Tuning the luminescence of transition metal complexes with acyclic diaminocarbene ligands

Mikhail A. Kinzhalov, Elena V. Grachova and Konstantin V. Luzyanin

Metal complexes featuring acyclic diaminocarbene ligands have recently emerged as powerful emitters for prospective use in OLEDs and other electroluminescent technologies. Owing to their strong π-donating properties and broad synthetic availability, acyclic diaminocarbenes have successfully challenged the application of other ancillary ligands for fine-tuning of luminescence emission. This review systematises the data reported concerning the preparation of luminescent metal complexes bearing acyclic diaminocarbene ligands, uncovers their photophysical properties as a function of the carbene ligand structure and metal characteristics, and draws attention to the potential of ADCs and the advantages that the application of metal-(acyclic diaminocarbenes) provides to the family of transition-metal-based phosphors.

1. Introduction

Luminescent transition metal complexes underpin the development of OLEDs and other electroluminescent technologies (for the main reviews on the topic see ref. 1–10), and a broad range of applications in the fields of luminescence chemosensing,11–13 photocatalysis,14–16 sensitisation,17–19 self-assembled materials,20 and bioimaging.20–23 Although pure organic light emitters are predominantly fluorescent (triplet excitons formed are deactivated by thermal processes), the presence of strong spin–orbit coupling in transition metal complexes upholds the intersystem crossing from the singlet to the triplet state enabling phosphorescent relaxation. An electroluminescent device doped with a transition metal complex has a maximum quantum yield Φ = 1 for the main photon absorption, and by placing electrons into metal–ligand σ*-antibonding orbitals, causing the gradual degradation of the compound by ligand loss.1 One of the solutions that emerged is the use of a secondary set of σ-donor ligands, which would additionally destabilize unoccupied d-orbitals and their corresponding metal-centred excited states, and improve the efficiency and stability of blue phosphorescence. As a result, strong σ-donor carbene ligands, i.e., N-heterocyclic carbenes (NHCs),42 acyclic diaminocarbenes (ADCs),165 and cyclic alkylaminocarbenes,166 have been evaluated as organometallic emitters (Fig. 1).

The prevailing part of examples discloses the photophysical properties of metal–NHC species (for reviews on the topic see ref. 30–33; for recent manuscripts see ref. 34–38); only a few reports describing the photoluminescence properties of cyclic alkylaminocarbenes have recently emerged.39–41 At the same time, a number of experimental studies considering luminescent metal–ADC species have now reached a level, where the first survey to consider these diverse ADC-containing organometallic phosphors is timely.

The essential goals of this review are (i) to systematise the reports published until now describing the preparation and the photophysical properties of metal–ADCs; (ii) to compare, whenever possible, the photophysical properties of metal–ADCs against metal–NHC species, and (iii) to draw attention to the potential of ADCs and the advantages that the application of ADCs gives to organometallic chemistry and materials science.
2. Acyclic diaminocarbenes as ligands

NHCs and ADCs are neutral ligands featuring divalent carbon centres with six-electron valence shells. Both carbene types exhibit a singlet ground-state electronic configuration with a HOMO (Highest Occupied Molecular Orbital) and LUMO (Lowest Unoccupied Molecular Orbital) best described as a formally sp2-hybridised lone pair and an unoccupied p-orbital at the carbene carbon, respectively. The relative stability of diaminocarbenes is ensured by the presence of mutually ω-electron-withdrawing and ρ-electron-donating nitrogens both inductively (by lowering the energy of the occupied ω-orbital) and mesomerically (by donating electron density to the empty ρ-orbital) (Fig. 2). The principal consequence of the ground-state electronic structure of diaminocarbenes is their propensity to act as ω-donors and bind to a wide range of metal centres. While strong ω-donor and comparatively weak π-accepting properties of aminocarbenes suggest the coordination properties closely resembling those of phosphines, the former are in general more electron-donating than the latter.  

This is reflected in a stronger and shorter metal–ligand bond (as judged by its length and greater bond dissociation energies) in diaminocarbone complexes when compared to their phosphine counterparts. Due to the lower electronegativity of carbon compared to nitrogen, the in-plane lone electron pair on the former possesses significantly higher energy than that in common N-donors such as pyridyl ligands. Fig. 3 shows the bonding interactions between diaminocarbenes and transition metals; the major contribution to bonding is due to ω-donation that effectively destabilises the transition metal Cσ ω σ* orbitals, with metal-to-π-backdonation interactions constituting relatively minor effects.

The absence of a cyclic fragment in the ADCs causes significant changes in their steric and electronic characteristics in comparison to NHCs. Owing to the lack of electron density delocalisation (as no aromatic heterocyclic fragment is present in the structure), ADCs are generally more electron-donating than aromatic-NHCs. Moreover, ADCs are more basic and nucleophilic species than their cyclic non-aromatic counterparts insofar as a wider N-Carbene–N bond angle decreases the s-character of the lone pair on the carbene carbon. The better donor ability of ADCs and an increased electronic density on a metal destabilise the ligand-field states, potentially leading to higher quantum yields for pure deep blue photoluminescence and expected performance in OLED devices (Fig. 4, left).

Due to the acyclic structure, rotation around the C-Carbene–N bonds is enabled allowing for the acyclic diaminocarbene ligand to adopt various conformations with differing steric and electronic characteristics. For instance, in carbene species having small substituents, a rotation barrier of the C-Carbene–N bond is lower than 13 kcal mol⁻¹ as indicated by experimental and theoretical studies. Rotational flexibility plays a significant role in the photoemissive properties; different conformers of ADCs exhibit different energies of the triplet metal-to-ligand charge-transfer (MLCT) states (Fig. 4, right).

Another important observation regarding metal–ADCs is their synthetic availability since several strategies for their preparation are established and widely used. Valuable properties of complexes of transition metals with diaminocarbene ligands associated with their photophysical applications are determined by the characteristics of the metal centre, as well as by the balance between the donor and steric properties of the diaminocarbene ligand. Each application requires complexes and, therefore, ligands with a different set of parameters, that are screened empirically, imposing stringent requirements on the method of their synthesis.

Fig. 1 Representative luminescent transition metal complexes supported by cyclometallated phenylpyridines, NHCs²⁶,²⁷ and ADCs.²⁸,²⁹

Fig. 2 Heterocyclic (NHC)³² and acyclic (ADC)¹⁶⁵ diaminocarbenes, and their corresponding metal complexes.
same time, the most widespread approach is based on nucleophilic addition to metal-bound isocyanides, leading to the production of diminocarbene metal complexes without the release of free carbene. This approach is stoichiometric, atom-efficient and high-yielding as a rule allowing for an easy variation of the periphery of ADCs, thus ensuring high synthetic flexibility (Scheme 1). The elegance of this approach consisting basically of one step and leading to diverse well-defined metal–diaminocarbene species is in sharp contrast to the strategies for the generation of custom-made N-heterocyclic carbenes and cyclic alkylaminocarbenes, where only the ligand synthesis may require several consecutive steps. Note that all luminescent ADC complexes known up to date and described in this review were obtained by the isocyanide pathway.

### 3. Luminescent [M]-ADC complexes

Luminescent complexes of four transition metals, viz. Ir
\(^{III}\), Pt
\(^{II}\), Re
\(^{I}\), and Au
\(^{I/III}\), bearing ADC ligands are considered in the

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**Scheme 1** Representative examples of Ir
\(^{III}\)–ADC complexes prepared by nucleophilic addition to metal-bound isocyanides.28,49,56,57
current review. All the examples discussed belong to the 5d row of transition metals, and although many 4d metal–ADC complexes are known, to the best of our knowledge none of them have been described as luminescent. While the excellent donor capacity and binding properties of ADC ligands make them suitable candidates for the construction of luminescent systems based even on first-row transition metal complexes, including iron, no significant progress has been made in this area to date. Nevertheless, it has been demonstrated that the donor properties of ADCs can destabilize the metal-localized ligand-field excited states in Fe complexes, affecting their optical properties.

The first example of an emissive transition metal ADC complex was reported by Che and coworkers for Pt species in the late 90s, but an explosion of research in this area took place several years ago, starting with the discovery of Ir←ADC complexes by Teets and Luzyanin groups. Of note, some of the complexes depicted below feature deprotonated forms of ADCs, that are called aminocarbenes for the sake of simplicity (Scheme 2).

The combination of ADCs and cyclometallating (C^N or C^C) ligands around the Ir← centre can potentially lead to bright triplet luminescence (phosphorescence) due to the strong spin–orbit coupling characteristics of Ir← complexes. Bis-C^N-cyclometallated Ir← complexes bearing neutral “Chugaev-type” deprotonated C^C-chelating bis(diaminocarbene) ligands 4–6 were prepared via the addition of excess hydrazine to isocyanides in 1–3 (Scheme 2). Protonation of 4–6 with one equivalent of HBF4 led to cationic bis(diaminocarbene) derivatives 7–9. Electrochemical studies were used to assess the energies of frontier orbitals; the oxidation potentials of cyclometallated Ir← complexes are linked to the HOMO energy levels consistent with the removal of an electron from an Ir←-centred orbital. The first oxidation potential of 4–9 corresponds to Ir←/Ir← transition (primarily metal-centred) and is located at 0.86–1.02 V. These electrochemical potentials are somewhat similar to those for cyclometallated Ir← complexes bearing NHClCs (0.97–1.10 V). When compared to starting bis-arylisocyanides species 1–3 (1.63 V, 1.68 V, and 1.44 V, respectively), Ir←-ADCs 4–9 are 0.58–0.66 V easier to oxidise, suggesting the destabilisation of the HOMO levels upon conversion of both isocyanides to a C^C-chelating ADC ligand.

Metal–ADCs 4–9 display 1,3-CT absorption bands tailing into the visible region (Fig 5) in contrast to bis-isocyanides 1–3, where those occur under 400 nm. In solution, 4 and 5 are blue emitters ($\lambda_{max}$ 473–474, $\Phi$ 0.044 for 4 or 0.010 for 5) (Fig 5). The emission spectra of 4–6 are considerably redshifted by ca. 8–12 nm (360–410 cm$^{-1}$) from those of the starting bis-isocyanides 1–3, which can be attributed to the decrease of the $\pi$-accepting properties and the increase of the $\sigma$-donating properties of the ADC ligand, leading to the destabilisation of the HOMO for 4–6. 6 emits in solution in the green to yellow regions with $\lambda_{max}$ at 519 nm and $\Phi$ of 0.19. Protonation of 4–6 results in a 2–4-fold increase in quantum yields for 7–9 ($\Phi$ 0.08 for 7, 0.02 for 8, and 0.60 for 9). Immobilisation of 4, 6, 7, and 9 in PMMA (2 wt% of the Ir← complex) brings a significant increase in $\Phi$ ($\Phi$ 0.56 for 4, 0.90 for 6, 0.68 for 7, and 0.86 for 9); both the emission energy and spectral profiles are retained. An increase in $\Phi$ can be justified as a result of decreasing the non-radiative vibrational relaxation of the excited state in the rigid environment. Even though 4–6 exhibit only moderate photoluminescence properties, this study illustrates that both electrochemical and photophysical properties are significantly impacted upon conversion of isocyanide to carbene species. As a result, this metal-mediated approach to ligand design features an alternative to a classic ligand exchange approach towards the tuning of excited states and subsequently the luminescent profiles of metal complexes.

The neutral tris-chelated cyclometallated Ir← complexes supported by the ADC ligands were generated via the coupling of amines with Ir←-isocyanides 10 and 11 (Scheme 3). Upon the nucleophilic attack by an amine, a subsequent base-promoted intramolecular activation of an aromatic C–H bond
yields the final ortho-metallated Ir\textsuperscript{III}–ADCs 12–15. The first oxidation potentials of 12 and 13 (0.52 and 0.54 V) are cathodically shifted by at least 0.25–0.28 V when compared to those of relevant Ir\textsuperscript{III}–NHC \([\text{Ir(Fppy)}_2(C,N\text{-NHC})]\) \((C,N\text{-NHC} \text{ is } C,C'\text{-cyclo-}
\text{metallated 3-methyl-1-((4-(trifluoromethyl)phenyl)phenyl)-1H-imida-
\text{zol-3-ium-2-ide}, 0.82 V or } C,C'\text{-cyclo-metallated } C,C'\text{-cyclo-
metallated 3-methyl-1-((4-(trifluoromethyl)phenyl)-1H-benzo[}\text{d}
\text{]imidazol-3-ium-2-ide}, 0.78 V).\textsuperscript{79} This comparison demonstrates
significantly higher-lying HOMO levels in the ADC complexes
than in their cyclic counterparts, consistent with the fact that
ADC is a more electron-rich donor than NHC.\textsuperscript{48}

Iridium(III)–ADCs 12–15 exhibit reduced lowest-energy absorption bands tailing to 450 nm for F\textsubscript{2}ppy (12, 13) and
500 nm for bt (14, 15) complexes attributed to both singlet
and triplet MLCT transitions (Fig. 6). The energies of the
MLCT transitions are affected by varying the cyclo-metallating
C\textsuperscript{N} ligand, while alteration of the ADC periphery induces a
minor change in the extinction coefficients without affecting
notably the MLCT bands. In CH\textsubscript{2}Cl\textsubscript{2} solution at RT, complexes
12 and 13 exhibit blue phosphorescence with \(\lambda_{\text{max}}\) values at
498 and 495 nm and moderate \(\Phi\) values (0.22 and 0.18,
respectively). In solution, the yellow-orange emission of 14
and 15 has significantly lower \(\Phi\) values (0.049 for 14, 0.047 for 15).
Immobilisation of metal–ADCs in the PMMA matrix increases
\(\Phi\) to 0.79 (for blue emitters 12 and 13) and 0.30–0.37 (for
orange emitters 14 and 15) (Fig. 6). Computational studies
showed that iridium-cyclo-metallated species possess typical
characteristics of thermally activated delayed fluorescence
emitters with potential to be used as novel electroluminescent
devices.\textsuperscript{82}

A combination of DFT and TD-DFT calculations were used
to predict the multi-state redox switchable nonlinear optics
response\textsuperscript{9} of iridium(III) ADC complexes like 7–9\textsuperscript{83}
and 12–15.\textsuperscript{84} The nonlinear optics response can be easily modu-
lated by one electron redox processes and Ir\textsuperscript{III} complexes like
7–15 can be applicable for redox second-order nonlinear optic
switches with two redox states.

An approach aimed towards the increase of the emission
energy of cyclo-metallated iridium complexes focused on the
destabilisation of the LUMO have been reported.\textsuperscript{24,33,85}
Replacing the pyridyl ring of the C\textsuperscript{N} ligands with an
N-heterocyclic carbene leads to a significant increase in the
LUMO energy and, consequently, increases the emission
energy of complexes. The majority of the recent achievements
depicting this approach involved homoletic tris-cyclo-
metallated Ir(C,N\text{-NHC})\textsubscript{3} complexes,\textsuperscript{34,38,85–89}
showing blue or near-
UV phosphorescence at room temperature. A cascade reaction
involving nucleophilic addition of propyl amine to a co-
ordinated isocyanide in 16–18 with a subsequent base-assisted
cyclometallation (Scheme 4) led to the heteroleptic cyclo-
metallated derivatives 19–21 featuring two C,C\text{-NHC} and one C,C\text{-}
ADCs.\textsuperscript{90} The first oxidation potential of 19–21 falls within the
0.16–0.25 V range, akin to those for homoletic mer-[Ir(C,C\text{-NHC})\textsubscript{3}]
complexes (0.14–0.23 V).\textsuperscript{34,85} Both 19–21 and the afore-
mentioned mer-[Ir(C,C\text{-NHC})\textsubscript{3}] are difficult to reduce due to
the high energy of the LUMO, which is being responsible for
the large HOMO–LUMO gap in this type of Ir\textsuperscript{III} compounds.
19 is not luminescent in solution at RT, while 20 shows a weak
blue (\(\lambda_{\text{max}}\) 418 nm, \(\Phi\) 0.013) and 21 green (\(\lambda_{\text{max}}\) 511 nm, \(\Phi\)
0.39) emissions. Immobilisation in the PMMA film (2 wt%) increases \(\Phi\) for 19 and 20 to 0.13 and 0.31 without affecting
\(\lambda_{\text{max}}\). For 21 in the PMMA film, a hypsochromic shift for
luminescence is accompanied by an increase in \(\Phi\) (\(\lambda_{\text{max}}\)
459 nm, \(\Phi\) 0.48), Fig. 7. When compared to homoletic mer-
[Ir(C,C\text{-NHC})\textsubscript{3}]\textsuperscript{34,85} already used for the manufacturing of deep
blue OLEDs, the mixed-carbene species 19–21 exhibit similar
or superior photoluminescence characteristics, i.e. higher \(\Phi\).

Metal-mediated addition of NH\textsubscript{3} to the coordinated isocy-
anides in 22–29 (Scheme 5) allows the preparation of another

![Fig. 6](image-url)
family of heteroleptic cyclometallated IrIII species bearing ancillary ADCs.\textsuperscript{28,68} Oxidation of the thus prepared ADCs 30–45 is manifested by an irreversible wave centred at metal (0.77–1.22 V). Conversion of one CNR in 26–29 to ADC in 34–37 brings about a decrease in the oxidation potential by ca. 0.2 V; generation of the second ADC in 38–41 induces a further drop of the oxidation potential by ca. 0.3 V.

Variation of the halogen substituent in the aryl fragment of the CNR or the corresponding ADC ligands induces no significant change in the positions of the singlet MLCT bands and only exerts a minimal influence on the extinction coefficients (Fig. 8, top). Conversion of the CNRs to ADCs impacts those transitions: a ca. 30 nm (ca. 2000 cm\(^{-1}\)) redshift of the singlet MLCT absorption bands occurs on going from 26–29 to 38–41 (Fig. 8, bottom, for representative examples of compounds with a bromo-substituted phenyl fragment).\textsuperscript{28}

Destabilisation of the HOMO as a result of the ADC ligand having lower \(\pi\)-accepting and higher \(\sigma\)-donating properties when compared to isocyanide is suggested as a reason for the singlet MLCT transition shift. Compounds 38–41 with two ADC ligands exhibit blueshifted singlet MLCT bands relative to the complexes with ADC/Cl (30–33) or ADC/CN (42–45) ligand environments insofar as strong \(\sigma\)-donation from ADC increases the HOMO–LUMO gap.

In solution, upon photoexcitation by UV light complexes all 22–45 exhibit a blue-green phosphorescence (Fig. 9).\textsuperscript{28} Conversion of 22–25 (\(\lambda_{\text{max}}\) 469 nm, \(\Phi\) 0.10 for 22) to 30–33 (\(\lambda_{\text{max}}\) 502 nm, \(\Phi\) 0.04 for 26) is accompanied by a bathochromic shift of emission energy and drop in the \(\Phi\). These relatively small \(\Phi\) values for 30–33 photoemission can be explained by the presence of chlorides which have a smaller ligand field splitting parameter\textsuperscript{41} and stabilise the non-emissive metal centred states.\textsuperscript{92,93} A redshift of the \(\lambda_{\text{max}}\) upon the transformation of coordinated CNR to ADC could be explained by the lower \(\pi\)-accepting and higher \(\sigma\)-donating properties of ADC\textsuperscript{59} that subsequently leads to the destabilisation of the HOMO,\textsuperscript{81} as confirmed by a cathode shift of the oxidation potential by ca. 0.3 V. A similar effect was observed for 26–29 (\(\lambda_{\text{max}}\) 451 nm, \(\Phi\) 0.65 for 27), 34–37 (\(\lambda_{\text{max}}\) 458 nm, \(\Phi\) 0.40 for 35), 38–41 (\(\lambda_{\text{max}}\) 472 nm, \(\Phi\) 0.31 for 39) and 42–45 (\(\lambda_{\text{max}}\) 473 nm, \(\Phi\) 0.45 for 43). A further theoretical investigation confirmed that the transformation of the isocyanide to the ADC ligand within the 22–45 family increases the relative contribution of the iridium d orbitals to the HOMO (23 31%, 31

**Fig. 7** Emission spectra of 20 and 21 in CH\(_2\)Cl\(_2\) and in the PMMA film at RT, and in 1:3 CH\(_2\)Cl\(_2\)/toluene glass at 77 K (\(\lambda_{\text{ext}}\) 310 nm). The insets show photographs of the PMMA films under UV illumination. Adapted with permission from ref. 90. Copyright (2019) The Royal Society of Chemistry.

**Fig. 8** UV-vis spectra of the representative Ir\textsuperscript{III} species of the 22–45 family in MeCN at RT. Adapted with permission from ref. 28. Copyright (2020) American Chemical Society.

**Fig. 9** Photoluminescence spectra of the representative Ir\textsuperscript{III} species of the 22–45 family recorded in MeCN at RT (\(\lambda_{\text{ext}}\) 350–400 nm). Adapted with permission from ref. 28. Copyright (2020) American Chemical Society.

**Scheme 5** Preparation of Ir\textsuperscript{III}–ADC complexes 30–45.\textsuperscript{28,68}
40%, 27 19%, 35 25%, 39 34%) inducing a drop in the HOMO–LUMO gap and enhanced mixing of the singlet MLCT and triplet LC states.\textsuperscript{28}

Comparison of 38–41 with the structurally related complexes with NHCS shown in Fig. 1 led to the conclusion of a rather similar $\lambda_{\text{max}}$, as well as lifetimes of the excited states. This suggests that the emission properties of [Ir(C$^N$)\textsubscript{2}L\textsubscript{2}]$^+$ complexes with carbene-based ancillary ligands are primarily controlled by the nature of C$^N$ ligands, in contrast to the classical approach of charged complexes [Ir(C$^N$)$_2$(N$^N$)]$^+$ bearing chemically tunable N$^N$ ancillary ligands with low-energy $\pi^*$ orbitals. The larger values of $\Phi$ for these ADC derivatives (0.43) when compared to NHC analogues (0.38) shown in Fig. 1 are explained by the less accessible higher-lying triplet MC excited states at room temperature for ADC complexes due to their superior donor properties.

Based on complexes 42–45 with ADC and cyano groups, ancillary ligands proposed a new class of organometallic chemosensors for the quantitative determination of mercury.\textsuperscript{28} 42–45 react selectively with Hg$^{2+}$ ions, which is accompanied by a linear decrease in the luminescence intensity; limit of detection of mercury(II) ions as low as $2.63 \times 10^{-7}$ M, which makes 42–45 one of the most sensitive photoluminescent chemosensors for the detection of mercury in solution.

Cyclometallated Ir$^{III}$ species 48–53 with deprotonated C$^N$N-chelating ADC species represent a novel class of luminescent mechanochromic systems. Species 48–53 were prepared by the replacement of the chloride ligand of isocyano precursors with substituted 2-aminopyridines followed by an intramolecular nucleophilic attack (Scheme 6).\textsuperscript{49} 48–53 exhibit two moderate absorption shoulders at 340–360 and 420–440 nm, ascribed to an admixture of LC and MLCT [$d_d$(Ir) $\rightarrow$ $\pi^*$(C,N-ADC)] transitions. In solution, complexes 48–53 display intensive triplet photoluminescence at RT with $\Phi$ ranging from 0.25 to 0.59 (Fig. 10). Solution emissions are attributed to phosphorescence derived from the predominant triplet MLCT [$d_d$(Ir) $\rightarrow$ $\pi^*$(C,N-ADC)] state mixed with triplet LC[ppy] character as confirmed by computational studies for 48.

In the solid state 48–53 emit strong sky blue to yellow luminescence. Mechanical stimulation of 48–53 leads to a redshift of the emission tentatively associated with a conformational change of the ADC ligand (Scheme 6 and Fig. 11). Different orientations of the N-aryl moiety of the ADC ligand is reflected in the crystal packing affecting the energy of MLCT transitions and subsequently the photophysical properties of the complexes. The contribution of this effect depends on the nature of the substituents in the N-aryl moiety and indicates that the luminescent mechanochromic properties of these emitters can be tailored through systematic modification of the bidentate ADC ligands.

It is worth mentioning that several different types of luminescent mechanochromic transition metal complexes have been previously reported.\textsuperscript{94–96} Luminescent mechanochro-
mism of most systems described is associated with a reversible switching between crystalline polymorphic modifications and the amorphous state, primarily through the intermolecular or intramolecular metal–metal interactions.\textsuperscript{97–100} This makes systematic fine-tuning and modifications of the mechanochromic properties including the emission characteristics of different mechanochromic forms challenging. In the cases of 48–53, the mechanochromic properties do not depend on their supramolecular structures supported by intermolecular interactions but are controlled by steric effects of the carbene ligands. This can be used as an alternative and additional tool for tuning the mechanochromic properties of organometallic emitters. The multiple stimuli responsive properties of 48–53 can be useful for smart materials in developing multifunctional films with potential applications as security storage materials.\textsuperscript{102}

### 3.2. Group 10: platinum(II) ADC complexes

Platinum as well as iridium has a high spin–orbit coupling constant. In contrast to d\textsuperscript{4} Ir\textsuperscript{II} complexes which prefer strongly an octahedral coordination geometry, the Pt\textsuperscript{II} ion has an intrinsic propensity to adopt a square-planar coordination geometry, even in the presence of weak field ligands.\textsuperscript{20,103,104} This structural bias is of key importance when considering radiationless deactivation pathways of their excited states. Population of the strongly antibonding \textit{d}_{\pi-\gamma} orbital, e.g. by absorption of light or by thermal activation, weakens the Pt-ligand bonds and induces structural distortion towards a tetrahedral geometry. Such structural deformation decreases the ligand-field splitting, thus facilitating non-radiative deactivation \textit{via} thermally excited Pt \textit{d} states. The way to circumvent this is to use rigid multidentate ligand scaffolds and/or strong \textit{σ}-donor atoms that are beneficial for the development of highly robust phosphorescent Pt\textsuperscript{II} complexes.

Pt\textsuperscript{II} complexes featuring Chugaev-type\textsuperscript{67,71–76,105–107} chelating bis-ACD ligands 54 and 55 were assembled \textit{via} the template reaction of ‘BuNC, excess hydrazine hydrate, and aqueous K\textsubscript{2}[PtCl\textsubscript{4}]; separation was achieved by fractional recrystallisation (Scheme 7, Route A).\textsuperscript{108} Through this reaction, two coordinated ‘BuNCS reacted with hydrazine to yield a deprotochelating dicyanamide ligand in 54 and 55. In the case of 54, the subsequent Pt\textsuperscript{II}-mediated dealkylation of ‘BuNC gives rise to the cyanide. Reversible protonation of 54 and 55 with aqueous HBF\textsubscript{4} generates 57 and 58 (Scheme 7, Route C).

The UV–vis spectra of 54 and 55 display a moderate absorption band at 390 nm and 406 nm, respectively, which was assigned to the MLCT transition \textit{Pt} → \pi*{ADC); protonation results in a significant blue shift for the lowest energy absorption in the UV–vis spectra (Fig. 12). The luminescence of 54 and 55 in solution is extremely weak, however, the protonation brings about a dramatic emission enhancement (ca. 12-fold for 54) accompanied by the hypochromatic shift (Fig. 12). High-energy emissions of 57 and 58 formed upon protonation of 54 and 55 are assigned to the triplet intraligand excited state of the bis-ACD moiety.

It is worth noting that homoleptic tetrakis(carbene) complexes with monodentate ADC \([\text{Pt}(\text{C(NHMe)}\textsubscript{2})\textsubscript{2}]\textsubscript{2}\text{Cl}\textsubscript{2}\textsuperscript{109–111} and NH\textsubscript{3}\textsuperscript{112} \([\text{Pt}(\text{NHMe})\textsubscript{2}]\textsubscript{2}[(\text{PF}_{6})\textsubscript{2} \text{CF}_{3} \text{SO}_{3} \text{CF}_{2}]\textsubscript{2} \text{PF}_{6} \text{CF}_{3} \text{SO}_{3} \text{CF}_{2}]\textsubscript{2} \text{PF}_{6} \text{CF}_{3} \text{SO}_{3} \text{CF}_{2}]\textsubscript{2}\) (NHC = imidazolidin-2-ylidene,\textsuperscript{112} tetrahydropyrimidine-2-ylidene,\textsuperscript{112} benzoimidazol-2-ylidene\textsuperscript{113}) ligands are not luminescent, while \textit{tetr}a(carbene) complexes with bis (NHC) \(C,C\)-chelating ligands\textsuperscript{114–116} emit in the visible and near-UV regions of the spectra. The main difference between the luminescent and non-luminescent platinum complexes discussed above is the existence of a \(d_{\pi} - P_{\pi}\) conjugated system in the case of complexes with \(C,C\)-chelating carbene ligands and the absence of conjugation in the case of complexes with monodentate carbene ligands due to the possibility of rotation around the metal–carbon bond.

A unique Pt\textsubscript{6} macrocycle 56 with chelating ADC and bridging CN ligands was prepared by a template reaction between...
K₂PtCl₄, ‘BuNC, and hydrazine hydrate in water under reflux (Scheme 7, Route B).⁵⁷ Compound 56 forms through the reaction of two coordinated ‘BuNC ligands with hydrazine to give a chelating dicarbene ligand followed by the Pt⁴⁺-mediated dealkylation of ‘BuNC to CN species. In comparison with 54 and 55, stronger heating of the reaction mixture leads to the complete dealkylation of the isocyanide.

The UV–vis spectrum of 56 contains an intense absorption band at ca. 350 nm assigned to the Pt–π*(ADC) spin-allowed charge transfer transition (Fig. 13). Upon excitation at 350 nm in solution and in the solid state, complex 56 emits at ca. 500 nm.

Luminescent Pt⁺⁻ADCs 61–67 were prepared by nucleophilic addition of hydrazine or amines to the coordinated isocyanide in starting from 59 and 60 (Scheme 8).⁶¹ Conversion of isocyanide ligands in 59 and 60 to the corresponding ADCs in 61–67 results in a significant bathochromic shift of emission maxima (λₘₐₓ ca. 530 nm for 59 and 60), and 61–67 display emission in solution with λₘₐₓ in the range 540–560 nm, that was assigned to the triplet MLCT excited states.

Another family of phospharescent cyclometallated Pt⁺⁻acetyl diaminocarbene complexes 70–73 was prepared by nucleophilic attack by the primary amines on the coordinated isocyanide in 68 and 69 (Scheme 9).¹¹⁸

ADC derivatives 70–73 exhibit characteristic absorption bands at ca. 400 (70, 71) and 440 nm (72, 73) assigned to the transitions into singlet mixed (IL/MLCT) excited states (Fig. 14). Very weak lowest-energy bands (at ca. 430 or 460 nm) were only discernible in very concentrated solution and were attributed to the direct population of mixed triplet IL/MLCT nature favoured by the high spin–orbit coupling of the metal centre. In the solid state and immobilised in the PS matrix (5 wt% of the Pt⁺⁻complex) 70–73 exhibit green (70 and 71, λₘₐₓ 470–475 nm) or orange (72 and 73, λₘₐₓ 550–590 nm) emission with high or moderate Φ values (0.70–0.74 for 70 and 71, and 0.15–0.19 for 72 and 73). Conversion of 68 and 69 to 70–73 is therefore not accompanied by a shift of emission energy, but a notable increase of Φ values (Φ 0.25 for 68 and 0.08 for 69). This observation can be justified by a stronger donicity of the ADC causing higher ligand field splitting and reducing the thermal population of non-radiative triplet metal-centred excited states. The emission energies in THF solutions are essentially identical, although, as expected, with lower lifetimes and lower quantum yields (Φ < 0.1) than in PS films and in the solid state. Compounds 70–73 bearing ADCs are shown to be valuable organometallic emitters for OLED devices alongside commonly used [Pt(C^N)Cl(NHC)] counterparts with NHC ligands.¹¹⁹

Bis-alkynyl species 76 and 77 with ADC ligands were prepared by nucleophilic addition of dimethylamine to the coordinated isocyanide in 74 and 75 (Scheme 10).¹²⁰ Despite a
large excess of diethylamine and a long time (3 days), addition to a solely coordinated isocyanide occurs. The authors associate the lack of reactivity of the second isocyanide to steric restrictions, which prevent the formation of bis-ADCs.

UV–Vis spectra of 76 and 77 contain two high-energy absorption resonances with maxima in the 252–313 region, assigned to transitions involving the intraligand π–π* excitation of the alkylnyl species and charge transfer in the Pt–C=CPh. The degree of conjugation within the complex is reduced upon generation of ADC in 76 and 77, leading to a minor decrease in the extinction coefficient for the most intense UV band and a significant blue-shift of the lower-energy band (Fig. 15). In solution, 74–77 are weakly luminescent; a significant enhancement is observed for 74–77 as frozen solvent glass (77 K) or in PMMA films. Both isocyanide and ADC species showed identical profiles of phosphorescence with a λmax of 430 nm, indicating that both aforementioned ligands in 74–77 have a cooperative effect on the energy of the emissive triplet states.

Conversion of metal-bound isocyanide to ADC results in a remarkable increase in Φ from 0.014 (for 74, PMMA) and 0.058 (for 75, PMMA) up to 0.23 (for 76, PMMA) and 0.15 (for 77, PMMA). The phosphorescence lifetime is longer for both ADC derivatives 76 and 77 than for the starting bis-isocyanides 74 and 75; the enhanced quantum efficiencies of the ADC complexes are attributed to the large destabilisation of the empty Pt-centred orbital by the strong σ-donation of ADC.

The photoluminescence properties of ADC derivatives 76 and 77 in PMMA are comparable to those of platinum(n) acetylide complexes with NHC ligands, viz. trans-[(NHC)2Pt(C≡CPh)], (NHC = 1,3-substituted-1H-imidazol-2(3H)-yl) or 1,3-substituted-1H-benzimidazol-2(3H)-yl), that have shown promise in the manufacturing of OLED devices.37,121

3.3. Group 7: rhenium(i) ADC complexes

Although the ADC complexes of both Mn and Re have been previously reported, a sole report of luminescent Re ADC complexes is disclosed.122 A series of luminescent Re complexes 81–88 with C,N-ADC was synthesised via the nucleophilic addition of amino-azaheterocycles, namely, 2-aminopyridine, 2-aminothiazole, and 2-aminobenzothiazole, to Re isocyanides. The reaction is facilitated by the halide abstraction with silver or thallium salts, and/or thermal or photochemical CO ligand substitution (Scheme 11). The UV–vis spectra of 81–88 show intense ligand centred π–π* transitions of the isocyanide and C,N-ADC ligands in the UV region (λ < 330 nm) and the lower-energy bands (350 nm > λ > 450 nm) assigned to the MLCT transitions dπ(Re) → π*(C,N-ADC) and dπ(Re) → π* (CNR) (Fig. 16). Upon excitation at a λext of 350 nm, 81–88 exhibit strong emission with the energy spanning from blue to red regions. Using DFT calculations, emissions of 81, 82, 85, and 87 were attributed to the mixed triplet MLCT and LC excited-state origin, while for 83, 84, 86, and 88 to the triplet MLCT [dπ(Re) → π*(C,N-ADC)]. Within the 81–88 family the emission energy is redshifted in-line with a decrease of π-conjugation of the C,N-ADC ligand (λmax 467 nm (81),

Fig. 15 UV–Vis spectra of 76 and 77 in CH2Cl2 solution; emission spectra in different media at variable temperature. λext 310 nm. Adapted with permission from ref. 120. Copyright (2020) John Wiley & Sons, Inc.

Scheme 11 Synthesis of ADC complexes 81–88.122

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Review
3.4. Group 11: gold(I/III) ADC complexes

Gold(I) complexes usually require the presence of chromophores or a group promoting the formation of dimeric or polymeric species through aurophilic interactions to impart the desired luminescence, with luminescence lifetimes depending on the origin of the emissions.\textsuperscript{130–132}

**ADC derivatives 90 and 91** were prepared by reacting H[AuCl\textsubscript{4}] or H[AuBr\textsubscript{4}] with methyl isocyano and methylamine in an aqueous solution (Scheme 13).\textsuperscript{133,134} In solution and in the solid state, protons of two ADC species interact with X\(^-\) via N–H⋯X–⋯H–N (X = Cl, Br) three-centred HBs, which prevent the X\(^-\) coordination to Au (Fig. 17). Addition of ammonia salts (NBu\textsubscript{4}BF\textsubscript{4}, NH\textsubscript{4}PF\textsubscript{6}, NH\textsubscript{4}AsF\textsubscript{6}, or NH\textsubscript{4}SbF\textsubscript{6}) to a solution of 90 results in counter-ion metathesis and precipitation of colourless 92–95 (anion = BF\textsubscript{4}, PF\textsubscript{6}, AsF\textsubscript{6}, SbF\textsubscript{6}) and (93, 94) × \(\frac{1}{3}\) (solvent) (solvent = benzene, acetone, chlorobenzene).\textsuperscript{133,135,136}

**Complexes 90–95** are not emissive in solution at RT.\textsuperscript{135} In frozen solutions and in the solid state, 92–95 show bright phosphorescence in the violet-green region (390–535 nm) associated with the formation of dimers (90, 91) or extended chains (92–95) supported by Au⋯Au attractions (Fig. 17 and 18).\textsuperscript{134–136} Although colourless crystals of 90 and 91 are isomorphous, they exhibit different emission maxima, \(\lambda_{\text{max}}\) 391 nm (90) vs. \(\lambda_{\text{max}}\) 412 nm (91), resulting from the larger distortion in the excited state for the bromide derivative due to the larger anion size. The crystal packing in both 93 and 94 is similar, and the anion interconnects with N–H groups of three ADC three organometallic cations via HBs facilitating the further self-association by aurophilic interactions (Fig. 18). The absence of luminescence in solutions of 90–95 is due to a breakdown of the supramolecular structure. The fact that the emission spectra of the frozen solutions strongly resemble the spectra of crystalline 90–95 suggest similarities in the respective supramolecular structures in both states.

![Scheme 13](image)

**Scheme 13** Synthesis of complexes 90 and 91.\textsuperscript{133,134}

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Fig. 16 (Top): UV–vis spectra of Re\textsuperscript{4+} ADC complexes 81–88 in CH\textsubscript{2}Cl\textsubscript{2} at 298 K; (bottom): emission spectra of 81–88 in CH\textsubscript{2}Cl\textsubscript{2} at 298 K. Adapted with permission from ref. 122 Copyright (2016) American Chemical Society.

489 nm (87), 499 nm (85), 570 nm (88), and 590 nm (86)). The authors justify this trend by the contribution of the empty \(\pi(C_{\text{ADC}})\), which becomes lower-lying in the presence of a less \(\pi\)-conjugated N-heterocycle due to the interaction with a lower-lying lone pair electrons in the LUMO \(\pi(C,N\text{-ADC})\).

Alkylation of 84 with Me\textsubscript{2}SO\textsubscript{4} gives 89 (Scheme 12),\textsuperscript{122} showing a noticeable redshift of the emission on going from \(C,N\text{-ADC} 84 (\lambda_{\text{max}} 547 \text{ nm})\) to the methylated derivative 89 (\(\lambda_{\text{max}} 672 \text{ nm} \)) is linked to the increased steric repulsion due to installation of methyl groups twisting the ADC moiety. It is noteworthy that while isocyano-(diimine) complexes [Re\{CNR\}_\text{2}(N\text{=N})\text{PF}_\text{6}\] display typically orange to red phosphorescence,\textsuperscript{129} a careful design of the \(C,N\text{-ADC}\) ligands enables the MLCT phosphorescence covering the entire visible region satisfying the requirements for different applications.

**Scheme 12** Methylation of the chelating \(C,N\text{-ADC}\) ligand in 78.\textsuperscript{122}

![Scheme 12](image)
The supramolecular structure of 93 and 94 is responsive to an external stimulus; 94 demonstrates a solvatochromic luminescent effect (Fig. 19). The difference in photoluminescence energy is an outcome of bending of the columns of cations and disorder in the cation positioning within the column. Thus, the synergistic action of HBs and aurophilic attractions allows overcoming the coulombic factors which should serve to separate the cations from each other, and the supramolecular interactions (aurophilic attractions) make compounds 92–95 emissive. These results show the significant impact of the even weakly coordinating anion or solvated molecule on the aurophilic interactions. Most likely, the differences in the N–H⋯X HBs with the anions are responsible for the variations in the solid-state structures and ultimately, the photoluminescence properties.

The reaction between [Au(CNMe)2]Cl formed in situ and 2-aminoethanol or 1,2-diaminoethane led to 96137 and 97138 (Scheme 14). The shorter Au⋯Au separation in 97 relative to 90 is facilitated by the fact that in 97 the three-centred N–H⋯Cl⋯H–N HBs connect with the NH groups of two neighbouring cations (Fig. 20). In contrast, for 90 the three-centred N–H⋯Cl⋯H–N HBs bind the NH groups of the two ADC ligands from the same cation (Fig. 17). Colourless 96 and 97 show no luminescence in solution, while solid 97 is strongly emissive at RT and at 77 K with \( \lambda_{\text{max}} \) 377 nm upon excitation with \( \lambda_{\text{ext}} \) 330 nm. The luminescence observed is attributed to the aurophilic interactions present in the solid (Fig. 20).

Changes in the backbone of the ADC species may induce dramatic changes in the emission associated with the change in the electron-richness of carbene species. Protic ADC ligands, i.e. those featuring one or two NH wingtips, possess Brønsted acidity, being able to act as donors of hydrogen bonding.62 For Au–ADC complexes, aurophilic interactions arise alongside the hydrogen bonding and their synergistic interactions make these complexes capable of luminescence. As an example, complex [Au(C(NMe2)(NHMe))2](PF6)135 bearing a sole NH proton on the ADC moiety is non-emissive in both the solution and in the solid state. Due to the absence of an effective matrix of HBs, the manifestation of Au⋯Au interactions associated with the luminescence of this class of compounds turns out to be impossible. In addition, the bisprotic gold(I)–NHC complex \([\text{Au(C3H4N2)}_2]\)Cl139 does not manifest hydrogen bonding and is therefore non-luminescent.
Switching and tuning the solid-state luminescence properties of molecular materials by modulating molecular packing through non-covalent routes is an attractive prospect.\textsuperscript{140–145} Herein, ADC ligands can uniquely play both roles enabling non-covalent binding as well as luminescence emission. Complexes 101–106 were prepared by the nucleophilic attack of primary or secondary amines or diamine on the corresponding Au\textsuperscript{I}-bound 2-pyridyl isocyanides in 98–100 (Scheme 15).\textsuperscript{146} Single-crystal XRD studies of 101 and 103 indicated the formation of supramolecular 1D polymers or dimers supported by Au⋯Au aurophilic interactions and N–H⋯Cl hydrogen bonds, while 104 and 105 do not exhibit aurophilic attractions in the solid state.

A similar strategy enabled the preparation of 109–112 starting from 107 and 108 bearing 4-pyridyl isocyanide (Scheme 16).\textsuperscript{147} 109–112 lack aurophilic contacts in the solid state. The absorption spectra of 101–106 and 109–112 are very similar and contain one or two bands (at ca. 250 nm for 109–112; 250 and 280 nm for 101–106, respectively), assigned to the ligand-centred π–π* transitions associated with aromatic groups in the ligand environment. Parent Au\textsuperscript{I}-CNRs 98–100, 107, and 108 are not luminescent either in the solid state or in solution, but all ADC derivatives 101–106 and 109–112 show intense yellow-green luminescence both in solution and in the solid state at RT. The emission profiles and energies of 104, 102–106, and 109–112 in solution and in the solid state are similar and are consistent with an intramolecular origin of the luminescence owing to the absence of Au⋯Au short contacts.

For 101 and 103, the emission energies in the solid state (λ\textsubscript{max} 562 nm, 101 and λ\textsubscript{max} 470 nm, 103) are redshifted in comparison to those recorded in CH\textsubscript{2}Cl\textsubscript{2} solution (λ\textsubscript{max} 457 nm, 101 and λ\textsubscript{max} 409 nm, 103), attributed to the intermolecular aurophilic interactions. A difference in the photophysical behaviour of isocyanide and ADC derivatives can be related to a higher donor capacity of ADC ligands when compared to the parent isocyanide.\textsuperscript{148,149} As a result, ADC introduces a more significant perturbation in gold(I) orbitals by enhancing the π back-bonding contribution from gold to ADC, linked to higher intersystem crossing and phosphorescence.

Trinuclear gold(I) aminocarbene and pyrazolate complexes that possess intramolecular aurophilic contacts are capable of displaying interesting luminescence associated with different extents of intermolecular aurophilic interactions.\textsuperscript{130} Thus, the deprotonation of 103 with KOH in MeOH generates formamidinyl intermediates spontaneously trimerising to cluster 113 (Scheme 17).\textsuperscript{27} Both in solution and in the solid state, 113 possesses a trigonal prismatic arrangement of six metal atoms stabilised by the Au⋯Au interactions between the two {Au\textsubscript{3}} triangles. 113 displays an intense luminescence both in the solid state with a λ\textsubscript{max} of 523 nm and in solution with a λ\textsubscript{max} of 563 nm. The long lifetime observed (84 µs) supports the phosphorescent nature of the emission that can be attributed to the presence of Au⋯Au aurophilic interactions, as reported for other trinuclear gold complexes.\textsuperscript{27,150–155}

116 and 117 with the perylene substituents at the periphery were obtained by the nucleophilic attack of NHEt\textsubscript{2} on the Au\textsuperscript{I}-
bound perylene isocyanide in 114 and 115 (Scheme 18).\textsuperscript{156} The UV-Vis absorption profiles of 114–117 are similar and dominated by the $\pi$-$\pi^*$ transition inside of the perylene moiety. The absorption maxima of 116 and 117 at 450 nm are blueshifted ($\Delta = 990$ cm$^{-1}$) from the parent isocyanides 114 and 115 and redshifted ($\Delta = 550$ cm$^{-1}$) from free perylene. ADCs 116 and 117, as well as the starting isocyanides 114 and 115, and free perylene isocyanide exhibit strong fluorescence in solution with a well-defined vibrational structure of the emission band (Fig. 21).

The emission energies of ADCs 116 and 117 ($\Phi = 0.69$–0.93) are blueshifted ($\Delta = 830$ cm$^{-1}$) relative to the parent isocyanide species 114 and 115 ($\Phi = 0.71$–0.88) and redshifted ($\Delta = 1150$ cm$^{-1}$) relative to free perylene in-line with absorption. The resemblance of the luminescence spectra in both the vibrational structure and the spectral position suggests an intraligand $\pi$-$\pi^*$ transition where the ligand orbitals have been marginally modified by heavy metals.

The ADC derivatives 122–125 containing perylene diimide moieties were obtained via the nucleophilic attack of dimethylamine on the Au'-coordinated perylene diimide-isocyanide in...
The reaction of 121 with ethylenediamine leads to the hexanuclear derivative 126 (Scheme 19). The presence of a bulky perylene diimide moiety defines the emission properties of 118–126. ADCs 122–126 demonstrate smaller quantum yields (Φ 0.10–0.21) in comparison to those of parent 118–120 (Φ 0.50–0.67), related to a photoinduced electron transfer effect involving the ADC moiety connected to perylene diimide. This result is consistent with a strong influence of the {Au−ADC} fragment on the electronic density on the perylene diimide moiety, which significantly affects the photo-physical properties of the complexes with charge transfer from metal to ligand (Fig. 22).

In sharp contrast to AuIII complexes, very few luminescent ADC complexes of AuI have been reported. This fact also contrasts with the luminescent examples found for the isoelectronic PtII complexes.2,20,159 Low energy d–d ligand field states, which would be reached by electrons instead of the emissive excited state, and the electrophilicity of AuIII have been claimed as the possible origin of this fact. The introduction of a strong σ-donating ADC ligand increases the electron density at the metal centre and increases the energy of the d–d state.160

Gold(III)−ADC derivatives 128–130 featuring a bidentate cyclometallated C,C′-ADC ligand are obtained by reaction of [Au(C,N-pap)Cl2] (pap = 2-(2-pyridylamino)phenyl) 127 with isocyanides (Scheme 20).161

Though the starting 127 is not emissive, its ADC derivatives 128–130 are weakly luminescent (λmax 525–508 nm) in the solid state at RT. The authors suggest that the generation of ADC increases the d–d ligand field states, avoiding the non-radiative deactivation of the 3IL excited state in 128–130 as a reason for the observed difference in the properties of the starting gold-CNR and gold-ADC product.161 Lowering the temperature to 77 K induces a bathochromic shift of the emission energy of 128–130 (λmax 540–500 nm) with the lifetime in the μs range. The authors described this emission as phosphorescence, although it is better defined as thermally activated delayed fluorescence (Fig. 23).162–164

4. Conclusions and outlook

In this review, we summarised experimental studies to date disclosing the photoluminescence properties of metal complexes with acyclic diaminocarbene ligands. We have illustrated that the intrinsic σ-donating properties of acyclic diaminocarbene ligands may be connected to an efficient destabilisation of the unoccupied d orbitals and their corresponding triplet MC states, improving the efficiency and stability of phosphorescence.

From this perspective, cyclometallated IrIII complexes with ancillary ADC ligands are fully coherent and support the idea that the introduction of ADC ligands leads to molecules capable of efficient luminescence. The main trend in the design of luminescent IrIII complexes is focused on the combination of properties of cyclometallating ligands with those of acyclic diaminocarbenes. The presence of cyclometallating species in the coordination sphere of the Ir-center ensures the strongest bonding interaction with transition metals, leading to a lower extent of non-emissive quenching and at the same time opening the possibility for the tuning of emission wavelengths. To overcome the problem with the thermal population of non-emissive higher-lying triplet metal-centred ligand-field states, restricting efficient blue luminescence, one of the solutions that emerged relies on the use of a secondary set of σ-donor ligands, such as acyclic diaminocarbenes. Adding ADCs to the ligands set in the coordination sphere of metal is shown to additionally destabilise the unoccupied d-orbitals and their
corresponding metal-centred ligand-field states, improving the efficiency and stability of phosphorescence. At the same time, variable steric and donor properties of acyclic dianinocarbenes associated with their synthetic accessibility provided an additional tool for fine-tuning of emission profiles.

On the other hand, luminescence of gold(i) complexes is usually linked to the presence of chromophores or a group promoting the formation of dimeric or polymeric species through non-covalent interactions with luminescence lifetimes and efficiencies depending on the origin of the emissions. Herein, strong intermolecular interactions and substituents with suitable steric hindrance, such as ADCs, may be required to produce an ideal distance between two conjugated molecules, resulting in efficient luminescent excimers. In this case, ADC ligands featuring NH groups were shown to act as donors of hydrogen bonding for a synergistic combination of weak interactions, i.e., hydrogen bonding and metallophilic interactions in the solid-state providing an efficient tool for the tuning of photophysical properties. Examples, based on other metals, viz. Pt and Re, are not yet sufficient to demonstrate general patterns, and further studies are expected to correct the status quo.

Finally, one of the advantages of the acyclic dianinocarbene ligands already evaluated for tuning the photophysical properties of organometallic emitters is the excellent synthetic availability and respective chemical stability of their metal complexes. Taking these into consideration, an expansion of the known application fields and emergence of new application fields can be expected, including but not limited to the development of chemosensing applications, and use in data protection, visible light photocatalysis, and clinical photodynamic therapy. Those may be enabled by the use of previously unexplored metal centres, and stabilisation of complexes featuring known metals in less common oxidation states, i.e. gold(n), platinum(n), and others.

Conflicts of interest

There are no conflicts to declare.

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