Plasmon properties of multilayer albumin/gold hybrid nanoparticles

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Abstract

Hybrid nanoparticles are mainly based on inorganic (metal, semiconductors) and organic (proteins, polymers) compounds. The inorganic compounds are metallic nanostructures with unique features such as localized surface plasmon resonance (LSPR) bands in the UV–visible–near IR spectral range. The LSPR frequency is extremely sensitive to small changes in the physicochemical environment surrounding NPs and is dependent on NP shape and size. Recently, we reported a novel preparation method of hybrid nanoparticles based on an Albumin multilayer coating of AuNPs, stabilized by radiation-induced crosslinking processes. With this method, we obtained core/shell nanoparticles composed by Au and albumins. The aim of this paper is the description of the plasmon properties of AuNPs and other core/shell gold/protein NPs prepared. UV–vis spectra were used to detect changes of the plasmon extinction of hybrid nanoparticles, in accordance to the Mie Theory. DLS and TEM measures were used to correlate the results obtained. UV–vis spectroscopy proved to be key to characterize protein layer through two parameters: thickness and refractive index. In conclusion, UV–vis spectroscopy can be a very useful quality control methodology to prepare hybrid Au/protein NPs.

1. Introduction

Inorganic NPs have been prepared from different materials, such as metals, oxides, semiconductors and polymers; however, gold NPs (AuNPs) are one of the nanoparticle systems most abundant in the literature [1]. Since the early stages of the ‘nanotechnology age’, AuNPs have been recognized as an excellent scaffold for the fabrication of novel chemical and biological sensors. Given their distinct physical and chemical attributes, availability of established protocols to prepare them at different sizes [2] as well as using ‘green’ reductants such as tea leaves [3] or plant extracts [4], AuNPs are a widespread nanomaterial.

AuNP surface capping using organic molecules, especially those containing sulphur atoms, are very well described in the literature [5]. This property of chemical interaction has been amply used for functionalization of the surface with different biomolecules. It has been used for several decades for preparation of rapid diagnostics tests [5], and it has become one of the most studied nanomaterials for (bio)chemical sensing and diagnosis [6, 7]. An exponential number of reports have been published in the last decade on this topic. A recent review in this field covers more than a thousand scientific papers [8].

Plasmonic nanoparticles are mainly constituted by Au or Ag cores, with unique features such as localized surface plasmon resonance (LSPR) bands in the UV–visible–near IR spectral range and fluorescence quenching. The LSPR frequency is characteristic of the size and shape of nanostructure and is also extremely sensitive to small changes in the physicochemical environment surrounding NPs. Some of the features of light–matter interaction can be explained through the Mie theory, which provides the optical properties of particles size in the nanometer size range and their optical constant. It is important to take into account that the LSPR of AuNPs has also been explained by means of assembly and surface chemistry [9]. Furthermore, AuNPs display localized surface plasmon resonance (LSPR), which can be used for biosensing and dark field imaging [1]. All the optical
properties of AuNPs will render a most relevant building block in nanostructures with two or more functions depending on the requirements of biomedical applications.

Inorganic materials in the composition of NPs confer them attractive features such as improved biomedical diagnosis and imaging sensitivity or magnetoresponsiveness [10–12], which has led to a growing interest in the development of novel ‘theranostic’ nanotechnology devices [13]. However, nanomaterials for in vivo applications require a special surface treatment to become compatible with the human body.

Analysis of interaction of NPs with biological media, such as blood, interstitial fluid or extracellular matrix, rapidly reveals a complex interaction between NPs and the environment. Their surface is rapidly covered by macromolecules, especially proteins, with the formation of a ‘corona’ shell which alters the particle size, stability and in consequence, the surface properties [14]. It has been recognized that the generation of corona shells around inorganic NPs and their dynamics is a very complex process to address [15–19]. Recently a centrifugation-based physicochemical method has been proposed to study the interaction of nanoparticles with proteins without labeling [20].

Several proteins, such as Albumin, are partially modified as a consequence of the strong interaction between some chemical moiety of the protein and the highly structured surface of the AuNPs. Recently, using XANES and molecular modeling simulation, Wang et al have demonstrated that albumin ‘spreads’ onto the inorganic surface and exposes eight internal disulphide bridge to a flat gold surface of Au-nanorod [18, 21].

Protein corona layer has a pivotal role in the final destination of the NPs within the body on account of their interactions the local environment [22]. Therefore, the development of highly structured surface topography onto NP surface is an important issue to address in the near future. In order to reduce the nonspecific adsorption and interactions with other proteins, our goal is to prepare AuNPs covered with a stable protein multilayer coating in which macromolecules preserve their native state. To this end, we select Albumin as building block to prepare the coating layer that could achieve the objectives above.

In a recent paper, we described a novel method of the synthesis of core/shell Au/Albumin NPs by radiation induced crosslinking [23]. Plasmon spectra of these hybrid NPs showed a shift of several nanometers to longer wavelength. This observation drove us to deep insights into these phenomena in order to understand and evaluate the chance of usefulness of this UV–vis spectroscopy to analyze these changes in the protein coating. In this paper, we applied the Mie theory to describe the plasmon properties of AuNPs and other core/shell gold/protein NPs.

2. Materials and methods

Gold (III) auric chloride hydrate (HAuCl₄ ⋅ H₂O) and commercial human serum albumin 20% (HSA) was kindly donated by Laboratorio de Hemoderivados UNC. The commercial serum albumin solution has N-acetyl tryptophan and sodium caprylate as additives and it was used as received. All other reagents were purchased from local suppliers an analytical grade and used as received.

2.1. Synthesis of Au/Albumin nanoparticles (Au/Alb NPs)

AuNPs were prepared according to the Frens method [24], which consists of the reduction of chloroauric acid with sodium citrate. For our assay, 100 ml of 1 mM chloroauric solution were boiled in a clean flask and subjected to continuous stirring. Then, 10 ml of sodium citrate (at 39 mM) were added to the boiling solution and kept for one hour under continuous stirring. This synthesis was carried out in an aqueous medium to obtain a nanostructured system compatible with biological media. The NPs were stored at 4°C in dark. The concentration of AuNPs was determined through UV–vis spectroscopy. The maximum Q₉₀₀ of the LSPR was recorded for serials dilutions of AuNPs in aqueous solutions. Assuming a spherical shape and size for observation of TEM images, the mass of each NP was calculated from the density of ffc gold (19.93 g/cm³) and Ar of atomic weight of gold (197 g/mol). This calculation and the linear fitting of Q₉₀₀ max of dilutions determined the initial concentration of AuNPs (assuming a complete reduction of gold salt).

In the second step, AuNPs (9.81 × 10¹¹ NPs/ml) were dispersed in different concentrations of albumins (HSA) from 0, 2.5, 5, 10 and 15 mg/ml protein. Following that, different amounts of ethanol were added drop-wise onto the protein solution, keeping the temperature at 0°C under constant stirring. Ethanol, Albumin and AuNPs mixtures were irradiated with gamma-rays from a 60 Co source (PISE CNEA-Ezeiza) at a dose rate 1 kGy/h and keeping sample temperature in the range of 5°C–10°C during the irradiation.

AuNPs with a monolayer of albumin are prepared with dispersion of AuNPs in a final concentration of 1 mg/ml protein.
2.2. Determination of size and morphology of NPs

Particle size was determined by means of two techniques: dynamic light scattering (DLS) and transmission electronic microscopy (TEM). For DLS, the samples were measured at 25°C using a 90Plus/Bi-MAS particle size analyzer, with a light source of 632.8 nm and a 10-mW laser. For TEM, samples were prepared by centrifugation at 14,000 rpm for 20 min in three cycles before measured. The measurements were carried out in Centro de Microscopias Avanzadas (CMA)-FCEyN-UBA. All measurements were carried out on days 1 and 30 after sample preparation. Atomic Force Microscopy (AFM) measures were performed with Au/chHSA NPs in Peak Force Mode. The samples were deposited in freshly cleaved mica, dried and imaging in water. Images were obtained using a Dimension Icon in Peak Force QNM (PFQNM) (Bruker®), (Institute of Biophysics, Universidade Federal do Rio de Janeiro, Rio de Janeiro, Brazil). Rectangular silicon tip with a nominal spring constant 0.8 N/m and with tip radius of 5 nm were used during the measurement.

2.3. Experimental and theoretical determination of plasmon spectra

Experimental determination of plasmon spectra was performed in a Shimadzu 1650PC spectrophotometer in the range 300 to 800 nm. Experimental plasmon spectra were parameterized by plasmon peak wavelength ($\lambda_p$) and wavelength at half-maximum ($\lambda_{0.5}$). Theoretical determination of extinction spectra were carried out using Matlab software onto desk computer based on the general Mie theory for single and core/shell spherical NPs. The metal dielectric function entering in Mie calculations was size-corrected for those cases under 10 nm particle radii [25]. For the case of core/shell NPs, extinction efficiency for each particle is defined by Mendoza et al [26]:

$$q_{ext} = \frac{2}{k^2 b^2} \sum_{n=1}^{\infty} (2n - 1) Re \{ a_n + b_n \}$$

where $a_n$ and $b_n$ are Mie coefficients written in terms of Bessel spherical functions, which in turn depend on NPs radii ($a$ and $b$ are radii parameters), $k$ are defined as $2/\lambda_0$ of and the wavelength dependent relative refractive indices of core and shell materials. In the presence of naked AuNPs, the equation \(q_{ext} = \frac{2}{k^2 b^2} \sum_{n=1}^{\infty} (2n - 1) Re \{ a_n + b_n \} \) could be replace for \(2/k^2 a^2\). For metals like gold, the plasmon spectra are represented for radii of $a$ and $b$, where there are wavelength values which, for given radii $a$ and $b$, yield maximum values of $a_n$ and $b_n$ thus producing a peak in $q_{ext}$ which is known as plasmon resonance.

The previous theoretical description holds for single particle size. However, real NPs samples show a size dispersion that is most frequently described by a log-normal function, which is characterized by a modal diameter ($D_m$) and a dispersion parameter ($\sigma$) related to the distribution width. Therefore, the final theoretical extinction spectrum ($Q_{ext}$) is calculated as the weighted sum of $q_{ext}$ for the different finite sizes NPs that define the distribution (see figure 1).

Response Surface Plots (RSP) were prepared from theoretical spectra calculated from log-normal population parameters or shell parameters, according to the proposed core/shell models (see below).

2.4. Electrophoretic analysis

Polyacrylamide gel electrophoresis (SDS-PAGE) was performed with a total of 30 $\mu$l of albumin and NPs diluted in deionized water and 10 $\mu$l of sample buffer with 5% of 2-mercaptoethanol, reaching a final concentration of 1.5 mg/ml of protein. Samples were heated up for 5 min. Following that, aliquots of 18 $\mu$l of the samples and protein markers were loaded on an 8% polyacrylamide gel and 5% stacking gel. A Biopack Electrophoresis Unit was used to run the SDS-PAGE at a constant voltage of 8 mA. Finally, the gel was stained using Comassie Blue and analyzed with Image J Gel Analyzes.

3. Results and discussion

Naked AuNPs are very sensitive to the changes of the environment caused by the addition of different organic solvents, pH, ionic strength and temperature [27]. Those changes easily induce nanoparticle agglomeration so the functionalization of AuNPs is mandatory to prepare useful nanomaterials. The functionalization of the surface of AuNPs using organic molecules containing sulphur atoms or polymers is very extended in the literature [15, 28, 29]. Monolayers of proteins, such as antibodies, serum albumins and other proteins have been prepared by physisorption method and applied to disposable diagnostic devices [30, 31].

AuNPs have been prepared according to Frens method to reach NPs in the range of 20 to 30 nm [24]. This synthesis is carried out in an aqueous medium to avoid the presence of organic solvents or other hydrophobic ligands onto the NP surface. Supplementary figures S1(a) and S1(b) is available online at stacks.iop.org/MRX/6/055005/mmedia show a TEM image and the corresponding histogram of size distribution (mean diameter 23 nm). Supplementary figure S1(c) plots the DLS histogram corresponding to the same sample (average
hydrodynamic size 33 nm). The differences of size measured by both techniques are under the error dispersion of the sample, in addition to the intrinsic differences between techniques. DLS also detected a small fraction of aggregated NPs.

The experimental plasmon spectra of NPs suspensions can be characterized by two parameters: plasmon peak wavelength ($\lambda_p$) and wavelength at half-maximum ($\lambda_m$). Figure 2(a) shows the experimental spectrum corresponding to AuNPs prepared in this work and their parameters. Meanwhile, considering a log-normal distribution of the NP population, theoretical extinction spectrum $Q_{ext}$ is calculated as the weighted sum of the extinction efficiencies ($q_{ext}$) calculated from equation [1]. Population distribution is characterized by a modal diameter ($D_m$) and a dispersion parameter ($\sigma$) (figure 1). A complete set of $Q_{ext}$ were calculated for a range of $D_m$ and $\sigma$ and this information was plotted as a Response Surface Plot (RSP) (see figure 1). Figure 2(b) shows in a 3D graph of surface responses of $\lambda_p$ and $\lambda_m$ of the $Q_{ext}$ as a functions of different NPs populations according to their parameters $D_m$ and $\sigma$. Values of $\lambda_p$ and $\lambda_m$ corresponding to the experimental spectrum shown in figure 1(a) ($\lambda_p = 522$ nm and $\lambda_m = 564$ nm) define two level curves in the 3D plot (figure 2(b)). The intersection of both curves (in the plane x-y) yield unique values $D_m = 26.74$ nm and $\sigma = 0.508$. These values characterize the NP population, which matches the experimental extinction values. This data was used to calculate the $Q_{ext}$ plotted in figure 2(a) (red dashed line). It can be seen that there is a very suitable matching between experimental and theoretical curves. In figure 2(b), in addition to the level curves corresponding to the solution values of $\lambda_p = 522$ nm and $\lambda_m = 564$ nm, those corresponding to $\pm 0.5$ nm for each parameter were included as blue and black curves respectively. In these cases, intersection points differ markedly from central values, illustrating the suitability of the fitting method.

Figure 3 shows a comparison of histograms which describe the naked AuNPs population measured from TEM images (red bars) and DLS measurements (blue bars). Each histogram is fitted with a log-normal curve (red long-dashed curve and blue short-dashed lines respectively). In addition, the black full line represents the size distribution calculated from the optimal parameters fitted, according to the Mie theory and log-normal nanoparticle distribution. The calculated log-normal bell agrees with the experimental data obtained by different techniques.

### 3.1. Protein coating in aqueous AuNPs suspension

The standard preparation of HSA monolayer coating of AuNPs (Au/mHSA NPs) is performed by adding small amounts of protein to the NPs suspension [32, 33]. Albumin spreads onto the gold surface, modifying its shape [18, 21]. Therefore, the surface exposed to the media is different from the native protein. It is our interest to prepare hybrid protein/gold NPs with a more ‘native’ surface resembling the Albumin surface. Our approach involves the preparation of a multilayer protein coating avoiding the surface crosslinking. Therefore, our first step will be to study the AuNPs/Albumin mixture under high protein concentrations. The system will be investigated by UV–vis spectroscopy (see figure 4 scheme of work), DLS and SDS-PAGE.
From the analysis of experimental plasmon spectra of aqueous AuNPs suspension in the presence of HSA, we detected modification of the plasmon peak position and width following increment of protein amount in the AuNPs suspension, in addition to a slight shift to higher wavelengths and changes in the peak height (figure 5(a)). As it was described in the previous section, experimental plasmon peaks are parameterized by $\lambda_p$ and $\lambda_m$ for the different HSA concentration in the media (see table 1).

The addition of different amounts of protein spontaneously generates core/shell NPs, of which the shell properties such as refractive index ($n_c$) and thickness ($\Delta$), are unknown. Furthermore, the alterations of the plasmon extinction are also suggests modifications in this shell structure. Figure 5(a) shows the experimental...
plasmon spectra corresponding to AuNPs suspension added with increasing amount of HSA and the variation of \( \lambda_p \) can be seen in the insert plot. In order to explain the plasmon dispersion of the coated NPs, experimental plasmon peaks will be fitted to a core/shell NP model using the Mie theory, recently described by Mendoza-Herrera et al. [26].

The fitting process involves the preparation of a RSP of the plasmon spectrum parameters, \( \lambda_p \) and \( \lambda_m \), as a function of shell parameters. The initial naked AuNPs population is defined by the log-normal distribution parameters, \( D_m \) and \( \sigma \) values. Protein shell will be defined by two new unknown parameters, \( n_c \) and \( \Delta \), which are the refractive index and thickness respectively.

The growth of protein shell will be considered according to two different coating models. See figure 5 for the two proposed models: the first approximation is a coating thickness proportional to AuNPs radii (Proportional model) and a second one in which it will be considered that all NPs will have the same coating thickness for a particular experimental condition, independent of the naked NP radii (Steady Model).

Considering both hypothesis, two RSP for \( \lambda_p \) and \( \lambda_m \) have been calculated for different values of \( n_c \) and \( \Delta \). In both cases, it was considered that the initial population of naked AuNPs are described by the values \( D_m = 26.74 \text{ nm} \) and \( \sigma = 0.508 \). Figure 5(b) shows the SRP of \( \lambda_p \) and \( \lambda_m \) corresponding to different values of \( n_c \) and \( \Delta \) of protein shell, in this case considered as a percentage of NP diameters (Model 1). In the same way, figure 5(c) shows the response surface for the second model where plasmon dispersion parameters are plotted as a function of \( n_c \) and \( \Delta \). Additionally, both figures show curves (onto the x-y plane) corresponding to solution values of \( \lambda_p = 523.1 \text{ nm} \) and \( \lambda_m = 566.5 \text{ nm} \) and those corresponding to \( \pm 0.5 \text{ nm} \) for each parameter were included as blue and orange curves respectively. These \( \lambda_p \) and \( \lambda_m \) values correspond to AuNPs population in a media containing 5 mg/mL of HSA. The Model 1 does not show a theoretical solution from the curves described in figure 5(b). Only figure 5(c) presents a solution (crossing lines) to the theoretical plasmon spectrum considering a single thickness values. The simulation reaches a solution for \( n_c = 1.345 \) and \( \Delta = 4.22 \text{ nm} \). All \( n_c \) and \( \Delta \) values corresponding to the solutions using different HSA concentrations are described in table 1.

Considering the experimental values which characterize the plasmon peaks, the simulation describes an increment in \( \Delta \) and a relative constant \( n_c \) value. It is to be expected therefore, that increases in the protein concentration media will lead to increments in the corona protein layer.

DLS analyzes of the samples were performed in order to find out additional experimental data to support these predictions. As a consequence of the important differences in the optical properties of both components (inorganic and organic), the light dispersion data of DLS was collected as % Intensity instead of % Relative number. In the analysis of multimodal samples, the former one does not keep the proportionality between populations of different sizes. However, it has been considered for a qualitative analysis as a consequence of bigger-sized particle populations exponentially overvalued the intensity of the dispersed light.
All histograms corresponding to DLS measurements of AuNPs in presence of HSA at high concentrations have shown three main particle populations (figure 6). Those consigned to free BSA ($R_H$ is 3.4 nm, [34]) or small size albumin aggregations ($R_H < 6$ nm); AuNPs coated by albumins; and aggregates. The intermediate size population has an average hydrodynamic size of approximately $44 \pm 9.7$ nm, assigned to a single AuNPs covered with HSA. The relative proportion of each population cannot be determined; however, the monolayer coated AuNPs are present in the aqueous systems and a minor amount of aggregates can be estimated.

The experimental data corresponding to particle size in solution is far more complicated than the predicted values. However, simulation would support the understanding of the main aggregation process involved. The differences in the $\Delta$ values predicted by the fitting data from the plasmon peak are smaller than the error in the

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**Figure 5.** (a) UV–vis experimental spectrum showing the plasmon peak of AuNPs dispersed in different solutions of HSA (0, 2.5, 5, 10 and 15 mg/ml albumin) are corresponded. The inset shows the Peak intensity versus HSA concentration. All samples are in water as solvent (b) Response surfaces of $\lambda_p$ and $\lambda_m$ corresponding de AuNPs with different refractive index ($n$) versus % of NPs diameters corresponding at first hypothesis and (c) refraction index ($n$) versus thickness ($\Delta$) corresponding at second hypothesis. In both cases in the presence of 5 mg/ml HSA.
DLS data, therefore it cannot used to con

firm the predictions. However, the model has been useful to discard the coating process in which the shell thickness is proportional to the NP size (Model 1).

3.2. Plasmon properties of aqueous-ethanol AuNPs suspensions

Low proportions of ethanol in water have a minor effect on AuNPs dispersion. The polarity of R-OH can play the role of the ionic double layer to keep the NP dispersion relatively stable, although not as well as water does [35]. The experimental plasmon peak shifts to high wavelength can be measured only for a solution containing 40% ethanol or higher (figure 7 and table 2). Considering a density of 0.789 g/mL for the absolute ethanol, the mass concentration will be 0.157 8, 0.236 7, 0.315 6 and 0.394 5 for these solutions respectively. From the fitting process, as described in figure 7(b), we determined a shifting of the $\lambda_p$ to higher wavelengths with the increment of the $n_r$ as a consequence of the ethanol concentration increase. However, these increments are lower than 1 nm and it is experimentally relevant at ethanol concentrations higher than 40%.

3.3. Protein coating in aqueous-ethanol AuNPs suspension

The ternary mixture prepared with albumin, ethanol and water is studied considering 30% of ethanol in the mixture. Under this experimental condition, the solvent will not disturb the plasmon peak. Therefore,
increasing concentrations of albumin from 2.5 to 15 mg/ml into the AuNPs suspension, prepared in aqueous ethanol media, were followed by visible spectrophotometry (figure 8). Experimentally, we found a slight shift of the AuNP plasmon peak to higher wavelengths and increased intensity, with the increase of protein concentration (see figure 8 insert). This behavior, which could be associated to the interaction of protein molecules with the gold surface, is independent of the solvent composition. Tsai et al reported an increase of...
plasmon intensity associated to an electrostatic interaction between them [37]. Also, it was observed, in both cases, the intensity of plasmon peak reaches a plate value from 5mg/ml of albumin. Under these experimental conditions the nc of the ethanol/water solution (30% ethanol) is comparable to the nc of the coated protein layer (nc(ethanol 30%) = 1.354 3 versus nc(AuNPs/albumin) = 1.345) (see table 3), therefore the plasmon spectra is similar for different protein amounts (similar values of λp and λm).

However, DLS analysis shows a completely different scenario for these samples (figure 9). The most important characteristic of these histograms is the absence of the free albumin population and low HSA concentrations do not have particles lower than 30 nm diameter. There are records of samples with 10 and 15 mg/ml HSA, with a particle population in the range of 6 to 20 nm, which can be assigned to an aggregation of only albumin molecules. This phenomenon has been recently described for BSA molecules in aqueous ethanol solution [38]. Albumin aggregation can generate small aggregates whose sizes have a direct correlation with the ethanol concentration [38].

The existence of high MW aggregates in the low-concentration samples can be explained by the presence of AuNPs aggregates where there are not enough albumin molecules to produce stable multilayer coating of AuNPs. Meanwhile, samples containing high albumin concentrations show aggregates of much lower MW than the low-concentration ones and in much lower proportion than the medium-size populations.

Table 3. Experimental and fitted data are corresponding to the extinction spectra of AuNPs in aqueous ethanol solution (30% v/v) with the addition of 2.5, 5, 10 and 15 mg/ml HSA. The experimental data λp and λm characterize the extinction.

| mg/ml | λp   | λm   |
|-------|------|------|
| 0     | 524.5| 571  |
| 2.5   | 524  | 566  |
| 5     | 524  | 566.5|
| 10    | 524.5| 567.5|
| 15    | 524.5| 567.5|

Figure 9. DLS histograms (Intensity % versus diameter size) corresponds to AuNPs in aqueous ethanol solution (30% v/v) with the addition of 2.5 mg/ml HSA (top left); 5 mg/ml (top right); 10 mg/ml (bottom left) and 15 mg/ml (bottom right).
Finally, the medium-size particles population has a diameter shifting from aggregates size lower than 100 nm to higher than 100 nm when the HSA increases from 2.5 mg/ml to 15 mg/ml. Meanwhile, the DLS histogram corresponding to 10 mg/ml suggests the presence of two medium-size populations. Table 4 summarizes the average diameters of these populations for the studied conditions. In relation to the sizes, these populations are compatible with NPs containing one AuNP core per particle and increasing amount of albumins in the shell when higher concentration of HSA and ethanol are present in the solution. This assumption can be supported by the simulation findings corresponding to the fact at higher protein concentration, higher shell thickness.

Considering the dynamic aggregation of albumins in aqueous ethanol solutions [38], it is to be expected that the hybrid Au/HSA NPs will be unstable in a media without ethanol. Therefore, a crosslinking process is mandatory to stabilize the suspension.

### 3.4. Protein shell crosslinking

Several chemical and physical treatments have been reported for preparation of albumin micro and nanoparticles [39, 40]. Regarding the physical treatments, our research group has developed a novel crosslinking method of proteins based on ionizing radiation [41]. Gamma rays are ionizing radiation that can induce chemical changing in the irradiated materials, where crosslinking and degradation effects onto the soft mater are achieved. The preponderance of either of the two effects depends on the chemistry of macromolecules, the physical state of the matter and the irradiation conditions. In an aqueous environment, the main effect is degradation; however, a high concentration of ethanol (>6 M) and irradiation conditions used in this work (low dose rate) reduce this effect as a consequence of avoiding the generation of hydroxyl radicals and hydrogen peroxide in the medium [42]. Therefore, a preponderance of crosslinking can be achieved. Additionally, the crosslinking process could be intermolecular or intramolecular depending on the tuning of the experimental conditions [43].

Recent studies have focused on irradiation in a gamma source of the ternary mixture of AuNPs, albumin in a mixture of ethanol and water [23]. Spherical shape Au/albumin NPs of average diameter of about 60nm was prepared with a core/shell structure. Only one kGy irradiation dose was enough to show changes in the plasmon peak characteristics. In addition, changes in the protein shell stability were shown by reduction in the dissolution rate of gold core under cyanate solutions [23].

Figure 10 shows the plasmon peak of three different Au/albumin NPs preparations. The standard Au/mHSA NPs in aqueous solution; multilayer albumin coated AuNPs on 30% v/v ethanol (Au/HSA NPs) and the radio-induced crosslinking one (Au/cHSA NPs) dispersed in aqueous solution. Au/cHSA NPs have a plasmon peak shift to higher wavelengths with respect to Au/HSA NPs and Au/mHSA NPs samples. The plasmon shift wavelength was 10 nm approximately, in addition to an increment of the peak intensity. In the case of multilayer coated NP (Au/HSA NPs) in ethanol/water mixture, we did not observe any change in the plasmon peak (see above). Table 5 depicts the data fitted (Δ and n,) from the spectra of the figure 10 (\(\lambda_p\) and \(\lambda_m\)) using the model described previously according to the Mie theory. In relation to the output fitted values, NPs coated with albumin without irradiation reached the same refractive index and have shown a significant shell thickness difference (longer ones in the presence of ethanol). Meanwhile, the irradiated sample shows an increase in both parameters. In the case of refractive index, the increment could be due to changing the chemical structure of the protein as a consequence of the ionizing radiation (table 5).

Considering the amount of initial protein concentration in the preparation of Au/cHSA NPs, the plasmon signal of the irradiated samples shows a maximum at 2.5 mg/ml of HSA in the suspension (Supplementary figure S2). This behavior is different to that determined for the non-irradiated samples. The irradiation process disrupts the dynamic aggregation of proteins onto NPs surface in the presence of ethanol, as a consequence of a crosslinking process induced by the reductive radicals from the solvent radiolysis, as it was described previously [38].

| Concentration of albumin (mg/ml) | DH (nm) in water non-irradiated | DH (nm) in 30% V/V EtOH non-irradiated |
|---------------------------------|---------------------------------|----------------------------------------|
| 2.5                             | 38                              | >5000                                  |
| 5                               | 35                              | >5000                                  |
| 10                              | 57                              | 149                                    |
| 15                              | 44                              | 120                                    |

Table 4. Hydrodynamic diameter of AuNPs/HSA aggregates in water and aqueous ethanol solution (30% v/v) determined by DLS in the presence of different amounts of protein.
Figure 10. UV–vis spectra showing the plasmon peak of AuNPs covered with a monolayer of HSA prepared by a standard method (Au/mHSA NPs); Au/HSA NPs with a multilayer coating without crosslinking (non-irradiated) and stabilized by radiation-induced crosslinking (Au/cHSA NPs). All samples are in water as solvent.

Table 5. Experimental and fitted data corresponds to the extinction spectra of Au/mHSA NPs (monolayer coated); Au/HSA NPs (multilayer coated without crosslinking (non-irradiated)) and stabilized by radiation-induced crosslinking (Au/cHSA NPs). The experimental data $\lambda_p$ and $\lambda_m$ characterize the extinction spectra and $n_c$ (refractive index) and $\Delta$ the calculated protein layer.

| NPs      | Coating  | Solvent  | Irradiation | $\lambda_p$ | $\lambda_m$ | $D_{av}$ (nm) | $n_c$ |
|----------|----------|----------|-------------|-------------|-------------|---------------|-------|
| Au/mHSA  | Monolayer| Water    | No          | 523         | 567.5       | 6.3           | 1.34  |
| Au/HSA   | Multilayer| Ethanol/Water | No      | 523.5       | 569         | 8.5           | 1.34  |
| Au/cHSA  | Multilayer| Water    | Yes         | 532         | 587.0       | 10.7          | 1.45  |

Figure 11. DLS histograms (Intensity % versus diameter) corresponding to Au/cHSA-NPs in water. These samples were prepared in aqueous ethanol solution 30% v/v with the addition of 2.5 mg/ml HSA (top left); 5 mg/ml (top right); 10 mg/ml (bottom left) and 15 mg/ml (bottom right).
The Au/cAlb NPs size was determined in aqueous solution by DLS. Considering the representation of DLS, the scattered intensity has been chosen on account of the heterogeneity of the different particles present in the system, some of which are pure protein and others have different ratios of protein and gold. Therefore, in order to avoid additional uncertainties of peak areas, the most straightforward variable was selected. Figure 11 shows the DLS histograms corresponding to systems prepared with HSA at 2.5, 5, 10 and 15 mg/ml. In most of the cases, two main populations were found. The average size of Au/cAlb NPs at 2.5 mg/ml is 52 nm. In other concentrations, the NPs size was slightly raised accordingly to increase albumin concentration. Those results confirmed the presence of albumins multilayer to generate an increase of NP size as it was found for Au/HSA NPs in ethanol water mixture (figure 9). Histograms at concentrations from 5 mg/ml to 15 mg/ml show the presence of albumins aggregates and/or nanoparticles (Alb NPs) and Au/albumin NPs aggregates. These results were confirmed by purification of NP samples by centrifugation and Size Exclusion Chromatography.

The presence of covalent protein crosslinking was evaluated by SDS-PAGE. Different NP samples were run in 8% acrylamide gel (2.5% stacking). Figure 12 shows the direct and Coomassie-dyed images of the stacking zone and the Coomassie-dyed running gel. Most of the AuNPs kept in seeding slots and naked and monolayer coated ones shift the color from red to blue (aggregation effect). The only exception is the Au/cHSA, which keeps its initial color. Albumin from Au/mHSA NPs runs in the gel as the HSA standard sample, and in the case of Au/cHSA NPs, a spread of proteins was found in the range of the high molecular weight. These multiple and random crosslinking effect found in the gel could be compatible with a non-specific radiation-induced crosslinking process. A densitometry analysis of the line corresponding to Au/cHSA NPs reveals a 34% of the protein is cross-linked.

3.5. Microscopy analysis
The NP shape of core/shell distribution of these NPs was determined by TEM. Proteins were visualized by staining with uranyl acetate. Figure 13 shows TEM image and its corresponding frequency histogram of Au/cHSA NPs. The presence of a halo around black points assigned to the albumin multilayer shell is found in all NPs. The average of NPs sizes determined from TEM data were 50 nm for Au/cHSA NPs. Taking into account the AuNPs average size of 33 nm, we speculated that one or two fold-size increase were due to one or two albumin layers.

An AFM image capture in solution shows a 3D shape of the protein shell. Figure 14 shows different kinds of structures in the plot: small tiny structures which are attributed to albumin (and small albumin aggregates); medium size spheres corresponding to AuNPs displaying very smooth surface and the big spheres attributed to the Au/cHSA-NPs. The latter have irregular surfaces which are assigned to the heterogeneity of the crosslinking process and the amount of protein in the shell.
4. Conclusions

Plasmon extinction spectra of AuNPs under different experimental conditions have been properly fitted using the Mie theory and a log-normal distribution of the NPs population. We have used a model of a single spherical NP in a surrounding media or a single NP coated with a layer of a soft material (protein layer) in aqueous or aqueous/ethanol solutions. The protein layer has been characterized by two parameters, thickness and refractive index under the assumption that the NP has a spherical shape. The theoretical results arising from the fitting process of naked AuNPs have an excellent correlation with TEM, DLS data in addition to other data available from literature. In accordance to the core/shell simulations, the fitting process was able to discriminate between

Figure 13. (a) TEM picture of Au/cHSA-NPs by negative tincion with uranil acetate; (b) Histogram corresponding to the TEM image; (c) LDS histogram of the same sample in water. The concentration of HSA in samples was 15 mg/ml of protein. This samples was purified by centrifugation before at measured.
different coating models and explain different experimental conditions. The Steady Model fits with the experimental plasmon spectrum of hybrid NPs and is also in accordance with our hypothesis about protein aggregation around AuNPs in the presence of ethanol \[^{38}\].

As predicted by the experimental and simulations data described in this work, UV–visible spectroscopy is a very useful technique to characterize the hybrid NPs and could be proposed as a quality control methodology to prepare these nanomaterials.

Additional characterization of the hybrid NPs has been extended by TEM and AFM microscopy and electrophoresis, confirming the core/shell structure of these novel nanoparticles.

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Figure 14. Image in Liquid AFM of Au/cHSA NPs. The concentration of HSA in samples was 15 mg/ml of protein. The measured are performed in aqueous solutions at room temperature.
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