Single-walled carbon nanohorn (SWNH) aggregates inhibited proliferation of human liver cell lines and promoted apoptosis, especially for hepatoma cell lines

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Abstract: Single-walled carbon nanohorns (SWNHs) may be useful as carriers for anticancer drugs due to their particular structure. However, the interactions between the material itself and cancerous or normal cells have seldom been studied. To address this problem, the effects of raw SWNH material on the biological functions of human liver cell lines were studied. Our results showed that unmodified SWNHs inhibited mitotic entry, growth, and proliferation of human liver cell lines and promoted their apoptosis, especially in hepatoma cell lines. Individual spherical SWNH particles were found inside the nuclei of human hepatoma HepG2 cells and the lysosomes of normal human liver L02 cells, implying that SWNH particles could penetrate into human liver cells and the different interacted mechanisms on human normal cell lines compared to hepatoma cell lines. Further research on the mechanisms and application in treatment of hepatocellular carcinoma with SWNHs is needed.

Keywords: single-walled carbon nanohorns, SWNHs, cell proliferation, cell apoptosis, hepatocellular carcinoma, nucleus, lysosome

Introduction

In 1999, Iijima et al1 first synthesized single-walled carbon nanohorns (SWNHs). Due to their large surface area, their particular surface structures, and especially their affinity for biomolecules, SWNHs may offer biomedical and pharmaceutical potential.2-3 Some researchers4 applied oxidized SWNHs (SWNHox) as carriers of the anticancer drug cisplatin and found that cisplatin released slowly from the SWNHs into aqueous solution and effectively inhibited the growth of human lung cancer NCI-H460 cells. They also found that as a carrier material, SWNHox itself did not promote or suppress the growth of lung cancer cells. Moreover, the roles of unmodified SWNHs on cells were not very clear up to now.5 Many scholars who have examined the biological functions between carbon nanotubes (CNTs), graphene, fullerene, and cells have found that carbon nanoparticles can enter into cells or subcellular organelles, such as nuclei and lysosomes.6-9 Carbon nanoparticles could induce apoptosis through oxidative stress.10-12

As a type of critical target cell line for the study of liver cancer therapy, HepG2 has been investigated by many researchers.13-15 How SWNHs affect the biological functions of human hepatoma cell lines such as HepG2 or normal liver cell lines such as L02 was unknown, and there were no reports on the differences in roles and mechanisms between the material and the normal or cancer cells. To address these
issues, the direct effects of SWNHs on human liver cell lines (human liver normal cell line L02 and human hepatoma cell line HepG2) were studied. We also investigated the internalization and localization of SWNH particles in liver cell lines. The remarkably different role on liver normal cell lines or hepatoma cell lines, induced by SWNHs, will be conducive for clinical treatment of hepatocellular carcinoma (HCC).

Materials and methods

Characterization of SWNHs

The SWNH material was synthesized by arc-discharge method in air and characterized by Raman spectra, thermogravimetry, and high-resolution transmission electron microscopy (HRTEM), as previously reported27 (partial figures of characterization of SWNHs are shown in Figure S1). The material is spherical aggregates of carbon nanohorns with dahlia-like structures; the diameters of aggregates ranged from 60–100 nm. The material dried at 100°C in air for 3 hours. The rapid N tube (Elementar Analysensysteme GmbH, Hanau, Germany) was used to detect the elemental contents of material. An S4 Explorer X-ray fluorescence spectrometer (Bruker AXS GmbH, Karlsruhe, Germany) was used to determine other elemental contents.

An ASAP 2010 V3.02E surface-area analyzer (Micromeritics Instrument Corporation, Norcross, GA, USA) was used to detect the surface area and mesopore size of SWNHs with Brunauer–Emmett–Teller (BET) theory, which aims to explain the physical adsorption of gas molecules on a solid surface and serves as the basis for an important analysis technique for the measurement of the specific surface area of a material. Particle density of materials was detected by AccuPyc 1330 Pycnometer (Micromeritics Instrument Corporation, Norcross, GA, USA) at 18°C using high-pressure helium (He) buoyancy effect. The particle size of SWNHs (10 µg/mL aqueous suspension) was detected by Zetasizer Nano ZS (Malvern Instruments, Malvern, UK) by means of dynamic light scattering at 25°C.24

Preparation of dishes coated with SWNHs

The dilute solution of SWNHs was suspended in ultrapure water from a Milli-Q Integral Water Purification System (EMD Millipore, Billerica, MA, USA). The dispersed SWNHs (10 µg/mL) were added to polystyrene (PS) dishes and then dried at 100°C for 2 hours in air, sterilized by ultraviolet irradiation for half an hour. The abbreviations represent SWNHs10 (0.21 µg/cm²), SWNHs20 (0.42 µg/cm²), SWNHs30 (0.64 µg/cm²), and SWNHs40 (0.85 µg/cm²), corresponding to 6 µg, 12 µg, 18 µg, and 24 µg in each 60 mm dish, respectively.24

The contact angles of water droplets (volume 2 µL) on SWNHs40-coated and noncoated PS surfaces were determined on the Dataphysics OCa 20 Contact angle Measuring System (DataPhysics Instruments gmbH, Filderstadt, Germany) at 20°C.

Cell culture

L02 and HepG2 cells were seeded onto PS dishes coated with SWNHs and were cultured with Dulbecco’s Modified Eagle’s Medium supplemented with 10% fetal bovine serum and 1% antibiotic (penicillin-streptomycin solution), and in 5% CO₂ at 37°C. There were clear similarities between the results from Chang liver cells and L02 cells, as well as those from Huh7 cells and HepG2 cells.24

Morphology and growth curve of cells cultured onto SWNH-coated dishes

L02 (3×10⁴) and HepG2 (3×10⁴) cells were added into dishes and cultured for 48 hours, respectively. The cells were subsequently observed by optical microscope, according to the general protocol. The cells were visualized and digital images were acquired using a Nikon camera (Nikon Corporation, Tokyo, Japan). The total numbers of cells at 48 hours were counted using a cell counter plate. The polynomial fitting for the relationship between number of cells (CN) and quantities (µg) of SWNHs (NH) was performed using Origin 8 software (OriginLab Corporation, Northampton, MA, USA).

Measurement of cell mitotic entry

Cells were synchronized by double thymidine block; the cells were incubated on SWNH-coated dishes for 48 hours, incubated with DNA–lipid mixture for 3 hours, washed twice, and incubated in fresh medium for an additional 5 hours. Subsequently, they were further cultured in a medium containing 2 mM thymidine and 2 µg/mL puromycin for the second arrest and drug selection. After 16 hours incubation, cells were released into the cell cycle by incubating in fresh medium. Finally, they were collected or fixed at indicated time points and subjected to specific analyses. DNA synthesis of cells was evaluated by bromodeoxyuridine (BrdU) labeling. After release from the block at each time point, cells grown in a 12-well plate were tagged with BrdU. After a further wash with phosphate-buffered saline (PBS), cells and BrdU-positive cells were counted. Mitotic events were evaluated with Openlab software (PerkinElmer, Waltham, MA, USA) by time-lapse videomicroscopy.38
The analysis of cell growth and proliferation

XTT (2,3-bis-(2-methoxy-4-nitro-5-sulfophenyl)-2H-tetrazolium-5-carboxanilide) assay (Roche Diagnostics Deutschland GmbH, Mannheim, Germany) was used to assess the cell growth, and DNA synthesis was evaluated with BrdU. After treatment for 48 hours, the cells of each group were reseeded onto 96-well plates with $3 \times 10^3$ cells per well, then counted with a plate reader at a wavelength of 450 nm.\(^\text{39}\)

Cell proliferation and viability assay

Cell Counting Kit-8 (CCK-8; Dojindo Laboratories, Kumamoto, Japan) was used to determine cell viability. Cells were pretreated using the above methods and then incubated in 96-well plates for 16 hours. In each well, 10 µL CCK-8 solution was added. Cells were incubated at $37^\circ\text{C}$ for 4 hours and absorbance values were measured with a plate reader at 450 nm.\(^\text{40}\)

Cell cycle analysis

Cells ($3 \times 10^5$) were cultured on noncoated and SWNH-coated dishes for 48 hours. After that, the cells were fixed and stained with propidium iodide (Sigma-Aldrich, St Louis, MO, USA). Cell cycle was determined by flow cytometry (BD FACSCalibur; BD Biosciences, San Jose, CA, USA).

Determination of apoptosis

Apoptotic cells were assessed with Alexa Fluor* 647 Annexin V (BioLegend, Inc, San Diego, CA, USA). Cultured cells were washed at $4^\circ\text{C}$ for 30 minutes in PBS and then stained with Annexin-V staining solution (Annexin-V-FLUOS Staining Kit; Roche Diagnostics Corporation, Indianapolis, IN, USA) at $4^\circ\text{C}$ for 3 hours. Gels were washed four times in PBS at $4^\circ\text{C}$ and fixed at room temperature with 1% paraformaldehyde (Sigma-Aldrich) in PBS for 15 minutes. For counterstaining, 7-Aminoactinomycin D (7-AAD; BD Biosciences) 2 µg/mL was added to the first washing step. The numbers of total, Annexin-V-positive, 7-AAD–positive, and double-positive cells were determined respectively. Apoptosis was verified by detection of activated caspases.\(^\text{51}\)

Antibodies and immunoblotting

Cells were lysed with buffer containing the cocktail of protease inhibitors. Equal amounts of protein were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis and transferred to a polyvinylidene difluoride membrane. After blocking with 5% nonfat dry milk, the membranes were reacted with primary antibodies. The following antibodies at indicated dilutions were used for immunoblotting: Silent information regulator 1 (SIRT1) and p53 antibodies (Santa Cruz Biotechnology Inc, Dallas, TX, USA; 1:1,000), β-actin (Cell Signaling Technology, Inc, Danvers, MA, USA; 1:1,000), and caspase-3 and caspase-7 (Santa Cruz Biotechnology Inc; 1:1,000). All affinity-purified and species-specific fluorophore-conjugated secondary antibodies were obtained from Santa Cruz Biotechnology Inc and used at dilutions between 1:500 and 1:800.

Cellular morphology observed by confocal microscope

Cells ($1 \times 10^5$) were grown on confocal dishes (10 mm), coated with or without SWNHs for 48 hours; they were subsequently treated with 4% paraformaldehyde for 10 minutes, washed three times with PBS, stained at $30^\circ\text{C}$, for 30 minutes with PBS containing 0.1 µg/mL 4′,6-diamidino-2-phenylindole. Confocal images were acquired using the Zeiss 510 META confocal microscope (Carl Zeiss Microscopy GmbH, Jena, Germany).

Cells observed by transmission electron microscope

Cells ($3 \times 10^5$) were cultured on non-coated and SWNHs-coated (0.85 µg/cm²) PS dishes, for 48 hours. The cells were collected and fixed. Then, samples were detected with a JEM-1400 transmission electron microscope (TEM) (JEOL, Tokyo, Japan).\(^\text{24}\)

Statistics

All data were presented as mean ± standard deviation (SD) from at least three independent experiments. For statistical comparison of quantitative data in the group, analysis of variance or Student’s t-test was performed. To determine differences between groups not normally distributed, medians were compared using Kruskal–Wallis analysis of variance. Significance was determined using the paired Student’s t-test for the mean of three different experiments. A P-value $<$0.05 was considered significant.

Results

Assessment of SWNHs

Elemental composition of SWNHs is shown in Table S1. It included 95.3% carbon and 0.25% total metal content (each metal less than 0.1%). The adsorptive isotherm plot of the SWNH material was similar to type III adsorptive isotherm.
curve (Figure S2), which indicated that SWNHs particles had a hydrophobic surface. The result showed that the surface area of BET was 631.55 m²/g, and larger than surface area reported by other research.²

The diameter was less than 308.7 nm at the corresponding relative pressure (P/P₀) 0.994, and the pore volume was 1.57 cm³/g. The single point total average pore diameter was 9.97 nm. The Barrett-Joyner-Halenda (BJH) adsorption pore size distribution demonstrated that most mesopores in the SWNH material were 2–8 nm in diameter (Figure S3). The particle density of SWNHs was 1.0077 g/cm³. The results indicated that SWNHs had many closed pores. The size of SWNH particles ranged from 295–615 nm suspended in aqueous solution (Figure S4).

Characterization of SWNH-coated dishes
On the surface of PS dishes, the scanning electron microscope (SEM) images exhibited single particles of SWNHs and their 60–100 nm diameters (Figure 1) (SEM measurements were carried out using a SIRION field emission scanning electronic microscope; FEI Corporation Ltd., Hillsboro, OR, USA). The secondary SWNH aggregates were dispersed in individual particles on the surface of dishes, which may have come from Λ-Λ stacking interactions between the benzene rings on the surface of PS and SWNHs, rather than those between SWNH aggregates.

On the surface of dishes, contact angle was 44.9° of water droplet (Figure S5A), less than that on the surface coated with dried SWNHs; 74.5° (Figure S5B); the hydrophobicity of SWNHs40/PS surface was higher than that of uncoated PS surface.

Morphology of liver cells observed by optical microscope
The morphology of liver cells cultured onto noncoated and SWNH-coated dishes under standard culture conditions for 48 hours was observed with an optical microscope. The results show that with increasing quantities of SWNHs on the dishes, the total counts of L02 cells (Figure 2A–C) and HepG2 cells (Figure 2D–F) decreased in a dose-dependent manner, the liver cells became smaller in size, and many more spherical cells were observed; the phenomena were more marked in the HepG2 cells. As a proliferation inhibitor, SWNHs had a much greater effect on the hepatoma cells than on the normal cells.

Role of SWNHs in inhibiting cell growth, proliferation, and mitotic entry of liver cells
Growth curves for L02 and HepG2 cells with increasing quantities of SWNHs are shown at 48 hours; both the numbers of L02 cells (Figure 2G, lower curve) and HepG2 cells (Figure 2H, upper curve) decreased significantly in a dose-dependent manner (P<0.01). Moreover, polynomial fitting for the relationships between CN and NH was performed. Although the HepG2 cells proliferated faster than the L02 cells did with the increase in NH, proliferation of HepG2 cells was significantly slowed down and gradually approached the rate of the L02 cells. The polynomial fitting formula for L02 (Figure 2G, lower dotted curve) was as follows:

\[
[CN] = -26.2 [NH]^4 + 1.23 \times 10^3 [NH]^3 - 1.29 \times 10^4 [NH]^2 - 1.20 \times 10^5 [NH] + 1.96 \times 10^6, R^2 = 1.0.
\]

The polynomial fitting formula for HepG2 (Figure 2G, upper dotted curve) was as follows:

\[
[CN] = -1.46 \times 10^2 [NH]^4 + 7.42 \times 10^3 [NH]^3 - 1.19 \times 10^4 [NH]^2 + 4.56 \times 10^5 [NH] + 4.33 \times 10^6, R^2 = 1.0.
\]

Both the relationships between the number of L02 or HepG2 cells and the quantities of SWNHs are quartic polynomial.

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**Figure 1** The films of SWNHs40/PS observed by scanning electron microscope.

**Notes:** PS dishes with a surface area of about 1 cm² and coated with SWNHs40 (0.85 µg/cm²) were prepared for SEM measurements. After being pretreated, by spraying gold on films of samples, SEM measurements were carried out using a SIRION field emission scanning electronic microscope (FEI Corporation Ltd., Hillsboro, OR, USA) with accelerating voltage of 10.0 kV. (A) 50,000x, scale bar represents 1 µm. (B) 100,000x, scale bar represents 500 nm. (C) 200,000x, scale bar represents 200 nm.

**Abbreviations:** SWNH, single-walled carbon nanohorn; PS, polystyrene; SEM, scanning electron microscope.
The first-order derivatives $d [CN]/d [NH]$ of the two formulas above were also deduced. The first-order derivative for polynomial fitting formula of L02 (Figure 2H, upper curve) was as follows:

$$d [CN]/d [NH] = -1.048 \times 10^7 [NH]^3 + 3.69 \times 10^5 [NH]^2 - 2.58 \times 10^4 [NH] - 1.20 \times 10^5 \text{ (3)}$$

The first-order derivative for polynomial fitting formula of HepG2 (Figure 2H, lower curve) was as follows:

$$d [CN]/d [NH] = -5.84 \times 10^2 [NH]^3 + 2.23 \times 10^4 [NH]^2 - 2.38 \times 10^3 [NH] + 4.56 \times 10^4 \text{ (4)}$$

First-order derivative indicated that the rate of change in cell numbers followed with the dose of SWNHs. The formula of the rate of change was a negative value, which revealed that cell proliferation was inhibited by SWNHs. Furthermore, the low rate of change in HepG2 cells compared with L02 cells demonstrated that SWNHs had a more marked inhibitory effect on hepatoma cell lines than on normal cell lines (Figure 2H).

To study the influence of SWNHs on cell mitosis, we mixed BrdU into cells. The results showed that mitotic entry of L02 and HepG2 cells was delayed significantly by SWNHs (Figure 3A and B) and that mitotic entry of cells was inhibited (Figure 3C and D) in a time- and dose-dependent manner ($P<0.01$). The effects on the delay of cell mitosis and that of SWNHs inhibited mitotic entrance into the cells were more significant for HepG2 cells than for L02 cells.

The effect on cell growth by SWNHs was studied using XTT assay. The results showed that both L02 cells (Figure 4A) and HepG2 cells (Figure 4B) were significantly inhibited in a time- and dose-dependent manner ($P<0.001$). The results of cell viability assayed with CCK-8 also indicated that SWNHs inhibited cell proliferation of L02 cells (Figure 4C) and HepG2 cells (Figure 4D) and was time- and dose-dependent.

**Figure 2.** Morphology and growth curves of L02 and HepG2 cells cultured onto noncoated and SWNH-coated dishes.

**Notes:** L02 cells (3x10^4) and HepG2 cells (3x10^4) were seeded onto 60 mm noncoated and SWNH-coated dishes, respectively, and cultured for 48 hours, and then images of the cells were observed by optical microscope according to the general protocol and the total numbers of L02 and HepG2 cells were counted. The cells were visualized and digital images were acquired using Nikon camera (Nikon Corporation, Tokyo, Japan). Magnification 400x. (A) Control, L02 not treated with SWNHs. (B) L02 treated with SWNHs20 (0.42 µg/cm²). (C) L02 treated with SWNHs40 (0.85 µg/cm²). (D) Control, HepG2 untreated with SWNHs. (E) HepG2 treated with SWNHs20 (0.42 µg/cm²). (F) HepG2 treated with SWNHs40 (0.85 µg/cm²). (G) Cell growth curves for L02 (lower curve) and HepG2 (upper curve). All data are represented as mean ± SEM. The polynomial fitting formulas for the relationship between CN and NH were performed using Origin 8 software (OriginLab Corporation, Northampton, MA, USA) (dotted curves). (H) First-order derivatives $d [CN]/d [NH]$ deduced from the two polynomial fitting formulas.

**Abbreviations:** L02, normal liver cell line; HepG2, human hepatoma cell line; SWNH, single-walled carbon nanohorns; SWNHs20, 0.42 µg of single-walled carbon nanohorns per cm², or 12 µg per 60 mm dish; SWNHs40, 0.85 µg of single-walled carbon nanohorns per cm², or 24 µg per 60 mm dish; CN, number of cells at 48 hours; NH, quantity (µg) of single-walled carbon nanohorns coated on culture dish; SEM, standard error of the mean.
SWNHs affected liver cell cycle

The role on cell cycle by the material was measured with flow cytometry. The results showed that the effect of SWNHs on the cell cycles of L02 (Figure 5A) and HepG2 (Figure 5B) was dose-dependent.

G1 phase increased and S phase decreased significantly in an SWNHs’ dose-dependent manner, in L02 cells (P<0.05) and HepG2 cells (P<0.01). But, G2 phase decreased in L02 and HepG2 cells; except it increased abruptly in HepG2 cells treated with SWNHs40 (P<0.05). The difference in G1 phase delay induced by SWNHs between L02 and HepG2 cells indicated that SWNHs had different cytotoxic effects on the two kinds of cells and led to much stronger inhibition of DNA replication in HepG2 cells than in L02 cells. The liver cells were blocked in G2 phase by SWNHs, and the gap between DNA synthesis and mitosis was delayed by the quantity of SWNHs40 in HepG2, which may result from the apoptosis of HepG2 cells. The cell cycle of liver cell lines was associated with the suppression of their proliferation induced by SWNHs.

SWNHs induced liver cell apoptosis

The cells were treated with SWNHs for 48 hours, and apoptotic cells were detected with Annexin-V. The results showed that apoptosis of L02 (Figure 6A) and HepG2 (Figure 6B) cells was improved in a SWNHs dose-dependent manner, especially for HepG2 cells (P<0.001).

Apoptosis is the process of programmed cell death that may occur in multicellular organisms. Biochemical events lead to characteristic cell changes (morphology) and death. These changes include blebbing, cell shrinkage, nuclear fragmentation, chromatin condensation, and chromosomal DNA fragmentation. Apoptosis of mammalian animal cells is attuned by p53 and SIRT1, among others. The group of cysteine...
proteases are the caspases, and they are key factors of cell apoptosis, especially for caspase-3 and caspase-7. The results of our work indicated that the expression of caspase-3, caspase-7, and p53 of L02 cells (Figure 6C) and HepG2 cells (Figure 6D) increased in a SWNH dose-dependent manner. However, the expression level of activation cleavage SIRT1 decreased and was dose-dependent on SWNHs, confirming that apoptosis of liver cells was advanced by SWNHs.

**Figure 4** SWNHs inhibited growth and proliferation of liver cells.

Notes: L02 cells (3\times 10^{4}) and HepG2 cells (3\times 10^{4}) were seeded onto 60 mm noncoated and SWNH-coated dishes, respectively, and cultured for 48 hours. Then the effects of SWNHs on the growth of L02 (A) and HepG2 (B) cells were investigated by XTT assays. L02 (Figure 4C, 3\times 10^{4}) and HepG2 (Figure 4D, 3\times 10^{4}) cells were cultured in 96-well plates (6 mm) treated with or without SWNHs, and the cell viabilities were evaluated by CCK-8 assay. (A) The XTT value of L02 cells treated with SWNHs decreased significantly from 36 hours after culture (P<0.05), especially after 48 hours (P<0.01). (B) The XTT value of HepG2 cells treated with SWNHs decreased significantly after 36 hours and 48 hours (P<0.01). (C) The absorbance of L02 cells decreased significantly from 24 hours and 36 hours after culture with SWNHs, and especially after 48 hours (P<0.01). (D) The absorbance of HepG2 cells decreased significantly from 24 hours after culture with SWNHs (P<0.05), and especially after 36 hours and 48 hours (P<0.01). All data are represented as mean ± SEM. *P<0.05; **P<0.01.

Abbreviations: SWNH, single-walled carbon nanohorn; XTT, 2,3-bis-(2-methoxy-4-nitro-5-sulfophenyl)-2H-tetrazolium-5-carboxanilide; L02, normal liver cell line; BrdU, bromodeoxyuridine; SWNHs10, 0.21 µg of single-walled carbon nanohorns per cm²; SWNHs20, 0.42 µg of single-walled carbon nanohorns per cm²; SWNHs30, 0.64 µg of single-walled carbon nanohorns per cm²; SWNHs40, 0.85 µg of single-walled carbon nanohorns per cm², or 24 µg per 60 mm dish; HepG2, human hepatoma cell line; CCK-8, Cell Counting Kit-8; SEM, standard error of the mean.

**Figure 5** SWNHs affected cell cycle of liver cells.

Notes: L02 cells (3\times 10^{4}) and HepG2 cells (3\times 10^{4}) were seeded onto noncoated and SWNH-coated 60 mm dishes, respectively, and cultured for 48 hours. Then the effects of SWNHs on L02 and HepG2 cell cycles were measured by flow cytometry. (A) G1 phase increased and S phase decreased significantly in an SWNH dose-dependent manner in L02 cells (P<0.05); G2 phase decreased in L02 cells. (B) G1 phase increased and S phase decreased significantly in an SWNH dose-dependent manner in HepG2 cells (P<0.01); G2 phase decreased in HepG2 cells, except it increased abruptly in HepG2 cells treated with SWNHs40 (P<0.05). All data are represented as mean ± SEM. *P<0.05; **P<0.01.

Abbreviations: SWNH, single-walled carbon nanohorn; L02, normal liver cell line; SWNHs10, 0.21 µg of single-walled carbon nanohorns per cm²; or 6 µg per 60 mm dish; SWNHs20, 0.42 µg of single-walled carbon nanohorns per cm²; or 12 µg per 60 mm dish; SWNHs30, 0.64 µg of single-walled carbon nanohorns per cm²; or 18 µg per 60 mm dish; SWNHs40, 0.85 µg of single-walled carbon nanohorns per cm², or 24 µg per 60 mm dish; HepG2, human hepatoma cell line; SEM, standard error of the mean.
Figure 6 SWNHs promoted cell apoptosis of liver cells, and apoptosis involves key factors in vivo.

Notes: L02 cells (1×10⁷) and HepG2 cells (3×10⁷) were seeded onto 60 mm noncoated and SWNH-coated dishes, respectively, and cultured for 48 hours. Then the effect of SWNHs on L02 (A) and HepG2 (B) cell apoptosis distribution was determined by flow cytometry. The expression levels of SIRT1 and those of activation cleavage of p53, caspase-3 and caspase-7 L02 (C) and HepG2 (D) were determined by Western Blotting. (A) The apoptotic L02 cells treated with SWNHs increased significantly at 48 hours after culture with SWNHs10 (P<0.05), and especially with SWNHs20, SWNHs30, and SWNHs40 (P<0.01). (B) The apoptotic HepG2 cells treated with SWNHs increased significantly at 48 hours after culture with SWNHs10 (P<0.05), and especially with SWNHs20, SWNHs30, and SWNHs40 (P<0.01). All data are represented as mean ± SEM. *P<0.05; **P<0.01.

Abbreviations: SWNH, single-walled carbon nanohorn; L02, normal liver cell line; HepG2, human hepatoma cell line; SEM, standard error of the mean; SIRT1, silent information regulator 1.

Figure 7 Morphology of L02 and HepG2 cells cultured onto SWNH-coated dishes observed by confocal microscope.

Notes: L02 cells (1×10⁷) and HepG2 cells (1×10⁷) were seeded onto 10 mm noncoated and SWNH-coated dishes, respectively, and cultured in Dulbecco’s Modified Eagle’s Medium with fetal bovine serum and penicillin–streptomycin solution at 37°C in humidified 5% CO₂/95% air for 48 hours. Then L02 and HepG2 cells were treated with paraformaldehyde and 4',6-diamidino-2-phenylindole. Confocal images were acquired by Zeiss 510 META confocal microscope (Carl Zeiss Microscopy GmbH, Jena, Germany). (A) Control, L02 not treated with SWNHs. (B) L02 treated with SWNHs20 (0.42 µg/cm²). (C) L02 treated with SWNHs40 (0.85 µg/cm²). (D) Control, HepG2 untreated with SWNHs. (E) HepG2 treated with SWNHs20 (0.42 µg/cm²). (F) HepG2 treated with SWNHs40 (0.85 µg/cm²). Scale bars represent 10 µm. The white arrows show the nuclei of cells.

Abbreviations: L02, normal liver cell line; HepG2, human hepatoma cell line; SWNH, single-walled carbon nanohorn; SWNHs20, 0.42 µg of single-walled carbon nanohorns per cm²; SWNHs40, 0.85 µg of single-walled carbon nanohorns per cm²; 12 µg per 60 mm dish; SWNHs30, 0.64 µg of single-walled carbon nanohorns per cm²; 18 µg per 60 mm dish; SWNHs10, 0.21 µg of single-walled carbon nanohorns per cm²; 6 µg per 60 mm dish; 0.05; **P<0.01). all data are represented as mean ± SEM. *P<0.05; **P<0.01.
Cellular morphology observed by confocal microscope

The confocal morphology of L02 cells showed that the shapes of their nuclei were similar to those of the control cells (Figure 7A), and the counts of cells decreased in an SWNH dose-dependent manner. The sizes of L02 cell nuclei treated with SWNHs were slightly larger than those of control cells (Figure 7B and C); the nuclei swelled in a dose-dependent manner.

The counts of viable HepG2 cells decreased significantly and those of apoptotic HepG2 cells increased in an SWNH dose-dependent manner. Apoptotic HepG2 cells can be observed in Figure 7F (indicated by arrow), but not in controls (Figure 7D). The images of HepG2 showed that the sizes of nuclei in HepG2 cells treated with SWNHs40 were much smaller than those of controls. This phenomenon may be explained by karyopyknosis (Figure 7F, indicated by arrows). As proposed by Romero et al,23 the expansive nuclei and karyopyknosis induced by SWNHs could be attributed to the fact that the protein content in nuclei increased in earlier apoptosis and decreased in late apoptosis, respectively. It could explain the abruptly increased ratio of G2 phase of HepG2 cells treated with SWNHs 40 (Figure 5B). Almost all HepG2 cells treated with SWNHs 40 were apoptotic, and the ratio of S phase in HepG2 cells decreased significantly when the cells were blocked at the G2/M phase.

The apoptosis effects on HepG2 cells induced by SWNHs were more significant than those on L02 cells. The cells treated with SWNHs40 showed typical apoptotic morphology; however, in the case of L02 cells, the apoptosis of cells was not so obvious. The apoptotic cells had typical features, which included chromatin condensation, cell shrinkage, scant cytoplasm, and membrane blebbing, especially for apoptotic body.43–45 These phenomena could be observed in L02 and HepG2 cells treated with SWNHs, and were more apparent in HepG2 cells.

Cells observed by TEM

Liver cells were cultured in dishes treated with SWNHs40 (0.85 µg/cm²) for 48 hours and then were collected for TEM measurement. SWNHs could be observed in cells by TEM. SWNHs aggregate was smaller than 100 nm, and localized at lysosomes of L02 (Figure 8B and C) and nuclei of HepG2 cells (Figure 8E and F), respectively. The untreated cells are shown in Figure 8A (L02) and Figure 8D (HepG2). The individual SWNH particles were easily distinguished from their neighbor molecules because of higher electronic densities on the surface of SWNHs than those of other molecules in organelles.

The TEM observation of organelles showed that both lysosomes and mitochondria of L02 cells (Figure 8G) and HepG2 cells (Figure 8H) treated with SWNHs were more expansive than those of controls, respectively, and the phenomena were more significant in HepG2 cells. There were manifold secretory vesicles outside the L02 cells (Figure 8G) and HepG2 cells (Figure 8H) treated with SWNHs, but these could not be observed outside the control cells. The secretory vesicles outside HepG2 cells (Figure 8H) treated with SWNHs were larger than those outside L02 cells (Figure 8G). SWNHs are heterogeneous materials; the response of cells was secreted ingested SWNHs. The more cells ingested SWNHs, the more there were secretory vesicles; the HepG2 cells may have ingested many more SWNHs than L02 cells did. In addition, SWNH aggregates could be found in cytoplasm by TEM.

Discussion

Previous works reported that some carbon nanomaterials inhibited proliferation of L02 or HepG2 cells. Fullerene C60 inhibited HepG2 proliferation at a dosage as low as 0.46 µg/mL.29 Liu et al demonstrated that unmodified multiwalled nanotubes (MWNTs), carboxyl modified MWNTs and hydroxyl modified MWNTs decreased the viability of L02 cells at dosages of 12.5–200.0 µg/mL in a dose-dependent manner. Romero et al reported oxidized or PSPM (poly [sulfopropyl methacrylate]), PAH (poly [allylamine hydrochloride]) and lipid-modified MWNTs inhibited proliferation of HepG2 at higher dosage (100 µg/mL). Single-walled carbon nanotubes (SWCNTs) are known to have great potential for biomedical applications such as photothermal ablation of tumor cells in combination with near-infrared irradiation. Hashida et al found that the photothermal activity of a novel SWCNT composite with a designed peptide having a repeated structure of H-(Lys-Phe-Lys-Ala)-7-OH ([KFKA]7) against tumor cells. The SWCNTs-[KFKA]7 composite demonstrated high aqueous dispersibility that enabled SWCNTs to be used in tumor ablation. This suggests the great potential of an SWCNT–peptide composite for use in photothermal cancer therapy.46 We obtained similar results using L02 and HepG2 cells treated with SWNHs at lower dosages (1.5–6.0 µg/mL, 4 mL of culture matrix).

Ajima et al showed that oxidized SWNHs (SWNHox) themselves did not promote or suppress the growth of lung cancer cells at dosages of 0–20 µg/mL, but their result in
Figure 8 TEM images of liver cells.

Notes: L02 cells (3×10⁵) and HepG2 cells (3×10⁵) were seeded onto 60 mm noncoated and SWNHs40-coated (0.85 µg/cm²) dishes, respectively, and cultured for 48 hours. The cells were collected and fixed with 3% glutaraldehyde. For TEM, ultra-thin cell slices of 100 nm thickness were cut using an ultramicrotome and mounted on grids. The slices were contrasted with aqueous solution of uranyl acetate and lead citrate, and examined with a JEM-1400 transmission electron microscope (JEOL Ltd, Tokyo, Japan) with accelerating voltage of 80 kV. (A) L02 not treated with SWNHs, as control (15,000×). Scale bar represents 1 µm. (B) L02 cultured onto SWNHs40-coated dishes (0.85 µg/cm²) for 48 hours (15,000×). Scale bar represents 1 µm. (C) L02 cultured onto SWNHs40-coated dishes (0.85 µg/cm²) for 48 hours (80,000×). Scale bar represents 100 nm (image was enlarged from the rectangular inset of panel B). The arrow shows that there were individual spherical SWNH particles with diameters less than 100 nm inside lysosomes of L02 cells. (D) HepG2 not treated with SWNHs, as control (15,000×). Scale bar represents 1 µm. (E) HepG2 cultured onto SWNHs40-coated dishes (0.85 µg/cm²) for 48 hours (15,000×). Scale bar represents 1 µm. (F) HepG2 cultured onto SWNHs40-coated dishes (0.85 µg/cm²) for 48 hours (80,000×). Scale bar represents 100 nm (image was enlarged from the rectangular inset of panel E). The arrow shows that there were individual spherical SWNH particles with diameters less than 100 nm inside nuclei of HepG2 cells. (G) L02 cultured onto SWNHs40-coated dishes (0.85 µg/cm²) for 48 hours (15,000×). Scale bar represents 1 µm. (H) HepG2 cells cultured onto SWNHs40-coated dishes (0.85 µg/cm²) for 48 hours (15,000×). Scale bar represents 1 µm.

Abbreviations: TEM, transmission electron microscope; L02, normal liver cell line; HepG2, human hepatoma cell line; SWNHs40, 0.85 µg of single-walled carbon nanohorns per cm², or 24 µg per 60 mm dish; SWNH, single-walled carbon nanohorn.

vivo showed that SWNHox itself had anticancer effects, which appeared to be contradictory to the result of cellular experiments. Fan et al showed that no obvious decrease of cell viability was observed for Hela cells incubated with gum arabic modified SWNHs (10–1000 µg/mL). This result suggested that the nanoparticles were nontoxic to Hela cells. Tahara et al reported recently that the proliferation of murine macrophage RAW 264.7 cells was inhibited by oxidized SWNHs coated with 1, 2-distearoyl-Sn-glycero-3-phosphoethanolamine-N-[amino (polyethylene, glycol) 2000] at higher dosages (10–300 µg/mL) in a dose-dependent manner. The modification of carbon nonmaterial with different chemical compounds could change various biological functions of cells.
As a powerful means for observation of micron and nanometer structures, TEM has been applied for the localization of nanoparticles in organelles. By means of TEM, oxidized SWNTs and unmodified and modified MWNTs have been found in cytoplasm, endosomes, or lysosomes of human macrophage cells. Porter et al\textsuperscript{18} observed that SWNTs dispersed in tetrahydrofuran could be localized within the lysosomes and nuclei of human macrophages; they also found fullerene $C_{60}$ in cytoplasm and nuclei of human macrophages.\textsuperscript{47} Besides being found in cytoplasm, endosomes, or lysosomes, MWNT-NH$_2$ was first discovered in nuclei of human embryonic kidney 293 cells in 2009.\textsuperscript{21} In 2011, oxidized or PSPM and lipid-modified MWNTs had been found in cytoplasm of HepG2 cells; however, the carbon nanoparticles had not been found in the nuclei of the cells from the images, despite the changes of DNA and protein in nuclei of HepG2 cells induced by modified MWNTs that had been acquired by confocal Raman microscope.\textsuperscript{23} Our results provide the first direct observation of SWNH particles in organelles of L02 and HepG2 cells.

Combining the results from confocal microscope and TEM, we assumed that there were different interactive mechanisms between SWNHs and L02 or HepG2 cells, respectively. SWNHs may directly react with nuclei of HepG2 cells and induce the apoptosis of the cells, whereas SWNHs may induce L02 cell apoptosis correlated to activation of lysosomal function.

Romero et al\textsuperscript{33} described the effects of the inhibited cell proliferation induced by apoptosis as “in vitro” toxicity of carbon nanotubes (CNTs) for HepG2 cells. Nevertheless, cytotoxicity for cancer cells may have therapeutic prospect. Our results showed that SWNHs were more toxic for HepG2 than for L02, and this prominent difference might be ascribed to selective affinity of SWNHs surface structure to subcellular structures in different cells. On the basis of these differences, new drugs or drug carriers with particular structures of SWNHs could be designed for clinical pharmaceutical therapy, as well as for cancer photothermal therapy.\textsuperscript{58–59} With the affinity of SWNHs to nuclei of HepG2, the prospect of designing cell-targeted drugs might be considered.

The fact that SWNHs themselves inhibited proliferation of HepG2 cells and promoted their apoptosis also reveals that when SWNH material is applied as an anti-HCC drug carrier, it may have a cooperative therapeutic role with anticancer drugs such as cisplatin carried by SWNHs.

The lysosomal compartment was the most common intracellular site of nanoparticle sequestration and degradation. Lysosomal dysfunctions and mitochondrial apoptosis result in toxicological consequences.\textsuperscript{51,52} According to the oxidative stress pathway for apoptosis induced by nanoparticle exposure,\textsuperscript{31} localization of nanoparticles may induce activation of lysosomal function, the “suicide bags” with high content of digestive enzyme, triggering mitochondrial apoptosis.\textsuperscript{34} Mitochondria, as the major source of reactive oxygen species (ROS) and redox machinery in cells, could activate inflammatory cytokines.\textsuperscript{32} Oxidative stress had been found in HepG2 cells treated with unmodified or oxidized SWNTs and MWNTs,\textsuperscript{25–28} L02 cells treated with unmodified, carboxyl-modified, or hydroxyl-modified MWNTs,\textsuperscript{29} and rat hepatocytes treated with fullerenes $C_{60}$, $C_{60}$ (OH)$_{12}$, and $C_{60}$ (OH)$_{24}$\textsuperscript{31} Gao et al\textsuperscript{19} found that modified MWNTs internalized in lysosomes of human macrophages (THP-1 cells), generated ROS, and lowered mitochondrial membrane potential. Tahara et al\textsuperscript{9} found that high accumulation of SWNHs induced lysosomal membrane destabilization and generated ROS in murine macrophage RAW 264.7 cells. Our results showed that individual SWNH particles were in lysosomes of L02 cells and mitochondria of L02 and HepG2 cells, suggesting that lysosomal dysfunctions and mitochondrial apoptosis of liver cells may play a key role in the cellular physiological pathways from SWNH exposure.

A model of MWNT transmembrane pathways divided MWNTs into two classes, clusters and singles. CNT clusters are taken up by cells through an energy-dependent endocytosis process, whereas highly dispersed single CNTs enter cells directly by penetrating cellular membranes.\textsuperscript{21} We are not sure which transmembrane pathway is adopted by SWNHs. Because about 2000 nanohorns were assembled to form one spherical SWNH aggregate, the aggregates would appear to be clusters of SWNHs.\textsuperscript{9} However, the aggregate was difficult to separate into individual nanohorns and was more like one single nanoparticle. Porter et al\textsuperscript{18} attributed the internalization of SWNTs inside cells and organelles to the smaller diameters of SWNTs (0.6–3.5 nm) rather than to nuclear pore complex and other pores in cells.\textsuperscript{18} A single primary SWNH aggregate has a larger diameter (60–100 nm) than that of single SWNTs (0.6–3.5 nm) and fullerene $C_{60}$ (0.7 nm). It is interesting how individual spherical SWNH aggregates with 2000 nanohorns internalize within cells and localize inside lysosomes of L02 cells and even enter into nuclei of HepG2 cells, as it was proposed previously that only nanoparticles less than 40 nm in diameter could enter into cell nuclei.\textsuperscript{53} The serum proteins of bovine used in cell culture and the low negative charge density on surfaces of SWNHs could be helpful for SWNH dispersion.
in culture fluid, or affect SWNHs recognized by receptors on cellular membranes and their transmembrane behavior.54

Many questions remain about this area of research. How could SWNHs enter into cells and go to different destinations (liposome or nuclei) in different cells? How is the apoptosis of different cells induced? Further research on the mechanisms and application in treatment of HCC with SWNHs should be carried out in animal models. We will use HCC mouse models to study the safety and anti-cancer role of SWNHs in detail.

**Conclusion**

In summary, we have for the first time demonstrated that small amounts of unmodified SWNHs could suppress cell proliferation in human liver cell lines, affect their cell cycle, and promote their apoptosis in a dose-dependent manner, especially in a hepatoma cell line. Apoptosis of liver cells induced by SWNHs was involved in caspase-3, caspase-7, p53, and SIRT1. During culture, individual SWNH particles could pass through the cellular membrane and enter the nuclei of HepG2 cells and the lysosomes of L02 cells. These discoveries imply different interaction mechanisms between L02 cells, HepG2 cells, and SWNHs.

In order to clarify the problems of SWNH transmembrane pathways and subcellular structure oriented interactive mechanisms between SWNHs and liver cells, it is evident that further research is needed on the basis of discoveries in this work.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**


Supplementary materials

Table S1  Elemental contents in single-walled carbon nanohorns

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Figure S1  Characterization of carbon nanohorns prepared in an atmosphere of air.
Notes: (A) Transmission electron microscopy (sample a detected) and high-resolution transmission electron microscopy (d); (B) Scanning electron microscopy; (C) Raman spectra; (D) Thermogravimetry. Upper curve: thermogravimetry curve; lower curve: decomposition thermogravimetry.
Abbreviations: AU, arbitrary unit; Deriv, derivation.

Figure S2  Adsorptive and desorptive isotherm plots of single-walled carbon nanohorns.
Abbreviations: SWNH, single-walled carbon nanohorn; STP, standard temperature; \( \frac{P}{P_0} \), the corresponding relative pressure.

Figure S3  BJH adsorption \( \frac{dV}{dD} \) – logD (nm) of single-walled carbon nanohorns.
Abbreviations: BJH, Barrett-Joyner-Halenda method; V, pore volume (cm\(^3\)/g); D, diameter (nm); SWNH, single-walled carbon nanohorn.
Effects of single-walled carbon nanohorns on human liver cells

Figure S5 Contact angles of water droplets on the surfaces of noncoated and SWNHs40-coated PS films.

Notes: Noncoated and SWNHs40-coated (0.85 µg/cm²) PS dishes with a surface area of about 1 cm² were prepared for contact angle measurements. The contact angles of water droplets (volume 2 µL) on SWNHs40-coated and noncoated PS surfaces were determined on the Dataphysics OCA 20 Contact Angle Measuring System (Dataphysics Instruments GmbH, Filderstadt, Germany) at 20°C. The figure shows the contact angles of water droplets on the surfaces of (A) noncoated PS, (B) SWNHs40-coated PS, and (C) SWNHs40-coated PS after a few minutes of contact with the water droplet.

Abbreviations: SWNHs40, 0.85 µg of single-walled carbon nanohorns per cm², or 24 µg per 60 mm dish; PS, polystyrene.

Figure S4 Single-walled carbon nanohorns particle size distribution in aqueous suspension.

Notes: The particle density was determined using the high pressure Helium buoyancy effect. This effect was measured gravimetrically by an electronic microbalance and pressure transducers. The filled pressure of Helium gas was 138.8 kPa. The particle intensity distribution showed that SWNH particle sizes in aqueous suspension (10 µg/mL) ranged from 295–615 nm. Determined on Zetasizer Nano ZS (Malvern Instruments, Malvern, UK) by means of dynamic light scattering.

Abbreviations: SWNHs40, 0.85 µg of single-walled carbon nanohorns per cm², or 24 µg per 60 mm dish; PS, polystyrene.