Click-Chemistry (CuAAC) Trimerization of an $\alpha_v\beta_6$ Integrin Targeting Ga-68-Peptide: Enhanced Contrast for in-Vivo PET Imaging of Human Lung Adenocarcinoma Xenografts

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$\alpha_v\beta_6$ Integrin is an epithelial transmembrane protein that recognizes latency-associated peptide (LAP) and primarily activates transforming growth factor beta (TGF-$\beta$). It is overexpressed in carcinomas (most notably, pancreatic) and other conditions associated with $\alpha_v\beta_6$ integrin-dependent TGF-$\beta$ dysregulation, such as fibrosis. We have designed a trimeric Ga-68-labeled TRAP conjugate of the $\alpha_v\beta_6$-specific cyclic pentapeptide SDM17 (cyclo[RGD-Chg-E]-CONH$_2$) to enhance $\alpha_v\beta_6$ integrin affinity as well as target-specific in-vivo uptake. Ga-68-TRAP (SDM17)$_3$ showed a 28-fold higher $\alpha_v\beta_6$ affinity than the corresponding monomer Ga-68-NOTA-SDM17 (IC$_{50}$ of 0.26 vs. 7.4 nM, respectively), a 13-fold higher IC$_{50}$-based selectivity over the related integrin $\alpha_v\beta_8$ (factors of 662 vs. 49), and a threefold higher tumor uptake (21 vs. 0.66%ID/g) in biodistribution experiments with H2009 tumor-bearing SCID mice. The remarkably high tumor/organ ratios (tumor-to-blood 11.2; -to-liver 8.7; -to-pancreas 29.7) enabled high-contrast tumor delineation in PET images. We conclude that Ga-68-TRAP(SDM17)$_3$ holds promise for improved clinical PET diagnostics of carcinomas and fibrosis.

Introduction

Integrins are a family of 24 heterodimeric transmembrane receptors, each comprising one out of 18 $\alpha$- and one out of eight $\beta$-subunits, which are expressed by almost all animal cells and fulfill a wide variety of biological purposes and functions. Integrins are primary adhesion receptors that facilitate the binding of cells to extracellular matrix (ECM) proteins, but they are also involved into a variety of signaling processes. A puzzling complexity is observed here because fundamentally different functions are sometimes promoted by binding to the same peptide sequences. For example, $\alpha_v\beta_6$ integrin shares its ability to recognize the arginine-glycine-aspartate (RGD) peptide sequence contained in many ECM proteins and clotting factors with seven other integrins$^{[1]}$ (such as the popular and well-characterized vitronectin receptor $\alpha_v\beta_3$ integrin), but its key function is a fundamentally different one, namely, activation of transforming growth factors $\beta$1 and $\beta$3 (herein further abbreviated as TGF-$\beta$)$^{[2]}$.

TGF-betas are a pleiotropic family of phylogenetically old cytokines, a class of signaling proteins that are produced by almost any mammalian cell type.$^{[3]}$ They are secreted into the extracellular space, albeit not in a freely diffusible form capable of binding to the respective receptors, but as complexes with a temporary inhibitor called latency-associated peptide (LAP). To exert its biological signaling functions, TGF-$\beta$ must be released from this aggregate. This occurs predominantly by binding of the extracellular domain of $\alpha_v\beta_6$ integrin to a RGD motif in LAP. Within the cell, the $\beta_6$ subunit is connected to the actin filaments of the cytoskeleton and basically acts as a rope to exert a pulling force on LAP, by which the latter is deformed and loses its ability to bind TGF-$\beta$. As a result, TGF-$\beta$ is released and influences gene expression and protein synthesis of adjacent cells.$^{[3]}$

$\alpha_v\beta_6$ Integrin expression and -signaling is thus an integral part of the TGF-$\beta$-mediated cell communication mechanism controlling tissue development and homeostasis, whose dysregulation is related to a wide variety of diseases.$^{[2]}$ Overexpression of $\alpha_v\beta_6$ integrin has particularly interesting implications in terms of cancer.$^{[4]}$ For example, TGF-$\beta$ acts as a growth suppressor by regulating the transcription of certain growth-promoting genes. Tumor cells can however acquire a reduced...
TGF-β sensitivity as a result of downstream mutations in the signaling pathway,
meaning that their growth is no more inhibited by increased TGF-β concentrations. Such tumor cells can benefit from a high TGF-β concentration in their surrounding because it limits the growth of normal cells and facilitates the tumor to infiltrate healthy tissue. Consequently, a high αvβ6 integrin expression is frequently observed on carcinoma cells, particularly on pancreatic carcinoma as well as its metastases and precursor lesions (PanIN). Likewise, αvβ6 integrin expression is connected to other carcinogenic processes associated with elevated TGF-β levels, such as epithelial mesenchymal transition (EMT) and metastasis, deposition of ECM proteins and activation of fibroblasts, or the suppression of T-cell-mediated immunosurveillance of tumor cells and resistance to immune checkpoint inhibitors. Hence, αvβ6 integrin represents a promising biomarker for the invasive potential and malignancy of carcinomas. This is however only part of a bigger picture, as dysregulation of the TGF-β pathway is involved into many other pathogenic processes, for example, inflammation and fibrosis of the lung.

Targeting αvβ6 integrin might therefore possess a high value for fundamental research and clinical reasoning. In particular, the analysis of spatiotemporal expression patterns in living subjects by means of quantitative noninvasive 3D imaging might guide the way towards a deeper understanding of the activation of TGF-β in the course of various diseases, or enable a better prognosis for carcinoma patients. To this end, we earlier reported SDM17 (cyclo-[(RGD-Chg-E)-CONH]2), a small, cyclic pentapeptide as a selective ligand for this target. Initially, we successfully used a radiolabeled conjugate of this peptide, 68Ga-NOTA-SDM17 (Figure 1), to demonstrate the feasibility of in-vivo imaging of αvβ6 integrin expressing subcutaneous xenografts in mice by means of positron emission tomography (PET). However, this compound exhibited an unsatisfyingly low affinity, resulting in a comparably low uptake in the tumor. Based on the observation that the affinity and the tumor uptake of ligands for RGD-binding integrins can be substantially increased by multimerization, we hypothesized that a trimeric conjugate of SDM17 would exhibit an improved uptake and imaging performance. Hence, we investigated the trimer 68Ga-TRAP(SDM17)3 (Figure 1) in terms of its suitability for in-vivo PET imaging of αvβ6 integrin.

Results and Discussion

Synthesis

In the monomer NOTA-SDM17, a single SDM17 cyclopeptide unit is linked via p-aminobenzoic amide to the 1,4,7-triazacyclononane-1,4,7-triacetic acid (NOTA) chelator via amide bonds (Figure 1). It is worth noticing that the actual chelating unit is NOTA but NOTA-monoamide, since the linker connects as an amide to one of the N pendant arms of NOTA.

In contrast, the trimer was elaborated on the basis of the triazacyclononane-triphosphinate (TRAP) chelator scaffold employing click chemistry (CuAAC). For this purpose, SDM17 was decorated on-resin with 4-pentyonic acid, employing ultrasound-enhanced coupling protocol which enables reaction times as short as 5 min. Following simultaneous cleavage off the resin/deprotection and purification, 3.3 equivalents of the threefold azide-decorated TRAP building block TRAP(azide)3 was reacted with the CuAAC trimerization was carried out applying a robust and convenient one-pot protocol which includes the final removal of any TRAP-bound copper by transchelation with NOTA.

Radiochemistry

A radiolabel for PET imaging was introduced by complexation of the positron emitter gallium-68 (t1/2 = 68 min) which is conveniently available from 68Ge/68Ga radionuclide generators (small, commercially available benchtop devices acting as long-lived regenerative sources for 68Ga in dilute HCl) and which is particularly suited for straightforward and cost-efficient development of PET radiopharmaceuticals. The TRAP structural motif is characterized by particularly robust and efficient
$^{68}$Ga labeling\cite{22,23} because its rapid Ga$^+$ incorporation reaction is comparably insensitive towards high concentrations of frequently occurring metal ion contaminants in generator eluates and $^{68}$Ga labeling solutions, such as Fe$^{3+}$, Zn$^+$, or Cu$^+$.\cite{24} Accordingly, $^{68}$Ga labeling of 5 nmol of TRAP(SDM17)$_1$ in a total reaction volume of approximately 1.5 mL for 2–3 min at pH 2 and 95 °C reliably afforded $^{68}$Ga-TRAP(SDM17)$_1$ in >96% radiochemical yield with a purity of >99.5% according to radio-TLC (Figure S3 in the Supporting Information).

Although being somewhat less efficient than TRAP,\cite{15,22} NOTA is also known as an excellent $^{68}$Ga$^+$ chelating system. However, we found that a relatively high concentration of NOTA-SDM17 (10 nmol in 1.5 mL, pH 3, 95 °C, 5 min) was necessary to obtain acceptable radiolabeling yields (averagely 76%, range 70–81%, $n = 4$). Actually, this is worse than what would be expected for an average DOTA peptide ($\text{DOTA} = 1,4,7,10$-tetraazacyclododecane-1,4,7,10-tetraacetic acid), as approximate amounts of only 5, 1, or 0.2 nmol of DOTA-, NODAGA-, or TRAP conjugates, respectively, are required under otherwise similar conditions to achieve a comparable percentage of $^{68}$Ga incorporation.\cite{22} Moreover, we found that the resulting radiopharmaceutical still contained a small fraction of noncomplexed $^{68}$Ga$^+$ (Figures S4 and S5); this had a noticeable impact on biodistribution (see below).

Apparently, the NOTA structural motif only possesses a high $^{68}$Ga$^+$ chelation efficiency if none of its N-acetic acid substituents is used for conjugation, that is, all three carboxylates are available for coordination. This might appear somewhat counterintuitive, since sacrificing one carboxylate for conjugation does not substantially impair the functionality of its larger congener DOTA.$^{22}$ This, however, works only because the resulting DOTA-monoamide motif retains three negative formal charges to counterbalance the +3 charge of the metal ion, keeping it capable of rapidly forming kinetically inert $^{68}$Ga$^{III}$ complexes.$^{[27]}$ Our observations suggest that for an optimal radiolabeling performance of NOTA in bioconjugates, a separate pendant arm with an additional functional group dedicated to conjugation is indispensable, preferably attached to one of its backbone or N-acetic acid methylene groups, such as in its bifunctional derivative NODAGA.$^{[28]}$ Conjugation of NOTA by amide bonding on one of the acetic acid $N$-substituents, essentially resulting in a NOTA-monoamide chelator structure, yields constructs with poor $^{68}$Ga labeling properties and is thus not recommendable, despite such compounds have even been translated into humans.$^{[29]}

### In-vitro characterization

Consistent with previous experience,$^{[30]}$ $^{68}$Ga-TRAP(SDM17)$_1$ possesses an approximately 28-fold higher $\alpha_\text{Ga} / \beta_\text{Ga}$ integrin affinity ($IC_{50} = 0.26$ nM) than the monomer $^{68}$Ga-NOTA-SDM17 (7.4 nM) as determined by ELISA on immobilized integrins$^{[31]}$ (Table 1). Interestingly, the trimer also exhibits a 13-fold higher selectivity over the functionally related integrin subtype $\alpha_\text{V} / \beta_\text{3}$ this underscores the benefits of trimerization. As the hydrophilicities (log $D_{2,4}$) of both compounds are virtually identical, any differences regarding their behavior in a biological context should therefore be considered a result of altered affinity or molecular size.

### In-vivo studies

In-vivo characterization of the radiolabeled compounds by PET and biodistribution studies was done using SCID mice bearing subcutaneous xenografts of the human lung adenocarcinoma

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### Table 1. Integrin affinities (expressed as 50% inhibition concentrations, $IC_{50}$), $IC_{95}$-based selectivities, and octanol/PBS distribution coefficients (log $D_{2,4}$)

| Compound               | $IC_{50}$ (95% confidence interval, nM) | Selectivity for $\alpha_\text{Ga} / \beta_\text{Ga}$ over $\alpha_\text{Ga} / \beta_\text{Ga}$ | log $D_{2,4}$ |
|------------------------|----------------------------------------|-----------------------------------------------|----------------|
| $^{68}$Ga-TRAP(SDM17)$_1$ | 0.26 (0.21–0.40)                      | 662                                          | $-4.08 \pm 0.07$ |
| $^{68}$Ga-NOTA-SDM17   | 7.4 ± 1.1$^*$                          | 49                                           | $-3.91 \pm 0.07$ |
| SDM17                  | 1.3 ± 0.2$^*$                          | 134                                          | n.a.           |

[a]: $IC_{50}$ data for Ga-NOTA-SDM17 and SDM17 were taken from the literature and are shown as reported therein.$^{[12]}$
null
Figure 5 shows that the higher tumor uptake of $^{68}$Ga-TRAP(SDM17)$_3$ enables a clear visualization of the subcutaneous xenograft, whereas the monomer $^{68}$Ga-NOTA-SDM17 is not capable of delineating the same lesion with satisfying contrast. Of note, the trimer also shows the advantage of much higher tumor-to-background ratios, which is of practical relevance for a possible clinical application. For example, its high tumor-to-pancreas and -to-liver ratios (29.7 and 8.7, respectively; Figure 4 and Table S1) appear much more suitable for detection of intrapancreatic or intrahepatic lesions, such as primaries and metastases of pancreatic carcinoma, as compared to the monomer (4.4 and 2.0, respectively; Figure 4 and Table S2).

Conclusion

$^{68}$Ga-TRAP(SDM17)$_3$ was synthesized by click-chemistry-driven trimerization of the $\alpha_\beta_\gamma$ integrin-selective cyclic pentapeptide SDM17 on the TRAP chelator. Compared to the monomer $^{68}$Ga-NOTA-SDM17, the trimer $^{68}$Ga-TRAP(SDM17)$_3$ showed a higher $\alpha_\beta_\gamma$ integrin affinity as well as a higher uptake and longer retention in $\alpha_\beta_\gamma$-positive H2009 tumor xenografts in mice. A high PET image contrast in living subjects (i.e., high tumor-to-background ratios) was achieved with $^{68}$Ga-TRAP(SDM17)$_3$, which encourages the translation into clinics. Notably, several recent reports underscored the potential clinical value of $\alpha_\beta_\gamma$-integrin PET imaging for diagnosis of various carcinomas$^{41–46}$ as well as pulmonary fibrosis$^{46}$ in humans. Hence, we are confident that in view of its favorable preclinical data, $^{68}$Ga-TRAP(SDM17)$_3$ can contribute to the improvement of healthcare schemes for diagnosis and therapy of carcinomas as well as other conditions associated with $\alpha_\beta_\gamma$ integrin dependent dysregulation of the TGF-$\beta$ signaling pathway, such as fibrotic diseases.

Experimental Section

Materials: Unless otherwise noted, all commercially available reagents and solvents were of analytical grade and were used without further purification. Protected amino acids were purchased from IRIS Biotech (Germany), Cu(OAc)$_2$·H$_2$O, 4-pentyneic acid, disopropylamine (DIPEA) and sodium ascorbate were purchased from Sigma-Aldrich. $1,4,7$-triazacyclononane-$1,4,7$-triacetic acid (NOTA) was purchased from Chematech (Dijon, France). HATU was obtained from Bachem Holding AG (Bubendorf, Switzerland). HOBr hydrate was obtained from Carbolution (St. Ingbert, Germany). For all radiochemical works, Tracepur® water (Merck) was used. TRAP (azide)$^{[41]–[45]}$ and o-NBS-L-Lys(Fmoc)-OH$^{[47]}$ were synthesized as described previously.

Instrumentation: The synthesis of SDM17-pentyenic amide was carried out in an ultrasonic bath SONOREX RK 52 H (interior dimensions 150 × 140 × 100 mm and operating volume 1.2 L) by BANDELIN electronic (Germany), equipped with timer control for 1–15 min and continuous ($\infty$) operations and built-in heating control (30–80 °C thermostatically adjustable). Semi-preparative reversed-phase HPLC was performed by using a Shimadzu system, consisting of two LC-20AP quaternary low-pressure gradient pumps, a SPD-M30A photodiode array detector, and a CBM-20 A system controller. Separations were performed by using a YMC-Pack ODS-A, 5 $\mu$m, 250 × 20 mm C$_{18}$ column. Analytical HESI-HPLC-MS (heated electrospray ionization mass spectrometry) was performed on a LCQ Fleet (Thermo Scientific) with a connected UltiMate 3000 UHPLC focused (Dionex) on C$_{18}$ columns: S1: Hypersil Gold AQ 175 Å, 3 $\mu$m, 150 × 2.1 mm (for 8 or 20 min measurements); S2: Accucore C18, 80 Å, 2.6 $\mu$m, 50 × 2.1 mm (for 5 min measurements; Thermo Scientific). Linear gradients (5%–95% acetonitrile content) with water (0.1% v/v formic acid) and acetonitrile (0.1% v/v formic acid) were used as eluents. Centrifugation was done with a Heraeus Biofuge 13 benchtop centrifuge. Activities were quantified with a Capintec CRC 15R dose calibrator. Small activities in tissue samples etc. were measured using a PerkinElmer Wizard 2 4200 automatic gamma counter. Radio-TLCs were evaluated using a Bioscan radio-TLC scanner, consisting of B-MS-1000 scanner, B-EC-1000 detector with a B-FC-3600GM tube.

Synthesis: SDM17-pentyenic amide: SDM17 functional monomer was synthesized on solid support by conventional Fmoc/Bu approach, employing an ultrasound-assisted solid-phase peptide synthesis (US-SPPS) protocol.$^{[47]}$ Rink amide resin (545 mg, 0.3 mmol) was functionalized with o-NBS-L-Lys(Fmoc)-OH (363 mg, 0.6 mmol, 2 equiv) using HBTU (227 mg, 0.6 mmol, 2 equiv) and HOBr (92 mg, 0.6 mmol, 2 equiv) as coupling partners, and DIPEA (209 $\mu$L, 1.2 mmol, 4 equiv) as base, in DMF (3.5 mL). The mixture of the resin was added to the resin in a SPPS reactor and then ultrasonicated for 5 min before washing. Fmoc protection was carried out by irradiating the resin with ultrasound in the presence of a 20% piperidine solution in DMF (2 × 1 min). The linear aminoacidic sequence was elongated by iterative cycles of the aforementioned amide bond coupling reactions and Fmoc deprotection; the completion of each step was qualitatively determined by Kaiser test or TNBS test. After loading the last amino acid, the resin-bound peptide underwent a Tsuji-Trost-mediated allyl ester removal on the glutamic acid side chain. The resin was treated with a solution of tetrakis(triphenylphosphine)palladium(0) (35 mg,
0.03 mmol, 10% mol) and DMBA (234 mg, 1.5 mmol, 5 equiv) in anhydrous THF (5 mL) for 1 h at rt under argon, and this procedure was repeated once. After being washed with DMF (3 × 1 min) and dichloromethane (3 × 1 min), the resin was suspended in a 0.06 M solution of potassium N,N-diethylthiocarbamate in DMF (38 mg in 3 mL of solvent) for 15 min in order to completely remove catalyst traces, and this procedure was repeated twice. At this stage, the α-amino group of the Arg residue was released when the resin was washed with MeOH (3 × 1 min) and CH₂Cl₂ (3 × 1 min). Final functionalization with the alkyln-bearing building block was carried out in a DMF solution (3.5 mL) of pentaerythritol (59 mg, 0.6 mmol, 2 equiv), HBTU (227 mg, 0.6 mmol, 2 equiv) and HOBt (92 mg, 0.6 mmol, 2 equiv) in the presence of DIPEA (313 µL, 1.8 mmol, 6 equiv) in DMF (5 mL) and allowing the resin to shake for 2 h. Next, the α-amino group on C-terminal lysine residue was released by removing the ortho-nitrobenzenesulfonyl (o-NBS) protecting group. This deprotection was performed by adding a clear solution (4 mL) of thiophenol in dry DMF (5% v/v) in the presence of 1.5 equiv. (relative to thiophenol) of ultrapure K₂CO₃. The obtained suspension was miniaturized by sonication and centrifuged, then the clear supernatant was added to the resin, which was allowed to shake for 10 min. This procedure was repeated a further two times and then the resin was washed exhaustively with DMF (3 × 1 min), MeOH (3 × 1 min) and CH₂Cl₂ (3 × 1 min). Final functionalization with the alkynyl-bearing building block was carried out in a DMF solution (3.5 mL) of pentaerythritol (59 mg, 0.6 mmol, 2 equiv), HBTU (227 mg, 0.6 mmol, 2 equiv) and HOBt (92 mg, 0.6 mmol, 2 equiv) in the presence of DIPEA (209 µL, 1.2 mmol, 4 equiv) and irradiating with ultrasound for 5 min.

The resin was washed with DMF (2 × 1 min), CH₂Cl₂ (2 × 1 min), and diethyl ether (3 × 1 min), and the peptide was cleaved from the solid support using a solution of TFA/TIS (95:5, 3 mL) for 3 h at room temperature. The suspension was filtered and the crude product precipitated from the TFA solution by diluting to 35 mL with cold diethyl ether, and then centrifuged (4400 g, 15 min). The supernatant was removed, and the precipitate was suspended again in 35 mL ether as described above. The wet solid was dried for 1 h under vacuum, re-dissolved in water/acetonitrile (9:1) and purified by RP-HPLC (solvent A: water + 0.1% TFA; solvent B: acetonitrile + 0.1% TFA; from 10 to 90% of solvent B over 20 min, flow rate: 10 mL min⁻¹). Product-containing fractions were identified by ESI-MS, concentrated in vacuo, and lyophilized. The product was characterized by analytical RP-HPLC (solvent A: water + 0.1% TFA; solvent B: acetonitrile + 0.1% TFA; from 10 to 90% of solvent B over 20 min, flow rate: 1 mL min⁻¹) and HRMS (ESI-MS) spectrometry. Overall yield: 179 mg (65%), purity: >95%, t₁/₂ = 12.45 min. MW (calcd for C₁₁₂H₁₈₁O₁₄N₃): 804.43. HRMS (ESI-MS): m/z = 804.43506 [M + H]+ (theoretical value: 804.43626; for MS spectra, see Figure S1).

TRAP(SDM17): The trimer was synthesized employing a previously established method.[24] SDM17-pentameric amide (161.1 mg, 20.0 µmol, 3.3 equiv) was added to a solution of TRAP(azide) (5.0 mg, 6.1 µmol, 1 equiv) and sodium ascorbate (60 mg, 303 µmol, 50 equiv) in a mixture of water and tert-butanol (3:1 by volumes, 400 µL). Copper(II) acetate hydrate (1.45 mg, 7.28 µmol, 1.2 equiv) was added, whereupon a brown precipitate formed immediately. Upon vortexing, the solution turned to a transparent green. The solution was allowed to react for 1 h at 60 °C without stirring. Then, all Cu species were sequestered from the TRAP(SDM17), compound and the reaction solution by addition of 1,4,7-triazacyclononane-1,4,7-triacetic acid (NOTA) (55 mg, 180 µmol, 30 equiv) dissolved in water (1.5 mL), adjusting to pH 2.2 by using 1 M aq. HCl, and reacted for 1 h at 60 °C. HPLC-MS was used in all steps for monitoring of reaction progress. TRAP(SDM17), was obtained as a colorless solid with a yield of 37% (7.2 mg, 2.2 µmol), RP-HPLC (gradient: 3–45% MeCN in water, both containing 0.1% trifluoroacetic acid, in 20 min, flow rate: 20 mL min⁻¹): t₁/₂ = 16.6 min. MW (calcd for C₁₁₂H₁₈₁O₁₄N₃P₃): 3235.63. MS (ESI, positive mode): m/z = 1618.8 [M + 2H]⁺, 1080.2 [M + 3H]³⁺, 810.3 [M + 4H]⁴⁺, 648.6 [M + 5H]⁵⁺, 540.6 [M + 6H]⁶⁺ (theoretical values: 1618.8, 1079.5, 800.9, 641.8, 540.3; for MS spectra, see Figure S2).

Affinity assays: The integrin affinities were determined by a solid-phase binding assay, applying a previously described protocol.[25] Briefly, flat-bottom 96-well enzyme-linked immunosorbent assay (ELISA) plates (BRAND, Wertheim, Germany) were coated with recombinant human LAPTGF-β (1) in carbonate buffer (pH 9.6, 35 mM Na₂CO₃, 35 mM NaHCO₃, pH 9.6) at 4 °C overnight. After washing the plates with PBS-T buffer (phosphate-buffered saline/Tween20), free binding sites were blocked with incubation with TBS buffer (Tris-buffered saline). Dilution series of non-radioactive Ga-TRAP(SDM17), (20 µM to 6.4 nM) were prepared and incubated in 1:1 mixtures with the respective integrin. Surface-bound integrin was detected by subsequent incubation with a specific primary antibody and a secondary peroxidase-labeled antibody (anti-mouse IgG-POD, Sigma–Aldrich). After addition of the dye SeramunBlau (Seramun Diagnostik, Heidelberg, Germany) and quenching of the reaction by addition of 3 M H₂SO₄, the absorbance at λ = 450 nm was measured with a microplate reader (Tecan Genesis, Männedorf, Switzerland). The IC₅₀ value for each compound was determined in duplicate and the inhibition curves were analyzed by using OriginPro 9.0 software. The measured IC₅₀ values were referenced to the activity of the internal standard RTDLDSLRT:[26] αIC₅₀ = 33 ± 6 nM, αIC₅₀ = 100 nM.

Radiochemistry: Fully automated Ga labeling was done in analogy to a previously described procedure[27] by using an accordingly programmed robotic system (GalEElit+, Scintomics, Fürstenfeldbruck, Germany) which carried out the following steps. A [68Ga]Ga-generator with TiO₂ matrix (Eckert & Ziegler, Berlin, Germany) was eluted with 0.1 M aq. HCl. A fraction containing the highest activity (1.4 mL, ca. 500 MBq) was collected in a 5 mL conical glass vial, containing 5 or 10 nmol of TRAP(SDM17), or NOTA-SDM17, respectively, as well as 50 or 100 µL, respectively, of a solution of 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) buffer (2.7 M, prepared from 14.4 g HEPES and 12 mL water), resulting in a labeling pH of approximately 2 or 3, respectively. The vial was heated for 5 min to 100 °C. Purification was done by passing the reaction mixture over a solid phase extraction cartridge (SepPak C8 light), which was purged with water (10 mL). The products were eluted with ethanol (0.3 mL), followed by an ethanol/water mixture (1:1 by volumes, 1 mL). The purity of the radiolabeled compounds was determined by HPLC-TLC, using silica impregnated glass fiber chromatography paper (ITLC® by Agilent) as stationary phase, and 0.1 M aq. sodium citrate or a mixture of 1 M aq. ammonium acetate and methanol (1:1 by volumes) as mobile phases.

To determine the n-octanol/PBS distribution coefficients (logDₐw), 650 µL octan-1-ol and 650 µL phosphate-buffered saline (PBS, pH 7.4) were combined in a 1.5 mL Eppendorf tube. Approximately 0.5 MBq of the radiolabeled compound was added and vortexed for 2 min at 2850 rpm using a Vortex Genie2 (Scientific Industries). The samples were centrifuged (11500g, 10 min), after which 100 µL of the organic phase and 10 µL of the aqueous phase were taken out and the activities of the aliquots were quantified in a γ-counter. The logD values were calculated from the quotients of the measured activities and given as averages ± standard deviation (n = 10).

Cell culture: H2009 human lung adenocarcinoma cells (CRL-5911; American Type Culture Collection (ATCC), Manassas, VA, USA) were cultivated as recommended by the distributor. Cells were subcultivated after trypsinization in a ratio of 1:2–1:5, two to three times weekly in culture medium (DMEM:F12, Biochrom FG4815; 5% fetal bovine serum, Biochrom Superior Biochrom S05615; 1% ITS-G, Thermofisher 41400045; 4.5 mM L-glutamine (final conc.), Biochrom K
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Conflict of Interest

The authors declare no conflict of interest.

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