

## EDGE ARTICLE

[View Article Online](#)  
[View Journal](#) | [View Issue](#)
Cite this: *Chem. Sci.*, 2019, **10**, 5603

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

Received 8th February 2019  
 Accepted 18th April 2019

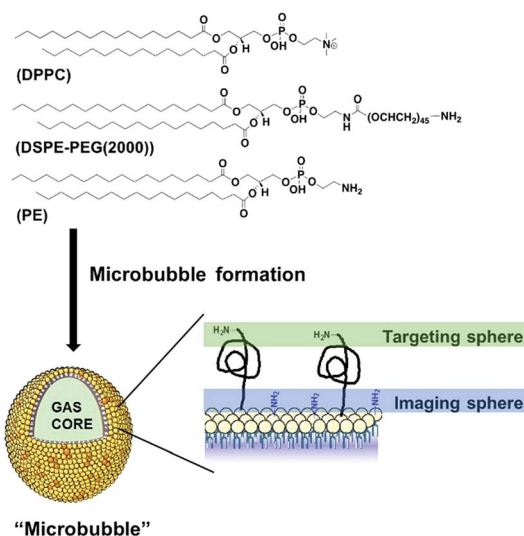
DOI: 10.1039/c9sc00684b  
[rsc.li/chemical-science](http://rsc.li/chemical-science)

## Introduction

The development and further refinement of contrast agents plays a central role in the fields of anatomical and molecular imaging, acting as a driving force to overcome limitations inherent to existing imaging modalities.<sup>1</sup> In ultrasound (US), the use of microbubbles (MBs, Scheme 1), which consist of a gas core stabilised by a phospholipid or polymer shell, has improved the poor ability of this technique to distinguish between tissues with similar acoustic responses.<sup>2</sup> In the presence of an acoustic field, these compressible MBs produce strong nonlinear signals, which can be differentiated from the surrounding tissue, enhancing soft-tissue contrast and signal-to-noise ratios in regions of interest. The fact that MBs typically present sizes of 1–5 µm, and therefore are restricted to intravascular targets, alongside their ability to incorporate disease-targeted ligands, has recently oriented the use of US imaging towards detecting and monitoring vascular pathologies at the molecular level.<sup>2c</sup> In this regard, different preclinical studies have successfully demonstrated the capability of MBs to visualise receptors overexpressed in inflammation, angiogenesis and/or early tumour formation,<sup>3a–c</sup> and a phospholipid-

based formulation has even recently entered first in-human clinical trials in various cancer types.<sup>3d,e</sup> These examples, together with the proven capacity of MBs to load drugs<sup>3f</sup> will further support the development of alternative MB agents for new diagnostic and therapeutic US applications in patients.

Clinical translation, however, is always a long, high-risk and expensive endeavour, and therefore, early evaluation of the



Scheme 1 Schematic structure/composition of the developed microbubbles (MBs), showing two surface conjugation sites (not to scale).

<sup>a</sup>Department of Chemistry, Imperial College London, UK. E-mail: [j.hernandez-gil@imperial.ac.uk](mailto:j.hernandez-gil@imperial.ac.uk); [n.long@imperial.ac.uk](mailto:n.long@imperial.ac.uk)

<sup>b</sup>Department of Surgery & Cancer, Imperial College London, UK. E-mail: [eric.aboagye@imperial.ac.uk](mailto:eric.aboagye@imperial.ac.uk)

<sup>c</sup>Department of Bioengineering, Imperial College London, UK

† Electronic supplementary information (ESI) available. See DOI: [10.1039/c9sc00684b](https://doi.org/10.1039/c9sc00684b)



biodistribution and/or pharmacokinetics of MBs could facilitate their (pre)clinical development.<sup>4a</sup> Whole-body imaging has long demonstrated utility in the evaluation of *in vivo* fates and/or possible off-target effects.<sup>4</sup> Unfortunately, contrast-enhanced US is limited to local imaging. This limitation has created a demand for new MB formulations that can provide whole-body imaging and therefore, offer a better understanding of diagnostic and/or drug activity during the initial stages of MB development.

Nuclear imaging techniques, namely positron emission tomography (PET) and single-photon-emission tomography (SPECT), provide non-invasive and whole-body imaging with high sensitivity and unlimited penetration depth.<sup>5</sup> Hence, a few groups have attempted to develop dual nuclear/US MB agents.<sup>6</sup> Most of these studies, however, have used streptavidin-biotin interactions to radiolabel MBs and/or employed SPECT tracers (iodine-125, technetium-99m or indium-111).<sup>6a-d</sup> Although elegant and encouraging, these approaches face disadvantages: (i) immunogenic streptavidin-coated MBs are not recommended for human use and (ii) SPECT cannot provide quantitative analysis *in vivo*. PET, unlike SPECT, allows determination of tracer concentration in tissue with higher sensitivity and image quality. Despite these advantages, only one group has incorporated a <sup>18</sup>F-labelled phospholipid onto MBs, allowing non-invasive assessment of the whole-body MB fates.<sup>6e</sup> However, <sup>18</sup>F-labelling usually requires time-consuming multi-step synthesis, and more importantly, depends on expensive cyclotron facilities. So, there is an unmet need for a rapid and robust method to generate radiolabelled MBs using broadly accessible PET isotopes.

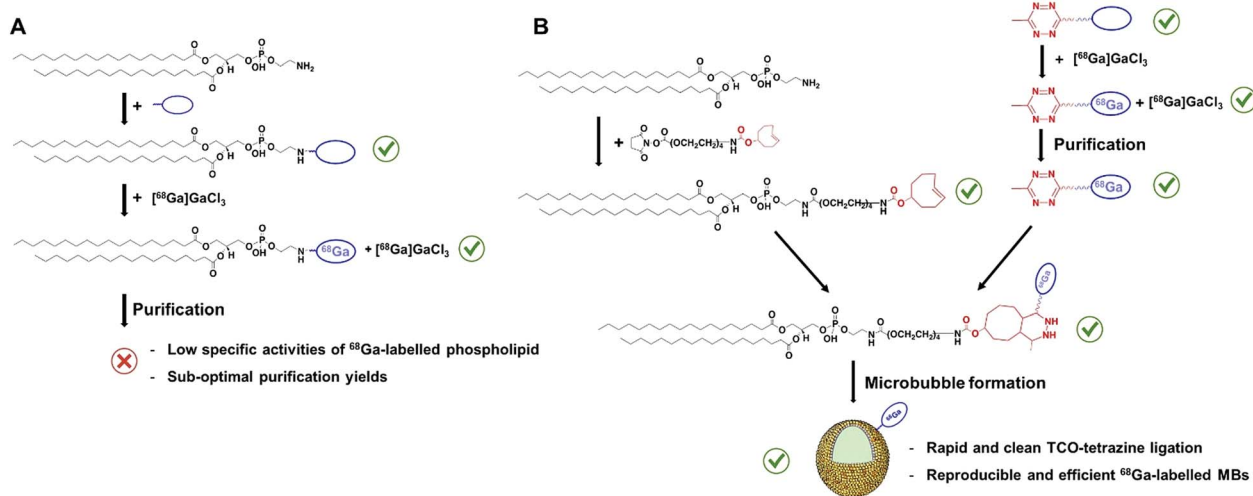
Gallium-68 (hereafter, <sup>68</sup>Ga) is a radiometal that offers aqueous single-step and high-yielding complexations with high positron emission (89%) and on-site availability, allowing hospitals economical production of a high-quality and generator-produced PET isotope.<sup>7</sup> In addition, its short half-life

(67.7 min) enables the perfect monitoring of MBs, which have circulation times of around 10–20 minutes.<sup>2</sup> However, the development of <sup>68</sup>Ga-labelled MBs (hereafter, <sup>68</sup>Ga-MBs) has never been attempted, which inspired us to focus on this particular problem in the field of US imaging. We reasoned that the optimised method should offer a fast, accessible and versatile way to radiolabel the shell components of MBs, as well as to provide reliable validation checkpoints with conventional To test this hypothesis, two <sup>68</sup>Ga labelling strategies were initially devised: (i) direct conjugation of a metal chelator to the phospholipid head-groups and, (ii) bioorthogonal 'click' ligation between a chelator-based prosthetic group and a complementary functionalised-phospholipid (Scheme 2A and B). Through a direct comparison between both approaches, we have developed a rapid and efficient method for labelling ultrasound MB agents with a generator-produced PET isotope, offering accessible solutions for *in vivo* tracking of MBs.

## Results and discussion

### Design and preparation of microbubbles

Here, we focused on phospholipid-shelled MBs, which represent the most common and versatile US contrast agent.<sup>2,3</sup> We recently demonstrated that a cyanine7.5-modified phospholipid could be successfully incorporated into a MB formulation of commercially available 1,2-dipalmitoyl-*sn*-glycero-3-phosphocholine (DPPC) and 1,2-distearoyl-*sn*-glycero-3-phosphoethanolamine-*N*-(amino (poly-ethylene glycol)-2000) [DSPE-PEG(2000)] phospholipids.<sup>8a</sup> We reasoned that 1,2-distearoyl-*sn*-glycero-3-phosphoethanolamine (PE), which shares DSPE-PEG(2000) backbone without its PEGylated moiety, could therefore, be combined with DPPC and DSPE-PEG(2000), and thus offer an alternative anchor point for the PET tracer. The absence of a bulky PEG<sub>45</sub> polymer on PE also facilitates the purification and characterisation of resulting products, while



**Scheme 2** (A) "Direct labelling approach": conjugation of a metal chelator to the headgroup of PE and subsequent labelling and purification. (B) "Click labelling approach": development of bioorthogonal components for 'click' ligation of a <sup>68</sup>Ga-chelator-tetrazine and a *trans*-cyclooctene-modified phospholipid to yield <sup>68</sup>Ga-labelled MBs.



maintaining DSPE-PEG(2000) available for subsequent conjugation of targeting ligands (Scheme 1). With such a rationale in mind, we prepared a new formulation containing DPPC, DSPE-PEG(2000) and PE at 85 : 5 : 10 molar ratios, following standard protocols (Table S1†).<sup>8</sup> Optical microscopy was used to visualise the size and concentration of the new MB formulation and, as shown in Table S2,† no significant difference was observed between PE-containing MBs and our previous formulation without PE. The stock MB solution containing 10% of PE yielded a mean bubble diameter of  $2.10 \pm 1.07 \mu\text{m}$  and a concentration of  $(1.22 \pm 0.68) \times 10^{10} \text{ MBs mL}^{-1}$  while the unmodified formulation generated bubbles with a mean size of  $2.18 \pm 1.06 \mu\text{m}$  and a concentration of  $(8.02 \pm 0.03) \times 10^9 \text{ MB mL}^{-1}$ . The acoustic response of both MBs was also compared. The two formulations were diluted to a concentration of approximately  $10^6 \text{ MBs mL}^{-1}$  in a 2 L water tank and their US contrast enhancements quantified (Table S2†). Both MBs produced a similar US contrast at 4.5 MHz, indicating that the new formulation can also be used for US imaging.

Both MBs produced a similar US contrast at 4.5 MHz, indicating that the new formulation can also be used for US imaging.

### Evaluation of the first approach: direct conjugation of a metal chelator to the phospholipid head-groups

To test the feasibility of our first approach (Scheme 2A), we functionalised PE with three chelators of known affinity for

$^{68}\text{Ga}$ :<sup>7a</sup> 1,4,7-triazacyclononane-1,4,7-triacetic acid (NOTA), 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA) and 2-(4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododecan-1-yl)- pentanedioic acid (DOTAGA). PE-NOTA (1) and PE-DOTA (2) were prepared by conjugation of the primary amine head-group of PE with a NHS-activated form of each chelator, whereas PE-DOTAGA (3) was obtained *via* reaction of a *p*-isothiocyanatobenzyl PE intermediate with an ethylenediamine-modified DOTAGA reagent. **1–3** were purified by preparative thin-layer chromatography (TLC) and obtained in high yields (Fig. 1A and ESI†).

To examine whether **1–3** can efficiently chelate  $^{68}\text{Ga}$ , a fixed amount of generator-produced  $^{68}\text{Ga}^{3+}$  ( $\sim 2 \text{ MBq}$ ) was separately reacted with **1–3** over a range of phospholipid concentrations and two pH values (3.5 and 4.5). All reactions were incubated for 10 min at 90 °C and the radiochemical yields (RCYs) were measured using radioTLC. **1** demonstrated the highest labelling efficiency, with RCYs greater than 80% at moderately low chelator concentrations (10  $\mu\text{M}$ ) for both pH conditions, corresponding to molar activities of 1–2 MBq nmol $^{-1}$  (Fig. 1B).

As previously reported, NOTA derivatives show better affinity for  $^{68}\text{Ga}^{3+}$  than DOTA conjugates as a consequence of the close fit of the relatively small Ga(II) metal ion in the pseudo octahedral cavity of the potentially six-coordinate 9-aneN<sub>3</sub>, with attached carbonyl and amide groups.<sup>9</sup> We then selected **1** for optimising a  $^{68}\text{Ga}$ -labelling method under *in vivo* conditions. Purified  $^{68}\text{GaCl}_3$  (70–75 MBq) was reacted with **1** ( $\sim 20 \text{ nmol}$ ) in acetate buffer (0.1 M, pH 4.0) for 15 min at 90 °C. The RCY of

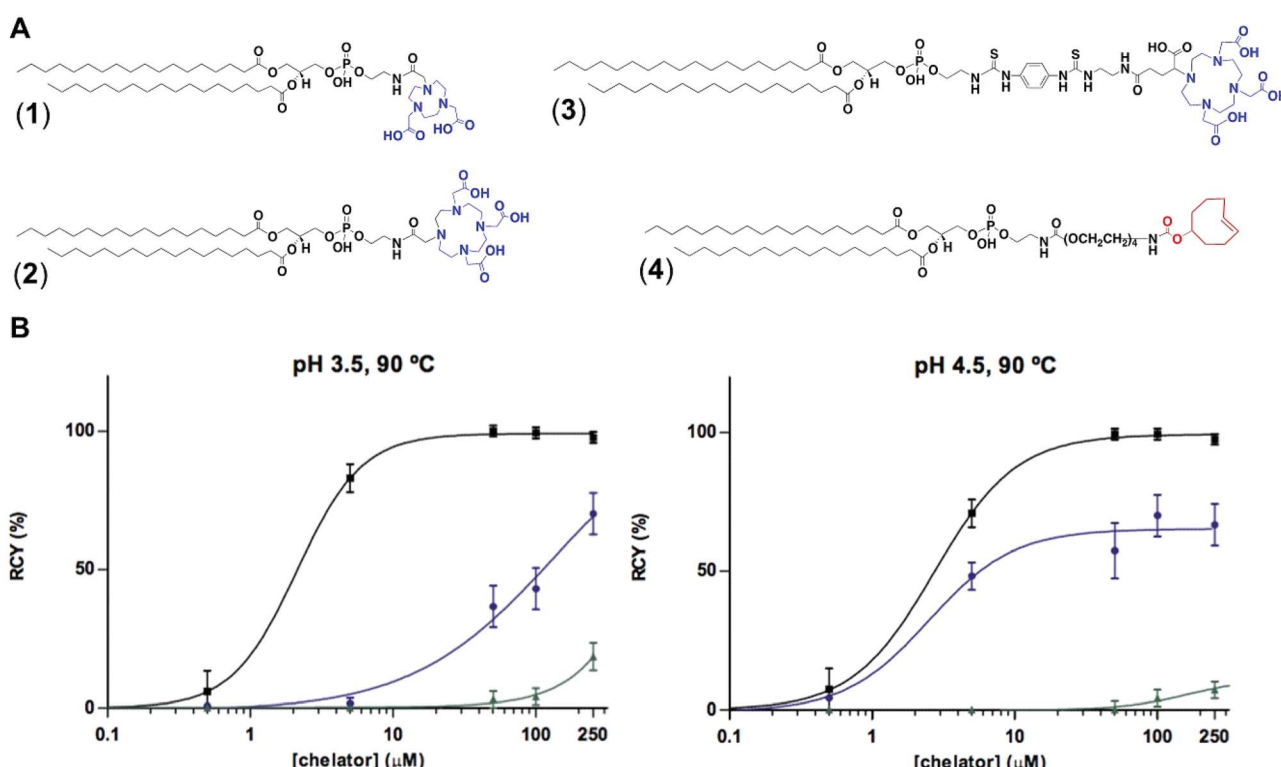


Fig. 1 (A) Structure of PE-NOTA (1), PE-DOTA (2), PE-DOTAGA (3) and PE-PEG<sub>4</sub>-TCO (4). (B) Radiochemical yields for the reaction of  $^{68}\text{Ga}^{3+}$  with **1** (black), **2** (green) and **3** (blue) under different phospholipid concentrations (0.05 to 250  $\mu\text{M}$ ) and pH values (3.5 and 4.5), after 10 min at 90 °C.



[<sup>68</sup>Ga][Ga(1)], under these conditions, was  $>85\%$  (non-isolated, estimated by radio-HPLC). Next, different solid-phase extraction (SPE) cartridges were used for [<sup>68</sup>Ga][Ga(1)] purification but, unfortunately all sorbents tested presented low recovery yields (Table S3†). The best results were obtained with Oasis® HLB cartridges with an average eluted activity of  $27 \pm 5\%$ . These all-purpose cartridges, with a strongly hydrophilic and reversed-phase polymer, allow acidic, basic and neutral analyte purifications from pH 0 to 14. Under our labelling conditions, pH = 3.5–4.0, [<sup>68</sup>Ga][Ga(1)] behaves as a zwitterionic phospholipid with a positively charged Ga-chelating moiety and a negatively charged phosphate group ( $pK_a \sim 1.5\text{--}2.5$ ), meeting *a priori* the manufacturer requirements for high recovery yields. Phospholipids, however, are a major concern for LC-MS analysis of small molecules in biological matrices and many SPE cartridges (in our case, HybridSPE®, Oasis® Prime and Oasis® HLB cartridges) contain sorbents that can retain phospholipids. This effect was clearly observed when using Oasis® Prime cartridges, retaining  $62 \pm 15\%$  of the initial loaded activity and, also noticeable with HybridSPE® and Oasis® HLB cartridges trapping  $32 \pm 7$  and  $17 \pm 3\%$ , respectively. Therefore, the sub-optimal purification yields obtained because of the lipid nature of 1 alongside its low molar activities prompted us to focus on the second approach.

### Evaluation of the second approach: bioorthogonal 'click' ligation between a chelator-based prosthetic group and a complementary functionalised-phospholipid

For the second strategy (Scheme 2B), we hypothesised that a tetrazine-bearing radioconjugate could combine high molar activities and straightforward work-up procedures from small-sized chelators with high-yielding and fast reactivities from 'click' ligation. In this regard, our group and others have been designing new probes suited for bioorthogonal radiolabelling *via* the inverse-electron-demand Diels–Alder (iEDDA) cycloadditions between tetrazines and olefins.<sup>10</sup> iEDDA reactions avoid the use of metal catalysts, which could compete with the far less abundant  $^{68}\text{Ga}^{3+}$  radiometal, as well as offer rapid kinetics at low concentrations, essential when working with short-lived radioisotopes.<sup>11</sup> Thus, a *trans*-cyclooctene (TCO)-modified phospholipid was synthesised using a commercially available TCO-PEG<sub>4</sub>-NHS ester and PE. TCO was chosen because of its higher reactivity compared to other strained olefins.<sup>12</sup> PE-PEG<sub>4</sub>-TCO (4) was purified by preparative-TLC and characterised by <sup>1</sup>H NMR spectroscopy and MALDI-TOF mass spectrometry (Fig. 1a and ESI†).

Compound 5, containing *N,N'*-bis[2-hydroxy-5-(carboxyethyl)-benzyl]ethylenediamine-*N,N*-diacetic acid (HBED-CC) and a tetrazine, was also prepared (Fig. 2 and ESI†). In a six-step sequence, the desired product was ultimately obtained by *in situ* coupling HATU-activated  $[\text{Fe}(\text{HBED-CC})]^-$  with (4-(6-methyl-1,2,4,5-tetrazin-3-yl)phenyl)methanamine. Here, we used HATU, which required a 15 minute activation, instead of the most common 2 day tetrafluorophenol/dicyclohexylcarbodiimide process.<sup>13</sup> The formation of the iron complex assures the protection of coordinating phenolic and carboxylate groups while exposing the two

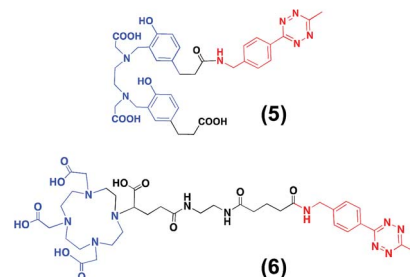


Fig. 2 (A) Structure of HBED-CC-tetrazine (5) and DOTA-GA-tetrazine (6).

propionic acid arms for amine conjugation. HBED-CC-tetrazine derivative 5 was purified by flash chromatography and its monosubstituted nature confirmed using <sup>1</sup>H, <sup>13</sup>C NMR spectroscopy and ESI mass spectrometry (see ESI†). As a reference, a tetrazine-bearing DOTA-GA tracer (6), previously designed by our group,<sup>10b</sup> was prepared (Fig. 2 and ESI†).

A head-to-head comparison of the <sup>68</sup>Ga-chelating efficiency between 5 and 6 was performed (Fig. 3). At room temperature, the difference between both chelators was remarkable. Whilst 5 performed extremely well, 6 presented poor ability to coordinate  $^{68}\text{Ga}^{3+}$ . This suggested that the labelling efficiency of 6 was limited by the higher kinetic barriers that macrocyclic-based chelators generally present compared to acyclic ligands.<sup>7a,14</sup> 5 also showed little pH-dependence, yielding near-quantitative RCYs at the low concentration of 5  $\mu\text{M}$  for all pH values tested (based on radioTLC analysis of the reaction solution). At high temperature, both chelators exhibited a similar efficiency, with conversions higher than 90% at 5  $\mu\text{M}$ . At lower concentrations, however, 5 again demonstrated better performance over 6. At 1  $\mu\text{M}$  and pH values of 3.5 and 4.5, RCYs of [<sup>68</sup>Ga][Ga(5)] were near quantitative while RCYs of [<sup>68</sup>Ga][Ga(6)] were lower than 30%. This corresponded to maximum molar activities for 5 of 15–20 MBq nmol<sup>-1</sup>. Next, we selected 5 to evaluate its <sup>68</sup>Ga-labelling efficiency under *in vivo* conditions. This time, [<sup>68</sup>Ga][Ga(5)], obtained after 10 min incubation at room temperature, could be successfully purified with conventional reverse-phase cartridges. The entire procedure was performed in less than 20 min, yielding purities  $>98\%$  and RCYs  $>95\%$  (determined by radio-HPLC analysis of the pure product).

### Preparation of <sup>68</sup>Ga-labelled MBs

Fig. 4A shows the chromatogram of purified [<sup>68</sup>Ga][Ga(5)] labelled at room temperature and pH  $4.2 \pm 0.3$ . As for other HBED-based chelators, 5 forms different geometric isomers when complexed to  $\text{Ga}^{3+}$ .<sup>15</sup> Although a mixture of species can potentially present different pharmaceutical profiles, a recent study measuring the cell binding and internalisation properties of [<sup>68</sup>Ga]Ga-PSMA-HBED-CC labelled at room temperature (containing a mixture of species) and at 95 °C (containing the thermodynamically most stable isomer) on PSMA-expressing LNCaP cells showed comparable results for both fractions, indicating a stereochemical independent receptor-interaction of the radioligand.<sup>16</sup>



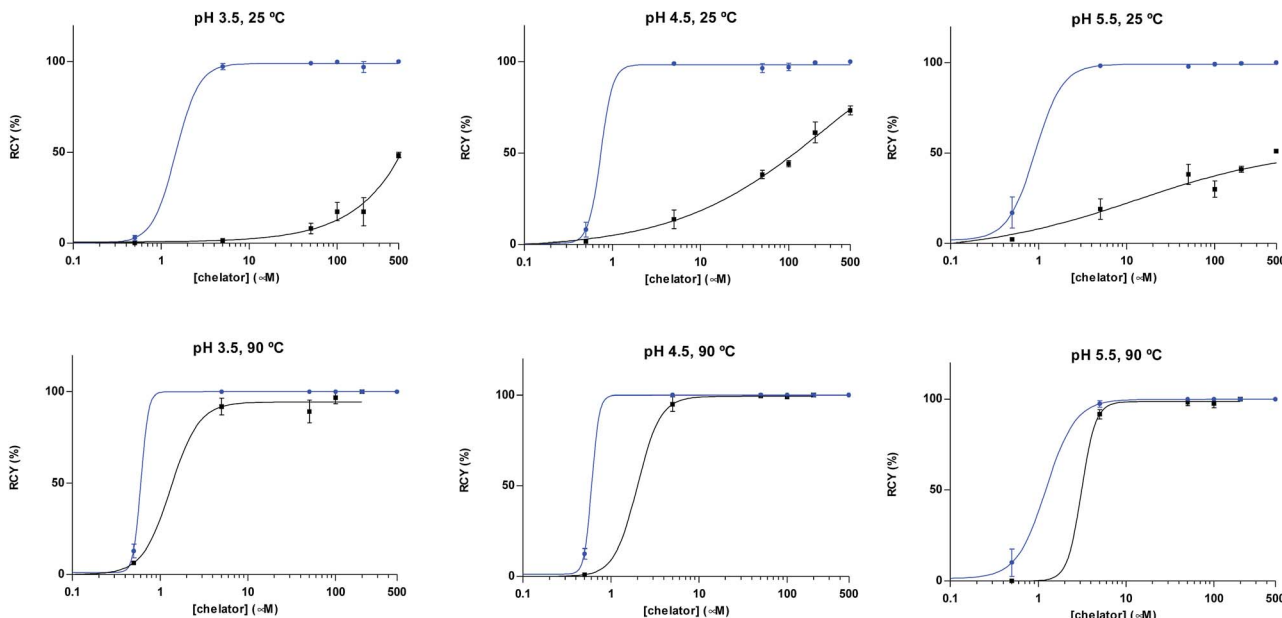


Fig. 3 Radiochemical yields for the reaction of  $^{68}\text{Ga}^{3+}$  with 5 (blue) and 6 (black) under different concentrations of chelator (0.05 to 500  $\mu\text{M}$ ), pH values (3.5, 4.5 and 5.5) and temperatures (25 and 90  $^{\circ}\text{C}$ ), after 10 min.

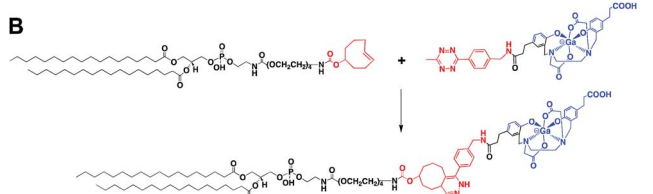
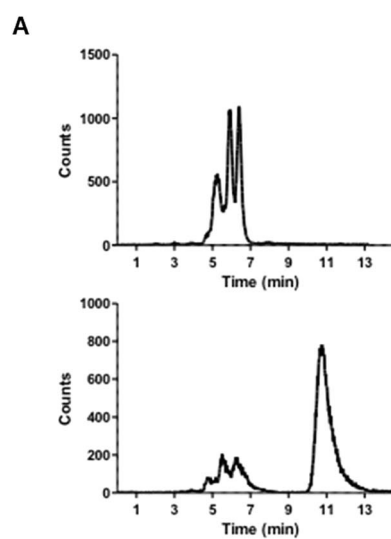


Fig. 4 (A) Radio-HPLC chromatograms of (top)  $^{68}\text{Ga}$ -Ga-HBED-CC-tetrazine precursor and (bottom) reaction between 4 and 5, after 15 min at 60  $^{\circ}\text{C}$ . (B) Scheme of the reaction between TCO-modified PE (4) and  $^{68}\text{Ga}$ -Ga-HBED-CC-tetrazine (5).

Next, we reacted a purified fraction of  $^{68}\text{Ga}$ [Ga(5)] ( $\sim 75$  MBq) with PE-PEG<sub>4</sub>-TCO (0.107  $\mu\text{mol}$ ) at 60  $^{\circ}\text{C}$  (Fig. 4A). After 15 minutes, the radio-HPLC chromatogram showed the

appearance of a new major peak at a slower retention time ( $\tau_r = 10.8$  min) than the labelled precursor ( $\tau_r = 5.3$ –6.4 min), which corresponded to radiochemical conversions of the dihydropyridazine products of  $\sim 80$ –85% (non-isolated, estimated by radio-HPLC) (Fig. 4A and B). Next,  $^{68}\text{Ga}$ -labelled PE (hereafter,  $^{68}\text{Ga}$ -PE) was combined with DPPC and DSPE-PEG(2000) and the resulting formulation was activated to form gas-filled MBs. Excess of unreacted  $^{68}\text{Ga}$ [Ga(5)] and free lipids were removed *via* a centrifugation/washing methodology.<sup>17</sup> The RCY of purified  $^{68}\text{Ga}$ -MBs was around 40–50%, generating samples with  $\sim 30$ –35 MBq and concentrations of  $(1.28 \pm 0.68) \cdot 10^9$  MB mL<sup>-1</sup>, in 40–50 minutes after  $^{68}\text{Ga}$  elution from the generator.

### In vivo study of $^{68}\text{Ga}$ -labelled MBs

Encouraged by the successful  $^{68}\text{Ga}$ -labelling of MBs, comparative dynamic PET imaging and biodistribution studies of  $^{68}\text{Ga}$ [Ga(5)],  $^{68}\text{Ga}$ -PE and  $^{68}\text{Ga}$ -MBs were undertaken on Balb/c nude mice ( $n = 5$ –6). The optimised labelling approach yielded MB stock solutions with high activity and concentration, allowing us to use concentrations of approximately  $5 \times 10^7$  MB per injection with activities between 0.37–0.74 MBq. These MB concentration values are in the same order of typical diagnostic doses used in US imaging, suggesting that this strategy generates  $^{68}\text{Ga}$ -MBs that can also be exploited for complementary US studies.

Kinetic analysis showed that the MBs had a pharmacokinetic profile different from that of freely-injected PE and 5 (Fig. 5 and S6–S8†). Following the injection of free  $^{68}\text{Ga}$ [Ga(5)], activity accumulated in the bladder, gallbladder and intestines within the first 5 minutes, which confirmed that  $^{68}\text{Ga}$ [Ga(5)] was characterised by a rapid tissue distribution and subsequent clearance through the urinary route.  $^{68}\text{Ga}$ -PE and  $^{68}\text{Ga}$ -MBs, on



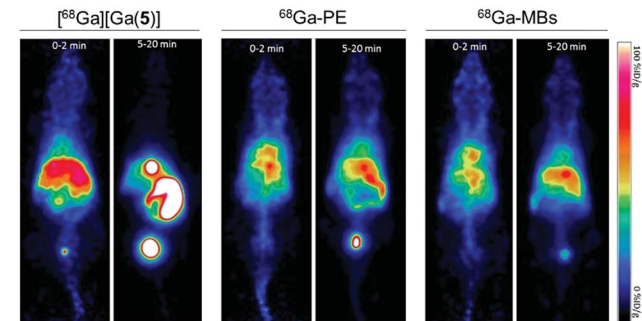


Fig. 5 Small-animal PET imaging of Balb/c nude mice after intravenous injection of  $[^{68}\text{Ga}][\text{Ga}(5)]$  (left),  $^{68}\text{Ga}$ -labelled PE (middle) and  $^{68}\text{Ga}$ -labelled MBs (right) at early (0–2 min) and late (5–20 min) time intervals.

the contrary, presented longer circulation times in the animals, with high blood pool mean values persisting for the entire imaging period. Both formulations, however, presented differences in their biodistribution, which became more evident at later time points. Between 0 to 2 min post-injection (PI),  $^{68}\text{Ga}$ -PE and  $^{68}\text{Ga}$ -MBs were mainly present in the heart and liver, whereas at 5–20 min time interval,  $^{68}\text{Ga}$ -PE was mainly accumulated in the bladder and gallbladder, and  $^{68}\text{Ga}$ -MBs were present in the liver, gallbladder and spleen. The uptake in the liver and spleen reached equilibrium around 3 min PI at approximately 66 and 40% ID  $\text{g}^{-1}$ , respectively. Previous studies have also indicated the liver and the spleen as the major organs for the accumulation of MBs with sizes around 1–10  $\mu\text{m}$ .<sup>6a,b</sup> In addition,  $^{68}\text{Ga}$ -MBs showed an initial peak of activity in the lungs of  $43 \pm 12\%$  ID  $\text{g}^{-1}$ , which decreased to  $20 \pm 5\%$  ID  $\text{g}^{-1}$  at 20 min, indicating that  $^{68}\text{Ga}$ -MBs were gradually redistributed into the systemic circulation after an initial accumulation in the lungs. Biodistribution data confirmed the PET results. At 20 min PI, both  $[^{68}\text{Ga}][\text{Ga}(5)]$  and  $^{68}\text{Ga}$ -PE were mostly present in the urine ( $689 \pm 64$  and  $146 \pm 26\%$  ID  $\text{g}^{-1}$ , respectively), consistent with an efficient urinary excretion route.  $^{68}\text{Ga}$ -MBs, on the other hand, showed a more distributed uptake within the animal models, with accumulation of radioactivity mainly observed in the liver ( $47 \pm 4\%$  ID  $\text{g}^{-1}$ ), spleen ( $40 \pm 2\%$  ID  $\text{g}^{-1}$ ) and urine ( $57 \pm 11\%$  ID  $\text{g}^{-1}$ ).

## Conclusions

In conclusion, we have developed an efficient method for labelling MBs with a generator-produced PET isotope using a rapid and clean ligation between a TCO-modified phospholipid and a  $^{68}\text{Ga}$ -HBED-CC-tetrazine tracer. Bioconjugation of TCO to phospholipids is simple and reproducible, and the new HBED-CC-tetrazine chelator provides efficient and high-yielding  $^{68}\text{Ga}$ -labelling, affording reproducible synthesis of  $^{68}\text{Ga}$ -MBs under mild conditions. This method offers real-time monitoring and the possibility of easily customising alternative phospholipid-based formulations. We also confirmed that the resulting  $^{68}\text{Ga}$ -MBs allow non-invasive study of the whole-body distribution of MBs in mice. As contrast-enhanced US

has emerged as a promising imaging modality for clinical applications, we believe that this strategy is a favourable way of non-invasively evaluating the pharmacokinetics and off-target accumulation of new MB formulations in their initial stages of (pre)clinical development. Therefore, this strategy can be easily implemented across multiple centres and hospitals, accelerating MB evaluation and development. Future work will explore the incorporation of targeted-modified phospholipids into the MB formulation and compare  $^{68}\text{Ga}$ -labelled MB containing targeted and non-targeted vectors.

## Experimental section

### Materials

All the syntheses were carried out with the following commercially available reagents used without further purification: 4-(aminomethyl)benzonitrile hydrochloride (97%, Sigma Aldrich), di-*tert*-butyl dicarbonate ( $\geq 99\%$ , Sigma Aldrich), nickel(II) trifluoromethanesulfonate (96%, Sigma Aldrich), hydrazine hydrate (50–60%, Sigma Aldrich), sodium nitrite ( $\text{NaNO}_2$ ,  $\geq 97\%$ , Sigma Aldrich), 3-(4-hydroxyphenyl)propionic acid (98%, Sigma Aldrich), sodium hydroxide (98.5–100.5%, VWR International), ethylenediamine-*N,N'*-diacetic acid ( $\geq 98\%$ , Sigma Aldrich), formaldehyde (37 wt% in  $\text{H}_2\text{O}$ , Sigma Aldrich), hydrochloric acid (37%, VWR International), trifluoroacetic acid ( $\geq 99.5\%$ , Tokyo Chemical Industry Co., Ltd.), *N,N*-diisopropylethylamine ( $\geq 99\%$ , Sigma Aldrich), iron(III) chloride (97%, Sigma Aldrich), 1-[Bis(dimethylamino)methylene]-1*H*-1,2,3-triazolo[4,5-*b*]pyridinium 3-oxid hexafluorophosphate, *N*-(dimethylamino)-1*H*-1,2,3-triazolo-[4,5-*b*]pyridin-1-ylmethylene-*N*-methylmethanaminium hexafluorophosphate N-oxide (HATU, 97%, Sigma Aldrich), glutaric anhydride (95%, Sigma Aldrich). Triethylamine ( $\geq 99\%$ , Sigma Aldrich), *N*-hydroxysuccinimide (98%, Sigma Aldrich), *N,N'*-Dicyclohexylcarbodiimide (DCC, 99%, Sigma Aldrich), *trans*-cyclooctene-PEG<sub>4</sub>-NHS ester (TCO-PEG<sub>4</sub>-NHS,  $>95\%$ , Jena Bioscience), *trans*-cyclooctene-NHS ester (E)-cyclooct-4-enyl-2,5-dioxo-1-pyrrolidinyl carbonate (TCO-NHS ester,  $>95\%$ , Jena Bioscience), 4-nitroaniline ( $\geq 99\%$ , Sigma Aldrich), *p*-phenylene diisothiocyanate (98%, Sigma Aldrich), 2,2',2''-(10-(4-((2-aminoethyl)amino)-1-carboxy-4-oxobutyl)-1,4,7,10-tetraazacyclododecane-1,4,7-triyl)triacetic acid (NH<sub>2</sub>-DOTA-GA, ChemMtech mdt), 2,2',2''-(10-(2-((2,5-dioxopyrrolidin-1-yl)oxy)-2-oxoethyl)-1,4,7,10-tetraazacyclododecane-1,4,7-triyl)triacetic acid (DOTA-NHS ester, ChemMtech mdt), 2,2'-7-(2-((2,5-dioxopyrrolidin-1-yl)oxy)-2-oxoethyl)-1,4,7-triazonane-1,4-diyl diacetic acid (NOTA-NHS ester, ChemMtech mdt), sodium acetate ( $\geq 99\%$ , Sigma Aldrich), acetic acid ( $\geq 99\%$ , Sigma Aldrich), ammonium acetate ( $\geq 98\%$ , Sigma Aldrich), decafluorobutane (99%, FluorMed, L.P.), 1,2-dipalmitoyl-*sn*-glycero-3-phosphocholine (DPPC,  $\geq 99\%$ , Avanti Polar Lipids), 1-palmitoyl-2-hydroxy-*sn*-glycero-3-phosphocholine (LPC,  $\geq 99\%$ , Avanti Polar Lipids), 1,2-distearoyl-*sn*-glycero-3-phosphoethanolamine (18:0 PE-NH<sub>2</sub>,  $\geq 99\%$ , Avanti Polar Lipids), 1,2-distearoyl-*sn*-glycero-3-phosphoethanolamine-*N*-(amino(polyethylene glycol)-2000) (ammonium salt) (DSPE-PEG(2000)-NH<sub>2</sub>,  $\geq 99\%$ , Avanti Polar Lipids), primuline (50%,



Sigma Aldrich), propylene glycol ( $\geq 99.5\%$ , Sigma Aldrich), glycerol (99%, Sigma Aldrich), Dulbecco's phosphate-buffered saline (10 $\times$ , Sigma Aldrich). Solvents were HPLC grade and obtained from VWR Chemicals. Anhydrous chloroform and methanol were purified and dried according to standard methods prior to use and, anhydrous dimethyl sulfoxide was obtained from Sigma Aldrich ( $\geq 99.5\%$ ).

For experiments working with radioactive substances, water (Ultrapur, VWR) was used for preparation of buffers, HCl and NaCl solutions. 5.5 M HCl prepared by dilution of hydrochloric acid 37% (Emprove, VWR).

## Methods

1D  $^1\text{H}$  and  $^{13}\text{C}$  NMR spectra were recorded on either a Bruker AV-500 or on a Bruker AV-400 spectrometer at room temperature using standard pulse programs. Chemical shifts ( $\delta$ ) are quoted in parts per million (ppm) and referenced to the appropriate residual solvent signal. Coupling constants ( $J$ ) are reported to the nearest 0.5 Hz. Mass spectra ( $m/z$ ) of phospholipids were run on a MALDI micro MX – TOF mass spectrometer from Waters. Samples were spotted 1 : 1 v/v with a matrix solution (4-nitroaniline, 10 mg mL $^{-1}$  in methanol) and measured in reflector mode. High resolution mass spectra ( $m/z$ ) were recorded on either a VG platform II or VG AutoSpec spectrometers, with only molecular ions ( $\text{M}^+$ ,  $\text{MH}^+$ ,  $\text{MNa}^+$ ,  $\text{MK}^+$ ,  $\text{MNH}_4^+$ ) and major peaks being reported. Flash chromatography (FC) was performed on silica gel (Merck Kieselgel 60 F<sub>254</sub> 320–400 mesh). Thin Layer Chromatography (TLC) was performed using TLC Silica Gel 60 F24 (aluminium sheets 20  $\times$  20 cm for analytical runs and glass plates 20  $\times$  20 cm for preparative TLC purifications of phospholipids) from Merck. Plates were visualised either by quenching of ultraviolet fluorescence ( $\lambda = 254$  and 366 nm) or by charring with 10% KMnO<sub>4</sub> in 1 M H<sub>2</sub>SO<sub>4</sub>. Phospholipids were additionally visualised by charring with 5% primuline in acetone : water (8 : 2 v/v), lipids appearing as yellow spots under 365 nm wavelength. Flash purifications were carried out in a Isolera<sup>TM</sup> Spektra System using Biotage<sup>®</sup> SNAP Ultra C18 for reverse phase conditions (12, 30 and 60 g sizes) and, Biotage<sup>®</sup> SNAP Ultra C18 for normal phase conditions (25, 50 and 100 g sizes). Gradients are indicated throughout the text. Analytical HPLC chromatograms were run in a Waters AutoPurification<sup>TM</sup> System equipped with a 2489 UV/vis Dectector and a 3100 Mass Detector, using a Phenomenex Gemini C18 column (150 mm  $\times$  4.6 mm) at a flow rate of 1.2 mL min $^{-1}$  and gradient A: (5–98% B over 12 min then 98% B for 3 min) buffer A = H<sub>2</sub>O (0.1% TFA), buffer B = MeCN (0.1% TFA). Supelco Iso-Disc<sup>TM</sup> Filters PFTE-4-4 (4 mm  $\times$  0.45  $\mu\text{m}$ ) from Sigma Aldrich were used for sample filtration prior injection.

The lipid-coated decafluorobutane-filled microbubbles (MBs) were produced using a modified formulation described previously.<sup>18</sup> 1 mL of the resulting lipid solution (refer to Section 4 in this document for further details) was sealed in a 2 mL glass vial and the headspace was then purged with decafluorobutane at room temperature. The microbubbles were produced *via* mechanical agitation using a dental HL-AH High Speed Digital

Amalgamator Amalgam Capsule Blend Mixer HL-AH (4000 rpm for 30 s, two cycles). Particle size of non-radioactive MBs was measured on a bright-field microscope (Nikon Eclipse 50 $\times$ i, 40 $\times$  objective), 10  $\mu\text{L}$  diluted samples (1 : 100 dilution factor from stock sample) were first introduced into a haemocytometer and then sized and counted according to our reported protocol.<sup>19</sup>  $\zeta$ -potential analyses of microbubbles in solution (1 : 1000 dilution factor from stock sample) were measured with a NanoSizer (Malvern Nano-Zs, UK) and using DTS1070 cuvettes. Alternatively,  $^{68}\text{Ga}$ -labelled MBs were counted and sized using a Countess II FL Automated Cell Counter from Life Technologies. Non-radioactive controls were run to confirm size and concentration of MBs presented non-significant differences from both methods used to count and size MBs. MBs were purified by centrifugation adapting a reported method using a ROTINA 35 R centrifuge from Hettich Zentrifugen.<sup>17</sup>

$^{68}\text{Ga}$  was eluted from an Eckert & Ziegler Radiopharma GmbH GalliaPharm<sup>®</sup>  $^{68}\text{Ge}/^{68}\text{Ga}$  generator (Berlin, Germany) using a fully automated Modular-Lab system. Aqueous HCl solutions (0.1 M, 5 mL) was passed through the generator and the eluate was collected in a single fraction and used directly for  $^{68}\text{Ga}$ -labeling reactions (refer to radiochemistry section below for further details).

Different purification cartridges were used for radioactive experiments: silica-based strong cation exchange cartridge (Bond Elut-SCX, 100 mg, 1 mL) purchased from Agilent (further referred as SCX cartridge); Sep-Pack<sup>®</sup> Light tC18 cartridge (145 mg sorbent per cartridge), Sep-Pack<sup>®</sup> Plus Light C8 cartridge (145 mg sorbent per cartridge), Oasis<sup>®</sup> HLB cartridge (1cc/30 mg) and Oasis<sup>®</sup> Prime HLB cartridge Plus Light (100 mg) acquired from Waters. HybridSPE<sup>®</sup> – Phospholipid (30 mg/1 mL SPE Tubes) acquired from SUPELCO. iTLC experiments were measured on a Scan-RAM<sup>TM</sup> PET/SPECT radio-TLC Scanner from LabLogic using iTLC-SG – Glass microfibre chromatography paper impregnated with silica gel from Agilent Technologies. Analysis data was performed with Laura 3 software (LabLogic, Sheffield, UK). Radio-HPLC was carried out on an Agilent 1100 series HPLC system (Agilent Technologies, Stockport, UK) equipped with a  $\gamma$ -RAM Model 3 gamma-detector (IN/US Systems Inc., Florida, USA) and Laura 3 software (LabLogic, Sheffield, UK) was used for processing all analytical HPLC chromatograms. For tetrazine chelators, a Phenomenex Germini 5 u C18 (150  $\times$  4.6 mm) was used and an acidic gradient: 5–95% B over 15 min, then 95% B for 5 min, where A = water (0.1% TFA) and B = acetonitrile (0.1% TFA). For phospholipids a Phenomenex Bondclone 10 C18 (150  $\times$  3.90 mm) was used, as well as a neutral gradient: 5–95% B over 7 min, then 95% B for 8 min, where A = 100 mM ammonium formate (H<sub>2</sub>O) and B = methanol.

$^{68}\text{GaCl}_3$  was eluted from a GalliaPharm<sup>®</sup>  $^{68}\text{Ge}/^{68}\text{Ga}$  Generator from Eckert & Ziegler Radiopharma GmbH (0.74 GBq) using a fully automated Modular-Lab system and an adapted NaCl based method reported previously.<sup>20</sup> Briefly,  $^{68}\text{GaCl}_3$  was eluted in 5 mL of a 0.1 M HCl solution, which was subsequently trapped on a SCX cartridge (pre-conditioned with 1 mL 5.5 M HCl and 10 mL water) and the activity was then eluted, with minimal loss (2–5%), using a mixture of 12.5  $\mu\text{L}$  of 5.5 M HCl



and 500  $\mu$ L of 5 M NaCl solution. This eluate was subsequently buffered for direct use in  $^{68}\text{Ga}$ -labelling experiments.

### Organic syntheses

All the syntheses and experiments containing NHS ester compounds were performed under anhydrous conditions and an atmosphere of nitrogen in flame-dried glassware. All purified products were kept in the freezer and found to be stable for more than six months.

#### Synthesis of phospholipids

*Synthesis of PE-NOTA (1) and PE-DOTA (2).* 18:0 PE- $\text{NH}_2$  (7.5 mg, 10  $\mu\text{mol}$ ) was dissolved in 3 mL of a chloroform:methanol mixture (0.6 : 0.4 v/v) and triethylamine (50  $\mu\text{L}$ ). To this solution, either DOTA-NHS (10.0 mg, 13.1  $\mu\text{mol}$ ) or NOTA-NHS (10.0 mg, 15.2  $\mu\text{mol}$ ) was added and the mixture was left to stir for 5 h. After that, the crude was concentrated in rotavapour and purified by preparative-TLC.  $R_f$  for PE-DOTA = 0.10–0.19 (CHCl<sub>3</sub> : MeOH : H<sub>2</sub>O 6 : 4 : 0.2 v/v) and,  $R_f$  for PE-NOTA = 0.08–0.16 (CHCl<sub>3</sub> : MeOH : H<sub>2</sub>O 6 : 4 : 0.2 v/v).

*PE-DOTA.* 9.2 mg, 8.2  $\mu\text{mol}$ , 82%. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>):  $\delta$  (ppm) 0.87 (t, CH<sub>3</sub>, 6H,  $^3J_{\text{H-H}} = 6.5$  Hz), 1.25 (s, CH<sub>2</sub>, 56H), 1.57 (m, CH<sub>2</sub>CH<sub>2</sub>CO, 4H), 2.27 (m, CH<sub>2</sub>CH<sub>2</sub>CO, 4H), 2.41–3.18 (m, NCH<sub>2</sub>CH<sub>2</sub>N + NCH<sub>2</sub>CH<sub>2</sub>N + NCH<sub>2</sub>COOH, 22H), 3.64–4.43 (m, CH<sub>2</sub>CH<sub>2</sub>N + CH<sub>2</sub>CH<sub>2</sub>N + PO<sub>4</sub>CH<sub>2</sub>CH + COOCH<sub>2</sub>COH + COCH<sub>2</sub>N, 10H), 5.17 (s, PO<sub>4</sub>CH<sub>2</sub>CH, 1H). MALDI (matrix: 4-nitroaniline);  $m/z$  for [C<sub>59</sub>H<sub>118</sub>N<sub>6</sub>O<sub>16</sub>PNa]<sup>+</sup> ([M – H + NH<sub>4</sub> + Na + CH<sub>3</sub>CH<sub>2</sub>OH]<sup>+</sup>) expected = 1220.8, found 1220.7 and [C<sub>59</sub>H<sub>117</sub>N<sub>6</sub>O<sub>16</sub>PNa<sub>2</sub>]<sup>+</sup> ([M – 2H + NH<sub>4</sub> + 2Na + CH<sub>3</sub>CH<sub>2</sub>OH]<sup>+</sup>) expected = 1242.8, found 1242.6.

*PE-NOTA.* 8.3 mg, 8.0  $\mu\text{mol}$ , 80%. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>):  $\delta$  (ppm) 0.87 (t, CH<sub>3</sub>, 6H,  $^3J_{\text{H-H}} = 6.6$  Hz), 1.25 (s, CH<sub>2</sub>, 56H), 1.56–1.90 (m, CH<sub>2</sub>CH<sub>2</sub>CO + NCH<sub>2</sub>CH<sub>2</sub>N + NCH<sub>2</sub>CH<sub>2</sub>N, 16H), 2.26 (m, CH<sub>2</sub>CH<sub>2</sub>CO, 4H), 2.64–2.92 (m, NCH<sub>2</sub>COOH, 6H), 3.25–3.45 (m, CH<sub>2</sub>CH<sub>2</sub>N, 2H), 3.56–4.41 (m, CH<sub>2</sub>CH<sub>2</sub>N + PO<sub>4</sub>CH<sub>2</sub>CH + COOCH<sub>2</sub>COH + 6H), 5.19 (s, PO<sub>4</sub>CH<sub>2</sub>CH, 1H). MALDI (matrix: 4-nitroaniline);  $m/z$  for [C<sub>55</sub>H<sub>109</sub>N<sub>5</sub>O<sub>14</sub>PNa]<sup>–</sup> ([M – 3H + NH<sub>4</sub> + Na + CH<sub>3</sub>CH<sub>2</sub>OH]<sup>–</sup>) expected = 1117.8, found 1117.5 and [C<sub>55</sub>H<sub>110</sub>N<sub>5</sub>O<sub>14</sub>P]<sup>–</sup> ([M – 2H + NH<sub>4</sub> + CH<sub>3</sub>CH<sub>2</sub>OH]<sup>–</sup>) expected = 1095.7, found 1095.4.

#### Synthesis of PE-isothiocyanate

18:0 PE- $\text{NH}_2$  (50 mg, 66.9  $\mu\text{mol}$ ) was dissolved in 15 mL of a chloroform:methanol mixture (0.8 : 0.2 v/v) and triethylamine (10  $\mu\text{L}$ ). To this solution, a solution of *p*-phenylene diisothiocyanate (130 mg, 690.0  $\mu\text{mol}$ , dissolved in 3 mL of chloroform) was dropwise added and the mixture was left to stir for 5 h. After that, the crude was concentrated in rotavapour and purified by flash column chromatography.  $R_f$  = 0.5 (CHCl<sub>3</sub> : MeOH : H<sub>2</sub>O 8 : 2 : 0.2 v/v). (57 mg, 60.2  $\mu\text{mol}$ , 90%). <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>):  $\delta$  (ppm) 0.88 (t, CH<sub>3</sub>, 6H,  $^3J_{\text{H-H}} = 6.5$  Hz), 1.25 (s, CH<sub>2</sub>, 56H), 1.53 (m, CH<sub>2</sub>CH<sub>2</sub>CO, 4H), 2.26 (m, CH<sub>2</sub>CH<sub>2</sub>CO, 4H), 3.70–4.20 (m, CH<sub>2</sub>CH<sub>2</sub>N + CH<sub>2</sub>CH<sub>2</sub>N + *trans*-PO<sub>4</sub>CH<sub>2</sub>CH + COOCH<sub>2</sub>COH, 7H), 4.36 (s, *cis*-PO<sub>4</sub>CH<sub>2</sub>CH, 1H), 5.19 (s, PO<sub>4</sub>CH<sub>2</sub>CH, 1H), 7.09 (d, CH, 2H,  $^3J_{\text{H-H}} = 8.3$  Hz), 7.47

(m, CH, 2H). MALDI (matrix: 4-nitroaniline);  $m/z$  for [C<sub>51</sub>H<sub>93</sub>N<sub>3</sub>O<sub>9</sub>PS<sub>2</sub>]<sup>+</sup> ([M + H + CH<sub>3</sub>CH<sub>2</sub>OH]<sup>+</sup>) expected = 986.6, found 986.6.

#### Synthesis of PE-iso-DOTA-GA (3)

PE-iso (7.0 mg, 7.45  $\mu\text{mol}$ ) was dissolved in 1 mL of a chloroform:methanol mixture (0.5 : 0.5 v/v) and triethylamine (50  $\mu\text{L}$ ). To this solution, 2,2',2''-(10-(4-((2-aminoethyl)amino)-1-carboxy-4-oxobutyl)-1,4,7,10-tetraazacyclododecane-1,4,7-triyl)triacetic acid (NH<sub>2</sub>-DOTA-GA, 10.0 mg, 19.28  $\mu\text{mol}$ ) was added and the mixture was left to stir for 5 h. After that, the crude was concentrated in rotavapour and purified by preparative TLC.  $R_f$  = 0.15–0.25 (CHCl<sub>3</sub> : MeOH : H<sub>2</sub>O 5 : 5 : 0.2 v/v). (6.8 mg, 4.65  $\mu\text{mol}$ , 62%). <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>):  $\delta$  (ppm) 0.71 (t, CH<sub>3</sub>, 6H,  $^3J_{\text{H-H}} = 6.5$  Hz), 1.09 (s, CH<sub>2</sub>, 56H), 1.44 (m, CH<sub>2</sub>CH<sub>2</sub>CO, 4H), 1.69 (m, CH<sub>2</sub>CH<sub>2</sub>, 2H), 1.93–2.55 (m, NCH<sub>2</sub>CH<sub>2</sub>N + NCH<sub>2</sub>CH<sub>2</sub>N + CH<sub>2</sub>CH<sub>2</sub>CO, 20H), 2.58–2.73 (m, NCH<sub>2</sub>COOH, 6H), 2.88 (m, CH<sub>2</sub>CH<sub>2</sub>, 2H), 3.62–3.88 (m, CH<sub>2</sub>CH<sub>2</sub>N + CH<sub>2</sub>CH<sub>2</sub>N + PO<sub>4</sub>CH<sub>2</sub>CH + COOCH<sub>2</sub>COH + CH<sub>2</sub>CH<sub>2</sub>, 8H), 4.17–4.27 (m, CH<sub>2</sub>CH<sub>2</sub>, 2H), 5.06 (s, PO<sub>4</sub>CH<sub>2</sub>CH, 1H), 7.21 (s, CH, 4H). MALDI (matrix: 4-nitroaniline);  $m/z$  for [C<sub>72</sub>H<sub>132</sub>N<sub>10</sub>O<sub>18</sub>PS<sub>2</sub>Na<sub>2</sub>]<sup>+</sup> ([M – 2H + NH<sub>4</sub> + 2Na + CH<sub>3</sub>CH<sub>2</sub>OH]<sup>+</sup>) expected = 1565.9, found 1566.0; [C<sub>72</sub>H<sub>133</sub>N<sub>10</sub>O<sub>18</sub>PS<sub>2</sub>Na]<sup>+</sup> ([M – H + NH<sub>4</sub> + Na + CH<sub>3</sub>CH<sub>2</sub>OH]<sup>+</sup>) expected = 1543.9, found 1544.0.

#### Synthesis of PE-PEG<sub>4</sub>-TCO (4) and PE-TCO (7)

18:0 PE- $\text{NH}_2$  (10 mg, 13.3  $\mu\text{mol}$ ) was dissolved in 7.5 mL of a chloroform : methanol mixture (0.56 : 0.44 v/v) and triethylamine (50  $\mu\text{L}$ ). To this solution, either a solution of TCO-PEG<sub>4</sub>-NHS (10 mg, 19.43  $\mu\text{mol}$  dissolved in 0.3 mL chloroform) or TCO-NHS (5 mg, 18.71  $\mu\text{mol}$ ) was added and the mixture was left to stir for 5 h. After that, the crude was concentrated *in vacuo* and purified by preparative TLC.  $R_f$  for PE-PEG<sub>4</sub>-TCO = 0.45 (CHCl<sub>3</sub> : MeOH : H<sub>2</sub>O 8 : 2 : 0.2 v/v) and,  $R_f$  for PE-TCO = 0.50 (CHCl<sub>3</sub> : MeOH : H<sub>2</sub>O 8 : 2 : 0.2 v/v).

**PE-PEG<sub>4</sub>-TCO.** 13.5 mg, 11.63  $\mu\text{mol}$ , 88%. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>):  $\delta$  (ppm) 0.88 (t, CH<sub>3</sub>, 6H,  $^3J_{\text{H-H}} = 6.5$  Hz), 1.26 (s, CH<sub>2</sub>, 56H), 1.56–1.64 (m, 2  $\times$  CH<sub>2</sub>CH<sub>2</sub>CO + 2  $\times$  CH<sub>2</sub>, 8H), 1.66–1.74 (m, CH<sub>2</sub>, 2H), 1.90–1.98 (m, CH<sub>2</sub>, 2H), 2.25–2.30 (m, 2  $\times$  CH<sub>2</sub>CH<sub>2</sub>CO, 4H), 2.33–2.40 (m, CH<sub>2</sub>, 2H), 2.41–2.52 (m, CH<sub>2</sub>, 2H), 3.46 (m, CH<sub>2</sub>CH<sub>2</sub>N, 2H), 3.52–3.84 (m, OCH<sub>2</sub>CH<sub>2</sub>O + OCH<sub>2</sub>CH<sub>2</sub>O, 18H), 4.00 (m, CH<sub>2</sub>CH<sub>2</sub>N + COOCH<sub>2</sub>COH, 4H), 4.14–4.21 (m, *trans*-PO<sub>4</sub>CH<sub>2</sub>CH, 1H), 4.23–4.30 (m, COOCH<sub>2</sub>, 1H), 4.35–4.41 (m, *cis*-PO<sub>4</sub>CH<sub>2</sub>CH, 1H), 5.23 (m, PO<sub>4</sub>CH<sub>2</sub>CH, 1H), 5.37–5.48 (m, CH, 1H), 5.53–5.64 (m, CH, 1H). MALDI (matrix: 4-nitroaniline);  $m/z$  for [C<sub>62</sub>H<sub>122</sub>N<sub>3</sub>O<sub>16</sub>PN]<sup>+</sup> ([M + NH<sub>4</sub> + CH<sub>3</sub>OH]<sup>+</sup>) expected = 1195.9, found 1195.4.

**PE-TCO.** 12.0 mg, 12.13  $\mu\text{mol}$ , 91%. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>):  $\delta$  (ppm) 0.88 (t, CH<sub>3</sub>, 6H,  $^3J_{\text{H-H}} = 6.5$  Hz), 1.26 (s, CH<sub>2</sub>, 56H), 1.58 (m, 2  $\times$  CH<sub>2</sub>CH<sub>2</sub>CO, 4H), 1.67 (m, CH<sub>2</sub>, 4H), 1.83–2.01 (m, CH<sub>2</sub>, 4H), 2.29 (m, CH<sub>2</sub>CH<sub>2</sub>CO, 4H + CH<sub>2</sub>, 2H), 3.33 (m, CH<sub>2</sub>CH<sub>2</sub>N, 2H), 3.85 (m, CH<sub>2</sub>CH<sub>2</sub>N + COOCH<sub>2</sub>COH, 4H), 4.15 (m, *trans*-PO<sub>4</sub>CH<sub>2</sub>CH, 1H), 4.30 (m, COOCH<sub>2</sub>, 1H), 4.38 (m, *cis*-PO<sub>4</sub>CH<sub>2</sub>CH, 1H), 5.20 (m, PO<sub>4</sub>CH<sub>2</sub>CH, 1H), 5.53 (m,



$CH_3$ , 2H). MALDI (matrix: 4-nitroaniline);  $m/z$  for  $[C_{51}H_{101}N_2O_{11}P]^+$  ( $[M + NH_4 + CH_3OH]^+$ ) expected = 948.7, found 948.4.

### Synthesis of bifunctional chelators

#### Synthesis of 3,3'-((ethane-1,2-diylbis((carboxymethyl)azanediyl))bis(methylene))bis(4-hydroxy-3,1-phenylene))

**dipropionic acid (HBED-CC).** HBED-CC was synthesized by modifying a reported method:<sup>21</sup>

3-(4-Hydroxyphenyl)propionic acid (3.39 g, 20.8 mmol) was dissolved in 21 mL of a solution of methanol:water: 4.0 N NaOH (0.71 : 0.19 : 0.1 v : v : v). To this solution, ethylenediamine-*N,N'*-diacetic acid (1.80 g, 10.4 mmol) suspended in 5 mL of NaOH (2 N) was added. The pH of the mixture was adjusted to 6.7 and the solution was cooled in an ice bath. Then, formaldehyde (aqueous solution at 37%, 20.8 mmol, 1.5 mL) was added dropwise while keeping the pH of the solution at 6.7 by intermittent addition of NaOH (4 N) and, the suspension was refluxed for 4 h at 70 °C. After that, the solution was cooled to room temperature and methanol was removed on a rotary evaporator. The remaining aqueous solution was diluted with water (15 mL), acidified with HCl (4 N) to pH 5.5 and extracted with diethyl ether (15 mL × 3) to remove unreacted 3-(4-hydroxyphenyl)propionic acid. The aqueous phase was then adjusted to pH 3 with HCl (6 N) and unreacted ethylenediamine-*N,N'*-diacetic acid was filtered off. The filtrate was then brought to neutrality with NaOH (4 N) and concentrated *in vacuo*. The crude was allowed to flash chromatography using an Isolera™ Spektra System (Biotage® SNAP Ultra C18 60 g cartridge, A: H<sub>2</sub>O with 0.1% TFA, B: CH<sub>3</sub>CN with 0.1% TFA. Gradient: 2–40% B, 55 CV). (277 mg, 0.52 mmol, 6%). <sup>1</sup>H NMR (400 MHz, dmsO-*d*<sub>6</sub>):  $\delta$  (ppm) 2.45 (t, CH<sub>2</sub>CH<sub>2</sub>COOH,  $^3J_{H-H}$  = 7.8 Hz, 4H), 2.70 (t, CH<sub>2</sub>CH<sub>2</sub>COOH,  $^3J_{H-H}$  = 7.8 Hz, 4H), 3.20 (s, CH<sub>2</sub>N, 4H), 3.65 (s, CH<sub>2</sub>COOH, 4H), 4.02 (s, CH<sub>2</sub>N, 4H), 6.79 (d, CH<sub>arom</sub>,  $^3J_{H-H}$  = 8.2 Hz, 2H), 7.07 (m, CH<sub>arom</sub>, 4H). <sup>13</sup>C NMR (100 MHz, dmsO-*d*<sub>6</sub>):  $\delta$  (ppm) 29.5 (2C), 35.6 (2C), 49.5 (CH<sub>2</sub>N, 2C), 52.2 (2C), 52.7 (2C), 115.4 (2C), 118.7 (2C), 129.8 (2C), 131.4 (2C), 131.9 (2C), 154.5 (2C), 170.2 (2C), 173.8 (2C). HPLC/MS  $\tau_r$  = 7.70 min (gradient A);  $m/z$  for  $[C_{26}H_{33}N_2O_{10}]^+$  ( $[M + H]^+$ ) expected = 533.2135, found 533.2142.

**Synthesis of 3-(3-((carboxymethyl) (2-((carboxymethyl) (2-hydroxy-5-(3-((4-(6-methyl-1,2,4,5-tetrazin-3-yl)benzyl)amino)-3-oxopropyl)benzyl)amino)ethyl)amino)methyl)-4-hydroxyphenyl)propanoic acid (HBED-CC-tetrazine, 5).** Compound 5 was synthesized by modifying a reported method:<sup>13</sup>

HBED-CC (5.3 mg, 9.4  $\mu$ mol) was dissolved in 0.7 mL of a mixture of H<sub>2</sub>O : CH<sub>3</sub>CN (50 : 50 v/v) containing 20  $\mu$ L in *N,N*-diisopropylethylamine. To this solution, 190  $\mu$ L of FeCl<sub>3</sub> (0.1 M in water) was added and the reaction mixture was stirred for 1 h and then evaporated under reduced pressure to dryness. The resulting solid was dissolved in CH<sub>3</sub>CN and evaporated to dryness again to remove free water. 1.25 mL of anhydrous dmsO was then added. To this solution, a solution of HATU (3.5 mg, 9.5  $\mu$ mol) in 0.25 mL of anhydrous dmsO and 0.050 mL *N,N*-diisopropylethylamine was added dropwise. After 15 min, (4-(6-methyl-1,2,4,5-tetrazin-3-yl)phenyl)methanamine (2.3 mg,

11.3  $\mu$ mol) dissolved in 0.5 mL of anhydrous dmsO was added dropwise and the reaction was left to react for 24 h. The reaction was then quenched with 2 mL of H<sub>2</sub>O and the solution was lyophilized. The remaining solid was dissolved in a mixture of H<sub>2</sub>O with 10–15% of dmsO and purified by flash chromatography (Isolera™ Spektra System). Briefly, the crude was loaded in a Biotage® SNAP Ultra C18 30 g cartridge and flushed with 25 mL of HCl (1 M) and washed with 5 mL of H<sub>2</sub>O. The disappearance of the violet colour indicated the removal of iron(III) species. The remaining mixture was eluted with the following gradient: A: H<sub>2</sub>O with 0.1% TFA, B: CH<sub>3</sub>CN with 0.1% TFA. A to B: 2 to 30%, 30 to 70%, 5 CV and 70 to 98%, 5 CV. The purified product was lyophilized and the residue dried *in vacuo* to give a pink solid as 7 (1.5 mg, 2.1  $\mu$ mol, 22%). The product was stable in the freezer for more than six months. <sup>1</sup>H NMR (400 MHz, methanol-*d*<sub>4</sub>):  $\delta$  (ppm) 6.82 (dd, CH,  $^3J_{H-H}$  = 22.3, 8.4 Hz, 2H), 7.18–7.11 (m, CH, 4H), 7.23 (d, CH,  $^3J_{H-H}$  = 8.2 Hz, 2H), 8.42 (d, CH,  $^3J_{H-H}$  = 8.2 Hz, 2H), 4.42 (s, NHCH<sub>2</sub>C, 2H), 4.21 (s, NCH<sub>2</sub>C, 2H), 4.11 (s, NCH<sub>2</sub>C, 2H), 3.78 (s, NCH<sub>2</sub>COOH, 2H), 3.64 (s, NCH<sub>2</sub>COOH, 2H), 3.41 (s, NCH<sub>2</sub>CH<sub>2</sub>N, 2H), 3.02 (s, CH<sub>3</sub>, 3H), 2.91 (t, CH<sub>2</sub>CH<sub>2</sub>CONH,  $^3J_{H-H}$  = 6.8 Hz, 2H), 2.81 (t, CH<sub>2</sub>CH<sub>2</sub>CONH,  $^3J_{H-H}$  = 7.5 Hz, 2H), 2.59 (t, CH<sub>2</sub>CH<sub>2</sub>CONH,  $^3J_{H-H}$  = 6.8 Hz, 2H), 2.59 (t, CH<sub>2</sub>CH<sub>2</sub>CONH,  $^3J_{H-H}$  = 7.5 Hz, 2H). <sup>13</sup>C NMR (100 MHz, methanol-*d*<sub>4</sub>):  $\delta$  (ppm) 21.1, 31.0, 31.8, 36.9, 38.8, 43.5, 48.4, 48.6, 48.8, 49.0, 49.2, 49.4, 49.6, 51.1, 51.4, 54.2, 116.6, 119.9, 120.5, 128.9, 129.0, 131.7, 131.8, 132.2, 133.2, 133.3, 133.6, 133.7, 144.9, 156.0, 165.2, 168.7, 171.6, 172.0, 175.2, 176.7. HPLC:  $\tau_r$  = 8.85 min (gradient A); ESI (negative mode):  $m/z$  for  $[C_{36}H_{40}N_7O_9]^-$  ( $[M - H]^-$ ) expected = 714.2888, found 714.2893.

**Synthesis of DOTA-GA-tetrazine (2,2',2''-(10-(1-carboxy-4-((2-(5-((4-(6-methyl-1,2,4,5-tetrazin-3-yl)benzyl)amino)-5-oxopentanamido)ethyl)amino)-4-oxobutyl)-1,4,7,10-tetraaza-cyclododecane-1,4,7-triyl)triacetic acid) (5)** was carried out according to a method reported by our group, with some minor modifications:<sup>10b</sup>

**Synthesis of *tert*-butyl (4-cyanobenzyl)carbamate.** 4-(Aminomethyl)benzonitrile hydrochloride (5.0 g, 30.0 mmol) was dissolved in CH<sub>2</sub>Cl<sub>2</sub> (4.0 mL), triethylamine (11.0 mL, 75.0 mmol) and di-*tert*-butyl dicarbonate (7.4 g, 33.0 mmol) were added and the resultant mixture stirred at room temperature for 16 h. The solvent was removed under reduced pressure, water added (100 mL) and the product extracted into DCM (3 × 100 mL). The organic layers were combined, washed with water (1 × 100 mL) and the solvent removed under reduced pressure to yield *tert*-butyl (4-cyanobenzyl)carbamate (5.8 g, 84% yield) as a white powder. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>):  $\delta$  (ppm) 1.46 (s, COOC(CH<sub>3</sub>)<sub>3</sub>, 9H), 4.37 (d, CH<sub>2</sub>,  $^3J_{H-H}$  = 6.1 Hz, 2H), 4.97 (s<sub>br</sub>, NH, 1H), 7.38 (d, CH<sub>arom</sub>,  $^3J_{H-H}$  = 8.4 Hz, 2H), 7.62 (d, CH<sub>arom</sub>,  $^3J_{H-H}$  = 8.4 Hz, 2H). <sup>13</sup>C NMR (100 MHz, CDCl<sub>3</sub>):  $\delta$  (ppm) 28.5 (3C), 44.4 (1C), 80.2 (1C), 111.3 (1C), 118.9 (1C), 127.9 (2C), 132.6 (2C), 144.8 (1C), 156.0 (1C). ESI (positive mode):  $m/z$  for  $[C_{13}H_{16}N_2O_2]^+$  ( $[M + H]^+$ ) expected = 233.1290, found 233.1296.

**Synthesis of *tert*-butyl (4-(6-methyl-1,2,4,5-tetrazin-3-yl)benzyl)carbamate.** A high pressure reaction tube was treated with *tert*-butyl (4-cyanobenzyl)carbamate (464 mg, 2.0 mmol), CH<sub>3</sub>CN (1050  $\mu$ L, 20.0 mmol), nickel(II) trifluoromethanesulfonate



(356 mg, 1.0 mmol) and hydrazine hydrate (50–60%  $\text{NH}_2\text{NH}_2$ ) (6.2 mL, 100.0 mmol). The tube was sealed and heated to 60 °C for 72 h, following which  $\text{H}_2\text{O}$  (10 mL) and sodium nitrite (2.82 g, 40.0 mmol) were added to the mixture.  $\text{HCl}$  (1 M) was added dropwise until the pH reached 3 and gases stopped evolving, at which point the mixture had turned bright red. The product was extracted with  $\text{EtOAc}$  (3 × 40 mL), and the combined organic layers were washed with  $\text{H}_2\text{O}$  (3 × 20 mL), dried over  $\text{MgSO}_4$ , filtered and concentrated under reduced pressure. The bright pink crude solid was purified by flash column chromatography (DCM/Et<sub>2</sub>O gradient 100 : 0 v/v to 96 : 4 v/v) to give *tert*-butyl (4-(6-methyl-1,2,4,5-tetrazin-3-yl)benzyl)carbamate (0.37 g, 62% yield) as a bright pink solid.  $R_f$  = 0.39 (2% Et<sub>2</sub>O/DCM). <sup>1</sup>H NMR (400 MHz,  $\text{CDCl}_3$ ):  $\delta$  (ppm) 1.46 (s,  $\text{COOC}(\text{CH}_3)_3$ , 9H), 3.07 (s,  $\text{CH}_3$ , 3H), 4.41 (d,  $\text{CH}_2$ ,  $^3J_{\text{H}-\text{H}} = 5.6$  Hz, 2H), 5.09 ( $s_{\text{br}}$ , NH, 1H), 7.47 (d,  $\text{CH}_{\text{arom}}$ ,  $^3J_{\text{H}-\text{H}} = 8.4$  Hz, 2H), 8.51 (d,  $\text{CH}_{\text{arom}}$ ,  $^3J_{\text{H}-\text{H}} = 8.4$  Hz, 2H). <sup>13</sup>C NMR (100 MHz,  $\text{CDCl}_3$ ):  $\delta$  (ppm) 21.2 (1C), 28.5 (3C), 44.5 (1C), 79.9 (1C), 128.1 (2C), 128.3 (2C), 130.9 (1C), 144.1 (1C), 156.1 (1C) 164.0 (1C), 167.3 (1C). ESI (positive mode):  $m/z$  for  $[\text{C}_{15}\text{H}_{20}\text{N}_5\text{O}_2]^{+}$  ( $[\text{M} + \text{H}]^+$ ) expected = 302.1617, found 302.1621.

**Synthesis of (4-(6-methyl-1,2,4,5-tetrazin-3-yl)phenyl)methanamine.** *Tert*-butyl (4-(6-methyl-1,2,4,5-tetrazin-3-yl)benzyl)carbamate (30.4 mg, 0.15 mmol) was treated with TFA/CH<sub>2</sub>Cl<sub>2</sub> (1/1, v/v, 3 mL) and stirred for 1.5 h at room temperature. The solvent was removed under reduced pressure to yield the TFA salt of 4-(6-methyl-1,2,4,5-tetrazin-3-yl)benzylamine 3 (26.4 mg, 84% yield) as a bright pink solid. <sup>1</sup>H NMR (400 MHz, methanol-*d*<sub>4</sub>):  $\delta$  (ppm) 3.06 (s,  $\text{CH}_3$ , 3H), 4.26 (s,  $\text{CH}_2$ , 2H), 7.72 (d,  $\text{CH}_{\text{arom}}$ ,  $^3J_{\text{H}-\text{H}} = 8.4$  Hz, 2H), 8.62 (d,  $\text{CH}_{\text{arom}}$ ,  $^3J_{\text{H}-\text{H}} = 8.4$  Hz, 2H). <sup>13</sup>C NMR (100 MHz, methanol-*d*<sub>4</sub>):  $\delta$  (ppm) 21.1 (1C), 43.9 (1C), 129.4 (2C), 130.8 (2C), 134.3 (1C), 138.8 (1C), 164.9 (1C), 169.1 (1C). ESI (positive mode):  $m/z$  for  $[\text{C}_{10}\text{H}_{12}\text{N}_5]^{+}$  ( $[\text{M} + \text{H}]^+$ ) expected = 202.1093, found 202.1083;  $m/z$  for  $[\text{C}_{12}\text{H}_{15}\text{N}_6]^{+}$  ( $[\text{M} + \text{H} + \text{CH}_3\text{CN}]^+$ ) expected = 243.138, found 243.1214.

**Synthesis of 5-((4-(6-methyl-1,2,4,5-tetrazin-3-yl)benzyl)amino)-5-oxopentanoic acid.** A dry flask was treated with (4-(6-methyl-1,2,4,5-tetrazin-3-yl)phenyl)methanamine (26.8 mg, 0.13 mmol), glutaric anhydride (76 mg, 10.67 mmol) and THF (2 mL), and the resultant mixture was heated to 70 °C for 4 h. The mixture was cooled to 50 °C, and was stirred at this temperature for a further 16 h. The solvent was evaporated *in vacuo*, to give the crude material as a pink oil. The product was purified by flash chromatography (Isolera™ Spektra System with a Biotage® SNAP Ultra C18 12 g cartridge) eluting with a gradient of 0–50%  $\text{H}_2\text{O}/\text{CH}_3\text{CN}$  (15 CV) to give 4 as a red solid (38.9 mg, 0.12 mmol, >90% yield).  $R_f$  = 0.2 (10% MeOH/CH<sub>2</sub>Cl<sub>2</sub>). HPLC:  $\tau_f$  = 6.50 min (gradient A). <sup>1</sup>H NMR (500 MHz, methanol-*d*<sub>4</sub>):  $\delta$  (ppm) 1.94 (m,  $\text{COCH}_2\text{CH}_2$ , 2H), 2.35 (m, 2 ×  $\text{COCH}_2\text{CH}_2$ , 4H), 3.03 (s,  $\text{CH}_3$ , 3H), 4.49 (s,  $\text{CH}_2\text{NH}$ , 2H), 7.54 (d,  $\text{CH}$ ,  $^3J_{\text{H}-\text{H}} = 8.8$  Hz, 2H), 8.51 (d,  $\text{CH}$ ,  $^3J_{\text{H}-\text{H}} = 8.5$  Hz, 2H). <sup>13</sup>C NMR (126 MHz, methanol-*d*<sub>4</sub>) (2 C<sub>quat</sub> not observed)  $\delta$  (ppm) 21.0 22.3, 34.1, 36.0, 43.8, 129.0, 129.3, 132.4, 145.1, 165.2, 168.7, 175.5, 176.8. ESI (positive mode):  $m/z$  calcd for  $[\text{C}_{15}\text{H}_{18}\text{N}_5\text{O}_3]^{+}$  ( $[\text{M} + \text{H}]^+$ ) 316.1410, found 316.1414 and, calcd for  $[\text{C}_{15}\text{H}_{17}\text{N}_5\text{O}_3\text{Na}]^{+}$  ( $[\text{M} + \text{Na}]^+$ ) 338.1229, found 338.1278.

**Synthesis of 2,2',2''-(10-(1-carboxy-4-((2-(5-((4-(6-methyl-1,2,4,5-tetrazin-3-yl)benzyl)amino)-5-oxopentanamido)ethyl)amino)-4-oxobutyl)-1,4,7,10-tetraazacyclododecane-1,4,7-triyl)**

**triacetic acid (DOTA-GA-tetrazine, 6).** To a stirred solution of 5-((4-(6-methyl-1,2,4,5-tetrazin-3-yl)benzyl)amino)-5-oxopentanoic acid (10.0 mg, 31.71  $\mu\text{mol}$ ) in anhydrous dmso (220  $\mu\text{L}$ ) and triethylamine (7  $\mu\text{L}$ ), was added a solution of *N*-hydroxysuccinimide (18.8 mg, 162.5  $\mu\text{mol}$ ) and *N,N'*-dicyclohexylcarbodiimide (33.5 mg, 162.5  $\mu\text{mol}$ ) in anhydrous dmso (80  $\mu\text{L}$ ). The resultant mixture was vigorously stirred for 16 h at room temperature in the absence of light. After that, the solution was passed through a 0.45  $\mu\text{m}$  filter and added to a solution containing DOTA-GA-NH<sub>2</sub> (17 mg, 32  $\mu\text{mol}$ ) in 100  $\mu\text{L}$  of anhydrous dmso. The reaction was stirred for 5 h at room temperature in the absence of light. The crude was lyophilized and the residue dried *in vacuo* to give a pink oil as crude. The product was purified by flash chromatography using an Isolera™ Spektra System (Biotage® SNAP Ultra C18 12 g cartridge, A:  $\text{H}_2\text{O}$  with 0.1% TFA, B:  $\text{CH}_3\text{CN}$  with 0.1% TFA. Gradient: A to B: 2 to 30%, 40 CV, 30 to 70%, 5 CV and 70 to 98%, 5 CV), to yield 6 as a pink solid (19.6 mg, 24.10  $\mu\text{mol}$ , 76% yield). HPLC:  $\tau_f$  = 7.78 min (gradient A). The product was stable in the freezer for more than six months. <sup>1</sup>H NMR (400 MHz, methanol-*d*<sub>4</sub>) 1.90–1.98 (m,  $\text{COCH}_2\text{CH}_2$ , 2H), 2.25 (t,  $\text{COCH}_2\text{CH}_2$ ,  $^3J_{\text{H}-\text{H}} = 7.6$  Hz, 2H), 2.33 (t,  $\text{COCH}_2\text{CH}_2$ ,  $^3J_{\text{H}-\text{H}} = 7.6$  Hz, 2H), 2.43–3.02 (m, 8H), 3.04 (s,  $\text{CH}_3$ , 3H), 3.12–3.29 (m, 8H), 3.39–4.31 (m, 15H), 4.50 (s,  $\text{NHCH}_2\text{C}$ , 2H), 7.55 (d,  $\text{CH}_{\text{arom}}$ ,  $^3J_{\text{H}-\text{H}} = 8.8$  Hz, 2H), 8.51 (d,  $\text{CH}_{\text{arom}}$ ,  $^3J_{\text{H}-\text{H}} = 8.4$  Hz, 2H). <sup>13</sup>C NMR (125 MHz, methanol-*d*<sub>4</sub>)  $\delta$  (ppm) 21.1, 23.2, 28.5, 30.7, 36.2, 36.3, 40.0, 40.2, 43.8, 49.6, 51.1, 54.3, 54.5, 56.2, 57.5, 129.1, 129.3, 132.4, 145.2, 165.2, 168.8, 175.5, 175.5, 175.6, 175.7, 175.7. ESI (positive mode):  $m/z$  calcd for  $[\text{C}_{36}\text{H}_{54}\text{N}_{11}\text{O}_{11}]^{+}$  ( $[\text{M} + \text{H}]^+$ ) 816.4004, found 816.4010.

## Radiochemistry

<sup>68</sup>GaCl<sub>3</sub> was eluted from a GalliaPharm® <sup>68</sup>Ge/<sup>68</sup>Ga Generator from Eckert & Ziegler Radiopharma GmbH (0.74 GBq) using a fully automated Modular-Lab system and an adapted NaCl-based method previously reported.<sup>20</sup> Briefly, <sup>68</sup>GaCl<sub>3</sub> was eluted in 5 mL of a 0.1 M HCl solution, which was subsequently trapped on a SCX cartridge (preconditioned with 1 mL 5.5 M HCl and 10 mL water) and the activity was then eluted, with minimal loss (2–5%), using a mixture of 12.5  $\mu\text{L}$  of 5.5 M HCl and 500  $\mu\text{L}$  of 5 M NaCl solution. This eluate was subsequently buffered for direct use in <sup>68</sup>Ga-labelling experiments.

**iTLC quantification.** Chelators, freshly prepared from stock solutions for each experiment, were dissolved in aqueous solutions of sodium acetate (0.2 M) to provide solutions with chelator concentrations ranging from 0.05 to 500  $\mu\text{M}$  (0.05, 0.5, 5, 50, 100, 250 and 500  $\mu\text{M}$ ). <sup>68</sup>Ga (15  $\mu\text{L}$ , approx. 2 MBq in 0.1 M aqueous HCl) was added to chelator solutions (100  $\mu\text{L}$ ) and the reaction solution was incubated at either 25 or 90 °C. The final pH of the reaction solutions was 3.5, 4.5 and 5.5. After 10 min, the reaction solution was analysed by iTLC (glass microfiber chromatography paper impregnated with silica gel, 100 × 10 mm). As controls, <sup>68</sup>Ga<sup>3+</sup> (15  $\mu\text{L}$ , approx. 2 MBq in 0.1 M aqueous HCl) was mixed with 100  $\mu\text{L}$  of sodium acetate (0.2 M) at the three tested pHs. The controls were incubated at either 25 or 90 °C for 10 min and then, analysed by iTLC and Laura 3 software (LabLogic, Sheffield, UK). The mobile phase used was ammonium acetate (1 M in



methanol : water 80 : 20 v/v) for all chelators. [ $^{68}\text{Ga}(\text{PE-NOTA})$ ]  $R_f$  = 0.66–0.76; [ $^{68}\text{Ga}(\text{PE-DOTA})$ ]  $R_f$  = 0.67–0.78; [ $^{68}\text{Ga}(\text{PE-iso-DOTAGA})$ ]  $R_f$  = 0.62–0.76; [ $^{68}\text{Ga}(\text{DOTAGA-Tetrazine})$ ]  $R_f$  = 0.67–0.77 [ $^{68}\text{Ga}(\text{HBED-CC-Tetrazine})$ ]  $R_f$  = 0.68–0.78; non-chelated  $^{68}\text{Ga}^{3+}$   $R_f$  < 0.3. iTLC plates were imaged and quantified by digital autoradiography using instruments and software described above.

**$^{68}\text{Ga}$ -labelling of 1.**  $^{68}\text{GaCl}_3$  eluate ( $\sim$ 75 MBq) was added to a solution of 2 mL sodium acetate buffer (0.1 M, pH = 4.0) and 100  $\mu\text{L}$  chelator (0.3 mg  $\text{mL}^{-1}$  in EtOH). During the labelling, the pH of the reaction mixture inside the reactor was determined to be  $3.7 \pm 0.2$ . After incubating the solution for 15 min at 90 °C, the reaction mixture was then diluted with 5 mL of water and passed through an Oasis® HLB (1cc/30 mg) cartridge (preconditioned with 10 mL ethanol and 10 mL water). The cartridge was washed with 5 mL of water and the labelled compound was eluted with 800  $\mu\text{L}$  of pure ethanol. **1** gave radiochemical conversions  $>80\%$ , and the product was obtained in  $>95\%$  radiochemical purity, and in 20–30% isolated RCY (n.d.c.). The synthesis was finished within about 23 min.

**$^{68}\text{Ga}$ -labelling of 5 and 6.**  $^{68}\text{GaCl}_3$  eluate (200–220 MBq) was added to a solution of 4.5 mL sodium acetate buffer (0.1 M, pH = 4.5) containing 30  $\mu\text{L}$  chelator (1 mg  $\text{mL}^{-1}$  in DMSO). During the labelling, the pH of the reaction mixture inside the reactor was determined to be  $4.2 \pm 0.3$ . After incubating the solution for 10 min either at room temperature for **5** or at 90 °C for **6**, the  $^{68}\text{Ga}$  chloride was quantitatively chelated to the precursor. The reaction mixture was then diluted with 5 mL of water and trapped on a Sep-Pack® Light tC18 cartridge (preconditioned with 10 mL ethanol and 10 mL water). The cartridge was washed with 5 mL of water and the labelled compound was eluted with 800  $\mu\text{L}$  of pure ethanol. Both chelators gave radiochemical conversions  $>95\%$ , and the product was obtained in  $>95\%$  radiochemical purity, and in 70–80% isolated RCY (n.d.c.). The synthesis was finished within about 15 min.

**Synthesis of  $^{68}\text{Ga}$ -labelled MBs.** An aliquot of purified [ $^{68}\text{Ga}][\text{Ga}(5)]$  ( $\sim$ 75 MBq) was added to a vial containing PE-PEG<sub>4</sub>-TCO (0.13 mg, 0.107 mol). The reaction vial was incubated for 20 min at 60 °C under a  $\text{N}_2$  stream. After that, the volume of the reaction was reduced to about 100  $\mu\text{L}$  and a solution containing the rest of lipids was added (concretely, 0.62 mg of DPPC and 0.140 mg DSPE-PEG2000-NH<sub>2</sub> dissolved in 1 mL of a solution of propylene glycol, glycerol, and phosphate-buffered saline (PBS) (15 : 5 : 80, v/v/v)). The vial was sealed and activated to form microbubbles (see in the next section the experimental details for this process).

In order to remove unreacted [ $^{68}\text{Ga}][\text{Ga}(5)]$ , microbubbles were purified by centrifugation adapting a reported method.<sup>17</sup> The vial containing the  $^{68}\text{Ga}$ -labelled microbubbles was put upside down and centrifuged at 350 g during 2 min using a ROTINA 35 R centrifuge. Then, the microbubbles were collected into a concentrated cake on the top of the vial and the remaining suspension (infranatant), which contained unreacted precursor and residual lipids and vesicles, which did not form part of the microbubble shell, was discarded. The microbubble cake was re-dispersed to a 1 mL volume of 20 vol% glycerol solution in PBS and stored in the same vial with PFB

headspace until further use. The washing step was repeated three times or until no activity was measured in the infranatant solution.  $^{68}\text{Ga}$ -labelled MBs were counted and sized on a Countess II FL Automated Cell Counter and stored in the same vial with PFB headspace until further use.

(\*) An alternative dienophile-modified phospholipid without the PEG<sub>4</sub> spacer, named PE-TCO (7), was also tested for reaction with [ $^{68}\text{Ga}][\text{Ga}(5)]$ . Under the same conditions, **7** showed low conversions (approx. 10%), probably as a consequence of its poor solubility in ethanol (data not shown).

### Microbubble production

The lipid-coated, decafluorobutane-filled microbubbles (MBs) were manufactured using a modified formulation.<sup>18</sup> Matsunaga *et al.* prepared MBs containing a lipid mixture of 85 mol% 1,2-dipalmitoyl-sn-glycero-3-phosphocholine (DPPC), 10 mol% 1-palmitoyl-2-hydroxy-sn-glycero-3-phosphocholine (LPC), and 5 mol% 1,2-dipalmitoyl-sn-glycero-3-phosphoethanolamine-N[methoxy(polyethylene glycol)-2000] (DSPE-PEG2000-NH<sub>2</sub>). In this paper, we studied the possibility of incorporating 1,2-distearyl-sn-glycero-3-phosphoethanolamine (PE-NH<sub>2</sub>) lipid into the MB shell. For that reason, we prepared MBs having a final lipid mixture of 85 mol% DPPC, 10 mol%, (either PE-NH<sub>2</sub> or PE-PEG<sub>4</sub>-TCO) and 5 mol% DSPE-PEG2000-NH<sub>2</sub>. In all cases, the total lipid concentration was between 0.86 and 0.88 mg  $\text{mL}^{-1}$ . The mixture of lipids was first dissolved in chloroform, dried over nitrogen gas, and then further dried in *vacuo* overnight to remove residual solvent from the lipid films. Then, 1 mL of a solution of propylene glycol, glycerol, and phosphate-buffered saline (PBS) (15 : 5 : 80, v/v/v) was used to rehydrate the films. This created a lipid suspension, which was immediately stirred for 10 min at 50–60 °C and later, decanted in a sealed 2 mL glass vial. Prior to activation, the headspace was purged with decafluorobutane at room temperature and, MBs were produced *via* mechanical agitation.

### Biology experimental procedures

All animal experiments were conducted by licensed investigators in accordance with the United Kingdom Home Office *Guidance on the Operation of the Animals (Scientific Procedures)* Act 1986 (HSMO, London, UK, 1990) and with the published guidelines for the Welfare of Use of Animals in Cancer Research Institute Committee on Welfare of Animals in Cancer Research.<sup>22</sup>

**PET imaging.** Dynamic PET imaging of [ $^{68}\text{Ga}(\text{5})$ ],  $^{68}\text{Ga}$ -PE and  $^{68}\text{Ga}$ -MBs was performed in non-tumour bearing Balb/c mice (5–6 mice/cohort, 6–8 weeks) (Charles River UK Ltd.). Mice were anesthetized with 2.5% isoflurane/O<sub>2</sub> and placed in a thermostatically controlled ring in a dedicated small animal Genisys<sup>4</sup> PET scanner (SOFIE Biosciences, Culver City, USA). Following injection of radioactivity (0.74 MBq in the case of non-MB tracers, and between 0.37–0.74 MBq for  $^{68}\text{Ga}$ -labelled MB, keeping a constant value of  $5 \times 10^7$  MB per injection) *via* lateral tail vein cannula, PET scans were acquired in a list-mode format over 0–30 min to give decay-corrected values of radioactivity accumulation in tissues. The collected data were



ordered into 11 time frames ( $4 \times 15$  s,  $4 \times 60$  s,  $3 \times 300$  s) and reconstructed with a 3-dimensional maximum likelihood estimation method (3D ML-EM). Volumes of interest (VOIs) for each tissue were defined using Siemens Inveon Research Workplace software (Siemens Molecular Imaging Inc. Knoxville, USA) and count densities (counts/min) were expressed as a percentage of the incubated dose (ID) of radioactivity and normalized to tissue weight. Tissue kinetics were calculated by averaging the count densities per timepoint.

**Biodistribution.** Biodistribution studies were carried out on the same cohort of animals after the end of PET scans (20 min after radioactivity injection). Tissue samples were quickly collected and radioactivity content was determined by  $\gamma$ -counting (LKB Wallac 1282 Compugamma laboratory  $\gamma$ -counter, PerkinElmer, Massachusetts, USA). Count densities (counts/min) were expressed as a percentage of the incubated dose (ID) of radioactivity and normalized to tissue weight.

## Conflicts of interest

The authors have no conflicts of interest to declare.

## Acknowledgements

This work was supported by the Cancer Research UK (CRUK Multidisciplinary Award – C53470/A22353).

## References

- (a) *The Chemistry of Molecular Imaging*, ed. N. J. Long and W. T. Wong, Wiley-VCH, Weinheim, 2014; (b) R. E. Mewis and S. J. Archibald, *Coord. Chem. Rev.*, 2010, **254**, 1686–1712.
- (a) L. Abou-Elkacem, S. V. Bachawal and J. K. Willmann, *Eur. J. Radiol.*, 2015, **84**, 1685–1693; (b) J. R. Lindner, *Nat. Rev. Drug Discovery*, 2004, **3**, 527–532; (c) F. Kiessling, S. Fokong, P. Koczera, W. Lederle and T. Lammers, *J. Nucl. Med.*, 2012, **53**, 345–348.
- (a) B. A. Kaufmann, J. M. Sanders, C. Davis, A. Xie, P. Aldred, I. J. Sarembok and J. R. Lindner, *Circulation*, 2007, **116**, 276–284; (b) G. Korpanyi, J. G. Carbon, P. A. Grayburn, J. B. Fleming and R. A. Brekken, *Clin. Cancer Res.*, 2007, **13**, 323–330; (c) G. E. R. Weller, M. K. K. Wong, R. A. Modzelewski, E. Lu, A. L. Klibanov, W. R. Wagner and F. S. Villanueva, *Cancer Res.*, 2005, **65**, 533–539; (d) M. Smeenge, F. Tranquart, C. K. Mannaerts, T. M. de Reijke, M. J. van de Vijver, M. P. Laguna, S. Pochon, J. J. M. C. H. de la Rosette and H. Wijkstra, *Invest. Radiol.*, 2017, **52**, 419–427; (e) J. K. Willmann, L. Bonomo, A. C. Testa, P. Rinaldi, G. Rindi, K. S. Valluru, G. Petrone, M. Martini, A. M. Lutz and S. S. Gambhir, *J. Clin. Oncol.*, 2017, **35**, 2133–2140; (f) K. Ferrara, R. Pollard and M. Borden, *Annu. Rev. Biomed. Eng.*, 2007, **9**, 415–447.
- (a) J. K. Willmann, N. van Bruggen, L. M. Dinkelborg and S. S. Gambhir, *Nat. Rev. Drug Discovery*, 2008, **7**, 591–607; (b) R. Weissleder and M. J. Pittet, *Nature*, 2008, **452**, 580–589.
- S. S. Gambhir, *Nat. Rev. Cancer*, 2002, **2**, 683–693.
- (a) N. Lazarova, P. W. Causey, J. A. Lemon, S. K. Czorny, J. R. Forbes, A. Zlitni, A. Genady, F. Stuart Foster and J. F. Valliant, *Nucl. Med. Biol.*, 2011, **38**, 1111–1118; (b) J. K. Willmann, Z. Cheng, C. Davis, A. M. Lutz, M. L. Schipper, C. H. Nielsen and S. S. Gambhir, *Radiology*, 2008, **249**, 212–219; (c) M. Palmowski, B. Morgenstern, P. Hauff, M. Reinhardt, J. Huppert, M. Maurer, E. C. Woenne, S. Doerk, G. Ladewig, J. W. Jenne, S. Delorme, L. Grenacher, P. Hallscheidt, G. W. Kauffmann, W. Semmler and F. Kiessling, *Invest. Radiol.*, 2008, **43**, 162–169; (d) P. Walday, H. Tolleshaug, T. Gjoen, G. M. Kindberg, T. Berg, T. Skotland and E. Holtz, *Biochem. J.*, 1994, **299**, 437–443; (e) M. S. Tantis, D. E. Kruse, H. Zheng, H. Zhang, A. Kheirolomoom, J. Marik and K. W. Ferrara, *J. Controlled Release*, 2008, **131**, 160–166.
- (a) P. Spang, C. Herrmann and F. Roesch, *Semin. Nucl. Med.*, 2016, **46**, 373–394; (b) M. Fani, J. P. André and H. R. Maecke, *Contrast Media Mol. Imaging*, 2008, **3**, 53–63.
- (a) S. Lin, A. Shah, J. Hernández-Gil, A. Stanzola, B. I. Harriss, T. O. Matsunaga, N. Long, J. Bamber and M.-X. Tang, *J. Photoacoust.*, 2017, **6**, 26–36; (b) P. S. Sheeran, S. Luois, P. A. Dayton and T. O. Matsunaga, *Langmuir*, 2011, **27**, 10412–10420.
- (a) A. Schmidtke, T. Lappchen, C. Weinmann, L. Bier-Schorr, M. Keller, Y. Kiefer, J. P. Holland and M. D. Bartholoma, *Inorg. Chem.*, 2017, **56**, 9097–9110; (b) E. T. Clarke and A. E. Martell, *Inorg. Chim. Acta*, 1991, **181**, 273–280; (c) E. T. Clarke and A. E. Martell, *Inorg. Chim. Acta*, 1991, **190**, 37–46; (d) R. D. Hancock and A. E. Martell, *Chem. Rev.*, 1989, **89**, 1875–1914.
- (a) H. L. Evans, L. Carroll, E. O. Aboagye and A. C. Spivey, *J. Labelled Compd. Radiopharm.*, 2014, **57**, 291–297; (b) H. L. Evans, Q.-D. Nguyen, L. S. Carroll, M. Kaliszczak, F. J. Twyman, A. C. Spivey and E. O. Aboagye, *Chem. Commun.*, 2014, **50**, 9557–9560; (c) J.-P. Meyer, P. Adumeau, J. S. Lewis and B. M. Zeglis, *Bioconjugate Chem.*, 2016, **27**, 2791–2807.
- A.-C. Knall and C. Slugovc, *Chem. Soc. Rev.*, 2013, **42**, 5131–5142.
- R. Selvaraj and J. M. Fox, *Curr. Opin. Chem. Biol.*, 2013, **17**, 753–760.
- M. Eder, B. Wängler, S. Knackmuss, F. LeGall, M. Little, U. Haberkorn, W. Mier and M. Eisenhut, *Eur. J. Nucl. Med. Mol. Imaging*, 2008, **35**, 1878–1886.
- S. R. Banerjee, Z. Chen, M. Pullambhatla, A. Lisok, J. Chen, R. C. Mease and M. G. Pomper, *Bioconjugate Chem.*, 2016, **27**, 1447–1455.
- (a) J. Schuhmacher, G. Klivenyi, R. Matys, M. Stadler, T. Regiert, H. Hauser, J. Doll, W. Maier-Borst and M. Zoller, *Cancer Res.*, 1995, **55**, 115–123; (b) J. Schuhmacher, G. Klivenyi, W. E. Hull, R. Matys, H. Hauser, H. Kalthoff, W. H. Schmiegel, W. Maier-Borst and S. A. Matzku, *Int. J. Radiat. Appl. Instrum., Part B*, 1992, **19**, 809–824.
- M. Eder, O. Neels, M. Müller, U. Bauder-Wüst, Y. Remde, M. Schäfer, U. Hennrich, M. Eisenhut, A. Afshar-Oromieh,



U. Haberkor and K. Kopka, *Pharmaceuticals*, 2014, **7**, 779–796.

17 J. A. Feshitan, C. C. Chen, J. J. Kwan and M. A. Borden, *J. Colloid Interface Sci.*, 2009, **329**, 316–324.

18 (a) S. Lin, A. Shah, J. Hernández-Gil, A. Stanziola, B. I. Harriss, T. O. Matsunaga, N. Long, J. Bamber and M.-X. Tang, *J. Photoacoust.*, 2017, **6**, 26–36; (b) P. S. Sheeran, S. Luois, P. A. Dayton and T. O. Matsunaga, *Langmuir*, 2011, **27**, 10412–10420.

19 C. A. Sennoga, V. Mahue, J. Loughran, J. Casey, J. M. Seddon, M. Tang and R. J. Eckersley, *Ultrasound Med. Biol.*, 2010, **36**, 2093–2096.

20 M. K. Schultza, D. Muellerc, R. P. Baumc, G. L. Watkinsa and W. A. P. Breeman, *Appl. Radiat. Isot.*, 2013, **76**, 46–54.

21 M. Zöller, J. Schuhmacher, J. Reed, W. Maier-Borst and S. Matzku, *J. Nucl. Med.*, 1992, **33**, 1366–1372.

22 P. Workman, E. O. Aboagye, F. Balkwill, A. Balmain, G. Bruder, D. J. Chaplin, J. A. Double, J. Everitt, D. A. H. Farningham, M. J. Glennie, L. R. Kelland, V. Robinson, I. J. Stratford, G. M. Tozer, S. Watson, S. R. Wedge, S. A. Eccles, V. Navaratnam and S. Ryder, *Br. J. Cancer*, 2010, **102**, 1555–1577.

