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CT Imaging of Enzymatic Activity in Cancer using Covalent Probes Reveal a Size-Dependent Pattern.

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ABSTRACT: X-ray CT instruments are among the most available, efficient and cost-effective imaging modalities in hospitals. The field of CT molecular imaging is emerging which relies mainly on the detection of gold nanoparticles and iodine containing compounds directed to tagging a variety of abundant biomolecules. Here, for the first time we attempted to detect enzymatic activity, while the low sensitivity of CT scanners to contrast reagents made this a challenging task. Therefore, we developed a new class of nano-sized cathepsin-targeted activity-based probes (ABPs) for functional CT imaging of cancer. ABPs are small molecules designed to covalently modify enzyme targets in an activity-dependent manner. Using a CT instrument, these novel probes enable detection of the elevated cathepsin activity within cancerous tissue, thus creating a direct link between biological processes and imaging signals. Here we present the generation and biochemical evaluation of a library of ABPs tagged with different sized gold nano-particles (GNPs), with various ratios of cathepsin-targeting moiety and a combination of different polyethylene glycol (PEG) protective layers. The most potent and stable GNP-ABPs were applied for non-invasive cancer imaging in mice. Surprisingly, detection of CT contrast from the tumor had revers correlation to GNP size and the amount of targeting moiety. Interestingly, TEM images of tumor sections show intercellular lysosomal sub-cellular localization of the GNP-ABPs. In conclusion, we demonstrate that the covalent linkage is key for detection using low sensitive imaging modalities and the utility of GNP-ABPs as a promising tool for enzymatic based CT imaging.

INTRODUCTION

One of the leading causes of death in the Western world is still cancer. Regardless of tremendous progress in pharmaceutical and nanomaterial chemistry, early and accurate detection of the disease remains a major challenge ¹. The ability to visualize, characterize and quantify specific proteins and genes, and evaluate molecular pathways within the living organism will certainly improve the diagnosis and prognosis of various conditions. Traditional pathological evaluation is often done by microscopy, in which excised tissues are typically examined to characterize histological changes within the disease process. Molecular imaging in contrast, targets distinct molecular pathways *in vivo*, providing non-invasive visual and quantitative information for diverse applications ².

Computed tomography (CT) is an imaging method that uses X-rays to create cross-sectional images of the body. CT visualizes the internal structures of the body providing three-dimensional (3D) anatomic details, and is routinely used for diagnosis of diseases, prediction of therapy, and treatment assessment. Currently, it is one of the most commonly used non-invasive clinical imaging modalities owing to its wide availability, high efficiency, and cost effectiveness ³⁻⁴. Although it has high spatial resolution, the main drawback of the modality for molecular imaging remains the low sensitivity ACS Paragon Plus

and difficulty to distinguish subtle changes in soft tissue, particularly tumors which have similar X-ray absorption as their surroundings. We strongly believe that exogenous CT detectable agents that enhance tumor contrast will lead to better characterization and detection of cancer ⁵.

A wide variety of CT contrast agents have been developed in the last decade which focus mainly on iodine-containing probes and gold nanoparticles (GNPs) ⁶. Although iodinebased molecules are clinically used, they have some limitations, such as fast renal clearance and renal toxicity, making them incompatible for image guided surgery (in humans) or preclinical studies by micro-CT. Additionally, blood pool agents such as iodine-based probes, exhibit fast body dissipation leading to uniform enhancement of the entire body. In the case of CT imaging contrast enhancement, GNPs attenuate Xrays more efficiently than the iodine by many orders of magnitude. This is due not only to the much stronger attenuation per atom (gold vs. iodine) but also to the large number of atoms per gold nanoparticle. In addition, a previous study showed a very high LD50 for gold nanoparticles of 3.2 g Au/kg⁷, which is a great advantage for CT imaging where high dosages of a contrast agent are required. Most of the nanoparticle-based contrast agents accumulate in the tumor via passive targeting due to enhanced permeability and retention (EPR) effect over the course of hours to days ⁸. Nevertheless, no optimal particle Environment size or surface charge was identified for maximizing this effect due to the significant variability between tumors. Nanoparticles in the range of 10 to 100 nm showed the highest passive tumor penetration in animal models with minimal sublethal toxicity effect ⁹⁻¹⁰. However, to achieve functional imaging using CT instruments, active targeting of nanoparticles is obligatory ¹¹. Targeting probes to specific pathologies can significantly reduce the injection dose and thus minimize side effects and toxicity. As recently revealed, many techniques are used for GNPs' surface modification for multiple tumor markers such as antibodies, peptides or small molecules ¹². In general, the targeting efficacy of functionalized nanoparticles depends on the nature of the ligand, the selected coupling reaction and the ligand surface density.

In this study, we evaluate three different sizes of GNPs stabilized by poly ethylene glycol (PEG) coating conjugated via an Au-S bond providing a stable CT tag. The tag was easily and reproducibly conjugated to a targeting moiety, an active site inhibitor specifically targeted to cathepsin proteases, thus generating an activity-based probe (ABP). ABPs are tagged molecules that are designed to covalently modify enzyme targets in an activity-dependent manner thus reporting their activity. To this end, ABPs have been developed for a number of enzyme families, including proteases, kinases, phosphatases, glycosidases, and so on 13. Proteases recognize their substrate target by the amino acid sequence around their cleavage site; in most cases the recognition is up to four amino acids on each side of the scissile bond (the target site for cleavage). The covalent nature of ABPs is generated by an electrophile termed "warhead" which is placed close to the recognition element. Several cysteine protease ABPs include an acyloxymethyl ketone (AOMK) chemical group as their warhead, since it is highly selective for cysteine proteases ¹⁴.

Cysteine cathepsins are highly upregulated in a wide range of cancer processes by mechanisms ranging from gene amplification to post transcriptional modification. Within the tumor microenvironment a large variety of cell types highly express cathepsins including fibroblast, T cells, endothelial, osteoclasts, neotrophils cells and predominantly, macrophages 15. We have previously reported on cathepsin fluorescent ABPs that allow for visualization of active cathepsins B, L and S in tumor-bearing mice 16,17. These ABPs were based on the covalent and specific cathepsin inhibitor GB111-NH2 as their targeting moiety¹⁸. Cathepsins B, L and S are highly abundant within intracellular endolysosomal vesicles and are constitutively expressed, while cathepsin B can reach a concentration as high as one millimolar ¹⁹. Cathepsin activity is highly elevated in metastatic tumors, where they increase motility and invasion of cancer cells, mediate degradation of extra cellular matrix as well as angiogenesis ²⁰. Furthermore, many studies link the high cathepsin activity with prognostic and predictive factors in human cancers, prominently breast, colorectal, pancreatic, ovarian, and lung cancers. Cathepsin expression is commonly elevated in tumors compared to normal tissue, thus recently cysteine cathepsins were targeted for anticancer strategies within the tumor microenvironment ¹⁵. Based on this evidence we hypothesized that cathepsin activity will be sufficiently elevated in tumors allowing for its detection by GNPlabeled ABPs, overcoming the low sensitivity of CT scanners. Here, we explore the newly generated cathepsin-targeted GNP-ABPs for their potency, overall permeability and imaging capabilities. We based the GNP probes on our published potent and selective GB111-NH₂ cathepsin inhibitor. Our novel probes are beneficial since they retain the contrast element at the desired site, increasing the imaging sensitivity. Moreover, reporting on the levels of activated cathepsins may reveal critical information regarding disease progression, enabling more accurate diagnoses and better targeted treatments using highly abundant CT scanners.

EXPERIMENTAL SECTION

Synthesis of GB111-NH₂: The synthesis was performed as previously reported ¹⁶ and is described in details in **Fig S1**. **Synthesis of Ac-DWK-amide:** Ac-DWK-amide was prepared

on Rink resin (Advanced Chemtech) using standard solid phase peptide synthesis method, detailed description in Fig S2. Synthesis of PEGylated gold nanoparticles

GNPs stabilized in citrate buffer were purchased from Cytodiagnostic Inc., at three different sizes: 10, 30, 100 nm. A protective layer of PEG was absorbed on the surface of the GNPs at different molar ratios and different lengths consisting of a mixture of thiol-polyethylene-glycol (mPEG-SH) and a heterofunctional thiol- polyethylene-acid (SH-PEG-COOH) (Creative PEGWorks, Winston Salem, NC). In a typical process, 11 µl of PEG mixture solution were added to 1.0 ml of GNP solution (OD 1), at different concentrations to obtain the following final molar concentration of PEG: 0.01, 0.1, 1, 10 mM. The reaction mixture was vortexed immediately and then incubated at room temperature for 2 hours. After PEGylation, each solution was centrifuged at 14000, 9400, 8600 rpm (10, 30, 100 nm respectively) for 20 min and re-dispersed with Milli-Q water by gentle shaking. This process was repeated 3 times. From the resulting solution, a small part was analyzed by measuring absorption spectra, hydrodynamic size and zeta potential. The mixture was stored in a refrigerator at 4°C. PEGylation at 0.01 mM final concentration was found to be most efficient and stable.

Conjugation of GB111-NH₂ to PEGylated gold nanoparticles

PEG-GNPs were washed in 1 ml of DMF: DDW (1:10) several times. EDC (1.3 μ mol, 0.21 mg) and NHS (1.3 μ mol, 0.13 mg) 10 eq relative to the PEG-COOH moiety, were dissolved together and added to the PEG-GNP solution for 30 min at room temperature, providing active sites on GNPs that further underwent an amidation reaction with GB111-NH₂. The amount of peptide added was 0.38 mg (75 nmol) per 2.5 mg of molecular gold (1:10 ratio to activated moiety). The mixture was incubated overnight at room temperature followed by centrifugation and DMF washes to remove unbound peptides in the solution.

Conjugation of Ac-DWK-amide to PEGylated gold nanoparticles: The conjugation synthesis was performed similarly to GB111-NH₂ described above.

Chemical evaluation

Dynamic light scattering (DLS) and ζ potential measurements were conducted in DDW using a Zetasizer nano-ZSP (Malvern). Transmission electron microscopy (TEM) analysis was performed using JEOL JEM-1400Plus by applying ~10 μ l of samples resuspended in DDW to a 200 or 400-mesh copper grid covered by carbon-stabilized Formvar film (SPI, West Chester, PA). The samples were dried overnight before scans were performed at different kV.

Thermogravimetric analysis (TGA), to establish the mass ratio between organic moiety (PEG) and gold atoms, was performed using a Mettler Toledo instrument. Temperature Program: heat from 30 °C to 450 °C with rate of 10 °C/min in nitrogen atmosphere with a purge rate of 20 mL/min.

Recombinant cathepsin labeling

Recombinant human Cathepsin B, 0.7 μg in reaction buffer (50 mM acetate, 2 mM DTT and 5 mM MgCl₂ at pH 5.5), was treated in room temperature (RT) with inhibitor (*GNP-PEG-GB111*) 0.01-10 μM probe or vehicle for 1 hour. Indicated concentrations of GB123¹⁷, a fluorescent labeled cathepsin ABP, were added to samples for 30 min at RT. The reaction was stopped by addition of sample buffer x4 (40% glycerol, 0.2 M Tris/HCl 6.8, 20% b-mercaptoethanol, 12% SDS and 0.4 mg/ml bromophenol blue), samples were boiled, separated on a 12.5% SDS gel and scanned for fluorescence by Typhoon scanner FLA 9500, excitation/emission 635/670 nm.

Evaluation of probes permeability to intact cells, competition assay

NIH-3T3 cells ($1x10^5$ cells/well) were seeded in a twelve-well plate one day before treatment. Cells were treated with vehicle or 0.1,1,10 μ M probes that were pre-dissolved in 0.1% DMF, 0.9% DDW. After 4 or 24 hours of probe incubation, residual cathepsin activity was labeled with GB123 (1 μ M). Cells were washed with PBS and lysed by addition of sample buffer. Lysates were boiled for 10 min and centrifuged. Protein determination was performed by butterfly assay and separated by 12.5% SDS gel. Residual labeled proteases in cells were visualized by scanning the gel with a Typhoon scanner FLA 9500, excitation/emission wavelengths of 635/670 nm.

Cells viability assay

NIH-3T3 mouse fibroblast cells $(5x10^3 - 24 \text{ h}, 3x10^3 - 48 \text{ h})$ and 4T1 murine mammary carcinoma cells $(6x10^3 - 24 \text{ h}, 4x10^3 - 48 \text{ h})$ were seeded in a 96-well plate and incubated for 24 hours for cell attachment. GNP-ABPs or control GNPs, dispersed in fresh culture media were added at equivalent concentrations to each well (200 µ1) and incubated for 24 or 48 hours at 37 °C. The medium was discarded to remove free particles in the solution and cell survival was investigated by standard methylene blue assay.

Animal Care

All animals were maintained on a 12:12 hours light/ dark cycle under fixed conditions of temperature (23 °C) and humidity (50%), with free access to food and water. All experimental procedures were approved by the Animal Care Committee of Hebrew University of Jerusalem and in accordance with the NIH Guide for the Care and Use of Laboratory Animals.

In vivo imaging

4T1 cells were grown to subconfluency, followed by detachment with trypsin, spin down and resuspension in 0.5% BSA in sterile PBS and 25% matrigel. Cells (1x106 per spot in a total volume of 20 µl) were injected subcutaneously into the back of 3-4 week-old male BALB/c mice under isoflurane anesthesia. Tumors were typically established 9-11 days after cells injections, compounds were injected intravenously via the tail vein as follows: in a total volume of 200 µl from stock solution of 25 mg/ml (5 mg of GNP per mouse). Mice anesthetized with isoflurane were imaged prior to injection, and at 24, 48 and 72 hours post injection. The number of mice used is indicated in the figures. Micro-CT scanner (Skyscan High Resolution Model 1176) equipped with 64 detectors with a nominal resolution of 35 µm was used (0.2 mm aluminum filter, a tube voltage of 40 kV and 500 mA). Reconstruction was performed using SkyScanNRecon software. Ring artifact reduction, Gaussian smoothing (3%), and beam hardening correction (25%) were applied. Volume rendered threedimensional (3D) images were generated using SkyScan CT-Volume ("CTVol") software. To quantify the exact amount of GNPs that reached the tumor tissue at each time point, we analyzed the tissue absorbance and GNP absorbance for the same ROI apart. The number of voxels for soft tissue absorbance was characterized by low energies, hence was defined by us as 25-255 in gray scale, while gold (and bone) 53-255 for higher energies. The minimum threshold for gold detection was derived from a pre-scan of the mouse, showing no voxels in the tumor at that grayscale index. After the total voxel number was derived from the image for each range, the labeling percentage of voxels per tumor was calculated (% labeling = num. of vox53-255/num. of vox25-255).

Iv vivo imaging with inhibitor pre-treatment: tumor bearing mice were injected intraperitoneal injection (IP) with cathepsin B inhibitor GB111-NH₂ 40 mg/kg 5 hours before T-GNPs injection, and an additional dose of 20 mg/kg 48 hours post injection of T-GNP. Mice in all other treatment groups (T, NT and NS GNPs) received saline solution injection (IP) at the same time points (5 and 48 hours) as the pre-treated group.

Flame atomic absorption spectroscopy

Flame atomic absorption spectroscopy (FAAS, SpectrAA 140, Agilent Technologies) was used to determine amounts of gold in the investigated organs. The tissues obtained from the experimental animals were melted with aqua regia acid (a mixture of nitric acid and hydrochloric acid in a volume ratio of 1:3). The liquids were then evaporated, filtered and diluted to a final volume of 7 mL. An Au wavelength lamp was used to determine the gold concentration in the samples. A calibration curve with known gold concentrations was prepared (commonly: 1, 2, 5 and 10 mg/L). The gold concentration in each sample was determined according to its absorbance value with correlation to the calibration curve. Each sample was analyzed in triplicate and averages and standard deviations were taken. The measured amount of the gold was divided by the organs' weight.

Preparation of tissues for transmission electron microscony

Tissue was dissected from animal into PBS (Phosphate Buffered Saline pH 7.4) and fixed in 2% paraformaldehyde, 2.5% glutaraldeyde in 0.1M cacodylate buffer (pH 7.4) for several hours at RT. The tissues were then rinsed 4 times, 10 min each, in cacodylate buffer and post fixed and stained with 1% osmium tetroxide, 1.5% potassium ferricyanide in 0.1 M cacodylate buffer for 1 hour. Tissues were then washed 4 times in cacodylate buffer followed by dehydration in increasing concentrations of ethanol consisting of 30%, 50%, 70%, 80%, 90%, 95%, for 10 min each step followed by 100% anhydrous ethanol 3 times, 20 min each, and propylene oxide twice, for 10 min each. Following dehydration, the tissues were infiltrated with increasing concentrations of agar 100 resin in propylene oxide, consisting of 25, 50, 75, and 100% resin for 16 hours each step. The tissues were then embedded in fresh resin and let to polymerize in an oven at 60 °C for 48 hours. Embedded tissues in blocks were sectioned with a diamond knife on an LKB 3 microtome and ultrathin sections (80 nm) were collected onto 200 Mesh, thin bar copper grids. The sections on grids were sequentially stained with uranyl acetate for 5 min and lead citrate for 2 min and viewed with Tecnai 12 TEM 100 kV (Phillips, Eindhoven, the Netherlands) equipped with MegaView II CCD camera and Analysis® version 3.0 software (SoftImaging System GmbH, Münstar, Germany).

Preparation of tumor tissue for pathological evaluation

To evaluate the tumor model, 65% of the resected tumors obtained from an *in-vivo* experiment were arbitrarily collected and fixed with 4% paraformaldehyde at 4°C, transferred to 10% formalin neutral buffer solution and embedded in paraffin. Sections (10 μ M thick) were obtained, stained by using Hematoxylin and Eosin and examined under a light microscope.

Statistics

Statistical analyses for in vitro experiments were performed using Excel Software, error bars represent Standard Deviation. A two-tailed t-test was performed to determine significance when comparing the means of two groups. Alternatively, oneway ANOVA with Dunett's correction for multiple comparisons was performed using GraphPad Prism software to compare the means of gold accumulation measured by FAAS as indicated in the figure legend. Independent experiments were conducted with a minimum of two biological replicates per condition to allow for statistical comparison. Repeated measures analysis of variance (ANOVA) was performed in SPSS and applied to examine effects of T-GNP-ABPs uptake in vivo and tumor volume containing gold due to time (in four levels), size (three levels) and targeting (targeted vs. nontargeted). In addition, in the cases of significant differences between more than two categories, post hoc pairwise comparisons (with Bonferroni correction) were conducted. The alpha level was set at p < 0.0083.

RESULTS

Micro CT sensitivity evaluation

To choose the optimal contrasting agent as an ABP tag, we first evaluated the detection level of the various elements using dual energy micro CT. Scanning each contrast agent separately demonstrated linearity of the contrast with concentrations, thus enabling us to determine both the lowest concentrations detected in DDW as well as the attenuation efficiency (derived from the attenuation slope). While the minimal concentration of iodine and gadolinium detected was lower, the GNPs showed X-ray attenuation three times more efficiently, making gold an excellent tag for our probes. This is due not only to the much stronger attenuation per atom but also to the large number of atoms per GNP, **Fig. S3**.

GNP-ABP synthesis and characterization

In our attempt to generate efficient GNP-ABPs we first designed a library of cathepsin-targeted nanoparticles with different particle sizes, and coatings that were evaluated for various characteristics such as potency, cell permeability and in vivo properties. We selected three different sizes of GNPs to be linked to the targeting moiety: 10, 30 and 100 nm which eventually enabled us to validate the influence of particle size on the uptake profile. As the targeting moiety of the ABP, we selected GB111-NH₂, a potent and selective cathepsin B, L and S inhibitor successfully used in our previous reports ¹⁸. GB111-NH₂ was synthesized using a combination of solution and solid phase chemistry as we previously published ¹⁶ Fig. 1A. To generate the GNP-ABP first, a protective layer of PEG with different lengths and different molar ratios of capped mPEG-SH and a heterofunctional COOH-PEG-SH were absorbed on the surface of the GNPs Fig. 1B. The monofunctional PEG (mPEG-), capped with methoxy group, was used for stability and biocompatibility of the particles ²¹ while the COOH-PEG- moiety provided a chemical handle for further binding of GB111-NH₂. The most stable coating was achieved

with 0.01 mM final molar concentration of PEG mixture. Overall, we generated a library of GNP-ABPs containing combinations of different percentages of targeting moiety 10. 50 and 100% on the particle surface with different PEG lengths (i-vi Fig. 1C). A control probe was synthesized for each size of GNP, lacking the targeting moiety (vii Fig. 1C). After synthesis, spectroscopy measurements were taken to characterize the GNP-ABPs, including Transmission Electron Microscopy (TEM), light absorption, size and zeta potential, with the TEM images revealing the gold core remained intact. The absorbance peak changed within 2-4 nm from the parent GNP core indicating aggregation did not occur. The hydrodynamic diameter (HD) measured by dynamic light scattering (DLS) revealed 10-30 nm increase in the (HD) after PEGylation and targeting moiety coupling. In addition, each step of the chemical modification and stability was confirmed by zeta potential measurements showing a decrease in the negative charge. Correlation between all methods provided reliable analysis of our probes confirming GNP-ABP formation and gold core stability Fig. 1D-F. We measured the stability of a selected GNP-ABP, DTS-iv, and found it exhibited good stability in 10% DMF:DDW over 10 weeks Fig. S4. Furthermore, particles could be easily re-suspended without notable aggregation, with no change in surface potential and size detected. In order to proceed with these GNP-ABPs to biochemical evaluation, a quantitative analysis to determine the amount of targeting moiety on each particle was performed. Thermogravimetric analysis (TGA) was used to establish the mass ratio between organic moiety (PEG) and gold atoms. Since PEG has a simple linear-chain-bond structure, thermal dissociation occurs in a narrow temperature range providing a reliable feature to measure its quantity. We thus measured the weight differences between GNP-PEGs before and after thermal dissociation. Calculations enabled assessment of the average PEG density being 0.21 PEG/ nm² on the GNP which corresponds to the low PEG density particle group (density lower than 1 PEG/nm²) having a "mushroom" configuration. This result is consistent with the low concentration of PEG mixture taken for the process 22-23

Biochemical evaluation of GNP-ABPs

To determine the potency of various derivatives, we performed competitive inhibition of recombinant human cathepsins B. Indicated concentrations of the various 30 nm GNP-ABPs were added to cathepsin B followed by labeling of the residual activity with GB123, a Cy5 fluorescent cathepsin ABP ¹⁷. The detection of the residual cathepsin B activity was visualized by fluorescent scanning of the samples run on SDS PAGE, demonstrating reactivity of the 10% and 100% targeting probes (DTS- i, iii, iv) each to different extents, Fig. 2A. Surprisingly, derivatives containing 50% of targeting moiety on the surface (DTS- ii, v) showed no potency at all. We then focused on derivative DTS-iv and investigated the inhibition of recombinant cathepsin B using different sized GNPs; we found an interesting correlation of potency with size, the potency being reduced with increase in GNP particle size, Fig. 2B. Evaluation of the probes' cell permeability and capability of labeling cellular cathepsin enzymes was performed using competitive inhibition assay by incubating selected compounds with NIH-3T3 or 4T1 cells for 4 and 24 hours. Next, residual cathepsin activity was labeled by GB123. To assure that the binding is dependent on protease activity, a nontargeted control (NT-GNP) was applied as well (DTS-vii), cell lysates were separated by SDS PAGE and scanned for fluores-

cence, Fig. 2C. Results for NIH-3T3 cells after 24 hours incubation and 4T1 cells 4, 24 hours incubation are presented in Fig. S5. Encouragingly, all 10 nm GNP-ABPs derivatives tested showed high cell permeability and inhibition of cathepsin activity, DTS-iii however was less potent than DTS-iv. Within the 30 nm probes, DTS-i and DTS-iv (consisting of 10% GB111), showed a clear preference to shorter PEG (3 kDa) as a targeting moiety linker. Both 10 and 30 nm GNP-ABP were more potent than the 100 nm probes. Similar data were observed for 24-hour-probe incubation, Fig. S5. Dose response inhibition of cathepsin activity was detected with targeted GNPs, revealing a size-dependent pattern. Again, derivative DTS-iv exhibited the highest potency therefore was selected to continue to *in vivo* evaluation. Importantly, since no inhibition was detected by the control DTS-vii, we concluded that the binding of the probes occurred in an activitydependent manner. Prior to examining the GNP-ABPs in noninvasive imaging experiments, their toxicity was evaluated on NIH-3T3 and 4T1 cells, selected as models for normal and cancer cells, respectively. We defined the targeted derivative DTS-iv, T-GNP-ABP, and the non-targeted control DTS-vii, NT-GNP. Viability was determined by methylene blue assay after incubating the cells with increasing concentrations of T-GNP-ABP and NT-GNP for 24 and 48 hours. Only minor cytotoxicity was detected with 10 nm T-GNP-ABP at the highest concentration tested for 48 hours while surprisingly, notable cytotoxicity was detected with 10 nm NT-GNP at 0.3 mg/ml at both 24 and 48 hours, **Fig. 2D.** We thus concluded that working concentrations of ~ 0.15 mg/ml Au should be safe for animal use.

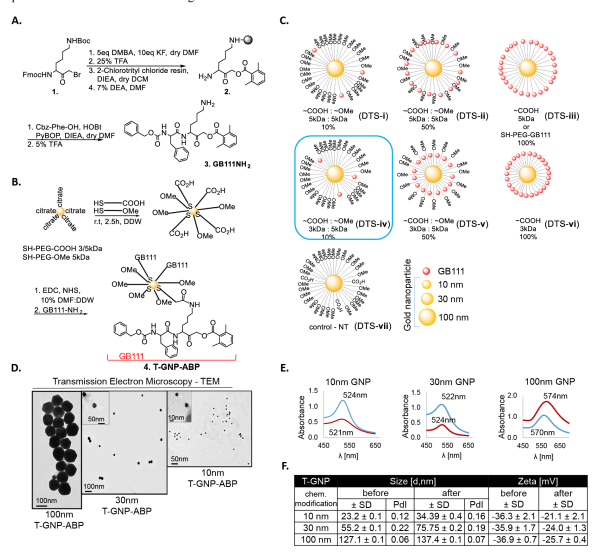


Fig. 1. Chemical synthesis and evaluation of Gold Nanoparticle Activity-Based Probes. (A) Synthesis of GB111-NH₂ as published in ¹⁶, lysine bromomethyl ketone was reacted with dimethylbenzoinc acid with KF, then the Boc protecting group was removed and the compound was loaded on resin to give (2). The Fmoc protecting group was removed with DEA and Phenylealanine capped with Cbz was coupled generating GB111-NH₂ (3). (B) Three different sizes of GNPs were coupled to combinations of different lengths of PEG protective layers, by PEG absorption on the surface of the GNPs. Different percentages of PEG-COOH moiety were used providing reactive sites on GNPs that were bound to (3) GB111-NH₂, to generate T-GNP-ABP. Non-functional PEG capped with methoxy group (OMe) were used for stability. (C) Schematic diagram representing various derivatives of GNP-ABPs that were generated. Characterization of selected GNP-ABPs: (D) TEM images of 10 nm, 30 nm, 100 nm GNP-ABPs. (E) Visible

spectra absorbance of GNP-ABPs exhibit a unique wavelength, before (bare GNP), red, and after chemical modifications, blue. (F) Size (DLS) and zeta-potential measurements of T-GNP-ABPs, represents the hydrodynamic diameter and surface charge. Correlation was found between all methods.

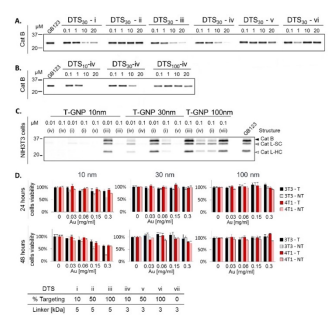


Fig. 2. Biochemical evaluation of GNP-ABPs. (A) Competitive inhibition of recombinant cathepsin B by the 30 nm GNP-ABP derivatives (Fig. 1). Indicated probe concentrations were incubated with recombinant cathepsin B for 1 h followed by 30 min labeling of the residual activity with GB123, a fluorescent ABP ¹⁷. Samples were separated on SDS gel and scanned for fluorescence at ex/em 635/670 nm. (B) Competitive inhibition of recombinant cathepsin B by DTS-iv derivatives of different particle sizes, size dependent inhibition was observed. (C) Inhibition of endogenous cathepsin activity in intact NIH-3T3 cells treated with probes in growth media for 4 h followed by labeling of residual enzyme activity by GB123. Control DTS-vii (NT-GNP) was applied. Whole cells lysates were separated on SDS gel and scanned for fluorescence. Highly potent inhibition of cathepsin activity was detected with DTS-i and DTS-iv probes. Size dependent pattern was observed, the smallest compounds showed highest potency. (D) NIH3T3 and 4T1 cells viability was determined by methylene blue assay after incubation with various concentrations of DTS-iv (T) or DTS-vii (NT) for 24 or 48 h. Only minor cytotoxity was observed with 10 nm T-GNP at the highest concentration tested. T-targeted, NT-non-targeted. Data are presented as mean \pm SD

Non-invasive imaging of cancer in a tumor mouse module

We continue with *in vivo* experiments to evaluate the efficacy of the ABP targeting methodology compared to passive accumulation of GNP alone. For this reason, we took a syngeneic mouse model in which 4T1 cells were injected subcutaneously to the back of male BALB/c mice. After tumors were established, each mouse received an intravenous (IV) injection (to the tail) of 5 mg of T-GNP-ABP (DTS-iv) or NT-GNP (DTS-vii) compounds of various sizes. Mice were scanned at 24, 48 and 72 hours post injection by a Micro-CT scanner.

To overcome the diversity in tumor size over the experiment duration we first measured the voxels specific to gold attenuation within the tumor (out of the full attenuation range), and then measured the total range of attenuations of the tissue (including gold) for the same ROI. We divided the gold voxels by the total voxels giving the percentage of gold containing voxels within each tumor, see Methods section for details. Similar to the in vitro data a marked size-dependent accumulation of both targeted and non-targeted GNPs was observed at 72 hours post injection as presented in volume-rendered 3D images. In general, accumulation of NT-GNPs in tumor tissues is thought to occur because of the EPR effect, slightly enhancing the CT signal ²⁴. Nevertheless, a pronounced enhancement of the CT signal was observed in tumors from T-GNP-ABP injected mice (10 nm and 30 nm) as compared to NT-GNP controls, Fig. 3A. The 2D axial cross section images revealed a similar pattern with a uniform accumulation of T-GNP-ABPs in the peripheral region of the tumor, presumably due to high interstitial pressure of solid tumors ²⁵, or higher cathepsin activity in that tumor region ²⁰, **Fig. 3B**. Furthermore, a distinct signal was already detected 24 hours post injection and became highly elevated at 72 hours post probe injection, excluding all 100 nm GNPs. The 100 nm GNPs exhibited poor active and passive accumulation in the tumor through all time points, Fig. 3C. While pronounced signals were detected in the tumors we also observed signals from the digestive system at all time points. This signal most likely results from the high mineral content of the chow as it was detected in the pre-scan images, Fig. 3D. Overall, a clear size-dependent accumulation was seen with the highest signal enhancement detected with 10 nm T-GNP-ABP and lowest tumor uptake with 100 nm.

Bio-distribution of GNP-ABPs

GNP bio-distribution was monitored to follow pharmacokinetics throughout the body and its accumulation in the tumors. After the last time point, the mice were sacrificed, then tumors and other selected organs were analyzed by flame atomic absorption spectroscopy (FAAS) determining the gold content in each organ, Fig. 4A. The T-GNP-ABPs and the control NT-GNPs showed uptake in the spleen, kidney and liver indicating both passive and active uptake. As previously published, contrast agents larger than 6 nm avoid renal clearance, hence are excreted from the blood pool by phagocytic cells in the reticuloendothelial system, occurring primarily in the liver and spleen, thus leading to accumulation of contrast in those organs 11. Based on FAAS analysis, GNP-ABPs excretion was detected over a 96-hour period. The GNP-ABPs all showed fecal excretion, with the 10 nm showing the highest clearance with a total 3.3% injected dose (ID), 30 nm 2.3% ID and 100 nm only 1.5% ID, Fig S6. Despite the targeting of our GNPs, the maximal tumor uptake was lower than in the liver or the spleen. Tumor uptake was prevalently stronger for T-GNP-ABPs than the NT-GNPs this finding was consistent both in the FAAS analysis and average tumor uptake, extracted from CT images Fig. 4A. A strong size-dependent uptake in favor of small GNPs was found in tumor tissue both in the T-GNP-ABPs and NT-GNPs. In a repeated measures ANOVA, comparing the changes in CT attenuation derived by time after administration, particle size, targeting a clear significant statistical difference between the targeted group and the nontargeted group in all sizes together was found (Wilk's Lambda P value = 0.034). However, when analyzing the 100 nm particles alone, there was no statistical difference, therefore we conclude that this effect is driven by the 10 and 30 nm size particles and not from the 100 nm particles. In a repeated measures ANOVA, a statistical significant difference was also found between the following parameters: Time and Size, Wilk's Lambda P value = 0.001, Time and Targeting, Wilk's Lambda P value = 0.034, and a trend when comparing Time, Size and Targeting Wilk's Lambda P value = 0.08. Furthermore, over 6% of the total tumor volume contained detectable

gold content both at 48 and 72 hours post injection of either 10 nm or 30 nm T-GNP-ABP while the maximal detectable gold content impressively reached over 10% at 72 hours post 10 nm T-GNP-ABPs injection, **Fig. 4B.**

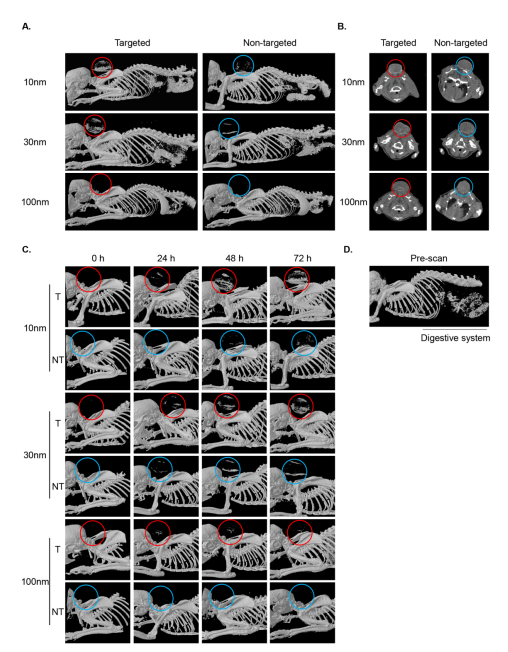


Fig. 3. Non-invasive micro-CT imaging of tumor bearing mice. (A) CT scans of mice 72 h post-injection of 10, 30 and 100 nm T or NT GNPs (5 mg gold per mouse). Representative CT images of volume rendered 3D images represent X-ray absorption of bones and gold, the circles indicate tumor location. (B) 2D axial cross section image of the same mice as in (A). Accumulation of T-GNP-ABPs at the tumor led to significant signal enhancement (shown in red circles). The highest enhancement was obtained with 10 nm T-GNP-ABPs. (C) Time course of all T-GNP-ABP and NT-GNP compounds. Images were acquired prior to probe injection (0 h) and 24, 48, and 72-h post probe injection. (D) Pre-scan image indicates contrast from the digestive system is attributed to the high mineral content of the chow.

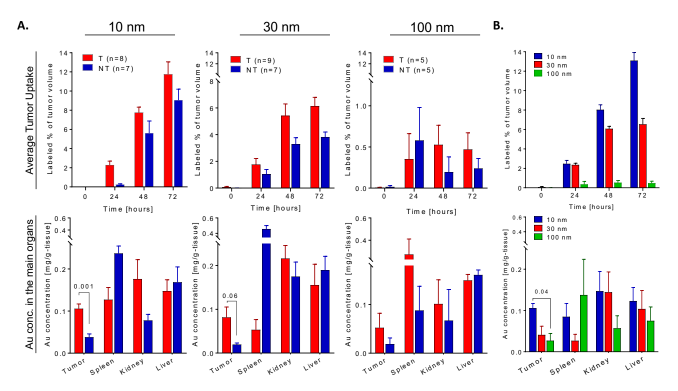


Fig. 4. T-GNP-ABPs uptake *in vivo* is size dependent. Quantification of gold attenuation in CT scans vs. gold concentration in main organs. (A) Top: Percentage of tumor voxels containing gold out of the total voxel in the tumors extracted from CT images of indicated times after IV injection of T-GNP-ABP or NT-GNP. Repeated measures ANOVA showed a significant (Wilk's Lambda P value = 0.034) difference between targeted and non-targeted groups across all measured sizes. Bottom: Average distribution of gold in the main organs 72 h post injection of T-GNP-ABP or NT-GNP detected by FAAS. T-Test analysis showed that both 10 nm and 30 nm T-GNP show higher accumulation in tumors compared to NT-GNP, P values are indicated. (B) Top: Comparison of different sized T-GNP-ABP, the percentage of tumor volume containing gold was analyzed from CT scans post injection as in (A); the highest signal was detected with the 10 nm T-GNP-ABP. The gold signal in the tumor increased over time and became highly elevated at 72-h post injection. Repeated measures ANOVA showed a significant difference between time and size (Wilk's Lambda P value = 0.001). Bottom: Gold accumulation in major organs detected by FAAS as in (A). Similar size-dependent accumulation was observed both by CT and FAAS. One way ANOVA showed a significant difference as indicated. The number in brackets represent the number of mice used. Bars represent standard error.

Since we detected high gold content in the tumors we attempted to visualize the cellular location of GNP within the cells. Tumors were resected 72 hours post IV injection of each goldcontaining compound and then subjected to TEM analysis. GNPs were clearly visualized in the lysosomes of tumor cells from 10 and 30 nm T-GNP-ABP injected mice, which is supported by the known cathepsins lysosomal cellular location ²⁶ . Small amounts of 10 and 30 nm NT-GNPs were also observed in the cells, most likely due to passive uptake. The 100 nm compounds however, showed low tumor cell presence with no difference observed between passive and active uptake, possibly due to low permeability and extensive spleen accumulation leading to short blood circulation; hence low availability of GNPs, Fig. 5. The TEM results further support the CT attenuation increase derived from T-GNP-ABPs that was detected by the CT scans localized within the tumor cells.

Evaluation of T-GNP-ABPs uptake efficiency

To further confirm that the signal detected and measured in the tumors was due to targeting moiety contribution and its covalent ability to target protease activity, we designed an additional control probe (in addition to the NT-GNP; DTS-vii). A non-specific GNP (NS-GNP; DTS-viii) bearing a short peptide, Ac-DWK-amide, (that lacks the ability to target cathepsin

B, L or S) was conjugated to 3 kDa PEG of a 30 nm GNP via an amide bond. This control was generated in order to mimic the surface potential of our leading compound T-GNP (DTS-iv), see schematic presentation in **Fig. 6A**.

Moreover, we applied an additional control group that included pre-treating animals with the inhibitor, GB111-NH₂, to block the cathepsin activity prior to T-GNP-ABP administration (T-GNP+GB111). All derivatives were designed based on 30 nm GNPs. To confirm our approach, we first evaluated endogenous cathepsin inhibition by the T-GNP compared to the new controls in 4T1 cells by competitive inhibition with fluorescent GB123 Fig. 6B. Encouragingly, 30 nm NT-GNPs and NS-GNPs derivatives showed no inhibition of cathepsin B as well as the Ac-DWK-amide peptide alone. The T-GNPs and the pre-treated (T+GB111) group showed similar inhibition efficiency as expected. Importantly, noninvasive CT imaging was studied in 4T1 tumor bearing mice to evaluate the efficacy of the ABPs' covalent targeting methodology compared to all three control groups (NT, NS and T+GB111) in a similar fashion as in Fig. 3. The inhibitor pre-treated group received the GB111-NH₂ inhibitor by intraperitoneal injections 5 hours before T-GNPs and an additional dose 48 h post injection of T-GNP. This inhibitor treatment regimen was chosen because

of the expected difference in pharmacokinetics, between the small molecule inhibitor (GB111-NH₂) and the nano-size probe. Mice were scanned at 0, 24, 48 and 72 hours post T-GNP injection by a Micro-CT scanner. In correlation with our in-vitro experiment, a significant CT signal was detected in the tumor of mice injected with T-GNP-ABPs while all threecontrol groups showed lower X-ray absorption Fig. 6C. We excluded that the differences in gold uptake was a result of tumor heterogeneity by pathological evaluation of the extracted tumors. All tumors demonstrated similar histological features without necrotic tissue except of one tumor (from the non-targeted group) Fig S7. Quantification of the gold attenuation in the CT scans as well as the FAAS analysis confirmed significant reduction in signal in the mice treated with the control systems, Fig.6 D, E. These results assure that the signal in T-GNPs injected mice was a result of the covalent modification of cathepsin activity. Overall, our data supports our hypothesis that cathepsin activity sufficiently elevated in tumors allows for its detection by T-GNP-ABPs.

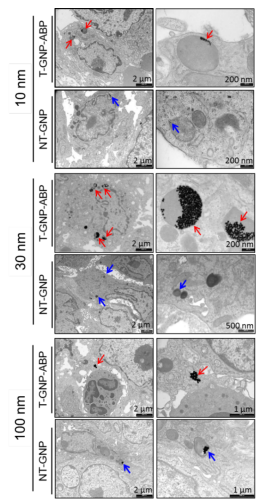


Fig. 5. TEM imaging of tumor tissue. TEM images of a section of tumor tissue taken from a tumor-bearing mouse 72-h post IV injection of 10 nm or 30 nm or 100 nm T-GNP-ABP or NT controls. Gold particles are marked with red arrows (T) and blue arrows (NT). Tumor uptake of T-GNP-ABP was clearly visualized for 10 nm, 30 nm and 100 nm. Lower amounts of NT-GNPs were observed in 10 nm and 30 nm; for

the 100 nm very little GNPs were detected with minor differences between T and NT compounds. Images depict localization of 10 nm and 30 nm T-GNP-ABP within the endosome/lysosome of tumor cells. Scale bars are marked.

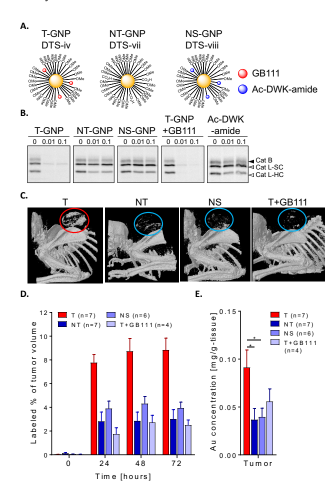


Fig. 6. Covalent targeting increases tumor uptake. (A) Schematic diagram representing GNP-ABPs DTS-iv (T-GNP) and control NT-GNP DTS-vii as in Fig. 1C. along with DTSviii, a non-specific (NS-GNP) control bearing off-target short peptide Ac-DWK-amide. (B) Activity of intact cathepsin activity within intact 4T1 cells was examined after GNP treatment in growth media for 24 h, followed by labeling of residual enzyme activity by GB123. In addition to T-GNP, NT-GNP and NS-GNP, cells treated with the cathepsin inhibitor GB111-NH₂ (1 µM) 2 hours prior to T-GNP addition (T+GB111) were examined. Whole cells lysates were separated on SDS gel and scanned for fluorescence as in Figure 2. Highly potent inhibition of cathepsin activity was detected after treatment with T-GNP and T+GB111 whereas NT and NS showed no cathepsin B inhibition. (C) Representative micro-CT scans (volume rendered 3D images) of tumor bearing mice 72 h post-injection of indicated GNPs (30 nm) is shown, circles indicate tumor location). Accumulation of T-GNP-ABPs at the tumor led to significant signal enhancement (shown in red circles) in all time points indicating the efficacy of covalent targeting of our probe. (D) Percentage of tumor voxels containing gold out of the total voxel in the tumors extracted from CT images of indicated times after IV injection indicated GNPs. Statistical analyses were performed by re-

peated measures ANOVA of the targeting group relative to all negative controls combined (NS, NT and T + Inhibitor), followed by Bonferroni post-Hoc tests, a significant difference was found with a Wilk's Lambda P Value <0.001. A graph depicting the statistical differences between the marginal means relative to the time is shown in **Fig S8**. (E) Average accumulation of gold in tumors 72 hours post injection of indicated GNPs detected by FAAS, One way ANOVA P*<0.05. The number in brackets represent the number of mice used. Data are represented as mean \pm SE.

DISCUSSION

The cysteine cathepsins are a family of proteases that, besides their main function of degrading proteases, play a critical role in several types of cancer such as melanoma, colorectal, glioma, breast, and lung ²⁸. Some members of the cathepsin family have redundant functions and some members, such as cathepsin B and L, have unique functions that play different roles in tumorigenic processes such as angiogenesis, metastasis, and invasion. The involvement of active proteases in various diseases led to the development of several protease-targeting clinical drugs, laying the basis for proteases as valid targets in pathological conditions. Here we exploited the highly overexpressed cysteine cathepsins to develop activity-dependent probes for molecular imaging of cancer. We described the development of a series of cysteine cathepsin-targeted ABPs linked to different sized GNPs coated with various combinations of PEG protecting layers. We showed the covalent binding of the T-GNP-ABPs to recombinant cathepsin and their ability to label and inhibit endogenous cathepsins within intact cells. After validating that T-GNP-ABPs have minimal toxicity, we showed a size-dependent uptake in subcutaneous tumors where the smallest particles showed significantly high accumulation, up to 12%, of the tumor volume (Fig. 4), thus enabling micro CT detection and overcoming its low sensitivity. The pronounced tumor accumulation was corroborated by FAAS studies and TEM images that clearly showed lysosomal localization within the cells. The significantly high amounts of T-GNP-ABP found in cancerous tissue are a combination of both the EPR effect and the covalent nature of the probes bound to the locally, highly-expressed cathepsin. Covalent suicide inhibitors however, often raise a potential problem lacking signal amplification since the target protease is inhibited upon binding. Nevertheless, the T-GNP-ABPs elegantly showed the potential use of the ABP methodology for functional CT imaging of cancer. Furthermore, by remaining bound to the target enzyme, a direct link between imaging signals and the biological process is made. One might suggest that smaller particles can't produce efficient contrast enhancement due to lower amounts of gold atoms per voxel, however our data suggest a higher accumulation of the smaller particles thus generating significant CT signals. This might be explained by the fact that smaller particles are engulfed more intensively to the lysosome compartment, leading to higher local concentrations of our probe, hence higher contrast 29-30 Active targeting of GNPs has grown extensively in the past few years, due to the ease of surface activation, with many pathological conditions being examined and evaluated by noninvasive CT imaging including multiple tumor receptors, lymph nodes, inflammation, and atherosclerosis 12. Furthermore, inherent targeting of high-density lipoprotein particles ³¹ and cell labeling by GNPs were also successfully accomplished ³². However, to the best of our knowledge there have been no reports on enzyme activity monitoring by the CT technique, most likely due to low sensitivity concerns. The goals of this work were to evaluate whether enzymatic activity can be detected by micro-CT, to assess the additional effect of protease targeting strategy compared to passive accumulation, and to select the optimal GNP size for visualization *in vivo*. We showed that cathepsin activity is sufficiently elevated in cancer to allow for its detection by T-GNP-

ABPs, generating additional contrast above the signal generated by passive uptake of similar sized GNPs. To generate functional compounds, various protective layers were used in our chemical scaffolds designed from two lengths of PEG, 3 or 5 kDa, in various ratios of reactive vs. capped PEG. The derivatives were studied in terms of their potency towards recombinant and endogenous cathepsins, and our results show that probes with 10% targeting moiety (GB111) linked to the shorter PEG (DTS-iv) possess high potency in all GNP sizes. The suggestive "mushroom" configuration may explain this enhanced potency of DTS-iv when GB111 is linked to the shorter PEG, enabling its exposure, rather than to the longer PEG that may be folded. As suspected, the similarity in sizes between GB111 and its PEG linker is advantageous 33. The bio-distribution studies of targeted GNP-ABPs reveal a sizedependent tumor uptake. Apart from the tumor, the spleen, liver and kidneys exhibited considerable amounts of GNP-ABPs yet, with different bio-distributions for each GNP size. Additionally, to take into account tumor variability and potential clinical applications, evaluation of the 10 nm and 30 nm T-GNP-ABPs should be considered in other tumor models. As previously suggested by S. Hirn et al., negatively-charged mono-sulfonated triphenylphosphine GNPs (TPPMS-GNPs) exhibited a hepatobiliary fecal clearance pathway from rat liver into the small intestine showing an inverse linear relationship to their diameter ⁹. The excretion kinetics assessment of our probes follows a similar tendency. Furthermore, the excretion of GNP-ABPs was slow but steady, thus partially reducing the concern of long term toxicity in the liver. Nevertheless, the 10 nm GNP-ABPs showed the highest liver uptake, as well as some toxicity in intact cells (Fig. 3D) raising the necessity to better understand long term effects.

CONCLUSION

In conclusion, we found our covalent GNP-ABPs as promising new tools for functional imaging of specific protease activity *in vivo* using the CT instrument. The covalent linkage is key for detection using the low sensitive imaging modalities. CT provides 3D anatomic details of the body, thus, it is widely used in clinical and preclinical practice for diagnosis of disease, prediction of therapy and treatment assessment. We screened and evaluated several types of GNP tags for most efficient and selective targeting of cysteine cathepsins. Tumor accumulation of the targeted GNP-ABPs is determined by the delicate balance between the targeting moieties' features and particle size. We show that the chemical structure of the particle strongly influences their biological behavior. We found

our lead compounds DTS-iv 10 nm and 30 nm to be highly effective for tumor imaging, its small size enabling excretion mainly through the hepatobiliary system, which appears to be a sufficient way to remove metal nanoparticles from the body. Still, the main limitation of our technology is the enhanced absorption in organs such as the liver, kidney and spleen where there is high cathepsin activity, making imaging of tumors in these organs challenging at this point. However, further developments in CT technology could improve the sensitivity and specificity limitations expanding the potential utility of our reagents. Additionally, Bindin et al. analyzed different stages of breast, lung and cervical tumors using multiplex zymography of elevated activities of cathepsins K, L and S 34 Thus, real-time measurements of tumor growth, metastasis formation and cancer staging could potentially be performed in large living organisms using our method of cathepsin activity measurements.

Moreover, after calibration of our system we suggest our platform to be adopted to target additional types of pathologies with high cathepsin activity such as atherosclerosis ³⁵ and arthritis ³⁶. The strategy described here overcomes a major limitation of signal detection in deep tissue that still exists with fluorescent protease substrates and fluorescent activity-based probes that have been extensively reported on during the last decade. Thus, the ABPs reported here have the potential to be developed for clinical use.

ASSOCIATED CONTENT

Supporting Information.

Figure S1. Chemical synthesis of GB111-NH₂

Figure S2. Chemical synthesis of Acetylated - Aspartic acid – Tryptophan – Lysine - amide

Figure S3. Detection level of various contrast agents by micro-CT

Figure S4. Characterization of GNP-ABPs

Figure S5. Biochemical evaluation of GNP-ABPs

Figure S6. Targeted GNP excretion is size depended in-vivo

Figure S7. Pathological sections of tumor tissue

Figure S8. Estimated Marginal Means of Measures

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Notes

The authors declare no competing financial interest.

Author Contributions

The manuscript was written through contributions of all authors, and all authors have given approval to the final version of the manuscript.

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