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Monoclonal antibodies for copper-64 PET dosimetry and radioimmunotherapy

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Key words: monoclonal antibodies, copper-64, positron emission tomography, tumor dosimetry, radioimmunotherapy, colon cancer, nude mice

Background: We previously described a two-antibody model of 64Cu radioimmunotherapy to evaluate low-dose, solid-tumor response. This model was designed to test the hypothesis that cellular internalization is critical in causing tumor cell death by mechanisms in addition to radiation damage. The purpose of the present study was to estimate radiation dosimetry for both antibodies (mAbs) using positron emission tomography (PET) imaging and evaluate the effect of internalization on tumor growth.

Results: Dosimetry was similar between therapy groups. Median time to tumor progression to 1 g ranged from 7–12 days for control groups and was 32 days for both treatment groups (p < 0.0001). No statistically significant difference existed between any control group or between the treatment groups.

Material and Methods: In female nude mice bearing LS174T colon carcinoma xenografts, tumor dosimetry was calculated using serial PET images of three mice in each group of either internalizing 64Cu-labeled DOTA-cBR96 (DOTA = 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid) or non-internalizing 64Cu-labeled DOTA-cT84.66 from 3 to 48 h. For the therapy study, controls (n = 10) received saline, DOTA-cBR96 or DOTA-cT84.66. Treatment animals (n = 9) received 0.890 mCi of 64Cu-labeled DOTA-cBR96 or 0.710 mCi of 64Cu-labeled DOTA-cT84.66. Tumors were measured daily.

Conclusions: PET imaging allows the use of 64Cu for pre-therapy calculation of tumor dosimetry. In spite of highly similar tumor dosimetry, an internalizing antibody did not improve the outcome of 64Cu radioimmunotherapy. Radio-resistance of this tumor cell line and copper efflux may have confounded the study. Further investigations of the therapeutic efficacy of 64Cu-labeled mAbs will focus on interaction between 64Cu and tumor suppressor genes and copper chaperones.

Introduction

Copper-64 is a radionuclide produced by a cyclotron with an intermediate half-life (T1/2 = 12.7 h) that decays by both β+ (655 keV, 17.4%) and β (573 keV, 39.0%) emission, making it suitable for labeling monoclonal antibodies (mAbs) for positron emission tomography (PET) imaging and radioimmunotherapy (RIT) of cancer. Previous experiments in xenograft-bearing rodent models have demonstrated tumor cytotoxicity of internalizing 64Cu radiopharmaceuticals superior to other nuclides, but at much lower tumor absorbed doses.

Two studies in particular offer tantalizing evidence of cytotoxicity in addition to traditional radiation damage mechanisms. Connett and others reported 82% complete tumor responses to the 64Cu-labeled mAb IA3 in Golden Syrian hamsters bearing GW39 xenografts, at a tumor absorbed dose of only 586 rad (5.86 Gy).1 Lewis and others reported complete, but temporary, tumor remissions using the somatostatin analogue 64Cu-TETA-Tyr³-octreotate in the highly aggressive CA20948 rat pancreatic tumor model at a low tumor absorbed dose.2 Evaluation of intracellular distribution of 64Cu offers some potential insight into additional cytotoxicity mechanisms. In vivo distribution studies in rats of 64Cu-TETA-/octreotide demonstrated transchelation of 64Cu to superoxide dismutase (SOD) in the liver.3 Other experiments identified 64Cu from 64Cu-TETA-octreotide in the nucleus (19.5%) and mitochondria (21.1%) of AR42J rat pancreatic tumor cells in vitro over a 24 h period.4 As there was no evidence that the somatostatin analogue itself had accumulated in these locations, it is possible that 64Cu transchelates to copper cofactor enzymes, metalloproteins and copper-handling chaperones following internalization.

We previously reported the development and characterization of a two-antibody model for comparison of 64Cu RIT.5 We confirmed the internalization of the mAb cBR96 which recognizes the Lewisy ceramide variant present in multiple human and veterinary carcinomas.6,7 We also confirmed that the mAb
tumor response to an internalizing versus a non-
internalizing mAb at the calculated tumor absorbed
dose of 10 Gy in a mouse xenograft model of cancer.

Results

PET/CT imaging. Representative PET/CT images for both $^{64}$Cu-labeled mAbs at time points from 3–48 h are shown in Figure 1. Tumor uptake was heterogeneous in most studies at the 24 and 48 h time points (Fig. 2). Tumor uptake of $^{64}$Cu-DOTA-cBR96 was 5.06% ID/organ at 3 h, 12.38% ID/organ at 26 h and 16.12% ID/organ at 48 h. Tumor uptake for $^{64}$Cu-DOTA-cT84.66 was 7.25% ID/organ at 3 h, 17.45% ID/organ at 25 h and 20.24% ID/organ at 49 h. There were no statistically significant differences between conjugates at any time point, although the power of the test is limited due to the small numbers of mice that could be imaged daily. This pattern of uptake was different from that seen in the traditional biodistribution studies in which tumor accumulation of $^{64}$Cu-DOTA-cBR96 was significantly more rapid at 3 h than that of $^{64}$Cu-DOTA-cT84.66. Using a Monte Carlo N-particle Transport Code, the calculated absorbed dose to the tumors was 484 rad/mCi (131 mGy/MBq) for $^{64}$Cu-DOTA-cBR96 and 643 rad/mCi (174 mGy/MBq) for $^{64}$Cu-DOTA-cT84.66.

Qualitative analysis of the PET images revealed few regions of radioactivity accumulation that were not predicted by the traditional biodistribution studies. What did appear unexpectedly in the 3 h images for $^{64}$Cu-DOTA-cBR96 was radioactivity associated with the vascular bed of the tail base. A lesser accumulation of radioactivity was visible in the tail of mice receiving $^{64}$Cu-DOTA-cT84.66, likely associated with small, perivascular leakage at the injection site. This radioactivity largely washed out in both cases over the subsequent 48 h period. Minor intestinal uptake was visible in the 48 h scan of mice receiving $^{64}$Cu-DOTA-cBR96. As with the traditional biodistribution studies, the hepatic radioactivity of $^{64}$Cu-DOTA-cT84.66 was clearly greater than that of $^{64}$Cu-DOTA-cBR96. This was best illustrated by the 48 h images depicted in Figure 1C. Intestinal radioactivity was clearly visible in mice imaged with $^{64}$Cu-DOTA-cT84.66 at 48 h.

Therapy study. The growth curves of the tumors are presented in Figure 3. Aggressive, unrestricted tumor growth was evident for the three control groups. Both experimental groups displayed a tumor growth delay, followed by unrestricted tumor growth. Abrupt decline in tumor volume was recorded for 22 of the 48 mice in the study. This decrease represented sudden ulceration and collapse of the tumor. The majority of these events occurred when tumor masses reached 1 g or larger. Mice were sacrificed when the tumor mass reached 3 g, the tumor developed severe ulceration or the tumor impaired the ability of the mouse to ambulate. Because of the tendency to ulcerate at a tumor burden

cT84.66, which recognizes carcinoembryonic antigen (CEA)
is non-internalizing. This antigen is also present on numerous
human carcinomas and reported in veterinary hepatocellular
carcinomas, rete testis mucinous adenocarcinomas and choroid
plexus carcinomas. The biodistributions of these antibodies
were characterized in an LS174T nude mouse model of colon
cancer and tumor dosimetry was estimated.

The purpose of these experiments was to test the hypothesis that internalization of $^{64}$Cu is the single necessary step in caus-
ing low-dose cytotoxicity with RIT of cancer. An imaging study
was performed to test the hypothesis that the actual tumor dose
received from the therapeutic administration would be equiva-

$^{64}$Cu-DOTA-cBR96. This was best illustrated by the 48 h images depicted in Figure 1C.
over 1 g, the most meaningful end point was the time to progression of tumor mass to 1 g. The Kaplan-Meier curve describing time to progression is presented in Figure 4. The median time to progression to 1 g ranged from 7–12 days for the three control populations. The median time to progression to 1 g was 32 days for both of the experimental groups. There was no statistically significant difference between the medians among the control groups of saline and unlabeled antibodies. The two experimental groups were also statistically indistinguishable; however, there was a statistically significant difference between the experimental and control groups (p = 0.0001). There was no evidence of toxicity in any of the mice in the experimental groups or the control groups receiving unlabeled mAbs.

Discussion

This therapy study yielded mixed results, some expected and some surprising. As hypothesized it is clear from the similarity between the saline control and the unlabeled DOTA-cBR96 and DOTA-cT84.66 groups that the mAbs did not initiate antibody-dependent cellular cytotoxicity (ADCC) in these animals. There was no significant difference between the times to progression to 1 g tumor mass, and no growth delay was observed in the mice receiving unlabeled antibodies compared to the mice receiving saline. Although an in vitro study previously demonstrated evidence for direct cytotoxicity as well as ADCC by unmodified BR96,7 to our knowledge there are no in vivo experiments that confirm this effect. The lack of direct cytotoxicity or ADCC in this study may be due to ambient concentrations below those which would be optimal for cell killing or potentially due to interference with this mechanism by DOTA conjugation. In either case antibody-induced immune response did not affect the outcome of this therapy study.

What was unexpected was the lack of support for the hypothesis that internalization is the single critical step in the unusual cytotoxicity previously demonstrated for internalizing 64Cu-labeled radiopharmaceuticals. Our in vitro studies confirmed the internalizing property of 64Cu-DOTA-cBR96 and the non-internalizing properties of 64Cu-DOTA-cT84.66.2 However, the 64Cu efflux studies previously published offer the possibility that a lack of persistence of 64Cu within the cells to interact with copper chaperones after internalization may explain the lack of differential therapeutic efficacy between these two mAbs.3 Cellular efflux has not been evaluated for 64Cu- and 131I-labeled 1A3 in GW39 cells, used in the experiment in which 64Cu-BAT-2IT-1A3 exhibited markedly superior tumor responses (Anderson CJ, personal communication). Rapid efflux of internalized 64Cu is a tempting explanation for the results of these experiments, but remains speculative as cellular efflux is difficult to quantify in vivo. What is likely, however, is that the trafficking of 64Cu into the nucleus and mitochondria previously described in reference 4, would cause accumulation, and thus residualization, of radioactivity within the cells. Recently, trafficking of 64Cu to the nucleus has been associated with the expression of wild-type p53.13 The LS174T cell line expresses wild-type p53, but did not show evidence of residualization.14 It is possible that chelation with a more stable ligand would have resulted in more favorable intracellular trafficking. Boswell and others have described a cross-bridged cyclam with greater stability than traditional chelators such as DOTA.15 Similarly, a bombesin peptide labeled with 64Cu using the chelator NOTA (NOTA = 1,4,7-triazacyclononane-1,4,7-triaacetic acid) demonstrated greater in vivo stability than a similar peptide conjugated with DOTA.16 However, intranuclear accumulation has been reported to be inhibited by more stable chelation in at least one model.17

The large difference in mAb dose between the biodistribution and therapy studies prompted us to confirm the tumor dosimetry by PET imaging studies. In the biodistribution experiments, different groups of mice were sacrificed at each time point. As such, mice exhibiting unusual clearance properties would be randomly distributed throughout the pool of animals, thwarting a systematic understanding of the pattern of microscopic antibody metabolism in the tumor. Furthermore, the injected dose of radiation and mAb in the biodistribution studies was nearly one log below the therapeutic dose, leaving open the possibility that a larger dose of antibody might be distributed differently.
A more ideal dosimetry study would follow the distribution of the $^{64}$Cu-labeled mAbs in the same group of living mice over a 48 h period. This experiment would allow a more complete understanding of the progression of distribution of the mAbs and $^{64}$Cu within the animals. An animal that cleared antibody or radioactivity rapidly in the reticuloendothelial system would be identified and accounted for in the modeling of the subsequent therapy experiment. The ability to image the distribution of the $^{64}$Cu-labeled mAbs with PET permits this quantification of biodistribution, including tumor targeting. Minor limitations exist in the ability to quantify the radioactivity within a region of the PET scan. However, because of the small size of the mouse, the energy of the annihilation photons detected ($511$ keV), as well as the subcutaneous location of the tumors, factors such as attenuation were likely to introduce negligible error into the calculations.

As previously discussed, the low abundance of $\beta^+$ emission from $^{64}$Cu decay allows for efficient imaging of therapeutic doses of the radiolabeled mAbs. Thus, a full therapy dose, rather than a tracer dose, may be administered to observe biodistribution under conditions of greater mAb concentration relative to antigen pool. Qualitative evaluation of the imaging studies detected little distribution to tissue that was not predicted by the results of the sacrificial studies. However, endothelial binding of $^{64}$Cu-DOTA-cBR96 in the tail vasculature was unexpected. In retrospect, this uptake may have been masked in the traditional biodistribution data because the tail was counted with the caudal half of the carcass. As with the circulating CEA pool

Figure 3. (A–E) Tumor growth curves for each control and experimental group. Tumor volumes are expressed in mm$^3$ and time is in days from treatment, after 14 days of tumor growth. Plot (A) represents the saline control mice. Plot (B) represents the unlabeled DOTA-cBR96 control mice. Plot (C) represents the unlabeled DOTA-cT84.66 control mice. Plot (D) are the $^{64}$Cu-DOTA-cBR96 experimental mice. Plot (E) are the $^{64}$Cu-DOTA-cT84.66 experimental mice. Large decreases in tumor volume were due to ulceration and necrosis of those tumors.
for $^{64}$Cu-DOTA-cT84.66, the endothelial sink for deposition of radioactivity may account for the delayed tumor accumulation of $^{64}$Cu-DOTA-cBR96, as detected by PET imaging, relative to the traditional biodistribution analysis. Vascular endothelial binding did not appear to diminish total tumor uptake, and the tumor distribution remained similar between the two mAbs at 24 and 48 h post-injection. Minor intestinal activity was observed in mice treated with $^{64}$Cu-DOTA-cBR96. Intestinal cross-reactivity with cBR96 has not been previously reported in mice. It has, however, been identified in the large intestine of dogs. It is not surprising that some cross-reactivity with normal tissue would exist. Consistent with sacrificial biodistribution results, $^{64}$Cu-DOTA-cT84.66 demonstrated increased liver and spleen activity over time, and intestinal activity became visible at 48 h as biliary excretion of $^{64}$Cu occurred. In the case of either mAb, it must be considered that copper could have been excreted into the intestinal tract unbound to antibody through the normal copper processing function of the liver. This series of non-invasive imaging studies gave striking confirmation of the biodistributions previously determined by more traditional means, with the added accuracy of repeated measures in the same living mice, and the potential to use fewer animals to obtain equivalent results.

The most valuable information gained from these images, however, was to estimate tumor dosimetry based on region of interest (ROI) evaluation. Because $^{64}$Cu was only available on a bi-monthly basis at the time, tumors were implanted 14 days prior to the experiment. During this time, tumors were expected to grow to approximately 200 mg in size. Prior to the imaging studies tumor growth was greater than expected, reaching 800–900 mg, four times the size of the tumors in the traditional biodistribution studies. On the two-dimensional PET images (Fig. 2), the tumors displayed heterogeneity of dose distribution within the parenchyma in most cases, suggesting that the large tumors had developed hypovascularized and likely necrotic areas within the tumors. Including the entire tumor area within the ROI, the larger tumors accumulated twice the radioactivity that was measured in the initial biodistribution studies. However, with four times the volume, the resulting dose to the tumors was approximately half that calculated previously. The therapy mice were injected with radiation calculated to deliver the same dose (10 Gy) to a 200 mg tumor, regardless of the mAb administered. Correcting for injected activity, for 0.89 mCi of $^{64}$Cu-DOTA-cBR96, the actual dose to the tumor was 4.31 Gy, and for 0.71 mCi of $^{64}$Cu-DOTA-cT84.66, the dose was 4.56 Gy in the larger tumors in the PET dosimetry experiment. Within 5% of each other, these doses are remarkably similar, given the potential for individual mouse variability, the limitations of sacrificial studies, and the great disparity in tumor size between the traditional biodistribution studies and PET imaging. Although not representative of the absolute dose administered in the therapy study, this result confirms that the dosimetry would have been similar between experimental groups in the therapy study.

PET dosimetry offers great advantages for future $^{64}$Cu therapy studies. Preliminary dose-finding biodistribution studies can be accomplished using far fewer mice, and allowing statistical evaluation with repeated measures to give greater power to the analysis. When therapy studies are performed, individual mice could be screened with a tracer dose and therapeutic mass, in order to compare biodistributions, metabolism and tumor uptakes. From this test dose, the appropriate dose to deliver the desired tumor absorbed dose could be calculated, ensuring that all mice receive the intended dose. These mice could then be imaged again after the therapy dose to confirm expected distributions. Using this method, future therapy studies of $^{64}$Cu-labeled delivery platforms could be better controlled, yielding more meaningful results.

In the case of this therapy study, the results of the imaging analysis suggested that the mice did, indeed, receive equivalent tumor absorbed doses from $^{64}$Cu-labeled mAbs. This lends further support to the veracity of the outcome, suggesting that the hypothesis tested cannot be supported by this two-antibody system and tumor model. Although internalization appeared in previous studies to be necessary for the enhanced cytotoxicity of $^{64}$Cu-labeled radiopharmaceuticals, the new information reported here suggests that cellular characteristics of copper handling or activity of apoptotic pathways may also play a large role in the cytotoxic mechanism. In vitro, LS174T cells clearly expressed mechanisms by which $^{64}$Cu from both mAbs was removed from the cell into the extracellular matrix. This would be expected for a non-internalizing mAb like cT84.66, as CEA is shed continuously, but internalization of cBR96 should have resulted in residualization of radioactivity within the cell. Future evaluation of these agents must continue to focus on the fate of $^{64}$Cu within the target cell.

**Material and Methods**

**Cell line.** The LS174T cell line was obtained from the American Type Culture Collection (Manassas, VA). Immediately prior
to implantation into nude mice, the cells were tested for mycoplasma and screened for a panel of 13 murine pathogens by PCR. All test results were negative, and all sentinel mice in the facility housing the nude mice tested negative for these pathogens during the course of the studies.

**Animal model.** The imaging and therapy studies were conducted in compliance with a protocol approved by the Animal Care and Use Committee of the University of Missouri-Columbia Animal Care Quality Assurance Office. Outbred female nu/nu mice (4–6 weeks of age) were obtained from Harlan Sprague Dawley (Indianapolis, IN). Mice were injected subcutaneously with 0.15 mL of Hank’s Balanced Salt Solution containing 2 x 10⁶ LS174T colon tumor cells in the right prefemoral region via a 23 gauge needle. Tumors for the imaging study reached a mean weight of 863 mg in 14 days. Tumors for the therapy study reached a mean weight of 171 mg in 14 days.

**Monoclonal antibodies.** The methods used to prepare the conjugates and perform quality control evaluation have been previously described in reference 5. Briefly, the antibodies used in these experiments were conjugated at a 10:1 molar ratio of DOTA-OSSu:mAb for the human/murine chimeric antibody cBR96 and a 20:1 molar ratio of DOTA-OSSu:mAb for the human/murine chimeric antibody cT84.66. The conjugates were purified by extensive dialysis against 0.1 M ammonium citrate buffer, pH 5.5. These conjugation ratios yielded 1.26 functional chelates per cBR96 mAb and 3.66 functional chelates per cT84.66 mAb. Immunoreactivity and serum stability were quantitative by ELISA methods for DOTA-cBR96 and by reaction with purified antigen and SE-HPLC for DOTA-cT84.66.⁸

**Antibody labeling.** Copper-64 was produced on a biomedical cyclotron at Washington University School of Medicine by previously published methods.⁸ Conjugates were labeled with ⁶⁴Cu to a specific activity of 10 μCi/µg in 0.1 M ammonium citrate, pH 5.5, for 1 h at 43°C.⁹ Diethylenetriaminepentaacetic acid (DTPA) was then added to a final concentration of 1 mM, and the reaction mixtures were left stand for 15 min at room temperature. The labeled mAbs were purified and exchanged into phosphate-buffered saline by gel-filtration spin column chromatography, using Bio-Spin 6 columns (Bio-Rad, Hercules, CA).⁹ Specific activity of labeling was determined prior to purification and radiochemical purity was determined after purification, by SE-HPLC using a Waters (Milford, MA) Delta 600 chromatograph equipped with a manual Rheodyne injector, a Waters 2487 dual wavelength UV detector, a Packard (Downers Grove, IL) 500TR Flow Scintillation Analyzer with a GAMMA-C flow cell for ⁶⁴Cu, a Waters busSAT/IN analog-digital interface, and the Waters Millennium 32 software package. A Phenomenex (Torrance, CA) BioSep-SEC-S 3000 column (7.8 x 300 mm, 5 μm, 290 Å), an isotropic mobile phase of 100 mM NaH₂PO₄/0.05% NaN₃, pH 6.8 and a flow rate of 1.0 mL/min were used.

**PET imaging.** Fourteen days prior to the start of the study, six outbred female nu/nu mice were inoculated with LS174T cells as described above. Mice were divided into two groups (n = 3). Based on previously reported data in reference 5, respective groups received either 0.89 mCi of ⁶⁴Cu-DOTA-cBR96 or 0.71 mCi of ⁶⁴Cu-DOTA-cT84.66 intravenously via the tail vein on day 1 of the study. Mice were anesthetized with isoflurane in a Plexiglas imaging chamber and lightly immobilized with gauze. PET imaging was performed using a Philips MOSAIC high resolution rodent PET scanner and CT imaging was performed using an ImTek microCAT II scanner. Each mouse was imaged for 15–60 min by PET and for 8 min by CT at 3, 24 and 48 h post-injection.

**PET tumor dosimetry.** High resolution imaging data was processed using the Philips Syntegra image fusion software for co-registration of anatomic (CT) and molecular (PET) imaging. Following previously published protocols,⁸,¹¹ a region of interest (ROI) was drawn around the complete area of the tumor, and the total number of counts/pixel/min was calculated. The total counts within the tumor area were compared to the total counts within a known standard dose of ⁶⁴Cu for each mAb to calculate the percent injected dose to each tumor (% ID/organ) at each time point, from which the doses of radiation delivered to the tumors over the imaging period were estimated in rad/mCi and mGy/MBq. Total absorbed dose was assumed to be averaged within the tumor without attempt to separate high and low-uptake regions of the tumors. The imaging results were also evaluated subjectively for distribution of radioactivity within soft tissues of the mouse body.

**Therapy study.** Fourteen days prior to the start of the study, outbred female nu/nu mice were inoculated with LS174T cells as described above. Mice were divided into three control groups (n = 10) and two experimental groups (n = 9). Control groups received either 0.15 mL saline, 0.089 mg of unlabeled DOTA-cBR96 or 0.071 mg of unlabeled DOTA-cT84.66, respectively, administered intravenously via the tail vein on day 1 of the study. A control group of ⁶⁴Cu-DOTA was not included, as an anticipated low calculated radiation dose would not be expected to be therapeutic if not targeted to the tumor and excretion would likely be extremely rapid. Experimental groups received either 0.89 mCi of ⁶⁴Cu-DOTA-cBR96 or 0.71 mCi of ⁶⁴Cu-DOTA-cT84.66 intravenously via the tail vein on day 1 of the study. Mice were weighed daily and examined for signs of overt systemic toxicity (e.g., weight loss >20%, lethargy, diarhea, cyanosis). Tumors were measured in three dimensions and tumor volume calculated by the formula length x width x depth x π/6 daily until tumor growth stabilized, then three times weekly. Mice were observed for adverse reactions on a twice daily basis. Mice were sacrificed if their tumor reached 3 g, body weight decreased >20%, the tumor ulcerated or the tumor interfered with normal ambulation.

**Statistical analysis.** Comparison of groups in the therapy study was accomplished using a Kaplan-Meier log-rank analysis. Comparison of % ID/organ values was accomplished using one-way analysis of variance (ANOVA). Differences were deemed significant at the 95% confidence level (p ≤ 0.05).

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