A Preclinical Model of CD38-Pretargeted Radioimmunotherapy for Plasma Cell Malignancies

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Abstract

The vast majority of patients with plasma cell neoplasms die of progressive disease despite high response rates to novel agents. Malignant plasma cells are very radiosensitive, but the potential role of radioimmunotherapy (RIT) in the management of plasmacytomas and multiple myeloma has undergone only limited evaluation. Furthermore, CD38 has not been explored as a RIT target despite its uniform high expression on malignant plasma cells. In this report, both conventional RIT (directly radiolabeled antibody) and streptavidin–biotin pretargeted RIT (PRIT) directed against the CD38 antigen were assessed as approaches to deliver radiation doses sufficient for multiple myeloma cell eradication. PRIT demonstrated biodistributions that were markedly superior to conventional RIT. Tumor-to-blood ratios as high as 638:1 were seen 24 hours after PRIT, whereas ratios never exceeded 1:1 with conventional RIT. 90Yttrium absorbed dose estimates demonstrated excellent target-to-normal organ ratios (6:1 for the kidney, lung, liver; 10:1 for the whole body). Objective remissions were observed within 7 days in 100% of the mice treated with doses ranging from 800 to 1,200 μCi of anti-CD38 pretargeted 90Y-DOTA–biotin, including 100% complete remissions (no detectable tumor in treated mice compared with tumors that were 2,834% of initial tumor volume in control animals) by day 23. Furthermore, 100% of animals bearing NCI-H929 multiple myeloma tumor xenografts treated with 800 μCi of anti-CD38 pretargeted 90Y-DOTA–biotin achieved long-term myeloma-free survival (>70 days) compared with none (0%) of the control animals. Cancer Res; 74(4); 1179–89. ©2013 AACR.

Introduction

Bortezomib, lenalidomide, and other novel agents have significantly improved the response rates, progression-free survival, and overall survival for patients with multiple myeloma in recent years (1, 2). Despite these advances, multiple myeloma remains incurable. With currently available therapies, the 77,000 patients with multiple myeloma living in the United States will almost universally relapse and die from progressive disease. Myeloma recurrence is presumably a function of malignant plasma cell clones, and possibly precursor stem cells (3, 4), that evade or develop resistance to available therapies. The efficacy of radioimmunotherapy (RIT) in the treatment of hematologic malignancies is well established (5–7). RIT selectively delivers radiation to target cells at multifocal disease sites and facilitates escalation to radiation doses not achievable through external beam therapy. The radiosensitivity of malignant plasma cells outside of the bone marrow has been well documented in clinical settings. Local recurrence of solitary extramedullary plasmacytomas occurs in less than 10% of cases after external beam radiation alone (8). Radiotherapy is also effective as a palliative measure in patients experiencing pain or other sequelae resulting from multiple myeloma–induced osteolysis. Steep dose–response relationships have been demonstrated for most hematologic malignancies, and the impact of radiation dose escalation may be of particular importance in the case of multiple myeloma (9).

A limited number of radionuclide-based therapies have been explored in the treatment of multiple myeloma (10–14). Although each of these radionuclide-based approaches has theoretical promise, none have directly targeted radiation to the CD38 antigen on multiple myeloma cells. The directed delivery of radionuclides to multiple myeloma cells requires an antigen target that is specific, stable, and uniformly expressed at high density. CD38 is a 45-kDa stable transmembrane glycoprotein receptor expressed at a high epitope density on malignant plasma cells. In this report, both conventional RIT (directly radiolabeled antibody) and streptavidin–biotin pretargeted RIT (PRIT) directed against the CD38 antigen were assessed as approaches to deliver radiation doses sufficient for multiple myeloma cell eradication. PRIT demonstrated biodistributions that were markedly superior to conventional RIT. Tumor-to-blood ratios as high as 638:1 were seen 24 hours after PRIT, whereas ratios never exceeded 1:1 with conventional RIT. 90Yttrium absorbed dose estimates demonstrated excellent target-to-normal organ ratios (6:1 for the kidney, lung, liver; 10:1 for the whole body). Objective remissions were observed within 7 days in 100% of the mice treated with doses ranging from 800 to 1,200 μCi of anti-CD38 pretargeted 90Y-DOTA–biotin, including 100% complete remissions (no detectable tumor in treated mice compared with tumors that were 2,834% of initial tumor volume in control animals) by day 23. Furthermore, 100% of animals bearing NCI-H929 multiple myeloma tumor xenografts treated with 800 μCi of anti-CD38 pretargeted 90Y-DOTA–biotin achieved long-term myeloma-free survival (>70 days) compared with none (0%) of the control animals. Cancer Res; 74(4); 1179–89. ©2013 AACR.
expressed in pancreatic, lung, brain, and even skeletal muscle (17); however, recent gene expression profiling of human tissues demonstrate that CD38 mRNA is largely restricted to cells of hematopoietic origin with minimal expression in nonhematopoietic derived tissue limited to the thymus and prostate (18).

Unconjugated anti-CD38 monoclonal antibodies (mAb) have induced myeloma cell killing both in culture and in xenograft mouse models (19, 20). CD38 mAbs have reportedly been well tolerated in a small number of patients (21), and a phase I/II clinical trial with the anti-CD38 mAb daratumumab is currently accruing patients with relapsed or refractory multiple myeloma. CD38 mAbs linked to cytotoxic agents have also demonstrated therapeutic activity in myeloma models (22, 23).

Although conventional one-step RIT with directly conjugated radiolabeled antibodies (Fig. 1A) is effective at achieving disease control, disease eradication is thought to be limited by relatively low tumor-to-normal organ ratios of absorbed radioactivity (e.g., 1.5:1 for tumor-to-lung using 131I-anti-CD20; ref. 6). Multistep pretargeting methods can optimize delivery of the therapeutic radionuclide to tumor targets, while limiting normal organs from radiation exposure. Several approaches to pretargeting have been described (24–26). One method uses an antibody–streptavidin (Ab-SA) construct, followed by administration of a small molecule, radio–DOTA–biotin. In the first step, the tumor-reactive Ab-SA localizes to tumor sites without subjecting the rest of the body to nonspecific irradiation. After maximal accumulation of Ab-SA in the tumor, a small molecular weight radioactive moiety (radio–DOTA–biotin) possessing a high affinity for the Ab-SA bound to tumor sites is administered as a second step. Because of its small size, the second-step reagent penetrates the tumors rapidly where it binds tightly to the pretargeted Ab-SA (Fig. 1B). Unbound radio–DOTA–biotin molecules are cleared from the blood and excreted in the urine within minutes. This rapid clearance of the second-step reagent limits the time during which normal tissues are exposed to nonspecific radiation. Dissociating the slow target antigen-binding distribution phase from the radionuclide delivery phase generates more favorable target-to-normal organ ratios. As a further refinement, a clearing agent can be injected shortly before the radiolabeled small molecule to remove unbound Ab-SA from the bloodstream and prevent it from complexing with the radiolabeled small molecule in the circulation (26, 27).

This report describes a new approach to the treatment of multiple myeloma. Anti-CD38 pretargeted RIT (PRIT) demonstrates superior biodistributions when compared with directly labeled conventional anti-CD38 antibody RIT and clear evidence of therapeutic efficacy, which provides a compelling rationale for further study.
Materials and Methods

Cell lines

The human multiple myeloma cell lines NCI-H929 and L363 were a gift from David Maloney (obtained from American Type Culture Collection [ATCC]). The human Ramos B-cell lymphoma and human RPMI 8226 multiple myeloma cell lines were also obtained from ATCC. The MM1R cell line was a gift from Steven Rosen (Northwestern University, Chicago, IL) and was generated in his laboratory. Cells were grown in RPMI-1640 supplemented with 10% FBS, 50 U/mL penicillin G and 50 μg/mL streptomycin sulfate. Following two passages, cells were frozen and stored in liquid nitrogen for future use. For all studies, a fresh vial of frozen cells was thawed and grown in culture for 7 to 21 days.

Antibodies, antibody conjugates, fusion proteins, and pretargeted reagents

The OKT10 hybridoma expressing murine IgG1 anti-CD38 Ab and nonspecific IgG1 control Ab, BHV1 (specific for bovine herpes virus 1), were obtained from ATCC. The OKT10-Ab and BHV1-Ab were produced from ascites in pristine-primed mice purified by Hitrap Protein G HP 5 mL column chromatography. DOTA-Ab reagents were generated as described previously (27). Each Ab-SA was also conjugated to streptavidin (SA) to form covalent synthetic chemical conjugates using previously described methods (27). The CC49 scFv-SA-FP (fusion protein), which recognizes the TAG-72 antigen on human adenocarcinomas, was a gift from NeoRx. Expression, purification, and characterization of CC49-FP have been previously described (28, 29). To produce OKT10-CC, intact OKT10 IgG1 anti-CD38 mAb was derivatized with either a DOTA (1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid) chelate (27, 30) or streptavidin using the heterobifunctional cross-linker SMCC, using methods previously described (27). Iminobiotin and cation exchange chromatography led to purities of >95% as determined by iILC and avidin bead assay. DOTA-biotin was synthesized as described in ref. 27.

Trichloroacetic acid precipitation

The amounts of intact and degraded radiolabeled antibody in culture supernatants were estimated by trichloroacetic acid (TCA) precipitation as previously described (Supplementary Methods).

Competitive cell binding assay

L363 cells (0.5 × 10⁶ cells per sample) were plated in a 96-well plate. Serial concentrations of either OKT10-Ab, OKT10 synthetic chemical conjugate (OKT10-CC), OKT10 scFv fusion protein (OKT10-FP), or isotype control, were added to each well followed by OKT10-APC [0.1 μg, Alexa Fluor 647 Protein Labeling Kit from Molecular Probes (Invitrogen)]. The cells were incubated at 4°C for 45 minutes, washed, and fluorescence intensity was measured on a flow cytometer (Becton Dickinson).

In vitro binding characterization

The CD38 binding ability of OKT10 was assessed using Ramos targeted cells. Cells (0.1 × 10⁶ per sample) were incubated with 30 μL of 10 μg/mL 1F5-Ab (positive control), OKT10-Ab, and BHV1-Ab (nonbinding isotype control) for 30 minutes at 4°C. The cells were washed, pelleted, and resuspended in PBS. Alexa Fluor 647–conjugated OKT10 mAb binding to CD38-expressing L363 multiple myeloma cells with escalating concentrations of unlabeled OKT10-FP (A) or OKT10-CC (B). No blocking is observed with escalating concentrations of nonbinding control Ab constructs.
minutes at 4°C. Cells were washed and incubated with 25 μL of 1:64 goat anti-mouse IgG (immunoglobulin G; Fab specific) fluorescein isothiocyanate conjugate (Sigma-Aldrich) in PBS. Cells were washed and fluorescence intensity was measured on a FACS Canto I flow cytometer (Becton Dickinson). Binding ability of OKT10 to L363 targeted cells was assessed using OKT10-APC.

**Cell binding assay**

Anti-CD38 (OKT10-FP/OKT10-CC; 20 μg/mL) or non-binding control (CC39-FP/BHV1-CC; 20 μg/mL) were added to 1 × 10⁶ cells pelleted in 96-well round-bottomed plates on ice. Pellets were resuspended in the antibody solution, incubated at 4°C for 1 hour, washed, and resuspended in 200 μL Dulbecco's Modified Eagle Medium (DMEM) for 24 hours and stored at 4°C or 37°C. Cells were pelleted and then resuspended in 25 μL of ¹¹¹In–DOTA–biotin (100 ng/mL), incubated at 4°C for 1 hour, washed twice, and activity was measured on a γ counter.

**Mouse RIT and PRIT studies**

Female athymic nude-Foxn¹nu mice, ages 5 to 6 weeks, were purchased from Harlan Sprague-Dawley. NCI-H929 and L363 cells (1 × 10⁶) were injected subcutaneously into the right flank 9 to 11 days before study start dates. Mice bearing palpable plasmacytoma xenografts measuring 100 mm² ± 10% were selected for the studies and randomly assigned to experimental groups. Multiple myeloma tumor-bearing mice were placed on a biotin-free diet for 5 days and injected with either 1.4 nmol of anti-CD38 OKT10–DOTA Ab or control BHV1–DOTA Ab, each directly labeled with ¹¹¹In, or 1.4 nmol of anti-CD38 OKT10 Ab-SA (OKT10-CC or OKT10-FP) or control Ab-SA (BHV1-CC or CC39-scFvSA-FP), followed 22 hours later by 5.8 nmol (50 μg) clearing agent and 2 hours later by 1.2 nmol (1 μg) of ¹¹¹In–DOTA–biotin for biodistributions or ⁹⁰Y–DOTA–biotin labeled with 400 μCi (14.9 MBq), 800 μCi (29.6 MBq), or 1,200 μCi (44.4 MBq) ⁹⁰Y for therapy studies. Mice were monitored three times weekly for general appearance, tumor volume measurements, and body weight. Mice were injected with anti-αlso1G1 antiserum (200 μL; Wako Chemicals USA, Inc.) 9 and 5 days before the injection of Ab-SA to abrogate NK cell activity and prevent spontaneous tumor regressions. Mice were euthanized when tumors reached a maximum bidirectional measurement of ≥20 × 20 mm, when tumor ulceration occurred, or when mice lost >30% of baseline body weight, as required by the institutional animal care guidelines.

**Blood clearance studies**

Blood clearance studies were conducted according to the double-label method of Pressman (31, 32). ¹²⁵Iodine (¹²⁵I)-OKT10-CC (1.4 nmol) and ⁹⁰Y-labeled small molecular weight fragments from cells (detailed in Supplementary Materials). To demonstrate effective PRIT targeting of ⁹⁰Y–DOTA–biotin, multiple myeloma cells were labeled with Ab-SA, washed, and incubated either at 4°C, a temperature that completely inhibits endocytosis, or at 37°C. After 24 hours of incubation, ⁹⁰Y–DOTA–biotin was added and the amount of radioactivity targeted to multiple myeloma cells was measured. The difference in cell-associated radioactivity was minor, indicating that the magnitude of the ¹²⁵I channel. Counts were also corrected for radioactive decay using an aliquot of the injectate.

**Dosimetry**

Absorbed radiation doses to organs were calculated for ⁹⁰Y using β kernel methods for localized β dosimetry expressly developed for accurately calculating the radiation doses to small organs and tissues of the mouse (33, 34). These methods account for energy losses by source and take into account the organ self-dose specific absorbed fractions and the β-particle cross-organ dose contributions. Femoral bone marrow doses were determined using a model that incorporates Monte Carlo calculations of the energy-absorbed fractions in the marrow shafts (35). This model also accounts for the contributions of ⁹⁰Y on bone surfaces, if any, that may contribute to bone marrow dose.

**Statistical analysis**

Differences in multiple myeloma tumor xenograft volumes were compared by computing the means and SDs of each treatment group and using the Student t test to determine statistical significance. For relatively large differences in tumor volume, 8 to 10 mice per group were projected to provide adequate power to detect statistically significant differences. Only the detection of large differences between treatment groups was considered to be of clinical interest.

**Results**

**OKT10 anti-CD38 reagents are cell-surface stable and enable excellent pretargeting**

Experiments assessing the binding and internalization of ⁹⁰Y-labeled OKT10-Ab and fusion protein were performed using four multiple myeloma cell lines (L363, NCI-H929, RPMI-8226, and MM1R), a CD38-expressing non-Hodgkin lymphoma (NHL) line (Ramos), and four multiple myeloma patient biopsy samples. These studies evaluated the degree of internalization, cell-surface retention, passive dissociation ("shedding"), intracellular metabolism, and exocytosis of ⁹⁰Y-labeled metabolites, using flow cytometric, cell binding, and "acid-wash" methods as previously published for other targets (36, 37). Only a minority (9% ± 3%; Supplementary Fig. S1) of CD38-targeted Ab or fusion protein was internalized by L363 multiple myeloma tumor cells after 24 hours (the time interval between the two reagents in PRIT). After 24 hours, approximately 60% of initially bound OKT10-Ab remained on the cell surface, whereas 30% to 40% dissociated passively from the cell surface into the culture medium. A TCA precipitation assay was used to demonstrate that the radioactivity released into the culture medium reflected passive "shedding" of intact Ab rather than exocytosis of ⁹⁰Y-labeled small molecular weight fragments from cells (38). To demonstrate effective PRIT targeting of ⁹⁰Y–DOTA–biotin, multiple myeloma cells were labeled with Ab-SA, washed, and incubated either at 4°C, a temperature that completely inhibits endocytosis, or at 37°C. After 24 hours of incubation, ⁹⁰Y–DOTA–biotin was added and the amount of radioactivity targeted to multiple myeloma cells was measured. The difference in cell-associated radioactivity was minor, indicating that the magnitude of
endocytosis of the first-step reagent in 24 hours was not sufficient to prevent effective targeting of the second-step reagent (90Y-DOTA–biotin). Concordant results were seen for three other cell lines and the patient samples tested. These conclusions also pertain in vivo since favorable in vivo targeting of 111In–DOTA–biotin to xenografts is observed (Fig. 3B–E).

**Comparative biodistributions of radioactivity demonstrate OKT10 PRIT improves the therapeutic index**

Blood, tumor, and nonspecific organ uptake of one-step directly radiolabeled (conventional) OKT10 (OKT10–DOTA) was compared with multistep PRIT in six biodistribution experiments involving athymic nude mice bearing subcutaneous CD38– human myeloma tumor xenografts (NCI-H929 and L363). Studies were designed to evaluate the relative merits of two distinct pretargeting constructs, OKT10-CC and OKT10-FP (administered at equimolar concentrations). Sixteen groups of 5 mice each were injected with 1.4 nmol of OKT10-DOTA, OKT10-FP, or OKT10-CC (including matched antibody isotype or fusion protein–matched controls). The PRIT reagents (OKT10-FP or OKT10-CC) were followed, 18 to 22 hours later, by 5.8 nmol clearing agent and 2 hours thereafter by 1.2 nmol 111In–DOTA–biotin (1 μg). Groups of animals were euthanized at 2, 24, 48, and 96 hours after injection of the radioactivity. Tumors excised from mice pretargeted with OKT10-CC contained 8.8% ± 2.8% of the injected dose of 111In–DOTA–biotin per gram (%ID/g) after 24 hours compared with 0.9 ± 0.4 %ID/g after 24 hours in tumors excised from control mice pretargeted with a control Ab-SA chemical conjugate (BHV1-CC; all values are mean ± SD; Fig. 3B). The 24-hour tumor-to-normal organ ratios of absorbed radioactivity were 10:1; 8:1; and 6:1, respectively for lung, liver, and kidney in mice pretargeted with OKT10-CC; compared with <1:1 for the lung, liver, and kidney in control mice pretargeted with BHV1-CC.

Tumors excised from mice pretargeted with OKT10-FP demonstrated uptakes of radioactivity in tumor sites (6.4 %ID/g ± 2.3% after 24 hours) that were similar to those observed with the chemical conjugate, whereas minimal tumor uptakes were seen in the controls treated with CC49-FP (0.55 %ID/g ± 0.21% after 24 hours; all values are mean ± SD; Fig. 3C). Tumor-to-organ ratios of absorbed radioactivity with OKT10-FP PRIT were 27:1; 10:1, and 3:1, respectively for lung, liver, and kidney at the 24-hour time point compared with ≤1:1 for the same organs in control mice pretargeted with the negative control CC49-FP. The minimal tumor uptake of radiobiotin detected in both the CC49-FP and BHV1-CC control groups demonstrates the specificity of targeting with OKT10-FP and OKT10-CC (Fig. 3B and C).
Although levels of radioactivity in tumor xenografts measured 24 hours after administration of the radioactive species were similar in the OKT10-FP, OKT10-CC, and OKT10–DOTA–treated mice (6.4% ± 2.3%, 8.8 ± 2.8 %ID/g, and 9.4% ± 2.2%, respectively), PRIT (OKT10-CC and OKT10-FP) led to biodistributions of radioactivity that were far superior to the conventional (OKT10–DOTA) approach (Fig. 3A–E). At 24 hours following injection of the radiolabeled species, the tumor-to-blood ratio for the conventional RIT was <1:1, whereas for the OKT10-FP and OKT10-CC, ratios were 63:1 and 9:1, respectively, confirming the capacity for PRIT to circumvent the tumor-to-normal organ distribution limitations seen with conventional one-step RIT (38, 39).

Excellent target specificity was demonstrated for both PRIT constructs. Although tumor-to-blood ratios were most dramatic with the fusion protein, this finding seems to be a consequence of very rapid circulatory clearance of the fusion protein (see blood clearance studies below) and did not lead to superior overall biodistributions. For both constructs, the kidney represented the normal organ with the largest nonspecific radiation uptake. Tumor-to-kidney ratios were 6:1 and 3:1 for the chemical conjugate and fusion protein, respectively, 24 hours after the 111In–DOTA–biotin infusion. Thus, the kidney was identified as the organ likely to define dose-limiting toxicity in high-dose therapy settings, and the findings supported an overall advantage for OKT10-CC pretargeting.

Blood clearance studies favor OKT10-CC

Optimal tumor target accretion in PRIT models requires a sustained concentration of the first-step streptavidin reagent in the bloodstream during the biodistribution phase of the treatment algorithm, followed by rapid blood clearance after administration of biotinylated polymeric N-acetyl-galactosamine clearing agent. The rapid clearance of OKT10-CC or fusion protein complexed with clearing agent is mediated by asialoglycoprotein receptors in the liver (26). Detailed blood clearance studies conducted in athymic mice both when administered separately and coinjected (32) with both 131I-OKT10-CC and 125I-OKT10-FP at equimolar concentrations revealed that 16% of the OKT10-CC remained in the circulation 24 hours after injection, compared with only 1.4% of the OKT10-FP (not shown).

Synthetic clearing agent reproducibly removed >85% of circulating OKT10-CC or OKT10-FP from the bloodstream within 60 minutes of administration in five blood clearance studies. In a representative experiment, clearing agent (5.8 nmol) was injected 24 hours after 131I-OKT10-CC (300 µg) into 5 athymic nude mice (intravenously). Retro-orbital venous sampling was conducted at serial time points up to 24 hours (Fig. 4).

Figure 4. Effects of biotinylated N-acetyl galactosamine clearing agent on circulating OKT10-CC. Clearing agent (CA; 5.8 nmol) was injected 24 hours after 131I-OKT10-CC (1.4 nmol) into 5 athymic nude mice (intravenously). Five control mice received 125I-OKT10-CC (1.4 nmol) and no clearing agent. Retro-orbital venous sampling was conducted at serial time points up to 24 hours.

Organ dosimetry

A detailed organ dosimetry analysis was performed to calculate 90Y absorbed dose estimates after OKT10-CC pretargeting (Fig. 5). Radiation-absorbed doses to tumors, whole body, and nine normal tissues were determined on the basis of integrating the areas under time–activity curves constructed by plotting the concentration of 111In–DOTA–biotin in tissues after various intervals of time (33). The dosimetry method used accounted for organ self-dose absorbed fractions as well as β-particle cross-organ dose contributions (33). The 90Y absorbed dose estimates (Gy) per millicurie administered, calculated from the 111In tracer, generated tumor-to-normal organ ratios of 6:1 for the kidney, lung, and liver, 15:1 for the spleen, and 10:1 for the whole body.

Therapy studies

90Y–DOTA–biotin pretargeted to CD38 eradicates plasmacytomas in vivo. Therapy studies were performed in athymic mice (n = 8–10 per group) bearing subcutaneous xenografts of either L363 or NCI-H929 malignant plasma cells, producing tumors similar to human plasmacytomas. Reagent concentrations and time-points for administration of OKT10-CC, BHV1-CC, and clearing agent were identical to those reported for the biodistribution studies.

90Y–DOTA–biotin (2 µg) was labeled with 400, 800, or 1,200 µCi per mouse in three OKT10-CC groups and three control groups (untreated control; 800 or 1,200 µCi 90Y–DOTA–biotin after BHV1-CC). All L363 tumor xenograft mice in the untreated control and BHV1-CC control groups experienced exponential multiple myeloma tumor growth and 78% of the untreated control animals required euthanasia within 17 days (Fig. 6A). After 22 days, 100% of the 800 µCi and 89% of 1,200 µCi BHV1-CC control groups had required euthanasia due to progressive tumor growth. All mice pretargeted with OKT10-CC followed by 90Y–DOTA–biotin demonstrated tumor shrinkage by day 6 at all dose...
Plasma cell malignancies, including multiple myeloma, are rarely cured. Despite the higher response rates and longer survival afforded by novel agents (1, 2), the vast majority of patients with plasma cell malignancies die of progressive disease, emphasizing the importance of continuing to investigate new treatment strategies. RIT is a therapeutic modality that is not cross-resistant with chemotherapy or “novel agents” and has been underexplored in this disease.

External beam radiation can cure isolated malignant plasmacytomas and radiotherapy is frequently used to palliate painful multiple myeloma bone lesions. Sustained local disease control and durable symptom relief has been reported for 98% of lesions receiving >10 Gy (40). Furthermore, the poor prognostic implications associated with high-risk bone marrow cytogenetics in active multiple myeloma are not predictive of a decrement in the very high rates of local control and cure after external beam radiotherapy is used to treat solitary extramedullary plasmacytomas with the same cytogenetic derangements (41). This suggests that the unique attributes associated with the targeted delivery of radiation may augur clinical
efficacy even among patients classified as "high risk". When ionizing radiation has been combined with bortezomib, synergistic inhibition of multiple myeloma cell proliferation has been reported (12). It has also been suggested that the efficacy of bortezomib can be improved when combined with skeletally targeted 153samarium lexidronam (42). These findings raise the possibility that anti-CD38 RIT may be enhanced when combined with novel therapies.

Although several plasma cell–associated antigens have been identified, we elected to target CD38 because it has a high density and consistency of expression on clonal plasma cells (15, 16). We demonstrate that the receptor has minimal internalization and is not shed from the plasma cell surface after binding to OKT10. The unmodified CD38 mAb daratumumab, which binds a unique epitope on the antigen receptor, has demonstrated anti-multiple myeloma tumor cell activity both in vitro and in mouse xenografts (19); however, perhaps as a consequence of binding a different epitope, OKT10-CC administered as a single agent without radioactivity demonstrates no antitumor effects. The promising results achieved with daratumumab, however, complement our approach targeting CD38 for radionuclide delivery and validate the antigen as a desirable target. Despite the salient favorable attributes enumerated for CD38, no target antigen is perfect and it must be acknowledged that CD38 expression is not limited exclusively to malignant plasma cells. Nevertheless, the expression of CD38 on normal plasma cells and some activated lymphoid cells must be viewed in the context of the successful selection strategies that have resulted in clinically approved RIT therapeutics for other hematologic malignancies, such as 131Iodine-tositumomab (Bexxar) and 90Yttrium-ibritumomab tiuxetan (Zevalin). The efficacy and favorable toxicity profiles of anti-CD20 RIT in B-cell lymphoma have been clearly established despite expression of this antigen on normal B cells (5–7). The CD38 antigen has many attributes parallel to CD20 to support its selection as a viable target for RIT.

Several therapeutic radionuclides might be considered for RIT of multiple myeloma. We selected 90Y as the radioisotope for these studies because it is a pure, high-energy β-emitter that is readily complexed by the DOTA chelate and is commercially
Figure 7. Tumor responses in athymic nude mice (n = 10 per group) bearing L363 right flank multiple myeloma xenografts. All mice received 1.4 nmol (300 μg) of OKT10-CC as a single agent (no radioactivity) or followed by clearing agent [5.8 nmol (50 μg)] and 90Y-DOTA–biotin (800 μCi (29.6 MBq)] 24 hours after the OKT10-CC. A, mice were monitored three times weekly for tumor volume measurements and were euthanized when tumors were >10% of body weight or when ulceration occurred, as mandated by the Institutional Animal Care Committee. Curves are truncated at the time the first animal was euthanized in each group. Complete regression of multiple myeloma tumor xenografts in the group receiving OKT10-CC followed by 800 μCi (29.6 MBq) is demonstrated. Animals receiving OKT10-CC as a single agent (no radioactivity) demonstrated rapid tumor progression. B, Kaplan–Meier analysis of cumulative survival of mice bearing L363 multiple myeloma xenografts.

available in high specific activity and purity. The long path length of 90Y’s β particles permits the delivery of lethal radiation across several cell diameters, resulting in a "crossfire effect" that can eradicate antigen-negative multiple myeloma cells (or multiple myeloma cells that are in the center of a tumor cluster) by lethal radiation emitted from surrounding antibody-binding CD38+ multiple myeloma cells. Furthermore, the current biotin reagents used for PRIT can only accommodate radiometals, not 131I, and attempts to produce effective 131I-labeled biotin have generally been disappointing, reinforcing the selection of 90Y.

In this report, we demonstrate that the 90Y-based anti-CD38 PRIT approach chosen, rapidly and completely eradicates plasmacytoma xenografts with minimal toxicity. This therapeutic efficacy was predicted by anti-CD38 PRIT biodistributions that were markedly superior to those achievable by conventional RIT, as confirmed by radiation dosimetry estimates demonstrating excellent tumor-to-normal organ ratios. Although both anti-CD38-SA constructs were effective, the rapid circulatory clearance of OKT10-FP resulted in a lower area under the curve (AUC) during the distribution phase; and, despite excellent blood-to-tumor ratios of activity, the fusion protein had lower tumor-to-kidney ratios of absorbed activity (3:1) when compared with OKT10-CC PRIT (6:1) after 24 hours. Consistent with prior published reports, biodistribution studies predicted the kidney to be the dose-limiting normal organ (43). The renal biodistribution profile for the chemical conjugate, therefore, provided a rationale for selection of OKT10-CC as the best pretargeting reagent for subsequent therapy studies.

PRIT enables the administration of 90Y doses not attainable using conventional one-step RIT and seems to improve the therapeutic index, enhance efficacy, and diminish toxicity. Removal of excess OKT10-CC from the vascular compartment by clearing agent, combined with rapid urinary excretion of excess 90Y-DOTA–biotin, limited nonspecific radiation exposure to normal organs even at doses of 1,200 μCi (three times the previously defined lethal doses in conventional 90Y labeled RIT; ref. 44). Ninety percent of animals in the highest dose group demonstrated only mild transient weight loss, and regained their normal weight within 15 days (Supplementary Fig. S2), whereas animals receiving either 400 or 800 μCi demonstrated no significant drop in weight. Among the OKT10-CC animals alive and disease free >70 days after therapy, no significant toxicity was detectable.

We elected to initiate preclinical studies of CD38-targeted RIT and PRIT using a plasmacytoma model because traditional radiation dosimetry and biodistribution methods can be most accurately performed using solid tumor nodules, and are much more difficult to apply in a quantitative fashion using marrow-based, disseminated disease. Tumor xenografts are, therefore, the standard for initial RIT validation studies (e.g., CD20 in NHL, CD45 in acute myeloid leukemia; refs. 27, 45–48). In addition, xenograft models have played a vital role in validating the impressive efficacy of proteasome inhibitor therapy in multiple myeloma and we have emulated this initial approach (49).

Despite serving as a well-validated model for the assessment of new RIT targets and multiple myeloma therapeutics in large cohorts of animals, multiple myeloma tumor xenografts do not fully recapitulate the clinical spectrum of the disease (49, 50). The absence of human CD38 antigen expression in murine tissues is an inherent limitation of most xenograft models, and the xenograft model applies most directly to soft tissue plasmacytomas rather than disseminated multiple myeloma. Furthermore, xenografts do not provide a bone-based platform that recapitulates the human marrow microenvironment. To address this concern, we are conducting studies in a SCID-hu multiple myeloma mouse model and have seen preliminary results that are encouraging, but beyond the scope of the current article.

In conclusion, this is the first report of anti-CD38 RIT in multiple myeloma and the first report documenting the efficacy of PRIT in this disease. The complete eradication of multiple myeloma tumor xenografts provides a scientifically compelling rationale for further exploring high-dose anti-CD38
PRIT as conditioning therapy before autologous stem-cell transplant for multiple myeloma. Although xenograft tumors are not ideal, they have been used to accurately predict the value of CD20 and CD45 directed RIT or PRIT in lymphoma and leukemia mouse models, and have been used to establish the value of "novel agents" to treat multiple myeloma despite the same limitations. Ultimately, clinical trials will be necessary to demonstrate safety and evaluate the potential for anti-CD38 PRIT to improve progression-free and long-term survival among patients with multiple myeloma.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

Authors’ Contributions
Conception and design: D.J. Green, J.C. Jones, J.M. Pagel, D.S. Wilbur, A.K. Gopal, O.W. Press

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