Volatile Fluoro- and Trifluoro-methyl Diphosphorus Compounds containing Phosphorus in Two Valencies

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RECENTLY reported attempts to prepare quinquevalent diphosphorus compounds from bis(trifluoromethyl)iodophosphine sulphide, (CF₃)₂P(S)I,¹ and iodothiophosphoryl difluoride, SPF₂I,² were unsuccessful. We have now found that the reaction of the above iodophosphorus compounds with *stoicheiometric* quantities of metallic mercury at room temperature gives the desired diphosphorus disulphide compounds

$$2X_2P(S)I + Hg \rightarrow X_4P_2S_2 + HgI_2 (X = F,CF_3)$$
 (1)

The reactions are slow and require vigorous agitation to renew the reactive metal surface. In the case of the trifluoromethyl compound the product contained appreciable amounts of $(CF_3)_2P\cdot S\cdot P(CF_3)_2$ and other impurities.

The new compounds are volatile liquids with moderate

stability at normal temperatures. Spectroscopic evidence suggests that the compounds do not have the expected phosphorus-phosphorus bonded structure but rather the isomeric structure with a sulphur bridge linking quinquevalent and tervalent phosphorus atoms:

The most compelling evidence for the mixed valence structure is provided by the $^{19}\mathrm{F}$ n.m.r. spectra. At room temperature the spectrum of $\mathrm{F_4P_2S_2}$ has the appearance of a doublet of doublets with parameters: ϕ_a 60·5 p.p.m. (vs. CCl₃F), 1J_a (FP^{III}) 1320 Hz.; ϕ_b 14·0 p.p.m., 1J_b (FPV) 1218 Hz. At -80° , further spin–spin splitting transforms

each doublet into a doublet of doublets of triplets yielding the following additional coupling constants: ³J_a (FPP^V) 22.0 Hz., 3J_b (FPPIII) 15.0 Hz., and 4J (FF) 5.9 Hz.). The trifluoromethyl compound (CF₃)₄P₂S₂ shows a doublet $[\phi_a \ 53.8 \ p.p.m., \ ^2J_a \ (FP^{III}) \ 82.5 \ Hz.]$ and a doublet of doublets [ϕ_b 58.9 p.p.m., $^2J_b(\mathrm{FP^V})$ 111.7 Hz., $^4J(\mathrm{FPPIII})$ 5.0 Hz.] at room temperature. Both regions of chemical shift in each compound exhibited equal total intensity, and the observed parameters are in good agreement with those of similar compounds1-3.

I.r. spectra, particularly that of F₄P₂S₂, also support the structures with both PIII and PV, showing bands which are readily associated with PIII_F stretching (847 and 834 cm.-1) and with PV-F stretching (925 and 898 cm.-1). The CF region of (CF₃)₄P₂S₂ is complex, and suggests that CF₃ absorptions arising from two different CF₃-P environments are overlapped.

Both compounds react with hydrogen chloride to form the quinquevalent thioacid and tervalent chlorophosphine in nearly quantitative yields:

$$X_2P \cdot S \cdot P(S)X_2 + HCl \rightarrow X_2PCl + X_2P(S)SH$$
 (2)

providing chemical evidence in support of the structure. Reaction 2 proceeds well with stoicheiometric quantities of HCl in the case of the fluoro-compound, but a large excess of HCl is required to consume the trifluoromethyl compound completely.

Additional chemical evidence in support of the structure is provided by the synthesis of the mixed-valence compounds in good yields from quinquevalent thioacid and aminophosphine:

$$2X_2P(S)SH + X_2PNMe_2 \rightarrow X_2P(S)\cdot S\cdot PX_2 + Me_2NH_2^{+}S_2PX_2^{-}$$
(3)

and, in the case of the trifluoromethyl compound, the reaction:

$$(CF_3)_2P(S)SH + (CF_3)_2PCl \rightarrow (CF_3)_2P(S)\cdot S\cdot P(CF_3)_2 + HCl$$

$$(4)$$

the latter using an excess of chlorophosphine.

Reaction of difluorophosphoric acid with dimethylaminodifluorophosphine proceeds analogously to equation 3 to give fair yields of the diphosphorus dioxide compound. Again, according to the 19F n.m.r. spectrum which shows a doublet of doublets at room temperature [ϕ_a 38·3 p.p.m., ${}^{1}J_{a}(\text{FP}^{\text{III}})$ 1396 Hz.; ϕ_{b} 68.9 p.pm., ${}^{1}J_{b}(\text{FP}^{\text{III}})$ 1032.5 Hz.), the compound has the mixed valence structure, $F_2P(O)\cdot O\cdot PF_2$.

Since all diphosphine disulphides and dioxides known at present have structures containing a direct phosphorusphosphorus bond,4 the adoption of mixed valence structures in this case is somewhat unexpected. While it is tempting to suggest that a stabilization of the tervalent state relative to the quinquevalent state by electronegative substituents such as CF₃ or F is responsible for the observed structure, it is apparent that the choice of the most stable structure must involve many different factors.

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