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Radical cations in mixtures of phosphorus trichloride and dimethyl sulfide. A combined ESR and quantum chemical study

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been neglected. For these compounds it has been shown that the latter "memory effects" may cancel the nearest neighbor control. Such processes have to be expected in compounds with "unstable" local electronic configurations in the absence of "memory effects".

VII. Final Outlook

The many-particle nature of π electron bonding in alternant hydrocarbon systems has been investigated by the method of the local approach supplemented by rather simple model Hamiltonians. The present authors believe that it is most convenient for the reader to review the principal findings in the form of characteristic catchwords. (i) The π electrons in annulenes and linear polyenes are sizably localized. It seems to be necessary to substitute widely accepted traditional models for π compounds. (ii) Bond alternation leads throughout to an increase in the π electron delocalization. (iii) The respective enhancement of the charge fluctuations is probably the microscopic origin for the distortive nature of π networks. (iv) An enhancement of the charge fluctuations (decreasing electronic correlation-strength Δ_i) and increasing interatomic π correlation energies caused by bond dimerization are not mutually exclusive in larger hydrocarbons. The out-of-phase modulation is obviously the rule. (v) Cyclic 6π electron structures allow for the optimum possible π delocalization in alternant hydrocarbons; but even this takes place far from the "free-electron" limit. The latter process may lead to the situation that extended π systems prefer to form more or less decoupled spatially localized 6π subunits. (vi) Extension of linear monocyclic or polycyclic π systems is frequently not coupled to an increasing delocalization of the π electrons (effects due to the end atoms neglected); their fluctuations saturate quite early. Exceptions have been discussed in the above sections. (vii) The interatomic π correlation energy is one driving force for the bond length alternation in linear and monocyclic hydrocarbons exceeding a certain threshold dimension. (viii) Strong electronic correlations tend to attenuate Jahn-Teller or Peierls instabilities. The attenuation is maximized with increasing correlation strength; see the series C_4H_4 , C_8H_8 , $C_{12}H_{12}$. (ix) The latter effect may be of some influence in "metastable" solids, i.e. solids where the instability is scarcely suppressed. In these compounds superconductivity under strong coupling conditions may become possible.

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Radical Cations in Mixtures of Cl₃P and Me₂S. A Combined ESR and Quantum Chemical Study

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Exposure of phosphorus trichloride (Cl_3P) and dimethyl sulfide (Me_2S) dissolved in halocarbons ($CFCl_3$, CF_3CCl_3 , $CF_2ClCFCl_2$, and CH_2Cl_2) to X rays at 77 K results in the corresponding parent cations and several cation—substrate adducts. The radicals are detected and identified by ESR spectroscopy. In dilute solution exclusive formation of the parent Cl_3P^{*+} and Me_2S^{*+} radical cations is observed. In $CFCl_3$, Me_2S^{*+} exhibits superhyperfine interactions due to chlorine and fluorine nuclei of the matrix molecule(s). At increased concentration, or on warming the sample, the parent radical cations readily react with dissolved Cl_3P or Me_2S molecules to form homodimeric $Cl_3P^+PCl_3^+$ and $Me_2S^+SMe_2^+$ and heterodimeric $Cl_3P^+SMe_2^+$ radical cations with a two-center three-electron $\sigma^2\sigma^{*+1}$ bond. The heterodimer is formed in spite of a significant difference between the ionization potentials of the two constituents in reduced form. On further annealing, the $Cl_3P^+PCl_3^+$ cation rearranges to the well-known trigonal-bipyramidal Cl_4P^* radical and an ayet unidentified configuration. Candidates for the latter are proposed. In concentrated frozen solutions an unexpected reaction of $Me_2S^+SMe_2^+$ and Cl_3P is observed, resulting in the heterotrimer $Cl_3P(SMe_2)_2^{*+}$ with an octahedral configuration, exhibiting a very large ^{31}P hyperfine interaction ($A_{150} = 5115 \text{ MHz}$). Extensive ab initio calculations at the $HF/3-21G^*$ level, including calculation of isotropic and dipolar electron—nuclear hyperfine interactions, confirm the assignments and provide detailed insight into the molecular geometry, electronic configuration, and stability of the radical products.

Introduction

The use of halocarbon matrices in combination with ESR spectroscopy has eminently contributed to the knowledge of radiogenic formation and reactivity of radical cations at low temperature. In general these experiments are conducted at low substrate concentration to ensure the detection of the parent radical cations and unimolecular decomposition reactions. However, at elevated concentration and depending on the temperature and mobility in the frozen halocarbon matrix, radical cations can react with free-electron pairs of other substrate molecules. Such ionmolecule reactions usually afford two-center three-electron bonds with a $\sigma^2 \sigma^{*1}$ configuration (σ^* bond). Recently an increased interest has been noted in the literature on the nature and stability of these three-electron bonds. On the basis of high-level quantum chemical calculations, Clark postulates that, in vacuo, the stability of a σ^* bond will diminish exponentially with increasing difference in ionization potential of the two reduced substrates $(\Delta_{IP})^{1,2}$ Therefore, formation of an ion-molecule adduct consisting of two equivalent molecular parts (homodimer) should be preferred over ion-molecule adducts comprised of two different molecular fragments (heterodimer). Accordingly, a large number of ESR studies describe homodimeric radical cations, whereas reports on heterodimers are few. However, this reflects the lack of systematic study rather than the alledged intrinsic instability of the heterodimers.

Recently we reported the results of radiogenic radical formation in a mixture of trimethylphosphine (Me₃P) and dimethyl sulfide (Me₂S) in Freon.³ We demonstrated that besides the well-known

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parent radical cations (Me₃P^{*+} and Me₂S^{*+})^{4,5} and homodimers (Me₃P÷PMe₃⁺ and Me₂S÷SMe₂⁺), 6,7 the heterodimeric ionmolecule adduct Me₃P+SMe₂+ was formed. Possibly as a result of the small Δ_{IP} (IP(Me₂S) = 8.68 eV, IP(Me₃P) = 8.60 eV, Δ_{IP} = 0.08 eV) the P-S σ^* bond is intrinsically stable. In fact, detection and expected hyperfine couplings of this species had been predicted from high-level quantum chemical calculations even before experimental verification.^{2,8} Nevertheless, experimental evidence also indicates that formation of a P-S σ^* bond in condensed phases is hardly affected by the Δ_{IP} of the two molecular fragments involved. As an example P-S σ^* bonds were also detected in X-irradiated single crystals of both trialkylphosphine sulfides (R₃P+S⁻), ^{8,9} and (methylthio)triethylphosphonium iodide (Et₃P-SMe, I⁻). In spite of the different coordination number of the sulfur atom, which leads to strongly different ionization potentials, both radicals are readily formed and the spin-density distribution throughout these species is very similar. In a related study Abu-Raqabah and Symons very recently demonstrated that Me₃P^{*+} or Ph₃P^{*+} radical cations react at elevated temperatures with a chloride ion released by the irradiated freon solvent.¹⁰ This results in neutral σ* Me₃P÷Cl and Ph₃P÷Cl structures. Evidently, the Δ_{IP} of Me₃P and Cl⁻ is too large to expect a stable σ^* bond in vacuo (IP(Cl⁻) = -EA(Cl⁺) = 3.61 eV, Δ_{IP} = 4.99 eV).

The objective of the present report is to study ion-molecule reactions and adducts in mixtures of phosphorus trichloride (Cl₃P) and Me₂S in frozen halocarbon solutions. The choice for this combination was initially based on the increased Δ_{IP} (IP(Cl₃P) = 9.81 eV, Δ_{IP} = 1.13 eV) compared to the Me₃P and Me₂S combination. Our interest in the system developed during the course of this study when a variety of radical cation products and reactions was observed. Among these is the detection and identification of a radical cation composed of one Cl₃P and two Me₂S units (heterotrimer). By changing the experimental conditions, e.g. ratio or concentration of the substrates and temperature we were able to study formation and reactivity of each individual radical product. By utilizing particular freons like CF₃CCl₃ and CF₂ClCFCl₂, which allow for increased molecular mobility in the temperature range between 90 and 130 K, 7k,h,11 it proved possible to study ion-molecule reactions in detail. Furthermore, we use ab initio quantum chemical calculations to gain additional insight in the electronic and geometric structure of the radical cations. Both geometry optimization and hyperfine coupling calculations were performed at 3-21G* HF/SCF level. Vibrational analyses

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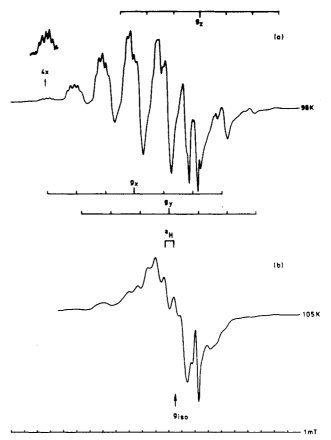


Figure 1. (a) ESR spectrum, recorded at 98 K, of the Me₂S^{*+} radical cation formed after X irradiation of a frozen 11% v/v Cl₃P/Me₂S (1:10) CFCl₃ solution. Additional ³⁵Cl and ¹⁹F superhyperfine couplings appear on the g_x tensor direction. (b) ESR spectrum, recorded at 105 K, of the Me₂S+SMe₂+ radical cation formed after X irradiation of a 30% v/v Cl₃P/Me₂S (1:1) CFCl₃ solution.

were employed to characterize the computed stationary points as local minima or saddle points.

Results and Assignment

A. Experimental. Cl₃P (Merck) was distilled prior to use. Me₂S, CFCl₃ (freon-11), CF₃CCl₃, and CF₂ClCFCl₂ (freon-113) (Aldrich) were used as received. CH₂Cl₂ (Merck) was purified over a basic-alumina column and stored on 4-Å molecular sieves. A quartz tube containing a freon or CH₂Cl₂ solution, ranging from 1 to 70% v/v Cl₃P and/or Me₂S, was degassed by repeated freeze-pump-thaw cycles in liquid nitrogen prior to irradiation and subsequently rapidly frozen at 77 K under argon. The quartz tube was transferred into a glass Dewar vessel and exposed, at 77 K, to unfiltered X irradiation from a Cu source operating at 40 kV and 20 mA for 2-6 h. The ESR spectra were recorded on a Bruker ER 200D spectrometer, operating with an X-band standard cavity and interfaced to a Bruker Aspect 3000 computer. In a typical run the field-sweep width of 0.3750 T (0.1 mT = 1 G) was sampled with 4k data points, resulting in a digital resolution of 0.09 mT (0.9 G). Expanded runs were measured with a 0.0375-T sweep width affording a 0.009-mT (0.09-G) digital resolution. The temperature was controlled with the aid of a Bruker ER 4111 variable-temperature unit. The klystron frequency was determined from ESR field positions of the hydrogen atom which is always present in X-irradiated quartz. The g values and hyperfine coupling constants, determined from the spectra, were corrected up to second order with the Breit-Rabi equations. 12 Since the powder ESR spectra can be complex due to the noncoincident g and hyperfine tensors, we expect the error in the powder assignments to be in the order of magnitude of the line width.

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TABLE I: ¹H Hyperfine Couplings^a and g Values of the Me₂S^{*+} Radical Cation, Its Homodimer Me₂S—SMe₂⁺ and Related Radical Cations Generated in Irradiated Halocarbon Solutions

radical	<i>T</i> , K	A, MHz	g _x	g _y	g z	8iso
Me ₂ O*+	77 ⁶	123; 119; 119	2.0138	2.0072	2.0045	2.0085
Me ₂ S*+	81°	57	2.019	2.019	2.0023	2.0134
_	124 ^d	57	2.019	2.0145	2.0076	2.0137
	77*	61; 60; 57	2.033	2.016	2.001	2.017
	77 ^f		2.032		2.002	2.012
	130√	59	2.019	2.0145	2.0076	2.0137
	988	60; 60; 56	2.032	2.015	2.001	2.016
	77ħ	58; 58; 56	2.033	2.017	2.002	2.017
	81 ⁱ	58	2.032	2.017	2.002	2.017
Me ₂ SSMe ₂ +	120°	18				2.010
	158 ^j	18				2.0102
	295*	19				2.0103
	100′	18				2.010
	105g	19				2.012
Me ₂ Se*+	77!	44	2.1316	2.0647	1.999	2.0651
Me ₂ Se_SeMe ₂ +	77‴	13	2.0702	2.0366	1.9964	2.0344

^aA₁(MHz) = 2.802 47 (g₁/g_e) a₁(G). ^bCFCl₃, ref 13. ^cCF₃CCl₃, ref 7h. ^dCFCl₃, refs 7f,g. ^cCFCl₃, ref 5b. ^fCFCl₃, ref 5a. ^gCFCl₃, this study. ^hCF₂ClCFCl₂, ref 5b. ^fCF₂ClCFCl₂, ref 7k. ^fCyclopropane, ref 7d. ^kH₂O, ref 7a. ^fCFCl₃, ref 1. ^mSingle crystal of Me₂Se, ref 21.

The Me₂S⁺ and Me₂S⁺SMe₂⁺ Radical Cations. ESR studies on both radicals are numerous.^{5,7} Table I lists some interesting literature data along with the results of the present study. The Me₂S⁺⁺ and Me₂S⁺SMe₂⁺ radical cations can be distinguished by the larger ¹H hyperfine coupling of the Me₂S⁺ radical cation (60 vs 18 MHz, Figure 1). Furthermore, the g tensor of Me₂S^{*+} is strongly anisotropic whereas the g value of Me₂S÷SMe₂⁺ seems isotropic (Figure 1). In Figure 1a the ESR spectrum of the Me₂S^{*+} radical cation in CFCl₃ at 98 K nicely illustrates the complex pattern caused by the overlapping ¹H hyperfine couplings and the anisotropic g tensor of the radical. Our assignment, indicated in Figure 1a, corresponds well with g values reported by Bonazzola et al.56 Prior to annealing, the spectrum at low temperature clearly exhibits additional hyperfine couplings on the g_x direction. These splittings have been attributed by Rao et al. 5a to weak σ* bonding between Me₂S^{*+} and a chlorine atom of CFCl₃. A closer look, however, reveals a five-line pattern (ratio 1:2:2:2:1) pointing to more nuclei contributing to the superhyperfine interaction. The pattern can be interpreted by a 7.0-MHz (2.5-G) superhyperfine interaction of one ³⁵Cl and one ¹⁹F nucleus. Likewise, extra hyperfine splittings of 4.2 MHz (1.5 G) were resolved on the g_z component, though no explicit pattern could be established. In agreement with previous studies, no extra coupling was resolved on the g_{ν} tensor component. In this respect, however, it is interesting to note that Rao et al.5a reported an extra 4.2-MHz (1.5-G) splitting on the g_y direction of the $(CD_3)_2S^{\bullet+}$ radical cation, which was well resolved as a result of an integer ratio of the ²H (8.4 MHz, 3.0 G) and ³⁵Cl (4.2 MHz, 1.5 G) couplings. Weak matrix interactions of radical cations to either ³⁵Cl or ¹⁹F are frequently observed in CFCl₃ solutions, ^{1,5a,14} but we believe that Me₂S^{*+} is the first example where ³⁵Cl and ¹⁹F interactions are observed simultaneously.

With respect to formation and stability of these cations we find that X irradiation of a mixture of Me₂S and Cl₃P invariably leads to Me₂S^{*+} at low Me₂S concentration, and to Me₂S⁺SMe₂⁺ dimers at increased concentration. It is interesting to note that, in contrast to their concurrent detection in the mobile CF₃CCl₃ and CF₂-ClCFCl₂ matrices, we were not able to observe the monomer and dimer radical cations simultaneously in the more rigid CFCl₃ or CH₂Cl₂ matrices. In CFCl₃, e.g., formation of either the monomer or dimer radical cation is determined by the Me₂S concentration. If the concentration is below 10% v/v, only Me₂S^{*+} is observed; above 10% v/v, solely the Me₂S-SMe₂⁺ radical cation is detected. A similar behavior was recently noted by Ambro2 et al. 7m In

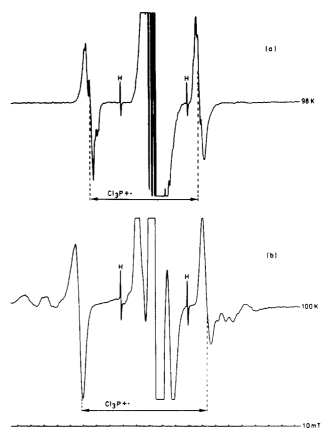


Figure 2. (a) ESR spectrum recorded at 98 K of the Cl_3P^{*+} radical cation formed radiogenically in a 11% v/v Cl_3P/Me_2S (1:10) CFCl₃ solution. Additional ³⁵Cl hyperfine interactions are present. (b) ESR spectrum of the Cl_3P^{*+} radical cation formed in a 10% v/v solution of Cl_3P in CH_2Cl_2 .

CFCl₃ the monomeric radical cation can be detected until the melting of the CFCl₃ matrix (145 K). The dimer cation is stable up to 135 K at which temperature the signal intensity rapidly decreases.

The Cl₃P*+ Radical Cation. X irradiation of Cl₃P results in the formation of the corresponding Cl₃P*+ radical cation in all halocarbons used. The ESR spectrum of this radical generated in CFCl₃ is depicted in Figure 2a. The spectrum reveals a large, nearly isotropic, ³¹P doublet with some partially resolved chlorine hyperfine splittings. The isotropic ³¹P coupling derived from this spectrum is in agreement with values established for the related (HO)₃P*+ and (MeO)₃P*+ radical cations. ^{4,6e,f} Interestingly, the ESR spectrum of the Cl₃P*+ radical cation generated from Cl₃P

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TABLE II: Hyperfine Couplingsa and g Values of the Radicals Detected in X-Irradiated Mixtures of Cl₃P and Me₂S, and Related Radical Structures Reported in the Literature

radical	<i>T</i> , K	x	³¹ P	35Cl	g _x
Cl ₂ P°	75 ^b	1	739	40	2.001
		\perp_1	-95	-14	2.023
		\perp_2	-75	-21	2.014
		iso	190	2	2.013
		2dip	549	38	
Cl₄P*	77°]	3553	178	2.00
		\perp	3391	56	2.02
		iso	3445	96	2.02
		2dip	108	82	
	130 ^{d,e}	iso	3638		2.024
"X"	130 ^{d,e}	iso	3945		2.022
Cl ₃ P*+	98 ^{d, f}	iso	2330	65	2.025
	98°	iso	2682		2.025
$(HO)_3P^{*+}$	778	1	2605		
		Τ	2039		
		iso	2227		
	_	2dip	378		
$(MeO)_3P^{*+}$	77ħ		2524		2.0038
		1	1998		2.0026
		iso	2173		2.003
		2dip	351		
	77'	H	2812		2.0049
		\perp	2255		2.004
		iso	2441		2.0043
		2dip	371		
Me ₃ P ^{•+}	100 ^j		1636		2.002
		\perp	813		2.007
		iso	1087		2.005
		2dip	549		

 $^aA_i(\text{MHz}) = 2.80247(g_i/g_e) \ a_i(G)$. $^b\text{Single crystal of Cl}_3\text{P, ref 15i.}$ Frozen Cl $_3\text{P, refs 15c,h.}$ $^d\text{CFCl}_3$, this study. $^s\text{CH}_2\text{Cl}_2$, this study. $^s\text{CH}_2\text{Cl}_3$ and CF $_2\text{ClCFCl}_2$, this study. $^s\text{H}_3\text{PO}_4$ in H $_2\text{SO}_4$, ref 4a. ^hCFCl₃, refs 6e,f. ¹CCl₄, refs 6e,f. ¹CFCl₃, ref 3.

in a CH₂Cl₂ matrix (Figure 2b) exhibits a ³¹P hyperfine interaction which is substantially larger (Table II), whereas no chlorine hyperfine interactions are resolved. A similar difference in hyperfine coupling constants and appearance of the spectrum has previously been reported by Symons and co-workers for the (MeO)₃P*+ radical cation in CFCl₃ and CCl₄ matrices. 6e,f They suggested that the (MeO), P*+ radical cation can adopt two different conformations: a spherical structure (C₃) in CCl₄ and an oblate structure (C_s) in CFCl₃. In case of Cl₃P^{*+}, different radical conformations are less obvious, and differences, when of an intramolecular nature, can only be rationalized by a change of hybridization of the central phosphorus atom, e.g. flattening of the radical structure (in extreme C_{3v} to D_{3h}). On the other hand, a specific electronic interaction with the matrix molecules cannot be ruled out.

The Cl₃P*+ radical cation is relatively stable in all halocarbon matrices up to high concentration of Cl₃P (20% v/v, Table II). However, it is absent if the cosolute Me₂S is present at a concentration over 10% v/v.

The Cl₄P Radical. The Cl₄P radical is readily detected after X irradiation of Cl₃P in a CH₂Cl₂ matrix. It is well-known that Cl₄P° possesses a trigonal-bipyramidal type structure in which the odd electron is located in a vacant equatorial position (TBP-e). The ³¹P hyperfine interaction (Table II) is in good accordance with literature values for this radical generated in pure Cl₃P.¹⁵

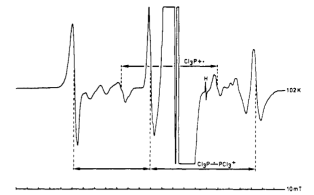


Figure 3. ESR spectrum, recorded at 102 K, of the Cl₃P+PCl₃+ radical cation formed in a 10% v/v solution of Cl₃P in CFCl₃.

TABLE III: 31P Hyperfine Couplingsa and g Values of Homo- and Heterodimeric Radical Cations Detected in X-Irradiated Mixtures of Cl₃P and Me₂S, and Related Radical Structures Reported in the Literature

radical	T, K	х	A_x , MHz	g _x
Cl ₃ PPCl ₃ +	102 ^b	iso	2260	2.020
Me ₃ P-PMe ₃ +	100°	1	1605	1.998
•		Ι.	1333	2.006
		iso	1424	2.003
		2dip	181	
$Cl_3P^{\bullet}SMe_2^+$	100^{d}	iso	3360	2.016
Me ₃ P-SMe ₂ +	100°	1	1840	2.003
•		Ĭ	1385	2.006
		iso	1537	2.005
		2dip	303	
Cl ₅ P*-	77°)	4619	2.00
-		1	4523	2.02
		iso	4555	2.02
		2dip	64	
$Cl_3P(SMe_2)_2$ **	$120^{d,f}$	iso	5115	2.023

 ${}^{a}A_{i}(MHz) = 2.80247(g_{i}/g_{e}) a_{i}(G)$. ${}^{b}CFCl_{3}$, and $CH_{2}Cl_{2}$, this study. °CFCl₃, ref 3. ^dCFCl₃, CH₂Cl₂, and CF₂ClCFCl₂, this study. Cl₅P, ref 15h. CF₃CCl₃, this study.

In the present case no chlorine hyperfine couplings were resolved. When generated in a CFCl₃ matrix, the ESR absorptions of Cl₄P^{*} increase upon annealing. In this case ClaP seems to be a secondary radical product because the rise of signal intensity coincides with the gradual loss of the Cl₃P-PCl₃+ radical (vide infra). The Cl₄P radical was not observed in CF₃CCl₃ and CF₂ClCFCl₂

The Cl₃P-PCl₃+ Radical Cation. X irradiation of Cl₃P in CFCl₃ or CH₂Cl₂ at relatively low concentration (>3% v/v) results in the formation of the homodimeric Cl₃P+PCl₃+ radical cation from an ion-molecule reaction of Cl₃P^{o+} with Cl₃P. The presence of Cl₃P-PCl₃+ cations is readily recognized by the degenerate triplet ESR spectrum shown in Figure 3, which is characteristic for a P-P σ^* type radical. As a result of the large hyperfine interaction to two identical ³¹P nuclei, the spectrum consists of four equally intense absorptions caused by a second-order splitting of the original degenerate central lines. In fact only three of the four lines can be identified because the fourth absorption is obscured by a strong central signal from the irradiated quartz tube. The ³¹P hyperfine coupling is listed in Table III. No ³⁵Cl or ³⁷Cl hyperfine interactions were resolved. Upon annealing, the radical can be detected up to 130-135 K. The decay of the radical cation seems to be coupled with the increase of the TBP-e Cl₄P* radical and another, as yet unidentified, phosphorus-centered radical (vide infra). In contrast to the parent radical cation Cl₃P^{*+}, the magnitude of the ³¹P hyperfine interaction is identical in CFCl₃ and CH₂Cl₂. Surprisingly, the Cl₃P-PCl₃+ radical cation is not observed after X irradiation of Cl₃P in the mobile CF₃CCl₃ and CF₂ClCFCl₂ matrices.

The Cl₃P+SMe₂+ Radical Cation. The Cl₃P+SMe₂+ radical

⁽¹⁵⁾ Cl₂P* and Cl₄P*: (a) Kokoszka, G. F.; Brinckman, F. E. J. Chem. Soc., Chem. Commun. 1968, 349. (b) Kokoszka, G. F.; Brinckman, F. E. J. Am. Chem. Soc. 1971, 92, 1199. (c) Begum, A.; Symons, M. C. R. J. Chem.
 Soc. A 1971, 2065. (d) Kerr, C. M. L.; Williams, F. J. Phys. Chem. 1971, 75, 3023. (e) Wei, M. S.; Current, J. H.; Gendell, J. J. Chem. Phys. 1972, 75, 2431. (f) Fullam, B. W.; Mishra, S. P.; Symons, M. C. R. J. Chem. Soc., Dalton Trans. 1974, 2145. (g) Kerr, C. M. L.; Webster, K.; Williams, F. J. Phys. Chem. 1975, 79, 2663. (h) Mishra, S. P.; Symons, M. C. R. J. Chem. Soc., Dalton Trans. 1976, 139. (i) Bonazzola, L.; Michaut, J. P.; Roncin, J. Chem. Blast 1861, 25, 4820. J. Chem. Phys. 1981, 75, 4829.

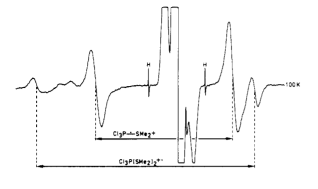


Figure 4. ESR spectrum, recorded at 100 K, of the heterodimer Cl₃P-SMe₂+ radical cation formed in a 70% v/v solution of Cl₃P and Me₂S (1:1) in CFCl₃.

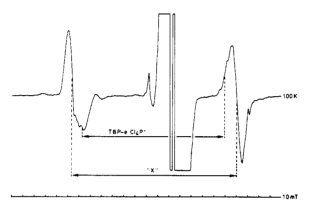


Figure 5. ESR spectrum, recorded at 100 K, of the ClaP and "X" radicals formed from Cl₃P+PCl₃+ after short annealing the sample to 148 K. The spectrum was obtained after X irradiation of a 10% v/v solution of Cl₃P in CFCl₃.

cation is readily produced when both Me₂S and Cl₃P are present in concentrations above 3% in spite of the nonzero Δ_{IP} of the two molecular fragments involved. The ³¹P hyperfine coupling (Table III) derived from the ESR spectrum shown in Figure 4 exhibits the expected increase with respect to the Cl₃P⁻⁺ and Me₃P-SMe₂⁺ radical cations. There is little doubt that the spectrum corresponds to the Cl₃P-SMe₂+ heterodimer radical cation. Analogous to radiogenic formation of Me₃P+SMe₂+ in CFCl₃ matrices,³ the structure of $\text{Cl}_3\text{P+SMe}_2^+$ is assigned to a P+S σ^* type radical. Evidently, formation of $\text{Cl}_3\text{P+SMe}_2^+$ can be rationalized by two different reaction pathways: $\text{Cl}_3\text{P*}^+ + \text{Me}_2\text{S} \rightarrow \text{Cl}_3\text{P+SMe}_2^+ \leftarrow$ Cl₃P + Me₂S^{*+}. It is experimentally not possible to distinguish between these alternatives. In CFCl₃, slight warming results in the irreversible loss of the radical at 120 K. This might be interpreted as an intrinsically lower stability of the heterodimer compared to the homodimers Me₂S÷SMe₂⁺ and Cl₃P÷PCl₃⁺. No other radicals are detected on further annealing.

The "X" Radical. Upon annealing an X-irradiated sample of 10% v/v Cl₃P in CFCl₃, we clearly detected a transition of Cl₃P-PCl₃+ into the TBP-e Cl₄P radical and a second unidentified phosphorus-centered radical (hereby called "X"). The rearrangement is also noted in CH₂Cl₂ but not in CF₃CCl₃ or CF₂-ClCFCl₂ (Table III). The corresponding ESR spectrum is shown in Figure 5. Both the TBP-e Cl₄P and "X" radicals are stable up to the melting of the matrix. The radicals are formed when Cl₃P is X irradiated without Me₂S as cosolute. Evidently, they are derived from Cl₃P alone, and a relatively low concentration of the parent substrate is sufficient (>3%). The magnitude of the ³¹P hyperfine coupling of "X" (Table II) indicates that the structure is similar to the TBP-e Cl₄P° radical. Essentially two candidates can be proposed for "X". First it can be assigned to a Cl₄P° trigonal-bipyramidal structure in which the odd electron occupies the vacant axial location (TBP-a). Ab initio quantum chemical calculations on $C_{2\nu}$ (TBP-e) and $C_{3\nu}$ (TBP-a) H_4P^* and F₄P• radicals¹⁶ predict that the TBP-a structure should exhibit



Figure 6. ESR spectrum, recorded at 120 K, of the heterotrimer Cl₃P-(SMe₂)₂*+ radical cation formed by ion-molecule reaction of Me₂S+ SMe₂⁺ and Cl₃P. The radicals are generated in a 70% v/v solution of Cl₃P and Me₂S (1:1) in CFCl₃.

a larger ³¹P hyperfine coupling than the TBP-e configuration. This would agree nicely with the present experiment (Figure 5, Table II). The second candidate is a σ^* Cl₃P+Cl radical structure. Abu-Raqabah and Symons reported formation of similar σ^* R₃P+Cl radicals by reaction of a R₃P*+ radical cation and a chloride ion descendent from the irradiated solvent. 10 However, at this moment no clear assignment can be made, and it is also conceivable that radical "X" does not correspond to either of the suggested structures.

The Cl₃P(SMe₂)₂⁴ Radical Cation. Upon annealing an Xirradiated frozen mixture of Cl₃P and Me₂S (for instance 70% v/v 1:1 in CFCl₃) a new phosphorus-centered radical is formed, exhibiting a very large 31P hyperfine coupling of 5115 MHz (Figure 6, Table III). The magnitude of the isotropic ³¹P hyperfine coupling points to a high contribution of the phosphorus 3s orbital to the SOMO.¹⁷ To our knowledge, the largest ³¹P hyperfine coupling reported until now amounts to 4555 MHz attributed to the Cl₂P^{*} radical anion which possesses an octahedral structure with the unpaired electron in a vacant position. 15h This suggests that in our case the central phosphorus atom bears at least five substituents. We assign the new formed radical to the Cl₃P-(SMe₂)₂*+ radical cation formed by reaction of Me₂S+SMe₂+ with a Cl₃P molecule. Our assignment is based on several observations. First, the larger ³¹P hyperfine coupling of Cl₃P(SMe₂)₂*+ compared to Cl₅P⁻⁻ can be explained by the fact that the positively charged SMe₂ substituents accommodate the valence electrons better than the chlorine atoms of Cl₅P^{•-}, thereby increasing spin density on the hypervalent phosphorus nucleus in the antibonding SOMO. Second, the Cl₃P(SMe₂)₂*+ radical is only detected when both Me₂S and Cl₃P are present, which points to a hetero ion-molecule reaction. Furthermore, the increase of the Cl₃P(SMe₂)₂*+ concentration is coupled with a rapid decrease of the Me₂S÷SMe₂+. In CFCl₃, the radical is absent if the concentration of Me₂S is below 10% v/v, regardless of the Cl₃P concentration (only formation of Me₂S⁺⁺!). In addition, the radical is formed in CF₃CCl₃ although the heterodimer Cl₃P+SMe₂+ is not observed. This signifies that the ion-molecule reaction between the heterodimer Cl₃P-SMe₂⁺ and a molecule of Me₂S is unlikely. Apparently, the Me₂S+SMe₂+ radical cation reacts with a Cl₃P molecule giving the trimer radical cation. Formation of this radical is observed in all halocarbon matrices used. The trimer vanishes before the matrices melt without generating new phosphorus-centered rad-

It is noted that Illies et al. 7i report a similar reaction. In a mass spectroscopic study on the homoassociation of Me₂S⁺ to Me₂S, they observed at -40 °C a prominent peak at m/z 182. It is assigned to the monopositive trimer of Me₂S less four amu.

^{(16) (}a) Howell, J. M.; Olsen, J. F. J. Am. Chem. Soc. 1976, 98, 7119.
(b) Janssen, R. A. J.; Visser, G. J.; Buck, H. M. J. Am. Chem. Soc. 1984, 106, 3429. (c) Janssen, R. A. J.; Buck, H. M. J. Mol. Struc. (THEOCHEM) 1984, 110, 139. (d) Cramer, C. J. J. Am. Chem. Soc. 1991, 113, 2439. (17) Morton, J. R.; Preston, K. F. J. Magn. Reson. 1978, 30, 577

TABLE IV: Total Energies (au)^a and $(S^2)_{UHF}$ Expectation Values of the Radical Precursors and Radicals at Their Equilibrium Geometries (Figures 7-11) (All Structures Were Calculated with the 3-21G* Basis Set)

compound	symmetry	UHF	ROHF	RHF	$\langle S^2 \rangle$
H ₂ S*+	C_{2v}	-396.476 386	-396.471 563		0.7611
H ₂ S	C_{2v}			-396.819637	
$H_2S \stackrel{\bullet}{-} SH_2^+$	C_{2h}	-793.332 709	-793.325 431		0.7717
Me ₂ S ⁺	$C_{2\nu}$	-474.174 441	-474.169 419		0.7628
Me ₂ S	C_{2v}			-474.456 985	
$Me_2S - SMe_2^+$	C_{2h}	-948.663 185	-948.656 055		0.7734
Cl ₂ P ⁺	C_{2v}			-1253.595 142	
Cl ₂ P*	$egin{array}{c} C_{2 u}^{\cdots} & & & & & \\ C_{2 u} & & & & & \\ C_{3 u} & & & & & & \end{array}$	-1253.922 205	-1253.917 335		0.7671
Cl ₃ P*+	C_{3v}	-1711.015 964	-1711.012755		0.7606
Cl ₃ P	C_{i_0}			-1711.356 885	
Cl ₄ P°	$C_{2v}^{c} \atop C_{3v}^{b}$	-2168.721 431	-2168.714 223		0.7874
	C_{3v}^{ab}	-2168.710450	-2168.703 551		0.7882
Cl ₅ P*-	C_{4v}	-2626.219 457	-2626.213718		0.7815
Cl ₃ P-SH ₂ +	C_s	-2107.859 525	-2107.853 847		0.7714
Cl ₃ PPCl ₃ •+	C_{2h}	-3422.387 232	-3422.380 975		0.7804
-	C_{2v}	-3422.321 490	e		0.7893
C_{\cdot}	C_s	-3422.382 358	e		0.7609
$Cl_3P(SH_2)_2^{*+}$	$C_{2v}(1)^c$	-2504.652 301	-2504.644 628		0.7928
	C_2	-2504.652 200	e		0.7942
	$C_{2v}(2)$ C_s^d	-2504.651 054	e		0.7936
	C_{c}^{d}	-2504.683 459	e		0.7609

^a1 au = 627.5 kcal/mol. ^bDegenerate imaginary frequency 118i cm⁻¹, E. ^cImaginary frequencies 92i cm⁻¹, B₂; 52i cm⁻¹, B₁. ^dImaginary frequencies 89i cm⁻¹ and 29i cm⁻¹, A". "No SCF convergence.

However, as they state: "the structure of this species is a tantalizing mystery".7i

B. Quantum Chemical Calculations. All calculations employed the GAUSSIAN88¹⁸ or GAMESS UK¹⁹ series of programs using the 3-21G* basis set. Odd-electron systems were calculated using the unrestricted and restricted open-shell Hartree-Fock (UHF and ROHF) formalisms, and closed-shell systems with restricted Hartree-Fock (RHF). For open-shell systems geometry optimization was started at the UHF level. The resulting structure then served as input to the ROHF geometry optimization. Harmonic vibrational frequencies were calculated at the 3-21G* level in order to characterize stationary points as minima (representing equilibrium structures) or saddle points (transition states). Magnetic properties such as the isotropic and anisotropic part of the electron-nuclear hyperfine interaction were determined, where possible, from the spin-density matrix. The isotropic hyperfine coupling was calculated from

$$A_{\rm iso} = \frac{8}{3}\pi g_{\rm e}\beta_{\rm e}g_{\rm N}\beta_{\rm N}|\psi_{\rm N}(0)|^2 \tag{1}$$

with

$$|\psi_{N}(0)|^{2} = \sum_{\mu\nu} P_{\mu\nu}^{\alpha-\beta} \langle \phi_{\mu} | \delta(r_{N}) | \phi_{\nu} \rangle \tag{2}$$

being the Fermi contact integral, and in which $P_{\mu\nu}^{\alpha-\beta}$ is the spindensity matrix. It is noted that the GAMESS UK program package gives the Fermi contact integral $^{1}/_{2}\pi$ times its true value (in au). The magnetic dipolar hyperfine tensor was calculated as

$$A_{\rm dip}(u,v) = -g_{\rm e}\beta_{\rm e}g_{\rm N}\beta_{\rm N}\sum_{\mu\nu}P_{\mu\nu}^{\alpha-\beta}\left(\phi_{\mu}\left|\frac{(r^2\delta_{uv}-3uv)}{r^5}\right|\phi_{\nu}\right)$$
(3)

where u, v are the Cartesian coordinates x, y, z. Subsequent

TABLE V: Calculated Isotropic and Anisotropic Hyperfine Couplings (MHz) of Nuclei with Sufficient Spin Density in the Calculated Radical Structures Using the ROHF Wave Function^a (Figures 7-11)

Radical Structures Using the ROHF Wave Function" (Figures 7–11)							
				$A_{ m dip}$			
radical	symmetry	nucleus	A_{iso}	1	2	3	
H ₂ S*+	C _{2v}	Н	0	-21	-2	23	
$H_2S \stackrel{\bullet}{-} SH_2^+$	C_{2h}	Н	0	-12	0	12	
Me ₂ S ^{•+}	C_{2v}	H_{avg}	20	-5	-4	9	
Me ₂ S-SMe ₂ +	C_{2h}	H_{avg}	8	-5	-3	8	
Cl ₃ P**	C_{3v}	P	1459	-196	-196	392	
Cl₄P•	C	Cl P	8 2151	-17 -98	-17 -98	34	
CI4F	C_{2v}	Cl ₁	36	-96 -34	-98 -34	196 68	
		Cl ₂	3	-3	-3	6	
	C_{3v}	P P	2537	-87	-87	174	
		Cl_1	0	-5	-5	10	
	_	Cl_2	40	-26	-26	52	
Cl ₅ P•-	C_{4v}	P	2870	-76	-76	152	
		Cl_1 Cl_2	0 18	-1 -18	-1 -18	2 36	
OLD SILL+							
Cl ₃ P-SH ₂ +	C_s	P Cl ₁	1644 0	-137 -17	−137 −17	274 34	
		Cl ₂	ő	-8	-8	16	
		H	ŏ	-8	1	7	
Cl ₃ PPCl ₃ **	C_{2h}	P	1274	-90	-90	180	
		Cl_1	36	-14	-14	28	
	- 1	Cl ₂	3	-6	-6	12	
	C_{2v}^b	P_1	2082	-210	-42	252	
	C_s^b	Cl ₁	47 1617	-58 -211	-57 -211	115 422	
	C _s	\mathbf{P}_1 \mathbf{Cl}_1	19	-211 -6	-211 -4	10	
		Cl ₂	14	-30	-25	55	
		Cl ₃	11	-27	-22	49	
$Cl_3P(SH_2)_2$ **	$C_{2v}(1)^b$	P	2694	-193	-19	212	
	C_2^b	P	2406	-191	-18	209	
	$C_{2b}^{\Sigma_b}$ $C_{2v}(2)^b$ C_s^b	P P	2393	-191	-23	214	
	C,	P	1870	-196	-191	387	

^a Conversion factors from au to MHz: ³¹P, $A_{iso}(MHz) = 1810A_{iso}$ (au) and $A_{\text{dip}}(\text{MHz}) = 215 A_{\text{dip}}(\text{au})$; ³⁵Cl, $A_{\text{iso}}(\text{MHz}) = 437 A_{\text{iso}}(\text{au})$ and $A_{\text{dip}} = 52 A_{\text{dip}}(\text{au})$; ¹H, $A_{\text{iso}}(\text{MHz}) = 4467 A_{\text{iso}}(\text{au})$ and $A_{\text{dip}}(\text{MHz}) = 531 A_{\text{dip}}(\text{au})$. ⁵UHF wave function.

diagonalization of the matrix results in the three principal values of A_{dip} and the corresponding direction cosines. Optimized geometries are given in Figures 7-11 and their total energies are listed in Table IV.

⁽¹⁸⁾ Frisch, M. J.; Head-Gordon, M.; Schlegel, H. N.; Raghavachari, R.; Binkley, J. S.; Gonzalez, C.; DeFrees, D. J.; Fox, D. J.; Whiteside, R. A.; Seeger, R.; Melius, C. F.; Baker, J.; Martin, R. L.; Kahn, L. R.; Stewart, J. J. P.; Fluder, E. M.; Topiol, S.; Pople, J. A. Gaussian 88; Gaussian, Inc.: Pittsburgh, PA, 1988. Calculations were performed on an Alliant FX/8

computer at our university.

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Lawrence Berkely Laboratory, Berkeley, 1980. Guest, M. F.; Kendrick, J.

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Laboratory, 1986. Guest, M. F.; Corley, 1980. Guest, M. F.; Kendrick, J. Games V. G. Laboratory, 1986. Guest, M. Theory, 1986. Guest, M. Theory, Chim. M. F.; Harrison, F. J.; van Lenthe, J. H.; van Corler, L. C. H. Theor. Chim. Acta 1987, 71, 117. Radial coefficients: P = 0.55, S = 0.65, and Cl = 0.75. Calculations were performed on a Convex C120 computer at the CAOS/CAMM center in Nijmegen.

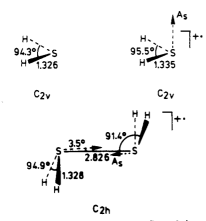


Figure 7. Optimized geometries of the H₂S, H₂S*+, and H₂S+SH₂+ structures calculated at ROHF/3-21G* SCF level within their respective symmetry constraints.

 $H_2S^{*+} + H_2S \rightarrow H_2S - SH_2^+$. The homodimer radical cation of H₂S has been subject of many high-level quantum chemical studies.^{2,20} The two stable minima found are C_{2v} and C_{2h} structures, of which the C_{2h} is the lowest in energy.² The structures resulting after geometry optimization within the $C_{2\nu}$ symmetry constraint for H_2S^{*+} and H_2S , and C_{2h} symmetry constraint for H₂S÷SH₂ are depicted in Figure 7. Characteristic for the dimer σ^* structure is the long S-S bond (2.826 Å). Furthermore, the H-S bond directions are nearly perpendicular to the S-S bond. The hydrogen hyperfine couplings calculated at ROHF/3-21G* level for both the monomer and dimer radical cations are listed in Table V. Sharing the odd electron between two sulfide moieties reduces the dipolar hyperfine coupling of the hydrogen atoms by half. Our calculations on the H₂S+SH₂+ radical reveal that the direction of the largest principal value of the sulfur dipolar hyperfine interaction (equivalent to the direction of the largest principal value of the 33S tensor) is not perfectly aligned with the S-S bond direction but makes a small angle (3.5°). The SOMO points away from the hydrogen substituents (Figure 7). Such a nonalignment is experimentally detected in the single-crystal ESR study on the Me₂S-SeMe₂+ radical cation.²¹ The direction of the largest principal value of the ⁷⁷Se hyperfine tensor, corresponding to the direction of the SOMO on that nucleus, is inclined by 2.5° to the direction of the minimal g tensor component, which is assumed to correspond to the Se-Se linkage.²¹

 $Me_2S^{*+} + Me_2S \rightarrow Me_2S - SMe_2^{+}$. Geometry optimization within C_{2v} symmetry constraint for $Me_2S^{\bullet+}$ and Me_2S and within C_{2h} for Me₂S÷SMe₂⁺ results in the structures depicted in Figure 8. All structures are true minima, and their total energies are listed in Table IV. Our results are in excellent agreement with the calculations performed by Illies and co-workers for the same systems.71 However, in their article they do not state the symmetry constraints used and give little details on the optimized structures (Figure 8).7i

The experimental ratio of the β -hydrogen hyperfine couplings of the monomer and dimer radical cations of Me₂S or Me₂Se is approximately 3.7f-h,22 The calculated hyperfine couplings are listed in Table V. The ratio of the isotropic couplings of the monomer and the C_{2h} dimer amounts to 2.5. Based on MNDO calculations,²² Williams suggested^{7f-h} that the increased ratio (>2) is a consequence of the nature of the σ^* orbital, rather than of a change in geometry at the heavy atoms. Our ab initio calculations on the C_{2h} structure of $Me_2S - SMe_2^+$, along with the computations performed by Illies and co-workers, 7i show indeed that the bond angles and bond distances of the heavy atoms do not vary much between the monomer and dimer structures. In-

Figure 8. Optimized geometries of the Me₂S, Me₂S*+, and Me₂S+SMe₂+ structures calculated at ROHF/3-21G* SCF level with their respective symmetry constraints.

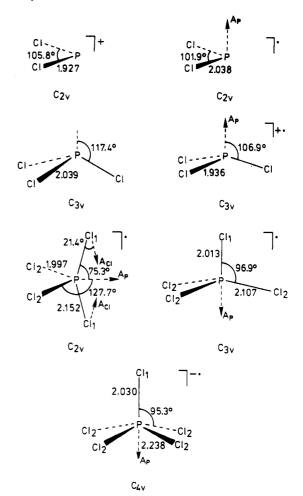


Figure 9. Optimized geometries, calculated at ROHF/3-21G* SCF level, of the structures containing one phosphorus and several chlorine atoms.

terestingly, in the C_{2h} dimer the angles between the S-S bond and the bisectors of the C-S-C bond angles are significantly larger than in $H_2S + SH_2 + (99.9^{\circ} \text{ vs } 91.4^{\circ}, \text{ Figures } 7 \text{ and } 8)$.

The principal direction of the SOMO on the sulfur atoms is almost aligned with the S-S bond. We calculated a small angle of 0.8° in which the orbital points away from the CH₃ substituents. In H₂S÷SH₂+ the angle of the H-S-H bisector to the SOMO amounts 93.9°, comparable to the 90° of the 3p_z orbital in the monomer H₂S⁺⁺. In Me₂S÷SMe₂⁺, however, this angle amounts

⁽²⁰⁾ H₂S-SH₂⁺: (a) Fernandez, P. F.; Ortiz, J. V.; Walters, E. A. J. Chem. Phys. 1986, 84, 1653. (b) Ortiz, J. V. Chem. Phys. Lett. 1987, 134, 366. (c) Gill, P. M. W.; Radom, L. J. Am. Chem. Soc. 1988, 110, 4931. (21) Nishikida, K.; Williams, F. Chem. Phys. Lett. 1975, 34, 302. (22) (a) Glidewell, C. J. Chem. Soc., Perkin Trans. 2 1985, 299. (b) Ibid.

^{1985, 551.}

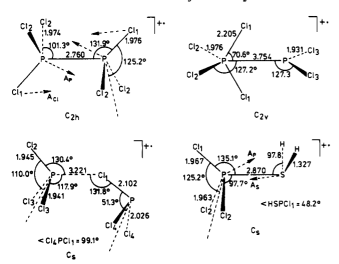


Figure 10. Optimized geometries of the three stationary points found for the Cl_3P+PCl_3+ structure (C_{2h} at ROHF, C_{2p} and C_s at UHF) and the heterodimer Cl₃P+SH₂+ radical cation (ROHF). Structures calculated with the 3-21G* basis set.

100.7°, which is considerable different. This could have a profound effect on the ratio of the β-hydrogen hyperfine couplings in Me₂S^{*+}

and $Me_2S+SMe_2^+$. $Cl_3P^{-+}+Cl_3P \rightarrow Cl_3P-PCl_3^+$. Vibrational analysis shows that the C_{3v} structure of the Cl_3P^{*+} cation is a true minimum. Radical formation shortens the P-Cl bonds, with respect to its C_{3v} precursor Cl₃P, by about 0.2 Å. A flattening of the tetrahedral structure is also noted (Figure 9). The calculated C_{2h} structure for Cl₃P-PCl₃+ is a true minimum, and the optimized geometry is again characterized by a long σ^* bond (2.760 Å, Figure 10). Formation of the homodimer results in a stabilization of 9.0 kcal/mol at the UHF level (7.1 kcal/mol at ROHF, Table IV), which is almost half of the stabilization calculated for H₃P+PH₃⁴ at different levels (e.g. $C_2 = 19.3 \text{ kcal/mol}$ at UHF/6-31G*). 2c,d

The calculated hyperfine couplings of Cl₃P⁺⁺ and Cl₃P⁺PCl₃⁺ are enumerated in Table V. The direction of the largest principal value of the ³⁵Cl tensor in Cl₃P⁴⁺ is inclined by 29.2° with respect to the largest ^{31}P hyperfine tensor component. In the dimer C_{2h} structure, the Cl₁ atoms in the mirror plane (Figure 10) bear a larger spin density than the Cl₂ atoms. The direction of the maximal principal value of the ³¹P hyperfine tensor is situated between the P-P and P-Cl₁ bonds, inclined by 24.9° to the P-P bond. A similar orientation was established in single-crystal ESR work on related σ^* radical anions of substituted diphosphine disulfides. Experiment and ab initio calculations revealed an angle of 25 to 30°.23

 $Cl_3P + H_2S^{*+} \rightarrow Cl_3P - SH_2^+ \leftarrow Cl_3P^{*+} + H_2S$. As a model for Cl₃P+SMe₂+ we optimized the Cl₃P+SH₂+ structure. The heterodimer radical cation Cl₃P+SH₂+ can be formed following the two different pathways indicated in the heading. The energetics have been envisaged and resulted for respectively the left or right side of the reaction in -16.5 and -15.0 kcal/mol, respectively (Table IV). The C_s staggered structure is computed to be a true minimum (see Figure 10), analogous to the H₃P-SH₂+ staggered C, structure.2c,d The stabilization is similar to or even slightly enhanced with respect to the latter one, depending on the theoretical level.

The calculated hyperfine couplings of the Cl₃P+SH₂+ radical are listed in Table V. In the C_s structure, atoms in the mirror plane contain the highest spin density. The elongation of the P-S σ^* bond is in accordance with its antibonding character. The directions of the SOMO contributions of phosphorus and sulfur are not aligned with the P-S bond direction. The orientation of

the anisotropic phosphorus and sulfur contributions to the SOMO (respectively 27.8° and 5.0°, Figure 10) are similar to the angular deviation noted earlier for their respective homodimers Cl₃P+PCl₃⁴

and $H_2S+SH_2^+$ (vide supra). **TBP-e** Cl_4P^* . Geometry optimization within $C_{2\nu}$ symmetry constraint results in a true minimum for Cl₄P^{*} (TBP-e, Figure 9). The hyperfine interactions are listed in Table V. In the TBP-e structure major spin density is primarily located on the axial chlorine atoms and the central phosphorus atom. In Cl₄P^{*} phosphorus is formally hypervalent, and consequently the bonds to the axial chorine atoms are elongated (Figure 9). Analogous to previous calculations on C_{2v} H₄P* and F₄P* 16 structures, the Cl_1 -P- Cl_1 bond angle in the C_{2v} structure is less than 180°. The direction of the largest principal value of the axial chlorine atoms is inclined to the P-Cl₁ bond direction by 21.4° and nearly orthogonal to the largest ³¹P hyperfine interaction (83.2°, Figure

Attempts To Calculate the Structure "X". Based on the experimental hyperfine coupling, the structure of the unassigned radical "X" seems similar to the TBP-e Cl₄P structure. Geometry optimization of Cl_4P^* within $C_{3\nu}$ symmetry constraint results in a stationary point (TBP-a, Figure 9). Unfortunately, vibrational analysis shows that it possesses a degenerate imaginary vibrational mode (118i cm⁻¹, E). The small value of the force constant of the imaginary mode encountered indicates that the local hypersurface along the direction of this vibrational mode is quite flat. Attempts to minimize the energy of a Cl₄P* structure within C, symmetry constraint or without symmetry (C_1) result in dissociation into Cl₃P and a Cl^{*} atom. Similarly, attempts to optimize the C_{3v} σ^* Cl_3P -Cl structure lead to such a dissociation. For the alternative conformation within C_{3v} symmetry, the TBP-a structure, maximal spin density is located at the equatorial chlorine atoms and the central phosphorus atom. The calculated ³¹P hyperfine coupling is larger than the interaction calculated for the $C_{2\nu}$ structure (Table V). This is in accordance with previous calculations¹⁶ and could possibly explain the experimental value of "X".

Since both TBP-e Cl₄P* and "X" radicals are formed on annealing by a rearrangement of the homodimer Cl₃P+PCl₃+ cation, we calculated several isomeric structures of Cl₃P+PCl₃+ by rearranging a chlorine atom from one phosphorus atom to the other. Only two stationary points were found, and their optimized geometries are depicted in Figure 10.

The first is a C_{2v} structure, composed of a TBP-e Cl₄P* molecular fragment and a Cl₂P⁺ residue. No vibrational analysis could be performed because of computer limitations. The P.--P distance of 3.75 Å clearly shows that the molecular fragments are loosely bound. The total energy of the C_{2v} structure has increased by 44.3 kcal/mol relative to optimized C_{2h} Cl₃P-PCl₃+.

The second stationary point found has been optimized within the C_s symmetry constraint. The resulting structure is composed of two Cl₃P fragments (Figure 10) and represents an intermediate prior to dissociation into a Cl₃P^{*+} radical and a Cl₃P molecule. Although the molecules are 3.22 Å apart, there is still mutual influence on the spin-density distribution. The chlorine atom of the Cl₃P molecule bears some spin density and likewise, the spin density on the central phosphorus atom of the Cl₃P⁺ radical cation is increased. The total energy lies 6 kcal/mol below the sum of the total energies of the respective $C_{3\nu}$ Cl₃P^{*+} and Cl₃P structures (Table IV).

Attempts to compute other stable geometries within C_s symmetry constraint failed because of SCF-convergence problems. $Cl_3P + H_2S - SH_2^+ \rightarrow Cl_3P(SH_2)_2^{\bullet+}$. As model compound for Cl₃P(SMe₂)₂*+ the structure of the Cl₃P(SH₂)₂*+ radical cation was optimized. Geometry optimization of the heterotrimer Cl₃P(SH₂)₂* resulted in four stationary points (Figure 11, Table IV). We performed vibrational analyses on the $C_{2\nu}(1)$ and C_s structures. Both resulted in two small imaginary frequencies, indicating no true minima $(C_{2\nu}(1), 92i \text{ cm}^{-1} \text{ B}_2, 52i \text{ cm}^{-1} \text{ B}_1; C_s, 89i \text{ cm}^{-1} \text{ and } 29i \text{ cm}^{-1} \text{ A''})$. The small size of the negative eigenvalue $C_{2\nu}(1)$ genvalues again indicates that the hypersurface for the corresponding vibrational modes is flat.

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Figure 11. Optimized geometries of the four stationary points found for the Cl₃P(SH₂)₂*+ structure. All structures calculated at the UHF/3-21G* SCF level.

< SPC12C11 = 85.6°

The experimental $\text{Cl}_3P(\text{SMe}_2)_2^{\bullet +}$ radical structure is analogous to the $\text{Cl}_5P^{\bullet -}$ radical anion. In order to provide a comparison, we computed the structure of the $\text{Cl}_5P^{\bullet -}$ radical anion. Geometry optimization within C_{4v} symmetry constraint gives a true minimum (Figure 9, Table IV). The spin density is located at the equatorial chlorine atoms and the phosphorus atom, similar to the C_{4v} $F_5P^{\bullet -}$ radical anion 16c and C_{3v} Cl_4P^{\bullet} radical. Therefore, the P-Cl₂ bonds are longer than the P-Cl₁ bond. The SOMO on phosphorus points away from the P-Cl₁ bond. The calculated hyperfine couplings are listed in Table V.

The calculated Cl₃P(SH₂)₂*+ radical structures are indeed reminiscent of the Cl₅P^{*} radical anion (Figure 11) with exception of the C_s structure. In the latter the P-S bond length of 3.16 Å indicates an intermediate structure prior to dissociation into Cl₃P^{o+} and 2H₂S. Likewise, the calculated ³¹P hyperfine coupling deviates seriously from the experimental value. The other three structures $(C_{2\nu}(1), C_{2\nu}(2), \text{ and } C_2)$ result in large ³¹P hyperfine couplings as also found experimentally. However, they are not larger than the hyperfine interaction calculated for Cl₅P*. The total energies are relatively high. In fact, the formation of the $C_{2\nu}(1)$ structure, by combining a Cl₂P molecule with the H₂S+SH₂+ radical cation, is disfavored by 23.4 kcal/mol at the UHF level (23.7 kcal/mol ROHF, Table IV). It should be taken into account that the simplification made by taking H₂S instead of Me₂S as substituent does rule out the possible stabilization via hyperconjugation by the β -hydrogen atoms.

Discussion

It has been shown that X irradiation at low temperature of mixtures of Me₂S and Cl₃P leads to several ion-molecule adducts. The adducts are detected and identified by ESR spectroscopy. In addition, ab initio quantum chemical calculations were performed to provide further insight into the radical structures.

The radiogenic production of the Cl₂P*+ radical cation from Cl₃P is straightforward. Analogous to trialkylphosphine radical cations, 4,6 Cl₃P^{o+} reacts with its precursor to yield the homodimeric Cl₃P-PCl₃+ radical cation. Likewise for Me₂S, we observe the well-known monopositive cation Me₂S^{*+} and the corresponding homodimer Me₂S+SMe₂⁺. In mixtures of Cl₃P and Me₂S the heterodimer Cl₃P+SMe₂⁺ radical cation is formed, in spite of a significant difference in ionization potential of the two participating molecular fragments. Its formation indicates that, in frozen solution, ionization potential differences are less important for the stability of σ^* bonds. At high solute concentrations we detected an ion-molecule reaction between the homodimer Me₂S÷SMe₂+ and a molecule of Cl₃P. The adduct is assigned to a heterotrimer Cl₃P(SMe₂)₂*+ radical cation in which the central phosphorus atom bears five substituents. The configuration of this radical is reminiscent of the octahedral Cl₅P* radical anion. The experimental ³¹P hyperfine coupling of the heterotrimer exceeds the value^{15h} of Cl₅P⁻⁻ (Table III). In addition, two radical structures arise

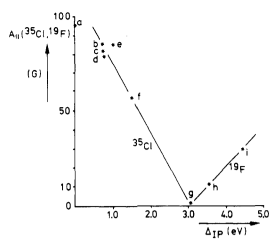


Figure 12. The parallel 35 Cl or 19 F superhyperfine couplings in σ^* ion-CFCl₃ adducts as a function of Δ_{IP} . (a) Cl₂FCCl*+, ref 1; (b) EtCl*+, ref 14b; (c) (MeO)₂P(H)O*+, ref 14d. (d) HCO₂Me*+, ref 14a; (e) (MeO)₃P=O*+, ref 14c; (f) EtBr*+, ref 14b. (g) Me₂S*+, ref 5a, and this study; (h) Me₂Se*+, ref 1; (i) Ph₃P*+, ref 6e.

from a rearrangement of the homodimer radical cation Cl_3P - PCl_3 ⁺. First, the TBP-e Cl_4P ⁺ radical can clearly be identified. Secondly, a possibly related radical structure can be observed ("X"). Until now no final assignment is made for "X". Candidates are a C_{3n} (TBP-a) Cl_4P ⁺ radical structure and a σ ^{*} Cl_3P -Cl radical.

Open-shell quantum chemical calculations on the expectation values of the isotropic and dipolar hyperfine coupling constants were performed on the optimized structures of the various monomer and dimer radical structures. The computed values are in general in qualitative agreement with the experimental data and confirm the assignments. Vibrational analyses show that the C_{2h} Cl₃P-PCl₃+ homodimer and the C_s heterodimer Cl₃P-SH₂+ are true minima. Calculations on the heterotrimer structure did not lead to a true minimum, although vibrational analysis reveals two small imaginary frequencies, indicating that the potential hypersurface for these vibrational modes is flat. It should be emphasized that substitution of Me₂S by H₂S in the calculations affects the electronic distribution within the radical due to the higher ionization potential of H_2S (IP = 10.4 eV). Likewise, it is well known that hyperconjugation of the methyl protons provides significant stabilization for such radical cation complexes despite long central bonds. Moreover, solvent interactions with radical structures can play a decisive role in stabilizing the radical configuration. However, such a stabilization is not yet amenable to computation. Efforts to find a satisfactory Cl₄P[•] structure for radical "X" did not yield a true minimum. For the C_{3n} (TBP-a) Cl₄P* structure a small degenerate negative eigenvalue is encountered. Geometry optimization of the $C_{3\nu}$ σ^* Cl_3P -Cl structure leads to a dissociation of the radical in Cl₃P and Cl^{*}.

Interestingly, we find that the extra CFCl₃ superhyperfine interactions on Me₂S^{*+} descend from both chlorine and fluorine of the CFCl₃ molecule(s). Usually such ion-molecule interaction is observed along the minimal g direction, 5a whereas it now coincides with the maximal g direction. Clark 1.2 suggested that the bond strength of a σ^* bond depends on Δ_{IP} . Likewise, Clark, Hasegawa, and Symons investigated the change in σ^* bond formation from chlorine to fluorine as a function of the difference in ionization potential between CFCl₃ and the dissolved molecule.¹ They predict that there should be a certain Δ_{IP} at which interaction switches from chlorine to fluorine. Indeed the Me₂S^{*+} radical cation seems to be this point ($\Delta_{IP} = 3.18 \text{ eV}$). This is illustrated in Figure 12, where the parallel superhyperfine coupling vs Δ_{IP} is displayed. Two straight lines can be drawn illustrating either superhyperfine interaction to either ³⁵Cl or ¹⁹F. However, there seems to be a paradox, since the ¹⁹F hyperfine coupling increases with increasing $\Delta_{\rm IP}$, i.e. with decreasing σ^* bond strength. Perhaps it is possible to rationalize this discrepancy by assuming a negative sign for the ¹⁹F hyperfine coupling. Yet we cannot explain the mechanism involved.

The type of matrix clearly influences radical formation. In the more mobile^{7h} matrices, CF₃CCl₃ and CF₂ClCFCl₂, the Me₂S' and Me₂S÷SMe₂+ radicals are observed simultaneously after X irradiation, whereas in the rigid, more crystalline, CFCl₃ or CH₂Cl₂ frozen solutions, they are detected separately depending on the concentration. Surprisingly, formation of the Cl₃P-PCl₃ homodimer is not observed in the mobile matrices, whereas in spite of ionization potential differences, the Cl₃P+SMe₂⁺ heterodimer is readily detected in CF₂ClCFCl₂ (but not in CF₃CCl₃!).

The exact mechanism controlling these phenomena is not clear. As a consequence of the absence of Cl₃P+PCl₃+ radical cation in the mobile matrices, the secondary TBP-e Cl4P and "X" radicals are also not detected in these frozen solvents. The experiments show that hyperfine couplings are fairly invariant to the type of freon. In CH₂Cl₂ the coupling for the same radical structure is usually slightly increased. It is noted that by following the reactions in different halocarbon solvents a consistent description

of radical formation and reactivity is obtained.

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Registry No. Cl₃P, 7719-12-2; Me₂S, 75-18-3; Cl₃P radical cation (1+), 136889-80-0; Me₂S radical cation (1+), 34480-65-4; Cl₃P dimer radical cation (1+), 136804-95-0; Me₂S dimer radical cation (1+), 76796-56-0; Cl₃P·Me₂S radical cation (1+), 136804-91-6; Cl₄P radical, 20762-59-8; Cl₃P·Me₂S·Me₂S radical cation (1+), 136804-92-7; Cl₃P· H_2S radical cation (1+), 136804-93-8; $Cl_3P \cdot H_2S \cdot H_2S$ radical cation (1+), 136804-94-9; H₂S radical cation (1+), 77544-69-5; H₂S, 7783-06-4; H₂S dimer radical cation (1+), 13604-96-1; Cl₅P, 58765-98-3; Cl₂P⁺, 75601-84-2; Cl₂P radical, 20762-58-7.

Electronic Structure and Aromaticity of 1,3-Azaphosphole and 1,3-Azarsole

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Three different aspects of the aromaticity have been studied for 1,3-azaphospholes and azarsoles, namely the molecular geometry, the bond separation reactions, and the UV photoelectron spectra. Ab initio molecular orbital theory has been used to calculate the geometrical and electronic structure of the molecules. Comparing the two possible tautomeric forms, the 1H derivatives (type I) have planar structures while the 3H derivatives (type II) show a puckered structure. The C-P or C-As bond lengths in the type I molecules are definitely shorter and the aromatic stabilization energies calculated from the bond separation reactions are larger than in type II molecules. Within each type I or type II molecules no characteristic difference in the aromaticity could be found. The band structure of the photoelectron spectra of type I molecules is similar to that of pyrrole, and the respective aromatic bands could be recognized.

The large variety of nitrogen-containing heteroaromatic systems inspired chemists dealing with organophosphorus compounds to synthesize heteroaromatic systems containing heavy atoms which replace the nitrogen. The achievements of the synthetic chemists in the 1980s have not been accompanied by widespread theoretical investigations. The photoelectron spectrum of the unstable parent compound phosphaethene1 has been observed, and numerous quantum-chemical calculations have been conducted on this Among the potentially aromatic metallocycles, phosphabenzene³ and phosphole⁴ have been studied. In our recent work, investigations of 1,3-benzazaphosphole and -benzoxaphosphole have been reported.⁵ The photoelectron spectra of methyl-substituted 1,2,4,3-triazaphosphole isomers have been studied, and on the basis of the change of some close-lying states the failure of Koopmans' theorem has been alleged.6

In a simple description of the electronic structure of nitrogen-containing heteroaromatic compounds two different types of nitrogen atoms are considered. For the first type the simplest example is pyrrole. The nitrogen atom in pyrrole is substituted by hydrogen that is easily removable by bases possessing acidic character to the parent molecule. In terms of MO theory the aromaticity and stability of pyrrole is the consequence of the interaction of the nitrogen lone pair with the cis-butadiene backbone forming a six-electron π -system. For the second type pyridine is the most common example. In this compound nitrogen possesses a basic character. In terms of MO theory this is a consequence of the free lone pair in the plane of the molecule.

The aromatic π -sextet can be formally constructed here from the interaction of a C=N double bond and the *cis*-butadiene systems. The least complex compound containing both types of nitrogens is imidazole. According to theoretical investigations the two different nitrogen atoms show similar conjugative ability. The partial charges and the linear coefficients in π -orbitals do not show significant differences between the two different types of nitrogen atoms of imidazole, benzimidazole, and perimidine.5,3

When replacing nitrogen by a heavier Group V/a element a diminished interaction is expected due to the small overlap between carbon 2p and phosphorus 3p (or arsenic 4p) orbitals. However, in a recent ab initio study,8 phosphabenzene was found as 88-90% aromatic as pyridine is, while phosphole is a nonplanar compound in contrast to pyrrole indicating that the aromatic stabilization of the planar state is insufficient to overcome the high inversion

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