Biomedical Applications of Lanthanide Nanomaterials, for Imaging, Sensing and Therapy

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Abstract

The application of nanomaterials made of rare earth elements within biomedical sciences continues to make significant progress. The rare earth elements, also called the lanthanides, play an essential role in modern life through materials and electronics. As we learn more about their utility, function, and underlying physics, we can contemplate extending their applications to biomedicine. This particularly applies to diagnosis and radiation therapy due to their relatively unique features, such as an ultra-wide Stokes shift in the luminescence, variable magnetism and potentially tunable properties, due to the library of lanthanides available and their multivalent oxidation state chemistry. The ability to prepare nanomaterials of relatively smaller sizes has increased the likelihood of use in vivo. In this review, we summarize the different emerging applications of nanoparticles with rare earth elements as the host or doped elements for biomedical applications in the past three to four years, especially in the area of imaging and disease diagnosis. Researchers have made progress in utilizing surfactants and polymers to modify the surface of lanthanide nanoparticles to enhance biocompatibility. At the same time, specific antibodies and proteins can also be conjugated to these nanoparticles to increase targeting efficiency for specific tumor models. Finally, in the near-infrared II imaging window, lanthanide nanoparticles have been shown to exhibit extraordinary bright emission, which is an exciting development for image-guided surgery.

Key words: Lanthanide Nanomaterials, Imaging, Sensing, Therapy

Introduction

This is a short review of the current use of lanthanide nanomaterials for biomedical applications. The word lanthanide is used colloquially over lanthanoid, which is the official IUPAC name. Collectively with Scandium and Yttrium, the term rare earth element (REE) is frequently used in geology, science and medicine; however, their relative natural abundance is not that scarce in most cases. The f-orbitals start with element 57, lanthanum, the first of the lanthanides and namesake of the elementary group. There are so many of these f-block elements (15, from Lanthanum to Lutetium) it is necessary for formatting/aesthetic reasons to place them at the bottom of the periodic table, with the actinides. Their discovery spans the 19th century, while the emergence of their widespread utility spans the latter half of the 20th century until the present day. As we have learned to harness the unique physics and chemistry characterized by their f-orbital behavior, there have been several key applications that have revolutionized industries, including the development of phosphors for fluorescent/LED lighting [1] and the development of rare earth magnets, which now dominate the design of personal headphones, motors for electric/hybrid vehicles, wind turbines, and portable MRI scanners [2-4]. Since 1959, when the
Dutch-American physicist and Nobel laureate Nicolaas Bloembergen observed nonlinear up-conversion of lanthanides in the Near IR region, optical properties (photonics) research chiefly focused on semiconductors and electronic equipment [5]. However, in more recent times, and primarily through adaptation enabled by nanotechnology, significant achievements have been made in basic research of rare earth element chemistry for biomedical applications [6].

In general, molecular-level based imaging is increasingly used for disease diagnosis and guidance of surgery [7-14]. Among the most widely used in vivo applications are planar and tomographic fluorescence and bioluminescence imaging. However, despite their utility, these techniques are primarily restricted to pre-clinical use. Factors that have prevented translation from the bench to the bedside include depth-penetration considerations, regulatory issues, and toxicity of the lanthanides [15, 16]. A recent development in lanthanide element based near IR-II nanoprobes [17] successfully overcame the shortcomings of traditional fluorescence contrast agents by extending the imaging window closer to the near IR-II region (1000-2000 nm). In parallel, nanotechnology researchers dedicated to developing rare earth nanoprobes with improved surface modification, morphology, and size, have enabled increased function for diagnosis or targeting.

In terms of treatment, especially tumor treatment, due to the powerful optical properties of lanthanides, there are remarkable effects when combining them with agents designed for photothermal therapy (PTT) or photodynamic therapy (PDT). Hereafter, we discuss the development of lanthanide element-based nanoparticles for in vivo studies. Next, we evaluate how researchers have been employed effectively as dual-modal imaging contrast agents for optical and MRI imaging in vivo[26]. In the recent work by Shane Plunkett et al., the procedure was improved by using Janus-type dendrimers, which have individual carboxylates on the surface of the ligand for conjugation to the near IR-II region (1000-2000 nm). In parallel, nanotechnology researchers dedicated to developing rare earth nanoprobes with improved surface modification, morphology, and size, have enabled increased function for diagnosis or targeting.

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Surface modification of lanthanide-based nanoparticles for in vivo studies

For pre-clinical nanomedical research, especially in inorganic metal/metal oxide nanoparticles, the highest priority is to enhance biocompatibility. Most biomaterials are considered as foreign matter to the host, especially the inorganic materials, and will induce the rejection reaction. The successful design of an in vivo used material must include low toxicity, longer circulation time, and even biodegradable or smaller enough size for extraction from the glomerulus. Making the lanthanide nanomaterial more biocompatible through surface modification became a popular research field for material scientists: Liu et al. developed a novel inorganic oxide nanoprobe based on amine-functionalized tetragonal ultrasmall ZrO2Ln3+ nanoparticles (< 5nm) synthesized through a simple solvothermal technique (Fig. 1C). To substitute the natural hydrophobic ligands on the surface of NPs, amine and phosphate groups were introduced for a ligand exchange reaction. They show the use of this hydrophilic Ln3+-based long-lived luminescence as a sensitive, time-resolved fluorescence resonance energy transfer (FRET) nanoprobe to identify avidin with a detection limit as low as 3.0 nM [24]. Fischer et al. reported synthesizing a series of sub-15 nm alkaline-earth lanthanide fluoride upconversion nanoparticles with different doping combinations (M1-x Ln3 Ln2x, MLnF) (Fig. 1B). Then, CaF2 shells, which have earlier been demonstrated to be biocompatible, are employed for passivation to increase the upconversion performance of these nanoparticles by more than five orders of magnitude while also enhancing their biocompatibility [25]. Zhijie Wang et al. presented a facile one-pot procedure for producing highly hydrophilic and biocompatible NaGdF4:Yb3+/Er3+ and NaGdF4:Yb3+/Tm3+ nanocrystal by thermal hydrolysis of lanthanide-oleate precursors and sodium fluoride by added α-dicarboxylic polyethylene glycol (HOOC-PEG-COOH) (Fig. 1D).

Moreover, NaGdF4:Yb3+/Er3+ nanocrystals have been employed effectively as dual-modal imaging contrast agents for optical and MRI imaging in vivo[26]. In the recent work by Shane Plunkett et al., the procedure was improved by using Janus-type dendrimers, which have individual carboxylates on the surface of the ligand for conjugation to NaYF4-based nanocrystals, doped with Yb3+ (20%) and Er3+ (20%), neutral polyethylene glycol (PEG) groups for water solubility. They also encapsulate a pH-sensitive dye, Eriochrome, as the center of the dendrimer. These Janus-dendrimers modified NaYF4 based nanocrystals were excellent vascular markers, enabling for depth-resolved mapping of specific capillaries in the mice brain achieved a depth of 1000 um under continuous wave (CW) stimulation with a relatively low energies level of no more than 20 mW (Fig. 1A) [27]. Another exciting aspect of lanthanide...
nano oxides is unusual water-solubility, which even could be used to functionize other materials to render them hydrophilic. In Wen Luo et al.'s recent work, they doped the Titanium−Oxo Cluster with Eu(III), Tb(III), and Yb(III) to form water-soluble lanthanide−titanium−oxo clusters (LnTOCs) which could quickly disperse in aqueous media to form a stable solution with aggregation of the clusters. The photocatalytic behavior, fluorescence sensing, and fluorescent labeling have all been retained and verified through in vitro cell imaging (Fig. 1E) [28].

Cancer imaging and therapy

Ren et al. developed Er-based lanthanide nanoparticles NaErF4:2.5 %Ce@NaYbF4, coupled with Dye-brush polymer (Dye-BP) to promote the $^4I_{13/2} \rightarrow ^4I_{15/2}$ transition, achieved 675-fold enhancement at 1525 nm wavelength near the IR II imaging window. They further conjugated a tumor-targeting angiopep-2 peptide, which could target the glioma in a mice model. Under the guidance of this near IR luminescence probe, they successfully removed the glioma from the mouse brain (Fig. 2A) [29]. Peroxynitrite(ONOO−) is over-expressed in the tumor-microenvironment. Zhao et al. developed a lanthanide-based β-NaYF4@NaYF4:1%Nd NIR-II nanosensor which attached an ONOO− responsive dye MY-1057. They utilize the specific energy transfer
between the ONOO- to the MY1057 dye and finally excite the β-NaYF₄@NaYF₄:1%Nd nanosensor in the NIR-II region. With the NIR light from the nanosensor, tumor lesions (Hepatocellular Carcinoma) could be distinguished from normal tissue (Fig. 2C) [30]. PDT and CDT (chemodynamic therapy) is usually limited by glutathione and hypoxia in the tumor. Jiating Xu et al. developed a copper/manganese silicate nanosphere (CMSN)–covered by Yb³⁺/Er³⁺/Tm³⁺shell nanoparticles which not only provide upconversion (UC) and downconversion (DC) emissions for NIR (1000 nm – 1700 nm wavelength, the significance of which is explained later in this section) imaging but also contribute to enhancing the potential for PDT through promoting the O₂ to singlet O₂ by the NIR light emitting from the lanthanide element (Fig. 2E) [31]. Since some of the lanthanides are also paramagnetic, they can be used as MRI contrast agents. Zhang et al. synthesized Lanthanide−Cyclen−Camptothecin nanocomposites encapsulating Gd(III) and Yb(III) complexes and the anticancer drug camptothecin into a micelle, offering real-time NIR / MR dual model imaging and chemotherapy (Fig. 2B, D) [32]. Lanthanide nanoparticles could be used as a platform incorporated with other nanomedicines for theranostic approaches. Xu et al. integrated CuS nanoparticles (PTT agents) and g-C₃N₄ QDs (PDT agents) on silica-coated NaGdF₄:Yb,Tm@NaGdF₄:Nd,Yb UCNPs. When they applied an 808 nm laser to this nanocomplex, the lanthanide UCNP would shift the light to UV/vis region and simultaneously active the PTT and PDT agent to achieve the effect of combination therapy [33]. Similarly, another group of researchers utilized the same material and 808 nm laser for image-guided tumor resection surgery by

Figure 2: A) Schematic illustration of T2-weighted magnetic resonance imaging (MRI), and noninvasive NIR IIb fluorescence imaging of a deep-seated and small-sized orthotopic glioma under the assistance of focused ultrasound (FUS) B, D) (i) In vivo T1-weighted MR images of the mice bearing HeLa xenograft from 0 to 24 h after the injection of Gd/YbNPs and Gd DOTA (control). (ii) Plots of 1/T₁ vs. [Gd] to determine r₁ of Gd-DOTA and Gd/YbNPs hybrid nanoparticles. C) Noninvasive intensity-based imaging, lifetime-based imaging of a mouse bearing multiple HCC lesions and optical photo of the dissected liver. E) Theranostic mechanism of PEG-LDNPs@CMSNs for TME and NIR laser co-enabled PDT/CDT and trimodal bioimaging F) Lanthanide–Cyclen–Camptothecin Nanocomposites for Cancer Theranostics Guided by Near-Infrared and Magnetic Resonance Imaging [15-19].
doping Ce$^{3+}$. They achieved an 11-fold enhancement of the NIR-II emission above 1500nm and successfully removed tumors in a colorectal cancer mice model [34]. Doping lanthanide elements into polymer nanoparticles is another strategy for cancer imaging or therapy since most organic/polymer nanoparticles can degrade over specific pH environments. Miao Feng et al. introduced hydrophobic NaGdF$_4$:2%Nd@NaLuF$_4$ nanocrystals ultrasmall nanoparticle and doxorubicin into a 300 nm pH-sensitive degradable mPEG-PLGA nanomicelles for imaging and chemotherapy application (Fig. 3A) [35]. While these approaches all used nontargeted agents, lanthanide nanoparticles were also targeted to a specific cancer model for imaging when coupled with targeted peptides. Jingxue Cao et al. designed a $\alpha$β3/$\alpha$β5 and aminopeptidase-N-receptor targeting peptides conjugated to NaYF$_4$:Yb nanoparticle as NIR-II nanoprobes for in vivo imaging of lung cancer (Fig. 3C) [36]. To maximally lower the toxicity of the inorganic lanthanide nanoparticle in vivo, an excretable lanthanide nanoprobe for imaging and surgical guidance in the NIR-II window has been developed by Daifen Li and his colleagues (Fig. 3B). They encapsulate ultra-small hydrophobic NaYF$_4$:Nd 7% into bio-degradable liposomes. This strategy can quickly clear their particle via the kidneys after intravenous injection with a half-life of 23.0 h for the liver and 14.9 h for the spleen [37]. Metal–organic frameworks (MOFs) are posited as excellent photosensitizers for photodynamic therapy (PDT) in cancer therapy. However, the efficacy of the MOFs PDT has been limited by the tissue penetration of the visible excitation light (Fig. 3F).

When coupled with upconversion lanthanide nanoparticles, a more penetrating NIR light source outside the body can be used. The NIR light can be transferred to visible light via upconversion lanthanide nanoparticles inside the body, which could further excite the MOFs to generate singlet oxygen and significantly increase PDT efficacy [38]. As an alternative to upconversion, one can also manipulate lanthanide nanoparticles for a down-converting application. Downconversion means shifting the UV/visible (200-400nm) or near IR (800-1000nm) to longer red light (600-700nm) or far IR region (beyond 1500 nm), respectively. Hongjie Dai’s group at Stanford designed a biocompatible cubic-phase (α-phase) erbium-based lanthanide nanoparticle attached with anti-PD-L1 antibodies that exhibits bright downconversion emission at ~1,600 nm for dynamic imaging of cancer immunotherapy in mice (Fig. 3D) [39]. Another group successfully converted a 980 nm laser emission to 1500 nm (NIR-II region) via poly(acrylic acid) (PAA)-modified NaLnF$_4$:40Gd/20Yb/2Er nanorods for cancer metastasis and vessel imaging (Fig. 3E) [40]. Downconverting technology could even be applied to nuclear imaging, especially Cerenkov imaging. A europium oxide nanoparticle was utilized to shift blue Cerenkov radiation to the more

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**Figure 3:** A) Schematic illustration of the bioapplication of DMDNs(DOX@mPEG-PLGA@DCNP nanoparticles). B) Schematic illustration of excretable lanthanide nanoparticles for multifunctional biomedical imaging and surgical navigation in the second near-infrared window. C) Treated group: tumor-bearing mice injected with UCNP@RGD-NGR. Digital photos were taken at 0, 15, 30, 60, 120 min after injection under 980 nm excitation. D) i), Schematic illustration of the hydrophilic ErNPs with cross-linking polymeric layers and amine groups on the surface as conjugation sites. ii), DLS spectra of hydrophilic ErNPs with polymeric cross-linked network and NIR-IIb cerebral vascular image (left) by intravenous injection of 200 μl ErNPs (40 mg ml$^{-1}$) and excitation by a 970 nm LED (30 fps). The luminescence intensity of an inferior cerebral vein was plotted as a function of time (iii), showing the cardiac cycles (iv) with a heartbeat frequency of 3.67 Hz by fast Fourier transformation (FFT, lower right). Scale bar, 5 mm E) i) Digital photograph of tumor-bearing mouse and in vivo NIR-IIb fluorescent imaging (the green circle indicated the tumor site). ii, iii) Digital photographs of the isolated organs/ tumor and the corresponding ex vivo NIR-IIb imaging, respectively. F) i) Schematic illustration of the synthesis of UCDs and UCSs through the conditional surface engineering of UCNPs. The structure of porphyrinic MOFs is illustrated in the right column. ii) TEM images from left to right is UCNPs, UCDs, and UCSs. Scale bar, 50 nm. iii) Elemental mapping images of a single (left to right) UC and (f) UCS. Scale bar, 50 nm. G) i-iv) Morphology of NdIII-IP virus-like nanodrugs and IP nanospheres observed by SEM, TEM [21-29].
penetrative red-light region, which significantly enhanced the Cerenkov light for lymph node and tumor imaging [41, 42]. Researchers have also designed rare-earth nanoparticles with different morphologies, resulting in unexpected and interesting effects under certain circumstances. Yang Li et al. created a virus-like nanoconjugate that included a NIR-II fluorescence agent (IR825), a chemotherapy drug (pemetrexed, PEM), and a lanthanide element (Nd(III))(Fig. 3G). The spike-like poly(ethylene glycol) shell protected the nanoparticle from immune clearance and enhanced the in vivo circulation time [43]. Xu et al. produced two-dimensional ultrathin lanthanide oxyiodide nanosheets for anticancer drug delivery. Those nanosheets exhibit remarkable drug loading capacity (300 wt.%) compared to other nanomaterial morphologies [44]. The unique paramagnetic property of gadolinium makes it an excellent MRI contrast agent among rare earth elements. Traditionally, Gd is usually chelated to small molecules. However, material scientists could also make a nanoparticle that uses Gd as the main or doped component. With the benefit of nanotechnology, the Gd nanoparticle could reach the targeted area, relying on the EPR effect, or the particle could be encapsulated with other functional elements to achieve different imaging modalities. Ye Kuang and colleagues doped Gd into HfO2 to create a Gd:HfO2 nanoparticle that utilizes the Gd as an MRI contrast agent and Hf as a radiosensitizer. They surface modified the nanoparticle with RGD peptides and loaded the particle with cisplatin to create a potential MRI-guided combined chemo-/ photothermal-/radiotherapy nanoplatfon for drug-resistant tumors [45]. McLeod et al. incorporated a Gd(III) complex into a Zr-based MOF (Metal-organic framework) through solvent-assisted ligand incorporation (SALI) and tested the combined MOFs as a potential MRI contrast agent [46]. By combining Gd and barium titanate, researchers Peng Wang et al. created a dual MRI and X-ray imaging agent [47]. Meanwhile, Yao Cai et al. labeled human heavy chain ferritin with Gd for enhanced MRI tumor imaging applications [48]. The most abundant lanthanide element, Cerium, is another potential candidate for biomedical applications such as sensors or imaging platforms. With the surface oxygen vacancies, CeO2 has been utilized as a contract agent or carrier for another rare earth element for imaging purposes. CeO2 also presents native fluorescence due to the presence of Ce3+ [49]. Meanwhile, CeO2 can also be used as a convenient host lattice with which to combine with another lanthanide element to achieve more imaging modalities. Anne et al. doped ytterbium and gadolinium into the CeO2 nano cubes for MRI and Near IR imaging on cancer cells [50]. Chulun Shao et al. utilize the abundant oxygen vacancies on the CeO2 surface and integrate the Gd for DCE-PWI (dynamic contrast-enhanced perfusion-weighted imaging) MRI [51]. Since lanthanide elements are in the d-block of the periodic table, they are also considered as high atomic number elements, which usually provide a good contrast for X-ray imaging. Zuwu Wei et al. developed a hollow CeO2-ZrO2 hybrid nanoplateform labeled with Gd for MRI/CT imaging and drug (Dox) delivery for heterotopic, subcutaneous human liver cancers [52].

Radiotherapy of 177 Lutetium

Lutetium is a unique member of the lanthanide family since one of its isotopes, Lutetium 177, intrinsically has electron and high energy gamma emission, making it a good candidate for targeted radiotherapy. 177 Lu has been used as a radiotherapy agent for a long time in the literature [53], and more recently has been chelated to small molecule peptides or prostate-specific membrane antigen (PSMA) inhibitors for targeted radiotherapy of neuroendocrine tumors and prostate cancers. Here, we touch on a subsection of the unique applications of 177Lu combined nanotechnology in recent years. Rodrigo et al. recently synthesized an Eu-doped Mesoporous SiO2 labeled with 177 Lu for HT-29 colorectal cancer imaging and radiotherapy. The author cleverly uses the beta emission from the 177Lu for a therapy study, and the gamma emission for SPECT imaging [54]. Another group at the University of Wisconsin (Yu et al.) developed a facile approach to assembling 177Lu-Porphyrin-PEG-Nanocomplexes which combined fluorescence imaging, photodynamic therapy with positron emission tomography imaging, internal radiotherapy and specifically targeted the tumor cell mitochondria [55]. Metal nanoparticles could also be labeled with 177Lu for therapy study. Pei Pei et al. chelated gold nanoclusters with 177Lu and 99mTc, and could strengthen the radioisotope therapy efficiency and induce anticancer immunity by activating dendritic cells [56].

Biosensing and other biomedical applications of lanthanide nanomaterials

Based upon excellent optical features, such as low background fluorescence, and high sensitivity, lanthanide nanomaterials can be applied to various biosensing areas, such as pH detection, essential physiological ions, ROS detection, temperature monitoring or even COVID-19 testing. A current, pandemic related application of lanthanide
nanoparticles is the utilization of self-assembled LNPs (lanthanide-doped polystyrene nanoparticles) as a fluorescent sensor to detect anti-SARS-CoV-2 IgG (Fig. 4C) [57]. Inflammation imaging has become, next to cancer imaging, an essential diagnostic tool for disease detection. Zhao et al. developed an ultra-small glutathione (GSH)-modified lanthanide downconversion nanoparticle system that could precisely target and detect ROS in the inflammation through a cross-linking strategy (Fig. 5C). Light and heat have always been a pair of inseparable factors. The intracellular thermal energy could affect the energy transfer of the outer shell electron of the lanthanide element. The outer shell electron configuration usually determines the light emission intensity of the rare earth element. Piñol et al. employed this theory to develop innovative Ln³⁺-doped polymeric micelles, which could achieve real-time 2D mapping of the intracellular temperature of breast metastatic adenocarcinoma cells (Fig. 5B) [59].

Figure 4: A) In vivo NIR-II imaging of C57BL/6 mice (n = 3) bone in the NIR-IIa 1345 nm and NIR-II 1064 nm windows. In vivo fluorescence of the whole body in (a) prone, (b) supine, and (c) lateral. B) Schematic diagram of a nude mouse by intragastric administration and intravenous injection with cssEr-PEG and cssNd-PEG aqueous dispersions, respectively. C) Design and fabrication of the developed assay. (i) Lateral flow test strip. (ii) Assay. D) Projections of a z-stack of images of a HeLa cell incubated with ECP-NPs (PMMA-COOH 10% at 40% [Eu(tta)3phen]). ECP-NPs are shown in magenta and the cell membranes in turquoise (costained with DiD). E) Photograph of papery patterns under 1540 nm excitation (power density: 10 W/cm²). (i) In vivo multiplexing upconversion imaging with aqueous CS-CS₄ nanoparticles under 1540 nm excitation. Bright-field (left) and dark-field (right) (power density: 0.5 W/cm²). (ii) Luminescence cell imaging of Ln₃⁺-MOFs. (above) Confocal images of the CT26 cells incubated with Eu-BDC-PEG and Tb-BDC-PEG nMOFs for 3h. Green and red fluorescence represent the emission from Tb³⁺ and Eu³⁺ the PEGylated nMOFs, respectively. Scale bars correspond to 50 μm. (below) In vitro RLI of CT26 cell pellets under X-ray excitation of (i) blank cells and cells incubated with (ii) Eu-BDC-PEG, (iii) Tb-BDC-PEG, and (iv) Eu₅TbₓBDC-PEG nMOFs for 24 h [30-36].
Similarly, the Mott-Seitz model could describe the light and temperature relationship as equation (1).

\[
\text{Mott-Seitz Model } \tau(T) = \frac{\tau_0}{1 + e^{\frac{-\Delta E}{kT}}} \tag{1}
\]

where \(T\) is the thermodynamic temperature in K, \(\tau_0\) is radiative lifetime, \(E\) the activation energy of the thermal-quenching effect, and \(k_b\) is the Boltzmann constant. By applying this theory, Yuen Pan et al. created \(\text{Eu}^{2+}/\text{Eu}^{3+}\) co-doped \(\text{Sc}_2\text{O}_3\) nanoparticles, which precisely monitor the temperature changing through the light emission intensity. The unique part of this study was using \(\text{Eu}^{2+}\) as the primary lattice host rather than \(\text{Eu}^{3+}\). Due to the lack of complete 5s and 5p orbitals protection, the \(\text{Eu}^{2+}\) electron configuration exhibits more sensitivity to the exterior environment, such as temperature [60]. Jianan Liu et al. integrate the upconversion nanoparticle with a commercial K+ indicator to detect the intracellular K+ level [61]. Chantal Galaup et al. evaluated a possibility of luminescent lanthanide material that could detect ROS in the context of Alzheimer’s disease [62]. Cyrille Charpentier et al. designed a particular ligand that could boost the brightness of the terbium nanoparticles and link the nanoparticle to different biological molecules to form specific biosensors to detect biotinylated dyes or even epidermal growth factor receptors (EGFR) [63]. For more biomedical applications, Shuqing He et al. found a unique property of DSPE-mPEG encapsulated rare-earth-doped nanoparticles, which surprisingly exhibited a high affinity to the bone. They used this particle for dedicated imaging of the bone and lymph nodes (Fig. 4A). The mechanism of this bone affinity is mainly caused by the hydroxyapatite mineral binding ability and white blood cell uptake [64]. Rather than laser excitation, lanthanide nanoparticles could also be excited by ionization radiation, like Cerenkov radiation, X-rays etc. [41, 42]. Neufeld et al., using X-ray as a radiation source, successfully excited nanoscale Ln-MOFs (Ln = Eu or Tb) that emit at 400-700 for radioluminescent cell imaging (Fig. 4F) [65]. Another benefit of lanthanide nanoparticles is the long luminescent half-life, eliminating the influence of autofluorescence from body tissue. Andreas Reisch’s group developed poly(methyl methacrylate)-based particles doped with a high percentage of europium complexes. By utilizing these rare earth nano complexes, they achieved background-free single-molecule imaging in live cells (Fig. 4D) [66]. By adjusting the doping percentage of different lanthanide element components, the emission of the lanthanide nanoparticle could easily be turned into the desired range. With this feature, the same wavelength excitation source could be used to achieve different emission wavelengths for multichannel or multiplexed imaging (Fig. 4B, E) [67, 68].

In an innovative and unusual exploration, Zhiming Deng et al. fed silkworms with \(\text{NaYF}_4\text{:Gd}^{3+}/\text{Yb}^{3+}/\text{Er}^{3+} @ \text{SiO}_2\) nanocrystals to produce a NIR-II fluorescent silk fibroin-based material (Fig. 5B). They further bioengineered the upconversion silk by attaching it to an implantable stent for mice to verify the potential for \textit{in vivo} applications [69].

\textbf{Toxicity} remains a primary challenge for \textit{in vivo} inorganic nanomedicine, and the lanthanide nanoparticles are no exception [70]. Gadolinium chelates are frequently used as MRI contrast agents and recently have been identified as causing renal impairment through slowly developing nephrogenic systemic fibrosis (NSF) in humans that had received certain gadolinium-based MR contrast agents [71]. The FDA also issued a safety warning for gadolinium-based contrast agents chelated by linear chelators such as DTPA after observations that Gd could be retained inside the body, even in the brain, for months or even years after the MRI scan. Other studies have shown there is nephrotoxicity of gadolinium-based contrast agents [72]. Meanwhile, other lanthanide ions, like Tb\(^{3+}\), Pm\(^{3+}\) or Eu\(^{3+}\) could bind to the Ca\(^{2+}\) binding sites of the intestinal membrane, affecting ion channel transportation, leading to toxicity. Sc\(^{3+}\), Y\(^{3+}\), and La\(^{+}\) have been found to bind to globulin and DNA, which could cause damage [73]. Encapsulating lanthanides into rapidly excretable nanoparticles and/or coating with a biocompatible polymer which decreases the time of the lanthanide inside the body, can minimize toxicity. Dana M. et al. tested their \(\text{NaGdF}_4\text{:Tm}^{3+}/\text{Yb}^{3+}\) nanoparticles on a variety of cancer cell lines and C.elegans. detected no toxicity over a wide concentration range [74]. For the same kind of particle, Huaiyong Xing et al. showed after seven days post-injection to the mice, the \(\text{NaGdF}_4\text{:Tm}^{3+}\) ultra-small nanoparticle could be excreted into feces and urine [75]. Jin Yan et al. developed a safe peptide-lanthanide nanocluster for cancer drug delivery. They functionalized 5 nm lanthanide cores with two types of targeting peptide, which increased the tumor-targeting efficacy and decreased the toxicity of the nano cluster [76]. Based on the issue of toxicity and the development of strategies to eliminate concerns with introducing lanthanides into the body, the design of lanthanide nanomaterials for biomedical applications continues to focus on developing...
ultra-small and clearable nanoparticle approaches, while still seeking to take advantage of the available optical properties to enhance sensitivity for disease diagnosis and future innovation in treatment of disease [35, 37, 41].

**Conclusion and Prospects**

Biomedical imaging is a multidisciplinary research field involving physics, chemistry, molecular biology, photonics and materials science. In this rapidly developing research field, new materials, new technologies, and new applications arise quickly. Due to the long time for clinical translation, it is essential that biomedical researchers become aware of new materials opportunities that enable improved sensitivity in diagnostic medicine, and increased efficacy/reduced toxicity in therapeutics. This review provided a short summary of the applications of lanthanide nanomaterials published over the past few years, including cell imaging, living tumor diagnosis, lymphatic and vascular imaging, and imaging guide surgery. Given the extensive literature on this global research topic, it is difficult to cover all aspects of rare earth nanomaterial science and biomedical applications. Lanthanide luminescent nanomaterials have made outstanding contributions to the field of biomedicine in the past few decades, mainly due to the apparent advantages of rare earth luminescent materials, such as stable photochemical properties, negligible photobleaching, and high fluorescence to noise ratio. The excitation light wavelength is in the ideal biological window (NIR region), which has better penetration, no damage to the normal tissue and high sensitivity. Because of the unique properties of rare earth luminescent materials, they could provide advantages that other probes cannot achieve and a valuable technical method to biologists and physicians. Optical imaging of rare earth materials is essential for various technologies required for modern bioanalysis and bioimaging. It has allowed for the creation of unusual nanosensors based on the unique properties of the lanthanides. Overall, lanthanide-based agents have shown remarkable application prospects in fluorescence imaging, medical diagnosis, and treatment.

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**Figure 5:**

A) Microscopy images (recorded in the Eu³⁺ channel) of the MDA-MB-468 cells incubated with DNPD-based polymeric micelles. i) The blue and red points mark, respectively, the locations used for temperature determination of the darker and brighter regions. ii) Temperature histograms obtained from the selected points indicated in panels B. i) A schematic illustration of the in vivo NIR-II imaging of the implanted biological stent model made from the hybrid silks. ii) Allergy test of a mouse by directly exposing the abdomen skin to the hybrid silk under 980 nm laser excitation. iii) The corresponding fluorescence intensity. iv) A digital photograph and in vitro phantom NIR-II imaging of other phantom biological stents with “H”, “N”, and “U” shapes. vi and vii. In vivo NIR-II imaging after subcutaneously implanting the models in the mouse abdomen under the excitation of a 980 nm laser. C) i) NIR-II fluorescence bioimaging of the acute local epidermal inflammation with various nanoprobes. ii) SNR as a function of time in the inflamed area. iii) In vivo I-PEM bioimaging. iv) I-PEM signal intensity as a function of time in the inflamed area with various nanoprobes. v) ICP-MS results of the retention of various nanoprobes in the inflamed area. vi) Pharmacokinetics of DCNP@GSH in blood and urine excretion after iv) injection. ICP-MS results of kidney biodistribution as a function of time. Insert shows NIR-II bioimaging of the bladder. vii) Excretion of DCNP@GSH and DCNP@OH in urine and feces as a function of time. h) Biodistribution of DCNP@GSH and DCNP@OH nanoprobes in the liver (Li), spleen (Sp), and kidney (Ki) after two weeks [37-39].
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Competing Interests

The authors have declared that no competing interest exists.

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