Dalton Transactions



PERSPECTIVE

View Article Online
View Journal | View Issue



Cite this: *Dalton Trans.*, 2025, **54**, 11464

Received 8th May 2025, Accepted 19th June 2025 DOI: 10.1039/d5dt01085c

Addressing misconceptions in dithiocarbamate chemistry

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Dithiocarbamates are monoanionic chelating ligands, easily prepared from CS_2 and secondary or primary amines, that find widespread use in agriculture, medicine, materials science and coordination chemistry. This is (in part) due to their ability to stabilise metals in a wide range of oxidation states, a result of *soft* (dithiocarbamate) and *hard* (thioureide) resonance forms. However, in the many thousands of publications on dithiocarbamate chemistry, some common misconceptions have arisen, often being accepted as truth. In this perspective we address some of these in the hope that, moving forward, the wider community will better grasp the nuances of the chemistry of this important ligand type.

1. Introduction

Dithiocarbamates, their complexes, and oxidised counterparts the thiuram disulfides (Fig. 1) find widespread uses in a diverse range of areas including: agriculture, 1,2 analytical chemistry,^{3–7} coordination chemistry,^{8–18} environmental remediation, ^{19–23} medicine, ^{24–29} enzyme inhibition, ^{30–36} medical imaging,^{37–39} living polymerisation,^{40–43} materials metal-sulfide and as precursors to nanomaterials. 49-52 They are a subset of the widely studied 1,1'dithiolate ligands and close relatives of thiocarbamates and carbamates.^{8,9} Despite being known for at least 150 years they remain an area of significant research activity. For example, with the recent identification of a new cell death mechanism termed cuprotosis,⁵³ the anti-cancer activity of [Cu(S₂CNEt₂)₂], which has been known for over 40 years, 54-59 has once again come under the spotlight.60-64 Thus, tetraethyl thiuram disulfide (Et4TDS), better known as Disulfiram (Antabuse), is a drug that finds widespread use in the treatment of alcoholism.65 It is rapidly metabolised in the gut (it is normally given orally) or in blood⁶⁶ to give diethyldithiocarbamate, which in turn can bind to Cu(II) to afford [Cu(S₂CNEt₂)₂] in situ.

Dithiocarbamates are easily prepared, generally in high yields, upon reaction of CS₂ with secondary or primary amines generally in the presence of an added base. Water is often used as the solvent, although reactions also proceed in MeOH and some other organic solvents. Reactions can be carried out in air, making dithiocarbamates easily accessible in less sophisticated lab environments. Consequently, this has led to an extremely large volume of research output: a search for "dithio-

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carbamate" in SciFinder® giving almost 25 000 hits and for "dithiocarbamato" another *ca.* 1500. Many of these publications are of excellent quality, some being cited thousands of times. However others contain errors and misconceptions, some of which are repeated so frequently as to be erroneously accepted as truth.

As researchers who are active in the area^{67–72} and have written significant reviews, ^{8,24,49} we have read and closely scrutinised a considerable number of papers on dithiocarbamate chemistry. In doing so we have seen some errors-misconceptions being regularly repeated. In this contribution we address these, with the hope that they may appear less frequently in the future. In general, our intention is not to highlight individual contributions where errors have been made or perpetuated, however at times this is unavoidable. We accept that the majority are made in good faith and hope to avoid demonising individuals.

2. Synthesis of dithiocarbamates

(i) Amines are deprotonated by base, and it is the amide that undergoes nucleophilic attack at CS₂ to afford the dithiocarbamate

Primary and secondary amines are extremely basic, having pK_bs of ca. 4. Consequently, they are not acidic and have pK_as of ca. 40. Thus, upon dissolution in water (or an organic solvent), even following the addition of a strong base such as NaOH, they are not deprotonated. They are, however, nucleophilic and like many N-based nucleophiles can react directly with the electrophilic carbon in CS_2 . This generates a zwitterion which cannot be isolated or spectroscopically identified, presumably since the equilibrium lies to the left-hand side. It can, however, be deprotonated by the added base, which if it is

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Fig. 1 Dithiocarbamates, their complexes and thiuram disulfides.

Fig. 2 Reaction of secondary amine with CS₂ in the presence of MOH as a base.

a hydroxide then simply affords water and the desired dithiocarbamate (Fig. 2).

Unfortunately, there are many examples in the literature that show the secondary (or primary) amine being deprotonated by the base and then suggesting that it is the generated amide [NR₂] or [NHR] that reacts with CS₂. As stated above, most bases used in the formation of dithiocarbamates are not basic enough to deprotonate the amine, and even if they were, the generated amides are very susceptible to hydrolysis and would immediately regenerate the amine. We note that alkali metal amides such as LiNMe₂ are highly pyrophoric and should be handled in a rigorously anhydrous environment (also see section 2-iii).

(ii) In the absence of an added base, CS2 reacts with a primary or secondary amine to afford the dithiocarbamic acid

Many secondary and primary amines react directly with CS2 and do not require added base. Thus, it is tempting to suggest that the reaction furnishes the free dithiocarbamic acid. This is, however, erroneous and rather this reaction generates the corresponding ammonium salt of the dithiocarbamate (Fig. 3).

Thus, it can be viewed in the same way as that above (Fig. 2), with the zwitterion being deprotonated by the amine which acts as both nucleophile and base. Thus, the overall stoichiometry is not 1:1 but rather 2:1, amine to CS₂. ⁷⁴⁻⁷⁶ Indeed, three purported crystal structures of dithiocarbamic acids have been reported⁷⁷⁻⁷⁹ (one later revised).80 The crystal structure of an adduct of hexylamine (HexNH₂) and the dithiocarbamic acid, HexNHCS2H 78 is really the ammonium salt (Fig. 4a), but the proton has been incorrectly located on sulfur rather than hexylamine. Likewise, in the proposed amino acid

derivative (Fig. 4b) the proton is actually located on the basic nitrogen.79

Sometimes, even when a second base has been added, the ammonium dithiocarbamate is (in part) generated in the reaction especially when the amine is a strong base, as the two compete to deprotonate the zwitterion. This is the case in the reaction of ¹Bu₂NH and CS₂ in water with added NaOH, a reaction we have carried out numerous times in our lab. Thus, the off-white ammonium salt has poor water solubility and can precipitate, often being observed as a "scum" in the reaction vessel. Simply stirring in a warm (ca. 50-60 °C) water bath for 20-30 min leads to complete dissolution and formation of a clear pale-yellow solution of NaS₂CNBuⁱ₂.

Nice illustrations of an amine acting as both a base and a nucleophile are in the reaction of diamines with CS2. For example, reaction of piperazine with CS2 affords as the major product a zwitterionic product (Fig. 5) together with a small amount (ca. 5-10%) of the (so-called) double salt, thus providing an elegant method of desymmetrising a cheap diamine which can then be used to build up novel multimetallic arrays.81-86

(iii) Reactions of diarylamines with CS2 in the presence of MOH afford diaryldithiocarbamates

Anilines are nucleophilic enough to react directly with CS2 and thus dithiocarbamates, [ArNHCS₂]⁻ are readily available.⁸⁷ In contrast, diphenylamine and other diarylamines are not nucleophilic enough and will not react directly with CS2 in the presence of standard bases. Thus, to prepare [Ar₂NCS₂]⁻, the corresponding amides [NAr2] need to be accessed; these are nucleophilic enough to react with CS₂. Reagents that are basic

$$R_2NH + CS_2$$
 $R_2NH + CS_2$ $R_2NH + CS_2$ R_2NH_2

Fig. 3 Reaction of secondary amine with CS₂ in the absence of a second base.

$$(b) \qquad \overset{\text{H}_2\text{N}}{\underset{\text{K}^{\bigoplus} \odot}{\text{O}_2\text{C}}} \overset{\text{S}}{\underset{\text{N}^{\bigoplus} \odot}{\text{N}}} \qquad \overset{\text{H}_3\text{N}}{\underset{\text{K}^{\bigoplus} \odot}{\text{O}_2\text{C}}} \overset{\text{S}}{\underset{\text{S}}{\text{N}}} = 0$$

Fig. 4 Actual and proposed forms of crystallographically characterised "dithiocarbamic acids".

Fig. 5 Reaction of piperazine with CS₂.

Fig. 6 Generation of diaryldithiocarbamate salts upon addition of CS₂ to LiNAr₂.

enough to deprotonate Ar₂NH include ⁿBuLi, ^{88,89} NaBH₄, ⁹⁰ sodium amide ^{91,92} and KOBu^t. ⁶⁷ All require water-free conditions otherwise the neutral amines will simply be regenerated. Thus, we have recently developed a preparation ⁸⁸ that uses ⁿBuLi in thf to afford LiNAr₂ at low temperatures, which in turn react with CS₂ to form LiS₂CNAr₂ in (essentially) quantitative yields ⁶⁷ (Fig. 6). Once formed these dithiocarbamate salts are very stable and can be stored indefinitely as solids in air. We have also had success with using KOBu^t in thf but found this not to be reproducible in regular lab grade solvents. As stated above, other authors have used NaBH₄ but in our hands this was not successful.

(iv) Amides such as phthalimide and succinimide react with CS₂ to generate dithiocarbamates

As discussed above, amines need to be relatively nucleophilic if a dithiocarbamate is to be made directly, and amides are not nucleophilic enough to react directly with CS_2 . Despite this, several publications claim the synthesis of amide-derived dithiocarbamates and their complexes, $^{93-97}$ some of which we have failed to reproduce. 93,94 As highlighted for diarylamines, a second approach is to initially deprotonate to yield the more nucleophilic anion. In this way, we have tried reacting commercially available sodium phthalimide with CS_2 under a

variety of conditions but in all instances, we found no evidence for the formation of the corresponding dithiocarbamate (Fig. 7).

In contrast, 2-pyrrolidone does react with CS₂ in the presence of bases such as KOH, to afford a dithiocarbamate which can be quenched with electrophiles to give the corresponding ester. ^{98–100} Formamide also reacts with CS₂ in the presence of base as confirmed by the crystal structure of KS₂CNH(CHO). ¹⁰¹ There is a short paper on metal complexes of 2-pyrrolidone dithiocarbamate, ¹⁰² which suggests this ligand type could be further developed, along with that of iso-indolinone, for which the dithiocarbamate has not been reported but is likely accessible.

There are several authenticated examples of amide-functionalised dithiocarbamate complexes, being accessible *via*

$$N \ominus + CS_2 \longrightarrow N \ominus S$$

Fig. 7 Unsuccessful reaction of phthalimide with CS₂.

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Fig. 8 Synthesis and molecular structure (R = ⁱBu) of amide-functionalised nickel bis(dithiocarbamate) complex.

dithiocarbimate complexes. 103 For example, deprotonation of the corresponding primary amine-derived nickel complexes, $[Ni(S_2CNHR)_2]$, followed by quenching with electrophiles such as acetic anhydride and benzoyl chloride (Fig. 8). $^{104-107}$ These amide-dithiocarbamate complexes are very stable and show interesting physical properties, something we have recently been reinvestigating in our own research. 107

3. Stability of dithiocarbamates and thiuram disulfides

(i) Dithiocarbamic acids are stable and isolable entities

A major misconception in dithiocarbamate chemistry, being especially prevalent in the biological domain, is that dithiocarbamic acids are stable entities. Thus, as has been well studied, disulfiram is rapidly reduced in vivo to give two equivalents of diethyldithiocarbamate, which is often erroneously written as the free acid. 108 Chemists are also not immune to this mistake and often (as noted above) the direct reaction of an amine with CS₂ is purported to give the dithiocarbamic acid rather than the ammonium salt of the dithiocarbamate. 77-79,109-112 Dithiocarbamates are basic, especially those with two alkyl substituents, and at pH 7 or below (indeed even above this in some cases) they are protonated to give the dithiocarbamic acids. However, these are unstable and, in most cases, decompose rapidly to give CS2 and the corresponding ammonium salt (Fig. 9). Many publications have addressed the decomposition process, and mechanistic aspects have been elucidated. 113-124 Here is not the place to go into detail but the main decomposition route involves a hydrogen-bonded intermediate, rather than zwitterion formation resulting from proton transfer from sulfur to nitrogen. Decomposition is accelerated upon lowering the pH and this is why dithiocarbamates are generated under basic conditions. Recently, the generation and decomposition of dithiocarbamic acids has been repurposed as a route for the release of CS_2 . These studies show that their lifetimes are highly dependent upon the nature of the substituents, those with two aryl groups being stable for up to 24 h at pH 7.4. Dithiocarbamic acids of primary amines are especially unstable 119 but still appear in the literature. Dithiocarbamates of primary amines are widely used as precursors for the generation of organic isothiocyanates 127,128 and other sulfur-containing organics. 129,130

(ii) Dithiocarbamate halides are accessible

A relatively uncommon misconception, but one that is increasingly appearing in the literature, is the idea that halides of dithiocarbamates, especially the iodide, are accessible. ¹³¹ It is well known that iodine acts as an oxidising agent, converting dithiocarbamates into thiuram disulfides, rather than forming the corresponding iodide (Fig. 10). With primary amine dithiocarbamates this (likely) generates the unstable thiuram disulfide (see below), and not the iodide. ^{132,133}

It is worth adding that, while not a misconception, the redox chemistry of the dithiocarbamate ligand is often overlooked, which can lead to an over-simplification of discussion of redox events when coordinated to metal centres. Examples are reactions of $[M(S_2CNR_2)_2]$ (M=Zn,Cd) with I_2 . 134,135 Thus, reduction of iodine is rapid at the non-redox active Zn(II) centre as it is the dithiocarbamate that is oxidised, on metal, to form the corresponding thiuram disulfide complexes (Fig. 11).

(iii) Thiuram disulfides of primary amines are stable

Thiuram disulfides generated from secondary amines are generally very stable and can be easily isolated, purified and stored. Indeed, they are often easier to store than the corresponding dithiocarbamates and can then be reduced *in situ* to

$$R_2N \xrightarrow{\begin{array}{c} \\ \\ \\ \end{array}} \xrightarrow{\begin{array}{c} \\ \\ \end{array}} \xrightarrow{\begin{array}{c} \\ \\ \end{array}} \xrightarrow{\begin{array}{c} \\ \\ \end{array}} R_2N \xrightarrow{\begin{array}{c} \\ \\ \end{array}} \xrightarrow{\begin{array}{c} \\ \\ \end{array}} R_2NH + CS_2 \xrightarrow{\begin{array}{c} \\ \\ \end{array}} \xrightarrow{\begin{array}{c} \\ \\ \end{array}} R_2NH_2$$

Fig. 9 In situ generation and decomposition of dithiocarbamic acids.

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$$R_2N \xrightarrow{S} I_2 \qquad R_2N \xrightarrow{S} I_2 \qquad R_2N \xrightarrow{S} NR_2$$

Fig. 10 | 12 oxidation of dithiocarbamate to thiuram disulfide

$$R_2N$$
 NR_2
 R_2N
 R_2N
 R_2N
 R_2N
 R_2N

Fig. 11 I₂ addition to [Zn(S₂CNR₂)₂] and formation of a thiuram disulfide complex.

provide a source of dithiocarbamate. In contrast, oxidation of primary amine dithiocarbamates affords thiuram disulfides with poor stability, decomposing to afford isothiocyanates and/or thioureas, depending upon the reaction conditions used^{127,136-140} (Fig. 12).

Accessibility and air-stability of [M $(S_2CNR_2)_2$

One of the beauties of doing transition metal dithiocarbamate chemistry is their ease of synthesis, isolation, and purification, much of which can be carried out in air. Thus, complexes normally have moderate to good solubility in polar organic solvents, and many can be crystallised in air, either by slow evaporation or the (careful) addition of an anti-solvent such as hexane or petroleum ether. In this way, the crystal structures of thousands of dithiocarbamate complexes have found their way into the Cambridge Crystallographic Data Centre (CCDC). 141 However, not all dithiocarbamate complexes are air stable. For example, while bis(dithiocarbamate) complexes, [M (S₂CNR₂)₂] (M = Ni, Cu, Zn) are air and moisture stable, in contrast those of the other first row transition elements can only be prepared (if at all) under rigorously oxygen-free conditions. 142-144 Thus, generally, if they can be prepared, [M (S₂CNR₂)₂] complexes are readily oxidised in air, with M(III) complexes being favoured. Below we discuss each metal type individually as their chemistry differs. We also describe some related chemistry (where appropriate) to provide context.

(i) M = Ti, V, Cr

The chemistry of titanium is dominated by the +4 and +3 oxidation states, and dithiocarbamates are no exception. A Ti(II)

complex has briefly been mentioned in the literature, 145 reaction of [Ti(NEt₂)₃] and CS₂ being reported to give a mixture of [Ti(S₂CNEt₂)₂] and [Ti(S₂CNEt₂)₄]. However, no characterising data or reaction details were given. The existence and stability of red Ti(iv) complexes [Ti(S2CNR2)4] are without doubt, being prepared upon insertion of CS₂ into [Ti(NR₂)₄],¹⁴⁶ or upon addition of LiS₂CNR₂ to TiBr₄. Both [Ti(S₂CNEt₂)₄]¹⁴⁸ and [Ti(S₂CNPrⁱ₂)₄]¹⁴⁷ have been crystallographically characterised. One report suggests that reaction of NaS2CNR2 with TiCl3 in EtOH affords [Ti(S₂CNR₂)₃] as brown- or yellow-green solids, ¹⁴⁹ but there are no later mentions of this type of complex in the literature. Thus, the stability of [Ti(S₂CNR₂)₃], their potential disproportionation to [Ti(S2CNR2)2] and [Ti(S2CNR2)4], and structure and stability of the former remain issues that require clarification. Interestingly, it has recently been established that dithiocarbamates can coordinate to Ti(0), thus, oxidation of $[Ti(CO)_6]^{2-}$ by thiuram disulfides affords $[Ti(CO)_4(S_2CNR_2)]^{-}$ which adopt an unexpected trigonal prismatic geometry. 150 There may yet be new things to discover in titanium dithiocarbamate chemistry.

The only V(II) dithiocarbamate complex reported is light brown [V(S2CNEt2)2], formed under rigorously oxygen-free conditions upon addition of two equivalents of NaS2CNEt2 to VCl2 in MeCN in a dry box. 143 Unfortunately, no characterising data was given, but it was reported to be soluble in warm 1,2dichlorobenzene, so may be worthy of reinvestigation. In air, addition of [R2NH2][S2CNR2] to VBr2·6H2O affords [V (S2CNR2)3]. The diethyl derivative has been crystallographically characterised 152 and magnetic measurements show a S = 1 ground state with two unpaired electrons. 153 Possibly, [V (S2CNR2)2] are fleetingly formed but readily oxidised, and putative $[V(S_2CNR_2)_2]^+$ reacts with further dithiocarbamate. The reduction chemistry of [V(S2CNEt2)3] has been investigated and reveals a reversible one-electron process. 154 Eight-coordinate V(v) complexes, $[V(S_2CNR_2)_4]$, are accessible from the insertion of CS₂ into [V(NR₂)₄], ¹⁵⁵ but upon heating they undergo intramolecular electron transfer resulting in elimination of thiuram disulfide and formation of [V(S₂CNR₂)₃]. Formal oxidation products of [V(S2CNR)2], namely vanadyl complexes [VO (S₂CNR₂)₂], are easily prepared¹⁵⁶ and have been extensively investigated as potential insulin mimetics¹⁵⁷ and molecular qubits. 158 Thus, akin to the titanium chemistry, even the

Fig. 12 Formation and decomposition of primary amine thiuram disulfides.

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Fig. 13 Formation of [Cr(S₂CNR₂)₂] via reduction of [Cr(S₂CNR₂)₃] and back oxidation.

simple dithiocarbamate chemistry of vanadium requires further investigation.

Chromium bis(dithiocarbamate) complexes are indisputably accessible, but extremely air sensitive, so unless rigorous experimental methods are utilised it is the analogous d³ Cr(III) complexes, [Cr(S₂CNR₂)₃] that are isolated. The first synthesis of [Cr(S2CNEt2)2] was reported by Fackler and Holah in 1966, 162 being described as a pyrophoric yellow-green solid. It has subsequently been prepared from thermolysis of [Cr(CO)₆] and [Hg(S2CNEt2)2].163 In solution [Cr(S2CNEt2)2] is extremely sensitive to air oxidation, converting rapidly to blue [Cr (S₂CNEt₂)₃]. Very few derivatives have been prepared and there are no crystal structure determinations, thus molecular structure(s) of [Cr(S₂CNR₂)₂] remain unknown, although most likely they are polymeric in the solid state, and tetrahedral in the gas phase or solution. Electrochemical studies of [Cr(S₂CNR₂)₃] show that they undergo a one-electron reduction, with the generated [Cr(S2CNR2)3] rapidly losing a dithiocarbamate to give $[Cr(S_2CNR_2)_2]^{164}$ (Fig. 13). However, all attempts to isolate or characterise [Cr(S2CNR2)2] generated in this way have led only to rapid back oxidation and regeneration of [Cr(S2CNR2)3], again highlighting their extreme air-sensitivity.

Non-homoleptic Cr(II) dithiocarbamate complexes can be prepared. For example, room temperature addition of [CpCr (CO)3 and tetraethyl thiuram disulfide affords [CpCr (CO)₂(S₂CNEt₂)] in good yields. 165

(ii) M = Mo, W

We briefly consider the heavier group 6 homologues, [Mo (S2CNR2)2] and [W(S2CNR2)2]. The synthesis and X-ray crystal structure of [Mo(S2CNEt2)2] has been claimed166 but is erroneous: the structure presented is that of [Ni(S2CNEt2)2] with a misassigned metal atom.167 Complexes of the stoichiometry Mo: dithiocarbamate 1:2 are known. Thus, reactions of [Mo₂(μ-OAc)₄] with four equivalents of dithiocarbamate afford [Mo₂(µ-S₂CNR₂)₄] as green solids. However, the latter are thermally unstable and upon heating undergo C-S bond scission to yield Mo(v) thiocarboxamide complexes, [Mo(μ-S)(S₂CNR₂)(SCNR₂)]₂ as shown crystallographically (R = iPr).169 Related W(II) complexes, [W(S2CNR2)2], remain unreported, but both cis-[W (CO)₂(S₂CNR₂)₂] and [W(CO)₃(S₂CNR₂)₂] are known¹⁷⁰ as are the corresponding Mo(II) carbonyl complexes. 171

(iii) $[Mn(S_2CNR_2)_2]$

Bright yellow [Mn(S2CNR2)2] complexes can easily be prepared, but both the reaction and work-up must be carried out under rigorously inert conditions. 172,173 The molecular structure of only one example, namely [Mn(S2CNEt2)2], has been reported (Fig. 14).¹⁷⁴ It is a coordination polymer containing octahedral Mn(II) centres resulting from the dithiocarbamates binding in a bridging manner.

Many papers detail reactions of two equivalents of dithiocarbamates with Mn(II) salts that have been carried out in water and air, and invariably the generated products are dark (often violet-purple) paramagnetic solids formulated as [Mn (S2CNR2)2]. Authors often cite elemental analysis data as proof of this formulation. Indeed, fits with calculated values are often good, albeit in many cases a small amount of bound water (between 0.5 and 2 molecules) are suggested. Recently, this issue has been addressed, and it is now unequivocally established that the dark product described is a Mn(III) oxygen adduct. 175 The precise nature of this adduct is less clear, with the authors suggesting it is monomeric, but a recent crystal structure (Fig. 15) has shown that (at least in part) it is a dimeric oxygen-bridged complex.176

The colour of the oxygen adducts is very similar to that for Mn(III) complexes, [Mn(S2CNR2)3], generated upon addition of three equivalents of dithiocarbamate to Mn(II) salts in air, as is the spectroscopic data. Thus, it is not easy to differentiate between the two. If two equivalents of dithiocarbamate are

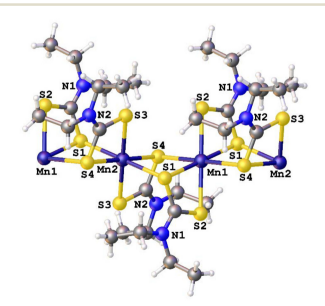


Fig. 14 Part of the polymeric structure of [Mn(S₂CNEt₂)₂].¹⁷⁴

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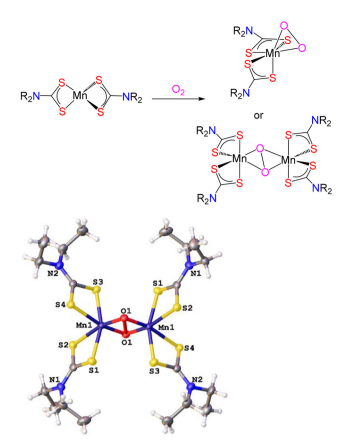


Fig. 15 Oxidation of [Mn(S₂CNR₂)₂] with proposed Mn(III) adducts and the molecular structure of [Mn₂(S₂CNEt₂)₄(μ -O₂)].^{175,176}

added to Mn(II) salts in water under argon then heavy yellow precipitates result, which are unequivocally [Mn(S₂CNR₂)₂]. Further, later exposure to oxygen (bubbling through solution works best to ensure full oxidation) reliably affords the pure oxygen adduct. Addition of more dithiocarbamate salt to the latter gives [Mn(S₂CNR₂)₃], although it is hard to know exactly when the reaction is complete and to ensure that it is, a slight excess of dithiocarbamate is recommended. In the salt of the sa

Manganese dithiocarbamate complexes are becoming widely used as single source precursors (SSPs) to manganese sulfide nanomaterials. In this respect, the accidental use of the oxygen adduct may be advantageous. Thus, oxygen adducts have a higher manganese content than $[Mn(S_2CNR_2)_3]$ and, as shown by thermogravimetric analysis (TGA) for the diethyl derivatives (Fig. 16), decompose at lower temperatures. Indeed, TGA is a good way of differentiating between these two Mn(III) dithiocarbamate complexes. Finally, we note that $[Mn(S_2CNR_2)_3]$ decomposes in a two-step process, the former being associated with loss of a dithiocarbamate and *in situ* formation of $[Mn(S_2CNR_2)_2]$, thus negating the need to isolate air sensitive Mn(III) complexes.

An early application of dithiocarbamates was as fungicides, with manganese containing Maneb© being widely used. It is formed from the reaction of $Mn(\pi)$ salts with Nabam©, the bis (dithiocarbamate) generated from ethylene diamine (Fig. 17).

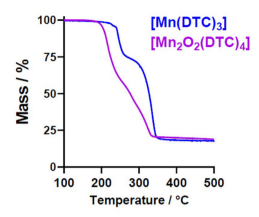


Fig. 16 TGA data for $[Mn(S_2CNEt_2)_3]$ and $[Mn_2(S_2CNEt_2)_4(\mu-O_2)]$. ¹⁷⁶

More recently it has been linked to the development of Parkinson's disease^{181,182} and since 2009 has been banned in the European Union, a fate also experienced by the zinc complex, Zineb®, which was also previously used as a pesticide.

Maneb© and Zineb© were used in agriculture for many years and are proposed to have polymeric structures, as has the double hydration product of Maneb© namely [Mn (S₂CNHC₂H₄NHCS₂)·2H₂O]_n. Recently a good quality molecular structure of the dimethylformamide (dmf) adduct [Mn (S₂CNHC₂H₄NHCS₂)·2dmf]_n was reported¹⁸³ (Fig. 18) and confirms this. Manganese is in the +2 oxidation state but, importantly, has an octahedral coordination geometry with coordination of two *cis* dmf molecules (there are another two non-coordinated dmfs not shown). Likely hydrated Maneb© has a very similar structure except with bound water. Another interesting observation is that while pure samples of this adduct and Maneb© are yellow, the latter ages over time, becoming dark.

(iv) $[Fe(S_2CNR_2)_2]$

Like their manganese counterparts, [Fe(S₂CNR₂)₂] can be prepared and isolated but are extremely air sensitive and should be handled in a dry box or a high quality Schlenk line environment. When prepared under the correct conditions they are described as chocolate brown, ¹⁶² pink ¹⁴² or red ¹⁸⁴ solids. Crystal structures of two polymorphs of [Fe(S₂CNEt₂)₂] ^{185,186} show that it adopts a dimeric structure in the solid state (Fig. 19). Each iron centre is 5-coordinate, being best described as a distorted trigonal bipyramid. No other molecular structures have been determined, but there is some early evidence from Mössbauer studies that some derivatives may be coordination polymers. ^{187,188}

As with the manganese chemistry described above, many authors describe the purported synthesis of Fe(n) bis(dithiocarbamate) complexes in air, and thus their assignment is clearly wrong. Unlike the manganese chemistry, however, it is less clear exactly what they have made if only two equivalents of dithiocarbamate are used. Thus, oxygen adducts of [Fe

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$$H_2N$$
 NH_2
 $NaOH, CS_2$
 Na

Fig. 17 Formation of Nabem@ and reaction with Mn(II) salts to produce Maneb@.

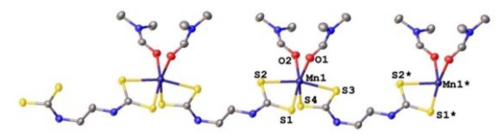


Fig. 18 Part of the polymeric structure of [Mn(S₂CNHC₂H₄NCS₂)·2dmf]_n. ¹⁸³

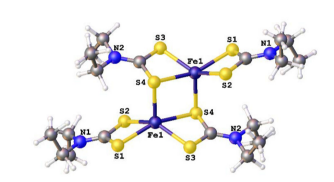


Fig. 19 The molecular structure of [Fe(S₂CNEt₂)₂]₂. ¹⁸⁵

 $(S_2CNR_2)_2]$ have not been reported. These complexes do undergo rapid addition, for example addition of bis(perfluoromethyl)-1,2-dithietene affords Fe(III) complexes (Fig. 20) showing how prone they are to oxidation. It is also known that [Fe(S_2CNR_2)₂] react with excess dithiocarbamate to afford yellow [Fe(S_2CNR_2)₃], which are also extremely air sensitive, being rapidly oxidised to brown Fe(III), [Fe(S_2CNR_2)₃]. Thus, it is most likely that reaction of two equivalents of dithiocarbamate with Fe(II) salts in air simply gives reduced yields of [Fe (S_2CNR_2)₃] along with some unreacted Fe(II) salt.

Bis(dithiocarbamate) complexes [Fe(S₂CNR₂)₂] also bind neutral donor ligands, as shown in crystal structures of a dmf adduct of the morpholine-dithiocarbamate complex (Fig. 21a)¹⁹¹ and a DABCO adduct of [Fe(S₂CNEt₂)₂] which cocrystallises with C_{60}^{144} (Fig. 21b). Thus, from aqueous solutions, it may be that a species [Fe(S₂CNR₂)₂·H₂O] precipitates out. Interestingly, an NMR spectrum of [Fe(S₂CNC₄H₈O)₂·dmf] run in CDCl₃ was found to be identical to that of paramagnetic [Fe(S₂CNC₄H₈O)₃], suggesting oxidation and ligand arrangement upon dissolution in this weakly acidic solvent. ¹⁹¹ Thus, it might be that Fe(II) is reducing enough to convert H⁺ into H₂ with concomitant formation of Fe(III).

The Fe(II) oxidation state can be also stabilised by binding of carbonyls in cis-[Fe(CO)₂(S₂CNR₂)₂]^{192,193} and NO in [Fe(NO) $(S_2CNR_2)_2$]. ^{194,195} Indeed, the binding of NO to $[Fe(S_2CNEt_2)_2]$, as determined by the distinctive EPR spectrum of [Fe(NO) (S₂CNEt₂)₂], is an accepted way of determining the presence of this important signalling gas. 196-198 It is noteworthy, however, that reaction of NO with [Fe(S2CNEt2)3] also affords [Fe(NO) (S₂CNEt₂)₂] (together with thiuram disulfide as a byproduct). Photolysis of [Fe(S2CNEt2)3] also results in elimination of thiuram disulfide and formation of [Fe(S₂CNEt₂)₂], which can be trapped with bis(diphenylphosphino)ethane (dppe) to afford $[Fe(\kappa^2-dppe)(S_2CNEt_2)_2]^{199}$ This suggests that there is a fine redox balance between the reducible Fe(III) centre and the oxidisable dithiocarbamate. Indeed, we have found using in situ EXAFS, that upon warming [Fe(S₂CNⁱBu₂)₃] in oleylamine at 60 °C, reductive elimination of thiuram disulfide results with concomitant formation of the corresponding Fe(II) complex. 193 Further, TGA measurements show that cis-[Fe(CO)₂(S₂CNR₂)₂] lose both carbonyls at relatively low temperatures to afford [Fe(S₂CNR₂)₂]. Thus, [Fe(S₂CNR₂)₂], [Fe (S2CNR2)3] and cis-[Fe(CO)2(S2CNR2)2] are all effectively equivalents when used as SSPs 193 thus negating the need to isolate air sensitive [Fe(S2CNR2)2].

Neither $[M(S_2CNR_2)_2]$ (M = Ru, Os) appear in the reliable literature, with both metals favouring the M(III) state [M (S₂CNR₂)₃], but like iron, carbonyl-stabilised M(II) complexes cis-[Ru(CO)₂(S₂CNR₂)₂]²⁰⁰ and cis-[Os(CO)₂(S₂CNR₂)₂]^{201–203} are accessible.

(v) $[Co(S_2CNR_2)_2]$

The theme continues into the chemistry of cobalt. Like iron and manganese, cobalt forms stable M(III) tris(dithiocarbamate) complexes [Co(S₂CNR₂)₃], the low spin d⁶ electronic configuration allowing NMR characterisation. $^{204-206}$ In contrast, while the corresponding M(II) species [Co(S₂CNR₂)₂] are accessible, they are extremely air-sensitive and paramagnetic, with [Co(S₂CNEt₂)₂] being described as a dark green-brown solid. 142

 R_2N R_2N

Fig. 20 Reactions of [Fe(S₂CNR₂)₂] with a dithietane resulting in oxidation, and addition of further dithiocarbamate followed by oxidation.

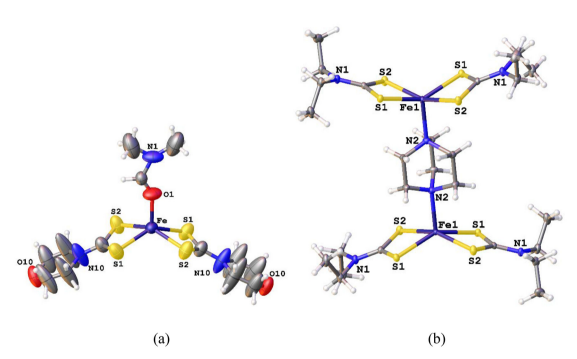


Fig. 21 Molecular structure of (a) $[Fe(S_2CNC_4H_8O)_2\cdot dmf]^{191}$ and (b) $[\{Fe(S_2CNEt_2)_2\}_2DABCO]$ as part of a large structure with $C_{60}\cdot 2DABCO$.

It is telling that there are no crystal structures of [Co(S₂CNR₂)₂] and examples of authentic syntheses and characterisation are rare. Most authors report the formation of green precipitates from reactions of dithiocarbamate salts with CoCl₂·6H₂O in air, *i.e.* they actually generate [Co(S₂CNR₂)₃]. Recently, Khitrich and co-workers provided some evidence for the formation of [Co(S₂CNR₂)₂],²⁰⁷ suggesting that, provided pH is kept between 6–7, then even in air aqueous solutions of [CoCl₂] react with two equivalents of NaS₂CNR₂ to afford [Co(S₂CNR₂)₂] and note that in the solid state they are air stable. Good elemental analysis data are presented, and powder X-ray diffraction studies reveal (for those that are not amorphous) the absence of [Co(S₂CNR₂)₃]. Further, magnetic moments of 2.19–2.45 BM suggest that they have one unpaired electron, consistent with a square planar geometry. Interestingly, when dissolved in

organic solvents they oxidise rapidly with formation of [Co $(S_2CNR_2)_3$] and [Co(OH)₂] (Fig. 22). Clearly more work in this area is warranted.

Electrochemical reduction of $[Co(S_2CNR_2)_3]$ leads to dithiocarbamate loss to give $[Co(S_2CNR_2)_2]$, but back oxidation is facile. Chemical reduction of $[Co(S_2CNR_2)_3]$ by excess Zn/Hg in CH_2Cl_2 results in a slow (ca. 3 h) darkening of the green solution and when canulated under nitrogen onto PhI = NTs, the Co(II) complexes react via insertion into the Co-S bonds to afford crystallographically characterised $[Co\{TsNSC(NR_2)SNTs\}_2]^{209}$ (Fig. 23). Thus, it is chemically possible to prepare $[Co(S_2CNR_2)_2]$ in organic solvents but oxygen must be rigorously excluded.

We note that there are no authentic examples of [Rh $(S_2CNR_2)_2$] or [Ir $(S_2CNR_2)_2$] complexes, ^{210,211} all purported

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3
$$R_2N$$
 O_2 , H_2O organic solvent O_2 , O_2

Fig. 22 Proposed decomposition of [Co(S₂CNR₂)₂] in organic solvents.²⁰⁷

Fig. 23 Reductive generation of [Co(S₂CNR₂)₂] and trapping via multiple NTs insertion in the Co-S bonds.

syntheses being likely that of the analogous [M(S₂CNR₂)₃] complexes. 212,213 Electrochemical reduction of [Rh(S₂CNR₂)₃] is irreversible and shown to be a two-electron process with formation of [Rh(S2CNR2)2] and dithiocarbamate being suggested.214

Structure and stability of dithiocarbamate complexes

(i) $[M(S_2CNR_2)_2]$ (M = Mn, Fe, Co) are all square planar

In solution or the gas phase, the molecular structures of group 10 bis(dithiocarbamate) complexes and [Cu(S₂CNR₂)₂] are square planar. In the solid state, the 4-coordinate square planar arrangement is maintained for $[M(S_2CNR_2)_2]$ (M = Ni,Pd, Pt), but the precise metal coordination environment for copper varies with substituents due to secondary Cu···S interactions.^{8,13} The solution and gas phase structures of [M $(S_2CNR_2)_2$ (M = Mn, Fe, Co) remain unknown, although some authors suggest that all are square planar. This seems highly unlikely for [Mn(S₂CNR₂)₂] as for a d⁵ electronic conformation one would not expect the difference in crystal field stabilisation energies (CFSEs) between tetrahedral and square planar geometries to override steric effects, even when the preferred bite angle for dithiocarbamates is less than 90°. An early communication suggested that, in ethanol, [Mn(S2CNEt2)2] adopts a spin quartet state, i.e. has 3 unpaired electrons, 215 although it was later shown that this result was erroneous, as measurements had been made on a partially oxidised sample. Upon rigorous exclusion of air, magnetic measurements on [Mn (S2CNEt2)2 and other derivatives show that they adopt a ground state with 5 unpaired electrons. 172 Thus, it seems likely that [Mn(S2CNR2)2] have a distorted tetrahedral coordination geometry, although a recent DFT study suggests a squareplanar arrangement, which may be a local minimum.²¹⁶ In a similar vein, iron complexes, [Fe(S₂CNR₂)₂], with a d⁶ electronic configuration have been shown to have 4 unpaired electrons, 186 being inconsistent with a square-planar array. Nevertheless, recent DFT calculations propose a local FeS₄ geometry with D_{2h} symmetry, consistent with a square planar geometry. 217 Clearly, accurate calculations are required to corroborate the actual coordination geometry. The absence of any crystallographic data makes it harder to predict the molecular geometry of d^7 [Co(S₂CNR₂)₂], but they are likely square-planar, as supported by the magnetic susceptibility measurements and DFT calculations. 218

(ii) Ru(III) complexes [Ru(S₂CNR₂)₃] and [Ru(S₂CNHR)₃] are diamagnetic

Iron tris(dialkylDTC) complexes, [Fe(S2CNR2)3], have been widely studied as spin crossover complexes as they can adopt either high spin (HS) ⁶A₁ or low spin (LS) ²T₂ electronic configurations, the position of the equilibrium being affected by the nature of the alkyl-substituents and external factors such as temperature and pressure. 219-222 Receiving less attention are analogous Ru(III) complexes, [Ru(S2CNR2)3], although a number of early papers established their fluxional distorted octahedral structures²²³ and paramagnetic nature, 224,225 the latter being as expected for a d⁵ electronic configuration. Nevertheless, several publications claim to have prepared diamagnetic Ru(III) tris(dithiocarbamate) complexes of both secondary and primary amines, characterised by (almost perfect) elemental analysis data and ¹H NMR peaks in the standard 0-10 ppm range. 226-228 Interestingly, ¹H NMR spectroscopy can be used to part-characterise [Ru(S₂CNR₂)₃], although peaks are observed across a ca. 40 ppm range, as expected for a weak (low spin) paramagnetic system. 229-231 Diamagnetic complexes can be formed from the reaction of RuCl3 and dithiocarbamates, indeed many studies have reported mixtures of paramagnetic $[Ru(S_2CNR_2)_3]$ and diamagnetic $[Ru_2(S_2CNR_2)_5]^+$ (which can exist in two isomeric forms) from these reactions. 229-232 Thus, it may be that such diamagnetic cations are the real products of the reactions described above. 226-228 Akin to their ruthenium analogues, osmium complexes, [Os

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(S₂CNR₂)₃], are also low spin d⁵ systems that show resonances across a wide chemical shift range in their ¹H NMR spectra and are easily oxidised.233

(iii) Stable dithiocarbamate complexes of high-valent metal oxides are always accessible

One of the major benefits of the dithiocarbamate ligand is its ability to stabilise metals in a wide range of oxidation states, being (in simple terms) attributed to the accessibility of dithiocarbamate and thioureide resonance forms. Thus, the former is a relatively soft (strong field) and the latter a hard (weak field) donor. While not unique to dithiocarbamates, some ligand sets, for example the related xanthates (ROCS₂⁻), cannot adopt the latter form as the oxygen is too electronegative to act as an electron-pair donor. 234 Adoption of the thioureide form also accounts for the partial double bond character of the backbone C-N vector which leads to hindered rotation about it. 235 Thus, rotamers of complexes $[M(S_2CNR^1R^2)_2]$ (M = Ni, Pd) can be distinguished, and their rate of interconversions measured.236 The thioureide form can also account for the stability of high-valent species such as of Mo(vi) and W(vi) complexes.^{8,9} However, one should not forget that the dithiocarbamate is redox active, being relatively easily oxidised, and thus when bound to a highly oxidised metal centre, the system is on a cliff edge, as beautifully illustrated by a series of publications from Stiefel and co-workers. 237-241

While a wide range of metal centres and oxidation states can be tolerated by dithiocarbamates,8 this does not extend to the most oxidising. For example, reaction of dithiocarbamate salts with potassium dichromate, K₂[Cr₂O₇], affords a mixture of two Cr(III) products: ring-expanded [Cr(S₂CNR₂)₂(OS₂CNR₂)] as the major component together with $[Cr(S_2CNR_2)_3]$ (R = Me, Et)²⁴² (Fig. 24). Thus, (at least formally) three equivalents of dithiocarbamate act as one-electron reducing agents and the other three as ligands. The precise mode of formation of the ring-expanded complex is not clear. In related work, Farmer and co-workers showed that addition of H_2O_2 to $[Ru(\kappa^2-2,2'-1)]$ bipy)₂(S₂CNMe₂)]⁺ afforded both ring-expanded sulfur-oxidised isomers but they did not interconvert, suggesting that they are formed via two separate pathways, 243 although in other work they showed that [Zn(S2CNEt2)(OS2CNEt2)] was a product of the oxygenation of [Zn(S2CNEt2)2], a transformation that can be reversed upon addition of a phosphine.²⁴⁴ Interestingly, reaction of chromate with excess dithiocarbamate has been repurposed as an analytical method of measuring relative amounts of Cr(III) and Cr(IV). 245-247 Thus, Cr(III) reacts to

form [Cr(S₂CNR₂)₃] and Cr(v₁) a mixture of this and the ringexpanded products. Importantly the two products have different chromatographic retention times and thus by understanding the precise ratio of products formed from the Cr(vi) reaction it is possible to determine relative amounts of Cr(III) and Cr(v₁), although not all publications seem to do it this way, with some suggesting that Cr(III) is unreactive towards dithiocarbamates.

Likely dithiocarbamates cannot stabilise high oxidation states such as Os(viii) and Re(vii), since these metal centres are powerful oxidising agents. There are reports of Os(v1) dithiocarbamate complexes, trans-[OsO₂(S₂CNR₂)₂], ^{248,249} which show distinctive vibrations at 839 and 888 cm⁻¹ in the IR spectrum, being assigned to asymmetric and symmetric trans-OsO₂ vibrations respectively.²⁴⁹ Further investigations are needed to unequivocally establish the validity of these claims, but they seem quite likely to be correct given that the trans-[Os(vi)O2] moiety is particularly stable, being found for example in complexes such as trans- $[OsO_2(CO)_4]^{2+}$ and trans- $[OsO_2(OH)_4]^{2-}$.

(iv) Au(III) complexes, [Au(S₂CNR₂)₂]⁺ and [AuCl₂(S₂CNR₂)], are easily prepared in their pure forms

Gold(III) dithiocarbamate complexes of the $[AuX_2(S_2CNR_2)]$ (where X = Cl, Br, I) have been widely studied as potential anti-cancer metallo-pharmaceuticals. Structurally they are comparable to *cis*-platin and the mechanism of action is suspected to be similar. 250 These compounds were first synthesised in 1964 by oxidation of the Au(1) complex [Au $(S_2CNR_2)_n$ with elemental X_2 .²⁵¹ This supposedly afforded the Au(III) dithiocarbamate [AuX₂(S₂CNR₂)] as a single compound in solution. Alternatively, the addition of one equivalent of dithiocarbamate to Au(III) halides is claimed to result in the sole formation of [AuX2(S2CNR2)]. In a similar fashion, addition of two equivalents of dithiocarbamate to Au(III) halides supposedly resulted in the sole formation of the cationic complex [Au(S2CNR2)2]+.9 Often in this system X is invoked as the counterion, but a haloaurate species is more likely to be present.

We recently investigated the [AuX₂(S₂CNR₂)] system thoroughly and discovered that the established reactivity is not correct. The series of compounds $[AuX_2(S_2CNR_2)]$ where X = halide cannot be isolated as a single species in solution; instead, they exist in an equilibrium with [Au(S2CNR2)2][AuX4] (Fig. 25). This equilibrium is present irrespective of the method of synthesis, or the ratio of dithiocarbamate to Au(III). A careful look at the older literature reveals that this equili-

Fig. 24 Cr(III) complexes from reaction of dithiocarbamates and K₂[Cr₂O₇].

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brium has always been present even though it had not previously been acknowledged. For example, the ¹H and ¹³C{¹H} NMR spectra of dibenzyl-dithiocarbamate- and methylsarcosine-dithiocarbamate-derived Au(III) complexes both clearly show the presence of two dithiocarbamate environments, one for the neutral species and one for the ion pair. 253,254 Mass spectrometry of [AuX₂(S₂CNR₂)] revealed that the cation [Au(S₂CNR₂)₂]⁺ was also present in the sample, although it was explained as decomposition in solution.²⁵⁵ A single crystal containing [AuCl₂(S₂CNiPr₂)], two [Au (S₂CNiPr₂)₂]⁺ cations and two [ClO₄]⁻ anions was obtained, showing that both components of the equilibrium were present in a single reaction mixture.²⁵⁶ Recently, it was shown that an equilibrium mixture where [Au(S2CNiPr2)2][AuCl4] was the majority product could be converted to a mixture where [AuCl₂(S₂CNiPr₂)] was the majority product by refluxing in acetone.257

We note here that it is possible to obtain $[Au(S_2CNR_2)_2]^+$ as a single species in solution, but the counterion must be chosen with care and haloaurate anions should be avoided. For example, use of a weakly coordinating anion such as [BF₄] led to the isolation of [Au(S2CNR2)2][BF4] which showed no sign of existing as an equilibrium, even upon standing in solution for several weeks. However, as soon as the anion was metathesized to [AuCl₄]⁻, the equilibrium restarted.²⁵² In a similar vein, it is possible to isolate [AuX₂(S₂CNR₂)] as a single pure product in solution by avoiding the use of halide ions: stable systems are known to exist for X = Me, C_6F_5 , mesityl and thiolate, for example. 258-261

The above notwithstanding, it is possible to fractionally crystallise [AuX₂(S₂CNR₂)] as a single compound. The neutral species forms orange crystals (in contrast to the ion pair, which is yellow); these are indefinitely stable in the solid phase, but as soon as the crystals are dissolved in an organic solvent then the equilibration process resumes. Researchers are urged to be cautious here by not mistaking this for a single pure product in solution.²⁶¹ It is also noteworthy that elemental analysis is not a useful characterization technique since the two components of the equilibrium, namely $[AuX_2(S_2CNR_2)]$ and $[Au(S_2CNR_2)_2][AuX_4]$, have identical empirical formulae and cannot be used to distinguish the two isomers.262

(v) Primary amine-derived dithiocarbamate complexes are always stable

The majority of dithiocarbamate chemistry focuses on those derived from secondary amines. There are several reasons for

this, one being their greater stability vs. those derived from primary amines. Thus, while all dithiocarbamates are unstable in acidic media, the relatively acidic nature of the unique proton in RNHCS₂ also makes them prone to instability in basic media. Nevertheless, there are some relatively simple and highly reproducible syntheses of primary amine dithiocarbamate salts. Examples of these are group 10²⁶³⁻²⁶⁵ and group $12^{266-269}$ $[M(S_2CNHR)_2]$ complexes, but also (S₂CNHR)₃|^{270,271} and ^{99m}Tc complexes with radiopharmaceutical applications.^{272–274} The authenticity of homoleptic primary amine dithiocarbamate complexes of other metals is far less certain and likely the majority are too unstable to be isolated.

Upon metal binding the acidity of the backbone proton is retained, its removal generating the dianionic dithiocarbimate (or imidomethanedithiolate) ligand, S2C=NR2-. Such ligands form complexes with a wide range of metals, 103 being especially prevalent for Group 10 elements and when the substituent is strongly electron-withdrawing, for example aryl or cyanide. For Group 10 elements, as discussed earlier (Fig. 8), double deprotonation upon addition of a strong base of [M (S₂CNHAr)₂] and [M(S₂CNHR)₂] affords the corresponding dithiocarbimate dianions that, provided they are kept under anhydrous and oxygen-free conditions, have significant lifetimes and can be used to prepare functionalised dithiocarbamate derivatives. Stability of the dithiocarbimate complexes (presumably) results from delocalisation of the excess electron density into the vacant d-orbital. However, this is not the case for most other metals. For example, addition of base to Group 12 complexes $[M(S_2CNHR)_2]$ and $[M(S_2CNHAr)_2]$ (M = Zn, Cd)results in rapid decomposition to give ZnS or CdS nanomaterials. 266-268 Indeed, this is advantageous, as is the case for [Ni(S2CNHR)2]275 as they provide low temperature routes to these nanomaterials (Fig. 26).

For organic chemists reading this it may ring bells as the base-induced decomposition of primary amine dithiocarbamates to afford organic isothiocyanates is a well-developed preparative method. 276-278 Indeed, often a metal ion is added to facilitate the process. Thus, such complexes are not stable, certainly under basic conditions, rapidly decomposing to afford the isothiocyanate. Nevertheless, there are (purported) examples of $[Fe(S_2CNHR)_3]^{279,280}$ and $[Cu(S_2CNHR)_2]^{281-285}$ in the literature, some of the latter being touted as enzyme inhibitors.285 We have recently looked more closely at both systems in our laboratory and will publish our results in due course. The iron system is complex and is not appropriate to discuss here, but [Fe(S2CNHR)3] have at best a fleeting stabi-

$$\begin{bmatrix} R_2 N & & & \\ & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

Fig. 25 Equilibrium between [Au(S₂CNR₂)₂][AuCl₄] and [AuCl₂(S₂CNR₂)].

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Fig. 26 Base-induced decomposition of [M(S₂CNHR)₂] to afford RNCS and nanoscale metal sulfides.

Fig. 27 Formation of [Cu(S₂CNHR)]_n and subsequent reaction with PPh₃

lity. Reactions of copper(II) salts with a range of primary amine derived dithiocarbamates in water leads to isolation of high yields of bright yellow polymeric Cu(1) complexes, [Cu $(S_2CNHR)_n$ (R = Bu, Cy) along with the corresponding thiourea.²⁸⁶ Their poor solubility hinders full characterisation, but simple addition of a slight excess of PPh3 affords the soluble and crystallographically characterised complexes, [Cu $(S_2CNHR)(PPh_3)_2^{286}$ (Fig. 27). In this regard we note that [Cu (S₂CNHPh)(PPh₃)₂] has previously been prepared from reaction of [Cu(BH₄)(PPh₃)₂] with PhNCS.²⁸⁷ Thus, primary amine dithiocarbamates are stable at a Cu(I) centre but not Cu(II).

We briefly note here that [NH₄][S₂CNH₂] is easily prepared^{288,289} and has been used to generate a range of stable homoleptic complexes, including crystallographically characterised: $[Ni(S_2CNH_2)_2]$, 290 $[Zn(S_2CNH_2)_2]$, 291 $[Cr(S_2CNH_2)_3]$, 292 $[Co(S_2CNH_2)_3]$, 288,293 $[Rh(S_2CNH_2)_3]$, 288 $[Ag(S_2CNH_2)]$, 294 [Au(S₂CNH₂)],²⁹⁵ [Au(S₂CNH₂)₂][SCN],²⁹⁶ [Cu(S₂CNH₂)]²⁹⁷ and [Cu (S₂CNH₂)₂[NH₄].²⁹⁷ Considering the discussion above, the finding that this ligand is only stabilised at the Cu(1) centre is interesting. Further several non-homoleptic Ru(II) derivatives have been prepared and crystallographically characterised²⁸⁸ suggesting that there is a relatively rich coordination chemistry of this ligand still to be explored.

(v) Complexes with two different dithiocarbamate ligands and dithiocarbamate-containing mixed-ligand complexes are (easily) accessible in a pure state

Many homoleptic dithiocarbamate complexes are known, being easy to prepare even in the most basic of laboratory

settings.^{8,9} In developing this chemistry further, it is tempting to consider preparing related complexes in which two (or more) different dithiocarbamates bind to a single metal centre, thus providing easy tuning of chemical and physiochemical properties. Consequently, over the past decade, there has been an increasing number of publications claiming the formation of pure complexes containing two different dithiocarbamate ligands of Ni(II), 298 Cu(II), 299–301 zinc and mercury³⁰²⁻³⁰⁶ and other transition metal³⁰⁷⁻³⁰⁹ and main group³¹⁰⁻³¹⁵ elements. In all these reports, the simple addition of equimolar equivalents of two different dithiocarbamates to a metal salt is reported to (cleanly) afford the mixed-ligand complex, without any explanation as to why a mixed-ligand complex might be thermodynamically preferable to either the two homoleptic complexes or a statistical mixture of homoand heteroleptic complexes. Very few of these reports are supported with single crystal X-ray data, or compelling characterising data. Crystal structures of [Hg(S2CNMePh)(S2CNEtPh)]304 and [Zn(S2CNMePh)(S2CNEtPh)(2,2'-bipy)]305 are in the literature but in both there is a significant disorder of the Me/Et groups. The only good quality (non-disordered) example of a molecular structure of a heteroleptic dithiocarbamate complex is that of the anion [Cd(S2CNPr2)2(S2CNMeBu)], formed (in a similar manner to other examples) upon addition of pyrrolidinium salts of [S2CNMeBu] to [Cd(S2CNPr2)2].316 However, while the crystal structure shows a well-ordered mixed-dithiocarbamate complex, both ¹H and ¹³C{¹H} NMR spectra show that in solution multiple products are present, resulting from ligand-exchange. Thus, while complexes with two different

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dithiocarbamates are accessible as part of a mixture and can (at least potentially) be selectively crystallised out, they are not accessible in pure form in solution.

For diamagnetic species, NMR spectroscopy should be a simple way of probing such equilibria, however, the (relatively) slow timescale coupled with low intensity of the quaternary backbone carbon signal in the 13C(1H) NMR spectrum and its insensitivity to substituent changes 317 renders it unsuitable. In contrast, HPLC has been shown to be an excellent analytical tool for investigating these systems, being used effectively, for example, by Moriyasu and Hashimoto, 318-320 Liska and coworkers^{321,322} and others.³²³ Thus, Morivasu and Hashimoto mixed homoleptic bis(dithiocarbamate) complexes of nickel and copper (which can't easily be studied by NMR spectroscopy) and in all cases found that they were labile and in equilibrium with the mixed-ligand counterpart (Fig. 28). Rate constants depend on both metal and substituents, for Ni(II) being of the order of 10^1 – 10^2 M⁻¹ s⁻¹ and for Cu(II) slower at ca. 10^3 M⁻¹ s⁻¹. 317-320 These studies were later extended to other metals and in all cases, the dithiocarbamates were found to be labile with equilibrium constants being ca. 4, i.e. a statistical mixture of the three complexes.³²⁴ Similar equilibria have also been noted between homoleptic nickel complexes of primary and secondary amines, affording [Ni(S2CNR12) (S₂CNHR²)], ³¹⁸⁻³²⁰ and also between [Ni(S₂CNR₂)₂] and xanthate complexes, [Ni(S2COR)2].325 Most studies support a bimolecular exchange process likely proceeding through a dimeric intermediate, the form of which is common in crystallographic studies of [Zn(S₂CNR₂)₂] and [Cu(S₂CNR₂)₂].⁸ Rapid dithiocarbamate exchange has also been noted at Fe(III)³²⁶⁻³²⁸ and Hg(II)^{329,330} centres.

For low spin d⁶ [Co(S₂CNR₂)₃] complexes, the CFSE is so high that exchange rates are slow331 and isolation of mixedligand complexes is possible. Thus, addition of a dithiocarbamate salt, NaS₂CNR¹₂ to [Co₂(S₂CNR²₂)₅]⁺ affords a mixture of $[Co(S_2CNR^1_2)(S_2CNR^2_2)_2]$ and $[Co_2(S_2CNR^2_2)_3]^{332}$ (Fig. 29). Further, adding equimolar amounts of two different dithiocarbamates to Co(III) salts in water affords a statistical mixture of products,³³³ and a similar mixture of products can be obtained upon heating equimolar amounts of [Co(S2CNEt2)3] and [Co (S₂CNⁱPr₂)₃] either in the solid-state or at 155 °C in chloronaphthalene for 4-5 h. 333 Importantly, these mixtures can be fully separated by column chromatography allowing their individual characterisation. Interestingly, while it is hard to differentiate them by ¹H or ¹³C{¹H} NMR spectroscopy, they can be distinguished by ⁵⁹Co NMR spectroscopy and mass spectrometry and have tuneable oxidation potentials. 333

The ease of accessibility of mixed-ligand complexes containing dithiocarbamates also needs some careful consideration. There are many such examples^{8,9} but for solution stability in their pure form they need to feature either a relatively non-labile metal centre with either bulky non-dithiocarbamate ligands that preclude formation of a bimetallic intermediate or secondary inter-ligand interactions (e.g. hydrogen-bonding). Following on from the discussion above, Co(III) is a good example of a nonlabile metal centre. Thus, $[Co_2(S_2CNR_2)_5]^+$ reacts with a range of nucleophiles to afford [Co(S2CNR2)3] and a mixed-ligand complex, a recent case being the formation of Co(III) bis(dithiocarbamate)dithiolane complexes from addition of dithiones (Fig. 29).³³⁴ Other examples of a high CFSE leading to stable mixed-ligand complexes relate to the iron dithietane complexes¹⁸⁹ (Fig. 20) which exist in temperature-dependent spinequilibrium between the singlet (S = 0) ground state and a lowlying triplet (S = 1) excited state. Thus, not only are these mixedligand complexes stable, but their NMR spectra are also accessible.

However, simultaneously adding a dithiocarbamate and a different ligand to a metal salt will not automatically afford the mixed-ligand species as has been suggested in several publications.335-338 Just as for complexes with two different dithiocarbamates, it is hard by simple spectroscopic methods to tell the difference between a 1:1 mixture of two homoleptic complexes and a mixed-ligand species, and in many cases the three likely co-exist. A recent publication highlights this nicely (Fig. 30).³³⁹ Thus, while insertion of an imide group into a Ni-S bond of [Ni(S2CNR2)2] affords mixed-ligand complexes, examples of which can be crystallographically characterised,

Fig. 28 Equilibrium between homoleptic and mixed ligand bis(dithiocarbamate) complexes

$$\begin{array}{c} R^{1}{}_{2}N \\ \\ S \\ \\ R^{1}{}_{2}N \\ \\ \end{array} \longrightarrow \begin{array}{c} R^{1}{}_{2}N \\ \\ \\ \\ \end{array} \longrightarrow \begin{array}{c} R^{1}{}_{2}N \\ \\$$

Synthesis of heteroleptic Co(III) dithiocarbamate complexes.

 $[Ni(S_2CNR_2)_2] \xrightarrow{ArN_3} R_2N \xrightarrow{S} NR_2 + R_2N \xrightarrow{S} NR_2 + R_2N \xrightarrow{S} NR_2$

Fig. 30 Formation of mixed-ligand complex in equilibrium with homoleptic complexes

upon dissolution they are shown to be in equilibrium with the two homoleptic complexes.

(vi) Reacting two different dithiocarbamate complexes gives a pure mixed metal product

Given the extremely large number of homonuclear transition metal dithiocarbamate complexes that are known, it is perhaps surprising that dithiocarbamate-bridged heteronuclear complexes are rare. Thus, as shown by Robinson, 340 complexes obtained by successive addition of metal salts MX₂ (M = Zn, Cd or Hg) and $M'Y_2$ (M' = VO, Mn, Fe, Co, Ni, Cu or Zn) to a solution of a sodium dithiocarbamate are not heterobimetallic $[\{M'M(S_2CNR_2)_4\}_n]$ but rather contain a mixture of homonuclear complexes. Likewise, reactions of Pd(II) salts [PdCl₂L₂] $(L = PhCN, L_2 = dppe)$ with a series of homoleptic dithiocarbamate complexes affords palladium-dithiocarbamate cations $[Pd(S_2CNR_2)L_2]^+$ with the second metal being incorporated into the counterion.341-343 More recently, Torimoto and coworkers added Na(S2CNEt2) to mixtures of metal salts and used the ensuing "complex" as a precursor to a range of nanoscale metal sulfides with quantum dot properties. 344-347 While the method is effective and efficient, the suggestion that a mixed-metal complex is formed is erroneous. Rather the precursor "complex" is an intimate mixture of several different complexes. There are some well-authenticated examples of dithiocarbamate-bridged heterobimetallic complexes, being especially prevalent amongst the coinage metals348-351 and there are also a small number of other dithiocarbamate-containing heteronuclear complexes that have been crystallographically characterised. 352,353 Nevertheless, the vast majority of dithiocarbamate complexes contain a single metal type.

6. Dithiocarbamate complexes as single source precursors

Dithiocarbamate complexes find widespread use as single source precursors for a range of nanoscale and thin film metal sulfides. 49–52 Thus, simply heating, either in the solid-state or solution, the complex or mixture of complexes, results in S–C and other bond scission(s) affording the thermodynamically stable metal-sulfide(s) and various organic species, the latter (normally) being easily removed either by washing or evaporation. The utility of this simple approach is that air-stable dithiocarbamate complexes from across the periodic table can be (relatively easily) prepared and stored, and by judicious control of reaction conditions and stoichiometry a wide range of binary, ternary and quaternary metal sulfides can be

accessed. A nice example of this is their use towards the synthesis of $\mathrm{Cu_2ZnSnS_4}$ (CZTS) a quaternary semiconductor with a power conversion efficiency of ca. 10% that has been suggested as a viable material for thin film solar cells. Thus, decomposing mixtures of copper, zinc and tin dithiocarbamates leads to the formation of CZTS thin films or quantum dots. 354,355

However, such transformations are not as simple as they might appear. Firstly, in the solid state, thermogravimetric analysis studies have shown that temperatures of between 200-400 °C are required to cleave the S-C (and other) bonds within dithiocarbamate complexes. 356-359 Decomposition temperatures can be tuned (normally within a small range) upon changing substituents, and there is also a correlation between the ease of thermal decomposition and ionic radius of the metal ion; the smaller the metallic ionic radius, the greater the thermal stability.360 In solution, decomposition temperatures can be significantly lower than in the solidstate. 275,361,362 For example, the water-soluble SSP [Cu{S₂CN (CH₂CO₂H)₂}₂] decomposes at 180 °C in the solid state but at ca. 80 °C in water. 362 Another issue to address when preparing ternary, quaternary or multinary sulfides, is matching of decomposition temperatures, such that all SSPs decompose within a relatively small range, thus ensuring that the different molecular building blocks are all available at the nucleation stage. This is nicely exemplified by O'Brien's synthesis of CZTS. 363,364 Thus, as the decomposition temperature of [Sn (S₂CNEt₂)₂] (174 °C) is significantly lower than those of [Cu $(S_2CNEt_2)_2$ (220 °C) and $[Zn(S_2CNEt_2)_2]$ (240 °C), heating this mixture will lead to the premature formation of tin sulfides and hence the tin precursors was replaced by the Sn(IV) SSPs, [Sn(S₂CNBu₂)₄]³⁶³ and [Bu₂Sn(S₂CNBu₂)₂]³⁶⁴ which decompose at higher temperatures. A study by Torimoto and co-workers nicely highlights the need to carefully control decomposition conditions.³⁶⁵ Thus, for the synthesis of $Ag(In_xGa_{1-x})S_2$ quantum dots, simply heating a mixture of silver, indium and gallium precursors initially results in formation of polydisperse Ag₂S as the silver dithiocarbamate decomposes at lower temperatures than others, and upon further heating as indium and gallium SSPs decompose they give a shell of these sulfides around the Ag₂S (Fig. 31a).³⁶⁶ However, when a silver source is injected into the decomposition reaction then co-nucleation of all components occurs, initially to give a core-shell structure, while further heating affords the desired ternary phase sulfide as quantum dots.³⁶⁵

Given these complications and constraints, it is surprising that there are an increasing number of reports seemingly suggesting that simply preparing mixtures of dithiocarbamates at room temperature results in the formation of metal sulfides **Dalton Transactions**

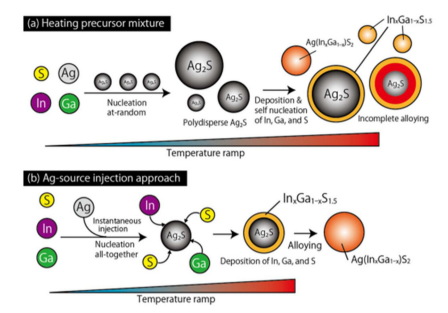


Fig. 31 Schematic illustrations of the synthesis of $Ag(In_xGa_{1-x})S_2$ quantum dots form (a) heating of the precursor mixture and (b) injecting a silver source into the decomposition mixture. 365

i.e. there is no mention of a heating or other decomposition process³⁶⁷⁻³⁷¹ although decomposition appears to be supported by powder X-ray diffraction data. Further, many of these "decompositions" are proposed to give composite mixtures of individual metal sulfides, some of which such as BaS3, are normally prepared at 400 °C. 372 Clearly such reports require some scrutiny and perhaps further explanation(s) from the authors.

Summary and conclusions

In this perspective we have tried to address common misconceptions and errors that we have found in the literature on dithiocarbamates and their complexes. Amongst the challenges faced by those working in this area are the vast number of publications and patents, and the widespread reach of this simple ligand type across many different fields of research. Thus, it is hard for anyone to fully keep up to date and understand all aspects of their chemistry. Likely many misconceptions-errors that have crept into the literature result are further exacerbated by the recent (seemingly unstoppable) proliferation of scientific journals. It is a testament to the wide applicability of dithiocarbamate chemistry that this perspective cites over 200 different journal titles: some of us long for the days when our library reading covered 10-15 journals, each with a clear remit. Thus, we need to understand that newcomers to this mature field of research have a lot of reading and understanding to do to get up to speed with what has gone before. Dithiocarbamate chemistry has been an active area of research for over 150 years and unlike some fields of scientific research, some early papers have relevance today. For example, while recently developing a primary-amine derived dithiocarbamate as a H₂S release-vector we came across a paper from 1891³⁷³ which had effectively already studied this and provided us with invaluable insight into our work.

Having said this, there is also some evidence of tardy work in the dithiocarbamate literature, with researchers either cutting corners, or not being totally honest in reporting their findings-data. Sadly, this appears to be an issue that has proliferated through all aspects of scientific research and one that seriously undermines the efforts of those who play fair. It is hard to understand why anyone would deliberately mis-represent their work, but undoubtably the high speed of modern living and (sometimes) excessive pressures put on academics to publish may be part of the problem. Linking publication to financial gain or making it a pre-requisite for promotion will tempt some to cut corners.

Conflicts of interest

There are no conflicts to declare.

Data availability

There is no data associated with this submission.

Acknowledgements

We thank Dr Jagodish C. Sarker for reading through the manuscript and providing valuable comments, and Dr Shishir Ghosh for attempting to reproduce a reaction discussed herein.

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