

80697-72-9; **4b**, 80697-73-0; **4c**, 80697-74-1; **4d**, 80719-01-3; **4e**, 80697-75-2; **5a**, 67583-11-3; **5b**, 80697-76-3; **5c**, 80052-14-8; **5d**, 69306-31-6; **6a**, 80697-77-4; **6b**, 67551-66-0; **6c**, 80697-78-5; **6d**, 80719-02-4; **6e**, 67670-43-3; **6f**, 80697-79-6; **6g**, 80719-03-5; Fe^{III}.

[TPP][Cl], 16456-81-8; **3a**, 133-07-3; **3b**, 133-06-2; Fe^{III}[T(*p*-Cl)-PP][Cl], 36965-70-5; Fe^{III}[TTP][Cl], 19496-18-5; Fe^{III}[OEP][Cl], 28755-93-3; *n*-C₄H₉NH₂, 109-73-9; Fe[TPP][CN-*n*-C₄H₉][NH₂-*n*-C₄H₉], 80719-68-2.

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Synthesis and Characterization of Some Ruthenium-Phosphoniodithiocarboxylate Complexes

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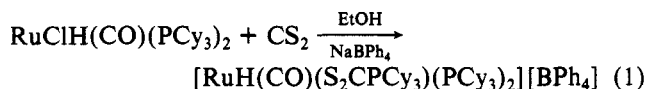
Received August 10, 1981

Addition of CS₂ to RuClH(CO)(PCy₃)₂ affords the cation [RuH(CO)(S₂CPCy₃)(PCy₃)₂]⁺, which has been isolated as the tetraphenylborate salt. The closely related complex [RuCl(CO)(S₂CPCy₃)(PCy₃)₂][BPh₄] is formed when the zwitterion ligand S₂CPCy₃ is added to a methanol suspension of RuCl₂(CO)(PCy₃)₂ and NaBPh₄. The reaction of carbonyl sulfide with RuClH(CO)(PCy₃)₂ results in the formation of RuClH(CO)₂(PCy₃)₂.

Carbon disulfide is known to insert into metal-hydride bonds to give metal dithioformates.¹⁻⁵ Recently it has become apparent that metal-phosphoniodithiocarboxylate complexes, M(S₂CPR₃)L_n, may be formed from the addition of CS₂ to metal-phosphine complexes.⁶⁻⁸ We report the syntheses of several ruthenium-phosphoniodithiocarboxylate complexes and one reaction in which formation of a phosphoniodithiocarboxylate ligand is favored over formation of a dithioformate ligand when a polar solvent is employed.

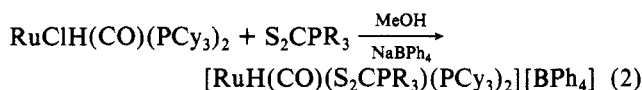
Results and Discussion

While CS₂ inserts into the RuH bond of RuClH(CO)(PCy₃)₂ (Cy = cyclohexyl) to afford RuCl(S₂CH)(CO)(PCy₃)₂,² we find that in a polar solvent (ethanol) a different reaction occurs. When CS₂ is added to an ethanol suspension of RuClH(CO)(PCy₃)₂, the yellow-orange solid dissolves and a purple solution is formed. Addition of NaBPh₄ to the solution precipitates [RuH(CO)(S₂CPCy₃)(PCy₃)₂][BPh₄] (eq 1). The yield of the salt is low, as expected since some of the RuClH(CO)(PCy₃)₂ starting material must serve as a source of PCy₃.



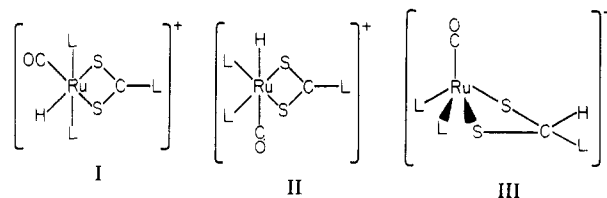
Coordination of CS₂ and subsequent transfer of a PCy₃ ligand to the carbon atom of CS₂ could lead to the formation of the phosphoniodithiocarboxylate ligand. Alternatively, phosphine dissociation could lead to the formation of the zwitterion adduct S₂CPR₃, which could then react with RuClH(CO)(PCy₃)₂ to give the insertion product. We have found that direct addition of a zwitterion adduct, S₂CPR₃ (R = Cy, Et), to RuClH(CO)(PCy₃)₂ results in the facile for-

mation of the cation [RuH(CO)(S₂CPR₃)(PCy₃)₂]⁺, which we have isolated as the tetraphenylborate salt (R = Cy, **1**; R = Et, **2**), as shown in eq 2. Although the formation of the



phosphoniodithiocarboxylate ligand could be regarded as an insertion of CS₂ into a RuP bond, we believe from (2) that it is more likely that reaction 1 proceeds via disproportionation.

The infrared spectra of **1** and **2** exhibit one terminal carbonyl stretching vibration and a band between 1050 and 950 cm⁻¹ that we suggest is ν(CS) of the S₂CPR₃ ligand⁷ (Table I). Each complex also exhibits a very weak band at ~2000 cm⁻¹ that may be attributed to ν(Ru-H), but the low intensity of this absorption precludes a definite assignment. The complexes exhibit several strong bands between 750 and 700 cm⁻¹ that might be attributed to ν(CS₂)_{sym}.⁵ However, these bands are apparently not observed in other phosphoniodithiocarboxylate complexes. The ³¹P{¹H} NMR spectra of **1** and **2** consist of an A₂X pattern, consistent with the presence of two magnetically equivalent and one magnetically inequivalent PR₃ group. The small values of the coupling constants (see Table I) are indicative of long-range coupling.^{6,7} Three isomers that should exhibit similar spectra are



Although phosphonium-betaine ligands (isomer III) are formed when CS₂ is added to similar metal complexes,^{7,9} the ¹H NMR spectra are consistent only with isomers I and II as the spectra exhibit a hydride resonance (Figure 1) that is split into a triplet by two equivalent PR₃ ligands and further split into a doublet by a more distant PR₃ group. The betaine proton of isomer III would be expected to appear further downfield^{7,9} (δ ≈ 6) and should couple more strongly to the

- (1) Moers, F. G.; ten Hoedt, R. W. M.; Langhout, J. P. *Inorg. Chem.* **1973**, *12*, 2196-2198.
- (2) Moers, F. G.; ten Hoedt, R. W. M.; Langhout, J. P. *J. Organomet. Chem.* **1974**, *65*, 93-98.
- (3) Robinson, S. D.; Sahajpal, A. *Inorg. Chem.* **1977**, *16*, 2718-2722.
- (4) Yaneff, P. V. *Coord. Chem. Rev.* **1977**, *23*, 183-220 and references therein.
- (5) Butler, I. S.; Fenster, A. E. *J. Organomet. Chem.* **1974**, *66*, 161-194.
- (6) Armit, P. W.; Sime, W. J.; Stephenson, T. A.; Scott, L. J. *J. Organomet. Chem.* **1978**, *161*, 391-406.
- (7) Werner, H.; Bertleff, W. *Chem. Ber.* **1980**, *113*, 267-273.
- (8) Clark, G. R.; Collins, T. J.; James, S. M.; Roper, W. R.; Town, K. G. *J. Chem. Soc., Chem. Commun.* **1976**, 475-476.
- (9) Ashworth, T. V.; Singleton, E.; Laing, M. *J. Chem. Soc., Chem. Commun.* **1976**, 875-876.

Table I. Infrared and NMR Data of the Complexes

compd	$^1\text{H NMR},^a \delta$		$^{31}\text{P NMR}^a$	IR absn ^b
	hydride	alkyl		
$[\text{RuH}(\text{CO})(\text{S}_2\text{CPCy}_3)_2(\text{PCy}_3)_2][\text{BPh}_4]$ (1)	-10.75 (dt, $^2J(\text{PH}) = 22.0$, $^4J(\text{PH}) = 6.1$)	1.88 (s) 1.48 (s) 1.24 (s)	66.3 (d) 22.8 (t, $^4J(\text{PP}) = 8.6$)	1998 (vw) $\nu(\text{RuH})?$ 1938 (vs) $\nu(\text{CO})$ 964 (m) $\nu(\text{CS})$ 741 (vs) 730 (vs) $\nu(\text{CS}_2)_{\text{sym}}?$ 703 (vs)
$[\text{RuH}(\text{CO})(\text{S}_2\text{CPEt}_3)(\text{PCy}_3)_2][\text{BPh}_4]$ (2)	-10.74 (dt, $^2J(\text{PH}) = 21.1$, $^4J(\text{PH}) = 6.1$)	1.82 (s) 1.50 (s) 1.18 (s)	63.9 (d) 28.7 (t, $^4J(\text{PP}) = 8.5$)	2010 (vw) $\nu(\text{RuH})?$ 1940 (vs) $\nu(\text{CO})$ 1043 (s) $\nu(\text{CS})$ 730 (s) $\nu(\text{CS}_2)_{\text{sym}}?$ 706 (s)
$[\text{RuCl}(\text{CO})(\text{S}_2\text{CPCy}_3)(\text{PCy}_3)_2][\text{BPh}_4]$ (3)		1.79 (s) 1.48 (s) 1.24 (s)	40.9 (d) 25.2 (t, $^4J(\text{PP}) = 5.7$)	1953 (vs) $\nu(\text{CO})$ 949 (m) $\nu(\text{CS})$ 738 (vs) 728 (vs) $\nu(\text{CS}_2)_{\text{sym}}?$ 705 (vs)
$\text{RuClH}(\text{CO})(\text{PCy}_3)_2$	-24.7 (t, $^2J(\text{PH}) = 18.3$)		45.5 (s)	1908 (vs) $\nu(\text{CO})$
$\text{RuClH}(\text{CO})_2(\text{PCy}_3)_2$	-5.3 (t, $^2J(\text{PH}) = 24.0$)		49.9 (s)	2030 (vs) $\nu(\text{CO})$ 1978 (m) $\nu(\text{RuH})$ 1945 (vs) $\nu(\text{CO})$
$\text{RuCl}_2(\text{CO})(\text{PCy}_3)_2$			34.5 (s)	1934 (vs) $\nu(\text{CO})$

^a In CDCl_3 . J values given in Hz. ^b In Nujol. Phosphine bands are not tabulated.

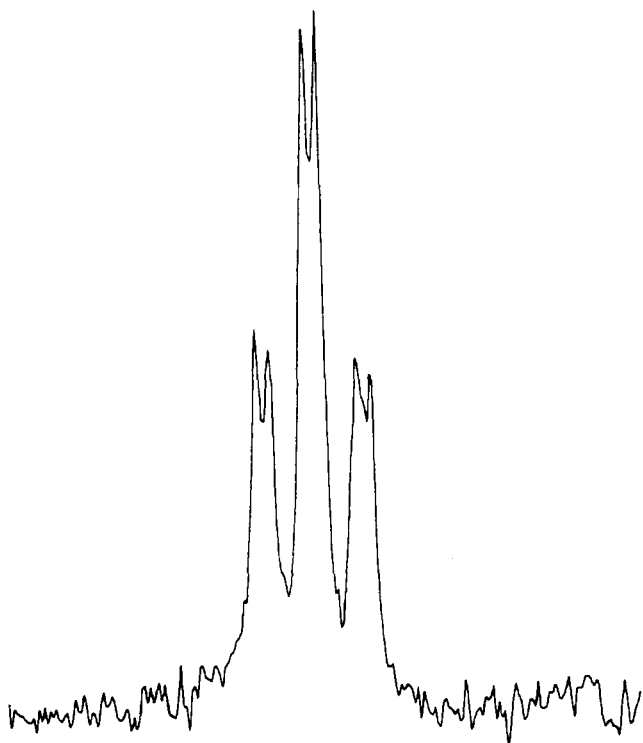


Figure 1. $^1\text{H NMR}$ spectrum of the hydride region of $[\text{RuH}(\text{CO})(\text{S}_2\text{CPCy}_3)(\text{PCy}_3)_2][\text{BPh}_4]$. The pattern is centered at $\delta -10.75$; the marker is 50 Hz wide.

betaine PR_3 group than to the PR_3 ligands.

The P-P coupling constant from PR_3 to S_2CPR_3 should be larger for isomer II, in which the S_2CPR_3 ligand is trans to the PR_3 ligands, than for isomer I, in which the S_2CPR_3 ligand is cis to the PR_3 ligands. For a similar complex, $[\text{RuCl}(\text{MeOH})(\text{S}_2\text{CPEt}_2\text{Ph})(\text{PEt}_2\text{Ph})_2][\text{BPh}_4]$, which is known to have PEt_2Ph ligands cis and trans to the $\text{S}_2\text{CPEt}_2\text{Ph}$ ligand,⁶ the coupling constants vary by a factor of 3 ($^4J(\text{cis PP}) = 3.5$ Hz, $^4J(\text{trans PP}) = 11.3$ Hz). The values of 4J for 1 and 2 are intermediate between these values, and we are unable to distinguish isomer I from II on the basis of the magnitude of 4J .

An analogue of 1, $[\text{RuCl}(\text{CO})(\text{S}_2\text{CPCy}_3)(\text{PCy}_3)_2][\text{BPh}_4]$ (3), may be prepared by the addition of S_2CPCy_3 to a meth-

anol suspension of $\text{RuCl}_2(\text{CO})(\text{PCy}_3)_2$ and NaBPh_4 . The ^{31}P and $^1\text{H NMR}$ spectra of 3 are similar to the spectra of 1 and 2, except that the hydride resonance is absent (see Table I). The analogous isomers (I, II) of 3 would have a sulfur atom or a carbonyl ligand trans to the chloro ligand, respectively, and the metal-chloride stretching frequency ($\nu(\text{RuCl}) = 274$ cm^{-1}) is consistent with either.^{2,10} Thus, we are unable to distinguish isomer I from isomer II on the basis of the metal-chlorine stretching frequency. The value of 4J for 3 is slightly smaller than for 1 or 2 and is nearer to $^4J(\text{cis PP})$ rather than $^4J(\text{trans PP})$ for $[\text{RuCl}(\text{MeOH})(\text{S}_2\text{CPEt}_2\text{Ph})(\text{PEt}_2\text{Ph})_2][\text{BPh}_4]$.⁶ Therefore, we propose that isomer I of 3 is formed. Isomer I should be favored over II for 1, 2, and 3 because of the sterically unfavorable cis arrangement of PCy_3 ligands in II.

Carbonyl sulfide reacts with $\text{RuClH}(\text{CO})(\text{PCy}_3)_2$ in polar and nonpolar solvents to afford $\text{RuClH}(\text{CO})_2(\text{PCy}_3)_2$. The fate of the sulfur in the reaction has not been determined, but we find no NMR spectroscopic evidence for the formation of SPCy_3 , even if free PCy_3 is present. It is not surprising that carbonylation, rather than insertion to give $[\text{RuH}(\text{CO})(\text{SOCPCy}_3)(\text{PCy}_3)_2]\text{Cl}$, occurs. Unlike CS_2 , COS does not form a stable zwitterion complex with PCy_3 , and metal-promoted $\text{C}=\text{S}$ bond cleavage has been shown to be more facile for COS than for CS_2 .¹¹

Experimental Section

$\text{RuCl}_2(\text{CO})(\text{PCy}_3)_2$ ² and $\text{RuClH}(\text{CO})(\text{PCy}_3)_2$ ¹² were prepared as previously described. Carbonyl sulfide was obtained from the Matheson Gas Co., East Rutherford, NJ. Infrared spectra were recorded on a Perkin-Elmer 283 spectrometer. Proton and ^{31}P NMR spectra were obtained on a JEOL FX90Q spectrometer. Peak positions are relative to tetramethylsilane and 85% phosphoric acid, respectively, with downfield values reported as positive. Elemental analyses were performed by Galbraith Laboratories, Inc., Knoxville, TN. All reactions were performed in air, unless otherwise noted.

$[\text{RuH}(\text{CO})(\text{S}_2\text{CPCy}_3)(\text{PCy}_3)_2][\text{BPh}_4]$ (1). Method A. Carbon disulfide (7 mL) was added to a suspension of $\text{RuClH}(\text{CO})(\text{PCy}_3)_2$ (0.100 g in 20 mL of methanol). Sodium tetraphenylborate (0.5 g) was added to the purple solution. The solution was concentrated by

(10) Lupin, M. S.; Shaw, B. L. *J. Chem. Soc. A* 1968, 741-749.

(11) Ibers, J. A.; Gaffney, T. R.; Schramm, K. D. "Coordination Chemistry—21" (IUPAC); Pergamon Press: Oxford and New York, 1981; pp 141-149.

(12) Moers, F. G.; Langhout, J. P. *Recl. Trav. Chim. Pays-Bas* 1972, 91, 591-600.

rapid purging with N₂ while warming to ~40 °C. The purple suspension was cooled to -30 °C, and the dark red solids were collected by filtration and dried under vacuum. The crude yield was 0.076 g. The predominant product (~40% of the solid) is [RuH(CO)(S₂CPCy₃)(PCy₃)₂][BPh₄], as determined by comparison of ³¹P and ¹H NMR spectra with those of a pure sample, which was prepared by method B. Additional products, including a ruthenium hydride complex (δ -12.6 (t, J = 20.7 Hz)), were detected via NMR spectroscopy.

Method B. A suspension of RuClH(CO)(PCy₃)₂ (0.160 g) and S₂CPCy₃ (0.0784 g in 35 mL of absolute ethanol) was stirred for 20 min. The purple solution was filtered, and 0.30 g NaBPh₄ was added to the filtrate. The purple suspension was cooled to -20 °C, and the purple solid was collected by filtration, washed with ethanol, and dried under vacuum. The yield was 0.177 g (59%). Anal. Calcd for C₈₀H₁₂₀BOP₃RuS₂: C, 70.31; H, 8.85; P, 6.80; S, 4.70. Found: C, 70.57; H, 9.02; P, 6.49; S, 4.55.

[RuH(CO)(S₂CPEt₃)(PCy₃)₂][BPh₄] (2). A suspension of RuClH(CO)(PCy₃)₂ (0.135 g) and S₂CPEt₃ (0.054 g in 20 mL of anhydrous methanol) was stirred for 5 min. Carbon disulfide (2.0 mL) was then added to the suspension, and the clear purple solution was stirred for 5 min. Addition of 0.20 g of NaBPh₄ gave a purple precipitate which was collected by filtration, washed with cold methanol, and dried under vacuum. The yield was 0.188 g (84%). Anal. Calcd for C₆₈H₁₀₂BOP₃RuS₂: C, 67.81; H, 8.54; P, 7.71; S, 5.32. Found: C, 67.69; H, 8.37; P, 7.45; S, 5.50.

[RuCl(CO)(S₂CPCy₃)₂(PCy₃)₂][BPh₄] (3). A suspension of RuCl₂(CO)(PCy₃)₂ (0.080 g) and S₂CPCy₃ (0.24 g in 40 mL of deoxygenated methanol) was stirred for 0.5 h under nitrogen at 40 °C. The purple solution was cooled to 20 °C and filtered. Addition of 0.40 g of NaBPh₄ to the filtrate gave a red-brown solid. The solid was recrystallized three times from chloroform-methanol to give a dark red solid which analyzed as [RuCl(CO)(S₂CPCy₃)(PCy₃)₂][BPh₄].CHCl₃. The yield was 0.065 g (40%). Anal. Calcd for

C₈₁H₁₂₀BCl₄OP₃RuS₂: C, 63.98; H, 7.95; Cl, 9.33; P, 6.11; S, 4.22. Found: C, 63.10; H, 8.07; Cl, 8.27; P, 6.59; S, 4.21.

RuClH(CO)₂(PCy₃)₂. Method A. Carbonyl sulfide was bubbled through a solution of RuClH(CO)(PCy₃)₂ (0.102 g in 10 mL of deoxygenated toluene) until the color faded from yellow to colorless. The solvent was removed under vacuum, and the white solid which remained was recrystallized from CHCl₃-MeOH to give the complex as white crystals. The yield was 0.069 g (60%). The complex was identified by comparison of ¹H NMR and IR spectra with those of an authentic sample.¹³ The complex has previously been assigned a stereochemistry with trans PCy₃ ligands and cis CO ligands.¹³ See Table I for spectral data.

Method B. Carbonyl sulfide (~10 g) was condensed into a liquid-nitrogen-cooled pressure reactor which contained 0.054 g of RuClH(CO)(PCy₃)₂, 0.10 g of PCy₃, and 8 mL of deoxygenated methanol. The vessel was allowed to warm to room temperature (Caution! 12 atm). The metal complex dissolved to give an orange solution which faded to colorless and precipitated a white solid. The carbonyl sulfide was distilled off, and the white solid was collected by filtration and dried under vacuum. The complex was identified by comparison of ¹H NMR and IR spectra with those of an authentic sample.

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Registry No. 1, 80697-67-2; 2, 80697-69-4; 3, 80697-71-8; RuClH(CO)(PCy₃)₂, 40935-25-9; RuClH(CO)₂(PCy₃)₂, 55100-76-0; RuCl₂(CO)(PCy₃)₂, 52524-94-4; CS₂, 75-15-0.

(13) Moers, F. G.; ten Hoedt, R. W. M.; Langhout, J. P. *J. Inorg. Nucl. Chem.* 1974, 36, 2279-2282.

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Stabilization of RN=NN=PR₃. Preparation and Structural Characterization of Stable Tetraarylphosphazide Complexes Containing Molybdenum and Tungsten

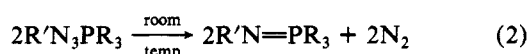
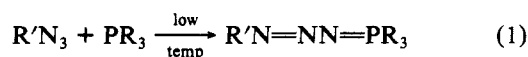
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Received October 15, 1981

The reaction of aromatic azides (R'N₃) with MBr₂(CO)₃(PPh₃)₂ (M = Mo, W; Ph = C₆H₅; tol = *p*-CH₃C₆H₄) in dry methylene chloride at 20 °C affords MBr₂(CO)₃(R'N₃PPh₃), N₂, and R'N=NN=PR₃ (R' = Ph, tol). The phosphazide complexes exhibit remarkable stability with respect to N₂ loss. In contrast to the Mo(II) and W(II) complexes, ReCl₃(CH₃CN)(PPh₃)₂ and ReCl₃(PPh₂Me)₃ yield ReCl₄(PR₃)₂ upon treatment with the same aryl azides. Triclinic needles of WBr₂(CO)₃(tolN₃PPh₃) were grown from chloroform-ether and crystallized in space group C₂h-P₁ with $Z = 2$, $a = 13.715$ (6) Å, $b = 9.904$ (5) Å, $c = 10.397$ (5) Å, $\alpha = 100.98$ (2)°, $\beta = 83.11$ (2)°, and $\gamma = 85.80$ (1)°. An X-ray diffraction study at -145 (5) °C showed the complex to be monomeric and seven-coordinate. The tolyl azide had inserted into the W-P bond, forming a phosphazide ligand (tolN₃PPh₃) which is bound to W in a chelating fashion through the α and γ nitrogen atoms; the N₃W metallacycle is nearly planar. Salient metrical parameters of the structure include the following: W-N(1) = 2.163 (4) Å, W-N(3) = 2.220 (5) Å, N(1)-N(2) = 1.279 (6) Å, N(2)-N(3) = 1.364 (6) Å, N(3)-P = 1.672 (5) Å, N(1)-C(41) = 1.423 (7) Å; N(1)-W-N(3) = 56.7 (2)°, W-N(1)-N(2) = 102.4 (3)°, W-N(3)-N(2) = 96.8 (3)°, N(1)-N(2)-N(3) = 103.8 (4)°. The full-matrix, least-squares refinement converged to $R(F) = 0.028$ and $R_w(F) = 0.037$ for 4066 unique data with $F_o^2 > 3\sigma(F_o^2)$.

Introduction

The reaction of tertiary phosphines with organic azides is known to proceed via a reactive intermediate, R'NNNPR₃, which decomposes in a bimolecular process to dinitrogen and the corresponding phosphoranamine¹ (eq 1 and 2). These



intermediates, which were originally named "phosphazides" by Staudinger,² are only rarely stable under ambient conditions; when R = R' = C₆H₅, the phosphazide decomposes rapidly at temperatures above -20 °C. Certain phosphazides

(1) Mosby, W. L.; Silva, M. L. *J. Chem. Soc.* 1965, 1003. Horner, L.; Gross, A. *Justus Liebigs Ann. Chem.* 1955, 591, 117.

(2) Staudinger, H.; Hauser, E. *Helv. Chim. Acta* 1921, 4, 861. Though unsystematic, the phosphazide nomenclature is easy to use and descriptive in its own right. Chemical Abstracts Service names RNNNPR₃ as a derivative of phosphoranylideneazide.