Anti-CD45 pretargeted radioimmunotherapy using bismuth-213: high rates of complete remission and long-term survival in a mouse myeloid leukemia xenograft model

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Introduction

For more than a decade, antibodies (Abs) conjugated to a radionuclide emitting particulate radiation have been used in the management of leukemia in an effort to deliver targeted doses of radiation to bone marrow, spleen, and other sites of disease while sparing normal organs. This radioimmunotherapy (RIT) approach has been used to achieve significant remissions in patients with acute myeloid leukemia (AML), particularly when used at high doses of radioactivity in conjunction with myeloablation.1-10 One of the major limitations of this approach, however, has been the pharmacokinetic properties of the Ab protein. Abs accrete slowly in solid tumors and are eliminated slowly from the circulation. Use of radionabeled Abs, therefore, results in prolonged exposure in radiosensitive tissues, particularly marrow, because of the extended time within the circulation. In addition, the extended time required for tumor localization of the Ab may result in loss of tumoricidal potency of the radionuclide because of ongoing isotopic decay. To address this shortcoming, the pretargeted (P)RIT system has been developed. This system differs from conventional RIT in that it uncouples the targeting agent from the radioisotope, which is administered in a separate step after facilitated clearance of non–tumor-bound targeting agent.11 Because the radioisotope can be delivered on a small molecule (<1 kDa) that is rapidly excreted through the kidneys, normal organ exposure to circulating radiation is effectively reduced by this approach. It has been demonstrated that PRIT technology can further amplify the amount of radiation delivered to CD45+ tissues and, at the same time, diminish the radiation dose to nontargeted cells.12-15

A variety of radionuclides have been investigated for RIT of leukemias, where the types of emissions used have primarily focused on the use of β-particles (131I, 90Y, and 188Re). Over the past several years, interest has developed in targeting α-emitters to leukemia cells for RIT.8,16 As opposed to the relative nonspecific cytoxicity of β-emitting constructs because of the crossfire effect, α-particle decay of radionuclides, such as 213Bi, 211At, and 225Ac, results in high-energy (6-8 MeV) delivery over a very short distance (50-80 μm). The short path length may provide a therapeutic advantage for targeting leukemic cells in the marrow and thus prevent the exposure of many normal hematopoietic stem cells to nonspecific irradiation. Therefore, the novel approach of PRIT combined with very short half-life of α-emitters may have the potential to further optimize the administration of radionuclide therapy and improve outcomes for leukemia patients.

To assess the merits of α-versus β-emitting CD45 PRIT for leukemia, we report here comparative biodistribution and therapy experiments using human leukemia xenografts implanted in athymic mice. We have demonstrated excellent localization to HEL leukemia tumor sites using both α- and β-emitting radionuclides with minimal uptake into normal organs because of elimination of nonspecific radiation exposure from blood-borne radiolabeled Ab after anti-CD45 Ab-SA pretargeting. The target-to-nontarget therapeutic...
ratios (based on radiation dose) obtained using PRIT with $^{213}$Bi were similar to those observed using $^{90}$Y. Using a novel $\alpha$-camera, we have also shown that $^{213}$Bi-DOTA-biotin uniformly distributes within tumor tissue 45 minutes after injection. Lastly, data from comparative PRIT experiments suggest that anti-CD45 PRIT using an $\alpha$-emitting radionuclide may allow for intensification of the targeted radiotherapy, with diminished toxicity, to sites of leukemic involvement to decrease the risk of relapse.

**Methods**

**Cell lines, antibodies, and production of Ab-SA conjugates**

All cell lines were obtained and maintained as described previously. The hybridoma cell lines expressing the murine anti-human IgG1, CD45 Ab BC8, and the isotype-matched human anti-bovine herpesvirus-1 Ab, used as nonspecific negative control, and all Ab-SA conjugates were produced as previously described.

**Radiolabeling**

DOTA-biotin was synthesized and labeled with either $^{90}$Y (PerkinElmer) or $^{213}$Bi (isolated from $^{225}$Ac; Department of Energy) as previously described. Radiochemical purity was typically >99% as determined by high performance liquid chromatography for each construct, and labeling efficiencies were > 90%.

**Biotinylated clearing agent**

A synthetic biotinylated CA containing 16 N-acetyl-galactosamine residues per dendrimeric molecule (Aletheon Corporation) was used to eliminate excess Ab-SA molecules from the circulation before the administration of a synthetic biotinylated CA containing 16 N-acetyl-galactosamine residues per dendrimeric molecule (Aletheon Corporation) was used to eliminate excess Ab-SA molecules from the circulation before the administration of a dendrimeric molecule (Aletheon Corporation). The N-acetyl-galactosamine residues have a high affinity for hepatic asialoglycoprotein receptors and thus facilitate the rapid hepatic clearance of residual Ab-SA conjugates from the bloodstream and their endocytosis into liver cells.

**Mice**

Female BALB/c mice, 6 to 12 weeks old, were purchased from Harlan Sprague Dawley. The animals were housed under protocols approved by the Fred Hutchinson Cancer Research Center (Seattle, WA) Institutional Animal Care and Use Committee. Results of all mouse studies are representative of at least 3 experiments.

**Biodistribution studies**

For murine biodistribution experiments, all mice were placed on a biotin-deficient diet (Animal Specialties) at least 5 days before injection of Ab-SA conjugates and radiobiody. Mice were subsequently injected with $10 \times 10^6$ HEL cells subcutaneously in the flank. Five days later, mice were given 1.4 nmol unlabeled BC8 (300 $\mu$g) or bovine herpesvirus-1 (isotype-matched negative control) Ab-SA conjugate intravenously via tail vein. Mice were then administered 5.8 nmol (50 $\mu$g) of CA intravenously 22 hours after each Ab-SA follow-up by delivery of 1.2 nmol (1 $\mu$g) of $^{90}$Y-DOTA-biotin or $^{213}$Bi-DOTA-biotin (50 $\mu$Ci) 2 hours after CA. At 10, 45, and 90 minutes after injection of $^{213}$Bi-DOTA-biotin and 1, 24, and 72 hours after $^{90}$Y-DOTA-biotin injection, tumors and normal organs were excised, weighed, and the percentage administered activities per gram of tumor or organ (% ID/g) were determined as previously described. In studies to reduce the nonspecific radionuclide uptake in the kidney, groups of 5 mice received 2,3-dimercapto-1-propanesulfonic acid (DMPS; Sigma-Aldrich) in the drinking water (1.2 mg/mL) 24 hours before $^{213}$Bi-DOTA-biotin.

**Radiation dosimetry**

Radiation absorbed doses were calculated for circulating blood and for each excised organ, tissue, and tumor based on wet tissue weights and time-activity plots that were constructed from the biodistribution data. To determine the total number of radioactive transformations (decay) in each tissue, we fit an exponential function to the time-activity data by least-squares regression analysis and then integrated the area under each curve from time of injection to infinity. The total number of decays were multiplied by the energy released from $^{90}$Y and $^{213}$Bi per decay. We accounted for the energy-absorbed fractions in that tissue and for all other neighboring tissues. This method is sensitive to the size, shape, and anatomic placement of organs and tissues in the mouse that can result in cross-organ doses from incorporated $^{90}$Y. For $\alpha$-particles from $^{213}$Bi, a self-organ absorbed fraction of 1.0 was assumed and a cross-organ absorbed fraction of 0 due the very short range of $\alpha$-particles.

**$\alpha$-camera imaging**

The $\alpha$-camera is a digital autoradiography technology using an optical registration of photons emitted from a scintillator dedicated to quantitative imaging of $\alpha$-particles in tissues ex vivo. To perform $\alpha$-camera experiments, tumors were placed in a cryomold containing a cryoprotective gel immediately after dissection. The molds were placed on a precooled aluminum block that was sunken in liquid nitrogen. The frozen tissues were placed in a HM 520 cryostat-microtome (Microm International GmbH) set to a temperature of $-19^\circ$C. Two consecutive cryosections, used for histologic comparison, were cut and transferred to a glass slide for H&E staining and for immunohistochemistry. Sections (10-16 $\mu$m) were cut and transferred to the top surface of a piece of scintillation sheet for the imaging procedure. Blue photos that impinged on the scintillator were optically imaged with a charge-coupled device detector. The pixel intensities in acquired images were linear to the activity of the $\alpha$-emitting radionuclide in the imaged tissue cryosection, allowing quantitative analysis of activity distribution at a spatial resolution approaching the microscopic level. The acquired and background-corrected $\alpha$-images were used quantitatively by analyzing and comparing the intratumoral activity distribution of the $\alpha$-emitter $^{213}$Bi at different micrometer scales. Circular regions of interest of different sizes were defined within the imaged tumors, and the mean ($\pm$ SD) pixel intensities were compared for regions of interest of different size and spatial location. Cryosections were cut and air-dried overnight. H&E sections were fixed in 10% neutral buffered formalin for 30 minutes and then stained. Immunohistochemistry sections were fixed in 10% neutral buffered formalin for 3-5 days and antigen retrieval was performed in a 65°C water bath overnight in Target Retrieval solution (pH 6.0; Dako). Slides were rinsed 3 times in TBS-T wash buffer and all subsequent staining steps were performed at room temperature using the Dako Autostainer. Endogenous peroxidase activity was blocked using 0.3% H$_2$O$_2$ for 8 minutes followed by protein blocking of 15% swine serum (Jackson ImmunoResearch Laboratories) in TBS containing 1% BSA for 10 minutes. CD45 LCA (M0701; Dako) was used at 7 $\mu$g/mL and detected with the Dako ARK kit per manufacturer’s methods. The staining was visualized with 3,3’-diaminobenzidine (DAB; Dako) for 8 minutes, and the sections were counter-stained with hematoxylin (Dako) for 2 minutes. Concentration-matched isotype control slides were run for each tissue sample (Jackson ImmunoResearch Laboratories).

**Predefined radioimmunotherapy studies using human leukemia xenografts**

To compare PRIT using either $^{90}$Y- or $^{213}$Bi-DOTA-biotin, groups of 10 mice were placed on biotin-deficient diet at least 5 days before study onset. Selected groups of mice were also treated with DMPS as described in “Biodistribution studies.” Mice were then injected subcutaneously with $10 \times 10^6$ HEL cells and 2 days after tumor injection, mice received 1.4 nmol unlabeled BC8 (300 $\mu$g) or bovine herpesvirus-1 Ab-SA conjugate intravenously. Mice were subsequently given 5.8 nmol (50 $\mu$g) of CA intravenously 21 hours later. Three hours after CA administration, groups of mice received 1.2 nmol (1 $\mu$g) of DOTA-biotin labeled with increasing doses of either $^{90}$Y (112, 800, and 1200 $\mu$Ci) or $^{213}$Bi (400 and 800 $\mu$Ci). Mice were monitored as previously described and were killed if xenografts exceeded 10% of total body weight, caused obvious discomfort, or impaired ambulation. Tumor volumes (mm$^3$) were calculated as previously...
Results

Biodistributions of radioactivity after PRIT using anti-CD45 Ab-SA conjugate followed by 90Y-DOTA-biotin or 213Bi-DOTA-biotin

The biodistributions of 90Y- and 213Bi-DOTA-biotin were evaluated after administration of an Ab-SA conjugate directed against human CD45+ leukemia xenografts given sequentially with a dendrimeric N-acetylgalactosamine-containing CA. These pretargeted biodistribution studies were performed using HEL leukemia-bearing mice injected with 1.4 nmol of anti-CD45 BC8 Ab-SA conjugate followed 22 hours later with 5.8 nmol of CA and subsequently 2 hours later with 1.2 nmol of radiolabeled DOTA-biotin. Groups of 5 mice were killed at 3 time points after injection. The radioactivity in blood, tumor, and normal organs was measured by γ-counting, corrected for decay, and expressed as %ID/g of tissue. (A) 90Y-DOTA-biotin uptake at 1 ( ), 24 ( ), and 72 ( ) hours after injection. (B-C) 213Bi-DOTA-biotin uptake at 10 ( ), 45 ( ), and 90 ( ) minutes after injection with anti-CD45 Ab-SA without (B) or with DMPS (C). (D) 213Bi-DOTA-biotin uptake at 10 ( ), 45 ( ), and 90 ( ) minutes after injection after delivery of nonbinding control Ab-SA conjugate.

Renal uptake and dosimetry

In an effort to reduce nonspecific uptake of radioactivity in the kidney, a metal chelator (DMPS) was administered in the drinking water for animals that subsequently received 213Bi-DOTA-biotin. In particular, DMPS has been shown to be effective in clearing the radioactive a-daughters of 225Ac, including 213Bi. Therefore, groups of 5 mice were treated with DMPS orally 24 hours before 213Bi-DOTA-biotin injection. DMPS-treated mice displayed significantly lower levels of radioactivity in the kidneys compared with the untreated controls at all time points (Figure 1C-D). For example, after 45 minutes, the renal uptake of 213Bi was 1.5 ± 0.2
for mice treated with DMPS, compared with 4.5 ± 0.9 for mice in the PRIT group that did not receive DMPS (P = .0006). Mice treated without DMPS displayed tumor-to-kidney ratios of radioactivity of approximately 1:1 at all time points after injection of 800 μCi of 213Bi-DOTA-biotin. In contrast, mice that were treated with DMPS and 800 μCi of 213Bi-DOTA-biotin demonstrated tumor-to-kidney ratios up to 3.5 by 45 minutes after injection.

Radiation absorbed doses for each organ per unit of administered activity were calculated based on the biodistribution data using standard Medical Internal Radiation Dose methods. The mean absorbed doses in all tissues other than kidney and tumor were similar either in the presence or absence of DMPS (data not shown because of space constraints). In DMPS-treated mice, however, the total absorbed dose of 213Bi per unit administered activity was found to be 0.7 cGy/μCi in the kidneys compared with 1.6 cGy/μCi for mice in the standard PRIT group without DMPS chelation. These data suggest a >60% reduction in the renal absorbed dose of 213Bi for DMPS-treated mice. Although the total absorbed dose in tumor was slightly lower in the DMPS-treated mice (2.0 cGy/μCi) compared with the untreated group (2.6 cGy/μCi), the tumor-to-kidney dose ratio remained approximately 2 times greater in the mice that received the renal protective agent.

α-camera imaging and quantitative autoradiography

Because α-particle emitters used in internal radiotherapy may exhibit heterogeneous distributions in tumors, a novel α-camera was used to detect α-particles in tissues ex vivo and thus assess for nonuniform activity distribution causing a nonuniform dose distribution.21 We used this quantitative imaging technique to obtain data on the 213Bi activity distribution on a suborgan level in cryosections of human HEL xenograft tumors at various times after injection of radioactivity. Nonuniform intratumoral activity distributions were found for tumor-specific 213Bi-DOTA-biotin at the 10-minute time point; however, 45 minutes after injection, the distribution was more uniform. At this 45-minute time point, the α-camera disclosed that the radionuclide was well and deeply distributed within the tumor tissue. The blue areas seen in the periphery of Figure 2 represent areas with lower activity concentration, whereas areas of red higher activity were distributed throughout the central parts of the tumor. H&E staining verified a highly proliferative tumor cell

Figure 2. Quantitative α-camera autoradiography. The intratumoral distribution of 213Bi-DOTA-biotin after anti-CD45 PRIT was analyzed using a novel digital autoradiography bio-imaging system. This α-camera system is dedicated for the ex vivo detection of α-particles in tissue and uses a scintillation setup. The technique is quantitative and fully linear toward activity content in the imaged sections. At serial times after injection (10, 45, and 90 minutes), tumors were cryosectioned (thickness 12-16 μm). Three serial, immediately adjacent sections were obtained: (A) the first for α-imaging, (B) the second for H&E staining, and (C) the third for immunohistochemistry of blood vessels using CD34. This representative, digitally collected α-camera image was obtained 45 minutes after 213Bi-DOTA-biotin injection and was color-coded to express the different levels of activity concentration of 213Bi. Insets represent smaller sections of the image shown in the next panel (inset in panel A comprises entire picture in panel B; inset in panel B comprises entire picture in panel C).
survival as a function of time. Deaths resulting from first mouse in each group. (B) Mice were analyzed for time of death because of excessive tumor growth in the injection. (A) Tumor volume curves are truncated at the time of death because of excessive tumor growth in the first mouse in each group. (B) Mice were analyzed for survival as a function of time. Deaths resulting from toxicity.

population in the imaged area, and corresponding immunohistochemistry confirmed a vascular net well established within the same area. Based on these data combined with its short path length and high linear energy transfer (LET), we further investigated the biologic outcomes of PRIT using $^{213}$Bi in a series of in vivo therapeutic studies.

**PRIT of human leukemia xenografts**

In light of favorable tumor biodistributions shown by both $\gamma$-counting and $\alpha$-camera methodologies, we performed therapy experiments to compare efficacy of $^{90}$Y and $^{213}$Bi in a minimal residual disease model of AML. Experimental groups of 10 mice were placed on a biotin-deficient diet and 2 days later were injected subcutaneously into the flank with $10^6$ HEL cells. Two days after leukemia cell injection, the mice were treated with 1.4 nmol anti-CD45 BC8 Ab-SA, followed 21 hours later by 5.8 nmol CA, and then 3 hours later with either 800 $\mu$Ci or 1200 $\mu$Ci of $^{90}$Y-DOTA-biotin or 400 $\mu$Ci or 800 $\mu$Ci of $^{213}$Bi-DOTA-biotin. Comparison groups were untreated mice that had received subcutaneous injection of HEL cells alone and mice that were injected with 1200 $\mu$Ci $^{90}$Y-DOTA-biotin or 800 $\mu$Ci $^{213}$Bi-DOTA-biotin after treatment with a nonbinding isotype-matched control Ab-SA conjugate all had exponential growth of tumors requiring death by 14 days after therapy (Figure 3). Mice that received 1200 $\mu$Ci of $^{90}$Y-DOTA-biotin developed tumors and were killed by day 64 and 1 mouse in this treatment group early exhibited regimen-related toxicity with huddling behavior and extensive loss of body weight necessitating death 14 days after therapy (Figure 3).

All $^{213}$Bi control groups exhibited anticipated growth of tumors requiring death by 16 days (untreated mice) or by 40 days (pretargeted isotype-matched negative control Ab-SA mice that received 800 $\mu$Ci of $^{213}$Bi-DOTA-biotin; Figure 4), similar to the control mice receiving $^{90}$Y-DOTA-biotin. We postulate that delayed tumor growth seen in the isotype-matched control mice injected with radiolabeled $^{213}$Bi-DOTA-biotin may have been the result of circulating $^{213}$Bi in the bloodstream, affecting the highly vascularized HEL xenografts. Overall, experimental mice pretargeted with anti-CD45 Ab-SA followed by either 400 or 800 $\mu$Ci of $^{213}$Bi-DOTA-biotin experienced superior survival rates compared with similar mice treated with higher doses of $^{90}$Y-DOTA-biotin in our...
Table 1. Comparative radiation absorbed doses after anti-CD45 PRIT using 90Y- or 213Bi-DOTA-biotin

<table>
<thead>
<tr>
<th>Tissue</th>
<th>90Y-DOTA-biotin, Gy</th>
<th>213Bi-DOTA-biotin, Gy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood</td>
<td>2.9</td>
<td>9.6</td>
</tr>
<tr>
<td>Spleen</td>
<td>2.3</td>
<td>2.3</td>
</tr>
<tr>
<td>Kidney</td>
<td>5.7</td>
<td>5.4</td>
</tr>
<tr>
<td>Tumor</td>
<td>13.1</td>
<td>17.0</td>
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Absorbed doses to blood, spleen, kidney, and tumor 10 minutes after delivery of 213Bi-DOTA-biotin and 1 hour after 90Y-DOTA-biotin was calculated for each radionuclide to deliver 5 Gy in the liver.

minimal residual leukemia model. Mice that received pretargeted anti-CD45 Ab-SA, CA, and 400 μCi of 213Bi-DOTA-biotin demonstrated a significant therapeutic benefit compared with the 800 μCi 213Bi-DOTA-biotin negative control mice (P = .002), as did anti-CD45 BC8 Ab-SA pretargeted mice that received 800 μCi of 213Bi-DOTA-biotin (Figure 4; P < .0001). One mouse pretargeted with anti-CD45 Ab-SA followed by 400 μCi of 213Bi-DOTA-biotin died by day 8 because of toxicity, whereas 2 other mice in this group were killed by day 64 because of progressive xenograft growth. In the group of mice that received anti-CD45 Ab-SA and 800 μCi of 213Bi-DOTA-biotin, 2 mice died of radiation-induced toxicity (one each on day 28 and day 49), and only one mouse developed overt leukemia leading to death on day 63. Both the 400 and 800 μCi of 213Bi-DOTA-biotin anti-CD45 pretargeted treatment groups had 70% long-term disease-free survival extending to > 120 days after therapy.

Direct comparison of anti-CD45 PRIT using 90Y- versus 213Bi-DOTA-biotin at equivalent absorbed dose

To further determine the therapeutic difference between the pretargeted α- and β-emitting radionuclide, equivalent absorbed doses to tissues were assessed (Table 1). These concentration ranges of radioactivity were determined based on equivalent absorbed doses over a specific organ volume. Thus, tissue dosimetry values for each radionuclide were normalized to an equivalent liver dose for comparative purposes. We estimated that the equivalent liver dose for mice bearing HEL tumors that received 800 μCi of 213Bi-DOTA-biotin would be a dose of 112 μCi 90Y-DOTA-biotin. Anti-CD45 Ab-SA PRIT was therefore subsequently performed using these respective doses in the model of minimal residual leukemia (Figure 5). Results from this experiment demonstrated no significant difference in survival detected between untreated mice and the group that received 112 μCi 90Y-DOTA-biotin; mice that received 112 μCi 90Y-DOTA-biotin survived 17 or fewer days, whereas mice that received no therapy survived a maximum of 14 days. There was a significant difference, however, between the group that was injected with 112 μCi 90Y-DOTA-biotin and the group that was injected with 800 μCi 213Bi-DOTA-biotin (P < .0001). Mice receiving 800 μCi 213Bi-DOTA-biotin had delayed tumor growth compared with control animals, with only one mouse requiring death (because of tumor growth on day 30 after therapy) and with 7 mice achieving long-term disease-free survival (at least 120 days).

Assessment of delayed radiation-induced toxicity after anti-CD45 Ab-SA and 213Bi-DOTA-biotin PRIT

Long-term radiotoxicity in mice surviving 120 days after PRIT with anti-CD45 Ab-SA followed by 213Bi-DOTA-biotin at 400 μCi or 800 μCi was assessed by evaluating hepatic and renal functions in addition to hematologic recovery (Table 2). Of at least 4 mice surviving in each group for 120 days, all anti-CD45 PRIT animals displayed normal body weights despite a slight nonsignificant increase in hepatic transaminase levels detected in the 400 μCi or 800 μCi 213Bi-DOTA-biotin groups compared with a group of untreated age-matched control mice. ALT, AST, and ALP levels were 389 ± 530, 214 ± 170, and 91 ± 41 U/L in the mice that received 400 μCi 213Bi-DOTA-biotin, respectively, and 371 ± 480, 385 ± 290, and 119 ± 63 U/L in those that were treated with 800 μCi 213Bi-DOTA-biotin, respectively. In contrast, untreated mice displayed ALT, AST, and ALP levels of 122 ± 84, 215 ± 94, and 69.7 ± 21 U/L (P = not significant, for all tests). Blood counts in pretargeted mice remained similar to those in untreated age-matched control mice. Mice that received 800 μCi 213Bi-DOTA-biotin had a white blood count (1000/μL), platelet count (1000/μL), hemoglobin (g/dL), and neutrophil count (per μL) of 9.9 ± 2.0, 983 ± 400, 13.6 ± 0.7, and 3145 ± 1300, respectively, whereas mice that received 400 μCi 213Bi-DOTA-biotin had counts of 12 ± 2.0, 1152 ± 170, 15.0 ± 0.2, and 5835 ± 630, respectively. Age-matched control mice had a white blood count of 14.2 ± 1.9, platelet count of 1732 ± 149, Hgb of 14.4 ± 1.2, and neutrophil count of 5820 ± 355 (P = not significant, for all tests). No significant differences were seen in BUN or creatinine levels between mice that received 800 μCi 213Bi-DOTA-biotin (34 ± 10 and 0.57 ± 0.1, respectively), 400 μCi 213Bi-DOTA-biotin (28 ± 3.0 and 0.48 ± 0.04, respectively), or age-matched control mice (26 ± 8.4 and 0.48 ± 0.01, respectively). Necropsies revealed no evidence of leukemia in any 213Bi-treated mice surviving beyond day 120, and histologic sections obtained from these mice did not show any evidence of disease or radiation-induced changes compared with age-matched controls (data not shown because of space constraints). We hypothesize that the minimal toxicity exhibited by anti-CD45 PRIT in this model may have been the result of the very

Figure 5. Leukemia xenograft regression and survival after PRIT with 213Bi- or 90Y-DOTA-biotin at equivalent absorbed doses. Athymic mice bearing HEL xenografts were injected intravenously via tail vein 2 days after tumor cell implantation with 1.4 nmol of anti-CD45 Ab-SA conjugate followed 22 hours later with 5.8 nmol of CA and subsequently with 1.2 nmol of radiolabeled DOTA-biotin 3 hours later to deliver an absorbed dose of 10 Gy from each radionuclide delivered to the liver (112 μCi 90Y-DOTA-biotin and 800 μCi 213Bi-DOTA-biotin). Control tumor-bearing control mice were untreated. (A) Tumor volume curves are not truncated. (B) Mice were also analyzed for survival as a function of time.
short path length (50 μm) of the α-particles from 213Bi compared with β-emitting isotopes, such as 90Y (~5000 μm).

### Discussion

This report critically evaluates the merits of PRIT using an α-emitting radionuclide in a myeloid leukemia xenograft model. The results described in this manuscript demonstrated that anti-CD45 PRIT using 213Bi-DOTA-biotin provided rapid tumor localization and excellent biodistributions of radioactivity, with significant improvements in efficacy compared with a PRIT approach using a β-emitting radionuclide (90Y). In vivo images using a novel α-camera displayed rapid tumor localization and favorable 213Bi radionuclide distribution at a microdosimetric level. Importantly, the pretargeted approach provided an excellent therapeutic index (target-to-nontarget ratio) for each targeting radionuclide with tumor-to-blood ratios of up to 13:1 and tumor-to-normal organ ratios of as high as 24:1, thus ensuring that the toxic effects on normal tissues were balanced by the cytotoxic effects on tumor cells. Because the therapeutic index of an antineoplastic agent depends on the balance between toxic effects on normal tissues and cells. Because the therapeutic index of an antineoplastic agent depends on the balance between toxic effects on normal tissues and cytotoxic effects on tumor cells, the pretargeted approach provided excellent tumor regressions and improved overall survival. The excellent tumor localization of the anti-CD45 Ab conjugate using PRIT and 213Bi-DOTA-biotin translated into improved xenograft survival rates compared with animals receiving 90Y-DOTA-biotin.

Although these experiments were encouraging, we recognize that this mouse leukemia xenograft model differs significantly from the state of naturally occurring leukemias in patients. In this xenograft system, the human target CD45+ cells are confined to tumors, which are more similar to chloromas than to typical disseminated leukemia. In addition, all other tissues in the xenograft model lack expression of human CD45 and will not bind anti–human CD45 Ab. Therefore, toxicity profiles in the human may not be reliably mimicked in our xenograft system. To overcome this limitation, we previously tested anti-CD45 PRIT in a murine syngeneic system in which CD45 was present on normal hematolymphoid tissues. The results of these complementary studies suggested that PRIT supported an improvement in the therapeutic index over that currently achievable with conventional RIT methodologies.

Although several clinical trials have provided encouraging results using β-emitting radionuclides (primarily 131I and 90Y) to treat malignancies, the relatively long path lengths of β-emissions can produce dose-limiting myelosuppression at conventional doses and nonhematologic dose-limiting toxicities at myeloblastic doses when used as part of a hematopoietic cell transplant conditioning regimen. Although the longer path length of these β-emitting radionuclides may be best suited for diseases with enlarged tumor masses, such as lymphomas, it has been postulated that alternative radionuclides with shorter path lengths may be better suited for clinical scenarios of minimal residual disease or where circulating isolated tumor cells are present to maintain radiation-induced toxicities at a minimum. In particular, α-emitting radionuclides with their very short path lengths may be particularly advantageous for leukemias where malignant cells are dispersed and the goal is rapid and efficient cell kill with minimal toxicities inflicted on surrounding tissues. It has been estimated that up to twice as much energy is deposited outside a 200-μm tumor when using an Ab labeled with a β-emitting radionuclide, such as 90Y or 131I, compared with an Ab conjugated to an α-emitting agent. In addition, the relatively low LET characteristics of β-particles (~0.2 keV/μm) may result in suboptimal killing of tumor cells and may ultimately contribute to the relapse of aggressive malignancies.

α-emitting radionuclides exhibit very high cytotoxicity, further making them attractive alternatives to the β-emitting radionuclides for RIT. The higher cytotoxicity delivered by an α-particle results from the large amount of energy that is emitted in a linear fashion within a few cell diameters (~50–90 μm). Moreover, the high LET energy transfer of α-emitters (~100 keV/μm) confers a high relative biologic effectiveness for cell killing compared with that for β-emitters. This high relative biologic effectiveness of α-particles results in double-strand DNA breaks that are commonly so great that cell repair mechanisms may not be effective. Not surprisingly, α-emitters have shown greater therapeutic efficacy in cell culture and animal studies than β-emitting radionuclides. Of particular relevance to our study, Friesen et al showed that a 213Bi-labeled anti-CD45 Ab, but not a 90Y-labeled anti-CD45 Ab, could overcome resistance of AML cells to β-irradiation and anthracyclines to achieve effective tumor cell killing. The 213Bi-labeled anti-CD45 Ab-induced DNA damage and apoptosis inflicted on AML cells could not be repaired, in contrast to DNA damage induced by either β- or γ-irradiation and doxorubicin. Taken together, these results suggest that α-emitting radionuclides may be superior to β-emitting radionuclides for treatment of leukemia.

Only a small number of α-emitting radionuclides have been considered suitable for in vivo applications, however, primarily because of availability and decay properties. The α-emitting radionuclides 211At (t1/2 = 7.2 hours), 212Bi (t1/2 = 60 minutes), and 225Ac (t1/2 = 45 minutes) have been favored because these radionuclides do not produce daughter radionuclides that also decay by α-emission. More recently, longer-lived α-emitting radionuclides, such as 222Ac (t1/2 = 10 days), 223Ra (t1/2 = 11.4 days), and 227Th (t1/2 = 18.7 days), have been investigated. These radionuclides do generate short-lived daughter therapeutic α-particles from 225Ac inside a targeted leukemia cell, which largely account for the increased potency of 225Ac-atomic nanogenerators over 211Bi constructs. Importantly, however, studies in mice and cynomolgous monkeys suggest that these α-daughters also have significant potential to inflict renal damage because of uptake of bismuth radioisotopes from the decay chain. A recent study by our group confirmed the efficacy of DMPS in animals treated with PRIT using an anti-CD20 Ab-SA conjugate followed by 213Bi-DOTA-biotin with a more than 50% reduction of the total radiation absorbed dose delivered to the
diseases, including hematologic as well as solid tumors. 

improvements targeting leukemic cells have the potential to 

challenges remain to achieve widespread applicability of this novel 

treatment RIT, which has been shown to result in an approximately 50% 

reduction in the glomerular filtration rate. 

In conclusion, our study confirms the encouraging results from 

pilot RIT studies using short-lived α-emitters for leukemia 

therapy. 

PRIT approach for mice with myeloid leukemia and the superiority 

of 213Bi compared with 90Y in our model. Although a number of 

challenges remain to achieve widespread applicability of this novel 

anti-CD45 PRIT approach using α-emitting radionuclides, further 

improvements targeting leukemic cells have the potential to 

increase the therapeutic index and form a basis for the treatment of 

a wide variety of malignant and life-threatening nonmalignant 

diseases, including hematologic as well as solid tumors. 

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Authorship

Contribution: J.M.P. contributed to the conception, design, analysis, 
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A.L.K. and S.I.P. contributed to the design of the research, performed 
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clinical reagents; D.S.W., Y.L., and D.R.F. contributed to the conception 
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References

1. Burke JM, Caron PC, Papadopoulos EB, et al. Cytoreduction with iodine-131-anti-CD33 antibody 
before bone marrow transplantation for advanced myeloid leukemias. Bone Marrow Transplant. 

2. Buchmann I, Bunjes D, Kolzerke J, et al. Myelo-
ablative radioimmunotherapy with Re-188-anti-
CD66-antibody for conditioning of high-risk leu-
kaemia patients prior to stem cell transplantation: 
biodistribution, biokinetics and immediate toxiciti-
Cancer Biother Radiopharm. 2002;17(2): 
151-163.

3. Bunjes D, Buchmann I, Duncker C, et al. Rhe-
ium 188-labeled anti-CD66 (a, b, c, e) monoclo-
nal antibody to intensify the conditioning regimen 
prior to stem cell transplantation for patients with 
high-risk acute myeloid leukemia or myelodys-
plastic syndrome: results of a phase I-I study. 

4. Ringhofer M, Blumentstein N, Neuhaier B, et al. 188Re or 90Y-labeled anti-CD66 antibody as part of 
a dose-reduced conditioning regimen for pa-
tients with acute leukemia or myelodysplastic syndrome 

5. Schwartz MA, Lovett DR, Redner A, et al. Dose-
escalation trial of M195 labeled with iodine 131 for 
cytoreduction and marrow ablation in relapsed 
or refractory myeloid leukemias. J Clin Oncol. 

6. Matthews DC, Appelbaum FR, Eary JF, et al. De-
velopment of a marrow transplant regimen for 
acute leukemia using targeted hematopoietic irra-
diation delivered by 131I-labeled anti-CD45 antibo-
dy, combined with cyclophosphamide and total 

7. Pagel JM, Appelbaum FR, Eary JF, et al. 131I-
anti-CD45 antibody plus busulfan and cyclophos-
phamide before allogeneic hematopoietic cell 
transplantation for treatment of acute myeloid leu-
2184-2191.

alpha particle immunotherapy for myeloid leuko-

cyclophosphamide and total body irradiation for 
advanced acute leukemia and myelodysplastic syndrome. Blood. 

10. Pagel JM, Gooley TA, Rajendran J, et al. Allogene-
ic hematopoietic cell transplantation after con-
ditioning with 131I-anti-CD45 antibodies plus flu- 
darabine and low-dose total body irradiation for 
elderly patients with advanced acute myeloid leu-
kemia or high-risk myelodysplastic syndrome. 

Chatel JF. Antibody pretargeting advances cancer 
radioimmunodetection and radioimmuno-

anti-CD20 and anti-CD45 antibodies for con-
ventional and pretargeted radioimmunotherapy of 
B-cell lymphomas. Blood. 2003;101(6):2340-
2348.

dissemianted leukemia in a syngeneic murine 
leukemia model using pretargeted anti-CD45 ra-
dioimmunotherapy. Blood. 2008;111(4):2261-
2268.

14. Pagel JM, Matthews DC, Knoyer A, et al. Preta-
greated radioimmunotherapy using anti-CD45 
monoclonal antibodies to deliver radiation to mu-
rine hematolymphoid tissues and human myeloid 

15. Lin Y, Pagel JM, Axworthy D, Pantelias A, Hedin 
N, Press OW. A genetically engineered anti-CD45 
single-chain antibody-streptavidin fusion protein for 
pretargeted radioimmunotherapy of hemato-
logic malignancies. Cancer Res. 2006;66(7): 
3884-3892.

16. Mulford DA, Scheinberg DA, Juric JC. The 
promise of targeted α-particle therapy. J Nucl 
Med. 2005;46(suppl 1):199S-204S.

17. Park SI, Shenoi J, Pagel JM, et al. Conventional 
and pretargeted radioimmunotherapy using 
bismuth-213 to target and treat non-Hodgkin lym-
phomas expressing CD20: a preclinical model 
toward optimal consolidation therapy to eradicate 
4231-4239.

18. Wilbur DS, Park SI, Chyan MK, et al. Design and 
synthesis of bis-biotin-containing reagents for 
applications utilizing monoclonal antibody-based 
pretargeting systems with streptavidin multimers. 
Bioconjug Chem. 2010;21(7):1225-1238.

of human carcinoma xenografts by a single dose of 
pretargeted yttrium-90 with negligible toxicity. 

model for calculating cross-organ beta doses 
from yttrium-90- labeled immunoconjugates. Can-

21. Back T, Jacobsson L. The alpha-camera: a quan-
titative digital autoradiography technique using 
a charge-coupled device for ex vivo high-resolution 
2010;51(10):1616-1623.

to control the errant products of a targeted in vivo 

radiation dosimetry for clinical testing of radia-
olemic monoclonal antibodies. Antibody Immuno-

24. Ballanger AM, Yang WH, Charlton DE, et al. Re-
sponse of LNCaP spheroids after treatment with
an alpha-particle emitter (213Bi)-labeled anti-
prostate-specific membrane antigen antibody


Anti-CD45 pretargeted radioimmunotherapy using bismuth-213: high rates of complete remission and long-term survival in a mouse myeloid leukemia xenograft model


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