Abstract: A magnesium complex incorporating a novel metal–CO₂ binding motif is spectroscopically identified. Here we show with the help of infrared photodissociation spectroscopy that the complex exists solely in the [ClMgCO₂]⁻ form. This bidentate double oxygen metal–CO₂ coordination has previously not been observed in neutral nor in charged unimetallic complexes. The antisymmetric CO₂ stretching mode in [ClMg(η²-O₂C)₂]⁻ is found at 1128 cm⁻¹, which is considerably redshifted from the corresponding mode in bare CO₂ at 2349 cm⁻¹, suggesting that the CO₂ moiety has a considerable negative charge (~1.8 e⁻). We also employed electronic structure calculations and kinetic analysis to support the interpretation of the experimental results.

Although not a new concept, the activation of carbon dioxide by metal (M) complexes is currently receiving much interest with regard to improving the feasibility of using CO₂ as a carbon feedstock in chemical synthesis. This is an attractive prospect with regard to CO₂’s abundance, and as an addition of a CO₂ molecule to a carbohydrate substrate. RuBisCO which catalyzes C₅ reactions, magnesium is of special interest. For example, in the Grignard reaction which is well known in organic chemistry, it plays a fundamental role in the classic Grignard reaction which is well known in organic chemistry.

**Scheme 1.** The metal–CO₂ coordination modes M(η¹-CO₂) (A) and M(η²-CO₂) (B) are the most common. Coordination as in (C) has only been observed in bimetallic complexes.

A key piece of information in understanding the reactivity of metal–CO₂ complexes is the coordination of CO₂ to the metal. In neutral complexes containing a single metal atom monodentate coordination M(η¹-CO₂) or bridging coordination M(η²-CO₂) are most common, corresponding to structures (A) and (B) in Scheme 1, respectively. Linear “end-on” coordination is also seen, but is less common. Work on isolated anionic metal–CO₂ complexes is scarce, but [M(–CO₂)₃]⁻ complexes also prefer the coordination modes A and B, retaining a bent CO₂ moiety as a result of the partial negative charge transfer into the π* orbital of CO₂. In contrast, cationic metal–CO₂ complexes, [M(–CO₂)₃]⁺, have so far shown to exclusively coordinate CO₂ “end-on” to the metal due to the electron deficiency resulting in electrostatic bonding between the positively charged metal atom and the partial negative charge on oxygen in CO₂. Here, we present first spectroscopic evidence of an anionic unimetallic Mg-containing complex with a bidentate double oxygen M(η²-O₂C) coordination, a binding motif that has, so far, only been encountered in bimetallic complexes (structure (C) in Scheme 1).

Mass spectrometric evidence was recently presented suggesting the presence of the bidentate motif (C) in anionic Mg-containing complexes of the type [XMgCO₂]⁻ in which X = Cl, Br, OH. These complexes are formed by electro-spraying a solution of oxalic acid and a Mg²⁺ salt (MgX₂, X = Cl, Br). The mass spectra show that the X = OH complex is reactive in S₂,2 gas phase reactions, providing support for the bidentate [HOMg(η²-O₂C)]⁻ structure analogous to structure 1 in Figure 1. However, the G4-level calculations included in that paper suggest the presence of an additional isomer [HOMg(η¹-CO₂)]⁻ with the more classic structure (B), analogous to structure 2 in Figure 1, which is 32 kJmol⁻¹ higher in energy than 1. Based on this information it is not clear which of the isomers is observed in the experiment, as not only the energetics of the minimum energy structures but also the barrier heights along the reaction coordinate need to
be considered. We have now conducted infrared photodissociation (IRPD) spectroscopy on the X = Cl complex, [ClMgCO2]−, to pin down the preferred binding motif of these complexes based on their vibrational fingerprint.

Vibrational spectra of [ClMgCO2]−, measured by way of IR vibrational predissociation spectroscopy of the [ClMgCO2]−·D2 complex (see the Experimental Section for details), are presented in the top part of Figure 1. Two IRPD spectra are shown probing overlapping regions, in which the higher energy region is covered with a tabletop laser system (I, 1050–2000 cm−1) and the lower energy region with a tabletop laser system (trace II) satisfactorily covers the region from 450 to 1140 cm−1 (II). The two strongest IRPD bands at 1128 cm−1 (a) and 570 cm−1 (d) correspond to the antisymmetric CO2 stretching (vnAS) and Cl-Mg-C stretching (vnCS) modes, respectively. Peaks characteristic for 1 are also detected at 1106 cm−1 (b), 814 cm−1 (c), and 462 cm−1 (e) and assigned to the CO2 symmetric stretch (vnS), bend (vnb), and rock (vnR), respectively. For comparison, the most intense IR transitions of 2 are predicted at 1700 cm−1 (free C=O stretch), and at 950 cm−1 (bound C=O stretch), at which no corresponding features are present in the experimental spectra.

Table 1: Experimental IRPD band positions (in cm−1), computed unscaled harmonic frequencies, and band assignments for [ClMg(η2-O2C)]−.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>CCSD(T)</th>
<th>MP2</th>
<th>wB97XD</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1128 (a)</td>
<td>1139</td>
<td>1155</td>
<td>1216</td>
<td>CO2 antisym. stretch (vnAS)</td>
</tr>
<tr>
<td>1106 (b)</td>
<td>1105</td>
<td>1108</td>
<td>1171</td>
<td>CO2 sym. stretch (vnS)</td>
</tr>
<tr>
<td>814 (c)</td>
<td>816</td>
<td>806</td>
<td>849</td>
<td>CO2 band (vnR)</td>
</tr>
<tr>
<td>570 (d)</td>
<td>560</td>
<td>560</td>
<td>568</td>
<td>ClMgC stretch (vnCS)</td>
</tr>
<tr>
<td>463 (e)</td>
<td>459</td>
<td>452</td>
<td>475</td>
<td>CO2 rock (vnR)</td>
</tr>
<tr>
<td>MSE[b]</td>
<td>5</td>
<td>10</td>
<td>41</td>
<td>n/a</td>
</tr>
</tbody>
</table>

[a] aug-cc-pVTZ, untagged; [b] aug-cc-pVTZ, tag on oxygen; [c] mean signed error; n/a not applicable.

The position of the antisymmetric CO2 stretching mode vnAS yields additional information on the extent of charge transfer in the complex. In bare CO2, vnAS is found at 2349 cm−1. Addition of an electron to CO2 leads to a weakening of the C=O bonds, a distortion from the linear geometry, and a redshift of vnAS, which is proportional to the extent of charge transfer. While CO2− by itself is not stable in the gas phase with respect to electron autodetachment (EA = −0.6 eV), its IR spectrum has been measured in a Ne matrix, yielding a value for vnAS of 1658.3 cm−1.[13] Intermediate values for vnAS (1865–1680 cm−1) have been found in CO2-solvated M(CO2)− complexes (M = Au, Ag), thus corresponding to partial charge transfers ranging from 0.5 to 0.9 e−.[8,9] The observation of a considerably more redshifted vnAS for 1 (1128 cm−1) suggests that substantially more negative charge (−1.8 e−) is transferred to the CO2 moiety in [ClMgCO2]−. This analysis then yields a CO2− 2−ligand with strong carbene character.

We also performed calculations on neutral [Mg−CO2]− to see if the chloride is directing the CO2 coordination in [ClMgCO2]−. Both MP2/aug-cc-pVTZ and G4 find only [Mg(η2-O2C)] as a stable complex, with no minimum found for [Mg(η2-CO2)]−, suggesting that the chloride does not alter the CO2-coordinating properties of magnesium.

Energetically, 1 is favored by 29 KJ mol−1 over 2; however, the formation mechanism and reaction barriers also need to be considered. The potential energy surface, calculated at the G4 level, along with the kinetic analysis presented in Figure 2 provide further support for the exclusive presence of isomer 1. Both isomers 1 and 2 follow similar mechanisms beginning...
with the isomerization of the $[\text{ClMgC}_2\text{O}_4]^{-}$ precursor. Once isomerization is complete via either $\text{TS A}$ or $\text{TS B}$, CO$_2$ is lost producing isomer 1 or 2, respectively. Formation of $[\text{ClMg}(\eta^1\cdot\text{O}_2\text{C})]$ via $\text{TS A}$ is greatly favored due to its much lower isomerization barrier of 133 kJ mol$^{-1}$, which is 100 kJ mol$^{-1}$ lower than the corresponding barrier of $\text{TS B}$ leading to $[\text{ClMg}(\eta^2\cdot\text{CO}_2)]^{-}$. The effect of this difference is clearly reflected in the kinetic analysis, because the formation of $[\text{ClMg}(\eta^2\cdot\text{CO}_2)]^{-}$ is not competitive until energies (>335 kJ mol$^{-1}$) are reached that significantly exceed the dissociation threshold of both complexes.

In summary, we have shown that the $[\text{ClMg}(\eta^2\cdot\text{O}_2\text{C})]^-$ complex exhibits a bidentate coordination of CO$_2$, in which both oxygen atoms are bound to Mg, a result which is consistent with the high oxophilicity and hardness of the Mg$^{2+}$ ion.$^{[12]}$ This may elucidate some of the intrinsic interactions of the biologically important Mg$^{2+}$ ion with an activated CO$_2$ molecule, because the CO$_2$ moiety in our complex has a considerable negative charge giving its carbon a highly nucleophilic character as a result of this binding scheme.

**Experimental Section**

The IRPD experiments were performed using the Berlin 6 K ion-trap triple mass spectrometer$^{[3]}$ in combination with the widely tunable and intense IR radiation from the free electron laser FHI-FEL.$^{[4]}$ Gas-phase anions were produced using a nanospray ion-source from a solution of oxalic acid (1 mM) and magnesium chloride (2 mM) in a 1:1 water/methanol mixture. $[\text{ClMgCO}_2]^{-}$ anions were obtained from $[\text{ClMgC}_2\text{O}_3]$ through skimmer collision-induced dissociation. Parent ions were mass-selected and focused into a cryogenically cooled radio frequency ring-electrode ion-trap. To allow for continuous ion loading and thermalization, the trap was continuously filled with D$_2$ gas at a trap temperature of 14.5 K. The untagged $[\text{ClMgCO}_2]^{-}$ and messenger-tagged $[\text{ClMgCO}_2]^{-}$(D$_2$) ions were then extracted from the ion trap and irradiated with the IR laser pulse prior mass spectrometric detection. The FHI-FEL was operated from 450–1200 cm$^{-1}$ with a bandwidth of approximately 4 cm$^{-1}$ and an average power of 6 mJ per macropulse. The IRPD spectra were recorded by monitoring all ion intensities simultaneously as the laser wavelength was scanned. The photodissociation cross-section $\sigma$ was determined as described previously.$^{[5]}$ Additional measurements in the range from 1000 to 2000 cm$^{-1}$ were performed using the radiation from an IR OPO/OPA table-top laser system.$^{[6]}$ The laser pulses had a bandwidth of ~3 cm$^{-1}$ and an average pulse energy of 0.7 mJ. The resulting IRPD spectra reflect the total parent ion signal (instead of $\sigma$), to avoid additional noise due to the normalization procedure at low laser pulse energies.

A range of density functional theory (DFT) functionals and the wavefunction methods MP2 and CCSD(T) were employed to simulate the experimental IRPD spectra. The DFT and MP2 calculations were performed with the Gaussian 09 program package, whereas the CCSD(T) calculations were performed with the MOLPRO and CFOUR program packages.$^{[7]}$ Briefly, the most accurate harmonic frequencies were obtained at the CCSD(T)/aug-cc-pVTZ level of theory, and the best relative intensities were obtained at the DFT levels of theory. The kinetic analysis was performed using the Rice–Ramsperger–Kassel–Markus (RRKM) theory using MP2/aug-cc-pVTZ rovibrational data and G4 energies.$^{[14,15]}$

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