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CO₂ and SO₂ activation by a Cr–Cr quintuple bond[†]

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A quintuply bonded dichromium complex stabilized by aminopyridinato ligands activates CO_2 and SO_2 by reducing the (formal) bond order of the metal-metal bonds. Oxygen abstraction is observed during CO_2 activation. SO_2 activation proceeds *via* formation of a unique dithionite complex/coordination. Furthermore, N₂O activation was investigated and the formation of a tetrameric Cr-oxo complex was observed.

 CO_2 and SO_2 are important molecules with, for instance, respect to global warming and acid rain formation, respectively. Thus, their emission has to be reduced with the search for utilization protocols of such molecules. Whereas SO_2 is diversely used in the food industry and for the production of sulfuric acid, the utilization of CO_2 as a C_1 -feedstock is still a highly attractive goal. CO_2 is thermodynamically stable and chemically inert and needs to be activated prior to chemical conversions. Stoichiometric CO_2 activation has been achieved (for instance) using complexes based on *f*-block metals,¹ main group metal compounds,² frustrated Lewis pairs,³ and most intensively using transition metal complexes. For the latter, mononuclear⁴ and bimetallic complexes⁵ have been investigated. In comparison, SO_2 activation has been mainly achieved by mononuclear transition metal complexes.⁶

Stable molecules containing a quintuple bond have been reported for chromium and molybdenum.^{7,8} The bimetallic platform in such quintuply bonded complexes stores 10 electrons and is well suited for small molecule activation.⁹ In addition, the Cr complexes show unusual short metal–metal bonds with the shortest bonds [1.7056(12) Å] matching by length with long C–C bonds.¹⁰

We have a rather long ongoing interest in CO_2 activation¹¹ and quintuple bond reactivity⁹ and report here on the activation of CO_2 and SO_2 by a chromium–chromium quintuple bond.



Scheme 1 Synthesis of complex **2**. In the course of the reaction with CO₂, an oxo-chromium byproduct is formed (*vide infra*).

 CO_2 activation proceeds with C–O-bond cleavage resulting in the formation of a carbonyl complex and a chromium oxo species that differs structurally from the O₂ activation product.^{9d} SO₂ activation leads to dithionite formation. Attempts to activate CO_2 using quintuple bonds have failed so far.^{9g}

The reaction of CO_2 with quintuply bonded chromium dimer 1 (Scheme 1) leads to C=O bond cleavage to give a CO bridged complex (2) (Scheme 1). Complex 2 shows evident reduction of CO₂ to CO. Complex 2 can also be prepared by reaction of 1 with an excess of carbon monoxide in toluene. Following this reaction by NMR, it was found that the formation of 2 is nearly quantitative. Complex 2 is diamagnetic and the ¹H NMR spectrum in benzene- d_6 shows the resonance of the methyl protons of the isopropyl group as two doublets at 1.06 and 1.36 ppm with a coupling constant ${}^{3}J_{\rm HH}$ = 6.8 Hz. The sharp singlet belonging to the methyl group of the 2,6-dimethylphenyl substituents appears at 2.11 ppm along with methyl protons of the co-crystallized toluene molecule. The resonance of the isopropyl CH-protons appears as a septet at 3.13 ppm. In ¹³C NMR, in addition to other signals, the peak for the carbon of the CO ligand is observed at 292 ppm. The infrared spectrum of complex 2 in the range of 4000-400 cm⁻¹ shows no absorption bands in the region $2300-2100 \text{ cm}^{-1}$. The strong absorption bands observed at 1924 and 1806 cm⁻¹ can be attributed to the carbonyl ligands. In solution (toluene) signals at 1942 and 1857 cm⁻¹ were recorded. The signals at higher wave numbers can be attributed to semi-bridged CO due to steric crowding, which is in accordance with the electronic structure of 2 (vide infra).

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Fig. 1 Molecular structure of 2 with hydrogen atoms and one toluene molecule omitted for clarity. Select bond lengths [Å] and angles [°]. Cr1–Cr1A 1.886(19), Cr1–N1 2.011(4), Cr1–N2 2.016(4), Cr1–C1 2.066(8), Cr1–C1 2.092(8), C1–O1 1.154(8), C1–Cr1 2.066(8); Cr1–Cr1A–N1 97.56(13), Cr1–Cr1A–N2 94.04(13), N1–Cr1–N2A 168.39(17), Cr1–Cr1A–C1 63.7(2), N1–Cr1–C1 92.0(2), N2–Cr1–C1 93.5(2), Cr1–Cr1A–C1 62.3(2), Cr1–C1–O1 152.9(6), Cr1–C1–O1 153.2(6), C1–Cr1–C1 126.1(2), O1–C1–Cr1 152.9(6), Cr1–C1–Cr1A 53.9(2).

The molecular structure of **2** is shown in Fig. 1. In **2**, the two Cr atoms are in the plane of the pyridine rings of the aminopyridinato ligands. The two CO ligands are coordinated perpendicular to this plane with a C–O bond length of 1.154(8) Å and Cr–C–O angles of 152.9(6) and $153.2(6)^{\circ}$ (Fig. 1). The Cr–C bond distances of the bridging carbonyl ligand are 2.092(8) and 2.066(8) Å. These values are comparable to Cr–Cr bonded complexes with bridging carbonyls.¹² The Cr–Cr distance of 1.886(19) Å is in the range known for Cr–Cr quadruple bonds but a quadruple bond is not in accordance with the electronic structure of **2** (*vide infra*).

To investigate the potential byproduct resulting from the oxygen released, NMR experiments were carried out. They showed 2 to be the major product along with some broad peaks attributable to a paramagnetic compound. This compound differs from the oxo complex isolated if 1 is reacted with O2.9d To get more insight we reacted N₂O with 1 (Scheme 2). The purple solution of 1 in toluene turned green upon exposure to N2O for half an hour at room temperature. Cooling of the solution under a N2O atmosphere leads to a cluster compound 3. The X-ray structural analysis of 3 shows a chair like conformation for the central [CrO]₄ fragment, in which two of the oxygens show (μ_2 -O) and the other two show (μ_3 -O) bridging coordination modes (Fig. 2). The Cr-O bond distances are as expected, longer for the $(\mu_2 - O)$ [1.839–2.050 Å] coordinated oxo-ligand than the (μ_3 -O) [1.706–1.965 Å] oxo-ligated moiety. Similarly, the Cr1-Cr1A bond length [2.7904(10) Å] for the Cr-Cr fragment joined by two (μ_3 -O) is shorter than the two terminal Cr–Cr fragments [2.8375(8) Å each] joined by one (μ_2 -O) and one (μ_3 -O) ligand. The aminopyridinato ligands are no more coordinated in a µ2-fashion but are coordinated in an η^2 -mode. The N_{amido}-Cr bond lengths for the aminopyridinato ligands coordinated to the two central Cr atoms are comprehensively shorter [1.898(3) Å] than Namido-Cr bond lengths [2.081(1) Å] for the aminopyridinato ligands coordinated to the two terminal Cr atoms. On the other hand, N_{pyridine}-Cr bond lengths for the two central Cr atoms are longer [2.179(3) Å] than N_{pvridine}-Cr bond lengths [2.113(3) Å] for the two terminal Cr atoms, highlighting the flexibility of the aminopyridinato ligands.¹³



Scheme 2 Synthesis of complex 3.



Fig. 2 Molecular structure of **3** with hydrogen atoms omitted for clarity. Select bond lengths [Å] and angles [°]. N1–Cr1 1.898(3), N2–Cr1 2.179(3), N3–Cr2 2.081(3), N4–Cr2 2.113(3), O3–Cr1 1.702(2), O3–Cr2 1.950(2), O4–Cr1 1.840(2), O4–Cr1A 1.949(2), O4–Cr2 2.043(2), Cr1–Cr1A 2.7904(10), Cr1–Cr2 2.8375(8); Cr1–O3–Cr2 101.74(10), Cr1–O4–Cr1A 94.80(10), Cr1–O4A–Cr2A 131.82(12), Cr1–O4–Cr2 90.55(9), O3–Cr1–O4A 122.43(10), O3–Cr1–N1 119.07(11), O4–Cr1–N1 118.37(11), O3–Cr1–O4A 87.55(10), O4–Cr1–O4A 85.20(10), N1–Cr1–O4A 101.29(10), O3–Cr1–N2 96.07(10), O4–Cr1–N2 103.41(10), N1–Cr1–N2 65.75(10), O4–Cr1–N2 166.68(10), O3–Cr1–Cr1A 108.90(8), O4–Cr1–Cr1A 44.11(7), N1–Cr1–Cr1A 116.75(8), N2–Cr1–Cr1A 146.37(8), O3–Cr1–Cr2 42.29(7), O4–Cr1A–Cr2A 115.01(7), N1–Cr1–Cr2 112.13(8), N2–Cr1–Cr2 134.05(7), Cr1–Cr1A–Cr2A 78.12(2), O3–Cr2–O4 78.68(9), O3–Cr2–N3 162.07(10), O4–Cr2–N3 114.10(10), O3–Cr2–Cr1 151.68(8), N4–Cr2–Cr1 137.74(8).

The room temperature magnetic moment of 3 is 6.16, too low for a Cr^{III} compound with four S = 3/2 centres (theoretical value: 7.74, assuming g = 2). The measured values are closer to the theoretically expected values for four S = 1 centres ($\mu_S = 5.65 \ \mu_B [g = 2]$) assuming some orbital momentum contribution. This is in agreement with a Cr–Cr-bond with a low bond order. Upon cooling, a slow decrease of the magnetic moment down to 3.84 μ_B at 25 K is observed. This behaviour is best explained with weak antiferromagnetic interactions between the remaining unpaired electrons of the four chromium centres. Indeed, the magnetic moment at 25 K is in the region expected for one unpaired electron per chromium ($\mu_S = 3.46 \ \mu_B$ assuming g = 2).



Scheme 3 Synthesis of 4.

The pronounced drop in the magnetic moment below 25 K can be explained using zero field splitting.

The introduction of SO₂ into a toluene solution of **1** led to an immediate colour change from purple to orange (Scheme 3). Orange crystals of 4 were grown at low temperature. X-ray analysis revealed the formation of a dithionite $(S_2O_4^{2-})$ ligand. In compound 4 the dithionite ligand lies on top of the Cr₂ moiety with a bridging O₂S-SO₂²⁻ group where the four oxygen atoms coordinate to the two Cr atoms with Cr-O bond distances between 2.0360(14) and 2.0698(14) Å (Fig. 3). This coordination mode of dithionite to transition metals has not been described yet (to the best of our knowledge). The long S-S bond distance of 2.3786(7) Å could be due to the π antibonding interactions and is comparable to that found in Na₂S₂O₄ [2.389 Å].¹⁴ A Cr–Cr bond distance of 2.3419(5) Å was observed. The S-O bond distances are in the range of 1.5045(15)-1.5189(15) Å. The O-S-O angles of 107.46(8) and 107.85(8) $^{\circ}$ are close to tetrahedral but the S-S-O angles (92.21(6)-93.90(6)°) deviate strongly. Compound 4 is a thiolate analogue of an oxalate, which has been previously observed due to the reductive coupling of CO₂ for Fe^I and Ni^I compounds.¹⁵ The ¹H NMR of 4 shows very broad peaks in the diamagnetic range of 0 to 8 ppm and could



Fig. 3 Molecular structure of **4** with hydrogen atoms omitted for clarity. Select bond lengths [Å] and angles [°]. N1–Cr1 2.0551(16), N2–Cr2 2.0594(16), O1–S1 1.5045(15), O1–Cr1 2.0360(14), O2–S1 1.5189(15), O2–Cr2 2.0698(14), S1–S2 2.3786(7), Cr1–Cr2 2.3419(5); S1–O1–Cr1 112.80(8), S1–O2–Cr2 114.73(8), S2–O3–Cr2 112.26(8), S2–O4–Cr1 115.36(8), O1–S1–O2 107.85(8), O1–S1–S2 92.21(6), O1–Cr1–N4 172.07(6), O1–Cr1–N1 89.23(6), N4–Cr1–N1 98.19(6), O1–Cr1–O4 76.65(6), N4–Cr1–O4 96.11(6), N1–Cr1–O4 165.40(6), O1–Cr1–Cr2 95.14(4), N1–Cr1–Cr2 90.47(5), O4–Cr1–Cr2 87.18(4), O3–Cr2–N2 172.16(6), O3–Cr2–N3 87.64(6), N2–Cr2–N3 99.76(6), N2–Cr2–Cr1 95.27(4), N2–Cr2–Cr1 87.39(5), O2–Cr2–Cr1 87.38(4).

be attributed to the steric hindrance of the molecule. Low temperature experiments could not be carried out since the compound precipitates quickly in non-polar solvents like toluene and decomposes in more polar solvents like THF. Compound **4** is stable like **2** without an atmosphere of SO₂ or CO₂, respectively, and does not convert back to **1** as evidenced by NMR studies.

Density functional theory (DFT) calculations were performed using the *TURBOMOLE*¹⁶ programme package. The geometry of 1, 2 and 4 was optimized starting from structural data obtained by X-ray single crystal structure analysis. Computational details are given in the ESI.† Calculations indicate the reduction of the formal bond order from five to four for the SO₂ activation product (4). Fig. S2 (ESI[†]) shows the metal centered HOMOs of 1 and 4. The formal bond order of the carbonyl complex 2 is lower than four. In addition to the consumption of the two δ-bonds, one of the π bonds is also significantly involved in CO bonding. The relevant frontier orbitals of 2 are shown in Fig. S3 (ESI[†]). The geometry optimization of 2 shows that the CO ligands form stronger bonds with one single Cr center (Table S4, ESI[†]). The energy of this non-centrosymmetric structure is 5 kcal mol⁻¹ lower than the energy of the centrosymmetric structure. Due to the symmetry of the molecule, the crystal structure of 2 shows two indistinguishable Cr centers.

In summary, we could show that quintuply bonded dichromium complexes can activate CO_2 and SO_2 in a controlled fashion. Oxygen removal is observed during CO_2 activation leading to a doubly CO bridged chromium complex. As an oxygen scavenger, an oxo-bridging tetrametallic Cr-complex could be identified. SO_2 activation proceeds *via* S–S-bond formation leading to a unique dithionite complex.

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