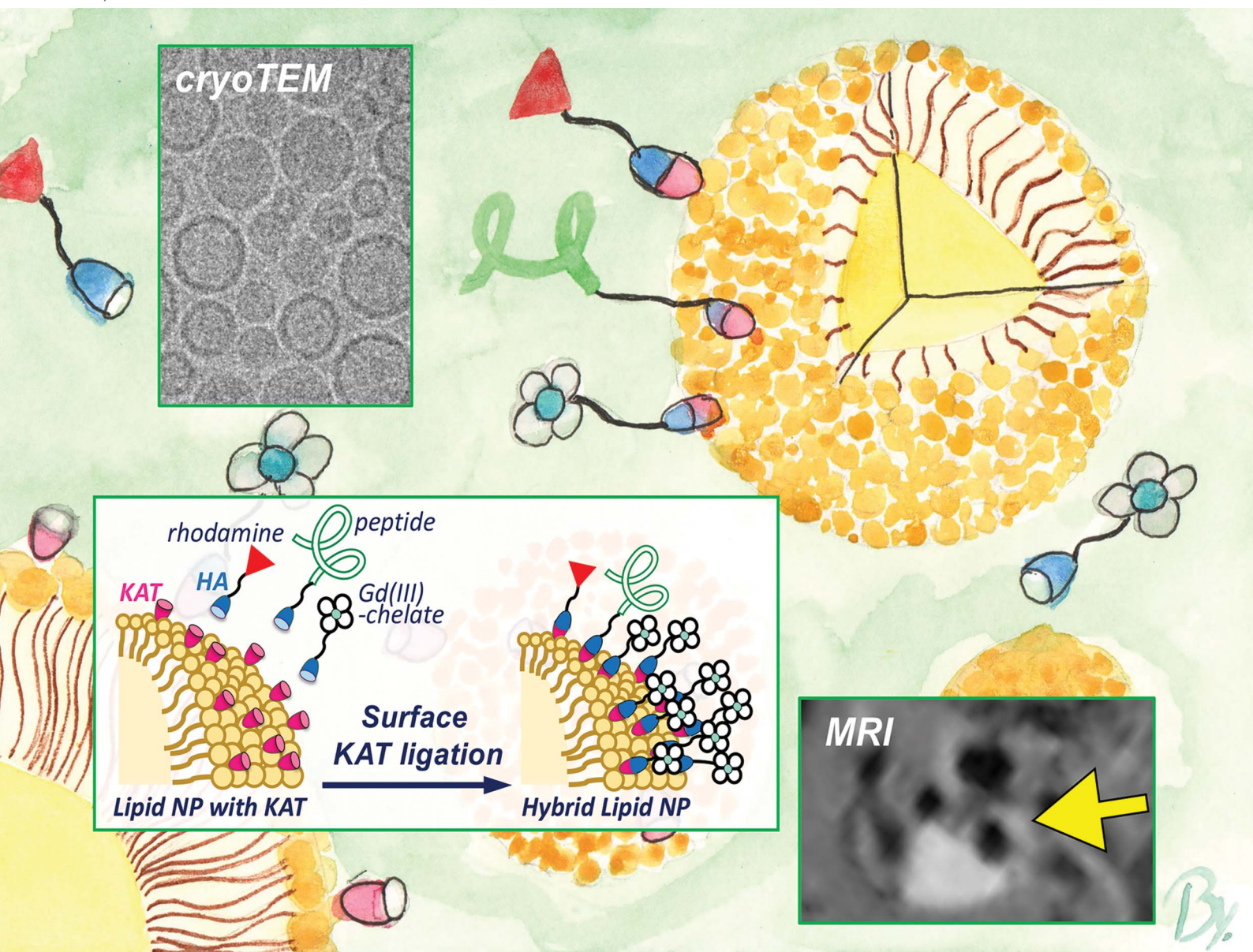


Chemical Science

rsc.li/chemical-science



ISSN 2041-6539

Cite this: *Chem. Sci.*, 2020, **11**, 11998

All publication charges for this article have been paid for by the Royal Society of Chemistry

LDL-mimetic lipid nanoparticles prepared by surface KAT ligation for *in vivo* MRI of atherosclerosis†

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Low-density lipoprotein (LDL)-mimetic lipid nanoparticles (LNPs), decorated with MRI contrast agents and fluorescent dyes, were prepared by the covalent attachment of apolipoprotein-mimetic peptide (P), Gd(III)-chelate (Gd), and sulforhodamine B (R) moieties on the LNP surface. The functionalized LNPs were prepared using the amide-forming potassium acyltrifluoroborate (KAT) ligation reaction. The KAT groups on the surface of LNPs were allowed to react with the corresponding hydroxylamine (HA) derivatives of P and Gd to provide bi-functionalized LNPs (PGd-LNP). The reaction proceeded with excellent yields, as observed by ICP-MS (for B and Gd amounts) and MALDI-TOF-MS data, and did not alter the morphology of the LNPs (mean diameter: ca. 50 nm), as shown by DLS and cryoTEM analyses. With the help of the efficient KAT ligation, a high payload of Gd(III)-chelate on the PGd-LNP surface (ca. 2800 Gd atoms per LNP) was successfully achieved and provided a high r_1 relaxivity ($r_1 = 22.0 \text{ s}^{-1} \text{ mM}^{-1}$ at 1.4 T/60 MHz and 25 °C; $r_1 = 8.2 \text{ s}^{-1} \text{ mM}^{-1}$ at 9.4 T/400 MHz and 37 °C). This bi-functionalized PGd-LNP was administered to three atherosclerotic *apoE*^{-/-} mice to reveal the clear enhancement of atherosclerotic plaques in the brachiocephalic artery (BA) by MRI, in good agreement with the high accumulation of Gd in the aortic arch as shown by ICP-MS. The parallel *in vivo* MRI and *ex vivo* studies of whole mouse cryo-imaging were performed using triply functionalized LNPs with P, Gd, and R (PGdR-LNP). The clear presence of atherosclerotic plaques in BA was observed by *ex vivo* bright field cryo-imaging, and they were also observed by high emission fluorescent imaging. These directly corresponded to the enhanced tissue in the *in vivo* MRI of the identical mouse.

Received 27th July 2020
Accepted 5th October 2020

DOI: 10.1039/d0sc04106h

rsc.li/chemical-science

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† Electronic supplementary information (ESI) available. See DOI: 10.1039/d0sc04106h

Introduction

Nanoparticles (NPs) are one of the most promising vehicles for delivery of drugs and imaging probes, and increasing numbers of NPs have recently been approved by the FDA or are currently under clinical trial.¹ Using the NP platform, larger amounts of imaging probes and drugs can be administered with a small volume *in vivo*.² By engineering the NP surface with disease-specific ligands, the local concentration of imaging probes can be increased by promoted accumulation of NPs to highlight the diseased tissue. Especially in the case of MRI contrast agents (MRI-CAs), NP-based Gd(III)-chelates generally reveal higher r_1 relaxivity in comparison to the small molecule MRI-CAs due to the decreased tumbling rate of Gd(III)-chelates, which are bound on the NP surface.²⁻⁵ For the development of such NP-based biomaterials, efficient and biocompatible surface functionalization reactions, which work under physiological conditions, are highly demanded.^{6,7} Such reactions should not cause either decomposition or aggregation of NPs,



and ideally proceed in an orthogonal manner under diluted conditions to be applicable for the attachment of biomolecules (e.g. unprotected peptide) which are often available only as diluted solutions due to their limited supply and/or insolubility.

In previous studies, NP surface functionalization was performed using chemical reactions such as Michael addition, Cu(I)-catalyzed azide-alkyne cycloaddition, and Cu-free strain-promoted click reaction.^{8–14} To complement to these reactions, in this study, we used a recently reported potassium acyltrifluoroborate (KAT) ligation.^{15,16} This amide-forming reaction between a KAT derivative and a hydroxylamine (HA) derivative was thought to be suitable for NP functionalization because of the following reasons: (1) the reaction can be performed under biorelevant conditions (in buffer, at near neutral pH, and at room temperature) with fast kinetics even under highly diluted conditions. (2) Only biocompatible reagents were involved in the reaction without providing any toxic side products.¹⁷ (3) The reaction works in an orthogonal manner and is applicable in the conjugation of biomolecules such as unprotected peptides.¹⁶ So far, KAT ligation has been mostly used in homogeneous reactions such as peptide-peptide conjugation and has not been used in non-homogeneous systems, except for our recent preliminary result of the surface functionalization of NPs with a fluorescent molecule.¹⁸

In this study, by the use of this KAT ligation, we prepared a low-density lipoprotein (LDL)-mimetic NP-type MRI-CA for the *in vivo* imaging of atherosclerotic plaques. In previous studies, we reported a natural LDL-based MRI-CA for efficient contrast enhancement of atherosclerotic plaques.^{19,20} The LDL, isolated from human blood, was modified with a synthesized Gd(III)-chelate derivative by intercalation into the lipid monolayer. The obtained LDL-based MRI-CA, with a payload of ca. 200 Gd(III)-chelates per particle, successfully visualized atherosclerotic plaques by *in vivo* MRI of *apoE*^{−/−} mice. However, in that natural LDL system, there were several disadvantages, such as infection risk and limited supply.

To make this system easily accessible, we planned to develop a synthetic LDL-mimetic lipid nanoparticle (LNP) by functionalizing the LNP core with an apolipoprotein-mimetic peptide. LNPs are biocompatible, inexpensive, and relatively easy to produce and handle, and can mimic the lipid core of LDL. They have been intensively used as a delivery vehicle for drugs and imaging probes, which were often entrapped in their hydrophobic cores,^{21–23} and have also been used in recent studies on the delivery of proteins and nucleic acids (such as siRNA).^{24–26}

Based on our preliminary results on LNP surface functionalization with fluorescein by KAT ligation,¹⁸ in the present study, we prepared a synthetic LDL-mimetic LNP. To mimic apolipoprotein in LDL (apoB100) with a large MW (514 kDa), a peptide motif with an LDL receptor-binding site^{27–29} was used. HA derivatives of this peptide (P), Gd(III)-chelate (Gd), and sulforhodamine B (R) were synthesized and subjected to the KAT ligation with the LNP core having KAT moieties (LNP-KAT) on the surface to provide covalently functionalized LNPs with peptide and imaging probes (PGd-LNP or PGdR-LNP) (Fig. 1). With the help of efficient KAT ligation, a high payload of MRI probes (ca. 2800 Gd(III)-chelates per particle) was successfully

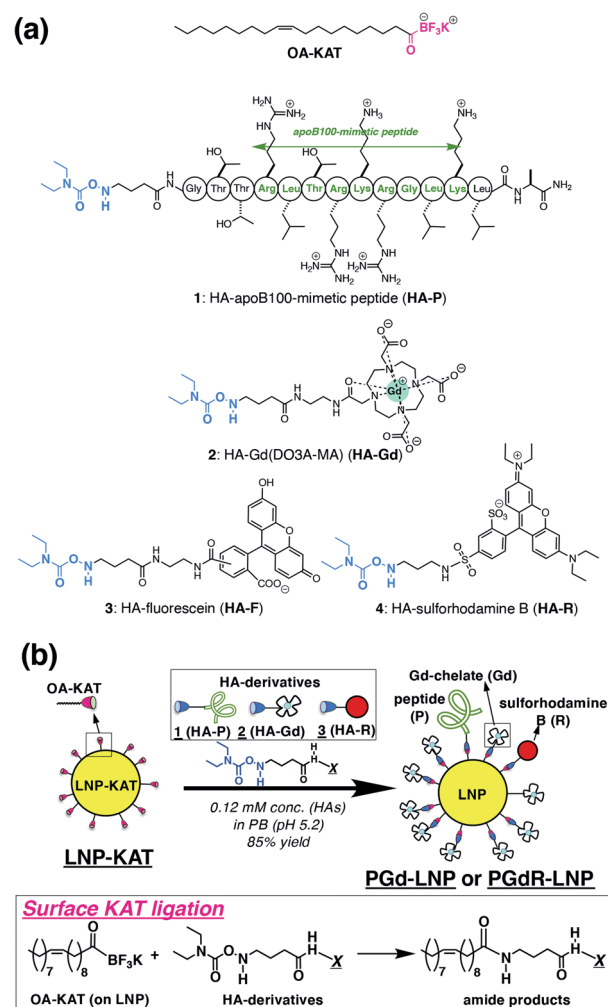


Fig. 1 (a) Chemical structures of OA-KAT and hydroxylamine derivatives; HA-apoB100-mimetic peptide (HA-P (1)), HA-Gd(DO3A-MA) (HA-Gd (2)), and HA-sulforhodamine B (HA-R (3)). (b) Schematic illustration for surface KAT ligation of the LNP-KAT nanoparticle for the preparation of the multi-functionalized LNPs (PGd-LNP or PGdR-LNP). Reactions proceeded under diluted conditions (0.12 mM) in the presence of 1.2 equiv. (in total) of HA-derivatives in pH 5.2 phosphate buffer at room temperature.

achieved providing a high r_1 relaxivity ($r_1 = 22.0$ or $8.2 \text{ s}^{-1} \text{ mM}^{-1}$ at 1.4 T/60 MHz (25 °C) or 9.4 T/400 MHz (37 °C)) and resulting in a strong enhancement of the atherosclerotic plaques in the MRI of the mouse model. The details of the preparation and characterization of the functionalized LNPs and results of *in vivo* MRI and *ex vivo* analyses are described below.

Results and discussion

Synthesis of HA derivatives of a peptide (P), an MRI-CA (Gd), and a fluorescent probe (R)

Three types of HA derivatives 1–3 were synthesized. HA-P (1), having a known binding motif to the LDL receptor in the apoB100 protein, was synthesized as an apolipoprotein-mimetic by a standard solid phase peptide synthesis (Scheme S1 in the ESI†). The synthesized *N*-Boc-HA-peptide on resin (S3) was treated with TFA



and TIPS, for simultaneous deprotection of the *N*-Boc group of the HA moiety on the *N*-terminus, deprotection of the amino acid side chains on the peptide, and cleavage of the whole peptide from resin, to successfully provide an all-deprotected **HA-P** (**1**).³⁰ The obtained **1** was purified by HPLC (Fig. S1 in the ESI†). The *N*-Boc precursor for HA-Gd(DO3A-monoamide) (or HA-Gd(DO3A-MA), **HA-Gd** (**2**)) was synthesized from cyclen *via* several synthetic intermediates (Scheme S3†). The *N*-Boc precursor of HA-sulforhodamine B (**HA-R** (**3**)) was synthesized as previously reported.³¹ These *N*-Boc HA derivatives (precursors for **2** and **3**) were purified by HPLC and treated with TFA for the *N*-Boc deprotection immediately before subjecting them to surface KAT ligation with 5% or 10% **LNP-KAT**.

Preparation of LNP-KAT and surface KAT ligation

The platform **LNP-KAT** particle was prepared from a mixture of three commercially available lipids (phosphatidylcholine (PC), triolein (TO), and cholesteryl oleate (CO)) and one synthetic lipid (**OA-KAT**, 5 or 10 mol% of total lipids) as previously reported (Section 2.1 in the ESI†).¹⁸ Briefly, a mixture of the four lipids above (Fig. S31 in the ESI†) was treated under sonication and extrusion in pH 8.0 Tris-HCl buffer to form **LNP-KAT** (5 or 10%).³² The obtained **LNP-KAT** dispersion was washed by spin filtration (50 kDa MW cut-off) to remove the free lipid molecules – which were not incorporated into the NPs – and stored in pH 8.0 Tris-HCl buffer (with 10 mM of KF) as a stable dispersion (Fig. S36†).

The surface KAT ligation of **LNP-KAT** and a mixture of HA derivatives (Fig. 1b) was carried out in pH 5.2 phosphate buffer due to its known faster reaction kinetics at lower pH.¹⁷ Particularly in this study, nanoparticle aggregation was suspected due to the electrostatic interaction of the negatively charged **LNP-KAT** surface and positively charged peptide (**HA-P**, Fig. 1a),³³ and the reaction was carried out under diluted conditions (120 μM of HA reagents) with a minimum amount of HA reagents (in total 1.2 equiv. relative to the **OA-KAT** molecules incorporated into the **LNP-KAT** particle) at room temperature. For the MRI tests, a bi-functionalized **PGd-LNP** with peptide (**P**) and Gd(DO3AMA) (**Gd**) was prepared by the reaction of **LNP-KAT** (5%) with a mixture of **HA-P** (**1**) and **HA-Gd** (**2**). For both MRI

and *ex vivo* cryo-fluorescence imaging, a triply functionalized **PGdR-LNP** was prepared by the addition of a mixture of **1**, **2**, and **HA-R** (**3**). The reaction was carried out simply mixing the HA derivatives with **LNP-KAT** (10%) in buffer at room temperature. After surface KAT ligation, each obtained LNP dispersion was thoroughly washed with PBS(–) by spin-filtration (MW cutoff: 50 kDa) to carefully remove the unreacted and non-specifically-bound HA derivatives **1–3** from the particle surface of **PGd-LNP** or **PGdR-LNP** before subjecting it to the following particle characterization and *in vivo* tests.

Reaction efficacy of surface KAT ligation

The reaction yield of surface KAT ligation was estimated by B and Gd quantification in **LNP-KAT** (5%) (starting material) and **PGd-LNP** (product) by ICP-MS analysis (Table 1). In the ligation reaction, a mixture of **1** and **2** was added in a mole ratio of 10 : 90. ICP-MS analyses of the particles were carried out after digestion by HNO₃. As shown in Table 1, the B contents in **LNP-KAT** (5%) and **PGd-LNP** were 514 and 227 μM suggesting that *ca.* 56% of the KAT group in **LNP-KAT** (B3 in the bottom figure) was consumed during the reaction. This result suggested that a sufficient number (more than half) of **OA-KAT** molecules were available for the reaction, presumably being embedded in the surface of the **LNP-KAT** particle with the help of amphiphilicity of the **OA-KAT** molecule, and efficiently exposed to the HA reactants in the external bulk solution. From the consumed B amount (B3 = (B1 + B2) – B4, 287 μM) and the gained Gd content (217 μM) in the reaction, taking into account that the HA reactant was a mixture of **1** and **2** (10 : 90), the conversion yield of surface KAT ligation was calculated to be 84% ((Gd × 100/90)/B3 × 100). The result indicated that the surface KAT ligation of the **LNP-KAT** (5%) particle proceeded in a sufficiently high yield under a diluted concentration (120 μM) and with a minimum amount of HA molecules (1.2 equiv.) without having side reactions. To confirm that these changes of B and Gd contents corresponded to the covalent reaction, the detection of ligation products was attempted by MALDI-MS as described below.

Table 1 Reaction yield of surface KAT ligation estimated from the contents of B and Gd in **LNP-KAT** (5%) and **PGd-LNP** measured by ICP-MS

	Content of B [μM]	Content of Gd [μM]	Consumed B [%]	Reaction yield [%]	Conversion yield [%]
LNP-KAT	514 ^a	—	—	—	—
PGd-LNP	227 ± 20 ^b	217 ± 5.2 ^c	56 ± 3.8 ^d	47 ± 1.0 ^e	85 ± 6.8 ^f

^a Total amount of B in the **LNP-KAT** (5%) before the reaction (B1 + B2). ^b Remaining amount of B in **PGd-LNP** after the reaction (B4). ^c Amount of Gd in the **PGd-LNP** after the reaction. ^d Calculated by (B1 + B2) – B4. ^e Reaction yield from Gd amount in **PGd-LNP** relative to the total amount of B in **LNP-KAT** (Gd/(B1 + B2) × 100). ^f Conversion yield from Gd amount in **PGd-LNP** relative to consumed B during the reaction (Gd/B3 × 100).



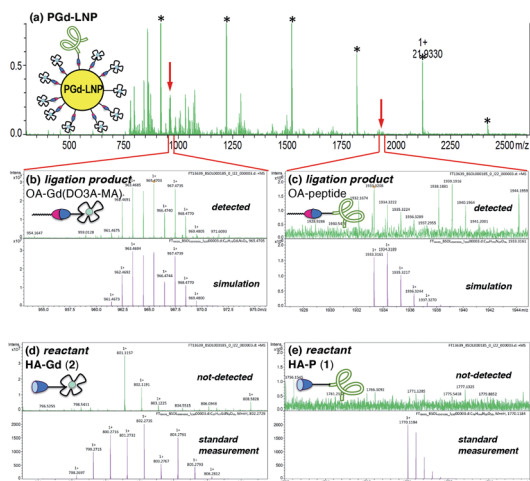


Fig. 2 (a) HR-MALDI-MS spectrum of PGd-LNP (matrix: HCCA, *standard for calibration). (b and c) Expansion at the molecular ion peak of OA-Gd(DO3A-MA) (m/z : $[M + H]^+$ found, 965.4703) (b) and OA-peptide (m/z : $[M + H]^+$ found, 1933.3208) (c) with simulation. (d and e) Expansion around the molecular ion peaks of HA-Gd(DO3A-MA) 2 (d), and HA-peptide 1 (e) with standard measurements of 2 (m/z : $[M + H]^+$ found, 802.2757) and 1 (m/z : $[M + H]^+$ found, 1770.1185).

Detection of ligation products in PGd-LNP by MALDI-MS

MALDI-MS analysis of PGd-LNP was employed for the detection of the ligation products (OA-peptide and OA-Gd(DO3A-MA)). An aliquot of PGd-LNP dispersion was analyzed using HCCA (α -cyano-4-hydroxycinnamic acid) as a matrix, and molecular ion peaks corresponding to the ligation products were clearly observed (Fig. 2a–c). Importantly, no trace of HA derivatives 1 and 2 was detected (Fig. 2d and e), indicating that there were no unreacted 1 and 2 that were physically adsorbed on the PGd-LNP surface. These results suggested that the ICP-MS results of the change in B and Gd contents shown in Table 1 corresponded to the covalent surface reaction.³⁴

Characterization of PGd-LNP by DLS and microscopy

The surface functionalization of NPs often alters their properties including charges and hydrophobicity or hydrophilicity, especially at the interface with the bulk water, causing damage to NPs such as aggregation and degradation.³⁵ Since this damage is critical in biological tests, the sizes and morphologies of the particles were analyzed by dynamic light scattering (DLS) and microscopic methods. As shown in DLS, both nanoparticles before (LNP-KAT (5 or 10%)) and after the reaction (PGd-LNP (5%) or PGdR-LNP (10%)) revealed a single distribution of diameters with a mean of *ca.* 50 nm (Fig. 3), which is in a suitable size range for *in vivo* application.³⁶ Importantly, no detectable aggregation was observed in all particles during the surface reaction and the LNPs were stably dispersed upon storage for at least 14 days at 4 °C in either pH 8.0 Tris-HCl buffer or pH 7.4 PBS(–) (Fig. S36†).

The dispersion status and the morphologies of the LNPs were further analyzed by AFM and cryoTEM (Fig. 4). For AFM imaging, an aliquot of LNP-KAT or PGd-LNP dispersion was

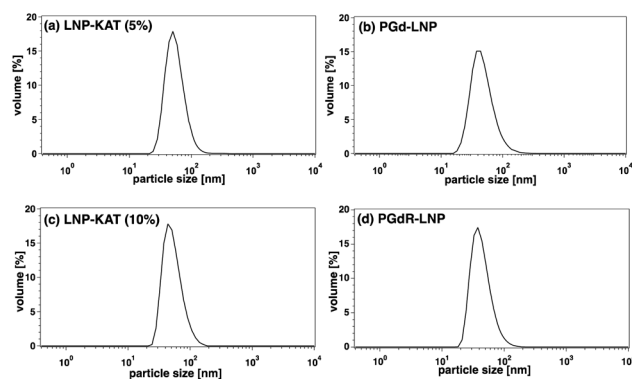


Fig. 3 DLS of LNPs before and after the surface KAT ligation. (a) LNP-KAT (5%) with 5 mol% of OA-KAT (mean: 54.5 nm, width: 20.8 nm). (b) PGd-LNP prepared from LNP-KAT (5%) (mean: 45.7 nm, width: 19.3 nm). (c) LNP-KAT (10%) with 10 mol% of OA-KAT (mean: 51.7 nm, width: 20.6 nm). (d) PGdR-LNP prepared from LNP-KAT (10%) (mean: 42.6 nm, width: 16.8 nm).

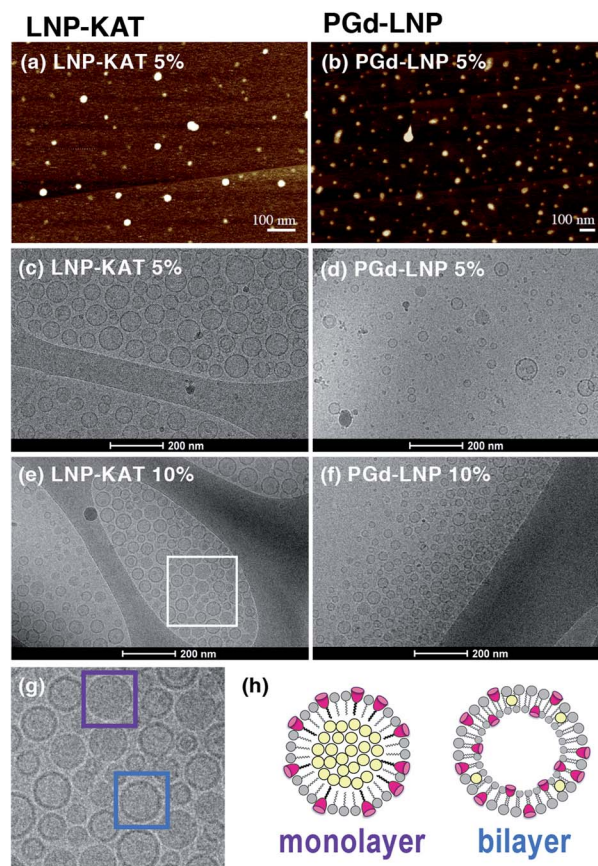


Fig. 4 Microscopic images of LNPs. (a and b) AFM images of LNP-KAT (5%) (a) and PGd-LNP prepared from LNP-KAT (5%) (b). (c–f) CryoTEM images of LNP-KAT (5%) (c), PGd-LNP prepared from LNP-KAT (5%) (d), LNP-KAT (10%) (e), and PGd-LNP prepared from LNP-KAT (10%) (f). (g) Expansion of the cryoTEM image of LNP-KAT (10%) (from the area marked as a white square in (e)). (h) Illustration of LNP-KAT with monolayer and bilayer morphologies. The particles marked with purple and blue squares revealed a relatively dark solid morphology and a relatively light inside with a darker edge, respectively.



Table 2 Determination of the Gd payloads in PGd-LNP (5%) and PGd-LNP (10%) prepared from LNP-KAT (5%) and LNP-KAT (10%)

Samples	No. of NPs per mL (NTA)	Conc. of Gd ³⁺ [μM] (ICP-MS)	No. of Gd ³⁺ species per LNP
PGd-LNP (5%) ^a	$4.6 \pm 0.26 \times 10^{13}$	216.9 ± 5.2	2841 ± 216
PGd-LNP (10%) ^b	$4.9 \pm 0.32 \times 10^{13}$	374.8 ± 7.3	4638 ± 200

^a Prepared from LNP-KAT (5%). ^b Prepared from LNP-KAT (10%).

deposited onto a mica surface and subsequently fixed with formalin. As shown in Fig. 4a and b, uniformly dispersed LNPs were observed by AFM without significant aggregation. The cryoTEM images were taken on the LNP samples, prepared by plunge freezing on grids in buffer. Measurements were performed at $-180\text{ }^\circ\text{C}$ without staining. As shown in Fig. 4c–f, all the LNP samples (LNP-KAT and PGd-LNP with 5 or 10% of KAT) exhibited well-dispersed spherical morphologies having *ca.* 30–70 nm diameter.³⁷ Upon closer observation of individual LNPs (Fig. 4g), interestingly, 2 types of LNPs were found, one with a relatively darker solid morphology (marked with a purple square) and the other with a lighter inside and darker outer edge with *ca.* 5–10 nm thickness (marked with a blue square). These two types of NPs presumably corresponded to the one with a lipid monolayer and the other with a lipid bilayer as illustrated in Fig. 4h (see also Fig. S37[†]).^{38,39} This cryoTEM observation of a mixture of monolayer- and bilayer-type LNPs was in line with the results of the yield of surface KAT ligation – about half of KAT in LNP-KAT (5%) was consumed during the surface KAT ligation (see Table 1). The other half of B, remaining after surface KAT ligation, at least in part, corresponded to a migrated KAT moiety intercalated in the inner layer of the bilayer-type LNP-KAT (Fig. 4h).

Estimation of the Gd(III)-chelate payload in PGd-LNP

The Gd(III)-chelate payload per particle is an important factor affecting the r_1 relaxivity and MRI contrast *in vivo*. The number of attached Gd(III)-chelates per PGd-LNP particle was estimated from (1) ICP-MS data of Gd concentration and (2) nanoparticle tracking analysis (NTA) data to count the number of particles (Tables 2 and S37[†]). From the results, Gd(III)-chelate payloads of PGd-LNP (5%) and PGd-LNP (10%) particles, prepared respectively from LNP-KAT (5%) and LNP-KAT (10%), were found to be *ca.* 2800 and 4600 per particle. Such large numbers of Gd(III)-chelate payload on PGd-LNP indicate the potential of these particles as an efficient delivery system of Gd(III)-chelates while providing high r_1 relaxivity and corresponding strong contrast enhancement in MRI.

Relaxivity of PGd-LNP

The proton longitudinal relaxation rate ($1/T_1$) of PGd-LNP (5%) was measured to obtain r_1 relaxivity. Under standard measurement conditions (1.4 T/60 MHz at $25\text{ }^\circ\text{C}$), the r_1 relaxivity of PGd-LNP (5%) was $22.0\text{ s}^{-1}\text{ mM}^{-1}$. This was much higher than those of small molecule MRI-CAs, such as ProHance® (Bracco Spa, Milano, Italy) with $r_1 = 4.0\text{ s}^{-1}\text{ mM}^{-1}$ (1.4 T/60 MHz).⁴⁰ It is known that the r_1 relaxivity value is related to the rotational

correlation time of the Gd(III)-chelate and a slowed rate causes higher r_1 in comparison to the freely rotating small molecule Gd(III)-chelate especially at a lower magnitude.⁴¹ This is commonly observed by linking the Gd(III)-chelate molecules to macromolecules, polymers, and nanoparticles.^{3,4} The r_1 value observed in the current PGd-LNP particle was also higher than that of most of the previously reported LNP-type Gd(III)-based MRI-CAs,^{42–44} including that of our previous natural LDL-based MRI-CA ($r_1 = 20.1\text{ s}^{-1}\text{ mM}^{-1}$) with a Gd payload of 210 per particle.²⁰ This was presumably, at least in part, due to the high payload of the Gd(III)-chelate (*ca.* 2800) per particle, which was successfully enabled by the fast kinetics of KAT ligation used in the current study. More importantly, the r_1 of PGd-LNP measured at 9.4 T/400 MHz (at $37\text{ }^\circ\text{C}$) was $8.2\text{ s}^{-1}\text{ mM}^{-1}$, showing significantly higher value than that of the small molecule ProHance® ($r_1 = 2.6\text{ s}^{-1}\text{ mM}^{-1}$ at 9.4 T/400 MHz at $37\text{ }^\circ\text{C}$),⁴⁶ although the NP effect providing a higher r_1 value is generally weaker at higher magnetic field.⁴⁵ These measurement conditions were the same as those for *in vivo* imaging in this study and sufficiently high contrast was expected in the *in vivo* imaging in the presence of PGd-LNP.

In vivo MRI of an atherosclerotic mouse model injected with PGd-LNP

Based on the successful LNP functionalization with apoB100-mimetic peptide (P) and Gd(III)-chelate (Gd) providing a high r_1 value, PGd-LNP (5%) was injected into an atherosclerotic mouse model for *in vivo* MRI. An *apoE*^{−/−} mouse fed on a high fat diet for 2 months was used as a mouse model.⁴⁷ After an initial baseline MRI measurement of three atherosclerotic *apoE*^{−/−} mice (no. 1–3), the PGd-LNP (5%) dispersion, which was prepared from LNP-KAT (5%), was injected *via* the tail vein in a Gd(III)-chelate dose of 0.018–0.051 mmol kg^{−1} (Table 3).⁴⁸ The MR images were acquired using a 9.4 T horizontal bore MR spectrometer and at 48 h post-injection, based on our previous results of atherosclerotic plaque imaging using a natural LDL-based MRI-CA.²⁰ The thoracic aorta, including the aortic arch

Table 3 *Ex vivo* ICP-MS analysis of the aortic arch from *apoE*^{−/−} (no. 1–3) 48 h post-injection of PGd-LNP (5%)

No.	Mouse body weight	Dose of Gd [mmol kg ^{−1}]	Dry weight of tissue used [mg]	Gd content by ICP-MS [ppm]
1	34 g	0.018	0.8	3.66
2	49 g	0.051	0.7	14.6
3	38 g	0.042	0.9	4.26



and its three branches (brachiocephalic artery (BA), left carotid artery (LC), and left subclavian artery (LS)), was visualized on coronal scout images, then multiple axial images were acquired.

As shown in Fig. 5, clear enhancements in the wall of the BA near its branching from the aortic arch were observed in all mice no. 1–3 (yellow arrow in Fig. 5c, d, g, h, k and l). These black blood (BB) images are useful for atherosclerotic plaque detection since the blood in the three arteries (BA, LC and LS, red arrows in Fig. 5) appear black due to the suppression of the blood signal, leading to clear delineation of the artery wall. The assignment of the arteries was confirmed by the comparison with white blood (WB) images, where the trachea appears dark and other arteries appear white (Fig. S39–S41 in the ESI†). In *apoE*^{−/−} mice, atherosclerotic plaques are often formed at the branching point of BA from the aortic arch, where the MRI enhancement was detected in this study. Also, a similar enhancement was previously observed in this region in our previous reports with a natural LDL-based MRI-CA.^{19,20}

The normalized signal enhancement (% NSE) in mice no. 1 and 2 was quite high (125 and 103%). The % NSE of mouse no. 3 was not obtained due to the overlapping of the artery wall with the enhancement of the thymus (pink arrows) although a strong

enhancement in the BA wall was apparent (Fig. 5i). These enhancements in BA were not observed in the MR images of the mice injected with small molecule ProHance® (Gd^{3+} dose = 0.085 mmol kg^{−1}) (Fig. S39–S41 in the ESI,† showing % NSE of 53 or 54%) and are apparently higher than those in our previous natural LDL-based particle experiment providing % NSE of ca. 75%.²⁰

Ex vivo ICP-MS analysis of the aorta of *apoE*^{−/−} model mice

After MRI, the aortic arch with its three branches (BA, LC, and LS) just above the branching points were harvested from mice no. 1–3 and analyzed by ICP-MS for Gd contents. During dissection, plaques were clearly observed in all mice especially in the BA region. As shown in Table 3, significant accumulation of Gd^{3+} in the aorta was observed in all three *apoE*^{−/−} mice. These data were comparable to or higher than our previous results on a natural LDL-based MRI-CA, showing the Gd content in the aortic arch (ca. 5.4 ppm).⁴⁹ Mouse no. 2 with the highest Gd accumulation (14.6 ppm) in its aorta had the highest body weight and atherosclerotic plaque burden, which was observed during dissection. The increased accumulation of Gd in the aorta was because of selective accumulation of the high payload of Gd-chelate on the **PGd-LNP** (5%), which was facilitated by KAT ligation, with affinity to the atherosclerotic plaques in the *in vivo* system.

In vivo MRI and ex vivo cryo-fluorescence imaging of *apoE*^{−/−} injected with **PGdR-LNP**

Although the atherosclerotic plaque progression in the region of enhancement was obvious during dissection of the mice, it is important to directly correlate MRI enhancement and accumulation of nanoparticles by an *ex vivo* study to show the direct connection between LNP accumulation in plaques and MRI enhancement. For this purpose, together with **P** and **Gd**, we added a fluorescent probe (sulforhodamine B, **R**) to the LNPs to prepare triply labelled **PGdR-LNP**, enabling the *ex vivo* fluorescent imaging of the mouse tissues. To achieve a higher payload of Gd(III)-chelate, **LNP-KAT** (10%) was used as a starting material and the obtained **PGdR-LNP** (10%) was characterized by fluorescence spectroscopy and ICP-MS for sulforhodamine B and Gd content, respectively (Table S4†). Also, DLS showed no aggregation of NPs (Fig. S35†).

An atherosclerotic *apoE*^{−/−} mouse (no. 4), injected with **PGdR-LNP**, in Gd(III)-chelate doses of 0.109 mmol kg^{−1}, was subjected to MRI. In this experiment, initially, MRI was compared at 24 and 48 h post-injection to optimize the MRI timing that provides the best enhancement. As shown in Fig. 6a, a clear enhancement of the BA branching region in the aortic arch was observed at both 24 and 48 h post-injection (yellow arrow). In comparison to the experiments of mice 1–3 that received **PGd-LNP** (5%) prepared from **LNP-KAT** (5%) (Fig. 5), a higher enhancement was observed in mouse 4 at 48 h post-injection of **PGdR-LNP** (10%) prepared from **LNP-KAT** (10%) (Fig. 6). By quantitative % NSE analysis, we found that the enhancement in BA was higher at 24 h post-injection than at 48 h (Fig. 6a), suggesting a partial washout of the contrast

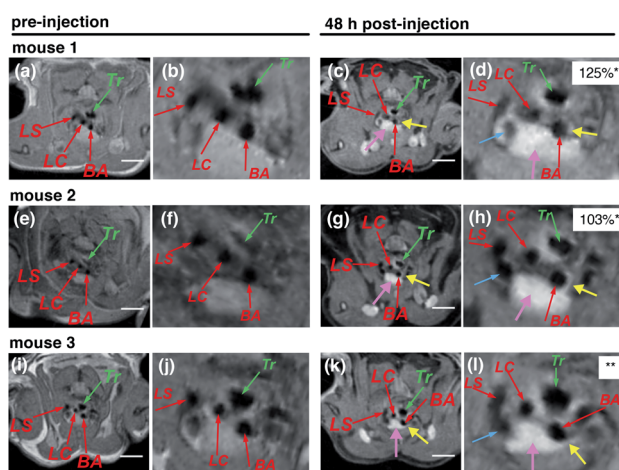


Fig. 5 *In vivo* MR images of *apoE*^{−/−} mice (no. 1–3) in the regions of arteries taken pre-injection (a, e and i) and 48 h post-injection of **PGd-LNP** (5%) with doses of 0.018, 0.051, and 0.042 mmol Gd(III)-chelate kg^{−1} (for no. 1, 2, and 3, respectively) (c, g and k) with their expanded images (b, f, j, d, h and l). Three branches of the aortic arch were identified by red arrows: BA = brachiocephalic artery; LC = left carotid artery; LS = left subclavian artery. Tr = trachea. Yellow arrows point to the enhancement of atherosclerotic plaques. Pink arrows point to the enhancement in the thymus. Green arrows point to the trachea. The enhancement surrounding the trachea is likely due to paratracheal lymph nodes, which contain macrophages taking up the **PGd-LNP**. Blue arrows point to the left cranial vena cava. *The normalized signal enhancement (% NSE), calculated using the following equation: % NSE = 100% × (($I_{\text{wallpost}}/I_{\text{musclepost}}$)/($I_{\text{wallpre}}/I_{\text{musclepre}}$) − 1). I_{wallpre} and I_{wallpost} are the signal intensity from the artery wall pre- and post-injection, respectively. $I_{\text{musclepre}}$ and $I_{\text{musclepost}}$ are defined similarly for the nearby skeletal muscle, which was used as reference tissue (see the ESI† for the details). **% NSE was not obtained due to the overlapping of the brachiocephalic artery wall with the enhanced thymus. White bars correspond to 5 mm.



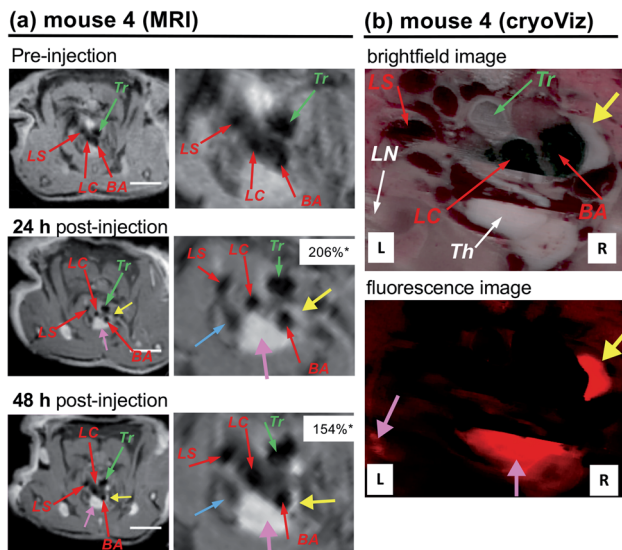


Fig. 6 (a) *In vivo* MRI images of atheroma in the arteries of an *apoE*^{-/-} mouse (no. 4) taken at pre-injection and at 24 and 48 h post-injection of PGdR-LNP (10%) (dose: 0.109 mmol Gd(III)-chelate kg⁻¹) with their expanded images. (b) CryoViz images of mouse no. 4 at 24 h post-injection of PGdR-LNP (10%) with brightfield (upper) and fluorescence (lower) images. Three branches of the aortic arch were identified by red arrows: BA = brachiocephalic artery; LC = left carotid artery; LS = left subclavian artery. Yellow arrows point to the enhancement of atherosclerotic plaques and corresponding fluorescence emission. Pink arrows point to enhancement in the thymus (Th) and lymph node (LN). Green arrows point to the trachea (Tr). Blue arrows point to the left cranial vena cava. *The normalized signal enhancement (% NSE), calculated using the following equation: % NSE = 100% × ((*I*_{wallpost}/*I*_{musclepost})/(*I*_{wallpre}/*I*_{musclepre}) - 1). *I*_{wallpre} and *I*_{wallpost} are the signal intensity from the artery wall pre- and post-injection, respectively. *I*_{musclepre} and *I*_{musclepost} are defined similarly from muscle (see the ESI† for the details). White bars correspond to 5 mm.

media between these time points. Based on this result, two weeks after the first MRI study, mouse no. 4 was injected again with PGdR-LNP (10%) and scanned by MRI at 24 h post-injection. After the MRI study, the mice were euthanized for *ex vivo* study by cryo-imaging of the whole mouse.

In MR images of mouse no. 4 (Fig. 6a), a strong enhancement in the BA wall at the aortic arch branching point was clearly observed (yellow arrows) presumably due to the significant accumulation of PGdR-LNP at this location. The brightfield cryo-imaging of identical mice (Fig. 6b, upper) clearly indicates plaque deposition in the inner wall of BA (yellow arrow) which completely overlapped with the MRI contrast enhancement location. By fluorescent cryo-imaging (Fig. 6b lower), a strong sulforhodamine B fluorescence was observed in the plaque position (yellow arrow). This result confirmed that PGdR-LNP particles were accumulated in the plaque of BA enabling the *in vivo* MRI diagnosis of atherosclerosis with high enhancement.

Other LNP-type MRI contrast agents, especially HDL-based ones, have been previously reported with successful detection of atheroplaques in abdominal aorta.^{50–52} In this study, the use of an LDL-mimetic LNP contrast agent with a high payload of Gd(III)-chelate produced by efficient KAT ligation and cardiac-

gated MRI to suppress the cardiac motion interference enabled the visualization of atherosclerotic plaques in the aortic arch by high-resolution MRI. Furthermore, we validated the MRI findings by confirming PGdR-LNP accumulation in enhanced anatomical structures using the state-of-the-art cryoViz technique.

Experimental

Synthesis

Details of the synthesis of OA-KAT and HA derivatives 1–3 are described in the ESI† with corresponding spectral data including all the synthetic intermediates. The HA-apoB100-mimetic peptide 1 was synthesized by a standard solid phase peptide synthesis (SPPS) and purified by reversed phase HPLC (Shiseido Capcell Pak C18 column, CH₃CN–H₂O in the presence of 0.1% TFA) to be obtained in a yield of 37% for the total peptide synthesis and characterized by MALDI-MS; HRMS (MALDI⁺) *m/z*: [M + H]⁺ calcd for C₇₆H₁₄₅N₂₈O₂₀⁺: 1770.1184; found, 1770.1185. HA-Gd(DO3A-MA) 2, and HA-sulforhodamine B 3 were synthesized by a common multistep organic synthesis.

Preparation of the LNP-KAT particle

The lipid nanoparticle (LNP) with KAT moieties (LNP-KAT) was prepared in a similar way to that in our previous report⁸ using the same lipid components for LNP preparation as those reported by Halbert and co-workers^{29,53–56} and Forte and co-workers.^{57,58} Briefly, to a mixture of phosphatidyl choline (33.8 mg, 44.3 μmol), triolein (29.2 mg, 33.0 μmol) and cholesterol oleate (9.4 mg, 16.5 μmol) in a mixture of CHCl₃, MeOH, and acetone (2 : 1 : 1, v/v, 90 mL), a solution of OA-KAT (1.9 mg, 5 μmol) in a mixture of MeOH and acetone (1 : 1, v/v, 30 mL) was added. The solvents were slowly evaporated to provide a lipid thin film. Subsequently, the film was hydrated with 4 mL of Tris–HCl buffer (10 mM, pH 8.0, with KF (10 mM) and BHT (1 mg L⁻¹)), sonicated, extruded with two stacked polycarbonate membrane filters (pore sizes: 0.05 and 0.1 μm) and filtered (0.45 μm). The filtrate was washed with Tris–HCl buffer by spin-filtration (50 kDa MW cut-off) to remove free lipid components, which were not incorporated in the particle, to yield 4 mL of LNP-KAT particle suspension in Tris–HCl buffer. The LNP-KAT was stored in Tris–HCl buffer at 4 °C until its use in the ligation reaction.

On-particle KAT ligation of LNP-KAT with HA derivatives

To a mixture of 10 mM KF in pH 5.8 phosphate buffer (10 mM) and 0.1 M HCl (400 μL), aqueous solutions of HA-peptide 1 (0.48 μmol) and HA-Gd(DO3A-MA) 2 (4.32 μmol) were added. Subsequently, the LNP-KAT particle suspension in Tris–HCl buffer was added and the reaction mixture (pH 5.2) was stirred overnight at room temperature. The resulting PGd-LNP particle was subjected to spin-filtration to provide 1.5 mL of suspension, which was washed with 5 mL of PBS (–) pH 7.4 three times to remove the unreacted 1 and 2 and subjected to particle characterization.



Quantification of B and Gd contents in LNP-KAT and PGd-LNP

ICP-MS and ICP-OES were employed for the quantification of B and Gd contents in LNP-KAT and PGd-LNP for the particle characterization and estimation of *on-surface* KAT ligation yield. Each particle suspension was dried, digested in conc. HNO₃, dried again, and dissolved in 2% HNO₃ before being subjected to ICP-MS or ICP-OES analyses. Measurements were performed on an Element XR sector-field ICP-MS (ThermoFisher Scientific, Waltham, MA, USA) or an Ultima Expert ICP-OES (Horiba, Ltd., Kyoto, Japan).

MALDI-MS analysis of PGd-LNP

The MALDI-MS analysis of PGd-LNP was performed on a Bruker Solarix-FTICR 9.4 T MS (Bruker Daltonics, Bremen, Germany) using α -cyano-4-hydroxycinnamic acid (HCCA) as a matrix.

DLS analysis of PGd-LNP

The hydrodynamic size of the nanoparticles was estimated by dynamic light scattering (DLS) on a Malvern Nano-ZetaSizer (Malvern Instruments Ltd., Worcestershire, UK) in Tris-HCl buffer (10 mM, pH 8.0) or phosphate buffer (10 mM, pH 8.0) at 25 °C.

AFM imaging of nanoparticles

Each nanoparticle suspension was deposited onto freshly cleaved mica (Ted Pella Inc., Redding, CA, USA) and fixed by the addition of 10% formalin. AFM imaging was performed on a Dimension Icon AFM (Bruker Nano Surfaces) in the Peak Force Tapping mode in liquid using an SNL 10-A cantilever (Bruker Co.).

CryoTEM imaging of nanoparticles

The microscopic characterization of the particles in their hydrated state was carried out on a Tecnai F20 cryo-TEM (FEI/Thermo Fisher Scientific) using samples on 300-mesh lacey carbon-coated copper grids (Quantifoil Micro Tools GmbH, Jena, Germany). A mixture of liquid ethane/propane was used for sample vitrification. The grids were then transferred on a Gatan cryo-transfer holder (Gatan, Inc., Pleasanton, CA, USA) into a microscope and kept at -180 °C during observation. Micrographs were recorded under low dose conditions (<500 e⁻ nm⁻²) using a Falcon II 4K Camera (FEI Thermo Fisher Scientific), operating the microscope at 200 kV acceleration voltage in bright field mode.

Relaxivity measurement of PGd-LNP

Proton relaxivities at 60 MHz were measured on a Bruker WP80 NMR electromagnet adapted to variable field measurements and controlled by a Stellar SMARTracer PC-NMR console (Stellar s.r.l., Pavia, Italy). The temperature was monitored by a VTC91 temperature control unit and maintained by a gas flow. The temperature was determined by previous calibration with a Pt resistance temperature probe. Relaxivities at 400 MHz were measured on a Bruker AVANCE NMR spectrometer. The

temperature was calculated according to previous calibration with ethylene glycol and methanol.⁵⁹ The Gd concentration of the sample was 193 μ M.

NTA measurement of PGd-LNP

The nanoparticle tracking analysis (NTA) was recorded on a Zetaview® (Particle Metrix, Meerbusch, Germany) to obtain the number of particle dispersions. An aliquot (2 mL) of sample was injected into the sample cell and illuminated with a 405 nm laser sheet (45 mW). The 11 independent positions in the sample cell were imaged by a video camera mounted on a 10 \times microscope objective oriented 90° to the laser sheet, probing 0.4 nL of volume at each position. The aspect ratio of the camera is 640 pixels \times 480 pixels, with an effective pixel size of 0.7 μ m per pixel.

ICP-MS analysis of the dissected aorta

All the animal procedures were approved by the Institutional Animal Care and Use Committee (IACUC) of the University of Pennsylvania. The male *apoE*^{-/-} mice were purchased from the Jackson Laboratory and fed with HFD (42% kcal from fat) for two months. To the *apoE*^{-/-} model mice, PGd-LNP was injected. After 48 h, the mice were euthanized and perfused with PBS before harvesting the aortas for the determination of Gd content by ICP-MS, performed in the Toxicology Core Lab at the New Bolton Center of Veterinary Hospital of the University of Pennsylvania.

In vivo imaging using the atherosclerotic mouse model

The *apoE*^{-/-} model mice were used for *in vivo* MRI. Under anaesthesia, the PGd-LNP or PGdR-LNP particle was injected *via* a catheter placed in the tail vein of the mouse. All MRI was performed on a 9.4 T horizontal bore MR spectrometer (DirectDrive®, Agilent, Palo Alto, CA) equipped with a 12 cm (ID) gradient coil. The mouse was positioned prone in a quadrature volume radio frequency (RF) coil (ID = 3.5 cm, length = 8 cm, m2m imaging/Polarean) tuned to ¹H resonance frequency (400 MHz). During imaging, the mouse was sedated with 0.8–1% of isoflurane mixed with air (flow rate = 1 L min⁻¹). ECG, respiration, and core temperature of the mouse were monitored using an MRI-compatible vital sign monitoring system (SA Inc, Long Island, New York). The rectal temperature was maintained at 37 \pm 0.2 °C by a feedback loop that turns on/off warm air directed into the magnet bore. Scout images were acquired to capture the aortic arch with three rising branches (LC, LS and BA). Then the image plane was placed to cut through the three aortic branching points. This image plane was used to acquire the “White Blood” (WB), where the blood signal in the aorta lumen is not suppressed, as well as “Black Blood” (BB) images, where the blood signal is suppressed (thus black). To obtain the WB image, an ECG-gated multi-slice gradient echo sequence (TR = 1 heartbeat, about 120 ms, TE = 2.4 ms) was employed with FOV = 26 \times 25 mm, matrix size = 192 \times 128, slice thickness = 0.8 mm. To obtain BB images, ECG-gated multi-slice FLASH (Fast imaging with Low Angle SHot) (TR = 76 ms, TE = 0.97 ms, FA = 40 degrees) was applied to the same slice. Due to the long



interval between MRI sessions, we relied on the unique anatomy of the aortic arch, its branching points, and other thoracic arteries in the imaging plane for the comparison of pre- and post-injection images. For images acquired after **PGd-LNP** injection, BB was further combined with a fat suppression option in the MRI protocol although such option is not necessary for the detection of enhancement.

Cryo-imaging analysis of the atherosclerotic mouse model after MRI

Immediately after euthanasia, the mouse was embedded in Tissue-Tek® Optimal Cutting Temperature (O. C. T.) solution (Sakura Finetek USA, Inc., Torrance, CA, USA) and frozen in liquid nitrogen before transfer to BioInVision Inc. (Cleveland, OH, USA). Concomitant cryosectioning and imaging was performed using an automated microtome-blockface episcopic imaging system (CryoViz™, BioInVision Inc.) that allows for microscopic, three-dimensional, high resolution brightfield and fluorescent imaging of macroscopic specimens. The mouse was sectioned from dorsal to ventral with a section thickness of 40 μm and an in-plane resolution of 10 μm × 10 μm, which were applied to the entire body.

Conclusions

In this study, we developed a new method for the efficient covalent surface functionalization of LNPs using the amide-forming KAT ligation to synthesize multi-functional hybrid LNPs. Under neutral and diluted aqueous conditions, surface functionalization proceeded in a high yield with several types of bioactive molecules including a peptide (**P**), a Gd(III)-chelate (**Gd**), and a fluorescent dye (sulforhodamine B, **R**). The resulting **PGd-LNP** and **PGdR-LNP** particles showed no significant aggregation with a hydrodynamic diameter of *ca.* 50 nm. The KAT ligation with fast reaction rate enabled the **LNP-KAT** surface functionalization, by using a minimum amount (1.2 equiv.) and low concentration (0.12 mM) of HA derivatives, to provide **PGd-LNP** or **PGdR-LNP** with outstanding loadings of Gd(III)-chelate (*ca.* 2300 per NP). The obtained LNPs revealed a high relaxivity ($r_1 = 22.0 \text{ s}^{-1} \text{ mM}^{-1}$ at 1.4 T/60 MHz at 25 °C), which was correlated with the strong enhancement in *in vivo* images in the atherosclerotic plaques of BA in *apoE*^{-/-}. Further *ex vivo* study by cryo-fluorescent imaging of a whole mouse injected with triply functionalized **PGdR-LNP** clearly suggested the strong accumulation of LNPs in atheroplaque in the BA region, completely overlapped with the observed MRI enhancement of the identical mice. This study demonstrated a powerful and versatile method to produce multifunctional LNPs with a sufficiently high payload of functional moieties on the surface on demand, such as peptides and imaging probes. By taking advantage of this efficient attachment of required functional groups by KAT ligation, further production of multi-functional LNPs that employ this **LNP-KAT** scaffold particle is currently in progress.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This study was performed in strict accordance with the NIH guidelines for the care and use of laboratory animals (NIH Publication No. 85-23 Rev. 1985) and was approved by the Institutional Animal Care and Use Committee of the University of Pennsylvania (Philadelphia, PA, USA). Authors thank Ms. Agnès Pallier of CNRS Orléans for performing relaxivity measurements. Authors thank Prof. Helm of EPFL for educational discussion. Authors thank Prof. Walde of ETH for intensive and fruitful discussion. Authors thank Prof. Leroux of ETH for his help in nanoparticle preparation and NTA measurements. Authors thank Prof. Spencer of ETH for his help in AFM imaging and Prof. Morbidelli of ETH for his help in DLS measurements. Authors thank Dr Pickup of the Small Animal Imaging Facility (SAIF), Department of Radiology, University of Pennsylvania for his expert assistance. Authors thank Mr Wirz and Dr Rubi of MoBIAS of ETH for their help in MALDI-MS measurements. The ScopeM facility in ETH is acknowledged for TEM measurements. Authors thank Prof. Lowell of Virginia Tech., and Dr Pattabiraman and Prof. Bode of ETH for fruitful discussion. This research was supported in part by an ETH Research Grant (ETH-25 11-1, ETH-45 19-1, YY), the Swiss National Foundation (200021_156097, 205321_173018, IZLJZ2_183660, YY), the Researcher's Developing Grant from the American Heart Association (0930140N, YY), and JSPS KAKENHI (Grant-in-Aid for Scientific Research [A], 6251004, HM).

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