Innovative NIR fluorescent probes for an improved tumor detection *in vivo*

Doctoral Thesis

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submitted by

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AFFIDAVIT

By this I declare that I independently authored the presented thesis:

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Furthermore I declare that I carried out the scientific experiments following the principles of Good Scientific Practice according to the valid "Richtlinien der Georg-August-Universität Göttingen zur Sicherung guter wissenschaftlicher Praxis".

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PUBLICATIONS

Pauli J, Brehm R, Grabolle M, Behnke T, <u>Mathejczyk J</u>, Hamann F, Alves F, Hilger I, Resch-Genger U. Dye-biomolecule conjugates and NIR-fluorescent particles for targeting of disease-related biomarkers. Proc SPIE. 2011;7910,791014

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PATENT APPLICATION

Resch-Genger U, Behnke T, Würth C, Hoffmann K, <u>Mathejczyk JE</u>, Alves F, Stühmer W. Breitbandig absorbierende und emittierende NIR-Fluorophore mit großem Stokes shift für den Einsatz als Farbstoffe in core-shell-Nanopartikeln und als optische Sonden und Sensoren für die Biomarkeranalytik und als Komponenten von FRETSystemen. Patent no. 16717P-DE, submitted October 2011

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ABSTRACT

There is a great demand for the development of highly fluorescent and specific probes that enable a sensitive and early detection of malignancies and a separation from healthy tissue in a non-invasive way by in vivo imaging. The main drawback of fluorescence imaging in vivo is the autofluorescence, which hampers the detection of probe-derived signals. In this study, two novel fluorescence probes were investigated concerning their suitability for visualization of tumors in mice: a pH-sensitive dye, CypHer5E conjugated to the tumor-specific antibody Herceptin (pH-Her) as well as polystyrene nanoparticles (PSNPs) of different sizes (15, 25, and 100 nm) and surface modifications (NH₂, polyethylene glycol, and Herceptin) loaded with the broadband fluorophore Itrybe. For this purpose, time-domain near-infrared (NIR) fluorescence imaging was applied in nude mice bearing orthotopic HER2-positive (KPL-4) and HER2-negative (MDA-MB-231) breast tumor xenografts. The pHsensitivity and functionality of pH-Her was confirmed by spectroscopic measurements and in cell-based assays. In HER2-positive tumor-bearing mice, pH-Her, which increases fluorescence in the acidic environment of the tumor only, combined with subtraction of autofluorescence led to an increased contrast to noise ratio (CNR) and thereby an improved sensitivity of tumor detection, when compared to the always-on fluorescent conjugate Alexa Fluor 647-Herceptin (Alexa-Her). In contrast, LT-gated imaging with pH-Her and Alexa-Her did not improve tumor-detection sensitivity in vivo. The broad excitation and emission spectra of Itrybe enclosed in PSNPs enable excitation and sensitive detection of this dye at different wavelengths in the NIR region, in vivo. Furthermore, Herceptin-conjugated 100 nm NPs bind specifically to HER2 as demonstrated in immunoassays as well as on KPL-4 tumor cells and sections in vitro. However, a certain amount of unspecific binding/uptake was observed. In vivo accumulation of PEGylated NPs of all sizes was detected in MDA-MB-231- but not in KPL-4 tumors as shown by ex vivo scans of excised tumors. Biodistribution analyses in healthy mice illustrated a high and rapid uptake of NPs of all sizes in the liver, accompanied by a fast elimination of NPs from the blood, thereby reducing efficient tumor accumulation. In conclusion, the pH-sensitive dye, CypHer5E in combination with tumor-specific ligands is highly suitable for sensitive in vivo monitoring of tumors and may also be a promising tool for the detection of weak signals deriving e.g. from small metastatic lesions. Highly fluorescent Itrybe-loaded NPs have great potential for imaging applications as they provide 3D platforms for various modifications and thus help to address different biological questions in vivo. However, the bioavailability of Itrybe-loaded NPs has to be improved by modification of their surface or composition to enable efficient tumor targeting *in vivo*.

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LIST OF ABBREVIATIONS

ADCC	Antibody dependent cellular cytotoxicity
AEC	3-amino-9-ethylcarbazole
Alexa-Her	Alexa Fluor 647-Herceptin
Alexa-OH	Hydrolyzed Alexa Fluor 647
ATCC	American type culture collection
BAM	Bundesanstalt für Materialforschung und -Prüfung
BCA	Bicinchoninic acid
ß-gal	ß-d-galactosidase
BSA	Bovine serum-albumin
CCD	Charge-coupled device
cDNA	Complementary DNA
CD31	Cluster of differentiation 31
CNR	Contrast to noise ratio
СТ	Computed tomoghraph(y)
Ct	Cycle threshold
Су	Cyanine
DAPI	4',6-diamidino-2-phenylindole
DEPC	Diethylpyrocarbonat
DMEM	Dulbecco's modified eagle medium
DMSO	Dimethyl sulfoxide
DMSZ	Deutsche Sammlung von Mikroorganismen und
	Zellkulturen
DNA	Deoxyribonucleic acid
dNTPs	Deoxynucleoside triphosphates
DP	Dye to protein
DPBS	Dulbecco's Phosphate Buffered Saline
dT	Deoxythymine
DTT	Dithiothreitol
ECD	Extracellular domain
EDTA	Ethylene-diamine-tetraacetic acid
EGFR	Epidermal growth factor receptor
EPR	Enhanced permeability and retention
FAM	Fluorescein amidite
FCS	Fetal calf serum
FDA	U.S. Food and Drug Administration
FMT	Fluorescence-mediated (-molecular) tomography(y)
FRET	Förster (-fluorescence) resonance energy transfer

FRI	Fluorescence reflectance imaging
GFP	Green fluorescent protein
GI	Gastrointestinal
HD	Hydrodynamic diameter
HE	Haematoxylin-Eosin
HEPES	4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid
HER2	Epidermal growth factor receptor 2
hHER2	Human HER2
HRP	Horseradish peroxidase
hTfR	Human transferrin receptor type 1
ICG	Indocyanine green
IgG	Immunoglobulin G
i <i>.p.</i>	Intraperitoneal(ly)
i.t.	Intratumoral
i.v.	Intravenous(ly)
LT	Lifetime
MPS	Mononuclear phagocytic system
MRI	Magnetic resonance imaging
mRNA	Messenger RNA
NC	Normalized counts
NIR	Near-infrared
NIRF	Near-infrared fluorescence
NPs	Nanoparticles
PAGE	Polyacrylamide gel electrophoresis
PBS	Phosphate buffered saline
PCR	Polymerase chain reaction
PEG	Polyethylene glycol
PeT	Photon induced electron transfer
PET	Positron emission tomography(y)
pH-Her	CypHer5E-Herceptin
рН-ОН	Hydrolyzed CypHer5E
PI	Total fluorescence intensity of the injected probe
PLGA	Poly(lactic- <i>co</i> -glycolic acid)
PS	Polystyrene
PSNPs	Polystyrene nanoparticles
QDs	Quantum dots
RNA	Ribonucleic acid
rpm	Rotations per minute

RT	Room temperature
RT-PCR	Reverse transcriptase-PCR
S.C.	Subcutaneous(ly)
SCID	Severe combined immunodeficiency
SDS	Sodium dodecyl sulfate
SPECT	Single photon emission computed tomograph(y)
TBS	TRIS buffered saline
TBST	TRIS buffered saline containing Tween-20
TIV	Total fluorescence intensity in the tumor normalized to
	the tumor volume
ТМ	Melting temperature
TPSF	Temporal point spread function
TRIS	Trishydroxymethylaminomethane
Tween 20	Polyoxyethylenesorbitan monolaurate
US	Ultrasonography
WST1	WST-1[2-(4-Iodophenyl)-3-(4-nitrophenyl)-5-(2,4-
	disulfophenyl)-2H-tetrazolium]
λ_{em}	Emission wavelength
λ_{ex}	Excitation wavelength
3D	Three-dimensional

1. INTRODUCTION

A considerable progress in the diagnostic and monitoring of cancer was the development of noninvasive imaging methods that enable detection and characterization of malignancies over time in living objects. For the anatomical illustration of tissue, structural imaging techniques are applied, like computed tomography (CT), ultra sonography (US), and magnetic resonance imaging (MRI). Characterization of cancerous tissue by its altered functionality, concerning e.g. irregular metabolic activity or altered expression of proteins is facilitated by functional imaging methods, such as positron emission tomography (PET), single photon emission computed tomography (SPECT) and optical imaging. Each technique has a combination of advantages and disadvantages affecting its selection for application in a particular study¹.

Optical imaging by the use of fluorescence light has become especially attractive due to its comparatively easy use and low costs as well as high sensitivity². In addition and in contrast to most other imaging techniques, fluorescence imaging, enables the visualization of multiple target structures or biological processes in one model organism by simultaneously using different fluorescence labels³. While the current clinical use of fluorescence imaging is mainly limited to microscopic and (micro)endoscopic techniques for diagnostic purposes and during surgical procedures, the routine application of fluorescence methods in the clinic and preclinic is certain to increase¹. In recent years the use of functional tumor-specific fluorescence probes has become focus of several studies that aimed to improve sensitivity of tumor detection in tumor-bearing mice⁴⁻⁸. However, the majority of these probes is still limited in their specificity and sensitivity, regarding penetration depth and real-time imaging of biological processes *in vivo*.

Therefore, there is an ever growing demand for the development of highly fluorescent and specific as well as activatable probes that enable a sensitive and early detection of malignancies and metastatic lesions and a precise separation from healthy tissue in a non-invasive way *in vivo*.

1.1. The principle of fluorescence imaging

Fluorescence imaging relies on the detection of photons, emitted by a substance that has absorbed light of a distinct wavelength. Emission of fluorescence takes place when an orbital electron of an atom, molecule or nanostructure, after being excited to a higher quantum state, relaxes to its ground state, thereby emitting light. Different techniques can be applied to measure fluorescence in living organisms. Most systems are based on planar measurements of fluorescence that are usually overlaid with photographic images of the surface of the scanned sample. Normally such devices consist of a multispectral light source or a laser for excitation of the fluorescent species, appropriate filters and a device for signal detection, like a CCD camera or a photo multiplier tube. Planar imaging can be performed in an epiillumination mode, known as fluorescence reflectance imaging (FRI), in which the excitation source and the detector are located on the same side thereby measuring the returning fluorescence response. When the excitation source and the detector are placed on the opposite sides of the scanned object, imaging is performed in a transillumination mode, in which the light penetrating the object is collected⁹. Recent developments in tomographic fluorescence imaging including fluorescence-mediated (-molecular) tomography (FMT) provide three-dimensional (3D) information about the fluorescence distribution, as well as probe concentration and depth^{1,9-11}.

The introduction of time-domain imaging has demonstrated a great improvement in fluorescence imaging as it allows the calculation of the fluorescence lifetime $(LT)^{12}$. This parameter characterizes a fluorescence probe by describing the average time a fluorophore spends in its excited state before returning to the ground state. In this method FRI is applied with a pulsed laser diode as an excitation source. After each single excitation pulse the emitted photons are measured time-resolved using e.g. a photomultiplier tube, thereby allowing calculations of the LT¹³. As each fluorescence probe has a characteristic LT, the determination of LTs permits the identification of probe-derived signals, e.g. originating from different fluorescent probes¹⁴ and/or their separation from autofluorescence^{1,2,15,16}. *In vivo* autofluorescence is defined as fluorescence that derives from unlabeled sources, like collagen, elastin, beta carotene, or oils, pigments, and proteins from ingested food¹⁷. This endogenous fluorescence poses one of the biggest problems in *in vivo* fluorescence imaging as it hampers the detection of probe-derived signals. This could be overcome by gating fluorescence intensity maps of scanned objects to the characteristic LT of the applied fluorescence probe, thereby only illustrating probe-derived signals. If the LT of the probe substantially differs from the LT of unspecific signals in this object, e.g. owing to autofluorescence, this method enables the selective illustration of only probederived signals and thus might lead to an improved sensitivity in detection of the target.

1.2. Fluorescence labels

For optimal detection of fluorescence light *in vivo* it is important to consider fluorescence probes that are very bright, relatively photostable and require minimal

exposure to excitation light in order to sustain cell viability¹⁸⁻²⁰. Of particular advantage for *in vivo* imaging are fluorophores absorbing and emitting in the near-infrared (NIR) wavelength region of 650 to 900 nm. The benefits of this spectral range are not only a decreased autofluorescence but also less absorbance by haemoglobin, water and lipids, allowing an increased tissue penetration of up to 2 cm^{1,21} (Figure 1). In addition, the considerably high absorption by melanin is decreasing towards longer wavelength^{1,21}.



Figure 1. Tissue optical window

The absorption (illustrated as absorption coefficient) by major light absorbers in tissue is illustrated at distinct wavelengths. In the NIR optical window (700 to 900 nm; light blue) absorption by hemoglobin (Hb), oxygenated hemoglobin (HbO₂), and water (H₂O) is low. Picture was taken from Phan et al.²².

Different types of fluorescence labels are applied for *in vivo* imaging and can be divided into endogenous fluorescence labels and exogenous fluorescence probes that can further be grouped in non-targeted contrast agents, target-specific fluorescence probes, and activatable probes.

1.2.1. Endogenous fluorescence labels

Endogenous fluorescent labels are produced by genetic manipulation of the model organism leading to the expression of a fluorescent protein in order to visualize distinct biological functions. A break-through in the application of fluorescence imaging was the identification of the first fluorescent protein, green fluorescent protein (GFP) that was isolated from jellyfish, and purified and characterized in the early 1970's²³. Today fluorescent proteins in many colors exist that fluoresce in the visible spectrum in colors of deep blue (~400 nm) to far red (~670 nm). In addition, NIR fluorescent proteins have been expressed in mammalian cells^{24,25}. Such genetically encoded probes have been shown to be suitable for *in vitro* and *in vivo* fluorescence imaging in a variety of different cellular systems and transgenic organisms.

1.2.2. Exogenous fluorescence labels

Most of the fluorescent probes are used to specifically target particular markers or molecular events in model organisms. Requirements for the efficient detection of exogenous labels are for instance, absorption- and emission-maxima in the NIR region, a high molar absorption coefficient at the excitation wavelength, a high fluorescence quantum yield, as well as sufficient thermal and photochemical stability^{19,20}. Furthermore, suitable fluorescence dyes have to be soluble and stable in buffers and body fluids and the resulting fluorescence probe should show a good targeting efficiency with minimal nonspecific binding as well as a low toxicity²⁰.

There are few classes of near-infrared fluorescence (NIRF) dyes available for *in vivo* applications. Most of them are cyanine dyes, such as indocyanine green (ICG), cypate, Alexa Fluor-dyes, or Cy-, DY-, and IR-dyes^{19,26}. Although a large variety of fluorescent probes is continuously developed and improved, ICG is the only dye approved by the FDA (Food and Drug Administration-USA) for use in patients^{27,28}.

To characterize or monitor malignancies *in vivo* by the use of fluorescence probes, appropriate model organisms have to be selected. In this regard, nude mice are very suitable. These hairless mice discovered in 1962²⁹ have a mutation in the forkhead box N1 (*Foxn1*) gene, leading to defective differentiation of hair and thymic disgenesis and therefore resulting in T cell deficiency and partial defects in B cell development^{30,31}. As the immune system of these mice is compromised, they do not reject human tumor cells and thereby provide excellent model organisms to study tumor biology *in vivo*³². In addition, the lack of fur presents an advantage for *in vivo* fluorescence imaging as fur can cause scattering and absorption of light, hampering the detection of probe-derived signals.

1.2.2.1. Non-targeted contrast agents

Non-targeted contrast agents are designed to passively accumulate in the tissue of interest. Such probes are mainly applied for imaging of blood vessels and/or tumors. An important prerequisite for such probes is a sufficiently long blood circulation time, allowing their accumulation at the site of interest. Tumor imaging by non-targeted probes is enabled by the enhanced permeability and retention (EPR) effect, involving hyperpermeable and leaky vasculature, as well as suppression of lymphatic function³³.

1.2.2.2. Target-specific fluorescence probes

Target-specific fluorescence probes are directed against molecular markers that are overexpressed or uniquely expressed on the target structure. The majority of such probes consists of fluorescence dyes or nanomaterials conjugates to ligands, like whole IgG antibodies, antibody fragments, or small peptides, all targeting a tissue or disease specific marker. Numerous aspects have to be considered when choosing the right ligand for the design of target-specific probes. Most important, the ligand should show maximal targeting efficiency and minimal unspecific binding²⁰. Furthermore, the biodistribution and pharmacokinetics of ligands are strongly influenced by their size. Molecules smaller than antibodies such as antibody fragments or peptides, can be advantageous for *in vivo* imaging due to a rapid clearance from blood, leading to a better signal to background contrast. However, if clearance is too fast, small ligands will have insufficient time to penetrate the target tissues before being cleared from the blood. In comparison, larger whole antibodies are cleared more slowly from the blood circulation and only slowly penetrate into tissues, which may result in a high background and subsequently a poor tissue to background contrast³⁴. Furthermore, the so called "binding site barrier", postulated by Weinstein in the early '90s^{35,36} can influence the diffusion of ligands into the target tissue. This may be caused by a (too) high affinity and/or interaction of a particular ligand for/with its antigen, which may result in a reduced or even inhibited tumor penetration³⁵.

1.2.2.3. Activatable fluorescence probes

Activatable fluorescence probes (also termed "smart probes") evolved to detect and monitor molecular events, such as the activity of enzymes, changes in pH, or oxygen concentration, either by switching their fluorescence "on" or "off" or due to shifts in fluorescence absorption and/or emission spectra in response to distinct molecular processes. Such probes may be applied to selectively illustrate target structures, such as tumor tissue, based on a selective activation at the targeted tumor site only, thereby circumventing background fluorescence caused by unbound probes in the blood stream.

Probes based on the "on" and "off" principle require efficient quenching in the "off"state. The term quenching refers to processes which lead to a decrease of the fluorescence intensity of a given probe and can be achieved by distinct photochemical mechanisms, such as Förster (or fluorescence) resonance energy transfer (FRET), Hdimer formation or photon induced electron transfer (PeT).

FRET takes place between two molecules that are in close contact to each other. Here, a donor chromophore, initially in its electronic excited state, may transfer energy to an acceptor chromophore while dropping down to its ground state. FRET may occur between two fluorophores of the same type (homo-FRET) which absorb energy from each other (self-quenching) that would have otherwise resulted in fluorescence³⁷, or between an excited donor fluorophore and a quencher (hetero-FRET) that absorbs the energy from the donor and gives it off as heat^{6,37}. Furthermore, some fluorophores, like ICG tend to perform autoquenching after conjugation to antibodies, most likely occurring via a noncovalent interaction between the ICG and hydrophobic amino acids on the IgG^{7,37}. In addition, some dyes, such as Xanthene derivates at high concentrations or protein-conjugated fluorophores when in close proximity to each other, like the rhodamin dye TAMRA, tend to form H-dimers⁵. This can lead to shifts of absorbance spectra resulting in quenched emission spectra^{5,37}. In PeT, electron transfer from a PeT donor to an excited acceptor fluorophore, diminishes the fluorescence signal of the acceptor but when the donor is cleaved from the fluorophore, fluorescence occurs. Intramolecular PeT occurs when one molecule comprises an electron donor as well as an acceptor fluorophore³⁷⁻³⁹.

Activatable NIR fluorophores are applied for instance for sensing of proteases, where signals are generated as a result of lysosomal protease activity after probe internalization in cells^{1,4}. Furthermore, activatable probes are used as substrates for bacterial β-d-galactosidase (β-gal), thereby enabling the measurement of β-gal activity *in vivo*^{1,40-42}. In addition, activatable NIR probes can be applied for sensing of metal ions, such as calcium, mercury, and zinc or for sensing of nitric oxide⁴³ as well as oxygen^{44,45}. NIR fluorescent lanthanide-based complexes (e.g. Tb³⁺ or Eu³⁺) exist that may be applied as markers of cellular stress⁴³. Another wide application area for activatable dyes is sensing of pH.

1.3. pH-sensing with fluorescence dyes

Although the detection of changes in pH is well established on tumor cells *in vitro*⁴⁶, pH sensing *in vivo* remains challenging. A change in pH is a characteristic of numerous diseases like inflammation or cancer. So far only few NIR-emitting pH-sensitive probes exist, but most of the pH probes consist of fluorophores with spectra in the visible region^{1,38}. pH-activatable NIRF probes can be divided in two groups. The first group comprises NIRF dyes coupled to target specific proteins in either a low dye to protein (DP) ratio, resulting in autoquenching⁷ or in a high DP ratio resulting in self-quenching by homo FRET⁸. Such probes are activated by enzymatic cleavage of fluorophores, e.g. by degradation in the lysosome, allowing spatial separation from each other⁴⁷. The second group consists of norcyanine- or norsquarine-based NIR pH probes which have a protonatable amine group within the fluorophore core. Examples are modified ICG, H-ICG^{43,48} or Square-650-pH⁴⁹, and also CypHer5E, applied in this study. Accordingly, such dyes are fluorescent when protonated in an acidic

environment but fluorescence is greatly reduced when non-protonated. Here, initiation of fluorescence is reversible, in contrast to activation via enzymatic cleavage of protein-fluorophore conjugates.

pH-responsive protonatable dyes may improve the sensitivity of detection and monitoring of acidic tumors *in vivo*. Following *intravenous* (*i.v.*) injection, always-on fluorescent probes systematically distribute in the entire object causing high background signals. Moreover, the tumor is undetectable until the fluorophore has concentrated in the tumor and the unbound probe is cleared from the object (Figure 2a). In contrast, pH-activatable dyes are non-fluorescent after application, thereby causing no background fluorescence of unbound probe and only become fluorescent in the acidic tumor environment, resulting in a high tumor contrast even before the unbound (non-fluorescent) probe is cleared from the object (Figure 2b).



Figure 2. Improvement of tumordetection sensitivity by the use of pHactivatable probes

a. Always-on fluorescence probes cause high background signals from the unbound probe when applied in vivo. After a certain time, following the probe's accumulation at the tumor, fluorescence intensity at the tumor site increases while background fluorescence decreases due to systemic clearance of the probe. Due to a certain amount of unbound fluorescent probe, the tumor to background ratio is low, especially at early time points after probe application. **b.** pH-activatable probes may avoid background fluorescence after application in vivo and only activate at the tumor site. This would result in an early detection of the fluorescent tumor even before the circulating probe is cleared from the organism, leading to a high tumor to background ratio. c. A mechanism of activation of pH-sensitive probes at the tumor site is internalization of tumorspecific ligands via receptor-mediated

endocytosis, exemplified here for Trastuzumab (Herceptin). Here, the conjugated pH-sensitive fluorophores are activated in the acidic pH of the late endosome and lysosome. Figure was taken from Urano et al.³⁸.

Activation of such pH-sensitive probes may also occur after internalization in the targeted tumor cell. This can be achieved, for example, by conjugation of the fluorophore to a tumor-specific antibody that binds to a receptor on the targeted tumor cell, followed by internalization of the antibody-conjugate receptor complex and localization in the endosomal and lysosomal compartments. There, the low pH of 5 to 6 leads to activation of the dye, resulting in fluorescence (Figure 2c)³⁸.

1.3.1. CypHer5E for sensitive tumor imaging

A promising candidate for *in vivo* imaging of tumors is the pH-activatable NIRF dye CypHer5 (1-(5-Carboxypentyl)-2-[1E,3E)-5-(3,3-Dimethyl-5-Sulfo-1,3-Dihydro-2H-Indol-2-Vlidene)-1,3-Pentadienyl]-3,3-Dimethyl-3H

Indolium-5-Sulfonate(V)). CypHer5 is a pentamethine cyanine (CyTM 5) dye derivative that increases its fluorescence intensity with decreasing pH. In addition, CypHer5 has a reactive functional group (hydroxyl succinimidyl ester) allowing covalent linking of proteins, like antibodies, via aliphatic amino groups. Coupling of CypHer5 to an antibody and thereby following internalization of G-protein-coupled receptor from the cell membrane into the endolysosomes was the first report of an application of a NIR pH probe^{39,43}.

CypHer5 has been further developed to CypHer5E enhanced pH-sensitive dye (4-{2-[(1E,3E,5E)-5-(3-{6-[(2,5-dioxopyrrolidin-1-yl)oxy]-6-oxohexylis e-3dihydro-2H-indol-2-ylidene)penta-1,3-dienyl]-3,3methyl-5-sulfo-1,3 dimethyl-5-sulfo-3H-indolium-1yl}butane-1-sulfonate) which is also commercially available. In comparison to CypHer5, CypHer5E comprises a better extinction coefficient resulting in increased excitability as well as an improved solubility by additional sulfonate groups. The structure of CypHer5E is shown in Figure 3. This pH-sensitive dye is capable of fluorescence emission when protonated at the nitrogen atom in an acidic environment but non-fluorescent in a basic to neutral environment when the amine group in not protonated. This deprotonation leads to a shift in the absorption maximum of the dye at 645 nm and subsequently to a decrease in fluorescence (Figure 4). As CypHer5E has a pKa in vitro of 7.3, it is perfectly suitable for pH sensing in a biological environment. CypHer5E has already been shown to be functional in cell assays for studying phagocytosis⁵⁰, for quantification of internalization of cellular target molecules^{51,52}, and for monitoring of synaptic vesicle recycling⁵³. Although CypHer5E, having NIR spectra, a good water solubility, and functional groups for bioconjugation, is a promising candidate for in vivo imaging of structures with a low pH, like a tumor, so far no reports have been published about the use of this dye in *in vivo* imaging.



Figure 3. Chemical structure of CypHer5E

In a basic to neutral solution, where the nitrogen atom of CypHer5E is not protonated, the dye is not fluorescent. When protonated in an acidic environment, the chromophore system of a classic cyanine dye is observed for CypHer5E, resulting in fluorescence. Picture was taken from GE website⁵⁴.



Figure 4. Absorption characteristics of CypHer5E at different pH

CypHer5E has two absorption bands that peak in the range of 490 and 645 nm. Photon absorption around 645 nm results in fluorescence emission with a peak intensity at 663 nm. The absorption band in the shorter wavelength region (~490 nm; not leading to fluorescence emission) increases with increasing pH of the PBS measurement solution, whereas the absorption band at longer wavelength (~645 nm) decreases resulting in decreased fluorescence. Image was provided by Jutta Pauli, BAM I.5, Berlin.

1.4. Nanoparticles as fluorescence probes

Even though organic fluorophores compose the major group of fluorescence probes for *in vivo* applications they often suffer from limited sensitivity, a low fluorescence quantum yield, and low photostability. Moreover, organic dyes may form quenched aggregates and are sensitive to environmental changes in tissue or blood, influencing their spectroscopic properties. Furthermore, many of them possess a certain level of toxicity that hampers their application *in vivo*²⁰.

A promising possibility to generate fluorescent probes for *in vivo* applications is to enclose fluorophores in nanomaterials, such as polymeric nanoparticles (NPs), which can be generated from a number of materials in diverse compositions and labeled with various fluorophores. This has the benefit of improved stability of the fluorophores due to reduced interactions with the environment, e.g. with quenching agents, solvent molecules, or chemically reactive species⁵⁵. In reverse, by enclosement in NPs, the environment can be protected against potentially toxic fluorophores. In most cases, enclosure in NPs does not alter the dye spectral characteristics, but it minimizes dye aggregation and subsequent dye quenching, and offers an improved photostability. As NPs are capable of carrying large numbers of fluorophores, very bright fluorescence probes can be produced, compared to a single dye molecule. NPs can further be applied for simultaneous imaging by different modalities, e.g. by generating NPs that are loaded with contrast agents for MRI and surface labeled with fluorescence dyes^{56,57}.

The core of NPs may be surrounded by a shell layer that insulates and protects the core and mediates solubility of NPs and/or linkage to bioligands⁵⁸. NPs with many different configurations have been designed which, due to their large surface area can be easily modified with polymeric compounds such as polyethylene glycol (PEG) or various target-specific biomolecules, like peptides or antibodies, thereby creating highly fluorescent target-specific 3D probes for *in vivo* imaging⁵⁹ (see Figure 5). Especially, PEGylation has a particular relevance as it can modify the pharmacokinetics of NPs by reducing their clearance through the mononuclear phagocytic system (MPS) and thereby not only increasing plasma half-lives of the NPs but also their stability and water solubility⁶⁰⁻⁶².

NPs of diverse compositions have been developed for imaging purposes. Most particles can be classified into two groups: NPs that are mostly composed of organic molecules, like dendrimers^{63,64}, liposomes^{64,65}, poly(lactic-*co*-glycolic acid) (PLGA)⁶⁶, polystyrene (PS)^{67,68}, or chitosan^{59,69}. And those that mainly consist of inorganic materials, such as silica⁷⁰, gold⁷¹, and semiconductor nanocristals, the so called

10



quantum dots (QDs) that comprise a core/shell structure, such as CdSe/ZnS, CdS/ZnS, or CdTe/ZnS⁷².

Figure 5. Compositions and surfacemodifications of nanoparticles

A. A scheme of NP components and possible configurations is shown. NPs can be composed of (i) a core directly surrounded by targeting biomolecules or (ii) surface ligands can be used to anchor the target molecules to the particle core. (iii) Biomolecules can also interact with a shell layer that surrounds the NP core or (iv) be anchored to the core/shell particle via intermediate ligands. On the other hand, (v) the targeting biomolecules can be located inside a porous NP core or (vi) be entrapped by a porous NP core surrounded by a shell layer. (vii) Furthermore, the NP core can be much smaller than the biomolecule and in addition (viii) be attached to the core by an intermediate ligand. **B.** A representative fluorophore-loaded NP, coated with a PEG layer is illustrated. Multiple target-specific molecules can be attached to the PEG chains in three dimensions. Specific targeting and detection of biomolecules by NPs may be achieved for instance via surface-conjugation with nucleic acids, target-specific small peptides, proteins, or antibodies. Figure was taken from Sapsford et al.⁵⁸.

The composition of the NPs should be selected according to the application purpose of the particle. NPs formulated from biodegradable substances, like liposomes are especially favorable for drug delivery and are already used as chemotherapeutic vehicles in different human tumors⁶⁴. Such drug vehicles are advantageous as they especially enable the delivery of hydrophobic agents, like the anti-cancer drugs paclitaxel and docetaxel that are not soluble in aqueous environments⁶⁴. In addition, other biodegradable NPs, composed of chitosan, gelatin, or PLGA are being investigated as non-viral gene-delivery systems⁷³. On the other hand, the use of non-biodegradable NPs, like PS, may be advantageous for imaging purposes as enclosed fluorophores are prevented from release into the surrounding environment, where they could potentially cause unknown toxic effects. Instead they are excreted from the body in the same composition as they were administered. Moreover, in the case of hydrophobic imaging agents, the incorporation in NPs enables these agents to be used in aqueous environments and consequently *in vivo*⁶⁸.

Despite the growing effort in the development of potent NPs for diagnosis and therapy of diseases, the biodistribution, behavior and risks of NPs in living organisms have to be examined further⁷⁴. In particular, most NP formulations, such as QDs²⁰, still suffer from potentially toxic effects or from a too rapid clearance from the body due to a low transport of relatively large particles across the endothelium and/or a rapid elimination of NPs by the MPS⁷⁵.

1.4.1. Polystyrene nanoparticles as potent imaging tools

Polystyrene nanoparticles (PSNPs; Figure 6) are especially advantageous for *in vivo* imaging purposes, as they are commercially available in different sizes from 15 nm to several micrometers. Moreover, in general PSNPs are regarded to have no impact on the metabolic activity of cells and low levels of cytotoxicity. Nevertheless, as is the case for other NP formulations, this may depend on the used concentrations of NPs and incubation times, as well as on particle size, and surface charge⁷⁶⁻⁷⁹. A further advantage is that the loading of PSNPs with inorganic and organic fluorophores is well established, thereby enabling a simple and homogeneous production of bright fluorescent reporters for imaging applications⁶⁸. As PSNPs are available with various surface chemistries, this further enables a simple and efficient modification of particles with target-specific biomolecules.



Figure 6. Transmission electron microscopy image of 100 nm PSNPs

The morphology of 100 nm PSNPs is illustrated. Scale bar represents 0.2 μ m. Image was taken from Liu et al.⁷⁶.

1.4.2. Highly fluorescent nanoparticles loaded with Itrybe

A very promising dye for imaging applications in combination with NPs is the broadband fluorophore Itrybe. This hydrophobic asymmetric cyanine dye has a broad absorption (450 to 750 nm) and emission spectrum (650 to 900 nm) in the NIR region making it especially favorable for *in vivo* imaging. Furthermore, compared to other NIR fluorophores, Itrybe has a large Strokes shift which is the difference between the positions of the band maxima of the absorption and emission spectra. This enables a good separation of excitation and emission light, using appropriate filter combinations. Due to the broad spectra of Itrybe it is also possible to achieve a high compatibility of the dye with many different imaging devices equipped with various laser and/or filter settings. In contrast, most other NIR dyes are characterized by a narrow absorption- and emission spectrum and a considerably smaller Strokes shift as well as highly overlapping absorption- and emission spectra.

In contrast to other NIR fluorophores, Itrybe, does also not form quenched aggregates at high concentrations, which otherwise could result in decreased fluorescence. This allows the loading of high amounts of Itrybe molecules into solid matrices, like PS, leading to a significantly higher brightness of these probes. A further advantage of this dye as a potent imaging tool in combination with NPs, is its outstanding thermal and photochemical stability (Thomas Behnke, BAM I.5, Berlin, personal communication).

On these grounds, a patent application has been submitted by the BAM in Berlin together with the Max-Planck-Institute for Experimental Medicine and the University Medicine in Göttingen on the application of Itrybe in combination with NPs for imaging purposes (see list of publications).

1.5. Specific targeting of tumors

NIRF imaging offers an inexpensive and safe alternative to imaging methods like PET and SPECT which use potentially harmful, highly energetic radiation. Therefore, NIRF imaging provides an attractive tool for preclinical studies of tumor development and progression and/or evaluation of cancer therapies in animal models. As cancer is the leading cause of death worldwide⁸⁰ there is a continuous need for the development of novel improved probes for an early detection and characterization of tumors as well as for a better understanding of cancer onset and progression. Cancer development is a multistep process during which cells acquire several capabilities, enabling them to become tumorigenic and finally malignant. One emerging hallmark of potential generality of cancer is reprogramming of energy metabolism which is needed to promote tumor cell growth and division⁸¹. The tumor microenvironment is known to be hypoxic, resulting from an inefficient perfusion due to a chaotic tumor vasculature. Under anaerobic conditions, cells favor glycolysis instead of energy production in the oxygen consuming mitochondria⁸¹. Otto Warburg first observed an abnormal behavior of energy metabolism in cancer cells, in that even in the presence of oxygen cancer cells can reprogram their energy metabolism to glycolysis^{82,83}. The upregulation of glycolysis results in an increased acid production and subsequently in microenvironmental acidosis⁸⁴. It is further known that the extracellular pH decreases with increasing tumor size and that gradients of interstitial pH exist in tumors. Moreover, the acidity can vary among different tumor types. For instance, the extracellular pH investigated in two breast cancer cell lines was determined to be 6.8 in MDA-MB-435 cells and 7.1 in MCF-7 cells when grown in SCID mice to a comparable size⁸⁵.

The characteristic of an acidic environment, that most tumors possess, can be effectively used to increase tumor contrast by using pH-sensitive fluorescence probes that are selectively activated in the acidic tumor environment.

1.5.1. The role of HER2 in cancer

Imaging probes, such as free fluorophores or dye-loaded NPs, which are systematically applied in order to detect tumors, have to be functionalized to specifically bind to tumor associated antigens on cancer cells. Easily accessible target structures might be for example membrane-associated proteins, like receptors⁸⁶, ion-channels⁸⁷, or cell-adhesion molecules, such as integrins².

In the case of breast cancer detection and monitoring, a well known biomarker is the human epithelial growth factor receptor 2 (HER2, also known as c-neu, Neu or ErbB-2). HER2 is highly expressed on 25 to 30 % of all breast cancers⁸⁸ but also in 10 to 15 % of ovarian cancers^{89,90} and in 5 to 15 % of other tumor types like gastric, cervix, colon, oral, endometrial, esophageal, lung, and pancreatic cancers⁹¹. Therefore, HER2 has become not only an interesting target structure for therapy, but due to its localization at the cell membrane, making it easily accessible by exogenously applied ligands, also an attractive marker for the diagnosis and monitoring of cancer.

HER2 is a member of the EGFR superfamily, also known as the ErbB protein family that form dimers upon ligand stimulation leading to activation of downstream signaling cascades associated with cell growth, proliferation, survival, cell motility, and angiogenesis^{91,92}. HER2 is a 185 kDa transmembrane glycoprotein consisting of an extracellular binding domain, a transmembrane segment, and a tyrosine kinase domain with a regulatory carboxyl terminal segment^{91,93,94}. In addition, a truncated
membrane-bound fragment of the HER2 protein, p95 exists that can be detected in breast tumors. p95 is generated after proteolytic cleavage of the extracellular domain of HER2 and is associated with an increased tyrosine kinase activity and thus an enhanced intracellular signaling^{95,96}. An overexpression or amplification of HER2 results in increased intracellular cell signaling, leading to a deregulation of cell proliferation and survival, which in turn may cause uncontrolled cell proliferation and subsequently malignancy⁹¹. The prolonged duration of intracellular signaling in HER2 overexpressing cells is caused by a slowdown of the dissociation of ligand-receptor complexes which results in slower internalization of the complexes through clathrin-coated pits and thereby slower degradation of the active receptors in lysosomes⁹².

1.5.2. Targeting of HER2 by Herceptin

Owing to its expression on the cell membrane of tumor cells, HER2 is an optimal target not only for therapy but also for tumor recognition. In this context a humanized monoclonal antibody, trastuzumab (Herceptin®, in the following referred to as Herceptin) was developed from the murine antibody 4D5. Herceptin was approved by the FDA in 1998 for treatment of patients with metastatic breast cancer overexpressing HER2⁹¹. In heavily pretreated patients, Herceptin has a response rate of 10 to 15 % and in untreated patients of 25 %⁹¹.

Herceptin is known to bind to the extracellular domain IV of HER2⁸⁵ and to induce formation of HER2 homodimers, leading to ligand-mediated endocytosis of the receptor and to a considerable removal of HER2 from the plasma membrane and subsequently to a decreased receptor-initiated constitutive signaling⁹⁷. However, the mechanism by which Herceptin causes its antitumor effect in cancer cells is not entirely understood. Other mechanisms of Herceptin action may include suppression of angiogenesis⁹⁸, sterical blocking of the proteolytic cleavage of the extracellular region of HER2⁹⁹, and disruption of interaction of HER2 with other proteins⁹⁹. Another hypothesis favors an immunological mechanism, where antibody dependent cellular cytotoxicity (ADCC) caused by Herceptin bound to the tumor cell results in tumor cell lysis⁹⁸.

As Herceptin is a very well evaluated and clinically proven antibody, it presents an attractive device for the detection and monitoring of breast tumors¹⁰⁰. Therefore, Herceptin is an optimal ligand for the bioconjugation of novel fluorescence probes, like pH-activatable dyes and dye-loaded NPs, to be evaluated as novel probes for *in vivo* imaging of breast tumors.

For the evaluation of tumor-detection sensitivity of fluorescence probes tumor models are of considerable importance. Human KPL-4 breast tumor cells were

selected as the orthotopic breast tumor model in this study as these cells are known to have an overamplification of the HER2 gene as well as a high HER2 protein expression¹⁰¹. Orthotopic MDA-MB-231 cell implantation was performed to obtain control tumors in nude mice that do not overexpress HER2¹⁰². Orthotopic breast tumor xenorafts in mice generated by KPL-4¹⁰³ as well as by MDA-MB-231¹⁰² cells have already been shown to be suitable model organisms for the evaluation of Herceptin binding *in vivo*. Therefore, both tumor models were used in this study to investigate *in vivo* binding and biodistribution of Herceptin-conjuagted fluorescence probes.

AIMS OF THE STUDY

The aim of the work presented here was to characterize and to evaluate novel and innovative fluorescent probes regarding their suitability for fluorescence imaging *in vivo*, in order to improve tumor-detection sensitivity in breast tumor-bearing mice.

For this purpose, 1.) a pH-sensitive NIR dye, CypHer5E and 2.) PSNPs loaded with a broadband NIR fluorophore, Itrybe, were investigated in orthotopic breast-tumor xenografts in nude mice. To achieve tumor targeting capacity, both probes were conjugated to Herceptin, in order to take advantage of the knowledge already available for the kinetics of the tumor targeting moiety Herceptin and thereby allowing me to focus on the behavior of the fluorescent labels. This is the first report of an *in vivo* imaging application and evaluation of CypHer5E as well as for Itrybe-loaded PSNPs.

The following issues will be addressed:

- 1.) The suitability of pH-activatable CypHer5E-Herceptin-conjugates to improve tumor-detection sensitivity *in vivo* in comparison to always-on fluorescence conjugates, AlexaFluor 647-Herceptin
 - a. The *in vitro* characterization of fluorescence probes upon the influence of different environmental factors
 - The functionality of pH-sensitive conjugates *in vitro* in cell based assays
 - c. The capability of CypHer5E-Herceptin probes to improve tumor contrast *in vivo* in KPL-4 and MDA-MB-231 breast tumor-bearing mice in combination with two different techniques for suppression of background fluorescence: subtraction of autofluorescence and LT-gated imaging
- 2.) The biodistribution of Itrybe-loaded PSNPs of distinct sizes (100, 25, and 15 nm) and with different surface modifications (amino groups, PEG chains of different sizes, or Herceptin coupled to PEG chains) and their feasibility for application in *in vivo* imaging of breast tumors in mice
 - The spectroscopic characterization of Itrybe-loaded PSNPs and the influence of environmental factors on fluorescence of the probes
 - The binding of Herceptin-modified NPs to HER2 in immuno- and cellbased assays
 - c. The influences of NPs on cell viability

- d. The biodistribution of probes *in vivo* and *ex vivo*
- e. The tumor-targeting capacity of PSNPs in KPL-4 and MDA-MB-231 breast-tumor bearing mice *in vivo* and *ex vivo*

The results of this study will contribute to an improved knowledge about the characteristics of pH-sensitive probes and Itybe-loaded NPs and their suitability for application in non-invasive detection of breast tumors.

2. MATERIAL AND METHODS

2.1. Material

2.1.1. Equipment

Analytical balance, Competence CP64 Anesthesia system

Apparatus for tissue processing, TP 1020 Camera, EOS 1000D Centrifuge 5810 Chemi-Doc luminescence detection system Centrifuge 5424 Cryostat, Jung Frigocut 2800E Extrudate 1439 Fluorescence microscope, Axiovert 200M Incubator, Heraeus BB6220 LightCycler 480 Light microscope, Axioscope 2 Light microscope, Telaval 31 Microtome HM 340E Nano photometer, Pearl Neubauer counting chamber Odyssey infrared imaging system Optix MX2 ORCA-ER digital camera

pH-meter, SevenEasy Quartz glass precision cuvette Scantainer Spectrofluorometer, QuantaMaster

Staining cuvette Steamer, MultiGourmet Thermomixer compact Tissue lyzer Ultrasonic cleaner Sartorius, Göttingen VisualSonics, Toronto, Ontario, Canada Leica, Bensheim Canon, Krefeld Eppendorf, Hamburg Bio-Rad, Munich Eppendorf, Hamburg Leica, Bensheim Altromin, Lage Carl Zeiss, Göttingen Kendro, Laboratory products, Hanau Roche, Mannheim Carl Zeiss, Göttingen Carl Zeiss, Göttingen Thermo Scientific, Braunschweig Implen, Munich Schütt Labortechnik, Göttingen Li-Cor Biosciences, Bad Homburg ART, Montreal, Canada Hamamatsu, Herrsching am Ammersee Mettler-Toledo, Giessen Helma, Müllheim Scanbur, Koge, Denmark Photon Technology International, Seefeld VWR International, Ismaning Braun, Kronberg Eppendorf, Hamburg Qiagen, Hilden Schütt Labortechnik, Göttingen

Vortexer, Vortex Genie 2 Wallac 1420 Victor 2 multilabel counter 2.1.2. **Consumable supplies** Cell culture flasks Chambered coverglasses, 4 well Cover glasses 12 mm Ø Cryotube vials Culture dishes Culture plates 24-wells, 96-wells Falcontubes 15 ml, 50 ml Filter tips, 10 µl, 100 µl, 1000 µl Insulin syringes with integrated needle $(30G \times \frac{1}{2})$ Insulin syringes with integrated needle (30G) Insulin syringes, Omnifix 40 solo Microscope slides Needles, Sterican, 26G x 1/2 Hybridization transfer membranes Nunc-Immuno modules, maxisorp loose NuPAGE Novex 3-8 % TRIS-acetate gels Pasteurpipettes Permanox cell culture slides Serological pipettes 5 ml, 10 ml Serological pipettes 25 ml Surgical suture materials, absorbable, Vicryl, 4/0 Tissue Tek cassettes

2.1.3. Chemicals and reagents

ABTS single solution

Bender & Hobein,Zurich, Switzerland American Instrument Exchange, Haverhill, MA, USA

Sarstedt, Nümbrecht Nalge Nunc International, Rochester, NY, USA Thermo Scientific, Bonn Nunc, Roskilde, Denmark Sarstedt, Nümbrecht Greiner Bio-One, Solingen BD, Heidelberg Sarstedt, Nümbrecht Braun, Melsungen

BD, Heidelberg

Braun, Melsungen Paul Marienfeld, Lauda-Königshofen Braun, Melsungen Amersham/GE Healthcare, Munich Nunc A/S, Roskilde, Denmark Invitrogen, Darmstadt Sarstedt, Nümbrecht Nalge Nunc International, Naper Ville, IL, USA Greiner Bio-One, Solingen Corning Incorporated, Corning, NY, USA Johnson & Johnson, Neuss

Vogel, Giessen

Invitrogen, Darmstadt

AEC peroxidase substrate Antibody diluent Aquatex, aqueous mounting medium Blocking buffer - fish BSA Casein Chemiluminescent HRP substrate Chloroform Coomassie R250 DEPC DePex, hydrophobic mounting medium DMEM DPBS EDTA Eosin G Ethanol FCS Formaldehyde GLP Nafaq feed 890 Goat serum Haematoxylin HEPES Hoechst 33342 trihydrochloride, trihydrate Hydrochloric acid, 37% Hydrogen peroxid Fluorescent mounting medium containing DAPI Isoflurane Ketamine 10% LDS sample buffer NuPAGE Antioxidant Paraffin (TM 60 °C) **PBS** tablets Poly-L-lysine Precision Plus protein standard

Vector, Peterborough, United Kingdom Dako, Hamburg Merck, Darmstadt SurModics, Eden Prairie, MN, USA Sigma-Aldrich, Seelze Roche, Mannheim Millipore, Billerica, MA, USA Merck, Darmstadt Biomol, Hamburg Sigma-Aldrich, Seelze Serva, Heidelberg Gibco, Invitrogen, Darmstadt Gibco, Invitrogen, Darmstadt Sigma-Aldrich, Seelze Merck, Darmstadt Merck, Darmstadt PAN Systems, Aidenbach Merck, Darmstadt Provimi Kliba AG, Kaiseraugst Gibco, Invitrogen, Darmstadt Merck, Darmstadt Carl Roth, Karlsruhe Invitrogen, Darmstadt Merck, Darmstadt Sigma-Aldrich, Seelze Merck, Darmstadt

Baxter, Unterschleißheim Medistar, Holzwickede Invitrogen, Darmstadt Invitrogen, Darmstadt Süsse, Guxhagen Gibco, Invitrogen, Darmstadt Sigma-Aldrich, Seelze BIO-RAD Laboratories, Munich ProLong gold antifade reagent Invitrogen, Darmstadt with DAPI Protease inhibitor cocktail Roche, Mannheim QIAzol lysis reagent Qiagen, Hilden Reducing agent Invitrogen, Darmstadt Regime 210, U8959A01R Scientific Animal Food and Engineering, Augy, France Sodium chloride Merck, Darmstadt Sodium chloride, isotone, 0.9% Braun, Melsungen Sodium hydroxide Merck, Darmstadt Target retrieval solution Dako, Hamburg TRIS Sigma-Aldrich, Seelze TRIS-acetate SDS running buffer Invitrogen, Darmstadt Triton X-100 Sigma-Aldrich, Seelze Tween-20 Sigma-Aldrich, Seelze Western blot stripping buffer Thermo Scientific, Bonn WST-1 cell proliferation reagent Clontech, Mountain View, CA, USA Xylazine 2% Riemser, Greifswald Xylol Merck, Darmstadt

2.1.4. Kits

BCA protein assay kitThermo Scientific, BonnMycoplasma PlusTM PCR primer setAgilent Technologies, WaldbronnReady-to-use hot-start PCR master mixRoche, MannheimRNase-free DNase setQiagen, HildenRNeasy mini kitQiagen, HildenSuperScript first-strand synthesis systemInvitrogen, Darmstadtfor RT-PCRKenter Scientific, Bonn

2.1.5. Probes

substance	application	provider
Itrybe-loaded NPs	fluorescence imaging	BAM I.5, Berlin
CypHer5E	fluorescence imaging	GE Healthcare, Munich
Alexa Fluor 647	fluorescence imaging	Invitrogen, Darmstadt
humanized anti-HER2 IgG, Herceptin (Trastuzumab)	tumor targeting	Roche Diagnostics, Mannheim

substance	application	provider
rabbit anti-HER2 IgG, Neu (C-18): sc-284	Western blot	Santa Cruz, Heidelberg
goat anti-actin IgG, (C-11): sc-1615	Western blot	Santa Cruz, Heidelberg
donkey anti-rabbit IgG, HRP-labeled: NA934	Western blot	GE Healthcare, Munich
rabbit anti-goat IgG, HRP- labeled: 172-1034	Western blot	BIO-RAD Laboratories, Munich
HER2 extracellular domain (ECD) protein	immunoassay	Roche Diagnostics, Mannheim
rat anti-mouse CD31 IgG: DIA 310 M	immunohistology	Dianova, Hamburg
goat anti-rat IgG, HRP labeled: AP136P	immunohistology	Merck, Millipore, Schwalbach

2.1.6. Enzymes

Trypsine-EDTA (0.25 %) DNase I RNase H Super Script II RT Uracil-DNA-glycosylase PAA Laboratories, Pasching, Austria Qiagen, Hilden Invitrogen, Darmstadt Invitrogen, Darmstadt New England BioLabs, Frankfurt am Main

2.1.7. Eukaryotic cell lines

• KPL-4, provided by J. Kurebayashi, Department of Breast and Thyroid Surgery, Kawasaki Medical School, Kurashiki, Japan

Human breast cancer cell line, isolated from the malignant pleural effusion of a breast cancer patient with an inflammatory skin metastasis¹⁰¹.

• MDA-MB-231, American type culture collection (ATCC)/LGC, Wesel

Human breast cancer cell line, adenocarcinoma from the mammary gland derived from a metastatic pleural effusion.

 3T3, Swiss albino mouse fibroblasts, Deutsche Sammlung von Mikroorganismen und Zellkulturen (DMSZ), Braunschweig

Fibroblasts, established from disaggregated Swiss albino mouse embryos.

2.1.8. Animals

Female athymic nude mice, strain NMRI-Fox1nu/nu, Central Institution of Animal Experiments of the University Medicine, Göttingen.

2.1.9. Application specific computer programs

Program:	Used for:	
Application Software, Version 2.1	Odyssey infrared imaging system	
AxioVision Rel.4.6	Axiovert 200M fluorescence	
	microscope	
FeliX32 Analysis 1.0	Spectroflourometer QuantaMaster	
Image 1.42I ¹⁰⁴	Image processing	
LightCycler 480 Software 1.5	LightCycler 480	
OptiView 1.00.00, ART, Montreal, Canada	Optix MX2 in vivo imaging system	
Quantity One 4.6.2	Chemi-doc luminescence detection	
	system	
Wallac 1420 Manager, Version 2.0	Victor 2 multilabel counter	

2.2. Methods

2.2.1. Molecular Biology

2.2.1.1. Total RNA isolation and cDNA synthesis

Purification of RNA from cells:

RNA from cells was isolated via use of RNeasy mini kit according to the instruction manual. Briefly, cells ($\sim 1 \times 10^6$) were disrupted in buffer containing guanidine isothiocyanate and homogenized. Ethanol was then added to the lysate to bind RNA to the RNeasy silica-gel membrane and contaminants were washed away. DNase treatment was performed to eliminate genomic DNA (according to the instruction manual). After wash steps, the RNA was eluted in 30 µl RNase-free diethylpyrocarbonat (DEPC) water.

Purification of RNA from tumor tissue:

RNA from tumor tissue was isolated via phenol-chloroform extraction in combination with use of RNeasy mini kit. Tumor tissue (< 100 mg) was homogenized with 1 ml QIA-zol lysis reagent containing phenol and guanidine thiocyanate using a tissue lyzer. 200 μ l chloroform was added and samples were vortexed and incubated for 5 min at room temperature (RT). After 15 min of centrifugation at 12,000 x g at 4 °C the aqueous phase was transferred in a new cup, supplemented with 600 μ l 70 % ethanol and vortexed. The suspension was transferred to a RNeasy mini spin column and centrifuged at maximal speed for 15 sec. Total RNA bound to the membrane and contaminants were washed away. DNase treatment and elution of RNA was performed as described above.

cDNA synthesis:

cDNA synthesis for reverse transcriptase-polymerase chain reaction (RT-PCR) was performed with the SuperScript first-strand synthesis system. cDNA was synthesized from 2.5 μ g of total RNA upon addition of 0.5 μ g oligo (dT) and DEPC-water to advice a RNA concentration of 0.2 μ g/ μ l in a volume of 12 μ l. At each step the samples were mixed by vortexing and collected by brief centrifugations. The reaction mixture was heated at 70 °C for 10 min and rapidly cooled on ice for 1 min. Subsequently reaction buffer was added containing: 2 μ l of reverse transcriptase buffer, 2 μ l MgCl2 25 mM, 2 μ l of dithiothreitol (DTT) 0.1 M and 1 μ l of dNTPs 10 mM. The reaction mixture was heated for 5 min at 42 °C. The negative samples were supplemented with 1 μ l of DEPC-water, whereas to the positive samples 200 units of reverse transcriptase enzyme were added. Probes were incubated for 50 min at 42 °C

following inactivation of the reverse transcriptase at 70 °C for 15 min. To degrade RNA present in the samples 2 units of RNase H were incubated with the mixture for 20 min at 37 °C. At the end of the procedure the mixture was diluted with DEPC-water to advice a reaction volume of 25 μ l with a final cDNA concentration of 0.1 μ g/ μ l.

RNA and cDNA concentration and purity were determined by optical density measurements at 260 and 280 nm using the spectrofluorometer.

2.2.1.2. Quantitative real-time PCR

Real-time PCR was performed using the TaqMan system in the LightCycler 480. Primers and probes were chosen from the Universal Probe Library from Roche¹⁰⁵. The TaqMan probes contain a reporter dye fluorescein amidite (FAM) at the 5 end of the probe and a Dark Quencher Dye at the 3 end of the probe. When cleaved by the Taq DNA polymerase during PCR the reporter and quencher dye of the probe are separated, resulting in fluorescence of the reporter. For analyzing the mRNA expression of the human epidermal growth factor receptor 2 (*hHER2*) and human transferrin receptor type 1 (*hTfR*), used to control for RNA integrity and quantification, the following primers and probes were used:

<u>Primer</u>	Sequence
hTfR forward	5'-TTG AGA AAA CAA TGC AAA ATG TG-3'
hTfR reverse	5'-CCC AGT TGC TGT CCT GAT ATA GA-3'
TaqMan probe # 61	5'-FAM CTG GGC AA Dark Quencher Dye-3'
hHER2 forward	5`-CAA GTA ATC CGG GGA CGA A-3`
hHER2 reverse	5'-CAC TGC CCA GTT CCC TCA-3'
TaqMan probe # 23	5'-FAM GGG CTG GG Dark Quencher Dye-3'

Real-time PCR was performed with 100 ng cDNA, 200 nM forward and reverse primer, 100 nM TaqMan probe, 0.2 U Uracil-DNA-glycosylase, and a ready-to-use hot-start PCR mix containing Taq DNA polymerase, dNTP mix, as well as buffer and MgCl₂.

PCR conditions were:

2 min 50 °C; 10 min 95 °C; (10 sec 95 °C; 30 sec 60 °C; and 10 sec 40 °C) x 45 cycles. The number of PCR cycles to reach cycle threshold (Ct) was used to determine the relative mRNA expression. The results were standardized to the amount of *hTfR* in the respective probe and to the amount of mRNA expression in the calibrator (human brain RNA).

For this purpose the $\Delta\Delta$ Ct-Method was applied:

Normalized ratio: $2^{\Delta Ct1} / 2^{\Delta Ct2}$

 $\Delta Ct1 = Ct hHER2$ (calibrator) - Ct hHER2 (sample)

 $\Delta Ct2 = Ct hTfR$ (calibrator) - Ct hTfR (sample)

2.2.2. Protein extraction, protein separation on protein gel, and Western blot

2.2.2.1. Protein extraction

To obtain cell lysates, cell cultures were detached from the dish as described in 2.2.5.1, transferred to a 15 ml falcon tube, washed twice with PBS, and resuspended in 3 ml of lysis buffer. After 30 min of incubation at RT, cell debris were centrifuged for 15 min at 18,000 g at 4 °C and the supernatant was used as total cell extract.

 Lysis buffer: 50 mM trishydroxymethylaminomethane (TRIS)-HCl, pH 7.4 300 mM NaCl
 5 mM ethylene-diamine-tetraacetic acid (EDTA) 1% Triton X-100 Protease inhibitor cocktail (1 tablet for 25 ml)

2.2.2.2. BCA protein assay

Protein concentration was determined using the bicinchoninic acid (BCA) protein assay kit that employs bovine serum-albumin (BSA) as a standard curve. The samples (triplicates) in a volume of 25 μ l were incubated for 30 min at 37 °C together with the BCA reagent, composed by solution A (sodium carbonate, sodium bicarbonate, bicinchoninic acid and sodium tartrate in 0.1 M sodium hydroxide) and solution B (4 % cupric sulfate). Absorbance was measured in the Wallac 1420 Victor 2 multilabel counter at 550 nm.

2.2.2.3. SDS-PAGE

50 µg of total protein extract per sample were separated on a gradient SDS-PAGE (sodium dodecyl sulfate-polyacrylamide gel electrophoresis) NuPAGE Novex TRISacetate 3-8 % gel, according to the instruction manual. Depending on the application the SDS-gels were either used for western blot analysis or stained with Coomassie for protein detection.

2.2.2.4. Western blot analysis

For Western blot analysis the proteins were transferred to nitrocellulose hybridization transfer membranes. Membranes were blocked for 1 h with 0.1 % casein in TRIS buffer containing Tween-20 (TBST) buffer, incubated with rabbit anti-HER2 antibody (1:200 dilution) in 0.1 % casein for 1 h, washed with deionized water for 7 times and then incubated in TBST for 5 min. Following, the membranes were incubated with horseradish peroxidase (HRP)-coupled donkey anti-rabbit antibody (1:8000 dilution) for 1 h. After washing, the bound antibody was visualized using chemiluminescent HRP substrate. Signals were detected in a Chemi-Doc luminescence detection system. To verify equal amounts of proteins in each loaded sample, the membranes were stained afterwards with anti-actin antibody. For this purpose, the membranes were stripped by incubation in stripping buffer at 37 °C for 20 min and washed and blocked as described above. The primary goat anti-actin antibody was used at a 1:200 dilution and the HRP-coupled rabbit anti-goat antibody was used at a 1:8000 dilution.

• <u>TBST:</u>

140 mM NaCl 20 mM TRIS-HCl, pH 7.5 0.05% Tween-20

2.2.3. Fluorescence probes

In the following we analyzed properties and characteristics of activatable and alwayson fluorescence probes as well as of the fluorophore, Itrybe encapsulated in functionalized NPs, in combination with the tumor-specific antibody Herceptin.

2.2.3.1. General information

The fluorescence spectra of the pH-sensitive hydrophilic fluorescence dye, <u>CypHer5E</u> can be seen in Figure 7 (left). The quantum yield of CypHer5E in PBS is 0.26 at a pH of 7.8 and 0.27 at a pH of 5.3 (experiments performed by Jutta Pauli, BAM I.5, Berlin). CypHer5E in its hydrolyzed form has a molecular weight of 751 g/mol.

The fluorescence spectra of the hydrophilic fluorescence dye <u>Alexa Fluor 647</u> are depicted in Figure 7 (right). The quantum yield of the dye in PBS is 0.33 and the LT in water is 1.0 ns^{106} . Alexa Fluor 647 in its hydrolyzed form has a molecular weight of ~1200 g/mol.



Figure 7. Normalized spectrum of wavelength of CypHer5E and Alexa Fluor 647

CypHer5E has an absorption maximum at 645 nm and an emission maximum at 663 nm. Alexa Fluor 647 has similar spectra with an absorption maximum at 650 nm and an emission maximum at 670 nm. Spectra were taken from GE website (left)⁵⁴ and Invitrogen website (right)¹⁰⁶ and were modified.

The hydrophobic asymmetric cyanine dye <u>Itrybe</u> has an absorption maximum at \sim 590 nm and an emission maximum at \sim 750 nm. The quantum yield is 0.23 in ethanol.

2.2.3.2. Coupling of Herceptin antibodies to fluorescence dyes

Herceptin was coupled to the near-infrared fluorescence (NIRF) dyes, CypHer5E or Alexa Fluor 647, via their succinimidyl ester. Coupling resulted in conjugates with different dye to protein (DP) ratios of 0.5 to 6.4 dye molecules per antibody. Coupling was performed by Squarix Biotechnology GmbH, Marl. Purification was done via gel chromatography followed by a preparative gel filtration. DP ratios were determined photometrically by Jutta Pauli from the Bundesanstalt für Materialforschung und -Prüfung (BAM) I.5, Berlin.

2.2.3.3. Modification of NPs

Loading of polystyrene nanoparticles (PSNPs)⁶⁸ with Itrybe as well as surface modification with polyethylene glycol (PEG) and Herceptin¹⁰⁷ were performed by Thomas Behnke, BAM I.5, Berlin. A scheme of surface-modified nanoparticles (NPs) is shown in Figure 8. For surface modification 500 nmol PEG was added to the reaction solution per 1 mg polystyrene (PS). For bioconjugation in a second step 0.48 nmol Herceptin was added per mg PEGylated PS.



Figure 8. Schematic illustration of surface-modified PSNPs

PEG chains were coupled to Itrybe-loaded PSNPs via surface amino groups (NH₂). Thiolated Herceptin antibodies were coupled to PEG via its maleimid group in a Michael-Addition¹⁰⁷. Figure was modified from Steinhauser et al.¹⁰⁷.

The amounts of PSNPs used in this study are given as mg PS. Therefore, the number of particles per mg PS as well as the surface area are listed for different NP sizes in Table 1.

size	15 nm	25 nm	100 nm
Itrybe molecules/ NP	10	46	3100
NPs/mg PS [1/mg]	5.4 x 10 ¹⁴	1.2 x 10 ¹⁴	1.8 x 10 ¹²
relative amount: NPs/mg PS*	296	64	1
surface area/ mg PS [m ² /mg]	0.38	0.23	0.06

Table 1. Characteristics of Itrybe-loaded PSNPs

(Calculations were performed by Thomas Behnke, BAM I.5., Berlin.)

* relative to 100 nm NPs

Itrybe-loaded NPs of distinct sizes and surface modifications were analyzed in this work. Table 2 shows the composition of different NPs and their abbreviations, used in this study.

NP size	surface group	PEG chain size	antibody	abbreviation
100 nm	$\rm NH_2$	-	-	100 nm-NH ₂
100 nm	$\rm NH_2$	1.5 kDa	-	100 nm-PEG 1.5 kDa
100 nm	$\rm NH_2$	1.5 kDa	Herceptin	100 nm-PEG 1.5 kDa-Her
100 nm	$\rm NH_2$	1.5 kDa	2-fold amount* Herceptin	100 nm-PEG 1.5 kDa-HER x2
100 nm	$\rm NH_2$	10 kDa	Herceptin	100 nm-PEG 10 kDa-Her
25 nm	NH ₂	-	-	25 nm-NH ₂
25 nm	NH ₂	1.5 kDa	-	25 nm-PEG 1.5 kDa
25 nm	NH ₂	1.5 kDa	Herceptin	25 nm-PEG 1.5 kDa-Her
25 nm	$\rm NH_2$	1.5 kDa	5-fold amount* Herceptin	25 nm-PEG 1.5 kDa-Her x5
25 nm	$\rm NH_2$	10 kDa	-	25 nm-PEG 10 kDa
25 nm	$\rm NH_2$	10 kDa	Herceptin	25 nm-PEG 10 kDa-Her
15 nm	NH ₂	-	-	15 nm-NH ₂
15 nm	$\rm NH_2$	10 kDa	-	15 nm-PEG 10 kDa
15 nm	NH ₂	10 kDa	5-fold amount* Herceptin	15 nm-PEG 10 kDa-Her x5
15 nm	NH ₂	1.5 kDa	5-fold amount* Herceptin	15 nm-PEG 1.5 kDa-Her x5
15 nm	NH ₂	1/10 amount*, 1.5 kDa	5-fold amount* Herceptin	15 nm-PEG 1.5 kDa-1/10-Her x5

Table 2.	Itrybe-loaded	NPs of different	sizes and	surface modifications
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* *X/X amount* and *X-fold amount* refers to variations in the amount of PEG (500 nmol/mg PS) or Herceptin (0.48 nmol/mg PS) used for coupling procedure.

The amount of Herecptin that was successfully coupled to the surface of 100 nm particles was determined in a BCA assay following measurement of absorption spectra and correction for absorbance by Itrybe and scattering of the particles. Experiments were performed by Thomas Behnke, BAM I.5, Berlin. Table 3 shows the amount of Herceptin molecules per 100 nm NP used in this study.

probe	Herceptin molecules/ NP	Herceptin molecules/ mg PS
100 nm-PEG 1.5 kDa-Her	~70	1.3 x 10 ¹⁴
100 nm-PEG 1.5 kDa-Her x2	~100	1.8×10^{14}
100 nm-PEG 10 kDa-Her, probe 1*	~10	1.8 x 10 ¹³
100 nm-PEG 10 kDa-Her, probe 2*	~10	1.8×10^{13}

Table 3. Number of Herceptin molecules successfully coupled on NPs

* *probe 1* and *2* refers to two independently produced probes.

2.2.3.4. Spectroscopic analysis

Fluorescence conjugates:

As dyes were coupled to Herceptin at different DP ratios, the amount of pH-Her and Alexa-Her used in the study always refers to the amount of Immunoglobulin G (IgG) in the respective conjugates. Fluorescence emission spectra of CypHer5E-Herceptin (in the following referred to as pH-Her) and Alexa Fluor 647-Herceptin (in the following referred to as Alexa-Her) conjugates were measured in a concentration of 5 µg IgG in 75 µl phosphate buffered saline (PBS; 0.07 g/l). In order to evaluate the influence of pH-changes on fluorescence intensity of the conjugates they were measured in a solution with a pH of 7.5 and 5.5. The pH of PBS was adjusted with 1 M NaOH and 1 M HCl solution and pH values were controlled again after addition of fluorescence probes and adjusted again to the respective value if necessary.

Itrybe-loaded NPs:

Excitation and emission spectra of Itrybe-loaded NPs with different surfacemodifications and sizes were recorded. For this purpose 0.5 g/l of 100 nm and 25 nm NPs and 2 g/l of 15 nm NPs were measured in 75 μ l H₂O.

Measurements were performed in a spectrofluorometer. The settings used for recording of spectra are listed in section 2.2.8.1.

2.2.3.5. In vitro determination of fluorescence intensities

Fluorescence intensities of <u>Itrybe-loaded NPs</u> were measured to analyze influences of the environment on fluorescence of NPs. Itrybe-loaded NPs were measured in PBS or PBS containing 5 % BSA with a pH of 7.5 and 5.5, respectively. The pH value of the solutions was adjusted as described in 2.2.3.4. For measurement 0.25 g/l of 100 nm and 25 nm and 0.5 g/l of 15 nm NPs in 150 μ l of the respective solution were placed

in 96-well plates. Measurements were done in the Optix MX2 as described in section 2.2.8.4.

2.2.3.6. *In vitro* determination of fluorescence lifetimes

Fluorescence lifetime (LT) analyses were performed *in vitro* in order to identify probe-derived signals *in vivo* according to their LT. LTs of the <u>fluorescence</u> <u>conjugates</u> pH-Her and Alexa-Her as well as of the respective free, hydrolyzed fluorescence dyes (in the following referred to as pH-OH and Alexa-OH) were determined in PBS or PBS containing 5 % BSA with a pH of 7.5 and 5.5, respectively. The pH of the solutions was adjusted as described in section 2.2.3.4. For measurement 3 μ g (0.02 g/l) IgG or 0.03 nmol (0.2 mM) free dye in 150 μ l of the respective solution were placed in 96-well plates. Measurements were performed in the Optix MX2 as described in section 2.2.8.4.

2.2.4. Immunological methods

2.2.4.1. Immunoassays

Functionality of Herceptin after surface-conjugation to Itrybe-loaded NPs was evaluated via immunoassays. Here, specific binding to antigen-coated plates was analyzed and compared to the unspecific binding to uncoated plates and to binding of non-Herceptin-modified NPs. The extracellular domain of HER2, in the following referred to as HER2 antigen, was used as antigen.

All steps of the immunoassay were performed at RT. 96-well plates were either coated over night with 2.5 g/l HER2 antigen in 70 μ l TRIS buffered saline (TBS) buffer or treated only with TBS. Unspecific binding sites were blocked with 3 % BSA in TBS for 1h. NP-probes were added to the wells at different concentrations (from 0.02 to 2 g/l) in 70 μ l TBS. For competitive binding experiments a mixture of 14 μ g Herceptin-labeled NP probe (0.2 g/l) and increasing amounts of free Herceptin (0.005 to 1 μ g) in 70 μ l TBS were added per well. Probes were incubated for 2 h. After three washing steps with TBST (see section 2.2.2.4) the wells were covered with 100 μ l TBS and fluorescence was directly measured in the Odyssey infrared imaging system.

• <u>TBS:</u>

140 mM NaCl 20 mM TRIS-HCl, pH 7.5

2.2.5. Cell biological methods

2.2.5.1. Cell culture conditions

Cultivation:

KPL-4, MDA-MB-231, and 3T3 cells were propagated in standard Dulbecco's Modified Eagle's Medium (DMEM) containing 4.5 g/l glucose and L-glutamine. The medium was supplemented with 10 % fetal calf serum (FCS). Cells were cultivated under standard cell culture conditions at 37 °C in a humidified atmosphere under 5 % CO2 in an incubator and grown up to about 80 % confluency.

For sub-cultivation medium was removed from the cells, cells were washed twice with sterile Dulbecco's Phosphate Buffered Saline (DPBS), and treated with 0.05 % trypsine-EDTA solution until they had detached from the dish. The enzyme was inhibited by addition of medium in which the cells were subsequently resuspended. When necessary, cells were counted in a Neubauer counting chamber and plated in the desired concentration.

Cryoconservation:

For cryoconservation eukaryotic cells were detached as described, centrifuged (1,200 x g for 2 min at RT), resuspended in freezing medium (1-5 x 10^6 cells/ml) and aliquoted into storage vials. Vials were placed in an insulated box in a -80 °C freezer overnight and finally stored in liquid nitrogen.

Freezing medium:
 90 % FCS
 10 % dimethyl sulfoxide (DMSO)

Revitalization:

For revitalization cryoconserved cells were quickly thawed in a 37 °C water bath after being removed from liquid nitrogen. Then, cells were pipetted in pre-warmed medium and centrifuged (1,200 x g for 2 min at RT). The medium was removed from the cells and cells were resuspended in medium and then placed into a flask containing medium.

Mycoplasma test:

Mycoplasma contamination may cause adverse effects, such as changes in metabolism, growth rate, viability, DNA, RNA and protein synthesis, morphology etc., which lead to unreliable experimental results. Therefore, cell cultures were regularly tested for the presence of mycoplasma using Mycoplasma Plus PCR Primer Set according to the manufacturer's instructions.

2.2.5.2. Incubation of cells with fluorescence probes

Fluorescence conjugates:

To ensure better attachment of KPL-4 and MDA-MB-231 breast cancer cells to 4-well chambered coverglasses, they were covered with sterile poly-L-lysine solution (0.1 g/l). After 3 h incubation at 37 °C, coverslips were washed several times with sterile PBS, and used for seeding of cells. 48 h prior to incubation with fluorescence probes cells were seeded at a density of 2 x 10^4 KPL-4 cells and of 1.2 x 10^4 MDA-MB-231 cells per well in 500 µl DMEM. After 48 h the medium was removed and cells were washed once with PBS. Cells were supplemented with 500 µl DMEM and 0.1 M 4-(2hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) buffer containing 5 µg fluorescence conjugates or 0.2 nmol of the free fluorescence dyes (corresponding to the amount of dye in 5 μ g conjugate with a DP ratio of 6) and incubated for 8 h at 37 °C. In order to analyze signals deriving from membrane bound conjugates, probes were also incubated with the cells for 8 h at 4 °C where internalization should be strongly reduced⁵. Afterwards, the medium was removed and the cells were washed several times with PBS to remove unbound substances. Subsequently, cells were incubated with 1 µg Hoechst 33342 in 500 µl PBS (2 mg/l) for 10 min in order to stain the nuclei of living cells. Hoechst solution was removed and cells were covered with 500 µl PBS and directly used for fluorescence microscopy. The settings used for detection of fluorophores are listed in section 2.2.8.3.

Itrybe-loaded NPs:

Adherent breast cancer cells were seeded on poly-L-lysine-coated cover slips (treated as described above) in 24-well chambers 48 h prior to incubation with Itrybe-loaded NPs. KPL-4 cells were seeded at a density of 3×10^4 cells and MDA-MB-231 cells were seeded at a density of 1.8×10^4 cells per well in 500 µl DMEM. After 48 h the medium was removed and cells were washed once with PBS and supplemented with 500 µl DMEM and 0.1 M HEPES buffer containing NPs at a concentration of 1 g/l. The cells were incubated for different time periods (30 min to 5 h) at 37 °C and 4 °C. Afterwards, the medium was removed and the cells were fixed with 4 % formaldehyde in PBS for 10 min on ice and then washed 3 times with PBS again. The cells were mounted with an antifade reagent supplemented with 4',6-diamidino-2-phenylindole (DAPI) for counter stain of cell nuclei. Samples were allowed to dry for 24 h at 4 °C and subsequently used for fluorescence microscopy. The settings used for detection of fluorophores are listed in section 2.2.8.3.

2.2.5.3. Assessment of cell viability

Effects of NPs on proliferation and viability of cells were quantified via use of WST-1 cell proliferation reagent. This assay is based on the enzymatic cleavage of the tetrazolium salt WST-1[2-(4-Iodophenyl)-3-(4-nitrophenyl)-5-(2,4-disulfophenyl)-2H-tetrazolium] (WST-1) to a water-soluble formazan dye by mitochondrial succinate-tetrazolium reductase which is active in viable cells. Absorbance of the dye is directly proportional to the amount of viable cells.

3T3 mouse fibroblasts were seeded at a density of 5 x 10^3 cells/well in 96 well plates 24 h before NP treatment. Cells were incubated with 0.01 to 2 g/l of 100 nm NPs or left untreated in DMEM for 24 h at 37 °C. To consider the absorbance caused by the Itrybe-loaded NPs itself, prior to addition of WST-1-reagent absorbance was measured in the Wallac 1420 Victor 2 Multilabel Counter at 450 nm. 10 µl of WST-1 ready-to-use solution was added to the wells and after 30 min of incubation at 37 °C absorbance was measured again at 450 nm. To obtain only the signals derived from the formazan dye, absorbance measured prior to addition of WST-1.

2.2.5.4. Cell preparation for animal experiments

One day before implantation, KPL-4 or MDA-MB-231 cells were sub-cultivated to achieve exponential growth. On the day of implantation cells were trypsinized and washed twice with medium. The cell number was determined as described in 2.2.5.1 and suspension volumes containing 5×10^6 KPL-4 cells or 1×10^6 MDA-MB-231 cells were pelletized under centrifugation at 1000 x g, the medium was removed, and cells were resuspended in 500 µl medium.

2.2.6. In vivo experiments

All animal experiments were according to § 8 of the animal protection law in the version of 2006 and all animal protocols were approved by the administration of lower Saxony with the animal experiment number 33.14-42502-04-10/0064.

2.2.6.1. Animal models

Experiments were performed on female athymic nude mice, strain NMRI-Fox1nu/nu. Feeding occurred via pelletized Extrudate 1439 food and acidulated water with a pH of 2.5. Two weeks before the mice were scanned in the Optix MX2 they received chlorophyll-depleted low fluorescent food, GLP Nafag feed 890 or Regime 210, U8959A01R. The mice were maintained in a sterile environment in special cages with

filter huts placed in a scantainer. Cages, bedding, and water were autoclaved and the food was gamma irradiated.

2.2.6.2. Orthotopic implantation of breast tumor cells in mice

KPL-4 and MDA-MB-231 breast tumor cells were implanted in the right abdominal mammary fat pad of female nude mice at the age of 6 to 12 weeks. For this purpose mice were anesthetized by intraperitoneal (i.p.) injection of 15 mg/kg body weight xylazine und 75 mg/kg body weight ketamine in a 1:1 dilution with isotone sodium chloride (NaCl, 0.9 %) solution using an insulin syringe (Ominifix 40 solo) and a sterican needle (26G x 1/2). The solution containing the respective amount of tumor cells (5 x 10^{6} KPL-4 cells or 1 x 10^{6} MDA-MB-231 cells, prepared as described in section 2.2.5.4) was centrifuged, the medium was discarded, and cells were resuspended in a volume of 30 µl PBS and taken up with an insulin syringe with integrated needle (30G). Anesthetized mice were placed under a laminar flow in a dorsal position and the skin was opened with a small incision in the region over the right abdominal mammary fat pad. Tumor cells were implanted very slowly into the mammary fat pad. The needle was slowly withdrawn after a 1-minute delay. The mammary gland was then returned, and the incision was closed using absorbable surgical suture material, 4/0. Afterwards, mice were placed in cages on a heating plate at 37 °C for 3 h. In time intervals of three days body weight and general condition of the mice were controlled.

2.2.6.3. Application of probes

Mice were used for *in vivo* fluorescence imaging experiments 1 to 4 month after tumor implantation when tumors reached sizes of approximately 0.3 cm³. Biodistribution experiments with NPs were carried out on tumor-free nude mice. All fluorescent probes were injected solved in NaCl, 0.9 % with an insulin syringe with integrated needle (30G x $\frac{1}{2}$). During injection, the animals were anesthetized with vaporized isoflurane at 0.8 – 1 % concentration.

The amounts of fluorescence probes injected in mice and the different application methods are listed in Table 4. Before and after probe injection, mice were imaged in the Optix MX2 as described in section 2.2.8.4.

Table 4. Amount and application methods of fluorescent probes injected in mice

Different amounts of fluorescent Herceptin-conjugates, the respective free fluorescence dyes, or Itrybeloaded NPs were injected in mice, either *intravenously* (*i.v.*), subcutaneously (*s.c.*), or *intratumoral* (*i.t.*).

probe	amount	volume	application
fluorescence conjugates	25 μg IgG	150 µl	i.v.
free dyes	0.8 nmol*	150 µl	i.v.
fluorescence conjugates	10 µg IgG	50 µl	5.C.
free dyes	0.1 nmol	50 µl	5.C.
100 nm NPs	400 µg	200 µl	i.v.
25 nm NPs	400 µg	200 µl	i.v.
15 nm NPs	1 mg	200 µl	i.v.
15 nm NPs	0.8 – 100 µg	30 µl	S.C.
15 nm NPs	1.6 µg	30 µl	i.t.

* Amount corresponds to the amount of dye in the Herceptin-conjugates with a DP ratio of 5.

2.2.6.4. Dissection

At the end of the *in vivo* experiments mice were sacrificed by cervical dislocation under high concentration of isoflurane anesthesia. The animals were examined macroscopically for noticeable abnormalities and tumor development. Subsequently, tumors were excised and tumor sizes were measured by a caliper. Under the assumption that the tumor shape is ellipsoid the tumor volume was calculated according to the formula V $[cm^3] = (length x width x height) / 2$. Furthermore, the abdomen and the thoracic cavity were opened and liver, kidneys, spleen, lung, heart, stomach, and gut were isolated and imaged *ex vivo*.

2.2.6.5. Tumor preparation

One half of the tumor was stored in a cryotube at - 80 °C and used for mRNA and/or protein analysis. The other half was fixed with formaldehyde (4 % in PBS) over night, embedded in paraffin and used for microscopic examination of tumor tissue. For this purpose, the formalin fixed tumors were placed in Tissue Tek cassettes, rinsed under the water-tap for half an hour and dehydrated and paraffinized in the apparatus for tissue processing over night.

applied substance repetition cycles time 1 h 75 % ethanol three 1 h 30 min 96 % ethanol two three 1 h 15 min 100 % ethanol 2 h xylol two two 1 h paraffin

The following steps were performed:

2.2.7. Histological examinations of tumor tissue

2.2.7.1. Generation of paraffin sections from tumor tissue

Sections of the paraffin-fixed tumors were made with the microtome in 2.5 to 5 μ m thickness. Sections were attached to object slides, and stored at RT until staining.

2.2.7.2. Haematoxylin-Eosin (HE) staining of paraffin-embedded tumor sections

Sections were incubated over night at 37 °C in the incubator to improve attachment of the tissue to the microscope slides. Afterwards the tumor sections were deparaffinized twice with xylol followed by rehydration in decreasing concentrations of ethanol (each for 2 min). The Haematoxylin staining of cell nuclei was done for 5 to 10 min and after washing with water the cell structures were stained with Eosin G (1: 10 dilution in H_2O) for 5 to 10 min. Tumor sections were washed with water and dehydrated with increasing concentrations of ethanol (50 %, 70 %, 90 %, 100 %) and xylol and mounted in hydrophobic mounting medium (DePex). Sections were analyzed under the light microscope.

2.2.7.3. CD31 staining

Paraffin-embedded tumor sections (2.5 µm thickness) were stained with the antiendothelial cell marker cluster of differentiation 31 (CD31) antibody after being deparaffinized as described in section 2.2.7.2. Prior to antibody treatment, sections were incubated for 20 min in target retrieval solution at 98 °C in a steamer. After cooling down, the sections were washed twice with TRIS buffer. The endogenous peroxidase was blocked with 3 % hydrogen peroxid in PBS for 10 min at RT. Afterwards unspecific binding sites were blocked with commercial blocking buffer from fish for 15 min at RT. Incubation of tissue with anti-mouse CD31 antibody at a 1:100 dilution was carried out in commercial background reducing antibody diluent over night at 4 °C. Afterwards, the sections were washed twice with TRIS buffer and incubated with the secondary HRP-coupled anti-rat antibody at a 1:200 dilution in TRIS-buffer for 1 h at RT. Sections were washed twice with TRIS buffer followed by incubation with peroxidase substrate, 3-amino-9-ethylcarbazole (AEC) for 30 min at RT, and washed twice with water. Staining of cell nuclei was performed by incubation in Haematoxylin for 10-20 sec. The tumor sections were washed with water and mounted with aqueous mounting medium (Aquatex). CD31 staining was analyzed under the light microscope.

<u>TRIS buffer</u>
 50 mM TRIS-HCl (pH 7.6)
 150 mM NaCl

2.2.7.4. Near-infrared (NIR)-microscopic analysis of tumor sections

Tumors treated in vivo with fluorescent probes:

Paraffin sections (2.5 µm thickness) of tumors of mice treated with Herceptinconjugates *in vivo* were deparaffinized as described in section 2.2.7.2. and mounted with hydrophobic mounting medium containing DAPI for counterstain of cell nuclei. Samples were allowed to dry for 24 h at 4 °C. Sections were analyzed under the fluorescence microscope as described in section 2.2.8.3.

Ex vivo treatment with fluorescent probes:

Paraffin embedded tumor sections (5 µm thickness) were deparaffinized as described in section 2.2.7.2. The sections were washed 3 times with TBS containing 1 % goat serum and blocked with TBS and 10 % goat serum for 1 h at RT. Tumor samples were incubated with Itrybe-loaded NPs in TBS and 10 % goat serum for 2 h at 4 °C. Afterwards, the sections were washed several times with TBST to remove unbound NPs. The cells were mounted with an antifade reagent supplemented with DAPI for counter stain of cell nuclei and allowed to dry for 24 h at 4 °C. Sections were analyzed under the fluorescence microscope as described in section 2.2.8.3.

2.2.8. Imaging devices

2.2.8.1. Spectrofluorometer

Excitation and emission spectra of fluorescence probes and dye-loaded NPs were recorded in a Spectrofluorometer QuantaMaster equipped with a xenon lamp as excitation source and two fluorescence detectors. The solutions were measured in a

quartz glass precision cuvette with a light pass of 3 x 3 mm. Spectra were recorded with a step size of 1 nm and an integration time of 1 sec. Wavelengths settings used for recording of spectra are listed in Table 5. Analysis of spectroscopic measurements was performed with the software FeliX32 Analysis 1.0. Measurements are quantified as the mean of signals recorded from both detectors.

	excitation scan		emission scan	
probe	λ _{em} [nm]	recording range [nm]	λ _{ex} [nm]	recording range [nm]
fluorescence conjugates	not analyzed	not analyzed	635	610 - 750
Itrybe-loaded NPs	740 - 770*	450 - 740	590	650 - 875

Table 5. Settings used for recording of spectra of fluorescent probes

 $\lambda_{\text{em}}:$ excitation wavelength; $\lambda_{\text{ex}}:$ emission wavelength

* λ_{em} was chosen according to the absorption maximum of the NPs with different sizes, which was 740 nm for 15 nm NPs, 750 nm for the 25 nm NPs, and 770 nm for the 100 nm NPs.

2.2.8.2. Odyssey infrared imaging system

Functionality of bioconjugated Itrybe-loaded NPs was evaluated in the Odyssey infrared imaging system (see section 2.2.3.3 for production of NPs) by exciting the probes with a 685 nm laser diode. Fluorescence intensity was detected in a raster mode with a resolution of 84 μ m. Analysis of fluorescence measurements was performed with the Odyssey Application Software, Version 2.1.

Average fluorescence intensity was calculated as the sum of the intensity in an area set in the respective well divided by the number of measurement points in the area.

2.2.8.3. Fluorescence microscope

For examination of fluorescence stained human KPL-4 and MDA-MB-231 breast tumor cells, the fluorescence microscope Axiovert 200M equipped with a NIR-sensitive ORCA-ER digital camera was used. Filter settings are illustrated in Table 6.

probe	λ _{ex} [nm]	beam splitter [nm]	λ _{em} [nm]
fluorescence probes	640 ± 15	660	690 ± 25
Itrybe-loaded NPs	620 ± 30	662	809 ± 45.5
DAPI, Hoechst	365 ± 12.5	395	445 ± 25

Table 6. Microscopy filter settings used for detection of fluorophores

Image generation and processing were performed with the software systems AxioVision Rel.4.6. and ImageJ, respectively.

2.2.8.4. In vivo NIRF imager, Optix MX2

The Optix MX2 system¹³ was used for characterization, quantification and illustration of NIRF probes *in vitro* and their localization in living animals. The system is a fluorescence imager working in reflectance mode and scanning in a raster scheme with a minimal step size of 0.5 mm. During the scan each raster point is irradiated with 9 ps laser impulses with a repetition rate of 80 MHz. The system utilizes a single photon counting detector, measuring the time between an excitation pulse and the detection of the first fluorescence photon. As illustrated in Figure 9, the generated temporal point spread function (TPSF) is a curve representing the distribution of the number of photons coming from a raster scan point on the sample over time. Under the assumption of a mono exponential decay of the measured intensity the fluorescence LT is calculated by linear regression in the slope of the logarithmic TPSF (Figure 9). Thus, the TPSF provides information about the fluorescence intensity and LT of the fluorescence transition.



Figure 9. TPSF of one raster scan point

Logarithmic photon time of flight histogram (TPSF) exemplarily shown for one scan point taken from a scan of a mouse injected with a fluorophore. The time between excitation pulse and detection of fluorescence photons is depicted on the x-axis. The

fluorescence intensity in a log_{10} scale is illustrated on the y-axis. The fluorescence intensity is determined by the area under the curve. From the decay of the curve the fluorescence LT is calculated.

For scanning mice were placed inside the imaging device on a table preheated to 37 °C and anesthetized with vaporized isoflurane at 0.8 to 1 % concentration throughout the imaging session.

The excitation lasers (λ_{ex}) and emission filters (λ_{em}) selected for detection of the fluorescence conjugates, free fluorescence dyes and Itrybe-loaded NPs are shown in Table 7.

probe	λ _{ex} [nm]	λ _{em} [nm]
fluorescence conjugates	635	670 ± 20
Itrybe-loaded NPs	635/670*	700 long pass

Table 7. Laser and filter settings used for detection of fluorophores in the Optix MX2

* Due to the very bright absorption spectrum of Itrybe, excitation was possible with the 635 and 670 nm laser.

Table 8 shows the measurement time per raster position (integration time) and step size generally used for the different scans in the Optix MX2.

situation:	in vivo		ex vivo		in vitro
scan:	tumor	whole body	tumor	other organs	solution
integration time [sec.]	1	0.7/0.2*	1	0.7	1
step size [mm]	1	1.5	1	1.5	0.5

 Table 8. Integration time and step sizes used for fluorescence measurements in the Optix MX2

* Biodistribution experiments in healthy mice were performed with an integration time of 0.2 for whole body scans.

The laser power was adapted automatically for optimal detection of the fluorescence yield. Analysis of the fluorescence measurements was performed with the software optiview 1.00.00 (ART).

The following parameters were calculated for data analysis:

Normalized counts (NC):

The measured fluorescence signals (counts) were normalized for varying laser power and integration times, thus allowing comparison of measurements with different settings.

Average fluorescence intensity:

The average fluorescence intensity over a certain area was determined as the sum of the NC in the area divided by the number of measurement points in the area.

Subtraction of autofluorescence:

In order to eliminate unspecific signals deriving from endogenous particles of the mouse, subtraction of autofluorescence was performed. For this purpose, the average

fluorescence intensity in a region of the prescan of the mouse (e.g. the tumor area) was calculated and subtracted from the intensity maps after probe application.

Contrast to noise ratio (CNR):

To consider the signal difference from the average intensity over a certain area and those over the background, the contrast to noise ratio (CNR) was calculated. This ratio provides information about the level of signal differences and the affection of the signal by the noise (e. g. electrical- or photon noise). The CNR was calculated as followed:

CNR = $\frac{(average intensity tumor area - average intensity backround area)}{standard deviation background area}$

Fluorescence LT:

Fluorescence LT is the average time an atom stays in the meta-stable state before its transition to the ground state under emission of a fluorescent photon. Thus the LT of the fluorescence transition is the characteristic value of a fluorescent probe. Fluorescence LT measurements enable a differentiation between fluorescence signals originating from the fluorescence probe and unspecific- or autofluorescence.

Average fluorescence LT:

The average fluorescence LT over a certain area was determined as the sum of the LTs in the area divided by the number of measurement points in the area.

LT goodness of fit:

For determination of the quality of the measured LT, the LT goodness of fit was calculated. Since, the decay of the fluorscence over a certain point is modeled with a mono exponential decay function, a signal derived from two or more fluorescence sources would no longer show a mono exponential decrease but an overlay of several signal decreases. Thus, the determined LT would arise from two or more fluorescent species leading to a high LT error and a poor accuracy of the LT estimation. A LT-calculation with a LT goodness of fit of 1 ± 0.2 is considered to have a high accuracy.

3. RESULTS

3.1. Characterization of tumor model organisms

Fluorescent probes evaluated in this study were coupled to Herceptin in order to achieve specific binding of probes to HER2-positive breast tumor cells *in vitro* and *in vivo*. Therefore, HER2-positive, KPL-4¹⁰¹ as well as HER2-negative, MDA-MB-231¹⁰² tumor models were established. Prior to evaluation of fluorescence probes *in vivo*, tumor cells were analyzed for HER2 expression levels and their suitability to bind Herceptin-conjugated probes.

3.1.1. Orthotopic breast tumor models in nude mice

Tumor-bearing mice were generated by orthotopic implantation of KPL-4 or MDA-MB-231 tumor cells in the right abdominal mammary fat pad of female nude mice at the age of 6-12 weeks. Imaging of mice bearing KPL-4 tumors started 8.3 ± 4.6 weeks after implantation when the tumors reached a mean size of 0.28 ± 0.19 cm³ (n = 24). MDA-MB-231 tumor-bearing mice were scanned after 9.9 ± 9.2 weeks, when tumors had a mean size of 0.25 ± 0.07 cm³ (n = 16). In Figure 10, representative KPL-4 (Figure 10A) and MDA-MB-231 (Figure 10B) tumor-bearing mice are shown. Due to the lack of fur in nude mice, the tumor is clearly visible *in vivo* (Figure 10A,B; upper part) and the growth rate can be assessed by caliper measurements easily. Section through the excised tumors show necrotic areas, which have been already developed at the described tumor size in both models (Figure 10A,B; lower part). In addition, for the KPL-4 tumors, a soft consistency at the tumor center was noted (Figure 10A, lower, right picture). No invasive growth and metastases were detected in both tumor types.



Figure 10. Macroscopic appearance of orthotopic breast tumor xenografts in nude mice

Human tumor cells were implanted orthotopically in the right abdominal mammary fat pad of female nude mice. **A,B.** Representative *ex vivo* picture of (**A**) a nude mouse bearing a KPL-4 tumor (T) with a size of 0.22 cm³ and (**B**) a nude mouse bearing a MDA-MB-231 tumor (T) with a size of 0.36 cm³. (**A,B**) Both tumors are supplied by large-diameter blood vessels (V; lower part, left). A section through the excised tumors (lower part, right) illustrates inhomogeneous tissue with necrotic areas.

Morphology of KPL-4- and MDA-MB-231 tumors was evaluated on sections stained with HE or with a CD-31 antibody for assessment of tumor blood vessels. For both tumor types, KPL-4 (Figure 11A-C; n = 12) and MDA-MB-231 (Figure 11D-F; n = 8) cancerous cells and also vital tumor tissue, such as mammary glands and ducts can be observed (Figure 11A,D; black arrow). Both tumors are supplied by several small-diameter capillaries (Figure 11C,F; red stars). Towards the tumor border a network of microvessels is observed whereas the vessel density seems to be lower towards the center (data not shown). Moreover, necrotic areas can be seen between vital tumor tissue (Figure 11A,B,D,E; white arrow). Here, less necrosis is observed for the MDA-MB-231 tumors (Figure 11A,B; white arrow) that is, in comparison to the KPL-4 tumors (Figure 11D,E) located more towards the tumor center.



Figure 11. Histology of breast tumor xenografts

A-C. Sections of a representative KPL-4 tumor (n = 12). **A.** HE staining shows several necrotic areas (white arrows) between vital tumor tissue and normal breast tissue (black arrow). **B.** HE staining at a higher magnification illustrates a necrotic area in detail (white arrow). **C.** CD31 staining marks endothelial cells (red stars). **D-E.** Sections of a representative MDA-MB-231 tumor (n = 8). **D.** HE staining shows some necrotic areas (white arrows) between vital tumor tissue (black arrow). **E.** HE staining at a higher magnification illustrates a necrotic area in detail (white arrow). **F.** CD31 staining marks endothelial cells (red stars). Images A,D. were taken with a x 2.5 objective and images B,C,E,F. were taken with a x 10 objective.

Taken together, KPL-4 and MDA-MB-231 tumor cells developed orthotopic breast tumors in nude mice that are partly necrotic but well vascularized.

3.1.2. HER2 expression in breast tumors

Tumors used in this study were analyzed for *HER2* mRNA expression levels via quantitative real-time PCR. In Figure 12, the normalized expression ratio of *HER2* in KPL-4 and MDA-MB-231 cells as well as in a corresponding tumor, are illustrated. High expression ratios were found in a KPL-4 tumor (sample no. 3383) and in KPL-4 cells whereas a MDA-MB-231 tumor (sample no.3450) and MDA-MB-231 cells show a very low *HER2* expression.



Figure 12. Expression levels of *HER2* mRNA in KPL-4 and MDA-MB-231 cell lines and tumors

Representative results of a quantitative real-time PCR (n = 2). Expression levels of *HER2* are normalized to the amount of human transferrin. The highest normalized ratio detected in the KPL-4 tumor (sample no. 3383)

was set to 100 %. All measurements were performed in triplicate.

In addition, the expression levels of the HER2 protein were determined. For this purpose, 50 µg of total protein amount of KPL-4 and MDA-MB-231 cells as well as of two tumors, from each cell type were separated via SDS-PAGE and analyzed for the presence of HER2 protein by Western blot. As illustrated in Figure 13, KPL-4 cells and -tumors both show high expression of HER2 whereas MDA-MB-231 cells and tumors show a very weak protein band at the size of HER2 (185 kDa) after incubation with the anti-HER2 antibody. Staining for actin revealed that for all samples a comparable amount of protein was used. (The Western blot experiments were performed by Dr. Fernanda Ramos.)



Figure 13. HER2 protein levels in KPL-4 and MDA-MB-231 cell lines and tumors

Western blot analysis revealed that KPL-4 cells (left) and tumors samples (right; no. 3383 and 3380) express HER2 protein showing a band with the size of 185 kDa after incubation with an anti-HER2 antibody (upper panel). In contrary,

almost no staining can be observed in the MDA-MB-231 cells (left) and tumors (right; samples no. 3444 and 3450). Staining of actin reveals comparable amounts of total protein in all samples (lower panel). Representative results of three independent experiments performed by Dr. Fernanda Ramos are shown.

Taken together, KPL-4 cells as well as tumors developed from these cells express HER2, on the mRNA and protein level. In contrary, MDA-MB-231 cells and tumors show a very low expression of HER2. Therefore, KPL-4 cells (defined as HER2-positive) provide an excellent model for the evaluation of Herceptin-conjugated fluorescence probes whereas MDA-MB-231 cells (defined as HER2-negative) were used as negative controls.

3.1.3. Validation of tumor models for evaluation of fluorescence probes

To evaluate the suitability of KPL-4 and MDA-MB-231 tumors, to be applied for evaluation of novel fluorescent conjugates *in vivo*, it is important to ensure that the dye-coupled Herceptin binds to HER2-expressing tumors that developed in mice. In order to enable a high sensitivity of signal detection, fluorescence distribution in the tumor was investigated by using an always-on fluorescent Herceptin-conjugate with high amounts of dyes per antibody. 24 h after *i.v.* injection of Alexa-Her at a DP ratio of 3.4, one KPL-4- and one MDA-MB-231 breast tumor were examined under a fluorescence microscope. As shown in Figure 14A, Alexa-Her-derived fluorescence can be detected inside the KPL-4 tumor, located around vital tumor cells. The antibody seems to be not equally distributed in the tumor tissue: Stronger fluorescence is observed in some clustered regions whereas signals in other areas are comparable weak. In contrary, as illustrated in Figure 14B, almost no Herceptin conjugate-derived signals can be detected inside the control MDA-MB-231 tumor.





Sections of paraffin fixed tumors were examined under a fluorescence microscope with a x 40 objective. A KPL-4 and a MDA-MB-231 tumor excised from mice 24 h after receiving *i.v.* injections of 25 µg Alexa-Her at a DP ratio of 3.4 is shown. For each box, on the left upper side, counter stain of cell nuclei with DAPI and on the left lower side, probe-derived signals are illustrated. Merged images of the cell nuclei (blue) and the probe (red) are shown on the right. **A.** The Herceptin-conjugates are detected inside the KPL-4 tumor. Some tumor areas show stronger accumulation of the conjugates then others. **B.** In the MDA-MB-231 tumor almost no Alexa-Her derived fluorescence is detected. The exposure time for detection of Alexa-Her was 300 ms. Bars represent 50 µm.

Here, we confirmed that breast tumors of HER2-positive, KPL-4 and HER2-negative, MDA-MB-231 cells can be used for the evaluation of Herceptin-conjugated fluorescence probes.
3.2. Improvement of tumor detection via pH-sensitive NIRF probes

There is a high demand for innovative probes with improved fluorescent properties to increase the sensitivity of tumor detection *in vivo*. In this context, functional fluorescence dyes that selectively activate at the targeted tumor site are promising tools to reduce unspecific signals in the background deriving from circulating and unbound fluorescence probes.

In this part of the work, the functionality and sensitivity of the pH-sensitive tumorspecific probe, pH-Her was examined and compared to the always-on fluorescence conjugate Alexa-Her.

3.2.1. pH-Her conjugates at different DP ratios increase fluorescence in an acidic environment

In order to investigate the spectroscopic properties of different fluorescence probes with distinct dye to protein (DP) ratios, ranging from statistically 0.5 up to 6.4 dye molecules per Herceptin, fluorescence emission spectra of Herceptin-conjuates were recorded. To illustrate influences of pH on fluorescence emission, the spectra of all probes were measured in a neutral pH of 7.5 and an acidic pH of 5.5.

As illustrated in Figure 15, the shapes of the emission spectra of the pH-sensitive antibody-dye-conjugates, pH-Her and the always-on control conjugates, Alexa-Her are comparable to those of the free fluorescence dyes (CypHer5E and Alexa Fluor 647) showing an emission maximum at about 665 nm for pH-Her and about 668 nm for Alexa-Her (for details see section 2.2.3.1, Figure 7). In an acidic environment (pH of 5.5; n = 3), all pH-Her probes increase their fluorescence compared to measurements at a neutral pH (of 7.5; n = 3). In contrast, the always-on Alexa-Her probes show no remarkable change in fluorescence in response to a low pH (n = 3).





Representative fluorescence emission spectra of pH-Her and Alexa-Her probes recorded from 620 to 750 nm at λ_{ex} of 635 nm in a spectrofluorometer (n = 3). Probes were measured in PBS at a pH of 7.5 (blue curve) and of 5.5 (red curve). **A-F.** Fluorescence of pH-Her with different DP ratios (from 0.5 to 6.4) remarkably increases in a pH of 5.5 compared to a pH of 7.5. **G,H.** No remarkable change in fluorescence can be observed for the Alexa-Her probes at low and high DP ratios (of 1.3 and 3.4) in response to an acidic pH.

Table 9, summarizes the fluorescence intensities of all probes calculated from the λ_{em} maximum of the spectra illustrated in Figure 15. The pH-Her and Alexa-Her conjugates both show a clear tendency of higher fluorescence with an increase in DP ratio. The later probe also demonstrates much higher intensity when compared to pH-Her at comparable DP ratios. For example, the mean intensity of pH-Her with a DP ratio of 1.2 at a pH of 5.5 is 44,446 counts/s whereas Alexa-Her with a DP ratio of 1.3 has an almost 4-fold higher mean fluorescence of 169,582 counts/s, under the same experimental conditions. pH-Her conjugates reveal a slightly less pH-sensitivity with increasing DP ratios. For example, pH-Her with a DP ratio of 5.0 shows a 3.9-fold increase in fluorescence when measured in a neutral pH in correspondence to a pH of 5.5 (factor). In comparison, pH-Her with a DP ratio of 1.6 shows the highest increase in fluorescence with a factor of 5.6. In contrast, both Alexa-Her conjugates do almost not change fluorescence in a low pH.

Table 9. Fluorescence intensities of Herceptin-conjugates in dependence of pH

Fluorescence of 0.07 g/l conjugates was recorded in a spectrofluorometer at λ_{em} of 665 nm for pH-Her and 668 nm for Alexa-Her. Measurements were performed in PBS at a pH of either 7.5 or 5.5. The fluorescence counts/s of three independently performed experiments (1 to 3) as well as the mean values are illustrated. The pH-sensitivity of the conjugates is shown as the factor of the mean fluorescence at a pH of 5.5 divided by the mean fluorescence at pH 7.5.

DP 0.5	pH 7.5 [counts/s]		pH 5.5 [counts/s]		DP 1.2	pH 7.5 [counts/s]		pH 5.5 [counts/s]
1	4,311		23,624	-	1	8,415		38,205
2	3,445		24,161		2	9,186		52,740
3	5,202		23,457		3	9,048		42,393
mean	4,319 ± 879		23,747 ± 368		mean	8,883 ± 411		44,446 ± 7,482
factor		5.5		-	factor		5.0	
DP 1.6	pH 7.5 [counts/s]		pH 5.5 [counts/s]		DP 1.8	pH 7.5 [counts/s]		pH 5.5 [counts/s]
1	11,890		67,832	-	1	12,963		58,480
2	10,411		59,381		2	11,534		56,458
3	9,751		51,224		3	13,136		52,796
mean	10,684 ± 1,095		59,479 ± 8,305		mean	12,544 ± 879		55,912 ± 2,881
factor		5.6			factor		4.5	

pH-Her conjugates

DP 5.0	pH 7.5 [counts/s]		pH 5.5 [counts/s]	DP 6.4	pH 7.5 [counts/s]		pH 5.5 [counts/s]
1	28,155		109,445	1	32,622		120,354
2	23,283		97,068	2	40,924		174,290
3	21,818		80,698	3	20,990		82,792
mean	24,418 ± 3,318		95,737 ± 14,419	mean	31,512 ± 10,013		125,812 ± 45,992
factor		3.9		factor		4.0	

pH-Her conjugates

Alexa-Her conjugates

DP 1.3	pH 7.5 [counts/s]		pH 5.5 [counts/s]	_	DP 3.4	pH 7.5 [counts/s]		pH 5.5 [counts/s]
1	157,824		152,775	_	1	317,535		264,915
2	152,639		163,651		2	332,074		302,574
3	169,503		192,321	_	3	338,816		310,969
mean	159,988 ± 8,638		169,582 ± 20,430		mean	329,475 ± 10,876		292,819 ± 24, 528
factor	•	1.1			factor		0.9	

After spectroscopic characterization, two pH-Her conjugates with distinct properties were chosen for further analysis: pH-Her with a low DP ratio of 1.6 (pH-Her DP 1.6) due to its high pH-sensitivity (5.6-fold increase in fluorescence from pH 7.5 to pH 5.5) and pH-Her with a high DP ratio of 5.0 (pH-Her DP 5.0) due to its relatively high fluorescence intensity (96,000 counts of 0.07 g/l probe at a pH of 5.5). As controls, the always-on conjugates Alexa-Her with a DP ratio of 1.3 (Alexa-Her DP 1.3) and Alexa-Her with a DP ratio of 3.4 (Alexa-Her DP 3.4) were used for further experiments in order to have probes with low and high DP ratios, comparable to the selected pH-Her probes.

3.2.2. LTs of fluorescence probes are influenced by the environment

Due to its comparable high pH-sensitivity, pH-Her DP 1.6 was applied for LT-gated analysis *in vivo*. Therefore, LTs of pH-Her DP 1.6 and of the control conjugate Alexa-Her DP 1.3 were analyzed in detail and compared to LTs of the respective free fluorescence dyes pH-OH and Alexa-OH.

The influence of the environment on the LT of pH-sensitive- and control-probes was analyzed by examination of the probes under different conditions that are known to influence LTs, like pH (pH of 7.5 and 5.5) or presence of proteins (PBS/BSA 5%)^{2,108,109}. Here, BSA was chosen as serum albumin presents the main plasma protein (60 %) in humans¹¹⁰. In addition, LTs were also determined under *in vivo* conditions after *s.c.* injection of probes in mice. The mean LT of all measurements for each probe was defined as the predicted LT and used for probe identification in *in vivo* LT-imaging.

As summarized in Table 10, pH-Her DP 1.6 has a mean LT of 1.3 ns and shows low variation upon changes in pH, in the presence of proteins, or after *s.c.* injection. Alexa-Her DP 1.3 has a mean LT of 1.5 ns and reveals slightly longer LTs in the presence of proteins (BSA). The respective free hydrolyzed fluorophores pH-OH and Alexa-OH, both have shorter mean LTs (of 1.1 and 1.3 ns) compared to the corresponding Herceptin-conjugated fluorophores, pH-Her and Alexa-Her. Moreover, LTs of both free dyes increase in the presence of proteins, especially at a low pH as well as after *s.c.* injection in mice.

Table 10. Determination of the probes' LT under different conditions

LTs of pH-sensitive- and control-probes were determined in PBS and PBS/BSA 5 % with a pH of 7.5 and 5.5, respectively as well as after *s.c.* injection in mice. Measurements were performed in triplicates in the Optix MX2. The mean LT of all measurements for each probe was defined as the predicted LT.

pH-Her	LT [ns]				
DP 1.6	pH 7.5		pH 5.5		
PBS	1.25 ± 0.01		1.26 ± 0.01		
PBS/BSA 5%	1.29 ± 0.03		1.33 ± 0.01		
in vivo s.c.		1.23 ± 0.01			
predicted LT		1.3 ns			

рН-ОН	LT [ns]				
	pH 7.5		рН 5.5		
PBS	0.81 ± 0.01		0.83 ± 0.00		
PBS/BSA 5%	1.00 ± 0.01		1.55 ± 0.02		
in vivo s.c.		1.14 ± 0.02			
predicted LT		1.1 ns			

Alexa-Her	LT [ns]					
DP 1.3	pH 7.5		рН 5.5			
PBS	1.43 ± 0.03		1.44 ± 0.03			
PBS/BSA 5%	1.53 ± 0.02		1.63 ± 0.07			
in vivo s.c.		1.36 ± 0.01				
predicted LT		1.5 ns				

Alexa-OH	LT [ns]			
	pH 7.5		рН 5.5	
PBS	1.17 ± 0.00		1.14 ± 0.00	
PBS/BSA 5%	1.20 ± 0.02		1.57 ± 0.02	
in vivo s.c.		1.27 ± 0.02		
predicted LT		1.3 ns		

Taken together, the influences of a changing environment on the probes' LTs are relatively small. However, the LTs of all probes show a tendency to increase in the presence of proteins. This increase is most prominent for the free dyes measured in BSA-containing solution with a low pH. Furthermore, probes' LTs become longer for both dyes after their conjugation to Herceptin.

3.2.3. pH-sensitive conjugates are functional in vitro

In order to test the functionality of the pH-sensitive probes of low and high DP ratios *in vitro*, fluorescence microscopy with HER2-positive (KPL-4) and HER2-negative (MDA-MB-231) breast cancer cells was performed after incubation with different probes. Herceptin-conjugates are known to internalize and localize in the acidic endosomes and lysosomes within 8 h of incubation at 37 °C^{7,38,92} whereas at 4 °C internalization is strongly reduced⁵. Therefore, cells were incubated with Herceptin-conjugates for 8 h at 37 °C and at 4 °C as a control, in order to measure only fluorescence of membrane bound probes, when located in an approximately neutral pH.

The pH-sensitive probe, pH-Her DP 1.6 shows fluorescence only after receptormediated internalization into KPL-4 cells (Figure 16A) whereas no signals are detected of membrane bound pH-Her (Figure 16B). In contrast, the always-on probe Alexa-Her DP 1.3, shows fluorescence after internalization (Figure 16C) as well as when bound to the cell membrane (Figure 16C,D). No fluorescence signals are detected on HER2-negative MDA-MB-231 cells after incubation with all probes (Figure



16E-H) at 37 °C and 4 °C showing that no unspecific binding and/or internalization occurred.

Figure 16. (legend see next page)

Figure 16. *In vitro* fluorescence microscopy demonstrates functionality of pH-sensitive probes with a low DP ratio

Cells grown on culture slides were examined under a fluorescence microscope with an x 40 objective. For this purpose breast cancer cells were seeded on chambered coverglasses and incubated for 8 h with the fluorescent probes pH-Her or Alexa-Her (5 µg IgG) with a low DP ratio. On the left, counter stain of cell nuclei with Hoechst 33342, in the middle, probe-derived signals, and on the right, merged images of the cell nuclei (blue) and the probe (red) are illustrated. For detection of pH-Her an exposure time of 1500 ms and for detection of Alexa-Her an exposure time of 200 ms was used. **A.** When incubated with KPL-4 cells at 37 °C, pH-Her DP 1.6 shows fluorescence only after receptor-mediated internalization (green arrow). **B.** No signals from the membrane-bound probes are observed, as can be seen by incubation of cells at 4 °C. **C,D.** In contrast, Alexa-Her DP 1.3 shows fluorescence from the internalized probe (green arrow) but also membrane-derived fluorescence after 8 h of incubation at (**C**) 37 °C and at (**D**) 4 °C (yellow arrow). **E-H.** No fluorescence signals can be detected after 8 h incubation of MDA-MB-231 cells with pH-Her at (**E**) 37 °C or (**F**) 4 °C as well as after incubation with Alexa-Her at (**G**) 37 °C and (**H**) 4 °C. Representative images of three independently performed experiments are presented. Bars represent 50 µm.

Incubation of breast cancer cells with pH-Her and Alexa-Her conjugates of high DP ratios (Figure 17) revealed similar results as the cell experiments performed with probes with low DP ratios (Figure 16). In KPL-4 cells pH-Her DP 5.0 shows fluorescence only after internalization (Figure 17A) but not when bound to the membrane (Figure 17B) whereas fluorescence of Alexa-Her DP 3.4 is independent on internalization (Figure 17C,D). MDA-MB-231 cells show no fluorescence signals upon incubation with probes with high DP ratios (Figure 17E-H).

In order to prove that fluorescence signals on the KPL-4 cells after incubation with pH-Her and Alexa-Her conjugates (Figure 16 and 17) do derive from specifically membrane-bound or internalized Herceptin, control experiments with the free fluorophores were carried out. No fluorescence signals are detected on HER2-positive KPL-4 cells after incubation with pH-OH neither at 37 °C nor at 4 °C (Figure 18A,B) or with Alexa-OH at 4 °C or at 37 °C (Figure 18C,D). MDA-MB-231 cells also show no detectable fluorescence when incubated with pH-OH or Alexa-OH (Figure 18E-H).



Figure 17. (legend see next page)

Figure 17. *In vitro* fluorescence microscopy demonstrates functionality of pH-sensitive probes with a high DP ratio

Cells grown on culture slides were examined under a fluorescence microscope with an x 40 objective. For this purpose cells were incubated for 8 h with pH-Her or Alexa-Her fluorescent probes (5 µg IgG) with a high DP ratio. On the left, counter stain of cell nuclei with Hoechst 33342, in the middle, probe-derived signals, and on the right, merged images of the cell nuclei (blue) and the probe (red) are illustrated. For detection of pH-Her an exposure time of 1000 ms and for detection of Alexa-Her an exposure time of 1000 ms was used. **A.** When incubated with KPL-4 cells at 37 °C, pH-Her DP 5.0 shows fluorescence only after receptor-mediated internalization (green arrow). **B.** No signals from the membrane-bound probes are observed at 4 °C. **C,D.** Alexa-Her DP 3.4 shows fluorescence from the internalized probe (green arrow) but also membrane-derived signals after 8 h of incubation at (**C**) 37 °C and at (**D**) 4 °C (yellow arrow). **E-H.** No fluorescence signals can be detected after 8 h incubation of MDA-MB-231 cells with pH-Her at (**E**) 37 °C or (**F**) 4 °C as well as after incubation with Alexa-Her at (**G**) 37 °C and (**H**) 4 °C. Representative results of three independently performed experiments are shown. Bars represent 50 µm.



Figure 18. *In vitro* fluorescence microscopy reveals no unspecific binding of free fluorescence dyes

Cells grown on culture slides were examined under a fluorescence microscope at with an x 40 objective. For this purpose cells were incubated for 8 h with 0.2 nmol of pH-OH or Alexa-OH. On the left, counter stain of cell nuclei with Hoechst 33342, in the middle, probe-derived signals, and on the right, merged images of the cell nuclei (blue) and the probe (red) are illustrated. For pH-OH an exposure time of 1500 ms and for Alexa-OH an exposure time of 200 ms was used, according to the exposure times used for Herceptin-conjugates with low DP ratios (Figure 16). **A.-H.** No fluorescence can be detected on KPL-4 cells after 8 h of incubation with pH-OH at (**A**) 37 °C and (**B**) 4 °C or with Alexa-OH at (**C**) 37 °C and (**D**) 4 °C. Incubation of MDA-MB-231 cells with pH-OH at (**E**) 37 °C or (**F**) 4 °C as well as with Alexa-OH at (**G**) 37 °C and (**H**) 4 °C also does not result in the detection of fluorescence signals. Representative results of three independently performed experiments are shown. Bars represent 50 µm.

The functionality of pH-Her was confirmed by showing pH-Her-derived fluorescence only after internalization in HER2-positive breast cancer cells presumably due to localization in acidic cell compartments. In contrast to that, fluorescence intensity of the always-on control probes, Alexa-Her was independent on pH and internalization.

3.2.4. Improved detection sensitivity of tumors by the combined use of pH-sensitive probes and subtraction of autofluorescence

pH-probes were tested for their ability to improve tumor-detection sensitivity in tumor-bearing mice. A further aim of this study was to eliminate autofluorescence signals deriving from the mouse itself, hampering the detection sensitivity of specific, probe-derived signals. Consequently, imaging data were processed by the use of two different techniques for background suppression, subtraction of autofluorescence and LT-gated imaging.

For this purpose, KPL-4 and MDA-MB-231 breast tumor-bearing nude mice were imaged before (prescan) and at different times up to 72 h after *i.v.* injection of 25 μ g IgG of the respective probe in the Optix MX2.

3.2.4.1. Background suppression via subtraction of autofluorescence in combination with pH-Her improves tumor-detection sensitivity

In order to eliminate autofluorescence of mice and thus to only illustrate signals deriving from the injected probes, subtraction of autofluorescence was performed. For this purpose, the average fluorescence intensity over the tumor area of the untreated mouse (prescan) was subtracted from all scans of the same mouse. In Figure 19, representative fluorescence intensity maps of KPL-4 tumor-bearing mice before and 1 to 72 h after injection of the probes are demonstrated. After i.v. application of pH-Her DP 1.6 (Figure 19A; n = 3) fluorescence in the tumor (white circle, shown at 24 h) can be clearly detected. The background fluorescence (white circle, shown at 24 h) is low and comparable to the fluorescence of the prescan, with exception of some signals deriving from the gastrointestinal (GI) tract of the mouse. In contrast, after injection of Alexa-Her DP 1.3 (Figure 19B; n = 3), fluorescence in the tumor (white circle, shown at 24 h) is also clearly visible but background signals (white circle, shown at 24 h) are high compared to signals of the prescan. Note also, that fluorescence intensities after injection of Alexa-Her in mice (Figure 19B) are much higher compared to signals after injection of pH-Her (Figure 19A) as indicated by the different scale bars in the figure.



Figure 19. Intensity maps after subtraction of autofluorescence after probe application *in vivo*

KPL-4 breast tumor-bearing mice were imaged before and at different times up to 72 h after probe injection (25 μ g IgG) in the Optix MX2 in a ventral position. Representative examples (n = 3) of fluorescence intensity images after subtraction of the mean autofluorescence, calculated over the tumor area of the respective untreated mouse (prescan) are shown. **A.** Images of a mouse receiving pH-Her DP 1.6 are illustrated. Probe-derived signals are mainly detected in the tumor. **B.** Images of a mouse receiving Alexa-Her DP 1.3 are illustrated. Probe-derived signals are detected in the tumor and in the background. Background- and tumor area are indicated with white circles. Note, that fluorescence intensities in B. are much higher than in A., as indicated by the different scale bars.

In summary, use of the pH-sensitive probe, pH-Her at a low DP ratio in combination with subtraction of autofluorescence resulted in elimination of major parts of background fluorescence in scans of tumor-bearing mice. In contrast, the intensity of the background strongly increased after injection of the always-on probe, Alexa-Her and could not be eliminated by subtraction of autofluorescence.

Contrast to noise ratios (CNRs) were calculated to quantify signal-detection sensitivity of the tumor in *in vivo* scans after background subtraction. The CNR provides information about the level of signal difference in the tumor and background and the affection of the signal by the background noise (see 2.2.8.4). CNRs were calculated in scans of KPL-4 tumor-bearing mice after treatment with Herceptin-conjugates. As controls, CNRs were also determined in scans of MDA-MB-231 tumor-bearing mice injected with Herceptin-conjugates and of KPL-4 tumor-bearing mice receiving the free fluorophores. As shown in Figure 20, *i.v.* injection of pH-Her DP 1.6 (Figure 20A) in HER2-positive KPL-4 tumor-bearing mice (n = 5) resulted in an continuous increase of CNRs within 24 h to a ratio of 171 that stays comparable high

up to 72 h. In contrast, Alexa-Her DP 1.3 (Figure 20B; n = 5) shows an increase in CNRs in these mice over time but only to a maximum CNR of 82 after 72 h. The average CNR over all times in KPL-4 mice treated with pH-Her DP 1.6 was about 2.5-fold higher than for mice treated with Alexa-Her.

Control experiments performed in HER2-negative MDA-MB-231 tumor-bearing mice injected with pH-Her DP 1.6 (Figure 20A; n = 3) or Alexa-Her DP 1.3 (Figure 20B; n = 3) revealed much lower CNRs, with an average of about 18, that is 9-times lower than in KPL-4 tumor-bearing mice treated with pH-Her. In contrast to that, for the same times, the average CNR after application of Alexa-Her DP 1.3 is only 4-times higher in KPL-4 tumor-bearing mice compared to MDA-MB-231 tumor-bearing mice. This illustrates that the differences of signals deriving from HER2-positive- and HER2-negative tumors can be enhanced by the use of the pH-sensitive probe.



Figure 20. Improved tumor-detection sensitivity *in vivo* by use of a pH-sensitive tumor-specific probe at a low DP ratio

A,B. CNRs calculated from *in vivo* intensity scans after subtraction of autofluorescence are shown. For calculation, the average intensity of the background was measured in the area over the lung. CNRs were calculated in scans of KPL-4 tumor-bearing mice before (0 h) and 1, 2, 4 (n = 3), 24, 28, and 72 h (n = 5) after injection of (**A**) pH-Her DP 1.6 or (**B**) Alexa-Her DP 1.3. As controls, CNRs were also calculated from scans of KPL-4 tumor-bearing mice 0 to 72 h after injection of the free fluorescence dyes (0.8 nmol; n = 3), (**A**) pH-OH and (**B**) Alexa-OH as well as from scans of MDA-MB-231 tumor-bearing mice 0, 24, 48, and 72 h (n = 3) after injection of (**A**) pH-Her DP 1.6 or (**B**) Alexa-Her DP 1.3. Standard deviations are indicated as black bars.

Further control experiments in KPL-4 tumor-bearing mice revealed that after injection of the free dyes, CNRs also are low after 24 to 72 h with a maximum ratio for pH-OH of 21 (Figure 20A; n = 3) and of 20 for Alexa-OH (Figure 20B; n = 3). These findings further suggest that signals after application of both Herceptin-conjugates, pH-Her and Alexa-Her to mice with KPL-4 tumors are deriving from specific interaction of the antibody with tumor cells, as here much higher CNRs were

calculated. Moreover, the fact that CNRs calculated for mice with MDA-MB-231 tumors after injection of pH-Her and Alexa-Her are comparable to the CNRs in mice that received free fluorophores, indicates that tumor signals in MDA-MB-231 tumor-bearing mice are deriving from unspecific accumulation of the Herceptin probe.

Surprisingly, for Alexa-OH administered to KPL-4 tumor-bearing mice, relatively high CNRs, of 72 were calculated 1 h after injection (Figure 20B). Furthermore, for both free dyes, pH-OH and Alexa-OH high intensity signals were observed over the bladder at early (1 and 4 h) times after probe injection, very likely deriving from a rapid excretion of the low molecular weight (~1000 Da) molecules via the urinary system.

In addition, to analyze in more detail whether the fluorescence probes unspecifically accumulate in any other organ than the tumor, KPL-4 and MDA-MB-231 tumorbearing mice were sacrificed 72 h after injection of pH-Her, Alexa-Her, or the free fluorescence dyes and organs were scanned *ex vivo*. When compared to organs of untreated mice (n = 4), no remarkable fluorescence signals could be detected in liver, kidneys, lung, heart, spleen, stomach, and gut (data not shown).

Taken together, subtraction of autofluorescence *in vivo* in combination with application of pH-Her at a low DP ratio led to an efficient elimination of background fluorescence in breast tumor-bearing mice and clearly improved tumor-detection sensitivity compared to injection of Alexa-Her.

3.2.4.2. Background suppression via LT-gating does not improve tumor-detection sensitivity *in vivo*

LT imaging *in vivo* not only illustrates specific signals deriving from the applied probe but can depict signals occurring from autofluorescence of the tissue and from unspecific fluorescence, e.g. from the food in the GI tract. First, in order to characterize this fluorescence from different origin, the LTs were analyzed in distinct areas of whole body scans of tumor-bearing mice prior to probe application. The representative histogram in Figure 21, on the left side shows the frequency of occurrence of different LTs measured in an untreated KPL-4 tumor-bearing mouse (n = 10). The most frequently detected LTs are shorter than 1 ns or longer than 2 ns. The corresponding color-coded LT map (Figure 21, right side) illustrates that the short LTs are deriving from autofluorescence in the background, which has a mean LT of 1.0 ns (n = 10) whereas the longer LTs correspond likely to the food in the GI tract, with a mean LT of 2.0 ns (n = 10). The LT of the autofluorescence in the tumor with a mean of 1.6 ns (n = 10) is higher than the LT in the background.



Figure 21. LTs of fluorescence signals in certain areas of untreated tumorbearing mice

On the left side, a representative histogram shows the frequency of occurrence of different LTs measured in an untreated KPL-4 tumorbearing mouse (n = 10) in the Optix MX2. The most frequently

measured LTs are shorter than 1 ns or longer than 2 ns. On the right side, the corresponding color-coded LT map of an untreated mouse is shown. The autofluorescence in the background has a mean LT of 1.0 ns (arrow; n = 10). In contrary, the GI tract (GI) has a mean LT of 2.0 ns (arrow; n = 10). The mean LT of the autofluorescence in the tumor is 1.6 ns (arrow).

With the aim to only illustrate fluorescence signals in vivo, that are deriving from the applied probe, intensity maps of mice were gated to the predicted LT of the applied probe \pm 0.2 ns in order to decouple them from unspecific background fluorescence. The predicted LT was calculated as the mean from the LTs of the respective probe determined in vitro under different conditions and after s.c. injection in mice (see section 3.2.2, Table 10). In Figure 22A, representative fluorescence intensity maps of a mouse bearing a KPL-4 tumor in the right abdominal mammary gland are illustrated at different times after receiving pH-Her DP 1.6 (n = 5). Only signals with LTs of 1.3 ns \pm 0.2 ns are shown, according to the predicted LT for pH-Her (Table 10). When comparing these LT-gated images to the ones after subtraction of background fluorescence of the same mice, in Figure 19A, some fluorescence is still visible in the background (white circle, shown at 24 h) whereas unspecific fluorescence from the GI tract of mice could be completely eliminated. Unexpectedly, the fluorescence signals of high intensity from pH-Her in the tumor (white circle, shown at 24 h) are also eliminated at most times (e.g. at 48 and 72 h). Figure 22B, shows representative fluorescence intensity maps of a mouse with a KPL-4 tumor receiving Alexa-Her DP 1.3 (n = 5). Only signals with LTs of 1.5 ns \pm 0.2 ns are shown, according to the predicted LT for Alexa-Her (Table 10). In contrary to LTgating after application of pH-Her, the major part of the background fluorescence is still visible (white circle, shown at 24 h) whereas fluorescence signals from Alexa-Her in the tumor (white circle, shown at 24h) are also partly eliminated at some time points (e.g. at 4 and 24 h; see also Figure 19B for comparison).





KPL-4 breast-tumor bearing mice were imaged before and from 1 to 72 h after probe injection (25 μ g IgG) in the Optix MX2 in a ventral position. Representative examples of LT-gated *in vivo* intensity maps are shown. The LT-gates were set according to the predicted LT of the respective probe ± 0.2 ns. **A**. A representative mouse receiving pH-Her DP 1.6 is illustrated (n = 3). After LT-gating (LT: 1.3 ± 0.2 ns) remaining background fluorescence and partial elimination of the tumor signals are observed at most time points (e.g. after 48h). **B**. A representative mouse receiving Alexa-Her DP 1.3 is illustrated (n = 3). After LT-gating (LT 1.5 ± 0.2 ns) a high amount of background fluorescence remains and part of the tumor signal is eliminated at some time points (e.g. after 4 and 48 h). Background- and tumor area are indicated with white circles at 24 h after probe injection. Note, that fluorescence intensities in B. are much higher than in A., as indicated by the different scale bars.

Here, LT-gating does not lead to an improved tumor-detection sensitivity *in vivo* as probe-derived signals in the tumor are eliminated, especially those of pH-Her. In addition, LT-gating does not reduce fluorescence in the background after injection of the always-on fluorescent probe, Alexa-Her.

The average LTs in the tumor at each time point were analyzed more precisely in order to evaluate why part of tumor signals was eliminated after LT-gating, although high intensities in the tumor were very likely deriving from the applied fluorescence probes. Figure 23 shows that the average LTs in the tumor area of KPL-4 tumor-bearing mice (M1 to M5), each treated with either pH-Her DP 1.6 (Figure 23A) or Alexa-Her DP 1.3 (Figure 23B) vary over time and in the different mice. Moreover, after application of pH-Her (Figure 23A) tumor LTs are clearly longer than the predicted LTs (black dashed line; Table 10) and are also partly outside the LT-range used for gating (1.3 \pm 0.2 ns, grey area) of pH-Her signals. The mean LT in the KPL-4 tumor area of all mice treated with pH-Her and at all times is 1.49 \pm 0.06 ns (n =

24). *Ex vivo* LT analyses of the excised tumors 72 h after injection of pH-Her reveal an even higher mean LT of 1.57 ± 0.13 ns (n = 5; data not shown). In contrary, the average LTs after injection of Alexa-Her (Figure 23B) are close to the predicted LT (black dashed line; Table 10) and are within the LT-gate (1.5 ± 0.2 ns, grey area) determined for Alexa-Her. Here, the mean LT in the tumor area over time of all mice is 1.48 ± 0.04 ns (n = 24). *Ex vivo* LT analyses of the tumors 72 h after injection of Alexa-Her also reveal a higher mean LT of 1.65 ± 0.05 ns (n = 5; data not shown). This demonstrates that for both Herceptin probes the LT does further increase when the tumor is measured *ex vivo* where its fluorescence signal is not overlayed with other signals deriving from mouse tissue.



Figure 23. Variations of LTs in the tumor area after probe application

In vivo average LTs in KPL-4 tumors of mice (M1 – M5) are shown. Tumor LTs were analyzed 1, 2, 4 (n = 3), 24, 48, and 72 h (n = 5) after *i.v.* injection of fluorescence probes. **A,B.** The predicted LT of the probe (determined in *in vitro* measurements and after *s.c.* injection) is indicated as a black dashed line and the LT-ranges used for LT-gating of *in vivo* intensity images are shown in grey. LTs after application of (**A**) pH-Her DP 1.6 and (**B**) Alexa-Her DP 1.3 are shown. LTs vary over time as well as between mice. Note, that tumor LTs after application of (**A**) pH-Her are partly longer than the predicted LT. Standard deviations are indicated as bars.

In summary, LT-analyses of tumor signals over time *in vivo* and *ex vivo* show that the predicted LT for pH-Her differs from the LT of the probe in the tumor. Therefore, a clear prediction of the *in vivo* LT of the probe in the KPL-4 tumor environment and thus an identification of probe-derived signals via LT-gating is not possible.

In order to examine the reliability of the LT calculations in the tumor, the LT goodness of fit was determined. This parameter determines the quality of the mean LTs calculated over the tumor area of the respective mice. A LT-calculation with a high accuracy is considered to have a LT goodness of fit between 0.8 and 1.2 (see also section 2.2.8.4 in Material and Methods). As can be seen in Figure 24, the LT goodness of fit does not significantly increase or decrease from 1 to 72 h after probe

application or between the mice. For tumor LTs after injection of pH-Her (Figure 24A) as well as of Alexa-Her (Figure 24B) the mean goodness of fit is 0.8 (n = 24) indicating that the LT calculations have a high accuracy.



Figure 24. Analysis of reliability of in vivo LT calculations

The LT goodness of fit of the *in vivo* LT calculation in the KPL-4 tumor area in mice (M1 – M5) at different times after injection of (**A**) pH-Her DP 1.6 and (**B**) Alexa-Her DP 1.3 shows slight variations over time as well as between the mice. The mean LT goodness of fit for the LT calculations of both probes is 0.8. Standard deviations are indicated as bars.

These results demonstrate that the *in vivo* LT calculations of the tumor area are reliable, even at early (1 h) and late (72 h) times after injection of the respective probe when the concentration of the fluorescent probe in the tumor might be low.

In summary, tumor-detection sensitivity *in vivo* was not enhanced after injection of Herceptin-conjugates in combination with LT-gating. As prediction of the pH-Her LT was not precise and background signals deriving from unbound Alexa-Her could not be efficiently eliminated, tumor-derived fluorescence was not effectively separated from background fluorescence.

3.2.5. pH-Her conjugates at high DP ratios are less suitable for *in vivo* imaging

Fluorescence probes with high DP ratios, pH-Her DP 5.0 and Alexa-Her DP 3.4 were injected in KPL-4 tumor-bearing mice (n = 3) to evaluate their tumor-detection sensitivity *in vivo* in comparison to fluorescence probes with low DP ratios. For this purpose, CNRs were calculated from *in vivo* fluorescence intensity scans of KPL-4 tumor-bearing mice receiving probes at high DP-ratios after background subtraction. As depicted in Figure 25, after injection of pH-Her DP 5.0, CNRs over time stay very low with a maximum ratio of 7 after 72 h (Figure 25A). In addition, during *in vivo* scans we could detect only weak tumor signals but interestingly strong fluorescence in the bladder of the mice especially at early times (1 to 4 h) after application of pH-

Her DP 5.0 (n = 3; data not shown). This was also observed after injection of pH-Her DP 6.4 (data not shown; n = 2) and after application of the free fluorescence dyes, pH-OH and Alexa-OH (see 3.2.4.1) in tumor-bearing mice.

CNRs after injection of Alexa-Her DP 3.4 in KPL-4 tumor-bearing mice increased over time to a maximum ratio of 47. However, the average CNR at all times was \sim 1.7 fold lower compared to the average CNRs in the same mice model after injection of the Alexa-Her conjugate with a low DP ratio (see 3.2.4.1, Figure 20B).

Human IgG has a hydrodynamic diameter (HD) of 10 nm and usually gets excreted via the liver⁷⁵. The observation of a rapid excretion of the pH-Her probe at high DP ratio via the kidneys and bladder which is usually the case for smaller molecules of 5 to 6 nm in HD⁷⁵ pointed already to a structural degradation of the Herceptin-conjugates into smaller fragments. As CNRs were low for Herceptin-conjugates with high DP ratios, when compared to CNRs of conjugates with low DP ratios, control studies on MDA-MB-231 tumor-bearing mice or with free fluorophores were not performed.



Figure 25. Fluorescence probes with high DP ratios reveal no improvement in tumor-detection sensitivity

CNRs were calculated from fluorescence intensity scans of KPL-4 tumor-bearing mice after background subtraction before (0 h) and 1 to 72 h (n = 3) after injection of probes (25 μ g IgG) at high DP ratios. CNRs after application of (**A**) pH-Her DP 5.0 and (**B**) Alexa-Her DP 3.4 are shown. For CNR-calculation the average intensity of the background was measured in the area of the lung. Standard deviations are indicated as black bars.

Taken together, the application of fluorescent probes with high DP ratios unexpectedly did not lead to an improved sensitivity of tumor detection *in vivo*, when compared to probes with low DP ratios.

In the following it was assessed, whether pH-Her at high DP ratios might have been decomposed to nonfunctional fragments at the time of *in vivo* application in tumorbaring mice, in comparison to their application *in vitro* on cells (see section 3.2.3, Figure 16 and 17).

For this purpose, first fluorescence probes were separated according to their size by SDS-PAGE under non-reducing conditions as shown in Figure 26. As a control, unlabeled Herceptin in its reduced and unreduced form was used. In order to separate the heavy (\sim 50 kDa) and the light chain (\sim 25 kDa) of the antibody, Herceptin was incubated with reducing agent and heated at 70 °C for 10 min (line A). Herceptin in its unreduced form has a size of ~146 kDa (line B). Compared to the size of this antibody, the CypHer5E- and Alexa Fluor 647 fluorophores are small (750 to 1200 Da) so that the size of the fluorescence conjugates does not significantly differ from that of the free IgG. Alexa-Her DP 1.3 (line C) and DP 3.4 (line D) as well as pH-Her DP 1.6 (line E) reveal a comparable strong band with the size of unreduced Herceptin. pH-Her DP 1.6 additionally shows two bands with the size of 250 kDa and larger which are likely to be deriving from agglomerates of the conjugates. In contrast, pH-Her DP 5.0 (line F) and DP 6.4 (line G), both show no bands at the size of unreduced Herceptin and only some very weak bands between 25 and 45 kDa, indicating that pH-Her at high DP ratios has been decomposed into smaller fragments.



Figure 26. Detection of conjugates in a nonreducing protein gel

Herceptin-dye-conjugates (5 IgG) were separated uа according to their size via SDS-PAGE under nondenaturating conditions and stained with Coomassie (n =1). (A) As a control, unlabeled Herceptin was treated with reducing agent and heated to 70 °C for 10 min. Heavy and light chains of the antibody can be seen. (B) Unlabeled Herceptin in its non-reduced form. (C) Alexa-Her DP 1.3 and (**D**) DP 3.4 as well as (**E**)

pH-Her DP 1.6 have a size comparable to the unlabeled Herceptin. (\mathbf{F}) pH-Her DP 5.0 and (\mathbf{G}) DP 6.4 show no bands at the size of Herceptin.

To verify whether the Herceptin-conjugates were destructed into smaller fragments, the protein content of the probes was determined in a BCA assay (n = 2; data not shown) revealing that pH-Her DP 5.0 (line F) and pH-Her DP 6.4 (line G) had an approximately 3- and 5-times lower protein content, respectively when compared to the other probes (line B-E). Via this method, proteins decomposed to very small peptides containing three or more aminoacid residues are still detectable. Therefore, it can be assumed that the low functionality of the pH-Her conjugates with high DP ratios *in vivo* might not solely been caused by a degradation of the antibody conjugates into smaller fragments, detected *in vivo* in the bladder but also by a partial precipitation, presumably caused by probe aggregation.

In general, fluorescence conjugates with low DP ratios tend to be more stable over time when compared to conjugates with high DP ratios.

3.3. Evaluation of Itrybe-loaded NPs as *in vivo* NIRF imaging agents

Nanosized carrier systems are promising imaging tools as they exhibit several advantages, such as loading with multiple fluorophores resulting in highly intense and stable fluorescent labels. Furthermore, they enable surface conjugation with multiple target-specific ligands, which compared to the commonly used single antibody-fluorophore units may be especially favorable for *in vivo* imaging applications.

In this context, NPs with a diameter of 100 nm, 25 nm, and 15 nm, loaded with the NIR fluorophore Itrybe were evaluated for their capacity to be used for *in vivo* monitoring of tumors. These particles were either unmodified, bearing NH₂ groups or PEG chains of 1.5 or 10 kDa, linked to the NPs via the NH₂ groups. In addition, Herceptin-modified (Her) NPs were examined, at which the antibody was conjugated to the particle surface via the PEG chains (dye-loading and surface conjugation were performed by Thomas Behnke, BAM I.5, Berlin, see section 2.2.3.3).

3.3.1. Characterization of Itrybe-loaded NPs

Fluorescence excitation and emission spectra of different Itrybe-loaded NPs were recorded, to analyze whether the different sizes (100, 25, and 15 nm) and surface modifications (NH_2 , PEG, Her) influence their spectral behavior.

Figure 27A-C illustrates no remarkable spectral differences of NPs with the same size but different surface-modifications. However, the excitation and emission spectra of the 100 nm particles (blue line) are slightly right shifted compared to the spectra of the smaller sized NPs (25 nm and 15 nm), as exemplarily shown for NH₂-modified probes in Figure 27. All spectra appear noisy which might be caused by a decreased emission intensity due to scattering effects of the NPs itself. Scattering of the NPs can lead to a size-dependent decrease of excitation intensity and to interreflections of emitted photons and subsequently to a stronger reabsorption by the dye. This may cause the slight right shifts in the spectra of the 100 nm NPs (Figure 1D) compared to the smaller sized 15 nm and 25 nm NPs. With an increase in NP size, scattering and reabsorption effects increase possibly leading to some measurement artifacts.





Representative normalized fluorescence spectra of Itrybe-loaded NPs of different sizes and surface modifications were measured in a spectrofluorometer. The excitation spectra range from ~450 to 750 nm. The emission spectra range from ~650 to 850 nm. Fluorescence spectra of (**A**) 100 nm (n = 3), (**B**) 25 nm (n = 3), and (**C**) 15 nm (n = 2) NPs are illustrated. No remarkable spectral changes are observed for the different surface-modifications of NPs (NH₂, PEG, and Her). **D.** Spectra of all three sizes of NPs with NH₂ groups are illustrated. Note that, excitation and emission spectra of 100 nm particles (blue line) are slightly right shifted compared to the 25 nm and 15 nm NPs (green and red line).

Taken together, spectral differences of NPs of the same size but with various surface modifications are only small showing that the surface moiety has no relevant impact on the detection of Itrybe-derived fluorescence using the same excitation and emission settings. As fluorescence intensities from differently sized particles were not directly compared, the slight spectral variations of NPs of different sizes were not of relevance for the results of the study.

The encapsulation of the NIR fluorescence dye, Itrybe in NPs provides a cover of the dye towards the environment. Common factors influencing the fluorescence properties of fluorophores are, among others, proteins and ions^{2,20,108}. Therefore, the influences of changes in the environment on Itrybe loaded in NPs were analyzed exemplarily for the different sizes of the NH₂-modified particles (n = 2, for each size and each condition). As depicted in Figure 28, a decrease in pH from 7.5 to 5.5 has

no influence on fluorescence properties of the dye enclosed in NPs of all sizes; however a slight (1.3-fold) increase was observed for the 100 and 25 nm NPs when 5 % BSA is added to the measurement solution (PBS), independent on pH. In contrary, as can be seen in Figure 28C, Itrybe loaded in the smallest, 15 nm particles shows a stronger, 5-fold increase in fluorescence in the presence of 5 % BSA at a neutral pH. At an acidic pH of 5.5 the fluorescence is still 4-fold higher compared to fluorescence measured in PBS.





The relative increase in fluorescence intensity of Itrybe-loaded $NH_2 NPs$ of different sizes in dependence on environmental factors was analyzed in the Optix MX2. For this purpose, NPs were measured in PBS with a pH of 7.5 and 5.5 as well as in the presence of proteins (5 % BSA). For relativization, the lowest mean intensity for each NP size was set to 1. **A.** 100 and **B.** 25 nm particles show no influence on fluorescence intensity with a change in pH but a slight (1.3-fold) increase in fluorescence in the presence of proteins. **C.** In contrary, 15 nm NPs revealed a high (5.3-fold) increase in fluorescence in the presence of proteins at a pH 7.5 that is slightly lower (4-fold) at a pH of 5.5. The mean of two measurements is illustrated for each probe and each condition.

In summary, the influence of proteins on Itrybe-derived fluorescence are relatively small in 100 nm and 25 nm particles but increase in the small, 15 nm NPs. Moreover, the pH shows no strong influence on fluorescence of Itrybe.

3.3.2. Functionality of bioconjugated NPs

Itrybe-loaded NPs were designed to specifically bind to tumor cells. For this purpose, Herceptin, which recognizes HER2, was bioconjugated to the 100, 25, and 15 nm NPs. Immunoassays were performed to evaluate whether 1.) bioconjugation of NPs was successful and 2.) Herceptin-conjugated NPs bind to the HER2 antigen, as well 3.) NH_2 -modified and PEGylated particles unspecifically bind to HER2 antigen-coated surfaces.

Figure 29 shows the ratio of fluorescence detected after incubation of HER2-coated and -uncoated wells with 15 nm particles. A ratio of 1.0 is determined for the control experiment in which coated and uncoated wells were incubated with TBS only.

Particles with NH₂ groups and PEG 10 kDa chains show no specific binding to antigencoated wells. To achieve a high conjugation efficiency of Herceptin to the relatively small 15 nm NPs und thus a specific binding to HER2, a 5-fold amount of Herceptin (Her x5; 2.4 nmol/mg PS; see section 2.2.3.3, Table 2) was used for coupling procedure. After Herceptin-conjugation (Her x5), NPs with 10 and 1.5 kDa PEG chains show a slightly increased binding ratio of 1.1 and 1.2, respectively. To further increase specific binding of 15 nm NPs by Herceptin, in a next step the amount of PEG for coupling procedure was reduced to 1/10 (PEG 1.5 kDa 1/10; 50 nmol/mg PS; see section 2.2.3.3, Table 2) of the original amount. This was done, as a too high density of PEG chains might probably lead to steric hindrances during antibody conjugation. In addition, a "blockade" of the already conjugated antibodies on the particle surface can occur, thereby hampering interaction of the antibody with its target. However, when reducing the PEG amount for coupling procedure, after conjugation to Herceptin (Her x5) again no specific binding to the HER2 antigen is observed (ratio 1.0).



Figure 29. Binding specificity of bioconjugated 15 nm NPs

Signals from immunoassays with 15 nm NPs were measured in the Odyssey infrared imaging system. The ratio of fluorescence detected after incubation with 0.2 g/l NPs on HER2 antigen-coated and -uncoated wells, is illustrated. The control experiments, after incubation with only TBS or NH₂

and PEG 10 kDa NPs all show a ratio of 1.0. After Herceptin conjugation (Her x5), NPs with 10 and 1.5 kDa PEG chains have a slightly increased ratio of 1.1 and 1.2, respectively. No binding of Herceptin-conjugated NPs (Her x5) is observed for NPs with PEG 1.5 kDa chains when lower amounts of PEG (1/10) were used for coupling (ratio of 1.0). The number of independently done experiments is listed under the probes. Standard errors are illustrated as black bars. Each experiment was performed in triplicate.

Further it was tested whether bioconjugation of 25 nm particles with Herceptin was successful and thus specific binding of these probes to HER2 occurs. As can be seen in Figure 30A, the controls, NH₂- or PEGylated NPs (PEG 1.5 kDa and PEG 10 kDa) do not specifically bind to antigen-coated wells. A weak binding to antigen-coated wells is observed after coupling of Herceptin to 25 nm particles with PEG 1.5 kDa chains (ratio of 1.7) that cannot be further enhanced when the amount of antibody was increased 5-fold for coupling procedure (Her x5; 2.4 nmol/mg PS, see section

2.2.3.3, Table 2; ratio of 1.7). The use of PEG 10 kDa chains leads to a reduced antigen binding after Herceptin conjugation, with a ratio of 1.4. The bioconjugated NPs with the highest ratio and therefore best HER2 binding, PEG 1.5 kDa-Her and PEG 1.5 kDa-Her x5 were selected for further analysis. Figure 30B, illustrates that binding of these probes to HER2-coated wells is saturated at concentrations of 1 g/l whereas binding to uncoated wells further increases up to 2 g/l (n = 1). The ratios of binding to coated and uncoated wells for both 25 nm-Her probes (Figure 30C) are comparable high at 0.2 to 1 g/l (ratios between 1.6 and 1.8). Therefore, a concentration of 0.2 g/l was chosen for further analysis of binding specificity of NPs which was evaluated for the 25 nm PEG 1.5 kDa-Her x5 probe.



Figure 30. Binding specificity of bioconjugated 25 nm NPs

Signals from immunoassays with 25 nm NPs were measured in the Odyssey infrared imaging system. **A.** The ratios of signals from HER2-coated and -uncoated wells incubated with 0.2 g/l of NPs are illustrated. The control experiments, after incubation with only TBS or NH₂, PEG 1.5 kDa, and 10 kDa NPs all show low ratios (of 1.1 and 1.0). Slightly higher ratios are observed for the Herceptin-conjugated NPs PEG 1.5 kDa-Her and -Her x5 (ratio of 1.7) and PEG 10 kDa-Her (ratio of 1.4). The number of independently done experiments is listed under the probes. **B.** Relative intensities of 25 nm PEG 1.5 kDa-Her and -Her x5 NPs incubated on HER2-coated and -uncoated wells at concentrations of 0.02 to 2 g/l are shown. **C.** The respective ratios of signals from coated and uncoated wells for both probes are illustrated. **D.** Relative intensities of 25 nm PEG 1.5 kDa-Her and -uncoated wells at concentrations of 0.2 g/l. Free Herceptin was added in increasing amounts (0.005 to 1 μ g) to the wells. Black bars indicate standard errors in **A,C.** and standard deviations in **B,D.** Each experiment was performed in triplicate.

Figure 30D shows, that binding of the probe to HER2 is specifically occurring via interaction of Herceptin and the antigen as this binding can be "blocked" by addition of increasing amounts of free Herceptin (n = 2). After co-incubation with 0.5 µg Herceptin per well, binding of 25 nm PEG 1.5 kDa-Her x5 NPs can be completely reduced to the level of unspecific binding to uncoated wells.

In Figure 31A, the ratios of signals from antigen-coated and -uncoated wells after incubation with 100 nm particles are illustrated. NPs without bioconjugation show no binding to the antigen (NH_2 , ratio of 1.0 and PEG 1.5 kDa, ratio of 1.1). In contrary, the Herceptin-conjugated NPs with PEG 1.5 kDa chains show a 4.7-fold increased ratio which is even slightly higher (ratio of 4.8) when the double amount of Herceptin was used for bioconjugation (Her x2; 0.96 nmol/mg PS, see section 2.2.3.3, Table 2). Antigen binding is reduced when PEG 10 kDa chains were used as a linker for Herceptin to the NPs, which is in agreement with the results observed for the 25 nm NPs (see Figure 30A). Here, a ratio of 2.9 and 2.7 was calculated for two independently produced 100 nm-PEG 10 kDa-Her probes. This effect can be explained by the differences in number of Herceptin molecules per 100 nm particle (listed in section 2.2.3.3., Table 3): The 100-PEG 1.5 kDa-Her and -Her x2 probes bear ~70 and 100 Herceptin molecules on the surface of one particle, whereas the 100-PEG 10 kDa-Her probes bear only ~10 antibodies per NP. Therefore, the 100 nm-PEG 1.5 kDa-Her and -Her x2 NPs, showing the highest ratios, were selected for further studies. As illustrated in Figure 31B, the binding of PEG 1.5 kDa-Her to the antigen is saturated at 0.2 g/l whereas binding of PEG 1.5 kDa-Her x2 increases up to 1 g/l as does the unspecific binding for both probes to the uncoated wells (n = 1). Analysis of the ratio of signals from coated and uncoated wells (Figure 31C) reveals best ratios for the PEG 1.5 kDa-Her and -Her x2 probes at 0.1 g/l (ratio of 4.8 and 4.1) and 0.2 g/l (ratio of 4.6 and 4.2). Exemplarily, the binding specificity is shown for the PEG 1.5 kDa-Her NPs at 0.2 g/l in Figure 31D. Binding of NPs to HER2 can be decreased via addition of increasing amounts of free Herceptin (n = 2). After coincubation with 0.5 µg Herceptin per well the signal from bound NPs is comparable low to the unspecific signal deriving from uncoated wells.

In summary, the highest binding specificity to HER2 was achieved after Herceptinconjugation to the 100 nm NPs, whereas bioconjugated 25 and 15 nm NPs showed weak or almost no binding to HER2, respectively. For the bioconjugated 25 nm as well as 100 nm particles stronger binding to HER2 was observed when short PEG chains (1.5 kDa) were linked to the NP-surface compared to longer PEG chains (10 kDa).





Signals from immunoassays with 100 nm NPs were measured in the Odyssey infrared imaging system. **A.** The ratios of signals from HER2-coated and -uncoated wells incubated with 0.2 g/l of NPs are illustrated. The control experiments, after incubation with only TBS or NH₂ and PEG 1.5 kDa NPs, all show low ratios (of 1.1 and 1.0). An increase in specific binding can be observed for 1.5 kDa PEG-Her and -Her x2 (ratio of 4.7 and 4.8, respectively). Use of 10 kDa PEG chains reduces binding of Herceptin-conjugated NPs to HER2 (ratio of 2.9 and 2.7 for two independently generated probes). The number of independently done experiments is listed under the respective probes. **B.** Relative intensities of 100 nm-1.5 kDa-Her and -Her x2 NPs incubated on coated and uncoated wells at concentrations of 0.02 to 1 g/l are shown. **C.** The respective ratios of signals from coated and uncoated wells for both probes are illustrated. **D.** Relative intensities of 100 nm-PEG 1.5 kDa-Her NPs (n = 2) incubated on HER2-coated and -uncoated wells at concentrations of 0.2 g/l. Free Herceptin was added in increasing amounts (0.005 to 1 μ g) to the wells. Black bars indicate standard errors in **A,C.** and standard deviations in **B,D.** Each experiment was performed in triplicate.

3.3.3. NPs show no influence on cell viability

As 100-PEG 1.5 kDa-Her and –Her x2 NPs showed the highest binding specificity to HER2, these probes were selected for further analyses of tumor cell binding specificity *in vitro* and *in vivo*.

Since, potentially cytotoxic effects of NPs, especially to non-cancerous cells had to be excluded, a cell viability assay was performed on 3T3 mouse fibroblasts. Fibroblasts were incubated for 24 h with 0.01 to 2 g/l 100 nm NPs and the amount of viable cells, proportional to the amount of formazan was measured by absorption at 450

nm. Figure 32 shows that 100 nm-NH₂ NPs (blue columns; n = 3) as well as Herceptin-conjugated NPs (PEG 1.5 kDa-Her, green columns, n = 2; PEG 1.5 kDa-Her x2, red columns, n = 1) had no effect on cell viability when compared to untreated cells (0 g/l, grey column, n = 3) with a confidence of 90 % (red lines). Control samples without cells, containing only medium (medium, grey column; n = 1) showed a low absorption, far outside the 90 % confidence interval of untreated 3T3 cells.



Figure 32. NPs do not decrease cell viability

3T3 mouse fibroblasts were incubated with 0.01 to 2 g/l of 100 nm NPs for 24 h and cell viability was analyzed via WST-1 assay and measurement of absorption at 450 nm. To exclude absorption effects by the Itrybe-loaded NPs, absorption was measured before addition of WST-1 and subtracted from absorption after WST-1 treatment. The mean values of the absorption by formazan are illustrated for untreated cells (0 g/l NPs; n = 3) and for medium only (grey column; n = 1) as well as for cells incubated with 100 nm NPs. Incubation with NPs is shown in blue columns for 100 nm-NH₂ (n = 3), in green columns for 100 nm-PEG 1.5 kDa-Her (n = 2), and in red columns for 100 nm-PEG 1.5 kDa-Her x2 (n = 1). All experiments were performed in triplicate. The 90 % confidence interval of untreated cells is depicted in red lines.

In summary, NPs at concentrations up to 2g/l incubated for 24 h do not affect cell viability of mouse fibroblasts.

3.3.4. Binding specificity of bioconjugated NPs to breast tumor cells *in vitro*

As 100-PEG 1.5 kDa-Her and –Her x2 NPs showed strong binding to the HER2 antigen in immunoassays (see section 3.3.2, Figure 31), they were analyzed for their binding capacity to KPL-4 and MDA-MB-231 tumor cells by fluorescence microscopy, in comparison to controls, the 100-NH₂ and -PEG 1.5 kDa NPs.

Specific binding of Herceptin-conjugated NPs to HER2-positive KPL-4 cells was observed, whereas the controls revealed also some unspecific binding. Figure 33 shows, that strongest fluorescence is detected from HER2-positive, KPL-4 cells after incubation with 100 nm-PEG 1.5 kDa-Her NPs (Her) whereas signals are low after incubation of the same cell type with the unspecific probe, 100 nm- PEG 1.5 kDa NPs (PEG). Incubation of both probes with HER2-negative, MDA-MB-231 cells also revealed comparable low fluorescence (n = 3). The differences of specific and unspecific binding to cells decreased with increasing incubation times: Whereas the differences of fluorescence of KPL-4 cells incubated with Her to the control samples (PEG on KPL-4 as well as Her and PEG on MDA-MB-231 cells) are relatively high after 30 min (Figure 33A) and 1 h (Figure 33B), unspecific uptake of the control samples in cells increased after 3 h (Figure 33C) and 5 h of incubation (Figure 33D).

The binding and/or uptake of 100 nm-PEG 1.5 kDa-Her NPs on KPL-4 cells over time is illustrated in Figure 34. Itrybe-derived signals continuously increased from 30 min to 5 h after incubation of the probe at 37 °C (Figure 34A; n = 3). At 4 °C incubation (Figure 34B; n = 1), no fluorescence can be detected on cells using the same exposure time (10 ms). Nevertheless, an increase of Itrybe-derived fluorescence over time is also detected at 4 °C with a higher exposure time of 20 ms (Figure 34C), however, cell binding and/or uptake of NPs is lower compared to incubation at 37 °C. The distribution of NPs on cells incubated at 37 °C (Figure 34A) and at 4 °C (Figure 34C) is relatively equal over time as no differences in location of NPs, e.g. predominantly membrane bound probe at early time points in comparison to internalized probe after longer incubation times, are observed. NPs seem to localize inside the cells after very short incubation times (Figure 33 and Figure 34).



Figure 33. Increased binding/uptake of Herceptin-conjugated NPs to HER2-positive cells *in vitro*

Cells were seeded on glass slides and incubated for 30 min, 1, 3, and 5 h at 37 °C with 1 g/l Itrybe-loaded 100 nm NPs and examined under a fluorescence microscope with an x 40 objective. For each box, on the left, counter stain of cell nuclei with Hoechst 33342 and in the middle probe-derived signals are illustrated. Merged images of the cell nuclei (blue) and the probe (red) can be seen on the right. The exposure time used for detection of Itrybe was 30 ms after 30 min, 10 ms after 1 and 3 h, and 4 ms after 5 h of incubation. **A,B.** After (**A**) 30 min and (**B**) 1 h of incubation strongest fluorescence is deriving from KPL-4 cells treated with 100 nm-PEG 1.5 kDa-Her (Her). The control probe, PEG 1.5 kDa (PEG) on KPL-4 cells as well as both probes on MDA-MB-231 cells reveal low fluorescence signals. **C,D.** When incubated (**C**) for 3 h and (**D**) 5 h with 100 nm NPs, still strongest fluorescence is deriving from 100 nm-Her on KPL-4 cells but signals from 100 nm-PEG on KPL-4 cells and from both probes on MDA-MB-231 cells are slightly increased, compared to A. and B. Representative results of three independently performed experiments are shown. Bars represent 50 µm.

Results



Figure 34. Increased binding/uptake of Herceptin-conjugated 100 nm NPs in HER2-positive cells over time at 37 and 4 $^{\circ}$ C

KPL-4 cells were incubated for 30 min, 1, 3, and 5 h at 37 °C and 4 °C with 1 g/l Itrybe-loaded 100 nm-PEG 1.5 kDa-Her NPs and examined under a fluorescence microscope with an x 40 objective. For each box, on the left, counter stain of cell nuclei with Hoechst 33342 and in the middle, probe-derived signals are illustrated. Merged images of the cell nuclei (blue) and the probe (red) can be seen on the right. **A.** Itrybederived fluorescence on cells, detected with a constant exposure time of 10 ms, continuously increases over time at 37 °C. Representative results of three independently done experiments are shown. **B.** No fluorescence signals on cells can be detected after incubation at 4 °C (n = 1) at the same exposure time as in A., of 10 ms. **C.** When using a higher exposure time of 20 ms, increased Itrybe-derived signals on cells can be detected over time. Bars represent 50 μ m. The binding specificity of Herceptin-conjugated NPs was further verified on KPL-4 and MDA-MB-231 tumor sections (two tumors of each type) that were incubated with different 100 nm probes. As illustrated in Figure 35, 100 nm-PEG 1.5 kDa-Her x2 NPs (n = 2) show strong binding to KPL-4 tumors (Figure 35A; n = 2), compared to the control probes, 100 nm-PEG 1.5 kDa and 100 nm-NH₂ (Figure 35B,C; n = 2).



Figure 35. *Ex vivo* fluorescence experiments demonstrate binding specificity of Herceptinconjugated NPs to HER2-positive tumors

Paraffin fixed KPL-4 and MDA-MB-231 tumor sections were incubated with 0.1 g/l Itrybe-loaded NPs and examined under a fluorescence microscope with a x 40 objective. For each box, on the left upper side, counter stain of cell nuclei with DAPI and on the left lower side, probe-derived signals are illustrated. Merged images of the cell nuclei (blue) and the probe (red) can be seen on the right. **A-C.** Representative results of two different KPL-4 tumors incubated with (**A**) 100 nm-PEG 1.5 kDa-Her x2 (Her; n = 2) show strong fluorescence in comparison to KPL-4 tumors incubated with (**B**) 100 nm-PEG 1.5 kDa (PEG; n = 1), or (**C**) 100 nm-NH₂ (NH₂; n = 2). **D-F.** Representative results of two different MDA-MB-231 tumors incubated with (**D**) 100 nm-PEG 1.5 kDa (PEG; n = 1), (**F**) 100 nm-NH₂ (NH₂; n = 1) reveal lower fluorescence compared to A. The exposure time for detection of NPs was 140 ms. Bars represent 50 µm.

Weak fluorescence could be observed in MDA-MB-231 tumors treated with 100 nm-PEG 1.5 kDa-Her x2 NPs (Figure 35D) that was slightly higher compared to both tumor types incubated with the control probes (Figure 35B-C,E-F).

Taken together, Herceptin-conjugated 100 nm NPs are functional on breast tumor cells, as they showed increased binding/uptake to HER2-positive KPL-4 cells and - tumors. In contrary, PEGylated or NH_2 -modified NPs showed lower binding to KPL-4 cells and -tumors as did all NP probes to HER2-negative, MDA-MB-231 cells and - tumors.

3.3.5. In vivo detection of Itrybe-loaded NPs

The suitability of fluorescence deriving from Itrybe-loaded NPs to be detected *in vivo* in tumor tissue and at other sites of the mice was investigated. For this purpose, KPL-4 and MDA-MB-231 tumor-bearing nude mice received *i.v.* injections of different Itrybe-loaded NPs and were imaged *in vivo* and *ex vivo* at diverse time points. Due to the broad absorption spectra of Itrybe-loaded NPs (see section 3.3.1, Figure 27), imaging was performed at two different λ_{ex} of 635 and 670 nm and fluorescence was detected at λ_{em} above 700 nm (n = 9).

Figure 36A shows representative in vivo images of a KPL-4 tumor-bearing nude mouse before and 1 to 24 h after *i.v.* injection of 400 µg 100 nm-PEG 1.5 kDa-Her at λ_{ex} 635 nm (n = 2). Autofluorescence (white circle) was assessed in prescans of mice before treatment. High fluorescence in the liver is seen shortly after NP-injection up to 24 h (white ellipse after 1 h). However, no tumor signals were detected within this time (white circle). The same mouse imaged at λ_{ex} 670 nm, is illustrated in Figure 36B. Here, autofluorescence is lower in the prescan but unspecific signals in the stomach (arrow) and gut (white ellipse) appear higher. NP-derived fluorescence in the liver was also observed but appears weaker than at λ_{ex} 635 nm (Figure 36A). This can be explained by the better excitability of Itrybe at λ_{ex} 635 nm, compared to λ_{ex} 670 nm (see Figure 27 in section 3.3.1). Ex vivo scans of excised organs of the same mouse 24 h after probe injection are depicted in Figure 36C,D. Here, the highest signals can be seen in the liver at λ_{ex} 635 nm (Figure 36C) as well as at λ_{ex} 670 nm (Figure 36D) compared to low signals at both wavelengths in the gut, stomach, kidneys, heart, spleen, lung, and tumor. In comparison to excitation at λ_{ex} 635 nm, Itrybe-derived fluorescence in the liver is weaker at λ_{ex} 670 nm but unspecific signals from the GI tract are slightly higher, which is in agreement with the in vivo measurements.





Fluorescence intensity scans taken by the Optix MX2 before (prescan) and after *i.v.* injection of 400 µg 100 nm-PEG 1.5 kDa-Her NPs in nude mice bearing KPL-4 tumors in the right abdominal mammary gland (n = 2). **A.** A representative mouse in a ventral position imaged before and 1, 4, and 24 h after probe application at λ_{ex} 635 nm and λ_{em} above 700 nm. Autofluorescence in the prescan over the lung area is illustrated with a white circle. No accumulation of NPs in the tumor (white circle, 24 h) but high liver signals (white ellipse, 1 h) are observed after probe application. **B.** The same mouse shown in A. imaged at λ_{ex} 670 nm and λ_{em} above 700 nm. In the prescan autofluorescence is lower compared to A. but unspecific signals from the stomach (arrow) and gut (white ellipse) are higher. **C.** *Ex vivo* scans of excised organs of the mouse shown in A. and B. at λ_{ex} 635 nm, 24 h after NP-injection confirm *in vivo* observations. **D.** The same organs seen in C. imaged at λ_{ex} of 670 nm.

In summary, Itrybe-loaded NPs can be excited at λ_{ex} 635 nm but also at λ_{ex} 670 nm and be detected well above λ_{em} 700 nm. Following injection of NPs, probe-derived fluorescence cannot be detected in the tumor, but strong signals are observed in the liver. Due to the higher fluorescence of Itrybe at λ_{ex} 635 nm making autofluorescence appear negligible, this excitation wavelength was used for all following *in vivo* and *ex vivo* scans performed in the Optix MX2.

In order to illustrate the brightness of Itrybe-loaded NPs and its detection limit for tumors *in vivo*, different concentrations (100 to 0.8 μ g) of 15 nm-PEG 10 kDa NPs were injected *s.c.* in mice at different sites (n = 2). 15 nm particles were selected due to their lowest fluorescence intensity per mg PS compared to other NPs (see Table 11). As illustrated in Figure 37, fluorescence intensities of NPs after *s.c.* injection (Figure 37A,B; white circles) are well detectable even at an amount of 1.6 μ g as NPs are still 4-times more fluorescent (Figure 37C; 1,200 NC) than the autofluorescence (Figure 37B,D; red circle; 327 NC). In comparison, fluorescence of 1.6 μ g of NPs injected in the MDA-MB-231 tumor (Figure 37C; 813 NC) is slightly weaker, as only 2.5-fold higher than the autofluorescence.

As a result, Itrybe-derived fluorescence can be detected in very low amounts *in vivo*, although fluorescence is lower within the tumor compared to *s.c.* injections near the surface of the mouse. Therefore, Itrybe-loaded NPs are suitable for *in vivo* tumor imaging in mice.


Figure 37. In vivo detection of Itrybe-derived fluorescence

Fluorescence intensity scans taken by the Optix MX2 directly after injection of different amounts of 15 nm-PEG 10 kDa NPs solved in 30 μ l NaCl. The areas of probe injections are indicated with white circles. **A.** Mouse in a dorsal view with *s.c.* injections of 100, 50, 25, and 12.5 μ g NPs (n = 2). **B.** Mouse in a dorsal view with *s.c.* injections of 6.3, 3.1, 1.6, and 0.8 μ g NPs (n = 2). Autofluorescence (red circle) was determined in an area outside the gut and stomach **C.** A MDA-MB-231 tumor-bearing mouse in a ventral position injected with 1.6 μ g NPs *i.t.* (red arrow) and *s.c.* (red arrow). Note that different intensity scales are shown in A-C. **D.** The log of the average intensity after *s.c.* injection of 100 to 0.8 μ g 15 nm NPs (A,B) and of the autofluorescence of the mouse (0.0 μ g NPs) is illustrated.

3.3.6. Biodistribution of NPs

In order to compare the *in vivo* behavior of NPs depending on different sizes and surface modifications, biodistribution studies were performed in healthy nude mice. As NPs seem to be captured in the liver rapidly after injection (see section 3.3.5, Figure 36), mice were analyzed at early time points, from 6 min to 4 h after injection of NH₂ and PEGylated NPs of different sizes (n = 28).

As illustrated in Figure 38, exemplarily for a mouse injected with 25 nm-PEG 10 kDa (n = 4), NPs accumulate in the liver (white ellipse) already 6 min after injection. Therefore, fluorescence is especially analyzed over the liver and compared to

fluorescence measured in the background of mice, representing the amount of circulating unbound NPs in the blood. An area over the hindlimb of mice (in a ventral position) was selected as a background region (white circle), since no unspecific accumulation of NPs is expected here. After injection of NPs, only a marginal increase in fluorescence is detected in the background (white circle) indicating a fast elimination of NPs from the blood pool.



Figure 38. Biodistribution of NPs in healthy mice over time

Fluorescence intensity maps of healthy, tumor free, female nude mice scanned *in vivo* in the Optix MX2 before (prescan) and 6 min to 4 h after injection of diverse NPs (n = 28). Exemplarily, a representative mouse before and after injection of 400 µg 25 nm-PEG 10 kDa NPs is illustrated (n = 4). High liver signals can be seen rapidly after probe application (white ellipse) whereas background fluorescence (white circle) remains low.

In Figure 39 average fluorescence intensities in liver (Figure 39A-C) and background regions (Figure 39D-F) of healthy nude mice treated with different NPs are illustrated before and 6 min to 4 h after treatment (n = 4; see also Figure 38). In order to eliminate signals deriving from autofluorescence of the mouse, the average fluorescence in the liver and background of the prescans of each mouse was subtracted from the respective regions after probe injection.

Note that intensities are not comparable between mice treated with NPs of varying sizes (15, 25, and 100 nm) since the fluorescence intensity of the injected probe varies dependent on the NP size. As illustrated in Table 11, per mg PS, the 25 nm NPs are 3-fold more fluorescent and the 15 nm particles are 5-fold less fluorescent compared to the 100 nm NPs.

Table 11. Fluorescence intensities of NPs of different sizes

Total fluorescence intensities of NP probes in 200 μ l NaCl were measured in the Optix MX2 prior to *i.v.* injection in mice. Measurements and calculations were performed for the PEGylated NPs (100 nm-PEG 1.5 kDa, 25 nm-PEG 10 kDa, and 15 nm-PEG 10 kDa).

probe	100 nm	25 nm	15 nm
injected amount [mg]	0.4	0.4	1
PI [NC]	1,900,000	5,630,000	790,000
relative* PI/mg	1.0	3.0	0.2

PI = Total fluorescence intensity of the injected probe

* relative to 100 nm NPs

As summarized in Figure 39A-C, NPs with increasing size accumulate slower in the liver: Fluorescence of 15 nm NPs already peaks ~6 min (Figure 39A), signals of 25 nm NPs peak ~30 min (Figure 39B), and fluorescence of 100 nm NPs peaks ~1 to 2 h after injection (Figure 39C). Furthermore, PEGylation of probes influences their accumulation in the liver, as 25 and 100 nm NPs demonstrate decreased liver signals after PEGylation whereas PEGylated 15 nm NPs show increased signals when compared to the NH₂-modified particles (Figure 39A-C). All probes show low fluorescence in the background with a maximal intensity within the first view minutes after injection of NPs that decreases over time (Figure 39D-F). Only the 100-PEG 1.5 kDa NPs (Figure 39F) revealed a well measurable and stable background signal over time (from 30 min to 4 h). In contrary, PEGylation of the 15 and 25 nm NPs had no impact on the prolongation of their blood circulation time (Figure 39D,E).

Healthy nude mice treated with NPs (see Figure 38 and 39) were sacrificed 4.5 h after injection and excised organs were scanned in the Optix MX2 *ex vivo*. Probederived fluorescence in the organs is displayed according to the presumable size of the injected NPs from the probe with the highest to the lowest diameter (100 nm-PEG 1.5 kDa > 100 nm-NH₂ > 25 nm-PEG 10 kDa > 25 nm-PEG 1.5 kDa > 25 nm-NH₂ > 15 nm-PEG 10 kDa > 15 nm-NH₂). To exclusively illustrate NP-derived signals, the autofluorescence from untreated organs (n = 4) was subtracted from the respective organs from mice that received NPs. To allow comparability of intensities from different probes, the average fluorescence intensity in particular organs is illustrated relative to the total intensity of all organs (liver, lung, heart, spleen, and kidneys) of mice treated with the same probe (n = 4).



Figure 39. NP distribution in liver and background of healthy mice in vivo over time

Average fluorescence intensities in liver (**A-C**) and background regions (**D-F**) of female healthy nude mice treated with distinct NPs are shown before and 6 min to 4 h after probe injection (n = 4). Mice received (A,D) 1 mg of 15 nm NPs, (B,E) 400 μ g of 25 nm, or (C,F) 400 μ g of 100 nm NPs. The average autofluorescence in the liver and background of the prescans of each mouse is subtracted from the respective regions after probe injection. Liver signals after injection of (**A**) 15 nm-NH₂ and -PEG 10 kDa, (**B**) 25 nm-NH₂, -PEG 1.5 kDa, and -10 kDa, as well as (**C**) 100 nm-NH₂ and -PEG 1.5 kDa are shown. Corresponding background signals after injection of (**D**) 15 nm-NH₂ and -PEG 10 kDa, (**E**) 25 nm-NH₂, -PEG 1.5 kDa, and -10 kDa, as well as (**F**) 100 nm-NH₂ and -PEG 1.5 kDa are illustrated. Mean values of 4 mice are shown. Standard deviations are indicated as black bars.

Figure 40A illustrates that all probes accumulate in the liver to a high extend after *i.v.* injection. For example, 25 nm NPs show highest liver uptake with up to 90 % (for the 25 nm-PEG 10 kDa probe) whereas lung, heart, spleen, and kidneys show comparable low fluorescence (Figure 40B). The second highest fluorescence after NP injection is seen in the spleen. Here, the 100 nm-NH₂ particles show the strongest fluorescence, with 9 % and a slightly decreasing tendency with a decrease in size of NPs. In contrary, highest fluorescence in the kidneys was found for the smallest (15 nm) NPs with 6 % whereas the larger probes show lower fluorescence of 2 to 3 %. Fluorescence in the heart was low for all probes (1.4 %).



Figure 40. Biodistribution of NPs by ex vivo analysis

Healthy nude mice injected with different NPs were sacrificed after 4.5 h and organs were scanned in the Optix MX2 (n = 4). Mice received 1 mg of 15 nm or 400 μ g of 25 or 100 nm NPs, with NH₂ or PEG (1.5 and 10 kDa) surface groups. Autofluorescence from the respective untreated organs (n = 4) is subtracted from the organs of treated mice. The relative fluorescence intensity of average signals over the respective organs is illustrated. This is calculated under the assumption that liver, lung, heart, spleen, and kidneys make up 100 % of the fluorescence. **A.** Relative fluorescence intensity in the liver is high compared to other organs. **B.** The spleen shows second highest uptake of NPs. In the kidneys, fluorescence increases after injection of 15 nm NPs.

Taken together, biodistribution studies in mice illustrate a rapid and high accumulation of all probes in the liver compared to the spleen, kidneys, lung, and heart. *Ex vivo*, 15 nm NPs showed the lowest liver accumulation but the highest signals in the kidney. With an increase in NP size, liver uptake and excretion seems to be slower over time. Blood circulation times for all NPs were low, showing peak intensity rapidly after injection, but are slightly enhanced after applying the PEGylated 100 nm NPs.

3.3.7. NPs accumulate in MDA-MB-231 breast tumors

In KPL-4 or MDA-MB-231 tumor-bearing mice injected with Herceptin-conjugated or PEGylated 15, 25, and 100 nm particles no NP-derived fluorescence could be detected in the tumors *in vivo* in whole-body scans up to 24 h (see Figure 36A,B; n = 19). Accumulation of NPs in a MDA-MB-231 tumor was also not observed *in vivo* up to 96 h after injection of 400 μ g 100 nm NPs with surface carboxyl-groups (-COOH) (data not shown).

Furthermore, tumors of mice treated with 1 mg of 15 nm NPs or with 400 μ g of 100 and 25 nm NPs were excised 24 h after injection and scanned *ex vivo* (n = 2). Figure

41A,B shows the "total fluorescence intensities in the tumor normalized to the tumor volume" (TIV) of KPL-4 tumors from mice treated with NPs. Compared to the average TIV of untreated mice (red line), no increase in fluorescence after NP-injection is observed. In contrary, TIVs in MDA-MB-231 tumors of mice treated with PEGylated NP show a clear increase with decreasing particle size compared to tumors of untreated mice (Figure 41C,D). Note, that untreated KPL-4 tumors (Figure 41A,B) have more than twice as high autofluorescence compared to MDA-MB-231 tumors (Figure 41C,D).

Ex vivo analysis of excised KPL-4 and MDA-MB-231 tumors of mice 24 h after *i.v.* injection of Herceptinconjugated and PEGylated 100, 25, and 15 nm NPs (n = 2) scanned in the Optix MX2. **A.** Total fluorescence intensities in the tumor normalized to the tumor volume (TIV) of KPL-4 tumors before (untreated, grey columns) and after treatment (white columns) with NPs are shown. From left to right: untreated, 100 nm-PEG 1.5 kDa-Her (100-Her), 100 nm-PEG 1.5 kDa (100-PEG), 25 nm-PEG 1.5 kDa-Her (25-Her), 25 nm-PEG 1.5 kDa (25-PEG), and 15 nm-PEG 10 kDa (15-PEG). The average TIV in untreated mice is indicated with a red line. No increase in fluorescence is observed in tumors after NP-treatment compared to tumors of untreated mice. **B.** Fluorescence intensity scans of KPL-4 tumors corresponding to A. are illustrated. **C.** TIV of MDA-MB-231 tumors before (untreated; grey columns) and after treatment (white columns) with NPs are shown. From left to right: untreated, 100-PEG, 25-PEG, and 15-PEG. The average TIV in untreated mice is indicated with a red line. Increased fluorescence is measured in tumors of mice treated with NPs. **D.** Fluorescence intensity scans of MDA-MB-231 tumors corresponding to C. are illustrated. In order to obtain only the fluorescence in the tumor deriving from injected NPs, the average TIV of untreated tumors was subtracted from the average TIV after probe injection; defined as the NP-derived TIV (Table 12). As the total fluorescence intensity of the injected probe (PI) is known for the 100, 25, and 15 nm PEGylated NPs, the NP-derived TIV, relative to the PI was calculated (NP-derived TIV/PI). Here, 15 nm-PEG 10 kDa NPs show a more than 4-fold higher NP-derived TIV/PI ratio than the 100 nm-PEG 1.5 kDa NPs (0.18 to 0.04) and a 9-fold higher ratio than the 25 nm-PEG 10 kDa NPs (0.18 to 0.02). This demonstrates that the 15 nm NPs show the highest fluorescence intensity in tumor tissue (normalized to the tumor volume) in relation to the intensity of the injected probe

Table 12. NP-derived signals in MDA-MB-231 tumors ex vivo

Total fluorescence intensities of NPs prior to injection in MDA-MB-231 tumor-bearing mice as well as of the excised tumors 24 h after injection were measured in the Optix MX2. The injected amounts of NPs were: 400 µg of 100 nm-PEG 1.5 kDa NPs (100-PEG), 400 µg of 25 nm-PEG 10 kDa NPs (25-PEG), and 1 mg of 15 nm-PEG 10 kDa NPs (15-PEG).

probe	100-PEG	25-PEG	15-PEG
NP-derived TIV [NC/cm ³]	79,251	125,695	143,754
PI [NC]	1,900,000	5,630,000	790,000
NP-derived TIV/PI [1/cm ³]	0.04	0.02	0.18

TIV = Total fluorescence intensity in the tumor normalized to the tumor volume

PI = Total fluorescence intensity of the injected probe

In summary, *ex vivo* analysis revealed that Herceptin-labeled and PEGylated NPs do not specifically bind to or accumulate in HER2-positive KPL-4 tumors. However, a passive accumulation of PEGylated NPs is seen in MDA-MB-231 breast tumors, especially for the smallest 15 nm-PEG 10 kDa NPs.

The origin of fluorescence signals after treatment with 15 nm-PEG 10 kDa NPs was further assessed on the cellular level by NIR fluorescence microscopy. Therefore, tumor sections of KPL-4 and MDA-MB-231 tumors excised 24 h after injection of 15 nm-PEG 10 kDa NPs were analyzed. KPL-4 tumors of untreated mice show a relatively high autofluorescence (Figure 42A; n = 2) whereas no fluorescece in KPL-4 tumors of mice treated with NP can be observed (Figure 42B; n = 2). Compared to untreated KPL-4 tumors, MDA-MB-231 tumors of untreated mice show lower autofluorescence (Figure 42C; n = 2). After treatment of mice with 15 nm-PEG 10 kDa NPs, MDA-MB-231 tumors (Figure 42D; n = 2) show fluorescence deriving from the tumor cells. The signal distribution within the tumor is not homogeneous with most parts showing comparably low fluorescence. In addition, more fluorescence was observed at the tumor border compared to the center (data not shown).

Figure 42. Detection of 15 nm NP-derived fluorescence in MDA-MB-231 tumor sections ex vivo

Sections of KPL-4 and MDA-MB-231 tumors of mice 24 h after *i.v.* injection of 1 mg 15 nm-PEG 10 kDa NPs were examined under a fluorescence microscope with a x 40 objective. For each box, on the left upper side, counter stain of cell nuclei with DAPI and on the left lower side, probe-derived signals are illustrated. Merged images of the cell nuclei (blue) and the probe (red) are shown on the right. **A.** In untreated KPL-4 tumors (n = 2) some autofluorescence can be detected. **B.** In KPL-4 tumors of mice treated with 15 nm-PEG 10 kDa NPs (n = 2) fluorescence is not higher compared to untreated tumors in A. **C.** MDA-MB-231 tumors show almost no autofluorescence (n = 2). **D.** MDA-MB-231 tumors of mice treated with 15 nm-PEG 10 kDa NPs (n = 2) show fluorescence located around the tumor cells. The exposure time for detection of NPs was 800 ms. Bars represent 50 μ m.

Best signal detection of Itrybe-loaded NPs was observed *ex vivo* for the PEGylated 15 nm particles after injection in MDA-MB-231 tumor bearing mice. Here, illustration of tumor signals was also possible *in vivo*, as illustrated in Figure 43 (n = 2). As tumor-derived fluorescence is relatively low after probe injection (compared to the prescan) this could only be achieved by scanning exclusively the tumor region. Thereby, the high probe-derived signals in the liver which were hampering the detection sensitivity of low fluorescent signals from the tumor are excluded (see Figure 36A for comparison). Although fluorescence in the tumor is relatively weak, the intensity in the tumor (black circle) is higher 1 h after injection of NPs and further increases up to 24 h when compared to the autofluorescence in the prescan.

Figure 43. PEGylated 15 nm NPs can be detected in the tumor in vivo

In vivo fluorescence intensity scan of a MDA-MB-231 breast tumor-bearing mouse before (prescan) and 1, 4, and 24 h after *i.v.* injection of 1 mg 15 nm-PEG 10 kDa NPs (n = 2) performed in the Optix MX2. Only the tumor region (black circle) was scanned. After injection of NPs, fluorescence over the tumor increases over time.

In summary, tumor-bearing mice treated with NPs of all sizes with PEG- and Herceptin surface-modifications showed no NP-derived fluorescence in the tumor *in vivo* in whole body scans. In contrary, a closer analysis of the tumor targeting ability of NPs *ex vivo* revealed low but clear accumulation of NPs in MDA-MB-231 breast tumors. Here, the 15 nm PEGylated NPs seem to be the most promising probes for tumor imaging *in vivo*.

4. DISCUSSION

4.1. Improvement of tumor detection via pH-sensitive NIRF probes

The present work illustrates the potential of the pH-sensitive tumor-specific NIRF conjugate, pH-Her (Cyp-Her5E-Herceptin) to improve tumor-detection sensitivity *in vivo* in comparison to the always-on probe Alexa-Her (Alexa Fluor 647-Herceptin).

4.1.1. Tumor imaging with target specific pH-activatable NIRF probes

The use of pH-Her enabled non-invasive detection of tumors in vivo with a 2.5-fold increased sensitivity compared to the always-on probe Alexa-Her. The very high tumor contrast shown here by the use of pH-Her was achieved by a considerable increase in fluorescence intensity of pH-Her in the acidic environment of the tumor and after internalization in targeted tumor cells. At the same time pH-Her produced a low background fluorescence in the blood, where it faces an approximately neutral pH. The remaining autofluorescence was efficiently eliminated by subtraction of autofluorescence from prescans of mice. In contrast, the always-on fluorescence conjugate, Alexa-Her was fluorescent in the tumor as well as in the blood, as this probe did not respond to pH changes in the environment. Although Alexa-Her had a higher fluorescence intensity, this resulted in 2.5-fold lower CNRs in mice, compared to pH-Her, despite subtraction of autofluorescence, which was negligible compared to the high background fluorescence caused by Alexa-Her in the blood. The higher fluorescence intensity of Alexa-Her and Alexa-OH compared to pH-Her and pH-OH observed in this study in *in vitro* and *in vivo* measurements is due to the considerably higher quantum yield of Alexa Fluor 647 of 0.33 compared to CypHer5E which has a guantum yield of 0.27 at a pH of 5.3.

Other groups, like Urano et al.³⁸ and Ogawa and colleagues⁶, have also achieved high tumor to background ratios with a pH-sensitive Herceptin-conjugate utilizing PeT for photo quenching and a TAMRA-QSY7 fluorophore-quencher pair conjugated to Herceptin, respectively. However, as these probes both emit in the visible wavelength region, the limited tissue penetration restricted the generation of *in vivo* data and allowed only *ex vivo* measurements of the opened abdominal cavity which is of low importance for noninvasive tumor detection in living organisms. Furthermore, some studies, like those of Ogawa and colleagues lack information such as CNRs or tumor to background ratios of their activatable probes, thereby giving no quantitative data about the ability of their probes to improve tumor detection⁷.

The very low background fluorescence of pH-Her, which contributed to high tumordetection sensitivity was further attained by the lack of major signals in other organs *in vivo* and *ex vivo*. In contrast to that, it has been reported that some quenched fluorescence probes, like ICG coupled to Herceptin⁷ or Cy5.5-Herceptin-conjugates⁸ at high labeling densities, produce high liver signals resulting in decreased tumor contrast. Furthermore, since the applied fluorophore CypHer5E in pH-Her emits fluorescence in the near-infrared region, the probe is well suited for *in vivo* imaging in deeper mouse tissues as less autofluorescence is detected and tissue penetration is increased compared to fluorophores that emit in the visible wavelength region²¹.

pH-Her was well detectable in the tumor as early as one hour after probe injection in KPL-4 tumor-bearing mice. This very early detection was probably mainly due to an increase in fluorescence of pH-Her in the acidic tumor interstitium, before receptorbased internalization in the target tumor cells occurred. It is known that CypHer5E changes its absorption spectrum and thus its excitability with changes in pH due to protonation/deprotonation at the nitrogen atom^{39,43} and therefore is able to increase its fluorescence in the presence of protons already in the acidic tumor environment. Different to pH-Her, activatable fluorescence probes based on quenching mechanisms induced by homo- or hetero-FRET^{6,8}, H-dimer formation⁵, or autoquenching⁷ are mainly dependent on lysosomal degradation which occurs after internalization in targeted cells. Here, processing by e.g. proteasomal cleavage leads to dequenching of the conjugates resulting in activation of fluorescence only within the cells.

In the present study the majority of Herceptin-conjugates internalized after 8 h of incubation at 37 °C in HER2-positive cells *in vitro*, confirming previous studies which illustrate high internalization of fluorescent Herceptin-conjugates in target cells after a similar period rather than after a short time of one hour^{6,7}. This indicates that the fluorescent Herceptin-conjugates activated by dequenching in the lysosomes, reported by others, need a longer time to be detected *in vivo* in the tumor, compared to the here used pH-Her. In the present work, the fluorescent signals in the tumor were well detectable as early as one hour after injection of pH-Her in mice and CNRs continuously increased in the first 24 h. In contrast to that, Ogawa and colleagues applied quenched fluorescence probes coupled to Herceptin for *in vivo* imaging of tumors, but did not show any fluorescence images of tumor-bearing mice, earlier than 24 h or 48 h after *i.v.* injection of the probes^{7,8}, suggesting that tumor contrast at early times were low.

The pH-sensitive probe used in the present study was earlier shown by others to be reversible activatable after protonation/deprotonation by changes in pH⁵³. Although not investigated in this study, this reversible fluorescence activation might be

especially advantageous for imaging of the localization of probes in real-time as fluorescence disappears as soon as pH-Her escapes the acidic environment of the tumor or of endolysosomal cell compartments, which might occur in case of cell damage or cell death. In contrast, activation of fluorescence-conjugates via dequenching by lysosomal processing is irreversible and thereby does not provide imaging of environmental changes in real time.

Herceptin presents an excellent tumor targeting moiety for this study, as binding specificity and kinetics of Herceptin are well characterized^{5,38,111} The Herceptinconjugate, pH-Her selectively activates its fluorescence after internalization into HER2-positive KPL-4 cells after 8 h at 37 °C, but not when bound to the cell membrane at 4 °C, when cell physiology is slowed down and the pH is neutral. It has also been described by others that Herceptin-fluorophore-conjugates are internalized to a high extend after 8 h of incubation at 37 °C but not at 4 °C⁵. The spectroscopic characterizations of the pH-Her probes, showing an increase in fluorescence only when the pH is decreased, are in accordance with the proposed pH-dependent increase in fluorescence of pH-Her after internalization in cells and point to the localization of these conjugates in acidic endolysosomal compartments. My means of lysosomal markers^{5,6,38}, this was earlier confirmed by showing that Herceptin-conjuated fluorescence dyes colocalize with lysosomal compartments, which are known to have an acidic pH of 5 to 6³⁸.

pH-Her is especially advantageous for imaging purposes as a high DP ratio is not necessary for quenching of the probe. In contrast, activatable probes that utilize selfquenching via conjugation of several fluorophores to one targeting ligand need, e.g. \sim 7 to 20 dyes to be linked to one ligand^{8,112}. The conjugation of many fluorophores to one ligand is challenging as, besides the higher costs for such an amount of hydrophilic dyes, some fluorophores, like ICG, form aggregates after conjugation to ligands¹¹³. The degree of aggregation is directly related to the amount of fluorophore loaded onto the protein¹¹³. In this study, a lower stability over time was observed for the pH-Her conjugates at high DP ratios of 5.0 and 6.4, compared to pH-Her at a low DP ratio of 1.6. The fact that pH-Her DP 5.0 and 6.4 showed very low tumor targeting capacity but high bladder signals early after application in tumor-bearing mice, pointed to a structural degradation of the antibody into non-functional smaller fragments (< 5.5 nm in HD) that are filtrated by the kidney rather than eliminated by the liver, representing the normal excretion route of IgGs⁷⁵. The structural instability of these conjugates is likely caused by the high labeling density of Herceptin, which may lead to fragmentation of the IgG especially at the disulfide bridges (Jutta Pauli, personal communication). However, the amount of protein was also lower in the pH-

Her DP 5.0 and 6.4 probes, compared to the pH-Her conjugate at a low DP ratio and the Alexa-Her conjugates. This suggests that a part of the pH-Her probes at high DP ratios had precipitated, probably due to enhanced agglomeration of these probes at a high labeling density (Jutta Pauli, personal communication). Most likely, both effects, structural degradation of Herceptin-conjugates into smaller non-functional fragments as well as partial precipitation of pH-Her with high DP ratios, occurred over time. The higher stability and the increased pH-sensitivity of pH-Her probes at a low DP ratio, observed in this work in spectroscopic measurements favor its use for *in vivo* tumor detection, when compared to pH-Her probes with high DP ratios.

KPL-4 cells, applied in this study to generate orthotopic breast tumor xenografts in nude mice, showed high expression of HER2 and were efficiently targeted by Herceptin-conjugated fluorophores. This confirmed the suitability of KPL-4 cells as an optimal tumor model for the evaluation of Herceptin-conjugated fluorescence labels, as already reported in previous studies¹⁰³.

4.1.2. Suppression of background fluorescence in vivo

In the present work, tumor-detection sensitivity was improved via elimination of background fluorescence by subtraction of autofluorescence but not via LT-gated imaging. Background fluorescence was eliminated in this work, as it presents one of the greatest problems in fluorescence imaging of living mice, hampering the detection sensitivity of specific probes *in vivo*^{1,2,15,16}.

Subtraction of autofluorescence was especially advantageous in combination with the pH-sensitive probe, pH-Her. This probe exhibited a low fluorescence outside the tumor area. Thereby, and in contrast to Alexa-Her, pH-Her had only a minor contribution to the background fluorescence and after injection of pH-Her, the majority of background signals were caused by autofluorescence of the mice. This could be measured in prescans and efficiently eliminated by a software-based subtraction, thereby leading to a high detection sensitivity of the pH-Her-derived fluorescence in the tumor. This straightforward technique is possible with the majority of imaging devices, as most imaging softwares are capable to subtract a given value of fluorescence from the generated scans. In comparison, suppression of autofluorescence by LT imaging or spectral unmixing, performed by scanning the excitation and emission spectra of signals in objects, followed by their separation via their spectroscopic signatures^{21,22}, both require a special hardware. However, only few commercially available imaging devices are equipped with photon-counting systems, needed for LT imaging or tunable excitation or emission filters, used for

spectral unmixing. The software based subtraction of autofluorescence represents a simple and effective alternative.

In the present study, subtraction of autofluorescence was performed by subtracting the average autofluorescence in the untreated tumor from whole scans of the same mice after probe injection. This technique may not precisely reflect the differences in autofluorescence in certain areas of the mice. As a second option, elimination of autofluorescence can be performed by subtraction of fluorescence images generated at two different excitation wavelengths, one at which mainly the autofluorescence is measured and another wavelength at which autofluorescence and probe signals can be measured^{114,115}. However, subtraction of signals taken at different wavelengths, although they are not too far apart, is also inaccurate as the amount of autofluorescence varies at different wavelength (see section 1.1, Figure 1).

For the probes applied in this study, pH-Her and Alexa-Her, LT-gated imaging, which offers the possibility to separate probe-derived fluorescence and autofluorescence^{2,15}, did not improve tumor-detection sensitivity in vivo. However, in other studies, LT imaging has been successfully applied as a tool to increase detection of specific signals *in vivo*^{2,15}, for example, for illustration of tumors after injection of a small peptide-fluorophore conjugate². Moreover, compared to fluorescence intensity imaging, LT measurements provide the advantage of being relatively independent on probe concentration, thereby increasing the sensitivity for the detection of weak signals¹⁰⁸. Nevertheless, several factors are known to influence the LT of fluorescent probes which can lead to misinterpretation of specific signals. As also shown in this study for the free dyes, pH-OH and Alexa-OH, the LTs of fluorophores change after conjugation to Herceptin. It has also been shown that the LT of fluorescent probes can be influenced by factors, such as the presence of proteins and varying pH^{2,108,109}. In this work, the pre-evaluation of probes under distinct conditions, such as varying pH, the presence of proteins, and after s.c. injection in vivo, was performed in order to predict a characteristic LT and thereby to identify the fluorescence probes when distributed in vivo after i.v. injection. However, for pH-Her in vivo, a higher LT of about 1.5 ns was measured at the tumor site compared to the predicted LT of 1.3 ns. Therefore, in pH-Her treated tumor-bearing mice, the majority of tumor signals was eliminated after gating of images to the predicted LT. These findings indicate, that some factors in the tumor environment influence the LT of pH-Her, which could not be assessed by the LT characterization in PBS and BSA, neither at different pH nor after s.c. injection in mice. Additional factors known to influence the LT are, amongst others, tissue depth, calcium concentration, and lipids^{16,108}. Therefore, further studies are required to examine the conditions in detail that might have led to an increase in

LT of pH-Her in the tumor in vivo. As the increase in LT at the tumor site, in comparison to the predicted LT was not observed for Alexa-Her, it is likely that the sensitivity of the LT of pH-Her to the tumor environment is a characteristic of the dye, CypHer5E. This would provide an additional potential to use CypHer5E as a highly sensitive means of identifying tumors via LT-gated imaging, due to the characteristic change in LT of the probe at the tumor site. Fluorescence probes that utilize such a principle would be highly advantageous compared to probes, like Alexa-Her which resulted in high background fluorescence when circulating in the blood of mice, thereby contributing to similar LTs in the background as well as at the tumor site. LT-gated imaging using Alexa-Her did not allow an efficient separation of background signals from tumor fluorescence and thereby did not lead to an increase in tumor-detection sensitivity. Furthermore, the high background signals of the Alexa-Her conjugates are caused by the prolonged blood circulation time of IgGs³⁴. In comparison, smaller molecules, such as peptides³⁴ are cleared more rapidly from the blood and might be a better tumor-targeting moiety to be conjugated to always-on probes, such as Alexa Fluor. This may enable a better separation of probe-derived signals in the tumor from background signals, mainly caused by autofluorescence, according to their different LTs² and thus improve tumor-detection sensitivity, compared to LT-gated imaging with IgG-conjugates.

In future, the pH-sensitive NIRF probe, pH-Her may be applied for a fast and sensitive detection of weak signals, deriving e.g. from small metastatic lesions *in vivo*.

4.2. Evaluation of Itrybe-loaded PSNPs as *in vivo* NIRF imaging agents

The broadband NIR fluorophore, Itrybe, enclosed in polystyrene nanoparticles (PSNPs) of different sizes (15, 25, and 100 nm) and surface modifications was evaluated, for the first time concerning its suitability for *in vivo* tumor imaging. The results presented here predict a great potential for the use of Itrybe-loaded NPs for application in highly sensitive *in vivo* NIRF imaging. Itrybe can be excited in the NIR region at two different wavelengths and is well detectable in mice down to amounts of 1.6 μ g (for 15 nm NPs).

4.2.1. Characterization of Itrybe-loaded NPs

Enclosure of Itrybe in PSNPs with different surface modifications (NH₂, PEG, or Herceptin) had no considerable influence on the shape of spectra of Itrybe, as shown by spectroscopic analyses. This ensured a good comparability of these probes when measurements were performed with the same excitation and emission settings. The enclosure of hydrophobic Itrybe fluorophores in PSNPs enabled its application in aqueous environments and therefore *in vivo*. In addition to the conservation of the spectral range of the dye after enclosure in NPs, a further advantage of this NP system is its protection of Itrybe against species in the environment that might impair its fluorescence properties⁵⁵.

In this study, with the exception of 15 nm NPs, none of the NH₂-modifed particles showed major alteration in fluorescence with changes in pH or protein concentration. In the presence of BSA, 25 and 100 nm NPs showed a slight increase in fluorescence. The unexpected high increase in fluorescence observed for the smallest, 15 nm NPs in the presence of BSA is likely caused by interaction of proteins with fluorophores that are close to the surface of the small particles. As the ratio of particle surface area of 0.38 m²/mg PS whereas 100 nm NPs only have a surface of 0.06 m²/mg PS), the fluorophores in the 15 nm NPs have the strongest contact to its surrounding. The here performed characterization of NPs by evaluation of factors known to influence fluorescence properties is an important investigation prior to preclinical studies, as NPs used for *in vivo* tumor imaging are exposed to the usually acidic pH in the tumor and to high amounts of protein in blood and tissue.

The high amount of Itrybe molecules per NPs (which is 10 for 15 nm, 46 for 25 nm, and 3100 for 100 nm) contributes to the high brightness of NPs. Hydrophobic dyes, like Itrybe are especially suitable for dense loading into NPs of hydrophobic PS. In

contrast, loading with hydrophilic dyes is challenging, as the dyes commonly applied in fluorescence imaging, such as Cy- or Alexa dyes don't incorporate efficiently in the hydrophobic PS matrix. Moreover, Itrybe is favorably incorporated in NPs in high amounts, because, in contrast to most other commonly used NIR fluorophores, a close proximity of Itrybe does not result in quenching of fluorescence (Thomas Behnke, personal communication).

4.2.2. Detection sensitivity of Itrybe enclosed in NPs

The broad absorption spectrum of Itrybe enabled excitation of dye-loaded NPs at 635 and 670 nm, both resulting in a well measureable fluorescence above 700 nm. *In vivo* scanning revealed a stronger autofluorescence of mice at 635 nm compared to 670 nm, which is in accordance with the fact that autofluorescence is reduced towards the infrared wavelength region^{1,21}. On the contrary, fluorescence of the gastrointestinal (GI) tract of mice increased at the longer wavelength of 670 nm compared to the measurement at 635 nm. Therefore, one can take advantage of the fact that Itrybe can be excited at different wavelengths. The excitation wavelength of the dye can be selected according to the imaging purpose of the probe. In the case of detection of gastric cancer for instance, an excitation wavelength of 635 nm may be favorable, as at this wavelength background fluorescence from food in the GI tract is minimized, that may otherwise hamper the specific detection of probe-derived signals.

Itrybe-loaded 15 nm NPs were well detectable *in vivo* down to amounts of 1.6 μ g when injected *s.c.* as well as *intratumorally*. For the latter, a slightly lower fluorescence intensity was measured in the tumor compared to *s.c.* application of the same amount of probe. This was presumably caused by absorption effects in the tumor tissue due to e.g. tumor blood vessels and the fact that the probe is distributed within the tumor in deeper layers of the mice^{9,116}. As the 15 nm NPs were shown to have the lowest fluorescence intensity per mg PS, compared to the other probes, the 25 and 100 nm NPs would even show higher detection sensitivity *in vivo*.

4.2.3. Surface functionalization of NPs

The successful surface conjugation of ligands to Itrybe-loaded NPs is important to ensure specific binding of these probes to target structures, like tumor cells. Here, of all the probes, the 100 nm Herceptin-conjugated NPs showed the highest binding specificity to the HER2 antigen.

By performing an immunoassay with HER2 antigen-coated plates, the reliable evaluation of the binding specificity of Herceptin-modified NPs was possible.

However, no absolute quantitative analysis of binding specificity of Herceptin-labeled NPs to HER2 (e.g. by determination of equilibrium constant or half maximal inhibitory concentration) was performed in this study. NPs are 3D structures allowing 3D surface conjugation of many Herceptin molecules per NP, whereas the immunoassay was performed on two-dimensional antigen-coated plates. Thus, due to sterical hindrances of the relatively large NPs bound to the antigen on the plate, the specific binding of Herceptin-coated NPs on the plate is presumably less effective than *in vivo*. In the animal, the 3D bioconjugation of NPs with many ligands may even result in a higher binding probability to target cells.

For the 100 nm NPs the number of Herceptin molecules conjugated to the surface was ~70 to 100 molecules, depending on the Herceptin amount used for coupling procedure. For these probes in particular, functionalization with Herceptin was successful, with ratios between specific binding to the HER2 antigen and unspecific binding of 4.8, as measured in immunoassays. Binding specificity could not be enhanced by increasing the amount of Herceptin during the coupling procedure. Most likely, sterical hindrances did not allow further coupling of NPs with Herceptin molecules. The smaller, 25 and 15 nm sized probes showed low (ratio of 1.7) and no (ratio of 1.2) binding to the HER2 antigen, respectively. Since the amount of Herceptin molecules per NP was not determined for these probes (Thomas Behnke, personal communication), it is not clear whether antibody coupling to 25 and 15 nm NPs was successful but the antibody was non-functional after conjugation or whether Herceptin conjugation was not achieved.

This study revealed that binding of NPs of all sizes to HER2 is more efficient when Herceptin is conjugated to the NP surface using 1.5 kDa- rather than 10 kDa PEG chains as linker. One possible explanation may be that sterical hindrance via the large 10 kDa PEG chains decreased binding efficiency of Herceptin to PEG. In addition, large PEG chains at a high density could form ravels with each other, further hampering the binding to Herceptin. On the other hand, conjugation of 1.5 kDa PEG chains, especially to small, 15 nm NPs was challenging as particles tended to agglomerate after conjugation to 1.5 kDa PEG (Thomas Behnke, personal communication). Agglomeration might have been caused by the low electrokinetic potential (zeta potential) of the small particles, due to their low surface area, leading to lower repulsion of the particles among each other and thus to a higher agglomeration probability.

Furthermore, a reduction of PEG amounts for coupling of 15 nm NPs to 1/10 (50 nmol/mg PS) of the original amount did not improve Herceptin conjugation or binding specificity of particles to HER2. It may be possible that PEG was still in excess,

leading to a too high density of PEG chains on the particle surface and thus a decreased mobility of PEG chains⁶². This probably resulted in steric hindrances during antibody conjugation. In future, the procedure of bioconjugation should be further optimized for small NPs, e.g. by further reducing the amount of PEG for coupling procedures.

4.2.4. Evaluation of PSNP toxicity

To evaluate toxicity, the impact of Itrybe-containing PSNPs on cell viability was investigated. The toxicity of the free fluorophore was not assessed, since Itrybe fluorophores were sterically incorporated in the matrix of non-biodegradable PSNPs and no leakage of the dye from the 100 nm NPs was observed when incubated with a 5 % BSA solution for 4 weeks (Thomas Behnke, personal communication). This implies that no release of free Itrybe fluorophores occurs in vivo, which could have potentially caused any unwanted effects. 100 nm PSNPs bearing NH₂ surface groups and PSNPs with Herceptin molecules linked to the particle surface via 1.5 kDa PEG chains at concentrations up to 2 g/l showed no impact on cell viability of 3T3 mouse fibroblasts after 24 h of incubation. These findings are in accordance with various reports in the literature, describing NPs which consist of PS as generally non-toxic and a safe tool for *in vivo* applications^{77,79}. However, particle size and surface charge have to be taken into account when analyzing toxicity¹¹⁷, as some studies report a higher toxicity for NH₂-modified PSNPs than e.g. for carboxylated PS particles, indicating that surface coating plays a prominent role in cytotoxicity^{66,78}. However, these findings could not be confirmed in the present work, as no impact of 100 nm-NH₂ PSNPs on cell viability was observed. An increase in cytotoxicity has also been reported with a decrease in particle size^{78,79}, as well as with increasing incubation times and concentration^{66,79}. These findings suggest that the reduction in cell viability may be caused by an overloading of cells with NPs leading to a disturbance of cellular functions rather than by toxicity of the NPs itself⁷⁹.

4.2.5. Binding specificity of bioconjugated NPs in vitro

Herceptin-modified 100 nm PSNPs specifically bound to HER2-positive, KPL-4 cells and -breast tumor tissue *in vitro*, whereas control probes, NH₂-modified or PEGylated NPs unspecifically bound to KP-4 cells or tumors to a low degree. All probes showed low binding to HER2-negative, MDA-MB-231 cells and tumors as investigated by fluorescence microscopy. These findings are in accordance with immunoassay results, verifying specific binding of Herceptin-conjugated 100 nm NPs to HER2 expressed on tumor cells. Binding specificity of Herceptin-modified 100 nm and 25 nm NPs to the HER2 antigen was further proven in immunoassays by "blocking" of HER2 with excess of unlabeled Herceptin, thereby prohibiting specific binding of Herceptinconjugated NPs.

Moreover, unspecific uptake of PEGylated and Herceptin-modified NPs in KPL-4 and MDA-MB-231 tumor cells was observed, which increased over time. Unspecific uptake of NPs over time was also reported by Steinhauser et al. who applied NPs consisting of human serum albumin and conjugated to Herceptin. This group reported, that the unspecific uptake of PEGylated NPs increased after 3 h. After 5 h of incubation, no differences between Herceptin-modified and PEGylated NPs on HER2-positive breast cancer cells could be observed. Steinhauser et al. also reported unspecific uptake of both probes in HER2-negative cells¹⁰⁷. These findings, together with the results reported here illustrate that unspecific uptake seems to be a common phenomenon for several NP formulations.

In general, uptake of all particles by both cell types, KPL-4 and MDA-MB-231, increased over time and was reduced when cells were kept at 4 °C. However, NPs showed no differences in their cellular localization under incubation at 4 °C and at 37 °C, which indicates no predominant binding or accumulation of NPs at the cell membrane or inside the tumor cells. The fluorescence microscope used for analysis of NP-derived signals on the cellular level revealed insufficient power of depth resolution to discriminate membrane-derived signals from fluorescence deriving from inside the cell.

Enhanced fluorescence was observed on KPL-4 cells treated with 100 nm-Herceptin NPs at early time points of incubation. Since the internalization rate of Herceptin into HER2-overexpressing cells has been reported as low as 4 % per hour¹¹⁸, this could have predominantly been caused by antibody-receptor interaction, leading to an increase of NPs on the surface and enhanced unspecific uptake. Nevertheless, the exact way of uptake of target-specific NPs in cells is not entirely known, as reports about NP formulations of various sizes are contradictory and uptake is also known to be influenced by particle charge and material. Reports in the literature indicate that endocytosis of NPs from 43 nm to 150 nm is driven by clathrin-mediated pathways or calveolae-mediated endocytosis^{66,119,120} whereas 24 nm NPs consisting of PS have been reported to be internalized in a clathrin- and calveolae independent way¹²⁰. Furthermore, 43 nm¹²⁰ and 100 nm sized PSNPs were shown to colocalize with lysosomes after internalization⁶⁶, whereas small, 24 nm PSNPs were shown to localize outside the lysosomes but near the perinuclear space¹²⁰. This indicates a dependence of NP cell uptake mechanisms on their size. Interestingly, very small NPs (8 nm) composed of ironoxide and conjugated to PEG and Herceptin localize in lysosomes after internalization in HER2-positive tumor cells in vivo¹²¹.

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As Itrybe-loaded NPs seem to be highly and rapidly internalized in cells, they may also present suitable tools for *in vivo* labeling of highly endocytotic cells, like macrophages and thereby may function as sensitive markers for imaging of inflammatory diseases. Furthermore, Itrybe-loaded NPs could be applied to monitor cell tracking *in vivo*. Stem cells for instance could be labeled *ex vivo* by endocytosis and then their migration followed *in vivo* by imaging.

4.2.1. Tumor targeting capacity of NPs in vivo

None of the NPs of different sizes, (100, 25, and 15 nm) or surface-modifications (PEG and Herceptin) were able to visualize KPL-4 or MDA-MB-231 tumors in whole body scans after *i.v.* injection in mice. However, imaging revealed that all NP probes accumulate rapidly and to a high extent in the liver. This indicates that these probes still need modification in order to increase their longevity in the blood and their accumulation in the targeted tumor *in vivo*. For efficient delivery to the tumor, NPs should present prolonged blood circulation times, reduced nonspecific uptake in normal tissue, as well as rapid accumulation in targeted tumor tissue¹²².

NPs of all sizes bearing PEG surface-chains passively accumulated in MDA-MB-231 tumors after injection in mice as shown by *ex vivo* imaging. The highest accumulation, relative to the injected amount of probe was found in tumors of mice that received 15 nm NPs. This is probably due to the size of these NPs, which are small enough to allow the most efficient extravasation from the vasculature into the tumor tissue⁷⁴, mediated by the enhanced permeability and retention (EPR) effect. Reports in the literature describe 20 to 100 nm to be the optimal size for efficient extravasation and retention of NP from capillaries to the tumor interstitium⁶⁵ as well as a cutoff size of >400 nm¹²³. Tumor targeting with smaller NPs was also reported, e.g. for QDs of sizes of ~5 to 15 nm or PEGylated ironoxide NPs of ~8 nm^{72,74,121}. Furthermore, PEGylated calcium phosphate NPs of 16 nm accumulated in the tumor of mice with a signal persisting up to 96 h after probe injection⁵⁵. Nevertheless, in accordance with the result of the present work, nonspecific uptake in other organs, such as liver, kidney, and spleen was also reported^{55,72,74,121}.

Based on the ratio of tumor intensity, relative to the total intensity of the injected amount, the smallest, 15 nm NPs have the highest capacity for tumor targeting compared to both larger NPs. Calculations of the fluorescence intensity in the tumor relative to the injected amount of the respective probe revealed a ratio for the 15 nm NPs that was 4-fold higher than for the 100 nm particles (0.18 to 0.04) and 9-fold higher than for the 25 nm NPs (0.18 to 0.02). However, it should be taken into consideration that the number of NPs injected *in vivo* was different for each probe of different size. Thus, the injected probe of 15 nm NPs contained more particles than

the 25 and 100 nm NP probes (in this order), causing a higher number of NPs in the blood. Therefore, the higher fluorescence observed in the tumor after injection of the 15 nm probe might have been influenced by the higher number of 15 nm NPs in the blood. However, this probe is nonetheless less fluorescent than the 25 and 100 nm NP probes (in this order). Therefore a direct comparison in absolute numbers of varying amounts of NPs with different sizes and fluorescence intensities is challenging and difficult. The ratio of tumor intensity, relative to the total intensity of the injected amount, was calculated to compare the tumor targeting capacity for different sizes of NPs. In addition, fluorescence reflectance imaging (FRI) of 3D structures, such as the tumor provides only semi-quantitative data, as fluorescence measurements are surface weighted⁹.

Even when increasing the affinity of NPs to HER2-positive KPL-4 tumors via conjugation to the tumor-specific ligand Herceptin, no clear accumulation of NPs was detected in KPL-4 tumors *in vivo* as well as *ex vivo* after 24 h. Specific tumor targeting was also not achieved by others, who reported no clear differences between *in vivo* tumor-targeting capacities of target-specific NPs compared to non-targeted NPs, such as gadolinium¹²⁴ or silica NPs¹²⁵ coated with folic acid, or RGD labeled lipid NPs⁶⁵. Specific targeting of tumors has mainly been achieved by use of QDs conjugated to tumor-specific antibodies⁷² or small ligands⁷⁴. However, QDs, containing heavy metals, suffer from potentially toxic effects^{20,75}, which limit their *in vivo* application.

PEGylated NPs of all sizes accumulated in MDA-MB-231 tumors when injected in mice and revealed a higher fluorescence in comparison to the autofluorescence in tumors from untreated mice. This could be shown in *ex vivo* measurements, where fluorescence is not absorbed by the surrounding tissue of the mouse, leading to a higher sensitivity of signal detection of excised tumors.

PSNPs used in this study are relatively inflexible spheres¹²⁶, which may be one reason why they were limited in efficient extravasation from the blood and tumor penetration. In addition to the size of NPs and surface coatings, it is likely that the NP-shape and deformability is important for the probes to efficiently pass the endothelial barrier towards the tumor interstitium. Compared to the relatively rigid spheres used in this study, other groups showed that more flexible lipid NPs¹²⁶ of 35 nm used *in vivo* accumulate clearly in the tumor and resulted also in high NP-derived signals in the blood plasma⁶⁵. Such liposomal NP-formulations of 100 nm are already used for efficient delivery of anthracyclines for cancer therapy in patients⁶⁴. Other examples are relatively large but deformable glycol chitosan NPs of ~260 nm, which are well detectable *in vivo* for up to 7 days and reveal the highest signal in the tumor compared to other organs. These flexible NPs also showed longer circulation times

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than the comparable unyielding PSNPs¹²⁶ of similar size¹²². These observations indicate that the PSNP-sizes of up to 100 nm, applied in the present work have the potential for tumor targeting when modified to achieve sufficient bioavailability and that the relatively low flexibility of PSNPs might have considerable influence on the tumor penetration capacity of the NPs. However, another report shows that mesoporous silica NPs of 100 to 130 can accumulate in the tumor¹²⁵. Interestingly, NPs formed of anorganic silica are expected to be relatively inflexible. However, these NPs showed stronger fluorescence in the tumor than in other organs up to 24 h after injection¹²⁵.

These findings underline the need for detailed and systematic investigations of the *in vivo* behavior of nanosized carriers to gain more knowledge about the influence of composition, size, surface charge, and deformability on biodistribution and tumor-targeting abilities of NPs. Therefore, all NP systems should be optimized prior to routine application in preclinical studies.

In this work, only the MDA-MB-231 tumor could be targeted by Itrybe-loaded NPs of all sizes. Histological examinations of both tumor types, MDA-MB-231 and KPL-4, revealed that vascularization was comparable, providing adequate conditions for NPs to reach both tumors through the blood system. However, in the present study, MDA-MB-231 tumors preferentially developed necrosis in the tumor center only, whereas KPL-4 tumors demonstrated necrotic areas in the tumor center as well as towards the tumor boarder. These findings confirmed previous observations for both tumor types of the frequent development of massive central necrosis^{101,127}. One might speculate that since the effectiveness of penetration of the tumor by NPs is dependent on the tumor characteristics, the more frequent and expanded necrosis observed in KPL-4 tumors could probably hinder the efficient distribution of NPs in these tumors to a higher extend when compared to MDA-MB-231 tumors.

These results further indicate that Itrybe-loaded PSNPs are in principle capable of reaching the tumor, but that tumor targeting with Itrybe-loaded PSNPs is not very effective, presumably due to 1.) fast elimination of NPs from the organism and 2.) relatively low flexibility of PSNPs probably impairing efficient tumor penetration. Consequently, NP formulations other than of PS should be used for incorporation of Itrybe fluorophores, e.g. built of more flexible glycol chitosan or lipids. These should be evaluated for their suitability to improve the longevity of Itrybe-loaded NPs in the blood and their accumulation in the targeted tumor *in vivo*.

4.2.2. In vivo biodistribution of NPs

Biodistribution studies of Herceptin-conjugated and PEGylated NPs in healthy mice revealed a rapid uptake of all probes in the liver which was accompanied by a fast decrease in background fluorescence, defined as the amount of circulating probe in the blood. Knowledge about the bioavailability of Itrybe-loaded NPs, is of considerable importance for their use in tumor imaging as NPs have to be present in the bloodstream long enough to reach the targeted tumor tissue⁶². Uptake in nontargeted tissue should be low or entirely prevented as it causes unspecific background signals, hampering the identification of target-specific signals, deriving from the tumor.

A rapid uptake by the liver^{62,71} and short blood-circulation times^{71,128,129} have been observed in the literature for many NPs of various materials, including PSNPs^{67,130,131} as well. The fast liver uptake of all NPs observed in this study can be explained, because the liver is part of the excretory system and its mononuclear phagocytic system (MPS) is the basic mechanism of excretion of NPs¹²². NPs can be bound by opsonins in the blood serum, most commonly by IgG and complement proteins. These opsonins are then recognized by macrophages, usually Kupffer cells or macrophages of the liver that clear the particles from the blood. The rapid elimination of Itrybe-loaded NPs from the blood was most likely caused by uptake in the MPS that can occur within minutes and presents a major drawback for the accumulation of NPs may result in short blood-circulation times, of a few minutes⁶².

Biodistribution was analyzed, at early time points (6 min to 4 h) as uptake of probes occurred very rapidly after NP injection. PSNPs of all three sizes showed a rapid increase in liver signals, which remained high for 4 h when compared to the low background observed for all probes. *Ex vivo* analyses of excised organs confirmed the *in vivo* observations with highest signals found in the liver for all probes (average of ~84 % of the total intensity measured in all tested organs).

The elimination of NPs from the blood was assessed in dependence of their size, and surface modification. By comparing the distribution of NPs of different sizes, a slower uptake of NPs in the liver was found with an increase in size of NPs. The 100 nm NPs showed a peak intensity in the liver after ~1 h, whereas the 15 nm NPs peaked as early as ~6 min after probe injection in mice. This indicates that elimination of larger, 100 nm Itrybe-loaded PSNPs from the blood by the liver is slower compared to 25 and 15 nm NPs.

Liver signals measured *ex vivo* 4.5 h after probe injection were slightly lower for the small 15 nm NPs (6 % and 2 % lower compared to the 25 and 100 nm NPs, respectively), presumably due to enhanced renal filtration of the smallest NPs by the kidney. This is demonstrated by a slight increased of the kidney signal, of ~3 % after injection of 15 nm NPs compared to larger NPs. These findings are in accordance with

the literature, showing a decrease in liver accumulation of smaller NPs⁷¹ especially at early time points after probe injection¹²⁸ and also for PSNPs⁶⁷. Furthermore, accumulation of small gandolinium oxide NPs of ~14 nm¹²⁶ or of gold NPs of 20 nm⁷¹ was reported to occur in the kidney rather than in liver. However, high liver accumulation was described for NP-formulations with a small diameter, such as QDs of ~5 to 15 nm^{72,74,116}. Although the results presented here, showed lower uptake in the liver of 15 nm NPs, compared to 25 and 100 nm NPs as confirmed by *ex vivo* analyses, the background fluorescence *in vivo* was also considerable low over time. This indicates that the small size of NPs did not contribute to an increased blood circulation. Work by others indicates that smaller particles tend to have longer bloodhalf-lives, presumably due to reduced plasma protein adsorption on their surface^{62,67,123}.

Furthermore, ex vivo analyses revealed low accumulation of NPs in the spleen and a slight tendency of decreased uptake with a decrease in NPs size, as shown by the average intensities in the spleen that were ~8 %, 6 %, and 5 % for the 100, 25, and 15 nm NPs, respectively. Accumulation of NPs, including PSNPs¹³⁰, in the spleen has been reported previously, and is likely mediated by macrophages as part of the MPS ¹¹⁶, but has also been shown to be reduced with a decrease in size⁷¹. In addition to liver and spleen, uptake in the lung has been observed in this study, with an average intensity in the lung for NPs of all sizes of ~3 %. Lung accumulation was also described for other NPs¹²⁸, including PSNPs¹³⁰ and may be caused by endocytosis of particles by pulmonary endothelial cells.

The results presented here reveal a change in the degree of liver uptake over time when NPs were PEGylated. For PEGylated 25 and 100 nm NPs, slightly higher liver signals were measured *in vivo* and *ex vivo* over time compared to the NH₂ probes. Therefore, a decrease of sequestration of NPs in the liver by surface PEGylation, compared to NPs with NH₂ surface groups could not be observed in the present study. These findings disaccord with the common believe that PEGylation leads to a decrease in liver signals due to avoidance of NP-elimination through the MPS, resulting in prolonged blood circulation and extravasation of NPs into permeable tissues, like the tumor^{62,123}. PEG belongs to a group of long hydrophilic, neutrally charged polymer chains shown to be very effective in creating a surface barrier between the NP and the opsonins by "blocking" electrostatic and hydrophobic interactions on the surface and thereby preventing elimination of NP from the blood stream⁶². It is known that neutrally charged particles have a lower opsonization rate than charged NPs, bearing for instance, positively charged NH₂ surface groups.

compared to hydrophobic ones, as blood serum proteins adsorb less on these surfaces. Apparently, as also shown in this work PEGylation does not definitively lead to a successful decrease in elimination of particles. This is in agreement with findings of Sun and colleagues who also observed no differences in biodistribution for PSNPs with sizes of 18 and 37 nm after PEGylation with 5 kDa PEG (except for a reduction in spleen uptake compared to non-PEGylated probes)⁶⁷.

In this work, no differences in biodistribution were observed for 25 nm NPs conjugated to 1.5 and 10 kDa PEG chains, both showing comparable high liver signals and low background signals over time *in vivo*. These results were confirmed by *ex vivo* analysis of organs, illustrating comparable high intensities (of 89 and 90 % of the total intensity of all organs) for the 25 nm-PEG 1.5 kDa and -10 kDa probes, respectively. Therefore, conjugation of NPs with different lengths of PEG chains had no impact on elimination of NPs by the MPS. These findings are in contrast to some reports in the literature, where PEG chains with a weight of at least 2 kDa are required for MPS-avoidance, presumably due to a reduced flexibility of shorter PEG chains^{62,126}.

In addition to the length of the PEG chains, their surface density may also influence recognition of probes by the MPS. As reported by Owens et al., very low surface densities of PEG result in gaps on the surface of the particle, where opsonins may bind, thereby initiating NP-elimination. On the other hand, a too high density of PEG on the surface leads to restriction of motion of the PEG chains, thereby decreasing PEG mobility and thus steric hindrance properties for opsonins. In this regard, a moderate density of PEG would be optimal, with no gaps on the surface but providing sufficient mobility of PEG⁶². Therefore, in future the effect of reduced amounts of surface-PEG on a prolonged bioavailability of Itrybe-loaded NPs in the blood should be investigated.

Independent on PEGylation of 15 and 25 nm NPs, the background fluorescence did not increase after probe injection. In contrast, application of the 100 nm-PEG 1.5 kDa probe resulted in a slightly enhanced background fluorescence compared to the 100 nm-NH₂ NPs. However, this is in contrast to the finding that the PEGylated 100 nm NPs also showed higher liver uptake over time compared to the NH₂-modified 100 nm NPs. Nevertheless, due to their enhanced fluorescence in the background, the PEGylated 100 nm NPs seem to have an increased blood circulation time, compared to all other NP probes.

Characteristic for all probes investigated here was the peak in background fluorescence within the first few minutes after probe injection, showing that the circulating NPs in the blood are rapidly eliminated. This is in accordance with reports of the literature for NPs built of PEGylated- (18 and 24 nm)⁶⁷, serum albumin-coated-(50 nm)¹³¹, or carboxylated PS (100 nm)¹³⁰, as well as of PEGylated gold (20 and 80 nm)⁷¹, of silica of different shape (185 and 720 nm in length) ¹²⁹, or of dextrancoated salt-bridged polymer aggregates (100 and 500 nm)¹²⁸. However, longer blood circulation times were observed for some probes, such as PEGylated gadolinium NPs of 85 nm, which in addition showed comparably low accumulation in liver, spleen, lungs and kidneys up to 8 h after injection¹²⁴. Similar results were obtained by use of 3 and 6 nm sized silica NPs up to 3 h after injection¹³². Furthermore, a long half-live of NPs in the blood was reported for lipid NPs of 35 nm with fluorescence clearly evident in the plasma after 24 h⁶⁵.

When compared to signals of single fluorophore-conjugates, like Alexa Fluor 647 coupled to Herceptin (Alexa-Her DP ratio 1.3), the background signals in this study and thus blood circulation times were relatively low for all NPs probes after *i.v.* injection in mice. This underlines the suggestion that the excretion of NPs was relatively fast, thereby not providing sufficient time for the NPs to effectively reach the targeted tumor.

Biodistribution analyses, performed systematically in this work, advanced the knowledge about the behavior and bioavailability of Itrybe-loaded PSNPs of different sizes and surface modifications. However, it should be mentioned, that fluorescence imaging performed in this study is not an absolutely quantitative method to assess biodistibution, mainly because different tissues (e.g. blood, plasma, and organs) absorb light in a different manner⁶⁵. Of further importance for the future might be the direct analysis of fluorescence in blood samples of mice at different time points after injection of diverse NPs, as this would provide a more precise calculation of probes present in the blood pool than measurement of background fluorescence.

However, the biodistribution study explored the mechanisms for the inefficient tumor-targeting by Itrybe-loaded NPs and helped to conclude which improvements are needed for the future design of these probes to mediate tumor-detection *in vivo*. The overall goal is for NPs to be present in the bloodstream long enough to reach the targeted tumor tissue⁶². Therefore, the results obtained in this analyses, concerning the yet low bioavailability and tumor-targeting capacity of Itrybe-loaded PSNPs are helpful to further modify Itrybe-loaded PSNP probes, in regards to surface density of PEG chains or the use of more flexible materials for their design. Systematically performed studies, addressing biodistribution and suitability of novel probes for efficient tumor targeting *in vivo*, including NP systems, are often missing in the literature, thereby impeding clear interpretations of obtained results.

Taken together, Itrybe-loaded NPs used in this study, demonstrate attractive probes in regard to their brightness and 3D appearance, allowing various surface modifications for *in vivo* imaging applications. However, the specificity and bioavailability of these probes has to be further improved to enable efficient tumortargeting *in vivo*.

5. SUMMARY AND CONCLUSIONS

The work presented here illustrates a detailed and systematic *in vitro* and *in vivo* analysis of novel and innovative fluorescence probes, pH-sensitive conjugates and PSNPs loaded with the broadband fluorophore Itrybe, concerning their suitability for *in vivo* tumor imaging. Coupled to the tumor-specific antibody Herceptin, the pH-sensitive dyes and Itrybe-loaded NPs were evaluated in orthotopic mouse breast tumor models for tumor-specific targeting.

This study illustrates, for the first time, the suitability of CypHer5E for its application in *in vivo* tumor imaging, with an enhanced sensitivity compared to commonly applied, commercially available always on fluorophores, such as AlexaFluor 647.

The pH-sensitive Herceptin conjugate, pH-Her, only activated in the acidic pH of the tumor, significantly enhanced detection sensitivity of breast tumors in combination with subtraction of autofluorescence. Analysis of different methods for elimination of unwanted and perturbing autofluorescence or background fluorescence showed that especially subtraction of autofluorescence is a reliable method of image data analysis and, in combination with pH-Her, improves tumor-detection sensitivity in vivo. In contrast, as shown for pH-Her and Alexa-Her, LT-gated imaging is more challenging, as it needs extensive pre-evaluation of LTs of applied probes under different conditions. Moreover, the results of LT-gated imaging have to be considered with caution as unknown influences, e.g. in the tumor environment may change the LT of some probes in vivo and therefore lead to misinterpretations of probe-derived signals. As a consequence, the unexpected changes in LT observed for pH-Her in tumor tissue require further investigations. Probes that display extensive drifts in LT upon distinct influences of a specific environment, such as a tumor, may have great potential for sensitive detection and monitoring of cancer *in vivo* when combined with LT-gated imaging. This would allow a simple separation of tumor-derived fluorescence from background signals caused by the fluorophore in the blood, according to different LTs of the probes in certain areas of the body. In future, the pH-sensitive fluorophore CypHer5E, in conjugation with other tumor-specific ligands, may be a promising tool for the sensitive detection of other malignancies as well as for monitoring of disease progression over time in vivo. The use of CypHer5E could also assist preclinical studies that evaluate novel anti-cancer therapies by detecting primary tumors as well as metastatic lesions.

This work is the first report about the *in vivo* application of Itrybe-loaded NPs. The results obtained here underline the potential of this probe as a highly sensitive imaging tool *in vivo*. As large numbers of Itrybe fluorophores can be loaded in NPs,

very bright fluorescence probes were generated in comparison to single dye molecules. The broad spectra of Itrybe in NPs enable excitation at different wavelengths in the NIRF region and a sensitive detection of the fluorophore in vivo, thereby allowing, for example to selectively minimize unspecific signals in the GI tract. Enhanced detection sensitivity is especially important in *in vivo* fluorescence imaging, where autofluorescence or unspecific fluorescence from unbound probes or food decrease the detection sensitivity of probe-derived signals. Moreover, by analysis of binding specificity of Herceptin-conjugated NPs in vitro, the functionality of these probes was confirmed, excluding the lack of binding specificity as a factor that could limit or reduce tumor targeting of NPs in vivo. Although Itrybe-loaded PSNPs may present attractive tools for labeling of cells due to their high cellular uptake, their application for in vivo imaging of tumors has to be improved by increasing their bioavailability and tumor targeting capacity. In this regard, use of materials other than PS, for the generation of NPs, which exhibit e.g. more flexibility, could be advantageous. In addition, further investigations should be performed on the influences of a lower PEG surface density on biodistribution of NPs of different sizes.

In this study, specific tumor targeting to KPL4 tumors, overexpressing HER2 was achieved by coupling the probes to the tumor-specific antibody Herceptin. The application of antibody-single dye-conjugates, such as pH-Her enabled specific targeting of tumor structures. This was possible due to a better bioavailability of these probes *in vivo* in comparison to Herceptin-conjugated or non-bioconjugated NPs. However, fluorophore-doped NPs also exhibit great potential for *in vivo* imaging and monitoring of tumors as they provide the possibility for various modifications depending on their application purpose. By combining different labels in/on one NP, such carriers could be applied for simultaneous monitoring by different imaging methods, such as fluorescence imaging and MRI^{56,57}. They can also be surface tagged with different markers to improve binding probability to the target structures *in vivo*. In addition, modifications of composition of NP-formulations can advance the biodegradability of NPs *in vivo* and by this means their use as drug carrier systems.

In future, Itrybe-loaded NPs and CypHer5E may be used in combination with other disease-specific markers, such as antibodies or peptides for detection and evaluation of various biological processes during disease progression.

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