A theranostic PSMA ligand for PET imaging and retargeting of T cells expressing the universal chimeric antigen receptor UniCAR

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ABSTRACT

Chimeric antigen receptor (CAR) T cells have shown impressive therapeutic potential. Due to the lack of direct control mechanisms, therapy-related adverse reactions including cytokine release- and tumor lysis syndrome can even become life-threatening. In case of target antigen expression on non-malignant cells, CAR T cells can also attack healthy tissues. To overcome such side effects, we have established a modular CAR platform termed UniCAR: UniCAR T cells per se are inert as they recognize a peptide epitope (UniCAR epitope) that is not accessible on the surface of living cells. Bifunctional adapter molecules termed target modules (TM) can cross-link UniCAR T cells with target cells. In the absence of TMs, UniCAR T cells automatically turn off. Until now, all UniCAR TMs were constructed by fusion of the UniCAR epitope to an antibody domain. To open up the wide field of low-molecular-weight compounds for retargeting of UniCAR T cells to tumor cells, and to follow in parallel the progress of UniCAR T cell therapy by PET imaging we challenged the idea to convert a PET tracer into a UniCAR-TM. For proof of concept, we selected the clinically used PET tracer PSMA-11, which binds to the prostate-specific membrane antigen overexpressed in prostate carcinoma. Here we show that fusion of the UniCAR epitope to PSMA-11 results in a low-molecular-weight theranostic compound that can be used for both retargeting of UniCAR T cells to tumor cells, and for non-invasive PET imaging and thus represents a member of a novel class of theranostics.

Introduction

Among tumor diseases prostate cancer (PCa) is still one of the major causes of cancer-related death in men worldwide. Even though primary localized tumors are well manageable, survival benefit of metastatic, castrate-resistant PCa patients is still limited. In order to improve the clinical outcome of these patients, on the one hand, novel innovative therapy options are needed. On the other hand, accurate imaging tools play a critical role to identify appropriate therapeutic measures. Thus, theranostics of PCa based on antibodies (Abs) or small molecules is an uprising and highly promising field.

The prostate-specific membrane antigen (PSMA) represents a valuable molecular target for PCa theranostics. Physiologically, the type II transmembrane glycoprotein PSMA is expressed on healthy prostate and to a lower extent also on other tissues including, e.g. kidneys, intestines, brain, lacrimal and salivary glands. PSMA levels increase up to 1000-fold on about 90% of primary and metastatic PCa cells including bone and lymph node metastases. Most importantly, there is a strong correlation of PSMA expression with high tumor-grade and stage as well as metastatic and hormone-refractory disease.

With the clinical introduction in 2012, the radiotracer Ga-PSMA-11 leveraged PSMA-directed theranostic radiopharmaceuticals. The small peptide ligand PSMA-11 is composed of a Glu-urea-Lys motif as pharmacophore and the Ga-complexing agent benzyl-ethylendiamine-N,N’-diacetic acid (HBED-CC). Among other Glu-ureido-based PSMA ligands, Ga-PSMA-11 has shown to be a widespread, clinically accepted positron
emission tomography (PET) imaging agent especially valuable for detection of lymph node and bone metastases as well as diagnosis of biochemical recurrence with low PSA levels (<2 ng/mL).\textsuperscript{5,6,17} Further modifications finally resulted in the development of theranostic PSMA radiotracers, e.g. PSMA-617\textsuperscript{19,20} and PSMA I&T,\textsuperscript{21} which can be labeled with both diagnostic (e.g. \textsuperscript{68}Ga) and therapeutic (e.g. \textsuperscript{177}Lu or \textsuperscript{225}Ac) radiometals. Depending on the employed radioisotope, these small molecules are successfully utilized in PCa patients for initial diagnosis, re-/staging, and monitoring as well as for salvage radioligand therapy of metastatic, castrate-resistant or progressive disease.\textsuperscript{6,22}

Besides radionuclide therapy, immunotherapeutic approaches based on chimeric antigen receptor (CAR) T cells play an increasingly important role for cancer therapy including PCa.\textsuperscript{23–35} CARs are genetically engineered synthetic receptors consisting of (i) an extracellular binding moiety most commonly derived from tumor-specific monoclonal Abs, (ii) a transmembrane domain, and (iii) intracellular signaling domain(s) taken from activating and optionally costimulatory immune receptors.\textsuperscript{23,26,31} After modification of T cells with CARs, they are able to bind tumor-associated antigen (TAA)-expressing cells independently of their MHC restriction, which finally results in T cell-mediated tumor cell killing.\textsuperscript{23,24,26,31} The pivotal role of CAR T cells for management of tumor diseases is clearly underlined by the approval of two different CD19-specific CAR T cell products in 2017.\textsuperscript{22,36–38} However, the broad clinical application of these living drugs also taught us that precise control mechanisms of CAR T cells after adoptive transfer will be highly important to improve their safety profile. Major side effects are mainly related to tumor lysis and cytokine release syndrome as well as on-target/off-tumor effects that are associated with the expression of the chosen target structure on healthy tissues.\textsuperscript{24,38} In order to rule out lifelong, target-specific off-tumor side effects and to facilitate regulation of adoptively transferred CAR T cells much progress is being undertaken to include additional safety switches [e.g. \textsuperscript{39,40,41,42}]. We therefore established a modular platform technology termed UniCAR, which is a two-component system consisting of UniCAR T cells and target modules (TMs).\textsuperscript{33–36} UniCARs are second-generation CARs, which target the La/SS-B-derived peptide epitope ESB9 (UniCAR epitope).\textsuperscript{53,52} As this epitope is physiologically not accessible on the surface of intact living cells, adoptively transferred UniCAR T cells are in contrast to TAA-specific CAR T cells per se inactive. Retargeting of UniCAR T cells to tumor cells is facilitated by a separate TM comprising a tumor-specific Ab binding moiety (e.g. single-chain fragment variable (scFv) or nanobody) and the UniCAR epitope (Figure 1a). Thus, UniCAR T cells are engaged for tumor cell killing only in the presence of TMs.\textsuperscript{42–50} Based on rapid TM elimination, adverse effects of redirected UniCAR T cells should be easily controlled by stopping TM infusion. After recovery of the patient, TM supply and hence immunotherapy can be restarted. The UniCAR platform technology allows moreover multi-tumor targeting approaches simply by combing UniCAR T cells subsequently or simultaneously with two or more TMs. Since the first presentation of the UniCAR idea in 2014,\textsuperscript{53} several, so far only Ab-based TMs were described including TMs directed against the TAAcs CD33, CD123, prostate stem cell antigen, PSMA, epidermal growth factor receptor, GD2, CD19, or St1.\textsuperscript{43–54}

The aim of the presented study was to develop a novel theranostic PSMA ligand for management of PCa patients that combines the fields of nuclear medicine and CAR T cell technology. Thus, we converted PSMA-11, one of the most commonly used PCa imaging agent, by fusion of the 10-amino acid peptide epitope ESB9 into a TM for the UniCAR system (Figure 1b). The resulting chemically synthesized, low-molecular-weight compound, termed PSMA peptide/ligand tracer TM (PSMA PLT-TM) was functionally characterized with respect to its diagnostic as well as therapeutic properties in vitro and in vivo.

Results

Synthesis and radiolabeling of PSMA PLT-TM

For theranostics of PCa, the novel low-molecular weight compound PSMA PLT-TM was chemically synthesized by fusion of the UniCAR epitope ESB9 to PSMA-11 (Figure 2). The final product (see Figure 2, product 5) was synthesized with purity greater than 98% (overall yield 15–20%), while radiolabeling resulted in one single species \([68\text{Ga}]\text{Ga-PSMA PLT-TM}\) as determined by analytical reversed-phase high-performance liquid chromatography (RP-HPLC) with radiochemical purity \(\geq 96\%\) (Figure S1).

Binding analysis of PSMA PLT-TM

First, an in vitro competitive cell-binding assay was performed for PSMA PLT-TM in order to determine its binding potential for the TAA PSMA in comparison with PSMA-11 using the PSMA-expressing LNCaP cell line. The results are expressed as percentage of cell-bound \([68\text{Ga}]\text{PSMA-10}\) in the presence of increasing

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**Figure 1. Retargeting of UniCAR T cells with the novel PET tracer-based PSMA PLT-TM.** (a) In previous studies, we showed that UniCAR T cells can be retargeted to tumor cells via single-chain fragment variable (scFv)-based target modules (TMs). As schematically shown such scFv-based TMs represent a fusion molecule consisting of the respective scFv and the UniCAR epitope (ESB9). Thus, UniCAR T cells can form an immune complex with the TM, which allows the UniCAR T cell to recognize a specific target on the surface of the tumor cells. (b) As the development of a scFv-based TM for a clinical application is time-consuming and cost-extensive, we wanted to learn whether or not functional TMs can also be constructed from low-molecular-weight molecules: E.g. based on the PSMA-specific ligand PSMA-HBED-CC (PSMA-11 analogue). In order to transform this tracer in a potential TM, we fused the ESB9 epitope during chemical synthesis with PSMA-HBED-CC (see also Figure 2). The resulting compound PSMA peptide/ligand tracer TM (PSMA PLT-TM) cannot only be used for PSMA-PET imaging but also for retargeting of UniCAR T cells to PSMA-positive tumor cells.
concentrations of the non-labeled competitors PSMA PLT-TM and PSMA-11 (Figure 3a). PSMA PLT-TM presented a higher IC$_{50}$ (50% inhibitory concentration) value (IC$_{50}$ = 30.3 ± 1.1 nM) than the reference compound (PSMA-11, IC$_{50}$ = 14.8 ± 1.2 nM).

With regard to UniCAR T cell immunotherapy, we further verified that both binding sites of the bifunctional PSMA PLT-TM are accessible and capable to simultaneously interact with the respective partner domain (Figure 3b). Experiments were conducted in comparison to the previously described Ab-based αPSMA scFv-TM, which was purified from cell culture supernatants of eukaryotic cells using NickelNTA affinity chromatography (Figure S2). As shown by immunofluorescent staining of LNCaP cells, binding of both PSMA PLT-TM and the αPSMA scFv-TM could be detected via the E5B9-tag (Figure 3b). Thus, the UniCAR epitope is still accessible for Ab binding which is the prerequisite for the interaction with UniCAR T cells. Using PC3 cells instead of LNCaP cells a binding of PSMA PLT-TM could hardly be detected (Figure 3b). As the staining of PC3 cells with both a commercial αPSMA mAb and the αPSMA

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**Figure 2.** Synthesis of PSMA PLT-TM. The (tBu protected) PSMA-binding motif and a PEG linker were coupled to either side of the Fe-protected HBED-CC double TFP ester (1) to give (2). The synthesis of the C-terminally modified E5B9 peptide (SKPLEVTDEY-Propargylglycine) (3) was performed manually on a 2CT-Resin under application of standard Fmoc-protocols and cleaved from it with a mixture of TFA/TIPS/H$_2$O (95/2.5/2.5, v/v/v). Then, (2) and (3) reacted via the Cu(I)-catalyzed alkyne-azide cycloaddition (CuAAC), the final product was isolated and the tBu groups were deprotected with TFA. Then, Fe was removed from the complex by applying 1 M HCl on a C-18 SEP-PAK cartridge. The final product was eluted from the cartridge with a mixture of CH$_3$CN/water (7/3, v/v, 20 mL).
scFv-TM also resulted in lower MFI values in comparison to LNCaP cells, this may be due to a low expression of PSMA on PC3 cells. Though the low expression level of PSMA on PC3 is still sufficient for retargeting of UniCAR T cells (as shown below) for technical reasons we selected the LNCaP cell line to estimate and compare $K_D$ values of the TM. For this purpose, increasing amounts of both TMs were incubated with LNCaP cells and the relative median of fluorescence intensity (MFI) values were determined by flow cytometry analysis as described previously.\textsuperscript{45,54} Based on the resulting binding curves (Figure 3c), we calculated $K_D$ values of 27 nM for the PSMA PLT-TM and 34 nM for the αPSMA scFv-TM. According to these data, PSMA PLT-TM and scFv-based αPSMA scFv-TM bind with similar affinity to PSMA.

**Figure 3.** Binding analysis of PSMA PLT-TM. (a) Displacement curves of $^{68}$Ga-PSMA-10 (30 nM) bound to PSMA expressed on LNCaP cells ($10^5$ cells per well). Results are expressed as % specific cell-bound radioactivity after incubation (45 min, RT) with increasing concentrations of non-radiolabeled PSMA-PLT TM or PSMA-11. The IC$_{50}$ values are expressed as mean ± SD. Experiments were performed in quintuplicate. (b) $2 \times 10^5$ LNCaP or PC3 cells were incubated with 20 ng/µL TM. Binding was detected using the mouse anti-ESB9 and PE-labeled goat anti-mouse IgG Abs. In addition, cells were stained with mouse anti-human PSMA Ab/PE as positive control. Histograms show stained cells (blue line) and respective negative controls (black line). Percentage indicate proportion of PSMA$^+$ cells under the marker. (c) For comparison of the binding affinity of the novel PSMA PLT-TM with the scFv-based αPSMA scFv-TM increasing amounts of the respective TM were incubated with LNCaP cells. The binding was estimated by flow cytometry. Relative median of fluorescence intensity (MFI) values were plotted against the concentration. Mean ± SEM of two different experiments is shown. $K_D$ values were calculated from the binding curves.
Killing of PSMA-positive PCa cells by retargeted UniCAR T cells via PSMA PLT-TM occurs in a TM-dependent and target-specific manner with high efficacy similar to the scFv-based αPSMA scFv-TM

In order to evaluate the therapeutic potential of the novel PSMA PLT-TM, we first analyzed its capability to redirect UniCAR T cells for tumor cell killing. Therefore, human T cells from healthy donors were transduced with three lentiviral vectors encoding (I) the signaling construct UniCAR 28/ζ or (II) the UniCAR Stop construct without intracellular signaling domains or (III) the vector control expressing EGFP marker protein. The anti-tumor activity of redirected UniCAR T cells was analyzed by performing chromium release assays (see also Materials and methods) using both the high and low PSMA-expressing cell lines, LNCaP (Figure 4a) and PC3 (Figure 4b). As shown in Figure 4a–b, both target cell lines were attacked by UniCAR 28/ζ T cells at effector-to-target cell (E:T) ratios between 5:1 and 1:5 only in the presence of TMs. Both, the PSMA PLT-TM and the scFv-based αPSMA scFv-TM worked comparably well. The

![Graphs and images](Figure 4. Killing capability of PSMA PLT-TM redirected UniCAR T cells. (a–d) In order to compare the killing capability of UniCAR T cells (UniCAR 28/ζ) armed with either the PSMA PLT-TM or the αPSMA scFv-TM, chromium release assays were performed using either PSMA-positive (a, c) LNCaP or (b, d) PC3 tumor cells. (a, b) As negative controls served UniCAR T cells transduced with an EGFP-expressing vector control (vector control) or UniCAR construct lacking the signaling domains (UniCAR Stop). Data are reported as mean ± SEM for three independent donors. (**p < .0002, *p < .002, *p < .033 with respect to UniCAR 28/ζ w/o TM; 2way ANOVA with Tukey’s multiple comparisons test). (c, d) In order to estimate the range of working concentration for the PSMA PLT-TM PSMA-positive (c) LNCaP cells or (d) PC3 cells were co-cultivated with human T cells modified with the UniCAR signaling construct at an E:T ratio of 5:1. The TM was added at indicated concentrations. Specific lysis of tumor cells was estimated by chromium release assays (see Materials and methods). Data are reported as mean ± SEM for indicated number of independent donors. (**p < .0002, *p < .002 with respect to UniCAR 28/ζ w/o TM; 2way ANOVA with Sidak’s multiple comparisons test). (e) PSMA-positive PC3 cells were incubated with UniCAR T cells in the presence (w/TM) or absence (w/o TM) of the PSMA PLT-TM. Images were taken at the start (10 min), after 60 min, 120 min, 180 min, and 210 min.)
retargeting of UniCAR T cells was TM dependent as tumor cell lysis was minimal and not statistically significant in the absence of TMs. As also expected, T cells modified with the UniCAR Stop vector (Figure 4a,b, UniCAR Stop) or the EGFP encoding vector control (Figure 4a,b, vector control) were neither in the presence nor in the absence of a TM able to induce significant tumor cell lysis. Using increasing amounts of the PSMA PLT-TM, we estimated EC₅₀ (half-maximal effective concentration) values of 0.5 nM for LNCaP (Figure 4c) and 9.2 nM for PC3 cells (Figure 4d).

In order to visualize the time kinetic of tumor cell elimination by redirected UniCAR T cells in more detail, time-lapse microscopy was performed over 4 h. For this purpose, UniCAR T cells were incubated with LNCaP or PC3 cells in the presence or absence of the PSMA PLT-TM. As shown exemplarily for PC3 cells at selected time points, UniCAR T cells armed with the PSMA PLT-TM rapidly bind to and attack tumor cells (Figure 4e, w/TM). In the absence of the TM no stable interaction between UniCAR T cells and target cells were observed (Figure 4e, w/o TM). Similar results were obtained for LNCaP cells (data not shown).

After cross-linkage of PCa cells with UniCAR T cells via the PSMA PLT-TM cytokines are released in a TM- and target-specific manner

An important effector mechanism of redirected UniCAR T cells is the secretion of various pro-inflammatory and growth-promoting cytokines. Thus, we analyzed in a next step the capability of PSMA PLT-TM to induce cytokine secretion in redirected UniCAR T cells. For this purpose, T cells were genetically engineered to express either only EGFP (vector control), UniCAR Stop or UniCAR 28/ζ. Modified T cells were cultivated with or without PC3 or LNCaP tumor cells at an E:T ratio of 5:1 in the presence or absence of the scFv-based aPSMA scFv-TM or the PSMA PLT-TM for 24 h. As shown in Figure S3, cross-linkage of UniCAR T cells with both LNCaP and PC3 cells via the PSMA scFv-TM or the PSMA PLT-TM resulted in release of pro-inflammatory cytokines (TNF and IFN-γ). By using the MACSplex Cytokine 12 Kit, human (see also Materials and methods), we could further show that only five of the 12 cytokines (GM-CSF, IFN-γ, IL-2, IL-4, and TNF-α) were detectable at relevant concentrations after retargeting of UniCAR T cells to PC3 cells via both TMs (Figure 5). Again, these observations substantiate that cytokine secretion occurs in a strictly TM-dependent and target-specific manner.

Figure 5. Analysis of cytokine release from redirected UniCAR T cells. Genetically engineered T cells (vector control, UniCAR Stop or UniCAR 28/ζ) were cultivated with or without PC3 cells (E:T = 5:1), either in the presence or in the absence of scFv-based aPSMA scFv-TM or PSMA PLT-TM. Data show concentrations of selected cytokines after 24 h and are reported as mean ± SD for four independent donors. (**p < .002, *p < .0002 with respect to UniCAR 28/ζ + tumor cells w/o TM; 2way ANOVA with Tukey’s multiple comparisons test).
No significant difference between PSMA PLT-TM and αPSMA scFv-TM could be observed.

**PSMA PLT-TM engages UniCAR T cells for tumor cell killing in vivo**

Recently, we have already successfully shown that the scFv-based αPSMA scFv-TM is able to redirect UniCAR T cells to eliminate PSMA-expressing tumor cells in experimental mice. Now here, we analyzed whether the PSMA PLT-TM is also able to redirect UniCAR T cells for killing of PCa cells in vivo. To reduce the number of experimental mice, the in vivo experiments were limited to LNCaP cells transduced to overexpress firefly luciferase (termed LNCaP-Luc) as target cells (see Materials and methods). 1 × 10⁶ LNCaP-Luc cells were mixed with 0.5 × 10⁶ UniCAR 28/ζ T cells and 3 μM of PSMA PLT-TM per mouse (Figure 6). As "untreated" controls served either 1 × 10⁶ LNCaP-Luc cells alone or 0.5 × 10⁶ UniCAR 28/ζ T cells without any TM. The respective mixture (100 μL) was injected subcutaneously into female NMRI nu/nu mice resulting in three groups of animals each consisting of five mice. Luminescence imaging of anesthetized mice was performed 10 min after i.p. injection of 200 μL of luciferin (15 mg/mL) starting at day zero (day 0) and followed at day two (day 2).

**PET imaging of PSMA-positive tumors using ⁶⁸Ga-labeled PSMA PLT-TM**

In order to show that the modified tracer PSMA PLT-TM can still be used for PET imaging, like PSMA-11, the PSMA PLT-TM was labeled with ⁶⁸Ga (see Materials and methods). As shown in Figure 7a,b, the modified PSMA tracer [⁶⁸Ga]Ga-PSMA PLT-TM specifically enriched within the tumor of experimental mice. The binding of the radiolabeled PET tracer can be blocked with simultaneously injected unlabeled precursor PSMA PLT-TM that results in low molar activity of the radiotracer (Figure 7b, c). As expected the tracer is rapidly eliminated and therefore also found in the kidneys and in the bladder. A detailed biodistribution analysis of the tracer is presented in Table 1 that confirmed the reduction of the ⁶⁸Ga-activity concentration in the tumor by low molar radiotracer concentration. Moreover, administration of the competing drug 2-(phosphonomethyl)-pentandioic acid (2-PMPA) 50 min after radiotracer injection resulted in fast clearance of [⁶⁸Ga]Ga-PSMA PLT-TM from the tumor indicating that [⁶⁸Ga]Ga-PSMA PLT-TM specifically binds to PSMA (Figure 7c).

In order to compare the diagnostic potential of the small molecule PSMA PLT-TM with the previously described αPSMA scFv-TM, we additionally labeled the scFv-based TM with ⁶⁴Cu. PET imaging of LNCaP tumor-bearing mice revealed that maximum molar activity of [⁶⁴Cu]Cu-NODAGA-αPSMA scFv-TM at the tumor site was detected after 24 h (Figure S4). Due to its small size and the extremely short half-life of the complexed radiometal ⁶⁸Ga, maximum molar activity of [⁶⁸Ga]Ga-PSMA PLT-TM was found only in the tumor and kidneys. These data indicate that UniCAR T cells can also be armed in vivo with the PSMA PLT-TM for efficient tumor cell killing in a TM-dependent manner.
Figure 7. (a) PET imaging of the novel $^{68}$Ga-Ga-PSMA PLT-TM is shown as maximum intensity projections (MIP) and orthogonal planes (axial, coronal, sagittal) of two LNCaP tumor-bearing NMRI nu/nu mice. The distribution kinetics of $^{68}$Ga-Ga-PSMA PLT-TM (injected activity 15 MBq, molar activity 18 GBq/µmol, injected peptide amount 0.52 nmol PSMA PLT-TM) after single intravenous injection was measured in a dynamic PET study over 2 h and the images were prepared as integral activity from 60 to 120 min (midframe time 90 min). To demonstrate the specific binding of $^{68}$Ga-Ga-PSMA PLT-TM, PSMA was competitively blocked by simultaneous injection of 86 nmol/animal unlabeled PSMA PLT-TM in one animal (blocked). The green arrows show the tumor of the control animal (SUV$_{\text{max}}$ 1.6 g/mL) and the red arrows indicate blocked tumors (SUV$_{\text{max}}$ 0.06 g/mL). The color scales are in SUV (SUV, 0–1.6) and Hounsfield Units (HU, −1000–0). (b) Blocking of $^{68}$Ga-Ga-PSMA PLT-TM tumor uptake. Estimation of (Bi) the blood and tumor uptake kinetics and (Bii) the corresponding tumor/blood and tumor/muscle ratios in the absence (control) or presence (blocked) of 86 nmol of unlabeled PSMA PLT-TM. Plots show mean ± SEM from experiments of four mice. (c) Kinetics of $^{68}$Ga-Ga-PSMA PLT-TM tumor uptake and blood activity concentration (SUV$_{\text{mean}}$). Comparison of $^{68}$Ga-Ga-PSMA PLT-TM in LNCaP tumor-bearing mice without (control, n = 4) or with simultaneous injection of 86 nmol of non-radiolabeled PSMA PLT-TM (TM simultaneous, n = 4) and 50 min after radiotracer injection, injection of 2 µmol 2-PMPA (50 min p.i. PMPA, n = 2). The activity concentrations were derived from the ROIs over the tumors. Plots are presented as mean ± SEM of n experiments, respectively. (d) PET imaging of a patient with metastasized PCa comparing $^{68}$Ga-Ga-PSMA PLT-TM with $^{68}$Ga-Ga-PSMA-11. (Di) $^{68}$Ga-Ga-PSMA-11 was used for PET of a patient with metastasized PCa. (Dii) The same patient has undergone PET imaging 2 weeks later using $^{68}$Ga-Ga-PSMA PLT-TM. (Diii) PET scan of a healthy volunteer using $^{68}$Ga-Ga-PSMA PLT-TM.
activity of $^{68}$Ga-Ga-PSMA PLT-TM could already be reached after 2 h (Figure S4B).

Finally, PET imaging results of the PSMA PLT-TM were compared with the original PET tracer PSMA-11 in order to further evaluate its potential as a valuable tool for tumor diagnosis in humans. For this purpose, the $^{68}$Ga-labeled PSMA PLT-TM was used for imaging of a tumor patient with metastasized PCa (Figure 7Dii). Two weeks before, the same patient had undergone diagnostic examination via PET imaging using the conventional PET tracer $^{68}$Ga-Ga-PSMA-11 (Figure 7Di). As shown in Figure 7d, the $^{68}$Ga-labeled PSMA PLT-TM (Figure 7Dii) gives almost the same PET imaging result as obtained with the already clinically used $^{68}$Ga-Ga-PSMA-11 (Figure 7Di). Thus, modification of the PET tracer PSMA-11 with the UniCAR epitope had not dramatically changed the capability to use it as a PET tracer. The minor differences seen between the two PET scans of the patient (Figure 7Di-ii) may be caused by the treatment of the patient between the two scanning events but could possibly also be related to modifications made in the imaging agent. The PET scan of a healthy volunteer indicates tracer uptake into salivary glands (Figure 7Diii) which is in agreement with the expression of PSMA in this tissue. As monitored by PET scans of the tumor patient, not only $^{68}$Ga-Ga-PSMA PLT-TM but also the conventional tracer $^{68}$Ga-Ga-PSMA-11 is able to accumulate in this gland tissue (Figure 7Di-ii).

**Discussion**

So far, theranostics in the context of PCa is based on nuclear medicine, namely the combination of small molecules or Abs for PET imaging and radioligand therapy.\(^{5,6,16,17,22}\) However, the here described PSMA PLT-TM goes beyond this strategy and represents for the first time a novel immuno-theranostic agent that allows both PCa imaging and T cell-based immunotherapy. For proof of concept, we selected the clinically used Glu-ureido-based PSMA-binding motif Glu-urea-Lys linked to the chelator HBED-CC. During chemical synthesis of the novel, PSMA PLT-TM tracer 11 amino acids representing the UniCAR epitope and additionally proparglycine for the C-terminal conjugation of the peptide were added to the PSMA-11 analogue by CuAAC to the mini-PEG linker attached to second propionic acid function of the chelator. Our data provide experimental evidence in mice and humans that PSMA-11 could be successfully converted into a UniCAR TM without affecting its use as tracer for PSMA-PET imaging.

From an immunotherapeutic point of view, the PSMA PLT-TM was highly effective as TM for retargeting of UniCAR T cells to kill PSMA-positive PCa cells both in vitro and in vivo. Thereby, UniCAR T cell efficacy and cytokine profile was comparable to the already established αPSMA scFv-TM. Low numbers of UniCAR T cells as well as low concentrations of PSMA PLT-TM were sufficient to engage UniCAR T cells for tumor cell lysis. After 24 h, already the majority of tumor cells was eliminated in vitro. In principle, redirected UniCAR T cells should be able to eradicate all target cells over time as CAR T cells are serial killers.\(^{55}\) This is in line with our in vivo experiment showing complete tumor destruction already after 2 days. Moreover, with EC\(_{50}\) values in a low nM range, killing efficiency of UniCAR T cells armed via the PSMA PLT-TM is comparable to all so far described combinations with Ab-based TMs.\(^{54-50}\) Furthermore, we have shown for the first time that not only Ab fragments but also peptide ligands can be successfully fused to the UniCAR epitope E5B9. In context of solid tumor UniCAR T cell therapy, such small molecules are advantageous due to increased tumor

**Table 1.** Biodistribution of $^{68}$Ga-Ga-PSMA PLT-TM. The biodistribution of $^{68}$Ga-Ga-PSMA PLT-TM and of $^{68}$Ga-Ga-PSMA-11 was determined for three mice inoculated with LNCaP tumors, injected with approximately 10 MBq radiotracer and low (high molar activity) and high (low molar activity) amounts of PSMA PLT-TM. Values are provided as SUV (g/g). BW: body weight; BAT: brown adipose tissue; PSMA PLT-TM: PSMA peptide/ligand tracer target module; WAT: white adipose tissue; Subm.gl.: submandibular glands.

| Precursor | PSMA PLT-TM (high molar activity) | PSMA PLT-TM (low molar activity) | PSMA-11 (high molar activity) |
|-----------|----------------------------------|---------------------------------|-----------------------------|
| | mean | SD | n | mean | SD | n | mean | SD | n |
| Injected activity (MBq) | 9.61 | 0.97 | 7 | 10.38 | 0.52 | 7 | 12.09 | 0.10 | 3 |
| Injected ligand (nmol/kg BW) | 5.06 | 0.34 | 7 | 135.34 | 8.82 | 7 | 6.04 | 0.64 | 3 |
| BW (g) | 29.68 | 0.68 | 7 | 30.60 | 1.55 | 7 | 24.07 | 2.28 | 3 |
| Blood | 0.16 | 0.06 | 7 | 0.12 | 0.05 | 7 | 0.06 | 0.01 | 3 |
| BAT | 0.12 | 0.04 | 7 | 0.06 | 0.01 | 7 | 0.10 | 0.03 | 3 |
| Skin | 0.13 | 0.05 | 7 | 0.09 | 0.01 | 7 | 0.09 | 0.02 | 3 |
| Brain | 0.01 | 0.00 | 7 | 0.01 | 0.00 | 7 | 0.01 | 0.00 | 3 |
| Pancreas | 0.23 | 0.12 | 7 | 0.05 | 0.02 | 7 | 0.29 | 0.14 | 3 |
| Spleen | 0.85 | 0.42 | 7 | 0.10 | 0.05 | 7 | 1.53 | 0.37 | 3 |
| Adrenals | 1.10 | 0.93 | 7 | 0.24 | 0.06 | 7 | 3.11 | 0.77 | 3 |
| Kidneys | 35.82 | 11.37 | 7 | 7.38 | 2.73 | 7 | 50.79 | 3.30 | 3 |
| WAT | 0.28 | 0.09 | 7 | 0.14 | 0.11 | 7 | 0.33 | 0.11 | 3 |
| Muscle | 0.04 | 0.02 | 7 | 0.02 | 0.00 | 7 | 0.04 | 0.00 | 3 |
| Heart | 0.12 | 0.09 | 7 | 0.05 | 0.02 | 7 | 0.09 | 0.04 | 3 |
| Lung | 0.25 | 0.09 | 7 | 0.09 | 0.02 | 7 | 0.45 | 0.14 | 3 |
| Subm. gl. | 0.09 | 0.02 | 7 | 0.05 | 0.00 | 3 | 0.29 | 0.10 | 3 |
| Liver | 0.11 | 0.05 | 7 | 0.07 | 0.02 | 7 | 0.11 | 0.03 | 3 |
| Femur | 0.06 | 0.02 | 7 | 0.04 | 0.01 | 7 | 0.04 | 0.01 | 3 |
| Testes | 0.18 | 0.12 | 7 | 0.05 | 0.01 | 7 | 0.22 | 0.13 | 3 |
| Tumor | 1.22 | 0.37 | 7 | 0.43 | 0.27 | 7 | 2.99 | 0.51 | 3 |
| Ratio tumor/muscle | 39.33 | 17.81 | 7 | 23.86 | 15.95 | 7 | 70.87 | 9.74 | 3 |
permeability and short serum half-lives. From an economic point of view, utilization of a PLT-TM is very favorable for drug development as GMP production is less cost-intensive and less time-consuming in comparison to Ab-based TMs. In a clinical setting, the UniCAR system seems to be a promising approach as it maintains high anti-tumor activity of conventional CARs while providing an additional safety switch. After intravenous injection of the UniCAR T cell product, the patient will receive continuous infusions of appropriate TM doses. For convenient treatment of PCa patients, it would be possible to administer first low doses of the rapidly eliminated PSMA PLT-TM until the majority of tumor cells have been eliminated. This allows a prompt intervention (stopping TM infusion) until the risk of severe acute side effects including tumor lysis syndrome and/or cytokine release syndrome becomes low due to reduced tumor burden. As shown by blocking experiments in experimental mice (see Figure 7c), administration of the competing agent 2-PMPA could also be used as an emergency medicine to rapidly eliminate TMs bound to PSMA-expressing tissues. During the course of treatment, the TM dose can be subsequently increased and adjusted in dependence on the occurrence of adverse reactions. Also, TMs with longer half-life or even a cell factory producing an antibody-based TM in vivo may be applied instead of a continuous TM infusion. In order to reduce the risk of tumor escape variants, a simultaneous application of two or even more TMs would be possible. If tumor cells up-regulate or already express resistant factors such as checkpoint inhibitors, TMs co-delivering stimulatory signals (as already shown for the modular bispecific T cell engager format [e.g. 52]) could help to circumvent this problem. Moreover, UniCAR T cell therapy can be shut down after molecular remission to avoid long-term destruction of healthy tissues, e.g. salivary glands and kidneys showing low PSMA expression. In case of tumor relapse, UniCAR T cells can be easily reactivated by TM infusion including to a different target by switching TM specificity.

With respect to its role as a potential PET radioligand, we have shown that the novel PSMA PLT-TM can be complexed rapidly and efficiently with $^{68}$Ga. The resulting $^{68}$Ga-PSMA PLT-TM was characterized by high purity underlining its suitability as PET radiotracer. It is well known that already small changes in the chemical structure of Glu-ureido-based PSMA ligands, can affect PSMA binding and thus their imaging properties. We could prove that after modification of PSMA-11 with the peptide E5B9 affinity toward PSMA could be maintained and is still in the nM range (IC_{50} value of 30.3 ± 1.1 nM). As required for high image quality in PCa patients, PSMA PLT-TM was also able to rapidly and specifically accumulate at the tumor site both in tumor-bearing mice and in one PCa patient. By performing a first in human study, we could show that it allows the detection of different tumor lesions in men with low imaging background similar to PSMA-11. Thereby, visualization of bone and lymph node metastases is especially important for high-risk PCa patients. Having used the PSMA PLT-TM for imaging of just one patient, these data are limited and need to be confirmed by imaging of further patients at best in the context of the planned phase 1 UniCAR trial. Nonetheless, the PSMA PLT-TM is a promising theranostic agent since we observed only minor differences of human PET/CT scans between $^{68}$Ga-PSMA PLT-TM and $^{68}$Ga-PSMA PLT-TM which may be explained by the fact that the patient was treated between the two measurements. Of course we currently cannot exclude that the observed minor differences, e.g. the more intense staining of the kidneys are caused by alterations in the chemical structure of the tracer molecule. Due to its small size and the short half-life of the radiometal $^{68}$Ga (68 min), the diagnostic window of $^{68}$Ga-PSMA PLT-TM seems to be suitable for fast patient examinations directly after radiotracer injections but might also be limited to a few hours. Thus, utilization of the $^{64}$Cu-labeled aPSMA scFv-TM represents one possibility to extend the diagnostic range of imaging up to 2 days after injection.

Like PSMA-11, the novel PSMA PLT-TM accumulated as expected also in non-cancerous organs, e.g. kidneys and salivary glands which are known to express lower PSMA levels. However, based on the herein presented data it is assumed that the PSMA-11-derived PSMA PLT-TM is still a reliable imaging and immunotherapeutic agent. In case severe side effects against healthy tissues occur, the PSMA PLT-TM can be rapidly removed from the body by both stopping TM supply and by administration of the competing agent 2-PMPA as already mentioned above.

Overall, the results indicate that the PSMA PLT-TM is an attractive candidate for an immunotheranostics in PCa patients. The radiolabeled compound might be a useful tool for initial diagnosis as well as for staging and monitoring of tumor disease during and after therapy. The unlabeled counterpart is instead intended as a TM for cancer immunotherapy with UniCAR T cells. Thereby, the radiolabeled PSMA PLT-TM can be used periodically for target verification and assessment of molecular response to therapy. To allow high-quality imaging and to avoid blockade of PSMA by unlabeled PSMA PLT-TM, TM infusions can briefly be stopped before performing $^{68}$Ga-PSMA PLT-TM PET imaging. In general, such a theranostic approach paves the way for personalized medicine in PCa and is especially important if PSMA-negative tumor escape variants occurs. It will therefore definitely help in decision-making and can guide an appropriate patient-specific treatment: e.g. switching to alternative TMs if TAA-negative tumor escape variants develop or stopping UniCAR T cell therapy in case severe side effects against healthy tissue occur. Local radiation of solid tumor lesions during PET imaging might also be supportive for UniCAR T cell imaging. As planned phase 1 UniCAR trial. Nonetheless, the PSMA PLT-TM is a highly promising theranostic compound that might be useful for retargeting of UniCAR T cells to PCa tissues and for following the progress of tumor treatment via
PET imaging. Moreover, other small molecule-based ligands binding to overexpressed tumor surface molecules may also be converted into such immunotheranostic compounds.

Materials and methods

Cell lines

Recombinantly or natively PSMA-expressing PC3 or LNCaP C4-2B PCa cells (termed here PC3 or LNCaP) were used as target cell lines and cultured in RPMI 1640 complete medium (Biochrom, Berlin, Germany) as described previously.54,59 3T3 cells modified to express the scFv-based aPSMA scFv-TM were kept in DMEM complete medium (Invitrogen, ThermoFisher Scientific, Schwerte, Germany).59 Incubation of all cell lines was done at 37°C and 5% CO₂.

Synthesis of PSMA PLT-TM

Commercially available analytical grade chemicals were used and not further purified. The chemical suppliers were: Sigma-Aldrich (Taufkirchen, Germany) and Merck (Darmstadt, Germany), unless otherwise indicated. Protected amino acids and resins were supplied from Novabiochem (Merck, Darmstadt, Germany) and IRIS Biotech (Marktredwitz, Germany).

The following RP-HPLC systems were used for analysis and purification: Agilent 1100 Series, equipped with a multi-wavelength-detector and Latek P402 (Latek Laborotechnik-Geraete, Eppelheim, Germany) equipped with a HITACHI variable UV detector (absorbance measurement at 214 and 254 nm) and a gamma detector (Bioscan, Washington, USA). Both RP-HPLC systems were equipped with either an analytical Chromolith RP-18e (4.6 x 100 mm; Merck, Darmstadt, Germany) for analysis or a semi-preparative column Chromolith RP-18e (10 x 100 mm; Merck, Darmstadt, Germany) for purifications.

The MALDI-MS Daltonics Microflex (Bruker Daltonics, Bremen, Germany) system was used for mass spectrometry. Full-scan single mass spectra were obtained by scanning m/z = 400–4000.

Synthesis of PSMA PLT-TM is summarized in Figure 2 and described in detail in the following chapters:

Synthesis of 1 and 2

The synthesis of the bis (TFP)ester of N,N'-bis-[2-hydroxy-5(carboxyethyl)benzyl] ethylenediamine- N,N'- diacetic acid, (1), [Fe(HBED-CC)][TFP]₂ iron complex was performed as described previously.60,61 Fifty-six milligrams of [Fe(HBED-CC)][TFP]₂ (0.095 mmol) was dissolved on 300 µL of dry DMF (Sigma-Aldrich, Taukirchen, Germany). After addition of 20 µL of DIPEA (1.2 equiv.) and 41.8 mg of di-tert-butyl (((S)-6-amino-1- (tert-butoxy)-1-oxohexan-2-yl)carbamoyl)-L-glutamate (ABX, Radeberg, Germany) (0.9 equiv.) the solution was stirred for 5 h at room temperature. Finally, 50 µL of 11-Azido-3,6,9-trioxo-aundecan-1-amine (Sigma-Aldrich, Taukirchen, Germany) (2.6 equiv.) was added and the mixture was stirred for 48 h at room temperature. The product (2) was isolated via RP-HPLC by using a linear gradient (A-B): 20–65% B in 16.2 min (Rₑ: 14.1 min); A: 0.1% TFA in H₂O and B: 0.1% TFA in CH₃CN, flow rate 5 mL/min. Maldi-MS (m/z) for [M + H]⁺ (calculated for C₉₀H₉₁N₁₈O₁₈): 1202.4 (1201.65).

Synthesis of C-terminally modified peptide (3)

The synthesis of the C-terminally modified UniCAR peptide (SKPLPEVTDEY-Propargylglycine) (3) was performed manually on a 2CT-Resin (Merck, Darmstadt, Germany) in a 0.15 mmol scale under application of standard Fmoc-protocols. After cleavage of the peptide from the resin with a mixture of TFA/TIS/Water (95/2.5/2.5, v/v/v) the peptide was purified via RP-HPLC with a linear gradient (A-B): 10–37% B in 12.3 min; A: 0.1% TFA in H₂O and B: 0.1% TFA in CH₃CN, flow rate 5 mL/min, Rₑ: 11.8 min. After lyophilization 30 mg of pure product were obtained (yield: 12.5%), Maldi-MS (m/z) for [M + H]⁺ (calculated for C₆₂H₉₃N₁₃O₂₂): 1371.7 (1371.66).

Synthesis of 4 and 5

The coupling of the alkyne bearing peptide (3) with the PSMA-ligand (intermediate 2) was performed via the Cu(I)-catalyzed alkyne-azide cycloaddition (CuAAC). 7.74 mg of 2 (0.0062 mmol) were dissolved together with 13.4 mg of peptide 3 (0.0098 mg; 1.4 equiv.) in 500 µL THF/H₂O. Then, 4.1 mg of CuSO₄ (0.0248 mmol, 4 equiv.) and 4.9 mg of sodium ascorbate (0.0248 mmol, 4 equiv.) were added and the mixture was left to stir for 16 h (RT). After purification by means of RP-HPLC the intermediate tris(tBu)-protected intermediate (4) was isolated and lyophilized. Gradient applied (A-B): 15–54% B in 14.2 min, flow: 5 mL/min, A: 0.1% TFA in H₂O and B: 0.1% TFA in CH₃CN, Rt: 13.4 min, Maldi-MS (m/z) [M + H]⁺ (calculated for C₁₂₀H₁₈₄N₂₂O₄₀) 2574.3 (2573.30). After cleaving the tert.-butyl-protecting groups from the PSMA-binding motif with TFA (3 mL, 3 h, RT) TFA was evaporated. The iron containing intermediate was dissolved in 2% CH₃CN in water (50 mL) and immobilized on a C18-SEP-PAK Cartridge (Waters, Eschborn, Germany). The iron was removed by eluting the C-18 cartridge with 10 mL of 1 M HCl. After washing with water the product 5 (PSMA PLT-TM) was eluted with CH₃CN/Water (7/3, v/v, 20 mL). After lyophilization 10 mg of the product were obtained as a white powder (yield 60%), Maldi-MS (m/z) for [M + H]⁺ (calculated for C₁₅₈H₁₆₀N₂₂O₄₀): 2406.5 (2405.12).

Radiolabeling of the final product with ⁶⁸Ga was accomplished according to previously published methods for ⁶⁸Ga-PSMA-11.60,61 Briefly, a solution of product 5 (1.0 nmol in 0.1 M HEPES buffer, pH 7.5, 100 µL) was added to a mixture of ⁶⁸[Ga]GaCl₃ (80–100 MBq, 40 µL) and HEPES (2.1 M, 10 µL), at pH 4.2 (98°C, 10 min). Radiochemical purity was analyzed by RP-HPLC (Zorbax C18 300SB 9.4 x 250, 4 µm) and gradient H₂O (1)/MeCN (2) containing 0.1% TFA with 5 min to 95% (1), during 10 min to 95% (2), 5 min to 95% (2) and recovery of the system in 5 min to 95% (1) and 5 min to 95% (1). The flow rate was 3 mL/min.

Determination of binding affinity for PSMA in LNCaP cells

Binding affinities (IC₅₀) for the compounds under study were determined by a competitive cell-binding assay with PSMA-
positive LNCaP cells, using known protocols. On a 96-well plate (MultiScreen HTS-DV 0.65 μm) LNCaP cells (10⁵ per well) were incubated with 0.75 nM [⁶⁸Ga]Ga-HBED-CC -[Glu-urea-Lys(Ahx)]₂ (⁶⁸Ga-PSMA-10) (16–20 MBq/nmol) in the presence of 12 different concentrations of PSMA PLT-TM or PSMA-11 ranging from 0 – 5000 nM (100 μL/well). After incubation at RT for 45 min with gentle agitation, the binding buffer was removed using a multiscreen vacuum manifold (Millipore, Billerica, MA). After washing twice with 100 μL and once with 200 μL ice-cold binding buffer, the cell-bound radioactivity was measured with a gamma counter. The 50% inhibitory concentration (IC₅₀) and values were calculated by fitting the data using a nonlinear regression algorithm (GraphPad Software, La Jolla, CA, USA). Experiments were performed in quintuplicate.

**Purification of recombinant antibodies**

The scFv-based anti-PSMA TM (here termed as αPSMA scFv-TM) was produced by a stable 3T3 cell line. For purification of recombinant Abs from cell culture supernatants, Ni-NTA affinity chromatography was performed. In order to evaluate purity and concentration of the αPSMA, scFv-TM SDS-PAGE and immunoblotting were done as already described.

**Generation of UniCAR vectors**

Cloning of the UniCAR 28/ζ vector as well as of the vector control encoding the EGFP marker protein and the UniCAR Stop construct lacking intracellular signaling domains has been described in detail elsewhere.

**Isolation of peripheral blood mononuclear cells and T cell subpopulations**

Isolation and cultivation of primary human T cells was done as already published. For this purpose, either buffy coats (supplied by German Red Cross, Dresden, Germany) or fresh blood of healthy donors was utilized. The study including the consent form was approved by the local ethics committee of the Medical Faculty of the University Hospital Carl Gustav Carus, TU Dresden (EK27022006).

**Lentiviral transduction of T cells**

Transduction of human T cells with lentiviral particles was conducted as described previously [e.g. 33, 45]. In order to ensure that all assays are performed with T cell populations consisting of >90% UniCAR 28/ζ T cells (UniCAR Stop T cells/vector control T cells), they were sorted using a BD FACSria Fusion (BD Biosciences Pharmingen, Heidelberg, Germany). Genetically modified T cells were cultured in RPMI 1640 complete media supplemented with cytokines IL-2 (Proleukin®, Novartis Pharmaceuticals, #2238131), IL-7 (ImmunoTools, #11340075) and IL-15 (ImmunoTools, #11340155). Media was replaced by RPMI 1640 complete media lacking any recombinant cytokines 24 h before experiments started.

**Cytokine release assay**

In a cytokine release assay, 2.5 × 10⁴ UniCAR T cells were incubated with or without 5 × 10⁴ PC3 or LNCaP cells in the presence or absence of the respective TM for 24 h. After centrifugation (360xg, 5 min), cell-free supernatants were collected. Cytokines were subsequently quantified by using the MACSplex Cytokine 12 Kit, human (Miltenyi Biotec GmbH, #130–099-169) or the OptEIA™ Human TNF (BD Biosciences, #555212) and OptEIA™ Human IFN-γ ELISA Kits (BD Biosciences, #555142).

**Flow cytometry analysis**

For TM binding analysis, tumor cells were firstly stained with different concentrations of TM, secondly a mouse anti-E5B9 mAb directed against the epitope E5B9 of the TM, and thirdly a PE-conjugated goat F(ab’)2 fragment anti-mouse IgG (Fcγ) PE (Beckmann Colter, #IM0551) detection Ab. The directly labeled mouse anti-human PSMA Ab/PE (MBL International, #K0142-5) served as positive control. Stained cells were analyzed using a MACSQuant® Analyzer and MACSQuantify™ software (Miltenyi Biotec GmbH, Bergisch Gladbach, Germany). Relative median of fluorescence intensities of the stained cells were used to create a binding curve and calculate the respective Kᵦ value with the GraphPad Prism software (GraphPad Software Inc., La Jolla, CA, USA) [e.g. 46].

**Cytotoxicity assay**

To analyze UniCAR T cell-mediated tumor cell killing standard chromium release assays were performed. For this purpose, genetically modified effector T cells were co-cultivated with 5 × 10⁵ ⁵¹Cr-labeled target cells (PSMA-expressing LNCaP or PC3 cells) in the presence or absence of either the scFv-based αPSMA scFv-TM or the PSMA PLT-TM at indicated E:T ratios. After 24 h or 48 h, specific lysis of tumor cells was determined as described previously.

**Time-lapse microscopy**

For live imaging, PC3 or LNCaP tumor cells were mixed with UniCAR T cells at an E:T ratio of 5:1 without TM or with 30 nM of the PSMA PLT-TM. Cells were cultured in RPMI complete media in an eight well chamber slide (ibidi, Martinsried, Germany). Imaging was performed over 4 h in a humidified atmosphere at 37°C with the all-in-one fluorescence microscope BZ-X700 (Keyence, Japan).

**Immuno-pet imaging, biodistribution analysis, and optical imaging of tumor xenograft models**

All animal experiments were carried out at the Helmholtz-Zentrum Dresden-Rossendorf (HZDR) according to the guidelines of German Regulations for Animal Welfare and have been approved by the Landesdirektion Dresden (24--9165.40–4, 24.9168.21–4/2004–1).
Immuno-PET imaging and biodistribution analysis was carried out in 5–8 week old naive, athymic male nude NMRI-nu Rj:NMRI-Foxn1nu/nu mice (Janvier Labs, France) (named NMRI nu/nu) as previously published. For this purpose, 1 × 10⁶ tumor cells (in PBS) were inoculated subcutaneously in the right hind flank or the right shoulder. Once tumors reached a size of 100–500 mm³, PET imaging or biodistribution analysis were performed. Specific tumor enrichment of ⁶⁸Ga-labeled PSMA PLT-TM radiotracer was blocked by high doses of unlabeled PSMA PLT-TM. Alternatively, specifically bound PSMA PLT-TM radiotracer was removed by intravenous administration of the competing drug 2-PMPA 50 min after radiotracer injection.

For in vivo analysis of tumor cell killing by TM-redirected UniCAR T cells, 1 × 10⁶ LNCaP-Luc cells were mixed with 0.5 × 10⁶ human UniCAR 28/1 T cells in the presence or absence of the PSMA PLT-TM (5 mice per group). As control, one experimental group (n = 5) received tumor cells alone. The cell mixtures (100 µL) were subcutaneously injected into the right thigh of experimental mice. Luminescence imaging (exposure times 1 s, 10 s, and 60 s) was performed as previously described.

**PET imaging of a human patient with metastasized PCA**

Radiolabeling of PSMA PLT-TM: For PET imaging of a patient with metastasized PCA PSMA-11 and PSMA PLT-TM were labeled with ⁶⁸Ga. Labeling of 30 µg PSMA PLT-TM and 20 µg PSMA-11 with ⁶⁸Ga was done similar to other peptides as previously described for ⁶⁸Ga-DOTA-TATE. Samples were taken for determination of identity, radiochemical purity and pH.

Imaging: ⁶⁸Ga-Ga-PSMA-11 and ⁶⁸Ga-Ga-PSMA PLT-TM PET/computed tomography (CT) scans were performed in a man with multiple metastases of PCA with 3-months interval with radionuclide therapy in between. Images were acquired 60 min after injection of 170 MBq ⁶⁸Ga-Ga-PSMA-11 and 65 min after injection of 164 MBq ⁶⁸Ga-Ga-PSMA PLT-TM. PET/CT scans were performed from the base of the skull to mid-thigh on a Biograph 16 HI-REZ scanner (Siemens/CTI, Knoxville, TN). Seven bed positions of 3 min each (axial field of view 15 cm, overlap 1 cm) were acquired. CT imaging was carried out as a low-dose scan without contrast enhancement (120 kV, 10 mAs, 512 × 512 matrix) for attenuation correction only. Images were reconstructed using the ordered subsets expectation maximization algorithm using six iterations and four subsets and CT-based attenuation correction. The study and consent were approved by the local ethics committee of the Technical University (TU) Dresden (EK491122018). A written informed consent was obtained from the patient as well as from the healthy volunteer (male).

**Statistical analysis**

Statistical analysis was performed with GraphPad Prism software version 7 (GraphPad Software Inc., La Jolla, CA, USA) as indicated in the figure legends.

**Acknowledgments**

We thank Julia Lagler, Regina Herrlich and Andrea Suhr for excellent technical assistance.

**Author contributions**

CA, AF, SK, MS, DB, AK, JAS, EP, NM performed experiments and analyzed the data. M. Schäfer synthesized the PSMA PLT-TM. RB and NB performed in vivo studies in experimental mice including the PET imaging. GW and JK performed the human PET analyses. IP, M. Schmitz, GE, KK, CL, M. Schäfer and MB provided critical reagents, know how, and materials. MB, SK, CA, and AF conceived and designed the experiments. MB had the idea to convert the PSMA-PET tracer into a TM, supervised the experiments, and wrote the manuscript.

**Disclosure of Potential Conflicts of Interest**

MB and GE have filed patents related to the UniCAR system. MB and GE are shareholders of the company GEMoAB which owns the IP related to the UniCAR system. GE is owner of the company Cellex PT. The other authors have declared that no competing interest exists.

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