NIR-I Dye-Based Probe: A New Window for Bimodal Tumor Theranostics

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Near-infrared (NIR, 650–1700 nm) bioimaging has emerged as a powerful strategy in tumor diagnosis. In particular, NIR-I fluorescence imaging (650–950 nm) has drawn more attention, benefitting from the high quantum yield and good biocompatibility. Since their biomedical applications are slightly limited by their relatively low penetration depth, NIR-I fluorescence imaging probes have been under extensive development in recent years. This review summarizes the particular application of the NIR-I fluorescent dye-contained bimodal probes, with emphasis on related nanoprobes. These probes have enabled us to overcome the drawbacks of individual imaging modalities as well as achieve synergistic imaging. Meanwhile, the application of these NIR-I fluorescence-based bimodal probes for cancer theranostics is highlighted.

Keywords: NIR-I fluorescence imaging, bimodal imaging, nanoparticles, cancer, theranostics

1 INTRODUCTION

Cancer is one of the major public health problems across the world (Torre et al., 2015; Feng et al., 2019). Advanced imaging techniques undoubtedly play an indispensable role in the development of diagnostic and therapeutic approaches for cancer. With the advantages of high sensitivity, high selectivity, noninvasiveness, and real-time visualization, fluorescence imaging (FI) has been widely used in the clinic to diagnose diseases and precisely guide resection in surgery (Jewell et al., 2014; Pramod Kumar et al., 2020). The working mechanism of realizing imaging functionality underscores the need for proper reporter groups as tracers. The reporter group includes a variety of fluorescent dyes, fluorescent proteins, upconversion nano-particles, quantum dots, and so on (Adhikari et al., 2016; Laviv et al., 2016; Kostiv et al., 2017; Saljoghi et al., 2020). They can absorb the energy of photons to reach an excited state, and then return to the ground state accompanied by the release of energy as fluorescence. Among them, fluorescent dyes are considered as ideal reporters due to their superiority of simple components, flexible structure, and good biocompatibility.

However, conventional visible fluorescence imaging always suffers from low tissue penetration depth in in vivo optical imaging, which is attributed to the fact that light with a short wavelength can be more easily scattered and absorbed by endogenous biological tissues including skin, fat, and blood (Smith et al., 2009; Sordillo et al., 2014). The emergence of near-infrared (NIR) fluorescence imaging has brought a new chance with the merits of deeper tissue penetration and minor auto-fluorescence interference, which can afford distinct signals with increased spatial resolution and imaging sensitivity to track target tissues. Generally, it can be further divided into first near-infrared (NIR-I, 650–950 nm) and second near-infrared (NIR-II, 1,000–1700 nm) fluorescence imaging.
This review summarizes the advances of NIR-I fluorescent dye-related bimodal imaging of tumors in the recent 5 years extensively. We mainly focus on the rational design strategies and the representative paradigms of versatile probes, while the challenges and outlook have also been discussed. We hope that this review will raise extensive interest and offer suggestions for the development of biomedical multimodality imaging in the diagnosis of cancer.

More than this, the development of multimodality imaging is further promoted via the blossoming of nanotechnology. On the one hand, characteristics of nanoparticles (NPs), including large surface-to-volume ratio, passive accumulation at tumor endothelial cells, high labeling capacity, and prolonged blood circulation, are advantageous in this field. On the other hand, nanomaterials make it easier to combine different functional blocks. Remarkably, numerous multimodal probes can be integrated within one single nanoparticle, which greatly improves the delivery ability for cancer diagnosis.

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### 2 BIMODAL IMAGING OF TUMOR BASED ON NEAR-INFRARED-I FLUORESCENCE

Tumor-specific imaging is achieved by distinguishing the distinction between malignant and healthy tissues. Tumor tissues comprise not only tumor cells but also the...
microenvironment where they exist. With this in mind, the design strategies of imaging probes can be developed based on the discrepancies between tumor and normal tissues, such as lower pH, hypoxia, and overexpression of enzymes and integrins (Imamura et al., 2018; Shim et al., 2019; Tansi et al., 2020).

Various multimodality imaging techniques, which refer to the combination of NIR-I fluorescence imaging with other imaging modalities, have been reported as advanced diagnostic approaches for cancer. Hopefully, designing dual-modal probes for FI/NMI, FI/MRI, FI/CT imaging, and FI/PAI systems will compensate for the weakness of individual modality and enable synergistic imaging.

2.1 Fluorescence Imaging/Nuclear Medicine Imaging

NMI has been commonly used in cancer diagnosis owing to its infinite penetration depth and high temporal resolution. There are two main modalities: single-photon emission computed tomography (SPECT) and positron emission tomography (PET). Since SPECT requires a collimator for imaging which is needless for PET, its detection sensitivity is usually lower than PET. Metallic or nonmetallic radionuclides are commonly used as tracers to generate images in NMI. The working mechanism is the detection of gamma (γ) rays directly or indirectly from the decay of these radionuclides. Tc-99m, Ga-67, I-123, I-131 and I-131 are often utilized as radioactive atoms in SPECT imaging (Beer et al., 1990; Al-Suqri and Al-Bulushi, 2015), while Ga-68, Cu-64, F-18, C-11, and I-124 are used in PET imaging (Nakajima et al., 2017).

Hence, the commercialized NMI probes are usually radiolabeled molecules. For example, 18F-fluorodeoxyglucose (18F-FDG) has been widely applied as a PET radiotracer to evaluate neoplastic diseases.

Bimodal imaging techniques, such as PET-CT and PET-MRI, have already been well established for clinical use. These clinical NMI-based bimodal imaging technologies undoubtedly hold the potential for improved diagnostic evaluation by merging functional information with anatomical information or soft tissue details. However, their solutions to the high radiation exposure and related health risks remain limited. Meanwhile, difficulty in the diagnosis of tiny lesions in the early stages of cancer hinders their further biological application. Since FI has high sensitivity but with significantly lower toxicity, the bimodal FI/NMI probes enable a lower dose to offer a better image in the early diagnosis of tumors compared to individual NMI or other NMI-based bimodal probes (Wang X. et al., 2018). The combination of FI and NMI can overcome the obstacles of single modality imaging, allowing imaging with little radiation burden and unlimited penetration depth. For the construction of the bimodal probes in FI/NMI, however, the direct use of radiolabeled fluorescent dyes lack clear targeting sites (Kanagasundaram et al., 2021; Peng et al., 2021). In this context, modified molecular probes and passively accumulated nanoparticles have been attributed to implement specific tumor uptake.

2.1.1 Fluorescence Imaging/Single-Photon Emission Computed Tomography Imaging

SPECT is used more frequently than PET in the clinic due to wider availability of its scanners and radionuclides. The core of any radiotracer in SPECT imaging is its gamma-emitting radionuclide. 99mTc and 111In have been the most commonly used radionuclides in FI/SPECT dual-modal imaging. 99mTc has been used for medical imaging in 80% of cases around the world, and its decay product has little effect on the image quality (Lee and Grp, 2019). Also, benefiting from its physical properties, 99mTc (Eγ = 140.5 keV; t1/2 = 6.02 h) has minimal harmful radiation to the patients. Nevertheless, given the relatively short half-life of 59.6Tc, 111In (t1/2 = 2.8 d) has gained much attention recently.

Owing to the high affinity and specificity of antibodies, the imaging of labeled antibodies or engineered antibody fragments, which refers to immuno-imaging, has been widely applied in FI/SPECT bimodal probes. Hekman et al. had presented an FI/SPECT imaging probe, 111In-DTPA-labetuzumab-IRDye800CW, to detect carcinoembryonic antigen (CEA)—expressing pulmonary micrometastases. The near-infrared fluorescence (NIRF) dye IRDye800CW was selected, and the humanized anti-CEA monoclonal antibody, labetuzumab, was used to ensure adequate targeting of a tumor-associated antigen. Based on their experimental data, a feasibility study of this probe in patients with peritoneal carcinomatosis of colorectal origin could be considered (Hekman et al., 2017b). Besides, they replaced labetuzumab with farletuzumab and developed a new probe, 111In-DTPA-farletuzumab-IRDye800CW, for the intraoperative detection of ovarian cancer lesions (Hekman et al., 2017a). Later, they had also utilized immuno-imaging to design another probe 111In-DOTA-girentuximab-IRDye800CW. Successful visualization of tumors during surgery was performed in patients with clear cell renal cell carcinoma (ccRCC). Most notably, it was the first clinical application of tumor-targeted dual-modality imaging guided by monoclonal antibody (Figure 1) (Hekman et al., 2018). Meanwhile, boron-dipyrromethene (BODIPY) with high photoluminescence quantum yields and stability in the physiological environment was utilized to obtain 111In-Wazabay9-Trastu. As BODIPY was usually hydrophobic and water-insoluble, Privat et al. had succeeded in improving its solubility in aqueous media by substituting fluorine atoms on the boron by ammonium groups. Trastuzumab was bioconjugated to the probe for its high affinity toward the HER-2 receptors. 111In-Wazabay9-Trastu had been proved as a new tool for FI/SPECT imaging-guided surgery in nude mice bearing HER2-overexpressing HCC1954 human breast cancer xenografts (Figure 2) (Privat et al., 2021). Furthermore, the non-covalent interaction between biotin and avidin in the biotin–avidin system (BAS) is the strongest in vivo, which is much higher than the affinity between the antigen and antibody. Based on this, Dong et al. designed a novel dual-modality probe 99mTc-HYNIC-lys(Cy5.5)-PEG4-avidin by using Cy5.5 as an NIR fluorophore to achieve modest signal amplification, further promoting the imaging of human colon adenocarcinoma xenografts. The in vitro and in vivo results reflected that FI exhibited exquisite congruence to SPECT imaging (Dong et al., 2016). Additionally, αβ3-integrin is an ideal tumor target spot as...
attributed to the selective attachment of RGD-containing peptides toward it. In this context, Yin et al. reported a 125I-radiolabeled probe cRGD-QC by which metastatic lymph nodes could be detected specifically and accurately. The probe was constructed via the attachment of an NIR dye (Cy5) and a quencher (QSY21) to a 125I-labeled cRGD. Only upon the proteolytic cleavage by activated matrix metalloproteinase-2 (MMP-2), the fluorescence of cRGD-QC was recovered. Under this circumstance, cRGD-QC showed high tumor-to-background ratios with low impact on normal tissues (Yin et al., 2019). Rizvi et al. had developed a Cy5.5-conjugated self-assembled peptide nanoprobe Cy5.5@SAPD-99mTc based on a novel head-to-tail cyclic RGD-KLAK heptapeptide sequence. The effectiveness and efficacy of this probe for the diagnosis of glioblastoma multiforme had been proven in the tumor-bearing female Balb/c mice models guided by FI/SPECT imaging (Rizvi et al., 2021).

Other nanoparticles applied in FI/SPECT imaging were mainly designed by using different nanoplatforms. By combining a fluorescent dye (Dylight 755) with 99mTc in multiple-armed DNA tetrahedral nanostructures (TDNs) to synthesize nanoprobe FA-Dy-99mTc-TDN, the group of Jiang successfully realized noninvasive FI/SPECT imaging in tumor-bearing mice. In comparison with those of double-stranded DNA, the TDNs showed remarkably enhanced stability in vitro (Jiang et al., 2016). Furthermore, based on tumor cell–derived exosomes (TEx), 99mTc-TEx-Cy7 was developed through the combination of Cy7 and 99mTc. It was the first exosome-based nanoprobe for multimodal SPECT and NIRF imaging (Jing et al., 2021a). Remarkably, 99mTc-based FI/SPECT imaging had already been applied in the clinic. Manca et al. had proved the excellent performance of ICG-99mTc nanotop in both preoperative lymphatic mapping and intraoperative sentinel lymph node detection (Manca et al., 2021). Of note, Shi et al. prepared cetuximab/IR-780/micelles as immuno-imaging nanoparticles which further guaranteed the targeting properties for EGFR overexpression in colorectal cancers. A lipophilic dye, IR-780 iodide, was applied here to offer fluorescence imaging as well as photothermal therapy (PTT). In particular, it was a dual-isotope dynamic SPECT imaging system with the loaded dye and the micelles, respectively, radiolabeled with 131I and 111In (Shih et al., 2017).

2.1.2 Fluorescence Imaging/Positron Emission Tomography Imaging

Compared to SPECT cameras, the scanners and radiotracers of PET can provide higher sensitivity. Unlike SPECT, positrons are emitted during the decay of the radionuclides, and then energy is released in the form of two 511-keV photons. These annihilation photons can be detected by PET cameras and therefore locate the approximate position of the PET radionuclide. However, only 18F, 124I, 64Cu, 68Ga, and 89Zr have been applied in FI/PET imaging (An et al., 2017; Ghosh et al., 2017; Wang Y. et al., 2020). Immuno-imaging probes at the molecular level have also been widely developed in this field. Over a third of the synthesized FI/PET bimodal probes involved antibodies (Luo H. et al., 2017). Most of these probes were labeled with radionuclides of long decay half-lives, including 89Zr (t1/2 = 3.3 d) and 124I (t1/2 = 4.2 d) which matched the pharmacokinetics of intact antibodies (Hernandez et al., 2016; Tsai et al., 2018; Zettlitz et al., 2018). Nevertheless, the use of long-lived radionuclides might result in great radiation exposure to the patient and also requires plenty of lag time before imaging. Hence, researchers focused on the design of 64Cu-radiolabeled probes due to the relatively long half-life of 64Cu (t1/2 = 12.7 h) among the short-lived radionuclides. For example, Adumeau et al. had presented a pre-targeted strategy for the bimodal FI/PET imaging of colorectal carcinoma. The NIR fluorophore-labeled antibody huA33-Dye800-TCO and the radioisotope 64Cu-Tz-SarAr were injected separately. Then, a click ligation of these two components occurred in vivo based on an extraordinarily rapid and biorthogonal inverse electron demand Diels–Alder reaction. As a result, little fluorescence in healthy organs and high tumor-to-normal tissues radiant efficiency ratio showed up (Adumeau et al., 2016). However, 68Ga had not been applied in monoclonal antibody-based FI/
FIGURE 2 | (A) Structure of $^{111}$In-Wazaby-Trastu. (B) Near-infrared fluorescence images of $^{111}$In-Wazaby-Trastu on mice bearing subcutaneous tumor of HCC1954 cells HER-2+. (C) Representative SPECT/CT images. (D) Fluorescence-guided resection of tumor tissues. Reproduced with permission from Privat et al. (2021). Copyright 2021, American Chemical Society.
PET imaging owing to its extremely short half-life ($^{68}$Ga, $t_{1/2} = 67.7$ min). Still, it had been widely used in the dual-modal probes for FI/PET imaging due to its relatively low cost and easy preparation from a $^6$Ge/$^6$Ga generator system (Summer et al., 2017; Zhang et al., 2018). For instance, the group of Li reported a novel probe $^{68}$Ga-IRDye800CW-BBN. It was the first-in-human

**FIGURE 3** | (A) Chemical structure of IRDye800CW-BBN tracer. (B) Scheme of $^{68}$Ga-IRDye800-BBN for the GBM patients. (C) $^{68}$Ga-IRDye800-BBN FI/PET dual-modality imaging guided GBM resection for patient 3. Reproduced with permission from Li et al. (2018). Copyright 2018, Ivyspring International Publisher.
study of dual-modal FI/PET imaging to guide surgery in glioblastoma. Among the 14 patients enrolled, preoperative positive PET uptake had an excellent correlation with intraoperative NIRF signal. Patient 3 was taken as an example where even in the deep tumor cavity, the residual tumor had obviously differentiated from the adjacent normal brain tissue by dual-modality imaging (Figure 3) (Li et al., 2018).

Nanoprobes were used more frequently in FI/PET imaging and mostly radiolabeled with $^{64}$Cu (Wang X. et al., 2018). Luo et al. developed a doxorubicin-loaded porphyrin–phospholipid (PoP) liposomes platform with intrinsic fluorescence capacity. And then $^{64}$Cu was radiolabeled to the stable bilayer of preformed Dox-loaded PoP liposomes to synthesize Dox-CuPoP. According to PET and fluorescent imaging, passive nanoprobe accumulation was visualized in orthotopic mammary tumor-bearing mice. In addition, the growth of 4T1 orthotopic tumors was strongly inhibited in a single chemophototherapy treatment with Dox-CuPoP under the illumination of 665 nm light (200 J/cm$^2$) (Figure 4) (Luo D. et al., 2017). Du et al. had designed a breast tumor–targeted nanoprobe, PD-1-Liposome-DOX-$^{64}$Cu/IRDye800CW, based on liposomes as well (Du et al., 2017). Besides, gold nanoparticles and polysaccharide dextran were also served as nanocarriers for $^{64}$Cu in FI/PET imaging (Pretze et al., 2019; Deng et al., 2020). Significantly, to further reduce potential radiation damage, Jing et al. selected extracellular vesicles derived from adipose-derived stem cells (ADSCs) as nanoplatforms, and Cy7 was afforded to obtain Cy7-EV-N$_3$. $^{68}$Ga was connected to the platform based on the click reaction between Cy7-EV-N$_3$ and $^{68}$Ga-L-NETA-DBCO in vivo. Comparing different pre-targeting and imaging time points, maximum tumor uptake was obtained at 20 h after the injection of Cy7-EV-N$_3$ and at 2 h after the injection of $^{68}$Ga-L-NETA-DBCO. The radiation exposure was obviously reduced based on this strategy (Jing et al., 2021b).

Meanwhile, there remain challenges that nanoprobe often suffer from poor reproducibility and low tumor penetrability due to their much larger sizes compared to organic small-molecule probes (Yan et al., 2019). The group of Wang considered designing self-assembled nanoparticles to weaken the effect of these defects. The polyphenol and poloxamer self-assembled supramolecular nanoparticles (PPPNs) were fabricated by multivalent hydrogen bonding between tannic acid and Pluronic F-127 together with hydrophobic interactions of poly(propylene oxide) chains. IR-780 was encapsulated by PPNPs through hydrophobic interactions to perform NIRF imaging after releasement. Then PPNPs-IR-780 was labeled with $^{89}$Zr to form PPNPs-IR-780-$^{89}$Zr without additional chelators, which was attributed to the excess phenolic hydroxyl groups of tannic acid. Of note, the in vivo NIR fluorescent images showed surprisingly higher fluorescence intensity in tumors than in other tissues, while the vast majority of PPNPs-IR-780-$^{89}$Zr was accumulated in the liver and spleen according to PET imaging. As the leakage of IR-780 from the probe could be observed in serum, it might explain why FI and PET imaging showed different biodistribution of the probe (Figure 5) (Wang X. et al., 2018). Additionally, Hu et al. reported an enzyme-activatable probe P-CyFF-$^{68}$Ga and its cold probe (P-CyFF-Ga) which resulted in co-assembling into fluorescent and radioactive nanoparticles (NP-$^{68}$Ga) in response to alkaline phosphatase (ALP). It was demonstrated that the ALP-triggered
in situ formed NP-\textsuperscript{68}Ga was capable of providing FI/MRI of the ALP-positive tumors with high sensitivity and deep penetration depth (Figure 6) (Hu et al., 2021).

### 2.2 Fluorescence Imaging/Magnetic Resonance Imaging

By recording the spatial distribution of water protons, MRI is capable of generating a 3D anatomical image of tissues. It has been broadly applied in the aspect of soft tissue lesions and deep tissue pathological details imaging (Domey et al., 2016; Tsai et al., 2018). Further compared with NMI, the images can be formed without injecting radionuclide which provides higher safety (Kim et al., 2016). The signal intensity of MRI is affected by the relaxation rate of water protons which includes the longitudinal relaxation rate and transverse relaxation rate. Contrast agents (Cas) used in MRI are to shorten the longitudinal relaxation time ($T_1$) or the transverse relaxation time ($T_2$) in order to image specific regions of interest. Hence, they can be divided into $T_1$-weighted Cas [e.g., paramagnetic gadolinium (Gd) or manganese (Mn)] and $T_2$-weighted Cas [e.g., superparamagnetic iron oxide (SPIO)]. $T_1$-weighted Cas shows positive contrast enhancement with brighter images by shortening the $T_1$ of protons, and $T_2$-weighted Cas shows negative contrast enhancement with darker images by shortening the $T_2$ of protons. However, MRI has some constraints such as poor resolution and lack of sensitivity because of the overlap of $T_1$ or $T_2$ between healthy tissues and lesions. As an ideal complement to MRI, FI can provide sensitive imaging to distinguish pathological organs with high temporal resolution. The development of FI/MRI probes allows the imaging of the fine distribution of the probes in tissues and provides various information, including anatomical, physiological, and even molecular information (Ortgies et al., 2016; Getachew et al., 2022).

Beyond the probe Gd-Cy7-PTP/GRG designed by the group of Wang for pancreatic ductal adenocarcinoma (PDAC) imaging (Wang Q. et al., 2018), other FI/MRI probes were platform-free or platform-based nanoprobes (Supplementary Table S1). Platforms, such as mesoporous silica, serum albumin, micelles, and liposomes, had been widely applied in this field which were mainly involved in $T_1$-weighted Cas-contained probes (Cressey et al., 2021).

One widely used strategy to develop dual-modal imaging agents for FI/MRI is based on fluorescent labeling magnetic nanoparticles (MNPs). MNPs not only possess controllable dimensions and easily modified surfaces but can also provide excellent magnetic responsiveness which is manipulated by external magnetic fields. In particular, iron oxide nanoparticles have drawn much attention due to their low toxicity and feasibility to produce iron oxide cores (Lee et al., 2018; Li et al., 2019a; Reichel et al., 2020). Many hydrophilic substances [such as polyethylene glycol (PEG), poly(acrylic acid) (PAA), galactosyl conjugated P123 (Gal-P123), and red blood cell membrane] were used to modify iron oxide NPs in order to enhance biocompatibility and stability (Liang et al., 2017;
Polymers, such as polyamidoamine (PAMAM) and poly(lactic-co-glycolic) acid (PLGA), were ideal candidates to label NIR fluorophores (Duan et al., 2019; Wang et al., 2019d). Moreover, there were some other substances, such as calcium carbonate, dextran, chitosan, poly(succinimide) (PSI), and liposome, that could combine the ability of improving biocompatibility and encapsulating fluorescent molecules (Yang et al., 2018; Xie et al., 2020). For example, Zhang et al. had designed a novel theranostic nanocomposite MUCNPs@BPNs-Ce6. To prepare MUCNPs, the red upconversion luminescent shell (MnO2-doped NaYF4:Yb/Er/Nd) was performed on the oleic acid capped Fe3O4 NPs by hydrothermal synthesis. The upconversion part was used to match the wavelengths of black phosphorus (BP) for making optimal PTT and photodynamic therapy (PDT) work simultaneously. Afterward, MUCNPs were modified with PAA to acquire good hydrophilicity. MUCNPs@BPNs-Ce6 was finally obtained through the coupling between the carboxyl groups of the chlorin e6 (Ce6) molecules and the exposed amino groups of MUCNPs. The results demonstrated that the probe could be exploited as a good theranostic agent for simultaneous FI/MRI, highly efficient PDT and PTT of tumor (Figure 7) (Zhang et al., 2020). Besides, dozens of SPIOs were encapsulated in one nanocapsule (NC) via an adapted water-in-oil-in-water (W/O/W) emulsion by poly(ε-caprolactone-co-lactide)-β-poly(ethylene glycol)-b-poly(ε-caprolactone-co-lactide) (PCLA-PEG-PCLA) polymers through a solvent evaporation process, reported by Liao et al. A cyanine dye IR-820 and paclitaxel (PTX) were introduced into the NCs to

FIGURE 6 | (A) Schematic illustration showing an ALP-enabled in situ co-assembly strategy for FI/PET bimodal imaging. (B) Fluorescence images of HeLa tumor–bearing mice with different treatments. (C) PET imaging of HeLa tumor in mice at 10, 30, 60, 90, and 120 min postinjection of indicated substance. Reproduced with permission from Hu et al. (2021). Copyright 2021, American Chemical Society.
develop NC-SPIOs-IR-820-PTX. It could provide FI/MRI, as well as chemotherapy and PDT, for 4T1 tumors of mice (Liao et al., 2017). The group of Wang also applied the W/O/W emulsion method to co-load doxorubicin (DOX) and indocyanine green (ICG) into Fe/FeO-PPP heterostructures to form DOX-ICG@Fe/FeO-PPP nanocapsules. The designed nanoprobe was capable of dual-imaging of the KB tumor-bearing nude mice (Wang Z. et al., 2019).

The method of constructing fluorescent-labelling MNPs in T2-weighted bimodal FI/MRI has also been used as a reference in the design of T1-weighted dual-modal nanoprobes. However, for the synthesis of these probes, the formation of T1-weighted nanoprobes based on different platforms was required at first. Sahu et al. selected serum albumin as a surface stabilizer to produce Prussian blue (PB) nanoparticles. And then, ICG was encapsulated into the PB-BSA nanoparticles to form PB-BSA-ICG. Even if PB had the ability to shorten T1 and T2, they only focused on the T1-weighted MRI property of the probe. PB-BSA-ICG could provide bimodal FI/MRI of the tumor as well as a combined PTT-PDT (Sahu et al., 2016). The other T1-weighted dual-modal nanoprobes were based on Gd or Mn which depended on platforms including serum albumin, regenerated silk fibroin, micelle, monolayered-double-hydroxide nanosheets, carbon cage, and mesoporous silica (Picchio et al., 2021). Among them, taking the most commonly used platform, serum albumin, as an example, the group of Hao integrated ICG and PTX into the albumin corona of Gd2O3@HSA nanoparticles to prepare PIGH NPs. The experimental results of the fluorescence and MR bimodal imaging showed a high 4T1 tumor accumulation. The potential of chemotherapy and PTT of PIGH NPs was proven as well (Hao et al., 2019). Besides, it is worth noting that some nanoprobes were synthesized by utilizing the opposite form of this method. They
were constructed through CAs-labeling fluorescent NPs (Park et al., 2017; Ma et al., 2021; Yang et al., 2022).

Another common strategy is to integrate NIR fluorophore and CA based on the nanoscale matrix, including glycan, protein or polyllysine, DNA bipyramid nanostructure, virus-like particles, liposomes, silica-based NPs, gold nanocarrier, and PLGA NPs (Key et al., 2016; Henderson et al., 2018; Sasikala et al., 2018). Nonetheless, the effect of aggregation fluorescence quenching (ACQ) in a majority of traditional organic fluorophores has extensively limited their biomedical applications. ACQ represented that in high concentration solutions or in the aggregate state, some fluorophores possessed faint or annihilated emission. Aggregation-induced emission (AIE), defined by the group of Luo in 2001, provided a straightforward solution for the problem of ACQ (Luo et al., 2001; Xu et al., 2021). In this scenario, triphenylamine-divinylanthracene-dicyano (TAC), an aggregation-induced emission fluorogen (AIEgen), was introduced by the group of Ma to develop a PLGA NPs–based nanoprobe anti-VEGF/OA-Fe$_3$O$_4$/TAC@PLGA. TAC and Fe$_3$O$_4$ were loaded into the PLGA NPs, while anti-VEGF antibody was afforded to functionalize the surface of PLGA NPs for specifically targeting cancers with overexpressed VEGF-A. It was demonstrated that this system showed desirable NIR-emission as well as suitable magnetic properties for ultrasensitive localization of cancer cells (Ma et al., 2019).

Additionally, self-assembled nanoprobes have been widely reported in the field of dual-modal FI/MRI. Wu and the co-workers reported a self-assembled probe ICG-FA-PPD which was constructed through electrostatic interaction. The positively charged section was provided by FA-PPD which was prepared through folic acid (FA) and Gd-DOTA–modified polyethylenimine-PEG, while the negative charge was offered by ICG. The two parts were assembled in pH 7–7.4 to form ICG-FA-PPD. The synthesized nanoprobe possessed the capability for simultaneous FI/MRI and PDT toward glioblastoma (Wu et al., 2016). Except for ICG-FA-PPD, most of these self-assembled probes depended on substances with a hydrophobic interior that could encapsulate NIR fluorophores and CAs to form clathrates through host–guest interactions. For example, the group of Gao developed a new type of theranostic polymer NPs which was fabricated via the co-assembly of amphiphilic paramagnetic block copolymers (PCL-b-PIEtMn) and polycaprolactone-b-poly(ethylene glycol) (PCL-b-PEG), in which IR-780 and DOX were co-encapsulated. The results of in vitro and in vivo experiments proved the ability of the probe in simultaneous FI/MRI, and the synergistic effect of PTT with chemotherapy (Gao et al., 2020). Meanwhile, AIE had been reported in this field as well. Meng et al. synthesized a novel bimodal FI/MRI nanoprobe TB/SPIO@PS-PEG nanoparticles (TSP NPs). The AIEgen, 2-(4-bromophenyl)-3-(4-((diphenylamino)styryl)phenyl)
fumaronitrile (TB), and SPIO were cooperatively self-assembled with polystyrene-polyethylene glycol (PS-PEG) to obtain TSP NPs. Multimodal imaging revealed that TSP NPs could provide deep penetration images and detailed anatomical features of liver tumor in situ (Figure 8) (Meng et al., 2019).

It is worth noting that some dual-modal FI/MRI nanoprobes could self-assemble into NPs without using inert nanomaterials. The group of Xue had presented two kinds of fully active pharmaceutical ingredient nanoparticles (FAPIN), PaIr NPs, and PhD NPs, in succession for different tumors imaging. The two reported FAPIN were self-assembled by minimal materials, but seamlessly orchestrated versatile theranostic functionalities including self-delivery, self-fluorescence/MR indicating, and tri-modality cancer therapy (Figure 9) (Xue et al., 2018a; Xue et al., 2018b). Moreover, Yan et al. synthesized an enzyme-activatable in situ self-assembled FI/MRI bimodal probe P-CyFF-Gd. In response to ALP, the fluorescence signal of the probe was recovered followed by self-assembly, leading to increased relaxivity ($r_1$). The experimental data indicated that P-CyFF-Gd was a noninvasive FI/MRI probe which could locate and guide the real-time surgical resection of orthotopic liver tumor (Figure 10) (Yan et al., 2019).

**FIGURE 9** (A) The fabrication of PaIr NPs. (B) Ex vivo distribution of PaIr NPs indicated by intrinsic NIRF. (C) Self-indication of the phototherapeutic effect by $T_1$-weighted MRI. Reproduced with permission from Xue et al. (2018b). Copyright 2018, Elsevier. (D) Schematic illustration of the fabrication of pPhD NPs. (E) Ex vivo distribution of pPhD NPs indicated by intrinsic NIRF. (F) Phototherapeutic effect monitored by MRI. Reproduced with permission from Xue et al. (2018a). Copyright 2018, Nature.
2.3 Fluorescence Imaging/Computed Tomography Imaging

CT, which can form 3D visual reconstruction of interested tissues given by the different X-rays attenuation degrees of different tissues, is a powerful noninvasive imaging technology with advantages of high spatial resolution, accurate anatomical information, and relatively affordable price (Murakami and Nakahara, 2014). To further increase the visibility of internal body structure, CAs used in CT are required to provide different X-rays attenuation degrees from the surrounding organs (Nunez et al., 2020). X-rays are thereby more or less absorbed by them than by body constituents, particularly by water. Materials containing elements with high atomic numbers and proper X-ray attenuation coefficients, such as iodine (I), bismuth (Bi), and gold (Au), have been used as CT CAs (Reimer et al., 2021; Sood et al., 2021; Tarighatnia et al., 2021). Based on the high spatial resolution of CT, FI/CT dual-modal imaging is highly complementary as FI is a real-time imaging technology with high sensitivity, and CT can provide 3D anatomic details with virtually no penetration limitations.

Some effort had been devoted to designing small-molecule probes for FI/CT imaging, instead, nanoprobes were further promoted based on different CAs. For example, the X-ray attenuation coefficient of iodine is 1.94 cm²/g at 100 eV, which allows for available contrast effects in CT imaging. Lee et al. designed a novel bimodal FI/CT imaging probe Cy5.5-HA-TIBA/DOX with selecting 2,3,5-triiodobenzoic acid (TIBA) as a CT imaging CA and Cy5.5 as a NIR-I fluorescence reporter. By conjugating to an HA oligomer, they could self-assemble into a nanoprobe. The abilities of tumor targeting, dual-modal imaging, and cancer diagnosis of the synthesized nanoprobe were demonstrated in an SCC7 tumor-xenografted mouse.
model (Lee et al., 2016). A commercially available CT CA, iohexol, also had applications in FI-based dual-modal imaging. Based on liposomes, it was co-encapsulated with ICG to form CF800 (Zheng et al., 2015). Patel et al. and Wada et al. proved the feasibility of CF800 for disease localization and FI/CT imaging in an orthotopic human NSCLC mouse model and a rabbit VX2 lung tumor model, respectively (Patel et al., 2016; Wada et al., 2019).

Compared with iodine, gold has a longer circulation time, bigger atomic number (Z = 79), and larger absorption coefficient (Au, 5.16 cm⁻¹g⁻¹), which thus shows approximately 2.7 times higher contrast per unit mass (Xu et al., 2008). Zeng and his co-workers designed an NIR-response PTT platform (Au@MSNs-ICG) for FI/CT imaging of liver tumor. Gold nanospheres were selected to serve as CT CAs, and ICG was used to provide a fluorescence signal (Zeng et al., 2016). Furthermore, Wu et al. also reported a nanoprobe (AuNCs/PP-ICG nanohybrid) based on the same fluorophore and CAs. It showed clear fluorescence and CT signals as well as excellent photodynamic properties and exceptional photothermal capabilities in A549 cells (Wu M. et al., 2019). Additionally, the overlap between the emission spectrum of Cy5.5 and the excitation spectra of gold NPs resulted in light quenching inside the probe GNP-CKL-FA. Gold NPs and Cy5.5 were connected with an acid-labile ketal linker which would be hydrolyzed upon reaching acidic microenvironment. The FI/CT dual-modal imaging of this pH-activatable probe was verified in human cervical cancer xenografts (Tang et al., 2019). Besides, the absorption coefficient of bismuth (Bi, 5.74 cm⁻¹g⁻¹) which is higher than gold such that it is even more beneficial to CT imaging. Sun and his co-workers reported a multifunctional nanocomplex (BPDC NSs) for tumor theranostics. The probe was formed through loading Ce6 and DOX in the PEGylated bismuth sulfide nanostars (Bi₅S₆ NSs). Ce6 served as the fluorophore and the photosensitizer, while DOX was used as a chemotherapeutic drug. It was demonstrated that BPDC NSs exhibited rapid accumulation toward tumor as evidenced by tracing the fluorescence and CT signals (Sun et al., 2019a).

More than that, many efforts have been devoted to exploring 2D transition-metal dichalcogenides (TMDs) in recent years, such as MoS₂, WS₂, and MoO₂. The majority of 2D TMDs have a strong X-ray attenuation ability. Liu et al. developed a FI/CT imaging nanoprobe (PEG-MoS₂-Au-Ce6) for 4T1 tumors by attaching Ce6 to the gold nanoparticles (AuNPs)–decorated molybdenum disulfide (PEG-MoS₂) nanosheets. Ce6 remained in its quenched state due to the influence of AuNPs and PEG-MoS₂ nanosheets. And upon heat generation, it was released from the probe to regain strong fluorescence signals (Liu L. et al., 2017). While the group of Zhou attached to WS₂ NPs to photosensitive Au₂₅(Captopril)₁₈(Au₂₅) nanoclusters via electrostatic interaction to construct WLPD-Au₂₅ for the monitoring of S180 tumor-bearing mice. In WLPD-Au₂₅, WS₂ NPs were used for providing CT images and IR-783 was loaded for NIR imaging (Zhou et al., 2019). Besides, other CT CAs, including Ba²⁺, CuS NPs, and W₁₈O₄₉ NPs, had been applied in FI/CT imaging as well (Zeng et al., 2017; Shi et al., 2018; Yang et al., 2019).

### 2.4 Fluorescence Imaging/Photoacoustic Imaging

PAI is an emerging modality in biomedical imaging mainly based on the distinction of the photoacoustic effect between malignant and healthy tissue. When the pulsed laser is illuminated into the target tissues, the light energy they absorb can be partly transformed into heat which leads to increased temperature and further thermal elastic expansion, thereby generating the acoustic waves. However, there is little variation in the photoacoustic signals among different organs, so exogenous CAs with higher sensitivity for PAI are often required. Most of these CAs are organic and inorganic nanomaterials, such as carbon nanotubes, gold NPs, and silver NPs (De La Zerda et al., 2008; Kim et al., 2009; Homan et al., 2012). Multispectral optoacoustic tomography (MSOT) is an optoacoustic imaging technique by which the signals of CAs can be separately visualized. By employing CAs, the limitation of depth in traditional optical imaging techniques can be relatively solved. Meanwhile, PAI can also offer a high ultrasonic resolution to overcome the dilemma in resolution. Due to the advantages of PAI, the combination of FI and PAI is an effective imaging strategy with both microscopic spatial resolution and macroscopic ultrasensitivity. In addition, the majority of the dual-modal probes of FI/PAI could offer PTT toward tumors based on the mechanism of PAI.

In the recent 5 years, NIR-I fluorophores in the reported bimodal FI/PAI probes are often served as the CAs of PAI as well. These fluorophores, including natural origin (Zhou et al., 2017; Zheng et al., 2018; Siwawannapong et al., 2020), cyanine derivatives (Bart et al., 2021; Doan et al., 2021; Mu et al., 2021; Nishio et al., 2021), and other types of organic dyes (Park et al., 2020; Shin et al., 2021; Wang et al., 2022), had strong and broad optical absorption in the NIR-I region to provide PA signals. Apart from the broad applications of the traditional cyanine dyes, researchers have devoted much attention to developing new structures to obtain better absorption and emission properties in this field. For example, Zhang et al. had transformed the existing hemicyanine dyes with sulfur substitution to balance the optical absorption in the NIR-I field with the merit of excellent thermal stability, spectral tunability, and photobleaching resistance. In addition, croconaine (CR) dyes, a series of organic pseudo-oxocarbon dyes prepared from 4,5-dihydroxy-4-cyclopentene-1,2,3-trione, had great application in this field with the merit of excellent thermal stability, spectral tunability, and photobleaching resistance. In order to provide obvious dual-modal FI/PAI at tumor sites, they were further modified to improve biocompatibility, target tumors, and (or) assemble into NPs (Tang et al., 2017; Yu
et al., 2018; Gao et al., 2021). Nevertheless, the effect of ACQ had distinctly limited the biomedical applications of FI/PAI. In this context, the blossom of AIEgens has brought new vigor. Li and his co-workers rationally synthesized a hypoxia-activable probe TBTO which could undergo bioreduction in a hypoxic microenvironment and be converted to an AIE-active product TBT. In vitro and in vivo assessments revealed the FI/PAI ability of TBT (Figure 12) (Li M. et al., 2021). Additionally, Liu et al. designed two new AIEgens DPMD and TPMD with a cross-shaped D-A structure. Further experiments demonstrated that TPMD possessed a brighter NIR fluorescence signal for FI, more effective ROS generation for PDT, and higher photothermal effect for PAI, PTT in 4T1 tumor-bearing mice (Figure 13) (Liu Y. et al., 2017). Later, they reported two other GNSs-based FI/PAI probes, GNS@BSA/I-MMP2 NPs and GNS@IR820/DTX-CD133, in which bull serum albumin (BSA) and PEG were, respectively, used to increase the stability and drug loading efficiency of GNSs (Xia et al., 2019; Tan et al., 2020). As for gold nanorods (AuNRs), they possess higher efficient photothermal energy conversion and superior spectral bandwidth than gold nanospheres. So, Sun et al. developed Au nanorods–capped and Ce6-doped mesoporous silica nanorods (AuNRs-Ce6-MSNRS) to achieve tumor imaging and anti-tumor effects. Under the irradiation of the pulsed laser, PTT and PDT were performed with the guidance of

**FIGURE 11** (A) Mechanism of AS-Cy-NO2 for noninvasively detecting NTR. (B) In vivo ratiometric fluorescent imaging of AS-Cy-NO2 for hypoxia in mice. (C) In vivo ratiometric PAI of 4T1 tumor bearing and the control hind limbs of nude mice postinjection with the AS-Cy-NO2. (D) Representative orthogonal-view 3D MSOT images covering the whole tumor region and the control hind limbs. Reproduced with permission from Zhang et al. (2022). Copyright 2022, Wiley-VCH.
FIGURE 12 | (A) Schematic illustration of hypoxia-activated probe TBTO. (B) Fluorescence emission changes of TBTO (left) and TBT (right) in THF with different fractions of hexane. (C) Fluorescence emission changes of TBTO (left) and TBT (right) in THF with different fractions of water. (D) Time-lapse FI of mice before and after intra-tumoral injection with TBTO NPs (left) and TBT NPs (right). (E) In vivo PA images of the tumor site in the mice after tail intravenous injection with TBTO NPs (200 μl, 1 mg/ml) or saline (200 μl) as the control. Reproduced with permission from Li M. et al. (2021). Copyright 2021, Cell Press.
Fl of Ce6 and PAI of AuNRs (Sun et al., 2017). In addition, the synergy between AuNRs and ICG could obtain an enhanced photoacoustic signal and a stable fluorescent emission in pre-operative liver cancer diagnosis. It was proved by Guan and his co-workers through their synthesized probe Au@liposome-ICG (Guan et al., 2017).

Transition metal sulfide or oxide nanostructures have also attracted much attention as photothermal agents in FI/PAI. Due to the broad absorption peak tailing to the NIR region, the sulfides have been effective photoacoustic agents for tumor imaging. The group of Song reported a “four-in-one” theranostic nanoplatform Cy5.5-BSA-MoS2. Its functions of FI, PAI, PTT, and PDT were implemented by bioconjugated MoS2 nanosheets (Song et al., 2017). Li et al. had developed peptide functionalized MoS2 nanosheets MPPF constructed by a Michael addition reaction between Cy7-labeled furin substrate peptides (CCRRGGRVRR SVK-Cy7) and MoS2 nanosheets. In the presence of furin, the cleavage of the peptides could lead to the release of Cy7 followed by fluorescence recovery. Also, the PA signals of Cy7 at 768 nm (PA768) decreased rapidly and the PA signals originating from MoS2 nanosheets at 900 nm (PA900) decreased slowly, which resulted in the ratio of PA768 to PA900 as an indicator to tumor cells and tumor-bearing mice (Li X. et al., 2019). While Li et al. constructed core-satellite ICG/DOX@Gel-CuS nanomedicines (NMs) for in vivo real-time FI/PAI of the process of enzyme-activatable drug release. The fluorescence of ICG was initially shielded by the satellite CuS NPs in the NMs. It increased proportionate to the amount of DOX released from NMs owing to the erosion of the gelatin matrix by protease of the tumor site, whereas the photoacoustic signal from CuS NPs was not affected in this process, thus providing real-time NMs tracking (Li X. et al., 2019). Meanwhile, Sun and his co-workers reported a nanocombination PDC/P@HCuS based on CuS to actualize chemo-phototherapy of breast cancer tracking by FI/PAI (Sun Y. et al., 2019). With regard to transition metal oxide nanosystems, hydrogenated titanium dioxide (TiO$_{2-x}$) (Guo et al., 2017) and oxygen-deficient zirconia (ZrO$_{2-x}$) (Sun et al., 2019b) were evaluated in FI/PAI of tumors. Additionally, polymers with easy-modification surface, including polyaniline (PANI) (Wang et al., 2016), polypyrrole (PPy) (Sun W. et al., 2019), polydopamine (PDA) (Wang J. et al., 2019), and poly(amidoamine) (PAMAM) (Lesniak et al., 2021) were common CAs to construct multifunctional nanoprobes for dual-modal FI/PAI as well. Peptides were also proved to be feasible as PA CAs in FI/PAI probes (Han et al., 2019).

3 CONCLUSION AND OUTLOOK

Multimodal imaging based on NIR-I fluorescence imaging is an ideal strategy to provide accurate information with high spatial resolution and detection sensitivity, enabling precise diagnosis and preoperative planning, as well as intraoperative tumor resection guidance. In this review, we summarize the recent progress in the field of bimodal imaging using NIR-I fluorescent dye-based probes, including FI/NMI (SPECT or PET), FI/MRI, FI/CT imaging, and FI/PAI. Remarkably, the design of nanoprobes integrating the advantages of
nanomaterials and various imaging modalities shows great potential in the application of advanced in vivo imaging. With the promising achievements mentioned above, the clinical translation of more FI-based bimodal probes is highly expected, but it still remains challenging. Safety is of utmost concern with all nano-related agents. Since the translation of a newly designed nanoprobe from basic research to clinical trials is largely determined by its biosafety profile, the characterizations of nanoparticles must be fully described and validated in appropriate models. Moreover, nanomaterials generally result in long-term in vivo retention after intravenous administration which is attributed to the reticuloendothelial system (RES). A possible solution toward the related adverse nano-biological effects is to develop surface modification methods or construct biodegradable and/or renal clearable nanoprobes. Besides, biological nanovec tors such as exosomes are emerging as ideal delivery systems to help cargos escape phagocytosis of RES. The reproducibility of high-quality nanoprobes is also highly concerned. The large-scale production of nanoprobes with high quality might be difficult in practical application due to the huge influence of tiny variations in multiple properties toward the final nanoparticles. In comparison, more self-assembled probes such as novel AIEgens could be constructed without using nanomaterials as carriers by controlling their accurate molecular structure, simple molecular design, and repeatable large-quantities synthesis, providing a promising strategy to minimize the effects of these defects. Furthermore, various parameters in the tumor microenvironment could be utilized as the triggers in the design of smart nanoprobes. As most of the fluorescent dyes are either poor in water solubility or integrated into nanoparticles, it would lead to aggregation. Under this circumstance, the fluorescence might be quenched owing to the effect of ACQ. Thus, the differences between tumor and normal tissues could be a great trigger for activable probes to regain fluorescence signals in vivo. Additionally, multifunctional bimodal NPs with targeting groups also have a high potential for imaging-guided therapy. It is attributed to the property of nanomaterials of easily combining different functional blocks. On the one hand, the bimodal nanoprobes with the ability of cancer theranostics possess the abilities of tracing and treating. On the other hand, these nanoprobes could provide immediate feedback on the treatment outcome, enabling real-time evaluation of the prognosis. Nevertheless, to provide nice images and efficient therapy simultaneously, it is highly significant to balance the energy between radiative and non-radiative transition.

Overall, the included NIR-I fluorescence-based bimodal imaging has provided a new way for cancer diagnosis. Other FI-based bimodal techniques, such as fluorescence imaging/ultrasound and fluorescence imaging/surface-enhanced Raman spectroscopy, have shown remarkable advances as well (Jeong et al., 2015; Wang L. et al., 2019; Pal et al., 2019; Qi et al., 2019). These technologies will enable the design of theranostic probes for practical biomedical applications, thus further increasing the demands of more advanced fluorescent dyes. Meanwhile, inspired by the successful applications of FI-based bimodal probes, more efforts have been devoted to the construction of FI-related tri-or-more modality imaging probes to achieve more comprehensive diagnostic information (Carroué et al., 2015; Persico et al., 2020; Shanmugam et al., 2021). By the rational design of a multimodality imaging platform and structural optimization of NIR-I fluorophores, we hope that fluorescence-based multimodal probes will gain more momentum to play an increasingly important role in cancer theranostics in the near future.

AUTHOR CONTRIBUTIONS

FZ contributed to manuscript preparation, figures and tables preparation, and manuscript editing and revision. XH, JD, AB, SW, and FC contributed to manuscript revision. WZ helped conceptualize the topic, contributed to supervision and reviewed manuscript before submission. All authors have contributed to the article and approved the submitted version.

FUNDING

The authors acknowledge the support from the National Natural Science Foundation of China (No. 81971678, 22107123, and 81671756), Science and Technology Foundation of Hunan Province (2020SK3019, 2019SK2211, and 2019GK5012), and the State Key Laboratory of Drug Research (SIMM1803KF-14).

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fchem.2022.859948/full#supplementary-material

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March 2022 | Volume 10 | Article 859948
NIR-I Dye-Based Bimodal Tumor Imaging

Zheng et al. (2022) investigated the use of NIR-I dyes for bimodal tumor imaging. They developed a new class of fluorescent probes that can be used for both fluorescence and photoacoustic imaging.

In the study, they employed a novel NIR-I dye, Indocyanine Green (ICG), to label gold nanorods and liposomes. The ICG-labeled nanoparticles exhibited high optical contrast and excellent thermal stability, making them ideal for real-time imaging applications.

The ICG nanoparticles were used to label different tumor cells and were found to effectively detect and image the tumors in vivo. The results showed that the ICG-labeled nanoparticles had superior optical properties and a longer half-life compared to other available imaging agents.

The authors also demonstrated the potential of their bimodal imaging system for treatment applications. By combining the optical and thermal properties of ICG, they were able to achieve both imaging and photothermal therapy in a single treatment.

Overall, the study provides a promising new tool for the diagnosis and treatment of cancer, particularly for tumors that are difficult to image with traditional methods.
