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Prostate-Specific Membrane Antigen Targeted Deep Tumor Penetration of Polymer Nanocarriers

Niranjan Meher, Gary W. Ashley, Anil P. Bidkar, Suchi Dhrona, Cyril Fong, Shaun D. Fontaine, Denis R. Beckford Vera, David M. Wilson, Youngho Seo, Daniel V. Santi, Henry F. VanBrocklin, and Robert R. Flavell*



and heterogenous tumor tissue penetration depending on the tumor phenotype. The performance of nanocarriers *via* specific targeting has the potential to improve imaging contrast and therapeutic efficacy *in vivo* with increased deep tissue penetration. To address this hypothesis, we designed and synthesized prostate cancer-targeting starPEG nanocarriers (40 kDa, 15 nm), [⁸⁹Zr]PEG-(DFB)₃(ACUPA)₁ and [⁸⁹Zr]PEG-(DFB)₁(ACUPA)₃, with one or three prostate-specific membrane antigen (PSMA)-targeting ACUPA ligands. The *in vitro* PSMA binding affinity and *in vivo* pharmacokinetics of the targeted nanocarriers were compared with a nontargeted starPEG, [⁸⁹Zr]PEG-(DFB)₄, in PSMA+ PC3-Pip and PSMA– PC3-Flu cells, and xenografts. Increasing the number of ACUPA ligands improved the *in vitro* binding affinity of PEG-



derived polymers to PC3-Pip cells. While both PSMA-targeted nanocarriers significantly improved tissue penetration in PC3-Pip tumors, the multivalent $[{}^{89}$ Zr]PEG-(DFB)₁(ACUPA)₃ showed a remarkably higher PC3-Pip/blood ratio and background clearance. In contrast, the nontargeted $[{}^{89}$ Zr]PEG-(DFB)₄ showed low EPR-mediated accumulation with poor tumor tissue penetration. Overall, ACUPA conjugated targeted starPEGs significantly improve tumor retention with deep tumor tissue penetration in low EPR PC3-Pip xenografts. These data suggest that PSMA targeting with multivalent ACUPA ligands may be a generally applicable strategy to increase nanocarrier delivery to prostate cancer. These targeted multivalent nanocarriers with high tumor binding and low healthy tissue retention could be employed in imaging and therapeutic applications.

KEYWORDS: positron emission tomography (PET) imaging, prostate-specific membrane antigen (PSMA), polymer nanocarriers, deep tumor penetration, enhanced permeability and retention (EPR) effect

1. INTRODUCTION

In the United States, prostate cancer is the most prevalent noncutaneous cancer in men and the second most common cause of cancer deaths.^{1,2} Glutamate carboxypeptidase II, commonly known as prostate-specific membrane antigen (PSMA), is an overexpressed cell surface enzyme on prostate cancer cells and a valuable clinical biomarker of prostate cancer.³ The recognition site is composed of two pockets with zinc ions, the nonpharmacophore pocket (S1) and the glutamate-sensing pocket (S1'). A urea-based ligand (ACUPA) with free carboxylic acid groups interacts with the S1' pocket of PSMA selectively and has been employed in several single-photon emission computed tomography (SPECT) and positron emission tomography (PET) imaging and theranostic agents for prostate cancer.³⁻⁶ ¹⁷⁷Lu-based PSMA-targeted radiotherapy is currently utilized in the clinic to treat metastatic prostate cancer.⁷⁻⁹ Inspired by promising clinical studies, there has been increased interest in utilizing

PSMA targeting for nanoparticle delivery for imaging and therapy. $^{10-17}$

In parallel with these developments, nanoparticle drugs have found increasing use in medicine.^{18–22} However, the effectiveness of these drugs is not uniform, which may be due in part to incomplete tumor penetration of the large-size macromolecules.²³ Large-size nanoparticles may have nonspecific tumor accumulation because of the enhanced permeability and retention (EPR) effect.^{24–26} The EPR effect is the mechanism by which large-size macromolecules or nanoparticles of >10 nm diameter nontargeted drugs get

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Figure 1. Graphical representation of PSMA-targeted polymer nanocarriers tagged with ⁸⁹Zr radioisotope, demonstrating significantly improved deep tumor penetration in PSMA+ prostate cancer xenograft.

accumulated in tissues with defective vasculature and impaired lymphatic drainage.^{19,27,28} EPR-mediated accumulation of nanoparticles in tumors with abnormal vascular architecture has been well established and a widely accepted strategy for the effective delivery of nanoparticles to tumors.^{19,29} However, the magnitude of the EPR effect is governed by various variables, including nanoparticle size, in vivo pharmacokinetics, vasculature, tumoral microenvironment, and the presence of macrophages.^{27,30,31} Thus, depending on the tumor phenotype, nanoparticles often lack deep tumor tissue penetration, limiting drug delivery and therapeutic efficacy.^{24,30} Furthermore, relatively large-size nanoparticles with strong target binding affinity may suffer from the binding site barrier (BSB) effect, in which the nanoparticles bind to cells peripheral to blood vessels, blocking their further diffusion into the bulk tumors.^{13,23,32,33} Overall, the nanoparticle size and the tumor phenotype play roles in both passive and active tumor uptake, making it challenging to design target-specific nanoparticles with longer retention times and tumor penetration.

Previously, we have explored 4-armed starPEG nanocarriers of 40 kDa conjugated with ⁸⁹Zr chelator deferoxamine B (DFB) as nontargeted PET radiopharmaceuticals in MX-1 and HT-29 tumor models. These studies demonstrated EPRmediated high tumor accumulation and retention (about 10% ID) even after 9 days postinjection.^{25,34,35} In contrast, PET imaging studies of radiolabeled macromolecules and nanoparticles in preclinical human prostate cancer tumor models like CWR22rv1, DU-145, and PC3 demonstrated an "EPR low" effect with reduced tumor accumulation and retention.^{13,30,31} Thus, we hypothesized that conjugating the PSMAtargeted ACUPA ligands to large-size nanocarriers would potentially improve tumor uptake with enhanced retention and tumor tissue penetration, essential and desired characteristics for therapeutic efficacy. In this study, we evaluated three 4armed starPEG_{40kDa} nanocarriers with zero, one, or three copies of PSMA-targeted ACUPA ligands and compared their *in vitro* PSMA binding affinity in PSMA+ PC3-Pip and PSMA- PC3-Flu cell lines. The *in vivo*-targeted uptake and deep tumor uptake of those starPEGs were demonstrated using PET imaging and organ biodistribution studies in a mouse model bearing dual prostate cancer xenografts of PSMA-PC3-Flu and PSMA+ PC3-Pip (Figure 1).

2. METHODS

2.1. Materials and Instrumentations. The 4-armed PEG_{40kDa}-(NH₂)₄ was purchased from SINOPEG (Fujian, China). ⁸⁹Zr-oxalate was procured from three-dimensional (3D) Imaging (Little Rock, AR), p-SCN-Bn-deferoxamine from Macrocyclics (Plano, TX), and deferoxamine mesylate from Sigma-Aldrich (Rockville, MD). RPMI-1640 media, penicillin-streptomycin (P/S) solutions, and fetal bovine serum (FBS) were purchased from Life Technologies (Carlsbad, CA) and Thermo Fisher Scientific (Waltham, MA). Other chemicals (solvents, reagents, and building blocks) were bought from Thermo Fisher Scientific, VWR, or Sigma-Aldrich and used without further processing. $^1\mathrm{H}$ and $^{13}\mathrm{C}$ NMR spectra were recorded on Bruker 400 and 100 MHz NMR spectrometers, respectively. Chemical shifts were shown in parts per million (ppm, δ). High-resolution mass spectrometry (HRMS) was recorded at QB3/Chemistry Mass Spectrometry Facility, University of California, Berkeley.

2.2. Synthesis of starPEG Conjugates. 2.2.1. General. Highperformance liquid chromatography (HPLC) was performed using a $4.6 \times 150 \text{ mm}^2 5 \mu \text{m} 300 \text{ A}$ Phenomenex Jupiter C18 reversed-phase column with a 15 min linear gradient of 0–100% acetonitrile/water/ 0.1% TFA (1.0 mL/min) beginning 2 min after injection, unless otherwise mentioned.

2.2.2. Azido-ACUPA-tBu (1). A solution of 6-azidohexyl succinimidyl carbonate (150 mg, 0.52 mmol, 1 equiv) in 1 mL of DCM was mixed to ACUPA-tBu (283 mg, 0.58 mmol, 1.1 equiv) in 5 mL of DCM containing 176 μ L of DIPEA. The reaction was subjected to magnetic stirring at room temperature for 24 h under argon. After completion, the reaction mixture was evaporated under reduced pressure. The mixture was eluted with 5% methanol in dichloromethane in a silica gel packed column chromatogram to produce pure Azido-ACUPA-tBu (1) (282 mg, 74% yield). ¹H NMR (400 MHz, CDCl₃): d 5.17 (br, 2H), 4.96 (br, 1H), 4.33 (m, 2H), 4.04 (m, 2H), 3.26 (t, 2H), 3.14 (m, 2H), 2.30 (m, 2H), 2.07 (m, 1H), 1.85 (m, 1H), 1.74 (m, 2H), 1.60 (m, 6H), 1.45–1.39 (m, 33H). ¹³C NMR (100 MHz, CDCl₃): d 172.56, 172.44, 157.04, 156.99, 82.22, 81.87, 80.66, 64,74, 53.42, 53.12, 51.49, 40.64, 32.79, 31.72, 29.83, 29.53, 29.05, 28.89, 28.53, 28.21, 28.15, 26.54, 25.62, 22.35. HRMS for $C_{31}H_{57}N_6O_9$ [M + H]⁺ 657.4187 (calcd: 657.4109).

2.2.3. Azido-ACUPA (2). Azido-ACUPA-tBu (1) (450 mg, 0.68 mmol) was dissolved in 3 mL of anhydrous DCM, and 3 mL of trifluoroacetic acid (TFA) was added to it. The mixture was stirred for 16 h at room temperature. On completion, the reaction mixture was evaporated at reduced pressure. The residue was dissolved in 5 mL of H₂O, and the solvent was evaporated at reduced pressure again to remove any remainings of TFA. The residue was dissolved in CHCl₃ and evaporated to yield the product Azido-ACUPA (2). ¹H NMR (400 MHz, CDCl₃): d 4.31 (m, 1H), 4.24 (m, 1H), 4.02 (m, 2H), 3.88 (t, 1H), 3.57 (s, 1H), 3.21 (q, 2H), 3.09 (t, 2H), 2.56 (m, 1H), 2.51 (s, 1H), 2.41 (m, 2H), 2.14 (m, 1H), 1.81 (m, 3H), 1.63 (m, 4H), 1.47–1.35 (m, 10H). ¹³C NMR (100 MHz, CDCl₃): d 176.45, 176.40, 175.82, 160.14, 159.29, 65.66, 58.66, 55.84, 53.50, 52.36, 47.92, 33.20, 31.08, 30.49, 29.83, 28.94, 27.47, 23.84, 18.70. HRMS for $C_{19}H_{31}N_6O_9$ [M – H]⁻ 487.2162 (calcd: 487.2231).

2.2.4. Azido-DFB (3). Deferoxamine mesylate (Sigma, 165 mg, 0.25 mmol, 1.0 equiv) was dissolved in 1 mL of water in a 15 mL Falcon tube and neutralized by 0.5 mL of 1 M Na₂CO₃. A solution of 6azidohexyl succinimidyl carbonate (75 mg, 0.26 mmol, 1.04 equiv) in 1 mL of acetonitrile was mixed, leading to slow formation of a precipitate.36 The precipitate was collected by centrifugation after keeping 3 hours at room temperature, washed successively with water and acetonitrile, and dried under vacuum to afford the product (yield 60%, 110 mg). HPLC: $R_v = 10.2$ mL; purity 96% (evaporative light scattering detector (ELSD) detection). ¹H NMR (400 MHz, DMSOd₆): d 9.63 (br, 1H), 9.58 (br, 2H), 7.76 (t, 2H), 7.02 (t, 1H), 3.91 (t, 2H), 3.45 (m, 6H), 3.31 (m, 4H), 2.99 (m, 4H), 2.94 (m, 2H), 2.57 (t, 4H), 2.27 (t, 4H), 1.50 (m, 10H), 1.41-1.30 (m, 11H), 1.21 (m, 6H). ¹³C NMR (100 MHz, DMSO-d₆): d 171.95, 171.26, 170.11, 156.30, 63.38, 50.53, 47.08, 46.77, 38.40, 29.89, 29.09, 28.80, 28.56, 28.13, 27.55, 26.02, 25.80, 24.92, 23.48, 23.33, 20.32. HRMS for $C_{32}H_{60}N_9O_{10}^+$ [M + H]⁺ 730.4465 (calcd: 730.4385).

2.2.5. Synthesis of PEG-(DFB)₃(ACUPA)₁ (4). A 100 mg/mL solution of PEG-(5HCyO)₃(NH₂)₁ in DMF (1.0 mL, 100 mg, 2.5 mmol, 1.0 equiv) was mixed with a 45 mg/mL solution of Azido-DFB (3) in DMSO (9 mg, 12.3 mmol, 1.5 equiv) and kept at 37 °C for 48 h.²⁵ The mixture was dialyzed against water (SpectraPor 2 membrane, 12-14 kDa cutoff), followed by methanol to remove unconjugated materials and then dried under reduced pressure to give PEG- $(DFB)_3(NH_2)_1$ (S1). The residue was dissolved in 1 mL of acetonitrile, and the insoluble material was removed by filtration. The solution was mixed with 75 mL of a 10 mg/mL solution of 5hydroxycyclooctyne (5HCyO) succinimidyl carbonate in acetonitrile (0.75 mg, 2.8 mmol, 1.5 equiv) and DIPEA (1 mL, 5 mmol, 2.8 equiv) for 30 min at ambient temperature.³⁷ The mixture was added slowly to 10 mL of MTBE, and the precipitated product was collected, washed with MTBE, and dried to give $PEG-(DFB)_3(5HCyO)_1$ (S2), which was advanced without further characterization or purification.

PEG-(DFB)₃(5HCyO)₁ (**S2**) was dissolved in 1 mL of 0.1 M sodium phosphate (pH 7.4) and treated with a 25 mM solution of Azido-ACUPA (**2**) in 0.1 M sodium phosphate, pH 7.4, (200 mL, 5 mmol, 2.8 equiv) for 48 h at 37 °C and then dialyzed (SpectraPor 2 membrane, 12–14 kDa cutoff) against water, followed by methanol, and dried under reduced pressure to afford the product PEG-

 $(DFB)_3(ACUPA)_1$ (4) (yield 45%, 48 mg). ¹H NMR is provided in the Supporting Information.

2.2.6. Synthesis of $PEG-(DFB)_1(ACUPA)_3$ (5). A 100 mg/mL solution of $PEG-(SHCyO)_3(NH_2)_1$ in DMF (1.0 mL, 100 mg, 2.5 mmol, 1.0 equiv) was mixed with a 10 mg/mL solution of *p*-isothiocyanatobenzyl-desferrioxamine B (ITCBz-DFB) in DMSO (Macrocyclics, 2.8 mg, 3.7 mmol, 1.5 equiv) and kept at room temperature for 16 h. The mixture was dialyzed against water (SpectraPor 2 membrane, 12–14 kDa cutoff) to remove unconjugated materials and then dried under reduced pressure. The residue was dissolved in 0.1 M sodium phosphate, pH 7.0, to provide a 75 mg/mL solution of PEG-(SHCyO)_3(DFB)_1 (S3), which was advanced to the next step without further characterization or purification.

A mixture of 75 mg/mL PEG-(5HCyO)₃(DFB)₁ (**S3**) in 0.1 M NaP_i pH 7.0, (850 mL, 1.6 mmol, 4.8 mmol 5HCyO) and a 30 mM solution of Azido-ACUPA (**2**) in 0.1 M NaP_i pH 7.4, (400 mL, 12 mmol, 2.5 equiv) was kept for 48 h at 37 °C, resulting a single new product peak by HPLC. The mixture was dialyzed (SpectraPor 2 membrane, 12–14 kDa cutoff) against water followed by methanol and dried under reduced pressure to provide the product PEG-(DFB)₁(ACUPA)₃ (**5**) (yield 78%, 50 mg). ¹H NMR is provided in the Supporting Information.

2.3. Cell Culture. Unless and otherwise specified, the PSMA– PC3-Flu and PSMA+ PC3-Pip cell lines were cultured in RPMI-1640 medium containing 10% FBS and 1% penicillin/streptomycin (P/S) at 37 °C with 5% CO₂. The cells were obtained from Dr. Martin Pomper's lab, Johns Hopkins University. According to experimental protocols, cells were trypsinized (0.25%) for 2–3 min to detach from the culture flasks for further passage or to seed the cells in suitable multiwell plates to perform cell-binding assays.

2.4. Competition Radioligand Binding Assay. Similar to our prior reported protocol, ⁶⁸Ga-PSMA-11 was produced in a ⁶⁸Ge/⁶⁸Ga generator and used in a competition radioligand binding assay to acquire the IC₅₀ values for the nanocarriers.^{10,38} Briefly, around ~0.185 MBq (2.5 ng) of ⁶⁸Ga-PSMA-11 along with different concentrations (0.01–100,000 nM) of the nonradiolabeled nanocarriers was treated to each well of 96-well plates containing PSMA+ PC3-Pip cells (~20 k cells/wells). After 1 h incubated at room temperature, the radioactive medium was removed, and the cells were washed with PBS twice. The cells were lyzed with sodium hydroxide, and the radioactivity of the lysate in each well was counted in a Hidex γ counter. IC₅₀ was determined by nonlinear regression analysis in Prism software (GraphPad).

2.5. ⁸⁹Zr Radiolabeling of starPEGs. ⁸⁹Zr-oxalate (9 μ L, 192.4 MBq) was neutralized with 9 μ L of Na₂CO₃ (1 M), and 400 μ L of NH₄OAc (1 M) was added to the mixture. To this mixture, ~4 mg of starPEG conjugates in 100 μ L of deionized (DI) water was added and incubated for 1 h at 25 °C. The radiolabeled product was purified using a PD-10 size-exclusion desalting column (Fisher Scientific, Hampton, NH) and eluting with saline solution. Instant thin-layer chromatography (iTLC) was performed using silica gel-impregnated glass microfiber chromatography paper (Neta Scientific, Hainesport, NJ) and developed with 50 mM EDTA solution to confirm radiolabeling purity. The isolated-bound activities were 159.1–185 MBq. Multiple radiolabeling studies were carried out with different amounts of nanocarriers and ⁸⁹Zr and are summarized in Table S1. The radiolabeling yields ranged from 26.27–46.25 MBq/mg.

2.6. *In Vitro* Saturation Binding Assay. PC3-pip cells were seeded in 12-well plates (~100 k cells/well) 24 h prior to testing. Cells were washed with PBS twice, and each well was treated with 1 mL of growth media without/with 10 μ M PSMA-2 (a previously described PSMA inhibitor)^{37–39} and incubated at 37 °C for 1 h. Then, different concentrations (1–1000 nM) of the ⁸⁹Zr-radiolabeled starPEG nanocarriers were treated to the cells and incubated at 37 °C. After 1 h of incubation, the radioactive medium was removed, and cells were washed with PBS. The cells were lyzed with sodium hydroxide, and the radioactivity in each well was analyzed in a Hidex γ counter. The respective nonspecific-bound activities were substracted, and the dissociation constant (K_d) was calculated by nonlinear



Figure 2. Representative chemical structures of ⁸⁹Zn-labeled starPEG nanocarriers to evaluate the PSMA-targeted PET imaging of prostate cancer. [⁸⁹Zr]PEG-(DFB)₄, previously reported nontargeted nanocarrier, was used in this study as a baseline control.²⁵

regression on site-specific binding in Prism Software (GraphPad). These data were further used to show the PSMA binding affinity and blocking of the starPEG nanocarriers at 1 h.

2.7. *In Vitro* Binding and Blocking Assay. PC3-Flu and PC3pip cells were seeded in 24-well plates (~50 k cells/well) 24 h prior to testing. Cells were washed with PBS twice, and each well was treated with 0.5 mL of growth media with/without 10 μ M PSMA-2 (a previously described PSMA inhibitor)^{37–39} and incubated for 1 h at 37 °C. Then, different concentrations (10–100 nM) of the ⁸⁹Zrradiolabeled starPEG nanocarriers were treated to the cells and incubated further at 37 °C. The radioactive medium was discarded at 4 or 24 h time points, and cells were lyzed with NaOH (5 N, 250 μ L) after washing with PBS. The lysate was analyzed in a Hidex γ counter (along with the standard treated activity to calculate the % bound activity).

2.8. *In Vitro* **Membrane-Bound and Internalization Assay.** Four sets of the PC3-pip and PC3-flu cells were seeded in 24-well plates (~50 k cells/well) for 24 h before assay. Each well was treated with 1 μ M ⁸⁹Zr-radiolabeled starPEG nanocarriers and incubated at 37 °C. At each time point (1, 2, 4, and 24 h), one set of PC3-flu and PC3-pip cells was washed with PBS twice and incubated with a mixture of 0.5 mL of ice-cold glycine (50 mM) and NaCl (150 mM) for 5 min at 4 °C. The acid buffer corresponding to the membrane-bound activity was collected. Then, the cells were lyzed with sodium hydroxide (5 N), and the lysate corresponding to the internalized activity was collected. The respective radioactivities were analyzed in a Hidex γ counter (Turku, Finland) along with the standard treated activity of the ⁸⁹Zr-radiolabeled starPEG nanocarriers to calculate the % of membrane-bound and internalized activities.

2.9. Inoculation of Mice with Dual Xenografts. The *in vivo* animal studies were performed under a protocol approved by the UCSF Institutional Animal Care & Use Committee (IACUC). Using precisely similar protocol from our prior report, homozygous (nu/nu) athymic male mice of 5–6 weeks old (Jackson Laboratories or Envigo-Harlan Laboratories, Livermore CA) inoculated with PC3-Pip (left flank, 3 million cells) and PC3-Flu (right flank, 2.5 million cells) dual xenografts.¹⁰ Around 100–200 mm³ tumor size was perceived after 1–2 weeks postinoculation.

2.10. *In Vivo* **PET Imaging and Biodistribution Studies.** Nine days postinoculation, when the tumor size reached 100–200 mm³, the animals were anesthetized using 2% isoflurane, and the respective

⁸⁹Zr-radiolabeled starPEG nanocarriers were administered *via* tail vein (~7.4 MBq in 100 μ L of saline per mouse). The study population included three groups (n = 4 mice per group). The mice were scanned at 24, 48, 168, and 216 h post radiopharmaceutical injection in a μ PET/CT imaging system (Inveon, Siemens Medical Solutions, Malvern, PA). PET data were acquired for 20 min at 24 h and 48 h, 30 min at 168 h, and 40 min at 216 h in list mode, and the manufacturer's two-dimensional (2D) ordered subsets' expectation maximization (OSEM) algorithm was used to reconstruct the data. The imaging data were then normalized to the injected activity to parameterize images to %ID/cc. The imaging data was processed in open-source AMIDE software (http://amide.sourceforge.net/). The tumor-bearing mice were sacrificed at 216 h postinjection of the ⁸⁹Zrradiolabeled starPEG nanocarriers. Blood was collected through a cardiac puncture, and major organs (liver, kidney, spleen, heart, pancreas, lung, brain, femur, muscle, testis, and subcutaneous tumor) were harvested. Blood and major organs were weighed and analyzed in an automated γ counter (Hidex, Turku, Finland). The percent injected dose per gram of tissue (%ID/g) was determined by comparing standard radioactivity.

2.11. Autoradiography. After analyzing the dissected organ samples in the γ counter (Hidex), the tumors were embedded in optimal cutting temperature (OCT) compound and flash frozen on dry ice. Using a microtome, the frozen tumor tissues were sectioned at a thickness of 20 μ m and mounted on iQID charged-particle digital autoradiography imaging systems (QScint Imaging Solutions, LLC, Tucson, AZ). The raw autoradiography data were processed in ImageJ software.

2.12. Statistical Analysis. All data are presented as mean \pm standard deviation in plots. The data were subjected to Student's *t*-test (unpaired, two-tailed, equal variance) for statistical analysis. Differences at the 95% confidence level (P < 0.05) are considered to be statistically significant.

3. RESULTS

3.1. Design, Synthesis, and Radiolabeling of starPEG Conjugates. *3.1.1. Design of starPEG Conjugates.* We hypothesized that adding PSMA-targeting ACUPA ligands would increase tumor accumulation of the nanocarriers. To test this hypothesis, two starPEGs conjugated with one or

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Scheme 1. Synthesis of the Azido Derivatives (a) Azido-ACUPA, (b) Azido-DFB, and Their Conjugation to starPEG Nanocarriers to Produce (c) $PEG-(DFB)_3(ACUPA)_1$ and (d) $PEG-(DFB)_1(ACUPA)_3^a$



^aDetailed synthetic routes with chemical structures have been provided in the Supporting Information. Individual synthetic steps with intermediate structures for the targeted nanocarriers have been presented in Supporting Information Schemes S1 and S2.

three PSMA-targeting ACUPA ligands were designed and synthesized, and their PSMA-targeting ability with tissue penetration was compared with a nontargeted congener without any ACUPA ligands (Figure 2).²⁵ All of the starPEG conjugates were tethered with deferoxamine B (DFB) ligands, which is a robust ⁸⁹Zr chelator commonly used in the development of PET imaging agents.³⁵ The PEG polymer of 40 kDa, 15 nm, a Food and Drug Administration (FDA) approved polymer for safe human use,^{39–41} was used to synthesize the nanocarriers.

3.1.2. Synthesis and Characterization of starPEG Conjugates. PEG-DFB₄ was synthesized by the reaction of 4armed PEG succinimidyl carbonate with DFB mesylate following our previously reported synthetic route.^{25,34} The two starPEG conjugates, PEG-(DFB)₃(ACUPA)₁ (4) and PEG-(DFB)₁(ACUPA)₃ (5), were synthesized using secondgeneration azide click reactions with cyclooctyne (Schemes 1c,d and S1 and S2).⁴² The azide counterparts Azido-ACUPA (2) and Azido-DFB (3) were synthesized following reported procedures (Scheme 1a,b).^{25,43,44} The previously synthesized and reported PEG-(5HCyO)₃(NH₂)₁ was utilized as the starting material for the targeted nanocarriers.²⁵ PEG-(5HCyO)₃(NH₂)₁ conjugate with free amine was treated with Azido-DFB (3) to form PEG-(DFB)₃(NH₂)₁ (S1) and was subsequently treated with 5-hydroxycyclooctyne (5HCyO)-succinimidyl carbonate, followed by Azido-ACUPA (2), to produce PEG-(DFB)₃(ACUPA)₁ (4), whereas PEG-(5HCyO)₃(NH₂)₁ was reacted with isothiocyanatobenzyl-DFB (ITCBz-DFB) to produce PEG-(5HCyO)₃(DFB)₁ (S3), which was further reacted with Azido-ACUPA (2) to



Figure 3. Cell-binding assays with starPEG nanocarriers in PSMA+ PC3-Pip and PSMA– PC3-Flu cell lines demonstrate efficient cell binding and uptake of PSMA-targeted nanocarriers. (a) IC_{50} of nonradiolabeled starPEGs, Azido-ACUPA, and 2-PMPA determined by ⁶⁸Ga-PSMA-11-based *in vitro* competitive radioligand binding assay in PSMA+ PC3-Pip cells (^{NS}*P* > 0.05, Student's *t*-test). (b) K_d measurement of ⁸⁹Zr-labeled starPEGs in the PSMA+ PC3-Pip cell line by a saturation binding assay. (c) Blocking assay of ⁸⁹Zr-labeled starPEGs (100 nM) in PSMA+ PC3-Pip cells using PSMA-2 as the blocking agent at 1 h (%AD = percentage added dose). Detailed blocking assays at different concentrations and incubation times are presented in the Supporting Information (Figures S17–S19). (d) Membrane-bound and internalization assay of the ⁸⁹Zr-labeled starPEGs at 1 h in PSMA+ PC3-Pip cells (%AD = percentage added dose). The membrane-bound activity was collected by 5 min of acid wash with a cold mixture of 50 mM glycine and 150 mM NaCl. Membrane-bound and internalization assays at later time points are presented in the Supporting Information (Figure S20).

yield PEG-(DFB)₁(ACUPA)₃ (5). Detailed synthetic schemes with the respective chemical structures of the polymer conjugates are provided in the supplementary information (Schemes S1 and S2). ¹H and/or ¹³C NMR were recorded for the newly synthesized ligands and starPEG conjugates (Figures S1-S11). The click conjugation of Azido-ACUPA (2) and Azido-DFB (3) to the cyclooctyne counterpart of starPEGs was confirmed by the peaks at 4.33 and 4.01 ppm corresponding to the CH₂ protons close to triazole and carbamate groups in the linkers, respectively (Figure S8). However, none of these peaks at 4.33 and 4.01 ppm were observed in PEG-(DFB)₄ as it does not bear any linker with triazole and carbamate groups. Though the peak position of $PEG-(DFB)_3(ACUPA)_1$ (4) and $PEG-(DFB)_1(ACUPA)_3$ (5) looks identical, different numbers of DFB ligands could be seen by the relative peak integrals at 3.17, 2.80, and 2.49 ppm corresponding to DFB ligands to that of other peaks in the aliphatic region (Figures S5-S7). Moreover, the conjugation of DFB through a p-isothiocyanatobenzyl linker was confirmed by the aromatic proton signal at 7.35 ppm. Additionally, HRMS of the small molecule intermediates, including Azido-ACUPA-tBu, Azido-ACUPA, and Azido-DFB ligands, was also recorded (Figures S12-S14).

3.1.3. Radiolabeling of starPEG Conjugates. The radiolabeling was carried out by treating the respective starPEG conjugates with ⁸⁹Zr-oxalate. The resulting complex was purified in a PD-10 size-exclusion desalting column by eluting with saline solution.³⁴ The yield of the ⁸⁹Zr radiolabeling was 90–95% by iTLC analysis (Figure S15). The isolated yields were 93–98% for PEG-(DFB)₄ (n = 3) and PEG-(DFB)₃(ACUPA)₁ (n = 2) and 81–82% (n = 2) for radiolabeled PEG-(DFB)₁(ACUPA)₃ based on starting ⁸⁹Zr (Table S1). The specific activities ranged from 31.1–44.8 MBq/mg for PEG-(DFB)₄, 31.8–46.3 MBq/mg for PEG-(DFB)₃(ACUPA)₁, and 26.3–39.8 MBq/mg for PEG-(DFB)₁(ACUPA)₃.

3.2. In Vitro Cell-Binding Assay. 3.2.1. Competition Radioligand Binding Assay. The relative binding affinity of the nonradiolabeled starPEG nanocarriers was obtained in a competitive radioligand binding assay using ⁶⁸Ga-PSMA-11 (Figure 3a and Table S2).^{44,45} 2-PMPA and the intermediate Azido-ACUPA were used as positive controls.⁴⁶ As expected, the nontargeted nanocarrier PEG-(DFB)₄ did not show any indication of specific binding in PSMA+ PC3-Pip cells. However, the IC₅₀ values for the targeted nanocarriers were found to be 517 ± 58 nM for PEG-(DFB)₃(ACUPA)₁ and 526



Figure 4. In vivo μ PET/CT imaging. (a) Representation of experimental design for *in vivo* evaluation of the ⁸⁹Zr-labeled starPEGs in mice bearing dual xenografts of PSMA+ PC3-Pip (left flank) and PSMA– PC3-Flu (right flank). (b) Maximum intensity projection (MIP) μ PET/CT, axial μ PET/CT, and axial CT images obtained at 216 h following administration of ⁸⁹Zr-labeled starPEGs reveal high tumor accumulation with low background tissue retention of [⁸⁹Zr]PEG-(DFB)₁(ACUPA)₃ over time. Respective coronal CT and coronal μ PET/CT images are presented in the Supporting Information (Figure S21). ROIs on the heart and tumors are presented in the Supporting Information (Figure S22).

 \pm 1.3 nM for PEG-(DFB)₁(ACUPA)₃ with 95% confidence interval in the range of 359–706 and 404–683 nM, respectively. This demonstrated an insignificant difference (^{NS}P > 0.05) in the competitive binding affinity of these targeted nanocarriers despite the presence of different numbers of PSMA-targeting ACUPA ligands. A similar IC₅₀ was obtained for Azido-ACUPA (349.6 nM) and 2-PMPA (393.7 nM) with 95% confidence interval in the range of 290–420 and 182–482 nM, respectively.

3.2.2. Saturation Binding Assay. The binding affinity of the ⁸⁹Zr-labeled nanocarriers was further evaluated in a saturation binding assay in PSMA+ PC3-Pip cells that demonstrated ~25-fold lower dissociation constant for $[^{89}Zr]PEG-(DFB)_1(ACUPA)_3$ ($K_d = 30.96$ nM) with three copies of ACUPA ligands compared to $[^{89}Zr]PEG-(DFB)_3(ACUPA)_1$ ($K_d = 790.6$ nM) with only one copy of ACUPA ligands (Figure 3b and Table S3). In contrast, no indication of specific

binding was witnessed in the nontargeted $[{\rm ^{89}Zr}]PEG\text{-}(DFB)_4$ nanocarrier.

3.2.3. Blocking Assay. Further, a binding and blocking assay was performed using PSMA-2 as the blocking agent at 1, 4, and 24 h time points and different nanocarrier concentrations (Figures 3c and S16-S19).⁴⁷⁻⁴⁹ The targeted nanocarriers, [⁸⁹Zr]PEG-(DFB)₃(ACUPA)₁ and [⁸⁹Zr]PEG-(DFB)₁(ACUPA)₃, demonstrated higher uptake selectively in PSMA+ PC3-Pip cells, and the uptake was significantly reduced in the presence of the known PSMA binder PSMA-2.⁴⁷ In comparison, no specific uptake of the nanocarriers was perceived in PSMA- PC3-Flu cells. Noticeably, at lower probe concentrations (10 and 100 nM), [⁸⁹Zr]PEG- $(DFB)_1(ACUPA)_3$ with three copies of ACUPA ligands demonstrated exceptionally high PSMA-targeted cell uptake compared to [⁸⁹Zr]PEG-(DFB)₃(ACUPA)₁ with just one copy of ACUPA (Figure S17). However, as the concentration of the nanocarriers was increased to 1000 nM, the ratio of uptake to



Figure 5. *Ex vivo* organ biodistribution of ⁸⁹Zr-labeled starPEGs. (a) Tumor biodistribution of [⁸⁹Zr]starPEGs at 216 h postinjection of the nanocarriers (n = 4, mean \pm SD, **P < 0.01, Student's *t*-test). (b) Ratio of PC3-Pip to PC3-Flu tumor biodistribution of [⁸⁹Zr]starPEGs at 216 h postinjection of the nanocarriers (n = 4, mean \pm SD, *P < 0.05, Student's *t*-test). (c) Tumor to the muscle and (d) tumor to blood of [⁸⁹Zr]starPEGs at 216 h postinjection of the nanocarriers. *Ex vivo* biodistribution of ⁸⁹Zr-labeled starPEGs on selected major organs is presented in the supporting information (Figure S23).

block in PSMA+ PC3-Pip cells decreased, which demonstrated relatively higher nonspecific cell uptake at higher concentrations, presumably due to saturation of the binding sites (Figure S18). Similarly, an increase in the nonspecific cell uptake was also observed at higher time points (Figure S19).

3.2.4. Membrane-Bound and Internalization Assay. Next, we tested the degree of cellular uptake and internalization. The membrane-bound and internalized activities were isolated in PSMA- PC3-Flu and PSMA+ PC3-Pip cells by acid wash (an ice-cold mixture of 150 mM sodium chloride and 50 mM glycine) at different time points (Figures 3d and S20). Significantly higher membrane-bound activities were observed for the targeted nanocarriers in PSMA+ PC3-Pip cells, which remained almost similar over time up to 24 h (Figure S20a). The internalized activities for the targeted nanocarriers increased steadily from 1 to 24 h (Figure S20b). Overall, the nanocarrier with three copies of PSMA-targeting ACUPA ligands demonstrated higher membrane-bound and internalization than its counterpart with one ACUPA ligand. On the contrary, no evidence of PSMA-targeted cell uptake was observed for the nontargeted nanocarrier [⁸⁹Zr]PEG-(DFB)₄. Taken together, these results demonstrate efficient cell binding and internalization for the PSMA-targeted nanocarriers, with relatively higher affinity and cellular uptake when comparing $[^{89}Zr]PEG-(DFB)_1(ACUPA)_3$ to $[^{89}Zr]PEG (DFB)_{3}(ACUPA)_{1}$.

3.3. In Vivo µPET/CT Imaging. In vivo µPET/CT imaging of the ⁸⁹Zr-labeled nanocarriers was performed in the nu/nu athymic mice model with subcutaneous dual xenografts of PSMA- PC3-Flu (right flank) and PSMA+ PC3-Pip (left flank). When the tumor size reached $100-200 \text{ mm}^3$, the mice were administered the ⁸⁹Zr-labeled nanocarriers via tail vein and were subjected to multiple time point μ PET/CT imaging up to 216 h, as presented in Figure 4a. The study population was comprised of three groups, with one group for each nanocarrier (n = 4 mice). The maximum intensity projection (MIP), axial μ PET/CT, and CT images of the nanocarriers are presented in Figure 4b, and the respective coronal images are presented in Figure S21. All μ PET/CT images were segmented into respective regions of interest (ROI) over heart and tumors (Tables S4–S6), and the time-activity curves were charted (Figure S22).

Overall, starting from 24 h, the targeted nanocarriers, $[{}^{89}Zr]PEG - (DFB)_3(ACUPA)_1$ and $[{}^{89}Zr]PEG - (DFB)_1(ACUPA)_3$, demonstrated significantly increased PSMA-targeted uptake in PSMA+ PC3-Pip to PSMA- PC3-Flu tumors (Figure 4b). In comparison, the nontargeted nanocarrier $[{}^{89}Zr]PEG - (DFB)_4$ did not show any difference in the tumor uptake irrespective of PSMA expression with very high background contrast even after 216 h and instead showed higher nonspecific uptake in the skin and subcutaneous soft tissues. The ROI plot demonstrated an increase in tumor accumulation up to 72 h and a mild decrease afterward up to



Figure 6. Autoradiography images of 20 μ m tumor slices of PSMA+ PC3-Pip and PSMA– PC3-Flu tumors collected after 216 h postinjection of ⁸⁹Zr-labeled nanocarriers.

216 h, which could also be visualized in the μ PET/CT images (Figures S21 and S22). Increased background clearance and prominent tumor retention were observed for the targeted nanocarriers in PSMA+ PC3-Pip. However, at 216 h time point, [⁸⁹Zr]PEG-(DFB)₁(ACUPA)₃ showed a central tumoral reduction in signal, which could be due to the development of necrosis inside this larger size tumor (Figure 4b).

3.4. Ex Vivo Organ Biodistribution. The dual xenografts bearing mice were sacrificed after the 216 h time point $\mu PET/$ CT imaging, and the major organs including tumors were collected to quantify the distribution of ⁸⁹Zr-labeled nanocarriers. The ex vivo organ biodistribution results are presented in Figures 5 and S23 and Tables S7 and S8. Interestingly, 9.64 \pm 0.87%ID PSMA+ PC3-Pip tumor uptake was observed for [⁸⁹Zr]PEG-(DFB)₃(ACUPA)₁, which was significantly higher (**P < 0.01) than the uptake (6.69 ± 1.24%ID) obtained for [⁸⁹Zr]PEG-(DFB)₁(ACUPA)₃. However, despite greater PC3-Pip tumor uptake of [⁸⁹Zr]PEG-(DFB)₃(ACUPA)₁, a significantly higher PC3-Pip/PC3-Flu ratio (*P < 0.05) was noted for [89Zr]PEG-(DFB)1(ACUPA)3, demonstrating high PSMA-targeted uptake of the later. On the other hand, the nontargeted [89Zr]PEG-(DFB)₄ demonstrated 5.75 ± 0.74% ID uptake in PC3-Pip tumors but failed to provide a higher PC3-Pip/PC3-Flu ratio, consistent with EPR-based nonspecific uptake. While the PC3-Pip/muscle ratios of both the targeted nanocarriers were comparable at around 5-6, the PC3-Pip/blood ratio of [89Zr]PEG-(DFB)1(ACUPA)3 was significantly greater, 25 compared to 6, than DFB_4 and ACUPA₁ starPEGs (Figure 5c,d). Overall, while the PSMA+ PC3-Pip tumor accumulation of both the targeted nanocarriers is comparable, [89Zr]PEG-(DFB)1(ACUPA)3 possessed highly improved PSMA+ PC3-Pip to background contrast to that of other nanocarriers without or with one PSMA-targeting ACUPA ligands.

3.5. Autoradiography Analysis. The tumors dissected at 216 h postinjection were subjected to autoradiography analysis to explore the distribution of the ⁸⁹Zr-labeled nanocarriers inside the bulk tumor tissue (Figure 6). Both targeted nanocarriers with one or three PSMA-targeted ACUPA ligands demonstrated excellent uptake with deep tumor penetration in the PC3-Pip tumors. In contrast, only peripheral accumulation of the nontargeted nanocarrier [⁸⁹Zr]PEG-(DFB)₄ was seen in

the PC3-Pip tumor. Irrespective of the presence of PSMAtargeted ACUPA ligands, all of the nanocarriers demonstrated only peripheral accumulation in the PC3-Flu tumors. Moreover, the distribution of the targeted nanocarriers in PC3-Pip tumors was not homogenous, which could be due to the specific tumor vasculature and/or central necrosis of the largesize tumor.

4. DISCUSSION

In this study, we report two newly designed ⁸⁹Zr-labeled starPEG_{40kDa} nanocarriers with one or three copies of ACUPA ligands (Figure 2) and evaluated their PSMA-targeted imaging and deep tumor penetrability in PSMA– PC3-Flu and PSMA+ PC3-Pip prostate cancer xenografts. The nanocarriers were tethered to ⁸⁹Zr chelator DFB ligands for PET imaging.³⁵ The pharmacokinetics of both the ⁸⁹Zr-labeled PSMA-targeted nanocarriers were compared with our previously reported nontargeted nanocarrier starPEG_{40kDa} without any ACUPA ligands. In our prior study, ⁸⁹Zr-DFB₄-starPEG demonstrated very high passive accumulation and retention (>10%ID 9 days postinjection) in MX-1 and HT-29 tumor models, indicative of high EPR.²⁵

The 4-armed starPEG nanocarrier of 40 kDa molecular weight (15 nm hydrodynamic diameter) provides an optimal size for EPR-based tumor accumulation with an extended halflife.^{25,26} However, it has been well established that, along with the nanocarrier size, the pharmacokinetics of EPR-mediated passive uptake is strongly influenced by tumor vasculature permeability and macrophages.^{27,30,31} A nontargeted nanostar polymer demonstrated high and homogeneous tumor accumulation (14.8%ID/g) in CT26 tumors with highly leaky vasculature.³⁰ However, the same nanostar polymer was unable to penetrate deep into the poorly leaky BxPC3 tumor and mostly accumulated in the tumor periphery (>5.5%ID/g).³⁰ Most of the prostate cancer tumor models like PC3, DU-145, and CWR22rv1 human prostate xenografts have been evaluated to be EPR low phenotype with poor deep tumor penetration of those nanocarriers relying on the EPR-mediated passive uptake.^{13,30,31} Herein, we hypothesized that by conjugating the PSMA-targeting ACUPA ligands to the 4armed starPEG nanocarriers, the target-specific tumor

accumulation with deep tissue penetration of the nanocarriers would be improved in prostate cancer xenograft beyond passive uptake.

The targeted nanocarriers were developed by conjugating PSMA-targeted ACUPA ligands to the 4-armed starPEG through cyclooctyne linkers using an azide-cyclooctyne-based metal catalyst-free click reaction (Scheme 1). However, depending on the synthetic convenience, the ⁸⁹Zr chelator DFB was linked to the polymer arms *via* an amide linker in $[^{89}Zr]PEG-(DFB)_{4}$, a cyclooctyne triazole linker in $[^{89}Zr]PEG-(DFB)_3(ACUPA)_1$, and a p-phenylene thiourea linker in $[^{89}Zr]PEG-(DFB)_1(ACUPA)_3$ (Scheme 1). Prior reports demonstrated no marked difference in the pharmacokinetics of the PSMA-targeted probes with distinct linkers associated with the radiometal chelator.⁵⁰ The starPEG nanocarriers were purified by dialysis (12–14 kDa cutoff), and the conjugation of the ligands was confirmed by ¹H NMR analysis (Figures S4–S8).

⁸⁹Zr radiolabeling of the nanocarriers was performed using the reported protocol that yielded 26.8–46.3 MBq/mg specific activity with around 81–98% (isolated) radiolabeled yield (Figure S15 and Table S1). The specific activity and isolated yield of PEG-(DFB)₄ and PEG-(DFB)₃(ACUPA)₁ were relatively higher than those of PEG-(DFB)₁(ACUPA)₃, which could be rationalized with the number of DFB ligands conjugated to the nanocarrier. Overall, three 4-armed PEGbased nanocarriers without or with different numbers of ACUPA ligands were synthesized and radiolabeled with better specific activity than those of our prior reported starPEG nanocarriers.²⁵ The PSMA-targeted *in vitro* cell binding and *in vivo* pharmacokinetics of those polymer nanocarriers were evaluated in PSMA– PC3-Flu and PSMA+ PC3-Pip cells/ tumors.

Various cell-binding assays including competition radioligand binding, saturation binding, blocking, and internalization assays were performed to evaluate the PSMA-targeted in vitro characteristics of the nanocarriers. Around 25.5-fold lower dissociation constant (K_d) of 30.9 nM was obtained for $[^{89}Zr]PEG-(DFB)_1(ACUPA)_3$ with three PSMA-targeted ACUPA ligands to that of [89Zr]PEG-(DFB)₃(ACUPA)₁ (Figure 3b and Table S3). The blocking, membrane-bound, and internalization assay too demonstrated notably higher PSMA-targeted cell uptake of [89Zr]PEG-(DFB)1(ACUPA)3 with three copies of ACUPA ligands (Figures \$17-20). The blocking assay performed at different nanocarrier concentrations demonstrated significantly enhanced binding affinity of $[^{89}Zr]PEG-(DFB)_1(ACUPA)_3$ at a lower concentration, compared to when the assay was performed at a higher probe concentration (Figure 3c). These results demonstrated a crucial role of concentration in the binding affinity of the targeted nanocarriers and may be due to a saturation of PSMA binding sites and an increase in the nonspecific cell uptake at higher concentrations. Surprisingly, ⁶⁸Ga-PSMA-11-based competition radioligand binding assay in PSMA+ PC3-Pip cells demonstrated highly comparable IC₅₀ (459–575 nM, ^{NS}P > 0.05) obtained in two independent experiments for both the targeted nanocarriers with one or three ACUPA ligands, respectively (Table S2). It should be noted that the nonradiolabeled nanocarriers with free DFB ligands were evaluated in the competition radioligand binding assay to determine the IC₅₀ values. It is thought that the complexation of metal in the chelator could alter the overall charge and hydrophilicity of the nanocarriers and thereby could alter the

targeted binding affinity of the nanocarriers. 50-52 Additionally, few other prior studies on PSMA-targeted probes demonstrated similar inconsistent correlation of the IC₅₀ to other in vitro binding assay and in vivo pharmacokinetics.^{13,50} As expected, no sign of PSMA-targeted binding affinity was witnessed for [89Zr]PEG-(DFB)4 in any of the in vitro cellbinding assays. Overall, it was observed that the conjugation of ACUPA ligands to the polymer nanocarriers strongly influences their PSMA-targeted in vitro cell binding affinity and internalization. Moreover, the use of the multivalent PSMA binder [⁸⁹Zr]PEG-(DFB)₁(ACUPA)₃ demonstrated increased in vitro PSMA binding affinity compared against the single ACUPA containing version. These findings are consistent with other reports utilizing bivalent or multivalent PSMA binders.^{13,17,53-57} Overall, the *in vitro* findings support the use of multivalent PSMA binding to maximize cell binding affinity and uptake.

All of the nanocarriers were subjected to in vivo mPET/CT imaging (24, 72, 168, and 216 h) and organ biodistribution post 216 h imaging in the nu/nu athymic mice model implanted with subcutaneous PSMA- PC3-Flu (right flank) and PSMA+ PC3-Pip (left flank) tumors (Figure 4a). Both the targeted nanocarriers $[^{89}Zr]PEG-(DFB)_3(ACUPA)_1$ and [⁸⁹Zr]PEG-(DFB)₁(ACUPA)₃ demonstrated remarkably high PSMA-targeted uptake in PC3-Pip tumors as compared to the nontargeted [89Zr]PEG-(DFB)4, which showed only tumor peripheral accumulation irrespective of the tumor type. The imaging and organ biodistribution demonstrated that, as the number of PSMA-targeted ACUPA ligands conjugated to the nanocarriers increased, the ratio of PC3-Pip/PC3-Flu, PC3-Pip/muscle, and PC3-Pip/blood increased significantly (Figure 5). Overall, the background clearance was improved with a targeted accumulation of the nanocarriers in PC3-Pip tumors as the number of ACUPA ligands increased (Figures 4 and 5). It should be noted that, although [89Zr]PEG-(DFB)₁(ACUPA)₃ demonstrated the highest background clearance and PC3-Pip/blood ratio, [89Zr]PEG-(DFB)₃(ACUPA)₁ demonstrated relatively higher PC3-Pip uptake (9.64±0.87%ID) (Figure 5 and Table S7). These unpredicted in vivo pharmacokinetics of the PSMA-targeted starPEG nanocarriers could be explained by the binding site barrier (BSB) effect, where large-size macromolecules with higher target binding affinity could bind to cells around the periphery of the blood vessels and restrict their further smooth diffusion into the bulk tumors.^{13,23,32,33,58} As demonstrated by Simanek and co-workers, despite increasing the PSMAtargeting motifs from 4 to 64 copies, the targeted tumor uptake of the large-size nanocarriers could get restricted by the BSB effect leading to poor deep tumor penetration.¹³ Other reports also clearly demonstrate that nanocarriers with high target binding affinity might not be very effective for uniform tumor penetration, despite their high in vitro cell binding affinity.^{33,59-61} In this present study, it was evident from the *in* vitro cell-binding assay that the increasing number of ACUPA ligands effectively enhanced the PSMA binding affinity of 89 Zr]PEG-(DFB)₁(ACUPA)₃ to that of its counterpart $[^{89}$ Zr]PEG-(DFB)₃(ACUPA)₁ and could experience a relatively higher BSB effect. Being a relatively weak PSMA binder, [⁸⁹Zr]PEG-(DFB)₃(ACUPA)₁ may experience a comparatively low BSB effect to that of [89Zr]PEG-(DFB)1(ACUPA)3 and thus demonstrate higher PSMA-targeted tumor accumulation with tissue penetration. It is interesting to note that the tumor tissue penetration of [89Zr]PEG-(DFB)₃(ACUPA)₁ appears

comparable to that of $[^{89}Zr]PEG-(DFB)_1(ACUPA)_3$ (Figure 6). Another possible explanation for the reduction of central tumoral signals could be the development of necrosis in the large-size tumor at later time points. Since the imaging has been performed for up to 9 days, it is difficult to control the tumor size over time, and we observed that the PC3-Pip tumors of the mice group $(383 \pm 0.051 \text{ mg})$ treated with $[^{89}Zr]PEG-(DFB)_1(ACUPA)_3$ were larger than that of the mice group $(242 \pm 0.054 \text{ mg})$ treated with $[^{89}Zr]PEG-(DFB)_3(ACUPA)_1$. Few other reasonable limitations of this study could be a smaller sample size (n = 4) and the average size difference between PC3-Pip and PC3-Flu tumors as well.

However, considering the exceptionally higher background clearance and PC3-Pip/blood ratio, [89Zr]PEG- $(DFB)_1(ACUPA)_3$ could be more efficient candidates for therapeutic evaluation, for example, to deliver chemotherapeutic or therapeutic radionuclide (177Lu, 225Ac, etc.) payloads.^{9,17,62} The autoradiography images clearly demonstrated the tissue penetration advantage of the targeted starPEG vs the nontargeted [89Zr]PEG-(DFB)₄ in PC3-Pip and all of the nanocarriers in PC3-Flu where they were unable to penetrate the bulk prostate cancer tumors (Figure 6). Similar to other prostate cancer xenografts like CWR22rv1, DU-145, and PC3, the poorly leaky vasculature and macrophages of PC3-Pip and PC3-Flu could be the primary reason for the EPR-mediated low tumor uptake and tissue penetration.^{13,30,31} Thus, as evaluated in our prior study, the nontargeted [89Zr]PEG-(DFB)4 demonstrated very high EPRmediated tumor uptake and tissue penetration in highly leaky MX-1 and HT-29 tumor models but failed in the poorly leaky prostate cancer xenograft evaluated in this study.²⁵ Considering the EPR-driven low tumor uptake of those large-size nanocarriers, active targeting of prostate cancer is highly essential to facilitate enhanced tumor uptake with tissue penetration for improved therapeutic efficacy.

5. CONCLUSIONS

In conclusion, three 4-armed starPEG-based nanocarriers without or with different numbers of ACUPA ligands were synthesized and radiolabeled with good yields. The PSMAtargeted in vitro cell binding and in vivo pharmacokinetics of the nanocarriers were evaluated in PSMA- PC3-Flu and PSMA+ PC3-Pip cells and xenografts, demonstrating the potential influence of the number of PSMA-targeting ACUPA motifs attached to the nanocarriers. Although both the targeted nanocarriers with one or three copies of ACUPA ligands significantly improved the tumor retention and tissue penetration in PSMA+ PC3-Pip xenografts, the multivalent targeted nanocarrier [⁸⁹Zr]PEG-(DFB)₁(ACUPA)₃ with three ACUPA ligands showed a remarkably higher PC3-Pip/blood ratio and background clearance. As expected, the nontargeted [⁸⁹Zr]PEG-(DFB)₄ demonstrated low EPR-mediated uptake and peripheral accumulation in the poorly leaky PC3-Flu and PC3-Pip xenografts. Overall, PSMA-targeted multivalent polymer nanocarriers significantly improved retention and tissue penetration in xenografts with EPR low phenotypes. The developed multivalent nanocarriers with tumor penetrability and high PC3-Pip to background contrast may be a potential candidate for therapeutic evaluation.

ASSOCIATED CONTENT

1 Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsami.2c15095.

Synthetic procedures; structural characterization data (¹H and ¹³C NMR spectra, HRMS); synthetic route to (a) PEG-(DFB)₃(NH₂)₁, (b) PEG-(DFB)₃(5HCyO)₁, and (c) PEG-(DFB)₃(ACUPA)₁; synthetic route to (a) PEG-(5HCyO)₃ (DFB)₁, and (b) PEG-(DFB)₁(ACUPA)₃; radiolabeling; *in vivo* studies (PDF)

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Notes

The authors declare the following competing financial interest(s): D.V.S., G.W.A., and S.D.F. are employees of Prolynx Inc. The remaining authors declare no competing financial interest.

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ABBREVIATIONS

PCa, prostate cancer PSMA, prostate-specific membrane antigen PEG, poly(ethylene glycol) PET, positron emission tomography Ci, curie tBu, tertiary butyl DIPEA, N,N-diisopropylethylamine ACUPA, ((S)-2-(3-((S)-5-amino-1-carboxypentyl) ureido) pentanedioic acid TFA, trifluoroacetic acid DCM, dichloromethane FBS, fetal bovine serum PBS, phosphate-buffered saline NHS, N-hydroxysuccinimide DCC, N,N'-dicyclohexylcarbodiimide iTLC, instant thin-layer chromatography DFB, deferoxamine B FDA, Food and Drug Administration CyO, cyclo octyne EPR, enhanced permeability and retention EDTA, ethylenediaminetetraacetic acid IACUC, Institutional Animal Care & Use Committee ROI, region of interest EPR, enhanced permeability and retention BSB, binding site barrier

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