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In vivo biodistribution of kinetically stable Pt₂L₄ nanospheres that show anti-cancer activity†

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There is an increasing interest in the application of metal–organic cages (MOCs) in a biomedical context, as they can offer non-classical distribution in organisms compared to molecular substrates, while revealing novel cytotoxicity mechanisms. Unfortunately, many MOCs are not sufficiently stable under *in vivo* conditions, making it difficult to study their structure–activity relationships in living cells. As such, it is currently unclear whether MOC cytotoxicity stems from supramolecular features or their decomposition products. Herein, we describe the toxicity and photophysical properties of highly-stable rhodamine functionalized platinum-based Pt₂L₄ nanospheres as well as their building blocks under *in vitro* and *in vivo* conditions. We show that in both zebrafish and human cancer cell lines, the Pt₂L₄ nanospheres demonstrate reduced cytotoxicity and altered biodistribution within the body of zebrafish embryos compared to the building blocks. We anticipate that the composition-dependent biodistribution of Pt₂L₄ spheres together with their cytotoxic and photophysical properties provides the fundament for MOC application in cancer therapy.

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Introduction

The biomedical application of metal–organic cages (MOCs) is getting increasing interest. Since the discovery and clinical success of the first inorganic anti-cancer drug cisplatin, research on more potent, selective, and multi-functional metalodrugs progressed.^{1,2} During the last 10 years, MOCs have emerged as potent alternatives to classical inorganic and organometallic complexes.^{3–5} MOCs are well-defined nanostructures, which are self-assembled by reversible coordination of multi-topic ligands (linkers) to metal (nodes).^{6–8} As MOCs are frequently composed of cytotoxic metals, they can act as anti-cancer metalodrugs. In this context, anticancer activity of many different MOCs in different shapes (spheres, prisms, helicates) and compositions (Ni₂L₃, Ni₄L₆, Pd₂L₄, Pd₁₂L₂₄, Pt₈L₂L₄^A₂^B₄, Ru₆L₂L₃^A₂^B₃) have been explored.^{9–22} MOCs have characteristic advantages over the use of molecular drugs, such as their large size (usually 1–10 nm), well-defined shape, and confined cavity

that can be exploited. These characteristics can improve their therapeutic profiles through alternative activity mechanisms. Amongst the cytotoxic MOC complexes, M₂L₄ nanospheres (Fig. 1) are especially interesting for reasons provided below.

M₂L₄ nanospheres are typically prepared by the combination of ditopic pyridine donors together with square planar palladium or platinum precursors (Fig. 1). This type of nanospheres have modulation possibilities by ligand variation or

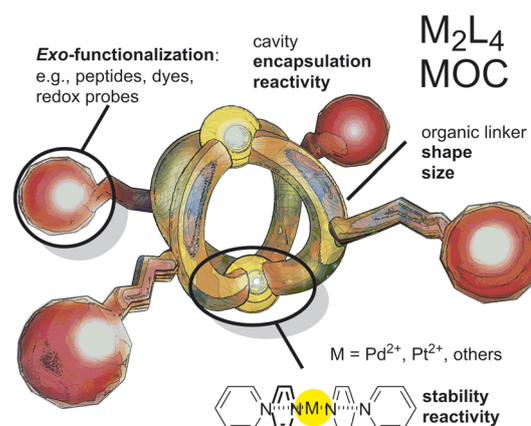


Fig. 1 Schematic representation of M₂L₄ spheres with different modulation possibilities. For example, outer functionalization for solubility; organic linker modification for size and shape modulation and metal modification for modulation of stability.

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functionalization and can be obtained in a variety of different shapes, which makes them interesting candidates for biological investigation.^{23–27} Furthermore, a number of reports describe the cytotoxicity of Pd₂L₄ nanospheres by disruption of cell membranes¹⁴ or alternative pathways^{28–30} or by encapsulation of toxic guest molecules (*i.e.* cisplatin),^{12,13,15,31} making further studies into this direction promising for the development of novel anti-cancer drugs. In this context, understanding how a certain shape of the nanosphere influences its cytotoxicity or the activity/toxicity of covalently attached drugs or encapsulated drugs against cancer (herein called structure–activity relationship) is important to design novel, potent and tumor-specific nanospheres. Unfortunately, Pd₂L₄ nanospheres were shown to decompose quickly when exposed to physiological conditions,^{12,15} hampering understanding of the cytotoxicity mechanism and development of structure–activity relationships. Despite numerous fascinating examples of *in vitro* application of M₂L₄ nanospheres, *in vivo* applications are rare.³² Imaging of nanospheres *in vivo* remains, due to the so called “heavy atom effect”, a problem and only a few fluorescent M₂L₄ spheres have been reported.^{33–35} Although, some M₂L₄ nanospheres have shown to be fluorescent, biological application requires a good fluorescent quantum yield and an emission wavelength preferably in the red light. Unfunctionalized nanospheres typically absorb and emit between 250 and 400 nm, which is impractical due to the absorption of living matter (*e.g.*, proteins) in the same region. Furthermore, red fluorescent probes allow for deeper tissue imaging and typically yield a brighter signal to background ratio.³⁶ Motivated by these challenges and by the potential of M₂L₄ nanospheres for therapeutic treatment, we explore in this contribution kinetically robust and fluorescent Pt₂L₄ nanospheres *in vitro* and *in vivo*. In a proof-of-concept study, we show that the well-defined arrangement of fluorescent drugs around nanospheres changes their biodistribution *in vivo* (using a zebrafish model), photophysical and cytotoxic properties, while the nanosphere itself (without covalent drug conjugation) acts as a delivery vehicle (for dyes or anti-cancer drugs) with little toxicity.

Results and discussion

Ligand design and nanosphere synthesis

Excellent photophysical properties of rhodamine B for *in vivo* imaging made this dye the subject of our studies leading to the design of two novel building blocks (Fig. 2). Next to the required spectroscopic features for *in vivo* detection, rhodamine B amide derivatives have shown recently potent cytotoxicity against different human cancer types.^{37,38} Furthermore, singlet oxygen formation using rhodamine dyes emerged as a tool in photodynamic therapy of tumor cells.^{39,40} Therefore, rhodamine functionalization of nanospheres can yield systems which are not only suitable for *in vivo* imaging, but also can have cytotoxic properties by themselves or through light driven singlet oxygen (¹O₂) formation. The rhodamine unit is attached *via* two different linker types to a sphere-forming building block (Fig. 2). We chose a long hydrophilic (tetra-ethylene glycol (TEG)) and a short linker (ethyl) to study the effect of the

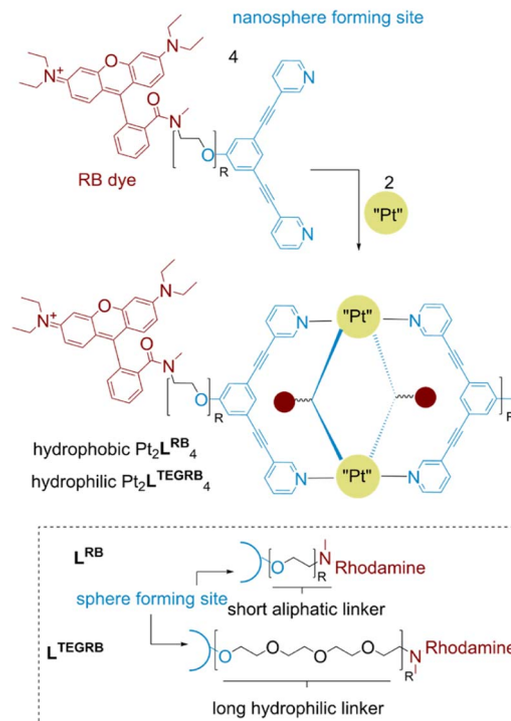


Fig. 2 Chemical structure of the two building blocks L^{RB} and L^{TEGRB} and the corresponding nanospheres Pt₂L^{RB}₄ and Pt₂L^{TEGRB}₄.

resulting well-defined 3-dimensional nanospheres on (A) photophysical properties, (B) cytotoxicity against cancer lines *in vitro* and (C) biodistribution within the body of zebrafish embryos. Both building blocks, L^{RB} and L^{TEGRB} are prepared according to standard organic synthesis protocols (summarized in Section SI1†).

To obtain kinetically robust nanospheres, we used platinum for the nanosphere formation, as the platinum–pyridine bond was demonstrated to be more kinetically inert compared to the conventionally applied palladium analogues.^{12,41,42} Pt₂L₄ spheres are prepared according to a recently published procedure.³⁸ L^{RB} (1 eq.), and [Pt(BF₄)(MeCN)₄] (0.6 eq.) were heated to 150 °C in acetonitrile-*d*₃ for 1 day (Fig. 3A). The observed downfield shifts of the pyridine protons in the ¹H-NMR spectra are in line with coordination to platinum (Fig. 3C). DOSY NMR spectra show the appearance of a slower diffusing species, with a calculated hydrodynamic radius of 1.1 nm, in line with the formation of Pt₂L₄ nanosphere (Fig. 3B). ESI-HR-MS analysis confirms selective formation of the desired structure by showing only signals corresponding to different charged states of the assembly with the general formula [Pt₂L^{RB}₄(BF₄)_x]^{(8–x)+} and matching isotope pattern for *x* = 2–8 (Fig. 3D). Crystals suitable for X-ray diffraction were grown from slow vapor diffusion of diethyl ether into a solution of the sphere in acetonitrile. The solid-state structure of Pt₂L^{RB}₄ confirms formation of the desired nanosphere (Fig. 4). The metal–metal distance and other structural properties are in accordance with similar M₂L₄ spheres. An interesting feature in the solid state is the bending of the rhodamine dyes towards the windows of the



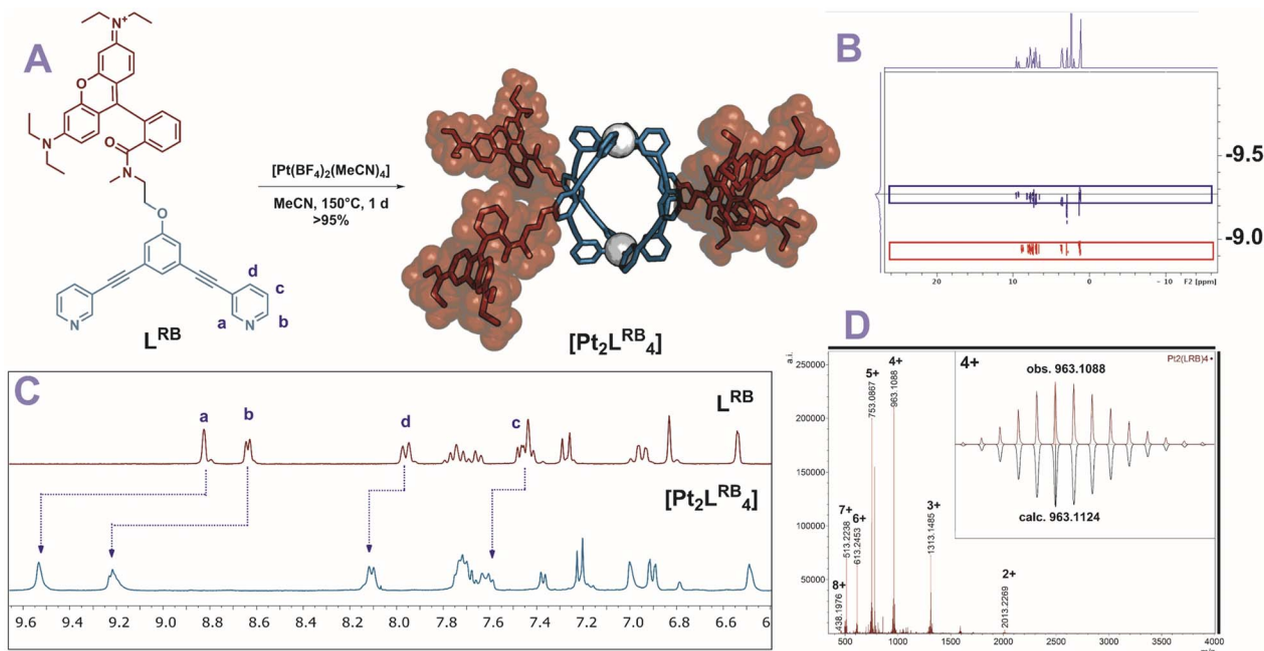


Fig. 3 Characterization of $[\text{Pt}_2\text{L}^{\text{RB}}_4]^{8+}$. (A) Reaction conditions for formation of nanospheres. Molecular structure of the displayed sphere was minimized on PM3 level. Carbon and nitrogen of nanosphere framework displayed in blue, rhodamine dye in red, platinum as white spheres. (B) Overlaid DOSY NMR of the $\text{Pt}_2\text{L}^{\text{RB}}_4$ nanosphere (blue) and the building block (red). (C) ^1H -NMR spectra of $[\text{Pt}_2\text{L}^{\text{RB}}_4]$ nanosphere and the corresponding building block. (D) ESI-MS spectrum of $\text{Pt}_2\text{L}^{\text{RB}}_4$.

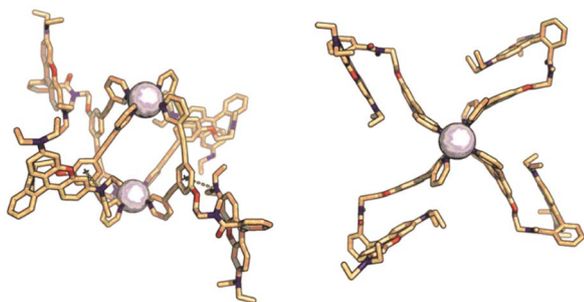


Fig. 4 Solid state structure of $\text{Pt}_2\text{L}^{\text{RB}}_4$, sphere framework displayed in stick form; carbon in beige, nitrogen in blue, oxygen in red, platinum as white spheres. Side view indicating cation- π interaction between rhodamine and central benzene group. Top view showing the densely packed arrangement of the sphere.

nanosphere. This bending provides the most compact packing and involves four π - π interactions between the rhodamine and the central aromatic ring of the building block with a distance of 3.7 Å (in line with 6 Å limit of other typical cation- π and π - π interactions).⁴³ The displayed densely packed form might occur in water as the hydrophobic surface area is minimized in this conformation.

$\text{Pt}_2\text{L}^{\text{TEGRB}}_4$ is obtained *via* an analogous experimental procedure to $\text{Pt}_2\text{L}^{\text{RB}}_4$ with similar characteristic spectroscopic features (Fig. S12–15†). Both spheres were precipitated in diethyl ether and redissolved in $\text{DMSO-}d_6$ for further application.

Stability and protein binding

A requirement for the development of structure-activity relationships in bio-medical application of MOCs is their stability under physiological conditions. Therefore, the kinetic stability of $\text{Pt}_2\text{L}^{\text{RB}}_4$ and $\text{Pt}_2\text{L}^{\text{TEGRB}}_4$ in the presence of common biological nucleophiles (Cl^- , imidazole) and a reducing agent/transition-metal ligand (glutathione) was investigated using ^1H -NMR and UV/vis spectroscopy. ^1H -NMR experiments were carried out in $\text{DMSO-}d_6$: D_2O 1:2 where 0.83 mM nanospheres were treated with 80 eq. NaCl, 30 eq. imidazole and 60 eq. of glutathione at 37 °C. Under these conditions, neither of the two nanospheres show any decomposition within 12 h (Fig. S16†). UV/vis absorbance spectra of the spheres were measured at 37 °C in phosphate-buffered saline (PBS, 1 M, pH 7.4). Absorbance spectra in water show characteristic absorbance at 570 nm and 535 nm (Fig. S16†). The $\text{Pt}_2\text{L}^{\text{TEGRB}}_4$ nanosphere shows no decrease in absorption after 12 h at 37 °C in PBS (Fig. S16†), providing support for excellent halide and buffer stability of this Pt_2L_4 nanosphere. In contrast, $\text{Pt}_2\text{L}^{\text{RB}}_4$ shows a decrease in absorbance after 12 h (approx. 20%). As the characteristic sphere absorption features (almost equivalent absorption at 535 and 570 nm) remain the same, we anticipate the decrease of $\text{Pt}_2\text{L}^{\text{RB}}_4$ absorption due to the observed adsorption of the sphere to the surface of the UV/vis cuvette.

Because these MOC's represent multi-charged species with different hydrophobic character (from the organic linkers), we also decided to study the stability of the two nanospheres in the presence of protein, which is typically overlooked in stability assays.⁴⁴ Although there are many proteins present *in vivo*, a very



prominent protein is albumin. This protein is present in all mammals (*e.g.*, bovine albumin or human albumin) in relatively high concentrations (60% of all plasma proteins) and has the main function to transport apolar and positively charged molecules. As these two functions (apolar and positively charged) are present in our structures (*e.g.*, the positive Pt corners and the organic linker or the positive charged aromatic rhodamine units), we anticipated a strong interaction of our nanospheres with albumin and studied these (Fig. 5B). When we mixed $\text{Pt}_2\text{L}^{\text{RB}}_4$ with bovine serum albumin (BSA), a change in rhodamine absorption was observed. The signal at 520 nm showed a decrease in intensity, whereas that at 570 nm was increased. This characteristic change can be attributed to an interaction between the sphere and BSA (Fig. 6). In order to investigate this further, we performed isothermal titration calorimetry (ITC) by addition of BSA to nanosphere solutions in HEPES (10 mM, pH = 7.2) (Fig. 5A). Analysis of the ITC thermogram of $\text{Pt}_2\text{L}^{\text{RB}}_4$ provides a high binding constant of the nanosphere to BSA ($K_D = 2.41 \pm 1.2 \times 10^{-7}$ M). The TEG extended sphere $\text{Pt}_2\text{L}^{\text{TEGRB}}_4$ is less hydrophobic relative to $\text{Pt}_2\text{L}^{\text{RB}}_4$ and exhibits a lower binding affinity to BSA ($K_D = 2.59 \pm 0.9 \times 10^{-6}$ M). In both scenarios binding is dominated by hydrophobic interactions (as judged by the signature plots, Fig. S19†). The binding affinities between the nanospheres and BSA are unexpectedly high compared to molecular rhodamine B ($K_D = 10^{-4}$ M). These high binding affinities suggest that albumin binding might contribute to the biodistribution of nanospheres in the body and influence their *in vivo* activity. We therefore included the BSA complexes in our subsequent biodistribution investigations (*vide infra*).§

Fluorescence studies

With stable nanospheres in hand, we continued to investigate their photophysical properties. The absorption profiles of ligands L^{RB} and L^{TEGRB} and nanospheres $\text{Pt}_2\text{L}^{\text{RB}}_4$ and $\text{Pt}_2\text{L}^{\text{TEGRB}}_4$ in PBS (1 M, pH = 7.4, 0.2% DMSO) are shown in Fig. 6. Both ligands L^{RB} and L^{TEGRB} display similar broad absorption bands

centered at 570 nm with molar absorption coefficients of $\epsilon = 18 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ and $60 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ respectively. After nanosphere formation, higher molar extinction coefficients of $\epsilon = 33 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ ($\text{Pt}_2\text{L}^{\text{RB}}_4$) and $88 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ ($\text{Pt}_2\text{L}^{\text{TEGRB}}_4$) and an extra absorption located around 535 nm appears for $\text{Pt}_2\text{L}^{\text{RB}}_4$. The increase in this absorption is an indication of rhodamine–rhodamine stacking, consistent with previous reports (*e.g.*, rhodamine 6G).⁴⁵ Rhodamine-dimer formation is less pronounced for $\text{Pt}_2\text{L}^{\text{TEGRB}}_4$. We anticipate that the hydrophilic TEG linkers reduce the hydrophobicity of the nanosphere in a significant manner, which results in less inter- and intramolecular rhodamine-stacking.

After exploring the absorption features of the two rhodamine containing building blocks and of the corresponding spheres, we studied the emission of the ligands and nanospheres (Fig. 6B). All investigated structures emit orange light with a center around 600 nm. L^{RB} displays a moderate quantum yield (QY) of 14%. L^{TEGRB} is four times more efficient with a QY of 55%. Following the same trend as the building blocks, $\text{Pt}_2\text{L}^{\text{RB}}_4$ and $\text{Pt}_2\text{L}^{\text{TEGRB}}_4$ displayed a QY of 4% and 29% respectively. The lower QY after nanosphere formation can be attributed to rhodamine-stacking, cation– π interactions and the heavy atom effect.

Addition of BSA enhanced the QY of $\text{Pt}_2\text{L}^{\text{RB}}_4$ by 3-fold from 4% to 12%, which can be attributed to rhodamine–rhodamine stack dissociation as supported by a decrease in absorbance at 535 nm (which is characteristic for rhodamine–rhodamine stacks). In a similar manner, but to a lesser extent, also the QY of $\text{Pt}_2\text{L}^{\text{TEGRB}}_4$ was enhanced by 20% (from 29% to 35%). The QY of the applied building blocks remained unchanged upon addition of BSA. In summary, a hydrophilic TEG linker between a dye and the nanosphere allows to maintain similar emissive properties of dyes in PBS buffer as before sphere formation. When the dye is placed close to the sphere, inter-/intramolecular dye stacking and the heavy atom effect led to a significant reduction of QY. In this context, BSA has shown to be a suitable candidate to break up rhodamine-dimer formation

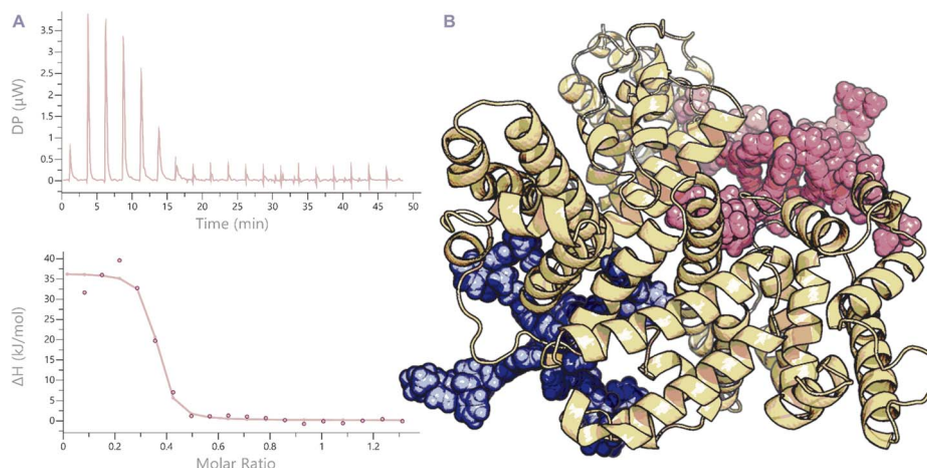


Fig. 5 ITC thermogram of the addition of BSA to $\text{Pt}_2\text{L}^{\text{RB}}_4$ (A). Below: fitted function to obtain the corresponding binding constants. (B) MD minimized model showing binding of $\text{Pt}_2\text{L}^{\text{RB}}_4$ into different hydrophobic pockets of BSA.



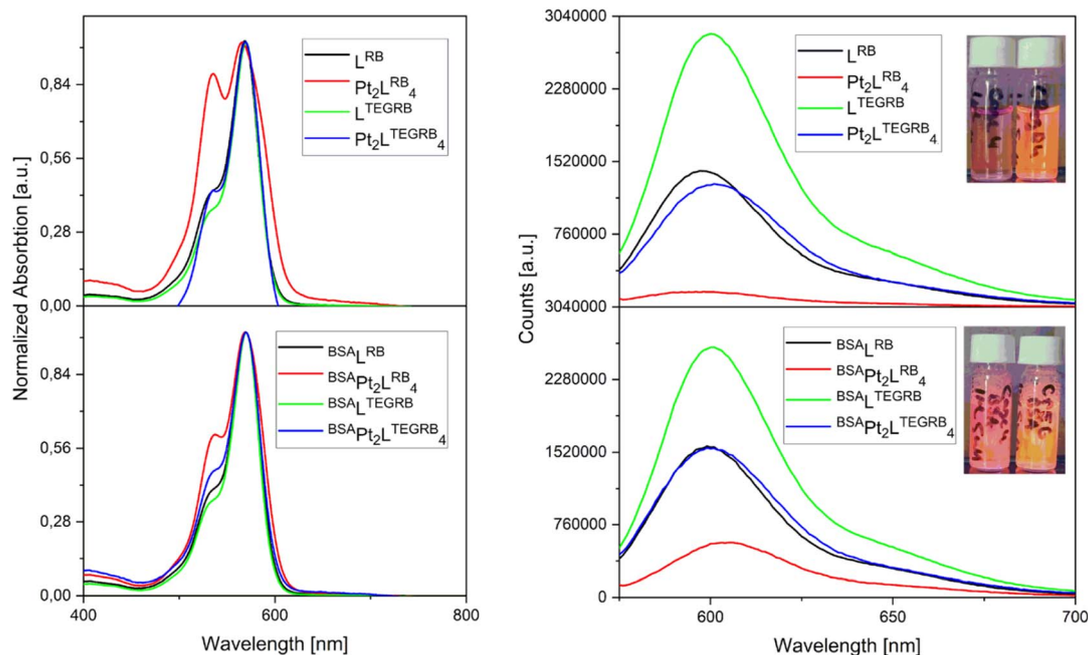


Fig. 6 Normalized (maximum absorption = 1) absorption spectra of the two building blocks L^{RB} and L^{TEGRB} and the corresponding nanospheres $Pt_2L^{RB}_4$ and $Pt_2L^{TEGRB}_4$ in PBS (left). Emission spectra of L^{RB} , L^{TEGRB} , $Pt_2L^{RB}_4$ and $Pt_2L^{TEGRB}_4$ in PBS (right). Insert: picture of the two nanospheres under UV irradiation (left: $Pt_2L^{RB}_4$; right: $Pt_2L^{TEGRB}_4$).

of sphere aggregates in PBS, improving their QY by a factor of 1.2 and 3.

Cytotoxicity

While some specific palladium-based Pd_2L_4 nanospheres are highly toxic,^{14,28–30} other structures, especially with similar design to our framework have shown little toxicity.^{12,15,31} Recent studies revealed the high toxicity of rhodamine derivatives,^{37,38} making our systems, with the well-defined nanosphere and linked toxic units, interesting candidates for structure-cytotoxicity assays. We anticipate differences between the molecular building blocks and their nanosphere analogues. We studied the building block L^{TEGRB} , the nanospheres $Pt_2L^{RB}_4$ and $Pt_2L^{TEGRB}_4$ as well as a dummy sphere $Pt_2L^{OMe}_4$ without rhodamine functionalization against two cell lines: PC3M-Pro4 (prostate carcinoma) and cisplatin-resistant MDA-MB-231 (breast cancer). The dummy sphere $Pt_2L^{OMe}_4$ shows no significant cytotoxicity within 24 h ($IC_{50} > 20 \mu M$, Table 1). The rhodamine-functionalized building block shows very high cytotoxicity against PC3M-Pro4 ($IC_{50} = 1.5 \mu M$ for L^{TEGRB}) and against MDA-MB-231 ($IC_{50} = 0.5 \mu M$ for L^{TEGRB}). The corresponding nanospheres show reduced toxicity compared to their molecular building blocks against both cell lines. The cytotoxicity of the rhodamine incorporated into the nanospheres is 4 to 13-times lower against both cell lines compared to the corresponding building blocks ($IC_{50} = 6.8–10 \mu M$ for $Pt_2L^{RB}_4$ and $IC_{50} = 6.8 \mu M$ for $Pt_2L^{TEGRB}_4$). Overall, the building block and the nanospheres display similar cytotoxicity for both cancer cell lines, regardless of the linker type between the nanosphere forming part of the building block and the dye. Although

rhodamines have emerged very recently as cytotoxic anti-cancer agents, currently the mechanism of action is not fully understood. Rhodamine-based dyes can possess cytotoxicity due to DNA complexation⁴⁶ by π -stacking and electrostatic attraction. More recent work describes rhodamine derivatives as cytotoxic agents which can disrupt/destroy mitochondrial membranes, causing apoptosis.^{37,38,47} Although, our study was performed thus far on only two different cell lines, it demonstrates how reduction of toxicity to an applicable level can be achieved by the formation of supramolecular nanospheres.

In vivo biodistribution

To investigate the influence of self-assembly of rhodamine building blocks to a nanosphere on the biodistribution of the

Table 1 IC_{50} values (per rhodamine in brackets) for ligand and nanospheres $Pt_2L^{RB}_4$, $Pt_2L^{TEGRB}_4$ and $Pt_2L^{OMe}_4$ against selected cancer cell lines^a

IC_{50} (μM)		
Compound	PC3M-Pro4 (prostate)	MDA-MB-231 (breast)
L^{TEGRB}	0.5 (0.5)	1.5 (1.5)
$Pt_2L^{RB}_4$	1.7 (6.8)	2.5 (10)
$Pt_2L^{TEGRB}_4$	1.7 (6.8)	1.7 (6.8)
$Pt_2L^{OMe}_4$	41.7 (n.a.)	>20 (n.a.)

^a The cells were incubated for 24 hours at 37 °C, 5% CO_2 . Then 9 μL WST-1 reagent (Sigma) was added to each well mixed, and incubated for 2 hours at 37 °C, 5% CO_2 . Then the absorbance at 400 nm was measured for each well, using the Infinite M1000Pro plate reader (Tecan).



rhodamine dyes, we used the zebrafish model using a transgenic *kdr1:EGFP* and *mpeg1:EGFP* zebrafish line (with green emissive endothelium of the vasculature and EGFP expressing macrophages, respectively, for schematic overview of zebrafish, see Fig. 8B).^{48,49} Acute toxicity studies of Pt₂L^{TEGRB}₄ in wild type zebrafish embryos over the course of 4 days revealed no apparent toxicity, abnormalities, or premature death in any of the injected embryos intravenously injected with concentrations up to 200 μM in 10 mM HEPES pH 7.4. Zebrafish do not express albumin, but instead a protein with similar function: vitamin D binding protein (VDBP),⁵⁰ making them also a good platform to study the biodistribution of albumin-bound spheres. The nanospheres were heated at 80 °C in DMSO (to break up possible aggregation), then diluted with HEPES (10 mM, pH 7.4) prior injection. 1 nL of 5 μM probe samples were directly injected intravenously into the duct of Cuvier in 48–80 hours post fertilization (hpf) zebrafish embryos. Initially, BSA-nanosphere complexes were prepared prior to injection by mixing 10 mM stock solutions of nanosphere in DMSO with PBS or HEPES buffer containing 150 μM BSA (15 eq.) to obtain a final concentration of 5 μM nanosphere bound to BSA. The BSA-nanosphere samples were directly injected without prior

heating. The biodistribution of the nanospheres and BSA-nanosphere complexes within the body of the zebrafish embryos was analyzed by confocal imaging at 90 minutes post injection (mpi) (Fig. S23†). After ITC studies revealed stoichiometry of roughly two spheres bound to one BSA-molecule (Fig. 5A), and with therapeutic application in mind, the complexes were prepared by injecting a higher 160 μM Pt₂-L^{TEGRB}₄ concentration into a 160 μM BSA solution in 10 mM HEPES, pH 7.4. The biodistribution of this higher concentration of the complex was studied in 72 hpf transgenic *kdr1:EGFP* zebrafish embryos.

A non-injected control showed red autofluorescence emission only in the yolk sac and eyes (Fig. S22†). Pt₂L^{RB}₄ shows rapid non-selective uptake within 30 min throughout the body by endothelial cells (Fig. 7B). The relatively high uptake in the scavenger endothelial cells (SECs) in the tail compared to endothelial cells in the arteries can be explained by the majority of spheres interacting with serum proteins (negatively charged) directly after entering the bloodstream. These sphere-protein complexes likely have a net negative charge, directing them to the SECs. The fluorescent intensity of Pt₂L^{RB}₄ is qualitatively much lower (Fig. 7B is 5 times less bright under identical laser

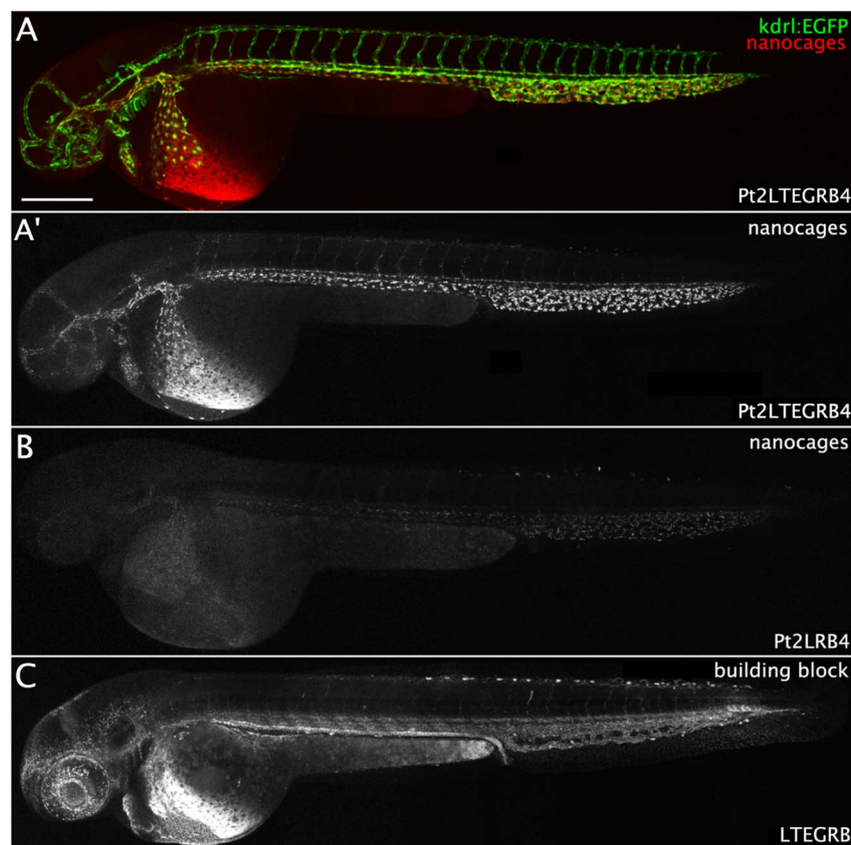


Fig. 7 Confocal images showing biodistribution of nanospheres and building block in 54 hours post fertilization (hpf) transgenic *kdr1:EGFP* zebrafish embryos injected intravenously in the duct of Cuvier with (A) Pt₂L^{TEGRB}₄ (shown in red), which is rapidly cleared from circulation into the venous and arterial vasculature (shown in green) within 30 minutes post injection (mpi); (A') nanocages (shown in white), showing the uptake pattern throughout the vascular endothelium; (B) Pt₂L^{RB}₄ showing lower fluorescence intensity, similar to Pt₂L^{TEGRB}₄, rapidly taken up in venous and arterial vessel walls within 30 mpi; (C) L^{TEGRB} remained primarily in circulation besides accumulation in the pronephric duct at 1.5 hours post injection (hpi). Scale bar represents 250 μm.



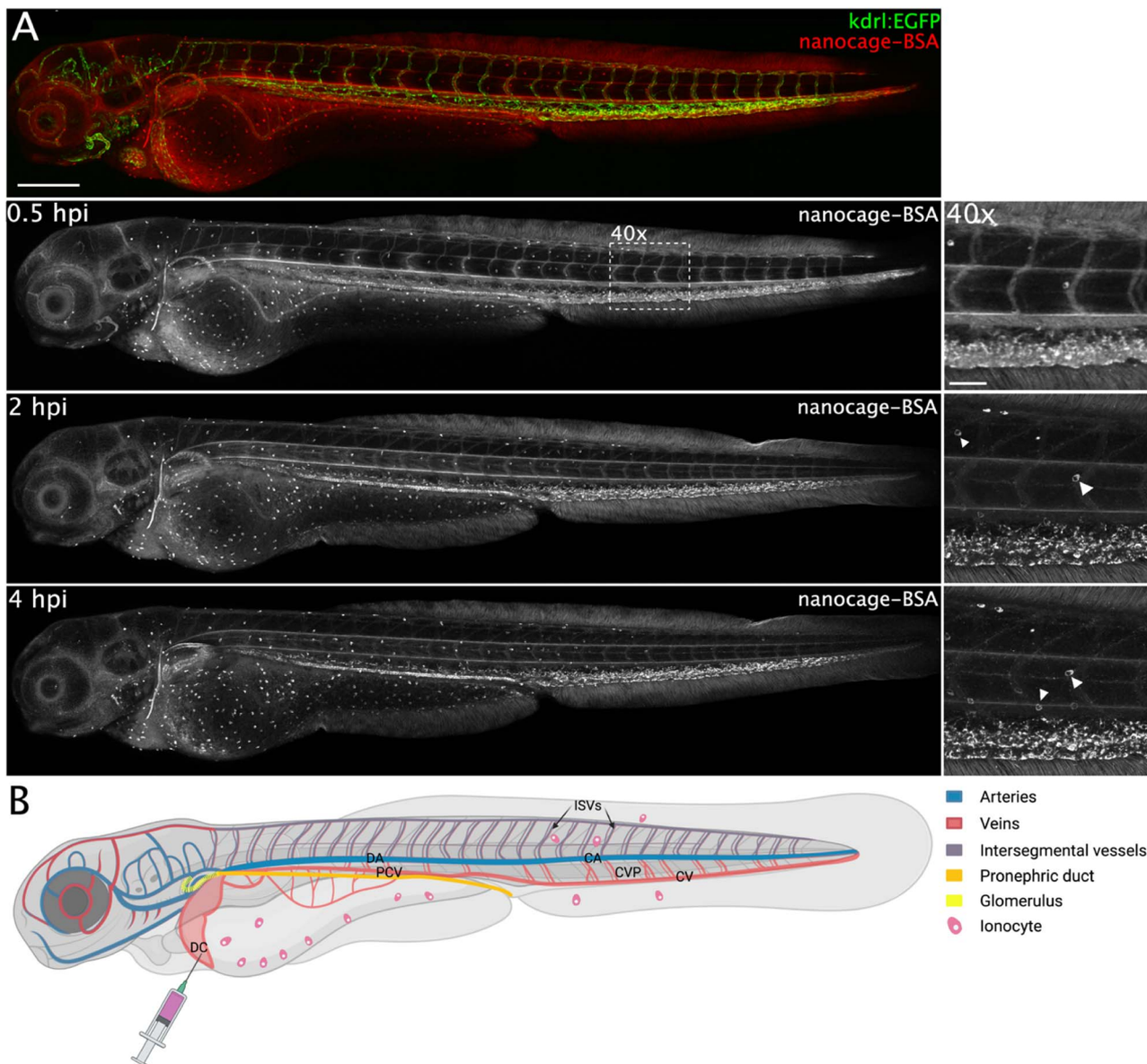


Fig. 8 (A) Confocal images showing biodistribution of $160 \mu\text{M}$ $\text{Pt}_2\text{L}^{\text{TEGRB}}_4 \supset \text{BSA}$ intravenously injected (1 nL) in 72 hpf transgenic *kdrl:EGFP* zebrafish embryos. At 30 minutes post injection (mpi) the complexes remained primarily in circulation, although accumulation in ionocytes, mainly around the yolk sac, and slight uptake in scavenger endothelial cells (SECs), as well as accumulation in the pronephric duct was already visible. At 2 hpi and 4 hpi this accumulation became increasingly pronounced, with clear accumulation in the glomerulus (schematically depicted on the right) visible at 4 hpi, although a (small) fraction remained in circulation. Accumulation in ionocytes is indicated in 2 hpi (40 \times) and 4 hpi (40 \times) by white arrow heads. Scale bars represent $250 \mu\text{m}$ (0.5 hpi) and $50 \mu\text{m}$ (0.5 hpi, 40 \times). (B) Schematic representation of accumulation of nanosphere materials in different tissues of zebrafish embryos. Intravenous injection of 1 nL nanosphere solution for biodistribution studies was performed in the Duct of Cuvier (DC). Upon injection, the nanospheres enter the heart, are pumped into the arterial vessels, among others the dorsal aorta (DA), followed by the caudal aorta (CA), intersegmental vessels (ISVs), caudal vein (CV) and caudal vein plexus (CVP) containing scavenger endothelial cells (SECs). Finally, the circulation continues back to the heart via the posterior cardinal vein (PCV) and the DC.

and detector settings compared to Fig. 7A). Similar endothelial cell uptake was observed when $\text{Pt}_2\text{L}^{\text{TEGRB}}_4$ was injected (Fig. 7A). $\text{Pt}_2\text{L}^{\text{TEGRB}}_4$ showed more intense fluorescence, comparable to free building block L^{TEGRB} (Fig. 7C compared to Fig. 7A, same settings were used when taking the pictures), which we attribute to the molecular behavior of dyes due to the long TEG linker between dye and nanosphere. As opposed to the spheres, building block L^{TEGRB} remained primarily in circulation,

besides some accumulation in the pronephric ducts (analogous to kidneys in mammals) at 1.5 hpi (Fig. 7C).

The presence of BSA changed the biodistribution of both nanospheres in a similar fashion (e.g., Fig. 8 and S23 \dagger). Both nanospheres show better fluorescence (especially $\text{Pt}_2\text{L}^{\text{RB}}_4 \supset \text{BSA}$) compared to the free nanospheres in the absence of BSA, in line with the enhanced fluorescent quantum yield under physiological conditions. Furthermore, the circulation of both



nanospheres was significantly enhanced (Fig. 8 and S23†). In line with anionic charged molecules, $\text{Pt}_2\text{L}^{\text{TEGRB}}_4 \supset \text{BSA}$ is taken up by scavenger endothelial cells in the tail (Fig. 8, likely due to the net negative charge of BSA (18^-)). In addition, the sphere–BSA complexes showed major accumulation in a different cell type identified as ionocytes located superficially on the skin (Fig. 8). Before full development of gills, kidneys, and intestine of fish embryos, both gas exchange and ion exchange take place through specialized cells in the skin. Ionocytes in the skin of fish embryos regulate body fluid pH and ionic composition.⁵¹ We assume that the very small size of the spheres allows them to diffuse relatively easily through the tissue of the fish, far beyond the bloodstream and the vascular endothelium allowing accumulation in ionocytes. Furthermore, the chemistry of rhodamine B might explain the bright staining of these cells. Under acidic conditions, the dye's fluorescence is bright.⁵² A certain type of ionocytes in embryonic zebrafish is involved in acid and ammonia secretion, providing an acidic environment in which rhodamine B fluorescence is brighter red compared to the intensity in physiological pH (in circulation).⁵¹ Likely, due to the high degree of H^+ present in these cells and the lipophilic nature of rhodamine B the (mitochondrial) membranes of these mitochondria-rich cells are brightly stained.

Additionally, we studied the possible uptake of nanosphere–BSA complexes by macrophages using transgenic zebrafish line *mpeg1:EGFP* carrying macrophages expressing EGFP. $\text{Pt}_2\text{L}^{\text{RB}}_4 \supset \text{BSA}$ showed no apparent uptake by macrophages (as the green and red emissions do not overlap, Fig. S23†). For both nanosphere–BSA adducts, as well as for the free building block, accumulation in the pronephric duct (part of the developing kidney) was observed, starting at around 45 mpi (Fig. 8 and S23†). Additionally, in case of nanosphere–BSA complexes, accumulation in the glomerulus (part of developing kidney) was observed starting at 2 hpi (Fig. 8). The developing kidney plays an important role in the clearance of various molecules, notably carrier proteins such as albumin and VDBP, likely explaining the accumulation of the BSA-bound spheres in this tissue. As for the building blocks, their positive charge might cause adsorption to (carrier) proteins in circulation, possibly including VDBP, which likely points at an accumulation mechanism similar to that of the nanosphere–BSA complexes. For a schematic representation of the tissues in which the injected samples accumulated refer to Fig. 8B.

With this set of experiments, we demonstrate a high degree of tunability of the biodistribution properties of Pt_2L_4 nanospheres by (A) their hydrophobic character and (B) by protein complexation. Depending on the probe formulation, we were able to achieve fast uptake by endothelial cells (plain nanospheres), longer circulation, uptake by scavenger endothelial cells in the tail (nanosphere–BSA complexes, anionic charged small particle) and accumulation in ionocytes and the pronephric duct (building blocks and nanosphere–BSA complexes). We anticipate that the high degree of tunable particle delivery together with their cytotoxic properties provides the fundament for new avenues of coordination-based spherical assembly application in cancer therapy. Depending on the cancer type which should be treated, sticky spheres (such

as $\text{Pt}_2\text{L}^{\text{RB}}_4$) can be useful when the drugs can be directly injected into the tumor. For other applications (*e.g.*, intravenous administration) more circulating materials are required (such as $\text{Pt}_2\text{L}^{\text{TEGRB}}_4 \supset \text{BSA}$). We have shown that simple ligand modification of the nanospheres has a huge influence on their properties *in vivo*.

Conclusions

While clinical use of coordination-based nanospheres is still in its infancy, they have promising properties, some of which are described in this paper. We have shown a clear step forward by designing two new *exo*-functionalized self-assembled Pt_2L_4 nanospheres with rhodamine groups, as was shown by NMR, mass spectroscopy, and single-crystal X-ray diffraction. The nanospheres were shown to be stable under physiological conditions for 12 h and exhibit good fluorescence, making Pt_2L_4 nanospheres suitable candidates for toxicity and the bio-distribution study in the body of zebrafish embryos. Previous studies on Pd_2L_4 nanospheres displayed decomposition under *in vivo* conditions, making it unclear what the effect of the supramolecular structure is on toxicity and biodistribution. With the first set of robust and fluorescent Pt_2L_4 nanospheres, we made some important observations on the effects of the supramolecular structure for *in vitro* and *in vivo* applications. We have demonstrated that albumin binding occurs with a very high binding constant in comparison to molecular analogues, which enhances the fluorescent properties and in turn, can play a role in the distribution of typically rather hydrophobic nanospheres *in vivo*. The toxicity of nano-assembled rhodamine drugs is downregulated towards specific cancer types in comparison to the molecular analogue which stems from the well-defined three-dimensional composition of the nanospheres, a handle that may come useful for specific drug design when too toxic materials are dealt with. The nanospheres show faster cell uptake than the building blocks they are assembled of and display alternated distributions in the presence of albumin. We foresee that Pt_2L_4 or similar systems could act as a universal platform for imaging purposes, which can be expanded to the delivery of therapeutically active drugs by encapsulation strategies (*e.g.*, ref. 12, 13, 15, and 31) or covalent attachment.

Data availability

All relevant data is provided in the ESI† of the paper. If other data is needed, it will be made available on reasonable request.

Author contributions

Conceptualization: EOB, RK, AK, JNHR; formal analysis: EOB, RK; funding acquisition: JNHR, BdB, AK, NIM; investigation: EOB, RK, SM, DAP, IK; supervision: JNHR, BdB, AK, NIM; validation: EOB, RK, SM, DAP, IK, JNHR, BdB, AK, NIM; visualization: EOB, RK; writing – original draft: EOB; writing – review & editing: EOB, RK, SM, DAP, IK, JNHR, BdB, AK, NIM. EOB and RK contributed equally to this work.



Conflicts of interest

There are no conflicts to declare.

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Notes and references

§ Although BSA and other albumins are biologically speaking not identical (ca. 75% identical), all of the albumins share the same function and possess pockets for positively charged and hydrophobic molecules. BSA as such acts chemically as a good estimation for other albumin forms which are less available for research (e.g., human serum albumin HSA).

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