Selective inhibition of liver cancer growth realized by the intrinsic toxicity of a quantum dot–lipid complex

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Abstract: Using the intrinsic toxicity of nanomaterials for anticancer therapy is an emerging concept. In this work, we discovered that CdTe/CdS quantum dots, when coated with lipids (QD-LC) instead of popular liposomes, polymers, or dendrimers, demonstrated extraordinarily high specificity for cancer cells, which was due to the difference in the macropinocytosis uptake pathways of QD-LC between the cancer cells and the normal cells. QD-LC-induced HepG2 cell apoptosis was concomitant with the activation of the JNK/caspase-3 signaling pathway. Moreover, QD-LC treatment resulted in a delay in the latent period for microtumor formation of mouse hepatocarcinoma H22 cells and inhibited tumor growth, with a reduction of 53.2% in tumor volume without toxicity in major organs after intratumoral administrations to tumor-bearing mice. Our results demonstrate that QD-LC could be a very promising theranostic agent against liver cancer.

Keywords: CdTe/CdS quantum dot–lipid complex, intrinsic nanotoxicity, selectivity, liver cancer therapy, macropinocytosis

Introduction

Recent advances in nanomedicine have brought new strategies for cancer diagnosis and treatment. Inorganic nanoparticles, such as gold, quantum dots (QDs), iron oxide, silica, and rare earth particles, are regarded as possible theranostic agents due to their unique material- and size-dependent physicochemical properties, such as an enhanced permeability and retention effect, photothermal effect, and optical tracking.1–3 However, the application of these materials, especially for biomedical purposes, is still limited by the potential toxicity of nanomaterials.4,5 Although the field of nanotoxicology is still developing, several common mechanisms have been found, and, in general, reasonable control over the extent of nanotoxicity is possible.6–8 Therefore, nanomaterial-based therapy utilizing the intrinsic toxicity of nanomaterials has been explored, and it was found that the uptake of some nanomaterials can be controlled to induce cellular autophagy and/or cell death in cancer cells.9–11 Several recent studies indicated great potential for degrading ZnO or silver nanoparticles as novel tools in cancer therapy.11–14 Sasidharan et al showed that ZnO nanoparticles degraded much faster in the low local pH microenvironment typically associated with tumors, thereby selectively destroying cancer cells.15 To date, the intrinsic toxicity of a wide variety of inorganic materials and the selectivity thereof toward cancer cells make this a facile and universal methodology for cancer therapy.

QDs, especially cadmium-based QDs like CdTe, CdTe/CdS, and CdSe/ZnS QDs, have been studied extensively in the past 2 decades. These compounds display unique...
optical properties, including a broad absorption with narrow photoluminescence spectra, high quantum yields (QYS), low photo bleaching, and size-dependent emission-wavelength tunability.16,17 These properties made QDs an appealing candidate for tumor-labeling probes, with significant advantages over conventional fluorescent dyes.18–20 However, safety concerns about the systematic toxicity of cadmium-based QDs has limited their application in biological research as well as their effectiveness as an imaging agent for cancer diagnosis; attempts have been made to derive effective means by which to attenuate their cellular toxicity.21–23 Functionalized QDs with polymers or dendrimers have been used successfully to improve biocompatibility and reduce cytotoxicity in normal tissues.24–26 Alternatively, for QDs modified with various materials, such as an antibody, aptamer, arginine-glycine-aspartic acid, or folic acid, to increase their targeting ability and biological compatibility, these modifications also offer the possibility of QD-based cancer imaging.27–30 The determinative factor of QD toxicity is the dose that refers to the intracellular levels of QD rather than its concentration in the extracellular fluids.31 Finding an appropriate method that facilitates QD uptake only in cancer cells, but not in normal cells, might selectively kill tumor cells without causing damage to normal tissues. Therefore, the aims of this study were to coat lipids on the CdTe/CdS QDs to form a complex labeled as QD-LC and to achieve the selective treatment of liver cancer in vivo and in vitro. This complex demonstrated striking susceptibility to cancer. We have executed in vitro and in vivo studies on the mechanism underlying the phenomenon. In animal experiments, QD-LC treatment resulted in a delay in the latent period for microtumor formation in mouse hepatocarcinoma H22 and inhibited tumor growth without toxicity in major organs. Our findings may enrich the nanocomplex spectrum of which nanotoxicity can be used for cancer therapy.

**Materials and methods**

**Reagents**

1-Ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (99%), sulfo-N-hydroxysulfo-sucinimidimer (98.5%), (3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide) (MTT), NaNO₃, amiloride, simvastatin, mycostatin, sucrose, and chlorpromazine were purchased from Sigma-Aldrich Co. (St Louis, MO, USA). Diagnostic kits for aspartate aminotransferase (AST), alanine aminotransferase (ALT), blood urea nitrogen (BUN), and creatinine (CRE) were purchased from the Nanjing Jiancheng Bioengineering Institute (Nanjing, People’s Republic of China). RPMI 1640 medium, fetal bovine serum, penicillin, streptomycin, and bovine serum albumin (BSA) protein assay kit were purchased from the Beyotime Institute of Biotechnology (Haimen, Jiangsu, People’s Republic of China). FuGENE® HD transfection reagents were purchased from Hoffman-La Roche Applied Science (Basel, Switzerland). The primary antibodies for JNK, p-JNK, Bax, cytochrome c, caspase-9, caspase-3, and GAPDH were obtained from Santa Cruz Biotechnology Inc. (Dallas, TX, USA). All the reagents were used without further purification.

**Synthesis and characterization of water-soluble core/shell CdTe/CdS QDs and CdTe/CdS QD-LC**

CdTe/CdS core/shell QDs were prepared according to our previous publications.20,32 The tailor-designed CdTe QDs (QY =27%) were capped with three monolayers of CdS shell to obtain the yellow emitting CdTe/CdS core/shell QDs with a QY of 69%. For the preparation of the QD-LC, 500 μL of QDs (10 μM) was mixed with 1 μL of FuGENE® HD transfection reagents in phosphate-buffered saline (PBS). The mixture was sonicated for 10 minutes in a bath sonicator at 37°C and then centrifuged in Microcon Centrifugal Filter Devices (50,000 nominal molecular weight limit; EMD Millipore, Billerica, MA, USA), and the resulting samples were then sterilized using a 0.22 μm filter. Finally, the concentration of QDs was determined via a UV-2550 spectrophotometer (Shimadzu, Kyoto, Japan). Transmission electron microscopy (TEM) images and energy-dispersive X-ray spectroscopy of QDs and QD-LC were performed using a Hitachi model H-7650 TEM (Hitachi Ltd., Tokyo, Japan) operated at 80 kV. Particle size distribution and zeta potential were measured with a Nano ZS90 Zetasizer (Malvern Instruments, Malvern, UK) according to the manufacturer’s instructions. Ultraviolet-visible absorption and fluorescent emission spectra were measured at room temperature using a UV-3101 spectrophotometer and a Hitachi F-4500 fluorochrome spectrophotometer, respectively.

**Cell viability determination by MTT assay**

Human hepatocarcinoma cell lines, including HepG2, Bel-7404, SMMC-7721, Huh-7, and Bel-7402, and normal cell lines, including hepatic embryo cells HL-7702, rat hepatic satellite cells CFSC-2G, myocardium cell line H9C2, neonatal rat myocardium cells, and human umbilical vein endothelial cells, were maintained in RPMI 1640 supplemented with 10% fetal bovine serum, 100 units/mL penicillin, and 100 μg/mL streptomycin in an atmosphere of 95% air and 5% CO₂ at 37°C. The MTT assay was performed in 96-well plates in sextuplicate. Cells were seeded at a density of 5×10³ cells/
well overnight and treated with the QD-LC at final concentrations of 0, 0.05, 0.25, 0.5, 1.0, 2.0, and 4.0 μM for 24 hours and also for different time intervals, including 6, 12, 24, and 48 hours, at 2.0 μM QD-LC. Twenty microliters of MTT (5 mg/mL) was added to each well for the last 3 hours of the incubation with the QD-LC. Afterward, the cell supernatants were discarded, the MTT crystals were dissolved in dimethyl sulfoxide, and the optical density was measured at the 490 nm wavelength. The ratio of cell proliferation to control group was calculated from the data obtained by the MTT assay.

Intracellular QD-LC quantification by flow cytometry
To quantitate the uptake dynamics of the QD-LC by fluorescence-activated cell sorting, HepG2 cells were seeded in 24-well plates and then incubated with 0.2 and 2 μM QD or QD-LC for 24 hours. The supernatant solution was removed, and the cells were washed twice with chilled PBS to remove any extracellular QD-LC. Then, the cells were trypsinized and resuspended in chilled PBS to yield a concentration of 10^6 cells/mL. The excitation wavelength was set at 360 nm, and the red fluorescence content was measured at 550–590 nm, the emission wavelength.

Identification of the cellular uptake pathways of QD-LC
To study the effect of different inhibitors on the cellular uptake of the QD-LC, HepG2 and HL-7702 cells were preincubated with different inhibitors for 30 minutes at 37°C. The cellular uptake mechanisms of the QD-LC were identified by blocking the uptake pathway with different treatments. HepG2 and HL-7702 cells were preincubated with different inhibitors for 60 minutes at 37°C. For adenosine triphosphate (ATP) depletion, the cells were preincubated with NaN_3 (3.0 mg/mL). For hindering the clathrin-mediated pathway, the cells were preincubated with chlorpromazine (20 μg/mL) and sucrose (500 mM). To block the caveolae/lipid rafts pathway, the cells were pretreated with simvastatin (10 μg/mL). To block the caveolae-mediated pathway, the cells were preincubated with mycostatin (5 μg/mL). To hinder the macrophagocytosis pathway, the cells were pretreated with amiloride (10 μg/mL). To block energy-dependent endocytosis, the cells were incubated at both 37°C and 4°C, then the inhibitor-containing culture medium was discarded, and QD-LC-containing culture medium was co-cultured with the cells for a 6-hour treatment. The cells were treated as described in “Intracellular QD-LC quantification by flow cytometry” and the fluorescence intensity was determined by fluorescence-activated cell sorting.

Detection of apoptosis by flow cytometry
Cell death and apoptosis was detected using an annexin V–fluorescein isothiocyanate apoptosis detection kit by flow cytometry. In brief, after HepG2 cells were treated with 2.0 μM QD-LC for 6 or 24 hours, the cells were harvested and washed with cold PBS twice, and the cell pellets were resuspended in binding buffer to cell suspension at a density of 1x10^6 cells/mL. Five microliters of fluorescein isothiocyanate-conjugated annexin V was added to the suspension, which was incubated for 15 minutes at 4°C in the dark, then 5 μL of propidium iodide was added and the solution was mixed for 5 minutes at 4°C in the dark. The samples were subsequently analyzed by flow cytometry.

Western blot for the apoptosis signaling pathway
HepG2 cells were treated with the QD-LC for 24 hours and lysed with RIPA cell lysis buffer (Cell Signaling Technology, Inc., Danvers, MA, USA) containing a phosphatase inhibitor and the protease inhibitor cocktail (Sigma-Aldrich Co.) by incubating on ice for 30 minutes. The lysates were collected by centrifugation, and the protein concentrations were determined by the BCA method. The samples were separated on 12% sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) gels and transferred onto polyvinylidene fluoride membranes. After blocking in Tris-buffered saline buffer (150 mmol/L NaCl, 10 mmol/L Tris, pH 7.4) containing 5% nonfat milk, the blots were incubated with a primary antibody (rabbit anti-JNK, p-JNK, Bax, Bcl-2, cytochrome c, caspase-9 or caspase-3) at 4°C overnight and with a corresponding horseradish peroxidase-conjugated secondary antibody (1:2,000 dilution) for 1 hour at room temperature. The blots were visualized by super-enhanced chemiluminescence and quantified by Quantity ONE software (Bio-Rad Laboratories Inc., Hercules, CA, USA). GAPDH was used as the internal control.

Tumor formation in vivo
The use of mice for this study was approved by the Animal Care and Use Committee of Jilin University (Changchun, People’s Republic of China). Six- to eight-week-old, 18–22 g male Institute of Cancer Research (ICR) mice were purchased from the Experimental Animal Center of the Norman Bethune College of Medicine, Jilin University. First, H22 cells were incubated with 0.25 μM QD-LC for 6 hours (cell viability was not decreased compared with the control group. Supplementary material; Figure S1). The complex was injected subcutaneously into the right side of the dorsal flank of the mice, with a
total cell number of $2 \times 10^6$ for five mice in each group. Tumor volumes were measured each day and calculated according to the formula $\text{length} \times \text{width}^2 \times 0.52$. The mice bearing more than 60 mm$^3$ tumor volume were considered to be undergoing microtumor formation. The mice were sacrificed on the tenth day after the tumor formation, and all tumors were removed and weighed.

**The inhibitory tumor growth effect of QD-LC in vivo**

Six- to eight-week-old, 18–22 g male ICR mice were used in this study. The H22 cell suspension was injected subcutaneously into the right side of the dorsal flank of the mice, with a total cell number of $2 \times 10^6$. When the tumor volume reached 100 mm$^3$, 12 mice were divided into two groups; 10 μL of the QD-LC PBS solution was injected into the center of the tumor in the QD-LC group, while 10 μL PBS was injected into the tumors in the control group. The administrations were conducted once every 3 days, and the mice were sacrificed on the 19th day.

**Toxicity assay in vivo**

To assess the toxicity of QD-LC, the safety profiling included an assessment of blood chemistry and a histological examination of the major organs. Following the animal experiments described above, the mice were sacrificed on the 19th day, and all tumors were removed and weighed. Serum was collected by centrifuging the whole blood at 3,000 rpm for 15 minutes. The biochemical parameters, including ALT, AST, CRE, and BUN, were assayed using diagnostic kits provided by the Nanjing Jiancheng Bioengineering Institute.

**Results**

**Characterization of QD-LC**

TEM micrographs of QDs and QD-LC are shown in Figure 1A and B. The QDs appeared as monodisperse dark dots approximately 10 nm in size, and the QD-LCs were spherical or ellipsoidal particles with an average size of 150 nm due to a number of QDs incorporating into the lipids, resulting in a rough particle surface. The dynamic light scattering data (Figure S2A and B) indicated that QDs and QD-LC had average diameters of approximately 7 nm and 162 nm, respectively. The zeta potential of the QD-LCs was evaluated to identify the formation of the QDs and lipids (Figure S2C–E). Furthermore, the formation of the QD-LCs was identified by the energy dispersive X-ray spectroscopy spectra (Figure S3). In addition, the optical characterization of free QDs and QD-LCs is shown in Figure 1C. Upon encapsulation within the lipids, then, appropriate-size sections of the tumor(s), liver, kidney, spleen, lung, and heart were fixed in 10% formalin and then embedded into paraffin. Tissue sections of 4 μm thickness were mounted on glass slides. The sections were stained with hematoxylin and eosin and examined by light microscopy. The slides were read by an experienced veterinary pathologist.

**Statistical analysis**

Data were expressed as the mean ± standard deviation. The statistical significance of the data between two groups was compared by Student’s $t$-test. An analysis of variance was used to analyze the differences among the different groups. The results were considered significant at $P < 0.05$.

**Notes**

- TEM images showing QDs (A) before and (B) after lipid encapsulation. (C) Ultraviolet-visible absorbance and photoluminescence emission spectra of QDs (yellow line) and QD-LC (red line).
- Abbreviations: PL, photoluminescence; QD, quantum dot; QD-LC, CdTe/CdS core/shell quantum dot–lipids complex; TEM, transmission electron microscopy.
there was some degree of aggregation, resulting in the emission peak shifting from 570 nm to 600 nm. No fluorescence quenching was observed when QD-LC was stored at 4°C for 3 months (data not shown).

**QD-LC selectively killed cancer cells in a dose- and time-dependent manner**

The MTT assay was used to evaluate the cell viability of human hepatocarcinoma HepG2 cells and human hepatic embryo HL-7702 cells after incubation with various concentrations of QD and QD-LC for 24 hours. As shown in Figure 2A and B, QD-LC significantly decreased cell viability of HepG2 cells when the concentration was more than 0.5 μM, and the killing effect of QD-LC was stronger than that of QDs. Interestingly, QD-LC, even at a concentration of 2 μM, had little influence on cell viability of HL-7702 cells, and an inhibitory rate of only 32.4% was observed at 4 μM. Moreover, we carried out a time-dependent study on 2 μM QDs or QD-LC. As depicted in Figure 2C and D, the cell viability of HepG2 significantly decreased with time. The reduction was 57% in 24 hours and 60% in 48 hours, whereas the cell viability of HL-7702 cells was affected less, with a reduction of 19% in 24 hours and 35% in 48 hours. In addition, transfection reagents had no intrinsic toxicity in either cell line. The difference in susceptibility to QD-LC in HepG2 and HL-7702 has attracted our attention.

To further confirm the selectivity of QD-LC toward cancer cells, several human hepatocarcinoma cell lines, including Hep3B, Bel-7404, SMMC-7721, Huh-7, and Bel-7402, as well as mouse hepatocarcinoma H22 cells, were incubated with 2 μM QD-LC for 24 hours. Cell lines derived from normal cells, including the hepatic satellite cell

![Figure 2](https://www.dovepress.com/)

**Figure 2 (Continued)**
Selective endocytosis in cancer cells

To elucidate the mechanisms for the selective toxicity in cancer cells, we chose HL-7702 cells as representative of the normal cell lines, and all these tests were conducted under identical conditions. As depicted in Figure 2E, all tumor cell lines were more susceptible to QD-LC than the normal cell lines, according to the reduced ratios of cell viability. Thus, it is confirmed that QD-LC selectively killed the cancer cells, while the damage to normal cells was negligible.

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line CFSC-2G, human cardiomyocyte line H9C2, primary cultured myocardiocytes (neonatal rat myocardium cells), and human umbilical vein endothelial cells, were used as normal cell controls, and all these tests were conducted under identical conditions. As depicted in Figure 2E, all tumor cell lines were more susceptible to QD-LC than the normal cell lines, according to the reduced ratios of cell viability. Thus, it is confirmed that QD-LC selectively killed the cancer cells, while the damage to normal cells was negligible.

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Several inhibitors targeting various cellular uptake pathways were utilized to block the energy-dependent endocytosis, ie, ATP-utilizing pathway, the clathrin-mediated pathway, the caveolae/lipid rafts pathway, the caveolae-mediated pathway, and the macropinocytosis pathway. The results indicated that the ATP-utilizing, the clathrin-mediated, and the macropinocytosis pathways were the main uptake pathways in the energy-dependent endocytosis of QD-LC. Interestingly, we found that the significant difference in uptake between the tumor cells and the normal cells was due to the macropinocytosis pathway (shown in Figure 3D).

**QD-LC-activated, reactive oxygen species-mediated JNK apoptosis pathway in HepG2 cells**

It has been documented that the intracellular uptake of CdTe/CdS QDs can disturb the oxidative balance of the cell and induce apoptosis through the release of cadmium ions. Therefore, the apoptotic rate was determined by flow cytometry, and the mechanism underlying QD-LC-induced apoptosis in HepG2 cells was investigated. As shown in Figure 4A, the apoptotic rate in HepG2 cells incubated with 2 μM QD-LC or QDs for 24 hours was 53.9%±5.9% and 10.8%±2.3%, respectively. Subsequently, the content of reactive oxygen species (ROS) in cell culture medium was determined. The ROS levels in the QD-LC group were more elevated compared to the QD group in cancer cells (Figure S6). Considering that ROS accumulation can induce cell cycle arrest and apoptosis through the activation of the JNK/mitochondrial apoptosis pathway in cancer cells, we detected the changes of JNK and key proteins in the intrinsic mitochondrial apoptosis signaling pathway. The results showed that the protein expression of p-JNK, Bax, cytochrome c, cleaved caspase-9, and cleaved caspase-3 were all increased (Figure 4B and C), indicating
that QD-LC had the ability to activate the ROS-mediated JNK apoptosis pathway in HepG2 cells.

**Effect of QD-LC on inhibiting tumor formation and growth in vivo**

To investigate the inhibitory effect of the QD-LC on tumor growth, both a microtumor formation model and a transplanted tumor model in nu/nu mice were employed. For the microtumor formation animal model, mouse hepatocarcinoma H22 cells were pretreated with or without 0.25 μM QD-LC for 6 hours in a flask before being injected subcutaneously into the right side of the dorsal flank of the mice at a total of $3 \times 10^5$ cells for each mouse. Tumor volumes were measured every day and calculated by the formula $\text{length} \times \text{width}^2 \times 0.52$. Microtumors (volume less than 60 mm$^3$) developed in 60% of the mice in the control group after 3 days of inoculation, and a 100% rate of tumor growth existed in the control group after 4 days. However, there were no microtumors detected at 4 days in the QD-LC group, and only 20% of the mice were found to have developed microtumors after 5 days of inoculation, while 100% of the mice exhibited these tumors after 9 days (as shown in Figure 5A, Kaplan–Meier curves). Moreover, the average tumor weight in mice that were sacrificed 10 days after the incubation was $0.315 \pm 0.085$ g in the control group, but only $0.102 \pm 0.023$ g in the QD-LC pretreatment group (Figure 5B and C). There was a significant difference in tumor-free time ($P=0.0023$) between the control group and QD-LC group, indicating that QD-LC pretreatment delayed the latent period of tumor development.

In the tumor transplant model, H22 cells ($2 \times 10^6$) were injected into the right side of the dorsal flank of the mice. When the tumor volume reached 100 mm$^3$, 10 μL of 2 μM QD-LC PBS solution was injected once every 3 days into the center of the tumor in the QD-LC group, and 10 μL PBS solution was injected once every 3 days into each tumor in the control group. The tumor volume was measured every day, and the mice were sacrificed at 19 days after inoculation. As depicted in Figure 6A and B, the tumor volume was significantly decreased after 11 days in the QD-LC treatment.
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Figure 5 In vivo monitoring of tumor formation of H22 liver cancer cells.
Notes: (A) Kaplan–Meier curves showing tumor-free mouse analysis using GraphPad Prism 4 software (GraphPad Software, Inc., La Jolla, CA, USA). (B) Tumor weight analysis at the end of the experiment. Data are presented as means ± SD, n=5, *P<0.01 versus control group. (C) Photograph of the dissected tumor tissues for each treatment group.
Abbreviations: QD-LC, CdTe/CdS core/shell quantum dot–lipids complex; SD, standard deviation.

group and a significant decrease in the final weight of the tumors was observed compared to the control group. The QD-LC suppressed tumor growth, with a 53.2% reduction in tumor weight. The average final weight of the tumors in the QD-LC treatment group was significantly lower than that of the control group (3.85±1.06 g versus 1.8±0.41 g). Interestingly, the fluorescence images of the removed tumors revealed that the fluorescence of the QD-LC covered 60% of tumor volume (Figure 6C), not the entire tumor, suggesting that it may be potentially useful as an optical imaging agent in cancer diagnoses.

Toxicological evaluation of the QD-LC in vivo
We performed a toxicological evaluation by comparing the effects of different treatments. The safety profiling included an assessment of the total body weight and blood chemistry and a histological examination of the major organs. Compared to saline-treated, tumor-bearing mice, no significant body weight changes were observed following the administration of the QD-LC (Figure 6D). Moreover, in the tumor transplant model, blood was collected at the end of the experiment. The levels of ALT, AST, BUN, and CRE in the serum were measured. The results showed that all the enzymes tested were not elevated compared to the control group (Figure 6E). In addition, the histological analysis of the tissues from five treated and control mice revealed that there were no observable pathological changes in the heart, lung, kidney, liver, or spleen. The hepatocytes in the liver were normal in both groups, and there were no signs of an inflammatory response. No pulmonary fibrosis was detected in the lung samples. The glomerulus structure in the kidney section was clearly identifiable, and necrosis was not observed in any of the histological sections (Figure 6F).

Discussion
The presence of systemic toxicity and the absence of cellular and/or organ targeting of antitumor agents are key factors limiting their widespread use in cancer chemotherapy. Recently, advances have been made in the employment of various nanoparticles as drug carriers, therapeutic agents, and imaging probes in cancer diagnosis. QDs have been used as an innovative fluorescence imaging probe for biomedical research and disease diagnosis, and, recently, their potential as antitumor agents has attracted more attention from researchers. The semiconductor nanocrystal core of QDs, however, generally contains toxic heavy metals, including Cd, Se, Te, and Hg, and their toxicities have caused safety
Figure 6 QD-LC inhibited tumor growth and toxicity detection.

Notes: (A) Tumor volume curve. (B) Tumor weight analysis after treatment. Data are presented as means ± SD, n=6, *P<0.01 versus control group. (C) Representative images of excised tumors from treated mice under white or UV light. (D) Body weight curve. (E) Effect of functional enzymes ([a] ALT; [b] AST; [c] CRE; [d] BUN) in liver and kidney after QD-LC treatment (n=6). (F) Histological images from the major organs of the treated mice. Images were taken at ×40 magnification with standard HE staining.

Abbreviations: ALT, alanine aminotransferase; AST, aspartate aminotransferase; BUN, blood urea nitrogen; CRE, creatinine; HE, hematoxylin and eosin; QD-LC, CdTe/CdS core/shell quantum dot-lipids complex; SD, standard deviation; UV, ultraviolet.
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cells but only a slight increase in the nonmalignant Hs27 cells,42 which is consistent with our results. Therefore, we suggest that macropinocytosis may be partially responsible for the difference in endocytosis capabilities in both cancer and normal cells, which subsequently results in the selective toxicity seen in the tumor cells. However, the possibility of enhanced uptake via the difference in the charge and fluidity of the plasma membrane or endocytosis receptors between the tumor and normal cells cannot be ruled out.

The production of ROS, such as superoxide, hydroxyl radicals, or peroxide radicals, can adversely affect cellular functions. At a high level of oxidative stress, the antioxidant defense system would be overwhelmed by the excessive ROS, which would eventually lead to mitochondrial dysfunction as well as apoptosis.43,44 The generation of ROS by cultured cells upon exposure to inorganic nanoparticles is a quite common phenomenon.45–47 In QDs, ROS can be induced by the reactivity of surface-located transition metals, the leaching of free Cd2+ ions, or the direct interaction of QDs with mitochondria.48–50 Our results show that the intracellular and extracellular ROS levels in HepG2 cells treated with QD-LC were significantly higher than those in the control HepG2 cells (without treatment of QD-LC). Our Western blot analysis indicated that the QD-LC treatment resulted in the overexpression of p-JNK, which would stimulate the release of cytochrome c and, subsequently, activate caspase-9 and caspase-3, which suggests that the QD-LC induced the activation of the intrinsic mitochondrial apoptosis signal pathway.

Our in vivo study showed that QD-LC treatment significantly delayed tumor development and suppressed tumor growth without any detrimental effect on functional enzymes in the liver and kidney or histopathology abnormalities in the major organs after intratumoral injection. It is clear from this study that there is a biosafety margin of QD-LC when employed as a plausible agent for tumor treatment. In addition, the accumulation of QD-LC in tumors could provide an alternative way to either label solid tumor cells or monitor their migratory patterns. It is entirely possible that this work may lead to the positive identification of well-localized solid tumors and aid in their early diagnosis and treatment in clinical situations. The metabolism of QD-LC and its degradation in both normal and cancer cells, however, would be an interesting topic for further investigation.

Conclusion

We have successfully prepared QD-LCs as cancer therapeutic nanoagents. It has been illustrated that QD-LC has distinct
effects on cell viability, killing cancer cells while having little impact on normal cells. Our study demonstrated that the macropinocytic uptake pathway determined the selective toxicity for the cancer cells, which induced intrinsic mitochondrial apoptosis by the activation of the ROS-mediated JNK signaling pathway. In vivo experiments with animal models indicated that the administration of QD-LC in a controlled dose prolonged the latent period of tumor development and significantly inhibited tumor growth without obvious toxicity. Our results provide a proof-of-concept for the intrinsic toxicity of QD-based cancer therapy strategies.

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Author contributions
Dan Shao and Jing Li contributed equally to this work; thus, both are co-first authors. All authors contributed toward data analysis, drafting, and revising the paper and agree to be accountable for all aspects of the work.

Disclosure
The authors report no conflicts of interest in this work. The authors alone are responsible for the content and writing of this paper.

References
Supplementary materials

Materials and methods

CdTe/CdS core/shell quantum dot–lipids complex fluorescence imaging in HepG2 cells by laser scanning confocal fluorescence microscopy

Quantum dots (QDs) and CdTe/CdS core/shell QD–lipids complex (QD-LC) were co-cultured with HepG2 and HL-7702 cells for 6 or 24 hours in a 24-well plate, then the cells were washed twice with chilled phosphate-buffered saline (PBS) and the nuclei were stained with Hoechst 33258 (5 mg mL^-1) for 5 minutes. The QD intracellular location and cytotoxicity were observed, after washing twice, by confocal laser scanning microscopy. The confocal laser scanning microscopy was carried out using an Olympus FV1000 microscope (Olympus Corporation, Tokyo, Japan) equipped with multiline argon laser, 405, 488 nm, and 30 mW Laserclass 3D laser.

Determination of intracellular reactive oxygen species

HepG2 cells were cultured in a 24-well plate overnight and then treated with QDs or QD-LC for 24 hours. The culture medium was collected and ultra-centrifuged at 100,000 rpm for 20 minutes to provide the supernatant for analysis, while the cells were washed with chilled PBS twice. Then, the cells with 20 µL PBS solution were frozen at −80°C and thawed at 25°C three times to break the cells. Reactive oxygen species content was then analyzed using a reactive oxygen species detection kit.

Figure S1 Cell viability of QD-LC on H22 cell lines at 6 and 24 hours.

Notes: The data represent three separate experiments and are presented as mean values ± SD. *P<0.05 versus control group.

Abbreviations: QD-LC, CdTe/CdS core/shell quantum dot–lipids complex; SD, standard deviation; T, transfection agent.

Figure S2 Characterization of QDs and QD-LC.

Notes: Average diameters, found by DLS, of QDs (A) and QD-LC (B). Zeta potentials of QDs (C), lipids (D), and QD-LC (E) were −40.4 mV, +34 mV, and −18.3 mV, respectively. These results indicated the conjugation of the QDs and lipids.

Abbreviations: DLS, dynamic light scattering; QD, quantum dot; QD-LC, CdTe/CdS core/shell quantum dot–lipids complex.
**Figure S3** Energy-dispersive X-ray (EDX) spectroscopy analysis of CdTe/CdS core/shell quantum dot–lipids complex.

**Figure S4** Intracellular distribution of QD-LC in HepG2 cells.

**Notes:** Confocal microscopy images of HepG2 cells after 24 hours’ incubation with different QD concentrations and different QD-LC concentrations (red). The cell nucleus is stained with Hoechst 33258 (blue). The scale bar represents 10 μm.

**Abbreviations:** QD, quantum dot; QD-LC, CdTe/CdS core/shell quantum dot–lipids complex.
**Figure S5** Intracellular distribution of QD-LC in HL-7702 cells.

**Notes:** Confocal microscopy images of HL-7702 cells after 24 hours' incubation with different QD concentrations and different QD-LC concentrations (red). The cell nucleus is stained with Hoechst 33358 (blue). The scale bar represents 10 μm.

**Abbreviations:** QD, quantum dot; QD-LC, CdTe/CdS core/shell quantum dot–lipids complex.

**Figure S6** Comparative effects of QDs and QD-LC on intracellular and extracellular ROS generation in HepG2 cells.

**Note:** *P < 0.05 and #P < 0.05 versus control group.

**Abbreviations:** L, lipids; QD, quantum dot; QD-LC, CdTe/CdS core/shell quantum dot–lipids complex; ROS, reactive oxygen species.
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