In vivo optical imaging of CD13/APN-expression in tumor xenografts

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Abstract. The metalloexopeptidase CD13/aminopeptidase N (APN) has been shown to be involved in cancer angiogenesis, invasion, and metastasis. Therefore, a CD13/APN–targeted NGR-peptide was labeled with the cyanine dye Cy 5.5 and applied to image tumor xenografts with different APN-expression levels using both planar and tomographic optical imaging methods. In vitro, the peptide-dye conjugate showed a clear binding affinity to APN-positive HT-1080 cells, while negative MCF-7 cells and predosing with the free NGR-peptide revealed little to no fluorescence. In vivo, tumor xenografts (n ≥ 5) were clearly visualized by two-dimensional (2-D) planar fluorescence reflectance imaging (FRI) and three-dimensional (3-D) fluorescence mediated tomography (FMT) up to 24 h after injection. FMT also allowed us to quantify fluorochrome distribution in deeper tissue sections, showing an average fluorochrome concentration of 306.7±54.3 nM Cy 5.5 (HT-1080) and 116.0±18.3 nM Cy 5.5 (MCF-7) in the target tissue after 5 h. Competition with the free NGR-peptide resulted in a reduction of fluorochrome concentration in HT-1080 tumor tissue (195.3±21.9 nM; 5 h). We thus conclude that NGR-Cy 5.5 combined with novel tomographic optical imaging methods allows us to image and quantify tumor-associated CD13/APN expression noninvasively. This may be a promising strategy for a sensitive evaluation of tumor angiogenesis in vivo. © 2008 Society of Photo-Optical Instrumentation Engineers. [DOI: 10.1117/1.2839046]

Keywords: optical imaging; CD13/APN; NGR-Cy 5.5; fluorescence mediated tomography; fluorescence reflectance imaging.

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1 Introduction

Optical imaging is a desirable technique for in vivo cancer diagnostics since it is a highly sensitive, inexpensive imaging method that can potentially resolve relevant oncological target structures in vivo.1 With the injection of a fluorescent compound (fluorochrome), optical imaging allows specific tagging of particular receptors, antibodies, genes, or drugs, thereby measuring compound biodistribution and pharmacokinetics and hence leading to a better understanding of these processes.1–4 Combining new quantitative fluorescence imaging methods with target specific fluorochromes facilitates the visualization and quantification of specific receptor expression in vivo. Especially, fluorescence mediated tomography (FMT) as a newly emerging optical imaging technique has been shown to be capable of resolving fluorescence signatures three-dimensionally and quantitatively.3–9

The CD13/aminopeptidase N (APN) is a metalloexopeptidase that removes unsubstituted, N-terminal amino acids with...
neutral side chains (Ala > Phe > Leu > Gly) from peptides and amide or arylamide derivatives of amino acids. APN has been shown to be involved in cancer angiogenesis, invasion, and metastasis. A unique peptide that contains the NGR (asparagine-glycine-arginine)-motif binds to APN in tumor vasculature. Tumor homing peptides containing the NGR motive, such as CNGR and GNGRG, have been used for delivering cytokines, chemotherapeutic drugs, apototic peptides, and liposomes to a CD13 isoform, highly secreted on the surface of various tumor cells and tumor blood vessels. Taking this into account, we chose a cyclic NGR peptide labeled with the cyanine dye Cy 5.5 for measuring and quantifying tumor homing of the conjugate with FMT, thereby comparing this technique with conventionally used fluorescence reflectance imaging (FRI) methods. Specific tumor targeting of the cyclo-[Cys-Asn-Gly-Arg-Cys]-Gly-Lys-Cy 5.5 conjugate (NGR-Cy 5.5) was measured on HT-1080 fibrosarcoma and MCF-7 adenocarcinoma cell assays in vitro and on tumor xenografts in vivo.

2 Materials and Methods

2.1 Synthesis and Characterization of NGR-Cy 5.5

Peptides were purchased as trifluoroacetate (TFA) salts in >90% purity from Bachem Distribution Services GmbH, Weil am Rhein, Germany. Cy 5.5 monoreactiv NHS-ester was purchased from Amersham Biosciences Europe GmbH, Freiburg, Germany. All chemicals were used as purchased without further purification. Peptides were dissolved in aqueous bicarbonate buffer (0.1 M, pH 8.4; 1.3 mg/500 μL) at room temperature. To this, a solution of the dye in DMSO (1.9 mg/200 μL) was added and stirred for 60 min at room temperature in the dark. Purification was accomplished by gradient HPLC using a Knauer system with two K-1800 pumps, an S-2500 UV-detector and a RP-HPLC Nucleosil 100-5 C18 column (eluents: A; water, 0.1% TFA; eluent B: acetonitrile, 0.1% TFA). Elution times were 40 to 60 min, dependent on solution quantity, at a flow rate of 1 mL/min. The product fractions were pooled, lyophilized, redissolved in phosphate-buffered saline (PBS) at 1.0 mg/500 μL, and frozen at −20°C. The identities of the labeled products were confirmed by high-resolution electrospray mass spectrometry (HR-ES-MS) on a QUATTRO LCZ (Waters Micromass, Manchester, United Kingdom) spectrometer with a nanospray capillary inlet, m/z=557.5 [M]3−, 837.0 [M+H]−. The fully assembled tracer was assessed by fluorescence and photometry for spectral characteristics (Hitachi, F-4500 fluorescence spectrometer and U-3310 UV/VIS spectrophotometer, Tokyo, Japan).

2.2 Cell Lines and Reagents

Human fibrosarcoma HT-1080 (ATCC: CCL-1321, Manassas, Virginia) and human adenocarcinoma MCF-7 (ATCC: HTB-22, Manassas, Virginia) cells were cultured in RPMI-1640 or MEM (Invitrogen Corporation, San Diego, California) supplemented with 10% fetal calf serum, penicillin and streptomycin (Biochrom AG, Berlin, Germany). Cells were grown routinely in a monolayer culture at 37°C in a 5% CO2 humidified air atmosphere. All antibodies were specific for CD13/APN, clone WM15 antibodies (phycoerythrin- and unconjugated) were from Pharmingen (BD Biosciences, San Jose, California), and antibody H-300 for Western blotting and immunohistochemistry was from Santa Cruz Biotechnology (Santa Cruz, California).

2.3 FACS Analysis by Flow Cytometry

Cells were washed with PBS and subsequently harvested in 5 mL Versen-buffer (13.7 mM NaCl, 10 mM EDTA, 2.6 mM KCl, 8.1 mM NaH2PO4, and 1.4 mM KH2PO4, pH 7.2). Aliquots of 2.5×105 cells were blocked with mouse IgG in PBS/BSA (0.1%) for 15 min at 4°C, washed twice with PBS with Ca2+ and Mg2+, and resuspended in 150 μL binding buffer (150 mM NaCl, 10 mM MgCl2, 10 mM CaCl2, 0.5 mM MnCl2, pH 7.2, diluted 1:20 in PBS/BSA 0.1%). Antibodies (3 μg/mL) were added, and cells were incubated for 45 min at 4°C. Cells were washed, resuspended in PBS with 0.1% BSA, and analyzed on a Becton Dickinson FACSDiCalibur (BD Biosciences, San Diego, California).

2.4 Western Blot

Confluent (80 to 100%) HT-1080 and MCF-7 cells were treated with 0.5 mL trypsin and harvested by scraping in cold PBS. After two centrifugation steps (1500 rpm, 5 min, 4°C), cells were resuspended in lysis buffer [0.5% Tween 20, 50 mM Tris-HCl pH 8, 250 mM NaCl, 5 mM EDTA pH 8, 50 mM NaF, 0.5 mM Na3VO4, 0.9 mg/mL protease inhibitor cocktail (Sigma, St. Louis, Missouri)], and the protein concentration of each sample was determined. To concentrate the protein, a two-fold volume of cold acetone was added to each sample and placed in −20°C for 16 h, followed by centrifugation at 10,000 g, 4°C for 30 min. Protein pellets were dissolved in 100 μL of 6× sodium dodecyl sulfate (SDS) sample buffer (New England Biolabs, Beverly, Massachusetts). Following precipitation, proteins were electrophoretically separated in nonreducing 10% SDS-PAGE and transferred to PVDF membranes (Millipore Corporation, Bedford, Massachusetts). After blocking, immunoblots were incubated with CD13/APN polyclonal antibody H-300 and with peroxidase-conjugated goat anti-rabbit IgG secondary antibody (Sigma, St. Louis, Missouri). Peroxidase activity was revealed using ECL chemiluminescence (Amersham, Buckinghamshire, United Kingdom) and x-ray film, as described by the manufacturers. THP-1 cell lysat (BD Biosciences, San Diego, California) was used as a positive control.

Tumor tissue was frozen in liquid nitrogen, and 0.1 to 0.5 g of each tissue was homogenized by ultrasound using an Ultra-Turrax T8 (IKA-Labortechnik, Staufen, Germany). The homogenate was resuspended in 300 μL lysis buffer, additionally sonicated twice at 20 kHz for 5 s, and centrifuged for 30 to 60 min at 10,000 g to remove cell debris. The supernatant was taken for determining protein concentration following SDS-PAGE and Western blotting.

2.5 Aminopeptidase N Activity Assay

Surface aminopeptidase activity of intact cells was measured by plating 5000 cells/well in a 96-well plate (Lumox-Multiwell, Greiner Bio-One GmbH, Frickenhausen, Germany). Cells were allowed to recover for 24 h, followed by washing with PBS and incubating with 100 μM L-alanine-4-methyl-7-coumarinylamide-(MCA) trifluoroacetate (Fluka) in
PBS with 0.5% BSA and 20 mM HEPES (pH 7.2).

The release of fluorescent product 7-amido-4-methylcoumarin was monitored on a Kodak Gel Logic 200 Imaging System (Kodak, New Haven, Connecticut), with $\lambda_{\text{exc}}$ of 360 nm and $\lambda_{\text{em}}$ of 460 nm after 90 min. To assay for inhibition of enzymatic activity due to binding of CD13-specific antibody WM15 or the NGR-peptide, cells were pretreated in PBS containing a excess of antibody (5 μg/mL) or peptide (25 μg/mL) at 37°C for 1 h.

2.6 In Vitro CD13/APN Ligand Binding Studies

Cells were seeded in 24-well plates and incubated in culture medium overnight. For binding studies, cells were washed twice with PBS and resuspended in 150 μL cation-buffer diluted 1:20 in PBS/BSA (0.1%). Cy 5.5 (up to 30 μM) or NGR-Cy 5.5 (12.5 μM) was added into each well. After an incubation period of 1 h at 37°C, cells were washed twice with PBS and resuspended in binding buffer. For blocking studies, free NGR-peptide (125 μM) was added to each well for 45 min at 37°C. Cells could be directly visualized by fluorescence microscopy (40X objective magnification, Nikon TE 2000-S, Tokyo, Japan). The microscope was equipped with a mercury vapor lamp, 620/775 nm and 545/675 nm (excitation/emission) filters (AHF Analysentechnik, Tuebingen, Germany), a Nikon DXM1200F camera, and ACT1/ DXM1200F software (Nikon, Tokyo, Japan).

2.7 Immunohistochemistry

Tumors were removed and embedded in buffered formalin (4%, pH 7.0). Immunohistochemistry has been carried out commercially by Habedank-LID (Berlin, Germany) using the specific anti-CD13mAb H-300. For the in vivo visualization of NGR-Cy 5.5 uptake in tumor tissue, tumors were removed 24 h p.i., embedded in Tissue-Tek O.C.T. (Sakura, Zoeterwoude, Netherlands), snap-frozen, and serial-sectioned (10 μm). After drying, tissue fluorescence was evaluated by fluorescence microscopy.

2.8 Tumor Xenografts

All animal studies were approved by the institutional review boards (# G76/2005). Female athymic nude mice (nu/nu) were obtained from Charles River Laboratories (Sulzfeld, Germany). At 4 to 6 weeks of age, $3 \times 10^6$ HT-1080 or MCF-7 cells suspended in 100 μL PBS and Matrigel (BD Biosciences, San Diego, California) were injected subcutaneously in the flank and thoracic wall, respectively. When tumors reached 4 to 6 mm in diameter, the tumor-bearing mice were anesthetized by an i.p. injection of ketamine (125 mg/kg bw) and xylazine (12.5 mg/kg bw) and subjected to in vivo imaging studies.

2.9 In Vivo Optical Imaging Systems – Fluorescence Reflectance Imaging

The combination of x-ray and optical fluorescence reflectance imaging (FRI) was performed using the Image Station In Vivo FX Imaging System (Kodak Molecular Imaging Systems, New Haven, Connecticut), which is equipped to provide multi-wavelength fluorescence, luminescence, x-ray, and radioisotopic imaging. Fluorescence is provided by a 150-W halogen illuminator with selectable bandpass excitation and emission filters. For Cy 5.5 excitation, 625 ± 18 nm and emission 700 ± 17.5 nm were employed. X-ray energy is provided by a 35-Kvp, 150-μamp, microfocus source in an enclosed chamber above the animal. A specialized radiographic phosphor screen is moved into the imaging path under the animal, which transduces the x-ray energy to light. Light from the fluorescence and/or the radiographic screen is captured with the 4-million-pixel cooled CCD camera equipped with a 10X zoom lens. Images were captured 1, 3, 5, and 24 h after probe injection. Image acquisition times were 30 s per animal at the emission wavelength. Optical images were co-registered with x-ray background images, and regions of interest were selected and analyzed with the Kodak MI 4.0 software.

2.10 Fluorescence Mediated Tomography

All tomographic optical imaging studies were performed with a small-animal fluorescence mediated tomography (FMT) system from VisEn Medical, Inc. (Woburn, Massachusetts) that has been described in detail elsewhere.

The central piece of the system is the animal-imaging chamber configured in a parallel plate geometry. The chamber is scanned by a movable optical fiber that can excite a user-defined matrix of typically 50,000 to 100,000 source-detector projection pairs. The light source is software-selectable from two high-power laser diodes at 670 nm and 745 nm, operating in the range of 5 mW to 150 mW. Detection is performed with a thermoelectrically cooled CCD camera by means of an imaging lens and appropriate bandpass filters (three-cavity interference at 675 nm or 755 nm for excitation measurements, and 705 nm or 770 nm for emission measurements). Tomographic reconstruction is based on the normalized Born approximation, a diffraction optical tomographic technique that uses a diffusion-type theoretical photon propagation model in biological tissue. The underlying assumption of the mathematical model has been propagation in a homogeneous diffuse medium with average optical absorption and scattering properties those of small rodents at the relevant wavelengths. However, it has been demonstrated that because of the appropriate data normalization scheme applied, the algorithm is very accurate even in highly heterogeneous media and in vivo. The instrument is calibrated on each channel using euthanized mice with surgical implants of known fluorochrome concentration and is able to recover fluorescence depth, size, and concentration in three dimensions from tissue at any depth. The system exhibits strict linearity over a broad range of biologically relevant concentration and detection sensitivities in the order of femtomoles.

Animal scan times are in the range of 2 to 5 min; tomographic reconstruction times on a standard desktop PC are about 1 to 3 min.

For in vivo experiments, 2 nmol NGR-Cy 5.5 were injected into the tail vein of each mouse ($n=5$ for MCF-7, $n=6$ for HT-1080 xenografts). FRI and FMT were performed 1, 2, 5, and 24 h (FRI additional 5, 10, and 30 min) post injection. Blocking studies were accomplished by injecting 10 mg/kg NGR-peptide in 100 μL PBS (200 nmol) 10 min before injection of the labeled peptide ($n=6$ for HT-1080 xenografts). For biodistribution studies, FRI images of isolated organs (tumor, muscle, heart, spleen, kidney, liver, and lung) were carried out 24 h after injection of the tracer.
2.11 Statistical Analysis

Data are presented as mean ± SEM (standard error of the mean). Statistical analysis of in vivo tumor fluorescence was conducted using a two-tailed, unpaired student t-test. A p-value ≤ 0.05 was considered to be significant.

3 Results

3.1 Synthesis and Characterization of NGR-Cy 5.5

The NHS ester of the near infrared fluorophore Cy 5.5 is reacted with an equimolar amount of the cyclo-[Cys-Asn-Gly-Arg-Cys]-Gly-Lys (NGR) peptide and purified by reversed phase HPLC (purity ≥ 95%). The retention time was 28.5 min. Yields of the NGR-Cy 5.5 conjugate were typically 75 to 90%, as calculated with $\varepsilon_{675} = 250,000$ (mol/L)$^{-1}$ cm$^{-1}$ from the absorption spectra measured with the PBS solution. The excitation and emission spectrum of the targeted probe remained unchanged as compared to the nonmodified fluorophore showing excitation and emission peaks at 675 nm or 694 nm, respectively.

3.2 CD13 Protein Expression in HT-1080 and MCF-7 Cells

To determine the cellular origin of the CD13/APN receptor and its activity in HT-1080 and MCF-7 cells, cell lines were analyzed for CD13 protein expression by FACS, Western blot, and immunohistochemistry as well as by measurement of the specific aminopeptidase activity. For flow cytometry, cells were incubated with PE-labeled WM15 antibody directed against the human APN, and binding was analyzed (Fig. 1). Approximately 87% of the HT-1080 cells expressed the APN-receptor. In contrast, the receptor was hardly detected on MCF-7 adenocarcinoma cells [0.24%; see also Fig. 1(b)]. CD13 Western blotting of HT-1080 and MCF-7 cells and tumor tissues with the H-300 antibody, recommended for detection of CD13 of mouse, rat, and human origin, confirmed the apparent Mr of APN at 150,000. THP-1 cell lysate was used as a positive control. The APN enzyme was expressed only on HT-1080 cells and tumor tissue, while MCF-7 cells and tumors are negative for CD13 expression (not shown). Proper catalytic function of the CD13 protein was verified using a specific neutral aminopeptidase activity assay [Ref. 23; Fig. 1(c)]. CD13 activity in HT-1080 and MCF-7 cells is correlated with the expression measured by flow cytometry and Western blot analysis. The enzyme activity in HT-1080 cells could be specifically blocked by the WM15 antibody and the NGR-peptide.

3.3 In Vitro NGR Binding Assays

Based on the CD13 protein expression data, the binding prop-
properties of cyclo-[Cys-Asn-Gly-Arg-Cys]-Gly-Lys-Cy 5.5 (NGR-Cy 5.5) to the cell lines were evaluated by fluorescence microscopy (Fig. 2). Negligible signals were detected when cells were incubated with the nonmodified Cy 5.5 dye alone (up to 30 μM; data not shown), and also for MCF-7 cells following incubation with up to 12.5 μM of the labeled tracer [Fig. 2(c)]. The labeled peptide bound distinctly to HT-1080 cells [Fig. 2(a)]. The fluorescence was distributed over the cell surface and membrane-associated. Blocking of the signal was possible with a tenfold excess of unlabeled NGR-peptide [125 μM; Fig. 2(b)].

3.4 Immunohistochemistry

Immunohistochemistry on tumor tissue slides with CD13 antibody H-300 [Figs. 3(c) and 3(d)] exhibited membranous CD13/APN antigen expression on HT-1080 cells and only background staining in MCF-7 xenografts. Specific in vivo binding of NGR-Cy 5.5 in HT-1080 tumor tissue could be visualized in Fig. 3(e), exhibiting an intense, membranous fluorescence in tumor tissue 24 h after injection of the optical tracer, whereas MCF-7 tissue showed only background staining [Fig. 3(f)].

3.5 In Vivo Target Assessment and Optical Imaging

Figure 4(a) shows typical NIR fluorescence reflectance images of mice with HT-1080 (n=6) and MCF-7 (n=5) tumors, respectively, 24 h after injection with NGR-Cy 5.5. After clearing of the unbound circulating fluorochromes, the HT-1080 tumor xenografts could be clearly delineated from the surrounding background tissue up to 24 h with maximal fluorescence intensities of 2756.2 ± 209.4 AU, visible 3 h post injection [Fig. 4(b)]. Compared to the targeted probe, the blocking experiment revealed lower fluorescence intensities (e.g., 1 h post injection: 2339.3 ± 269.9 AU versus 1999.5 ± 213.5 AU; n=6; p<0.05). Target-to-background ratios (TBR) after tracer injection ranged around 1.61 ± 0.52 (1.54 ± 0.56 for the blocking experiment). MCF-7 xenografts showed a weaker fluorescence, indicating a lower expression of the CD13/APN receptor [1407.5 ± 122.0 AU 3 h p.i.; Fig. 4(b)]. In HT-1080 tumor xenografts, the nonmodified Cy 5.5 revealed a fluorescence intensity of 1404.4 ± 243.8 AU (n=7) 24 h post injection (data not shown).

FMT confirmed the FRI results, exhibiting a strong fluorochrome distribution in the target tissue throughout the tumor (Fig. 5). Quantitative data analysis revealed a fluorochrome concentration 5 h after injection of 306.7 ± 54.3 nM NGR-Cy 5.5 (Fig. 5), while blocking with the free peptide
resulted in an average fluorochrome concentration of 195.3 ± 21.9 nM NGR-Cy 5.5 (Fig. 5). TBR values for this time-point ranged around 7.53 ± 4.04 and 3.33 ± 2.13 for the blocking experiment. Tumor fluorescence reached maximum intensities after 5 h and showed a slow clearance over the time [Fig. 5(g)]. Fluorochrome concentrations in nontarget tissue were significantly lower and ranged around 116.0 ± 18.3 nM.

3.6 **Biodistribution**

The distribution of Cy 5.5-labeled NGR-peptide was measured in tumor, muscle, heart, lung, spleen, kidneys, and liver from nude mice bearing HT-1080 fibrosarcoma (n=6) and MCF-7 adenocarcinoma (n=5) tumors. Data were obtained by FRI measuring 24 h post injection of the tracer. A representative example of NGR-Cy 5.5 biodistribution in HT-1080 tumor–bearing mice is shown in Fig. 6. The highest concentration of optical tracer 24 h after injection was determined in kidneys (792.6 ± 272.6 AU for HT-1080 and 725.3 ± 52.6 AU for MCF-7), followed by the accumulation in liver (398.0 ± 86.9 AU), tumor (331.9 ± 151.7 AU), and lung (306.5 ± 38.6 AU). Preinjection of the unlabeled peptide resulted in decreased uptake in all dissected tissues (data not shown).

4 **Discussion**

Targets that are located on the endothelial surface are ideal candidates for in vivo molecular imaging approaches because pharmacological barriers are virtually nonexistent. CD13/APN is an ectoenzyme located in the outer cell membranes of various cells like epithelial cells, macrophages, and endothelial cells. Therapeutic targeting of the CD13/APN receptor has successfully been performed using, e.g., NGR-fusion peptides with related anticancer prodrugs exhibiting selective cytotoxicity toward CD13/APN positive tumor cells or angiogenic endothelial cells.14,15,18,24–26 More recently, fluorochrome labeling of NGR-peptides has been shown to be feasible.17

For our study, we chose a cyclic NGR-peptide with a cysteine group and a gly-lys carboxy terminus as a spacer function. The use of a cysteine moiety for cyclization was chosen because of the reported enhanced tumor targeting efficiency of such peptides.15 CD13/APN activity of positive HT-1080 cells16 and negative MCF-7 cells were demonstrated by FACS, Western blotting, and an APN activity assay using Ala-MCA as substrate. Blocking of CD13-specific aminopeptidase activity with the NGR-peptide and the peptide-dye conjugate was in line with the activity-blocking antibody WM15,23,24 indicating the specific binding of the labeled NGR-peptide to the target structure.

In in vitro binding studies evaluated by fluorescence microscopy, the peptide-dye conjugate showed distinct binding affinity to HT-1080 cells, while MCF-7 cells and incubation of cells with Cy 5.5 revealed little to no fluorescence. Initially, the fluorescence was membrane associated – a phenomenon also observable in fluorescence microscopic analysis of the tumor xenografts; prolonged incubation times resulted in endocytosis of the tracer and nuclear staining. The specificity of the probe could be confirmed by a competition assay with a tenfold excess of unlabeled NGR-peptide, which resulted in a reduction of cellular fluorescence. Thus, in vitro tracer binding results closely correlate with FACS and APN activity assays and suggest a high binding affinity of the peptide-dye conjugate to the target.

In vivo, HT-1080 xenografts were clearly visualized by FRI up to 24 h after i.v. injection of NGR-Cy 5.5, showing a maximum TBR of 1.61 ± 0.52 after 1 h. Blocking with free peptide resulted in a decreased uptake of the optical tracer (TBR 1.54 ± 0.56). MCF-7 tumors showed a TBR of 0.96 ± 0.23. To overcome FRI-related quantification problems, FMT was performed in a group of animals, allowing the three-dimensional (3-D) reconstruction and quantification of fluorochrome distribution in vivo. At maximum tumor tracer uptake after 5 h, FMT data correlates nicely with in vitro and FRI data, reflecting the different APN cellular expression level for HT-1080 and MCF-7 tumors. Blocking with the unlabeled peptide led to a decreased uptake of optical tracer (e.g., for HT-1080 after 24 h 112.8 ± 25.7 nM). Besides target-specific fluorochrome accumulation, nonspecific accumulation of the tracer occurs due to perfusion and vascular permeability effects as well as endocytosis of the peptides
over time. This effect was also observed here by imaging the nonmodified fluorophore and has been described previously for other non-target-specific NIR-fluorochromes. These non-specific, contrasting mechanisms can also explain the relatively high remaining fluorescence signal after blocking with the free peptide. Necrotic areas within the tumor also lead to a heterogeneous tracer uptake and will influence the correlation between NGR-Cy 5.5 and APN expression. TBR measured by FMT for HT-1080 xenografts (e.g., 5 h post injection 7.53 ± 4.04) was significantly higher as compared to corresponding FRI data (1.61 ± 0.52). This may be on one hand attributed to a lower amount of bleed-through between the excitation and emission spectrum in the FMT. On the other hand, this demonstrates the capability of resolving and differentiating deeply located tissue fluorescence (e.g., deriving from the liver) from more superficially located tumor-associated fluorochrome accumulation. While FRI cannot differentiate between deeply located, strongly fluorescent organs (e.g., liver) and superficial fluorochrome tumoral accumulation, cross-sectional imaging with FMT can. Thus, scattered fluorescence from deep-seated structures can significantly spoil the TBR in FRI compared to FMT.

Direct imaging of excised organs 24 h p.i. exhibited a high accumulation of the optical tracer not in the tumor but in the kidneys and liver of all tested mice. That has a lot to commend a rapid blood clearance and a renal excretion of the NGR-Cy 5.5 peptide, a feature that is preferable for further clinical applications. Nevertheless, CD13 expression by the reabsorptive epithelium of the kidney was also reported, but NGR drug conjugates like NGR-TNF failed to bind to CD13 expressed in normal kidney, indicating a selective tumor-homing property to a specific CD13 isofrom. Further studies are necessary to clarify the binding properties of NGR-conjugated optical imaging probes to different APN isoforms. However, the immunochemical data suggest that the NGR-Cy 5.5 probe binds to both the membrane-associated protein of the tumor cells as well as endothelial targets.

For in vivo applications, specifically for tomographic imaging in deep tissue sections, a maximum TBR is desirable.

Fig. 5 Fluorescence-mediated tomography of HT-1080 and MCF-7 tumor-bearing nude mice. FMT experiments were performed to visualize and quantify the three-dimensional (3-D) fluorochrome distribution in deeper tissue sections. The fluorochrome concentration is displayed color encoded (scale: 1 to 256 nM). FMT was carried out 24 h after injection of 2 nmol NGR-Cy 5.5 probe in HT-1080 (a) and MCF-7 (e) tumor-bearing nude mice. In order to assess probe specificity, 200 nmol of the unlabeled peptide were i.v. injected 10 min before application of the NGR-Cy 5.5 probe (c). FMT revealed intense fluorochrome retention 60 min after probe injection in HT-1080 tumors (b), which could be significantly reduced after pretreatment with the nonmodified peptide (d). Compared to the fluorescence measured in HT-1080 tumors, the fluorochrome accumulation in MCF-7 tumors (f) was substantially lower. Tumor fluorescence washout over the time was slower in HT-1080 than in MCF-7 tumor tissue (g), black bar = native HT-1080; gray bar = blocked HT-1080; white bar = MCF-7.

Fig. 6 Representative overlay of white light and color-encoded FRI images of dissected organs of HT-1080 tumor-bearing nude mice. Animals (n=6) were sacrificed 24 h after intravenous injection of 2 nmol NGR-Cy 5.5. Enhanced fluorochrome accumulation was detected in kidneys, liver, and lung tissue [(1): HT-1080 tumor, (2) heart, (3) spleen, (4) lung, (5) liver, (6) kidneys], indicating a high tracer washout of tumor tissue and a renal excretion. The fluorochrome concentration is displayed color encoded (scale: 1 to 256 nM).
The affinity of the currently available NGR-NIR-dye conjugate is moderate. Different strategies have been proposed for other peptidic tracers, especially the RGD-peptide directed against the integrin \(\alpha_\beta_2\)-receptor, to increase the target affinity of the imaging agent and decrease a high accumulation in other organs such as liver and kidneys. Possible methods are glycosylation of the tracer, peptide carbohydrate conjugates, and peptidomimetics based on the NGR-structure as well as heterodimeric, homodimeric, and homotetrameric ligand systems as tested for the RGD-peptide.\(^{30-32}\) These modifications have already been carried out with radio-labeled\(^{30,33}\) and Cy 5.5-labeled RGD probes.\(^{34}\) The results could be a useful source for the development of NGR optical imaging probes.

A clinical translation of this optical imaging technology is conceivable for certain organs such as breast or head and neck tumors. The feasibility of contrast-enhanced diffuse optical mammography in humans has been shown by Ntziachristos et al.\(^{25}\) Combining this imaging technique with target-specific fluorochromes as described in this work should greatly enhance the diagnostic accuracy for detecting breast cancer at an early tumor stage.\(^{36}\) Besides tomographic imaging, planar imaging techniques such as FRI can easily be translated for clinical use. A tumor binding fluorochrome could for instance be applied to preoperatively “tag” the tissue to be resected. The NIR signal deriving from the tumor could then easily be detected with planar optical imaging techniques in the operating room, and thus tumor margins could be visualized for the surgeon.\(^{37,38}\)

In conclusion, our study shows that CD13/APN expression can be imaged in vivo using fluorescence reflectance imaging and fluorescence-mediated tomography approaches. Since FMT allowed 3-D reconstruction of fluorochrome distribution, the tumor-to-background ratio could be significantly enhanced and furthermore quantified noninvasively. This imaging technique is therefore a versatile tool not only for experimental small animal molecular imaging but potentially also for noninvasive target detection, e.g., in clinical breast cancer imaging.\(^{39}\)

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