Research Article


Octreotide-conjugated silver nanoparticles for active targeting of somatostatin receptors and their application in a nebulized rat model

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Abstract: Drug uptake and distribution through cell–receptor interactions are of prime interest in reducing the adverse effects and increasing the therapeutic effectiveness of delivered formulations. This study aimed to formulate silver nanoparticles (AgNPs) conjugated to somatostatin analogs for specific delivery through somatostatin receptors (SSTRs) expressed on cells and by nebulizing the prepared AgNPs formulations into lung cells for in vivo application. AgNPs were prepared using the citrate reduction method, yielding AgNPs–CTT, which was further chemically conjugated to octreotide (OCT) to form AgNPs–OCT through an amide linkage. The AgNPs–OCT formulation was coated using alginate to yield a carrier, AgNPs–OCT–Alg, feasible for drug delivery through nebulization. AgNPs were uniform in size with an acceptable range of zeta potential. Furthermore, the concentrations of AgNP formulations were found safe for the model cell lines used, and cell proliferation was significantly reduced in a dose-dependent manner (p < 0.05). In the healthy lung tissues, AgNPs–OCT–Alg accumulated at a concentration of 0.416 ± 5.7 mg/kg tissues, as determined via inductively coupled plasma optical emission spectrometry. This study established the accumulation of AgNPs, specifically the AgNPs–OCT–Alg, in lung tissues, and substantiated the active, specific, and selective targeting of SSTRs at pulmonary sites. The antitumor efficacy of the formulations was in vitro tested and confirmed in the MCF-7 cell lines. Owing to the delivery suitability and cytotoxic effects of the AgNPs–OCT–Alg formulation, it is a potential drug delivery formulation for lung cancer therapy in the future.

Keywords: silver nanoparticle, somatostatin receptor, active targeting, octreotide, in vivo lung cancer delivery, molecular modeling, receptor binding

1 Introduction

It is difficult to diagnose lung cancer because it has no early observable symptoms. Thus, the tumor is typically advanced at the time of primary diagnosis and continues to progress [1,2]. Currently, although radiotherapeutic and chemotherapeutic interventions are available as part of lung cancer treatment modalities, their harmful effects on normal and other noncancerous cells have limited their role, and research has been conducted to determine alternative treatment approaches [3]. These limitations and applications to treatment, through radio-oncologic modalities, have also led to dose perturbations with uncertain modulations during drug and radiological dose delivery.
However, the side effects of these treatment modalities have not yet rendered them ineffective; therefore, even with putative damage to normal or non-cancerous cells, studies on improving the efficacy and drug- and radiation-dose controls are under development. Although the effectiveness of a drug increases as its dose increases, the increase in the dose is limited by the inherent toxicity of the drug and the radio-intervention, leading to undesired physiological and morphological changes and the destruction of normal cells [4]. The inhalation route of drug delivery is considered an effective strategy for the treatment of lung diseases [5]. This route of drug administration shows favorable advantages in targeted organs such as the lungs because the drug can reach the sites directly through inhalation, provided that the carrier-based formulation and the drug to be delivered are feasible for inhalation. Both show receptivity to diseased tissue parts. In this case, the direct delivery of the drug to the lungs has the advantage of lowering the required dose of the drug, which would otherwise have been higher and required more frequent administrations. It also specifically targets cancer sites and reduces the toxic effects on normal/non-cancerous lung cells, which is feasible [6].

Breast cancer is the commonest cancer among women and the leading cause of cancer-related deaths worldwide. Breast cancer is initiated when the cells of the breast undergo uncontrolled division. Symptoms of breast disease include breast masses, changes in the volume of the breast, breast discomfort, and nipple discharge. A hard, painless mass in one of the nipples is the most common symptom of breast disease in males [7]. Currently, available treatment modalities for breast cancer include radiotherapy and chemotherapy; however, radiotherapy damages normal cells in addition to cancer cells. Chemotherapy can also be administered after surgery and radiotherapy [3]. However, cancer chemotherapy is limited due to its toxicity to normal cells. Moreover, increasing the dose can increase the effectiveness of chemotherapy [4]. However, many new drugs result in severe forms of toxicity, mainly in the form of myelo-suppression [8]. Delivering cytotoxic agents directly to tumor cells is more beneficial in terms of dose increase, reduction in the occurrence of peripheral toxicity, and improvement of therapeutic efficiency. Active targeting is considered an effective strategy for curing breast cancer. Further, there is a need to reduce the systemic side effects of parenteral and oral chemotherapy by using site-specific formulas.

Silver nanoparticles (AgNPs) are considered one of the most promising nanostructures. Their favorable physicochemical characteristics and bioactive properties have been found to be appropriate for several biomedical applications in vitro and in vivo. Intense interest in AgNPs and their derivatives has improved the treatment choices for currently available oncological interventions [9]. AgNPs have many applications in the medical, biomedical, and biotechnical fields, including cancer cell targeting therapy, and several other biomedical treatment arenas [10]. Although all studied AgNPs were previously formulated only for inhalation as a passive targeting strategy or to determine their cytotoxicity [11–13], AgNPs can be applied in active targeting via receptors. Moreover, ligand-coated AgNPs can target specific receptors in the human body, such as somatostatin (SST) receptors (SSTRs), which are among the most abundant receptors expressed by lung tumor cells [14]. High- and low-affinity SSTRs have been identified in numerous cancer cells, such as those of the lung, ovaries, pancreas, colorectum, and prostate [15,16]. Furthermore, SST analogs, such as octreotide (OCT) and vapreotide (VAP), have also been shown to have high and specific affinities to SSTR subtypes 2 and 5 [17,18]. OCT binds directly to receptors on tumor cells, as a majority of human tumor cells, either benign or malignant, usually express SSTRs [19]. Nilsson et al. have reported that numerous normal and infected human cells overexpress SSTRs. This makes SSTRs (particularly, SSTR3) feasible and promising target ligands using OCT or VAP-conjugated nanoparticulate delivery systems to reduce side effects and increase the accumulation of drugs in tumor cells [20]. Dasgupta et al. confirmed the delivery of OCT to cancer cells that express SSTRs, while Norenb erg et al. reported that SST analogs widely used to target cancer cells express SSTRs for therapy with 90Y or 177Lu radionuclides [21].

Currently, no studies have reported the use of an AgNPs-conjugated SST analog OCT to target lung cancer via inhalation. This study aimed to develop a simple, non-invasive, aerosol-based, site-specific method to deliver a payload (drug) to the lung tissues for delivery by nebulization for in vivo inhalation and subsequently targeted delivery.

2 Methods

2.1 Materials

Silver nitrate (AgNO3), tri-sodium citrate (TSC), sodium alginate (Alg), sodium chloride, potassium chloride, potassium dihydrogen phosphate, sodium hydroxide, and disodium hydrogen phosphate were purchased from El Nasr Pharmaceuticals Chemicals Company (Abu Zaabal, Cairo,
OCT acetate, Gibco Roswell Park Memorial Institute (RPMI) 1640 medium with 10% fetal bovine serum (FBS), and antibodies were purchased from Santa Cruz Biotechnology Inc. (Heidelberg, Germany). Kits for cytokine-specific enzyme-linked immunosorbent assays (ELISAs) were purchased from R&D Systems, Inc. (St. Paul, MN, USA). The Michigan Cancer Foundation-7 (MCF-7) cell line was obtained from Vacsera (Dokki, Giza, Egypt). All chemicals were of analytical grade.

### 2.2 Synthesis of silver citrate nanoparticles by the reduction technique

Citrate-reduced and capped AgNPs (AgNPs–CTT) were prepared using the citrate reduction method following a previously reported protocol [22] with some modifications using 100 mL of aqueous AgNO₃ (molecular weight, 169.87; 0.05 mol; 8.4935 g) solution. Briefly, 28 mg of AgNO₃ was diluted in 50 mL of Millipore-purified water (stock solution I). The stock solution I (6.5 mL) was diluted to 100 mL with Millipore water. The solution was stirred vigorously at 6,000 rpm to 100°C. About 2 mL of 2% TSC solution was slowly and carefully added to the boiling AgNO₃ solution in an Erlenmeyer flask and allowed to continue boiling for 15 min. The colorless solution changed to yellow, bright red, red, and brown, indicating completion of the reaction after 15 min. The AgNPs–CTTs were stored at room temperature (27°C) in amber-colored bottles. The particles were purified by centrifugation at 1,200 rpm for 6 min to remove large particulate matter.

### 2.3 Preparation of AgNPs–CTT capped with OCT and wrapped with alginate

The OCT was selectively conjugated to AgNPs–CTT through its N-terminus [23]. The AgNPs–CTT formed at a concentration of 0.1 mM was reacted with a two-fold excess of OCT. Then, 100 µL of 0.2 mM OCT was added to the reaction mixture, followed by the addition of 0.15 mM N-(3-dimethylaminopropyl)-N′-ethyl carbodiimide hydrochloride (EDC) and 0.2 mM N-hydroxsuccinimide, and stirred for an additional 3 h at room temperature. The final product, AgNPs–OCT, was purified by centrifugation at 1,200 rpm for 5 min to remove large particulate matter [24]. The AgNPs–OCT formulation was optimized for lung delivery by stirring AgNPs through dispersion in a sodium alginate aqueous solution (%), which generated a coat of alginate around the AgNPs–OCT. The formulated AgNPs–OCT–Alg was prepared to enhance AgNPs–OCT delivery into the lungs through inhalation.

We calculated the amount of Ag⁺ transformed to Ag⁰, which represents the percent yield of the produced AgNPs, using the previously described approach with some minor modifications [25,26]. Inductively coupled plasma optical emission spectrometry (JY-70 PLUS, Jobin Yvon Instruments SA Inc., Edison, NJ, USA) was used to calculate the amount of Ag⁰ in all AgNP formulated. Each formula was centrifuged at 1,200 rpm for 5 min in order to separate the large particulate and remove the impurities. The supernatant was evaluated by ICP-OES. The quantity of silver Ag⁰ was measured in the supernatant and then divided by the initial concentration of AgNO₃. Each solution was scanned times. The % yield of the produced AgNPs was calculated by dividing the obtained concentration by the initial concentration for AgNO₃ using the following equation:

\[
\% \text{ AgNPs yield} = \left( \frac{\text{Concentration of each sample by ICP OES}}{\text{Initial concentration of AgNO}_3} \right) \times 100. \tag{1}
\]

### 2.4 Size and zeta potential measurements

A Malvern Zetasizer Nano 6.01 analyzer (Malvern Instruments GmbH, Herrenberg, Germany) was used to measure the particle size, polydispersity index, and zeta (ζ) potential. Sampling times were automatically set on the instrument. Three measurements were permitted for every 10 subruns. All calculations were performed according to previously described protocols [14,27].

### 2.5 UV-Vis spectroscopy

The absorption spectra of AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg formulations were recorded. The absorption spectrum for each formulation was scanned in the range of 300–700 nm in a 2/cm plastic cuvette using the Uvikon-941 spectrophotometer (Kontron Device GmbH, Herrenberg, Germany). Surface plasmon resonance (SPR) peaks were obtained using Microsoft Excel for Macintosh 2019 (Microsoft Corporation, Redmond, WA, USA) [14].
2.6 Cell culture

2.6.1 Toxicity study

MCF-7 cells were grown in the Gibco RPMI 1640 medium containing 10% FBS and 1% streptomycin and penicillin at 37°C in 5% carbon dioxide, as described previously [28,29]. Toxicity studies were performed on the optimized formulations, and the results were compared with those obtained among the control cells. The cells were treated with various concentrations (0.1–10 µM) of AgNPs–CTT [30], AgNPs–OCT, and AgNPs–OCT–Alg for 24 h. The concentrations of all formulas were calculated depending on the molecular weight of silver. Cytotoxicity was examined using a kit (Promega, Madison, WI, USA), as previously described [31].

2.6.2 TNF-α ELISA

After pretreatment with different doses of AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg in MCF-7 cells for 2 h, MCF-7 cells were examined. The TNF-α content in the culture medium was quantified using a TNF-α-specific ELISA kit according to the manufacturer’s instructions (R&D Systems Inc., Minneapolis, MN, USA) [30].

2.6.3 Cell uptake and displacement studies

The uptake and transport of AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg in MCF-7 cells were investigated to determine the internalization of SSTR-based AgNPs–OCT into the cells. Furthermore, the AgNPs coupled with OCT and alginate ensured targeting of the nanoformulation, while the alginate wrapping ensured its feasibility for nebulization. A competitive replacement experiment was performed by incubating the AgNPs–OCT for 1 h at 37°C [32]. In brief, MCF-7 cells (1 × 10^5/mL) were placed in 35 mm culture dishes containing Dulbecco’s modified Eagle’s medium and incubated at 37°C overnight. The cells were incubated with 10 µM each of AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg. MCF-7 cells were used as a model for SSTR expression [33]. Moreover, the groups that received AgNPs–OCT and AgNPs–OCT–Alg were also incubated with 100 µM free OCT 30 min prior to incubation with the nanoparticles, which led to receptor saturation with OCT via competitive replacement. This step aimed to confirm that AgNPs–OCT and AgNPs–OCT–Alg could be displaced from SSTRs by free OCT. The cells were then washed twice with Dulbecco’s phosphate-buffered saline, centrifuged at 300 g (180.5°C) for 10 min, and placed in 5 mL glass vials. ICP-OES was used to assess the concentration of silver ions (Ag⁺) [34,35]. Briefly, each AgNP sample (0.5 mL) was mixed with 0.2 mL of freshly prepared aqua regia and diluted to 5 mL with Millipore water. ICP-OES was performed using a JY-70 PLUS (Jobin Yvon Instruments SA Inc.) to determine the quantity of Ag⁺ in the digested cells. Ag⁺ concentrations of 1, 10, and 100 ppm were used as standards. The concentration of AgNPs in solution was calculated by dividing the number of silver atoms per particle by the estimated concentration of Ag⁺ [14,36]. The ICP-OES principle is used to determine the quantity of metals in a sample by measuring the amount of light emitted at different wavelengths [37]. The amount of Ag⁺ was calculated using a Microsoft Excel sheet, as reported previously [35].

2.7 Experimental animals

The animal experiments were conducted at the Pharmacology Research Laboratory, College of Pharmacy, Qassim University (Buraydah, Saudi Arabia). The study protocol was approved by the Institutional Animal Ethics Committee Deanship of Scientific Research, Qassim University, Saudi Arabia (approval number: 2019-2-2.1-5551). A histamine chamber (Plexiglas) was prepared (with the following dimensions: length 32 cm, width 15 cm, and height 28 cm) to expose four rats to the prepared AgNP formulations. The nebulizer inlet carried AgNPs to the rats, and the outlet tube maintained the optimum pressure inside the chamber for inhalation. Millipore water was used to prepare all nebulizing solutions. This experiment was carried out on 12 Sprague–Dawley rats of both sexes obtained from the College of Pharmacy Animal House. The rats were 7 weeks old, with body weights of 200–250 g, and were provided a rodent chow diet and water ad libitum. All rats were acclimatized to the laboratory environment in clean cages for 5 days and maintained daily with a 10–14 h photoperiod. The rats were nebulized using a System LS 290 aerosol nebulizer (System Assistance Medical SAS, Villeneuve-sur-Lot, France). This system used room air with appropriate pressures and temperatures to vaporize solutions into a micro-fine aerosol mist that flowed into the tubes of the Plexiglas chamber for the animals to inhale. The rats were distributed equally into groups (the control, AgNPs–CTT, AgNPs–OCT–Alg groups). Rats in the control group were nebulized with Millipore water, those in the AgNPs–CTT group with AgNPs–CTT, and those in the AgNPs–OCT–Alg group with AgNPs–OCT–Alg. The concentrations of the nanoformulations were 10 µM each. The total dose of each formulation was 1.27 mg/kg [38], and 50 mL of each formulation was used for nebulization.
at a rate of 5 mL/h. In each experiment, the nebulizer dosing cup was filled with 10 mL of liquid and nebulized until it dried out; the nebulization was continued for 3 days (within 10 h/day). On the same day, lung tissues were excised and processed [39,40].

2.7.1 Analysis of the biodistribution of the AgNPs–OCT–Alg

The quantity of Ag⁺ was analyzed using ICP-OES to determine the amount of inhaled AgNPs–OCT. Lung tissues were homogenized in 86% nitric acid and hydrochloric acid. The homogenized tissue was diluted with phosphate-buffered saline (pH 7.4). The AgNPs–OCT concentration was determined by quantifying the amount of Ag⁺ using the ICP-OES technique according to a protocol described in previous studies [41,42].

2.7.2 Ultrastructural examination by transmission electron microscopy (TEM)

Lung specimens were excised and processed (fixed in 2.5% glutaraldehyde for 2 h, post-fixed in 1% osmium tetroxide for 1 h, dehydrated in ethanol, cleared in propylene oxide, and embedded in epoxy resin). After thermal polymerization, ultrathin sections were cut with an ultramicrotome and stained with uranyl acetate and lead citrate [43]. The grids of the ultrathin sections were examined and photographed using a transmission electron microscope (JEOL 1010; JEOL Ltd, Tokyo, Japan) at the Electron Microscopy Unit of the College of Applied Medical Sciences (Qassim University) to identify the ultrastructural details of the alveolar cells [44]. The nanoparticles were electron-dense with characteristic round shapes and equal in size.

2.7.3 Statistical analysis

The data are presented as the mean of at least three experiments with their standard deviations. Comparisons of single or multiple samples were performed using t-tests. The analyses were performed using SPSS Statistics version 27 (IBM Corporation, Armonk, NY, USA). Analysis items with p < 0.05 were considered statistically significant.

3 Results

The AgNPs–CTT formulation was prepared and conjugated to OCT to enable it to specifically target SSTR₂. The prepared AgNPs–CTT–OCT or AgNPs–OCT was coated with alginate (via a 1% aqueous sodium alginate solution) to yield AgNPs–OCT–Alg for easy delivery to the lungs by inhalation in rat models in a histamine chamber (Figure 1a–c) (Plexiglas). The nebulizer inlet transported the AgNPs–OCT–Alg formulation to the rats (Figure 1b and c). The prepared AgNPs–CTTs were uniformly shaped and brown in color. The brown color turned darker after conjugating the AgNPs–CTT with OCT. The AgNPs–CTT was coupled with OCT at its N-terminus, forming an amide bond between the OCT and the citrate carboxyl group of the AgNPs–CTT. The reaction can be progressed at a slightly acidic or neutral pH to avoid reaction with the lysine side-chain amino group (NH₃) of OCT, which (lysine-NH₂) is essential for activity (Figure 1d) [17,45].

Table 1 shows the initial Ag⁺ and Ag⁰ concentrations in AgNPs following NP formation, as measured by ICP-OES. The yield% was calculated for Ag⁰ synthesized in the AgNPs–CTT, AgNPs–OCT, AgNPs–OCT–Alg to be 2.82 ± 0.016 mg/100 mL (≈ 77.9%), 2.43 ± 0.21 mg/100 mL (≈ 67.12%), and 2.166 ± 0.19 mg/100 mL (≈ 59.83%), respectively.

3.1 Size, ζ potential, and surface morphology

A dynamic light scattering zetasizer nano was used to record the sizes of the AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg, which were determined to be 22.77 ± 1.1, 78.77 ± 2.3, and 155.99 ± 5.2 nm, respectively. The ζ potentials of the three formulations had surface charge values of −40.17 ± 2.0, −14.13 ± 1.3, and +16.17 ± 1.3 mV, respectively (Figure 2).

3.2 Ultraviolet-visible spectroscopy

Ultraviolet-visible spectroscopy (UV-VIS) was used to record the wavelengths of maximum absorption (λₘₐₓ) of the AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg, which were determined as 430.7 ± 2.99, 447.8 ± 2.93, and 450.8 ± 3.5 nm, respectively. The UV-VIS spectrum of the AgNPs–CTT showed redshifts of 17 and 3 nm in the SPR peaks for both AgNPs–OCT and AgNPs–OCT–Alg, respectively (Figure 3) [35].

3.3 In vitro treatment of MCF-7 cells with AgNP formulations

The cytotoxic effects of the AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg on MCF-7 cells were evaluated using
Figure 1: (a) Schematic diagram of AgNPs–OCT–Alg preparation and delivery to lung cells. (b and c) Nebulization of AgNPs–OCT–Alg among rats. (d) Schematic diagram of AgNPs–OCT synthesis; the AgNPs–OCT formed were coated with alginate for delivery to the lung cells and the proposed manner in which OCT fits into the SSTRs by interactions between lysine and phenylalanine.
different concentrations (0.1–10 µM) of the nanoparticle formulations. The extent of cancer cell proliferation significantly decreased after treatment with different AgNP formulations compared to the proliferation of control cells (Figure 4). In the model cytotoxicity evaluation experiment involving the MCF-7 cell lines, the results showed that treatment of the breast cancer cell lines with AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg significantly reduced cell proliferation in a concentration-dependent manner (Figure 4). The degree of proliferation

<table>
<thead>
<tr>
<th>Sample</th>
<th>Concentration (mg/100 mL)</th>
<th>Yield % (conversion%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ag⁺ (initial concentration)</td>
<td>3.62 ± 0.1</td>
<td>—</td>
</tr>
<tr>
<td>AgNPs–CTT</td>
<td>2.82 ± 0.016</td>
<td>77.9</td>
</tr>
<tr>
<td>AgNPs–OCT</td>
<td>2.43 ± 0.21</td>
<td>67.12</td>
</tr>
<tr>
<td>AgNPs–OCT–Alg</td>
<td>2.166 ± 0.19</td>
<td>59.83</td>
</tr>
</tbody>
</table>

Table 1: The yield percentage of Ag⁺ in AgNPs after conversion of Ag⁺ to Ag⁰

Figure 2: Size distributions (nm), ζ potentials (mV), and polydispersity indices of the AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg.

Figure 3: UV-Vis spectra of AgNPs–CTT, AgNPs–OCT and AgNPs–OCT–Alg. The AgNPs–OCT and AgNPs–OCT–Alg showed redshifts in their SPR peaks.
was significantly reduced when MCF-7 cells were treated with each of the AgNP formulations for 24 h. The AgNPs–OCT–Alg formulation was more efficient in reducing cell proliferation than the AgNPs–OCT, and AgNPs–CTT formulations after 24 h of incubation (Figure 4). Treatment of the MCF-7 cell lines with 5–10 μM of each of the AgNP formulations revealed that treatment with AgNPs–OCT and AgNPs–OCT–Alg inhibited the proliferation of all cells compared to the cells treated with only the AgNPs–CTT [46].

In addition, the protein levels of TNF-α were determined via ELISA. Treatment with 0.1–10 μM of AgNP preparations significantly decreased the production of TNF-α in the MCF-7 cell lines compared with control cells (Figure 5). The cell cultures treated with 5–10 μM each of the AgNP formulations showed significant suppression ($p < 0.05$). The level of TNF-α production was higher in the 0.1 μM group than in the control group. The levels of TNF-α decreased significantly in the cells treated with 5 and 10 μM each of
the AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg (p < 0.05) formulations compared with the cells in the control group. These results confirmed that the nanoparticles could be used as cancer formulations at concentrations >5 µM.

Moreover, ICP-OES showed higher levels of internalization of AgNPs–OCT also into the MCF-7 cells. The cellular uptakes of 10 µM formulations of AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg were determined via ICP-OES after 60 min of incubation. The control MCF-7 cell lines showed negligible quantities of internalized AgNPs, and AgNPs–CTT showed nonsignificant levels of random internalization (p > 0.05). AgNPs–OCT showed more significant levels of internalization at 387 ± 23.6 × 10³ per cell (p < 0.001). The amount of accumulated AgNPs decreased to 243 ± 12.6 × 10³ in the cells treated with AgNPs–OCT–Alg (p < 0.05). Furthermore, the amounts of internalized AgNPs decreased to 156.88 ± 21.4 × 10³ and 126 ± 11.7 × 10³ in the AgNPs–OCT and AgNPs–OCT–Alg treated groups, respectively (p < 0.001), when the receptors were occupied by 100 µM of free OCT (Figure 6).

3.4 In vivo inhalation of AgNP formulations

The images of inhaled AgNPs in different groups are presented as photomicrographs from lung sections stained with toluidine blue (Figure 7a–c). Figure 7b shows an image of inhaled AgNPs–CTT (arrow) and Figure 7c shows an image of inhaled AgNPs–OCT–Alg (arrow). Also, Figure 7d–f presents the TEM photomicrographs of the ultrathin lung sections. Figure 7e presents an image of inhaled AgNPs–CTT (arrow) and Figure 7f shows an image of inhaled AgNPs–OCT–Alg (arrow). Figure 7g shows the proportions of inhaled AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg as determined via ICP-OES. Figure 7a shows the absence of inhaled AgNPs in the lungs; Figure 7b shows an image with a low amount of inhaled AgNPs–CTT (red arrow), while Figure 7c shows an image with a significant amount of inhaled AgNPs–OCT–Alg (red arrow) (p < 0.05). The results were confirmed by TEM photomicrographs of ultrathin lung sections (Figure 7d–f). Figure 7d shows the absence of inhaled AgNPs, Figure 7e displays a low amount of inhaled AgNPs–CTT (red arrow), while Figure 7f shows a higher amount of inhaled AgNPs–OCT–Alg (red arrow), which confirms our hypothesis. In addition, the nanoparticles shown in Figure 7e and f were smaller than the artifacts appearing as irregular black spots (blue arrows) in Figure 7d. These results were also confirmed by ICP-OES, which determined the amount of inhaled AgNPs in lung cells. Figure 7g demonstrates the proportion of inhaled AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg as determined by ICP-OES. The proportion of inhaled AgNPs–OCT–Alg was 0.416 ± 5.7 mg/kg tissue, which confirmed active targeting of the lung tissue, while the proportion

![Figure 6](image_url)
AgNPs–OCT–Alg formulations were all inhaled after nebulization, and the lung tissues were histologically examined for the presence of AgNPs. Nonspecific and random accumulation of AgNPs–CTT in the lung occurred due to rapid uptake by alveolar cells [47].

4 Discussion

In the present study, we modified a previously described chemical reduction technique to synthesize citrate-reduced and capped nanoparticles (AgNPs–CTT) [48]. Due to the reduction of Ag⁺, neutral silver atoms were formed by the nucleation of Ag⁺ [49] according to the redox equation, based on the reduction potential of the bulk starting material, as shown below:

\[ \text{Ag}^+ + e^- \rightarrow \text{Ag}(s) \]

Seeding and nucleus production were more likely to occur when Ag⁺ was used at low concentrations (sub-millimolar range), as used in this study. Constant stirring of the reaction solution was necessary to prevent collision and the formation of silver nuclei in the solution. Temperature and pH are also important factors in the preparation; however, they also affect synthesis and product yields, resulting in changes in the nanoparticle size, charge, and polydispersity [36]. The reduced AgNPs–CTT provided strongly reactive, thermodynamically stable, fully reduced, appropriately citrate-capped AgNPs with high antioxidant potential, functional, specific, and ready-to-conjugate nanopreparations. The reduction process yielded characteristically superior quality by designing nanoparticles for further use in the preparation. Bulk materials are very sluggish in reacting and producing conjugates, have been improved to highly chemically reactive, energy-absorbing nanopreparations, and have provided the functional material for further use in preparing desired nanoparticles with site-specific targeting. The yield% was calculated for Ag⁰ synthesized in the AgNPs–CTT, AgNPs–OCT, AgNPs–OCT–Alg using ICP-OES to be 2.82 ± 0.016 mg/100 mL (≈ 77.9%), 2.43 ± 0.21 mg/100 mL (≈ 67.12%), and 2.16 ± 0.19 mg/100 mL (≈ 59.83%), respectively. Although the yield percentage does not reach 100% with the current synthesis method, while an acceptable yield percentage was attained for all kinds of AgNPs. Inconsistencies between reported and theoretical Ag inputs can also be ascribed to Ag⁺ or AgNPs adsorption on centrifuge tubes or pipettes, volume loss, and transportation during sample transfer for dilution and quantification [25].
Nanoparticle delivery to the lungs suffers from two significant drawbacks, one of which occurs prior to administration and the other after administration. First, nanoparticle aggregation occurs more frequently in an aqueous medium after long-term storage. Second, the smaller-sized nanoparticles are quickly exhaled unless they are delivered and deposited deep inside the lungs [50,51]. The AgNPs–OCT–Alg was designed as a formulation carrier to overcome the problems associated with pulmonary delivery and the aggregation of nanoparticles to achieve better delivery and deposition in the lungs. The formulated nanoparticles incorporated an alginate layer, which acted as an outer coat and worked as a better carrier than the nanoparticles themselves and the alginate coating and the nanoparticles themselves, as well as noncovalent coating between the AgNPs–OCT and AgNPs–Alg, which are easily exhaled upon administration among the animals. The alginate coating was only used to facilitate the inhalation route and nearly diminished the exhalation of the AgNPs–OCT formulation. The alginate wrapping was replaced in aqueous media, both in the in vivo (inhalation among the rats) and in vitro (MCF7 cancer cells) models, and resulted in internalization at higher levels than the non-OCT citrate-capped formulation (AgNPs–CTT) (Figure 8). The SPR redshifts were observed due to the coupling of AgNPs–CTT with the OCT and alginate wrapping of the AgNPs–OCT. The optical properties of all AgNPs depended on the diameters of the preparations, and the peaks shifted toward longer wavelengths (redshift); therefore, the large-sized particles had larger optical cross-sections, and their albedo (the reflected fraction of SPR) increased similar to the redshifts with the increased size [35]. In the case of the AgNPs–OCT–Alg, the redshift was 3 nm, which signified the least changes in the SPR shifts (nearly no structural change, except noncovalent coating between the AgNPs–OCT and the AgNPs–OCT–Alg). The final products were citrate-capped nanoparticles (AgNPs–CTT), OCT-chemically-conjugated-to-citrate nanoparticles (AgNPs–OCT), and the alginate-wrapped AgNPs–OCT–Alg. The absorption for the covalently linked alginate was much higher. However, in this case, the AgNPs–OCT was treated in an alginate solution, dried to produce the alginate layer, which was easily exhaled upon administration among the animals. The alginate wrapping was replaced in aqueous media, both in the in vivo (inhalation among the rats) and in vitro (MCF7 cancer cells) models, and resulted in internalization at higher levels than the non-OCT citrate-capped formulation (AgNPs–CTT) (Figure 8). The SPR redshifts were observed due to the coupling of AgNPs–CTT with the OCT and alginate wrapping of the AgNPs–OCT. The optical properties of all AgNPs depended on the diameters of the preparations, and the peaks shifted toward longer wavelengths (redshift); therefore, the large-sized particles had larger optical cross-sections, and their albedo (the reflected fraction of SPR) increased similar to the redshifts with the increased size [35]. In the case of the AgNPs–OCT–Alg, the redshift was 3 nm, which signified the least changes in the SPR shifts (nearly no structural change, except noncovalent coating between the AgNPs–OCT and the AgNPs–OCT–Alg). The final products were citrate-capped nanoparticles (AgNPs–CTT), OCT-chemically-conjugated-to-citrate nanoparticles (AgNPs–OCT), and the alginate-wrapped AgNPs–OCT–Alg. The absorption for the covalently linked alginate was much higher. However, in this case, the AgNPs–OCT was treated in an alginate solution, dried to produce the alginate-wrapped nanoparticles (a thin coating/film around the AgNPs–OCT) as AgNPs–OCT–Alg, and the chemically conjugated (absorption being higher) and nonconjugated (with no chemical bond) formulations showed comparatively low levels of absorption. As the nanoparticles were in aqueous media, alginate displacement yielded very little change in the

**Figure 8:** Schematic representation of the different types of AgNPs applied in *in vitro* and *in vivo* experiments.
absorption, as in this spectrum recording event, alginate was displaced with enough water in the medium. All other UV recordings were also performed using water as the medium. All three preparations were also of different colors. The smaller difference of 3 nm was a result of the effect of alginate. UV spectra were recorded for the AgNPs–OCT–Alg product, a characterized material with size, charge, and shape specifications. The small shift (redshift) of <3 nm was in agreement with the layer-by-layer coating of AgNPs reported by Detsi et al. [53].

Therefore, due to nanoparticles, AgNPs–OCT–Alg increased in size and was found to be more effectively delivered into the lungs through inhalation. As reported previously, for a carrier of suitable (nano-metric) size, entrapment in the lungs is better for achieving pulmonary deposition (Figure 9) [54]. The size of the AgNPs–OCT–Alg was intended to be increased, and the size of AgNPs increased from 22.77 ± 1.1 nm for the AgNPs–CTT to 78.77 ± 2.3 nm for the AgNPs–OCT. Finally, the coated nanoformulation, AgNPs–OCT–Alg, resulted in an average size of 155.99 ± 5.2 nm, which could be effectively delivered to the lungs. The alginate coating has been proposed to be dissolved after formulation delivery since there is no chemical bonding between the alginate and the surface-located OCT of the AgNPs–OCT, except for the physical wrappings of the alginate coating around the AgNPs–OCT. However, a chemical bond forms between the AgNPs–CTT and OCT to produce AgNPs–OCT from the AgNPs–CTT based carboxylate and the NH₂ group of the OCT forming an amide linkage, which is stable enough for delivery and deposition [45,55]. The Ag⁺ internalizations of the formulations, AgNPs–CTT and AgNPs–OCT–Alg, were compared in terms of site-specific delivery and Ag⁺ internalizations by the alginate-coated nanoparticles and noncoated, non-OCT-capped AgNPs–CTT. There was no need to compare the AgNPs–CTT and AgNPs–OCT, as the OCT coating itself was target-specific (targeting SSTRs) and was expected to yield higher levels of internalization of Ag⁺

Figure 9: Structural representations of the AgNP nanocarriers; (a–c) 2D renderings, (a) AgNPs–CTT (citrate bound to the nanoparticle, nanoparticle depicted in black). (b) AgNPs–OCT (OCT covalently bound to citrate, CTT). (c) AgNPs–OCT–Alg (no chemical bonding; instead, physical wrapping of the alginate around the AgNPs–OCT. Alg: alginate, coatings shown in blue). 3D renderings of the AgNPs. (d–f) 3D renderings, (d) AgNPs–CTT. (e) AgNPs–CTT–OCT. (f) AgNPs–OCT–Alg. Alginate coatings are not chemically bound. The core of the formulated AgNPs–OCT is shown in red, while the surrounding alginate wrapping is shown in blue.
The AgNPs–OCT–Alg could be considered as an innovative therapy for lung diseases, as it imparts SSTR analogs and several drugs with the potential for delivery due to the capacity for attachment to the AgNPs–CTT and alginate coating of alginate onto AgNPs, respectively. The AgNPs (their stages of preparation and developmental processes) formulation to cell lines under in vitro conditions, and successfully and effectively delivered the AgNPs–OCT–Alg formulation to effectively deposit into the lung tissues within the optimal concentration limits, between 5 and 10 µM, for in vivo anticancer activity in rat models. The lowest concentration that induced anticancer activity in vivo has also been investigated [46,60]. The half-maximal inhibitory concentration (IC50) values were observed at 5–10 and 10 µM under in vitro and in vivo conditions, respectively. Moreover, the IC50 for AgNPs–CTT was also found to be less than 20 µM [30]. TNF-α promotes the proliferation of cancer cells [61], it was also observed that AgNPs–OCT significantly inhibited TNF-α production. Moreover, the cells were affected by the minimum concentration of AgNPs–OCT. A higher concentration of all AgNP preparations (10 µM) was used to assess their antiproliferative activity, as suggested by Xu et al. [62]. The researchers confirmed that the toxicity of AgNPs–CTT increased at concentrations >10 µg/mL, while AgNPs–CTT (5 µg/mL) tended to show comparatively reduced levels of cytotoxicity. Moreover, ICP-OES showed higher levels of internalization of AgNPs in the form of AgNPs–OCT in in vitro tests due to the direct interaction between OCT and SSTR2 expressed on the surfaces of the MCF-7 cells. In addition, the weak interactions between AgNPs–OCT–Alg and MCF-7 cells due to alginate coating on the surface of the nanoparticles initially formed a barrier between the OCT binding sites and SSTR2.

The selectivity for SSTR2 was confirmed based on the occupation of the receptors with 100 µM free OCT, and the binding of the nanoparticulate formulations to the receptor was observed to be significantly reduced under these conditions [59,63,64]. Moreover, these results showed lower levels of internalization of the nanoparticle formulations in the cells, as binding was reduced in the presence of free OCT, which preferentially and quickly blocked and occupied the SSTR2. The AgNP formulations were applied at concentrations described in a previous report published by Xu et al. [62]. The levels of TNF-α also showed a marked decline, as determined via ELISA. The AgNP formulations at concentrations of 0.1–10 µM significantly decreased TNF-α production in MCF-7 cells. Moreover, the 10 µM AgNP formulations showed the most significant levels of suppression of TNF-α, with no additional decrease in TNF-α production at higher nanoparticle concentrations [30]. This significant decline in TNF-α production indicated that AgNPs–OCT could be an effective material to target cancer cells, particularly, breast cancer cells, while decreasing TNF-α production. The results obtained also showed promise in targeting breast cancer since the AgNPs–OCT and AgNPs–OCT–Alg were better designed for active targeting with an increased cellular accumulation of nanoparticles. Active targeting of
breast cancer is a promising strategy for efficient drug delivery [65]. Many target sites can be reached in breast tissue for site-specific and active targeting [65]. SSTR2 and SSTR5 are expressed in neuroendocrine breast cancer and are considered potential targets for the treatment of breast cancer in the future [66,67]. Li et al. [68] reported a novel active targeting approach established on L-type amino acid transporter 1 (LAT1) expressed in a range of cancers. Poly (lactic-co-glycolic acid) nanoparticles have also been investigated among MCF-7 cells. Compared to unmodified nanoparticles, LAT1-targeting nanoparticles showed a considerable increase in cellular uptake and cytotoxicity.

The AgNPs–OCT–Alg formulation is stabilized primarily by the adsorption of alginate (Alg) onto the surface of AgNPs–OCT [69]. The produced nanomaterials (AgNPs–CTT and AgNPs–OCT) are comparatively smaller in size and lighter in weight, which implies that upon inhalation, these AgNPs preparations will be either partially or completely exhaled without deposition into the lungs. The alginate coating provides additional increases in layering, weightage, thickness, and size to yield a larger and effective nanoscale carrier (approximately, 155.99 nm) with good flow properties to target the required regions in the lungs and with minimum or no exhalation of the formulation. After accumulating in the lungs, the AgNPs–OCT–Alg attaches to the surface of the pulmonary alveoli, and the coated nanoparticle formulation is presented in the vicinity of the epithelial cells for delivery into the lung cells [70]. These sizes of deposition are in agreement with the report published by Brown [71], which stated that particle deposition in the respiratory tract occurs mainly by diffusion, sedimentation, and impaction. The total amount deposited in the respiratory tract follows a “U-shaped” curve, with a minimum deposition of nanoparticles with sizes ranging between 0.1 and 1.0 μm, with enhanced diffusive deposition. Han et al. [72], formulated gold nanoparticles (AuNPs) and studied the effect of size on the biodistribution of AuNPs among Sprague–Dawley rats via inhalation. The researchers used small AuNPs (with diameters of 13–105 nm and concentrations of 12.8–13.7 μg/m). The animals were exposed to AuNPs for 5 days (6 h/day). They found that the exposed rats exhibited no toxic responses to AuNPs. Regardless of size, both small and large AuNPs were deposited in the lungs of rats. The biodistribution of small AuNPs from the lungs to secondary target organs was significantly higher when compared to those of large AuNPs. Moreover, nanoparticles with diameters smaller than 100 nm were also probably deposited [73,74].

Upon reaching the lung tissues, the alginate layer (Alg) is expected to dissolve slowly in the presence of aqueous media based on the presence of water/moisture, swelling, and removal of the alginate coating, and AgNPs–OCT is released from the wrapped coating of alginate. The AgNPs–OCT escapes the initial mucociliary clearance of the lungs and crosses the epithelial layer to target lung cancer through OCT, and cancer cell surfaces present SSTR interactions for deposition deep in the lung tissues. Alginate, a highly hydrophilic ionic material (in the form of Alg), is expected to dissolve within a short time, as recorded in the UV-VIS absorption experiments as well as in the tests of the accumulation of AgNPs in nanoformulation, wherein the coated material accumulated to a lesser extent than did the free OCT.

Healthy rats were used for the in vivo study as reported to express the SSTR2 [75], which was our target to prove. The OCT ligand-coated AgNPs–OCT was capable of specifically targeting SSTRs located in the lung tissues. The ability of these AgNP formulations to target SSTRs expressed in lungs is also potentially beneficial in the future treatment of other cancers that express SSTRs. The results obtained in targeting the lungs in the present study also corroborated a previous finding that SST analogs coated with AuNPs target SSTR-expressing cancer cells [14]. Other studies also provided details on VAP-coated quantum dots (QDs) and showed that the formulated VAP-QDs can be delivered to BON-1 cells overexpressing SSTR2 [17]. SST has a longer peptide sequence, while OCT has a shorter sequence that is feasible in many ways for conjugation, targeting, and delivery. OCT also selectively binds to SSTR2 [76]. Biomechanical considerations for current targeting by OCT have been proposed, wherein a binding interface has been depicted that exhibits both hydrophobic and hydrophilic interactions between the peptide residues. Ala-Gly-*Cys(cyclo)-Lys-Asn-Phe-Trp-Lys-Thr-Phe-Ser-*Cys-(*cyclo)-OH is the amino acid sequence of SSTR2 that is involved in binding with the Phe-Lys residues of OCT. A preliminary binding model of the interaction between the SSTR and OCT is shown in Figure 1. The pharmacophore-containing residues Phe-Trp-Lys-Thr of the SSTR bind with the incoming ligand to exert its biological activity. A parallel and detailed mechanistic study of the binding interactions is available [77].

The AgNPs–CTT, AgNPs–OCT, and AgNPs–OCT–Alg formulations were inhaled via nebulization at a dose for each formulation 1.27 mg/kg. This dose was selected based on the results of in vitro cytotoxicity and previously reported results of in vivo experiments [12,38,78,79]. All formulations were used in the liquid form to be nebulized and not in dry form, as the dried form of AgNPs has the potential for aggregation and cannot be resuspended. Freeze-drying of AgNPs induces aggregation [80]. The animals successfully inhaled the formulated AgNPs
without experiencing drowsiness or coma. The AgNPs–OCT–Alg was more viscous than the other formulas but was successfully nebulized. To determine the amounts of all AgNPs, the lungs in each group of animals were digested and examined for the inhalation of AgNPs. Upon pulmonary delivery, the AgNPs–CTT accumulated in the lungs due to rapid lymphatic uptake through non-specific binding interactions, while the AgNPs–OCT and AgNPs–OCT–Alg formulations accumulated via specific and targeted delivery. The proportion of inhaled AgNPs–OCT–Alg was 0.416 ± 5.7 mg/kg tissu, which confirmed the successful active targeting of the lung tissue, while only 0.199 ± 3.21 mg/kg tissu was detected for AgNPs–CTT, indicating nonspecific uptake of Ag⁺. The lungs showed a higher accumulation of AgNPs–OCT and AgNPs–OCT–Alg due to the expression of SSTRs on the lung cells, which specifically bound to the AgNPs-conjugated OCT [47,81,82]. This form of targeting is active, specific, and selective for lung cancer cells, such as those of non-small cell lung cancer. The AgNPs–OCT–Alg was internalized via active targeting, while a parallel report described the internalization of AgNPs into A549 human alveolar cells via nonspecific targeting [83].

However, numerous new drugs have shown high levels of cytotoxicity, mainly due to myelosuppression. Indeed, the delivery of cytotoxic agents that can selectively target tumor cells is useful in terms of controlling concentration by way of frequency and quantity. A reduction in peripheral toxicity also improves the therapeutic efficacy of chemotherapeutic agents [8]. Previous studies have reported a number of approaches to targeted drug delivery through specific tags on delivery carriers or at the site, compatible with the double-ended molecular structure. Reubi demonstrated that certain SST analogs labeled with radionuclides (111In, 90Y, 177Lu, and 68Ga) and injected intravenously in mice resulted in reductions in tumor masses [84]. Laznickova et al. used materials labeled with radioactive compounds to target SST analogs for diagnostic imaging and the treatment of tumors expressing SSTRs [85]. Moreover, it has also been reported that nanoparticulate formulations provide controlled and sustained drug release in the lung tissue, which supplements reduce the frequency of dosing and complement patient compliance. These approaches of targeted delivery, particularly, delivery to the lungs, further provided impetus to our proposal for the alginate-masked and OCT-mediated site-specific and well-targeted delivery of the lungs mediated by SSTRs.

In conclusion, the functionalized AgNPs were successfully produced using the citrate reduction method, and the synthesized AgNPs–CTT was chemically conjugated with OCT to target SSTRs, particularly, the SSTR₂. We developed and evaluated a formulation (AgNPs–OCT–Alg) that was capable of delivery to and internalization in the healthy lung tissues through the nebulization route in the animal model. An array of better prospects exists for the AgNPs–OCT–Alg formulation to serve as a specific, active, site-directed, and on-site method of drug delivery by using the currently presented noninvasive drug delivery method, which is plausible and holds further promise. The prepared AgNPs–OCT–Alg successfully targeted healthy lung tissues in vivo in a rat model, and in in vitro condition in the model breast cancer cell line, MCF-7, through binding to SSTRs. We analytically confirmed that the AgNPs–CTT was coated with OCT and the alginate wrappings. The presence of OCT in the nanoformulations, AgNPs–OCT and AgNPs–OCT–Alg, assisted in on-site delivery and cell targeting. MCF-7 cells showed specific interactions with the prepared AgNPs–OCT–Alg formulation. Moreover, the AgNPs–OCT and AgNPs–OCT–Alg exhibited higher internalization in the lung tissues due to the facile inhalation strategy and subsequent internaliza tion. The in vivo experiments also exhibited high levels of biodistribution and high concentrations of AgNPs in the lung tissue, as confirmed by the presence of Ag⁺ in ICP-OES analyses of several lung tissue samples obtained from various animal experimental groups treated with different nanoformulations and their varying dose concentrations. Further studies are needed for future lung cancer targeting. Also, more studies of animal lung cancer models are required to reflect the accuracy of active targeting of SSTRs by the AgNPs–OCT–Alg in lung tissues and lung cancer.

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