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> Studies of Some Thermally Unstable Complexes of Group V Trihalides with Trimethylamine and [2H,]Trimethylamine by Infrared, Raman, and Matrix-isolation Raman Spectroscopy, Normal Co-ordinate Analysis, and Structural Methods

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> Low-temperature i.r. and Raman spectroscopy, ligand deuteriation, and matrix-isolation Raman techniques are used to determine the shapes of the thermally unstable complexes PCl₃,NMe₃ and PBr₃,NMe₃. Comparison of the observed and calculated vibrational spectra for the phosphine complexes with the similar data for AsCla, NMea (whose crystal structure is known) suggest that they are all isostructural and form pseudotrigonal bipyramidal molecular complexes having a stereochemically active lone pair of electrons in the equatorial plane, which can be considered to be replacing an equatorial chlorine atom in the isoelectronic Group IV complexes MX4,NMe3.

OF particular interest is the effect of the lone pair of electrons on same Group V trihalides in determining the stereochemistry of their complexes with trimethylamine. Many mono- and bis-trimethylamine complexes are known, for example, MCl4,NMe3 (where M = Si, Ge, Sn, and Ti); 1 SiCl₃ \hat{X} , NMe₃ (where $\hat{X} = H$, F, Br, or I); 2 and MCl₃,2NMe₃ (where M = Al, Ga, or In),3 and have been shown to have a trigonal bipyramidal shape around the central metal atom, with the NMe₃ group always lying in the axial position.

Vibrational data have not yet been reported for complexes of the type MX3,NMe3 although it is known that PCl₃, PBr₃, and AsCl₃ form unstable 1:1 adducts with

One can consider the Group V problem in the light of our knowledge of the isoelectronic Group IV species SiCl₄,NMe₃ and GeCl₄,NMe₃ which are known to have Can trigonal bipyramidal stereochemistries.1 The replacement of a chlorine atom in SiCl4, NMe3 or GeCl4,-NMe₃ by a lone pair of electrons to yield PCl₃,NMe₃ or AsCl₃, NMe₃ might not be expected to alter the basic stereochemistry. Thus the axial NMe3 is expected to persist in the Group V complexes, with the lone pair of electrons in either an axial or equatorial site.

The X-ray structures for AsCl₃,NMe₃ 6 and SbCl₃,-¹ I. R. Beattie and T. R. Gilson, J. Chem. Soc. (A), 1965,

² I. R. Beattie, D. H. Boal, and G. A. Ozin, to be published. I. R. Beattle, D. H. Boal, and G. A. Ozin, J. Chem. Soc.
 I. R. Beattle, T. R. Gilson, and G. A. Ozin, J. Chem. Soc.
 (A), 1968, 1993.
 R. R. Holmes, J. Phys. Chem., 1960, 64, 1295.

PhNH₂ 7 have recently been reported and show the complexes to have pseudotrigonal bipyramidal coordination about the As and Sb, with the ligand lying

axially as shown in structure (1). There is no known example of pseudotrigonal bipyramidal species (e.g., ClF₃, SF₄, ICl₂-, etc.) in which the stereochemically active lone pair(s) of electrons lies anywhere but in the equatorial plane.

Our vibrational study for the PCl3-NMe3 system can be divided into four parts: (a) the low-temperature spectra of (i) the pure 1:1 complex, (ii) the 1:1 complex in a CO₂ matrix, and (iii) the 1:1 complex in a NMe₃ matrix; (b) isotope substitution data for the completely deuteriated complex PCl₃, N(CD₃)₃; (c) comparison of these data with the analogous systems PBr₃,-NMe₃ and AsCl₃,NMe₃; and (d) normal co-ordinate analysis of PCl₃, NMe₃, PBr₃, NMe₃, and AsCl₃, NMe₃ and their perdeuterio-analogues.

In considering the trigonal bipyramidal structure of the Group V complexes from a general point of view, the

⁵ R. R. Holmes and E. F. Bertant, J. Amer. Chem. Soc., 1958,

R. R. Holmes and E. F. Bertant, J. Amer. Chem. Soc., 1968, 80, 2980; 1960, 82, 5285.
 M. Webster and S. Keats, J. Chem. Soc. (A), 1971, 836; J. Amer. Chem. Soc., 1958, 80, 2980.
 R. Hulme and J. C. Scruton, J. Chem. Soc. (A), 1968, 2448.

atoms could arrange themselves around the central atom in four possible configurations (A)—(D). In the examination of the vibrational spectra which are expected for these models, the modes which involve mainly

the trimethylamine group will vary little for the four models. Normal modes that will vary substantially from (A) to (D) will be the metal-halogen stretching modes.

It has been extensively proven that main-group metal-halogen stretching modes appear at *low* frequency when the central metal atom is light (e.g., Al, Si, P, or S) and does not move appreciably during the

from ones involving the NMe₃ ligand modes as the latter experience 30—70 cm⁻¹ shifts on deuteriation of the ligand whereas the metal-halogen stretching modes generally show only small shifts (2—5 cm⁻¹). Additional information can be obtained by replacing one halogen by another as in PCl₃,NMe₃ and PBr₃,NMe₃.

AsCl₃,N(CH₃)₃ and AsCl₃,N(CD₃)₃.—The former compound is considered in the light of its known X-ray structure.⁶ The spectra for the complex and its perdeuterio-analogue are in Table 1, together with the computed frequencies for C_s symmetry. Arsenic-chlorine totally symmetric stretching modes (A') are observed at 384 and 363 cm⁻¹. A more explicit assignment to axial and equatorial motions is not possible owing to the highly mixed character (determined from the calculated potential energy contributions) of these modes. The strong i.r. band observed at 354—356 cm⁻¹ (which is very weak in the Raman) is assigned to the A" (equatorial) arsenic-chlorine stretching mode.

TABLE 1 Vibrational spectrum a of AsCl₃,NMe₃ b

AsCl _a ,N	Me ₃	AsCl ₂ ,1	$V(CD_3)_3$	Calculated for	Approximate
I.r.	Raman	Raman	Ţ.r.	AsCl _a ,NMe _a	description of mode
466s	463mw			411	A" 8NMe.
	427vvw			$\frac{420}{2}$	A' SNMe,
430vw	427VVW	400s	397w,sh		$A' \delta N(CD_3)_3$
200	384s	385ms	388s	390	A' [vAsCl
387s	363s	362m	365w.sh	349	A' \vAsCl •
0.00	2028	354w,sh	354vs	376	$A''' \vee AsCl_{equ} + \delta N(CD_3)_3$
356s	262vvw	33±w,3n	250s,br	•	22 7
255s,br	232mw	226w sh	220w,br	216	A'
220w,br	232mw	208ms		212	A' A'
	100	2081118	190m	.212	. **
190w	199ms	184s	. 130111	175	A'
*	184mw	1848		. 110	
	170w		160w.br	171	A''
160w		184	100W,D1	157	A'
	155mw	154ms		107	21
		144w,sh	100	108	A"
102w	102mw	100m	102mw	73	A'
	56s			10	A

Arsenic-chlorine stretching modes are in italics. b Frequencies are reported in units of cm⁻¹. c Very mixed modes.

mode.⁸ Modes of this type are usually totally symmetrical and are expected to occur at ca. 300—380 and 200—250 cm⁻¹ for light metal-atom chlorides and bromides respectively. Thus, if models (B) and (D) are the correct structures they should exhibit one intense, low-frequency Raman line at 300—380 and 200—250 cm⁻¹ corresponding to the PCl and PBr totally symmetrical stretching modes for PCl₃,NMe₃ and PBr₃-NMe₃ respectively.

However, the metal-halogen symmetric stretching modes in models (A) and (C) involve motion of the metal atom (which will make a contribution to the G matrix and raise the frequency) and are expected at high frequency, that is ca. 420—510 cm⁻¹ for PCl₃,NMe₃ and 300—400 cm⁻¹ for PBr₃,NMe₃.

One should thus be able to deduce the stereochemistry of PCl₃,NMe₃ and PBr₃,NMe₃ on the basis of the frequencies and intensities of the metal-halogen stretching modes. These skeletal modes can easily be distinguished

The observed and calculated frequencies for AsCl₃,-NMe₃ are found to be more compatible with models (A) or (C) than with (B) or (D). Experimentally, of course, (A) is the correct structure.

PCl₃,N(CH₃)₃ and PCl₃,N(CD₃)₃.—Holmes ^{4,5} has reported that PCl₃ and NMe₃ form a single 1:1 addition complex at low temperatures. Evidence was not cited for any complexes of higher stoicheiometries in this system.

The low-temperature (-196 °C) spectra for the pure 1:1 complex and its perdeuterio-analogue are in Table 2. Unlike AsCl₃,NMe₃ which exists as a volatile solid at room temperature, PCl₃,NMe₃ is completely dissociated into its constituent species. Once again, low-frequency stretching modes are not observed, leaving the choice between models (A) and (C). The stretching modes observed in the Raman spectrum of PCl₃,NMe₃ are, as expected, of higher frequency (496 and 471 cm⁻¹) than the AsCl₃,NMe₃ complex. As with the arsenic compound,

TABLE 2 Vibrational spectrum a of PCl₃, NMe₃

			•	•
PCl ₃ ,NMe ₃		PCl _s ,N(CD _s) _s	Calculated	Approximate
Infra-red	Raman	Raman	for	description
(−196 °C)	(196 °C)	(−196 °C)	PCl ₃ ,NMe ₃	of mode
500w,sh	500vs	498vs	493	A' vPCl°
480s		483ms	511	A" vPCl
	476ms	474ms	482	A′ vPCl °
	453ms		454	A' δNMe_s
431ms	429w		419	A' Darre
•	421w,sh		410	A'' δ NMe ₃
	,	388ms		A') where
		340w	1 1 1 1 1 1 1 1 1 1	$\begin{bmatrix} A \\ A' \end{bmatrix}$ $\delta N(CD_3)$
	293vw	01011		o.NMe
277ms	270w,sh			
	263w	253w	269	A'
	243m,sh	246w,sh		
236m	236vs	228vs	235	A
		214mw	222	\overline{A}'
	197w.sh	192mw		
	189s		183	· A'
	1000	168ms	177	\overrightarrow{A}'
	159m	156m,br		**
•	141m	140m	121	A'' ·
	117mw	116m	121 .	А
	67w	110111	87	A'
	53vs		78	A'_{\cdot}
	46s			
	35s			

Phosphorus-chlorine stretching modes are in italics.
 All frequencies are in units of cm⁻¹.
 Highly coupled.

the small separation of the high-frequency P-Cl stretching modes (25 cm⁻¹), together with the satisfactory agreement between the observed and calculated spectra (Table 2) lend strong support to model (A) as being correct.

Matrix Raman Studies of PCl₃, NMe₃.—It is possible to apply the matrix isolation technique to the complex TABLE 3

Raman spectra of matrix-isolated PCl₃,NMe₃ a

_		o. 0
$PCl_3 : NMe_3 : CO_2$	$PCl_3 : NMe_3$	Approximate b
(1:1:100)	(1:100)	description of mode
505s		νPCl (1:1)
	486s	νPCl (1:2)
480m,sh		νPCl (1:1)
439m	435m	δNMe_3 (1:1) and (1:2)
426m		
	412w,sh	$\delta NMe_3 (1:2)$
	402m	νPCI (1:2)
	373 ms	νPCI (1:2)

In this Table, only the P-Cl and N-Me stretches and All frequencies are in cm⁻¹. b In deformations are shown. All frequencies are in cm⁻¹, b In these assignments (1:1) refers to PCl₃,NMe₃ and (1:2) refers to PCl, 2NMe,

PCl₃, NMe₃. This was performed (Table 3) with carbon dioxide and with trimethylamine as matrix gases. The

- ⁸ I. R. Beattie and G. A. Ozin, J. Chem. Soc. (A), 1969, 370
- D. Frieson, M.Sc. Thesis, University of Toronto, 1972.

(100:1) spectrum in CO₂ shows little change from the 1:1 complex, indicating that the complex is unimolecular rather than polymeric. It is interesting to compare the shapes of the known isoelectronic species shown as (E)—(H).8,9 Our data thus imply that on replacing a chlorine atom in the silicon complexes (E) and (F) by a chemically active lone pair of electrons in the isoelectronic phosphorus complex (G) and (H) respectively, the basic molecular framework remains unchanged.

Drastic spectral changes are observed when trimethylamine (a reactive matrix) was used both as ligand and matrix diluent. The presence of new bands and the shifting to lower frequencies of all the P-CI stretches in

the trimethylamine matrix indicates that a new compound has been formed, most likely by the co-ordination of a second trimethylamine ligand. Although it is difficult to determine the stereochemistry of the new complex on the basis of these data alone, a useful comparison can be made with SbCl₃. Antimony tri-chloride is known to react with PhNH₂ to form a 1:1 adduct whose structure 7 is known to be the same as configuration (A). The only known X-ray structure of a 1:2 adduct of SbCl₃ 10 is with triphenylarsine oxide which has the cis-pseudo-octahedral stereochemistry shown as (I). Our spectra suggest that a similar reaction

has occurred with PCl₃ to yield PCl₃,2NMe₃ since there are only high-frequency P-Cl stretching modes present [indicating a cis-arrangement of the PCl_a residue as shown in (J)] as well as a shift to lower frequencies, compatible with increased negative charge being placed on the phosphorus by an additional trimethylamine ligand.

PBr₃, NMe₃.—The spectra, as with the PCl₃ complex, show only high-frequency bands in the P-Br stretching region. The 361 and 341 cm⁻¹ peaks (strong in the Raman) are assigned as the A' P-Br stretching modes, while the 391 cm⁻¹ absorption (strong in the i.r.), is assigned as the A'' P-Br asymmetric stretching mode.

¹⁰ I. Lindquist, 'Inorganic Adduct Molecules of Oxo-compounds,' Springer-Verlag, Berlin, 1963, p. 71.

The frequencies of these bands also serves to distinguish phosphorus-halogen stretching modes from ligand deformational modes in the compounds PCl3,NMe3 and PBr₃, NMe₃. The small separation (20 cm⁻¹) between the A' stretching modes, and the greater intensity of the higher-frequency mode indicates that PBr₃, NMe₃ is isostructural with the previous 1:1 compounds.

Normal Co-ordinate Analysis.—In the absence of structural parameters for the unstable phosphorus complexes we tested the suitability of the transference force field approximation by computing initially the vibrational spectrum of AsCl₃, NMe₃ whose symmetry is known to be Cs [model (A)]. Arsenic-chlorine force constants for the approximately pyramidal AsCl₃ residue in the complex were transfered from free AsCl3. Trimethylamine force constants were the same as those used previously.1,3 The metal-ligand bond stretching force constant was set at 1.25 mdyne Å-1 which is in the range calculated from single-crystal Raman polarisation

TABLE 4 Vibrational spectrum a of PBr₃,NMe₃ b

I.r. —196°C)	Raman (—196°C)	Calculated	Approximate description of mode
	451mw,br	423	A' δNMe,
433m	423w,br	416	$A' + A''$ δNMe_3
390s	391m	399	A" vPBr
3905		406	A' vPBr
	361s		
	341m	385	A' νPBr
290s			
214w			
208w		194	$A^{\prime\prime}$
	193m	194	A'
	200	167	A'
	1 55 s		
	1000	141	A'
	100	147	n
	132m		
		119	A'
	101w	95	$A^{\prime\prime}$
		67	$A^{\prime\prime}$
		50	Ā'
		90	**

Phosphorus-bromine stretching modes are in italics. b All frequencies in cm⁻¹.

TABLE 5 Bond lengths used in vibrational analyses

Bond	Distance/Å	Bond	Distance/Å
P-Cl	2.043	P-N	2.010
P–Br	2.200	As-N	2.289
As-Cl	$2 \cdot 160$	N-Me	1.470

TABLE 6

Force constants used in vibrational analyses of MX3, NMe3 PCl₃ Force constant * 1.32 2.05 1.84 1.321.85 1.84 3·12 0·23 3.12 3.12 $=f_{RR}=f_{Rr}$ 0.200.10 0.29 0.090.060.06. 0.06^{2}

1.01

0.065

0.425 0.42 "In units of mdyne Å-1, mdyne radian-1, or mdyne-Å radian-2.

1.01

1.01

0.42

data for main-group trimethylamine complexes.11 The weakening of the arsenic-chlorine bonds in AsCl₃,NMe₃ through co-ordination to NMe₃, relative to those in free AsCl₃, could be satisfactorily accounted for by reducing the $k_{\rm AsCl}$ bond stretching force constants by 10%. The reasonably close agreement between the observed and calculated spectra for AsCl₃,NMe₃ (Table 1) lend support to the approximations made in the force field, the basis

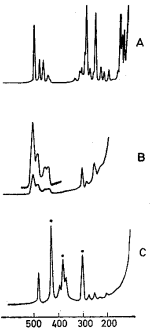


Figure 1 The low-temperature Raman spectra of A, pure PCl_3,NMe_3 (1:1); B, PCl_3,NMe_3 in CO_2 (1:100); and C, PCl_3,NMe_3 in NMe_3 (1:100)

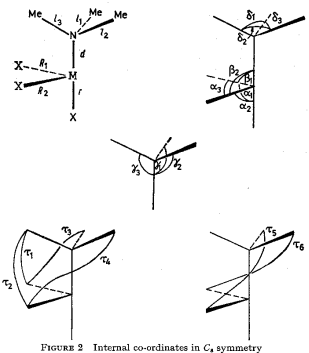
of which is used in the similar computations of the phosphorus complexes (see text and Tables 2, 4, 5, and 6). The internal and symmetry co-ordinates used in the calculations are shown in Figure 2 and Table 7.

TABLE 7 Symmetry co-ordinates (see Figure 2)

Type A' $S_{1} = (1/\sqrt{2})(\Delta R_{1} + \Delta R_{2})$ $S_{2} = \Delta r$ $S_{3} = \Delta d$ $S_{4} = (1/\sqrt{2})(\Delta l_{1} + \Delta l_{2})$ $S_{5} = \Delta l_{3}$ $S_{6} = (1/\sqrt{2})(\Delta \alpha_{1} + \Delta \alpha_{2})$ $S_{7} = \Delta \alpha_{3}$ $S_{8} = (1/\sqrt{2})(\Delta \beta_{1} + \Delta \beta_{3})$ $S_{9} = (1/\sqrt{2})(\Delta \gamma_{1} + \Delta \gamma_{2})$	Type A'' $S_{18} = (1/\sqrt{2})(\Delta R_1 - \Delta R_2)$ $S_{17} = (1/\sqrt{2})(\Delta I_1 - \Delta I_2)$ $S_{18} = (1/\sqrt{2})(\Delta \alpha_1 - \Delta \alpha_2)$ $S_{19} = (1/\sqrt{2})(\Delta \beta_1 - \Delta \beta_2)$ $S_{20} = (1/\sqrt{2})(\Delta \tau_1 - \Delta \tau_2)$ $S_{21} = (1/\sqrt{2})(\Delta \tau_1 - \Delta \tau_2)$ $S_{22} = (1/\sqrt{2})(\Delta \tau_1 + \Delta \tau_2)$ $S_{23} = (1/\sqrt{2})(\Delta \tau_1 + \Delta \tau_2)$ $S_{24} = (1/\sqrt{2})(\Delta \tau_1 + \Delta \tau_2)$ $S_{24} = (1/\sqrt{2})(\Delta \tau_3 + \Delta \tau_4)$ $S_{24} = (1/\sqrt{2})(\Delta \tau_5 + \Delta \tau_6)$
$S_{7} = \Delta \alpha_{3}$	$S_{22} = (1/\sqrt{2})(\Delta \tau_1 + \Delta \tau_2)$
$S_9 = (1/\sqrt{2})(\Delta \gamma_1 + \Delta \gamma_2)$	
$S_{10} = \Delta \gamma_3 S_{11} = (1/\sqrt{2})(\Delta \delta_1 + \Delta \delta_2)$	
$S_{12} = \Delta \delta_3$ $S_{13} = (1/\sqrt{2})(\Delta \tau_1 - \Delta \tau_2)$	
$S_{14} = (1/\sqrt{2})(\Delta \tau_3 - \Delta \tau_4)$	
$S_{15} = (1/\sqrt{2})(\Delta\tau_5 - \Delta\tau_6)$	

The normal co-ordinate calculations for the phosphorus complexes provide evidence in support of the 11 G. A. Ozin and I. R. Beattie, J. Chem. Soc. (A), 1969, 2535, and references therein.

structural assignment to model (A) for which only high frequency metal-halogen stretching modes are calculated and observed (Tables 2 and 4).



EXPERIMENTAL

All manipulations were carried out under vacuum, except for the preparation of mulls for the i.r. spectra of AsCl₃,NMe₃ (which were done in a glove box). The phosphorus and arsenic halides were outgassed and vacuum-distilled. Trimethylamine was purified by the standard procedure of treatment with silicon tetrachloride and pyridine. It was then outgassed at -78 °C and vacuum-distilled. The CO₂ used in the matrix work was research grade supplied by Matheson Co.

Holmes has used vapour pressure—composition studies to show the PCl₃, PBr₃, and AsCl₃ form 1:1 addition compounds with NMe₃ at low temperatures.^{4,5} The method of preparation of the complexes for low-temperature spectral analysis in the present study was similar to that of Holmes work. Mixtures of the appropriate trihalide and trimethylamine (1:1) were prepared in the vapour phase. The gaseous mixture was then slowly deposited through a needle valve on the central cold plate of either a low-temperature i.r. or Raman cell. Each experiment was repeated at least twice to ensure reproducibility of the spectroscopic work.

In the case of AsCl₃,NMe₃ the compound was sufficiently stable to permit analysis by conventional argentometric and pH titration techniques (Found: Cl, 46·5%; As: NMe₃, 1:1·01. Calc. for AsCl₃,NMe₃, Cl, 46·9%). The Raman spectra were recorded on a Spex 1401 spectrometer with Carson argon and krypton ion laser excitation. I.r. spectra were recorded on a Beckmann Ir 11 instrument.

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