

CCA-1800

YU ISSN 0011-1643

UDC 543.42

Original Scientific Paper

The Conformations and Vibrational Spectra Including Matrix Isolation of 1,3-Dibromo-2,2-dimethylpropane

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Received November 19, 1987

The IR spectra of 1,3-Dibromo-2,2-dimethylpropane as a liquid and as two crystalline solids obtained at low temperature and at high pressure were recorded. Additional IR spectra of this compound, matrix isolated in argon and nitrogen at 14 K, were obtained using nozzle temperatures of 300, 450 and 700 K. Raman spectra including polarization measurements were recorded at various temperatures between 340 and 230 K. Crystalline solids were obtained by freezing the liquid and by shock freezing the vapour at 85 K with subsequent annealing.

The GG and AG conformers were present in the low temperature and in the high pressure crystals, respectively, and the enthalpy differences were 5.6 (liquid) and 4.2 kJ mol⁻¹ (vapour) with GG being the more stable. An additional conformer AA was detected in the liquid and in the matrix isolated spectra, being approximately 6.4 and 6.5 kJ mol⁻¹ less stable than GG in the liquid and vapour, respectively. Vibrational assignments of the GG and AG spectra are presented, supported by a normal coordinate analysis.

INTRODUCTION

Neopentane (C(CH₃)₄) has a spherical shape and represents an interesting derivative of propane. If at least two hydrogens attached to different carbon atoms are substituted with halogens, the molecules will have conformers of different abundance.¹ Neopentane² and some³ of the halogenated derivatives have plastic phases, stable in the temperature range between the low temperature anisotropic crystal phase and the liquid, while other neopentanes have no plastic phases.⁴

The vibrational spectra of various halogenated neopentanes have recently been investigated in our laboratory, including 1-chloro-2,2-dimethylpropane,⁵ 1,3-dichloro-2,2-dimethylpropane (DCDMP),⁶ 2-(chloromethyl)-2-methyl-1,3-dichloropropane⁷ and 2,2-di(chloromethyl)-1,3-dichloropropane⁸ and 2,2-di(bromomethyl)-1,3-dibromopropane.⁹ Apart from the latter two compounds, all of these exhibited plastic phases. Due to the interesting conformational properties of these molecules we decided to extend our studies to include 1,3-di-

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bromo-2,2-dimethylpropane (hereafter abbreviated DBDMP); certain preliminary observations have recently been published.⁹ A full account of our results for DBDMP, including conformational stabilities, the conformation present in the crystal phases and complete interpretations of the spectra are given in the present paper.

EXPERIMENTAL

DBDMP was purchased from ICN Pharmaceuticals Inc. and purified by preparative gas chromatography. Its identity was confirmed by mass spectroscopy. IR spectra of the low temperature amorphous and annealed crystalline solids deposited on a CsI window ($4000\text{--}200\text{ cm}^{-1}$) at 85 K were obtained. The sample was isolated in argon and nitrogen matrices (1:500), deposited at 14 K from vapour mixtures at 300, 450 and 700 K by means of an electrically heated quartz nozzle. High pressure spectra were obtained with a diamond anvil cell (DAC) with diamonds of type IIa, equipped with a spacer of bronze and interfaced with a 4x beam condenser from P.-E. The sample was carefully observed in a polarization microscope during the increase and decrease of the pressure.

The IR spectra were recorded with a Perkin-Elmer model 225 spectrometer ($4000\text{--}200\text{ cm}^{-1}$) and an evacuable FTIR model 114c from Bruker ($4000\text{--}40\text{ cm}^{-1}$). A closed cycle helium cooled cryostat from Air Products was employed for the matrix isolation experiments.

The Raman spectra were recorded with a Dilor triple monochromator spectrometer RT 35 interfaced with the Aspect 2000 computer of the Bruker spectrometer. A Spectra Physics argon ion laser model 2000 was employed using the 514.5 nm line with 90° and 180° illumination geometries. Low temperature Raman spectra of the liquid and of the crystalline solid were obtained in a glass tube of 2 mm inner diameter surrounded by a Dewar cooled by gaseous nitrogen.¹⁰ Conventional cryostats cooled with liquid nitrogen consisting of either a cooled copper finger (Raman) or a CsI (mid IR) or Si window (far IR) were employed for the amorphous and crystalline solids.

RESULTS

Among the large number of the recorded spectra, we have reproduced IR spectra of the amorphous and crystalline solids at 85 K in the regions $1500\text{--}1200\text{ cm}^{-1}$ (Figure 1) and $1000\text{--}400\text{ cm}^{-1}$ (Figure 2). An IR spectrum of a high pressure crystal is reproduced in Figure 3, whereas those of DBDMP matrix isolated in nitrogen with nozzle temperatures of 300 and 700 K are given in Figure 4. Parts of the Raman spectra of DBDMP as a liquid and

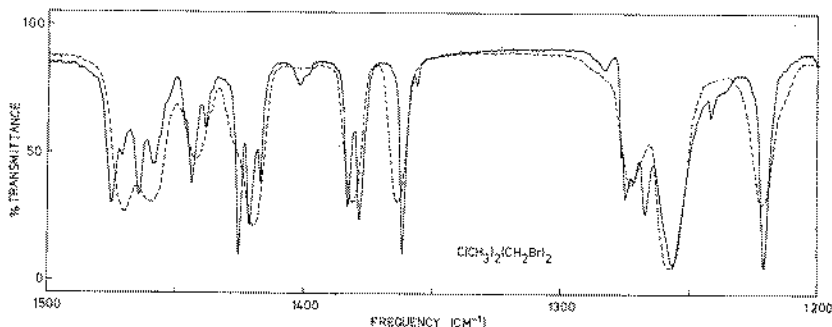


Figure 1. IR spectrum of 1,3-Dibromo-2,2-dimethylpropane (DBDMP) in the $1500\text{--}1200\text{ cm}^{-1}$ range as an amorphous (dotted line) and annealed crystalline sample (solid line) at 85 K.

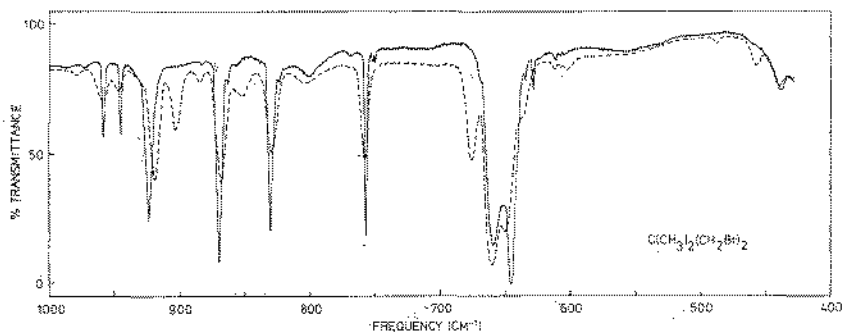


Figure 2. IR spectrum of DBDMP in the 1000—400 cm^{-1} range as an amorphous (dotted line) and annealed crystalline sample (solid line) at 85 K.

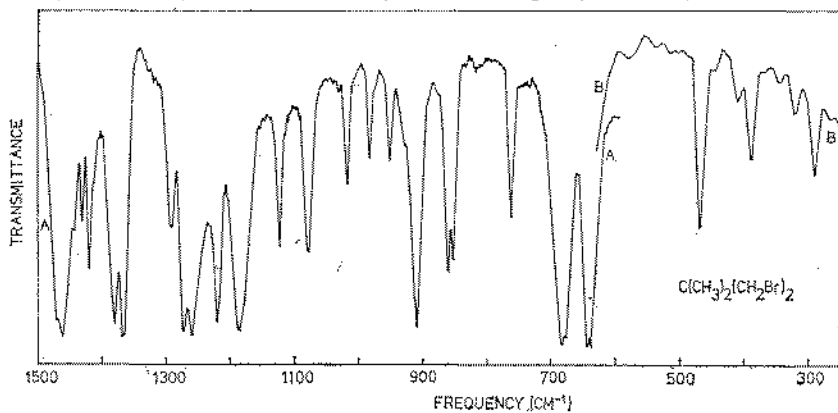


Figure 3. IR spectra of DBDMP as a high pressure crystal at ca. 20 kbar, curve A, resolution 2 cm^{-1} , curve B resolution 8 cm^{-1} .

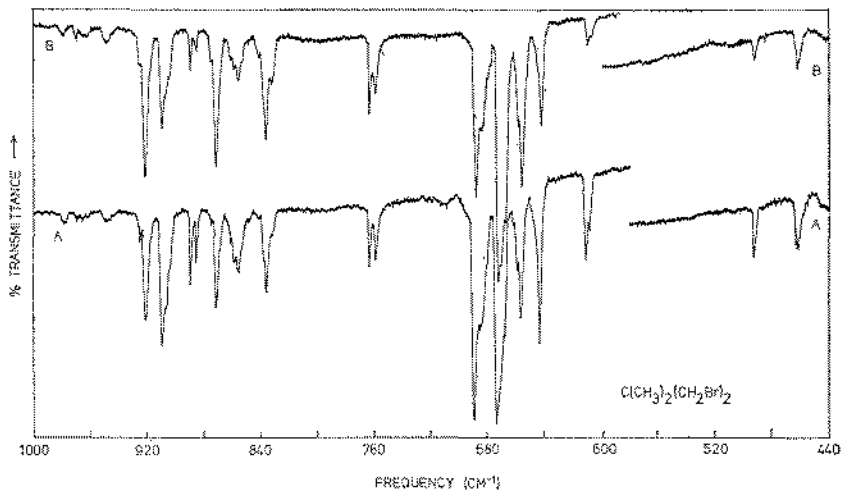


Figure 4. IR spectra of DBDMP in nitrogen matrices (1:500) recorded at 14 K, nozzle temperatures 700 K (curve A) and 300 K (curve B) in the range 1000—400 cm^{-1} .

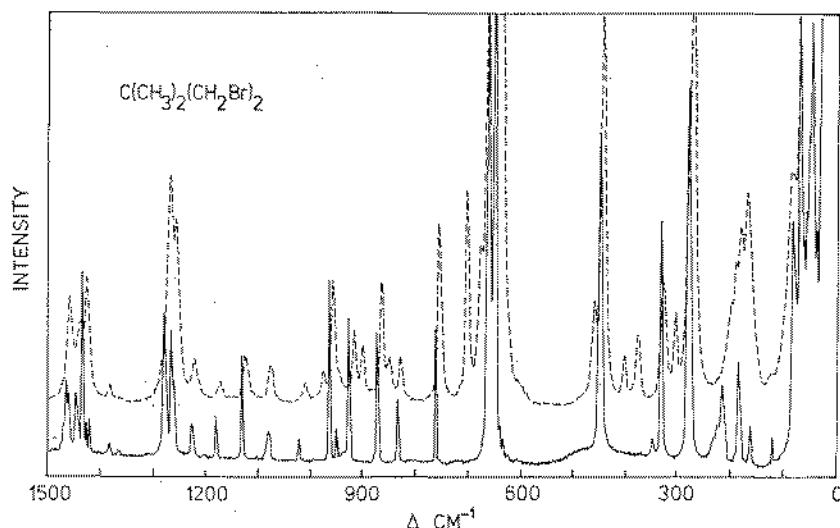


Figure 5. Raman spectra of DBDMP as a liquid (dotted line) at 285 K and as a crystalline solid at 85 K (solid line).

as a crystalline solid are given together in Figure 5. The spectral results essential for the interpretations are listed in Table I. Additional spectral data (e. g. IR matrix isolated spectra in the nitrogen matrix as well as IR and Raman spectra of the amorphous solid at 85 K) are omitted from the Table for the sake of brevity.

TABLE I

Infrared and Raman Spectral Data for 1,3-Dibromo-2,2-Dimethylpropane (DBDMP)^a

Infrared				Raman		Assignment ^b			
Liquid	Ar-matrix	Low. temp. crystal	High pressure crystal	Liquid	Low temp. crystal	GG	AG	AA	
3006w	3018vw ^c	3015vw	3031mw	3008w, D	3015mw	ν_1	ν_1		
		3009vw	3019mw		3008mw	ν_{24}	ν_2		
	3002vw								
	2991mw		2988m						
	2981w								
2966s	2976s	2977m	2984s		2977m	ν_2	ν_3		
	2971s	2973s	2978s	2959s, P	2974s	ν_{25}	ν_4		
		2965w			2966w				
	2964m	2962s	2966mw		2963s	ν_3, ν_{26}	ν_5, ν_6		
	2959vw	2956m			2957w				
2941w	2947w	2948w	2941s		2942w, P	2946w	ν_4	ν_7	
2931w	2935w	2933vw	2928vw	2928w, P	2933vw	ν_{27}	ν_8		
		2921vw	2918w		2923vw				
		2907vw	2913vw	2908w, P	2910w				
2894vw	2894vw	2892vww	2896w	2893vw, P	2893vw	ν_5	ν_9		
		2881vww			2883vww				
		2874vw							

Table I cont'd.

Infrared				Raman		Assignment ^b		
Liquid	Ar-matrix	Low. temp. crystal	High pressure crystal	Liquid	Low temp. crystal	GG	AG	AA
2872w	2875vw	2871w 2859w	2877w	2878vw, P	2872w 2860w	ν_{28}	ν_{10}	
2837w	2847vw 1473s	2850w 2838w 1475s 1471w	2835w	2835w	2837vww 1472vww	ν_{29}		
1470s	1467m \uparrow		1471s					
1463m	1464w	1465s	1461s	1460s, P	1464m	ν_6	ν_{11} ν_{12}	
1460w	1459w	1459w	1456w		1459w	ν_{30}	ν_{13}	
1443w	1444w	1444m	1446w	1441m, P	1445m	ν_7	ν_{14}	
1428w	1428m	1426vs	1431m	1428s, P	1433s	ν_8	ν_{15}	
1423s	1425s	1422s 1418m	1420s		1426w 1420w	ν_{31}	ν_{16}	
1384s	1386s	1383s 1379s	1380s	1385vw, P	1382vww	ν_9	ν_{17}	
1366s	1384w 1372vw 1369m \uparrow	*	1367s	1367vww			ν_{18}	
	1367s 1362vww 1289vw \uparrow	1362vs *	* 1291m		1365vww	ν_{32}		
1286w	1278mw 1271vw \uparrow	1275m *	* 1275s	1287vww	1278s	ν_{33}	ν_{19}	
1270m	1267m 1262vs	1267s 1256vs	* *	1269vs, P	1263s 1259m	ν_{10} ν_{34}	ν_{20}	
1257vs	1257m \uparrow 1231m	*	1260s	1258m, D			ν_{21}	
1223s	1225m	1221vs	*	1222w, P	1224w	ν_{11}		
1214m	1215w \uparrow 1213vw 1183mw	*	1220s	1215vw, P \uparrow	*		ν_{22}	
1180m	1181mw 1178m	\uparrow *	1185s				ν_{23}	
1172s	1176m	1176m	*	1173w, D	1177mw	ν_{35}		
1153w	1155vw \uparrow	*	*					AA
1126s	1128w	1126s	*	1126w, P?	1127m	ν_{12}		
1120w	1119vw \uparrow 1089vw	* 1079w	1123m	1119vw \uparrow	*		ν_{34}	
1079m	1083w	1076m		1077w, D	1078w	ν_{36}		
1076m	1077vw 1073vw	\uparrow *	1079m				ν_{25}	

Table I cont'd.

Infrared				Raman		Assignment ^h		
Liquid	Ar-matrix	Low. temp. crystal	High pressure crystal	Liquid	Low temp. crystal	GG	AG	AA
1012w	1011w	1016m	1018mw	1011w, P	1017w	ν_{13}	ν_{26}	
		1013m						
977w	978vw \uparrow	*	983mw	976w, P \uparrow	*		ν_{27}	
958mw	960vw	959m	*	958m, P	960m	ν_{14}		
947w	946vw	946m	951mw	947vw, D	947w	ν_{37}	ν_{28}	
917s	921s	924s	*	918mw, D	924m	ν_{38}		
901s	904m \uparrow	*	910s	902w, P? \uparrow	*		ν_{29}	
884w	888w \uparrow	*	*	886vwv	*			AA
866s	872m	870vs	*	866m, P	870m	ν_{15}		
850s	855mw \uparrow	*	861m	851w, P \uparrow	*		ν_{30}	
	838w	830s	853m	830w, D	830mw	ν_{39}	ν_{31}	
830s	833m							
	768vw	757s	763m	757m, P	758m	ν_{16}	ν_{32}	
756s	761mw							
702vw	710vw \uparrow	*	*	704m, P \uparrow	*			AA
679vs	683s \uparrow	*	685vs	677m, P \uparrow	*		ν_{33}	
	679s	659vs	*	663s, D	657vs	ν_{40}		
666vs	673vs							
654vs	660s	645vs	*	650vs, P	645vs	ν_{17}		
	647vw	\uparrow *	643vs	640vs, P \uparrow	*		ν_{34}	
642m	645mw							
629vw		629w						
605w	611w \uparrow	*	*	605vw, D \uparrow	*			AA
		602vvw						
492w	494w \uparrow	*	*					AA
460m	461w \uparrow	*	469s	460w, P \uparrow	*		ν_{35}	
		447vw	*	443m, P	443s	ν_{41}, ν_{18}		
443w	443vvw	441vw						
421vvw								
397vw	402vvw \uparrow	*	411w	403w, P \uparrow	*		ν_{36}	
375vw	376vw \uparrow	*	390m	376w, P \uparrow	*		ν_{37}	
352vw	~340vvw?	341w	*		342vvw	ν_{19}		
327w	328vw	327mw	*	329mw, D	327s	ν_{42}		
317vw	304vvw	*	320w	306w, P	*		ν_{38}	
283vw	284vw \uparrow	*	289m	282w, P? \uparrow	*		ν_{39}	
272vw	277vw	275w	*	272s, P	275s	ν_{20}		
251vvw		266vw						
224vw	~226vvw?	222w	241w		221vw	ν_{43}	ν_{40}	
		211m	224		212w	ν_{21}	ν_{41}	
203w		181mw	*	201w, D	180mw	ν_{44}		
~190?		*	199w	190w, P \uparrow	*		ν_{42}	
~180?		*	*	181w, P \uparrow	*			AA?
169vw		157vw	*	168m, P	158w	ν_{22}		
~152?		*	145w	~155?	*		ν_{43}	
123vvw		109?	*	123vw, D	116vw	ν_{45}		

Table I cont'd.

Infrared				Raman		Assignment ^b		
Liquid	Ar-matrix	Low temp. crystal	High pressure crystal	Liquid	Low temp. crystal	GG	AG	AA
~108?		*	114w 95w					^v 44 lat.
88vw		*	~75w	83w, P?				^v 45
			72w		76w	lat.		lat.
			~50w					
60vw			55w 40w 31w	50w	62m 39mw 21s	^v 23		

^a Many weak bands, particularly in the regions 4000—3100 cm^{-1} and 2800—1500 cm^{-1} were omitted.

^b For the meaning of conformers GG, AG and AA see Figure 6.

^c Abbreviations: s, strong; m, medium; w, weak; v, very; P, polarized; D, depolarized; asterisks (*) denote vanishing bands; arrows (\uparrow) denote bands enhanced at higher temperatures.

DISCUSSION

Conformers

Four staggered conformers of DBDMP are theoretically possible with symmetries and notation C_2 (GG), C_1 (AG), C_{2v} (AA) and C_s (GG₁), as depicted in Figure 6. The conformer GG₁ of C_s symmetry has one parallel 1,3-interaction between the C-Br bonds, leading to a decreased stability of the order of 15¹¹ to 35¹² kJ mol⁻¹, as calculated by molecular mechanics calculations. The GG₁ conformer is accordingly of no interest when the conformational abundance in DBDMP is discussed, and attention will be focused upon the GG, AG and AA conformers as previously done for DCDMP.⁶

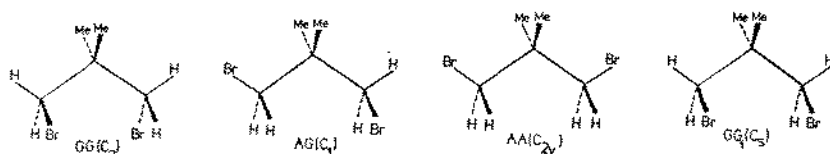


Figure 6. Possible conformers of DBDMP.

Low Temperature Spectra

Deposition of DBDMP from vapour in either an infrared or a Raman cryostat led to an initially amorphous film. The spectra of this film were quite similar to those of the liquid. When the sample was annealed to approximately 180 K, it became crystalline and many bands vanished (Figures 1, 2, 5). Also when the liquid was slowly cooled in the Raman capillary,¹⁰ the sample readily crystallized after an initial supercooling. The wave numbers of the vanishing bands are marked with asterisks in Table I. From the intensities of the disappearing bands in IR as well as in Raman, we feel

that the vanishing conformer(s) are present in less than 30% abundance at room temperature.

In the earlier studies of DBDMP and the corresponding chloro and iodo compounds dissolved in freon (CBrF_3) by NMR spectroscopy^{11,13} only the GG conformers were detected at the coalescence temperatures. Moreover, both of the two molecular mechanics calculations^{11,12} list the relative stabilities in the order GG, AG, AA and GG₂ for DBDMP as well as for DCDMP although the relative values differ considerably between the two studies. From these results, from the close similarity of the persistent conformer spectra with those of DCDMP⁶ and from the results of the normal coordinate analysis, it can be concluded without doubt that the GG conformer is present in the low temperature crystal. Given all these reasons we are also confident that the main conformer which disappears on crystallization is AG.

Since the complete IR spectrum of the AG conformer alone was recorded as a high pressure crystal (see below), we could detect some weak IR and Raman bands present in the liquid and in the amorphous phase which were present neither in the low temperature nor in the high pressure crystals. In the matrix isolated IR spectra these bands, although weak, showed large enhancements with higher nozzle temperatures, in accordance with a high ΔH° difference from the stable GG conformer. Undoubtedly, these bands are due to the AA conformer. The two most prominent bands at 704 and 605 cm^{-1} attributed to the AA conformer (see Table I) fit quite well with the calculated frequencies of species a_1 and b_2 at 712 and 595 cm^{-1} , respectively. In our preliminary paper⁹, submitted before the complete set of data was collected, we failed to recognize the existence of the third conformer (AA). Moreover, no third conformer was observed in our spectra of DCDMP⁶ although ca. 7% of the AA conformer was suggested in the gaseous electron diffraction measurements at 333 K.¹ We now plan to record hot nozzle matrix spectra of DCDMP as well, in order to detect possible bands of the AA conformer in this molecule.

The enthalpy differences ΔH° between the three conformers were calculated from five pairs of liquid Raman bands recorded at different temperatures between 340 K and 230 K. A numerical separation of the overlapping bands was achieved by defining band profiles as intermediate between the Lorentz and Gauss functions.¹⁴ The results based upon band areas are listed in Table II, together with the enthalpies obtained from the matrix hot nozzle experiments in the vapour phase.

High Pressure Spectra

Application of approximately 15 kbars pressure in the diamond anvil cell was sufficient to produce an anisotropic crystal. Just as it had been earlier observed for DCDMP,⁶ the conformer present at high pressure was not the one stable in the low temperature crystal. Rather, the high pressure crystal conformer was the major conformer which vanished when the low temperature crystal was formed. As it was mentioned earlier, we are confident that it is the AG conformer.

The variable temperature measurements of the liquid and of the vapours (observed in the matrices) reveal enthalpy differences of 4–5 kJ mol^{-1}

TABLE II

Thermodynamic Parameters Related to the Conformers of DBDMP

Conformers	Band pairs (cm ⁻¹)	Calculated ΔH° (kJ mol ⁻¹) from spectra		
		N ₂ -matrix ^a	Ar-matrix ^a	Liquid ^b
AG/GG	1182/1177	4.51	4.02	—
	1080/1087	4.26	—	—
	904/ 921	4.22	3.52	5.93
	855/ 872	3.88	3.87	—
	683/ 673	4.89	4.44	5.32
	645/ 660	4.64	4.54	5.95
	460/ 443	—	—	5.02
	306/ 329	—	—	5.72
283/ 273	3.96	4.37	—	
ΔH° (AG—GG)	[kJ mol ⁻¹]	4.2 ± 0.4	—	5.6 ± 0.6
K_{298} (AG—GG)		0.37		
ΔG°_{298} (AG—GG)	[kJ mol ⁻¹]	2.5		
ΔS° (AG—GG)	[J K ⁻¹ mol ⁻¹]	5.8		
Conformers	Band pairs (cm ⁻¹)	Calculated ΔH° (kJ mol ⁻¹) from spectra		
		N ₂ -matrix ^a	Ar-matrix ^a	Liquid ^b
AA/GG	1155/1177	5.95	6.66	—
	888/ 921	5.81	6.12	—
	611/ 660	7.14	7.27	—
	704/921	—	—	5.68
	704/673	—	—	6.77
	704/ 660	—	—	6.74
ΔH° (AA—GG)	[kJ mol ⁻¹]	6.5 ± 1.0	—	6.4 ± 0.8
K_{298} (AA—GG)		0.036		
ΔG°_{298} (AA—GG)	[kJ mol ⁻¹]	8.2		
ΔS° (AA—GG)	[J K ⁻¹ mol ⁻¹]	-5.8		

^a Calculated from absorbance values.^b Calculated from band areas, see text.

between GG and AG (Table II). The same was true for DCDMP⁶ although the exact values were not measured for this compound. The GG and AG conformers have statistical weights 2 and 4, respectively.

The ΔG° (AG—GG) values calculated in Table II at atmospheric pressure may change considerably with pressure. From the expression $(\partial G/\partial p)_T = V$ we can derive $\Delta G(P) - \Delta G(1) = \int_1^P \Delta V dP$ where $\Delta G(P)$ and $\Delta G(1)$ are the ΔG (AG—GG) at pressure P and at 1 bar, respectively. Considering ΔV (AG—GG) as constant in the pressure range we get:

$$\Delta G(P) \approx \Delta G(1) + (P - 1) \Delta V \approx \Delta G(1) + P \Delta V.$$

$\Delta G(P)$ will be different from $\Delta G(1)$ depending upon ΔV . Obviously, if ΔV (AG—GG) = V_m (AG) - V_m (GG), in which V_m , the molar volume of each conformer, is negative, $\Delta G(P)$ would be smaller than $\Delta G(1)$ and the equilibrium displaced towards AG with increasing pressure. However, it is by no means obvious that the AG conformer should have a smaller volume

that the GG. It has been suggested¹⁵ that the number of *gauche* interactions in each conformer should to a first level of approximation determine the relative molar volume of haloalkanes. However, the number of C...Br *gauche* interactions is the same (four) for GG and AG as well as for AA. It seems significant that the two dihaloneopentanes DBDMP and DCDMP⁶ both crystallize in the AG conformer under pressure. The corresponding 1,3-dibromopropane¹⁶ as well as 1,2,2,3-tetrachloropropane⁶ with hydrogens and chlorines in the 2-positions, respectively, crystallize in the GG conformer in the low temperature as well as in the high pressure crystals.

Twice during the release of pressure, a phase was observed in the microscope which was almost certainly plastic, as judged from the appearance of »wavy lines«. However, this phase was apparently stable only in a quite small pressure range and we did not succeed in recording a spectrum. It is our experience from halogenated neopentanes⁵⁻⁷ that the molecular motions within the plastic phase permit the presence of all conformers in equilibrium similar to the conditions in the vapour and liquid states.

Matrix Spectra

Depositions of DBDMP both in the argon and nitrogen matrices (1:500 ratio) were made at nozzle temperatures of 300, 450 and 700 K. Implicit in the hot nozzle technique used for conformational analysis is the assumption that the thermodynamic equilibrium of the vapour at the nozzle temperature will be maintained¹⁷ when the vapour mixture is trapped on the window at 14 K. It is our experience that for molecules with a high conformational barrier, such as the saturated halogenated hydrocarbons, the vapour conformational equilibrium is upheld in the matrix. On the other hand, for molecules with low conformational barriers such as ethylazide¹⁸ the equilibrium is found to be quite different in the argon and in the nitrogen matrices. Moreover, if the barrier is very low (less than 5 kJ mol⁻¹), a window temperature of 14 K is not sufficient to trap the unstable conformers. As recently observed for monohalogenated cyclobutanes¹⁹, the axial conformer was completely absent in the matrix spectra.

As seen from Figure 4, a number of IR bands of DBDMP in a nitrogen matrix change significantly in relative intensities from nozzle temperatures 300 to 700 K. The intensity variations with temperature were quite similar in the two matrices, and only the argon matrix data are included in Table I in which the enhanced bands are marked with vertical arrows. As it is apparent from Table I, the bands present in the high pressure crystal (AG) are enhanced with temperature; those present in the low temperature crystal (GG) diminish in relative intensities. In addition, the weak IR bands at 1155, 888, 710, 611 and 494 cm⁻¹ which are absent in both crystal spectra are enhanced with temperature and must belong to the AA conformer.

Thermodynamic Functions

In order to calculate the enthalpy difference ΔH° between the conformers from band intensities at various temperatures, it is essential that the bands employed are »pure« (belong to one conformer only). Since DBDMP has 45 fundamentals for each conformer, fundamentals (or combination bands or

overtone) of one of the three conformers present can easily overlap with those of another conformer, leading to erroneous values. The conformational purity of the band pairs used in the calculations was therefore checked by the complete appearance/disappearance in the two crystal spectra and by the wavenumber separation obtained in the force constant calculations. As apparent from Table II, 13 band pairs (7 in the N₂ and 6 in the Ar matrices) were employed for GG/AG and 6 for the GG/AA calculations. Absorbance values, rather than band areas were used, and the results for each band pair treated by a least squares procedure. The ΔH° values between AG and GG calculated from the vapour phase (IR matrix) show slightly lower values than those obtained in the liquid (Raman).

From the average values of ΔH° (XX—GG) (in which XX = AG or AA) in Table II the equilibrium constants:

$$1) K(XX/GG) = [XX]/[GG] = (g(XX)/g(GG)) e^{-\Delta H^\circ(XX-GG)/RT}$$

were calculated. The Gibbs free energy functions $\Delta G^\circ(XX-GG) = -RT \ln K(XX-GG)$ and the entropy $\Delta S^\circ(XX-GG) = -[\Delta G^\circ(XX-GG) - \Delta H^\circ(XX-GG)]/T$ are obtained. From these equations we get $\Delta S^\circ(XX-GG) = R \ln [g(XX)/g(GG)]$ in which $g(XX)$ and $g(GG)$ are the statistical weights of conformers XX and GG, respectively. Thus, if $\Delta H^\circ(XX-GG)$ is temperature independent, the same is true for $\Delta S^\circ(XX-GG)$.

The total entropy difference $\Delta S^\circ(XX-GG) = S^\circ(XX) - S^\circ(GG)$ between two spectroscopically distinguishable conformers in the vapour phase is:

$$2) \Delta S^\circ(XX-GG) = \Delta S^\circ_{\text{rot}}(XX-GG) + \Delta S^\circ_{\text{vib}}(XX-GG) + \Delta S^\circ_{\text{sym}}(XX-GG).$$

The terms $\Delta S^\circ_{\text{rot}}$ and $\Delta S^\circ_{\text{vib}}$ are the rotational and vibrational entropy differences, respectively, while $\Delta S^\circ_{\text{sym}}$ include the changes in symmetry and optical activity. The rotational and vibrational entropy can be expressed as $S_{\text{rot}} = R(3/2 + \ln Q_{\text{rot}})$ and $S_{\text{vib}} = R[u/(e^u - 1) - \ln(1 - e^{-u})]$ in which Q_{rot} is the classical rotational partition function and $u = hc\nu/kT$ where ν are the wavenumbers of the normal vibrations (Table IV). The symmetry term of the entropy is $S_{\text{sym}} = R \ln n/\sigma$ in which n and σ are defined in Table III. The S_{rot} and S_{vib} vary with temperature, but the calculations showed that they were nearly identical for the GG, AG and AA conformers. Therefore, their differences made negligible contributions to the entropy of Eq. 2, and the main part is connected with $\Delta S^\circ_{\text{sym}}(XX-GG)$.

Two of the argon matrices were annealed for 45 minutes at 35 K with no consistent change in the spectra, aside from minor differences ascribed to matrix effects. Even 12 hours annealing of the nitrogen matrix at 33 K led to no change in the conformational abundance. Thus, the barrier between the conformers must be at least 11 kJ mol⁻¹, using the value 11.2 for log A²⁰ in the Arrhenius equation and assuming that a change as low as 5% would be detected.

Normal Coordinate Analysis

Force fields for the various conformers of DBDMP were derived by transferring force constants employed in the studies of chloro-,⁵ bromo-,²¹ dichloro-,⁶ trichloro-,⁷ tetrachloro-⁸ and tetrabromoneopentane.⁸ We have used dimensionless internal stretching coordinates $\Delta r/r_0$, whereby it is possible to

TABLE III
Principal Moments of Inertia, Symmetry Parameters and Entropy Terms for Conformers of DBDMP

Conformers:	GG	AG	AA
I_a [amu Å ²] ^a	313.05	214.58	115.98
I_b [amu Å ²]	754.46	1066.56	1378.67
I_c [amu Å ²]	954.57	1167.69	1380.95
g (XX) ^b	2	4	1
n	2	2	1
σ_c	2	1	2
$\pi\sigma_1$	9	9	9
$\sigma = \sigma_c \cdot \pi\sigma_1$	18	9	18
Q_{rot} (298K)	1.160×10^6	1.262×10^6	1.148×10^6
Q_{rot} (700K)	4.171×10^6	4.542×10^6	4.128×10^6
S_{rot}^0 (298K) ^c	128.56	129.27	128.48
S_{rot}^0 (700K)	139.21	139.91	139.12
S_{vib}^0 (298K)	100.44	99.58	100.41
S_{vib}^0 (700K)	241.51	240.69	241.50
S_{sym}^0	-18.27	-12.50	-24.03
ΔS_{298}^0 (XX—GG)	—	5.62	-5.87
ΔS_{700}^0 (XX—GG)	—	5.65	-5.86

^a Principal moment of inertia; assuming C—H, 1.093Å; C—C, 1.54Å; C—Br, 1.94Å and all angles tetrahedral.

^b g (XX) — statistical weight of conformers (XX = GG, AG, AA).

n — number of distinguishable conformers.

σ_c — symmetry number refers to the symmetry of the entire molecule.

$\pi\sigma_1$ — product of symmetry numbers of those parts of the molecule which can rotate around its single bonds.

^c All entropy terms are in [J K⁻¹ mol⁻¹].

reduce²² considerably the number of different force constants in a series of halogen substituents. The procedure has been described^{7,22} earlier. A 41 parameter force field was derived by a least squares treatment of *ca.* 280 observed frequencies below 1500 cm⁻¹, taken from the various conformers of the halogenated neopentanes.^{8-8,21} Our results for the three important conformers GG, AG and AA are listed in Table IV together with the wave-numbers of the observed vibrational bands assigned as fundamentals.

Spectral Interpretation

The observed vibrational bands assigned as fundamentals for the GG, AG and AA conformers are listed in Table IV, together with the results of the force constant calculations. Since spectra of the isolated GG and AG conformers were recorded in the low temperature and in the high pressure crystals, respectively, these interpretations should be quite reliable. A large number of bands were present in both crystals, revealing coinciding GG and AG fundamentals. Frequently, this was supported by the matrix spectra, showing separate peaks for each conformer in one or both matrices. To keep the size of Table I within reasonable limits, the nitrogen matrix data were omitted, as were certain weak peaks in the argon matrix spectra.

With low symmetry of the two most abundant conformers GG (two-fold axis) and AG (no symmetry), the Raman polarization data had limited significance. However, all the polarized bands were attributed either to *a*-modes

TABLE IV

Observed and Calculated Fundamental Frequencies for the Conformers of DBDMP

ν	GG		PED ^b	AG		AA			
	Obs. ^a	Calc.		Obs. ^c	Calc.	Obs.	Calc.		
1	3006	3009	a	CH ₂ as(99)	3031	3008		3007	b ₁
24	3006	3007	b	CH ₂ as(99)	3019	3007		3007	a ₂
2	2966	2967	a	CH ₂ ss(95)	2984	2967		2966	b ₂
25	2966	2966	b	CH ₂ ss(96)	2978	2966		2966	a ₁
3	2966	2963	a	CH ₃ as(97)	2966	2964		2963	a ₁
26	2966	2961	b	CH ₃ as(99)	2966	2962		2963	b ₂
4	2941	2961	a	CH ₃ as(96)	2941	2961		2961	a ₂
27	2931	2960	b	CH ₃ as(96)	2928	2960		2960	b ₁
5	2894	2883	a	CH ₂ ss(99)	2896	2884		2884	a ₁
28	2872	2883	b	CH ₂ ss(99)	2877	2883		2883	b ₁
29	1470	1464	b	CH ₃ ad(84), CH ₃ r(10)	1471	1468		1475	a ₁
6	1463	1461	a	CH ₃ ad(85), CH ₃ r(9)	1461	1463		1463	b ₂
30	1460	1458	b	CH ₃ ad(87), CH ₃ r(8)	1456	1457		1456	b ₁
7	1443	1454	a	CH ₃ ad(91), CH ₃ r(7)	1446	1453		1452	a ₂
8	1428	1412	a	CH ₂ δ(66), BrCHδ(27)	1431	1407		1407	b ₂
31	1423	1410	b	CH ₂ δ(66), BrCHδ(28)	1420	1406		1407	a ₁
9	1384	1380	a	CH ₃ sδ(97), CCs(5)	1380	1386		1389	a ₁
32	1366	1377	b	CH ₃ sδ(99), CCs(4)	1367	1375		1373	b ₁
33	1286	1296	b	CCs(39), BrCHδ(19)	1291	1297		1297	b ₁
10	1270	1292	a	CCs(43), CH ₂ tw(18)	1275	1282		1289	b ₂
34	1257	1267	b	CH ₃ w(24), CH ₂ r(19)	1260	1281		1270	a ₁
11	1223	1230	a	CH ₂ w(40), BrCHδ(39)	1220	1222		1208	a ₁
35	1172	1181	b	CCs(38), BrCHδ(23)	1185	1192		1191	b ₂
12	1126	1111	a	BrCHδ(32), CH ₂ tw(28)	1123	1120	1155*	1141	a ₂
36	1079	1095	b	BrCHδ(36), CH ₃ r(21)	1079	1050		1066	b ₁
13	1012	1018	a	CH ₃ r(85), CH ₃ ad(5)	1018	1019		1019	a ₂
37	947	952	b	CH ₃ r(57), CCs(36)	983	953		953	a ₁
14	958	947	a	CCs(35), CH ₃ r(31)	951	949		952	b ₁
38	917	912	b	CCs(52), CH ₃ r(39)	910	914	868*	911	b ₂
15	866	860	a	CCs(33), CH ₃ r(26)	861	854		848	a ₂
39	830	838	b	CH ₂ r(36), BrCHδ(34)	853	842		843	b ₁
16	756	760	a	CCs(81), CBrs(7)	763	758		757	a ₁
40	666	674	b	CBrs(72), CCCδ(22)	685	679	710*	712	a ₁
17	654	650	a	CBrs(74), CCBrd(19)	643	644	611*	595	b ₂
18	443	445	a	CCCδ(67), CBrs(6)	469	454	494*	493	b ₂
41	443	443	b	CCCδ(67), CBrs(7)	411	402		402	b ₁
19	352	340	a	CCCδ(92), CH ₂ Brτ(9)	390	382		377	a ₁
42	327	323	b	CCCδ(60), CBrs(40)	320	308		342	a ₂
20	272	263	a	CCCδ(47), CBrs(24)	289	274		268	b ₂
43	224	246	b	CH ₂ r(91), Br..Hs(5)	241	255		238	a ₂
21	211	238	a	CH ₂ r(78), CCCδ(10)	224	239		214	b ₂
44	203	188	b	CCBrδ(42), Br..Hs(25)	199	176	181?	194	a ₁
22	169	167	a	CCBrδ(44), Br..Hs(24)	145	155		137	a ₂
45	123	138	b	CH ₂ Brτ(70), BrCHδ(9)	114	105		108	a ₁
23	60	47	a	CH ₂ Brτ(73), BrCHδ(12)	~75	73		71	b ₁

^a IR liquid phase values.^b Potential energy distribution defined as $X_{ik} = 100F_{ik}L^2/\lambda_{ik}$; a, asymmetric; s, symmetric; s, stretch; δ, deformation; r, rock; w, wag.; tw, twist.; τ, torsion; Br, Hs, non bonded stretch interaction.^c IR high pressure crystal values.

* Values from Ar-matrix.

of GG, or to AG or overlapping GG and AG fundamentals. An exception is the polarized band at 704 cm^{-1} attributed to the a_1 mode ν_{11} of the AA conformer, associated with C—Br stretch.

The two torsional modes of the CH_2Br groups were calculated to lie above and below 100 cm^{-1} for each of the conformers. They were tentatively assigned to weak IR and Raman bands in the low frequency region (Tables I and IV) and correspond to similar data recorded for 1,3-dibromopropane.²³

The agreement between the observed and calculated fundamentals is satisfactory for all the conformers. It should be stressed that the force field was transferred from the work on other neopentanes and was not at all adjusted to the present spectra. Some of the weaker bands in Table I which were not assigned as fundamentals are interpreted as combination bands or overtones. Certain low frequency bands observed in the crystal spectra are undoubtedly lattice modes.

Acknowledgement. The authors are grateful to A. Horn for drawing the figures and to G. Isaksen for purifying the sample. A. G. received a post-doctorate fellowship from NTNF. D.L.P. received support from the Norwegian Marshall Fund and from NAVF.

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SAŽETAK

Konformacije i vibracijski spektri, uključujući matičnu izolaciju, 1,3-dibromo-2,2-dimetilpropana

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Snimljeni su infracrveni spektri 1,3-dibromo-2,2-dimetilpropana u tekućini i dva kristalna stanja pri niskoj temperaturi i pri visokom tlaku. Dodatni infracrveni spektri ovoga spoja, izoliranog u matricama argona i dušika pri 14 K, dobiveni su korištenjem temperatura sapnice od 300, 400 i 700 K. Ramanovi spektri, uključujući polarizacijska mjerenja, snimljeni su pri različitim temperaturama između 340 i 230 K. Kristali su dobiveni smrzavanjem tekućine i smrzavanjem šokom pri 85 K uz napuštanje.

Konformer *GG* i *AG* prisutni su u kristalima pri niskoj temperaturi, odnosno visokom tlaku, dok je razlika u entalpiji 5,6 (tekućina) i 4,2 kJ mol⁻¹ (para), pri čemu je *GG* stabilniji. U spektrima tekućine i matrici detektiran je dodatni konformer *AA*, koji je nestabilniji od konformera *GG* za približno 6,4 kJ mol⁻¹ u tekućini, odnosno 6,5 kJ mol⁻¹ u pari. Prikazana je vibracijska asignacija *GG* i *AG* spektara, podržana analizom normalnih koordinata.