1	Gold Nanoparticles Functionalised with Fast Water Exchanging Gd ³⁺ Chelates:
2	Linker Effects on the Relaxivity
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4	Miguel F. Ferreira, ^a Janaina Gonçalves, ^a B. Mousavi, ^b M. I. M. Prata, ^c S. P. J.
5	Rodrigues, ^d Daniel Calle, ^e Pilar López-Larrubia, ^e Sebastian Cerdan, ^e Tiago B.
6	Rodrigues, fg Paula M. Ferreira, L. Helm, S José A. Martins and Carlos F. G. C.
7	Geraldes ^h
8	
9	^a Centro de Química, Campus de Gualtar, Universidade do Minho, 4710-057 Braga,
10	Portugal. E-mail: jmartins@quimica.uminho.pt
11	^b Laboratoire de Chimie Inorganique et Bioinorganique, Ecole Polytechnique Fédérale
12	de Lausanne, EPFL-BCH CH-1015 Lausanne, Switzerland. E-mail:
13	lothar.helm@epfl.ch; Fax: +41 (0)21 693 98 95;Tel: +41 (0)21 693 98 76
14	^c IBILI and ICNAS, Universidade de Coimbra, Coimbra, Portugal.
15	^d Chemistry Center and Department of Chemistry, University of Coimbra, 3004-535,
16	Coimbra, Portugal.
17	^e Instituto de Investigaciones Biomédicas "Alberto Sols", CSIC-UAM, Madrid, Spain
18	^f Cancer Research UK Cambridge Research Institute, Li KaShing Centre, Cambridge
19	CB2 0RE, United Kingdom.
20	^g Department of Biochemistry, University of Cambridge, Cambridge CB2 1GA, United
21	Kingdom.
22	^h Chemistry Center and Department of Life Sciences, Faculty of Science and
23	Technology, University of Coimbra, Calçada Martim de Freitas, 3000-393 Coimbra,
24	Portugal.
25	
26	Corresponding authors:
27	José A. Martins, Centro de Química, Campus de Gualtar, Universidade do Minho,
28	4710-057 Braga, Portugal.
29	Lothar Helm, École Polytechnique Féderale de Lausanne, EPFL-BCH CH-1015
30	Lausanne, Switzerland.

- 1 Keywords: Gold nanoparticles; Gd^{3+} chelates; Contrast Agents; MRI; linkers; ω -thiol
- 2 functionalized DO3A-N-(α-amido)propionate chelators; fast water exchange; stability;
- 3 biodistribution; *in vivo* MRI.

Abstract

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The relaxivity displayed by Gd³⁺ chelates immobilized onto gold nanoparticles is the result of complex interplay between nanoparticle size, water exchange rate and chelate structure. In this work we study the effect of the length of ω-thioalkyl linkers, anchoring fast water exchanging Gd³⁺ chelates onto gold nanoparticles, on the relaxivity of the immobilized chelates. Gold nanoparticles functionalized with Gd3+ chelates of mercaptoundecanoyl and lipoyl amide conjugates of the DO3A-N-(α-amino)propionate chelator were prepared and studied as potential CA for MRI. High relaxivities per chelate, of the order of magnitude 28-38 mM⁻¹s⁻¹ (30 MHz, 25 °C) were attained thanks to simultaneous optimization of the rotational correlation time and of the water exchange rate. Fast local rotational motions of the immobilized chelates around connecting linkers (internal flexibility) still limit the attainable relaxivity. The degree of internal flexibility of the immobilized chelates seems not to be correlated with the length of the connecting linkers. Biodistribution and MRI studies in mice suggest that the in vivo behavior of the gold nanoparticles is determined mainly by size. Small nanoparticles (HD= 3.9 nm) undergo fast renal clearance and avoidance of the RES organs while larger nanoparticles (HD= 4.8 nm) undergo predominantly hepatobiliary excretion. High relaxivities, allied to chelate and nanoparticle stability and fast renal clearance in vivo suggests that functionalized gold nanoparticles hold great potential for further investigation as MRI Contrast Agents. This study contributes to understand the effect of linker length on the relaxivity of gold nanoparticles functionalized with Gd³⁺ complexes. It is a relevant contribution towards "design rules" for nanostructures functionalized with Gd³⁺ chelates as Contrast Agents for MRI and multimodal imaging.

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Introduction

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MRI is becoming the "central imaging modality" in clinical diagnostic. MRI is based 2 on the nuclear magnetic resonance phenomenon (NMR). In MRI scans, essentially, the 3 4 relaxation times (T_1 and T_2) of the water protons of tissues (intrinsically different) are acquired and reconstructed into tridimensional anatomical images.^{2,3} MRI is inherently 5 non-invasive, makes use of (benign) non-ionizing radiation (static and radiofrequency 6 magnetic fields), is depth independent and displays superb spatial resolution. Low 7 detection sensitivity (inherent to the NMR phenomenon) is the main limitation of MRI.⁴ 8 Contrast Agents (CA) are paramagnetic species (Gd³⁺, Mn²⁺, Fe³⁺, stable organic 9 radicals, iron oxide nanoparticles, etc.) that by promoting selective reduction of T_1 or T_2 10 of the water protons of tissues can generate dramatic contrast enhancements.^{5,6} The 11 selective enhancement of the relaxation rates, $R_{L,2}$, $(R_{L,2} = 1/T_{L,2})$, normalized to 1 mM 12 concentration of paramagnetic centres - relaxivity $(r_{1,2}, \text{ units mM}^{-1} \text{ s}^{-1})$, measures CA 13 efficacy.^{4,7} Approved CA for clinical MRI are either Gd³⁺ complexes with linear 14 (DTPA-type) and macrocyclic (DOTA-type) poly(aminocarboxylate) ligands (T_I -15 weighed MRI)⁸ or iron oxide nanoparticles (IONPS) stabilized with dextran $(T_2-T_2*$ -16 weighed MRI). Low molecular weight Gd³⁺-based CA display relaxivities of the order 17 of magnitude 3-5 mM⁻¹ s⁻¹ at magnetic fields relevant (currently) for clinical MRI (20-18 120 MHz). The Solomon-Bloembergen-Morgan (SBM) theory predicts that very high 19 relaxivities, of the order of magnitude 100 mM⁻¹ s⁻¹ at magnetic fields relevant for 20 clinical imaging (20-120 MHz), are attainable by Gd³⁺ chelates displaying simultaneous 21 optimization of the main parameters that govern relaxivity: rotational correlation times 22 (τ_R) , water exchange rate constant (k_{ex}) and electron relaxation parameters $(\tau_v \text{ and } \Delta^2)$. 23 Despite great advancements in the design and synthesis of CA during the past two 24 25 decades, the ideal CA- displaying very high relaxivity and safety in vivo, targeting capability and responsiveness coupled to the rapeutic properties, is still elusive. 10 26 The "nanotechnology revolution" is underway with a dramatic impact in many 27 fields, particularly in medical imaging. 11 Gold nanostructures (nanoparticles, 28 29 nanoclusters, nanorods, etc) are finding many applications in chemistry, medicine, biotechnology and other fields, owing to intrinsic reporting properties (localized 30 surface plasmon resonance, fluorescence, X-Ray attenuation)¹² coupled to therapeutic 31 properties (hyperthermia, X-ray sensitization), ¹³ biocompatibility and safety in vivo ¹⁴ 32 and facile preparation with tunable size and surface properties by bottom-up 33

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methodologies. 15 The first generation of gold nanoparticles (AuNPs) CA made use of, easy to synthesize, thiol-functionalised Gd(DTPA-bis-amide)¹⁶⁻¹⁸ and Gd(DO3A)-type chelates. 19 Superb relaxivities (per nanoparticle) were attained thanks to chelate clustering. 16 In vivo MRI studies established the merits of AuNPs as CA for MRI, bimodal MRI/X-ray imaging and as theragnostics (MRI/X-ray sensitization). 16-18,20 Slow water exchange and fast local rotational motions of the immobilized chelates around linkers/spacers (chelate flexibility) result in relaxivity enhancements (per chelate) lower than those expected for Gd³⁺ chelates appended to rigid nanosized objects. 16-18,21,22 Helm and co-workers reported very high relaxivity per chelate immobilized onto AuNPs (60 mM⁻¹ s⁻¹; 30 MHz, 25 °C), attributed to two exchanging inner sphere water molecules in Gd(DTTA)-type chelates and complete rigidity of the chelates immobilized via a short aromatic linker.²³ The relaxivity was however, still limited by slow water exchange. We have demonstrated in previous studies that the Gd[DO3A-N-(α-amino)propionate] chelate and Gd³⁺ complexes of amide conjugates of the DO3A-N-(α-amino)propionate chelator display water exchange rates within the range considered ideal for attaining high relaxivities at intermediate fields, thanks to "steric compression around the water binding site". 24,25,26 AuNPs functionalized with the fast water exchanging chelate Gd[DO3A-N-(α-cystamido)propionate] display high relaxivities at intermediate and high fields (27 and 8.0 mM⁻¹ s⁻¹; 20 and 200 MHz, respectively, 25 °C) as the result of simultaneous optimization of the rotational dynamics and water exchange rate.²⁷ Fast local rotational motions around the cysteine linker still limit the attainable relaxivity, as demonstrated before for other macromolecular/nanosized objects such as micelles, dendrimers, polymers, etc. 26,27,28 In this work we address the effect of the length of the ω-thioalkyl linker, anchoring fast water exchanging Gd[DO3A-N-(α -amido)propionate] chelates to gold nanoparticles, on the relaxivity. Biodistribution and in vivo MRI studies with the functionalized AuNPs as CA are reported also.

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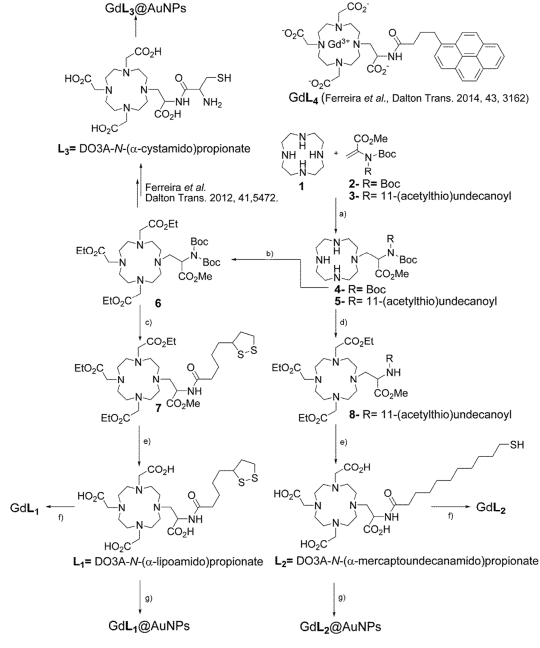
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Synthesis and characterization

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- 3 Mercaptoundecanoyl and lipoyl conjugates of the DO3A-N-(α -amino)propionate
- 4 chelator were synthesized to study the effect of the length of the ω-thioalkyl linker on
- 5 the relaxivity of AuNPs functionalized with Gd³⁺ chelates (Scheme1).



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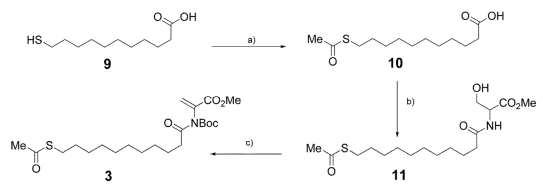
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Scheme 1. Synthetic pathway for ω-thioalkyl conjugates of the DO3A-N-(α-amino)propionate chelator: a) $K_2CO_3/MeCN$; b) ethyl bromoacetate, $K_2CO_3/MeCN$; c) i. TFA/CH₂Cl₂, ii. DIPEA/CH₂Cl₂, lipoic acid, DCC/HOBt; d) i) TFA/DCM, ii. Ethyl bromoacetate, $K_2CO_3/MeCN$; e) i. NaOH aq., ii. Flash chromatography silica gel, iii. SEM (Sephadex G10); f) GdCl₃.6H₂O; g) i. HAuCl₄, NaBH₄, ii. GdCl₃.6H₂O, iii. SEM (Sephadex G10) followed by dialysis (cellulose tubing MWCO 10 KDa).

The lipoic acid conjugate (**L**₁) was prepared following the synthetic methodology reported before for the cysteine conjugate of the DO3A-*N*-(α-amino)propionate chelator (**L**₃).²⁷ The synthetic pathway excludes, all along, acidic conditions likely to promote oligomerization of the chelator through the lipoic acid moiety.²⁹ Deprotection of the fully alkylated orthogonally protected intermediate **6** allows direct conjugation of lipoic acid to the preformed DO3A-*N*-α-(amino)propionate scaffold.²⁷ For preparing the 11-mercaptoundecanoyl conjugate (**L**₂) the preformed amide was introduced into the *cyclen* scaffold *via* Michael addition of the *N*-Boc,*N*-(11-(acetylthio)undecanoyl)dehydroalanine methyl ester electrophile (**3**).^{25,26} Reactive block **3** was prepared over 3 steps in 48 % overall yield (Scheme 2).³⁰



Scheme 2. Synthetic route for Michael electrophile *N*-Boc,*N*-(11-(acetylthio)undecanoyl)dehydroalanine methyl ester (3): a) acetic anhydride/pyridine; b) serine methyl ester hydrochloride, HOBt/DCC/NEt₃; c) Boc₂O/DMAP, dry acetonitrile.

The thioacetyl protecting group proved easy to install and stable under mild alkaline and strong acidic conditions *en route* to L_2 . Final deprotection was performed in one step by saponification with ethanolic NaOH. Following pH adjustment to neutrality with diluted hydrochloric acid, chelators L_1 and L_2 were adsorbed onto silica and purified by flash chromatography followed by Size Exclusion Chromatography (SEC) on Sepahdex G10 with water elution.

Relaxometric studies of GdL₁ and GdL₂

- 4 The concentration dependence of the paramagnetic longitudinal water proton relaxation
- rate (R_{Ip}) was measured for GdL_1 and GdL_2 (20 MHz, 25 °C, pH 7.1) (Figure 1).

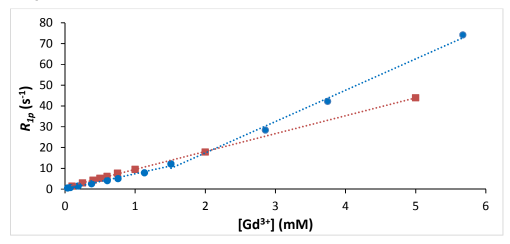


Figure 1. Concentration dependence of the paramagnetic water proton relaxation rate $R_{Ip} = (R_{lobs} - R_{Id})$ for GdL_1 (\blacksquare) and GdL_2 (\bullet) (20 MHz, 25 °C, pH 7.1).

The relaxation rate data for GdL_1 can be well fitted to a straight line (Equation 1) affording a relaxivity of 8.6 ± 0.9 mM⁻¹ s⁻¹ (20 MHz, 25 °C, pH 7.1), characteristic of low/intermediate molecular weight chelates in fast rotation in solution. Fitting the relaxation rate data of GdL_2 requires two straight lines with different slopes (Equation 1 and 2). The concentration at the interception of the two lines defines the critical micelle concentration- cmc ($cmc = 1.5 \pm 0.3$ mM).³¹

$$R_{1p} = R_1^{obs} - R_1^d = r_1^{na} \times c_{Gd}$$
 Eq. 1
 $R_{1p} = R_1^{obs} - R_1^d = (r_1^{na} - r_1^a) \text{cmc} + r_1^a \times c_{Gd}$ Eq. 2

 R_I^{obs} is the longitudinal relaxation rate measured for the solution, R_I^d is the diamagnetic contribution to the longitudinal relaxation rate (the relaxation rate of pure water) and C_{Gd} is the analytical Gd^{3+} concentration.

Below the *cmc* GdL₂ is present in solution as monomers (non-aggregated), displaying a relaxivity (r_I^{na} = 6.6 ± 0.1 mM⁻¹ s⁻¹; 20 MHz, 25 °C, pH 7.1) characteristic of low molecular weight chelates (Equation 1).

1 For GdL₂ at concentrations above the *cmc* the relaxation rate has a contribution 2 from monomers (at a concentration equal to the *cmc* value) and from the (aggregated) micellar form $(r_I^a = 15.1 \pm 0.8 \text{ mM}^{-1} \text{ s}^{-1}; 20 \text{ MHz}, 25 ^{\circ}\text{C}, \text{ pH } 7.1)$ (Eq. 2). Self-association 3 4 of GdL₂ into micelle-type structures leads to an increase of the effective molecular volume of the chelate. Slow tumbling in solution (longer rotational correlation times) 5 translates into substantially higher relaxivity for the aggregated form of GdL₂ 6 7 comparing to its monomeric (non-aggregated) form. The relaxivity enhancement for GdL₂ upon self-assembly is of the same order of magnitude as that reported for the 8 DOTA-type Gd(DOTASA-C12) chelate functionalized with a C_{12} alkyl chain ($r_I^a = 18.0$ 9 mM⁻¹ s⁻¹, 20 MHz, 25 °C)³², but significantly lower than that attained by the aggregated 10 form of the fast water exchanging Gd[DO3A-N-(α-pyrenebutanamido)propionate] 11 chelate (GdL₄ in Scheme 1) ($r_1^a = 32 \text{ mM}^{-1} \text{ s}^{-1}$; 20 MHz; 25 °C).²⁶ The temperature 12 13 dependence of the water proton longitudinal relaxation rate for GdL₁ and GdL₂ (20 MHz, 25 °C) (Figure SI1) indicates that the relaxivity is not limited by slow water 14 exchange, as demonstrated before for other Gd3+ chelates of amide conjugates of the 15 DO3A-N-(α -amino)propionate chelator. ^{24,27} As both GdL_2 and $Gd[DO3A-N-(\alpha-$ 16 pyrenebutanamido)propionate] chelates display fast water exchange, the lower 17 relaxivity attained by GdL₂ has to be ascribed to higher internal flexibility and/or 18 smaller size of the GdL₂ micelles (Figure SI2). The pH dependence of the proton 19 relaxation rate (Figure SI3) and the transmetallation study (Figure SI4) show that GdL₁ 20 21 and GdL₂ are stable in the physiological pH range and kinetically inert towards transmetallation against Zn²⁺.³³ 22

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Preparation of gold nanoparticles functionalized with GdL₁ and GdL₂ chelates

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A modified Brust's methodology in aqueous solution was employed for preparing 26 AuNPs functionalized with GdL₁ and GdL₂ chelates.^{27,34} Using directly the GdL₁ and 27 GdL₂ chelates as nanoparticle stabilizers resulted in extensive precipitation upon 28 29 addition of the reducing agent (NaBH₄). Attempts to functionalize citrate-stabilized AuNPs with GdL₁ and GdL₂ chelates via place exchange revealed also unsuccessful. 18 30 A two-step methodology, using the L_1 and L_2 chelators as NPs stabilizers, followed by 31 Gd3+ complexation, revealed successful for preparing AuNPs functionalized with GdL1 32 and GdL₂ chelates (Scheme 1).²⁷ 33

Solutions containing equimolar amounts of L_1 or L_2 and HAuCl₄ turned immediately dark brown upon addition, in one aliquot, of one molar equivalent of NaBH₄. Adding a molar equivalent of Gd^{3+} , in relation to the total amount of L_1 or L_2 in the crude mixture, resulted in stable NPs. Size Exclusion Chromatography (SEC) (Sephadex G10, elution with water) followed by extensive dialysis against water (cellulose tubing MWCO 10 000), afforded stable AuNPs functionalized with GdL_1 and GdL_2 chelates. A single fraction, including the broad colored band eluting on SEC, was collected. The absence of (free) uncomplexed Gd^{3+} was confirmed by the xylenol orange test.³⁵

Characterization of GdL₁@AuNPs and GdL₂@AuNPs preparations

(DLS) (Figure 2) and Transmission Electron Microscopy (TEM) (Figure 3).

The Gd content of the functionalized AuNPs was estimated by bulk magnetic susceptibility measurements³⁶ and further confirmed by ICP-OES following sample digestion with *aqua regia* (HCl/HNO₃; 3/1 v/v) (Table 1).^{27,37}
The AuNPs were characterized regarding size distribution by Dynamic Light Scattering

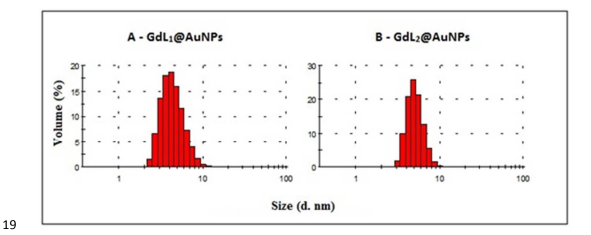


Figure 2. Size distribution (% volume) for GdL₁@AuNPs (A) and GdL₂@AuNPs (B).

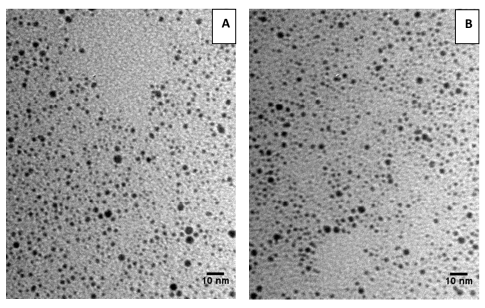


Figure 3. Transmission Electron Microscopy (TEM) for GdL₁@AuNPs (A) and GdL₂@AuNPs (B).

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DLS measurements give the hydrodynamic diameter (HD) of NPs, including the Au nanocrystal core, the chelate monolayer and the immobile ionic layer surrounding the NPs. An average HD of 4.8 nm (NPs distribution in the range 3-10 nm with a maximum at 3-4 nm) and 5.9 nm (NPs distribution in the range 2-11 nm with a maximum at 4-5 nm), was measured for GdL₁@AuNps and GdL₂@AuNPs, respectively. TEM measurements reveal only the nanocrystal core. The TEM images obtained for GdL₁@AuNPs and GdL₂@AuNPs do not allow to determine the average diameter of the Au core owing to the very small size of the NPs. From the TEM images one can only conclude that most GdL_1 @AuNPs and GdL_2 @AuNPs have a nanocrystal core with a diameter under 2 nm. The average diameter of the Au core of GdL₁@AuNPs and GdL₂@AuNPs (~ 1.0 and 0.9 nm, respectively) was estimated from the DLS measurements taking into account the thickness of the chelate monolayer, calculated by PM6 semi-empirical calculations for the most likely conformations of GdL₁ and GdL₂ bonded to one or two Au atoms (1.9 and 2.5 nm, respectively) (Figure SI5, Table SI2). 23,24 The absence of a well-defined plasmon absorption band in the UV-Vis spectra of GdL₁@AuNPs and GdL₂@AuNPs (Figure SI6 and SI7), corroborates the very small size of the NPs core.³⁸

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1 Table 1. Characterization of GdL₁@AuNPs and GdL₂@AuNPs

	GdL ₁ @AuNPs ^a	GdL ₂ @AuNPs ^a	GdL ₃ @AuNPs ^b
[Gd] (mM) ^c	0.57	1.30	1.24
HD (nm) ^d	4.8	5.9	3.9
Zeta potential (mV)	-6.3	-13,7	-12.3
\mathbf{D}_{Au}^{e}	1.0	0.9	0.7^f

^aThis work; ^bRef ²⁷; ^cDetermined by ICP-OES; ^dDLS measurements; ^eEstimated from the HD and semi-empirical calculations of the chelate monolayer thicknes- see Table SI2; ^fRevised value according to semi-empirical calculations for the length of GdL₃.

 As L_1 , L_2 and L_3 share the same coordination cage, the length of the linker defines the overall wedge-like geometry of the chelator. Shorter linkers originate bulkier thiol ligands. Ligand bulkiness increases in the series $L_3>L_1>L_2$ (Figure SI5). Bulkier thiols are likely to terminate the growth of AuNPs earlier than less bulky ligands, resulting in AuNPs displaying smaller Au cores associated to higher surface curvature.³⁹ This correlation ($D_{Au} = 1.0$, 0.9 and 0.7 nm, for Gd L_1 , Gd L_2 and Gd L_3 @AuNPs) is followed roughly by L_2 and L_3 . The discrepancy observed for L_1 can be due to the different sulfur binding mode.

 $GdL_1@AuNPs$ and $GdL_2@AuNPs$ were found to be stable in solution for extended periods. The NPs could be freeze-dried and re-dissolved without aggregation/precipitation. This can be ascribed to the overall negative charge (-1) of the immobilized Gd^{3+} complexes, resulting in NPs displaying negative zeta-potential (Figure SI8).

Relaxometric characterization of GdL₁@AuNPs and GdL₂@AuNPs

The concentration dependence of the proton longitudinal relaxation rate (R_{Ip}) was evaluated for GdL₁@AuNPs and GdL₂@AuNPs ($r_I = 29$ and 38 mM⁻¹ s⁻¹, respectively, 20 MHz, 25 °C, pH 7.1) (Figure SI9).

For relevant clinical applications chelates immobilized onto NPs must be stable regarding demetallation and inert towards transmetallation with physiological metal ions, mainly Zn^{2+,33} In addition to releasing toxic Gd³⁺, demetallation and transmetallation processes of immobilized chelates are likely to trigger particle aggregation and precipitation *in vivo*. Stability at low pH is particularly important as protonation-assisted mechanisms have been implicated in demetallation, presumably

followed by transmetallation with serum ions, of macrocyclic Gd(DOTA)-type chelates. ^{33,40,41} The pH dependence of the protonic relaxation rate (R_{Ip}) was evaluated for GdL₁@AuNPs and GdL₂@AuNPs in the pH range 3-10 (Figure SI10).

The kinetic stability of the immobilized chelates (and entire nanoparticles) was evaluated by challenging $GdL_1@AuNPs$ and $GdL_2@AuNPs$ with Zn^{2+} ions in phosphate buffer (Figure SI11 and Figure SI12).³³ According to the criteria set by Muller and co-workers, the immobilized chelates (and whole NPs) can be classified as kinetically inert and thermodynamically stable.³³ The pH stability and kinetic inertness indicate that the NPs are potentially safe for *in vivo* applications.

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Nuclear Magnetic Relaxation Dispersion Profiles

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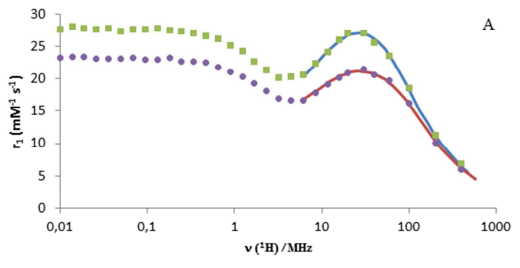
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The magnetic field dependence of the proton relaxivity (r_l) (Nuclear Magnetic Relaxation Dispersion - NMRD profiles) was obtained for GdL₁@AuNPs and GdL₂@AuNPs in the *Larmor* frequency range 0.01-400 MHz. The most important parameters that govern relaxivity are the hydration number (q), the water exchange rate constant $(k_{\rm ex})$, the rotational correlation time $(\tau_{\rm R})$ and the electron relaxation parameters $(\tau_v \text{ and } \Delta^2)^{4,7}$. The number of water molecules in the first coordination sphere, the water exchange rate and the rotational correlation time can be tuned by chelate design. Clear rules to tune the electron relaxation parameters are still elusive. 42 Treating chelates immobilised onto macromolecular/nanosized objects (micelles, proteins, polymers, dendrimers, nanoparticles, viral particles) as rigid entities, often fails to deliver reliable parameters from the fitting of the NMRD profiles to the SBM theory. In fact, it is necessary to assume in the fittings that the interactions that generate the relaxation are influenced by both fast local rotational motions (τ_{Rlocal}) of the immobilized chelates around linkers/spacers and a slower, global motion, common to the entire object $(\tau_{Rglobal})$. The degree of spatial restriction of the local motion (interpreted as chelate flexibility), is measured by the generalized, model independent order parameter- S^2 . The order parameter can assume values in the range 0-1: $S^2 = 0$ if the internal motions are isotropic, $S^2 = 1$ if the internal motions are completely restricted.⁴³

In this work, it was assumed in the fittings that the immobilised GdL_1 and GdL_2 complexes have one inner sphere water molecule (q = 1) like other Gd^{3+} complexes of the $DO_3A-N-(\alpha-amino/amido)$ propionate family. ²⁴⁻²⁶ The water exchange rate constant

and its activation enthalpy $(k_{\rm ex}^{298}, \Delta H^{\ddagger})$ were fixed to values determined for the analogous Gd[(DO3A-*N*-(α -benzoylamido)propionate] chelate.²⁵

 The fittings (continuous lines in Figure 4) are restricted to frequencies above 6 MHz as the SBM theory is not suitable for describing the rotational dynamics of slow-rotating objects at low magnetic fields. The best fit parameters for $GdL_1@AuNPs$ and $GdL_2@AuNPs$, obtained from the analysis of 1H NMRD data, are represented in Figure 4 and summarized in Table 2.



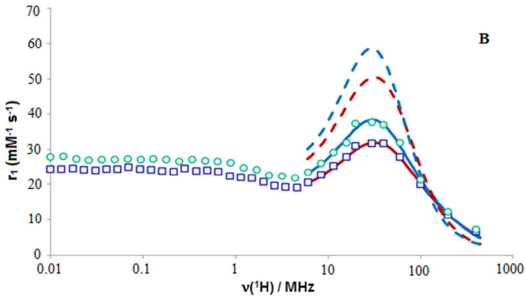


Figure 4. ¹H Nuclear Magnetic Relaxation Dispersion (NMRD) profiles for: **A-** GdL₁@AuNPs (0.56 mM; pH 7.0) 25 °C (\blacksquare) and 37 °C (\bullet); **B-** GdL₂@AuNPs (1.30 mM; pH 7.0); 25 °C (\circ) and 37 °C (\square). The fitted curves are represented as continuous lines. The broken lines are the result of simulations using the same parameters as in Table 2, but assuming total rigidity ($S^2 = 1$) of the immobilized chelates.

Table 2. Best fit parameters obtained for $GdL_1@AuNPs$ and $GdL_2@AuNPs$ from the fitting of the 1H NMRD profiles to the SMB theory, including the Lipari-Szabo approach for internal flexibility.

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	GdL ₁ @AuNPs	GdL ₂ @AuNPs
Parameters	Value	Value
q	<u>1</u>	1
ΔH^{\ddagger} [J/mol]	<u>17</u>	<u>17</u>
$k_{\rm ex}^{298} [10^7 {\rm s}^{-1}]$	<u>5.14</u>	<u>5.14</u>
E _R [kJ/mol] (global)	19.4±1.1	18±3.6
$ au_{\rm RH}^{298}[\rm ps]$ (global)	1900±140	3500±940
E _R [kJ/mol] (local)	<u>20</u>	<u>18</u>
$\tau_{\rm RH}^{298}[\rm ps]$ (local)	460±50	970±230
S ²	0.41 ± 0.04	0.42±0.12
$E_{\rm V}$ [kJ/mol]	1	1
$ au_{ m V}^{298} [m ps]$	27±4	17± 3
$E_{\rm H}^{298} [10^{-10} {\rm m}^2 {\rm s}^{-1}]$	23	<u>23</u>
E _{DGH} [kJ/mol]	<u>20</u>	<u>20</u>
Gd-O [Å]	2.5	2.5
$\Delta^2 [10^{20} \text{ s}^{-2}]$	0.044±0.002	0.065±0.004
Gd-HW 1 st [Å]	<u>3.1</u>	<u>3.1</u>
Gd-HW 2 nd [Å]	3.6	3.6

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Table 3. Selected molecular parameters for $GdL_1@AuNPs$ and $GdL_2@AuNPs$ and other systems reported in the literature and discussed in the manuscript.

Parameter	GdL ₁ @AuNPs ^a	GdL ₂ @AuNPs ^a	GdL ₃ @AuNPs ^b	$\operatorname{GdL_4}^c$
q	<u>1</u>	<u>1</u>	1	<u>1</u>
$k_{ex}^{298} [10^7 \text{ s}^{-1}]$	<u>5.14</u>	5.14	5.14	6.2
$\tau_g^{298} [ps]$	1900	3500	2470	3780
$\tau_{lo}^{298} [ps]$	460	970	177	930
S^2	0.41	0.42	0.48	0.24
$HD (nm)^d$	4.8	5.9	3.9	49 ^e
	27 ^{f,h}	38 ^{f,h}	28 ^{f,h}	$32^{g,h}$
$r_I \text{ (mM}^{-1} \text{ s}^{-1})$	11.2 ^{f,i}	$8.4^{f,i}$	$8.5^{f,i}$	n.d.

^a This work; ^bRef 27; ^cRef 26; ^d From DLS measurements; ^eZ-average from a bimodal distribution of particles; ^f Relaxivity per chelate; ^g Relaxivity of the aggregated form; ^h 20 MHz, 25 °C; ^l 200 MHz, 37 °C.

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The NMRD profiles are characteristic of macromolecular objects in slow rotation, confirming the immobilization of the GdL_1 and GdL_2 chelates onto gold

nanocrystals: a plateau in the frequency range 0.01 to 1 MHz, a simple dispersion at about 1-10 MHz and a broad hump centered at 20-60 MHz.

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The AuNPs prepared in this work display exceptional relaxivities (per Gd3++ 3 chelate) (r_{lmax} = 27 and 38 mM⁻¹ s⁻¹ for GdL₁@AuNPs and GdL₂@AuNPs, respectively; 4 30 MHz, 25 °C), much higher than those reported by other authors for AuNPs 5 functionalized with monoaquated (q = 1) Gd^{3+} complexes. 16,19,21 The temperature 6 dependence of the relaxivity, higher relaxivity at lower temperature, for both 7 GdL₁@AuNPs and GdL₂@AuNPs, indicates that the water exchange rate is not limiting 8 9 the relaxivity. The superb relaxivities attained can be ascribed to simultaneous optimization of the water exchange rate (fast water exchange regime) and of the 10 rotational correlation time. The value obtained for the order parameter ($S^2 \sim 0.40$) 11 indicates that fast local motions of the chelates anchored onto the Au core are still 12 limiting the relaxivity. Simulations, using the same parameters as on Table 3, but 13 assuming total rigidity of the immobilized chelates ($S^2 = 1$), afford much higher 14 relaxivities (of the order of magnitude 60 mM⁻¹ s⁻¹; 20 MHz; 25 °C) for GdL₂@AuNPs. 15 The higher relaxivity attained by GdL₂@AuNPs, comparing to GdL₁@AuNPs and 16 GdL₃@AuNPs, has to be ascribed to its significantly larger global rotational correlation 17 time (τ_{Rg}) , reflecting the larger size (hydrodynamic diameter) of the GdL₂@AuNPs 18 nanoparticles. In fact, the length of the thioalkyl linker seems not to have much 19 influence on the internal flexibility of the immobilized chelates: the order parameter S^2 20 is identical for GdL₁@AuNPs and GdL₂@AuNPs, despite the longer linker anchoring 21 GdL₂ to the Au core. Moreover, the shorter cysteine linker anchoring GdL₃ to the Au 22 23 core (possibly in a bidentate (N, S) fashion similar to GdL₁ (S,S)), results only in slightly higher rigidity of the immobilized chelates. Despite the limiting effect of the 24 internal rotational motions, chelate immobilization onto AuNPs results in relaxivity 25 enhancements of more than 300% for GdL₁ and over 500% for GdL₂ (comparing to its 26 monomeric form), attributed to simultaneous optimization of τ_R and k_{ex} . Moreover, 27 GdL₂ immobilized onto AuNPs displays substantially higher relaxivity than the 28 aggregated (micellar) form of GdL₂ (38 vs 15 mM⁻¹ s⁻¹; 20 MHz, 25 °C). The micellar 29 30 form of the $Gd[DO3A-N-(\alpha-pyrenebutanamido)propionate]$ chelate (GdL_4) (sharing the same coordination cage with $GdL_{1,2,3}$) is significantly more flexible ($S^2 = 0.24 \text{ vs } 0.42$) 31 than the Au-anchored GdL₁ and GdL₂ chelates. Accordingly, GdL₂@AuNPs, displaying 32 a τ_{Rg} value similar to the aggregated form of GdL₄, exhibits higher relaxivity (38 vs 32 33

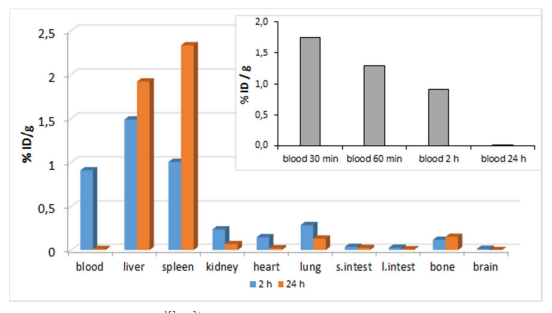
mM⁻¹ s⁻¹; 20 MHz, 25 °C, for GdL₂@AuNPs and for the aggregated form of GdL₄, respectively).²⁶

Covalent immobilization of Gd³⁺ chelates onto AuNPs seems more effective in attaining high relaxivities, than chelate self-assembly into micelle-type structures, owing to higher restriction of internal rotational motions.

The work reported here addresses explicitly the effect of linker length on the relaxivity of AuNPs functionalized with Gd³⁺ chelates, contributing to the "rational design" of nanomaterials as CA for MRI/multimodal imaging.⁴⁴

Biodistribution studies

The biodistribution of $[^{153}Sm]L_1@AuNPs$ in Wistar rats was obtained at 2 and 24 hours post-injection (Figure 6).



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Figure 6. Biodistribution of $[^{153}Sm^{3+}]L_1@AuNPs$ in Wistar rats, stated as percent of injected dose per gram of organ (% ID/g): a) 2 and b) 24 hour post-injection. Inset - time evolution of the activity in the blood. The results are from a group of four animals in each experiment.

The activity in the blood was measured after 30 minutes, 60 minutes, 2 and 24 hours (inset in Figure 6) revealing fast clearance of activity from the blood with a reduction of approximately 50% between 30 minutes and two hours. After 2 hours post-injection, the NPs are mainly found in the organs of the reticulo-endothelial system (RES), liver and spleen, and to a lesser extent in the blood and lungs. These results suggest that the nanoparticles are cleared mainly by phagocytosis by the macrophage

At 24 hours post-injection significant activity is found only in the organs of the

MRI studies were performed in male Swiss mice (~ 20 g) in a preclinical

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rich organs, liver and spleen, with a less important contribution from renal 3

elimination. 20,26,45 This is in accordance with what was found by MRI for GdL₁@AuNPs (see below).

4 reticulo-endothelial system, RES. The activity approximately doubled in the spleen, 5 showing only a slight increase in the liver. The activity in the bones at 24 hours post-6 7 injection is very low, suggesting that the rate of chelate demetallation and formation of

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MRI Studies

insoluble metal colloids in vivo is very low.

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imaging platform (PharmaScan) operating at 7.0 Tesla (300 MHz). A Dynamic Contrast Enhancement (DCE) study was performed with GdL₁@AuNPs (0.1 mmol Gd/kg body weight) and GdL₃@AuNPs (0.1 and 0.05 mmol Gd/kg body weight) and for comparison purposes with Gd(DTPA) (Magnetvist[®], Bayer) at the same doses (Figure 7). Figure 7 shows a representative series of T₁-weighted spin-echo coronal images. In the pre-injection images, the kidney structures (cortex, inner and outer medulla) and adjacent tissues appear dark. After bolus injection in the vascular system, a strong signal enhancement was observed in the kidneys for Gd(DTPA), GdL₁@AuNPs and GdL₃@AuNPs as result of T₁ shortening. A much slighter signal enhancement was observed in the liver. Both NPs follow mainly renal elimination by glomerular filtration, with significant hepatobiliary contribution to excretion seen for GdL₁@AuNPs only.

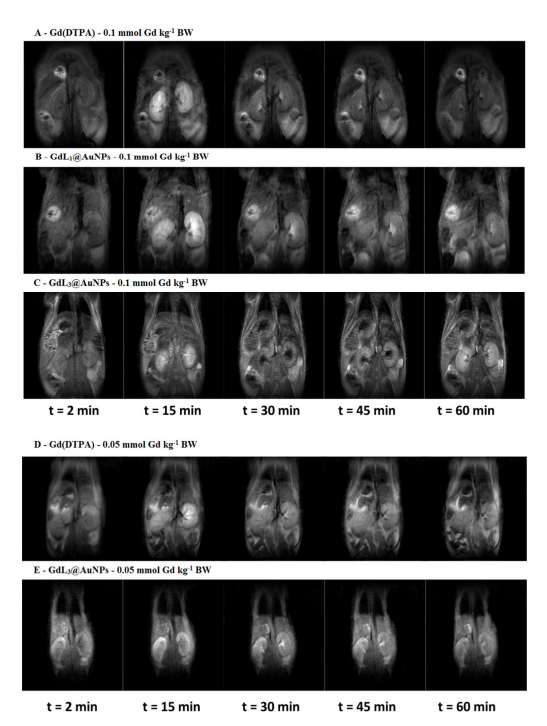


Figure 7. Representative coronal T₁-weighted spin echo MR images of mice before and after injection of contrast agents: (**A**) Gd(DTPA) (0.1 mmol Gd kg⁻¹ BW), (**B**) GdL₁@AuNPs (0.1 mmol Gd kg⁻¹ BW), (**C**) GdL₃@AuNPs (0.1 mmol Gd kg⁻¹ BW); (**D**) Gd(DTPA) (0.05 mmol Gd kg⁻¹ BW), (**E**) GdL₃@AuNPs (0.05 mmol Gd kg⁻¹ BW).

The time course of the average intensity (mean values of groups of four animals) within different regions of interest (ROIs) placed on the several organs (Figure 8) allows to understand better the features of Figure 7.

Figure 8. Time course of signal intensity, up to 60 min post-injection, for several regions of interest, relative to the initial value, during dynamic contrast enhancement MRI experiments in rats administrated with: (A) Gd(DTPA) (0.1 mmol kg⁻¹ BW) and (B) GdL₁@AuNPs (0.1 mmol kg⁻¹ BW), (C) GdL₃@AuNPs (0.1 mmol kg⁻¹ BW); (D) Gd(DTPA) (0.05 mmol kg⁻¹ BW); (E) GdL₃@AuNPs (0.05 mmol kg⁻¹ BW). The time courses are data from mean values of four animals.

In order to compare the results for all the animals under study (n = 4), the data were normalized by calculating the mean relative enhancement of each ROI. The scattering in the time course curves was caused by animal respiratory motion. The relative enhancement obtained with Gd(**DTPA**) at 0.1 mmol kg⁻¹ BW dose (Figure 8A), increased almost immediately after intravenous injection, from 0 up to about 160% in the kidney medulla and 100% in the kidney cortex, followed by a steady decrease to values around 60% and 30%, respectively, within 60 minutes. This time course is in agreement with the literature for the Gd(DTPA) and Gd(DOTA) low molecular weight CA.^{46,47}

1	The enhancement profiles of GdL ₁ @AuNPs (Figure 8B) and GdL ₃ @AuNPs
2	(Figure 8C) at 0.1 mol Gd kg ⁻¹ BW dose are considerably different from the
3	enhancement profile of Gd(DTPA) at the same concentration: there is an immediate
4	enhancement of the kidney structures (cortex and medulla) followed by a steady liver
5	enhancement. For GdL ₃ @AuNPs at 0.1 mmol Gd kg ⁻¹ BW dose (Figure 8C) is
6	noticeable a fast and strong enhancement of the kidney medulla and kidney cortex (~
7	150%) which slowly decreases to \sim 100% over the time course of the experiment. A
8	much lower muscle and liver enhancement is also noticeable. Reducing the dose of
9	GdL ₃ @AuNPs to 0.05 mmol Gd Kg ⁻¹ BW results in an imaging profile virtually
10	equivalent to Gd(DTPA): fast renal elimination with negligible hepatobiliary
11	contribution (Figure 8D and 8E for $Gd(DTPA)$ and $GdL_3@AuNPs$, respectively). There
12	is a fast enhancement of the kidney cortex (~150 % at 20 minutes) which steadily
13	decreases over the time course of the experiment. These results strongly suggest that
14	while $\text{Gd}L_1@\text{AuNPs}$ is mostly eliminated through hepatobiliary excretion, or is taken
15	up by resident macrophages (Kupfer cells) in liver, GdL ₃ @AuNPs behaves in vivo as a
16	low molecular weight CA following mainly renal elimination. The steady, liver and
17	presumably spleen enhancement observed with $GdL_1@AuNPs$ is in sharp contrast to
18	the "clean" renal elimination observed for $GdL_3@AuNPs$. This behaviour can only be
19	explained by the difference in size between $\text{Gd}L_1@\text{AuNPs}$ and $\text{Gd}L_3@\text{AuNPs}\text{-}$ average
20	HD 4.8 and 3.9 nm, respectively, stressing the complex interplay between the physical-
21	chemical properties of nanostructures and in vivo behaviour.
22	The animal MRI studies were performed at high field (300 MHz, 7 Tesla). This study
23	illustrates the mismatch between the performance of macromolecular/nanosized CA
24	optimized for intermediate fields (20-60 MHz), and the trend for increasingly higher
25	magnetic field imagers. The overwhelming advantage of the AuNPs, over low
26	molecular weight CA at intermediate fields (20-60 MHz), is partially eroded at higher
27	magnetic fields (Figure 4). ⁴⁸ Nonetheless, the AuNPs studied in this work still exhibit
28	relaxivities significantly higher than $Gd(\mathbf{DTPA})$ at high fields (11.2, 8.4 $vs \sim 2 \text{ mM}^{-1} \text{ s}^{-1}$
29	for GdL1@AuNPs and GdL2@AuNPs, respectively, and Gd(DTPA), 200 MHz, 25 $^{\circ}\text{C}$).

Conclusions

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In this work we extend the synthetic methodologies developed before for AuNPs functionalized with stable fast water exchanging Gd³⁺ chelates as high relaxivity, potentially safe CA for in vivo MRI. Two novel ligands were designed to investigate the role of the length of the ω-thioalkyl linker, anchoring the coordination cage to the gold nanocrystal, on the relaxivity. Superb relaxivities at magnetic fields relevant for clinical imaging (27 and 38 mM⁻¹ s⁻¹, 30 MHz, 25 °C, for GdL₁@AuNPs and GdL₂@AuNPs, respectively) were obtained attained thanks to simultaneous optimization of the rotational correlation time and of the water exchange rate. Relaxivities, still relevant for clinical high field applications (of the order of magnitude 10 mM⁻¹ s⁻¹; 200 MHz, 37 °C) were attained also. The relaxivity is still limited by internal flexibility of the immobilized chelates. The degree of internal flexibility of the immobilized chelates (measured by the order parameter S^2) seems not to be determined by the length of the linker, presumably owing to the high surface curvature of the NPs. A MRI study in mice demonstrated that while GdL_3 @AuNPs (HD = 3.9 nm) behaves in vivo much like the low molecular weight CA Gd(DTPA), undergoing fast renal elimination without liver (and presumably spleen) uptake, $GdL_1@AuNPs$ (HD = 4.8 nm) shows considerable hepatobiliary contribution to elimination. A biodistribution study in rats using the ¹⁵³SmL₁@AuNPs tracer confirmed extensive activity uptake and accumulation over time in the liver and spleen.

The GdL₃@AuNPs CA, amenable to further elaboration with targeting moieties, seems particularly promising for *in vivo* MRI applications.

The work reported is a relevant contribution towards the design of nanomaterials functionalized with Gd³⁺ chelates as very high relaxivity/multimodal CA for MRI.⁴⁴

Experimental

Materials and methods

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- 4 Chemicals were purchased from Sigma-Aldrich and used without further purification.
- 5 Cyclen was purchased from Chematech, France. Analytical grade solvents were used
- and not further purified, unless specified. Reactions were monitored by TLC on silica
- 7 gel by examination under UV light (250 and 365 nm) and staining with iodine vapour
- 8 and Ellman's reagent. Preparative chromatography was carried out on Silica Gel 60
- 9 (230-400 mesh). Ion exchange chromatography was performed on Dowex 1X2-100-
- 10 OH (50-100 mesh) resin. Size Exclusion Chromatography (SEC) was performed on
- 11 Sephadex G10 (40-120 µm) with water elution. Dialysis was performed against water
- 12 on cellulose membranes (MWCO 10 KDa). UV-VIS spectra were acquired with a
- 13 Shimadzu UV-2501PC spectrophotometer. The size distribution and zeta potential of
- the AuNPs was determined with a Malvern Zetasizer, NANO ZS (Malvern Instruments
- Limited, UK), using a He-Ne laser (wavelength of 633 nm) and a detector angle of 173°.
- 16 TEM experiments were performed with a JEOL JEM1200EXII microscope at Bath
- 17 University, UK. Mass spectrometry was performed at CACTI Vigo, Spain.
- ¹H and ¹³C NMR spectra were run on Varian Unity Plus 300, Bruker Avance-3
- 19 400 Plus and Varian VNMRS 600 NMR spectrometers. Chemical shifts (δ) are given in
- 20 ppm relative to the CDCl₃ solvent (¹H, δ 7.27; ¹³C 77.36) as internal standard. For ¹H
- and 13 C NMR spectra recorded in D₂O, chemical shifts (δ) are given in ppm, relative to
- TSP as internal reference (${}^{1}H$, δ 0.0) and *tert*-butanol as external reference (${}^{13}C$, CH₃ δ
- 23 30.29), respectively.

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Preparation of Lipoic acid conjugate DO₃A-N-(α-lipoamido) propionate - L_1

- 27 Synthesis of ((5-(1,2-dithiolan-3-yl)-2-pentanamido)methoxycarbonylethyl)-4,7,10-
- 28 tris-(ethoxycarbonylmethyl) -1,4,7,10-tetraazacyclododecane fully protected
- 29 conjugate 7.
- 30 Orthogonally protected compound 6 was synthesized as described before by us.²⁷ A
- 31 solution of compound 6 (85 mg; 1.12 mmol) in a mixture DCM/TFA (24 ml, 3:1, v/v)
- 32 was stirred at room overnight. The solvent was removed under reduced pressure, the
- 33 residue was re-dissolved in DCM and the solvent was evaporated. This procedure was
- repeated several times.

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- The resulting oil was dried under *vacuum* to afford a white foam. ¹H NMR (CDCl₃) 1 2 revealed the disappearance of the signals assigned to the Boc groups on compound 6. Quantitative deprotection was assumed. The residue (1.12 mmol, assuming quantitative 3 4 deprotection) was dissolved in DCM (20 ml) and the solution was adjusted to pH 9-10 (pH paper) by drop-wise addition of DIPEA. To this solution was added sequentially 5 Lipoic acid (288 mg; 1.40 mmol), HOBt (214 mg; 1.40 mmol) and a solution of DCC 6 (288 mg; 1.40 mmol) in DCM (5 ml). The solution was stirred at room temperature 7 overnight. The DCU byproduct was removed by filtration and the reaction mixture was 8 9 concentrated under reduced pressure. The residue was re-dissolved in ethyl acetate (100 ml), and the solution was washed with NaHCO₃ (50 ml, saturated solution) and brine 10 (3x50 ml). The organic phase was dried (MgSO₄) and concentrated under reduced 11 pressure to afford the title compound (7) (358 mg; 44 %). ¹H NMR (300 MHz, CDCl₃): 12 δ = 1.28 (m, 9 H, C(O)OCH₂CH₃), 1.48 (m, 2 H, NHC(O)CH₂CH₂CH₂), 1.70 (m, 2 H, 13 NHC(O)CH₂CH₂), 2.18 (m, 2 H, NHC(O)CH₂CH₂CH₂CH₂), 2.30 (m, 4 H, NHC(O)CH₂ 14 and CHCH₂CH₂S), 2.60-3.60 (broad overlapped signals, integrating for 16 H, 15 N(CH₂)₂N; 2 H, ABX; 2 H, CHCH₂CH₂S, 1 H, CHSCH₂CH₂S), 3.73 (m, 6 H, 16 $C(O)CH_2N$), 3.97 (s, 3 H, $C(O)OCH_3$), 4.19 (m, 6 H, $C(O)CH_2CH_3$), 4.90 (dd,1 H, 17 ABX). HRMS (ESI): m/z: cacd. for $C_{32}H_{58}N_5O_9S_2[M+H]^+$: 720.3676, found: 720.3645. 18 19 20 Preparation of ((5-(1,2-dithiolan-3-yl)-2-pentanamido)carboxyethyl)-4,7,10-tris-(carboxymethyl)-1,4,7,10-tetraazacyclododecane - fully deprotected DO3A-N-(α-21 lipoamido) propionate chelator (L_1) . 22 Compound (7) (2.26 g, 3.15 mmol) was dissolved in a mixture water/ethanol (40 ml, 1/1 23 v/v). The solution was adjusted to pH ~ 11 with aqueous NaOH 1 M (pH paper) and 24 was left stirring at room temperature overnight. Then, the reaction mixture was adjusted 25 to pH ~7 with hydrochloric acid 1 M (pH paper) and concentrated under reduced 26 pressure. The residue was adsorbed onto silica and purified by flash chromatography 27 $(CH_2Cl_2 \rightarrow CH_2Cl_2/EtOH \ 1/1 \rightarrow EtOH \rightarrow EtOH/H_2O \ 1/1 \rightarrow H_2O)$ to afford a light yellow 28 foam. The final compond (L₁) was further purified by size exclusion chromatography 29
- on Sephadex G10 (0.42 μ m) with elution with water. The conductivity of the collected fractions was measured and were also tested by TLC (ethanol/water (1/1), revelation with iodine vapor). The high conductivity fractions (salt) were discarded and the

medium/low conductivity fractions showing a signal on the TLC were pooled,

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concentrated at room temperature and further dried under vacuum to afford the final

- deprotected compound as a light yellow solid (L₁) (0.685 g, 35%). ¹H NMR (300 MHz,
- 2 D_2O): $\delta = 1.46$ (m, J = 7.8 Hz, 2 H, NHC(O)CH₂CH₂CH₂), 1.64-1.72 (m, 6 H,
- 3 NHC(O)CH₂CH₂, NHC(O)CH₂CH₂CH₂CH₂, CHCH₂CH₂S), 2.01 (m, 2 H,
- 4 NHC(O)C H_2), 2.34 (t, J=7.5 Hz 2 H, CHC H_2 C H_2 S), 2.49 (m, 1 H, CHSC H_2 C H_2 S),
- 5 2.10-3.40 (broad overlapped signals integrating to 16 H, 4 x $N(CH_2)_2N$, 6 H, 3x
- 6 NCH₂C(O) and 2H, ABX), 4.49 (m, 1 H, ABX). ¹³C NMR (75.4 MHz, D₂O): 25.02 (1
- 7 C, CH₂), 28.32 (1 C, CH₂), 33.93 (1 C, CH₂), 35.73 (1 C, CH₂), 38.22 (1 C, CH₂), 40.46
- 8 (2 C, 2xCH₂), 47.81 (3 C, 3xCH₂), 49.46 (1 C, CH₂), 51.37 (1 C, CHCH₂), 51.94 (2 C,
- 9 CH₂), 54.51 (2 C, CH₂), 56.08 (2 C, CH₂), 56.76 (2 C, CH₂), 170.94 (1 C, C(O), 176.65
- 10 (2 C, 2xC(O), 177.36 (2 C, 2xC(O)). HRMS (ESI): m/z: cacd. for C₂₅H₄₄N₅O₉S₂
- 11 [M+H]⁺: 622.2580, found: 622.2572.

- 13 Preparation of 11-mercaptoundecanoic acid conjugate DO3A-N-(α-
- 14 mercaptoundecanamido) propionate L₂

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Synthesis of 11-(acetylthio)undecanoic acid (10)

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- To an ice-cooled solution of 11-mercaptoundecanoic acid (9) (2.00 g, 9.17 mmol) in
- 19 pyridine (2.6 ml) was added acetic anhydride (2.6 ml, 2.81 g, 27.5 mmol). The solution
- 20 was left stirring at room temperature overnight. Ice was directly added to the reaction
- 21 mixture, followed by magnetic stirring until complete melting of the ice. The mixture
- was extracted with ethyl acetate (3×150 ml). The organic phase was washed with brine
- 23 (3x30 ml), dried (MgSO₄) and the solvent was removed under reduced pressure. The
- 24 residue was further dried under vacuum to afford the final compound as an off-white
- solid (2.12 g, 89%). ¹H NMR (400 MHz, CDCl₃): δ = 1.27 (s (*br*),12 H, 6×CH₂), 1.58-
- 26 1.52 (m, 2 H, SCH₂CH₂), 1.66-1.59 (m, 2 H, (CO₂H)CH₂CH₂), 2.32 (s, 3 H, C(O)CH₃),
- 27 2.35 (t, J= 7.2 Hz, 2 H, CH_2COOH), 2.86 (t, J= 7.2 Hz, 2 H, SCH_2). ¹³C NMR (100.62)
- 28 MHz, CDCl₃): δ = 24.61 (C(O)CH₂CH₂), 28.73 (SCH₂), 28.97, 29.00, 29.10, 29.13,
- 29 29.26, 29.32 (overlapped inner CH₂ signals), 29.43 (SCH₂CH₂), 30.59 (CH₃), 33.99
- 30 (C(O)CH₂), 179.89 (COOH), 196.11 (SC(O)Me). HRMS (ESI): m/z: calcd. for
- 31 $C_{13}H_{24}NaO_3S[M+Na]^+$: 283.1338, found.: 283.1339.

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Synthesis of (11-(acetylthio)undecanoyl)serine methyl ester (11)

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3 To an ice-cooled solution of compound 10 (2.12 g, 8.14 mmol) in acetonitrile (70 ml) 4 was added HOBt (1.85 g, 8.95 mmol) and a solution of DCC (1.25 g, 8.14 mmol) in 5 6 acetonitrile (10 ml). The mixture was left stirring at the ice bath temperature and after 7 15 minutes L-serine methyl ester hydrochloride (1.27 g, 8.14 mmol) and triethylamine 8 (1.13 ml, 0.82 g, 8.14 mmol) were added. The reaction mixture was left stirring at room 9 temperature overnight. The DCU byproduct was removed by filtration and the sample was concentrated under reduced pressure. The residue was re-dissolved in ethyl acetate 10 (100 ml) and the solution was washed sequentially with KHSO₄ (1 M, 3x50 ml), 11 NaHCO₃ (saturated solution, 50 ml) and brine (3x50 ml). The organic phase was dried 12 (MgSO₄) and concentrated under reduced pressure to afford title compound 11 (2.73 g; 13 93%). H-NMR (CDCl₃, 400 MHz): δ = 1,25 (s (*br*), 12 H, 6 × CH₂), 1.52-1.57 (m, 2 H, 14 SCH_2CH_2), 1.58-1.65 (m, 2 H, $NHC(O)CH_2CH_2$), 2.25 (t, J=7.2 Hz, 2 H, 15 NHC(O)C H_2), 2.30 (s, 3 H, SC(O)C H_3), 2.84 (t, J=7.2 Hz, 2 H, SC H_2), 3.76 (s, 3 H, 16 OCH_3), 3.87 (ddd, J=3.6, 11.2, 29.8 Hz, 1 H, CH_aH_bOH), 3.96 (ddd, J=4.0, 11.0 and 17 29.8 Hz, 1 H, CH_aH_bOH), 4.67-4.63 (m, 1 H, CH), 6.62 (s (*br*), 1 H, NH). ¹³C-NMR 18 19 $(CDCl_3, 100.62 \text{ MHz}): \delta = 24.80 \text{ (NHC(O)CH}_2\text{CH}_2), 28.67, 28.95, 29.06, 29.09$ (overlapped CH₂ signals), 29.17 (SCH₂), 29.25, 29.27 (overlapped CH₂ signals), 29.37 20

 (SCH_2CH_2) , 30.55 (SC(O)Me), 36.37 $(C(O)CH_2)$, 52.54 (OMe), 54.54 (CH), 63.20

(CH₂OH), 171.03 (C(O)OMe), 173.78 (NHC(O)), 196.19 (SC(O)).

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of
 1
      Synthesis
                                      N-(tert-butoxycarbonyl), N-(11-(acetylthio)undecanoyl)
      dehydroalanine methyl ester (3)
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      To a solution of compound (11) (0.866 g, 2.40 mmol) in dry acetonitrile (15 ml) was
      sequentially added DMAP (0.081 g, 0.66 mmol) and Boc<sub>2</sub>O (1.44 g, 6.6 mmol). The
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      mixture was left stirring for 5 days at room temperature. The reaction progress was
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      monitored by <sup>1</sup>H NMR. A small volume of reaction mixture was removed, worked-up
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      was as described below, and analyzed by <sup>1</sup>H NMR by monitoring the disappearance of
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      the signal of the intermediate carbonate ester and the appearance of the alkenic signals
      at \delta= 5.62 and 6.44 ppm. The solid residues were removed by filtration and the sample
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11
      was concentrated under reduce pressure. The residue was re-dissolved in ethyl acetate
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      (150 ml), and the solution was washed sequentially with KHSO<sub>4</sub> (1 M, 3x50 ml),
      NaHCO<sub>3</sub> (saturated solution, 50 ml) and brine (3x50 cm<sup>3</sup>). The organic phase was
13
      concentrated under reduced pressure and the residue was purified by a flash
14
      chromatography (n-hexane\rightarrow n-hexane/ethyl acetate
                                                                   (70:30)) to afford the title
15
      compound as a thick reddish oil (0.445 \text{ g}, 58\%). H-NMR (CDCl<sub>3</sub>, 400 MHz) \delta= 1.27 (s
16
      (br), 12 H, 6 \times CH_2), 1.46 (s, 9 H, C(CH_3)_3), 1.49-1.57 (m, 2 H, SCH_2CH_2), 1.65 (m, 2 H,
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      C(O)CH_2CH_2), 2.32 (s, 3 H, SC(O)CH_3), 2.86 (t, J=7.6 Hz, 2 H, NHC(O)CH_2), 2.93
18
      (t, J = 7.6 \text{ Hz}, 2 \text{ H}, \text{SC}H_2), 3.78 \text{ (s, 3 H, C(O)OC}H_3), 5.62 \text{ (s, 1 H, CCH}_3H_b), 6.44 \text{ (s, 1)}
19
      H, CCH_aH_b). <sup>13</sup>C-NMR (CDCl<sub>3</sub>, 100.62 MHz): \delta = 24.71 (C(O)CH<sub>2</sub>CH<sub>2</sub>), 27.40
20
      (SCH<sub>2</sub>CH<sub>2</sub>), 27.80 (3×OCCH<sub>3</sub>), 28.75, 29.05, 29.08, 29.11, 29.35, 29.44 (overlapped
21
22
      CH<sub>2</sub> signals), 29.38 (C(O)CH<sub>2</sub>), 30.59 (SC(O)CH<sub>3</sub>), 37.66 (SCH<sub>2</sub>), 52.38 (C(O)OCH<sub>3</sub>),
      83.53 (OCCH<sub>3</sub>), 125.71 (CCH<sub>2</sub>), 135.56 (CCH<sub>2</sub>), 151.52 (NC(O)O), 163.68
23
      (C(O)OCH_3), 175.66 (NC(O)), 196.00 (SC(O)). HRMS (ESI): m/z: calcd. for
24
      C_{22}H_{37}NaO_6S [M+Na]^+: 466.2234, found: 466.2223.
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27
      Synthesis of (11-(acetylthio)-2-N-(tert-butoxycarbonyl)undecanamido-
28
      methoxycarbonylethyl)-1,4,7,10-tetraazacyclododecane - monoalkylated cyclen (5)
29
      To a solution of cyclen (0.260 g, 1.5 mmol) in acetonitrile (30 ml) was added K<sub>2</sub>CO<sub>3</sub>
30
31
      (0.83 g, 6.0 mmol) and in several portions compound 3 (0.445 g, 1.0 mmol). The
      suspension was vigorously stirred at room temperature for 4 hours. The suspended solid
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      was removed by filtration and the solvent was evaporated under reduced pressure.
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The
                        residue
                                                                          flash
 1
                                                 purified
                                                                by
                                                                                      chromatography
                                      was
 2
      (CH_2Cl_2 \rightarrow CH_2Cl_2/EtOH/NH_3/H_2O (70:30:5:5)) to the afford the title compound 5 as a
      white foam (0.451 \text{ g}, 73.0 \%). H-NMR (CDCl<sub>3</sub>, 400 MHz): \delta = 1.33 (s (br), 12 H,
 3
 4
      6 \times CH_2), 1.47 (s, 9 H, OC(CH_3)<sub>3</sub>), 1.60 (m, 2 H, C(O)CH<sub>2</sub>CH<sub>2</sub>), 1.60-1.65 (m, 2 H,
      SCH_2CH_2), 2.33 (s, 3 H, SC(O)CH_3), 2.51-2.65 (m, 16 H, 4\times N(CH_2)_2N), 2.74-2.79 (m,
 5
      2 H, (NHC(O)CH_2)), 2.74-2.79 (m, 1 H, NCH_aCH_bCH), 2.85 (t, J=7.2 Hz, 2 H,
 6
 7
      CH_3C(O)SCH_2, 3.45 (dd, J=5.2 and 14.4 Hz, 1 H, NCH_0CH_0CH_0, 3.68 (s, 3 H,
      C(O)OCH_3), 5.46 (t, J=5.2 Hz, 1 H, NCH_2CH). <sup>13</sup>C-NMR (CDCl<sub>3</sub>, 100.62 MHz): \delta=
 8
      25.01 (C(O)CH<sub>2</sub>CH<sub>2</sub>), 27.90 (OCCH<sub>3</sub>), 28.75 (SCH<sub>2</sub>), 29.05, 29.10, 29.17, 29.37, 29.39,
 9
      29.40 (overlapped CH<sub>2</sub> signals), 29.43 (SCH<sub>2</sub>CH<sub>2</sub>), 30.58 (SC(O)Me), 40.06
10
      (C(O)CH_2), 46.92 (6\times NHCH_2), 51.07 (2\times CH_2NCH_2CH), 52.16 (C(O)OMe), 53.56
11
      (NCH_2CH), 58.06 (NCH_2CH), 83.99 (C), 151.96 (NC(O)O), 170.84 (C(O)OMe),
12
      175.52 \text{ (N(Boc)}C(O)CH_2), 195.99 \text{ (SC(O))}. HRMS \text{ (ESI): m/z: calcd. for } C_{30}H_{58}N_5O_6S
13
      [M+H]<sup>+</sup>: 616.4102, found.: 616.4100.
14
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Synthesis of (11-(acetylthio)-2-undecanamido-methoxycarbonylethyl)-4,7,10tris-(ethoxycarbonylmethyl)-1,4,7,10-tetraazacyclododecane - fully alkylated cyclen **(8)**

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> A solution of monoalkylated cyclen 5 (0.451 g, 0.87 mmol) in trifluoroacetic acid in dichlorometane (33%, 24 ml) was stirred overnight at room temperature. The solvent was evaporated at reduced pressure and the residue was re-dissolved in dichlorometane. The solvent was evaporated again, and this procedure was repeated several times to give a light thick yellow oil which was further dried under vacuum. ¹H NMR spectroscopy (CDCl₃) revealed the disappearance of the signal assigned to the Boc group in the precursor compound 5. The deprotected compound (0.87 mmol, assuming quantitative deprotection) was re-dissolved in MeCN (20 ml), K₂CO₃ (1.17 g, 8.46 mmol) was added and the suspension was left under vigorous stirring at room temperature for 30 minutes. Ethyl bromoacetate (0.29 ml, 2.61 mmol) was added, and the suspension was further stirred for 2 hours. The suspended solids were removed by filtration, the solvent was evaporated under reduce pressure and the residue was purified by flash chromatography (CH₂Cl₂→ CH₂Cl₂/EtOH (7:3)) to afford compound **8** (0.218 g, 32 %) as a white foam.

Dalton Transactions

1	¹ H-NMR (CDCl ₃ , 400 MHz): δ = 1.33 (m, 12 H, 6×CH ₂), 1.47 (s, 9 H, OC(<i>CH</i> ₃) ₃), 1.60
2	(m, 2 H, C(O)CH ₂ CH ₂), 1.60-1.65 (m, 2 H, SCH ₂ CH ₂), 2.32 (s, 3 H, SC(O)Me), 2.70-
3	2.90 (m, 16 H, $4 \times N(CH_2)_2N$), 2.74-2.79 (m, 2 H, $(C(O)CH_2)$), 2.74-2.79 (m, 1 H,
4	NCH_aCH_bCH), 2.85 (t, $J=7.2$ Hz, 2 H, SCH_2), 3.49 (m, 6 H, $3\times C(O)CH_2CH_3$ and 1 H,
5	NCH_aCH_bCH), 3.72 (s, 3 H, C(O)OC H_3), 4.19 (m, 6 H, C(O)OC H_2CH_3), 5.46 (t, $J=5.2$
6	Hz, 1 H, NCH ₂ CH). HRMS (ESI): m/z: calcd. for $C_{37}H_{68}N_5O_{10}S$ [M+H] ⁺ : 774.4681,
7	found: 774.4684.
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- 1 Synthesis of (11-mercapto-2-undecanamido-carboxyethyl)-4,7,10-tris-
- 2 (carboxymethyl)-1,4,7,10-tetraazacyclododecane Fully deprotected DO3A-N-(α-
- 3 mercaptoundecanamido)propionate chelator (L₂)
- 4 Compound 8 (0.218 g, 0.28 mmol) was dissolved in a mixture EtOH/H₂O (20 ml, 1:1
- 5 (v/v)). The solution was adjusted to pH \sim 10-11 (pH paper) with aqueous NaOH (1 M)
- and left stirring at room temperature overnight. The solution was adjusted to pH \sim 7 (pH
- 7 paper) with diluted hydrochloric acid (1M) and was evaporated at reduced pressure
- 8 (temperature < 40 °C). The residue was adsorbed onto silica and purified by flash
- 9 chromatography (CH₂Cl₂ \rightarrow CH₂Cl₂/EtOH 1/1 \rightarrow EtOH \rightarrow EtOH/H₂O 1:1 \rightarrow H₂O) to afford
- 10 a light yellow foam. The residue was re-dissolved in water and was purified by size
- exclusion chromatography (Sephadex G10). The relevant fractions were pooled together
- and the solvent was removed under reduced pressure to give chelator L₂ (0.074 g, 42
- 13 %). H-NMR (CDCl₃, 400 MHz): δ = 1.33 (s (*br*), 10 H, 5×CH₂),1.61 (s (*br*), 2 H,
- 14 SCH₂CH₂CH₂), 1.63 (s (*br*), 2 H, (C(O)CH₂CH₂), 1.72 (s (*br*), 2 H, SCH₂CH₂), 2.33 (s
- 15 (br), 2 H, (C(O)C H_2), 2.58 (t, J=6.8 Hz, 2 H, SC H_2), 2.80 (s (br), 1 H, NC H_aH_b CH),
- 2.80 (s (*br*), 1 H, NC*H_a*H_bCH), 3.18 (s, 4 H, NCH₂C*H*₂NCH₂CH), 3.39-3.36 (m, 4 H,
- 17 N(CH₂)₂N), 3.43 (s, 8 H, $2\times$ N(CH₂)₂N), 3.76 (s (*br*), 6 H, $3\times$ NCH₂(C(O)OH), 4.52 (s
- 18 (*br*), 1 H, C*H*). ¹³C-NMR (CDCl₃, 100.62 MHz): δ = 23.82 (SCH₂), 25.13
- 19 (C(O)CH₂CH₂), 27.73 (SCH₂CH₂), 28.11, 28.31, 28.39, 28.52, 28.61 (overlapped CH₂
- 20 signals), 33.08 (SCH₂CH₂CH₂), 35.84 (C(O)CH₂), 38.51 (NCH₂CH), 48.79 (2×
- 21 NCH₂CH₂NCH₂CH), 51.20 (CH), 51.62 ($2 \times NCH_2CH_2NCH_2CH$), 56.56 ($3 \times$
- 22 NCH₂(C(O)OH), 56.09 (2×NCH₂CH₂N), 177.14 (NHC(O)), 177.14 (CHC(O)OH),
- 23 177.30 ($3 \times NCH_2C(O)OH$). HRMS (ESI): m/z: calcd for $C_{28}H_{51}N_5NaO_9S$ [M+Na]⁺:
- 24 656.3305, found: 656.3300.

Preparation of GdL₁ and GdL₂ complexes for relaxometric measurements

- 28 A solution of GdCl₃.6H₂O was added drop-wise, under magnetic stirring, to an
- equimolar solution of L₁ or L₂ (5% excess), while keeping the solution pH around 5.8
- 30 (pH meter) by the addition of diluted NaOH. The solution was left stirring at room
- 31 temperature overnight. Then, the solution was adjusted to pH 7.0 with NaOH (0.1 M)
- and filtered through a 0.2 μm syringe filter. The absence of free Gd³⁺ was confirmed by
- 33 the xylenol orange test. The final concentration of Gd was determined by ICP-OES
- 34 following sample digestion with nitric acid.

Preparation gold nanoparticles functionalized with GdL₁ and GdL₂ chelates.

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An aqueous solution of ligand DO3A-N-(α-Lipoamido)propionate (L₁) (20.5 mM, 4.53 4 ml, 0.091 mmol) was added drop-wise, under magnetic stirring at room temperature, to 5 an aqueous solution of HAuCl₄ (58.86 mM, 1.54 ml, 0.091 mmol). During the initial 6 7 stages of the addition of L₁, the light yellow HAuCl₄ solution turned dark orange, 8 fading away in color to light yellow with further addition of ligand. To the reaction 9 mixture was added, in one aliquot, a freshly prepared aqueous NaBH₄ solution (522) mM, 0.179 ml, 0.093 mmol). The reaction mixture turned instantaneously dark brown 10 and was left stirring at room temperature for 16 hours. The NPs solution was adjusted to 11 12 pH \sim 7 (pH meter) by adding aqueous NaOH (0.1 M) and was filtered through a 0.20 µm PTFE syringe filter. A small volume of solution (1 ml) was kept for further 13 characterization. To remaining NPs solution (~ 5.3 ml) was added slowly a solution of 14 15 GdCl₃.6H₂O in water (303 µM, 0.300 ml, 0.091 mmol) while keeping the solution pH around 5.5 (pH meter) by adding aqueous NaOH 0.1 M. The NPs solution was left 16 stirring at room temperature for 16 hours and was adjusted to pH \sim 7 with aqueous 17 NaOH (1 M solution). The nanoparticles were purified by size exclusion 18 19 chromatography (Sephadex G10, 0.42 µm) with elution with water. The entire colored broad band eluting from the column was collected without attempting to fractionate the 20 sample. The nanoparticles were further purified by extensive dialysis against water 21 using a 10 KDa MWCO cellulose membrane. The xylenol orange test indicated the 22 absence of free Gd³⁺ in the gold nanoparticles preparation. 23

The same procedure was followed for the preparation of GdL2@AuNPs starting from ligand L2 (20.5 mg/mL, 2 ml, 0.0647 mmol) and HAuCl4 (22 mg/ml, 1 ml, 0.0647 mmol).

The Gd and Au content of the NPs preparations ([Gd]= 0.57 mM and 1.30 mM; [Au]/[Gd]= 1.40 and 0.87, for GdL₁@AuNPs and GdL₂@AuNPs, respectively) was determined by ICP-OES analysis following sample digestion with *aqua regia*.

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NMRD measurements

- 1 The NMD measurements were performed using a StelarSpinmaster FFC NMR
- 2 relaxometer (0.01-20 MHz) equipped with a VTC90 temperature control unit. At higher
- 3 fields, the ¹H relaxivity measurements were performed on Bruker Minispecs mq30 (30
- 4 MHz), mq40 (40 MHz) and mq60 (60 MHz), as well as Bruker Advance spectrometers
- 5 connected to 2.35 T, 4.7 T and 9.4 T superconducting magnets. In each case, the
- 6 temperature was measured by a substitution technique. Variable temperature
- 7 measurements were performed at 25 and 37 °C. The NMRD profiles were analysed
- 8 using the Visualiseur/Optimiseur 3.6 program running on a Matlab® 6.5 platform. 49

Relaxivity studies of pH dependence and Zn²⁺ transmetallation

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- The transmetallation reaction of the GdL_1 and GdL_2 chelates and of the metal chelate-
- 13 decorated NPs GdL_1 @AuNps and GdL_2 @AuNPs against Zn^{2+} , was studied by
- following the time-dependent decrease of the protonic longitudinal relaxation rate, R_1 ,
- 15 (20 MHz, 25 °C) of phosphate-buffered saline solutions (PBS, pH 7.1, 10 mM),
- containing GdL_1 , GdL_2 , GdL_1 @AuNPs and GdL_2 @AuNPs ([Gd]= 1.0, 1.13, 0.42, 1.33
- 17 mM, respectively), before and after adding an equimolar amount of ZnCl₂, while
- vigorously stirring the solutions.
- The pH dependence of the relaxivity was measured by adjusting the solution pH
- 20 with aqueous diluted NaOH (0.1 M) or diluted hydrochloric acid (0.1 M), using a
- 21 Crison micro TT 2050 pH meter equipped with a Mettler Toledo 422 electrode. A
- 22 Bruker Minispec mq20 relaxometer was used for all measurements (20 MHz, 25 °C).

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MRI studies

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- Preparation of the GdL₁@AuNPs and GdL₂@AuNPs CA solutions for MRI
- 27 studies.

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- 29 The GdL₁@AuNPs and GdL₂@AuNPs CA for the MRI studies were prepared
- 30 following the procedure described above. The final nanoparticles solutions were freeze-
- 31 dried and their Gd and Au content (per mg of solid material) were determined by ICP-
- 32 OES following digestion with *aqua regia*.

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In vivo MRI studies.

environment.

The experimental protocols were approved by the appropriate institutional review committees and meet the guidelines of their responsible governmental agency. The Magnetic Resonance Imaging (MRI) experiments were all performed on a Bruker Pharmascan platform (Bruker Medical GmbH, Ettlingen, Germany) using a 7.0 T (300 MHz) horizontal-bore superconducting magnet, equipped with a ¹H selective 60 mm birdcage resonator and a Bruker gradient insert with 90 mm diameter (maximum intensity 360 mT/m). Data were acquired using a Hewlett-Packard console running Paravision software (Bruker Medical Gmbh, Ettlingen, Germany) under a LINUX

All MRI examinations were carried out on mice (n = 4, \sim 20 g body weight) anaesthetized initially by inhalation in an induction box with O_2 (1 L/min) containing 3 % isoflurane, and maintained during the experiment using a face mask allowing free breathing and 1-2 % isoflurane on O_2 . Animals were taped down into a holder, to minimize breathing - related motion, and were then placed in a heated probe, which maintained the core body temperature at 37 \pm 0.5 °C, monitored by a rectal probe. The physiological state of the animal was monitored throughout the entire experiment by a Biotrig physiological monitor (Bruker Medical GmbH, Ettlingen, Germany), using the respiratory rate and body temperature.

Solutions of $GdL_1@AuNPs$ and $GdL_2@AuNPs$ 10 mM in [Gd] were prepared by dissolving the freeze-dryed NPs in the appropriate volume of PBS buffer. The solutions were filtered through a 0.2 μ m seringe filter before injection. 10 mM Gd(DTPA) (Magnevist[®], Schering, Berlin, Germany) solutions were also prepared. The solutions were injected into the catheterized tail vein as a bolus in 20 s (0.05 and 0.1 mmol Gd kg^{-1} body weight) using an infusion pump (Panlab, Barcelona, Spain).

Regional contrast agent uptake was assessed using Dynamic Contrast Enhanced (DCE) MRI. DCE MRI experiments were performed with series of T_1 -weighted spin echo images sequentially acquired over 1 h, before and following the injection of the contrast agent 10 min after the beginning of the study. The acquisition parameters were: TR = 310 ms, TE = 10.58 ms, number of averages = 2, ten coronal slices, slice thickness = 2 mm, FOV = 5.0 x 5.0 cm, matrix = 256 x 256, 30 repetitions with a total acquision time of 119 s.

MRI data analysis.

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- 3 Data were analyzed with the public domain software Image J (http://rsbweb.nih.gov/ij/).
- 4 With the aim of comparing the pharmacokinetics obtained from different animals, the
- data were normalized by calculating the percentage of relative, rather than absolute,
- 6 enhancement:

$$RE = \frac{(I - I_0)}{I_0} \times 100$$

- 8 where I is the signal intensity at any given time after CA injection and I_0 is the intensity
- 9 before injection. Pharmacokinetic behaviour was analyzed by calculating the average
- enhancements within the different regions of interest (ROIs) placed on each one of the
- following regions: liver, kidney medulla, kidney cortex and muscle.

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Biodistribution of radiolabeled nanoparticles

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Preparation of [153Sm]L₁@AuNPs chelates for the biodistribution studies

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- 17 In these studies ¹⁵³Sm³⁺ was used as a radioactive surrogate of Gd³⁺. [¹⁵³SmCl₃] (1 mCi)
- was added to a solution of L₁@AuNPs (5 mg freeze-dried NPs) in sodium acetate buffer
- 19 (400 μL, 0.4 M, pH 5). The solution was stirred at 80 °C for 5 hours. After that, cold
- 20 SmCl₃ was added to each solution in order to obtain an equimolar Sm³⁺:chelator ratio.
- 21 The final solution was heated at 80 °C for 2 hours and left overnight at room
- 22 temperature. The radiolabeled nanoparticles were purified by size exclusion
- chromatography using a Sephadex G-10 column eluted with 0.4 M acetate buffer. The
- 24 whole colored broad band eluting from the column was collected and concentrated by
- centrifugal filtration (Centricon 10 kDa **MWCO** membrane, Millipore).

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Biodistribution studies

- 29 Groups of four animals (Wistar rat males weighting ca 200 g) were anaesthetized with
- 30 Ketamine (50.0 mg/mL)/chloropromazine (2.5%) (10:3) and injected in the femoral vein
- with ca 100 μ Ci of [153 Sm]L₁@AuNPs and sacrificed 2 and 24 hours later. The major
- organs were excised, weighed and the tissue radioactivity was measured in a γ well-

- 1 counter. Blood samples were obtained at appropriate periods of time, weighted and
- 2 radioactivity counted.
- 3 National regulations for the care and use of laboratory animals were strictly followed in
- 4 this study.

Semiempirical calculations, molecular modelling and NPs size estimates

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- 8 All calculations were performed with Mopac code⁵⁰ using the semiempirical model
- 9 Hamiltonian PM6⁵¹ and COSMO⁵² implicit water solvent model (ε = 74.8 with Gd and
- 10 Au tesserae radius taken as 0.2 nm). The length of the chelates was estimated from
- various chelate conformers averaged over several S...O and S...H top bottom distances
- within conformers (Figure SI5); ascribing an error of 0.1 nm to the estimates seems
- 13 reasonable for this methodology. The average AuNPs diameter is estimated from the
- diameter exclusion of the left and right chelates (Table SI2).

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