Research Article

Interaction of Solid Lipid Nanoparticles and Specific Proteins of the Corona Studied by Surface Plasmon Resonance

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The applications of pharmaceutical and medical nanosystems are among the most intensively investigated fields in nanotechnology. A relevant point to be considered in the design and development of nanovehicles intended for medical use is the formation of the “protein corona” around the nanoparticle, that is, a complex biomolecular layer formed when the nanovehicle is exposed to biological fluids. The chemical nature of the protein corona determines the biological identity of the nanoparticle and influences, among others, the recognition of the nanocarrier by the mononuclear phagocytic system (MPS). After exposure of nanocarriers to biological fluids, biomolecules are rapidly adsorbed onto their surface forming the protein corona. The composition and characteristics of solid lipid nanoparticles (SLN) coated with polymers of different chemical nature (e.g., polyvinyl alcohol, chitosans) are reported. The proof-of-concept for the use of SPR technique in characterizing protein-nanoparticle interactions of surface-immobilized proteins (immunoglobulin G and bovine serum albumin, both involved in the formation of the corona) subjected to flowing SLN is demonstrated for non-chitosan-coated nanoparticles. All assayed nanosystems show more preference for IgG than for BSA, such preference being more pronounced in the case of polyvinyl-alcohol-coated SLN.

1. Introduction

Pharmaceutical and medical applications of nanosystems represent a growing area in the flourishing field of nanotechnology, including developments in advanced pharmaceutical nanocarriers and nanodiagnostics [1–3]. Nanosized carriers of very diverse composition are being actively investigated, including metallic, polymeric, ceramic, and lipidic nanoparticles and also hybrid systems by combining such materials [4, 5].

One of the main limitations of these kinds of delivery systems is their short circulation time due to the opsonization phenomenon and subsequent uptake by the mononuclear phagocytic system (MPS). After exposure of nanocarriers to biological fluids, biomolecules are rapidly adsorbed onto their surface forming the protein corona. The corona composition determines the biological identity of the nanoparticle and influences its toxicity, biological fate, circulation time, cell uptake, and drug release kinetics [6, 7]. At first, the protein corona includes considerable amounts of low-affinity proteins which abound in the biological medium. Over time, the composition of the corona is enriched with high affinity proteins [7]. Interestingly, some of the adsorbed proteins are opsonins which promote the uptake and degradation of the
nanoparticles by the cells of the mononuclear phagocytic system (MPS) (mainly located in the liver, spleen, and lymph nodes). Among the proteins commonly found in the corona, IgG and proteins involved in the complement pathway typically act as opsonins (promoting uptake by MPS) whereas serum albumin and apolipoproteins enhance circulation time [8]. The clearance of the nanoparticles by the MPS reduces the bioavailability of the encapsulated drug [9, 10]. A considerable amount of evidence, however, indicates that this phenomenon can be reduced by coating the nanoparticles with different polymeric substances [11–13].

Lipid-based nanosystems stand out because of their nontoxic, biocompatible and environmentally friendly composition. In fact, most of the pharmaceutical nanosystems that have received approval from national health authorities and are already in use in the clinics are of lipidic nature (liposomes) [14]. Solid lipid nanoparticles (SLN), consisting in a solid lipid core matrix stabilized by surfactants, are of particular interest to encapsulate lipophilic drugs [15]. SLN are not only versatile with respect to their composition, but it is also possible to regulate both their particle size and surface characteristics [15]. They are relatively easy to prepare with “green” procedures, resulting in a final product that displays good physicochemical stability compatible with long-term storage, lyophilization, and sterilization processes [16, 17].

Surface Plasmon Resonance (SPR) is a label-free detection method which is particularly suitable for biomolecular interactions studies, allowing high sensitivity measurements in real time. This optical technique is based on the changes that occur in the refractive index due to modifications in the chemical composition in the proximity of thin metal layers (typically gold and silver) that constitute the sensor chips of the SPR equipment [18–21]. The study of protein–nanoparticles interactions by SPR can be performed in two ways; one entails the flow of proteins onto NP immobilized on the SPR sensor surface and the other involves the immobilization of the protein and the injection of nanoparticles throughout the fluidic system. A higher sensitivity is expected for the latter option due to the higher mass of the NP providing that the binding of the proteins to the sensor surface does not alter their typical configuration regarding protein/NP interactions. Previous SPR studies on the interactions of nanovehicles with proteins of the bloodstream include the investigation of the corona formation by employing serum or plasma [22, 23] and the interaction of nanoparticles with individual proteins known to adsorb onto the surface of nanocarriers [24], an approach that has been considered for the present experimental design.

In this work the synthesis of SLN by ultrasonication of cetyl esters mixture Crodamol™ SS (melting point ~ 44°C) was kindly donated by Croda (Argentina). Pluronic® F68, polyvinyl alcohol (PVA, mol wt = 13–23kDa, 98.0 hydrolyzed), low and medium molecular weights chitosans (50–190 kDa and 190–310 kDa, resp., 75–85% deacetylated), Bovine Serum Albumin (BSA, MW = 66.5 kDa.), 11-mercaptoundecanoic acid (MUA), N-hydroxysuccinimide (NHS), ethanamine hydrochloride, and N-(3-dimethylaminopropyl)-N′-ethylcarbodiimide hydrochloride (EDC) were purchased from Sigma-Aldrich (St. Louis, Mo, USA). Potassium chloride, sodium chloride, sodium dihydrogen phosphate, and disodium hydrogen phosphate were obtained from J. T. Baker (Pasadena, Ca, USA). Sodium carbonate was purchased from Biopack (Buenos Aires, Argentina). Commercial gold substrates (SPR102-AU) were obtained from BioNanos™ (Tampere, Finland). The IgG antibody used in the experiments is monoclonal IgG1 anti-DNP 112D5 [25] and it was kindly provided by Drs. Guillermo Docena and Martin Rumbo from IIFP (CONICET-UNLP), La Plata (Argentina).

2.2. Synthesis of Solid Lipid Nanoparticles. SLN were prepared by the ultrasonication method [26]. Briefly, 400 mg of lipid (2.0% w/v) was melted in a water bath at 70°C. Later, hot aqueous solution (70°C, 20 ml) containing 3% (w/v) of Pluronic F68 was added to the lipid phase. Immediately, the mixture was ultrasonicated for 30 min (50% amplitude) using an ultrasonic processor (130 W, Cole-Parmer, USA) equipped with a 3 mm titanium tip. Then, the dispersion was cooled down at room temperature and stored at 5°C [27, 28]. The obtained SLN formulation is denoted, from here, P188. The same synthesis, but using 4.5% (w/v) of Pluronic F68, yielded a SLN formulation called P188+. Three SLN formulations including coating polymers were prepared, by adding polyvinyl alcohol, low molecular weight chitosan, and medium molecular weight chitosan to the aqueous phase (called PVA, CL, and CM, resp.). In the case of SLN formulations CL and CM, the procedure involved the dissolution of 2.0% (w/v) chitosan in 0.1% acetic acid at pH 4.0 and the preheating to 70°C before the addition of the polymer to the aqueous phase. The compositions of all formulations are shown in Table I.

2.3. Particle Size, Zeta Potential (ζ), and Polydispersity Index (PDI). The mean diameter and size distribution were measured by photon correlation spectroscopy (PCS) (Nano ZS Zetasizer, Malvern Instruments Corp, UK) at 25°C in polystyrene cuvettes with a path length of 10 mm. The zeta potential was determined by laser Doppler anemometry also
using the Nano ZS Zetasizer. Measurements were performed in capillary cells with path lengths of 10 mm, using deionized water obtained from a Milli-Q system and phosphate buffered saline (PBS) as diluting media. Additional measurements were accomplished in SLN formulations preincubated in Fetal Bovine Serum (FBS) during one hour [29]. In all cases the measurements were performed in triplicate.

2.4. Transmission Electron Microscopy (TEM). The nanoparticle dispersion was ten times diluted with ultrapure water and a drop of the dispersion was spread onto a collodion-coated Cu grid (400 mesh). Liquid excess was drained with filter paper. One drop of phosphotungstic acid was added to the dispersion, for contrast enhancement. Finally, TEM analysis was performed using Jeol-1200 EX II-TEM microscope (Jeol, MA, USA.).

2.5. Preparation of the Sensor Surfaces. Single thiol SAMs were obtained by overnight incubation of the gold substrates in 50 μM MUA ethanolic solution at room temperature. The obtained surfaces were washed with Milli-Q* water and dried in a stream of nitrogen before SPR measurements.

2.6. SPR Measurements. Measurements were performed with a SPR Navit™ 210A (BioNavis) instrument. The setup is equipped with two incident laser wavelengths, 670 nm and 785 nm, two independent flow channels, and inlet tubing and outlet (waste) tubing. In this work, both of the flow channels were measured in parallel with 670 nm incident light. The measurement temperature was kept constant at 22 °C, and the flow rate used for protein immobilization and for monitoring the SLN interactions with the immobilized proteins was 10 μL/min.

The working mode was “in parallel,” by immobilizing IgG in one flow cell and BSA in the other. The immobilization of the proteins to the single SAMs was performed by the well-known carbodiimide coupling reaction [30]. The first step is the activation of the MUA-carboxyl groups on the surface with a mixture of 0.1 M EDC and 0.1 M NHS to give reactive succinimide esters. IgG 50 μg/mL in 10 mM sodium acetate pH 4.6 and BSA 100 μg/mL in 10 mM sodium acetate pH 4 are then passed through flow cells 1 and 2, respectively, over the surface, and the esters react spontaneously with primary amine groups of the proteins. The immobilization process concludes by blocking remaining succinimide esters using 1 M ethanolamine, pH 8.5. A simplified scheme of the sensor surface is shown in Figure 1. The same schedule of carbodiimide coupling reaction, but flowing buffer solution instead of protein solution, was performed to obtain control surfaces.

SLN solutions (2 mg/ml in PBS pH 7.4) were passed during 10 minutes followed by an injection of PBS pH 7.4 as running buffer and, to remove the bound SLN from the sensor surfaces, 1-minute regeneration steps with Triton X-100 1%. The SPR signal assigned to each SLN solution corresponded to the plateau of the SPR response in the time in between the end of the SLN injection and the start of the regeneration step. Experiments were performed in triplicate.

Table 1: Composition of the aqueous phase of SLN formulations*. In all cases, the lipid used was Crodamol SS.

<table>
<thead>
<tr>
<th>Formulation*</th>
<th>Aqueous phase components</th>
<th>Composition (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SLN P188</td>
<td>Pluronic F68</td>
<td>3</td>
</tr>
<tr>
<td>SLN PVA</td>
<td>Pluronic F68 : PVA</td>
<td>2.5 : 0.5</td>
</tr>
<tr>
<td>SLN CM</td>
<td>Pluronic F68 : MMWC</td>
<td>2.5 : 0.5</td>
</tr>
<tr>
<td>SLN CL</td>
<td>Pluronic F68 : LMWC</td>
<td>2.5 : 0.5</td>
</tr>
<tr>
<td>SLN P188+</td>
<td>Pluronic F68</td>
<td>4.5</td>
</tr>
</tbody>
</table>

3. Results and Discussion

3.1. Synthesis and Characterization of the SLN. SLN of different chemical composition with a common lipidic core composed of cetyl esters mixture and dispersed in poloxamer 188 as emulsifying agent were prepared by sonication method and, in some cases, modified by polymer coatings with PVA or chitosans. The mean diameter, zeta potential, and polydispersity index of the prepared SLN formulations were measured in aqueous solution and in PBS 1x as diluting medium (Tables 2(a) and 2(b), resp.). The particle size of the SLN ranges between 140 and 200 nm, being comparable to the mean diameter measured in water or PBS in the case of P188+, CL, and CM. An increase in the particle size when the SLN is measured in PBS is observed in P188 and PVA, probably due to the change in the hydrodynamic diameter of these nanovehicles in the high ionic strength medium. The polydispersity index of the obtained nanoparticles ranged between 0.14 and 0.23, verifying in all cases the small polydispersity of the SLN solutions. It is well known that size influences both cell uptake and drug release kinetics in nanoparticles and other particulate systems [31–33]. Accordingly, it is desirable to attain low polydispersity since this is expected to result in a more predictable and homogeneous behavior within the targeted biological system.

In relation to the surface charge of the formulations, P188, P188+, and PVA appear to be negatively charged in both media, which is an interesting feature to maintain their stability and avoid possible aggregations. On the other side, significant changes in zeta potential values for chitosan-coated nanoparticles are observed by changing deionized water for PBS in PCS measurement. Both CL and CM are characterized by positive zeta potentials in deionized water, in accordance with the positive surface charge expected from the chemical polymer structure of the coating. However, determination of zeta potential in PBS reflects a shift to more neutral values, which can be explained considering the
Table 2

(a) Mean diameter (Z-average), polydispersity index (PI), and zeta potential (ζ) of different formulations of solid lipid nanoparticles measured in deionized water. Data shown as mean ± standard deviation, n = 3.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Mean diameter (nm)</th>
<th>PDI</th>
<th>ζ (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SLN P188</td>
<td>199 ± 3</td>
<td>0.10 ± 0.03</td>
<td>−6.4 ± 0.9</td>
</tr>
<tr>
<td>SLN PVA</td>
<td>161 ± 1</td>
<td>0.213 ± 0.006</td>
<td>−1.5 ± 0.1</td>
</tr>
<tr>
<td>SLN CM</td>
<td>154 ± 3</td>
<td>0.183 ± 0.002</td>
<td>22 ± 1</td>
</tr>
<tr>
<td>SLN CL</td>
<td>151 ± 3</td>
<td>0.18 ± 0.02</td>
<td>13 ± 2</td>
</tr>
<tr>
<td>SLN P188+</td>
<td>142 ± 1</td>
<td>0.154 ± 0.008</td>
<td>−1.1 ± 0.1</td>
</tr>
</tbody>
</table>

(b) Mean diameter (Z-average), polydispersity index (PI), and zeta potential (ζ) of different formulations of solid lipid nanoparticles measured in PBS as diluting medium. Data shown as mean ± standard deviation, n = 3.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Mean diameter (nm)</th>
<th>PDI</th>
<th>ζ (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SLN P188</td>
<td>207 ± 2</td>
<td>0.16 ± 0.02</td>
<td>−2.8 ± 0.5</td>
</tr>
<tr>
<td>SLN PVA</td>
<td>174 ± 1</td>
<td>0.237 ± 0.009</td>
<td>−6.1 ± 0.1</td>
</tr>
<tr>
<td>SLN CM</td>
<td>152.3 ± 0.3</td>
<td>0.175 ± 0.007</td>
<td>0.7 ± 0.5</td>
</tr>
<tr>
<td>SLN CL</td>
<td>146.5 ± 0.8</td>
<td>0.16 ± 0.01</td>
<td>2.2 ± 0.2</td>
</tr>
<tr>
<td>SLN P188+</td>
<td>142 ± 1</td>
<td>0.16 ± 0.01</td>
<td>−5.2 ± 0.4</td>
</tr>
</tbody>
</table>

Figure 1: Simplified representation of the sensor surface to demonstrate the distinct layers of interest in the sensor surface and the correlation of the interactions that occurs in the sensor surface and the observed sensorgram. Roman numerals indicate (I) the protein layer after immobilization onto the thiol self-assembled monolayer, (II and III) SLN interaction with immobilized proteins at different times of the flow of SLN solutions, and (IV) the interaction of SLN-protein after restoration of buffer flow.
interaction between the positive surface charges of chitosan with negatively charged phosphate ions [34]. This interaction masks the positive charge of chitosan-coated SLN and produces particles with zeta potential values close to neutrality.

More information about the size, distribution, and morphology of SLN was obtained by TEM observations. An effect of increasing 50% of the surfactant concentration during SLN preparation was observed (Figure 2). While SLN P188 formulation exhibited nanoparticles around 200 nm in a relative narrow size distribution, the SLN P188+ preparation produced a bimodal mixture with similar nanoparticles of 200 nm in diameter but with the additional presence of small micelles (mean diameter ≤ 50 nm), possibly due to the surfactant excess. This result is consistent with PCS measure, which indicated that the increase in surfactant concentration produces a lower mean diameter and higher PDI of nanoparticles.

On the other side, the effect of biopolymers addition during SLN preparation was also studied by TEM (Figure 3). The SLN PVA were characterized by nanoparticles in the range of 100–200 nm and covered by PVA layers. Interconnected nanoparticles maybe by PVA bridges are observed (Figures 3(a) and 3(b)). PCS experiments showed that PVA formulation exhibits mean diameters of 161 ± 1 nm in water and 174 ± 1 nm in PBS indicating that these bridges are consequence of the dryness process in the TEM chamber. The morphology, size, and distribution of chitosan nanoparticles seem not to be modified by changes in the MW of the biopolymer. In this sense, spherical SLN with a mean diameter around 200 nm were observed (Figures 3(c)–3(f)).

3.2. DLS Measurements of SLN-Protein Corona Complexes.

In order to investigate the change in the nanoparticle size due to the adsorption of serum proteins, DLS measurements of SLN formulations incubated with Fetal Bovine Serum (FBS), as a model system of biological fluid, were performed (see Figure S1 in Supplementary Material available online at https://doi.org/10.1155/2017/6509184). In all cases, the same increment of the particle size is observed (around 100 nm) when the formulations are in contact with FBS, a finding that can be ascribed to the adsorption of serum proteins over the SLN independently of the differences in surface charge, chemical composition, and molecular weight of the coating in all tested SLN formulations.

The identity of the protein corona is known to play a fundamental role in biological adhesion [35], but due to the complexity of its composition, we propose to study the interaction of the nanovehicles with specific corona proteins like BSA and IgG. Due to the well-known ability of SPR technique to study biomolecular interactions, next we will discuss experimental results involving the immobilization of the proteins on the sensor surface and flowing SLN through the microfluidic system.

3.3. SPR Study of SLN Interaction with Immobilized IgG and BSA.

Typical SPR sensorgrams resulting from the passage of SLN solutions over immobilized IgG and BSA are shown in Figures 4 and 5, respectively. In the figures, the SPR signal corresponding to each formulation is indicated in red. The SPR signal corresponding to the interaction of positively charged chitosan-coated nanoparticles with IgG or BSA, showed in (d) and (e) of Figures 4 and 5, is more intense in comparison with the SPR signal resulting from the other SLN. Even more, the binding of the positively charged chitosan-coated formulations to the immobilized proteins cannot be regenerated with Triton X-100. However, chemical treatment with the nonionic surfactant allows recovering the sensor surface in the case of the other SLN. It is interesting to notice that this finding is independent of the chemical nature of the protein immobilized on the sensor surface, suggesting that the interaction of the SLN with proteins depends strongly on the physicochemical surface properties of the polymer coating. Considering that both immobilized proteins, BSA (isoelectric point = 4.7) and IgG (isoelectric point = 7.0), are negatively charged at the working pH of 7.4 and taking into account the fact that non-chitosan-coated SLN are also negatively charged, it could be inferred that electrostatic forces would not play an important role in the SLN-protein interactions.
Control experiments, flowing SLN over reference surfaces without immobilized proteins, showed negligible interaction in the case of P188, P188+, and PVA formulations (Figure 6). However, the chitosan-coated SLN showed roughly the same SPR signal change in the control experiments and in the measurements involving immobilized proteins. It is well known that due to its chemical nature, chitosan exhibits interactions with several surfaces and strong mucoadhesivity [36–39]. Thus, the interaction of chitosan-coated SLN with the surface in control experiments can be related to the interaction between the hydrophilic surface of the ester formed when MUA was subjected to carbodiimide coupling and the positively charged hydrophilic CL and CM. Another possibility is that in case that some carboxyl activated MUA terminal groups were not blocked by ethanolamine they can react with the nonprotonated amine groups of chitosan. It is well known that amines and hydroxyl groups situated on chitosan chains are responsible for chemical cross-linking. Hydrogen bonding can also contribute to the affinity of CL and CM with the MUA blocked surface.

In order to investigate if P188, P188+, and PVA formulations show preference to interact with IgG or with BSA, it is necessary to make a correction of the SPR signal that gives account of the amount of immobilized protein in each

Figure 3: TEM images showing the effect on SLN preparation by addition of polymers observed by TEM: SLN PVA (a-b), SLN QL (c-d), and SLN QM (e-f) at 40,000 (a, c, and e) and 60,000 (b, d, f) magnifications.
Figure 4: Sensorgrams showing the SPR signal resulting from the injection of the solid lipid nanoparticles (a) P188, (b) P188+, (c) PVA, (d) CL, and (e) CM over immobilized IgG (sensor surface: Au-MUA-IgG). Each cycle included a ten-minute injection of the SLN solution followed by a washing step with running buffer (PBS pH 7.4). In order to remove the bound SLN from the sensor surfaces, 1-minute regeneration steps with Triton X-100 were performed. SPR signal corresponding to each SLN was measured in a time in between the end of the SLN injection and the start of the regeneration step (around ~ 1250 s of each cycle, except CL nanoparticles that the SPR signal was measured at around 750 s).

4. Conclusions

Solid lipid nanoparticles ranging 140–200 nm were obtained by the ultrasonication method. Distinct surface charges characterized the formulations, according to the chemical coating. In all cases a small polydispersity of the SLN solutions could be verified by photon correlation spectroscopy measurements. An increment of around 100 nm in the particle diameter was observed when the SLN formulations are incubated in FBS, finding that it can be ascribed to protein adsorption around the nanoparticles.
The SPR experiments performed in this work verified different proclivity of non-chitosan-coated SLN to interact with IgG with respect to BSA, a fact that can be related to different circulation times in the bloodstream. Due to the complexity and the relative abundance of proteins of the plasma, this hypothesis should be confirmed in future in vivo studies. In the case of chitosan-coated SLN, the proposed strategy did not yield results that could gain information about the affinity of the nanoparticles and the studied proteins. However, the finding that these nanoparticles strongly interact with the sensor surface is in accordance with the fact that chitosan shows affinity with a variety of surfaces.

We demonstrate that SPR is a suitable technique to evaluate the interactions between blood proteins and non-chitosan-coated SLN. In these cases it is possible to obtain information about the different relative affinities of the formulations and the studied proteins, by correcting the SPR signal change for the amount of immobilized protein in the sensor surface. The approach discussed in this paper can be extended to study the interactions of non-chitosan-coated SLN with other components of the protein corona like proteins of the complement pathway and fibrinogen. Therefore, the application of SPR can be considered as a promising tool for the evaluation of SLN-protein interactions.
Table 3: Normalized SPR signal for each SLN formulation (for more detail concerning the calculation see Figure S2 in Supplementary Information). *Ratio between the normalized SPR signal for immobilized IgG and the normalized SPR signal for immobilized BSA.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Au-MUA-IgG</th>
<th>Au-MUA-BSA</th>
<th>IgG/BSA*</th>
</tr>
</thead>
<tbody>
<tr>
<td>P188</td>
<td>0.78</td>
<td>0.13</td>
<td>6.00</td>
</tr>
<tr>
<td>P188+</td>
<td>1.71</td>
<td>0.17</td>
<td>10.05</td>
</tr>
<tr>
<td>PVA</td>
<td>1.44</td>
<td>0.20</td>
<td>7.20</td>
</tr>
</tbody>
</table>

Figure 6: Sensorgrams showing the SPR signal resulting from the injection of the solid lipid nanoparticles (a) P188, (b) P188+, (c) PVA, (d) CL, and (e) CM over control surface without immobilized protein (sensor surface: Au-MUA). Each cycle included a ten-minute injection of the SLN solution followed by a washing step with running buffer (PBS pH 7.4). SPR response arising from the interaction of chitosan-coated SLN and the control surface corresponds to the stabilization of the signal after the end of the SLN injection (at around 1000 s).
in vitro, as well as for predicting the blood clearance of liposomes and likely other nanocarriers in vivo.

Conflicts of Interest

The authors declare that they have no conflicts of interest regarding the publication of this paper.

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