Copper-64 radiopharmaceuticals for PET imaging of cancer: Advances in preclinical and clinical research

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Carolyn J. Anderson1–3 and Riccardo Ferdani1

Summation

Copper-64 (T1/2 = 12.7 hours; β+, 0.653 MeV [17.8 %]; β−, 0.579 MeV [38.4 %]) has decay characteristics that allow for positron emission tomography (PET) imaging and targeted radiotherapy of cancer. The well-established coordination chemistry of copper allows for its reaction with a wide variety of chelator systems that can potentially be linked to peptides and other biologically relevant small molecules, antibodies, proteins, and nanoparticles. The 12.7-hours half-life of 64Cu provides the flexibility to image both smaller molecules and larger, slower clearing proteins and nanoparticles. In a practical sense, the radionuclide or the 64Cu-radiopharmaceuticals can be easily shipped for PET imaging studies at sites remote to the production facility. Due to the versatility of 64Cu, there has been an abundance of novel research in this area over the past 20 years, primarily in the area of PET imaging, but also for the targeted radiotherapy of cancer. The biologic activity of the hypoxia imaging agent, 60Cu-ATSM, has been described in great detail in animal models and in clinical PET studies. An investigational new drug application for 64Cu-ATSM was recently approved by the U.S. Food and Drug Administration (FDA) in the United States, paving the way for a multicenter trial to validate the utility of this agent, with the hopeful result being FDA approval for routine clinical use. This article discusses state-of-the-art cancer imaging with 64Cu radiopharmaceuticals, including 64Cu-labeled peptides for tumor-receptor targeting, 64Cu-labeled monoclonal antibodies for targeting tumor antigens, and 64Cu-labeled nanoparticles for cancer targeting. The emphasis of this article will be on the new scientific discoveries involving 64Cu radiopharmaceuticals, as well as the translation of these into human studies.

Key words: antibody, bone metastases, cancer, PET, molecular imaging

Introduction

A significant research effort has been devoted to the copper radionuclides because they offer a varying range of half-lives and positron energies (Table 1). In addition, the well-established coordination chemistry of copper allows for its reaction with a wide variety of chelator systems that can be linked to antibodies, proteins, peptides, and other biologically relevant small molecules. This update will focus on 64Cu radiopharmaceuticals for positron emission tomography (PET) imaging applications. The longer half-life allows 64Cu to be produced at regional or national cyclotron facilities and distributed to local nuclear medicine departments with the loss of approximately one half-life. In addition, the longer half-life is compatible with the time scales required for the optimal biodistribution of slower clearing agents, such as monoclonal antibodies (mAbs), nanoparticles, and higher molecular weight polypeptides requiring longer imaging times.

Production of Copper Radionuclides

The production of no-carrier-added 64Cu via the 64Ni(p,n) 64Cu reaction on a biomedical cyclotron was proposed by Szelecsenyi et al. In this study, small irradiations were performed demonstrating the feasibility of 64Cu production

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activity samples of $^{64}$Cu-$\text{CuCl}_2$ were obtained (95–310 mCi purification by using an ion-exchange column, high-specific-

- donating ligands. Copper (II) is a d$^9$ metal of bor-

- to the thermodynamic

<table>
<thead>
<tr>
<th>Isotope</th>
<th>$t_{1/2}$</th>
<th>$\beta^-$ MeV (%)</th>
<th>$\beta^+$ MeV (%)</th>
<th>EC (%)</th>
<th>$\gamma$ MeV (%)</th>
</tr>
</thead>
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<td>$^{60}$Cu</td>
<td>23.4 minutes</td>
<td>—</td>
<td>2.00 (69)</td>
<td>7.0</td>
<td>0.511 (186)</td>
</tr>
<tr>
<td>$^{61}$Cu</td>
<td>3.32 hours</td>
<td>—</td>
<td>1.22 (60%)</td>
<td>40</td>
<td>0.511 (120)</td>
</tr>
<tr>
<td>$^{62}$Cu</td>
<td>9.76 minutes</td>
<td>—</td>
<td>2.91 (97%)</td>
<td>2</td>
<td>0.511 (194)</td>
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<tr>
<td>$^{64}$Cu</td>
<td>12.7 hours</td>
<td>0.573 (38.4)</td>
<td>0.655 (17.8%)</td>
<td>43.8</td>
<td>0.511 (35.6)</td>
</tr>
<tr>
<td>$^{67}$Cu</td>
<td>62.0 hours</td>
<td>0.395 (45)</td>
<td>—</td>
<td>—</td>
<td>0.184 (40)</td>
</tr>
</tbody>
</table>

Coordination Chemistry of Copper(II)

The aqueous-solution coordination chemistry of copper is limited to three oxidation states (I–III). Due to the lability of most Cu(I) complexes, they typically lack sufficient kinetic stability for radiopharmaceutical applications, while Cu(III) is relatively rare and difficult to attain without the use of strong $\pi$-donating ligands. Copper (II) is a d$^9$ metal of borderline softness, which favors amines, imines, and bidentate ligands, such as bipyridine to form square planar, distorted square planar, trigonal pyramidal, square pyramidal, as well as distorted octahedral geometries. Cu(II) is generally less labile toward ligand exchange and is the best candidate for incorporation into radiopharmaceuticals. Jahn-Teller distortions in six-coordinate Cu(II) complexes are often observed as an axial elongation or a tetragonal compression. Although Cu(II) is less labile than Cu(I) and $^{64}$Cu is a good radionuclide for PET imaging, the kinetic stability of Cu(II) complexes in vivo is very different from the thermodynamic stability in aqueous solution. Therefore, the development of Cu(II) complexes for radiopharmaceutical applications has been an active area of research.

Chelators based on cyclam and cyclen backbones

The most widely used chelators for attaching $^{64}$Cu to biological ligands are tetraazamacrocyclic ligands with pendant arms that utilize both the macrocyclic and chelate effects to enhance stability. By far, the most extensively used class of chelators for $^{64}$Cu has been the macrocyclic polyaminocarboxylates shown in Figure 1. Two of the most widely studied chelators are DOTA (1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid) and TETA (1,4,8,11-tetraazacyclotetradecane-1,4,8,11-tetraacetic acid). While DOTA has been used as a BFC (bifunctional chelator) for $^{64}$Cu, its ability to bind many different metal ions and its decreased stability, compared to TETA, makes it less than ideal. The tetraazamacrocyclic ligand, TETA, therefore, has been extensively used as a chelator for $^{64}$Cu, and successful derivatization of this ligand has allowed researchers to conjugate it to antibodies, proteins, and peptides.

Although $^{64}$Cu-TETA complexes are more stable than $^{64}$Cu-DOTA and $^{64}$Cu-labeled complexes of acyclic ligands, their instability in vivo has been well documented by our lab. Bass et al. demonstrated that when $^{64}$Cu-TETA-octreotide (OC) was injected into normal Sprague-Dawley rats, nearly 70% of the $^{64}$Cu from $^{64}$Cu-TETA-OC was transchelated to a 35-kDa species believed to be superoxide dismutase (SOD) in the liver 20 hour postinjection. These results are supported by the observations of Boswell et al.

Sarcophagine chelators

Another class of ligands that has gained attention as potential $^{64}$Cu chelators are the hexaazamacroyclic cage-type ligands, which are based upon the sepulchrate or sarcophagine cage motifs (Fig. 1) and whose syntheses were first described by Sargeson. Both cage systems are synthesized by reaction of the inert tris-ethylenediamine cobalt (III) complex with formaldehyde, followed by reaction with ammonia/formaldehyde or nitromethane/formaldehyde under basic conditions to generate the sepulchrate or sarcophagine (Sar) ligands, respectively. Smith et al. investigated a family of Sar
FIG. 1. Structures of macrocyclic chelators for complexing copper radionuclides.
derivatives with various functional groups at the apical sites, while the SarAr ligand was used to determine the $^{64}$Cu complexation rates from pH 4 to 9. From the data presented, complexation was 100% complete within several minutes at 25°C over the entire pH range. Biodistribution data was collected by using $^{64}$Cu-Sar, $^{64}$Cu-diamSar, and $^{64}$Cu-SarAr in Balb/c mice. All three complexes cleared from the blood rapidly, and uptake was low in bone, heart, stomach, spleen, muscle, lungs, and the gastrointestinal tract. Liver clearance was observed to be good over the 30-minute time course of this study, demonstrating that the $^{64}$Cu complexes are initially stable in vivo, but clearance of all three $^{64}$Cu complexes is much slower through the kidney. Activity levels increased in the case of the $^{64}$Cu-Sar complex, though this type of accumulation is not uncommon for positively charged complexes.

The cross-bridged tetraamine ligands

This class of chelators was first conceived of and synthesized by Weisman et al. in the 1990s and they were originally designed to complex metal cations, such as Li$^+$, Cu$^{2+}$, and Zn$^{2+}$, within their clamshell-like clefts. Numerous copper complexes of these and related ligands have since been prepared and studied by the Wong and Weisman labs as well as other research groups. The expected cis-fused coordination geometry of these chelators has been confirmed in all cases via the available structural data. The attachment of two carboxymethyl pendant arms to CB-cyclam to give CB-TE2A (4,11-bis(carboxymethyl)-1,4,8,11-tetraazacyclotetradecane) further ensures the complete envelopment of a six-coordinate Cu(II).

While the measurement of stability constants of Cu(II)-CB complexes have been limited by the proton-sponge nature of these chelators, available data for Cu(II)-CB-cyclam (log $K_f = 27.1$) revealed very similar values to nonbridged Cu(II)-cyclam (log $K_f = 27.2$) and related complexes. On the other hand, their kinetic inertness, especially in aqueous solution, has been shown to be truly exceptional. Proton-assisted decomplexation is one indicator of solution inertness. Under pseudo-first-order conditions of high-acid concentration (e.g., 5 M HCl), decomplexation half-lives can provide a comparative gauge. For example, Cu-CB-cyclam is almost 1 order of magnitude more inert than Cu (II)-cyclam in 5 M HCl at 90°C, while Cu(II)-CB-TE2A is 4 orders of magnitude more inert (T$_{1/2} = 154$ hour). Impressively, the latter complex resists acid decomplexation even better than the fully encapsulated sarcophagine complex, Cu(II)-diamSar (3,6,10,13,16,19-hexaaazabicyclo[6.6.6]hexadecane) (T$_{1/2} = 40$ hours). It was confirmed that both the cross-bridged cyclam backbone as well as the presence of two enveloping carboxymethyl arms are required for this unusual kinetic inertness.

Biologic stability of $^{64}$Cu-labeled cross-bridged complexes, including CB-cyclam, $^{64}$Cu-CB-TE2A, and CB-DO2A (10-bis (carboxymethyl)-1,4,7,10-tetraazabicyclo[5.5.2]octadecane), has been investigated. The biodistribution of these $^{64}$Cu complexes in female Sprague-Dawley rats were highly dependent upon the chelator. Based on the rapid clearance from the blood, liver, and kidney, $^{64}$Cu-CB-TE2A was thought to be the most stable. Follow-up metabolism studies of $^{64}$Cu-CB-TE2A and $^{64}$Cu-CB-DO2A, compared to $^{64}$Cu-DOTA and $^{64}$Cu-TETA, demonstrated the robust stability of $^{64}$Cu-CB-TE2A in vivo, with low amounts of transchelation to the liver and blood proteins.

In order to find chelators that complex Cu(II) with faster kinetics while retaining the high stability and the significant inertness observed with CB-TE2A, phosphonic-acid (-CH$_2$-PO$_2$H$_2$) donor groups were investigated as pendant arms. It has been shown previously that chelators with phosphonic-acid pendant arms have higher selectivity as well as increased thermodynamic and kinetic stability, compared to their acetic acid analogs. Cross-bridged 1,4,6,11-tetraazacyclotetradecane-1,8-bis(methanephosphonic acid) (CB-TE2P) and 4,8,11-tetraazacyclotetradecane-1-(methanephosphonic acid) 8-(methanecarboxylic acid) (CB-TE1A1P) were synthesized, radiolabeled with $^{64}$Cu, and their in vivo behavior was investigated. While CB-TE2P labeling with $^{64}$Cu was complete within 1 hour in buffer at higher temperatures, radiolabeling yields above 90% were observed even at 37°C. CB-TE1A1P had 100% radiolabeling yields at 37°C. Preliminary biodistribution studies showed that the biodistribution of $^{64}$Cu-CB-TE2P and $^{64}$Cu-CB-TE1A1P compared favorably to $^{64}$Cu-CB-TE2A.

Imaging Tumor Hypoxia with $^{64/60}$Cu-ATSM

There is one class of copper radiopharmaceuticals where the stability of the Cu(II) complex is not essential for successful targeting. Cu(II) thiosemicarbazones have been evaluated as blood-flow agents and for imaging tumor hypoxia. In this article, we discuss the most recent developments for imaging tumor hypoxia with this class of agents.

It was well established that hypoxia is an important determinant of the overall response of the tumor to conventional therapy. The presence of hypoxia can result in an increase in tumor aggressiveness, failure of local control, and activation of transcription factors that support cell survival and migration. The ability to locate and quantify the extent of hypoxia within solid tumors by using noninvasive nuclear imaging would facilitate early diagnosis and help clinicians select the most appropriate treatment for each individual patient.

In 1997, Fujibayashi et al. discovered that the neutral, lipophilic copper(II) complex of the N$_2$S$_2$ tetradentate ligand, diacetyl-2,3-bis(N$^3$-methyl-3-thiosemicarbazono), commonly referred to as Cu-ATSM, showed hypoxia-selective uptake in ex vivo ischemic, perfused, isolated rat-heart models. Cu-ATSM was later shown to be hypoxia selective in vitro and for tumor hypoxia. Recent experimental and computational work provided the first experimental evidence directly probing the reduction, reoxidation, and pH-mediated ligand dissociation reactions of Cu-ATSM and their relationship to hypoxia selectivity.

The thiosemicarbazones have been evaluated with the short-lived copper radionuclides. $^{60}$Cu and $^{68}$Cu (T$_{1/2} = 0.16$ hours, $\beta^+ = 98\%$, EC = 2%). Takahashi et al. reported
the first human studies of the uptake of $^{62}$Cu-ATSM in 10 patients: 4 normal patients and 6 with lung cancer. High tumor uptake was observed (uptake ratio, 3.00 ± 1.50) in all patients with lung cancer. Dehdashti et al. reported the first correlative studies comparing the uptake of $^{60}$Cu-ATSM ($T_1=0.16$ hour, $\beta^+=98\%$, EC = 2\%) with response to conventional therapies in patients with non-small-cell lung cancer (NSCLC) and cervical cancer. In the NSCLC study, response to therapy was evaluated by using $^{60}$Cu-ATSM tumor-to-muscle ($T/M$) uptake ratios. Imaging with $[^{18}$F]-FDG was also conducted as part of the routine clinical evaluation. Of the 14 patients studied, 8 responded to radiotherapy (5 showed a complete response with 3 partial responders) and 6 showed no response. The mean $^{60}$Cu-ATSM $T/M$ ratio of nonresponders (3.4 ± 0.8) was found to be much larger than uptake observed in responders (1.5 ± 0.4) [$p=0.002$]. However, no significant differences were observed in the standardized uptake values (SUVs) between the tumors of responders (3.5 ± 1.0) and nonresponders (2.8 ± 1.1) [$p=0.2$]. The threshold $T/M$ value of 3.0 was identified as an accurate cut-off value for distinguishing responders from nonresponders. In contrast to the results with $^{60}$Cu-ATSM, no significant differences were observed in either the mean $T/M$ ratios or SUVs for the uptake of $[^{18}$F]-FDG (2-Fluoro-2-deoxy-D-glucose) in responders (12.7 ± 10.4) and nonresponders (10.9 ± 4.1) [$p=0.7]$. In addition, no statistically significant correlation between $^{60}$Cu-ATSM and $[^{18}$F]-FDG uptake was observed.

Before radiolabeled Cu-ATSM could be used for routine clinical analysis, accurate dosimetry measurements were required. In 2005, Laforest et al. used the Medical Internal Radionuclide Dose (MIRD) approach to provided estimates of human absorbed doses from $^{60}/^{61}/^{62}/^{64}$Cu-ATSM by extrapolating data acquired from biodistribution data in rat models. Calculated organ doses for $^{61}$Cu, $^{62}$Cu, and $^{64}$Cu were extrapolated from the results obtained for $^{60}$Cu-ATSM dosimetry. The estimated human dose for safe injection into an adult was predicted to lie between 500 and 800 MBq.

Human doses using $^{64}$Cu-ATSM have also been estimated from biodistribution data in non-tumor-bearing hamsters.

Lewis et al. reported the first clinical comparison between the imaging characteristics of $^{60}$Cu-ATSM and $^{64}$Cu-ATSM (and $[^{18}$F]-FDG) in cancers of the uterine cervix conducted after Cu-ATSM was approved for study as an investigational new drug (IND 62,675) (Figure 2). The study concluded that tumor uptake of Cu-ATSM as measured in images recorded between 1 and 9 days was reproducible, irrespective of the radionuclide used. This important result showed that Cu-ATSM is a marker for chronic tumor hypoxia, as opposed to acute hypoxia. Pretherapy imaging has also confirmed previous results indicating that the PET imaging of Cu-ATSM provides clinically relevant information about tumor oxygenation and is predictive of the likelihood of disease-free survival post-treatment in patients with cervical cancer.

**Copper-64-Labeled Somatostatin Analogs for Targeting Neuroendocrine Tumors**

Somatostatin is a 14-amino-acid peptide that is involved in the regulation and release of a number of hormones, and somatostatin receptors (SSRs) are present in many different normal organ systems, such as the central nervous system (CNS), the gastrointestinal tract, and the exocrine and endocrine pancreas. Several human tumors of the neuroendocrine system, CNS, breast, and lung are SSR positive, making it a viable disease target. Further, the presence of SSRs in a tumor is predictive of a good therapeutic response. An 8-amino-acid analog of somatostatin, octreotide (OC) has a longer biologic half-life and is shown to be several times more effective than somatostatin in the suppression of growth-hormone secretion in animals. Somatostatin analogs that have been conjugated with various metal chelators and labeled with $^{64}$Cu for evaluating SSR-positive tumors in rodent models and humans are represented in Figure 3.

![FIG. 2. Transaxial positron emission tomography/computed tomography (PET/CT) images showing the CT image (top left), $[^{18}$F]-FDG (Fluorine-18-2-fluoro-2-deoxy-D-glucose) image, $^{60}$Cu-ATSM and $^{64}$Cu-ATSM images recorded between 30 and 60 minutes in 2 patients with known cervical cancers. (A) Images recorded for a patient who responded to conventional radiotherapy and (B) images from a nonresponder. Reprinted by permission of the Society of Nuclear Medicine from reference 64.](image-url)
In one of earlier studies with SSRs, \textit{in vitro} and \textit{in vivo} evaluation of $^{64}$Cu-labeled OC conjugates was performed.\textsuperscript{20} OC was conjugated with TETA for labeling with $^{64}$Cu, and this agent was compared with $^{111}$In-DTPA-D-Phe\textsubscript{1}-OC ($^{111}$In-DTPA-OC; Octreoscan,\textsuperscript{66} Coviden, Hazelwood, MO), a single-positron emission computed tomography (SPECT) imaging agent approved for routine clinical use as a diagnostic agent for neuroendocrine cancer in the United States and Europe.\textsuperscript{67} $^{64}$Cu-TETA-OC was evaluated as a PET imaging agent in humans (8 subjects) and compared to $^{111}$In-DTPA-OC with gamma scintigraphy and SPECT imaging.\textsuperscript{22} $^{64}$Cu-TETA-OC and PET imaged more tumors in 2 patients, compared to $^{111}$In-DTPA-OC and SPECT, and in 1 patient, $^{111}$In-DTPA-OC and SPECT weakly imaged a lung lesion that was not detected with $^{64}$Cu-TETA-OC. Overall, $^{64}$Cu-TETA-OC and PET showed greater sensitivity for imaging neuroendocrine tumors, in part due to the greater sensitivity of PET, compared to SPECT.

In \textit{in vitro} and \textit{in vivo} evaluation of a second-generation somatostatin analog, $^{64}$Cu-TETA-Y\textsubscript{3}-TATE (Y\textsubscript{3}-TATE: tyrosine-3-octreotate), were conducted, where Y\textsubscript{3}-TATE differs from OC in that tyrosine (Tyr) replaces phenylalanine (Phe) in the 3-position, and the C-terminal threonine (Thr) is an acid rather than an alcohol. Y\textsubscript{3}-TATE previously showed improved targeting of somatostatin-rich tissues.\textsuperscript{24,68} $^{64}$Cu-TETA-Y\textsubscript{3}-TATE had high binding affinity to somatostatin in receptor-positive rat pancreatic tumor-cell membranes, while in rat pancreatic tumor models, $^{64}$Cu-TETA-Y\textsubscript{3}-TATE had twice as much uptake as $^{64}$Cu-TETA-OC. This reagent demonstrated superior potential as a radiopharmaceutical for the imaging and therapy of SSR-positive tumors.

After demonstrating the superiority of CB-TE2A, compared to TETA, for stably chelating $^{64}$Cu \textit{in vivo},\textsuperscript{28} CB-TE2A was conjugated to Y\textsubscript{3}-TATE and directly compared to the $^{64}$Cu-TETA-Y\textsubscript{3}-TATE conjugate.\textsuperscript{69} $^{64}$Cu-CB-TE2A-Y\textsubscript{3}-TATE was radiolabeled in high radiochemical purity with specific activities of 1.3–5.1 mCi/\textmu g of peptide at 95°C and pH 8.0.\textsuperscript{70} Biodistribution studies, using AR42J tumors implanted in male Lewis rats, revealed that this complex had higher uptake in somatostatin-positive tissues, compared to the TETA conjugate. Accumulation of $^{64}$Cu-CB-TE2A-Y\textsubscript{3}-TATE was lower at all time points, in blood and liver, and less accumulation was observed in the kidney at earlier time points, when compared to $^{64}$Cu-TETA-Y\textsubscript{3}-TATE. For example, the tumor-to-blood (T/B) ratio at 4 hours for $^{64}$Cu-CB-TE2A-Y\textsubscript{3}-TATE was 156 ± 55; for $^{64}$Cu-TETA-Y\textsubscript{3}-TATE, the T/B ratio was 8.2 ± 1.6 ($p < 0.001$). These data suggest that the $^{64}$Cu-CB-TE2A-Y\textsubscript{3}-TATE is more resistant to transchelation than the TETA analog.

The majority of somatostatin analogs that have been evaluated for PET and SPECT imaging are somatostatin agonists, and as such, they are internalized into cells via receptor-mediated endocytosis and mimic the behavior of somatostatin itself. The belief has been that greater cellular internalization of a radiolabeled somatostatin analog \textit{in vitro} is a predictor of improved tumor uptake \textit{in vivo}. This has been demonstrated by the group at Rotterdam for $^{111}$In-labeled somatostatin analogs\textsuperscript{71,72} as well as by our group.\textsuperscript{24,69} In 2006, Ginj et al. showed that an $^{111}$In-labeled somatostatin receptor type 2 (SST\textsubscript{r}2) antagonist, sst2-ANT, had improved uptake, compared to $^{111}$In-DTPA-Y\textsubscript{3}-TATE,\textsuperscript{73} in mice bearing SST\textsubscript{r}2-transfected HEK-cell tumors. The researchers showed that sst2-ANT was not internalized in the HEK cells and demonstrated classical antagonist behavior. $^{64}$Cu-CB-TE2A-sst2-ANT was compared with $^{64}$Cu-CB-TE2A-Y\textsubscript{3}-TATE in AR42J tumor-bearing rats.\textsuperscript{74} $^{64}$Cu-CB-TE2A-sst2-ANT showed low levels of internalization in AR42J cells and similar uptake to $^{64}$Cu-CB-TE2A-Y\textsubscript{3}-TATE \textit{in vivo} at early time points. An interesting characteristic of the SST\textsubscript{r}2 antagonist is that it appears to bind to ~15-fold higher number of receptors than the agonist (23,000 versus 1551 fmol/\textmu g protein), but with ~17-fold decreased affinity (26 vs 1.5 nM). However, $^{64}$Cu-CB-TE2A-sst2-ANT showed longer retention in the AR42J tumor, resulting in improved T/B (72) and T/M (93) ratios at 24 hour postinjection, compared to $^{64}$Cu-CB-TE2A-Y\textsubscript{3}-TATE (T/B, 20; T/M, 45).\textsuperscript{74}

**Copper-64-Labeled Integrin-Targeting Peptides**

Integrins are transmembrane proteins that regulate cell-cell and cell-matrix interactions. They are dimers that consist of two noncovalently bound subunits ($\alpha$ and $\beta$) that have an extra-
cellular domain arranged in a characteristic way that imparts different adhesion properties to the cell. Integrin proteins have been found to play important roles in angiogenesis and tumor metastasis. So far, 24 different integrins have been identified, constituted by combinations of 18-α and 8-β subunits. Alpha v beta 3 (αvβ3) is one of the most widely studied integrins, since it is upregulated in endothelial cells involved in active angiogenesis but not in quiescent endothelial cells, making it an ideal biomarker for angiogenesis and tumor imaging. Tumors where αvβ3 are found to be highly expressed include glioblastomas, breast and prostate tumors, malignant melanomas, and ovarian carcinomas. The αvβ3 integrin binds to extracellular proteins through a specific binding pocket that recognizes the three-amino-acid sequence, arginine-glycine-aspartic acid (Arg-Gly-Asp or RGD). This discovery has led to the design of many RGD-based imaging agents, and several investigations involving the 64Cu radiolabeled complexes have been reported (Figure 4).

Chen et al. conjugated 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA) to c(RGDyK) and labeled it with 64Cu for breast-cancer imaging studies but found only moderate uptake in U87MG human glioma tumors [1.44 ± 0.09 percent injected dose per gram [%ID/g] at 4 hour postinjection] with relatively high liver and kidney retention (2.84 ± 0.17 and 1.98 ± 0.06 %ID/g at 4 hour postinjection, respectively). In order to improve tumor uptake and in vivo kinetics, they substituted the monomeric RGD derivative for dimeric compounds (E[c(RGDyK)2 and E[c(RGDfK)2]) and observed improved tumor targeting. However, kidney uptake remained too high for the compounds to be considered for further clinical studies. In an attempt to modulate kidney retention, polyethylene glycol (PEG) groups were added to the monomeric RGD peptide derivative, and it was observed that 64Cu-DOTA-c(RGDyK)-PEG had very similar uptake in brain tumors, compared to 64Cu-DOTA-c(RGDyK), but a much lower liver uptake and a faster clearance from blood and kidneys. By using tetrameric and octameric RGD derivatives, binding affinity and tumor uptake in glioblastoma cells improved; however, liver and kidney uptake were also increased. Shi et al. examined the effects of linkages (Gly-Gly-Gly and PEG4) between cyclic RGD dimers for agents labeled with 64Cu by using the DOTA chelator. This group showed that these linkages improved the tumor uptake, compared to simple RGD dimers, potentially due to having the appropriate distance between the two RGD peptides that allows binding to two different receptors simultaneously. This strategy can be applied to other receptors as well using molecular modeling to determine the distances between receptors on tumor cells.

In a recent patent, Kimura et al. reported on the conjugation of DOTA to a library of many “miniproteins” derived from knottin peptides whose 25–40-amino-acid sequences have been enriched by an RGD loop. After screening for initial
integrin-binding ability, some of the chelators were labeled with $^{64}$Cu. Biodistribution and micro-PET imaging studies showed good specific uptake in U87MG tumors (glioblastoma). However, kidney uptake was consistently higher than tumor uptake over a 25-hour period.96 Sprague et al. conjugated c(RGDyK) to a different chelator, CB-TE2A, and found that the corresponding $^{64}$Cu complex was taken up specifically by osteoclasts,93 which are upregulated in osteolytic lesions and bone metastases.92 These investigations open the possibility of other applications for imaging $x$, $\beta_1$ in diseases, such as osteoarthritis or osteoporosis, as well as imaging osteolytic bone metastases.

Wei et al. compared two RGD peptides labeled with two highly stable chelating systems, CB-TE2A-c(RGDyK) and diamsar-c(RGDfD), in M21 and M21L human melanoma tumor-bearing mice.93 This study showed that although both chelator-peptide conjugates had similar binding affinity for isolated $x$, $\beta_1$, the tumor targeting in vivo was better for $^{64}$Cu-CB-TE2A-c(RGDyK) than the diamsar analog. There was also improved blood and liver clearance for $^{64}$Cu-CB-TE2A-c(RGDyK). Some of these differences could be due to the differences in the peptides used, as well as the fact that diamsar had a very short linkage between the aspartic acid in the 5-position and the chelator.

$^{64}$Cu-Labeled Antibodies for Tumor Targeting

Targeting epidermal growth factor receptor 1

The epidermal growth factor (EGF) family of membrane receptors (EGFR) is one of the most relevant targets in the tyrosine kinase family. EGFR expression is increased in many human tumors such as breast cancer, squamous-cell carcinoma of the head and neck, and prostate cancer.94 Activation of EGFR contributes to several tumorigenic mechanisms, and in many tumors, EGFR expression may act as a prognostic indicator, predicting patient survival and/or more of the presence of diseases in advanced stages.94 At present, monoclonal antibodies (mAbs), which block the binding of EGF to the extracellular ligand-binding domain of the receptor, have shown promise from a therapeutic standpoint. Cetuximab (C225; Erbitux,94 Bristol-Myers Squibb, New York, NY) was the first mAb targeted against the EGFR approved by the U.S. Food and Drug Administration (FDA) for the treatment of patients with EGFR-expressing, metastatic colorectal carcinoma. Cetuximab binds competitively to the extracellular domain of EGFR with an affinity comparable to the activating ligand to the receptor.95,96 Cai et al. reported the evaluation of $^{64}$Cu-DOTA-cetuximab in several tumor-bearing mouse models.97 Using Western blot analysis, a positive correlation was shown to exist between the expression of EGFR and uptake of $^{64}$Cu-DOTA-cetuximab in several different EGFR-expressing tumor-bearing mouse models. At Washington University, St. Louis, MO, $^{64}$Cu-DOTA-cetuximab was synthesized for the small-animal PET imaging of EGFR expression in A431 tumor-bearing mice.98 Highly EGFR-expressing A431 and low-EGFR-expressing MDA-MB-435 cells were compared. An equilibrium dissociation constant ($K_D$) of 0.28 nM was obtained with the A431 cells, and the $K_D$ and $B_{max}$ (maximum receptor density) were in agreement with the reported literature values of unlabeled cetuximab with A431 cells.98 In vivo evaluation of $^{64}$Cu-DOTA-cetuximab was performed in A431 and MDA-MB-435 tumor-bearing mice. Both biodistribution and micro-PET data showed a higher uptake in the EGFR-positive A431 (Figure 5) tumor than in the EGFR-negative MDA-MB-435 tumor. Metabolism experiments were also performed to determine the extent of $^{64}$Cu transchelation to blood, liver, and tumor proteins in A431 tumor-bearing mice. The results showed minimal metabolism of $^{64}$Cu-DOTA-cetuximab in the blood out to 24 hours postinjection. Liver metabolism studies, using size-exclusion chromatography, demonstrated that transchelation of $^{64}$Cu to three proteins occurs; these were identified as SOD and metallothionein, while the third metabolite was believed to be a protein aggregate.

$^{64}$Cu-DOTA-cetuximab has also been evaluated for correlating EGFR densities on the surface of five different cervical cancer lines with the EGFR-messenger RNA (mRNA) expression. Based on the cellular data, micro-PET imaging was performed on tumor-bearing mice, using the highest expressing cervical cancer cell line, CaSki. For the in vitro analysis, five cervical cancer cell lines were selected after a screen of 23 human cervical cancer lines, based on their level of EGFR gene expression by gene-expression microarray analysis. The five cell lines had different ranges of EGFR expression with the following order: CaSki (high), ME-180 and DcTc2 4510 (both midrange), HeLa (low), and C-33A (negative). The cell-surface EGFR expression was evaluated by conducting saturation binding assays at 4°C, and the results paralleled the levels of EGFR expression determined by microarray analysis. In vivo biodistribution and small-animal PET studies with $^{64}$Cu-DOTA-cetuximab in CaSki tumor-bearing nude mice showed relatively high tumor uptake at 24 hour after injection (13.2 %ID/g), with significant retention of radioactivity in blood and liver as well. Overall, this study demonstrated that $^{64}$Cu-DOTA-cetuximab is a useful marker of EGFR-expression levels, as well as a potential PET agent for determining patient-specific therapies and therapeutic monitoring.

Other $^{64}$Cu-labeled mAbs for tumor targeting

The SarAr chelator was attached to the anti-GD2 mAb, 14.G2a, and its chimeric analog, ch14.18, that target disialogangliosides overexpressed on neuroblastoma and melanoma.99 Biodistribution studies in athymic nude mice bearing subcutaneous (s.c.) neuroblastoma (IMR-6, NMB-7) and melanoma (M21) xenografts showed that 15%–20% of the ID/G accumulated in the tumor at 24 hours after injection, and only 5%–10% of the ID accumulated in the liver, a lower value than typically seen with other chelators. Uptake by a GD2-negative tumor xenograft was significantly lower (<5 %ID/G). This study demonstrates the utility of the highly stable SarAr chelation system, which enables the formation of stable $^{64}$Cu complexes attached to mAbs by using mild radiolabeling conditions.

Copper-64-Labeled Nanoparticles

Nanotechnology is an applied science that creates and studies molecules or aggregates that have an overall size in the 1–1000-nm range (<1 μm). In the last few years, nanodevices and -particles have been used in biomedical studies
FIG. 5.  (A) Projection micro-PET (positron emission tomography) images of A431 tumor-bearing nude mice after 20 and 46 hours postadministration of $^{64}$Cu-DOTA-cetuximab, with and without an injected blocking dose 20 hours prior to the imaging dose (5.6 MBq, 6 g, left; 5.6 MBq, 1 mg of cetuximab, right). (B) Coronal micro-PET images of $^{64}$Cu-DOTA-cetuximab in A431 [epidermal growth factor receptor (EGFR)-positive] and MDA-MB-435 (EGFR-negative) tumor-bearing mice after 19 and 48 hours postadministration of $^{64}$Cu-DOTA-cetuximab. (C) micro-PET/computed tomography coregistration images of $^{64}$Cu-DOTA-cetuximab in a mouse bearing both A431 and MDA-MB-435 tumors (arrow) at 24 hours postinjection. Reprinted by permission of the Mary Ann Liebert, Inc., publishers from reference 98.
investigating new and improved diagnosis and therapy agents. Oncology is one of the disciplines that has benefited the most from nanotechnology. Several nanoparticles are used in diagnostic assays for cancer, as contrast agents for MRI, as drug-delivery agents, as tumor visualization agents during surgery, and as therapeutic agents. Several types of nanoparticle platforms have been evaluated for imaging applications, including iron-oxide nanoparticles, gold nanoparticles, liposomes, emulsions, dendrimers, and nanotubes (see Figure 6 for some examples). Nanoparticles conjugated with bifunctional chelators and targeting ligands are particularly useful for PET imaging purposes because their higher surface area per volume allows a higher number of targeting residues and radionuclides per particle, which, in turn, translates into higher affinity and higher specific activity, respectively.

Studies have been performed to determine the pharmacokinetics of nontargeted nanostructures labeled with $^{64}$Cu by using the DOTA chelator. Pressly et al. prepared well-defined amphiphilic copolymers with a predetermined number of reactive functionalities, with PEG chains of variable length and low polydispersity. Upon collapsing in water, these polymers formed three-dimensional, three-layered nanoparticles with a hydrophobic inner core surrounded by a hydrophilic shell where the functional groups are located and, finally, a PEG outer shell. The thickness of each layer, the number of reactive sites, and the dimension of the particle are determined by the composition of the initial linear polymer. When DOTA molecules were conjugated to these nanoparticles, $^{64}$Cu labeling was achieved and biodistribution studies were conducted. Not surprisingly, particles with longer PEG-chain length had longer circulation in blood and lower liver uptake.

Sun et al. synthesized shell-cross-linked nanoparticles (SCKs) by cross-linking to different degree micelles formed by amphiphilic block copolymers. When TETA was incorporated onto the final SCKs, the yield was low and the labeling efficiency was unsatisfactory. This problem was solved by preincorporating the copper chelator (DOTA in this case) into the copolymer before the nanoparticles were formed. Tuning of the pharmacokinetics of these particles was performed by introducing different numbers and different lengths of PEG chains. The extent of cross-linking and the dimensions of the linker between nanoparticle and
copper chelator were found to have a dramatic impact on the specific activity of the radiolabeled particle.\textsuperscript{122}

The majority of targeted nanoparticles that have been evaluated have been conjugated with RGD peptide for the targeting of \(\alpha_v\beta_3\) integrin. Cai et al. conjugated c(RGDyK) and DOTA to quantum dots (QD), obtaining a 20-nm nanoparticle having about 28 DOTA and 90 RGD residues on its surface.\textsuperscript{123} They observed selective targeting of the vasculature of \(\alpha_v\beta_3\)-positive tumors, such as U87MG human glioblastoma, with minimal extravasation, which would be necessary for high tumor uptake. This led to a lower than expected tumor uptake, with most of the \(\text{\textsuperscript{64}}\text{Cu}\text{-DOTA-QD-RGD}\) being taken up by the liver, spleen, and bone marrow. The researchers concluded that smaller particles would probably have improved tumor-targeting properties due to easier extravasation and lower reticuloendothelial system uptake.\textsuperscript{123}

Lee et al. reported 5-nm iron-oxide nanoparticles coated with polyaspartic acid functionalized with an estimated 35 RGD peptides and 30 DOTA macrocycles per particle.\textsuperscript{124} PET studies gave high contrast images of the tumor; however, liver uptake was still high. This behavior may be explained by the fact that while the core diameter of the particles was 5 nm, their hydrodynamic size was much larger (45 nm), so the same problems observed with the QD nanoparticles persisted.\textsuperscript{124}

One of the most successful examples of tumor targeting with \(\text{\textsuperscript{64}}\text{Cu}\)-labeled RGD-conjugated nanoparticles involves the use of single-walled carbon nanotubes (SWNTs).\textsuperscript{133} Here, a comparison of SWNT that contained different sizes of PEG gave high contrast images of the tumor; however, liver uptake was still high. This behavior may be explained by the fact that while the core diameter of the particles was 5 nm, their hydrodynamic size was much larger (45 nm), so the same problems observed with the QD nanoparticles persisted.\textsuperscript{124}

Conclusions

\(\text{\textsuperscript{64}}\text{Cu}\)-based radiopharmaceuticals are being explored as agents for the delineation of disease in humans. By exploitation of the chemistry of Cu(II) and the decay characteristics of \(\text{\textsuperscript{64}}\text{Cu}\), agents based on small molecules, peptides, and larger biomolecules, such as antibodies and nanoparticles, are in development for clinical translation. A diverse array of highly specific molecular \(\text{\textsuperscript{64}}\text{Cu}\)-radiopharmaceutical imaging probes will inevitably lead to improved patient-specific treatments.

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Disclosure Statement

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