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Design of porous self-assembled homoleptic and heteroleptic Pd²⁺ cages incorporating siliconbased fluoride acceptors: the way towards nuclear imaging applications†

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Supramolecular self-assembly is a promising tool to develop multifunctional theranostic agents and has recently entered the field of radiochemistry. In this work, lantern-shaped [Pd₂L₄]⁴⁺ metallacages featuring a blood-brain barrier-penetrating peptide were designed for dual-modality imaging. Two silicon-based fluoride acceptors were incorporated in the cage ligand for radiolabeling with fluorine-18, an isotope used in positron emission tomography (PET). Ligands (L1, L2) and the respective homoleptic cages ($C1_{hom}$ $[Pd_2(\mathbf{L1})_4]^{4+}$, $\mathbf{C2_{hom}}$ $[Pd_2(\mathbf{L2})_4]^{4+}$) were fully characterized by NMR spectroscopy, liquid chromatography and high-resolution electrospray mass spectrometry (HR-ESI-MS). Cage stability was assessed in different solvents and concentrations by HPLC. ¹⁸F-labelled cages were obtained by radiolabeling the ligands preassembly under mild conditions within four hours via [19F]-to-[18F] isotopic exchange. The high lipophilicity of the ligands was also assessed in vitro ($log D_{pH7.4}$) and in ovo, using the chick chorioallantoic membrane (CAM) model. Furthermore, formation of host-guest complexes between the metallacages and perrhenate (ReO₄⁻), the 'cold' surrogate of radioactive ^{99m}TcO₄⁻ (used for single photon computed tomography, SPECT), could be detected by mass spectrometry, although the adduct did not sustain chromatographic separation. To improve stability of the supramolecular system, heteroleptic cages of general formula $[Pd_2(L)_m(\mathbf{L0})_n]^{4+}$ (m = 1, 2, n = 4 - m) were synthesized by statistical self-assembly and separated by liquid chromatography in good yield. Overall, this study demonstrates the feasible adaptation of supramolecular principles to achieve innovative theranostic agents.

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Introduction

In the last few decades, supramolecular coordination complexes (SCCs) have been explored in different fields of chemistry due to the broad span of possible applications, including in medicine.^{1–3} SCCs are discrete molecular arrangements composed by metal nodes bound to organic ligands that come

together via coordination-driven self-assembly (CDSA).4,5 A plethora of two-dimensional and three-dimensional shapes can be created upon thoughtful combination of the aforementioned building blocks, which morphologically are classified broadly into three main types, namely metallacycles, metallacages and helicates. 6,7 The coordination geometry of the metal ion, ligand curvature, size and number of coordination sites encode the self-assembly of the proposed architecture.3,8 In this context, the case of the metallacages (MCgs) is particularly appealing since their porous nature favors their use to encapsulate and transport small molecules.9 Moreover, exo- and endo-functionalization of the ligands pre- and post-CDSA allows for further tuning of the supramolecular system. The host-guest chemistry and versatility of MCgs towards functionalization led to their investigation as catalysts, 10,11 sensors^{12,13} and gas storage systems¹⁴ amongst other possible uses. 15 Concerning biomedical applications, MCgs have been initially explored as potential carriers of cargos of pharmaceutical interest¹⁶⁻²⁰ either for therapy or imaging (Fig. 1A), or

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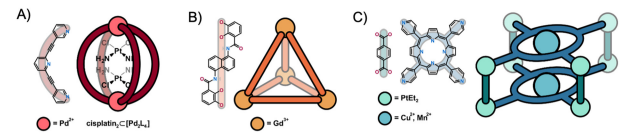


Fig. 1 Literature examples of MCgs designed for biomedical applications. (A) Self-assembled lantern shaped- $[Pd_2L_2]^{4+}$ MCqs encapsulating the anticancer drug cisplatin; (B) tetrahedral Gd(III) MCgs used for MRI; (C) porphyrin-based MCgs designed for trimodal imaging (MRI, PET or aggregationinduced emission) in vivo

for their intrinsic cytotoxic properties. 21-23 Some MCgs structures have also been incorporated into nanoparticles for multimodal imaging and to improve their stability and targeting capability.^{24,25} Besides the cytotoxic potential of SCCs, their design can be directed to incorporate other interesting therapeutic modalities, such as photodynamic therapy via generation of reactive oxygen species at the target site.26-28 Recently, an Au(III)-porphyrin-based Rug-cage was reported for G-quadruplex-DNA-targeted photodynamic therapy, which displays higher cytotoxicity upon irradiation compared to the free porphyrin.²⁹ Exo-functionalization of the cage ligand is a suitable strategy for the conjugation of targeting vectors towards cancer cells. As an example, $[Pd_2L_4]^{4+}$ metallacages with ligands bioconjugated to integrin-binding RGD-peptides were developed.30 Similarly, MCgs can also be designed to display luminescent properties useful to study their fate in biological systems by optical imaging methods.31-34

Concerning clinical imaging applications, self-assembled Gd(III)/Fe(III) tetrahedral cages (Fig. 1B), originally designed by Raymond et al.,35 were investigated by Morrow and coworkers as T1 MRI probes in vivo, 36-39 showing resistance to dissociation in biological media. In 2018, the group of Lusby and Archibald made use of the encapsulation properties of SCCs and designed a Co(III)4L6 rhomboidal MCg for the incorporation of 99mTcO₄-, a γ-emitter that is commonly used for single photon emission computed tomography (SPECT) imaging in the clinic.40 While free 99m-pertechnetate exhibited high thyroid uptake in healthy mice, encapsulation of the oxo-anion resulted in a distinct biodistribution profile, showcasing the utility of drug delivery properties also for radionuclides. Representative examples of MCgs for medicinal applications are included in Fig. 1 and the reader is directed to more comprehensive reviews for further insights. 2,41,42

Amongst the most intriguing aspects of MCgs is the possibility to design them as multifunctional systems for targeted concomitant diagnostic and therapeutic (theranostic) applications. 43 For example, initial work by Stang, Chen, Cook and coworkers reported on different metallacage systems for therapy and imaging via different approaches, including nuclear imaging modalities or aggregation-induced emission (Fig. 1C). 27,44 In this context, the case of "lantern-shaped" cationic [Pd₂L₄]⁴⁺ cages (L = bitopic monodentate N-donor ligand)

is noticeable since already a few reports exist on their application in fluorescence microscopy and nuclear imaging (Table S1†). 32,33,45-50

In the case of radiolabeled MCgs, three strategies are applicable to integrate the radionuclide in the system: ligand functionalization, encapsulation via host-guest chemistry, and complexation of the radiometal as part of the supramolecular assembly. Thus, our group reported on $[Pd_2L_4]^{4+}$ cages whereby the ligand L was bioconjugated to a peptide (AGILKRW, PepH3) capable to cross the blood-brain barrier (BBB), and assembled into a [Pd₂L₄]⁴⁺ cage. 48 Upon encapsulation of 99mTcO₄-, the MCg was injected in mice in vivo, and brain accumulation of the host-guest complex was detected at 0.42% ID per g 5 min p.i., supporting cage application as brain delivery vehicle.

Previously, a [Pd₂L₄]⁴⁺ cage had already been modified via click chemistry with 18F-trifluoroborate moieties for positron emission tomography (PET) imaging. 49 Here, the intactness of the MCg in vivo was corroborated by its different biodistribution with respect to the free ligand by PET imaging. Interestingly, the zwitterionic nature of the trifluoroborate moiety rendered the cage sufficiently soluble in aqueous environment. More recently, (Tyr³)-octreotate, a somatostatinreceptor 2 addressing peptide for targeting neuroendocrine tumors was tethered via amide bond formation to the bispyridyl L ligand. 50 A DOTA chelator (1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid) was also inserted into the ligand sequence to bind Lu-177, a β^- and γ -emitting radionuclide used for cancer treatment and therapy control via SPECT. While the sterically demanding ligand only allowed the formation of a [Pd2L2]4+ homoleptic metallacycle, formation of a heteroleptic MCg was achieved by adding a non-functionalized less sterically hindering ligand.50

Thriving to implement self-assembled MCgs as new generation radiopharmaceuticals, we herein report on the synthesis by CDSA of homoleptic and heteroleptic Pd²⁺-cages featuring a 3,5-bis(3-ethynylpyridine)phenyl backbone and exo-functionalized with the BBB-targeting peptide PepH3 tethered to two different silicon-based fluoride acceptors (SiFAs) moieties for radiolabeling with fluorine-18 for PET imaging (Fig. 2A). PepH3 has already been studied for its BBB translocation capability in vitro and in vivo, 51 including

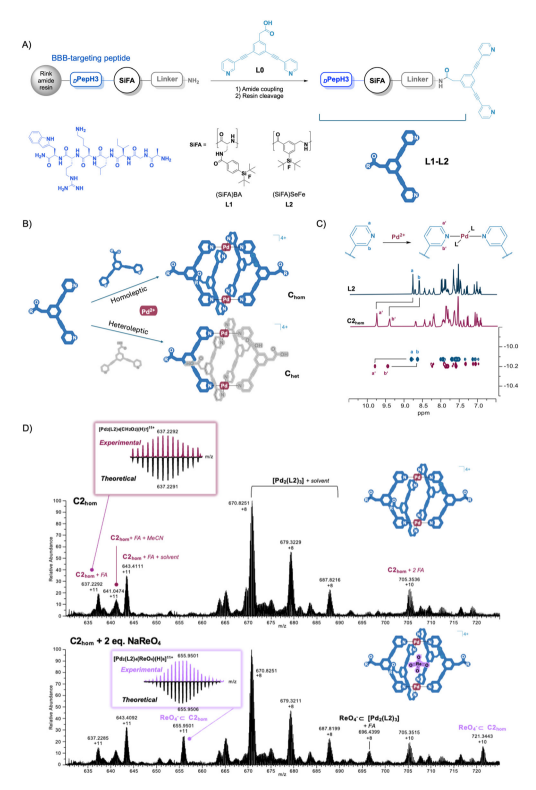


Fig. 2 (A) Design and synthesis of BBB-targeted ligands L1 and L2 by Fmoc-SPPS. The exo-functionalization consists of a BBB shuttle peptide PepH3, a silicon-based fluoride acceptor (SiFA) for radiolabeling with the PET nuclide fluoride-18 and a PEG linker (4-((2-(2-aminoethoxy) ethoxy)ethyl)amino)-4-oxobutanoic acid) that acts as a spacer. (B) Coordination-driven self-assembly of homoleptic and heteroleptic MCgs, C_{hom} and C_{het} . Reaction conditions: 4 eq. ligand (in total), 2 eq. [Pd(MeCN)₄](BF₄)₂, DMSO, RT, 1 h. (C) ¹H NMR (top) and ¹H-DOSY NMR (bottom) spectra in DMSO- d_6 (400 MHz) of C2_{hom} (purple) compared to L2 (blue). The α -pyridinyl protons a and b shift upon metal coordination as explained by the scheme. (D) Mass spectra of $C2_{hom}$ (top) and the host-guest complex $ReO_4^- \subset C2_{hom}$ (bottom) obtained by DI-HR-ESI-MS. Corresponding theoretical and experimental isotopic patterns are shown in the graph insets.

bioconjugated to MCgs structures. 48 Interestingly, the peptide has been reported to cross the BBB via an active transport mechanism, namely adsorptive-mediated transcytosis (AMT), due to the presence of positively charged amino acid residues in its sequence.⁵² Here, the physiologically more stable *D*-enantiomer of PepH3 (*p*PepH3) was applied.⁵³ The suitability of this design strategy was challenged by exploring the fundamentals of homoleptic and heteroleptic CDSA (Fig. 2B). The lipophilic character of the ligands $(\log D_{\rm pH7.4})$ was determined, and in one case, also investigated in ovo in glioblastoma U87 tumors engrafted on the chicken chorioallantoic membrane (CAM) model by PET. Further, the obtained homoleptic MCgs were characterized by different methods, including ¹H NMR spectroscopy and high-resolution electrospray ionization mass spectrometry (HR-ESI-MS). In addition, the encapsulation properties of the homoleptic systems were investigated by using the guest molecule perrhenate as a cold surrogate for 99mTcO₄-. Further, a protocol for the synthesis of ¹⁸F-labeled MCgs via pre-assembly labeling was optimized, which held radiochemically pure MCgs. In parallel, the synthesis of heteroleptic MCgs, of general formula $[Pd_2(L)_m(L0)_n]^{4+}$ (m = 1, 2, n = 4 - 1)m) was achieved by statistical self-assembly of two different types of ligands,⁵⁴ followed by chromatographic separation.

Experimental

Research Article

General information

All reagents and solvents were purchased commercially and were used without further purification. The synthesis of SiFAs building block was achieved as previously reported, 55,56 and their purity >95% were determined by reverse-phase high performance liquid chromatography (RP-HPLC). Ligand L0 (2-(3,5-bis(pyridin-3-ylethynyl)phenyl)acetic acid) was synthesized according to literature-known procedure.⁵⁷ The product was obtained as a white solid in a total yield of 66% and 97% purity by RP-HPLC.

Fluoride-18 in target water ([18O]H2O) was obtained from Klinikum rechts der Isar (Munich, Germany) and Guys and St Thomas' Hospital (London, UK).

¹H NMR and ¹H-DOSY NMR spectra were recorded on a Bruker Avance III HD 400 NMR instrument by Bruker Cooperations (Billerica, USA). Chemical shifts are given in parts per million (ppm) and are referenced to the residual proton signal of the solvent. The coupling constants J are reported in hertz (Hz) and the signal splitting patterns are expressed as singlet (s), doublet (d), doublet of doublets (dd), triplet (t), doublet of triplets (dt), doublet of quartets (dq) and multiplet (m).

Analytical and semi-preparative RP-HPLC measurements were performed using gradient systems by Shimadzu Corp. (Kyoto, Japan), consisting of a SPD-20A dual wavelength UV/Vis detector (l = 220 nm, l = 254 nm), two LC-20AD gradient pumps, a CBM-20A communication unit and a CTO-20A column oven. For analytical RP-HPLC measurements a flow

rate of 1 mL min⁻¹ and for semi-preparative RP-HPLC measurements a flow rate of 10 mL min⁻¹ was used. Quality controls of the ligands and analysis of cage formation reactions were performed using a MultoKrom® 100-5 C-18 column (150 × 4.6 mm, 5 µm particle size) from CS-Chromatographie Service GmbH (Langerwehe, Germany). For analytical separation of ligands and cages, different gradients of H2O (0.1% trifluoroacetic acid (TFA)) (A) and MeCN (2% H₂O, 0.1% TFA) (B) were used during RP-HPLC measurements. For semi-preparative purification of ligands, different gradients of H2O (0.1% TFA) (A) and MeCN (5% H₂O, 0.1% TFA) (B) were used.

Quality controls of radiolabeled products was conducted using a gradient system by Shimadzu Corp. (Kyoto, Japan), consisting of an SPD-20A dual wavelength UV/Vis detector (l =220 nm, l = 254 nm), two LC-20AD gradient pumps, a HERM LB500 (NaI scintillation crystal) as well as a FlowStar² LB514 radio detector from Berthold Technologies GmbH (Bad Wilbad, Germany) and a CBM-20A communication unit. Radiolabeled ligands were analysed using a MultoKrom® 100-5 C-18 column (125 \times 4 mm, 5 μ m particle size) with a constant flow rate of 1 mL min⁻¹. Analysis of the radiolabeled cages was performed using a MultoHigh® Bio 300-5 C4 column (150 × 4.6 mm, 5 μm particle size) at a flow rate of 1 mL min⁻¹. Radio thin-layer chromatography (TLC) was performed on a ScanRAM radio TLC detector by LabLogic (Sheffield, UK) and analysed with the Laura software.

Direct-infusion (DI) and liquid chromatography (LC) highelectrospray-ionization spectrometry resolution mass (HR-ESI-MS) was applied for the analysis of ligands and MCgs using an Orbitrap Q-Exactive Plus Mass Spectrometer from Thermo Fisher Scientific. For LC-MS, the MS system was coupled to Dionex ultimate 3000 series UHPLC from Thermo Fisher Scientific for LC-MS. The column used for the C2hom analysis was a SUPELCO BIOshell A400 Protein C4 (3.4 µm, 400 Å, 15 cm \times 1 mm) from Sigma-Aldrich. The mobile phases were A: H₂O and B: MeCN, both with 0.1% formic acid (FA). The flow rate was 0.05 mL min⁻¹ and sample elution was performed by using a gradient from 20% to 40% of B over 3 min, followed by a gradient from 40% to 80% of B over 15 min. Sample injection volume was 10 µL. For the C1_{het} analysis the column used was an AcclaimTM 120: C18 (5 µm, 120 Å, 4.6 mm × 100 mm), Dionex Bonded Silica Products from Thermo Fisher Scientific. The mobile phases were A: H₂O and B: ACN, both with 0.1% formic acid (FA). The flow rate was 0.5 mL min⁻¹ and sample elution was performed by using a gradient from 20% to 40% of B over 3 min, followed by a gradient from 40% to 80% of B over 15 min. Ionization was performed using an ESI source operating in positive ion mode with a capillary voltage of 3.50 kV and capillary temperature 320 °C. Sheath gas, auxiliary gas and sweep gas flow rate were set at 20, 12 and 1 (arbitrary units), respectively. Auxiliary gas temperature was set at 100 °C.

For DI, the sample was infused at 0.01 mL min⁻¹ and ionization was performed using an ESI source operating in positive ion mode with a capillary voltage of 3.50 kV and capillary temperature of 300 °C. Sheath gas, auxiliary gas and sweep gas flow rate were set at 20, 12 and 0 (arbitrary units), respectively. Auxiliary gas temperature was set at 50 $^{\circ}$ C.

Ligand synthesis

Synthesis of the BBB-shuttle peptide $_D$ PepH3. $_D$ PepH3 (AGILKRW) was synthesized via microwave-assisted solid phase peptide synthesis on a Liberty BlueTM instrument manufactured by CEM (Matthews, USA). ProTide Rink Amide resin (loading capacity = 0.65 mmol g⁻¹) in combination with Fmocprotected amino acids were used for the synthesis. No final Fmoc-deprotection was performed and the peptide was stored on the resin until further use.

Synthesis of pPepH3-pDap((SiFA)BA)-Ebes-L0 (L1)

Fmoc-deprotection. The N-terminal Fmoc-protected $_D$ PepH3, immobilized on resin, (60.0 µmol, 1.0 eq.) was deprotected by addition of 20% piperidine in dimethyl formamide (DMF, 5 mL) at RT. The deprotection step was carried out twice (1 × 15 min, 1 × 5 min) and afterwards the resin was washed with DMF (6 × 5 mL).

Fmoc-Dap(Dde)-OH (N-alpha-(9-fluorenylmethyloxycarbonyl)-N-beta-[(4,4-dimethyl-2,6-dioxocyclohex-1-ylidene) ethyl]-D-2,3-diaminopropionic acid) (90.0 µmol, 1.5 eq.), TBTU (N,N,N',N'-tetramethyluronium-tetrafluorborate) (90.0 µmol, 1.5 eq.), HOAt (1-hydroxy-7-azabenzotriazole) (90.0 µmol, 1.5 eq.) and 2,4,6-trimethylpyridine (300 µmol, 5.0 eq.) were dissolved in 5 mL DMF. After 2 min preincubation, the solution was added to the deprotected resin-bound peptide and shaken for 2 h at RT. Afterwards the resin was washed with DMF (6 \times 5 mL) and N-methyl-2-pyrrolidone (NMP, 6×5 mL). For a coupling in branched position the Dde-protection group was cleaved by adding a solution of NH2OH·HCl (6.0 mmol, 100 eq.) and imidazole (4.5 mmol, 75 eq.) in 7 mL NMP/DCM (5/2). After shaking for 2 h at RT, the resin was washed with NMP (6 \times 5 mL) and DMF (6 \times 5 mL).

Amide coupling. Fmoc-(SiFA)BA (72.0 μ mol, 1.2 eq.), TBTU (90.0 μ mol, 1.5 eq.) and HOAt (90.0 μ mol, 1.5 eq.) were dissolved in 5 mL DMF. After the addition of DIPEA (*N*,*N*-diisopropylethylamine) (210 μ mol, 3.5 eq.) the solution was preincubated for 30 min. The coupling solution was added to the deprotected resin-bound compound and the reaction mixture was shaken at RT for 2.5 h. Afterwards the resin was washed with DMF (6 × 5 mL).

Fmoc-Ebes (N-[8-(9-fluorenylmethyloxycarbonyl)amino-3,6-dioxaoctyl]succinamic acid) (90.0 μ mol, 1.5 eq.) and **L0** (90.0 μ mol, 1.5 eq.) were coupled consecutively to the resinbound compound by first performing Fmoc-deprotection and then amide coupling.

Resin cleavage. The ligand was cleaved off the resin by shaking with 2 \times 5 mL of a mixture of TFA/triisopropyl silane (TIPS)/H₂O (95/2.5/2.5) for 2 \times 1 h at RT. The ligand solution was collected in a round-bottom flask and stirred for 1 h before being concentrated in an N₂ stream. Then, 5 mL of TFA was added and the mixture was stirred at RT for another 2 h. Afterwards, the TFA was removed again in an N₂ stream and the ligand was purified via semi-preparative RP-HPLC and lyophilized.

 $_D$ PepH3- $_D$ Dap((SiFA)BA)-Ebes-**L0** was obtained as a white solid (11.5 mg, 11% yield, 98% purity).

RP-HPLC (10–90% B): t_R = 9.7 min.

¹H NMR (400 MHz, DMSO- d_6) δ (ppm) = 10.78 (s), 8.78 (d, J = 3.1 Hz, 2H), 8.61 (d, J = 6.5 Hz, 2H), 8.49 (d, J = 4.4 Hz), 8.26 (d, J = 6.6 Hz), 8.21 (d, J = 5.9 Hz), 8.14–8.08 (m), 8.04–7.97 (m), 7.98–7.92 (m), 7.89–7.79 (m), 7.68 (s), 7.67–7.61 (m), 7.61–7.53 (m), 7.52–7.43 (m), 7.39 (s), 7.32 (d, J = 7.1 Hz), 7.11 (d, J = 2.6 Hz), 7.08–7.01 (m), 7.00–6.92 (m), 4.43 (dq, J = 13.6, 6.7 Hz), 4.33–3.92 (m), 3.81–3.58 (m), 3.53–3.46 (m), 3.43 (t, J = 5.8 Hz), 3.36 (t, J = 5.9 Hz), 3.20 (dt, J = 30.4, 5.7 Hz), 3.12–2.92 (m), 2.72 (d, J = 7.5 Hz), 2.36 (s), 1.85–1.36 (m), 1.25 (d, J = 7.4 Hz), 1.11 (s), 1.01 (d, J = 1.1 Hz), 0.85 (d, J = 6.4 Hz), 0.83–0.73 (m).

MS (ESI positive): calculated monoisotopic mass for $C_{90}H_{124}FN_{19}O_{14}Si: 1741.93$; found by ESI-MS: $m/z = 582 [M + 3H]^{3+}, 872 [M + 2H]^{2+}$.

Synthesis of $_D$ PepH3-(SiFA)SeFe-Ebes-L0 (L2). Fmoc-SiFA (SeFe) (72.0 μ mol, 1.2 eq.), Fmoc-Ebes (90.0 μ mol, 1.5 eq.) and L0 (90.0 μ mol, 1.5 eq.) were coupled consecutively to the resinbound peptide by first performing Fmoc-deprotection and then amide coupling as described above. A resin cleavage was carried out to obtain $_D$ PepH3-(SiFA)SeFe-Ebes-L0 as a white solid (17.6 mg, 17% yield, 98% purity) as described for the synthesis of L1.

RP-HPLC (10–90% B): t_R = 9.5 min.

¹H NMR (400 MHz, DMSO- d_6) δ (ppm) = 10.79 (s), 8.79 (s, 2H), 8.73 (d, J = 6.5 Hz), 8.61 (dd, J = 4.9, 1.7 Hz, 2H), 8.52–8.43 (m), 8.39–8.30 (m), 8.28–8.17 (m), 8.00 (dt, J = 7.9, 1.8 Hz), 7.98–7.78 (m), 7.75–7.62 (m), 7.61–7.53 (m), 7.53–7.45 (m), 7.39 (s), 7.32 (d, J = 7.9 Hz), 7.12 (d, J = 2.4 Hz), 7.09–7.01 (m), 7.00–6.92 (m), 4.44 (dt, J = 19.4, 6.9 Hz), 4.33 (d, J = 5.9 Hz), 4.30–3.93 (m), 3.74 (d, J = 5.8 Hz), 3.50 (dd, J = 4.3, 2.6 Hz), 3.43 (t, J = 5.7 Hz), 3.37 (t, J = 5.9 Hz), 3.29–3.14 (m), 3.14–2.94 (m), 2.73 (d, J = 7.5 Hz), 2.46–2.29 (m), 2.24 (d, J = 0.6 Hz), 2.08 (s), 1.97–1.90 (m), 1.71–1.58 (m), 1.56–1.42 (m), 1.37 (d, J = 7.1 Hz), 1.23 (s), 1.11 (s), 1.03–0.97 (m), 0.88–0.70 (m).

MS (ESI positive): calculated monoisotopic mass for $C_{88}H_{121}FN_{18}O_{13}Si: 1684.91$; found by ESI-MS: $m/z = 562 \text{ [M + 3H]}^{3+}, 844 \text{ [M + 2H]}^{2+}, 1686 \text{ [M + H]}^{+}.$

Synthesis of metallacages by CDSA and RP-HPLC analysis

To achieve homoleptic MCgs assembly, ligand L1 or L2 (2.0 eq., 5 mM in DMSO) and $[Pd(MeCN)_4](BF_4)_2$ (1.0 eq., 20 mM in DMSO) were mixed in DMSO to reach a theoretical cage concentration of 1 mM. Instead, CDSA of the heteroleptic cages was achieved by either mixing 1.0 eq. of L1 (5 mM in DMSO) and 3.0 eq. of L0 (5 mM in DMSO) for C1_{het}, or 2.7 eq. of L2 (5 mM in DMSO) and 1.3 eq. of L0 (5 mM in DMSO) for C2_{het}, with 2.0 eq. of $[Pd(MeCN)_4](BF_4)_2$ (20 mM in DMSO) and adding DMSO to reach a theoretical cage concentration of 1 mM

In all cases, the reaction was complete after 1 h at RT and analysed by analytical RP-HPLC. For the homoleptic cages a gradient of 40–70% B in 15 min was applied. In the case of the

heteroleptic systems, the reaction resulted in a mixture of heteroleptic coordination species with different ligand ratios. Therefore, two different gradients were tested to achieve the best separation: (i) 30–60% B over 15 min with a flow rate of 1 mL min⁻¹ and 0.1% TFA as additive, and (ii) 20%–40% B over 3 min, followed by a gradient 40%–80% over 15 min with a flow rate of 0.5 mL min⁻¹ and 0.1% FA as additive.

 $[Pd_2(L1)_4](BF_4)_4(C1_{hom})$

RP-HPLC (40–70% B in 15 min): t_R = 8.9 min. Yield 88.7%. [$Pd_2(L2)_4$](BF_4)₄ ($C2_{hom}$)

RP-HPLC (40-70% B in 15 min): t_R = 7.3 min. Yield 82.1%.

Stability studies of homoleptic cages

For stability studies, $C1_{hom}$ and $C2_{hom}$ (20 μ L, 1 mM) were added to 180 μ L of different solvents (DMSO, H_2O , MeCN, MeCN/ H_2O (1/1, +0.1% TFA)), phosphate buffered saline (PBS, pH 7.4) and saline solution (0.9 wt% NaCl), respectively, to achieve a cage concentration of 0.1 mM. The stability was determined by RP-HPLC after incubation at different times (0 min, 30 min, 60 min, 4 h, 24 h).

CDSA and encapsulation studies by ¹H and ¹H-DOSY NMR spectroscopy

The NMR samples were prepared using \sim 7 mg of ligand L1/L2 (2.0 eq.) dissolved in 400 μ L of DMSO- d_6 , and 1 H NMR as well as 1 H-DOSY NMR spectra were recorded. To the ligand solution, [Pd(MeCN)₄](BF₄)₂ (1.0 eq.) was added in 25 μ L DMSO- d_6 . 1 H NMR and 1 H-DOSY NMR spectra were recorded after 1 h at RT to observe homoleptic cage formation.

To monitor CDSA of the heteroleptic cages, \sim 2 mg of L1 (1.0 eq.) and 3.0 eq. of L0 for C1_{het}, or \sim 6 mg of L2 (2.7 eq.) and 1.3 eq. of L0 for C2_{het}, were dissolved in 400 μ L of DMSO- d_6 in an NMR tube. Thus, 1 H NMR as well as 1 H-DOSY NMR spectra were recorded. To the ligand solution was added 2.0 eq. of [Pd(MeCN)₄](BF₄)₂ in 25 μ L DMSO- d_6 and after 1 h at RT, 1 H NMR and 1 H-DOSY NMR spectra of the formed homoleptic cages C1_{het} and C2_{het} were measured. Note that a mixture of different heteroleptic coordination species results from the reaction.

To study the encapsulation properties of the homoleptic MCgs, the cages were prepared in DMSO- $d_6/\mathrm{D_2O}$ (1/1). 2.0 eq. of NaReO₄ in 25 μ L DMSO- $d_6/\mathrm{D_2O}$ (1/1) was added to the cage solution, and after 30 min at RT, ¹H NMR spectra were recorded.

 $[Pd_2(L1)_4](BF_4)_4$ (C1_{hom})

¹H NMR (400 MHz, DMSO- d_6) δ (ppm) = 10.78 (s), 9.77 (s, 2H), 9.41 (s, 2H), 8.52–8.44 (m), 8.27–8.17 (m), 8.13–8.04 (m), 8.04–7.91 (m), 7.90–7.75 (m), 7.73–7.60 (m), 7.58 (d, J = 4.8 Hz), 7.47 (t, J = 5.4 Hz), 7.40 (s), 7.32 (d, J = 8.1 Hz), 7.11 (s), 7.09–7.01 (m), 7.00–6.92 (m), 4.50–4.37 (m), 4.33–3.84 (m), 3.46 (s), 3.42–3.28 (m), 3.19–2.92 (m), 2.75–2.69 (m), 2.34 (s), 2.07 (s), 1.75 (s), 1.46 (dt, J = 20.2, 7.5 Hz), 1.25 (t, J = 7.1 Hz), 1.11 (s), 1.06–0.93 (m), 0.88–0.72 (m).

 $[Pd_2(L2)_4](BF_4)_4$ (C2_{hom})

¹H NMR (400 MHz, DMSO- d_6) δ (ppm) = 10.80 (s), 9.78 (s, 2H), 9.43 (s, 2H), 8.72 (d, J = 6.2 Hz), 8.48 (t, J = 5.8 Hz), 8.33

(d, J = 5.6 Hz), 8.24 (d, J = 7.8 Hz), 8.06–7.77 (m), 7.68 (d, J = 5.6 Hz), 7.58 (d, J = 7.6 Hz), 7.51 (d, J = 5.4 Hz), 7.42 (s), 7.32 (d, J = 8.1 Hz), 7.11 (d, J = 2.3 Hz), 7.10–7.01 (m), 7.01–6.92 (m), 4.51–4.38 (m), 4.32 (d, J = 5.7 Hz), 4.29–4.11 (m), 3.76 (s), 3.46 (s), 3.41–3.29 (m), 3.24–2.92 (m), 2.72 (d, J = 7.2 Hz), 2.42–2.27 (m), 2.24 (s), 2.08 (s), 1.92 (s), 1.75 (s), 1.71–1.56 (m), 1.57–1.41 (m), 1.36 (d, J = 7.0 Hz), 1.21 (d, J = 3.0 Hz), 1.11 (s), 0.99 (s), 0.88–0.70 (m).

Synthesis of metallacages by CDSA and analysis by mass spectrometry

Stock solutions of L0, L1, L2 and [Pd(MeCN)₄](BF₄)₂ (20 mM) were prepared in DMSO. The reaction of ligand L1 or L2 with the Pd(II) precursor [Pd(MeCN)₄](BF₄)₂ mixed in a 2:1 ratio in DMSO resulted in the homoleptic coordination cage $C1_{hom}$ or C2_{hom} within 1 h at RT. To achieve heteroleptic C1_{het} system, the stock solutions (20 mM) of L1 and L0 (20 mM) were mixed in a 1:3 ratio in DMSO (final concentrations 3 and 9 mM, respectively). Similarly, for C2_{het} stock solutions (20 mM) of L2 and LO were mixed in a 1:2 ratio in DMSO (final concentrations 4 and 8 mM, respectively). The ligand mixtures were stirred for 1 h at RT with 0.5 eq. (6 mM) of the Pd(II) precursor [Pd(MeCN)₄](BF₄)₂ to achieve a theoretical cage concentration of 3 mM. The cage mixtures were then diluted to a final cage concentration of 2 µM, using H₂O with 5% (v/v) MeCN and 0.1% (v/v) FA, and used for RP-HPLC, DI- and LC-HR-ESI-MS analysis.

Encapsulation studies by mass spectrometry

For the encapsulation of ReO_4^- , $C2_{hom}$ (3 mM in DMSO) was diluted with MilliQ water in 67% of H_2O . Stock solution of $NaReO_4$ (10 mM) in MilliQ water was freshly prepared, and an aliquot added to each cage solution to achieve cage: $NaReO_4$ ratio of 1:2 (0.9 mM:1.8 mM) in 72% of H_2O . After 30 min incubation at RT, the mixtures were diluted to a cage concentration of 4 μ M, using H_2O with 5% (v/v) of MeCN and 0.1% (v/v) FA, and used for DI-HR-ESI-MS or LC-HR-ESI-MS analysis.

Radiolabeling of ligands and cages

¹⁸F-labeling of the ligands. Radiolabeling of the SiFA moiety was carried out via isotopic exchange reaction.⁵⁸ Therefore, the required amount of fluoride-18 (approximately 800 MBq in [¹⁸O]H₂O) was loaded onto a SEP-Pak® Light (46 mg) Accell™ Plus QMA Cartridge (preconditioned with 10 mL H₂O) and dried with 8 mL anhydrous DMSO. The cartridge was then eluted into a LoBind Eppendorf tube using 40 mg of NH₄HCOO dissolved in 500 μL anhydrous DMSO. ~250 MBq of ¹⁸F⁻ eluent was added to 15 nmol of the ligand (1 mM in DMSO) and the labeling was complete after 10 min at RT. The reaction mixture was quenched using 10 mL H2O and the labeled ligand was fixed on an Oasis® HLB (30 mg) Light cartridge (preconditioned with 10 mL EtOH and 10 mL H₂O). The cartridge was washed with 10 mL H2O and the purified ligand was inversely eluted using 300 µL EtOH. Analysis of the labeled ligand was performed via radio RP-HPLC (10-80% B in 15 min, C18 column) and radio TLC (silica gel coated aluminum TLC plated F_{254} , MeCN/ H_2O , 8/2, +10% sodium acetate (2 M), +1% TFA).

¹⁸F-_DPepH3-_DDap((SiFA)BA)-Ebes-L0 (¹⁸F-L1)

RCY: 44%, RCP: 98%.

Radio RP-HPLC (10-80% B in 15 min): $t_R = 12.4$ min.

¹⁸F-_DPepH3-(SiFA)SeFe-Ebes-L0 (¹⁸F-L2)

RCY: 47%, RCP: 99%.

Radio RP-HPLC (30–60% B in 15 min): t_R = 10.4 min.

Self-assembly of $^{18}\text{F-}[Pd_2(\text{L1})_4][BF_4)_4$ ($^{18}\text{F-C1}_{hom}$) and $^{18}\text{F-}[Pd_2(\text{L2})_4][BF_4)_4$ ($^{18}\text{F-C2}_{hom}$). 30 nmol of L1 or L2 (2.0 eq., 5 mM in DMSO) were labeled according to the previously described procedure. The resulting solution of $^{18}\text{F-L1}$ or $^{18}\text{F-L2}$ was evaporated at 80 °C with an applied N₂ flow. 3 μ L of [Pd (MeCN)₄](BF₄)₂ (4.0 eq., 20 mM in DMSO) and 3 μ L of DMSO were added to the labeled ligand. After 4 h, the reaction was complete and analysed via radio RP-HPLC (40–70% B in 15 min, C4 column).

 $^{18}F-[Pd_2(\text{L1})_4](BF_4)_4$ ($^{18}F-\text{C1}_{\text{hom}}$) Radio RP-HPLC (40–70% B in 15 min): t_R = 9.6 min. $^{18}F-[Pd_2(\text{L2})_4](BF_4)_4$ ($^{18}F-\text{C2}_{\text{hom}}$) Radio RP-HPLC (40–70% B in 15 min): t_R = 7.7 min.

Lipophilicity determination ($\log D_{\rm pH7.4}$)

The octanol-PBS partition coefficient ($\log D_{\mathrm{pH7.4}}$ values) were determined by addition of 0.5 MBq of the labeled ligand to a 1.5 mL LoBind Eppendorf tube, filled with 500 µL octanol and 500 µL PBS (n=8). The tubes were vortexed for 3 min at RT and centrifuged (9000 rpm, 5 min, RT) using a Heraeus Pico 17 centrifuge by Thermo Scientific (Waltham, USA) ($24 \times 1.5/2.0$ mL rotor with ClickSeal-Lid). Out of each tube 200 µL of each solvent layer were taken out and the activity was quantified separately using a γ -counter by PerkinElmer Inc. (Langerwehe, Germany). Lipophilicity of the cages could not be determined due to stability issues upon vortexing of their solution.

$$\begin{split} \log D_{\rm pH7.4} \left(^{\mathbf{18}}\text{F-L1} \right) &= 2.53 \pm 0.05. \\ \log D_{\rm pH7.4} \left(^{\mathbf{18}}\text{F-L2} \right) &= 2.07 \pm 0.07. \end{split}$$

Cell culture

U87 cells were cultivated at 37 °C (+5% CO₂) in MEME (Sigma Life Science) containing 10% fetal bovine serum (Thermo Fisher), 1% L-glutamine (Sigma Life Science) and 1% penicillin–streptomycin (Sigma Aldrich). The cells were split 1/6 twice a week using trypsin–EDTA solution (Thermo Fisher) for detachment. The cells were tested Mycoplasma-free every month.

In ovo studies

All *in ovo* experiments were carried out on fertilized Dekalb white or brown eggs (Henry Stewart & co. Ltd, UK). Experiments took place before embryonic day 14 (E14). The eggs were incubated for up to 14 days at 12–14 $^{\circ}$ C in a wine cooler (Haller) with humidified atmosphere until use. To grow tumors on the chick chorioallantoic membrane (CAM), eggs were cleaned with Brinsea disinfectant (100×) and moved to an incubator (Brinsea) with a temperature of 38.7 $^{\circ}$ C and 48%

humidity. The first day of incubation at this temperature was embryonic day 0 (E0). To loosen the CAM from the eggshell the incubator trays were slowly tilted from one side to the other until E3. On E3, eggs were taken from the incubator for window cutting. The eggs were rolled to prevent the CAM sticking to the shell and placed on a cushioned holder. They were pierced at the wide base where the air cell is located and approximately 5 mL of albumin was removed through the hole using a syringe with a 19G needle. The hole was then resealed with Scotch Magic tape. Next, a square of tape was placed onto one side of the egg and four rectangularly arranged holes were punched into the shell through the tape. A rectangular window (1 × 2 cm) was made with sharp dissection scissors by carefully cutting three sides of a rectangle into the shell using the holes for orientation and without damaging the inner shell membrane. The window was closed with tape and the eggs placed in the incubator again until E7. U87 cells were maintained as described previously and cell culture media was renewed 24 h prior to harvesting. On the day of inoculations (E7), cells were detached from the flask, resuspended in media and aliquots with 3×10^6 cells were prepared. The aliquots were centrifuged for 3 min at 500g, 4 °C in a Centrifuge 5424 R (Eppendorf) and stored on ice. Meanwhile, Matrigel® Matrix Basement Membrane (Corning) was defrosted on ice. Eggs were taken from the incubator, placed on an egg holder and the windows were opened to locate the CAM. After dabbing the CAM dry with a sterile lens tissue, a suspension of the U87 cell pellet in 20 µL of Matrigel was pipetted onto the CAM. The eggs were closed with tape, labelled accordingly and placed in the incubator for a further 7 days. On E14 the eggs were removed from the incubator and placed on a cushioned holder. For tumor imaging the shell window was enlarged to allow for direct injection of the radiotracer into a CAM blood

Preparation of the radiotracer for *in ovo* studies. The radiolabeled ligand ¹⁸F-L1 was prepared as described before. The obtained solution in EtOH was reduced to 100 μL in an N₂ stream at 60 °C and then diluted with PBS to obtain until EtOH <25%. A sample was taken for quality control *via* radio RP-HPLC on a 1260 Infinity II LC-instrument (Agilent Technologies, MeCN (+5% $\rm H_2O$ + 0.1% $\rm TFA$)/ $\rm H_2O$ (+0.1% $\rm TFA$), $10\% \rightarrow 80\%$ MeCN gradient over 15 min, 1.5 mL min⁻¹ flow rate) with a GABI Nova detector (Elysia-Raytest) on a ZORBAX Exclipse XDB-C18 Analytical 4.6×250 mm 5 Micron column (Agilent Technologies). For *in ovo* injection, 150 μL an activity of 5 MBq were prepared by further dilution with saline (0.9 wt% NaCl).

In ovo PET imaging. On E14 a CAM vein was cannulated and 90 μ L of a 1 mg mL⁻¹ solution of the anaesthetic medetomidine (Virbac) was pipetted onto the surface of the CAM. Eggs were left for 15 min at RT before receiving an intravenous bolus injection of ~3 MBq of the labelled radiotracer on the imaging bed (<150 μ L), followed by 50 μ L PBS (Sigma Life Science). A 60 min dynamic PET scan was performed using a Mediso NanoScan PET/CT system (1–5 coincidence mode; 3D reconstruction; computed tomography (CT) attenuation cor-

rected; scatter corrected). Throughout the scan the eggs were kept at 37 °C. CT images were obtained for attenuation correction (180 projections; semi-circular acquisition; 50 kVp; 300 ms exposure time). The acquired PET data was reconstructed into 19 bins of 4 × 15 seconds, 4 × 60 seconds, and 11 × 300 seconds (Tera-Tomo 3D reconstructed algorithm; 4 iterations; 6 subjects; 400–600 keV; 0.3 mm³ voxel size). VivoQuant software (v2.5, Invicro Ltd) was used for analysis of the reconstructed images. Regions of interest (ROIs) were drawn manually using PET signal and 5–30 min summed dynamic PET to decide location of the brain. Time *versus* radioactivity curves (TACs) were generated using the percentage injected dose per mL (% ID per mL) and from this the area under the time *versus* radioactivity curve (AUC) was generated.

Ex ovo biodistribution. Biodistribution studies were carried out right after the PET scans. The embryos were sacrificed through cervical dislocation, and tumor, brain, liver and bone samples were collected. The organs were placed in weighed scintillation vials, weighed and counts were measured using a gamma-counter. The radioactivity was calculated as the percentage uptake of injected dose per gram of tissue (% ID per g). A dilution series of the labelled radiotracer in PBS was prepared, the activity determined with a Carpentec and measured in the γ-counter together with the organ samples to serve as calibration curve.

Results and discussion

Ligands synthesis and characterization

To achieve the assembly of the desired radiolabelled metallacages, the bispyridinyl ligand L0 (3,5-bis(3-ethynylpyridine) phenyl)acetic acid (Fig. S1-S4†) was functionalized via amide bond formation with a PEG linker (Ebes) acting as spacer and further attached to a SiFA synthon for ¹⁸F-radiolabeling, as well as tethered to the BBB-shuttle peptide DPepH3 (Fig. 2A). Two different SiFAs were selected: a monofunctional SiFAbenzoic acid ((SiFA)BA) prosthetic group⁵⁵ which can be attached in a branched fashion between the peptide and the PEG linker as⁵⁵ in L1, and a new bifunctional silicon-based fluoride acceptor ((SiFA)SeFe) recently developed in our group.⁵⁶ SiFAs are important synthons in radiochemistry since they can undergo ¹⁸F-fluorination under mild conditions compatible, for example, with the presence of bioactive molecules in the radiotracer. Unfortunately, a major challenge is the high lipophilic character of the SiFA moiety, which can lead to an unfavourable slow hepatic excretion.^{59,60} The latter decreases the image quality and increases off-target radiation dose to the abdomen. To partly overcome this problem, ((SiFA)BA) was developed featuring a carboxylic functionality.⁵⁵ An important progress was achieved with the ((SiFA)SeFe) synthon, whereby the amino acid-like bifunctionality increases its hydrophilic character, and also enables its direct incorporation in a ligand sequence in a bridged fashion as in L2. This feature reduces the steric demand and further counterbalances the SiFA lipophilicity.56

Ligands L1 and L2 were synthesized *via* semi-automated fluorenylmethoxycarbonyl solid-phase peptide synthesis (Fmoc-SPPS) by consecutive amide coupling on a Rink amide resin starting from _DPepH3 (Fig. 2A). After resin cleavage and semi-preparative RP-HPLC purification, the ligands were obtained in moderate yield of 11% and 17% for L1 and L2, respectively and excellent RP-HPLC purity (98%). Full characterization of L1 and L2 was carried out by HR-ESI-MS as well as by ¹H NMR and ¹H-DOSY NMR spectroscopy (Fig. S5–S8 and S9–S12,† for L1 and L2, respectively).

Metallacage synthesis and characterization

Homoleptic metallacages. One advantage of Pd_2L_4 -type cages is their straightforward synthesis, achievable by mixing the ligand and the metal precursor in stoichiometric amounts at RT for less than one hour. Thus, addition of ligands **L1** or **L2** to $[Pd(MeCN)_4](BF_4)_2$ in DMSO afforded the homoleptic cages $C1_{hom}$ and $C2_{hom}$, respectively (see Experimental for details) which were fully characterized by different methods (Fig. S13–S16 and S17–S19,† for $C1_{hom}$ and $C2_{hom}$, respectively). Analysis via RP-HPLC showed a shifted retention time t_R with respect to the ligands from 6.4 to 8.9 min for $C1_{hom}$ (Fig. S13†), and from 5.2 to 7.3 min for $C2_{hom}$ (Fig. S17†). Moreover, only traces of the free ligand were detected indicating full conversion at this concentration (mM). Also, the cages remained intact for at least 24 h under these conditions (Fig. S20†).

NMR spectroscopy is the most applied method to assess the formation of MCgs by CDSA. The ¹H NMR spectra of C1_{hom} and C2hom are shown in Fig. S15 and S18,† respectively. In both cases, the ligand's α-pyridyl protons H_a and H_b are exposed to a new electronic environment upon metal coordination, which results in their deshielding shift in the ¹H NMR spectrum.⁶¹ Exemplary, the pyridyl protons of L2 shift from 8.79 ppm and 8.61 ppm to 9.78 ppm and 9.43 ppm, respectively, indicating the formation of a supramolecular species (Fig. 2C). Additional ¹H-DOSY NMR spectra of the cages revealed, as expected, a decrease in the diffusion coefficient upon CDSA (Fig. S16, S19† and Fig. 2C). The SEGWE calculator⁶² was used to determine the cages' solvodynamic radii from the corresponding diffusion coefficients, which amount to 2.29 and 1.80 nm for C1hom and C2hom, respectively. The Pd₂L₄ assemblies were further investigated by DI-HR-ESI-MS or LC-HR-ESI-MS. Successful CDSA of C1hom was confirmed by a peak at 661.8748 m/z, attributed to the 11⁺ charge state of C1_{hom} adducts with formic acid (FA) and acetonitrile (MeCN) molecules from sample preparation (Fig. S14† and Table 1). Concerning cage C2hom, the mass spectrum obtained by DI shows a number of peaks corresponding to C1_{hom} adducts with FA and solvent molecules, or to $Pd_2(L2)_3$ species (Fig. 2D top). Formation of the coordination cage C2hom was also revealed by LC-HR-ESI-MS analysis, whereby a peak at t_R = 13.94 min in the total ion chromatogram (TIC) corresponds to species with m/z value of 637.3228, attributed to the 11⁺ charge state of the C2hom adduct with formic scid related mass (Fig. 3 and Table 1). The LC-HR-ESI-MS also shows the presence of a

Table 1 Calculated and experimental m/z values (most abundant isotope) for MCg species and host-guest adducts with perrhenate

Compound	Formula	m/z		
		Theoretical	Measured	Observed adducts
C1 _{hom}	$C_{360}H_{496}F_{4}N_{76}O_{56}Pd_{2}Si_{4} \\$	661.8758 (z = 11)	661.8747 664.2367	$[M^{4+} + FA + ACN + 7H^{+}]^{11+}$ $[M^{4+} + 2FA + Na + 7H^{+}]^{11+}$
[Pd ₂ (L1) ₃]	$C_{270}H_{372}F_3N_{57}O_{42}Pd_2Si_3$	686.4560 (z = 8)	686.4570 692.3329 695.2059 700.5811 704.9545 709.2046	$\begin{split} & \big[M^{4+} + FA + 4H^+ \big]^{8+} \\ & \big[M^{4+} + 2FA + 4H^+ \big]^{8+} \\ & \big[M^{4+} + 2FA + Na^+ + 4H^+ \big]^{8+} \\ & \big[M^{4+} + 2DMSO + 4H^+ \big]^{8+} \\ & \big[M^{4+} + Cl^- + 2DMSO + 5H^+ \big]^{8+} \\ & \big[M^{4+} + 2Cl^- + 2DMSO + 6H^+ \big]^{8+} \end{split}$
C2 _{hom}	$C_{352}H_{484}F_{4}N_{72}O_{52}Si_{4}Pd_{2} \\$	637.3200 (z = 11)	637.3203 641.0483 643.5020	$\begin{aligned} & \left[\mathbf{M}^{4+} + \mathbf{FA} + 7\mathbf{H}^{+} \right]^{11+} \\ & \left[\mathbf{M}^{4+} + \mathbf{FA} + \mathbf{ACN} + 7\mathbf{H}^{+} \right]^{11+} \\ & \left[\mathbf{M}^{4+} + 2\mathbf{FA} + \mathbf{Na} + 7\mathbf{H}^{+} \right]^{11+} \end{aligned}$
$ReO_4 \subset C2_{hom}$	$C_{352}H_{484}F_{4}N_{72}O_{56}Si_{4}Pd_{2}Re$	705.5519 (z = 10) $655.9506 (z = 11)$	705.5536 655.9501	$[M^{4^{+}} + 2FA + 6H^{+}]^{10^{+}}$ $[M_{3}^{+} + 8H^{+}]^{11^{+}}$
$[Pd_2(\mathbf{L2})_3]$	$C_{264}H_{363}F_3N_{54}O_{39}Si_3Pd_2\\$	670.8237 (z = 8)	670.8237 683.4458 687.8202	$[M^{4+} + 2FA + 4H^{+}]^{8+}$ $[M^{4+} + 2DMSO + Cl^{-} + 5H^{+}]^{8+}$ $[M^{4+} + 2DMSO + 2Cl^{-} + 6H^{+}]^{8+}$
$ReO_4{\subset}\left[Pd_2(\mathbf{L2})_3\right]$	$C_{264}H_{363}F_3N_{54}O_{43}Si_3Pd_2Re$	696.4408 (z = 8)	696.4399	$[M^{3+} + FA + 5H^{+}]^{8+}$
C1 _{het}	$C_{156}H_{166}FN_{25}O_{20}Pd_{2}Si$	594.2134 (z = 5) $742.7652 (z = 4)$	594.2142 742.7651	$ig[M^{4+} + H^+ ig]^{5+} \ ig[M^{4+} ig]^{4+}$
C2 _{het}	$C_{154}H_{163}FN_{24}O_{19}SiPd_2$	582.8092 (z = 5)	582.8102 605.8088	$[M^{4+} + H^{+}]^{5+}$ $[M^{4+} + 2FA + Na]^{5+}$
$[\mathrm{Pd}_2(\mathrm{L2})_2(\mathrm{L0})_2]$	$C_{220}H_{270}F_2N_{40}O_{30}Si_2Pd_2$	609.2671 (z = 7)	609.2677 625.4096	$[M^{4+} + 3H^{+}]^{7+}$ $[M^{4+} + 2FA + Na^{+} + 2H^{+}]^{7+}$

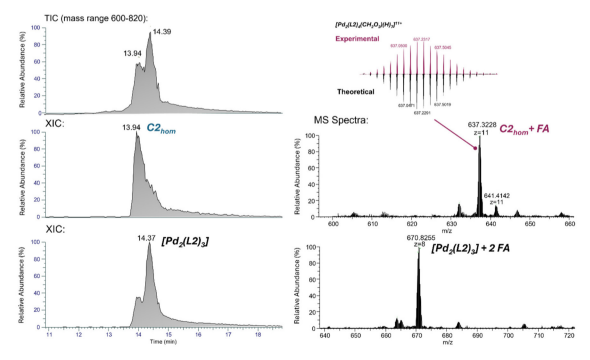


Fig. 3 Characterization of the homoleptic cage C2_{hom} by LC-HR-ESI-MS. Left: TIC (mass range between 600–820 m/z) and extracted ion chromatograms (XIC) of ions m/z 637.32 (z = 11, C2_{hom} at $t_R = 13.94$ min) and 670.83 (z = 8, [Pd₂(L2)₃] + 2 FA at $t_R = 14.37$ min). Right: mass spectrum of $C2_{hom}$ + FA at t_R = 13.94 min (comparison of theoretical and isotopic pattern is provided as insert) and $[Pd_2(L2)_3]$ + 2 FA at t_R = 14.37 min. The mobile phases were A: H₂O and B: MeCN, both with 0.1% formic acid (FA). The flow rate was 0.05 mL min⁻¹ and sample elution was performed by using a gradient from 20% to 40% of B over 3 min, followed by a gradient from 40% to 80% of B over 15 min.

 $Pd_2(L2)_3$ coordination complex at $t_R = 14.37 \text{ min } (m/z 670.8255, z = +8)$. Finally, the TIC spectrum demonstrates presence of free ligand L2 at $t_R = 14.60 \text{ min } (\text{data not shown})$. These results suggest the possible detachment of the fourth ligand in the MCg structure induced by the chromatographic elution. Nevertheless, it is evident that all the possible supramolecular species can be fully separated and properly characterized.

Next, stability studies were conducted in solvents relevant for different assays, radiolabeling and analytic procedures. Therefore, cage solutions were prepared as described in the Experimental section (1 mM theoretical concentration) and diluted 10-fold with DMSO, H2O, MeCN, HPLC solvent (MeCN/ H₂O (1/1) +0.1% TFA), 0.9 wt% NaCl, and PBS (pH 7.4), respectively, and analysed via RP-HPLC (Fig. 4 and Fig. S21, S22†). At this concentration, precipitation and partial disassembly of the cage were observed in PBS and isotonic NaCl solution, although less pronounced in the latter case over 1 h. This poses a problem for preclinical studies as salts are naturally present in physiological conditions, as well as for encapsulation of 99mTcO4-, which is eluted from the 99Mo/99Tc-generator with saline. Instead, both MCgs exhibited sufficient stability in water with 97% of C1hom and 75% of C2hom being intact after 4 h. In general, C2hom appears to be less stable than C1_{hom} despite the predicted reduced steric hindrance of the exo-functionalities. The presence of coordinating solvents like DMSO or MeCN induced formation of new supramolecular assemblies over 24 h, leaving only ca. 50% (or lower) remaining intact cage (Fig. 4). Of note, addition of water to MeCN as used in the HPLC conditions, increased the stability of the cages over time. Interestingly, an effect of concentration on cage stability was also recorded. For example, a 10-fold more concentrated solution of C2hom (1 mM) remained intact in DMSO solution for at least 24 h (Fig. S20†). Overall, these studies show that many factors can affect the stability of Pd2L4 MCgs, including steric effects of the exo-functionalities, cage surface charges which can be sensitive to the ionic strength of the medium, as well as interactions of the Pd(II) centres with

nucleophilic molecules (e.g. DMSO). In the specific case of $C1_{hom}$ and $C2_{hom}$, the obtained results suggest that both electrostatic and nucleophilic interactions may play a major role in destabilizing the cages in solution.

To get further insights into the effect of ligand concentration on the CDSA, $C1_{hom}$ and $C2_{hom}$ were self-assembled in DMSO directly to reach a theoretical concentration of 0.1 mM, which resulted in only partial cage formation (65% and 57% conversion to $C1_{hom}$ and $C2_{hom}$, respectively) (Fig. S23 and S24†), hence indicating that CDSA is concentration dependent and can result in different coordination species. This finding needs to be kept in mind especially when entering new fields such as radiochemistry, where reactions are performed in μ mol or even nmol scale.

Overall, both homoleptic metallacages $C1_{hom}$ and $C2_{hom}$ were successfully assembled and characterized using three different analytical techniques. However, further improvement in cage stability, which might be affected by the complex and sterically demanding peptidic ligands, is needed.

Heteroleptic metallacages. Heterolepticity in MCgs design is a promising strategy to increase multi-functionalization as ligands with different properties are combined in one unique structure. In recent work, the CDSA of a heteroleptic Pd₂L (L0)₃-type cage was a viable solution to introduce a peptidefunctionalized ligand featuring DOTA as chelator for lutetium-177.50 In order to favor the self-assembly of the heteroleptic cage C_{1_{het}} featuring a L₁/L₀ ratio of 1:3 - $[Pd_2(L_1)(L_0)_3]^{4+}$ - 1 eq. of L1 and 3 eq. of L0 were reacted with [Pd(MeCN)₄](BF₄)₄ in DMSO for 1 hour at RT. 1H NMR spectroscopy revealed a deshielding of the α -pyridinyl protons to 9.76 and 9.40 ppm, respectively, compared to the reference spectra of L1 and L0 (Fig. S25†). Moreover, the ¹H-DOSY NMR spectrum showed more than one newly formed species, upon addition of the Pd²⁺ precursor, suggesting a mixture of supramolecular assemblies (Fig. S26†). The self-assembly process was further investigated by RP-HPLC, where the species could be separated and analysed by MS (Fig. S27A and B†). By comparing the chroma-

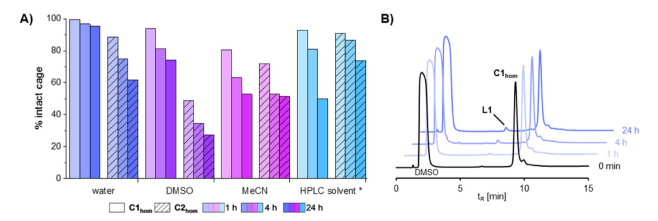


Fig. 4 (A) Stability study of $C1_{hom}$ and $C2_{hom}$ in different solvents analysed by RP-HPLC. The theoretical cage concentration after dilution with the respective solvent was 0.1 mM. *MeCN/H₂O (1/1) +0.1% TFA. (B) Stability of $C1_{hom}$ in Milli-Q H₂O monitored over 24 h by RP-HPLC (40–70% B in 15 min).

togram with those of the corresponding ligands and homoleptic cages – including $C0_{hom}~[\text{Pd}_2(L0)_4]^{4+}$ and $C1_{hom}$ – several peaks of similar intensity could already be attributed ($t_R(\mathbf{L0})$ = 3.2 min, $t_R(L1) = 11.5$ min, $t_R(C0_{hom}) = 8.3$ min). MS analysis of the peak at 11.1 min (25%) associated it to the target C1het species. The remaining peaks at 9.4 and 12.7 min could not be clearly identified by MS, but were likely to originate from heteroleptic CDSA leading to a statistical distribution of metallacages with different L1/L0 ratios (Fig. S27†). Interestingly, when changing the HPLC gradient from 30-60% B in 15 min (1 mL min⁻¹) to 20-40% B in 3 min and 40-80% B in 15 min (0.5 mL min⁻¹), a different chromatogram was obtained (Fig. S28†), whereby only a mean peak at $t_R = 6.7 \text{ min } (62.4\%)$ was observed, corresponding to C1_{het}. In order to identify the different assemblies, the mixture was further analysed by LC-HR-ESI-MS, using the optimized HPLC gradient (Fig. 5 and Table 1). In the obtained TIC graph, three main peaks could be resolved. At $t_R = 7.5$ min, the desired $[Pd_2(L1)(L0)_3]^{4+}$ $C1_{het}$ was the dominant species with m/z of 594.2142 and 742.7651 corresponding to the 5⁺ and 4⁺ charge state, respectively. Additional species eluting at t_R = 8.4 and 9.3 min corresponded to free L0 and L1 ligands, respectively.

For the assembly of the heteroleptic $[Pd_2(L2)(L0)_3]^{4+}$ $(C2_{het})$ cage, tuning of the ratio L2/L0 was required. Eventually, a mixture of heteroleptic species was obtained by combining 2.7 eq. of L2 and 1.3 eq. of L0 and reacting with [Pd(MeCN)₄]

(BF₄)₄ in DMSO for 1 hour at RT, as evidenced by ¹H and ¹H-DOSY NMR spectroscopy (Fig. S29 and S30†). The mixture of heteroleptic assemblies could be unpicked again by RP-HPLC using different gradients (Fig. S31 and S32†). Even in this case, the optimized gradient (20-40% B in 3 min and 40-80% B in 15 min (0.5 mL min⁻¹)) enabled the separation of a main peak at t_R = 6.8 min (58.7%) corresponding to $C2_{het}$ (Fig. S32†). Analysing directly the mixture by DI-HR-ESI-MS, the main desired species $(C2_{het})$ $[Pd_2(L2)(L0)_3]^{4+}$ (m/z =582.8102, z = +5) and $[Pd_2(\mathbf{L1})_2(\mathbf{L0})_2]^{4+}$ (m/z = 609.2677, z = +7) were identified, including comparison of the experimental and theoretical isotopic patterns (Fig. S33† and Table 1).

Encapsulation studies

One of the strategies to radiolabel metallacages is via encapsulation of the radioactive guest molecule. The encapsulation of ^{99m}TcO₄ and stability of different host-guest systems in vivo have already been demonstrated successfully by the groups of Lusby and Archibald as well as by Correia and Casini.⁴⁸ In both studies perrhenate was used as a 'cold' surrogate for pertechnetate and the host-guest interaction was investigated either by NMR spectroscopy or via in silico methods. Following these promising results, we investigated the encapsulation of perrhenate in our MCgs. Thus, 2 eq. of NaReO₄ were added to solutions of $C1_{hom}$ and $C2_{hom}$ in $D_2O/DMSO-d_6$ (1/1) and 1H NMR spectra were recorded after 30 min. Deuterated water was

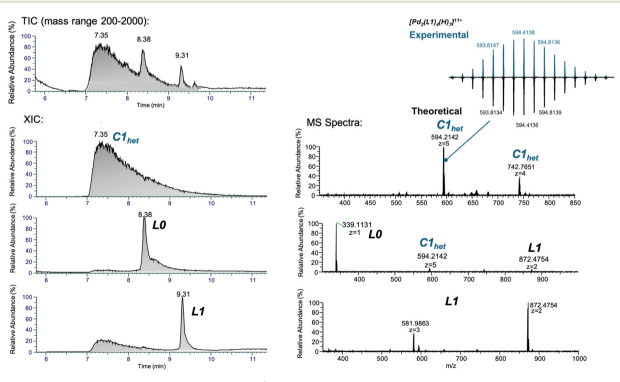


Fig. 5 Characterization of the heteroleptic cage $[Pd_2(L1)(L0)_3]^{4+}$ (C1_{het}) by LC-HR-ESI-MS. TIC (mass range between 200–2000 m/z) and XIC of ions m/z 594.21 (z = 5, C1_{het} at $t_R = 7.37$ min), m/z 399.1131 (z = 1, L0 at $t_R = 8.38$ min) and m/z 872.48 (z = 2, L1, $t_R = 9.31$ min). Right: mass specinosmic spec trum of $C1_{het}$ at $t_R = 7.37$ min (comparison of theoretical and isotopic pattern is provided as insert), L0 at $t_R = 8.38$ min and L1 at $t_R = 9.31$ min. The mobile phases were A: H₂O and B: ACN, both with 0.1% formic acid (FA). The flow rate used was 0.5 mL min⁻¹ and sample elution was performed by using a gradient from 20% to 40% of B over 3 min, followed by a gradient from 40% to 80% of B over 15 min.

added to support the encapsulation of the oxo-anion, which interacts with the host cavity via electrostatic forces and hydrogen bonding. 63 For C1_{hom} a deshielding of 0.01 ppm of the H_a proton was observed upon perrhenate addition and in the case of C2_{hom} the H_a proton was shifted by 0.03 ppm (Fig. S34 and S35†). No significant H_b shift was detected for both cage-perrhenate mixtures. Compared to the proton upfield shifts of a similar host-guest system (H_a: ~0.08 ppm, H_b: ~0.04 ppm), ⁴⁸ this result indicates poor encapsulation of perrhenate under these conditions, due to the high concentration of DMSO in the media, which does not reflect the clinical setting.

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Therefore, the ability of the cages to encapsulate perrhenate in predominantly aqueous media was additionally evaluated by DI-HR-ESI-MS in the case of C2hom. The comparison of the mass spectra of C2hom and the host-guest system (ReO₄⁻)⊂C2_{hom} in Fig. 2D, unambiguously revealed the formation of the host-guest adduct through the appearance of signals at m/z 655.9501 and 721.3443, which can be attributed to the 11⁺ and 10⁺ charge state of (ReO₄⁻)⊂C2_{hom}, respectively (Table 1). While the encapsulation is not quantitative, this result confirms some affinity of ReO₄⁻ for the cage cavity in aqueous solution, at concentrations more relevant to the clini-

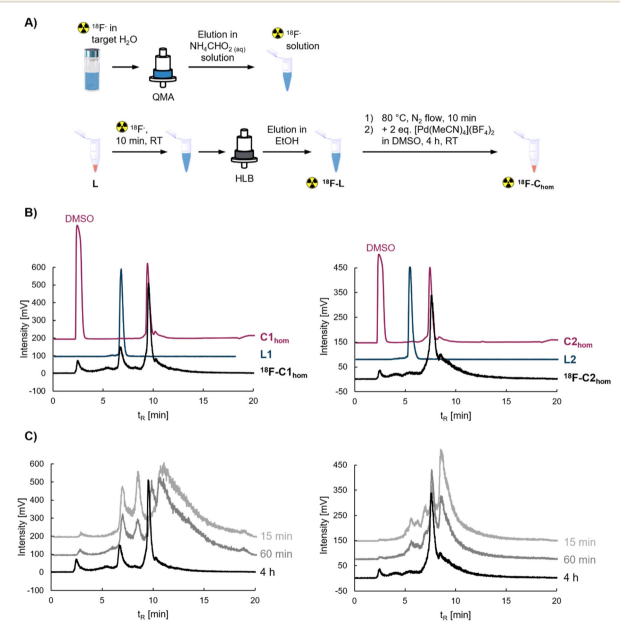


Fig. 6 (A) Schematic representation of the ¹⁸F-radiolabeling procedure of the ligands and CDSA of radiolabeled homoleptic MCgs. For fixation either a silica-based ion exchange cartridge (QMA) or a copolymer-based reverse phase cartridge (HLB) was used. (B) Radio chromatograms of ¹⁸F-C1_{hom} and ¹⁸F-C2_{hom} after 4 h. Chromatograms of the ligands (blue) and the co-injected non-radiolabelled cages (purple) are inserted as comparison. (C) CDSA of ¹⁸F-C1_{hom} and ¹⁸F-C2_{hom} monitored over 4 h by radio RP-HPLC.

cal setting, in the μ M range. However, it should also be noted that at variance with previously reported studies on similar MCgs, ⁴⁸ we were not able to preserve any host-guest complex following separation by LC.

Fluoride-18 radiolabeling and $log D_{pH7.4}$ determination

Next, the ligands were radiolabeled with fluoride-18 by [19F]-to-[18F] isotopic exchange. A Kryptofix-free synthesis route using cartridges as depicted in Fig. 6A was chosen.⁵⁸ This method achieved 60-70% radiochemical conversion (RCC) and resulted in the radiolabeled compounds ¹⁸F-L1 and ¹⁸F-L2 obtained in 44-47% radiochemical yield (RCY) and 98-99% radiochemical purity (RCP) as evidenced by radio TLC and radio RP-HPLC (Fig. S36-S39†). Since lipophilicity is an important parameter in radiotracers development, the distribution coefficient $(\log D_{\rm pH7.4})$ was determined for both ligands. High lipophilicity could cause low tumor accumulation in vivo due to an increased nonspecific liver uptake that elevates the off-target radiation level to the abdomen. 60 On the other hand, a study, in which the physicochemical properties of around 300 central nervous system (CNS) drugs were analysed and compared to non-CNS drugs, revealed that positive $log D_{pH7.4}$ values $(\log D_{\rm pH7.4}$ in the range 2-3) can be beneficial for BBBcrossing.64

In the case of L1, a $\log D_{\rm pH7.4}$ of 2.53 \pm 0.05 of the radiolabeled ligand was determined. To assess the effects of such high lipophilicity *in vivo*, the organ uptake of ¹⁸F-L1 was evaluated in the CAM model (Fig. 7A). This preclinical tool can be used for high-throughput tumor imaging in a time- and costeffective way and is in line with the ethical principles of reduction, refinement and replacement (3Rs).⁶⁵ Glioblastoma tumors from U87 cells were grown on the CAM of fertilized chicken eggs. After injection of approximately 3 MBq of radiolabeled ligand, a 60 min dynamic PET scan was performed, and representative time points are depicted in Fig. 7B. While, this model was not intended for BBB-translocation experiments, evaluation of the maximal brain uptake was attempted over time up to 60 min p.i. (Fig. 7C and D). Overall, the obtained values are in accordance with published data for the BBB shuttle peptide PepH3 (0.31 \pm 0.07% ID per g),⁵¹ as well as a PepH3-conjugated MCgs $(0.42 \pm 0.06\% \text{ ID per g})^{48}$ evaluated in mice models in vivo. However, further studies are necessary to validate these data, possibly using bimodal PET/ MRI imaging to better delineate the brain region. Moreover, long blood circulation and very high liver uptake demand further optimization of the ligand. In addition, bone uptake of $3.55 \pm 0.91\%$ ID per g might indicate defluorination of the ligand.66 Despite these hurdles, good tumor uptake was measured ex vivo, reaching 3.94 ± 1.31% ID per g at 60 min p.i. In the future, tumor accumulation could even be enhanced by combining the BBB-penetrating peptide with a glioblastomatargeting ligand in a heteroleptic approach.

In the case of **L2**, as expected, the $\log D_{\rm pH7.4}$ was 2.07 \pm 0.07, which is a substantial reduction compared to the (SiFA) BA ligand. Nevertheless, its evaluation in the CAM model was not performed, due to the predicted high liver uptake; moreover, our focus was to investigate the formation of radiolabeled cages.

Synthesis of ¹⁸F-labeled homoleptic metallacages

Since the _DPepH3-conjugated ligands **L1** and **L2** are modified with a SiFA moiety, labeling of the metallacage structure with

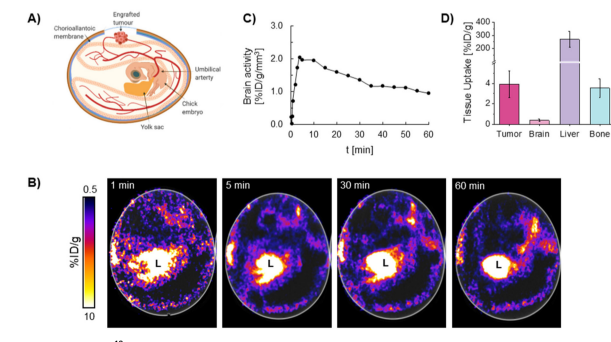


Fig. 7 In ovo evaluation of ^{18}F -L1. (A) Schematic representation of the chick CAM model. (B) Representative in ovo PET images from a 60 min dynamic scan (n = 4). The liver (L) is indicated. (C) Time-activity curve of the chick embryo brain. (D) Ex ovo biodistribution one hour p.i.

F-18 is possible via exo-functionalization. Here, the radiolabeling can either proceed before CDSA (pre-assembly), during CDSA (one-pot) or after CDSA (post-assembly). 43 In our case, to avoid cage disassembly in the relatively 'harsh' radiofluorination conditions, the ligands were radiolabeled pre-assembly and then CDSA was performed to form the homoleptic cages. Thus, after labeling 30 nmol of ligand (1 eq.) with ~200 MBq of fluoride-18 and elution in EtOH, the solvent was fully evaporated and CDSA was conducted by adding 2 eq. of [Pd (MeCN)₄](BF₄)₂ in 6 μL of DMSO (Fig. 6A). These conditions enabled the optimal concentration of each component (mM range) for successful CDSA. The self-assembly was monitored by radio RP-HPLC at 15 min, 60 min and 4 h (Fig. 6C). As depicted in Fig. 6B, 73% conversion to 18 F-C1_{hom} ($t_R = 9.6 \text{ min}$) was reached after 4 h with 18% remaining 18 F-L1 ($t_{\rm R}$ = 6.7 min), while full conversion to 18 F-C2_{hom} (t_R = 7.7 min) was achieved. In both cases slight defluorination (18F-C1hom: 9%, ¹⁸F-C2_{hom}: 3%) occurred as evidenced by the F-18 peak at t_R = 2.4 min. Further purification of the reaction mixtures by analytical radio RP-HPLC or via cartridges was not carried out due to the short half-life of F-18 of 109.7 min, as well as possible instability of the radiolabeled cages.

Conclusions

In this work, we aimed at further substantiating the applicability of supramolecular coordination complexes, namely porous MCgs, for radiopharmaceutical applications. These nano-structures can benefit from the self-assembly multicomponent design for functionalization and pharmacokinetics modulation, but at the same time their size range (1-few nm) enables their precise characterization by classical separation and analytical methods; thus, offering an ideal versatile chemical space to achieve next generation nuclear imaging agents.

Here, two bispyridinyl ligands featuring a BBB shuttle peptide- and different silicon-based fluoride acceptor synthons (L1 and L2) were successfully synthesized by solid-phase peptide synthesis and radiolabeled with fluorine-18. The lipophilicity of the ligands was determined in vitro, via standard $\log D_{\rm pH7.4}$ determination, and for L1 also ex vivo using the CAM model. Afterwards, formation of the respective homoleptic metallacages C1_{hom} and C2_{hom} was achieved by CDSA and confirmed by NMR spectroscopy, RP-HPLC and HR-ESI-MS. Importantly, ¹⁸F-radiolabeled homoleptic cages were achieved by labeling the ligands pre-assembly and via formation of the cages in situ. The optimization of the isotopic exchange protocol was an important step towards the future applicability of MCgs for PET imaging in a clinical setting.

Additionally, partial encapsulation of ReO₄⁻ into the cages following mixing of the components was evidenced by HR-ESI-MS; however, the extent and stability of the (ReO₄[−])⊂MCg host-guest adducts appeared less pronounced with respect to previously reported Pd₂L₄ cage structures.⁴⁸ Thus, further ligand optimization is necessary to validate the suitability of these porous systems for encapsulation of

anionic radioactive counterparts, such as 99mTcO4 for SPECT imaging or the β --emitting ¹⁸⁸ReO₄-.

In order to improve the stability of the homoleptic Pd-based cages in solution, heteroleptic MCgs were also synthesized by statistical self-assembly and characterized by ESI-MS. The obtained results showed that $[Pd_2L(L0)_3]^{4+}$ $(C1/2_{het})$ and $[Pd_2L_2(\mathbf{L0})_2]^{4+}$ were the main assembled species.

In the future, an option to increase MCg's stability and to enable optimized radiofluorination of the cages post-assembly, could be replacing Pd²⁺ with the more kinetically inert Pt²⁺ to achieve [Pt2L4]4+ cages. The latter have already shown to be more stable with respect to ligand exchange than their Pdbased counterparts, 67,68 which would also improve their stability in physiological conditions. Alternatively, organometallic building blocks endowed with higher metal-carbon bond stability could also be integrated in the cage scaffold.1

Overall, our results highlight the current challenges in the construction of targeted theranostics based on SCCs for future clinical applications in nuclear medicine. While much effort still needs to be spent on the development of metallacages for biomedical applications, their design flexibility and easiness of functionalization offer a unique toolbox for the obtainment of multimodal theranostics.

Author contributions

A. C. and L. R. were responsible for the project conceptualization and supervision. M. D. and A. C. wrote the first draft of the manuscript and all co-authors contributed to the Results and discussion sections. M. D., I. K. and A. C. designed the graphics. M. D. and D. K. performed the synthesis and radiolabeling of ligands and cages. I. K., L. R. and R. L. carried out high-resolution mass spectrometry studies and performed the data analysis. G. M.-A. and A. C. provided expertise in MS and NMR data analysis. M. D., L. M. S., M. E. G. and T. H. W. contributed to, and supervised, the in ovo studies.

Data availability

The data supporting this article have been included as part of the ESI.† Raw data for this article, including chromatograms, NMR spectra, MS data, gamma counter data for $\log D$ determination, in ovo imaging data are available at Zenodo at [https:// doi.org/10.5281/zenodo.15358484].

Conflicts of interest

The authors declare no conflict of interest.

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