Positron Emission Tomography Image-Guided Drug Delivery: Current Status and Future Perspectives

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ABSTRACT: Positron emission tomography (PET) is an important modality in the field of molecular imaging, which is gradually impacting patient care by providing safe, fast, and reliable techniques that help to alter the course of patient care by revealing invasive, de facto procedures to be unnecessary or rendering them obsolete. Also, PET provides a key connection between the molecular mechanisms involved in the pathophysiology of disease and the according targeted therapies. Recently, PET imaging is also gaining ground in the field of drug delivery. Current drug delivery research is focused on developing novel drug delivery systems with emphasis on precise targeting, accurate dose delivery, and minimal toxicity in order to achieve maximum therapeutic efficacy. At the intersection between PET imaging and controlled drug delivery, interest has grown in combining both these paradigms into clinically effective formulations. PET image-guided drug delivery has great potential to revolutionize patient care by in vivo assessment of drug biodistribution and accumulation at the target site and real-time monitoring of the therapeutic outcome. The expected end point of this approach is to provide fundamental support for the optimization of innovative diagnostic and therapeutic strategies that could contribute to emerging concepts in the field of “personalized medicine”. This review focuses on the recent developments in PET image-guided drug delivery and discusses intriguing opportunities for future development. The preclinical data reported to date are quite promising, and it is evident that such strategies in cancer management hold promise for clinically translatable advances that can positively impact the overall diagnostic and therapeutic processes and result in enhanced quality of life for cancer patients.

KEYWORDS: positron emission tomography, image-guided drug delivery, cancer, theranostics, molecular imaging, personalized medicine

INTRODUCTION

Targeted drug delivery guided by molecular imaging approaches is a burgeoning area of clinical research, particularly for the treatment of cancer.1−5 This approach involves an optimized delivery of a therapeutic molecule and an imaging probe to the disease site, thereby using selective diagnosis and effective pharmacotherapy in unison for management of several diseases. Successful utilization of this strategy requires integrated knowledge and versatile approaches in multidisciplinary fields such as cell and molecular biology, chemistry, material science, and physics and has opened up vast prospects in pharmacokinetics, therapeutic target discovery, drug delivery research, and quantification of multiple biomarkers in diseases. The major goal of this approach is to use molecular imaging to maximize effective therapy in diseased tissues and to minimize systemic drug exposure in order to reduce toxicities. In the past decade, innumerable studies have been reported on the synergistic use of molecular imaging with targeted drug delivery, and this strategy has now matured with promises to fulfill the vision of “personalized” medical treatment in the near future.5−9

In order to minimize the effects of toxicity and improve therapeutic effects, it is essential to deliver the therapeutic drugs to the right site, in the right time, and in the right concentration. Ideally, the drug should act as a “magic bullet” that possesses perfect specificity to targeted lesions and has no side effect on the rest of the body. Controllable and selective delivery of drugs improves bioavailability by preventing premature degradation and enhancing uptake, maintains drug concentration within the therapeutic window by adjusting the drug release rate, and reduces side effects by targeting to disease site and target cells.1,10 The ability to deliver therapeutic drugs
locally, in a minimally invasive manner, has advanced drastically with the growth of molecular imaging techniques.\textsuperscript{11–17} Molecular imaging approaches have been implemented in areas ranging from new therapeutic target discovery to effectively monitoring tumor pharmacokinetics and drug distribution to modulation of drug release at the target site.\textsuperscript{1,4,6,18} When molecular imaging probes are coadministered as part of the drug delivery system, it can help to achieve multiple goals, such as real-time and concurrent assessment of drug delivery efficiency/targeting, \textit{in vivo} fate of drug and sites of localization/accumulation, modes of excretion, imaging, and monitoring the progress of drug treatment, in a single dosing. When an image-guided approach is not used, there is neither any means to track or image the \textit{in vivo} fate nor the ability to measure the delivery efficiency of drugs. Also, the bioavailability, therapeutic efficacy, and dose response of drug treatment has to be estimated based on separate sets of experiments which might render the process cumbersome and cost-ineffective. However, molecular imaging of the drug delivery process involves several challenges and is affected by several factors such as target expression, type of drug, \textit{in vivo} accessibility of the receptor (e.g., vascular density, vascular permeability, and interstitial pressure), enhanced permeability and retention (EPR) effect, receptor internalization, tracer protein dose, and timing of imaging.\textsuperscript{5,6,7,10,11} Nevertheless, this approach has the potential for patient selection for targeted therapy and monitoring the therapeutic response after the drug is delivered.

Currently, several noninvasive image-guided modalities are being used in biomedical and clinical settings, which include magnetic resonance imaging (MRI), computed tomography (CT), positron emission tomography (PET), single photon emission computed tomography (SPECT), optical imaging, and ultrasonography.\textsuperscript{14,16,19–21} Among these, PET, SPECT, and optical imaging are regarded as quantitative or semi-quantitative imaging modalities, whereas CT and MRI are normally used for anatomical imaging. The relative advantages and limitations of these imaging modalities have been elaborately discussed in several review articles.\textsuperscript{14,16,19–21} In particular, PET offers picomolar sensitivity and is a fully translational noninvasive functional imaging technique with high sensitivity and accurate quantification and thus helps in measuring biological processes at the molecular and the metabolic levels \textit{in vivo}. However, the limited spatial resolution of the PET images might sometimes make it difficult to accurately define the regions of interest (ROIs).\textsuperscript{19} Unnecessary radiation exposure to the nontargeted organs due to highly energetic \textit{γ}-rays (511 keV) emitted by the PET radioisotopes is also a cause of concern. Nevertheless, one has to acknowledge that no single imaging modality can provide information on all aspects of structure and function. Therefore, investigation of a subject using multiple imaging modalities is highly desirable and is rapidly gaining popularity.\textsuperscript{20–23}

In this review, we aim to provide a timely and comprehensive overview of the PET image-guided drug delivery approaches reported to date, with focus on quantitative assessments of tumor-targeted therapeutic delivery, distribution, uptake, and response. The development of various carriers for site- and event-specific targeting and controlled drug release are summarized, and the great potential and intriguing opportunities for future development which might help in bringing this exciting research avenue closer to a clinical reality are discussed.

\section*{PET IMAGING}

The interest in using PET as a molecular imaging modality in clinical research has steadily grown during the last 2–3 decades, and has now gained considerable importance in routine hospital practices because of its ability to diagnose diseases in early stages and monitor therapeutic responses.\textsuperscript{13,19,24–27} In a typical scenario of PET imaging, a suitable compound is radioabeled with positron-emitting radionuclides such as \(^{18}\text{F}\), \(^{64}\text{Cu}\), \(^{89}\text{Ga}\), or \(^{89}\text{Zr}\) and administered to a living subject.\textsuperscript{19} The positron that emerges from the radionuclide decay travels a short distance before being annihilated with an electron to release two 511 keV \textit{γ}-rays, which are approximately 180⁰ apart. The 511 keV \textit{γ}-rays can be detected by a ring of detectors configured in the coincidence mode in the PET camera. The registered events are reconstructed into a three-dimensional image which provides information on the spatial distribution of the radioactivity as a function of time in the living subject.\textsuperscript{19} Nowadays, PET is increasingly used in combination with CT as a hybrid imaging modality in clinical settings, to obtain higher resolution by fusing both functional and anatomical information at the same time.

PET also plays an important role in the process of drug development and evaluation, whereby understanding drug action and establishing dosage regimens and treatment strategies have been most crucial.\textsuperscript{28–32} Positron-emitting radionuclides of elements such as C, N, O can replace the stable analogues in drugs and biomolecules, and hence it is possible to synthesize PET probes with the same chemical structure as the parent unlabeled molecules without altering their biological activity. Low bioavailability, insufficient targeting, and poor localization in desired tissue/organ, adverse side effects, etc. are some of major concerns with most of the systemic drug delivery approaches.\textsuperscript{10} Targeted drug delivery systems have the potential to improve these undesirable features, and when used in conjunction with PET imaging, they are effective in increasing safety to efficacy ratio and decreasing dose, which in turn reduces adverse reactions and toxicity of drugs. PET can also provide information on the kinetics, dosimetry, and distribution of drugs in the diseased and normal tissues within the field of view as well as the clearance pattern in a biological system.

PET image-guided drug delivery is expected to play an increasingly important role in realizing the full potential of the next generation of therapeutics. For this purpose, it is essential to choose radioisotopes of appropriate half-lives to match the pharmacokinetics of the drug carriers used. Generally, for inorganic drug carriers (such as silica nanoparticles, superparamagnetic iron oxide nanoparticle, gold nanoparticles, quantum dots etc.) which are expected to have circulation half-lives of a few hours,\textsuperscript{33–36} short-lived or intermediate-lived radioisotopes such as \(^{64}\text{Ga}\) (\(t_{1/2} = 68\) min), \(^{15}\text{F}\) (\(t_{1/2} = 109.8\) min), \(^{48}\text{Sc}\) (\(t_{1/2} = 3.9\) h), \(^{68}\text{Ga}\) (\(t_{1/2} = 9.7\) h), \(^{68}\text{Cu}\) (\(t_{1/2} = 12.7\) h), etc. are more suitable. However, for organic drug carriers (such as carbon nanotubes, polymeric nanoparticles, micelles, liposomes, etc.), which can circulate \textit{in vivo} for more than 1 day,\textsuperscript{37–39} intermediate-lived or long-lived radioisotopes such as \(^{68}\text{Ga}\) (\(t_{1/2} = 9.7\) h), \(^{64}\text{Cu}\) (\(t_{1/2} = 12.7\) h), \(^{89}\text{Zr}\) (\(t_{1/2} = 78.4\) h), or \(^{124}\text{I}\) (\(t_{1/2} = 4.17\) day) would be the ideal choices for PET image-guided drug delivery. The choice of suitable radioisotopes is also governed by the conjugation strategies adopted for radiolabeling the drug carrier. The radiolabeled agent must
peptides, and other small molecules.9,40,41 The carriers are ligands such as monoclonal antibodies, antibody fragments, carrier systems, they can be conjugated with various targeting convenience in modifying the surface properties of these interactions with speciﬁc and antibodies.42 In addition to delivery of chemotherapeutic targeting vectors. Streptavidin/biotin interaction is one good interaction in response to speciﬁc biological system without causing any adverse effects), targeting, and stimulus-responsive (which can control drug biodistribution in response to speciﬁc stimuli).10 In addition to increased selectivity against diseased cells, these delivery systems can also solve problems associated with drug instability in the biological environment as well as issues related to the modulation of drug. Two different approaches are used for drug loading and delivery for pinpoint targeted treatment of cancer cells. In the ﬁrst approach, chemotherapeutic drugs are loaded onto multifunctional drug carriers such as liposomes, micelles, nanoparticles, microparticles, microbubbles, dendrimers, co-polymers, intestinal pathogen, etc.9,40,41 Owing to the convenience in modifying the surface properties of these carrier systems, they can be conjugated with various targeting ligands such as monoclonal antibodies, antibody fragments, peptides, and other small molecules.9,40,41 The carriers are either directly conjugated to targeting ligands or derivatized for interactions with speciﬁc adapters that are conjugated to the targeting vectors. Streptavidin/biotin interaction is one good example used for binding various carriers to targeting proteins and antibodies.42 In addition to delivery of chemotherapeutic drug molecules for therapy, these carriers also carry PET radionuclides or other contrast agents for diagnosis of the diseases. Such drug delivery strategies are an important move toward achieving simultaneous diagnosis and therapy of diseases, which have recently been termed as “theranostics”.9,43

In the second approach, drugs (e.g., therapeutic radionuclides) are conjugated with the targeting ligands using suitable bifunctional linkers.9,44 Unlike the ﬁrst approach, here the drug and the imaging label (PET radionuclide) do not necessarily share the same delivery carrier. For diagnosis or monitoring therapeutic response, PET imaging is carried out separately in this case by conjugation of the targeting ligands with suitable PET radioisotopes. Another striking difference between the two approaches is that, in the former, the delivery of the drug to the target tissue can be achieved by both passive and active targeting, while, in the latter, the drug is delivered primarily due to active targeting.45 In passive targeting, the drug carriers such as nanoparticles, liposomes, micelles, etc. can reach the tumor sites through the EPR effect.45 Also, therapeutic concentrations can be much lower than optimal at the tumor site by simply relying on EPR-mediated accumulation, and therefore passive targeting is generally not preferred for drug delivery. More efﬁcient and selective uptake of drug into the target cells is achieved by active targeting wherein the drug carriers are conjugated with targeting ligands, as mentioned earlier. Active targeting requires careful identiﬁcation of tumor biomarkers, as well as selection of speciﬁc molecules that can bind to such markers in a selective and directed manner. Targeted drug delivery vehicles can then be internalized by tumor cells via receptor-mediated endocytosis/phagocytosis, resulting in elevated concentration of drugs in tumor tissue.

Thus, the concept of “theranostic agent” is not just limited to chemotherapy but also has a relevant role to guide in radiation-based targeted therapies. Various drug carrier systems have been radiolabeled with different positron emitter radionuclides for image-guided drug delivery, most of which are summarized in Table 1 and discussed in the following text.

**CARRIERS FOR PET IMAGE-GUIDED DRUG DELIVERY**

The current revolution in targeted drug delivery is fueled by the innovations in material science, organic chemistry, functional genomics, and proteomics which have created carriers that are biodegradable (which can be slowly dissolved in vivo by biological means), biocompatible (which can remain in a biological system without causing any adverse effect), targeting, and stimulus-responsive (which can control drug biodistribution in response to speciﬁc stimuli).9 In addition to increased selectivity against diseased cells, these delivery systems can also solve problems associated with drug instability in the biological environment as well as issues related to the modulation of drug. Two different approaches are used for drug loading and delivery for pinpoint targeted treatment of cancer cells. In the ﬁrst approach, chemotherapeutic drugs are loaded onto multifunctional drug carriers such as liposomes, micelles, nanoparticles, microparticles, microbubbles, dendrimers, co-polymers, intestinal pathogen, etc.9,40,41 Owing to the convenience in modifying the surface properties of these carrier systems, they can be conjugated with various targeting ligands such as monoclonal antibodies, antibody fragments, peptides, and other small molecules.9,40,41 The carriers are either directly conjugated to targeting ligands or derivatized for interactions with speciﬁc adapters that are conjugated to the targeting vectors. Streptavidin/biotin interaction is one good example used for binding various carriers to targeting proteins and antibodies.42 In addition to delivery of chemotherapeutic drug molecules for therapy, these carriers also carry PET radionuclides or other contrast agents for diagnosis of the diseases. Such drug delivery strategies are an important move toward achieving simultaneous diagnosis and therapy of diseases, which have recently been termed as “theranostics”.9,43

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**Alumun-Based Delivery Approach.** Albumin is an attractive macromolecular carrier that may be modiﬁed suitably for biomedical imaging applications.9 Such carriers have also been studied for drug and gene delivery in vitro and in vivo, through cavitation.46-49 Generally, albumin-based carriers are biodegradable, nontoxic, metabolized in vivo to produce harmless degradation products, nonimmunogenic, easy to purify, and soluble in water allowing ease of delivery by
injection and thus ideal candidates for image-guided drug delivery procedures. A significant amount of drug can be incorporated in the albumin based carrier systems because of different binding sites present in the albumin molecule. Owing to the defined albumin primary structure and high content of charged amino acids (e.g., lysine) on the surface, albumin-based carriers offer the possibility of direct electrostatic adsorption of positively (e.g., ganciclovir) or negatively charged (e.g., oligonucleotide) molecules without the requirement of any other compound.57 In addition, these carriers can easily be prepared under mild conditions by coacervation, controlled desolvation, or emulsion formation.57 Commercially, albumins are obtained with significant quantities from egg white (ovalbumin), bovine serum (bovine serum albumin, BSA), and human serum (human serum albumin, HSA) and also available from soybeans, milk, and grains.57

The chelator-free radiolabeling of macroaggregated human serum albumin with 68Ga (t1/2 = 68 min) for PET imaging was first described by Even et al.50 Subsequently, this procedure was improved, and development of a kit for labeling macroaggregated human serum albumin with 68Ga for PET imaging of liver anomalies was reported by Okada et al.51 The kit was clinically tested and was found useful in the evaluation of the function of the reticuloendothelial system. In a similar study, Maus et al. reported the radiolabeling of different commercially available human serum albumin kits with 68Ga.52 In vivo PET imaging showed that 68Ga-labeled human serum albumin was mainly retained in the lungs. No decrease in activity or migration of particles from the lungs was observed during the first 1 h (~1 half-life of 68Ga), which demonstrated the in vivo stability of the radiolabeled albumin over that period of time. Also, no significant retention of 68Ga-labeled human serum albumin particles in the liver was detected. The authors concluded that this approach could be used to estimate the liver-to-lung shunt and eliminate extrahepatic macroaggregate deposition in patients with primary and secondary liver malignancies, warranting 90Y-based radioembolization therapy.53–55

In a recent development, Liao et al. prepared albumin shielded microbubbles filled with perfluorocarbon (C3F8) gas to enhance the contrast in ultrasound imaging.56 Additionally, the microbubbles were radiolabeled with N-succinimidyl-4-[18F]-fluorobenzoate ([18F-SFB]) and also conjugated with antibodies targeting vascular endothelial growth factor receptor 2 (VEGFR2) using avidin–biotin interaction. The radiolabeled microbubble shells could thus be used as dual-modality (PET and ultrasound) imaging agent. The 18F-labeled, albumin-shelled, VEGFR2-targeted microbubbles had a lifetime of 30 min in the blood pool and demonstrated a highly specific adherence to tumor vessels in mice bearing human breast cancer. The size of the microbubbles was on the order of several micrometers and therefore should be retained in the tumor vasculature after intravenous injection. However, dynamic microPET imaging showed a relatively low tumor uptake of ~1% ID/g, even 1 h post injection. The low tumor uptake might be attributed to attachment of 18F-SFB on the surface of the microbubble, which might have influenced the targeting efficiency of the antibody. The targeted microbubbles accumulated rapidly in both the liver and lung and cleared slowly from the blood circulation. The trends found in microPET imaging were further corroborated by ex vivo biodistribution studies. The specificity of the binding of targeted microbubbles to endothelial VEGFR2 was further validated by comparing the results of targeted and nontargeted contrast-enhanced ultrasound imaging. The authors concluded that the 18F-labeled albumin-shelled microbubbles can be used for targeted drug delivery to VEGFR2 in breast cancer, guided by the dual-modality (PET/ultrasound) functional imaging approach.

In all these studies, development of only imaging strategies using albumin-based platforms have been described without direct relation to drug delivery. However, there are several other reports on utility of drug-loaded albumin-based carriers and controlling drug release using ultrasound energy in such systems.57 Therefore, it was expected that tracking disease progression would be analogous to tracking drug delivery using albumin-based carriers. This hypothesis might also be valid for other drug carrier systems described below.

**Liposome-Based Delivery Approach.** Liposomes are concentric, closed bilayer membranes of water-insoluble polar lipids that can that can be used to encapsulate biomolecules and drugs for targeted delivery while protecting their bioactivity. Soluble drugs can be loaded in the aqueous core and the hydrophobic drugs partitioned in the lipid bilayer. Liposomal carriers are the earliest and the most extensively studied drug delivery carriers,57–58 They are widely used not only in delivery of a variety of anticancer drugs but also in delivery of antineoplastic agents, antimicrobial compounds, immunomodulators, anti-inflammatory agents, cardiovascular drugs, etc.57,58 The widespread interest in the use of liposomal systems for drug delivery stems from their biocompatibility, biodegradability, and nontoxicity and the ease of controlling their size during the preparation process. Currently, there are several commercially available liposomal formulations for cancer therapy, including doxorubicin (Doxil), daunorubicin (Daunoxome), cytarabine (Depocyt), Myocet, and vincristine (ONCO-TCS).57,58 Many liposomal-based drug delivery systems are currently undergoing global clinical trials.57,58 The recent advances in the use of radiolabeled liposomes for imaging as a tool in personalized medicine have been summarized in a recent review.59

Generally, liposomal systems are coated with poly(ethylene glycol) (PEG) to increase the circulation time in blood and decrease uptake in the reticuloendothelial system (RES).50–52 Radiolabeling of liposomes with PET radioisotopes generally requires the use of chelator molecules in the aqueous core or conjugation on the lipid bilayer. The radiolabeled liposomal systems employed in PET studies must be carefully designed as slower stability of radiolabeled agent might obscure image-based assessment of particle pharmacokinetics. Seo et al. reported the development of a method for radiolabeling liposomes with 64Cu for imaging and drug delivery monitoring using PET.65,66 Bifunctional chelators, such as, 6-[(bromoacetamido)benzyl]-1,4,8,11-tetraazacyclotetradecane-N,N’,N”,N”’-tetraacetic acid (TETA-PDP), and 4-(2-(2-pyridyldithioethyl)-ethanamido)-11-carboxymethyl-1,4,8,11-tetraazacyclotetradecane-1,4,8,11-tetraacetic acid (TETA-PDP), and 4-(2-(2-pyridyldithioethyl)-ethanamido)-11-carboxymethyl-1,4,8,11-tetraazabicyclo[6.6.2]hexadecane (CB-TE2A-PDEA) were radiolabeled with 64Cu, and the radiolabeled conjugates were attached to maleimide lipids in the liposome. The radiolabeled liposomes were found to be stable in mouse serum even after 48 h of incubation. In vivo PET studies demonstrated that liposomal activity was high in the blood pool from 0 to 6 h and slowly cleared out through the RES. The presence of the PEG spacer between the chelator and the lipid did not significantly alter the labeling efficiency.
and the clearance rate of liposomes from the blood pool. The study was further extended by the same group of authors to characterize the in vivo clearance of 64Cu-labeled distearoyl and dipalmitoyl lipids included within PEGylated liposomes.\textsuperscript{65} In vivo PET imaging studies established that changes in lipidacyl chain length can result in desorption of lipid from the liposomal anchorage and interaction with blood components. Therefore, this factor should be considered for liposomal PET studies as desorption can rapidly alter the apparent pharmacokinetics.

In another study, Peterson et al. developed a remote loading method using 2-hydroxyquinoline ionophore, to carry 64Cu across the membrane of preformed liposomes and deliver it to an encapsulated copper-chelator.\textsuperscript{66} A highly efficient loading (>95%) and retention stability (>99%) was obtained adopting this approach. In vivo PET imaging studies demonstrated that a maximum tumor uptake of \( \sim 5\% \) ID/g with high tumor to muscle ratio could be achieved. The 64Cu-liposomes reached a maximum level in the liver and spleen after 4 h and subsequently remained at a constant level. Also, the 64Cu-liposomes remained in the blood pool for \( >24 \) h. The method provided 64Cu-labeled liposomes with excellent imaging properties due to the high concentration of 64Cu inside the liposomes and restricted exchange of 64Cu with the biological environment due to the protective barrier constituted by the liposomal membrane. The same group of authors investigated the suitability of 64Cu-labeled liposomes for imaging somatostatin receptor expression in neuroendocrine tumor model.\textsuperscript{67} The peptide octreotate (TATE) was covalently attached to the PEGylated liposomes with an encapsulated positron emitter 64Cu. This peptide is routinely used in clinic for imaging somatostatin receptor-positive tumors by scintigraphy.\textsuperscript{68} In vivo PET imaging and biodistribution studies revealed that the presence of TATE on the liposomes resulted in a significantly faster initial blood clearance in comparison to control liposomes without TATE. There was no significant difference in tumor uptake (\( \sim 5\% \) ID/g in both cases) on using 64Cu-labeled PEGylated liposomes with or without TATE, suggesting that the uptake was mainly due to passive targeting. However, 64Cu-loaded PEGylated liposomes with TATE showed significantly higher tumor-to-muscle (T/M) ratio (12.7 \( \pm 1.0 \)) than the control-liposomes without TATE (8.9 \( \pm 0.9 \)). The tumor accumulation and T/M ratio achieved in this study suggest that liposomal systems might be used as carriers of radionuclides for therapeutic use and also for delivery of chemotherapeutic drugs.

Tumor associated macrophages (TAMs) have been shown to play a major role in the growth and spread of several types of cancer.\textsuperscript{69,70} Locke et al. reported PET imaging of TAMs in a mouse model of pulmonary adenocarcinoma, using mannosone coated liposomes radiolabeled with 64Cu.\textsuperscript{71} In vivo PET imaging and biodistribution studies revealed that radiolabeled mannosylated liposome accumulated in TAMs and exhibited little accumulation in remote lung areas at \( 6 \) h post injection. Further, it was verified by confocal microscopy that the PET signal was due to liposome internalization by TAMs. Urakami et al. developed a methodology for one-step labeling of liposomes with 18F.\textsuperscript{72} Solid-phase transition method was utilized, and high labeling efficiency and visualization of liposomal trafficking in mice by real-time analysis were obtained by PET. The same group reported the development of an efficient method for preparation of 18F-labeled liposome-encapsulated hemoglobin.\textsuperscript{73} Using the radiolabeled liposome, the oxygen transfer even in an ischemic brain could be monitored by dynamic PET. In another study, radiolabeling of PEGylated liposomes with [\( ^{18}\text{F} \)]fluorodipalmitin ([\( ^{18}\text{F} \)]FDP) was reported by Marik et al.\textsuperscript{74} Radiolabeled diglyceride was synthesized by the incorporation of \( ^{18}\text{F} \) into the lipid molecule by nucleophilic substitution of p-toluenesulfonyl moiety. While free [\( ^{18}\text{F} \)]FDP was rapidly taken by the liver, spleen, and lungs, liposome incorporated [\( ^{18}\text{F} \)]FDP was observed to circulate in blood vessels for nearly 90 min.

Adopting the previously reported procedures,\textsuperscript{63,74} 18F and 64Cu-labeled liposomes were prepared by Paoli et al.\textsuperscript{75} The liposomes were preconjugated with suitable fluorophores (calcein or AF-750), for dual-modality PET/optical imaging. A model hydrophilic drug was encapsulated in the liposomal system and administered in mice bearing bilateral Met-1 tumors. Using in vivo PET imaging and ex vivo fluorescent imaging of tumors, the authors could demonstrate that the accumulation of the drug was increased by up to 177-fold by liposomal encapsulation. Recently, Oku et al. reported the radiolabeling of liposomes [modified with PEG or Ala-Pro-Arg-Pro-Gly (APRPG) peptide] with 1-[\( ^{18}\text{F} \)]fluoro-3,6-dioxatetracosane, which enabled imaging of gliomas by PET with higher contrast than that obtained with [\( ^{18}\text{F} \)]fluorodeoxyglucose ([\( ^{18}\text{F} \)]FDG).\textsuperscript{76} The liposomes did not accumulate in the normal surrounding brain tissue due to blood–brain barrier protection, and using this approach, even a very small sized (\( \sim 1 \) mm) brain tumor could be specifically imaged with the radiolabeled liposome (Figure 1).

![Figure 1. PET image-guided tumor targeting using liposome based carrier. PET imaging of brain tumor using PEG-modified liposomes (top panel) and APRPG-modified liposomes (middle panel), labeled with 1-[\( ^{18}\text{F} \)]fluoro-3,6-dioxatetracosane. The other regions of the brain showed a low background. On the contrary, [\( ^{18}\text{F} \)]FDG imaged the whole brain, although the accumulation was higher in the tumor region (bottom panel). Autoradiograms shown in the right panel confirmed the region of tumor. Adapted with permission from ref 76. Copyright 2011 Elsevier.](image-url)
Micelle-Based Delivery Approach. A micelle is an aggregate of surfactant molecules dispersed in a liquid colloid. Micellar structures are important carriers for drug delivery because they can form relatively small and uniform size structures, be prepared from a variety of amphiphatic materials, increase solubility of hydrophobic molecules, and incorporate multiple functionalities into a single structure.79–80 When tagged with suitable contrast agents, these systems can also be used for molecular imaging as well as image-guided drug delivery. Among the various micellar structures, the polymeric micelles are the most extensively used for drug delivery applications.81 The polymeric micelles generally consist of a unique core–shell structure. The inner core is the hydrophobic part of the block copolymer, which encapsulates the water-insoluble drug. The outer shell or corona of the hydrophilic block of the copolymer is often composed of PEG, and it protects the drug from the aqueous environment and also imparts particle stability and excellent dispersibility in an aqueous solution. Owing to these characteristics, polymeric micelles have several advantages as drug carriers such as enhancing the aqueous solubility of hydrophobic drugs, prolonging the circulation time of the drug in the blood, improving the in vivo stability of the drug, providing both passive and active tumor targeting abilities, and reducing nonspecific uptake by the reticuloendothelial system.82

In vivo tumor targeting and drug delivery properties of polydiacetylene (PDA) micelles (diameter ~10 nm) were investigated by Mackiewicz et al. in a breast cancer model.81 Such small sized micelles can better diffuse through blood vessel walls and reach deeper tumor tissues due to the EPR effect. The authors synthesized different micelles with coatings consisting of either nitrioltriacetic acids (NTA) or PEG chains of variable lengths and tested for their ability to passively target tumor. Among them, 2 kDa PEG-coated micelle (PDA-PEG2000) was identified as the most promising carrier in terms of longer blood residence time, higher tumor uptake, and better imaging contrast. Fluorescence diffuse optical tomographic imaging indicated a tumor uptake of ∼3% of the injected dose of PDA-PEG2000. The diffusion of PDA-PEG2000 micelles inside the tumor was further evidenced and quantified by PET imaging using 18F-FDG colocalization. Drug delivery application of the cargo was also assessed using micelles loaded with paclitaxel, a hydrophobic anticancer drug, which showed good in vitro cytotoxicity and in vivo tumor growth inhibition. Thus, the potential of PDA-micelles for drug delivery could be successfully demonstrated in this study. In another study, Cho et al. reported a novel drug delivery strategy using poly(ethylene glycol)-block-poly(ε-caprolactone) (PEG-b-PCL) micelles.82 Three different drugs, namely, paclitaxel (cytotoxic agent), cyclopamine (hedgehog inhibitor), and gossypol (Bcl-2 inhibitor), were loaded on PEG-b-PCL micelles and evaluated in xenograft models of ovarian cancer. Multi-drug-loaded PEG-b-PCL micelles were nanoscopic, fairly stable in aqueous solution, and capable of simultaneous as well as sustained release of each of the three drugs in vitro. In vivo studies based on bioluminescence imaging and 3′-deoxy-3′-[18F]fluorothymidine ([18F-FLT]) PET imaging revealed that multi-drug-loaded PEG-b-PCL micelles had significantly less tumor burden than use of paclitaxel alone. Also, [18F-FLT]-PET images clearly showed that multi-drug-loaded PEG-b-PCL micelles significantly reduced tumor volumes over paclitaxel and vehicle controls and could thus prolong the overall survival. Thus, the authors could establish that the strategy of concurrent delivery of drug combinations of cytotoxic agents and molecular targeted agents using a micellar-based drug delivery vehicle is effective for the treatment of ovarian cancer.

Recently, Benezra et al. evaluated the potential of 18F-labeled dasatinib derivative (SKI249380, a new-generation Src and platelet-derived growth factor receptor (PDGFR) inhibitor83)

SPECT), or 64Cu2+ (for PET) and used for multimodal imaging. The effective radiolabeling and noninvasive imaging strategies developed thus far might aid further research on PET image-guided drug delivery using liposomal carriers in the near future.
Review

In vivo PET imaging studies demonstrated a significantly higher tumor uptake for 18F-SKI249380-loaded micellar formulations (4.9% ID/g) compared to control group (1.6% ID/g). Saturation studies using excess cold dasatinib showed marked reduction of tumor uptake values to levels in normal brain (1.5% ID/g), consistent with in vivo binding specificity. The improved drug solubility, delivery, and kinetic behavior conferred by the use of these micellar 18F-SKI249380 preparations might find utility in treatment of various types of gliomas.

Despite the excellent attributes of the polymeric micelles as carriers for drug delivery and uptake in xenograft models of high-grade glioma,85 In vivo PET imaging studies demonstrated a significantly higher tumor uptake for 18F-SKI249380-loaded micellar formulations (4.9% ID/g) compared to control group (1.6% ID/g). Saturation studies using excess cold dasatinib showed marked reduction of tumor uptake values to levels in normal brain (1.5% ID/g), consistent with in vivo binding specificity. The improved drug solubility, delivery, and kinetic behavior conferred by the use of these micellar 18F-SKI249380 preparations might find utility in treatment of various types of gliomas.

These limitations could be circumvented with the use of suitably engineered unimolecular micelles possessing excellent in vitro and in vivo stability.86,87 The synthesis of a multifunctional unimolecular micelle made of a hyperbranched amphiphilic block copolymer, Boltorn H40-poly(1-glutamate-hydrazone-doxorubicin)-b-poly(ethylene glycol), for PET image-guided drug delivery was reported by Xiao et al.88 The copolymer was conjugated with cyclo(Arg-Gly-Asp-D-Phe-Cys) peptides (cRGD) for integrin αβ3 targeting and macrocyclic chelators (1,4,7-triazacyclononane-N,N′,N″-triacetic acid [NOTA]) for 64Cu-labeling and PET imaging. The anticancer drug doxorubicin (DOX) was covalently conjugated onto the hydrophobic segments of the amphiphilic block copolymer arms via a pH-labile hydrazone linkage to enable pH-controlled drug release. In vivo PET imaging and biodistribution studies in U87MG tumor-bearing mice showed higher tumor uptake for cRGD-conjugated unimolecular micelles (~7% ID/g) than nontargeted micelles (~2.5% ID/g) (Figure 2). Additionally, cRGD-conjugated unimolecular micelles exhibited a much higher cellular uptake in U87MG human glioblastoma cells than nontargeted unimolecular micelles due to integrin αβ3-mediated endocytosis, thereby leading to a significantly higher cytotoxicity when the micellar systems were conjugated with DOX. The same group of authors reported the synthesis of another unimolecular micelle formed by dendritic amphiphilic block copolymers poly(amideamine)-poly(l-lactide-b-poly(ethylene glycol) conjugated with an anti-CD105 monoclonal antibody (TRC105) and NOTA for PET image-guided drug delivery.87 DOX was loaded into the hydrophobic core of the unimolecular micelles. As observed in the previous study, 64Cu-labeled targeted micelles exhibited a much higher level of tumor accumulation than 64Cu-labeled nontargeted micelles, measured by serial noninvasive PET imaging and confirmed by biodistribution studies in murine breast tumor-bearing mice. Thus, these multifunctional unimolecular micelles possessing passive and active tumor-targeting abilities, pH-controlled drug release, and PET imaging capabilities are potentially important drug delivery vehicles for image-guided therapy.

**Enzyme/Prodrug-Based Delivery Approach.** Enzyme/prodrug therapy is one of the most promising strategies where systemic toxicity can be minimized while maintaining the therapeutic efficacy.88–90 In this process, a drug-activating enzyme is targeted or expressed in cancer cells, following which a nontoxic prodrug is administered systemically.91,92 The enzyme converts the prodrug to an active anticancer drug, achieving high concentrations in the tumor and sparing the normal tissues. However, there are certain requirements for this strategy to work in clinical context. The enzyme should be non-human or expressed at very low concentrations in the normal tissue and should have high enzymatic activity. The prodrug should be a good substrate for the enzyme but should not be activated in nontumor tissues. While the prodrug should be.

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**Figure 3.** PET image-guided tumor targeting using enzyme/prodrug approach. (A) PET imaging using 18F-FIAU to identify the location, magnitude, and extent of vector-mediated gene expression in gene therapy for recurrent glioblastoma. Treatment responses were recorded by PET imaging with 11C-MET. The region of specific 11C-MET retention within the tumor after HSV-1-tk-transduction (white arrow) showed the signs of necrosis (cross hairs, right column and reduced methionine uptake [MET]) after ganciclovir treatment. Adapted with permission from ref 98. Copyright 2001 Elsevier. (B) PET imaging of HSV-1-tk activity in tumors after Sindbis/tk infection. Tumor-bearing mice either received no vector treatment (Tumor +, Sindbis/tk −) or received 3 Sindbis/tk treatments via intraperitoneal injection far away from sites of tumor inoculation (Tumor +, Sindbis/tk +). HSV-1-tk activity was determined after intravenous administration of 18F-FEAU as tracer. Tumors on the right shoulder of SCID mice are indicated by yellow arrows, and white arrows indicate activity in urinary bladder. Adapted with permission from ref 102. Copyright 2006 Society of Nuclear Medicine and Molecular Imaging.
nontoxic, the activated drug should be highly toxic and diffusible to be taken up by the adjacent cells for a “bystander cell kill effect”. Ideally, the activated drug should not leak out into the systemic circulation. Currently, there are three major categories of enzyme/prodrug strategies: (a) delivery of genes that encode prodrug-activating enzymes into tumor tissue (gene encoding prodrug activating enzyme therapy, GDEPT, and virus-directed enzyme prodrug therapy, VDEPT), (b) targeted delivery of active enzymes in tumor tissue where the therapeutic enzyme is conjugated with an antibody, small molecular ligand, or peptide that binds to antigens preferentially expressed on the surface of tumor cells or in the tumor vasculature or interstitium (targeting group-directed enzyme/prodrug therapy, TDDEPT), and (c) vasculature permeability-dependent enzyme/prodrug therapy (VPDDEPT) in which the intratumoral delivery of the enzyme is realized through the higher permeability of tumor vasculature. 91-94 PET image-guided enzyme/prodrug strategies have been extensively reviewed and hence will be discussed briefly in the following text.99-96

Most of the studies in PET guided enzyme/prodrug based cancer therapy are based on the GDEPT approach. Bankiewicz et al. developed a strategy which combined gene therapy with aromatic L-amino acid decarboxylase (AADC) gene and a prodrug, dopamine.97 Using this approach, the authors could synthesize and regulate the neurotransmitters involved in Parkinson’s disease. In vivo PET imaging using AADC tracer, 6-18F]fluoro-L-m-tyrosine (FMT) could measure the gene expression and thus establish the potential of enzyme/prodrug approach in delivery of therapeutic agents to the central nervous system. Also, the extent of gene expression could be effectively used to predict the therapeutic response. This approach could be further validated in another study where PET imaging with 124I-labeled 2’-fluoro-2’-deoxy-1b-d-arabinofuranosyl-5-iodo-uracil (124I-FIAU), a specific marker substrate for expression of the herpes simplex virus type 1 thymidine kinase (HSV-1-tk) gene, was used to identify the location, magnitude, and extent of vector-mediated gene expression in a phase 1/II clinical trial of gene therapy for recurrent glioblastoma.98 In this study, dynamic 18F-FIAU-PET scans were done before gene transduction to assess the basal state of FIAU-accumulation and washout of the tumor, and also after vector application to investigate whether specific FIAU-accumulation did occur (Figure 3A). Ganciclovir treatment (5 mg per kg twice a day over 14 days) was done starting 4 days after vector infusion. Treatment responses were recorded by repeated MRI as well as PET with 18F-FDG and 11C-labeled methionine (11C-MET). The same PET tracer (124I-FIAU) was used by Hackman et al. to assess the potential of double prodrug activation gene therapy using the Escherichia coli cytosine deaminase (CD)-HSV-1-tk fusion gene (CD/TK) for treatment of different tumors.99 PET imaging was used for monitoring expression of the CD/TK fusion gene, and the different levels of CD/TK expression in tumor models could be imaged quantitatively. The results of these studies could be utilized to develop standardized gene therapy protocols adopting enzyme/prodrug strategy for human subjects.

The synthesis of half-mustard prodrug, 4-[2-(2-chloroethyl)(2-ethyl) amino]-phenoxycarbonyl-L-glutamic acid, by reductive alkylation of 4-[2-(chloroethyl)amino]-phenoxycarbonyl-L-glutamic acid was reported by Malik et al.100 The prodrug was radiolabeled with 13C, and its potential for imaging antibody- and gene-directed enzyme prodrug therapy with PET could be established. The use of another prodrug, 1-(2-deoxy-2-fluoro-β-β-arabinofuranosyl)uracil (FAU), for treatment of tumors with high thymidylate synthase catalytic activity was reported by Eiseman et al.101 This prodrug was activated by thymidylate synthase enzyme. PET imaging using 18F-FAU was used to visualize tumors that have high thymidylate synthase catalytic activity. However, the authors did not observe high localization of 18F-FAU in tumors compared with background, which might limit further utilization of this PET probe in clinical context. In another study, selective tumor targeting and quantitative in vivo monitoring using PET of a commonly applied GDEPT, based on HSV-1-tk and ganciclovir (GCV), was reported by Tseng et al.102 Sindbis virus was used to deliver the HSV-1-tk suicide gene to tumor cells for subsequent GCV activation and tumor killing. PET imaging using 18F-labeled fluoro-ethyl-arabinosyluridine (18F-FAU) was used to monitor the HSV-1-tk activity in tumor cells after parenteral administration of Sindbis virus (Figure 3B). High tumor uptake of 18F-FAU (~3% ID/g) proved that the Sindbis vector efficiently targeted the HSV-1-tk enzyme gene into the infected tumor cells. Also, PET imaging could be used to monitor HSV-1-tk activities after systemic Sindbis vector treatments for determining the levels and tissue distribution of the vector and optimizing efficient prodrug activation for more accurate treatment planning and monitoring. This study was further extended by Stelter et al., where different molecular imaging strategies, such as bioluminescence, fluorescence molecular tomography, and PET, were used to evaluate Sindbis virus mediated infection of tumor cells in vitro and in vivo.103 The authors concluded that the Sindbis virus infection rates were not solely dependent on cellular laminin receptor expression and other factors such as cellular infection and viral replication might also be responsible. In another similar study, Wang et al. evaluated the efficacy of 4 different radiotracers, 123I-S-iodo-29-fluoro-1-b-d-arabinofuranosyluracil (123I-FIAU), 5,18F-fluoro-29-deoxyuridine (18F-FUdR), 2,18F-fluoroethyl-L-tyrosine (18F-FET), and 18F-FDG for monitoring tumor responses using SPECT or PET during prodrug activation gene therapy with HSV-1-tk and GCV.104 Based on tumor uptake of the radiotracers, 18F-FUdR was identified as the most suitable radiotracer for assessment of responses in tumors undergoing HSV-1-tk and GCV prodrug activation gene therapy.

The enzyme β-glucuronidase (β-GUS) has recently been investigated as a target in prodrug therapy for cancer.105-107 In order to optimize β-GUS-based prodrug therapies, a PET tracer, 18F-labeled 1-O-(4-(2-fluoroethyl-carbamoyloxymethyl)-2-nitrophenyl)-O-β-d-glucopyronuronate (18F-FEA) was evaluated for imaging of β-GUS in tumor (C6 gliomas) and inflammation models.106 In vivo PET imaging and biodistribution studies showed high uptake of the radiotracer in tumor, high target to nontarget ratio, and rapid renal clearance. In inflammation model, the uptake of the radiotracer in inflamed muscle was significantly higher than in control muscle, thereby establishing the potential of this radiotracer to detect increased activity of β-GUS. The extent of β-GUS release in small C6 glioma tumors after a single treatment of doxorubicin (DOX), carmustine (BCNU), and tumor necrosis factor α (TNF-α) with 18F-FEA PET was evaluated by the same group of authors.107 PET studies confirmed that β-GUS was released in vivo and the distribution volume of 18F-FEA in C6 gliomas was increased significantly. These results were further confirmed by histochemical analysis and flow cytometry. The results obtained in this study demonstrate the potential of
a two-step chemotherapy−prodrug approach, in which tumors are treated with a single dose of a cytostatic drug before prodrug treatment. Recently, Moon et al. reported the synthesis of 18F labeled 1-(3-furyl)-4-hydroxy-5-fluoro-1-pentanone (18F-F-4-IM), which can be metabolized by the CYP4B1 enzyme and used for PET imaging of tumors and monitoring enzyme-activating anticancer prodrugs. Biodistribution studies in normal rats showed that the uptake of 18F-F-4-IM was high in the lung, where CYP4B1 gene is preferentially expressed. The results were further confirmed by in vitro cell assays, and the potential of 18F-F-4-IM for imaging of CYP4B1-transfected tumor cells and monitoring CYP4B1 enzyme/prodrug interactions could be demonstrated. It is envisaged that further development of PET guided enzyme/prodrug protocols would significantly facilitate their clinical translation with high safety and reliability.

**Nanoparticle-Based Delivery Approach.** In the past two decades, the applications of nanotechnology in cancer diagnostics and therapy have attracted widespread interest and a variety of functional nanoparticles have been developed and evaluated for drug delivery, diagnostic sensors, imaging agents, and labeling probes. Nanoparticles used for this purpose vary with a size from 1 nm to few hundred nanometers and surface charge varying from negative to positive and even...

![Figure 4. PET image-guided tumor targeting using nanoparticle based carrier.](image-url)
neutral. Particularly as drug delivery vehicles and molecular imaging tools, targeted nanoparticle based systems hold significant promise by virtue of their controllable size, high surface area to volume ratio, and customized internal and external chemistries. The major advantages of using the engineered nanoparticle based systems for such applications include (a) the ease of particle functionalization for conjugation with suitable targeting vectors such as peptides or antibodies, (b) the ability to deliver a higher concentration of contrast agent for every targeted binding event to achieve higher detection sensitivity which might permit diagnosis of the disease in its very early stage, and (c) improved treatment effects when used as drug carriers by protecting entrapped drugs from degradation, enhancing tumor uptake through the enhanced permeability and retention effect as well as receptor-mediated endocytosis and thereby achieving increased exposure of the tumor to therapeutic drugs. A variety of drug delivery systems based on metallic nanoparticles, oxide nanoparticles, polymeric nanoparticles, carbon nanostructures, biodegradable nanoparticles, etc. have been developed for molecular imaging as well as drug delivery.121−125 For a given system, multiple factors determine the stability and fate of the delivery vehicle during storage and after administration, including size, rigidity, charge, solubility, and surface modifications of the nanoparticles. Therefore, the choice of a nanoparticle based delivery system is guided by the biodistribution, types of drugs that can be delivered using that system, and the specificity and pharmacokinetics of delivery. The different nanoparticle based systems which can be radiolabeled with suitable positron emitting radioisotopes for PET image-guided drug delivery are discussed in the following text.

**Metallic Nanoparticles.** Among the various metallic nanoparticles reported to date, gold nanoparticles are most widely used for biomedical applications, including drug delivery and novel diagnostic and therapeutic approaches, due to their biocompatibility, small size, ease of characterization, and rich surface chemistry.126−129 The utilization of 18F-labeled gold nanoparticles for PET imaging was first reported by Guerrero et al.130 in which the gold nanoparticles of ∼12 nm were synthesized by citrate reduction of HAuCl₄. The nanoparticles were functionalized with two different peptides, CK and CLPFFD, and 18F-SFB was covalently bound to the nanoparticle conjugate. After intravenous administration of the radiolabeled nanoparticles in normal rats, in vivo PET imaging showed highest uptake of the radioactivity in the bladder. The lungs, liver, and spleen were the organs with the next highest levels of radioactivity, followed by the intestine, kidneys, and blood. The pancreas and brain, however, accumulated very low concentrations of radiolabeled nanoparticles. Clearance of the nanoparticles from the biological system took place by both renal and biliary excretions.

Gold nanorods with suitable aspect ratios can absorb and strongly scatter light in the near-infrared region, which can be used for enhanced optical imaging and photothermal cancer therapy.131 The development of a multifunctional gold nanorod-based nanoplatform for targeted anticancer drug delivery and PET imaging of tumors was reported by Xiao et al.132 The bare gold nanorods had a length and diameter of approximately 45 and 10 nm, respectively. An anticancer drug (DOX) and tumor targeting agent (cRGD) were conjugated to the PEGylated gold nanorods. Also, NOTA was attached onto the distal ends of the PEG arms for complexation with ⁶⁴Cu. Based on flow cytometry analysis, cRGD-conjugated gold nanorods exhibited a higher cellular uptake and cytotoxicity than nontargeted ones in vitro. However, in vivo PET imaging and biodistribution studies showed that targeted and nontargeted gold nanorods had similar distribution pattern especially in the tumor (Figure 4A). Despite this limitation, this initial attempt provided a suitable nanoplatform for possible integration of multifunctionality including molecular targeting, chemotherapy, and photothermal therapy, as well as multimodality imaging, which can potentially lead to improved therapeutic efficacy and cancer monitoring. To achieve a similar goal, Xie et al. reported the preparation of ⁶⁴Cu-labeled gold nanoshells conjugated with cRGD and studied the in vivo biodistribution and tumor specificity using PET.133 The nanoshell used in this study was composed of a silica core (∼120 nm in diameter) and a gold shell (8−10 nm) to absorb light at near-infrared wavelengths. In vivo PET imaging suggested that tumor targeting was improved by conjugation of gold nanoshells to cRGD, which was advantageous over the previous study by Xiao et al.132 Both targeted and nontargeted gold nanoshells were cleared from the circulation by the liver and spleen. In the subablative thermal therapy study, enhanced biological effectiveness of targeted gold nanoshell was shown by the higher degree of tumor necrosis compared with nontargeted nanoshell. The promising results obtained from this study might lead to advancement of gold nanoshells as theranostic platforms for effective cancer diagnosis and therapy.

**Oxide Nanoparticles.** Another category of promising nanoplatforms which has drawn substantial interest recently is the oxide nanoparticles (such as mesoporous silica and iron oxide nanoparticles) due to their nontoxic nature, easily modifiable surface, and good biocompatibility.134,135 The utilization of core−shell silica nanoparticles as targeted PET/ optical multimodal imaging probes was reported by Benezra et al.136 Near-infrared fluorescent, Cy5 dye-encapsulated, core−shell silica-based nanoparticles were prepared and coated with PEG as per the method reported by Burns et al.137 The nanoparticles was conjugated with cRGD and radiolabeled with ¹²⁴I through a tyrosine linker. In vitro cell binding assays demonstrated the specificity of the nanoplatform toward integrin αvβ₃ expression. In vivo PET/optical imaging and biodistribution studies showed a tumor uptake of ∼1.5% ID/g at 4 h post injection, with high tumor to background ratio and rapid renal clearance. Owing to their favorable characteristics, such as bulk renal clearance, favorable targeting kinetics, lack of acute toxicity, superior photophysical features, and multimodal (PET/optical) imaging capabilities, such nanoparticles have received the United States Food and Drug Administration (US FDA)-investigational new drug approval for a first-in-human clinical trial.138

The synthesis of ultrasmall, monodisperse silica nanoconjugates for targeted dual-modal imaging of lymph nodes with metastatic tumors was reported by Tang et al.139 The nanoparticles were functionalized with an aptamer derivative having high binding affinity for nucleolin, a protein that is overexpressed in the cytoplasm and on the plasma membrane of several cancer cells. Also, a near-infrared (NIR) dye and DOTA were conjugated on the surface of the functionalized nanoparticle. The aptamer functionalized silica nanoconjugate was radiolabeled with ⁶⁴Cu, and in vivo PET/optical imaging studies showed markedly enhanced uptake of the radiolabeled agent in lymph nodes with metastatic tumors in a murine breast tumor model. In a similar study, Kim et al. reported the development of a core−shell silica nanoprobe for multimodal
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PET (optical/MRI) imaging of the sentinel lymph node.\textsuperscript{140} Magnetic silica nanoparticles with cobalt ferrite core and silica shell were synthesized which encapsulated NIR dye on the silica shell. The surface of the nanoparticle was modified with amino group and PEG for conjugation with NOTA, which was used for chelating \textsuperscript{68}Ga. The triple modality nanoprobe could be successfully utilized to visualize the sentinel lymph node in mice. Thus, these multimodal silica nanostructures hold great potential for improving the accuracy of clinical tumor staging by serving as probes for efficient noninvasive targeted imaging of metastatic lymph nodes. Different chemotherapeutic drugs can also be attached on the surface of the functionalized nanconjunctures for prevention of metastases.

In an interesting study, Di Pascua et al. utilized commercially available mesoporous silica nanoparticles as a carrier material for the therapeutic radioisotope \textsuperscript{166}Ho ($t_{1/2} = 26.8$ h, \textit{E}_{\text{fiss}} = 1.84 MeV).\textsuperscript{141} A lipophilic acetylacetone complex of \textsuperscript{165}Ho was incorporated in mesoporous silica nanoparticles (80–100 nm in diameter), which were subsequently irradiated in a neutron flux to produce particles containing \textsuperscript{165}Ho by ($n$,\gamma) reaction. These radioactive nanoparticles were utilized to deliver effective therapeutic doses for treating ovarian cancer metastases after intraperitoneal delivery in SKOV-3 ovarian tumor-bearing mice. \textit{In vivo} SPECT imaging demonstrated that most of the \textsuperscript{166}Ho-containing mesoporous silica nanoparticles administered to ovarian tumor-bearing mice were retained in the peritoneal cavity and selectively accumulated in the tumors (33% ID/g after 24 h). Radiotherapeutic efficacy was monitored using PET/CT using \textsuperscript{18}F-FDG which showed a decrease in tumor volume, which correlated with a marked increase in survival after treatment with $\sim$4 MBq of the radioactive nanoparticles. Though the authors could not explain the reason for the high uptake of mesoporous silica nanoparticles in ovarian tumor, this strategy might find utility in incorporation with other therapeutical radionuclides in nanostructured materials for treatment of various types of cancer. In another approach, Chen et al. reported the development of biocompatible functionalized mesoporous silica nanoparticles for actively targeted \textit{PET} imaging and chemotherapeutic drug delivery.\textsuperscript{142} Mesoporous silica nanoparticles were surface functionalized with thiol groups, PEGylated, conjugated with NOTA chelator and TRC105 antibody (specific for CD105/endoglin), and radio labeled with \textsuperscript{64}Cu. \textit{In vivo} PET imaging and biodistribution studies in 4T1 breast tumor bearing mice showed high tumor uptake ($\sim$6% ID/g) at 5 h post injection (Figure 4B). The tumor uptake of radio labeled nanoparticles not conjugated with TRC105 was much lower than the tumor uptake observed with TRC105 conjugated nanoparticles, indicating that active targeting was responsible for the enhanced tumor uptake. The authors also demonstrated the feasibility of enhanced tumor targeted drug delivery \textit{in vivo} using TRC105 conjugated mesoporous silica loaded with an anticancer drug, DOX. The encouraging results obtained in this study hold promise for future image-guided drug delivery and targeted cancer therapy using this class of nanomaterials.

Recently, iron oxide nanoparticles have been actively investigated as nanoplatforms for multimodal molecular imaging.\textsuperscript{145–145} The conventional drug loading approach by covalent linkage on such nanoparticles is inefficient and suboptimal for drug release.\textsuperscript{146} In order to circumvent this limitation, Xie et al. synthesized iron oxide nanoparticles, modified their surface using dopamine, and encapsulated them into human serum albumin matrices, which are clinically utilized as drug carriers.\textsuperscript{147} The human serum albumin coated iron oxide nanoparticles were dually labeled with \textsuperscript{64}Cu-DOTA and Cy5.5 dye, and tested in a subcutaneous U87MG xenograft mouse model. \textit{In vivo} PET/optical/MR imaging showed a high tumor uptake ($\sim$5% ID/g at 4 h post injection) with high tumor to background ratio. An inhomogeneous particle distribution pattern was observed with MRI, but PET and optical imaging showed homogeneous intensities at the tumor area. The human serum albumin coated nanoparticles manifested a prolonged circulation half-life. Adopting this strategy, small drug molecules can be coloaded with iron oxide nanoparticles into human serum albumin to yield theranostic agents. Recently, Chen et al. reported a chelator free approach for preparation of radiolesnic labeled iron oxide nanoparticles.\textsuperscript{148} The radiolabeled nanoparticle was used as PET/MRI agent for dual-modality imaging \textit{in vivo} and lymph node mapping. This strategy can be extended for radio labeling iron oxide nanoparticles with \textsuperscript{77}As for radiotherapeutic applications.\textsuperscript{149} In another study, Yang et al. reported the synthesis of cRGD-functionalized, DOX-conjugated, and \textsuperscript{64}Cu-labeled iron oxide nanoparticles for targeted anticancer drug delivery and PET/MR imaging.\textsuperscript{150} \textit{In vivo} PET imaging and biodistribution studies in U87 tumor bearing mice showed that cRGD-conjugated iron oxide nanocarriers showed a much higher level of tumor accumulation ($\sim$5% ID/g) than cRGD-free ones ($\sim$2% ID/g). Also, cRGD-conjugated nanocarriers induced a significant amount of cytotoxicity in the U87MG tumor cells, suggesting that DOX was released from the iron oxide nanocarrier and entered the cell nucleus. Thus, the potential of iron oxide nanoparticles for combined tumor-targeting drug delivery as well as multimodal imaging could be amply demonstrated.

**Polymeric Nanoparticles.** In recent times, there has been widespread interest in the use of biocompatible and biodegradable polymer nanoparticles for drug delivery.\textsuperscript{151} Bartlett et al. reported the synthesis of nanoparticles using cyclodextrin-containing polycations and siRNA sequence targeting luciferase mRNA.\textsuperscript{152} A bifunctional chelator, DOTA, was conjugated to the S’ end of siRNA and used for labeling with \textsuperscript{64}Cu. A dual-modality (PET/optical) imaging approach was used to investigate the biodistribution and functional activity of siRNA delivered by the nanoparticles. \textit{In vivo} PET/CT imaging in mice bearing luciferase-expressing Neuro2A tumors was used to analyze the biodistribution and tumor localization of the siRNA nanoparticles. Also, bioluminescent imaging was used before and after PET imaging to enable correlation of functional efficacy with biodistribution data. It was observed that both nontargeted and transferrin-targeted siRNA nanoparticles exhibited similar biodistribution and tumor localization. However, transferrin-targeted siRNA nanoparticles could reduce tumor luciferase activity by $\sim$50% relative to nontargeted siRNA nanoparticles, 1 d after injection. Compartmental modeling was used to demonstrate that the primary advantage of targeted nanoparticles was associated with processes involved in cellular uptake in tumor cells rather than overall tumor localization. The authors inferred that optimization of internalization might be the key factor for effective targeted therapy using this class of nanoparticles.

The utilization of PET to quantify the uptake of intercellular adhesion molecule 1 (ICAM-1) targeted, \textsuperscript{64}Cu-labeled polymeric nanoparticles by the pulmonary endothelium was reported by Rossin et al.\textsuperscript{153} \textit{In vivo} PET imaging and
biodistribution studies showed a 3- to 4-fold higher uptake in the lungs of mice injected with ICAM-targeted nanoparticles compared to that of the control group (Figure 4C). The lung uptake could be further enhanced by pretreating the mice with lipopolysaccharides probably due to ICAM-1 upregulation. However, a considerable release of small 64Cu-radiometabolites from the nanoparticles beginning as early as 1 h after injection was observed, suggesting poor in vivo stability of the radiolabeled conjugate. An improved strategy where the radiolabeled nanoparticle remained stable in vivo was reported by Simone et al.154 The authors developed a polymeric nanoparticle using a poly(4-vinylphenol) polymer backbone which could directly be radiolabeled with 124I. The polymeric nanoparticles were coated with monoclonal antibodies targeting endothelial determinants. The radiolabeled nanoparticles were used for imaging the pulmonary vasculature and also for tracking the nanoparticle pharmacokinetics. This approach might find utility in image-guided delivery of therapeutics to the pulmonary endothelium in patients with acute and chronic respiratory diseases.

The synthesis and utilization of poly(N-vinylpyrrolidone)-b-poly(ε-caprolactone) nanoparticles (∼100 nm diameter) for drug delivery was reported by Zhu et al.155 The nanoparticles were conjugated with a near-infrared fluorescent dye, NIR-797, for in vivo optical imaging. An anticancer drug, paclitaxel (PTX), was loaded in the polymeric nanoparticles with high drug loading content (>25%) and encapsulation efficiency (>85%). The antitumor effect of PTX-loaded nanoparticles was evaluated, both in vitro on three different cancer cell lines and in vivo on a hepatic H22 tumor bearing mouse model using optical imaging. The antitumor effects of the PTX-loaded nanoparticles were further visualized using 18F-FDG PET scans. By combining the tumor volumes and survival rate measurements, it could be confirmed that PTX-loaded nanoparticles exhibited superior in vivo antitumor effect than Taxol (commercially available formulation of paclitaxel). In a similar study, Liu et al. reported the synthesis of poly(ethylene glycol)-poly(caprolactone) nanoparticles (∼70 nm diameter) and evaluated their efficacy for drug delivery.156 In the previous case, the polymeric nanoparticles were conjugated with NIR-797 dye for investigating the biodistribution of the drug-loaded nanoparticles using in vivo optical imaging. An anticancer drug, docetaxel (DOC), could be encapsulated into the polymeric nanoparticles with a high drug loading content (∼20%) and encapsulation efficiency (>80%). In vitro cytotoxicity test showed that DOC-loaded nanoparticles inhibited the murine hepatic carcinoma cell line H22 in a dose-dependent manner, which was similar to Taxotere, the commercialized formulation of docetaxel. However, in vivo tumor evaluation using optical imaging and 18F-FDG PET scans demonstrated the superiority of DOC-loaded polymeric nanoparticles over Taxotere. Therefore, it could be envisaged that these highly efficient and biodegradable nanoparticles might find clinical utility in PET image-guided anticancer drug delivery in the near future.

In a pioneering study, Zhou et al. reported the synthesis of poly(lactide-co-glycolide) nanoparticles (∼70 nm diameter), which could be utilized as brain penetrating nanocarriers for the treatment of glioblastoma.157 In order to illustrate the translational potential of brain-penetrating nanoparticles, the authors conducted a screen of ∼2,000 compounds that were previously approved by US FDA to inhibit patient-derived brain cancer stem cells and encapsulated the best agent (dithiazanine iodide) into the nanocarrier. The nanoparticles were radio-labeled with 18F using streptavidin/biotin chemistry. The radiolabeled brain-penetrating nanocarriers were administered by convection-enhanced delivery in rats bearing brain cancer stem cell derived xenografts. In vivo PET imaging demonstrated accumulation of the nanoparticles in the brain. Also, a significantly increased survival in rats bearing brain cancer xenografts was observed, which demonstrated the potential of such brain-penetrating nanoparticles for targeted image-guided drug delivery for treatment of brain tumors.

With a different strategy, Chen et al. employed anionic poly(ε-glutamic acid) as a carrier to covalently link with camptothecin (an anticancer drug), enabling encapsulation into supramolecular nanoparticle vectors.158 Approximately five camptothecin molecules were conjugated to each polymer chain by ester bond formation, which could be degraded via esterase-mediated hydrolysis to allow controlled release of camptothecin under physiological conditions. The supramolecular nanoparticle was further conjugated with DOTA for 64Cu labeling. The authors synthesized nanoparticles of two different sizes (37 and 104 nm), both of which were radiolabeled with 64Cu and administered in mice bearing Lewis lung carcinoma xenografts. In vivo PET imaging and biodistribution studies revealed that the smaller sized (37 nm) nanoparticles exhibited higher tumor accumulation due to the EPR effect. The superior in vivo antitumor efficacy of the 37 nm supramolecular nanoparticles was further validated by tumor reduction/inhibition studies. Despite the encouraging results, the tumor uptake of the supramolecular nanoparticles was not impressive, which might be a deterrent for their use as potential drug delivery vehicles. However, the tumor uptake might be improved on conjugation with suitable targeting ligands.

Homopolymers or copolymers have long been explored as potential carriers in targeted drug delivery.159 The synthesis and characterization of PEGylated star-shaped copolymer nanoparticles (25–70 nm size) containing core–shell morphology for in vivo PET imaging was reported by Fukukawa et al.160 These copolymers possessed a hydrophilic inner shell bearing reactive functional groups, and a central hydrophobic core. DOTA was conjugated to the functional groups in the inner shell for 64Cu labeling. In vivo PET imaging and biodistribution studies in normal rats showed that copolymers with increasing PEG shell thickness showed increased blood circulation and low accumulation in excretory organs. This preliminary study suggested the potential of such systems as for in vivo tumor imaging and targeted drug delivery. The synthesis of N-(2-hydroxypropyl)methacrylamide (HPMA) copolymers for PET imaging and image-guided chemotherapy of prostate cancer was reported by Yuan et al.161 The HPMA copolymer was conjugated with DOTA for 64Cu labeling and also with cRGD peptide for targeting αvβ3 integrin in tumor neovasculature. The tumor localization of the radiolabeled copolymer was visualized by PET in a mouse model bearing human prostate cancer xenografts. A time-dependent increase in radioactivity uptake in tumor-bearing mice injected with the HPMA-cRGD-DOTA-64Cu copolymer was observed, but this phenomenon was not seen in mice injected with control HPMA-DOTA-64Cu copolymer. However, the tumor uptake observed for the targeted copolymer (2.75% ID/g) at 3 h post injection was slightly higher than what was observed with the non-targeted copolymer (1.29% ID/g), suggesting that, along with active targeting, passive EPR effect also plays a partial role in tumor localization of the radiolabeled copolymer. The findings from these studies might set the stage for further optimization and
evaluation of the copolymer constructs for image-guided drug delivery in various tumor models.

**Carbon-Based Nanomaterials.** Owing to their unique physical and chemical properties, the use of functionalized carbon-based nanomaterials is gaining popularity in many areas of biomedical research, including molecular imaging and drug delivery.\(^{162}\)−\(^{164}\) PET imaging using PEGylated single walled carbon nanotubes (1−5 nm diameter, 100−300 nm length) conjugated with RGD peptides was reported by Liu et al.\(^{165}\) DOTA was attached to the termini of the PEG chains and used to conjugate \(^{64}\)Cu. In vivo PET imaging and biodistribution studies showed that PEG\(_{5000}\) modified single walled carbon nanotubes conjugated with cRGD exhibited a high tumor uptake of 10−15% ID/g, with high target to nontarget ratio (Figure 4D). High tumor uptake of the radiolabeled single walled carbon nanotubes was observed over long periods (>24 h). In another study, McDevitt et al. studied the biodistribution pattern of \(^{89}\)Y labeled carbon nanotubes (~1 nm diameter, ~50 nm length) without PEG modification in normal mice.\(^{166}\) The radiolabeled agent cleared from the blood within 3 h and distributed predominantly to the kidneys, liver, spleen, and bone. Suitable PEG modification of carbon nanotubes is of paramount importance in order to reduce reticuloendothelial system uptake and prolong blood circulation time of the single walled nanotubes for finding utility in drug delivery approaches.

Graphene is another structurally robust, yet highly flexible, nanoplateform with potential for use as a drug delivery vehicle.\(^{162}\) Hong et al. reported the synthesis of covalently functionalized nanographene oxide sheets (10−50 nm), which were PEGylated and conjugated with anti-CD105 monoclonal antibody (TRC105) for imaging tumor angiogenesis.\(^{167}\) The PEGylated graphene oxide was also conjugated with NOTA for \(^{64}\)Cu labeling. In vitro studies using human umbilical vein endothelial cells (HUVECs, high CD105 expression) demonstrated strong and specific CD105-binding by the TRC105 conjugated nanographene. Also, in vivo PET imaging and biodistribution studies in 4T1 tumor bearing mice showed high tumor uptake (~6% ID/g) within 0.5 h post injection, which remained fairly stable over time. These studies were further validated by ex vivo histological analyses, and vasculature specific targeting with little extravasation of TRC105

Figure 5. PET image-guided tumor targeting using conventional radiopharmaceuticals. (A) \(^{68}\)Ga-DOTATATE PET/CT. (i) Images showing \(^{68}\)Ga-DOTATATE avid lesions in T4 vertebral body and 3 metastases in liver (arrow). Physiologic uptake is seen in pituitary, kidneys, bladder, stomach wall, liver, and spleen. (ii) Images from repeated \(^{68}\)Ga-DOTATATE PET/CT 1 y later after 3 administrations of \(^{177}\)Lu-DOTATATE, showing metabolic partial response with reduction in SUV\(_{\text{max}}\) of lesion in T4 and liver and no new lesions. Adapted with permission from ref 190. Copyright 2011 Society of Nuclear Medicine and Molecular Imaging. (B) \(^{89}\)Zr-trastuzumab PET/CT. (i) A patient with liver and bone metastases, and (ii and iii) two patients with multiple bone metastases. A number of lesions have been specifically indicated by arrows. Adapted with permission from ref 198. Copyright 2010 Nature Publishing Group.
Conventional Radiopharmaceutical-Based Delivery Approach. The drug delivery systems discussed thus far were mostly limited to chemotherapy. The concept of “theranostics” has also played a vital role in radiation-based therapies, especially, using targeted radiopharmaceuticals. In this approach, a radiation dose is specifically administered to the cancerous lesions using peptides, proteins, or antibodies radiolabeled with suitable therapeutic radionuclides such as $^{90}$Y, $^{131}$I, or $^{177}$Lu. The same targeting ligands can also be conveniently radiolabeled with suitable positron emitters such as $^{18}$F, $^{68}$Ga, $^{64}$Cu, or $^{89}$Zr, thereby providing exciting opportunities to guide such therapies using PET.

This approach plays a dominant role in diagnosis of the disease in its early stage, validation of the targeting strategy, and development of novel therapeutic radiopharmaceuticals. Thus, it facilitates better, faster, and cost-effective decision making, helping to eliminate failures in the targeted radiotherapy pathway and advance only with the promising candidates to receive such therapies. Despite the availability of a wide variety of PET radiopharmaceuticals, $^{18}$F-FDG is still the most widely used radiotracer in cancer management, and the pivotal role of $^{18}$F-FDG-PET/CT in modern nuclear medicine needs hardly to be reiterated. Excellent review articles have appeared in recent times which have summarized the clinical diagnosis and therapeutic response evaluation using $^{18}$F-FDG and other $^{18}$F-based radiotracers, and hence these are not discussed here further.

Another PET radioisotope which is gaining significant clinical attention in recent times is $^{68}$Ga ($T_{1/2} = 68$ min). The convenient availability of this radioisotope from $^{68}$Ge/$^{68}$Ga generators without the dependence on onsite cyclotrons makes it economical to use for a wide variety of PET scans. In particular, from the perspective of image-guided therapy, the radiolabeled peptides targeting somatostatin receptors, overexpressed in the majority of neuroendocrine malignancies, have a great potential for both imaging and therapy of tumors where other therapies fail. The development of $^{68}$Ga-labeled somatostatin analogues such as DOTANOC (DOTA-1-Nal$^3$-octreotide), DOTATOC (DOTA-Ph$^1$-Tyr$^3$-octreotide), or DOTATATE (DOTA-Ph$^1$-Tyr$^3$-Thr$^8$-octreotide) for PET/CT imaging has significantly improved the diagnosis of neuroendocrine tumors.

In a typical example, Gains et al. investigated the efficacy of PET/CT using $^{68}$Ga-DOTATATE to select children with primary refractory or relapsed high-risk neuroblastoma for treatment with $^{177}$Lu-DOTATATE and also evaluated whether this is a viable therapeutic option for those children. Imaging with $^{68}$Ga-DOTATATE could indicate the expression of somatostatin receptors and was of paramount importance for guiding radiotherapy (Figure 5A). Post-therapy of $^{177}$Lu-DOTATATE, $^{68}$Ga-DOTATATE scans was used to assess the response (Figure 5A), and a positive therapeutic outcome with a regression in some lesions and an apparent block of the metastatic activity could be observed in many cases. In a similar study, Budiawan et al. investigated the application and role of $^{68}$Ga-DOTATATE PET/CT imaging in non-radioiodine-avid refractory thyroid cancer patients who have undergone peptide receptor radionuclide therapy (PRRT) with $^{90}$Y/$^{177}$Lu-DOTATATE. The authors concluded that PRRT guided by PET imaging is an effective therapeutic option with minimal toxicity, good response rate, and excellent survival benefits. In a different study, Wu et al. reported the use of $^{68}$Ga-labeled SIR spheres as PET imaging surrogate for distribution assessment and radiation dose estimation of $^{90}$Y-SIR-Spheres, which are currently used in the treatment of solid liver tumors. The authors observed that $^{68}$Ga-SIR-Spheres had good in vivo stability and localization of the radiolabeled microparticles in the liver could be observed using PET imaging. Similar results were obtained by Avila-Rodriguez et al. on using $^{64}$Cu or $^{86}$Y labeled SIR spheres.

Another example, close to clinical translation, is represented by PET imaging of human epidermal growth factor receptor 2 (HER2) expressions. The overexpression of the HER2 is observed in ~15% of breast cancers and associated with poor prognosis in terms of overall survival. Generally, $^{90}$Y or $^{131}$I labeled trastuzumab is used for targeted radiotherapy of primary or metastatic breast cancer, in experimental, preclinical, or clinical settings. PET imaging of HER2-positive lesions in patients with metastatic breast cancer was reported by Dijikers et al. The antibody trastuzumab was radiolabeled with $^{89}$Zr, a PET imaging surrogate for $^{90}$Y. Administration of $^{89}$Zr-trastuzumab at appropriate doses allowed visualization and quantification of uptake in HER2-positive lesions in patients with metastatic breast cancer by PET (Figure 5B). The authors concluded that PET imaging of HER2 expression would aid in improving diagnosis, staging of the disease, guiding trastuzumab therapy, and monitoring the therapeutic response. A large number of other PET radiopharmaceuticals for molecular imaging and personalized cancer management have also been reported, details of which can be found in recent review articles.

We would apologize to those whose work could not be presented here, mainly due to the vastness of the field and availability of enormous literature which could not be summarized in a single review.

### SUMMARY AND FUTURE PERSPECTIVES

In the last several decades, major milestones have been reached in every area of cancer care and a variety of anticancer drugs have been developed which are now commercially available worldwide. Though most of these anticancer agents have the potential to be effective at sufficiently high doses, they are often associated with severe systemic side effects that cannot be tolerated by patients who are already weak due to effects of cancer. In a majority of the cases, the success of cancer therapy typically hinges upon circumventing the dose-limited toxicity while administering such drugs. Additionally, many conventional therapeutic agents often fail due to their limited ability to reach the target tissue and their poor selectivity against...
cancerous lesions. Ideally a drug should possess perfect specificity to cancerous cells and have no effect on the rest of the body. To achieve these objectives, a variety of drug delivery systems have been engineered for the targeted delivery and controlled release of therapeutic agents to specifically kill the cancerous cells. The full potential of the drug delivery systems extends beyond treatment, and several image-guided approaches have now been implemented in areas ranging from new therapeutic target discovery to effectively monitoring tumor pharmacokinetics and drug distribution to modulation of drug release at the target site. In particular, PET image-guided drug delivery provides a means for treating a variety of diseases with minimal systemic involvement while concurrently monitoring therapeutic efficacy. This minimally invasive approach provides a comprehensive answer to many challenges with conventional therapeutic approaches and is expected to lead to a paradigm shift in cancer patient care.

The major advances in targeted drug delivery have been attributed to the recent progress in nanotechnology that has resulted in the development of nanosized drug delivery platforms having distinct advantages in cancer therapy. The versatile nanoplatforms provide opportunities for multifunctionalization so that a single platform can be used to detect and treat tumors, monitor treatment response, and thus guide therapeutic regimes. The nanoformulations can be functionalyzed to minimize clearance by the immune system and prolong circulation times, and also for attachment of suitable vectors (peptides, proteins, antibodies, etc.) targeting specific receptors, thereby enhancing tumor uptake through EPR effect as well as receptor-mediated endocytosis. The nanomaterial based delivery systems also result in improved treatment effects by protecting entrapped drugs from degradation during their delivery. The multifunctionality of the nanoplatforms offers possibility for multimodality molecular imaging. When two or more molecular imaging techniques are used in conjunction, they would provide synergistic information when compared with any single imaging modality. Additionally, multiple therapeutic agents such as chemotherapy, antiangiogenic, or gene therapy agents can be simultaneously delivered by nanocarriers to tumor sites to enhance the effectiveness of therapy.

Despite these advantages, it must be admitted that the field of PET image-guided drug delivery is still in its infancy and more systematic studies to understand the mechanisms for targeting and drug delivery would be required in order to translate these novel discoveries into clinical impact. The potential challenges to clinical translation of the drug delivery systems are in vivo characterization of the drug carriers, preclinical validation of targeting and delivery, studies of biodistribution, pharmacokinetics, pharmacodynamics, and toxicity, and scale-up manufacturing of delivery systems. Moreover, the performance of the drug delivery carriers depends on several limiting factors which include the synthesis method adopted, their size and shape, internal structure of the drug carrier, drug loading methodology, surface functionalization and conjugation strategy adopted for attaching targeting ligands to the carrier platforms, etc., thereby making relative evaluation of different drug delivery platforms difficult. The issues related to toxicity of nanosized drug delivery platforms can be addressed by the use of biocompatible and biodegradable polymeric nanoparticles for drug delivery, which have received considerable attention in the recent times. Also, stimulus-responsive polymer nanomaterials can be synthesized which mimic the behavior of biological molecules, where external stimuli or changes in local environment can trigger a change in property to regulate drug release. Such “smart” nanoparticle systems when coupled with suitable targeting ligands can probably best minimize off-target effects and maximize programmability, thereby offering the possibility of radically changing the practice of drug delivery.

While numerous obstacles face all new technologies, materializing the opportunities presented by PET image-guided drug delivery requires addressing the significant interdisciplinary challenges and biological barriers. Besides these, several other complex factors, such as considerable regulatory hurdles, limited potential market, lobbying by the manufacturers of established anticancer drugs, lack of reimbursement strategies by the insurance agencies for such novel strategies, etc., might impede the “bench to bedside” translation of this promising approach. The concerted efforts of all stakeholders, which include scientists from academia as well as industries, progressive and technology savvy physicians, radiologists, surgeons, program advisory boards, regulatory authorities, and grant review panels, would be required, both to create enthusiasm for developing these new concepts and also to prevent adverse messaging based on myths, conjectures, hyperbole, and bias. This in turn would provide impetus to further research which might aid in clinical translation of PET image-guided drug delivery approaches and thus achieve the ultimate goal of “personalized medicine” in the near future.

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Notes
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