Self-assembled polymeric nanoparticles as new, smart contrast agents for cancer early detection using magnetic resonance imaging

Abstract: Early cancer detection is a major factor in the reduction of mortality and cancer management cost. Here we developed a smart and targeted micelle-based contrast agent for magnetic resonance imaging (MRI), able to turn on its imaging capability in the presence of acidic cancer tissues. This smart contrast agent consists of pH-sensitive polymeric micelles formed by self-assembly of a diblock copolymer (poly(ethylene glycol-b-trimethylsilyl methacrylate)), loaded with a gadolinium hydrophobic complex (BuBipyGd) and exploits the acidic pH in cancer tissues. In vitro MRI experiments showed that BuBipyGd-loaded micelles were pH-sensitive, as they turned on their imaging capability only in an acidic microenvironment. The micelle-targeting ability toward cancer cells was enhanced by conjugation with an antibody against the MUC1 protein. The ability of our antibody-decorated micelles to be switched on in acidic microenvironments and to target cancer cells expressing specific antigens, together with its high Gd(III) content and its small size (35–40 nm) reveals their potential use for early cancer detection by MRI.

Keywords: micelle, pH-sensitive, self-assembly, smart contrast agent, cancer detection

Introduction

The recent developments in imaging techniques such as computed tomography, fluorescence imaging, positron emission spectroscopy, and magnetic resonance imaging (MRI) have led to major advances in the field of biomedical diagnosis and therapy. Currently, MRI is one of the most powerful diagnostic techniques for molecular imaging, mainly due to its noninvasive nature allied to other benefits, such as the ability to obtain direct multiplanar images, differentiate soft tissues, and obtain anatomical data while avoiding ionizing radiation. In addition, MRI allows sub-millimeter spatial resolution and has an excellent contrast resolution. However, in terms of sensitivity, MRI still lags behind other imaging tools. For example, MRI has a sensitivity of approximately $10^{-3}$ mol L$^{-1}$ to $10^{-4}$ mol L$^{-1}$, which compared to other types of imaging, can be very limiting. Also, because healthy and pathological tissues as well as distinct diseases show similar magnetic moments, they produce a poor image contrast. Thus, in order to overcome these shortcomings and obtain more informative images, a substantial quantity of contrast agent (CA) needs to be injected. CA molecules are paramagnetic chemical substances that act by shortening the longitudinal and transversal relaxation times of water protons, $T_1$ and $T_2$, respectively. The most frequently used in clinical MRI are small molecules of Gd(III) chelates, which are $T_1$ agents that due to their low molecular weight can distribute uniformly to all perfused tissues throughout the vasculature and can diffuse across endothelial wall vessels into the extracellular spaces.
Free Gd(III) is toxic because its diameter is close to that of calcium ions and, for this reason, it has to be chelated to a ligand. These ligands reduce Gd(III) toxicity significantly and influence the pharmacokinetics of the complex. Then again, these low-molecular weight CAs suffer from some weaknesses such as lack of selectivity toward particular organs or pathological areas, absence of response toward cellular microenvironment, short half-life in blood (a large part of administered CA is eliminated before images are taken), and potential toxic side effects for some patients.

Recently, scientists have developed new classes of medical imaging probes with low molecular weight called MRI smart CAs to overcome the classic CAs limitations. Unlike standard CAs, which enhance MR image constantly, smart CAs change between two states, depending on the type of the stimulus. This change is detected as an alteration in signal, because one of the states has low or no enhancement effect (“off state”), while the other induces a high enhancement (“on state”). These stimulus-sensitive agents can be detected when switching from one state to another after exposure to a metabolic or physiological event in a specific molecular target. A variety of smart CAs have been developed that become activated in response to changes in pH, to the presence of metal ions through enzymatic activation, and to the presence of oxygenated hemoglobin or radicals or upon exposure to ultrasounds or changes in temperature. These smart CAs have proven to be excellent MRI tools for very specific targets in vitro and in vivo. On the other hand, targeted nanoparticles such as liposomes, dendrimers, and micelles provide several unique features and capabilities such as to house a large number or different types of functional groups that can be used in multiple diagnostic (eg, gadolinium complex) and therapeutic (eg, anticancer) agents. Moreover, nanoparticles with the size ranging from 10 nm to 100 nm accumulate preferentially at tumor sites through an effect called enhanced permeability and retention (EPR). Finally, the capability of these nanoparticles of encapsulating a large amount of molecules (eg, gadolinium complex) inside their core will allow the detection of cancer at early stages (0, I, and II) due to the signal amplification generated by the large number of CAs released from the nanoparticles at the cancer tissue sites.

In this work, we aimed to develop a new type of smart CAs based on polymeric nanoparticles capable of amplifying the MRI signal chemically and at the same time have the ability to remain silent during circulation and active at the target site. These newly developed nanoparticle-based CAs consist of pH-sensitive polymeric stealth PEGylated micelles formed by self-assembly of a diblock copolymer, poly(ethylene glycol-b-trimethylsilyl methacrylate) (PEG-b-PTMSMA), loaded with the hydrophobic complex tetraaquodichloro(4,4′-di-tert-butyl-2,2′-bipyridine) gadolinium(III) chloride (BuBipyGd). To enhance the targeting ability of these polymeric nanoparticles toward cancer cells, we decorated micelles with monoclonal antibody (mAb) against the human MUC1 protein, which is aberrantly expressed in many epithelial cancers, including breast cancer.

Materials and methods

Reagents and equipment

Ethanol (96%) was from AGA, tetrahydrofuran (THF), HPCL grade, from Ridel-de-Haën, hydrochloric acid (37%) from Merck, N,N-dimethylformamide (DMF) and isopropanol from BDH Prolabo, methoxy(polyethylene glycol) N-hydroxysuccinimide ester and paraformaldehyde from Sigma-Aldrich, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT) from Applichem, and Trizol reagent from Invitrogen. For cell culture, complete Dulbecco’s Modified Eagle’s Medium or RPMI1640 culture media (DMEM or RPMI1640 basic medium [Lonza] supplemented with 10% fetal bovine serum [PAA], 2 mM glutamine, 100 units mL⁻¹ penicillin, and 100 μg mL⁻¹ streptomycin) were prepared. We purchased mAbs against the human MUC1 protein, C595 (Santa Cruz Biotechnology) and SM3 (eBioscience), and phycoerythrin-conjugated goat anti-mouse secondary antibody (Biolegend). Oligonucleotide primers were synthesized by Sigma-Aldrich. Milli-Q water (Millipore) was used in all procedures. For dialysis, cassettes were purchased from Pierce Biotech, 10-kDa MWCO. For ¹H nuclear magnetic resonance (NMR) experiments, a Bruker ARX 400 (400 MHz) spectrometer was used, and MR images were obtained in a Signa 1.5T SYS#GEMSOW, GE Healthcare, clinical MRI. Fourier transform infrared spectroscopy spectrum was recorded in a KBr pellet within the range of 4,000–600 cm⁻¹ in a Bruker, Tensor 27 spectrophotometer. Elemental analysis was performed on a Truspec 630-200-200 apparatus. Steady-state fluorescence spectra were recorded on a SPEX Fluoromax-3 spectrophotometer (Jobin Yvon–Horiba). All emission spectra were carried out in 1-cm quartz cuvettes (Hellma Quartz-Suprasil), using λₑₒ =334 nm, collecting the emission from 360 nm to 510 nm (increment of 1 nm), using 1-nm slit widths in excitation and emission (wavelength resolution of 1 nm), and corrected for nonlinear instrument response. For MTT assays, a Tecan Infinite M200 monochromatic-based multifunction microplate reader was used.
Nucleic acid concentration was determined on a NanoDrop 2000c Spectrophotometer (Thermo Scientific). Polymerase chain reaction (PCR) amplification was performed on a CFX96 Real-Time PCR Detection System (Bio-Rad). Flow cytometry was performed on a FACS Calibur analyzer, and data were processed with Cell Quest software (BD Biosciences). White light and fluorescence observation and image acquisition of living cells was performed with an inverted fluorescence microscope (Leica DM IL) coupled to a digital CCD camera (Evolution MP-5.1, Media Cybernetics). Observation and image acquisition of fixed cells was performed on an AxioObserver Z2 Fluorescence microscope (Zeiss), coupled to a digital AxioCam camera (Zeiss). Images were processed using AxioVision software (Zeiss). All light-scattering experiments were performed using Brookhaven Instruments BI-200SM Multiangle DLS & SLS Light Scattering System.

**Diblock co-polymer synthesis**

The macro-CTA weighing 0.8 g was placed in a Schlenk tube, previously charged with nitrogen and dissolved in 10 mL of THF. A volume of 3.5 mL (45 eq) of trimethylsilyl methacrylate and 0.030 g (1 mol% of the monomer) of AIBN were added. After three freeze–pump–thaw cycles, the mixture was incubated in an oil bath at 60° C, under stirring, for 24 hours. The flask was then cooled to room temperature, 10 mL of THF added to dilute the mixture, which had become very viscous, and its content transferred to a round-bottomed flask. After evaporation of the solvent, the white residue was dissolved in methanol and the polymer precipitated by the addition of water. The polymer was separated from the supernatant by centrifugation at 15,344 × g during 30 minutes, followed by decantation (Figure 1A). This procedure was repeated one more time, affording a white solid that was vacuum dried, at 60°C, overnight, yielding 1.47 g (40%) of the desired product. 1H NMR (DMSO) δ 0.056 (s, (CH$_3$)$_3$Si), 1.086 (s, CH$_3$C–C=O), 1.177 (s, CH$_3$C–C=O), 1.878 (s, CH$_2$C–C=O), 1.992 (s, CH$_2$C–C=O), 3.356 (s, CH$_2$–(CH$_2$O)$_n$), 3.639 (s, (CH$_2$CH$_2$O)$_n$), and 4.660 (q, CH$_3$CH$_2$O–C=S); DP$_n$(NMR) = 45. Gel permeation chromatography (GPC) analysis (PSSNa standards) reveals a monomodal molecular weight distribution, from which a value of $M_n$ = 9 kDa was predicted.

**Preparation of tBuBipyGd complex**

The preparation of this complex (Figure 1B) was based on a well-described method.$^{29,30}$ A mixture of 0.500 g (0.830 mmol) of GdCl$_3$·4H$_2$O and 0.223 g (0.830 mmol) of tBuBipy in 25 mL of ethanol was stirred and refluxed overnight. The resulting solution was concentrated, and the final product was obtained by the addition of diethyl ether. The white solid was washed with diethyl ether to remove unreacted tBuBipy and dried under reduced pressure to afford 0.450 g (90%) of tBuBipyGd complex. Elemental analysis

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**Figure 1** (A) Synthesis strategy of the co-polymer poly(ethylene glycol-b-trimethylsilyl methacrylate); (B) synthesis strategy of tBuBipyGd complex; and (C) encapsulation of contrast agent during the self-assembly of polymeric nanoparticles (micelles).

**Abbreviations:** THF, tetrahydrofuran; h, hour; AIBN, azobisisobutyronitrile; DMSO, dimethyl sulfoxide.
theoretical values are as follows: 35.77% C; 5.34% H; 4.64% N; and the obtained values were as follows: 35.99% C; 5.59% H; 4.69% N. Fourier transform infrared spectroscopy (KBr) values were as follows: 2,963 cm\(^{-1}\) (s), 2,866 cm\(^{-1}\) (s), 1,594 cm\(^{-1}\) (vs), 1,545 cm\(^{-1}\) (vs), 1,374 cm\(^{-1}\) (s), 837 cm\(^{-1}\) (s), 605 cm\(^{-1}\) (s). To obtain a 50 mM stock solution, 5.1 mg of the complex was dissolved in 169 μL of ethanol. The solution was kept at −20°C.

**Preparation of fluorophore-loaded and \textsuperscript{t}BuBipyGd complex-loaded polymeric micelles**

Polymeric micelles loaded with \textsuperscript{t}BuBipyGd complex were obtained as follows: 30 mg of PEG-\textsuperscript{b}-PTMSMA and 0.5 mg of either 1-methylpyrene or \textsuperscript{t}BuBipyGd complex was dissolved in 0.3 mL of DMF. The mixture was stirred for 3 hours at room temperature. Subsequently, 0.7–1.0 mL of pure Milli-Q water was added at a rate of one drop every 10 seconds to induce micellization. The resulting micelle solution was then dialyzed against 4 L of Milli-Q water for 3 days at room temperature. Subsequently, 0.7–1.0 mL of pure Milli-Q water was added at a rate of one drop every 10 seconds to induce micellization. The resulting micelle solution was then dialyzed against 4 L of Milli-Q water for 3 days at room temperature, using a 10-kDa cut-off dialysis cassette to exclude unencapsulated compound or nonself-assembled polymer molecules. The Milli-Q water was replaced twice a day.

**Measurement of Gd(III) complex and fluorophore concentrations loaded in micelles**

A 6.62×10\(^{-5}\) M stock solution of Gd(III) complex in ethanol was diluted to obtain standards with the following concentrations: 1.63×10\(^{-5}\) M, 1.66×10\(^{-5}\) M, 3.31×10\(^{-6}\) M, 1.66×10\(^{-6}\) M, and 3.31×10\(^{-7}\) M, and then obtain a Gd(III) complex calibration curve (Figure 2A). The corresponding absorbance values were recorded at 282 nm, and the molar absorption coefficient was 1.30×10\(^4\) dm\(^3\) mol\(^{-1}\) cm\(^{-1}\). From a 2.17×10\(^{-4}\) M stock solution of 1-methylpyrene in THF, five standards of concentrations 1.63×10\(^{-5}\) M, 1.09×10\(^{-5}\) M, 8.68×10\(^{-6}\) M, 5.43×10\(^{-6}\) M, and 1.09×10\(^{-6}\) M were prepared and a 1-methylpyrene calibration curve was obtained (Figure 2B). The absorbance values were measured at 344 nm and a value of 3.01×10\(^4\) dm\(^3\) mol\(^{-1}\) cm\(^{-1}\) for the molar absorption coefficient was obtained. For the determination of Gd(III) complex or fluorophore concentrations in loaded micelles, each solution was diluted (30 μL of solution to a total volume of 3 mL) in a spectrophotometer cuvette containing the appropriate solvent (ethanol for the former and THF for the latter) and the absorbance was read at 282 nm or 344 nm, respectively. The average concentration of \textsuperscript{t}BuBipyGd complex in the micelles was 0.065 mM, which represents 70% yield of contrast encapsulation. The average concentration of 1-methylpyrene in micelle solution was found to be 0.96 mM.

**Bioconjugation of polyethylene glycol with MUC1 antibodies**

Bioconjugation of PEG with C595 or SM3 mAb was performed as follows: 3 mg of methoxypolyethylene glycol N-hydroxysuccinimide ester was added to a solution containing 3 μg of mAb in 0.1 M bicarbonate buffer (500 μL, pH 8.3). The mixture was gently stirred for 5 days in slow tilt rotation at 4°C.

**Preparation of fluorophore-loaded bioconjugated micelles**

Fluorophore-loaded bioconjugated micelles were prepared by mixing 30 mg of PEG-\textsuperscript{b}-PTMSMA with 0.5 mg of

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**Figure 2** Relationship between the absorbance intensity and concentration of \textsuperscript{t}BuBipyGd complex and 1-methylpyrene at the optimum wavelength.

**Notes:** (A) Absorbance of \textsuperscript{t}BuBipyGd complex at 282 nm. (B) Absorbance of 1-methylpyrene at 344 nm.
1-methylpyrene and dissolved in 0.4 mL of DMF. The mixture was stirred for 3 hours at room temperature. Subsequently, 0.5 mL PEG-Anti-MUC1 aqueous solution was added at a rate of one drop every 10 seconds and, immediately after, 0.5 mL of cold Milli-Q water was added at the same rate to induce micellization. The resulting micelle solution was then placed in a dialysis cassette with a 10-kDa cut-off, to exclude unencapsulated compounds (either 1-methylpyrene or Gd complex) or possibly nonself-assembled polymer molecules and dialyzed against cold Milli-Q water for 3 days at 4°C. The recovered solution was then centrifuged to remove the free PEG-Anti-MUC1 and then the pellet was diluted once again with Milli-Q water.

Assessment of micelle stability in water and cell culture medium
1-Methylpyrene–loaded micelle solution (200 μL) was added to a cuvette containing 800 μL of either water or complete DMEM culture medium, and the fluorescence spectrum was acquired. The spectrum of 1-methylpyrene–loaded micelles biocjugated with antibody in culture medium was also obtained.

Determination of the ‘BuBipyGd complex in the $T_1$ value of water
‘BuBipyGd complex–loaded micelle aqueous solution (10 μL) was placed in an NMR tube containing deuterated water, and $T_1$ of water protons was measured using the inversion recovery method. The sequence was [180°, (16 μs) − τ (0.05, 0.25, 0.5, 1, 1.5, 2, 3, 5, and 10 seconds) − 90°, (8 μs) − $T_1$ (equal to τ) − D (12 seconds)]r. Subsequently, approximately 10 μL of acidic solution (pH 5.5) was added to the tube to induce micelle bursting, and thus ‘BuBipyGd complex release. The above procedure was repeated. Data were fit to an exponential, from which $T_1$ values were derived, using Bruker Topspin 2.0 software.

Determination of the ‘BuBipyGd complex imaging ability in vitro
To image ‘BuBipyGd complex released from micelles, the same volume of ‘BuBipyGd complex–loaded micelles in serum was dispensed in two microfuge tubes. Next, hydrochloric acid was added to only one of them to obtain approximately pH 5.5. Both tubes were positioned on top of an MRI phantom for calibration. MR images were subsequently acquired with a Spin Echo sequence (TR/TE=300/12 ms; slice thickness, 3 mm; field of view, 18×18 cm; matrix, 256×256).

Cell lines
The MCF-7 and MDA-MB-468 (human breast adenocarcinoma), HEK293T (Human Embryonic Kidney), and S17 (mouse bone marrow stroma) cell lines were maintained in complete DMEM culture medium. The Jurkat (human T-cell leukemia) cell line was maintained in complete RPMI1640 culture medium. Cells were cultured at 37°C under humidified atmosphere of 5% CO₂, unless stated otherwise.

In vitro cytotoxicity tests
Jurkat or MCF-7 cells were seeded on 96-well culture plates. Different concentrations (5 μM, 10 μM, 15 μM, and 20 μM) of ‘BuBipyGd complex buffer solution was added to the culture medium and incubated for 3 days. In addition, different concentrations of free and encapsulated ‘BuBipyGd complex (at concentrations of 10 μM or 50 μM) were added to the cells and incubated for 3 days. These assays included ‘BuBipyGd-loaded micelles dialyzed for 3 days and also nonencapsulated micelles. After incubation, toxicity was evaluated by the MTT assay. Briefly, 10 μL of 5 mg mL⁻¹ MTT solution was added to each well followed by further incubation for 4 hours. Then, 100 μL of 0.04 N HCl in isopropanol was added to the wells and formazan crystals were allowed to solubilize for 1 hour at room temperature. The formazan concentration was quantified by measuring the absorbance at 570 nm and 630 nm. The final optical density was calculated following the formula: optical density = $L_{570\text{nm}} - L_{630\text{nm}}$, where L was the obtained absorbance value for each wavelength. The toxicity was assessed by calculating the percentage of cell survival in relation to the control untreated group. The results were statistically compared by Student’s t-test using GraphPad Prism software.

Detection of MUC1 expression by reverse-transcriptase PCR and immunofluorescent flow cytometry
Total RNA was extracted from cultured cell lines using Trizol following the manufacturer’s instructions. The concentration and purity of isolated RNA was measured by spectrophotometry, and its quality assessed by agarose gel migration. RNA treated with RNase-free DNase I (Fermentas) was reversed transcribed using the RevertAid First Strand cDNA Synthesis Kit (Fermentas), and quantitative PCR was performed using the iQ SYBR-Green Supermix kit (Bio-Rad). Primers to detect expression of MUC1 and the ACTB reference gene were used at a 200 mM concentration: MUC1-Forward, 5’-GTG CCC CCT AGC AGT
ACC G-3' ; MUC1-Reverse, 5'-GAC GTG CCC CTA CAA GTT GG-3'; ACTB-Forward, 5'-AGG CCA ACC GCG AGA AGA TGA C-3'; and ACTB-Reverse, 5'-AGG TCC AGA CGC AGG ATG GCA T-3'. Relative mRNA expression levels were calculated through the 2-ΔΔCt method. For MUC1 protein detection by flow cytometry, HEK293T, MCF-7, and MDA-MB-468 cells were dissociated by 0.02% EDTA treatment, washed, and labeled with SM3 mAb (1/500 dilution), C595 mAb (1/100 dilution), or an isotopic nonspecific antibody and phycoerythrin-conjugated goat anti-mouse secondary antibody. Each antibody incubation was performed for 1 hour on ice and was followed by PBS/3% fetal bovine serum/5 mM NaN3 washes, before flow cytometry analysis.

Assessment of polymeric micelle cellular uptake by microscopy
MCF-7 or MDA-MB-468 cells were seeded in culture plates and allowed to attach for 24 hours at 37°C. Micelles bioconjugated with C595 or SM3 antibody and loaded with 1-methylpyrene were added to the cells and incubated for 1 hour at 37°C. As a control, we used non-conjugated 1-methylpyrene–loaded polymeric micelles. Excess micelles were removed by washing additional three times with PBS. For mixed cell cultures, MDA-MB-468 and S17 cells were either grown together (proportion of 1–5, respectively) or alone on glass cover slips. The cells were allowed to adhere for 24 hours at 37°C, and then, anti-MUC1–conjugated 1-methylpyrene–loaded micelles were added to the cells and incubated for 1 hour at 37°C. Cells were then washed three times with PBS and fixed with 4% paraformaldehyde for 30 minutes. After washing, the cover slips were mounted on microscope slides and observed on a fluorescence microscope.

Results
Polymeric micelle preparation
Poly(ethylene glycol-b-trimethylsilyl methacrylate), which contains silicon moieties that can be cleaved under mildly acidic conditions (pH between 5 and 6), was synthesized by Reversible Addition-Fragmentation chain Transfer method (Figure 1A). The amphiphilic diblock co-polymer (PEG-b-PTMSMA) micelles were designed to accommodate the hydrophobic CA (BuBipyGd) within its core and to release its load at the site of interest, via the hydrolysis of its silyl ester groups located in the hydrophobic moiety of the copolymer in a mildly acidic medium, such as cancer tissue.

The ‘BuBipyGd complex was obtained from the reflux of GdCl3·4H2O and ‘BuBipy in ethanol. The structure [Gd/BuBipy(H2O)4(Cl)2]Cl was assessed based on the literature data for the complex [GdBipy(H2O)4(Cl)2]Cl (Figure 1B). Elemental analysis and Fourier transform infrared spectroscopy spectrum were in accordance with the proposed structure. We chose this complex as a model compound because of its hydrophobicity, conferred by the tert-butyl group in the bipyridine core, as well as the expected strong MR signal due to the high number of coordinated water molecules.

In the encapsulation process, ‘BuBipyGd complex and PEG-b-PTMSMA were co-dissolved in DMF and self-assembly was induced by slow addition of water. Due to its hydrophobic nature, the complex tends to lay within the micelle core (Figure 1C). Organic solvent was then removed by dialysis to yield micelles with the size ranging from 35 nm to 40 nm (Figure 3A, B). The ‘BuBipyGd complex concentration in micelles determined by spectrophotometry was on average of 0.065 mM.

The approximate micelle aggregation number (Nagg) can be calculated from the ratio of the average molecular weight of the micelle to the molecular weight of the block copolymer. Light-scattering experiments (SLS and DLS) indicated that the average molecular weight of the micelle is 8.1×106 Da (individual co-polymer weight of around 9 kDa). Thus, we estimated an aggregation number of 9. Accordingly, the maximum number of micelles formed in a solution of 1 mL was found to be approximately 4.2×1010. By correlating the dye concentration with its absorbance at 344 nm, we quantified the total number of dye molecules as 6.3×1010 after they were released into the solution. Because we washed off any unencapsulated dye in the solution before the destruction of micelles, we can estimate that one micelle with an average diameter (Rh=30 nm, Nagg = 9) can load at least 150,000 dye molecules and around 75,000 Gd(III) complex if we assume that the volume of Gd(III) doubles the volume of 1-methylpyrene. Therefore, the release of these dye molecules into solution should provide a significant signal amplification effect for better resolution and sensitivity.

Polymeric micelle stability and selectivity study
To test the in vitro stability and uptake of micelles by cells, 1-methylpyrene fluorophore was loaded into micelles instead of ‘BuBipyGd complex, because of the feasibility of fluorescence detection and quantification in cells. The concentration of 1-methylpyrene in these micelles, as determined
by spectrophotometry, was on average of 0.96 mM. In the fluorescence spectrum of 1-methylpyrene–loaded micelles (Figure 4A), the observation of both monomer (375 nm) and excimer (480 nm) emission bands clearly confirms the encapsulation of the fluorophore. When nonconjugated and bioconjugated micelles were diluted in cell culture medium, there were no significant changes in the fluorophore fluorescence spectrum (Figure 4A, B, C), which is indicative of maintenance of micelle integrity. However, and in spite of being stable in aqueous solutions for weeks, when the solution pH was lowered to values between 5 and 6, a progressive release of the fluorophore was observed.

Figure 3 (A) Dynamic light-scattering results for loaded polymeric micelles after the encapsulation of the contrast agent. (B) TEM of contrast-loaded polymeric micelles after the self-assembly of poly(ethylene glycol-b-trimethylsilyl methacrylate).

Note: Magnification: ×200,000. Bar scale: 50 nm.

Abbreviation: TEM, transmission electron microscopy.

Figure 4 1-Methylpyrene (MePy) fluorophore-loaded polymeric micelle stability in water and cell culture medium.

Notes: Fluorescence intensity spectra of 1-methylpyrene monomer (375 nm) and excimer (480 nm) was measured to detect disintegrated and intact micelles, respectively, and these were found to be stable in (A) water and (B) culture medium. (C) Micelle bioconjugation with the SM3 MUC1 monoclonal antibody did not alter stability in culture medium.
Determination of Peg-b-PTMSMA micelle-based CA imaging ability

The paramagnetic Gd(III) ions in complexes used as CA in MRI shorten the relaxation time $T_1$ of the water protons in their proximity, leading to an increase of the image contrast. Therefore, restricting water access to those chelates should prevent them from decreasing the $T_1$ of the surrounding water molecules. As a result, the CA remains switched-off in neutral pH but switched-on at acidic pH, which is a typical characteristic of malignant tumors. This feature was first demonstrated in vitro using 1H NMR, where the $T_1$ of water protons was initially measured in a sample containing intact ‘BuBipyGd-loaded micelles and it was found to be 3.4 seconds, but upon micelle disruption, by the addition of an acid solution (pH 5.5), the $T_1$ value dropped to 1.2 seconds (Figure 5A).

Further testing using a clinical MRI was performed. This experiment consisted of imaging two microfuge tubes containing ‘BuBipyGd complex–loaded micelle in water, one at neutral pH and the second with pH 5.5. The first sample, with neutral pH, did not change the background image signal and no contrast was observed (Figure 5B, square a), while the second tube presented a significant signal enhancement visible as a bright area (Figure 5B, square b). These results indicate that the micelles disassembled at low pH and released the ‘BuBipyGd complex. Consequently, the complex interacted with the surrounding water molecules and decreased the $T_1$ relaxation time, which leads to an increase in the image signal intensity from that specific area. Together, these experiments show the ability of these ‘BuBipyGd-loaded micelles to respond to an acidic condition by changing their states (on and off), as indicated by the alteration in $T_1$ time and image signal intensity.

In vitro cytotoxicity tests

To assess the cytotoxicity of free ‘BuBipyGd complex, we tested its effect on two different human cell lines, the Jurkat leukemia cell line and the MCF-7 breast cancer cell line. By performing MTT assays, we observed that the median lethal dose (LD50) of ‘BuBipyGd complex for the MCF-7 and Jurkat cell lines was, respectively, 12.8 μM and 10.2 μM (Figure 6A), indicating that the ‘BuBipyGd complex is toxic at the micromolar range. Further cytotoxicity studies demonstrated that encapsulation of ‘BuBipyGd complex inside polymeric PEG-b-PTMSMA micelles greatly decreased its toxicity (Figure 6B). The difference in cytotoxicity was most evident at 50 μM of ‘BuBipyGd, the highest concentration tested, where most cells treated with encapsulated complex for 3 days were still viable, while most cells treated with free ‘BuBipyGd complex were dead (Figure 6B). The observation that Gd(III) complex–loaded micelles were not cytotoxic corroborated their observed stability in vitro (Figure 4), because neither during dialysis nor during contact with cell culture medium there appeared to occur significant release of the complex and consequent increased cell death. Cytotoxicity tests also showed that up to 1 mg mL$^{-1}$ of PEG-b-PTMSMA micelles alone were not toxic to Jurkat cells (data not shown), further supporting the notion that encapsulation of ‘BuBipyGd complex within PEG-b-PTMSMA micelles prevents

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**Figure 5** (A) Scheme of 1H NMR relaxometric experiment to measure $T_1$ of water before and after the release of the contrast agent from the polymeric micelles. (B) Magnetic resonance image of microfuge tubes containing ‘BuBipyGd complex–loaded micelles, at pH 7 (square a) and at pH 5.5 (square b).

Abbreviation: NMR, nuclear magnetic resonance.
cytotoxicity. The observed protection from content toxicity provided by our polymeric micelles, an important property for their in vivo administration as nanocarriers, should lead to reduced or no toxicity to healthy tissues, at least until the micelles reach the appropriate acidic microenvironment and disassemble. This important feature will open the door to new types of dual-function CAs that detect and kill at the same time the cancer cells in early stages, these dual-function contrasts are currently under investigation in our group.

Targeting of MUC1-expressing breast cancer cells by bioconjugated micelles

An important desired feature of the designed micelles, besides the pH-sensitive release mechanism, is the ability to specifically target cancer cells. To obtain this capability we bioconjugated micelles with an antibody against a cell surface tumor antigen. For this work, the MUC1 glycoprotein was chosen as a target for our new MRI CA because it is often overexpressed in human cancers, notably in breast cancer.27 Furthermore, MUC1 has shown a great potential for early detection and staging of tumors, as well as for the assessment of tumor responses to therapy and detection of disease recurrence.27 In addition MUC1 has been tested as target for quantum dot and gold nanoparticle–mediated cancer diagnosis or therapy.31–33 Confirming previously published data,34 we verified that the MDA-MB-468 and MCF-7 human breast adenocarcinoma cell lines, but not the unrelated HEK293T cell line, expressed high levels of MUC1 mRNA (Figure 7A)
and MUC1 protein at the cell surface (Figure 7B), indicating their suitability as cancer models to test our bioconjugated micelles. Of two different mAbs tested by flow cytometry, SM3 showed higher affinity for surface MUC1 protein in both cell lines, indicating its potential to be more suitable for breast cancer cell targeting (Figure 7B).

To test the ability of bioconjugated micelles to target breast cancer cells, we loaded micelles with the 1-methylpyrene fluorophore instead of BuBipyGd complex, because of the fast and ready detection by microscopy of internalized fluorescence in targeted cells. After 1 hour incubation with micelles loaded with this fluorophore, followed by unbound micelle removal, cells were observed on a fluorescence microscope. Thus, MCF-7 cells incubated with C595 anti-MUC1–conjugated micelles presented more fluorescent staining than cells incubated with nontargeted micelles, in which no specific fluorescence was observed (Figure 8A). C595-conjugated micelles targeted both MCF-7 and MDA-MB-468 breast cancer cells, but micelles conjugated with the higher affinity SM3 mAb conferred increased fluorescence to target cells (Figure 8B). Together, these results demonstrated the specificity of the targeted micelles to MUC1-expressing cells and demonstrated that micelles released their content within cells.

Next we used another approach to test the targeting specificity of anti-MUC1 micelles. The goal was to observe whether anti-human MUC1 micelles were able to specifically target MDA-MB-468 cells, even when these cells were
mixed in a lower proportion (1:5) with unrelated cells from another species, the mouse S17 bone marrow stromal cell line. By incubating this mixed culture with anti-MUC1 polymeric micelles, we observed that MDA-MB-468 cells, which display spherical morphology, incorporated far more 1-methylpyrene fluorescence than S17 cells (Figure 9). In this experiment little fluorescence of the elongated morphology cells (S17) was observed, in contrast to the larger and brighter spots in round-shaped MDA-MB-468 cells (Figure 9A).
**Discussion**

The present study reports the development of a smart CA composed of pH-sensitive micelles containing a hydrophobic Gd(III) complex with the aim of specifically detecting cancer by MRI. Two main features confer high specificity for these CAs to detect cancer tissues: (i) pH sensitivity, which leads to a contrast signal only in acidic microenvironments, and (ii) antibody conjugation, which leads to preferential targeting to tumor tissues expressing the respective antigen.

Our experiments demonstrated that when encapsulated in these micelles, the CA remained in the off state, being activated only upon micelle disruption in an acidic medium. Several researchers have reported pH-sensitive particles such as liposomes with the aim of detecting cancer tissues or other acidic pathologic areas by MRI. However, pH sensitivity was the only feature in these tools for cancer targeting, besides the enhanced permeability and retention effect. Other researchers have developed nanoparticles conjugated to biomolecules (eg, antibody or peptides) to target and image cancer. Our results with MUC1-expressing cell lines indicate that the conjugation of self-assembled polymeric micelles with a tumor-specific mAb increases significantly their affinity toward cancer cells. Therefore, our bioconjugated pH-sensitive micelles present two features enabling targeting and imaging of cancer. This characteristic is instrumental for the application of these smart micelles as carriers to deliver CAs to MUC1-expressing cancer tissues. In this work we have tested two mAbs targeting the MUC1 surface protein, often overexpressed in mammary and other adenocarcinomas, such as gastric, colorectal, lung, prostate, ovarian, pancreatic, and bladder cancers. The C595 mAb, which reacts with the tetrameric motif RPAP present on the tandem repeats in the extracellular region of the MUC1 protein, has already been proven to be an efficient and specific targeting vector in preclinical and clinical trials of radioimmunoscintigraphy and radioimmunotherapy. The SM3 mAb reacts against the APDTRP sequence, which is exposed only in underglycosylated MUC1 protein. Because underglycosylated MUC1 is very frequent and specific for breast and other epithelial cancers, the SM3 mAb should be highly suitable to conjugate the micelles to target cancer tissues. The possibility of conjugating our polymeric micelles with other antibodies specific for different tumor antigens should enable their application to a wide range of human cancers. The identification of cancer biomarkers is an area of intense research, so novel cell surface biomarkers may become potential targets for our nanoparticle-based smart CAs.

**Figure 9** Increased targeting of anti-MUC1 1-methylpyrene-loaded micelles to MDa-MB-468 breast cancer cells.

**Notes:** (A) MDa-MB-468 and s17 cells cultured together in a 1:5 proportion were incubated for 1 hour with C595 mAb–bioconjugated micelles. (B) Bright field microscopy of MDa-MB-468 cells. (C) Bright field microscopy of S17 cells.
Conclusion

The presented work demonstrated that the developed polymeric micelle-based CA is able to turn on its imaging ability upon a pH decrease. The encapsulation of the active complex within micelles decreased its toxicity and allows the use of small amounts of CA (1.08 μmol kg⁻¹) in the clinic, as compared to current protocols, to avoid potential side effects. These features allied with the finding that micelle conjugation with an antibody specific for a tumor antigen increased their uptake by cancer cells and with their small size, turns this polymeric nanoparticles into a promising new tool for cancer early detection by MRI.

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Disclosure

The authors report no conflicts of interest in this work.

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