Original Article

Design, synthesis and in vitro characterization of fluorescent and paramagnetic CXCR4-targeted imaging agents

Ole Tietz1*, Nazila Kamaly1*, Graham Smith1, Elham Shamsaei1, Kishore K Bhakoo2, Nicholas J Long1,3, Eric O Aboagye1

1Department of Surgery and Cancer, Comprehensive Cancer Imaging Centre, Hammersmith Hospital, Imperial College London, London, W12 0NN, UK; 2Translational Molecular Imaging Group, Singapore Bioimaging Consortium, Agency for Science, Technology and Research, Singapore 138667; 3Department of Chemistry, Imperial College London, South Kensington, London, SW7 2AZ, UK. *These authors contributed equally to this work.

Received February 27, 2013; Accepted June 21, 2013; Epub July 10, 2013; Published July 15, 2013

Abstract: The G-protein coupled C-X-C chemokine receptor type 4 (CXCR4) is highly overexpressed in a range of cancers and is therefore an excellent biomarker for cancer imaging. To this end targeted iron oxide nanoparticles were developed and utilised for in vitro imaging of MDA-MB-231 breast cancer cells overexpressing the CXCR4 receptor. Nanoparticles comprising an iron oxide core, encapsulated in a stabilising epichlorohydrin crossed-linked dextran polymer, were conjugated to a cyclopentapeptide with affinity to the CXCR4 receptor. The particles were characterized for their size, surface charge and r2 relaxivity at 4.7 T. MR imaging of the CXCR4 receptor with targeted iron oxide nanoparticles revealed an approximately 3-fold increase in T2 signal enhancement of MDA-MB-231 cells compared to non-targeted controls. Prussian blue staining of labeled MDA-MB-231 cells revealed darker and more intense staining of the cellular membrane. This study demonstrates the potential of targeted iron oxide nanoparticles for the imaging of the CXCR4 receptor by magnetic resonance imaging (MRI).

Keywords: CXCR4, iron oxide nanoparticles, tumor MRI, targeted nanoparticles, T2 imaging, cyclopentapeptide

Introduction

Superparamagnetic iron oxide nanoparticles (IONPs) are used in MRI due to their biocompatible nature and strong effect on T2 relaxation. IONPs are commonly composed of a monocrystalline iron oxide core with a polysaccharide coating such as dextran [1-3]. Clinical IONPs are non-targeted and are generally used to image the blood pool, macrophage activity and sites of macrophage infiltration such as the liver, spleen and lymph nodes [4, 5]. IONPs have been used for in vivo imaging, in particular for stem cell tracking, as excellent signal-to-noise ratios are obtainable [6, 7]. Another attractive feature of IONPs is the relative ease of modifying their surface chemistry and thus the possibility of targeting specific biological targets by functionalization with targeting ligands [8-13]. In this study, we created a CXCR4 receptor targeted IONP by conjugating a cyclic pentapeptide moiety to epichlorohydrin cross-linked dextran coated IONPs and used these particles to detect MDA-MB-231 cells overexpressing the CXCR4 receptor.

The chemokine receptor CXCR4 [14] is a seven trans-membrane G-protein coupled receptor that is overexpressed in numerous types of cancers [15], including breast, brain [16], ovarian, pancreatic and prostate [17]. It has been shown to be involved in the metastasis of cancers such as breast, kidney, prostate, lung, pancreas, melanoma, neuroblastoma, non-Hodgkin’s lymphoma, multiple myeloma, ovarian cancer and malignant gliomas [15]. The interaction of the CXCR4 receptor with its natural ligand, stromal cell-derived factor 1α (SDF-1α) is critical for tumor development, growth and metastasis [18]. SDF-1α is the only major ligand of the CXCR4 chemokine receptor thus making this receptor-ligand pair an attractive target for...
CXCR4 targeted MRI contrast agent

imaging studies [15]. The receptor’s role in breast cancer and metastasis is particularly significant [19]. Metastases express high levels of CXCR4; it has therefore been suggested that CXCR4 levels could be predictive of metastatic potential, particularly in breast cancers [18, 19]. A number of studies have explored CXCR4 as a biomarker target over the past decade. Various radionuclides have been used to label CXCR4’s natural ligand SDF-1α [20], as well as anti-CXCR4 antibodies [16], peptide-based inhibitors [21-25] and small molecule inhibitors of CXCR4 [26-28] for nuclear imaging by single photon emission computed tomography (SPECT) and positron emission tomography (PET). Fluorescence and bioluminescence imaging probes have been developed using SDF-1α [29, 30], various peptides [31, 32], as well as small molecule inhibitors [33, 34]. Recent studies have experimented with more innovative imaging concepts, using metal nanoshells [35] and bimodal (fluorescence and SPECT) imaging agents for the detection of CXCR4 [36]. The advances in the field are summarized in two recent reviews by Knight et al [37] and van Leeuwen et al [38].

Since the discovery of the CXCR4 receptor and its role in HIV infection and cancer, several antagonists to the receptor have been identified and developed. One of the most potent compounds identified is cyclo-[Nal1-Gly2-(D-Tyr7)-Arg4-Arg5].

Figure 1. Structures of the original cyclopentapeptide CXCR4 antagonist FC131 (A), novel analogue with ornithine substitution and polyethylene glycol (PEG)-thiol linker (B) and synthesis of CXCR4 targeted fluorescent probe NalGYOrn(PEG4SH)R-TxR; (DMF/Phosphate buffer (pH 7), room temperature, 21 h) followed by semi-preparative HPLC purification (C).
CXCR4 targeted MRI contrast agent

Tyr$_3$-Arg$_5$ with an IC$_{50}$ of 4 nM, which is a cyclic peptide commonly known as FC131 (Figure 1A) [39]. Based on the structure of FC131, an ornithine substituted cyclic pentapeptide was developed [40], which possesses CXCR4 antagonist activity and provides an amine group for functionalization and subsequent bioconjugation with imaging probes.

In the first instance we conjugated the peptide cyclo-[Nal$_1$-Gly$_2$-(D-Tyr$_3$)-Orn$_4$-(PEG$_4$-SH)-Arg$_5$], (NalGYOrn(PEG$_4$SH)R, Figure 1B) to Texas Red and assessed the binding of the fluorescent probe to MDA-MB-231 cells. MDA-MB-231 cells have been shown to overexpress CXCR4 [41, 42]. Subsequently the NalGYOrn(PEG$_4$SH)R peptide was conjugated to IONPs for MR imaging of the CXCR4 receptor in vitro.

He et al have recently reported the development of anti-CXCR4 monoclonal antibody-labeled ultrasmall superparamagnetic iron oxide (USPIO) nanoparticles [43]. To the best of our knowledge, our research herein represents the first attempt to label an iron oxide-based MRI contrast agent with a small molecule CXCR4 specific targeting ligand.

Materials and methods

General methods

The NalGYOrn(PEG$_4$SH)R peptide was obtained from Cambridge Research Chemicals (UK), Texas red bromoacetamide was purchased from Invitrogen (UK) and all other chemicals and reagents were purchased from Sigma-Aldrich (UK). High performance liquid chromatography (HPLC) studies were conducted on a Waters system with the following modules: Waters 600 Controller module, a Waters 2487 Dual λ Absorbance Detector and a Waters Fraction Collector using Empower Software for analysis. MRI experiments were conducted on a 4.7 T Magnex magnet (Oxford, UK) Varian Unity Inova console (Palo Alto, CA).

Synthesis of NalGYOrn(PEG$_4$SH)R-TxR peptide probe

NalGYOrn(PEG$_4$SH)R (0.6 mg, 0.616 µmol) was dissolved in phosphate buffer (500 µL, K$_2$HPO$_4$ 10 mM, pH 7.0), to which was added Texas red bromoacetamide in dimethylformamide (DMF) (800 µL). The reaction was stirred for 21 h at room temperature. The mixture was then purified by semi-preparative HPLC on a Waters system with absorbance detection at 230 and 320 nm. A Waters Bondapak C18 column (7.8 x 300 mm) was used; t$_r$: 19 min, gradient mixture: 0-15 min: 10/90 MeOH/H$_2$O, 15-30 min: 75/25 MeOH/H$_2$O, 30-45 min: 82/18 MeOH/H$_2$O. HRMS: (TOF-MS ES+) calcd for C$_{84}$H$_{107}$N$_{13}$O$_{18}$S$_3$, 1681.70 found 1683.7 (M$^+$H$^+$).

In vitro incubation assay with NalGYOrn(PEG$_4$SH)R-TxR

MDA-MB-231 (ATCC, USA) cells were cultured in L-15 growth medium (ATCC, USA) supplemented with 10% Fetal Calf Serum (FCS), 2% Penicillin/Streptomycin (Sigma) solution and 0.2% fungicide and incubated at 37°C in a 5% CO$_2$/95% air atmosphere. NalGYOrn(PEG$_4$SH)R-TxR was dissolved in dimethyl sulfoxide (DMSO) to give a 2 mg/mL stock solution. Stock solution was diluted to give desired concentrations in L-15 growth medium. Cells were seeded (1.2 x 10$^3$ cells/well) 24 hours prior to experiment in 100 µL growth medium. The peptide conjugate was then added to each well at the appropriate dose. Cells were incubated for 60 min in a 5% CO$_2$/95% air atmosphere. The media was removed and the cells washed twice using ice-cold water. Phosphate-buffered saline (PBS) (150 µl) was added to each well and a fluorescent signal detected using a fluorescent plate reader (Victor) fitted with a Texas Red filter.

In vitro time course incubation assay with NalGYOrn(PEG$_4$SH)R-TxR

MDA-MB-231 cells were cultured as described above, the cells were trypsinized (1 mM ethylenediaminetetraacetic acid (EDTA)), washed and centrifuged (2000 rpm, 3 min); cells were re-suspended in PBS, and centrifuged again (× 2). Cells were counted using Trypan blue staining and 2 x 10$^5$ cells were added to each Eppendorf containing 1000 nM of NalGYOrn(PEG$_4$SH)R-TxR in PBS. Eppendorf tubes were incubated for 5, 30 or 60 min in a 5% CO$_2$/95% air atmosphere at 37°C. The reaction was stopped by adding 500 µL ice-cold PBS and keeping the Eppendorf tubes on ice. Samples were then spun down (13000 rpm, 4°C, 5 min) and supernatant removed using a fine glass needle under vacuum. Cell pellets were then washed using 500 µL ice-cold PBS and sam-
samples further spun down (6,500 rpm, 4°C, 5 min). Supernatant was removed and cells re-suspended in 150 µL PBS. Cell suspensions were transferred to 96 well plates and fluorescence detected using a Victor fluorescence plate reader fitted with a Texas Red filter.

**Fluorescence-activated cell sorting (FACS) analysis of MDA-MD-231 cells labeled with NaIGYOrn(PEG$_4$SH)R-TxR**

NaIGYOrn(PEG$_4$SH)R-TxR was dissolved in DMSO to give 2 mg/mL stock solution. Stock solution was diluted to give 100 nM and 1000 nM peptide conjugate in PBS. Eppendorf tubes were prepared with peptide conjugate diluted to give a final volume of 500 µL/tube. MDA-MB-231 cells were cultured as mentioned previously, washed, trypsanized (1 mM EDTA), centrifuged (2000 rpm, 3 min), and washed with PBS (× 2). The cells were then re-suspended in PBS and counted using Trypan blue staining. Eppendorf tubes were incubated for 60 min in a 5% CO$_2$/95% air atmosphere at 37°C. The reaction was stopped by adding 500 µL ice-cold PBS and keeping the Eppendorf tubes on ice. Samples were then spun down (13,000 rpm, 4°C, 5 min) and supernatant removed using a fine glass needle under vacuum. Cell pellets were washed using 500 µL ice-cold PBS and samples spun down (6,500 rpm, 4°C, 5 min). Supernatant was removed and cells re-suspended in PBS (1 mL) and transferred to FACS tubes. Unlabeled cells were used as a standard. Samples were kept on ice post incubation and analyzed using a BD LSR2 FACS analysis machine. Fluorescence was measured using a 610/620 YG-A filter. Results were analyzed using FlowJo 8.8.6 software.

**Ferrozine assay**

Iron content of IONPs was determined using a previously published protocol [44]. The ferrozine reagent was made by dissolving ferrozine (51.4 mg) and hydroxylamine hydrochloride (1 g) in a minimal amount of water, followed by the addition of concentrated (37%) hydrochloric acid (5 mL) and diluting the solution to a final volume of 10 mL using distilled water. An ammonium buffer was made by dissolving ammonium acetate (10 g) in ammonium hydroxide (8.75 mL) and diluting with distilled water to give a final volume of 25 mL. Hydrochloric acid was used to titrate the buffer to pH 5.5. Ferrozine (100 µL) reagent was added to iron oxide nanoparticle sample (50 µL) in an Eppendorf tube. The tube was heated using an oil bath at 60°C for 1 h. Samples were then treated with ammonium buffer (200 µL) and left for full color development (30 min). Samples were diluted to a final volume of 2 mL using distilled water for UV-Vis spectrometry (Perkin-Elmer Lambda 2 UV-Vis spectrometer). Endorem® (Guerbet, France) was used as a standard to produce a calibration curve for comparison.

**Synthesis of iodo-functionalized IONPs**

Succinimide iodoacetate (8.49 mg, 0.03 mmol) was dissolved in DMSO (30 µL) and added to amine-functionalized iron oxide nanoparticles (total volume 3 mL, 1.5 mg/mL) in HEPES buffer (0.01 M pH 8.5) and the reaction mixture stirred at room temperature for 1 h. The sample was dialyzed using a spectropore dialysis membrane (MWCO 3000, SpectrumLabs; Rancho Dominguez, CA, USA) in HEPES buffer (0.01 M
CXCR4 targeted MRI contrast agent

pH 7.0) overnight to remove unbound iodoacetate.

**Conjugation of NaI GyOrn(PEG₄SH)R to iodoIONPs**

NaI GyOrn(PEG·SH) (31.40 mg, 0.033 mmol) was dissolved in DMSO (70 µL) and added to succinimide iodoacetate-functionalized iron oxide nanoparticles (IONP) (4.5 mg) in 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) buffer (7 mL, 0.01 M pH 7.0). The reaction mixture was stirred gently for 21 h and thereafter purified by dialysis using a spectropore dialysis membrane (MWCO 3000, SpectrumLabs; Rancho Dominguez, CA, USA) in HEPES buffer (0.01 M pH 7.0) overnight in order to remove unbound material.

**Prussian blue staining of MDA-MB-231’s incubated with NaI GyOrn(PEG·SH)R-IONPs**

MDA-MB-231 cells were seeded on glass plates 24 h prior to experiment and cultured in L-15 growth medium (ATCC) supplemented with 10% Fetal Calf Serum (FCS), 2% Penicillin/Streptomycin solution and 0.2% fungicide (Sigma) and incubated at 37°C in a 5% CO₂/95% air atmosphere. NaI GyOrn(PEG·SH)R-IONPs, in addition to non-targeted IONPs (NH₂·IONPs) and Endorem®, were diluted in HEPES buffer (0.01 M, pH 7.0) to give [Fe] = 0.2 mg/mL. Cells were then incubated with 500 µL nanoparticle solution and 4500 µL growth medium to give a final concentration of 0.02 mg/mL. Cells were incubated for 240 min at 37°C in a 5% CO₂/95% air atmosphere. After incubation cells were washed with PBS (× 3) and detached using trypsin (500 µL). Trypsin was neutralized using 1 mL media and cells were centrifuged (13,000 rpm, 5 min). Media was removed and cells resuspended in 200 µL PBS. Samples were transferred to 200 µL Eppendorf tubes and cells were centrifuged again (2500 rpm, 5 min) in PBS (200 µL) and were placed in a quadrature ¹H coil and measurements taken at room temperature. MRI experiments were carried out as described above.

**MRI relaxivity measurements**

NaI GyOrn(PEG·SH)R-IONPs and Endorem® were diluted in HEPES (0.01 M, pH 7.0) to obtain 4 samples with varying amounts of Fe content ([Fe]: 0.24 mg/mL; 0.12 mg/mL; 0.06 mg/mL; 0.03 mg/mL). Samples were then added to Eppendorf tubes (200 µL) and placed in a quadrature ¹H coil and measurements taken at room temperature. MRI experiments were carried out using a 4.7 T Magnex magnet (Oxford, UK) Varian Unity Inova console (Palo Alto, CA, USA). T₂ values were obtained using saturation recovery experiments performed with a standard spin-echo sequence and a 9 mm single slice acquisition (TR = 3000 ms, TE values = 11.18, 15, 30, 60 and 120 ms), slice thickness: 9 mm, number of signal averages: 10, FOV: 100 × 50 mm², collected into a matrix of 256 × 128. MR images were analyzed using ImageJ (National Institutes of Health, USA), with a region of interest (ROI) drawn around the sample area. The T₂ values were calculated by plotting the mean signal intensity (Y) against the ROI taken at each TR and fitted to the equation: Y = M₀*exp(-x/T2) using GraphPad Prism software (San Diego, USA). The r₂ relaxivity was obtained from the linear fit of 1/T₂ vs [Fe] mM [45].

**MRI analysis of MDA-MB-231 cells incubated with NaI GyOrn(PEG·SH)R-IONPs**

MDA-MB-231 cells were seeded in T25 flasks (7.5 × 10⁴ cells/flask) and incubated at 37°C for 24 hours in a 5% CO₂/95% air atmosphere. NaI GyOrn(PEG·SH)R-IONPs, non-targeted IONPs and Endorem® were diluted in HEPES buffer (0.01 M, pH 7.0) to give [Fe] = 0.2 mg/mL. Cells were then incubated with 500 µL nanoparticle solution and 4500 µL growth medium to give a final concentration of 0.02 mg/mL. Cells were incubated for 240 min at 37°C in a 5% CO₂/95% air atmosphere. After incubation cells were washed with PBS (× 3) and detached using trypsin (500 µL). Trypsin was neutralized using 1 mL media and cells were centrifuged (13,000 rpm, 5 min). Media was removed and cells resuspended in 200 µL PBS. Samples were transferred to 200 µL Eppendorf tubes and cells were centrifuged again (2500 rpm, 5 min) in PBS (200 µL) and were placed in a quadrature ¹H coil and measurements taken at room temperature. MRI experiments were carried out as described above.

**Results**

**In vitro evaluation of NaI GyOrn(PEG·SH)R-TxR**

The cyclopentapeptide NaI GyOrn(PEG₄SH)R bearing a short PEG linker and reactive thiol functionality was coupled to Texas red bromoacetamide and dissolved in DMF to obtain NaI GyOrn(PEG₄SH)R-TxR (Figure 1C). The cou-
The fluorescent conjugated peptide were further assessed by flow cytometry experiments, where a concentration-dependent fluorescence shift from the MDA-MB-231 cells was observed following labeling with NaIGYOrn(PEG$_4$SH)R-TxR (Figure 3A).

Cellular localization of the conjugate MDA-MB-231 cells was investigated by fluorescence microscopy imaging; in these experiments, cells were incubated with 1000 nM of NaIGYOrn(PEG$_4$SH)R-TxR for 60 min at 37°C and then fixed and imaged. The fluorescence microscopy studies confirmed NaIGYOrn(PEG$_4$SH)R-TxR retention by MDA-MB-231 cells (Figure 3B and 3C) with a ubiquitous level of fluorescence observed throughout the cells indicating internalization of the peptide conjugate.

To establish MRI capability of the cyclic pentapeptide, NaIGYOrn(PEG$_4$SH)R (Figure 1B) was additionally conjugated to the surface of IONPs for MR imaging of the CXCR4 receptor.

IONPs with terminal amines (NH$_2$-IONPs) were prepared using the method described by Palmicci et al [72]. The iron and amine content of the nanoparticles was determined using ferrozine and fluorescamine assays respectively [44, 47]. Conjugation of the peptide to the surface of the IONPs was carried out as described previously [48]. Briefly, terminal amines on the surface of the IONPs were chemically modified to produce iodo-iron oxide nanoparticles (I-IONPs) using succinimide iodoacetate under basic conditions. The NaIGYOrn(PEG$_4$SH)R peptide was subsequently coupled to I-IONPs at neutral pH in (Figure 4). HEPES buffer yielding NaIGYOrn(PEG$_4$SH)R-IONPs. The IONPs were purified using Spectrapore dialysis membranes (MWCO 3000) to remove unbound peptide. IONPs were analyzed for their size and zetapotential following each modification procedure (Table 1) with NaIGYOrn(PEG$_4$SH)R-IONPs diameter measuring ~55 nm. The size and zetapo-
CXCR4 targeted MRI contrast agent

Figure 3. FACS analysis of MDA-MB-231 cells alone (red), incubated with a 100 nM (blue) and 1000 nM (green) dose of NalGYOrn(PEG_SH)R-TxR for 60 min at 37 °C (A), fluorescence microscopy images of MDA-MB-231 cells after incubation with NalGYOrn(PEG_SH)R-TxR (1000 nM, 60 min, 37 °C) (B), co-localization with DAPI (C).

Figure 4. Scheme of NalGYOrn(PEG_SH)R peptide conjugation to IONPs; Step 1: succinimide iodoacetate, amine functionalized IONPs, HEPES buffer (pH 8.5), room temperature, 1 hour; Step 2: NalGYOrn(PEG_SH)R, SIA-functionalized IONPs, HEPES buffer (pH 7), room temperature, 21 hours; Purification by dialysis.

Table 1. Iron oxide nanoparticles (IONPs) and physical characteristics

<table>
<thead>
<tr>
<th>Nanoparticle</th>
<th>Buffer</th>
<th>Size (nm)</th>
<th>Zeta Potential (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NH2-IONPs</td>
<td>HEPES, 0.01 M, pH 8.5</td>
<td>39.64</td>
<td>-0.75</td>
</tr>
<tr>
<td>I-IONPs</td>
<td>HEPES, 0.01 M, pH 7.0</td>
<td>21.45</td>
<td>0.29</td>
</tr>
<tr>
<td>NalGYOrn(PEG_SH)R-IONPs</td>
<td>HEPES, 0.01 M, pH 7.0</td>
<td>55.49</td>
<td>8.06</td>
</tr>
</tbody>
</table>

tential of the NalGYOrn(PEG_SH)R-IONPs increased due to aggregation effects induced by the peptide conjugation and the presence of the guanidinium group of arginine respectively. The guanidinium group (pKa 12.48) is positively charged under neutral conditions making the zetapotential of the IONPs more basic.

The concentration of peptide bound to the surface of the particles was determined using UV-Vis spectroscopy and the concentration of peptides was determined to be 6.0 mM, corresponding to approximately 90% peptide conjugation to the IONPs.

In vitro MRI of CXCR4 with targeted IONPs

To assess cellular binding of the NalGYOrn(PEG_SH)R peptide conjugated IONPs (NalGYOrn(PEG_SH)R-IONPs), MDA-MB-231 cells were incubated with nanoparticles for 60 min at 37 °C. To compare specificity of binding,
CXCR4 targeted MRI contrast agent

CXCR4 overexpressing MDA-MB-231 cells were incubated with NalGYOrn(PEG_SH)R-IONPs, non-peptide conjugated particles (IONPs) or the clinically used iron oxide nanoparticle Endorem® (total [Fe]: 0.1 mg/mL). To visualize the nanoparticles, cells were then incubated with a Prussian blue reagent and cell nuclei were counterstained with DAPI (Figure 5). Darker staining was observed from cells stained with NalGYOrn(PEG_SH)R-IONPs, which revealed dark blue rings around the cellular membrane, confirming NalGYOrn(PEG_SH)R-IONPs targeting to MDA-MB-231 cells. Minimal staining was observed for the non-targeted IONPs and Endorem controls (Figure 5B and 5D). Cell surface binding of the targeted IONPs to the cells and lack of intracellular uptake of the particles suggest/confirm that the peptide acts as an antagonist of the CXCR4 receptor. Therefore, the internalization of nanoparticles does not occur to any appreciable extent within the 60 min incubation period.

The MRI efficacy of the synthesized NalGYOrn(PEG_SH)R-IONPs was investigated by measuring $r_2$ relaxivity at 4.7 T and ambient temperature. Both NalGYOrn(PEG_SH)R-IONPs and non-targeted IONPs were incubated with CXCR4 overexpressing MDA-MB-231 cells in order to assess their MR signal enhancing potential. The particles were incubated with the cells for 4 h at $37^\circ$C at a total [Fe] 0.01 mg/mL. Following this incubation period, the cells were washed, trypsinized, pelleted in PBS and imaged at 4.7 T. The $r_2$ relaxivity of NalGYOrn(PEG_SH)R-IONPs was measured to be 3.66 mM$^{-1}$s$^{-1}$ which is comparable to other commercially available $T_2$ contrast agents.

MRI relaxation data obtained using region of interest quantification showed approximately

![Figure 5. Bright field images of MDA-MB-231 cells incubated with NalGYOrn(PEG_SH)R-IONPs (A), IONPs (B), Endorem® (C) ([Fe]=0.1 mg/ml, 60 min, 37°C) and cells alone (D); Iron was stained using Prussian Blue, cell nuclei were stained using DAPI, white arrows point to cell membrane stained areas.](image-url)
3-fold increase in $T_2$ signal from MDA-MB-231 cells labeled with NalGYOrn(PEG$_4$SH)R-IONPs in comparison to control non-targeted IONPs (Table 2). Additionally, Endorem® did not reduce $T_2$ signal post incubation with MDA-MB-231 cells suggesting no binding or uptake of this non-targeted agent with these cells.

**Discussion**

CXCR4 expression has been found to be an important biomarker for several cancers [49]; and CXCR4 up-regulation has been implicated in metastases and tumor invasion [19]. Therefore CXCR4 targeted imaging agents can be valuable for both the detection of primary tumors and secondary tumors arising as a result of metastasis.

In order to establish the targeting capability of the selected NalGYOrn(PEG$_4$SH)R for binding to the CXCR4 receptor, the peptide was conjugated to Texas Red to produce a NalGYOrn(PEG$_4$SH)R-TxR probe for fluorescence imaging. MDA-MB-231 cell associated fluorescence was measured post incubation with the fluorescent probe in a concentration dependent manner (Figure 2A). Interestingly, the time course incubation assay did not reveal a significant increase in the level of uptake of the probe from 5 min to 60 min suggesting rapid saturation of binding sites (Figure 2B). To assess binding of the peptide conjugate to CXCR4 cells, flow cytometry experiments were carried out and a concentration-dependent increase in fluorescence signal was observed (Figure 3A). Fluorescence microscopy studies of MDA-MB-231 cells labeled with the peptide conjugate revealed a uniform level of fluorescence visible in the cell cytoplasm, indicating the likely internalization of the probe into cells. It was postulated that intracellular fluorescence should not be observable as the targeting peptide is an antagonist of the CXCR4 receptor. It is likely, however, that the observed internalization is due to the lipophilic nature of the Texas Red conjugate. Although fluorescent probe conjugation to active targeting ligands is a relatively cheap and facile method of establishing specificity, it is important to note that cellular internalization due to the lipophilicity of aromatic structures in fluorophores via diffusion or endocytosis is a limiting factor. It is possible that active transport mechanisms lead to the internalization of the peptide conjugate as other fluorescent CXCR4 receptor antagonists bearing a similar fluorophore have also been shown to become internalized into cells as a result of active transport mechanisms [33].

Incubation with targeted nanoparticles, on the other hand, showed distinct cell surface association of the IONPs on MDA-MB-231 cells. For these studies, amino-coated IONPs were synthesized and characterized for their charge and size. Incubation of MDA-MB-231 cells with NalGYOrn(PEG$_4$SH)R-IONPs led to the appearance of intense blue rings around the Prussian blue stained cells (Figure 5). This intense stain-
ing was not visible in cells incubated with non-targeted IONPs or Endorem®. These images confirm the cell surface binding of the targeted nanoparticles. Non-targeted nanoparticles below 60 nm in diameter have been shown to enter cells through receptor-mediated endocytosis [50], with ~50 nm being an optimal nanoparticle size for cellular entry [51], however, we do not observe this behavior with NalGYOrn(PEG$_4$SH)R-IONPs suggesting “antagonistic” binding to the cell surface membrane as a dominant binding mechanism.

MRI of MDA-MB-231 cells incubated with NalGYOrn(PEG$_4$SH)R-IONPs, control non-targeted-IONPs, and Endorem® was carried out and $T_2$ measurements obtained at 4.7 T (Table 2). A hypointense signal was observed from the cell pellet region post incubation with NalGYOrn(PEG$_4$SH)R-IONPs, where the susceptibility artifact of the superparamagnetic IONPs was also apparent (Table 2). The measured $T_2$ % signal enhancement was up to 3-fold higher from MDA-MB-231 cells labeled with the targeted nanoparticles than with non-targeted-IONPs or Endorem®. The measured $T_2$ signal from non-targeted IONPs was lower than from Endorem®. This is likely to be due to the fact that the control IONPs, although non-targeted, have a surface decorated with amino groups which may become positively charged in cell culture media leading to attractive forces with the negatively charged phosphate anions of the cell surface membrane. The $T_2$ signal intensity of cells labeled with Endorem® did not decrease because the surface properties of particles in this formulation are neutral, detering interactions with the cell membrane.

Further work to establish the cytotoxicity of the targeted-IONPs on MDA-MB-231 and other cell lines overexpressing CXCR4 is currently underway, furthermore we aim to establish the utility of NalGYOrn(PEG$_4$SH)R-IONPs for MR imaging of CXCR4 overexpressing tumors in vivo.

**Conclusion**

In summary, we have synthesized a novel MRI contrast agent for the imaging of the CXCR4 receptor. Using an in vitro model based on the MDA-MB-231 cell line, we have demonstrated the feasibility of using targeted-fluorescent or -IONPs for this purpose. The majority of commonly used superparamagnetic IONPs in the clinic are non-targeted agents. Targeted imaging of specific receptors on cancer cells allows for more effective detection of cancer.

**Acknowledgments**

The authors thank Dr. William Jones for helpful discussions. We acknowledge support by staff at the Biological Imaging Centre at Imperial College London. Funding for this project was provided from CRUK, EPSRC, MRC and the Department of Health Centre grant (C2536/A10337). NJL would like to thank the Leverhulme Trust for a Research Fellowship.

**Address correspondence to:** Dr. Nicholas J Long, Department of Chemistry, Imperial College London, South Kensington, London, SW7 2AZ, UK. Phone: +44 (0) 20 7594 5781; Fax: +44 (0) 20 7594 5804; E-mail: n.long@imperial.ac.uk; Dr. Eric O Aboagye, Department of Surgery and Cancer, Comprehensive Cancer Imaging Centre, Imperial College London, Hammersmith Hospital, Du Cane Road, East Acton, London, W12 0NN, UK. Phone: +44 (0) 20 8383 3759; Fax: +44 (0) 20 8383 1783; E-mail: eric.aboagye@imperial.ac.uk

**References**


CXCR4 targeted MRI contrast agent


CXCR4 targeted MRI contrast agent


