Research paper

Novel theranostic agent for PET imaging and targeted radiopharmaceutical therapy of tumour-infiltrating immune cells in glioma

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ARTICLE INFO

Article History:
Received 18 February 2021
Revised 17 August 2021
Accepted 19 August 2021
Available online xxx

ABSTRACT

Background: Malignant gliomas are deadly tumours with few therapeutic options. Although immunotherapy may be a promising therapeutic strategy for treating gliomas, a significant barrier is the CD11b+ tumour-associated myeloid cells (TAMCs), a heterogeneous glioma infiltrate comprising up to 40% of a glioma’s cellular mass that inhibits anti-tumour T-cell function and promotes tumour progression. A theranostic approach uses a single molecule for targeted radiopharmaceutical therapy (TRT) and diagnostic imaging; however, there are few reports of theranostics targeting the tumour microenvironment.

Methods: Utilizing a newly developed bifunctional chelator, Lumi804, an anti-CD11b antibody (aCD11b) was readily labelled with either Zr-89 or Lu-177, yielding functional radiolabelled conjugates for PET, SPECT, and TRT.

Findings: 89Zr/177Lu-labeled Lumi804-aCD11b enabled non-invasive imaging of TAMCs in murine gliomas. Additionally, 177Lu-Lumi804-aCD11b treatment reduced TAMC populations in the spleen and tumour and improved the efficacy of checkpoint immunotherapy.

Interpretation: 89Zr- and 177Lu-labeled Lumi804-aCD11b may be a promising theranostic pair for monitoring and reducing TAMCs in gliomas to improve immunotherapy responses.

Funding: A full list of funding bodies that contributed to this study can be found in the Acknowledgements section.

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Keywords: Theranostics targeted radiotherapy immunoPET checkpoint immunotherapy gliomas bifunctional chelator

1. Introduction

Gliomas are the most common primary malignant brain tumour [1]. The current standard of care for gliomas includes a combination of surgery, chemotherapy, and radiation therapy, depending on the specific type of tumour. However, even with current treatments, the median survival times for patients with high-grade gliomas (HGG), including the World Health Organization (WHO) grade IV glioma
A theranostic approach that uses a single molecule for targeted radiopharmaceutical therapy (TRT) and diagnostic imaging could improve outcomes for malignant glioma by quantifying a pool of immunosuppressive cells bearing the biomarker CD11b and selectively targeting eradication of those cells.

Added value of this study
A single molecule, Lumi804-anti-CD11b, labelled with either 89Zr or 177Lu, enabled non-invasive imaging of TAMCs in murine gliomas. Additionally, 177Lu-Lumi804-anti-CD11b treatment reduced TAMC populations in the spleen and tumour and improved the efficacy of checkpoint immunotherapy.

Implications of all the available evidence
Together, 89Zr- and 177Lu-labeled Lumi804-antiCD11b may be a promising theranostic pair for monitoring and reducing TAMCs in gliomas to improve immunotherapy responses.

Research in context
Evidence before this study
A theranostic approach that uses a single molecule for targeted radiopharmaceutical therapy (TRT) and diagnostic imaging could improve outcomes for malignant glioma by quantifying a pool of immunosuppressive cells bearing the biomarker CD11b and selectively targeting eradication of those cells.

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A single molecule, Lumi804-anti-CD11b, labelled with either 89Zr or 177Lu, enabled non-invasive imaging of TAMCs in murine gliomas. Additionally, 177Lu-Lumi804-anti-CD11b treatment reduced TAMC populations in the spleen and tumour and improved the efficacy of checkpoint immunotherapy.

Implications of all the available evidence
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(glioblastoma), have held steady at approximately 15 months in both adults and children [1,2]. Therefore, more effective treatment approaches are urgently needed.

Theranostics are agents that enable both therapy and diagnostic imaging utilizing a single targeting molecule (e.g. an antibody or peptide). Recent theranostics have incorporated radiometals for PET imaging and targeted radiopharmaceutical therapy (TRT) [3,4]. For example, DOTA-oxalate (dotate) radiolabelled with Lu-177 (177Lu-dotate; Lutathera) or Ga-68 (68Ga-dotate; NetSpot) are FDA-approved for TRT and PET imaging, respectively [5,6], emphasizing the translational potential of theranostic agents with a single chelator for both therapy and PET.

Preclinical and clinical studies support the safety and early efficacy of immunotherapies for gliomas in both adults and children. However, the immunosuppressive tumour microenvironment (TME), largely mediated by CD11b+ tumour-associated myeloid cells (TAMCs), remains a barrier to successful glioma immunotherapy. In fact, CD11b+ myeloid-derived cells are the predominant subset of non-malignant cells in the glioma TME. These cells are considered as a high priority target for glioma treatment and improving immunotherapy responses [7,8]. TAMCs include subpopulations of tumour-associated macrophages (TAMs) and myeloid-derived suppressor cells (MDSCs), which together account for up to 40% of a glioma’s mass [9,10] and are associated with decreased survival [11,12]. Additionally, TAMCs express many inhibitory molecules that promote tumorigenesis, maintain the immunosuppressive TME, and suppress T-cell mediated anti-tumour immunity [13]. Recent translational research efforts targeting TAMCs have focused on blocking specific TAMC-expressed cell mechanisms including colony-stimulating factor 1 receptor (CSF-1R), CD47 signalling pathways, and immune-checkpoint molecules such as programmed cell death 1 programmed cell death ligand 1 (PD-1/PD-L1) [14-17]. Importantly, decreased TAMC levels in murine gliomas are associated with extended survival times [17]. Thus, selective eradication of these cells may improve patient overall survival as well as response to immunotherapy [13,18]. Moreover, monitoring TAMC levels within the glioma TME is essential to assess the efficacy of TACM-targeted therapies. Along this line, since CD11b is a predominant marker expressed by cells of the myeloid lineage including TAMCs, previous reports of immunoPET of CD11b have shown promising results in assessing myeloid cell levels in bone marrow, spleen, and gut in murine models of colitis, melanoma, and systemic inflammation [19-21]. We previously showed that 89Zr-DFO-αCD11b exhibits high tumour uptake and enables visualization of TAMCs in an orthotopic syngeneic mouse glioma using preclinical immunoPET imaging [22], thus demonstrating the potential of 89Zr-αCD11b conjugates to monitor CD11b levels in the central nervous system (CNS) within the glioma TME. The high tumour uptake of 89Zr-DFO-αCD11b, coupled with the critical importance of targeting TAMCs, led us to hypothesize that an anti-CD11b antibody conjugate could be developed into an immuno-theranostic for gliomas. However, while Zr-89 is a positron emitter useful for immunoPET diagnostic agents, it lacks cytotoxicity appropriate for TRT. We therefore selected Lu-177, a clinically relevant cytotoxic β+ emitter, to target TAMCs and TAMC-adjacent cancer cells with TRT.

As bifunctional chelators (BFCs) that can bind both Zr-89 and Lu-177 radionuclides are not available, we report here on the use of a new macrocyclic BFC, Lumi804, developed for that purpose which thus may facilitate translation. Moreover, we evaluated the immunotheranostatic targeting of TAMCs in a murine glioma model using an anti-CD11b antibody (αCD11b) conjugated to Lumi804 for the use of both TRT and immuno-PET. We showed that Lumi804 rapidly and stably chelates both Zr-89 and Lu-177 at ambient temperature, which is an essential feature for radiolabelling a heat-sensitive antibody-based theranostic agent. The data presented here suggest 89Zr/177Lu-Lumi804-αCD11b can be used as novel immuno-theranostic agent for gliomas, which can both improve immunotherapy responses (Lu-177) while visualizing response to therapy (Zr-89).

2. Methods

2.1. Reagents

Chemicals were purchased from Sigma-Aldrich unless otherwise specified. Antibodies for flow cytometry and αCD11b (clone M1/70) were purchased from BioLegend. Immune checkpoint inhibitor antibodies, αPD-1 (clone RMP1-14) and αCTLA-4 (clone 9H10), were purchased from BioXCell. The deferoxamine (DFO) BFC (p-NCS-Bz-DFO) was purchased from CheMatech. Ca-Lumi804-NHS was prepared as previously described. Zirconium-89 oxalate was purchased from the University of Wisconsin. Lutetium-177 chloride was obtained from the Department of Energy National Isotope Development Center. Lumi804 is a proprietary bifunctional chelator patented by Lumiphore.

2.2. Instrumentation and software

Radiochemistry reaction progress and yield were monitored using an Agilent 1260 infinity HPLC (Agilent Technologies) equipped with a SuperoseTM 12 10/300 GL SEC column (GE Healthcare). Immunoreactivity and biodistribution samples were counted using a PerkinElmer 2470 WIZARD2 Automatic Gamma Counter. Flow cytometry studies were performed on a BD LSRII (BD Biosciences), and data were analysed using FlowJo software (version 10.6.1, Flowjo, LLC). T2-weighted MRI scans were acquired with a multi slice RARE sequence using a 7T/30cm Bruker Biospec AVANCE 3 scanner equipped with a 12-cm gradient set, 40mm quadrature birdcage RF resonator with custom mouse bed and ParaVision 6.0.1 (Bruker BioSpin, Billerica MA). Tumour volumes were determined by manual segmentation using ITK-SNAP (version 3.8.0) and DS1 Studio software. PET and T2-weighted MRI scans were co-registered using 3D Slicer (version 4.10.2). PET/CT and SPECT/CT scans were performed on Inveon Preclinical Imaging Stations (Siemens Medical Solutions) with Inveon Research Workstation (IRW) software (version 4.2, Siemens Healthcare, Germany). Complete blood count (CBC) studies were performed using the Abaxis VetScan HMS Hematology Analyzer (Allied Analytic).
2.3. Preparation of Lumi804–αCD11b and DFO–αCD11b conjugates

Aliquots of αCD11b (1.00 mL, 10.1 mg/mL stock) were buffer exchanged using size exclusion columns (SEC) prepared with sodium carbonate buffer (100 mM, pH 9.0) and Sephadex G50 Fine (GE Healthcare), and the combined eluents were quantified by UV-vis spectroscopy (ɛ_{280}=1.33 mL/mg). Aliquots of the resulting αCD11b solution (1 mL, 7.34 mg/mL in sodium carbonate buffer) were treated with Ca-Lumi804-NHS (16.4 µL of 4.48 mM stock solution in anhydrous DME, 1.5 molar equivalents) and shaken at 1000 rpm on a microtube shaker for 1 h. The calcium complex of Lumi804 was used in the conjugation reaction to prevent reaction of the NHS ester with chloride to 10 mL 90 nM Ca-Lumi804-(20 µL 385 (Kd + [conjugate]), where y is mean luminescence (AU), Bmax is maximum luminescence, and Kd is the dissociation constant). Kd was reported with Ca-Lumi804-NHS (16.4 mM stock solution, 20 µL 385 solution (1 mL, 7.34 mg/mL in sodium carbonate buffer) were treated and Lu-177 (56 mM stock solution, 2.5 µL TRF europium settings). Background luminescence was subtracted rinsed thrice (cold HBSS), and time-resolved luminescence was measured for 1 h at room temperature with and without a blocking dose (100 µg αCD11b) and then centrifuged at 1500 rpm for 5 min to separate cell-bound vs. unbound radiolabelled αCD11b. Cells were washed twice with PBS, and the CPM of the supernatant and cell pellet were quantified by gamma counting for each radiolabelled conjugate. The cell-bound: supernatant ratio was used to determine the percentage of bound radiotracer (CPM pellet / CPM supernatant + CPM washes) x 100). Total binding was corrected with non-specific binding to compute immunoreactivity.

2.6. Immunoreactivity assay for 89Zr- and 177Lu-Lumi804–αCD11b antibodies

RAW264.7 cells (ATCC, #TIB-67) were treated with poly-D-lysine (6.4 µg/mL, 50 µL/well, 18 h, Advanced Biomatrix 5049). After 48 h, Fc receptors were blocked with TruStain fcX (100 nM anti-CD16/32, BioLegend, cat#101302, 20 µL/well) and cells were treated with either αCD11b (20 µL/well, 7.5 µM), to determine the non-specific binding, or vehicle control (50 mM HEPES, 150 mM sodium chloride, pH 7.4) and incubated for 30 min at 37°C followed by 15 min at 4°C. After incubation, Eu-Lumi804–αCD11b (prepared by adding 3 µL of 50 mM europium chloride to 10 mL 90 nM Ca-Lumi804–αCD11b) was added at varying concentrations to the plate and incubated (1 h, 4°C). The cells were rinsed thrice (cold HBSS), and time-resolved luminescence was measured on a microplate reader (Perkin Elmer Envision 2104, Wallac TRF europium settings). Background luminescence was subtracted from all wells, and non-specific binding was subtracted from total binding and fit by non-linear regression (y = B_{max} * [conjugate] / (K_d + [conjugate]), where y is mean luminescence (AU), B_{max} is maximum luminescence, and K_d is the dissociation constant). K_d is reported as the mean ± standard deviation using values obtained from three microplates.

2.5. Radiolabelling of DFO- and Lumi804–αCD11b with Zr-89 and Lu-177

89Zr-DFO–αCD11b was prepared and its purity determined by SEC-HPLC as previously described [24]. For radiolabelling of Lumi804–αCD11b, the antibody (10 nmol, 187 µL) was added to Zr-89 oxide (111 MBq, 20 µL) or Lu-177 chloride (111 MBq, 7 µL) in 0.5 M ammonium acetate buffer pH 5.0 (200 µL) with gentisic acid (56 mM stock solution, 2 µL) and incubated for 30 min at 25 °C (Fig. 1). For all radiolabelled antibody conjugates, determination of radiochemical yield was performed by first diluting a small aliquot of the reaction mixture with 50 mM DTPA, pH 5.5, and then monitoring the reaction progress by SEC-HPLC (Superose, PBS, 0.6 mL/min). Radiochemical yield for all conjugates was >99% with a molar activity of 11.1 GBq/µmol. Radiolabelling with high molar activities 22.2 GBq/µmol and 33.3 GBq/µmol was also achieved for 177Lu-Lumi804–αCD11b with >99% radiochemical yield. Serum stability of both 89Zr- or 177Lu-Lumi804–αCD11b (n=2 each) was monitored by radio-SEC-HPLC (Superose 6 Increase (Cytiva), 10/300, 1 mL/min, PBS) over 24, 48, 72, 96 h in mouse serum. For in vivo studies, the radiolabelled antibody conjugates were diluted in the sterile saline before i.v. injections.

2.7. Mouse syngeneic glioma model

The murine immunocompetent glioma 261 (GL261) model was used as previously described [22]. GL261 cells were obtained from the Division of Cancer Treatment and Diagnosis (DCTD) tumour repository of the National Cancer Institute (NCI). Cells were tested for mycoplasma periodically and prior to tumour injections. Intraocular injections of GL261 cells were performed on 5-6-week-old female C57Bl6/J mice (Jackson Labs). Mice received intraocular injection of 1 × 10^5 GL261 cells at coordinates +2.50 mm medial/lateral, +1.50 mm anterior/posterior, and -3.50 mm dorsal/ventral from the bregma.

2.8. SPECT/CT, PET/CT, and biodistribution studies

SPECT/CT was performed at 24, 48 and 72 h post-i.v. injection of 177Lu-Lumi804–αCD11b (100 µg, 7.4 MBq; molar activity: 11.1 GBq/µmol). 177LuCl3 (3.7 MBq) was used for SPECT scanner signal calibration. For PET/CT imaging, mice were injected i.v. with 89Zr-Lumi804–αCD11b (50 µg, 3.7 MBq; molar activity: 11.1 GBq/µmol). For blocking studies, 89Zr-Lumi804–αCD11b was co-injected with 500 µg of αCD11b antibody (unlabelled) to reduce molar activity 10-fold (1.1 GBq/µmol). 89Zr-Lumi804–αCD11b uptake is expressed as SUVmean.

Biodistribution studies of 89Zr-Lumi804–αCD11b and 177Lu-Lumi804–αCD11b were performed immediately after the 72h imaging. Mice were euthanized, major organs were collected, weighed, and the tissue-associated radioactivity was assessed in a gamma counter and expressed as % ID/g. The biodistribution of 89Zr-Lumi804–αCD11b was also compared with that of 89Zr-DFO–αCD11b at 216 h post injection. To determine the biodistribution of 89Zr-DFO–αCD11b, serial PET imaging studies were performed at 1, 24, 48, 72, and 96 hours. Regions of interest were drawn around the tumour and within the contralateral brain, along with muscle, liver, and heart as a surrogate for blood. The SUVmean values were calculated (VivoQuant 2020, Invicro) and plotted against time.

2.9. Flow cytometry

For ex vivo analysis of immune cell populations, randomized day 7 glioma-bearing mice were treated with a single i.v. dose of Lumi804–αCD11b (100 µg) labelled with either 14.8 or 22.2 MBq Lu-177. At
region was harvested, dissected, and processed for flow cytometry analysis as previously described [24]. Dissociated tumour cells were incubated in TruStain fcx at 1 μg/million × 10^4 cells and isolated splenocyte cells at 0.5 μg/10^6 cells in 100 μL staining buffer for 10 min on ice. Suspensions were then stained with 1 μL of each of the following fluorescent-labelled antibodies (Biolegend): CD45-PerCP (clone 30-F11), CD11b-APC (M1/70), Gr1-APC (RB6-8C5), and CD14-PE (sa14-2) or CD45-PerCP/Cy5.5, CD3-APC (17A2), CD4-PE/Cy7 (GK1.5), and CD8-PE (53-617) for 30 min at 4°C. After staining, cells were fixed with Fixation Buffer (BioLegend) and resuspended in Cyto-Last Buffer (BioLegend) for flow cytometry according to the manufacturer’s protocol.

2.10. Targeted radiopharmaceutical therapy (TRT)

For survival studies, at 6 days after tumour cell injections mice were randomized for treatment (day 0). Mice received vehicle (control), two i.v. doses of Lumi804-αCD11b (100 μg) labelled with 7.4 MBq Lu-177 (TRT; days 0 and 7), i.p. injections of 200 μg of each αCTLA-4 and αPD-1 immune checkpoint inhibitors [25] on days 0, 3, 7, and 10, or combination TRT and ICI. MRI was performed one day before treatment to verify solid tumour formation and again on day 16 to quantify tumour volume (DSI Studio software). CBC analysis was performed on day 15 (after treatment). Whole blood (50-500 μL) was harvested by inserting a 4 mm GoldenRod lancet (Fisher Scientific) into the superficial temporal vein and collecting it directly into a potassium-EDTA treated Sarstedt CB300 capillary tube (Fisher Scientific). Samples were mixed and analysed on an Abaxis VetScan HM5 Hematology Analyzer to acquire red blood cell parameters and 3-part white cell differentials. Mice were monitored for symptom-free survival (SFS) and were euthanized if >20% weight loss and neurological symptoms were observed.

3.1. Lumi804-αCD11b is radiolabelled with Zr-89 or Lu-177 at room temperature and demonstrates robust serum stability

An ideal radionuclide combination for antibody-based theranostics is Zr-89 (for PET) and Lu-177 (for TRT) due to relatively long half-lives (t1/2: Zr-89 = 78 h; Lu-177 = 6.7 d), which are compatible with relatively slow antibody localization and clearance rates [26-29]. The recently developed BFC, Lumi804, contains an active ester for conjugation to ε-amino groups of lysine and the N-terminal amines of an antibody. Lumi804 is an octadentate, macrocyclic compound based on four 1-hydroxypropyridin-2-one (1,2-HOPO) coordinating units suitable for binding both Zr-89 and Lu-177 (Fig. 1; left panel).

To synthesize 177Lu[ 89Zr-Lumi804-αCD11b, Ca-Lumi804 was conjugated with αCD11b to generate Ca-Lumi804-αCD11b, followed by trans-chelation of Ca(II) with Zr-89 or Lu-177 to generate the final radioimmuno-theranostic agents (Fig. 1). The molar ratio of Lumi804:αCD11b was found to be 1.3 by UV-vis spectroscopy. We did not observe any degradation or aggregation of Lumi804-αCD11b within 20 days. First, we demonstrated that the photoluminescent Eu-Lumi804-αCD11b agent bound CD11b with high affinity (Kd = 2.6 nM; Supplementary Fig. 1). We then tested the labelling conditions for Ca-Lumi804-αCD11b with both Zr-89 and Lu-177 and achieved facile labelling at room temperature in high radiochemical yields and purity (>98% by size-exclusion chromatography-high performance liquid chromatography (SEC-HPLC)) with 11.1 GBq/μmol molar activity at end-of-synthesis for both agents (Supplementary Fig. 2).

To establish serum stability, we evaluated the stability of 89Zr-Lumi804-αCD11b and 177Lu-Lumi804-αCD11b at 37°C daily for 96 hours. Both agents demonstrated high stability (>98%) with no evidence of radionuclide dissociation, proteolysis, or antibody aggregation at any timepoint evaluated (Supplementary Fig. 3). The immunoreactive fractions of 89Zr-Lumi804-αCD11b and 177Lu-Lumi804-αCD11b were 82% ±4.1% and 69% ±3.1%, respectively. Together, these data demonstrate that Ca-Lumi804-NHS is readily conjugated to the αCD11b antibody, and this conjugate can be radio-labelled with Lu-177 and Zr-89 in high radiochemical yield, purity, molar activity, and immunoreactivity at ambient temperatures.

3.2. 89Zr-Lumi804-αCD11b demonstrates receptor-mediated tumour localization within glioma bearing mice

The GL261 orthotopic C57BL/6j-syngeneic glioma model is widely used for preclinical testing of immunotherapies and exhibits robust TACM accumulation, similar to that of patient gliomas [30-32]. A limitation of many radiometal ion chelates, including Zr-89 bound to the widely used DFO chelator, is poor in vivo stability, which leads to free radionuclide uptake in the bone [29,33]. To evaluate the in vivo stability of Lumi804 with Zr-89, the biodistribution of 89Zr-Lumi804-αCD11b was compared with 89Zr-DFO-αCD11b at 216 h post injection into GL261-glioma bearing mice (Fig. 2). Blood clearance of 89Zr-DFO-αCD11b (0.09 ± 0.03 %ID/g) was significantly different from 89Zr-Lumi804-αCD11b (0.01 ± 0.05 %ID/g; p<0.01, unpaired Student’s t-test), and higher retention in bone was observed for 89Zr-DFO-αCD11b (6.3 ± 2.3 %ID/g) compared with 89Zr-Lumi804-αCD11b (0.50 ± 0.05; p<0.01, unpaired Student’s t-test), putatively due to accumulation of released free Zr-89. 89Zr-DFO-αCD11b accumulation was also significantly higher in the kidney (2.6 ± 1.9 %ID/g) and intestine (1.2 ± 0.5 %ID/g) compared to 89Zr-Lumi804-αCD11b (0.5 ± 0.03 %ID/g; p<0.01 and 0.5 ± 0.02 %ID/g; p<0.01, respectively. Unpaired Student’s t-test was used), suggesting renal and hepatobiliary excretion of 89Zr-DFO-αCD11b. Retention of both agents in the liver was similar. There was greater residual radioactivity in the muscle, heart, thymus, lungs, and ipsilateral and contralateral hemispheres in mice receiving 89Zr-DFO-αCD11b compared with 89Zr-
Lumi804-αCD11b. Overall, these data demonstrate reduced bone uptake of Zr-89 following administration of 89Zr-Lumi804-αCD11b compared to 89Zr-DFO-αCD11b, likely due to the ability of Lumi804 to form a more stable complex with Zr-89 than DFO.

Having established the improved long-term in vivo stability of 89Zr-Lumi804-αCD11b, we then evaluated its utility for immunoPET imaging of TAMCs in glioma-bearing mice at 72 h post tracer injection (Fig. 3a). PET images, co-registered with MRI images with a transparency of 50% PET over MRI [34], verified tracer localization within the tumour (Supplementary Fig. 4). Consistent with our previous report using DFO [24], 89Zr-Lumi804-αCD11b demonstrated significant uptake in the tumour-containing ipsilateral hemisphere (SUVmean = 1.4 ± 0.26) compared with the contralateral hemisphere (SUVmean = 0.21 ± 0.14; p = 0.001, unpaired Student’s t-test; Fig. 3b). To determine the specificity of 89Zr-Lumi804-αCD11b for CD11b, a group of mice received agent that was 10-fold reduced in molar activity (i.e., a greater mass of Lumi804-αCD11b as a competitive block) (Supplementary Fig. 5). The group receiving agent with low molar activity showed significantly less uptake in the tumour ipsilateral hemisphere (SUVmean = 0.35 ± 0.29) compared with the non-blocked group (p = 0.0001, unpaired Student’s t-test; Fig. 3b). Conversely, no significant decrease in SUV was observed in the contralateral hemisphere of mice in the blocked group compared with non-blocked mice (p = 0.68; Fig. 3b). After confirming that the tumour accumulations were measurable with pre-clinical PET, we then determined the biodistribution of 89Zr-Lumi804-αCD11b into dissected major organs at the point of PET imaging (72 h post tracer injection) to corroborate the observed PET data. We confirmed a significantly higher uptake of 89Zr in the tumour ipsilateral hemisphere (3.4 ± 1.0 %ID/g) compared with blocked mice (1.3 ± 0.56 %ID/g; p < 0.01, unpaired Student’s t-test) and compared with the paired contralateral hemisphere (0.4 ± 0.2 %ID/g; p < 0.01, unpaired Student’s t-test; Fig. 3c). Significant differences were also noted in the contralateral hemisphere, blood, and lungs between the non-blocked and the blocked mice. No significant decreases were observed in the bone, muscle, liver, spleen, thymus, or intestine uptake between non-blocked and blocked mice (Fig. 3c).

Due to high and variable blood activity, tumour-to-blood ratios were relatively low (T:B, 0.43 ± 0.32) but tumour-to-muscle ratios were higher (T:M, 5.5 ± 4.29). Our data suggest that tumour uptake of 89Zr-Lumi804-αCD11b is likely reduced by high receptor-specific accumulation in peripheral organs such as blood. Future experiments, including a molar activity study would determine an optimal value to reduce receptor-mediated radiotracer uptake in blood, spleen, and other organs to improve tumour contrast.

**Figure 3d and 3e** show a time course of 89Zr-DFO-αCD11b at 1, 24, 72, and 96 h, demonstrating rapid blood clearance by 48 h (SUVmean = 0.46 ± 0.21) through 96 h (SUVmean = 0.35 ± 0.15). Liver accumulation was relatively high, typical of an IgG-based radiotracer (SUVmean = 0.89 ± 0.17, 72 h and SUVmean = 1.1 ± 0.50, 96 h). 89Zr-Lumi804-αCD11b demonstrated significant uptake within GL261 tumour implants (72 h post injection, SUVmean = 0.52 ± 0.20) compared to contralateral brain (SUVmean = 0.20 ± 0.87; p < 0.05, unpaired Student’s t-test). These values are consistent with the ratio of GL261 tumour accumulation to contralateral brain tissue (tumour: contralateral brain, 2.8 ± 0.6, 72 h). The tumour-to-blood ratios were highest at 72 h post injection (T:B, 1.8 ± 0.62, 72 h) but were moderated due to persistence of tracer in the blood. However, tumour-to-muscle ratios were high (T:M, 6.6 ± 7.0). We note a discrepancy in values in SUVmean of the tumour and tumour:tissues ratios compared with the prior single timepoint PET (Figure 3a-c) experiment. In this experiment tumours grew more rapidly (determined by MRI) and higher interstitial pressures and vascular disruptions may have reduced tracer uptake.

### 3.3. 177Lu-Lumi804-αCD11b has high tumour localization

While Zr-89 is a positron emitter suitable only for immunoPET imaging, Lu-177 is a β+ emitter which can kill target cells by TRT but provides poor in vivo visualization of tumour uptake by SPECT as the γ photons are in low abundance. To demonstrate tumour uptake of 177Lu-Lumi804-αCD11b, SPECT/CT imaging of glioma-bearing mice was performed (7.4 MBq 177Lu-Lumi804-αCD11b). We demonstrated greater uptake of 177Lu-Lumi804-αCD11b in the tumour ipsilateral hemisphere (tumour site) compared with the contralateral hemisphere (tumour site) (SUVmean = 0.70 ± 0.15, 72 h). The tumour-to-blood ratios were high at 72 h post injection (T:B, 1.8 ± 0.62, 72 h) but were moderated due to persistence of tracer in the blood. However, tumour-to-muscle ratios were high (T:M, 6.6 ± 7.0). We note a discrepancy in values in SUVmean of the tumour and tumour:tissues ratios compared with the prior single timepoint PET (Figure 3a-c) experiment. In this experiment tumours grew more rapidly (determined by MRI) and higher interstitial pressures and vascular disruptions may have reduced tracer uptake.
analysis was performed to validate the observed higher uptake of the tracer in the tumour ipsilateral hemisphere (5.2 ± 0.27 %ID/g) relative to the contralateral hemisphere (0.71 ± 0.35 %ID/g; p < 0.0001, paired Student’s t-test; Fig. 4c). Significant uptake was noted in immune cell rich organs including spleen (34.1 ± 5.7 %ID/g) and thymus (10.2 ± 2.8 %ID/g) (Fig. 4c). Blood radioactivity was very low in all mice (0.01 ± 0.01 %ID/g), and consequently very high tumour-to-blood and tumour-to-muscle ratios were also observed (525 ± 270 and 69 ± 33, respectively). We speculate that the reason for the lower blood activity of 177Lu-Lumi804-αCD11b relative to 89Zr-Lumi804-αCD11b is due to eradication of CD11b-positive cells in the serum and subsequent liberation of the tracer from those cells for accumulation in liver and spleen, organs that are relatively resistant to radiation toxicity. The favourable biodistribution, rapid blood clearance, and high tumour ipsilateral hemisphere accumulation then led us to evaluate 177Lu-Lumi804-αCD11b for TRT in glioma-bearing mice.

3.4. αCD11b TRT selectively depletes CD11b+ cell populations and improves response to checkpoint inhibitor immunotherapy therapy

To determine whether a therapeutic dose of 177Lu-Lumi804-αCD11b would decrease TAMCs in the spleen and tumour, glioma-bearing mice were treated with unlabelled αCD11b as a control (100 μg, n=5) or a single dose of Lumi804-αCD11b (100 μg) labelled with either 14.8 (n=4) or 22.2 (n=5) MBq of Lu-177. Five days post-TRT, tumours and splenocytes were isolated and analysed by flow cytometry for a panel of immune cell types. MDSCs and macrophages (CD11b-positive) were significantly reduced in the spleens of mice treated with 177Lu-Lumi804-αCD11b (p < 0.001, p < 0.05, one-way ANOVA with Tukey’s multiple comparisons test; Fig. 5a). Within the tumour, MDSCs were significantly reduced in mice treated with 22.2 MBq 177Lu-Lumi804-αCD11b (p < 0.05, one-way ANOVA with Tukey’s multiple comparisons test), while TAMs trended downward (Fig. 5b).
However, CD11b-negative cell populations (granulocytes, CD4+ helper T-cells [Th], and CD8+ cytotoxic T-lymphocytes [CTLs]) in the spleen and tumour remained unchanged among treatment groups compared with cold antibody controls (Fig. 5c, d). Of note, no signs of therapy-related toxicity were apparent at time of autopsy.

While our data showed that CD11b+ cells were more strongly depleted in lymphoid organs (e.g., spleen) following treatment with $^{177}$Lu-Lumi804-αCD11b, even a moderate reduction in tumour infiltrating TAMCs or peripheral immunosuppression could improve anti-tumour T cell responses and may therefore improve immunotherapy responses. Moreover, circulating monocytes, such as MDSCs and monocyte-derived dendritic-cells (DCs), could affect anti-tumour T cell responses through several mechanisms, including inhibiting T cell priming, maturation, and effector phenotype. Thus, we tested the ability of TRT to improve immunotherapy responses to checkpoint inhibitor immunotherapy (ICI; Fig. 6a), using a fractioned TRT dosing regimen (2 × 7.4 MBq) that we anticipated to maximize safety and therapeutic index. Importantly, we found that glioma-bearing mice

![Figure 5](https://example.com/figure5.png)

**FIGURE 5.** $^{177}$Lu-Lumi804-αCD11b TRT depletes CD11b+ cells in the immune-periphery and within the TME. Glioma-bearing mice were treated with αCD11b (control) or $^{177}$Lu-Lumi804-αCD11b TRT (either 14.8 MBq or 22.2 MBq). Spleen and tumour were harvested 5 days post-TRT, dissociated, and analysed by flow cytometry. (a) Myeloid cell populations, including MDSC, macrophages/TAM, and granulocytes, shown as a % of CD45+ leukocyte-gated cells in spleen, and (b) in tumour. (c) Lymphoid cell populations, including CTL and Th, shown as a % of CD45+ lymphocyte-gated cells in spleen, and (d) in tumour. Data are represented as the mean ± SD. (One-way ANOVA with Tukey’s multiple comparisons test was used to assess significance).

![Figure 6](https://example.com/figure6.png)

**FIGURE 6.** Combination TRT and ICI therapy in syngeneic glioma model reduces tumour volume and improves survival. (a) Schematic of survival study. (b) Kaplan-Meier survival curve of mice treated with vehicle (control, n=10), two doses of Lumi804-αCD11b (100 μg) labeled with 7.4 MBq Lu-177 (TRT, n=4; days 6 and 13 post-tumour cell injection), αPD-1 and αCTLA-4 antibody (ICI, n=8; days 6, 9, 13, and 16 post-tumour cell injection), or both TRT and ICI (n=5). X-axis indicates days post-initiation of therapy. Log-rank survival analysis was used to calculate p-values. (c) Quantification of tumour volume from MRI scans performed 16 days post-initiation of therapy. Unpaired t-test was used to assess significance. (d) CBC performed on superficial temporal vein blood 15 days post-initiation of therapy showed no leukodepletion from TRT. Data are represented as the mean ± SD. (One-way ANOVA with Tukey’s multiple comparisons test was used to assess significance).
receiving a combination of TRT and ICI exhibited significantly pro-
longed survival (median SFS 26 days) compared with control
(p < 0.01, Kaplan Meier analysis with a log-rank test; median SFS 19
days), ICI only (p < 0.05; Kaplan Meier analysis with a log-rank test,
median SFS 20 days), or TRT only (p < 0.05, Kaplan Meier analysis
with a log-rank test; median SFS 19 days; Fig. 6a-b). Tumour volume,
analysed by MRI, was reduced in glioma-bearing mice treated
with combination TRT and ICI compared with control (p < 0.05,
one-way ANOVA with Tukey’s multiple comparisons test; Fig. 6c).
Complete blood count (CBC) analysis demonstrated no
leukodepletion in any treatment group or controls (Fig. 6d, Sup-
plementary Fig. 7). Taken together, our data demonstrate that
\(^{177}\)Lu-Lumi804-\(\alpha\)-CD11b TRT can significantly improve \(\alpha\)CTLA4/
\(\alpha\)PD1 immunotherapy and deplete CD11b+ cells in both the
immune-periphery and within the TME.

4. Discussion

THERANOSTICS, the combination of targeted molecular therapy with
diagnostic imaging, is an evolving field that encompasses nuclear
medicine, radiation oncology, and medical oncology. Theranostics
may be particularly well suited for malignant glioma patients, given
the current lack of effective treatment options and current challenges
in monitoring of therapeutic efficacy (e.g., tumour heterogeneity,
treatment-response heterogeneity, and challenges acquiring tumour
tissue for analysis). Development of theranostic agents for these
tumours requires careful consideration of molecular targets and ther-
apeutic radionuclides.

Radioisotope selection for theranostic pairing of agents must
account for half-life that is suitable for the serum clearance of target-
ing agent, chelator compatibility, positron abundance for PET imag-
ing, therapeutic efficacy, and availability of the radionuclide.
Antibodies are highly effective targeting agents, but the radionuclide
no chelator in widespread use which can rapidly and stably complex
both Zr-89 and Lu-177, two of the most promising radionuclides for
an antibody-based theranostic. Previously reported Lu-177-labeled anti-
body conjugates primarily utilize DOTA or DOTA-derivatives as the
chelator [35] [36], but the slow kinetics of Lu-177 incorporation into
DOTA and DOTA-derivatives remain a significant challenge [37-44].
Furthermore, DOTA analogues do not bind Zr-89 under conditions
that are suitable for antibodies. Previously reported Zr-89-labeled
antibody conjugates primarily utilize DFO as the chelator, but DFO
does not stably chelate Lu-177 and there remain significant stability
issues for Zr-89 labelled DFO agents. Therefore, a single chelator that
readily and stably binds both Zr-89 and Lu-177 under mild conditions
is highly desirable for an effective antibody-based theranostic
because it would facilitate translation of a single Lumi804-antibody
immunoconjugate for FDA approval for both TRT and PET. The combi-
nation of Zr-89 (for PET) and Lu-177 (for TRT) is ideal for antibody-
based theranostics due widespread availability and relatively long
physical half-lives of these isotopes [26-29,45].

Here we utilize a newly developed macrocyclic BFC, Lumi804,
which readily chelates metal ions that prefer 8-coordination, such as
Zr(IV) and Lu(II). Our data demonstrate that Lumi804-\(\alpha\)CD11b is
radiolabelled with either Zr-89 or Lu-177 in 30 minutes at room tem-
perature with high molar activity and high radiochemical yield
(requiring no purification steps). The radiolabelled conjugates
retained high immunoreactivity and demonstrated receptor specific
accumulation in a preclinical animal glioma model. A significant con-
cern of radiometal chelate incorporation into theranostics is bone
deposition of free radiometal due to the instability of the chelate in vivo.
Comparing the \(^{90}\)Zr-Lumi804- and \(^{90}\)Zr-DFO-\(\alpha\)CD11b conjugates
in vivo, the Lumi804 conjugate resulted in significantly less Zr-89
activity in the bone relative to the DFO conjugate, demonstrating the
feasibility of Lumi804-radiolabeled conjugates for in vivo PET imaging
of TAMCs.

Survival times for adults and children with malignant gliomas
remain dismal [30], and an effective theranostic could be a highly
attractive approach for treatment. TAMCs express CD11b on their
surface and represent a major depot of immunosuppressive cells
within gliomas that can directly promote tumour growth, angiogene-
sis, immunosuppression, immunotherapy resistance, and recruit-
ment of immunoregulatory cells [9,10,46]. Our results showing tracer
uptake within brain tumour are consistent with previous observa-
tions that GL261 tumours have a compromised BBB [47]. In the pre-
clinical glioma model, used in this study stereotactic tumour
injection allows rapid reestablishment of BBB integrity due to injec-
tion trauma within a week, and further damage to the BBB is physio-
logic from tumour growth [48,49]. Of note, circulating CD11b+ myeloid-derived cells can migrate through the BBB into the CNS
upon tissue damage [50-52], thereby facilitating transport of tracer-
payload into the TME. Our data show that \(^{177}\)Lu-Lumi804-\(\alpha\)CD11b eradicates circulating CD11b+ cells. As circulating CD11b+ cells are
immunosuppressive and migratory, this reduction may account for
reduced numbers of these cells in the TME, and less immunosuppres-
sion within and outside the TME, leading to enhanced anti-tumour
immunity. Together, these highlight a benefit of CD11b as a target,
independent of BBB integrity. We recently demonstrated that an anti-
body-based immunoPET tracer \(^{89}\)Zr-DFO-\(\alpha\)CD11b had high receptor
specific uptake within murine gliomas [22]. Following treatment
with \(^{177}\)Lu-Lumi804-\(\alpha\)CD11b, we observed reductions of both MDSCs
and CD11b+ macrophages in splenocytes, while CD8+ CTLs which
are important for tumour cell killing, were not depleted. Furthermore,
MDSCs were reduced in tumour samples, although to a lesser extent
than in splenocytes. TAMs were the least affected by \(^{177}\)Lu-Lumi804-
-\(\alpha\)CD11b treatment, potentially due to their innate resistance to radia-
tion, thus explaining their persistence in the TME [53]. Additionally,
as oxygen is critical for TRT effectiveness [54,55], it is possible that
TAMs are residing in hypoxic areas of the tumour. Moreover, lower
radiation exposure in these poorly vascularized tumour regions may
also contribute to greater reduction of splenic macrophages relative
to TAMs [54,55]. An important future direction should explore radio-
sensitizing TAMs to further improve the efficacy of TAMC-targeted
TRT.

While immunotherapies have proven remarkably effective in
some cancer types, a large population of TAMCs contributing to an
immunosuppressive TME has likely contributed to low efficacy of this
treatment in glioma. Our demonstration that targeting TAMCs
with \(^{177}\)Lu-Lumi804-\(\alpha\)CD11b in combination with ICI immunother-
apy significantly prolongs animal survival shows the potential for
TRT of TAMCs. Moreover, our TRT approach did not cause non-spe-
cific peripheral immune cell depletion (determined by CBC analysis)
which is a significant element of safety for TRT. Moreover, immuno-
suppressive CD11b+ myeloid cells, particularly MDSC, have been
observed in many other types of cancers and are also a main target
of immunotherapies [56,57]. Thus, our Lumi804-anti-CD11b could
potentially be used as a theranostic agent in other types of CD11b+
cell-enriched cancers.

A limitation of the current work is the remaining need to perform
dosimetry for our tracers. Future dosimetry studies will allow for
the prediction of off-target toxicity and safety due to tracer accumula-
tion in organs such as the liver, due to IgG clearance, or spleen, given the
presence of CD11b+ cells. Off-target tracer accumulation is a
common problem in TRT, ameliorated through dosimetric calcula-
tions to guide efficacious dosing. Bexxar, an I-I31-labeled antibody
TRT agent for non-Hodgkin lymphoma, demonstrated high liver
and spleen accumulation, similar to our tracer. However, image-
guided dosimetry determined a regimen for safe levels prior to treat-
ment [58,59]. Similarly, we envision image-guided dosimetry with
\(^{89}\)Zr-Lumi804-\(\alpha\)CD11b to set safe levels for \(^{177}\)Lu-Lumi804-\(\alpha\)CD11b.
Advances in imaging technology, particularly in the clinical setting,
such as the incorporation of cadmium-zinc-telluride (CZT) cameras
into modern SPECT are allowing for increased sensitivity and imaging speed to calculate dosimetry based upon $^{177}$Lu-Lumi804-αCD11b, without the need for a separate PET radiotracer [60]. Some promising radiopharmaceuticals for radioimmunotherapy have been developed recently, such as the $^{177}$Lu-labeled αPD-1 antibody [61,62]. similarly, a synergistic effect was noted through TRT with a Lu-177-labeled peptide targeting $\alpha\beta/3$ integrin in combination with anti-PD-1 checkpoint immunotherapy [63].

Here we report on the feasibility of Lumi804-αCD11b as a theranostic agent, with Lumi804 representing the first chelator readily complexing both Zr-89 and Lu-177 for immunopET and TRT, respectively. To our knowledge, this is also the first report of a TRT or theranostic targeting immunosuppressive cells in any solid tumour. Utilizing Lumi804-αCD11 as a theranostic to reduce immunosuppression in combination with immune-stimulating therapies, while also allowing for immunoPET monitoring in glioma patients, may be a promising future clinical approach.

Contributors

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Data sharing statement

Data of this study are available from the corresponding authors upon reasonable request.

Declaration of Competing Interest

DM, DST, and JX are employed and own stock options in Lumiphore, Inc. CJA is on the scientific advisory board and has funding from Lumiphore, Inc. GK receives research support from Lumiphore, Inc. All other authors have no conflict of interest in respect to the work presented in this manuscript.

Acknowledgements

The authors would like to thank Yijen Wu and Nathan Salamacha for the assistance with SPECT/CT imaging, and Kathryn Day for the assistance with SPECT/CT data analysis. The UPMC Hillman Cancer Center (P30 CA047904) supported shared resources were employed in this research, including the Animal Facility and the In Vivo Imaging Facility. Lutetium-177 used in this research was supplied by the U.S. Department of Energy Isotope Program, managed by the Office of Science for Nuclear Physics. This work was funded by Lumiphore Inc., the American Brain Tumour Association (to GK), the Pittsburgh Foundation Copeland Fund (to GK, and to IR), and a generous gift from the Kavalieros Family through Children’s Hospital Foundation. Italy Raphael (IR) was supported by a fellowship from UPMC Children’s Hospital of Pittsburgh and the Walter L. Copeland Foundation.

Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.ebiom.2021.103571.

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