CALCULATION OF THE EQUILIBRIUM CONFIGURATION AND INTERMOLECULAR FREQUENCIES OF WATER DIMERS AND HEXAGONAL ICE

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The equilibrium structure and vibrational frequencies of the water dimer and hexagonal ice have been calculated using the Hartree-Fock potential of Clementi and coworkers and the correction for dispersion interactions of Kolos and coworkers. This correction term is proven to improve substantially the calculated results in the solid. The results obtained for the dimer were compared to other semiempirical and ab initio calculations and converging trends of the different studies are pointed to the convergence of the convergence of the convergence of the popular behavior of the lattice frequencies as a function of the

1. Introduction

Recently there has been increased interest in the study of the interaction between water molecules. Research in this field has been carried out mainly in two directions: semiempirical studies in which an intermolecular potential function is assumed and ab

function in the semiempirical calculations are selected so that the calculated values of some observable quantities in the liquid or solid phase, reproduce the measured values. Among the semiempirical potentials frequently employed are the Ben-Naim-Stillinger potential (BNS) [1], the Shipman-Scheraga potential (SS) [2] and the modification by Stillinger and

analytical representation of these potentials makes

types of calculations [7] which make use of a closed form representation of the potential.

In the field of ab initio calculations several studies have been performed which show general agreement on the equilibrium structure of the water dimer [8-10]. Among these calculations, that of Popkie et al. [9] was carried out near the equilibrium configuration, close to the Hartreen Fock limit Marsayer Popkie et al.

to fit an analytical representation to this surface. In a more recent work by these authors [11] the ab initio calculations were extended to less constrained configurations of the dimer and the parameters of the analytical representation of the surface were changed accordingly. Ab initio calculations, however, neglect the correlation energy contribution to the potential surface. To correct

[12,13]. The expression proposed by Kolos and co-

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accurate quantum mechanical calculations makes them of particular interest.

The quality of such ab initio interaction potentials has been examined by using them to compute bulk features such as the cohesive energy of water dimers [11] and orientationally averaged properties such as the virial coefficient of steam [7] and pair correlation functions of liquid water [4]. We feel that a rather sensitive test of the quanty of the potentials regarding their anisotropy, the contribution of electron correlation, the importance of many body forces, etc. could

teraction centers. The parameters in eq. (1) (in angstroms and kilocalories per mole) are:

$$q_{\rm O} = 0, q_{\rm H} = 11.801, q_{\rm M} = -2q_{\rm H}$$
,

$$A_{OM} = A_{HM} = A_{MM} = 0$$
,

$$A_{OO} = 71533$$
, $A_{OH} = 4084.0$, $A_{HH} = 779.8$,

$$b_{OO} = 3.969$$
, $b_{OH} = 3.914$, $b_{HH} = 3.125$.

scattering). In this paper we report the results of a preliminary study on the binding energies, equilibrium structure and intermolecular frequencies of the water dimer and hexagonal ice (Ih) using the Hartree—Fock interaction potential (HF potential) of Popkie et al.

those yielded by various phenomenological potentials and the results for the solid are compared with available experimental data.

2. The potentials

The restrict fit to the United Fresh out food of

i=1 i=1

Here R_{ij} is the distance between point i in molecule

the hydrogen positions. A fourth interaction point (M)

molecular plane, at a distance of 0.2259 Å from the oxygen [11]. The O-H bond length was taken to be 0.957 Å and the HOH bond angle 105°. It should be stressed that no special meaning was attributed by

the interaction potential or to the position of the in-

$$V_{\alpha,\beta}^{(2)} = C_6/R_{\text{O}\cdots\text{O}}^6 - C_8/R_{\text{O}\cdots\text{O}}^8 + C_{10}/R_{\text{O}\cdots\text{O}}^{10}$$
, (2) with

 $C_6 = 922.78$, $C_8 = 17283.5$ and $C_{10} = 24119.7$. The potentials $V^{(1)}$ and $V^{(1)} + V^{(2)}$ are used here to

3. The water dimer

Six degrees of freedom are necessary to describe the conformation of the system composed of two rigid monomers. A configuration of minimum energy has to be found in this six dimensional space. For our

Cartesian apardinates of the N - & mainta which re

rigid geometry for the individual monomers we chose an artificially stiff intramolecular potential to yield the correct hand lengths and hand analyse for the between inter- and intramolecular requestors.

approximation should not appreciable affect the

Starting from an arbitrary initial configuration we reach a configuration of minimum energy using steepest

The final results were shown to be independent of the



Fig. 1. Intermalecular coordinates for the water dimer

Table 1
Calculated conformation and stabilization energy of the water dimer

	Potential				
	CPa)	HF	HFK	SSp)	ST2c)
R (A)	2.974	2.98	2.73	2.85	2.852
θ (°)	4.6	3.6	i 4	5.8	-1.0
φ(°)	119.0	124.8	103	100.2	128.2
Stabilization	5.6	4.89	6.48	5.76	6.839
energy (kcal/mole)					

a) From ref. [10]. Calculations carried out with the 6-31G* basis set.

initial configuration in the aslaulations we servind out

one minimum in either energy surface although several saddle point configurations (which exhibit imaginary frequencies in the dynamic calculations) have been found in both surfaces.

Our results with both potentials agree with other ab initio [10] and semiempirical [15,16] calculations which show that the minimum energy configuration is the one for which one water molecule and the bisector of the HOH and a of the coord malacula lie in one

The configuration of the dimension there has

of minimum energy for both potentials, and compare them with the results of the study carried out with ab initio methods by Curtiss and Pople (CP) [10] using the 6-31 G^* basis set and the results obtained with the semiempirical potentials SS [16] and ST2 [15]. It can be seen that all potentials predict, within 15°, a linear hydrogen bond ($\theta = 0$). It should be mentioned

that the stable dimer structure predicted by hath the

initio HF and HFK potentials differs markedly from the structure which would be stable for dipole and made the interestions [15]. Thus, accordingly the range polarization effects seem to contribute to determining the equilibrium conformation.

The O···O distance, which seems to be the most important conformational parameter, is longer for the HF potential $(R_{O···O} \approx 2.98 \text{ Å})$ than for the semi-

material (D - 2.72 8)

presented in figs. 2a and 2b. Basically, the correction term provides an extra attractive force between the oxygen atoms. This shortens the $O_1\cdots O_2$ distance, permitting better electrostatic interaction between the H(2) of the donor molecule and the M center of the acceptor molecule. This in turn allows a reduction in the ϕ angle from 124° to 103°, resulting in close re-

b) From ref. [16]. c) From ref. [15].

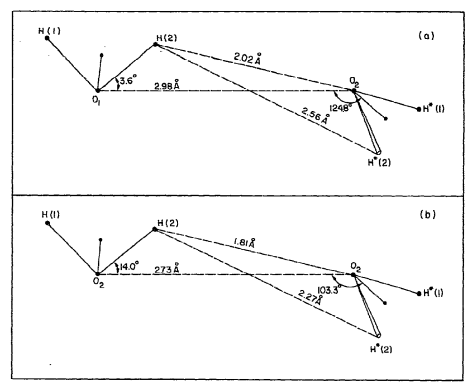


Fig. 2. Interatomic distances at the equilibrium conformation of the water dimer calculated with (a) the HF potential; (b) the HFK

Table 2					dimer is the creation of closer interatomic di-
					sms Aliszada tili ar illikulunasmar ni
A'	452 204 185	343.8 174.6 126.8	427.9 238.1 79.4	451 183 106	for all potentials the equilibrium geometry on has a C_s symmetry, the frequencies can be different case into three planar modes of symmetry.
A"	536 118	552.5 141.6	635.6 121.7	681 113	and 3 out-of-plane modes of symmetry, A".

a) From ref. [10]. Calculations carried out with the 4-31G

pulsive contacts between H(2)-O2, H(1)*--H(2) and $H(2)^* \cdots H(2)$ which are relieved by widening the θ angle from about 4° to 14°. Thus the overall effect of the correction term on the equilibrium geometry of the of the dimer lividied in netry, A', The analysis

istances

upper A mode has, basicany, the character of an hydrogen bond bend whereas the two lower A' modes The upper A" mode is clearly a hydrogen bond out-ofplane bend, a motion in which the H...O distance changes substantially. The second A" mode is a torsion of the two monomers around the hydrogen bond, and the third mode is a mixture of bending and the torsional king of displacement.

b) From ref. [16].

(a) The calculation by curriss and ropic [10] with

they recalculated the diagonal force field with the more extended 6-31G* basis set, they obtained smaller diagonal A' force constants. In an approximate calculation (using only diagonal terms) they carried out with these force constants, the hydrogen bond stretch frequency of 204 cm⁻¹ (column 2, table 2) was reduced to 170 cm⁻¹. This compares very well with the 174 cm⁻¹ value obtained in the HE calculation.

(b) When comparing the frequencies calculated with

cies increase, whereas the three low frequencies decrease. The character of the normal modes and their relative order remain unchanged by introducing the dispersion forces. The frequencies are rather sensitive to the inclusion of the correlation contribution. (Changes of up to 50% are seen.) The measurement

tive correction to the HF surface.

much closer than to the pure restricted HF calculations. This is the behavior one would expect from an empirical potential which implicitly includes the effect of the dispersion forces.

4. Hexagonal ice

method [17], in which one parametrizes the force constants to obtain agreement between calculated and experimental normal modes [18,19] or on phenomenology which deduce the elements

[20,21]. In this section we report the initiated and

the ab initio interaction potentials. The calculations

molecules. This assures that the calculated frequencies

by locating 4 oxygen atoms of a primitive unit cell in special positions $\pm (1/3, 2/3, z; 2/3, 1/3, 1/2 + z)$ of the P63/ $_{mmc}$ (D $_{6h}^4$) hexagonal space group [24]. According to the experimental results z = 1/16 and the unit cell dimensions a and c are such that all the O···O distances are equal and the sublattice formed by the oxygen atoms is tetrahedral [25]. The hydrogen atoms were located 0.957 Å from the oxygens and allowed to depart from the O···O line so as to form an HOH angle of 105° . (If the hydrogens would be on the O···O line they would also form a tetrahedral

is given by

$$V_{\text{u.cell}} = \frac{1}{2} \sum_{\alpha=1}^{4} \sum_{\beta=\alpha}^{M} V_{\alpha,\beta}$$
 (3)

Here $V_{\alpha,\beta}$ is the interaction energy between monomer

potential (eq. (1)) or as the sum of eqs. (1) and (2)

action with the 124 unit cells surrounding the basic unit cell and hence M=496. The energy per molecule, which is the quentity to be related to the sublimation energy of ice, is one-fourth of $V_{\rm u.cell}$. For hexagonal ice we carried out two parallel studies. In a first approximation we neglected zero point energy effects and looked for a crystal configuration which minimizes the static energy given by eq. (3). The energy was minimized with second to the continuous of the continuous of

the unit cell parameters a, b, c, α, β and γ . No constraints were imposed on the minimization path except to assume that the crystal is built by translating the basic unit cell [26]. In other words, the four monomer in

unit cens rengins and angles were anowed to change

zation was carried out in the 3N cartesian coordinates

tion

Table 3 Conformation of ice Ih at which the lattice energy is a mimi-

	Potential	
	HF	HFK
a (A)	4.857	4.530
b (A)	4.812	4.541
c (A)	7.852	7.403
α	89.5	89.7
β	88.9	90.2
Υ	117.9	1199

Lattice energy

Frequencies (cm-1):

		002	ı
	692	789	
	660	762	
	653	679	
	569	635	
	544	556	
	465	445	
	423	410	
•	382	325	
	375	320	
	341	281	
	339	274	
	221	221	
Translations	234	271	
	217	269	

38	37
31	28

space of the N = 16 centers of interaction of the four monomers in the unit cell. Results for the calculated unit cell parameters, lattice energy and k = 0 lattice frequencies for the configuration which minimizes the static energy are given in table 3. It can be seen that with

pendent units no single value of the calculated O...O bond length, or HO...O hydrogen bond angle can be given. Dut the differences in the O... O length and the angle of the hydrogen bond formed by the different monomers are definitely small. From table 3 it can be seen that the HFK potential predicts 0... 0 distances 0.2 Å smaller than the HF calculations and that in both cases the bond is almost linear

It is difficult to describe the exact character of the normal modes, since symmetry rules are not operative here. But even though the eigenvectors show that there

groups: higher frequencies, at which the molecules of the frequencies and lowering others, caused by the correlation notential in the dimer can be noticed clearly here in the librational modes. The higher librational frequencies calculated with the HFK potential are about 100 cm⁻¹ higher than the high frequencies yielded by the HF potential. On the other hand the lower librational modes are about 60 cm⁻¹ lower with the HFK potential.

In the second part of our study we estimated the influence of the zero point energy in determining the equilibrium configuration of the crystal, by repeating our calculations at a series of preestablished molar volumes. In every case the starting conformation of the critetal was taken as the tetrahedral arrangement of th

this case the unit cell axes were kept at fixed values which satisfy the relation c/a = 1.628 with angles $\alpha = \beta = 90^{\circ}$, $\gamma = 120^{\circ}$. We omitted a complete calculation of the density of states to evaluate the zero point energy and assumed in this preliminary study that the average of each one of the optical modes over the whole Brillouin zone equals its k = 0 value, and that the three acoustical modes can be estimated by a Debye model

the unit cell parameters are substantially shortened when the correlation energy term is included in the calculation.

mentioned that such an approximation was shown to be very good in the case of N_2 crystals [22,23]. The results at the crystal configuration which minimizes the

Table 4
Conformation of ice Ih which minimizes the total energy of the crystal. Frequencies calculated at this conformation

Table 5
Comparison between calculated and experimental data for ice

	Potential			Coloulated		Evporimental	
	m	nrk		ar	nrx		
a = b (A)	4.854	4.602	a (A)	4.854	4.602	4.4968 a)	
c (A)	7.927	7.516	c (A)	7.927	7.516	7.3198 a)	
Energy per molec	6.20	8.53	Sublimation energy	7.39 b)	9.72 b)	11.31°)	
			(cm ')				
Librations	753	835	Lower libration	339	267	~525 d)	
	715	778	(cm ⁻¹)	307	207	020	
	704 616	762 701	Librational bandwidth	414	568	~500 d)	
	592	624	(cm ⁻¹)	414	300	~300 07	
	549	570	Higher translation				
	444	452	(cm ⁻¹)	237	260		
	<u> </u>	HOW	i .		Y	_	
		V	/our \				
	375	306	Translational				
	344	283	bandwidth	203	232		
	339	267	(cm ⁻¹)				
Translations	237	260	a) From the low temperature results of ref. [27]. b) These values were obtained by adding to the results in				
	220	259					
	202	253					
	201	246	table 4, +1.19 kcal/mole (ref. [28]) to account for the				
	190-	าษ	intramolecular zer	o point ene	rgy.		
	131	100					

data. It can be seen from the table that the dispersion

Alexander of the second second

volume in such a way that their average is a function which depends only weakly on the volume. Another interesting effect we noticed is that the gap between transfer and libraries all modes (defined and back)

the reduce et which the calculations are carried out

39 A³/molec for the HF potential and 38 A³/molec for the HFK potential. This prediction could be tested by infrared and Raman pressure-dependent experiments. In table 5 we compare some values calculated with

dimensions larger and sublimation energy smaller than the experimental values. This could be due to the fact

ration anarmy of water trimage and totromage [11] A

sublimation energy within 5% of the experimental value and would cause a shortening of the unit cell parameters, thus bringing them to closer agreement with the experimental results.

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