Synthesis and biological evaluation of 2.4 nm thiolate-protected gold nanoparticles conjugated to Cetuximab for targeting glioblastoma cancer cells via the EGFR

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Abstract
Therapeutic monoclonal antibodies benefit to patients and the conjugation to gold nanoparticles (AuNPs) might bring additional activities to these macromolecules. However, the behavior of the conjugate will largely depend on the bulkiness of the AuNP and small sizes are moreover preferable for diffusion. Water-soluble thiolate-protected AuNPs having diameters of 2–3 nm can be synthesized with narrow polydispersity and can selectively react with incoming organic thiols via a $S_N2$-like mechanism. We therefore synthesized a mixed thionitrobenzoic acid-, thioaminobenzoic acid-monolayered AuNP of 2.4 nm in diameter and developed a site-selective conjugation strategy to link the AuNP to Cetuximab, an anti-epidermal growth factor receptor (EGFR) antibody used in clinic. The water-soluble 80 kDa AuNP was fully characterized and then reacted to the hinge area of Cetuximab, which was selectively reduced using mild concentration of TCEP. The conjugation proceeded smoothly and could be analyzed by polyacrylamide gel electrophoresis, indicating the formation of a 1:1 AuNP-IgG conjugate as the main product. When added to EGFR expressing glioblastoma cells, the AuNP-Cetuximab conjugate selectively bound to the cell surface receptor, inhibited EGFR autophosphorylation and entered into endosomes like Cetuximab. Altogether, we describe a simple and robust protocol for a site-directed conjugation of a thiolate-protected AuNP to Cetuximab, which could be easily monitored, thereby allowing to assess the quality of the product formation. The conjugated 2.4 nm AuNP did not majorly affect the biological behavior of Cetuximab, but provided it with the electronic properties of the AuNP. This offers the ability to detect the tagged antibody and opens application for targeted cancer radiotherapy.

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(Some figures may appear in colour only in the online journal)

1. Introduction

Nanoparticles (NPs) are particles of sizes ranging between 1 and 100 nm that have important biomedical applications [1]. Some NPs can be functionalized with multiple elements, which permits to provide the nanomaterial with new properties. The coalescence of several functions allows dealing with the complexity of biological systems and might help for diagnosing and treating diseases [2, 3]. Several sophisticated systems demonstrated some efficiencies at preclinical stages for imaging modalities [4], nucleic acid delivery [5], protein delivery [6], tissue-targeted drug delivery [7], hyperthermia and photoablation therapy [8]. Gold nanoparticles (AuNPs) have been extensively investigated for biomedical application, because they have a low toxicity profile and their unique optic and electronic properties can trigger cellular damage upon application of light [9] or radiation [10, 11]. Furthermore, AuNPs can be equipped with organic molecules, including antibodies, which facilitate accumulation of the AuNPs within selected tissues or cancer lesions [12]. AuNPs with diameters above 5 nm display a large surface area that can be used for tight adsorption of antibodies and other proteins [13, 14]. For example, El-Sayed et al coated 40 nm AuNPs with monoclonal antibodies targeting the epidermal growth factor receptor (EGFR) by random adsorption in order to target oral squamous carcinoma cells. The antibody-mediated accumulation of these AuNPs into the cancer cells was then used to promote cell death via a photothermal treatment [9]. Patra et al synthesized 5 nm AuNPs that were also surface-coated with anti-EGFR antibodies, as well as with gemcitabine for targeting the drug to cancer cells [15]. Although of straightforward practicability, the functionalization of AuNPs via adsorption to the particle’s surface has limitations. Firstly, a control over the orientation and stoichiometry of the adsorbed molecules onto the AuNPs is challenging [16]. Secondly, the physicochemical properties of the antibody and its subsequent cellular response are impacted by the AuNP’s size [17, 18]. When the properties of an AuNP-IgG conjugate should resemble the ones of an antibody, AuNPs of smaller sizes should be selected. However, since antibodies do not tightly adsorb to the surface of small-sized AuNPs, the functionalization method must be adjusted by the formation of an Au–S coordination bond.

Small-sized and uniform AuNPs with diameters between 0.8 and 2 nm are easily prepared by reduction of chloroauric acid in the presence of organic thiols [19]. Thiobenzoate-protected AuNPs of such small sizes and of rather precise chemical composition can be directly prepared in aqueous solutions leading to water-soluble AuNPs. These AuNPs can be further grafted with biological macromolecules, such as oligonucleotides, peptides and proteins [20, 21], or viruses [22] by exchanging the thiobenzoate ligands with incoming thiol-containing macromolecules. To diminish unspecific association to cellular constituents and to enhance the ligand exchange reaction, we have previously developed a mixed thionitrobenzoic acid (TNBA), thioaminobenzoic acid (TABA) protected-AuNP of 1.4 nm diameter that showed diffusion abilities inside living cells after grafting with bioactive peptides [23]. This type of AuNP appeared to be particularly suited for the site-directed conjugation to an IgG at the antibody’s hinge region. The hinge region of an IgG connects the complement-activating Fc domain to the antigen-binding (Fab) domain and contains disulfide bonds that can be selectively reduced to liberate nucleophilic thiols. These liberated thiols can then react with electrophiles, thereby forming covalent bonds [24, 25]. Moreover, they can also exchange with the ligands of thiolate-protected AuNPs [26]. When the antibody is tagged at the hinge area, the antibody functionality is generally untouched since the Fab and the Fc domain, which both are implicated in the IgG cellular action, remain unmodified [27]. It should be however mentioned that the thiol-specific conjugation of AuNPs [28] and thiolate-protected AuNPs [26] at the hinge area is not always easy to achieve, likely due to steric hindrance.

In the presented study, we first modified a synthetic protocol for making a TNBA-, TABA-protected AuNP of 2.4 nm. Secondly, we evaluated the ability of this AuNP to react with the thiols of reduced IgGs at the hinge region (AuNP synthesis scheme and bioconjugation strategy illustrated in figure 1). As models, we selected the anti-EGFR antibody Cetuximab (Cmab) and the anti-vascular endothelial growth factor (VEGF) antibody Bevacizumab (Bmab). The direct ligand exchange proceeded seamlessly at near stoichiometric ratio and the AuNP-antibody link remained intact, even after addition of the CALNNG peptide in large excess, which served the purpose of exchanging the remaining reactive TNBA/TABA ligands with a passive CALNNG layer [29]. Thirdly, the ability of the AuNP-Cetuximab conjugate to bind to its cellular target was assayed using cell line models. Biological evaluation using living cells with or without cell surface EGFR demonstrated that the AuNP-Cetuximab conjugate behaved very similarly to Cetuximab, despite being tagged with a 2.4 nm AuNP.

2. Materials and methods

2.1. Chemicals

Water was purified with a Millipore Q-POD apparatus. The paraformaldehyde (PFA 16% solution) and the glutaraldehyde (25% solution) solutions were of Electron Microscopy quality grade and purchased from electron microscopy sciences. The jet PRIME siRNA transfection reagent was
2.4. Synthesis of AuNP-antibody conjugate

A 2 mg ml⁻¹ antibody solution (225 μl, 0.45 mg) was treated with a 7 mM Tris(2-carboxyethyl)phosphine-HCl (TCEP) solution, pH 7.0 (90 μl, 0.63 μmol) for 1.5 h at 37 °C. The AuG (73 μl of a 42 μM, 3.06 nmol) was then added to the reduced antibody (297 μl, 0.42 mg) in 0.1 M HEPES buffer, pH 7.5 at 25 °C and the reaction was let to proceed overnight.

The next day the AuNP-antibody conjugate was passivated with a 1 mM solution of peptide CALNNG (123 μl, 123 nmol or 40 molar eq. of AuNP-antibody conjugate) for 4 h at 25 °C in 0.1 M HEPES buffer, pH 7.5. The exchanged AuG-ligands (TNBA and TABA) and excess CALNNG peptides were removed by ultrafiltration using Amicon 100 K ultra centrifugal devices.

2.5. Mass spectrometry analysis

Mass spectra were recorded with a MALDI-TOF MS operating in positive ion mode on an Autoflex™ system (Bruker Daltonics GmbH, Bremen, Germany). The system was used at an accelerating potential of 20 kDa in linear and reflector mode. The nitrogen laser (337 nm) was used at a frequency of 5 Hz and the acquisition mass range was set to 5000–30 000 m/z with a matrix suppression deflection of 500 m/z. Samples were prepared by the dried droplet method. The matrix solution consisted of a saturated solution of α-cyano-4-hydroxycinnamic acid in H₂O/CH₃CN (50:50), which was threefold diluted in H₂O/CH₃CN/TFA (50:49:9.0:1).

2.6. Electron microscopy and EDX analysis

Images of the AuNPs were obtained by performing microscopy experiments using a C₄-corrected JEOL JEM-2100F scanning transmission electron microscope (STEM) operating at 200 keV. Energy dispersive x-ray (EDX) analysis was carried out on the same instrument, being equipped with an EDX detector. Samples were prepared by adding 10 μl of a 5 μM AuNP solution onto the Carbon filter devices. 3rd step: passivation of the AuNP-IgG conjugate using excess of peptide CALNNG (HS-Cap).

2.7. FTIR analysis

Fourier-transform infrared (FTIR) spectrum of AuNPs was recorded using a Nicolet 380 FTIR spectrometer and a diamond ATR by Thermo Fisher Scientific (supporting...
information, figure S1 is available online at stacks.iop.org/NANO/30/184005/mmedia).

2.8. SDS-PAGE

SDS-PAGE was performed according to a published protocol of Laemmli et al on 10% and 15% acrylamide gels [30]. The gels were pre-run for 20 min in a tris-glycine buffer (0.25 M Tris, 1.92 M glycine, 1% SDS, pH 8.5) at 20 mA. For loading 50% (v/v) glycerol solution was added to the AuNP solutions to a 5% final proportion. After electrophoresis, the AuNPs were seen as black-brown bands. Few amounts of AuNPs could be further visualized by silver enhancement. Proteins were revealed by Coomassie blue staining.

2.9. Cell culture

Cell lines were maintained in a 37°C humidified incubator with 5% CO₂. The human U87 glioblastoma cells (U87 MG, ATCC HTB-14) and the human fibrosarcoma cells (HT-1080, ATCC CCL-121) were maintained in Eagle’s minimum essential medium containing 10% fetal bovine serum, 1% sodium pyruvate and 1% nonessential amino acids. Human foreskin fibroblast (HFF) cells (HFF-1, ATCC SCRC-1041) were cultured in Dulbecco’s modified eagle medium supplemented with 2 mM L-glutamine, HEPES buffer, 10% heat inactivated fetal calf serum (FCS) and 50 μg ml⁻¹ gentamycine. The U87 and HFF cells co-culture was done in Opti-MEM cell culture medium containing 10% FCS. The EGFR (+) U87 cell line was a gift from Professor Furnari [31]. The MTT assay was performed according to a published procedure [32].

2.10. Downregulation of EGFR expression in U87 cells

Expression of EGFR was down-regulated using the synthetic interfering RNAs (siRNAs) methodology. The U87 cell line was seeded in 6-well plates at 250 000 cells/well the day before the siRNA transfection experiment. For one well, a 50 nM siRNA solution (200 μl jetPrime buffer, 10 pmol siEGFR) was mixed with 4 μl of jetPrime reagent. After 10 min incubation at room temperature, the complexes were added to the cells by dilution into the cell culture medium. To ensure maximum gene silencing the cells were incubated at 37°C for 30 min This starvation step aimed at optimizing EGFR presentation on the cell surface [34]. Culture medium was then carefully removed and replaced with a serum-free medium containing the AuNP-antibody conjugate. After 30 min of incubation, the cell culture medium was removed. Cells were washed with PBS and then fixed with either 4% PFA in PBS (10 min) or 2.5% glutaraldehyde in Sorenson’s Buffer (1 h).

2.12. Assay of EGFR-mediated endocytosis

Cells were seeded in 24-well plates and let to adhere on fibronectin-coated (20 μg ml⁻¹) glass coverslips the day before the assay. The cell culture medium was then replaced with serum-depleted culture medium and the cells were let in this medium for 30 min at 37°C. After serum-depletion, cells were incubated in ice-cooled serum-free medium containing 167 nM of the AuNP-antibody conjugate. After 30 min of incubation on ice, the cell culture medium was replaced with pre-warmed serum-containing cell culture medium and the cells were incubated at 37°C for different time periods. The cell surface-bound antibodies were detached with a 0.2 M sodium acetate solution (pH 2.7). The cells were then washed with PBS and fixed with 4% PFA.

2.13. Preparation of the cell specimen for AuNP detection

The AuNPs were detected using a modified Danscher method [23, 35]. Briefly, after the 2.5% glutaraldehyde fixation step, the cells were incubated with a 0.1 M Sorenson’s buffer, pH 7.4 containing 50 mM glycine for 20 min. The cell membrane was then permeabilized using a Sorenson’s buffer, pH 7.4 containing 0.05% (w/v) saponine. The buffered solution was then replaced by a 0.1 M citrate solution, pH 6.7 containing 2% (w/v) sucrose. Development of the silver-mediated AuNP staining was stopped by washing the cell specimen with 0.16 M sodium citrate solution, pH 6.7.

2.14. Western blot

Cells were lysed in Laemmli loading buffer, the lysate was fractionated by SDS-PAGE and transferred onto a poly-vinylidene difluoride membrane. The anti-EGFR D38B1, anti-pEGFR Tyr1068 and anti-GAPDH antibodies were used to detect EGFR, phosphorylated EGFR and GAPDH respectively.

3. Results

3.1. Gold nanoparticle synthesis and characterization

We previously described the synthesis of a TNBA-, TABA-protected AuNP of circa 102 gold atoms that could be grafted with thiolated peptides by exchanging most of the
TNBA-ligands, leaving a surrounding zwitterionic protecting shell consisting of gold-coordinated TABAs [23]. In an initial stage, we explored the possibility of preparing the same type of TNBA-, TABA-protected AuNP, but of larger diameter. The nature and proportion of the solvents were seen to dramatically alter the production of thiolate-protected AuNPs [36]. We therefore assayed the reduction of HAuCl₄ with NaBH₄ and DTNB in various co-solvents. It was observed that a HAuCl₄/DTNB/NaBH₄ ratio of 1:1:5.5 in a solvent mixture of CH₃CN/H₂O (80:20) yielded to a AuNP population migrating as a discrete band when subjected to a sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) analysis, suggesting a homogenous population (figure 2(a)). This AuNP population (named AuG) was further characterized by UV–vis spectroscopy (figure 2(b)). Data showed that the absorption gradually increases for decreasing wavelengths. The spectrum contains a hump with a maximum absorption at 520 nm, corresponding to the weak surface plasmon resonance absorption of 2 nm diameter AuNPs [37]. The STEM analysis of the AuNP revealed a homogenous population of spherical particles (figure 2(c)) with a mean diameter of 2.4 ± 0.28 nm (n = 61). The observation of a crystalline lattice at high resolution (inset image in figure 2(c)) confirmed that the metallic core of AuG was massive (Au⁰). A MALDI-TOF mass spectrometry analysis of the AuNP (figure 2(d)) displayed a narrow distribution of masses at 80 kDa, confirming the SDS-PAGE and EM data. By combining the different data and a volumetric density of 19.3 g cm⁻³ for Au, we estimated that the AuNP contains on average about 420 gold atoms and 130 ligands. Further calculations and a test reaction using increasing ratios of a thiol-containing cationic peptide to the AuNP suggested that the ligand to peptide substitution saturates at about 35 exchanges per particle (figure S2, supporting information). EDX analysis was also performed (figures 2(e) and (f)). The spectrum displayed the characteristic peaks of gold (Au₉₀, at 2.12 keV; Au₅₄, at 9.712 eV) along with peaks corresponding to carbon and copper resulting from the carbon film-coated copper grid, on which the AuNPs were deposited for the analysis.

### 3.2. Conjugation to antibodies

The weakly nucleophilic and thiol-free reducing agent TCEP was used to reduce the antibody disulfide bridges [38]. For optimizing the reduction condition, the Cetuximab antibody (Cmab) was incubated with increasing TCEP concentrations and the reactions were monitored by SDS-PAGE analysis using non-reducing conditions (supporting information, figure S3). Data showed that a final 2 mM TCEP concentration produced a complete reduction of the 150 kDa band to the expected 75 kDa band. Cmab was hence reduced with 2 mM TCEP in PBS for 90 min and the water-soluble 2.4 nm AuNP (AuG) was then directly added to the TCEP-reduced antibody mixture at a 1:1.2 (Cmab:AuNP) stoichiometry. The formation of the AuNP-Cetuximab conjugate (Au-Cmab) was monitored by SDS-PAGE using 10% acrylamide gels (figure 3). To enable dual detection of the protein and the AuNP, the gel was firstly stained using Coomassie blue and then silver ions. The conjugation reaction based on the substitution of an AuG ligand with a thiol group of the antibody’s hinge area proceeded seamlessly, which could be concluded from the observation of a black colored 150 kDa band and no remaining band at 75 kDa in the lane of Au-Cmab (figure 3, lane 5). An apparent 250 kDa band was also observed suggesting dimerization of the IgG, but the proportion was estimated to be lower than 10%. We assumed that the major apparent 150 kDa band corresponds to a 1:1 AuNP-IgG conjugate, whereas the 250 kDa species possibly represents either a AuNP-IgG₂ product or an aggregate of two 1:1 AuNP-IgGs. The observation that the electrophoretic mobility of the main Au-Cmab conjugate was similar to the one of unreduced 150 kDa Cmab is puzzling. However, the AuNP migrated within the migration front and not as classical 80 kDa protein. This high electrophoretic mobility likely results from the high volumetric mass density of gold (19.3 g cm⁻³) and the electronegative charge of AuG. A small amount of unreduced AuG was still detectable in the crude Au-Cmab solution (figure 3, lane 5, faint band at the bottom of the gel), which likely resulted from the slight excess of AuG used for the reaction. Finally, the released ligands, as well as excess peptides and AuNPs were removed using a 100 kDa cut-off ultracentrifugation device. At the present stage, we were unable to remove all the AuNPs as judged by SDS-PAGE analysis, but obtained a batch with less than 5% of free AuNPs.

The conjugation of AuG to Bmab and the purification procedure were performed in a similar manner, but using a TCEP concentration of 0.1 mM for reduction of the hinge disulfide bonds (supporting information, figure S4).

### 3.3. Biological evaluation of the Au-Cmab conjugate

The ability of the Au-Cmab to bind to EGFR, present on the surface of various cancer cells, was examined using a U87 glioblastoma cell line overexpressing the EGFR [31], hereafter referred to as EGFR(+) U87 cells. The Au-Bmab conjugate, which does not target the EGFR, but the (VEGF), was used as the control. In a parallel control experiment, the EGFR expression of U87 wild type cells was almost abolished using the siRNA-mediated gene silencing technology to obtain EGFR(−) U87 cells (western blot confirming the successful downregulation of EGFR depicted in figure S5, supporting information). The Cmab, Au-Cmab and Au-Bmab were added to living cells at a concentration of 167 nM by dilution into the cell culture medium. After 30 min of incubation, the cells were fixed and each domain of the conjugate was separately tracked (figure 4). The antibody was detected by immunofluorescence (IF) [39] (figure 4(a)). Green fluorescence (IgG) was only observed when Cmab and Au-Cmab were added to EGFR(+) U87 cells. Next, the AuNP moiety was revealed by gold-induced silver staining (figure 4(b)). Analogous to the IF results, the strongest silver staining pattern was only seen for Au-Cmab-treated EGFR(+) U87 cells. Some silver staining was nonetheless observed within the endosomes of EGFR(−) and EGFR(+) cells for Au-Bmab and for Au-Cmab, suggesting that the AuNP domain
Figure 2. Characterization of 2.4 nm AuNP (AuG) (a) structure and SDS-PAGE analysis of AuG (15% acrylamide gel). Structure of organothiolate ligands building the surface coating of AuG are depicted below the nanoparticle: R₁ = TNBA (thionitrobenzoic acid), R₂ = TABA (thioaminobenzoic acid); (b) UV-vis spectrum of AuG (small peak at 520 nm corresponding to weak surface plasmon resonance absorption); (c) scanning transmission electron microscopy image of AuG particles. Inset image in right corner shows magnification (scale bar of main image: 20 nm, scale bar of inset image: 2 nm); (d) MALDI-TOF mass spectrum of AuG (MW\text{obs} = 80 kDa); (e) elemental EDX mapping of AuG (scale bar: 20 nm); (f) EDX spectrum of AuG (C\text{Kα} = 0.277 keV, Cu\text{Lα} = 0.93 keV, Au\text{Mα} = 2.12 keV, Cu\text{Kα} = 8.04 keV, Cu\text{Kβ} = 8.9 keV, Au\text{Lα} = 9.71 eV).
somehow favors adherence to cell surfaces and subsequent endocytosis. It should be however mentioned that the silver-enhancement procedure is highly sensitive and not a quantitative method.

The ability of Au-Cmab to bind to EGFR-overexpressing cells was confirmed using an other EGFR-expressing cancer cell line (human fibrosarcoma cells, HT-1080; Supporting Information, Figure S6). As previously described for binding experiments using EGFR(+) U87 cells, the Au-Cmab bound to the surface of HT-1080 cells, whereas the control conjugate Au-Bmab did not show this pattern. Here again, we noticed some silver staining of the cells incubated with Au-Bmab, reinforcing the assumption that the AuNP domain slightly promotes adherence to the cell surface and subsequent endocytosis.

These experiments convincingly demonstrated that Cmab and the Au-Cmab conjugate selectively bind to EGFR of living EGFR-presenting cells. We then evaluated the impact of the AuNP on the ability of Cmab to bind to the cell surface receptors. Cmab and Au-Cmab were incubated with the EGFR(+) U87 cells at concentrations ranging from 0.67 pM to 167 nM. The cells were fixed and the cell-attached antibodies were qualitatively detected by IF. An on/off fluorescence detection threshold was used and the on/off detection data were plotted as a function of the initial material concentration (supporting information, figure S7). This rough quantitative analysis showed that the detection of the EGFR onto the cells required 10 times more of the Au-Cmab conjugate, than of Cmab, suggesting that appending the 2.4 nm AuNP at the hinge area may not be fully innocuous.

Next, we assayed the ability of the Au-Cmab to get internalized into cells, as it is described for Cmab [40]. Both compounds (Cmab and Au-Cmab) were incubated with living serum-starved EGFR(+) U87 cells for 30 min on ice to allow for receptor binding, but not for internalization. Afterwards, the sample- and non-serum-containing medium was exchanged for serum-containing cell culture medium and the cells were incubated at 37 °C for 30 and 60 min, to allow internalization. At the end of the incubation, the nanomaterials bound to the cell surface receptors were washed away using a mild acidic treatment [41]. The cells were then fixed, the plasma membrane permeabilized with detergent, and the components detected by IF (figure 5). The time-course experiment showed that binding of Cmab and Au-Cmab to the cell surface receptors is followed by internalization into intracellular vesicular compartments. Although the intracellular fate of Cmab and Au-Cmab was similar, slight differences were observed at the 30 min incubation time-point. Cmab mainly localized into perinuclear compartments, whereas the Au-Cmab was still seen inside vesicles closer to the plasma membrane.

To further examine whether the biological function of Cmab was affected by the conjugation to AuG, we compared the ability of Cmab and Au-Cmab to inhibit EGFR autophosphorylation after induction with EGF. The serum-starved EGFR(+) U87 cells were incubated with Cmab and Au-Cmab together with EGF for 15 min at 37 °C. Afterwards, the cells were lysed and the cell extracts were fractionated by SDS-PAGE to quantify the intracytosolic levels of EGFR and EGFR-pTyr1068 by western blot analysis (figure 6). Data showed that the Au-Cmab inhibited the phosphorylation of EGFR similarly to Cmab [42].

Even though Cmab and the Au-Cmab inhibited EGFR phosphorylation, their addition to EGFR(+) U87 cells at a concentration of 167 nM did not apparently impact the cellular viability, as judged by a MTT assay (supporting information, figure S8).

Finally, we examined whether the Au-Cmab conjugate is able to distinguish between EGFR-overexpressing cancer cells and non-cancerous cells. The EGFR(+) U87 cells were co-cultured with the non-cancerous HFF cells and the Au-Cmab was then added to the cell culture medium. After 30 min the cells were fixed and the presence of the Au-Cmab conjugate was revealed by IF and silver staining (figure 7). The two cell types were easily distinguishable by their cell morphology. EGFR(+) U87 cells (figure 7(a): black arrows) are much smaller and thinner than HFF cells (figure 7(a): red arrow; figure 7(b): cells encircled in red). Only the EGFR(+) U87 cells were engulfing a large proportion of Cmab and Au-Cmab, confirming that the Au-Cmab conjugate might be useful to selectively target EGFR-overexpressing tumor cells while not affecting non-cancerous cells.

4. Discussion

NPs, including AuNPs, can be prepared at various sizes and be equipped with functional organic components, which makes them useful for a multitude of different applications [43–48]. For biological applications, the AuNP size plays a major role. Particles having sizes above 4–5 nm offer the advantage to be easily detectable by electron microscopy and they can also be easily surface-coated with several antibodies using strong non-covalent binding or be coordinated to
organic molecules via an Au–S coordination [49, 50]. However, presentation of a large surface to macromolecules present in the solvent is not without consequence. When AuNPs are mixed with serum, a large protein corona is forming around the AuNPs [51] that can impact cellular interactions [52]. Beside these variations in physicochemical properties, the size plays an important role for the ADME (absorption, distribution, metabolism, excretion) profile of the particles [53–55]. Another parameter that is clearly impacting the ADME profile of NPs, such as elimination from the body, is the particle’s coverage [56–58]. For inorganic non-biodegradable AuNPs, renal excretion should be undoubtedly favored, giving priority to the development of small AuNPs. Based on the work of Ackerson [59] we have prepared a novel type of AuNPs containing a mixed TABA, TNBA layer of circa 102 gold atoms that showed extremely promising

**Figure 4.** Analysis of the EGFR binding ability of the anti-EGFR Cmab, Au-Cmab and Au-Bmab to living EGFR(+) U87 glioblastoma cells and EGFR(−) U87 cells. (a) Detection of the antibody domain of the nanomaterial by immunofluorescence; (b) detection of the AuNP domain by silver staining. Cells were incubated with 167 nM of antibody or AuNP-antibody conjugate for 30 min at 37 °C. Scale bar: 20 μm.
usage for biological application, due to its abilities to be functionalized with peptides and to be stabilized with zwitterionic ligands [23]. Although we could have used this AuNP for conjugation, we wished to prepare slightly larger NPs for increasing the quantity of gold atoms within the system on the one side, but also to increase the conjugation challenge as bulkiness provides steric hindrance and unspecific interactions [60, 61]. While it is well described that increasing the NP size can increase the formation of protein corona [62], the size-threshold for the occurrence of protein corona for thiolate-protected AuNPs and the associated change in the particle’s physicochemical properties, is unknown. To start answering to this question, we have hence privileged to work with AuNPs of 2.4 nm, instead of with AuNPs of 1.4 nm.

A previous investigation showed that mercaptobenzoic acid-protected AuNPs can be prepared at various sizes by adjusting the type and composition of the solvent mixture used for the particle synthesis [36]. In our case, a solvation of the gold-DTNB complex in an acetonitrile/water (80:20) mixture led to 2.4 nm AuNPs showing a high degree of monodispersity that could be characterized by SDS-PAGE analysis, MALDI-TOF mass spectrometry, STEM, EDX and UV–vis spectroscopy.

The site-directed bioconjugation of the antibodies Cmab and Bmab to the AuG via simple thiolate-for-thiolate ligand exchange proceeded smoothly and could be monitored by non-reducing SDS-PAGE. Until today there are very few reports about the controlled conjugation of large biomolecules to small-sized AuNPs. Ackerson and coworkers attempted the ‘direct’ labeling of cysteine-containing proteins with Au_{144}NPs, however the reaction seemed to require a large excess of NPs, as a large quantity of unreacted AuNPs could be detected on the SDS gels, indicating that the reaction did not proceed as straightforward as it was the case in the present study [26]. The following reason could be hypothesized. The

Figure 5. Assay of EGFR-mediated endocytosis. Cetuximab (Cmab) and AuNP-Cetuximab conjugate (Au-Cmab) were added to living EGFR (+) U87 cells for 30 min (37 °C) at concentrations of 167 nM. The cell endocytosis was then evaluated immediately (image on top: 0’), as well as after 30 and 60 min of further incubation in complete medium (not containing antibody and conjugate samples). The nanomaterial was detected by immunofluorescence. Scale bar: 20 μm.
AuNPs produced by Ackerson et al were coated with mercaptobenzoic acid, while the particles of the present study contained zwitterionic thioaminobenzoate ligands. This zwitterionic coating might diminish unspecific associations between the NP and the biomolecule, thereby favoring the accessibility of the AuNP to the antibody’s hinge thiols and consequently the S2-like substitution.

The ability of Au-Cmab to selectively bind to EGFR present on living cells was assayed using U87 glioblastoma cells that were engineered to overexpress the EGFR, as well as using the human HT-1080 fibrosarcoma cell line, which also overexpresses the EGFR. The glioblastoma cell model system was chosen, because 40% of all glioblastoma patients overexpress the EGFR, however the response to any EGFR-based therapeutic treatment is extremely low, an issue, which remains unresolved until today [63–65]. As a consequence, approaches have been developed to use the anti-EGFR antibody Cmab as a cancer targeting agent to deliver active payloads [63]. These active payloads can hence induce cell damage of the targeted cells, without relying on a ‘normal functioning’ EGFR signaling pathway. The data obtained from the EGFR binding assays of the present study showed that the Au-Cmab conjugate selectively binds to the EGFR on living cells in an analogous, but not identical manner than Cmab. The following AuNP-mediated differences were observed. First, an AuNP-mediated endocytosis was noted, suggesting that the 2.4 nm AuNPs slightly bind by themselves to cell surfaces. The association of the AuNP to the cell membrane was moreover promoting a small change in the intracellular trafficking of the Au-Cmab, confirming some AuNP-mediated non-selective associations to cell surface membranes. The relevance of this slight, but apparent difference between Cmab and AuCmab is unclear but deserves careful attention. Finally, the conjugation of the AuNP decreased the apparent binding affinity. However, it should be emphasized that we have not comprehensively optimized the quality of the Au-Cmab conjugate and the magnitude in decrease of binding affinity, which we have observed (10 times difference), might be reduced.

The cell viability of the EGFR(+) U87 cells was not diminished by incubation with Cmab or Au-Cmab. This absence of toxicity has already been reported for cultured glioblastoma cells [66] and we hypothesize that this issue has the same background as the resistance of glioblastoma tumors to EGFR-based therapies.

5. Conclusion

A highly defined 2.4 nm AuNP, displaying an inner metallic core and an Au-S coordinated organic ligand shell, was synthesized by NaBH4 reduction of chloroauroic acid in the presence of the Ellman’s reagent in a 80:20 acetonitrile/water mixture. This 2.4 nm AuNP could be characterized using several methods including MALDI-TOF mass spectrometry, SDS-PAGE, UV–vis spectroscopy, electron microscopy, FTR and EDX analysis, thereby facilitating the reproducibility of production. The AuNP was subsequently functionalized with the anti-EGFR antibody Cmab via a simple thiolate-for-thiolate exchange of the AuG ligands (TNBA and TABA) and the hinge thiols of the AuNPs produced by Ackerson et al were coated with mercaptobenzoic acid, while the particles of the present study contained zwitterionic thioaminobenzoate ligands. This zwitterionic coating might diminish unspecific associations between the NP and the biomolecule, thereby favoring the accessibility of the AuNP to the antibody’s hinge thiols and consequently the S2-like substitution.

The ability of Au-Cmab to selectively bind to EGFR present on living cells was assayed using U87 glioblastoma cells that were engineered to overexpress the EGFR, as well as using the human HT-1080 fibrosarcoma cell line, which also overexpresses the EGFR. The glioblastoma cell model system was chosen, because 40% of all glioblastoma patients overexpress the EGFR, however the response to any EGFR-based therapeutic treatment is extremely low, an issue, which remains unresolved until today [63–65]. As a consequence, approaches have been developed to use the anti-EGFR antibody Cmab as a cancer targeting agent to deliver active payloads [63]. These active payloads can hence induce cell damage of the targeted cells, without relying on a ‘normal functioning’ EGFR signaling pathway. The data obtained from the EGFR binding assays of the present study showed that the Au-Cmab conjugate selectively binds to the EGFR on living cells in an analogous, but not identical manner than Cmab. The following AuNP-mediated differences were observed. First, an AuNP-mediated endocytosis was noted, suggesting that the 2.4 nm AuNPs slightly bind by themselves to cell surfaces. The association of the AuNP to the cell membrane was moreover promoting a small change in the intracellular trafficking of the Au-Cmab, confirming some AuNP-mediated non-selective associations to cell surface membranes. The relevance of this slight, but apparent difference between Cmab and AuCmab is unclear but deserves careful attention. Finally, the conjugation of the AuNP decreased the apparent binding affinity. However, it should be emphasized that we have not comprehensively optimized the quality of the Au-Cmab conjugate and the magnitude in decrease of binding affinity, which we have observed (10 times difference), might be reduced.

The cell viability of the EGFR(+) U87 cells was not diminished by incubation with Cmab or Au-Cmab. This absence of toxicity has already been reported for cultured glioblastoma cells [66] and we hypothesize that this issue has the same background as the resistance of glioblastoma tumors to EGFR-based therapies.

5. Conclusion

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selectively reduced antibody—a site-directed conjugation strategy, which has not been explored before for antibodies and small-sized AuNPs. To minimize the formation of protein corona and to prevent NP aggregation, the Au-Cmab conjugate was passivated with peptide CALNNG in a second step. To demonstrate that the conjugation strategy is generally applicable, the AuNP was also conjugated to the VEGF-targeting antibody Bmab. Besides, the Au-Bmab conjugate served as control in the EGFR binding assays. The conjugation reactions could be readily visualized using non-reductive SDS-PAGE analysis, from which it was assessed that the major conjugation products consist of one IgG and one AuNP. The generated Au-Cmab conjugate behaves similar to Cmab and Au-Cmab concentrations used for incubation: 167 nM. Scale bar in (a): 40 μm, scale bar in (b) : 20 μm.

Figure 7. Evaluation of cell selectivity towards EGFR using a co-culture of EGFR(+) U87 cancer cells and non-cancerous human foreskin fibroblast (HFF) cells. Cmab or Au-Cmab were added to the co-culture by dilution in the cell culture medium. After 30 min incubation at 37 °C the antibody domain and the AuNP moiety were detected by immunofluorescence (a) and silver staining (b). The nuclei were stained in blue (DAPI). (a) HFF cells indicated by red arrow, EGFR(+) U87 cells indicated by black arrow. (b) HFF cells are encircled in red, EGFR (+) U87 cells are encircled in blue (in blank image only). Cmab and Au-Cmab concentrations used for incubation: 167 nM. Scale bar in (a): 40 μm, scale bar in (b) : 20 μm.

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