Phototherapy and Sonotherapy of Melanoma Cancer Cells Using Nanoparticles of Selenium-Polyethylene Glycol-Curcumin as a Dual-Mode Sensitizer

Mohammadi S.1,2, Soratijahromi E.1,2, Dehdari Vais R.2, Sattarahmady N.3,4*

ABSTRACT

Background: As an alternative form of cancer therapy, photothermal therapy (PTT) and sonodynamic therapy (SDT) using nanomaterials are in development. Nanomaterials can act as energy absorber as well as anti-cancer agent.

Objective: In this study, the effects of laser and ultrasound irradiation with Se-PEG-Cur nanoparticles were investigated on melanoma cancer.

Material and Methods: In this experimental study, nanoparticles of selenium-polyethylene glycol-curcumin (Se-PEG-Cur) were synthesized, and their UV-vis absorption, particle size, zeta potential and photothermal conversion efficiency were determined. Se-PEG-Cur was then introduced as a novel 808-nm laser light absorbing agent as well as ultrasound (US) wave for treatment of C540 (B16/F10) cancer cells. Also, ROS generation in C540 (B16/F10) cancer cells was measured upon PTT and SDT using Se-PEG-Cur.

Results: Mean size, zeta potential and photothermal conversion efficiency of Se-PEG-Cur were obtained as ~300 nm, 42.7 mV and 16.7%, respectively. Cell viability upon irradiation of the laser light or US waves with 100 µg mL⁻¹ Se-PEG-Cur were decreased to 33.9 and 22.9%, respectively.

Conclusion: Intracellular ROS detection indicated that dual PTT and SDT in the presence of Se-PEG-Cur induced the highest ROS production. Se-PEG-Cur was therefore introduced as an absorbing agent of both laser light and US waves for cancer treatment.

Keywords

Hyperthermia; Skin Cancer; Diode Laser; Ultrasound; Nanomedicine; Reactive Oxygen Species; Cell Survival

Introduction

Losing normal regulation and extreme cell proliferation can cause cancers. The lymphatic and circulatory systems can produce cancer cells that transferred to other parts of the body and invade tissues in living systems. They can even lead into lesions or patients death [1]. There are some limitations such as high side effects and low therapy...
efficiency for common cancer therapies of chemotherapy, radiotherapy and surgery [2]. To treat cancers, two alternative phototherapy methods including photothermal therapy (PTT) and photodynamic therapy (PDT) have been developed [3-5]. Phototherapy has some advantages including spatial and temporal control, minimal invasion and low toxicity in comparison with the conventional treatments [6]. Converting near infrared (NIR) light into heat can be done during PTT as a new technology to treat cancers [4, 5, 7]. Through this route, cancer cells are killed by rising temperature, while, it has the least side effects on normal cells [8]. One of the treatment modalities in PDT is combination of a proper and suitable light wavelength with a photosensitizing agent for selective killing the cancer cells. Formation of reactive oxygen species (ROS) such as singlet oxygen (\(O_2^\cdot\)) or free radicals rendering necrotic and apoptotic cell death can be made by the interaction of photons with photosensitizers in the presence of oxygen molecules. ROS are correlated with deleterious effects of cancer cells including DNA fragmentation, membrane perturbation and cytoskeletal damages [9-12]. Following PDT, sonodynamic therapy (SDT) has been developed as a new noninvasive approach [13]. Beside the important advantages of SDT including less invasiveness, deeper penetration of ultrasound (US, in comparison with PDT) enables noninvasive treatment of deep-seated tumors [14]. In SDT, US activation of an agent that is called sonosensitizer usually leads to generation of ROS.

Nanotechnology has caused criteria of diagnostics and treatment to be elevated and is also considered as the best compared to the most encouraging exploration introduction for oncotherapy [15-19]. Owing to the novel bioactivities in nanostructures, inorganic nanoparticles (NPs) of metals and metal oxides have a noticeable place for biomedical applications [20-22]. Up to now, metal nanomaterials have been served as proper activator agents of laser light, US and X-ray that increase quality of cancer treatment strategies [23, 24]. One of the important nanostructures for cancer cell treatment is selenium nanoparticles with low toxicity against healthy cells [15, 25, 26]. It exhibits anticancer effect by introducing ROS-mediated apoptosis in the cancerous cells [15, 25, 26].

Curcumin that is derived from the rhizomes of turmeric [27] is a medication for treatment of different illnesses with anti-inflammatory [28], antifungal and antioxidant [28], antimicrobial [29] and anti-cancer toward various tumor cells [30-33] activities. In vitro studies showed that curcumin is more efficient against tumor cells compared to drugs such as doxorubicin and 5-fluorouracil [34]. Curcumin has a light absorption in a range of 300 to 500 nm enable its applications in phototherapy [35], and studies have shown that phototherapy makes an improvement in the therapeutic property of curcumin [36-40]. Nevertheless, due to its lipophilicity, poor pharmacokinetics and instability, its application is limited in medical aims. Therefore, synthesis of curcumin in the forms of nanostructures accompanied by biocompatible polymers is in progress to elevate its treatment outcome [41].

This study aims to synthesize selenium-polyethylene glycol 600-curcumin nanoparticles (Se-PEG-Cur) as a dual-mode sensitizer for phototherapy and sonotherapy using a continuous-wave laser working at 808 nm as a source of light and a US wave source, respectively of C540 (B16/F10) cell line. Some items, including ROS production, cell viability, and photothermal conversion efficiency were evaluated.

**Material and Methods**

**Materials**

In this experimental study, PEG600, dimethyl sulfoxide (DMSO), ascorbic acid and sodium selenite were purchased from Scharlu (Spain). 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazoliumbromide (MTT) and 2',7'-dichlorodi-

---

Mohammadi S. et al
hydrofluorescein diacetate (DCHF-DA) were prepared from Sigma (USA). Curcumin was purchased from Merck (Germany). Deionized (DI) water was used for solutions preparation.

**Synthesis and characterization of Se-PEG-Cur**

Sodium selenite (500 mg) was dissolved in water (500 mL). Then, PEG 600 (1000 mg) was added. Curcumin (400 mg) was dissolved in acetone (5 mL), and added to the aforementioned solution. Ascorbic acid (12 g) was dissolved in water (100 mL), and dropwise added to that mixture and mixed by stirring for 24 h. A solution with a brick red color was obtained that centrifuged at 8000 rpm during 30 min. Then it was washed with cold ethanol and washed multiple times with water.

UV-visible absorption spectra of the synthesized nanostructure were recorded using a Rayleigh UV2601 spectrophotometer (China). Analysis of particle size and zeta potential of the Se-PEG-Cur suspensions were performed by a SZ-100 HORIBA instrument (Japan).

**NIR diode laser and US instruments**

Irradiation with 808-nm light was done using a Thorlabs diode laser (USA), which has an output power of 1000 mW. By changing the lens distance to the target, power density of laser radiation was set at 1.0 W cm⁻². Time of laser light irradiation was 10 min.

An ultrasonic instrument of Novin (Iran) was utilized for US irradiation and the US transducer was located under 96-well culture plates. A gel covered the transducer surface. US was irradiated from the bottom of plates with the parameters of output powers of 1.0 W cm⁻² in a duty ratio of 100%, frequency of 1 MHz, and time of irradiation of 1 min.

**Cell line preparation**

A cell line of malignant melanoma C540 (B16/F10) was prepared from Pasteur Institute (Iran). The cells were cultured in Roswell Park Memorial Institute-1640 (RPMI) medium from Gibco (USA) with 1% antibiotic (penicillin streptomycin) from Gibco (USA) and 10% fetal bovine serum from Gibco (USA) at temperature 37 °C in a humidified atmosphere of cell incubator, which contained 5% CO₂.

**Calculation of photothermal conversion efficiency (η) of Se-PEG-Cur**

A Se-PEG-Cur suspension (100 µg mL⁻¹, 0.5 mL) was exposed to NIR laser to investigate its photothermal effect in the solvent. A glass cuvette cell was utilized and real time monitoring of temperature changes upon laser irradiation were measured using a thermoprobe of Lutron (Taiwan) keeping away from the laser beam. After reaching temperature to a plateau, the laser was turned off following by permitting the cuvette to cool down to room temperature. The photothermal conversion efficiency of Se-PEG-Cur was calculated based on the method previously reported [42, 43].

**Effects of laser and US irradiation with Se-PEG-Cur on cancer cell viability**

C540 (B16/F10) cells with a density of 1.0 ×10⁴ cell well⁻¹ were seeded into 96-well plates during 24 h for cell adhesion. Then, the wells were incubated in a 5% CO₂ atmosphere at 37 °C and divided into the following groups:

- L⁻N⁻ or U⁻N⁻: control C540 (B16/F10) cells without any light or Se-PEG-Cur treatment
- L⁻N⁺ or U⁻N⁺: C540 (B16/F10) cells treated with Se-PEG-Cur of various concentrations including 5, 10, 100, 250, 500 µg mL⁻¹ without any irradiation
- L⁻N⁻: C540 (B16/F10) cells incubated in the culture medium for 4 h followed by irradiation with laser light (1.0 W cm⁻²) without any Se-PEG-Cur treatment. DI water of equal volume to that employed for Se-PEG-Cur dispersion in N⁺ groups was added.
- L⁻N⁺: C540 (B16/F10) cells treated with 100 µg mL⁻¹ of Se-PEG-Cur for 4 h followed by irradiation with laser light (1.0 W cm⁻²)
- U⁻N⁻: C540 (B16/F10) cells incubated in the
Mohammadi S. et al

The culture medium for 4 h followed by irradiation with US wave (1.0 W cm⁻²) without any Se-PEG-Cur treatment. DI water of equal volume to that employed for Se-PEG-Cur dispersion in N⁺ groups was added.

U⁺N⁺: C540 (B16/F10) cells treated with 100 µg mL⁻¹ of Se-PEG-Cur for 4 h followed by irradiation with US wave (1.0 W cm⁻²), and then immediately irradiated with US (1.0 W cm⁻²) without any Se-PEG-Cur treatment. DI water of equal volume to that employed for Se-PEG-Cur dispersion in N⁺ groups was added.

L⁺N⁺: C540 (B16/F10) cells incubated in the culture medium for 4 h followed by irradiation with laser light (1.0 W cm⁻²), and then immediately irradiated with US (1.0 W cm⁻²) without any Se-PEG-Cur treatment. DI water of equal volume to that employed for Se-PEG-Cur dispersion in N⁺ groups was added.

L⁺U⁺N⁺: C540 (B16/F10) cells treated with 100 µg mL⁻¹ of Se-PEG-Cur for 4 h followed by irradiation with laser light (1.0 W cm⁻²), and then immediately irradiated with US (1.0 W cm⁻²).

The time of incubation for various groups of the cells was 24 or 72 hours at 37 °C and 5% CO₂. It should be noted that for 24 h of incubation, the cells were incubated with Se-PEG-Cur for 24 h following by washing and performing the MTT assay, as described below. However, for 72 h of incubation, the cells were incubated with Se-PEG-Cur for 24 h, then they washed off from Se-PEG-Cur, a fresh culture medium was added, and incubation was continued for another 48 h. Finally, following by washing the cells, the MTT assay was performed as described below. The cell viability was assayed by the MTT method. The cell medium was replaced with 100 µL of 0.5 mg mL⁻¹ MTT, which was incubated at 37 °C for 4 h in dark and dissolved in phosphate buffer saline (PBS). Then, the plates were centrifuged for 15 min to remove the supernatants. Next, 100 µL of DMSO was added in order to dissolve the MTT formazan crystals. After centrifuging for 10 min, the supernatant was eliminated from every well, and the optical density (OD) was measured at 570 nm using a microplate reader from Biotek (USA). Finally, the viability of the cells was stated as the ratio of the 570 nm absorbance of the treated cells and control ones.

Detection of intracellular ROS

The DCF assay was applied to measure intracellular ROS level. Briefly, C540 (B16/F10) cells at a density of 1.0 ×10⁴ cell well⁻¹ were seeded in 96-well plates. Various cell groups (Section 2.6) were treated after 24 h. After incubation with 100 µg mL⁻¹ of Se-PEG-Cur for 3.5 h, 100 µL of a fresh DCHF-DA solution (50 µmol L⁻¹) was added to these treated cells. After laser or US irradiation, the cells were incubated for 30 min followed by washing (3 times) with PBS to remove the extracellular DCF. Then, 100 µL of a lysis buffer containing 150 mmol L⁻¹ NaCl+0.1% Triton X-100+50 mmol L⁻¹ Tris-HCl at pH=8.0 was added to the wells. The intensity of fluorescence emission at 520 nm was then measured upon excitation at 485 nm after 30 min by amicroplate reader of Biotek (USA).

Statistical analysis

For each sample, at least three parallel experiments were performed. p-values less than 0.05 were considered statistically significant through the t-test.

Results

An absorption spectrum of Se-PEG-Cur is shown in Figure 1A. To determine the Se-PEG-Cur size, dynamic light scattering was performed, and the results are shown in Figure 1B. The mean particle size of Se-PEG-Cur was obtained as ~300 nm with a uniform dispersion index (PDI~0.1). Zeta potential of Se-PEG-Cur was also measured to be -42.7 mV, as shown in Figure 1C. This value of zeta potential led to a high stability of Se-PEG-Cur without formation of aggregates [44].

To investigate the photothermal conversion efficiency (η) of Se-PEG-Cur, a rise in temperature was measured for a 100 µg mL⁻¹ Se-PEG-Cur suspension with 808-nm irradiation at a power density of 1.0 W cm⁻². The results
Selenium Nanoparticle as Photo/Sono-Sensitizer

showed a temperature increment of 11.5 °C for the Se-PEG-Cur suspension, as shown in Figure 1D, while, the control had a small temperature increment. η was calculated according to the equation:

\[ η = \frac{hs (T_{MAX} - T_{SURR}) - Q_{DIS}}{I (1 - 10^{-A})} \]  

where \( h \), \( s \), \( I \), and \( A \) are the coefficient of heat transfer, the container surface area, power density of laser (equal to 1.0 W cm\(^{-2}\)) and absorbance at wavelength of 808 nm, respectively, and \( T_{MAX} \) and \( T_{SURR} \) are the equilibrium and ambient temperatures, respectively. \( Q_{DIS} \) is the heat dispersed from light, is absorbed by the cuvette, and was independently measured to be 7.0 mW as the cuvette contained pure water (without Se-PEG-Cur NPs). The product of \( hs \) is:

\[ hs = mC / τ \]  

where \( m \) and \( C \) are the mass and specific heat of pure water, and \( τ \) is slope of dependency of cooling time on \( -\ln θ \), and θ is:

\[ θ = (T - T_{SURR}) / (T_{MAX} - T_{SURR}) \]  

**Figure 1:** UV-vis spectra recorded for the Se-PEG-Cur (A), size distribution (B), zeta potential (C), and temperature changes of Se-PEG-Cur of 100 µg mL\(^{-1}\) in water upon laser irradiation (to reach \( T_{MAX} \)) subsequent by laser light shutting off (cooling). Inset: Dependency of \( -\ln θ \) on \( t \)
Dependency of cooling time on \(-\ln \theta\) is shown in the inset of Figure 1D, and considering a value of 0.93 for \(A_{sos}\) for the Se-PEG-Cur, \(\eta\) for Se-PEG-Cur was obtained as 16.7%.

To evaluate the efficacy of Se-PEG-Cur upon PTT, SDT and PTT/SDT on the C540 (B16/F10) cells, the cell viability was assessed by the MTT assay. First, in vitro cytotoxicity of various concentrations (5-500 µg mL\(^{-1}\)) of Se-PEG-Cur without any US or laser radiation was measured, and the viability percentages of the cells treated with Se-PEG-Cur after 24 or 72 h of incubation (in comparison with untreated ones) are shown in Figure 2A, B. Figure 2A indicated upon increasing Se-PEG-Cur concentration, the cell viability decreased after 24 h incubation, and reached 30% in the presence of 500 µg mL\(^{-1}\) Se-PEG-Cur. Figure 2B also indicated that the cell viability decreased upon increasing in the Se-PEG-Cur concentration after 72 h incubation, and the viability reached 22% in the presence of 500 µg mL\(^{-1}\) Se-PEG-Cur.

In following, PTT and SDT effects of Se-PEG-Cur on viability of C540 (B16/F10) cells were investigated. Figure 2C shows viabilities of different groups of cell including L-N, U-N, L’N, U’N, L-N+, U-N+, L’N+, U’N+ and L’U’N+ after 72 h of incubation.

Figure 3 shows the fluorescence intensity

![Graph A](image1.png)  
![Graph B](image2.png)  
![Graph C](image3.png)

**Figure 2:** Viability of C540 (B16/F10) cell after exposure to Se-PEG-Cur (5-500 µg mL\(^{-1}\)) after 24 h (A), after 72 h (B) incubation time, and Viability of C540 (B16/F10) cell after exposure to Se-PEG-Cur (100 µg mL\(^{-1}\)) and radiation with laser and/or US (C).
Selenium Nanoparticle as Photo/Sono-Sensitizer

Discussion

Recently, melanoma incidence has nearly reached epidemic proportion. By current clinical tools, it is not possible to cure late-stage melanoma [45]. For treatment of cancer, it is necessary to replace conventional therapeutic strategies with non-invasive methods. These novel methods aim to early cancer diagnosis, reduce side effects and preserve normal tissues [46, 47]. An absorption peak of Se-PEG-Cur at about 212 nm can be related to PEG, and the broad peaks at longer wavelengths are due to overlapping the selenium and curcumin of Se-PEG-Cur. The ability of Se-PEG-Cur to absorb 808-nm light is witnessed from the spectrum. The value of zeta potential led to a high stability of Se-PEG-Cur without formation of aggregates [44]. Also, η value for Se-PEG-Cur was obtained as 16.7%.

The viability results (Figures 2A and 2B) showed the toxicity behavior of Se-PEG-Cur. Statistical analysis of the results indicated a noticeable influence of Se-PEG-Cur at concentrations more than 10 μg mL\(^{-1}\) on the C540 (B16/F10) cells viability, compared to the untreated cells. However, the cytotoxicity of Se-PEG-Cur at high concentrations of 50 to 500 μg mL\(^{-1}\) was almost the same (with no significant differences) maybe due to an opposite behavior of selenium and curcumin parts of Se-PEG-Cur; the former would generated ROS and the later would act as an antioxidant. This hypothesis needs more investigations. Anyway, 100 μg mL\(^{-1}\) Se-PEG-Cur and 72 h of incubation were selected for further investigations. It should also be added that the results obtained for 72 h of incubation indicated that although the cells were washed off from Se-PEG-Cur after 24 h, continuing the incubation without Se-PEG-Cur represented decrement in the cell viability. This pointed out that Se-PEG-Cur penetrated inside the cells, and its effect was prolonged to the end of incubation time.

The viability results in Figure 2C, revealed that irradiation of laser in L\(^-\)N\(^-\) led to a 7% decrement in the C540 (B16/F10) cell viability, compared to L\(^-\)N\(^-\) with no significant difference. The viability of cells in L\(^-\)N\(^+\) had a noticeable decrease (p<0.05), in comparison with the cell viability of L\(^-\)N\(^-\), L\(^-\)N\(^+\) and L\(^-\)N\(^-\). The difference between the cell viabilities of L\(^-\)N\(^-\)
and L‘N‘ was about 14%. Thus, laser irradiation made an effective photothermal treatment with the assistant of Se-PEG-Cur as a photosensitizer. On the other hand, the results indicated that US irradiation in U‘N‘ reduced the C540 (B16/F10) cell viability about 23% in comparison with the control cells (U‘N‘). Besides, the cell viability in U‘N‘ noticeably decreased (p<0.05), in comparison with the cell viability of U‘N‘, U‘N‘ and U‘N‘. There was a 12% difference between the cell viabilities in U‘N‘ and U‘N‘. In order to evaluate the combined effect of laser and US irradiation in the presence of Se-PEG-Cur, the cell viability of L‘U‘N‘ was evaluated. It was found that there was no significant decrease in the cell viability of L‘U‘N‘ (p>0.05), compared to the cell viabilities of U‘N‘ and L‘N‘. Thus, to attain a significant efficiency from dual radiation of laser and US in the presence of Se-PEG-Cur, repeated treatments by laser or US would be useful.

The results in Figure 3 showed that ROS production in different groups had the order of (with significant differences) L‘N‘ or U‘N‘<L‘N‘<U‘N‘<L‘N‘<U‘N‘<L‘U‘N‘. Although laser radiation alone induced a very low ROS level compared to untreated cells, its radiation in the presence of Se-PEG-Cur produced ROS upon the PTT process. On the other hand, either US irradiation or Se-PEG-Cur alone induced ROS production in C540 (B16/F10) cells. US radiation in the presence of Se-PEG-Cur exacerbated ROS production along with the SDT process. The ROS generation upon dual treatment of laser and US exposure in the absence of Se-PEG-Cur was lower than that generated upon dual treatment in the presence of Se-PEG-Cur; dual PTT and SDT induced the highest ROS production, and this was arose from the sensitizing role of Se-PEG-Cur.

**Conclusion**

We synthesized and characterized Se-PEG-Cur as an efficient NIR and US absorbing agent, and therefore as a sensitizer for both PTT and SDT to kill melanoma cancer cells via thermal effects and ROS generation, respectively. More investigations are needed to explore the role of the sensitizer components (Se, PEG and Cur) to identify the ROS generating species as well as antioxidant active component(s). In vivo applicability of Se-PEG-Cur would be another subject of further investigations.

**Acknowledgment**

This paper has been extracted from S. Mohammadi’s MSc thesis supported by the Research Council of Shiraz University of Medical Sciences (21051).

**Conflict of Interest**

None

**References**


7. Lal S, Clare SE, Halas NJ. Nanoshell-enabled pho-
Selenium Nanoparticle as Photo/Sono-Sensitizer


28. Martins CVB, Da Silva DL, Neres ATM, Magalhaes
Mohammadi S. et al


