Recent Progress in NIR-II Contrast Agent for Biological Imaging

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Fluorescence imaging technology has gradually become a new and promising tool for \textit{in vivo} visualization detection. Because it can provide real-time sub-cellular resolution imaging results, it can be widely used in the field of biological detection and medical detection and treatment. However, due to the limited imaging depth (1–2 mm) and self-fluorescence background of tissue emitted in the visible region (400–700 nm), it fails to reveal biological complexity in deep tissues. The traditional near infrared wavelength (NIR-I, 650–950 nm) is considered as the first biological window, because it reduces the NIR absorption and scattering from blood and water in organisms. NIR-II fluorescence bioimaging’s penetration is larger than that of visible light. In fact, NIR-I fluorescence bioimaging is still interfered by tissue autofluorescence (background noise), and the existence of photon scattering, which limits the depth of tissue penetration. Recent experimental and simulation results show that the signal-to-noise ratio (SNR) of bioimaging can be significantly improved at the second region near infrared (NIR-II, 1,000–1,700 nm), also known as the second biological window. NIR-II bioimaging is able to explore deep-tissues information in the range of centimeter, and to obtain micron-level resolution at the millimeter depth, which surpass the performance of NIR-I fluorescence imaging. The key of fluorescence bioimaging is to achieve highly selective imaging thanks to the functional/targeting contrast agent (probe). However, the progress of NIR-II probes is very limited. To date, there are a few reports about NIR-II fluorescence probes, such as carbon nanotubes, \textit{Ag}_{2} \textit{S} quantum dots, and organic small molecular dyes. In this paper, we surveyed the development of NIR-II imaging contrast agents and their application in cancer imaging, medical detection, vascular bioimaging, and cancer diagnosis. In addition, the hotspots and challenges of NIR-II bioimaging are discussed. It is expected that our findings will lay a foundation for further theoretical research and practical application of NIR-II bioimaging, as well as the inspiration of new ideas in this field.

Keywords: fluorescence imaging technology, the second region near infrared (NIR-II), biological imaging, contrast agents, biomedical applications
INTRODUCTION

Optical imaging technology has become a very important research method in the field of biomedicine, especially for its ability to monitor biomolecules, cells, tissues, and living organisms in real-time and multi-dimensional visualization. Optical imaging technology has many advantages, in that it is non-invasive and safe, has visualization capabilities and high spatial resolution, has high rapid output, and is a low-cost method. It has been widely used in biomolecular detection imaging, drug distribution metabolic tracking, disease detection and diagnosis (Sevick-Muraca et al., 2002; Hilderbrand and Weissleder, 2010; Chen et al., 2014). Especially in early cancer diagnosis and imaging-guided treatment, optical imaging technology have a good application prospect.

In the process of optical in vivo bioimaging, the penetration depth of photons primarily depends on the absorption and scattering of tissue elements. Meanwhile, the fluorescence and scattering photons generated by tissue itself will cause interference noise and background radiation to the photon penetration process. Therefore, fluorescence imaging technology also has limitation in its practical applications in that some active components in organisms (such as melanin, hemoglobin, cytochrome, etc.) have higher light absorption and light scattering within the visible band (400–700 nm), which will reduce the penetration depth of visible light. Because organisms are rich in many luminescent macromolecules (usually located in the visible region), these biomolecules can also produce non-specific fluorescence emissions under visible light excitation, thus interfering with imaging results (Weissleder, 2001).

Because near-infrared light (NIR) is less absorbed and scattered in biological tissues than visible light, the former can penetrate biological tissues, such as skin, more effectively. As shown in Figure 1A (top), the effective attenuation coefficients (EACs) of tissue components, such as whole blood (oxygenated or deoxygenated), skin, and fat are significantly lower in the range of 650–950 nm than that in visible light range, which is considered as the “first (optical) window” (NIR-I) (Frangioni, 2003; Smith et al., 2009). In the practice of NIR-I biological imaging, the imaging quality in deep tissue is far from adequate resolution. Due to the large amount of background noise generated by tissue’s auto-fluorescence, the tissue penetration depth of photons in NIR-I is only 1–2 cm (Croce and Bottiroli, 2014).

Typically, the NIR-II window that may be suitable for bioimaging application is from 1,000 to 1,700 nm. Tissue components show a strong EAC from 1,350 to 1,700 nm, as shown in Figure 1A (top). We herein narrowed down the wavelength range to 1,000–1,350 nm, as labeled as “second window” in Figure 1A (top). In this review, the “NIR-II” in this manuscript only refers to the wavelength range of 1,000–1,350 nm. The relationship between wavelength and the absorption coefficient or the light scattering coefficient in the skin is shown in Figures 1B, C respectively. The absorption to the skin in NIR-I decreases slowly with the increase of wavelength, and the trend continues into NIR-II. It reaches the lowest point at ∼1,125 nm and starts spiking after 1,250 nm. No significant difference is found for the light scattering coefficients in NIR-I and NIR-II regions. The penetration depth of light to the skin can be seen from Figure 1D, from which we could find a sweet spot (largest penetration depth) of 1,125 nm in NIR-II. Overall, the range of 1,000–1,125 nm in NIR-II shows a better performance than NIR-I for skin. We may get the similar conclusions for human subcutaneous fat tissue (Figures 1E, F) and human subcutaneous adipose tissue (Figures 1G–I). Again, it is difficult to compare directly the performance of NIR-I and NIR-II because the absorption coefficients in the latter shows a turning point in the middle of NIR-II (at ∼1,125 nm). The sweet spot of 1,000–1,125 nm of NIR-II shows the lowest absorption coefficients for both red blood cells (RBCs) and hemoglobin with or without saturated oxygen, which is also applicable for water (see Figure 1J). In fact, it can be seen from Figure 1J of the relationship between water absorption coefficient with wavelength that the wavelength below 800 nm is the least absorbed by water, which is the optimal excitation wavelength. The closest point in NIR-II to eliminate the absorption from water is at 1,100 nm. It also can be seen from Figure 1K that the effective light scattering of 100% and 0% oxygen saturation RBCs shows no significant differences in NIR-I and NIR-II. However, the autofluorescence from tissue or endogenous compounds limits the signal-to-noise ratio (SNR) of NIR-I fluorescence imaging. The SNR of NIR-II fluorescence imaging, on the other hand, is enhanced due to the elimination of autofluorescence noises. For example, Lim et al. (2003) studied the optical imaging process in different biological media (including tissue and blood) and simulated the process and established relevant models. From the results of this study, it can be concluded that the SNR of quantum dot fluorescent clusters emitting at 1,320 nm is more than 100 times higher than that of quantum dots emitting at 850 nm. This result aroused great interest of the researchers in exploring biocompatible NIR-II fluorescence probes. Therefore, compared with visible light (400–700 nm) and traditional NIR-I window, NIR-II can avoid background interference such as spontaneous fluorescence and photon scattering (Smith et al., 2009; Welscher et al., 2011). In 2009, Welscher et al. (2009) first reported on a class of single-walled carbon nanotubes (SWCNTs), whose fluorescence emission wavelength is 950–1,600 nm. Further experiments show that such material can be used for imaging in living mice. The fluorescence signal of SWCNTs at 950–1,400 nm can be detected to obtain high-resolution imaging of blood vessels under deep skin tissue. Since then, more and more NIR-II fluorescent probes have been successfully developed and have been applied to biomedical imaging field.

At present, the application of NIR-II imaging window is still not fully utilized, due to the lack of research on probe materials and the restriction of imaging equipment. The Figure 1A (bottom) shows the relationship between incident light wavelength and quantum efficiency when cameras with different sensors are used, including silicon (Si), InGaAs, and HgCdTe. Furthermore, Si camera is the most sensitive to the light of NIR-I, while InGaAs camera is the most sensitive to the light of NIR-II. In addition, HgCdTe camera is less sensitive to the light of these two windows although it has a higher resolution array, and...
Prahl et al. (1993) reported the inverse double increase (IAD) method and the power law approximation are used to process experimental data and determine the optical properties of tissues. In this figure, \( \mu_s' \) is calculated as \( \mu_s' = \mu_s(1-g) \), where \( \mu_s \) is the scattering coefficient and \( g \) is the anisotropic coefficient of scattering. The solid line corresponds to the average experimental data, and the vertical line shows the SD value. (A) NIR-I (first window) and NIR-II (second window) imaging windows (Smith et al., 2009). The effective attenuation coefficient represents how easily a volume of material can be penetrated by a beam of light. (A) Bottom) The sensitivity curves of the sensor in the signal detector camera with silicon (Si), indium gallium arsenic (InGaAs), and mercury telluride cadmium (HgCdTe). Unlike the charge coupled device (CCD) camera using silicon sensor, the core component of near infrared camera is semiconductor alloy sensor, including InGaAs and HgCdTe, which has a narrower band gap. In particular, InGaAs cameras exhibit high quantum efficiency when used in the NIR-II window, i.e., high sensitivity. Adapted from Smith et al. (2009) written by Smith, A.M., etc. with permission. (B,C) Show the relationship between the incident light wavelength and absorption coefficient \( \mu_a \) or the reduced light scattering coefficient \( \mu_s' \) in human skin in vitro, respectively. (B) Adapted from Bashkatov et al. (2005) with permission. In (C), except for the solid line, the remaining data marker points correspond to the experimental data obtained in reference (Chan et al., 1996; Simpson et al., 1998; Du et al., 2001; Troy and Thennadil, 2001; Bashkatov et al., 2005). Adapted from Chan et al. (1996), Simpson et al. (1998), Du et al. (2001), Troy and Thennadil (2001), and Bashkatov et al. (2005) with permission. (D,F) Show the relationship between wavelength and \( \mu_a \) or \( \mu_s' \) in subcutaneous adipose tissue, respectively. In (E), all the data markers except the solid line correspond to the experimental data obtained in Peters et al. (1990).
it is most sensitive to the light of longer wavelength. Silicon-based CCD (Si-CCD) is usually used to collect fluorescence signals from NIR-I window. However, the quantum yield of this kind of CCD in NIR-II window is low, which is not sufficient for signal acquisition. Near infrared CCD based on InGaAs is commonly used as a NIR-II window detector, but the high cost of use restricts its widespread use in research (Smith et al., 2009).

In recent years, there have been a few successful applications of Nanofluorescent probes developed by NIR-II technology in bioimaging, which include the use of carbon nanotubes (Hong et al., 2014; Diao et al., 2015), AgS quantum dots (Hong et al., 2012b; Zhang et al., 2012) and small organic molecules (Antaris et al., 2016). Next, the applications of various near infrared nanomaterials and bioimaging will be introduced in detail.

**CATEGORIES OF REPORTED NIR-II NANOMATERIALS**

**Single-Walled Carbon Nanotubes**

Semiconductor-based single-walled carbon nanotubes (SWCNTs) have unique optical properties due to the existence of the bandgap. When photons interact with SWCNTs, they are absorbed and released as fluorescence. It is worth noting that the absorption wavelength of SWCNTs is generally in the visible region (400–700 nm) and NIR-I (650–950 nm). However, their emission wavelength is in NIR-II (1,000–1,700 nm), and the energy from absorbed photons is not released by radiation relaxation in the form of heat (O’Connell et al., 2002). SWCNTs have been widely used in the fields of photothermal and photodynamic therapy (Robinson et al., 2010; Murakami et al., 2012), fluorescence labeling, and fluorescence imaging of deep tissue in vivo (Welsher et al., 2011; Hong et al., 2012a, 2014). In addition, the use of SWCNT surface functionalization (Zheng et al., 2003) using coating surfactants, polymers, DNA, proteins, and even viruses can greatly enrich the scope of application for SWCNTs in the biological field. However, the characteristically low fluorescence quantum efficiency (<1%) of SWCNTs is a major issue still needing to be addressed.

SWCNTs are used in vivo imaging because of their inherently wide NIR-II fluorescence emissions (Yi et al., 2012; Ghosh et al., 2014). In 2014, Belcher’s research team coated SWCNTs with an M13 virus, using the M13 virus surface polypeptide, which targets tumor cells, to specifically bind SWCNTs to tumor tissue. Using SWCNTs, Belcher was able to achieve targeted fluorescence imaging of tumor tissues in mice, which proved that the system could be applied to clinical diagnosis (Ghosh et al., 2014). In the same year, Hong et al. combined an IRDye-800 fluorescent group with SWCNTs and intravenously injected the IRDye-800 fluorescent group into mice. Using a NIR-II fluorescence imaging in vivo (Hong et al., 2014), they found that the fluorescence of carbon nanotubes could penetrate 2.6 mm deep into a mouse skull, which not only provided imaging of the distribution of blood vessels in the head of mice, but also clearly showed the fine structure of the brain capillaries. In 2019, Toshiya Okazaki and other Japanese scholars used oxygen-doped SWCNTs and enveloped it with phospholipid polyethylene glycol (o-SWCNT-PEG). They found that it has special potential and can emit 1,300 nm NIR-II fluorescence under 980 nm light excitation. Therefore, it is considered to be a promising angiographic imaging probe (Takeuchi et al., 2019). In addition, o-SWCNTs were injected intravenously into living mice as a contrast agent for vascular imaging. The biological factors of the angiographic probe in vivo, including retention time, biological distribution, and toxicity, were studied. The results showed that o-SWCNT-PEG, an angiographic probe, had low toxicity in vivo, and that it can be successfully used in angiographic imaging within the NIR-II wavelength range.

**Semiconductor Quantum Dots**

A second successful application of Nanofluorescent probes developed by NIR-II technology is the use of semiconductor quantum dots. Among the many kinds of semiconductor fluorescent quantum dots available, AgS quantum dots have been most widely used in near infrared imaging because of their strong NIR-II fluorescence and low biological toxicity. Their biological toxicity is lower than other quantum dots containing Se, Te, Cd, Pb, As, and other acute or chronic toxic elements, and the fluorescence can be tuned from 687 nm in NIR-I to 1,294 nm in NIR-II (Yang et al., 2013; Gui et al., 2014a,b; Zhao and Song, 2014). At the same time, it also has high fluorescence stability and fluorescence quantum efficiency. AgS quantum dots have smaller particle sizes [3.7 nm (Zhao and Song, 2014), 1.6–6.8 nm (Yang et al., 2013), 2.6–3.7 nm (Gui et al., 2014b), 3.5 nm (Gui et al., 2014a)], which are very suitable for bioimaging applications.

For example, in 2012 Wang’s and Dai’s research teams collaborated to report on NIR-II AgS quantum dots with a fluorescence of 1,200 nm. The solubility of AgS QDs was changed by using a surface modification of water-soluble PEG molecules. And then AgS QDs can be dissolved in water. AgS was first used in NIR-II cell imaging and non-specific tumor detection (Hong et al., 2012b). Additionally, in 2012, they applied PEG modified AgS quantum dots with high quantum efficiency to NIR-II imaging in vivo (Zhang et al., 2012), using nude mice inoculated with 4T1 tumors. Not only did the AgS quantum dots prove to be a good NIR-II fluorescence contrast agent, but interestingly, after injection into the tail of nude mice, AgS quantum dots gradually accumulated in the tumors of...
nude mice as the circulation time increases. They believe that the reason for above observation is the non-specific enhanced permeability and retention effect (EPR effect) of cancer tissue. Subsequently, Wang’s team did an in-depth study on the long period cytotoxicity of Ag$_2$S quantum dots in vivo (Zhang et al., 2013). The reticuloendothelial system (RES), such as liver and spleen, is the main accumulation site of Ag$_2$S QDs in vivo, but it can be gradually metabolized or excreted over time. Additionally, the results of the blood biochemical analysis and histological examination of the rats given the Ag$_2$S quantum dots for 2 months showed that Ag$_2$S quantum dots had no obvious toxicity.

**Nanoparticle Alloys**
In materials science, mixing various metal elements to form intermetallic compounds or alloys can greatly expand the properties of metals. Binary or ternary metal nanoparticles (or nanoalloys) can synthesize intermetallic compounds with controllable properties and structures on a nanoscale, which has attracted extensive attention from researchers (Ferrando et al., 2008). The main reason why alloy nanoparticles are fascinating is that the chemical and physical properties of alloy materials change with changes in composition, atom distribution, and particle size.

Researchers who have devoted themselves to in-depth studies of these metal nanoalloys have found that metal alloy materials not only have excellent catalytic properties, but also have excellent fluorescence properties. Further research shows that these fluorescent alloy nanoparticles inherit many advantages of the original metal nanoparticles, such as fluorescence properties, water solubility, and biocompatibility. They also can adjust the optical properties of the main metal nanoparticles or develop new functions through the introduction of another metal. For example, in 2013, Millstone’s team (Andolina et al., 2013) introduced copper into fluorescent Au nanodots to form Au/Cu alloy nanodots. By adjusting the content of copper in the alloy nanodots, the fluorescence of Au/Cu alloy nanodots gradually shifted from NIR-I to NIR-II. Their group further introduced the Co element into fluorescent Au nanodots to prepare Au/Co alloy nanodots, which had both magnetic and near infrared fluorescence tunable functions (Marbella et al., 2014). Compared to the traditional near infrared quantum dots, the multifunctional alloy nanodots with near infrared fluorescence not only free of toxic elements, such as heavy metals, but also have many functions in the same material. These mutli-functional alloy nanodots have wide prospects for application in bioimaging, especially in the field of multi-mode imaging.

**Down-Conversion Rare Earth Nanoparticles**
Recent research results have shown that rare earth nanomaterials have the ability of down-conversion luminescence, which is, when near-infrared light (980 nm) is used as excitation light to irradiate nanoparticles, its emission light is in the NIR-II spectrum range (Tan et al., 2009). The structure of these nano-materials is very similar to that of up-conversion nanoparticles. Nanocrystals such as lanthanide nanocrystals (NaYF$_4$) are used as primary materials, which doped with lanthanide elements such as Ho$^{3+}$, Tm$^{3+}$, and Nd$^{3+}$ (Naczynski et al., 2013; van Saders et al., 2013). Using the basic structure of rare earth nanocrystals, multi-shell complex core-shell rare earth nanoparticles have been developed. For example, in 2013, Moghe team reported on a core-shell structure nanoparticle of NaYF$_4$ which had been regenerated on the surface of existing NaYF$_4$ Yb:Ln nanocrystals (Naczynski et al., 2013). By changing the doped lanthanides (Er, Ho, Tm, Pr), NIR-II fluorescence with different wavelengths can be obtained under 800 nm excitation, and the longest wavelength can reach up to 1,500 nm. Compared with up-conversion rare earth nanoparticles, down-conversion nanoparticles have received less attention, but they are still widely applied in vivo bioimaging research (Naczynski et al., 2013, 2014; Jiang et al., 2016). Li et al. reported on a dual-function particle system using up-conversion and down-conversion (Li et al., 2013), where the particle can emit visible (800 nm) and near-infrared light (980 nm) according to the excitation light of different wavelengths and can be used in biological imaging.

In recent years, it has been found that Nd ion not only has NIR-II fluorescence emission with wavelength of 1,050 or 1,300 nm, but also has the sensitization effect on Ytterbium ion. It is excited at a biocompatibility wavelength below 800 nm, which has the lowest absorption of water. Nd$^{3+}$ has been recognized gradually as a photosensitizer (Hammer et al., 2016), and Nd$^{3+}$ doped near infrared fluorescent nanoparticles have attracted great interest from researchers who study lanthanide-doped biological probes. One kind of Nd$^{3+}$ doped near infrared fluorescent materials is an NaYF$_4$ type down-conversion lanthanide nanocrystal (Chen et al., 2012), and the other is an Nd$^{3+}$ doped fluoride nanoparticle (Pokhrela et al., 2014). For example, in 2015, Garcia’s group synthesized Nd$^{3+}$ doped SrF$_2$ nanoparticles and used them reduce background fluorescence (Villa et al., 2015), since minimizing any background fluorescence is essential for high contrast bioimaging. The excitation wavelength of the nanoparticles is 808 nm, and the emission wavelength is 1,100 nm, which belongs to NIR-II fluorescence. The nanoparticles are mixed into the feed and fed to mice. After the nanoparticles enter the mice through the digestive system, in vivo fluorescence imaging is performed. It was found that an NIR-II fluorescence of 1,300 nm generated by the $^4F_{3/2}$-$^4I_{13/2}$ orbital transition of Nd$^{3+}$ effectively eliminated the background fluorescence. It can realize the NIR-II imaging of deeper structure, inorganic spontaneous fluorescence and high distinguishability in vivo. In addition, in 2016 Prasad’s team prepared a hybrid organic-inorganic system to form an epitaxy of NaYF$_4$:Yb$^{3+}$/X$^{3+}$@NaYF$_4$@NaYF$_4$:Nd$^{3+}$ (X = null, Er, Ho, Tm or Pr) core/shell/shell (CSS) nanocrystals, and coated ICG on the external layer of CSS nanocrystals. This hybrid system can capture NIR light in a wide excitation spectrum range (700–860 nm) by ICG and produce effective polychromatic narrowband NIR-II emission light from 1,000 to 1,600 nm according to the different doping elements in the nucleus. Further experiments show that the NIR-II emission fluorescence can be used to image clearly at a tissue depth of 9 mm and detect optical signal at a tissue depth of 23 mm (Shao et al., 2016).
Nanoparticles Based on Organic Fluorescent Dyes

Until now, organic fluorescent dyes are still the most widely used luminescent markers in fluorescence imaging. Organic fluorescent dyes have attracted much attention due to their high fluorescence quantum efficiency, easy functionalization, and adjustable luminescence spectra (Thekkkek and Richards-Kortum, 2008; Willmann et al., 2008; Kobayashi et al., 2010; Sinkeldam et al., 2010). The commonly used organic fluorescent dyes presently include naphthalimides, coumarins, fluoresceins, rhodamine, anthocyanins, BODIPY (Lu et al., 2014), porphyrin phthalocyanine, and other macrocyclic molecules. The absorption and emission wavelengths of these commonly used organic fluorescent dyes cover the ultraviolet, visible, and near infrared regions (Mishra et al., 2000; Lavis and Raines, 2008; Ma and Su, 2010).

Over the years, how to convert NIR-I fluorescent probes directly into NIR-II fluorescent probes by molecular engineering methods and design principles has been a highly discussed topic among investigators. According to the available literature (Yang et al., 2017; Zhu et al., 2018, 2019; Wu et al., 2019), the specific design principles are as follows: (1) The typical structure of NIR-II small molecule fluorescent dyes is a molecular structure composed of donor-acceptor-donor (D-A-D) (e.g., CH1055-PEG), which is modified to form a class of dyes with similar properties and emission wavelengths. The specific emission principle uses aromatic π-bridge connectors as molecular wiring. These electron-supplying groups can produce enhanced electron shifts and low energy gaps to the central electron acceptor, resulting with the dye molecule having the capability of NIR-II emission. (2) By systematically adjusting the electronic donor part, the π-bridge connector and the functional groups at the end of the fluorescent dye molecule promote quantum yield. Specific strategies to improve the brightness of small molecular dyes include enhancing intramolecular charge transfer and molecular stiffness, creating complexes with serum proteins, such as human serum albumin (HSA), and introducing a protecting group (S) to produce a S-D-A-D-S structure at the end of the dye, which protects the dye's pillars from intermolecular interactions and fluorescence quenching polymerization. (3) By increasing the conjugate length to separate electron donor/acceptor and heteroatom substitution, the fluorescence emission of existing polymethine dyes (i.e., cyanine dye) can be re-shifted to produce a new fluorescent dye that can emit NIR-II fluorescence with high quantum yield and absorption coefficient (ε) (Figure 2). At present, polymethine molecular dyes with emission wavelength more than 1,000 nm are available on the market, including IR-1040, IR-1048, IR-1051, and IR-1061. Unlike the previous method of for red-shifting anthocyanin dyes by simply increasing the length of polymethine chain, by Cosco et al. (2017) propose of a new method of extending heterocyclic conjugation and adding new electron donor groups. This new method has been proven as a feasible method for red-shifting anthocyanin dyes. In addition, most of the NIR-II small molecule fluorescent dyes mentioned above are organic products with low water solubility, which need to be encapsulated in a hydrophilic matrix to enhance biocompatibility in vivo imaging. Recently, Li et al. (2018) developed a novel small molecule fluorescent dye FD-1080 through a structural redesign of a typical anthocyanin dye. Its excitation and emission wavelength are both in the range of NIR-II. Besides there are reports about indocyanine green (ICG). ICG is a NIR fluorescent dye with strong absorption, low toxicity, no involvement in biological transformation in vivo and rapid excretion. It is the only NIR optical imaging contrast agent approved by the US Food and Drug Administration (FDA) for clinical use. Because the absorption and fluorescence spectra of ICG are in the NIR-I window (650–950 nm), ICG has been widely used in cardiovascular system, liver function evaluation, visualization of retina and choroid, ophthalmic angiography, cerebral angiography, and other clinical fields. Recently Starosolski et al. (2017) found that ICG dye also has NIR-II fluorescence characteristics in the NIR-II window. This discovery has opened up new uses of ICG and will greatly expand the application scope of ICG. In the experiment, the absorbance and NIR-II fluorescence emission of ICG were detected in different concentrations of ICG media, including PBS, plasma and ethanol. The customized spectral NIR module is used for in vitro and in vivo tests, and the images of NIR-I and NIR-II windows are obtained at the same time. The results show that ICG has a significant fluorescence emission in the NIR-II window at the wavelength of about 1,100 nm, and this emission (similar to the absorption curve) is essentially affected by the molecular environment of ICG. The results of in vivo imaging further illustrated that ICG can be used as NIR-II fluorescent dye and the contrast-to-noise ratios (CNR) value was twice that in NIR-I window. Clinical transformation of NIR-II imaging technology can be accelerated when ICG, an FDA approved imaging agent, is used. Later, the research team published another paper on NIR-II fluorescence imaging using indocyanine green nanoparticles (Bhavane et al., 2018). In this study, they collected the fluorescence spectra of ICG liposomes in PBS and plasma. In vivo imaging research was carried out to observe the vascular structure of the hind limbs and the intracranial region in real time. Free ICG, NIR-I imaging and cross-sectional imaging (MRI and CT) were used as controls. The results showed that the liposome ICG had strong NIR-II fluorescence, similar to the free ICG in plasma. In vitro studies have shown that liposome ICG has better performance than free ICG in the NIR-II imaging of deep (≥4 mm) vascular analog structure. In vivo, NIR-II fluorescence imaging with liposome ICG can significantly improve the contrast (P < 0.05) compared with long-term free ICG, and make the hind limbs and intracranial vessels visualized within 4 h after injection. Compared with NIR-I imaging, liposome ICG enhanced NIR-II imaging has better vascular clarity. Subsequently, several research papers on tumor imaging using Second-Window-ICG (SWIG) technique were published, and this field has become the latest research hotspot of NIR-II imaging (Zeh et al., 2017; Cho et al., 2019a,b; Suo et al., 2019; Wang et al., 2019b).
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FIGURE 2 | Increasing the length of polymethine chain was proved to result in the red shift of cyanine dye’s emission. Cosco et al. developed new methods of heterocyclic conjugation and added new electron donor groups (Cosco et al., 2017; Zhu et al., 2018). (A) Molecular structure of cyanine dye series of compounds for optical fluorescence imaging. (B) The blue structure is the compound of dimethyl-flavylium heterocycles, which can be used to replace the indolenines to prepare flavylium polymethine fluorophores. (C) The emission and absorption of Flav7, the modified organic small molecule fluorescent dye, was in the NIR-II window. Adapted from Cosco et al. (2017) and Schnermann (2017) with permission.

APPLICATIONS OF NIR-II WINDOW BIOIMAGING IN BIOMEDICINE

Tumor Imaging and Image-Guided Surgery

Solid tumor patients often fail in treatment due to local recurrence. Cytopenic surgery is targeted to improve staging and reduce tumors using fluorescence imaging to guide the specific resection of tumors and significantly improve the prognosis. Recently, this application field has become a high-interest topic in research, resulting in the publishing of many high-level papers.

Papers published by Wang et al. (2018) in Nature Communications, for example, report on a novel down conversion nanoparticle emitted in the NIR-II window for living assembly. In metastatic ovarian cancer, fluorescent nanobeads modified with DNA and targeted polypeptides are used to ensure the precision of cytoreduction surgery. The image distinguishability of NIR-II nanoparticles is better than that of clinically recognized ICG, mainly due to its better light resistance and deeper tissue penetration (8 mm). After the nanoprobe is assembled in vivo, the stable preservation time of the nanoprobe on tumors is up to 6 h, so the nanoprobe can be used in precise tumor resection surgeries. The probe successfully obtained a superior ratio of tumors to normal tissues, which can help to delineate the resection profile in the surgery for metastatic abdominal ovarian cancer. The results also showed that metastases <1 mm could be completely removed under the guidance of NIR-II bioimaging. This novel method provides a new idea for the design of nanomaterials for medical applications in the future.

The key to the implementation of above-mentioned study is the in vivo assembly of novel NIR-II fluorescent nanoparticles, which is also the ingenuity of this research design. The specific assembly principle is shown in Figure 3. At the top of Figure 3 is a schematic of the step-by-step construction of DNA and FSHβ modified down-conversion nanoparticles (DCNPs-L1-FSHβ). The core of the nanoprobe are lanthanide doped core-shell nanoparticles, consisting of a 5.0 nm luminescent gadolinium tetrafluoride sodium doped 5% neodymium (NaGdF4:5% Nd) encapsulated by a 2.5 nm thick homogeneous gadolinium sodium inert shell. This type of a structure can avoid fluorescence quenching by water and bind complementary DNA (L1 or L2) on its surface. FSHβ is a follicle stimulating hormone peptide that specifically binds to ovarian cancer epithelium to enhance the targeting efficiency of the fluorescent nanoprobe. Because of the great potential of DNA complementary strand hybridization, the nanoprobe can be stable hybridized on the surface of tumors in vivo for bioimaging. Next, a process diagram of the fluorescent nanoprobe assembly in vivo is presented in the lower part of Figure 3. It greatly enhances the tumor targeting of the probe, but also enables the probe to be excluded from the body by the liver and kidney more quickly after the operation. The specific process is to implement two-step injection of DCNPs-L1-FSHβ and DCNPs-L2-FSHβ, in succession. In order to verify the performance of two-step in vivo simultaneous assembly of fluorescent nanoprobes, the fluorescent dye Cy5 was grafted into the probe assembly structure of the first needle, i.e.,
FIGURE 3 | Construction of NIR-II nanoprobe schematic diagram in surgery for metastatic ovarian cancer guided by NIR-II bioimaging (Wang et al., 2018). The diagram above shows the preparation of DCNPs (DCNPs-L1-FSHβ) modified by DNA and FSHβ. The schematic diagram below shows the further assembly of the NIR-II nanoprobe in vivo by a two-step sequential injection of DCNPs-L1-FSHβ (the first injection) and DCNPs-L2-FSHβ (the second injection), which is beneficial to improve tumor targeting and rapid liver and kidney clearance. Under the guidance of NIR-II imaging, metastatic ovarian tumors can be clearly observed and accurately removed. Adapted from Wang et al. (2018) written by Fan Zhang etc. with permission.

DCNPs-L1-(Cy5)-FSHβ, and Cy7 were combined to the probe assembly structure of the second needle, i.e., DCNPs-L2-(Cy7)-FSHβ. When the complementary DNA fragments L1 and L2 in the two components of the probe are hybridized close to each other, the two dyes will cause fluorescence resonance transfer (FRET), which is more than twice as strong as two dyes without FRET. The results show that the two components of the probe can be specifically assembled on the surface of tumors in vivo. Ultimately, surgery guided by NIR-II imaging can clearly observe metastatic ovarian cancer and precisely resection it.

Other recent representative work includes the development of Ag2Te quantum dots by Zhang et al. (2019), which can emit fluorescence at 1,300 nm wavelength after assembly. The assembly process is divided into two steps; first, polymerization is conducted with polylactic-co-glycolic acid (PLGA), and second, further packaging is completed with the cancer cell membrane. The fabricated biomimetic nano-biological probes have a bright and highly stable fluorescence in the NIR-II window. Through the second step of surface encapsulation of active homologous tumor membranes, the probe enhances the ability to target tumors. Tumor targeting occurs through the passive enhancement of permeability and retention of homologous cancer cell membranes, which can increase the accumulation of probes in the tumor sites. Zhang et al. successfully prepared a novel biomimetic NIR-II fluorescence nanoprobe which showed characteristics of ultra-bright, stable fluorescence, homologous targeting and high biocompatibility, which can significantly enhance the imaging of living tumors. In 2018, Yang et al. (2018) reported on a NIR-II lanthanide complex (Nd-DOTA) probe that can be rapidly excreted. Within
3 h after injection, more than 50% of the probe can be excreted through the kidney. The molecular weight of the probe is only 0.54 KDa, and in terms of light resistance and tissue penetration, the NIR-II imaging quality of the new probe is much better than that of the clinically approved ICG. Yang et al. also demonstrated the ability to obtain an excellent tumor-to-normal tissue ratio, which can help to accurately mark the profile of micro-tumors in surgery for abdominal metastatic ovarian cancer. Metastatic tumors <1 mm can be completely removed under the guidance of NIR-II bioimaging. In addition, the probe Nd-DOTA has the same structure as Gd-DOTA, and Gd-DOTA is a clinically recognized MRI contrast agent. Therefore, it may be a straightforward path for the clinical transformation of the new NIR-II probe. In 2018, Shou et al. (2018) prepared a semiconductor polymer nanoparticle of diketopyrrolopyrrole (PDFT1032) and developed it as a NIR-II (near infrared window 1,000–1,700 nm) fluorescent probe for in vivo tumor imaging and image-guided tumor re-sectioning. The NIR-II probe has many advantages, including stable fluorescence emission, high absorption efficiency at 809 nm, large Stokes shifting, biocompatibility, and lower toxicity in vivo. Moreover, research has shown this NIR-II probe has a wide range of applications in the biomedical field. For example, tumor imaging in terms of subcutaneous osteosarcoma patterns, evaluating vascular embolization treatment on tumors, in-situ cytoreduction surgery guided by NIR-II images, and sentinel lymph node biopsy with superior temporal-spatial resolution (SLNB). In general, PDFT1032 not only has good biocompatibility and hydrophilicity, but also excellent chemical and optical properties. The NIR-II fluorescent probe has a wide application prospect in the imaging of malignant tumors and cytoreduction surgery.

**Medical Testing**

Fluorescence bioimaging can detect deep tissues in the NIR-II window, and it has the smallest self-fluorescence and tissue scattering. Nevertheless, in vivo NIR-II fluorophores detection is merely concentrated on direct disease lesions or living organ imaging, and producing real-time dynamic NIR-II fluorescence biosensor has been challenging. In recent years, new methods and breakthroughs have been emerging in this field, such as the conception and progress of quantitative detection with fluorescent probe of ratio meter.

In Fan Zhang’s paper (Liu et al., 2018) published in Angewandte Chemie International Edition in 2018, he reports a novel Erbium-sensitized up-conversion of nanoparticles, which have the characteristics of both 1,530 nm excitation and 1,180 nm emission in the NIR-II window, which can be used in vivo biosensors. This research studies a micro-manipulation needle debris sensor for in vivo inspection of inflammation in real-time, where the detection principle of the sensor is based on the ratio meter fluorescence combined with efficient NIR-II fluorescent emission and the organic probe sensitive to hydrogen peroxide under the action of Fenton catalyst Fe$^{3+}$. Finally, the dynamic detection of inflammation in vivo is successfully produced. This NIR-II radiometric probe has the following advantages: large anti-Stokes displacement, low background fluorescence, low absorption, and scattering in biological tissues. Therefore, the probe can detect and evaluate inflammation in vivo in real time with high imaging quality. The specific process of in vivo inspection of inflammation in real time by the fluorescence sensor of the ratio meter is shown in Figure 4, including the device diagram of fluorescence probe used for confirmatory testing in vivo, the fluorescence value of the ratio meter ($I_{980}/I_{1180}$) channel at different times, the quantitative curve of ratiometer fluorescence ($I_{980}/I_{1180}$), and the corresponding concentration of hydrogen dioxide at different times.

Figure 4A is the device diagram of the ratiometer fluorescent probe used for confirmatory testing in vivo, and Figure 4B is a photograph of a mouse treated with fragments of a micromanipulator needle. Figures 4C–E show the 980, 1,180, and ratio gauge ($I_{980}/I_{1180}$) channels, respectively. The micromanipulator needle fragments obtained fluorescence values at different times after LPS-induced inflammation. Figure 4F is a quantitative curve of the ratio meter fluorescence ($I_{980}/I_{1180}$) of the micromanipulation needle fragments and the corresponding concentration of hydrogen peroxide at different times. It can be used to quantify the degree of actual inflammation in clinical tests.

Other recent studies in this field include Zhao et al. (2019) which reported on an original method for the accurate imaging of inflammation in vivo using in situ cross bonding of glutathione-combined ultrafine lanthanum nanoparticles with NIR-II fluorescent emission. Although nanoparticles have been proven to be promising bioimaging platforms by reason of their EPR effects, to the inability to enrich nanoprobe at target position has been a key bottleneck to improve detection ability and effect. While cross bonding of nanoparticles in vivo can increase the enrichment of EPR region (e.g., inflammatory areas), nanoparticles are absorbed by RES, resulting in unidirectional cross bonding in non-target organs. Based on these difficulties, the principle of this strategy was developed to enhance the in vivo imaging by using sub-10 nanometer glutathione (GSH) combined lanthanide nanoparticles, which react with reactive oxygen species (ROS) in the inflammation region to locate and image reactive oxygen species rapidly in the NIR-II window. At the same time, these nanoprobes can be excreted quickly because of their ultrafine magnitude. Based on the in-situ cross bonding and rapid excretion ability of the probe, this method can achieve accurate biological imaging and is suitable for other ultrafine contrast agents.

In 2019, Wang et al. (2019a) developed a reverse quenched NIR-II molecular fluorophore and applied it to high contrast imaging and pH sensing in vivo. Currently, the molecular fluorophore with contrast and sensitivity in NIR-II window (1,000–1,700 nm) have both been developed in vivo fluorescence imaging. However, the solvation of long-wavelength absorbed fluorophore in aqueous solution, results in quenching, which is challenging to avoid. Therefore, a series of reverse-quenched pentamethine cyanine fluorophore have been developed, which significantly overcome the serious solvation coloration, thus providing a stable absorption/emission of fluorescence over 1,000 nm in aqueous solution. The fluorescence intensity increases up to ~44 times and has excellent photostability. The conditions for lymphatic imaging can be met thanks to these
advantages, including the need for tissue penetration up to 8 mm, high definition, and optical stability. Its imaging effect is better than the clinically approved indocyanine green. In addition, the fluorophore display pH-responsive fluorescence allows for non-invasive ratio meter fluorescence imaging and gastric pH quantification in vivo. The results show that this method is consistently accurate when tissue depth is 4 mm compared to the standard pH electrode method. This work opens up the potential of reverse quenching pentamethylene cyanine in NIR-II biological applications.

In 2019, Li et al. (2019c) developed a NIR-II fluorescence molecular probe activated by peroxynitrate for drug-induced hepatotoxicity detection. Drug-generated liver injury is a key problem to the safety of drug research and use. The emergence of peroxynitrite (ONOOC−) is considered to be an initial signal of drug-generated liver injury. Therefore, the structure of the fluorescent probe is designed as the combination of benzothiophenacyl cyanine skeleton and phenyl borate group. The NIR-II fluorescence of this probe can be activated by ONOO− and can detect ONOO− sensitively. When the probe IRBTP-B and the target ONOO− exist at the same time, the structure of probe changes to produce the fluorescent group IRBTP-O to turn on the NIR-II fluorescence. The linear relationship between the NIR-II fluorescence intensity and the concentration of ONOO− is acceptable. Tissue model studies confirm that reliable activation signals can be obtained at penetration depths up to 5 mm. With this probe, the up-regulation of ONOO− in the model of incubation period drug-generated hepatotoxicity and the repair effect of N-acetylcysteine (NAC) in vivo could be seen. In conclusion, this method will
be used as a general method to develop activated NIR-II probes triggered by specific analytes based on hydroxyl functionalization reaction sites.

**Vascular Biological Imaging**

The number of people with cardiovascular and cerebrovascular diseases has increased, as has the ability to detect these diseases using examination technology, which, in turn, has led to an increase in incidence of cardiovascular and cerebrovascular diseases. The commonly used techniques in clinical angiography include B-mode ultrasonography, CT scans, and nuclear magnetic resonance (NMR). However, these techniques are non-tumor-specific, have low-sensitivity and high-cost, and can’t be used for real-time detection or clinical practice in surgical operations. In recent years, some research groups have designed NIR-II fluorescence probes and injected them into mice intravenously, after which the NIR-II fluorescence is used to perform vascular imaging *in vivo*. The results showed that the NIR-II fluorescence could penetrate the skull of mice to a deeper level, not only imaging the distribution of blood vessels in the heads of mice, but also clearly showing the fine structure of capillaries.

In 2018, Li et al. (2018) published papers within this field of study in the Angewandte Chemie, regarding the successful synthetization of a small molecule fluorescent probe FD-1080, which has excitation and emission both in the NIR-II region and it has excellent bioimaging capability of the vascular system *in vivo*. The structure of heptamethine in fluorescent probe FD-1080 has been ingeniously designed as a key group for the conversion of absorption and emission into the NIR-II region. An acidic sulfuric acid group and cyclohexene group were applied to intensify the water dissolvability and chemical durability of the probe. The quantum yield of FD-1080 was 0.31% and improved to 5.94% when combined with bovine fetal serum (FBS). It is noteworthy that compared with the excitation wavelength from 650 to 980 nm in NIR-I region in the previous work, the excitation wavelength at 1,064 nm in NIR-II region has proved to have stronger penetration in biological media and excellent imaging quality. FD-1080 can not only achieve non-invasive, high-resolution, in-depth tissue bioimaging of the vascular and cerebrovascular systems, but it also quantifies respiratory frequencies of awake and anesthetized mice according to dynamic images of respiratory cranio-caudal motion within the liver. The key steps in this study included the schematic diagram of the vascular imaging device, the fluorescence imaging of the rat cerebral vascular system with the NIR-II fluorescence probe FD-1080, and the breathing rate detection graph of the rat, as shown in Figure 5.

**Figure 5A** is a device diagram of NIR-II fluorescence penetrating the scalp and skull brain tissue for an angiography. **Figure 5B** is the disappearance spectra of the scalp (black) and skull (red). It has been found that the strongest penetration is at 1,064 nm. **Figure 5C** contrasts the fluorescence images of FD-1080-FBS complex in a rat brain vasculature system under different excitation lengths (1,300 nm long pass filter), where it was found that the clearest angiogram was at the wavelength of 1,064 nm in NIR-II region. In **Figure 5D**, a red line of interest is marked at the Gaussian fitting fluorescence intensity profile at the 808 and 1,064 nm excitation wavelengths, respectively. It is obvious that at 1,064 nm the fluorescence intensity is higher and the peak is narrower. When stimulated at 1,064 nm (1,300 nm long pass filter), the clear emission of the FD-1080-FBS complex enables the imaging of an alert or anesthetized mouse to detect the signal fluctuations produced by liver movement (Figure 5E). **Figure 5F** is a spectrogram of the breathing rate of alert (above) and anesthetized (below) anesthetized mice. The graph shows that there are obvious differences in the spectrogram of breathing rate between the two groups of mice, which can be used to quantify the respiratory rate of mice under different conditions.

Other studies have also been recently conducted in this field, including Li et al. (2019a) in 2019, who developed a rare earth nanoprobe triggered by 808 nm laser, which emitted NIR-II fluorescence for small tumor detection and angiography. In this study, polyacrylic acid (PAA) modified sodium tetrafluorolumetum: gadolinium/neodymium nanorods (PAA-NRs) were prepared into single crystal hexagonal phase and unified magnitude, and then further developed as a highly sensitive NIR-II imaging probe for optical imaging navigated detection of tiny tumors, angiogenesis-related diseases, and angiogenesis diagnosis. The NIR-II emission wavelength with sodium tetrafluorolumetum: gadolinium as the main body can be easily adjusted by changing neodymium doping, which makes it hopeful that the emission center will have high optical stability at 1,056 and 1,328 nm. The probe has a high spatial resolution (~105 μm) for small blood vessels *in vivo* and can detect them clearly. The *in vivo* tracking experiments with time changes confirmed that PAA-NRs probe was mostly collected in the RES, and discharged from the body by the liver. Histological examination revealed that the hydrophilic nanorods had very low toxicity and great biological compatibility in living animals.

In the same year, Li et al. (2019b) reported a polydopamine-coated multifunctional lanthanide diagnostic agent, which can be used for angiography of vascular malformations and tumors, as well as photothermal therapy guided by imaging at wavelengths above 1,500 nm. NIR-II optical imaging with emission wavelengths >1,500 nm can be used as the next generation fluorescence imaging technology to guide the display technology of tumor vessels and vascular malformations, which can then be used for early tumor diagnosis and recognition of tumor-related vascular characteristics. This study is based on the core-shell structure of NaLuF<sub>4</sub> nanorods@polydopamine (NRs@PDA), which combines advanced NIR-II fluorescence imaging ultra 1,500 nm wavelength and photothermal (PTT) functions. This is a high-performance integrated nano-platform for diagnosis and treatment developed for visualization of tumor-related vascular malformations and imaging-guided photothermal therapy.

In 2018, Wan et al. (2018) developed a bright organic NIR-II fluorescent nanocluster (p-FE) for three-dimensional imaging of biological tissues, which can provide bright NIR-II fluorescent emission light with a wavelength >1,100 nm for non-invasive *in vivo* blood flow tracking in the rat cerebrovascular system. Moreover, p-FE can produce a layer-by-layer image of the
vascular system based on single-photon and three-dimensional confocal imaging in a fixed rat brain tissue. Its depth is up to 1.3 mm, and it has a spatial resolution of up to 10 microns. The study also completed in vivo dual-color fluorescence imaging in the NIR-II window, using p-FE emitted between 1,100 and 1,300 nm as a vascular imaging reagent, as well as single-walled carbon nanotubes (CNTs), which emitted more than 1,500 nm as a contrast agent to highlight tumors in mice. Ultimately, the results demonstrated excellent dual imaging of vascular system and tumors.

**Diagnosis and Treatment of Cancer**

Each imaging technology has its own unique advantages and inherent limitations. For example, fluorescence imaging has excellent sensitivity, but low tissue penetration and spatial resolution in turbid media. MRIs, CT scans, and ultrasound...
imaging have better spatial resolution, but their sensitivity is limited. The sensitivity of PET-CT scan is relatively high, but it can't provide the structural information of the imaging material. Therefore, multi-mode imaging, which integrates two or more imaging technologies, can supplement the imaging information and diagnose diseases with reliable accuracy. In order to meet the needs of multi-mode imaging, researchers usually combine fluorescent nanomaterials with functionalized small molecules or particles by doping, bonding and coating to make them multi-functional nanoparticles, which can be used in multi-mode biological imaging.

Deep tissue imaging within the NIR-II window has great prospects in physiological research and biomedical applications. However, the inhomogeneous signal attenuation in biological material limits the application of multi-wavelength NIR-II probes in the imaging of various biomarkers and cancer diagnoses. For example, Fan et al. reported in 2018 (Fan et al., 2018), that NIR-II nanoparticles doped with lanthanides have the ability to design the luminescence time for quantitative imaging in vivo after using multiplexing in the time domain. To achieve this, a systematic research method based on controllable energy transfer was designed to create an adjustable lifetime range of nanoprobes, which had a luminescence time spanning three orders of magnitude and had only one emission band. When the depth of the NIR-II nanoparticle probe in biological tissue reached 8 mm, the selected nanoparticle duration was continuously resolved, and the SNR was taken from the time node when light intensity was <1.5. Powerful time-length editors have been shown to be independent of tissue penetration depth, and in vivo multiplexing technology (IVM) has been used to diagnose tumor hypotye in living animals. The data in this paper are in good agreement with those of a standard ex vivo immunohistochemistry assay, which proves that the luminescent time-length imaging can serve as a microinvasive method for tumor definite diagnosis. The research process and key points of IVM for tumor biomarkers using specific luminescence time-length imaging are shown in Figure 6.

Figure 6A illustrates the schematic diagram of conducted animal experiments. Three batches of Erbium nanoparticles were bound to three different kinds of antibodies (anti-ER, anti-PR, anti-HER2) and were transplanted into mice via the caudal vessel and resulted showed that each had a different luminescence
duration. Then, time-resolved images using in vivo multiple techniques were used to quantitatively analyze the expression of biomarkers in the tumors. In vivo multiple technology devices include CCD and charge coupling devices. Figure 6B shows the time-processing image, which decomposes MCF-7 and BT-474 tumors into three time-long paths, indicated by the red, green, and blue monochrome image collections. The expression of biomarkers was determined by integrating the intensity of each component and normalizing the total intensity to synthesize all tumor regions. Figures 6C,D are the results of in vitro protein imprinting: Figure 6C shows the result of immunohistochemical experiment of indirect in vitro therapy, While Figure 6D shows the result of determined biomarker expression modes of the two tumor hypotypes by using IVM.

At present, this field has become a vital resource for biomedical diagnosis and treatment, and many research groups are competing to continue pushing forward in expanding its study. Several high-level research papers and reviews have been reported recently, such as Guo et al. (2019), who used NIR-II fluorescence and photoacoustic imaging to study the precise identification of the vascular system and micro-tumors in 2019. It is well-known that the diagnosis of cerebrovascular structure with complete blood-brain barrier and microtumors is of great significance for the timely treatment of patients with nervous system diseases. The combined diagnosis and treatment of NIR-II fluorescence and photoacoustic imaging (PAI) is anticipated to provide improving performance, for instance, excellent spatial and temporal distinguishability, large penetration depth and great SNR, and accurate brain diagnosis. In this study, conjugated polymer nanoparticles (CP NPs) with biocompatibility and photostability were prepared, and bimodal brain imaging was achieved using its NIR-II window. The nanoparticles with a uniform size of 50 nm can be prepared by microfluidic device. The emission peak was at 1,156 nm and had strong absorption of 35.2 Lg−1cm−1 at 1,000 nm. NIR-II fluorescence imaging provides a solution to the depth and resolution issues previously experienced with the hemodynamic and cerebral vascular systems, providing an imaging depth of 600 microns and a spatial resolution of 23 microns. After an ultrasound-induced opening of the blood-brain barrier, NIR-II PAI could successfully perform non-invasive imaging of deep micro-brain tumors (2.4 mm below dense skull and scalp <2 mm) with a SNR of 7.2. This study showed that CP-NPs is a promising brain diagnostic contrast agent.

In 2019, Fan et al. (2019) published an overview of optical multiplex techniques for biological detection to improve biomedical diagnosis, arguing that traditional methods according to the detection of single disease markers may not be precise enough, since disease progression usually involves a variety of chemicals and biomolecules. Multi-target simultaneous analysis is of great importance in basic biomedical research and clinical application, promoting simultaneous multi-target analysis that requires the development of a high-throughput multi-target biological analysis technology. In order to improve the level of biomedical diagnosis, this paper reviewed the research progress of optical multiplexing analysis technology used in biomedical diagnosis over the recent years. The review primarily focused on the Fluorescence and Surface Enhanced Raman Scattering (SERS) technique which has unique optical characteristics as a main signal reader. The paper also focused on the review of the multiplexing strategy in biomedical field and the recent advances in biosensors from multi-analyte and multi-color cell tracking to multi-channel bioimaging in vivo. Finally, the paper provided forecasting of future challenges and opportunities of multi-bio-analysis.

In 2018, He et al. (2018) proposed that the structural design and synthesis of fluorophores emitted in NIR-II biological window are moving toward multi-mode imaging, as well as toward multimodal diagnosis and treatment. He et al. reviewed some progress of NIR-II fluorophores and molecular probes and considered the synthesis of NIR-II fluorescent group fit for multimode imaging will become a new way to obtain high-resolution images through the structural design. NIR-II fluorophores can convert NIR-II photons into heat required for photothermal therapy and can also be stimulated by NIR-II light to generate singlet oxygen for photodynamic therapy. The single probe has both diagnostic and therapeutic functions which can be used for precise treatment. He et al. details the latest development in structural design and synthesis of various NIR-II fluorophores and provides a discussion on the similarities and differences in known NIR-II imaging systems and the recent research on NIR-II imaging in biomedical applications.

### Tracking the Transplanted Stem Cells

Stem cell (SC) is one type of pluripotent cells with self-replication ability, which can differentiate into a variety of functional cells under certain conditions. SC therapy is showing the hope of curing major human diseases. In the process of SC therapy, how to observe the transplanted SCs non-invasively in real time has become a challenge in experimental and clinical research. NIR imaging based on exogenous markers has always been one of the best methods to trace stem cells in vivo.

In recent years, NIR fluorescent probes, with their strong biological penetration ability, low photon detection domain value, high detection sensitivity, relatively simple operation, and advantages of real-time imaging during surgery, have made the shift of emission from NIR-I window to NIR-II window to track transplanted stem cells. For instance, Chen et al. (2018b) has reviewed the research progress of tracking transplanted SCs with NIR fluorescence NPs. The latest development of NIR-II window fluorescence imaging technology was introduced. The emergence of new fluorescent functional NPs (QDs, RE-doped NPs, organic fluorescent NPs, etc.) makes stem cells easy to be labeled and tracked, greatly promoting the further development of stem cell therapy. NIR-II fluorescence emission is absorbed little by tissues, and the scattering and spontaneous fluorescence in NIR-II region are the least in tissues. Therefore, the signal-to-background ratio can be greatly improved, yielding to a large tissue penetration depths and a good spatial-temporal resolution. Moreover, NIR-II fluorescence imaging technology for accurate optical tracking can improve our comprehending of the fate and regeneration ability of transplanted SCs and provide immense potential development of SC-based regenerative medicine. Up to now, only a few research labs are exploring the feasibility of
FIGURE 7 | Ag₃S quantum dots of NIR-II emitting were used to track the transplanted MSCs (Chen et al., 2018b). (A) Representative TEM pictures and fluorescence emission spectra of Ag₃S QDs with diverse sizes. Adapted from Zhang et al. (2014) written by Qiangbin Wang etc. with permission. (B) Ag₃S QDs labeled MSCs were (Continued)
using NPs emitted in NIR-II to track SCs in vivo. Ag2S QDs-based NIR-II imaging is currently the only method that has succeeded in SC tracking research in vivo. In order to greater improve the performance of NPs emitted in NIR-II in SC-based regeneration research, more efforts should be made to conquer the defects (such as water solubility, physiological stability, biocompatibility, and metabolic capacity) of current NIR-II NP-based imaging technology. In addition, new strategies are needed to take full advantage of NIR-II excitation and emission. The comparison diagram of transplanted mesenchymal stem cells (MSCs) tracked by Ag2S quantum dots emitted NIR-II fluorescence is shown in Figure 7.

Figure 7A shows typical TEM pictures and fluorescence emission spectrum of Ag2S quantum dots with different particle sizes. Figure 7B shows the real-time NIR-II fluorescence imaging of Ag2S QDs labeled MSCs after intravenous injection into mice exposed for 100 ms. When excited at 808 nm, NIR-II fluorescence images were collected with InGaAs/SWIR camera. The fluorescence intensity of NIR-II in liver and lung of mice at diverse time points was measured, and then quantitative analysis was performed. In Figure 7C, Ag2S quantum dots were labeled on MSCs and then injected into mice with acute liver failure for tracking study. MSCs were combined with or without heparin and then transplanted intravenously into mice and imaged. Figures 7D,E, respectively show in vivo imaging and fluorescence quantitative homing behavior of MSCs to the wound in a mouse model with skin trauma. MSCs were transplanted intravenously into model mice. The left and right wounds were treated with SDF1-α-loaded collagen scaffolds and collagen scaffolds, respectively.

Other related researches on the tracking of transplanted CS with NIR-II quantum dots include an article published in 2019 by Chen’s team about the fate of intraarticular injection of MSCs in vivo for the useful treatment of supraspinatus tendon tears (Yang et al., 2019). This paper reports that MSCs have a strong therapeutic potential in the treatment of supraspinatus tendon tears. However, by reason of the finite evidence of dynamic visualization of cell behavior in vivo, MSC therapy has not been fully utilized and may even be underestimated. Here, PbS QD labeled MSCs can treat supraspinatus tendon tears in mice. PbS QD is a biocompatible NIR-II fluorescence imaging probe, which can provide a cell migration map and information about the biological distribution and clearance process of MSCs injected into the joints with three densities. Intra-articular injection can avoid MSCs being wrapped by filtered organs and reduces organ toxicity induced by quantum dots. It is worth noting that MSCs have similar migration directions, but the migration efficiency of the medium density group is higher. In the repair stage, MSCs stay around the footprint for the longest time, with the highest cell retention rate. In addition, quantitative kinetics studies showed that labeled MSCs were cleared by feces and urine. Histomorphological analysis showed that therapeutic effect of the medium density group was the best, and the labeled MSCs showed no damage or inflammatory response to the main organs, suggesting that too much or too little MSCs administration might reduce its efficacy. This imaging method provides spatio-temporal evidence for the response of MSCs therapy in vivo, which is helpful for the optimization of MSCs therapy.

Moreover, in 2018, Huang et al. published a paper on NIR-II fluorescence and bioluminescence multiple imaging for in vivo observation of the location, survival and differentiation of transplanted stem cells (Huang et al., 2019). An NIR-II fluorescence/bioluminescence composite imaging method is successfully developed covering 400–1,700 nm visible light and NIR-II window for in vivo monitoring of location, survival, and osteogenic differentiation of human bone marrow mesenchymal stem cells (hMSCs) in a mouse model of skull defect. The long-term biological distribution of transplanted hMSCs was observed by using Ag2S quantum dot with NIR-II window. Bioluminescent imaging (BLI) based on endogenous red firefly luciferase (rfluc) and gaussia luciferase (gluc) driven by collagen type 1 promoter were used to report the survival and osteogenic differentiation status of transplanted hMSCs. At the same time, by combining the three imaging channels, they can not only directly observe the various dynamic biological behaviors of the transplanted hMSCs, but also further observe the promoting effects of immunosuppression and bone morphogenetic protein 2 on the survival and osteogenic differentiation of the transplanted hMSCs. This new multifunctional imaging technique can widely extend the analysis of the fate and therapeutic capacity of transplanted stem cells and help to promote stem cell-based regenerative therapy and its transformation in the clinical applications.

CONCLUSION

In summary, compared with traditional NIR-I imaging technology and other medical imaging methods, NIR-II bioimaging technology not only has a deeper imaging depth, but also can better avoid background interference such as spontaneous fluorescence and photon scattering of tissue. So far, a variety of NIR-II dyes, such as inorganic nanomaterials (single-walled carbon nanotubes, semiconductor quantum dots, rare earth nanomaterials), conjugated polymers and organic small-molecule materials, have been successfully synthesized and prepared. Because of its special properties, NIR-II dyes can
TABLE 1 | NIR-II contrast agent and its biomedical application.

| Fluorophore                  | Fluorescence property | Biomedical applications                                                                 | References                  |
|-----------------------------|-----------------------|-----------------------------------------------------------------------------------------|-----------------------------|
| SWCNPs                     | 808/950–1,400 nm (excitation/emission) | High-resolution imaging of blood vessels                                               | Welsher et al., 2009        |
| M13-SWCNTs                 | 808/950–1,100 nm (excitation/emission) | Targeted fluorescence imaging of tumor tissues in mice                                 | Ghosh et al., 2014          |
| IRDye-800- SWCNTs          | 808/1,300–1,400 nm (excitation/emission) | Imaging of the distribution of veins in the head of mice and the fine structure of the brain capillaries | Hong et al., 2014           |
| A synthetic organic molecule (CH1055) | 750/1,055 nm (excitation/emission) | Tumor imaging and precise image-guided tumor removal surgery                           | Antaris et al., 2016        |
| o-SWCNT-PEG                | 980/1,300 nm (excitation/emission) | Used in angiographic imaging                                                          | Takeuchi et al., 2019       |
| Ag2S QD                    | 800/200 nm (excitation/emission) | Used in NIR-II cell imaging and non-specific tumor detection                             | Hong et al., 2012b; Zhang et al., 2012, 2013 |
| Au/Cu alloy nanodots or Au/Co alloy nanodots | 808/950–1,100 nm (excitation/emission) | It is a non-toxic and multifunctional nanodots for use in biological imaging, especially in multi-mode imaging | Andolina et al., 2013; Marbella et al., 2014 |
| NaLuF4:Yb/Nd@NaYF4 (L1= Er, Ho, Trm, Pr) nanocrystals | 800/1–1,500 nm (excitation/emission) | Widely applied in vivo biomaging research                                              | Naczynski et al., 2013      |
| SrF2:Nd3+/NP                | 808/1,100 nm (excitation/emission) | In vivo NIR-II optical imaging                                                        | Villa et al., 2015          |
| NaYF4:Yb3+/x@NaYF4@NaYF4: Nd3+/x (x = null, Er, Ho, Trm or Pr) | 700–860/1,000–1,600 nm (excitation/emission) | NIR-II in vivo imaging of deep tissue                                                | Shao et al., 2016           |
| FD-1080                    | 1,046/1,080 nm (excitation/emission) | Non-intrusive high distinguishability depth tissue                                       | Li et al., 2018             |
| Free ICG                   | 810–830/1,100 nm (excitation/emission) | Utility in NIR-II navigated cytoreductive surgery of cerebral cancers and accelerate the clinical translation of NIR-II imaging technology | Starosolski et al., 2017; Zeh et al., 2017; Cho et al., 2019a,b; Wang et al., 2019b |
| liposomal-ICG (Lip-ICG)     | 782/950–1500 nm (excitation/emission) | Long-term imaging of hind limb and intracranial vessels in vivo                         | Bhavane et al., 2018        |
| ICG conjugated bevacizumab (Bev-ICG) | 808 nm/900 nm (excitation/emission) | Immensely facilitate the application of NIR-II clinical biomaging                     | Suo et al., 2019            |
| Core:DCNPs(NaGdF4:5%Nd@NaGdF4) | 808/1,060 nm (excitation/emission) | Tumor imaging and image-guided surgery                                                   | Wang et al., 2018           |
| In vivo assembly nanoprobe: DCNPs-L1@F3H4L1 and L2 are conjugated DNA: F3H4 is a follicle stimulating hormone peptide | 808/1,060 nm (excitation/emission) | Tumor imaging and image-guided surgery                                                   | Wang et al., 2018           |
| Ag2Te quantum dots (QDs)   | 808/1,300 nm (excitation/emission) | Tumor imaging and image-guided surgery                                                   | Zhang et al., 2019          |
| lanthanide complex (Nd-DOTA) | 808/1,060 and 1,330 nm (excitation/emission) | Accurately mark the profile of micro-tumors in surgery and thoroughly removed under the guidance of NIR-II biological imaging | Yang et al., 2018           |
| semiconductor polymer nanoparticle of diketopyrrolopyrrole (PDP1032) | 809/1,032 nm (excitation/emission) | Widely used in clinical imaging and surgical treatment of malignant tumors             | Shou et al., 2018           |
| NaErF4:2%Ho@NaYF4:UCNPs the sensor is based on the ratio meter fluorescence | 1,530/1,180 nm (excitation/emission) | Used to quantify the degree of actual inflammation in clinical tests                   | Liu et al., 2018            |
| DCNPs(NaGdF4:5%Nd@NaGdF4)@GSH | 808/1,060 nm (excitation/emission) | Accurate imaging of inflammation in vivo                                               | Zhao et al., 2019           |
| pentamethine cyanine NIR-II molecular fluorophore | 1,015/1,065 nm (excitation/emission) | High-quality imaging and pH sensing in vivo                                             | Wang et al., 2019a           |
| Peroxymtrate NIR-II molecular fluorophore IRBTP-O | 575 and 905/1,000 nm (excitation/emission) | Drug-induced hepatotoxicity detection                                                  | Li et al., 2019c             |
| Polyaacryl acid (PAA) modified NaLuF4:Gd@NaLuF4:Gd@NaLuF4:Gd | 808/1,058 and 1,328 nm (excitation/emission) | For Fluorescence imaging-guided detection of small tumors, angiogenesis-related diseases, and angiogenesis diagnosis | Li et al., 2019a             |
| NaLuF4:Gd/Yb/Er NRs@POA | 980/1,525 nm (excitation/emission) | Visualization of tumor-related vascular malformations and imaging-guided photothermal therapy | Li et al., 2019b             |
| A bright organic NIR-II fluorescent nanocluster (p-FE) | 980/1,100 nm (excitation/emission) | As a vascular imaging reagent for non-invasive in vivo blood flow tracking in the rat cerebrovascular system | Wan et al., 2018             |

(Continued)
be used not only as biomedical contrast agent, but also in the fields of photothermal and photodynamic therapy, drug delivery, surgical guidance and tracking the transplanted stem cells. In this article, the reported NIR-II fluorescent probes and their biomedical applications are summarized in Table 1.

With the development of molecular design theory and nanomaterials, more and more NIR-II imaging systems will be designed, developed and promoted into clinical trials. It is worth noting that at present, most of NIR-II fluorescent molecules have poor water solubility and physiological stability, and low fluorescence quantum yield. For polymers and inorganic nano systems, problems, such as slow metabolism, high toxicity, and lack of specific tissue targeting, still exist. New methods and materials will be helpful to promote the application of NIR-II imaging technology in the field of biology. In general, the following points should be paid attention to in the development and design of NIR-II dyes. Firstly, in order to improve the water solubility and physiological stability of organic NIR-II fluorescent dyes, reduce the band gap when choosing the appropriate space configuration, and consider the influence of the group on the water solubility of dyes when modifying the group. Secondly, in order to improve the biocompatibility and metabolic capacity of the probe, the molecular structure should be designed reasonably. In particular, when the probe is a polymer molecule dye and nanoparticles, the design and synthesis of the degradation of NIR-II dye probe is the best. Thirdly, dye is endowed with specific functions, such as combining dye with targeted molecules, improving the aggregation ability of dye in specific parts of the organism, and promoting the development of disease targeting and early diagnosis in clinical practice.

**AUTHOR CONTRIBUTIONS**

JC wrote the draft and corresponded to submit, revision, and coordination. BZ, KZ, SH, LM, JS, and HY participated in manuscript revision. JS and HY oversaw the design, quality assurance, and finalization of the manuscript.

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