Targeted peptide-modified oxidized mesoporous carbon nanospheres for chemo-thermo combined therapy of ovarian cancer in vitro

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ABSTRACT

Ovarian cancer remains one of serious hazards to human health due to many drawbacks of existing available treatment options. In this study, a multifunctional chemo-thermo combined therapy nanoplatform (OMCNPID) was successfully prepared, which is composed of I₆P₈ peptide as a targeting moiety to interleukin-6 receptors (IL-6Rs), oxidized mesoporous carbon nanospheres (OMCN) as a near infrared (NIR)-triggered drug carrier and doxorubicin (DOX) as a chemotherapeutic drug and fluorescent agent. The synthesized multifunctional nanoplatform displayed high storage capacity for drugs and excellent photothermal properties. Besides, DOX was rapidly released from OMCNPID at the condition of low pH and NIR laser irradiation due to the dissociation of DOX from graphitic cores of OMCN. In vitro experimental results verified that OMCNPID could be markedly taken up by SKOV-3 monolayer cells and tumor spheroids, and revealed a remarkable synergistic chemo-photothermal effect against ovarian cancer. All the results demonstrated that OMCNPID is a pH/NIR dual-stimulus responsive nanoplatform and can achieve efficient chemo-thermo combined therapy.

1. Introduction

Ovarian cancer is a dominant cause of gynecologic malignancy-related death worldwide (Menon et al., 2021). Despite cytoreductive surgery and conventional chemotherapy, clinical data have indicated that the five-year survival rate with advanced stage is no more than 25% (Lheureux et al., 2019). This high mortality rate is attributed to lots of factors, including lack of effective treatment modalities, insufficient drug dosage to diseased regions, severe side effects, multidrug resistance and nonspecific drug delivery (Dai et al., 2020; Kim et al., 2021; Zhang et al., 2021). With this understanding, it is urgent to explore innovative therapeutic strategies to address these challenges for therapy of ovarian cancer.

Recently, multimodal therapy which simultaneously administers two or more monotherapies like chemotherapy, photothermal therapy (PTT), immunotherapy, gene therapy and photodynamic therapy, have offered a myriad of opportunities to achieve excellent outcomes (Gotwals et al., 2017; Wang et al., 2019; Zhan et al., 2020; Hao et al., 2021; Tang et al., 2021). Among the many cancer treatments, chemotherapy, which uses drugs such as DOX, platinum and paclitaxel to destroy or damage cancer cells, is remain the indispensable treatment for ovarian cancer (Harter et al., 2010; Chuan et al., 2021; Uruski et al., 2021). Hence, the burgeoning research interests have been focused on combining chemotherapy with other types of antitumor treatment including PTT. PTT, a well-established treatment mode, has recently emerged as an efficient minimally invasive therapy. PTT employs hyperthermia generated from the absorbed near-infrared (NIR) light by photothermal agents to ablate cancer cells and offers a series of distinct merits, including deep penetration, precise spatial-temporal selectivity, minimal trauma to health tissues and enhanced sensitivity of chemotherapy (Kim et al., 2016; Pallavicini et al., 2021). With advances in nanotechnology, the therapeutic efficacy can be further enhanced by employment of multifunctional nanomaterials as PTT agents, which will enable additional drug carrier or/and cancer imaging functionalities and make the increase of the drug accumulation at tumor sites possible. As such, PTT agents mainly focus on various noble-metal nanomaterials, 2D-transition metal dichalcogenides, Cu-based semiconductor nanomaterials and carbon-based nanomaterials (Jain et al., 2008; Huang et al., 2017; Levin et al., 2019; Murugan et al., 2019; Wang et al., 2020). In our previous work, oxidized mesoporous carbon nanospheres (OMCN) could been regarded as prospective PTT reagents and nanocarriers due to their outstanding photothermal conversion ability and excellent drug loading capacity (Wang et al., 2017). Therefore, it is
reasonable to expect that the synergistic therapy of PTT and chemotherapy based on OMCN could enhance the effectiveness of ovarian cancer treatment. Besides, internal (acidic condition) and external (NIR) stimulation are beneficial to improve the drug release from carbon-based nanomaterials (including OMCN), which relies on the absorption of some aromatic drugs via simple π-stacking and the remarkable photothermal sensitivity (Yaghoubi & Ramazani, 2020; Do et al., 2021). However, no study was reported to investigate the employment of OMCN-based nanoplatform for treating ovarian cancer.

Besides, some nanoplatforms prone to accumulate inside tumor sites with a passive manner through the enhanced permeability and retention (EPR) effect. However, this passive manner has limitations due to its random delivery form. Active tumor targeting has been achieved by surface modification of nanoplatforms with ligands or antibodies, which may realize more accurate tumor delivery. A novel peptide I6P8 (sequenced CLSLITRL) was screened by a phage display technology on the cells expressing interleukin-6 receptor (IL-6R) (Wang et al., 2017). In this regard, for IL-6R overexpressed tumors, I6P8 may be a potential ligand for targeting delivery of drugs. Also, studies have revealed that IL-6R is significantly overexpressed in ovarian cancer cells, but is sub-expressed in normal tissues (Rath et al., 2010; Coward et al., 2011). Hence, I6P8 peptide was selected for designing nanoplatform to target ovarian cancer.

In this work, we designed a simple but versatile DOX-loaded OMCN nanoplatform to realize combined chemotherapy and PTT, by decorating the I6P8 peptide to target the IL-6R over-expressed in ovarian cancer. Our results demonstrated that the nanoplatform displayed high drug loading ability, excellent photothermal effect and pH- and thermo-triggered drug release. Furthermore, in vitro ovarian cell culture models were used to verify the tumor specificity, cellular uptake pathway and antitumor activity of the nanoplatform.

2. Materials and methods

2.1. Materials

SINOPHARM (Shanghai, China) provided the raw ingredients for the production of OMCNPID, including phenol, NaOH, formalin, N-Hydroxysuccinimide (NHS) and other reagents, while Sigma-Aldrich (St. Louis, USA) provided the triblock copolymer Pluronic F127 and the live-dead kit. JenKem Technology (Beijing, China) provided Maleimide PEG Amine (NH2-PEG3500-MAL) and Methoxy PEG Amine (NH2-PEG3500-OMe); ZiYu Biotech (Beijing, China) supplied the I6P8 peptide; Huafeng United Technology (Beijing, China) supplied the doxorubicin (DOX); and Dojindo Laboratories (Kumamoto, Japan) provided the Cell Counting Kit-8 (CCK-8) for this study.

2.2. Synthesis and surface modification of OMCN

In a typical synthesis (Wang et al., 2017), phenol (0.6 g) and formalin aqueous phase (37 wt percent, 2.1 mL) were mixed in a NaOH aqueous medium (0.1 M, 15 mL) and agitated for 0.5 hours at 70 °C. Then triblock copolymer Pluronic F127 aqueous solution (0.064 g/mL, 15 mL) and deionized water (50 mL) were then added, agitating for 2 hours and 18 hours respectively. The reaction solution was then diluted with distilled water (56 mL) and heated for 24 hours at 130 °C. Ultra-centrifugation was used to collect the product, which was then cleaned with deionized water before being freeze-dried. Finally, for carbonization, the freeze-dried powder was put in a nitrogen flow and heated up programmatic. To obtain OMCN, the aforementioned synthesized carbon powder was refluxed at 60 °C for 4 hours in a mixed acid solution (VH2SO4/VHNO3 =3/1).

Two-step processes were used to modify the surface of OMCN. To activate the carboxylic group, OMCN (4 mg) was submerged in an EDC/NHS/MES solution (40 mg EDC and 72 mg NHS in 4 mL MES buffer, pH 6.0) and swirled at room temperature for 2 hours. Then, ultra-centrifugation at 4500 g for 5 minutes was used to purify the product. Excessive NH2-PEG3500-MAL (5.6 mg) and NH2-PEG3500-OMe (5.6 mg) were then added to the MES buffer solution (pH 6.0, 8 mL) and mixed for 4 hours, harvesting PEGylation OMCN (OMCNP). The I6P8 peptide (2 mg) was then added to OMCNP suspension (PBS buffer, pH 7.0) and stirred for 24 hours at ambient temperature, obtaining I6P8 peptide-modified OMCNP (OMCNPID). Finally, ultra-centrifugation was used to eliminate any remaining compounds.

2.3. DOX loading

DOX was loaded prior to the above-mentioned OMCN surface modification for maximal drug loading. In short, the prepared OMCN and DOX with equal mass were submerged in PBS buffer (pH 7.0) and swirled in the dark for 24 hours. Ultra-centrifugation was used to collect the product, and the unloaded DOX remained in the supernatant. Subtracting the supernatant DOX from the total DOX measured using a fluorescence spectrophotometer yielded the mass of DOX loaded in OMCN. The following mathematical formulae were used to determine OMCNPID's drug loading capacity (DLC, %) and encapsulation efficiency (EE, %):

\[
DLC(\%) = \frac{\text{mass of DOX loaded in OMCN}}{\text{mass of nanocarrier}} \times 100\%
\]

\[
EE(\%) = \frac{\text{mass of DOX loaded in OMCN}}{\text{mass of total DOX}} \times 100\%
\]

2.4. Characterizations

A dynamic light scattering (DLS) spectrometer and a transmission electron microscope (TEM) were used to measure particle size, zeta potential and morphology, respectively. A Fourier transform infrared (FT-IR) spectrometer was used to measure the FT-IR spectra. A UV-vis spectrophotometry was used to measure the absorption spectra. A fluorescence spectrophotometer was used to record the fluorescence...
spectra. A laser scanning confocal microscope or an inverted fluorescence microscope was used to obtain all cell pictures.

2.5. Photothermal effect

Various concentrations of OMCNPID aqueous solution (0, 15, 30, 45, 60, and 75 μg/mL) were placed in plastic tubes and irradiated with an 808-nm laser at a series of power densities (2, 3, and 4 W/cm²) for 5 minutes to determine the photothermal effect of OMCNPID. An infrared thermal camera was used to continually monitor the temperature of OMCNPID aqueous solution.

2.6. In vitro drug release profiles

Equal volumes of OMCNPID solutions (2 mg/ml) were put in dialysis tubing and then submerged in 10 mL three different PBS with pH of 5.0, 6.0, and 7.4 in the dark with moderate shaking at 37 °C to investigate DOX releasing amount. At the specified intervals, aliquots (1 mL) were withdrawn and replaced with an equivalent volume of fresh PBS. A similar method was followed for the NIR-triggered groups, but the solution was irradiated with an NIR laser (3 W/cm²) for 5 minutes before the release solution was withdrawn. A microplate reader was used to track the amount of DOX release.

2.7. Cell culture

The Cell Bank of the CAS provided two distinct cell lines for this study: human ovarian cancer epithelial cells (SKOV-3 cells) and normal human ovarian epithelial cells (IOSE-80 cells). SKOV-3 cells were typically cultured in McCoy’s 5A medium, which included 10% heat-inactivated fetal bovine serum (FBS), 2.2 g/L NaHCO₃, 1% L-glutamine and penicillin-streptomycin. IOSE-80 cells were cultured in RPMI 1640 medium supplemented by 10% FBS, 1% L-glutamine and penicillin-streptomycin. In a 5% CO₂ water-saturated incubator, cells were kept at 37 °C.

2.8. Western blot analysis

In 6-well plates, SKOV-3 and IOSE-80 cells were cultured at a density of 1 × 10⁵ cells per well, respectively. The cells in each well were collected and lysed with cell lysis buffer after 24 hours of culturing. Cell lysates were centrifuged at 13500 rpm for 10 minutes at 4 °C and then the supernatant was collected and total protein content was determined using a BCA Protein Assay Kit. Following that, 30 μg proteins were added to 10% SDS-PAGE gels and then transferred to PVDF membranes. The membrane was treated overnight with a primary rabbit anti-IL-6R antibody (1:1000 dilution) or mouse anti-GAPDH antibody (1:1000 dilution) and further cultured with HRP-conjugated secondary goat anti-rabbit antibody (1:1000 dilution) or goat anti-mouse antibody (1:1000 dilution) for 1 hour. A chemiluminescence technique was used to measure the amounts of IL-6R expression in SKOV-3 and IOSE-80 cells.

2.9. Cellular uptake

SKOV-3 and IOSE-80 cells were cultured at 1 × 10⁵ cells/dish in confocal cell culture dish to test targeting cellular uptake. 1.5 mL media containing OMCNPD and OMCNPID with a DOX content of 50 μg/mL was individually introduced to the cells after 24-hours of adhesion and cultured for 1 hour at 37 °C. SKOV-3 cells were treated with excess I₆P₈ peptide for 20 minutes at 37 °C before aforesaid 1 hour incubation with OMCNPID to examine I₆P₈ peptide-mediated cellular uptake by ovarian cancer cells. SKOV-3 cells were treated with 50 μg/mL OMCNPID for 1 hour at 37 °C and 4 °C respectively to investigate energy-dependent cellular uptake. The cells were seen using a laser scanning confocal microscope after being rinsed and immobilized.

2.10. Tumor spheroid penetration

SKOV-3 cells were cultured in 48-well plates covered with 2% sterile low-melting-point agarose to create three-dimensional (3D) tumor spheroids. Tumor spheroids were treated for 12 hours with OMCNPID and OMCNPD, respectively, seven days later. After being rinsed with cold PBS and fixed with 4% paraformaldehyde, a laser scanning confocal microscope was used to scan the tumor microspheres every 20 μm from top to bottom.

2.11. In vitro antitumor activity

A live-dead kit was used to test the cytotoxicity of the drug-loaded nanoplatform in SKOV-3 cells. Briefly, SKOV-3 cells were cultured at a density of 1 × 10⁴ cells/well in 96-well plates and incubated for 24 hours. Then cells were treated with saline, OMCNPID, OMCNPID + NIR, OMCNPD and OMCNPID + NIR with OMCN concentration at 45 μg/mL. After a 6-hour incubation period, the photothermal treatment groups’ cells were subjected to a 3 W/cm² NIR laser at 808 nm for 5 minutes. All the cells were then rinsed and cultured for another 42 hours in new complete medium. After the incubation period, a live-dead kit was used to assay cytotoxicity, and a CCK-8 kit was used for quantitative evaluation.

2.12. Statistical analysis

The mean ± standard deviation was displayed for each value (SD). Analysis of variance (ANOVA) was used to conduct statistical analysis of multigroup trials, with a P value < 0.05 regarded statistically significant.

3. Results and discussion

3.1. Synthesis and characterizations

As shown in Scheme 1, the fabrication procedure of OMCNPID included two main steps. The first step was DOX loading into the mesopores of OMCN, and the second step was I₆P₈ peptide fuzing on the surface of OMCN via the PEG modification. By calculating with fluorescence, the DLC (%) and EE (%) of
OMCNPID reached up to approximately 86.4%, which were much higher than those of other carriers (Xu et al., 2020; Dinçer et al., 2022; Galhano et al., 2022).

To confirm the synthesis, TEM, particle size, zeta potential, FT-IR, UV-vis-NIR and fluorescence spectra were carried out. The morphology of OMCN was characterized by TEM. Figure 1A revealed OMCN had a regular near-spherical shape with mesoporous structure and were well-dispersed in water. Surface decoration with PEG and I₆P₈ peptides increased the average hydrodynamic size determined by dynamic laser scattering (DLS) from 131.6 nm for OMCN to 164.1 nm for OMCNP and 198.3 nm for OMCNPID (Figure 1B). Successful modification was further proved by the surface zeta potential in water. The zeta potential of OMCN was highly negative (-46.5 ± 2.3 mV) attributable to the fact that there are a number of carboxy groups on their surface (Figure 1C). On account of the PEGylation shielding effect and the covalent binding of positive-charged I₆P₈ peptide, the zeta potentials demonstrated a growing tendency (-32.4 ± 1.6 mV for OMCNP and −27.6 ± 2.9 mV for OMCNPID).

Moreover, the successful coupling of PEG and conjugation of I₆P₈ peptide on OMCN were evaluated by FT-IR analysis. In the FT-IR spectrum of OMCN, the broad absorbance appeared in around 3400 cm⁻¹ was ascribed to the secondary amine stretching vibration, while the peaks at 1735 cm⁻¹ was designated to carbonyl stretching vibration of the carboxy groups (Figure 1D). After OMCN reacted with PEG and I₆P₈, the characteristic bands at 3445 cm⁻¹ and 3516 cm⁻¹ (primary amide stretching vibration, νN-H) which belong to PEG and 2550 cm⁻¹ (S-H stretching vibration, νS-H) which belongs to I₆P₈ were disappeared. Meanwhile, some new characteristic peaks appeared at 1620 cm⁻¹ (amide carbonyl stretching vibration, νC = O(NH)), 1540 cm⁻¹ (amide II band, δN-H + νC-N) and 1013 cm⁻¹ (C-S-C stretching vibration, νC-S-C) indicated the successful synthesis of the OMCNPID. As can been seen in Figure 1E, the UV−vis absorption curve of OMCNPID exhibited a relatively strong peak at 488 nm, which belongs to the characteristic absorption of DOX, suggesting that DOX was successfully loaded into OMCNPID. In addition, obvious absorption in the NIR region (including 808 nm) was exhibited in the UV−vis spectra of OMCN, OMCNPID and OMCNPID, indicating that OMCN-based carriers were suitable candidates for PTT upon 808 nm laser irradiation. Figure 1F demonstrated the fluorescence spectra of various products in the process of

Scheme 1. Schematic illustration of synthetic process of OMCNPID and its drug release.

Figure 1. Characterizations of nanoparticles. (A) TEM image of OMCN. Bar = 50 nm. (B) DLS and (C) zeta potential measurements of OMCN, OMCNP and OMCNPID. Data were presented as mean ± S.D. (n = 3). (D) FT-IR spectra of OMCN, PEG, I₆P₈ and OMCNPID. (E) UV-vis-NIR spectra of OMCN, OMCNPID, DOX and OMCNPID. (F) Fluorescence spectra of free DOX, the mixture of OMCNPID and DOX, supernatant and OMCNPID.
fabricating OMCNPID (Ex = 488 nm). It can be noted that the fluorescence intensity of OMCNPID was much inferior to that of free DOX and the mixture of OMCNP and DOX at equal concentrations, which might be owing to the fact that the fluorescence signal of DOX would be quenched by OMCN when they aggregated at high concentrations.

3.2. Photothermal effect

The in vitro photothermal performance of OMCNP was evaluated by detecting the temperature change of OMCNP aqueous solutions at different concentrations during 5 minutes laser irradiation. The temperature profiles and photothermal images were showed in Figure 2. As expected, the NIR-triggered photothermal effect was highly dependent on the concentration of OMCNP and laser power intensity. For pure water (0 μg/mL OMCNP), the temperature increased minimally after irradiation for 5 minutes. On the contrary, the temperature of OMCNP aqueous solutions increased rapidly when the concentration increased from 15 to 75 μg/mL at the same power intensity (Figure 2A). In the meantime, the temperature of OMCNP aqueous solutions gradually increased as the power intensity ranging from 2 W/cm² to 4 W/cm² (Figure 2B). Moreover, the temperature of 45 μg/mL OMCNP aqueous solution could exceed 50 °C under a power density of 3 W/cm², reaching the biological photolethal limit of the cells. These findings revealed that OMCNP could serve as a great potential near-infrared agent for PTT.

3.3. In vitro drug release profiles

As illustrated in Figure 3, the release of drug from OMCNPID triggered by pH change and NIR laser irradiation were investigated. It was observed that only about 19.5% DOX was released from OMCNPID in pH 7.4 PBS after 48 hours, which validated the stable loading of DOX into OMCNPID under physiological condition (Figure 3A). However, the cumulative drug release in pH 6.0 PBS and pH 5.0 PBS increased up to 47.2% and 66.9% within the same time period, suggesting that the DOX release from OMCNPID was significantly pH-triggered. This phenomenon could be ascribed to the protonation of amino groups on DOX under acidic conditions, thus increasing electrostatic repulsion between DOX and graphitic cores of OMCN. Additionally, the amount of DOX released from OMCNPID after exposure to NIR laser for 5 minutes at predetermined time demonstrated a noticeable increase in comparison with that without NIR laser exposure in all release conditions (Figure 3B). The cumulative release amount was 82.4% under the condition of pH 5.0 PBS and NIR laser irradiation, which was more than that under the single condition of pH 5.0 PBS (66.9%), implying that OMCNPID had NIR-triggered drug release properties. Consequently, OMCNPID
could realize remotely controlled drug release under the dual-stimulus including tumor microenvironment (acidic condition) and outer environment (NIR) to decrease the side effect and enhance the synergistic therapeutic effect.

3.4. IL-6R expression level in specific cells

IL-6R expression level was testified on both SKOV-3 cells and IOSE-80 cells by western blot (Figure 4). Densitometry studies revealed the high IL-6R expression on SKOV-3 cells, which was consistent with previous reports (Coward et al., 2011), making SKOV-3 cells an ideal model for ovarian cancer targeting delivery and treatment. Nevertheless, there was low IL-6R expression on IOSE-80 cells, which might be deemed as a negative control to assess the function of the targeted IL-6R. Thus, this study utilized SKOV-3 cells and IOSE-80 cells for the evaluation of targeting efficiency of OMCNPID.

3.5. Cellular uptake

The tumor-targeting capability of different formulations was assessed by the uptake in SKOV-3 cells and IOSE-80 cells. As displayed in Figure 5A–L, higher fluorescence intensity of OMCNPID was found in SKOV-3 cells as compared with that of OMCNPD. Also, the fluorescence intensity of OMCNPID in SKOV-3 cells was far superior to that in IOSE-80 cells, suggesting that I₆P₈ on nanoplatform surface could significantly facilitate the nanoplatform targeting to SKOV-3 cells. However, there was no striking difference in the fluorescence intensity between the OMCNPID and OMCNPD in IOSE-80 cells, indicating the existence of I₆P₈ did not influence the uptake behavior of the nanoplatform by IOSE-80 cells. This was most likely due to the apparently higher expression on IL-6R in SKOV-3 cells than that on IOSE-80 cells.

3.6. Permeation into 3D tumor spheroids

The above findings confirmed the potential of OMCNPID as an efficient formulation with excellent cellular internalization
in a 2D SKOV-3 cell cultured monolayer model. However, the cellular uptake by monolayer cells may not reflect the tumor-accumulating effect of different formulations accurately because the drugs must be transported into the cells in the core regions of tumor, not just the surface cells. Considering this, 3D tumor spheroid could serve as a reliable in vitro evaluation model for investigating penetration behavior of different formulations. The penetrating ability of different formulations was monitored by confocal microscopy after 12 h incubation with SKOV-3 tumor spheroids. For qualitative analysis, the fluorescence intensity observed in SKOV-3 tumor spheroids treated with OMCNPID was much more prominent than that of OMCNPD (Figure 6). For quantitative analysis, OMCNPID penetrated around 73.3 μm into SKOV-3 tumor spheroids, whereas OMCNPD showed an average penetration depth of only 48.4 μm. These results suggested that I6P8 dec- oration could effectively improve nanoplatform penetration into SKOV-3 tumor spheroids. The excellent permeation ability was indispensable to ovarian cancer treatment.

3.7. In vitro antitumor activity

The in vitro antitumor activity of different formulations was assessed via live-dead assay by staining the tumor cells with Calcein-AM and PI. In this assay, live cells stained by Calcein-AM showed green fluorescence, whereas dead cells stained by PI exhibited red fluorescence. As visualized in Figure 7A, cells treated with saline or OMCNPID showed the darkest red fluorescence indicating that there were no significant dead cells. This suggested that the drug nanocarrier, OMCNPID, could cause negligible cytotoxicity. Notably, chemo-thermo combined therapy (OMCNPID + NIR group) displayed higher cell death compared to that in monotherapy (OMCNPID group or OMCNPID + NIR group). Then the antitumor activity was further quantitative validated by CCK-8 assay. As seen in Figure 7B, chemo-thermo combined therapy group resulted in lowest SKOV-3 cell viability and the cytotoxicity was about 1.69- and 1.58-fold of that of OMCNPID group and OMCNPID + NIR group, respectively. Overall, the results clearly revealed that the chemo-thermo
combined therapy exhibited significantly enhanced antitumor efficacy in vitro, which was probably due to the targeting I_P8 peptide for better cellular uptake and synergistic effect of different therapies. Certainly, further in vivo experiment is necessary to evaluate the antitumor efficacy before a final statement.

4. Conclusion

This study demonstrated that OMCNPID was fabricated as a multifunctional anti-cancer nanoplatform for chemo-thermo combined therapy of ovarian tumors. Based on the in vitro observations, the as-prepared OMCNPID revealed suitable size, high storage capacity for drugs, satisfying photothermal ability and pH- and NIR-triggered DOX release effect. Here, DOX loaded in the nanoplatform and OMCNI were utilized for chemotherapy and PTT, respectively. Cellular uptake and tumor spheroid penetration results respectively indicated that the conjugation of I_P8 peptide to OMCN significantly enhanced the IL-6R-mediated accumulation and penetration of DOX in both SKOV-3 monolayer cell and tumor spheroids. More importantly, OMCNPID showed remarkable antitumor efficacy in vitro, owing to the surface conjugation of targeting I_P8 peptide and excellent synergistic therapeutic effects. Therefore, the investigation validated that the multifunctional nanoplatform would open a potential new avenue for ovarian tumor therapy.

Disclosure statement

No potential conflict of interest was reported by the authors.

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