Targeted co-delivery of Beclin 1 siRNA and FTY720 to hepatocellular carcinoma by calcium phosphate nanoparticles for enhanced anticancer efficacy

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Purpose: FTY720, known as fingolimod, is a new immunosuppressive agent with effective anticancer properties. Although it was recently confirmed that FTY720 inhibits cancer cell proliferation, FTY720 can also induce protective autophagy and reduce cytotoxicity. Blocking autophagy with Beclin 1 siRNA after treatment with FTY720 promotes apoptosis. The objective of this study was to enhance the anticancer effect of FTY720 in hepatocellular carcinoma (HCC) by targeted co-delivery of FTY720 and Beclin 1 siRNA using calcium phosphate (CaP) nanoparticles (NPs).

Materials and methods: First, the siRNA was encapsulated within the CaP core. To form an asymmetric lipid bilayer structure, we then used an anionic lipid for the inner leaflet and a cationic lipid for the outer leaflet; after removing chloroform by rotary evaporation, these lipids were dispersed in a saline solution with FTY720. The NPs were analyzed by transmission electron microscopy, dynamic light scattering and ultraviolet–visible spectrophotometry. Cancer cell viability and cell death were analyzed by MTT assays, fluorescence-activated cell sorting analysis and Western blotting. In addition, the in vivo effects of the NPs were investigated using an athymic nude mouse subcutaneous transplantation tumor model.

Results: When the CaP NPs, called LCP-II NPs, were loaded with FTY720 and siRNA, they exhibited the expected size and were internalized by cells. These NPs were stable in systemic circulation. Furthermore, co-delivery of FTY720 and Beclin 1 siRNA significantly increased cytotoxicity in vitro and in vivo compared with that caused by treatment with the free drug alone.

Conclusion: The CaP NP system can be further developed for co-delivery of FTY720 and Beclin 1 siRNA to treat HCC, enhancing the anticancer efficacy of FTY720. Our findings provide a new insight into HCC treatment with co-delivered small molecules and siRNA, and these results can be readily translated into cancer clinical trials.

Keywords: LCP-II NPs, autophagy, FTY720, Beclin 1, co-delivery

Introduction

With ~700,000 deaths annually, hepatocellular carcinoma (HCC) is the fifth most common malignancy in the world and a serious threat to human health.1 The development and progression of HCC are associated with multiple genetic mutations and genetic heterogeneity.2 At present, the most effective treatment for HCC is surgical resection and liver transplantation.3–6 However, surgical resection and liver transplantation still faces major problems, such as tumor recurrence and metastasis.4 In addition, surgical therapy is always ineffective for patients with advanced HCC.5 Therefore, it is necessary to develop novel therapies for HCC treatment.6
FTY720 (fingolimod), a Federal Drug Administration–approved drug and the first oral therapeutic for multiple sclerosis, was recently confirmed to possess strong antitumor effects.\textsuperscript{5,7,8} Compared with traditional chemotherapies, FTY720 may be a beneficial anticancer drug because it is less toxic and has better oral bioavailability. Although FTY720 has been widely used in various cancers, an increasing number of studies have found that FTY720 induces cell autophagy, adversely affecting apoptosis and contributing to tumor resistance.\textsuperscript{9,10} Autophagy in tumor therapy is associated with chemotherapy, radiation and immune tolerance.\textsuperscript{11,12} In addition, increased autophagy after treatment with anticancer drugs reduces the effect of drugs.\textsuperscript{13,14} Furthermore, some studies have confirmed that blocking autophagy through Beclin 1 siRNA (siBeclin 1) or ATG 5 siRNA (siATG 5) increases cancer cell sensitivity to FTY720.\textsuperscript{15,16} Gene silencing strategies, particularly RNA interference, have been widely applied to inhibit the expression of cancer-related proteins.\textsuperscript{17,18} Combining drug and siRNA therapy to downregulate cancer-related gene expression, promoting the anticancer drug effects at the tumor site can effectively suppress tumor progression, and an increasing number of studies have focused on using nonviral vectors to co-deliver drugs and siRNA.\textsuperscript{19,20} Incorporating drugs and siRNA into one vector is the most obvious method for achieving a true synergistic effect from simultaneous delivery of both a drug and an siRNA.\textsuperscript{21} The objective of this study was to co-deliver FTY720 and siRNA to enhance the effect of this anticancer drug.

Constructing a co-delivery vector has several challenges, which are as follows: the drug and siRNA must be loaded at a high entrapment rate, the carrier needs to remain stable in systemic circulation and targetted release of the drug and siRNA at the tumor site must be achieved.\textsuperscript{22} In recent studies, nanoparticles (NPs), which are nanoscale carriers, have demonstrated great value and potential for promoting drug/gene delivery.\textsuperscript{23,24} For instance, Kang et al confirmed that cationic liposomal co-delivery of siRNA and an MEK inhibitor improved anticancer efficacy.\textsuperscript{25} In addition, Ding et al designed a nanostructure-based drug and siRNA delivery system to knock down the expression of Rap2b, a novel p53 target, enhancing adriamycin-based anticancer therapeutics.\textsuperscript{26} Furthermore, the use of a novel liposome to efficiently deliver siRNA against MGMT, a DNA repair protein, promoted temozolomide cytotoxicity,\textsuperscript{27} and the strategy of simultaneously co-delivering epirubicin and BCL-2 siRNA with lipid-NPs significantly reversed tumor cell multidrug resistance.\textsuperscript{28} In summary, these combined therapies provide an experimental basis for their use to reduce clinical drug resistance.

Recently, Li et al designed calcium phosphate (CaP) NPs with an asymmetric lipid bilayer coating for targeted delivery of siRNA to tumors.\textsuperscript{30} An anionic lipid, dioleoylphosphatidic acid (DOPE), was used as the inner leaflet lipid to coat the nanosized CaP cores, thereby entrapping the siRNA and making the coated cores soluble in organic solvent; a suitable neutral or cationic lipid was used as the outer leaflet lipid to form an asymmetric lipid bilayer structure. The formulation was named lipid/calcium/phosphate type II (LCP-II). The results showed that the improved formulation effectively delivered siRNA in vitro and to the tumor site in vivo. In this study, we used LCP-II with minor modifications to co-deliver Beclin 1 siRNA and FTY720 (termed LCP-II-siBeclin 1-FTY720) to enhance cytotoxicity by blocking the protective autophagy induced by FTY720. LCP-II has great potential for simultaneously transporting siRNA and FTY720 across the cell membrane to reduce Beclin 1 expression and to induce more FTY720-mediated cytotoxicity than free FTY720 alone.

**Materials and methods**

**Materials**

DOPA, 1,2-dioleoyl-3-trimethylammonium propane (DOTAP), 1,2-distearoyl-sn-glycero-3-phosphoethanolamine-N-methoxy(polyethylene glycol-2000) ammonium salt (DSPE-PEG) and DSPE-PEG-folic acid (DSPE-PEG-FA) were purchased from Avanti Polar Lipids, Inc. (Alabaster, AL, USA). Cholesterol was purchased from Sigma-Aldrich. Beclin 1 siRNA (target sequence 5′-CAGTTTGGCACAATC AATA-3′) labeled with or without fluorescein (FAM or Cy5) and control siRNA (target sequence 5′-AAATCTCAGAACGT GTCACGT-3′) labeled with or without FAM were purchased from Shanghai GenePharma Co., Ltd. (Shanghai, China).

**Cell lines and cell culture**

The cell lines SMMC-7721 and A549 were purchased from the Cell Bank of the Chinese Academy of Sciences (Shanghai, China) and were cultured in complete DMEM (Wisent Inc., St-Bruno, QC, Canada) supplemented with 10% fetal bovine serum (Wisent Inc.) at 37°C in a humidified incubator with 5% CO\textsubscript{2}.

**Preparation and characterization of LCP-II NPs**

LCP-II NPs were prepared according to Huang’s protocol with minor modifications as previously described.\textsuperscript{31} Figure 1
illustrates a flow diagram of the preparation of LCP-II-siBeclin 1-FTY720. First, 300 μL of CaCl₂ (500 mM) and 100 μL of siBeclin 1 (2 mg/mL) were dispersed in 15 mL of a cyclohexane/Igepal CO-520 (71/29 volume/volume [V/V]) solution to prepare the water-in-oil microemulsion. To form the phosphate portion, 300 μL of Na₂HPO₄ (25 mM, pH = 9.0) solution was added to another 15 mL of cyclohexane/Igepal CO-520 (71/29 V/V) solution, and then, 200 μL of DOPA (20 mg/mL) in chloroform was added to the phosphate phase. After the two solutions were mixed for 20 min, 30 mL of absolute ethanol was added to the mixture, and the resulting mixture was centrifuged at 12,000× g for 20 min to remove the cyclohexane/Igepal and to pellet the CaP core. The pellets were washed with ethanol two to three times, dissolved in 1 mL of chloroform and then stored in a glass vial for the upcoming procedures.

Figure 1 Schematic for the preparation of LCP-II NPs for co-delivery of siRNA and FTY720. Abbreviations: DOPA, dioleoylphosphatidic acid; FTY720, fingolimod; LCP-II NPs, lipid/calcium/phosphate type II nanoparticles.
without FA were prepared by the method mentioned above. LCP-II-siRNA (LCP-II–entrapped siRNA NPs with FA) was used as a negative control. LCP-II-siBeclin 1 represents LCP–II–entrapped siBeclin 1 NPs with FA. LCP-II-siRNA-FTY720 represents LCP–II–entrapped siRNA and FTY720 NPs with FA, which was used as a negative control. LCP-II-siBeclin 1-FTY720 represents LCP–II–entrapped siBeclin 1 and FTY720 NPs with FA. Except where noted, the different formulations were all prepared with the FA ligand.

To measure the encapsulation efficiency (EE) of FTY720 in the LCP-II NPs after they were dispersed in saline solution with 10 mM FTY720, the LCP-II NP–containing solution was centrifuged at 12,000 × g for 20 min and the supernatant was harvested for absorbance detection with an ultraviolet–visible spectrophotometer (V500; Jasco, Tokyo, Japan). FTY720 in saline solution had a maximum absorption wavelength of 220 nm. The absorbance of different samples was obtained at 220 nm. The concentration of FTY720 was measured according to the Beer–Lambert law, and the EE was determined according to the following equation: EE=(1–concentration of free FTY720/concentration of total FTY720)×100%.

LCP-II NPs were examined by transmission electron microscopy (TEM) as previously described with small modifications. LCP-II NPs (1 μL) were resuspended in 1 mL of PBS and applied to copper-mesh formvar grids (Beijing Zhongxingkeyi Technology Co., Ltd., Beijing, China) with or without negative staining with phosphotungstic acid. TEM images were obtained with a JEM-2100 transmission electron microscope (Jeol, Tokyo, Japan). For size and zeta potential measurements, 1 μL of LCP-II NPs was resuspended in 1 mL of PBS, and this LCP-II NP–containing PBS solution was used for size and zeta potential measurements with a Zetasizer Nano ZS90 instrument (Malvern Instruments, Malvern, UK).

In vitro release of FTY720 from LCP-II NPs

To measure the release rate of FTY720 from LCP-II-siRNA-FTY720 and LCP-II-siBeclin 1-FTY720, the NPs were added to PBS (pH=7.4) at 37°C. The unreleased FTY720, which was retained within the LCP-II-siRNA-FTY720 and LCP-II-siBeclin 1-FTY720 NPs, was removed at different time points (0, 6, 12, 24, 36 and 48 h) by centrifugation at 12,000× g for 20 min, and then, the supernatant (the released portion) was harvested for absorbance detection by a ultraviolet–visible spectrophotometer. The concentration of FTY720 was measured as described above. The percentage of released FTY720 was calculated by comparing the released portion to the original total FTY720 amount. To measure the stability of LCP-II NPs in culture medium, the NPs were added to the culture medium for 6 and 12 h and the sizes of LCP-II-siBeclin 1 and LCP-II-siBeclin 1-FTY720 were measured with a Zetasizer Nano ZS90 instrument (Malvern Instruments).

Cell uptake of LCP-II NPs

SMMC-7721 cells (1×10^4 per well) were cultured on coverslips in 12-well plates. After 24 h, the cells were treated with LCP-II-siRNA with or without FA, LCP-II-siBeclin 1 with or without FA or LCP-II-siBeclin 1-FTY720 with or without FA at 37°C for 4 h (100 nM total siRNA labeled with FAM) and then washed with PBS twice. The cells were further fixed with 4% paraformaldehyde at room temperature for 20 min and, again, were washed with PBS twice. After being counterstained with DAPI, cell images were obtained using a confocal microscope (FV10i; Olympus Corporation, Tokyo, Japan).

Cell viability analysis

To measure the cell viability in vitro, SMMC-7721 cells were cultured in 96-well plates at a concentration of 1×10^4 cells/well. After 24 h, the cells were treated with PBS, free FTY720 solution, solution containing free FTY720 and lipo2000-siBeclin 1, LCP-II-siRNA, LCP-II-siBeclin 1, LCP-II-siRNA-FTY720, LCP-II-siBeclin 1-FTY720 without FA or LCP-II-siRNA-FTY720-siBeclin 1 with FA for 24 h (100 nM total siRNA). Lipo2000-siBeclin 1 represents the cells transfected with siBeclin 1 using Lipofectamine 2000 (Thermo Fisher Scientific, Waltham, MA, USA) in accordance with the manufacturer’s instructions. Then, MTT solution (5 mg/mL, 20 μL/well) was added to the 96-well plates. After incubation for 4 h, 150 μL of dimethyl sulfoxide was added to dissolve the insoluble crystals. Absorbance values were measured with a spectrophotometer at 490 nm. All the experiments were performed with three replicates.

Flow cytometric analysis

SMMC-7721 cells (1×10^4) were seeded in six-well plates for 24 h and treated with PBS control, LCP-II-siBeclin 1, LCP-IIsiBeclin 1-FTY720 with or without FA or LCP-II-siBeclin 1-FTY720 plus 1 mM FA at 37°C for 4 h (100 nM total siRNA labeled with FAM). A549 cells (1×10^4) were seeded in six-well plates for 24 h and treated with PBS control or LCP-II-siBeclin 1-FTY720 with FA at 37°C for 4 h (100 nM total siRNA labeled with FAM). The uptake of LCP-II NPs
by SMMC-7721 and A549 cells was confirmed qualitatively by flow cytometry (FACSCalibur; BD, Franklin Lakes, NJ, USA). To evaluate the apoptosis rate, SMMC-7721 cells (1×10^4) were seeded in six-well plates for 24 h and treated with a solution of free FTY720, a solution containing both free FTY720 and lipo2000-siBeclin 1, or one of the following NP solutions for 24 h: LCP-II-siRNA, LCP-II-siBeclin 1, LCP-II-siRNA-FTY720, LCP-II-siBeclin 1-FTY720 without FA or LCP-II-siRNA-LCP-II-siBeclin 1 with FA (100 nM total siRNA, 7.5 μM FTY720). Cells, including apoptotic, dead and adherent cells, were collected and washed with cold PBS and then resuspended in annexin V-fluorescein isothiocyanate (FITC) binding buffer. After incubation with annexin V-FITC/propidium iodide (PI), apoptosis was detected by flow cytometry.

**Western blot analysis**

After being treated with free FTY720 solution, solution containing both free FTY720 and lipo2000-siBeclin 1, or LCP-II-siRNA, LCP-II-siBeclin 1, LCP-II-siRNA-FTY720, LCP-II-siBeclin 1-FTY720 without FA or LCP-II-siRNA-LCP-II-siBeclin 1 with FA (100 nM total siRNA, 7.5 μM FTY720), the SMMC-7721 cells were lysed with radioimmunoprecipitation assay peptide lysis buffer (Beyotime Biotech, Haimen, China) containing 1% protease inhibitors (Thermo Fisher Scientific). The proteins were quantified through a BCA protein assay kit (Thermo Scientific). After the proteins were quantified, they were fractionated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis. Then, the proteins were stained with antibodies targeting Beclin 1, LC3B, PARP, cleaved PARP, cleaved caspase-9, and cleaved caspase-3 from Cell Signaling Technology (Beverly, MA, USA); the dilution of the antibodies used was 1:1,000. The control antibody (anti-GAPDH, 1:10,000) was from Bio-world Technology, Inc. (St. Louis Park, MN, USA). The protein bands were detected using enhanced chemiluminescence (EMD Millipore, Billerica, MA, USA) after incubation with horseradish peroxidase–conjugated secondary antibodies.

**Quantitative real-time-polymerase chain reaction analysis**

SMMC-7721 cells were seeded and transfected with liposome-siBeclin 1, LCP-II-siRNA, LCP-II-siBeclin 1, LCP-II-siRNA-FTY720, LCP-II-siBeclin 1-FTY720 without FA or LCP-II-siRNA-LCP-II-siBeclin 1 with FA NPs for 24 h (100 nM total siRNA, 7.5 μM FTY720). RNA was extracted, purified, and converted to cDNA using an RNAiso Plus kit and PrimeScript RT Master Mix (Takara Bio Inc., Naha, Japan). Quantitative polymerase chain reaction was performed using SYBR Premix Ex Taq II (Takara Bio Inc.).

**Tumor accumulation studies**

All animal studies were conducted according to the national guidelines of the Institutional Animal Care and Use Committee and were approved by the Animal Care Ethics Committee of Nanjing Drum Tower Hospital. Four- to six-week-old male nude mice (BALB/c nude mice) were purchased from the Laboratory Animal Center of Shanghai, Academy of Science. To evaluate tumor-targeting ability, SMMC-7721 cells were resuspended in PBS and injected into the right axillary region of mice after infiltration anesthesia with isoflurane (1×10^7 cells in 200 μL of PBS per mouse). When the tumor grew to ~800 mm³, the mice were injected intravenously with PBS, LCP-II-siBeclin 1 (Cy5 labeled), LCP-II-siBeclin 1-FTY720 (Cy5 labeled) or LCP-II-siBeclin 1-FTY720 (Cy5 labeled) without FA at the given FTY720 (5 mg/kg) and siRNA (2 mg/kg) dose. After 12 h, the Cy5 fluorescence signals throughout the whole body were observed with an IVIS Lumina III system (PerkinElmer, Akron, OH, USA).

**Mice and in vivo tumor studies**

SMMC-7721 cells were resuspended in PBS and injected in the right axillary region of mice (1×10^6 cells in 200 μL of PBS per mouse). After 2 weeks, when the tumor volumes were ~100 mm³, the mice were randomly divided into five groups (25 mice in total, with five in each of the following groups: PBS, LCP-II negative control, LCP-II-siRNA-FTY720, LCP-II-siBeclin 1-FTY720 without FA and LCP-II-siBeclin 1-FTY720 with FA) and were injected daily through the tail vein with different complexes (5 mg/kg total FTY720, 2 mg/kg siRNA) for 3 weeks. The longest (a) and the shortest (b) tumor diameters were examined every 4 days with a Vernier caliper. The tumor volume (mm³) = 1/2axbxb, and all the mice were sacrificed 35 days after the subcutaneous SMMC-7721 cell injection.

**Statistical analysis**

All results are expressed as the mean ± SD. The data were analyzed with Student’s t-test. When P<0.05, the data were considered significantly different.

**Results**

**Preparation and characterization of LCP-II NPs**

LCP-II NPs were prepared as previously described, with minor modifications. Briefly, we prepared a pH-sensitive CaP core encapsulating siRNA and added the amphiphilic
phospholipid DOPA as a precoating reagent. The CaP core was mixed with DOTAP/cholesterol (1:1) and DSPE-PEG-2000 with or without FA to synthesize the LCP-II NPs. After removing chloroform with rotary evaporation, the remaining lipid was dispersed in saline with or without FTY720 to form the LCP-II-siBeclin 1-FTY720 co-delivery NPs (Figure 1). Finally, the NPs were extruded through 200 and 100 nm polycarbonate membranes three times. The size, morphology and surface charge of the LCP-II NPs were determined by TEM (Figure 2A) and a Zetasizer Nano ZS90 instrument. After three extrusion cycles through the 200 and 100 nm polycarbonate membranes, the average particle size of the LCP-II-siRNA NPs was 57±6.4 nm (Figure 2B), and after loading with FTY720, the size was 76±3.9 nm (Figure 2B). The zeta potential was ~5 mV for LCP-II-siRNA NPs and 12 mV for LCP-II-siRNA-FTY720 NPs (Figure 2C), suggesting that FTY720 loading slightly influenced the surface zeta potential of the complexes. The EE of LCP-II NPs loaded with FTY720 was 36.1±2.4%.

To determine the in vitro release kinetics of FTY720 from the LCP-II-siRNA-FTY720 NPs, the formulation was incubated in PBS (pH=7.4) at 37°C. The FTY720 released from the LCP-II-siRNA-FTY720 NPs into the medium after 12 h was <30%, but after 24 h, the LCP-II NPs still retained over 45% of the FTY720 (Figure 2D). In addition, the amount of FTY720 released from LCP-II-siBeclin 1-FTY720 NPs was similar to the amount released from LCP-II-siRNA-FTY720 NPs (Figure 2D). Furthermore, the sizes of LCP-II-siBeclin 1 and LCP-II-siBeclin 1-FTY720 remained stable in culture medium (Figure 2E). These results indicated that the LCP-II-siRNA-FTY720 and LCP-II-siBeclin 1-FTY720 NPs were stable in PBS and culture medium.

**Figure 2** Characterization of LCP-II NPs loaded with siRNA and FTY720.

**Notes:** (A) Representative TEM images of LCP-II-siRNA (a) without and (b) with negative staining and LCP-II-siRNA-FTY720 NPs (c) without and (d) with negative staining. Arrows in (b, d) show the lipid bilayer surrounding the CaP core. (B, C) Size and zeta potential distribution of LCP-II-siRNA and LCP-II-siRNA-FTY720. (D) FTY720 released from LCP-II-siRNA-FTY720 and LCP-II-siBeclin 1-FTY720 NPs over time in vitro. (E) The sizes of LCP-II-siBeclin 1 and LCP-II-siBeclin 1-FTY720 in the culture medium.

**Abbreviations:** FTY720, fingolimod; LCP-II NPs, lipid/calcium/phosphate type II nanoparticles; siBeclin 1, Beclin 1 siRNA; TEM, transmission electron microscopy.
Cellular uptake of LCP-II NPs and the in vitro gene silencing activity of LCP-II NP–delivered siBeclin 1

The folate receptor (FR) is highly expressed in various cancers, including HCC, and has low expression levels in normal tissue.37 FA, a necessary vitamin for human cells, has a high affinity for the FR. In addition, coupling FA to other molecules maintains the affinity of FA for FR, and FA can be internalized through cell endocytosis. To achieve ligand-mediated siRNA and drug uptake in target cells, FA was coupled to the distal end of DSPE-PEG, which is compatible with FR as a targeting ligand. The cellular uptake of LCP-II NPs by SMMC-7721 cells was evaluated using fluorescence microscopy. siRNA and siBeclin 1 were labeled with FAM. LCP-II-siRNA NPs with or without FA, LCP-II-siBeclin 1 NPs with or without FA or LCP-II-siBeclin 1-FTY720 NPs with or without FA were added to SMMC-7721 cells. After 4 h, images of the SMMC-7721 cells were collected. The results demonstrated that SMMC-7721 cells presented with green intracellular fluorescence following incubation with FAM-siRNA–loaded LCP-II NPs with FA, indicating that FAM-labeled siRNA was evenly distributed throughout the cytoplasm of SMMC-7721 cells (Figure 3A). However, when the LCP-II NPs that were not tethered to FA were used, they were not easily internalized by the cells (Figure 3B). In addition, LCP-II-siBeclin 1 and LCP-II-siBeclin 1-FTY720 were effectively internalized by the cells, indicating that co-loading siRNA and FTY720 into the LCP-II NPs did not affect NP function (Figure 3C–F). In addition, the uptake of LCP-II-siBeclin 1-FTY720 by SMMC-7721 cells was confirmed qualitatively by flow cytometry (Figure 3G). The results showed that the co-delivery nanocarrier was effectively taken up by SMMC-7721 cells, but not when LCP-II was not tethered with FA. Moreover, to confirm that FA is a targeting ligand that improves cellular uptake, 1 mM free FA was added to the cells, and the results showed that the FR targeting effect was blocked by 1 mM free FA (Figure 3G). Furthermore, we used A549 cells, which are FR-negative tumor cells,38,39 to measure the cellular uptake of LCP-II NPs. The results showed that the uptake of LCP-II-siBeclin 1-FTY720 by A549 cells was very low (Figure 3H), which confirmed that the FR-targeted nanocarriers showed delivery specificity to FR-positive cells, but not FR-negative cells. Moreover, the expression of Beclin 1 was significantly inhibited after treatment with LCP-II-siBeclin 1 and LCP-II-siBeclin 1-FTY720, as measured by Western blotting (Figure 3I). In addition, the Beclin 1 mRNA level was inhibited (Figure 3J). Most interestingly, the effect of LCP-II-siBeclin 1 and LCP-II-siBeclin 1-FTY720 on Beclin 1 expression was greater than that of lipo2000-siBeclin 1. Furthermore, the gene silencing activity of siBeclin 1 was greatly reduced when we used LCP-II-siBeclin 1-FTY720 without FA, instead of LCP-II-siBeclin 1-FTY720. In conclusion, the results confirmed that LCP-II NPs can be internalized by cells.

In vitro cytotoxicity of LCP-II NPs

After treatment with the PBS control, LCP-II-siRNA or LCP-II-siBeclin 1 NPs for 24 h, cell viability was determined by MTT assays. The results indicated that LCP-II NPs without FTY720 did not affect cell proliferation (Figure 4A). To evaluate the cytotoxicity of co-delivered siBeclin 1 and FTY720 with LCP-II NPs compared with that of the other formulations, SMMC-7721 cells were treated with different formulations at different concentrations for 24 h, and the inhibitory effect on cell growth was determined by MTT assays. The results showed that compared with free FTY720 or the different formulations used to treat SMMC-7721 cells, the co-delivery NPs induced the lowest cell viability. Cell growth was inhibited by 33% with 7.5 μM free FTY720. When Beclin 1 expression, which is a mediator of cell autophagy, was blocked with lipo2000-siBeclin 1, the autophagy induced by FTY720 was significantly inhibited and cell cytotoxicity was further increased. The cell growth inhibition increased to ~47% after protective autophagy was impaired. After treatment with LCP-II–entrapped FTY720 NPs, cell growth was inhibited by ~50%. More importantly, cell growth was inhibited by 65% after treatment with LCP-II-siBeclin 1-FTY720 NPs, which effectively co-delivered siBeclin 1 and drug to the target cells. However, without the FA ligand, the cytotoxicity of the co-delivery LCP-II NPs decreased significantly. In summary, the results showed that co-delivery of FTY720 and siBeclin 1 with LCP-II NPs induced the highest level of inhibition and the highest levels of cytotoxicity (Figure 4B).

GFP-LC3–transfected SMMC-7721 cells treated with LCP-II NPs

To investigate whether LCP-II-siBeclin 1-FTY720 NPs effectively blocked the cell autophagy induced by FTY720, SMMC-7721 cells were transfected with GFP-LC3, which is a well-known indicator of cell autophagy. Endogenous LC3 that is diffusely distributed in the cytoplasm organizes into punctate structures, with the occurrence of autophagy. The results showed that after treatment with free FTY720 and LCP-II-siRNA-FTY720 NPs, bright green fluorescent dots were present in SMMC-7721 cells, confirming that FTY720
induced cell autophagy (Figure 5A). After treatment with LCP-II-FTY720 NPs for 24 h, the proportion of SMMC-7721 cells forming LC3 puncta was 63.3%, which was higher than the percentage of apoptotic cells resulting from treatment with free FTY720. Compared with LCP-II-siRNA-FTY720 NPs, treatment with LCP-II-siBeclin 1-FTY720 observably reduced the number of green fluorescent aggregates of endogenous LC3 in SMMC-7721 cells, which confirmed that LCP-II-siBeclin 1-FTY720 NPs blocked FTY720-induced autophagy. In addition, when LCP-II-siBeclin 1-FTY720 NPs without FA were added to the cells, the proportion of SMMC-7721 cells with typical LC3 puncta was reduced
to 34.6%, which indicated that without FA targeting, the co-delivered LCP-II NPs were not internalized by the cells (Figure 5B). In addition, in the presence of autophagy, LC3-I is converted to LC3-II, an autophagy indicator. The level of LC3-II was determined by Western blotting, and the results showed that after treatment with free FTY720 and LCP-II-siRNA-FTY720 NPs, much of the LC3-I had been converted to LC3-II. When cells were treated with LCP-II-siBeclin 1-FTY720 NPs, the levels of LC3-II and Beclin 1 were significantly decreased (Figure 5C). In summary, co-delivery of siBeclin 1 and FTY720 effectively blocked the cell autophagy induced by FTY720.

Apoptosis assay
To evaluate the synergistic effects of LCP-II-siBeclin 1-FTY720 co-delivery NPs on SMMC-7721 cell apoptosis, we assessed apoptosis in SMMC-7721 cells after treatment with different NP formulations. Annexin V-FITC/PI apoptosis staining was performed, and the percentage of apoptotic cells was analyzed by flow cytometry. The apoptotic rate of SMMC-7721 cells after treatment with free FTY720 was ~22%. After treatment with LCP-II-siRNA-FTY720 NPs, the percentage of apoptotic cells increased to 36%. When Beclin 1 protein was suppressed in SMMC-7721 cells with lipo2000-siBeclin 1 and LCP-II-siBeclin 1-FTY720, the apoptotic rates were 38% and 51%, respectively. The co-delivery of siBeclin 1 and FTY720 with LCP-II NPs significantly increased apoptosis compared with that of the other groups (Figure 6A). Without FA, the effect of LCP-II-siBeclin 1-FTY720 was significantly reduced to even less than that of free FTY720. The results showed that the suppression of autophagy by LCP-II-siBeclin 1-FTY720 NPs significantly increased cell death, which was associated with apoptosis induction.

Moreover, to further evaluate the levels of apoptotic proteins, such as cleaved caspase-3, cleaved caspase-9 and cleaved PARP, in SMMC-7721 cells, we conducted Western blotting analyses of the protein extracts from the SMMC-7721 cells after treatment with different complexes. Consistent with the flow cytometry analysis results, treatment with free FTY720 and LCP-II-siRNA-FTY720 induced apoptosis. Additionally, compared with the levels in LCP-II-siRNA-FTY720- or free FTY720-treated cells, the addition of LCP-II-siBeclin 1-FTY720 to SMMC-7721 cells significantly increased the levels of cleaved caspase-3, caspase-9 and PARP, indicating that apoptosis had been induced (Figure 6B). In summary, these results indicated that co-delivery of siBeclin 1 and FTY720 with LCP-II NPs offers great potential for treatment of HCC.

Tumor accumulation studies of LCP-II NPs
To measure the effect of LCP-II NPs on tumor accumulation, we added Cy5-labeled siBeclin 1 to the LCP-II-siBeclin 1 and LCP-II-siBeclin 1-FTY720 NPs with or without FA to track tumor targeting with an in vivo imaging system. Tumor-bearing male BALB/c nude mice were intravenously injected...
with PBS, LCP-II-siBeclin 1 or LCP-II-siBeclin 1-FTY720 NPs with or without FA at the given doses. After 12 h, the Cy5 signal at the tumor site was much higher in mice treated with LCP-II-siBeclin 1 and LCP-II-siBeclin 1-FTY720 NPs with FA than in those administered LCP-II-siBeclin 1-FTY720 NPs without FA. Without FA, the LCP-II-siBeclin 1-FTY720 remained in other parts of the body and did not target the tumor (Figure 7A). We also investigated the effect of the co-delivery nanocarrier on biodistribution. After 4 h, the Cy5 signal at the tumor site was much higher than in the...
liver and kidneys (Figure 7B), which confirmed that LCP-II was effectively delivered to the tumor site and preferred to accumulate in the tumor. The results indicated that targeting of FTY720 and siBeclin 1 to the tumor by co-delivery with LCP-II-siBeclin 1-FTY720 was feasible in vivo.

**In vivo antitumor efficacy of LCP-II NPs**

To determine whether co-delivery of siBeclin 1 and FTY720 with LCP-II NPs significantly enhanced the sensitivity of SMMC-7721 cells to FTY720 in vivo, we established a subcutaneous xenograft model in nude mice by intravenous...
Tail injection of different NP complexes to evaluate the antitumor efficacy. The results showed that LCP-II-siRNA-FTY720 NP treatment caused a mild reduction in tumor volume and weight compared with the PBS-treated group. In addition, after treatment with LCP-II-siBeclin 1-FTY720 NPs, the tumors were significantly smaller than those in mice treated with LCP-II-siRNA-FTY720 NPs, indicating that the autophagy induced by FTY720 protected the cells from apoptosis. The volume and weight of tumors from nude mice treated with LCP-II-siBeclin 1-FTY720 NPs were four-fold less than those from nude mice treated with the PBS control. However, when we used LCP-II-siBeclin 1-FTY720 NPs without FA in mice, the antitumor efficacy was markedly reduced and even lower than that of LCP-II-siRNA-FTY720 NPs with FA, which indicated that the FA ligand effectively identified and targeted SMMC-7721 cells and tumor xenografts in the nude mice (Figure 8A). In addition, there was no difference in the body weight of mice after the different treatments (Figure 8B), and no mice died in the experiment, which confirmed the low toxicity.

Figure 7 Evaluation of LCP-II NP tumor targeting.
Notes: (A) Whole body fluorescence imaging of mice after intravenous injection of the indicated formulations and quantitative analysis of fluorescence intensity at the tumor site (blue circle). (B) After 4 and 12 h, the biodistribution of fluorescence-labeled siRNA delivered by LCP-II in the livers, kidneys and tumors of mice and quantitative analysis of fluorescence intensity at the tumor site. (n=3; ***P<0.001).
Abbreviations: FA, folic acid; FTY720, fingolimod; LCP-II NPs, lipid/calcium/phosphate type II nanoparticles; siBeclin 1, Beclin 1 siRNA.
of LCP-II. These changes in tumor weight and volume are shown in Figure 8C and D, confirming that co-delivering siRNA and drug with LCP-II NPs had a greater effect on inhibiting the growth of solid tumors than that induced by the FTY720-loaded LCP-II NPs.

However, more follow-up research is still needed to verify the toxic effects and mortality induced by the nanocarrier. In summary, these results indicated that co-delivery with nanocarriers and a targeting ligand is an effective new strategy for inhibiting tumor growth.

Discussion

FTY720 is a new immunomodulator that has been used with multiple sclerosis and multiple organ transplantation.10 Recently, FTY720 was found to inhibit the growth of various tumors, including HCC tumors.40,41 Because of its nontoxicity, specific cytotoxicity toward tumors and high oral bioavailability, FTY720 has great potential for tumor therapy. However, after exploring the use of FTY720 in various cancers, studies have shown that this drug substantially induces autophagy in tumor cells, which protects them from damage and reduces apoptosis.10–12 Autophagy is a process of cell self-degradation that plays an important role in adapting to metabolic stress and maintaining genomic integrity and internal stability.42 When tumor cells experience prolonged starvation, hypoxia, radiotherapy or chemotherapeutics, they induce autophagy to reduce cellular damage and maintain cell stability. In some anticancer drug therapies, autophagy reduces apoptosis and promotes tumor resistance.

Recently, many studies have shown that inhibiting autophagy effectively enhances the cytotoxicity of FTY720.10,15
Other studies have found that suppressing Beclin 1 protein expression, which inhibits autophagy,\textsuperscript{43} with siRNA, substantially improves FTY720 sensitivity in cancer cells. In addition, an increasing number of studies have shown that the combined use of siRNA and anticancer drugs effectively reverses multidrug resistance and inhibits tumor growth. However, one of the major difficulties in combining drugs and siRNA therapy is targeting the tumor tissue. To solve the problem, studies have focused on using liposomal vehicles to co-deliver siRNA and anticancer drugs.\textsuperscript{28–30} For example, the co-delivery of doxorubicin and Bmil-siRNA with FA-modified cationic liposomes significantly enhanced tumor accumulation and improved the antitumor effect.\textsuperscript{44} Additionally, Yang et al designed tumor-targeting cationic liposomes modified with dual peptides for co-delivery of siRNA and doxorubicin to gliomas.\textsuperscript{45} In summary, combining siRNA and anticancer drugs into the same vector has unique advantages.

In our study, we tested the ability of a novel siRNA delivery vector, LCP-II NPs, to co-deliver Beclin 1 siRNA and FTY720, and we demonstrated the protective role of FTY720-induced autophagy in reducing apoptosis. The results revealed that free FTY720 and LCP-II-siRNA-FTY720 NPs induced cell death and that the LCP-II-siRNA-FTY720 NPs were more effective than the free drug. As previously shown, free FTY720 and LCP-II-siRNA-FTY720 NPs also induced cell autophagy. After introduction of Beclin 1 siRNA, the Beclin 1 protein expression was effectively reduced, leading to autophagy inhibition. A series of experiments, such as confocal microscopy of GFP-LC3–transfected SMMC-7721 cells and Western blot analysis of the LC3-II to LC3-I ratio, confirmed the role of siBeclin 1 in LCP-II-siBeclin 1-FTY720 NPs. The results showed that after treatment with LCP-II-siBeclin 1-FTY720 NPs, the proportion of SMMC-7721 cells forming LC3 puncta and the ratio of LC3-II to LC3-I were significantly decreased, indicating that autophagy was suppressed by the co-delivery NPs. In addition, after treatment with siBeclin 1 and FTY720 through the co-delivery LCP-II NPs, cell viability was significantly decreased. Furthermore, the ratio of AV-positive and PI-positive cells increased by 50% after treatment with the LCP-II-siBeclin 1-FTY720 NPs for 24 h, and the expression levels of cleaved caspase-3, caspase-9 and PARP were also increased compared with the levels in cells treated with free FTY720 and LCP-II-siRNA-FTY720 NPs. Consistent with the in vitro experiments, co-delivery of siBeclin 1 and FTY720 in vivo exhibited the most powerful antitumor efficacy. Tumor weight and volume were significantly smaller in the LCP-II-siBeclin 1-FTY720 NP–treated group than in the other groups. However, without FA targeting, the cytotoxicity of LCP-II-siBeclin 1-FTY720 NPs was substantially decreased, indicating that this ligand was essential for targeted delivery to the tumor site. Consequently, blocking autophagy enhanced the cytotoxicity of FTY720. Autophagy is most likely an important factor in HCC resistance to FTY720. Combined treatment with siBeclin 1 and FTY720 using LCP-II NPs may effectively promote the antitumor efficacy of FTY720 and has great potential for HCC treatment.

Conclusion
LCP-II NPs effectively encapsulate siRNA and anticancer drugs, resulting in particles with a suitable average size that are internalized by SMMC-7721 cells. Treatment with these NPs silences Beclin 1 expression, blocking the autophagy induced by FTY720. Compared with free FTY720 and LCP-II-siRNA-FTY720 NPs alone, combined delivery of siBeclin 1 and FTY720 more efficiently inhibited HCC cell progression by suppressing protective autophagy and increasing apoptosis. This is the first time we have used LCP-II NPs as a co-delivery system for a small molecule and siRNA, and we found that this system has favorable synergistic effects for cancer treatment. Combining gene therapy and anticancer drugs in a nanocarrier is a popular focus of nonsurgical tumor treatment.

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Authors contributions
All authors contributed toward data analysis, drafting and revising the paper and agree to be accountable for all aspects of the work. JYW and ZWX contributed equally to this work.
Disclosure

The authors report no competing interests in this work.

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