Towards the stable chelation of radium for biomedical applications with an 18-membered macrocyclic ligand†

Diane S. Abou, †a,b,c Nikki A. Thiele, †d,e Nicholas T. Gutsche, † Alexandria Villmer, ab Hanwen Zhang, † ab Joshua J. Woods, † dmg Kwamena E. Baidoo, † Freddy E. Escorcia, †ff Justin J. Wilson †df and Daniel L. J. Thorek †a,bhi

Targeted alpha therapy is an emerging strategy for the treatment of disseminated cancer. [223Ra]RaCl2 is the only clinically approved alpha particle-emitting drug, and it is used to treat castrate-resistant prostate cancer bone metastases, to which [223Ra]Ra2+ localizes. To specifically direct [223Ra]Ra2+ to non-osseous disease sites, chelation and conjugation to a cancer-targeting moiety is necessary. Although previous efforts to stably chelate [223Ra]Ra2+ for this purpose have had limited success, here we report a biologically stable radiocomplex with the 18-membered macrocyclic chelator macropa. Quantitative labeling of macropa with [223Ra]Ra2+ was accomplished within 5 min at room temperature with a radiolabeling efficiency of >95%, representing a significant advancement over conventional chelators such as DOTA and EDTA, which were unable to completely complex [223Ra]Ra2+ under these conditions. [223Ra][Ra(macropa)] was highly stable in human serum and exhibited dramatically reduced bone and spleen uptake in mice in comparison to bone-targeted [223Ra]RaCl2, signifying that [223Ra][Ra(macropa)] remains intact in vivo. Upon conjugation of macropa to a single amino acid β-alanine as well as to the prostate-specific membrane antigen-targeting peptide DUPA, both constructs retained high afinity for 223Ra, complexing >95% of Ra2+ in solution. Furthermore, [223Ra][Ra(macropa-β-alanine)] was rapidly cleared from mice and showed low 223Ra bone absorption, indicating that this conjugate is stable under biological conditions. Unexpectedly, this stability was lost upon conjugation of macropa to DUPA, which suggests a role of targeting vectors in complex stability in vivo for this system. Nonetheless, our successful demonstration of efficient radiolabeling of the β-alanine conjugate with 223Ra and its subsequent stability in vivo establishes for the first time the possibility of delivering [223Ra]Ra2+ to metastases outside of the bone using functionalized chelators, marking a significant expansion of the therapeutic utility of this radionuclide in the clinic.

Introduction

[223Ra]RaCl2 is the first and currently only approved α particle-emitting radiopharmaceutical, with an indication for men with bone-metastatic castrate-resistant prostate cancer.1 Since its approval in 2013, [223Ra]RaCl2 has been used to treat over 18,000 patients, substantially improving the quality of life for those suffering from bone pain, reducing fracture risk, and extending overall survival.2,4 This radionuclide is administered as a chloride salt in aqueous citrate buffer without a biological
targeting vector or chelating agent. As a bone seeker, [\(^{223}\text{Ra}]\text{Ra}^{2+}\) is readily incorporated at sites of bone turnover,\(^1\)–\(^7\) including sites of osseous metastases, where it subsequently decays to irradiate the surrounding malignant tissue. Through its decay to stable \(^{207}\text{Pb}\), the 4 high-energy, short-range sites of osseous metastases, where it subsequently decays to 

PSMA-617, the limited availability of \(^{225}\text{Ac}\) and the challenges applied universally. need to be overcome before this therapeutic isotope can be 

in scaling up its production remain signiﬁcant hurdles that 

ions.

substantially weaker than those of the smaller alkaline earth and thus possesses a low charge-to-ionic radius ratio that 

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effectively use \(^{223}\text{Ra}\) for this purpose, however, this radiometal must be stably conjugated to tumor-targeting vectors via 

a bifunctional chelating agent (BFC). An effective BFC must match the coordination chemistry of the radiometal of interest 

such that the ion remains stably bound to the targeting vector in vivo.\(^{16,18}\) Despite decades of interest in Ra\(^{2+}\) for biomedical and environmental applications, no effective chelator for \(^{223}\text{Ra}\) has been identified that is suitable for biological applications.\(^{19}–\(^21\)

The challenge in ﬁnding an efﬁcient chelator for the stable chelation of this ion arises from its chemical properties. As an s-block ion, its interactions with ligands are predominantly electrostatic. Furthermore, Ra\(^{2+}\) is the largest +2 ion in the periodic table (eight-coordinate ionic radius = 1.48 Å (ref. 22)), and thus possesses a low charge-to-ionic radius ratio that results in electrostatic metal–ligand interactions that are substantially weaker than those of the smaller alkaline earth ions.\(^{23}\) Previous efforts to investigate the aqueous coordination chemistry of Ra\(^{2+}\) have focused on linear polyaminocarboxylate ligands, such as ethylenediamine(tetraacetic) acid (EDTA).\(^{24}–\(^28\)

These studies, which employed either long-lived \(^{226}\text{Ra}\) or tracer \(^{228}\text{Ra}\), demonstrate that complexes of Ra\(^{2+}\) with these ligands form, albeit with lower stability constants than the lighter alkaline earth ions. Competition extraction studies have also shown that macrocycles such as DOTA and Kryptofix 2.2.2 can bind to \(^{223}\text{Ra}\), although the biological stability of these complexes has not been evaluated.\(^{29}\) A further body of work in this field has been directed towards the development of organo-soluble extractants that partition this ion to the organic phase in biphasic systems.\(^{30}–\(^34\) Of these extractants, the most common structural motif explored has been that containing a calixarene core. Although calixarene-based ligands have demonstrated efﬁciency for the selective extraction of Ra\(^{2+}\) in the context of environmental remediation,\(^{32}–\(^35\) attempts to use these ligands for Ra\(^{2+}\) complexation for nuclear medicine applications have been unsuccessful due to either failed radiolabeling\(^{36}\) or poor kinetic stability of the resulting complex.\(^{39}\) These unsuccessful efforts have led many in the radiopharmaceutical community to the conclusion that stable chelation of \(^{223}\text{Ra}\) sufﬁcient for its in vivo use cannot be achieved. In turn, signiﬁcant work has been directed towards the incorporation of this radionuclide into various nanoparticle constructs.\(^{39}–\(^40\) Doping of the Ra\(^{2+}\) ion into solid-state nanoparticles, such as those of BaSO\(_4\), LaPO\(_4\), Fe\(_2\)O\(_3\), TiO\(_2\), hydroxyapatite, and nanozelites, has been shown to be an effective means of stabilizing this radionuclide and altering its biodistribution properties.\(^{41}–\(^46\) However, the complementary use of molecularly-targeted constructs using a BFC would enable more possibilities for radiopharmaceutical optimization via appropriate chemical modiﬁcations.

In this work, we show that macropa, an 18-membered bis-picolinate diazacycrown macrocyle (Fig. 1a), is an effective chelator of \([^{223}\text{Ra}]\text{Ra}^{2+}\), demonstrating rapid complexation kinetics and profound in vivo stability. This ligand is the ﬁrst chelator, to the best of our knowledge, that can stabilize this large ion in vivo. Further, we investigated Ra\(^{2+}\) chelation utilizing a bifunctional derivative of macropa conjugated to a single amino acid, β-alanine, or a prostate cancer-targeting agent, DUPA (Fig. 1b), to probe the effects of chelator...
functionalization on Ra$^{2+}$-ligand complex stability. Ultimately, this work garners a better understanding of the elusive coordination chemistry of the Ra$^{2+}$ ion and opens the path for the generation of targeted $^{223}$Ra therapeutics for novel biomedical applications.

Results and discussion

$^{223}$Ra labeling and in vitro characterization

Macropa is an expanded 18-membered macrocyclic ligand that contains two pendant picolinate donor arms appended to a diazao-18-crown-6 core (Fig. 1a).46 We and others have previously shown that ligands based on this macrocycle exhibit an unusual preference for binding to large over small ions.47–50 This property has made macropa useful for chelating the large radiometals $^{132/135}$La and $^{225}$Ac for therapeutic nuclear medicine applications.7–9,46 Furthermore, we have shown that this ligand forms a complex of high thermodynamic stability with Ba$^{2+}$, the largest non-radioactive +2 ion (eight-coordinate ionic radius (ref. 22) = 1.42 Å),51 prompting its recent investigation as a chelator for the diagnostic isotope $^{131}$Ba.52 In aqueous solution at pH 7.4, the conditional stability constant ($K_{ML}^{C_{\text{II}}}$) of the [Ba(macropa)] complex is 10.74 ($I = 0.1$ M KCl, 25 °C), indicating that macropa is the highest-affinity chelator for Ba$^{2+}$ at physiological pH reported to date. Based on these properties, we hypothesized that macropa might also be suitable for binding to Ra$^{2+}$, the heavier alkaline earth ion congener of Ba$^{2+}$, for TAT applications (Fig. 1a). The ionic radius of Ra$^{2+}$ is only marginally larger than that of Ba$^{2+}$, and as members of the same group in the periodic table, their donor atom preferences are expected to be similar. Density functional theory calculations (Fig. S1 and Tables S1–S3, ESI†) support this assertion, as both Ba$^{2+}$ and Ra$^{2+}$ are computed to form structurally analogous complexes with macropa.

To probe the suitability of macropa for the stable chelation of Ra$^{2+}$, we first sought to establish conditions for radiolabeling macropa with $^{223}$Ra.[Ra$^{2+}$]. The reaction was carried out at room temperature by mixing a solution of macropa with 3.7 kBq (0.1 μCi) of $^{223}$Ra[Ra$^{2+}$], obtained from an $^{222}$Ac/$^{222}$Th generator (Section 1.2 of the ESI†). The complexation was conducted at room temperature and buffered in metal-free ammonium acetate (0.1 M) at pH 6. Aliquots from these radiolabeling reactions were removed at different time points and analyzed by radio-instant thin layer chromatography (radioTLC) to assess complex formation. Under these TLC conditions, which used diglycolamide (DGA)-impregnated silica as the solid support and 0.1 M NaOH as the mobile phase, free $^{223}$Ra[Ra$^{2+}$] remained at the baseline ($R_t = 0$), whereas the complexed species migrated with the solvent front ($R_t = 1$) (Fig. 2a). The TLC strips were visualized using a phosphorimager after radioactive equilibration was reached to quantify the extent of $^{223}$Ra complexation. Within 5 min at room temperature, $^{223}$Ra[Macropa] was formed with a radiolabeling efficiency (RL%) of 95 ± 1.05% ($n = 5$) (Fig. 2a). These mild radiolabeling conditions are highly favorable for use with temperature-sensitive biological targeting vectors like antibodies, which degrade at elevated temperatures. By contrast, the most widely used chelator in nuclear medicine, $^{1,4,7,10}$-tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA, Fig. 1b), fails to completely coordinate $^{223}$Ra[Ra$^{2+}$] under similar conditions, as does the common polyaminocarboxylate ligand EDTA (see Section 1.3 and Fig. S2 in the ESI†). These results indicate that macropa provides an effective coordination environment for Ra$^{2+}$.

Next, the concentration of macropa was varied over a range of 187 nM to 3.74 mM with a fixed amount of $^{223}$Ra activity (3.7 kBq) to determine the maximum apparent molar activity that we could obtain with this system. Above a ligand concentration of 18 μM, the RL%’s were greater than 80% with no change over the 1 h duration (Fig. 2b). At a ligand concentration of less than 18 μM, however, the RL% dropped significantly to values of <20%. A plot of the RL% versus ligand concentration as determined at 5 min is shown in Fig. 2c. Interpolation of these data indicate that 50% RL% is achieved with a ligand concentration of 13 μM. Radiolabeling data obtained at the 1 h time point also gave the same ligand concentration required for 50% RL% (Fig. S3†), further showing that complexation efficiencies remain constant after the initial 5 min. Collectively, these data show that $^{223}$Ra[Macropa] is rapidly and quantitatively formed with molar activities ranging from 0.26 to 55.5 Ci mol$^{-1}$ or 2.05 to 9.62 × 10$^9$ MBq mol$^{-1}$, utilizing very low activity amounts (<5 μCi or <185 kBq) with radiochemical purity >95%. Although these molar activities are generally lower than those obtained for $^{225}$Ac-DOTA constructs,58,59 the high purity of $^{223}$Ra[Macropa] precludes the needs for additional chromatography of this radiocomplex prior to in vivo administration.

Having established that macropa can rapidly bind $^{223}$Ra under mild conditions, we next sought to determine the stability of the complex in conditions that are encountered in vivo using human serum at 37 °C over 12 days. Although radioTLC proved useful for assessing RL% in aqueous buffer, this method (silica or DGA-coated TLCs) failed when analyzing $^{223}$Ra speciation in the complex media of serum. Specifically, TLC plates analyzed from serum gave rise to multiple peaks and streaking, which prevented unambiguous assignment of free versus complexed $^{223}$Ra[Macropa] (Fig. S4†). To overcome this limitation, we turned to size exclusion chromatography (SEC) using dual detection of absorbance (at 280 nm UV) and gamma counting of collected fractions at radioactive equilibrium, a method that we have validated to be sufficient for unambiguously distinguishing between free and complexed $^{223}$Ra[Macropa] (Section 1.4 and Fig. S5 in ESI†). Upon incubation of $^{223}$Ra RaCl$_2$ in human serum at 37 °C (Fig. 3a), the UV chromatogram of this mixture shows several peaks below 20 mL, corresponding to serum proteins. The radiochromatogram shows only a broad peak at 30 mL, which is indicative of free Ra$^{2+}$. Importantly, no activity is observed to coincide with the protein fractions, indicating that free Ra$^{2+}$ does not strongly interact with other components in serum. Having validated this method for evaluating serum stability, we incubated $^{223}$Ra[Macropa] in human serum at 37 °C and analyzed aliquots of this mixture by SEC. A representative chromatogram of this mixture after 2 h is shown in Fig. 3b. At this time point, the radiochromatogram
shows only a single peak at 20 mL, which matches that of [Ba(macropa)] and [223Ra][Ra(macropa)] (Fig. S5†). The absence of the broad peak near 30 mL confirms that the [223Ra][Ra2+] ion remains bound within the macrocycle over this time period and that the complex is stable. After 12 days in serum, the span of approximately one half-life of 223Ra, [223Ra][Ra(macropa)] remains approximately 90% intact (Fig. 3c). This high degree of stability was also found for the non-radioactive [Ba(macropa)] complex, which remained intact in the presence of hydroxyapatite, a major constituent of bone matrix that also binds these metals (Section 1.5 and Fig. S6 in ESI†). The robustness of [223Ra][Ra(macropa)] compares favorably with other a particle-emitting radiopharmaceuticals and their respective chelation platforms (e.g. 225Ac-DOTA-antibody).58

In vivo biodistribution of [223Ra][Ra(macropa)]

Based upon the promising in vitro stability of [223Ra][Ra(macropa)], we proceeded to evaluate the biodistribution of this complex. Previously, we have shown that [223Ra]RaCl2 distribution in mice closely resembles that observed in humans, with rapid blood clearance, localization to sites of active bone remodeling, and excretion predominantly occurring through the small bowel and kidneys.6 To evaluate the in vivo stability of [223Ra][Ra(macropa)], its biodistribution was compared to the FDA/EMA approved formulation of [223Ra]RaCl2 administered in citrate buffer. We utilized healthy, skeletally mature rodent models combined with quantitative gamma counting measurements of organ activity normalized to weight. Organs were excised at 15 min and 24 h post-injection (p.i.) (Fig. 4). Consistent with its bone-seeking properties, a significant amount of [223Ra][Ra2+] was detected in bones (>10% IA per g) for the control [223Ra]RaCl2 citrate-administered groups at 15 min and 24 h p.i. (Fig. 4a). By contrast, an order of magnitude decrease in bone uptake was observed for animals injected with [223Ra][Ra(macropa)]. The latter demonstrated a bone uptake of 1.6 ± 0.2% IA per g versus 22 ± 1% IA per g for the control group at 24 h (****p < 0.0001; Fig. 4b and c).

In addition to decreasing the bone uptake, macropa markedly altered the excretion profile of this radiometal. The majority of [223Ra][Ra(macropa)] is rapidly cleared from the blood (2.55 ± 0.85% IA per g at 15 min to 0.007 ± 0.006 at 24 h) through renal excretion. Gut and spleen uptake were reduced (1.67 ± 1.5 and 0.7 ± 0.14% IA per g, respectively) as compared...
to the $^{223}$RaCl$_2$ control (4.08 $\pm$ 1.9 and 11.6 $\pm$ 6.1% IA per g). Preventing splenic accumulation (previously noted as a site of uptake of $^{223}$RaCl$_2$ in the rodent$^6$) and gut accumulation are meaningful improvements as gastrointestinal distress is a commonly reported symptom from $^{223}$RaCl$_2$ treatment in patients.$^2,3$ At 15 min p.i., we measured a 300-fold higher activity in the urine, sampled from the bladder, in mice treated with $^{223}$Ra[Ra(macropa)] compared to those treated with $^{223}$RaCl$_2$ (Fig. S7a†). At 24 h, all other organs showed negligible counts (<0.37 Bq or 0.01% IA).

The lower kidney activity combined with increased bladder accumulation of $^{223}$Ra[Ra(macropa)] compared to $^{223}$RaCl$_2$ (Fig. S7a†) indicates that the complex is rapidly cleared, thus providing further support that macropa retains this radiometal in vivo. As a final confirmation of the in vivo stability of $^{223}$Ra[Ra(macropa)], we analyzed urine samples of mice treated with this compound by SEC. The resulting radiochromatogram shows that the excreted species has the same retention time (20 mL) as $^{223}$Ra[Ra(macropa)], with no evidence of free or protein-bound activity (Fig. S7b†). In conjunction with the biodistribution data, this chromatogram shows conclusively that the $^{223}$Ra[Ra(macropa)] complex is stable in vivo, marking the first observation of a stable Ra$^{2+}$ coordination complex in this setting.

**Radiolabeling and testing of macropa conjugates**

Following our demonstration of $^{223}$Ra[Ra(macropa)] in vivo stability, the next objective was to explore conjugated forms of this chelator. We have previously reported macropa-NCS, a BFC version of macropa containing an amine-reactive isothiocyanate group appended to one of the picolinate pendant arms.$^{51}$ Using this bifunctional ligand, we first prepared a simple amino acid conjugate, macropa-$\beta$-alanine (see Section 1.7 and Fig. S8–S14 in the ESI†). In this construct, $\beta$-alanine is linked to macropa via a thioura bond. This compound provides a straightforward...
starting point to evaluate the effects of the thiourea conjugation strategy on Ra²⁺ complex stability. As a first means of evaluating the effects of this group, we carried out pH-potentiometric titrations to determine both the pH values and metal stability constants of this ligand for comparison to unfunctionalized macropa. The results of these titrations are compiled in Tables 1, S4 and Fig. S15, S16.† The pH values of macropa-β-alanine are very similar to those previously reported for macropa (Table 1). As expected, two additional protonation constants for the deprotonation of the NH thiourea and β-alanine carboxylic acid are present. The similarity of the other pH values, however, suggests that the donor properties of macropa-β-alanine are largely unchanged from those of macropa. The stability constants of macropa-β-alanine for Ca²⁺, Sr²⁺, and Ba²⁺ confirm this assertion. These values are scarcely perturbed from those measured for macropa. For example, we measured log K₅ values for macropa-β-alanine to be 11.06(0.10), whereas this value is 11.11 for macropa. Based on these values, both [Ba(macropa)] and [Ba(macropa-β-alanine)] are completely complexed at physiological pH (Fig. S5a). Thus, these thermodynamic data show that the addition of the thiourea group has little effect on the binding affinity of macropa for the alkaline earth ions.

To further confirm that the thiourea group has no negative effects on alkaline earth coordination, the Ba²⁺ complex of macropa-β-alanine was synthesized (see Section 1.7 and Fig. S17–S19 in the ESI†). To evaluate its kinetic stability, this complex was challenged with hydroxyapatite, a model of the Ca²⁺ center is 11-coordinate with 10 donors that the thiourea and β-alanine moieties were heavily disordered in the [Ba(macropa-β-alanine)[DMSO]] crystal structure, but in both possible configurations these groups were positioned far from the direct coordination sphere of the Ba²⁺ ion. These structural data further support the potentiometric titrations by showing that thiourea conjugation has no significant effects on the Ba²⁺ coordination properties of this ligand, suggesting that this BFC will retain high affinity for the Ba²⁺ ion.

Table 1 Protonation constants of macropa and macropa-β-alanine and thermodynamic stability constants of their alkaline earth complexes determined by pH potentiometry (25 °C and I = 0.1 M KCl)

| Ligand | Log K₅ | Log K₆ | Log K₇ | Log K₈ | Log K₉ | Log K₁₀ |
|--------|--------|--------|--------|--------|--------|--------|
| Macropa-β-alanine | 11.25(0.11), 11.36(0.01) | 7.41⁴, 7.41⁵ | 6.90⁴, 6.85⁵ | 3.25, 3.32⁵ | 2.45, 2.36⁵ | 1.69⁵ |
| Macropa | 2.51(0.05) | 11.18(0.10) | 3.76⁴ | 4.54(0.03) | 3.67(0.03) | 9.58(0.13) |

Table 1 Protonation constants of macropa and macropa-β-alanine and thermodynamic stability constants of their alkaline earth complexes determined by pH potentiometry (25 °C and I = 0.1 M KCl)

Although this structure suffers from significant crystallographic disorder and was weakly diffracting, the overall atomic connectivity and coordination geometry of Ba²⁺ could be discerned. The Ba²⁺ center is 11-coordinate with 10 donors provided by macropa and an 11th afforded by a disordered DMSO solvent molecule that interpenetrates the macrocycle base. In general, this structure resembles closely that of the unfunctionalized [Ba(macropa)] complex, which we previously reported. The conformation of the macrocycle about the Ba²⁺ center is similar for both structures (Fig. 5c). The thiourea and β-alanine moieties are heavily disordered in the [Ba(macropa-β-alanine)[DMSO]] crystal structure, but in both possible configurations these groups were positioned far from the direct coordination sphere of the Ba²⁺ ion. These structural data further support the potentiometric titrations by showing that thiourea conjugation has no significant effects on the Ba²⁺ coordination properties of this ligand, suggesting that this BFC will retain high affinity for the Ba²⁺ ion.

To further probe the utility of this BFC for [²²⁳Ra]Ra²⁺ chelation, we evaluated its ability to bind this radiometal. Using conditions identical to those employed for macropa, RL%’s of >90% were obtained for macropa-β-alanine with [²²⁳Ra]Ra²⁺ (n = 5), and the resulting complex was characterized with SEC (Fig. S20a). We tested the in vitro stability of the radiolabeled conjugate in human serum by SEC. These studies revealed that >70% of the complex remained intact after 12 days (Fig. 6a and S20b). Although this stability is slightly lower than what we observed for unfunctionalized [²²⁰Ra][Ra(macropa)], these data show that this ion can be kinetically stabilized in different macropa-like ligands. In the case of [²²⁰Ra][Ra(macropa-β-alanine)], the SEC chromatogram revealed a small amount of activity (~5% of initial content) in the elution volume that is characteristic of proteins (10–15 mL). Because we had shown that free [²²⁰Ra]Ra²⁺ does not bind directly to proteins as determined by SEC, we hypothesize that the presence of activity in the protein fractions for [²²⁰Ra][Ra(macropa-β-alanine)] is due to outer sphere interactions of this intact complex with proteins and not due to an inherent instability of the Ra²⁺ coordination sphere.

To assess the in vivo stability of [²²⁰Ra][Ra(macropa-β-alanine)], its biodistribution was investigated in mice. Similarly to [²²⁰Ra][Ra(macropa)], major differences in the organ distribution were observed as compared to [²²⁰Ra]RaCl₂. Notably, at 24 h p.i., it showed very low bone uptake (2.69 ± 0.24% IA per g) in comparison to [²²⁰Ra]RaCl₂ (9.7 ± 1.66%, *p = 0.0027) (Fig. 6b). Furthermore, [²²⁰Ra][Ra(macropa-β-alanine)] displayed approximately 5-fold lower splenic uptake (0.95 ± 0.5% IA per g vs. 5.52 ± 2.3% IA per g) and ~7-fold lower kidney uptake compared to [²²⁰Ra]RaCl₂, results that can be explained by the faster and more favorable renal clearance of the β-alanine conjugate. At 24 h, significantly less radioactivity is present in all tissues for [²²⁰Ra][Ra(macropa-β-alanine)] in comparison to mice treated with [²²⁰Ra]RaCl₂, for which significant activity was detected in the bone, spleen, kidney and intestine. Thus, these data show that conjugates of macropa have the potential to be highly stable in vivo. Furthermore, these results suggest that with the choice of an appropriate biological targeting vector, targeted therapy employing [²²⁰Ra]Ra is feasible.
To probe the potential of TAT using $^{223}$Ra, macropa-NCS was conjugated to a glutamate–urea–glutamate (DUPA) targeting vector. This DUPA targeting vector binds with high affinity to the PSMA, which is overexpressed on prostate cancer adenocarcinoma cells. Macropa-DUPA was found to chelate Ba$^{2+}$ effectively (Fig. S21–S23†) and prevent this ion from adsorbing Fig. 5 Alkaline earth metal complexes of macropa-β-alanine. (a) Species distribution diagrams of macropa (left) and macropa-β-alanine (right) in the presence of Ba$^{2+}$ at [Ba$^{2+}$]$_{tot}$ = [L]$_{tot}$ = 1.0 mM, $\mu$ = 0.1 M KCl, and 25 °C. (b) X-ray crystal structure of [Ba(macropa-β-alanine)(DMSO)]. Ellipsoids are drawn at the 50% probability level. Nonacidic hydrogen atoms are omitted for clarity. A full discussion of the crystallographic disorder of the structure is provided in the ESI.† (c) Comparison of the immediate coordination sphere of the Ba$^{2+}$ center in the crystal structures of [Ba(Hmacropa)(DMF)]$^+$ and [Ba(macropa-β-alanine)(DMSO)].

Fig. 6 In vitro and in vivo evaluation of $^{223}$Ra[Ra(macropa-β-alanine)]. (a) Stability of $^{223}$Ra[Ra(macropa-β-alanine)] over the course of 12 days in human serum at 37 °C. (b) Organ distribution of $^{223}$Ra[Ra(macropa-β-alanine)] at 24 h p.i. Significant differences in osseous (⋆p < 0.005), splenic, and renal uptake were observed in comparison to the control $^{223}$RaRaCl$_2$. © 2021 The Author(s). Published by the Royal Society of Chemistry.
onto hydroxyapatite, albeit to a slightly lower extent than macro-
ropa-β-alanine (Fig. S6†). Having confirmed its ability to
complex Ba²⁺, radiolabeling with [²²³Ra][Ra²⁺] was carried out.
Incorporation of this radiometal proceeded efficiently, giving
RL% of >95% (Fig. S24a†). Furthermore, the radiolabeled
conjugate remained >90% intact upon incubation in serum at
37 °C for 12 days (Fig. S24b†).

Based on these promising in vitro data, we evaluated the
biodistribution of this conjugate in mice. Unexpectedly, the
macropa-DUPA conjugate exhibited no difference in bi-
distribution in comparison to free [²²³Ra][RaCl₂] (Fig. S25†).
Notably, high levels of bone uptake consistent with the
release of free [²²³Ra]Ra²⁺ were detected. Thus, despite the
promising stability of macropa and macropa-β-alanine with
²²³Ra, this stability is lost upon attachment to the DUPA
PSMA-targeting vector. This observation contrasts with our
previous report of stable chelation with radioactive
lanthanum utilizing macropa-DUPA,⁴† highlighting the
unique requirements for successful chelation of radium.
Although the reasons for the lack of stability in the DUPA
conjugate, in comparison to the β-alanine conjugate, are not
fully understood, these data show that targeting vectors can
affect metal-chelate stabilities. This phenomenon warrants
further investigation, as it may have significant implications
in the development of new metal-based radiopharmaceutical
agents.

Conclusions

We have demonstrated the robust, rapid, and stable chelation
of radium in aqueous solution using the macrocyclic ligand
macropa. Upon complexation at room temperature, the
[²²³Ra[macropa]] complex displays a high degree of long-term
stability both in vitro under physiological conditions and in vivo
in murine models. Although our efforts to target ²²³Ra to
soft-tissue prostate cancers using the PSMA-binding ligand
DUPA were unsuccessful, the in vitro and in vivo success of
both [²²³Ra][Ra(macropa)] and [²²³Ra][Ra(macropa-β-alanine)]
opens the path forward for possible cancer-targeted z-particle
radiotherapy with ²²³Ra. Considering that within the radio-
pharmaceutical community, targeted forms of ²²³Ra were
considered inaccessible due to the lack of biologically stable
chemator complexes, our demonstration that macropa and
macropa-β-alanine can retain this ion in vivo will motivate
continued efforts towards applying this radionuclide for new
applications in TAT. We are currently focusing on efforts to
develop TAT constructs that are sufficiently stable in vivo for
such applications.

Conflicts of interest

Wilson and Thiele hold intellectual property rights on macropa;
Abou and Thorek have filed provisional patent protection for
radium-related production and utilization through WUSTL.
Justin Wilson holds equity in Noria Therapeutics, Inc., which
has licensed this technology.

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