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Introduction

Many malignancies are distinguished by profound alterations in the extracellular matrix (ECM).¹ Pathological remodeling of the ECM leads to the development of a desmoplastic microenvironment characterized by the deposition of fibrillar collagens and other structural proteins, including fibronectin, hyaluronan and

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Contrast enhanced photoacoustic detection of fibrillar collagen in the near infrared region-I†

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Fibrillar collagen accumulation emerges as a promising biomarker in several diseases, such as desmoplastic tumors and unstable atherosclerotic plaque. Gold nanorods (GNRs) hold great potential as contrast agents in high-resolution, biomedically safe, and non-invasive photoacoustic imaging (PAI). This study presents the design and characterization of a specialized imaging tool which exploits GNR assisted targeted photoacoustic imaging that is tailored for the identification of fibrillar collagen. In addition to the photoacoustic characterization of collagen in the NIR 1 and 2 regions, we demonstrate the detailed steps of conjugating a decoy to GNRs. This study serves as a proof of concept, that demonstrates that conjugated collagenase-1 (MMP-1) generates a distinct and collagen-specific photoacoustic signal, facilitating real-time visualization in the wavelength range of 700–970 nm (NIR I). As most of the reported studies utilized the endogenous contrast of collagen in the NIR II wavelength that has major limitations to perform *in vivo* deep tissue imaging, the approach that we are proposing is unique and it highlights the promise of MMP-1 decoy-functionalized GNRs as novel contrast agents for photoacoustic imaging of collagen in the NIR 1 region. To our knowledge this is the first time functionalized GNRs are optimized for the detection of fibrillar collagen and utilized in the field of non-invasive photoacoustic imaging that can facilitate a better prognosis of desmoplastic tumors and broken atherosclerotic plaques.

tenascin C, among others.²⁻⁵ Notably, fibrillar collagens generate structurally specific complex filaments or bundles called fibers, contrasting with non-fibrillar collagens and other ECM proteins that form networks within the ECM with less pronounced features. This process triggers drastic changes in the biophysical and biochemical properties of the tissue, altering cell signaling, migration and proliferation, as well as promoting tumor growth and metastasis.1 Consequently, the detection of desmoplasia, either preceding tumor formation or present in the early stages of desmoplastic tumors, holds immense potential for enhancing patient outcomes and optimizing treatment efficacy. Notably, the excessive deposition of fibrillar collagen is a hallmark feature of the desmoplastic microenvironment in various malignancies, such as bladder cancer, breast cancer, and gastric cancer.^{2,6,7} Other than oncological diseases, exposure of collagen by atherosclerotic plaques is an indicator of plaque progression and instability.8 With this understanding, the exploration of desmoplastic microenvironments for early prognostic indicators was deemed imperative, particularly leveraging fibrillar collagens as a disease biomarker.

Over the past decade, photoacoustic imaging (PAI) has risen as a rapid, non-invasive, and safe biomedical imaging modality, synergizing the advantages of optical absorption and

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ultrasound resolution. Operating on the principle of pulsedlaser light absorption by endogenous or exogenous contrast agents within tissues, PAI generates an acoustic wave collected using a transducer, thereby furnishing real-time functional and molecular insights with exceptional resolution and tissuepenetration capabilities.9 The first window of the near infrared wavelength region (NIR I, 700-1000 nm) is ideal for deep tissue in vivo PA imaging as it has compromised optical absorption and scattering that can enable the detection of oxydeoxy hemoglobin. In addition to hemoglobin, other prominent chromophores in this window that can be visualized are melanin and lipids.^{10,11} To date, most of the reported studies utilized the second window of NIR (NIR-II, 1100 to 2000 nm) for the detection of collagen. While collagen has very distinct characteristics in the NIR II window, and intravascular PA imaging has demonstrated visualization possibilities,12 this wavelength range is not ideal for in vivo deep tissue imaging due to the limited energy of the commonly used light sources for PA imaging in this window. However, this limitation can be mitigated by utilizing contrast agents. Recent study demonstrated the feasibility of deeper tissue imaging in the NIR-II region by utilizing specially designed responsive optical probes as contrast agents and imaging deep up to 116 mm.13

Targeted exogenous contrast agents offer the dual advantage of amplifying the PAI response and selectively targeting regions of interest, thus expanding PAI's scope to molecular imaging.14-17 Several contrast agents have been developed, but the majority of these are based on small molecules that are difficult to adapt for the real targeting of specific tissues or lesions. Nanoparticles, and in particular gold nanoparticles, are of interest as promising contrast agents for their unique optical properties, possibility to tune the light absorption wavelength and safety profile due to their inert nature.18 Several types of gold nanoparticles have been designed and explored for this purpose, and among them, gold nanorods (GNRs) have shown the highest extinction coefficient and a good PA conversion efficiency. Furthermore, by modifying their aspect ratio (ratio between length and width) it is possible to tune the wavelength of maximum absorption of light, thereby enabling a perfect match between contrast agent properties and PAI application.¹⁹ In particular, the near-infrared region (NIR I), which is placed above 750 nm, is to be preferred for PAI because it leverages the optical window that allows for deeper tissue penetration and overcomes the interference from different endogenous contrast molecules present in tissues:20 GNRs with the maximum absorption wavelength peaking at 800 nm represent the best candidates for this scope.21

In this study we aimed to demonstrate that the steps for conjugating a decoy to GNRs and thus functionalized GNRs can be used to detect collagen in the NIR-I wavelength range. In addition to the *ex vivo* characterization of collagen in both NIR-I and NIR-II regions, here, we demonstrated the feasibility of an innovative approach and visualization strategy for the noninvasive detection of the desmoplastic ECM, by exploiting fibrillar collagens as prime candidates of desmoplastic areas and the utilization of GNRs as plasmonic nanoantennas for PAI.

To facilitate GNR localization to fibrillar collagens, these nanoparticles were functionalized with a highly specific and selective decoy receptor. We needed to engineer a molecule that would recognize fibrillar collagens with high affinity and specificity, without affecting their structure or functions and without exhibiting proteolytic activity. This collagen decoy receptor was engineered by inactivating human collagenase-1, *i.e.*, matrix metalloproteinase 1 (MMP-1), which demonstrates remarkable affinity for fibrillar collagens, primarily degrading collagen types I and III with high specificity.^{22,23} Additionally, MMP-1 specifically cleaves interstitial collagen types I, II, and III within the triple-helical collagen chain, yielding classic $\frac{3}{4}$ and $\frac{1}{4}$ fragments.^{24,25}

In vitro experiments confirmed the binding of the MMP-1 decoy to collagen type I from both human and rat sources. Subsequently, the localization of GNRs conjugated with the MMP-1 decoy was evaluated ex vivo using a rat model of bladder carcinoma, demonstrating the maintenance of MMP-1 decoy receptor conformation and its ability to mediate targeted delivery of GNRs to collagen-rich sites within the neoplastic ECM. Finally, we validated the decoy's ability to facilitate GNRassisted photoacoustic signal generation specifically for fibrillary collagens. To the best of our knowledge, the presented study is pioneering in demonstrating the capability of detecting collagen in the NIR-I range. Given the widespread utilization of the NIR-I range for the in vivo imaging of various tissue chromophores such as oxy/deoxy hemoglobin, lipids, and melanin, our study offers an additional value for comprehensive tissue characterization as it integrates collagen detection by utilizing specifically tailored GNRs as a contrast agent.

Results

Expression, purification and characterization of the MMP1decoy

To develop a specific and selective receptor decoy for human fibrillary collagens, the scaffold of one of the major enzymes capable of degrading this protein, such as MMP-1, was chosen. Previous reports have shown that both the catalytic and hemopexin domains of MMP-1 are required for strong MMP-1-collagen interactions.^{26–29} Thus, the zymogen form of MMP-1 was inactivated due to known mutation E200A³⁰ and expressed in HEK293-6E cells, allowing the generation of soluble secreted proteins with >95% and a yield of 21 \pm 4 mg L⁻¹ (Fig. 1A).

The thermostability analysis of the MMP-1 decoy was conducted using differential scanning fluorimetry (DSF). The melting temperature ($T_{\rm m}$) of the MMP-1 decoy was quantified to be 71.7 \pm 0.03 °C (Fig. 1B). Notably, this Tm value closely resembles that of active enzyme zymogens such as MMP-2 (71.3 °C) and MMP-8 (74.8 °C),³¹ indicating a high level of protein stability for the MMP-1 decoy.

To predict the recognition and binding of the MMP-1 decoy to collagens from other origins, the sequence similarities of collagen fragments around cleavage sites, as defined in human fibrillar collagens,^{32,33} were compared with those from various species, using basic local alignment search tool BLAST and Clustal Omega (Fig. 1C).³⁴ A high degree of sequence similarity and sequence identity in the MMP interaction domain was revealed for collagen type I from different origins, for both the

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Fig.1 Characterization of the MMP-1 decoy. (A) SDS-PAGE analysis of MMP-1 decoy purification. Lane labels: molecular mass standards; L, load; F-T, flow-through; His-Trap, protein eluted using imidazole from the Ni column; SEC, size exclusion chromatography using a Superdex 75. (B) MMP-1 decoy thermostability by DSF, showing the melting curve and derivative plot used to determine the T_m of the decoy. (C) BLAST search results; red rectangles show the alignment of collagen fragments from different origins with human collagen MMP interaction domains identified³³ or supposed³² to be MMP interaction domains. (D) Direct specific binding of the MMP1-decoy to collagen type I from different species revealed by pull down assay. Left: the SDS-PAGE image of the decoy pulling down different collagens. lanes 1 and 2: negative control and beads immobilized with the MMP14 based decoy (MMP-14 SIA). Lanes 3 and 4: the MMP-1 decoy binds to reconstituted collagen type I from rats and humans, respectively. Right: the densitometry analysis of total pulled down collagen (all bands above 139 kDa). The bars represent the normalized integrated density of collagen compared to that of the decoy.

 α 1 and α 2 chains. However, a single substitution of Ala with Ser was observed at position 945 in the α 1 chain of rat and mouse collagen, differing from human collagen type I. Additionally, one substitution (P856A) and two substitutions (L866I and I872F) were identified in the α 2 chain of collagen type I in mice and pigs respectively. These differences indicate that the MMP-1 decoy will interact with rat collagen type I, but its binding ability has to be checked for that of mice and pigs, which harbor two substitutions in the collagen interaction domain. Furthermore, only one substitution (P861A) is observed in rat collagen type III compared to that of humans, suggesting the interaction between the decoy and collagen type III in rats. Finally, two substitutions are detected in the MMP interaction domain of collagen type II in rats (A967S and E968D), mice (A867L and E968D) and pigs (A967P and E968D) and in collagen type III in mice (P857A and P861A) and pigs (L866I and I872F) indicating weakened interactions with the decoy molecule.

To confirm the predicted binding of the MMP-1 decoy to collagen type I from different origins, a pull-down assay with a His-tagged MMP-1 decoy and reconstituted collagens from humans and rats, was conducted. The eluted MMP-1-decoy complexes were analyzed by SDS-PAGE, followed by densitometry analysis (Fig. 1D). As predicted, the MMP-1 decoy binds both rat and human collagen type I. Remarkably, in comparison to published data,27 we observed the binding of the latent form of inactivated collagenase to collagen type I obtained from different origins. The data strongly suggest that the decoy binding occurs due to the interaction of the hemopexin domain with collagens, since the MMP-1 decoy exists in zymogen form. The analysis of collagen bands (>139 kDa) obtained as a result of the pull-down procedure shows that the integrated density value for human collagen is higher than that of rat collagen. These results confirm BLAST prediction, pointing to a stronger binding of the decoy to human collagen type I than to rat collagen. Remarkably, the inactive variant mutant of the MMP14 catalytic domain,35 which is non-specific to fibrillar collagens, does not bind to human or rat collagen type I, further indicating the specific binding of the MMP-1 decoy to collagen I.

Synthesis and characterization of GNRs@Chit-Dec ex vivo

With the aim to provide a biocompatible system for future *in vivo* applications, we opted to use GNRs coated with chitosan, whose biocompatibility of the entire nanosystem (GNRs@Chit) or of the single components of the nanosystem has been previously demonstrated.^{21,36-38}

The presence of a N-term 8xHis-tag in the MMP1-decoy sequence, exploited for performing decoy purification, was also utilized in a conjugation approach based on "Ni-NTA labelling" for coupling to GNRs. Chitosan-coated GNRs (GNRs@Chit) were prepared from CTAB-coated GNRs and characterized as we recently reported.²¹ A summary of the materials characterization from previously published data is available (Fig. S1 in the ESI†). The nanosystem displayed free amino groups on chitosan chains, which were exploited for binding a bifunctional linker able to couple to the biopolymeric matrix with the targeting decoy

protein. For this purpose (*N*-[5-(4-isothiocyanatobenzyl)amido-1carboxypentyl]iminodiacetic acid (isothiocyanobenzyl-NTA)) was selected. The isothiocyanate group is known to readily react with free amino groups forming stable thiourea linkages, which have been widely used to couple biomolecules, dyes, and nanostructures with a click chemistry approach.^{39,40} On the other hand, the nitrilotriacetic acid (NTA) terminus of the linker can be activated for proteins containing six adjacent histidine residues (His-tagged proteins), by complexation of Ni²⁺ ions, as previously reported (Fig. 2).^{41,42}

His-tagged green fluorescent protein (GFP) was used as a model protein to verify the effectiveness of the developed GNRs@Chit functionalization approach. Fluorescence emission spectra were recorded on purified GNRs@Chit-GFP and on the wastewaters collected during purification, revealing the quantitative binding of the His-tagged protein to the nanosystem (Fig. 3A) and no fluorescence signal arising from the wastewaters. Therefore, this approach was adopted for the Histagged decoy molecule whose conjugation to the GNRs did not induce aggregation or reshaping of the nanoparticles (Fig. 3B), neither modify the properties of the metallic core of the nanosystem (Fig. 3C and D). The functionalization with the MMP1decoy did not significantly affect the ζ-potential, which shifted from +40 \pm 6 mV for GNRs@Chit to +37 \pm 5 mV for GNRs@Chit-Dec, and there was a small shift in the position of the longitudinal surface plasmon resonance band from 800 nm for GNRs@Chit to 803 nm for GNRs@Chit-Dec, with no appreciable reduction of peak intensity or increase in the values of full width at half-maximum (Table 1).

We also assessed whether the conjugation of the decoy to the GNRs could affect the PA spectra of GNRs@Chit, which occurs in NIR-I. The PA spectra of GNRs in the absence or presence of the MMP-1 decoy were obtained, by embedding GNRs in agar drops. The GNRs were identified in B mode and the PA spectra were acquired in NIR-I, demonstrating that the conjugation of the decoy to the GNRs has not modified their expected PA spectra that peaked at 810 nm (Fig. 3E and F), as previously reported for the conjugation with a small peptide.²¹



Fig. 2 Coupling of the His-tagged decoy to chitosan-coated GNRs. First, the free amino groups of GNRs@Chit are reacted with the iso-thiocyanate terminus of the bifunctional linker. Then, the NTA terminus is exploited for Ni²⁺ complexation, thus forming a Ni-NTA complex able to bind His-tagged proteins.

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Fig. 3 Effectiveness of the conjugation approach and characterization of GNRs@Chit-Dec. (A) Fluorescence emission spectrum of GNRs@Chit-GFP compared to the wastewaters collected during its purification, revealing quantitative attachment of the His-tagged protein to the Niactivated nanosystem. (B) Representative TEM image of GNRs@Chit-Dec. (C) Normalized Uv-vis spectrum of GNRs@Chit-Dec compared to GNRs@Chit. (D) Distribution of widths and lengths of n = 55 GNRs@Chit-Dec measured by TEM. The solid lines represent the Gaussian fit for both dimensions. Photoacoustic imaging of an agar drop containing GNR@Chit-NTA-Ni (6 nmol Au) (E) and its associated PA spectra in NIR-I (680–970 nm), followed by the overlay with the PA spectra of GNR@Chit-Dec (6 nmol Au) (F).

To visualize the targeting ability of the MMP-1 decoy, a rat bladder carcinoma model was first established. Among the various bladder cancer models available, the orthotopic bladder cancer model mediated by the bladder-specific carcinogen nitrosamine (*N*-(4-hydroxybutyl)nitrosamine: BBN) was opted. Employing this compound allows all stages of bladder cancer development and progression to be monitored, thus mimicking the pathological processes that happen in humans,^{43,44} which include the formation of desmoplastic stroma (Fig. 4a and b). To assess the specificity of GNRs@Chit-Dec to recognize fibrillar collagen we used electron microscopy, because this technique allows the visualization of both collagen fibers and GNRs at the same time and could be used to demonstrate the colocalization of gold with the collagen. The ECM scaffolds obtained by decellularization of rat bladder and rat bladder cancer enriched with desmoplastic areas were subjected to high resolution scanning electron microscopy (Fig. 4c and d), confirming the desmoplastic modification that was detected using

Table 1 Physical-chemical properties of the nanostructure after various surface modification steps^a

	GNRs@CTAB	GNRs@Chit	GNRs@Chit-Dec
VIS-NIR			
$\lambda_{\rm max} ({\rm nm})$	798	800	803
FWHM (nm)	—	113	112
TEM			
GNRs length ^{b} (nm)	_	88.2 ± 6.4	91.8 ± 2.2
GNRs width ^b (nm)	_	25.7 ± 2.0	26.9 ± 2.3
Aspect ratio ^b	_	3.4 ± 0.5	3.4 ± 0.4
Zeta potential (mV)	+29.8	$+40\pm 6$	$+37.0\pm5$

hematoxylin eosin stain (Fig. 4a). In particular, distinctive features of the bladder cancer ECM were the thickening and linearization of collagen fibers, in agreement with previous information reported for human bladder cancer.45

We investigated the MMP1-decoy's targeting activity for delivering GNRs to collagen fibrils using ECM scaffolds derived from rat bladder cancer, as they maintain the structure and morphology of the tumor microenvironment (Fig. 4c and d). Scanning electron microscopy images captured after incubation of GNRs@Chit-Dec with the ECM from bladder cancer showed direct adsorption of the decoy onto the fibrils. Conversely, the control nanoparticles lacking the decoy (GNRs@Chit-NTA-Ni) did not bind to the fibrils (Fig. 4e-h).

Identification of the PA spectra of collagen I in NIR-II

To evaluate the recognition of fibrillar collagens, mainly type I collagen, by PAI of GNRs@Chit-Dec, we set up in vitro and ex vivo experimental conditions. A sterile, neutral purified porcine gelatin absorbable sponge made of collagen I, Spongostan[™], was used to identify the PA spectra of collagen I in the NIR I and NIR-II regions. B-mode US imaging was used to locate Spongostan[™] below the US gel (Fig. 5A) and to identify its PA spectra (Fig. 5B). The specific shape of the PA spectra of collagen was identified by superimposing the spectra of a water-based US gel on the spectra of Spongostan[™] (Fig. 5B). For collagen type I we ascribed a characteristic shape of the spectra in NIR-II (1200-1350 nm wavelengths) and two peaks at 1450 and 1920 nm that we ascribed to the ultrasound gel. Rat tail tendons (Fig. 5C) were used to establish the in vivo spectra of collagen I in NIR-II (Fig. 5D), which was superimposable to that of Spongostan[™] (Fig. 5D). This information provided the reference spectra of collagen I in the NIR-II range, with a characteristic three-peak shape of the PA spectra of collagen, with peaks at 1210/1250/1280 nm wavelengths.

The MMP-1 decoy delivers a GNR-assisted photoacoustic signal to collagen in NIR-I

We then investigated the PA spectra of the rat tail tendons and GNRs@Chit-Dec in NIR-I. The rat tail tendons were characterized by an unspecific spectrum and low absorbance in NIR-I,



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Fig. 4 Images illustrating the progression of bladder carcinoma in rats and the interactions of GNRs with collagen fibrils enriched in ECM scaffolds derived from bladder carcinoma tissues. Representative H&E section bladder from an 8-month rat (a) and rat bladder carcinoma after 6 months of BBN treatment (b), used for the isolation of the ECM; the red asterisk shows a typical desmoplastic stroma, which applies to all tumors present in that bladder. Representative SEM images of the ECM isolated from healthy (c) and carcinoma rat bladder (d) tissues; ECM scaffolds from healthy tissues consist of tightly packed collagen fibers with similar diameters, creating a wavy pattern. In contrast, bladder carcinoma scaffolds exhibit disrupted collagen packing, characterized by a disorganized composition of fibrils (thinnest fibrils indicated by asterisks) and fibers with varying diameters. Linearization of thick collagen fibers, as expected in solid tumors, is also observed (indicated by the arrow). The images of at least 7 fields in different locations within the desmoplastic region of each decellularized bladder from three rats were captured. Topography of the tumor ECM captured in the secondary electron mode (e) and acquired in the backscattered electron mode (f) to highlight the binding of GNRs@Chit-Dec to collagen fibrils (indicated by white dots). Panels (g) and (h) refer to the same experiment shown in panels (e) and (f), but with the ECM incubated with the control nanoparticles GNRs@Chit-NTA-Ni.

compared with the specific and intense bell-shape spectra that were acquired for GNRs@Chit-Dec in NIR-I (Fig. 6A).

The PA spectra in NIR-I was then acquired for the rat tail tendons alone, incubated with either GNRs@Chit-NTA-Ni or GNRs@Chit-Dec. The tendons did not show specific PA spectra,



Fig. 5 PA spectra of collagen I in NIR-II. Photoacoustic imaging (overlay of B mode in grayscale and photoacoustic signal in red scale) of SpongostanTM (A) and its associated PA spectra in NIR-II (1200–2000 nm) followed by the overlay with the PA spectra of the ultrasound gel (B); the wavelength of the three peaks of the collagen spectra and of the two peaks of the ultrasound gel are reported. Photoacoustic imaging of rat tail tendons (C) and their associated PA spectra in NIR-II, followed by the overlay with the PA spectra of SpongostanTM (D); the PA spectra of rat tail tendons and SpongostanTM were normalized to highlight the similarities. The right axis of US imaging shows (i) the acquisition depth in millimeters (mm), (ii) the yellow arrow the focus of the US imaging, and (iii) the red line the time gain compensation.

while the PA spectra of GNRs were detected in the samples incubated with either GNRs@Chit-NTA-Ni or GNRs@Chit-Dec (Fig. 6B). Indeed, an increased PA signal of the GNRs was acquired from tendons incubated with GNRs@Chit-Dec νs . GNRs@Chit-NTA-Ni (Fig. 6B).

The PA reference spectra of rat tail tendons and GNRs@Chit-Dec were then used to unmix the PA spectra in NIR-I (within 680–970 nm) of rat tail tendons alone, incubated with GNRs@Chit-NTA-Ni or GNRs@Chit-Dec. After spectral unmixing collagen visualization was poor in rat tail tendons alone or incubated with GNRs@Chit-NTA-Ni, while the incubation with GNRs@Chit-Dec allowed better visualization of collagen (Fig. 6C). This information demonstrated the specificity of the decoy to collagen I in its natural microenvironment. This feasibility test also proved that the Ni-NTA chemical approach, applied for decoy conjugation through histidines, maintains the MMP-1 decoy conformation, allowing GNR delivery to fibrillar collagens and giving the PA signal. GNRs@Chit-Dec show the potential to be used as contrast agents for photoacoustic imaging of collagen I in NIR-I.

Discussion

This study demonstrated the feasibility of detecting fibrillar collagen in NIR-I, using a novel contrast agent for PAI represented by GNRs@Chit conjugated with an MMP-1 derived decoy (GNRs@Chit-Dec).

We investigated the feasibility of this innovative visualization strategy for the non-invasive detection of the desmoplastic ECM, by assessing fibrillar collagens as prime candidates of desmoplastic areas and the utilization of GNRs as plasmonic nanoantennas for PAI. We also demonstrated that the steps for conjugating the decoy to GNRs@Chit did not affect the properties of neither GNRs nor the decoy. Worth noting is that the long-term storage of GNRs@Chit-Dec was ensured by the lyophilization step, as previously demonstrated by us in a related study.²¹ In fact, in this study, the photoacoustic properties and the collagen-binding capabilities of GNRs@Chit-Dec have been assessed after lyophilization and prolonged storage of the materials (up to 12 months).

This study presents a proof of concept for an innovative noninvasive approach aimed at early detecting extracellular matrix proteins within desmoplastic microenvironments, which are a distinctive pathological feature in many cancers. Our investigation demonstrates the successful targeting of collagen in both homeostatic and desmoplastic microenvironments through the conjugation of the MMP-1 decoy with innovative GNRs. Effective targeting is achieved through the distinctive structure of collagenase MMP-1 that allows recognition and binding to the collagen binding site of the hemopexin domain, which is distant from cross-linking sites.³³ This capability allows the MMP-1 decoy to deliver GNRs to collagens with varying degrees of cross-linking, including linearized collagen fibrils in the desmoplastic region.



C. Spectral unmixing; Rat tail tendons; NIR-I, GNRs@Chit-Dec; NIR-I



+GNRs@Chit-Dec; NIR-I



Fig. 6 GNRs@Chit-Dec allows photoacoustic detection of collagen in NIR-I. Reference spectra of GNRs@Chit-Dec and rat tail tendons in NIR-I (A). PA spectra or rat tail tendons alone, incubated with GNRs@Chit-NTA-Ni, or GNRs@Chit-Dec (B). PA imaging of rat tail tendons alone, incubated with GNRs@Chit-Dec (100 nmol Au), after spectral unmixing using the reference spectra of rat tail tendons and GNRs@Chit-Dec acquired in NIR-I (C); the right axis of US imaging shows (i) the acquisition depth in millimeters (mm), (ii) the yellow arrow the focus of the US imaging, and (iii) the red line the time gain compensation.

GNRs@Chit-Dec function effectively as photoacoustic antennas, generating GNR-assisted photoacoustic signals specific to collagen I, which were visualized in real-time. To investigate the feasibility of GNRs@Chit-Dec as contrast agents for PAI, the acquisition was performed in NIR-I. In general, PA imaging in NIR I normally facilitates the detection of endogenous tissue chromophores such as oxy/deoxygenated hemoglobin, while the detection of collagen is limited as it has a modest optical contrast in this wavelength range.⁴⁶ With the help of functionalized GNRs this limitation can be vanquished and the collagen can be visualized in the NIR I range. As the utilization of NIR-II wavelength has energy impairment for *in vivo* deep tissue imaging the use of GNRs@Chit-Dec for PAI in NIR-I can serve as a potential candidate for the exploration of desmoplastic microenvironments.

In addition to imaging in the NIR-I range, we also report a characteristic shape of the PA spectra of collagen in the range of 1200-1300 nm wavelength. This range of wavelengths was identified using a pure gelatin sponge, which allows the separation of the PA contribution of the water and, potentially, other contrast agents that could be present in the acellular dermal matrix recently used to derive the PA spectra of collagen.47 Furthermore, the use of a dry sponge allows the shape of the spectra of collagen vs. the shape of the spectra of water present in the ultrasound gel to be magnified, avoiding interference by the use of acetic acid commonly used to dissolve collagen.48 The range of 1200-1300 nm wavelength for the PA spectra of collagen, and that of the water in NIR-II, is in agreement with previous reports.48-52 What is relevant in the PA spectra of collagen identified in our study is the characteristic three-peak shape in the range of 1200-1300 nm wavelengths, a resolution that was not shown in the above mentioned studies. Indeed, the shape of PA spectra is at the basis of the spectral unmix approach used to differentiate specific tissue components from the overall PA signal of the tissue and thus to provide detailed molecular imaging.53 The characteristic three-peak shape of the PA spectra of collagen, with peaks at 1210/1250/1280 nm wavelengths, provides a more accurate identification of collagen-rich tissue area. The spectrum of collagen was reported to share many features similar to that of another structural protein that is elastin, although the peak of elastin spectra is at 1275 nm.⁵⁴ The three-peak shape of the PA spectra of collagen in NIR-II lays the groundwork for better discrimination of the two structural components in vivo. Although conventional spectral unmixing facilitates the detection of specific chromophores by leveraging user input spectral profiles, the presence of highly absorbing chromophores such as oxy/deoxy hemoglobin can hamper the sensitivity of lower absorbing chromophore visualizations. However, a recent study showcased the possibility of detecting prominent and non-prominent absorbers through data-driven spectral unmixing.53 Incorporating such artificial intelligence (AI) based approaches can further enhance the detection of collagen in the NIR-I region.

The versatility of GNRs extends beyond their current application, as they can be functionalized with other highly specific and selectively designed decoy molecules, enabling recognition of various extracellular matrix biomarkers within the desmoplastic microenvironment. These molecular probes bear paramount significance in the development of non-invasive, nonionizing, and highly sensitive contrast agents tailored for precise photoacoustic imaging. This approach holds immense promise for the prognosis of a multitude of solid tumors, including bladder carcinoma, pancreatic cancer, colon cancer, breast cancer, and prostate cancer. Other than desmoplastic areas, another potential application of GNRs@Chit-Dec is for the detection of disrupted atherosclerotic plaque that by exposing the fibrillary collagen ignites the formation of coronary thrombosis presenting as an acute coronary syndrome, a life-threatening disease.⁵⁵ The potential applications of decoy receptors conjugated with GNRs may be extended into multiple medical applications. These constructs may find utility in specific drug delivery, photothermal therapy, photodynamic therapy, and radiation therapy.^{56–59} Moreover, their implications in diagnostic imaging continue to hold immense prospects.

The successful conjugation of the MMP-1 decoy and GNRs offers a powerful tool for non-invasive detection and targeted intervention within the desmoplastic microenvironment. This approach may be easily adapted to other ECM biomarkers and its potential integration into a spectrum of medical applications underscores its significance in both the diagnostic and therapeutic realms.

Experimental

Materials and methods

Reagents. All the reagents, if not specified, were purchased from Sigma Aldrich (St. Louis, MO). Chitosan-coated gold nanorods (GNRs@Chit) have been prepared from CTAB-coated GNRs and thiolated chitosan. Isothiocyanobenzyl-NTA (N-[5-(4-isothiocyanatobenzyl)amido-1-carboxypentyl]iminodiacetic acid, 99%) was purchased from Dojindo EU GmbH (München, Germany). All aqueous solutions were prepared with deionized water obtained using an ultrafiltration system (Milli-Q, Millipore, Burlington, MA, USA) with a measured resistivity above 18 M Ω cm⁻¹.

MMP-1 decoy expression and purification. The zymogen form of the human MMP-1 decoy (E200A) with an N-terminal 8xHis-tag was cloned into the pTTVH8G plasmid and transformed into E. coli strain DH5a for plasmid expression. Plasmids were purified according to the protocol of a PureLinkTM HiPure plasmid filter midiprep kit. The purified plasmids were then transfected into HEK293-6E cells with polyethylenimine (PEI) for protein expression. Prior to transfection, cells were diluted to 0.5×10^6 cells per mL with FreeStyle F17 expression medium. For 1 mL of transfection, 1 μ g μ L⁻¹ of the purified plasmid was mixed with PEI in a ratio of 1:3 to form a DNA-PEI complex. After adding the DNA-PEI complex to the cells, the culture was incubated on an orbital shaker at 37 °C in a humidified incubator containing 5% CO₂ for 5 days. Tryptone N1 was introduced to the cells at a final concentration of 0.5% 24 h post-transfection.

The expressed cells were harvested and centrifuged at 300g for 20 min at 4 °C. The supernatant was collected, filtered through a 0.22 μ m membrane, and loaded onto a His-Trap 5 mL column in an AKTA purification system. The column was equilibrated with an equilibration buffer (TNC buffer: 50 mM Tris–HCl (pH 7.5), 150 mM NaCl and 10 mM CaCl₂) with 20 mM imidazole and eluted using an elution buffer (TNC buffer with 250 mM imidazole). The folded proteins were concentrated to ~10 mL using a 10 kDa cut-off Amicon (Merck) and further purified through size-exclusion chromatography (SEC) using a Superdex 75 pre-equilibrated with TNC buffer. The fraction eluted at 120–140 mL of the SEC column was concentrated to

approximately 1 mg mL⁻¹ using a 10 kDa cut-off Amicon (Merck), filtered through a 0.22 μm PVDF membrane, and stored at -80 °C. The purification protocol yielded 21 \pm 4 mg L⁻¹ of purity >95%.

Protein thermo-stability. The melting point of the MMP-1 decoy was determined using NanoDSF Prometheus NT.48. A solution of the protein was diluted to a concentration of 0.5 mg mL⁻¹, and triplicate samples were loaded into a nanodifferential scanning fluorimeter (DSF). The temperature was ramped from 20 °C to 95 °C at a rate of 1 °C min⁻¹. The fluorescent signal intensity was recorded at one-minute intervals. The obtained fluorescent ratios (F350/330) were then plotted against temperature and fitted to the Boltzmann equation using Origin 8.

Analysis of collagen MMP interaction domains. Sequence homologies between the human MMP interaction domain of collagen types I, II, and III^{60,61} and fibrillar collagen types I, II, and III from different origins were analyzed using BLAST.

The sequence analysis with BLAST was performed using the NCBI BLAST web interface (https://blast.ncbi.nlm.nih.gov/ Blast.cgi). This search involved comparing the sequence of human collagen type I (genes COL1A and COL1A2 with accession numbers P02452 and P08123, respectively) to collagens of different types from different origins in the public NCBI database. Accession numbers for fibrillar collagens examined are: Homo sapiens COL2A1, P02458; Homo sapiens COL3A1, P02461; Mus musculus COL1A1, P11087.4; Mus musculus COL1A2, Q01149; Mus musculus COL2A1, P28481; Mus musculus COL3A1, P08121; Rattus norvegicus COL1A1, P02454; Rattus norvegicus COL1A2, P02466; Rattus norvegicus COL2A1, P05539; Rattus norvegicus COL3A1, P13941; Sus scrofa (pig) COL1A1, A0A5G2QQE9; Sus scrofa (pig) COL1A2, F1SFA7; Sus scrofa (pig) COL2A1, A0A286ZWS8; Sus scrofa (pig) COL3A1, A0A286ZQ85. Alignment of the genes of interest was performed using Clustal Omega,62 and the figure was generated using ESPript 3.0.63

Pull down assay. HisPur™ Ni-NTA magnetic beads were purchased from Thermo Fisher Scientific, Israel. MMP1-decoys or negative control (inactive variant mutant of the MMP14 catalytic domain, i.e. MMP-14 SIA)64 in TNC buffer at a concentration of $\sim 8 \ \mu M$ were immobilized on Ni-NTA beads (5 μL) through incubation for 1 h at 4 °C. Subsequently, the beads were precipitated to separate the unbound decoy from the molecules bound to Ni-NTA and washed 3 times with binding buffer to remove all "unbound" decoys. For the pull-down assay, human and rat collagen type I were diluted to a concentration of 0.1 mg mL⁻¹ in binding buffer, containing 50 mM Tris, 150 mM NaCl, 10 mM CaCl₂ and 10 mM imidazole, pH 7.5 at 4 °C. Collagen solutions (100 μ L) were added to the tubes containing the Ni-NTA-MMP-1 decoy or MMP-14 SIA and mixed on an endover-end rotator for 1 h at 4 °C. The resin was then washed with 500 µL of buffer containing 50 mM Tris, 150 mM NaCl, 10 mM CaCl₂ and 25 mM imidazole, pH 7.5 at 4 °C, followed by centrifugation for 2 min at 700g. The wash step was repeated three times (vortex was employed after adding buffer to resuspend the beads adhering to the tube walls). After the final wash, the tube was centrifuged again without adding any buffer and residual buffer was removed from the pellet. The His-tagged

MMP-1 decoy–collagen complexes bound to the resin were eluted using 20 μ L of elution buffer (50 mM Tris, 150 mM NaCl, 10 mM CaCl₂ and 250 mM imidazole, pH 7.5 at 4 °C). The tubes containing the samples were briefly vortexed to resuspend any beads sticking to the walls. The reducing SDS buffer was added to each sample, which were then boiled at 95 °C for 3 min to denature the samples. The samples were loaded on 12% SDS PAGE gels for pull-down assessment.

Densitometric analysis. SDS-PAGE gels were scanned and integrated density of collagen bands was calculated using ImageJ. The integrated density was then normalized to that of the decoy.

Activation of GNRs@Chit with nickel-NTA. GNRs@Chit, prepared by binding thiolated chitosan to CTAB-coated GNRs with a surface plasmon resonance absorption peak centered at 800 nm, were diluted in water up to a gold concentration of 0.25 mM (total volume 10 mL). Parallelly, 20 mg of isothiocyanobenzyl-NTA (45.7 µmol) were dissolved in 1 mL of DMSO, and this was added to the GNRs@Chit solution under stirring and incubated at 37 °C for 1 h, during which the isothiocyanate groups of the linker bound to free amino groups on chitosan leading to functionalized GNRs@Chit-NTA. The nanosystem was purified by centrifugation and washes over Amicon membranes (MWCO 300 kDa) to remove the unbound NTA linker up to a final volume of 25 mL. At this point, 5 mL of an aqueous solution of NiCl₂ 2 mM were added to activate the NTA functionality, and a complexation reaction was allowed to take place at room temperature for 90 min. Excess of Ni(II) ions was removed by centrifugation and washes over Amicon membranes (MWCO 300 kDa), and the purified solution was freeze-dried to obtain a solid cake of GNRs@Chit-NTA-Ni which was stored at +4 °C.

Binding of His-tagged proteins to the nanosystem: GNRs@Chit-GFP and GNRs@Chit-Dec. GNRs@Chit-NTA-Ni are dissolved to achieve 10 mL of a solution with a gold concentration of 0.25 mM, and then the His-tagged protein (either 1.30 mL 100 μ g mL⁻¹ of green fluorescent protein, GFP or 1.04 mL 100 μ g mL⁻¹ of decoy) was slowly added as an aqueous solution, with the target to bind 6000 protein molecules per nanorod. After incubation at +4 °C for 60 min, the unbound protein is removed by centrifugation over Amicon membranes (MWCO 300 kDa). GNRs@Chit-GFP and GNRs@Chit-Dec were then freeze-dried and stored at +4 °C.

Characterization of GNRs@Chit-GFP and GNRs@Chit-Dec. The gold concentration was determined in all solutions *via* flame atomic absorption spectroscopy (FAAS) using a SpectraAA 100 Varian spectrometer (Agilent Technologies, Santa Clara, USA). Before performing the analysis, the solution containing gold nanorods (100μ L) were first dissolved in aqua regia (3 mL), and then diluted to 10 mL with ultrapure water. For the calibration of the FAAS measurement, Au standard solutions (concentrations: 1, 2, 5 and 10 mg L⁻¹) were prepared by diluting appropriate amounts of 1000 mg mL⁻¹ TraceCERT® solutions in 30% aqua regia. VIS-NIR absorption spectra (range 400–1100 nm) were acquired using a Cary 3500 UV-VIS-NIR modular spectrometer (Agilent Technologies, Santa Clara, USA) using a 1 cm path-length plastic cuvette. Fluorometric

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analysis was performed with an Edinburgh FLSP920 (Livingston, UK) spectrofluorometer equipped with a 450 W xenon arc lamp. Spectra were recorded with an excitation wavelength of 395 nm.

High resolution SEM imaging of extracellular matrices from rat bladder tumor. Rat bladders embedded in optimal cutting temperature (OCT) were sliced into 400 μ m sections and washed with ice-cold deuterium-deleted water (DDW) three times for 3 min each. The samples were then incubated under gentle shaking overnight at 4 °C in solution containing 30 mM EDTA, 2% Triton X-100 and 1.5 M NaCl and Complete Protease Inhibitor Cocktail (Roche). The solution was replaced twice during this process. The samples were washed five times with DDW for 1 h each at 4 °C. The tissue-derived ECM (Fig. S2 in the ESI†) was stored in DDW at 4 °C in the presence of 0.02% NaN₃ for no longer than a week.

To assess the delivery activity of the MMP-1 decoy the ECM scaffolds were incubated with a fresh solution of GNRs@Chit-Dec for 24 h at 4 °C. Subsequently, the scaffolds were gently washed with DDW three times for 30 min each at 4 °C. The samples were then fixed in a 0.1 M cacodylate buffer solution (pH 7.4) containing 2.5% paraformaldehyde and 2.5% glutaraldehyde (pH 7.2) for 60 min at room temperature followed by three washes in the same buffer. The samples were dehydrated through a series of ascending ethanol concentrations up to 100% ethanol. The dehydrated samples were observed using a Zeiss FEG Ultra55 SEM operating at 5 kV.

Histological analysis. Bladder cryosections of 10 μ m thick consecutive to the ones used for SEM were prepared. OCT was washed away with PBS and tissue slides were fixed on 4% PFA. Then hematoxylin eosin (HE) staining was performed as follows: slides were washed twice with MilliQ water and cell nuclei were stained with hematoxylin for 50 seconds, next washed for 5 min in MilliQ water and incubated on eosin for 15 seconds. After washing, tissue slides were dehydrated on an increasing gradient of ethanol and then incubated on xylene as a clearing agent. Samples were then mounted with Eukit.

GNR suspension. Freeze-dried GNRs were suspended in 0.5 mL of 1% acetic acid/phosphate buffer to obtain 1 mM Au solution, gently vortexed, and left for 15 minutes at room temperature before use.

Set up for *in vitro* detection of the PA signal of GNRs@Chit-NTA-Ni and GNRs@Chit-Dec. Since GNRs are susceptible to shape change and a consequent loss of photoacoustic properties when stimulated with pulsed light above a certain energy threshold, light attenuators (1% agar added to 0.4% intralipids) mounted on the tip of the optical fibers of a Vevo LAZR-X were used to reduce laser fluence to avoid the reshaping of the GNRs, as previously reported.²¹ The *in vitro* PA properties of the GNRs were investigated using 1% agar drops containing GNRs@Chit-NTA-Ni or GNRs@Chit-Dec (6 nmol Au).

High-resolution ultrasound (US) and photoacoustic (PA) imaging have been performed using the Vevo LAZR-X platform (FUJIFILM VisualSonics, Inc., Toronto, ON, Canada). The imaging platform includes a high frequency US system (Vevo 3100) combined with an Nd:YAG nano-second pulsed laser with a repetition rate of 20 Hz. The linear US transducer array

MX550D consists of 256 elements with a nominal center frequency of 40 MHz (25–55 MHz bandwidth) and a spatial resolution of 40 μ m with a maximum imaging depth of 15 mm. Light from the laser is delivered to the tissue through optical fibers mounted on either side of the transducer. The photo-acoustic spectra have been acquired between 680 nm and 970 nm (NIR-I) or between 1200 and 2000 nm (NIR-II) with a step size of 5 nm. The parameters used for the B-mode were: 2D power 100% and 2D gain 13 dB and for the PA-mode were: PA power 100% and PA gain 44 dB.

PAUS imaging of the binding of GNRs@Chit-NTA-Ni and GNRs@Chit-Dec to rat tail tendons. Clews of tendons were isolated from the tail of a female rat (Fischer 344) (Fig. S3A in the ESI†) and incubated with GNRs@Chit-NTA-Ni or GNRs@Chit-Dec (100 nmol Au; 0.5 mL of 0.2 mM solution in a 2 mL Eppendorf tube) for 15 minutes at room temperature under horizontal shaking (600 rpm). Control tendons were incubated with the GNR diluent under the same conditions. After two washes with PBS, the tendon clews were dried on a paper and PAUS imaging was performed using the same settings used for the agar drops. Data analysis was conducted using the VevoLab 3.2.5 software. The specific binding of GNRs to rat tail tendons was demonstrated through spectral unmixing using GNRs, melanin, oxy- and deoxy-haemoglobin spectra. The algorithm reported by Luke et al. was used to select these wavelengths which is ideal for separating the signal from GNRs from those of other endogenous absorbers.65

PA spectra of Spongostan[™] and rat tail tendons. Spongostan[™] (Fig. S3B in the ESI†) and tail tendons from female rats (Fischer 344) were placed below ultrasound gel and acquisition of the PA signal was performed in the near-infrared 1200–2000 nm optical wavelength range (NIR-II), scanned with a step size of 10 nm. The parameters used for the B-mode were: 2D power 100% and 2D gain 13 dB and for the PA-mode were: PA power 100% and PA gain 41 dB.

Rat model of bladder carcinogenesis. Forty-two eight-week old, female Fisher rats (Charles River, Germany) were housed in the animal facility at IRCCS San Raffaele Hospital under standard conditions (temperature: 22 °C \pm 2; humidity: 50 \pm 10%; light/dark cycle: 12 h light and 12 h dark). After a 1-week period of acclimatization, the rats were watered with 0.05% N-(4-hydroxybutyl)nitrosamine (BBN; Sigma-Aldrich). After 6 months of BBN treatment, the animals were euthanized using CO_2 , the bladder was filled with the cryoprotectant OCT (Bio-Optica, Milan, Italy) using a 22 G cannula (BD, Italy). The urethra was closed and the bladder was explanted, embedded in OCT, snap-frozen (isopentane and dry ice) and then stored at -80 °C, as recently reported.44 All procedures and studies involving animals were performed under protocols approved by the IRCCS Ospedale San Raffaele Animal Care and Use Committee and in accordance with national and international standard guidelines (IACUC, approval number 942).

Author contributions

Writing the original draft of the manuscript: MA. Review and editing: IS, EL, MM, AS, VG, GM, JJ, MCF, IS, and MA.

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Conceptualization: MCF, IS, and MA. Data curation and data analysis: IS, IL, ST, MU, SLA, EA, EL, MM, CV, RL, FC, VG, GM, JJ, MCF, IS, and MA. Investigation: IS, IL, ST, MU, SLA, EA, EL, MM, CV, and FC. Methodology; IS, IL, ST, MU, SLA, EA, EL, MM, CV, FC, VG, GM, and JJ. Funding acquisition: MA.

Conflicts of interest

The authors declare no competing financial interests; GM and JJ are employees of FujiFilm VisualSonics.

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