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Comparison of MRI Properties between Multimeric DOTAGA and **DO3A Gadolinium-Dendron Conjugates**

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S Supporting Information

ABSTRACT: The inherent lack of sensitivity of MRI needs the development of new Gd contrast agents in order to extend the application of this technique to cellular imaging. For this purpose, two multimeric MR contrast agents obtained by peptidic coupling between an amido amine dendron and GdDOTAGA chelates (premetalation strategy, G1-4GdDO-TAGA) or DO3A derivatives which then were postmetalated (G1-4GdDO3A) have been prepared. By comparison to the monomers, an increase of longitudinal relaxivity has been observed for both structures. Especially for G1-4GdDO3A, a marked increase is observed between 20 and 60 MHz. This structure differs from G1-4GdDOTAGA by an increased rigidity due to the aromatic linker between each chelate and



the organic framework. This has the effect of limiting local rotational movements, which has a positive impact on relaxivity.

INTRODUCTION

Due to its high spatial and temporal resolutions, magnetic resonance imaging (MRI) is a mainstream method for clinical diagnosis and therapeutic decision-making. The MRI signals arise from the ¹H abundancy in the body. When the body is exposed to a strong magnetic field, the ¹H nuclei, which are magnetically active, orient themselves within this magnetic field. When the body is then submitted to radiofrequency pulses, the energy as well as the orientation of the nuclei change and as they relax to their initial states, a resonance radio-wave is emitted. The recovery of this MRI signal and its spatial encoding leads to image production.¹ In MRI, the main contrast parameters are ¹H density, longitudinal T_{1} , and transversal T_2 proton relaxation times. Although T_1 and T_2 are generally reduced in injured tissues, there is considerable overlap of relaxation times between healthy and injured damaged tissues on native MRI. This stresses the need for MRI contrast agent injection to enhance the contrast.

Paramagnetic contrast agents are known to exert a strong stimulation to T_1 relaxation, and gadolinium-based contrast agents (GBCAs) are ideal candidates for producing MRI contrastophores that affect T_1 of water protons in tissues.² The MRI effectiveness of GBCAs is evaluated from their longitudinal r_1 relaxivity value, which refers to the increase in

the inverse of T_1 per millimolar metal ion. The GBCAs efficacy to affect the relaxation of protons in tissues is related to several parameters such as its molecular rotational correlation time τ_{R} the number of inner-sphere water molecules q, or the exchange rate of these inner-sphere water molecules $(k_{ex})^{2,3}$ Low molecular weight GBCAs are the active substances of the commercially and clinically available MRI contrast agents.⁴ Their low molecular weight favors their rapid elimination from the body through the renal pathway, which effectively avoids deleterious side effects associated with gadolinium (Gd) body accumulation (e.g. nephrogenic systemic fibrosis (NSF)).5 Moreover, the relaxivity of small GBCAs is limited by their rapid tumbling motions which results in a fast rotational correlation time τ_R (~10⁻¹⁰ s) and then a low relaxivity between 20 and 60 MHz (3–5 mM $^{-1}$ s $^{-1}$).

In order to amplify the MRI efficacy, the conjugation of small GBCAs with macromolecular carriers has been investigated.^{2,3,6} Among them, dendrimers and particularly polyamidoamines (PAMAM) have been explored because of their more rigid structure, and GBCA conjugation with PAMAM dendrimers of different generations was carried out.⁸

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As the result of the reduced rotational tumbling rate $(1/\tau_R)$, the proton relaxivity was improved at intermediate fields, but the efficiency of dendrimeric GBCAs did not show a linear dependence with their molecular mass. Besides, the enhancement clearly tends to diminish for larger systems. Moreover, high-generation dendrimers are not easy to prepare and to purify,⁸ and a latent cytotoxicity is observed due to a slow degradation *in vivo.*⁹

In this report, we propose to take advantage of the dendronization strategy to design multimeric GBCAs. The dendritic structures prepared in this work (Scheme 1) rely on a common bifunctional amido amine core (G1-4NH₂). The installation of MRI contrastophores will herein be carried out by peptidic coupling with the amino groups of the G1-4NH₂ core. On the resulting G1-4GdDOTAGA and G1-4GdDO3A structures, there will remain an activatable group (benzyl ester group), which may be released in the future, in order to introduce either a targeting ligand or a second imaging probe. Indeed, in order to reduce ambiguities in the MRI signal analysis, it will be interesting to develop contrast agents able to multiplex an MRI probe to a more sensitive one (for instance an optical imaging probe),¹⁰ provided that differences in concentration imposed by differences in sensitivity are taken into account in the system design.

Two different approaches were used to obtain the desired multimeric structures. For the first one, G1-4GdDOTAGA, four premetalated GdDOTAGA chelates were grafted on the central G1-4NH₂ core. The second one G1-4GdDO3A, was obtained by covalent linking of four DO3A macrocyclic units on the same central G1-4NH₂ core, followed by Gd metalation. In order to ensure a good thermodynamic stability of the corresponding metal complexes, these two structures were designed on the basis of macrocyclic GBCAs, and their synthesis is described in the paper. Before considering any additional functionalization of these multimeric GBCAs, we have evaluated their respective efficacy to enhance MRI

sensitivity as well as their kinetic inertness. Finally, their harmlessness toward fibroblasts was investigated.

EXPERIMENTAL SECTION

All reagents were purchased from commercial suppliers and used without further purification. ¹H NMR and ¹³C NMR spectra were recorded on a 600 MHz Bruker Avance III spectrometer equipped with a TCI cryoprobe and on a Bruker Avance Neo 600 MHz. The chemical shifts were reported in parts per million (δ scale). Traces of residual solvents were used as internal standards for ¹H and ¹³C chemical shift referencing. Abbreviations: s: singlet, d: doublet, t: triplet, q: quadruplet, m: multiplet, ws: wide signal.

FTIR absorption spectra were obtained on a Thermo Scientific Nicolet iS5 spectrometer equipped with ATR iD5 accessory, and wavenumbers are reported in cm^{-1} .

All mass spectra were collected by using a SYNAPT G2-Si HDMS (Waters Corp., Manchester, UK) equipped with an electrospray ionization source. ESI mass spectra were acquired over the m/z 100–4000 range. The desolvation gas was set to 800 L·h⁻¹ at a temperature of 200 °C, the cone gas was set to 40 L·h⁻¹, and the source temperature was set to 120 °C. The capillary voltage was set to 2500 V. Spectra were acquired in continuum and negative mode.

Column chromatographies were performed using silica gel (230–400 mesh) (Macherey-Nagel GmbH, Germany). Inversed phase chromatography was performed on C18 bonded silica (45–75 μ m, VersaFlash). The excess of Gd³⁺ ions was removed on Chelex 100 sodium form resin (50–100 mesh, purchased from Sigma).

Syntheses. The syntheses of starting dendrimer G1-4NH₂, DO3A-based ligand in the form of activated ester DO3A(*t*Bu)₃NHS, as well as mononuclear complexes GdDOTAGA and GdDO3A-COOH were performed as described in the Supporting Information (SI 1, 2, Supporting Information).

G1-4GdDOTAGA Synthesis. The **G1-4NH**₂ dendrimer (200 mg, 0.20 mmol, 1 equiv) and Gd-chelate **Na**₂**GdDOTAGA 5** (84% purity (SI 2, Supporting Information) 955 mg, 1.19 mmol, 6 equiv) were loaded into a flask flashed with argon. Anhydrous DMSO (50 mL) and triethylamine (166 μ L, 1.19 mmol, 6 equiv) were added, and the mixture was homogenized in ultrasonic bath.

After addition of 1-[bis(dimethylamino)methylene]-1H-1,2,3triazolo[4,5-*b*]pyridinium3-oxidhexafluorophosphate (HATU, 452

mg, 1.19 mmol, 6 equiv), the mixture was stirred at room temperature for 1 day. Addition of ethyl acetate (450 mL) provoked the formation of a precipitate that was recovered and washed two times with ethyl acetate (30 mL). The precipitate was taken up with 10 mL of distilled water; the resulting solution was split into two, diluted to 12 mL with water and loaded into two Vivaspin 15R tubes (MWCO 2000 g/mol) for ultrafiltration (6000 g for 45 min). Three ultrafiltration cycles were performed afterward; the remaining solutions were transferred into a round-bottom flask and evaporated under reduced pressure to afford G1-4GdDOTAGA (499 mg, Gd-purity found by relaxometric study 89%, 0.14 mmol, yield 73%) as a crystalline solid. The absence of free gadolinium ions was checked by a colorimetric test with xylenol orange as a dye.¹¹ FTIR: 1595 cm⁻¹. HR-MS (ESI): m/z calculated for [M - H]-: 3071.8726/found: 3071.8738; calculated for [M -2H]²⁻: 1535.4324/found: 1535.4336 (values are given for the highest component of isotopic profile; M≡C₁₀₇H₁₆₇Gd₄N₂₅O₄₀). Detailed HRMS spectra with simulated isotopic distribution profiles are given in SI 3, Supporting Information.

G1-4GdDO3A Synthesis. $G1-4[DO3A(tBu)_3]$. To a solution of **G1-4NH**₂ dendrimer (0.411 g, 0.42 mmol, 1 equiv) in DMF (5 mL) and DIPEA (0.99 mL, 5.83 mmol, 14 equiv) was added the solution of the activated ester **6** (2.112 g, 2.67 mmol, 6.3 equiv, prepared as described in SI 4, Supporting Information) in dichloromethane (5 mL), and the mixture was stirred for 48 h under a nitrogen atmosphere at room temperature. The solution was diluted with ethyl acetate (15 mL) and extracted three times with distilled water (3 × 30 mL) and brine (3 × 30 mL). Organic phase was dried over sodium sulfate, filtered, and concentrated under reduced pressure. A white solid was isolated (0.48 g) containing the desired compound as revealed by mass spectrometry. It was used in the further synthetic step without purification.

HR-MS (ESI): m/z calculated for $[C_{171}H_{279}N_{29}O_{36}+Cl]^-$: 3351.0620/found: 3351.0625.

G1-4DO3A. The compound G1-4[DO3A(tBu)₃] (0.48 g) was dissolved in a mixture of dichloromethane and trifluoroacetic acid (20 mL, 1:1 v/v), and this solution was stirred at room temperature for 48 h. The reaction mixture was concentrated under reduced pressure, and the residue precipitated in diethyl ether and washed three times with this solvent (3 × 30 mL). The obtained product was solubilized in distilled water (10 mL) and purified by dialysis (dialysis membrane MWCO 500–1000 Da, VWR) against 1 L of distilled water (which was changed four times, after 2.5, 6.5, 11.5, and 18 h). After dialysis, the resulting solution was freeze-dried and the desired dendron G1-4DO3A obtained (0.40 g, 0.15 mmol, yield 36% over two steps from G1-4NH₂).

¹H NMR (600.16 MHz, DMSO-*d*₆, 25 °C): δ 1.81 (ws, 4H, -NHCH₂C<u>H₂CH₂N<)</u>, 1.90 (ws, NHCH₂C<u>H₂CH₂N<)</u>, 2.8–5.4 (ws, dendrimer CH₂ and cyclen CH₂), 3.60 (s, 2H, >NC<u>H₂COOBn</u>), 5.21 (s, 2H, $-\text{COOCH}_2\text{Ph}$), 7.32–7.44 (m, 5H, C<u>H</u>), 7.66 (d, 8H, ³*J* = 8.4 Hz, C<u>H</u>), 7.83 (d, 8H, ³*J* = 8.4 Hz, C<u>H</u>), 8.63 (s, 4H, N<u>H</u>), 8.86 (s, 2H, N<u>H</u>) 10.78 (ws, 4H, N<u>H</u>).

¹³C NMR (150.9 MHz, DMSO- d_6 , 25 °C): δ 24.03, 26.81, 28.27, 36.35, 36.53, 51.40, 51.51, 51.74, 52.35, 53.11, 53.47, 54.91, 55.05, 113.54, 115.51, 117.47, 118.58, 119.00, 119.43, 128.19, 128.45, 128.54, 129.15, 130.82, 141.09, 157.87, 158.09, 158.32, 158.54, 163.66, 164.60, 165.93. FTIR: 1733 cm⁻¹, 1669 cm⁻¹.

HR-MS (ESI): m/z calculated for $[C_{123}H_{183}N_{29}O_{36}-H]^-$: 2642.3333/found: 2642.3303; calculated for $[C_{123}H_{183}N_{29}O_{36}-2H]^{2-}$: 1320.6627/found: 1320.6633.

G1-4GdDO3A. The dendron **G1-4DO3A** (0.40g, 0.15 mmol, 1 equiv) was dissolved in distilled water (2.5 mL), and the pH of the corresponding solution was adjusted to 5.5 with 1 M solution of sodium hydroxide. Gadolinium chloride ($GdCl_3 \cdot 6H_2O$, 0.34 g, 0.91 mmol, 6 equiv) was dissolved in distilled water (10 mL) and added to the solution of dendron **G1-4DO3A**. The pH of the mixture was adjusted again to 5.5, and the reaction mixture was stirred at room temperature for 1 day. The solution was treated for 1 h with Chelex 100 resin to remove the excess of gadolinium ions (in **G1-4GdDO3A** sample, the absence of free gadolinium ions was checked as previously, by complexometry with xylenol orange as a dye).¹¹ The

resin was separated by centrifugation and the solution was concentrated under reduced pressure. The resulting crude product was dissolved in water (10 mL) and purified by dialysis (membrane with MWCO 500–1000 Da, VWR) against 1 L of distilled water (which was changed after 2.5, 6.5, 11.5, and 18 h). The dialyzed solution was freeze-dried leading to **G1-4GdDO3A** (0.12 g, 0.038 mmol, yield 25%). FTIR: 1600 cm⁻¹. HR-MS (ESI): m/z calculated for $[M - H]^{-1}$: 3259.9368/found: 3259.9368; calculated for $[M - 2H]^{2^{-1}}$: 1629.4645/found: 1629.4651 (values are given for the highest component of isotopic profile; $M \equiv C_{123}H_{171}Gd_4N_{29}O_{36}$). Detailed HRMS spectra with simulated isotopic distribution profiles are given in SI 5, Supporting Information.

Determination of Gd Concentration for Relaxometric Measurements. To determine Gd³⁺ concentration, either ICP-OES or relaxometric measurements were performed. Previously, the absence of free Gd³⁺ ion in the analyzed samples was checked by complexometry with xylenol orange as a dye.¹ Furthermore, each sample (200 μ L) was submitted to an acidic digestion (H₂O/HNO₃ 1:1 v/v, 200 μ L) to release Gd³⁺. For ICP-OES measurements, volumetric dilutions were carried out to achieve an appropriate Gd concentration within the working range of the method. Samples were analyzed using a Varian Liberty Series II spectrometer. Counts of Gd were correlated to a Gd calibration curve generated by mixing $Gd(NO_3)_3$ standard prepared under the same acidic conditions. For relaxometry measurements, the longitudinal relaxation rate of the digested acidic solution was then determined after subtraction of the diamagnetic contribution (0.35 s⁻¹ at 37 °C for the H₂O/HNO₃ 1:1 v/v solution). This value was compared to the one of a 1 mM GdCl₃ solution (11.76 s⁻¹), and after taking into account the dilution factor (2), the Gd^{3+} concentration was obtained for the sample of interest. Both methods gave similar results since for a solution of G1-4GdDOTAGA, a Gd concentration of 1.68 mM was found by ICP-OES while the same solution analyzed by relaxometry gave a concentration of 1.63 mM. Similar results were obtained for a G1-4GdDO3A solution, i.e. 1.51 mM and 1.54 mM in Gd by ICP-OES and relaxometry respectively (between 2 and 3% error between the two methods). For determination of r_1 relaxivities Gd concentration determined by relaxometry was used.

Relaxometric Measurements. T_1 Measurements at 20 MHz (0.47 T). T_1 measurements were performed in water on a Bruker mq20 minispec relaxometer (0.47 T) using an inversion recovery pulse sequence. The measurements were performed at nine different temperatures between 5 and 45 °C. The temperature was equilibrated before each scan. The inverse of the paramagnetic longitudinal relaxation time $(1/T_{1parar} s^{-1})$ of each sample was calculated according to

$$(1/T_{1para}) = (1/T_{1observed}) - (1/T_{1dia})$$

 $1/T_{1para}$ values were divided by Gd³⁺ concentration (mM) to determine G1-4GdDOTAGA and G1-4GdDO3A relaxivities r_1 (s⁻¹ mM⁻¹).

NMRD Profiles. ¹H NMRD profiles were measured on a Stelar Spinmaster FFC fast field cycling NMR relaxometer (Stelar, Mede, Pavia, Italy) over a range of magnetic fields extending from 0.24 mT to 0.7 T (which corresponds to ¹H Larmor frequencies from 0.01 to 30 MHz) using 0.5 mL samples in 7.5 mm o.d. tubes. The temperature was kept constant at 37 °C. Additional relaxation rates at 60 and 300 MHz were obtained with Bruker Minispec mq60 and Bruker Avance-300 MHz spectrometers, respectively (Bruker, Karlsruhe, Germany). The least-squares fitting of the ¹H NMRD data was performed using a homemade program (Fitting2000) and equations from the Solomon and Bloembergen (SBM) theory for the small gadolinium complexes or from the Lipari-Szabo theory for the slow rotating macromolecular contrast agents. In the fitting procedure, the $r_{\rm GdH}$ distance was fixed to 3.10 Å and the closest approach of the bulk water protons to the Gd³⁺ ion, d_{GdH} to 3.6 Å.¹² The diffusion constant *D* has been fixed to $3.0 \times 10^{-9} \text{ m}^2 \text{ s}^{-1}$,¹³ and the hydration number q was 1, as determined by luminescence lifetime measurements on the Eu³⁺ analogues (SI 6, Supporting Information). The water exchange rate $k_{ex} = 1/\tau_m$ where τ_m is the

Table 1. Best-Fit Parameters Obtained from the Fitting of the ¹H NMRD Profiles to the SBM Theory, Including the Lipari– Szabo Approach for Internal Flexibility^a

	$d (nm)^{b}$	$r (nm)^{b}$	$ au_M (\mathrm{ns})^c$	q^{b}	$ au_{SO}~(\mathrm{ps})$	$ au_{V} (\mathrm{ps})$	$ au_{RG}$ (ps)	$ au_{RL}$ (ps)	S^2
G1-4GdDOTAG	A 0.36	0.31	97	1	332 ± 11	31.1 ± 4.3	700 ± 112	150.0 ± 0.9	0.051 ± 0.015
G1-4GdDO3A	0.36	0.31	505	1	126 ± 3	37.4 ± 1.8	800 ± 94	167 ± 13	0.20 ± 0.02
^{<i>a</i>} The definition of	f the paramete	rs is given ir	the Experin	nental	Section. ^b Fixed	parameters. "	Value obtained fr	om ¹⁷ O experin	nents.

Scheme 2. Synthesis of G1-4NH₂ Dendron



residence time of the coordinated water molecule in the inner sphere, was fixed to the value obtained by independent ¹⁷O measurements of the transverse relaxation time of water $(1/T_2)$ according to the temperature. The fitting procedure allowed the determination of τ_{SO} , the gadolinium electronic relaxation time at zero field, τ_V , the correlation time that modulates the electronic relaxation, and τ_{RJ} the rotational correlation time (if the SBM theory is used), or τ_{RG} and τ_{RIJ} , *i.e.* the rotational correlation times describing the global and local motions, respectively, and S^2 that reflect the degree of spatial restriction of the local motion with respect to the global motion ($0 \leq S^2 \leq 1, S^2 = 0$ for isotropic internal motions, $S^2 = 1$ for completely restricted internal motions), if the Lipari–Szabo approach is used. The best-fit parameters obtained from the analysis of ¹H NMRD data are summarized in Table 1.

Temperature Dependent ¹⁷O NMR Measurements. The transverse ¹⁷O relaxation rates $(1/T_2)$ were measured in aqueous solutions of G1-4GdDOTAGA and G1-4GdDO3A (12 mM) in the temperature range 7-77 °C, on a Bruker Avance 500 (11.7 T, 67.8 MHz) spectrometer. The temperature was calculated according to a previous calibration with ethylene glycol and methanol. Proton decoupling was applied during all the acquisitions. Transverse relaxation times (T_2) were obtained by the measurement of the signal width at midheight. The data are presented as the reduced transverse relaxation rate $1/T_2^{R}$ = $55.55/([Gd_{complex}] \cdot q \cdot T_2^{P})$, where $[Gd_{complex}]$ is the molar concentration of the complex, q is the number of coordinated water molecules, and T_2^{P} is the paramagnetic transverse relaxation rate obtained after subtraction of the diamagnetic contribution from the observed relaxation rate. The treatment of the experimental data was performed as described elsewhere.¹⁴ The following parameters were determined: A/\hbar , the hyperfine coupling constant between the oxygen nucleus of the bound water molecule and the Gd³⁺ ion; τ_{vt} the correlation time modulating the electronic relaxation of Gd^{3+} ; E_{y} , the

activation energy related to τ_v ; *B*, related to the mean-square of the zero field splitting energy Δ ($B = 2.4\Delta^2$); and ΔH^{\ddagger} and ΔS^{\ddagger} the enthalpy and entropy of activation, respectively, of the water exchange process. The number of coordinated water molecules was set to 1.

T1-Weighted Images. MR imaging of G1-4GdDOTAGA and G1-4GdDO3A were performed using a 1.0 T MRI device (ICON, Bruker) using a T1-weighted spin–echo sequence (TR = 300 ms, TE = 12 ms, resolution = $208 \times 195 \ \mu$ m, slice thickness = 1.25 mm, 4 averages). Solutions of G1-4GdDOTAGA and G1-4GdDO3A were tested in the 0.1–1 mM range.

Transmetalation Experiments. The reactions of G1-4GdDOTA-GA and G1-4GdDO3A with Zn^{2+} ions were monitored by measuring the longitudinal relaxation rates R_1 (1/ T_1) of water protons on a Bruker mq20 minispec relaxometer (0.47 T), by the inversion recovery method. 12 μ L of a 2.5 × 10⁻¹ mol L⁻¹ solution of ZnCl₂ was added to 300 μ L of a phosphate buffered solution (pH 7) of the paramagnetic complex (2.5 × 10⁻³ mol L⁻¹). The temperature was equilibrated and maintained at 37 °C during the experiments. The mixture was stirred, and the measurements were started immediately and followed during 5 days. The $R_1^P(t)$ relaxation rate was obtained after subtraction of the diamagnetic contribution of the proton water relaxation (0.2826 s⁻¹) from the observed relaxation rate $R_1(t)$.

In Vitro Cytotoxicity Studies. NIH/3T3 (ATCC CRL-1658) cell viability was evaluated with an MTT assay (Sigma-Aldrich) in the presence of G1-4GdDOTAGA, G1-4GdDO3A, Na₂GdDOTAGA, and GdDO3ACOOH. NIH/3T3 cells were chosen as they are commonly used to screen the toxicity of chemicals¹⁵ and nanomaterials.¹⁶ They are well characterized, uniform, and known to provide reproducible results.¹⁷ NIH/3T3 cells were cultured in Dulbecco's Modified Eagle Medium GlutaMax supplemented with 2 mM L-glutamine, 100 U/mL penicillin, 100 μ g/mL streptomycin, and 10% fetal bovine serum and maintained at 37 °C under a humidified

Scheme 3. Synthesis of G1-4GdDOTAGA



atmosphere containing 5% CO₂. The cells were seeded in a 96 well plate (15.000 cells per well) and were allowed to settle overnight. The next day, **G1-4GdDOTAGA**, **G1-4GdDO3A**, **Na₂GdDOTAGA**, and **GdDO3ACOOH** were added on cells in the following concentrations 0.1, 0.3, 0.5, 0.75, and 1.05 mM of Gd and incubated for 24 h at 37 °C. The medium containing the dendrimers was then removed, and the MTT reagent (100 μ L, 1.5 mM in cell medium) was added to each well for 3 h at 37 °C. After taking off the reagents, DMSO was added under shaking for 1 h to dissolve the formazan crystals. The absorbance of each plate was measured at 595 nm with a plate spectrophotometer (PerkinElmer VICTOR³ 1420 Multilabel Counter).

RESULTS AND DISCUSSION

Preparation of G1-4GdDOTAGA and G1-4GdDO3A. *G1-4NH*₂ *Synthesis.* **G0** (Scheme 2, SI 1, Supporting Information) that will become the core of **G1** dendrimers was synthesized according to a procedure inspired from a published approach,^{18,19} where the primary amino groups of 3,3'-iminobis(propylamine) 1 were regioselectively protected by *tert*-butoxycarbonyl groups²⁰ to give amine 2 (88% yield). Amine 2 was further alkylated with benzylbromoacetate²⁰ (63% yield) to provide the core structure **G0** with Bocprotected terminal amine functions and a central carboxylic acid function protected by a benzyl (Bn) group. Hydrogenolysis²⁰ of **G0** gave acid 3 (98% yield) while acidolysis¹⁸ of G0 gave the bis-amine 4 (92% yield). Acid 3 and amine 4 were then coupled using 2-(1H-benzotriazol-1-yl)-1,1,3,3-tetrame-thyluronium hexafluorophosphate (TBTU) to give the pentavalent protected dendrimer G1 (28% yield after silica chromatography). Finally, deprotection of *tert*-butoxycarbonyl groups under acidic conditions afforded dendron G1-4NH₂ (35% yield).

G1-4GdDOTAGA Synthesis. G1-4GdDOTAGA synthesis was achieved from synthon G1-4NH₂ in the presence of Gd-DOTAGA 5 (Scheme 3). DOTAGA complexation with Gd₂O₃ afforded complex 5, and the excess of Gd³⁺ was removed by precipitation after NaOH addition. The peptidic coupling between 5 and G1-4NH₂ was then performed in the presence of 1-[bis(dimethylamino)methylene]-1H-1,2,3triazolo[4,5-*b*]pyridinium 3-oxid hexafluorophosphate (HATU). In this approach, the chelate was then metalated before being grafted to the dendron (premetalation strategy).

G1-4GdDOTAGA synthesis was confirmed by means of ESI-MS spectroscopy. The ESI-MS spectrum of **G1-4GdDOTAGA** clearly showed intense signals for molecular ions at $[M - H]^-$ and $[M - 2H]^{2-}/2$, with an isotopically resolved profile matching well with the corresponding simulated signals (Figure 1 and SI 3, Supporting Information).

Scheme 4. Synthesis of G1-4GdDO3A



Figure 2. ESI-HRMS spectrum of G1-4GdDO3A with $[M - H]^-$ (inset: signal at 3260 m/z) and $[M - 2H]^{2-}/2$ signals.

Therefore, **G1-4GdDOTAGA**, that exhibited 4 Gd^{3+} chelates/per molecule was obtained and readily isolated after ultrafiltration in 73% yield.

G1-4GdDO3A Synthesis. To obtain **G1-4GdDO3A**, a different approach, based on DO3A ligand grafting to **G1-4NH**₂, followed by insertion of Gd^{3+} into DO3A chelators of the corresponding dendrimeric ligand, was followed (post-metalation strategy). In a preliminary step, the well-known DO3A-tris-tBu-ester²¹ was alkylated with methyl 4-[(bromoacetyl)amino]benzoate. Then the methyl ester group was selectively saponified with lithium hydroxide²² and the corresponding benzylic acid was treated with EDC and NHS to afford the activated ester **6** (50% yield over 3 steps after a flash chromatography) (SI 4, Supporting Information). The ester **6** was then attached to **G1-4NH**₂ to provide tBuprotected **G1-4**[**DO3A**(tBu)₃] (Scheme 4). Trifluoroacetolysis of **G1-4**[**DO3A**(tBu)₃] gave **G1-4DO3A** dendron that was finally treated with GdCl₃ to afford **G1-4GdDO3A**

dendron (9% yield over 3 steps). Excess of Gd^{3+} was removed using a Chelex 100 resin followed by dialysis.

The ESI-MS spectrum of **G1-4GdDO3A** (Figure 2 and SI 5, Supporting Information) clearly showed an intense signal for bis-anionic molecular ion $[M - 2H]^{2-}/2$, with an isotopically resolved profile matching well with its corresponding simulated signals.

To have an insight of G1-4GdDOTAGA and G1-4GdDO3A hydrodynamic diameters, dynamic light scattering (DLS) was used (SI 7, Supporting Information). DLS analysis indicated Z-ave diameters of 1.70 and 2.50 nm for G1-4GdDOTAGA and G1-4GdDO3A, respectively. In parallel, DOSY experiments carried out on diamagnetic analogues (G1-4LaDOTAGA and G1-4LaDO3A) revealed hydrodynamic diameters of 2.08 and 1.75 nm, respectively (SI 7, Supporting Information). Given the uncertainties affecting each of these measurements, it can be concluded that the hydrodynamic diameters of G1-4GdDOTAGA and G1-4GdDO3A were inferior to 2.6 nm.



Figure 3. r₁ relaxivity at different temperatures (20 MHz) for (a) G1-4GdDOTAGA and (b) G1-4GdDO3A.

Finally, to determine the hydration number of Gd^{3+} in G1-4GdDOTAGA and G1-4GdDO3A, the luminescence lifetimes of the corresponding Eu³⁺ complexes in H₂O and D₂O were measured.²³ τ_{H2O} lifetimes were 0.649 and 0.523 ms for Na₂EuDOTAGA and EuDO3ACOOH, respectively, while τ_{D2O} lifetimes were 2.410 and 1.346 ms, giving a number of water molecules directly coordinated to Eu³⁺ of 1.18 for Na₂EuDOTAGA and 1.23 for EuDO3ACOOH *i.e.* very close to 1 (SI 6, Supporting Information). Assuming the fact that luminescence lifetimes are similar for mononuclear and multimeric complexes, one can conclude that each Gd³⁺ ion in G1-4GdDOTAGA and G1-4GdDO3A structures exhibited one coordinated water molecule in the inner sphere.

Relaxometric Properties of G1-4GdDOTAGA and G1-4GdDO3A. The proton relaxivity, which describes the MRI efficacy of chelates as contrast agents was first recorded at 20 MHz and 37 °C for both structures. On a per millimolar Gd basis, G1-4GdDOTAGA and G1-4GdDO3A presented r_1 relaxivities of 6.6 mM⁻¹ s⁻¹ and 8.1 mM⁻¹ s⁻¹, respectively. These values are significantly higher than relaxivities found for the corresponding mononuclear chelates Na₂GdDOTAGA and GdDO3ACO₂H (3.9 and 4.6 mM⁻¹ s⁻¹, respectively), as expected for a longer value of the rotational correlation time associated with the increased molecular size of G1-4GdDOTAGA and G1-4GdDO3A.⁸ In order to characterize the parameters that govern proton relaxivity, r_1 longitudinal relaxivity evolution with the temperature was recorded at 20 MHz for both complexes (Figure 3).

For G1-4GdDOTAGA, the continuous decrease of the relaxivity with increasing temperature is characteristic of a fast exchange of the coordinated water molecule in the inner sphere, which means in this case that water exchange will not limit the relaxivity. On the contrary, the observed evolution for the compound G1-4GdDO3A is typical of a slower exchange of the coordinated water molecule, limiting the relaxivity. In order to confirm these results and to have an insight into the microscopic parameters governing the dynamics of the system, variable-temperature ¹⁷O T_2 measurements were performed to have access to the water exchange rate k_{ex} (equal to the reciprocal lifetime of the water molecule in the inner sphere of the complex, $1/\tau_m$) for G1-4GdDO3A and G1-4GdDOTAGA (Figure 4).

The observed behavior was completely different for the two complexes and confirmed the results obtained by the evolution of the relaxivity with the temperature. Indeed, a decrease of the transverse relaxation rate was observed with increasing temperature on most of the temperature domain for G1-4GdDOTAGA, which is typical of a fast exchange of the coordinated water molecule. The opposite was observed for G1-4GdDO3A, indicating a slower exchange of the coordi-



Figure 4. $^{17}\mathrm{O}$ NMR measurements for (a) G1-4GdDOTAGA and (b) G1-4GdDO3A

nated water molecule. The experimental data were fitted with a theoretical model as described previously,¹⁴ which allowed extraction of τ_M values of 97 ± 4 ns ($k_{ex} = 1.03 \times 10^7 \text{ s}^{-1}$) and 505 ± 17 ns ($k_{ex} = 1.99 \times 10^6 \text{ s}^{-1}$) at 37 °C for G1-4GdDOTAGA and G1-4GdDO3A, respectively, as well as the thermodynamic parameters governing the exchange (SI 8, Supporting Information). The lower value of k_{ex} obtained for G1-4GdDO3A is consistent with data from the literature,²⁴ where a similar Gd-complex was grafted on a polysaccharide derived from inulin, giving a similar k_{ex} of 1.6 × 10⁶ s⁻¹. This difference can be correlated with the structure of the two complexes and more precisely to the functional groups directly coordinated to the Gd center. It is indeed well-known that the presence of an amide group, as in G1-4GdDO3A, slows down the exchange of the coordinated water molecule.²⁵

Nuclear magnetic relaxation dispersion (NMRD) profiles were also recorded in the field range 0.01–300 MHz for all the complexes.

The **G1-4GdDOTAGA** NMRD profile (Figure 5a) was analyzed using the classical inner sphere and outer sphere theories, which led to a rotational correlation time $\tau_{\rm R}$ of 147.0 \pm 2.0 ps (SI 9, Supporting Information), *i.e.* almost two times higher than $\tau_{\rm R}$ of GdDOTAGA (65.1 \pm 1.9 ps). This NMRD profile is nevertheless characteristic of a system in a fast rotational motion, which is in relative contradiction with its high molecular weight. When the Lipari–Szabo treatment was incorporated in the fit (Table 1), the order parameter S², which describes the degree of spatial restriction of the local rotational motion of the Gd³⁺-waterproton axis with respect to the global system motion, was almost 0 (S² = 0.05), *i.e.* typical of isotropic internal motions. This explains the low value of $\tau_{\rm R}$ extracted from the fitting with the SBM theory.

The **G1-4GdDO3A** NMRD profile (Figure 5b) was analyzed in a similar way, which led to a rotational correlation time $\tau_{\rm R}$ of 242.0 \pm 10.0 ps (SI 9, Supporting Information), *i.e.* almost three times higher than $\tau_{\rm R}$ of **GdDO3ACO**₂H (71.2 \pm 0.8 ps). The profile exhibited a hump at medium field which is typical of slowly rotating species. When the Lipari–Szabo

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Figure 5. NMRD profiles at 37 $^{\circ}$ C for (a) G1-4GdDOTAGA and (b) G1-4GdDO3A (the line represents the least-square fit to the experimental data point as described in the text).



Figure 6. T₁-weighted images of G1-4GdDOTAGA and G1-4GdDO3A with DOTAREM and water as controls. Samples imaged at 1 T, 37 °C, and with a standard spin echo (SE) sequence.

treatment was incorporated in the fit, the order parameter S^2 was 0.20, which indicated a higher rigidity of this system by comparison to **G1-4GdDOTAGA** (Table 1). This could be correlated with the presence in **G1-4GdDO3A** of the short and rigid aromatic linker,^{8e} which was a positive feature to limit the rotation of the small Gd³⁺ chelates. This has thus a positive impact on the relaxivity of this compound, especially at medium field where a hump is observed on the NMRD profile.

To demonstrate how relaxivity enhancements were translated into contrast, T_1 -weighted images of phantoms containing solutions of **G1-4GdDOTAGA** and **G1-4GdDO3A** were acquired at 1T with DOTAREM as the control (Figure 6).

For both systems, the bright signal enhancement progressively increased with Gd concentration. Furthermore, for each Gd concentration, G1-4GdDO3A or G1-4GdDOTAGA signals were brighter than those of the control, corroborating the relaxometric results.

Finally, relaxivity studies of G1-4GdDOTAGA and G1-4GdDO3A in the presence of human serum albumin (HSA) were performed (Table 2).

G1-4GdDOTAGA demonstrated a very small increase of relaxivity (9%) which could be related to an increase of the

Table 2. G1-4GdDOTAGA and G1-4GdDO3A (1 mM, 37 °C, 20 MHz) Relaxivities in the Presence of HSA (4%)

	$r_1 (s^{-1} \text{ mM}^{-1}, \text{H}_2\text{O})$	$r_1 (s^{-1} \text{ mM}^{-1}, \text{HSA})$	$r_{1_{\mathrm{H_2O}}}^{\mathrm{HSA}}/r_1$
G1-4GdDOTAGA	6.90	7.50	1.09
G1-4GdDO3A	8.96	13.37	1.49

medium viscosity. For G1-4GdDO3A a more significant difference (49%) was observed, but it is still not high enough to conclude to a noncovalent association to HSA. This should thus allow a fast elimination of the compounds from the body, probably *via* glomerular filtration, given the estimated hydrodynamic diameter of both structures (SI 7, Supporting Information).

Kinetic Inertness and Cytotoxicity Evaluations of G1-4GdDOTAGA and G1-4GdDO3A. Before testing G1-4GdDOTAGA and G1-4GdDO3A safety against cells, their kinetic inertness was evaluated by relaxometry in the presence of Zn^{2+} in a phosphate-buffered solution.²⁶

If the transmetalation reaction occurs, it will result in Gd^{3+} release which then will precipitate into $GdPO_4$. Consequently, it will lead to a subsequent decrease of the proton paramagnetic relaxation rate (R_1^p) .²⁶ **G1-4GdDOTAGA** and

G1-4GdDO3A R_1^p evolutions gave therefore a good indication of potential transmetalation. Figure 7 showed the evolution of their normalized paramagnetic relaxation rates $R_1^p(t)/R_1^p(0)$.



Figure 7. Evolution of $R_1^{p}(t)/R_1^{p}(0)$ vs time (at 20 MHz, 37 °C) for (a) **G1-4GdDOTAGA** and (b) **G1-4GdDO3A** in the presence of Zn^{2+} in phosphate buffer (pH = 7)

By comparison to the initial relaxation rate $R_1^{p}(0)$, no significant changes were observed over a 4 days period for both dendrons, indicating that for **G1-4GdDOTAGA** and **G1-4GdDO3A** no Gd³⁺ release occurred.

G1-4GdDOTAGA and G1-4GdDO3A in vitro safety was then tested against NIH/3T3 cells (Figure 8).



Figure 8. Metabolic activity of fibroblasts (NIH/3T3) incubated with (a) G1-4GdDOTAGA and (b) G1-4GdDO3A for 24 h (Na₂GdDOTAGA and GdDO3ACOOH being used respectively as controls).

For G1-4GdDO3A no significant decrease in metabolic activity was observed up to a Gd concentration of 1.05 mM (>90% viability for both), and this behavior is similar to that of control. However, a slight and progressive decrease was observed with increasing concentrations of G1-4GdDOTAGA to reach 75% at a Gd concentration of 1.05 mM. This follows the same trend as for Na₂GdDOTAGA alone, which likely means the decreasing in metabolic activity cannot be totally attributed to the dendrimer. Na₂GdDOTAGA is ionic while GdDO3ACOOH is neutral which implies differences in osmolarity that can likely explain these discrepancies in toxicity between the two compounds.

CONCLUSION

Two G1-4GdCA assemblies containing GdDOTAGA and GdDO3A units were prepared and subjected to a ¹H and ¹⁷O NMR relaxometric investigation. These compounds have similar mass and molecular sizes but differ by the linkers used to graft the chelates to the organic framework. The multimeric nature of G1-4GdCA assemblies allowed an increase in molecular size which was translated into a corresponding increase of $\tau_{\rm R}$ and hence of relaxivity (between

70 and 76% over the monomer units). It should be emphasized that the two multimeric systems were characterized by exchange regimes of water molecule not limiting the relaxivity for **G1-4GdDOTAGA** while limiting for **G1-4GdDO3A**, in agreement with the presence of amide groups at the vicinity of the lanthanide ion. Nevertheless, **G1-4GdDOTAGA** exhibited an NMRD profile typical of a system in a fast rotational motion while the **G1-4GdDO3A** NMRD profile was characteristic of a system in a slower rotational motion. The analysis of NMRD profiles by the Lipari–Szabo theory showed for **G1-4GdDOTAGA** a significant limiting role of the local rotational motions of the chelates which resulted in an effective value of $\tau_{\rm R}$ lower than that expected based on molecular dimensions.

On the other hand, the S^2 value determined for G1-4GdDO3A suggested a more restricted motion of the chelates located at the periphery of the assembly. This local rigidity was induced by the aromatic linker used to hook the chelate on the organic framework. Therefore, G1-4GdDO3A that still exhibits a derivatizable site is a good candidate for the design of targeting paramagnetic probes for molecular imaging or multimodal probes that combine at least two different imaging techniques. Further works are in progress in these two directions.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.inorg-chem.9b01747.

Experimental details of new compounds' synthesis and analytical data (HR-ESI-MS mass spectra, luminescence lifetime measurements, hydrodynamic diameters, and fitted parameters of 17 O and NMRD data) (PDF)

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Notes

The authors declare no competing financial interest.

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