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ORIGINAL RESEARCH

Externally Triggered Novel Rapid-Release Sonosensitive Folate-Modified Liposomes for Gemcitabine: Development and Characteristics

This article was published in the following Dove Press journal: International Journal of Nanomedicine

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Purpose: To develop an externally triggered rapid-release targeted system for treating ovarian cancer, gemcitabine (GMC) was entrapped into sonosensitive (SoS) folate (Fo)-modified liposomes (LPs).

Methods: GMC-loaded LPs (GMC LPs), GMC-loaded Fo-targeted LPs (GMC-Fo LPs), and GMC-loaded Fo-targeted SoS LPs (GMC-SoS Fo LPs) were prepared utilizing a filmhydration technique and evaluated based on particle size, ζ -potential, and percentage entrapped drug. Cellular uptake of the fluorescent delivery systems in Fo-expressing ovarian cancer cells was quantified using flow cytometry. Finally, tumor-targeting ability, in vivo evaluation, and pharmacokinetic studies were performed.

Results: GMC LPs, GMC-Fo LPs, and GMC-SoS Fo LPs were successfully prepared, with sizes of <120.3±2.4 nm, 39.7 mV ζ -potential, and 86.3%±1.84% entrapped drug. Cellular uptake of GMC-SoS Fo LPs improved 6.51-fold over GMC LPs (under ultrasonic irradiation — p<0.05). However, cellular uptake of GMC-Fo LPs improved just 1.24-fold over GMC LPs (p>0.05). Biodistribution study showed that of GMC concentration in tumors treated with GMC-SoS-Fo LPs (with ultrasound) improved 2.89-fold that of free GMC (p<0.05). In vivo, GMC-SoS Fo LPs showed the highest antiproliferative and antitumor action on ovarian cancer.

Conclusion: These findings showed that externally triggered rapid-release SoS Fo-modified LPs are a promising system for delivering rapid-release drugs into tumors.

Keywords: sonosensitive liposome, gemcitabine, folate-modified liposomes, externally triggered, ovarian cancer

Introduction

Cancer is one of the most dangerous abnormalities threatening human life today. In Egyptian females, ovarian cancer is the fifth-commonest cause of death.¹ Because of an inability to detect and diagnose ovarian cancer early, it spreads within the internal viscera. As such, most ovarian cancer patients cannot be diagnosed, leading to advanced stages (III or IV).² Late diagnosis of ovarian cancer may lead to death within 5 years. Surgical removal of ovarian tumors and chemotherapy are basic approaches in standard treatment. Chemotherapy is a significant factor in treating cancer. There are many anticancer drugs, such as adriamycin, topotecan, and gemcitabine (GMC).^{3,4}

Adriamycin may act by intercalating into DNA, leading to disruption of topoisomerase II-mediated DNA repair, or by producing free radicals, leading to

683

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GMC is an attractive choice in ovarian cancer management,¹⁰ because of its well-known efficacy in platinum/paclitaxel-resistant cancer, no cross-resistance with platinum compounds, and ease in combining it with other chemotherapeutics for treating recurrent ovarian cancer.¹¹ It can be utilized for treating cancer of the ovaries, lungs, and pancreas.¹² GMC penetrates the cell membrane and transforms into GMC diphosphate and GMC triphosphate through deoxycytidine kinase. Ribonucleotide reductase is inhibited by GMC diphosphate, leading to a halt in the synthesis of deoxynucleoside triphosphates required for DNA synthesis. At the same, GMC triphosphate is incorporated into DNA instead of endogenous deoxynucleoside triphosphates.¹³ The GMC-triphosphate metabolite is also utilized RNA synthesis, inhibiting it.¹⁴ However, the hydrophilicity, short plasma half-life, and rapid metabolism of GMC lead to increases doses required for effective plasma concentration for cancer-cell death, causing many side effects, such as hematologic neutropenia occurring more often than thrombocytopenia.¹⁵ GMC is a hydrophilic drug (soluble in water at 25 mg/mL) with an octanolwater partition coefficient (logP) of 1.123. Moreover, incorporation of GMC in liposomes (LPs) enhances its cytotoxic effects compared to free drug.¹⁶

In an attempt to diminish the side effects of GMC, many delivery systems have been used, such as nanoparticles, transferosomes, and LPs.¹⁷⁻¹⁹ LPs are biocompatible, biodegradable, and safe and can be used to improve efficacy, decrease side effects, and enhance the stability of encapsulated drugs. Phospholipid bilayers of LPs may be used as membranes into which hydrophobic moieties can be inserted to target LPs toward the desired site.^{20,21} LPs are widely used as delivery systems because of relatively long circulation time, enhanced permeability-and-retention effect (EPR), and enhanced accumulation of drug into cancerous tissue, leading to improved therapeutic action and diminished toxic effects.²² Moreover, LPs have the capability of entrapping both hydrophobic (within the bilayer) and hydrophilic (within the core) drugs. They are atoxic and do not activate the immune system.²¹ Many attempts have been made to enhance the targeting ability of LPs by modifying, the surface such as PEGylated LPs and Fo-modified LPs.^{23,24} To improve cell-specific and intracellular delivery. LPs may be further conjugated with targeting ligands, such as Fo.²⁵ Currently, Fo-modified LPs are utilized based on expression of Fo receptors, which is very high in most types of cancer, while expression is low in healthy tissue.²⁶ These modified LPs have been characterized as having sustained release.²⁷ However, there is great demand for developing a delivery systemwith an enhanced targeting index and an externally triggered rapid release of drug cargo within the cancerous tissue in ovaries.²⁸

Because Fo-targeted LPs have proved to be highly effective in many medical applications, easily disrupted LPs have rarely been studied. Fo-conjugated LPs containing chlorin e_6 (Ce₆) could be studied for rapid-release capability after binding with cancerous cells.²⁹ Encapsulation of drug in long-circulating PEGylated LPs has overcome limitations experienced with other non-PEGylated LPs.³⁰ Moreover, proper PEGylation is also required, as a result of initial (and overall) tumor accumulation that still depends on passive extravasation.³¹ Active targeting (such as conjugation with Fo) is utilized to decrease disadvantages associated with passive targeting (such as decoration PEG conjugation and long circulation).³²

The usse of most antitumor drugs for treatment of cancerous tissue is restricted, as a result of their low specificity.³³ To counter this, antitumor drugs could be targeted to cancerous tissue. Moreover, high levels of

antitumor drugs are desired within cancerous tissue, in order to kill abnormal cells. To achieve this aim, externally triggered rapid-release drug carriers should be developed.

The present study was carried out to achieve rapid release of the loaded drug within tumors. Formulated LPs should be sensitive to external stimuli, such as ultrasonic waves, and once applied, the drug should be released. The present study aimed to target a drug to the ovaries utilizing externally triggered novel rapid-release sonosensitive (SoS) Fo-modified LPs. SoS Fo-modified LPs were characterized by measuring particle size, ζ -potential, and encapsulation efficiency (EE). Moreover, ultrasound sensitivity, cellular association using flow cytometry, in vivo antitumor efficacy, and pharmacokinetics of the liposomal drug were examined.

Methods

Soybean phospholipids (L- α -phosphatidylcholine [PC, purity \geq 99%), folic acid–polyethylene glycol–stearylamine (FA-PEG-SA) and cholesterol (purity \geq 99%) were purchased from Xian Ruixi Biological Technology. Ce₆ was purchased from Frontier Scientific (Salt Lake City, UT, USA). The CAOV3 cell line was obtained from the Chinese Academy of Sciences (Shanghai, China). RPMI 1640 medium and FBS were purchased from Sigma Chemical.

All reagents were of HPLC or analytical grade. Sprague Dawley rats were purchased from the animal house of the Faculty of Medicine, Assiut University, Egypt.

Preparation of Liposomal Samples

GMC LPs (GMC LPs) were prepared by the pH-gradient method using PC and cholesterol (PC:cholesterol 5.5:4.5).^{15,34} Briefly, LPs were prepared using thin-film hydration. To prepare non-Fo-modified LPs (500 mg lipid), a lipid composition of PC and cholesterol (55% and 45% molar ratio) were completely dissolved in a mixture of ether and chloroform (1:3 v:v). With a rotary evaporator (Büchi), the organic solution was evaporated under vacuum into a sterile conical flask, forming a dried thin membrane. Phosphate citrate buffer (50 mM, 10 mL, pH 2.2) was utilized to hydrate the membrane at 55°C (50 mg/mL), resulting in a colloidal solution, then the pH was raised to 7.2 using disodium hydrogen phosphate. The colloidal solution was extruded 12 times (mini-extruder, Avanti) through 100 nm polycarbonate filters, aiming to optimize the particle size, as shown in Figure S1 and S2

(supplementary section). GMC (concentration 0.9–1.6 mg/ mL) was then incubated in the LPs at 65°C for 4 hours to get active loading. Separation of free GMC was carried out using extensive dialysis against buffer (20 mM HEPES, pH 7.4, containing 150 mM NaCl and 0.1 mM EDTA). GMC-Fo LPs were prepared by the same procedure, except FA-PEG-SA was added to the lipid mixture at 0.1% molar ratio. To prepare GMC-SoS Fo LPs, a lipid composition of PC and cholesterol (55% and 45% molar ratio) and FA-PEG-SA (0.1%) was completely dissolved in a mixture of ether and chloroform (1:3 v:v). The lipophilic sonosensitizer Ce₆ ester (3%, 6% or 9%) was dissolved in the prepared organic phase. With the rotary evaporator, the organic solution was evaporated under vacuum into a sterile conical flask, forming a dried thin membrane. Then, all preparation steps used the same procedures mentioned previously. These procedures were performed under aseptic conditions and vertical laminar flow (vertical laminar flow hood, BZ series, Germfree).

Determination of Entrapment Efficiency and Drug Loading

GMC-EE and drug loading (DL) were calculated after purifying the LPs by ultrafiltration in centrifuge tubes with an Ultrafilter (Amicon Ultra, molecular weight cutoff kDa. Millipore). Quantification was 30 done spectrophotometrically.³⁵ Briefly, LP formulations (0.5 mL) were placed in the upper part of the centrifuge tubes, then centrifuged at 6,000 rpm, 4°C for 30 minutes to separate free drug from the encapsulated LPs. The last step was repeated until obtaining all LP quantities. The amount of free drug in the filtrate was estimated at 267 nm spectrophotometrically (LC20A, Shimadzu). EE and DL were calculated using Equations 1 and 2:

$$EE\% = (A_{total} - A_{free})A_{total} \times 100$$
(1)

$$DL\% = (A_{total} - A_{free})A_{lipsomes} \times 100$$
(2)

where A_{total} , A_{free} , and $A_{liposomes}$ represent GMC, free GMC, and the total LPs recovered, respectively.

Characterization of Liposomes

Morphology of GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs was examined utilizing transmission electron microscopy. Briefly, aliquots were placed on hydrophobic copper grids and dried, then negatively stained with uranyl acetate (50 μ L, 2.5% w:v). A Malvern Zetasizer Nano ZS Zen 3600 was utilized to investigate the colloidal dispersion to determine ζ -potential, particle size, and polydispersity index (PDI) values.³⁶ Briefly, LP samples (200 µL) were diluted with deionized water (800 µL) to be analyzed based on dynamic light scattering using the Zetasizer.

Evaluation of Ultrasound Sensitivity Acoustic Parameter Selection

GMC-SoS-Fo LPs were diluted eightfold with PBS (0.1 M, pH 7.4) and kept at 37°C for 10 minutes, then transferred into an ultrasound sonication system (2 MHz, 1.5 W/cm² power intensity, 10 seconds' ultrasonic irradiation with 10-second rests for a total of 0–150 seconds). Temperature of the LP preparations was fixed during ultrasonic irradiation utilizing a water bath containing a thermocouple. Drug release was determined after each ultrasonic irradiation and recorded. Colloidal dispersions were left for 10 minutes before they were purified by ultrafiltration in the Amicon centrifuge tubes.³⁵ Quantification was done spectrophotometrically at 267 nm spectrophotometrically (Shimadzu LC20A), as per determination of EE.

Molar Ratio of Sonosensitizer

GMC release from GMC-SoS Fo LPs prepared withmolar ratios of 3%, 6%, or 9% Ce₆ of total lipid was measured under the same procedures. However, the total period of ultrasonic irradiation was changed to 130 seconds.

In Vitro Drug Release

In vitro GMC release from GMC-loaded LPs was determined using dialysis bags.³⁷ Briefly, 5 mL GMC-loaded LPs was placed into a cellophane bag (molecular weight cutoff 12–14 kDa), then inserted into 100 mL PBS (pH 7.4, 0.1 M, 37°C). Dissolution media were stirred gently and treated with ultrasonic irradiation (2 MHz, 1.5 W/cm² power intensity, 10 seconds' ultrasonic irradiation with a 10-second rests for a total of 0–130 seconds) or not treated. Aliquots (2 mL) replaced with equal volume of fresh prepared PBS were withdrawn from the dissolution medium to be spectrophotometrically analyzed at at λ_{max} =267 nm for measuring released GMC. Measurements were repeated three times, and are represented as means ± SD.

Storage Stability of Liposomes

To examine the stability of colloidal dispersions, LP dispersions were kept at different temperatures ($4^{\circ}C\pm2^{\circ}C$, $25^{\circ}C\pm2^{\circ}C$, $37^{\circ}C\pm2^{\circ}C$, 65% relative humidity) for 3 months. Aliquots were taken every month to investigate particle size, ζ -potential and EE.

Cell Culture

CAOV3 cells (adherent type) were incubated in RPMI 1640 media (50 mL) containing FBS (10%, 5 mL) 1%, streptomycin (0.1 mg/mL), and penicillin (200 IU, 1 mL) into flasks at $37\pm2^{\circ}C/5\%$ CO₂. Cells were reseeded and suspended every 4 days and used for study at passages 5–20.

Quantitative Cellular Uptake of Liposomes Using Flow Cytometry

To determine the intercellular uptake of the GMC LPs, GMC-Fo LPs, and GMC-SoS Fo LPs, flow cytometry was utilized based on measuring mean fluorescence intensity.³⁸ Briefly, cells were incubated into 48-well plates at a density 6×10^5 cells/well. Incubated cells were withdrawn from the medium after 24 hours, then washed with PBS and reseeded and suspended in fresh medium for 4 hours at 37° C±2°C/5% CO2 atmosphere. Cell suspensions were incubated for 48 hours with calcein LPs, calcein-Fo LPs, and calcein-SoS Fo LPs, respectively. Subsequently, they were treated with ultrasonic irradiation (2 MHz, 1.5 W/cm² power intensity, 10 seconds' ultrasonic irradiation and with 10-second rests for a total of 0-130 seconds) or not treated, then incubated for 30 minutes at $37\pm2^{\circ}C/5\%$ CO₂. Thereafter, culture media weres removed and sterile PBS used to rinse adherent monolayers. Trypsin enzyme (10 mL, 0.25% v:v) was utilized to liberate cells from the adhesion state. Cells were suspended in sterile PBS and analyzed with flow cytometry (Thermo Fisher Scientific) at excitation wavelength 495 nm and emission wavelength 516 nm. The LP dispersions were prepared to encapsulate calcein instead of GMC following the same aforementioned procedures.

Cell suspensions incubated with calcein-free LPs were treated withultrasonic irradiation (2 MHz, 1.5 W/cm² power intensity, 10 seconds' ultrasonic irradiation and with 10-second rests for a total of 0–130 seconds) or not treated under the aforementioned conditions. The first sample (incubated with empty-LPs) was treated with ultrasonic irradiation, so was considered a control (control 1) for the tested samples treated with ultrasonic irradiation. The second sample (incubated with empty LPs) was not treated with ultrasonic irradiation. The second sample (incubated with empty LPs) was not treated with ultrasonic irradiation. The second sample (incubated with empty LPs) was not treated with ultrasonic irradiation. The second sample (incubated with empty LPs) was considered a control (control 2) for tested samples that did not receive-ultrasonic

irradiation. These controls were used to subtract the effects of autofluorescence and ultrasonic irradiation.

After that, cells were washed with PBS twice and suspended in PBS before subjecting them to flow cytometry to calculate cellular binding.measurements were repeated three times.

Cytotoxicity

To examine the cell-killing effect of SoS Fo LPs, free GMC, GMC LPs, GMC-Fo LPs, GMC-SoS Fo LPs (no ultrasonic irradiation) and GMC-SoS Fo LPs (ultrasonic irradiation), cytotoxicity was assessed. Preparations were examined by MTS assays in CAOV3 cells. Briefly, cells were seeded in 96-well plates at 105 cells/well and incubated at 37°C in 5% CO₂ for 48 hours. Thereafter, cells were treated with different concentrations (0–150 μ M) of GMC from the formulations for 72 hours. Ultrasonic irradiation was applied for 150 seconds only to the last preparation after 48 hours' incubation, then incubated for 24 hours (total incubation 72 hours). Subsequently, 20 µL MTS reagent (tetrazolium inner salt) per well was added and incubated for another 4 hours at 37°C in 5% CO₂. Absorbance at 490 nm was measured using a multimode microplate reader to calculate cell viability:

$$Cell \, viability(\%) = \left(S_{[Test]} - S_{[Blank]}\right) / \left(S_{[Control]} - S_{[Blank]}\right) \\ \times 100\%$$

where $S_{[Test]}$, $S_{[Control]}$, and $S_{[Blank]}$ represent the absorbance of the treatment group, untreated control group, and blank-culture medium group, respectively. Moreover,IC₅₀ was calculated using an online calculator (<u>https://www.aatbio.com/tools/ic50-calculator</u>).

Establishment of Ovarian Cancer Model in Sprague Dawley Rats

A xenograft ovarian cancer model in rats obtained from the Faculty of Medicine, Assiut University was produced by surgical orthotopic implantation.³⁹ Briefly, subcutaneous tumors were induced by injecting 10⁶ CAOV3 cells into the right axillary regions of immunocompromised Sprague Dawley rats aged 30 days using 0.25 mL dexamethasone acetate (4 mg/mL) daily intramuscular injections. Immunosuppressive treatment was also started 5 days before the injection of CAOV3 cells and maintained throughout the experiment.

The resulting subcutaneous tumors were allowed to grow about 0.5 cm^3 , then isolated as small pieces

(1 mm³). One piece was incubated surgically into the ovary. Development of ovarian cancer was monitored by ultrasound imaging.⁴⁰ The study was conducted under license of the ethical approval committee of the Faculty of Medicine, Assiut University (Assiut 155–020, January 20, 2020), and in aqccordance with the *Guide for Care and Use of Laboratory Animals* published by the US National Institutes of Health (eighth edition, revised 2011).

Biodistribution Studies

To evaluate in vivo targeting efficiency of GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs, a biodistribution and pharmacokinetic study was performed using 40 ovarian cancer xenograft models in the rats. Rats were randomly distributed into fife groups: the first (n=8) was intravenously injected with free GMC (2 mL, 4 mg/mL), and the second (n=8), third, fourth, and fifth were intravenously injected with LPs GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs (without ultrasound) and GMC-SoS-Fo LPs (with ultrasound) at 12.5 mg/kg. At 30 minutes after injection, the rats treated with GMC-SoS-Fo LPs were subjected to ultrasonic irradiation above the ovaries at both abdomen sides (sonication area of 2 cm² and ultrasound of 2 MHz, 1.5 W/cm² power intensity, 10 seconds' ultrasonic irradiation with 10-second rests for a total of 0-150 seconds). At 1 hour after injection, three rats from each group were humanely killed, and ovaries, kidneys, lungs, spleens, livers, and hearts collected. The drug was measured in the plasma as reported in the literature with some modification.⁴¹ Collected organs were kept at -20°C until further analysis. To detect GMC concentration, the organs collected were homogenized in fourfold volumes of double-distilled water. The homogenates (2 mL) spiked with 100 µL floxuridine (20 ng/mL) as internal standard were treated and vortexed with acetonitrile (1 mL) and centrifuged (2,000 rpm, 3 minutes) to precipitate the protein. The supernatants were evaporated under vacuum at 50°C. The dried residue was reconstituted with 2 mL mobile phase, which consisted of 40 mmol/L acetate ammonium buffer solution (pH 5.5):acetonitrile (97.5:2.5, v:v). Aliquots of reconstituted solution (50 µL) were injected into the column (C18, 250 mm×4.6 mm ID, 5 μm), and flow rate was set at 0.8 mL/minute for detection of the drug at λ_{max} =268 nm.

Pharmacokinetic Studies

Twelve female Sprague Dawley rats weighing 250–280 g were utilized for the present pharmacokinetic studies and

distributed randomly into four groups (n=3). The rats were kept under standard conditions at 25°C, relative humidity 55%±5% and free access to drinking water Pharmacokinetic study of GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs were conducted utilizing a single dose of 4 mg/kg GMC (intravenously via tail vain, diluted to 1 mL with isotonic saline solution 0.09% w:v). Blood samples (0.3 mL) were withdrawn at predetermined time intervals (5, 15, 30, 45, 60, 120, 180, 240, and 300 minutes) from the tail vein to separate the plasma. To conduct the analysis, 100 µL of floxuridine solution as internal standard (20 ng/mL) was added to each plasma sample. Plasma protein was precipitated in all samples by adding 1 mL acetonitrile to each, then vortexing for 10 minutes and centrifuging at 2,000 rpm for 4 minutes. Supernatants were withdrawn and evaporated under vacuum at 50°C. Dried residues were reconstituted with 2 mL mobile phase (40 mmol/L acetate ammonium buffer solution, pH 5.5): acetonitrile, 97.5:2.5, v:v). Analysis was performed as per the biodistribution study.

In Vivo Antitumor Efficacy

Ovarian cancer rat xenograft models were distributed into various groups, each with ten rats. The first was intravenously injected with GMC-free LPs (control group) and subjected to ultrasonic irradiation. The second, third, fourth and fifth were intravenously injected with 4 mg/kg free GMC solution, GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs, respectively. The sixth group, injected with GMC-SoS-Fo LPs, was subjected to ultrasonic irradiation. The dose regimen was intravenous injecton 4 mg drug equivalent per kilogram on days 1, 7, 10, 13, 16, and 20. Tumors in each group were measured every 3 days up to 27 days using a commercial three-dimensional sonar (Artida, Toshiba Medical Systems).⁴² Survival time (from inoculation of tumors until death) was recorded for each group and plotted. Kaplan-Meier survival curves were plotted for each group. The body weight of each mouse was measured daily.

Statistical Analysis

All findings are presented as means \pm SD. One way-ANOVAs were applied to calculate differences between any two groups. Significance was regarded as p < 0.05, with Tukey–Kramer multiple assessments or two-sided Student *t*-tests for pairwise comparisons (GraphPad Prism 6.0, GraphPad, San Diego, CA, USA).

Results and Discussion

Targeting of drug-delivery systems into tumors is very important. There is currently huge demand to achieve rapid release of drugs after targeting tumor sites.

Characterization of Liposomes

Physical properties of the prepared LPs were investigated by measuring particle size, EE, and DL (Table 1). Mean particle sizes of GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs were 120.3±2.4, 119.1±3.5, and 109.5±2.9 nm, respectively. Although a mini-extruder (100 nm polycarbonate filter) was used to optimize particle size, mean particle size was >100 nm. These results may be interpreted on the basis of elasticity of the LPs: they were squeezed through the polycarbonate filter under high pressure. After the LPs had passed the extruder, they attained their final size. Moreover, PDI values of all LPs was 0.209-0.272, suggesting high homogeneity. The EE of the prepared LPs was 82.8%-86.3%, confirming that conjugation of LPs with Fo moiety (GMC-Fo LPs) or Fo and Ce₆ (GMC-SoS-Fo LPs) did not affect EE significantly (p>0.05). The ζ -potential of all LP formulations was more than $>34.52\pm1.01$ mV, as presented in Table 1, confirming stability of all formulations. Finally, DL of LPs was 3.45%±0.12%-3.73%±0.27%, showing that conjugation of the LPs had no effect on DL, as presented in Table 1. The small differences among GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs could be interpreted on the basis of the same lipid composition, while the other ingredients ----Fo moiety (GMC-Fo LPs) or with Fo and Ce₆ — were formulated with neglected ratio. Moreover, most Fo

Table	I.	Physical	Properties	of the	Nanocarriers
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	Particle Size (nm)	PDI	ZP ± SD (mV)	EE (%)	Drug Concentration (mg/mL)	Drug Loading (%)
GMC LPs	120.3±2.4	0.272±0.005	37.24±0.97	86.3±1.84	0.1865	3.73±0.12
GMC-Fo LPs	9. ±3.5	0.228±0.009	34.52±1.01	82.8±1.12	0.1725	3.45±0.31
GMC-SoS-Fo LPs	109.5±2.9	0.209±0.002	39.7±0.51	84.6±2.04	0.1805	3.61±0.27

Abbreviations: PDI, polydispersity index; ZP, ζ-potential; EE, encapsulation efficiency. Values are means of three experiments.

moieties were outside the phospholipid-bilayer membrane, so they may not have interfered with or had little effect on the phospholipid-bilayer membrane.

The ζ-potential of LPs may have great influence on drug accumulation within cells. Accumulation of cationic LPs is high after intravenous injection. Cationic macromolecules show higher glomerular permeability than anionic macromolecules of similar molecular weight. Larger cationic macromolecules accumulate in the kidney and liver as a result of many factors. The predominant charge of the cell surface is negative, leading to binding of cationic molecules to the cells. Interaction of negatively charged components of the blood with cationic macromolecules leads to embolization of aggregates. Moreover, cationic LPs are excellent as gene-delivery carriers and for liver, lung, and tumor targeting. Cationic LPs may be used as carriers to target cell nuclei. In addition, cationic LPs can cross the blood-brain barrier more easily than anionic LPs. Despite the many advantages of cationic LPs over viral vectors in gene delivery, the transfection efficiency of DNA-cationic LPs (lipoplexes) is too low and they have more toxicity than engineered viral vectors. Cationic LPs have been reported to target tumor vasculature selectively, due to an inherent yet unexplained mechanism.⁴³ Moreover, a wide area of scientific research has focused on formulating cationic LPs to enhance their vascular targeting efficiency and diminish toxicity-related reactions.

Morphology of Liposomes

TEM of GMC-SoS-Fo LPs showed that spherical singlebilayer vesicles (unilamellar LPs) were formed, as shown in Figure 1, and their size concurrent with those obtained with the Zetasizer.

Evaluation of Ultrasound Sensitivity of Sonosensitive Liposomes

The release of GMC from GMC-SoS-Fo LPs was investigated under various periods of ultrasonic irradiation (2 MHz, 1.5 W/cm^2 power intensity, 10 seconds' ultrasonic irradiation with 10-second rests for a total of 0–150 seconds), as shown in Figure 2. GMC release increased as a function of time, reaching 90.7%±3.7% at 130 seconds (10 seconds' ultrasonic irradiation, 10-second pauses). However, an insignificant increase in GMC release was noted with an increase to 150 seconds, (92.4%±3.9%), so the period of ultrasonic irradiation was adjusted to 130 seconds in the following experiments.

To confirm the role of Ce_6 as activator of SoS GMC LPs, GMC-Fo LPs, and SoS LPs with various



Figure I Transmission electron micrography of the GMC LPs when stained with uranyl acetate 10%.

concentrations of Ce₆ (3%, 6%, and 9%) incubated at 37°C for 20 minutes were examined under the specific ultrasonic irradiation condition of 2 MHz, 1.5 W/cm² power intensity, 10 seconds' ultrasonic irradiation and 10-second rests for a total of 0-130 seconds), as shown in Figure 3. Minimal GMC release was recorded for all LPs untreated with ultrasonic irradiation. GMC release from conventional liposomes (GMC LPs) and GMC-Fo LPs was also unchanged under the same condition, as shown in Figure 3. However, GMC release from GMC-SoS-Fo LPs with 3% Ce₆, GMC-SoS-Fo LPs with 6% Ce₆, and GMC-SoS-Fo LPs with 9% Ce6 treated with ultrasonic irradiation was 48.7%±3.9%, 89.7%±4.2%, and 90.1% $\pm 2.7\%$, respectively. The results demonstrated that Ce₆ content was a fundamental factor in releasing GMC from relevant LPs, while increased GMC release resulted from increasing the Ce₆ content to 9% compared to the release from GMC-SoS-Fo LPs with 6% Ce₆ (p<0.05). Ultrasonic irradiation conditions (2 MHz, 1.5 W/cm² power intensity, 10 seconds' ultrasonic irradiation and 10-second rests for a total of 0–130 seconds) may be insufficient to completely activate SoS LPs GMC-SoS-Fo LPs with 9% Ce6. However, 6% Ce₆ was the optimum concentration under the same ultrasonic irradiation conditions to produce sonodynamic activity, resulting in burst release of GMC from GMC-SoS-Fo LPs. These results may be attributed to the ability of the SoS activator (Ce_6) to form pores or



Figure 2 In vitro release of GMC from GMC-SoS-Fo LPs (6% Ce₆) with various ultrasonic irradiation periods in PBS (0.1 M, pH 7.4) at 37° C. Results depicted as means ± SD (n=3).

irreparably damage the LP bilayer.^{44,45} Sonosensitivity of DSPE-containing LPs has been attributed to DSPE possibly forming an inverted hexagonal frame within the liposome bilayer as a response to high temperature or pressure.⁴⁶ As a result of ultrasonic irradiation, DSPE in the lamellar liquid crystalline converted to a hexagonal phase, forming cavities because the polar head groups occupied less space than theapolar moieties.⁴⁷

In Vitro Release of Liposomes

In vitro release of GMC from GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs treated with sonication or not is represented in Figure 4. GMC release from GMC-SoS-Fo LPs showed ultrasounddependence: as GMC-SoS-Fo LPs were exposed to ultrasonic irradiation, they were disrupted, releasing GMC promptly. However, GMC release from GMC-SoS-Fo LPs not treated with ultrasonic irradiation was 9.5%±1.5% after incubation of 50 minutes. Moreover, GMC LPs and GMC-Fo LPs represented ultrasound independence and minimal release change was observed, as shown in Figure 4.

GMC release from GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs incubated for 24 hours in isotonic PBS (pH 7.4, 37°C) was minimal (<23%). However, subjecting GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs to ultrasonicirradiation at 25 hours led to burst release (93.6%±3.9) only with GMC-SoS-Fo LPs, while insignificant release changes were observed with GMC LPs and GMC-Fo LPs, as shown in Figure 5. The results confirmed that release from GMC-SoS-Fo LPs was low until LPs were in contact with ultrasonic irradiation: burst release was achieved, suggesting little drug released in blood until the LPs reached the desired site (ovarian tumors) and were subjected to ultrasound. GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs showed a strong tendency to be stable upon no ultrasonic wave exposure. There were no changes in drug release from GMC LPs or GMC-Fo LPs in the absence or presence of ultrasonic irradiation. These findings could be explained by insensitivity of GMC LPs and GMC-Fo LPs to ultrasonic irradiation. However, GMC-







Figure 4 In vitro release of GMC from GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs untreated or treated with ultrasonic irradiation in isotonic PBS) pH 7.4, 37°C). Results depicted as means ±SD (n=3).



Figure 5 In vitro release of GMC from GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPsuntreated or treated with ultrasonic irradiation after 24 hours' incubation in isotonic PBS (pH 7.4, 37°C). Results depicted as means ± SD (n=3).

SoS-Fo LPs were disrupted, releasing the encapsulated drug once they were stimulated with acoustic energy. The sensitizer Ce_6 was a key component, whereas disruption of the LP membrane may have occurred as a result of vibration of Ce_6 to an extent that was sufficient to destroy the bilayer membrane or form sufficient pores for escaping

the encapsulated drug. SoS LPs for ultrasound-triggered release have been developed.⁴⁸ As ultrasonic irradiation was applied, gas bubbles oscillated and collapsed in media, so introducing high mechanical pressure increases release substantially. This could be explained by the formation and collapse of small gas nuclei in the hydrophobic

region of the lipid bilayer during exposure to ultrasonic irradiation, thereby inducing the formation of transient pores through which drugs are released.⁴⁴

Stability

GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs were physically and chemically stable in the conditions examined (4 C, up to 3 months), as presented in Table 2. EE change forGMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs) was insignificant. Particle size growth, ζ -potential reduction, and PDI values of the LPs showed insignificant changes, proving that monodispersion of LPs was stable. Finally, ultrasonic sensitivity of GMC-SoS-Fo LPs continued without change during storage.

Cellular Uptake of Liposomes on Flow Cytometry

Flow-cytometry analysis was utilized to quantify cellular uptake of GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs with or without ultrasonic irradiation. The cell populations that bound and absorbed released calcein were identified by green fluorescence. Calcein acetoxymethyl molecules have no green fluorescence, but were converted to green fluorescent calceinacetoxymethyl by intracellular metabolism. Cellular uptake of GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs incubated with the cell line and untreated with ultrasonic irradiation was 12.5%±5.7%, 15.85%±3.9%, and 16.12%±4.9%, respectively. Additionally, cellular uptake of GMC LPs and GMC-Fo LPs incubated with the cell line and treated with ultrasonic irradiation (2 MHz, 1.5 W/cm² power intensity, 10 seconds' ultrasonic irradiation, and 10-second rests for a total of 0-130 seconds), was 13.4%±4.5% and 15.45%±5.2%, respectively. However, cellular uptake of SoS targeted LPs (GMC-SoS-Fo LPs) incubated with the cell line treated with ultrasonic irradiation (2 MHz, 1.5 W/cm² power intensity, 10 seconds' ultrasonic irradiation, and 10-second rests for a total of 0-130 seconds), was 81.4%±8.5%, as shown in Figure 6. Compared to the cellular uptake achieved using GMC LPs not treated with ultrasonic irradiation, that achieved using GMC-SoS-Fo LPs treated with ultrasonic irradiation was improved 6.51-fold. These findings can be explained on the basis of intercellular uptake based on the Fo ligand followed by burst release as a response to ultrasonic irradiation. Ce₆ molecules inserted into SoS targeted LPs were vibrated under the effect of ultrasonic irradiation, creating pores or rupturing the nanocarriers. Moreover, the extracellular liberated calcein acetoxymethyl can be easily engulfed by cells, converting it to have greenfluorescence. These results prove that SoS targeted LPs can bind strongly with Fo-expressing cells and release cargo promptly as a result of ultrasonic irradiation.

In Vitro Cytotoxicity Assays

The cytotoxic effects of SoS Fo LPs, free GMC, GMC LPs, GMC-Fo LPs, GMC-SoS Fo LPs (no ultrasonic irradiation), and GMC-SoS Fo LPs (ultrasonic irradiation) against the CAOV3 cells are illustrated in Figure 7. The

Table 2 Particle Size, PDI, Encapsulation Efficiency, and Effect of Ultrasonic Irradiation on Release of GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs) Over 3 Months

	Time (Months)	Particle Size, Mean ± SD (nm)	PDI, Mean ± SD	ζ-Potential, Mean ± SD (mV)	EE, Mean ± SD (%)	Ultrasonic Irradiation Effect on Release Rate
GMC LPs	0	120.3±2.4	0.272±0.005	37.24±0.97	86.3±1.84	No
	1	125.7±1.9	0.282±0.003	33.74±0.84	85.6±2.35	No
	2	127.5±1.1	0.280±0.006	32.51±0.86	84.8±2.01	No
	3	129.1±3.1	0.279±0.004	30.24±1.04	84.1±1.45	No
GMC-Fo LPs	0	119.1±3.5	0.228±0.009	34.52±1.31	82.8±1.12	No
	1	120.8±2.2	0.231±0.004	32.42±1.21	82.0±2.33	No
	2	122.1±2.8	0.229±0.006	31.61±0.82	80.9±1.99	No
	3	124.1±2.4	0.232±0.005	30.37±0.83	80.1±1.54	No
GMC-SoS-Fo LPs	0	109.5±2.9	0.209±0.002	39.70±0.71	84.6±2.04	Burst release promptly
	1	113.1±2.5	0.201±0.006	38.14±0.59	83.9±1.72	Burst release promptly
	2	114.9±2.7	0.202±0.005	36.41±0.93	82.8±2.77	Burst release promptly
	3	7. ±3.3	0.211±0.004	36.01±1.15	81.9±2.45	Burst release promptly

Abbreviation: PDI, polydispersity index.



Figure 6 Flow-cytometry findings for GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs) when incubated with the CAOV3 cell line and treated or untreated with US-IR. Notes: Measurements estimated after 30 minutes' incubation, and are averages of three results ± SD. Abbreviation: US-IR, ultrasonic irradiation.



Figure 7 Cytotoxic effect of GMC. CAOV3 cells were grown and treated with increasing concentrations of GMC in different liposomal formulations. Cell viability was evaluated by MTS reagent(see Methods). Values expressed asmeans ± SEM of at least three different experiments in triplicate.

CAOV3 cell line was chosen for this study because its viability is unaffected by the chemotherapeutic agent cisplatin at concentrations >200 µM and GMC >1,000 nM.⁴⁹ The drug-free carrier system (SoS Fo LPs) alone showed no effect on cell viability, confirming the biocompatibility of SoS Fo LPs. CAOV3 cells were treated with various concentrations of GMC. Free GMC, GMC LPs, GMC-Fo LPs, and GMC-SoS Fo LPs (untreated) reduced cell viability in a dose-dependent manner up to 80 µM. However, GMC-SoS Fo LPs (treated with ultrasonic irradiation) reduced cell viability in a dose-dependent manner up to 40 µM. The highest cellular cytotoxic effect was reported with GMC-SoS Fo LPs (treated with ultrasonic irradiation, 40 µM drug equivalent). Our results showed that GMC-SoS Fo LPs (treated with ultrasonic irradiation, 80 µM drug equivalent) had an insignificant increase in cellular cytotoxic effect compared to GMC-SoS Fo LPs (treated with ultrasonic irradiation, 40 μ M drug equivalent).

Pharmacokinetic and Biodistribution Studies

Pharmacokinetic and biodistribution evaluations of the formulations were performed to examine in vivo ovarian cancer–targeting efficiency. GMC concentration in plasma after intravenous injection in LPs was measured, as presented in Figure 8. Limits of determination detection were 0.1617 and 0.4903 μ g/mL, respectively, calculated using calibration curves as per International Council for Harmonisation guidelines. Accuracy and precision of analytical assays were achieved across their linear range.

GMC plasma concentration reached its peak (24.1 ± 1.2 µg/mL) in five minutes after a single GMC injection. It quickly decreased to be around 8.78% of the peak value within 1 hour, suggesting rapidly eliminationt. The rapid plasma elimination of GMC following intravenous injection of the pure form of the drug could be explained by the rapid metabolism of GMC and conversion into its inactive metabolites. These metabolites are more susceptible to elimination, and this was confirmed by increased concentration of GMC in kidneys, as shown in Figure 8.

GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs showed insignificant differences in plasma concentration. These results confirmed that the Ce_6 component did not interfere with long circulation in LPs (Table 3).

However, pharmacokinetic parameters of free GMC — $t_{\frac{1}{2}}$, AUC_{0-t}, AUC_{0- ∞}, mean residence time, and clearance



Figure 8 Plasma concentration of GMC versus time after IV injection of free drug, GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs in rats (n=3). Results are depicted as means ± SD (n=3).

— were significantly different from GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs (p < 0.05, Table 3).

Selective targeting of drug to tumors was confirmed by the concentration of GMC in tumors following intravenous injection of GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs without ultrasound and GMC-SoS-Fo LPs with ultrasound, as presented in Figure 9. Concentration of GMC in tumors treated with GMC-SoS-Fo LPs with ultrasound was 2.89-fold that of free GMC, while concentration of GMC in tumors treated with GMC-SoS-Fo LPs with ultrasound was 1.98-fold that of GMC LPs, 1.44-fold that of GMC-Fo LPs, and 1.42-foldthat of GMC-SoS-Fo LPs with no ultrasound. The highest concentration of GMC was achieved in tumors treated with GMC-SoS-Fo LPs with ultrasound, leading not only to improved specificity of the drug and killing of cancerous cells but also to decreased side effects. These benefits of using GMC-SoS-Fo LPs with ultrasound encourage further application of SoS delivery systems for treating tumors.

These findings could be explained by the targeting efficiency of GMC-SoS-Fo LPs and their externally triggered rapid release, producing high concentrations within tumor tissue. The dual mechanism of GMC-SoS-Fo LPs may potentially support anticancer efficiency of GMC in vivo. To confirm these results clinically, evaluation of antitumor activity of various LPs was performed in vivo.

In Vivo Antitumor Activity

Repression of tumor growth following intravenous injection of free GMC, GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs with or without ultrasonic irradiation was

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	Free Drug	GMC LPs	GMC-Fo LPs	GMC-SoS-Fo LPs
C _{max} (μg/mL)	25.95	27.75	28.5	27.15
AUC₀₋₊(µg· min/mL)	766.875	3,678.375	3,538.5	3,387.375
AUC₀_∞ (µg· min/mL)	806.27	4,272.91	4,016.32	3,824.11
AUMC₀ _{−t} (µg· min²/mL)	14,388.75	437,186.3	402,480	379,428.75
AUMC₀–∞ (µg· min²/mL)	17,491.93	686,238.61	600,183.72	559,353.74
MRT (minutes)	18.76	118.85	113.74	112.013
K _e (minutes)	0.05330	0.00841	0.00879	0.00893
t _{1/2} (minutes)	11.9805423	113.5290319	98.12962104	94.51163601
CI (mL/µg)	5.22	1.09	1.13	1.18

Table 3 Pharmacokinetic Parameters of GMC After IV Injection in Different Formulations (Free Drug, GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs)

Abbreviations: AUC, area under the plasma concentration-time curve; AUMC, area under the moment plasma concentration-time curve; C_{max} , maximum plasma concentration; MRT, mean residence time; K_{e} , elimination-rate constant; Cl, clearance; t_{ij} , half-life.



Figure 9 Biodistribution of GMC in heart, liver, spleen, lung, and kidney and ovarian cancer 30minutes after IV injection of free GMC, GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs without ultrasonic irradiation and GMC-Fo LPs and GMC-SoS-Fo LPs with ultrasonic irradiation. Results depicted as means ± SD (n=10). *p<0.05.

evaluated, as shown in Figure 10. To determine tumorrepression effects, the ratio of ovary volume at the end of the experiment to ovary volume at the beginning of the experiment was measured. The higher the ovary-growth ratio is, the lower the tumor-repression effect. Ovarygrowth ratio for free GMC, GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs with or without ultrasonic irradiation were 3.46, 2.76, 2.25, 1.75, 1.71, and 1.21,



Figure 10 Antitumor activity of GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs without ultrasonic irradiation and GMC-SoS-Fo LPs with ultrasonic irradiation against isotonic saline and free drug in xenograft ovarian cancer rat models. Results depicted as means \pm SD (n=10). Arrows show the days when treatments were given. *p<0.5.

respectively. As predicted, the tumor-growth ratio forGMC-SoS-Fo LPs with ultrasonic irradiation was lowest — 1.21. This can be attributed to the targeting efficiency of the drug-delivery system, relatively long circulation time of the nanocarriers, and external triggering by ultrasonic irradiation, resulting in rapid release of the drug cargo.

As for the Kaplan–Meier survival curves, the median survival of GMC-SoS-Fo LPs (G5) with ultrasonic irradiation–treated rats was compared to all other groups. Significant improvement in median survival and thus slowdown of tumor recurrence was observed in groups treated with GMC-SoS-Fo LPs (G5) with ultrasonic irradiation compared to untreated and other groups (Figure 11).

Finally, body weight in the groups was measured to evaluate the safety of the dosages. Insignificant bodyweight changes were recorded for groups injected with GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs with or without ultrasonic irradiation, as shown in Figure 12. However, the control group and that intravenously injected with free GMC were characterized by acute body-weight reduction because of cancer cachexia and the toxicity of untargeted dosage forms. Repression of tumor growth inhibited weight loss.

In cancer-suffering living bodies, loss of body weight with specific loss of skeletal muscle and adipose tissue is

recorded, which is called cachexia. This is highly associated with cancers of the pancreas, ovaries, stomach, lungs, and bowels. Weight loss may be due to processes related to metabolic changes mediated by excess release of proinflammatory cytokines and increased activity of the sympathetic nervous system. Both catecholamines and proinflammatory cytokines promote catabolic processes.³⁶ Proinflammatory cytokines, in addition to decreasing the efficacy of growth hormones, work in the central nervous system as mediators of inflammation and act as catabolic factors stimulating proteolytic pathways, leading to muscle and increased adipose-tissue breakdown.⁵⁰ atrophy Proinflammatory cytokines, including IL6, IL10, IL18, and TNFa, stimulate muscle-protein degradation, cause contractile dysfunction, and inhibit myogenesis, in addition to promoting adipose-tissue waste, inhibition of adipocyte differentiation, lipolysis stimulation, and increased apoptosis in adipocytes.51

Finally, these formulations succeeded in managing ovarian cancer because of their accumulation within tumors via passive (GMC LPs) or active (GMC-Fo LPs and GMC-SoS-Fo LPs) targeting. Accumulation of nano-carriers in tumors has been reported as due to the EPR effect,⁵² because of leaky vasculature and lymphatic drainage within cancerous tissue. Physicochemical properties of nanocarriers, such as particle size, charge of surface, and circulation time(longevity), can influence their



Figure 11 Kaplan–Meier survival curves for rats treated intravenously with GMC-free LPs (control group), 4 mg/kg of free GMC solution (G1), GMC LPs (G2), GMC-Fo LPs (G3), GMC-SoS-Fo LPs (G4), and GMC-SoS-Fo LPs (G5) subjected to ultrasonic irradiation. Drug dose administered was 4 mg/kg (n=3 for all groups).



Figure 12 Weight loss recorded for groups treated with control (isotonic saline solution), free GMC, GMC LPs, GMC-Fo LPs, and GMC-SoS-Fo LPs without ultrasonic irradiation and GMC-SoS-Fo LPs with ultrasonic irradiation) in xenograft ovarian cancer rat models. Results depicted as the means ± SD (n=10).

accumulation within tumors. Particle size of nanocarriers is an important parameter affecting the EPR effect of nanocarriers.⁵³ To achieve extravasation of drug in cancerous tissue, the mean size of drug carriers should be about 100 nm. At the same time, EE should be also be sufficient. These technological requirements are important for effective in vivo–delivered drugs.^{54,55}

The negative charge of the luminal surface of blood vessels, due to the presence of carboxylate sugars and the sulfate group, interact with positively charged nanocarriers, preventing them from recirculation.⁵⁶ However, a high positive charge may lead to opsonization and clearance of nanocarriers from the circulation⁵⁷ and may be the reason for insignificant difference in tumor accumulation found between GMT LPs and GMT-Fo LPs.

Longevity can influence both passive and active accumulation of nanocarriers, because of the large number of passages achieved via the target. To prevent the capture and clearance of nanocarriers, steric protection should be achieved by grafting polymers, such as PEG, on the surface of nanocarriers. Steric protection can be explained on the basis of shielding surface charges, improved hydrophilicity, and formation of a polymeric layer covering the nanocarriers' surfaces, leading to repulsion between blood components and nanocarriers.⁵⁸

Active targeting can be achieved via two techniques. The first is decorating the nanocarrier's surface with ligands, such as Fo, that interact with specific receptors. Fo receptors are extensively expressed in tumor tissue. The choice of the targeting ligand represents a significant factor in the engineering of targeted nanocarriers.⁵⁹ The second technique is achieved via formulating stimulus-sensitive nanocarriers that respond to internal stimuli, such as changes in the pathological area or external stimuli ultrasonic irradiation.

Ultrasonic waves are mechanical waves with periodic continuous vibrations at frequencies of 20 kHz or more. They are safe; however, they have exceptional tissue-penetrating ability without major attenuation of energy.⁶⁰ The effect of ultrasonic waves can be localized by irradiating specific areas and choosing an SoS drug-delivery system with cancer affinity.⁶¹ In the present study, the sensitizer was a key component. Distribution and cellular uptake of Ce_6 was the most important factor in its therapeutic action, due to short life and very short diffusion distance of some radical products, such as peroxides, derived from the sensitizer and produced after its cellular uptake.⁶² Li et al reported that Ce₆ enhanced cytotoxicity and apoptotic response as a result of Ce6 accumulation within mitochondria, leading to damage and apoptosis.63 Subsequently, the damaged mitochondria released many chemical factors, eg, cytochrome C. The factors released stimulate activation of the caspase 9 proenzyme to its active form, causing cell apoptosis.63

The present study used ultrasonic irradiation to enhance delivery of GMC to ovarian cancer upon GMC- SoS-Fo LPs. This technique used ultrasonic irradiation to increase the permeabilization of phospholipid membranes, inducing GMC release from nanocarriers at cancerous ovaries and increasing drug uptake into cancer cells.

Conclusion

Targeted SoS LPs (GMC-SoS-Fo LPs) have a spherical shape and are nanosized with optimum ζ -potential. Compared to untargeted LPs (GMC LPs) and targeted nonultrasonic sensitive LPs (GMC-Fo LPs), the targeted SoS LPs succeeded in releasing the drug cargo as a response to ultrasonic irradiation as external stimuli. Moreover, targeted SoS LPs enhanced intracellular uptake, leading to significant inhibition of tumor growth.

The study data showed that GMC-SoS-Fo LPs are a promising anticancer drug- delivery system. To enhance the targeting efficiency of SoS LPs, Fo ligands were inserted into the the prepared LPs (active targeting) and SoS LPs were prepared in the nanosize range (passive targeting).

In future, we need to develop SoS theranostic LPs to explore the usage of these drug-delivery systems not only for oncotherapy but also for diagnosis.

Abbreviations

GMC, gemcitabine; GMC LPs, GMC-loaded LPs; GMC-Fo LPs, GMC-loaded Fo-targeted LPs; GMC-SoS Fo LPs, GMC-loaded Fo-targeted SoS LPs; PC, L- α -phosphatidylcholine; EE, entrapment efficiency); DL, drug loading; PDI, polydispersity index; EPR, enhanced permeability and retention; Ce₆, chlorin e₆.

Disclosure

The authors report no conflicts of interest in this work.

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