Original article

Lipidomics reveals carnitine palmitoyltransferase 1C protects cancer cells from lipotoxicity and senescence

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Lipotoxicity, caused by intracellular lipid accumulation, accelerates the degenerative process of cellular senescence, which has implications in cancer development and therapy. Previously, carnitine palmitoyltransferase 1C (CPT1C), a mitochondrial enzyme that catalyzes carnitinylation of fatty acids, was found to be a critical regulator of cancer cell senescence. However, whether loss of CPT1C could induce senescence as a result of lipotoxicity remains unknown. An LC/MS-based lipidomic analysis of Panc-1, MDA-MB-231, HCT-116 and A549 cancer cells was conducted after siRNA depletion of CPT1C. Cellular lipotoxicity was further confirmed by lipotoxicity assays. Significant changes were found in the lipidome of CPT1C-depleted cells, including major alterations in fatty acid, diacylglycerol, triacylglycerol, oxidative lipids, cardiolipin, phosphatidylglycerol, phosphatidylcholine/phosphatidylethanolamine ratio and sphingomyelin. This was coincident with changes in expressions of mRNAs involved in lipogenesis. Histological and biochemical analyses revealed higher lipid accumulation and increased malondialdehyde and reactive oxygen species, signatures of lipid peroxidation and oxidative stress. Reduction of ATP synthesis, loss of mitochondrial transmembrane potential and down-regulation of expression of mitochondrial genes indicated mitochondrial dysfunction induced by lipotoxicity, which could further result in cellular senescence. Taken together, this study demonstrated CPT1C plays a critical role in the regulation of cancer cell lipotoxicity and cell senescence, suggesting that inhibition of CPT1C may serve as a new therapeutic strategy through induction of tumor lipotoxicity and senescence.

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1. Introduction

The concept of cellular senescence was originally proposed in 1961 by Hayflick and Moorhead [1] who discovered that cells could not proliferate indefinitely under in vitro culture conditions. Currently, cellular senescence refers to the irreversible arrest of cell proliferation, characterized by inhibited proliferation ability, DNA replication, increased senescence-associated-β-gal (SA-β-gal) activity and senescence-associated secretory phenotype (SASP) [2,3]. Senescence response has profound implications in the suppression of cancer development, and it is a complex process caused by numerous triggers, including telomere shortening, genomic damage, and mitochondrial dysfunction [4–6]. Recently, a close link between cellular senescence and lipotoxicity was reported [7]. Lipotoxicity is caused by the accumulation of lipids, particularly fatty acids, in nonadipose cells or tissues that are not capable of metabolizing or storing lipids [8,9]. Disruption of lipid metabolism and lipotoxicity could lead to cellular dysfunction, which is involved in the development of various diseases, such as diabetes, Alzheimer’s disease and cancer [10,11]. A collective review of recent studies has proven that the main mechanism of senescence is lipid peroxidation and oxidative stress [12,13]. Excess lipids are vulnerable to the attack by reactive oxygen species (ROS) and conversion into toxic reactive lipid peroxides and oxidized lipids (oxi-lipid), which could have lipotoxic effects on mitochondrial DNA (mtDNA), RNA and proteins of mitochondrial machinery, and oxidation equilibrium, further leading to mitochondrial dysfunction and senescence [8,14,15]. In addition, it was reported that senescence could be a physiological degenerative process accelerated by...
lipotoxic insults, while conversely, the process of senescence can make cells more vulnerable to lipotoxicity [16].

Carnitine palmitoyltransferase 1C (CPT1C), a member of the CPT1 family, is an enzyme located in the outer mitochondrial membrane that catalyzes the acylation of long-chain fatty acids and their entry into mitochondria for fatty acid β-oxidation [17]. The CPT1 family consists of three members, namely, CPT1A, mainly expressed in the liver; CPT1B highly distributed in muscle; and CPT1C specifically expressed in brain and tumors [18]. Despite the great similarity of gene sequences to that of CPT1A and CPT1B, CPT1C exerts unique physiological functions, such as participating in learning and cognition, regulating the emotional state, and it is necessary for energy homeostasis as well as food intake [19–21]. Recently, it was found that compared with paracancerous tissues, CPT1C expression was abnormally increased in malignant tumors such as human lung cancer and breast cancer, and CPT1C could promote the proliferation and growth of tumor cells under metabolic pressure such as glucose deprivation and hypoxia, suggesting that CPT1C may serve as a new therapeutic target for anti-tumor treatment [22,23].

Most recently, extended culturing of pancreatic carcinoma PANC-1 cells was found to cause growth arrest and severe cellular senescence [24,25]. Low CPT1C expression is involved in replicative senescence and CPT1C is a novel regulator of cancer cell senescence. Moreover, CPT1C exerts a significant effect on tumor cellular senescence among the other members of the CPT family [26]. In addition, abnormal lipid metabolism was preliminarily found in replicative senescent PANC-1 cells [27]. However, whether loss of CPT1C could induce senescence via lipotoxicity remains unknown. Therefore, the current study aimed to elucidate whether knockdown of CPT1C could induce lipotoxicity resulting in cancer cell senescence. These results demonstrated that CPT1C plays a crucial role in the regulation of cancer cell lipotoxicity and cell senescence, suggesting that inhibition of CPT1C may serve as a new therapeutic strategy through induction of tumor lipotoxicity and senescence.

2. Materials and methods

2.1. Chemicals and reagents

Dulbecco’s modified Eagle’s medium (DMEM) and Roswell Park Memorial Institute (RPMI) 1640 medium were purchased from Corning Inc. (New York, USA) and HyClone Laboratories, Inc. (Los Angeles, CA, USA). Fetal bovine serum (FBS), penicillin sodium, and streptomycin sulfate solution were purchased from Gibco Life Technologies (Grand Island, NE, USA). All the other chemicals and solvents were of analytical grade or HPLC grade when appropriate and commercially available.

2.2. Cell culture and transfections with siRNA

Human pancreatic cancer PANC-1 cells were provided by the American Type Culture Collection (Manassas, VA, USA), human breast cancer cell line MDA-MB-231, human colorectal carcinoma cell line HCT-116 and human lung cancer cell line A549 were obtained from Dr. Jun Du (Sun Yat-sen University, China). Cells were cultured in DMEM or RPMI 1640 medium containing 10% FBS, 100 IU/mL of penicillin sodium, and 100 μg/mL of streptomycin sulfate at 37 °C in a 5% CO2 atmosphere with saturated humidity.

PANC-1, MDA-MB-231, HCT-116 and A549 cells were transfected with 5 nM small interfering RNA (siRNA) (Ribobio, Guangzhou, China) using Lipofectamine RNAiMAX transfection reagent (Invitrogen Corporation, Carlsbad, CA, USA) according to a previous report [24]. The siRNAs sequences are listed in Table S1. Each cancer cell line was transfected with siRNAs for 72 h prior to the following experiments.

2.3. Bromodeoxyuridine (BrdU), colony formation, and SA-β-gal assay

As previously reported [24], for BrdU incorporation measurement, a cell proliferation ELISA BrdU kit (Roche, CA, USA) was applied according to the manufacturer’s protocol. Briefly, after adding 10 μL of BrdU labeling solution, the cells were incubated for 2 h at 37 °C. A specific antibody anti-BrdU-POD working solution was employed and the absorbance wavelength was set at 370 nm. For colony formation assay, different densities (5000, 2500, and 1250 cells/well) of cells were seeded into 6-well plates and cultured at 37 °C for 10–14 days. 4% formaldehyde was used to fix the cells and 1% crystal violet was used to stain the cell colonies. For senescence-associated β-galactosidase activity assay, following the manufacturer’s protocol (Beyotime, China), cells were fixed with gluteraldehyde for 10–15 min and then washed twice with PBS before staining with X-gal staining solution overnight. Inverted fluorescence microscope was used to obtain photograph images.

2.4. Real-time PCR analysis

Total cellular RNAs from each cancer cell line were isolated using Trizol reagent (Takara Bio Inc., Shiga, Japan). Extracted RNA was reversely transcribed to cDNA by using Prime Script RT Reagent Kit (Takara Bio Inc., Shiga, Japan). Real-time PCR analysis was carried out in Applied Biosystems 7500 real-time PCR System by using SYBR Premix Ex-Taq II Kit (Takara Bio Inc., Shiga, Japan). Data were analyzed using the 2ΔΔCt method. The specific primers used for real-time PCR analysis were demonstrated in Table S2.

2.5. Lipid extraction, chromatographic and mass spectrometric conditions

For lipidomic analysis, the cells were grown in 6-well plates to approximately 90% confluence as previously described [27]. Briefly, the cells were quenched with liquid nitrogen and detached using cell scrapers. After two cycles of freeze–thaw, 1.2 mL of chilled extraction solvent (methanol/MTBE/H2O, 4:5:5, V/V/V) was added into 200 μL of cell suspension. After centrifugation at 1,000 rpm for 5 min at 4 °C, the mixtures were vortexed vigorously for 2 min and left on ice for 10 min. The supernatants were collected, and 200 μL of each sample was dried under nitrogen flow at room temperature, and stored at −20 °C until analysis. Finally, the residuals were resuspended with 100 μL of isopropanol/methanol (1:1, V/V) and injected for further UHPLC-ESI-MS-based lipidomics analysis.

Lipidomic chromatographic separation was carried out by using a reverse-phase Ascentis Express C18 100 mm × 2.1 mm 2.7 μm column (Sigma-Aldrich, St. Louis, MO, USA) on a Thermo Scientific Dionex Ultimate 3000 UHPLC system. Flow rate was set at 300 μL/min and column temperature was maintained at 45 °C. A gradient elution was employed with 50% acetonitrile in water with 5 mM ammonium formate and 0.1% formic acid (mobile phase A) and 5% acetonitrile in isopropanol with 5 mM ammonium formate and 0.1% formic acid (mobile phase B): 80% mobile phase A for 0.5 min, 20%–50% mobile phase B in 7 min, 50%–80% mobile phase B in the next 10 min, 80%–100% mobile phase B in 10 min and maintained 100% mobile phase B for 1.5 min, following 100%–20% mobile phase B for 0.1 min and 80% mobile phase A for 3 min. The quality controls for each cancer cell line were generated by pooling 20 μL of each sample and were injected intermittently during the whole process of sample injections.

Analysis of metabolites was performed with a Thermo Scientific Q Exactive™ benchtop Orbitrap mass spectrometer with a heated ESI source in both positive and negative modes. Untargeted
lipidomic profiling analyses acquired the data in the full MS-ddMS² mode (100–1000 m/z), followed by top-10 data-dependent MS/MS at 17,500 FWHM resolution. The other main parameters for MS/MS included AGC target 10⁵; maximum IT 65 ms; normalized step collision energy 25 and 35 for positive ion mode and 20, 30, and 40 for negative ion mode; apex trigger 5–10 s; and dynamic exclusion 10 s. The spray voltage was operated at 3.5 kV, and ion transfer capillary was set to 300 °C. Chromatograms and mass spectra from the UHPLC-ESI-MS runs were processed as raw files in Xcalibur (Thermo Scientific, San Jose, CA, USA).

2.6. Lipidomic data processing

Identification and relative quantification of the lipid molecular species were assisted by use of LipidSearch software (Thermo Scientific, San Jose, CA, USA) as reported previously [27]. The precursor tolerance was set to 5 ppm mass window while 10 ppm mass window was chosen as the product tolerance. The m-score threshold was set to 5.0. For multivariate principal component analysis (PCA), the raw data containing aligned peak areas with matched mass-to-charge ratios and retention times were transformed into a multivariate matrix and analyzed by SIMCA 13.0 software (Umetrics, Kinnelon, NJ, USA). Then supervised orthogonal partial least squares-discriminant analysis (OPLS-DA) was performed. The major lipid ions that contributed to the separation of sample groups were shown in the loading scatter S-plots.

2.7. Oil red O staining and Nile red fluorescence measurement

For oil red O staining, the cells were washed with PBS and then fixed with 10% formaldehyde for 30 min at room temperature. Stock solution of oil red O (0.5%) was diluted to 60% isopropl alcohol with water (3:2, V/V) and used to stain cells for 15 min in the dark, followed by washing carefully with water twice. After dyeing, images were obtained immediately under an inverted fluorescence microscope (Olympus, Tokyo, Japan) at 200 × magnification.

For Nile red fluorescence measurement, the cells were transfected for 72 h, washed with PBS and then fixed with 4% paraformaldehyde for 10 min. After adding 100 μL of 5 μg/mL Nile red solution and incubating for 15 min, the cells were washed gently with PBS twice and the fluorescence intensity was determined immediately (excitation wavelength at 543 nm and emission wavelength at 598 nm).

2.8. Malondialdehyde (MDA) and ROS assay

MDA assay was performed according to the manufacturer’s protocol (Nanjing Jiancheng Bioengineering Institute, Nanjing, China). Seventy-two h after transfection, a specific extraction solvent was added and cells were harvested after centrifugation and the protein was quantified. Glacial acetic acid and other reagents in the kit were added before heating for 40 min in a 95 °C water bath. The absorbance was measured at 530 nm wavelength. ROS assay was conducted using a DCFH-DA fluorescent probe (Nanjing Jiancheng Bioengineering Institute, Nanjing, China) and the final concentration was 10 μM. After incubation at 37 °C for 60 min, the fluorescence intensity was detected. The excitation wavelength was set at 485 nm while the emission wavelength was 525 nm.

2.9. ATP luminescent cell viability assay and mitochondrial transmembrane potential experiment

For ATP production measurements, 72 h after siRNA transfection, the cell plates were taken out at room temperature for 30 min and then 100 μL of CellTiter-Glo was added according to the manufacturer’s protocol (Promega Corporation, Madison, MI, USA). Cells were fully cracked by slowly shaking for 2 min and incubated for 10 min at room temperature to stabilize the luminescence signal. Luminescence intensity was recorded by a Flex Station 3 multifunctional enzyme marker device. The mitochondrial transmembrane potential was analyzed by fluorescence dequenching of rhodamine (rh) 123 (Sigma, St. Louis, MO, USA) with ArrayScanVTI High Content Application (Thermo Fisher, Waltham, MA, USA). The excitation wavelength was set at 485 nm while the emission wavelength was 535 nm.

2.10. Statistical analysis

All values of this study were expressed as mean ± SEM. Statistical analysis was conducted by SPSS and Two-tailed Student’s t tests. Graphs were performed using GraphPad Prism v6.0c software (San Diego, CA, USA). Significance was presented by *P < 0.05, **P < 0.01, and ***P < 0.001 vs. the control.

3. Results

3.1. Knockdown of CPT1C induces cellular senescence in cancer cells

Transfection with siRNA for 72 h was applied to knock down CPT1C in PANC-1, MDA-MB-231, HCT-116 and A549 cells and significant reduction of CPT1C mRNA level was found in all cell lines (Fig. S1). Combined with previous interfering efficiency results [24], CPT1C siRNA-3 for PANC-1, HCT-116 and A549 cells and CPT1C siRNA-1 for MDA-MB-231 cells were chosen for the following experiments.

Cellular senescence was monitored by BrdU incorporation, colony forming ability, SA-β-gal activity and mRNA expression of SASP. Lower colorimetric immunoassay, weaker capacity to form colonies, and enhanced SA-β-gal activity were observed in CPT1C siRNA-treated group (Fig. 1). Additionally, the expression levels of typical SASP factors, including interleukin 8 (IL8), IL1A, IL1B, matrix metallopeptidase 1 (MMP1) and MMP3 mRNAs were markedly increased when CPT1C was knocked down in cancer cells. Taken together, these results confirmed that knockdown of CPT1C induced cellular senescence in cancer cells.

3.2. Knockdown of CPT1C significantly changes the lipidome of cancer cells

To comprehensively investigate whether knockdown of CPT1C could affect the lipidome of these cell lines, lipidomic analysis was carried out in CPT1C siRNA-treated cancer cells. PCA was performed to uncover the differences in the lipidomic profiles under both positive and negative ion modes. Data points of quality controls coalesced, which indicated stability and reproducibility of the UHPLC-ESI-HRMS operation system (Fig. 2A). PCA showed clear separation between siControl-treated and CPT1C siRNA-treated cancer cells, indicating distinct discrimination in the lipidomes between the two groups. OPLS-DA and loading S-plots were conducted to reveal the contribution of each detected ion to the lipidomic distinctions. OPLS-DA separated well the siRNA CPT1C group from the siControl group among all samples (Fig. 2B). In the S-plot, the ions that contributed most to the discrimination were selected and identified with the aid of Lipidsearch software (Fig. 3).

The major lipid ions were identified as fatty acids (FA), triacylglycerol (TG), sphingomyelin (SM) and cardiolipin (CL).

The levels of specific lipid species and mRNA involved in lipid metabolism were measured and compared with the negative control. The total FA contents were remarkably increased in cancer cells...
when knocking down CPT1C (Fig. 4). The mRNA levels of genes related to FA synthesis and oxidation, including fatty acid synthase (FASN), sterol regulatory element-binding transcription factor 1 (SREBF1), acyl-CoA dehydrogenase long chain (ACADL), acyl-CoA oxidase 1 (ACOX1) and Acox2, were decreased in different degrees (Fig. 5). The levels of diacylglycerol (DG) were decreased whereas the levels of TG were increased, which was consistent with the elevated expressions of diacylglycerol acyltransferase 1 (DGAT1) and diacylglycerol acyltransferase 2 (DGAT2) and decreased expressions of patatin-like phospholipase domain-containing protein 2 (PNPLA2) and lipase E (LIPE). Higher levels of FA and TG revealed lipid accumulation after knockdown of CPT1C. Furthermore, total contents of oxidized lipids showed trends of increase, suggesting possible lipid peroxidation and oxidative stress. Moreover, reduced levels of mitochondrial characteristic lipids, CL and phosphatidylglycerol (PG), were found in the CPT1C siRNA group, indicating impaired mitochondrial function. In addition, lower ratios of phosphatidylcholine/phosphatidylethanolamine (PC/PE) and higher levels of SM were also observed in CPT1C knockdown cells (Fig. 4). The detailed information of FA, TG, oxo-lipid, CL and SM (different FA chains of carbon number and degree of unsaturation) with significant difference between the siRNA CPT1C group and the siControl group in four cancer cell lines is listed in Tables S3-S6.

Overall, these data suggested that knockdown of CPT1C could significantly change the lipidome and affect cancer cell lipid homeostasis. The dysregulation of lipid homeostasis could impact the normal physiological function of cells.

3.3. Knockdown of CPT1C induces lipid accumulation, lipotoxicity and mitochondrial dysfunction

Since the lipidomic data indicated that knockdown of CPT1C changed lipid metabolism, cellular lipotoxic assays were performed to further understand the relationship between CPT1C and lipotoxicity. Oil red O staining showed a significant increase of lipid droplets in CPT1C siRNA-treated cancer cells (Fig. 6A). Nile red fluorescence intensity measurements revealed elevated lipid contents in CPT1C siRNA-treated cells as compared with the siControl-treated cells (Fig. 6B). Contents of the typical toxic lipid marker malondialdehyde (MDA) were increased significantly, suggesting higher levels of lipid peroxidation (Fig. 6C). Also, higher level of ROS was observed, which indicated disturbed balance of oxidation and reduction, and oxidative stress (Fig. 6D). In brief, these results demonstrated that knockdown of CPT1C could induce lipid accumulation and lipotoxicity.

Since mitochondria is the main site of lipid metabolism and ROS production, the accumulation of lipid peroxides in mitochondria and the occurrence of oxidative stress can cause damage to mitochondrial structure and function [14]. ATP is the most direct energy source of cells and most of ATP is generated in mitochondria, while mitochondrial membrane potential is a necessary prerequisite to guarantee mitochondrial respiratory function [28]. Therefore, ATP production and mitochondrial membrane potential are important...
indicators of mitochondrial function. Lower levels of ATP and decreased mitochondrial membrane potential indicated mitochondrial dysfunction (Figs. 7A and B). Furthermore, lipid peroxides could have lipotoxic effects on mitochondrial DNA (mtDNA). Thus, mRNAs encoded by genes related to mtDNA synthesis were measured. Expressions of peroxisome-proliferator-activated receptor γ coactivator-1α (PGC1A), nuclear respiratory factor 1 (NRF1), transcription factor A mitochondrial (TFAM) and cytochrome b (CYBA) mRNAs were decreased to different extents after knockdown of CPT1C, indicating impaired mtDNA function (Fig. 7C).

4. Discussion

Cellular senescence can suppress the proliferation of tumor cells and inhibit the occurrence and development of tumors [29]. Our previous studies revealed that knockdown of CPT1C could induce cellular senescence in PANC-1, MDA-MB-231, HCT-116 and A549 cells [24,26]. In the current study, lipidomic analysis of these four cancer cell lines after CPT1C knockdown was performed. Cellular lipotoxicity was detected through a series of lipotoxic assays. The results demonstrated that loss of CPT1C could significantly change the lipidome, induce lipotoxicity, and further lead to mitochondrial dysfunction and senescence, suggesting universality in relationship between CPT1C, lipotoxicity and senescence in different cancer cells. CPT1C depletion may serve as a new therapeutic strategy through induction of cellular lipotoxicity and senescence.

It has been reported that CPT1C is an outer integral membrane protein enriched in hypothalamus with mitochondrial localization [30]. In our previous study [24], co-localization of CPT1C with mitochondria in PANC-1 cells was detected with immunofluorescence assays, which supports our conclusion that CPT1C is a key regulator of mitochondrial dysfunction and cellular senescence [24,25]. However, it is also found that CPT1C is localized in the endoplasmic reticulum (ER) of neurons [30]. Although the protein sequences of CPT1C containing necessary catalytic reaction sites share high similarity with CPT1A and CPT1B, the subcellular localization results showed CPT1C is also likely to be located in ER and microsomal structure [17]. Furthermore, whether CPT1C has CPT capacity remains controversial. When large amounts of CPT classic catalytic substrates such as carnitine and acyl-CoA esters were added into CPT1C-high expressed yeast, no acyltransferase activity was observed [31]. However, it is proved that CPT activity could be detected in overexpression of CPT1C PC12 cells and acyl-CoA is the catalytic substrate of CPT1C, but the catalytic activity of CPT1C was much lower than that of CPT1A and CPT1B [32]. Therefore, compared with CPT1A and CPT1B, CPT1C may possess other unique physiological functions, which needs to be further clarified.

Senescent cells exert several recognized features such as enhanced SA-β-gal activity, lower proliferation capacity, and weaker capacity to form colonies. However, not all aging cells exhibit all known senescent characteristics. Therefore, in general, at
least three indicators should be combined to investigate whether cell senescence occurs [3]. In the present study, lower BrdU proliferation capacity, weaker capacity to form colonies, and enhanced SA-β-gal activity and SASP expression levels were observed in PANC-1, MDA-MB-231, HCT-116 and A549 cells after CPT1C knockdown.

Fig. 3. OPLS-DA loading S-plots of PANC-1 (A), MDA-MB-231 (B), HCT-116 (C) and A549 (D) cells after knockdown of CPT1C. Data were obtained from LC-MS/MS under positive and negative ion modes, n = 6/group. Each point represents an individual lipid ion. The major significant lipid ions are labeled in the S-plots.
Compared with that of normal cells, lipid homeostasis of senescent cells also changes significantly [7]. Earlier studies have shown that abnormal lipid metabolism can serve as a new mechanism of metabolic transformation of tumor cells, and importantly, small changes in abundance, composition, or location of lipids can have profound effects on cell viability and function [33]. Lipidomics can systematically and comprehensively analyze the species and content of most lipid molecules, and thus has been widely used in the discovery of disease biomarkers, drug targets and related mechanisms [34]. Given the close relationship between senescence, CPT1C, fatty acid oxidation and transportation, a more extensive lipidomic analysis was carried out to elucidate the lipids affected by CPT1C levels.

PCA and OPLS-DA showed that knockdown of CPT1C could change the cancer cell lipidomes. S-plot analysis further revealed that the major lipid ions contributing to the difference between control and CPT1C siRNA-treated cells were FA, TG, SM and CL. Free FA refers to non-esterified FA which can be released by hydrolysis of TG, phospholipid and other lipids. FA can also be synthesized by FASN, which is regulated by SREBF1, and SREBF1 could be a potential gatekeeper for lipotoxicity [35]. FA oxidation is mainly carried out in mitochondria by ACADL, ACADM, ACOX1, ACOX2 and other FA oxidases. The major physiological functions of FA include energy supply via \( \beta \)-oxidation, structural composition of biofilm and formation of lipid signaling molecules [33,36]. Although at present, while it is not known whether CPT1C participates in FA oxidation, there are data supporting the view that CPT1C can affect FA metabolism. For example, after 14 weeks of high-fat diet, Cpt1c knockout mice was both increased, but the increase rate in cpt1c-null group was much lower than that in wild-type group [19,37]. Moreover, it was found that the FA oxidation rate of MCF7 cells was significantly increased after overexpression of CPT1C [22]. The current study showed that total FA content of tumor cells increased dramatically after CPT1C was knocked down, particularly of FA (16:0), FA (18:0) and FA (18:1). The accumulation of these saturated FA and mono-unsaturated FA in cells may cause mitochondrial damage. Excess FA and oxidative damage could induce biofilm damage and associated aging and metabolic diseases [38].

On the other hand, FASN, SREBF1, ACADL, ACADM, ACOX1 and ACOX2 mRNAs related to FA synthesis and oxidative metabolism were significantly reduced in the CPT1C siRNA cells, consistent with the report that excess saturated FA could inhibit the synthesis of endogenous FA through negative feedback regulation and disrupt FA generation and metabolic balance [36]. TG is a conjugate of glycerol with FA of various lengths and compositions that store and provide energy for cells. When cellular FA exceeds the requirement of synthesis and catabolism, FA can be converted into TG by esterification enzymes and stored in lipid droplets. Therefore, the increase of TG content can also indicate the accumulation of FA in cells to some extent [39]. TG accumulation during replicative senescence and the observation of excessive TGs in a nonadipose tissue were identified as markers of lipotoxicity [7]. CPT1C promotes the storage of acyl-CoA of TG in neurons, rather than the oxidation process [40]. A previous study reported that FA and TG accumulated in senescent PANC-1 cells [27]. In the present study, the total TG contents in CPT1C knockdown cells showed an increasing trend while lower levels of DG were observed. After CPT1C knockdown, the DGAT1 and DGAT2 mRNAs involved in synthesis were increased, while PNPLA2 and LIPE mRNAs encoding...
Gene expressions related to the homeostasis of the above lipids in PANC-1 (A), MDA-MB-231 (B), HCT-116 (C) and A549 cells (D) upon knockdown of CPT1C. Data are represented as mean ± SEM, n = 4/group (*P < 0.05; **P < 0.01; ***P < 0.001 vs. siControl group). Fold change means the siRNA CPT1C vs. the siControl group. *FASN: fatty acid synthase; SREBF1: sterol regulatory element-binding transcription factor 1; ACADL: acyl-CoA dehydrogenase long chain; ACOX: acyl-CoA oxidase; DGAT: diacylglycerol acyltransferase; PNPLA2: patatin-like phospholipase domain containing protein 2; LIPE: lipase E; TAMM41: TAM41 mitochondrial translocator assembly and maintenance homolog; CDS1: CDP-diacylglycerol synthase 1; TAZ: tafazzin.

higher levels of MDA and ROS were observed, indicating that knockdown of CPT1C could induce lipid peroxidation and oxidative stress.

Since mitochondria are the main site of FA oxidation and ROS production, the accumulation of lipid peroxides in mitochondria can cause serious lipid toxic damage to mitochondrial structure and function, and lead to cell dysfunction [14]. In the current study, a series of assays were conducted to evaluate mitochondrial function. A reduction of mitochondrial characteristic lipids, CL and PG, were found in the siRNA CPT1C-treated cells. Diphosphatidyl glycerol (CL), also known as cardiac phospholipids, is a characteristic lipid associated with mitochondrial function (oxidative phosphorylation, respiratory chain) and apoptosis [44]. Reduction of CL in senescent cells was also reported, indicating that CL content is related to cell senescence [45]. PG is a precursor molecule for the synthesis of CL and is responsible for the mitochondrial respiratory chain activity. Since CL is located in the mitochondrial inner membrane, it is particularly susceptible to lipid peroxidation. After the peroxidation of the most common mitochondrial CL-tetramethylacyl CL in mammalian tissues, not only the lipid hydrogen peroxide in the membrane accumulated, but also the mitochondrial toxic metabolite 4-hydroxynonenal was produced. Peroxidation of CL is often accompanied by a decrease in the activity of complexes I and IV, resulting in impaired mitochondrial respiration [46]. CL synthesis is mainly catalyzed by TAM41 mitochondrial translocator assembly and maintenance homolog (TAMM41), CDP-diacylglycerol synthase 1 (CDS1), and the spatial configuration arrangement of CL needs to be catalyzed by tafazzin (TAZ) [44]. The current study showed that after CPT1C knockdown, TAMM41, CDS1 and TAZ mRNAs were decreased, which was consistent with significantly decreased levels of ATP and decreased mitochondrial membrane potential were also found upon CPT1C knockdown. Furthermore, genes related to mtDNA synthesis were detected. Biosynthesis of mtDNA was mainly regulated by the PGC1A-NRF1-TFAM pathway. When the expressions of PGC1A was inhibited, the expressions of its downstream signaling molecules, such as NRF1 and TFAM, were inhibited, which affected mtDNA replication.
Cytochrome b, a representative subunit of mtDNA encoding gene, can reflect the copy capacity of mtDNA. The position of mtDNA in mitochondria is relatively bare and vulnerable to oxidative damage, and therefore the decreased biosynthesis capacity of mtDNA indicates mtDNA damage [47,48]. In the present study, the expressions of PGC1A, NRF1, TFAM and CYBA (cytochrome b) mRNAs were decreased to various extents, reflecting impaired mtDNA function.

Taken together, these results showed that knockdown of CPT1C could induce lipotoxicity and mitochondrial dysfunction, which is consistent with the results of a previous finding that CPT1C regulates cancer cell senescence through mitochondria-associated metabolic reprogramming [24].

In addition, a lower ratio of PC/PE and higher levels of SM were also observed in CPT1C knockdown cells. PC and PE, the most common phospholipids within cells, are the main structural lipids in biofilms. Specifically, membrane fluidity and permeability can be damaged by oxidation of lipid membranes. Alteration of membrane components (PC/PE) can reduce membrane mobility, increase stiffness and permeability, resulting in senescence [49,50]. SM is also the major structural component of cell membrane, and its metabolites, ceramide and sphingosine, are bioactive signaling molecules [51]. It was reported that the SM content was increased during senescence, but the specific relationship and mechanism of SM and senescence remains unknown [52]. Ceramide can regulate cell proliferation and apoptosis, and neuronal ceramide metabolism can be regulated by CPT1C [53]. However, in this study, there was no statistical difference of the ceramide after CPT1C knockdown, suggesting that silencing CPT1C had little effect on tumor cell ceramide metabolism.

Knockdown of other CPT members also has been investigated. Lipid accumulation was related to CPT1A mRNA expression in skeletal cells [54]. Lipids accumulated and abnormal mitochondrial function was noted in cardiac muscle of Cpt1b+/− mice, demonstrating that absence of Cpt1b can aggravate the cardiac hypertrophy possibly due to lipotoxicity [55]. While in cancer cells, combined with previous results [26], CPT1C exerted a significant role in mitochondria-associated senescence via lipotoxicity compared with other CPT isoforms.

In summary, the present study demonstrates that CPT1C plays a crucial role in the regulation of cancer cell lipotoxicity and cell senescence. For the reason that the role of Cpt1c in FA metabolism remains largely unresolved, further studies are warranted to provide more information about the relationship and molecular
mechanisms of CPT1C, lipotoxicity and senescence. This study suggests that inhibition of CPT1C may serve as a new therapeutic strategy through induction of tumor lipotoxicity and senescence.

Declaration of competing interest

The authors declare that there are no conflicts of interest.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jpha.2020.04.004.

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