in native laccase. The similar optical absorption spectra associated with the two intermediates (vide supra) despite the difference in their reduction state suggests that the absorption features are not directly associated with the oxygen intermediates per se but arise from a similarly perturbed trinuclear Cu site, i.e., with respect to Cu oxidation states or ligand binding mode.

Another difference between the two intermediates is that native but not T1Hg laccase exhibits a broad EPR signal observable at liquid He temperature. This is consistent with the extra reducing equivalent derived from the type 1 Cu(I) in native laccase. The signal is further broadened when reduced laccase is oxidized by ^{[17}O]dioxygen,^{10b} suggesting that it is associated with an oxygen radical, and it has been assigned to O^{•-} on the basis of its rapid electron spin relaxation, the assumed transfer of three electrons to dioxygen,⁸ and the uptake of two protons concomitant to its formation.³⁶ However, note that the significant divergence of the lowest g value of this signal (1.7) from 2.0 is not consistent with this assignment to an O⁻⁻ species, which would not have sufficient orbital angular momentum due to the low spin-orbit coupling constant of oxygen. Also it cannot arise from a magnetically isolated tetragonal Cu(II) ion,37 because a system with a $d_{x^2-y^2}$ ground state cannot exhibit g values below 2.0. However, it does show some similarity to the triplet state EPR spectra associated with a pair of dipolar-coupled S = 1/2 centers;³⁸ such signals can arise from Cu(II) dimers and have been observed upon N_3^- -induced uncoupling of the type 3 site in native⁴ and met T2D laccase.²⁴ Thus, although the absence of the EPR signal in T1Hg is consistent with the formulation of this intermediate as an even-electron species, the assignment of the intermediate EPR

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(37) MCD and CD studies of T1Hg laccase in the ligand field region indicate that the type 2 and type 3 sites in T1Hg have tetragonal effective geometries.²²
(20) Control of C

(38) Smith, T. D.; Pilbrow, J. R. Coord. Chem. Rev. 1974, 13, 173-278.

signal in native laccase to O^{•-} requires detailed spectral study.

In summary, it is clear that the type 2 Cu is required for dioxygen reactivity in laccase and that dioxygen reduction occurs in the absence of the type 1 Cu. This demonstrates that the type 2-type 3 trinuclear Cu site represents the active site for the binding and multielectron reduction of dioxygen. Previous observation that N_3^- can bind as a bridging ligand between the type 2 and type 3 sites⁴ suggests that a similar binding mode may be relevant to the reactivity of the trinuclear site with dioxygen. The type 1 Cu is clearly not necessary for reactivity with dioxygen, and in its absence, an intermediate is formed that shares some properties with the oxygen intermediate previously described in native laccase. Spectroscopic characterization of the oxygen intermediates in native and T1Hg laccase and X-ray edge determination of the associated Cu redox states should provide a detailed description of the mechanism of the irreversible multielectron reduction of dioxygen to water catalyzed by the multicopper oxidases.

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Heterocyclic 1,2,4,6-Thia- and 1,2,4,6-Selenatriazinyl Radicals. Spin Distributions and Modes of Association

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Abstract: Synthetic routes to the heterocyclic 1-chloro-1,2,4,6-thia- and -selenatriazines $[(Ph_2P)_{2-x}(PhC)_xN_3EC]$ (x = 0, 1; E = S, Se) have been developed. For x = 0 the method involves the coupling of $[ClPh_2PNPPh_2Cl]^+Cl^-$ with the N,N'-disilylated diimides Me₃SiNENSiMe₃ ($\dot{E} = S$, Se), while for x = 1 the preparation uses the condensation of (Me₃Si)₂NPN(Ph)CNSiMe₃ with either SCl₂ or SeCl₄. Reduction of the 1-chloro derivatives with triphenylantimony affords the thia- and selenatriazinyl radicals $[(Ph_2P)_{2-x}(PhC)_xN_3E]^{\bullet}$, all of which have been characterized by ESR spectroscopy. Hyperfine coupling constant data are interpreted in the light of MNDO calculations on model structures. X-ray structural analysis of the radical dimers [Ph2(tol)PCN3S]2 and [Ph3PCN3Se]2 CH3CN reveals significantly different modes of association; for the sulfur-based radical, association occurs through a long (average (range) = 2.489 (9) Å) sulfur-sulfur bond, while in the selenium species the dimer pair is coupled by a selenium-nitrogen linkage, the structural parameters within the two rings resembling those expected for a charge-transfer interaction.

Many of the recent advances in the chemistry of sulfur nitrogen compounds have stemmed from the study of radical species.²

Interest has focused on the use of such compounds in the design of synthetic metals,³ and also on how radical association can

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^{(1) (}a) University of Guelph. (b) University of Arkansas.





provide pathways for structural change.^{2,4,5} As part of our work on both topics, we have been studying the properties of heterocyclic radicals based on the 7π -electron 1,2,4,6-thiatriazinyl framework (Chart I).⁶⁻⁸ A variety of systems with similar topology, but with quite different electronic structures and chemistry, has emerged. The primary system 1 can been generated with a wide range of R groups^{6,9} and is stable to oxidation to a 6π -electron cation and reduction to an 8π -electron anion.¹⁰ ESR studies reveal a uniform spin distribution around the ring, with minimal spin leakage onto the exocyclic ligands. In the solid state, 1 (R = Ph) associates to the eclipsed dimer 2 in which the closest interannular contact is a long (2.666 Å) sulfur-sulfur "bond". Replacement of both endocyclic carbons in 1 by phosphorus affords the thiadiphosphatriazinyl system 3. Preliminary ESR data for 3 (R = Ph)suggest a heavily localized spin distribution, indicative of the dipolar formulation shown.8 Like 1, 3 forms a dimer in the solid state, but the structure of dimer 4 is not the simple cofacial arrangement typified by 2. Instead the cyclic framework of 3 is cleaved as two six-membered rings mutate into a single twelvemembered ring possessing a transannular sulfur-sulfur (2.385 (1) Å) contact.¹¹

The marked structural dichotomy represented by 2 and 4 has prompted us to explore the entire range of phosphorus-containing heterocyclic radicals based on the thiatriazinyl and selenatriazinyl frameworks. Our aim has been to investigate how variations in

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Table 1. Hyperfine Coupling Constants (mT) and g Values in Thiaand Selenatriazinyls

	$[Ph_2C_2N_3E]^{\bullet}$		[Ph ₃ PCN ₃ E] [•] ^a		$[Ph_4P_2N_3E]$	
E	S ^b	Se ^c	S	Se	S	Se
g	2.0059	2.0169	2.0064	2.0171	2.0058	2.0156
a(N2)	0.397	0.430	0.555	0.630	0.410	0.480
a(N4)	0.397	0.380	0.111	0.141	0.000	0.000
a(N6)	0.397	0.430	0.187	0.186	0.410	0.480
a(P)			0.420	0.544	0.050	0.259

^a Data obtained from 5-C₆D₅ derivative. ^b Data from ref 6. ^c Data from ref 13.

electronic structure (RC vs R₂P and S vs Se) and steric bulk can influence spin distributions and modes of association. In the present paper we provide a comprehensive account of the preparation and ESR analysis¹² of 1,2,4,6-thiatriazinyls 1, 3, and 5 (R = R' = Ph) and the corresponding 1,2,4,6-selenatriazinyls 6-8 (R = R' = Ph). Within this latter group only the purely carbon based system, i.e., 6 and its dimer 9, have hitherto been reported.¹³ The molecular structures of 10 (R = Ph, R' = tol) and 11 (R =R' = Ph), the radical dimers of 5 and 7, are also described.

Results and Discussion

Preparation of $R_2R'PCN_3ECl$ (E = S, Se). 1-Chloro-1,2,4,6thiatriazines can be prepared in several ways,² the most convenient being the condensation of an imidoylamidine, or its hydrochloride, with SCl₂.¹³ 1-Chloroselenatriazines can be obtained similarly with SeCl_{4}^{1} , although more forcing conditions are required to effect complete elimination of HCl.¹³ The corresponding thia- and selenatriazinyls 1 and 6 can then be generated by reduction, typically with triphenylantimony. Our proposed synthetic route to the mixed thia- and selenaphosphatriazine ring system was developed as an extension of this condensation approach, this time using the persilvlated phosphimidoylamidine 12 (eq 1). Ex-



periment fulfilled expectation; the reaction of 12 (R = R' = Ph; R = Ph, R' = tol) with SCl_2 provides the required 1-chloro-1,3,2,4,6-thiaphosphatriazine 13, while with SeCl₄ the corresponding 1-chloroselenaphosphatriazine 14 (R = R' = Ph) is produced. Both reactions proceed in high (>80%) yield.

Preparation of $R_4P_2N_3ECl$ (E = S, Se). 1-Chloro-1,3,5,2,4,6-thiadiphosphatriazines have been known for some time; the chloro (R = Cl) and phenyl (R = Ph) derivaties can both be prepared by the reaction of S_4N_4 with the appropriate R_2PCl compound.¹⁴ The method, however, cannot be readily extended to selenium chemistry.¹⁵ We have therefore sought a preparative route suitable for both $R_4P_2N_3ECl$ (15, E = S; 16, E = Se). This proved to be a nontrivial task; while sulfur dichloride can be condensed with both $[NH_2(Ph_2)PN(Ph_2)PNH_2]^+Cl^-$ and Me₃SiNPN(H)PNSiMe₃¹⁶ to afford Ph₄P₂N₃SCl in good yield, similar condensations with SeCl₄ fail to give Ph₄P₂N₃SeCl.¹⁷

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Chimia 1988, 42, 70. (16) This compound ($\delta(P) = 11.3$) can be made in situ by the reaction of Ph₂PN(H)PPh₂ with trimethylsilyl azide in toluene at 100 °C. (17) [NH₂(Ph₂)PNP(Ph₂)NH₂]⁺Cl⁻ condenses with a variety of transition-metal halides (e.g., WCl₆) to give the heterocyclic systems (e.g., Ph₄P₂N₃WCl₃); see: Katti, K. V.; Roesky, H. W.; Rietzel, M. *Inorg. Chem.* 1987, 26, 4032.

Eventually we pursued the use of the well-known N,N'-bis(trimethylsilyl)sulfur diimide (Me₃SiNSNSiMe₃ (17, E = S)) and its recently reported¹⁸ selenium analogue Me₃SiNSeNSiMe₃ (17, E = Se) as coupling reagents with the bis(chlorodiphenylphosphine)iminium salt [ClPh₂PNPPh₂Cl]⁺Cl⁻ (eq 2). This



approach proved to be surprisingly effective for both the sulfur and selenium systems 15 and 16, the latter preparation working in spite of the thermal instability of Me₃SiNSeNSiMe₃ and the consequent need to use (impure) material prepared in situ.¹⁹ In view of the success of this particular reaction, it is interesting to speculate to what extent the extensive chemistry of Me₃SiNSNSiMe₃ can be replicated by Me₃SiNSeNSiMe₃.

Generation and ESR Spectra of Radicals. All of the thia- and selenatriazines 13-16 described above can be reduced by triphenylantimony in oxygen-free methylene chloride to give strong and persistent ESR signals of the correponding thia- and selenatriazinyls 3, 5, 7, and 8^{20} There are some systematic variations in signal intensity, g value, and line width as a function of electronic structure. Thus, replacement of carbon by phosphorus and of sulfur by selenium both lead to weaker ESR signals, presumably as a result of stronger radical/radical interactions. In an earlier paper we assessed this trend quantitatively for 1 and $6^{.13}$ Replacement of sulfur by selenium also has the predictable effect of increasing line widths and g values (Table I).

Initial attempts to analyze the ESR spectra of $[Ph_3PCN_3E]^*$ 5 and 7 (R = R' = Ph) were hindered by slight spin leakage onto the C-phenyl group, which led to additional hyperfine splitting of an already complex spectrum.^{12,21} In order to simplify the spectrum, we prepared (starting from C₆D₅CN, see Experimental Section) samples of 5 and 7 in which the 5-phenyl group (R') was fully deuteriated. The ESR spectra of these deuteriated derivatives are shown in Figure 1. Hyperfine coupling constants (Table I) to phosphorus *a*(P) and nitrogen *a*(N) were obtained by full spectral simulation. Analysis of the ESR spectra of $[Ph_4P_2N_3E]^*$ 3 and 8 (Figure 1) was more straightforward. The presence of a molecular plane of symmetry and the apparent lack of coupling to N4 reduced the simulation problem to only two variables; the hyperfine coupling data provided in Table I for these two radicals were nonetheless confirmed by spectral simulation.

Spin and Charge Distributions in $[(Ph_2P)_{2-x}(PhC)_xN_3E]^*$ (E = S, Se; x = 0-2). ESR hyperfine coupling constant data for both the sulfur (1, 3, 5) and selenium (6-8) based radicals are compiled in Table I. Interpretation of the data in terms of variations in electronic structure is facilitated by comparison with spin densities (ρ) and net ($\sigma + \pi$) atomic charges (q) obtained from restricted Hartree-Fock (RHF) MNDO calculations on a series of model thiatriazinyl radicals (R = R' = H). The results (Figure 2), particularly the differences in the charge densities at carbon and phosphorus, illustrate how the delocalized [H₂C₂N₃S]* radical evolves to the highly polarized zwitterionic structure of [H₄P₂N₃S]*. Consistent with the internal salt description for the latter species, the a(N) values are of the same magnitude as those found in simple acyclic sulfur diimide radical anions [RNSNR]*-.22



Figure 1. ESR spectra of 5 and 7 ($R = C_6H_5$, $R' = C_6D_5$) and 3 and 8 ($R = C_6H_5$).

In the hybrid system $[H_3PCN_3S]^*$ both the spin densities and atomic charges suggest a saltlike formulation, with spin density increasing along the nitrogen sites from N4 to N6 to N2; indeed, given the limited capability of the MNDO method to model hypervalent phosphorus, the calculated spin densities concur remarkably well with the observed nitrogen hyperfine coupling constants, a(N). The high spin densities at sulfur are likewise reflected in the deviation of the g values from the free-electron value. The calculated spin density at carbon in $[H_3PCN_3S]^*$ is significantly greater than in $[H_2C_2N_3S]^*$, thus accounting for the more extensive spin leakage onto the C-phenyl groups of 5 and 7.

We have been unable, within the confines of the MNDO method, to examine in detail the electronic effects of replacement of sulfur by selenium. The ESR data can nonetheless be interpreted qualitatively with simple perturbation arguments based on the expected differences in electronegativity of sulfur and selenium. In short, sulfur appears to be a better acceptor of spin density than selenium; as a result, the coupling constants a(N)

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⁽²⁰⁾ The selenium-based radical 8 is the least stable. Solutions of this radical slowly deposit a red precipitate of amorphous selenium. Attempts to isolate a dimer of this species were unsuccessful.

⁽²¹⁾ Proton ENDOR spectroscopy and ESR simulation experiments for 5, $R' = C_6H_5$, revealed coupling to three exocyclic protons with a(H) = 0.035 mT.

⁽²²⁾ Hunter, J. A.; King, B.; Lindsell, W. E.; Neish, M. A. J. Chem. Soc., Dalton Trans. 1980, 880.



Figure 2. MNDO π -molecular orbital energies and distributions for 1, 5, and 3 (R = R' = H). Hyperconjugative out-of-phase PH₂ σ -combinations are not shown. Spin (ρ) and total ($\sigma + \pi$) charge (q) densities shown below.

for nitrogens adjacent to selenium are raised relative to the a(N) values found in the corresponding sulfur-based radicals. The effect is least apparent in delocalized radicals 1 and 6, as expected.

The relatively small a(P) values observed are typical of those found for four-coordinate phosphorus-containing radicals in which the phosphorus center represents a phosphonium cation rather than a neutral phosphoranyl unit.²³ For radicals such as **18**, which



bear a close structural resemblance to those reported here, it is well recognized that the value of a(P) can be influenced by several factors, including (i) spin polarization in the P-N σ -bond, (ii) π -spin density at phosphorus, and (iii) torsional motion about the C-N bond leading to hyperconjugation with the P-N σ -bond. The



Figure 3. ORTEP drawing of one (of two unique) molecule of 10 (R = C_6H_5 , R' = C_6D_5) showing the atom-numbering scheme.



Figure 4. ORTEP drawing of 11 ($R = R' = C_6H_5$) showing the atomnumbering scheme.

significant variation in the a(P) value in the present systems is most probably a result of changes in the degree of hyperconjugation rather than polarization or π -effects; in essence, the value of a(P) can be viewed as a measure of the distortion from planarity of the heterocyclic ring. Consistently, the a(P) value found in the puckered eight-membered heterocyclic radical 19, in which spin polarization and π -effects are expected to be similar to 3, is much larger (2.45 mT) than those of any of the six-membered radicals.²⁴

In summary, the extent of spin delocalization in heterocyclic thia- and selenatriazinyl radicals is restricted to the CNS backbone. Conjugation to include phosphorus is severely limited, the observed spin distributions giving little direct evidence of a genuine π -effect.²⁵

Solid-State Structures of $[Ph_2(tol)PCN_3S]_2$ and $[Ph_3PCN_3Se]_2$ ·CH₃CN. Attempts to obtain crystallographic data from the dimer of $[Ph_3PCN_3S]$ ·(5, R = R' = Ph) were fruitless; the compound was amorphous to X-rays. We therefore prepared (see the Experimental Section) a modified derivative with a tolyl group rather than a phenyl group at the 5-position. This slight change afforded a crystalline dimer, the structure of which has been determined by X-ray diffraction. Compound 10 crystallizes in the triclinic space group PI with two dimers per asymmetric unit. The conformational and structural features of these two dimers, however, are very similar, and for present purposes we restrict our discussion to the average of these units. A compilation of pertinent bond lengths and angles is provided in Table II. Structural data for the other dimer are deposited with the sup-

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Table 11.	Selected Bond	l Lengths	(Å) and a	Angles (deg)	with Esd's

[Ph ₂ (tol)PC	N ₃ S] ₂	[Ph ₃ PCN ₃ Se] ₂ ·CH ₃ CN						
Distances								
S(1)-S(2)	2.481 (3)	Se(1) - N(4)	1.990 (13)					
S(1) - N(1)	1.611 (5)	Se(1)-N(1)	1.736 (15)					
S(1) - N(3)	1.606 (6)	Se(1) - N(3)	1.743 (15)					
S(2) - N(4)	1.600 (5)	Se(2) - N(4)	1.888 (13)					
S(2) - N(6)	1.588 (6)	Se(2) - N(6)	1.83 (2)					
P(1) - N(1)	1.614 (5)	P(1) - N(1)	1.628 (13)					
P(1)-N(2)	1.611 (6)	P(1) - N(2)	1.594 (15)					
P(2)-N(4)	1.633 (6)	P(2)-N(4)	1.654 (14)					
P(2) - N(5)	1.604 (6)	P(2) - N(5)	1.565 (15)					
N(2)-C(1)	1.32 (2)	N(2)-C(1)	1.340 (8)					
N(3)-C(1)	1.36 (2)	N(3)-C(1)	1.331 (8)					
N(5)-C(2)	1.36 (2)	N(5)-C(2)	1.322 (8)					
N(6)-C(2)	1.31 (2)	N(6)-C(2)	1.348 (8)					
	Α	ngles						
S(1)-S(1)-N(1)	102.82 (3)	N(4)-Se(1)-N(1)	101.7 (7)					
S(2)-S(1)-N(3)	96.15 (3)	N(4)-Se(1)-N(2)	89.4 (7)					
S(1)-S(2)-N(4)	102.50 (3)	Se(1)-N(4)-P(2)	118.7 (8)					
S(1)-S(2)-N(6)	96.75 (3)	Se(1)-N(4)-Se(2)	118.4 (7)					
N(1)-S(1)-N(3)	114.4 (3)	N(1)-Se(1)-N(3)	111.5 (7)					
N(4)-S(2)-N(6)	115.4 (3)	N(4)-Se(2)-N(6)	103.4 (6)					
N(1)-P(1)-N(2)	115.8 (3)	N(1)-P(1)-N(2)	117.5 (9)					
N(4)-P(2)-N(5)	115.4 (3)	N(4)-P(2)-N(5)	115.7 (8)					
S(1)-N(1)-P(1)	114.9 (3)	Se(1)-N(1)-P(1)	112.9 (8)					
P(1)-N(2)-C(1)	120.1 (5)	P(1)-N(2)-C(1)	123 (1)					
S(1)-N(3)-C(1)	122.1 (5)	Se(1)-N(3)-C(1)	117 (1)					
S(2)-N(4)-P(2)	115.7 (3)	Se(2)-N(4)-P(2)	103.6 (7)					
P(2)-N(5)-C(2)	121.4 (5)	P(2)-N(5)-C(2)	124 (1)					
S(2)-N(6)-C(2)	122.4 (5)	Se(2)-N(6)-C(2)	115 (1)					
N(2)-C(1)-N(3)	131 (2)	N(2)-C(1)-N(3)	128.9 (7)					
N(5)-C(2)-N(6)	131 (2)	N(5)-C(2)-N(6)	128.3 (6)					

plementary material. The structure of 11, the dimer of $[Ph_3PCN_3Se]^{\bullet}$ (as its CH₃CN solvate), has also been determined, and pertinent positional and structural data for this compound are available in Table II. ORTEP drawings of the dimer units in $[Ph_2(tol)PCN_3S]_2$ (10) and $[Ph_3PCN_3Se]_2 CH_3CN$ (11) are shown in Figures 3 and 4.

In the sulfur-based dimer 10, the mode of association, i.e., through sulfur, is similar to that observed for 2,⁶ although the interannular S-S contact (the average of S1-S2 and S3-S4 is 2.489 Å) is somewhat shorter (2.666 Å in 2). Also in contrast to 2 the two rings are not eclipsed. Instead, and probably to alleviate steric interactions caused by the presence of the phenyl groups on phosphorus, one ring rotates with respect to the other around the S-S contact to make an NSSN torsion angle of 84° (average of four values). Within the two PCN₃S rings the bond lengths and angles are comparable and similar to those found in the parent chloro derivative 10.¹²

The structure of the selenium-based dimer 11 is markedly different, both from its sulfur analogue described above and dimer 9,¹³ which adopts a cofacial Se-Se structure similar to 2 with d(Se-Se) = 2.792 (3) Å. In 11 there is no formal Se-Se bond; instead the two rings are linked by a Se-N bond (Sel-N4 = (1.990 (13) Å). The mutual disposition of the rings, illustrated in Figure 4, is such that the four-atom sequence Se1-N4-Se2-N3 is almost coplanar (to within 0.06 Å). Within this sequence, the remaining interannular contact (Se2-N3) is 3.106 (15) Å. Both rings show marked deviations from planarity. Within the Sebonding ring, the five-atom N-C-N-P-N sequence is planar to within 0.05 Å, with the selenium atom lying 0.410 Å out of this plane (toward N4). In the N-bonded ring the Se-N-C-N-P unit forms a near-planar sequence (to within 0.12 Å), with N4 tipped 0.82 (1) Å away (toward Se1). The geometry at the three-cooordinate nitrogen N4, i.e., \sum (bond angles) = 340.7°, deviates substantially from sp² hybridization, but the extent of puckering falls short of that expected for a pure sp³ configuration.

The two PCN₃Se rings in 11 have totally different structural features. The Se-N bonds of the N-bonded ring are much longer than in the Se-bonded ring, and the C=N(Se) bond is much shorter. Taken collectively, the endocyclic structural parameters are broadly consistent with electron transfer from one PCN₃Se

Scheme 1



ring (Se-bonded) to the other (N-bonded); instead of the dimer unit being two loosely connected radicals, it more nearly resembles a cation/anion pair. In molecular orbital terms the gross structural features can be rationalized in terms of a simple model in which the SOMO of one ring is oxidized (to give a 6π -cation) while that of the other is reduced (to give an 8π -anion). In valence bond parlance, the resonance formulations shown in Scheme I reflect the observed endocyclic bond length variations surprisingly well. Similar structural differences have been observed between the 6π -cation **20** and the 8π -imide **21** (the disproportionation products of **1**).¹⁰



The above electronic description of the selenium-based dimer as a cation/anion pair allows some qualitative insight into the reason for the structural dichotomy observed for 10 and 11. As illustrated in Scheme I, the energetics of heteronuclear association of two radicals (5 or 7) can be viewed as the net result of two steps, (i) the disproportionation of two radicals, i.e., the sum of the ionization potential (IP) and electron affinity (EA), and (ii) the subsequent formation of a donor-acceptor (E-N) bond between the cation and anion, here denoted by $\Delta H(E-N)$. This process must then be compared with the simple homonuclear association enthalpy $\Delta H(E-E)$. The strong ESR signals observed for both 5 and 7 indicate that $\Delta H(E-E)$ and [$\Delta H(E-N) + IP + EA$] are small.

While the overall balance between the two modes of association clearly depends on several variables, the easiest to specify, and certainly the largest, is the IP of the radical.²⁶ On the basis of the ESR evidence outlined above, which suggests that selenium is slightly more electropositive than sulfur, we anticipate that 7 should have a slightly lower IP than 5. The preference for the Se-N dimerization mode for 7 can then be viewed as a result of a more favorable (less endothermic) disproportionation energy. Increasing the IP of 7, through judicious modification of the exocyclic R groups, may allow isolation of a Se-Se-coupled system. Reciprocally, decreasing the IP of 5 may favor a charge-transfer (S-N) dimer. Experiments to test this hypothesis are in progress.

⁽²⁶⁾ MNDO calculations on 1 (R = H)¹⁰ indicate that the IP is the major contributor to the disproportionation energy. Measurements on the dissociation constants for 1 and 6¹³ suggest that, entropy effects notwithstanding, the Se-Se bond is very slightly stronger than the S-S bond.



Figure 5. Qualitative potential energy hypersurface for Se-N-bonded dimers of 7.

Radical Association and Rearrangement. One of the major objectives of our ongoing studies of thia- and selenatriazinyl radicals has been to map out the relationships between electronic structure and mode of association. Our initial efforts were prompted in part by the early report of the structure of the dimer of 3, which revealed the possibility of association with rearrangement (to 4).¹¹ The result provided a direct analogy with the scrambling reaction of sulfur diimide radical anions (eq 3)^{2,5}



and further substantiated the description of 3 as an internal salt of a radical anion. In the case of the acyclic radical anion scrambling, the position of the equilibrium is determined by the nature of the groups R and R', while the expansion 3 to 4 is most probably driven by a release of ring strain.

In the present work we have sought to isolate and characterize dimers of the novel radicals 5, 7, and 8. Our initial hope was to find a radical dimer whose structure mimicked to some degree the putative head-to-tail four-center transition state in eq 3. While the instability of the [Ph₄P₂N₃Se][•] radical (with respect to decomposition to selenium) and the consequent difficulties in isolating a radical dimer for this species precluded completion of the desired "hat-trick", characterization of the two dimers 10 and 11 has successfully demonstrated how sensitive the preferred mode of binding for two radical units can be to minor electronic changes. In the context of the rearrangement shown in eq 3, the selenium dimer can be viewed as an example of a severely distorted fourcenter interaction between two radical anions. Qualitative consideration of the global potential energy hypersurface available to the undistorted arrangement 22 (Figure 5) reveals, however, several other distorted Se-N-bonded structures, i.e., 23 and 24. The preference for observed structure 25 can be understood in terms of the charge-transfer model developed above. Accordingly, the most stable coupling modes should be those that involve the most negatively charged nitrogen (N2) on the anion, i.e., 23 and 25 as opposed to $24.^{27}$ Secondary interactions in 23 and 25 between a (donor) selenium of the anion ring can take place to two different (acceptor) nitrogens of the cation, N6 in 25 and N2 in 23. The preference for 25 over 23 stems from the relative (electrostatic) acidities of these two sites; calculated MNDO charge densities on a model $[H_3PCN_3S]^+$ cation²⁸ confirm that N6 (q = -0.45) rather than N2 (q = -0.56) is the better acceptor.

Experimental Section

Starting Materials and General Procedures.²⁹ Benzonitrile (Aldrich), benzonitrile- d_5 (MSD), tolunitrile (Aldrich), chlorodiphenylphosphine (Aldrich), butyllithium (1.6 M in hexane, Aldrich), trimethylsilyl azide (Aldrich), hexamethyldisilazane (Aldrich), selenium powder (Alfa), sulfur dichloride (Aldrich), chlorine gas (Matheson), thionyl chloride (Aldrich), triphenylantimony (Aldrich), and selenium oxychloride (Alfa) were all commercial products; all were used as received, save thionyl and sulfur dichloride, both of which were freshly distilled before use. Acetonitrile (Fisher HPLC grade) was also freshly distilled from P_2O_5 before use. LiN(SiMe)2. Et2O, 30 Me3SiNSNSiMe3, 31 and Ph2PN(H)PPh232 were prepared according to standard literature methods. The N, N, N'tris(trimethylsilyl)phosphimidoylamidine [(Me₃Si)N(Ph₂)PN(Ph)CN- $(SiMe_3)_2$, $\delta(P) = -3.6$ (lit. -4.2)] was prepared as recently described,³³ i.e., by the reaction of benzonitrile with LiN(SiMe₃)₂ with PhCN and treatment of the so-formed amidine with Ph2PCl to produce Ph2PN-(Ph)CN(SiMe₃)₂, followed by in situ reaction of the latter intermediate with Me₃SiN₃. The product was crystallized from acetonitrile. Deuteriation of the phenyl group (at carbon) was achieved with benzonitrile-d₅. The corresponding C-tolylphosphimidoylamidine [(Me₃Si)N- $(Ph_2)PN(tol)CN(SiMe_3)_2$, mp = 95 °C, $\delta(P) = -4.0$] was made in an analogous fashion from tolunitrile. Anal. Calcd for $C_{29}H_{44}N_3PSi_3$: C, 63.34; H, 8.06; N, 7.64. Found: C, 63.32; H, 8.09; 7.80. [ClPh₂PNPPh₂Cl]⁺Cl⁻ was prepared by direct oxidation of Ph₂PN(H)-PPh₂ with chlorine; ³¹P NMR datum ($\delta(P) = 44.4$) of the product so obtained was in agreement with the literature $(\delta(P) = 44.3)$.³⁴ The selenium diimide Me₃SiNSeNSiMe₃ was prepared from SeOCl₂ and $LiN(SiMe_3)_2^{18}$ and used in situ. All reactions and experimental manipulations were performed under an atmosphere of nitrogen or argon. NMR spectra were recorded (in CDCl₃ reference H₃PO₄ unless otherwise indicated) on a Bruker WH-400 spectrometer, and infrared spectra (1600–250-cm⁻¹ region, Cs1 optics) were obtained on a Nicolet 20SX/CFT1R instrument. Chemical analyses were performed by MHW Laboratories, Phoenix, AZ.

Preparation of Ph₃PCN₃SCl (13, $\mathbf{R} = \mathbf{R}' = \mathbf{Ph}$). An excess (2 mL) of sulfur dichloride in 20 mL of CH₃CN was added slowly to a stirred solution of (Me₃Si)₂N(Ph₂)PN(Ph)CNSiMe₃ (1.4 g, 2.6 mmol) in 40 mL of acetonitrile. In the course of the addition the reaction mixture turned dark red and then golden yellow. The mixture was heated at reflux for 1 h, cooled to room temperature, and filtered. The filtrate was concentrated in vacuo to a volume of ca. 20 mL and cooled to -25 °C overnight. Subsequent filtration of this mixture afforded bright yellow blocks of Ph₃PCN₃SCl (0.81 g, 2.1 mmol, 81%) that were recrystallized from acetonitrile: dec > 157 °C; ³¹P NMR δ (P) = 10.9; 1R 1494 (w), 1438 (w), 1318 (s), 1256 (m), 1179 (m), 1119 (m), 1035 (m), 1022 (m), 749 (w), 730 (s), 703 (s), 682 (s), 543 (s), 488 (w), 377 (w) cm⁻¹. Anal. Calcd for C19H15N3PSCI: C, 59.46; H, 3.94; N, 10.95; Cl, 9.24. Found: C, 59.26; H, 4.06; N, 10.70; Cl, 9.09. A sample of this material, fully deuteriated at the 5-position, was made on a small scale from (Me₃Si)₂N(Ph₂)PN(C₆D₅)CNSiMe₃ and SCl₂

Preparation of $Ph_2(tol)PCN_3SCl$ (13, R = Ph, R' = tol). This compound was made (in 69% yield) with a procedure identical with that described above for Ph3PCN3SC1 from the reaction of SCl3 and (Me₃Si)N(Ph₂)PN(tol)CN(SiMe₃)₂. Ph₂(tol)PCN₃SCl forms pale yellow crystalline blocks from acetonitrile: mp 165-70 °C; ³¹P NMR $\delta(P)$ = 10.3; 1R 1436 (m), 1405 (w), 1314 (m), 1296 (m), 1186 (m), 1125 (sh), 1115 (m), 1035 (m), 1016 (m), 934 (w), 882 (w), 814 (m), 740 (s), 692 (m), 540 (s), 520 (m), 490 (m), 446 (w), 397 (m) cm⁻¹. Anal. Calcd for C₂₀H₁₇N₃PSCI: C, 60.38; H, 4.31; N, 10.56; Cl, 8.91. Found: C, 60.11; H, 4.30; N, 10.56; Cl, 9.11.

Preparation of Ph_3PCN_3SeCl (14, R = R' = Ph). Selenium tetrachloride was prepared in situ by passing chlorine gas over a slurry of selenium powder (0.59 g, 7.5 mmol) in 35 mL of acetonitrile. A solution

⁽²⁷⁾ Structure 23 is directly accessible by distortion of the centrosymmetric four-center interaction 22. Generation of the other two structures, 24 and 25, requires that one ring of 22 be inverted with respect to the other.

⁽²⁸⁾ In the absence of MNDO parameters for selenium, the calculation was performed on the sulfur-containing cation.

⁽²⁹⁾ The health hazards of the selenium compounds prepared in this work are not known. Caution should be exercised in handling these potentially toxic materials

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of $(Me_3Si)_2N(Ph_2)PN(Ph)CNSiMe_3$ (4.0 g, 7.5 mmol) in 15 mL of acetonitrile was then added dropwise to the previously generated slurry of SeCl₄; the SeCl₄ rapidly dissolved to give a bright yellow solution. When the addition was complete, the mixture was heated at reflux for 30 min and then cooled to -25 °C overnight. The canary yellow blocks of Ph₃PCN₃SeCl (2.7 g, 6.3 mmol, 85%) so obtained were filtered and recrytallized from acetonitrile: mp 147-50 °C; ³¹P NMR $\delta(P) = 12.0$; 1R 1590 (w), 1438 (w), 1423 (m), 1323 (s), 1284 (m), 1778 (m), 1129 (s), 1115 (s), 1068 (w), 1026 (w), 947 (s), 783 (w), 755 (w), 724 (s), 705 (s), 690 (s), 528 (s), 445 (w), 353 (w), 322 (w) cm⁻¹. Anal. Calcd for C₁₉H₁₅N₃PSeCl: C, 52.98; H, 3.51; N, 9.76; Cl, 8.23. Found: C, 53.05; H 3.65: N, 9.85; Cl, 8.16

H, 3.65; N, 9.85; Cl, 8.16. **Preparation of Ph₄P₂N₃SCl (15, R = R' = Ph).** A solution of Me₃SiNSNSiMe₃ (1.3 g, 6.3 mmol) in 10 mL of acetonitrile was added dropwise to a cold (0 °C) solution of [ClPh₂PNPPh₂Cl]⁺Cl (3.0 g, 6.1 mmol) in 25 mL of acetonitrile. The mixture was stirred for 30 min at room temperature, then concentrated in vacuo to ca. 25 mL, and cooled to -25 °C overnight. Subsequent filtration affored pale yellow blocks of Ph₄P₂N₃SCl (1.4 g, 2.9 mmol, 48%), the identity of which was confirmed by comparison of its infrared spectrum¹⁴ and ³¹P chemical shift (δ (P) = 8.1, lit.¹⁴ 7.8) with the literature data.

Preparation of Ph₄P₂N₃SeCl (16, R = Ph). A solution of crude Me₃SiNSeNSiMe₃, prepared in situ by the reaction of SeOCl₂ (4.2 g, 25.5 mmol) with LiN(SiMe₃)₂ (8.0 g, 47.9 mmol) in 50 mL of diethyl ester, was dissolved (after removal of the ether) in 75 mL of CH₃CN. To this solution was added solid [ClPh₂PNPPh₂Cl]⁺Cl⁻ (7.0 g, 14.3 mmol). The mixture was stirred at room temperature for 30 min, then filtered, and concentrated in vacuo to a volume of ca. 40 mL, and the filtrate was cooled to -5 °C overnight. The large pale yellow blocks of Ph₄P₂N₃SeCl (3.0 g, 5.7 mmol, 40% based on phosphorus) so obtained were recrystallized from acetonitrile: dec > 143 °C; ³¹P NMR δ (P) = 12.5;¹⁹ IR 1437 (7), 199 (m), 1172 (w), 1036 (m), 1025 (m), 910 (s), 722 (s), 691 (s), 674 (m), 584 (m), 539 (s), 516 (s), 423 (m), 331 (w) cm⁻¹. Anal. Calcd for C₂₄H₂₀N₃P₂SeCl: C, 54.72; H, 3.83; N, 7.98; Cl, 6.73. Found: C, 54.62; H, 4.08; N, 7.83; Cl, 6.56.

Preparation of [Ph₃PCN₃S]₂ (10, R = R' = Ph), Solid triphenylantimony (0.25 g, 0.71 mmol) was added to a hot (70 °C) solution of Ph₃PCN₃SCl (0.50 g, 1.3 mmol) in 10 mL of triply degassed (freezepump-thaw) acetonitrile. The solution was slowly cooled to room temperature, and the bright red glassy nodules of [Ph₃PCN₃S]₂ (0.31 g, 0.45 mmol, 68%) were separated by filtration: dec > 90 °C; 1R 1491 (m), 1435 (w), 1338 (s), 1322 (s), 1295 9s), 1180 (m), 1123 (s), 1112 (m), 853 (s), 926 (m), 884 (m), 817 (m), 743 (s), 724 (s), 686 (s), 533 (s), 528 (s) cm⁻¹. Anal. Calcd for $C_{38}H_{30}N_6P_2S_2$: C, 65.51; H, 4.34; N, 12.06. Found: C, 65.29; H, 4.54; N, 11.84.

Preparation of [Ph₂(tol)PCN₃S]₂ (10, R = Ph, R' = tol). Solid triphenylantimony (0.38 g, 1.1 mmol) was added to hot (70 °C) solution of Ph₂(tol)PCN₃SCl (0.75 g, 1.9 mmol) in 50 mL of acetonitrile. The solution was slowly cooled, and the resultant bright red crystals of [Ph₂(tol)PCN₃S]₂ (0.28 g, 0.4 mmol, 41%), separated by filtration: dec > 153 °C; IR 1421 (m), 1338 (s), 1315 (m), 1293 (m), 1184 (w), 1122 (m), 1115 (m), 977 (s), 926 (m), 862 (w), 811 (w), 737 (m), 722 (w), 693 (m), 535 (m), 523 (m) cm⁻¹. Anal. Calcd for C₄₀H₃₄N₆P₂S₂: C, 66.28; H, 4.73; N, 11.59. Found: C, 65.40; H, 4.89; N, 11.81.

66.28; H, 4.73; N, 11.59. Found: C, 65.40; H, 4.89; N, 11.81. **Preparation of [Ph₃PCN₃Se]₂ (11, \mathbf{R} = \mathbf{R}' = \mathbf{Ph}).** Solid triphenylantimony (0.45 g, 1.3 mmol) was added to hot (70 °C) solution of Ph₃PCN₃SeCl (1.0 g, 2.3 mmol) in 25 mL of triply degassed (freezepump-thaw) acetonitrile. The deep red solution so formed was then filtered (to remove a trace of red selenium) and allowed to cool slowly to room temperature. The resulting deep red crystals of [Ph₃PCN₃Se]₂·CH₃CN (0.56 g, 0.71 mmol, 62%) were isolated by filtration: dec > 109 °C; IR 1436 (w), 1424 (m), 1319 (s), 1293 (m), 1177 (m), 1120 (m), 1118 (m), 1067 (w), 1024 (w), 946 (m), 880 (m), 819 (w), 753 (w), 724 (s), 700 (s), 692 (s), 612 (w), 597 (s), 525 (s), 517 (s) cm⁻¹. Anal. Calcd for C₄₀H₃₃N₇P₂Se₂: C, 57.77; H, 4.00; N, 11.79. Found: C, 57.49; H, 4.10; N, 11.79. **Preparation of** $[Ph_4P_2N_3S]_2$ (4, R = Ph). Solid triphenylantimony (0.52 g, 1.5 mmol) was added to a hot (70 °C) solution of $Ph_4P_2N_3SCI$ (0.70 g, 1.5 mmol) in 30 mL of triply degassed (freeze-pump-thaw) CH₃CN. The orange solution was left to stand overnight, and the resulting orange crystals of $[Ph_4P_2N_3S]_2$ (0.33 g, 0.37 mmol, 51%) were isolated by filtration. The identity of this material, dec > 210 °C, was confirmed by comparison with the known IR spectrum¹¹ and ³¹P chemical shift ($\delta(P) = 15.8$; lit.¹¹ 15.8). Anal. Calcd for C₄₈H₄₀P₄N₆S₂: C, 64.86; H, 4.54; N, 9.45; S, 7.21. Found: C, 64.76; H, 4.66; N, 9.57; S, 7.13.

H, 4.54; N, 9.45; S, 7.21. Found: C, 64.76; H, 4.66; N, 9.57; S, 7.13. Attempted Preparation of $[Ph_4P_2N_3Se]_2$. Repeated attempts to generate a radical dimer of $[Ph_4P_2N_3Se]^*$ by reduction of $Ph_4P_2N_3SeCl$ with Ph_3Sb with a procedure analogous to that described above for $[Ph_4P_2N_3S]_2$ were unsuccessful. Such reactions afforded pale yellow solutions (from which ESR data were obtained) that slowly deposited a red precipitate of amorphous selenium.

ESR Spectra. All ESR spectra were recorded on samples in degassed methylene chloride at ambient temperature with a Varian E109 ESR spectrometer. Samples were prepared either by in situ reduction of the corresponding chloro derivative with triphenylantimony or by dissolving the radical dimer. Spectral simulations, from which coupling constant data were extracted, were performed with the program ESR42, written by Dr. U. M. Oehler, University of Guelph.³⁵

Molecular Orbital Calculations. Restricted open-shell Hartree-Fock (RHF) MNDO molecular orbital calculations were performed with the MOPAC suite of programs³⁶ operating on a SUN-3 workstation. Geometries of all model structures were optimized with a minimum requirement of C_s symmetry (i.e., ring planarity).

X-ray Structure Determinations. Crystals of $[Ph_2(tol)PCN_3S]_2$ and [Ph₃PCN₃Se]₂·CH₃CN suitable for X-ray work were grown from acetonitrile. All X-ray data were collected on an Enraf-Nonius CAD-4 at 293 K with monochromated Mo K_{α} ($\lambda = 0.71073$ Å) radiation. Crystals were mounted on glass fibers coated with epoxy. Data were collected by using a $\theta/2\theta$ technique with a scan width of $(1.0 + 0.3 \tan \theta)$. Data collection, structure solution, and refinement parameters for both structures are available as supplementary material. Both structures were solved with MULTAN and refined by full-matrix least squares that minimized $\Sigma w(\Delta F)_2$. In the selenium-based dimer the solvent molecule was best modeled by five partially occupied positions centered in crystal cavities at 0.0, 0.5, 0.0 and 0.3, 0.5, 0.0. The paucity of data with I > $3\sigma(I)$ for the sulfur dimer is apparently due to lack of long-range order; the number of reflections with $I > 3\sigma(I)$ dropped off rapidly at 19° in θ and comprised only 12% of the reflections scanned in the shell of θ from 21 to 22°. No extinction correction was applied to either structure. Atomic scattering factors and anomalous dispersion corrections were taken from ref 37. All programs used were those provided by the Enraf-Nonius Structure Determination Package.

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Supplementary Material Available: Tables of crystal data, structure solution, and refinement (S1), atomic coordinates for $[Ph_2(tol)PCN_3S]_2$ (S2) and $[Ph_3PCN_3Se]_2$ ·CH₃CN (S3), bond lengths and angles (S4, S5), and anisotropic thermal parameters (S6, S7) for both structures 10 and 11 (26 pages). Ordering information is given on any current masthead page.

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