Calix[4]pyrrolato gallate: square planar–coordinated gallium(III) and its metal–ligand cooperative reactivity with CO₂ and alcohols†

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Forcing a priori tetracoordinate atoms into planar configuration represents a promising concept for enhanced reactivity of p-block element-based systems. Herein, the synthesis, characterization, and reactivity of calix[4]pyrrolato gallates, constituting square planar–coordinated gallium(III) atoms, are reported. Unusual structural constraint-induced Lewis acidity against neutral and anionic donors is disclosed by experiment and rationalized by computations. An energetically balanced dearomatization/rearomatization of a pyrrole unit enables fully reversible metal–ligand cooperative capture of CO₂. While alcohols are found unreactive against the gallates, a rapid and selective OH-bond activation can be triggered upon protonation of the ligand. Secondary ligand–sphere modification adds a new avenue to structurally-constrained complexes that unites functional group tolerance with unconventional reactivity.

Results and discussion

Reacting the tetrasodium salts of two differently substituted calix[4]pyrrolato ligands with the gallium(III) source tetrakis[tris(phenyl)[1]phosphonium]tetrachlorogallate ([PPh₃][GaCl₄]) in 1,2-dimethoxyethane (DME) at room temperature over 3 h resulted in a colorless precipitate (Fig. 2A). After extraction with dichloromethane, pale rose solids were isolated in fair yields (69 and 57%). The ¹H NMR spectra of the obtained compounds showed a singlet at around 5.7 ppm and two sets of aliphatic resonances originating from the methyl and ethyl groups, respectively. These characteristics indicate effective solution-phase ⁷d molecular symmetry, consistent with a square planar environment around the gallium atom. Single crystals obtained from dichloromethane/n-pentane mixtures at −40 °C suitable for SCXRD analyses confirmed the NMR spectroscopical data and unveiled the square planar arrangement of the four N-donor sites enclosing the gallium(III) atom (Fig. 2B). The mean Ga–N bond lengths are 193.8(1) and 195.0(1) pm (the data for [³¹⁷¹Ga] is always given first), which are larger compared to the analogous

Introduction

Steric and electronic effects dictate the tetrahedral arrangement of substituents around tetrasubstituted p-block elements with a complete valence electron octet. While these so-called van’t Hoff–Le-Bel configurations are found in most chemical compounds, precise ligand design allowed unique exceptions to this paradigm. Most recent additions are the square planar anti-van’t Hoff–Le-Bel cases for aluminum and silicon by using the macrocyclic calix[4]pyrrolato ligand (Fig. 1A). While a close-to-planar neutral tetracoordinated gallium atom was recently reported in a pyridine diimine-supported complex, anionic and truly square planar–coordinated gallium(III) atoms and the reactivity of this compound class, remains elusive.

Planarization results in a distinct alteration of the electronic structure, namely rising the energy of the highest occupied molecular orbital (HOMO) and lowering the energy of the lowest unoccupied molecular orbital (LUMO) (Fig. 1B). These features equip the compounds with biphilic properties, ideal for element-ligand cooperative reactivity toward various functional groups (Fig. 1A). The potential of gallium–ligand cooperativity (GaLC) was so far exploited with binuclear Ga(II)–element-ligand cooperative reactivity features equip the compounds with biphilic properties, ideal for this paradigm.

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aluminum compounds (189.3(2) and 188.3(1) pm). The average cis- and trans-N-Ga-N angles are 90.0(1) and 90.0(1)°, and 178.6(1) and 178.7(1)°, respectively. The τ value of 0.02 for both gallates indicates essentially perfect square planarity. Hence, the calix[4]pyrrolato gallates represent – compared with the neutral gallium pyridine diimine systems (τ = 0.13–0.22) (ref. 18) – the first examples for gallium(m) with ideal square planar, anti-van’t-Hoff–Le-Bel configuration.

This structural arrangement was analyzed by various computational techniques (for all computational details see the ESI). Structural optimization with density functional theory (DFT) confirmed the square planar structure as the energetic global minimum. Constrained structural optimization of the neutral gallium pyridine diimine systems gave a conformer 581 kJ mol⁻¹ higher in energy as the fully relaxed structure (see Fig. S8 in the ESI). In comparison, constraining the non-cyclic tetrapyrrylato gallate, which has an effectively tetrahedral ground state, to square planarity costs 166 kJ mol⁻¹. This nicely illustrates the substantial support of the macrocyclic calix[4]pyrrolato ligand and its electronic properties (σ-accepting, π-donating) as being crucial for the experimental realization of this configuration.

The anion [MeI⁻] features an energetically low-lying LUMO (1.48 eV), having its largest coefficient (0.754 within a natural atomic orbital basis) for the 4pₓ orbital of the gallium atom (Fig. 2C). In contrast, the HOMO is entirely located in the ligand backbone (Fig. 2C). This agrees with the findings for the corresponding aluminum(m) and silicon(n) compound and matches early theoretical work from the von Schleyer group. This electronic structure should impart the isolated gallates with biphilic character (vide infra). To probe the global Lewis acidity of [MeI⁻], its solution phase (dichloromethane) fluoride ion affinity (FIAₐₙ) was computed. It amounts to 149 kJ mol⁻¹ ([F]⁻ : 132 kJ mol⁻¹). This value is slightly reduced compared to the analogous aluminates (196 kJ mol⁻¹), though still substantial given the anionic charge state of [MeI⁻]. For comparison, the FIAₐₙ value of the tetrahedral tetrapyrrylato gallate is 58 kJ mol⁻¹ – corresponding to an increase in Ga-F bond strength of 90 kJ mol⁻¹ mainly based on geometric effects.

Following these computational results [PPh₄]([MeI]) was reacted with one equivalent of the fluoride ion donor tetrabutyrammonium difluorotriphenylsilicate to tetrabutylammonium difluorotriphenylsilicate (TBAT) in acetonitrile at room temperature (Fig. 3A). Workup after 4 h afforded an orange solid in 80% yield. Its ¹H NMR spectrum showed two doublets for the pyrrolic β-protons and four signals for the methyl groups. Intriguingly, the signal of the endo-methyl groups next to the fluoride is split into a doublet (3.1 Hz), indicating through-space fluoride-proton spin–spin coupling. This doublet splitting was also found for the signal of the carbon atom of the respective methyl groups (18.3 Hz). The ¹⁹F NMR spectrum contained a resonance at −146 ppm, significantly broadened due to this coupling. In total, this is consistent with the formation of the dianionic [MeI-F]⁻, which was further confirmed by mass spectrometry. These results also agree with the calculated lower FIAₐₙ of difluorotriphenylsilane (85 kJ mol⁻¹) in comparison to the value of [MeI⁻] (see above) and with the overall negative solution-phase Gibbs free reaction energy of −65 kJ mol⁻¹ for the fluoride transfer reaction from difluorotriphenylsilicate to [MeI⁻] in acetonitrile. Additional experimental proof for the Lewis acidic reactivity of [MeI⁻] was found when [PPh₄][MeI] was crystallized from tetrahydrofuran (THF) as solvent.
One THF coordinating to the gallium atom was observed in the solid-state structure with a Ga–O distance of 202.3 pm (Fig. 2B). Interestingly, the computed solution-phase affinity toward THF is low for both gallates ([Me1][Ga3]1: 0 kJ mol$^{-1}$, [Et1]$^{-1}$: $-4$ kJ mol$^{-1}$), allowing for THF removal during the workup of the synthesis of [PPh4][Me1] and [PPh4][Et1], respectively.

Having established the Lewis acidity of the gallates by computations and experiment, it was tested whether their properties arising from the square planar-coordinated Ga(n) atoms serve to facilitate reactions with other substrate molecules. The analogous calix[4]pyrrolato aluminate showed manifold reactivity with carbonyl-containing compounds (see Fig. 1A)$^{23}$ so we reacted [PPh4][Me1] and [PPh4][Et1] with carbon dioxide.

After exposing both gallates dissolved in dichloromethane-$d_2$ to 5 bar of CO$_2$ at room temperature, the solutions turned intensively yellow within several hours. This color change indicates the dearomatization of one pyrrole moiety (marked with an asterisk in the following formulae). The $^1$H NMR spectra showed the clean emergence of a new $C_1$ symmetric species, which was unambiguously identified as the gallium–ligand cooperative addition product of CO$_2$ to [Me1]$^{-1}$ and [Et1]$^{-1}$, respectively (Fig. 4, see the ESI for details).$^1$ Interestingly, the conversion to [Me1$^\star$-CO$_2$]$^{-1}$ was found almost quantitative for the methyl case (96%), whereas a 2 : 1 ratio of starting material to addition product was observed for the ethyl gallate under the chosen reaction conditions. The Gibbs free activation energy for the addition reaction of CO$_2$ to [Me1]$^{-1}$ was calculated to $\Delta G_r = 90$ kJ mol$^{-1}$, and the overall Gibbs free reaction energy to $\Delta G_r = 6$ kJ mol$^{-1}$, which conforms with the experimental observations. Further, the computational simulations reproduced the decreased binding strength of CO$_2$ to [Et1]$^{-1}$ ( $\Delta G_r = 18$ kJ mol$^{-1}$). This may be rationalized by the slightly decreased Lewis acidity on the FIA$_{solv}$ scale of the gallium atom in [Et1]$^{-1}$ (see above) as well as by a larger stabilization of [Me1$^\star$-CO$_2$]$^{-1}$ in a dichloromethane solution as found by COSMO-RS$^{41-43}$ calculations (see Chapter S11 in the ESI).$^1$
In agreement with these thermoneutral computed values, the application of reduced pressure to solutions of \([\text{PPh}_4]^{[\text{Me1}]}\)–CO\(_2\) over several hours at room temperature resulted in the quantitative removal of CO\(_2\) from the systems and in the selective reformation of \([\text{H}^{-}\text{Me1}]]\) (see Fig. S1 and S2 in the ESI).\(^\dagger\)

Element-ligand cooperative (ELC) binding of CO\(_2\) is a widely studied reaction with various transition metals\(^{44-50}\) and p-block elements\(^{51-59}\) and was also investigated in small molecule contexts other than ELC.\(^{35,60-66}\) To the best of our knowledge, the described reaction represents the first example for reversible CO\(_2\) fixation by a gallium(n) platform. This is achieved by the balanced energetics in dearomatization/rearomatization of one of the pyrrole rings of the calix[4]pyrrolato ligand.

We next turned our attention to protic substrates, namely alcohols, as their ready activation by the calix[4]pyrrolato aluminate was already noticed by our group.\(^{24}\) Upon reacting one equivalent of \(\text{PrOH}\) with \([\text{PPh}_4]^{[\text{Me1}]}\) and \([\text{PPh}_4]^{[\text{Et1}]}\), respectively, no reaction occurred, contrasting the behavior of the corresponding aluminates, i.e., the formation of an oxygen–element bond with the proton being transferred to the 2-position of one of the pyrrole rings (Fig. 5B, left part). We reasoned that protonation of the ligand backbone might serve as a powerful strategy for second-sphere reactivity modulation in structurally constrained complexes. Such tactics are well-known in an alternative context,\(^{67-70}\) for example, applied by the Corey group in enantioselective Diels–Alder catalysis.\(^{68-70}\) To probe this hypothesis computationally, the \(\alpha\)-pyrrole protonated, neutral \([\text{H}^{-}\text{Me1}^*]\) was optimized by DFT and its electronic structure was examined. The HOMO is now concentrated almost exclusively in the pyrrole unit opposite the protonation site, which itself is hosting the LUMO (Fig. 5A). The LUMO+1 resembles the former LUMO (cf. Fig. 2C) with the dominant contribution of the \(4p_z\) orbital of the gallium atom. The calculated solution phase FIA of \([\text{H}^{-}\text{Me1}^*]\), which was 149 kJ mol\(^{-1}\) for \([\text{Me1}^*]\), is now boosted to 191 kJ mol\(^{-1}\), pointing to an augmented Lewis acidity.

Motivated by these computational results \([\text{PPh}_4]^{[\text{Me1}]}\) and \([\text{PPh}_4]^{[\text{Et1}]}\) were treated with bis(trifluoromethyl)sulfonimide (HNTf\(_2\)) in dichloromethane–\(\text{d}_2\) as solvent. The \(^1\text{H}\) NMR spectra of the reaction mixtures showed an unselective product mixture. Computations disclosed a potential decomposition channel initiated after the formation of \([\text{H}^{-}\text{Me1}^*]\) by the cleavage of the exopyrrolic C–C bond originating from the proton-bearing carbon atom of the dearomatized pyrrole ring (see Chapter S8 in the ESI for details).\(^\ddagger\) However, a selective transformation was observed when THF–\(\text{d}_4\) was used as solvent. The \(^1\text{H}\) NMR data revealed the dearomatization of one of the pyrrole rings, besides a slightly broadened singlet resonance at 5.02 ppm, assigned to the new proton in the 2-position of the dearomatized pyrrole ring. Apparently, the Lewis basic properties of THF suppress the undesired pathway mentioned above. Indeed, remodeling the C–C bond cleavage transition state in the presence of an explicit THF molecule coordinating to Ga in \([\text{H}^{-}\text{Me1}^*]\), resulted in a significant increase in Gibbs free activation energy of 64 kJ mol\(^{-1}\) (see the ESI for details).\(^\ddagger\) When repeating the reaction with HNTf\(_2\) in a mixture of dichloromethane–\(\text{d}_2\) and diethyl ether (2 : 1 % vol), the same \(^1\text{H}\) NMR characteristics were found, supporting the Lewis base stabilization rationale.

Subsequently, the reaction with \(\text{PrOH}\) was repeated with the \(\text{in situ}\) generated \([\text{H}^{-}\text{Me1}^*]\)–Et\(_2\)O in the dichloromethane/diethyl ether mixture. After stirring at room temperature for 1 min, removing the solvent, and extracting with \(n\)-pentane/diethyl ether (4 : 1 % vol), a pale-yellow solid was isolated in 72% yield. Its \(^3\text{H}\) NMR spectrum showed the formation of a symmetric species with two dearomatized pyrrole rings due to protonation at the 2-positions and an isopropyl group with shifted signals compared to free \(\text{PrOH}\). These features align

Fig. 5 (a) Kohn–Sham frontier molecular orbital representations of \([\text{H}^{-}\text{Me1}^*]\). Hydrogen atoms, except for the one inducing dearomatization of one pyrrole ring, are not shown for clarity. (b) Reactivity of isopropanol with \([\text{PPh}_4]^{[\text{Me1}]}\) (left part) and \([\text{H}^{-}\text{Me1}^*]\)–Et\(_2\)O (right part), \(\text{in situ}\) generated by treatment of \([\text{PPh}_4]^{[\text{Me1}]}\) with bis(trifluoromethyl)sulfonimide (HNTf\(_2\)).
with the formation of the neutral O–H bond addition product \([\text{HH}^{\text{Me}}]^{+} - \text{OPr}\) by gallium–ligand cooperativity (Fig. 5B, right part). Hence, whereas \([\text{Me}^{\text{Me}}]^{+}\) itself is not capable of splitting the O–H bond by GaLC, the protonated, neutral \([\text{H}^{\text{Me}}]^{+}\)–Et\(_{2}\)O instantly adds 'PrOH. It confirms the hypothesis that the reactivity of \([\text{Me}^{\text{Me}}]^{+}\) can be enhanced by protonation of the ligand backbone.

Conclusions

In conclusion, we report on the synthesis, characterization, and reactivity of two calix[4]pyrrolato gallates(m), which feature a nearly ideal square planar coordination environment around their gallium atoms. Structural constraint empowers Lewis acidic reactivity by \(p_{z}\)-type LUMO-lowering, primarily at gallium. This was experimentally proven with donors like THF or fluoride anions, opposing Coulomb repulsion in the latter case. Reversible metal–ligand cooperative addition of CO\(_{2}\) is achieved via the dearomatization/rearomatization of one of the pyrrole units. The associated equilibrium can be fine-tuned by substituent variation in the ligand backbone. Isopropanol as a model for protic substrates did not react with the gallates in the first place, but a rapid reaction was triggered upon activation with a Brønsted acid. Overall, the reactivity of \(p\)-block element complexes of the calix[4]pyrrolato ligand and broadens the substrate scope for more challenging bond activation reactions.

Data availability

Crystallographic data has been deposited at the CCDC under 2175981–2175983. Further datasets supporting this article have been uploaded as part of the ESI.†

Author contributions

L. G. and L. M. S. devised the project and designed the experiments. L. M. S carried out the calculations, did the experimental work together with E. E. and N. R., and analyzed the data. L. G. and L. M. S. wrote the manuscript.

Conflicts of interest

There are no conflicts to declare.

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