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Neutral cyclic alkyl amino carbenes stabilize rare chromium(I) centres

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Cr(I)Cl is a very unstable species. The present work describes the stabilization of Cr(I)Cl in the low-coordinate environment of cyclic alkyl(amino) carbone ligands and its synthetic application to yield an unprecedented cationic complex with a two-coordinate Cr(I). One electron reduction of (cAAC)2CrCl2 (1) with equivalent amounts of KC8 results in the formation of (cAAC)2CrCl (2), with a distorted trigonal planar configuration at the metal. SQUID, EPR and theoretical studies reveal a Cr(I) centre with S = 3/2 spin ground state for 2. It represents the first example of a mononuclear Cr complex showing slow magnetic relaxation in an applied magnetic field. The chloride atom in 2 is expected to be prone to further reactions with appropriate reagents. This qualifies 2 as a promising precursor for the preparation of various interesting complexes with Cr(I) in low-coordinate environment. The first example of this metathesis reaction is observed when 2 is treated with Na[B(C8H8(CF3)2)2] resulting in [(cAAC)2Cr][B(C8H8(CF3)2)2]1, a linear cationic complex with two-coordinate Cr(I) and an S = 3/2 spin ground state.

Introduction

Chromium is an important first-row transition element because of the broad utility of its compounds in catalysis and numerous other applications.1 In addition to this, the magnetic properties of chromium (Cr) are unique in the sense that it is the only element which shows antiferromagnetic ordering at temperatures below 38°C while above this temperature it exhibits paramagnetism.2 A recent study revealed that elemental Cr exhibits special electrical properties similar to a magnet which would enable the application of antiferromagnets in spintronics.3 High-coordinate, especially octahedral complexes dominate the coordination chemistry of Cr, but recent years have witnessed an interest in complexes with low-coordinate chromium as well.4 One of the main strategies to obtain complexes with low-valent low-coordinate chromium is to use sterically demanding ligands, which restrict metal atoms from achieving high-coordination polyhedra. Power and coworkers used a bulky monoanionic terphenyl ligand to synthesize ArCrCrAr (Ar = C6H5-2,6-(C6H5-2,6-η2-P2))3, the first stable binuclear compound with five-fold bonding between two Cr atoms.5,6 However, a number of compounds with quintuply bonded Cr(I) centres have been synthesized since then using various ligands.5b-8 In a different strategy, other binuclear Cr(I) complexes of composition [(η2-P2)C6H5-NacnacCr]2(μ-X) (X = N3, C6H6; nacnac = CH(CMeNAr)2, Ar = C6H5-2,6-η2-P2) were synthesized by Theopold and coworkers using a monoanionic chelating β-diketiminate ligand.6 Each Cr atom in these compounds is chelated by a terminal bidentate β-diketiminate and the two metal ions are bridged by a central N3 or benzene moiety. However mononuclear β-diketiminate Cr(I) complexes were obtained by coordinating various acetylene derivatives to the Cr centre.7 Utilizing an even more crowded terphenyl ligand Ar′ (Ar′ = C6H5-2,6-(C6H5-2,4,6-η3-P3)) led to the formation of mononuclear compounds with two-coordinate Cr(I) of compositions Ar′Cr(TTHF) and Ar′Cr(PPh3).8

In all of the above compounds the Cr(I) atoms are bound to a monoanionic and relatively bulky ligand. This consequently leaves the Cr(I) atom without a functional group, which could be further transformed to other important compounds by appropriate metathesis reactions. In this respect our aim was to stabilize Cr(I)Cl species which, unlike Cr(II) or Cr(III) halides, do not exist under normal conditions.3 Various N-heterocyclic carbenes (NHC) have been reported to form adducts with Cr(II)Cl2 and Cr(III)Cl3, but they are not known to form stable adducts with Cr(I)Cl.10 At this juncture, it is worth noting that NHC has carbene carbon atom bonded to two N atoms which are σ-withdrawing and π-donating where as in cyclic alkyl(amino)carbenes (caacs) one of the nitrogen atoms is replaced by a σ-donating quaternary carbon atom. This allows caac to exhibit more nucleophilic and electrophilic character than NHC and hence utilizing caac will be advantageous to stabilize unstable species with low-valent metal centres.11 In this direction, we have prepared (cAAC)2CrCl, a compound in which Cr(I)Cl is stabilized by two flanking caacs. This compound features the first stable Cr(I)Cl entity having Cr(I) in a three-coordinate non-chelating ligand environment. Such a compound is anticipated to offer enormous scope for the preparation of various other Cr(I) derivatives by replacing the chloride atom with suitable functional groups of interest. Consequently an attempt to
substitute the chlorine atom with an anionic group of extremely high steric hindrance may result in the stabilization of a Cr⁷⁺ species in the coordination environment of two neutral cAACs. A recent report showed an anionic complex with Cr(I) in the two-coordinate environment of monoanionic amido ligand, -N(SiPr₃)Dipp (Dipp = C₈H₈-2,6-3-iPr₂). However, Cr(I) cationic complexes so far known are stabilized only in high-coordinate environment. The best known of this kind are π-arene complexes of Cr(I) as well as chromium carbonyl complexes with phosphorus donor ligands. These Cr(I) cationic complexes have been found to be effective in the catalytic ethylene tetramerization and trimerization reactions. However, there is no report of a cationic compound with a two-coordinate Cr(I) ion and so we became interested in synthesizing such a novel compound. Reaction of (cAAC)₂CrCl with Na[B(C₆H₆(CF₂)₃)] resulted in the formation of [(cAAC)₂Cr][B(C₆H₆(CF₂)₃)], a cationic two-coordinate Cr(I) complex. It should also be noted that low-coordinate transition metal complexes with high symmetry and low metal oxidation state attract a lot of interest because of their potentially interesting magnetic properties, including large magnetic relaxation barriers. This is because the remaining near-degeneracy of d-orbitals in such low-coordinate systems, in combination with a proper number of d-electrons, can give rise to significant unquenched orbital angular momentum. Hence the magnetic properties of some of the new Cr complexes have been investigated.

Results and discussion

Reaction of two equivalents of cAAC with CrCl₂ in THF results in the formation of (cAAC)₂CrCl₂ (1), a pink compound in high yield (Scheme 1). The high exothermicity (-102.1 kcal/mol) for the formation of 1 from CrCl₂ and cAAC supports the experiment. The molecular structure of 1 shows Cr-C (2.180 Å) and Cr-Cl (2.339 Å) bond distances comparable to the corresponding bond lengths in (NHC)₂CrCl₂ reported earlier. (see SI for structural details of 1) The reaction of equivalent amounts of 1 and KC₁ in THF afforded (cAAC)₂CrCl (2) as a green product. The abstraction of Cl from 1 by K is calculated to be exothermic by -13.1 kcal/mol. 2 crystallizes in the triclinic space group P-1. The Cr atom in 2 features a distorted trigonal planar coordination geometry. The molecular structure of 2 is shown in Figure 1. The distance of the chromium atom from the Cl-C1-C1’ plane is 0.018 Å, which is very close to the ideal planar geometry around the Cr atom. The Cr-C bond lengths in 2 are 2.084(2) and 2.093(2) Å which are shorter than the corresponding distances in 1. However, the Cr-Cl bond length is 2.366(1) Å which is close to the corresponding value for 1. Unlike in the case of 1, the cAAC ligands in 2 are arranged trans with respect to the position of the N atoms.

Scheme 1: Synthesis of compounds 1-3. Gas phase reaction energies, ∆E in kcal/mol are calculated at the M06/def2-TZVPP//BP86/def2-SVP level of theory. ∆E₂ is the reaction energy of 1 with K, considering KC₁ as a source of later.

Figure 1. Molecular structure of 2. Hydrogen atoms are omitted for clarity. Anisotropic displacement parameters are depicted at the 50% probability level. Selected bond lengths [Å] and angles [°]. Calculated values at the BP86/def2-SVP level of theory are given in square brackets. Cr–Cl, 2.366(1) [2.332]; Cr–C1’, 2.084(2) [2.091]; Cr–Cl, 110.11(5) [106.3]; C1–N1, 1.333(2) [1.350]; C1’–N1’, 1.333(2) [1.350]; C1–Cr–Cl, 112.70(5) [106.3]; C1’–Cr–Cl, 110.11(5) [106.3]; C1–Cr–C1’, 137.17(6) [147.3].

Figure 2. χ₀T versus T plot for 2. Inset: VTVH magnetization measurements as M_{mol} versus μ₀B/kT. Solid lines represent the calculated curve fits (see text)
A magnetic susceptibility measurement of 2 (Figure 2) shows a $\chi_M T$ value of 4.81 cm$^3$mol$^{-1}$K at 210 K, which is slightly higher than the expected spin-only value for a $S = 3/2$ system (4.375 cm$^3$mol$^{-1}$K). $\chi_M T$ remains nearly constant down to 20 K but drops to 4.00 cm$^3$mol$^{-1}$K at 2 K, which is likely due to zero field splitting (vide infra). No temperature dependence of $\chi_M T$ was observed between 20 K and 210 K, indicating that only the $S = 3/2$ spin state is populated in this temperature range. The experimental $\chi_M T$ vs. $T$ as well as the variable temperature-variable field (VTTH) magnetization data were simultaneously modelled using the anisotropic spin Hamiltonian with Zeeman splitting as well as axial ($D$) and rhombic ($E$) zero-field splitting as given in eq. 1.

$$\hat{H} = g \mathbf{B} \cdot \mathbf{S} + D \left( S_z^2 - \frac{S^2}{3} \right) + E \left( S_x^2 - S_y^2 \right) \quad \cdots (1)$$

In this Hamiltonian, $\mathbf{H}$ is the magnetic field vector, $\mathbf{g}$ is the Zeeman tensor, and the other terms pertain their usual meaning. The best fit values are $g_x = g_y = 1.93$, $g_z = 2.41$ and $D = 0.5$ cm$^{-1}$. The rhombic ZFS parameter $E$ was fixed to zero to avoid any overparametrisation. It was also possible to simulate the experimental data using a negative $D$ with $g_x = g_y > g_z$. Thus, the SQUID measurements were inconclusive with respect to a proper assignment of the sign of $D$ in this case, likely due to small magnitude of this parameter (~1 cm$^{-1}$).

Figure 3. Temperature dependence of $\chi''$ for 2 at various frequencies with an applied dc field of $H_{dc} = 500$ Oe.

Alternating current (ac) magnetic susceptibility measurements at various frequencies were performed, both in the absence of a direct current (dc) magnetic field as well as with applied dc fields ($H_{dc} = 250 - 3000$ Oe). Application of dc fields revealed a frequency dependence in the imaginary part of the magnetic susceptibility ($\chi''$) (Figure 3, Figures S6), thus indicating slow magnetic relaxation in 2 at low temperatures. Although 3d transition metal based single ion magnets (SIMs) have been known since 2010, so far, only SIMs containing Fe(I,II,III), Co(II), Ni(I) or Mn(III) ions have been reported. Significant efforts have been made to understand the magnetic behavior of homometallic and heterometallic clusters of chromium in recent past. To the best of our knowledge, this is the first report on slow relaxation of magnetization for a mononuclear Cr complex suggesting SIM behavior.
The valence electron (VE) count of Cr(II) in 1, Cr(I) in 2 and Cr(I) in 3 are 12, 11 and 9, respectively. Apart from the Cr-ligand bond formation in these high spin complexes, four, five and five valence electrons occupy Cr-based d-orbitals in 1, 2 and 3 respectively (Figure S9, S10, S11). These singly occupied d-orbitals are responsible for Cr→π^∗_cAAC back donation. Hence, the extent of Cr→π^∗_cAAC back donation can be understood from NBO spin density. The calculated NBO spin density on the Cr atom in complexes 1, 2 and 3 are 3.85, 4.25 and 4.65 respectively (Table S9). The relatively less spin density on the Cr-atom in 2 as compared to that of 3 can be attributed to relatively higher Cr→π^∗_cAAC back donation in 2, which is also shown in SOMO+3 and SOMO+4 (Figure S10). However, only one MO shows Cr→π^∗_cAAC back donation in 3 (SOMO+2 in Figure S11) and in 1 (SOMO in Figure S9). The lowest positive group charge of cAAC in 2 (0.01 e) as compared to 1 (0.23 e) and 3 (0.23 e) also corroborates well with the MO analysis.

The dissociation energy of one Cr–C_cAAC bond in 1, 2 and 3 are 51.1, 48.6 and 79.7 kcal/mol, respectively. The highest Cr–C_cAAC bond strength in 3 can be qualitatively attributed to the higher charge of the complex and the weaker Cr–C_cAAC bond strength in 2 as compared to 1 can be attributed to the lower oxidation state of Cr in 2. Moreover, the SOMOs showing the σ-antibonding interaction between Cr and Cl⁻ in 2 indicate easiness of the removal of Cl⁻ ligand for the formation of complex 3. The dissociation energies of a Au–C_cAAC bond in Au(cAAC)_2, Cu–C_cAAC bond in Cu(cAAC)_2 and Co–C_cAAC in Co(cAAC)_2 are 45.6, 48.7 and 64.3 kcal/mol, respectively. These molecules show significant amount of π-back donation, a situation very different from the present Cr complexes. Note that the NBO data such as C–N bond order, partial charges and spin density give only the relative strength of donor-acceptor interactions in these complexes. It does not necessarily correlate with the dissociation energy of the Cr–C_cAAC bond, which depends on the other energy components as well. Thus, 1, 2 and 3 are examples of complexes with low-coordinate and low-valent Cr in their highest spin states mainly stabilized by the σ-donation of the cAAC ligands.

In order to understand the extent of Cr→π^∗_cAAC back donation as a function of spin states, we have calculated the C–N bond order and cAAC group charges of complex 3 in the sextet, quartet and doublet spin states (Table S8). The C–N bond order in the sextet state (1.50) is the same as that of free cAAC, whereas the corresponding bond orders in quartet and doublet states are 1.34 and 1.38, respectively. If the C–N bond order can be considered as the measure of Cr→π^∗_cAAC back donation, the relative back donation is in the order: quartet > doublet > sextet. This is in-line with the calculated cAAC group charges viz., 0.23 for sextet, 0.05 for quartet and 0.12 for the doublet. Hence, rather than the Cr→π^∗_cAAC back donation, the sextet spin state plays a more significant role for the stability of the low coordinate

Quantum mechanical calculations at the M06/def2-TZVPP/BP86/def2-SVP level of theory show that the quintet state of the complex 1 is lower in energy than the triplet and singlet electronic states by 59.2 and 63.6 kcal/mol, respectively. The sextet state of 2 is more stable by 5.8 and 38.5 kcal/mol whereas the sextet state of 3 is more stable by 33.3 and 66.6 kcal/mol as compared to their quartet and doublet states respectively. The calculated geometrical parameters of 1, 2 and 3 in their respective high-spin states are also closest to those of the crystal structures (Figures S12-S14, Tables S3-S5). The C–N bond lengths and Wiberg bond indices in the free cAAC ligand (1.320 Å, 1.50), complexes 1 (1.330 Å, 1.47), 2 (1.350 Å, 1.37) and 3 (1.324 Å, 1.50) suggest that the Cr→π^∗_cAAC back donation is less in these complexes (Table S6-S8), while in 2 a relatively higher back donation is observed. The geometrical analysis is in line with the following bonding description based on molecular orbital and NBO analyses.

The molecular structure of cAAC is shown in Figure 5. Hydrogen atoms are omitted for clarity. Anisotropic displacement parameters are depicted at the 50% probability level. Primes (') represent the atoms of the second molecule present in the asymmetric unit. Selected bond lengths [Å] and angles [']. Calculated values at the BP86/def2-SVP level of theory are given in square brackets. Cr–C1, 2.134(2) [2.152]; Cr’–C1’, 2.136(2) [2.152]; C1–N1, 1.303(2) [1.324]; C1–N1’, 1.305(2) [1.324]; C1–Cr–C1A, 180.0 [180.0].

The above discussion indicates that in 2, the Cr(I)Cl species which would otherwise be very unstable is stabilized by the donor electrons of cAAC. This is further supported by the complementary theoretical calculations (see below). Replacement of chlorine atom in 2 with other functional groups is an open challenge which may lead to a bouquet of new complexes with low coordinate Cr(I) with potential applications. In the first reactivity study of this series, we have found that the reaction of 2 with equivalent amounts of Na[B(C_6H_5)(CF_3)A] results in the elimination of NaCl and thereby the formation of the ionic compound [(cAAC)Cr][B(C_6H_5)(CF_3)A] (3) as a pale green solid. The formation of 3 by the abstraction of Cl⁻ by Na⁺ is exothermic by -28.6 kcal/mol. Single crystals of 3 suitable for X-ray diffraction were obtained from a saturated solution in toluene at ~35 °C. 3 crystallizes in the triclinic space group P-1. The molecular structure of the cation in 3 is given in Figure 5 (see S1 for full molecular structure). The geometry around Cr is strictly linear and the Cr atom is located on a crystallographic inversion centre. However, there are two crystallographically different molecules with Cr–C bond distances of 2.134(2) and 2.136(2) Å which are a little longer than the corresponding bond distances of 2. The EPR spectrum of 3 was measured and simulated similarly to 2 and resulted in an excellent fit using the parameters: S = ½/2, g_anisotropic = 2.00, |D| = 0.37 cm⁻¹ and E/D = 0.06, as shown in Figure 4 (right).
Conclusion

This work presents two novel low-coordinate Cr(I) complexes 1 and 2. The X-ray crystal structure, magnetic, EPR and theoretical studies show a rare Cr(II) species is stabilized by the coordination of two cAAC ligands. The presence of a chlorine atom in the three-coordinate Cr(II) compound 2 is expected to make it a good precursor for the preparation of various Cr(III) compounds of potential interest. Attempts are in progress in our laboratory to replace the chlorine atom with other important elements such as fluorine and hydrogen. 2 represents the first example of a mononuclear Cr complex showing slow magnetic relaxation in an applied magnetic field. This work also shows the utility of 2 as a precursor for preparing 3, a hitherto elusive Cr(I) catonic compound in a two-coordinate neutral ligand environment. The magnetic parameters for 3 are surprisingly quite different from those of 2, while the total spin stays the same, S = 1/2, thus exhibiting the possible control over the magnetism that can be synthetically achieved in the compounds. However, potential applications of the coordinately unsaturated Cr(I) sites in 2 and 3 are still to be investigated.

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See SI for the details of computational methodology, related references, and MO analysis.