Luminogenic iridium azide complexes

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Abstract

The synthesis and characterization of luminogenic, bioorthogonal iridium probes is described. These probes exhibit long fluorescent lifetimes amenable to time-resolved applications. A simple, modular synthesis via 5-azidophenanthroline allows structural variation and allows optimization of cell labeling.

Luminogenic bioorthogonal probes have emerged as essential tools to visualize specific biomolecules in complex and confined environments, by virtue of “turn-on” emission and exquisite selectivity toward unique functional groups. Transition-metal complexes as labeling dyes have potential benefits such as red to near-IR emission, long photoluminescence lifetime, low photobleaching, stability in oxidative environments, and synthetic ease. Although they complement organic fluorogenic probes, transition-metal luminogenic probes remain relatively little-studied. In this paper, we describe the first transition-metal complex-based luminogenic azide probe appropriate for biological imaging: an iridium emitter with red photoluminescence, long emission lifetimes, efficient “turn-on” photoluminescence, and cell penetration and labeling capabilities.

Much of the development of luminogenic transition-metal complexes has focused on sensor development, while bioorthogonal probe development remains less studied. Yet metal-based emitters have tunable emission and are prepared by simple synthesis. Furthermore, transition-metal complexes can have superior performance in two-photon imaging. Octahedral metal complexes are 3D objects, less prone to aggregation, membrane association, and DNA interactions common with planar organic fluorophores. Indeed, appending polyarene units to octahedral complexes is a common method to induce DNA interactions in otherwise inert complexes. Perhaps most significantly, the intrinsic triplet excited–state of transition–metal complexes results in long photoluminescence lifetime (10 ns to 100 \( \mu \)s or longer). Together with “time-gated” detection methods, photoluminescent probes would allow independent analysis of multiple dyes with similar emission profiles and/or images with significantly lower background noise. Recently, a study described turn-on imaging with rhenium compounds by means of [4 + 2] cycloaddition reaction. A recent

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report of DNA staining with dinuclear ruthenium complexes confirms the potential of time-
resolved imaging.\textsuperscript{16} Likewise, pH-responsive iridium complexes are effective for time-
resolved imaging of the cytoplasm.\textsuperscript{17}

Our probe design was guided by our work\textsuperscript{7} indicating that the photoluminescence quantum
efficiency of an octahedral phenanthroline–iridium complex is affected by substituents at the
phenanthroline 5-position, which seem capable of modulating the contribution of non-
radiative pathways to the relaxation of a metal-to-ligand charge transfer (MLCT) excited
state.\textsuperscript{19} Similar octahedral iridium complexes are well-suited to cellular and sub-cellular
imaging.\textsuperscript{20-23} We hypothesized that an azide might serve as a similarly non-radiative
quencher for Ir(ppy)\textsubscript{2}(phen) (ppy = 2-phenylpyridine and phen = 1,10-phenanthroline).
Although the efficacy of azide-based quenching can be unpredictable even in relatively well-
studied organic fluorophores,\textsuperscript{24} we designed azide-substituted complexes 5. Three azide
complexes 5a-5c were isolated (78-93\%) by reaction of amine precursors 4 with t-butyl
nitrite and trimethylsilyl azide\textsuperscript{25} after precipitation from ether (Fig. 1a).

A more convergent and efficient preparation of the desired complexes (5) was also
developed from 5-azidophenanthroline (3), prepared for the first time here by epoxide ring-
opening and subsequent elimination\textsuperscript{1} of an epoxide precursor 1 (all attempts at diazotization
of 5-aminophenanthroline were unsuccessful). In this way, 5-azidophenanthroline (3) was
purified in high yield, and complexation with [Ir(ppy)\textsubscript{2}(MeCN)\textsubscript{2}][PF\textsubscript{6}] afforded
5a (74\%). The preparation of complex 5 from azidophenanthroline 3 is a more convergent route that
facilitates variation on the 2-phenylpyridine ligand. With an eye toward investigating the
effects of different substituents, we synthesized complexes 5b, 5c, 5d, incorporating water-
solubilizing and anionic groups.

Consistent with photoluminescence “turn-on” behavior, the azide complexes show very
weak photoluminescence, while the triazole products (6) of a cycloaddition reaction with
phenylacetylene show bright emission. Fig. 1b shows the emission spectra of the
carboxyazide complex 5b (red line) and the carboxytriazole complex 6b (blue line). The
photophysical properties of these complexes are summarized in Table 1. Both parent and
carboxylate-functionalized complexes show significant enhanced photoluminescence upon
the triazole ring formation (13\times for 6a, 19\times for 6b, relative to the azide complexes).
Gratifyingly, the amine complexes 4a and 4b also show minimal luminescence. As expected,
the triazole complex 6a has significant emission in the region of relative tissue transparency
above 650 nm (emission maximum: 637 nm).\textsuperscript{27,28} Photoluminescence lifetimes of the
triazole complexes (~60 ns) are significantly longer than that of typical organic fluorophores
(<5 ns). The triazole 6b exhibited useful quantum efficiency in aqueous solution (4.5\%), in
contrast to lower efficiencies often observed with transition-metal emitters due to a large
non-radiative rate constant. Moreover, the lifetime and brightness are greatly increased in a
non-solution application; the lifetime of a carboxytriazole complex immobilized on PVDF
membrane (vide infra) is \~1 \mu s, rendering these complexes useful for surface imaging.

The kinetics of azide-alkyne cycloadditions were straightforward and consistent with an
electron-deficient azide. We first tested copper-catalyzed reactions with a triazole ligand
using a microplate reader (Fig. 2a).\textsuperscript{26} Complete “turn-on” of the photoluminescence was
observed within 20 min (Fig. 2a, red rectangles), while no reaction occurred in the absence of copper catalyst (blue triangles). Similar trends were observed for the other azide complexes 5a-5c (ESI). A recent report suggests that the kinetics of metal-free cycloaddition reactions are governed by electronics matching of the two reactants to favor normal-demand or inverse-demand cycloaddition pathways. To further investigate reactivity of the azido Ir complex, we monitored the photoluminescence enhancement of reaction of 5d with an electron-rich cycloalkyne (BCN, 8a) and electron-poor dibenzocyclooctyne (DBCO, 9) (Fig. 2b, c). Although both of reactions shows increase of photoluminescence, the reaction of 5d with 8a proceeded considerably faster than 9; Assuming a pseudo-first-order reaction, the apparent half-life of the reaction of 8a was much shorter than that of 9 (19 min vs 200 min). These results identify possible optimal alkyne partners for labeling applications, and imply that the complexes act as electron-deficient azides in cycloaddition reactions.

Having confirmed the luminogenic properties of the iridium complexes, we examined their suitability for protein labeling. Alkyne-tagged BSA was treated with the iridium-azide complex 5b in the presence of CuSO₄, sodium ascorbate, and THPTA. After SDS-PAGE and membrane transfer, a strong photoluminescence band was observed only in the presence of a copper (II) salt and alkyne tag, consistent with selective tagging (Fig. 3b).

The long photoluminescence lifetime of the Ir complexes could be used to discriminate among emissive dyes and remove background emission. We analyzed mixtures of two dye-labeled proteins, the BSA-Ir conjugate and maltose binding protein labeled with rhodamine (MBP-TAMRA), which has an emission profile similar to BSA-Ir. A gel blot analysis of the mixture revealed two photoluminescent bands with similar steady-state intensities (Fig. 3c and ESI). However, the emission lifetimes were markedly different (~1 μs vs <100 ns, Fig. 4a). Following concepts we outlined previously, a 1000–1500-ns time window was chosen to minimize background MBP-TAMRA. Fig. 4b compares the emission intensity of the proteins under steady-state (left) and time-resolved (right) analysis. Time-gating and the unique photophysical properties of the iridium complex allow a 20-fold diminution in signal from the model background TAMRA signal. Interestingly, time-gating methods also eliminated a significant background emission from the PVDF membrane (ESI).

Finally, luminogenic iridium azides proved capable cellular imaging agents. U2OS cells were pre-treated with reactive alkyne 8b, washed, and fixed. Attempted imaging with complex 5a was disappointing, as photoluminescence was observed in negative controls (Fig. 5a). Accumulation and membrane association of hydrophobic, cationic dyes is well known, and thus we had incorporated carboxylate or sulfate groups (5b and 5d) to alter the charge. The anionic sulfate complex 5d succeeded in minimizing background staining (Fig. 5a). The observation of background non-covalent staining with 5a and 5b could be due to background photoluminescence of either the unreacted azide or (more probably) the reduced aniline compound, possibly amplified by solvochromatic behavior of hydrophilic dyes. Through chemical manipulation of the physical properties of the azide complexes, we successfully imaged alkyne-modified biopolymers while preventing background photoluminescence.
Two-photon excitation is a powerful imaging tool, and iridium complex $5d$ exhibited robust two-photon imaging. Motivated in part by the large two-photon cross-section of similar octahedral metal complexes, we irradiated alkyne-modified cells with a 860-nm laser and saw clean cell images with minimal background in the absence of alkyne (Fig. 5b). Similar to results from one-photon excitation, the best images with minimal background were obtained with anionic complex $5d$. Gratifyingly and consistent with the primary motivation for two-photon excitation in general, our initial studies indicate improved signal-to-noise with 860-nm excitation.

In summary, the facile synthesis of luminogenic azido iridium complexes enables their use for biomolecule labeling and imaging. These iridium complexes have large Stokes shifts, exhibit photoluminescence “turn-on” behaviour upon triazole formation, have long photoluminescence lifetimes, and are amenable to two-photon imaging. The use of time-gating removes unwanted signal from spurious emitters. The simple, modular synthesis allowed access to an anionic derivative that prevents background emission. Since phenanthroline is a common ligand motif for photoactive transition-metal complexes (e.g. Ru, Re, Pt), the facile synthesis of 5-azidophenanthroline described here could help expand the portfolio of functionalized photoactive complexes.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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References

Fig. 1.
a) Synthesis of compounds. i) NaN₃; then Ac₂O, 83%. ii) DBU, 83%. iii) [Ir(ppy)_2(MeCN)_2][PF₆], 74%. iv) tBuONO and TMS-N₃, 88% (5a), 78% (5b), 93% (5c). v) phenylacetylene, Cul, 42% (6a), 74% (6b). (vi) SO₃·NMe₃, 44%. b) Emission spectra of 5b (blue) and 6b (red). c) Solution of 5a (left) and 6a (right) under UV lamp.
Fig. 2.
Kinetics complex 5d cycloadditon, assessed by micro plate reader. Emission measured at 600 nm. a) reaction with pentynoic acid in the presence or absence of CuSO₄, sodium ascorbate, and a triazole ligand THPTA. b) Cu-free reaction with bicyclo[6.1.0]non-4-yne (BCN) derivative 8a and dibenzocyclooctyne (DBCO) derivative 9. c) Structure of alkynes used.
Fig. 3.
a) Labeling alkyne-tagged BSA with the carboxy complex $5b$ in PBS buffer/DMSO (95:5).
b) Analysis of reactions by photoluminescence imaging and total protein stain. c) Luminescence imaging of a mixture of BSA-Ir (alkyne-tagged BSA reacted with Ir complex $5b$) and MBP-TAMRA (fusion protein of Yes-SH3 and maltose binding protein (MBP) modified with TAMRA-NHS ester).
Fig. 4.

a) Photoluminescence time decay for the two bands in Fig. 3c on the PVDF membrane. Excitation with a picosecond 370-nm laser diode and emission collected at 570 nm. b) Comparison of emission intensity using steady-state (left) and time-resolved (1000-1500 ns, right) spectroscopy.
Fig. 5.
Labeling U2OS cells after treatment with BCN NHS-carbonate 9b. a) Single-photon excitation luminescence image after treatment with complexes 5a, 5b, or 5d. The emission observed in the absence of BCN varies among three dyes while is comparable in the presence of BCN. b) Two-photon excitation luminescence image of the cells after treatment with azide complexes 5a, 5b, and 5d. (Scale bar: 100 μm).
Table 1

Photoluminescence properties of Ir complexes.

<table>
<thead>
<tr>
<th>complex</th>
<th>$\lambda_{em}$ (nm)</th>
<th>$\epsilon^a,b$ (M$^{-1}$cm$^{-1}$)</th>
<th>$\phi^a,c$ (%)</th>
<th>$\tau^a,d$ (ns)</th>
<th>brightness$^e$</th>
<th>turn-on ratio$^f$</th>
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<tr>
<td>amine 4a</td>
<td>601</td>
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<td>7600</td>
<td>0.13</td>
<td>69</td>
<td>9.5</td>
<td>–</td>
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<tr>
<td>triazole 6a</td>
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<td>8000</td>
<td>1.50</td>
<td>62</td>
<td>119.0</td>
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</tr>
<tr>
<td>amine 4b</td>
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<td>9400</td>
<td>0.15</td>
<td>719</td>
<td>14.2</td>
<td>0.80</td>
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<tr>
<td>azide 5b</td>
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<td>0.18</td>
<td>104</td>
<td>17.0</td>
<td>–</td>
</tr>
<tr>
<td>triazole 6b</td>
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<td>4.48</td>
<td>104</td>
<td>322.6</td>
<td>19.0</td>
</tr>
<tr>
<td>BSA-Ir$^g$</td>
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<td>–</td>
<td>–</td>
<td>1151</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

$^a$measured in PBS/MeOH (4:1) for 4a-6a and in PBS/MeOH (95:5) for 4b-6b.

$^b$extinction coefficient at 370 nm.

$^c$quantum yields relative to [Ru(bpy)$_3$]Cl$_2$ reference ($\phi = 4.0\%$) in air-saturated aq soln.$^{18}$

$^d$photoluminescence lifetime.

$^e$brightness = $\epsilon \times \phi$.

$^f$Relative brightness of triazole/azide.

$^g$carboxylate complex bound to bovine serum albumin on PVDF membrane.