A Thiol-Reactive ¹⁸F-Labeling Agent, *N*-[2-(4-¹⁸F-Fluorobenzamido)Ethyl]Maleimide, and Synthesis of RGD Peptide-Based Tracer for PET Imaging of $\alpha_v \beta_3$ Integrin Expression

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The cell adhesion molecule integrin $\alpha_v \beta_3$ plays a key role in tumor angiogenesis and metastasis. A series of ¹⁸F-labeled RGD peptides have been developed for PET of integrin expression based on primary amine-reactive prosthetic groups. In this study we introduced a new method of labeling RGD peptides through a thiolreactive synthon, N-[2-(4-18F-fluorobenzamido)ethyl]maleimide (18F-FBEM). Methods: 18F-FBEM was synthesized by coupling N-succinimidyl 4-18F-fluorobenzoate (18F-SFB) with N-(2-aminoethyl)maleimide. After high-pressure liquid chromatography purification, it was allowed to react with thiolated RGD peptides. and the resulting tracers were subjected to receptor-binding assay, in vivo metabolic stability assessment, biodistribution, and microPET studies in murine xenograft models. Results: Conjugation of monomeric and dimeric sulfhydryl-RGD peptides with ¹⁸F-FBEM was achieved in high yields (85% \pm 5% nondecaycorrected on the basis of ¹⁸F-FBEM). The radiochemical purity of the ¹⁸F-labeled peptides was >98% and the specific activity was 100~150 TBq/mmol. Noninvasive microPET and direct tissue sampling experiments demonstrated that both ¹⁸F-FBEM-SRGD (RGD monomer) and ¹⁸F-FBEM-SRGD2 (RGD dimer) had integrin-specific tumor uptake in subcutaneous U87MG glioma and orthotopic MDA-MB-435 breast cancer xenografts. Conclusion: The new tracer ¹⁸F-FBEM-SRGD2 was synthesized with high specific activity via ¹⁸F-FBEM and the tracer exhibited high receptor-binding affinity, tumor-targeting efficacy, metabolic stability, as well as favorable in vivo pharmacokinetics. The new synthon ¹⁸F-FBEM developed in this study will also be useful for radiolabeling of other thiolated biomolecules.

Key Words: thiol-reactive synthon; ¹⁸F-FBEM; microPET; ¹⁸F labeling; integrin $\alpha_v \beta_3$

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The $\alpha_v \beta_3$ integrin, which binds to arginine-glycineaspartic acid (RGD)-containing components of the interstitial matrix, such as vitronectin, fibronectin, and thrombospondin, is significantly upregulated on endothelium during angiogenesis but not in quiescent endothelium (1,2). The special role of integrin $\alpha_v \beta_3$ in tumor invasion and metastasis arises from its ability to recruit and activate matrix metalloproteinase MMP-2 and plasmin, which degrade components of the basement membrane and interstitial matrix (3). Integrins expressed on endothelial cells modulate cell migration and survival during angiogenesis, whereas integrins expressed on carcinoma cells potentiate metastasis by facilitating invasion and movement across blood vessels (2,4). Antagonists of integrin $\alpha_v \beta_3$ (antibodies, peptides, and peptidomimetics) can inhibit tumor angiogenesis, tumor growth, and metastasis in vivo (5). The ability to noninvasively visualize and quantify tumor integrin $\alpha_v \beta_3$ expression level will provide new opportunities to document tumor (tumor cells and sprouting tumor vasculature) integrin expression, to more appropriately select patients for antiintegrin treatment, and to monitor treatment efficacy in integrin-positive patients.

Over the last several years, significant advances have been achieved in developing novel probes for multimodality molecular imaging of tumor integrin expression (6). Small molecules, peptides, peptidomimetic integrin $\alpha_v \beta_3$ antagonists, and antibodies have been labeled with radioisotopes, superparamagnetic nanoparticles, fluorescent dyes, quantum dots, and microbubbles for PET, SPECT, MRI, near-infrared fluorescence, and ultrasound imaging of small animals, mostly tumor models (6,7). Because of the high sensitivity and adequate spatial and temporal resolution of PET, development of PET probes for integrin expression imaging is currently the most active among all of these modalities.

Cyclic RGD peptide was first labeled with ¹⁸F by Haubner et al. and the resulting ¹⁸F-galacto-RGD exhibited integrin $\alpha_v \beta_3$ – specific tumor uptake in an integrin-positive

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M21 melanoma xenograft model (8,9). Initial clinical trials in healthy volunteers and a limited number of cancer patients revealed that this tracer can be administered safely to patients and is capable of delineating certain lesions that are integrin positive with some indication of integrin $\alpha_v \beta_3$ expression level in vivo (9,10). We labeled a series of mono-, di-, and tetrameric RGD peptides with ¹⁸F or ⁶⁴Cu for integrin-positive tumor targeting (11–19). In particular, the dimeric RGD peptide-based tracer ¹⁸F-FRGD2 was found to be able to visualize and quantify the integrin expression level in vivo (11,12).

Radiofluorination of RGD peptides generally uses 18Fsynthons such as N-succinimidyl 4-18F-fluorobenzoate (18F-SFB) (20,21) or p-nitrophenyl ¹⁸F-fluoropropionate (¹⁸F-NFP) (22) to form a stable amide bond by reacting with primary amino groups of RGD peptides. It was also reported that oxoamino derivatives of RGD peptides react with 4-18Ffluorobenzaldehyde (¹⁸F-FBA) under acidic condition to form an oxime (23). A few ¹⁸F labeled thiol-reactive reagents have been reported in the literature—namely, 1-(4-18F-fluorophenyl)pyrrole-2,5-dione (18F-FPPD) (24), N-[3-(2,5-dioxo-2,5dihydropyrrol-1-yl)phenyl]-4-18F-fluorobenzamide (18F-DDPFB) (24), 1-[3-(2-(¹⁸F-fluoropyridin-3-yloxy)propyl]pyrrole-2, 5-dione (^{18}F -FPyME) (25), and N-[4-[(4- ^{18}F -fluorobenzylidene)aminooxyl]butyl]maleimide (18F-FBABM) (26). However, no in vivo microPET data have been reported on tracers synthesized using these prosthetic groups. In this study, we developed a new thiol-reactive synthon, N-[2-(4-18F-fluorobenzamido)ethyl]maleimide (18F-FBEM), for ¹⁸F labeling of thiol-containing molecules. Two thiolated RGD peptides were labeled with ¹⁸F through ¹⁸F-FBEM and tested in murine xenograft models.

MATERIALS AND METHODS

Unless otherwise specified, all chemicals were of analytic grade and commercially available. RGD peptides were synthesized as previously reported (11,12). N-Succinimidyl S-acetylthioacetate (SATA), hydroxylamine·HCl, and tris(2-carboxyethyl)phosphine hydrochloride (TCEP·HCl) were purchased from Pierce Biotechnology, Inc. N-(2-Aminoethyl)maleimide trifluoroacetate salt, 4-fluorobenzonic acid, N,N,N',N'-tetramethyl-O-(N-succinimidyl) uronium tetrafluoroborate (TSTU), N-hydroxysuccinimide (NHS), and N,N-diisopropylethylamine (DIPEA) were purchased from Sigma-Aldrich. No-carrier-added ¹⁸F-F⁻ was obtained from the in-house PETtrace cyclotron (GE Healthcare). The ¹⁸F-F⁻ was provided in a mixture solution of K₂CO₃ (2 mg/mL in water) and Kryptofix 2.2.2. (Sigma-Aldrich; 10 mg/mL in acetonitrile). The semipreparative reversed-phase high-pressure liquid chromatography (RP-HPLC) system was reported earlier (12). Reversedphase extraction C₁₈ Sep-Pak cartridges (Waters) were pretreated with methanol and water before use.

The SATA-RGD peptides were prepared following the protocol supplied by the vender. Briefly, c(RGDyK) or $E[c(RGDyK)]_2$ (5 μ mol) in 1 mL 50 mmol/L $Na_2B_4O_7$ buffer (pH 8.5) was mixed with 100 μ L SATA solution in dimethyl sulfoxide (DMSO; 6 μ mol). After the reaction had gone to completion as shown by analytic RP-HPLC, it was quenched by 100 μ L 2% trifluoroacetic

acid (TFA) in water. The crude product was lyophilized without purification. The yield of SATA-c(RGDyK) (HPLC retention time [R_t], 12.1 min) was 95% and that of SATA-E[c(RGDyK)]₂ (R_t, 13.6 min) was 65% on the basis of analytic RP-HPLC.

The crude SATA-RGD peptides (20 mg) were dissolved in 1 mL water and 100 μL of 0.5 mol/L hydroxylamine solution were added. The pH was adjusted to 6.0. After 2 h, sulfhydryl-c(RGDyK) and sulfhydryl-E[c(RGDyK)]_2 (denoted as SRGD and SRGD2, respectively) were purified by semipreparative RP-HPLC. The overall yield was 80% and 50% for SRGD (R_t, 10.7 min) and SRGD2 (R_t, 13.1 min), respectively; little or no dimerization was observed for either peptide when stored under acidic condition (pH 3). MALDITOF MS (matrix-assisted laser desorption/ionization time-of-flight mass spectroscopy): SRGD, $C_{29}H_{43}N_9O_9S$, calculated 693.3, observed 694.5 ([M+H]^+); SRGD2, $C_{61}H_{89}N_{19}O_{19}S$, calculated 1,423.6, observed 1,422.7 ([M+H]^+).

N-(2-Aminoethyl)maleimide (5 μmol) in 200 μL acetonitrile, N-succinimidyl 4-fluorobenzoate (4.5 μmol) in 100 μL acetonitrile, and 500 μL 50 mmol/L Na₂B₄O₇ buffer (pH 8.5) were mixed and reacted at 50°C for 20 min. The reaction was quenched by adding 100 μL 2% TFA in water. HPLC purification gave FBEM in 85% yield. 1 H NMR (chloroform-d, 400 MHz): 7.79 –7.76 (m, 2H, phenyl o-H); 7.26 –7.09 (m, 2H, phenyl m-H); 6.75 (2H, CH=CH); 3.86 – 3.65 (m, 4H, CH₂–CH₂).

The sulfhydryl-RGD peptides (0.5 μ mol) were dissolved in 0.5 mL phosphate-buffered saline (PBS, pH 7.4; Invitrogen Corp.). FBEM (0.55 μ mol) was dissolved in 200 μ L acetonitrile and added to the solution. After 30 min, the reaction mixture was subjected to HPLC purification. FBEM-SRGD (R_t, 14.8 min) and FBEM-SRGD2 (R_t, 15.2 min) were obtained with 80% yield. MALDI-TOF MS: FBEM-SRGD, C₄₂H₅₄FN₁₁O₁₂S, calculated 956.1, observed 956.7 ([M+H]⁺); FBEM-SRGD2, C₇₃H₉₈FN₂₁O₂₂S, calculated 1,672.5, observed 1,673.5 ([M+H]⁺).

Radiochemistry

¹⁸F-SFB was synthesized as previously reported (*12,15*) with C₁₈ Sep-Pak cartridge purification. It was dissolved in acetonitrile (300 μL) and 1 mg N-(2-aminoethyl)maleimide in 500 μL acetonitrile and 20 μL DIPEA were added. The reaction mixture was heated to 40°C for 20 min and then quenched by addition of 50 μL TFA. HPLC purification gave ¹⁸F-FBEM (R_t, 13.9 min; total reaction time, 150 \pm 20 min with nondecay-corrected radiochemical yield of 5% \pm 2% on the basis of ¹⁸F-F⁻; specific activity, 150~200 TBq/mmol).

¹⁸F-FBEM was dissolved in 600 μL PBS buffer (pH 7.4); 0.2 mg of sulfhydryl-RGD peptide in 50 μL DMSO and 1.0 mg TCEP·HCl in 0.1 mL water were then added. The pH was adjusted to 7.0~7.5 using 0.2 mol/L NaOH solution. The reaction mixture was kept at room temperature (r.t.) for 20 min. HPLC purification gave ¹⁸F-FBEM-SRGD (R_t, 14.9 min) and ¹⁸F-FBEM-SRGD2 (R_t, 15.3 min) in 80% nondecay-corrected yield in both cases. The radiotracers were reconstituted in PBS and passed through a 0.22-μm Millipore filter into a sterile vial for in vivo applications. Nonreacted sulfhydryl-RGD peptides were baseline-separated from the desired products during HPLC and the specific activity of the tracers was 100-150 TBq/mmol.

Eppendorf microcentrifuge tubes containing 500 μ L of octanol, 500 μ L of normal saline, and \sim 370 kBq of ¹⁸F-FBEM-SRGD or ¹⁸F-FBEM-SRGD2 were vortexed vigorously for 1 min. Each tube was centrifuged at 14,000 rpm for 5 min and the activities in 20- μ L aliquots of both organic and aqueous layers were measured

by a γ -counter (GMI, Inc.). The reported octanol/water partition coefficient represents the mean \pm SD of 6 measurements.

In Vitro Cell-Binding Assay

Both U87MG and MDA-MB-435 cell lines were purchased from American Type Culture Collection and the culture media were obtained from Invitrogen Co. U87MG glioblastoma cells were grown in Dulbecco's modified Eagle medium (DMEM, low glucose) and MDA-MB-435 breast cancer carcinoma cells were grown in Leibovitz's L-15 medium. Both cell lines were cultured in the medium supplemented with 10% (v/v) fetal bovine serum (FBS) at 37°C. In vitro integrin-binding affinities and specificities were assessed via displacement cell-binding assays using 125Iechistatin as the integrin $\alpha_v \beta_3$ -specific radioligand. Both U87MG and MDA-MB-435 cells are integrin $\alpha_v \beta_3$ positive (12). Cellbinding assay were performed using U87MG cells. The cells were harvested, washed twice with PBS, and resuspended $(2 \times 10^6 \text{ cells/})$ mL) in binding buffer (20 mmol/L Tris, pH 7.4, 150 mmol/L NaCl, 2 mmol/L CaCl₂, 1 mmol/L MgCl₂, 1 mmol/L MnCl₂, 0.1% bovine serum albumin). Millipore 96-well filter multiscreen DV plates (pore size, 0.65 µm) were seeded with 10⁵ cells per well and incubated with ¹²⁵I-echistatin (30,000 cpm/well) in the presence of increasing concentrations of different RGD peptide analogs (0-1,000 nmol/L). The total volume in each well was adjusted to 200 µL. After incubation at r.t. for 2 h, the plates were filtered through a multiscreen vacuum manifold and washed twice with cold binding buffer. The filters were collected and the radioactivity was measured using a γ-counter. The best-fit IC₅₀ (inhibitory concentration of 50%) values for U87MG cells were calculated by fitting the data by nonlinear regression using GraphPad Prism (GraphPad Software, Inc.). Experiments were performed twice with triplicate samples.

Animal Models

All animal experiments were performed under a protocol approved by the Stanford University Administrative Panel on Laboratory Animal Care. The MDA-MB-435 breast cancer model was established by orthotopic injection of 5×10^6 cells into the left mammary fat pad of female athymic nude mice, whereas the U87MG tumor model was obtained by injecting a mixture of 5×10^6 cells suspended in 50 μL medium and 50 μL Matrigel (BD Biosciences) into the right front leg. The mice were used for biodistribution and microPET imaging studies when the tumor volume reached $300-400~mm^3~(3-4~wk~after~inoculation~for~both~U87MG~and~MDA-MB-435~tumors).$

Biodistribution Studies

Female athymic nude mice bearing both U87MG and MDA-MB-435 tumors were injected with 1 MBq of $^{18}\text{F-FBEM-SRGD}$ or $^{18}\text{F-FBEM-SRGD}$. The mice were sacrificed and dissected at 10, 30, and 60 min after injection. Blocking experiment was performed by coinjecting radiotracer with a saturating dose of c(RGDyK) (10 mg/kg mouse body weight) and the mice were sacrificed at 60 min after injection. Blood, tumor, major organs, and tissues were collected and wet weighed (contents in the intestines were removed before weighing). The radioactivity in the tissues was measured using a γ -counter. The results were calculated as percentage injected dose per gram (%ID/g). For each mouse, the radioactivity of the tissue samples was calibrated against a known aliquot of the injectate and normalized to a body weight of 20 g. Values are expressed as mean \pm SD for 3 animals per group.

microPET and Image Analysis

PET scans were performed using a microPET R4 rodent model scanner (Concorde Microsystems Inc.). The scanner has a computercontrolled bed and 10.8-cm transaxial and 8-cm axial fields of view (FOVs). It has no septa and operates exclusively in the 3-dimensional list mode. Animals were placed near the center of FOV of the microPET scanner, where the highest image resolution and sensitivity are available. Mice were injected with 3.7 MBq of ¹⁸F-FBEM-SRGD or ¹⁸F-FBEM-SRGD2 via tail vein under isoflurane anesthesia. The 60-min dynamic (5 \times 60 s, 5 \times 120 s, 5 \times 180 s, 6×300 s) microPET data acquisition (total of 21 frames) was started about 3 min after radiotracer injection. Later timepoint static images were also acquired as 10-min static images after obtaining a 1-h dynamic scan. The images were reconstructed by a 2-dimensional ordered-subsets expectation maximum algorithm and no correction was applied for attenuation or scatter (27).

For each microPET scan, regions of interests (ROIs) were drawn over each tumor, normal tissue, and major organs by using vendor software ASI Pro 5.2.4.0 on decay-corrected whole-body coronal images. The maximum radioactivity concentration (accumulation) within a tumor or an organ was obtained from mean pixel values within the multiple ROI volume, which were converted to MBq/mL/min by using a conversion factor. Assuming a tissue density of 1 g/mL, the ROIs were converted to MBq/g/min, and then divided by the administered activity to obtain an imaging ROI-derived %ID/g.

Metabolic Stability

Athymic nude mice bearing U87MG tumor were intravenously injected 3.7 MBq of ¹⁸F-FBEM-SRGD2. The animals were sacrificed 60 min after tracer injection. Blood, urine, liver, kidneys, and tumor were collected. Blood was immediately centrifuged for 5 min at 13,200 rpm. Organs were homogenized using an IKA Ultra-Turrax T8 homogenizer (IKA Works Inc.), suspended in 1 mL of PBS buffer, and centrifuged for 5 min at 13,200 rpm. After removal of the supernatant, the pellets were washed with 500 µL of PBS. For each sample, supernatants of both centrifugation steps were combined and passed through Sep-Pak C₁₈ cartridges. The urine sample was directly diluted with 1 mL of PBS and then passed through the cartridge. The cartridges were washed with 2 mL of water and eluted with 2 mL of acetonitrile containing 0.1% TFA. After removal of acetonitrile, the residue was redissolved in 1 mL of water and injected onto an analytic HPLC column. Radioactivity was monitored using a solid-state radiation detector. The eluent was also collected using a fraction collector (0.5 min/fraction) and the activity of each fraction was measured by a γ-counter.

Statistical Analysis

Quantitative data are expressed as mean \pm SD. Means were compared using 1-way ANOVA and a Student t test. P values < 0.05 were considered significant.

RESULTS

Chemistry

Both monomeric peptide SRGD and the dimeric peptide SRGD2 were synthesized with good overall yield (80% for SRGD and 50% for SRGD2, respectively). FBEM was prepared by reacting SFB with *N*-(2-aminoethyl)maleimide (90% yield). The conjugation of SRGD or SRGD2 with

FBEM was performed at r.t. The resulting conjugates FBEM-SRGD and FBEM-SRGD2 were purified by RP-HPLC and confirmed by MALDI-TOF mass spectrometry.

 18 F-SFB was synthesized following a previously reported procedure (*12,15*). 18 F-FBEM was obtained by coupling 18 F-SFB with *N*-(2-aminoethyl)maleimide (Fig. 1A). After HPLC purification, the thiol-reactive synthon was allowed to react with SRGD or SRGD2 at r.t. for 20 min. Starting from 18 F-F $^-$, the total reaction time including final HPLC purification was about 200 \pm 25 min. The overall decay-corrected radiochemical yield was 20% \pm 4% (n = 5) for both 18 F-FBEM-SRGD (Fig. 1B) and 18 F-FBEM-SRGD2 (Fig. 1C). On the basis of 18 F-FBEM, both reactions were

achieved in high yields (85% \pm 5% nondecay corrected), virtually quantitative. The radiochemical purity of the ¹⁸F-labeled peptides was >98% according to analytic HPLC. The octanol/water partition coefficient (log *P*) for ¹⁸F-FBEM-SRGD was determined to be 0.93 \pm 0.02, indicating the hydrophobic character of this tracer, whereas ¹⁸F-FBEM-SRGD2 was hydrophilic (log *P* = -1.69 ± 0.02).

In Vitro Cell-Binding Assay

The cell-binding affinity studies of c(RGDyK), E[c(RGDyK)]₂, FBEM-SRGD, and FBEM-SRGD2 with U87MG cells are summarized in Figure 2. All 4 peptides inhibited the binding of ¹²⁵I-echistatin to U87MG cells in a

FIGURE 1. (A) Synthetic route for *N*-[2-(4-¹⁸F-fluorobenzamido)ethyl]maleimide (¹⁸F-FBEM). (B) Structure of ¹⁸F-FBEM-SRGD. (C) Structure of ¹⁸F-FBEM-SRGD2.

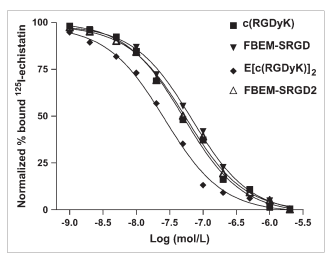


FIGURE 2. Cell-binding assay of c(RGDyK), E[c(RGDyK)]₂, FBEM-SRGD, and FBEM-SRGD2 using U87MG cells (integrin $\alpha_{\rm v}\beta_3$ –positive human glioblastoma). The cell-binding affinity of the peptides was determined by performing competitive displacement studies with 125 l-echistatin. IC₅₀ values for c(RGDyK), E[c(RGDyK)]₂, FBEM-SRGD, and FBEM-SRGD2 were 51.3 \pm 4.2, 26.1 \pm 3.2, 66.8 \pm 5.1, and 55.1 \pm 6.5 nmol/L, respectively (n=6).

dose-dependent manner. The IC₅₀ values for c(RGDyK), E[c(RGDyK)]₂, FBEM-SRGD, and FBEM-SRGD2 were 51.3 ± 4.2 , 26.1 ± 3.2 , 66.8 ± 5.1 , and 55.1 ± 6.5 nmol/L, respectively, indicating that FBEM conjugation had minimal effect on the integrin-binding avidity of the RGD peptides.

Biodistribution

Biodistribution of ¹⁸F-FBEM-SRGD and ¹⁸F-FBEM-SRGD2 was determined in athymic nude mice bearing both U87MG and MDA-MB-435 tumors and the results are shown in Figure 3. For ¹⁸F-FBEM-SRGD (Fig. 3A), the U87MG and MDA-MB-435 tumor uptakes were 1.33 \pm 0.28 and 1.43 ± 0.11 %ID/g, respectively, at 60 min after injection. When blocked by coinjection of c(RGDyK) at a dose of 10 mg/kg body weight, the tumor uptake decreased to 0.40 \pm 0.02 %ID/g for the U87MG tumor (P < 0.05when compared with the U87MG tumor without blocking) and 0.77 \pm 0.04 %ID/g for the MDA-MB-435 tumor (P <0.05 when compared with the MDA-MB-435 tumor without blocking), respectively. Intestine exhibited a high uptake of $^{18}\text{F-FBEM-SRGD}$ (16.57 \pm 0.81 %ID/g when c(RGDyK) was coinjected), most probably due to the relatively hydrophobic nature of the tracer, which is consistent with our previous result for ¹⁸F-FB-RGD (18). For ¹⁸F-FBEM-SRGD2 (Fig. 3B), the U87MG and MDA-MB-435 tumor uptakes were 2.71 ± 0.19 and 5.25 ± 0.17 %ID/g, respectively, at 60 min after injection (P < 0.01 for U87MG and P < 0.001 for MDA-MB-435 when compared with ¹⁸F-FBEM-SRGD). When blocked by coinjection of c(RGDyK) at 10 mg/kg body weight, the tumor uptake of U87MG tumor decreased >5-fold to 0.52 \pm 0.26 %ID/g (P < 0.01). Blocking reduced

the tumor uptake of both tracers to the background level (due to nonspecific binding in normal organs), clearly indicating integrin-specific binding. Kidney uptake of ¹⁸F-FBEM-SRGD2 was high at an early time point (11.40 \pm 0.22 %ID/g at 10 min after injection), but the washout was also fast. The 2 tracers exhibited different excretion routes due to the difference in hydrophilicity. ¹⁸F-FBEM-SRGD is relatively hydrophobic, therefore exhibiting mainly hepatobiliary excretion, whereas ¹⁸F-FBEM-SRGD2 is more hydrophilic and mainly excreted through the kidney. It is worth noting that more radioactivity accumulated in the MDA-MB-435 tumor with time for ¹⁸F-FBEM-SRGD2 (Fig. 3B), which might be due to the internalization of the tracer. A similar phenomenon was observed in our previous studies for the ¹⁸F-FRGD2 tracer (12). Comparing these 2 radiotracers, ¹⁸F-FBEM-SRGD2 has significantly higher kidney and tumor uptake, whereas ¹⁸F-FBEM-SRGD has much higher intestine uptake. There was no major difference in the tracer uptake by other organs between the 2 tracers.

microPET

Dynamic microPET scans were performed for both radiotracers. Selected coronal images at different time points after injection in a mouse bearing both subcutaneous U87MG and orthotopic MDA-MB-435 tumors are shown in Figure 4. High tumor activity accumulation was observed as early as 6 min after injection for both tracers. For ¹⁸F-FBEM-SRGD, the U87MG and MDA-MB-435 tumor uptakes were 1.27 and 1.04 %ID/g at 60 min after injection, respectively, whereas the liver and kidney uptake were much higher than that of the tumors. Up to 4 h after injection, there is still a fair amount of activity accumulated in the abdomen. 18F-FBEM-SRGD was excreted through both liver and kidneys (Fig. 5A). For ¹⁸F-FBEM-SRGD2, most radioactivity in nontargeted tissues was cleared at 60 min after injection. The uptakes in the U87MG, MDA-MB-435, kidneys, liver, and lung at 60 min after injection were 2.14, 2.11, 4.00, 1.46, and 0.58 %ID/g, respectively. Time–activity curves showed that this tracer excreted predominantly through the renal route (Fig. 5B). When the microPET images of these 2 radiotracers were compared, ¹⁸F-FBEM-SRGD2 had a much better tumor-to-background contrast and higher tumor uptake, which makes it more suitable for future clinical studies.

Static microPET scans with blocking were then performed for $^{18}\text{F-FBEM-SRGD2}$ on U87MG tumor-bearing mice (Fig. 6). When coinjected with 10 mg/kg of c(RGDyK), the tracer uptake in the U87MG tumor dropped from 2.21 %ID/g to 0.94 %ID/g at 30 min after injection and from 1.73 %ID/g to 0.44 %ID/g at 60 min after injection, which are essentially at the background level. Successful blocking again confirmed the integrin $\alpha_{v}\beta_{3}$ –specific binding of the radiotracer $^{18}\text{F-FBEM-SRGD2}.$

Metabolism of ¹⁸F-FBEM-SRGD2

The metabolic stability of ¹⁸F-FBEM-SRGD2 was also determined in mouse blood and urine samples and in liver,

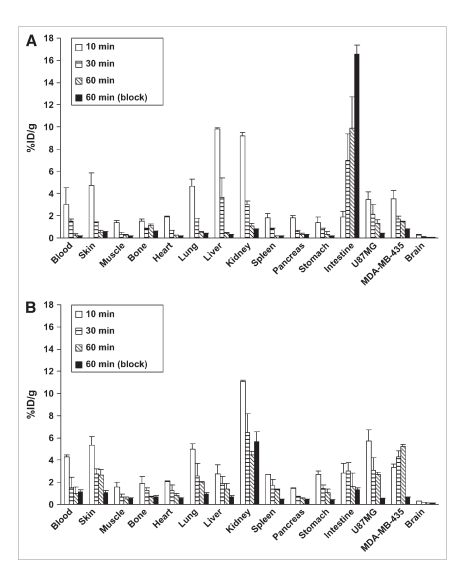


FIGURE 3. Biodistribution of 18 F-FBEM-SRGD (A) and 18 F-FBEM-SRGD2 (B) in athymic nude mice bearing both U87MG and MDA-MB-435 tumors at 10, 30, and 60 min after injection (n=3). Biodistribution of both tracers at 60 min after injection when coinjected with 10 mg/kg mice body weight of c(RGDyK) is also shown.

kidneys, and U87MG tumor homogenates at 60 min after tracer injection. The extraction efficiency for all organs was >90% (Table 1). The lowest elution efficiency was found for the U87MG tumor and the liver. HPLC analysis results of the soluble fractions of all samples are shown in Figure 7. The percentage of intact tracer was between 41.7% (urine) and 85.8% (kidney). Although we did not identify the composition of the metabolites, we found that all metabolites came out earlier from the HPLC column than the parent compound. The major metabolite peak was found at about 13 min for almost all organs except the kidney. No defluoridation of ¹⁸F-FBEM-SRGD2 was observed, as there is no bone uptake visible in all of the microPET scans.

DISCUSSION

¹⁸F labeling is generally achieved through 3 types of functional groups: amino group, carboxylic acid group, and sulfhydryl group. Most known ¹⁸F-labeling reagents for peptides and protein react with the primary amino groups at the N terminus or the lysine side chain. This can be

achieved through active esters, aldehydes, imidates, or azido functionalities. ¹⁸F-SFB is probably the most-oftenused active ester for ¹⁸F labeling via an acylation reaction (12,15,21). Reductive amination using 4-18F-fluorobenzaldehyde (18F-FBA) (28), oxime formation using 18F-FBA (23,29), imidation reaction using 3-18F-fluoro-5-nitrobenzimidate (¹⁸F-FNB) (30), photochemical conjugation using 4-azidophenacyl ¹⁸F-fluoride (¹⁸F-APF) (22), and alkylation reactions using 4-18F-fluorophenacyl bromide (18F-FPB) have been reported earlier (30). The major concern for protein labeling using these reagents is the possible interference with biologic activity: modification of one or more lysines located at or near the active site could reduce the binding affinity through steric hindrance if a bulky group is added. ¹⁸F labeling of peptide or protein via the carboxylic acid group at the C terminus or internal glutamic/aspartic acid side chain is less common. Only 2 amines, 1-[4-(18Ffluoromethyl)benzoyl]aminobutane-4-amine (18F-FMBAA) (31) and 4-18F-fluorophenylhydrazine (18F-FPH) (32), have been reported for these amidation reactions.

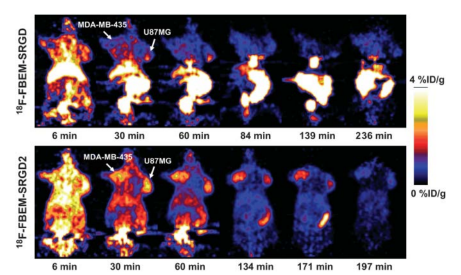


FIGURE 4. Dynamic microPET scans using both radiotracers at different time points in a mouse bearing both U87MG and MDA-MB-435 tumors. Ten-minute static scans at several later time points were also conducted to complete the tracer kinetic study.

Thiol-reactive agents have been used to modify peptides and proteins at specific sites, providing high chemoselectivity as compared with amine and carboxylate-reactive reagents (32,33). The disulfide bonds of a protein can be reduced to enable modification using thiol-specific reagents (34–36). More recently, site-directed mutagenesis was used to place cysteine residues on the surface of proteins to

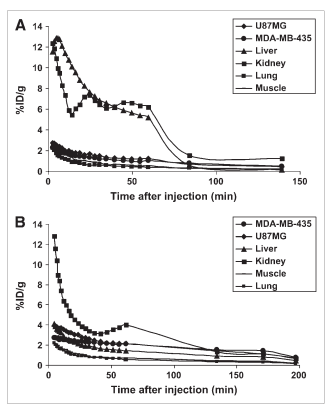


FIGURE 5. Time-activity curves of ¹⁸F-FBEM-SRGD (A) and ¹⁸F-FBEM-SRGD2 (B) obtained from microPET scans. The inflection point for tracer clearance is most likely due to the slower metabolism during the dynamic scan when mice were under anesthesia and body temperature was lowered.

provide reactive sulfhydryl groups (*37,38*). Several thiol-reactive ¹⁸F-synthons have been described (*24–26*), all of which bear a maleimide group allowing for thiol-specific Michael addition reaction. The total synthesis time (100–150 min) and radiochemical yield (10%–20% non-decay corrected) of these synthons are comparable to ¹⁸F-FBEM. However, no in vivo data have been reported on tracers synthesized using these prosthetic groups.

We have labeled c(RGDyK) with 18 F using 18 F-SFB as a prosthetic group (18). The labeling yield was reasonably good for in vivo imaging applications. The resulting 18 F-FB-RGD had good tumor-to-blood and tumor-to-muscle ratios but also rapid tumor washout and unfavorable hepatobiliary excretion, making it suitable only for visualizing lesions above the liver (e.g., breast cancer, head and neck cancer, and brain tumor). Because the bent conformation of c(RGDyK) has been optimized to fit into the deep cleft between the α - and β -units of integrin $\alpha_v\beta_3$, it is unlikely that one can further improve integrin affinity and selectivity of the monomeric RGD peptide by fine tuning the pentapeptide configuration (39). Therefore, the polyvalency effect has been applied to develop dimeric and multimeric RGD peptides, with repeating cyclic pentapeptide

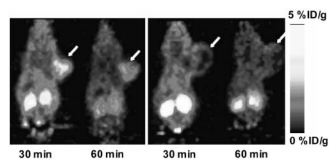


FIGURE 6. Ten-minute static microPET scans of U87MG tumor-bearing mice (arrows) injected with 3.7 MBq of ¹⁸F-FBEM-SRGD2. (Left) Control mouse. (Right) Blocking with 10 mg/kg mouse body weight of c(RGDyK).

TABLE 1Extraction and Elution Efficiency Data and HPLC Analysis of Soluble Fraction of Tissue Samples at 60 Minutes
After Injection of ¹⁸F-FBEM-SRGD2

Organ or tissue	Extraction efficiency (%)	Elution efficiency (%)	Intact fraction (%)
Blood	91.5	94.9	80.7
Urine	100.0	99.3	41.7
Liver	91.4	66.2	59.7
U87MG	91.0	66.2	77.5
Kidney	93.5	86.0	85.8

units connected by glutamates (11,12,14,19,40). We have found that ¹⁸F-FRGD2 (¹⁸F-FB-E[c(RGDyK)]₂) had predominant renal excretion and almost twice as much tumor uptake in the same animal model as compared with the monomeric tracer ¹⁸F-FB-RGD (11,12). The synergistic effect of polyvalency and improved pharmacokinetics may be responsible for the excellent imaging characteristics of ¹⁸F-FRGD2. At late time points when most of the nonspecific binding had been cleared, the tumor-to-background ratio also had a linear relationship with the tumor integrin levels, thus making it possible to quantify the tumor integrin expression level in vivo with static PET scans using 18F-FRGD2. We are currently in the process of translating ¹⁸F-FRGD2 into clinical trials. In parallel, we developed ¹⁸F-FBEM as a prosthetic group for dimeric RGD peptide labeling as well as for protein or engineered antibody labeling through site-specific Michael addition with the sulfhydryl group. The reaction between ¹⁸F-FBEM and the thiolated RGD peptides was virtually quantitative. Although ¹⁸F-FBEM-SRGD2 demonstrated integrin specificity, as evidenced by effective inhibition of tumor activity accumulation in the presence of a blocking dose of integrin $\alpha_v \beta_3$ antagonist c(RGDyK), whether it is able to quantify integrin expression in vivo remains to be determined in future studies. It is also worth noting that addition of the thiolated RGD peptides to ¹⁸F-FBEM generates a new

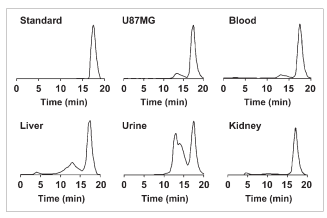


FIGURE 7. Metabolic stability of ¹⁸F-FBEM-SRGD2 in mouse blood and urine samples and in liver, kidneys, and U87MG tumor homogenates 60 min after injection. HPLC profile of tracer itself (Standard) is also shown.

asymmetric center, resulting in 2 diastereomeric products. Because we only observed one sharp peak in the analytic HPLC for both ¹⁸F-FBEM-SRGD and ¹⁸F-FBEM-SRGD2, it is very likely that such a small change was not detectable by the HPLC system used.

This study demonstrated that ¹⁸F-FBEM could be used to efficiently label peptides through the sulfhydryl group. The major advantage of ¹⁸F-FBEM lies in the fact that it can be applied to label a variety of peptides, proteins, or oligonucleotides containing sulfhydryl groups. Because most proteins contain Cys residues, whereas those that do not can be easily engineered to incorporate a Cys residue without compromising the biologic activity, it is expected that ¹⁸F-FBEM will have wide applications in the near future for development of novel tracers for in vivo PET.

CONCLUSION

The objective of the present work was to develop a new thiol-reactive ¹⁸F-labeling reagent for the prosthetic labeling of peptides and proteins via selective conjugation with a sulfhydryl group. ¹⁸F-FBEM was thus incorporated with thiolated monomeric and dimeric RGD peptides via efficient alkylation of the free thiol group though the maleimido function. The advantage of labeling at the sulfhydryl group using ¹⁸F-FBEM over labeling at the primary amino group using ¹⁸F-SFB was confirmed. The dimeric RGD peptide labeled through the ¹⁸F-FBEM strategy showed high integrin affinity in vitro and effective tumor targeting in vivo. The fast tracer clearance, good tumor-to-background contrast, relatively good metabolic stability, and favorable pharmacokinetics of ¹⁸F-FBEM-SRGD2 promise further investigation of this tracer in both preclinical and clinical settings for documenting tumor integrin expression (such as the correlation between the tumor uptake and the integrin $\alpha_v \beta_3$ expression level in vivo). ¹⁸F-FBEM also provides a general method of labeling thiol-containing peptides, proteins, antibodies, as well as 5'-thio-functionalized oligonucleotides in high radiochemical yield and high specific activity for successful PET applications.

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