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molecular pharmaceutics

Novel Bifunctional Cyclic Chelator for ⁸⁹Zr Labeling–Radiolabeling and Targeting Properties of RGD Conjugates

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Supporting Information

ABSTRACT: Within the last years ⁸⁹Zr has attracted considerable attention as long-lived radionuclide for positron emission tomography (PET) applications. So far desferrioxamine B (DFO) has been mainly used as bifunctional chelating system. Fusarinine C (FSC), having complexing properties comparable to DFO, was expected to be an alternative with potentially higher stability due to its cyclic structure. In this study, as proof of principle, various FSC-RGD conjugates targeting $\alpha_v \beta_3$ integrins were synthesized using different conjugation strategies and labeled with ⁸⁹Zr. *In vitro* stability, biodistribution, and microPET/CT imaging were evaluated using [⁸⁹Zr]FSC-RGD conjugates or [⁸⁹Zr]triacetylfusarinine C (TAFC). Quantitative ⁸⁹Zr labeling was achieved within 90 min at room temperature. The distribution coefficients of the different radioligands indicate hydrophilic character. Compared to [⁸⁹Zr]DFO, [⁸⁹Zr]FSC derivatives showed excellent *in vitro* stability and resistance against



transchelation in phosphate buffered saline (PBS), ethylenediaminetetraacetic acid solution (EDTA), and human serum for up to 7 days. Cell binding studies and biodistribution as well as microPET/CT imaging experiments showed efficient receptor-specific targeting of [⁸⁹Zr]FSC-RGD conjugates. No bone uptake was observed analyzing PET images indicating high *in vivo* stability. These findings indicate that FSC is a highly promising chelator for the development of ⁸⁹Zr-based PET imaging agents. **KEYWORDS:** *fusarinine C, triacetylfusarinine C, ⁸⁹Zr, RGD peptide, positron emission tomography (PET)*

INTRODUCTION

Over the last years the positron-emitter ⁸⁹Zr has attracted considerable interest for molecular imaging applications using positron emission tomography (PET). In comparison to the commonly used radionuclides for PET (¹⁸F, ⁶⁸Ga, ¹¹C, etc.), ⁸⁹Zr with its relatively long half-life of 78.4 h is particularly suited for *in vivo* imaging at late time-points. With a mean positron energy of 0.395 MeV, which is between the positron energies of ¹⁸F (0.250 MeV) and ⁶⁸Ga (0.836 MeV), it allows for high resolution PET images for small animal imaging applications.¹

For imaging applications at late time points ⁸⁹Zr has to be stably bound to a chelator to minimize dissociation *in vivo*, as free ⁸⁹Zr can accumulate in the bone or associate with plasma proteins. Over the years, several chelators, such as diethylenetriaminepentaacetic acid (DTPA), ethylenediaminetetraacetic acid (EDTA), and 1,4,7,10-tetraazacyclododecane *N*,*N'*,*N''*,*N'''*, tetraacetic acid (DOTA), have been evaluated with limited success.² Today desferrioxamine B (DFO) is the most prominent chelator for ⁸⁹Zr labeled biomolecules (Figure 1). $[^{89}Zr]$ DFO mAb-conjugates have been used in a number of clinical trials for *in vivo* tracking and quantification of monoclonal antibodies (mAbs) by PET with good imaging quality. However, there are concerns about the stability of the complexes *in vivo* not only in relation to image quality but in particular to radiation dose. Several preclinical studies reported bone accumulation of dissociated ⁸⁹Zr ranging from 3 to 15% after 3 to 7 days.^{3–8} Alternative chelators for ⁸⁹Zr⁴⁺ that could eliminate the release of osteophilic ⁸⁹Zr⁴⁺ and lead to safer PET procedures with reduced radiation dose and optimal image quality are required for the development of novel targeted PET-tracers.

Several studies have focused on the conjugation moiety^{6,9,10} and recently also on the ligand itself.^{11–15} Considering that DFO is a linear ligand and exposed to endogenous competitive cations and natural chelators *in vivo* that may challenge the

 Received:
 February 11, 2015

 Revised:
 April 8, 2015

 Accepted:
 May 5, 2015

 Published:
 May 5, 2015



Figure 1. Chemical structures of DFO and FSC derivatives.

stability of the metal chelate, macrocyclic structures may favor a higher kinetic inertness of their metallic complexes.¹⁶

Recently we demonstrated that fusarinine C (FSC), a representative of the class of hydroxamate siderophores, is a promising bifunctional chelator for ⁶⁸Ga-radiolabeling (Figure 1).^{17,18} FSC has three primary amines, which can be derivatized in a number of ways also applying the concept of multivalency. This is the combination of several targeting units in one single molecule. By attaching a cyclic RGD peptide, binding to $\alpha_{v}\beta_{3}$ integrins that are expressed during angiogenesis, via a succinic acid linker, stable [68Ga]FSC(succ-RGD)3 with excellent receptor binding properties and in vivo targeting was prepared. ⁶⁸Ga]FSC(succ-RGD)₃ showed superiority to monomeric cyclic RGD peptides, such as [18F]Galacto-RGD and [68Ga]-NODAGA-RGD. FSC has a similar structure like DFO with three hydroxamate groups providing six oxygen donors for metal binding. We postulated that FSC could be an alternative for ⁸⁹Zr labeling with complex formation properties comparable to DFO but potentially higher stability due to its cyclic structure. Thus, increased image contrasts at delayed time points are possible. In an initial study we could demonstrate that cyclic siderophores can be labeled with ⁸⁹Zr displaying comparable properties to its ⁶⁸Ga counterparts.¹⁹

In this study triacetylfusarinine C (TAFC), the triacetylated form of FSC, was investigated, and the labeling procedure and *in vitro* stability were compared with [⁸⁹Zr]DFO. Additionally, the feasibility of different FSC conjugation strategies, where the RGD peptides have been bound either directly or through a linker, has been evaluated. Moreover, radiolabeling properties, stability, and *in vivo* behavior of [⁸⁹Zr]FSC(succ-RGD)₃ in comparison to its ⁶⁸Ga labeled counterpart were compared. The study also included microPET/CT imaging in mice bearing $\alpha_{y}\beta_{3}$ integrin expressing tumor xenografts.

EXPERIMENTAL SECTION

General. All substances described were of reagent grade and were used without further purification. Human melanoma M21 and M21-L cells were a kind gift from D. A. Cheresh, Departments of Immunology and Vascular Biology, The Scripps Research Institute, La Jolla, CA, USA. 89Zr was purchased from PerkinElmer, Inc. (USA). Analytical reversedphase high performance liquid chromatography (RP-HPLC) analysis was performed using a Vydac 218 TP5215, 150×3.0 mm column (SRD, Vienna, Austria) at a flow rate of 1.0 mL/ min with the following acetonitrile $(CH_3CN)/H_2O/0.1\%$ trifluoroacetic acid (TFA) gradients: 0-0.5 min 0% CH₃CN, 0.5-7.0 min 0-55% CH₃CN. Radiolabeling efficiency and radiochemical purity were determined by instant thin-layer chromatography (ITLC) on ITLC silica gel strips (Agilent Technologies) with 50 mM EDTA (pH 7, adjust pH using 10 M NaOH) as mobile phase. The strips were scanned using a mini-scan radio TLC scanner with flow-count detector (LabLogic, Sheffield, UK).

FSC and TAFC were prepared as described previously.²⁰ Briefly, the *Aspergillus* strains (*A. fumigatus* strain for FSC, *A. nidulans* strain for TAFC) were cultured in iron free minimal medium with 1% glucose for 36 h at 37 °C and 200 rpm. Biomass was removed by filtration, and the media-containing

siderophores were collected. TAFC was directly isolated by preparative RP-HPLC.

[Fe]Fusarinine C ([Fe]FSC). [Fe]FSC was prepared by saturating FSC solution with iron followed by purification using Amberlite XAD18 beads (Dow Chemical Company, Philadelphia, PA, USA) as column matrix. [Fe]FSC was further purified via preparative RP-HPLC and dried by lyophilization. Approximately 500 mg of a red colored solid was obtained. MALDI TOF-MS m/z [M + H]⁺ = 780.4 [C₃₃H₅₁FeN₆O₁₂; exact mass, 779.3 (calculated)].

Peptide Synthesis. The cyclic pentapeptides cyclo(-Arg(Pbf)-Gly-Asp(OtBu)-DPhe-Lys-) (c(RGDfK) (Pbf, OtBu)) and cyclo(-Arg(Pbf)-Gly-Asp(OtBu)-DPhe-Glu-) (c-(RGDfE) (Pbf, OtBu)) (protecting groups: Pbf = 2,2,4,5,7pentamethyl-3-hydrobenzofuran-6-sulfonyl; OtBu = O-tertbutyl) were prepared by cyclization of the side chain protected linear precursors, which were synthesized by Fmoc-based solid phase peptide synthesis (SPPS) as previously reported.²¹ The Z-protecting group of lysine and ODmab of glutamic acid were cleaved by hydrogen with the Pd/C catalyst or 2% hydrazine in DMF, respectively. Pbf/OtBu was cleaved with a TFA/H₂O solution (v/v 95:5). A succinylated-derivative of c(RGDfK) (Pbf, OtBu) (succ-c(RGDfK) (Pbf, OtBu)) was synthesized using c(RGDfK) (Pbf, OtBu) reacting with molar excess of succinic anhydride.¹⁷ The products were purified by preparative RP-HPLC, and their identities were confirmed by ESI-MS or MALDI TOF-MS. succ-c(RGDfK) (Pbf, OtBu), ESI-MS m/z $[M + H]^+ = 1012.6 [C_{48}H_{69}N_9O_{13}S;$ exact mass, 1011.5 (calculated)]; c(RGDfE) (Pbf, OtBu), ESI-MS $m/z [M + H]^+$ = 913.4 $[C_{43}H_{60}N_8O_{12}S;$ exact mass, 912.4 (calculated)]; c(RGDfK), MALDI TOF-MS $[M + H]^+ = 604.9$ $[C_{27}H_{41}N_9O_7;$ exact mass, 603.3 (calculated)].

[Fe]FSC(RGDfE)₃ and [Fe]FSC(succ-RGD)₃. [Fe]FSC (10 mg, 12.8 µmol) and an excess of 5 equiv of succ-c(RGDfK) (Pbf, OtBu) (64.9 mg, 64.2 µmol) or c(RGDfE) (Pbf, OtBu) (58.5 mg, 64.2 μ mol) were dissolved in 2 mL of DMF. After addition of 24.4 mg of HATU (64.2 $\mu mol)$ and 8.7 mg of HOAt (64.2 μ mol) and adjustment of the pH to 9 using DIPEA, the reaction mixture was stirred for 72 h at room temperature (RT). Hereafter, the volume of the solvent was reduced in vacuo, and the conjugate was precipitated using water. The crude [Fe]FSC(succ-RGD)₃ (Pbf, OtBu)) and [Fe]FSC(RGDfE)₃ (Pbf, OtBu) were dissolved in 2 mL of a solution composed of TFA/H₂O/triisopropylsilan in a ratio of 38:1:1. The reaction mixture was allowed to react for 1 h at RT. Subsequently, the solvent was reduced, and the crude product was obtained by precipitation using diethyl ether. Finally, the product was dried and purified via preparative RP-HPLC. $[Fe]FSC(RGDfE)_3$, yield 14.9 mg (5.9 μ mol), 46%; MALDI TOF-MS $[M + H]^+ = 2539.2 [C_{111}H_{153}FeN_{30}O_{36}$; exact mass, 2538.0 (calculated)]; [Fe]FSC(succ-RGD)₃, yield 23.5 mg (8.3 μ mol), 65%, MALDI TOF-MS [M + H]⁺ = 2836.1 $[C_{126}H_{180}FeN_{33}O_{39};$ exact mass, 2835.2 (calculated)].

SAT(PEG)₄-**RGD.** c(RGDfK) (107 mg, 0.13 mmol) in 2 mL of DMF was mixed with 855 μ L of *N*-succidimidyl-*S*-acetyl(thiotetraethylene glycol) (SAT(PEG)₄) stock solution in dimethyl sulfoxide (DMSO; 0.16 mmol, 1.2 equiv). The reaction mixture was alkalized to a pH of 9 using DIPEA and stirred for 2 h at RT. Subsequently, the solvent was reduced, and the conjugate was precipitated by addition of diethyl ether. Then, SAT-(PEG)₄-RGD was washed thrice with diethyl ether and was dried *in vacuo* to give the product as yellow oil with honey-like consistency. Yield 122 mg (0.13 mmol).

HS-(PEG)₄-RGD. SAT-(PEG)₄-RGD (12 mg, 55 μ mol) was dissolved in 0.5 mL of phosphate buffered saline (PBS, pH 7.4), and 2 mL of 0.5 M hydroxylamine solution was added. The pH was adjusted to 6.0. After the mixture was stirred for 2 h at RT, HS-(PEG)₄-RGD was purified via preparative RP-HPLC and dried by lyophilization. Yield 42.9 mg (49.5 μ mol), 90%, ESI-MS m/z [M + H]⁺ = 868.5 [C₃₈H₆₁N₉O₁₂S; exact mass, 867.4 (calculated)].

Maleimide Derivatized [Fe]FSC ([Fe]FSC(Mal)₃). [Fe]FSC (10 mg, 12.8 μ mol) and an excess of 5 equiv of 3maleimidopropionic acid *N*-hydroxysuccinimide ester (17.1 mg, 64.2 μ mol) was dissolved in 2 mL of DMF. The pH was adjusted to 8.0 using DIPEA, and the reaction mixture was stirred for 1.5 h at RT. Hereafter, [Fe]FSC(Mal)₃ was purified via preparative RP-HPLC. [Fe]FSC(Mal)₃, yield 12.7 mg (10.3 μ mol), 80%, MALDI TOF-MS [M + H]⁺ = 1234.1 [C₅₄H₆₆FeN₉O₂₁; exact mass, 1232.4 (calculated)].

[Fe]FSC(Mal-RGD)₃. HS-(PEG)4-c(RGD) (30 mg, 34.6 μ mol) was dissolved in 1 mL PBS. [Fe]FSC(Mal)₃ (12.7 mg, 10.3 μ mol) was dissolved in 200 μ L of CH₃CN and added to the solution. After 30 min, the product was purified via preparative RP-HPLC and dried by lyophilization. Yield 30.5 mg (8.0 μ mol), 78%.

Demetalation. [Fe]FSC(RGDfE)₃ (10 mg, 3.9 μ mol), [Fe]FSC(succ-RGD)₃ (10 mg, 3.5 μ mol) or [Fe]FSC(Mal-RGD)₃ (10 mg, 2.6 μ mol) was dissolved in 40-fold of a disodium EDTA solution (25 mM, pH 4). After 1 h stirring at RT the solution was directly isolated via preparative RP-HPLC and dried by lyophilization. FSC(RGDfE)₃, yield 6.2 mg (2.5 μ mol), 64%, MALDI-TOF-MS [M + H]⁺ = 2487.8, [C₁₁₁H₁₅₆N₃₀O₃₆; exact mass, 2485.1 (calculated)]; FSC(succ-RGD)₃, yield 6.4 mg (2.3 μ mol), 66%, MALDI-TOF-MS [M + H]⁺ = 2785.0, [C₁₂₆H₁₈₃N₃₃O₃₉; exact mass, 2782.3-(calculated)]; FSC(Mal-RGD)₃, yield 6.8 mg (1.8 μ mol), 69%, MALDI-TOF-MS [M + H]⁺ = 3785.2 [C₁₆₈H₂₅₂N₃₆O₅₇S₃; exact mass, 3781.7 (calculated)]. ⁸⁹Zr Radiolabeling. Approximately 30 MBq (30 μ L) of

⁸⁹Zr-oxalate was transferred into a low-protein binding Eppendorf tube, followed by addition of 27 μ L of 1 M sodium carbonate. The reaction solution was incubated for 3 min at RT. Thereafter, 100 μ L of HEPES buffer (0.5 M, pH 7.0) was added to the reaction vial. DFO (32.8 μ g), TAFC (42.6 μ g), or the FSC-RGD conjugates [(FSC(RGDfE)₃, FSC(succ-RGD)₃, or FSC(Mal-RGD)₃); 30 μ g] was added to the reaction vial. The pH of the reaction vial was measured to be sure that it was between 6.8 and 7.2. The solution was allowed to react at RT for 90 min. The reaction was monitored via ITLC (89Zrcomplexes remained at the origin and free ⁸⁹Zr migrates with the solvent front, see Supporting Information, Figure S1) and confirmed by HPLC. For animal experiments, after radiolabeling 60 μ L of CaCl₂ (0.5 M) was added to the reaction vial, which resulted in a precipitate of Ca-oxalate. The solution was passed through a 0.2 μ m filter to remove Ca-oxalate. The filtrate was diluted to an appropriate volume using saline for further evaluation.

Distribution Coefficient (logD). Aliquots of 50 μ L of radioligand ([⁸⁹Zr]DFO, [⁸⁹Zr]TAFC, or [⁸⁹Zr]FSC-RGD conjugates) were diluted in 450 μ L of PBS. Then 500 μ L of octanol was added, and the mixture was vortexed for 15 min at 1400 rpm and centrifuged for 2 min at 2000 rcf. Subsequently, 50 μ L aliquots of the aqueous and the octanol layer were collected, measured in the gamma counter, and logD values were calculated using Excel (n = 5).

Stability Assay. Determination of the stability of [⁸⁹Zr]-DFO, [⁸⁹Zr]TAFC, and [⁸⁹Zr]FSC-RGD conjugates was carried out by incubating the radioligands in PBS, 1000-fold molar excess of EDTA solution with different pH (pH 7 and pH 6; radioligand vs EDTA: 25 μ M vs 25 mM) as well as in human serum for 7 days at 37 °C, respectively. At selected time points three aliquots of PBS and EDTA solution were analyzed directly via ITLC, while the three serum aliquots were mixed with 500 μ L of CH₃CN, vortexed, and centrifuged at 20,000 rcf for 2 min prior to analysis.

Transchelation Study. [⁸⁹Zr]DFO and [⁸⁹Zr]TAFC (50 μ L each) were mixed with a 1000-fold molar excess of either TAFC or DFO for 7 days at RT. At selected time points aliquots of the solutions were analyzed directly via RP-HPLC. ([⁸⁹Zr]DFO $t_{\rm R} = 4.8$ min; [⁸⁹Zr]TAFC $t_{\rm R} = 5.9$ min). The transchelation was determined by the ratio of [⁸⁹Zr]DFO and [⁸⁹Zr]TAFC. Three replicates were analyzed for each radioligand.

Protein Binding Assay. Protein binding abilities were evaluated by incubating [⁸⁹Zr]DFO and [⁸⁹Zr]TAFC for 7 days and [⁸⁹Zr]FSC-RGD conjugates for 4 h at 37 °C in fresh human serum. Subsequently, 30 μ L of the solution was passed through a size exclusion spin column (MicroSpin G-50 column, GE healthcare, Buckinghamshire, UK) via centrifugation at 2000 rcf for 2 min. Protein binding of the conjugates in triplicates was determined by measuring the activity on the column (nonprotein bound) and the activity in the eluate (protein bound) in a gamma counter.

Internalization Assay. M21 cells ($\alpha_{\nu}\beta_{3}$ integrin positive) were diluted with RPMI 1640 (Gibco, Invitrogen Corporation, Paisley, UK) containing 1% glutamine (m/v), 1% bovine serum albumin (BSA) (m/v), CaCl₂ (1 mM), MgCl₂ (1 mM), and MnCl₂ (10 mM) to a concentration of 2.0 \times 10⁶ cells/mL. Aliquots of 1 mL cell solution were transferred to Eppendorf tubes for incubation at 37 °C for 1 h. After addition of [⁸⁹Zr]FSC(RGDfE)₃, [⁸⁹Zr]FSC(succ-RGD)₃, or [⁸⁹Zr]FSC- $(Mal-RGD)_3$ (approximately 1.5×10^6 cpm), the cells were incubated in triplicates with either PBS with 0.5% BSA (150 μ L, total series) or with 10 μ M c(RGDyV) in PBS/0.5% BSA (150 μ L, nonspecific series) at 37 °C for 90 min. After the incubation time the tubes were centrifuged, and incubation was stopped by removal of the medium. Then the cells were washed twice with ice-cold TRIS-buffered saline. Subsequently, the cells were incubated two times in acid wash buffer (20 mM acetate buffer, pH 4.5) at 37 °C for 5 min. Then the tubes were centrifuged, and the supernatant was collected in plastic vials (membrane bound activity). In a last step the cells were lysed by addition of 2 M NaOH, and the radioactivity associated with cells was collected in plastic vials (internalized radioligand fraction). Protein content in the NaOH fraction was determined using a spectrometric method (Bradford assay). The internalized activity was calculated and expressed as percentage of total activity per milligram protein.

Biodistribution Studies. All animal experiments were conducted in compliance with the Austrian animal protection laws and with approval of the Austrian Ministry of Science (BMWF-66.011/000604-II/3b/2012 and BMWFW-66.011/0049-WF/II/3b/2014).

For the evaluation of biodistribution, four healthy female BALB/c mice (Charles River Laboratories, Sulzfeld, GER) were intravenously injected with [89 Zr]TAFC (~1.5 MBq/mouse, ~4 μ g of precursor) into the tail vein and sacrificed at 6 h postinjection (p.i.). For the evaluation of tumor uptake of

[⁸⁹Zr]FSC(succ-RGD)₃ female, athymic BALB/c nude mice (Charles River Laboratories) were used. For the induction of tumor xenografts mice were injected subcutaneously with 5 \times $10^6 \alpha_{\rm v} \beta_3$ integrin positive M21 cells into the right hind limb and with 5 \times 10⁶ $\alpha_{,\beta_{3}}$ integrin negative M21-L cells (negative control) into the left hind limb of the same mouse (n = 4). The tumors were allowed to grow until they had reached a volume of 0.3 to 0.6 cm³. On the day of the experiment [⁸⁹Zr]FSC- $(succ-RGD)_3$ (~0.5 MBq/mouse, ~0.5 µg of peptide) was intravenously injected in the lateral tail vein. Mice were sacrificed by cervical dislocation at 1, 2, and 4 h after injection. Organs (spleen, pancreas, stomach, intestine, kidney, liver, heart, and lung), blood, muscle tissue, bone, and tumors were dissected and weighted. As a comparison, four tumor-bearing mice were intravenously injected in the tail vein with $[^{68}Ga]FSC(succ-RGD)_3$ (~0.5 MBq/mouse, ~0.5 μ g of precursor) and sacrificed at 1 h p.i. Activity of the different samples was measured in the gamma counter. Results were expressed as percentage of injected dose per gram tissue (% ID/g).

MicroPET/CT Imaging. MicroPET/CT imaging experiments were conducted on an Inveon microPET/CT scanner (Siemens Preclinical Solutions, Knoxville, USA). A group of four M21-tumor xenograft bearing BALB/c nude mice were administered with [⁸⁹Zr]FSC(succ-RGD)₃ (~5 MBq/mouse, ~5 μ g of peptide) via intravenous tail vein injection. MicroPET images were acquired under general anesthesia (isoflurane/O₂) for 20 min. PET data for each mouse were recorded via static scans at 1, 4, and 24 h p.i. The microPET scans were reconstructed with Inveon Acquisition Workplace software (version 1.5; Siemens Preclinical Solutions), using a 3-dimensional fast maximum *a posteriori* algorithm with the following parameters: matrix, 256 × 256 × 161; pixel size, 0.4 × 0.4 × 0.8 mm; β -value of 1.5 mm resolution with uniform variance.

Statistical Analysis. Statistical analysis was performed using SPSS 17.0 software. The biodistribution data was analyzed using Student's t test. The level of significance was set at P = 0.05.

RESULTS

Precursor Synthesis. RGD peptides could be synthesized in good yields following standardized protocols.²¹ Assemblies of the linear RGD peptides were accomplished on solid phase using Fmoc-protocols. Cyclization was carried out under high dilution conditions and subsequent Z-deprotection under hydrogen atmosphere for c(RGDfK)(Pbf, OtBu) or ODmabdeprotection with 2% hydrazine in DMF for c(RGDfE)(Pbf, OtBu). The lysine side chain was modified via addition of succinic anhydride enabling the coupling of the RGD peptide to the primary amino functions of [Fe]FSC. Amidation of succc(RGDfK)(Pbf, OtBu) or c(RGDfE)(Pbf, OtBu) with [Fe]FSC was accomplished via in situ activation using HATU/HOAt and DIPEA. Complete deprotection and subsequent demetalation of the organometallic complexes resulted in an average yield of approximately 40% and 30% for FSC(succ-RGD)₃ and FSC(RGDfE)₃, respectively. [Fe]-FSC(Mal)₃ was synthesized by [Fe]FSC reaction with 3maleimidopropionic acid N-hydroxysuccinimide ester. The deprotected c(RGDfK) was modified with $SAT(PEG)_4$ and then treated with hydroxylamine for 20 min to expose the sulfhydryl group for coupling [Fe]FSC(Mal)₃ at neutral pH, resulting in an average yield of 55%. The removal of [Fe],

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which was used to protect the complexing moiety, from [Fe]FSC-RGD conjugates was carried out using 40-fold molar excess of EDTA solution at pH 4 within 25 to 60 min with high yield (>90%). The rapid disappearance of the intense red color of the solution indicated the successful removal of [Fe]. After HPLC purification the chemical purity of the final compounds was >95%.

⁸⁹Zr Radiolabeling. Labeling of DFO, TAFC, and the FSC-RGD conjugates with ⁸⁹Zr-oxalate was performed with slight modifications as described recently.²² Radiolabeling was carried out in HEPES buffer at RT at pH 6.8 to 7.2. FSC could acquire ⁸⁹Zr quantitatively from ⁸⁹Zr-oxalate at RT with incubation times between 30 and 90 min depending on the concentration of precursor. Moreover, 1 μg of TAFC could be quantitatively labeled with 30 MBq of ⁸⁹Zr-oxalate leading to a specific activity of 25 GBq/μmol (without optimization). The time course of ⁸⁹Zr-complexation of FSC(succ-RGD)₃ (30 μg, 58 μM) is shown in Figure 2. Oxalate, which could be toxic due to the



Figure 2. Time course of ⁸⁹Zr complexation of FSC(succ-RGD)₃ (58 μ M, pH 7, at RT).

production of solid calcium oxalate causing kidney failure, was readily removed by preprecipitation with $CaCl_2$ and filtration using a 0.2 μ m filter for *in vivo* experiments.

In Vitro Characterization. [⁸⁹Zr]DFO, [⁸⁹Zr]TAFC, and [⁸⁹Zr]FSC-RGD conjugates showed high stability in PBS and human serum over a period of 7 days, and no demetalation was observed. LogD and protein binding data are summarized in Table 1. High hydrophilic character was observed for [⁸⁹Zr]FSC-RGD conjugates (logD was between -2.9 and -3.0). The lower hydrophilicity of [⁸⁹Zr]TAFC (logD -2.0) is in accordance with the data found for [⁶⁸Ga]TAFC.²³ The low protein-bound activity (<10%) over the whole monitoring period further confirmed the stability of FSC in human serum.

Table 1. LogD and Binding Properties of [89Zr]DFO,[89Zr]TAFC, and [89Zr]FSC-RGD Conjugates

⁸⁹ Zr complex	logD (pH 7.4)	incubation time (h)	protein binding (%)
[⁸⁹ Zr]DFO	-3.0 ± 0.1	168	8.7 ± 1.0
[⁸⁹ Zr]TAFC	-2.0 ± 0.0	168	6.8 ± 0.5
[⁸⁹ Zr]FSC(RGDfE) ₃	-3.0 ± 0.0	4	5.0 ± 0.8
[⁸⁹ Zr]FSC(succ-RGD) ₃	-2.9 ± 0.1	4	7.3 ± 0.5
[⁸⁹ Zr]FSC(Mal-RGD) ₃	-2.9 ± 0.2	4	6.0 ± 1.5

Stability data of [⁸⁹Zr]DFO, [⁸⁹Zr]TAFC, and [⁸⁹Zr]FSC(succ-RGD)₃ in 1000-fold molar excess of EDTA solution with different pH and transchelation data are summarized in Table 2. Based on ITLC analysis TAFC showed no transchelation in EDTA solution at pH 7 and minor transchelation (5.2%) at pH 6 after 7 days. [⁸⁹Zr]FSC(succ-RGD)₃ exhibited 4.4% transchelation, indicating high *in vitro* stability of the ⁸⁹Zr-FSC complex. In contrast, [⁸⁹Zr]DFO showed nearly 60% transchelation in EDTA at pH 7 after 7 days and more than 90% transchelation at pH 6 after 3 days. The superior stability of [⁸⁹Zr]FSC was also confirmed by the transchelation study, in which 100% of [⁸⁹Zr]DFO transchelated in 1000-fold molar excess of TAFC within 1 h, whereas 40% of [⁸⁹Zr]TAFC remained intact in 1000-fold molar excess of DFO after 7 days.

By incubating [⁸⁹Zr]FSC-RGD conjugates with $\alpha_{\gamma}\beta_{3}$ integrin positive M21 cells and carrying out corresponding blocking studies [22 µmol of c(RGDyV)] the internalization ability of the multimeric RGD peptides was determined (Figure 3). The internalized activities were 2.3 ± 0.3% for [⁸⁹Zr]FSC(RGDfE)₃, 2.1 ± 0.3% for [⁸⁹Zr]FSC(succ-RGD)₃, and 1.6 ± 0.1% for [⁸⁹Zr]FSC(Mal-RGD)₃ of total activity per milligram protein (% cpm/mg). The corresponding activities were reduced to 0.2 to 0.3% cpm/mg of the reference activity via addition of excess of c(RGDyV), which demonstrated receptor-specific binding.

Biodistribution Studies. [89Zr]TAFC showed rapid pharmacokinetics in normal mice at 6 h p.i. with fast blood clearance $(0.05 \pm 0.01\% \text{ ID/g})$, low uptake of bone $(0.04 \pm$ 0.02% ID/g), and predominant renal excretion with some retention at this late time point $(0.86 \pm 0.48\% \text{ ID/g})$. Biodistribution and specific tumor uptake of [89Zr]FSC(succ-RGD)₃ were investigated at 1, 2, and 4 h postinjection in BALB/c nude mice bearing M21 ($\alpha_v \beta_3$ integrin positive) as well as M21-L ($\alpha_v \beta_3$ integrin negative) tumor xenografts (n = 4) (Figure 4). The uptake of $[^{89}Zr]FSC(succ-RGD)_3$ in the $\alpha_v\beta_3$ integrin positive M21 tumor was $2.83 \pm 0.60\%$ ID/g at 1 h p.i. At 2 h a slight increase to the maximum uptake of $3.48 \pm 0.45\%$ ID/g was seen and a slow clearance to $2.86 \pm 0.30\%$ ID/g at 4 h p.i. In contrast, a constant but low uptake (0.5% ID/g) was observed at all time points in the contralateral $\alpha_v \beta_3$ integrin negative M21-L tumors. This highly significant difference in tumor uptake between receptor-positive and receptor-negative tumors confirmed the receptor-selective uptake found in vitro. The fast blood clearance $(0.16 \pm 0.04\% \text{ ID/g}, 0.15 \pm 0.0\% \text{ ID/g})$ g, and 0.10 \pm 0.01% ID/g at 1, 2, and 4 h p.i., respectively) led to an increased tumor/blood ratio. The comparably high activities in kidneys (6.42 \pm 0.72% ID/g, 5.77 \pm 0.20% ID/g, and 5.26 \pm 0.90% ID/g at 1, 2 and 4 h p. i., respectively) could be related to renal excretion and partial retention. Notably, the low amount of activity in the bone remained stable (0.71 \pm 0.05% ID/g, $0.79 \pm 0.28\%$ ID/g, and $0.73 \pm 0.27\%$ ID/g at 1, 2, and 4 h p. i., respectively) indicating the high *in vivo* stability of [⁸⁹Zr]FSC(succ-RGD)₃. Compared with the biodistribution data of [⁶⁸Ga]FSC(succ-RGD)₃ at 1 h, significant differences of uptake in blood, spleen, liver, and M21-L tumors were observed. The higher uptakes of $[{\rm ^{68}Ga}]FSC(succ\text{-}RGD)_3$ in the spleen (3.27 \pm 0.64% ID/g vs 1.78 \pm 0.22% ID/g) and in M21-L tumors (1.10 \pm 0.30% ID/g vs 0.50 \pm 0.12% ID/g) are related to the prolonged presence of the 68Ga-tracer in the blood $(0.63 \pm 0.26\% \text{ ID/g vs } 0.16 \pm 0.04\% \text{ ID/g})$.

A microPET/CT imaging study with tumor-bearing nude mice was performed in order to investigate the *in vivo* pharmacokinetics of $[^{89}Zr]FSC(succ-RGD)_3$ (Figure 5). Again images confirmed receptor-specific activity accumulation

Table 2. Stability and Transchelation Studies of $[^{89}Zr]DFO$, $[^{89}Zr]TAFC$, and $[^{89}Zr]FSC$ -RGD Conjugates (n = 3)

			% intact starting species by incubation time					
starting complex	competitor (1000-fold excess)	pН	1 h	4 h	1 d	3 d	5 d	7 d
[⁸⁹ Zr]DFO	EDTA	7	95.5 ± 0.5	76.0 ± 1.0	55.4 ± 3.2	41.0 ± 5.2	43.3 ± 2.5	42.2 ± 2.3
		6	94.6 ± 0.4	63.5 ± 3.6	38.2 ± 8.4	6.4 ± 2.0	6.0 ± 1.8	
	TAFC	6	0.0 ± 0.0					
[⁸⁹ Zr]TAFC	EDTA	7	99.6 ± 0.1	99.6 ± 0.1	99.3 ± 0.2	96.9 ± 0.3	98.3 ± 0.1	97.2 ± 0.2
		6	99.0 ± 0.1	98.1 ± 0.2	96.0 ± 0.6	94.7 ± 0.3	95.9 ± 0.3	94.4 ± 0.5
	DFO	6	96.7 ± 0.2	91.9 ± 0.3	74.2 ± 3.0	53.0 ± 4.1	43.3 ± 3.5	39.8 ± 2.0
[⁸⁹ Zr]FSC(succ-RGD) ₃	EDTA	7	98.2 ± 0.1	98.2 ± 0.1	96.5 ± 0.5	96.0 ± 0.4	94.5 ± 0.3	93.9 ± 0.7



Figure 3. Internalization of $[{}^{89}\text{Zr}]\text{FSC}(\text{RGDfE})_3$, $[{}^{89}\text{Zr}]\text{FSC}(\text{succ-RGD})_3$, and $[{}^{89}\text{Zr}]\text{FSC}(\text{Mal-RGD})_3$ in $\alpha_v\beta_3$ integrin positive M21 tumor cells.

in the $\alpha_v \beta_3$ integrin positive M21 tumors, and more importantly, no bone uptake was observed, confirming high *in vivo* stability of the compound. The M21 tumors could be clearly visualized over the whole monitoring period of 24 h, whereas the $\alpha_v \beta_3$ integrin negative M21-L tumors showed no uptake confirming the high specificity of [⁸⁹Zr]FSC(succ-RGD)₃. Besides the M21 tumors, only kidneys were visible, which is related to excretion and kidney retention.

DISCUSSION

The positron-emitting radionuclide ⁸⁹Zr is increasingly used for molecular imaging with PET; its clinical application is currently limited to conjugates using DFO as chelator. Even though several antibodies conjugated to DFO and radiolabeled with ⁸⁹Zr achieved some successful results, dissociation of ⁸⁹Zr *in vivo* has been observed indicating limited *in vivo* stability. Therefore, increasing efforts are made to design and synthesize novel robust and stable chelators for ⁸⁹Zr. The design of an ideal chelator is affected by many factors, such as the type and number of coordinated atoms, acyclic vs cyclic constructs, ring size, etc. Even the charge of the metal-complex may influence the biodistribution of a corresponding tracer labeled with a radiometal.

Based on the knowledge on structure, synthesis, and labeling of DFO-conjugates, the natural siderophore FSC was chosen as the starting point for our research. FSC is produced by fungi for acquiring iron from the environment and has been proven to be an excellent bifunctional chelator for ⁶⁸Ga in our laboratory.¹⁷ Therefore, FSC, having three hydroxamic acid moieties similar to those of DFO, embedded in a cyclic structure, was expected





Figure 4. Biodistribution of $[{}^{89}\text{Zr}]/[{}^{68}\text{Ga}]\text{FSC}(\text{succ-RGD})_3$ at 1, 2, and 4 h p.i. in BALB/C nude mice bearing the $\alpha_{y}\beta_3$ integrin positive M21 cell tumor on the right flank and the $\alpha_{y}\beta_3$ integrin negative control tumor M21-L on the left flank, and $[{}^{89}\text{Zr}]\text{TAFC}$ at 6 h in normal BALB/c mice. Significant differences in uptake are marked with an asterisk (P = 0.05). Each data point represents an average of biodistribution data in four animals (n = 4).

to be an interesting alternative for $^{89}\mathrm{Zr}$ labeling. In our study, TAFC was initially chosen to explore the labeling procedure and to evaluate the *in vitro* stability. It was labeled with $^{89}\mathrm{Zr}$

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Figure 5. Three-dimensional volume projections of fused microPET/ CT images of M21/M21-L tumor xenograft bearing nude mouse $([^{89}Zr]FSC(succ-RGD)_3, 5 \ \mu g, 5 \ MBq)$ at 1, 4, and 24 h p.i. Red arrow, $\alpha_{\nu}\beta_3$ integrin positive M21 tumor; blue arrow, $\alpha_{\nu}\beta_3$ integrin negative M21-L tumor.

oxalate following the established protocol found for $[^{89}Zr]$ -DFO.²² The cyclic structure of TAFC exhibits excellent complexation ability with quantitative complexation of ^{89}Zr from ^{89}Zr -oxalate between 30 to 90 min. This resulted in highly hydrophilic $[^{89}Zr]$ TAFC and $[^{89}Zr]$ FSC-RGD conjugates with low partition coefficients, indicating advantageous properties of these compounds for PET imaging. However, initial experiments using macroscale amounts of ZrCl₄ demonstrate that TAFC can, in contrast to DFO and other chelators that use ^{89}Zr -oxalate at pH 6.8–7.2 and reaction times between 1 to 2 h,¹¹ quantitatively form complexes at acidic pH (1–5) within 5 min (data not shown). This indicates that the chloride form of ^{89}Zr could be the better choice for labeling reactions. Moreover, it would avoid the use of highly concentrated solutions of oxalate, which is potentially toxic.

One of the main attractions of FSC-based constructs for ⁸⁹Zr radiolabeling lies in its excellent stability in challenging solution. [⁸⁹Zr]TAFC remained stable toward challenge with 1000-fold molar excess of EDTA and only limited transchelation was found after days of incubation with 1000-fold molar excess of DFO. In comparison [⁸⁹Zr]DFO revealed rapid transchelation toward EDTA even at pH 7 and complete transmetalation toward TAFC (both 1000-fold molar excess) within hours. This high stability of the [⁸⁹Zr]FSC structure was further confirmed by the studies with the [⁸⁹Zr]FSC-RGD conjugates. Additionally, as the pH in tumors (approximately pH 6.5) is lower than in healthy tissue, the observed high stability of [⁸⁹Zr]TAFC in slightly acidic conditions (pH 6) may be beneficial in particular for cancer imaging.

Inspired by the promising results found for [⁸⁹Zr]TAFC, the potential use of FSC conjugated to targeting vectors was investigated. For several reasons cyclic RGD pentapeptides were chosen as model vectors. First, RGD-peptides are very stable *in vitro* and *in vivo* and, more important, tolerate a variety of modifications at position 5 in the amino acid sequence without losing binding affinity.²⁴ Second, rapid predominantly renal excretion with low background activity is described for a big variety of different tracer based on this pentapeptide allowing an easier monitoring of the distribution of fragments based on rapid degradation in the body and especially in the bone.²⁵ Third, it is a well-known class of tracer where a great set of data is available.^{26,27} Fourth, it is known that binding

affinity and imaging properties of this tracer class benefit from the so-called multimerization approach.²⁸ Due to the fact that FSC includes three primary amine functions, this chelator allows conjugation of up to three ligands making them, besides working as chelator for the radiometal, an ideal scaffold for such approaches. Thus, this option additionally increases potential applications of FSC and derivatives. Anyway, it has to be pointed out that only small peptides may benefit from such a multimerization approach and not antibodies, which is the main field of application of ⁸⁹Zr-labeling. Nevertheless, it has been demonstrated by using, e.g., ⁶⁴Cu-NODAGA-RGD that tumorto-background ratios further improve with prolonged imaging time making labeling of this peptide class with ⁸⁹Zr not *per se* uninteresting.²⁹

As a proof of concept, different cyclic RGD pentapeptides were synthesized, and versatile conjugation strategies were explored. $FSC(succ-RGD)_3$ was synthesized using a succinic acid linker by amidation with a yield of 40%. In another approach c(RGDfE) was directly conjugated via the glutamic acid carboxylate to the chelator. This approach reduces the synthesis steps but also resulted in a lower yield possibly related to steric effects. In a third approach the maleimide—thiol strategy was introduced. This is a well-established technique and allows fast and mild reaction conditions in aqueous solution, facilitating the site specific conjugation of peptides and antibodies, which might be not stable under harsh conditions. Thus, the PEG-modified RGD-peptide could be coupled with even higher yield as found for the other two derivatives under mild conditions with the FSC derivative.

Biological properties of [89Zr]FSC(succ-RGD)₃ were comparable to the properties of [⁶⁸Ga]FSC(succ-RGD)₃, which has clearly shown the advantage of multivalency through remarkable improvement of internalized activity in vitro and tumor uptake in vivo in our previous study.¹⁷ An exception was the lower activity concentration in blood resulting in better tumor/blood ratios for the 89Zr compound in comparison with the ⁶⁸Ga-labeled analogue, indicating an excellent in vivo stability of [89Zr]FSC(succ-RGD)3. The high stability was confirmed by microPET/CT imaging using the same murine tumor model. These studies clearly demonstrated no uptake in bone and high uptake in $\alpha_v \beta_3$ integrin positive M21 tumors. A reason for the superior properties of [89Zr]FSC(succ-RGD)₃ compared with [⁶⁸Ga]FSC(succ-RGD)₃ could partially be to the difference in the charge, having been reported to potentially influence excretion patterns of such biomolecules.³⁰

The preferred coordination number for ⁸⁹Zr is eight, which was confirmed recently.³¹ Certain developments aim on the design and synthesis of novel both octadentate and oxygen-rich Zr⁴⁺ chelates to achieve superior stabilities to DFO. Both Deri et al. and Patra et al. reported two kinds of linear octadentate chelators named HOPO and DFO*.^{11,12} Guérard et al. also described a cyclic octadentate tetrahydroxamic acid chelator (C7).¹³ In vitro studies of all three octadentate ligands showed improved stability. However, HOPO and C7 lack the pendant group for coupling it to a targeting molecule. DFO* was linked to the bombesin peptide analogue [Nle¹⁴]BBS(7-14), but was only investigated in vitro. [89Zr]TAFC and C7, both have a 36membered ring but different coordination numbers for ⁸⁹Zr. Nevertheless, transchelation found in an EDTA challenging experiment was lower for TAFC (3-5% TAFC vs 13% C7 after 7 days) despite its lower coordination number (6 vs 8).

Although further research is warranted, our study shows that the use of macrocyclic structure leads to considerably improved

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in vitro stability compared to DFO. A limitation of the study is that DFO cannot be used to generate a trimeric structure. Thus, in vivo comparison of both chelating systems is difficult because not only the chelator system would change but also the number of targeting moieties. Based on these changes the in vivo properties are not only correlated to the different chelators but also are influenced by the different structures. Therefore, it was decided for this proof of principle study to restrict comparison to in vitro assays where it is assumed that these differences have only minor effects. Anyway, to our knowledge this is the first report where a ⁸⁹Zr-labeled macrocyclic bifunctional chelating system conjugated to a target vector is studied in vivo including biodistribution and PET imaging. Both, the in vitro and in vivo studies of [89Zr]TAFC and [89Zr]FSC(succ-RGD)₃ demonstrated very promising results. However, small peptides, even though an excellent model for this evaluation based on their rapid pharmacokinetics, are certainly not the most suitable application for this concept. Nevertheless, for larger targeting vectors (such as antibodies or antibody fragments), due to their slower pharmacokinetics, FSC system could be a very interesting alternative for ⁸⁹Zr-labeling to currently used bifunctional chelators. At the moment, mono- and diacetylated FSC are under investigation, which may, due to the reduced free valences, better suit the application with higher molecular weight targeting vector systems.

CONCLUSION

Versatile conjugation strategies of FSC were explored using RGD as model peptide and quantitative labeling with ⁸⁹Zr was achieved successfully. Excellent *in vitro* and *in vivo* stability of [⁸⁹Zr]TAFC and [⁸⁹Zr]FSC-RGD conjugates were demonstrated. These results led us to conclude that FSC is a novel promising chelator for the development of ⁸⁹Zr-based PET imaging agents.

ASSOCIATED CONTENT

S Supporting Information

Figure S1 showing the representative chromatograms of ITLC of ⁸⁹Zr-oxalate and [⁸⁹Zr]TAFC. The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.molpharmaceut.Sb00128.

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Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

We acknowledge the financial support of the Austrian Science Foundation (FWF) grant P 25899-B23 and the China Scholarship Council.

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