Somatostatin receptor-targeted organometallic iridium(III) complexes as novel theranostic agents

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A novel somatostatin receptor-targeted anticancer agent based on the conjugation of a highly cytotoxic and luminescent cyclometalated iridium(III) complex to tumor-targeting vectors based on octreotide peptide has been described, and its potential for targeted theranostic applications has been demonstrated.

During the fight against cancer, chemotherapeutic agents have to overcome many obstacles that often prevent a successful outcome of the disease. Indeed, low molecular weight cytotoxic drugs, either organic molecules (e.g. camptothecin and doxorubicin) or metal complexes (e.g. cisplatin and derivatives), cause severe toxic side effects in patients owing to their poor tumor tissue selectivity. Low tumor accumulation, poor aqueous solubility and intrinsic or acquired resistance also contribute to reducing their anticancer efficacy. In such a context, targeted delivery approaches have emerged as a promising strategy to overcome these drawbacks, particularly those based on ligands whose receptors are overexpressed on the surface of malignant cells compared with healthy cells. The conjugation of therapeutic agents to targeting vectors based on small regulatory peptides offers several advantages including the disposal of efficient solid-phase procedures for synthesizing drug conjugates with improved pharmacological properties.

Iridium complexes have recently emerged as promising alternatives to platinum-based metallo-anticancer drugs. Meanwhile, cyclometalated Ir(III) complexes have gained attention as imaging and sensing probes due to their rich photophysical properties and good cell permeability, which can be fine-tuned by the modification of the ligands. For example, the use of the pharmacophore benzimidazole as a ligand has given rise to a wide variety of metal compounds that act either as anti-angiogenic and/or anti-tumor agents, or inhibitors of amyloid-β aggregation. The integration of anticancer activities into cyclometalated Ir(III) complexes provides, therefore, an opportunity for the construction of novel theranostic platforms. Ir(III) complexes can also function as efficient photosensitizers for producing singlet oxygen, 1O2, and can even be developed as organelle-targeted PDT agents. Despite these promising achievements, further work is necessary to improve the pharmacological properties of Ir(III) metallo drugs, such as aqueous solubility and selectivity against cancer cells. In this context, targeted approaches based on peptide vectors whose receptors are overexpressed on the membrane of tumor cells compared with normal cells in combination with light activation open the door to a new generation of metallo-anticancer agents with a dual mechanism of selectivity.

Herein, we have conjugated for the first time a highly cytotoxic and luminescent Ir(III) complex, [Ir(ppz)2(N ppm)2] (Hppz = 1-phenylpyrazole; N ppm = methyl 1-butyl-2-pyridyl-benzimidazole-5-carboxylate), to tumor-targeting vectors based on octreotide peptide with the aim of increasing cancer cell selectivity and exploring their potential as novel theranostic agents (Scheme 1).

Octreotide (OCT) is a FDA-approved synthetic cyclooctapeptide agonist of the endocrine hormone somatostatin that displays high affinity for the somatostatin subtype-2 receptor (SSTR2). It is metabolically more stable than somatostatin since it incorporates α-amino acids, and the cysteine bridge stabilizes the pharmacophore sequence (Phe7-α-Try8-Lys9-Thr10) in a β-turn. As a result, octreotide binds with high affinity and selectivity to somatostatin receptors (SSTRs), mainly SSTR2. Precisely, the fact that SSTR2 is the most frequently overexpressed somatostatin receptor on the membrane of many tumor cells led to the use in the clinics of the [111In-DTPA]-octreotide and [90Y-DOTA-Tyr3]-octreotide conjugates in molecular imaging and therapy of neuroendocrine tumors, respectively, and several other SSTR2-targeted radiotherapeutics are currently under clinical evaluation. Octreotide has also been conjugated to cytotoxic organic drugs and metal complexes,
The attachment of the Ir(III) complex to octreotide was designed through the formation of an amide bond between a carboxylic function in the benzimidazole diimine ligand and the N-terminal end of the peptide sequence (Scheme 1). 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of 5 and 6 was reduced in SSTR2+ HeLa cells at 4 °C, thereby indicating internalization through an SSTR2-mediated energy-dependent endocytic mechanism. However, the accumulation of the Ir–octreotide conjugates in the SSTR2− MDA-MB-231 cells was not influenced when lowering the temperature, which could suggest the participation of other penetration mechanisms. These results are not surprising since the nature of the targeted drug and of the linker can strongly influence not only peptide–receptor binding but also tumor penetration, metabolism and excretion of the conjugate.1a

Taking into account the above-mentioned results, we carried out competitive studies with somatostatin to further confirm the involvement of SSTR2 in the internalization of the conjugates. As shown in Fig. 1C, the pre-treatment of HeLa cells with somatostatin led to a concentration-dependent reduction of the accumulation of both conjugates, which confirms the participation of SSTR2. As expected, neither Ir accumulation from complex 1 or 2 nor Pt accumulation from cisplatin was affected by the presence of somatostatin. Interestingly, accumulation in HeLa cells at different time periods revealed a significant drop in the total cellular Ir accumulation after 24 h, which suggests that the Ir compounds or their metabolites are good substrates for p-glycoprotein or some other alternative detoxification mechanisms (Fig. 1D).

Cytotoxicity studies were performed in HeLa and MDA-MB-231 cells to estimate the in vitro antitumor potential of Ir–octreotide conjugates (5 and 6) and of the parent complexes (1 and 2) and peptides (OCT and dcOCT). The photobiological activity of the compounds was also assessed via irradiation with visible light. In both cases, the MTT assay was performed after 24 h (Table 1) or 72 h (Table S4, ESI†). Not surprisingly, complex 1 displayed the highest activity under all the tested conditions,5a and visible light irradiation leads to a slight improvement of the IC50. In contrast, complex 2 did not display antiproliferative activity. These results correlate well with the accumulation data of both complexes determined by ICP-MS. The conjugation of the Ir(III) complex to the peptide moieties reduced the cytotoxic activity of 1 in both cell lines, which again correlates with the reduced accumulation. However, the activity that the conjugates retain is still reasonable for a drug–peptide conjugate, specially taking into account that their efficacy depends not only on the potency of the therapeutic cargo but also on several factors such as the number of receptors available to mediate internalization, the receptor recycling rate, the binding affinity of the peptide for its receptor when conjugated to the drug cargo or endosomal sequestration, among others.1a Overall, both conjugates showed similar antitumor activities which attest the CH2–CH2 linkage as a suitable isostere for the disulfide bond. Even so, Ir–OCT was about 2–3 times more active than Ir–dcOCT and was capable of distinguishing between HeLa and MDA-MB-231 cells after 24 h, being more active in the SSTR2-overexpressing cell line. This is particularly appealing since the parent complex 1 was more active (about 2.5-fold) in the SSTR2− MDA-MB-231 cells. Similarly to complex 1, visible light irradiation drives to a moderate improvement of the antitumor activities of the conjugates, enough however to balance the activity reduction that conjugation involves. Interestingly, the cytotoxic activity of conjugate 5 after 24 h was slightly lower in the non-malignant CHO-K1 cell line than in tumor HeLa cells (Table S4, ESI†). Octahedral Ir(III) complexes are characterized for being highly photostable, so that they are rarely used in PACT7 because of their high photostability and low phototoxicity. 5b,6b,14 Overall, the results obtained in this study confirm the potential of these compounds as photodynamic agents.

Table 1 | IC50 (μM) of the compounds tested after 2 h of incubation with the cells followed by 24 h of the recovery time in drug-free media\(^a\)

|          | Dark | Irradiated | PI\(^b\) Dark | Irradiated | PI\(^b\) |
|----------|------|------------|---------------|------------|---------|
| HeLa     |      |            |               |            |         |
| 1        | 3.13 ± 0.21 | 1.25 ± 0.11 | 2.5 | 1.23 ± 0.09 | 0.91 ± 0.08 | 1.4 |
| 2        | > 200 | > 200      | nd            | > 200      | nd      |
| 5        | 36.9 ± 2.7 | 15.5 ± 2.5 | 2.0 | 49.2 ± 4.1 | 41.9 ± 4.4 | 1.2 |
| 6        | 57.8 ± 4.9 | 33.1 ± 5.1 | 1.1 | 31.0 ± 3.3 | 36.7 ± 2.8 | 1.4 |
| OCT      | > 200 | > 200      | nd            | > 200      | nd      |
| dcOCT    | > 200 | > 200      | nd            | > 200      | nd      |

\(^a\) Results are the means ± SDs from three independent experiments. 
\(^b\) PI: phototoxic index (IC50 of non-irradiated cells/IC50 irradiated cells).
was studied by laser scanning confocal microscopy in HeLa cells with the aim of exploring their potential as targeted theranostic agents. As shown in Fig. 2 and Fig. S12 [ESI†], the emission of the Ir(III) complex allowed the visualization of luminescent vesicles in the cytoplasm, most likely endosomes, confirming the cellular uptake of the Ir–octreotide conjugates by HeLa cells. Slightly lower cellular accumulation in MDA-MB-231 cells was reflected from the reduced intensity of the luminescence signal compared to that of HeLa cells (Fig. S13, ESI†).

In summary, we have described the synthesis of novel somatostatin-targeted anticancer agents based on the conjugation of a cyclometalated luminescent Ir(III) complex to octreotide vehicles, and demonstrated their potential as targeted theranostic agents. On the one hand, Ir–octreotide conjugates accumulate in cancer cells overexpressing SSTR2, and the participation of the receptor was confirmed by competitive experiments. Such differences in accumulation between SSTR2+ and SSTR2− cancer cell lines allowed the modification of the cytotoxicity of the parent Ir complex, since the conjugates were more active in HeLa cells than in MDA-MB-231 cells, which is the opposite tendency found with 1. Notably, peptide vehicles (OCT and dcOCT) were non-cytotoxic and the cytotoxicity was increased in all cases upon visible light irradiation and ROS production was confirmed. On the other hand, the internalization of the Ir–octreotide conjugates could be easily visualized by confocal microscopy owing to the luminescence properties of the Ir(III) complex. Overall, these results open the door to the design of novel theranostic agents based on Ir–peptide conjugates with improved tumor selectivity. Future work is directed to the optimization of the compounds by improving the potency and photophysical properties of the cyclometalated Ir(III) complex through ligand modifications as well as by exploring the use of more hydrophilic or cleavable linkers to improve their pharmacological properties.

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