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ORIGINAL RESEARCH

A pH-Responsive System Based on Fluorescence Enhanced Gold Nanoparticles for Renal Targeting Drug Delivery and Fibrosis Therapy

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Background: Stimuli-responsive gold nano-assemblies have attracted attention as drug delivery systems in the biomedical field. However, there are challenges achieving targeted delivery and controllable drug release for specific diseases.

Materials and Methods: In this study, a glutathione (GSH)-modified fluorescent gold nanoparticle termed AuLA-GSH was prepared and a Co^{2+} -induced self-assembly drug delivery platform termed AuLA-GSH-Co was constructed. Both the pH-responsive character and drug loading behavior of AuLA-GSH-Co were studied in vitro. Kidney-targeting capability was investigated in vitro and in vivo. Finally, the anti-fibrosis efficiency of AuLA-GSH-Co in a mouse model of unilateral ureteral obstruction (UUO) was explored.

Results: AuLA-GSH-Co was sensitive to pH changes and released Co^{2+} in acidic conditions, allowing it to have controllable drug release abilities. AuLA-GSH-Co was found to improve cellular uptake of Co^{2+} ions compared to $CoCl_2$ in vitro. AuLA-GSH exhibited specific renal targeting and prolonged renal retention time with low non-specific accumulation in vivo. Moreover, the anti-fibrosis efficiency of AuLA-GSH-Co was higher compared to $CoCl_2$ in a mouse model of unilateral ureteral obstruction (UUO).

Conclusion: AuLA-GSH-Co could greatly enhance drug delivery efficiency with renal targeting capability and obviously relieve renal fibrosis, providing a promising strategy for renal fibrosis therapy.

Keywords: gold nanoparticles, self-assembly, pH sensitivity, drug delivery, renal fibrosis

Introduction

Nanotechnology has promoted new vitality in the biomedicine field.¹ Various nanoparticles (NPs), such as polymer NPs, magnetic NPs, quantum dots, carbon nanotubes and metal NPs have been studied in relation to antibacterial effects, leukemia treatment and drug delivery.^{2,3} Specifically, the application of metal NPs, especially gold NPs (AuNPs), has attracted attention due to versatile surface modifications, simple preparation and good biocompatibility.^{4,5} However, AuNPs as drug carriers still face the challenges of low drug loading efficiency, short half-life and high non-specific accumulation.^{6,7} Thus, appropriate surface modifications are necessary to broaden the application of AuNPs.^{6,8–10} Small molecule polymers and polypeptides were added to the surfaces of AuNPs with the goal of optimizing their properties.^{8,11,12} Glutathione (GSH) is an ideal surface coating for AuNPs that prolongs the half-life in blood circulation and possess particles with negative charges that escape macrophage phagocytosis.^{13,14} At the same time, abundant

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Xuandi Lai¹ Xinran Geng² Lishan Tan¹ Jianqiang Hu² Shubin Wang¹ ¹Department of Oncology, Shenzhen Key

Laboratory of Gastrointestinal Cancer Translational Research, Peking University Shenzhen Hospital, Shenzhen Peking University-the Hong Kong University of Science and Technology Medical Center, Shenzhen 518036, People's Republic of China; ²Nanobiological Medicine Center, Key Laboratory of Fuel Cell Technology of Guangdong Province, School of Chemistry and Chemical Engineering, South China University of Technology, Guangzhou 510640, People's Republic of China

Correspondence: Jianqiang Hu Nanobiological Medicine Center, Key Laboratory of Fuel Cell Technology of Guangdong Province, School of Chemistry and Chemical Engineering, South China University of Technology, No. 381, Wushan Road, Tianhe District, Guangzhou 510640, People's Republic of China

Tel +86-186 6571 1833 Email jqhusc@scut.edu.cn

Shubin Wang

Department of Oncology, Shenzhen Key Laboratory of Gastrointestinal Cancer Translational Research, Peking University Shenzhen Hospital, Shenzhen Peking University-the Hong Kong University of Science and Technology Medical Center, No. 1120, Lianhua Road, Futian District, Shenzhen 518036, People's Republic of China Tal +86-755-83923333

Tel +86-755-83923333 Fax +86-755-83061340 Email wangshubin2013@163.com



lone pair electrons from -COOH and -NH₂ residues in GSH-modified AuNPs may interact with drugs through coordination interaction, especially in the case of transition metal ions, to form a self-assembled drug delivery system.¹⁵ Since the coordinate bond forms a weak interaction, it is sensitive to pH and can be beneficial to realize controllable drug release.¹⁶ Thus, GSH-modified AuNPs exhibit superior properties when it comes to drug delivery.

Renal fibrosis is a common, final phase of chronic kidney disease (CKD). The progression of fibrosis will increase renal dysfunction, leading to end-stage renal disease.^{17,18} Traditionally, controlling systemic blood pressure, hyperglycemia or dyslipidemia are strategies for renal fibrosis therapy.¹⁹ However, treatment to slow down or even reverse the development of renal fibrosis remains a challenge. Recently, work has focused on genes or proteins that may play a role in alleviating fibrosis, leading to complex and expensive studies.²⁰⁻²² However. biomolecules are prone to degradation or non-specific enrichment during circulation in vivo, resulting in low drug concentrations in kidney and increased side effects. Therefore, designing a renal targeting delivery system containing an accessible drug with controllable release is crucial to solve these challenges.

Studies have confirmed that GSH specifically targets the kidney.^{23,24} Moreover, cobalt ion (Co^{2+}) is reported to be a hypoxia-inducible factor (HIF) activator^{25,26} Cobalt also slows down tubulointerstitial damage and maintains the integrity of capillary networks in rat fibrosis models.²⁷ Thus, taking advantage of GSH, which possesses renal targeting ability and abundant lone pairs, as well as Co^{2+} may contribute to an efficient drug delivery system for renal fibrosis therapy.

In this study, a GSH-modified fluorescent gold nanoparticle termed AuLA-GSH was prepared and a Co^{2+} induced self-assembly drug delivery platform termed AuLA-GSH-Co was constructed for renal fibrosis targeting therapy. GSH modifications enhanced the fluorescence of AuLA, which is beneficial for tracing drug delivery. In addition, AuLA possesses renal targeting ability, which can decrease non-specific drug accumulation. AuLA-GSH-Co was shown to be sensitive to pH due to the relatively weak coordination interaction between AuLA-GSH and Co^{2+} . The drug was released under acidic conditions, which is consistent with the microenvironment of the fibrotic site. The cellular uptake of Co^{2+} in AuLA-GSH-Co was significantly higher than $CoCl_2$ uptake by cells. AuLA-GSH showed noticeable renal accumulation for an extended period of time (at least 12 h) with low reticuloendothelial system (RES) retention in vivo. Meanwhile, AuLA-GSH-Co significantly improved the anti-fibrosis efficiency in unilateral ureteral obstruction (UUO) mice.

Materials and Methods Materials

Chloroauric acid (HAuCl₄·3H₂O), reduced glutathione (GSH), copper chloride dihydrate (CuCl₂·2H₂O), cobalt chloride hexahydrate (CoCl₂·6H₂O) and nickel chloride hexahydrate (NiCl₂·6H₂O) were purchased from Sigma-Aldrich (USA). 1-ethyl-3-[3-(dimethylamino)propyl] carbodiimide hydrochloride (EDC) and N-hydroxysuccinimide (sulfo-NHS) were purchased from Sangon Biotech Co., Ltd. (Shanghai, China). D,L-thioctic acid (LA) was purchased from Bioscience & Technology Co., Ltd. All reagents were used as received unless stated otherwise.

Luminescence spectra were collected using a Hitachi F-4500 Fluorescence Spectrophotometer (Japan). The hydrodynamic size of nanoparticles was analyzed using a Malvern Nano-ZS Laser Particle Size Analyzer (UK). Transmission electron microscopy (TEM) measurements were performed using a 200 kV JEOL 2010-F transmission electron microscope (Japan). Ultraviolet-visible (UVvis) absorption spectra were recorded using a Hitachi U-3010 UV-vis spectrophotometer. Thermogravimetric analyzer (TGA) curves were obtained using a Q600 SDT DTA-TG thermal analyzer (USA). Cellular uptake was measured using Agilent 7700S inductively coupled plasma-mass spectrometry (ICP-MS, USA). Fluorescent and hematoxylin and eosin (H&E) stained images were captured using a Zeiss digital camera Axio Imager (Germany). In vivo and ex vivo fluorescent images were taken using a Bruker FX PRO imaging system (Rochester, NY, USA). Real-time polymerase chain reaction (RT-PCR) analysis using SYBRGreen assays (TaKaRa, Japan) was measured using a LightCycler 96 (Roche, Switzerland).

Preparation of Fluorescent Gold Nanoparticles

The synthesis of near-infrared fluorescent gold nanoparticles was performed as previously described.²⁸ Briefly, 1.3 mg of LA powder was dissolved in 3.9 mL of aqueous solution containing 10 μ L of 2 M NaOH. The mixture was stirred at room temperature for 15 min and 40 μ L of HAuCl₄ (2%, w/v) was added dropwise under vigorous stirring. After another 5

min of stirring, the mixture changed from light yellow to colorless, which indicated that Au^{3+} was reduced to $Au^{+.29}$ Then, 80 µL of ice-cold sodium borohydride aqueous solution (50 mM) was slowly added to the mixture, and it gradually changed from colorless to reddish brown. The reaction solution was stirred overnight at room temperature to obtain LA covered gold nanoparticles AuLAs. The resulting solution was centrifuged and purified using a 10 kDa ultrafiltration centrifugation tube to remove excess LA molecules. Purified nanoparticles were re-dispersed in pH 7.4 PBS stored in a 4°C refrigerator for later use. The concentration of AuLA solution was determined using a UV-vis spectrophotometer.

GSH Modification of AuLAs

Surface modifications of AuLAs containing GSH were performed using a covalent cross-linking reaction between the -COOH group on AuLA and the -NH₂ group on GSH. First, 20.0 mg of GSH powder, 18.8 mg of EDC powder and 28.0 mg of sulfo-NHS powder were dissolved in 10 mL of pure water. The mixture was gently stirred for 12 h in a 50°C water bath. Then, the solution was cooled to room temperature and dropped to freshly prepared AuLA solution. GSH-modified gold nanoparticles AuLA-GSHs were obtained after 12 h of a light-sensitive reaction in an ice bath. Finally, the pH of the product solution was adjusted to 4–6 following 10,000 rpm centrifugation for 10 min to remove excess reactants. Purified AuLA-GSHs were re-dispersed in pH 7.4 PBS.

Ion-Induced Fluorescence Switch System and Drug Loading

CuCl₂·2H₂O, CoCl₂·6H₂O and NiCl₂·6H₂O solids were weighed and dissolved in ultra-pure water to form a 10 mM of metal ion solution. Then, 10 μ L of the metal ion solution was added to 100 μ L of AuLA-GSH (20 mM) under ultrasonic and the fluorescence spectra of the mixed solution were characterized (voltage: 750 V, excitation: 540 nm). The pH of each mixture was set to 6.0 or 7.4 using HCl or NaOH. Fluorescence intensities were measured using the same conditions mentioned above.

Under ultrasonic conditions, different volumes (0–100 μ L) of 10 mM CoCl₂ were mixed in AuLA-GSH solution (100 μ L, 20 mM), and mixtures were incubated at room temperature for 30 min. Then, the fluorescence spectra of each mixture were measured. The loading amount and rate of Co²⁺ for each sample were obtained by dissolving prepared AuLA-GSH-Co(x) (x represents the adding volume Co²⁺) in fresh aqua

regia and determining the concentrations of Au^{3+} and Co^{2+} through ICP-AES.

Cell Lines and Animals

The proximal tubule NRK-52E and renal fibroblast NRK-49F cell lines were purchased from American Type Culture Collection (ATCC, VA, USA). Balb/c and nude mice were purchased from the Guangdong Experimental Animal Center. All experiments were approved by the Animal Research Ethics Committee of Peking University Shenzhen Hospital and were conducted in accordance with the regulations and protocols of the Animal Laboratory Center of Peking University Shenzhen Hospital.

Cytotoxicity Assays

NRK-52E and NRK-49F cells were cultured in Dulbecco's Modified Eagle Medium (DMEM) containing 10% FBS, penicillin (100 U/mL) and streptomycin (100 mg/mL) in a 5% CO₂, 95% humidity and 37°C incubator. Logarithmically growing cells were detached from plates using trypsin and plated to 96-well plates. After 24 h, AuLA-GSH stock solutions with different concentrations (10-400 mM) were added to the cells and co-cultured for 12, 24 or 36 h. Finally, 10 µL of CCK-8 solution was added to each well and cultured for 2 h at 37°C. Cell viability based on optical density (OD) was calculated as follows: cell viability = (OD in the intervention wells blank wells)/(OD in the control wells - blank wells) \times 100%.³⁰ Viabilities of cells co-cultured with different concentration of $CoCl_2$ (0–40 μ M) for 12, 24 or 36 h were also measured. Three sets of parallel samples were included in all experiments and each experiment was repeated three times. Wells not containing nanoparticles served as controls.

Cellular Uptake Assays

NRK-52E cells were co-cultured with AuLA or AuLA-GSH at 37°C for 3 h and then washed in ice-cold PBS 5 times to remove free nanoparticles. Subsequently, cells were fixed with a mixture of acetone and methanol at a 1:1 ratio for 10 min and stained with phenylindoles (DAPI) for 10 min. Nanoparticle uptake was observed using a fluorescence microscope. Cell slices were prepared and observed using transmission electron microscopy (TEM).

To investigate the cellular uptake of Co,²⁺ NRK-52E cells co-cultured with AuLA-GSH-Co or CoCl₂ for different periods (10, 20, 30, 60, 120, 180, 360, 720, 1440 min) were collected. Then, cells were washed with ice-cold PBS to remove extra nanoparticles or Co^{2+} and digested with

aqua regia. Concentrations of Co^{2+} were determined using inductively coupled plasma mass spectrometry (ICP-MS).

Kidney Targeting Tests

AuLA-GSH or AuLA solution was injected into balb/c mice intraperitoneally. The fluorescence distribution was observed on the in vivo imaging system 1 h post injection (p.i.). Mice injected with AuLA-GSH or AuLA were sacrificed at different time points (1, 3, 6 and 12 h) to collect main organs (heart, liver, spleen, lung, kidney, thymus, small intestine, muscle and brain). The fluorescence of each organ was observed using an in vivo imaging system.

To further observe the specific location of AuLA-GSH in the kidney, mice injected with AuLA-GSH were sacrificed 6 h p.i. to harvest kidneys and frozen sections were immediately generated. Kidney slices were stained with AQP-1 antibody. The expression of AQP-1 antibody was detected by Alexa 488. Subsequently, the cell nucleus was stained with DAPI and observed using a fluorescence microscope.

Modeling and Grouping of Unilateral Ureteral Obstruction (UUO) Mice

Briefly, the modeling process of UUO mice was as follows:³¹ mice were anesthetized and excised on their left-back to remove the left kidney. Then, a 5–0 nylon surgical suture was tied at the end of the kidney and ligated twice. Finally, the ligated kidney was placed back into the mouse and the wound was sewn. Mice that underwent the same procedure but did not have their kidneys ligated were treated as the Sham operation (Sham) group. Specifically, mice were divided into the following groups: 1) Sham; 2) UUO mice + saline; 3) UUO mice + CoCl₂; 4) UUO mice + AuLA-GSH-Co. The intervention material was intraperitoneally injected into the mice at a dose of 5 mg/kg/day. Mice were injected continuously for 7 days before surgery and sacrificed. Relevant samples were collected 14 days post operation.

Histological Analysis

Kidney samples were fixed in 10% paraformaldehyde at 4° C overnight and embedded in paraffin for subsequent tissue staining. Tissue slices were stained using hematoxylin-eosin (H&E), Masson and Sirius red. Moreover, Fibronectin and Collagen-I expression levels in kidney slices were stained with corresponding antibodies for

each group. Tubular injury was scored ranging from 0 to 4: 0, <5%; 1, 5–25%; 2, 25–50%; 3, 50–75%; 4, >75%.

Western Blotting

Fresh kidney tissues were grounded and lysed using 1 mM protease inhibitor lysate (RIPA) on ice. Total protein concentrations for each sample were quantified using the BCA Protein Assay Kit (EMD Millipore Corp, USA). Then, all protein samples were electrophoresed using 10% SDS-PAGE gels and transferred to PVDF membranes (pore size 0.2 µm. Merck Millipore Ltd, Germany). Membranes were blocked in 5% skimmed milk for 1 h and incubated with relevant primary antibodies at 4°C overnight. Finally, membranes were incubated with secondary antibodies labelled with HPR for 1 h and signals were detected by ECL (Thermo Fisher Scientific, Shanghai). Antibodies used in this study included anti-fibronectin (Sigma-Aldrich), anticollagen-I (Calbiochem), anti-NOX2, -NOX4 and anti-IL-6 (Santa Cruz Biotechnology), anti-MCP-1, anti-p-ERK, anti-ERK, anti-BAX (Cell Signaling Technology, Beverly, MA) and anti-ICAM (Abcam, USA).32

RNA Extraction and Quantification

RNA extraction and quantification were performed as follows:³³ fresh kidney samples were grounded and lysed in TRIzol. Then, chloroform (1:5 volume ratio with TRIzol) was added, samples were mixed for 15 s and tissue homogenates were centrifuged in 12,000 g for 15 min at 4°C. Subsequently, isopropyl alcohol and ethanol were added to precipitate and purify RNA. The mRNA levels were quantitatively detected through RT-PCR and calculated after normalization with the endogenous control, GAPDH. <u>Table S1</u> includes primer sequences used in RT-PCR analyses.

Results and Discussion

Morphology and Structure

Characterization of AuLA and AuLA-GSH AuLA was prepared using a "green" method where D, L-lipoic acid was used as reductant and surfactant.²⁸ Asprepared AuLAs were uniform spherical particles with diameters of 2.5 ± 0.3 nm observed by TEM (Figure S1). To target AuLA carrying medicine to the kidney, GSH was cross-linked to the surface of AuLA to obtain kidneytargeting nanoparticles AuLA-GSH. The morphology of AuLA-GSH was nearly the same as that of AuLA, which was spherical and contained a diameter of 2.9 ± 0.6 nm

Lai et al

(Figure 1A and B). The hydrodynamic size of AuLA-GSH (6.7 \pm 0.8 nm, PDI%: 20.5) measured by DLS was slightly larger than AuLA (5.4 \pm 0.7 nm, PDI%: 13.5), as shown in Figure 1C, indicating successful modification of GSH onto AuLA. This was due to the fact that the modified GSH molecule could interact with water molecules through hydrogen bonds in aqueous solution, so that nanoparticles were covered by a layer of hydrated molecules.³⁴ AuLA-GSH yielded red fluorescence with emission excitation

peaks at 710 nm and 395 nm, respectively (Figure 1D). Although the emission wavelengths of AuLA-GSH and AuLA were nearly the same (Figure S2), the fluorescence intensity of AuLA-GSH was significantly enhanced compared to AuLA under the same conditions (Figure 2A and B). This feature of AuLA-GSH is beneficial to its application in non-invasive tracking of drug delivery.

The FTIR spectra of GSH, AuLA and AuLA-GSH were measured (Figure 2C). A new vibration peak at



Figure I Characterization of AuLA and AuLA-GSH. TEM images of AuLA-GSH using a magnification of (A) 5000 and (B) 20,000, the inset of B is particle size distribution counted from the corresponding TEM image. (C) Hydrodynamic size distributions of AuLA and AuLA-GSH. (D) UV-vis absorption, fluorescence excitation and emission spectra of AuLA-GSH, insets are digital photos of AuLA-GSH solution taken under white light and UV light.



Figure 2 Fluorescence enhancement properties and structure characterization of AuLA-GSH. (A) The LA-coated gold nanoparticles (AuLAs) exhibit weak fluorescence while GSH-modified AuLA-GSH perform enhanced fluorescence. (B) Fluorescence emission spectra of AuLA and AuLA-GSH under the same particle concentration. The successful modification of GSH in AuLA was confirmed by (C) fourier transform infrared spectroscopy (FTIR) and (D) thermogravimetric analysis (TGA).

2300 cm⁻¹ appeared in the FTIR spectrum of AuLA-GSH compared to AuLA, which was consistent with the characteristic peak of the –SH group in GSH.^{35,36} Moreover, two weight-loss platforms for AuLA-GSH detected by TGA indicated that there were two types of molecules in the nanoparticles (Figure 2D). These results further confirmed the successful modification of GSH on AuLA.

Fluorescence Switch Features and Co²⁺ Loading of AuLA-GSH

Hypoxia and inflammatory responses usually cause changes in microenvironment pH from neutral to acidic. This characteristic of lesions provides an intrinsic basis for a pHresponsive DDS design.³⁷ Buffer solutions with a pH 7.4 and 6.0 PBS were introduced to mimic normal physiological and pathological environments, respectively. Abundant lone pair electrons of –SH and –COOH residues on the surface of AuLA-GSH could easily coordinate with transition metal ions.^{38,39} which enabled AuLA-GSH to be an ideal metal ion carrier. By co-incubation with Co²⁺, Ni²⁺ or Cu²⁺ under a pH of 7.4, the fluorescence intensities of AuLA-GSH

-x (x represents Co or Ni or Cu) at the maximum emission wavelength (705 nm) were reduced by 31.4%, 52.8% and 89.4%, respectively (Figure 3A). This was attributed to the ion-induced coordinate self-assembly of AuLA-GSH.^{40,41} When changing the pH values of these mixtures from 7.4 to 6.0, the fluorescence intensities of AuLA-GSH-Co and AuLA-GSH-Ni recovered by ~30%, while the fluorescence intensity of AuLA-GSH-Cu did not change (Figure 3B, S3). This was due to the fact that the strength of coordination bonds between Cu²⁺ ions and residues in AuLA-GSH was more robust than those of Co²⁺ and Ni²⁺ ions, which would not be influenced by pH. The fluorescence switch performance of AuLA-GSH-Co and AuLA-GSH-Ni could be generated by tuning pH values between 7.4 and 6.0, which meant that Co and Ni ions would be released in an acidic environment. This is a fundamental characteristic for AuLA-GSH to be a controllable drug carrier.

Cobalt has been reported as a promising drug for kidney fibrosis.²⁶ Thus, AuLA-GSH-Co was further studied as a drug delivery system targeting renal fibrosis. As shown in Figure 3C, the hydrodynamic size of AuLA-GSH-Co



Figure 3 Fluorescence switches and drug loading measurements. (**A**) Fluorescence emission spectra of AuLA-GSH incubated with Co^{2+} , Ni^{2+} or Cu^{2+} ions, respectively. (**B**) The fluorescence intensities at 705 nm of AuLA-GSH or AuLA-GSH-Co/Ni/Cu under pH 7.4 or 6.0. (**C**) Hydrodynamic sizes of 100 μ L AuLA-GSH incubated with different volumes (0–100 μ L) of 10 mM CoCl₂ aqueous solution and PBS (100–0 μ L) for 2 h and (**D**) TEM image of AuLA-GSH-Co under neutral conditions. (**E**) Loading amount and correlated (**F**) loading percentage curves of Co^{2+} on AuLA-GSH-Co.

increased when the added amount of Co^{2+} increased. At the same time, the fluorescence intensity of AuLA-GSH diminished gradually along with increasing Co^{2+} volume (Figure S4). AuLA-GSH-Co assemblies were composed of several nanoparticles with Co^{2+} ions entrapped (Figure 3D). The loading amount of Co^{2+} in AuLA-GSH-Co samples was determined using ICP-MS (Figure 3E and F). When the amount added was lower than 10 µg, the loading percentage of Co^{2+} was higher than 80%, indicating that most Co^{2+} was entrapped in the DDS. The loading percentage gradually decreased when the amount of Co^{2+} was increased, indicating a saturation state of AuLA-GSH in loading Co^{2+} . Thus, the most effective loading point of Co^{2+} was 10 µL adding volume.

Cellular Toxicity and Uptake Assays

The proximal tubule NRK-52E and renal fibroblast NRK-49F cell lines were introduced to evaluate cellular toxicity of AuLA-GSH. Cells were incubated with different concentrations of AuLA-GSH (0-4000 µM) and CoCl₂ (0-40 µM). CCK-8 results revealed that cell viabilities cocultured with AuLA-GSH were greater than 90% at different time points (6, 12, 24 h), indicating that AuLA-GSH had good biocompatibility (Figure 4A and B), which laid a solid foundation for its biomedical application. In contrast, CoCl₂ showed considerable cytotoxicity at high concentrations (Figure S5). The uptake efficiencies of AuLA and AuLA-GSH in NRK-52E cells were next investigated. As shown in Figure 4C and D, the red fluorescence was much stronger in cells incubated with AuLA-GSH compared to cells incubated with AuLA. The fluorescence intensity of AuLA-GSH incubated cells was 1.6 times more than AuLA incubated cells. This may be due to the enhanced fluorescence of AuLA-GSH and the specific targeting ability of GSH to proximal tubules. Cell slices observed by TEM confirmed that AuLA-GSH could be taken up through endocytosis after a period of co-culturing (Figure S6).

To further quantify cellular uptake of Co,²⁺ cell samples incubated with CoCl₂ or AuLA-GSH-Co for 0–48 h were digested and measured by ICP-MS (Figure 4E). The concentrations of Co^{2+} in AuLA-GSH-Co incubated cells were much higher than in CoCl₂ incubated cells within 1 h, which may be attributed to the nanoencapsulation of Co^{2+} ions in AuLA-GSH-Co, enhancing endocytosis and slowing exocytosis. This feature allows AuLA-GSH as an ideal candidate for Co^{2+} delivery.

To observe the distribution of AuLA-GSH in cells, the two specific fluorescent tracer molecules LAMP-1-FITC (green fluorescence) and DAPI (blue fluorescence) were introduced to label lysosomes and nuclei, respectively.⁴² Co-location imaging results showed that the position of AuLA-GSH (red fluorescence) mostly overlapped with lysosomes (green fluorescence), while red fluorescence was almost absent in the nucleus (blue fluorescence) (Figure S7). This indicated that AuLA-GSH was mostly present in lysosomes.

Targeting AuLA-GSH to the Kidney

To verify the kidney-targeting ability of AuLA-GSH, nude mice were injected with PBS, AuLA or AuLA-GSH intraperitoneally. AuLA and AuLA-GSH showed red fluorescence under the in vivo imaging system (Figure S8). After injecting, mice were sacrificed at different time points to harvest major organs (heart, liver, spleen, lung, kidneys, brain, small intestine and muscle) and observed using the imaging system. Organs collected from mice injected with PBS served as controls and showed weak autofluorescence (Figure S9). AuLA-GSH injected mice exhibited strong fluorescence in the kidneys for at least 12 h, whereas AuLA injected mice showed weak fluorescence intensity in the kidneys at 6 h p.i. (Figure 5A). Most of the AuLA-GSHs accumulated in the kidney and bladder at 1 h p.i., indicated that AuLA-GSH could specifically target the kidney and be excreted in urine (Figure 5B). It is well known that the liver is the largest RES organ and nonspecific accumulation of nanoparticles are most likely to occur in this organ.43 Here, an intensity contrast index $(CI)^{44}$ was introduced to visually describe the kidneytargeting efficiency of nanoparticles, which represented the contrast intensity of kidneys to the liver and was calculated as follows:

$$CI = \frac{\text{Fluorescence intensity of kidneys}}{\text{Fluorescence intensity of liver}} - 1$$

When the CI value was positive, it indicated that the fluorescence intensity of the kidneys was stronger than the liver. Otherwise, the florescence intensity of the liver was stronger than the kidneys. As shown in Figure 5C, CI values remained positive within 6 h p.i. for AuLA-GSH injected mice. In contrast, CI values for AuLA injected mice were almost all negative. This result indicated a remarkable advantage of AuLA-GSH in kidney targeting.



Figure 4 Cytotoxicity and cellular uptake analysis. Cell viability of (A) NRK-52E and (B) NRK-49F cell lines treated with AuLA-GSH for 6, 12 or 24 h. (C) Fluorescence images of NRK-52E incubated with PBS, AuLA or AuLA-GSH for 3 h. (D) Corresponding fluorescence intensities of nanoparticles in cells. (E) Cellular uptake of Co^{2+} in NRK-52E cells incubated with AuLA-GSH-Co or $CoCl_2$ within 48 h. Data are expressed as mean \pm SD, $n \ge 3$. *p < 0.05, ***p < 0.001.





Figure 5 In vivo and ex vivo bio-distribution of AuLA and AuLA-GSH. (\mathbf{A}) Ex vivo fluorescent images of major organs collected from mice injected with AuLA or AuLA-GSH at different time intervals (1, 6 and 12 h) (1–9, heart, liver, spleen, lung, kidneys, thymus, small intestine, muscle, brain). (\mathbf{B}) In vivo fluorescence distribution of nude mice after 1 h AuLA-GSH injection. (\mathbf{C}) Relative contrast indexes representing contrast fluorescence intensity of kidneys to liver were calculated at different time points. (\mathbf{D}) Co-localization images of AQP-1 (green), nucleus (blue) and AuLA-GSH (red) in kidney sections.

To further verify the targeting location of AuLA-GSH, kidney frozen sections were collected and stained with AQP-1, which mainly was expressed in the epithelial cell membrane of proximal tubules but not in the glomerulus or other types of tubules.⁴⁵ Figure 5D reveals that the position of AuLA-GSH (red fluorescence) and AQP-1 (green fluorescence) almost completely overlapped and the fluorescence of AuLA-GSH outlined the shape of renal tubules. This indicated



Figure 6 Histology and pathology analysis of kidneys in different treatment groups. (A) Histological sections of kidneys collected from different treatment groups stained with H&E, Sirius red and immunohistochemical staining of collagen-1 and fibronectin. (B) Quantitative assessment of tubular injury score, (C) infiltrated cells and (D) tubulointerstitial fibrosis area in different treatment groups. Data are expressed as mean \pm SD, n = 5. *p< 0.05 versus UUO+ CoCl₂ group. **p< 0.01 versus UUO+ CoCl₂ group.

that AuLA-GSH specifically targeted proximal tubular cells, which could be the internal reason for the significant renal targeting of AuLA-GSH.

In vivo Antifibrosis Efficacy

Sham mice and UUO mice injected with AuLA-GSH, $CoCl_2$ or AuLA-GSH-Co for 7 days with a Co^{2+} dose of 5 mg/kg

mouse were weighed every day. All mice were sacrificed on the 7th day p.i., and main organs including the heart, liver, spleen, lung and kidneys were collected for histological staining. H&E staining of the heart, liver, spleen and lung for each mouse indicated that all interventions did not harm normal organs (Figure S10). Compared to the Sham group, the kidneys of UUO mice treated with AuLA-GSH showed apparent tubular cell damage accompanied by tubular expansion, inflammatory cell infiltration and severe interstitial fibrosis (Figure 6A). Moreover, collagen-I and fibronectin staining revealed that the kidneys of UUO mice showed obvious collagen deposition and fibrosis. The degree of kidney damage in UUO mice treated with CoCl₂ was slightly relieved and the deformation of renal tubules and the lumen was smaller than what was observed in the AuLA-GSH group. Surprisingly, kidney morphology of UUO mice treated with AuLA-GSH-Co was significantly restored, the number of inflammatory cells was reduced and the tubular shape was similar to what was observed in the Sham group. The Masson staining results showed the same trend (Figure S11).

Figure 6B-D includes a semi-quantitative analysis of tubular damage index, the number of infiltrated inflammatory cells and interstitial fibrosis area in the kidney for each mouse group. Inflammatory cells reduced from 80%





to 50% and 30% in $CoCl_2$ and AuLA-GSH-Co treated mice, respectively. Furthermore, the ratio of interstitial fibrosis was decreased to 16.2% in the AuLA-GSH-Co group, while ratios were 40.5% and 31.3% in the AuLA-GSH and $CoCl_2$ groups, respectively. Their results indicated that AuLA-GSH-Co played a prominent role in maintaining tissue morphology of the damaged kidney and reducing infiltration.

Molecular Biology Assays Confirming Anti-Fibrosis Efficiency

A variety of fibro-related proteins in the kidney were analyzed by Western blotting. As shown in Figure 7A and B, AuLA-GSH-Co significantly down-regulated collagen-I and fibronectin expression levels by 2.5 and 1.4-fold lower, respectively, than what was observed in the saline group. The expression levels of other representative proteins including Nox2, Nox4, Bax, p-ERK and ERK, were also dramatically reduced in the AuLA-GSH-Co group (Figure 7C-E).

To further confirm anti-fibrosis efficiency, real-time PCR testing was used. Figure 7F reveals that fibrorelated mRNA expression levels of collagen-I, fibronectin and IL-6 were significantly reduced in the AuLA-GSH-Co group compared to the saline and CoCl₂ groups, which was consistent with Western blotting results. These findings demonstrated that AuLA-GSH-Co has the ability to reduce extracellular matrix deposition and act as an ideal strategy for renal fibrosis therapy.

Conclusion

Here, we have prepared GSH-modified gold nanoparticles, AuLA-GSHs containing enhanced fluorescence properties and remarkable renal targeting ability. Plentiful lone pair electrons on the surface of AuLA-GSHs possessed them with excellent metal coordination ability, which was beneficial to load and deliver Co²⁺ as a potential anti-fibrosis drug. In vitro and in vivo experiments revealed that the AuLA-GSH significantly improved kidney targeting compared to AuLA. In addition, self-assembled AuLA-GSH-Co showed a significant advantage in anti-fibrosis efficiency compared to free Co.²⁺ With these attractive properties, AuLA-GSH-Co is a promising drug delivery platform for renal fibrosis and other diseases.

Abbreviations

AuNPs, gold nanoparticles; GSH, glutathione; CKD, chronic kidney disease; HIF, hypoxia-inducible factor; RES,

reticuloendothelial system; UUO, unilateral ureteral obstruction; OD, optical density; DAPI, phenylindoles; CI, contrast index.

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Disclosure

The authors report no conflicts of interest in this work.

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