A Spectroscopic Study of Molecular Interaction with Phosphoryl Chloride

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Raman and infrared spectra of mixtures between POCl₃ and a number of other molecules have been studied. Two types of interaction have been found, one tentatively described as dipole-dipole, the other as electron-pair acceptor-donor interaction.

The phosphoryl (P—O) bond frequency in addition compounds with POCl₃ was first studied by Sheldon and Tyre1,2 who correlated the decrease in frequency with the donor function of oxygen in the adduct formation with different acceptor molecules. In some cases, e.g. with SnCl₄ in SnCl₄·2POCl₃, they obtained more than one infrared band in the expected phosphoryl frequency range, and explained this fact as due to the co-existence of two types of POCl₃, one bonded and one free. A recent crystal structure determination 3 proves, however, that both POCl₃ molecules in the solid compound are bonded in a very similar way, and another explanation must be sought. We therefore decided to make a more complete spectroscopic study of the molecular interaction with POCl₃, going from very weak to very strong interaction, covering a wider spectral range, using infrared and Raman spectroscopy. The present paper mainly deals with weak interaction.

EXPERIMENTAL

Chemicals. POCl₃, AsCl₃, and C₆H₅COCl were purified by distillation. (B. p. +105°, +129°, and +197°C.) CCl₄, CS₂, CHCl₃ and 1,2-C₆H₄Cl₂ were purified by fractional distillation (76°—77°, 46.5°, 61°—63° and 83°—84°C). SnCl₄ and SbCl₅ were purified according to Dodd and Robinson 4. SbCl₄ was purified by sublimation.

Spectroscopic measurements. The Raman spectra, excited by Hg 4 358 Å, were photographed using an E 612 Hilger spectrograph equipped with two glass prisms and a camera having an aperture of f/5.7 (Exposure times 1—2 h). The illuminating system consisted of a Toronto type helical mercury lamp. A saturated solution of sodium nitrite in water was used as a filter. The temperature was 20°C. The spectroscopic arrangement is de-

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scribed by Kinell. Normally a Raman tube containing 20 ml was used. When the amount of solution was small, a 5 ml tube could be utilized. The tubes were filled very carefully in order to avoid as far as possible any contact between the liquids and moisture. For that purpose each tube had a ground joint by means of which it could be attached to the container holding the liquids.

The Ilford Zenith super sensitive plates used were processed with Kodak D-76 fine grain developer. The positions of the Raman lines were determined using Hartmann's dispersions formula. As reference lines Hg 4 077, 4 339 and 4 916 Å were selected. Measurements were as far as possible made on several exposures for the same solution. The accuracy of the wave numbers is ±1 – 3 cm⁻¹, depending upon the shape of the line. In a few cases the lines were too weak to be measured in the comparator. The positions of such lines have been estimated approximately and are especially marked (with circles) in the figures. No measurements of the band width and the intensities were made. The spectral slit width was 5—10 cm⁻¹.

The infrared spectra were recorded on a Perkin-Elmer Model 21 Spectrophotometer equipped with NaCl, KBr and CsBr prisms (in the Institute of Biochemistry). The spectra were run with the solutions in thin layers between two KBr plates. The spectral slit width was 3 cm⁻¹.

RAMAN SPECTRA OF POC13 MIXED WITH SOME OTHER MOLECULES

The following molecules, in order of increasing dipole moment and dielectric constant, have been studied: CS₂ = CCl₄ < CHCl₃ < 1,2-C₂H₄Cl₂ < AsCl₃ = C₆H₅N < C₆H₅COCl < SbCl₅. POC1₃ would come between C₆H₅N and C₆H₅COCl in this sequence and the spectrum of pure liquid POC1₃ has been placed accordingly in Fig. 1 showing the POC1₃ frequencies in the different mixtures. (A few lines could not be measured because of overlap with the spectrum of the other molecules.)

The very interesting molecules SO(CH₃)₂, SeOCl₂, CH₃CN and C₆H₅CN were also tried but gave complicating chemical reactions with POC1₃.

A brief summary and discussion of the results is given in the following.

1. The shifts in the four lowest frequencies are very small. The largest shift is obtained with SbCl₅ for the symmetrical P-Cl bond vibration frequency, 486 cm⁻¹, (± 7 cm⁻¹).

2. The antisymmetrical P-Cl bond vibration frequency, 584 cm⁻¹, is shifted towards higher frequencies with all solvents except C₆H₅N, the largest shifts being found with AsCl₃ (13 cm⁻¹) and SbCl₅ (20 cm⁻¹).

3. The symmetrical phosphoryl bond vibration frequency, 1 297 cm⁻¹, shows both positive and negative shifts. The largest positive shift is obtained with the nonpolar CCl₄ (7 cm⁻¹), the largest negative shifts with AsCl₃ (13 cm⁻¹) and with SbCl₅ (29 cm⁻¹).

4. There is obviously no direct correlation between dipole moment and shift. It has been pointed out, however, that a dipole-dipole interaction depends also upon the charge distribution in the molecules.

5. Fig. 1 shows that AsCl₃ and SbCl₅ are exceptional compared with the other molecules. These two molecules differ in two respects from the others. Their molecular shapes offer good opportunities for approach to the center of the positive charge in the dipole, their charge distributions are thus favourable for dipole-dipole interaction. They are, however, also the only possible acceptor molecules (Lewis acids) among the compounds studied.

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These two attributes are thus inseparable in this case. This is not the rule, however, as most Lewis acids are nonpolar molecules like SnCl₄ and SbCl₅.

6. The increase in the antisymmetrical P—Cl bond frequency has an analogy in the increases of the antisymmetrical C—C chain vibration frequency obtained by compound formation between acetone and acceptor molecules like BF₃ (52 cm⁻¹) or AlBr₃ (40 cm⁻¹) ⁷,⁸.

7. The large decreases in the symmetrical phosphoryl bond frequency by adduct formation were first measured by Sheldon and Tyree ¹,², who also found large negative shifts of the Se—O bond frequency in TiCl₄·2SeOCl₂. Similar effects on the carbonyl bond frequency in ketones have also been reported ⁷,⁹. In all these cases the shift has been caused by interaction with Lewis acids. This does not exclude the possibility, however, that the same effect can be obtained by favourable dipole-dipole interaction.

8. The spectrum with C₅H₅N is of particular interest because of the chloride ion transfer processes caused by C₅H₅N in POCl₃. Gutmann and Baaz ¹⁰ have recently studied the conductances in the similar system (C₅H₅)₃N·POCl₃ and conclude that the following equilibria exist:

\[
\text{POCl}_3 + \text{Et}_3\text{N} = [\text{POCl}_3(\text{Et}_3\text{N})^+\text{Cl}^-] \quad \text{(ion pair)} = \text{POCl}_3(\text{Et}_3\text{N})^+ + \text{Cl}^- 
\]

The corresponding K values are 2.3 × 10⁻² and 1.2 × 10⁻³, and it is thus probable that the effect on the spectrum of POCl₃ should be small. (The ionic transfer process can in this case not be preceded by a compound formation POCl₃·Et₃N because POCl₃ has no acceptor properties. The conditions are quite different with SeOCl₂ or AsCl₃ and amines ¹¹.) The small shifts obtained with C₅H₅N in this case are thus easy to understand. The possibility cannot
be excluded, however, that the exceptional influence of \( C_6H_5N \) on the antisymmetrical P-Cl bond vibration (decrease in frequency instead of increase as obtained with all the other molecules) is associated with a tendency to loosen the P-Cl bond.

**RAMAN SPECTRA OF THE SYSTEMS AsCl₃-POCl₃ AND SbCl₅-POCl₃**

The Raman spectra of solutions with varying mole ratios AsCl₃:POCl₃ and SbCl₅:POCl₃ are given in Fig. 2. A brief summary and discussion of the results follows.

1. In both systems a *continuous* change in band positions is found.
2. This change is not connected with any obvious tendency to band broadening. The band profile of the antisymmetrical P-Cl band is slightly altered, however.
3. The continuous change would be expected for a dipole-dipole interaction without formation of stable molecules. The same effect has been reported for the system \( H_2O_2-C_4H_8O_2 \) where \( H_2O_2 \) has an exceptional influence upon the dioxane spectrum (over a range of 19 cm⁻¹). The authors comment: "Die Wechselwirkung muss also dynamischer Art sein und von der ganzen Umgebung ausgehen".
4. The AsCl₃ spectrum is almost unchanged in the mixtures.
5. The SbCl₅ spectrum is changed more in the concentration range studied but not appreciably. The spectrum of liquid SbCl₅ (from Kohlrausch, *Hand- und Jahrbuch der chemischen Physik*, 9:IV, Leipzig 1943) is taken at \( +250°C \) and cannot safely be used for comparisons.
Fig. 3. Infrared spectra, in the phosphoryl band region, of the system SbCl₅—POCl₃. The shoulder near 1200 cm⁻¹ is a POCl₃ overtone band.

6. The typical Lewis acids usually change their structures by adduct formation with donor molecules like POCl₃ (cf. the structure of SbCl₅·POCl₃[12]). As a result the spectrum of the acceptor molecule disappears by the adduct formation.

7. In these cases therefore everything indicates dipole-dipole interaction. It would be very interesting to know the structure of the compound AsCl₃·POCl₃[11] (M.p. −24ºC) to find out whether the formation of the solid compound is also due to electrostatic interaction.

8. The only point of evidence indicating the existence of free and bonded molecules is the double phosphoryl band obtained with excess of POCl₃. One band is shifted towards lower frequencies, the other is almost fixed. The latter successively disappears with increasing concentration of SbCl₅ and cannot be seen at the composition SbCl₅·2POCl₃. This evidence will be discussed in the next section.

INFRARED SPECTRA OF THE SYSTEM SbCl₅-POCl₃

Infrared spectra have been taken of all the systems discussed in the preceding section. They have confirmed the Raman spectroscopic results in all cases but one, and are therefore not reproduced here. The exception is the disappearance of the double phosphoryl band at mole ratios POCl₃:SbCl₅ smaller than 2:1 (Fig. 3). The trend in the intensity ratio is the same but the intensity of the high frequency band is not reduced as much as in the Raman spectrum. It is actually as strong as the low frequency band even at concentrations where

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it has disappeared in the Raman spectrum. This indicates that the disappearance in the latter at a composition SbCl$_3$·2POCl$_3$ is purely incidental and is unrelated to compound formation. This result shows the importance of parallel work with Raman and infrared spectra. A study of the phosphoryl band only in the Raman spectrum would in this case have led to a completely erroneous interpretation.

**RAMAN SPECTRA OF MIXTURES POCl$_3$-AsCl$_3$-CCl$_4$ AND POCl$_3$-SbCl$_5$-CCl$_4$**

The Raman spectra of the solutions POCl$_3$:AsCl$_3$:CCl$_4$ = 1:4:10 and POCl$_3$:SbCl$_5$:CCl$_4$ = 1:1:5 have been studied. The POCl$_3$ bands are shown in Fig. 4 compared with the CCl$_4$-free mixtures.

The spectra indicate a diluting effect of CCl$_4$, reducing the shifts caused by AsCl$_3$ and SbCl$_5$ in agreement with the interpretation as dipole-dipole interaction.

**RAMAN SPECTRA OF SOLUTIONS OF STRONG LEWIS ACIDS IN POCl$_3$**

A preliminary study has been made of solutions of strong acceptor molecules in POCl$_3$ (SbCl$_5$:POCl$_3$ = 1:20 and SnCl$_4$:POCl$_3$ = 1:9). The results are shown in Fig. 5 together with the spectra of SbCl$_5$ (from Kohlrausch, *Handbuch der chemischen Physik*, 9:IV, Leipzig 1943), POCl$_3$ and SnCl$_4$.

A summary and discussion is given in the following:

1. The spectra of SnCl$_4$ and SbCl$_5$ have disappeared; as expected for strong Lewis acids.
2. The spectrum of free POCl$_3$ (in excess) is found in both cases as expected for strong compound formation of acceptor-donor type (in contrast to the continuous effect by electrostatic interaction).
3. New bands are formed. These have not been analysed but the knowledge of the exact crystal structure has opened possibilities for a complete analysis.
4. Some of the free POCl$_3$ bands may also belong to the spectra of the adduct molecules SbCl$_5$·POCl$_3$ and SnCl$_4$·2POCl$_3$ formed. Infrared spectra

of the solid compounds actually show that the phosphoryl band near 1297 cm\(^{-1}\) is also present in the adduct spectra.

5. The phosphoryl band region thus shows two bands with SbCl\(_5\) (the maximum shift is 117 cm\(^{-1}\)), and three bands with SnCl\(_4\) (the maximum shift is 85 cm\(^{-1}\)).

6. Similar large negative shifts with strong Lewis acids have been mentioned earlier in this paper\(^1,2,7,9\) and the splitting has also been found earlier. Examples of such splitting of carbonyl bands has been given by Susz and Chelandon \(^7,9\). It has also been reported for the Se—O band \(^2\) in, e.g., SnCl\(_4\) · 2SeOCl\(_2\).

7. The opinion that the splitting is due to the existence of two types of donor molecules in the adduct has already been rejected in the introduction to this paper. We have no alternative explanation to offer at present but think that the common occurrence of the splitting should be emphasized.

8. A complicating fact is that the phosphoryl band splitting has also been found in some pure phosphoryl compounds \(^14\). In those cases the variation is largest in the high-frequency band which has therefore been considered as the typical phosphoryl band.

9. The magnitude of the shift for the lowest frequency phosphoryl band (obtained by the splitting by adduct formation) seems to be a rough measure of the acceptor strength of the acceptor molecule. An interesting survey of the shifts in the carbonyl band has been published recently \(^8\). If this relation has general validity a very simple method of determining relative acceptor strengths is available. It can, however, only be used if the acceptor molecules to be compared are under very similar conditions.

10. The donor strength of the oxygen atom is not reflected directly in the shifts. The weak donor molecule SOCl\(_2\) does not show any shift \(^2\), but POCl\(_3\) gives a larger shift than POR\(_3\)\(^1\) with the same acceptor molecule although the latter without doubt is a stronger donor (cf. the recent assessment of the relative donor strengths of POCl\(_3\) and PO(CH\(_3\))\(_3\)\(^15\)).
CONCLUDING DISCUSSION

The preceding discussions can be summarized as follows: Negative shifts and splitting of the phosphoryl band can be caused by dipole-dipole interaction as well as by Lewis acid-base reactions. The former will give rise to a general medium effect (the shifts vary continuously with the mole ratios). The latter will lead to formation of kinetically stable adduct molecules (spectra of free and bonded molecules). The spectrum of the acceptor molecule will disappear if its structure is changed by the adduct formation. (This is not a necessary condition with molecules like HgCl₂).

If this description is correct (it can only be considered as a tentative suggestion) AsCl₃ and SbCl₅ give favourable dipole-dipole interaction with POCl₃ (in contrast to C₆H₁₂COCl), while SnCl₄ and SbCl₅ give adduct formation by acceptor-donor reactions. It is possible that there exist intermediate cases. We have here deliberately not discussed the nature of the acceptor-donor bond, pending the accumulation of more structural informations, but we intend to study these problems further by spectroscopic and X-ray methods.

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