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# A new ligand skeleton for imaging applications with d-f complexes: combined lifetime imaging and high relaxivity in an Ir/Gd dyad $\dagger$ 

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#### Abstract

A new rigid and conjugated ligand structure connecting phenanthroline and poly(amino-carboxylate) binding sites provides d-f complexes which show high potential for use in dual (luminescence + magnetic resonance) imaging and for optimisation of $d$ - f photoinduced energy-transfer.


Phosphorescent metal complexes offer major advantages over conventional fluorescent organic molecules as the basis of luminescent probes for cell imaging. ${ }^{1-3}$ The long luminescence lifetimes associated with triplet emission from complexes of e.g. $\operatorname{Pt}(\mathrm{II}),{ }^{2 \mathrm{a}}$ $\operatorname{Re}(\mathrm{I}),{ }^{2 \mathrm{~b}} \operatorname{Ir}(\mathrm{III}),{ }^{2 \mathrm{c}} \mathrm{Ru}(\mathrm{II}),{ }^{2 \mathrm{~d}}$ and lanthanides, ${ }^{2 \mathrm{e}}$ allow simple rejection of short-lived background autofluorescence which might otherwise interfere. In addition, variations in luminescence lifetimes of such complexes (or 'probes') in different cellular regions, caused by the presence of different analytes such as $\mathrm{O}_{2}$, provide the basis of the recently-developed microsecond-scale lifetime mapping techniques phosphorescence lifetime imaging (PLIM) ${ }^{2 \mathrm{~d}, 3}$ and time-resolved emission microscopy (TREM). ${ }^{2 \mathrm{a}, 3 \mathrm{~b}, \mathrm{c}}$

In addition, the use of highly paramagnetic complexes - often of $\mathrm{Gd}(\mathrm{III})$ - for magnetic resonance imaging (MRI) is now well established. ${ }^{4}$ Compared to luminescence-based imaging, MRI is quite complementary. Confocal microscopy offers excellent sensitivity and spatial resolution (particularly when two-photon excitation is used), but is limited in terms of tissue penetration; it is excellent for providing cellular-level detail. In contrast, MRI is capable of imaging whole bodies but with much lower spatial resolution. The combination of MRI with luminescence imaging methods using a single molecule is appealing as such a probe

[^0]would combine the broad scope of MRI with the fine detail allowed by luminescence imaging. ${ }^{5}$

This possibility has stimulated interest in a range of heteronuclear d-f complexes in which one or more phosphorescent d-block units is connected to one or more stable Gd (III) units. Notable recent examples have come from the groups of Faulkner ${ }^{6}$ and Parac-Vogt ${ }^{7}$ amongst others. ${ }^{5,8}$ A common feature of these is that the $\mathrm{Gd}($ (iII) unit is coordinated by a saturated poly-amino/ carboxylate ligand of the 'DTPA' or 'DOTA' types as these provide the necessary high kinetic and thermodynamic stability in aqueous media. A disadvantage of these however is that the saturated skeletons can permit free rotation of the $\operatorname{Gd}$ (III) unit independently of the rest of the molecule, which limits relaxivity: high relaxivities arise from slow molecular tumbling in solution which gives long rotational correlation times, and many synthetic strategies have been employed specifically to rigidify $\operatorname{Gd}($ III $)$ complexes to increase their relaxivity. ${ }^{4,5}$

We report here a new ligand architecture (Fig. 1), which allows a strongly phosphorescent $\operatorname{Ir}($ III $)$ unit to be connected to a water-stable $\mathrm{Gd}($ III ) unit via a fully conjugated and rigid connector.




Fig. 1 Structural formulae of the complexes Ir Ln and of the starting materials A and B (see ESI, $\dagger$ for full synthetic scheme).

This results in both (i) long-lived luminescence which can be used in PLIM imaging under one-photon or two-photon excitation, and (ii) unusually long relaxivity from a single $\mathrm{Gd}(\mathrm{III})$ centre as a consequence of the rigid design. The combination of $\operatorname{Ir}$ (III) and $\operatorname{Gd}($ III $)$ components for dual-imaging purposes has been very little explor$\mathrm{ed}^{8 \mathrm{c}}$ and this report is the first demonstration of PLIM using a complex that also has high relaxivity for MRI purposes. As an additional benefit, the same ligand architecture provides an effective through-bond coupling pathway for efficient Dexter $\operatorname{Ir}($ III) -Eu (iiI) energy-transfer (EnT) in the isostructural Ir Eu complex. Dualluminescent d-f complexes are of interest for a range of applications from imaging ${ }^{9}$ to white-light emission ${ }^{10}$ and many of these applications hinge on the extent of $\mathrm{d}-\mathrm{f}$ EnT which controls the balance of luminescence output from the two components. ${ }^{11,12}$ We prepared Ir Eu as an adjunct to Ir Gd to allow measurement of the q value around the $\operatorname{Ln}$ (III) centre, but its properties arising from the ligand structure are of significant interest in their own right.

The complexes Ir Ln [where $\mathrm{Ln}=\mathrm{Gd}($ III $)$ and Eu (III) respectively] are shown in Fig. 1. The $\operatorname{Ir}($ III $)$ unit is one of the well-known $\left\{\operatorname{Ir}\left(\mathrm{F}_{2} \mathrm{phpy}\right)_{2}(\mathrm{NN})\right\}^{+}$units based on cyclometallating fluorinated phenyl-pyridine ligands. ${ }^{13}$ The $\operatorname{Gd}($ III $)$ coordination is provided by a heptadentate pyridine-2,6-bis(amino-diacetate) chelating unit, ${ }^{14}$ connected to the $\left\{\operatorname{Ir}\left(\mathrm{F}_{2} \mathrm{ppy}\right)_{2}(\mathrm{phen})\right\}^{+}$chromophore via an alkynyl linkage, providing the rigid, fully conjugated pathway containing no $\mathrm{sp}^{3}$-hybridised atoms. The key step is a Sonogashira coupling reaction between compounds A (the 4-bromopyridine with two pendant, protected, amino-diacetate arms) and 3-ethynyl-1,10phenanthroline (B). After assembling the ligand skeleton, coordination of the phen to an $\left\{\operatorname{Ir}\left(\mathrm{F}_{2} \mathrm{ppy}\right) 2\right\}^{+}$unit, unmasking of the amino/ carboxylate binding site by removal of the esters, and finally incorporation of $\operatorname{Ln}($ III $)$, all used standard methods (see ESI $\dagger$ ); the final products Ir Ln were purified by HPLC and characterised by mass spectrometry and elemental analysis.

The UV/Vis absorption spectra of Ir A [the free tetracarboxylic acid complex with no $\mathrm{Gd}(\mathrm{III})$ ] and Ir Ln show the usual intense absorptions in the visible region associated with ligand-centred p p* transitions (see Fig. S14, ESI $\dagger$ ). In addition the weak shoulder and long tail between 400 nm and 550 nm is ascribed to the $\operatorname{Ir}(\mathrm{III})$ phen MLCT transition. ${ }^{13}$ The luminescence of Ir A at 530 nm , and Ir Gd at 560 nm , are broad and featureless, indicative of ${ }^{3}$ MLCT luminescence (Fig. 2a): the red-shift in Ir Gd may be ascribed to the eff ect of the $\operatorname{Gd}($ III $)$ ion whose positive charge stabilises the LUMO of the conjugated phen/alkyne/pyridyl ligand. Assignment of the luminescence as ${ }^{3}$ MLCT is supported by the substantial rigidichromism: at 77 K ( $\mathrm{MeOH} / \mathrm{EtOH}$ glass) the highest-energy feature in the luminescence spectrum of $\operatorname{Ir} \mathrm{Gd}$ (which now shows a clear sequence of vibronic components, Fig. 2a) is blue-shifted from 560 nm to 495 nm , giving an energy of $20200 \mathrm{~cm}^{1}$ for the $\operatorname{Ir}$ (III)based ${ }^{3}$ MLCT excited state. In aqueous solution at RT the $\operatorname{Ir}$ (III)based emission of $\operatorname{Ir} \operatorname{Gd}(\mathrm{f}=4 \%)$ shows two decay components with lifetimes of $\mathrm{t}_{1}: 1100 \mathrm{~ns}(56 \%)$ and $\mathrm{t}_{2}: 450 \mathrm{~ns}(44 \%)$. The presence of two components is a common consequence of aggregation in solution, ${ }^{12 \mathrm{a}, \mathrm{b}}$ possibly associated with the hydro-phobic $\left\{\operatorname{Ir}\left(\mathrm{F}_{2}-\right.\right.$ phpy)(phen) $\}^{+}$units.

The luminescence properties of Ir Eu are also of interest. The ${ }^{3}$ MLCT excited-state energy of the $\operatorname{Ir}(\mathrm{III})$-component at $20200 \mathrm{~cm}{ }^{1}$


Fig. 2 Luminescence spectra in $\mathrm{MeOH} / \mathrm{EtOH}$ (1:4) of (a) Ir Gd, in fluid solution at RT (black) and as a frozen glass at 77 K (red); and (b) Ir Eu in fluid solution at RT (purple) and as a frozen glass at 77 K (green). lex $=400$ nm in all cases.
is sufficient to allow sensitisation of the $\mathrm{Eu}(\text { III })^{5} \mathrm{D}_{0}$ state which lies at ca. $17500 \mathrm{~cm}{ }^{1}$; at RT a gradient for EnT between donor (Ir) and acceptor (Eu) of ca. $2000 \mathrm{~cm}^{1}$ is required. ${ }^{15}$ The luminescence spectrum of Ir Eu in solution (Fig. 2b) shows how partial $\operatorname{Ir}(\mathrm{III})$ Eu (III) EnT has occurred, with the $\operatorname{Ir}($ (III)-based luminescence reduced in intensity by $22 \%$ compared to what was observed for Ir Gd, and five sharp luminescence lines at $580,590,615,687$ and 700 nm from the $\mathrm{Eu}($ III $){ }^{5} \mathrm{D}_{0}-{ }^{7} \mathrm{D}_{\mathrm{n}}$ transitions superimposed on the low-energy tail of the $\operatorname{Ir}($ (III)-based luminescence making it appear red (Fig. S13, ESI $\dagger$ ). At 77 K the two emission components are more clearly separated because of the rigidochromic blue-shift of the $\operatorname{Ir}(\mathrm{III})$-based emission component (Fig. 2b).

The $\operatorname{Ir}$ (III) - Eu (III) EnT reduces the $\operatorname{Ir(III)-based~luminescence~}$ lifetime (compared to Ir Gd) to $\mathrm{t}_{1}=780$ and $\mathrm{t}_{2}=116 \mathrm{~ns}$ (again, we see two components). If we make the reasonable assumption that $\operatorname{Ir}($ III $)-\mathrm{Eu}($ III) EnT provides the best additional deactivation pathway for $\operatorname{Ir}(\mathrm{III})$-based luminescence in Ir Eu compared to what is possible in Ir Gd, then the shortest luminescence component of 116 ns in Ir Eu is associated with intramolecular quenching by $\operatorname{Ir}$ (III) -Eu (III) EnT. This gives from eqn (1) (where $t_{u}$ is the 'unquenched' lifetime from $\operatorname{Ir} \mathrm{Gd}$ and $\mathrm{t}_{\mathrm{q}}$ is the 'quenched' lifetime from Ir Eu ) an EnT rate kEnT of ca. $610^{6} \mathrm{~s}{ }^{1}$. Significantly this is an order of magnitude faster than we observed in our previous 'rod-like' water-soluble $\operatorname{Ir}($ III $)-\mathrm{Eu}($ III $)$ dyad that was investigated for cell imaging, despite the greater Ir Eu separation. The markedly superior $\operatorname{Ir}($ III $)-\mathrm{Eu}($ III $) \mathrm{EnT}$ in Ir Eu can be ascribed to the fully conjugated pathway facilitating Dexter energy-transfer ${ }^{12}$ in this present system. All the photophysical results are summarised in Table S1 in ESI. $\dagger$

$$
\begin{equation*}
\mathrm{kenT}^{\mathrm{En}}=1 / \mathrm{t}_{\mathrm{q}} \quad 1 / \mathrm{t}_{\mathrm{u}} \tag{1}
\end{equation*}
$$

To assess the suitability of the complexes as probes for PLIM imaging, their cellular localization, emission properties and toxicity were evaluated in live MCF7 cells. Cells were incubated with Ir Ln at $25 \mathrm{mM}, 50 \mathrm{mM}$ and 100 mM for 4 and 24 hours in fully supplemented Roswell Park Memorial Institute (RPMI) media at 37 1C. Steady-state confocal microscopy (typical images in Fig. 3A), shows that Ir Gd exhibits punctate cytoplasmic staining with some accumulation in the perinuclear region, the latter being most notable at high concentrations and long incubation times.


Fig. 3 Two-photon (lex $=780 \mathrm{~nm}$ ) steady-state confocal imaging of Ir Ln complexes. Column (A), in descending order: DIC, emission and overlay images showing typical staining pattern of Ir Ln dyads in live MCF7 cells (Ir Gd, $50 \mathrm{mM}, 4 \mathrm{~h}$ ). Column (B): cellular uptake comparison after 24 hour incubation with (in descending order): RPMI media only, Ir Eu ( 100 mM ), Ir Gd (100 mM).
$\operatorname{Ir}$ (III)-based emission was observed under both one-photon (458 nm) and two-photon ( 780 nm ) excitation, consistent with the known modest two-photon absorption ability of $\operatorname{Ir}(\mathrm{III})$ complexes of this family. ${ }^{9,16}$ Optical sectioning (Fig. S21, ESI $\dagger$ ) and co-staining with the commercial nuclear stain DAPI (Fig. S22, ESI $\dagger$ ) confirm that Ir Gd was internalized into the cell cytoplasm, but did not cross the nuclear membrane.

Interestingly, Ir Gd appeared to be internalized more rapidly than the isostructural Ir Eu complex. Fig. 3B shows steady-state confocal images after 24 hours incubation at 100 mM , recorded with the same laser power and detector gain. Emission from Ir Gd incubated cells is significantly brighter than that of Ir Eu, to the extent that the detail of the staining pattern cannot be clearly distinguished (due to the high detector gain). This difference is not solely due to the inherently brighter $\operatorname{Ir}($ III $)$-based emission in $\operatorname{Ir}$ Gd. The significant eff ect Ir Gd has on the metabolic activity of MCF cells in comparison to Ir Eu suggest that considerably more $\operatorname{Ir} \mathrm{Gd}$ is taken up by the cells (see Fig. S23, ESI $\dagger$ ). The reason for this diff erence in uptake between Ir Gd and Ir Eu is not obvious but the eff ect is clear, with lower concentrations/shorter incubation times being typically preferred for Ir Gd .

Lifetime mapping of the $\operatorname{Ir}($ III $)$-based emission from both dyads was carried out using TP-PLIM (Fig. 4). In both cases, emission decays were best fit to a double exponential and only


Fig. 4 Two-photon PLIM imaging ( $\mathrm{lex}^{=}=780 \mathrm{~nm}, 12 \mathrm{~ms}$ imaging window) of Ir Eu ( $100 \mathrm{mM}, 20 \mathrm{~h}$ ) and Ir Gd ( $25 \mathrm{mM}, 20 \mathrm{~h}$ ) in live MCF7 cells. Top: intensity images, where all emitted photons are binned into one channel. Middle: t 1 lifetime maps with rainbow legend set to $0-600 \mathrm{~ns}$ for both images, showing unifomity of cellular lifetime for each compound and the difference in $\operatorname{Ir}(ו 11)$-based emission lifetime between Ir Eu and Ir Gd. Bottom: emission decay traces and lifetime dsitributions of the $\operatorname{Ir}(111)$-based emission from Ir Gd and Ir Eu in the cells.
the major component ( $486 \%$ ) $\mathrm{t}_{1}$ was used for plotting lifetime maps. The $\operatorname{Ir}($ III $)$-based luminescence lifetimes are uniform across the cells for both dyads, with the lifetime values being comparable to those observed in aerated solution. Fig. 4 also highlights the clear diff erence in $\operatorname{Ir}(\mathrm{III})$-based luminescence lifetimes between Ir Gd and Ir Eu, brought about by energy transfer, by showing both lifetime maps set to the same parameters (rainbow chart $=0-600 \mathrm{~ns}$ ). Ir Gd appears green (longer lifetime), whereas Ir Eu appears orange due a shorter lifetime. Example decay traces for each dyad were exported and overlaid for comparison (Fig. 4).

From Ir Eu we found that the $\mathrm{Eu}(i \mathrm{iII})$-based luminescence lifetimes were 0.42 ns in water and 1.14 ms in $\mathrm{D}_{2} \mathrm{O}$, giving a value for q of 1.6 $0.5,{ }^{17}$ comparable to what is observed with other $\operatorname{Gd}($ III $)$ complexes of heptadentate ligands used for MRI. ${ }^{4,5}$ Despite this, at 20 MHz and 371 C the relaxivity of Ir Gd is $11.9 \mathrm{mM}^{1} \mathrm{~s}^{1}$, measured over a range of concentrations (see ESI $\dagger$ ). This is considerably higher than that of typical mono-nuclear $\operatorname{Gd}\left(\text { (II) complexes (typically, } 4-5 \mathrm{mM}^{1} \mathrm{~s}^{1}\right)^{4,5}$ and must
be a consequence of the rigidity imposed on the complex by the conjugated linkage.

Notably, this is comparable to relaxivity values observed in other d-f hybrids which contain three or four $\mathrm{Gd}(\mathrm{III})$ centres that are individually more flexible due to the saturated ligand skeletons. ${ }^{5 \mathrm{c}}$ Thus the ligand design in Ir Gd is clearly effective at providing high relaxivity for a relatively low molecular weight complex without the need to incorporate several $\mathrm{Gd}($ (III ) centres, or to conjugate the probe to a biomolecule to slow down its rotational correlation time.

For imaging purposes with Ln (iiI)-containing complexes, kinetic stability is important due to the toxicity of free Ln (iiI) ions. The luminescence spectra of Ir Gd and Ir Eu showed no change after prolonged storage in aqueous solution: loss of the $\operatorname{Ln}($ III $)$ ion would result in each case in a blue shift of the $\operatorname{Ir}(\mathrm{III})$-based emission maximum from 564 nm to 532 nm due to the generation of free Ir A (Fig. S15, ESI $\dagger$ ). In addition, the kinetic stability of Ir Eu was measured by luminescence spectroscopy in the presence of 1 equivalent of the competing ligand DOTA [the octadentate macrocyclic ligand system cyclen-1,4,7,10-tetraacetic acid, used as a Ln (III) receptor] at a concen-tration of 0.1 mM , in both water and in PBS buff er (see Fig. S19 and S20 respectively, ESI $\dagger$ ). If the Eu(III) ion were extracted from Ir Eu by the competing DOTA ligand, we would see a steady loss of sensitised $\mathrm{Eu}(\mathrm{III})$-based luminescence as well as the blue-shift of the $\operatorname{Ir}(\mathrm{III})$-based emission component. In the presence of DOTA, the $\operatorname{Ir}($ III $)$-based emission showed no significant change in profile, and the sensitised $\mathrm{Eu}(\mathrm{III})$-based luminescence remained almost intact ( $05 \%$ decrease in intensity after 3 days), confirming the integrity of the complex even under these challenging conditions (Fig. S19, ESI $\dagger$ ). When PBS buff er was used as the medium, greater loss of Eu-based luminescence intensity was observed (Fig. S20, ESI $\dagger$ ) presumably associated with the presence of phosphate.

In conclusion, this ligand architecture off ers substantial scope for dual (luminescence + magnetic resonance) imaging using d-f complexes because of its rigidity; and for applications requiring d-f energy-transfer because of the conjugated pathway. Thus Ir Gd provides both the capacity for PLIM measurements as well as unusually high relaxivity for a mono-nuclear $\mathrm{Gd}($ (II) complex; and Ir Eu demonstrates unusually effective d-f Dexter energy-transfer.

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    $\dagger$ Electronic supplementary information (ESI) available: Full details of (i) synthesis and characterisation of all new compounds and intermediates; (ii) photophysical studies including additional images of UV/Vis and luminescence spectra, and tabulated data; (iii) imaging experiments, including cell culturing, confocal microscopy, PLIM measurements and toxicity studies; and (iv) relaxivity measurements.

