## **Reactivity Studies of Mononuclear and Dinuclear Vanadium-Sulfide-Thiolate Compounds**

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The synthesis, structure, and properties of  $(Me_3NBz)_{2}[V(S_2)S_2(SPh)]$  (1) and  $(Me_3NBz)_{4}[V_2(S_2)_{2}(CS_3)_{4}]$  (2) are described. Compound **1** is produced via oxidation of V(II1) by elemental sulfur in a reaction mixture comprising VCl<sub>3</sub>, LiSPh, S, and NEt<sub>4</sub>Cl in a molar ratio of 1:7:3:2. Compound 1 reacts with excess CS<sub>2</sub> to produce compound **2** in ca. **65%** yield. Compound **1** crystallizes in monoclinic space group *ml/n* with the following unit cell dimensions at  $-158$  °C:  $a = 18.178$  (8) A,  $b = 8.843$  (2) A,  $c = 18.852$  (7) A,  $\beta = 107.06$  (2)°, and  $Z = 4$ . A total of 2807 unique reflections with  $F > 3.00\sigma(F)$  were employed, and the structure solution was refined to conventional indices *R* and *R,* of **5.65** and **5.63,** respectively. The vanadium(V) center is ligated by three types of sulfur atoms, viz. one PhS<sup>-</sup>, one persulfido (S<sub>2</sub><sup>2-</sup>), and two sulfido (S<sup>2-</sup>); the coordination geometry is approximately tetrahedral, with the  $S_2^2$ - considered as occupying one site. Compound 2 crystallizes in monoclinic space group  $P_1/c$  with the following unit cell dimensions at  $-114$  °C:  $a = 20.978$  (4) Å,  $b = 13.384$  (2) Å,  $c = 21.613$  (4) Å,  $\beta = 93.14$  (1)°, and  $Z = 4$ . A total of 5651 unique reflections with  $F > 2.33\sigma(F)$  were employed, and the structure solution was refined to conventional *R* and  $R<sub>w</sub>$  indices of 6.32 and 6.75, respectively. The vanadium(IV) centers are bridged by two symmetry-related  $\eta^2:\eta^2:\mu_2-S_2^{2-}$  ligands, and two bidentate chelating trithiocarbonate ligands complete the coordination sphere about each metal. The metal centers are eight-coordinate, and the coordination geometry is best described as distorted dodecahedral. The V-V distances in the two independent molecules present in the unit cell are **2.872 (6)** and **2.841 (6)** A, respectively, and the complex is diamagnetic. *An* EHMO calculation was performed in order to assess the extent of metal-metal bonding in **2.** Comparisons with the more highly developed area of Mo/S chemistry are made. The terminal trithiocarbonate sulfur atoms of **2** are nucleophilic and can be alkylated with MeI to produce the neutral compound  $V_2(S_2)_2(S_2CSMe)_4(3)$  in high yield. Compound 3 reacts with secondary amines to form the dialkyldithiocarbamate compounds  $V_2(S_2)_2(S_2CNR_2)_4$  ( $R = Et (4a); R = Nu (4b)$ ). Solution-phase **IH** and 13C NMR studies of dimers 2-4 indicate the presence of two isomers whose interconversion is slow on the NMR time scale. Solution-phase s'V NMR data are reported for compounds **14** and are compared to literature values for other V/S complexes.

We have been studying vanadium-sulfur-thiolate chemistry for a number of years because of the relevance and importance of these compounds to a wide variety of chemical, industrial, and biological systems. These complexes may serve as models for the transformation of vanadyl  $(VO^{2+})$  impurities present in crude oils to polymeric vanadium sulfides (e.g.,  $V_2S_3$ ,  $V_3S_4$ ) under the reducing and sulfur-rich conditions of hydrodesulfurization.<sup>1</sup> The sulfides thus produced decrease the activity and lifetime of the heterogeneous catalyst employed (e.g., Mo/Co/S on alumina). Vanadium-sulfide-thiolate compounds can also **be** viewed as soluble, discrete structural counterparts of polymeric vanadium sulfide phases that are of interest for their magnetic and electric properties, catalytic activity, and use as electrodes in solid-state batteries.2 The availability of molecular compounds allows for characterization of the reactivity and other intrinsic properties of the basic building block of the extended lattice. Furthermore, **EXAFS** results obtained for a vanadium-containing nitrogenase

**Introduction suggest that the metal center is located in a complete or partial** sulfur environment, so the study of model compounds may help delineate biochemical structural aspects and reaction pathways.3 A more basic reason to study these compounds is to explore their reactivity characteristics and compare these results to the betterdeveloped area of molybdenum-sulfur chemistry.

> In recent reports, we have described the preparation and properties of a variety of discrete vanadium-sulfide-thiolate complexes of nuclearity  $1-4$  and oxidation level III-V.<sup>4</sup> With a number of compounds in hand, we have recently turned our attention **to** an exploration of the reactivity of these compounds with small organic molecules. A particularly attractivecandidate for reactivity studies is the vanadium(V) monomer  $(Me<sub>3</sub>NBz)<sub>2</sub>$ - $[V(S_2)S_2(SPh)]$ <sup>k</sup> In addition to a variety of nucleophilic sites, the presence of an easily oxidizable thiolate in conjunction with

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**a metal in its highest oxidation state suggests that this compound should display high reactivity. In fact, the compound is metastable**  in solution  $(t_{1/2}$  of ca. 4 h in MeCN) and reacts with a variety **of substances, including elemental S and Se, electron-deficient**  acetylenes, and heterocumulenes such as CS<sub>2</sub>. The production of a vanadium(IV) dimer from the reaction between  $[V(S_2)S_2]$ -**(SPh)] 2- and CS2 and subsequent reactivity studies on this dimer are the subject of the current report.** 

## **Experimental Seetion**

Syntheses. All manipulations were performed under a purified nitrogen atmosphere employing standard Schlenk techniques and a Vacuum Atmospheres glovebox. Tetrahydrofuran, diethyl ether, and hexanes were distilled from purple sodium benzophenone ketyl. Acetonitrile and chloroform were distilled from CaH<sub>2</sub>. Carbon disulfide, "Bu<sub>2</sub>NH, Et<sub>2</sub>-NH, MeI, and DMSO were stored over activated 4-A molecular sieves. Anhydrous VCI<sub>3</sub> was used as received (Aldrich). LiSPh and NaSPh were prepared by addition of the alkali metal toa THF solution containing a slight excess of PhSH; the white precipitate was collected by filtration, washed with Et<sub>2</sub>O, and dried under vacuum.

**A.**  $(Me_3NBz)gV(S_2)S_2(SPh)$  (1). This material was previously synthesized in 30-40% yield.<sup>4c</sup> An improved procedure uses LiSPh in place of NaSPh. Hence, to a stirred slurry of VCl<sub>3</sub> (1.57 g, 9.98 mmol), LiSPh (8.13 **g,** 70.0 mmol), and EkNCI (3.30 **g,** 19.9 mmol) in MeCN (250 mL) was added **S** (0.96 **g,** 30 mmol). The initial orange-brown solution quickly darkened upon **S** addition to a deep green coloration. After being stirred for 1 h, the mixture was filtered, leaving a small amount of purple precipitate and giving an intense green filtrate. Addition of Me<sub>3</sub>NBzCl (3.70 g, 19.9 mmol), followed by Et<sub>2</sub>O (ca. 30 mL) to the filtrate initiated crystallization of **1.** After overnight storage of the flask at-10 $\degree$ C, the black crystalline product was collected by filtration, washed with MeCN/Et<sub>2</sub>O (1:3), and dried under vacuum (yield  $4.13$  g,  $70\%$ based on V). IR (Nujol, cm<sup>-1</sup>): 1574 (m), 1312 (w), 1265 (w), 1215 (w), 1156 (w), 1078 (m), 1024 (m), 988 (w), 976 (m), 930 (w), 918 (w), 891 (m), 831 (w), 779 (m), 741 **(s),** 723 **(s),** 700 **(s,** sh), 694 **(s,** sh), 529 (vs), 496 (m), 453 (m), 421 (w). Electronic spectrum in MeCN,  $\lambda_{\text{max}}/ \text{nm}$  $(\epsilon_m/L \text{ mol}^{-1} \text{ cm}^{-1})$ : 206 (sh, 52 000), 230 (sh, 30 000), 296 (14 000). <sup>51</sup>V NMR (35.1 mM in 1:1 CD<sub>3</sub>CN/DMSO):  $\delta$  +970 ppm.

**B.**  $(Me_3NBz)(V_2(S_2)_2(CS_3)_4]$  (2). To a stirred slurry of 1 (0.40 g, 0.68 mmol) in MeCN (45 mL) was added  $CS_2$  (0.41 mL, 6.8 mmol) via syringe. The solution quickly turned deep red. After being stirred for **1** h, the mixture was filtered, leaving a small amount of brown powder and a deep **red** filtrate. Storage of the filtrate at ambient temperature for several days gave **2** as blackcrystals. Recrystallization can be effected from warm MeCN. Typical yields are 60-70%. Analysis of material dried under vacuum indicates the nonsolvated form. Anal. Calcd for CaHwN&V2: C, 41.81; H, **5.10;** N, 4.43; **S,** 40.59; V, 8.06. Found: C, 41.50; H, 5.28; N, 4.51; **S,** 41.15; V, 8.14. IR (Nujol, cm-I): 1559 (w), 1215 (w), 1157 (w), 1034 (w), 990 **(s), 885 (s),** 779 (m), 727 **(s),**  702 (m), 667 (w), 613 (w), 575 (w), 517 (m), 484 (w), 453 (w), 410 (w), 360 (m). Electronic spectrum in MeCN,  $\lambda_{max}$ , nm ( $\epsilon_{m}/V_2$ , L mol<sup>-1</sup> cm-I): 206 (91 OOO), 242 (39 OOO), 290 (34 OOO), 336 (54 OOO), 490 (6200). I3C NMR (iH-decoupled, DMSO-ds): 6 253.0, 252.8 ppm  $(CS_3)^2$ . <sup>51</sup>V NMR (8.8 mM in 1:7 CD<sub>3</sub>CN/DMSO):  $\delta$  +135 ppm.

**C.**  $V_2(S_2)_2(S_2CSMe)_4$  (3). To a stirred slurry of 2 (0.200 g, 0.158 mmol) in MeCN  $(50 \text{ mL})$  was added MeI $(60 \mu L, 0.96 \text{ mmol})$  via syringe. Over a period of 2 days, the solution became colorless and a brown precipitate was observed. This material was collected by filtration, washed with MeCN/Et<sub>2</sub>O (1:3), and dried under vacuum (yield 0.100 mg, 88%). Anal. Calcd for  $C_8H_{12}S_{16}V_2$ : C, 13.29; H, 1.67; V, 14.09. Found: C, 13.32; H, 1.71; V, 13.27. IR (Nujol, cm-I): 1407 (w), 1308 (w), 1159 (w), 1015 (m), 965 **(s),** 953 **(s),** 938 **(s),** 926 **(s),** 721 (m), 583 (m), 515 (m), 465 (w), 413 (w). Electronic spectrum in CHCl<sub>3</sub>, λ<sub>max</sub>, nm (ε<sub>M</sub>/V<sub>2</sub>, L mol<sup>-1</sup> cm<sup>-1</sup>): 286 (34 000), 318 (42 000), 480 (4300). <sup>1</sup>H NMR (CDCI,): **d** 2.662 **(s),** 2.658 ppm **(s). 51V** NMR **(8.8** mM in 1:7 CD<sub>3</sub>CN:DMSO):  $\delta$  +173 ppm.

**D.**  $V_2(S_2)_2(S_2CNR_2)_4$  ( $R = Et (4a)$ ). To a stirred slurry of 3 (0.400 **g,** 0.553 mmol) in THF **(50** mL) was added HNEt2 (0.37 mL, 3.6 mmol) via syringe. Over a period of ca. 18 h, the solution became homogeneous and dark red-brown. The solvent was removed, and the resulting dark brown material was washed with 3 **X 10** mL portions of hexane. Black crystals were obtained after recrystallization from THF/hexane at -10 °C (yield 225 mg, 50%). Anal. Calcd for  $C_{20}H_{40}N_4S_{12}V_2$ : C, 29.18; H,4.90;N,6.80;V,12.38. Found: **C.29.25;H,4.98;N,6.54V,12.68.**  IR (Nujol, cm-I): 1485 **(s),** 1426 **(s),** 1354 **(s),** 1306 (w), 1269 **(s),** 121 1 (m), 1140 (m), 1073 (w), 1038 (w), 1001 (w), 914 (w), 882 (w), 849 (m),

**779(m),723(m),577(~),554(m),498(~),438(~),349(m).** Electronic spectrum in THF,  $\lambda_{\text{max}}$ , nm ( $\epsilon_M/V_2$ , L mol<sup>-1</sup> cm<sup>-1</sup>): 246 (36 000), 278 **3.77(m,16H,CH2),1.24(t,12H,J=6.6Hz,CH3),1.16ppm(t,12H,**  43.99, 43.86 *(CHz),* 12.74, 12.45 ppm **(CH3).** 51V NMR (CDCI3): 6 +lo3 ppm. (32 OOO), 364 (9600), 478 (2OOO), 550 (1800). 'H NMR (CDCI3): 6  $J = 7.2$  Hz, CH<sub>3</sub>). <sup>13</sup>C NMR (<sup>1</sup>H-decoupled, CDCl<sub>3</sub>):  $\delta$  44.43, 44.35,

**E.**  $V_2(S_2)_2(S_2CNR_2)_4$  ( $R = Bu(4b)$ ). To a stirred slurry of 3 (0.340) **g,** 0.470 mmol) in THF (50 mL) was added "Bu2NH (2.4 mL, 14.2 mmol) via syringe. After 2 h, the solution became homogeneous and deep red. The solvent was removed, leaving a dark oil that was repeatedly washed with IO-mL portions of hexane until a brown powder was obtained. Recrystallization from THF/hexanes at -10 °C gave dark brown crystals (yield 210 mg, 43%). Anal. Calcd for  $C_{36}H_{72}N_4S_{12}V_2$ : C, 41.28; H, 6.93; N, 5.35. Found: C, 41.48; H, 7.19; N, 5.24. IR (Nujol, cm<sup>-1</sup>): 1491 **(s),** 1424 **(s),** 1366 **(s),** 1304 (m), 1289 (m), 1258 (w), 1219 **(s),**  1183 (w), 1142 (m), 1109 (w), 1092 (w), 1049 (w), 1013 (w), 974 (w), 951 (w), 930 (w), 909 (w), 779 (w), 723 **(s),** 604 (m), 586 (m), 559 (w), 515 (w), 436 (w), 377 (w), 364 (m). Electronic spectrum in THF,  $\lambda_{\text{max}}$ , nm (e<sub>M</sub>/V<sub>2</sub>, L mol<sup>-1</sup> cm<sup>-1</sup>): 242 (35 000), 288 (32 000), 370 (9600), 472 (2700), 560 (1500). <sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  3.66 (m, 16H, NCH<sub>2</sub>), 1.61 (m, 16H, NCH<sub>2</sub>CH<sub>2</sub>), 1.31 (m, 16H, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>), 0.956 (t, 6H, *J*  $= 7.33$  Hz, CH<sub>3</sub>), 0.947 (t, 6H,  $J = 7.34$  Hz, CH<sub>3</sub>), 0.887 ppm (t, 12H,  $J = 7.35$  Hz, CH<sub>3</sub>). <sup>13</sup>C NMR (<sup>1</sup>H-decoupled, CDCl<sub>3</sub>):  $\delta$  49.81, 49.69, 49.33, 49.14 (NCH<sub>2</sub>), 29.46, 29.24 (NCH<sub>2</sub>CH<sub>2</sub>), 20.15, 20.06 (NCH<sub>2</sub>-CH<sub>2</sub>CH<sub>2</sub>), 13.83 ppm (CH<sub>3</sub>). <sup>51</sup>V NMR (CDCl<sub>3</sub>):  $\delta$  +101 ppm.

**Other Measurements.** Infrared spectra (Nujol mull) were recorded on a Nicolet 510P FTIR or on a Perkin-Elmer 283. Cyclic voltammetry experiments were performed in a nitrogen-filled glovebox using a BAS CV-27 potentiostat and an IBM Model 7424M **X-Y** recorder. The reference electrodes used consisted of a cracked glass bead outer shell (Fisher) containing a Ag wire in contact with a saturated solution (MeCN or THF) of AgCl/MedNCl. A **Pt** working electrode and a **Pt** mesh auxiliary electrode were also used. The concentration of electroactive species was ca. 2 mM with a supporting electrolyte  $(\text{``Bu}_4 \text{NPF}_6)$ concentration of 100 mM. Potentials are referenced vs  $FeCp_2/FeCp_2^+$ . No IR compensation was employed. The scan rate was 200 mV/s. 'H and I3C NMR spectra were recorded on a Varian XL-300 (299.94 MHz,  $H$ ; 75.47 MHz, <sup>13</sup>C) or a Bruker AM-500 (500.137 MHz, <sup>1</sup>H; 125.73 MHz, 13C) at Indiana University. Chemical shifts were referenced to the carbons or residual protons of the solvent. 51V NMR spectra (94.73 MHz) were obtained on a Bruker AM-360 spectrometer using 10 mm diameter sample tubes at the University of Hamburg. Chemical shifts were referenced to an external standard of VOCl<sub>3</sub> (0 ppm). UV-visible spectra were recorded on a Hewlett-Packard 8452A diode array spectrophotometer. Elemental analyses were performed by Galbraith Laboratories, Knoxville, TN.

X-my **CrystaUograpby and** Structure **Solution.** Data were collected on a Picker four-circle diffractometer by using standard low-temperature procedures; details of the diffractometry, low-temperature facilities, and computational procedures employed by the Molecular Structure Center are available elsewhere.<sup>5a</sup> Data collection parameters are summarized in Table I. The structures were solved by a combination of direct methods (MULTAN78) and Fourier techniques and refined by full-matrix leastsquares methods. No absorption correction was performed.

 $(Me<sub>3</sub>NBz)<sub>2</sub>V(S<sub>2</sub>)S<sub>2</sub>SPh$ ] (1). A systematic search of a limited hemisphere of reciprocal space located a set of diffractional maxima with monoclinic symmetry and **systematicabsencescorresponding** to the unique space group  $P2_1/n$ . The positions of all hydrogen atoms were clearly visible in a difference Fourier map phased on the non-hydrogen atoms, and the coordinates and isotropic thermal parameters for hydrogens were varied in the final cycles of refinement. A final difference Fourier was essentially featureless, with the largest peak being  $0.5 \frac{e}{A^3}$ .

(Me<sub>3</sub>NBz)<sub>4</sub>V(S<sub>2</sub>)<sub>2</sub>(CS<sub>3</sub>)<sub>4</sub>] (2). A suitable crystal was kept in contact with the mother liquor until mounted toavoid solvent **loss** problems noticed with earlier crystals, which were found to diffract weakly or not at all. A systematic search of a limited hemisphere of reciprocal space revealed symmetry and systematic absences corresponding to the unique monoclinic space group  $P2_1/c$ . The V and S atom positions were obtained from an initial E-map. The remainder of the non-hydrogen atoms were found in subsequent iterations of least-squares refinement and difference Fourier

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Table I. Crystallographic Data for Complexes 1 and 2

parameters		2
formula	$C_{26}H_{37}N_{2}S_{5}V$	$C_{46}H_{67}N_5S_{16}V_2$
molar mass, g mol <sup>-1</sup>	588.85	1304.91
cryst syst	monoclinic	monoclinic
space group	$P2/$ n	P2 <sub>1</sub> /c
temp, °C	$-158$	$-114$
a, A	$18.178(8)^a$	20.978 (4) <sup>b</sup>
b, A	8.843(2)	13.384(2)
$c, \Lambda$	18.852(7)	21.613(4)
$\beta$ , deg	107.06 (2)	93.14(1)
z	4	4
$V, \mathbf{A}^3$	2897.16	6059.27
radiation. A	0.71069	0.710 69
abs coeff, cm <sup>-1</sup>	6.951	8.643
$R, \%$	5.65	6.32
$R_w$ , % <sup>d</sup>	5.63	6.75

<sup>*a*</sup> 34 reflections. <sup>*b*</sup> 50 reflections.  $^c R = \sum ||F_0| - |F_c||/\sum |F_0|$ . *d*  $R_w$  =  $[\sum w(|F_o| - [F_c])^2 / \sum w|F_o|^2]^{1/2}$ , where  $w = 1/\sigma^2(|F_o|)$ .



**Figure 1.** ORTEP representation of the anion of complex **1** at the **50%**  probability level.

calculations. After partial refinement of the non-hydrogen atoms, **a**  difference Fourier map revealed some, but not all, hydrogen atoms. All hydrogen atoms except those of the MeCN solvate molecule were introduced in fixed calculated positions. In the final cycles of Ieastsquares refinement, all non-hydrogen atoms were varied with anisotropic thermal parameters. A final difference Fourier map was featureless, with the largest peak being 0.99  $e/\text{\AA}^3$ .

**Molecular Orbital Calculations.** Calculations were performed using the EHMO method with weighted  $H_{ij}$ 's.<sup>5b</sup> The parameters for V,<sup>5c</sup> S,<sup>5d</sup> and C<sup>5e</sup> were taken from the literature. Atomic coordinates were taken from the crystallographic data.

## **Results and Discussion**

**Syntheses and Structures.** The original synthesis of  $[V(S_2)S_2-$ (SPh)]2- **(1)** was plagued with concomitant formation of a purple oligomeric material of as-yet undetermined formulation.<sup>4c</sup> We have since discovered that replacement of NaSPh with LiSPh in a reaction mixture comprising VC13, LiSPh, **S,** and NEt4CI in a 1:7:3:2 molar ratio in MeCN avoids the formation of the purple material and increases the yield of **1** from ca. 35% to ca. *65%.*  The material crystallizes directly from the reaction solution in a form suitable for crystallography. Table **I1** lists atomic coordinates, and Figure 1 shows an ORTEP representation of the anionic portion of compound **1.** Selected bond distances and angles are listed in Table 111. The vanadium(V) center is ligated by three types of sulfur atoms, viz. one PhS<sup>-</sup>, one persulfido  $(S_2^2)$ , and two sulfido  $(S^2)$  groups, yielding a  $VS_5$  core of idealized  $C_5$ symmetry. The coordination geometry is best described as tetrahedral with the  $S_2^2$ - considered as occupying one site. Although many vanadium(II1 or IV)/thiolate linkages are known,<sup>4a,6,7</sup> compound 1 is the only structurally-characterized

Table II. Selected Fractional Coordinates (X10<sup>4</sup>) and Isotropic Thermal Parameters (A2 **X** 10) for Complexes **1** and **2** 

atom	x	y	z	$B_{\rm iso}$
Complex 1				
V1	9226 (1)	1923 (1)	1687 (1)	20
S <sub>2</sub>	9948 (1)	2043 (2)	1005 (1)	24
S3	9958 (1)	3049 (2)	2802 (1)	38
S4	9119(1)	$-305(2)$	2027 (1)	23
S5	8995(1)	4151 (2)	2239 (1)	35
S6	7965 (1)	2384 (2)	907(6)	28
		Complex 2		
V <sub>1</sub>	272(1)	979 (1)	4953 (1)	15
V <sub>2</sub>	5452(1)	677(1)	9771 (1)	16
S3	406 (1)	$-141(2)$	5813(1)	21
S4	$-477(1)$	445 (2)	5707 (1)	18
S5	713(1)	2237 (2)	5711 (1)	23
S6	1459(1)	931 (2)	5034 (1)	22
S7	505 (1)	2156 (2)	4097 (1)	24
S8	$-575(1)$	2291 (2)	4804 (1)	23
S9	$-560(1)$	3556 (2)	3662(1)	37
S10	2126 (1)	2011 (2)	6090 (1)	32
S11	5633(1)	$-445(2)$	10644 (1)	20
S12	5039(1)	704 (2)	10787(1)	21
S13	5773 (1)	1251 (2)	8744 (1)	23
S14	6448 (1)	$-188(2)$	9463 (1)	24
S15	5044 (1)	2428 (2)	9756 (1)	21
S16	6278 (1)	1869 (2)	10201 (1)	20
S17	5934 (1)	4070 (2)	10179(1)	25
S <sub>18</sub>	6872(1)	236(2)	8173(1)	37
C <sub>23</sub>	1461 (4)	1757 (7)	5634 (4)	22
C <sub>24</sub>	$-228(4)$	2729 (7)	4152 (4)	21
C <sub>25</sub>	6392 (5)	415(7)	8750 (5)	27
C <sub>26</sub>	5764 (5)	2857 (7)	10062 (5)	25

complex with this linkage at the *+S* metal oxidation state. The V-S6-C7 angle of 111.67 (24)<sup>o</sup> is very close to that found in **vanadium(III)/benzenethiolate** compounds.' Unlike the case of alkoxides, this angle is not indicative of the amount of metalsulfur  $d\pi$ -p $\pi$  bonding, since the sulfur 3p orbital is oriented perpendicular to the M-S-C plane.<sup>8</sup> The S-S bond length in the  $S_2^2$ - ligand of 2.013 (3) Å is typical<sup>9</sup> for bound persulfide and is consistent (vide infra) with ligand *r\** to empty metal d-orbital electron donation (free  $S_2^2$ ;  $\bar{S}-S = 2.13 \text{ Å}$ ).<sup>10</sup> The V-S2 and V-S4 distances of 2.092 (2) and 2.099 (2) Å are somewhat shorter than those in  $VS_4^{3-}$  (2.135–2.175 (1) Å),<sup>11</sup> the only other known vanadium(V) monomer with terminal sulfide ligation.

The unusual nature of **1,** particularly the VV/PhS- linkage, suggested that it might be highly reactive, and we have therefore investigated its reactions with, among other things, carbon disulfide. Addition of excess  $CS<sub>2</sub>$  to an MeCN solution of compound **1** rapidly formed a deep red color. Following filtration, well-formed crystals of 2 grew slowly from this solution at ambient temperature in good yield. Elemental analysis indicated two cations and eight sulfur atoms per vanadium. The formula and nuclearity of this product were determined by X-ray crystallography to be  $(Me_3NBz)_4[V_2(S_2)_2(CS_3)_4]$  (2). The formation of **2** can be summarized as shown in *eq* 1. Oxidation of the thiolate

$$
2(Me_3NBz)_2[V(S_2)S_2(SPh)] + 4CS_2 \xrightarrow{MeCN} 1
$$
  
\n
$$
(Me_3NBz)_4[V_2(S_2)_2(CS_3)_4] + PhSSPh (1)
$$

in compound **1** to thedisulfide provides for concomitant reduction of vanadium(V) to vanadium(1V).

The crystallographic study of **2** revealed two independent molecules in the unit cell with each dimeric molecule residing on

~~ ~~~ ~

- 
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Table III. Selected Distances (Å) and Angles (deg) for Complexes **1** and **2** 

Complex 1					
$V1 - S2$	2.0919 (20)	V1–S6	2.3674(23)		
V1–S3	2.3549 (23)	S3–S5	2.013(3)		
$V1 - S4$	2.0988 (20)	S6-C7	1.765(7)		
$V1 - S5$	2.3230 (23)				
$S2-V1-S3$	103.57 (9)	S4-V1-S5	127.88 (8)		
$S2-V1-S4$	111.52(8)	S4-V1-S6	101.08 (8)		
$S2-V1-S5$	117.40 (8)	S5-V1-S6	82.11(8)		
$S2-V1-S6$	106.14(8)	$V1 - S3 - S5$	63.70(8)		
$S3-V1-S4$	101.69 (8)	$V1 - S5 - S3$	65.33 (9)		
$S3-V1-S5$	50.97 (8)	V1-S6-C7	111.67 (24)		
$S3-V1-S6$	132.24 (8)				
		Complex 2 <sup>a</sup>			
$V1-V1'$	2.8720 (59)	V2-V2′	2.8411 (57)		
$V1-S3$	2.3930 (28)	V2–S11	2.4205 (27)		
$V1-S3'$	2.4009 (28)	$V2 - S11'$	2.4258 (27)		
$V1-S4$		$V2-S12$			
	2.4324 (27)		2.406(3)		
$V1-S4'$ $V1 - S5$	2.4325 (27)	$V2-S12'$ $V2-S13$	2.4070 (28)		
	2.492(3)		2.476(3)		
$V1 - S6$	2.4859 (27)	$V2-S14$	2.510(3)		
$V1 - S7$	2.4973 (28)	$V2 - S15$	2.4949 (28)		
V1–S8	2.5069 (28)	$V2 - S16$	2.4968 (27)		
$S3 - S4$	2.012(3)	S11-S12	2.014(3)		
$S3-V1-S3'$	106.39 (9)	$S4-V1-S7$	147.26 (10)		
$S3' - V1 - S4$	86.79 (9)	$S4-V1-S8$	79.15 (9)		
$S3-V1-S4$	49.29 (8)	$S4' - V1 - S8$	128.35 (10)		
$S3-V1-S4'$	86.97 (9)	$S5-V1-S6$	68.72 (9)		
S3'-V1-S4'	49.20 (8)	$S5-V1-S7$	88.92 (10)		
S3-V1-S5	83.59 (9)	S5-V1-S8	81.16 (9)		
$S3' - V1 - S5$	162.51 (10)	$S6-V1-S7$	80.44(9)		
$S3-V1-S6$	81.60 (9)	$S6-V1-S8$	136.72 (10)		
$S3' - V1 - S6$	126.15 (10)	S7-V1-S8	68.27 (9)		
$S3-V1-S7$	162.01 (10)	$V1 - S3 - V1'$	73.61 (9)		
$S3' - V1 - S7$	85.24 (9)	$V1-S3-S4$	66.38 (10)		
$S3' - V1 - S8$	81.36 (9)	$V1 - S3' - S4$	66.22 (10)		
$S3-V1-S8$	126.17 (10)	$V1 - S4 - V1'$	72.36 (8)		
S4-V1-S4'	107.64 (8)	$V1 - S4' - S3$	64.58 (10)		
S4-V1-S5	89.28 (9)	$V1 - S4 - S3$	64.34 (10)		
$S4' - V1 - S5$	147.56 (10)	$V1 - S5 - C23$	89.5 (3)		
$S4-V1-S6$	128.71 (10)	$V1 - S6 - C23$	89.9 (3)		
S4'-V1-S6	79.24 (9)	V1-S7-C24	91.0(3)		
$S4' - V1 - S7$	90.72 (9)	$V1 - S8 - C24$	90.8(3)		

<sup>a</sup> Only limited data are listed for the second independent anion (V2, V2'); full listings are available in the supplementary material.



**Figure 2.** ORTEP representation of the anion of complex 2 at the 50% probability level.

an inversion center. An ORTEP projection of one of the independent anions is shown in Figure 2; atomic coordinates and structural parameters are listed in Tables I1 and 111, respectively. The metal centers are bridged by two symmetry-related  $\eta^2$ : $\eta^2$ : $\mu_2$ -S<sup>2-</sup> ligands, and two bidentate chelating trithiocarbonate ligands



**Figure 3.** Portion **of** the infinite linear chain present in the mineral patronite.

**Table IV.** Comparison of Core Distances in  $V_2(S_2)_2Z_4$ 

	z				
	$(CS_3)^{2-}$	$(S_2CN^{i}Bu_2)^{-1}(S_2CNEt_2)^{-1}$		$(S_2CCH_3)$	patronite, $[V(S_2)_2]$
v-v	2.872	2.851	2.884	2.800	2.83
$V-S(av)$	2.415	2.405	2.409	2.392	2.41
s s	2.012	1.998	1.984	1.997	2.03
ref	α	12	14	13	15

This **work.** 

complete the coordination sphere about each vanadium. The metal centers are eight-coordinate, and the coordination geometry is best described as distorted dodecahedral. The V-V distances in the two independent molecules are 2.872 *(6)* and 2.841 *(6)* A, respectively, which is within bonding range (vide infra). Similar  $[V_2(S_2)_2]^{4+}$  cores have been observed in the molecular compounds (S2CNEt2)4.14 These complexes, including **2,** are all related to the polymeric, solid-state mineral known as patronite. The latter consists of linear chains of vanadium(1V) ions bridged by two **S22-** ligands, as illustrated in Figure **3."** The metal centers in patronite have pairwise bonding interactions (2.83 **A)** and longer, nonbonding distances  $(3.22 \text{ Å})$ . A comparison of these core distances is tabulated in Table IV.  $V_2(S_2)_2(S_2CN^iBu_2)_4$ ,<sup>12</sup>  $V_2(S_2)_2(S_2CMe)_4$ ,<sup>13</sup> and  $V_2(S_2)_2$ -

The rapidity of the reaction and lack of suitable monitoring techniques make it difficult to deduce the mechanism of the conversion of **1** to **2.** Of particular interest would **be** the initial site of attack of  $CS_2$  on the  $[V(S_2)S_2(SPh)]^{2-}$  anion. Given the retention of the  $V: S_2^2$  ratio (1:1) of 1 in the product 2 and the loss of terminal sulfides, it is tempting to postulate attack of  $CS<sub>2</sub>$ at theVS groups **todirectlygivechelatingCS32-groupsvia** [2+2] cycloaddition *(eq* 2). Although rare, such cycloaddition reactions

 $\begin{array}{c}\n\circ \\
\circ \\
\circ \\
\circ \\
\circ\n\end{array} \longrightarrow \begin{array}{c}\n\circ \\
\circ \\
\circ \\
\circ \\
\circ \\
\circ\n\end{array}$  $(2)$ 

involving  $M=X$   $(X = 0, S)$  are known. For example,  $(CF_3)_2$ C=O cycloadds<sup>16</sup> across the W=O group of Cp<sub>2</sub>W=O and, in a similar fashion, SO<sub>2</sub> cycloadds across a Re-O bond on each Re atom of  $Cp^*{}_2Re_2O_4$ .<sup>17</sup> Most recently, the complex  $(tmtaa)$ Ti=O has been shown to undergo many  $[2+2]$  cycloadditions with a number of reagents such as ketones,  $SO_2$ ,  $SO_3$ , and  $CS<sub>2</sub>$ . Similarly, the thio complex (tmtaa)Ti=S is reported to react with  $CS_2$  to give (tmtaa)Ti( $CS_3$ ).<sup>18</sup> Thus, the reaction of **1** with CS<sub>2</sub> might also be proceeding via a cycloaddition reaction with the terminal  $V=$ S groups; major loss of the  $\pi$ -donation to the metal from these multiply-bonded **S2-** groups would then trigger both reduction of the metal by the PhS- group and

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aggregation via conversion of terminal  $S_2^2$  groups into bridging modes  $(eq 3)$ . Alternatively, it cannot be ruled out that the  $CS<sub>2</sub>$ 



is attacking at other sites on the  $[V(S_2)S_2(SPh)]^{2-}$  ion, most obviously the  $S_2^2$ -group, and that internal electron redistribution then oxidizes the  $S<sup>2</sup>$  groups to  $S<sub>2</sub><sup>2</sup>$ , which adopts the bridging mode in **2.** However, this possibility should initially generate a  $CS<sub>4</sub><sup>2-</sup>$  unit that would then have to transfer a sulfur atom to either a second  $CS_2$  or a  $S^{2-}$ . At the very least, this mechanism would be more complex. Unfortunately, we do not have access to suitable derivatives of 1 (e.g., containing the  $[V(S_2)O_2(SPh)]^{2-}$ ion) to probe this matter further.

Extended Hikkel Molecular **Orbital** Calculations. Both  $V_2(S_2)(S_2CN^iBu_2)_4$  and  $V_2(S_2)_2(S_2CMe)_4$  were reported to be diamagnetic,<sup>12,13</sup> as is  $[V_2(S_2)_2(CS_3)_4]^{4-}$  on the basis of its <sup>1</sup>H, <sup>13</sup>C, and <sup>51</sup>V NMR spectra (vide infra). In an effort to address the extent of metal-metal bonding in **2** (as opposed to strong antiferromagnetic coupling via the bridging ligands), EHMO calculations have been performed on the  $[V_2(S_2)_2(CS_3)_4]^{\text{4-}}$  ion. (Calculations have previously been reported on 'CpMS4MCp" systems employing metal fragments of  $C_{3v}$  symmetry.<sup>19</sup>) Of the coordinates of two independent molecules present in the unit cell of **2,** the (nonidealized) coordinates of the one with the long vanadium-vanadium distance were used. The bridging persulfides were treated as one fragment and the two " $V(CS<sub>3</sub>)<sub>2</sub>$ " units were viewed as distorted ML<sub>4</sub> fragments with  $C_{2v}$  symmetry.<sup>20</sup> Symmetry labels were assigned according to the  $D_{2h}$  symmetry of the entire molecule. As shown in Figure **4,** there is a metalbased orbital match for each of the persulfide  $\pi$  and  $\pi^*$  orbitals. This match leads to agood interaction between the two fragments and a large HOMO/LUMO gap (ca. 1.2 eV); the HOMO possesses approximately 13% metal 3d character. Significant electron density is transferred from the high-lying persulfide  $\pi^*$ orbitals to empty metal-based orbitals, thus decreasing the S-S bond distance (2.012 (3)  $\hat{A}$  in 2 vs 2.13  $\hat{A}$  in free  $S_2^{2-}$ ).<sup>10</sup> In regard to a metal-metal  $\sigma$  bond, there is a greater amount of electron density transferred from the lower lying persulfide  $\pi$ orbitals into the metal-metal bonding,  $d_{z^2}$  combination  $(\sigma)$  than is transferred into the out-of-phase  $(\sigma^*)$   $d_{z^2}$  combination. These results lend support for the existence of a weak metal-metal bond, although it should be noted that there is extensive mixing of sulfur character into this molecular orbital. Indeed, many quadruply- and triply-sulfur-bridged early-transition-metal compounds possess a formal metal-metal bond.19

Comparisons with Mo/S Systems. It has always been of interest **tocompareresultsobtained** inour V/Sstudies with thoseavailable from better-understood Mo/S chemistry. Coucouvanis and *co*workers have shown that the Mo- $\eta^2$ -S<sub>4</sub> group present in both the molybdenum(IV) monomer  $[(S_4)_2M_0=S]^2$  and the molybdenum(V) dimer  $[Mo<sub>2</sub>S<sub>4</sub>(S<sub>4</sub>)<sub>2</sub>]<sup>2-</sup>$  is in equilibrium with the Mo- $\eta^2$ -S<sub>2</sub> (persulfide) group and elemental sulfur, as shown in eqs 4 and  $5.21$  The reaction chemistry of the Mo- $\eta^2$ -S<sub>4</sub> group is dominated by the presence of the  $Mo-\eta^2-S_2$  group. Since the  $[(S_4)Mo(S)(S_2)]^2$  and  $[(S_4)Mo_2S_4(S_2)]^2$  compounds contain



**Figure 4.** Interaction diagram for  $[V_2(S_2)_2(CS_3)_4]^{4-}$  (anion of 2). Labels are for  $D_{2k}$  symmetry.



structural elements<sup>22</sup> in common with the vanadium(V) monomer  $[V(S_2)S_2(SPh)]^2$  (terminal sulfido and persulfido), it is of interest to compare their reactivity characteristics. Both of the molybdenum species react with CS<sub>2</sub> to form "(CS<sub>4</sub>)<sup>2-"</sup> units as shown in eqs 4 and  $5^{10,23}$  It should be emphasized that the Mo- $\eta^2$ -S<sub>2</sub> unit is not 'pure" in the sense that it is in equilibrium with the Mo- $\eta^2$ -S<sub>4</sub> group and it is not known with which species the CS<sub>2</sub> reacts. Another possibility is that  $CS<sub>2</sub>$  undergoes cycloaddition with the Mo<del>S</del> unit followed by intramolecular S addition, forming the perthiocarbonate ligand.23b However, it is apparent in both cases that the terminal sulfido group *uppcars* not to react and is retained in the structure of the product. The different reactivity pattern observed with the vanadium specias **may** be due to greater nucleophilicity of **a** thiovanadyl unit: i.e., the **V--S**  group is more reactive toward electrophiles than the Mo-S group.

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**Spectroscopic Properties and Reactivity Characteristics of**   $[V_2(S_2)_2(CS_3)_4]^+$  (2). The <sup>1</sup>H NMR spectrum of complex 2 in  $CD<sub>3</sub>CN$  displays only sharp signals for the Me<sub>3</sub>NBz<sup>+</sup> cations at the expected chemical shifts, in accord with a diamagnetic nature for 2. In the <sup>13</sup>C<sup>[1</sup>H] NMR spectrum, there are the expected peaks for the cation and two peaks of comparable intensity at **253.0** and **252.8** ppm assigned to the carbon atoms of *two*  inequivalent  $CS_3^2$ - groups. This was an unexpected observation. The  $[V_2(S_2)_2(CS_3)_4]^+$  anion has virtual  $C_{2h}$  symmetry, and one would expect only a single  $CS<sub>3</sub><sup>2-13</sup>C NMR$  signal in solution. Twice as many signals as expected were also observed in the spectra of derivatives of **2** (vide infra) and it appeared that there was probably a common explanation. We believe this explanation to be that there are two isomers in solution and that their interconversion is slow on the NMR time scale, giving multiple signals. Thus, on dissolution of the *C2h* solid-state structure, an equilibrium is established in which the  $D_2$  isomer is also present. The two isomers differ only in the relative orientation of the  $CS<sub>3</sub><sup>2-</sup>$  chelates at the two ends of the molecule; the essence of this difference is indicated in simplified form in I. Each isomer of



**2** should give a single 13C resonance, resulting in the two observed signals of similar chemical shift. Theinterconversion could readily occur by, for example, partial dissociation of  $CS<sub>3</sub><sup>2-</sup>$  groups to monodentate modes followed by reattachment. Given the expected lability of an eight-coordinate VIv complex, it is reasonable that two isomers should exist in solution. Indeed, we note that  $V_2(S_2)_2(S_2CN^iBu_2)_4$  crystallizes in the  $D_2$  isomeric form, with the molecule lying on a crystallographic 2-fold rotation axis.<sup>12</sup> Thus, *both* isomers have been demonstrated to be possible. It is therefore conceivable that all members of this family of  $[V_2(S_2)_2]^{4+}$ -containing species might exist in isomeric forms in solution with the solid-state form dictated by lattice energies, kinetics of crystallization, and the space group of the crystal phase. Complex **2,** remember, crystallizes in *Pi* with the anion on an inversion center, dictating the  $C_{2h}$  isomer. As previously observed in other **trithiocarbonate-containing 2**  exhibits strong absorptions at **990** and **885** cm-l assigned to the *v*(C=S) and *v*(C $\cdot$ S) stretching vibrations, respectively.

Previous work has demonstrated that the terminal sulfur of a metal-bound trithiocarbonate complex is nucleophilic and can react with alkyl halides.<sup>24c,d,25</sup> Addition of excess MeI to an MeCN solution of **2** readily yields the methylated product 3 (as summarized in **eq 6).** which can be obtained in excellent yield

$$
(Me3NBz)4[V2(S2)2(CS3)4] + 4MeI V2(S2)2(S2CSMe)4 + 4Me3NBzI (6)
$$

and analytical purity. The  $H$  NMR spectrum of 3 in CDCl<sub>3</sub> **shows** two sharp singlets of comparable intensity at **2.662** and 2.658 ppm, in contrast to the single resonance expected simply from methylation of the solid-state structure. This is again consistent with two isomers being present in solution. Compound 3 also exhibits IR absorptions similar to other bound alkyl trithiocarbonates,<sup>24c,d,f,25,26</sup> with the major band occurring at 965  $cm^{-1}$ .

Alkyl trithiocarbonate compounds are known to undergo nucleophilic substitution reactions with amines to form dithiocarbamate compounds with concomitant elimination of the We have found complex 3 to undergo similar chemistry. Thus, 3 reacts with secondary amines in THF to give the corresponding dialkyldithiocarbamate **(eq 7).** The structures of these compounds have been discussed above. Alternatively, compounds **4a** and **4b** can be prepared directly from compound **2** via sequential additions of Me1 and the amine, as shown in **eq**  8. The coplanarity of the six atoms  $(S_2C=NC_2)$  of a dialkyl-

$$
V_{2}(S_{2})_{2}(S_{2}CSMe)_{4} + 4HNR_{2} \xrightarrow{\text{THE}} V_{2}(S_{2})_{2}(S_{2}CNR_{2})_{4} + 4MeSH (7) \xrightarrow{\text{4a (R = E)}} \text{4b (R = E)} \text{4b (R = "Bu)}
$$
\n
$$
[V_{2}(S_{2})_{2}(CS_{3})_{4}]^{4} \xrightarrow{\text{(2) HNR}_{2}} \text{4a or 4b} \tag{8}
$$

dithiocarbamate group and absence of rotation about the  $C=N$ double bond cause the methylene protons of the ethyl derivative **4a** to be diastereotopic. Thus, the 'H NMR spectrum of **4a** in CDCl3 exhibits a complex multiplet at ca. **3.7** ppm. When the methyl protons are decoupled, there are still complications from second-order effects, and **so** we resorted to the 13C spectrum. *As*  expected by now, four (instead of two) different 13C chemical shifts are observed from the methylene carbons in the  $^{13}C(^{1}H)$ NMR spectrum of **4a,** consistent with the presence of two isomers. The methyl region shows only two resonances, suggesting accidental overlap. High-temperature <sup>1</sup>H NMR spectra of compound **4a** in CDCl<sub>3</sub> remained unchanged up to 70 °C, the point at which decomposition began to occur. This observation suggests interconversion between the  $C_{2h}$  and  $D_2$  isomers is still slow on the NMR time scale even at 70 °C. Infrared spectra of the dithiocarbamate dimers **4.** and **4b** display strong absorptions at ca. **1490** cm-l, in accord with other metal dialkyldithiocarbamate complexes.<sup>24e,d,27</sup>

Electrochemistry. The electrochemical behavior of compounds **1-4** was investigated using cyclic voltammetry. None of the complexes exhibited any reversible features. **For** compound **1,**  cathodic scans in MeCN revealed an irreversible process at **-1.7**  V. Under similar conditions, compound **2** gave an irreversible wave at **-2.2** V. Cathodic scans of compound 3 in THF exhibited peaks at **-0.86, -1.18,** and **-1.88 V;** no corresponding oxidation peaks were observed on the return scan. An anodic scan of compound **4a** in THF revealed peaks at **-0.42** and **0.36** V with additional peaks at **0.33** and **-0.32** V observed upon the return scan. Similarly, compound **4b** exhibited a peak at **0.082** V upon anodicscanning, with a peak at **0.22** V observed during thereturn scan.

**Solution <sup>51</sup>V NMR Results.** <sup>51</sup>V chemical shifts of complexes **1–4 are listed in Table V, together with data from the literature** for comparison. Figure **5** is a stack plot of selected spectra.

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**Table V.** <sup>51</sup>V NMR Shifts (Relative to VOCl<sub>3</sub>) for 1–4 and for **Selected Complexes from the Literature** 

		δ( <sup>51</sup> V),	
complex	solvent	ppm	ref
$[V_2S_6-\mu-S]^+$	$H_2O$	+1457	28
$[VS3SH]2-$	H,O	+1392	28
$[VS(SSiPh_3)_3]$	$C_7H_8/Me_2CO$	$+1410$	29
$[Cp^*2V_2(\mu-S)_3]$	THF	$+1630$	30
$[Cp^*2V_2(\mu-S)(\mu-S_2)_2]$	THF	+596	30
$[VO(OH)(S_2CNEt_2)_2]$	CDCl <sub>3</sub>	–468	31
$[VS2(S2)SPh]2$ (anion of 1)	<b>DMSO/</b>	+970*	ь
	CD <sub>3</sub> CN (1:1)		
$[V_2(S_2CS)_4(\mu-S_2)_2]$ <sup>+</sup> (anion of 2)	<b>DMSO/</b>	$+125$	b
	CD <sub>3</sub> CN (1:1)		
	<b>DMSO/</b>	+135	b
	CD <sub>3</sub> CN (7:1)		
$[V_2(S_2CZ)_{4}(\mu-S_2)_2]$			
$Z = SMe(3)$	DMSO/	$+173c$	b
	CD <sub>3</sub> CN (7:1)		
$Z = \text{NEt} (4a)$	CDCI <sub>3</sub>	$+103$	b
$Z = NnBu$ (4b)	CDCI.	$+101$	Ь
$[VO3(OH)]2-$	H <sub>2</sub> O	$-537$	32
$[VO2(O2)OH]2-$	H <sub>2</sub> O	$-623$	32
$[VO2(dipic)]^-$	$H_2O$	-529	33
[VO(O <sub>2</sub> )dipic]	H <sub>2</sub> O	-595	33
[VO(ONH2)dipic]	H <sub>2</sub> O	$-672$	33

**Two additional signals are observed at -385 and 414 ppm (see**  Figure 5), possibly corresponding to vanadium complexes where most of **the thio ligands arc replaced by DMSO and/or acetonitrile. From the**  relation between  $\delta({}^{51}V)$ , the ligand electronegativity, and the coordination number,<sup>31,34</sup> possible compositions are  $V(O/N)_xS$ , where  $x = 5, 4,$  or 3. *b* This work. <sup>*c*</sup> Prepared from 2 in DMSO/CD<sub>3</sub>CN by addition of a 10**fold molar excess of McI.** 

Line widths of 51V NMR signals of diamagnetic vanadium complexes in solution aredominated by thequadrupole relaxation mechanism ( $51V$ : 99.78% natural abundance,  $I = \frac{7}{2}$ ,  $Q = -0.04 \times 10^{-28}$  m<sup>2</sup>, relative receptivity ( ${}^{1}H = 1$ ) = 0.38) and hence depend on, inter alia, the local symmetry, the size and bulkiness of the complex, the electronic nature of the ligand, and the viscosity of the solution. In compounds  $1-4$ ,  $W_{1/2}$  values are typically around *0.7* **kHz,** which corresponds to relaxation times in the millisecond range. The fact that the  $V(IV)$  complexes 2-4 show relatively sharp signals indicates that thereis strong coupling between the two metal centers (negligible population of a triplet spin state), thus supporting the EHMO calculation results discussed previously. In contrast to the  ${}^{1}H$  and  ${}^{13}C$  spectra, no resolution of more than one peak is observed.

In compounds 1-4, the <sup>51</sup>V nucleus is deshielded with respect to VOCl<sub>3</sub>, the standard adopted in  $51V$  NMR spectroscopy, and also with **respect** to all common V(V) complexescontaining oxygen and/or nitrogen ligands in the coordination sphere. As a d<sup>o</sup> system, the  $V(V)$  complex 1 exhibits the so-called "inverse polarizability" (electronegativity) dependence of metal shielding"; $^{31,34,35}$  i.e., shielding decreases as the polarizability of the ligand function increases (or its electronegativity decreases). Hence, generally speaking, sulfur ligands give rise to low shielding, as documented by the shift values of our complexes and the other thiovanadium complexes listed **in** Table V. But there are also distinct differences within the group of vanadium complexes with coordination spheres exclusively built up of sulfur functionalities.

Thus, in compound **1,** a complex of tetrahedral geometry (if the persulfide is considered to occupy a single coordination site), the vanadium nucleus is less effectively deshielded than, e.g., in the tetrahedral thiovanadates  $[VS_3SH]^2$  and  $[V_2S_6(\mu-S)]^+$  (cf. Table V). Apparently this is a consequence of the presence of persulfide; it has been previously demonstrated that replacement of an oxo by a peroxo group increases metal shielding.32 The

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400 1000 800 600 200 ò -200 -400 PPM **Figure 5.** Stack plot of selected <sup>51</sup>V NMR spectra: (a)  $[V(S_2)S_2(SPh)]^2$  $({\rm anion\ of\ 1});$  **(b)**  $[V_2(S_2)_2(CS_3)_4]^+$  (anion of 2); **(c)**  $V_2(S_2)_2(S_2CN^*Bu_2)_4$  $(4b)$ ; (d)  $V_2(S_2)_2(S_2CSMe)_4$  (3).

same is true for hydroxylamine  $(H_2NO^-)$  coordinating in the  $\eta^2$ mode (cf. Table V). This effect has been rationalized in terms of the strain (nonoptimal overlap) present in a three-membered ring<sup>31,34</sup> but may also, in the case of  $S_2^2$  vs  $S_2^2$  complexes, and in the light of the EHMO studies (vide supra), be traced back to a relatively large HOMO/LUMO gap for the  $V_2(S_2)_2$  moiety.

Compounds 2-4 all have  $\delta$ (<sup>51</sup>V) values between +103 and +173 ppm. With respect to 1, the <sup>51</sup>V nucleus is more effectively shielded by ca. 800 ppm. Again, this reflects the strain present in a chelate structure arising from replacement of the sulfide by the bidentate trithiocarbonato ligand, which gives **rise** to a rather **small S-VS**  angle of ca. *69O. An* even higher shielding **has been obeerved** for  $[VO(OH)(S_2CNEt_2)_2]$ ,<sup>31</sup> where, in addition to two four-membered rings, the **02-** and OH- ligands induce an extra upfield shift. Within the group of thiocarbonato  $(ZCS_2^{2-})$  complexes 2-4, the variation of  $\delta$ <sup>(51</sup>V) is relatively small, indicating that, in solution, the thiocarbonate appears to coordinate in a bidentate fashion in all cases, and the electronic second-sphere influences from the group **Z** are small.

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**Supplementary Material Available: Textural and tabular summaries** of the structure determinations, tables of atomic coordinates, thermal **parameters, and bond diatancos and angles, and additional structural diagrams for complexes 1 and 2 (31 pages). Ordering information is given on any current masthead page.** 

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