Chondroitin sulfate functionalized mesostructured silica nanoparticles as biocompatible carriers for drug delivery

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Abstract: Mesoporous silica nanoparticles (MSNs) have garnered a great deal of attention as potential carriers for therapeutic payloads. Here, we report a pH-responsive drug-carrier based on chondroitin sulfate functionalized mesostructured silica nanoparticles (NMChS-MSNs) ie, the amidation between NMChS macromer and amino group functionalized MSNs. The prepared nanoparticles were characterized using dynamic light scattering, fourier transform infrared spectroscopy and transmission electron microscopy. The resultant NMChS-MSNs were uniform spherical nanoparticles with a mean diameter of approximately 74 nm. Due to the covalent graft of hydrophilic and pH responsive NMChS, the NMChS-MSNs could be well dispersed in aqueous solution, which is favorable to being utilized as drug carriers to construct a pH-responsive controlled drug delivery system. Doxorubicin hydrochloride (DOX), a well-known anticancer drug, could be effectively loaded into the channels of NMChS-MSNs through electrostatic interactions between drug and matrix. The drug release rate of DOX@NMChS-MSNs was pH dependent and increased with the decrease of pH. The in vitro cytotoxicity test indicated that NMChS-MSNs were highly biocompatible and suitable to use as drug carriers. Our results imply that chondroitin sulfate functionalized nanoparticles are promising platforms to construct the pH-responsive controlled drug delivery systems for cancer therapy.

Keywords: mesoporous silica nanoparticle, pH sensitive, chondroitin sulfate, drug delivery

Introduction

Nanoscale drug delivery systems (DDS) have achieved advantages by overcoming the challenges of common cancer treatments,¹,² they can perform as a vehicle protecting the therapeutic agent from the biological milieu and improve cellular uptake and accumulation inside tumor sites. Up to date, many materials, eg, liposomes, block copolymers, dendrimers, and various inorganic nanomaterials, have been utilized as drug carriers in DDS.³–⁶ Among them, mesoporous silica nanoparticles (MSNs) have attracted more and more attention as a promising component,⁷–¹³ they are excellent candidates for many biomedical applications owing to their high specific surface area, large pore volume, tunable pore structures and well-defined surface property for modification.

Despite their promise, however, recent reports highlight the potential toxicity of unmodified MSNs due to the interactions of surface silanols with cellular membranes.¹⁴–¹⁶ This toxicity can be reduced by coating the nanoparticles with polymer shells. The polymer shells can provide colloidal stability, handle for chemoligation (targeting moieties) and improve the blood circulation lifetimes, which are crucial for efficient in vivo drug delivery. Various polymers have been reported as coating shells.
of MSNs, such as chitosan,\(^{17,18}\) poly(ethylene glycol) (PEG),\(^{19}\) poly(acrylic acid) (PAA),\(^{20}\) poly(4-vinyl pyridine) (PVP),\(^{21}\) poly(L-lysine) (PLL),\(^{22}\) poly(N-isopropylacrylamide),\(^{23}\) poly (lactic acid) (PLA)\(^{24}\) either in their native forms or as conjugates. However, the applicability of these functionalized MSNs is largely restricted to their compositions and the coating materials, whereby the loading efficiency is usually not high enough for a practical usage and the polymer shell also limits both drug loading and release from MSNs. Therefore, efforts should still be made to develop safe and facile approaches for the fabrication of functionalized MSNs with a variety of potential biomolecular payloads.

Chondroitin sulfate (ChS) is an important structural component in connective tissues and cartilage. It is a copolymer of D-glucuronic acid and sulfated N-acetyl-D-galactosamine in C4 or C6 and belongs to the group of glycosaminoglycans (GAGs), which are primarily located on the surface of cells or in the extracellular matrix.\(^{25,26}\) ChS can be degraded by anaerobic bacteria, namely Bacteroides thetaiotaomicron and Bacteroides ovatus, which are residents of the large intestine. This character suggests that ChS is potentially a good candidate for usage as a drug carrier.\(^{27-30}\) Recently, we used a method of interfacial polymerization in emulsion to fabricate chondroitin sulfate-based nanocapsules for loading of indomethacin.\(^{31}\) The shell of nanocapsules consisted of the cross-linked ChS chains, and showed a much higher drug loading capacity. Our recent study also showed that the nanogels fabricated of chondroitin sulfate could exhibit promising pH-responsive properties.\(^{32}\) Therefore, it can be imagined that the combination of MSNs and ChS polymer as carriers could realize the high loading capacity and pH-triggered controlled release of drug simultaneously, which would be a potential and promising strategy to design a stimuli-responsive codelivery system of drug or gene.

Herein, we report a new strategy to prepare a pH-responsive NMCChS functionalized MSNs (NMCChS-MSNs) by the amidation reaction of NMCChS macromers with amino groups modified MSNs. (Scheme 1). We first synthesized a macromer named as O-maleyl chondroitin sulfate (NMCChS) (Scheme S1). To obtain the animated MSNs (MSNs-NH\(_2\)), we used bifunctional (3-aminopropyl) triethoxysilane (APTES), and then MSNs-NH\(_2\) were activated by 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide (EDC), then followed by the addition of positively charged NMCChS to form covalent bonding between primary amine of MSNs-NH\(_2\) and carbonyl groups of the NMCChS macromers. An acrylamide group in NMCChS macromer made the polymer chains around the silica surface to be cross-linked possible. Subsequently, a covalently cross-linked NMCChS shell was synthesized at room temperature by addition of K\(_2\)S\(_2\)O\(_7\)/NaHSO\(_3\), and then NMCChS-MSNs could be obtained. Doxorubicin hydrochloride (DOX) was chosen as a model drug in this study to assess the drug loading and releasing behaviors of NMCChS-MSNs. In vitro cellular cytotoxicity test was performed to evaluate the biocompatibility of NMCChS-MSNs, and the cytotoxic effect of drug-loaded DOX@ NMCChS-MSNs to HeLa cells was also investigated.

**Materials and methods**

**Materials**

Chondroitin sulfate (ChS, oral grade, Mn = 50000 by GPC) powder was supplied by Shanxi Scipharm Biotechnology Co, Ltd (Xi’an, China). The analytical grade reagents of maleic anhydride (MA), NaHSO\(_3\), (3-aminopropyl) triethoxysilane (APTES), tetraethoxysilane (TEOS), cetyltrimethylammonium bromide (CTAB), potassium persulfate (KPS) and NaOH were obtained from Sinopharm Chemical Reagents Co, Ltd (Shanghai, China). KPS was re-crystallized from

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**Scheme 1** Schematic procedure for preparation of O-maleyl chondroitin sulfate functionalized mesoporous silica nanoparticles (NMCChS-MSNs) for drug loading. *Abbreviations: APTES, (3-aminopropyl) triethoxysilane; EDC, 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride; NMCChS, O-maleyl chondroitin sulfate; KPS, potassium persulfate; NaHSO\(_3\), sodium bisulfite; DOX, doxorubicin.*
water and used after vacuum drying. Doxorubicin (DOX) in the form of hydrochloride salt was obtained from Beijing Huafeng United Technology Company (Beijing, China). MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) assay and other biological reagents were purchased from Invitrogen Corp. (USA). All other reagents were of analytical grade and used without further purification.

Methods

Synthesis of NMChS

O-maleyl chondroitin sulfate (NMChS) was prepared as similar to the method of Zhu A. (Scheme S1). In a typical synthesis, chondroitin sulfate powder (1.0 g) was first dissolved in 100 mL aqueous solution and transferred to a three necked flask. The maleic anhydride (MA) were dissolved in 5 mL of acetone, and then the maleic anhydride-acetone solution was added to the reaction flask drop wise over 20 min at 65°C. After that, the reaction was allowed to stand at 70°C for another 12 h. Later, the reaction mixture was cooled to room temperature and precipitated by adding excess amount of acetone several times. After washing by acetone, the precipitate was collected and dried at 45°C in vacuum for 24 h.

Synthesis of MSNs

MSNs were synthesized via a sol-gel chemistry using cetyltrimethylammonium bromide (CTAB) as templates. In a flask, 0.28 g of NaOH and 1.0 g of CTAB in sequence were completely dissolved into 480 mL of deionized water under vigorous stirring at 80°C. After the solution became clear, 5 mL of TEOS was added dropwise with the whole dropping time of about 15 min. The vigorous stirring was continued for 20 h, and then milk-white as-synthesized materials were collected by centrifugation. In order to remove the surfactant, the as-synthesized materials were refluxed in a mixed solution of 500 mL ethanol and 5 mL hydrochloric acid at 78°C for 12 h, then centrifuged and washed for several times with ethanol. The same reflux operation was repeated for three times. The final products were dried at 80°C in vacuum for 24 h.

Synthesis of NMChS-MSNs

Firstly, 0.2 g of MSNs was added to 40 mL of ethanol containing 0.5 mL of APTES. After being stirred at room temperature for 24 h, the mixture was extensively washed with ethanol and dried at 80°C to obtain the aminated MSNs (MSNs-NH$_2$). Next, to activate the amine groups, EDC (0.1 mol·L$^{-1}$) was added to the MSNs-NH$_2$ (5.0 mg·mL$^{-1}$ in water, 20 mL), the above mixture was kept for 1 h, and then NMChS solution (2.5 mg·mL$^{-1}$) was added. After 8 h of stirring at room temperature, the mixture was centrifuged, washed twice with water to remove excess EDC and un-reacted NMChS, and re-dispersed in water. Subsequently, the polymerization cross linking reaction was initiated by addition of 25 µL of KPS/NaHSO$_3$ solution (1.0 mg/1.0 mg) under a nitrogen atmosphere at 40°C, and performed for 6 h. After that, the NMChS functionalized MSNs were collected by centrifugation for further use.

Drug adsorption into MSNs and NMChS-MSNs

In a typical MSNs drug loading process: 12.5 milligrams of MSNs was mixed with 5 mL of DOX solution in PBS. After stirring for 24 h under dark conditions, the DOX-loaded MSNs (DOX@MSNs) were centrifuged and washed with PBS. In a typical NMChS-MSNs drug loading process (Scheme 1): 12.5 milligrams of NMChS-MSNs before cross-linking were mixed with 5 mL of DOX solution in PBS, after stirring for 24 h under dark conditions, the DOX-loaded nanoparticles were centrifuged, washed with PBS and re-dispersed in 5 mL PBS by addition of KPS/NaHSO$_3$ solution for cross linking reaction. Finally, the DOX@NMChS-MSNs were collected by centrifugation. The obtained DOX@MSNs or DOX@NMChS-MSNs were used for subsequent tests of in vitro DOX release and cytotoxicities against HeLa cells. To evaluate the DOX-loading efficiency, the supernatant and washed solutions were collected and the residual DOX content was measured by using UV-vis measurement at a wavelength of 490 nm. The loading content (LC%) and loading efficiency (LE%) of DOX can be calculated as follows:

\[
\text{Drug loading content} (\%) = \frac{\text{weight of drug in MSNs}}{\text{weight of MSNs}} \times 100\% \quad (1)
\]

\[
\text{Drug loading efficiency} (\%) = \frac{\text{weight of drug in MSNs}}{\text{weight of feed drug}} \times 100\% \quad (2)
\]

In vitro release of drug

The in vitro simulated release of DOX was executed in pH 5.0, 6.8 and 7.4 buffer solutions, respectively. The above-prepared DOX@MSNs or DOX@NMChS-MSNs (2.5 mg·L$^{-1}$, 2.5 mL) was immersed in 20 mL PBS solutions at 37°C and shaken at 100 rpm. At certain time intervals, 2 mL of PBS was taken out by centrifugation to test the concentration of the released DOX and 2 mL of fresh PBS was added to the tube containing the DOX@MSNs.
or DOX@NMChS-MSNs. The aliquot was centrifuged to obtain a clear supernatant and analyzed using UV-visible spectroscopy at 490 nm.

**In vitro cytotoxicity**

The cytotoxicity of NMChS-MSNs or DOX@NMChS-MSNs against Hela cells was evaluated by MTT assay. Hela cells were seeded in 96-well plates at 10 000 cells per well in 100 µL of DMEM medium and incubated at 37°C in 5% CO₂ atmosphere for 24 h. The culture medium was replaced with 100 µL of fresh medium containing DOX@NMChS-MSNs or free DOX at pH 7.4. After 24 h treatment, the medium was replaced by fresh DMEM at pH 7.4 and followed by addition of 25 µL of MTT stock solution (5 mg · mL⁻¹ in PBS). After incubation for an additional 2 h, 100 µL of the extraction buffer (20% SDS in 50% DMF, pH 4.7) was added to the wells and incubated for 4 h. The absorbance of the solution was measured at 570 nm using a Bio-Rad 680 microplate reader and cell viability was normalized to that of Hela cells cultured with blank culture medium. The results were compared with those of the control wells to determine relative cell viability.

**Characterization**

Transmission electron microscopy (TEM) images were obtained on a TECNAI-12 transmission electron microscope (Philips, Netherland), and samples for TEM measurements were made by casting one drop of the sample’s ethanol solution on copper grids. Field emission scanning electron microscopy (FE-SEM) observation was carried out on an Hitachi S4800 electron microscope (Hitachi, Japan). Powder X-ray diffraction (XRD) patterns were collected for the powdered samples using an AXS D8 diffractometer (Bruker, Germany). The particle size was characterized by photon correlation spectroscopy (Zetasizer Nano ZS-90, Malvern Instruments, UK). Raw data were subsequently correlated to mean hydrodynamic size by cumulants analysis (Z-average mean). The Zeta potentials of all nanoparticles were analyzed by laser doppler anemometry (Zetasizer Nano ZS-90, Malvern Instruments, UK). All samples were run in triplicate, and the mean averages and standard deviations were calculated. Fourier transform infrared (FT-IR) spectra were recorded on a Tensor 27 spectrophotometer (Bruker, Germany). The UV-vis spectra were recorded by using a UV-2550 spectrophotometer (Shimadzu, Japan) in the wavelength range of 200–600 nm. ¹H NMR spectra were recorded on a NMR spectrometer (ANAVCE DMX600, Bruker, Germany) with D₂O as solvent, working at 600 MHz.

**Results and discussion**

**Structure of NMChS macromer**

Structure changes of chondroitin sulfate after maleoyl-acylation were first confirmed by FT-IR spectra (Figure S1). The representative absorption peaks for nascent ChS (Figure S1A) can be adapted from the publication.² OH and NH stretching (3426 cm⁻¹), CH₂ stretching (2926 cm⁻¹), amide I (1634 cm⁻¹), COO⁻ vₛ stretching (1560 cm⁻¹), COO⁻ v₁ stretching (1412 cm⁻¹), SO₄²⁻ related modes (1248 cm⁻¹), C–O stretching (1088 cm⁻¹ and 1068 cm⁻¹). In the spectrum of maleic anhydride (Figure S1B), there are representative absorption peaks for anhydrides at 1849 and 1778 cm⁻¹. Accompanying the disappearances of the peaks for anhydrides, we find a new absorption peak at 1738 cm⁻¹ in the spectra of NMChS (Figure S1C), which is due to the characteristic ester absorption band. Meanwhile, significant enhanced absorption peaks at 1560 cm⁻¹ (COO⁻ v₃ stretching) and at 1068 cm⁻¹ (C–O stretching) are obvious. These changes shown in FT-IR spectra indicate that maleoyl-acylation has successfully bonded with the ChS molecule.

The evidence of MA substitution on ChS was also observed by ¹H-NMR as shown in Figure S2. A peak at 6.23 ppm is due to the two protons attached to the double bond (CH=CH), which is not present in the virgin chondroitin sulfate molecule. So, the results of ¹H NMR also illuminate that the maleoyl-acylation has successfully bonded with the ChS molecule. Furthermore, according to the N/C weight ratios of ChS (0.121) and NMChS (0.075) detected by elemental analysis, the MA grafting degree is calculated as 62%. After completed the synthesis, the acrylamide group is brought to the NMChS macromer, which makes the polymer chains around the silica surface to be cross-linked possible.

**The morphology and mesostructure of MSNs and NMChS-MSNs**

The schematic preparation process of NMChS-MSNs is illustrated in Scheme 1. Firstly, MSNs were synthesized via the sol-gel chemistry using CTAB as templates. Subsequently, MSNs were functionalized with amino groups by a reaction with APTES, which would afford targets to the consequent modification with NMChS. Then, the aminated MSNs were activated by EDC, and followed by the addition of negatively charged NMChS to form covalent bonding between primary amine and carboxyl groups in NMChS. Finally, the polymerization-cross linking was initiated by the addition of KPS/NaHSO₃ and the cross-linked chondroitin sulfate functionalized mesostructured silica nanoparticles could be
obtained. This synthetic strategy provides a covalent polymer shell without the use of catalysts and surfactants and requires mild conditions compatible with a variety of potential biomolecular payloads. According to our previous work, we hypothesize that NMChS wrapping around MSNs will act as pH responsive shell ensuring release of guest molecules in acidic pH conditions, and this pH responsiveness is believed due to the differences in protonation state of chondroitin sulfate under different pH conditions.

As shown in transmission electron microscopy (TEM) images (Figure 1A), the prepared MSNs are uniform spherical nanoparticles with a mean diameter of approximately 60 nm. FE-SEM images (Figure S3) confirm that MSNs are uniform spheres, which are suitable for further modification. The ordered mesoporous structure of the nanoparticles is further confirmed by a small angle XRD measurement (Figure 1B). Three well-resolved diffraction peaks, assigned as (100), (110) and (200) planes respectively, are clearly observed in the XRD pattern of MSNs, which is consistent with the characteristic diffraction patterns of MCM-41 type MSNs.

After the MSNs were functionalized by NMChS (Figure 2) indicate that the synthesis yielded individually encapsulated MSNs displaying a thin polymeric shell, which is absent on uncoated nanoparticles, and an about 10 nm polymer coating can be observed around the silica particle after polymerization. TEM measurements indicate that the diameters of the uncoated and polymer-coated MSNs are 60 ± 8 and 74 ± 16 nm, respectively. Particle size distribution of MSNs in PBS solution measured by dynamic light scattering (DLS) is relatively broad (mean dimmer = 212.6 ± 13.9, PDI = 0.684) (Figure 3A). The particle size is larger than that of the TEM observation, which may have resulted from the aggregation of MSNs in PBS solution. Compared with the size distribution of MSNs, the NMChS-MSNs have a relatively narrow particle size distribution (mean dimmer = 255.4 ± 9.1 PDI = 0.245) (Figure 3B). This suggests that the NMChS modification improves the dispersity of MSNs in PBS solution. After MSNs and NMChS-MSNs solutions in PBS are statically placed for 12 h, almost all MSNs are located in the bottom, while most of NMChS-MSNs still suspend in PBS solution (Figure 3C and D). Therefore, the DLS and TEM data indicate that this polymer coating procedure can avoid agglomeration of the nanoparticles into the larger (micrometer-scale) aggregates, which is a critical factor in biomedicine applications.
Successful coating of NMChS onto MSNs was also detected by zeta potential measurements. Figure 4 shows zeta potentials of MSNs, MSNs-NH$_2$ and NMChS-MSNs in PBS solution, respectively. It can be found that zeta potentials are changed after the NMChS modification. The zeta potential of MSNs is $-11.9 \pm 3.5$ mV, while it is $8.6 \pm 4.6$ mV for MSNs-NH$_2$ and $-35.1 \pm 5.8$ mV for NMChS-MSNs. This suggests that NMChS have bounded to MSNs, and change the surface potential of MSNs. Such high zeta potential value of in PBS solution also explains that particles are very well dispersed in the solution and do not aggregate so much.

Here, FT-IR spectra were used to further confirm the NMChS modification. The FT-IR spectra for coated and uncoated MSNs display the absorption band at around 1071 cm$^{-1}$, which is assigned to Si–O bond in silica (Figure 5). The peaks at 1648, 796, 967, 1087 and 1237 cm$^{-1}$ in all curves are attributed to the stretching vibration of O–H, the flexible vibrations of Si–O, the stretching vibration of Si–OH and Si–O–Si, respectively. NMChS-MSNs show the adsorption bands of native NMChS at about 1416 cm$^{-1}$. That is attributed to the COO$^-$ symmetry stretching, which is not absent in MSNs FI-IR spectra. Meanwhile, a significant enhanced absorption peak at 1634 cm$^{-1}$ (amide I) is obvious. All the above results indicate the presence of NMChS in modified MSNs, and the covalently cross-linked polymer shell can improve the stability of MSNs.

The physicochemical properties of NMChS-MSNs were also tested at different pH values and temperatures. When the NMChS-MSNs are dispersed in buffer solution with various pH values, the diameter and zeta potential of the NMChS-MSNs are increased with an increase of the pH value (Figure 6A). At pH = 1.0 the diameter of...
the NMChS-MSNs is 164 ± 7.1 nm and the zeta potential is −18.3 ± 3.4 mV. When the pH value is changed to 7.4, the diameter of the NMChS-MSNs increases to 255± nm and the zeta potential increases to −35.9 ± 2.7 mV. When the pH value of solution is lower, the carboxylic groups (−COOH) are located on the surface of NMChS-MSNs. The network among molecular chains exhibits intense structure and shape under strong acidic condition and shows the smaller diameter and the lower zeta potential. However, with increasing the media pH values, carboxylic groups are dissociated into the ionized form (−COO−). Static repulsion increases on account of the dissociation of the carboxylic acid groups under alkaline conditions, leading to the swelling of the NMChS cross-linked layer and increase in size. Likewise, no significantly further condensation occurred below pH < 3 because of complete protonation of the carboxyl groups. These results indicate that the NMChS-MSNs can exhibit promising pH-responsive properties.

The effect of the temperature on the NMChS-MSNs stability was evaluated in a range from 15°C to 50°C. For this study, a solution of NMChS-MSNs was placed in the DLS cell, and the temperature in the chamber was scanned with increments of 5°C and 20 min stabilization time at each temperature. Afterward, the cell was left to cool to 15°C, and the size distribution was again recorded. A plot of average particle size versus temperature is shown in Figure 6B. A modest decrease in the average particle diameter with increasing the temperature is observed. These results show that a rise in temperature does not destroy the polymer network between segments consisting of NMChS monomer units, and the NMChS polymer shells keep their structure well and do not collapse even at 50°C. This result indicates that NMChS-MSNs are stable in the temperature range of 15°C–50°C.

**Calculation of loading capacity and release behavior**

We chose doxorubicin (DOX) as a model drug due to its well-characterized spectral characteristics and its use in chemotherapy. As it is known, the pKa of DOX is 8.6. At pH 7.4, the positively charged DOX will bind with the negatively charged NMChS to form the DOX@MSNs-NMChS complex by electrostatic interaction, which can enhance the loading of drug into the MSNs pores. We first investigated the drug-loading capacity of the polymer-coated MSNs by comparing the total amount of drug loaded before and after polymer coating. When we fixed the concentration of DOX to be 0.25 mg⋅mL−1, the loading of DOX in polymer-coated MSNs is only slightly lower (about 68% of total DOX added) compared with the uncoated MSNs (about −74% of total DOX added). DOX loading contents in MSNs and NMChS-MSNs are 68.1 and 74.0 µg DOX per 1.0 mg of MSNs, respectively. This suggests that the polymer shell does not reduce the drug loading capacity of MSNs as drastically as other reported polymer shell MSNs systems, which is an additional advantage of our technique over previously reported coating methods.35,36

We then determined the loading content (LC%) and the loading efficiency (LE%) of DOX at different weight ratios.

**Table 1** DOX loading content and loading efficiency of NMChS-MSNs

<table>
<thead>
<tr>
<th>DOX (mg⋅mL−1)</th>
<th>Loading content (%)</th>
<th>Loading efficiency (%)</th>
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<tr>
<td>0.25</td>
<td>6.8</td>
<td>68.1</td>
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<tr>
<td>0.5</td>
<td>11.9</td>
<td>59.5</td>
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<tr>
<td>1.0</td>
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**Note:** The concentration of NMChS-MSNs is 2.5 mg⋅mL−1.

**Abbreviations:** DOX, doxorubicin; MNChS-MSNs, O-maleyl chondroitin sulfate functionalized mesoporous silica nanoparticles.
of DOX/NMChS-MSNs. The results are listed in Table 1. The LC% increases with the DOX concentration in the PBS solution due to the increased concentration gradient. The maximum LC% of 14.1% is reached at the DOX concentration of 1.0 mg·mL⁻¹. The value of LE% decreases with the increasing of DOX concentration presumably, because the maximum absorption of the drug is reached. The maximum LE% is about 68.1%. These results indicate that NMChS-MSNs possess high drug loading efficiency to DOX.

Besides drug loading, drug release is another important parameter for drug efficacy. The release profiles of DOX@MSNs and DOX@NMChS-MSNs were performed at pH values of 7.4 (blood circulation), 6.8 (tumor extracellular) and 5.5 (endosomes) for 24 h at 37°C. Figure 7A demonstrates that the DOX release rate is obviously pH dependent and increases with the decrease of pH. At pH 7.4, the release amount is quite low and only approximately 16% is released in 24 h. At pH 6.8, the release profile curve levels off after 12 h and the release amount is 37% in 24 h. Intriguingly note that when pH decreases to 5.5, a faster release behavior is obtained and the release amount reaches 61% in 24 h, which is almost four times of that at pH 7.4. This is because the drug release from the NMChS-MSNs is mainly determined by the effect of electrostatic interaction between DOX and polymer nanoshell. The pKa of DOX is 8.3, so it is positively charged at the selected three pH values. On the whole, the higher the pH of the medium is, the more negative charges are on the surface of the NMChS-MSNs, the stronger attraction exists between the positively charged DOX and the negatively charged carriers, so the more difficult for the loaded DOX to be released out of the NMChS functionalized MSNs.

As a comparison, the drug release experiments of DOX@MSN were also carried out. Figure 7B shows the drug release of DOX@MSNs at pH values of 7.4, 6.8 and 5.5. Obviously, the differences in release amount among different pH are not as significant as that of DOX@NMChS-MSNs with the released drug amount of 51%, 54% and 58% in corresponding pH values after 24 h. Therefore, DOX@NMChS-MSNs display a more pronounced pH dependent property than DOX@MSNs, and more drug molecules could be released for DOX@NMChS-MSNs system at lower pH solution. Considering the fact that the tumor tissues are more acidic than the normal tissues, the prepared DOX@NMChS-MSNs would be able to minimize the side effect of DOX. Combined with the specific drug release behavior and simultaneous high loading capacity and encapsulation efficiency for DOX, the NMChS-MSNs are highly expected to be used in cancer treatment without frequent interval medication administrations.

**In vitro cell assay**

The results of DOX loading and releasing studies motivated us to further investigate the in vitro cellular cytotoxicity. The in vitro cytotoxicity of DOX@NMChS-MSNs to HeLa cells was investigated by MTT assay. When the pure NMChS-MSNs are 0.5, 5, 10, 100 µg·mL⁻¹, the cell viabilities are 98% ± 5%, 95% ± 5%, 94% ± 12%, 92% ± 9% for 24 h, respectively. As shown in Figure 8, the blank carrier NMChS-MSNs show no cytotoxicity on the HeLa cells even at 100 µg·mL⁻¹ after incubation for 24 h, suggesting that the NMChS-MSNs are bio compatible. Conversely, when the HeLa cells are treated with either the suspension of DOX-loaded NMChS-MSNs or a solution of pure DOX at high DOX concentration, significant decrease of the cancer
cell viability is presented. DOX@NMChS-MSNs exhibit higher inhibition activity after 24 h culture in comparison with DOX in free form. The DOX dose required for 50% cellular growth inhibition (IC\textsubscript{50}) is only 5.2 µg·mL\textsuperscript{-1} with DOX-loaded nanogels, which is 7.1 µg·mL\textsuperscript{-1} of that required for treatment with DOX in free form. Enhanced inhibition of DOX-loaded nanogels to tumor cell growth would be advantageous in lowering the dose of DOX in application. This result indicates that the NNMChS-MSNs may be a good carrier for drug delivery to the cell.

**Conclusion**

In conclusion, we presented a facile and effective method to coat chondroitin sulfate-based macromer onto aminated mesoporous silica nanoparticles to generate pH responsive nanocarriers. The loading and releasing behaviors of DOX were investigated. NMChS-MSNs exhibit a high drug loading efficiency, due to the strong electrostatic interaction between NMChS and DOX. Detailed characterization studies show that chondroitin sulfate is effectively anchored on the surface of the silica preventing release of DOX under undesirable conditions. It is apparent from release studies that DOX releases under mild acidic conditions due to the decrease of the electrostatic interactions between drug and matrix. The in vitro cellular cytotoxicity test demonstrates that the NMChS-MSNs are highly biocompatible and suitable to utilize as drug carriers. Furthermore, DOX@NMChS-MSNs show a more efficient cytotoxicity than free DOX to HeLa cells. The unique polymer coated MSN enables it not only to be a pH-responsive nanocarrier, but also to possess tailored release profile. Therefore, the polymer modified MSNs system is a promising strategy toward their potential application as drug delivery vehicles.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**


Supplementary figures

Scheme S1 Synthesis pathway of macromer NMChS.
Abbreviation: NMChS, O-maleyl chondroitin sulfate.

Figure S1 FT-IR spectra of ChS, maleic anhydride and macromer NMChS. (A) rare ChS; (B) maleic anhydride and (C) macromer NMChS.
Abbreviations: FT-IR, fourier transform infrared; NMChS, O-maleyl chondroitin sulfate; ChS, chondroitin sulfate.
Figure S2 ¹H NMR spectra of the rare ChS (A) and synthesized NMChS (B).
Abbreviations: ¹H NMR, nuclear magnetic resonance spectroscopy; NMChS, O-maleyl chondroitin sulfate; ChS, chondroitin sulfate.

Figure S3 FESEM image of MSNs. (B) is higher magnification than (A).
Abbreviations: FESEM, field emission scanning electron microscopy; MSN, mesoporous silica nanoparticles.
Chondroitin sulfate–functionalized nanoparticles for drug delivery