Aptamer Grafting onto (on) and into (in) Pegylated Gold Nanoparticles: Physicochemical Characterization and In vitro Cytotoxicity Investigation in Renal Cells

Celia Arib1, Serena Milano2, Andrea Gerbino2 and Jolanda Spadavecchia**

1CNRS, UMR 7244, CSFBAT, Laboratoire de Chimie, Structures et Propriétés de Biomatériaux et d’Agents Therapeutiques, Université Paris 13, Sorbonne Paris Cité, Bobigny, France
2Department of Biosciences, Biotechnologies and Biopharmaceutics, University of Bari, 70126 Bari, Italy

Abstract

Gold Nanoparticles (AuNPs) have already a remarkable interest as viable biomedical materials. Additionally, the strategy of using biomolecules to modify their surface properties is a very attractive as it leads to the generation of new nanometric hybrid materials. In this respect, aptamers, functional small single-strand oligonucleotides (DNA or RNA), are ideal candidates for molecular targeting applications since the high affinity to their target molecules. Thus, the urge of new and effective methodologies to graft aptamers on AuNPs is rapidly increasing especially for applications in bioanalysis and biomedicine, including early diagnosis and drug delivery. Here we used two chemical methodologies to conjugate the aptamer (APT) onto pegylated gold nanoparticles (PEG-AuNPs): the carbodiimide chemistry (EDC/NHS) (methodology ON) and the chelation-bond (R-Au bond) (methodology IN). The aptamer's conjugations with the PEG-AuNPs were characterized by UV-Vis absorption, Raman Spectroscopy and transmission electron microscopy (TEM).

In addition, the potential nanotoxicity of the two aptamer-conjugated AuNPs was evaluated on two different renal cell lines, being the kidneys one of the most important site of bioaccumulation upon systemic circulation. Interestingly, the two aptamer-conjugated AuNPs showed different cytotoxicity when exposed to human embryonic kidney (HEK293) and mouse collecting duct cells (M-1), indicating that cell viability has to be taken into account when choosing the proper strategy for NPs production. In conclusion this study provides two effective methods to graft aptamers on NPs and important insights regarding NPs conformation and the relative cell viability.

Keywords: Colloids; Functionalization; Biological interaction; Biomarker; PEG-AuNPs; Raman Spectroscopy; Aptamer; SERS

Introduction

During the last years, the nanoparticles were intensively used in industrial and medical applications [1-3]. Among them, Gold Nanoparticles (AuNPs) have a remarkable interest as viable biomedical materials and the research efforts about this subject are continuously growing due to their unique physical and chemical properties [4]. By using some of the biomolecules to modify the surface properties is a very attractive strategy, as it leads to a new generation of nanometric hybrid materials [5-7]. Several characteristics of such nanostructures have been identified as providing a better biocompatibility and furtivity to the nanoparticles or a better targeting of some specific disease biomarkers for nanosensor development or some of biostructures as tumours or organs that have to be detected or treated [8]. Additionally, the physicochemical characteristics of NPs deeply impact their ability to migrate away from the organism’s entry site [9]. Liver, spleen and kidney possess a fenestrated endothelium that allows for passive accumulation of NPs and are potential sites for NPs off-target toxicity or target of the therapy [10]. In particular, biodistribution studies in vivo confirmed renal distribution of NPs [11-14] and this highlight the need of evaluating nanotoxicity at renal level. One of the most promising bioreceptors is the DNA aptamer, a DNA single strand (ssDNA) specially designed to have a high affinity to proteins [15] or other kinds of analytes as toxins [16,17]. Based on their high binding affinity and specificity towards other molecules, aptamers, can be utilized in applications ranging from biosensing to diagnostics and therapeutics [18-20]. Meanwhile, advances in nanotechnology have led to new and improved materials for biomedical applications [21]. Specifically, nanoparticles can readily interact with both intra- and extra-cellular biomolecules to yield improved signal amplification and target recognition [22]. To accomplish this, several parameters were investigated, including nanoparticle size, conjugation chemistry, use of multiple aptamer sequences on the nanoparticles, and use of multiple nanoparticles with different aptamer sequences [16,18]. In this context the study of the interaction between the bioreceptor (e.g. aptamer) and proteins is essential to improve the biomedical application of nanoparticles [23,24]. Among various analytes of interest, aquaporins (AQPs) are membrane water/glycerol channels that plays a key role in many physiological functions [25], as well as chemical tools to assess AQPs function in biological systems [25,26]. In recent years several gold (III) compounds have shown promising anticanerous effects related to the inhibition of different protein targets, such as the proteasome and specific zinc finger proteins [27,28].

In particular, Soveral G et al. have described the selective and potent inhibitory effect (in the nanomolar range) of Au(III) complexes bearing nitrogen donor ligands on AQPs proteins, which together with their limited toxicity and high water solubility makes them suitable candidates for future in vivo studies [27]. The aim of the current paper, is to preliminary inquire the possibility to exploit the bioconjugation
of aptamer AQP-2 onto gold nanoparticle surface as two different chemical strategies in order to evaluate their biological interactions with kidney cells as well as the chemical physical characterization of AQP2 aptamer onto gold nanoparticles. In particular, two methods for the aptamer grafting onto pegylated gold nanoparticles (PEG-AuNPs) was established: the carbodiimide chemistry [29] (reaction between COOH and NH₂ chemical groups) and, for the first time, the chelation reaction concerning the complexation between Au-Cl and aptamer functional groups (PO₄³⁻, CO and/or NH₂).

Both aptamer-conjugated PEG-AuNPs underwent physicochemical characterization and tested for cytotoxicity on two different renal cell lines, human embryonic kidney (HEK293) and mouse collecting duct cells (M-1), respectively.

This study open the way to develop original protocols will be employed for decoration of hybrid nanoparticles in order to create aptamer-based nanomaterials, with high affinity cancer biomarkers.

Materials and Methods

Tetrachloroauric acid (HAuCl₄), sodium borohydride (NaBH₄), dicarboxylic PolyEthylene Glycol (PEG)-600 (PEG), N-hydroxysuccinimide (NHS), 1-(3-dimethylaminopropyl)-N’-ethylcarbodiimide hydrochloride (EDC) and phosphate-buffered solution (PBS) were all provided by Sigma Aldrich at maximum purity grade. Aquaporin 2 aptamer was purchased from Eurogentec.

Aquaporin 2 aptamer (APT)

Aquaporin 2 Homo sapiens aptamer was purchased from Eurogentec. The aptamer was dispersed in PBS buffer (pH = 9).

Cell culture and in vitro cytotoxicity assay

Human embryonic kidney cells HEK293 cells were cultured in DMEM Glutamax supplemented with 10% fetal bovine serum and 100 i.u/ml penicillin, 100 µg/ml streptomycin at 5% CO₂ and 37°C. Mouse cortical collecting duct M-1 cells [30] were maintained in DMEM/F12 (1:1) supplemented with 5% fetal bovine serum, 2 mM L-glutamine, 100 i.u/ml penicillin, 100 µg/ml streptomycin, and 5 µM dexamethasone. Cells were either left under basal condition or exposed to PEG-AuNPs (1 µM), APT ON PEG-AuNPs (1 µM) and APT IN PEG-AuNPs (1 µM) for 6, 12, 24, 48 and 72 hours at 37°C to evaluate cytotoxicity using trypan blue exclusion assay. Briefly, an aliquot of the cell suspension was diluted 1:1 (v/v) with 0.4% trypan blue and viable cells were counted with a hemocytometer Luna II automated cell counter (Logos Biosystem). The percentage viability was calculated by dividing the number of viable cells by the number of all counted cells. Three independent experiments were carried out. Cell viability data were analyzed using Prism software (version 6.0; www.graphpad.com). Statistical differences between experimental conditions were determined by One-way Anova with Tukey’s multiple comparisons post-test. Data were expressed as means ± SEM and p<0.05 was considered statistically significant.

Synthesis of PEG-AuNPs

Colloids of COOH-terminated PEG-coated AuNPs (PEG-AuNPs) were prepared by a well assessed chemical reduction process according to previously described procedure (Scheme 1) [2].

Determination of PEG-AuNPs concentration

AuNPs concentration was determined by exploiting standard mathematical calculations in colloidal solution as described previously [31].

Bioconjugation of PEG-AuNPs with aptamers

PEG-AuNPs surface was modified by APT aptamer, according to the grafting procedures depicted in Scheme 1.

Aptamer graftings onto peg-aunps by carbodiimide cross-linker chemistry (apt on peg-aunps)

The first grafting strategy consists of the conjugation of the AQP2 aptamer (APT) or on the PEG-AuNPs surface by carbodiimide chemistry (a) and chelation reaction (b) (Please note that drawings are not in scale and are not intended to be representative of the full samples composition).
chemistry. Briefly, 50 μl of EDC/NHS (40/10 w/w ratio) aqueous solution was added into 2 ml PEG-AuNPs dispersion (42 mM). After 2 h, 50 μl of aptamer solution (10 μM in PBS pH 9) was added in 2 ml of the reaction mixture and stirred for 2 h at room temperature. The APT-Peg-AuNPs (6APT/NPs) thus obtained were centrifuged two times at 9,000 rpm for 10 min to remove excess of no-conjugated APT and dried under nitrogen.

Aptamer graftings into peg-aunps by complexation methodology (apt in peg-aunps)

The second grafting strategy consists of the complexation of the AQP2 APT aptamer (50 μl of APT 10 μM; PBS pH 9)(APT) with gold salt (HAuCl4) (5 ml 2.6*10^-4 M), and 50 μl of PEG-diacide (PEG)(1 mM) thought chelation reaction, in order to form APT IN PEG-AuNPs. After 18 h the APT IN PEG-AuNPs suspension was centrifuged twice at 6000 rpm for 10 min to remove excess of aptamer and then the pellets were redispersed in 1 ml MilliQ water. The resultant colloidal solution was sonicated for 5 minutes and then stirred for 1 h at room temperature.

Physico-chemical characterization

All the measurements were performed in triplicate in order to validate the reproducibility of the synthetic and analytical procedures.

UV/Vis measurements: Absorption spectra were recorded using a Perkin Elmer Lambda UV/Vis 950 spectrophotometer in plastic cuvettes with an optical path of 10 mm. The wavelength range was 200–900 nm.

Transmission electron microscopy (TEM): TEM images were acquired with a JEOL JEM 1011 microscope (JEOL, USA) at an accelerating voltage of 100 kV. 2 μl of the particle suspension was placed on a carbon coated copper grid (Smethurst High-Light Ltd) and dried at room temperature.

Raman spectroscopy: The Raman experiments have been performed on an Xplora spectrometer (Horiba Scientifics-France). The Raman spectra have been recorded using an excitation wavelength of 785 nm (diode laser) at room temperature. For measurements in solution, a macro-objective with a focal length of 40 mm (NA = 0.18) was used in backscattering configuration. The achieved spectral resolution is close to 2 cm⁻¹.

Dynamic light scattering (DLS) and Zeta potential measurements: The size and zeta potential measurements were performed using a Zetasizer Nano ZS (Malvern Instruments, Malvern, UK) equipped with a He-Ne laser (633 nm, fixed scattering angle of 173°) at room temperature.

Stability of APT AuNPs (APT ON PEG-AuNPs; APT IN PEG-AuNPs) as a function of pH

For stability studies, APT AuNPs were dispersed in PBS (0.1 M; pH 7 and 5.5), and absorption spectra collected over 3 months (Figure S1).

Aptamer loading efficiency: The amount of the AQP2 aptamer (APT) incorporated onto APT ON PEG-AuNPs and APT IN PEG-AuNPs, were measured by UV-Vis absorption spectroscopy. Absorption at 260 nm was used to extrapolate APT concentrations based on a calibration curve (Figure S2). The aptamer loading efficiency was calculated as follows Equation 1:

\[
\text{APT loading efficiency} = \frac{C_1 - C_2}{C_1} \times 100
\]

Where \(C_1\) is the initial drug content and \(C_2\) is the amount of free

![Figure S1](image-url): Changes in the UV-Vis absorption spectra of APT AuNPs when incubated in buffer solution at pH 7 (A: APT ON PEG-AuNPs; C: APT IN PEG-AuNPs) and pH 5.5 (B: APT ON PEG-AuNPs; D: APT IN PEG-AuNPs) up to three months.
APT in the filtrate after separation of the nanoparticles by ultrafiltration with an Ultrafree MC centrifugal filter units, (30,000 NMWL, Millipore, USA).

**Aptamer release from AuNPs:** APT releases was evaluated at physiological temperature (37°C). aptamer-loaded AuNPs (APT ON PEG-AuNPs; APT IN PEG-AuNPs) were dispersed at concentration of 1.5 × 10¹² particles/ml in 1.5 ml PBS and performed by dialysis as previously described [32]. The concentration of aptamer released from AuNPs was expressed as a percentage of the total aptamer concentration present in the sample Equation 2 and plotted as a function of time.

\[ \% \text{APT released} = \frac{\text{Concentration of aptamer released}}{\text{Initial aptamer concentration}} \times 100 \]  

(2)

**Endotoxin content in drug-gold nanoparticles**

As routine quality control of the materials prior to biological testing, the Pierce LAL Cromogenic Endotoxin Quantitation kit (Thermoscientific, France) was used to determine the LPS contamination in pegylated gold nanoparticles (PEG-AuNPs) before and after conjugation with AQP2 aptamer(APT ON PEG-AuNPs; APT IN PEG-AuNPs) as described previously [33]. Results are reported in Table 1 as average (n=2) ± standard deviation. Endotoxin contamination results were kept into account when defining the type of cytotoxic endpoint to be examined.

**Results and Discussion**

**Formation mechanism of APT AuNPs:** (APT ON PEG-AuNPs; APT IN PEG-AuNPs)

Previously, Spadavecchia et al. have synthetized and conjugated pegylated gold nanoparticles (PEG-AuNPs) with biomolecules (protein, antibody, peptides, aptamer) by several grafting methodologies [2,5,7,31]. Chemical-physical evaluations and biological “in vitro” studies of biomolecules onto pegylated-gold-nanoparticles have successfully showed a large influence in its therapeutic activity [32-35].

Recently, new nano-therapeutic agents based on drugs-gold-complex such as doxorubicin, docetaxel, paclitaxel, gemcitabine has been generated [32,36,37]. According to this experimental approach called “Method IN”, the drug interacts actively with gold salt (HAuCl₄) by chelation bonding and a biocompatible polymer molecules (PEG diacid) as surfactant, in order to form hybrid gold nanoparticles under reduction with NaBH₄.

Other authors conjugated many type of aptamers in order to study the interaction with specific biomarkers [38]. Herein we designed and characterized the grafting of the AQP2 aptamer (APT) at the surface of the PEG-AuNPs for carbodiimide chemistry (method ON) and chelation bond (method IN). In the first case, APT was grafted on the surface of pegylated nanoparticles through the formation of amide links between the COOH groups embedding the surface of the PEG-AuNPs and the NH₂ groups of the APTamer (Scheme 1 panel a previous activation of EDC/NHS). In the second case, APT participate to the stabilization of AuNPs via complexation and electrostatic interaction between its chetone and amino groups with chloride auric ions and dicarboxylic pegylated chains.

In particular, the formation of gold NPs from AuCl₄⁻ includes some mains steps (Scheme 1b):

1. Complexation of APT-AuCl₄⁻ and generation of gold clusters [5].
2. Initial reduction of APT metal complex to form Au II;
3. Staking process of polymer molecules by electrostatic adsorption onto the PEGylayed gold clusters;
4. Final reduction of metal ions to form gold particles stabilized by molecules of APT.

In the first step, a mixture of APT molecules was added in HAuCl₄ aqueous solution to induce formation of complexes. Then, the positively charged APT in water solution shows strong electrostatic interaction with the negatively charged AuCl₄⁻ ions leading to formation of a hybrid complex APT-AuCl₄⁻ (scheme 1b). The addition of PEG-COOH on the hybrid complex stabilizes it through electrostatic interaction between carbonyl and amino groups, improving the kinetics of reduction by complexation of Au ions [39] and controlling the growth process of the final nanoparticles (APT IN PEG-AuNPs). The steric arrangement of APT INPEG-Au clusters before reduction of HAuCl₄ was confirmed by UV-Vis absorption spectroscopy, TEM and Raman Spectroscopy, as reported in the following sections.

**Comparative Aptamer grafting: physicochemical evaluation**

The UV-visible spectra of PEG-AuNPs exhibited a surface plasmon band at 515 nm (Figure 1, black line). After the APT binding onto the PEG-AuNPs (APT ON PEG-AuNPs) via EDC/NHS, the plasmon band is red-shifted to 523 nm (Figure 1, red line). Besides, we assume that
the red-shift amplitude depends on both the type of bioconjugation method and the APT model. Concerning the chelation method, the red-shift is bigger (to 543 nm) with a decrease in intensity and a significant broadening (Figure 1, green line). The red-shift can be displayed by modification of dielectric environment of the nanoparticles thus confirming the successful functionalization of the AuNPs surface in both cases. In particular, the prominent red shift and the broadening observed for APT IN PEG-AuNPs can be due to agglomeration between the pegylated gold nanoparticles combined to the formation of classical van der Waals inter-aptamer interactions [40], thus the different steric conformation of APT during the chelation method. The different red-shift of the plasmon band suggests that each chemical methodology (ON and IN) induce a specific chemical behavior of the APT from PEG-AuNPs. This will have considerable repercussions on the steric conformation of the APT at the PEG-AuNP surface as well as different immobilization kinetic [41,42]. Indeed, the chemical conjugation will influence the grafting position of the APT. With the carbodiimide (EDC/NHS) grafting, the APT is located directly at the gold surface decorated by a PEG layer whereas for the chelation method, the APT will be located into the core of gold, surface embedded of the PEG layer. This chemical behavior is due to a different adsorption onto gold facets of APT, based on their different steric conformation of chemical groups during nucleation and growth process of PEG-AuNPs. Consequently, the color bright violet-blue of the hybrid nanoparticles and the UV-Vis spectra remain unaltered after storage for more than three months at room temperature suggesting the formation of stable colloids suspension. Such observation is very remarkable since the nature of capping layer onto AuNPs plays a key role on the success of bioconjugation and comparative kinetic [5]. This behavior was confirmed by TEM Images. Figure 1-B panel 1 shows a spherical pegylated gold nanoparticles utilized as control with a diameter about 10± 2 nm as described previously [2,5]. After carbodiimide grafting, the resulting APT ON PEG-AuNPs appears as circular agglomerate of nanoparticles embedded of a dense layer of polymer (Figure 1-B panel 2). Considering that the size of single aptamer was about 5 nm [43], we suppose that APT assume a preferential location of PEG molecules onto the particles, with a specific steric arrangement. Contrarily to shape of APT ON PEG-AuNPs, APT IN PEG-AuNPs displays a snow-like shape, embedded in a shell of PEG, with a similar diameter around 50 ± 2 nm (Figure 1-B panel 3. Spadavecchia et al. have realized the synthesis of analogous nanostructures using dicarboxylic PEG [32,33] and drugs, while characteristic snowflakes nanoparticles were obtained.
by incorporating protoporphyrin molecules in the growth solution of AuNPs [44].

According to previous studies [5,32,45], when APT was mixed to gold salt (AuCl₄⁻) complex, the PEG adapt its chemical conformation switching from mushroom to brush mode conformation.

The APT binding is confirmed by DLS and Zeta potential measurements (Table 1). The hydrodynamic diameter is increased by around 10 nm which means an increase of 5 nm of the radius. This distance should correspond to a superior value of the aptamer length since the hydrodynamic radius includes the interaction with water and could not reflect the exact diameter. Moreover, Zeta potential measurements show that PEG-AuNPs and APT PEG-AuNPs were colloidal stable at physiological pH (z-potential = -28 ± 1 MV, -30 nV for APT ON -PEG-AuNPs and -33 ± 1 mV for APT IN PEG-AuNPs with a PdI equal to 0.3) (Table 1). This stability was improved by the presence of PEG coating [2] [42].

The APT AuNPs (APT ON PEG-AuNPs; APT IN PEG-AuNPs) were also characterized by TEM and have a spherical shape (Figure 1B) with a diameter of 10 nm, for PEG-AuNPs confirmed by the hydrodynamic diameter measured by zetasizer (Table 2). The stability of APT AuNPs in solution, was monitored by the Localized Surface Plasmon (LSP) band at 523 nm (APT ON PEG-AuNPs) and 543 nm (APT IN PEG-AuNPs). Analysis was carried out at pH 5.5 and electrolytic conditions. The synthesized APT AuNPs did shows an almost negligible change in the LSP band position over a period of three months (Figure S1). Some differences in terms of LSP were observed between APT ON PEG-AuNPs and APT IN PEG-AuNPs, due to different chemical behavior of APT AuNPs in function of size and shape respectively.

The successful immobilization of AQP2 aptamer onto and/or into PEG-AuNPs surface was also demonstrated by Raman spectroscopy (Figure 2 A-B). Moreover, we have access to some information regarding the aptamer structure with the Raman bands. The APT AuNPs surface was characterized by the APT grafting method impacting the conformation of the aptamer. In particular the peak at 1058 cm⁻¹ typical of phosphate group (PO₄³⁻) and the vibration of the sugar, one at 1536 cm⁻¹ which corresponds to the Adenine base and one at 1731 cm⁻¹ which corresponds to ketones in the different bases of the aptamer [46]. Some bands are specific to the APT as the peak at 1058 cm⁻¹ typical of phosphate group (PO₄³⁻) [46] or the intense peak at 1371 cm⁻¹ which corresponds to thymine base. Its high intensity is due to the good alignment of the aptamer with the Au surface of nanoparticles [49], in contrast to the lower signal recorded for the case of the free thymine DNA in water [50].

If we compare the Raman results of bioconjugation of APT with PEG-AuNPs (APT ON PEG-AuNPs; APT IN PEG-AuNPs) it confirms that the chemical and steric modification of the aptamer depending of the grafting method impacting the conformation of the aptamer. In particular the peak at 1528 cm⁻¹ corresponding to Adenine and NH out plane, is more pronounced in APT IN PEG-AuNPs compared to APT ON PEG-AuNPs, confirming a different steric disposition of APT onto PEG-AuNPs. An effect SERS was observed in the range 300-3000 cm⁻¹ corresponding to vibration Au-O-C, Au-Cl, O-O and 2500 -3500 cm⁻¹ due to C-C aliphatic stretching (Figure 2 panel B). However, for each grafting method, we can assume that the aptamer different packing density and as a consequence its conformation will be largely influence by such parameters.

### Table 1: Endotoxin units detected by means of the LAL Chromogenic Endotoxin Quantitation kit (ND = not detectable; >1 = above detection limit of the assay).

<table>
<thead>
<tr>
<th>Sample</th>
<th>Endotoxin Units (EU/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PEG-AuNPs (control)</td>
<td>ND</td>
</tr>
<tr>
<td>APT ON PEG-AuNPs</td>
<td>&gt;1</td>
</tr>
<tr>
<td>APT IN PEG-AuNPs</td>
<td>ND</td>
</tr>
<tr>
<td>APT</td>
<td>&gt;1</td>
</tr>
</tbody>
</table>

### Table 2: z-potential and hydrodynamic diameter of PEG-AuNPs, APT on PEG-AuNPs and APT IN PEG-AuNPs by carbodiimide and chelation chemistry.

<table>
<thead>
<tr>
<th>Synthetic product</th>
<th>Zeta potential (mV)</th>
<th>Hydrodynamic diameter (nm)</th>
<th>PdI</th>
</tr>
</thead>
<tbody>
<tr>
<td>PEG-AuNPs</td>
<td>-28 ± 1</td>
<td>10 ± 2</td>
<td>0.3</td>
</tr>
<tr>
<td>APT ON PEG-AuNPs</td>
<td>-30 ± 1</td>
<td>10 ± 2</td>
<td>0.3</td>
</tr>
<tr>
<td>APT IN PEG-AuNPs</td>
<td>-33 ± 1</td>
<td>50 ± 2</td>
<td>0.3</td>
</tr>
</tbody>
</table>

### Table S1: Raman band assignment for the APT and vibrational modes in the range spectral 800-1800 cm⁻¹.
In this study, after complexation of AuCl₄⁻ with PEG diacid chains, APT was covalently bound onto complex after reduction of NaBH₄ to obtain APT ON PEG-AuNPs. In this novel formulation APT assume a different chemical and sterical conformation compared to recent synthesis approach in which APT molecules reside inside the AuNP core (APT IN PEG-AuNPs) [32,55]. The successful loading ratio of APT onto PEG-AuNPs, were evidenced by the characteristic absorption peaks at 260 nm from APT (Figure S2). The standard absorption of APT was plotted in the inset of figure (Figure S2) according to UV-Vis absorbance spectra of APT at various concentrations. The loading efficiencies were estimated to be 85% with 8.2 µg present in 2.5 × 10⁻⁴ mol of NPs (data not shown). A sustained APT release was observed in the first 5 h for both nanoparticles (APT ON PEG-AuNPs; APT IN PEG-AuNPs) at pH 4, which is highly favorable for delivery in the cancer cells [56]. We observed that the delivery of APT was favorable when the nano-formulation was obtained by method ON (APT ON PEG-AuNPs). Therefore, we supposed, that the release of APT was checked by a ionic equilibrium between Au (III)-complexes trapped into AuNPs by the hydrophobic interactions between PEG chains and APT. As previously described, the mechanism by which acidic pH triggers drug release is probably associated to the presence of carboxylate groups in the chemical structure of PEG molecules [32]. Such groups become protonated at acidic pH. This reduces the electrostatic interactions occurring between PEG and drugs that stabilize the APT ON PEG-AuNPs structure (Figure 3).

![Figure 3: A) APT release percentage (%) of APT ON PEG-AuNPs and APT IN PEG-AuNPs time in PBS (37°C) at pH = 4 B) Schematic diagram of APT release under pH conditions.](image)

![Figure 4: Cytotoxicity of APT ON PEG-AuNPs and APT IN PEG-AuNPs in renal cells: A) HEK 293 and B) M-1 cells were either left under basal condition (CTR) or exposed to PEG-AuNPs (1 µM), APT ON PEG-AuNPs (1 µM) and APT IN PEG-AuNPs (1 µM) for 6, 12, 24, 48 and 72 hours at 37°C to evaluate NPs cytotoxicity using trypan blue exclusion assay. The cell viability was expressed as percentage of live cells vs total cells and normalized in respect to the CTR for each time point. Data were expressed as means ± SEM of three independent experiments. p<0.05 was considered statistically significant.](image)
We suppose that, the ionic strength and protonation of APT on PEG chains improve the water solubility of the APT molecules, taking advantage of the drug release from PEG-AuNPs. Indeed, a remarkable moiety of APT electrostatically binding onto dissociated complex is released, through PEG chain and diffusion phenomena [32,39].

Cytotoxicity of apt on PEG-AuNPs and apt in PEG-AuNPs in renal cells

Finally, the potential cytotoxic effect of these NPs on two different renal cell lines was evaluated. Human embryonic kidney and (HEK293) and mouse collecting duct (M1) cells [30] were exposed to medium containing vehicle alone, nanoparticles (PEG-AuNPs i.e. control), and nanoparticles produced with the aptamer either within (APT IN PEG-AuNPs) or outside (APT ON PEG-AuNPs) the nanoparticles. Whenever the aptamer was present the final concentration was 1 µM, which was the higher concentration possible according to the initial concentration of the solution containing the NPs. Cell viability was monitored for up to 72 hours. We found significant cell death, both for HEK293 (Figure 4A)

M-1 (Figure 4B) cells, after 24 h of exposure to APT ON PEG-AuNPs. No significant cell death was evident for the other experimental conditions tested. The apoptosis inducer Staurosporin (5 µM for 6 h) has been used as positive control in each experiment (data not shown).

The evaluation of cells viability under exposure of exogenous NPs is extremely important considering that, after NPs administration, they may migrate to different body sites through the systemic circulation. Indeed, several studied demonstrated NPs distribution and bioaccumulation within several organs such as liver, spleen, kidneys, brain or heart [57-59].

A number of recent studies have shown the impact of NPs accumulation in the kidneys. Chen et al. [60] described damage to proximal tubular cells in mice exposed to copper NPs while Wang et al. [61] Observed signs of glomerulonephritis and pathological degeneration in renal proximal convoluted tubules after oral titanium dioxide administration. These evidence together with our in vitro results highlight the urge for a careful risk assessment regarding all bioaccumulation sites within the body.

Here, we showed the ability of APT ON PEG-AuNPs to exert a significant albeit modest cytotoxic effect. However, further in vitro studies are needed to specifically investigate the mechanisms through which APT ON PEG-AuNPs induce cell death after 24 h exposure. The dissection of the cellular mechanisms leading to nanoparticles-induced cell death might help the design of nano-molecules and nano-complexes whose physicochemical characteristic shape, size, chemical or physical interactions do not interfere with cellular health. This will pave the way for a safer use of these new hybrid nanoparticles in biomedical approaches such as drugs delivery and gene therapy. Under this scenario, nanoparticles whose size allows renal filtration might represent a fascinating strategy to specifically target different parts of the nephron segment. In this study, we designed a putative aptamer against AQP2, which is a water channel responsible of facultative renal water reabsorption. Further investigations are further needed to exploit AQP2- nanoparticles based gene therapy to specifically rescue AQP2 physiological functions in a genetic disease whose main hallmark is impaired water reabsorption [62-66].

Conclusion

The success of this study was to provide the design, the synthesis and the characterization of new hybrid nanoparticles based on the incorporation of AQP2 aptamer (APT) PEGylated in gold nanoparticles under two chemical methodologies. Chemical physics characterizations were extensively achieved and fully elucidated the formation mechanism of the nanostructure and the APT conformational changes associated with such processes. Aptamer-conjugated PEG-AuNPs were assessed for in vitro study on two different renal cell lines, human embryonic kidney (HEK293) and mouse collecting duct cells (M-1), respectively. Based on different shape and size of different aptamer based gold nanoparticles, we will provide in the further, a selective study on several renal cells and therapeutic application thought photothermal (PTT) studies.

Reference


