_6H_6C_6D_6$  at the cluster  $0_0^0$ , and calculations employing the observed molecular quadrupole moment of C<sub>6</sub>H<sub>6</sub> to set partial and atomic charges and multipolar terms. In all instances, the experiments and calculations appear to arrive at self-consistent and independent conclusions which are in agreement with one another.

Molecular jet studies of other isolated dimers have also been reported. Rotationally resolved fluorescence excitation and dispersed emission spectra of tetrazine,<sup>3</sup> phenyltetrazine,<sup>4</sup> and dimethyltetrazine<sup>3</sup> dimers have been reported. Hydrogen bonded benzoic acid<sup>6</sup> and benzoic acid-p-toluic acid<sup>7</sup> dimers have also been studied.

In this paper we report rotationally resolved 2-color TOFMS of pyrazine and pyrimidine isolated molecules at a resolution of 0.08 cm<sup>-1</sup>. Unfortunately, this is insufficient resolution to obtain rotationally resolved 2-color TOFMS of the pyrazine and pyrimidine dimers. Computer simulations, based on a reasonable symmetric top algorithm, predict a resolution of at least 0.005 cm<sup>-1</sup> (150 MHz) is needed to observe rotational structure for the dimers, assuming they are rigid.

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In addition to the experimental spectroscopic methods used to study these dimers, potential energy calculations utilizing a Lennard-Jones (LJ: 6-12-1) potential are performed to yield minimum energy geometries and binding energies. This potential form is replaced with a LJ-hydrogen bonding (HB; 10-12-1) form for the appropriate set of atoms. The calculation and parameters employed are discussed and presented in an earlier publication.3 These potentials, with literature parameter values, give the same results as the exp-6 with dipole-dipole, dipole-quadrupole, and quadrupolequadrupole terms for the benzene, toluene, and benzene-toluene (and pyrazine and pyrimidine) dimers. The major advantage of the LJ form presently used in our studies is that more atomic parameters are available in the literature' and experimental multipole moments are not required for each system.

Dimer geometry is determined through analysis of experimental (e.g., shifts, ionization energies, origin identification, vibronic analyses, etc.) and calculational results.

#### **II. EXPERIMENTAL PROCEDURES**

A pulsed valve supersonic molecular jet system is used to generate the dimers. The pulsed valve is mounted in the mass spectroscopy chamber of a two chambered vaccum system. Since the duty cycle of the valve is roughly  $10^{-3}$ , the 10 in. diffusion pump on the chamber and the 6 in. diffusion pump on the TOFMS flight tube adequately handle the gas load and maintain the chamber pressure below  $2 \times 10^{-6}$ Torr. The system is described in previous publications.<sup>1,8</sup> The beam passes through a skimmer and then into the ionization region of a TOFMS. Two separately tunable lasers provide the photons for the  $S_0 \rightarrow S_1$  transition and the  $S_1 \rightarrow$  cluster ion transition.

Rotationally resolved 2-color TOFMS are obtained through pressure tuning of the grating box of the pump  $(S_0 \rightarrow S_1)$  dye laser oscillator cavity. The output of this laser is narrowed by an etalon placed between the dye cell and the grating in the oscillator cavity. The doubled output from this dye laser is 0.08 cm<sup>-1</sup> in width. The laser can be scanned over roughly 20 cm<sup>-1</sup> for a N<sub>2</sub> pressure variation of 10 to 1500 Torr.

Pyrazine and pyrimidine are obtained from Aldrich Chemical Company and used without purification. The sample is placed in a trap behind the pulsed valve through which He flows at 120 psig.

The LJ potential energy function and calculational procedure have previously been described.<sup>8</sup> The additional constants needed for this work are the (aromatic) N···H hydrogen bonding values:  $B = 8.244 \times 10^3$  kcal Å<sup>10</sup>/mol and  $A' = 3.2897 \times 10^4$  kcal Å<sup>12</sup>/mol. In order to check the LJ potential form, in particular for the pyrimidine dimer, LJ plus multipolar (i.e., dipole-dipole, dipole-quadrupole. and quadrupole-quadrupole) potential calculations are also performed.<sup>2</sup> The pyrimidine dipole and quadrupole moments<sup>10</sup> are taken to be  $-2.97 \times 10^{-18}$  esu cm and  $-1.91 \times 10^{-26}$ esu cm<sup>2</sup>, respectively.

Calculations are also reported which simulate the rotational structure of pyrazine and pyrimidine monomers and dimers. A symmetric top model is employed for this fit because it is simple, reasonably accurate, and in general is well suited to the purpose of roughly predicting the unresolved dimer structure. Both molecules are nearly symmetric tops ( $\kappa \approx 0.9$ ). The form of the equations and methods employed are given by Herzberg.<sup>11</sup> The rotational temperature achieved in our system is  $\sim 2$  K. The rotational constants used in the dimer rotational spectra calculations are found from the calculated LJ geometries. The molecular geometries can be found in Ref. 12 for pyrazine and Ref. 13 for pyrimidine.

## III. RESULTS

#### A. Pyrazine dimer

The spectrum of the pyrazine dimer at the  $0_0^0$  transition is presented in Fig. 1 at two ionization energies, both of which are lower than the minimum ionization (second photon) energy of 44 000 cm<sup>-1</sup> required to observe the 2-color TOFMS of the pyrazine monomer. Lowering the ionization energy from 43 182 to 42 185 cm<sup>-1</sup> causes three of the dimer related peaks to disappear: these features are found at -11.0, 12.0, and 26.2 cm<sup>-1</sup> on the scale of Fig. 1. From the nature and appearance of these features, the + 12 and + 26 cm<sup>-1</sup> peaks are quite likely vibrations built on the -11 cm<sup>-1</sup> origin of a given configuration cluster. The intense features that remain at the lower ionization energy are found at - 26.3, -5.8, + 34.1, and 50.7 cm<sup>-1</sup>. From this variation of ionization energy one can determine that at least two configurations of the pyrazine dimer are present in the su-





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personic jet expansion. The pyrazine- $d_4$  dimer  $0_0^9$  spectrum at two different ionization energies is presented in Fig. 2: the similarity between the pyrazine- $h_4$  and  $-d_4$  dimer spectra is quite striking and reinforces our identification of origins and vdW vibronic features. The features that vanish at lower ionization energy are found at -11.5, 11.5, 25.5, and 64.1 cm<sup>-1</sup> with respect to the pyrazine- $d_4 0_0^9$  transition (31 030.4 cm<sup>-1</sup>). These should be compared with the numbers in Table I. The features that remain with lower ionization energy are iocated at -26.9, -6.5, 31.0, and 45.9 cm<sup>-1</sup>.

The spectra of the pyrazine dimer at other pyrazine vibronic origins are presented, along with the  $\theta_0^0$  spectrum for comparison, in Fig. 3. The feature at roughly + 61 cm<sup>-1</sup> in this figure (+34.1 cm<sup>-1</sup> in Fig. 1) is clearly an additional origin. The energy values and shifts for these features are presented in Table I. Since the pyrazie dimer is still observed at  $10a_0^2$ , its binding energy is greater than 800 cm<sup>-1</sup>. No other features, appearing in the dimer mass channel, are found within - 400 cm<sup>-1</sup> of the pyrazine  $\theta_0^0$  transition. The spectra of the pyrazine- $d_4$  dimer at these other vibronic monomer origins are again very similar to those of the pyrazine- $h_4$  dimer. We concluded from these spectra (not presented) that the third origin for the deuterated dimer lies at 31.0 cm<sup>-1</sup> from the  $\theta_0^0$  of the deuterated monomer.

Utilizing a Lennard-Jones potential function with a hydrogen bonding form, three configurations for the pyrazine dimer are calculated. Two of these configurations, a planar hydrogen bonded form and a perpendicular form, are displayed in Fig. 4. A parallel, stacked and 90° rotated structure



FIG. 3. Two-color TOFMS of the pyrazine dimer in the  $0^6_{11}$ ,  $10a^5_{11}$ , and  $6a^6_{12}$  regions. These spectra are taken at high (  $\sim$ 43 200 cm  $^{-1}$ ) ionization energy.





FIG -4. Minimum energy configurations and binding energies for parables dimer as obtained with a LJ plus HB potential calculation.

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is also found. The binding energies for the former two configurations are presented in the figure. In the planar geometry the ring centers are at 5.6 Å from each other and the suggested C-H...N hydrogen bonds are 2.3 Å long. The pyrazine at the stem of the perpendicular configuration is situated in a symmetric position above the base pyrazine, with two of its hydrogens pointing to the base pyrazine nitrogens and equidistant from them. This perpendicular configuration has a 4.6 Å pyrazine center to center separation and apparent 2.6 Å H...N bond lengths.

#### **B.** Pyrimidine dimer

Three segments of the pyrimidine dimer 2-color TOFMS spectrum in the  $0_0^0$  region are displayed in Fig. 5 for two different ionization energies. Based only on the position and appearance of these segments we suggest that the features at -168 and +296 cm<sup>-1</sup> are each associated with different geometries. The grouping of features in the +170 cm<sup>-1</sup> region must be associated with more than one dimer configuration, as these eight sharp, relatively intense features are clearly not vdW vibronic progressions built on a single  $0_0^0$  origin. The minimum ionization energy for the pyrimidine monomer is near 44 090 cm<sup>-1</sup> above the  ${}^{18}_{1}(n\pi^{\circ})$  excited state at 31 073 cm<sup>-1</sup>. As can be seen from Fig. 5, features in two of the three regions displayed disappear as the ionization energy is lowered from 44 363 to 42 320 cm<sup>-1</sup>.

Calculations using a LJ-HB potential function yield four planar configurations, a parallel stacked head-to-tail displaced configuration, and a parallel stacked undisplaced configuration with the two pyrimidine molecules rotated 90° with respect to each other. The latter configuration most likely does not contribute to the observed dimer spectrum since it will probably form an excimer. No perpendicular geometry is calculated even with the LJ-HB potential augmented with multipolar terms. Figures 6 and 7 give those calculated geometries for the pyrimidine dimer which can produce the observed spectral features. The parallel displaced geometry shown in Fig. 6 is head-to-tail displaced by



FIG. 5. Three segments of the 2-color TOFMS spectra of the pyrimidine dimer at two ionization energies, top trace at an ionization energy of 44 363 cm<sup>-1</sup> and the lower trace at an ionization energy of 42 320 cm<sup>-1</sup>. The energy scale is relative to the pyrimidine monomer  $0_0^6$ .



PYRIMIDINE DIMER

#### -1478 cm<sup>-1</sup>

FIG. 6. Minimum energy configuration and binding energy for the stacked pyrimidine dimer as obtained with a LJ plus HB potential calculation.

0.6 Å along the CH–CH line from the molecular center; the interplane separation is 3.3 Å. The calculated binding energy for this dimer is  $1478 \text{ cm}^{-1}$ .

The planar configurations are displayed in Fig. 7. Configuration I has a center to center distance of 5.5 Å, two N---H hydrogen bonds (2.3 Å separation), and a calculated binding energy of 709 cm<sup>-1</sup>. Configurations II, III, and IV also display some hydrogen bonding but to a lesser extent than that displayed in configuration I. In these latter three cases, the less "acidic" hydrogens, not between the two N atoms, are involved in the "hydrogen bonds": the pyrimidine molecule center to center distance is  $\sim 6.0$  Å and the calculated binding energies range from 400 to 430 cm<sup>-1</sup>, substantially less than the binding energy for configuration I. Configuration II has two N--H hydrogen bonds each of 2.9 Å. Configurations III and IV have a nitrogen atom of one pyrimidine equidistant from two hydrogens of the other pyrimidine with an apparent hydrogen bond distance of 2.9 Å. Planar configurations in which two nitrogens are facing each other are not stable.



FIG. 7 Minimum energy configurations and binding energies for the planar pyrimidine dimers as obtained with a LJ plus HB potential calculation

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## C. Rotational structure

Rotationally resolved 2-color TOFMS data can be obtained for the pyrazine and pyrimidine monomers using the resolution presently available in our laboratory ( $\Delta v = 0.08$ cm<sup>-1</sup>). The spectra are presented in Fig. 8. These well resolved spectra evidence a central Q branch with well developed R and P branches to the high and low energy sides, respectively. The calculated rotational structures for these transitions are presented in Fig. 9. In order to make the fit look reasonable, a 0.1 cm<sup>-1</sup> Gaussian linewidth was incorporated in the calculated spectrum.

Considering that a symmetric top equation is used for the fit, the agreement between the calculated and experimental results is excellent. The purpose of this exercise is to observe and calculate the rotational spectrum of a dimer. One can see from Fig. 10 (top) that the rotational structure of the pyrazine dimer is not evident at this laser linewidth. Similar conclusions arise from the spectra of the benzene dimer (see Fig. 11). Computer simulations of the pyrazine dimer spectrum (based on the symmetric top calculations), show that a ~0.005 cm<sup>-1</sup> laser linewidth is required to resolve rotational transitions for the aromatic dimers (Fig. 10 bottom). An attempt to fit the rotational contours to parallel or perpendicular transitions of parallel or perpendicular dimers demonstrates that no convincing conclusions concerning dimer geometry can be reached in this manner. In fact, the spectra of the parallel and perpendicular pyrazine dimer origins do not appear different at this resolution. The calculated contours are found using rotational constants of  $A^* = 0.0611, B^* = 0.0157, \text{ and } C^* = 0.0141 \text{ cm}^{-1}$  for the perpendicular dimer and  $A^* = 0.0381$ ,  $B^* = 0.0153$ , and  $C^* = 0.0112 \text{ cm}^{-1}$  for the parallel planar dimer.



FIG 8 Two-color TOFMS rotational spectra of the origins of pyrazine (top) and pyrimidine (bottom) monomers.



FIO. 9. Simulated rotational spectra of the pyrazine and pyrimidine origins.

## IV. DISCUSSION

## A. Pyrazine dimer

In the following paragraphs, only the pyrazine- $h_4$  dimer results will be discussed in detail. The similarity between the pyrazine- $h_4$  and  $-d_4$  dimer results obviates the need for discussion of these data separately.

One of the most important experimental observations concerning the pyrazine dimer is the change in the spectrum as a function of ionization laser or second photon energy. Lowering the ionization laser energy by 1000 cm<sup>-1</sup> to 42 185 cm<sup>-1</sup> causes three features to disappear: two of these are assigned as vibrations built on a single origin (see Table I and Fig. 1). Further reduction of the ionization energy to 41 721 cm<sup>-1</sup> results in no observed TOFMS spectrum for the pyrazine dimer. At least two different geometries of the pyrazine dimer are therefore present in the beam. The dimer with the higher ionization energy is probably a symmetrical dimer with two symmetry equivalent molecules because only one origin is associated with the high ionization energy spectrum.

Different geometries will possess different ionization energies depending on the involvement of the  $\pi$  clouds in the overall dimer interaction. For example, a planar hydrogen bonded dimer would probably have a poor ion "solvation" or stabilization and might therefore have a higher ionization energy. This geometry would in addition have only one spec-

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TABLE I. Pyrazine dimer.

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FIG. 10. Two-color TOFMS rotational spectrum of the pyrazine dimer origin (top) and computer simulated rotational spectra of the pyrazine dimer origin (bottom). The origin is at 30 849.5 cm<sup>-1</sup> (-26.5 cm<sup>-1</sup> origin in Fig. 1). The FWHM is roughly 2 cm<sup>-1</sup>. The 30 865.0 cm<sup>-1</sup> origin looks nearly identical to this one at the experimental resolution.

troscopic origin. A perpendicular geometry dimer would, on the other hand, probably evidence two features and a lower ionization energy due to the  $\pi$ -cloud involvement of the "horizontal" pyrazine in the stabilization of the ion.

The calculations give three general geometries for the pyrazine dimer: a parallel planar, a perpendicular, and a parallel stacked on 90° rotated geometry. The latter geometry is not discussed in this work because it likely is not important for any of the spectroscopic observations presented earlier. The remaining two configurations give rise to three separate spectra: one for the parallel planar geometry (1) and two for the perpendicular geometry (base IIa and stem IIb).

BENZENE DIMER O:



FIG 11. Two-color TOFMS of the benzene dimer  $0^0_5$  transition at 0.08 cm  $^+$  resolution. Most of the "features" in this trace are noise and are not reproducible

Energy (vac. cm <sup>- 1</sup> )	Energy relative to corresponding pyrazine feature (cm <sup>-1</sup> )	Energy relative to corresponding pyrazine dimer feature (cm =)	Assignments*
30 849.5	- 26.5	0	II base 0%
30 865.0	- 11 0	0	105
30 870.2	- 5.8	20.7	
30 879.1	3.1	29 6	
30 888.0	12.0	23.0	
30 891.4	15.4	41.9	
30 898.7	22.7	49 2	
30 902.3	26.3	37.3	_
39 910.1	34.1	0	II stem 05
30 914.0	38.0	3.9	
30 918.2	42.2	8.1	
30 926.7	50.7	16.6	
31 228.5	- 30.5	0	II base 10a
31 238.5	- 20.5	0	I 10a0
31 245.4	- 13.6		
31 254.8	- 4.2		
31 289.4	30.4	0	II stem 10a
31 297.6	38.6		
31 314.4	55.4		
31 433.3	- 26.4	0	II base 6a
31 445.1	- 14.6	0	1600
31 456.8	- 2.9		
31 467.8	8.1		
31 481.7	22.0		
31 488.9	29.2	_	
31 493.2	33.5	0	II stem 640
31 507.9	48.2		
31 565.7	106.0		
31 677.9	- 21.1	0	II base 10a
31 686.1	- 12.9	0	I 10az

\*See Fig. 4.

Table I gives the assignment of the dimer spectra. The planar geometry (1) is assigned to the origin at -11 cm<sup>--</sup> (Fig. 1) since this single origin feature is associated with the higher ionization energy. The other two origins at -26.5and  $+ 34.1 \text{ cm}^{-1}$  (Fig. 1) with respect to the pyrazine monomer origin are assigned to the perpendicular dimer because they both show the same low ionization energy. The base (IIa) is associated with the red shifted origin and the stem (IIb) is associated with the blue shifted origin. This latter correlation between spectra and calculated structures is based on the argument presented in previous publications<sup>1,2</sup> relating solvent cluster shifts and  $\pi$ -cloud involvement in the solute-solvent interaction: the larger the red shift, the more direct is the interaction between the system and the solvent. Thus the base molecule should be expected to have a larger red shift than the stem molecule.

The pyrazine  $6a^{1}$  (in-plane C-C stretch) and  $10a^{2}$  (outof-plane C-C bend) vibrational modes show strong interaction with the van der Waals modes (Fig. 3).

## **B. Pyrimidine dimer**

The ionization energy for the pyrimidine dimer system is again an important piece of information used to help determine the number of different configurations responsible for

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S-TETRAZINE DIMER

FIG. 12. Minimum energy configurations and binding energies for the tetrazine dimer as obtained with a LJ plus HB potential calculation.

the observed spectra and, perhaps, their geometry. Lowering the ionization by 2 043 cm<sup>-1</sup> to 42 320 cm<sup>-1</sup> causes the feature at -168 cm<sup>-1</sup> to disappear, the features at  $\sim +175$ cm<sup>-1</sup> nearly to disappear, and the feature at +296 cm<sup>-1</sup> to reduce in intensity. In addition, dimer spectral shifts can also be employed to associate calculated geometries with spectroscopic features: red shifted origins can be assigned to parallel stacked geometries, and blue shifted origins to planar hydrogen bonded forms. Perpendicular geometries can be responsible for both red and blue shifts depending on which molecule of the dimer is involved.<sup>1</sup>

LJ-HB potential calculations suggest one parallel stacked head-to-tail displaced, one parallel stacked rotated, and four parallel planar configurations for the pyrimidine dimer. No perpendicular geometry can be calculated using LJ-HB or a multipolar form.<sup>1,2</sup> All calculations give nearly identical geometries and binding energies for the parallel planar and stacked configurations.

The features at -168 cm<sup>-1</sup> is suggested to be due to the parallel stacked and displaced head-to-tail geometry. One would expect this structure to have only one spectroscopically observed 0<sup>0</sup><sub>0</sub> transition and a substantial red shift. The remaining features in the spectra, due to their significant blue shifts, must be attributed to planar hydrogen bonded dimers. The feature at +296 cm<sup>-1</sup> is suggested to be due to configuration I shown in Fig. 7. This configuration of the pyrimidine dimer forms two hydrogen bonds both of which involve the hydrogen atoms between the two ring nitrogen atoms on each pyrimidine: these hydrogens are the most electropositive (acidic) hydrogens on the ring. This configuration also has the monomers closest to each other (5.5 Å compared to 6.0 Å in the others). These factors suggest that configuration I gives rise to the most blue shifted feature in the spectra. The remaining three configurations II, III, and IV must generate the features in the +175 cm<sup>-1</sup> region. Configuration II is a symmetrical dimer and will account for one feature while configurations III and IV each will account for two features since the pyrimidines in these last two configurations are not symmetry equivalent. Assigning these features to configurations II, III, and IV is a difficult task without further information: five of the eight major features in this region can, however, be associated with origins of configurations II, III, and IV.

The parallel planar hydrogen bonded configuration I is assigned to the large blue shift, low ionization energy feature, and the parallel stacked displaced geometry is assigned to the large red shifted, high ionization energy feature. On the other hand, the parallel planar pyrazine dimer is assigned to the feature with the higher ionization energy (and also a small red shift). Clearly the two dimers have a very different electronic structure and the component monomers must interact in a different manner. A possible explanation for these apparent differences is that the N-C-N moiety of the pyrimidine system becomes the positive end of the molecular ion which is in turn well solvated in the parallel planar dimer thus lowering the ionization energy, and that the loss of electron density in the N-C-N region in the  $n\pi^{\bullet}$ excited state reduces the hydrogen bond energy thus increasing the energy of the excited  $S_1$  state and causing a dimer blue shift. Similar arguments can be rendered to rationalize a negligible shift for the pyrazine system. We caution, however, that all such qualitative reasoning is subject to verification by more rigorous quantum mechanical calculations.

To ensure that our LJ-HB potential can produce other perpendicular dimer configurations, we have calculated the geometries expected for the tetrazine system. The tetrazine dimer has been studied by Levy and co-workers,<sup>3</sup> who have reported two geometries: a parallel planar configuration and a perpendicular configuratiou. These experiments involve rotationally resolved fluorescence excitation spectra. Our calculations generate three geometries for this dimer: a parallel planar configuration, a parallel, stacked and 90° rotated configuration, and a perpendicular configuration, as shown in Fig. 12. The calculated perpendicular configuration has one hydrogen of the stem tetrazine pointing towards an N-N

## TETRAZINE DIMER



FIG. 13. Calculated rotational contours of the parallel polarized transition of the perpendicular tetrazine dimer centered at 18.272.0 cm<sup>-1+1</sup>(0 cm<sup>-1+1</sup>in the figure). The upper trace is calculated using the rotational constants obtained from the perpendicular configuration reported in Ref. 3(a). The lower trace is calculated using the rotational constants obtained from the perpendicular configuration reported in this work. A symmetric top model is assumed for both calculations and the intensities used are those of Ref. 3(a). The rotational constants employed are given in the text, in the symmetric top approximation  $\overline{B}^{+} = (B^{+} + C^{+})/2$ .

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bond of the base tetrazine. In this configuration the plane of the stem tetrazine bisects the two N-N bonds of the base tetrazine. Levy's published perpendicular configuration<sup>3(a)</sup> has one hydrogen of the two C-H bonds of the stem tetrazine pointing toward one C-H bond of the base tetrazine and the plane of the stem tetrazine passing through the two C-H bonds of the base tetrazine. Rotational constants reported by Levy and obtained from a rotational analysis of the perpendicular configuration for parallel polarization are  $A^{*} = 0.072 \ 87 \ \text{cm}^{-1}$ ,  $B^{*} = 0.017 \ 22 \ \text{cm}^{-1}$ , and  $C^* = 0.01649$  cm<sup>-1</sup>. Rotational constants obtained from our calculated perpendicular configuration are A'' = 0.06458 cm<sup>-1</sup>, B'' = 0.01644 cm<sup>-1</sup>, and  $C^* = 0.018$  466. These two sets of rotational constants render reasonably similar spectra, as can be seen in Fig. 13.

#### **V. CONCLUSION**

The analysis of the structure and properties of the pyrazine and pyrimidine dimers is based on an interpretation of ionization energy dependence, van der Waals vibronic structure, dimer spectral shifts, and potential energy calculations with LJ-HB and multipolar forms.

Variation of the ionizing laser energy allows different configuration dimers of a particular species to be identified. The pyrazine ( $h_4$  and  $d_4$ ) dimer has two identified geometries based on ionization energy dependence and vibronic analysis. Given the dimer spectral shifts and calculations, these have been associated with a planar parallel hydrogen bonded configuration and a perpendicular configuration. A third geometry, planar stacked and rotated 90°, is calculated but not observed probably due to excimer formation.

The pyrimidine dimers absorb in three spectral regions. The lowest energy feature is thought to be the calculated head-to-tail parallel stacked displaced geometry, the highest energy feature is assigned as a parallel planar strongly hydrogen bonded form in which the most electropositive H atoms are involved in the hydrogen bonding. The features at  $+ 175 \text{ cm}^{-1}$  are attributed to different planar configurations which are only weakly hydrogen bonded through the less acidic hydrogens on the rings. Dimer spectral shifts are expected to follow the rules determined previously for solute-solvent clusters: the major red shift mechanism is  $\pi$ -system coordination or overlap between the two molecules and the major blue shift mechanism is hydrogen bonding. Ionization energies can be rationalized in accordance with the general notions of ion solvation by either the  $\pi$  system in the case of pyrazine or the N-C-N hydrogen bonding region in the case of pyrimidine. AND ALL DAMAGE AND A CONTRACT OF A CONTRACT OF

Calculations are also presented for the tetrazine dimers to compare parallel planar and perpendicular spectroscopically assigned geometries and our calculations. Calculations predict, and experiments suggest, perpendicular geometries for toluene, benzene-toluene, pyrazine, and tetrazine but not for benzene. Moreover, calculations predict, and experiments are consistent with, the absence of perpendicular geometries for the pyrimidine dimer. Perhaps one of the most remarkable results of this study is the rather large number of different, roughly equal binding energy configurations found for the N-heterocyclic aromatic dimers in general.

- <sup>1</sup>K. S. Law, M. Schauer, and E. R. Bernstein, J. Chem. Phys. 81, 4871 (1984).
- <sup>2</sup>M. Schauer and E. R. Bernstein, J. Chem. Phys. 82, 3722 (1985).
- <sup>3</sup>(a), C. A. Haynam, D. V. Brumbaugh, and D. H. Levy, J. Chem. Phys. 79, 1581 (1983); (b), L. Young, C. A. Haynam, and D. H. Levy, *ibid*. 79, 1592 (1983).
- <sup>4</sup>Y. D. Park and D. H. Levy, J. Chem. Phys. 81, 5527 (1984).
- <sup>5</sup>D. V. Brumbaugh, C. A. Haynam, and D. H. Levy, J. Chem. Phys. 73, 5380 (1980).
- <sup>6</sup>D. E. Poelti and J. K. McVey, J. Chem. Phys. 78, 4349 (1983).
- Y. Tomioka, H. Abe, N. Mikami, and M. Ico, J. Phys. Chem. 89, 5186 (1984).
- <sup>4</sup>J. Wanna and E. R. Bernstein, J. Chem. Phys. 84, 927 (1986).
- <sup>9</sup>(a). F. A. Momany, L. M. Carruthers, R. F. McGuire, and H. A. Scheraga, J. Phys. Chem. 78, 1595 (1974); (b) G. Nemethy, M. S. Pottle, and H. A. Scheraga, *ibid.* 87, 1883 (1983).
- <sup>10</sup>F. Mulder, G. Van Dijk, and C. Huiszoon, Mol. Phys. 38, 577 (1979).
   <sup>11</sup>G. Herzberg, Molecular Spectra and Molecular Structure. II. Infrared and Raman Spectra of Polyatomic Molecules (Van Nostrand Reinhold,
- New York, 1945), Chaps. I and IV.
- <sup>12</sup>P. J. Wheatley, Acta Crystallogr. 10, 182 (1957).
- <sup>13</sup>P. J. Wheatley, Acta Crystallogr. 13, 80 (1960).

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APPENDIX THREE

"ECCEMP2"

20 + Program name: ECCEMP2 30 1 40 ) This program calculates cluster binding energy and geometry using an empiri 50 / cal atom-atom Lennard-Jones potential. The potential includes 6-12 general 60 / non-bonding, 10-12 general hydrogen bonding , and 1 monopole charge terms. 70 <sup>1</sup> The program will perform calculations on clusters composed of up to three 80 / molecules (subunits). 90 1 1001 Geometry and binding energy optimization is accomplished by analyzing the 110 I forces and torques on the cluster subunits which result from the intermole 120 I cular potential between the cluster subunits. The molecules are translated 130 I in three dimensions and rotated about the their centers-of-mass in three 140 I dimensions in response to the forces and torques. Binding energy and 150 I geometry optimization is achieved by simultaneously manipulating the six 150 ! degrees of freedom until the forces and torques are zero; that is, until 170 | minimum energy and optimal geometry are obtained. 180 1 190 / This program also calculates the intermolecular force field for van der 200 <sup>1</sup> Waals vibrational mode modeling which can be conducted using the VDWNCA 210 | program. The ECCEMP2 program will determine the intermolecular force 220 <sup>1</sup> field for clusters containing up to 25 atoms total (solute+solvent). 240 1 250 OPTION BASE 1 260 PRINTER IS 1 270 OUTPUT 2; CHR\$(255)&"K"; 280 DE6 290 1 310 | Coordinate matrices for atoms composing cluster subunits. 320 1 330 ! Co(+)=Carbon coordinates. 340 1 Hh(+)=Hydrogen coordinates. 350 ! Nn(+)=Nitrogen coordinates. 360 ! Oo(+)=0xygen coordinates. 370 ! Zn(+)=User defined coordinates. 380 | (+)=(subunit,atom \$,atom type,coordinates and charges and hydrogen 390 ! bonding flags). 410 1 420 DIM Co(3,44,5,5),Hh(3,44,5,5),Nh(3,12,5,5),Oo(3,12,5,5),2h(3,12,5,5) 430 1 450 i N(+)=Number of atoms in cluster subunit. (+)=(subunit,atom,atom type . 460 | Ljpar(+)=non-bonding parameter matrix.(+)=(atom,atom type,polarizatili\*.es 470 | and electrons and intermolecular distances). 480 | Fdir(+)=Force matrix. (+)=(subunit,component,force and direction 490 | indicator and last move direction indicator). 500 ! Mdir(+)=Moment(torque) matrix. (+)=(subunit,component,moment and direction 510 | indicator and last direction indicator). 520 | Tx(+),Ty(+),Tz(+)=Translation matrices. (+)=(subunit).

530 | Rr(+),Ry(+),Rr(+)=Rotation matrices. (+)=(subunit). 540 · Cg(+)=Center-of-mass matrix. (+)=(subunit,coordinates). 550 | Rstep(+)=Rotation step matrix. (+)=(subunit,direction). 560 = Tstep(+)=Translation step matrix. (+)=(subunit,direction). 570 ) T1(+/=Temporary translation step matrix. (+)=/suburit,direction . 530 ( R!(+)=Temporary rotation step matrix. (+)=(subunit,direction). 600 -610 DIM N(3,5,5),L;par(5,5,3) 620 DIM Fdir(3,3,3),Mdir(3,3,3) 630 DIM Tx(3) Ty(3) Tz(3) Rx(3) Ry(3) Rz(3) 540 DIM Cq(3,3), Rstep(3,3), Tstep(3,3), T1(3,3), R1(3,3) 650 GCLEAR 660 Look=7 : Default graphics display limits set at 7 angstroms. 670 Sol=! ! Default flag which draws solute and solvents. 680 Do=2 | Effective dielectric constant for monopole charge interaction. 690 BEEP 700 Q\$="0" 710 INPUT "Do you want expanded graphics?",Q\$ 720 IF QS="Y" THEN DUMP DEVICE IS 710, EXPANDED 730 IF QS="N" THEN DUMP DEVICE IS 710 740 IF Q\$<>"Y" AND Q\$<>"N" THEN 700 750 PRINT \*\* 770 PRINT \*\* Empirical vdW Cluster 780 PRINT \*\* • \* Configuration Energy 790 PRINT \*\* • • Minimization Program (ECCEMP) 800 PRINT \*\* • • • " 810 PRINT \*\* • \* 820 PRINT \*\* Version ECCEMP/2 830 PRINT \*\* by 840 PRINT "+ Joseph A. Menapace 850 PRINT \*\* 5 October 1986 850 PRINT \*\* 870 PRINT \*\* 880 PRINT \*\* Developed from ECEPP/2 890 PRINT ". G. Nemethy, M.S. Pottle, and H.A. Scheraga ... 900 PRINT "+ • " J. Phys. Chem. 87, 1883 (1983) 910 PRINT \*\* • " 920 PRINT "...... 930 1 950 / Main Menu: 960 1 970 | In Coef - Inputs non-bonding parameters from disc storage file LJPARAM or 980 ! a file created by the user. Subroutine used - Inputcf. 990 1 1000! In Coord - Inputs a cluster coordinate file from disc storage or from 1010) the keyboard. Subroutine used - Inputco. 10201 1030) Energy - Calculates the cluster binding energy at any geometry. Subroutine 10401 used - Energy. 10501 1050! Minimize - Performs cluster geometry and binding energy optimization. 1070 - Subroutine used - Minimize. 1080 | 1090 ( Move - Translates and rotates cluster subunits in three dimensions.

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1100 - Subroutine used - Hand. 1112 1 1120 : Draw - Draws the cluster geometry in four perspectives. Subroutire 1130 | used - Draw. 1140 1150 - NCA - Generates the intermolecular force field for the intermolecular (160 ) normal coordinate analysis performed by VDWNCA. Subroutine used - Eigen. 1170 1 1180 | Quit - Stops the subroutines and returns execution to the Main Menu. 1190 | Subroutine used - Quit. 1200 1 1210 ! Stor Con - Stores the cluster coordinates in user defined storage files. 1220 | Subroutine used - File. 1240 1 1250 ON KEY 0 LABEL "In Coef",3 GOSUB Inputcf 1260 ON KEY 1 LABEL "In Coord",3 GOSUB Inputco 1270 ON KEY 2 LABEL "Energy", 3 GOSUB Energy 1280 ON KEY 3 LABEL "Minimize",3 GOTO Minimize 1290 ON KEY 4 LABEL "Move",3 GOSUB Hand 1300 ON KEY 5 LABEL "Draw",3 GOSUB Draw 1310 ON KEY & LABEL "NCA",3 GOSUB Eigen 1320 ON KEY 9 LABEL "Quit",3 GOTO Quit 1330 ON KEY 7 LABEL "Stor Con",3 GOSUB File 1340 BEEP 1350 GOTO 1350 1360 1 1380 ! Subroutine: Inputcf 1390 1 1400 ! Inputs non-bonding parameter data file from disc storage or from 1410 ! keyboard. Parameters are stored in Ljpar(+). Parameters include polariza-1420 | bilities, number of electrons, and intermolecular distances for the atoms 1430 ! in question. 1450 1 1460 Inputcf: PRINTER IS 1 ! 1470 OUTPUT 2:CHR\$(255)&"K"; 1480 GOTO 1500 1490 CAT 1500 05="0" 1510 INPUT "Do you want to start another COEFICIENT data file?",Q\$ 1520 IF QS="N" THEN 1830 1530 IF Q\$<>"Y" THEN 1500 1540 ON ERROR 60TO 1490 1550 1 1570 ! This routine allows the user to enter and update data from the keyboard. 1590 1 1500 INPUT "What is your IN E W COEFICIENT file name?",Coef\$ 1610 CREATE BDAT Coefs&":INTERNAL".75.8 1620 OFF ERPOR 1630 GOSUB Atommenu 1640 INPUT "Select the atom." Atomis 1650 GOSUB Attpmenu 1660 INPUT "Select the atom type.",Attp1\$

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1670 GOSUB Atomnumbers 1580 IF Atom1=0 OR Attp1=0 THEN DISP "BAD SELECTION (1)) 1690 IF Atom1=0 OR Atto1=0 THEN WAIT 2 1700 IF Atomi=0 OR Attp1=0 THEN 1630 1710 INPUT "Enter the atom-atom polarizability (10°24 cm°3).",Ljpar-Atom),Atopl. 13 1720 INPUT "Enter the atom-atom effective electron number.",Ljpar(Atom),Attp1,20 1730 INPUT "Enter the atom-atom bond distance (Angstroms),",Ljpan(Atom1,Atip1,3) 1740 Q\$="Q" 1750 INPUT "Are you done?".Q\$ 1760 IF Q\$="Y" THEN 1790 1770 IF Q\$<>"N" THEN 1740 1780 GOTO 1630 1790 ASSIGN @Coef TO Coefs 1800 OUTPUT @CoefiLjpar(+) 1810 ASSIGN @Coef TO + 1820 GOTO 1910 1830 -1850 ! This routine inputs LJPARAM or user defined non-bonding parameter file 1860 ! from disc storage and displays it on the screen. 1880 ! 1890 ON ERROR GOTO 1490 1900 INPUT "What is your COEFICIENT data file name?",Coef\$ 1910 OUTPUT 21CHR\$(255)&"K"1 1920 PRINT "Your COEFICIENT data file name is",Coefs 1930 PRINT 1940 ASSIGN @Coef TO Coefs 1950 ENTER @Coef;Ljpar(+) 1960 ASSIGN @Coef TO . 1970 OFF ERROR 1980 PRINT USING 1990 1990 IMAGE "Atom",5X,"Type",6X,"Polar",4X,"Elect",4X,"Dist" 2000 PRINT USING 2010 2010 IMAGE 18X,"(10^24)",12X,"(Ang)" 2020 PRINT USING 2030 2030 IMAGE 18X,"(cm ^3)" 2040 PRINT 2050 PRINT 2050 FOR Atom1=1 TO 5 2070 FOR Attp1=1 TO 5 2080 GOSUB Atomnames 2090 IF Ljpar(Atom1,Attp1,1)=0 THEN 2120 2100 PRINT USING 2110(Atomis, Attp:s,Ljpar(Atomi, Attp:),L),Ljpar(Atom), Attp:, 2),Lj par(Atom1,Attp1,3) 2110 IMAGE 24,6X,5A,6X,2D.2D,4X,2D.2D,4X,2D.2D 2120 NEXT Attp1 2130 NEXT Atom1 2140 DISP "Here are your coeficients." 2150 WAIT 1 2160 1 2180 ! This routine allows for parameter changes and/or additions. 2200 1 2210 Q\$="Q"

2220 INPUT "Do you want to make changes or additions?",Q\$ 2230 IF 9\$="N" THEN 2250 2240 IF Q\$<>"Y" THEN 2210 2250 GOTO 1630 2260 BEEP 2270 OUTPUT 2; CHR\$(255)&"K"; 2290 RETURN 2290 1 2310 | Subroutine: Inputco 2320 1 2330 ! Inputs cluster coordinates, atomic charges, and hydrogen bonding flags 2340 | into coordinate matrices from disc storage or from the keyboard. 2360 1 2370 Inputco: ! 2380 OUTPUT 2:CHR\$(255)&"K"; 2390 GOTO 2410 2400 CAT 2410 Q\$="Q" 2420 INPUT "Do you want to start another COORDINATE file name?",Q\$ 2430 IF QS="N" THEN 3890 2440 IF Q\$<>"Y" THEN 2410 2450 ON ERROR GOTO 2400 2460 ! 2480 ! This routine allows the user to enter the coordinates, charges, and 2490 ! hydrogen bonding flags from the keyboard. 2510 1 2520 INPUT "What is your N E W COORDINATE file name?" Coords 2530 CREATE BDAT Coord\$&":INTERNAL",9400,8 2540 OFF ERROR 2550 INPUT "Enter the total number of molecules.",L 2560 IF L<1 OR L>3 THEN 2550 2570 INPUT "What ligand are you working on (1 for solute)?",K 2580 IF K<1 OR K>3 THEN 2570 2590 INPUT "Center of mass X coordinate?",Cq(K,1) 2500 INPUT "Center of mass Y coordinate?" ,Cq(K,2) 2610 INPUT "Center of mass Z coordinate?",Cq(K.3) 2620 GOSUB Atommenu 2630 INPUT "What atom coords are you inputting?",Atomis 2640 GOSUB Attpmenu 2650 INPUT "What is the atom type?",Attp1\$ 2650 605UB Atomnumbers 2670 IF Atomi=0 OR Attpl=0 THEN DISP "BAD SELECTION VEVY" 2680 IF Atom1=0 OR Attp1=0 THEN WAIT 2 2690 IF Atom1=0 OR Attp1=0 THEN 2620 2700 INPUT "How many?",N(K,Atom1,Attp1) 2710 FOR J=1 TO N(K,Atom1,Attp1) 2720 OUTPUT 2:CHR\$(255)&"K"; 2730 GOSUB Atomnames 2740 PRINT Attp:\$;" ":Atom:\$;" Number";J 2750 PRINT 2750 1 2780 / Inputs carbon coordinates, charges, and hydrogen bonding flags from the

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2790 ! keyboard. 28:0 1 2620 IF Atom1=1 THEN INPUT "Enter X coordinate.",Co(R,J,Attp1," -2830 IF Atom1=1 THEN PRINT "X Coordinate":Co(H.J.Attp1,1) 2840 IF Atom1=1 THEN INPUT "Enter Y coordinate.", Co(K,J,Attp1,2) 2850 IF Atom1=1 THEN PRINT "Y Coordinate";Co(K,J,Attp1,2) 2860 IF Atomi=1 THEN INPUT "Enter Z coordinate.",Co(K,J,Attp1,3) 2870 IF Atomi=1 THEN PRINT "Z Coordinate":Co(K,J,Attp1.3) 2880 IF Atom1=1 THEN INPUT "Enter atom charge.",Co(K,J,Attp1,4) 2890 IF Atomi=1 THEN PRINT "Atom Charge";Co(K,J,Attp1.4) 2900 05="0" 2910 IF Atom1=1 THEN INPUT "Is the atom a hydrogen bonder?",Q\$ 2920 IF Atom = 1 AND QS="N" THEN Co(K, J, Attp1, 5)=0 2930 IF Atom1=1 AND QS="Y" THEN Co(K,J,Attp1,5)=1 2940 IF Atom1=1 AND Q\$<>"Y" AND Q\$<>"N" THEN 2900 2950 IF Atomi=1 THEN PRINT "Hydrogen Bonding";Co(K,J,Attp1,5) 2960 ! 2980 | Inputs hydrogen coordinates, charges, and hydrogen bonding flags from the 2990 | keyboard. 3010 ! 3020 IF Atom1=2 THEN INPUT "Enter X coordinate.", Hh(K, J, Attp1, 1) 3030 IF Atom1=2 THEN PRINT "X Coordinate";Hh(K,J,Attp1,1) 3040 IF Atom1=2 THEN INPUT "Entor Y coordinate.", Hh(K,J,Attp1,2) 3050 IF Atom1=2 THEN PRINT "Y Coordinate";Hh(K,J,Attp1,2) 3050 IF Atom1=2 THEN INPUT "Enter 2 coordinate.", Hh(K,J,Attp1,3) 3070 IF Atom1=2 THEN PRINT "Z Coordinate";Hh(K,J,Attp1.3) 3080 IF Atom1=2 THEN INPUT "Enter atom charge.", Hh(K,J,Attp1,4) 3090 IF Atom1=2 THEN PRINT "Atom Charge"; Hh(K, J, Attp1, 4) 3100 Q\$="Q" 3110 IF Atom1=2 THEN INPUT "Is the atom a hydrogen bonder?".QS 3120 IF Atom1=2 AND Q\$="N" THEN Hh(K,J,Attp1,5)=0 3130 IF Atom1=2 AND QS="Y" THEN Hh(K, J, Attp1, 5)=1 3140 IF Atom1=2 AND Q\$ <> "Y" AND Q\$ <> "N" THEN 3100 3150 IF Atom1=2 THEN PRINT "Hydrogen Bonding";Hh(K,J,Attp1,5) 3160 ! 3180 | Inputs nitrogen coordinates, charges, and hydrogen bonding flags from the 3190 + keyboard. 3210 1 3220 IF Atomi=3 THEN INPUT "Enter X coordinate.", Nn(K, J, Attpi, 1) 3230 IF Atom1=3 THEN PRINT "X Coordinate";Nn(K,J,Attp1,1) 3240 IF Atom1=3 THEN INPUT "Enter Y coordinate.", Nn(K,J,Attp1,2) 3250 IF Atom1=3 THEN PRINT "Y Coordinate":Nn(K,J,Attp1,2) 3250 IF Atom!=3 THEN INPUT "Enter Z coordinate.", Nn(K, J, Attp1, 3) 3270 IF Atom1=3 THEN PRINT "Z Coordinate";Nn(K,J,Attp1,3) 3280 IF Atom)=3 THEN INPUT "Enter atom charge.",Nn(K,J,Attp1,4) 3290 IF Atom!=3 THEN PRINT "Atom Charge";Nn(K,J,Attp1,4) 3300 0\$="0" 3310 IF Atom1=3 THEN INPUT "Is the atom a hydrogen bonder?",Gs 3320 IF Atom!=3 AND QS="N" THEN Nn(K,J,Attp1,5)=0 3330 IF Atom1=3 AND Q\$="Y" THEN Nn(K,J,Attp1,5)=1 3340 IF Atom!=3 AND Q\$<>"Y" AND Q\$<>"N" THEN 3300 3350 IF Atomi=3 THEN PRINT "Hydrogen Bonding";Nn(K,J,Attp1,5)

3260 / 3330 | Inputs oxygen coordinates, charges, and hydrogen bonding flags from the 3390 · Feyboard. 3410 1 3420 IF Atom1=4 THEN INPUT "Enter X coordinate.",00(K,J,Attp1,i) 3430 IF Atomi=4 THEN PRINT "X Coordinate":00(K, J, Attp1, 1) 3440 IF Atomi=4 THEN INPUT "Enter Y coordinate.",00(K,J,Attp1,2) 3450 IF Atom1=4 THEN PRINT "Y Coordinate":00(K,J,Attp1.2) 3460 IF Atom!=4 THEN INPUT "Enter Z coordinate.", 00(K, J, Attp1, 3) 3470 IF Atom1=4 THEN PRINT "Z Coordinate":00(K.J.Attp1.3) 3480 IF Atom!=4 THEN INPUT "Enter atom charge.",00(K,J,Attp!,4) 3490 IF Atom!=4 THEN PRINT "Atom Charge":Oo(K,J,Attp!,4) 3500 Q\$="Q" 3510 IF Atom1=4 THEN INPUT "Is the atom a hydrogen bonder?", Q\$ 3520 IF Atom1=4 AND QS="N" THEN Oo(K,J,Attp1,5)=0 3530 IF Atom1=4 AND Q\$="Y" THEN Oo(K,J,Attp1,5)=1 3540 IF Atom1=4 AND Q\$<>"Y" AND Q\$<>"N" THEN 3500 3550 IF Atom1=4 THEN PRINT "Hydrogen Bonding" (Oo(K.J.Attp1.5) 3560 1 3580 | Inputs user defined coordinates, charges, and hydrogen bonding flags from 3590 ! the keyboard. 3610 1 3620 IF Atomi=5 THEN INPUT "Enter X coordinate.", Zn(K, J, Attpl, 1) 3630 IF Atom1=5 THEN PRINT "X Coordinate"; Zn(K.J.Attp1,1) 3640 IF Atom1=5 THEN INPUT "Enter Y coordinate.", Zn(K, J, Attp1, 2) 3650 IF Atom1=5 THEN PRINT "Y Coordinate" (Zn(K, J, Attp1, 2) 3660 IF Atom1=5 THEN INPUT "Enter Z coordinate.",Zn(K,J,Attp1,3) 3670 IF Atom1=5 THEN PRINT 'Z Coordinate';Zn(K,J,Attp1,3) 3680 IF Atomi=5 THEN INPUT "Enter atom charge.", Zn(K, J, Attpl, 4) 3690 IF Atom1=5 THEN PRINT "Atom Charge"; Zn(K,J,Attp1,4) 3700 QS="Q" 3710 IF Atom1=5 THEN INPUT "Is the atom a hydrogen bonder?",Q\$ 3720 IF Atom)=5 AND QS="N" THEN Zn(K,J,Attp1,5)=0 3730 IF Atom1=5 AND QS="Y" THEN Zn(K,J,Attp1,5)=1 3740 IF Atom1=5 AND Q\$<>"Y" AND Q\$<>"N" THEN 3700 3750 IF Atom1=5 THEN PRINT "Hydrogen Bonding"; Zn(K, J, Attp1, S) 3760 NEXT J 3770 Q\$="Q" 3780 INPUT "Are you done with this molecule?",QS 3790 IF QS="N" THEN 2620 3800 IF Q\$<>"Y" THEN 3770 3810 Q\$="Q" 3820 INPUT "Are you done inputting?",Q\$ 3830 IF QS="N" THEN 2570 3840 IF Q\$<>"Y" THEN 3810 3850 ASSIGN @Coord TO Coord\$ 3860 OUTPUT @Coord:L,Cq(+),N(+),Co(+),Hh(+),Nh(+),Oo(+),Zh(+) 3870 ASSIGN @Coord TO . 3880 GOTO 3970 3890 ON ERROR GOTO 2400 3900 1 3920 ! This routine inputs a user defined cluster coordinate file from

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3930 I disc storage and displays it on the screen. 3950 + 3960 INPUT "What is your COORDINATE data file name?",Coords 3970 OUTPUT 2:CHR\$(255)&"K"; 3980 PRINT "Your COCRDINATE data file name is",Coord\$ 3990 ASSIGN @Coord TO Coords 4000 ENTER @Coord:L,Cq(+),N(+),Co(+),Hh(+),Nn(+),Co(+),Zn(+) 4010 ASSIGN @Coord TO . 4020 OFF ERROR 4030 FOR K=1 TO L 4040 PRINT 4050 IF K=! THEN PRINT "Solute" 4060 IF K>1 THEN PRINT "Ligand ";K-1 4070 PRINT 4080 PRINT "Center of mass",Cq(K,1);Cq(K,2);Cq(K,3) 4090 PRINT 4100 FOR Atom1=1 TO 5 4110 FOR Attp1=1 TO 5 4120 FOR J=1 TO N(K Atom1 Attp1) 4130 GOSUB Atomnames 4140 IF Atomi=1 THEN PRINT Attp1\$;" "Atom1\$;J;Co(K,J,Attp1,1);Co(K,J,Attp1,2);C o(K.J.Attp1.3): 4150 IF Atomi=1 THEN PRINT Co(K, J, Attp1, 4) (Co(K, J, Attp1, 5) 4160 IF Atom1=2 THEN PRINT Attp1\$;" ";Atom1\$;J;Hh(K,J,Attp1,1);Hh(K,J,Attp1,2);H h(K.J.Attp1.3) 4170 IF Atom1=2 THEN PRINT Hh(K,J,Attp1,4);Hh(K,J,Attp1,5) 4180 IF Atomi=3 THEN PRINT Attpls: "iAtomisijiNn(K, J, Attpl, 1);Nn(K, J, Attpl, 2);N n(K,J,Attp1,3); 4190 IF Atomi=3 THEN PRINT Nn(K, J, Attp1, 4); Nn(K, J, Attp1, 5) 4200 IF Atom1=4 THEN PRINT Attp15: ";Atom15:J:00(K,J,Attp1,1):00(K,J,Attp1,2):0 o(K,J,Attp1,3); 4210 IF Atom1=4 THEN PRINT Oo(K, J, Attp1, 4): Oo(K, J, Attp1, 5) 4220 IF Atom1=5 THEN PRINT Attp15: ":Atom)5:J:Zn(K,J,Attp1,1):Zn(K,J,Attp1,2):Z n(K,J,Attp1,3); 4230 IF Atom1=5 THEN PRINT Zn(K,J,Attp1,4);Zn(K,J,Attp1,5) 4240 NEXT J 4250 NEXT Attp1 4260 NEXT Atom1 4270 NEXT K 4280 DISP "Here are your coordinates." 4290 WAIT 1 4300 05="0" 4310 1 4330 ! This routine allows for coordinate, charge, and hydrogen bonding flag 4340 1 changes and/or additions. 4360 1 4370 INPUT "Do you want to make changes or additions?",G\$ 4380 IF Q\$="N" THEN 5390 4390 IF Q\$<>"Y" THEN 4300 4400 05="0" 4410 INPUT "Do you want to add or change ENTIRE ligands?",Q\$ 4420 IF Q\$="N" THEN 4450 4430 IF QS- "Y" THEN 4400 4440 GOTO 2450

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4450 INPUT "Enter molecule number (1 for solute).",K
4460 IF KK1 OR KX3 THEN 4450
4470 Q$="Q"
4480 INPUT "Do you want to change the center of mass?",Q$
4490 IF Q$="N" THEN 4550
4500 IF Q$<>"Y" THEN 4470
4510 INPUT "Center of mass X coordinate?",Cq(K,1)
4520 INPUT "Center of mass Y coordinate?",Cq(K,2)
4530 INPUT "Center of mass Z coordinate?",Cq(K,3)
4540 6010 4300
4550 GUSUB Atommenu
4560 (NPUT "What atom coordinates?",Atom1$
4578 GOSUB Attpmenu
4580 INPUT "What is the atom type?",Attp1$
4590 INPUT "What number is the atom chosen?",J
4600 GOSUB Atomnumbers
4510 IF Atom1=0 OR Attp1=0 THEN DISP "BAD SELECTION !!!!"
4620 IF Atom1=0 OR Attp1=0 THEN WAIT 2
4630 IF Atom1=0 OR Attp1=0 THEN 4550
4640 OUTPUT 2:CHR$(255)&"K";
4650 GOSUB Atomnames
4660 PRINT Attp1$;" ";Atom1$;" Number";J
4670 Q$="Q"
4680 INPUT "Do you want to change the charge?",Q$
4690 IF QS="N" THEN 4810
4700 IF Q$<>"Y" THEN 4570
4710 IF Atom1=1 THEN INPUT "Enter atom charge.",Co(K,J,Attp1,4)
4720 IF Atom1=1 THEN PRINT "Atom Charge":Co(K,J,Attp1,4)
4730 IF Atom1=2 THEN INPUT "Enter atom charge.", Hh(K,J,Attp:,4)
4740 IF Atom1=2 THEN PRINT "Atom Charge";Hh(K,J,Attp1,4)
4750 IF Atom1≈3 THEN INPUT "Enter atom charge.",Nn(K,J,Attp1,4)
4750 IF Atom!=3 THEN PRINT "Atom Charge";Nn(K,J,Attp1,4)
4770 IF Atom1=4 THEN INPUT "Enter atom charge.",Oo(K,J,Attp1,4)
4780 IF Atom1=4 THEN PRINT "Atom Charge";00(K,J,Attp1,4)
4790 IF Atom!=5 THEN INPUT "Enter atom charge.",Zn(K,J,Attp!,4)
4800 IF Atom1=5 THEN PRINT "Atom Charge"; Zn(K, J, Attp1, 4)
4810 05="0"
4820 INPUT "Do you want to change hydrogen bonding capability?",Q$
4830 IF QS="N" THEN 4990
4840 IF Q$<>"Y" THEN 4810
4850 Q$="Q"
4850 INPUT "Is the atom a hydrogen bonder?",Q$
4870 IF QS="Y" AND Atom!=1 THEN Co(K, J, Attp1, 5)=1
4880 IF QS="Y" AND Atom1=2 THEN Hh(K,J,Attp1,5)=1
4890 IF QS="Y" AND Atom1=3 THEN Nn(K,J,Attp1,5)=1
4900 IF QS="Y" AND Atomi=4 THEN Qo(K,J,Attp1,5)=1
4910 IF QS="Y" AND Atom1=5 THEN Zn(K.J.Attp1.5)=1
4920 IF QS="Y" THEN 4990
4930 IF Q$<>"N" THEN 4850
4940 IF Atom1=1 THEN Co(K, J, Attp1, 5)=0
4950 IF Atom1=2 THEN Hh(K,J,Attp1,5)=0
4960 IF Atom1=3 THEN Nn(K, J, Attp1, 5)=0
4970 IF Atom1=4 THEN Oo(K, J, Attp1, 5)=0
4980 IF Atom1=5 THEN Zn(K, J, Attp1, 5)=0
4990 IF Atom1=1 THEN PRINT "Hydrogen Bonding":Co(K,J,Attp1,S)
5000 IF Atom1=2 THEN PRINT "Hydrogen Bonding";Hh(K,J,Attp1,S)
5010 IF Atom!=3 THEN PRINT "Hydrogen Bonding";Nn(K,J,Attp1,S)
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5020 IF Atom1=4 THEN FRINT "Hydrogen Bonding":00(K,J,Attp1.5) 5030 IF Atom)=5 THEN FRINT "Hydrogen Bonding (Inth.),Attp/.57 5040 Q\$="Q" 5050 INPUT "Do you want to change the coordinates?", Ca 5060 IF Q\$="N" THEN 4300 5070 IF Q\$<>"Y" THEN 5040 5080 IF Atomi=1 THEN INPUT "Enter X coordinate.",Co(E.J.Attpl.') 5050 IF Atomi≈! THEN PRINT "X Coordinate";Co(K,J,Attp1,1) 5100 IF Atomi=1 THEN INPUT "Enter Y coordinate.", Co(K,J,Attp1,2) 5110 IF Atom1=1 THEN PRINT "Y Coordinate":Co(K,J,Attp1,2) 5120 IF Atom1=1 THEN INPUT "Enter Z coordinate." (Co(K,J,Attp1,3) 5130 IF Atom1=1 THEN PRINT "Z Coordinate":Co(K,J,Attp1,3) 5140 IF Atom1=2 THEN INPUT "Enter X coordinate.", Hh(K,J,Atip1,1) 5150 IF Atom1=2 THEN PRINT "X Coordinate":Hh(K,J,Attp),1) 5160 IF Atom1=2 THEN INPUT "Enter Y coordinate.", Hh(K,J,Attp1,2) 5170 IF Atom1=2 THEN PRINT "Y Coordinate";Hh(K,J,Attp1,2) 5180 IF Atom1=2 THEN INPUT "Enter Z coordinate.", Hh(K,J,Attp1,3) 5190 IF Atomi=2 THEN PRINT "Z Coordinate"(Hh(K,J,Attp1,3) 5200 IF Atom!=3 THEN INPUT "Enter X coordinate.", Nn(K, J, Attp1, 1) 5210 IF Atom1=3 THEN PRINT "X Coordinate";Nn(K,J,Attp1,1) 5220 IF Atom1=3 THEN INPUT "Enter Y coordinate.", Nn(K,J,Attp1,2) 5230 IF Atom1=3 THEN PRINT "Y Coordinate"; Nn(K, J, Attp1, 2) 5240 IF Atom1=3 THEN INPUT "Enter 2 coordinate.", Nn(K,J,Attp1,3) 5250 IF Atom1=3 THEN PRINT "Z Coordinate"(Nn(K,J,Attp1,3) 5260 IF Atomi=4 THEN INPUT "Enter X coordinate.", 00(K, J, Attpl, 1) 5270 IF Atom1=4 THEN PRINT "X Coordinate":Do(K,J,Attp1,1) 5280 IF Atom!=4 THEN INPUT "Enter Y coordinate.", Oo(K,J,Attp1,2) 5290 IF Atom1=4 THEN PRINT "Y Coordinate"; Oo(K,J,Attp1,2) 5300 IF Atom1=4 THEN INPUT "Enter Z coordinate.", Oo(K,J,Attp1,3) 5310 IF Atom!=4 THEN PRINT "Z Coordinate":Oo(K,J,Attp:,3) 5320 IF Atom1=5 THEN INPUT "Enter X coordinate.", Zn(K,J,Attp1,1) 5330 IF Atom1=5 THEN PRINT "X Coordinate";Zn(K,J,Attp1,1) 5340 IF Atom1=5 THEN INPUT "Enter Y coordinate.", Zn(K,J,Attp1,2) 5350 IF Atom!=5 THEN PRINT "Y Coordinate";Zn(K,J,Attp1,2) 5360 IF Atom1=5 THEN INPUT "Enter Z coordinate.", Zn(K, J, Attp1, 3) 5370 IF Atom1=5 THEN PRINT "Z Coordinate"; Zn(K,J,Attp1,3) 5380 GOTO 4300 5390 ASSIGN @Coord TO Coords 5400 OUTPUT @CoordiL,Cq(+),N(+),Co(+),Hh(+),Nn(+),Oo(+),Zn(+) 5410 ASSIGN @Coord TO + 5420 BEEP 5430 OUTPUT 2:CHR\$(255)&"K": 5440 RETURN 5450 5470 ! Subroutine: Print 5480 ! 5490 | Prints the binding energy, forces, moments(torques), and coordinates for 5500 ! the optimized cluster geometry on the paper printer. 5520 1 5530 Print: 1 5540 ALPHA ON 5550 GRAPHICS OFF 5560 PRINT 5570 PRINT "Filename: ";Coord\$ 5580 PRINT

5590 PRINT 5600 IF Bod=1 AND Bhb=1 THEN PRINT "Potential=Lennard-Jones+Charges+Hydrogen Eur JING 5513 IF Bcc=1 AND Bhb=0 THEN PRINT "Potential=Lennard-Jones+Charges 5620 IF Boom#O AND Bhb=1 THEN PRINT "Potential#Lennard-Jones+Hydrogen Bonding 5630 IF Bcc=0 AND Bhb=0 THEN PRINT "Potential=Lennard-Jones" 5640 PRINT 5650 FOR K=1 TO L 5560 IF K#1 THEN PRINT "Solute" 5670 IF K>1 THEN PRINT "Ligand";K-1 5680 PRINT 5690 PRINT "Center of Mass=";Cq(K,1);Cq(K,2);Cq(K,3) 5700 PRINT 5710 FOR Atom1=1 TO 5 5720 FOR Attp1=1 TO 5 5730 FOR J=1 TO N(K,Atom1,Attpi) 5740 GOSUB Atomnames 5750 PRINT Attp1\$1" ":Atom1\$1 5750 IF Atom1=1 THEN PRINT Co(K, J, Attp1, 1);Co(K, J, Attp1, 2);Co(K, J, Attp1, 3); 5770 IF AtomI=1 THEN PRINT Co(K, J, Attp1, 4); Co(K, J, Attp1, 5) 5780 IF Atom1=2 THEN PRINT Hh(K,J,Attp1,1);Hh(K,J,Attp1,2);Hh(K,J,Attp1,3); 5790 IF Atom1=2 THEN PRINT Hh(K,J,Attp1,4);Hh(K,J,Attp1,5) S800 IF Atom1=3 THEN PRINT Nn(K,J,Attp1,1);Nn(K,J,Attp1,2);Nn(K,J,Attp1,3); 5810 IF Atom1=3 THEN PRINT Nn(K,J,Attp1,4):Nn(K,J,Attp1,5) 5820 IF Atom!=4 THEN PRINT 00(K,J,Attp1,1);00(K,J,Attp1,2);00(K,J,Attp1,3); 5830 IF Atom)=4 THEN PRINT Oo(K,J,Attp1,4):Oo(K,J,Attp1,5) 5840 IF Atom1=5 THEN PRINT Zn(K,J,Attp1,1);Zn(K,J,Attp1,2);Zn(K,J,Attp1,3); 5850 IF Atom1=5 THEN PRINT Zn(K,J,Attp1,4);Zn(K,J,Attp1,5) 5860 NEXT J 5870 NEXT Attp1 5880 NEXT Atom! 5890 PRINT 5900 NEXT K 5910 PRINT "Minimum Energy="(E 5920 PRINT 5930 FOR J=1 TO L 5940 IF J=1 THEN PRINT "Solute Forces "+"Fx="+Fdir(J,1,1)+"Fy="+Fdir(J,2,1)+"Fz= ":Fdir(J,3,1) 5950 IF J>1 THEN PRINT "Ligand";J-1; "Forces "; "Fx=";Edir(J,1,1); "Fy=";Edir(J,2,1 ); "Fz="; Fdir(J,3,1) 5960 IF J=1 THEN PRINT "Solute Moments ";"Mx=";Mdir(J,1,1);"My=";Mdir(J,2,1);"Mz =":Mdir(J.3.1) 5970 IF J>1 THEN PRINT "Ligand"+J-++ "Moments "+"Mx="+Mdin(J,1,1)+"My="+Fdin(J,2, 1); "Mz="; Mdir(J.3.1) 5980 PRINT 5990 NEXT J 5000 PRINT "Potential Curvature ";"Kxx=";Cvx;"Kyy=";Cvy;"Kzz=";Cvz 6010 PRINT 6020 PRINT 6030 PRINTER IS 1 6040 RETURN 6050 1 6070 | Subroutine: Energy 6080 1 5090 - Calculates the cluster binding energy, forces, and torques for a specific 5100 ' cluster geometry. Also calculates the intermolecular force field

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6110 / for VEWNCA. 5120 INCOMENTATION OF A DETAIL OF A 5130 1 6140 Energy: 1 6150 1 6170 4 Initialization of variables to start routine. 6190 1 6200 E=0 ! Cluster binding energy. 6210 Elj=0 ! non-bonding term energy. 5220 Eqq=0 ! Monopole charge term energy. 6230 Ehb=0 ! Hydrogen bonding term energy. 6240 Cvx=0 ! Curvature x=x direction. 5250 Cvy=0 ! Curvature y-y direction. 6260 Cvz=0 ! Curvature z-z direction. 5270 FOR J=1 TO L 6280 FOR Dir1=1 TO 3 6290 Fdir(J\_Dir1,1)=0 6300 Mdir(J,Dir1,1)≈0 6310 NEXT Dirl 6320 NEXT J **5330** Row=1 ! Control variables for intermolecular force field disc storage. **5340** Fa=75 ! routine. Force field is stored in file: H20EIG. 6350 ! 6370 ! Summation over all atom-atom interactions between solute and solvents for 6380 ! intermolecular potential calculation. 6400 ! 6410 FOR K1=1 TO L 6420 FOR C1=1 TO 5 6430 FOR A1=1 TO 5 6440 FOR J1=1 TO N(K1,C1,A1) 6450 FOR K2=2 TO L 6460 FOR C2=1 TO 5 6470 FOR A2=1 TO 5 6480 FOR J2=1 TO N(K2,C2,A2) 6490 IF K2<=K1 THEN 11630 6500 D=0 ! Intermolecular distance initialization for atoms in question. 6510 ! 6530 | Determines coordinates, charges, and hydrogen bonding flags for atoms 6540 ! involved in atom-atom potential terms. **6550** 6560 **5570 1113111311414444444** 6580 <sup>1</sup> Carbon coordinates, charges, and hydrogen bonding flags. 6600 1 6510 IF C1=1 THEN Coord11=Co(K1,J1,A1,1) 6620 IF C1=1 THEN Coord12=Co(K1, J1, A1, 2) 6630 IF C1=1 THEN Coord13=Co(K1, J1, A1, 3) 5640 IF C1=1 THEN Coord14=Co(K1,J1,A1,4) 6650 IF C1=1 THEN Coord15=Co(K1, J1, A1, 5) 5550 IF C2=1 THEN Coord21=Co(K2,J2,A2,1) 6670 IF C2=1 THEN Coord22=Co(K2,J2,A2,2)

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6580 IF C2=1 THEN Coord23=Co(K2, J2, A2, 3) 5590 IF C2\*) THEN Coord24=Co(K2,J2,A2,4) 6700 IF C2=1 THEN Coord25=Co(K2,J2,A2,5) 5710 1 6730 Hydrogen coordinates, charges, and hydrogen bonding flags. **5740** (CONTRACTOR CONTRACTOR CONTRA TOR CONTRA TOR CON 6750 / 6760 IF C1=2 THEN Coordil=Hh(K1,J1,A1,1) 6770 IF C1=2 THEN Coord12=Hh(K1, J1, A1, 2) 6780 IF C1=2 THEN Coord13=Hh(K1,J1,A1,3) 6790 IF C1=2 THEN Coord14=Hh(K1,J1,A1,4) 6800 IF C1=2 THEN Coord15=Hh(K1,J1,A1,5) 6810 IF C2=2 THEN Coord21=Hh(K2, J2, A2, 1) 6820 IF C2=2 THEN Coord22=Hh(K2, J2, A2, 2) 6830 IF C2=2 THEN Coord23=Hh(K2, J2, A2, 3) 6840 IF C2=2 THEN Coord24=Hh(K2, J2, A2, 4) 6850 IF C2=2 THEN Coord25=Hh(K2.J2.A2.S) 6860 1 5880 ! Nitrogen coordinates, charges, and hydrogen bonding flags. 6900 1 6910 IF C1=3 THEN Coord11=Nn(K1,J1,A1,1) 6920 IF C1=3 THEN Coord12=Nn(K1,J1,A1,2) 6930 IF CI=3 THEN Coord13=Nn(K1, J1, A1, 3) 6940 IF C1=3 THEN Coord14=Nn(K1,J1,A1,4) 5950 IF C1=3 THEN Coord15=Nn(K1,J1,A1,5) 5950 IF C2=3 THEN Coord21=Nn(K2,J2,A2,1) 6970 IF C2=3 THEN Coord22=Nn(K2, J2, A2, 2) 6980 IF C2=3 THEN Coord23=Nn(K2, J2, A2, 3) 6990 IF C2=3 THEN Coord24=Nn(K2,J2,A2,4) 7000 IF C2=3 THEN Coord25=Nn(K2, J2, A2, 5) 7010 1 7030 ! Oxygen coordinates, charges, and hydrogen bonding flags. 7050 7060 IF C1=4 THEN Coord11=0o(K1, J1, A1, 1) 7070 IF C1=4 THEN Coord12=Do(K1, J1, A1, 2) 7080 IF C1=4 THEN Coord13=0o(K1, J1, A1, 3) '7090 IF C1=4 THEN Coord14=0o(K1,J1,A1,4) 7100 IF C1=4 THEN Coord15=00(K1,J1,A1,5) 7110 IF C2=4 THEN Coord21=0o(K2.J2.A2.1) 7120 IF C2=4 THEN Coord22=0o(K2, J2, A2, 2) 7130 IF C2=4 THEN Coord23=00(K2, J2, A2, 3) 7140 IF C2=4 THEN Coord24=0o(K2, J2, A2, 4) 7150 IF C2=4 THEN Coord25=0o(K2, J2, A2, 5) 7160 1 7180 | User defined atom coordinates, charges, and hydrogen bonding flags. 7200 1 7210 IF C1=5 THEN Coord11=Zn(K1,J1,A1,1) 7220 IF C1=5 THEN Coord12=Zn(K1,J1,A1,2) 7230 IF C1=5 THEN Coord13=Zn(K1,J1,A1,3) 7240 IF C1=5 THEN Coord14=Zn(K1,J1,A1,4)

7750 IF C1=5 THEN Coord15=Zn(K1.J1.A1.5) 7250 IF 02=5 THEN Coord21=Zn(H2,J2,A2,1) 7270 IF 62=5 THEN Coord22=Zn(K2,J2,A2,2) 7230 IF 02=5 THEN Coord23=Zn(K2,J2,A2,3) 7290 IF C2=5 THEN Coord24=Zn(K2,J2,A2,4) 7300 IF C2=5 THEN Coord25=Zn(K2,J2,A2,5) 7310 1 7330 + Calculates atom-atom distances for atoms in question. 7350 1 7360 D=(Coord21-Coord11)^2+(Coord22-Coord12)^2+(Coord23-Coord13)^2 7370 Dr=01.5 7380 IF Dr=0 THEN 11600 7390 ! 7410 ! Calculates unit position vectors in each cartesian direction. 7430 1 7440 Dx=(Coord21-Coord11)/Dr 7450 Dv=(Coord22-Coord12)/Dr 7450 Dz=(Coord23-Coord13)/Dr 7470 1 7490 ! Calculates moment arm components in each cartesian direction. Moment arms 7500 ! are calculated relative to the cluster subunit center-of-mass. 7520 1 7530 Dmx2=Coord21-Cq(K2,1) 7540 Dmy2=Coord22-Cq(K2,2) 7550 Dmz2=Coord23-Cq(K2,3) 7560 Dmx1=Coord11-Cq(K1,1) 7570 Dmy1=Coord12-Cq(K1,2) 7580 Dmz1=Coord13-Cq(K1,3) 7590 1 7510 | Initialization of variables for routine looping. 7630 1 7640 Ehb=0 7650 Fhb=0 ! Force of hydrogen bonding interaction. 7650 Chb=0 + Curvature of hydrogen bonding interaction. 7670 Elj=0 7680 Flj=0 ! Force of non-bonding interaction. 7690 Clj=0 ! Curvature of non-bonding interaction. 7700 Eqq=0 7710 Eqg=0 ! Force of monopole charge interaction. 7720 Cqqv=0 | Curvature of monopole charge interaction. 7730 Ha=0 + Hydrogen bonding repulsive parameter. 7740 Hb=0 ! Hydrogen bonding attractive parameter. 7750 Aa=0 | Non-bonding repulsive parameter. 7750 Cc=0 | Non-bonding attractive parameter. 7770 1 7790 / Calculates atom-atom non-bonding parameters using the Slater-Kirkwood 7900 / approximation. 

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7820
7930 IF Ljpan(C),A),Z(=0 OR Ljpan(C2,A2,2)=0 THEN 7900
7640 Ljdenom=(Ljpan(C1,A1,1)+1.E-24/Ljpan(C1,A1,2))1.5+(Ljpan(C2,A2,1)+1.E-24 Lj
par(12,A2,2)) .5
7350 Ljnum=Ljpan(C1,A1,1)+1.E-24+Ljpan(C2,A2,1/+1.E-24
7850 Ljconst=3/2+4.803242E-12+1.054887E-27/(9.109534E-28)1.5+1.43881512E+12+1.E+
48+349.54
7970 Co=Ljconst+Ljnum/Ljdenom
7980 Ljrad=(Ljpar(C1,A1,3)+Ljpar(C2,A2,3))/2
7890 Aa=Ce/2+(Ljrad)16
7300 IF Coord15=0 OR Coord25=0 THEN 8560
7910 Ha=0
7920 Hb=0
7930 1
7950 | Selects hydrogen bonding parameters to use in calculations.
7970 1
7990 <sup>1</sup> Amin hydrogen/amin nitrogen hydrogen bonding parameters.
8010 1
8020 IF C1=2 AND A1=2 AND C2=3 AND A2=1 THEN Hb=8244+349.64
8030 IF C1=2 AND A1=2 AND C2=3 AND A2=1 THEN Ha=3.2897E+4+349.64
8040 IF C1=3 AND A1=1 AND C2=2 AND A2=2 THEN Hb=8244+349.64
8050 IF C1=3 AND A1=1 AND C2=2 AND A2=2 THEN Ha=3.2897E+4.349.54
8060 1
8080 ! Carb hydrogen/amin nitrogen hydrogen bonding parameters.
8100 1
8110 IF C1=2 AND A1=4 AND C2=3 AND A2=1 THEN Hb=8244+349.64
8120 IF C1=2 AND A1=4 AND C2=3 AND A2=1 THEN Ha=3.2897E+4+349.54
8130 IF C1=3 AND A1=1 AND C2=2 AND A2=4 THEN Hb=8244+349.64
8140 IF C1-3 AND A1=1 AND C2=2 AND A2=4 THEN Ha=3.2897E+4+349.64
8150 1
8170 <sup>1</sup> Amin hydrogen/carb oxygen hydrogen bonding parameters.
8190 1
8200 IF C1=2 AND A1=2 AND C2=4 AND A2=1 THEN Hb=4014+349.54
8210 IF C1=2 AND A1=2 AND C2=4 AND A2=1 THEN Ha=1.2040E+4+349.54
8220 IF C1=4 AND A1=1 AND C2=2 AND A2=2 THEN Hb=4014+349.64
9230 IF C1=4 AND A1=1 AND C2=2 AND A2=2 THEN Ha=1.2040E+4+349.64
8240 1
8260 - Hydr hydrogen/carb oxygen hydrogen bonding parameters.
8280 1
8290 IF C1=2 AND A1=4 AND C2=4 AND A2=1 THEN Hb=5783+349.64
8300 IF C1=2 AND A1=4 AND C2=4 AND A2=1 THEN Ha=1.3344E+4+349.64
8310 IF C1=4 AND A1=1 AND C2=2 AND A2=4 THEN HD=5783+349.64
8320 IF C1=4 AND A1=1 AND C2=2 AND A2=4 THEN Ha=1.3344E+4.349.54
8330 1
8350 / Amin hydrogen/hydr oxygen hydrogen bonding parameters.
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8370 ' 8380 IF 01=2 AND A1=2 AND 02=4 AND A2=2 THEN H5=2524+349.54 8390 IF C1=2 AND A1=2 AND C2=4 AND A2=2 THEN Ha=5.530E+3.349.54 8400 IF 01=4 AND A1=2 AND 02=2 AND A2=2 THEN H6=2524+349.64 3410 IF 01=4 AND A1=2 AND 02=2 AND A2=2 THEN Ha=5.890E+3+349.54 8423 4 8440 / Hydr hydrogen/hydr oxygen hydrogen bonding parameters. 8460 1 8470 IF C1=2 AND A1=4 AND C2=4 AND A2=2 THEN Hb=4610+349.64 8480 IF C1=2 AND A1=4 AND C2=4 AND A2=2 THEN Ha=1.122E+4+349.64 8490 IF C1=4 AND A1=2 AND C2=2 AND A2=4 THEN Hb=4610+349.64 8500 IF C1=4 AND A1=2 AND C2=2 AND A2=4 THEN Ha=1.122E+4+349.64 8510 1 8530 | Calculates hydrogen bonding interaction between atoms in question. 8550 1 8550 Ehb=8hb+(Ha/Dr^12-Hb/Dr^10) 8570 Fhb=Bhb+(12+Ha/Dr^13-10+Hb/Dr^11) 8580 Chb=8hb+(12+13+Ha/Or\*14-10+11+Hb/Or\*12) 8590 IF Ehb<>0 THEN 8730 8600 1 8620 | Calculates non-bonding interaction between atoms in question. 8640 1 8650 Elj=Aa/Dr^12-Cc/Dr^6 8650 Flj=12+Aa/Dr^13-6+Cc/Dr^7 8670 Clj=12+13+Aa/Dr^14-6+7+Cc/Dr^8 8680 1 8700 | Calculates monopole charge interaction between atoms in question. 8720 1 8730 Eqg=Bcc+Coord14+Coord24/Do/Dr+332+349.64 8740 Fqq=Bcc+Coord14+Coord24/Do/Dr^2+332+349.64 8750 Cqqv=Bcc+Coord14+Coord24+2/Do/Dr^3+332+349.64 8760 IF Elj=0 AND END=0 THEN DISP "NO PARAMETER IN THE POTENTIAL IIII" 8770 IF Elj=0 AND Ehb=0 THEN WAIT 2 8780 IF Elj=0 AND Enb=0 THEN GOTO Quit 8790 ! 8810 ! Determines total interaction energy, force, and curvature. 8830 1 8840 E=E+E1j+Eqq+Ehb 8850 Force=Flj+Fqq+Fhb 8860 IF Cutoff=0 THEN 8900 8870 IF Clj≤0 THEN Clj=0 8880 IF Coov<0 THEN Coov=0 8890 IF Chb<0 THEN Chb=0 8300 Curve=Clj+Cagv+Chb 8910 IF Eigen=0 THEN 11360 8920 1 

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8940 ' Generates intermolecular force field for VDWNCA and stores the field in 3950 ) file H2DEIG. The force field is calculated using the central force 8960 & approximation. The calculations are conducted in the cartesian operainate 8970 | system. 8980 1 8990 | Matel=Force constant matrix element in file H20EIG. 9000 1 Forcon=Intermolecular force constant for atom in question. 9020 1 9040 | Force constant in x-x direction. 9060 1 9070 ASSIGN @Dest TO "H2DEIG" 9080 CONTROL @Dest.5;(Column-1)\*Fa+Column 9090 ENTER @Dest;Matel 9100 CONTROL @Dest 5; (Column-1)+Fa+Column 9110 Forcon=Curve+(Dx^2) 9120 OUTPUT @Dest:Forcon+Matel 9130 CONTROL @Dest,5:(Row-1)\*Fg+Column 9140 ENTER @Dest(Matel 9150 CONTROL @Dest,5:(Row-1)\*Fg+Column 9160 Forcon=-Curve+(Dx^2) 9170 OUTPUT @Dest:Forcon+Matel 9180 CONTROL @Dest,5:(Column-1)\*Fg+Row 9190 ENTER @Dest:Matel 9200 CONTROL @Dest.5;(Column-1)+Fa+Row 9210 Forcon=-Curve+(Dx)^2 9220 OUTPUT @Dest;Forcon+Matel 9230 CONTROL @Dest,5:(Row-1)+Fg+Row 9240 ENTER @Dest:Matel 9250 CONTROL @Dest,5:(Row-1)\*Fg+Row 9260 Forcon=Curve+(Dx^2) 9270 1 9290 ! Force constant in x-y direction. 9310 1 9320 OUTPUT @Dest:Forcon+Matel 9330 CONTROL @Dest,5:(Column-1)\*Fg+Column+1 9340 ENTER @Dest;Matel 9350 CONTROL @Dest,5:(Column-1)\*Fg+Column+1 9360 Forcon=Curve+(Dx+Dy) 9370 OUTPUT @Dest:Forcon+Matel 9380 CONTROL @Dest\_5:(Column-1)\*Fa+Row+1 9390 ENTER @Dest:Matel 9400 CONTROL @Dest,5:(Column-1)\*Fg+Row+1 9410 Forcon=-Curve\*(Dx+Dy) 9420 OUTPUT @Dest:Forcon+Matel 9430 CONTROL @Dest,5:(Row-1)\*Fg+Column+1 9440 ENTER @Dest:Matel 9450 CONTROL @Dest,5:(Row-1)+Fg+Column+1 9460 Forcon=-Curve+(Ox+Dy) 9470 OUTPUT @Dest;Forcon+Matel 9480 CONTROL @Dest S: (Row-1)+Fg+Row+1 9490 ENTER @Dest;Matel 9500 CONTROL @Dest, 5: (Row-1)+Fg+Row+1

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9510 Forcen=Curve+(D<+Dy) 9520 1 9540 | Force constant in K-2 direction. 9560 1 9570 OUTPUT @Dest:Forcon+Matel 9580 CONTROL @Dest,5;(Column-1)\*Fg+Column+2 9590 ENTER @Dest;Matel 9500 CONTROL @Dest,5;(Column-1)\*Fg+Column+2 9510 Forcon=Curve+(Dx+Dz) 9620 OUTPUT @Dest:Forcon+Matel 9630 CONTROL @Dest,S:(Column-1)+Fg+Row+2 9640 ENTER @Dest:Matel 9650 CONTROL @Dest,5:(Column-1)\*Fg+Row+2 9660 Forcon=-Curve+(Dx+Dz) 9670 OUTPUT @Dest:Forcon+Matel 9680 CONTROL @Dest,5:(Row-1)+Fg+Column+2 9690 ENTER @Dest;Matel 9700 CONTROL @Dest,5:(Row-1)\*Fg+Column+2 9710 Forcon=-Curve+(Dx+Dz) 9720 OUTPUT @Dest:Forcon+Matel 9730 CONTROL @Dest ,5; (Row-1)+Fg+Row+2 9740 ENTER @Dest:Matel 9750 CONTROL @Dest,5;(Row-1)\*Fg+Row+2 9760 Forcon=Curve+(Dx+Dz) 9770 9780 9790 I Force constant in y-x direction. 9800 9810 1 9820 OUTPUT @Dest;Forcon+Matel 9830 CONTROL @Dest\_5;(Column+1-1)\*Fg+Column ENTER @Dest;Matel 9840 9850 CONTROL @Dest.5;(Column+1-1)\*Fg+Column 9850 Forcon=Curve+(Dy+Dx) 9870 OUTPUT @Dest:Forcon+Matel 9880 CONTROL @Dest,5:(Column+1-1)+Fg+Row 9890 ENTER @DestiMatel 9900 CONTROL @Dest,5:(Column+1-1)\*Fg+Row 9910 Forcon=-Curve+(Dy+Dx) 9920 OUTPUT @Dest;Forcon+Matel 9930 CONTROL @Dest,5:(Row+1-1)\*Fg+Column 9940 ENTER @DestiMatel 9950 CONTROL @Dest,5;(Row+1-1)+Fg+Column 9960 Forcon=-Curve\*(Dy\*Dx) 9970 OUTPUT @Dest:Forcon+Matel 9980 CONTROL @Dest,5:(Row+1-1)+Fg+Row 9990 ENTER @Dest;Matel 10000 CONTROL @Dest S:(Row+1-1)+Fg+Row 10010 Forcon=Curve+(Dy+Dx) 10020 1 10040 | Force constant in y-y direction. 10060 1 10070 OUTPUT @Dest;Forcon+Matel

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10080 CONTROL @Dest,5:(Column+1-1)\*Fg+Column+1 10090 ENTER @Dest;Matel 10100 CONTROL @Dest,5;(Column+1-1)+Fg+Column+1 10110 Forcon=Curve+(Dy^2) 10120 OUTPUT @Dest;Forcon+Matel 10130 CONTROL @Dest,5:(Column+1-1)\*Fg+Row+1 10140 ENTER @Dest:Matel 10150 CONTROL @Dest,5:(Column+1-1)+Fg+Row+1 10160 Forcon=-Curve+(By^2) 10170 OUTPUT @Dest:Forcon+Matel 10180 CONTROL @Dest 5:(Row+1-1)+Fg+Column+1 10190 ENTER @Dest:Matel 10200 CONTROL @Dest,5;(Row+1-1)+Fg+Column+1 10210 Forcon=-Curve+(Dy^2) 10220 OUTPUT @Dest:Forcon+Matel 10230 CONTROL @Dest\_5:(Row+1-1)\*Fg+Row+1 10240 ENTER @Dest:Matel 10250 CONTROL @Dest,5:(Row+1-1)\*Fg+Row+1 10260 Forcon=Curve+(Dy^2) 10270 1 10290 | Force constant in y-z direction. 10310 1 10320 OUTPUT @Dest;Forcon+Matel 10330 CONTROL @Dest\_5;(Column+1-1)+Fg+Column+2 10340 ENTER @Dest:Matel 10350 CONTROL @Dest,5;(Column+1+1)\*Fg+Column+2 10360 Forcon=Curve+(Dy+Dz) 10370 OUTPUT @Dest:Forcon+Matel 10380 CONTROL @Dest,5:(Column+1+1)\*Fg+Row+2 10390 ENTER @Dest;Matel 10400 CONTROL @Dest,5:(Column+1-1)+Fg+Row+2 10410 Forcon=-Curve+(Dy+Dz) 10420 OUTPUT @Dest:Forcon+Matel 10430 CONTROL @Dest,5:(Row+1~1)\*Fg+Column+2 10440 ENTER @Dest;Matel 10450 CONTROL @Dest,5;(Row+1~1)+Fg+Column+2 10460 Forcon=-Curve\*(Dy\*Dz) 10470 OUTPUT @Dest:Forcon+Matel 10480 CONTROL @Dest\_5:(Row+1-1)\*Fg+Row+2 10490 ENTER @Dest:Matel 10500 CONTROL @Dest,5;(Row+1~1)+Fg+Row+2 10510 Forcon=Curve+(Dy+Dz) 10520 ! 10540 | Force constant in z-x direction. 10560 1 10570 OUTPUT @Dest;Forcon+Matel 10580 CONTROL @Dest.5:(Column+2-1)+Fa+Column 10590 ENTER @Dest;Matel 10600 CONTROL @Dest,5;(Column+2-1)\*Fg+Column 10510 Forcon=Curve+(Dz+Dx) 10520 OUTPUT @Dest:Forcon+Matel 10530 CONTROL @Dest,5:(Column+2-1)+Fg+Row 10640 ENTER @Dest;Matel

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10650 CONTROL @Dest 5;(Column+2-1)\*Fg+Row 10650 Forcon=-Curve+(D:+Ok) 10570 OUTPUT @Dest;Forcon+Matel 10580 CONTROL @Dest 5: (Row+2-1)+Fg+Column 10530 ENTER @Dest;Matel 10700 CONTROL @Dest,5;(Row+2-1)+Fg+Column 10710 Forcon=-Curve+(D2+Dx) 10720 OUTPUT @Dest:Forcon+Matel 10730 CONTROL @Dest,5;(Row+2-1)+Fg+Row 10740 ENTER @Dest;Matel 10750 CONTROL @Dest,S:(Row+2-1)+Fg+Row 10750 Forcon=Curve+(Dz+Dx) 10770 1 10790 ! Force constant in x+y direction. 10810 1 10820 OUTPUT @Dest:Forcon+Matel 10830 CONTROL @Dest,5;(Column+2-1)\*Fg+Column+1 10840 ENTER @Dest:Matel 10850 CONTROL @Dest,5;(Column+2-1)\*Fg+Column+1 10860 Forcon=Curve+(Dz+Dv) 10870 OUTPUT @Dest:Forcon+Matel 10880 CONTROL @Dest.5;(Column+2-1)+Fa+Row+1 10890 ENTER @Dest;Matel 10900 CONTROL @Dest .5; (Column+2-1)+Fa+Row+1 10910 Forcon=-Curve\*(Dz\*Dy) 10920 OUTPUT @Dest;Forcon+Matel 10930 CONTROL @Dest,5:(Row+2-1)+Fg+Column+1 10940 ENTER @Dest;Matel 10950 CONTROL @Dest,S:(Row+2-1)\*Fg+Column+1 10960 Forcon=-Curve+(Dz+Dy) 10970 OUTPUT @Dest:Forcon+Matel 10980 CONTROL @Dest,5:(Row+2-1)+Fg+Row+1 10990 ENTER @DestiMatel 11000 CONTROL @Dest\_5:(Row+2-1)\*Fg+Row+1 11010 Forcon=Curve+(Dz+Dy) 11020 11040 ! Force constant in z-z direction. 11060 1 11070 OUTPUT @Dest;Forcon+Matel 11080 CONTROL @Dest,S:(Column+2-1)+Fg+Column+2 11090 ENTER @Dest:Matel 11100 CONTROL @Dest,S:(Column+2-1)\*Fg+Column+2 11110 Forcon=Curve\*(Dz^2) 11120 OUTPUT @Dest;Forcon+Matel 11130 CONTROL @Dest,S:(Row+2-1)\*Fg+Column+2 11140 ENTER @Dest:Matel 11150 CONTROL @Dest,S:(Row+2-1)\*Fg+Column+2 11160 Forcon=-Curve+(Dz^2) 11170 OUTPUT @Dest:Forcon+Matel 11180 CONTROL @Dest 5: (Column+2-1)\*Fg+Row+2 11190 ENTER @Dest:Matel 11200 CONTROL @Dest 5:(Column+2-1)+Fa+Row+2

11210 Forcon=-Curve+(Dz)^2

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a share

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11220 DUTPUT @Dest:Forcor+Matel
11220 CONTROL @Dest,5:(Row+2-1)*Fg+Row+2
11240 ENTER @Dest;Matel
11250 CONTROL @Dest,5;(Row+2-1)+Fg+Row+2
11260 Forcon=Curve+(Dz^2)
11270 OUTPUT @Dest:Forcon+Matel
11280 Column=Column+3
11290 IF Column=Last THEN Column=Begin
11300 1
11320 I Calculates force and curvature components in the three cartesian
11330 ! directions.
11350 1
11360 Fdir(K2,1,1)=Fdir(K2,1,1)+Force+Dx
11370 Edir(K2,2,1)=Edir(K2,2,1)+Force+Dy
11380 Fdir(K2,3,1)=Fdir(K2,3,1)+Force+Dz
11390 Cvx=Cvx+Curve+Dx^2
11400 Cvy=Cvy+Curve+Dy^2
11410 Cvz=Cvz+Curve+Dz^2
11420 IF K1<>1 THEN Cvx=Cvx+Curve+Ox^2
11430 IF K1<>1 THEN Cvy=Cvy+Curve+Dy^2
11440 IF K1<>} THEN Cvz=Cvz+Curve+Dz^2
11450 IF KI<>1 THEN Fdir(KI,I,I)=Fdir(KI,I,I)-Force+Dx
11460 IF K1<>1 THEN Fdir(K1,2,1)=Fdir(K1,2,1)-Force+Dy
11470 IF K1<>1 THEN Fdir(K1,3,1)=Fdir(K1,3,1)-Force+Dz
11480 1
11500 / Calculates the moments (torques) about the cartesian axes. Moments are
11510 ! determined relative to the cluster subunit center-of-mass.
11530
11540 Mdir(K2,1,1)=Mdir(K2,1,1)+(Dmy2*Force*Dz-Dmz2*Force*Dy)
11550 Mdir(K2,2,1)=Mdir(K2,2,1)+(Dmz2*Force*Dx-Dmx2*Force*Dz)
11560 Mdir(K2,3,1)=Mdir(K2,3,1)+(Dmx2+Force+Dy-Dmy2+Force+Dx)
11570 IF K1<>1 THEN Mdir(K1,1,1)=Mdir(K1,1,1)+(Dmy1+(-Force)+Dz-Dmz1+(-Force)+Dy
11580 IF K1<>1 THEN Mdir(K1,2,1)=Mdir(K1,2,1)+(Dmz1+(-Force)+Dx-Dmx1+(-Force)+Dz
11590 IF K1<>1 THEN Mdir(K1.3.1)=Mdir(K1.3.1)+(Dmx1+(-Force)+Dy-Dmy1+(-Force)+Dx
11600 NEXT J2
11610 NEXT A2
11620 NEXT C2
11630 NEXT K2
11640 Row=Row+3
11650 NEXT J1
11660 NEXT A1
11670 NEXT C1
11580 NEXT K1
11690 ASSIGN @Dest TO +
11700 IF Eigen=1 THEN RETURN
11710 IF Done=1 THEN RETURN
11720 IF Hand1=0 THEN OUTPUT 2;CHR$(255)&"K";
11730 1
11750 - Prints the binding energy, forces, moments, and curvature on the screen.
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11773 1 ∵;E 11782 PRINT "Energy 11790 PRINT 11300 FOR J=2 TO L 11810 PRINT "Forces on Ligand ";J-1;" (X,Y,Z)" 11820 PRINT Fdir(J,1,1); Fdir(J,2,1); Fdir(J,3,1) 11830 PRINT 11840 PRINT "Moments on Ligand ";J~1;" (X,Y,Z)" 11850 PRINT Mdir(J,1,1)(Mdir(J,2,1)(Mdir(J,3,1)) 11850 PRINT 11870 NEXT J 11880 PRINT "Potential Curvature (X,Y,Z)" 11890 PRINT CVX1CVV1CVZ 11900 PRINT 11910 RETURN 11920 1 11940 / Subroutine: Minimize 11950 1 11950 | Performs binding energy and geometry optimization by analyzing the 11970 / forces and moments on cluster subunits. 11990 1 12000 Minimize: PRINTER IS 1 12010 Eskio=1 12020 Passes=0 ! Initialize number of interations. 12030 Flag2=1 12040 OUTPUT 2:CHR\$(255)&"K": 12050 GRAPHICS OFF 12060 OFF KEY 12070 ! 12090 / Optimization Graphics Menu: 12100 1 12110 | Sketch Y - Activates cluster geometry drawing routine during 12120 ! optimization. Subroutine used - Sketchy. 12130 1 12140 | Sketch N - Deactivates cluster geometry drawing routine during 12150 | optimization. Subroutine used - Sketchn. 12160 1 12170 / Gelear - Clears the graphics dispay. Subroutine used - Wipe. 12180 1 12190 / Solute Y - Draws solute and solvent during optimization. Subroutine 12200 / used - Solutey. 12210 1 12220 ! Solute N - Draws solvent only during optimization. Subroutine 12230 ! used - Soluten. 12240 1 12250 / Quit - Stops optimization and returns program to Main Menu. Subroutine 2 sed - Quit. 12260 ' Subroutine used - Quit. 12270 / 12280 / Inc Pig - Decreases graphics display limits by 1 angstrom. Subroutire 12290 ! used - Incpic. 12300 1 12310 / Dec Pic - Increases graphics display limits by 1 angstrom. Subroutine

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12320 + used - Decpic.
12340 1
12350 ON KEY 0 LABEL "Sketch Y",3 GOSUB Sketchy
12350 ON KEY 5 LABEL "Sketch N",3 GOSUB Sketchn
12370 ON KEY 2 LABEL "Gclear",3 GOSUB Wipe
12380 ON KEY 3 LABEL "Solute Y",3 GOSUB Solutey
12390 ON KEY 8 LABEL "Solute N",3 GOSUB Soluten
12400 ON KEY 9 LABEL "Quit",3 GOTO Quit
12400 ON KEY 9 LABEL "Quit",3 GOTO Quit
12410 ON KEY 1 LABEL "Inc Pic",3 GOSUB Incpic
12420 ON KEY 5 LABEL "Dec Pic",3 GOSUB Decpic
12430 !
12450 | Initialization of variables before optimization.
12470 1
12480 FOR J=1 TO L
12490 Tx(J)=0
12500 Ty(J)=0
12510 Tz(J)=0
12520 Rx(J)=0
12530 Ry(J)=0
12540 Rz(J)=0
12550 FOR Dir1=1 TO 3
12560 FOR Dir2=1 TO 3
12570 Fdir(J_Dir1,Dir2)=0
12580 Mdir(J_Dir1_Dir2)=0
12590 NEXT Dir2
12500 NEXT Dir1
12610 NEXT J
12620 FOR J=1 TO L
12630 FOR Dir1=1 TO 3
12640 T1(J,Dir1)=1
12650 R1(J,Dir1)=1
12560 Tstep(J,Dir1)=0
12670 Rstep(J,Dirl)=0
12680 NEXT Dir1
12690 NEXT J
12700 Cutoff=0
12710 Bcc=0
12720 Bhb=0
12730 QS="Q"
12740 INPUT "Do you want to include charges?",Q$
12750 IF QS="Y" THEN Bcc=1
12750 IF QS="Y" THEN 12780
12770 IF Q$<>"N" THEN 12730
12780 0$="0"
12790 INPUT "Do you want to include hydrogen bonding?",Q$
12800 IF OS="Y" THEN Bhb=1
12810 IF QS="Y" THEN 12830
12820 IF Q$<>"N" THEN 12780
12830 Q$="Q"
12840 INPUT "Do you want to cut off the negative curvature?",Q$
12850 IF QS="Y" THEN Cutoff=1
12860 IF QS="Y" THEN 12880
12870 IF Q$<>"N" THEN 12830
12880 SET TIME 0
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12890 GOSUB Energy 12900 1 12920 > Determines force and moment direction change between current move 12930 F and last move to calculate translation and rotation step sizes for 12940 ! cluster subunits. 12960 1 12970 FOR J=2 TO L 12980 FOR Dir1=1 TO 3 12990 IF Fdir(J.Dir1.2)=0 THEN 13020 13000 IF ABS(Fdir(J,Dir1,2)-Fdir(J,Dir1,3))=2 THEN T1(J,Dir1)=T1(J,Dir1)+.2 13010 IF ABS(Fdir(J,Dir1,2)-Fdir(J,Dir1,3))<>2 THEN T1(J,Dir1)=T1(J,Dir1)-.05 13020 IF Mdir(J,Dir!,2)=0 THEN 13110 13030 IF ABS(Mdir(J,Dir1,2)-Mdir(J,Dir1,3))=2 THEN R1(J,Dir1)=R1(J,Dir1)+.2 13040 IF ABS(Mdir(J,Dir1,2)-Mdir(J,Dir1,3))<>2 THEN R1(J,Dir1)=R1(J,Dir1)=.05 13050 ! 13070 ! Determines translation and rotation step sizes. Maximum translation step 13080 ! size is .1 angstroms. Maximum rotation step size is 10 degrees. 13100 ! 13110 IF T1(J,Dir1)<1 THEN T1(J,Dir1)=1 13120 IF R1(J,Dir1)<1 THEN R1(J,Dir1)=1 13130 Tstep(J\_Dirl)=1/1000+10^(-.5+T1(J\_Dirl)+2.5) 13140 Rstep(J,Dir1)=1/10+10^(-.5+R1(J,Dir1)+2.5) 13150 Fdir(J,Dir1,3)=Fdir(J,Dir1,2) 13160 Mdir(J,Dir1,3)=Mdir(J,Dir1,2) 13170 NEXT Dir1 13180 NEXT J 13190 FOR S=2 TO L 13200 GOSUB Trans 13210 NEXT S 13220 605UB Energy 13230 Min=0 13240 Passes=Passes+1 13250 FOR J=2 TO L 13260 FOR Dir1=1 TO 3 13270 1 13290 ! Determines current force and moment directions and determines which 13300 ! degrees of freedom are optimized. Optimization is accomplished when 13310 / forces and torques are less that or equal to +/-.01. 13330 ! 13340 IF Fdir(J,Dir1,1)>0 THEN Fdir(J,Dir1,2)=1 13350 IF Fdir(J,Dir1,1)<0 THEN Fdir(J,Dir1,2)=-1 13360 IF ABS(Fdir(J,Dir1,1))<=.01 THEN Fdir(J,Dir1,2)=0 13370 IF Fdir(J,Dir1,2)=0 THEN Min=Min+1 13380 IF Mdir(J,Dir1,1)>0 THEN Mdir(J,Dir1,2)=1 13390 IF Mdir(J.Dir1.1)<0 THEN Mdir(J.Dir1.2)=-1 13400 IF ABS(Mdir(J\_Dir1,1))<=.01 THEN Mdir(J\_Dir1,2)=0 13410 IF Mdir(J,Dir1,2)=0 THEN Min=Min+1 13420 NEXT Dirl 13430 NEXT J 13440 1 

13460 + Determines if all degrees of freedom are optimized and if so terminates 13470 ' the routine. 13490 1 13500 IF Min=6+L-6 THEN 13520 13510 IF Sk=1 THEN GOSUB Draw 13520 IF Min=6+L-6 THEN GOSUB Draw 13530 IF Min=6+L-6 THEN 13550 13540 GOTO 12970 13550 BEEP 13560 DISP "Done 1111" 13570 Tim=(TIMEDATE MOD 86400)/60 13580 Done=1 13590 Eck=E 13600 Iter=1 13610 Handl=1 13620 1 13640 ! Checks final geometry for minimum energy by translating and rotating 13650 ! about the minimum energy cluster configuration. 13670 1 13680 FOR S=2 TO L 13690 FOR Xxx=1 TO 3 13700 Tstep(S,Xxx)=.001 13710 GOSUB 22000 13720 1 13740 ! Translation checks in three directions. 13760 13770 IF Xxx=1 AND E>=Eck THEN PRINT " +X Minimum"; 13780 IF Xxx=1 AND E<Eck THEN PRINT " +X Not Min"; 13790 IF Xxx=2 AND E>=Eck THEN PRINT " +Y Minimum"; 13800 IF Xxx=2 AND EKEck THEN PRINT " +Y Not Min": 13810 IF Xxx=3 AND E>=Eck THEN PRINT \* +Z Minimum\* 13820 IF Xxx+3 AND E<Eck THEN PRINT " +2 Not Min" 13830 Tstep(5,Xxx)=-.002 13840 GOSUB 22000 13850 IF Xxx=1 AND E>=Eck THEN PRINT " -X Minimum"; 13860 IF Xxx=1 AND EKEck THEN PRINT " -X Not Min": 13870 IF Xxx=2 AND E>=Eck THEN PRINT \* -Y Minimum\* 13880 IF Xxx=2 AND EKEck THEN PRINT " -Y Not Min" 13890 IF Xxx=3 AND E>=Eck THEN PRINT " -Z Minimum"; 13900 IF Xxx=3 AND E<Eck THEN PRINT " -Z Not Min"; 13910 Tstep(S,Xxx)=.001 13920 GOSUB 22000 13930 Rstep(S,Xxx)=.001 13940 GOSUB 22380 13950 13970 - Rotation checks in three directions. 13990 1 14000 IF Xxx=1 AND E = Eck THEN PRINT " +Rx Minimum 14010 IF Xxx=1 AND E'Eck THEN PRINT " +Rx Not Min" 14020 IF Xxx=2 AND E = Eck THEN PRINT " +Ry Minimum";

14030 IF Xxx=2 AND ECEck THEN PRINT " +Ry Not Min"; 14040 IF Xxx=3 AND EN=Eck THEN PRINT " +Rz Minimum"; 14050 IF Xxx=3 AND ErEck THEN PRINT " +R: Not Min"; 14060 Rstep(S,Xxx)=-.002 14070 GOSUB 22380 14080 IF Xxx=1 AND E>=Eck THEN PRINT " ~RA Minimum"; 14090 IF Xxx=1 AND E'Eck THEN PRINT " -Rx Not Min"; 14100 IF Xxx=2 AND E>=Eck THEN PRINT " -Ry Minimum"; 14110 IF Xxx=2 AND E Eck THEN PRINT " -Ry Not Min"; 14120 IF Xxx=3 AND E>=Eck THEN PRINT " -Rz Minimum" 14130 IF Xxx=3 AND EKEck THEN PRINT " -Rz Not Min" 14140 Rstep(\$,Xxx)=.001 14150 GOSUB 22380 14160 NEXT Xxx 14170 NEXT S 14180 Done=0 14190 Hand1=0 14200 Min=6+L-6 14210 Q\$="Q" 14220 ! 14240 ! Prepares calculation for paper printout and/or disc storage. 14260 ! 14270 INPUT "Do you want a printout?",Q\$ 14280 IF QS="N" THEN 14520 14290 IF Q\$<>"Y" THEN 14210 14300 Q\$="Q" 14310 INPUT "Do you want to store the configuration?",Q\$ 14320 IF QS="Y" THEN GOSUB File 14330 IF Q\$<>"N" AND Q\$<>"Y" THEN 14300 14340 PRINTER IS 710 14350 GOSUB Print 14360 PRINTER IS 710 14370 PRINT "Calculation Time (Min:Sec)=";INT(Tim);":";INT((Tim-INT(Tim))+60);" Iterations=":Passe 14380 PRINT 14390 Q\$="Q" 14400 INPUT "Do you want to change the drawing?",Q\$ 14410 IF QS="N" THEN 14470 14420 IF Q\$<>"Y" THEN 14390 14430 Flag2=0 14440 GOSUB Draw 14450 Flag2=1 14460 GOTO 14390 14470 DUMP GRAPHICS #710 14480 PRINT 14490 PRINT 14500 PRINT 14510 PRINTER IS 1 14520 Flag2=0 14530 Min=0 14540 OFF KEY 14550 GOTO 1250 14560 1 14580 | Subroutine: Trans

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14590 1 14600 | Translates and notates the cluster subunits in three dimensions. 14620 1 14630 Trans: ! 14640 1 14660 3 Determines the force and moment magnitudes. 14680 1 14690 Fmag=(Fdir(S,1,1)^2+Fdir(S,2,1)^2+Fdir(S,3,1)^2)^.5 14700 Mmag=(Mdir(S,1,1)^2+Mdir(S,2,1)^2+Mdir(S,3,1)^2)^.5 14710 IF Fmag=0 THEN 14800 14720 ! 14740 | Translates the cluster subunit screen parameters. 14760 1 14770 Tx(S)=Tx(S)+Fdir(S,1,2)+Tstep(S,1)+ABS(Fdir(S,1,1)/Fmag) 14780 Ty(S)=Ty(S)+Fdir(S,2,2)+Tstep(S,2)+ABS(Fdir(S,2,1)/Fmag) 14790 Tz(S)=Tz(S)+Fdir(S,3,2)+Tstep(S,3)+ABS(Fdir(S,3,1)/Fmag) 14800 IF Mmag=0 THEN 14890 14810 1 14830 ! Rotates the cluster subunit screen parameters. 14850 14860 Rx(S)=Rx(S)+Mdir(S,1,2)+Rstep(S,1)+ABS(Mdir(S,1,1)/Mmag) 14870 Ry(S)=Ry(S)+Mdir(S,2,2)+Rstep(S,2)+ABS(Mdir(S,2,1)/Mmag) 14880 Rz(S)=Rz(S)+Mdir(S,3,2)+Rstep(S,3)+ABS(Mdir(S,3,1)/Mmag) 14890 IF Rx(S)>=360 THEN Rx(S)=Rx(S)-360 14900 IF Rx(S)<=-360 THEN Rx(S)=Rx(S)+360 14910 IF Ry(S)>=360 THEN Rx(S)=Rx(S)-360 14920 IF Ry(S)<=-360 THEN Rx(S)=Rx(S)+360 14930 IF Rz(S)>=360 THEN Rx(S)=Rx(S)~360 14940 IF Rz(S)<=-360 THEN Rx(S)=Rx(S)+360 14950 IF Done=1 THEN 15080 14950 IF Hand1=1 THEN OUTPUT 2;CHR\$(255)&"K"; 14970 1 14990 ! Prints the translation and rotation of the cluster subunits on the 15000 ! screen display. 15020 1 15030 PRINT "Translation of Ligand":S-1 15040 PRINT USING "53D.4D";Tx(S);Ty(S);Tz(S) 15050 PRINT "Rotation of Ligand" (S-1 15060 PRINT USING "\$30.40";Rx(\$);Ry(\$);Rz(\$) 15070 PRINT 15080 IF Fmag=0 THEN 15500 15090 1 15110 / Translates the cluster subunit coordinates. 15130 1 15140 FOR C=1 TO 5 15150 FOR A=1 TO 5

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15'60 FOR J=1 TO N(S,C,A) 15170 FOR X=1 TO 3 15190 1 15200 / Translates carbon atoms. 15220 1 15230 IF C=1 THEN Co(S,J,A,X)=Co(S,J,A,X)+Fdir(S,X,2)+Tstep(S,X)+ABB(Fdir(B,K)) /Fmag) 15240 1 15260 - Translates hydrogen atoms. 15280 1 15290 IF C=2 THEN Hh(S,J,A,X)=Hh(S,J,A,X)+Fdir(S,X,2)+Tstep(S,X)+ABS(Fdir(S,X,1) /Fmag) 15300 1 15320 / Translates nitrogen atoms. 15340 1 15350 IF C=3 THEN Nn(S,J,A,X)=Nn(S,J,A,X)+Fdir(S,X,2)+Tstep(S,X)+ABS(Fdir(S,X,1) /Fmag) 15360 15380 ! Translates orygen atoms. 15400 1 15410 IF C=4 THEN 0o(S,J,A,X)=0o(S,J,A,X)+Fdir(S,X,2)+Tstep(S,X)+ABS(Fdir(S,X,1) /Fmag) 15420 1 15440 ! Translates user defined atoms. 15460 1 15470 IF C=5 THEN Zn(S,J,A,X)=Zn(S,J,A,X)+Fdir(S,X,2)+Tstep(S,X)+ABS(Fdir(S,X,1) /Fmac) 15480 NEXT X 15490 NEXT J 15500 NEXT A 15510 NEXT C 15520 15540 / Translates the cluster subunit center-of-mass. 15560 1 15570 FOR X=1 TO 3 15580 Cq(5,X)=Cq(5,X)+Fdir(5,X,2)+Tstep(5,X)+ABS(Fdir(5,X,1)/Fmap) 15590 NEXT X 15600 IF Mmag=0 THEN 16990 15610 Munitx=ABS(Mdir(S,1,1)/Mmag) 15520 Munity=ABS(Mdir(5,2,1)/Mmag) 15630 Munitz=ABS(Mdir(S,3,1)/Mmag) 15640 1 15660 ! Rotates the cluster subunits about their centers-of-mass. 

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15630 1 15690 FOR C=1 TO 5 15700 FOR A=1 TO 5 15710 FOR J=1 TO N(S.C.A) 15720 IF C=1 THEN 15820 15730 IF C=2 THEN 16060 15740 IF C=3 THEN 16300 15750 IF C=4 THEN 16780 15760 IF C=5 THEN 16540 15770 1 15790 | Rotates carbon atoms about molecular center-of-mass. 15810 1 15820 Col=Co(S,J,A,2)-Cq(S,2) 15830 Co2=Co(S,J,A,3)-Cq(S,3) 15840 Co(5,J,A,2)=Co1+CO5(Mdir(5,1,2)+Rstep(5,1)+Munitx)-Co2+SIN(Mdir(5,1,2)+Rst ep(S,1)\*Munitx) 15850 Co(5, J, A, 3)=Co1+SIN(Mdir(5, 1, 2)+Rstep(5, 1)+Munitx)+Co2+COS(Mdir(5, 1, 2)+Rst ep(S,1)+Munitx) 15860 Co(S,J,A,2)=Co(S,J,A,2)+Cq(S,2) 15870 Co(S,J,A,3)=Co(S,J,A,3)+Cq(S,3) 15880 Col=Co(S,J,A,3)-Cq(S,3) 15890 Co2=Co(S,J,A,1)-Cq(S,1) 15900 Co(S,J,A,3)=Co1+COS(Mdir(S,2,2)+Rstep(S,2)+Munity)-Co2+SIN(Mdir(S,2,2)+Rst ep(S,2)+Munity) 15910 Co(5, J, A, 1)=Co1+SIN(Mdir(5, 2, 2)+Rstep(5, 2)+Munity)+Co2+COS(Mdir(5, 2, 2)+Rst ep(S,2)+Munity) 15920 Co(S,J,A,3)=Co(S,J,A,3)+Cq(S,3) 15930 Co(S,J,A,1)=Co(S,J,A,1)+Cq(S,1) 15940 Co1=Co(S,J,A,1)-Cq(S,1) 15950 Co2=Co(S,J,A,2)-Cq(S,2) 15960 Co(S,J,A,1)=Co1+COS(Mdir(S,3,2)+Rstep(S,3)+Munitz)-Co2+SIN(Mdir(S,3,2)+Rst ep(S.3)+Munitz) 15970 Co(5,J,A,2)=Co1+5IN(Mdir(5,3,2)+Rstep(5,3)+Munitz)+Co2+CO5(Mdir(5,3,2)+Rst ep(S.3)#Munitz) 15980 Co(S,J,A,1)=Co(S,J,A,1)+Cq(S,1)15990 Co(S,J,A,2)=Co(S,J,A,2)+Cq(S,2) 16000 GOTO 16960 16010 1 16030 <sup>1</sup> Rotates hydrogen atoms about molecular center-of-mass. 16050 + 16060 Col=Hh(S,J,A,2)-Cq(S,2) 16070 Co2=Hh(S,J,A,3)-Cq(S,3) 15080 Hh(S,J,A,Z)=Co1+COS(Mdir(S,1,Z)+Rstep(S,1)+Munitx)-Co2+SIN(Mdir(S,1,Z)+Rst ep(S,1)+Munitx) 16090 Hh(S,J,A,3)=Co1+SIN(Mdir(S,1,2)+Rstep(S,1)+Munitx)+Co2+COS(Mdir(S,1,2)+Rst ep(S,1)+Munitx) 16100 Hh(S,J,A,2)=Hh(S,J,A,2)+Cq(S,2) 15110 Hh(S,J,A,3)=Hh(S,J,A,3)+Cq(S,3) 16120 Co1=Hh(S,J,A,3)-Cq(S,3) 16130 Co2=Hh(S,J,A,1)-Cq(S,1) 16140 Hh(S,J,A,3)=Co1+COS(Mdir(S,2,2)+Rstep(S,2)+Munity)-Co2+SIN(Mdir(S,2,2)+Ast ep(S,2)+Munity) 15150 Hh(S,J,A,1)=Col+SIN(Mdir(S,2,2)+Rstep(S,2)+Munity)+Co2+COS(Mdir(S,2,2)+Rst

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ep(S.21+Munity)
16160 HH: 5, J, A, 30 HH: 5, J, A, 30 HCa: 5, 30
16170 Hn(S,J,A,1)=Hh(S,J,A,1)+Ca(S,1)
15180 Col=Hh(S,J,A,1)-Ca(5,1)
16190 Co2=Hn(S,J,A,Z)-Cq(S,2)
15230 Hh(S,J,A,1)=Col+COS(Mdin(S,3,2)+Rstep(S,3)+Munitz)-Co2+SIN(Mdin(S,3,2)+Pst
ep(S,3)+Munitz)
16210 Hn(S,J,A,2)=001+SIN(Mdir(3,3,2)+Rstep(S,3)+Munitz)+602+605(Mdir(5,3,2)+Fst
ep(S,3)+Munitz)
16220 Hh(S,J,A,1)=Hh(S,J,A,1)+Cq(S,1)
16230 Hh(S,J,A,2)=Hh(S,J,A,2)+Cq(S,2)
16240 GOTO 16960
16250 +
15270 | Rotates nitrogen atoms about molecular center-of-mass.
16290 1
16300 Col=Nn(5,J,A,2)-Cq(5,2)
16310 Co2=Nn(5, J, A, 3)-Cq(5, 3)
16320 Nn(S,J,A,2)=Col+COS(Mdir(S,1,2)+Rstep(S,1)+Munitx)-Co2+SIN(Mdir(S,1,2)+Rst
ep(S,1)+Munitx)
15330 Nn(S,J,A,3)=Col+SIN(Mdir(S,L,2)+Rstep(S,1)+Munitx)+Co2+CO5(Mdir(S,L,2)+Rst
ep(S_1)+Munitx)
16340 Nn(S,J,A,2)=Nn(S,J,A,2)+Cq(S,2)
16350 Nn(S,J,A,3)=Nn(S,J,A,3)+Cq(S,3)
16360 Col=Nn(S,J,A,3)-Cq(S,3)
15370 Co2=Nn(S,J,A,1)-Cq(S,1)
16380 Nn(5,J,A,3)=Col+COS(Mdir(5,2,2)+Rstep(5,2)+Munity)-Co2+SIN(Mdir(5,2,2)+Rst
ep(S.2)+Munity)
16390 Nn(S,J,A,1)=Co1+SIN(Mdir(S,Z,Z)+Rstep(S,Z)+Munity)+Co2+COS(Mdir(S,Z,2)+Rst
ep(S,2)+Munity)
16400 Nn(S,J,A,3)=Nn(S,J,A,3)+Cq(S,3)
16410 Nn(S,J,A,1)=Nn(S,J,A,1)+Cq(S,1)
15420 Col=Nn(S,J,A,1)-Cq(S,1)
15430 Co2=Nn(S,J,A,2)-Cq(S,2)
16440 Nn(5,J,A,1)=Co1+COS(Mdir(5,3,2)+Rstep(5,3)+Munitz)-Co2+SIN(Mdir(5,3,2)+Rst
ep(S,3)+Munitz)
16450 Nn(S,J,A,Z)=Co1+SIN(Mdir(S,3,Z)+Rstep(S,3)+Munitz)+Co2+CO5(Mdir(S,3,2)+Rst
ep(S.3)+Munitz)
16460 Nn(S,J,A,1)=Nn(S,J,A,1)+Cq(S,1)
16470 Nn(S,J,A,2)=Nn(S,J,A,2)+Cq(S,2)
16480 GOTO 16960
15490
15510 ! Rotates user defined atoms about molecular center-of-mass.
16530
16540 Co1=Zn(S,J,A,2)-Cq(S,2)
15550 Co2=Zn(S,J,A,3)-Ca(S,3)
16560 7n(5,J,A,2)=Co1+COS(Mdir(5,1,2)+Rstep(5,1)+Munitk)=Co2+SIN(Mdir(5,1,2)+Rst
ep(S,1)+Munita)
16570 Zn(S,J,A,3)=Co1+SIN(Mdir(S,1,2)+Rstep(S,1)+Munitx)+Co2+COS(Mdir(S,1,2)+Ret
ep(S.1)+Munits)
16580 Zn(S,J,A,2)=Zn(S,J,A,2)+Cq(S,2)
16590 Zn(S,J,A,3)=Zn(S,J,A,3)+Cq(S,3)
16600 Col=Zn(S,J,A,3)~Ca(S,3)
16610 Co2=Zn(S,J,A,1)~Cq(S,1)
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16620 Zn(S,J,A,3)=Col+COS(Mdir(S,2,2)+Rstep(S,2)+Munity)-Co2+SIN(Mdir(S,2,2)+Rst ep(S,2)+Munity) 16630 Zn(S,J,A,1)=Co1+SIN(Mdir(S,2,2)+Rstep(S,2)+Munity)+Co2+CO5(Mdir(S,2,2)+Rst ep(S,2)+Munity) 16640 Zn(S,J,A,3)=Zn(S,J,A,3)+Cq(S,3) 16650 Zn(S,J,A,1)=Zn(S,J,A,1)+Cq(S,1) 16660 Col=Zn(S,J,A,1)-Cq(S,1) 16670 Co2=Zn(S,J,A,2)-Cq(S,2) 16680 Zn(5,J,A,1)=Col+COS(Mdir(5,3,2)+Rstep(5,3)+Munitz)-Co2+SIN(Mdir(5,3,2)+Rst ep(S,3)+Munitz) 16690 Zn(S,J,A,2)=Col+SIN(Mdir(S,3,2)+Rstep(S,3)+Munitz)+Co2+CO5(Mdir(S,3,2)+Rst ep(S,3)+Munitz) 16700 Zn(S,J,A,I)=Zn(S,J,A,I)+Cq(S,I)16710 Zn(S,J,A,2)=Zn(S,J,A,2)+Cq(S,2) 16720 GOTO 16960 16730 1 16750 ( Rotates oxygen atoms about molecular center-of-mass, 16770 ! 16780 Co1=0o(S,J,A,2)-Cq(S,2) 16790 Co2=0o(S,J,A,3)-Cq(S,3) 16800 Oo(S,J,A,2)=Co1+COS(Mdir(S,1,2)+Rstep(S,1)+Munitx)-Co2+SIN(Mdir(S,1,2)+Rst ep(S,1)+Munitx) 15810 0o(\$,J,A,3)=Col+SIN(Mdir(S,1,2)+Rstep(S,1)+Munitx)+Co2+COS(Mdir(S,1,2)+Rst ep(S,1)+Munitx) 16820 Oo(S,J,A,2)=Oo(S,J,A,2)+Cq(S,2) 15830 Oo(S,J,A,3)=Oo(S,J,A,3)+Cq(S,3) 16840 Co1=Oo(S,J,A,3)-Cq(S,3) 16850 Co2=Oo(S,J,A,1)-Cq(S,1) 16860 Oo(S,J,A,3)=Col+COS(Mdir(S,2,2)+Rstep(S,2)+Munity)-Co2+SIN(Mdir(S,2,2)+Rst ep(S.2)+Munity) 16870 Oo(S,J,A,1)=Co1+SIN(Mdir(S,2,2)+Rstep(S,2)+Munity)+Co2+COS(Mdir(S,2,2)+Rst ep(S,2)+Munity) 16880 Oo(S,J,A,3)=Oo(S,J,A,3)+Cq(5,3) 16890 Oo(S,J,A,1)=Oo(S,J,A,1)+Cq(S,1) 15900 Col=Oo(S,J,A,1)-Cq(S,1) 15910 Co2=Oo(S,J,A,2)-Cq(S,2) 16920 Oo(5,J,A,1)=Co1+COS(Mdir(5,3,2)+Rstep(5,3)+Munitz)-Co2+SIN(Mdir(5,3,2)+Rst ep(S,3)+Munitz) 15930 Oo(5,J,A,2)=Co1+SIN(Mdir(5,3,2)+Rstep(5,3)+Munitz)+Co2+COS(Mdir(5,3,2)+Rst ep(S.3)+Munitz) 16940 Do(S,J,A,1)=Oo(S,J,A,1)+Cq(S,1) 16950 Oo(S,J,A,2)=Oo(S,J,A,2)+Cq(S,2) 16960 NEXT J 16970 NEXT A 16980 NEXT C 16990 RETURN 17000 1 17020 ! Subroutine: Draw 17030 1 17040  $\pm$  Draws the cluster geometry in four perspectives: 3-dimensional, top view 17050 / end view, and side view. 17070 1 17080 Draw: 1

17090 IF Flag2=1 THEN 17400 17100 OFF KEY 17110 1 17130 / Drawing Menu: 17140 + 17150 - Gelear - Clears the graphics display. Subroutine used - Wipe. 17160 -17170 | Solute Y - Draws the solute and solvent. Subroutine used - Solutey. 17180 1 17190 ! Solute N - Draws the solvent only. Sibroutine used - Soluten. 17200 1 17210 ! Inc Pic ~ Decreases the graphics display limits by 1 angstrom. 17220 / Subroutine used - Incpic. 17230 1 17240 + Dec Pic ~ Increases the graphics display limits by 1 angstrom. 17250 | Subroutine used - Decpic. 17250 1 17270 - Draw - Draws the cluster geometries. 17280 1 17290 ! No Draw - Stops drawing at any time. 17310 1 17320 ON KEY 0 LABEL "Gclear",9 GOSUB Wipe 17330 ON KEY 1 LABEL "Solute Y",9 GOSUB Solutey 17340 ON KEY 5 LABEL "Solute N",9 GOSUB Soluten 17350 ON KEY 2 LABEL "Inc Pic",9 GOSUB Incpic 17350 ON KEY 7 LABEL "Dec Pic",9 GOSUB Decpic 17370 ON KEY 5 LABEL "Draw ",9 GOTO 17400 17380 ON KEY 9 LABEL "No Draw",9 GOTO 19240 17390 GOTO 17390 17400 GRAPHICS ON 17410 ALPHA OFF 17420 IF Min=L+6-6 THEN GINIT 17430 IF Min=L+6-6 THEN Sol=1 17440 FOR Pic=1 TO 4 17450 1 17470 ! 3 dimensional perspective. 17490 ! 17500 IF Pic=1 THEN VIEWPORT 0,66,0,50 17510 IF Pic=1 THEN Dwx=-10 17520 IF Pic=1 THEN Duz=15 17530 1 17550 | Top view perspective. 17570 ! 17580 IF Pic=2 THEN VIEWPORT 66,133,0,50 17590 IF Pic=2 THEN Dwx=0 17600 IF Pic=2 THEN Dwz=90 17610 1 17630 | End view perspective. 17550 1

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17660 IF Pic=3 THEN VIEWPORT 0,66,50,100
17570 IF Pic=3 THEN DWx=-90
17580 IF Pic=3 THEN Dwz=0
17690 -
17710 / Side view perspective.
17730 1
17740 IF Pic=4 THEN VIEWPORT 66,133,50,100
17750 IF Pic=4 THEN Dwx=0
17750 IF Pic=4 THEN Dwz=0
17770 SHOW -LOOK,LOOK,-LOOK,LOOK
17780 FRAME
17790 1
17810 | Draws cartesian axes on graphics display.
17830
17840 LINE TYPE 3
17850 Cf1(1)=-5+Look+COS(Dwx)
17860 Cel(2)=-5+Look+SIN(Dwx)
17870 Cf2(1)=5+Look+COS(Dwx)
17880 Ce2(2)=5*Look+SIN(Dwx)
17890 Cf1(3)=Cel(2)+SIN(Dwz)
17900 Cf2(3)=Ce2(2)+SIN(Dwz)
17910 MOVE Cf1(1),Cf1(3)
17920 DRAW Cf2(1),Cf2(3)
17930 Cft(1)=5+Look+SIN(Dwx)
17940 Cel(2)=-5+Look+COS(Dwx)
17950 Cf2(1)=-5+Look+SIN(Dwx)
17950 Ce2(2)=5+Look+COS(Dwx)
17970 Cf1(3)=Ce1(2)+SIN(Dwz)
17980 Cf2(3)=Ce2(2)+SIN(Dwz)
17990 MOVE Cf1(1),Cf1(3)
18000 DRAW Cf2(1),Cf2(3)
18010 Cf1(1)=0
18020 Cel(2)=0
18030 Cf2(1)=0
18040 Ce2(2)=0
18050 Cf1(3)=-5+Look+COS(Dwz)
18060 Cf2(3)=5+Look+COS(Dwz)
18070 MOVE Cf1(1), Cf1(3)
18080 DRAW Cf2(1)_Cf2(3)
18090 LINE TYPE 1
18100 1
18120 <sup>1</sup> Draws cluster geometry using balls and sticks.
18140 1
18150 FOR K=Sol TO L
18160 FOR C1=1 TO 5
18170 FOR A1=1 TO 5
18180 FOR J1=1 TO N(K,C1,A1)
18190 FOR C2=1 TO 5
18200 FOR A2=1 TO 5
18210 FOR J2=1 TO N(K,C2,A2)
18220 0=0
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18230 1 13250 - Determines atoms to be connected by bonds. 19270 1 18230 | Carbon atom coordinates. 18310 1 18320 IF C1=1 THEN Coord11=Co(K, J1, A1, 1) 18330 IF C1=1 THEN Coord12=Co(K, J1, A1, 2) 18340 IF C1=1 THEN Coord13=Co(K, J1, A1, 3) 18350 IF C2=1 THEN Coord21=Co(K, J2, A2, 1) 18360 IF C2=1 THEN Coord22=Co(K, J2, A2, 2) 18370 IF C2=1 THEN Coord23=Co(K, J2, A2, 3) 18380 | 18400 | Hydrogen atom coordinates. 18420 ! 18430 IF C1=Z THEN Coord11=Hh(K,J1,A1,1) 18440 IF C1=2 THEN Coord12=Hh(K, J1, A1, 2) 18450 IF C1=2 THEN Coord13=Hh(K, J1, A1, 3) 18460 IF C2=2 THEN Coord21=Hh(K, J2, A2, 1) 18470 IF C2=2 THEN Coord22=Hh(K, J2, A2, 2) 18480 IF C2=2 THEN Coord23=Hh(K, J2, A2, 3) 18490 ! 18510 ! Nitrogen atom coordinates. 18530 1 18540 IF C1=3 THEN Coord11=Nn(K, J1, A1, 1) 18550 IF C1=3 THEN Coord12=Nn(K, J1, A1, 2) 18560 IF C1=3 THEN Coord13=Nn(K, J1, A1, 3) 18570 IF C2=3 THEN Coord21=Nn(K, J2, A2, 1) 18580 IF C2=3 THEN Coord22=Nn(K, J2, A2, 2) 18590 IF C2=3 THEN Coord23=Nn(K, J2, A2, 3) 18600 1 18520 · Oxygen atom coordinates. 18640 1 18550 IF C1=4 THEN Coord11=0o(K,J1,A1,1) 18560 IF C1=4 THEN Coord12=0o(K, J1, A1, 2) 18570 IF C1=4 THEN Coord13=0o(K.J1.A1.3) 18680 IF C2=4 THEN Coord21=00(K, J2, A2, 1) 18590 IF C2=4 THEN Coord22=00(K, J2, A2, 2) 18700 IF C2=4 THEN Coord23=0o(K, J2, A2, 3) 18710 1 18730 / User defined atom coordinates. 18750 1 18760 IF C1=5 THEN Coord11=Zn(K,J1,A1,1) 13770 IF C1=5 THEN Coord12=Zn(K, J1, A1, 2) 18780 IF C1=5 THEN Coord13=Zn(K, J1, A1, 3) 13790 IF C2=5 THEN Coord21=Zn(K, J2, A2, 1)

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18800 IF C2=5 THEN Coord22=Zn(K, J2, A2, 2) 18810 IF C2=5 THEN Coord23=Zn(K,J2,A2,3) 18820 1 18840 | Determines ball sizes for specific atoms. 18860 1 18870 IF CT=1 THEN Siz=.3 18880 IF C2=2 THEN 512=.18 18890 IF C2=3 THEN Siz=.24 18900 IF C2=4 THEN S12=.28 18910 IF C2=5 THEN Siz=.32 18920 D=(Coord21-Coord11)^2+(Coord22-Coord12)^2+(Coord23-Coord13)^2 18930 Dr=D^.5 18940 ! 18960 | Determines Z dimensional projections for drawing 3 dimensional 18970 ! representations of clusters. 18990 ! 19000 IF C1=2 THEN GOTO 19190 19010 Cf1(1)=Coord11+COS(Dwx)-Coord12+SIN(Dwx) 19020 Ce1(2)=Coord11+SIN(Dwx)+Coord12+COS(Dwx) 19030 Cf2(1)=Coord21+COS(Dwx)-Coord22+SIN(Dwx) 19040 Ce2(2)=Coord21+SIN(Dwx)+Coord22+COS(Dwx) 19050 Cf1(3)=Ce1(2)+SIN(Dwz)+Coord13+COS(Dwz) 19060 Cf2(3)=Ce2(2)+SIN(Dwz)+Coord23+COS(Dwz) 19070 MOVE Cf1(1), Cf1(3) 19080 IF Dr>1.7 THEN 19140 19090 DRAW Cf2(1),Cf2(3) 19100 FOR Cir=0 TO 360 STEP 30 19110 IF Cir=0 THEN MOVE Cf2(1)+Siz+SIN(Cir),Cf2(3)+Siz+COS(Cir) 19120 DRAW Cf2(1)+51z+5IN(Cir),Cf2(3)+51z+CO5(Cir) 19130 NEXT Cir 19140 NEXT J2 19150 NEXT A2 19160 NEXT C2 19170 NEXT J1 19180 NEXT A1 19190 NEXT C1 19200 NEXT K 19210 NEXT Pic 19220 GRAPHICS OFF 19230 ALPHA ON 19240 IF Flag2=1 THEN 19410 19250 OFF KEY 19260 | 19280 / Sets up Main Menu for use after geometry and binding energy optimization 19290 <sup>1</sup> is complete. 19310 1 19320 ON KEY 0 LABEL "In Coef",3 GOSUB Inputcf 19330 ON KEY 1 LABEL "In Coord",3 GOSUB Inputco 19340 ON KEY 2 LABEL "Energy",3 GOSUB Energy 19350 ON HEY 3 LABEL "Minimize",3 GOTO Minimize 19360 ON KEY 4 LABEL "Move",3 GOSUB Hand

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19370 ON KEY 5 LABEL "Draw" ,3 GOSUB Draw 19350 ON KEY & LABEL "NCA",3 GOSUB Eigen 19390 ON KEY 9 LABEL "Quit",3 GOTO Quit 19400 ON KEY 7 LABEL "Stor Con" 3 GOSUB File 19410 RETURN 19420 -19440 | Subroutine: Atomnames 19450 19460 ! Assigns atomic symbols and atom types to the atoms composing the 19470 / clusters. Used in paper print and screen display routines. 19490 / 19500 Atomnames: 4 19510 ! 19530 / Assigns atomic symbols. 19550 19560 IF Atom1=1 THEN Atom1\$=" C" 19570 IF Atom1=2 THEN Atom1\$=" H" 19580 IF Atom1=3 THEN Atom1\$=" N" 19590 IF Atom1=4 THEN Atom1\$=" 0" 19600 IF Atom1=5 THEN Atom1\$=" Zn" 19610 IF Atom1=1 THEN 19720 19520 IF Atom1=2 THEN 19780 19630 IF Atom1=3 THEN 19840 19640 IF Atom1=4 THEN 19900 19650 IF Atom1=5 THEN 19960 19660 RETURN 19670 19690 | Assigns atom types. 19710 19720 IF Attp1=1 THEN Attp1\$=" Alip" 19730 IF Attp1=2 THEN Attp1s=" Carb" 19740 IF Attp1=3 THEN Attp1\$=" Arom" 19750 IF Attp!=4 THEN Attp1\$=" YYYY" 19760 IF Attp1=5 THEN Attp1\$=" ZZZZ" 19770 RETURN 19780 IF Attp1=1 THEN Attp1S=" Alip" 19790 IF Attp1=2 THEN Attp1\$=" Amin" 19800 IF Attp:=3 THEN Attp:s=" Arom" 19810 IF Attp:=4 THEN Attp:s=" Carb" 19820 IF Attp1=5 THEN Attp1\$=" ZZZZ" 19830 RETURN 19840 IF Attp1=1 THEN Attp1\$=" Amin" 19850 IF Attp1=2 THEN Attp1\$=" WWWW" 19860 IF Attp:=3 THEN Attp:\$=" XXXX" 19870 IF Attp1=4 THEN Attp1\$=" YYYY" 19880 IF Attpl=5 THEN Attpl\$=" ZZZZ" 19890 RETURN 19900 IF Attp:=: THEN Attp:\$\*" Carb" 19910 IF Attpl=2 THEN Attpl\$=" Hydr" 19920 IF Attp1=3 THEN Attp15=" XXXX" 19930 IF Attp1=4 THEN Attp1\$=" YYYY"

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19940 IF Attp1=5 THEN Attp1\$=" ZZZZ" 19950 PETURN 19960 IF Attpl=1 THEN Attpls=" Alip" 19970 IF Attp1=2 THEN Attp1\$=" WWWW" 19980 IF Attp1=3 THEN Attp1\$=" XXXX" 19990 IF Attp1=4 THEN Attp1\$=" YYYY" 20000 IF Attp1=5 THEN Attp1\$=" ZZZZ" 20010 RETURN 20020 ! 20040 | Subroutine: Atomnumbers 20050 ! 20060 ! Assigns atoms and atom types to numbers for array element 20070 | identification. 20090 20100 Atomnumbers: 1 20110 Atom1=0 20120 Attp1=0 20130 ! 20150 ! Assign numbers to atoms. 20170 1 20180 IF Atom1s="C" THEN Atom1=1 20190 IF Atom1s="H" THEN Atom1=2 20200 IF Atom1S="N" THEN Atom1=3 20210 IF Atom1\$="0" THEN Atom1=4 20220 IF Atom1 \*\*\* Z" THEN Atom1=5 20230 IF Atom1\$="C" THEN 20340 20240 IF Atom15="H" THEN 20400 20250 IF Atom1\$="N" THEN 20460 20260 IF Atom1s="0" THEN 20520 20270 IF Atom: \$="Z" THEN 20580 20280 RETURN 20290 1 20310 / Assign numbers to atom types. 20330 ! 20340 IF Attp:s="ALIP" THEN Attp:=1 20350 IF Attp:s="CARB" THEN Attp:=2 20360 IF Attp1s="AROM" THEN Attp1=3 20370 IF Attp:s="YYYY" THEN Attp:=4 20380 IF Attp1\$="ZZZZ" THEN Attp1=5 20390 RETURN 20400 IF Attp1s="ALIP" THEN Attp1=1 20410 IF Attp:s="AMIN" THEN Attp:=2 20420 IF Attp:s="AROM" THEN Attp:=3 20430 IF Attols="CARB" THEN Attol=4 20440 IF Attp1\$="ZZZZ" THEN Attp1=5 20450 RETURN 20460 IF Attp:s="AMIN" THEN Attp:=) 20470 IF Attpl\$="WWWW" THEN Attpl=2 20480 IF Attp:s="XXXX" THEN Attp:=3 20490 IF Attp1s="YYYY" THEN Attp1=4 20500 IF Attp1\$="2222" THEN Attp1=5

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20510 RETURN
18520 IF Atto1$≠"CARB" THEN Atto1=1
20530 IF Attp:s="HYDR" THEN Attp:=2
20540 IF Attp1$="XXXX" THEN Attp1=3
20550 IF Attpls="YYYY" THEN Attpl=4
20550 IF Attp1$="2222" THEN Attp1=5
20570 RETURN
20580 IF Attols="ALIP" THEN Attol=1
20590 IF Attols="WWWW" THEN Attol=2
20600 IF Attp1$="XXXX" THEN Attp1=3
20510 IF Attp:s="YYYY" THEN Attp:=4
20520 IF Attp1$="ZZZZ" THEN Attp1=5
20630 RETURN
20640 1
20550 ! Subroutine: Atommenu
20670 |
20580 | Atom menu for parameter and coordinate input routines.
20700 1
20710 Atommenu: | Selects atoms
20720 OUTPUT 2:CHR$(255)&"K";
20730 PRINT
                         Atom Selection"
20740 PRINT
20750 PRINT
20750 PRINT "
                         C - Carbon"
20770 PRINT
20780 PRINT *
                         H - Hydrogen"
20790 PRINT
20800 PRINT "
                         N - Nitrogen"
20810 PRINT
20820 PRINT *
                         0 - Oxygen"
20830 PRINT
20840 PRINT *
                         Z - Zinc/Other"
20850 RETURN
20860 1
20880 | Subroutine: Attpmenu
20890 1
20900 ! Atom type menu for parameter and coordinate input routines.
20920 1
20930 Attpmenu: |
20940 OUTPUT 2; CHR$(255)&"K";
20950 IF Atom15="C" THEN 21000
20960 IF Atom1s="H" THEN 21130
20970 IF Atom15="N" THEN 21260
20980 IF Atom: $="0" THEN 21390
20990 IF Atom15= 2" THEN 21520
21000 PRINT -
                         Larbon Atom Types"
21010 PRINT -
21020 PPINT
21030 PRINT "ALIP - Tetranedral Aliphatic
21040 PPINT
21050 PRINT "CARB - Carbonyl, Carboxylic Acid, Carboxylate, or
                                                          Peptide
21050 PRINT
21070 PRINT "AROM - Anomatic on C=C"
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21220 PRINT 2:090 PRINT "YYYY - YYYY" 21100 PRINT 21110 PRINT "ZZZZ - ZZZZ" 21120 RETURN 21130 PRINT Hydrogen Atom Types" 21140 PRINT "\_ 21150 PRINT 21160 PRINT "ALIP - Aliphatic" 21170 PRINT 21180 PRINT "AMIN - Amine or 1(2) Degree Amide" 21190 PRINT 21200 PRINT "AROM - Aromatic or Sulfhydryl" 21210 PRINT 21220 PRINT "CARB - Hydroxyl or Carboxylic Acid" 21230 PRINT 21240 PRINT "ZZZZ - ZZZZ" 21250 RETURN 21260 PRINT " Nitrogen Atom Types" 21270 PRINT 21280 PRINT 21290 PRINT "AMIN - Amine or Amide" 21300 PRINT 21310 PRINT "WWWW - WWWW" 21320 PRINT 21330 PRINT "XXXX - XXXX" 21340 PRINT 21350 PRINT "YYYY - YYYY" 21360 PRINT 21370 PRINT "2222 - 2222" 21380 RETURN 21390 PRINT Oxygen Atom Types" 21400 PRINT 21410 PRINT 21420 PRINT "CARB - Carbonyl or Carboxylic Acid" 21430 PRINT 21440 PRINT "HYDR - Hydroxyl, Carboxylic Acid, or Ester" 21450 PRINT 21460 PRINT "XXXX - XXXX" 21470 PRINT 21480 PRINT "YYYY - YYYY" 21490 PRINT 21500 PRINT "2222 - 2222" 21510 RETURN 21520 PRINT -Zinc/Other Atom Types" 21530 PRINT -21540 PRINT 21550 PRINT "Alis - Aliphatic" 21550 PRINT 2:578 PRINT ់មាលាលា – សាសាសាសី 21580 PRINT CIESO PRINT x+\*\* - \*\*\*\*\*\* 21600 PRINT CIEIS PRIME 21528 PRINT 21620 PRINT 2222 - 2222 21640 RETURN

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21650 ! 21670 | Subroutine: Hand 21680 1 21690 ! Allows user to translate and rotate cluster subunits. 21710 1 21720 Hand: 1 21730 ON ERROR GOTO 21750 21740 OUTPUT 2:CHR\$(255)&"K"; 21750 INPUT "What molecule are you working on (1 for solute)?",S 21750 INPUT "Do you want to rotate or translate?",Rt\$ 21770 Hand1=1 21780 IF Rts="R" THEN 22250 21790 IF Rts="T" THEN 21850 21800 IF Rt\$<>"T" AND Rt\$<>"R" THEN 21760 21810 1 21830 ! Cluster subunit translation routine. 21850 ! 21850 INPUT "What direction for translation?" Dts 21870 IF Dt\$="X" THEN Xxx=1 21880 IF Dts="Y" THEN Xxx=2 21890 IF Dts="Z" THEN Xxx=3 21900 IF Dts<>"X" AND Dts<>"Y" AND Dts<>"Z" THEN 21860 21910 INPUT "What interval?", Tstep(S,Xxx) 21920 INPUT "How many iterations?", Iter 21930 Eskip=0 21940 Q\$="Q" 21950 INPUT "Do you want to calculate the energy?",Q\$ 21960 IF QS="Y" THEN Eskip=1 21970 IF QS="Y" THEN 21990 21980 IF Q\$<>"N" THEN 21940 21990 OFF ERROR 22000 FOR 6=1 TO Iter 22010 Fmag=1 22020 Mmag=1 22030 FOR J=1 TO L 22040 FOR Xx=1 TO 3 22050 FOR Y=1 TO 3 22060 Fdir(J,Xx,Y)=0 22070 Mdir(J,Xx,Y)=0 22080 NEXT Y 22090 NEXT Xx 22100 NEXT J 22110 Fdir(S,Xxx,1)=1 22120 Fdir(S,Xxx,2)=1 22130 GOSUB 14710 22140 IF Eskip=0 THEN 22160 22150 GOSUB Energy 22160 IF Done=1 THEN 22190 22170 GOSUB Draw 22180 NEXT 6 22190 GOTO 22570 22200 + 

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22220 - Cluster subunit rotation routine. 22240 + 22250 INPUT "What axis of rotation?",Dts 22260 IF Dts="X" THEN Xxx=1 22270 IF Dts="Y" THEN Xxx=2 22280 IF Dt\$="Z" THEN Xxx=3 22290 IF Dt\$<>"X" AND Dt\$<>"Y" AND Dt\$<>"Z" THEN 22250 22300 INPUT "What interval?",Rstep(S,Xxx) 22310 INPUT "How many iterations?", Iter 22320 Eskip=0 22330 Q\$="Q" 22340 INPUT "Do you want to calculate the energy?.",Q\$ 22350 IF QS="Y" THEN Eskip=1 22360 IF QS="Y" THEN 22380 22370 IF Q\$<>"N" THEN 22330 22380 FOR G=1 TO Iter 22390 Fmag=1 22400 Mmag=1 22410 FOR J=1 TO L 22420 FOR Xx=1 TO 3 22430 FOR Y=1 TO 3 22440 Fdir(J,Xx,Y)=0 22450 Mdir(J,Xx,Y)=0 22460 NEXT Y 22470 NEXT Xx 22480 NEXT J 22490 Mdir(S,Xxx,1)=1 22500 Mdir(S,Xxx,2)=1 22510 605UB 14710 22520 IF Eskip=0 THEN 22540 22530 GOSUB Energy 22540 IF Done=1 THEN 22570 22550 GOSUB Draw 22560 NEXT 6 22570 Hand1=0 22580 RETURN 22590 ! 22610 ! Subroutine: Eigen 22620 1 22630 ! Sets up the parameters for calculation and storage of the intermolecular 22640 | force field for use in VDWNCA. 22660 ! 22670 Eigen: 4 22680 OUTPUT 2:CHR\$(255)&"K": 22590 PRINT "Place in storage disc containing H20EIG or a disc with space to cre ate H2OEIG." 22700 PRINT 22710 PRINT "To create a new H20EIG, Type in the following command at this time: 22720 PRINT 22730 PRINT "CREATE BDAT (QUOTES)H20EIG:INTERNAL(QUOTES),7300,8" 22740 DISP "Press continue to proceed." 22750 PAUSE 22760 Eigen=}

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22770 INPUT "What matrix row does solvent begin?",Begin 22780 Column=Begin 22790 INPUT "What matrix row does solvent end?" Lastm 22800 Last=Lastm+1 22810 Q\$="Q" 22820 INPUT "Do you want charges?",Q\$ 22830 IF Q\$="Y" THEN Bcc=1 22840 IF Q\$<>"Y" AND Q\$(0 "N" THEN 22810 22850 Q\$="0" 22860 INPUT "Do you want hydrogen bonding?",Q\$ 22870 IF Q\$="Y" THEN Bhb=1 22880 IF Q\$<>"Y" AND Q\$<>"N" THEN 22850 22890 GOSUB Energy 22900 Eigen=0 22910 RETURN 22920 1 22940 ! Subroutine: File 22950 ! 22960 ! Stores the coordinates of a specific cluster geometry in a disc file. 22980 1 22990 File: ! 23000 Q\$="Q" 23010 ON ERROR GOTO 23000 23020 INPUT "Do you want to start another COORDINATE file?",Q\$ 23030 IF QS="N" THEN 23070 23040 IF Q\$<>"Y" THEN 23020 23050 INPUT "Enter your N E W COORDINATE file name.",Coord\$ 23060 CREATE BDAT Coord\$&":INTERNAL",9400.8 23070 60SUB 5390 23080 OFF ERROR 23090 RETURN 23100 1 23120 ! Subroutine: Quit 23130 ! 23140 ! Stops subroutine execution and returns program to Main Menu. 23160 1 23170 Quit: 1 23180 OFF KEY 23190 Flac2=0 23200 GOTO 1250 23210 1 23230 / Subroutine: Sketchy 23240 1 23250 3 Activated the graphics display during geometry and binding energy 23250 ! optimization. 23280 + 23290 Sketchy: 4 23300 S+=1 23310 DISP "Graphics activated." 23320 PETURN 23330 +

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13350 - Subroutine: Sketchn 23360 / 23370 + Deactivates the graphics display during geometry and binding energy 23380 / optimization. 23400 -23410 Sketchn: 1 23420 Sk=0 23430 DISP "Graphics deactivated." 23440 RETURN 23450 23470 | Subroutine: Wipe 23480 1 23490 <sup>1</sup> Clears the graphics display. 23510 1 23520 Wipe: ! 23530 GINIT 23540 DISP "Clear screen." 23550 RETURN 23560 1 23580 ! Subroutine: Solutey 23590 1 23600 ' Allows he user to draw the cluster solute and solvent. 23620 1 Z3630 Solutey: ! 23540 Sol=1 23650 DISP "Draw solvent and solute." 23660 RETURN 23670 ! 23590 | Subroutine: Soluten 23700 ! 23710 <sup>1</sup> Allows the user to draw the solvent only. 23730 . 23740 Soluten: 1 23750 Sol=2 23760 DISP "Draw solvent only." 23770 RETURN 23780 1 23800 ' Subroutine: Incpic 23810 -23820 ! Decreases the graphics display limits by 1 angstrom. 22840 : 13850 Incold: 4 23860 LOOK=LOOK-1 3970 IF Look 1 THEN Look=1 23860 ALPHA ON 23890 DISP "Graphics limits are ";Look;" angstroms. 23900 RETURN

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APPENDIX FOUR

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"VDWNCA"

20 / Program name: VDWNCA 30 1 40 / This program calculates van der Waals cluster intermolecular vibrational 50 F modes using a normal coordinate analysis (NCA). The NCA is conducted using 60 I the FG matrix method. The clusters are treated as "giant molecules" in the 70 | NCA for which the intra- and intermolecular motions are treated 80 I simultaneously. The intermolecular force field used in the calculations is 90 ! generated using ECCEMP2 and is stored on disc as H20EIG. The intramolecular 1001 force fields are taken as those of each cluster sumubit. Both the intra-110! and intermolecular force fields are treated using the central force and 120! harmonic oscillator approximations. 130 1 140 1 150 ! 160 ! NOTICE: This program will not run with documentation contained in program 170 ! if not more than 255 Kbyte RAM is available. 190 1 200 OPTION BASE 1 210 PRINTER IS 1 220 1 240 | Matrices used: 250 1 260 ! A(+)=Energy (FG) matrix. 270 | Evr(+)=Real eigenvalue matrix. 280 | Evi(+)=Imaginary eigenvalue matrix. 290 + Vecr(+)=Real eigenvector matrix. 300 ! Veci(+)=Imaginary eigenvector matrix. 310 | Indic(+)= Matrix diagonalization indicator matrix. 320 ! Mol(+)=Cluster coordinates matrix. 340 1 350 DIM A(75,75), Evr(75), Evi(75), Vecr(75,75), Veci(75,75), Indic(75), Mol(75,4) 360 DEG 370 VIEWPORT 0,134,0,100 380 LORG 5 390 Paper=1 400 Look=6 + Default graphics display limits set at 6 angstroms. 410 Dex=-45 i Default X axis orientation for graphics display. 420 Dwz=20 | Default Z exis orientation for graphics display. 430 Dwxy=1 + Default top view for force field graphics display. 430 450 ! Main Menu: 470 1 480 'LJ MAT - Inputs intermolecular force field into matrix A(\*) from disc 490 ' storage file H2GEIG. Subroutine used - Inply. 500 1 510 - OWN VAL - Allows the user to input F matrix elements from keyboard. 520 \* Subroutine used - Inpown.

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530 1 540 ( INT DATA - Inputs the intramolecular force fields into A(+) from data 550 / statements. Subroutine used - Inpdat. 550 1 570 | EIGVALUE - Diagonalizes the FG matrix to yield eigenvalues frequencies and 580 <sup>1</sup> eigenvector normal modes describing cluster motion. Subroutire 590 | used - Eigen. 602 1 610 ! OUTPUT - Print the NCA results on the screen or on a paper printer. 620 | Subroutine used - Output. 630 1 640 ! INITIALI - Initializes the A(+) matrix and stores it in H20EIG. Used when 650 ! generating a new force field. Subroutine used - Zero. 670 ! 580 ON KEY 0 LABEL "LJ MAT ",1 GOSUB Inplj 590 ON KEY 1 LABEL "OWN VAL ",1 GOSUB Inpown 700 ON KEY 2 LABEL "INT DATA",1 GOSUB Inpdat 710 ON KEY 3 LABEL "EIGVALUE",1 GOSUB Eigen 720 ON KEY 7 LABEL "OUTPUT ".4 GOSUB Output 730 ON KEY 9 LABEL "INITIALI", 1 GOSUB Zero 740 BEEP 750 GOTO 750 760 1 780 ! Subroutine: Inplj 790 1 800 ! Inputs the intermolecular force field into A(\*) from disc storage file -810 ! H20EIG. 830 ! 840 Inplj: ! 850 ASSIGN @Mat TO "H2OEIG" 850 ENTER @Mat;A(+),Mdim 870 ASSIGN @Mat TO + 880 BEEP 890 RETURN 900 1 920 | Subroutine: Inpown 930 1 940 | Inputs F matrix elements from the keyboard into A(+). 960 1 970 Inpown: 1 980 INPUT "ENTER MATRIX DIMENSION", Mdim 990 INPUT "ENTER ROW, COLUMN", N.M. 1000 INPUT "ENTER ELEMENT", A(N,M) 1010 PRINT "ELEMENT ROW":N:"COLUMN":M: =":A(N,M) 1020 INPUT "ARE YOU DONE",Q\$ 1030 IF 0\$="Y" THEN 1050 1040 GOTO 990 1050 BEEP 1060 RETURN 1070 1 

1090 | Subroutine: Inpdat 1100 + 1110 | Inputs the intramolecular force fields from data statements. Trese force 1120 ) fields are added to the intermolecular force field placed in  $A(\star)_{\star}$ 1140 1 1150 Inpdat: | 1160 WINDOW -Look, Look, -Look, Look 1170 GCLEAR 1180 RESTORE 1190 INPUT "ENTER MATRIX DIMENSION" Mdim 1200 READ Numberdat 1210 FOR Y=1 TO Numberdat 1220 1 1240 ! Reads coordinates and force constants from data statements. 1260 1 1270 READ X1.Y1.Z1 1280 READ X2.Y2.Z2 1290 READ Forc 1300 1 1320 ! Converts force constants from dynes/cm to wavenumbers/square angstrom. 1340 1 1350 Forc=Forc/3.E+10/6.626E-27/1.E+16 1360 1370 12112212212222231122222222 1380 | Reads matrix row and column from data statements. 1490 ! 1410 READ Row 1420 READ Col 1430 1 1450 / Calculates unit position vectors. 1470 1 1480 D=((X2-X1)^2+(Y2-Y1)^2+(Z2-Z1)^2)^.5 1490 Dx=(X2-X1)/D 1500 Dy=(Y2-Y1)/D 1510 Dz=(Z2-Z1)/D 1520 PRINT XILYHIZI 1530 PRINT X2: Y2:Z2 1540 PRINT DxiDyiDz 1550 PRINT Forc D 1560 PRINT Row.Col 1570 1 1590 <sup>1</sup> Draws the intramolecular force field. 1610 1 1520 GRAPHICS ON 1630 IF DWxy=1 THEN MOVE X1,Y1 1640 IF DWAY=1 THEN DRAW X2,Y2 1650 IF Dwxy=0 THEN MOVE X1.21

1550 IF CWKy=0 THEN DRAW X2,22 1570 1 1880 - Force constant in x-- direction. 1710 1 1720 A(Col,Col)=Forc+(Dx)^2+A(Col,Col) 1730 A(Row,Col)=-Forc+(Dx)^2+A(Row,Col) 1740 A(Cal.Row)=-Forc\*(Dx)^2+A(Cal.Row) 1750 A(Row Row)=Forc+(Dx)^2+A(Row Row) 1760 1 1780 | Force constant in x-y direction. 1800 1 1810 A(Col.Col+1)=Forc+(Dx+Dy)+A(Col,Col+1) 1820 A(Col,Row+1)=-Forc\*(Dx+Dy)+A(Col,Row+1) 1830 A(Row,Col+1)=-Forc\*(Dx+Dy)+A(Row,Col+1) 1840 1 1860 | Force constant in x-z direction. 1880 1 1890 A(Row,Row+1)=Forc+(Dx+Dy)+A(Row,Row+1) 1900 A(Col.Col+2)=Forc\*(Dx+Dz)+A(Col.Col+2) 1910 A(Col.Row+2)=-Forc+(Dx+Dz)+A(Col.Row+2) 1920 A(Row\_Col+2)=-Forc+(Dx+Dz)+A(Row\_Col+2) 1930 A(Row\_Row+2)=Forc+(Dx+Dz)+A(Row\_Row+2) 1940 / 1960 | Force constant in v-x direction. 1980 1 1990 A(Col+1,Col)=Forc+(Dy+Dx)+A(Col+1,Col) 2000 A(Col+1,Row)=-Forc+(Dy+Dx)+A(Col+1,Row) 2010 A(Row+1,Col)=-Forc+(Dy+Dx)+A(Row+1,Col) 2020 A(Row+1,Row)=Forc+(Dy+Dx)+A(Row+1,Row) 2030 1 2050 | Force constant in ymy direction. 2070 4 2080 A(Col+1,Col+1)=Forc+(Dy^2)+A(Col+1,Col+1) 2090 A(Col+1,Row+1)=-Forc+(Dy^2)+A(Col+1,Row+1) 2100 A(Row+1,Col+1)=-Forc+(Dy^2)+A(Row+1,Col+1) 2110 A(Row+1,Row+1)=Forc+(Cy^2)+A(Row+1,Row+1) 2120 1 2140 ' Force constant in ymp direction. 2150 1 2170 A(Col+1,Col+2)=Forc+(Dy+Dz)+A(Col+1,Col+2 2180 A(Col+1,Row+2)=-Forc+(Dy+Cz)+A(Col+1,Row+2 2190 A(Row+1,Col+2)==Forc+(Dy+Dz)+A(Row+1,Col+2 2200 A(Row+1,Row+2)=Forc+(Dy+Cz/+A(Row+1,Row+2) 2210 1 

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2020 | Force constant in z=k direction. 2250 1 2050 A(Col+2.Col)=Forc+(Dz+Dx)+A(Col+2.Col) 2273 A(Col+2,Row)==Forc+(D:+D+)+A(Col+2,Row) 2280 A(Row+2 Col)=-Forc+(Dz+0x)+A(Row+2,Col) 2290 A(Row+2,Row)=Forc+(Dz+D+)+A(Row+2,Row) 2300 1 2320 / Force constant in z-y direction. 2340 1 2350 A(Col+2,Col+1)=Forc\*(Dz\*Dy)+A(Col+2,Col+1) 2360 A(Col+2,Row+1)=-Forc+(Dz+Dy)+A(Col+2,Row+1) 2370 A(Row+2,Col+1)=-Forc+(Dz+Dy)+A(Row+2,Col+1) 2380 A(Row+2,Row+1)=Forc+(Dz+Dy)+A(Row+2,Row+1) 2390 1 2410 | Force constant in z-z direction. 2430 + 2440 A(Col+2,Col+2)=Forc+(Dz^2)+A(Col+2,Col+2) 2450 A(Co1+2, Pow+2)=-Forc+(Dz"2)+A(Co1+2, Row+2) 2450 A(Row+2,Col+2)=-Forc+(Dz^2)+A(Row+2,Col+2) 2470 A(Row+2,Row+2)=Forc+(Dz^2)+A(Row+2,Row+2) 2480 NEXT Y 2490 BEEP 2500 RETURN 2510 / 2520 References results and results 2530 - Subroutine: Eigen 2540 2550 Clagonalizes the F5 matrix. 1552 3578 2558 Eliger 0558 \$5.30 26.2 matrix and must bules it by the Preserv 1615 21.72 1948 - E N.+ M - .... LEND PRINT ENTER RIGH . . . **W** 4 see te tagt ene escar au ۰. रेण्ड्र दे के मंद्र · · · الم رد شهر مر و شر فر کر - . 22.0414.00 3 1.00 ; ٠, 4.2 . . . . . . . . · · · / ° .2 ÷ . . . . - .\* 0 . 32

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3370 | Subroutine used - Deck. 2260 H 3390 F INC I - Increases the z axis prientation by F degree. 3400 ' Subroutine used - Incz. 3410 + 3420 | DEC Z - Decreases the class orientation by 1 degree. 3430 | Subroutine used - Decz. 3440 1 3450 + DONE - Stops drawing the clusters. Subroutine used - Flag. 3470 1 3480 OFF KEY 3490 ON KEY 0 LABEL "INC X",5 GOSUB Incx 3500 ON KEY 5 LABEL "DEC X",5 GOSUB Deck 3510 ON KEY 1 LABEL "INC Z",5 GOSUB Incz 3520 ON KEY 6 LABEL "DEC I",5 GOSUB Decz 3530 ON KEY 9 LABEL "DONE",5 GOSUB Flag 3540 1 3550 ! Reads cluster subunit coordinates from data statements. 3580 ! 3590 FOR N=1 TO Md1m/3 3600 FOR M=1 TO 4 3610 READ Mol(N.M) 3520 NEXT M 3630 NEXT N 3540 Q\$="Q" 3650 INPUT "OUTPUT TO PRINTER",Q\$ 3660 IF Q\$="Y" THEN Paper=710 3570 IF QS="Y" THEN 3700 3680 Paper=1 3690 IF Q\$<>"N" THEN 3650 3700 PRINTER IS Paper 3710 1 3730 ! Prints eigenvalues. 3750 1 3760 PRINT 3770 PRINT "------- EIGENVALUES (cm-1) -----------3780 PRINT 3790 FOR N=1 TO Mdim STEP 3 3800 IMAGE K, A, X, MZ. 9DE, 3X, K, A, X, MZ. 9DE, 3X, K, A, X, MZ. 9DE 3810 IF Paper=710 THEN PRINT USING 3800:N.".".Evr(N),N+1.".".Evr(N+1),N+2.".".E. r(N+2) 3820 IF Paper=1 THEN PRINT USING 3830; Evr(N); Evr(N+1); Evr(N+2) 3830 IMAGE MZ. 9DE, MZ. 9DE, MZ. 9DE 3840 NEXT N 3850 PRINT 3860 Q\$="0" 3870 INPUT "DO YOU WANT EIGENVECTORS",QS 3880 IF QS="N" THEN 5080 3890 IF G\$<>"Y" THEN 3870 3900 INPUT "WHAT EIGENVECTOR DO YOU WANT",M 3310 IF M/0 OR M Mdim THEN 3900

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3920 ( 3940 / Prints eigenvector for selected eigenvalue. 3960 1 -----3980 PRINT 3990 PRINT "EIGENVECTOR FOR EIGENVALUE"; M; Evr(M) 4000 PRINT 4010 FOR N=1 TO Mdim STEP 3 4020 IF Paper=710 THEN PRINT USING 3800; N, ".", Vecr(N,M), N+1, ".", Vecr(N+1,M), N+2, ".".Vecr(N+2.M) 4030 IF Paper=1 THEN PRINT USING 3830;Vecr(N\_M);Vecr(N+1\_M);Vecr(N+2\_M) 4040 NEXT N 4050 PRINT 4060 IF Paper=710 THEN 3870 4070 GCLEAR 4080 Flag=0 4090 ! 4110 ! Draws cluster geometry and eigenvector diplacement vectors using a stick 4120 ! model. 4140 ! 4150 GRAPHICS ON 4160 ALPHA OFF 4170 GCLEAR 4180 VIEWPORT 0,123,0,100 4190 LORG 1 4200 SHOW -Look,Look,-Look,Look 4210 MOVE -Look,Look-1 4220 LABEL Dwx:Dwz:M:Evr(M) 4230 LORG 5 4240 LINE TYPE 3 4250 ! 4270 ! Draws cartesian axes on graphics display. 4290 1 4300 Cf1(1)=-5+Look+COS(Dwx) 4310 Cel(2)=-5+Look+SIN(Dux) 4320 Cf2(1)=5+Look+COS(Dwx) 4330 Ce2(2)=5+Look+SIN(Dwx) 4340 Cf1(3)=Ce1(2)=SIN(Dwz) 4350 Cf2(3)=Ce2(2)+SIN(Dwz) 4360 MOVE Cf1(1),Cf1(3) 4370 DRAW Cf2(1)\_Cf2(3) 4380 Cfl(1)=5+Look+SIN(Dwx) 4390 Cel(2)=-5+Look+COS(Dwx) 4400 Cf2(1)=-S+Look+SIN(Dwx) 4410 Ce2(2)=5+Look+COS(Dwx) 4420 Cf1(3)=Ce1(2)+SIN(Dwz) 4430 Cf2(3)=Ce2(2)+SIN(Dwz) 4440 MOVE Cf1(1),Cf1(3) 4450 DRAW Cf2(1),Cf2(3) 4460 Cf1(1)=0

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4470 Cel(2)=0
4480 Cf2(1)=0
4490 Ce2(2)≠0
4520 0f1(3)=-5+Look+COS(Dw2)
4510 Cf2(3)=5+Look+COS(Dwz)
4520 MOVE Cf1(1).Cf1(3)
4530 DRAW Cf2(1), Cf2(3)
4540 LINE TYPE 1
4550 1
4570 | Determines atoms in cluster subunits to be connected by bonds.
4590 1
4600 FOR C1=1 TO Md1m/3
4610 FOR C2=1 TO Mdim/3
4620 D=0
4630 Crd11=Mol(C1,1)
4640 Crd12=Mo1(C1,2)
4550 Crd13=Mol(C1,3)
4660 Crd21=Mol(C2,1)
4670 Crd22=Mol(C2,2)
4680 Crd23=Mol(C2,3)
4690 D=(Crd21-Crd11)^2+(Crd22-Crd12)^2+(Crd23-Crd13)^2
4700 Or=0^.5
4710 IF Mol(C1,4)=2 THEN 4820
4720 Cfl(1)=Crdll+COS(Dwx)-Crdl2+SIN(Dwx)
4730 Cel(2)=Crd11+SIN(Dwx)+Crd12+COS(Dwx)
4740 Cf2(1)=Crd21+COS(Dwx)-Crd22+SIN(Dwx)
4750 Ce2(2)=Crd21+SIN(Dwx)+Crd22+COS(Dwx)
4750 Cfl(3)=Cel(2)+SIN(Dwz)+Crd13+COS(Dwz)
4770 Cf2(3)=Ce2(2)+SIN(Dwz)+Crd23+COS(Dwz)
4780 MOVE Cf1(1),Cf1(3)
4790 IF Dr>1.6 THEN 4820
4800 DRAW Cf2(1),Cf2(3)
4810 LABEL "0"
4820 NEXT C2
4830 NEXT C1
4840
4850 | Draws displacement vectors on atoms.
4880 1
4890 FOR C1=1 TO Mdim/3
4900 Crd11=Mol(C1,1)
4910 Crd12=Mol(C1,2)
4920 Crd13=Mol(C1,3)
4930 Crd21=Mol(C1,1)+Vecr(3+C1-2,M)
4940 Crd22=Mol(C1,2)+Vecr(3+C1-1,M)
4950 Crd23=Mol(C1,3)+Vecr(3+C1,M)
4950 Cf1(1)=Crd11+COS(Dwx)-Crd12+SIN(Dwx)
4970 Cel(2)=Crd11+SIN(Dwx)+Crd12+COS(Dwx)
4980 Cf2(1)=Crd21+COS(Dwx)-Crd22+SIN(Dwx)
4990 Ce2(2)=Crd21+SIN(Dwx)+Crd22+COS(Dwx)
5000 Cf1(3)=Ce1(2)+SIN(Dwz)+Crd13+COS(Dwz)
5010 Cf2(3)=Ce2(2)+SIN(Dwz)+Crd23+COS(Dwz)
5020 MOVE Cf1(1),Cf1(3)
5030 DRAW Cf2(1),Cf2(3)
```

5040 LABEL "+" 5050 NEXT C1 5060 IF Flag=0 THEN 4080 5070 GOTO 3870 5080 Q\$="Q" 5090 INPUT "DO YOU WANT TO LOOK AGAIN",Q\$ 5100 IF QS="N" THEN 5130 5110 IF Q\$ >"Y" THEN 5090 5120 GOTO 3640 5130 PRINTER IS 1 5140 BEEP 5150 STOP 5160 RETURN 5170 ! 5190 ! Subroutine: Zero 5200 + 5210 ! Initializes the A(+) matrix and stores it in H20EIG. 5230 1 5240 Zero: ! 5250 FOR N=1 TO Mdim 5260 FOR M=1 TO Mdim 5270 A(N,M)=0 5280 NEXT M 5290 NEXT N 5300 INPUT "ENTER MATRIX ORDER", Mdim 5310 ASSIGN @Mat TO "H20EIG" 5320 OUTPUT @Mat:A(+),Mdim 5330 ASSIGN @Mat TO . 5340 BEEP 5350 RETURN 5360 1 5380 ! Subroutine: Incx 5390 1 5400 I Increases x axis orientation by 1 degree. 5410 5420 1 5430 Incx: 1 5440 Dwx=Dwx+1 5450 RETURN 5460 1 5480 | Subroutine: Decx 5490 ! 5500 <sup>1</sup> Decreases x axis orientation by 1 degree. 5520 1 5530 Decx: 1 5540 Dwx=0wx-1 5550 RETURN 5560 + 5580 | Subroutine: Incz 5590 1 5600 | Increases z axis orientation by 1 degree.

5615 1 5620 Incz: 1 5625 Dwz=Dwz+1 5630 RETURN 5635 4 5645 | Subroutine: Decz 5650 1 5655 | Decreases z axis orientation by 1 degree. 5665 1 5670 Decz: 1 5675 Dwz=Dwz-1 5680 RETURN 5685 ( 5695 | Subroutine: Flag 5700 1 5705 | Stops drawing cluster geometry and displacement vectors. 5715 1 5720 Flag: 1 5725 Flac=1 5730 RETURN 5735 5745 ! Intramolecular force fields for cluster subunits. 5750 1 5755 1 The force fields are generated using the central force approximation and 5760 | expressed as force constants in a cartesian coordinate system. The 5765 ! following data statements are for the intramolecular force fields in 5770 ! Benzene-(Methane)1. They are included to give an example on how the 5775 ! force fields are generated. Since the benzene subunit is planar, the 5780 ' central force field constants must be modified to incorporate 5785 ! out-of-plane motion. This is accomplished by setting the atoms composing 5790 | benzene slightly out-of-plane. One angstrom provides sufficient 5795 ! out-of-plane displacement to account for the out-of-plane force field. 5800 1 5805 1 The force fields are entered as force constants between adjacent atoms 5810 <sup>1</sup> and between atoms displaced by one intervening atom. The atoms in 5815 I each cluster subunit are numbered in the same order as used in the 5820 | ECCEMP2 calculation. The force constants used are those corresponding to 5825 <sup>1</sup> general functional group stretches and bends. 5830 1 5835 / 5840 | Order of data entries: 5845 1 5850 | DATA X-Coord, Y-Coord, Z-Coord for atom 1. 5855 ! DATA X-Coord, Y-Coord, Z-Coord for atom j. 5860 | DATA Force Constant, Atom i X-Coord Matrix Location, Atom j X-Coord Matri-5865 + Location. 5875 1 5880 DATA 52 | Number of force constants used in fields. 5985 1 

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5895 | Benzene C-C force constant. 5905 / 5910 DATA -1.395.0,0 5915 DATA -.6975.1.208.0 5920 DATA 6.9465,1,4 5925 ( 5935 | Benzene C-C force constant. 5945 ( 5950 DATA -1.395.0.0 5955 DATA .6975,1.208.0 5960 DATA 1.054E5.1.7 5965 1 5975 ! Benzene C-C force constant. 5985 1 5990 DATA -.6975.1.208.0 5995 DATA .6975,1.208,0 6000 DATA 6.9465.4.7 6005 1 6015 ! Benzene C-C force constant. 6025 1 6030 DATA -.6975,1.208,0 6035 DATA 1.395,0.0 6040 DATA 1.054E5,4,10 6045 6055 ! Benzene C-C force constant. 6065 ! 6070 DATA .6975.1.208.0 6075 DATA 1.395,0.0 6080 DATA 5.94E5.7.10 6085 1 6095 | Benzene C-C force constant. 6105 1 6110 DATA .6975,1.208,0 6115 DATA .6975,-1.208,0 6120 DATA 1.054E5,7,13 6125 1 6135 <sup>1</sup> Benzene C-C force constant. 6145 1 6150 DATA 1.395.0.0 6155 DATA .6975,-1.208,0 6150 DATA 6.9465,10,13 6165 1 5175 | Benzene C-C force constant.

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╋╬┑╒╊╬╕╪┇┥╸╊╬╕╱┫╘╖╱┫╘╖┙┇┝┙┚╞┝╌╒╞┑┙╞╬┱╱┟┑┙┠┟┱╱┟╖┍╞┟┑╸╞┟┑┥╞╎┑┥╞┝┑┥╞╵┑┥╞╵┑┥╞╵┑╡┝┑╞╵┑╝╵╸╞╵╸ 285 6185 1 6190 DATA 1.395.0.0 5195 DATA -.6975,-1.208,0 6200 DATA 1.054E5,10,16 6225 + 6215 | Benzene C-C force constant. 6225 + 6230 DATA .6975,-1.208,0 6235 DATA -.6975.-1.208.0 6240 DATA 6.94E5,13,16 6245 ! 6255 ! Benzene C-C force constant. 6265 1 6270 DATA .6975,-1.208,0 6275 DATA -1.395,0,0 6280 DATA 1.05465,13,1 6285 ! **5290** 6295 / Benzene C-C force constant. 6305 ! 6310 DATA -. 6975, -1.208,0 6315 DATA -1.395.0.0 6320 DATA 6.94E5,16,1 6325 / 6335 | Benzene C-C force constant. 5340 \*\*!!\*\*!\*\*\*\*!!\*\*\*\* 6345 ! 6350 DATA -.6975,-1.208,0 6355 DATA -.6975.1.208.0 6360 DATA 1.054E5,16,4 6365 ! 6375 <sup>1</sup> Benzene C-H force constant. 6385 1 6390 DATA -2.479,0,0 6395 DATA -1.395.0.0 6400 DATA 5.508E5,19,1 6405 ! 6415 | Benzene C-H force constant. 6425 1 6430 DATA -2.479.0.0 6435 DATA -.6975.1.208.0 6440 DATA 1.093E5.19.4 6445 1 6455 <sup>1</sup> Benzere C-H force constant. 

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286 6465 1 6470 DATA -2.479,0,0 5475 CATA -.5375,-1.208.0 6480 DATA 1.09365,19,16 6485 1 6495 | Benzene C-H force constant. 6505 1 6510 DATA -1.24.2.147.0 6515 DATA -.6975.1.208.0 6520 DATA 5,508E5.22.4 6525 + 6535 | Benzene C-H force constant. 6545 1 6550 DATA -1.24,2.147,0 6555 DATA .6975,1.208,0 6560 DATA 1.093E5,22,7 6565 1 6575 | Benzene C-H force constant. 6585 I 6590 DATA -1.24,2.147,0 6595 DATA -1.395.0.0 6600 DATA 1.093E5.22.1 6605 ! 6615 ! Benzene C-H force constant. 6625 ! 6630 DATA 1.24,2.147,0 6635 DATA .6975.1.208.0 6640 DATA 5.508E5,25,7 6645 ! 6655 | Benzene C-H force constant. 6665 ! 6670 DATA 1.24,2.147,0 6675 DATA 1.395,0,0 6680 DATA 1.093E5,25,10 6685 ! 6695 / Benzene C-H force constant. 6705 1 6710 DATA 1.24,2.147,0 6715 DATA -.6975,1.208,0 6720 DATA 1.093E5,25,4 6725 + 6735 | Benzene C-H force constant. 6745 +

5750 CATA 2.479.0.0 5755 DATA 1,395.0.0 S760 DATA 5,50365,28,10 6765 / 5775 | Benzene C-H force constant. 6785 + 6790 DATA 2.479,0,0 6795 DATA .6975,1.208,0 6800 DATA 1.093E5.28.7 6805 4 6815 | Benzene C-H force constant. 6825 1 6830 DATA 2.479.0.0 6835 DATA .6975,-1.208,0 6840 DATA 1.093E5,28,13 6845 ! 6855 ! Benzene C-H force constant. 6865 1 6870 DATA 1.24 .- 2.147 .0 6875 DATA .6975.-1.208.0 6880 DATA 5.508E5.31.13 6885 1 **5890** 6895 | Benzene C-H force constant. 6905 1 6910 DATA 1.24 .- 2.147.0 6915 DATA 1.395.0.0 6920 DATA 1.093E5.31.10 8925 / 6935 | Benzene C-H force constant. 6945 I 6950 DATA 1.24 .- 2.147.0 6955 DATA -.6975.-1.208.0 6960 DATA 1.093E5,31,16 6965 I 6975 | Benzene C-H force constant. 6985 I 6990 DATA -1.24,-2.147,0 6995 DATA -. 6975 .- 1. 208 .0 7000 DATA 5.508E5.34.16 7005 1 7015 + Benzene C-H force constant. 7025 1 7030 DATA -1.24,-2.147,0

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7035 DATA .5975,-1.208,0 7040 DATA 1.093E5,34,13 1345 1055 - Benzene C-H force constant. 7065 1 7070 DATA -1.24,-2.147,0 7075 DATA -1.395,0.0 7080 DATA 1.093E5,34,1 7085 1 7095 | Benzene C-C out-of-plane force constant. 7105 1 7110 DATA -1.395.0.1 7115 DATA -.6975.1.208.0 7120 DATA 165,1,4 7125 + 7135 / Benzene C-C out-of-plane force constant. 7145 1 7150 DATA -.6975,1.208,1 7155 DATA .6975,1.208,0 7160 DATA 165.4.7 7165 ! 7175 | Benzene C-C out-of-plane force constant. 7185 1 7190 DATA .6975,1.208,1 7195 DATA 1.395,0,0 7200 DATA 165,7,10 7205 1 7215 | Benzene C-C out-of-plane force constant. 7225 1 7230 DATA 1.395.0.1 7235 DATA .6975.-1.208.0 7240 DATA 165,10,13 7245 1 7255 | Berzene C-C out-of-plane force constant. 7265 1 7270 DATA .6975.-1.208.1 7275 DATA -.6975,-1.208.0 7280 DATA 165,13,16 7295 1 7295 | Benzene C-C out-of-plane force constant. 7205 1 7310 DATA -.5975.-1.208.1 7315 DATA -1.395.0.0

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7505 1 7510 (0.000) (0 7515 ! Methane C-H force constant. 1575 1 7630 DATA 0.7.79346319178E-5.3.46941045213 7540 DATA 4.79E5,37,43 7545 1 7655 | Methane C-H force constant. 7665 L 7670 DATA 0.7.78346319178E-5.3.46941045213 7675 DATA 0.1.03987841578.3.1043966958 7680 DATA 4.79E5.37.46 7685 ! 7695 ! Methane C-H force constant. 7705 1 7710 DATA 0,7.78346319178E-5,3.46941045213 7715 DATA 6.77616018046E-21..000106109702385.4.56937173828 7720 DATA 4.79E5,37,49 7725 ! 7735 | Methane H-H force constant. 7745 1 7750 DATA -.8989, -.518821585724, 3.10438237398 7755 DATA .8989,-.518821585724,3.10438237398 7760 DATA 1.85E5.40.43 7765 1 7775 1 Methane H-H force constant. 7785 1 7790 DATA .8989.-.518821585724.3.10438237398 7795 DATA 0,1.03987841578,3.1043966958 7800 DATA 1.85E5,43,46 7805 1 7915 ! Methane H-H force constant. 7825 1 7830 DATA 0,1.03987841578,3.1043966958 -. 3989 .-. 518821585724 .3. 10438237398 7835 DATA 7840 DATA 1.85E5,46,40 7845 | 7955 <sup>1</sup> Methane H-H force constant. 7965 / 7870 DATA 0,1.03987841578,3.1043966958 7875 DATA 6.77616018046E-21,.000106109702385,4.56937172929 7880 DATA 1.85E5.46.49 7885 1

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7395 - Methane H-H force constant. 1985 1 7910 DATA 6.77515018046E-21,.000106109702385,4.56937173828 7920 DATA 1.8565,49,43 7925 1 7935 | Methane H-H force constant. 7945 ( 7950 DATA 6.77616018046E-21,.000106109702385,4.56937173828 7955 DATA -. 8989, -. 518821585724, 3.10438237398 7950 DATA 1.85E5,49,40 7965 ( 7975 / Data statements containing cluster subunit coordinates. 7980 ! The data is used to draw the cluster geometry and vdW mode displacement 7985 ! vectors. The coordinates are entered as they are numbered in ECCEMP2. 7990 -7995 1 8000 ! Order of data entries: 8005 1 8010 | DATA X-Coord, Y-Coord, Z-Coord, Atom #. 8015 1 8020 | Atom #: 8025 1 1 - Carbon. 8030 1 2 - Hydrogen. 3 - Nitrogen. 8035 1 4 - Oxygen. 8040 1 8045 1 5 - User defined atom. 8055 1 8060 DATA ~1.395,0,0,1 8065 DATA ~. 6975, 1.208,0,1 8070 DATA .6975,1.208,0,1 8075 DATA 1.395,0,0,1 8080 DATA .6975,-1.208,0,1 8085 DATA -.6975,-1.208,0,1 8090 DATA -2.479,0,0,2 8095 DATA -1.24,2.147,0.2 8100 DATA 1.24,2.147,0,2 9105 DATA 2.479.0.0,2 8110 DATA 1.24,-2.147,0,2 8115 DATA -1.24 -2.147,0,2 8120 DATA 0.0.3.4694.1 8125 DATA -. 8989, -. 5188, 3.10438, 2 9135 DATA 0,1.0398,3.10439,2 3140 DATA 0,.000106,4.56937,2 8145 END 8150 1 9155 4 8160 1 9165 1 8170 1

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9175 1 9190 1 2135 3190 1 3195 1 3200 + 3205 1 8215 / Subprogram: Eigen 8220 1 8225 | This subprogram determines all the eigenvalues and eigenvectors of a real 8230 | general matrix. The eigenvalues are computed by the QR double-step method 8235 ) and the eigenvectors by inverse interation. 8240 1 8245 1 8250 / This subprogram was taken from HP 98821A BASIC Numerical Analysis Library 8255 ! which is designed to be executed using HP series 200 computers. 8265 1 8270 1 8275 1 8280 1 8285 ! 8290 1 8295 1 8300 1 9860 SUB Eigen(N,A(+),Evr(+),Evi(+),Vecr(+),Veci(+),Indic(+)) 9870 Baddta=(N<=0) 9880 IF Baddta=0 THEN 9930 9890 PRINT FNLin\$(2); "ERROR IN SUBPROGRAM Eigen." 9900 PRINT "N=":N:FNLin\$(2) 9910 PAUSE 9920 GOTO 9870 9930 OPTION BASE 1 9940 ALLOCATE INTEGER Local(N) 9950 ALLOCATE Prfact(N),Subdia(N),Work(N) 9960 IF N<>1 THEN 10030 9970 Evr(1)=A(1,1) 9980 Ev1(1)=0 9990 Vecr(1,1)=1 10000 Veci(1,1)=0 10010 Indic(1)=2 10020 GOTO 11200 10030 CALL Scale(N,A(+),Veci(+),Prfact(+),Enorm) 10040 Ex=EXP(-39+L06(2)) 10050 CALL Hesqr(N,A(+),Veci(+),Evr(+),Evi(+),Subdia(+),Indic(+),Eps,Ex) 10060 J=N 10070 I=1 10080 Local(1)=1 10090 IF J=1 THEN 10160 10100 IF ABS(Subdia(J-1)) Eps THEN 10130 10110 [=I+1 10120 Local(I)=0 10130 J=J-1 10140 Local(I)=Local(I)+1 10150 IF J. 1 THEN 10100 10150 H=1

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10170 Kon=0 10180 L=Local(1) 10190 M=N 10200 FOR I=1 TO N 10210 Ivec=N-I+1 10220 IF I<=L THEN 10260 10230 K=K+1 10240 M=N-L 10250 L=L+Local(K) IF Indic(Ivec)=0 THEN 10420 10250 IF Evi(Ivec)<>0 THEN 10370 10270 10280 FOR KI=1 TO M 10290 FOR LI=K1 TO M 10300 A(K1,L1)=Veci(K1,L1) 10310 NEXT L1 10320 IF K1=1 THEN 10340 A(K1,K1-1)=Subdia(K1-1) 10330 NEXT K1 10340 10350 CALL Realve(N,M,Ivec,A(+),Vecr(+),Evr(+),Evi(+),Work(+),Indic(+),Eps,Ex ) 10360 60T0 10420 10370 IF Kon<>0 THEN 10410 10380 Kon=1 10390 CALL Compve(N,M,Ivec,A(+),Vecr(+),Veci(+),Evr(+),Evi(+),Indic(+),Subdia (+),Work(+),Eps,Ex) 10400 GOTO 10420 10410 Kon=0 10420 NEXT I 10430 FOR I=1 TO N 10440 FOR J=1 TO N 10450 A(I,J)=0 10460 NEXT J 10470 A(I,I)=1 10480 NEXT I 10490 IF N<=2 THEN 10640 10500 M=N-2 10510 FOR K=1 TO M 10520 L=K+1 10530 FOR J=2 TO N 10540 D1=0 FOR I=L TO N 10550 10560 D2=Veci(I\_K) D1=D1+D2+A(J,I)10570 10580 NEXT I FOR I=L TO N 10590 A(J,I)=A(J,I)-Veci(I,K)+D1 10500 10510 NEXT I NEXT J 10520 10630 NEXT K 10540 Kon=1 10650 FOR I=1 TO N 10660 L=0 IF Evi(I)=0 THEN 10720 10570 10680 L=1 IF Kon=0 THEN 10720 10690 10700 Kon=Ø 10710 GOTO 11190

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10720 FOR J=1 TO N D1=0 10730 10740 D2=0 FOR K=1 TO N 10750 10760 D3=A(J.K) D1=D1+D3+Vecr(K,I) 10770 IF L=0 THEN 10800 10780 10790 D2=D2+D3+Vecr(K,I-1) 10800 NEXT K Work(J)=D1/Prfact(J) 10810 IF L=0 THEN 10840 10820 Subdia(J)=D2/Prfact(J) 10830 10840 NEXT J IF L=1 THEN 10970 10850 D1=0 10860 FOR M=1 TO N 10870 D1=D1+Work(M)^2 10880 NEXT M 10890 D1=SQR(D1)10900 FOR M=1 TO N 10910 Vec1(M.I)=0 10920 Vecr(M,I)=Work(M)/D1 10930 10940 NEXT M 10950 Evr(I)=Evr(I)+Enorm 60T0 11190 10960 Kon=1 10970 Evr(I)=Evr(I)+Enorm 10980 10990 Evr(I-1)=Evr(I) Evi(I)=Evi(I)+Enorm 11000 11010 Evi(I-1)=-Evi(I) 11020 R=0 11030 FOR J=1 TO N R1=Work(J)^2+Subdia(J)^2 11040 IF R>=R1 THEN 11080 11050 11060 R=R1 11070 L=J NEXT J 11080 D3=Work(L) 11090 11100 R1=Subdia(L) FOR J=1 TO N 11110 D1=Work(J) 11120 11130 D2=Subdia(J) Vecr(J,I)=(D1+D3+D2+R1)/R 11140 Vec1(J,I)=(D2+D3-D1+R1)/R 11150 Vecr(J,I-1)=Vecr(J,I) 11160 Veci(J,I-t) = -Veci(J,I)11170 11180 NEXT J 11190 NEXT I 11200 SUBEXIT 11210 SUBENDI 11220 SUE Scale(N,A(+),H(+),Prfact(+),Enorm) 11230 OPTION BASE 1 11240 INTEGER I, J, Iter, Noount 11250 FOF I=1 TO N FOR J=1 TO N 11260 11270 H(I,J)=A(I,J)

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NEXT J
11280
         Prfact(I)=1
11290
11300 NEXT I
11310 Bound1*.75
11320 Bound2=1.33
11330 Iter=0
11340 Ncount=0
      FOR I=1 TO N
11350
         Column=0
11360
         Row=0
11370
         FOR J=1 TO N
11380
11390
           IF I=J THEN 11420
           Column=Column+ABS(A(J,I))
11400
           Row=Row+ABS(A(I,J))
11410
         NEXT J
11420
         IF Column=0 THEN 11480
11430
         IF Row=0 THEN 11480
11440
11450
         Q=Column/Row
         IF QKBound1 THEN 11500
11460
         IF Q>Bound2 THEN 11500
11470
11480
         Ncount=Ncount+1
11490
         60T0 11570
11500
         Factor=SQR(Q)
         FOR J=1 TO N
11510
           IF I=J THEN 11550
11520
           A(I,J)=A(I,J)+Factor
11530
11540
           A(J,I)=A(J,I)/Factor
         NEXT J
11550
         Prfact(I)=Prfact(I)+Factor
11560
       NEXT I
11570
       Iter=Iter+1
11580
       IF Iter>30 THEN 11750
11590
       IF Ncount (N THEN 11340
11600
11510 Fnorm=0
       FOR I=1 TO N
11620
         FOR J=1 TO N
11630
           Q=A(I,J)
11640
           Fnorm=Fnorm+Q+Q
11650
         NEXT J
11660
11570 NEXT I
      Fnorm=SQR(Fnorm)
11680
11590 FOR I=1 TO N
         FOR J=1 TO N
11700
           A(I,J)=A(I,J)/Fnorm
11710
         NEXT J
11720
11730 NEXT I
11740 Enorm=Fnorm
 11750 6010 11830
 11760 FOR I=1 TO N
         Prfact(I)=1
 11770
         FOR J=1 TO N
 11780
 11790
           A(I,J)=H(I,J)
11800
         NEXT J
11510 NEXT I
11820 Enormal
11830 SUEEKIT
11840 SUBEND+
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11850 SUB Hesgr(N,A(+),H(+),Evr(+),Evi(+),Subdia(+),Indic(+),Eps.Ex) 11860 OPTION BASE 1 11870 INTEGER I.J.K.L.M.Maxst.MI.Ns IF N-2<0 THEN 12460 11880 11890 IF N-2>0 THEN 11920 Subdia(1)=A(2,1) 11900 11910 60T0 12460 11920 M=N-2 FOR K=1 TO M 11930 L=K+1 11940 5=0 11950 FOR I=L TO N 11960 H(I,K)=A(I,K)11970 11980 S=S+ABS(A(I,K)) 11990 NEXT I IF 5<>ABS(A(K+1,K)) THEN 12040 12000 Subdia(K)=A(K+1,K) 12010 H(K+1.K)=0 12020 12030 60T0 12410 12040 Sr2=0 12050 FOR I=L TO N 12060 Sr=A(I,K) 12070 Sr=Sr/S A(I,K)=Sr 12080 Sr2=Sr2+Sr+Sr 12090 NEXT I 12100 Sr=SQR(Sr2) 12110 IF A(L,K)<0 THEN 12140 12120 Sr=-Sr 12130 Sr2=Sr2-Sr+A(L,K) 12140 A(L,K)=A(L,K)-Sr 12150 H(L,K)=H(L,K)-5r+S 12160 Subdia(K)=Sr+S 12170 12180 X=S+SQR(Sr2) 12190 FOR I=L TO N H(I,K)=H(I,K)/X 12200 Subdie(I)=A(I,K)/Sr2 12210 NEXT I 12220 12230 FOR J=L TO N 12240 Sr=0 FOR I=L TO N 12250 Sr=Sr+A(I,K)+A(I,J) 12260 NEXT I 12270 FOR I=L TO N 12280 A(I,J)=A(I,J)-Subdia(I)+Sr12290 NEXT I 12300 12310 NEXT J FOR J=1 TO N 12320 12230 5-=0 FOR I=L TO N 12340 12250 Sr=Sr+A(J,I)+A(I,K) NERT I 12352 FOR I=L TO N 12278 12380 A J,I = A(J,I)~Subdia(I)\*Sr 12390 NEXT 1 12400 NEXT J

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12420 FOR K=1 TO M
12430
        A(K+1,K)≠Subdia(K)
12440 NEXT K
12450 Subdia(N-1)=A(N,N-1)
12450 Eps=0
12470 FOR K=1 TO N
        Indic(K)≠0
12480
        IF K<>N THEN Eps=Eps+Subdia(K)^2
12490
        FOR I=K TO N
12500
          H(K,I)=A(K,I)
12510
          Eps=Eps+A(K,I)"2
12520
        NEXT I
12530
12540 NEXT K
12550 Eps=Ex+SQR(Eps)
12560 Shift=A(N_N-1)
12570 IF N<=2 THEN Shift=0
       IF A(N,N)<>0 THEN Shift=0
12580
       IF A(N-1,N)<>0 THEN Shift=0
12590
       IF A(N-1,N-1)<>0 THEN Shift=0
12500
12610 M=N
12620 Ns=0
12630 Maxst=N+10
12640 FOR I=2 TO N
         FOR K=I TO N
12650
           IF A(I-1,K)<>0 THEN 12750
12660
         NEXT K
12670
12680 NEXT I
12690 FOR I=1 TO N
        Indic(I)=1
12700
         Evr(I)=A(I,I)
12710
        Evi(I)=0
12720
12730 NEXT I
12740 60TO 13780
12750 K=M-1
12760 M1-K
12770 I-K
12780 IF K<0 THEN 13780
12790 IF K=0 THEN 13530
12800 IF ABS(A(M,K)) <= Eps THEN 13530
 12810 IF M-2=0 THEN 13580
 12820 I=I-1
 12830 IF ABS(A(K,I)) <= Eps THEN 12860
 12840 ¥=I
12850 IF KAL THEN 12820
 12860 IF K-M1 THEN 13580
 12870 S=A(M,M)+A(M),M1)+Shift
 12880 Sr=A(M,M)+A(M),M1)-A(M,M1)+A(M),M)+.25+Shift'2
 12890 A(++2,+)=0
12900 X=A(F,F)+(A(F,F)-S)+A(F,F+1)+A(F+1,K)+Sr
 128'8 - Y=4-++1, F )+(A(F, F)+A(F+1, F+1)-5)
12920 R#463+X /+485-Y
 SCEDE IF FEE THEN SHIFTER(M.M-1)
 12940 IF F=C THEN 12860
 12952 2=4-1+2,1+1 1+4-1+1,1+1
12960 Shift#0
12970 Ns=Ns+1
```

12980 FOR I=K TO M1 IF I=K THEN 13050 12990 X=A(I,I-1)13000 Y=A(I+1,I-1) 13010 13020 Z=0 IF I+2>M THEN 13050 13030 Z=A(I+2,I-1) 13040 Sr2=ABS(X)+ABS(Y)+ABS(Z) 13050 IF 5-2=0 THEN 13100 13060 13070 X=X/Sr2 13080 Y=Y/Sr2 Z=Z/Sr2 13090 S=SQR(X+X+Y+Y+Z+Z) 13100 IF X<0 THEN 13130 13110 13120 S=-S IF I=K THEN 13150 13130 A(I,I-1)=S\*Sr2 13140 IF 5r2<>0 THEN 13180 13150 IF I+3>M THEN 13500 13160 13170 60T0 13470 13180 Sr=1-X/S 13190 5=X-S 13200 X=Y/S 13210 Y=Z/S 13220 FOR J=I TO M 13230 S=A(I,J)+A(I+1,J)+XIF I+2>M THEN 13260 13240 S=S+A(I+2,J)+Y 13250 13260 5=5+5r A(I,J)=A(I,J)-S13270 A(I+1,J)=A(I+1,J)-S=X13280 IF 1+2>M THEN 13310 13290 A(I+2,J)=A(I+2,J)-S+Y 13300 NEXT J 13310 13320 L=I+2 IF IKMI THEN 13350 13330 13340 L=M FOR J=K TO L 13350 S=A(J,I)+A(J,I+1)+X 13360 13370 IF I+2>M THEN 13390 13380 S=S+A(J,I+2)+Y 13390 S=S+Sr A(J,I)=A(J,I)-S13400 A(J,I+1)=A(J,I+1)-S+X13410 IF I+2>M THEN 13440 13420 A(J,I+2)=A(J,I+2)-5+Y 13430 NEXT J 13440 IF 1+3.M THEN 13500 13450 S=-A(I+3,I+2)\*Y\*5r 13450 13470 A(1+3,1)=5 A(1+3,1+1)=5+X 13492 A'I+3,I+2)=5+Y+A(I+3,I+2) 13452 13500 NEXT I :3512 IF No Marst THEN 13780 GOTE 12750 13520 13530 EVE (M/=A(M,M) 12540 Evi(M)=0

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13550 Indic(M)=1
13550 M=K
13570
      GOTO 12750
13580 R=.5+(A(K,K)+A(M,M))
13590
      S=.5+(A(M,M)-A(K,K))
      S=5+5+A(K.M)+A(M.K)
13600
13610
      Indic(K)=1
13620
      Indic(M)=1
13630 IF S<0 THEN 13710
13640 T=5QR(S)
13650 Evr(K)=R-T
13660 Evr(M)=R+T
13670 Evi(K)=0
13680 Evi(M)=0
13690 M=M-2
13700 GOTO 12750
13710 T=SQR(-S)
13720 Evr(K)=R
13730 Evi(K)=T
13740 Evr(M)=R
13750 Evi(M)=-T
13760 M=M-2
13770 GOTO 12750
13780 SUBEXIT
13790 SUBEND!
13800 SUB Realve(N,M,Ivec,A(*),Vecr(*),Evr(*),Evi(*),Work(*),Indic(*),Eps,Ex)
13810 Baddta=(N<=0) OR (M<=0) OR (Ivec<=0)
      IF Baddta=0 THEN 13860
13820
13830 PRINT FNLin$(2); "ERROR IN SUBPROGRAM Realve."
13840 PRINT "N=";N,"M=";M,"Ivec=";Ivec;FNLin$(2)
13850 PAUSE
13860 OPTION BASE 1
13870 ALLOCATE INTEGER Iwork(N)
13880 INTEGER I, Iter, J, K, L, Ns
13890 Vecr(1,Ivec)=1
13900 IF M=1 THEN 14850
13910 Evalue=Evr(Ivec)
13920 IF Ivec=M THEN 14010
13930 K=Ivec+1
13940 R=0
13950 FOR I=K TO M
13960
        IF Evalue<>Evr(I) THEN 13990
13970
         IF Ev1(I)<>0 THEN 13990
13980
         R=R+3
13990 NEXT I
14000 Evalue=Evalue+R+Ex
14010 FOR K=1 TO M
14020
       A(E,E)=A(E,E)-Evalue
14030 NEXT H
14040 H=M-1
14050 FOF I=1 TO K
14080
        i_ = ] + 1
14070
         lwor+(l/=€
14080
        IF A(1+1,1) 0 THEN 14120
14092
        IF A'I, I: -0 THEN 14240
14100
         A(I,I)=Eps
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14110 GOTO 14240 IF ABS(A(I,I))>=ABS(A(I+1,I)) THEN 14190 14120 Iwork(I)=1 14130 FOR J=I TO M 14140 R=A(I,J) 14150 A(I,J)=A(I+1,J)14160 14170 A(I+1,J)=R NEXT J 14180 14190 R=-A(I+1,I)/A(I,I)14200 A(I+1,I)=P FOR J=L TO M 14210 A(I+1,J)=A(I+1,J)+R+A(I,J)14220 14230 NEXT J 14240 NEXT I 14250 IF A(M,M) >0 THEN 14270 14260 A(M,M)=Eps 14270 FOR I=1 TO N IF 1>M THEN 14310 14280 14290 Work(I)=1 60T0 14320 14300 Work(I)=0 14310 14320 NEXT I 14330 Bound=.01/(Ex+N) 14340 Ns=0 14350 Iter=1 14360 R=0 14370 FOR I=1 TO M 14380 J=M-I+1 14390 S=Work(J) IF J=M THEN 14460 14400 L=J+1 14410 14420 FOR K=L TO M 14430 Sr=Work(K) 14440 S=5-5r+A(J,K) 14450 NEXT K Work(J)=S/A(J,J)14460 14470 T=ABS(Work(J)) IF R>=T THEN 14500 14480 14490 R=T 14500 NEXT I 14510 FOR I=1 TO M Work(I)=Work(I)/R 14520 14530 NEXT I 14540 R1=0 14550 FOR I=1 TO M 14560 T=0 FOR J=I TO M 14570 14580 T=T+A(I,J)+Wort(J) NEXT J 14590

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14642 IF Iter=1 THEN 14660 14650 IF Frevis =F1 THEN 14650

F : = \*

14650 FOR I=1 TO M

IF FI #T THEN 14ED0

Vech(I,Ivec)=Work(I)

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14522

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14600 NEYT 1

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14680 NEXT I
14690 Previs=R1
14700 IF Ns=1 THEN 14850
14710 IF Iter/6 THEN 14860
14720 Iter=Iter+1
14730 IF R'Bound THEN 14750
14740 Ns=1
14750 H=M-1
14760 FOR I=1 TO K
        R=Work(I+1)
14770
        IF Iwork(I)=0 THEN 14820
14780
        Work(I+1) = Work(I) + Work(I+1) + A(I+1,I)
14790
        Wort(I)=R
14800
14810
        GOTO 14830
        Work(I+1)=Work(I)+A(I+1,I)+Work(I+1)
14820
14830 NEXT I
14840 60T0 14360
14850 Indic(Ivec)=2
14860 IF M=N THEN 14910
14870 J=M+1
14880 FOR I=J TO N
14890
       Vecr(I,Ivec)=0
14900 NEXT I
14910 SUBEXIT
14920 SUBEND!
14930 SUB Compve(N,M,Ivec,A(+),Vecr(+),H(+),Evr(+),Evi(+),Indic(+),Subdia(+),Wor
k(+).Eps.Ex)
14940 Baddta=(N<=0) OR (M<=0) OR (Ivec<=0)
14950 IF Baddta=0 THEN 14990
14950 PRINT FNLin$(2); "ERROR IN SUBPROGRAM Compve."
14970 PRINT "N=";N."M=";M."Ivec=";Ivec;FNLin$(2)
14980 PAUSE
14990 OPTION BASE 1
15000 ALLOCATE INTEGER Iwork(N)
15010 ALLOCATE Work1(N), Work2(N)
15020 INTEGER I.II.IZ, Iter, J.K. L. Ns
15030 Fksi=Evr(Ivec)
15040 Eta=Evi(Ivec)
15050 IF Ivec=M THEN 15160
15060 K=Ivec+1
15070 R=0
15080 FOR I=K TO M
       IF FFS1 ()EVr(I) THEN 15120
15090
        IF ABS(Eta)<>ABS(Evi(I)) THEN 15120
15100
15110
         R=R+3
15120 NEXT I
15130 P=R+E-
15140 E+51=F+51+R
15:50 Eta=Eta+R
15160 R=F+s1+F+s1+Eta+Eta
15178 Sel+Frei
15182 L=M-1
15150 FCF I=1 TC M
        - FÇ≂ (±1 t0 M
15202
15218
         5=3
15220
           A'J.I)=0
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FOR K=I TO J 15230 D=D+H(I,K)+H(K,J) 15240 15250 NEXT K A(I,J)=D-S+H(I,J)15260 NEXT J 15270 15280 A(I,I)=A(I,I)+R15290 NEXT I 15300 FOR I=1 TO L 15310 R=Subdia(I) A(I+1,I)=-S\*R15320 I1=I+115330 15340 FOR J=1 TO I1 A(J,I)=A(J,I)+R+H(J,I+1)15350 NEXT J 15360 15370 IF I=1 THEN 15390 A(I+1,I-1)=R+Supdia(I-1) 15380 FOR J=I TO M 15390 A(I+1,J)=A(I+1,J)+R\*H(I,J)15400 15410 NEXT J NEXT I 15420 15430 ¥=M-1 FOR I=1 TO K 15440 15450  $I_{1=I+1}$ 15460 12=1+2 Iwork(I)=0 15470 IF I=K THEN 15500 15480 IF A(I+2,I)<>0 THEN 15540 15490 IF A(I+1,I)<>0 THEN 15540 15500 IF A(I,I)<>0 THEN 15770 15510 15520 A(I,I)=Eps 15530 60T0 15770 IF I=K THEN 15600 15540 IF ABS(A(I+1,I))>=ABS(A(I+2,I)) THEN 15600 15550 IF ABS(A(I,I))>=ABS(A(I+2,I)) THEN 15700 15560 L=I+2 15570 15580 Iwork(I)=2 60T0 15630 15590 IF ABS(A(I,I))>=ABS(A(I+1,I)) THEN 15680 15600 15610 L=I+115620 Iwor+(I)=1 15630 FOR J=I TO M 15640 R=A(I,J) A(I,J)=A(L,J)15650 15660 A(L,J)=R15670 NEXT J IF ICK THEN 15700 15580 12=11 15690 FOR L=I1 TO I2 15700 15710 R=-A(L,I)/A I,I) 15720 A-L,I-F 15732 FOR JEIN TO M A'E,J'=A'E,J'+R+A'I,J' 15742 NEXT 15752 15752 NERT -15770 NE+T : 15782 IF AMM, MU & THEN 15800 ALM MIEECS 15752

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15822 FDF 1+1 TO N
15510 IF INM THEN 15850
15520 Veer(1,1vec1+)
15820 Veer(1,1vec1+)
15830 Veer(1,1vec1+0
15850 Veer(1,1vec1+0
15850 Ne+0
15850 Ne+0
15850 Ne+0
15850 Veer(1,1vec1+1)
15850 Veer(1,1vec1+1)
15850 Iter+)
15850 D=Veer(1,1vec1,1vec(1,1vec)
15850 IF 1+1 THEN 15980
15970 DD+Sudula(1-)+veer(1,1vec)
15850 IF L=1 THEN 15980
15970 DD+Sudula(1-)+veer(K,1ve
15800 FOR K=L TO M
15810 D=D+HI,K)+veer(K,1ve
15820 Veer(1,1vec-1)=D=Eteve
15820 Veer(1,1vec-1)=D=Eteve
15820 Veer(1,1vec-1)=Veer(1,1vec))
15850 R=Veer(L,1vec-1)=Veer(1,1vec))
15850 Veer(1,1vec-1)=Veer(1,1vec))
15850 Veer(1,1vec-1)=Veer(1,1vec))
15850 Veer(1,1vec-1)=Veer(1,1vec))
15850 FOR I=1 TO K
15870 Veer(1,1vec-1)=Veer(1,1vec))
15850 FOR I=1 TO K
15870 Veer(1,1vec-1)=Veer(1,1vec))
15860 FOR K=L TO M
15810 Veer(1,1vec-1)=Veer(1,1vec))
15860 FOR K=L TO M
15810 Ueer(1,1vec-1)=Veer(1,1vec))
15860 FOR I=1 TO M
15810 IF J=H THEN 15140
15820 IF J=H THEN 15240
15820 NEXT I
15250 REXT I
15250 FOR I=1 TO M
15210 D1=A(J,K)
15220 D=D+Veer(J,1vec-1)
15220 IF J=1 THEN 1528
15220 IE J=1 THEN 1528
15220 Veer(1,1vec-1)=Veer(1,1vec))
15220 REXT I
15250 REXT D=D+Subdia(I-1)+Vecr(I-1,Ivec) D=D+H(I,K)+Vecr(K,Ivec) Vecr(I,Ivec-1)=D-Eta+Vecr(I,Ivec-1) Vecr(L,Ivec-1)=Vecr(I,Ivec-1) Vecr(I+1,Ivec-1)=Vecr(I+1,Ivec-1)+A(I+1,I)+R Vecr(I+2,Ivec-1)=Vecr(I+2,Ivec-1)+A(I+2,I)+R D=D-D1+Vecr(K,Ivec-1) Vecr(J,Ivec-1)=D/A(J,J) D=Work(I)\*Vecr(I,Ivec-1) D=D+Subdia(I=E)+vecr(I=!,Ivec=1) C=C+H+I,F3+Jecr(F,Ivec-F) Vech I, Ived >= (Vech(I, Ived)-D)/Eta

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 16370 L=1 16380 5=0 16390 FOR I=1 TO M R=Vecr(I,Ivec)^2+Vecr(I,Ivec-1)~2 IF R =S THEN 16440 S≈R L=I 15440 NEXT I 16450 U=Vecr(L,Ivec-1) V=Vecr(L,Ivec) FOR I=1 TO M B=Vecr(I,Ivec) R=Vecr(I,Ivec-1) Vecr(I,Ivec)=(R+U+B+V)/S Vecr(I,Ivec-1)=(B+U-R+V)/S 16520 NEXT I B=0 FOR I=1 TO M R=Work(I)+Vecr(I,Ivec-1)-Eta+Vecr(I,Ivec) U=Work(I)+Vecr(I,Ivec)+Eta+Vecr(I,Ivec-1) IF I=1 THEN 16600 R=R+Subdia(I-1)=Vecr(I-1,Ivec-1) U=U+Subdia(I-1)\*Vecr(I-1,Ivec) L=I+1 IF L>M THEN 16660 FOR J=L TO M R=R+H(I,J)+Vecr(J,Ivec-)) U=U+H(I,J)+Vecr(J,Ivec) NEXT J U=R+R+U+U IF B>=U THEN 16690 8=U 16690 NEXT I IF Iter=1 THEN 16720 IF Previs<=8 THEN 16830 FOR I=1 TO N Work1(I)=Vecr(I.Ivec) Work2(I)=Vecr(I,Ivec-1) 16750 NEXT I Previs=8 IF Ns=1 THEN 15870 IF Iter>6 THEN 15890 Iter=Iter+1 IF Bound>SQR(S) THEN 15940 Ns=1 GOTO 15940 FOR I=1 TO N Vecr(I,Ivec)=Work1(I) Vecr(I,Ivec-1)=WorkZ(I) 16850 NEXT I 16870 Indic(Ivec-1)#2 16880 Indic(Ivec)=2 16890 SUBEND! 15300 DEF FNLin\$(X1) 16910 X=INT(X1+.5) 16920 IF X=0 THEN RETURN CHR\$(13)

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16930 Eol\$=CHR\$(13)&CHR\$(10) 16940 IF X-0 THEN Eol\$=CHR\$(10) 16950 ALLOCATE R\$[X+LEN(Eol\$/] 16950 R\$="" 16970 FOR I=1 TO X 15980 R\$=R\$&Eol\$ 16990 NEXT I 17000 RETURN R\$ 17010 FNEND:

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APPENDIX FIVE

"H2PCNCA"

20 / Program name: H2PCNCA 30 1 40 | This program performs an out-of-plane normal coordinate analysis (NCA) on 50 I free base phthalocyanine. The NCA is conducted using the FG matrix method. 50 | The out-of-plane motion is modeled using the internal coordinates described 70 1 in chapter 7 of this dissertation. This particular program calculates the 80 | Au vibrations. The Blu, B2g, and B3g vibrations can be calculated by 90 ) replacing the data statements with those appropriate for the specific 100 I vibration symmetry. The data statements for the Blu, 82g, and 83g 110 Uvibrations are included at the end of this program. 130 1 140 OPTION BASE 1 150 PRINTER IS 1 160 1 180 ! Matrices used: 190 1 200 ! A(+)=Energy (FG) matrix. 210 ! Evr(+)=Real eigenvalue matrix. 220 ! Evi(+)=Imaginary eigenvalue matrix. 230 ! Vecr(+)=Real eigenvector matrix. 240 ! Veci(+)=Imaginary eigenvector matrix. 250 ! Indic(+)=Matrix diagonalization indicator matrix. 260 ! F(+)=Potential energy (F) matrix. 270 ! G(+)=Kinetic energy (G) matrix. 280 ! Vib(+)=Vibrational frequency matrix 290 + Fpert(+)=Perturbation matrix for vibrational mode fitting. 310 1 320 DIM A(30,30),Evr(30),Evi(30),Vecr(30,30),Veci(30,30),Indic(30),F(30,30),G(30 ,30),V15(30) 330 DIM Fpert(30,2) 340 OUTPUT 2:CHR\$(255)&"K"; 350 Amp=1 ! Amplification factor for vibrational fitting routine. 360 Dev=1 / Default printer set to the screen display. 370 N=23 1 Number of symmetry coordinates used in the NCA. 380 1 400 | Selects those coordinates not needed in a particular symmetry in 410 ! the F and 6 matrices. 430 1 440 Y1=19 450 YZ=0 460 Y3=0 470 Y4=0 480 Y5=0 490 Y5=0 500 Y7=0 510 Y8=0

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520 Y9=0 530 Y10=0 540 1 560 ! Reads in force field into F matrix from data statements. 570 1 580 | R=Row. 590 | C=Column. 600 | Sd=Flag - 0 to add, 1 to subract. E10 | Fell=Force constant 1. 620 | Fel2=Force constant 2. 640 1 650 READ R 660 IF R=9999 THEN 870 670 READ C,Sd,Fell,Fel2 580 IF 5d=0 THEN Fel=Fel1+Fel2 690 IF Sd=1 THEN Fe1=Fe11-Fe12 700 1 720 ! Deletes rows and columns in F matrix which are not needed. 740 1 750 IF Y1=R OR Y1=C THEN Fe1=0 760 IF Y2=R OR Y2=C THEN Fel=0 770 IF Y3=R OR Y3=C THEN Fel=0 780 IF Y4=R OR Y4=C THEN Fe1=0 790 IF Y5=R OR Y5=C THEN Fel=0 800 IF Y5=R OR Y6=C THEN Fel=0 810 IF Y7=R OR Y7=C THEN Fe1=0 820 IF Y8=R OR Y8=C THEN Fe1=0 830 IF Y9=R OR Y9=C THEN Fe1=0 840 IF Y10=R OR Y10=C THEN Fe1=0 850 F(R\_C)=Fel 860 6010 650 870 READ Fins 880 PRINT Fin\$ 890 PRINT 900 H 920 <sup>1</sup> Reads in 6 matrix elements from data statements. 930 1 940 + R=Row. 950 ! C=Column. 960 | Sd=Flag - 0 to add, 1 to subtract. 970 · Gell=G matrix element 1. 980 / Gel2=G matrix element 2. 1000 1010 READ P 1020 JF R=9999 THEN 1230 1030 FEAD C,Sd,Gel1,Gel2 1040 IF Scat THEN Gel=Gel1+Gel2 1052 IF Scal THEN Gel=Gel1-Gel2 1050 1082 " Deletes row and columns in G matrix which are not meeded.

1100 1 1110 IF Y1=R OR Y1=C THEN Gel=0 1120 IF Y2=R OR Y2=C THEN Gel=0 1130 IF Y3=R OR Y3=C THEN Gel=0 1140 IF Y4=R OR Y4=C THEN Ge1=0 1150 IF YS=R CR YS=C THEN Gel=0 1160 IF YE=R OR YE=C THEN Gel=0 1170 IF Y7=R OR Y7=C THEN Gel=0 1180 IF Y8=R OR Y8=C THEN Ge1=0 1190 IF Y9=R OR Y9=C THEN Gel=0 1200 IF Y10=R OR Y10=C THEN Ge1=0 1210 G(R\_C)=Gel 1220 GOTO 1010 1230 READ Fin\$ 1240 PRINT Fin\$ 1250 PRINT 1260 1 1280 | Generates symmetric F and G matrices. 1300 1 1310 FOR R=1 TO N 1320 FOR C=R TO N 1330 F(C,R)=F(R,C) 1340 6(C,R)=6(R,C) 1350 NEXT C 1360 NEXT R 1370 1 1390 | Removes zero rows and columns from F and 6 matrices and reduces their 1400 ) dimensions. 1420 1 1430 FOR C=1 TO N 1440 Sum=0 1450 FOR R=1 TO N 1460 Sum=6(R,C)+Sum 1470 NEXT R 1480 IF Sum<>0 THEN 1690 1490 FOR Z=C TO N 1500 FOR R=1 TO N 1510 IF Z=N THEN F(R,Z)=0 1520 IF Z=N THEN 6(R,Z)=0 1530 IF Z=N THEN 1560 1540 F(R,Z)=F(R,Z+1) 1550 G(R,Z)=G(R,Z+1) 1560 NEXT R 1570 NEXT Z 1580 FOF Z=C TO N 1590 FOR F=1 TO N 1600 IF 2=N THEN F(2,P)=0 1610 JF J=N THEN & J.R =0 1620 IF 2=N THEN 1650 1630 F 2 P = F 2+1 F 1640 6 2 F += 6 (2+1 F 1650 NEFT F

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310 1660 NEXT 2 1670 N=N-1 1650 6070 1430 1690 NEXT C 1700 -1720 - Multiplies the F and G matrices to yield the FG matrix for 1730 - diagonalization. 1750 + 1760 FOR R=1 TO N 1770 FOR C=1 TO N 1780 Sum=0 1790 FOR E1=1 TO N 1800 Sum=6(R,E1)+F(E1,C)+Sum 1810 NEXT E1 1820 A(R\_C)=Sum 1830 NEXT C 1840 NEXT R 1850 1 1870 - Call subprogram Eigen to diagonalize the F6 matrix. 1890 1 1900 CALL Eigen(N,A(+),Evr(+),Evi(+),Vecr(+),Veci(+),Indic(+)) 1910 BEEP 1920 -1940 / Shell sorts the eigenvalues and eigenvectors. 1950 | Sorted in increasing order with respect to eigenvalue. 1970 1 1980 Pass=N 1990 Pass=INT(Pass/2) 2000 IF Pass=0 THEN 2220 2010 FOR St=1 TO Pass 2020 I1=St 2030 Jj=St+Pass 2040 Sw=0 2050 IF Evr(I1) (= Evr(J3) THEN 2150 2060 Sw=1 2070 Avr=Evr(11) 2080 Evr(I1)=Evr(J3) 2090 Evr(J) = Avr 2100 FOR R=1 TO N 2110 Avc=Vecr(R,I1) 2120 Vecr(R,Ii)=Vecr(R,Jj) 2130 Vecr(R,J))=Avc 2140 NEXT R 2150 Ii=Jj 2160 Jj=Jj+Pass 2170 IF JJ N+1 THEN 2050 2180 IF Sw=0 THEN 2200 2190 GOTO 2020 2200 NEXT St 2210 GOTO 1990 2220 IF Pertflg=1 THEN 4490

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2230 -contracte NCA results on the scheen on on a paper printer. 2250 2250 2278 -1190 PRINTER IS Dev 2290 PRINT 2200 PRINT 2310 FRINT "Au Vibrational Modes for Free Base Phthalocyanine" 2320 PRINT 2330 PRINT 2340 1 2360 - Prints the F matrix. 2380 / 2390 PRINT "----- F Matrix -----------2400 PRINT 2410 PRINT 2420 L=INT(N/8)+1 2430 E=8 2440 Z=1 2450 FOR B=1 TO L 2460 IF B=L THEN E=N 2470 FOR C=2 TO E 2480 PRINT USING 2490:C 2490 IMAGE XX,DDD,XXXXX,# 2500 NEXT C 2510 PRINT 2520 FOR R=1 TO N 2530 J=Z 2540 FOR C=Z TO E 2550 IF J>=E THEN PRINT USING 2560:F(R.C) 2560 IMAGE 00.00000 2570 IF J>=E THEN 2600 2580 PRINT USING 2590 (F(R.C) 2590 IMAGE DD.DDDDD.XX.\* 2600 J=J+1 2610 IF JOE THEN J=Z 2620 NEXT C 2630 NEXT R 2640 PRINT 2650 PRINT 2660 Z=Z+8 2670 E=E+8 2580 NEXT 8 2630 PRINT 2700 PRINT 1730 ' Prints the G matrix. 2752 -2"60 PPINT ----- 6 Matrix -----2770 PRINT

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2780 PRINT 2790 L=INT(N/8)+1 2800 E=8 2310 2=1 2820 FOR B=1 TO L 2830 IF B=L THEN E=N 2840 FOR C=Z TO E 2850 PRINT USING 2860:C 2860 IMAGE XX,DDD,XXXXX,# 2870 NEXT C 2880 PRINT 2890 FOR R=1 TO N 2900 J=Z 2910 FOR C=Z TO E 2920 IF J>=E THEN PRINT USING 2930;G(R,C) 2930 IMAGE DD.DDDDD 2940 IF J>=E THEN 2970 2950 PRINT USING 2960:6(R,C) 2960 IMAGE 00.00000,XX,# 2970 J=J+1 2980 IF J>E THEN J=Z 2990 NEXT C 3000 NEXT R 3010 PRINT 3020 PRINT 3030 2=2+8 3040 E=E+8 3050 NEXT 8 3060 PRINT 3070 PRINT 3080 1 3100 | Prints the 6 matrix row/column sums. 3120 1 ----- G Matrix Row Sums ------3130 PRINT "-------3140 PRINT 3150 PRINT 3160 FOR C=1 TO N 3170 Sum=0 3180 FOR R=1 TO N 3190 Sum=Sum+6(R,C) 3290 NEXT R 3210 PRINT USING 32201C, Sum 3220 IMAGE 0000,".",X,SD.110E 3230 NEXT C 3240 PRINT 3250 PRINT 3260 1 3280 + Prints the vibrational frequencies. 3300 1 3310 PRINT "------- Normal Coordinate Analysis Results -----------3320 PRINT

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3330 PRINT 3340 PRINT "Vibrational Eigenvalues: Numbers 1 70":N;"in cm-}" 3350 PRINT 3360 FOR R=1 TO N 3370 IF Evr(R)>=0 THEN Vib(R)=(Evr(R)/1000+1.E+8+6.02E+23)^.5/2/PI/3.E+10 3360 IF Evr(R)<0 THEN UIb(R)=-(ABS(Evr(R))/1000+1.E+8+6.02E+23)^.5/2/PI/3.E+10 3390 PRINT USING 3400; R, VIb(R) 3400 IMAGE DODD, ". ", 1X, DDDDD.DDD 3410 NEXT R 3420 IF Dev=710 THEN 3450 3430 DISP "Press Continue To Proceed" 3440 PAUSE 3450 DISP 3460 FOR C=1 TO N 3470 PRINT 3480 PRINT 3490 J=1 3500 FOR R=1 TO N 3510 IF R>1 THEN 3590 3520 1 3540 Prints the eigenvector normal modes. 3550 1 3570 PRINT "Vibrational Eigenvector for Eigenvalue";C;":";Vib(C);"cm-1" 3580 PRINT 3590 IF Vecr(R,C)>=0 THEN Mot=100+Vecr(R,C)^2 3600 IF Vecr(R,C)<0 THEN Mot=-100+Vecr(R,C)^2 3510 IF J>=8 THEN PRINT USING 3520; Mot 3620 IMAGE DDDD.DDD 3630 IF J>=8 THEN 3660 3640 PRINT USING 3650:Mot 3650 IMAGE DDDD.DDD.2X.# 3550 J=J+1 3570 IF J>8 THEN J=1 3680 NEXT R 3690 PRINT 3700 IF Dev=710 THEN 3730 3710 DISP "Press Continue To Proceed" 3720 PAUSE 3730 NEXT C 3740 OUTPUT 2:CHR\$(255)& K": 3750 05="0" 3760 INPUT "Do you want to fit vibrations",Q\$ 3770 IF QS="Y" THEN 3970 3780 IF Q\$ "N" AND Q\$ > "Y" THEN 3750 3790 Q\$="Q" 3800 INPUT "Do you want a paper print",Q\$ 3810 1F 05='Y" THEN Dev=710 2820 IF G\$="N" THEN Dev=1 BETC IF OS "Y" AND OS - "N" THEN 3790 2840 GOTO 2280 2950 2570 - Performs a vibrational frequency fit to observed vibrations. Fit is 3880  $\pm$  performed using perturbation theory. The approach assumes that the  ${\mathbb P}$ 3892 ' matrix is correct and refines the F matrix using the eigenvectors until

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3900 ! a eigenvalue fit is obtained. 1970 1 3940 ( Prints the eigenvalues to be selected in the fitting routine. 3960 1 3970 PRINT "Vibrational Eigenvalues: Numbers 1 to";N;"in cm-1" 3980 PRINT 3990 FOR R=1 TO N 4000 PRINT USING 4010; R, V15(R) 4010 IMAGE DDDD,".",1X,DDDDD.DDD 4020 NEXT R 4030 PRINT 4040 R=0 4050 1 4070 <sup>+</sup> Selects the eigenvalues to be used in fit and inputs the values which 4080 ! are to be fit to. 4100 4110 INPUT "Eigenvalue Number(s) to be Fit: [CONT] to End".R 4120 IF R=0 THEN 4210 4130 IF R>N THEN 4040 4140 Fpert(R,2)=1 4150 PRINT "Eigenvalue":R:" is to be fit" 4160 INPUT "Enter the NEW Eigenvalue", Fpert(R,1) 4170 PRINT "New:":Fpert(R,));" 01d:";Vib(R) 4180 IF Fpert(R,1)>=0 THEN Fpert(R,1)=(Fpert(R,1)+2+PI+3.E+10)^2/6.02E+23/1.E+8+ 1000 4190 IF Fpert(R,1)<0 THEN Fpert(R,1)=-(ABS(Fpert(R,1))+2+PI+3.E+10)^2/6.02E+23/1 .E+8+1000 4200 GOTO 4040 4210 PRINT 4220 FOR P=1 TO N 4230 IF Fpert(P,2)=0 THEN 4350 4240 4260 ! Refines force constants using eigenvectors. 4280 1 4290 FOR R=1 TO N 4300 FOR C=1 TO N 4310 IF R<>C THEN 4330 4320 F(R,C)=F(R,C)+(Fpert(P,1)-Evr(P))+Vecr(R,P)+Vecr(C,P)+Amp 4330 NEXT C 4340 NEXT R 4350 NEXT P 4360 Pertfla=1 4370 1 4390 <sup>1</sup> Returns execution to part of program where F and G matrices are 4400 | multiplied and the FG matrix is diagonalized. 4420 1 4430 GOTO 1750 4440 1

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4450 + Companes calculated and sought eigenvalues, and if less than .01 4470 ! wavenumbers difference exists, the routine is terminated. 4430 FOR R=1 TO N 4500 IF Fpert(R,2)=0 THEN 4560 4510 IF Fpert(R,1)>≈0 THEN Pert#(Fpert(R,1)/1000+1.E+8+5.02E+23)\*.5/2 FI/3.E+\*0 4520 IF Fpert(R,1)<0 THEN Pert=-(ABS(Fpert(R,1))/1000+1.E+8+6.02E+23+.5-2.PI/3. E+10 4530 IF Evr(R)>=0 THEN Chk=(Evr(R)/1000+1.E+8+6.02E+23)^.5/2/PI/3.E+10 4540 IF Evr(R)<0 THEN Chk=-(ABS(Evr(R))/1000+1.E+8+6.02E+23)\*.5/2/PI/3.E+10 4550 PRINT "Sought:";Pert;" Present:":Chk 4560 NEXT R 4570 PRINT 4580 FOR R=1 TO N 4590 IF Fpert(R,2)=0 THEN 4650 4500 IF Fpert(R.1)>=0 THEN Pert=(Fpert(R.1)/1000+1.E+8+6.02E+23)^.S/2/PI/3.E+10 4610 IF Fpert(R.1)<0 THEN Pert=-(ABS(Fpert(R.1))/1000+1.E+8+6.02E+23)^.5/2/PI/3. E+10 4620 IF Evr(R)>=0 THEN Chk=(Evr(R)/1000+1.E+8+6.02E+23)^.5/2/PI/3.E+10 4630 IF Evr(R)<0 THEN Chk=-(ABS(Evr(R))/1000+1.E+8+6.02E+23)^.5/2/PI/3.E+10 4640 IF ABS(Pert-Chk)>.01 THEN 4210 4650 NEXT R 4660 Pertflo=0 4570 GOTO 2280 4680 4690 4700 I F matrix elements. 4710 4720 1 4730 DATA 1,1,0,.119,0 | Row/Column A. DATA 2,2,0,.119,0 1 Row/Column 8. 4740 DATA 3,3,0,.119,0 ' Row/Column C. 4750 DATA 4,4,0,.119,0 | Row/Column D. 4750 4770 DATA 5,5,0,.119,0 ! Row/Column E. DATA 5,5,0,.119,0 | Row/Column F. 4780 4790 DATA 7,7,0,.119,0 1 Row/Column 6. 4800 DATA 8,8,0,.119,0 ! Row/Column H. 4810 DATA 9,9,0,.119,0 ! Row/Column I. 4820 DATA 10,10,0,.119,0 + Row/Column J. 4830 DATA 11,11,0,.119,0 ! Row/Column K. 4840 DATA 12,12,0,.119,0 ! Row/Column L. 4850 DATA 13,13,0,.119,0 | Row/Column M. 4860 DATA 14,14,0,.119,0 ' Row/Column N. 4870 DATA 15,15,0,.3237,0 + Row/Column 0. DATA 15,16,0,.3237,0 1 Row/Column P. 4880 DATA 17,17,0,.3237,0 + Row/Column Q. 4890 4900 DATA 18,18,0,.3237,0 1 Row/Column R. DATA 19,19,0,.3237,0 1 Row/Column S. 4910 4920 DATA 20,20,0,.3237,0 / Row/Column T. 4930 DATA 21,21,0,.3237,0 + Row/Column U. DATA 22,22,0,.3237,0 + Row/Column V. 4940 4950 DATA 23,23,0,.3237,0 ' Row/Column W. 4960 DATA 9999 4970 DATA F Matrix Loaded 4980 4990 

- G matrix elements. 5000 50:0 5220 5232 5243 CATA 1,1,0,.565893424036,0 5050 CATA 1,2,0,-.320683347477,-.320683347477 5260 CATA 1,3,0,.160341673739,.160341673739 5070 CATA 1,4,0,-.113378684807,0 DATA 1,15,1,.333006762504,-.333006762504 5280 5090 DATA 1,16,1,-.183960876595,.183860876595 DATA 1,20,1,.0347149906857,-.0347149906857 5100 SECONDENSE Row/Column B PERFECTATION STATEMENT STATEMENTS STATEMENTS 5110 DATA 2,2.0,.566893424036,.226757369615 5120 5130 DATA 2,3,0,-.453514739229,-.113378684807 5140 DATA 2,4,0,.320583347478,0 5150 DATA 2,5,0,.0805797038531,.0510728876983 5160 DATA 2,5,0,-.0494507276469,0 DATA 2,15,1,-.470942679895,.26001854527 5170 DATA 2,15,1,.470942679895,-.0490944106454 5180 DATA 2,20,1,-.226146200577,.0367387608396 5190 DATA 2,21,1,.0389677051594,0 5200 5210 DATA 3,3,0,.566893424036,.226757369615 5220 DATA 3,4,0,-.641366694954,0 5230 DATA 3,5,0,-.212232295405,-.18272547925 5240 5250 DATA 3,5,0,.0989014552938,.0494507276469 DATA 3,15,1,.26001854527,-.0490944106454 5260 DATA 3,16,1,-.470942679895.0490944106454 5270 DATA 3,20,1,.390842340703.-.201434900966 5280 5290 DATA 3.21,1,-.0779354103188..0389677051594 5300 5310 DATA 4,4,0,.566893424036,0 DATA 4,5,0..185184880494,.185184880494 5320 5330 DATA 4,6,0,-.0699338897075,-.0699338897075 5340 DATA 4,15,1,-.0347149906857,.0347149906857 5350 DATA 4,15,1,.183860876595,-.183860876595 5360 DATA 4,20,1,-.267630505722,.267630505722 5370 DATA 4,21,1,.055108657131,-.055108657131 5380 5390 DATA 5.5.0..32337728364..10262392793 5400 DATA 5,6,0,-.262922216048,.103016745283 5410 DATA 5,7,0,-.254211554548..0510773809802 5420 DATA 5,8,0,.0914781725126,0 5430 DATA 5,16,1,.0697840705662,-.0442304181914 5440 DATA 5,20,1,-.276334047244,.167501525476 5450 DATA 5,21,1,.28849914471,.000594132862558 5470 DATA 6,6,0,.333321393988,-.273298586424 5480 DATA 6,7,0,.290995674509,-.214876283361 5490 DATA 6,8,0,-.093065914028,.0587868211474 5500 DATA 5,15,1,-.0428255863778,0 5510 DATA 6,20,1,.174036720164,.00737992993753 5520 DATA 6,21,1,-.29377409633,-.185808381849 5530 5540 DATA 7,7,0,.754497398141,-.0612857429547 5550 DATA 7,8,0,-.539541002456,0 5560 DATA 7,13,0,0,.0914781725126 HILL 1 AND 4

5570 DATA 7,14,0,.0587868211474,-.0930659514028 ..... 3 AND 4 DATA 7,19,1,0,.0575832193445 5520 DATA 7,20,1,.103535317135.0 5530 DATA 7,21,1,-.485418546828,-.0428370961158 5600 DATA 7,22,1,0,.0466704801694 ..... AND 4 56:0 DATA 7,23,1,0,-.259293054322 FFFFF 1 AND 4 5620 TITITITITE Row/Column H FATERIAL COLUMN H 5630 DATA 8,8,0,.764497398141,~.0612867429647 (110) 1 AND 4 5540 DATA 8,13,0,.0510773809802,-.254211554548 ++++ 3 AND 4 5650 DATA 8,14,0,-.214875283361,.290995574509 +++++ 3 AND 4 5660 DATA 8,19,1,0,-.366404054596 5670 DATA 8,20,1,-.0466704801694,0 5680 5690 DATA 8,21,1,.259293054322,0 DATA 8,22,1,0,-.103635317135 1111 1 AND 4 5700 DATA 8,23,1,.0428370961158,.485418546828 (1111 3 AND 4 5710 5720 5730 DATA 9,9,0,.566893424036,0 5740 DATA 9,10,0,-.320683347477,-.320683347477 5750 DATA 9,11,0,.150341673739,.160341673739 5760 DATA 9,12,0,-.113378684807,0 5770 DATA 9,17,1,.333006762504,-.333006762504 5780 DATA 9,18,1,-.183860876595..183860876595 5790 DATA 9,22,1,.0347149906857,-.0347149906857 5800 5810 DATA 10,10,0,.566893424036,.226757369615 5820 DATA 10.11.0.-.453514739229.-.113378684807 5830 DATA 10,12,0,.320683347478,0 5840 DATA 10,13,0,.0805797038531,.0510728876983 DATA 10,14,0,-.0494507276469,0 5850 DATA 10,17,1,-.470942679895,.26001854527 5860 5870 DATA 10,18,1,.470942679895,-.0490944106454 5880 DATA 10,22,1,-.226146200577,.0367387608396 5890 DATA 10,23,1,.0389677051594,0 5900 5910 DATA 11,11,0,.566893424036,.226757369615 5920 DATA 11,12,0,-.641366694954,0 5930 DATA 11,13,0,-.212232295405,-.18272547925 5940 DATA 11,14,0,.0989014552938,.0494507276469 5950 DATA 11,17,1,.26001854527,-.0490944106454 5960 DATA 11,18,1,-.470942679895,.0490944106454 5970 DATA 11,22,1,.390842340703,-.201434900966 5980 DATA 11,23,1,-.0779354103188..0389677051594 5990 5000 DATA 12,12,0..566893424036,0 6010 DATA 12,13,0,.186184880494,.186184880494 DATA 12,14,0,-.0699338897075,-.0699338897075 6020 5030 DATA 12,17,1,-.0347149906857,.0347149906857 DATA 12,18,1,.183860876595,~.183860876595 5040 6050 DATA 12,22,1,-.267630505722,.267630505722 6060 DATA 12,23,1,.055108657131,-.0551086571318 TEEPEREN Row/Column Mterrererererererererere 6070 DATA 13,13,0,.32337728364,.10262392793 5080 6090 DATA 13,14,0,-.262922216048,.103016745283 DATA 13,18,1,.0697840705662,-.0442304181914 6100 6110 DATA 13,19,1,-.269084916899,0 DATA 13,22,1,-.276234047244,.167501525476 6120

6130 DATA 13,23,1,.28849914471,.000594132852558

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6150 DATA 14,14,0..333321393988,-.273298586424 6150 DATA 14,18,1,-.0428255863778,0 6170 DATA 14,19,1,.445083571317,0 5180 DATA 14,22,1,.174036720164,.00737992993753 6190 DATA 14,23,1,-.29377409633,-.185808381849 6210 DATA 15,15,1,1.4043443166,-.280297304107 6220 DATA 15,16,1,-.280297304107,.0425170068027 6230 DATA 15,20,1,.0425170068027,0 6240 6250 DATA 16,16,1,1.4043443166.0 DATA 16,20,1,-.250962993117,.0318167001907 6260 DATA 16,21,1,.0337470225952,0 6270 6280 6290 DATA 17,17,1,1.4043443166,-.280297304107 6300 DATA 17,18,1,-.280297304107,.0425170058027 DATA 17,22,1,.0425170058027,0 6310 6320 DATA 18,18,1,1.4043443156,0 6330 6340 DATA 18,22,1,-.250962993117,.0318167001907 6350 DATA 18,23,1,.0337470225952,0 6360 6370 DATA 19,19,1,1.55488461455,0 6380 DATA 19,22,0,.022697123287,.022697123287 6390 DATA 19,23,0,-.148555354853,-.148555354853 6400 6410 DATA 20,20,1,.386448961524,-.16585112319 6420 DATA 20,21,1,~.188670742999,.0252538685337 6430 6440 DATA 21,21,1,.421776560109,.0299414965597 6450 DATA 21,23,1,0,.0409943591762 !!!!! 1 AND 4 6460 5470 DATA 22,22,1,.386448961524,-.16585112319 6480 DATA 22,23,1,-.188670742999,.0252538685337 6490 6500 DATA 23,23,1,.421776560109,.0299414965597 DATA 9999 6510 DATA G Matrix Loaded 6520 END 6530 6540 6550 <sup>3</sup> Subprogram Eigen is attached at this point in the program. A listing 6560 ! of Eigen is given at the end of VDWNCA. 6570 5580 

Distant Production

10 1 F and G matrix elements for free base phthalocyanine used to 20 <sup>1</sup> calculate B1u vibrations in program H2PCNCA. 30 40 50 60 70 F matrix elements. 80 90 DATA 1,1,0,.119,0 + Row/Column A. 100 DATA 2,2,0,.119,0 + Row/Column B. 110 DATA 3,3,0,.119,0 ' Row/Column C. 120 130 DATA 4,4,0,.119,0 ' Row/Column D. DATA 5,5,0,.119,0 1 Row/Column E. 140 150 DATA 5,5,0,.119,0 + Row/Column F. DATA 7,7,0,.119,0 + Row/Column 6. 160 DATA 8,8,0,.119,0 1 Row/Column H. 170 DATA 9,9,0,.119,0 ! Row/Column I. 180 190 DATA 10,10,0,.119,0 + Row/Column J. DATA 11,11,0,.119,0 + Row/Column K. 200 210 DATA 12,12,0,.119,0 ! Row/Column M. 220 DATA 13,13,0,.119,0 + Row/Column N. 230 DATA 14,14,0,.119,0 1 Row/Column 0. 240 DATA 15,15,0,.3237,0 ! Row/Column P. 250 DATA 15,15,0,.3237,0 ! Row/Column Q. 260 DATA 17,17,0,.3237,0 / Row/Column R. 270 DATA 18,18,0,.3237,0 + Row/Column S. DATA 19,19,0,.3237,0 1 Row/Column T. 280 290 DATA 20,20,0,.3237,0 | Row/Column U. DATA 21,21,0,.3237,0 1 Row/Column V. 300 DATA 22,22,0,.3237,0 1 Row/Column W. 310 DATA 23,23,0,.3237,0 ! 320 DATA 9999 330 DATA F Matrix Loaded 340 350 360 370 1 6 matrix elements. 380 390 400 410 DATA 1,1,0,.566893424036,0 DATA 1,2,0,-.320683347477,-.320683347477 420 430 DATA 1,3,0,.150341673739,.150341673739 DATA 1,4,0,~.113378584807,0 440 450 DATA 1,15,1,.333006762504,-.333006762504 DATA 1,16,1,-.183860876595..183860876595 460 470 DATA 1,20,1,.0347149906857,-.0347149906857 480 490 DATA 2,2,1,.565893424036,.226757369615 DATA 2,3,1,-.453514739229,-.113378684807 500 510 DATA 2.4.1..320683347478.0 DATA 2,5,1,.0805797038531,.0510728876953 522

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530 DATA 2.6.1.-.0494507276469.0 540 DATA 2.15.0.-.470942679895..26001854527 550 DATA 2,16,0..470942679895,-.0490944106454 560 DATA 2,20,0,-.226146200577,.0367387608396 570 DATA 2,21,0,.0389677051594,0 580 DATA 3,3,1,.566893424036,.226757369615 590 600 DATA 3,4,0,-.641366694954,0 DATA 3,5,1,-.212232295405,-.18272547925 610 620 DATA 3,6,1,.0989014552938,.0494507276469 DATA 3,15,0,.26001854527,-.0490944106454 630 540 DATA 3,16,0,-.470942679895,.0490944106454 650 DATA 3,20,0,.390842340703,-.201434900966 660 DATA 3,21,0,-.0779354103188,.0389677051594 670 680 DATA 4,4,0,.566893424036,0 690 DATA 4,5,0,.186184880494,.186184880494 700 DATA 4,6,0,-.0699338897075,-.0699338897075 710 DATA 4,15,1,-.0347149906857,.0347149906857 720 DATA 4,15,1,.183860876595,-.183860876595 730 DATA 4,20,1,-.267630505722,.267630505722 740 DATA 4,21,1,.055108657131,-.055108657131 750 750 DATA 5.5.1.32337728364.10262392793 770 DATA 5,6,1,-.262922216048,.103016745283 780 DATA 5,7,1,-.254211554548,.0510773809802 790 DATA 5,8,1,.0914781725126,0 800 DATA 5,16,0,.0697840705662,-.0442304181914 810 DATA 5,20,0,-.276334047244,.167501525476 820 DATA 5,21,0,.28849914471,.000594132862558 830 DATA 5,6,1,.333321393988,-.273298586424 840 DATA 5,7,1,.290995674509,-.214876283361 850 860 DATA 6,8,1,-.093065914028,.0587868211474 870 DATA 6,16,0,-.0428255863778,0 880 DATA 6.20.0..174036720164..00737992993753 DATA 6,21,0,-.29377409633,-.185808381849 890 900 910 DATA 7.7.1..764497398141,-.0612867429647 920 DATA 7.8.1.-.59954100245E.0 930 DATA 7,13,1,0,.0914781725126 1111 1 AND 4 940 DATA 7,14,1,.0587868211474,-.0930659514028 \*\*\*\*\* 3 AND 4 950 DATA 7,19.0,0,.0676832193445 960 DATA 7,20,0,.103635317135,0 970 DATA 7,21,0,-.485418546828,-.0428370961158 980 DATA 7,22,0,0,.0466704801694 11111 1 AND 4 DATA 7.23.0,0,-.259293054322 ..... 1 AND 4 990 1000 DATA 8,8,1,.764497398141,-.0612867429647 .... 1 AND 4 1010 DATA 8,13,1,.0510773809802,-.254211554548 ++++ 3 AND 4 1020 DATA 8,14,1,-.214875283361,.290995674509 11111 3 AND 4 1030 1040 DATA 8,19,0,0,-.366404054596 1050 DATA 8,20,0,-.0465704801694,0 DATA 8,21,0,.259293054322,0 1050 DATA 8,22,0,0,-.103635317135 (1)11 1 AND 4 1070 DATA 8,23,0,.0428370961158..485418546828 +++++ 3 AND 4 1032 ARCONTRACTOR Row/Column I FILIPPETER HELPPETER DEFENSION 1090

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1100 DATA 9,9,1,.566893424036.0 DATA 9,10,1,-.320583347477,-.320683347477 1110 DATA 9,11,1,.160341673739,.160341673739 1120 1130 DATA 9,12,1,-.113378684807,0 1140 DATA 9,17,0,.333006762504,-.333006762504 DATA 9,18,0,-.183860876595,.183860876595 1150 DATA 9,22.0.0347149906857.-.0347149906857 1160 1170 DATA 10,10,1,.566893424036,.226757369615 1160 1190 DATA 10,11,1,-.453514739229,-.113378684807 DATA 10,12,1,.320683347478,0 1200 DATA 10.13,1,.0805797038531,.0510728876983 1210 1220 DATA 10,14,1,-.0494507276469,0 1230 DATA 10,17,0,+.470942679895,.26001854527 DATA 10,18,0,.470942579895,-.0490944105454 1240 DATA 10,22.0.-.226146200577..0367387608396 1250 1260 DATA 10,23,0,.0389677051594,0 1270 1280 DATA 11,11,1,566893424036,.226757369615 1290 DATA 11,12,1,-.641366694954,0 1300 DATA 11,13,1,-.212232295405,-.18272547925 DATA 11,14,1,.0989014552938,.0494507276469 1310 DATA 11,17,0,.26001854527,~.0490944106454 1320 1330 DATA 11,18,0,-.470942679895,.0490944106454 1340 DATA 11,22,0,.390842340703,-.201434900966 DATA 11,23,0,-.0779354103188,.0389677051594 1350 1360 1370 DATA 12,12,1,.566893424036,0 1380 DATA 12,13,1,.186184880494,.186184880494 1390 DATA 12,14,1,-.0699338897075,-.0699338897075 1400 DATA 12,17,0,-.0347149906857,.0347149906857 1410 DATA 12,18,0,.183860876595,-.183860876595 1420 DATA 12,22,0,-.267630505722,.267630505722 1430 DATA 12,23,0,.055108657131,-.0551086571318 1440 1450 DATA 13,13,1,.32337728364,.10262392793 1460 DATA 13,14,1,-.262922216048,.103016745283 1470 DATA 13,18,0,.0697840705662,-.0442304181914 1480 DATA 13,19,0,-.269084916899,0 DATA 13,22,0,-.276334047244..167501525476 1490 DATA 13,23,0,.28849914471,.000594132862558 1500 1520 DATA 14,14,1,.333321393988,-.273298586424 1530 DATA 14,18,0,-.0428255863778,0 1540 DATA 14,19,0,.445083571317,0 1550 DATA 14,22,0,.174036720164,.00737992993753 1560 DATA 14,23,0,-.29377409633,-.185808381849 1580 DATA 15,15,0,1.4043443166,-.280297304107 1590 DATA 15,16,0,-.280297304107,.0425170068027 1600 DATA 15,20,0,.0425170068027,0 1E10 1620 DATA 16,16,0,1.4043443166,0 1600 DATA 16,20,0,-.250962993117,.0318167001907 1540 DATA 18,21,0,.0337470225952,0 - FILERALLER FL Row/Column C FILERER FLEERER FLEERER FLEERER FLEERER 1650 1660 DATA 17,17,0,1.4043443166,-.280297304107

1670 DATA 17,18,0,-.280297304107,.0425170068027 1690 DATA 17,22,0,.0425170068027,0 1700 DATA 18,18,0,1.4043443166,0 1710 DATA 18,22,0,-.250962993117,.0318167001907 1720 DATA 18,23,0,.0337470225952,0 1740 DATA 19,19,0,1.55488461455,0 1750 DATA 19,19,0,1.00086799405,0 DEUTERATED 1760 DATA 19,22,0..022697123287,.022697123287 1770 DATA 19,23,0,-.148555354853,-.148555354853 1790 DATA 20,20,0,.386448961524,-.16585112319 1800 DATA 20,21,0,-.188670742999,.0252538685337 1810 1820 DATA 21,21,0,.421776560109,.0299414965597 1830 DATA 21,23,0,0,.0409943591762 1111 AND 4 1840 1850 DATA 22,22,0,.386448961524,-.16585112319 1850 DATA 22,23,0,-.188670742999,.0252538685337 1870 1880 DATA 23,23,0,.421776560109,.0299414965597 1890 DATA 9999 1900 DATA 6 Matrix Loaded 1910 END

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10 F and G matrix elements for free base phthalocyanine used to calculate 20 30 B3g vibrations in program H2PCNCA. 40 50 Ŧ 50 70 F F matrix elements. 80 90 1 DATA 1,1,0,.119,0 + Row/Column A. 100 DATA 2,2,0,.119,0 | Row/Column B. 110 DATA 3,3,0,.119,0 1 Row/Column C. 120 130 DATA 4,4,0,.119,0 | Row/Column D. 140 DATA 5,5,0,.119,0 ! Row/Column E. 150 DATA 5,6,0,.113,0 + Row/Column F. 160 DATA 7,7,0,.119,0 ! Row/Column G. 170 DATA 8,8,0,.119,0 ! Row/Column H. 180 DATA 9,9,0,.119,0 ! Row/Column I. 190 DATA 10,10,0,.119,0 ! Row/Column J. 200 DATA 11,11,0,.119,0 ! Row/Column K. 210 DATA 12,12,0,.119,0 ! Row/Column L. DATA 13,13,0,.119,0 ! Row/Column M. 220 DATA 14,14,0,.119,0 ' Row/Column N. 230 240 DATA 15,15,0,.3237,0 ! Row/Column 0. 250 DATA 16,16,0,.3237,0 | Row/Column P. DATA 17,17,0,.3237,0 ' Row/Column Q. 260 270 DATA 18,18,0,.3237,0 ! Row/Column R. 280 DATA 19,19,0,.3237,0 ! Row/Column S. 290 DATA 20,20,0,.3237,0 + Row/Column T. 300 DATA 21,21,0,.3237,0 ! Row/Column U. 310 DATA 22,22,0,.3237,0 + Row/Column V. 320 DATA 23,23,0,.3237,0 ! Row/Column W. 330 DATA 9999 340 DATA F Matrix Loaded 350 TERETER CONTRACTOR CONTRA 360 370 1 6 matrix elements. 380 390 400 410 DATA 1,1,1,.566893424036.0 420 DATA 1,2,0,-.320683347477,-.320683347477 430 DATA 1,3,0,.160341673739,.160341673739 440 DATA 1,4,1,-.113378684807,0 450 DATA 1,15,1,.333006762504,-.333006762504 460 DATA 1,16,1,-.183860876595,.183860876595 470 DATA 1,20,1,.0347149906857,-.0347149906857 480 ITTELETER Row/Column B Itteleters in the construction of the second 490 DATA 2,2,0,.566893424036,.226757369615 500 DATA 2,3,0,-.453514739229,-.113378684807 510 DATA 2,4,1,.320583347478,0 520 DATA 2,5,0,.0805797038531,.0510728876983

DATA 2,5,0,-.0494507275469,0 530 DATA 2,15,1,-.470942579895,.26001854527 540 DATA 2,16,1,.470942679995,-.0490944106454 550 DATA 2,20,1,-.226146200577,.0367387608336 550 DATA 2,21,1,.0389677051594,0 570 LIBELERIER Row/Column C. SECTION SECTION 530 DATA 3,3,0,.566893424035,.226757369615 590 600 DATA 3,4,1,-.641366694954,0 DATA 3,5,0,-.212232295405,-.18272547925 610 DATA 3,5,0,.0989014552938,.0494507275469 620 630 DATA 3,15,1,.26001854527,-.0490944106454 DATA 3,16,1,-.470942679895,.0490944106454 540 DATA 3,20,1,.390842340703,-.201434900966 650 660 DATA 3,21,1,-.0779354103188,.0389677051594 670 680 DATA 4,4,1,.565893424036,0 690 DATA 4,5,0,.186184880494,.186184880494 700 DATA 4.6.0.-.0699338897075.-.0699338897075 DATA 4,15,1,-.0347149906857,.0347149906857 710 720 DATA 4,16,1,.183860876595,-.183860876595 730 DATA 4,20,1,-.267630505722,.267630505722 740 DATA 4,21,1,.055108657131,-.055108657131 750 760 DATA 5,5,0,.32337728364..10262392793 770 DATA 5.6.0.-.262922216048..103016745283 780 DATA 5.7.0.-.254211554548..0510773809802 790 DATA 5.8.0..0914781725126.0 800 DATA 5,16,1,.0697840705662,-.0442304181914 810 DATA 5,20,1,-.276334047244,.167501525476 820 DATA 5,21,1,.28849914471,.000594132862558 830 840 DATA 6,6,0,.333321393988,-.273298586424 850 DATA 6,7,0,.290995674509,-.214876283361 860 DATA 6.8.0,-.093065914028.0587868211474 870 DATA 6,16,1,-.0428255863778,0 880 DATA 6,20,1,.174036720164..00737992993753 890 DATA 6,21,1,-.29377409633,-.185808381849 900 910 DATA 7,7,0,.764497398141,-.0612867429647 920 DATA 7.8.0,-.599541002456.0 930 DATA 7,13,0,0,.0914781725126 + ++++ + AND 4 940 DATA 7,14,0,-.0587868211474,-.0930659514028 +++++ 3 AND 4 950 DATA 7,19,1,0..0676832193445 960 DATA 7,20,1,.103635317135,0 970 DATA 7,21,1,-.485418546828,-.0428370961158 980 DATA 7,22,1,0,.0466704801594 1111 1 AND 4 990 DATA 7,23,1,0,-.259293054322 +++++ 1 AND 4 - TEFFETTER FILL Row/Column H. FELTERETER FELTER FLETTER FELTER FELTER FELTER FELTER FE 1000 1010 DATA 8,8,1,.764497398141,-.0612867429647 (1919) 1 AND 4 1020 DATA 8,13,0,-.0510773809802,-.254211554548 1111 3 AND 4 1030 DATA 8,14,0,.214876283361,.290995674509 (1111 3 AND 4 1040 DATA 8,19,1,0,-.365404054596 1050 DATA 8,20,1,-.0466704801694,0 1060 DATA 8,21,1,.259293054322,0 1070 DATA 8,22,1,0,-.103635317135 (11) 1 AND 4 1080 DATA 8,23,1,-.0428370961158..485418546828 ..... 3 AND 4 1090 

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1100 DATA 9,3,0,.565893424036,0 DATA 9,10,0,-.320693347477,-.320683247477 1110 DATA 9,11,0,.160341673739,.160341673739 1120 DATA 9,12,0,-.112378684807.0 1120 DATA 9,17.1,.333006762504,-.332006752504 1140 DATA 9,18,1,-.183860876595,.183860876595 1150 DATA 9,22,1,.0347149906857,-.0347149906857 1160 1170 DATA 10,10,1.,566893424036,.226757369615 1180 DATA 10,11,1,-.453514739229,-.113378684807 1190 DATA 10,13,1,.0805797038531,.0510728876983 1200 1210 DATA 10,14,1,-.0494507276469,0 DATA 10,17,0,-.470942679895,.26001854527 1220 DATA 10,18,0,.470942679895,-.0490944106454 1230 DATA 10,22,0,-.226146200577,.0367387608396 1240 1250 DATA 10,23,0,.0389677051594,0 1250 DATA 11,11,1.566893424036,.226757369615 1270 1280 DATA 11,13,1,-.212232295405,-.18272547925 1290 DATA 11,14,1,.0989014552938..0494507276469 DATA 11,17,0,.26001854527,-.0490944106454 1300 DATA 11,18,0,-.470942679895,.0490944106454 1310 1320 DATA 11,22,0..390842340703,-.201434900966 1330 DATA 11,23,0,-.0779354103188,.0389677051594 1340 1350 DATA 12,12,0,.566893426036,0 DATA 12,13,0,.186184880494,.186184880494 1360 1370 DATA 12,14,0,-.0699338897075,-.0699338897075 1380 DATA 12,17,1,-.0347149906857,.0347149906857 1390 DATA 12,18,1,.183860876595,-.183860876595 1400 DATA 12,22,1,-.267630505722,.267530505722 1410 DATA 12,23,1,.055108657131,-.0551086571318 1420 1430 DATA 13,13,1,.32337728364,.10262392793 1440 DATA 13,14,1,-.262922216048,.103016745283 1450 DATA 13,18,0,.0597840705662,-.0442304181914 1460 DATA 13,19,1,-.269084916899,0 1470 DATA 13,22,0,-.276334047244,.167501525476 DATA 13,23,0,.28849914471,.000594132862558 1480 1490 DATA 14,14,1,.333321393988,-.273298586424 1500 1510 DATA 14,18,0,-.0428255863778,0 1520 DATA 14,19,1..445083571317,0 1530 DATA 14,22,0,.174036720164,.00737992993753 1540 DATA 14,23,0,-.29377409633,~.185808381849 1550 1560 DATA 15,15,1,1.4043443166,-.280297304107 DATA 15,16,1,-.280297304107,.0425170068027 1570 1580 DATA 15,20,1,.0425170068027,0 1590 THE CONTRACTOR Row/Column P CONTRACTOR DISTORTS FOR DISTORTS FOR DISTORTS 1600 DATA 15,15,1,1.4043443165,0 1610 DATA 16,20,1,-.250962993117,.0318167001907 1620 DATA 16,21,1,.0337470225952,0 INTEREST Row/Column Q Institutes interesting the second se 1630 DATA 17,17,0,1.4043443166,-.280297304107 1640 1650 DATA 17,18,0,-.280297304107,.0425170068027 1660 DATA 17,22,0,.0425170068027.0

1590 DATA 18,19,0,1.4043443166,0 1530 CATA 15,22,0.-.250962993117,.0318167001907 1700 CATA 18,23,0,.0337470225952,0 statettivette Row/Column 5 villeventarettarettarettarettarettaret 6171 1720 DATA 19,19,1,1.55488461455,0 1720 DATA 19,22,0,.022697123287,.022697123287 1740 DATA 13,23,0,-.148555354853,-.148555354853 1750 DATA 20,20,1,.386448961524,-.16585112319 1750 DATA 20,21,1,-.188670742999,.0252538685337 1770 1780 1730 DATA 21,21,1,.421776560109,.0299414965597 1800 DATA 21,23,1,0,.0409943591762 +++++ 1 AND 4 1810 DATA 22,22,0,.386448961524,-.16585112319 1820 1830 DATA 22,23,0,-.188670742999.0252538685337 1840 DATA 23,23,0,.421776560109,.0299414965597 1850 DATA 9999 1860 DATA G Matrix Loaded 1870 1880 END

10 F and G matrix elements for free base phthalocyanine used to calculate 20 B2g vibrations in H2PCNCA. 30 40 50 60 70 F matrix elements. 80 90 ŧ 100 DATA 1,1,0,.119,0 + Row/Column A. 110 DATA 2,2,0,.119,0 + Row/Column B. 120 DATA 3,3,0,.119,0 + Row/Column C. DATA 4,4,0,.119,0 + Row/Column D. 130 140 DATA 5,5,0,.119,0 ! Row/Column E. 150 DATA 5,6,0,.119,0 ! Row/Column F. DATA 7,7,0,.119,0 ! Row/Column 6. 160 170 DATA 8,8,0,.119,0 / Row/Column H. 180 DATA 9,9,0,.119,0 ! Row/Column I. 190 DATA 10,10,0,.119,0 ! Row/Column J. 200 DATA 11,11,0,.119,0 ! Row/Column K. 210 DATA 12,12,0,.119,0 ! Row/Column L. 220 DATA 13,13,0,.119,0 ! Row/Column M. 230 DATA 14,14,0,.119,0 1 Row/Column N. DATA 15,15,0,.3237,0 ! Row/Column 0. 240 DATA 15,15,0,.3237,0 ! Row/Column P. 250 260 DATA 17,17,0,.3237,0 ! Row/Column Q. 270 DATA 18,18,0,.3237,0 ! Row/Column R. DATA 19,19,0,.3237,0 ! Row/Column S. 280 290 DATA 20,20,0,.3237,0 ! Row/Column T. DATA 21,21,0,.3237,0 ! Row/Column U. 300 310 DATA 22,22,0,.3237,0 + Row/Column V. 320 DATA 23,23,0,.3237,0 ! Row/Column W. 330 DATA 9999 340 DATA F Matrix Loaded 350 360 370 ! 6 matrix elements. 380 390 400 410 DATA 1,1,1,.566893424036,0 420 DATA 1,2,0,-.320683347477,-.320683347477 430 DATA 1,3,0,.160341673739,.160341673739 440 DATA 1,4,1,-.113378684807,0 450 DATA 1,15,1,.333006762504,-.333006762504 DATA 1,16,1,-.183860876595,.183860876595 460 DATA 1,20,1,.0347149906857,-.0347149906857 470 480 DATA 2,2,1,.566893424036,.226757369615 490 500 DATA 2,3,1,-.453514739229,-.113378684807 DATA 2,4,1,.320683347478,0 510 520 DATA 2,5,1,.0805797038531,.0510728876983

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DATA 2,6,1,-.0494507276469,0 530 DATA 2,15,0,-.470942679895,.25001854527 340 DATA 2,16,0,.470942679895.-.0430944106454 550 DATA 2,20,0,-.225146200577,.0367387608396 560 DATA 2.21,0,.0389677051594,0 570 TELEVISION Row/Column C 1994 Freedore Freedore Provide Freedore Free 580 DATA 3,3,1,.566893424036,.226757369615 550 600 DATA 3,4,1,-.641366694954,0 DATA 3,5,1,-.212232295405,-.18272547925 610 DATA 3,6,1,.0989014552938..0494507276469 620 DATA 3,15,0,.26001854527,-.0490944106454 630 DATA 3,16,0,-.470942679895,.0490944106454 640 650 DATA 3,20,0,.390842340703,-.201434900966 660 DATA 3,21,0,-.0779354103188,.0389677051594 670 680 DATA 4,4,1,.566893424036,0 690 DATA 4,5,0,.186184880494..186184880494 DATA 4,6,0,-.0699338897075,-.0699338897075 700 DATA 4,15,1,~.0347149906857,.0347149906857 710 720 DATA 4,16,1,.183860876595,-.183860876595 730 DATA 4,20,1,-.267630505722,.267630505722 740 DATA 4,21,1,.055108657131,-.055108657131 750 HILLING Row/Column E HILLING HILLING HILLING HILLING HILLING HILLING 760 DATA 5,5,1,.32337728364,.10262392793 770 DATA 5,6,1,-.262922216048,.103016745283 780 DATA 5,7,1,-.254211554548,.0510773809802 790 DATA 5,8,1,.0914781725126,0 800 DATA 5,15,0,.0597840705652,-.0442304181914 810 DATA 5,20,0,-.275334047244,.167501525476 820 DATA 5,21,0,.28849914471,.000594132862558 830 840 DATA 5,6,1,.333321393988,-.273298586424 850 DATA 6,7,1,.290995674509,-.214876283361 860 DATA 5,8,1,-.093055914028,.0587868211474 870 DATA 5,15,0,-.0428255863778,0 880 DATA 5,20,0,.174036720164,.00737992993753 890 DATA 5.21.0.-.29377409533.-.185808381849 900 DATA 7,7,1,.764497398141,-.0612867429647 910 920 DATA 7.8.1.-.599541002456.0 930 DATA 7,13,1,0,.0914781725125 1111 1 AND 4 940 DATA 7,14,1,-.0587868211474,-.0930659514028 +++++ 3 AND 4 DATA 7,19,0,0,.0676832193445 950 DATA 7,20,0,.103635317135,0 960 970 DATA 7,21,0,-.485418546828,-.0428370961158 980 DATA 7,22.0.0..0456704801694 1111 1 AND 4 990 DATA 7,23,0,0,-.259293054322 1111 1 AND 4 1000 DATA 8.8.0..764497398141.-.0612867429647 IIII I AND 4 1010 1020 DATA 8,13,1,-.0510773809802,-.254211554548 1111 3 AND 4 DATA 8,14,1,.214876283361,.290995674509 (111) 3 AND 4 1030 1040 DATA 8,19,0,0,-.366404054596 1050 DATA 8,20,0,-.0466704801694,0 1060 DATA 8.21.0..259293054322.0 1070 DATA 8,22,0,0,-.103635317135 1111 1 AND 4 1080 DATA 8,23,0,-.0428370961158,.485418546828 1111 3 AND 4 1090 

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1100 DATA 9,9,1,.566893424036,0 1110 DATA 9,10,0,-.320683347477,-.320683347477 1120 DATA 9,11,0,.150341673739,.150341673739 1120 DATA 9,12,1,-.113378694807,0 1140 DATA 9,17,1,.333006762504,-.333006762504 1150 DATA 9,18,1,-.183860876595,.183860876595 1160 DATA 9.22.1.0347149906857.-.0347149906857 1160 DATA 10,10,0..566893424036..226757369615 1190 DATA 10,11,0,-.453514739229,-.113378684807 1200 DATA 10,12,1,.320683347478,0 1210 DATA 10,13,0..0805797038531,.0510729876983 1220 DATA 10,14,0,-.0494507276469,0 1230 DATA 10,17,1,-.470942679895,.26001854527 1240 DATA 10,18,1,.470942679895,-.0490944106454 1250 DATA 10,22,1,-.226146200577,.0367387608396 1260 DATA 10,23,1,.0389677051594,0 1270 1280 DATA 11,11,0,.566893424036,.226757369615 1290 DATA 11,12,1,-.641366694954,0 1300 DATA 11,13,0,-.212232295405,-.18272547925 1310 DATA 11,14,0..0989014552938,.0494507276469 1320 DATA 11,17,1..26001854527,-.0490944106454 1330 DATA 11,18,1,-.470942679895,.0490944106454 1340 DATA 11,22,1,.390842340703,-.201434900966 DATA 11,23,1,-.0779354103188,.0389677051594 1350 1360 1370 DATA 12,12,1,.566633426036,0 1380 DATA 12,13,0,.185184880494,.186184880494 1390 DATA 12,14,0,-.0699338897075,-.0699338897075 1400 DATA 12,17,1,-.0347149906857,.0347149906857 1410 DATA 12,18,1,.183860876595,-.183860876595 1420 DATA 12,22,1,-.267630505722..267630505722 1430 DATA 12.23.1..055108657131.-.0551086571318 1440 1450 DATA 13,13,0,.32337728364,.10262392793 1460 DATA 13,14,0,-.262922216048,.103016745283 1470 DATA 13,18,1,.0697840705662,-.0442304181914 1480 DATA 13,19,1,-.269084916899,0 DATA 13,22,1,-.276334047244,.167501525476 1490 1500 DATA 13,23,1,.28849914471,.000594132862558 1510 1520 DATA 14,14,0,.333321393988,-.273298586424 1530 DATA 14,18,1,-.0428255863778,0 1540 DATA 14,19,1,.445083571317,0 DATA 14,22,1,.174036720164,.00737992993753 1550 1560 DATA 14,23,1,-.29377409533,-.185808381849 1570 1580 DATA 15,15,0,1.4043443166,-.280297304107 DATA 15,16,0,-.280297304107,.0425170068027 1590 DATA 15,20,0,.0425170068027,0 1600 1610 1620 DATA 15,15,0,1.4043443166,0 DATA 16,20,0,-.250962993117,.0318167001907 1630 1640 DATA 16,21,0,.0337470225952,0 1650 1560 DATA 17,17,1,1.4043443166,-.280297304107

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1570 DATA 17,13,1,-.280297304107,.0425170058027 1580 DATA 17,22,1,.0425170068027,0 1530 FRANKELLE Row/Column R FIELEREFELLEREFELLEREFELLER 1720 DATA 18,18,1,1.4043443166,0 1710 DATA 18,22,1,-.250962993117,.0318167001907 1720 EATA 18,23,1,.0337470225952,0 1733 CHARLEN Row/Column S JUNETIAL CHARLEN CHARLEN CONTRACTORS 1740 DATA 19,19,0,1.55488461455,0 1750 DATA 19,22,0,.022697123287,.022697123287 1760 DATA 19,23.0,-.148555354853.-.148555354853 1790 DATA 20,20,0,.386448961524,-.16585112319 1790 DATA 20,21,0,-.188670742999,.0252538685337 1800 1810 DATA 21,21,0,.421776560109,.0299414965597 1820 DATA 21,23,0,0,.0409943591762 +++++ 1 AND 4 1830 1840 DATA 22,22,1,.386448961524,-.16585112319 DATA 22,23,1,-.188670742999,.0252538685337 1850 1850 1870 DATA 23,23,1,.421776560109,.0299414965597 1880 DATA 9999

- 1890 DATA G Matrix Loaded
- 1900 END

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APPENDIX SIX

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COCO SCALE

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"3DHRRA"

10 -C-----20 E C Program: 3DHRRA 30 1 C This program calculates vdW torsional modes for benzene(methane)) type 40 1 C clusters. The torsional modes are calculated for a specified 50 1 C torsional barrier height and torsional potential form. The 60 1 70 1 C torsional modes are treated using rigid rotor C symmetric top wavefunctions as a basis set. Under free rotation, these 80 1 C wavefunctions are solutions to the spherical top Schrödinger equation. 90 1 C Upon application of the torsional potential, the torsional mode 1001 C eigenvalues are obtained by diagonalizing the energy matrix 110! C which describes the hindered rotational/librational motion. 1201 C-----130! 1401 С PROGRAM 3DHRRA 150 ! 160 ! C 170 · C-----180 ! C A(+)=Energy matrix to be diagonalized. 190 | C D(+)=Diagonal elements of tridiagonal matrix, or eigenvalues of 200 ! C diagonalized matrix. C E(+)=Subdiagonal elements of tridiagonal matrix. 210 1 220 ! C ZAP(+)=Orthogonal transformation matrix to tridiagonalize A(+). C-----230 240 ! C 250 ! DIMENSION A(680,680),D(680),E(680),ZAP(680) 250 ! С 270 1 C---280 ! C Identifies external functions for integrations subroutine. 290 ! C-----300 ! C EXTERNAL THINT, XINT, PHINT 310 1 С 320 ! 330 C------\_\_\_\_\_\_ C Sets up common blocks. 340 1 350 1 360 ! C COMMON /BIA/A/B2ZAP/ZAP/B3D/D/B4E/E/B5F/RJ,RK,RM,CJ,CK,CM 370 1 380 1 + ,ZA,ZB,ZC,ZD,ZE,ZF,ZG,ZH,FLG1 390 I C C-----400 1 410 : C Declares integers. 420 · C-----\_\_\_\_\_ 430 I C 440 1 INTEGER RE, CE, ROWTOT, ORDER, EJ, EIGVLE, SJ, Z, FLG1 450 I C -450 · C-----470 ' C Determines the order of the energy matrix in terms of J quantum number 480 · C SJ=Starting J level. 490 · C EJ Ending J level. 500 · C-----510 / C 520 1 SJ=0

EJ=7 530 1 540 i ORDER=0 550 / DO 19 Z=0.EJ ORDER=ORDER+(2+Z+1)++2 560 1 570 1 19 CONTINUE 580 · C 590 | C-----\_\_\_\_\_ 500 ! C Initializes the energy matrix. 610 / C-----620 ! С 630 I DO 85 I=1,0RDER 640 ! 00 86 J=1,I 650 1 A(I,J)=0.0 86 CONTINUE 660 ! 670 1 85 CONTINUE 680 ! C 690 ! C---700 ! C Maximum size of energy matrix. 710 I C---720 ! C 730 ! ROWTOT=680 740 ! C 750 ! C-----750 ! C Torsional barrier height in wavenumbers. 770 ! C-----789 I C 790 1 VPERT=500 800 I C 819 | C-----820 ! C Effective internal rotational constant for cluster in wavenumbers. 839 ! C -840 1 С 850 ! ROTCA=2.6025 860 1 PI=3.1415927 С 870 1 880 1 -----890 C Determines matrix row element in question. 900 1 C------910 I C 920 1 DO I RJ=SJ,EJ 930 1 DO 2 RM=-RJ\_RJ 940 1 DO 3 RK=-RJ,RJ 950 1 RE-0 960 1 DO 4 Z=0,RJ 970 ! RE=RE+(2+(Z-1)+1)++2 980 ! 4 CONTINUE 990 1 RE=RE+RK+RJ+(2+RJ+1)+(RM+RJ) 10001 С 1010 C---1020! C Determines matrix column element in question. 10301 C-----10401 C 10501 DO 5 CJ=SJ\_EJ 10601 00 6 CM=-CJ,CJ D0 7 CK=-CJ,CJ 10701 10801 CE-0 10901 00 8 Z=0,CJ 11001 CE=CE+(2+(Z-1)+1)++2 11101 8 CONTINUE 11201 CE=CE+CK+CJ+(2+CJ+1)+(CM+CJ)

ويديك المحدجية بالالمحجج فكظ

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1130 C 11401 C---C Inputs diagonal matrix elements into A(+). 11501 C------11601 1170 C IF (RE.EQ.CE) A(RE,CE)=ROTCA+RJ+(RJ+1.)+VPERT/2. 11901 IF (CE.GT.RE) GO TO 7 11901 С 12001 C-----12101 C Selects possible nonzero off-diagonal matrix elements in A(+). 12201 C-----12301 12401 C IF(ABS(RJ-CJ).LE.2.AND.ABS(RK-CK).EQ.2.AND.ABS(RM-CM).EQ. 12501 12601 + 1) GO TO 18 GO TO 7 12701 С 12801 C------12901 C Calculates normalization constants for matrix elements. 13001 C------13101 1320 C 18 ZA=(2+RJ+1)/8/PI/PI 13301 13401 DO 9 Z=RJ+RM.2.-1 1350 ZA=ZA+Z 13601 9 CONTINUE ZA=SQRT(ZA) 13701 1380! Z8=1 DO 10 Z=RJ-RM.2.-1 13901 14001 ZB=ZB+Z 14101 10 CONTINUE 14201 ZB=SQRT(ZB) 14301 ZC=1 DO 11 Z=RJ+RK, 2,-1 14401 14501 ZC=ZC+Z 1460! 11 CONTINUE ZC=SQRT(ZC) 14701 1480! ZD=1 DO 12 Z=RJ-RK .2 .-1 1490! 1500 ZD=ZD+Z 15101 12 CONTINUE ZD=SQRT(ZD) 15201 ZE=(2+CJ+1)/8/PI/PI 15301 1540! DO 13 Z=CJ+CM,2,-1 1550 ZE=ZE+Z 15601 13 CONTINUE 15701 ZE=SQRT(ZE) 15801 ZF=1 DO 14 Z=CJ-CM,2,-1 15901 16001 ZF=ZF+Z 16101 14 CONTINUE ZF=SQRT(ZF) 16201 16301 ZG=1 16401 DO 15 Z=CJ+CK,2,-1 16501 ZG=ZG+Z 16601 15 CONTINUE 1670 ZG=SQRT(ZG) 1680+ ZH=1 DO 16 Z=CJ-CK,2,-1 16901 1700+ ZH=ZH+Z 17101 16 CONTINUE 17201 ZH=SQRT(ZH)

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17301 C 1740! C-----1750) C Integrates over theta, phi, and chi coordinates. 17501 17701 C 00 17 FLG1=0.0 17801 17901 C \*\*=====\*\*\*\* 18001 C-C Integration over theta coordinate between 0 and pi. 18101 C-----1820 1830 C P=0 1840! B=PI 18501 18601 EPSABS=1.E-6 18701 EPSREL=1.E-6 1880! C 1890! C------1900! C Subroutine: STIT 1910! C 1920! C This subroutine calculates an approximation result to a given definite 1930! C integral. The function to be integrated is named THINT. 1940! C The subroutine is from QUADPACK, a FORTRAN subroutine package for the 1950! C numerical computation of definite 1 dimensional integrals. 1960! C Authors: Robert Piessens and Elise de Doncker, Appl. Math. and Progr. 1970! C Div. - K.U. Leuven. 1980! C The subroutine name in QUADPACK is QNG. 19901 C-----2000! C CALL STIT(THINT, P, B, EPSABS, EPSREL, RESULT, ABSERR, NEVAL, IER) 20101 CADRE1=RESULT 2020! 20301 IF (CADREI.EQ.0) GO TO 20 2040! C 2050 C-----20501 C Integration over chi coordinate between 0 and 2\*pi. 2070 C-----2080! С 2090! P=0 B=PI+2 21001 21101 EPSABS=1.E-6 21201 EPSREL=1.E-6 21301 С 2140! C---C Call subroutine STIT to integrate over the function XINT. 21501 21601 21701 С 21801 CALL STIT(XINT.P.B.EPSABS, EPSREL.RESULT, ABSERR, NEVAL, IER) 21901 CADRE2=RESULT 2200! IF (CADREZ.EQ.0) GO TO 20 С 22101 2220! C-2230! C Integration over phi coordinate between 0 and 2\*pi. 2240! C------22501 P=0 2250 B=PI+2 2270 | EPSABS=1.E-6 2280 ( EPSREL=1.E-6 2290 ! C 2300 · C-----2310 + C Call subroutine STIT to integrate over the function PHINT. 2320 | C-----

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2330 · C CALL STIT(PHINT, P, B, EPSABS, EPSREL, RESULT, ABSERR, NEVAL, IER) 2340 1 2350 + CADRE3=RESULT 2360 | 20 CADRE=CADRE1+CADRE2+CADRE3 IF (ABS(CADRE).LT.1.E-5) CADRE=0 2370 1 IF (A(RE,CE).NE.0) A(RE,CE)=A(RE,CE)-CADRE+VPERT/2. 2380 1 IF (A(RE,CE).EQ.0) A(RE,CE)=CADRE+(-1.)+VPERT/2. 2390 + 2400 | 17 CONTINUE 2410 1 7 CONTINUE 2420 1 6 CONTINUE 2430 1 5 CONTINUE 2440 ! 3 CONTINUE 2450 | 2 CONTINUE 2460 | 1 CONTINUE 2470 1 EIGVLE=ORDER IF (ORDER.GT.200) EIGVLE=200 2480 ! C 2490 1 C 2500 ! C-------------------2510 / C Subroutine: TDIAG 2520 ! C 2536 ! C This subroutine reduces a real symmetric matrix to a symmetric 2540 ! C tridiagonal matrix using and accumulating orthogonal similarity 2550 ! C transformations. 2560 | C The subroutine is from EISPACK, a collection of FORTRAN subroutines 2570 ! C for eigenanalysis of matrices. 2580 ! C The program is called TRED2 in EISPACK. 2590 | C Authors: Martin, Reinsch, and Wilkinson, Num. Math. 11, 181-195(1968). 2600 / С-----2610 ! C 2620 1 CALL TDIAG(ROWTOT, ORDER) 2630 ! C 2640 I C-----2650 | C Subroutine: DIAGIT 2660 ! C 2670 ! C This subroutine finds the eigenvalues of a symmetric tridiagonal 2580 | C matrix by the implicit QL method. 2690 ! C This subroutine is from EISPACK. 2700 ! C The subroutine is named IMTQL1 in EISPACK. 2710 ! C Authors: Martin and Wilkinson, Num. Math. 12, 377-383(1968). Modified 2720 ! C by DuBrille, Num. Math. 15, 450(1970). 2730 ! C------2740 ! C 2750 ! CALL DIAGIT(ORDER, IERR) 2760 ! C 2770 I C------2780 1 C Prints the results of the diagonalization. 2790 ! C-----2800 ! C 2810 1 PRINT 90 ORDER , VPERT , ROTCA 2820 + 90 FORMAT(///,1X, MATRIX ORDER = ',15,10X,' POTENTIAL = ',F10.5. 2830 1 + 10X, 'ROTATIONAL CONSTANT = ',F10.4./) 2840 1 PRINT 91, EIGVLE 2850 | 91 FORMAT(1X, 'FIRST ', I5, ' EIGENVALUES:',/) 2860 1 PRINT 80, (D(I), I=1, EIGVLE) 2870 1 80 FORMAT (10(1X,F10.4)) 2880 1 PRINT 96 EIGULE 2890 · 95 FORMAT (///,)X,'FIRST ',IS,' EIGENVALUES RELATIVE TO Z.P.:'./) 2900 DIFF=D(1) 2910 1 PRINT 97.(D(I)-DIFF.I=1.EIGVLE) 2920 + 97 FORMAT (10(1X,F10.4))

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STOP 2330 1 2940 1 END 2950 ! C 2960 · C---2970 / C Function: THINT 2980 / C 2390 | C Part of the symmetric top wavefunction that depends on theta. 3000 + C Also contains the theta component of the torsional potential. 3010 + C This function is integrated between the range of 0 and pi by STIT. 3020 | C------3030 · C 3040 1 FUNCTION THINT(X) 3050 COMMON/BSF/RJ,RK,RM,CJ,CK,CM,ZA,ZB,ZC,ZD,ZE,ZF,ZG,ZH,FLG1 3060 | INTEGER FIRSTR, FINR, S.Z. FIRSTC, FINC 3070 ! C 3080 ! C-----3090 / C Theta component of the symmetric top wavefunction. 3100 ! C--3110 ! C 3120 1 FR=0 3130 ! FIRSTR=0 IF(-1+(RK+RM).GT.FIRSTR) FIRSTR=-1+(RK+RM) 3140 1 3150 FINR=RJ-RM 3160 IF(RJ-RK.LT.FINR) FINR=RJ-RK 3170 1 DO 1 S=FIRSTR\_FINR 3180 1 SI=RJ-RM-S 3190 ! SZ=RM+RK+Z+S 3200 1 53=2+RJ-RM-RK-2+5 3210 ! F1=-1 3220 ! IF(INT(S1/2).EQ.S1/2) F1=1 3230 1 F2=COS(.5+X) IF(ABS(F2).LT.1.E-9) F2=0 3240 ! 3250 1 IF(F2.EQ.0.AND.S2.EQ.0.) F2=1 IF(F2.EQ.0.AND.S2.EQ.0.) 60 TO 2 3260 ! 3270 ! F2=F2++52 3280 ! 2 F3=SIN(.5+X) 3290 1 IF(ABS(F3).LT.1.E-9) F3=0 3300 IF(F3.EQ.0.AND.S3.EQ.0) F3=1 3310 1 IF(F3.EQ.0.AND.S3.EQ.0) 60 TO 3 3320 F3=F3++S3 3330 1 3 ZZ=F1+F2+F3 3340 1 ZZR=ZA 3350 1 DO 4 Z=5,2,-1 3360 1 ZZR=ZZR/Z 3370 4 CONTINUE 3380 ! ZZR=ZZR+ZB 3390 ! 00 5 Z=RJ-RM-5,2,-1 3400 1 ZZR=ZZR/Z 3410 5 CONTINUE 3420 ZZR=ZZR+ZC DO 6 Z=RM+RK+S,2,-1 3430 | 3440 1 ZZR=ZZR/Z 3450 5 CONTINUE 3460 ZZR=ZZR+ZD 3470 1 DO 7 Z=RJ-RK-S,2,-1 3480 1 ZZR=ZZR/Z 3490 1 7 CONTINUE 3500 + FR=FR+ZZR+ZZ 3510 1 CONTINUE 3520 1 FC=0

338 FIRSTC=0 3530 ! IF(-1+(CK+CM.GT.FIRSTC) FIRSTC=-1+(CK+CM) 3540 3550 1 FINC=CJ-CM IF(CJ-CK.LT.FINC) FINC=CJ-CK 3550 -DO 8 S=FIRSTC,FINC 3570 / S1=CJ-CM-S 3580 1 3590 1 S2=CM+CK+2+S 3600 / S3=2+CJ-CM-CK-2+5 3610 -F1=-1 3620 1 IF(INT(S1/2).EQ.S1/2) F1=1 3630 1 F2=COS(.5+X) IF(ABS(F2).LT.1.E-9) F2=0 3640 ! 3650 1 IF(F2.EQ.0.AND.S2.EQ.0) FZ=1 IF(F2.E0.0.AND.52.E0.0) GO TO 9 3660 3670 ! F2=F2++S2 3680 1 9 F3=SIN(.5+X) IF(A85(F3).LT.1.E-9) F3=0 3690 IF(F3.EQ.0.AND.S3.EQ.0) F3=1 3700 ! 3710 / IF(F3.EQ.0.AND.53.EQ.0) 60 TO 10 F3=F3++S3 3720 1 3730 10 ZZ=F1+F2+F3 3740 ZZR=ZE 3750 1 DO 11 Z=S.2.-1 3760 ZZR=ZZR/Z 3770 ! 11 CONTINUE 3780 1 ZZR=ZZR+ZF 3790 1 DO 12 Z=CJ-CM-S.2.-1 3800 1 ZZR=ZZR/Z 3810 1 12 CONTINUE 3820 1 ZZR=ZZR+ZG 3830 1 DO 13 Z=CM+CK+S.2.-1 3810 1 ZZR=ZZR/Z 3850 13 CONTINUE 3369 ! ZZR=ZZR+ZH 3870 1 DO 14 Z=CJ-CK-S.2.-1 3880 | ZZR=ZZR/Z 3890 1 14 CONTINUE 3900 FC=FC+ZZR+ZZ 3910 1 8 CONTINUE 3920 ! C 3930 I C------3940 ! C Theta component of the torsional potential. 3950 · C-----3960 I C IF(FLG1.EQ.0) THINT=FR+FC+SIN(X)+COS(2+X) 3970 / IF(FL61.EQ.1) THINT=FR+FC+SIN(X)+SIN(2+X) 3980 1 IF(FLG1.EQ.2) THINT=FR+FC+SIN(X)+SIN(2+X) 3990 1 IF(FL61.EQ.3) THINT=-1.+FR+FC+SIN(X)+COS(2+X) 4000 1 4010 -RETURN 4020 1 END 4030 1 C 4040 ! C-----4050 | C Function: XINT 4060 . 0 4070 | C Part of the symmetric top wavefunction that depends on chi. 4080 ( C Also contains the chi component of the torsional potential. 4090 + C This function is integrated between the range 0 and 2+pi by STIT. 4100 | C-----4110 I C 4120 1 FUNCTION XINT(X)

COMMMON/BSF/RJ .RK .RM ,CJ ,CK ,CM ,ZA ,ZB ,ZC ,ZD ,ZE ,ZF ,ZG ,ZH ,FLG1 4130 1 4140 1 C 4150 / C-----4150 1 C Chi component of the symmetric top wavefunction. 4170 / C-----4180 C XINT=COS(RK+X)+COS(CK+X)+SIN(RK+X)+SIN(CK+X) 4190 1 4200 C 4210 · C-----4220 + C Chi component of the torsional potential. 4230 ! C-----4240 C IF (FLG1.EQ.0) XINT=XINT+COS(2+X) 4250 IF (FLG1.EQ.1) XINT=XINT+SIN(2+X) 4260 ! IF (FLG1.EQ.2) XINT=XINT+COS(2+X) 4270 1 IF (FLG1.EQ.3) XINT=XINT+SIN(2+X) 4280 ! RETURN 4290 1 4300 1 END 4310 ! C 4320 ! C-----4330 ! C Function: PHINT 4340 ! C 4350 ! C Part of the symmetric top wavefunction that depends on phi. 4360 ! C Also contains the phi component of the torsional potential. 4370 ! C This function is integrated between the range of 0 and 2+pi by STIT. 4380 | C------4390 ! C 4400 FUNCTION PHINT(X) 4410 ! COMMON/B5F/RJ ,RK ,RM ,CJ ,CK ,CM ,ZA ,ZB ,ZC ,ZD ,ZE ,ZF ,ZG ,ZH ,FLG1 4420 I C 4430 I C------4440 ! C Phi component of symmetric top wavefunction. 4460 ! C PHINT=COS(RM+X)+COS(CM+X)+SIN(RM+X)+SIN(CM+X) 4470 | 4480 ! C 4500 / C Phi component of the torsional potential. 4510 / C-----4520 I C IF (FLG1.EQ.0) PHINT-PHINT+COS(X) 4530 ( 4540 1 IF (FLG1.EQ.1) PHINT=PHINT+COS(X) 4550 / IF (FLG1.EQ.2) PHINT=PHINT+SIN(X) 4560 1 IF (FL6).EQ.3) PHINT=PHINT+SIN(X) 4570 1 RETURN 4580 1 END 4590 + C 4510 / C Subroutines STIT, TDIAG, and DIAGIT are attached at this point 4620 <sup>|</sup> C in the program. 4630 1 0-----

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