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Citation: The Journal of Chemical Physics 82, 736 (1985); doi: 10.1063/1.448497

View online: http://dx.doi.org/10.1063/1.448497

View Table of Contents: http://aip.scitation.org/toc/jcp/82/2

Published by the American Institute of Physics



Molecular jet study of the solvation of toluene by methane, ethane, and propane^{a)}

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(Recieved 6 July 1984; accepted 1 October 1984)

Two color time of flight mass spectroscopy studies of toluene solvated by methane, ethane, and propane in a supersonic molecular jet have been carried out. This work is quite similar to the studies in the preceding paper on benzene. The conclusions and finding in the benzene investigation are strengthened and elaborated. The comparison of calculations and experiments has yielded information on binding energy, geometry, and spectral shift. A strong correlation is found between observed cluster transition intensity and cluster nucleation processes and a tentative nucleation scheme for the molecular jet formation of solute-solvent clusters is presented

I. INTRODUCTION

The spectroscopic study of van der Waals (vdW) clusters in a molecular jet supersonic expansion has greatly increased our understanding of these species. Additionally, clusters composed of an aromatic molecule (solute) surrounded by small hydrocarbons (solvent) have been discussed as models for condensed phase systems. Initial studies of these models have been focused on relatively small clusters with few solvent molecules:^{2,3} geometries, energetics, and energy dynamics have all been explored in these vdW systems.

In a companion paper (hereafter referred to as I), information on benzene solvated with methane, ethane, and propane is presented.4 Based on this and previous work from our laboratory, the nature of solvation and the solvation process has been explored. The solvent shift in the benzene clusters is understood to arise primarily from the interaction of the solvent with the π cloud of the aromatic ring: hence, the strong dependence of spectral shift on geometry. Identification of cluster geometry and energy levels has also led to understanding of the nucleation processes important in the formation of van der Waals clusters. Homogeneous nucleation can occur through the vibrational predissociation (VP) of a solvent dimer as it forms a complex with the aromatic solute. Inhomogeneous nucleation (the addition of two or more solvent molecules at a time to a solute) can be the dominant process in the beam. The relative amount of homogeneous and inhomogeneous nucleation is proportional to the ratio of solvent-solvent binding energy to solvent-solute binding energy.

The present paper reports the results of a study of the vdW clusters of toluene with methane, ethane, and propane. Due to the reduction of the solute symmetry compared to benzene, the geometry and energetics of toluene clusters are different from those of benzene

Both large $[Tol(S)_x, x > 3]$ and small (x < 3) clusters have been studied for the toluene-solvent systems. The changes in the spectra of these clusters as larger and larger clusters are probed, indicate possible relationships between the clusters and condensed phase systems. Comparisons of the large cluster spectra with the spectra of cryogenic solutions⁶ suggest how well and to what extent the jet generated clusters model condensed phase systems.

Computer modeling of toluene-hydrocarbon clusters is extremely helpful in supplementing the small cluster spectroscopic data in order to determine cluster geometry and binding energy. A number of local minima are found on the potential surface for toluene interacting with these hydrocarbon solvents. Only the lowest energy computer generated geometries which are consistent with the observed spectra are taken as acceptable solute-solvent configurations. If the computer generated potential energy surface has a number of minima with relatively low energy, that geometry which best agrees with the toluene and benzene spectra is accepted as the appropriate one.

The toluene cluster data will be discussed in two parts. The data for small clusters will be analyzed first and will be compared to the benzene small cluster data. The modeling of condensed phase systems by large clusters will be dealt with at the end of Sec. IV.

II. EXPERIMENTAL PROCEDURES

The apparatus and techniques employed in this work are similar to those described in I. Mass selective absorption (two-color MS) is the only spectroscopic technique used in this study. The optimal backing conditions for each nozzle are found experimentally and are reported in the figure captions.

clusters.⁵ Comparison of the benzene and toluene results for similar beam conditions and the same solvents strengthens and generalizes the conclusions reached previously.

Both large $[Tol(S)_x, x > 3]$ and small (x < 3) clusters

a) Supported in part by grants from ONR and ARO-D.

Three different nozzles are used in these experiments. The pulsed nozzle with a 500 μ m orifice produced a factor of ~15 larger intensity than the 50 μ m continuous (cw) nozzle which in turn produces signals three to four times more intense than the 25 μ m cw nozzle. Greater enhancements in the signals were expected for the larger diameter nozzels⁷; several possible loss mechanisms may contribute to this reduced signal improvement. In general, however, direct comparison between signals from the various nozzles is difficult because the optimum backing conditions for clustering are different for each nozzle.

Lowering the backing pressure and raising the pulsed nozzle temperature reduces the cooling in the expansion and produces hot clusters which evidence spectral hot bands. Although the (calculated) translational temperature of the expansion at the most extreme warming conditions $(P_0 = 10 \text{ psi}, T_0 = 70 \,^{\circ}\text{C})$ is still quite low ($\sim 1 \, \text{K}$), the vibrational temperature of the clusters is $\sim 50 \, \text{to} \, 100 \, \text{K}$, based on the observed spectral intensities.

Calculations of the minimum energy configurations of toluene clusters are performed as described in I. The coefficients for the interactions between the methyl group of toluene and the solvents are taken to be the same as those between solvents. The coefficients are reproduced in Table I for convenience.

Minimization of the energy of a given cluster is accomplished by moving the ligands in a stepwise fashion, calculating the energy at each step, and comparing the two most recent values of the energy. The procedure will terminate each time a local minimum in the energy has been found. To ensure that all minima are found and, in particular, that the lowest minimum is located, all reasonable starting geometries must be employed. Low energy and agreement with the spectroscopic data are the two criteria used to settle on "correct" configurations for the clusters.

TABLE I. Parameters for the energy expression in the computer modeling of the vdW clusters of toluene.

$E_{ij} = -A_{ij}/R_{ij}^{6} + B_{ij} \exp[-C_{ij}R_{ij}]$ $A\left(\frac{\text{cm}^{-1} \mathring{A}}{\text{mol}}\right)^{6} B\left(\frac{\text{cm}^{-1}}{\text{mol}}\right) C(\mathring{A}^{-1})$						
Aliphatic-aliphati	Ca.					
C-C	131 096	8 890 353	3.421			
C-H	47 830	7 562 708	3.94			
H-H	15 028	6 390 528	4.643			
Aliphatic-aromati	c ^b					
C-C	156 654	15 219 240	3.5105			
C-H	48 231	4 990 321	3.805			
H–H	13 758	2 990 682	4.1915			
Aromatic-aromati	ic ^c					
C-C	187 196.5	26 053 554	3.6			
C-H	48 636.1	3 292 908.9	3.67			
Н-Н	12 596.4	1 399 600	3.74			

From the methane-methane parameters of Ref. 11 in paper I.

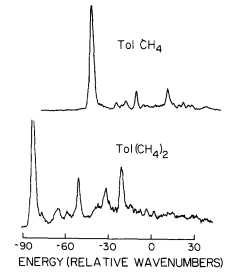


FIG. 1. Two-color TOFMS of $Tol(CH_4)_1$ (upper trace) and $Tol(CH_4)_2$ in the 0^0_0 region. The energy scale is relative to the $Tol~0^0_0~(37~477.5~cm^{-1})$. Assignments are given in Table II. This figure is reproduced from Ref. 5.

III. RESULTS

A. Toluene-methane

The Tol(CH₄)_{1,2} absorption spectra have been published previously and are reproduced here in Fig. 1 and Table II for convenience. These spectra exhibit sharp origins and features due to van der Waals (vdW) stretching and bending modes. The spectral shift for the Tol(CH₄)₂ 0_0^0 is twice that of the Tol(CH₄)₁ 0_0^0 . The additive shift indicates addition of the two methane molecules to equivalent sites on the toluene. The stretching mode for Tol(CH₄)₁ has a considerable anharmonicity as pointed out previously.⁵ Tol(CH₄)₂ evidences a similar progression in the stretching mode. The Tol(CH₄)₂ peak at -20 cm^{-1} is probably associated with a second configuration of Tol(CH₄)₂.

Large clusters of toluene with methane produce broad spectra which become more diffuse and structureless as the clusters increase in size (see Fig. 2). The red shift of the most prominent feature in the $Tol(CH_4)_3$ spectrum is nonadditive; this feature is actually less red shifted than the $Tol(CH_4)_2$ 0_0^0 . Nonetheless, intensity is still evident in the $Tol(CH_4)_3$ spectrum far to the red of the $Tol(CH_4)_2$ 0_0^0 . The absorption spectra of clusters as large as $Tol(CH_4)_{10}$ have been observed. The spectra become very broad and unstructured with a maximum about -50 cm⁻¹ from the Tol 0_0^0 .

The binding energy of these clusters can be estimated by observing the presence or absence of higher vibronic bands in the 2-color MS spectra. If the first photon induces a transition to a level which results in vibrational predissociation of the cluster, the transition will not be observed in the 2-color MS spectrum. For $Tol(CH_4)_{1,2}$ the $6b_0^1$ transition is observed but the 1_0^1 transition is not. This sets the limits on the binding energy at $534 \le D_0 \le 740 \text{ cm}^{-1}$.

The binding energy is the same within these limits for $Tol(CH_4)_{1,2}$. The binding energy for larger clusters is

^b The parameters are found by combining the benzene-benzene and methane-methane parameters as follows: $A_{\rm Bm} = (A_{\rm BB}A_{\rm mm})^{1/2}$, $B_{\rm Bm} = (B_{\rm BB}B_{\rm mm})^{1/2}$, $C_{\rm Bm} = \frac{1}{2}(C_{\rm BB} + C_{\rm mm})$.

TABLE II. Two-color TOFMS of $Tol(CH_4)$ and $Tol(CH_4)_2$ in the 0_0^0 region. Only the prominent peaks are tabulated and tentative assignments are given (see Fig. 1).

Species	Energy (vac cm ⁻¹)	Energy relative to Tol 00	Energy relative to cluster 00	Assignment ^a
Tol(CH ₄)				
	37 434.4	-43	0	00
	37 451.8		17	A_0^1
	37 458.7		24	B_0^1
	37 466.2		32	V_0^1
	37 488.5		54	V_0^2
Tol(CH ₄) ₂ I				
	37 394.1	-83	0	00
	37 412.6		18	A_0^1
	37 418.5		24	B_0^1
	37 426.7		32	V_0^1
	37 440.5		46	$V_0^1A_0^1$
	37 446.2		52	V_0^2
II				
	37 456.9	-21	0	08

Tentative assignments are as follows: V stands for the Tol-CH₄ stretch and A and B are different vdW bends. I and II represent different configurations.

difficult to estimate since the spectra of large clusters are less well defined and weak.

The computer modeling of $Tol(CH_4)_1$ shows a minimum energy configuration in which the methane is shifted away from the center of the aromatic ring toward the methyl group (see Fig. 3 and Table III). The methane should interact strongly with the aromatic π cloud of toluene in this predicted geometry. Only one stable configuration is calculated for $Tol(CH_4)_1$: This agrees with the observation of one origin in the spectrum. The calculated binding energy of 667 cm⁻¹ is consistent with the limits obtained spectroscopically.

Two acceptable geometries are found in the computer modeling of Tol(CH₄)₂. The configuration labeled I in Fig. 3 features a methane on either side of the aromatic

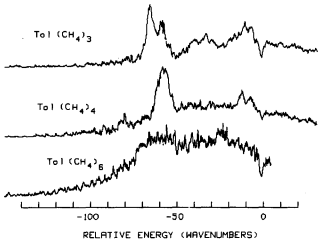
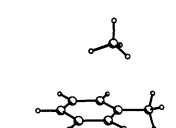


FIG. 2. Two-color TOFMS of toluene-methane large clusters in the 0_0^0 region. The energy scale is relative to the Tol 0_0^0 (37 477.5 cm⁻¹). The concentration of methane in the beam is 1% for Tol(CH₄)₃ and 2.5% for Tol(CH₄)₄ and Tol(CH₄)₆. The pulsed nozzle is cooled to 4 °C and $P_0 = 100$ psia.

ring, forming an isotropic cluster. The second configuration predicted by the computer modeling (labeled II in Fig. 3) is anisotropic with both methane molecules on the same side of the ring. In the isotropic geometry both of the solvent molecules occupy equivalent sites similar to the site occupied by methane in Tol(CH₄)₁. Conse-



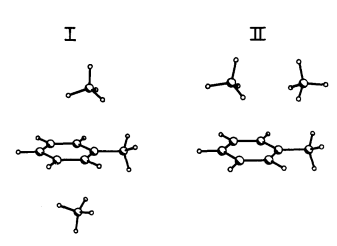


FIG. 3. Minimum energy configurations of Tol(CH₄)₁ and Tol(CH₄)₂. Atomic positions are given in Table III. Note that both isotropic (I) and anisotropic (II) clusters of Tol(CH₄)₂ are predicted in the computer modeling.

TABLE III. The atomic positions for the minimum energy configurations of $Tol(CH_4)_1$ and $Tol(CH_4)_2$ (see Fig. 3). The origin is at the center of the aromatic ring of toluene. The +X axis passes through the methyl group, the Y axis is in the plane of the ring and the Z axis is out of the plane of the ring.

Atom	X (Å)	Y (Å)	Z (Å)	Energy (cm ⁻¹)
Toluene				
C1	2.925	0	0	
C2	-1.395	0	0	
C3	-0.6975	1.208	Ō	
C4	0.6975	1.208	ŏ	
C5	1.395	0	ŏ	
C6			0	
	0.6975	-1.208		
C 7	-0.6975	-1.208	0	
HI	3.29	0	-1.038	
H2	3.29	0.8985	0.518	
Н3	3.29	-0.8985	0.518	
H4	-2.479	0	0	
H5	-1.24	2.147	0	
H6	1.24	2.147	0	
H7	1.24	-2.147	0	
Н8	-1.24	-2.147	Ō	
Tol(CH ₄) ₁				-667
Cl	0.921	0	3.384	007
H1	-0.104	0	2.985	
H2	1.451	-0.8985	3.036	
H3	1.451	-0.8985	3.036	
H4	0.884	0	4.483	
Tol(CH ₄) ₂ C	onfiguration I			-1348
Ligand 1				
C1	0.922	0	3.381	
HI	-0.103	Ö	2.982	
H2	1.452	-0.8985	3.033	
H3	1.452	0.8985	3.033	
H4	0.886	0.0703	4.48	
Ligand 2				
C1	0.767	0	-3.391	
HI	-0.255	Ö	-2.983	
H2	1.300	-0.8985	-3.048	
H3	1.300	0.8985	-3.048	
H4	0.721	0.0703	-4.49	
Configuratio	on II			-1360
Ligand 1				
-	2.372	0.003	3.433	
Cl		0.003		
H1	1.977	-0.8968	2.938	
H2	3.470	0.005	3.371	
H3	1.973	0.900	2.938	
H4	2.066	0.003	4.490	
Ligand 2			_	
C1	-1.089	-0.003	3.454	
H 1	-2.172	-0.005	3.260	
H2	-0.634	-0.901	3.011	
Н3	-0.638	0.896	3.011	
H4	-0.913	-0.003	4.540	
				

quently, this geometry should have the larger (additive) spectral shift in the Tol(CH₄)₂ spectrum (Fig. 1).

B. Toluene-ethane

The small clusters of toluene and ethane $[Tol(C_2H_6)_{1,2}]$ exhibit sharp, assignable structure (see Fig.

4 and Table IV). $Tol(C_2H_6)_1$ shows a progression in the vdW stretch, as well as two intense bends. The $Tol(C_2H_6)_2$ spectrum clearly suggests that two geometries exist for this cluster. One geometry generates a spectral shift which is twice that found for $Tol(C_2H_6)_1$. The vdW vibrational features of this additive shift configuration consist of a long progression in the stretching mode. The second geometry is characterized by a small (-10 cm^{-1}) solvent shift and a short progression of doublets after the origin peak.

The absorption spectra of larger $Tol(C_2H_6)_x$ clusters become broad with very little well-defined structure. The maximum in a typical absorption spectrum occurs at about 30 cm⁻¹ to the low energy side of the Tol 0_0^0 , although significant measurable intensity extends to roughly -200 cm^{-1} .

The effect of warming the $Tol(C_2H_6)_x$ clusters is similar to that of warming the $Tol(CH_4)_x$ clusters. The hot bands produce a broad background somewhat to the low energy side of the cold bands. As judged by the decrease in the signal intensities, continued warming (P < 20 psi, $T_0 \sim 70$ °C) begins to destroy the vdW complexes.

Vibrational predissociation of $Tol(C_2H_6)_{1,2}$ clusters occurs at the 12^1 toluene excitation but not at 1^1 : the binding energy of these clusters therefore lies between 740 and 933 cm⁻¹. Both geometries of $Tol(C_2H_6)_2$ seem to have the same binding energy within this experimentally defined range.

The computer modeling of $Tol(C_2H_6)_1$ shows two minima in the potential surface above toluene. Since only one geometry is obvious experimentally only the lower energy computer generated configuration is presented in Fig. 5 and Table V. The less stable geometry ($\sim 180 \text{ cm}^{-1}$ higher in energy) is probably not observed spectroscopically, although one of the peaks assigned to a vdW bending mode is possibly the origin of a second configu-

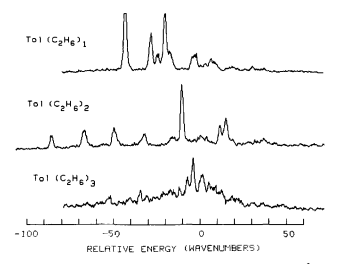
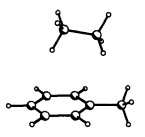


FIG. 4. Two-color TOFMS of toluene-ethane clusters in the 0_0^0 region. The energy scale is relative to the Tol 0_0^0 (37 477.5 cm⁻¹). The concentration of ethane in the beam is 0.5% for Tol(C_2H_6)₁ and Tol(C_2H_6)₂ and 2.0% for Tol(C_2H_6)₃. The backing conditions for the pulsed nozzle are P_0 = 100 psia and T_0 = 4 °C. Data for Tol(C_2H_6)₁ and Tol(C_2H_6)₂ are presented in Table IV.

TABLE IV. Two-color TOFMS of $Tol(C_2H_6)_1$ and $Tol(C_2H_6)_2$. Only the prominent peaks are tabulated and tentative assignments are given (see Fig. 4).

Species	Energy (vac cm ⁻¹)	Energy origin relative to Tol 00 (cm ⁻¹)	Energy relative to cluster 0% (cm ⁻¹)	Assignment ^a
Tol(C ₂ H ₆)				
. 2 0	37 435	-43	0	0%
	37 450		15	A_0^1
	37 454		19	B_0^1
	37 458		23	V_0^1
	37 460		25	A_0^2
	37 475		40	$V_0^2(V_0^1A_0^1)$
	37 485		50	$V_0^1 A_0^2$
	37 486		51	V_0^3
Tol(C ₂ H ₆) ₂ I				
	37 392	-86	0	00
	37 411		19	V_0^1
	47 428		36	V_0^2
	37 445		53	V_0^3
	37 461		69	V_0^4
	37 478		86	V_0^5
II				
	37 467	-11	0	00
	37 481		14	A_0^1
	37 489		22	B_0^1
	37 492		25	V_0^1

^a Tentative assignments are as follows: V stands for the fundamental vdW stretch A and B are different vdW bends. I and II represent different geometries. In $Tol(C_2H_6)_2$ the vdW modes may involve solvent-solvent motions as well as solvent-solute motions.



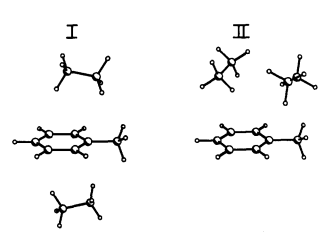


FIG. 5. Minimum energy configurations of $Tol(C_2H_6)_1$ and $Tol(C_2H_6)_2$. Atomic positions are given in Table V. $Tol(C_2H_6)_2$ also shows isotropic and anisotropic clusters.

ration. The binding energy predicted by the computer modeling is within the spectroscopically determined limits.

For $Tol(C_2H_6)_2$ several possible geometries are calculated. Since the spectrum clearly shows only two geometries, the most probable configurations must be chosen from this group. One of the geometries must be an isotropic configuration with an ethane on either side of the ring and the other configuration must involve both ethane molecules on the same side of the ring. Configurations consistent with the spectrum are shown in Fig. 5.

C. Toluene-propane

The spectra of the small clusters of toluene and propane $[Tol(C_3H_8)_{1,2,3}]$ also consist of well-resolved features (Fig. 6 and Table VI). The $Tol(C_3H_8)_1$ cluster spectrum strongly suggests two configurations exist for this cluster. The geometry with a relatively large spectral shift shows a progression in the vdW stretch as well as several vdW bends to the high energy side of its 0^0_0 transition. The second configuration gives rise to a small spectral shift of the origin: the origin is followed by a progression of doublets. This pattern is quite reminiscent of that found for the second (small shift) configuration of $Tol(C_2H_6)_2$.

The vdW cluster spectrum of $Tol(C_3H_8)_2$ has a complicated structure indicating three or four possible different configuration for this mass species. These ge-

TABLE V. The atomic positions for the minimum energy configurations of $Tol(C_2H_6)_1$ and $Tol(C_2H_6)_2$ (see Fig. 5). Axes and solute coordinates are the same as in Table III.

Atom	X (Å)	Y (Å)	Z (Å)	Energy (cm ⁻¹)
Tol(C ₂ H ₆) ₁				-967
Ligand				
Cı	0.166	-0.001	3.742	
C2	1.672	-0.001	3.478	
Hi	-0.374	0	2.783	
H2	-0.104	0.897	4.317	
H3	-0.105	-0.900	4.315	
H4	2.212	-0.002	4.437	
H5	1.943	0.898	2.905	
Н6	1.942	-0.899	2.903	
Tol(C ₂ H ₆) ₂ Co	onfiguration I			-1663
Ligand 1				
Cl	0.017	0.041	3.877	
C2	1.515	0.185	3.599	
HI	-0.524	-0.051	2.924	
H2	-0.340	0.929	4.418	
H3	-0.156	-0.857	4.487	
H4	2.056	0.277	4.553	
H5	1.688	1.083	2.989	
H6	1.574	-0.734	3.061	
Ligand 2				
CI	0.302	0	-3.845	
C2	1.802	0	-3.541	
Hi	-0.262	0.001	-2.900	
H2	0.048	0.898	-4.426	
H3	0.047	-0.899	-4.425	
H4	2.366	-0.001	-4.486	
H5	2.057	0.899	-2.961	
H6	2.056	-0.898	-2.960	
Configuration	II			-1834
Ligand 1				
C1	-0.628	1.561	3.933	
C2	-1.263	0.311	3.555	
H1	-1.355	2.254	4.496	
H2	-0.336	2.187	3.018	
H3	0.261	1.471	4.554	
H4	-1.554	-0.226	4.469	
H5	-2.151	0.490	2.933	
H6	-0.535	-0.292	2.991	
Ligand 2				
C1	2.294	-1.263	3.754	
C2	2.820	0.157	3.538	
H 1	2.139	-1.748	2.778	
H2	3.027	-1.838	4.338	
H3	1.341	-1.221	4.299	
H4	2.087	0.732	2.953	
H5	3.774	0.115	2.993	
H6	2.975	0.642	4.513	

ometries are generally distinguishable based on relative intensities, spectral shifts, qualitative vibrational analysis, and comparison with the $Tol(CH_4)_{1,2}$, $Tol(C_2H_6)_{1,2}$, and $Tol(C_3H_8)_1$ data.

The $Tol(C_3H_8)_3$ spectrum is actually less congested than the $Tol(C_3H_8)_2$ spectrum, presumably because some of the possible configurations are not observed in the beam. Two sets of peaks, corresponding to two different geometries of $Tol(C_3H_8)_3$, are present in the spectrum as

shown in Fig. 6. One set of peaks features a regular progression in an apparently single vdW mode and the other features a short progression of doublets.

Warming the toluene-propane clusters destroys the well-resolved structure in the $Tol(C_3H_8)_{2,3}$ spectra (Fig. 7). As in the other systems, hot bands of the $Tol(C_3H_8)_x$ clusters form an unstructured, broad background to the red of the cold spectrum.

The large clusters of toluene and propane produce spectra qualitatively similar to those of the other systems (Fig. 8). Although some structure exists in the $Tol(C_3H_8)_4$ spectrum, the $Tol(C_3H_8)_6$ spectrum is largely unstructured with an intensity maximum around -60 cm^{-1} relative to the $Tol(0_0^0)$.

The binding energies of the $Tol(C_3H_8)_{1,2,3}$ clusters are all greater than 933 cm⁻¹ as all three clusters show clear spectra in the 12_0^1 region. No toluene features above 12_0^1 are either intense enough or free enough from interferences to allow further studies; therefore, no upper limit to the binding energy can be quoted at this time.

Computer modeling predicts several stable geometries for $Tol(C_3H_8)_1$. As noted earlier, the $Tol(C_3H_8)_1$ spectrum indicates that two configurations with very different spectral shifts exist (see Fig. 6). Note the similarity in the spectral shifts of configuration II of $Tol(C_3H_8)_1$ and configuration II of $Tol(C_2H_6)_2$ (Fig. 4). This similarity in spectral shift indicates a similarity in the overlap of the solvent with the aromatic ring of toluene. Figure 9 and Table VII present the lowest energy geometries which should give rise to spectral shifts consistent with the $Tol(C_3H_8)_1$ spectrum base on the correspondence between solvent–aromatic ring overlap and spectral shift (I). Other less stable geometries may be producing some of the observed peaks in the spectrum, but further assignments cannot be made with the existing data.

Similar considerations are used to choose those geometries of $Tol(C_3H_8)_2$ which are consistent with the spectrum. Possible minimum energy geometries are pic-

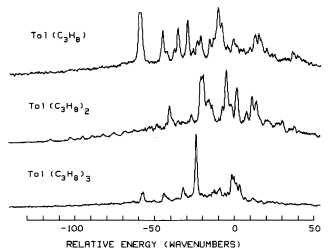


FIG. 6. Two-color TOFMS of toluene-propane small clusters in the 0_0^0 region. The energy scale is relative to the Tol 0_0^0 (37 477.5 cm⁻¹). The concentration of propane in the beam is 1% for all three spectra. The pulsed nozzle conditions are $P_0 = 100$ psia and $T_0 = 4$ °C. Data for these spectra are presented in Table VI.

TABLE VI. Two-color TOFMS of $Tol(C_3H_8)_1$, $Tol(C_3H_8)_2$ and $Tol(C_3H_8)_3$. Only the prominent peaks are tabulated and tentative assignments are given (see Fig. 6).

Species	Energy (cm ⁻¹)	Energy origin relative to Tol 00 (cm ⁻¹)	Energy relative to cluster 0_0^0 (cm ⁻¹)	Assignment
Tol(C ₃ H ₈) I				
	37 419	-59	0	O_0^0
	37 433		14	A_0^1
	37 435		17	B_0^1
	37 440		22	C_0^1
	37 442		24	D_0^1
	37 448		30	${V}^1_0$
	37 452		33	$B_0^1 A_0^1$
	37 454		36	$C_0^1 A_0^1$
	37 456		38	$D_0^1\!A_0^1$
	37 462		43	$V_0^1 A_0^1$
	37 461		51	V_0^2
	37 473		54	$V_0^1 D_0^1$
II				ο0
	37 467	-10	0	00
	37 477		10	A_0^1
	37 491		24	B_0^1
	37 493		26	V_0^1
Tol(C ₃ H ₈) ₂ I				
	37 361	-117	0	00
	37 374		13	V_0^1
	37 388		27	V_0^2
	37 402		41	V_0^3
II				00
	37 436	-41	0	00
	37 456		14	V_0^1
III				
	37 457	-20	0	O_0^0
	37 479		22	V_0^1

IV	27 472	•	^	00
	37 472	-5	0	B_0^1
	37 489		16	V_0^1
	37 492		20	V 0
Tol(C ₃ H ₈) ₃ I				
	37 421	-57	0	O_0^0
	37 434		13	$egin{array}{c} V_0^1 \ V_0^2 \end{array}$
	37 446		25	V_0^2
**				
II	27.454	24	0	00
	37 454	-24	0	A_0^1
	37 469		15	V_0^1
	37 477		23	r 0

^a A through D represent different vdW bends and V represents the fundamental vdW stretch. I through IV represent different geometries.

tured in Fig. 10 and the atomic positions of one geometry are tabulated in Table VIII. Computer modeling of $Tol(C_3H_8)_3$ was not done.

IV. DISCUSSION

The spectral shift of a cluster depends primarily on the polarizability of the solvent along the intermolecular bond and on the change in polarizability of the solute upon excitation. One would expect that the change in polarizability upon excitation is larger for toluene¹⁰ than benzene and thus, that the cluster shifts are larger for toluene than benzene clusters. However, most of the change in polarizability upon $(\pi \rightarrow \pi^*)$ excitation occurs in the aromatic π system; moving the solvent away from the π system must significantly reduce the cluster shift. Thus, two competing effects, increased polarizability change and reduced interaction with the aromatic π system, influence the relative spectral shifts of benzene and toluene clusters with small hydrocarbons. In the ensuing discussion we will treat large and small clusters separately and discuss shifts, geometries, and nucleation processes.

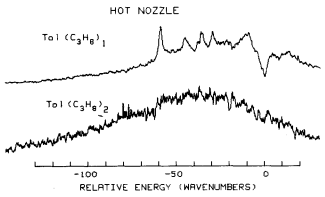


FIG. 7. Two-color TOFMS of $Tol(C_3H_8)_1$ and $Tol(C_3H_8)_2$ under hot nozzle conditions. The energy scale is relative to the Tol 0_0^0 (37 477.5 cm⁻¹). The concentration of propane is 1%, $P_0 = 40$ psia and $T_0 = 70$ °C. Note the dramatic increase in the background relative to the spectra in Fig. 6. This broad background is due to hot bands. The negative peak at 0 cm⁻¹ in the $Tol(C_3H_8)_1$ spectrum is due to detector overload at $Tol(0_0^0)$.

A. Small clusters

1. Methane

The computer modeling of the $Tol(CH_4)_1$ cluster shows that the methane is displaced 0.921 Å towards the methyl group of toluene (see Fig. 3 and Table III). This calculated geometry presents a rationale for the essential (unexpected) equivalence of the cluster spectral shifts for $Tol(CH_4)_1$ and $Ben(CH_4)_1$, even though toluene has a (presumed) larger change in polarizability upon excitation than does benzene. Thus, the abovementioned two competing effects tend to cancel one another for $Tol(CH_4)_1$.

These basic ideas also help to rationalize the similarities and differences in the Tol(CH₄)₂ and Ben(CH₄)₂ spectra. Spectra of the isotropic clusters (a solvent on each side of the aromatic ring) of Tol(CH₄)₂ and Ben(CH₄)₂ are nearly identical, as would be surmized based on the above arguments for the Tol(CH₄)₁ clusters. However, anisotropic clusters of toluene and benzene with two methanes exhibit significantly different shifts. Computer

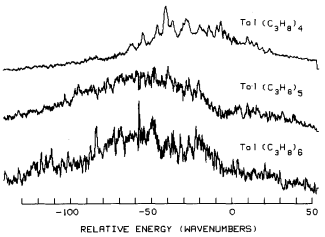


FIG. 8. Two-color TOFMS of toluene-propane large clusters. The energy scale is relative to the Tol 0_0^0 . The concentration of propane in the beam is 2% and the nozzle backing conditions are $P_0 = 100$ psia and $T_0 = 4$ °C.

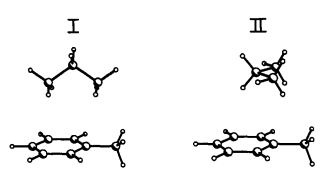


FIG. 9. Minimum energy configurations of Tol(C₃H₈)₁. Atomic positions are given in Table VII. In configuration II the propane interacts less with the aromatic cloud than in configuration I.

modeling of the anisotropic cluster of Ben(CH₄)₂ suggests that one methane moiety interacts strongly with the π system (I), thus yielding a spectral shift similar to that observed for Ben(CH₄)₁. On the other hand, computer modeling of the anisotropic cluster of Tol(CH₄)₂ shows that both methane molecules are considerably displaced from the center of the aromatic ring (1.09 and 2.37 Å). The observed small spectral shift for Tol(CH₄)₂ (-21 cm⁻¹) is thereby consistent with the previously determined (I) relationship between spectral shift and geometry.

The ratios of intensities of the isotropic to anisotropic cluster peaks are different for $Tol(CH_4)_2$ and $Ben(CH_4)_2$: $I_{iso}/I_{aniso}[Tol(CH_4)_2, \ 0_0^0] \sim 3$ (Fig. 1), whereas $I_{iso}/I_{aniso}[Ben(CH_4)_2, \ 6_0^1] \sim 1$. As pointed out in I, isotropic clusters can only be produced by homogeneous nucleation. Most homogeneously nucleated clusters are argued to

TABLE VII. The atomic positions for the minimum energy configurations of $Tol(C_3H_8)_1$ (see Fig. 9). The origin and the coordinates of the solute are the same as in Table III.

Atom	X (Å)	Y (Å)	Z (Å)	Energy (cm ⁻¹)
Configuration	on I			-1196
Ligand				
Cl	-0.438	0	3.418	
C2	0.819	0	4.29	
C3	2.060	0	3.396	
H1	-1.33	0	4.061	
H2	-0.360	0.774	2.970	
H3	-0.360	-0.774	2.970	
H4	0.825	0.898	4.925	
H5	0.825	-0.898	4.925	
H6	2.964	0	4.023	
H7	1.974	0.774	2.950	
Н8	1.974	-0.774	2.950	
Configurati	on II			-1128
Ligand				
Cl	1.521	1.248	3.617	
C2	0.638	100.0-	3.636	
C3	1.520	-1.250	3.616	
Hı	0.887	2.146	3.632	
H2	1.985	1.165	4.381	
H3	1.951	1.166	2.834	
H4	0.020	-0.001	4.424	
H5	-0.014	0	2.876	
H6	0.886	-2.148	3.629	
H 7	1.984	-1.169	4.380	
H8	1.950	-1.168	2.832	

TABLE VIII. The atomic positions for the minimum energy geometry of configuration IV of Tol(C₃H₈)₂ (see Fig. 10). The origin and coordinates of the solute are the same as in Table III.

Atom	<i>X</i> (Å)	Y (Å)	Z (Å)	Energy (cm ⁻¹)
Configuration	on IV	-		-2376
Ligand 1				
C1	1.992	1.278	3.661	
C2	0.954	0.155	3.643	
C3	1.667	-1.198	3.633	
H 1	1.480	2.251	3.668	
H2	2.420	1.131	4.435	
H3	2.429	1.147	2.888	
H4	0.320	0.230	4.414	
H5	0.329	0.246	2.866	
H6	0.920	-2.006	3.620	
H7	2.116	-1.183	4.410	
H8	2.125	-1.167	2.862	
Ligand 2				
C1	-2.720	0.898	3.577	
C2	-3.350	-0.490	3.439	
C3	-2.263	-1.560	3.585	
H1	-3.502	1.665	3.472	
H2	-2.248	0.987	4.566	
H3	-1.963	1.034	2.792	
H4	-3.823	-0.580	2.450	
H5	-4.108	-0.627	4.224	
H6	-2.716	-2.558	3.486	
H7	-1.505	-1.423	2.800	
H8	-1.790	-1.471	4.574	

form through the collision of a solvent dimer with the solute; the binding energy of the solute with one of the solvent molecules is dissipated, in part, by VP of the solvent dimer bond. Apparently, the larger solute-solvent binding energy of the toluene clusters (as predicted by the computer modeling) facilitates breaking of the solvent dimer bond and allows homogeneous nucleation to dominate the formation of Tol(CH₄)₂ clusters. Nucleation processes will be treated in more detail at the end of this section.

2. Ethane

Figure 5 depicts the computer calculated geometry for $Tol(C_2H_6)_1$: the ethane molecule is displaced toward the methyl group of toluene and inclined with respect to the plane of the ring. Assuming this geometry to be correct, it results in a spectral shift smaller by roughly 5 cm⁻¹ than that found for the comparable $Ben(C_2H_6)_1$ configuration. The methyl group of toluene, according to the potential calculations, draws the ethane solvent molecule away from the ring center, thereby reducing the toluene spectral shift relative to the comparable benzene cluster.

An isotropic $Tol(C_2H_6)_2$ cluster is clearly indicated in the observed spectrum based on an origin shift twice that of the $Tol(C_2H_6)_1$ origin (Fig. 4). The intensity of features due to isotropic clusters relative to those due to anisotropic clusters is much less than seen in the $Tol(CH_4)_2$ spectrum. The ratio is consistent with the trend report in I; homogeneous nucleation can be less probable with solvents having larger binding energies than with solvents

having smaller binding energies. However, more homogeneous nucleation is evident for the toluene-ethane system than for the benzene-ethane systems. This same trend was found for the solute-methane clusters and is related to the strength of the solvent-solute bond.

3. Propane

Arguments put forth above concerning spectral shift, solvent-solute geometry, binding energies, etc., can also be applied for the somewhat more complicated toluene-propane clusters. The spectral shifts of the two configurations determined by computer modeling of $Tol(C_3H_8)_1$ (Fig. 9) can thereby be qualitatively predicted. Configuration I, having a significant interaction of a terminal methyl group of propane with the aromatic π system of toluene, should have a spectral shift greater than that of $Tol(CH_4)_1$. On the other hand, configuration II (Fig. 9) would be expected to have a small spectral shift since the propane molecule is close to the methyl group of toluene. Origins with the expected shifts are indeed observed in the spectrum of $Tol(C_3H_8)_1$ as depicted in Fig. 6.

The $Tol(C_3H_8)_2$ spectrum (Fig. 6) has substantial intensity assigned to isotropic clusters (see origin peaks labeled I, II, III in Table VI). This observation apparently does not fit the trend established above if the computer predicted values of the solvent dimer and solute-solvent

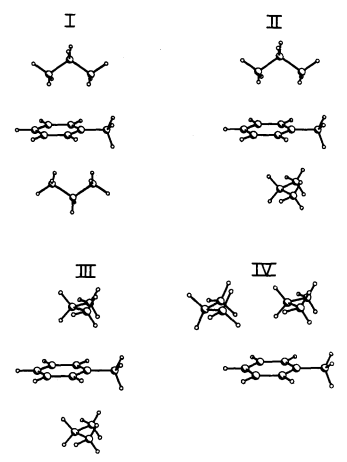


FIG. 10. Minimum energy configurations of Tol(C₃H₈)₂. Only configuration IV was minimized in the computer modeling. The other configurations are composites of the geometries of Tol(C₃H₈)₁ shown in Fig. 9. The atomic positions of configurations IV are given in Table VIII. Other configurations are possible.

binding energies are used. However, computer modeling of the propane dimer suggests that several geometries are stable, some of which have relatively small binding energies. If a reasonable fraction of the propane dimers in the beam are hot or are in geometries with small binding energies (i.e., $\sim 500~\rm cm^{-1}$), the relative number of isotropic clusters in the beam would be larger than otherwise expected. Additionally, some of the intense features assigned to isotropic clusters could be associated with anisotropic geometries not yet located by the computer modeling.

4. Nucleation

The nucleation process in the jet expansion can be better understood by employing a simple kinetic scheme. We assume, as has been previously suggested (I), that small vdW clusters are formed by solute collisions with solvent dimers (or possibly larger species). While this may not be the only cluster formation mechanism, it is consistent with all benzene and toluene cluster data and can be justified by reasonability arguments. A solvent dimer colliding with the solute produces an "activated complex" in much the same fashion as is typically considered in a bimolecular reaction. This complex can then decay through three different pathways: (1) vibrational predissociation of the solvent dimer to form a homogeneously nucleated solute(solvent)₁ vdW cluster; (2) collision with a third body (He, typically) to generate a inhomogeneously nucleated solute(solvent)₂ vdW cluster; or (3) decay back to the reactants (solvent)2 and solute.

Data presented in the Sec. III indicate that details of the solvent dimer binding energy and solute-solvent binding energy are important for the relative concentrations of homogeneously and inhomogeneously nucleated products. Thus, finer detail in the overall kinetic scheme should be considered. The mechanism suggested for the reaction is represented as follows:

in which m indicates a vdW bond with significant vibrational excitation, O indicates a solvent molecule, M stands for an inert third-body collision partner, and represents the solute. Assuming d[II]/dt = 0, and ignoring the first step in the reaction chain, the ratio of concentration [IV] to [III] is given by

$$\frac{[IV]}{[III]} \simeq \frac{k_4}{k_3} \frac{(k_1 + k_2 + k_3 + k_4)}{k_1}.$$
 (1)

The amount of inhomogeneous nucleation relative to homogeneous nucleation is proportional to this ratio. Two reasonable approximations can be made for this expression. First, $k_4 < k_1$, k_2 as k_4 is at most $\sim 10^{11}$ s⁻¹ in the backing region (and certainly much less in the beam) while k_1 , $k_2 \sim 10^{12}$ s⁻¹. If this were not the case, inhomogeneously nucleated clusters would always dominate the intensity. Second, $k_1 \simeq k_2$ because the density of states is high for both species at the given energies (300–900 cm⁻¹) and energy is not expected to concentrate in one bond or the other over many vibrational periods. With these approximations Eq. (1) becomes

$$\frac{[\text{IV}]}{[\text{III}]} \propto \frac{k_4}{k_3},\tag{2}$$

where k_4 depends only on the collision rate of third bodies with species I and II and should be roughly the same for all the systems studied here. The difference in the relative numbers of isotropic species for the various systems must then be associated with the changes in k_3 from cluster to cluster.

The transformation of species II to species III can be viewed as a reaction with a potential barrier equal to the binding energy of the solvent dimer. The rate constant k_3 can be written as

$$k_3 \sim \exp[-BE_{LL}/E_{LL}],\tag{3}$$

in which BE_{LL} is the solvent dimer binding energy and E_{LL} is the energy in the solvent dimer bond for species II. If equilibrium is established $(k_1, k_2 \gg k_3)$ before VP occurs and the excess energy in the intermediate is statistically distributed between the solvent-solute and solvent-solvent bonds, then $E_{LL} \sim \frac{1}{2}BE_{SL}$, in which BE_{SL} is the solute-solvent binding energy. This latter quantity takes on the role of kT for the reaction II \rightarrow III. Thus k_3 becomes a function of both the solvent-solvent and solute-solvent binding energies. Table IX shows that the relative number of isotropic clusters in the system scales approximately with this form for k_3 . Of course, the detailed behavior of the ratio [III]/[II] depends on the appropriateness of the various approximations for each system.

B. Large clusters

The intensity maximum in the spectrum of clusters with more than three solvent molecules is much less red shifted with respect to the free molecule than the intensity maximum in the comparable cryogenic solution. This observation is directly attributable to the conditions of the experiment and the nature of the nucleation processes in the expansion. Inhomogeneous nucleation dominates the formation of large clusters, especially for clusters involving better solvents such as propane. Since inhomogeneous nucleation produces more anisotropic clusters, most of the large clusters formed in the jet will be anisotropic and will exhibit small spectral shifts; larger spectral shifts are found for isotropic (solution-like) clusters.

TABLE IX. The ratio of the solvent dimer binding energy (BE_{LL}) to the solute-solvent binding energy (BE_{SL}), the evaluation of Eq. (3) in the text, and the experimental intensity ratio for isotopic and anisotopic clusters. All energies are in cm⁻¹ based on the calculations.

Solute	Solvent	BE _{LL}	$\mathrm{BE}_{\mathit{SL}}$	$\frac{\mathrm{BE}_{LL}}{\mathrm{BE}_{SL}}$	$rac{I_{ m iso}}{I_{ m aniso}}$
Toluene	Methane	227	667	0.34	3
Benzene	Methane	227	589	0.39	1
Toluene	Ethane	503	926	0.54	0.2
Toluene	Propane	786	1128	0.70	
Benzene	Ethane	503	778	0.65	0
Benzene	Propane	786	1040	0.76	0

The spectral shifts in the jet generated clusters have been determined to be largely due to interaction between the solvent and the aromatic π system of the solute. If the clusters are to serve as potential models of condensed phase systems, the difference between the principal gas to cluster and gas to liquid shifts must be explored. (Note nonetheless, that some cluster intensity is found at the same spectral shift as that observed for the analogous solution.^{1,4,5}) While in general the shifts for clusters are saturated with a solvent molecule on each side of the aromatic ring, larger shifts might arise through two mechanisms: forced crowding of the solvent around the solute so that more molecules effectively interact with the π system of the solute, and interaction of the solvent with the solute in the plane of the ring. These latter positions are not local minima in the potential surface of small clusters. However, the cage structure of the liquid would force solvent molecules to interact with the solute in the ring plane. This in-plane interaction may also contribute to the spectral shift of larger vdW clusters and, in particular, could be responsible for the weak intensity at approximately the value of the solution spectral shift reported for these systems previously.⁶ Since we have no direct information on the in-plane interactions, the relative contribution to the shift from these two possible sources is difficult to assess.

V. CONCLUSIONS

Two-color time-of-flight mass spectroscopy and simple potential calculations have been employed to determine cluster binding energy, spectral shift, geometry, and nucleation processes for toluene solvated by methane, ethane, and propane. The major findings of this effort can be summarized as follows:

- (1) Cluster binding energy is related to the polarizability of the two species in the cluster and is apparently not a sensitive function of solute-solvent geometry.
- (2) The observed spectral shift for a cluster is due mostly to the solvent interaction with the solute aromatic

 π system and therefore is a sensitive function of cluster geometry.

- (3) The nucleation of clusters under the experimental beam conditions is suggested to occur through the interaction of solvent dimers or larger aggregates and the solute monomer—whether isotropic clusters (homogeneous nucleation) or anisotropic clusters (inhomogeneous nucleation) arise depends on the relative size of the solvent dimer binding energy and the solute-solvent binding energy.
- (4) The benzene and toluene cluster systems appear to behave in a consistent and predictable fashion—the data analysis for both systems has been carried out such that trends are transferable from one system of clusters to another.
- (5) The exponential-6, atom-atom potentials employed in the computer modeling of cluster geometry and binding energy, which have been quite successfully used over the years in treating condensed phases, are apparently reasonable potentials for the modeling of clusters.

ACKNOWLEDGMENTS

We are grateful to Professor O. Anderson and C. Schauer for help with the computer simulated drawing and for use of the crystallographic computing system.

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