

# **Intraoperative fluorescence imaging : clinical translation of targeted and non-targeted tracers**

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# **chapter 10**

Real-time near-infrared fluorescence imaging using CRGD-ZW800-1 for intraoperative visualization of multiple cancer types

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#### **ABSTRACT**

Incomplete resections and damage to critical structures increase morbidity and mortality of patients with cancer. Targeted intraoperative fluorescence imaging aids surgeons by providing real-time visualization of tumors and vital structures. This study evaluated the tumor-targeted zwitterionic near-infrared fluorescent peptide CRGD-ZW800-1 as tracer for intraoperative imaging of multiple cancer types. crgd-zw800-1 was validated *in vitro* on glioblastoma (u-87 mg) and colorectal (ht-29) cell lines. Subsequently, the tracer was tested in orthotopic mouse models with HT-29, breast (MCF-7), pancreatic (BXPC-3), and oral (OSC-19) tumors. Dose-ranging studies, including doses of 0.25, 1.0, 10, and 30 nmol, in xenograft tumor models suggest an optimal dose of 10 nmol, corresponding to a human equivalent dose of 63 µg/kg, and an optimal imaging window between 2 and 24 h post-injection. The mean half-life of crgd-zw800-1 in blood was 25 min. Biodistribution at  $4$  h showed the highest fluorescence signals in tumors and kidneys. *In vitro* and *in vivo* competition experiments showed significantly lower fluorescence signals when u-87 mg cells (minus 36%, p=0.02) or ht-29 tumor bearing mice (TBR at 4 h 3.2±0.5 vs 1.8±0.4, p=0.03) were simultaneously treated with unlabeled crgd. crgd-zw800-1 visualized *in vivo* all colorectal, breast, pancreatic, and oral tumor xenografts in mice. Screening for off-target interactions, crgd-zw800-1 showed only inhibition of cox-2, likely due to binding of CRGD-ZW800-1 to integrin  $a_V\beta_3$ . Due to its recognition of various integrins, which are expressed on malignant and neoangiogenic cells, it is expected that crgd-zw800-1 will provide a sensitive and generic tool to visualize cancer during surgery.

#### introduction

Accurate and real-time detection of tumors and vital structures during surgery remains challenging.Similarto radioisotopes for single-photon emission computed tomography (SPECT) and positron emission tomography (PET), tumor targeting ligands can also be conjugated to near-infrared (nir, 700-900 nm) fluorophores. These fluorescent tracers allow real-time visualization of tumors during surgery using dedicated intraoperative imaging systems [1, 2]. The advantage of nir fluorescence is that it can penetrate through 8 mm into tissue, allowing identification of targets even when they are not yet fully exposed [3]. Several nir fluorophore-labeled ligands have been introduced in clinical trials, including the labeled antibodies cetuximab-IRDYE800CW [4] for the epidermal growth factor receptor (egfr), bevacizumab-irdye800cw (ClinicalTrials.gov Identifier nct01508572) for vascular endothelial growth factor (VEGF), and SGM-101 (Netherlands Trial Register number NTR5673) for carcinoembryonic antigen (CEA). Antibodies have high affinity to specific targets, but they can cause immunogenic reactions and have a relatively slow biodistribution and low clearance rates [5, 6]. On the other hand, peptides possess more favorable pharmacokinetic properties, but often have lower affinity compared to antibodies. Clinical trials using the small NIR fluorescent molecules otl-38 (1.4 kDa) targeting the folate receptor α (frα) and ge-137 (3.7 kDa) targeting human tyrosine kinase c-met [7, 8] have already been reported. However, these targets are selectively overexpressed and may only be applicable for a limited number of tumor types.

In this study we evaluate crgd-zw800-1, a cyclic pentapeptide conjugated to the 800 nm NIR fluorophore zw800-1. RGD is clinically well known and binds to various integrins ( $\alpha_V\beta_1$ ,  $\alpha_V\beta_3$ ,  $\alpha_V\beta_5$ ,  $\alpha_V\beta_6$ ,  $\alpha_V\beta_8$ ,  $\alpha_5\beta_1$ ,  $\alpha_8\beta_1$  and  $\alpha_{IIB}\beta_3$ ). Expression of particular integrins can be found on tumor cells and tumor-associated vascular endothelium and correlates with neoangiogenesis [9]. Overexpression is found in almost all solid tumors, including breast, colorectal, pancreas, brain, lung, and other cancers [10]. CRGD-ZW800-1 may therefore act as a generic tracer for a broad variety of solid tumors. Indeed, various phase I and ii clinical trials with rgd-based pet tracers have shown uptake in multiple tumor types, including melanomas, glioblastomas, breast, colorectal, ovarian, cervical, non-small cell lung, neuroendocrine, head and neck, and pancreatic cancer [11-14]. Preliminary work on the development of crgd-zw800-1 showed low non-specific uptake and excellent *in vivo* properties in tumor xenograft mouse models [15, 16]. Moreover, due to its predominant renal clearance, ureters could also be recognized, which can prevent damage to these structures during surgery in the lower pelvis.

This study aims to characterize the *in vitro* properties, optimize dosage and timing, study pharmacokinetics, and to evaluate the feasibility of using CRGDzw800-1 as a generic tracerforintraoperative near-infraredfluorescence imaging of solid tumors.

### results

## **The expression of αVβ3, αVβ5 and αVβ6 on tumor cell lines**

Table 1 reports the measured antigenic sites per cell for three RGD binding integrins on clinically relevant tumor cell lines. Pancreatic bxpc-3 cells expressed all 3 integrins, while colorectal HT-29, glioblastoma u-87 MG, and oral OSC-19-GFP cells expressed 2 of the integrins evaluated.Breast mcf-7 cells showed only expression of integrin α<sub>ν</sub>β<sub>5</sub>. Overall, u-87 MG cells showed the highest and MCF-7 cells the lowest number of the three integrin antigenic sites combined.

## **Binding assay and** *in vitro* **and** *in vivo* **competition experiments**

Binding assays with crgp-zw800-1 on intermediate integrin-expressing HT-29 cells showed an almost linear increase in fluorescence intensity with increasing concentrations (Figure 1a). Significantly lower signals were observed when cells were incubated at 4°C, which prevents internalization of the agent. *In vitro* competition for binding crgd-zw800-1 (500 nM) with a 1:1 molar ratio of unlabeled crgd (500 nM) resulted in a reduction of 32% on the ht-29 cells and 36% on the high integrin-expressing u-87 Mg cells compared to cells treated without unlabeled CRGD (Figure 1B).

*In vivo* competition using a 200 times higher dose of unlabeled CRGD compared to CRGD-ZW800-1 (2.0 µMol vs. 10 nmol) in the orthotopic HT-29 model (Figure 1c) resulted in a significant decrease in tumor-to-background ratio (tbr) at 4 and 24 h (42%, p=0.02 and 54%, p=0.007,respectively).

## **Identification of off-target interactions**

Out of  $44$  selected targets, crgp-zw800-1 resulted in significant inhibition (62%) of control-specific binding of cyclooxygenase-2 (cox-2). Weak to moderate inhibition was seen in 5-hydroxytryptamine (serotonin) receptor 2B (5-HT2B, 29%). lymphocyte-specific protein tyrosine kinase (lck, 28%), and dopamine receptor D<sub>25</sub> (22%) (Table s1, supplementary data).

## **In vivo validation using ht-29 (αVβ3 negative) colorectal tumors**

Four doses were explored using both the Pearl® Impulse small animal imaging system (li-cor, Biosciences, Lincoln, ne, u.s.a.) and the prototype Fluorescence-Assisted Resection and Exploration (flare®, Curadel, llc, ma, u.s.a.) system as indicated (Figure 2A). TBRS at 4 and 24 h post injection were sufficient (i.e.  $>$ 2) in the 1, 10 and 30 nmol dose groups (Figure 2B and 2C). The FLARE showed clear fluorescence signals in the 10 and 30 nmol dose groups. The optimal dose was therefore set at 10 nmol.

## In vivo validation using **BXPC-3** (α<sub>V</sub>β<sub>3</sub> positive) pancreatic tumors

Tumor specific signals (TBRS >2) were observed in all three dose groups using FLARE® and Pearl® at 4 h post injection (Figure  $\overline{5}$ A and  $\overline{3}$ B). Fluorescence microscopy of the pancreatic tumors showed that nir fluorescence signals originate from inside malignant cells.

## **In vivo validation using osc-19 (α<sub>V</sub>β<sub>3</sub> negative) oral tumors**

The optimal dose of 10 nmol CRGD-ZW800-1 was administered in the well-established tongue orthotopic cancer model and resulted in clear tumor demarcation with a mean TBR of 5.4  $\pm$  1.1 using Pearl® (Figure 3c and 3D). Fluorescence microscopy of the tumor showed a clear demarcation between malignant and normal tissue.

## **In vivo validation using MCF-7 (α<sub>V</sub>β<sub>3</sub> negative) breast tumors**

The optimal dose of 10 nmol CRGD-ZW800-1 was administered to mice bearing orthotopic breast tumors. Compared to 0.5 h post injection fluorescence intensity of tumors decreased by 61%  $\pm$  8% and by 77%  $\pm$  4% at 4 and 24 h post injection (Figure  $\Delta C$ ) Using the Pearl® and the skin as background tissue TBRs reached values slightly higher than 2 between 1 and  $\Delta 8$  h post injection (Figure  $\Delta B$ ). These relatively low that compared to the other models were due to the relatively high fluorescence of the skin. Removal of the skin in a single mouse resulted in significant higher TBRs:  $3.0 \pm 0.3$  vs. 1.5  $\pm$  0.1 at 4 h post injection (p=0.02, Figure S1, supplementary data).

## **In vivo biodistribution**

In both ht-29 and bxpc-3 tumor models an increased dose of crgd-zw800-1 resulted in stronger fluorescence signals in the tumors (Figure 5a). The calculated organ-to-tumor ratios in these mice showed that for all doses the bxpc-3 tumor was at least twice as fluorescent as the skin, intestines, lungs, muscle, gallbladder, brain, and blood at  $\Delta$  h post injection (Figure 5B). The large intestines and liver showed fluorescence ratios of around half compared with bxpc-3 tumors. Fluorescence signals in kidneys were equal to or higher than tumors, reflecting renal clearance of CRGD-ZW800-1.

As described before, at 24 h the intensity of fluorescence signals in HT-29 tumors decreased compared to  $\Delta$  h (Figure 5c). However, in all dose groups the ht-29 tumor was still more than twice as fluorescent compared to signals observed in lungs, pancreas, muscle, gallbladder, brain, and blood (Figure 5d). Compared to the ht-29 tumor, the fluorescence signal was higher in kidneys (all dose groups), and liver and skin (only 10 and 30 nmol dose groups). Significant differences between dose groups were seen in kidneys only.

## **Pharmacokinetics**

Peripheral blood concentrations were measured after injection of 10 nmol crgdzw800-1 per mouse (n=5). These mice did not have tumors. The measured fluorescence values were validated by a calibration curve (Figure s2, supplementary data). The maximal concentration of CRGD-ZW800-1 in serum measured at 1 minute post injection was 1.17 µM (Figure 6). Decrease of CRGD-ZW800-1 serum concentrations followed a biphasic pattern with a distribution and elimination phase with a mean terminal half-life of 71.1  $\pm$  9.4 min. Mean systemic clearance was  $0.25 \pm 0.06$  ml/min.

#### discussion

nir fluorescent tracers offer significant potential for accurate and real-time tumor visualization during resection. This study shows that CRGD-ZW800-1 is specific for integrins expressed on multiple tumor types providing an imaging window between 2 to 24 h post injection. Due to its clearance via kidneys rather than the liver, crgd-zw800-1 could be used for nir fluorescence imaging of almost all solid cancers and their metastases.

Research on RGD peptides for cancer targeting has mainly focused on the  $q_v\beta_3$ integrin and neoangiogenic endothelial cells, often using the glioblastoma u-87 mg cell line as proof of concept in animal models. Indeed, our inventory shows that u-87 mg cells contain the highest level of  $a_0$  $\beta_3$  integrin, confirming its frequent use as positive control for rgd binding. However, 7 other integrins are also able to bind the RGD sequence, including  $q_y\beta_5$  and  $q_y\beta_6$  which are overexpressed on malignant cells from various tumor types [17-19]. This explains why crgdzw800-1 accumulates in HT-29, MCF-7, and OSC-19 tumors: All are  $q_v\beta_3$  negative, but express integrins  $a_V\beta_5$  and/or  $a_V\beta_6$ . The specificity of the fluorescence signals was previously demonstrated [16] and could also be extracted from *in vitro* and *in vivo* competition experiments, ruling out possible non-specific accumulation, e.g. due to the enhanced permeability and retention (EPR) effect [20]. crGD thus offers potential for visualization of almost any type of solid cancer.

crgd has previously been conjugated to the fluorophores indocyanine green (icg), irdye800cw, and cy5.5 for optical imaging applications [21-25]. A study using CRGD-IRDYE800CW showed delineation of U-87 MG tumors in mice and underscored the relative simplicity and low cost of the synthesis [21]. Another study showed *in vitro* the binding capacity of crgd-icg using mcf-7 cells as a negative control [24]. In our study, we showed that crgd-zw800-1 targets *in vivo* even the low integrin expressing cell line MCF-7. In addition to expression of integrin  $a_V\beta_5$  by MCF-7 cancer cells, this could also be explained by expression of RGD-binding integrins by tumor-associated vascular endothelium and stromal cells. NIR fluorescence imaging of tumors using CRGD conjugated to

irdye800cw, cy5.5, or icg is feasible, but compared to crgd-zw800-1 these tracers show higher non-specific uptake in normal tissues and organs due to their anionic charge [16]. For example, crGD-ICG showed significant uptake in liver and colon, which can hamper identification of gastrointestinal tumors [25]. CRGDzw800-1 has a balanced charge distribution and a net charge of 0, which likely shields its hydrophobicity and minimizes interactions with serum proteins [26]. In our biodistribution study ratios between tumor and relevant organs, such as intestines, lung, pancreas, muscle, or brain, were more than sufficient for accurate discrimination. Moreover, increasing the dose to 30 nmol did not result in more non-specific uptake, whereas fluorescence in tumors was enhanced. Our group previously showed that crgd-zw800-1 is renally cleared, allowing identification of ureters [15]. This study confirms that kidneys are the exclusive clearance route. First, the measured clearance of CRGD-ZW800-1 is identical to basal glomerular filtration rate in mice (i.e. 0.25 ml/min) [27]. Second, there was a significant difference in fluorescence signals between doses in the kidneys.Third, all mice showed high signals in the bladder up to 4 h post injection, as shown for example in MCF-7 model. In contrast, no significant differences were seen between the dose groups in the gallbladder or intestines, making hepatic clearance unlikely. Clearance patterns are important, because limited target expression often constrains maximal tumor specific uptake [26]. For imaging colorectal or pancreatic tumors, renal clearance is beneficial, as it reduces background signals from liver and intestines and thereby improves TBRS. On the other hand, imaging tumors of the genitourinary tract might be impeded due to the renal excretion of CRGD-ZW800-1.

crgd-zw800-1 is internalized into cells after binding as shown in previous studies and recognizable in the *in vitro* binding assays at 4 and 37°C. The continuous process of internalization and recruitment of fresh integrins to the cell membrane results in a relatively large imaging window between 2 to 24 h post injection. After internalization, the majority of integrins are recycled back to the cellular membrane in approximately 45 min [28]. Based on the mean blood halflife of 25 min, we can rationalize that the majority of crGD-ZW800-1 is internalized during the first hours. The MCF-7 model confirms this; TBRS increase up to 2 h post injection. The absolute signal in tumors depends therefore on the amount of initially available agent and integrins rather than accumulation of the agent over time. Although the central oxygen linking modality of zw800-1 might be vulnerable for degradation *in vivo*, the intracellular signals remained visible up to 48 h post dosing [29]. For optimal results in a clinical study CRGD-ZW800-1 should be administered several hours prior to imaging.This is a major advantage compared to labeled antibodies, which have to be administered several days prior to surgery, requiring an additional visit or earlier admission to the hospital.

Besides interacting with extracellular matrix proteins, integrin  $\alpha_V \beta_3$  can also modulate intracellular pathways essential for cox-2 expression [30]. The inhibition of cox-2 as shown in the *in vitro* off-target screening may be the result of binding crασ-zw800-1 to integrin α<sub>ν</sub>β<sub>3</sub>. Potentially, inhibition of cox-2 in blood vessels can lead to higher blood pressure and increase the risk of cardiovascular events because of decreased prostacyclin production [31].This seems unlikely to occur after a single low dose in humans, also because cox-2 inhibition by crgdzw800-1 was observed in the micromolar range. Following these preclinical findings a toxicology study in rats (unpublished data) is already performed and neither clinical symptoms, nor treatment-related histopathological, biochemical, and hematological abnormalities were seen after intravenous injection of 5.0 mg/ kg crgp-zw800-1. We can therefore conclude that the no observable adverse event level (NOAEL) in rats is at least 5.0 mg/kg. This confirms the absence of significant off-target inhibition by CRGD-ZW800-1, even at relatively high doses. A single, low dose of CRGD-ZW800-1 is therefore not expected to result in relevant clinical effects. The NOAEL of 5.0 mg/kg in rats is adjusted for body surface area a human equivalent dose (HED) of 0.80 mg/kg [32]. The us Food and Drug Administration (FDA) recommends to use 10% of the HED of the NOAEL to determine a safe starting dose [33]. A dose of 80 µg/kg can therefore be considered as a safe starting dose for first in-human studies. Furthermore, the dye zw800-1 was separately evaluated (unpublished data) and showed no toxicity or adverse events in animal toxicity studies using doses of 24.5 mg/kg (HED 3.9 mg/kg). The rgd integrin-binding site is evolutionary conserved in vertebrates, including humans, mice, and rats [34]. Due to these similarities, animal models provide realistic expectations for first-in-patient studies. Adjusting the optimal dose in mice (10 nmol) into the hed shows the same dose range (63 µg/kg). In a clinical study with a RGD-PET tracer the tumor uptake was  $5.1 \pm 3.6\%$  injected dose per liter [35]. Assuming that the tumor uptake of CRGD-ZW800-1 will be comparable, the concentration inside tumors after administering 63 µg/kg cRGD-ZW800-1 will be approximately 0.2 µM. The FLARE® prototype system and other clinical systems can easily detect such concentrations (See also Figure s2, Supplementary data) [36]. Furthermore, this dose range is comparable to the optimal clinical dose of the similar sized fluorescent peptide OTL-38 for intraoperative fluorescence imaging of frα-positive ovarian cancer: 12.5 to 50 µg/kg [8]. We conclude that targeting RGD-binding integrins appears to be safe. This is also suggested by a previous clinical study in which the crgd-based integrin inhibitor cilengitide was intravenously administrated at a dose of 2.0 g twice a week for a period up to 2 years without serious side effects [37].

#### conclusion

In conclusion, due to its recognition of multiple integrins and its high affinity binding to a wide range of malignant and neoangiogenic cells, CRGD-ZW800-1 has the potential to become a sensitive and generic tool to visualize cancer during surgery.

#### materials and methods

### **crgd-zw800-1 synthesis**

The synthesis of crgp-zw800-1 was previously described [16].

## **Human cancer cell-lines**

Human colorectal (HT-29-LUC2), pancreatic (BXPC-3-LUC2), squamous tonque (OSC-19-luc2-cgfp),breast carcinoma (mcf-7-luc2-cgfp), and glioblastoma (u-87 mg) cell line have been described previously [38, 39] (see also Supplementary Data).

## **Flow cytometry**

Qifikit (dako) was used to determine the number of antigenic sites for the three most commonly described integrins  $a_V\beta_3$ ,  $a_V\beta_5$ , and  $a_V\beta_6$  in the four cell lines with flow cytometry. Cells were grown to 90% confluence, detached with trypin/EDTA (PAA), and counted using trypan blue. Cells were resuspended in DMEM/0.1%BSA /0.1%NaN3 (wash- and dilution buffer), adjusted to 0.25\*106 cells per tube, and incubated with anti-ανβ3 (Millipore, Clone LM609), ανβ<sub>5</sub> (Millipore, Clone P1F6), αvβ6 (Millipore Clone10D5), and isotype control antibody mopc21 (BioXcell) at saturated conditions, for 30 min on ice. Cells and set-up and calibration beads were washed twice with wash buffer, followed by incubation with fitc-conjugated secondary antibody for 45 min on ice. After washing twice, propidium iodide was added to the cells. Cells and beads were measured on a BD LSRII flow cytometer (bd Biosciences). With the mean fluorescence intensity values of the calibration beads, a calibration curve was constructed. Using this curve the antigen density per cell for the different integrins was calculated.

## **In vitro binding and competition experiments**

To evaluate the binding capacity of crgd-zw800-1,u-87 mg cells were plated in a 96-well plate at a density of ~40,000 cells per well. At 90-100% confluence, cells were washed and incubated with various concentrations of crosp-zw800-1 at 4°C or 37°C for 1 h (binding assay). crgp-zw800-1 was added to the u-87 MG cells in various concentrations: 0,125, 250, 500, and 1000 nM.

For *in vitro* competition, u-87 MG (α<sub>V</sub>β<sub>3</sub> and α<sub>V</sub>β<sub>5</sub> positive, α<sub>V</sub>β<sub>6</sub> negative) and HT-29 (α<sub>V</sub>β<sub>3</sub> negative, α<sub>V</sub>β<sub>5</sub> and α<sub>V</sub>β<sub>6</sub> positive) cells were plated in a 96-well plate at a density of  $\sim$ 40,000 and  $\sim$ 60,000 cells per well, respectively. Subsequently, 500 nM of crgd-zw800-1 was simultaneously added to the cells and incubated at 37°C for 2 h. At the same time, half of the cells were also incubated with 500 nM unlabeled ("cold") crgp.

Both the cells in the binding and the competition experiment were then washed twice and imaged with the Odyssey NIR scanner (LI-COR Biosciences, Lincoln, Nebraska: focus offset 3 mm; 800-nm channel). Next, cells were permeabilized with a 40/60 mixture of acetone and methanol followed by a washing step and a 5 min incubation with ToPro3 (1/2000, Invitrogen), a far red fluorescent dye (642/661nm). The wells were then washed and again imaged with the Odyssey scanner (focus offset 3 mm; 700-nm channel) to quantify the number of cells in each well. The experiments were performed in triplicate.

## **Identification of off-target interactions**

Off-target interactions are binding of tracers to other components on the cell surface than the actual target site. These non-specific interactions are often of low affinity and easily missed, but can have significant effects. We outsourced the screening for *in vitro* for off-target interactions of CRGD-ZW800-1 to Eurofins Cerep sa, Le bois l'Evêque, France. They assessed the percentage of inhibition of a reference compound in 44 selected targets, including G protein-coupled receptors (gpcrs), transporters, ion channels, nuclear receptors, kinases and other non-kinase enzymes. These targets were recommended by four major pharmaceutical companies [40]. The complete list is included in Table s1 of the supplementary data.Each assaywasperformedtwice using10µMcrgd-zw800-1 anda reference compound. Inhibition higher than 50% was considered to represent a significant effect of crgp-zw800-1. Results showing between 20% and 50% were considered weak to moderate effects, while lower than 20% was not considered significant and mostly attributable to variability of the signal around the control level.

## **Animal models**

Six week-old athymic female mice ( $CD1$ -Foxn $1^{NU}$ , Charles River Laboratories, Wilmington, ma, u.s.a.) were used and housed in ventilated cages. Normal pellet food and sterilized water were provided *ad libitum*.Throughout tumorinoculation and imaging procedures, animals were anesthetized with isoflurane. Each group consisted of 3 or more mice. The Animal Welfare Committee of Leiden University Medical Center approved all animal experiments for animal health, ethics, and research. All animals received humane care and maintenance in compliance with the "Code of Practice Use of Laboratory Animals in Cancer Research" (Inspectie WandV, July 1999).

To induce subcutaneous tumors, colorectal cancer cells (ht-29) were injected at 4 sites on the back (500,000cells per spot). Subsequently,these colorectal tumor cells were transplanted on the colon of other mice, as described by Tseng *et al*. [41]. Approximately 500,000 pancreatic tumor cells (bxpc-3) were injected into the pancreas, as previously described by Kim *et al*. [42]. Approximately 40,000

oral tumor cells (osc-19) were injected in the tongue, as previously described by van Driel et al. [39]. Approximately 500,000 breast tumor cells (mcf-7) were injected in both sides of the mammary fat pad to induce breast cancer tumors.

All animals were imaged using a Pearl® imager (LI-COR, Lincoln, NE) and an original flare® prototype. Animals with ht-29 tumors were imaged at 4 and 24 h post injection; animals with bxpc-3 and osc-19 tumors at 4 h only. Animals with MCF-7 tumors were imaged at 0.5, 1, 2, 4, 6, 8, 24 and 48 h post injection. The specific and control images were normalized and regions of interest (rois) were selected using associated software. TBRS were calculated by dividing the tumor signal by the background signal. Based on our experience a TBR ≥ 2 is sufficient to discriminate between tumor and surrounding tissue [38].

After imaging, mice were sacrificed and OSC-19 and BXPC-3 tumors were quickly frozen in isopentane at -80°C for histological evaluation.Tissues were sectioned at 10 µM and fluorescence imaging was performed using fluorescence microscopy (Nikon eclipse e800, Nikon, Amsterdam, The Netherlands). All histologic sections were stained with standard hematoxylin-eosin stain (he). Organs of mice with colorectal and pancreatic cancer were collected and imaged using the Pearl® to evaluate biodistribution.

## *In vivo* **specificity**

The binding specificity of crgp-zw800-1 was explored in a competition experiment using an excess of unlabeled crgp (200-fold excess) co-injected with a standard dose of 10 nmol  $CRGD-ZW8OO-1$  (n=4). Simultaneously, mice (n=3) were injected with 10 nmol crgd-zw800-1 as positive control. All mice had orthotopic ht-29 colon tumors. Images were acquired at 4 and 24 h post-injection after which animals were sacrificed and organs were resected and scanned with Pearl®.

## *In vivo* **binding characteristics and biodistribution**

Mice were administered with 0.25 (HT-29 model only), 1.0, 10, or 30 nmol CRGD $zw800-1$  to evaluate tumor signals, biodistribution, and to calculate TBRS (n=3 per dose group). At  $\Delta$  h (BXPC-3 model) and 2 $\Delta$  h (HT-29 model) after injection, mice were fully dissected and fluorescence intensity of their organs was measured.

Mice bearing orthotopic osc-19 tumors (n=2) were administrated with 10 nmol crgp-zw800-1 and imaged at  $4 \text{ h}$ . Mice bearing orthotopic MCF-7 tumors (n=2) were injected with 10 nmol crgp-zw800-1 and imaged at 0.5, 1, 2, 4, 6, 8, 24, and 48 h post injection. One mouse with 2 mcf-7 tumors was administered with 10 nmol crgD-zw800-1 and imaged at 4 h post injection while its skin was partially removed. In vivo TBRS were calculated by dividing the mean fluorescence signals of tumors by the mean of surrounding tissue. Ex vivo organ-to-tumor ratios were calculated by dividing the fluorescence intensity of the organ by that of the tumor.

## **Pharmacokinetics**

crgd-zw800-1 was diluted in 0.05 mM hepes buffered mouse serum to the concentrations 5.00, 4.50, 4.00, 3.50, 3.00, 2.50, 2.00, 1.50, 1.00, 0.50, 0.25, 0.13, 0.06,0.03, and 0.02 µM. A calibration curve was created by measuring each concentration in 60 µL capillary tubes (Hirschmann® Laborgeräte GmbH and co. kg, Eberstadt, Germany) using the FLARE® with exposure times of: 1, 2, 5, 10, 25, 50, 100, 200, 500,1000, and 2000 msec. Data were plotted in fluorescence intensity over concentration (µM) (Figure s2, supplementary data). For each exposure time, the formula of the linear regression line was calculated. Subsequently, 5 mice without tumors were injected with 10 nmol CRGD-ZW800-1 via the lateral tail vein. The contralateral lateral tail vein was used to draw blood on the time points -5, 1, 6, 10, 20, 30, 40, 50, 60, 90, 120, and 240 min. post injection. Blood samples were absorbed using 60 µL capillary tubes and immediately measured using the flare® with the same exposure times. Saturated images and those under the detection limit were excluded from analysis. Each measurement was calculated back to its concentration using the matching formulas (Figure s3, supplementary data). The mean of each formula output was used as the concentration. Data was analyzed using NONMEM® software (Icon Development Solutions, Ellicott City, MD, u.s.a.). Data were plotted in logarithmic and linear concentration (µM) over time (min).

## **Statistical analysis**

For statistical analysis and the generation of graphs, GraphPad Prism software (version 5.01, GraphPad Software Inc., La Jolla, California, u.s.a.) was used. All values were reported using mean and standard deviation. Statistical significance for binding assay, *in vitro* and *in vivo* competition experiments, comparison of tbrs of the Pearl® and flare®, and comparison of fluorescence intensity of tumors was determined using the Holm-Sidak method. Differences in the biodistribution were calculated by two-way analysis of variance (anova) and Dunnett for posthoc testing. Group means were calculated for continuous data and medians were calculated for discrete data (scores). Test statistics were calculated on the basis of exact values for means and pooled variances.All tests were two-sided and in all an alpha of 5.0% was used.

**Figure 1** Binding assay and in vitro and in vivo competition experiments. Binding assay of crgd-zw800-1 on intermediate integrin-expressing ht-29 cells shows an almost linear increase in fluorescence intensity with increasing applied concentrations. In addition, fluorescence signals are increased when cells are incubated at 37°C compared to 4°C with the various concentrations ( $p < 0.0001$ ). This may be the result of enhanced internalization of the agent. *In vitro* competition experiment (1:1 ratio) using unlabeled crgd (competition group) showing decreased NIR fluorescent signals in both the  $a_V\beta_3$  positive u-87 MG (\* p < 0.05) as well as the α<sub>ν</sub>β<sub>3</sub> negative HT-29 cell line compared to the control without unlabeled CRGD. Healthy colon was used as background. *In vivo* competition experiment using a 200 times higher dose of unlabeled crgp compared to the dose of crgp-zw800-1 (1.24 mg, 2.0 µMol vs.15.5 µg,10 nmol) in the orthotopic ht-29 model. Compared to mice in the control group, a significant decrease in TBR was seen at  $4 h (* p=0.02)$  and  $24 h (* p=0.007)$ . Healthy colon was used as background.



**Figure 2** Near-infrared fluorescence imaging of colorectal cancer (ht-29)**.** Representative images captured using the original  $FLARE<sup>®</sup>$  prototype of the various concentrations at 4 h post injection. For the 30 nmol dose group the exposure time had to be set at 50 msec due to saturation of the near-infrared fluorescence images. For the other dose groups 200 msec was sufficient. nir fluorescence signals are clearly visible in 10 and 30 nmol groups only. Arrow=tumor; arrowhead=bladder. Near-infrared fluorescence imaging at 4 h post injection showed sufficient tumor-to-background ratios (i.e. ≥2) already in the 0.25 nmol (0.39 µg) dose group using the Pearl®. Surrounding tissue was used as background.Near-infrared fluorescence imaging using the Pearl® at 24 h post injection still showed sufficient tumor-tobackground ratios in the 1.0, 10 and 30 nmol dose groups. Healthy tongue tissue was used as background.

> 0.25 nmol 200 ms



b. imaging at 4 h post injection (pearl®)



1.0 nmol 200 ms 10 nmol 200 ms 30 nmol 4 200 ms  $\begin{aligned} \mathsf{r} & \mathsf{r} \mathsf{$ 



**Figure 3** Near-infrared fluorescence imaging of pancreatic (BXPC-3) and oral cancer (osc-19) . Upper panel: example showing the in vivo images of 30 nmol crgd-zw800-1 in the orthotopic pancreas model at  $4 h$  post injection using the original FLARE® prototype. Arrow=tumor. Lower panel: nir fluorescence microscopy of the tumor shows fluorescence inside the cells. Mean tumor-to-background ratios at 4 h post injection in the bxpc-3 model with the Pearl®. Skin was used as background. Upper panel: example of images from the orthotopic osc-19 model at 4 h using the original flare® prototype. Arrow=tumor. Lower panel: nir fluorescence microscopy shows the border of a resected osc-19 tumor. Clear fluorescent demarcation between normal and tumor tissue is shown (white dotted line). T=tumor; N=normal tissue surrounding the tumor. Mean tumor-to-background ratios of the optimal dose 10 nmol at 4 h in the orthotopic head-and-neck cancer model using the Pearl®.

#### a. imaging of pancreatic cancer (flare®)

#### b. imaging at 4 h post injection (pearl®)





 color-nir color nir merge c. imaging of oral cancer (flare®)



d. imaging at 24 h post injection (pearl®)







**Figure 4** Near-infrared fluorescence imaging of breast cancer (mcf-7)**.** Representative images captured with the Pearl® of the various time points after administration of 10 nmol crgd-zw800-1 in a xenograft breast cancer model. Arrowhead: bladder, arrows: tumors. All images are identically scaled.Tumor-to-background ratios over time measured using the Pearl®. Fluorescence intensity of tumors compared with 0.5 h post injection using the Pearl®.



Fluorescence

**Figure 5** Biodistribution of crgd-zw800-1**.** *In vivo* fluorescence intensity of bxpc-3 and ht-29 tumors at 4 h post injection of several doses of crgd-zw800-1 using the Pearl®. Significant differences (\* p<0.05; \*\* p<0.01) were seen in the 10 nmol dose group (p < 0.05). The *ex vivo* fluorescence intensity of BXPC-3 tumors were used to calculate the organ-to-tumorratios at 4 h post injection. Compared to the 10 nmol dose group, significant differences were seen between kidneys (1.0 nmol: p < 0.05; 30 nmol: p < 0.01 ), liver (1.0 nmol: p < 0.05) and large intestines (1.0 nmol: p < 0.05). In all dose groups, tumors were more than twice as fluorescent compared to skin, intestines, lungs, muscle, brain, gallbladder, and blood. *In vivo* fluorescence intensity of ht-29 tumors at 4 and 24 h post injection of several doses. Significant differences (\*) were seen in the 10 nmol dose group (p < 0.05).The *ex vivo* fluorescence intensity of ht-29 tumors were used to calculate the organ-to-tumor ratios at 24 h post injection. Compared to the 10 nmol dose group, significant differences (\*) were seen between kidneys (1.0 nmol and 30 nmol: p < 0.05). In all dose groups, tumors were more than twice as fluorescent compared to lungs, pancreas, muscle, gallbladder, brain, and blood.



**Figure 6** Pharmacokinetics of 10 nmol crgp-zw800-1. Graph shows the mean absolute concentrations (in µM) after intravenous administration of 10 nmol crGD-zw800-1 up to 240 min post injection. 1) auc calculated via trapezoid rule. 2) Half-life calculated via ln(2)/k10.



**Table 1** The expression of integrins αvβ3, αvβ5 and αvβ6 on tumor cell lines



The number of antigenic sites for  $\alpha_V\beta_3$ ,  $\alpha_V\beta_5$ , and  $\alpha_V\beta_6$  integrins per individual cell of the u-87 mg, HT-29, BXPC3, MCF-7, and OSC-19 cell lines using Qifikit (DAKO) for quantitative flow cytometric analysis.

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