RSC Advances



PAPER

View Article Online
View Journal | View Issue



Cite this: RSC Adv., 2017, 7, 49586

Comparison of macrocyclic and acyclic chelators for gallium-68 radiolabelling†

Maria Iris Tsionou,^a Caroline E. Knapp, ^b Calum A. Foley,^a Catherine R. Munteanu,^a Andrew Cakebread,^c Cinzia Imberti, ^a Thomas R. Eykyn,^a Jennifer D. Young,^a Brett M. Paterson,^d Philip J. Blower ^a and Michelle T. Ma ^b*

Gallium-68 (⁶⁸Ga) is a positron-emitting isotope used for clinical PET imaging of peptide receptor expression. ⁶⁸Ga radiopharmaceuticals used in molecular PET imaging consist of disease-targeting biomolecules tethered to chelators that complex ⁶⁸Ga³⁺. Ideally, the chelator will rapidly, quantitatively and stably coordinate ⁶⁸Ga³⁺ at room temperature, near neutral pH and low chelator concentration, allowing for simple routine radiopharmaceutical formulation. Identification of chelators that fulfil these requirements will facilitate development of kit-based ⁶⁸Ga radiopharmaceuticals. Herein the reaction of a range of widely used macrocyclic and acyclic chelators with ⁶⁸Ga³⁺ is reported. Radiochemical yields have been measured under conditions of varying chelator concentrations, pH (3.5 and 6.5) and temperature (25 and 90 °C). These chelators are: 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA), 1,4,7-triazacyclononane-1,4,7triacetic acid (NOTA), 1.4.7-triazacyclononane macrocycles substituted with phosphonic (NOTP) and phosphinic (TRAP) groups at the amine, bis(2-hydroxybenzyl)ethylenediaminediacetic acid (HBED), a tris(hydroxypyridinone) containing three 1,6-dimethyl-3-hydroxypyridin-4-one groups (THP) and the hexadentate tris(hydroxamate) siderophore desferrioxamine-B (DFO). Competition studies have also been undertaken to assess relative complexation efficiencies of each chelator for ⁶⁸Ga³⁺ under different pH and temperature conditions. Performing radiolabelling reactions at pH 6.5, 25 °C and 5-50 µM chelator concentration resulted in near quantitative radiochemical yields for all chelators, except DOTA. Radiochemical yields either decreased or were not substantially improved when the reactions were undertaken at lower pH or at higher temperature, except in the case of DOTA. THP and DFO were the most effective ⁶⁸Ga³⁺ chelators at near-neutral pH and 25 °C, rapidly providing near-quantitative radiochemical yields at very low chelator concentrations. NOTP and HBED were only slightly less effective under these conditions. In competition studies with all other chelators, THP demonstrated highest reactivity for ⁶⁸Ga³⁺ complexation under all conditions. These data point to THP possessing ideal properties for rapid, one-step kit-based syntheses of ⁶⁸Ga-biomolecules for molecular PET imaging. LC-MS and ¹H, ¹³C{¹H} and ⁷¹Ga NMR studies of HBED complexes of Ga³⁺ showed that under the analytical conditions employed in this study, multiple HBED-bound Ga complexes exist. X-ray diffraction data indicated that crystals isolated from these solutions contained octahedral [Ga(HBED)(H₂O)], with HBED coordinated in a pentadentate N₂O₃ mode, with only one phenolic group coordinated to Ga³⁺, and the remaining coordination site occupied by a water molecule.

Received 16th August 2017 Accepted 6th October 2017

DOI: 10.1039/c7ra09076e

rsc.li/rsc-advances

Introduction

Gallium-68 (68 Ga) is a positron-emitting isotope with emission properties ($t_{1/2}=68$ min, β^+ 90%, $E_{\rm max}=1880$ keV) that make it suitable for diagnostic imaging with positron emission tomography (PET). A pharmaceutical grade 68 Ge/ 68 Ga generator has recently become commercially available, 1 providing hospitals with on-site access to a GMP-grade diagnostic PET radionuclide without the need for local cyclotron facilities. The most widely utilised 68 Ga radiopharmaceuticals consist of 68 Ga coordinated to a chelator that is attached to a peptide for targeting cell-surface receptors of tumours. Numerous centres already routinely produce diagnostic 68 Ga-HBED-PSMA^{2,3} and 68 Ga-

[&]quot;King's College London, Division of Imaging Sciences and Biomedical Engineering, St Thomas' Hospital, London SE1 7EH, UK. E-mail: michelle.ma@kcl.ac.uk

^bDepartment of Chemistry, University College London, 20 Gordon Street, London WC1H 0AJ, UK

Division of Analytical and Environmental Sciences, King's College London, Franklin Wilkin's Building, London SE1 9NH, UK

^aSchool of Chemistry, Bio21 Molecular Science and Biotechnology Institute, The University of Melbourne, 3010, Victoria, Australia

[†] Electronic supplementary information (ESI) available. CCDC 1564603. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c7ra09076e

RSC Advances Paper

DOTA-TATE4-6 for whole-body PET imaging of prostate and neuroendocrine cancers respectively. These radiotracers have had a significant impact on patient management in centres where they are available, but the complexity of their radiosynthesis in hospitals is a barrier to widespread implementation.

Radiosynthesis of ⁶⁸Ga-DOTA-TATE requires heating at 80-100 °C in order for the DOTA chelator to chelate radiopharmaceutical concentrations of ⁶⁸Ga³⁺ with yields greater than 80%.⁷⁻⁹ On the other hand, ⁶⁸Ga-HBED-PSMA can be prepared at ambient temperatures, but HBED forms multiple species when complexed to Ga³⁺. ¹⁰ This is undesirable as it is possible that the different species have different pharmacological profiles. Heating is employed to increase formation of the most thermodynamically favoured compound, although the structure of this complex has not been defined. Even with heating, populations of other isomers are observed. 10 The radiosyntheses of ⁶⁸Ga-DOTA-TATE and ⁶⁸Ga-HBED-PSMA are undertaken at pH 3-5.

As a result of heating requirements at acidic pH, clinical radiosyntheses of both ⁶⁸Ga-DOTA-TATE and ⁶⁸Ga-HBED-PSMA require multiple manipulations or complex automated equipment.^{9,10} Typical ⁶⁸Ga radiopharmaceutical syntheses involve (i) elution of ⁶⁸Ga from a generator, (ii) pretreatment of eluate to remove contaminating metal impurities that interfere with radiolabelling, as well as ⁶⁸Ge "breakthrough", (iii) addition of ⁶⁸Ga to aqueous solutions of peptide-chelator precursor at pH 3-5, (iv) heating for 5-10 min (followed by cooling) (v) removal of unreacted 68Ga and buffering salts (using solid phase extraction cartridges) and (vi) reconstitution in physiologically compatible solutions for patient administration. In centres that are equipped for more complex preparations of ¹⁸F radiopharmaceuticals, this is not a barrier to routine radiosynthesis but it is time-consuming and costly. However, in regional healthcare centres, or hospitals in countries with developing healthcare systems, such complexity will be a barrier to widespread implementation.

Chelators that quantitatively coordinate 68Ga3+ at nearneutral pH, room temperature and low concentrations of chelator-bioconjugate will enable one-step, kit-based radiolabelling protocols, with concomitant widespread patient benefit. Such radiosyntheses would ideally only require a kit vial containing bioconjugate and buffer components, ⁶⁸Ga generator eluate, a syringe and appropriate radiation shielding. Chelators that fulfil these requirements would also be useful for radiolabelling of small proteins that are susceptible to unfolding or degradation at extremes of pH and temperature. Over the past decade, several chelators have been evaluated and/or developed for ⁶⁸Ga radiolabelling of biomolecules, to overcome the limitations of DOTA (1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid). These include chelators based on 1,4,7-triazacyclononane-1,4,7-triacetic acid (NOTA and its derivative NODAGA),11-15 1,4,7-triazacyclononane (tacn) macrocycles substituted with phosphonic (NOTP16,17) and phosphinic (TRAP18,19) groups at the amine, hexaazamacrobicycles,20 a pyrmacrocycle (PCTA), 15,21 idyl-substituted DOTA hydroxybenzyl)ethylenediaminediacetic acid (HBED) and related compounds possessing phenol, amine and carboxyl

donor groups,3,22 6-amino-1,4-diazepanes with acetate substituents at the amines (DATA),23-26 a siderophore-derived macrocyclic chelator with hydroxamate groups (FSC),27 the acyclic siderophore desferrioxamine-B (which also contains hydroxamates),28,29 and an acyclic chelator based on a substituted pyridine carboxylate with an N₄O₂ binding mode (DEDPA) (Chart 1).30-32 Our research group has recently developed tris(hydroxypyridinone) (THP) derivatives based on 1,6dimethyl-3-hydroxypyridin-4-one units.33-39 Of these chelators, NOTA/NODAGA, TRAP/NOPO, HBED, FSC, DATA, DFO, DEDPA and THP can reportedly be radiolabelled with ⁶⁸Ga³⁺ at ambient temperature. Only DATA derivatives24 and THP derivatives^{33–35,37,38} have been reported to complex ⁶⁸Ga³⁺ above pH 5 at ambient temperature. Many of these ligands provide highly rigid and inert Ga³⁺-chelator complexes. Rigidity is imparted by both selection of appropriate "hard" donor atoms with high affinity for Ga³⁺, which has a relatively high charge density, and the geometry or topology of the chelator itself. Chelators with pre-arranged conformations^{18,26,32} that accommodate octahedral binding of Ga³⁺ favour very high complex rigidity, contributing to kinetic stability of the resulting Ga³⁺ complex.

Most radiosyntheses of ⁶⁸Ga-chelator complexes have been undertaken in acidic solution, below pH 5. This is because hydrated Ga3+ species such as [Ga(H2O)6]3+ predominate in solution below pH 4 but as the pH is raised above 4, the poorly soluble hydroxide species Ga(OH)3 is formed, until the pH exceeds 6.3, where tetracoordinate [Ga(OH)₄] predominates, 40-43 although this is strongly dependant on temperature and is influenced by concentrations of other metal ions and coordinating molecules in solution.44,45 For efficient 68Ga3+ radiolabelling of chelate-peptide conjugates above pH 4, chelate complex formation must effectively compete with 68Gacolloid formation. Preferably, the rate of chelation will be diffusion-controlled, so that complex formation outcompetes ⁶⁸Ga³⁺ colloid formation. The amounts of ⁶⁸Ga eluted from clinical generators are in the range of 400-2000 MBq, approximately equivalent to 4-20 pmol of ⁶⁸Ga³⁺ in 1-5 mL of solution, i.e. nanomolar concentrations. Highly efficient chelators are required to quantitatively coordinate such low concentrations of metal ion without excessively high chelator concentrations.

For most of the chelators mentioned above, their Ga³⁺ complexes and their complexes with some other metal ions, metal stability constants and protonation constants have been reported (Table 1). These data are very useful in predicting the ability of a chelator to coordinate Ga³⁺, the selectivity of a chelator for Ga³⁺ over other metal ions, and the ability of other ligands such as hydroxide ions, to compete for Ga³⁺ binding under physiological conditions. These data do not, however, predict the kinetics of complexation, and without very detailed speciation studies, they do not describe the complexity of the reaction matrix in 68Ga radiolabelling solutions where other adventitious metal ions are present, as well as buffer components.

To the best of our knowledge, there have been no comprehensive side-by-side comparisons of Ga³⁺ chelators to evaluate relative radiolabelling efficiencies. Several prior studies have compared ⁶⁸Ga radiolabelling for a limited number of

chelators. 16,21,24,33,35,46,47 Some of these studies have compared different chelator concentrations or amounts, and all of these studies only explore one or two specific reaction conditions. In

Table 1 $\,$ Proton and $\,$ Ga $^{3+}$ affinity constants of chelators used in this study

Chelator	$\log K_{\rm a}$	$\log K_1$
DOTA ^{48,49}	11.74, 9.76, 4.68, 4.11, 2.37	26.05
NOTA ^{18,50}	13.17, 5.74, 3.22, 1.96	29.63
$NOTP^{51}$	11.7, 9.1, 7.5, 5.8, 3.1, 0.9	_
TRAP ¹⁹	11.48, 5.44, 4.84, 4.23, 3.45, 1.66	26.24
${ m HBED}^{52}$	12.60, 11.00, 8.44, 4.72, 2.53, 1.74	39.57
DFO^{53}	10.79, 9.55, 8.96, 8.32	28.65
Deferiprone ^{a,54,55}	9.86, 3.70; ⁵⁴ $9.78, 3.61$ ⁵⁵	$(\log \beta_3)$ 38.42; ⁵⁴
		37.35 ⁵⁵

^a In the absence of published stability constants for THP, we have included $\log \beta_3$ values for the related [Ga(deferiprone)₃] complex (deferiprone = 1,2-dimethyl-3-hydroxypyridin-4-one).

some of these studies including our own, the ⁶⁸Ga radiolabelling conditions used for each chelator are not identical.^{33,35,46} Therefore, to identify the most suitable chelators to take forward for kit-based ⁶⁸Ga radiolabelling, we have compared the efficiency of ⁶⁸Ga radiolabelling of commercially available DOTA, NOTA, NOTP, TRAP, HBED, THP and DFO, under four different reaction conditions (high and ambient temperatures, and neutral and low pH conditions), and across five orders of magnitude of chelator concentration (50 nM to 500 mM). We also report data that reveal the complexity of Ga³⁺– HBED coordination.

Experimental

Materials and instrumentation

All solvents and reagents were obtained from Sigma-Aldrich (Dorset, UK) unless otherwise indicated. DOTA, NOTA and NOTP were purchased from Macrocyclics (Dallas, USA). TRAP was purchased from CheMatech (Dijon, France). HBED was

purchased from Santa Cruz Biotechnology (Dallas, USA). DFO was purchased from Sigma-Aldrich. THP was synthesised in our laboratory according to a previously reported procedure.33,56 The purchased chemicals were used without further purification. NMR spectra were acquired on Bruker Avance 400 MHz spectrometers (either narrow-bore or wide-bore) (Bruker, Germany) equipped with either a 5 mm QNP probe or a 5 mm BBO probe at 298 K. Spectra were referenced to residual solvent signals or TMS. High-performance liquid chromatography (HPLC) analysis was carried out using an Agilent 1200 LC system with in-line UV and gamma detection (Flow-Count, LabLogic). Instant thin layer chromatography plates (iTLC-SG) were obtained from Agilent Technologies (California, USA) and iTLC strips were visualized and quantified using a Cyclone Plus Storage Phosphor System (Perkin Elmer) interfaced with OptiQuant V5.0 software (Perkin Elmer). Analytical reverse phase HPLC were acquired using an Agilent Eclipse XDB-C₁₈ column (9.4 \times 250 mm, 5 µm) and UV spectroscopic detection at 220 nm. Aliquots (50 µL) of each radiolabelled sample were injected onto the column, using a flow rate of 1 mL min⁻¹, and the following gradient: 0-5 min: 100% A/0% B; 5-25 min: 100% A/0% B to 60% A/40% B. Mobile phase A comprised water with 0.1% trifluoroacetic acid and mobile phase B comprised acetonitrile with 0.1% trifluoroacetic acid. Analytical LC-MS were recorded in the positive ion mode on an Agilent 6510 Q-TOF LC/MS mass spectrometer coupled to an Agilent 1200 LC system (Agilent, Palo Alto, CA) and a LabLogic scintillation detector Flow-count system (Sheffield, UK). An Agilent Eclipse XDB-C₁₈ column (9.4 imes 250 mm, 5 μ m) and UV spectroscopic detection at 220 nm was used with a flow rate of 1 mL min⁻¹, and the following gradient: 0-5 min: 100% A/0% B; 5-25 min: 100% A/0% B to 0% A/100% B. Mobile phase A comprised water with 0.1% formic acid and mobile phase B comprised acetonitrile with 0.1% formic acid. Elemental analysis was performed by the Science Centre, London Metropolitan University.

⁶⁸Ga radiolabelling and iTLC quantification

 ^{68}Ga was eluted from an Eckert & Ziegler $^{68}\text{Ge}/^{68}\text{Ga}$ generator system (Berlin, Germany). Aqueous HCl solution (0.1 M, 5 mL) was passed through the generator and the eluate was collected in 5 \times 1 mL fractions. Aliquots of the second fraction (1 mL, containing 130–230 MBq ^{68}Ga) were used directly for radio-labelling reactions.

Chelators were dissolved in aqueous solutions of sodium acetate (0.2 M) or ammonium acetate (0.2 M) to provide solutions with chelator concentrations ranging from 50 nM to 1 mM (50 nM, 500 nM, 5 μM , 50 μM , 500 μM , and 1 mM). Ligand solutions were freshly prepared from stock solutions for each experiment. 68 Ga (10 μL , approx. 2 MBq in 0.1 M aqueous HCl) was added to chelator solutions (100 μL) and the reaction solution was incubated at either 25 or 90 °C. The final pH of the reaction solutions was 3.5 and pH 6.5 for the sodium acetate and ammonium acetate solutions respectively. After 10 min, the reaction solution was analysed by iTLC (glass microfiber chromatography paper impregnated with silica gel, 80 \times 10 mm).

Separately, solutions of 68 Ga $^{3+}$ (10 μ L, approx. 2 MBq in 0.1 M aqueous HCl) were added to aqueous solutions of sodium acetate (0.2 M) or ammonium acetate (0.2 M), and incubated at either 25 or 90 °C. After 10 min, the solutions were analysed by iTLC

Three different mobile phases were employed for iTLC:

- (1) For THP, DFO and chelator-free reactions under all reaction conditions, aqueous sodium citrate solution (0.1 M, pH 5.5) was used. [68 Ga(chelator)] $R_{\rm f} < 0.1$; non-chelated, soluble 68 Ga $^{3+}$ $R_{\rm f} > 0.9$; 68 Ga colloids: <0.1. Based on quantification of 68 Ga colloid and soluble 68 Ga $^{3+}$ in chelator-free reactions, RCY values were adjusted to account for coincident $R_{\rm f}$ values of 68 Ga colloid, [68 Ga(THP)] and [68 Ga(DFO)].
- (2) For NOTP under all reaction conditions, and DOTA, NOTA, TRAP and HBED, under all conditions except pH 3.5, 90 °C, aqueous sodium phosphate solution (0.4 M, pH 4) was used. [68 Ga(NOTP)] $R_f = 0.6$ –0.7; DOTA, NOTA, TRAP and HBED: [68 Ga(chelator)] $R_f = 0.8$ –1; non-chelated 68 Ga $^{3+}$ $R_f < 0.1$.
- (3) For DOTA, NOTA, TRAP and HBED at pH 3.5 and 90 °C, an ammonium acetate solution (1 M in 80% methanol, 20% water) was used. [68 Ga(DOTA)] $R_{\rm f}=0.65$ –0.75; [68 Ga(NOTA)] $R_{\rm f}=0.8$ –0.9; [68 Ga(HBED)] $R_{\rm f}=0.9$ –1; [68 Ga(TRAP)] $R_{\rm f}=0.4$ –0.6; nonchelated 68 Ga $^{3+}$ $R_{\rm f}=<0.3$. These conditions were selected because after heating 68 Ga solutions at pH 3.5 and 90 °C, two distinct compounds were observed for non-chelated 68 Ga $^{3+}$: $R_{\rm f}=0$ –0.1 and 0.7–0.9 using mobile phase (2). Thus, non-chelated 68 Ga $^{3+}$ could not be distinguished from [68 Ga(chelator)] using mobile phase (2) after heating at pH 3.5 and 90 °C (except in the case of [68 Ga(NOTP)]).

iTLC conditions and $R_{\rm f}$ values are also summarised in Tables S3–S6.†

iTLC plates were imaged and quantified by digital autoradiography using instruments and software described above.

ICP-MS analysis of ⁶⁸Ga generator eluate

Fractionated eluate (as described above) was allowed to decay for several days before it was analysed by ICP-MS. The quantification of metal contaminants was carried out on a PerkinElmer NexION 350D Inductively coupled plasma mass spectrometer (ICP-MS) running Syngistix v1.0 software with a CETAC ASX520 autosampler (King's College London, UK). The acquisition mode included 5 replicates averaged to give reported values (Fig. 2, S12 and S13†) for ²⁷Al, ⁵⁹Co, ⁵²Cr, ⁶⁵Cu, ⁵⁶Fe, ⁶⁹Ga, ⁷²Ge, ⁵⁵Mn, ⁶⁰Ni, ²⁰⁸Pb, ⁴⁵Sc, ¹¹⁸Sn, ⁴⁷Ti, ⁵¹V, ⁶⁶Zn and ⁶⁸Zn. The dwell time was 50 ms per isotope, with 18 L min⁻¹ main argon flow, 1.2 L min⁻¹ auxiliary argon flow, 0.97 L min⁻¹ nebuliser argon flow (optimised), 1600 W RF power, 0.2 mL min⁻¹ sample flow, and KED cell mode with 1.2 mL min⁻¹ helium flow.

HPLC analysis of [⁶⁸Ga(chelator)] complexes and competition studies

 68 Ga³⁺ generator eluate (10 μ L in 0.1 M aqueous HCl, approx. 2 MBq) was added to chelator solutions (1 mM chelator, 100 μ L in either 0.2 M ammonium acetate, or 0.2 M sodium acetate) and the reaction mixtures were incubated at either 25 °C or 90 °C for

RSC Advances Paper

10 min, after which they were applied to an analytical reverse phase C₁₈ HPLC column.

For competition studies, ⁶⁸Ga³⁺ generator eluate (20 µL in 0.1 M aqueous HCl, approx. 4 MBq) was added to a solution containing equimolar concentrations of two chelators (each 500 μM in either 200 μL 0.2 M ammonium acetate, or 200 μL 0.2 M sodium acetate) and the reaction mixtures were incubated at either 25 °C or 90 °C for 40 min, after which they were applied to an analytical reverse phase C18 HPLC column. Data were processed and analysed using Laura Radiochromatography Software (LabLogic).

Preparation of [natGa(HBED)]

A sample of HBED (40 mg, 0.09 mmol) was reacted with Ga(NO₃)₃·xH₂O (40 mg, 0.15 mmol, 1.6 equiv.) in aqueous ammonium acetate solution (0.2 M, 5-10 mL) and heated at 90 °C for 30 min. The solution was then applied to an Agilent Eclipse semi-preparative reverse phase XDB-C₁₈ column $(9.4 \times 250 \text{ mm}, 5 \text{ } \mu\text{m}) \text{ with a 3 mL min}^{-1} \text{ flow rate, and the}$ reaction solution was purified using a gradient elution, in which mobile phase A consisted of water containing 0.1% TFA and mobile phase B consisted of acetonitrile containing 0.1% TFA. The concentration of B increased at a rate of $1\% \text{ min}^{-1}$. [Ga(HBED)] eluted with a retention time of 32 min. Fractions containing pure [Ga(HBED)] were combined, lyophilised, and ¹H and ¹³C{¹H} NMR spectra (in both D₂O/CD₃OD (50%/50%) and D₂O/CD₃CN (60%/40%)), and LC-MS chromatograms were acquired. Data are reported in Fig. 4, 5 and S4-S8.†

Crystals of Ga-HBED of were obtained from a solution of D2O CD₃CN. Anal. calc. for [Ga(HBED)(H₂O)]·CH₃CN (C₂₂H₂₆GaN₃O₇): C, 51.39; H, 5.10; N, 8.17. Found: C, 51.23; H, 5.25; N, 8.10. A suitable crystal containing [Ga(HBED)(H₂O)] was selected and mounted on a nylon loop on a SuperNova Atlas diffractometer with Cu K α radiation ($\lambda = 1.54184$ Å). The crystal was kept at 150.4(5) K during data collection. Using Olex2,57 the structure was solved with the ShelXS58 structure solution program using Direct Methods, and refined with the ShelXL59 refinement package using Least Squares minimisation.

Crystal structure determination

Crystal data for [Ga(HBED)(H2O)]·CH3CN (C22H26GaN3O7) $(M = 514.18 \text{ g mol}^{-1})$: monoclinic, space group $P2_1/c$ (no. 14), a = 12.96001(14) Å, b = 7.01939(10) Å, c = 25.0910(3) Å, $\beta = 97.7700(10)^{\circ}, V = 2261.60(5) \text{ Å}^3, Z = 4, T = 150.4(5) \text{ K},$ $\mu(\text{CuK}\alpha) = 2.093 \text{ mm}^{-1}, D_{\text{calc}} = 1.510 \text{ g cm}^{-3}, 34 357 \text{ reflections}$ measured (6.884° $\leq 2\Theta \leq 147.354^{\circ}$), 4541 unique ($R_{\text{int}} = 0.0461$, $R_{\text{sigma}} = 0.0263$) which were used in all calculations. The final R_1 was 0.0292 $(I > 2\sigma(I))$ and w R_2 was 0.0747 (all data). The identification code is xstr0762. Deposit number for [Ga(HBED)-(H2O)]·CH3CN: CCDC 1564603.†

Results

A comprehensive selection of chelators was reacted with generator-produced solutions of ⁶⁸Ga³⁺ at a range of chelator concentrations, at high (90 °C) and room (25 °C) temperatures,

and in acidic (pH 3.5) and near neutral (pH 6.5) aqueous acetate solutions. In all cases, the reaction time was 10 min. The chelators are: macrocyclic DOTA, NOTA, NOTP and TRAP, and acyclic HBED, DFO and THP (Chart 1). [Note: for ease of nomenclature, charge and protonation states are not included in these abbreviations of the ligands or complexes.

Quantifying the efficiency of ⁶⁸Ga³⁺ chelation

Whilst many ligands will chelate a metal quantitatively if the concentration is high enough, only the most efficient will continue to do so as the concentration is reduced. 12,21,36,46,60,61 A series of reactions was undertaken in which a solution of generator-produced ⁶⁸Ga³⁺ (approx. 2 MBq in 0.1 M aqueous HCl, 10 µL) was added to a solution of chelator at a concentration in the range 500 μ M to 50 nM (100 μ L). The final pH of the reaction solution was 3.5 (using 0.2 M sodium acetate) or 6.5 (using 0.2 M ammonium acetate). After 10 min reaction time, radiochemical yields (RCY) were measured using instant thin layer chromatography (iTLC). Experimental data are supplied for each chelator in Fig. 1, S1, S2† and Table 2. A summary of iTLC conditions is provided in Tables S3-S6.†

Room temperature radiolabelling. At pH 3.5, 25 °C, and a chelator concentration of 50 μM , RCYs were greater than 85% for all chelators except DFO (Fig. 1, S1, S2† and Table 2). At 5 μM and 50 µM chelator concentrations, the best performing chelators (i.e. that demonstrated highest labelling efficiency) were NOTA and TRAP. At concentrations of 50 µM, RCY of $^{[68}$ Ga(NOTA)] was 97 \pm 1.7%, and RCY of $^{[68}$ Ga(TRAP)] was 95 \pm 1.3%. At 5 μ M, RCY of [68 Ga(NOTA)] was 93 \pm 2.0%, and RCY of [68 Ga(TRAP)] was 92 \pm 2.4%.

At pH 6.5, 25 °C, and a chelator concentration of 5 μM, RCYs were greater than 85% for all chelators except for DOTA. At pH 6.5 and 25 °C, at very low chelator concentrations of 500 nM and 5 µM, the best performing chelators were DFO and THP. At concentrations of 5 μ M, RCY of [68 Ga(DFO)] was 97 \pm 1.0%, and RCY of $[^{68}$ Ga(THP)] was 97 \pm 0.1%. At 500 nM, RCY of [68 Ga(DFO)] was 96 \pm 1.5% and the RCY of [68 Ga(THP)] was $97 \pm 0.6\%$.

The pK_a values for deprotonation of coordinating O donor atoms of NOTA and TRAP are substantially lower than those of DFO and THP (Table 1). At pH 3.5, both NOTA and TRAP complexed 68Ga3+ with efficiency comparable to that achieved at pH 6.5. At pH 3.5, at concentrations below 500 μM, RCYs of THP and DFO with ⁶⁸Ga³⁺ were relatively poor, however with an increase in pH, Ga³⁺ competed more effectively with protons for coordination to THP and DFO at lower concentrations of chelator. Thus, at pH 6.5, the lowest concentration at which a RCY greater than 95% was reached was 500 nM, achieved using DFO and THP.

High temperature radiolabelling. For some macrocycles, the energies of activation for chelation of metal ions are significantly higher than those of linear chelators. To overcome these substantial kinetic barriers when radiolabelling DOTA conjugates with ⁶⁸Ga³⁺, reaction solutions are heated. On the other hand, studies have demonstrated that radioisotopes of Ga³⁺ bind to NOTA and its derivatives at room temperature 12,15 - the

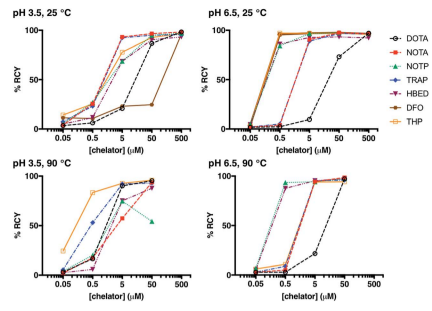


Fig. 1 Radiochemical yields for the reaction of 68 Ga $^{3+}$ with DOTA, NOTA, NOTP, TRAP, HBED, DFO and THP under different concentrations of chelator (500 μ M to 50 nM); different pH conditions (pH 3.5 or pH 6.5); and different temperatures (25 °C or 90 °C), after 10 min reaction.

kinetic barriers to ${\rm Ga}^{3^+}$ complexation are likely lower for NOTA than DOTA.^{11,49} To evaluate the contribution of kinetics to radiolabelling efficiencies at room temperature, radiolabelling reactions (at chelator concentrations of 50 μ M to 50 nM) were also undertaken at 90 °C for all chelators except DFO (Fig. 1, S1, S2† and Table 2). HPLC studies (described below) suggested that either DFO or its ${\rm Ga}^{3^+}$ complex decompose at 90 °C.

At pH 3.5, 90 °C, and a chelator concentration of 50 μM , RCYs were greater than 85% for all chelators except NOTP, and the most efficient chelators were THP, DOTA and TRAP. RCY of $[^{68}\text{Ga}(\text{THP})]$ was $96\pm0.5\%$, RCY of $[^{68}\text{Ga}(\text{DOTA})]$ was $95\pm0.9\%$ and RCY of $[^{68}\text{Ga}(\text{TRAP})]$ was $93\pm0.6\%$. At 5 μM , RCY of $[^{68}\text{Ga}(\text{THP})]$ was $93\pm1.1\%$, RCY of $[^{68}\text{Ga}(\text{DOTA})]$ was $90\pm1.5\%$ and RCY of $[^{68}\text{Ga}(\text{TRAP})]$ was $92\pm1.5\%$. Thus, at pH 3.5, heating substantially improves RCY at chelator concentrations of 500 nM to 5 μM for DOTA and THP. The increased RCY observed for $[^{68}\text{Ga}(\text{DOTA})]$ at 5 μM at 90 °C (90 \pm 1.5%) compared to 25 °C (21 \pm 5.6%) is consistent with previous reports. 8,9,19 This suggests that the labelling efficiencies of these ligands at pH 3.5 at room temperature are limited by kinetic barriers.

Interestingly, the RCY (57 \pm 2.6%) of $[^{68}\text{Ga}(\text{NOTA})]$ at 5 $\mu\text{M},$ 90 °C, pH 3.5 was substantially decreased compared to that observed at 25 °C (93 \pm 2.0%). It is possible that contaminating metal ions present in generator eluate (see below) effectively compete with Ga^{3+} for NOTA binding at high temperature, but that at lower temperature, the kinetic barriers to complexation of these other metal ions prevent them from competing with $\text{Ga}^{3+}.$

At pH 6.5, 90 $^{\circ}$ C, and a chelator concentration of 5 μ M, RCYs were greater than 94% for all chelators except DOTA. In contrast to results observed at room temperature, the best performing chelators were NOTP and HBED. At a chelator concentration of

500 nM, RCY of [68 Ga(NOTP)] was 94 \pm 0.8% and RCY of [68 Ga(HBED)] was 88 \pm 4.4%.

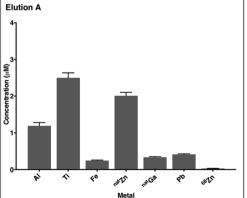
The RCY of $[^{68}Ga(THP)]$ at 500 nM and pH 6.5 was substantially reduced at 90 °C (11 \pm 2.6%) compared to RCY at 25 °C (97 \pm 0.6%). Again, it is possible that THP complexes of metal ion contaminants are formed at high temperature, but not low temperature.

⁶⁸Ga generator ICP-MS eluate analysis. Different batches of ⁶⁸Ga eluate were used for all of the above experiments. Prior work has shown that the concentrations of contaminating metal ions increase with increasing time between 68Ga generator elutions.^{9,62} In our experiments, the time between elutions was 2-24 hours. To better characterise these reaction solutions, and identify metal ions that compete with ⁶⁸Ga³⁺ for chelator complexation, the concentration of selected metals ions in two batches of generator eluate was quantified, with each eluate fractionated into five samples (each 1 mL). Eluates from two separate elutions were assessed: elution A was obtained 5 hours after the previous elution, and elution B, 150 hours (five days) after the previous elution. The concentrations of Al, Sc, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, Zn (nat Zn and ⁶⁸Zn), Ga, Ge, Sn and Pb in generator eluate solution and the hydrochloric acid solution used as eluate were quantified using ICP-MS.

In most eluate fractions, including the second fraction used for radiolabelling, Al, Ti, Fe, Zn, Ga and Pb were present at concentrations of 0.1 μ M to 5 μ M (Fig. 2, S12 and S13†). Concentrations of 68 Zn (arising from decay of 68 Ga) were significantly higher in eluate B than eluate A. This is expected: eluate B contained decay products of 150 h of 68 Ge/ 68 Ga decay, whereas eluate A contained decay products of only 5 h of 68 Ge/ 68 Ga decay. Concentrations of nat Ga were also higher in eluate B compared to eluate A. Although the sampling size here

Table 2 Radiochemical yields (±standard deviation) for the reactions of ⁶⁸Ga³⁺ with DOTA, NOTA, NOTP, TRAP, HBED, DFO and THP. Experiments were undertaken in triplicate

Chelator	Concentration (µM)	pH 3.5, 25 °C	рН 3.5, 90 °C	pH 6.5, 25 °C	pH 6.5, 90 °C
DOTA	500	98.3 ± 0.4		97.0 ± 0.7	
	50	86.7 ± 5.0	95.3 ± 0.9	73.2 ± 6.4	97.2 ± 0.3
	5	20.7 ± 5.6	90.1 ± 1.5	9.6 ± 5.9	21.6 ± 2.5
	0.5	6.2 ± 2.8	16.5 ± 2.2	2.3 ± 0.7	2.8 ± 0.2
	0.05	3.5 ± 1.1	3.1 ± 0.7	1.7 ± 0.5	2.8 ± 0.5
NOTA	500	98.2 ± 0.6		96.6 ± 1.2	
	50	96.2 ± 1.7	93.7 ± 0.3	97.5 ± 0.1	98.5 ± 0.1
	5	93.2 ± 2.0	57.3 ± 2.6	90.6 ± 4.5	94.4 ± 0.4
	0.5	25.4 ± 35.6	17.4 ± 2.1	3.8 ± 0.3	4.9 ± 0.4
	0.05	4.0 ± 2.2	2.2 ± 0.5	1.9 ± 0.7	2.8 ± 0.2
NOTP	500	97.0 ± 0.8		96.7 ± 1.8	
	50	95.0 ± 1.8	54.3 ± 2.4	97.5 ± 0.1	97.3 ± 0.4
	5	68.7 ± 25.0	74.9 ± 3.4	96.6 ± 0.4	94.2 ± 0.9
	0.5	26.8 ± 33.0	20.0 ± 1.9	84.3 ± 0.9	93.5 ± 0.8
	0.05	5.2 ± 3.7	3.9 ± 0.7	4.8 ± 0.4	6.8 ± 1.2
TRAP	500	95.6 ± 0.7		96.5 ± 0.4	
	50	95.0 ± 1.3	93.0 ± 0.6	96.6 ± 0.1	96.0 ± 1.4
	5	92.5 ± 2.4	92.0 ± 1.5	89.0 ± 1.4	95.2 ± 0.4
	0.5	23.0 ± 9.5	53.1 ± 4.8	5.1 ± 0.8	8.8 ± 2.0
	0.05	7.4 ± 1.5	5.5 ± 0.6	2.0 ± 0.5	2.8 ± 0.5
HBED	500	93.2 ± 3.8		92.2 ± 0.6	
	50	90.4 ± 8.1	87.7 ± 0.5	93.3 ± 3.0	97.2 ± 0.4
	5	68.6 ± 23.5	75.2 ± 0.8	92.7 ± 2.2	95.4 ± 0.8
	0.5	11.6 ± 11.9	5.8 ± 0.4	86.0 ± 3.9	87.5 ± 4.4
	0.05	4.9 ± 2.2	2.5 ± 0.4	4.6 ± 0.7	5.1 ± 0.8
DFO	500	96.0 ± 0.8		96.4 ± 1.4	
	50	24.5 ± 1.9		97.5 ± 0.7	
	5	23.0 ± 12.9		97.0 ± 1.0	
	0.5	10.5 ± 5.4		95.8 ± 1.5	
	0.05	11.2 ± 5.0		3.6 ± 1.3	
THP	500	96.7 ± 1.4	96.2 ± 0.8	95.8 ± 1.6	94.3 ± 0.7
	50	92.7 ± 4.2	95.9 ± 0.5	97.1 ± 1.1	94.2 ± 0.6
	5	77.8 ± 13.5	92.7 ± 1.1	97.1 ± 0.1	94.0 ± 0.3
	0.5	25.4 ± 6.8	83.4 ± 1.1	97.1 ± 0.6	10.7 ± 2.6
	0.05	14.0 ± 6.8	24.3 ± 3.4	3.8 ± 3.6	6.1 ± 1.1



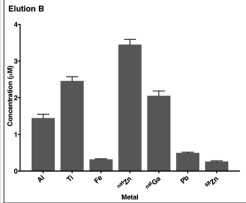


Fig. 2 The concentrations of selected metals in 68 Ga generator eluate from the second 1 mL fraction (measured by ICP-MS). Elution A was obtained 5 hours after the previous elution, and elution B, 150 hours (five days) after the previous elution. Error bars represent standard deviation of the measurement (n = 5). 68 Zn concentrations correspond to 68 Zn arising from 68 Ga decay.

is very low, the measured metal contaminant levels fall within a similar range to that of prior reports. The source of these metal ion contaminants includes the HCl solution used as eluate

(contributing a proportion of measured Al, Fe, Zn and Pb), and components of the ⁶⁸Ga generator including titanium dioxide on a borosilicate glass column, lead shielding and tubing.

Complexes of Al³⁺, Ti⁴⁺, Fe³⁺, Zn²⁺, Ga³⁺ and Pb²⁺ with many of these chelators or their derivatives have been described and characterised. Some of these chelators (for example, HBED,⁵² hydroxypyridinones,⁶³ NOTA¹⁸ and TRAP^{18,19}) have demonstrated selectivity for Ga³⁺ over divalent ions. However, it is likely that the presence of Al³⁺, Fe³⁺ and ^{nat}Ga³⁺, and the high concentrations of Ti⁴⁺, Zn²⁺ and Pb²⁺, will decrease RCY of the desired ⁶⁸Ga-chelator complex for all chelators used in these experiments. It is also likely that differences in metal concentrations in different eluates leads to variability in RCYs.

HPLC radiochromatograms of ⁶⁸Ga complexes

Before performing ⁶⁸Ga competition experiments with reverse-phase HPLC, each chelator was reacted with ⁶⁸Ga³⁺ solution, and HPLC radiochromatograms (Fig. S3†) were acquired to determine the retention times and chromatographic behaviour of each complex. For [⁶⁸Ga(DFO)] at 90 °C, multiple signals were observed in the radiochromatograms at both pH 3.5 and 6.5, with wide-ranging retention times. We did not pursue further experiments to elucidate the nature of these species, nor did we study any further reactions of DFO at 90 °C. Others have previously described the structure of [Ga(DFO)]. ⁶⁴ We postulate that the ligand or complex are not stable at 90 °C, and that the signals correspond to decomposition products.

Ga-HBED complexes

Prior studies have reported that derivatives of the HBED chelator form isomers when complexed to Ga³⁺ in solution, ^{10,65} and we have previously suggested that these correspond to geometric isomers (Fig. 3), ⁶⁶ however little empirical evidence is available to support this. Consistent with all previous reports of Ga³⁺-bound HBED (Ga–HBED) derivatives, multiple species with distinct HPLC retention times were formed (Fig. 4). At 25 °C, at least three distinct signals in the radiochromatogram of ⁶⁸Ga³⁺–HBED could be distinguished but at 90 °C, only two of these signals were observed (Fig. S4†). The distribution of these species at each temperature was the same whether they were synthesised at pH 3.5 or 6.5.

To further characterise these reaction products, a solution of HBED was reacted with 1.6 equivalents $Ga(NO_3)_3$ at 90 °C, and the resulting complex was isolated (using semi-preparative reverse phase C_{18} HPLC as a mixture of species, including stereo- and possible geometric-isomers). The isolated material was characterised by NMR and LC-MS.

The LC-MS retention times (absorbance at $\lambda = 220$ nm and total ion count) were coincident with HPLC signals observed from reaction solutions of ⁶⁸Ga³⁺ with HBED at 90 °C (Fig. 4). The product was resolved into two distinct molecular ions by LC-MS, with both corresponding to the expected isotopic pattern for $\{[Ga(HBED)]^- + 2H^+\}$ (Fig. 4c inset). In the ${}^{13}C\{{}^{1}H\}$ spectrum of Ga-HBED, there are four signals corresponding to C=O groups, four signals corresponding to C-O phenolic groups and ten signals (rather than the expected twelve) corresponding to methylene groups. Presumably there are two pairs of coincident signals in the case of the methylene groups. In the ¹H NMR spectra (including COSY and heteronuclear HSQC ¹H-¹³C spectra, Fig. S5-S8†), the chemically distinct methylene protons display geminal coupling, and the spectra are also consistent with formation of multiple species. ¹³C{¹H} and ¹H spectra were acquired in a mixture of D₂O/CD₃CN (60%/40%), and separately, in a mixture of D₂O/CD₃OD (50%/50%). Similar spectra were observed for both samples, with no notable differences in the number of signals, nor their chemical shifts and relative intensities. In the 71 Ga spectrum (acquired in D₂O/ CD₃CN), a very broad signal is observed, likely arising from overlaid broad resonances corresponding to different Ga-HBED complexes (Fig. 4). A small amount of unchelated Ga3+, presumably $[Ga(H_2O)_6]^{3+}$, is also present. The broad signal from ⁷¹Ga reflects the asymmetric environment in all Ga³⁺-HBED species, causing significant quadrupolar relaxation, compared to the symmetric agua ion.

¹H, ¹³C(¹H), COSY, HSQC and ⁷¹Ga NMR spectra (Fig. 5 and S5-S8†) are consistent with formation of at least two different species. We postulate that these species could include the three possible geometric isomers of a hexadentate N2O4 species (Fig. 3), as well as complexes in which the HBED ligand coordinates to Ga³⁺ with lower density, (with each species consisting of NMR-indistinguishable Δ and Λ enantiomers). For each of geometric isomers designated [Ga(HBED)]-1 and [Ga(HBED)]-2, there are three chemically distinct methylene environments, one phenolic environment and one carboxylate environment. For the geometric isomer designated [Ga(HBED)]-3, there are six chemically distinct methylene environments, two phenolic environments and two carboxylate environments. For a Ga-HBED pentadentate species (with a monodentate ligand occupying the sixth coordination site), there are six chemically distinct methylene environments, two phenolic environments and two carboxylate environments.

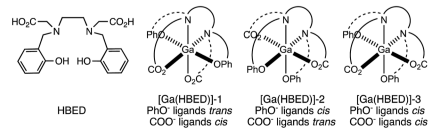


Fig. 3 Possible geometric isomers for hexadentate [Ga(HBED)]. Note that each isomer depicted here is one of a pair of enantiomers.

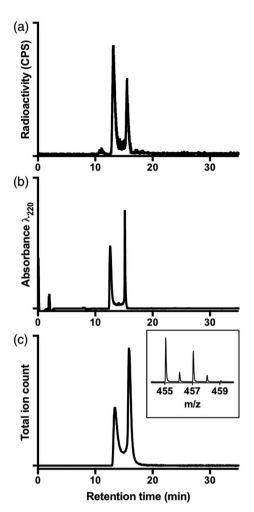


Fig. 4 (a) Radiochromatogram of [68 Ga(HBED)] (prepared at 90 °C); (b) UV chromatogram of [Ga(HBED)]; (c) extracted ion chromatogram of isolated [Ga(HBED)] (m/z of monoisotopic signal = 455.07); inset positive ion MS spectrum of {[Ga(HBED)] + 2H⁺}: [Ga(C₂₀H₂₂N₂O₆)]⁺.

Slow evaporation of a solution of isolated Ga-HBED material in water and acetonitrile provided crystals suitable for X-ray diffraction (Table S1†). The selected crystal contained the neutral [Ga(HBED)(H₂O)] complex crystallised with a molecule of acetonitrile, and the unit cell contained four symmetryrelated equivalents of [Ga(HBED)(H2O)]·CH3CN, including both Δ and Λ enantiomers of the Ga^{3+} complex. In [Ga(HBED)(H₂O)], HBED is bound to Ga³⁺ in a pentadentate N₂O₃ environment, with only one phenolic group coordinated to Ga³⁺ (Fig. 6). The non-coordinating phenolic group is protonated and uncharged. A water molecule occupies the remaining Ga³⁺ coordination site to give an octahedral Ga³⁺ complex in which the two carboxylate groups are coordinated trans to each other, and the single coordinating phenolic group and H₂O ligand are cis to each other. There is some distortion of the octahedral environment. N-Ga-O bond angles of the fivemembered chelate rings, formed by the coordinating carboxylate and amine ligands, are significantly smaller (81.73 $^{\circ}$ and 82.30°) than other bond angles about the metal centre (Table S2†), likely owing to the steric strain in the ligand.

Elemental analysis of the isolated crystalline material indicated that the bulk crystalline material has the same elemental (C, H, N) composition as the single crystal used to acquire X-ray diffraction data (see Experimental section).

Similar geometric arrangements of the HBED ligand have been observed for HBED complexes of Ti^{4+} and Fe^{3+} , although in these structures, both phenolic groups are coordinated to the metal centre. In hexadentate [Ti(HBED)] and [Fe(HBED)], the two carboxylate groups are *trans* to each other, and the two phenolic groups are *cis* to each other. The metal-HBED ligand bond lengths in these octahedral complexes are similar to corresponding metal-HBED ligand bond lengths of [Ga(HBED)(H₂O)].

HPLC chromatograms of solutions of isolated $[Ga(HBED)(H_2O)] \cdot CH_3CN$ gave the same peak shapes as previously observed (Fig. S4†).

Competition studies

Competition studies³⁶ were undertaken, in which two chelators were allowed to compete for $^{68}Ga^{3^+}$ binding, at different temperatures (90 °C and 25 °C) and different pH conditions (pH 3.5 and 6.5). Generator eluate containing $^{68}Ga^{3^+}$ was added to solutions containing equimolar concentrations of two chelators. The concentration of each chelator was 500 μM . At this concentration, the chelators were in large excess over $^{68}Ga^{3^+}$ and the above iTLC studies demonstrated that RCYs for every chelator were near-quantitative. After 40 min reaction (to allow for equilibration), solutions were analysed by reverse phase HPLC (for example, Fig. 7). Radio-chromatographic signals for each ^{68}Ga species were integrated, and results summarised as a percentage of total radioactivity (Tables 3 and 4).

Relative labelling efficiency among DOTA, NOTA, NOTP and TRAP was not compared, as their complexes could not be adequately separated from each other by HPLC or iTLC. ⁶⁸Ga radiolabelled complexes of HBED, DFO and THP showed distinct retention times under the HPLC conditions employed, and labelling efficiencies of these chelators could be compared with each other and with each of DOTA, NOTA, NOTP and TRAP. These competitive comparisons were repeated at pH 3.5 and 6.5, and 25 and 90 °C, except for those involving DFO at 90 °C.

Under all conditions, THP "competed" most effectively for 68 Ga $^{3+}$ in comparison with all other chelators (Fig. S9†). In each competition study involving THP at 25 °C, RCY of 68 Ga(THP)] was in the range 92–100%. At 90 °C, the proportion of 68 Ga $^{3+}$ complexed by the three triazacyclononane (tacn) derivatives (NOTA, NOTP and TRAP) in competition with THP was significantly higher than at 25 °C, although RCYs of 68 Ga(THP)] still exceeded 60% in these reactions. This suggests that, to some extent, these 68 Ga reaction products at 25 °C are a result of kinetic preferences.

For competition reactions between THP and DOTA, and between THP and HBED, RCYs of $[^{68}$ Ga(THP)] were 100% under all conditions.

The three tacn derivatives competed favourably for ⁶⁸Ga³⁺ in reactions with either HBED or DFO (Fig. S10 and S11†). Indeed in all such competitive reactions, RCYs of ⁶⁸Ga with either

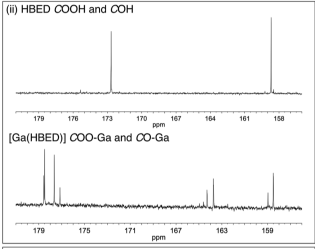
Paper

(a) ¹³C NMR: (i) HBED *C*H₂

69 67 65 63 61 ppm 59 57 55 53 51

[Ga(HBED)] CH₂

69 67 65 63 61 ppm 59 57 55 53 51



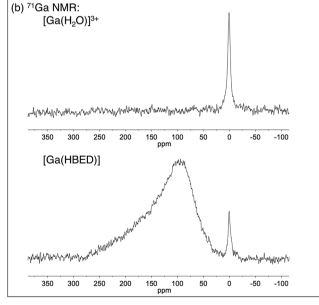


Fig. 5 Regions from (a) 13 C{ 1 H} NMR spectra of HBED and [Ga(HBED)], and (b) 71 Ga NMR spectrum of [Ga(HBED)] (60% D₂O/40% CD₃CN). In the 13 C{ 1 H} NMR spectrum of HBED, three CH_2 resonances, one COH resonance and one COOH resonance are detected, but upon coordination to Ga^{3+} , an increase in the number of signals is observed. A residual methanol signal is marked (x). In the 71 Ga NMR spectrum of [Ga(HBED)], a broad, asymmetric peak is observed, distinct from that of unchelated [Ga(H_2O)₆] $^{3+}$.

NOTA, NOTP or TRAP exceeded 55%. Generally, of the three tacn derivatives, radiochemical yields of [⁶⁸Ga(NOTP)] were highest across all competition studies.

Under the tested conditions, HBED "outcompeted" only DOTA and DFO (Fig. S9 and S10†). At 25 °C under both pH conditions, DFO "outcompeted" only DOTA (Fig. S9–S11†). Compared to all other chelators and under all tested conditions, DOTA was least able to compete for ⁶⁸Ga³⁺ complexation.

There were no remarkable differences between competition studies undertaken at pH 3.5 and pH 6.5, except in the case of reactions of ⁶⁸Ga³⁺ with HBED and TRAP (Fig. S10†). At pH 3.5, all ⁶⁸Ga³⁺ was bound to the TRAP chelator, whereas at pH 6.5, only 60–80% of added ⁶⁸Ga³⁺ was bound to TRAP (depending on the temperature).

Discussion

There is a prevailing notion in the radiochemical literature, based largely on knowledge of the pH-dependence of the hydrolytic behaviour of the Ga³⁺ aqua ion in forming relatively insoluble hydroxides, that ^{68/67}Ga³⁺ chelation is most effective at pH 5 or lower. Most reactions that assess ^{68/67}Ga complexation are undertaken at low pH values. Our results demonstrate that for ⁶⁸Ga-radiolabelling of NOTA, NOTP, TRAP, HBED, DFO and THP at 25 °C under these specific reaction conditions, RCYs at pH 6.5 are equal to or greater than RCYs achieved at pH 3.5.

Many reported 68 Ga radiolabelling experiments have been conducted at temperatures greater than 50 °C. Our results demonstrate that for the majority of chelators, heating the reaction does not significantly increase RCY. The exceptions to this are reactions of 68 Ga³⁺ with DOTA at both pH 3.5 and pH 6.5, and THP at pH 3.5. In our hands, heating and low pH conditions are only favourable in the case of DOTA – the chelator that is currently used the most for clinical 68 Ga biomolecule labelling. Even under low pH and high temperature conditions, near-quantitative RCYs for DOTA labelling (greater than 95%) were only achieved at concentrations of 50 μ M and above. In these studies, in which low amounts of 68 Ga were used, this corresponded to maximum specific activities of 20–40 MBq μ mol $^{-1}$.

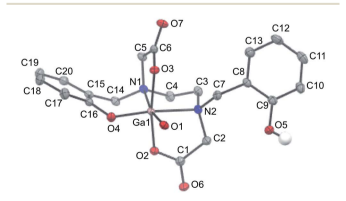


Fig. 6 ORTEP representation of $[Ga(HBED)(H_2O)] \cdot CH_3CN$. Ellipsoids are at the 50% probability level. Hydrogen atoms (with the exception of the proton of the non-coordinating phenolic group) and solvent molecules are omitted for clarity.

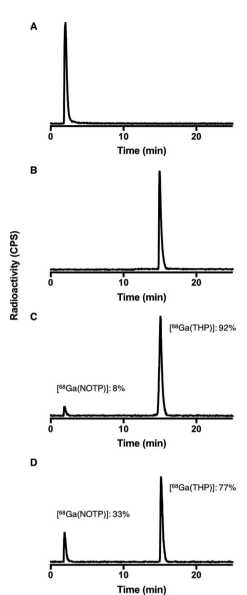


Fig. 7 Exemplar HPLC radiochromatograms from competition studies: (A) [68 Ga(NOTP)] standard; (B) [68 Ga(THP)] standard; reaction solutions (pH 6.5) containing equimolar concentrations (500 μ M) of THP and NOTP with 68 Ga $^{3+}$ eluate at (C) 25 °C and (D) 90 °C. Radiochromatograms from competition studies are included in ESI, Fig. S9–S11.†

In contrast, near-quantitative RCYs were achieved for THP and DFO at pH 6.5 and 25 °C, at chelator concentrations as low as 500 nM, and in aqueous solutions (ammonium acetate solution) that are physiologically compatible. This corresponded to specific activities of approximately 2–4 GBq μmol⁻¹. For reactions that achieved near quantitative RCYs, the maximum specific activity for [⁶⁸Ga(THP)] and [⁶⁸Ga(DFO)] (under mild conditions) was two orders of magnitude higher than that achieved for [⁶⁸Ga(DOTA)] (under low pH and high temperature conditions). High radiochemical yields (>80%) were achieved for NOTP and HBED under the same mild ⁶⁸Galabelling conditions. Identifying chelators such as these, that enable reproducible and near-quantitative ⁶⁸Ga biomolecular

Table 3 ⁶⁸Ga competition studies at pH 3.5: equimolar solutions of HBED, THP or DFO with either DOTA, NOTA, NOTP, TRAP, HBED, THP or DFO, were reacted with ⁶⁸Ga³⁺. The RCYs of [⁶⁸Ga(HBED)], [⁶⁸Ga(THP)] and [⁶⁸Ga(DFO)] for each of these reactions are given in the below table

	25 °C			90 °C		
	HBED	THP	DFO	HBED	THP	
DOTA NOTA NOTP TRAP HBED THP	99% HBED 43% HBED 0% HBED 0% HBED — 0% HBED	100% THP 97% THP 94% THP 100% THP 100% THP	100% DFO 38% DFO 24% DFO 30% DFO 17% DFO 0% DFO	96% HBED 32% HBED 0% HBED 0% HBED — 0% HBED	100% THP 79% THP 64% THP 82% THP 100% THP	

Table 4 68 Ga competition studies at pH 6.5: equimolar solutions of HBED, THP or DFO with either DOTA, NOTA, NOTP, TRAP, HBED, THP or DFO, were reacted with 68 Ga $^{3+}$. The RCYs of [68 Ga(HBED)], [68 Ga(THP)] and [68 Ga(DFO)] for each of these reactions are given in the below table

	25 °C			90 °C	
	HBED	THP	DFO	HBED	THP
DOTA NOTA NOTP TRAP HBED THP	99% HBED 41% HBED 0% HBED 38% HBED —	100% THP 99% THP 92% THP 100% THP 100% THP	100% DFO 33% DFO 26% DFO 25% DFO 18% DFO 0% DFO	96% HBED 38% HBED 0% HBED 20% HBED —	100% THP 87% THP 77% THP 88% THP 100% THP

labelling under low chelator concentration, mild conditions and in physiologically compatible solutions will facilitate (i) one-step, kit-based radiosynthesis of ⁶⁸Ga radiopharmaceuticals; and (ii) ⁶⁸Ga radiolabelling of small proteins (<50 kDa). Proteins that accumulate at target tissue and clear circulation in less than four hours (including engineered antibody derivatives and recombinant proteins) have utility in imaging *in vivo* receptor expression, ^{47,69–71} but many are likely to be sensitive to extremes of pH and temperature.

⁶⁷Ga-labelled DFO-protein conjugates have previously demonstrated *in vivo* and *in vitro* instability, with DFO releasing ⁶⁷Ga³⁺ (between 20–60% dissociation of ⁶⁷Ga³⁺ to serum proteins over three days in solutions containing serum proteins). ²⁸ This is likely a result of kinetic instability of the [Ga(DFO)] complex, leading to transmetallation of Ga³⁺ to endogenous ligands (proteins, peptides and bone mineral) *in vivo* and *in vitro*. However, the shorter half-life of ⁶⁸Ga renders prolonged *in vivo* stability unnecessary. As DFO complexes ⁶⁸Ga³⁺ in near quantitative yields under mild conditions at low concentrations, DFO is possibly very useful for molecular imaging with ⁶⁸Ga.⁷²

At pH 6.5 and 25 °C, NOTP is more efficient at chelating $^{68}\text{Ga}^{3^+}$ than either of the other tacn derivatives, NOTA and TRAP, and the radiochemical yield for $[^{68}\text{Ga}(\text{NOTP})]$ is greater than 95% at chelator concentrations of 5 μ M. This corresponded to specific activities of 200–400 MBq μ mol⁻¹. NOTP is

also potentially very useful for ⁶⁸Ga biomolecule labelling under mild conditions. Additionally, out of all chelators except THP, NOTP competes most effectively for ⁶⁸Ga³⁺ in competition reactions. Whilst other tacn derivatives, TRAP and NOTA, have been extensively studied for ⁶⁸Ga biomolecule labelling, ⁶⁸Ga bioconjugates of NOTP have not been reported. Reassessment of its utility would be timely.

At pH 6.5 at both high and low temperatures, HBED efficiently chelates ⁶⁸Ga³⁺, achieving 85% RCY at concentrations as low as 500 nM. LC-MS and NMR data for HBED complexes of ^{68/nat}Ga³⁺ indicate that under the analytical conditions described here, more than one chemically distinct species exists in solution. It is possible that neutral pentadentate complexes, such as [Ga(HBED)(H₂O)] observed by single crystal X-ray diffraction (Fig. 6), exist within this population of Ga-HBED species. In light of the geometric arrangement of donor atoms in crystals of [Ga(HBED)(H₂O)]·CH₃CN, and existing data on hexadentate HBED complexes of Ti⁴⁺ and Fe³⁺, ^{67,68} we postulate that hexadentate [Ga(HBED)]-2 (Fig. 3) is also present in solution.

The analytical chromatographic and spectroscopic conditions used in this study (and in prior studies 10,65) do not mimic physiological conditions. Nonetheless, the data demonstrate the intricate speciation of HBED complexes of Ga^{3+} . Detailed speciation studies under physiologically-relevant conditions are required. The $\log K_a$ values for phenolic protons of HBED are 12.6 and $11.0,^{52}$ and it is possible that the different species of Ga-HBED (and indeed Ga-HBED-PSMA) arise from fluxionality in the coordination/dissociation of the phenolic oxygen ligands. It is likely that in solutions at higher pH values than those studied here, Ga-HBED exists exclusively as a hexadentate complex. It is also possible that a hexadentate complex would be observed in the solid state, if proton counter ions were substituted for other cations.

Competition studies, in which an equimolar solution of two chelators was reacted with ⁶⁸Ga³⁺, indicated that out of all tested chelators and under all tested conditions (which were not concentration-limited), THP competed most effectively for Ga³⁺. These data, alongside THP radiolabelling studies that demonstrate very high ⁶⁸Ga radiolabelling efficiency, point to THP possessing ideal properties for rapid radiolabelling under mild conditions.33-35,37,39 It is likely that other chelators that have demonstrated suitable properties for kit-based radiolabelling, such as DEDPA,30-32 FSC,27 and DATA,24,25 which we have not tested in this study, possess similar properties. Such properties will enable rapid, one-step kit-based syntheses of ⁶⁸Gabiomolecules for molecular PET imaging without the requirement for complex automated equipment or specialist radiochemistry expertise. This will be key to providing many more hospitals and patients with access to ⁶⁸Ga radiopharmaceuticals.

Whilst thermodynamic and kinetic studies can predict the utility of a chelator for binding very low concentrations of metal ions, it is difficult to compile all necessary data to reliably model the complex reaction matrix of radiolabelling solutions. These solutions contain adventitious metal ions present in concentrations exceeding that of ⁶⁸Ga, and buffer/salt components, which can coordinate both Ga³⁺ and other metal ions, and determine metal ion speciation and reactivity. The simple

radiolabelling experiments that we have described here enable identification of suitable and efficient chelators for kit-based ⁶⁸Ga³⁺ radiolabelling, in a reaction matrix typical of radio-pharmaceutical formulations.

Author contributions

M. I. T. and M. T. M. performed radiolabelling experiments, C. E. K. performed crystallography, C. A. F. and M. T. M. synthesised HBED complexes of Ga(III), C. R. M. and A. C. performed ICP-MS analyses, C. I., J. D. Y. and B. M. P. helped to devise experiments and co-authored the manuscript, T. R. E. performed NMR analysis, P. J. B. and M. T. M. conceived of this study and authored the manuscript.

Conflicts of interest

P. J. B. is a named inventor on related patents. All other authors have no conflicts to declare.

Acknowledgements

This research was supported by the Wellcome Trust through the London Metallomics Facility funded by the Wellcome Trust Multi-user Equipment Grant (202902/Z/16/Z), the Centre of Excellence in Medical Engineering Centre funded by the Wellcome Trust and EPSRC (203148/Z/16/Z), the KCL and UCL Comprehensive Cancer Imaging Centre funded by CRUK and EPSRC in association with the MRC and DoH (England), and by the NIHR Biomedical Research Centre at Guy's and St Thomas' NHS Foundation Trust and King's College London. B. M. P. was supported by a Victorian Postdoctoral Research Fellowship (Victorian Government). The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the DoH. We thank Stephen Boyer, Science Centre, London Metropolitan University, for acquiring elemental microanalysis data.

References

- 1 I. Velikyan, Molecules, 2015, 20, 12913-12943.
- W. P. Fendler, D. F. Schmidt, V. Wenter, C. Zach,
 P. Bartenstein, F. J. Gildehaus, K. M. Thierfelder, C. Stief,
 C. Gratzke, T. Kirchner and C. Faber, *J. Nucl. Med.*, 2016,
 57, 1720–1725.
- 3 A. Afshar-Oromieh, C. M. Zechmann, A. Malcher, M. Eder,
 M. Eisenhut, H. G. Linhart, T. Holland-Letz,
 B. A. Hadaschik, F. L. Giesel, J. Debus and U. Haberkorn,
 Eur. J. Nucl. Med. Mol. Imaging, 2014, 41, 11–20.
- 4 A. R. Haug, C. J. Auernhammer, B. Waengler, G. P. Schmidt, C. Uebleis, B. Goeke, P. Cumming, P. Bartenstein, R. Tiling and M. Hacker, *J. Nucl. Med.*, 2010, 51, 1349–1356.
- 5 M. S. Hofman, G. Kong, O. C. Neels, P. Eu, E. Hong and R. J. Hicks, *J. Med. Imaging Radiat. Oncol.*, 2012, **56**, 40–47.
- 6 R. Srirajaskanthan, I. Kayani, A. M. Quigley, J. Soh, M. E. Caplin and J. Bomanji, *J. Nucl. Med.*, 2010, **51**, 875–882.

- 7 K. P. Zhernosekov, D. V. Filosofov, R. P. Baum, P. Aschoff, H. Bihl, A. A. Razbash, M. Jahn, M. Jennewein and F. Roesch, J. Nucl. Med., 2007, 48, 1741–1748.
- 8 E. Eppard, M. Wuttke, P. L. Nicodemus and F. Roesch, *J. Nucl. Med.*, 2014, 55, 1023–1028.
- 9 I. Velikyan, G. J. Beyer and B. Langstroem, *Bioconjugate Chem.*, 2004, **15**, 554–560.
- 10 M. Eder, O. Neels, M. Mueller, U. Bauder-Wuest, Y. Remde, M. Schaefer, U. Hennrich, M. Eisenhut, A. Afshar-Oromieh, U. Haberkorn and K. Kopka, *Pharmaceuticals*, 2014, 7, 779–796.
- 11 J.-F. Morfin and É. Tóth, *Inorg. Chem.*, 2011, **50**, 10371-10378.
- 12 I. Velikyan, H. Maecke and B. Langstrom, *Bioconjugate Chem.*, 2008, **19**, 569–573.
- 13 K.-P. Eisenwiener, M. I. M. Prata, I. Buschmann, H.-W. Zhang, A. C. Santos, S. Wenger, J. C. Reubi and H. R. Maecke, *Bioconjugate Chem.*, 2002, 13, 530–541.
- 14 R. A. Dumont, F. Deininger, R. Haubner, H. R. Maecke, W. A. Weber and M. Fani, *J. Nucl. Med.*, 2011, 52, 1276–1284.
- 15 C. L. Ferreira, D. T. T. Yapp, D. Mandel, R. K. Gill, E. Boros, M. Q. Wong, P. Jurek and G. E. Kiefer, *Bioconjugate Chem.*, 2012, 23, 2239–2246.
- 16 M. Fellner, P. Riss, N. S. Loktionova, K. P. Zhernosekov, O. Thews, F. G. C. Geraldes Carlos, Z. Kovacs, I. Lukes and F. Rösch, *Radiochim. Acta*, 2011, 99, 43–51.
- 17 M. I. M. Prata, A. C. Santos, C. F. G. C. Geraldes and J. J. P. De Lima, *J. Inorg. Biochem.*, 2000, **79**, 359–363.
- 18 J. Simecek, M. Schulz, J. Notni, J. Plutnar, V. Kubicek, J. Havlickova and P. Hermann, *Inorg. Chem.*, 2012, 51, 577– 590.
- 19 J. Notni, P. Hermann, J. Havlickova, J. Kotek, V. Kubicek, J. Plutnar, N. Loktionova, J. Riss Patrick, F. Rosch and I. Lukes, *Chem.-Eur. J.*, 2010, 16, 7174–7185.
- 20 M. T. Ma, O. C. Neels, D. Denoyer, P. Roselt, J. A. Karas, D. B. Scanlon, J. M. White, R. J. Hicks and P. S. Donnelly, *Bioconjugate Chem.*, 2011, 22, 2093–2103.
- 21 C. L. Ferreira, E. Lamsa, M. Woods, Y. Duan, P. Fernando, C. Bensimon, M. Kordos, K. Guenther, P. Jurek and G. E. Kiefer, *Bioconjugate Chem.*, 2010, 21, 531–536.
- 22 M. Eder, M. Schaefer, U. Bauder-Wuest, W.-E. Hull, C. Waengler, W. Mier, U. Haberkorn and M. Eisenhut, *Bioconjugate Chem.*, 2012, 23, 688–697.
- 23 B. P. Waldron, D. Parker, C. Burchardt, D. S. Yufit, M. Zimny and F. Roesch, *Chem. Commun.*, 2013, 49, 579–581.
- 24 J. Seemann, B. P. Waldron, F. Roesch and D. Parker, *ChemMedChem*, 2015, **10**, 1019–1026.
- 25 J. Seemann, B. Waldron, D. Parker and F. Roesch, *EJNMMI Radiopharmacy and Chemistry*, 2016, 1, 4.
- 26 D. Parker, B. P. Waldron and D. S. Yufit, *Dalton Trans.*, 2013, **42**, 8001–8008.
- 27 P. A. Knetsch, C. Zhai, C. Rangger, M. Blatzer, H. Haas, P. Kaeopookum, R. Haubner and C. Decristoforo, *Nucl. Med. Biol.*, 2015, 42, 115–122.
- 28 S. V. Govindan, R. B. Michel, G. L. Griffiths, D. M. Goldenberg and M. J. Mattes, *Nucl. Med. Biol.*, 2005, 32, 513–519.

- 29 J. E. Ryser, K. Rose, R. Jones, A. Pelegrin, A. Donath, R. Egeli, A. Smith and R. E. Offord, *Nucl. Med. Biol.*, 1998, 25, 261–265.
- 30 E. Boros, C. L. Ferreira, J. F. Cawthray, E. W. Price, B. O. Patrick, D. W. Wester, M. J. Adam and C. Orvig, *J. Am. Chem. Soc.*, 2010, 132, 15726–15733.
- 31 E. Boros, C. L. Ferreira, D. T. T. Yapp, R. K. Gill, E. W. Price, M. J. Adam and C. Orvig, *Nucl. Med. Biol.*, 2012, 39, 785–794.
- 32 C. F. Ramogida, J. F. Cawthray, E. Boros, C. L. Ferreira, B. O. Patrick, M. J. Adam and C. Orvig, *Inorg. Chem.*, 2015, 54, 2017–2031.
- 33 D. J. Berry, Y. Ma, J. R. Ballinger, R. Tavare, A. Koers, K. Sunassee, T. Zhou, S. Nawaz, G. E. D. Mullen, R. C. Hider and P. J. Blower, *Chem. Commun.*, 2011, 47, 7068–7070.
- 34 M. T. Ma, C. Cullinane, C. Imberti, J. Baguna Torres, S. Y. A. Terry, P. Roselt, R. J. Hicks and P. J. Blower, *Bioconjugate Chem.*, 2016, 27, 309–318.
- 35 M. T. Ma, C. Cullinane, K. Waldeck, P. Roselt, R. J. Hicks and P. J. Blower, *EJNMMI Res.*, 2015, 5, 52.
- 36 M. T. Ma, L. K. Meszaros, B. M. Paterson, D. J. Berry, M. S. Cooper, Y. Ma, R. C. Hider and P. J. Blower, *Dalton Trans.*, 2015, 44, 4884–4900.
- 37 C. Imberti, S. Y. A. Terry, C. Cullinane, F. Clarke, G. H. Cornish, N. K. Ramakrishnan, P. Roselt, A. P. Cope, R. J. Hicks, P. J. Blower and M. T. Ma, *Bioconjugate Chem.*, 2017, 28, 481–495.
- 38 R. Cusnir, C. Imberti, P. J. Blower, M. T. Ma and R. C. Hider, *Int. J. Mol. Sci.*, 2017, **18**, 116.
- 39 J. D. Young, V. Abbate, C. Imberti, L. K. Meszaros, M. T. Ma, S. Y. A. Terry, R. C. Hider, G. E. Mullen and P. J. Blower, J. Nucl. Med., 2017, 58, 1270–1277.
- 40 B. Hacht, Bull. Korean Chem. Soc., 2008, 29, 372-376.
- 41 R. E. Weiner and M. L. Thakur, in *Handbook of Radiopharmaceuticals: Radiochemistry and Applications*, ed. M. J. Welch and C. S. Redvanly, John Wiley & Sons Ltd., 2003, pp. 363–399.
- 42 G. E. Jackson and M. J. Byrne, *J. Nucl. Med.*, 1996, 37, 379–386.
- 43 S. M. Moerlein and M. J. Welch, *Int. J. Nucl. Med. Biol.*, 1981, **8**, 277–287.
- 44 S. A. Wood and I. M. Samson, *Ore Geol. Rev.*, 2006, **28**, 57–102.
- 45 L. E. McInnes, S. E. Rudd and P. S. Donnelly, *Coord. Chem. Rev.*, 2017, DOI: 10.1016/j.ccr.2017.05.011, advance web publication.
- 46 J. Notni, K. Pohle and H.-J. Wester, *EJNMMI Res.*, 2012, 2(28), 25.
- 47 M. Eder, A. V. Krivoshein, M. Backer, J. M. Backer, U. Haberkorn and M. Eisenhut, *Nucl. Med. Biol.*, 2010, 37, 405–412.
- 48 A. Bianchi, L. Calabi, C. Giorgi, P. Losi, P. Mariani, P. Paoli, P. Rossi, B. Valtancoli and M. Virtuani, *J. Chem. Soc., Dalton Trans.*, 2000, 697–705.
- 49 V. Kubíček, J. Havlíčková, J. Kotek, G. Tircsó, P. Hermann, É. Tóth and I. Lukeš, *Inorg. Chem.*, 2010, **49**, 10960–10969.
- 50 B. Drahos, V. Kubicek, C. S. Bonnet, P. Hermann, I. Lukes and E. Toth, *Dalton Trans.*, 2011, **40**, 1945–1951.

- 51 C. F. G. C. Geraldes, A. D. Sherry and W. P. Cacheris, *Inorg. Chem.*, 1989, 28, 3336–3341.
- 52 C. H. Taliaferro and A. Martell, *Inorg. Chim. Acta*, 1984, **85**, 9–15.
- 53 A. Evers, R. D. Hancock, A. E. Martell and R. J. Motekaitis, *Inorg. Chem.*, 1989, **28**, 2189–2195.
- 54 D. J. Clevette, D. M. Lyster, W. O. Nelson, T. Rihela, G. A. Webb and C. Orvig, *Inorg. Chem.*, 1990, **29**, 667–672.
- 55 Y.-Y. Xie, Z. Lu, X.-L. Kong, T. Zhou, S. Bansal and R. Hider, *Eur. J. Med. Chem.*, 2016, **115**, 132–140.
- 56 T. Zhou, H. Neubert, D. Y. Liu, Z. D. Liu, Y. M. Ma, X. L. Kong, W. Luo, S. Mark and R. C. Hider, *J. Med. Chem.*, 2006, 49, 4171–4182.
- 57 O. V. Dolomanov, L. J. Bourhis, R. J. Gildea, J. A. K. Howard and H. Puschmann, *J. Appl. Crystallogr.*, 2009, **42**, 339–341.
- 58 G. M. Sheldrick, Acta Crystallogr., Sect. A: Found. Crystallogr., 2008, 64, 112–122.
- 59 G. M. Sheldrick, Acta Crystallogr., Sect. C: Struct. Chem., 2015, 71, 3–8.
- 60 M. S. Cooper, M. T. Ma, K. Sunassee, K. P. Shaw, J. D. Williams, R. L. Paul, P. S. Donnelly and P. J. Blower, *Bioconjugate Chem.*, 2012, 23, 1029–1039.
- 61 M. T. Ma, M. S. Cooper, R. L. Paul, K. P. Shaw, J. A. Karas, D. Scanlon, J. M. White, P. J. Blower and P. S. Donnelly, *Inorg. Chem.*, 2011, 50, 6701–6710.

- 62 E. Oehlke, V. S. Le, N. Lengkeek, P. Pellegrini, T. Jackson, I. Greguric and R. Weiner, *Appl. Radiat. Isot.*, 2013, 82, 232–238.
- 63 E. T. Clarke and A. E. Martell, *Inorg. Chim. Acta*, 1992, 191, 56–63.
- 64 B. Borgias, A. D. Hugi and K. N. Raymond, *Inorg. Chem.*, 1989, 28, 3538–3545.
- 65 J. Schuhmacher, G. Klivenyi, W. E. Hull, R. Matys, H. Hauser, H. Kalthoff, W. H. Schmiegel, W. Maier-Borst and S. Matzku, *Nucl. Med. Biol.*, 1992, 19, 809–824.
- 66 M. T. Ma and P. J. Blower, in *Metal Chelation in Medicine*, ed. R. R. Crichton, R. J. Ward and R. C. Hider, The Royal Society of Chemistry, 2017, pp. 260–312.
- 67 S. K. Larsen, B. G. Jenkins, N. G. Memon and R. B. Lauffer, Inorg. Chem., 1990, 29, 1147–1152.
- 68 A. D. Tinoco, C. D. Incarvito and A. M. Valentine, J. Am. Chem. Soc., 2007, 129, 3444–3454.
- 69 A. C. Freise and A. M. Wu, Mol. Immunol., 2015, 67, 142-152.
- 70 M. Rosestedt, K. G. Andersson, B. Mitran, V. Tolmachev, J. Loefblom, A. Orlova and S. Staahl, Sci. Rep., 2015, 5, 15226.
- 71 P. M. Smith-Jones, D. B. Solit, T. Akhurst, F. Afroze, N. Rosen and S. M. Larson, *Nat. Biotechnol.*, 2004, 22, 701–706.
- 72 P. M. Smith-Jones, B. Stolz, C. Bruns, R. Albert, H. W. Reist, R. Fridrich and H. R. Maecke, *J. Nucl. Med.*, 1994, 35, 317–325.