Synthesis and Preclinical Evaluation of a ⁶⁸Ga-Labeled Adnectin, ⁶⁸Ga-BMS-986192, as a PET Agent for Imaging PD-L1 Expression

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Blocking the interaction of the immune checkpoint molecule programmed cell death protein-1 and its ligand, PD-L1, using specific antibodies has been a major breakthrough for immune oncology. Whole-body PD-L1 expression PET imaging may potentially allow for a better prediction of response to programmed cell death protein-1-targeted therapies. Imaging of PD-L1 expression is feasible by PET with the adnectin protein ¹⁸F-BMS-986192. However, radiofluorination of proteins such as BMS-986192 remains complex and labeling yields are low. The goal of this study was therefore the development and preclinical evaluation of a ⁶⁸Ga-labeled adnectin protein (⁶⁸Ga-BMS-986192) to facilitate clinical trials. Methods: ⁶⁸Ga labeling of DOTAconjugated adnectin (BXA-206362) was performed in NaOAc-buffer at pH 5.5 (50°C, 15 min). In vitro stability in human serum at 37°C was analyzed using radio-thin layer chromatography and radio-high-performance liquid chromatography. PD-L1 binding assays were performed using the transduced PD-L1-expressing lymphoma cell line U-698-M and wild-type U-698-M cells as a negative control. Immunohistochemical staining studies, biodistribution studies, and small-animal PET studies of ⁶⁸Ga-BMS-986192 were performed using PD-L1-positive and PD-L1-negative U-698-M-bearing NSG mice. Results: ⁶⁸Ga-BMS-986192 was obtained with quantitative radiochemical yields of more than 97% and with high radiochemical purity. In vitro stability in human serum was at least 95% after 4 h of incubation. High and specific binding of ⁶⁸Ga-BMS-986192 to human PD-L1-expressing cancer cells was confirmed, which closely correlates with the respective PD-L1 expression level determined by flow cytometry and immunohistochemistry staining. In vivo, ⁶⁸Ga-BMS-986192 uptake was high at 1 h after injection in PD-L1-positive tumors (9.0 \pm 2.1 percentage injected dose [%ID]/g) and kidneys (56.9 \pm 9.2 %ID/g), with negligible uptake in other tissues. PD-L1-negative tumors demonstrated only background uptake of radioactivity (0.6 ± 0.1 %ID/g). Coinjection of an excess of unlabeled adnectin reduced tumor uptake of PD-L1 by more than 80%. Conclusion: ⁶⁸Ga-BMS-986192 enables easy radiosynthesis and shows excellent in vitro and in vivo PD-L1-targeting characteristics. The high tumor uptake combined with low background accumulation at early imaging time points demonstrates the feasibility of ⁶⁸Ga-BMS-

986192 for imaging of PD-L1 expression in tumors and is encouraging for further clinical applications of PD-L1 ligands.

Key Words: PD-1/PD-L1 checkpoint inhibitors; PD-L1 PET imaging; ⁶⁸Ga-adnectin; ⁶⁸Ga-BMS-986192; ¹⁸F-BMS-986192

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he immune system is in principle capable of recognizing and destroying cancer cells even in the presence of larger tumor masses and multiple metastases (1). However, cells of the innate and adaptive immune systems are frequently inhibited by molecular pathways that suppress their activation and effector functions and allow tumor cells to escape immune recognition and attack (2). One major checkpoint that is exploited by cancer cells to evade the immune system is the programmed death protein-1 (PD-1) pathway. The negative costimulatory receptor PD-1 is expressed on the surface of activated T cells (3,4). Its ligand, the programmed death protein ligand-1 (PD-L1), a surface protein, is expressed on both antigen-presenting cells and tumor cells of a variety of human cancers (5–7). PD-1/PD-L1 interaction induces downregulation of T-cell activation and allows tumor cells to evade immune recognition and elimination (8).

Inhibition of the PD-1/PD-L1 interaction by antibodies has been a breakthrough for the treatment of several common malignancies such as melanoma, non-small cell lung cancer, renal cell carcinoma, and urothelial cancer (9–12). However, only a subgroup of patients responds to this immune checkpoint inhibitor treatment, and the underlying reasons are not well understood (13–16). Several studies demonstrated that response to PD-1/PD-L1-targeted immunotherapy correlates with PD-L1 expression levels on tumor tissue of diverse malignancies as determined by immunohistochemistry (17). However, the quantitative analysis of PD-L1 expression in tumor tissue is challenging because of the heterogeneous and dynamic expression of such immune checkpoint molecules (18–21). Prediction of response to PD-1-targeted therapy by immunohistochemistry is therefore limited, and in several tumor types, patients benefit from PD-1-targeted therapies in the absence of significant PD-L1 expression in the tumor tissue (17,22,23).

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Imaging of PD-L1 expression can overcome several of the fundamental limitations of PD-L1 immunohistochemistry. First, imaging can provide a 3-dimensional measure of the overall PD-L1 expression of a tumor, whereas only a very small fraction of the tumor tissues can be routinely studied by immunohistochemistry. Second, imaging can assess PD-L1 expression at the whole-body level and therefore allows for studies of the heterogeneity of PD-L1 expression across multiple metastases in an individual patient. Third, imaging is noninvasive and therefore not only can provide information on PD-L1 status before the start of PD-L1 treatment but also monitor changes in PD-L1 expression at multiple times during treatment. Thus, PD-L1 imaging enables new approaches for studying PD1/PD-L1 pathophysiology in patients and may potentially allow for a better prediction of response to PD-1-targeted therapies than immunohistochemistry (24).

It is feasible to image and quantify PD-L1 expression within the tumor in patients with radiolabeled antibodies (25,26). However, antibody-based imaging has several disadvantages such as the long circulatory half-life of antibodies and their slow penetration in the tumor tissue, which necessitates imaging several days after injection. This requires the use of long-lived radioisotopes for radiolabeling, which cause the radiation exposure of antibody PET to be severalfold higher than that of PET with ¹⁸F-FDG.

Adnectins are a family of engineered, target-binding proteins with a size of about 10 kDa, derived from the tenth type III domain of human fibronectin (27). Donnelly et al. synthesized and evaluated an ¹⁸F-labeled anti-PD-L1 adnectin (¹⁸F-BMS-986192) for PET imaging of the PD-L1 expression in vivo (28). ¹⁸F-BMS-986192 showed high in vivo stability and excellent anti-PD-L1-targeting characteristics combined with fast renal clearance, resulting in high-contrast imaging of PD-L1-positive lesions up to hours after injection (28). A first-in-humans study with ¹⁸F-BMS-986192 in advanced non-small cell lung carcinoma demonstrated a correlation of ¹⁸F-adnectin tumor uptake with PD-L1 expression confirmed by immunohistochemistry and response to nivolumab treatment on a lesional basis. The authors illustrated that ¹⁸F-BMS-986192 SUVneak was higher for responding lesions than nonresponding, indicating a correlation of ¹⁸F-BMS-986192 PD-L1 PET signal with PD-L1 expression and tumor-level response (29).

However, the multistep synthesis of ¹⁸F-BMS-986192 with only moderate radiochemical yields is challenging and time-consuming (28,30). To facilitate a broader clinical application of anti-PD-L1 PET imaging, the development of a corresponding ⁶⁸Ga-labeled analog based on the adnectin scaffold seemed the conclusive next step, because of the commercial availability of Food and Drug Administration–approved ⁶⁸Ga/⁶⁸Ga-generators, the possibility of large-scale cyclotron production of ⁶⁸Ga, and the fast and robust ⁶⁸Ga-labeling technique. Herein, we report the synthesis and preclinical evaluation of ⁶⁸Ga-BMS-986192 in terms of PD-L1-affinity, metabolic stability, micro-PET imaging, and in vivo biodistribution in PD-L1–positive and PD-L1–negative xenografts to investigate the feasibility of this tracer for in vivo imaging of PD-L1 expression in tumors.

MATERIALS AND METHODS

Materials

All reagents were obtained from Sigma Aldrich unless otherwise stated. The DOTA-conjugated antihPD-L1 adnectin (BXA-206362) was kindly provided by Bristol-Myers-Squibb Pharmaceutical Research Institute, formulated in phosphate-buffered saline (PBS) buffer (0.9 mg/

mL, pH 7.4), and tested to be endotoxin-free (1.5 EU/mg). 68 Ga was obtained from a 68 Ge/ 68 Ga generator (Galliapharm; Eckert and Ziegler AG).

Synthesis of ⁶⁸Ga-BMS-986192 and Quality Control

The generator was eluted in 1-mL fractions with 0.05 M aqueous HCl (4 mL) containing 170-250 MBq of ⁶⁸GaCl₃. To this eluate, 1 M NaOAc (100 μL, pH 5.5) and 200 μg of DOTA-adnectin (222 μL in PBS) were added, resulting in a labeling solution at pH 5.5. The solution was mixed briefly and incubated for 15 min at 50°C. ⁶⁸Ga-BMS-986192 was purified by gel filtration on a PD-10 column (GE Healthcare). Radiochemical yield and radiochemical purity was analyzed by radio-thin-layer chromatography (radio-TLC) and radio-high-performance liquid chromatography. Radio-TLC was performed using Varian silica impregnated instant TLC paper (Varian Inc.) and 0.1 M aqueous sodium citrate buffer (pH 5.5) as the mobile phase. TLC strips were analyzed on a B-FC-3600 TLC Scanner (Bioscan). Analytic radio-size-exclusion chromatography of ⁶⁸Ga-BMS-986192 was performed using a bioZen SEC-2 (300 \times 4.6 mm) column (Phenomenex LTD.) on a Shimadzu high-performance liquid chromatography system (0.1 M phosphate buffer; pH 6.8; flow, 0.35 mL/min) equipped with a NaI(TI) scintillation detector (5 \times 5 cm [2 \times 2 in]) and a SPD M20A diode array ultraviolet/visible-light detector.

In Vitro Stability of ⁶⁸Ga-BMS-986192 in Human Serum

An in vitro stability study was performed by adding $^{68}\text{Ga-BMS-986192}\,(\sim\!18\,\text{MBq}$ to freshly prepared human serum (1/8, v/v; Seronorm Human; IGZ Instruments AG), followed by incubation at 37°C for up to 4 h. To investigate the stability of $^{68}\text{Ga-BMS-986192}$ in human serum, radio-high-performance liquid chromatography and radio-TLC were performed at 0, 1, 2, 3, and 4 h.

Cell Culture

The PD-L1–negative B-cell lymphoma cell line U-698-M was purchased from ATCC. Cultures were maintained in RPMI medium supplemented with 10% fetal bovine serum and penicillin/streptomycin (100 IU/mL). Cells were grown at 37°C in a humidified atmosphere of 5% CO₂. To stably express PD-L1, the U-698-M cells were retrovirally transduced with a construct containing genes for human PD-L1 and green fluorescent protein, as previously described (31). Quantification of PD-L1 expression on transduced and wild-type U-698-M cells was determined by fluorescence-activated cell sorting (FACS) analysis using an antihuman CD274 antibody (Clone MIH1; BD Bioscience) as previously described (32).

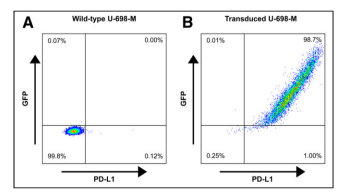


FIGURE 1. Representative dot-plots of expression of green fluorescent protein and PD-L1 in wild-type U-698-M cells (A) and in U-698-M cells transduced by retroviral vector MP71 containing PD-L1 and green fluorescent protein (GFP) separated by P2A element (B).

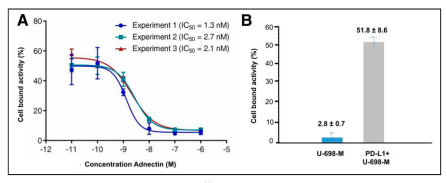


FIGURE 2. Binding affinity and specificity of 68 Ga-BMS-986192 toward PD-L1 in competitive radioligand binding assays. (A) Cell-bound activity in presence of increasing concentrations of cold ligand (BXA-206362). (B) Cell-bound activity in presence of cold ligand (0.1 nM) on PD-L1-positive (gray bar) and PD-L1-negative (blue bar) U-698-M cells. $IC_{50} = half$ -maximal inhibitory concentration.

Assessment of PD-L1 Binding Affinity and Specificity

The binding affinity of 68 Ga-BMS-986192 toward human PD-L1 was determined in a competitive binding experiment using stable transduced U-698-M PD-L1 cells with elevated PD-L1 expression. Briefly, a solution containing a mixture of 68 Ga-labeled (25 μ L) and unlabeled DOTA-conjugated antihPD-L1 adnectin (25 μ L, competitor) with increasing concentrations (10^{-10} – 10^{-6} M) was added to the cells (400,000 cells per vial). After 1 h of incubation at 37°C, the cells were centrifuged at 600g (1,200 rpm; Biofuge 15) for 5 min, the supernatant of each vial was removed, and the cells were thoroughly washed 2 times with 250

MIP Α 0-30 s 0.5-4 min 4-12 min 12-20 min 20-40 min 40-60 min В C Kidneys PET/CT 0-9 %ID/g 1 h p.i. 60 30 PD-L1+ tumor U-698-M PD-L1* U-698-M wild-type Blood 90 Muscle 50 Time (min)

FIGURE 3. Dynamic ⁶⁸Ga-BMS-986192 PET/CT imaging in anesthetized PD-L1-positive and wild-type U-698-M xenograft-bearing NSG mice. (A) Summation maximum-intensity projections (MIP) of different time frames during 90-min acquisition. (B) Time-activity curves for blood pool (heart), kidneys, muscle, and PD-L1-positive tumor derived from dynamic data. (C) Axial PET/CT scan at 1 h after injection (p.i.).

 μL of PBS. The washing medium was combined with the previously removed supernatant, representing the amount of free radioligand. The amount of cell-bound activity (cell pellet), as well as the amount of free radioligand, was measured in a 2470 Wizard² γ -counter (PerkinElmer). Because of the high structural similarity of ⁶⁸Ga-labeled and unlabeled ligand, a nearly identical affinity for PD-L1 is assumed, resulting in homologous competitive binding. Specific binding was confirmed using nontransduced U-689-M cells as a negative control.

U-698-M Tumor Model

All animal experiments were conducted according to the current animal welfare regulations in Germany (DeutschesTierschutzgesetz, approval ROB-55.2–2532.-

Vet_02–15–216). The transduced U-698-M PD-L1–positive and U-698-M wild-type cell lines were detached from the surface of the culture flask using trypsin/ethylenediaminetetraacetic acid (0.05% and 0.02%) in PBS, centrifuged, and resuspended in PBS. Approximately 1×10^7 cells/200 μL of the U-698-M PD-L1–positive cell line were inoculated subcutaneously on the right flank, and U-698-M wild-type cells on the left flank, of 6- to 8-wk-old male NSG mice (Charles River WIGA GmbH). Tumors were grown for 2–3 wk to reach 0.6–1 cm in diameter.

Small-Animal PET Imaging

The mice were anesthetized with isoflurane and intravenously injected via the tail vein with approximately 5–7 MBq (\sim 4.5–5.5 μ g) of ⁶⁸Ga-BMS-986192. In vivo imaging studies were performed using a Siemens Inveon smallanimal PET/CT scanner. Static images were recorded at 1 h and 2 h after injection with an acquisition time of 20 min. For blocking studies, unlabeled BXA-206362 (9 mg/kg) was coinjected with ⁶⁸Ga-BMS-986192. Dynamic imaging was performed after on-bed injection for 1.5 h under isoflurane anesthesia. Reconstruction of the images was performed using a 3-dimensional ordered-subsets expectation maximum algorithm with scanner and attenuation correction. Data analysis was performed using Inveon Workplace software (Siemens). Regions of interest were drawn around areas with increased uptake in transaxial slices for calculation of the average tracer concentration as percentage injected dose (%ID)/g.

Ex Vivo Histology and Immunohistochemistry

Tumor tissues were fixed in 10% neutral-buffered formalin solution for at least 48 h, dehydrated under standard conditions (Leica ASP300S), and embedded in paraffin. Serial 2- μ m-thick sections prepared with a rotary microtome (HM355S; Thermo Fisher Scientific) were collected and subjected to histologic and immunohistochemical analysis. Hematoxylin–eosin staining was performed on deparaffinized sections with eosin and

Mayer hematoxylin according to a standard protocol.

Immunohistochemistry was performed using a Bond RXm system (Leica) with primary antibodies against human PD-L1 (clone 28-8, ab205921, 1:500).

Ex Vivo Biodistribution

About 5–7 MBq of 68 Ga-BMS-986192 (4.5–5.5 μ g) were injected into the tail vein of the U-698-M-PD-L1-positive and U-698-M wild-type tumor-bearing NSG mice under isoflurane anesthesia. The animals were sacrificed at 1 h after injection (n=4) or 2 h after injection (n=4), the organs of interest were dissected, and the activity in the weighed tissues samples was quantified using a γ -counter.

RESULTS

⁶⁸Ga Labeling and In Vitro Stability of ⁶⁸Ga-BMS-986192 in Human Serum

High labeling efficiencies with quantitative radiochemical yields of more than 97% were obtained after 15 min. After purification, moderate specific activities of $11-16~\mathrm{GBq/\mu mol}$ and radiochemical purity of more than 98% was achieved.

Radio-high-performance liquid chromatography of ⁶⁸Ga-BMS-986192 at various time points after mixing with human serum revealed monomeric elution profiles with a minimal radioactive impurity of higher molecular weight (8.97 min), which slightly increased up to 5% after 4 h. Radio-TLC analysis revealed moderate transmetallation (5%) for ⁶⁸Ga-BMS-986192 within 4 h (Supplemental Fig. 1; supplemental materials are available at http://jnm.snmjournals.org).

Characterization of PD-L1 Expression in Transduced U-698-M Cells

Quantification of PD-L1 expression on transduced and wild-type U-698-M cells was determined by FACS analysis. A low PD-L1 expression of approximately 4,000 molecules per cell was observed for wild-type U-698-M cells (Fig. 1). PD-L1 expression was significantly increased by stable transduction of U-698-M cells, with approximately 155,000 PD-L1 molecules per cell.

Competitive Binding Assay of ⁶⁸Ga-BMS-986192 to PD-L1

Binding affinity of 68 Ga-BMS-986192 to human PD-L1 was determined in a competitive radioligand binding experiment using PD-L1–positive and wild-type U-698-M cells. 68 Ga-BMS-986192 showed high affinity toward human PD-L1 with a half-maximal inhibitory concentration of 2.0 \pm 0.6 nM (Fig. 2A). Specific binding was confirmed using nontransduced U-698-M cells as a negative control (Fig. 2B).

PET/CT Imaging

⁶⁸Ga-BMS-986192 showed a rapid clearance from the blood pool and from nontarget tissues, with continuously increasing accumulation in the kidneys over time (Figs. 3A and 3B). ⁶⁸Ga-BMS-986192 uptake in PD-L1–positive tumor was fast within 4 min after injection, and tracer accumulation in PD-L1–expressing tumor tissue increased within 60 min, with high retention over 90 min after injection (Figs. 3A and 3C). No uptake in the PD-L1–negative U-698-M tumor was observed.

Additional static small-animal PET scans were performed with ⁶⁸Ga-BMS-986192 in PD-L1-positive and PD-L1 wild-type tumor-bearing NSG mice at 1 and 2 h after injection (Fig. 4A). ⁶⁸Ga-BMS-986192 showed comparably high tumor uptake in PDL-1-positive tumors after 1 and 2 h after injection. As shown in Figure 4A, in contrast to the high uptake of ⁶⁸Ga-BMS-986192

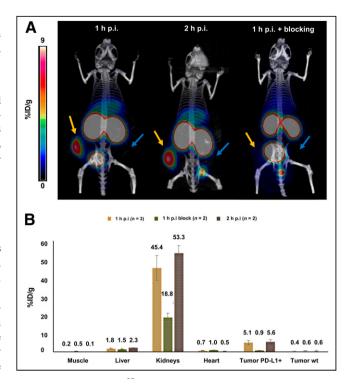


FIGURE 4. (A) Static ⁶⁸Ga-BMS-986192 PET imaging examples in PD-L1-positive (yellow arrows) and wild-type (blue arrows) U-698-M xenograft-bearing NSG mice. Images are maximum-intensity projections (5–6 MBq, $4.5–5.5~\mu g$) obtained at 1 and 2 h after injection (p.i.). Mouse at far right received ⁶⁸Ga-BMS-986192 plus blocking with unlabeled adnectin, 9 mg/kg. (B) ROI quantification of static PET scans (mean %ID/g).

in PD-L1–positive U-698-M tumors, no accumulation was observed in U-698-M wild-type xenografts, confirming PD-L1 specific binding of ⁶⁸Ga-BMS-986192. Additionally, blocking experiments with an excess of unlabeled adnectin (9 mg/kg) demonstrated that ⁶⁸Ga-BMS-986192 uptake is specific and mediated by PD-L1 (Fig. 4B).

Ex Vivo FACS Analysis, Histology, and Immunohistochemistry

Ex vivo FACS analysis of dissected tumors confirmed the correlation of ⁶⁸Ga-BMS-986192 uptake with PD-L1 expression levels in tumor tissue (Figs. 5A and 5B). PD-L1 expression levels in transduced U-698-M tumors was highly increased in comparison to non-transduced U-698-M wild-type xenografts, confirming the generation of a stable PD-L1–expressing tumor cell line in vivo and PD-L1–mediated uptake of ⁶⁸Ga-BMS-986192.

Ex vivo histology and immunohistochemistry of the xenograft tissues also revealed high and homogeneous PD-L1 expression in U-698-M PD-L1–positive tumors, whereas no PD-L1 expression was found in U-698-M wild-type tissues (Fig. 5C). These results confirm the PD-L1–specific uptake of ⁶⁸Ga-BMS-986192 in small-animal PET imaging in vivo and correlate with ex vivo flow cytometry analysis.

Biodistribution

The biodistribution data of ⁶⁸Ga-BMS-986192 in PD-L1-positive-U-698-M and U-698-M wild-type tumor-bearing mice (1 and 2 h after injection) are shown in Figure 6 and Supplemental Table 1. The data reflect the results of small-animal PET/CT imaging. ⁶⁸Ga-BMS-986192 exhibited predominant renal clearance, with only slight uptake in the liver and nontargeted tissues. The significant uptake of ⁶⁸Ga-BMS-986192 in PD-L1-positive tumor

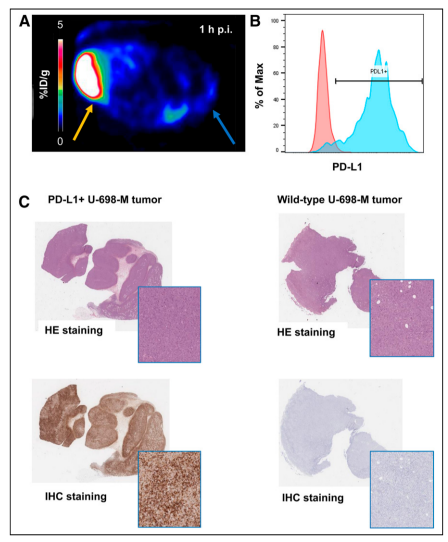


FIGURE 5. (A) Axial ⁶⁸Ga-BMS-986192 PET scan at 1 h after injection in PD-L1–positive (yellow arrow) and wild-type (blue arrow) U-698-M xenograft–bearing NSG mouse. (B) Ex vivo FACS analysis of PD-L1 expression on wild-type (red) and PD-L1 transduced (blue) tumor cells. (C) Ex vivo hematoxylin–eosin (HE) and anti-PD-L1 immunohistochemistry (IHC) staining of PD-L1–positive and wild-type xenograft tissues.

xenografts is highly specific, as accumulation in PD-L1-negative tumors was less than blood-pool activity (Supplemental Table 1).

DISCUSSION

This study revealed that the high-affinity PD-L1-binding adnectin protein BMS-986192, conjugated at the C terminal position with DOTA via maleimide linkage (BXA-206362), can be labeled with ⁶⁸Ga while retaining high target binding affinity (half-maximal inhibitory concentration, 2.0 ± 0.6 nM determined in a competitive radioligand binding assay using PD-L1-positive U-698-M cells). The DOTA-maleimide moiety was conjugated to the cysteine residue engineered into the C terminus of the adnectin (28,30). The same position within the adnectin scaffold has been used previously for site-specific conjugation of the ¹⁸F-labeled prosthetic group of ¹⁸F-BMS-986192 without a significant effect on binding affinity (28). ¹⁹F-BMS-986192, an unmodified and DOTA-conjugated adnectin, has displayed picomolar dissociation constants against human PD-L1 determined by surface plasmon

resonance (28). This indicates that conjugation of small molecules to the C terminus of adnectin is well tolerated, without observable effects on human PD-L1 binding. The ease of tracer preparation and the quantitative radiochemical yields with high radiochemical purity make the synthesis of ⁶⁸Ga-BMS-986192 fully compatible with the everyday workflow in a clinical radiopharmacy and are well suited for automated radiosynthesis via a module or a kit formulation. Furthermore, the pharmacokinetics of ⁶⁸Ga-BMS-986192 match very well the 68-min physical half-life of ⁶⁸Ga, as peak tumor uptake reached a plateau within 60 min after injection. Collectively, these characteristics are highly encouraging for using ⁶⁸Ga-BMS-986192 to image PD-L1 expression in humans.

Small-animal PET and ex vivo biodistribution studies demonstrated excellent targeting of PD-L1-expressing xenografts, whereas tracer uptake was low in tumors without PD-L1 expression on immunohistochemistry (Figs. 4 and 5). The specificity of tracer binding was further confirmed by coinjection of an excess of unlabeled BMS-986192, which decreased tumor uptake of ⁶⁸Ga-BMS-986192 by 80% (Fig. 6). As expected for a 10-kDa protein, ⁶⁸Ga-BMS-986192 showed fast renal clearance from the blood. As observed for other small proteins, peptides and peptide-like molecules, ⁶⁸Ga-BMS-986192 showed significant retention of activity in the kidneys at 1 and 2 h after injection. However, the activity concentration was similar to that of clinically used PET imaging agents such as prostate-specific membrane ligands (33,34). Therefore, we do not expect renal uptake of ⁶⁸Ga-BMS-986192 to limit its clinical use. Injection of an excess of unla-

beled adnectin significantly decreased renal uptake of ⁶⁸Ga-BMS-986192. This is unlikely to be due to blocking of PD-L1 binding since BMS986192 has only low affinity for murine PD-L1 (28). Nevertheless, it indicates that renal uptake of radioactivity is a partially saturable process that could potentially be reduced by injection of proteins or peptides that do not interfere with the binding of ⁶⁸Ga-BMS-986192 to its target. The activity concentration in the kidneys increased from 1 to 2 h after injection, probably reflecting redistribution of the ligand from other tissues to the bloodstream (Fig. 6).

A comparison of the preclinical results obtained for ¹⁸F-BMS986192 and ⁶⁸Ga-BMS-986192 revealed similar specific activities and similar in vitro and in vivo behavior for both tracers (28). Although comparable protein and activity doses of ¹⁸F-BMS986192 and ⁶⁸Ga-BMS-986192 were administered to mice, a direct comparison of the tumor uptake of both radiolabeled ligands is not feasible as different cell lines were used in the 2 studies, but both tracers showed similar pharmacokinetics, with fast renal

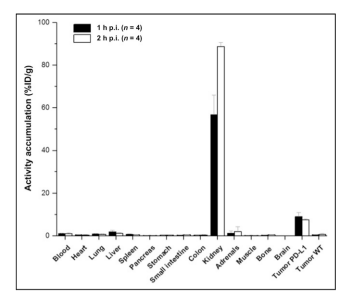


FIGURE 6. Biodistribution data for 68 Ga-BMS-986192 in PD-L1-positive and wild-type (WT) U-698-M xenograft-bearing mice at 1 and 3 h after injection (p.i.). Data are %ID/g (mean \pm SD).

clearance and similarly low background activity in all other organs. This is encouraging for the clinical translation of ⁶⁸Ga-BMS-986192 because PET imaging of PD-L1 expression with ¹⁸F-BMS-986192 has already been shown to be feasible in human pilot studies (29,35).

The following limitation of the study should be noted. ⁶⁸Ga-BMS-986192 exclusively binds to human PD-L1, with no relevant affinity toward its murine counterpart (28). Tumor-to-normal organ uptake ratios may therefore be lower in humans than in our mouse model. Furthermore, we used a PD-L1 transduced cell line to study the binding of ⁶⁸Ga-BMS-986192. The advantages of this approach are that the parent cell line can be used as a negative control to assess the specificity of ligand binding. Furthermore, the stable expression of PD-L1 improves the reproducibility of these results to ensure having a tool for measurement PD-L1 expression of cancer cells, as endogenous cancer cell PD-L1 expression is known to vary significantly over time (36). However, PD-L1 expression levels may be higher in PD-L1 transduced U698-M cells than in human tumors.

CONCLUSION

The novel PD-L1 imaging agent ⁶⁸Ga-BMS-986192 shows similar PD-L1-targeting characteristics and pharmacokinetic properties to ¹⁸F-BMS-986192, which has already been successfully used to image PD-L1 expression in cancer patients. In contrast to ¹⁸F-BMS-986192, which requires a 2-step synthesis with low radiochemical yields, radiolabeling of BMS-986192 with ⁶⁸Ga is a straightforward 1-step process that is easy to automate. ⁶⁸Ga-BMS-986192 therefore has the potential to significantly facilitate preclinical and clinical imaging of PD-L1 expression.

DISCLOSURE

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WO2016086021A1, WO2016086036A2, WO2017/210302, and WO2017/210335. No other potential conflict of interest relevant to this article was reported.

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KEY POINTS

QUESTION: Can adnectins be efficiently radiolabeled with ⁶⁸Ga to allow the in vivo assessment and quantification of PD-L1 expression levels on tumor tissue with PET?

PERTINENT FINDINGS: The expansion of the adnectin-based concept for PD-L1 expression PET imaging to the requirements of ⁶⁸Ga chemistry enables fast and efficient radiolabeling. ⁶⁸Ga-BMS-986192 showed comparable in vitro and in vivo PD-L1-targeting characteristics to its counterpart, ¹⁸F-BMS986192. Both tracers showed favorable pharmacokinetics, with fast renal clearance and similarly low background activity in nontargeted tissues.

IMPLICATIONS FOR PATIENT CARE: The preclinical results are encouraging for the clinical translation of ⁶⁸Ga-BMS-986192 because PET imaging of PD-L1 expression with ¹⁸F-BMS-986192 has already been shown to be feasible in human pilot studies.

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