Bright luminescent lithium and magnesium carbene complexes†‡

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We report on the convenient synthesis of a CNC pincer ligand composed of carbazole and two mesoionic carbenes, as well as the corresponding lithium- and magnesium complexes. Mono-deprotonation affords a rare “naked” amide anion. In contrast to the proligand and its mono-deprotonated form, tri-deprotonated s-block complexes show bright luminescence, and their photophysical properties were therefore investigated by absorption- and luminescence spectroscopy. They reveal a quantum yield of 16% in solution at ambient temperature. Detailed quantum-chemical calculations assist in rationalizing the emissive properties based on an Intra-Ligand-Charge-Transfer (ILCT) between the carbazolido- and mesionic carbene ligands. (Earth-)alkali metals prevent the distortion of the ligand following excitation and, thus, by avoiding non-radiative deactivation support bright luminescence.

Introduction

Carbazole-based dyes have a rich history as photo-sensitizers,1–3 photo-initiators and -catalysts,4–7 host materials for Organic Light-Emitting Diodes (OLEDs),8,9 triplet emitters,10,11 and Thermally Activated Delayed Fluorescence (TADF)12–18 materials. Recent remarkable achievements comprise the emissive properties of two-coordinate coinage metal complexes.19–32 There, embedding copper(i) in a push–pull electronic environment provided by a π-acidic carbene19 and a π-donating carbazolido ligand (Fig. 1, I), resulted in a 100% quantum yield of the 474 nm luminescence and an excited state lifetime of 2.8 μs.32 The surprising efficiency of these “carbene-metal-amido” complexes relies on a prompt Reverse Inter System Crossing (RISC) mechanism.34–36 Furthermore, the donor–bridge–acceptor substitution pattern, as it is known from organic dyes, enhances the transition probability, and thus favors fluorescence over non-radiative decay channels.37 We became interested in carbazolyl bridged pincer-type NHC ligands38–41 as promising candidates to stabilize multiple bonded late transition metal complexes.42–44 Similar ligands, which had been studied mainly as NNN pincers (Fig. 1, II),45–57 have been

![Fig. 1](image-url) Previously studied related complexes.
pioneered by Bezuidenhout58–62 (CNC pincer, C: mesoionic carbene MIC; III and IV) and Kunz63–68 (CNC pincer, C: N-heterocyclic carbene NHCl).69–73 In general, and 1,2,3-triazolylidenes, in particular, excel through their donating properties76–77 and, as such, are expected to stabilize high-valent transition metals.

Despite the fact that they are much less popular than conventional NHCs78 and less explored in photochemistry, they increasingly attract attention.52,79–89 During our initial coordination experiments with the 1,2,3-triazolylidene decorated carbazolide obtained by deprotonation of 3, we noticed strong luminescence upon deprotonation (vide infra). Intriguingly, Kunz had already noticed luminescence for lithium carbazolyl bridged dicarbones, however, these systems have not been studied spectroscopically yet (Fig. 1, III and V).59,63,64,90 Bezuidenhout and co-workers reported the photochemical properties of T-shaped and linear coinage metal complexes (Fig. 1, IV).52 They found that the luminescence wavelength is tunable by the judicious choice of the metal. Thereby, the copper(i) complex emitted in the blue- (quantum yield $\Phi_{\text{em}} = 0.8\%$), the silver(i) in the orange- ($\Phi_{\text{em}} = 2.4\%$), and the gold(i) complex in the green- ($\Phi_{\text{em}} = 0.6\%$) regions of the spectrum. The protonated pigrongid showed emissive properties as well (green, $\Phi_{\text{em}} = 2.0\%$). Heinze et al.91 described, for example, “alkaliblue” emissive pyrrolates (Fig. 1, VI). A quantum yield of 1% was achieved through an efficient ILCT (Intra-Ligand-Charge-Transfer) thanks to the templating effect of the alkali metals.92 Blue luminescence was also observed by Roesky et al. in the case of iminophosphonamide alkali metal complexes (Fig. 1, VII) with a solid-state quantum yield of up to 36%.93 Agapie and co-workers introduced lithium bridged dipyridyl dipentacene pyrrolates as efficient singlet fission molecules.94

Inspired by these results and the surge of interest in photochemistry with complexes of earth-abundant metals,95 we report herein a carbazolide bridged mesoionic biscarbene pincer ligand and a detailed investigation on the luminescent properties of its (earth-)alkali complexes. These complexes show excellent quantum yields of up to 16% at ambient temperatures in solution. Using quantum chemical calculations, we elucidate the effects of rigidity and planarity on the luminescence quantum yield and the excitation/luminescence wavelengths.

Results and discussion

Pigrrongid synthesis

Searching for an alternate route to design tridentate carbazole-triazolylidene ligands, avoiding the use of potentially hazardous and explosive tert-butylhypochloride as suggested by Bezuidenhout and co-workers,96 we initially examined the cyclation of classical triazoles, as has been reported by Limberg, Hecht and Brooker.97,98 This strategy proved also successful in case of carbenaporphyrins.99 In the present case, neither the use of methyl iodide nor Meerwein’s salt (triethyloxonium tetrafluoroborate) or methyl triflate yielded the desired carbazolebistriazolium salts in reasonable yields. Instead, mixtures with predominantly N-carbazole-methylation were observed.

To avoid the undesired N-carbazole alkylation, we consequently adopted an intramolecular alkylation strategy (Scheme 1).96 The synthesis of 1 was achieved following nitration, reduction, and azotation of commercially available 3,6-di-tert-butylcarbazole.91 Using standard CuAAC (Cu-catalyzed Azide–Alkyne Cycloaddition) conditions with 6-chlorohex-1-yne led to clean 2. The formation of the product was apparent from the disappearance of the characteristic azide stretching resonance at 2099 cm$^{-1}$ (Fig. S1†) and by the characteristic low-field $^1$H NMR resonance at $\delta = 7.98$ ppm indicative for a triazole heterocycle (Fig. S2†). Adding excess of potassium iodide and heating the mixture in acetonitrile to reflux for two days gave the $N$-fused triazolium salt 3 in essentially quantitative yields. The cyclization was evident by several features in the NMR spectra, namely (i) the low-field shift of the triazolium-$SH$ resonance in the $^1$H NMR spectrum from $\delta = 7.98$ ppm (2) to $\delta = 8.91$ ppm (3), (ii) the low-field shift of the methylene protons’ resonance of the former $–CH_2Cl$ group from $\delta = 3.64$ ppm (2) to $\delta = 4.79$ ppm (3), and (iii) the coupling of this methylene group to one of the triazol nitrogen atoms according to two-dimensional $^1$H−$^1$N HMBC spectra (Fig. S6 and S11†). X-ray quality crystals90 of 3 were obtained by slow evaporation of a saturated chloroform
solution (Fig. 2). The triazolium salt 3 crystallized with four strongly disordered molecules of chloroform in the lattice in the orthorhombic space group Pbcm with half a molecule of 3 in the asymmetric unit.

The structural metrics of the dication in 3 (Table S3†) resemble those of previously reported triazolium salts.98,99 In the solid-state structure of 3, strong hydrogen bonding interactions between the central triazolium nitrogen atoms N3 and the carbazole proton H10 were observed. Due to sterically bulk, these interactions are not feasible for the other reported MIC-CNc pincer ligands (Fig. 1 III and IV), but have been observed in a bis[pyrazolyl]carbazole derivative.100 Additionally, weak hydrogen bonding between H10–12 and H1–12 were observed in the solid-state structure of 3 (Fig. S29†).

Complex synthesis

The reaction of pincer ligand 3 with one equivalent of lithium hexamethyl disilazide [LiHMDS; LiN(SiMe3)2] led to deprotonation of the carbazole (4, Scheme 2). Evidence for the latter came from the disappearance of the resonance for the carbazole N–H group in the 1H NMR spectrum (Fig. S12†). X-ray quality crystals of deep-orange and air-stable 4, which crystallized in the P1 space group, were obtained by diffusion of hexane into a THF/benzene solution (Fig. 3). Notably, the carbazolide in 4 does not coordinate a lithium cation. Instead, the latter precipitated from the reaction mixture in the form of lithium iodide. Compound 4, therefore, contains a rare “naked” amide anion, as was also corroborated by calculations (Fig. S31†).101–103 The structure in the solid-state reveals a weak hydrogen bond [2.399(2) Å] between N10 and H2, which might be the reason for the surprising stability of 4 towards moisture. Proligand 3 was also deprotonated thrice by three equivalents of LiHMDS, as was confirmed by the 1H NMR spectroscopic analysis of 1–5 (Fig. S14†). The 7Li NMR of 1–5 showed a signal at δ = 1.43 ppm, which corroborates the presence of lithium cations (Fig. S16†). Immediately upon deprotonation of 3, strong blue luminescence was observed (vide infra) even in dim light. X-ray quality crystals of 1–5 could be obtained by slow diffusion of pentane into a diethyl ether solution of the complex (Fig. 4A).

Fig. 3 Compound 4 is devoid of N-coordinated lithium in the solid-state. Thermal ellipsoids are shown at the 50% probability level; hydrogen atoms (except the ones bonded to the triazolium heterocycles) are omitted for clarity. Selected bond lengths [Å], angles [°], and dihedral angles [°]: N10–C13 1.3734(14), N10–C19 1.3681(15), C24–N2 1.434(4), N2–C2 1.351(4), C2–C4 1.361(5), C4–N6 1.354(4), N6–N4 1.322(4), N4–N2 1.335(4), N10–H2 2.397(2), N10–I2 2.909(4), C18–N1 1.437(4), N1–N3 1.324(4), N5–N3 1.324(4), N5–C3 1.351(5), C3–C1 1.361(5), C13–N10–C19 103.1(3), N2–C2–C4 106.0(3), N1–C1–C3 105.7(3), C13–C18–N1–N3 –137.7(4), C19–C24–N2–N4 162.9(4).

Fig. 4 (A) The molecular structure of dimeric Li5 contains six lithium atoms and four bridging μ-iodo ligands. Thermal ellipsoids are shown at the 50% probability level; hydrogen atoms are omitted for clarity. The monomers are shown from different views in (B) and (C), for which the annulated six-membered rings, 1Bu groups, ethyl fragments of the diethyl ether and all hydrogen atoms have been omitted for clarity. Selected bond lengths [Å], angles [°], and dihedral angles [°]: N1–C1 1.3857(19), N1–C8 1.3899(19), C9–N5 1.4302(19), N5–N6 1.3396(18), N6–N7 1.3255(19), N7–C28 1.356(2), C28–C27 1.392(2), C27–N5 1.373(2), C27–Li2 2.136(3), Li2–I1 2.776(3), I1–Li1 2.911(3), Li1–I2 2.748(3), Li1–I2 2.911(3), C27–Li1 2.633(3), Li3–I2 2.850(3), Li3–I2 2.970(3), C21–Li1 2.391(3), C21–Li3 2.165(3), C21–N2 1.377(4), N2–N3 1.3414(17), N3–N4 1.3249(18), N4–C22 1.357(2), C22–C21 1.398(2), C2–N2 1.4348(19), N1–Li1 2.052(3), N1–Li2 2.188(3), C1–N1–C8 103.18(12), N2–C21–C22 100.22(12), N5–C27–C28 100.25(13), C1–C2–N2–N3 137.4(1), C8–C9–N5–N6 151.3(1).

Scheme 2 Deprotonation of 3 to 4 and Li5 and subsequent transmetalation to MgBr5.
I1 and Li3–I2 units could be understood as lithium iodide molecules.

The Li2–I1 bond length [2.776(3) Å] is comparable with that observed in lithium iodide adducts (2.70–2.80 Å). Also the larger Li3–I2 distance [2.970(3) Å] is in the range of previously reported lithium iodide clusters (2.98 Å). The N1 nitrogen atom of the carbazolido- (Fig. 4B and C), as well as the MIC-ligands, thus coordinate one lithium atom (Li1) and formally another molecule of lithium iodide (Li2–I1 and Li3–I2, respectively). Overall, this arrangement locks the central Li1 atom in place. Eventually, the lability of the lithium iodide was corroborated by elemental analysis (ESI†). Repeated re-crystallization lowered the equivalents of lithium iodide from two, as present in the solid-state structure shown in Fig. 4, to 0.3 equivalents.

Complex MgBr5 was prepared by transmetalation of Li5 with MgBr2, but may be as well prepared directly from 3 and the Grignard reagent MeMgBr. We were not able to obtain single crystals of sufficient quality for elucidation of the structure in the solid-state. However, a diffusion NMR experiment (DOSY) revealed that solutions of 4 and MgBr5 are mononuclear in deuterated benzene, whereas Li5 remains a dimer (page S17). Like blue-green luminescent Li5 (and in contrast to non-emissive 3 and weakly emissive 4) complex MgBr5 showed intense, lime-green luminescence in solution, whereas all investigated compounds were essentially non-luminescent in the solid state.

Luminescent properties

The bright luminescence motivated more detailed photo-physical studies of all compounds. In benzene solutions, very strong luminescence is seen for Li5, strong luminescence for MgBr5, and undetectable to low luminescence for 3 and 4 (Fig. 5). Pertinent spectroscopic features are summarized in Table 1, normalized steady-state absorption and luminescence spectra are shown in Fig. 6.

Although the bisscaticonic proligand 3 is poorly soluble in organic solvents, maxima evolved at 297 and 360 nm. It features a plateau-like shoulder peaking at 492 nm and reaching up to approximately 570 nm. Upon photoexcitation in benzene, 3 is found to be non-luminescent. In contrast, in the absorption spectra of 4, we find maxima at 352, 445, and 487 nm. Photoexcitation of 4 at, for example, 390 nm leads to a broad and undefined luminescence with a maximum at 565 nm and a 2% emission quantum yield.

| Compound | λabs,max/nm | λem/nm | Φem/% | ΕStokes/eV (cm⁻¹) |
|----------|-------------|--------|--------|-------------------|
| 3        | 360, (492)b | —      | —      | —                 |
| 4        | 352, 445, 487 | 565 | 2 | 0.35 (2835) |
| Li5      | 325, 402, 465 | 506 | 16 | 0.23 (2016) |
| MgBr5    | 339, 380, 431 | 482 | 14 | 0.30 (2455) |

a Spectroscopic data were obtained at room-temperature for 1 × 10⁻² M benzene solutions of 3, 4, Li5, and MgBr5. The emission wavelengths and quantum yields were obtained after excitation at 390 nm. b The plateau is assigned to vibronic transitions as discussed below.
\( \phi_{\text{em}} \) (Fig. 6). Turning to the absorption spectrum of \( \text{Li}^5 \), two maxima are discernable at 325 and 402 nm. A tail is superimposed onto the latter all the way to 600 nm including a minor shoulder centered at around 465 nm. Excitation spectra for \( \text{Li}^5 \) and \( \text{MgBr}^5 \) reveal that the lowest absorption wavelength which leads to luminescence is 465 nm for \( \text{Li}^5 \) and 434 nm for \( \text{MgBr}^5 \). In both cases, this is in line with their absorption maxima (Fig. S21†). We determined Stokes shifts of 0.35, 0.30, and 0.23 eV for \( \text{Li}^5 \), \( \text{MgBr}^5 \) and \( \text{Li}^5 \), respectively.

Excitation of \( \text{Li}^5 \) at 390 nm gives a luminescence maximum at 506 nm and a luminescence quantum yield of 16%. The luminescence spectrum of \( \text{Li}^5 \) also shows a tail up to 725 nm. Turning finally to \( \text{MgBr}^5 \), we note absorption maxima at 339, 380, and 431 nm. Once again, the latter features a tail up to approximately 560 nm.

Photoexcitation of \( \text{MgBr}^5 \) at 390 nm leads to a luminescence that reaches a maximum at 487 nm with a quantum yield of 14%. Again, a tail up to 730 nm is noted. Upon exposure of solutions of \( \text{Li}^5 \) and \( \text{MgBr}^5 \) to air, the spectroscopic signatures of mono-deprotonated \( \text{Li} \) were regenerated (Fig. S21 and S22†). Time-Correlated Single-Photon Counting (TCSPC) experiments on \( \text{Li}^5 \) and \( \text{MgBr}^5 \) revealed a luminescence decay with lifetimes of (11.8 ± 1.6 \( \times \) 10\(^{-2} \)) and (10.7 ± 4.8 \( \times \) 10\(^{-2} \)) ns, respectively (Fig. S23 and S26†). By virtue of lifetime components which are in the typical range for singlet excited states, we rule out the involvement of triplet excited state species as is the case for TADF or phosphorescence. In other words, the luminescent deactivation is fluorescence.

### Computational analysis

Puzzled by the optical properties, we investigated the electronic and structural properties of all compounds in their excited states. Exploratory TD-DFT calculations revealed that the transitions to the S\(_1\) state of 3 and 4 are ILCT (approximated HOMO \( \rightarrow \) LUMO, Fig. 7A and C) processes.

The HOMO is located at the carboxalid ligand, whereas the LUMO is associated with the MICs (cf. Fig. S32†). For an accurate description of the charge-transfer states, which is challenging for TD-DFT\(^{111-113}\) (Fig. S33–S35†), absorption spectra were computed with the more suitable \( \text{ab initio} \) method “Similiarly Transformed Equation of Motion Coupled Cluster Singles and Doubles” (STEOM-CCSD)\(^{114-116}\) using the “Domain-based Local Pair Natural Orbital (DLPNO)” approximation.\(^{117,118}\) Indeed, this method reproduced the absorption spectra including the transitions to the \( \text{S}_1 \) states best (3, \( f_{\text{osc}} = 0.13 \), \( \text{calcd}_{\text{abs}} = 357 \text{ nm} \), \( \text{exp}_{\text{abs}} = 360 \text{ nm} \), Fig. 7B; 4, \( f_{\text{osc}} = 0.19 \), \( \text{calcd}_{\text{abs}} = 466 \text{ nm} \), \( \text{exp}_{\text{abs}} = 487 \text{ nm} \), Fig. 7D).

The luminescence from the \( \text{S}_1 \) state of 4 is predicted to be weak (\( f_{\text{em}} = 0.01 \), \( \text{calcd}_{\text{em}} = 528 \text{ nm} \), \( \text{exp}_{\text{em}} = 565 \text{ nm} \), Fig. 7D) which is in agreement with the observation that 4 is weakly luminescent with a 2% quantum yield. For 3, neither TD-DFT nor STEOM-CCSD reproduced the plateau between 425 and 525 nm. However, the calculation of the vibronically resolved absorption spectrum using excited state dynamics (Fig. S36†) shows that this shoulder arises from molecular vibrations.

Subsequently, we investigated the electronic structures of the complexes 5. The calculations revealed that the dimeric structure of \( \text{Li}^5 \) could be well modelled by calculating the electronic structure of the truncated monomer, thereby omitting the diethyl ether molecules as well (Fig. S32†).

A bright transition to the \( \text{S}_1 \) state \( \text{Li}^5_{\text{truncated}} \) (\( f_{\text{osc}} = 0.17 \), \( \text{calcd}_{\text{abs}} = 338 \text{ nm} \), \( \text{exp}_{\text{abs}} = 402 \text{ nm} \)) was also predicted for \( \text{Li}^5 \) (Fig. 8B). This transition originates also from the ILCT from the carboxalido- to the MIC ligand (Fig. 8). The same is true for \( \text{MgBr}^5 \) (\( f_{\text{osc}} = 0.17 \), \( \text{calcd}_{\text{abs}} = 407 \text{ nm} \), \( \text{exp}_{\text{abs}} = 431 \text{ nm} \)) (Fig. 8D).

Luminescence from the \( \text{S}_1 \) state of the complexes 5 is bright \( \text{Li}^5 \), \( f_{\text{osc}} = 0.10 \), \( \text{calcd}_{\text{em}} = 408 \text{ nm} \), \( \text{exp}_{\text{em}} = 506 \text{ nm} \), \( \text{MgBr}^5 \) \( f_{\text{osc}} = 0.20 \), \( \text{calcd}_{\text{em}} = 435 \text{ nm} \), \( \text{exp}_{\text{em}} = 482 \text{ nm} \), Fig. 8B and D, respectively) and in agreement with 5 being luminescent with quantum yields of 16% for \( \text{Li}^5 \) and 14% for \( \text{MgBr}^5 \). The STEOM-CCSD calculations also reproduce the experimental Stokes shifts (\( \text{Li}^5_{\text{calcd}} = 0.31 \), \( \text{exp}_{\text{calcd}} = 0.35 \); \( \text{Li}^5_{\text{truncated}} \) \( \text{calcd} = 0.40 \), \( \text{exp} = 0.25 \); and \( \text{MgBr}^5_{\text{calcd}} = 0.20 \), \( \text{exp} = 0.30 \) eV).
Fig. 8 The dominant absorptions and concomitant excitations to the S1 states of \(^{15}\)S and \(^{19}\)BuS belong to the HOMO \(\rightarrow\) LUMO transitions (B and D, red). For both \(^{15}\)LiS and \(^{19}\)MgBrS, the HOMOs (left) are located at the carbazole, whereas the LUMOs (right) are mainly localized on both MIC units (A and C). For both \(^{15}\)LiS and \(^{19}\)MgBrS, the luminescence from the S1 states is bright (B and C, blue). Orbitals were obtained at the TD-DFT(SMD=C\(_6\)H\(_6\))/def2-TZVPP//B3LYP-D3(BJ)/def2-SVP level of theory, whereas the luminescence spectra were obtained at the STEOM-CCSD(SMD=C\(_6\)H\(_6\))/def2-TZVPP//B3LYP-D3(BJ)/def2-SVP level of theory. Hydrogen atoms are omitted for clarity.

Considering the vastly differing luminescent properties of 3, 4, and complexes 5, we evaluated the geometries of the first S\(_1\) excited states. The proligand 3 is distorted in the relaxed S\(_1\) state, in which a triazolium group bends out of plane (Fig. 9, left). This means that the excitation is followed by a change of the mean value of the C=C=N-N dihedral angles before and after excitation (\(\Delta\angle_{\text{C-C-N-N}}\)) between the carbazole and the triazolium unit of 90\(^\circ\) (S\(_0\); \(\angle_{\text{C-C-N-N}} = 102^\circ\) and S\(_1\); \(\angle_{\text{C-C-N-N}} = 100^\circ\)). A comparable degree of distortion was also derived from the root-mean-square deviation for the change of the positions of all atoms (RMSD\(_{S0-S1}\); Fig. S38).

Upon the photoexcitation of 4, a MIC group bends out of plane as well (Fig. 9, right) with \(\Delta\angle_{\text{C-C-N-N}} = 42^\circ\) (S\(_0\); \(\angle_{\text{C-C-N-N}} = 177^\circ\) and S\(_1\); \(\angle_{\text{C-C-N-N}} = 135^\circ\)). \(^{15}\)Li essentially retains the ground-state geometry in the relaxed S\(_1\) state, due to the fact that the coordinated metal ions lock the MIC ligands in place (Fig. 10, left). Throughout the excitation, the \(\Delta\angle_{\text{C-C-N-N}}\) changes only by 7\(^\circ\) (S\(_0\); \(\angle_{\text{C-C-N-N}} = 139^\circ\) and S\(_1\); \(\angle_{\text{C-C-N-N}} = 146^\circ\)).

The anchoring effect of the metal is also evident for \(^{19}\)MgBrS, which retains a pseudo-square-planar coordination around the magnesium ion (Fig. 10, right). Here, the \(\Delta\angle_{\text{C-C-N-N}}\) is 11\(^\circ\) (S\(_0\); \(\angle_{\text{C-C-N-N}} = 177^\circ\) and S\(_1\); \(\angle_{\text{C-C-N-N}} = 166^\circ\)). Intriguingly, the computationally determined degree of distortion in the excited S\(_1\) state correlates well with the experimental Stokes shifts and quantum yields.
quantum yields (Fig. 11). For instance, 4 shows a large Stokes shift of $\Delta E = 0.35$ eV and $\Delta \angle_{C-C-N-N} = 42^\circ$, while $^{115}$S exhibits a Stokes shift of $\Delta E = 0.25$ eV and $\Delta \angle_{C-C-N-N} = 7^\circ$.

Berlman suggested that chromophores, in which the ground and first excited states are planar, show high quantum yields,119,120 because molecular distortions and rotations in the relaxed excited state often lead to non-radiative relaxation.121-123

This rigidification or Restriction of Intramolecular Motions (RIM) principle is of course crucial for organic fluorophores including ubiquitous BODIPY.124,125 It has also been exploited in sensing and coordination chemistry as the Chelation-Enhanced Fluorescence (CHEF) effect, in which the ground and excited geometries of a molecule are locked into a planar conformation by coordinating metals.126-128

Accordingly, complexes with comparably small structural distortions in the excited state are more efficient emitters ($^{115}$S

$\Delta \angle_{C-C-N-N} = 7^\circ$, $\phi^{em} = 16\%$; $^{115}$MgBr

$\Delta \angle_{C-C-N-N} = 11^\circ$, $\phi^{em} = 14\%$; 4 $\Delta \angle_{C-C-N-N} = 42^\circ$, $\phi^{em} = 2\%$; 3 $\Delta \angle_{C-C-N-N} = 90^\circ$, non-emissive). Therefore, we conclude that the structural relaxation of the excited S$_1$ state quenches the luminescence. In contrast, the alkali- and earth-alkali metals lock the conformation and, hence, allow for bright luminescence with comparably small Stokes shifts.

Conclusions

We report on the synthesis of an N-fused CNC pincer ligand composed of carbazole and two triazolium units. The synthetic approach, scalable to multigram quantities, avoids the use of hazardous tert-butylhypochlorite, which had found use for related ligand systems. The pincer ligand undergoes single deprotonation to afford a rare “naked” amide, which is air-stable due to intramolecular hydrogen-type bonding interactions. Triple deprotonation by a lithium base affords a chelated, binuclear lithium complex, which undergoes transmetalation with magnesium. Photophysical investigations show that the s-block complexes excel with luminescence quantum yields of up to 16% at ambient temperature and in solution, whereas the pro- and mono-deprotonated ligands are essentially non-luminescent. Detailed quantum-chemical calculations helped to rationalize the luminescent properties with an Intra-Ligand-Charge-Transfer (ILCT) from the carbazole to the mesoionic carbene. (Earth-)alkali metals prevent the distortion of the ligand following excitation and, in turn, enable bright luminescence in the blue to green region of the spectrum.

Author contributions

PP and SH prepared the compound 4, PP prepared as well complexes $^{115}$Li5, $^{115}$MgBr5 and ran the computations, CMS supported by TU performed the photochemical studies, FAW and ND synthesized the compounds 1, 2 and 3, RHI performed the XRD analysis of 3, SH performed the XRD analysis of 4, BM performed the XRD analysis of $^{115}$S, AG contributed to the synthesis of $^{115}$S and $^{115}$MgBr5, MZ performed the DOSY experiments, DMG, SH and DM coordinated the project, DM conceived the idea. The manuscript was proof-read and approved by all authors.

Conflicts of interest

There are no conflicts to declare.

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