Chem Soc Rev



TUTORIAL REVIEW

View Article Online



Cite this: Chem. Soc. Rev., 2020, 49 2886

Received 9th December 2019 DOI: 10.1039/c8cs00986d

rsc.li/chem-soc-rev

Metal-based imaging agents: progress towards interrogating neurodegenerative disease

Adam C. Sedgwick, Da James T. Brewster II, Da Peter Harvey, D*bc Diana A. Iovan, Od Graham Smith, Od Xiao-Peng He, Od * He Tian, Od * Jonathan L. Sessler * and Tony D. James * * and Tony D. James * * *

Central nervous system (CNS) neurodegeneration is defined by a complex series of pathological processes that ultimately lead to death. The precise etiology of these disorders remains unknown. Recent efforts show that a mechanistic understanding of the malfunctions underpinning disease progression will prove requisite in developing new treatments and cures. Transition metals and lanthanide ions display unique characteristics (i.e., magnetism, radioactivity, and luminescence), often with biological relevance, allowing for direct application in CNS focused imaging modalities. These techniques include positron emission tomography (PET), single-photon emission computed tomography (SPECT), magnetic resonance imaging (MRI), and luminescent-based imaging (Luml). In this Tutorial Review, we have aimed to highlight the various metalbased imaging techniques developed in the effort to understand the pathophysiological processes associated with neurodegeneration. Each section has been divided so as to include an introduction to the particular imaging technique in question. This is then followed by a summary of key demonstrations that have enabled visualization of a specific neuropathological biomarker. These strategies have either exploited the high binding affinity of a receptor for its corresponding biomarker or a specific molecular transformation caused by a target species, all of which produce a concomitant change in diagnostic signal. Advantages and disadvantages of each method with perspectives on the utility of molecular imaging agents for understanding the complexities of neurodegenerative disease are discussed.

Key learning points

- (1) The importance of molecular imaging for the understanding of neurodegenerative disease
- (2) To understand the importance of a specific metal in each imaging modality
- (3) The strategies to design an effective metal-based molecular imaging agent
- (4) Current limitations of each imaging modality
- (5) Future perspectives

Introduction

Central Nervous System (CNS) degeneration involves a series of neurological conditions that affect millions of people worldwide. These diseases stem from the progressive dysfunction and death of neuronal cells, which lead to loss of motor function and memory. A common theme amongst neurodegenerative processes is the progressive misfolding, aggregation, and accumulation of protein and peptide aggregates resulting in cellular dysfunction, loss of synaptic connections, and brain damage. The most common disorders, Alzheimer's disease (AD), Parkinson's disease (PD), Huntington's disease (HD), Lewy Body dementia, prion diseases, and amyotrophic lateral sclerosis (ALS), also share phenotypic similarities and all result

^a Department of Chemistry, The University of Texas at Austin, Austin, Texas 78712-1224, USA. E-mail: sessler@cm.utexas.edu

^b Department of Biological Engineering, Massachusetts Institute of Technology, Cambridge, Massachusetts 02139, USA

^c Sir Peter Mansfield Imaging Centre, Division of Clinical Neuroscience, School of Medicine, University of Nottingham, Nottingham NG7 2RD, UK. E-mail: peter.harvey@nottingham.ac.uk

^d Department of Chemistry, University of California, Berkeley, California, USA

^e Division of Radiotherapy & Imaging, Institute of Cancer Research, 123 Old Brompton Road, London, SW7 3RP, UK

^f Key Laboratory for Advanced Materials and Joint International Research Laboratory of Precision Chemistry and Molecular Engineering, Feringa Nobel Prize Scientist Joint Research Center, School of Chemistry and Molecular Engineering, East China University of Science and Technology, 130 Meilong Rd, Shanghai 200237, China. E-mail: xphe@ecust.edu.cn, tianhe@ecust.edu.cn

^g Department of Chemistry, University of Bath, Claverton Down, Bath, BA2 7AY, UK. E-mail: t.d.james@bath.ac.uk

in death.¹ Current treatment protocols offer short-term palliative benefits with no long-term improvement for patients or relief for caregivers. Not surprisingly, the pharmaceutical industry and academic institutions alike are seeking to develop new therapies, as well as working to elucidate the biochemical processes responsible for these disorders. High attrition rates

in clinical trials have led to the belief that understanding the precise etiology of CNS disease is critical to developing an effective treatment or cure. As discussed below, metal-based imaging modalities may have a critical role to play in this search. The goal of this tutorial review is to summarize recent progress in the area.



From left to right: Adam C. Sedgwick and Tony D. James

Adam C. Sedgwick received his MChem (2014) and PhD (2018) with Tony D. James and Steven D. Bull at the University of Bath. He is now a postdoctoral research fellow working under the supervision of Prof. Jonathan L. Sessler at The University of Texas in Austin. His research interests are in the realms of stimuli-responsive materials, molecular imaging agents, and theranostic agents.

Tony D. James is a Professor at the University of Bath and Fellow of the Royal Society of Chemistry. He received his BSc from the University of East Anglia (1986), PhD from the University of Victoria (1991), was Postdoctoral Research with Seiji Shinkai (1991–1995) and Royal Society University Research Fellow at the University of Birmingham (1995–2000). He was awarded the Daiwa-Adrian Prize (2013), Inaugural CASE Prize (2015), MSMLG Czarnik Award (2018) and currently holds a prestigious Royal Society Wolfson Research Merit Award. (2017–2022).



James T. Brewster

James T. Brewster II grew up in Valley Center, California and obtained his BS with honours in chemistry from the University of California, Irvine with mentorship from Professor Kelvin W. Gee, Dr Derk J. Hogenkamp, and Dr Timothy Johnstone. James recently finished his PhD with Professor Jonathan L. Sessler at The University of Texas. James is currently conducting his post-doctoral work with Professor Andrew G. Myers at Harvard University.



Peter Harvey

Pete Harvey received his MChem (2009) and PhD (2013) with Prof David Parker FRS in the Chemistry Department at Durham University. He was then awarded an ESPRC Doctoral Prize Fellowship to work at The University of Manchester. In 2015, he was awarded a Wellcome Trust-MIT Postdoctoral Fellowship in the Department of Biological Engineering at MIT. He recently began his independent career in molecular MRI as a Nottingham Research Fellow.



Diana A. Iovan

Diana A. Iovan received her BS in chemistry in 2012 from the University of Richmond. She then moved to Harvard University for her graduate studies with Prof. Ted Betley. After earning her PhD in inorganic chemistry in 2017, Diana joined Prof. Chris Chang's laboratory at UC Berkeley as a Howard Hughes Medical Institute Fellow of the Life Sciences Research Foundation.



Graham Smith

Graham Smith obtained his PhD from the University of Surrey and was a PDRA at Newcastle University and Imperial College. In 2011 he became a Lecturer in Molecular Imaging at the University of Hull where he helped establish a new PET research facility. In 2013 he moved to the Institute of Cancer Research to lead radiochemistry development within the Division of Radiotherapy and Imaging.

To date, a variety of metal-based approaches have proved useful in illuminating the biochemical malfunctions that lead to neurodegeneration. These methods include: (1) positron emission tomography (PET), (2) single-photon emission computed tomography (SPECT), (3) magnetic resonance imaging (MRI), and (4) luminescent imaging. This tutorial review will highlight metal-based imaging techniques used to understand aberrant biochemical processes involved in neurodegenerative diseases. Particular focus will be placed on a general overview of each imaging technique, the purpose of each method, and its implementation by highlighting recent demonstrations in the literature.

Positron emission tomography (PET) & single photon emission computed tomography (SPECT)

Major advances in scanner technology over the last 20 years have facilitated adoption of PET and SPECT in the biomedicalimaging field. PET/SPECT-based imaging requires the use of a radiopharmaceutical or radiotracer. Typically, this is a molecule containing a radioisotope within its structure. For PET, the radiotracer contains a positron-emitting isotope. The emitted positron (β^+) or "anti-electron" collides with nearby electrons resulting in an annihilation process that produces two γ -rays at exactly 511 keV emitted 180° apart. The specific energy of these



Xiao-Peng He

Xiao-Peng He is professor at Feringa Nobel Prize Scientists Research Center, School of Chemistry and Molecular Engineering, ECUST. He obtained his BSc (2006) and PhD (2011) from ECUST. He completed a co-tutored doctoral program at ENS Cachan (France) (2008 to 2009) and postdoctoral research with Kaixian Chen (SIMM, CAS) from 2011 to 2013 at ECUST.

 γ -rays is dissimilar to other modes of decay possible for most radionuclides. The PET derived γ -rays (511 keV) are detected by specialized instrumentation and used to form the image for diagnosis. SPECT utilizes a similar approach wherein the radiotracer contains a γ-ray emitting short-lived isotope (ideal γ -ray energies: 100–250 keV), which is detected to provide the image for diagnosis. The required dosage for PET/SPECT is commonly within picomolar (pM; 10^{-12} mol L⁻¹) to nanomolar (nM; 10⁻⁹ mol L⁻¹) concentrations reducing any potential toxicology risk arising from radioactive isotope and nonradioactive "carrier" material.

The requirement for highly specialized and expensive instruments had previously limited the widespread adoption of PET. Due to early adoption and cheaper instrumentation, more hospitals are equipped with SPECT scanners. As a result, SPECT is more widespread, both in terms of clinical use and current imaging research. However, modern PET setups permit remarkable sensitivity compared to the analogous magnetic resonance imaging (MRI) and X-ray computer tomography (CT) approach, and arguably provides a slight advantage over SPECT-based methods. This has resulted in a shift towards PET-based technology in recent years.

For a PET-based imaging agent to achieve the spatial resolution needed for use in neurological or small animal imaging, a radioisotope with low energy positron emissions is required. The use of such isotopes enables the production of higher resolution images, i.e., lower positron energies correlate to a shorter positron distance giving a more specific location. Tables 1 and 2 highlight the current radioisotopes that have been used towards the development of radiotracers for PET/ SPECT imaging in neurodegenerative diseases with corresponding decay energies, half-lives, and production methods. It is important to note, each radioisotope may be subject to one or more decay modes; this can require additional consideration when designing a PET/SPECT radiotracer. The reader is directed to an excellent review by Blower for a more extensive list of radioisotopes used in medical imaging and therapy.²

The PET/SPECT-based imaging section will be divided according to each metal-based radioisotope with attendant consideration of the neurodegenerative disease-related biomarker target. For use of other radioisotopes and non-metal-based



He Tian

He Tian was born in China in 1962. In 1999, he was appointed Cheung Kong Distinguished Professor by the Education Ministry of China. In 2011, he was selected as a member of the Chinese Academy of Science. He has been listed by Thomson a Highly Cited Reuters as Researcher in Chemistry (2014) and Materials Science (2014-2019).



Jonathan L. Sessler

Jonathan L. Sessler received a BSc degree in Chemistry in 1977 from University of California, Berkeley. He obtained his PhD from Stanford University in 1982. After postdoctoral stays Strasbourg and Kyoto, he accepted a position as an Assistant Professor of Chemistry at the University of Texas at Austin, where he is currently the Doherty-Welch Chair.

Table 1 Radioisotopes currently used in PET-based imaging of neurodegenerative disease related biomarkers³

Tutorial Review

$t_{1/2}$	Production methods	Energy (β^+) – MeV	Energy (γ) – MeV	Decay type
20 min	Cyclotron	1.0	_	β^+
110 min	Cyclotron	0.6	_	$\beta^+ \ \beta^+$
9.7 min	⁶² Zn/ ⁶² Cu	2.9	0.12	β^+/γ
	generator			
12.7 h	Cyclotron	0.7	0.13	β^+ , β^- , γ , EC
1.1 h	⁶⁸ Ge/ ⁶⁸ Ga	1.9	0.11	β^+ , β^- , γ , EC β^+ , γ , EC
78.5 h	generator Cyclotron	0.9	_	β ⁺ , EC
	20 min 110 min 9.7 min 12.7 h 1.1 h	$t_{1/2}$ methods 20 min Cyclotron 110 min Cyclotron 9.7 min $^{62}Zn/^{62}Cu$ generator 12.7 h Cyclotron 1.1 h $^{68}Ge/^{68}Ga$ generator	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$

Table 2 Current radioisotopes used in SPECT-based imaging towards neurodegenerative disease related biomarkers

Radioisotope	$t_{1/2}$ (h)	Production methods	γ-ray – keV	Decay type
^{99m} Tc	6	^{99m} Tc/ ⁹⁹ Mo generator	141	γ
¹²³ I	13	Cyclotron	159	γ, EC
¹¹¹ In	67.9	Cyclotron	245	γ, EC

systems for PET/SPECT-based imaging and therapy the reader is directed to a number of excellent reviews.^{2,4}

The development of metal-based radiotracers requires knowledge of metal coordination chemistry and appropriate ligand design. Moreover, the radiotracer itself must:

- Have a high selectivity and binding affinity for the desired target with rapid binding kinetics.
- Target a specific receptor or biochemical process occurring in sufficient concentration so as to provide a good signal-tonoise ratio.
- Permit rapid insertion of the metal radioisotope and produce a complex that does not result in the loss of binding affinity to the biological target. Protocols that allow for metal complexation in aqueous conditions are particularly desired since they may allow for the facile development of "kit-based technology" for use in a clinical setting.
- The imaging agent or administration technique must allow for sufficient localization within the brain, i.e., penetrate the blood brain barrier (BBB). This remains a challenge when designing CNS agents.

Copper (60Cu, 61Cu, 62Cu, 64Cu) - PET

Copper (Cu) has two key biologically relevant oxidation states, Cu⁺ and Cu²⁺. Several decades of research from the inorganic chemistry community have resulted in a number of effective chelators able to form stable complexes with both Cu oxidation states. The imaging community has largely relied on Cu²⁺ systems containing mixed donors (e.g., N and S) with coordination numbers (CNs) of 4-6. To date, a number of Cu positronemitting isotopes have found utility in PET-based imaging. These include: 60 Cu ($t_{1/2}$ = 20 min), 62 Cu ($t_{1/2}$ = 9.7 min), 61 Cu $(t_{1/2} = 3.4 \text{ h})$, and ⁶⁴Cu $(t_{1/2} = 12.7 \text{ h})$. Copper-62 is an attractive option despite having a relatively short half-life ($t_{1/2} = 9.7 \text{ min}$) as it can be readily produced via an on-site generator. In contrast, the other Cu isotopes are currently produced from

solid targets at cyclotron facilities. Copper-64 ($t_{1/2}$ = 12.7 h) has a longer half-life and has been used extensively to radiolabel a range of small molecules, antibodies, proteins, and nanoparticles.⁵ The low energy positron emission of ⁶⁴Cu also enables the production of high-quality images comparable to the gold standard ¹⁸F PET (Table 1). To date, [¹⁸F]FDG (2-[¹⁸F]fluoro-2-deoxy-Dglucose) is the only PET imaging agent that has seen widespread clinical use. It has been extensively used as an imaging agent for studying glucose metabolism in the brain and identifying potential neurological disorders through reduced glucose usage.4 In the following section, we have highlighted select strategies implementing the Cu²⁺ radioisotopes for the imaging of neurodegenerative disease-related biomarkers. The reader is also directed to an excellent review by Cui and co-workers for more examples of Cu2+-based radiotracers explored in the context of neurodegenerative disease.5

To be viable as clinical candidates, metal-based imaging agents must have the appropriate physicochemical properties $(e.g., \log P)$ so as to balance the need for cell permeability and water solubility. Optimizing this features often requires planned synthetic efforts and an ability to derivatize easily the core metalchelating scaffold. Early efforts to develop Cu-based PET agents focused on square planar metal complexes, particularly $Cu(\pi)$ (diacetyl-bis(N^4 -methylthiosemicarbazone)) [Cu^{2+} (atsm)] and $Cu(\pi)(glyoxal-bis(N^4-methyl-3-thiosemicarbazonato)) [Cu^{2+}(gtsm)]$ (Fig. 1).

Seminal efforts by Yokoyama and co-workers demonstrated that $Cu(\pi)$ (diacetyl-bis(N^4 -methylthiosemicarbazone)) Cu(atsm)can be utilized as a PET radiotracer for the imaging of hypoxia in brain and neck cancers.6 Cu(atsm) displayed good stability and blood brain barrier (BBB) permeability with attendant accumulation in regions of the brain suffering from oxidative stress. Under these latter conditions, Cu(atsm) was found to undergo a superoxide oxidase-type conversion wherein superoxide $(O_2^{\bullet-})$ is oxidized to molecular oxygen (O_2) and the Cu^{2+} center is reduced to Cu⁺.7 Taking advantage of this redox chemistry, Yoneda and co-workers applied 62Cu(atsm) to image the striatum brain region (i.e., the motor and reward system) in patients with Parkinson's disease (PD).6 High accumulation of 62Cu(atsm) was seen in the striatum of the PD patients compared to the cerebral cortex. This finding was considered consistent with a localized enhancement of oxidative stress due to an increased concentration of superoxide. These efforts by Yoneda and associates are credited with inspiring further imaging research targeting the aberrant oxidative and nitrative processes associated with neurodegeneration.

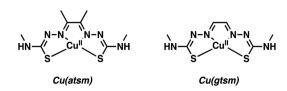


Fig. 1 Cu-Based $Cu(II)(diacetyl-bis(N^4$ agents methylthiosemicarbazone)) [Cu(atsm)] and Cu(π)(glyoxal-bis(N^4 -methyl-3-thiosemicarbazonato)) [Cu(qtsm)]

Scheme 1 Intracellular reduction of Cu²⁺ (glyoxal-bis(N⁴-methyl-3thiosemicarbazonato)) [Cu(gtsm)], which leads to the release of free Cu⁺ ions

Continued interest in Cu²⁺ systems has provided an impetus to develop Cu(atsm) analogues with stimuli-specific properties. Blower and co-workers recently developed an elegant method utilizing Cu(gtsm), the more easily reduced variant of Cu(atsm).8 The use of a glyoxal backbone, as found in gtsm, instead of the diacetyl in atsm, increased the reduction potential of the coordinated Cu2+ making it more susceptible towards intracellular reduction; $E_{1/2} = -0.44$ mV versus Ag/AgCl for Cu(gtsm) compared to $E_{1/2} = -0.60$ mV versus Ag/AgCl for Cu(atsm). In these studies, it was proposed that the Cu(gtsm) complex passes first through the cell membrane via a combination of passive diffusion and protein-carrier mechanisms. Subsequently, the Cu(gtsm) complex disassociates to yield free Cu⁺ (Scheme 1). In strictly operational terms, these efforts allowed for the PETbased imaging of free Cu⁺ ions in a transgenic AD mouse model and an initial assessment of the potential of Cu(gtsm) to act as a useful diagnostic biomarker.9

Notably, 64Cu(gtsm) was able to penetrate the BBB and increase 64Cu concentrations in the brain. PET imaging revealed statistically significant differences in ⁶⁴Cu global brain uptake and clearance, as well as in the brain regional ⁶⁴Cu concentration between AD transgenic mice and age-matched controls.8 These findings were among the first to show that ⁶⁴Cu could be used as an imaging tool to track changes in the trafficking of free Cu ions in the context of neurodegenerative disease studies.

Free metal ions, such as Cu²⁺, Fe^{2+/3+}, Mg²⁺, Mn²⁺, and Zn²⁺ are known to induce aggregation of amyloid-beta (Aβ) peptide and support formation of AB plaques, a key pathological biomarker of AD. 10 Imaging tools enabling the detection of AB aggregates are thus of great interest for understanding AD progression. Non-metal based PET imaging of AB plaques is possible through the use of radiolabelled benzothiazole and stilbene derivatives, [18F]florbetapir, [18F]flutemetamol,

Fig. 2 Non-metal-based PET imaging agents for the visualisation of Aβ plaques in vivo.

and [18F]florbetaben (Fig. 2).5,11 However, 18F radiotracers (as well as those based on ¹¹C) have a number of disadvantages, such as relatively short half-lives ($[^{18}F]t_{1/2} = 110 \text{ min}$; $[^{11}C]t_{1/2} =$ 20 min) that require late stage synthetic incorporation of the radiolabel, which is typically produced on site in highly specialized facilities. The development of metal-based strategies that rely on longer lived radionuclides and the insertion of a metal radioisotope into a ligand-conjugate structure with a predetermined high target affinity and selectivity are thus attractive.

Donnelly and co-workers have recently reported an elegant demonstration of ⁶⁴Cu-based PET imaging of Aβ plaques. They did this by coupling the AB targeting motifs, benzothiazole and dimethylaminostyrenyl, to Cu(atsm) to generate complexes CuL₁-CuL₃ (Fig. 3). 11 The resultant complexes proved stable in aqueous solutions over a range of pH and avoided the unwanted release of Cu. The authors noted that CuL₁ displayed poor aqueous solubility. The majority of their studies thus focused on CuL2 and CuL3. Co-localization experiments with immunostained Aß plaques provided evidence that CuL2 and CuL₃ would bind effectively to Aβ plaques in post-mortem human brain tissue (Fig. 4).

Subsequent radiolabelling experiments demonstrated marked differences between the aqueous soluble 64CuL2 and -L3, in which 64CuL2 produced only a weak PET signal in the brain after administration, a finding interpreted in terms of poor brain uptake. However, with CuL3, the complex incorporating an appended N,N-dimethylaminoethylene unit, a marked increase in BBB penetration was inferred. Biodistribution experiments

Fig. 3 Cu(atsm) hybrids functionalised with the benzothiazole and dimethylaminostyrenyl targeting Aβ ligands highlighted in blue.

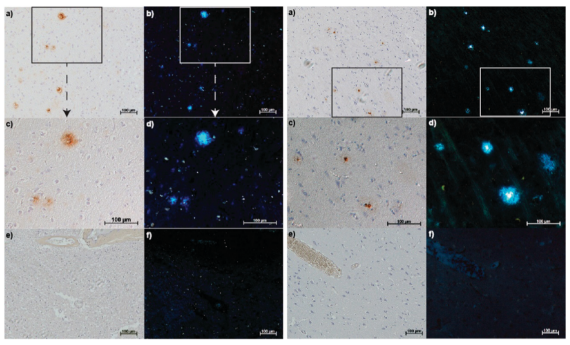


Fig. 4 AD human brain sections with 1E8 antibody stained Aβ plaques and fluorescence microscopy of CuL₂ (left) and CuL₃ (right) binding to Aβ plaques. Reproduced with permission from (J. Am. Chem. Soc., 2013, 135, 16120-16132). Copyright (2019) American Chemical Society.

in normal mice showed CuL3 had excellent brain uptake: 1.11% ID g⁻¹ at 2 min post injection, and excellent washout with limited accumulation, 0.38% ID g^{-1} at 30 min. This work highlights some of the challenges associated with developing imaging agents that can effectively penetrate the BBB.

More recently, Mirica and co-workers reported a series of ⁶⁴Cu-PET agents (TACNL₁ and PPL₂₋₅) for the imaging of Aβ plaques.¹² These imaging agents consisted of a 1,4,7triazacylononane (TACN) or 2,11-diaza[3.3](2,6)-pyridinophane (PP) ligands conjugated to 2-phenyl-benzothiazole, which resembles the well-known Aβ-binding dye Thioflavin T (ThT) (Fig. 5). The corresponding Cu complexes were shown to exhibit low nanomolar affinity for AB aggregates in vitro and ex vivo with the best binding system proving to be with PPL2. Radiolabelling experiments were then carried out under aqueous conditions (45 °C, 20 minutes, 0.1 M NH₄OAc, pH 7). Radiochemical yields of greater than 95% were obtained for L₁-L₅. The resultant complexes were then used for imaging studies without

additional purification. Biodistribution studies demonstrated promising brain uptake in wild-type mice followed by rapid clearance. In vivo PET imaging experiments in AD (Tg2576) transgenic mice revealed radiotracer accumulation in the neck and head for ⁶⁴Cu-PPL₂, ⁶⁴Cu-PPL₄, and ⁶⁴Cu-PPL₅. In contrast, no appreciable brain uptake was seen in the case of 64Cu-TACNL1. The highest brain uptake and tracer accumulation was seen in the case of ⁶⁴Cu-PPL₂, a finding that mirrored what was seen in the *in vitro* Aβ experiments.

Efforts with Cu-based PET agents have provided insight into the design criteria that need to be addressed to obtain an effective in vivo imaging agent. In the case of the Donnelly and Mirica studies summarised above, synthetic modification of the chelating motif and judicious choice of targeting unit were crucial in endowing the most effective system with the appropriate properties, particularly BBB penetration. However, continued efforts will be required to evaluate further the potential of these promising candidates as markers for AD. We also believe that additional effort focused on creating next

Chem Soc Rev

 PPL_2 ; R = HTACNL₁ PPL₅ PPL_3 ; R = Me PPL_4 ; $R = CH_2CO_2H$ > 9.0 Coronal

Fig. 5 (top) Structures of Cu chelators (TACNL₁ and PPL₂₋₅) that are conjugated to a series of Aβ ligands, targeting moieties (highlighted in blue) and (bottom) *in vivo* PET imaging experiments in AD (Tg2576) transgenic mice carried out using 64 Cu-TACNL₁. 64 Cu-PPL₂, 64 Cu-PPL₄, and 64 Cu-PPL₅. The best uptake was seen for 64 Cu-PPL₂ with little or no brain uptake being seen in the case of 64 Cu-TACNL₁. Adapted with permission from (*J. Am. Chem. Soc.*, 2017, **139**, 36, 12550–12558, https://pubs.acs.org/doi/10.1021/jacs.7b05937). Copyright (2019) American Chemical Society.

generation Cu-based PET imaging will allow the scope of utility to be expanded to include other neurodegenerative diseases (e.g., Huntington's, ALS, etc.). That researchers will be able to capitalize upon the plethora of Cu ligands found within the bioinorganic and inorganic communities is expected to facilitate these efforts and codify Cu-based PET as a tool to understand better and diagnose more effectively CNS neurodegeneration.

Gallium (⁶⁸Ga) – PET

Gallium-68 (68 Ga, $t_{1/2}$ = 1.13 h)-based PET radiotracers are becoming attractive as imaging agents owing to the ready availability of gallium-68 from a 68 Ge/ 68 Ga generator and an ideal half-life, which complements the biodistribution times

typically seen for small molecules. However, widespread adoption of ⁶⁸Ga-based PET radiotracers has been slow due to concerns associated with the high energy positron emission characteristic of gallium-68 and the resulting high levels of radioactivity (Table 1). However, these concerns are balanced by the commercial availability of ⁶⁸Ge/⁶⁸Ga generators, leading us to predict that gallium-68 radiotracers will continue to find utility in the clinic.

0.0

The most prevalent oxidation state of gallium is the charge dense Ga³⁺ ion, which forms complexes of CN 4-6. Typical Ga³⁺ complexes are based on the use of hexadentate ligands containing multiple hard anionic donors, such as 1,4,7,10-tetra-azacyclododecane-1,4,7,10-tetraacetic acid (DOTA), 1,4,7-triazacyclononane-1,4,7-triaceticacid (NOTA), and diethylene-triaminepentaacetic acid (DTPA) (Fig. 6).⁵ The metallation of these ligands often requires relatively harsh conditions

Fig. 6 Chemical structures of ligands commonly used to prepare complexes of Ga^{3+} .

(elevated temperatures, prolonged reaction times) to achieve Ga^{3+} chelation. Alternative chelators have thus been developed, allowing for improved radiolabelling conditions. In this section, we highlight select examples using gallium-68 for the development of PET-based imaging agents that are designed to target neurodegenerative disease.

Seeking to take advantage of the properties of gallium-68 for use in PET imaging and drawing inspiration from previous small molecule AB aggregate probe structure-activity relationship studies (Fig. 2), Saji et al. developed a benzofuran-Ga-DOTA conjugate displaying a high affinity towards AB aggregates in vitro and in vivo (Fig. 7).13 Here, the radiolabelled benzofuran-Ga-DOTA conjugate was isolated in 26% yield (>99% purity) by heating the ligand and Ga(NO₃)₃ in an ammonium acetate buffer (pH 5.8) at 100 °C for 10 min. In vitro binding assays provided support for the conclusion that appending a ⁶⁸Ga-DOTA metal complex to the benzofuran targeting moiety had little effect on the binding affinity towards $A\beta_{42}$ aggregates $(K_i = 10.8 \text{ nM})$. Subsequent in vitro staining of brain tissue revealed effective binding to AB plaques in a tissue section derived from an AD mouse model (Tg2576). Unfortunately, biodistribution experiments in a normal mouse displayed low initial uptake (0.45% ID g^{-1}) in the brain at 2 min post-injection. Currently, it is an open question as to whether additional synthetic modifications will allow for brain uptake. Nevertheless,

Fig. 7 Benzofuran-Ga-DOTA conjugate reported by Saji and co-workers The benzofuran A β targeting ligand is highlighted in blue.

Fig. 8 A multi-component complex, Ga–DTPA chalcone, reported by Mishra and co-workers. The chalcone A β targeting ligand is highlighted in blue.

these results serve to illustrate a simple strategy for developing ⁶⁸Ga complexes that bind to Aβ aggregates *in vitro*.

Mishra and co-workers reported a ⁶⁸Ga chelator-targeting system obtained by conjugating the Aß binding dimethylamino chalcone motif to DTPA (Fig. 8).14 In this case, rapid ⁶⁸Ga³⁺ insertion was achieved in acidic sodium acetate buffer within 10 min (65-70 °C, pH 4.5-5.0). In vitro binding assays demonstrated a high binding affinity towards AB aggregates $(K_i = 4.18 \text{ nM})$, as monitored by the formation of the fluorescent chalcone-bound Aß aggregates. In particular, upon presumed binding, a blue shift and a 5.5-fold increase in fluorescence emission intensity ascribed to the Ga-DTPAchalcone complex was observed. This study demonstrated the advantages of a dualmodal PET and fluorescence imaging system as a more refined technique in disease imaging. Animal studies and biodistribution analysis of Ga-DTPA_{chalcone} revealed BBB penetration with brain uptake of 1.24% ID g⁻¹ at 2 minutes post injection and rapid washout with negligible activity (0.36% ID g^{-1}) after 30 minutes. No further biological experiments were performed. Thus, it is difficult at present to assess fully the potential of this PET imaging agent. However, the work as reported does serve to underscore a potentially exciting new approach to developing PET imaging probes for AD.

Asti and co-workers reported a 68 Ga-labeled curcumin complex for the imaging of A β plaques (Fig. 9). Curcumin is a phytocompound commonly used in traditional Eastern medicines to which salutary antioxidant, anti-inflammatory, antimicrobial, and anticancer activities have been ascribed. 15 The relative planarity of curcumin endows this natural product with an inherent affinity towards A β plaques. In the Asti study, curcumin and two synthesised curcumin derivatives, diacetyl-curcumin (DAC) and bis(dihydroxy)curcumin (bDHC), were used to form 2:1 complexes with 68 Ga $^{3+}$ in good-to-excellent yields with a short reaction time when heated at 100 $^{\circ}$ C in a sodium acetate buffer (pH 5). *In vitro* studies revealed moderate to high affinities for A β plaques. The inherent fluorescence properties of these complexes allowed their cellular uptake to

OMe
$$X = H, OH, OAc$$

$$X = CI$$

Fig. 9 Ga-curcumin complex for the imaging of A β plaques reported by Asti and co-workers. The curcumin A β targeting ligand is highlighted in blue.

be monitored readily in the A549 human lung cancer cell line. No further biological experiments were carried out; however, this research underscores the potential that ⁶⁸Ga-curcumin-based systems may hold as potential diagnostic tools in the area of neurodegenerative disease imaging.

Zirconium (89Zr) – PET

Zirconium-89 (89 Zr) has become one of the most popular PET radioisotopes for use with biological targeting/therapeutic agents due to its long half-life ($t_{1/2}$ = 78.4 h). This has enabled the use of 89 Zr monoclonal antibodies (mAb) conjugates (typical biological half-lives = 7 days) for the development of "immuno-PET" radiotracers. 16 The use of immuno-PET radiotracers has allowed for more precise targeting of disease-related biomarkers, such as specific neurotoxic protein aggregate conformations. This specificity is expected to provide new insights into neurodegenerative disease and guide the development of treatment protocols that could be individualised from patient to patient. 5

The high charge and small radius of Zr⁴⁺ (CN 4–9) forces a preference toward polyanionic hard donor chelators, such as ethylenediaminetetraacetic acid (EDTA) and diethylenetriaminepentaacetic acid (DTPA). However, the well-known iron (Fe) chelator desferrioxamine B (DFO) has also found utility in *in vivo* zirconium-89 PET-based applications (Fig. 10).

Fissers *et al.* have recently reported studies detailing the development of a 89 Zr immuno-PET strategy for the targeted imaging of A β plaques in AD (Fig. 11). This work relies on the use of the bifunctional chelator *p*-isothiocyanatobenzyl

Fig. 10 Chemical structures of commonly used ⁸⁹Zr chelators, ethylene-diaminetetraacetic acid (EDTA) and desferrioxamine B (Df).

Fig. 11 DFO-based ⁸⁹Zr immuno-PET imaging agent developed by Fissers *et al.* with the goal of achieving enhanced $A\beta$ plaque imaging. The monoclonal antibody (mAb) $A\beta$ plaque targeting unit is highlighted in blue.

[89Zr]DFO-JRF/AβN/25

desferrioxamine (Df-Bz-NCS), a system routinely used for the rapid and efficient preparation of Zr-89-labeled mAbs. Taking advantage of the isothiocyanate moiety, Df-Bz-NCS was coupled to the amino residue of the antibody JRF/AβN/25, an N-terminal end-specific mAb targeting the full-length human Aβ epitope. Complexation of the resulting construct, Df-Bz-JRF/AβN/25, with 89Zr was successfully achieved under aqueous conditions (HEPES buffer) at room temperature to afford [89Zr]Df-Bz-JRF/ A β N/25 in an average yield of 48.39 \pm 12.33%. An in vitro binding assay provided evidence of a high binding affinity towards $A\beta_{40}$ ($K_d = 0.85$ nM) with no non-specific binding being observed. This high binding affinity was further validated by in vitro autoradiography studies on brain sections from AD Tg mice models. No obvious binding was observed in brain sections of normal mice. [89Zr]Df-Bz-JRF/AβN/25 demonstrated good in vivo stability (i.e., up to 48 hours post injection). [⁸⁹Zr]Df-Bz-JRF/AβN/25 demonstrated the ability to penetrate the BBB resulting in modest brain uptake in normal mice, thus showing promise as an ⁸⁹Zr immuno-PET strategy. Unfortunately, no further biological experiments were carried out. Moreover, in vitro studies revealed instability in plasma.

Further studies within the context of developing Zr-based imaging systems should focus on functionalising the best extant systems to improve brain uptake, as well as the synthesis and study of new Zr chelators with improved physicochemical properties. Different mice models should also be screened to allow the utility of these systems across a range of neurodegenerative disease indications.

Technetium (99mTc) – SPECT

The invention of ^{99m}Tc/⁹⁹Mo generators revolutionised nuclear medicine, facilitating the production of the technitium-99m radioisotope and lead to its daily availability in major hospitals. Not surprisingly, this led to a surge in the development of ^{99m}Tc-based radiotracers and use of ^{99m}Tc as a popular choice for the imaging of a range of diseases. One of the key attributes of ^{99m}Tc-based radiotracers, which has led to the predominance of this isotope in the field of nuclear medicine, is the ability to synthesise a range of technitium-99m complexes *via* a one-pot, one-step "kit" process. This simplicity meant that many sterile ^{99m}Tc-based radiotracers could be produced daily with minimal technologist input or without costly equipment.

None of the isotopes of Tc are stable enough for routine bench chemistry. However, the similar ionic radius of rhenium (Re) to that of Tc has led to its use as a cold-laboratory (nonradioactive) surrogate for Tc-based complexes. On this basis, various core Tc ion complexes, such as: 99mTc(v)O3+, $CpTc(v)(CO)_3^{3+}$, $Tc(v)N^{2+}$, $Tc(u)(CO)_3^{3+}$, and $Tc(i)(CNR)_6^{+}$, have been characterised to date.5 One of the most commonly used ligands is the N₂S₂ donor, bis-aminoethanethiol (BAT); it forms a complex with 99mTc(vi)O3+ that is stable in vivo. For more indepth information on the history and various biological targets of 99mTc SPECT imaging agents, the reader is directed to the excellent review by Blower.2 Analogous to the metal complextargeting conjugate systems used in PET, motifs capable of detecting specific biomarkers in neurodegeneration have been developed utilising [99mTc]-SPECT conjugates. In this section, we highlight selected examples of technetium-99m-based systems being explored as SPECT imaging agents for neurodegenerative disease.

An important application of SPECT-based imaging agents has been towards understanding the role of the dopamine transporter (DAT) in a number of neurological processes.

Fig. 12 Chemical structure of cocaine, a well-known DAT antagonist, and TRODAT-1, developed as a potential DAT SPECT imaging agent. The tropane core targeting unit is highlighted in blue.

Catecholamine neurotransmitters are crucial for maintaining homeostasis throughout the autonomic nervous system. Altered DAT activity has been shown in a number of neurological disorders, with a reduction in DAT being observed in postmortem examinations of patients with PD and AD. Patients with PD have also been shown to have a selective loss of dopamine neurons in the basal ganglia and substantia nigra.¹⁸

DAT antagonists that display a high binding affinity, such as cocaine, are generally constructed around a tropane scaffold (Fig. 12). Influential efforts by Kung and co-workers utilized the tropane motif to design [99mTc]TRODAT-1, a BAT-tropane conjugate (Fig. 12). [99mTc]TRODAT-1 demonstrated nanomolar binding affinity towards DAT in rat striatal homogenate $(K_i = 8.7 \text{ nM})$. Ex vivo autoradiography results further confirmed high uptake and retention in the striatal region. These promising results led to the study of [99mTc]TRODAT-1 in normal healthy human subjects. The radiotracer was found to localize in the basal ganglia, a finding consistent with selective DAT binding. Dosimetry studies confirmed that [99mTc]TRODAT-1 was safe for administration to humans and that it could be used in imaging DAT for the diagnosis of PD.¹⁹ However, [99mTc]TRODAT-1 has now been largely supplanted by the alternative iodine-123 imaging agent, [123I]FP-β-CIT (DaTSCAN), due to a world-wide shortage of the parent isotope, ⁹⁹Mo, needed to produce ^{99m}Tc.⁴

In continuation of their efforts involving the use of labelled benzofuran compounds for AB detection, Saji and co-workers reported three [99mTc]-pyridyl benzofuran complexes (Fig. 13).²⁰ The ^{99m}Tc and corresponding Re complexes were synthesized using BAT as the chelating ligand. In vitro assays provided evidence of strong affinity towards $A\beta_{42}$ aggregates ($K_i = 13.6$ -149.6 nM) for the Re complexes. Biodistribution analyses of [99mTc]BAT-Bp-2 in normal mice (ddy mice) revealed brain uptake post injection, $1.80\%~{
m ID}~{
m g}^{-1}$ at 2 minutes, with moderate

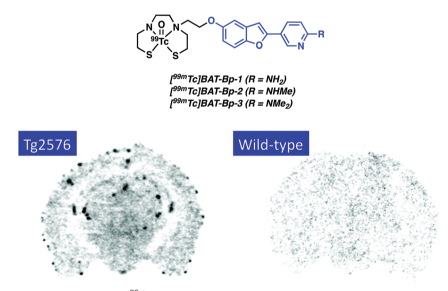


Fig. 13 The [99m Tc]-pyridyl benzofuran complexes ([99m Tc]BAT-Bp-n, where n = 1-3) developed by Saji and co-workers. The 2-pyridylbenzofuran- A β targeting ligands are highlighted in blue. Ex vivo autoradiography with [99mTc]BAT-Bp-2 illustrating labelling of Aß plagues in sections of brain tissue from an AD transgenic (Tg2576) mouse versus age-matched controls (wild-type). Reprinted (adapted) with permission from (J. Med. Chem., 2012, 55, 2279-2286). Copyright (2019) American Chemical Society

washout, 0.79% ID g^{-1} in the brain at 60 minutes, that was deemed sufficient to encourage further study. Furthermore, ex vivo autoradiography with [99mTc]BAT-Bp-2 revealed substantial labelling of AB plaques in sections of brain tissue from an AD transgenic (Tg2576) mouse versus age-matched controls (Fig. 13).

Zhang and co-workers later reported a structural variant of the BAT conjugate containing a 2-arylbenzothiazole moiety. 21 A competitive binding assay identified seven Re complexes with potent binding affinity toward $A\beta_{42}$ aggregates ($K_i < 50$ nM). Radiolabelling with 99mTc yielded no significant differences in the binding of the [99mTc]BAT conjugate probes with Aβ₄₂ plaques. [99mTc]BAT-benzothiazole conjugate 1 displayed ideal pharmacokinetics with a brain uptake of 2.11% ID g⁻¹ at 2 min and good washout of 0.62% ID g⁻¹ at 60 min in normal mice. Subsequent ex vivo autoradiography experiments revealed extensive labelling of AB plaques in brain tissue from AD Tg mice versus age-matched healthy controls. SPECT imaging studies in non-human primates (rhesus monkeys) were also performed. The semiquantitative data resulting from these studies provided support for the notion that [99mTc]BATbenzothiazole (1) penetrates the BBB of healthy rhesus monkeys (Fig. 14).

Despite metal-based SPECT imaging agents having several advantages, [99mTc]TRODAT-1 is among a select few agents that

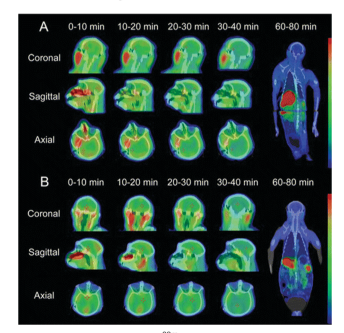


Fig. 14 Chemical structure of [99mTc]BAT-benzothiazole conjugate 1 reported by Zhang and co-workers with the benzofuran targeting moiety highlighted in blue. SPECT images of [99mTc]1 in rhesus monkeys (A: 4 years old, male; B: 27 years old, female); SPECT images are overlaid on individual co-registered CT images. Reprinted (adapted) with permission from (Bioconjugate Chem., 2016, 10, 2493-2504). Copyright (2019) American Chemical Society

have actually been used for the clinical diagnosis of neurodegenerative disease. However, as summarised above, a number of promising agents have been reported that underscore how [99mTc]-based systems may have an increased future role in creating improved SPECT imaging agents for CNS disorders. However, balanced against promise are the current supply problems associated with the required parent isotope, 99Mo.

Indium (In) - SPECT

Indium-111 is an isotope characterized by a long half-life ($t_{1/2}$ = 67.3 h, γ-rays emission - 245 keV). It is thus attractive for conjugation to targeting biomolecules with slow pharmacokinetics, such as mAb. Indium-111 has been the radioisotope of choice for this purpose since the 1980s and was the first radioisotope to be commercially and clinically approved for bioconjugation. Similar to Ga³⁺, the predominant oxidation state of indium under most aqueous conditions is In³⁺. Polyanionic, hard donor chelators, such as EDTA, DTPA, and DOTA, have been the ligands of choice for forming In-complexes with good aqueous stability (Fig. 6 and 10).

The use of 111 In-based SPECT to study neurodegenerative disease has focused on attempts to address issues with BBB penetration and lack of in vivo binding seen with other small-molecule and mAb conjugate systems. For instance, Pardridge et al. reported an Aβ-peptide based radiotracer, which was designed to deposit ¹¹¹In at pre-existing Aβ plaques. The Pardridge system was constructed by conjugating biotin at the N-terminus of AB peptide and attaching DTPA chelator subunits to key lysine residues (Lys16 or Lys28) (Fig. 15).22 Ex vivo autoradiography studies provided support for the expectation that significant binding to AB plaques from brain sections of patients with AD could be achieved. However, additional quantitative binding studies and in vivo kinetic analysis were not reported. Despite indium-111 being well established in the field of immuno-based imaging, the rise of 89Zr as a more readily available and longer-lived surrogate is seeing 111 In-based SPECT agents being increasingly replaced by their 89Zr-based immuno-PET counterparts.

Magnetic resonance imaging (MRI) in neurodegenerative disease

Optimization of imaging instruments and the availability of a number of small molecule probes, known as contrast agents (CAs)

[111 In]A β_{40} -biotin

Fig. 15 Aβ-peptide based ¹¹¹In radiotracer developed by Pardridge et al. The biotin targeting unit is highlighted in blue.

has made magnetic resonance imaging (MRI) a powerful technique within the imaging paradigm. In comparison to PET and SPECT, MRI is generally cheaper, more widely accessible, and provides higher spatial and temporal resolution (>0.01-1 mm and > 0.01-1 s) with enhanced (total) depth penetration through biological tissue. These properties make MRI an excellent tool for in vivo studies and clinical use. A major advantage of MRI over radiotracer (PET/SPECT) approaches is that MRI CA signals can be modulated by environmental changes or via exposure to specific analytes, allowing for realtime functional imaging (i.e., fMRI) in living subjects. However, as a general rule MRI lacks the sensitivity seen for PET and SPECT. Recent efforts within the MR imaging community have focused on the development of tools that enable the imaging of neurodegenerative diseases.²³ For extensive coverage of the theory and development of MRI contrast agents, current directions, and challenges for the field, the reader is directed to two excellent reviews by Meade²³ and Caravan.²⁴

The most common contrast agents for MRI utilize a polydentate ligand with a coordinated paramagnetic metal ion (e.g., Gd^{3+} , $Fe^{2+/3+}$, and $Mn^{2+/3+}$). The operational mechanism for MRI involves administration of the contrast agent followed by the application of a short radiofrequency (RF) pulse to a sample (e.g., human, mouse, or test-tube) placed within an external magnetic field (i.e., the scanner). This RF pulse results in the excitation of the nuclear spins of surrounding water protons, which then relax to the equilibrium of the external field. The rates of the relaxation, based upon the longitudinal (T_1) and transverse (T_2) time constants, are dependent upon the specific environment of the water molecules. It is the variation of these rates that forms the basis of the signal intensity in typical anatomical and contrast-enhanced MRI.

Paramagnetic metal ions, such as Gd^{3+} , $Fe^{2+/3+}$, and $Mn^{2+/3+}$, contain unpaired electrons. These ions generate magnetic moments that interact with nearby water molecules to increase proton nuclear relaxation rates. The extent to which a contrast agent can change the T_1 or T_2 of water is generally defined by relaxivity, r_1 or r_2 , respectively. Relaxivity is the change in bulk relaxation rate per unit of contrast agent, normally in units of mM^{-1} s⁻¹, and is defined by eqn (1). Briefly, $(1/T_i^0)$ is the relaxation rate in the absence of the contrast agent, $(1/T_i)$ is the relaxation rate in the presence of contrast agent, and [CA] is the concentration of the MRI contrast agent.

$$\frac{1}{T_i} = \frac{1}{T_i^0} + r_i[CA]; \ i = 1, 2$$
 (1)

As a general rule, the greater the relaxivity (r_i) of a contrast agent, the better the CA increases the rate of relaxation of nearby protons. This enhancement generally manifests as an improved resolution contrast image. Additional changes to the contrast image can also occur depending upon the local environment. Indeed, the relaxation times of endogenous water molecules vary upon interacting with macromolecules (e.g., proteins and DNA) generating a contrast difference in T_1 - and T_2 -weighted images compared to standard conditions.

Due to the physical processes underlying relaxation rates, T_2 is always shorter than T_1 . The T_2 is thus more difficult to

modulate via the addition of contrast agents. In the case of standard Gd^{3+} contrast agents with relatively similar r_1 and r_2 values, the slower T_1 will display a larger percent modulation than the faster T_2 . Traditionally, small molecule CAs have been used to increase the signal change in the T_1 -weighted image (Fig. 16a). For transverse relaxation of these small molecule probes, the observed T_2 or T_2^* is shorter than the true T_2 due to field inhomogeneities and other factors. Efforts to generate meaningful changes in T_2 have focused on the use of nanoparticle systems. The resulting systems are referred to as T_2 , or T_2^* MRI contrast agents; they generally lead to a decrease in the signal of T_2/T_2^* -weighted images.

For T_1 -weighted contrast agents, three main factors determine the relaxivity of the agent:

- \bullet q average number of water molecules bound directly to the paramagnetic metal centre.
- $\tau_{\rm m}$ mean residence lifetime of the bound water molecule, *i.e.*, the inverse of the water exchange rate ($k_{\rm ex}$).
- \bullet $\tau_{\rm r}$ rotational correlation time, *i.e.*, how rapidly the molecule tumbles (rotates) in solution.

In the case of parameter q, water molecules in the first coordination sphere of the metal complex experience the greatest T_1 enhancement. Thus, the larger the q-value the higher the r_1 value (Fig. 16b). For τ_m , an exchange rate optimised to the timescale of relaxation will generate the highest relaxation enhancement. To date, $\tau_{\rm m}$ remains difficult to predict, measure, and implement in CA design (Fig. 16c). However, manipulation of τ_r is a commonly employed strategy and achieved by increasing the molecular weight of the MRI contrast agent (e.g., by binding to a protein, as larger molecules tend to tumble and rotate much slower) (Fig. 16d). For T_1 contrast agents, changes in τ_r have minimal effects at higher field strengths (>3 T), limiting use of such systems in high-field preclinical and ultra-modern clinical scanners. For T_2 -weighted agents, changes in tumbling rates are still an effective way to modulate contrast at any field strength. Aggregation remains the primary method for functional imaging with T_2 -probes. Alternative methods, such as chemical exchange saturation transfer (CEST) and paramagnetically-enhanced chemical shift imaging, are available, but due to the limited applications in neurodegenerative disease imaging they will not be covered in this review. For further details on applications of these techniques, we refer the reader to the aforementioned reviews.23,24

The MRI imaging section will be divided by each paramagnetic metal ions Gd^{3+} , $Fe^{2+/3+}$, and $Mn^{2+/3+}$ with focus on the precise method and biomarker relevant to neurodegenerative diseases. These examples will demonstrate both targeted and functional approaches.

Gadolinium (Gd³⁺) - MRI

 Gd^{3+} has the largest number of unpaired electrons for an aqueous stable ion (S = 7/2), a high available coordination number (CN 7–10), and close-to-ideal electronic relaxation rate that optimises its ability to increase the relaxation rate of water

Chem Soc Rev

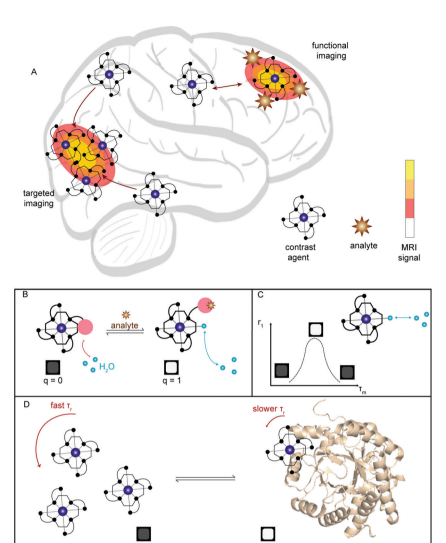


Fig. 16 (A) Simplified illustration showing the two main ways in which MRI has been applied towards neurodegenerative imaging; targeted imaging by selective accumulation of contrast agent and functional imaging, where the signal is modulated by presence of an analyte. (B–D) Illustration of the common methods for developing functional T_1 contrast agents: (B) modulation of bound water, q; (C) optimisation of water exchange rate, τ_m ; (D) binding to a higher molecular weight conjugate to alter rotational correlation time, τ_r .

protons through simple water–metal interactions.²⁴ However, free Gd³⁺ metal ions display off-target toxicity, requiring the use of a tight binding chelator. Polydentate and hard anionic donors, such as those found in diethylenetriaminepentaacetic acid (DTPA) and 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA), remain the ligands-of-choice for Gd-based MRI contrast agents in academic research and clinical use. Due to its higher thermal stability and worries of systemic toxicity resulting from leaching of Gd³⁺ from linear chelators (*e.g.*, DTPA), the macrocyclic DOTA is now preferred for new applications, as illustrated throughout this MRI section.²³

MRI-based imaging of CNS disorders has focused on methods of targeted (MRI) or functional imaging (fMRI). Recent efforts towards targeted imaging have utilized metal complex-targeting conjugates analogous to PET/SPECT. An exemplary demonstration of such efforts was recently reported by DeBeof and co-workers using arecoline- and tropane-based

Gd-DOTA metal complexes for the selective MRI imaging of DAT.²⁵ DeBeof and associates recognized that small molecule binding within a macromolecular biomarker would yield an enhancement in T_1 via modulation of the molecular rotation (τ_r) . The tropane-derived CA-1 (Fig. 17) produced a 38% increase in the r_1 value from 6.32 mM⁻¹ s⁻¹ to 8.71 mM⁻¹ s⁻¹ in the presence of DAT-transfected HEK293 cells at a 7 T magnetic field. Binding studies with tropane-derived CA-2 revealed a high binding affinity to DAT, similar to that of the DAT antagonist cocaine (303 nM). Unfortunately, due to the relative insensitivity of MRI, which typically requires $> 10 \mu M$ detectable concentration of contrast agent, and DAT only being present at nM concentrations in vivo, these probes were not pursued further in the context of potential biological applications. However, this work provided an elegant demonstration of combining targeted molecules with manipulation of τ_r towards the detection of neurodegenerative biomarkers.

Tutorial Review

Chemical structures of tropane and arecoline MRI-analogues, CA-1 and CA-2.

A similar Gd-DOTA-targeting moiety conjugate strategy was recently used by Wang and co-workers to effect the MR imaging of myelin.²⁶ Myelination is a fundamental biological process within the nervous system, which involves the formation of myelin sheaths along neuronal axons for rapid and efficient conduction of nerve impulses. Abnormalities or destruction of myelin results in the neurodegenerative disorder Multiple Sclerosis (MS). The ability to visualise and discriminate between healthy and abnormal myelin in vivo could provide insight into this disease and mechanisms behind its progression.

In an attempt to visualize myelination with MRI, Wang and co-workers developed a first-generation conjugate that was prepared by appending a known myelin selective coumarinbased fluorescent probe to Gd-DOTA to form the MRI contrast agent, myelin dye-1 (cf. Fig. 18). Ex vivo imaging of partially demyelinated mouse and rat brain tissue stained with myelin dye-1 revealed discrimination between regions that were high and low in myelin. Subsequent MRI T_1 mapping of brain tissue complemented the fluorescence data, and provided support for the suggestion that retention of myelin dye-1 is proportional to the level of myelination in the brain. Due to the polar nature of myelin dye-1, intra-cerebroventricular infusion was required to bypass the BBB and enable in vivo imaging of demyelinated lesions in L-α-lysophosphatidylcholine (LPC) treated rats. MRI mapping yielded relatively long T_1 values in demyelinated lesions, a result ascribed to little or no binding of myelin dye-1 in these latter samples. These efforts highlight the potential of MRI contrast agents that can selectively bind to myelin in vivo and underscores the promise they may have in terms of facilitating early diagnoses of MS and monitoring treatments aimed at myelin repair.

Analogous to the PET and SPECT conjugate literature, Ferroud and co-workers designed a series of Gd3+-based MRI conjugated contrast agents comprising of an AB aggregate targeting moiety, a linker, and a Gd3+ chelator (Fig. 19).27 Ferroud et al. carried out a structure activity relationship (SAR) study on 16 different contrast agents, varying the Gd³⁺ chelator from DOTA to PCTA (3,6,9,15-tetraazabicyclo[9.3.1]pentadeca-1(15),11,13-triene-3,6,9-triacetic acid), the Aβ targeting ligand between 2-arylbenzothiazole, 2-arylbenzoxazole, and various stilbene derivatives, and the charge of the linker from neutral to overall positive or negative while likewise adjusting its length. The resultant complexes showed good water solubility with nanomolar binding affinities to AB aggregates. However, they failed to pass through an intact (healthy) BBB. This shortcoming again highlights the significant challenges associated with developing effective imaging agents that have both high affinity for a neurodegenerative biomarker and the ability to penetrate the BBB.

DOTA-type macrocycles ligate to eight of the nine available Gd³⁺ coordination sites, leaving one site available for an inner sphere water molecule.23 The Meade group has exploited this free binding site by pioneering the development of *q*-modulated MRI contrast agents. Such systems are designed to only allow inner-sphere access of water upon binding to the target analyte (biomarker). A seminal contribution by Meade and co-workers was the development of a calcium ion (Ca2+) responsive MRI agent, in which the relaxivity was controlled by the reversible binding of Ca2+ ions.28

Ca²⁺ ions play a vital role in maintaining proper cell function with deviations in neuronal Ca2+ signalling being correlated with a number of neurodegenerative diseases. The Ca²⁺-responsive

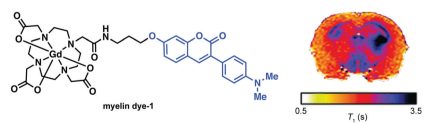


Fig. 18 (left) Chemical structure of myelin dye-1 with the coumarin targeting moiety highlighted in blue and (right) T_1 mapping of a Sprague-Dawley rat brain proximal after 5 h post injection. The dark regions (short T₁) are indicative of increased binding in areas of myelination. Reprinted (adapted) with permission from (J. Med. Chem., 2012, 55, 2279-2286). Copyright (2019) American Chemical Society

$$X = 0, S \qquad K_i \ge 1000 \text{ nM}$$

$$\text{Gd(III) chelate} \qquad \text{linker} \qquad \text{targeting moiety}$$

$$\text{Optimization}$$

$$\text{NH}_2$$

$$\text{NH}_2$$

$$\text{NH}_2$$

$$\text{NH}_2$$

$$\text{NH}_2$$

$$\text{NH}_2$$

$$\text{NH}_2$$

$$\text{NH}_3$$

$$\text{NH}_4$$

$$\text{NH}_4$$

$$\text{NH}_4$$

$$\text{NH}_5$$

$$\text{NH}_6$$

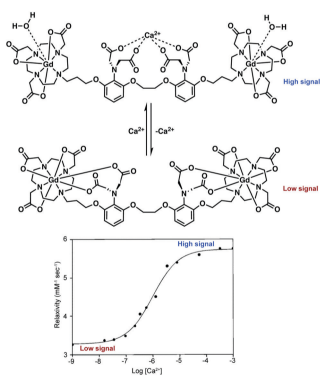
$$\text{NH}_6$$

$$\text{NH}_7$$

Fig. 19 Structural optimisation of a Gd-DOTA complex for the imaging of Aβ plaques

MRI contrast agent developed by Meade and co-workers incorporated a seven-coordinate DOTA analogue, 1,4,7-tris-(carboxymethyl)-1,4,7,10-tetraazacyclododecane (DOPTA), wherein q can vary from 0 to 2 and was linked to the 1,2-bis(o-aminophenoxy)ethane-N,N,N',N' tetraacetic acid (BAPTA) Ca²⁺ binding subunit via a propyl chain (Scheme 2). In the absence of Ca2+, the aromatic iminoacetates of the BAPTA moiety interact with the Gd³⁺ centre. The presence of Ca²⁺ modulated the binding of the aromatic iminoacetates such that they interacted with the Ca2+ ions, rather than the Gd³⁺ centre. The resulting exposure of the Gd³⁺ centre to water molecules caused a change in relaxivity. Thus a stimulus-responsive (Gd-DOPTA)₂-Ca²⁺ sensor was produced that was found to have high sensitivity toward Ca2+ (detection range: 0.1 to 10 µM). This MR-based sensor was found to display a 1:1 stoichiometry for the Ca²⁺ ion triggers and produce an 80% overall increase in relaxivity upon Ca²⁺ complexation. This work stands as an excellent initial example of how to manipulate various parameters (e.g., q) to enact an effective change in T_1 signal. Unfortunately, application of these agents in the context of neurodegenerative disease has been limited due to difficulties in tuning the calcium affinity and MRI sensitivity to match the relative intracellular (0–100 μM) and extracellular (0.1-2 mM) calcium fluctuations typically seen in the brain. Applications may also prove limited by an apparent lack of cell permeability.

Chang and co-workers developed a class of Cu⁺ responsive *q*-modulated MRI contrast agents as potential tools for studying copper dysregulation in neurodegenerative diseases.²⁹ The general scaffold was a Gd–DOTA analogue featuring a thioether-rich conjugate (Scheme 3). In the absence of Cu⁺ ions, the nitrogen on the pyridine ring unit coordinates the Gd³⁺ centre, restricting inner sphere water coordination. Upon addition of Cu⁺, the pyridine nitrogen and thioether moiety undergoes a conformational change,



Scheme 2 $\,q$ -Modulating Gd-DOPTA-BAPTA contrast reported by Meade and coworkers that permits the MR-monitored reversible detection of Ca²⁺ ions. Upon binding of Ca²⁺, water access to the Gd³⁺ ion is increased, raising the relaxivity and MRI signal of the contrast agent. The relationship between relaxivity and Ca²⁺ concentration is displayed in the lower plot. Reprinted (adapted) with permission from (*J. Am. Chem. Soc.*, 1999, **121**, 1413–1414). Copyright (2019) American Chemical Society.

a chemical switching event that serves to "turn off" the Gd-pyridine binding interaction. The increased water access

$$Cu^{+}$$
 Cu^{+}
 C

Scheme 3 Chang's q-modulating Gd³⁺-DO3A contrast agent that permits the reversible detection of Cu²⁺ ions via MRI.

to the Gd3+ centre leads to a marked enhancement in the relaxivity (r_1) , with the largest overall change in r_1 (360%; from 1.5 to 6.9 mM^{-1} s⁻¹) being seen in the case of compound *CG2*.

In order to improve cellular uptake, Chang and coworkers included an octa-arginine unit to yield Arg₈CG2 (Fig. 20).²⁹ This system was applied towards imaging a Menkes disease model fibroblast cell line and control fibroblasts. Menkes disease is a disorder characterized by a progressive deterioration of the nervous system. In comparison to the untagged CG2, the arginine tagged complex showed a remarkable 9-fold greater accumulation in both cell lines. The overall relaxivity change of Arg₈CG2 was slightly lower than with CG2, but gave a higher absolute r_1 for both the 'off' and 'on' states (3.9 to 12.5 mM⁻¹ s⁻¹). However, it should be noted that the buffer system was different for these two studies. While a higher absolute signal will lower the detection threshold, a larger relative change between the bound and unbound conditions should lead to greater sensitivity in determining analyte binding. Phantom MR images of the two cell lines revealed a higher contrast when Arg₈CG2 was present than in its absence. Treatment of the cells with a copper chelator allowed monitoring of the depletion of intracellular copper through MRI monitoring, indicating the potential of such an approach for in vivo biological MRI studies.

Modification of Gd³⁺-based contrast agents has also yielded functional systems for monitoring enzymatic function (e.g., glutamate decarboxylase (GAD) activity). GAD is an essential enzyme converting L-glutamate (L-Glu), an excitatory neurotransmitter, to

γ-aminobutyric acid (GABA), a major inhibitory neurotransmitter. Abnormal variations in GABA-mediated signalling have been linked to a number of neuropathological conditions, such as PD, epilepsy, and various psychiatric disorders. Traditional methods for GAD identification and quantification are based on postmortem examinations. The development of non-invasive agents that would allow GAD activity to be monitored in vivo would thus be highly desirable; ultimately, they might make it possible to correlate precise deviations in activity with specific neurodegenerative phenotypes.

Hoehn and co-workers recently reported a q-modulated MRI contrast agent activated by GAD.30 The system was constructed from Gd-DO3A conjugated to flexible linkers with pendant glutamate residues (Fig. 21). GAD-mediated decarboxylation of the pendant glutamates unveiled the Gd³⁺ centre allowing inner sphere water coordination. This increase in the hydration state of the Gd^{3+} ion (q = 0 to 2) resulted in enhanced relaxivity with attendant complexation to endogenous macromolecules via increased electrostatic interactions. The Gd-DO3A-glutamate conjugate-1 was used to achieve the in vivo imaging of implanted GABAergic neurons that were differentiated from embryonic stem cells. Faster relaxation was observed for the differentiated neurons relative to undifferentiated cells in vitro and in vivo, a result considered consistent with GAD-specific activation of the contrast agent in those neurons.

Manganese (Mn²⁺ and Mn³⁺) - MRI

Health concerns regarding the use of Gd-based contrast agents have become a focal point in MRI research. Researchers are thus actively searching for non-Gd-based alternatives.24 Manganese (Mn^{2+/3+}) has shown promise as a viable alternative with relatively low inherent toxicity. Manganese is an essential metal ion in biological systems. Indeed, various inherent biochemical mechanisms enable the uptake and cell internalisation of Mn within the brain.31 Even the simple MnCl2 salt has found use in the manganese-enhanced MRI (MEMRI) visualisation of brain architectures, neuronal tracts, and brain activity through T_1 and T_2 imaging (Fig. 22). However, the relaxivity of Mn²⁺ is low compared to Gd3+. As a result, significant amounts

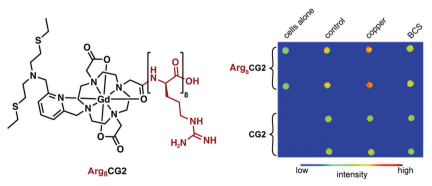


Fig. 20 Chemical structure of Arg₈CG2 and phantom images of both Arg₈CG2 and CG2 in two different cell lines, revealing an increase in signal intensity in the presence of copper. Reproduced from (Chem. Sci., 2012, 3, 1829-1834) with permission from The Royal Society of Chemistry.

Fig. 21 g-Modulated MRI contrast agent reported by Hoehn and co-workers. This system proved responsive towards glutamate decarboxylase (GAD), a finding rationalized in terms of the enzyme serving to cleave the terminal acid group and increase the relaxivity of the contrast agent as the result of an increase in the metal-bound water number, q

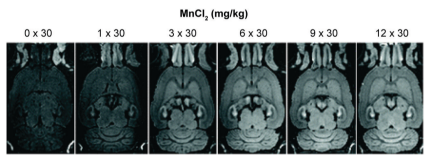


Fig. 22 Example of manganese-enhanced MRI (MEMRI), showing an increased signal intensity throughout the brain with increasing concentration of MnCl₂. Reprinted (adapted) with permission from (Schizophr. Bull., 2008, 34, 595-604). Copyright (2019) Oxford University Press.

of Mn²⁺ are required to produce an effective MRI signal. This can become a limitation since the Mn2+ concentrations in question can produce toxic effects, at least in the case of the

In an attempt to address the toxicity of free Mn²⁺ at high concentrations, Flesch and co-workers developed a firstgeneration dendritic DTPA-based Mn²⁺ complex. The polymeric network acted as a time-release vehicle for the progressive delivery of Mn²⁺ in the brain (Fig. 23).³² This dendrimeric Mn²⁺ complex displayed excellent aqueous solubility (0.1 M) with minimal in vitro neuronal toxicity. For instance, a 100% rat survival rate was observed when concentrations as high as 20 mM were given via intraperitoneal (i.p.) administration; presumably, these favourable findings reflect the slow diffusion of the released Mn²⁺ cation in vivo. The obtained in vivo images in the context of this study were also consistent with what was seen when MnCl2 was injected intravenously. These efforts serve to outline what might emerge as an effective strategy to overcome the unwanted toxicity of free Mn2+ at the concentrations needed to effect the MEMRI-based visualisation of brain structures.

To explore imaging beyond MEMRI and compete with current Gd3+-based contrast agents, stable small molecule Mn contrast agents must be developed.²⁴ Jasanoff and co-workers recently

Fig. 23 A dendritic DTPA-based complex reported by Flesch and coworkers that was designed to improved delivery of Mn²⁺ while decreasing toxic side effects.

reported the preparation of three in vivo stable, cell-permeable phenylenediamido (PDA)-based Mn³⁺ complexes and showed

that they could be used as T_1 MRI contrast agents for selective cell labelling (Fig. 24).33 These Mn3+ complexes displayed a higher stability compared to Mn²⁺ complexes due to favourable chelation between the hard donor, anionic PDA ligand, and the high-spin 3d⁴ Mn³⁺ cation. The open coordination sphere of the PDA-based Mn(III) complexes also facilitated exchange with inner-sphere water molecules. Subsequent studies focused on modification of the ligand scaffold with a BAPTA Ca²⁺ chelating unit. These efforts afforded an effective T_1 MRI contrast agent that allowed for the monitoring of intracellular Ca2+ concentrations (Mn-L3-BAPTA). Facile modification of the BAPTA scaffold allowed this system to be tuned to produce an analogue, ManICS1-AM, that displayed good cell permeability. Upon cellular uptake in HEK293 cells, ManICS1-AM undergoes esterase-mediated cleavage to reveal the BAPTA and enable monitoring of intracellular Ca²⁺ concentrations. *In vivo* studies provided support for the design expectation that ManICS1-AM could be used to monitor changes in Ca²⁺ concentrations induced via stimulation with potassium ions (K⁺). This latter stimulation induces neuronal cell depolarization and gives rise to changes in the Ca²⁺ concentrations in brain tissue. ManICS1-AM apparently constitutes the first example wherein calcium fluctuations in a living rat brain could be monitored directly at the intracellular level via fMRI.

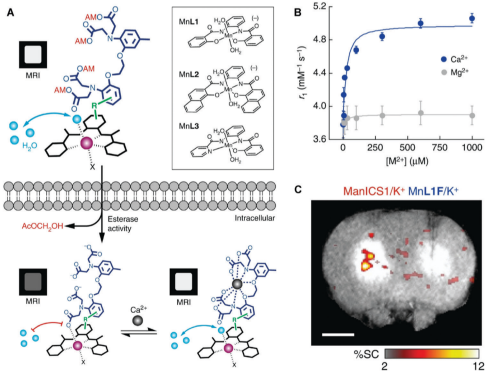


Fig. 24 Phenylenediamido (PDA)-based Mn^{3+} complexes reported by Jasanoff and co-workers. These complexes act as T_1 MRI contrast agents and permit the imaging of intracellular and in vivo Ca²⁺ concentrations. (A) When the cell permeable AM-ester protected complex (ManICS1) is internalised in cells, esterases cleave the AM-ester groups to release the active complex, which is cell permeable and trapped. Binding of Ca²⁺ causes an increase in signal intensity due to increased water access to the Mn³⁺ centre. Early iterations of cell permeable Mn³⁺ complexes are shown in the inset. (B) The relaxivity response of ManICS1 to increasing Ca²⁺ concentration, in comparison to the similar divalent cation, Mg²⁺. (C) In vivo signal change in a rat model upon stimulation with potassium, which results in increased neural activity and a spike in intracellular calcium levels. A calcium-insensitive analogue (MnL1F) give rise to no detectable response under otherwise identical experimental conditions. Reproduced from (Nat. Commun., 2019, 10, DOI: 10.1038/s41467-019-08558-7) - Open Access.

Iron ($Fe^{2+/3+}$) – MRI

Chem Soc Rev

Iron, able to form in vivo stable metal complexes in both its divalent and trivalent oxidation states, is also being explored actively as a relaxation agent for use in MRI.24 In fact, one of the most widespread uses of Fe in MRI involves the use of endogenous haemoglobin (Hb). The difference between paramagnetic deoxygenated Hb and diamagnetic oxygenated Hb can be observed in T2*-weighted images and permits a functional readout through blood oxygenation level-dependent (BOLD) fMRI (Fig. 25). This method provides real-time information on blood oxygenation, blood flow, and blood volume changes. Data gathered through BOLD fMRI can be used to obtain indirect insights into the underlying neural activity, as activation of neurons requires an increased usage of oxygen and blood flow to the firing cells. This technique has shown promise as a biomarker for Alzheimer's disease and other neurodegenerative disorders.³⁴ However, BOLD fMRI remains limited by the indirect coupling between neural activity and vascular response.

Currently, synthetic iron-based contrast agents are being explored as an alternative to BOLD. In an attempt to gain direct information on neuronal activity, Jasanoff and co-workers recently reported an Fe3+-based MRI strategy that relies on a directed evolution approach utilising bacterial cytochrome P450-BM3 to afford a series of protein-based MRI contrast agents that permitted dopamine imaging. Dopamine is a monoamine neurotransmitter vital for proper neurological function.³⁵ Therefore, real time imaging of dopamine activity in vivo is particularly desirable as malfunction of the dopaminergic system has been associated with several neurodegenerative diseases. Dopamine is a substrate (endogenous ligand) for the bacterial cytochrome P450-BM3. However, in the absence of this substrate, a water molecule can fill the remaining coordination site of the Fe3+ atom. Subsequent binding of dopamine results in an attenuated MRI signal (0.83 \pm $0.01 \text{ mM}^{-1} \text{ s}^{-1} \rightarrow 0.10 \pm 0.00 \text{ mM}^{-1} \text{ s}^{-1}$). This change in signal could be used to achieve the spatiotemporal imaging of dopamine release from PC12 cells and in the ventral striatum of brains in live rats following reward-related lateral hypothalamic brain stimulation.35 This seminal work by Jasanoff

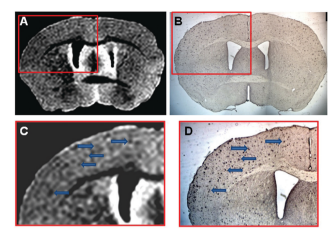


Fig. 26 Ex vivo μMRI, at the level of the area cinguli of the cerebral cortex, of a 14 month-old APP/PS1 Tg injected with USPIO-PEG-Aβ1-42, showing numerous dark spots indicating location of Aβ plaques. Reproduced from (*PLoS One*, 2013, **8**, DOI: 10.1371/journal.pone.0057097) – Open Access.

and co-workers established a new paradigm in fMRI and demonstrated directly how it could be applied to the problem of mapping neuronal function.

For MRI contrast applications, iron is commonly used in the form of T_2 superparamagnetic iron oxide nanoparticles (SPIONs). Wisniewski and co-workers employed bis-functionalised ultrasmall SPIO (USPIO) nanoparticles with sufficient BBB permeability for the targeted imaging of A β plaques. These USPIO nanoparticles were constructed *via* conjugation to the A β_{42} peptide through a polyethylene glycol (PEG) linker. *In vivo* and *ex vivo* imaging of the resulting USPIO-PEG-A β_{42} nanoparticles revealed an ability to image A β plaques *in vivo* by means of T_2 *-weighted MRI on the μ m scale. These imaging-based conclusions were supported by histological analyses (Fig. 26).

Jasanoff and co-workers have recently reported an elegant SPION-based imaging strategy for the detection of extracellular Ca²⁺ fluctuations that exploits the Ca²⁺ dependent interactions between the calcium-binding protein moiety C2AB and a lipid-coated iron oxide nanoparticle (LCIO).³⁷ In the presence of Ca²⁺, the magnetic calcium-responsive nanoparticles (MaCaReNa) undergo clustering which serves to enhance the protein–LCIO

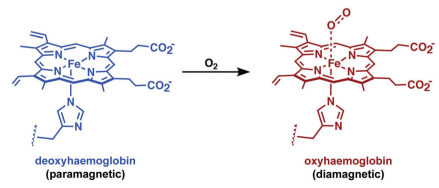


Fig. 25 The chemistry underlying blood oxygenation level-dependent (BOLD) functional MRI (fMRI). Oxygenation of haemoglobin causes an decrease in paramagnetism. Neural activity leads to changes in blood flow/oxygenation levels in nearby vessels, which leads to signal changes in MRI.

interactions. Structural changes occur (i.e., aggregation) that lead to an increase in the hydrodynamic diameter from 35 \pm 1 nm in the low calcium regime to 262 \pm 14 nm in the presence of high calcium concentrations. An almost 2-fold change in r_2 accompanies these changes in the Ca²⁺ concentrations and considered most relevant to extracellular studies (i.e., 0.1 to 1 mM). The possibility of studying brain activation in response to chemical (K⁺) and electrical stimuli in vivo was then explored using rats. Observed MRI response times were on the order of seconds. This work by Jasanoff and co-workers is arguably the first example of molecular fMRI using an MRI calcium sensor in a living brain model.

Luminescence-based imaging agents for the detection of neurodegenerative disease related biomarkers

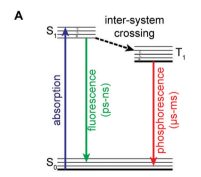
Advances in the chemistry of luminescent small molecules and coordination complexes have provided systems that may be used for luminescence-based imaging, either alone or in conjunction with PET, SPECT, or MRI techniques. Luminescence is defined as the emission of light from a substance (luminophore) in its electronically excited state. Upon exciting the luminescent optical imaging agent, the corresponding emission signal can be used to ascertain the location and concentration of the luminophore within a specific biological sample. The resultant emission can be characterised by its wavelength, intensity, and the type of emission - Fig. 26A (i.e., length of the luminescence lifetime; fluorescence < 10 ns versus phosphorescence $\sim \mu s$ to ms). A number of optical imaging agents have been designed that may be used to selectively detect an analyte of interest, either through a binding event or by means of a chemical reaction. In comparison to MRI and PET/SPECT, these luminescent optical agents display unrivalled sensitivity, specificity, and temporal and spatial resolution. Optical imaging agents have thus begun to find use in the direct quantification of specific biomarkers involved in physiological or pathological processes in real time at the cellular and subcellular levels. However, due to the limited depth of light penetration through

tissue, luminescence-based imaging is currently restricted to small live animals for research purposes only.

To achieve optical-based bioimaging, researchers have utilised luminescent imaging agents in the form of organic compounds, transition metal complexes, lanthanide chelates, quantum dots, nanoparticles, and fluorescent proteins. 38 In the context of reviewing efforts to image and understand better CNS neurodegeneration, it is convenient to divide the general class of metal-based luminescent imaging agents into those based on transition metal complexes and those that rely on lanthanide cations.

Transition metal-based imaging agents

The photophysical properties of transition metal coordination complexes with luminescent properties have been extensively detailed within the inorganic and material science literature. These complexes often display large Stokes shifts, high photochemical stability, and tuneable chromatic properties achievable via ligand modification. Transition-metal complexes applied to luminescence-based imaging include: Re+, Ru2+, Os2+, Ir3+, Rh3+, Pt²⁺, Au⁺, and Cu⁺. These systems all exhibit spin-forbidden triplet-state phosphorescence involving electronic transitions from the metal ion and ligand (i.e., metal-to-ligand chargetransfer (MLCT) and ligand-to-metal charge-transfer (LMCT)). Intra-ligand (IL) and ligand-to-ligand charge-transfer (LLCT) can also occur and are the result of intra- and inter-ligand transitions, respectively. Many complexes are subject to metal-metal interactions or ligand-ligand stacking that favour triplet-state phosphorescence, dominated by metal-metal-to-ligand charge-transfer (MMLCT), ligand-to-metal-metal charge-transfer (LMMCT), and metal-to-ligand-ligand charge-transfer (MLLCT). As a result of the spin-forbidden nature of the associated relaxation from the triplet back to the ground state long-lived luminescence emission lifetimes are observed for many of these complexes. Researchers have developed various photoluminescence time-resolved techniques, such as time-gated luminescence and photoluminescence lifetime imaging microscopy (TGL and PLIM), that permit discrimination relative to other interfering fluorescent signals with similar wavelengths (Fig. 27).38 This represents a significant advantage for



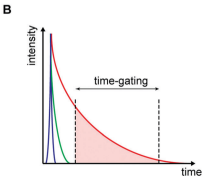


Fig. 27 (A) Simplified Jablonksi diagram illustrating the difference between fluorescence and phosphorescence, with representative emission lifetimes. (B) Illustration of the principle behind time-gated imaging and spectroscopy. By applying a time-gate, the excitation pulse (blue) and shorter-lived fluorescence (green) can be removed from the signal to allow selective detection of long-lived phosphorescence (red).

metal-based luminescent species compared to, for instance, traditional organic fluorophores. On the other hand, as a result of the numerous electronic transitions involving both the metal and the ligand, the excited states of transition metal complexes are often sensitive to changes in the surrounding environment and the nature of the coordinating ligand. These combined set of attributes makes transition metal-based luminescence-based imaging agents attractive for the selective detection of analytes. both in vitro and in vivo.

For an extensive coverage of time-resolved photoluminescence techniques and the development of transition-metal complexes for bioimaging applications, the reader is directed to two significant reviews by Huang and Li. 38,39 In this section, we present selected examples of transition metal-based luminescent imaging agents (Ir3+, Ru2+, and Re+) that have been used to detect biomarkers implicated in the development of neurodegenerative disease.

Iridium (Ir³⁺)

Octahedral cyclometallated iridium (Ir3+) complexes remain a current gold-standard in the area of luminescence. They are typically characterized by high photoluminescence efficiencies, good aqueous stability, and excellent colour tunability through synthetic modification of the coordinating ligand. To date, a large number of cyclometallated Ir3+-based complexes have been developed for the imaging of disease-specific biomarkers in vitro and in vivo.38

As underscored by the imaging agent development efforts summarised in the earlier portions of this review, the so-called amyloid hypothesis, focused on the role of AB plaques in AD progression, remains a focal point in AD research. Within this overall paradigm, researchers have set out recently to develop dual modal tools, so-called theranostics, that are able to detect Aß peptide and inhibit Aß plaque formation. Ma and co-workers recently reported a series of Ir3+ and Rh3+ metal

complexes capable of inhibiting amyloid fibrillogenesis while permitting the attendant luminescence detection of AB fibrils.40 These metal complexes are comprised of two cyclometallated (2-phenyl pyridine, C^N) ligands with two bound solvent molecules (i.e., H2O) (Fig. 28). The labile solvent molecules allow for coordination to the histidine residues (H6, H13, or H14) present within the Aβ fibril framework. A brief structure activity relationship analysis led to the conclusion that Ir^{3+} and Rh³⁺ metal complexes bearing C^N phenyl-pyridine (ppy) ligands (i.e., 2a and 2b) provided the greatest inhibition of AB aggregation.

In comparison to 2a, the Rh³⁺-complex 2b was found to be more active with near complete inhibition of AB peptide aggregation at 5 μM, demonstrating its potential as an Aβ peptide inhibitor for the treatment of AD. Unfortunately, 2b proved weakly emissive. On the other hand, the Ir³⁺-complexes 2a, 3, and 4 displayed luminescence turn-on in the presence of Aß peptide. This enabled the use of these latter complexes as tools to distinguish between the different states of aggregation, with a higher luminescence emission being observed for Aβ fibril versus Aβ monomers.

In follow up work, Ma and co-workers showed that appropriately designed Ir3+ complexes could function as dual AB fibrillogenesis inhibitors and luminescent imaging agents. 41 Toward this end, Ma et al. prepared twelve different luminescent Ir³⁺ complexes and evaluated their ability to bind to the AB peptide. It was found that complex 5, comprised of a 2-phenyl-1*H*-imidazo[4,5-*f*][1,10]phenanthroline (phenyl-imidazo-phen) (N^N) ligand, was effective in distinguishing between Aβ fibrils over Aß monomers. Continued optimization yielded complexes 6 and 7, with 6 demonstrating partial inhibition and 7 demonstrating complete inhibition of AB fibrillogenesis (Fig. 29). Complex 6 provided for a 6- and 12-fold enhancement in luminescence emission intensity when contacted with AB monomers and fibrils, respectively. Complex 7 proved even more effective, providing an 11- and 18-fold luminescence emission enhancement upon treatment with AB monomers

Fig. 28 Octahedral cyclometallated iridium (lr^{3+}) complexes developed by Ma and co-workers for the simultaneous detection of A β peptide and inhibition of Aß plaque formation.

Fig. 29 Optimised octahedral cyclometallated iridium (Ir³⁺) phenyl-imidazo-phen-based complexes (5-7) developed by Ma and co-workers for the simultaneous detection of Aß peptide and inhibition of Aß plaque formation.

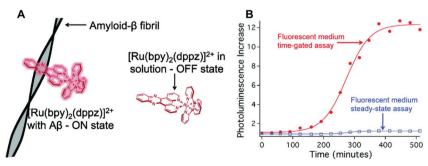
and fibrils, respectively. In vitro analysis of complex 7 revealed a neuroprotective effect against Aβ-induced cytotoxicity in SH-SY5Y cells and mouse primary cortical cells. The complexes generated by Ma and coworkers thus provide an early demonstration that it is possible to create theranostic systems, namely agents able to inhibit (therapeutic) and monitor (diagnostic) AB peptide formation via luminescence.

Ruthenium (Ru²⁺)

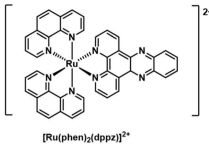
As true for Ir-systems, ruthenium (Ru2+) complexes have found extensive use in bioimaging applications. Recent efforts have been devoted to exploring cellular imaging and sensing applications wherein octahedral dipyridophenazine (dppz) Ru²⁺ complexes are used as molecular "light-switches". For instance, Marti and co-workers reported a Ru2+-based complex $[Ru(bpy)_2(dppz)]^{2+}$ (bpy = bipyridyl) that undergoes a photoswitching process in the presence of $A\beta$ aggregates (Fig. 30).⁴² Specifically, upon binding to A β fibrils, $[Ru(bpy)_2(dppz)]^{2+}$

displayed a 50-fold increase in luminescence intensity ("turn-on") as compared to the free Ru complex in aqueous solution. In the presence of monomeric AB only a minimal increase in luminescence intensity was observed. By exploiting the long emission lifetimes of [Ru(bpy)₂(dppz)]²⁺, time-resolved experiments enabled the monitoring of AB binding within a fluorescent environment. This work demonstrates the potential to overcome a major limitation associated with most current ThT-AB inhibition screening assays, since it shows that it is possible to use an appropriately chosen metal complex to screen potential AB inhibitors with inherent fluorescent properties.

PD is often characterized by the abnormal accumulation of Lewy body protein aggregates comprised primarily of the alpha-synuclein (αS) peptide. The role of αS in the development of PD has been studied with data showing a strong correlation between as misfolding and aggregation and the progression of PD pathogenesis. However, the direct cause of αS misfolding and aggregation as well as the role of as accumulation in the development of PD has yet to be elucidated. The development of an effective imaging agent that allows αS misfolding and



 $\textbf{Fig. 30} \hspace{0.2in} \textbf{(A) } \hspace{0.2in} \text{Ru}^{2+} - \text{Based dipyridophenazine (dppz) complex reported by Marti and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} \text{time-gated assay for the monitoring and co-workers.} \\ \textbf{(B) } \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+} \hspace{0.2in} [\text{Ru}(\text{bpy})_2(\text$ of Aß fibril formation in a fluorescent medium (rhodamine B). Reprinted (adapted) with permission from (J. Am. Chem. Soc., 2011, 133, 11121-11123). Copyright (2019) American Chemical Society



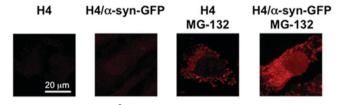


Fig. 31 [Ru(phen)₂dppz]²⁺ complex used by Marti and co-workers to monitor in real-time αS fibrillization in human neuroglioma cells (H4). Reprinted (adapted) with permission from (J. Am. Chem. Soc., 2012, 134, 20776-20782). Copyright (2019) American Chemical Society

aggregation progression to be monitored readily is thus viewed as being highly desirable.

Building on their earlier work summarised above, Marti and co-workers recently reported a 1,10-phenathroline (phen) supported Ru complex, [Ru(phen)2dppz]2+ and showed it could be used to monitor αS fibrillization. ⁴³ Upon binding to αS fibrils, an 18-fold increase in luminescence emission intensity was observed with a remarkable selectivity. [Ru(phen)2dppz]2+ also displayed good cell permeability with low toxicity in human neuroglioma H4 cells, as well as an ability to distinguish between healthy cells and cells overexpressing αS (H4-MG-132) (Fig. 31).

The Ru²⁺ complexes prepared by Marti and associates provide an excellent demonstration of an optical imaging agent. They retain the benefits of the ThT displacement assay currently used in AD research in that they are amenable to use in the highthroughput screening of potential therapeutics targeting αS fibrillization in PD. However, they may be employed under conditions complicated by the presence of a fluorescent background. In addition, these luminescent-based imaging systems may allow the general monitoring of protein aggregation in vitro and do so with high spatial and temporal resolution.

Rhenium (Re⁺)

Rhenium (Re⁺) tricarbonyl bipyridine [Re(N^N)(CO)₃X]ⁿ⁺ [X = anionic (n = 0) or neutral (n = 1)] complexes have foundextensive use in bioimaging applications.³⁹ As true for the [Ru(bpy)₂(dppz)]²⁺ complexes discussed immediately above, [Re(dppz)(CO)₃]⁺ also displays light-switching properties and is capable of detecting biologically important species. In continuation of their efforts on luminescent transition metal probes, Marti and co-workers reported that [Re(CO)₃(dppz)(Py)]⁺ could be used to effect the luminescent-based detection of Aß aggregates (Fig. 32).⁴⁴ [Re(CO)₃(dppz)(Py)]⁺ was shown to be non-emissive

[Re(CO)3(dppz)(py)]

Fig. 32 [Re(CO)₃(dppz)(Py)]⁺ complex Marti and co-workers that allows for the luminescence detection of AB aggregates. As noted in the main text, this system undergoes an unprecedented second light-switching event upon light irradiation.

in aqueous buffer. However, upon exposure to fibrillar Aβ, an instantaneous 18-fold increase in luminescence emission intensity was observed. Upon light irradiation for 6000 s in the presence of oxygen, the AB peptide underwent oxidation. The resultant modifications of the AB peptide induced an unprecedented second light-switching event with an attendant 105-fold increase in luminescence emission intensity. As used by Marti and coworkers, [Re(CO)₃(dppz)(Py)]⁺ provides an elegant demonstration of a luminescent-based imaging system that may be used to monitor AB plaque formation and to obtain insights into the local environment within the AB plaque that results from oxidation.

More recently, Pigge and co-workers reported a series of oligothiophene benzothiazole-based tetraarylethylene (TAE) Re(CO)₃ complexes (8–13) and showed that these systems were capable of monitoring Aβ aggregate formation in real time.⁴⁵ The benzothiazole TAE chelating ligands used in this study contain a different number of thiophene units, a variation that serves to modulate the photophysical properties and the binding affinities toward Aβ fibrils. In aqueous solution, complexes 8-10 displayed minimal luminescence emission intensity (Fig. 33). However, in the presence of AB aggregates, a significant increase in the luminescence emission intensity was observed. Complex 8 displayed the smallest response in the presence of aggregated A β fibrils (~13-fold), complex 9 exhibited the largest response with a 34-fold increase in luminescence intensity, and complex 10 demonstrated the largest Stokes shift (105 nm) with a red-shifted emission in the NIR region (λ_{em} = 641 nm). Complex 9 was also evaluated against

$$Ar = \begin{cases} Ar \\ S \\ OC \\ CO \\ CO \end{cases}$$

$$Ar = \begin{cases} Ar \\ S \\ S \end{cases}$$

$$Ar = \begin{cases} Ar \\ S \\ S \end{cases}$$

$$Ar = \begin{cases} Ar \\ S \\ S \end{cases}$$

$$Ar = \begin{cases} Ar \\ S \\ S \end{cases}$$

Fig. 33 Oligothiophene benzothiazole-based tetraarylethylene (TAE) Re(CO)₃ complexes (8-10) reported by Pigge and co-workers.

Fig. 34 Benzothiazole-based tetraarylethylene (TAE) $Re(CO)_3$ complexes (11–13).

other bio-macromolecules and displayed excellent selectivity towards the $A\beta$ peptide.

To confirm further the structural importance of both thiophene and benzothiazole functionalities, complexes **11–13** were synthesised (Fig. 34). In the case of complex **11**, each (oligo)thiophene moiety was replaced by a phenyl ring. In **12** and **13** the bis(bithiophene) units were retained, but the benzothiazole core was replaced by ones based on bis(2-pyridyl) or bis(2-quinolyl)Re(CO)₃ subunits, respectively. As anticipated, complexes **11–13** displayed only a modest increase in emission intensity in the presence of A β fibrils. The high binding affinities and desirable photophysical properties of the oligothiophene benzothiazole-based Re complexes **8–10** supports the contention that these systems may have a role to

play as fluorescent $A\beta$ probes, perhaps as an alternative to ThT. Additionally, the fact that this class of complexes is easy to modify leads us to suggest that they could inspire the development of new second-generation TAE-based diagnostic tools for investigating protein aggregation in AD and other neurological diseases.

Lanthanide-based imaging agents

Trivalent lanthanide ions (Ln3+) display unique intrinsic luminescence properties that arise from the f-f electron transitions. However, as transitions occurring within the same orbitals (4f-4f) are Laporte forbidden, Ln3+ ions display low absorption coefficients. This complicates the efficient harnessing of their luminescence properties. Free Ln3+ ions are also toxic, which makes them unsuitable for bioimaging applications per se. To circumvent toxicity and their poor absorptivity, Ln3+ ions are typically chelated by polyanionic macrocycles, such as DOTA and DTPA, and functionalised with various organic chromophores for use as light excitation "antennas". These "antennas" enable efficient Ln-based luminescence proceeding via excitation of the organic antennas to their respective S1 excited states, followed by intersystem crossing to their T1 states. Subsequent intramolecular charge transfer populates the excited state of the Ln³⁺ ion; relaxation of the ion to the ground state then produces a unique fingerprint emission characteristic of the Ln³⁺ ion in question (Fig. 35).

Unlike transition metal complexes, lanthanide-based emissions are not significantly affected by the surrounding environment or ligands. Indeed, even under widely different conditions, almost no variation is observed in the emission/excitation wavelengths of the Ln³⁺ ions. The lack of such interactions is attributed to the valence 4f orbitals being core-like with shielding from

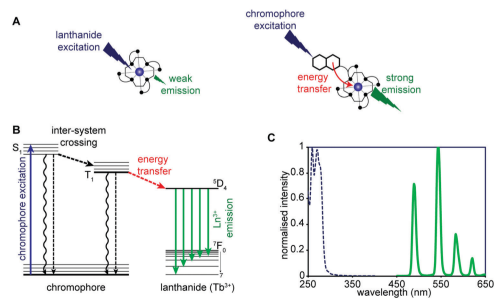


Fig. 35 (A) Sensitisation through a chromophore generally leads to more intense emission in comparison to direct excitation of the Ln^{3+} ion. (B) Simplified Jablonski diagram illustrating the sensitisation of Ln^{3+} ions, in this case Tb^{3+} . (C) A typical chromophore-sensitised Tb^{3+} emission spectrum (solid green trace), showing the large Stokes shift relative to the chromophore excitation spectra (dashed blue line) that is typically observed.

chemical bonding by the populated 5s and 5p orbitals. This feature, coupled with the spin and Laporte forbidden nature of these electronic transitions, yields long luminescence lifetimes in the micro- to millisecond range. Luminescent-based lanthanide imaging agents have thus found utility in time-gated or timeresolved live-cell and in vivo imaging. A significant advantage of these lanthanide based optical imaging agents is that varying the Ln³⁺ centre yields only minor changes in the chemical properties enabling multimodal and multiplexed applications. However, since each Ln3+ ion gives rise to unique luminescent features, in principle it is possible to generate several similar imaging agents based on the same organic framework.

While the wavelengths of Ln3+ transitions are typically fixed, it is possible to modulate the emission intensity and, for some of the ions, the ratio between different emission bands. To date, three main determinants have been used to modulate the luminescence signal of lanthanide-based optical imaging agents for the detection of disease related biomarkers; these are:

- Varying q
- Adjusting the antenna-Ln³⁺ ion distance
- Modulating the antenna excited state

Varying q, number of bound water molecules, influences the emission intensity, luminescence lifetime, and spectral form of otherwise similar Ln complexes. This influence is due to quenching of the excited state(s) caused by the O-H bond vibrations. Design considerations used to adjust the extent of sensitization produced by the antenna stem from an appreciation that the energy transfer from the triplet excited state (T1) of the antenna to the metal ion is both distance and energy dependent. Current systems have focused on altering the Lnantenna distance through chemical reactions or binding events. Finally, modulation of the antenna excited state has been achieved. Typically, this has been done by exploiting aqueousbased chemoselective reactions with kinetics that are fast even in complex biological environments to achieve chemical modification to the antenna. This latter approach is particularly attractive in the context of designing agents for the selective detection of neurodegenerative specific analytes.

For the development of molecular lanthanide-based optical imaging agents, three major factors need to be taken into consideration: The Ln³⁺ ion itself, the photophysical and chemical properties of the chelating ligand, and the photophysical features of the organic-based sensitising antenna. Historically, applications in lanthanide-sensing agents have utilized europium (Eu³⁺) and terbium (Tb3+), owing to their high intrinsic quantum yields, long-luminescence lifetimes, and emission bands that fall in the visible region. Recent efforts have also focused on neodymium (Nd3+) and ytterbium (Yb3+) as near IR-emitting analogues. For additional and extensive overviews on the use of lanthanidebased imaging agents for general bioimaging applications, the reader is directed to two excellent reviews by Meade and Huang.23,38

Sénèque and co-workers recently reported a responsive system that combines functional readout with multiplexing capabilities. In this system, the authors relied on a bioinspired zinc finger peptide strategy for the selective luminescent detection of Zn2+ ions through the modulation of the antenna-Ln³⁺ ion distance (Fig. 36).⁴⁶ They exploited the tendency of $\beta\beta\alpha$ zinc finger (ZF) peptides to change conformation and fold in the presence of Zn²⁺ over other biologically relevant metal ions. Four different ββα zinc finger (ZF) peptides containing a conjugated Ln3+-DOTA complex and a sensitizing antenna were prepared. Chelation of the Zn²⁺ cation resulted in a conformational change, basically a folding to the system, which served to decrease the distance between the small molecule sensitizer and the reporter Ln³⁺-DOTA complex leading to an increase in the luminescence intensity.

Initial efforts were focused on a Tb³⁺-based peptide (LZF1^{Trp|Tb}) with a tryptophan antenna. This framework yielded a time-delayed 8.5-fold enhancement in the emission intensity in the presence of Zn2+ (1 equiv.). Luminescence lifetime measurements revealed no change in the number of water molecules within the Tb³⁺ coordination sphere, providing support for the contention that the observed luminescence enhancement was due to zinc-induced peptide folding. In a similar vein, a neodymium (Nd³⁺) LZF1^{NBD|Nd} probe was developed

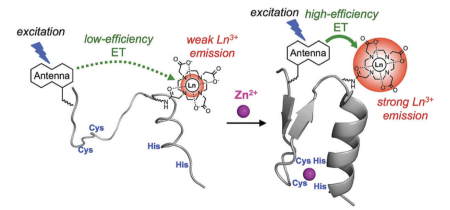


Fig. 36 Binding of Zn^{2+} results in a conformation change of the zinc-finger peptides, which bring the chromophore antenna spatially closer to the Ln^{3+} ion, resulting in an increase in Ln³⁺ emission intensity. Reprinted (adapted) with permission from (Chem. - Eur. J., 2017, 23, 10992-10996). Copyright (2019) John Wiley and Sons.

Tutorial Review

В R' = O, NBn $+\bar{N}H_3$ glutamate HO dopamine NH_2 serotonin NdL³

Fig. 37 (A) Glutamate-responsive contrast agents. Here $Ln = Gd^{3+}$, Eu^{3+} , Yb^{3+} , Nd^{3+} . (B) The chemical structure of glutamate is shown along with that of two common neurotransmitters, dopamine and serotonin. (C) Longitudinal relaxivity change seen for the Gd3+ complex as a function of increasing glutamate (Glu) concentration. (D) Excitation and emission spectra of the Nd³⁺ and Yb³⁺ analogues.

300 400 800

Yhl 3

1000

ntensity (a.u.)

that gave rise to a NIR emission upon exposure to the Zn²⁺ cation in aqueous conditions.

r, (mM⁻¹s⁻¹)

K. = 14.8 M

[Glu]/[GdL3]

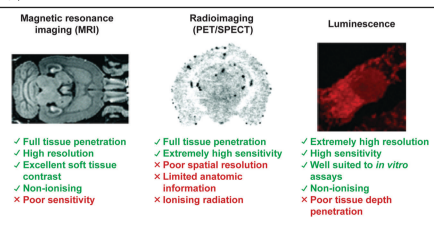
200 400 600

Multimodal systems that combine the emissive Tb3+/Eu3+ ions with the MRI active Gd3+ cation have been studied in an effort to generate imaging agents that permit both luminescence and MRI readouts. 47 Tóth and co-workers recently employed this bimodal approach in an effort to image neurotransmitter signalling in neurodegenerative disease models.⁴⁸ In particular, a

Gd-DO3A complex functionalised with a triazacrown ether (Gd-DO3A-3AC) was prepared. This framework enabled the ditopic binding of the excitatory neurotransmitter L-glutamate (L-Glu), comprised of cationic ammonium (-NH₃⁺) and negatively charged carboxylate (-COO⁻) groups (Fig. 37). The coordinatively unsaturated Gd³⁺ metal centre (positively charged) provided a binding site for the carboxylate functionality (negatively charged), while the appended triazacrown ether unit enabled binding of the

1400 \(\lambda\) (nm)

Table 3 General characteristics of the imaging modalities presented in this review. Combination of the various approaches may alleviate some of the limitations associated with employing each technique individually. MRI image adapted with permission from (Schizophr. Bull., 2008, 34, 595-604). PET/ SPECT image adapted with permission from (J. Med. Chem., 2012, 55, 2279-2286). Luminescent image adapted with permission from (J. Am. Chem. Soc., 2012, 134, 20776-20782)



cationic $-NH_3^+$ group. The dual coordination of Gd-DO3A-3AC rendered it capable of discriminating zwitterionic neurotransmitters from monoamine neurotransmitter (*e.g.*, serotonin νs . noradrenaline). Upon coordination of the appropriate neurotransmitter, a marked change in relaxivity was observed. This was believed to arise from a decrease in the hydration number (q). Gd-DO3A-3AC was also used to monitor neural activity in

Chem Soc Rev

ex vivo mouse brain slices by MRI.

More recently, Toth and co-workers extended this design strategy by attaching a benzophenone chromophore to the lanthanide ligand site. 49 Incorporation of this flat aromatic substituent led to an increased interaction with albumin causing an enhancement in the overall relaxivity. Unfortunately, this came with an overall decrease in the neurotransmitter response. On the other hand, the presence of the benzophenone chromophore allowed for efficient sensitisation of Eu³⁺, Nd³⁺, and Yb³⁺, thus paying the way towards both the visible and nearinfrared (NIR) detection of neurotransmitters. These additional studies by Toth and associates provide an illustrative example of the multimodal capabilities possible by creating a system that can function with different Ln³⁺ ions. It is to be noted, however, that the instability (i.e., photoreactivity) of these first generation systems have precluded their use in more advanced application scenarios.

The efforts with Ln³⁺ luminescent systems summarised above have provided initial support for the notion that such systems may have utility in the context of elucidating the biochemical mechanisms associated with various disease states. However, this area remains in its infancy in terms of potential applications involving neurodegenerative disorders.

Conclusion and future perspectives

As shown throughout this review, PET/SPECT, MRI and luminescence are non-invasive molecular imaging techniques that show particular promise in terms of visualising various pathological biomarkers that may play a role in the development of neurodegenerative diseases. We have to highlight the fact that each metal-based imaging technique has a set of unique advantages and disadvantages (Table 3). For example, metal complex luminescence-based imaging agents offer a method to visualise specific biomarkers with high sensitivity, specificity, and spatial resolution at the cellular/subcellular level (i.e., mitochondria). On the other hand, they generally suffer from poor tissue penetration leading to limited in vitro and in vivo applications. In contrast, PET/SPECT-based imaging agents have been used extensively to monitor various metabolic/ physiological processes and disease related biomarkers with high sensitivity (pM/nM range). However, PET/SPECT provide images with relatively poor resolution and require the use of ionising radiation. On the other hand, small molecule MRI imaging agents have been widely used to effect clinical diagnoses and in the context of research related to neurodegenerative disease. Various small molecule MRI imaging agents have been used to visualise various anatomical architects and pathological biomarkers with high spatial resolution with nearly unlimited depth penetration. However, MRI-based imaging approaches are typically hampered by relatively poor sensitivity (in the μ M range); this makes the detection of biologically important species (*i.e.*, DAT) present at nM concentrations *in vivo* difficult, if not impossible.

Increasingly, researchers in the field are coming to appreciate that no single imaging modality will typically suffice to obtain all the necessary information needed for a clinical diagnosis or for understanding of a specific pathological process. Dedicated efforts in a number of laboratories are thus being made to combine various imaging methods to develop so-called multimodal approaches. For instance, PET/SPECT is being used in conjunction with MRI in an effort to overcome the limitations of each imaging modality so as to unravel more effectively the determinants of various disease states.⁵⁰ We anticipate a continued growth in metal-based imaging agents, particularly multimodal systems, that will permit a deeper understanding of neurodegenerative disease and its underlying causes. Ultimately, the knowledge gained from the design and study of metal-based imaging agents is expected to translate into new therapeutics and cures for these frightening and debilitating disorders.

Abbreviations

Alpha-synuclein

β^+	Beta positive decay
β^-	Beta minus decay
μΜ	Micrometre
γ	Gamma ray emission
¹¹ C	Carbon-11
⁶¹ Cu	Copper-61
⁶² Cu	Copper-62
⁶⁴ Cu	Copper-64
¹⁸ F	Fluorine-18
⁶⁸ Ga	Gallium-68
^{123}I	Iodine-123
⁹⁹ Mo	Molybdenum-99
⁸⁹ Zr	Zirconium-89
ւ-Glu	Glutamic acid
$ au_{\mathbf{m}}$	Mean residence lifetime
$\tau_{ m r}$	Rotational correlation time
Αβ	Amyloid beta
AD	Alzheimer's disease
Ag	Silver
AgCl	Silver chloride
Au ⁺	Gold ion
ALS	Amyotrophic lateral sclerosis
Atsm	$Diacetyl-bis(N^4-methylthiosemicarbazone)$
BBB	Blood brain barrier
bDHC	Bis(dihydroxy)curcumin
BAPTA	1,2-Bis(o-aminophenoxy)ethane-N,N,N',N'-tetra-
	acetic acid
BOLD	Blood-oxygenation level dependent

Tutorial Review

Millimolar BPy Bipyridyl mM Metal-metal-to-ligand charge-transfer Bzq Benzoquinoline MMLCT CAContrast agent MLCT Metal-to-ligand charge-transfer Ca2+ Calcium ion Metal-to-ligand-ligand charge-transfer MLLCT $Mn^{2+/3+}$ CEST Chemical exchange saturation transfer Manganese ion CN Coordination number MRI Magnetic resonance imaging **CNS** Minute Central nervous system Min CO Carbon monoxide ms Millisecond Cp Cyclopentyl mV MilliVolt $Cu^{+/2+}$ Nd^{3+} Copper ion Neodymium ion CTComputer tomography nM Nanomolar DAC NOTA Diacetylcurcumin 1,4,7-Triazacyclononane-1,4,7-trisacetic acid DAT Dopamine active transporter ns Nanosecond DFO Desferrioxamine B O_2 Oxygen O_2^{\bullet} DO3A 1,4,7,10-Tetraazacvclododecane-1,4,7-triacetic Superoxide anion Os^{2+} acid Osmium ion DOTA PCTA 1,4,7,10-Tetraazacyclododecane-1,4,7,10-3,6,9,15-Tetraazabicyclo[9.3.1]-pentadecatetraacetic acid 1(15),11,13-triene-3,6,9-triacetic acid DOPTA 1,4,7-Tris-(carboxymethyl)-1,4,7,10-PD Parkinson's disease tetraazacyclododecane PDA Phenylenediamido dppz Dipyridophenazine PEG Polyethylene glycol EC PET Electron capture Positron emission tomography Pt2+ **EDTA** Ethylenediaminetetraacetic acid Platinum ion Eu Europium Phen 1,10-Phenathroline FDA Food and drug administration PiB Pittsburgh compound B $Fe^{2+/3+}$ **PLIM** Phosphorescence lifetime imaging **fMRI** Functional magnetic resonance imaging Phq 2-Phenylquinoline **GABA** γ-Aminobutyric acid 2-Phenylpyridine Ppy GAD Glutamate decarboxylase Py Pyridine Gd^{3+} Gadolinium ion pM Picomolar Ge Germanium ps Picoseconds GFP Green fluorescent protein Pyridine Py Glyoxal-bis(N^4 -methyl-3-thiosemicarbazonato) Average number of water molecules bound gtsm q h Hours directly to the paramagnetic metal centre Hb Haemoglobin Relaxivity $r_{1/2}$ HDHuntington's disease Re⁴ Rhenium ion Rh³⁺ Rhodium ion $ID g^{-1}$ Injected does per gram tissue ILIntraligand Radiofrequency RF In³⁺ Ru²⁺ Indium ion Ruthenium ion Ir^{3+} Iridium ion SPECT Single photon emission computed tomography K^{+} Potassium ion **SPION** Superparamagnetic iron oxide nanoparticles $K_{\rm d}$ Dissociation constant **USPIO** Ultra-small superparamagnetic iron oxide keV Kiloelectron volts nanoparticles Inhibition constant Half-life K_{i} $t_{1/2}$ L Ligand \mathbf{T} Tesla T_1 LCIO Lipid coated iron oxide nanoparticles Longitudinal time constant Ligand-to-ligand charge-transfer Transverse time constant LLCT T_2 TAE Tetraarylethylene LMMCT Ligand-to-metal-metal charge-transfer Tb³⁺ Ligand-to-metal charge-transfer LMCT Terbium ion ^{99m}Tc Ln³⁺ Lanthanide ion Technetium-99m Lysine Transgenic Lys Tg TGL Time-gated luminescence mAb Monoclonal antibody MaCaReNa Magnetic calcium-responsive nanoparticles ThT Thioflavin T

 Yb^{3+}

Zn

⁸⁹Zr

Ytterbium ion

Zirconium-89

Zinc

Megaelectron volt

imaging

Manganese-enhanced magnetic resonance

MeV

MEMRI

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

DAI thanks the Life Sciences Research Foundation for a post-doctoral fellowship sponsored by the Howard Hughes Medical Institute. PH thanks the University of Nottingham for support through a Nottingham Research Fellowship. XPH would like to thank The National Key Sci-Tech Special Projects of Infectious Diseases of China (2018ZX10732202). XPH and HT thank financial support from Natural Science Foundation of China (No. 21788102, 91853201, 21722801 and 21776078) and the Shanghai Municipal Science and Technology Major Project (No. 2018SHZDZX03). A. C. S. and J. T. B. would like to thank Gregory D. Thiabaud (UT Austin) for helpful discussions. JLS would like to thank the National Institutes of Health (grant CA 68682) and the Robert A. Welch Foundation (F-0018) for support. TDJ wishes to thank the Royal Society for a Wolfson Research Merit Award.

References

- 1 C. Soto and S. Pritzkow, Nat. Neurosci., 2018, 21, 1332-1340.
- 2 P. J. Blower, Dalton Trans., 2015, 44, 4819-4844.
- 3 C. Grupen, *Introduction to Radiation Protection*, Springer, Berlin, 2010.
- 4 L. Zhu, K. Ploessl and H. F. Kung, *Chem. Soc. Rev.*, 2014, 43, 6683–6691.
- 5 K. H. Chen and M. C. Cui, *MedChemComm*, 2017, 8, 1393–1407.
- 6 M. Ikawa, H. Okazawa, T. Kudo, M. Kuriyama, Y. Fujibayashi and M. Yoneda, *Nucl. Med. Biol.*, 2011, **38**, 945–951.
- 7 K. Wada, Y. Fujibayashi, N. Tajima and A. Yokoyama, *Biol. Pharm. Bull.*, 1994, 17, 701–704.
- 8 J. B. Torres, E. M. Andreozzi, J. T. Dunn, M. Siddique, I. Szanda, D. R. Howlett, K. Sunassee and P. J. Blower, *J. Nucl. Med.*, 2016, 57, 109–114.
- 9 Z. Xiao, P. S. Donnelly, M. Zimmermann and A. G. Wedd, *Inorg. Chem.*, 2008, 47, 4338–4347.
- 10 C. J. Maynard, A. I. Bush, C. L. Masters, R. Cappai and Q. X. Li, *Int. J. Exp. Pathol.*, 2005, **86**, 147–159.
- 11 J. L. Hickey, S. Lim, D. J. Hayne, B. M. Paterson, J. M. White, V. L. Villemagne, P. Roselt, D. Binns, C. Cullinane, C. M. Jeffery, R. I. Price, K. J. Barnham and P. S. Donnelly, J. Am. Chem. Soc., 2013, 135, 16120–16132.
- 12 N. Bandara, A. K. Sharma, S. Krieger, J. W. Schultz, B. H. Han, B. E. Rogers and L. M. Mirica, *J. Am. Chem. Soc.*, 2017, 139, 12550–12558.
- 13 H. Watanabe, M. Ono, S. Iikuni, M. Yoshimura, K. Matsumura, H. Kimura and H. Saji, *Bioorg. Med. Chem. Lett.*, 2014, 24, 4834–4837.
- 14 K. Chauhan, A. Datta, A. Adhikari, K. Chuttani, A. K. Singh and A. K. Mishra, *Org. Biomol. Chem.*, 2014, 12, 7328–7337.

- 15 M. Asti, E. Ferrari, S. Croci, G. Atti, S. Rubagotti, M. Iori, P. C. Capponi, A. Zerbini, M. Saladini and A. Versari, *Inorg. Chem.*, 2014, 53, 4922–4933.
- 16 G. Van Dongen, M. C. Huisman, R. Boellaard, N. H. Hendrikse, A. D. Windhorst, G. W. M. Visser, C. F. M. Molthoff and D. J. Vugts, Q. J. Nucl. Med. Mol. Imaging, 2015, 59, 18–38.
- 17 J. Fissers, A. M. Waldron, T. De Vijlder, B. Van Broeck, D. J. Pemberton, M. Mercken, P. Van der Veken, J. Joossens, K. Augustyns, S. Dedeurwaerdere, S. Stroobants, S. Staelens and L. wyffels, *Mol. Imaging Biol.*, 2016, 18, 598–605.
- 18 M. P. Kung, D. A. Stevenson, K. Plossl, S. K. Meegalla, A. Beckwith, W. D. Essman, M. Mu, I. Lucki and H. F. Kung, Eur. J. Nucl. Med., 1997, 24, 372–380.
- 19 P. D. Mozley, J. B. Stubbs, K. Plossl, S. H. Dresel, E. D. Barraclough, A. Alavi, L. I. Araujo and H. F. Kung, *J. Nucl. Med.*, 1998, 39, 2069–2076.
- 20 Y. Cheng, M. Ono, H. Kimura, M. Ueda and H. Saji, *J. Med. Chem.*, 2012, **55**, 2279–2286.
- 21 X. Y. Zhang, P. R. Yu, Y. P. Yang, Y. Q. Hou, C. Peng, Z. G. Liang, J. Lu, B. A. Chen, J. P. Dai, B. L. Liu and M. C. Cui, *Bioconjugate Chem.*, 2016, 27, 2493–2504.
- 22 A. Kurihara and W. M. Pardridge, *Bioconjugate Chem.*, 2000, 11, 380–386.
- 23 M. C. Heffern, L. M. Matosziuk and T. J. Meade, *Chem. Rev.*, 2014, **114**, 4496–4539.
- 24 J. Wahsner, E. M. Gale, A. Rodriguez-Rodriguez and P. Caravan, *Chem. Rev.*, 2019, **119**, 957–1057.
- 25 G. R. Naumiec, G. Lincourt, J. P. Clever, M. A. McGregor, A. Kovoor and B. DeBoef, *Org. Biomol. Chem.*, 2015, 13, 2537–2540.
- 26 L. Frullano, J. Q. Zhu, C. N. Wang, C. Y. Wu, R. H. Miller and Y. M. Wang, J. Med. Chem., 2012, 55, 94–105.
- 27 G. Bort, S. Catoen, H. Borderies, A. Kebsi, S. Ballet, G. Louin, M. Port and C. Ferroud, *Eur. J. Med. Chem.*, 2014, **87**, 843–861.
- 28 W. H. Li, S. E. Fraser and T. J. Meade, *J. Am. Chem. Soc.*, 1999, **121**, 1413–1414.
- 29 E. L. Que, E. J. New and C. J. Chang, *Chem. Sci.*, 2012, 3, 1829–1834.
- 30 M. Aswendt, E. Gianolio, G. Pariani, R. Napolitano, F. Fedeli, U. Himmelreich, S. Aime and M. Hoehn, *NeuroImage*, 2012, **62**, 1685–1693.
- 31 A. C. Silva and N. A. Bock, Schizophr. Bull., 2008, 34, 595-604.
- 32 A. Bertin, J. Steibel, A. I. Michou-Gallani, J. L. Gallani and D. Felder-Flesch, *Bioconjugate Chem.*, 2009, **20**, 760–767.
- 33 A. Barandov, B. B. Badelle, C. G. Williamson, E. S. Loucks, S. J. Lippard and A. Jasanoff, *Nat. Commun.*, 2019, 10, 897.
- 34 R. Sperling, Neurobiol. Aging, 2011, 32, S37-S43.
- 35 T. Lee, L. X. Cai, V. S. Lelyveld, A. Hai and A. Jasanoff, Science, 2014, 344, 533–535.
- 36 Y. Z. Wadghiri, J. L. Li, J. H. Wang, D. M. Hoang, Y. J. Sun, H. Xu, W. Tsui, Y. S. Li, A. Boutajangout, A. Wang, M. de Leon and T. Wisniewski, *PLoS One*, 2013, 8, DOI: 10.1371/journal.pone.0057097.
- 37 S. Okada, B. B. Bartelle, N. Li, V. Breton-Provencher, J. J. Lee, E. Rodriguez, J. Melican, M. Sur and A. Jasanoff, *Nat. Nanotechnol.*, 2018, **13**, 473–477.

38 K. Y. Zhang, Q. Yu, H. J. Wei, S. J. Liu, Q. Zhao and W. Huang, Chem. Rev., 2018, 118, 1770-1839.

Tutorial Review

- 39 Q. Zhao, C. H. Huang and F. Y. Li, Chem. Soc. Rev., 2011, 40, 2508-2524.
- 40 B. Y. W. Man, H. M. Chan, C. H. Leung, D. S. H. Chan, L. P. Bai, Z. H. Jiang, H. W. Li and D. L. Ma, Chem. Sci., 2011, 2, 917-921.
- 41 L. H. Lu, H. J. Zhong, M. D. Wang, S. L. Ho, H. W. Li, C. H. Leung and D. L. Ma, Sci. Rep., 2015, 5, 25774, DOI: 10.1038/srep25774.
- 42 N. P. Cook, V. Torres, D. Jain and A. A. Marti, J. Am. Chem. Soc., 2011, 133, 11121-11123.
- 43 N. P. Cook, K. Kilpatrick, L. Segatori and A. A. Marti, J. Am. Chem. Soc., 2012, 134, 20776-20782.
- 44 A. Aliyan, B. Kirby, C. Pennington and A. A. Marti, I. Am. Chem. Soc., 2016, 138, 8686-8689.

- 45 M. T. Gabr and F. C. Pigge, Chem. Eur. J., 2018, 24, 11729-11737.
- 46 M. Isaac, L. Raibaut, C. Cepeda, A. Roux, D. Boturyn, S. V. Eliseeva, S. Petoud and O. Seneque, Chem. - Eur. J., 2017, 23, 10992-10996.
- 47 E. Debroye and T. N. Parac-Vogt, Chem. Soc. Rev., 2014, 43, 8178-8192.
- 48 F. Oukhatar, S. Meme, W. Meme, F. Szeremeta, N. K. Logothetis, G. Angelovski and E. Toth, ACS Chem. Neurosci., 2015, 6, 219-225.
- 49 F. Oukhatar, S. V. Eliseeva, C. S. Bonnet, M. Placidi, N. K. Logothetis, S. Petoud, G. Angelovski and É. Tóth, Inorg. Chem., 2019, 58, 13619-13630, DOI: 10.1021/acs.inorgchem.9b00887.
- 50 C. T. Yang, K. K. Ghosh, P. Padmanabhan, O. Langer, J. Liu, D. N. C. Eng, C. Halldin and B. Gulyas, Theranostics, 2018, 8, 6210-6232.