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Deoxygenation of Vanadyl Schiff's Base Complexes: Conversion of Oxocation Complexes into Halogeno-derivatives

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Summary Deoxygenation of vanadyl Schiff's base complexes by SOCl₂ gave the corresponding dichloro-derivatives of vanadium(IV).

The chemistry of vanadium(IV) is dominated by the stable VO²⁺ ion,¹ which forms a wide variety of complexes, with the VO²⁺ entity maintaining a discrete existence in each complex.² Six-co-ordinate complexes of the simple V⁴⁺ ion are rare, especially with chelating ligands.³

We report a route for converting the VO²⁺ unit into the more reactive VCl₂²⁺ unit, which can be used to transform many of the existent vanadyl complexes into the corresponding dichlorovanadium(iv) six-co-ordinate complexes, and may be useful for oxygen substitution in other oxocationic complexes.

There is a large amount of data on vanadyl Schiff's base complexes containing tetradentate⁴ or bidentate Schiff's base ligand.⁵ Most of the oxovanadium(IV) Schiff base complexes are monomeric, $v_{V=0}$ ca. $960 \, \mathrm{cm}^{-1}$, 2 except for NN'-propylenebis(salicylideneiminato)oxovanadium(IV)⁶ and its derivatives,⁴ which can be envisaged as having polymerised through the V=O unit. We have studied the electronic and/or steric conditions under which polymerisation of vanadyl complexes can occur. The only two examples of polymeric complexes seem to be those containing N-phenyl-p-chloro- and N-phenyl-p-nitro-(salicylaldimine) as ligands, which show an anomalously low V=O stretching frequency at 885 (p-Cl) and 875 cm⁻¹ (p-NO₂), and this seems to be diagnostic of a polymeric form via V-O-V interaction.⁴

Treating a benzene solution of bis-N-n-butyl(salicylideneiminato)oxovanadium(IV), (I) [VO(salNR)2; R = Bun] with a stoicheiometric amount of SOCl₂ at room temperature gave green-blue crystals of bis-N-n-butyl(salicylideneiminato)dichlorovanadium(IV), (II), quantitatively (reaction 1). In reactions under similar conditions with other vanadyl

$$[VO(salNBu^n)_2] + SOCl_2 \rightarrow [VCl_2(salNBu^n)_2] + SO_2$$
 (1)

complexes with bidentate ligands, R = PhCH₂, Ph, p-ClC₆H₄, the dichloro-derivatives may sometimes include benzene which can be lost by heating or recrystallisation from acetone. The resulting dichloro-complexes were identified via analysis, magnetic moment (μ_{eff} 1.7—1.8 B.M. at 292 K for all the complexes), and i.r. spectra.

The bis(acetylacetonato)oxovanadium(IV) complex [VO-(acac)2], treated in similar way with SOCl2, gave [VCl2-(acac)2], prepared previously by reaction of the diketone with [VCl₄] in anhydrous benzene.3 In this way the relatively unreactive VO2+ unit can be converted into the very reactive VCl_2^{2+} . The $[VCl_2(salNR)_2]$ complexes can be used as precursors to other substituted complexes and organovanadium(IV) compounds in which the V-C bond can be stabilized by the Schiff's base ligands.7 The Cl ligands exchange rapidly with other anionic ligands such as SCN-.8

Treating a benzene suspension of NN'-ethylenebis(salicylideneiminato)oxovanadium(IV), (III), with SOCl, at 40 °C for I h gave deep blue crystals of the dichloro-derivative, [V(salen)Cl₂], (IV), which was identified via analysis magnetic (1.72 BM at 292 K), and i.r. data, and shown to be identical to a sample of [V(salen)Cl2] obtained by reaction (2). Similar deoxygenation of NN'-o-phenylenebis(salicylideneiminato)oxovanadium(IV) gave the corresponding

$$[VCl_4] + salenH_2 \longrightarrow [VCl_2(salen)] + 2HCl$$
 (2)

dichloro-derivative. When the ligand around VO2+ is a relatively rigid tetradentate one the reaction with SOCI, seems to be more difficult. This was confirmed by the absence of reaction between vanadylphthalocyanine and SOCl₂, even under drastic conditions (excess of SOCl₂ in boiling benzene).

These results may be of interest with regard to the rôle of SOCl₂ in the deoxygenation.⁹ The presence of flexible ligands around the metal allows a multicentre, concerted attack by SOCl₂ and decomposition of the intermediates formed. A completely closed-ring equatorial ligand, such as the phthalocyanine, may hinder SOCl₂ attack and the opening of the V=O double bond (reaction 3).

Transfer of oxygen from an oxotransition metal compound or, vice-versa, deoxygenation of an organic substrate by transition metal complexes may be useful synthetically. Satisfactory elemental analytical data were obtained for all the compounds reported.

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