Sterol Regulatory Element-binding Protein (SREBP)-1-mediated Lipogenesis Is Involved in Cell Senescence

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Increased cell mass is one of the characteristics of senescent cells, but this event has not been clearly defined. When subcellular organelar mass was estimated with organelle-specific fluorescence dyes, we observed that most membranous organelles progressively increase in mass during cell senescence. This increase was accompanied by an increase in membrane lipids and augmented expression of lipogenic enzymes, such as fatty acid synthase (FAS), ATP citrate lyase, and acetyl-CoA carboxylase. The mature form of sterol regulatory element-binding protein (SREBP)-1 was also elevated. Increased expression of these lipogenic effectors was further observed in the liver tissues of aging Fischer 344 rats. Ectopic expression of mature form of SREBP-1 in both Chang cells and primary young human diploid fibroblasts was enough to induce senescence. Blocking lipogenesis with FAS inhibitors (cerulenin and C75) and via siRNA-mediated silencing of SREBP-1 and ATP citrate lyase significantly attenuated H2O2-induced senescence. Finally, old human diploid fibroblasts were effectively reversed to young-like cells by challenging with FAS inhibitors. Our results suggest that enhanced lipogenesis is not only a common event, but also critically involved in senescence via SREBP-1 induction, thereby contributing to the increase in organelle mass (as a part of cell mass), a novel indicator of senescence.

Enlarged cell morphology is a typical characteristic of senescent cells, reflecting increased cell mass (1–3). However, the detailed components of the mass are not clearly defined, and whether the increase in mass is due to the increase of any specific cellular compartment(s), or all, is unknown. Until now, the lysosome has been the most recognized organelle increased in both senescent and aged cells (4, 5). However, senescent lysosomes are filled with indigestible autofluorescent pigments, such as lipofuscin, and displaying nonfunctionality (6). Recently, an increase in mitochondrial mass has been reported in stress-induced senescence (7) and replicative senescence (8, 9), which was generally explained to be the result of its compensatory biogenesis to cope with the persistent functional defects of mitochondria (10, 11). An increase in mitochondrial mass is further augmented by the inability of the lysosome to remove damaged mitochondria, thereby allowing the damaged mitochondria to generate hazardous reactive oxygen species (ROS) continuously. This sequence suggests that most senescent mitochondria are also not functionally intact. The functional deterioration of mitochondrial respiration accompanied by a dramatic increase in their mass during cell senescence and aging has often been found, supporting the hypothesis (7, 12).

The close link between the accumulated nonfunctional organelles, mitochondria and lysosomes, yields the “mitochondrial-lysosomal axis theory of aging” (13, 14). This theory sufficiently supports the generation of persistent oxidative stress during the progression of senescence. Analogous to this theory, the inevitable increase in nonfunctional organelles due to the combined actions of the uncontrolled ROS generator (mitochondria) and the nonfunctional eradicator (lysosome) is likely not restricted to mitochondria, but may be applied to all subcellular components. Therefore, whether all cellular organelles, especially membranous ones, are also under the compensatory enhancement of their biosynthesis and, if so, whether these features are genuinely involved in the progression of cell senescence should be evaluated. Monitoring the biogenic activity of each organelle is not simple; however, lipid biosynthesis is a critical element for the biogenesis of all membranous organelles and may be a plausible barometer of overall membranous organelle biogenesis.

Lipid biosynthesis, called lipogenesis, includes fatty acid synthesis and the subsequent synthesis of triglycerides and membrane phosphoglycerides. Lipogenesis is activated during cell growth and proliferation to provide membranous components for subcellular organelle formation (15, 16). Lipogenesis comprises multiple sequential steps governed by several key enzymes, including ATP citrate lyase (ACL), acetyl-CoA carboxylase (ACC), and fatty acid synthase (FAS).

The abbreviations used are: ROS, reactive oxygen species; ACC, acetyl-CoA carboxylase; ACL, ATP citrate lyase; ad-mSREBP, mature SREBP-1 adenovirus; DCFH-DA, 2′,7′-dichlorodihydrofluorescein diacetate; DFO, deferoxamine; DT, doubling time; FAS, fatty acid synthase; FSC, forward scatter; HDF, human diploid fibroblast; PD, population doubling; SA-β-galactosidase, senescence-associated β-galactosidase; SREBP, sterol regulatory element-binding protein; SSC, side scatter.

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carboxylase (ACC), and fatty acid synthase (FAS) (17). ACL converts cytosolic citrate into acetyl-CoA and oxaloacetate, supplying essential metabolites for lipid biosynthesis. ACC catalyzes the irreversible carboxylation of acetyl-CoA to produce malonyl-CoA, and FAS subsequently condenses acetyl-CoA and malonyl-CoA to generate long chain fatty acids. Interestingly, the expression of these lipogenic enzymes is controlled by the common lipogenic transcription factors, sterol regulatory element-binding proteins (SREBPs). Three SREBP isoforms (SREBP-1a, -1c, and -2) have been identified. SREBP-1a and -1c are alternative spliced forms transcribed from two distinct promoters on the same gene, whereas SREBP-2 is encoded from a different gene. SREBP-1 plays a greater role in regulating fatty acid synthesis than cholesterol synthesis, whereas SREBP-2 is relatively selective in transcriptionally activating genes involved in cholesterol biosynthesis (18, 19). SREBPs are initially bound to the rough endoplasmic reticulum membrane but, upon activation, are cleaved to liberate the N-terminal portion containing a basic helix-loop-helix leucine zipper domain. The cleaved active fragment enters the nucleus, where it binds to specific sterol response elements within the promoter regions of the lipogenic genes and activates their transcription (17, 18). Therefore, SREBP activation can be used as a good indicator of the lipogenic status of the cell. In addition, SREBP is the best target for controlling cellular lipogenesis and exploring the relationship between lipogenesis and senescence.

In this study, we show that all subcellular organelles are generally increased with membrane lipids in both stress-induced senescence and replicative senescence. In addition, we demonstrate that lipid biosynthesis is directly involved in senescence-associated organelle biogenesis via SREBP-1 activation. These results suggest that SREBP-1-mediated lipogenesis is an important modulator of senescence and provide new insight into the metabolic background of aging and aging-related pathogenesis. The overall organelle increase is also indicated as a novel indicator of senescence.

**EXPERIMENTAL PROCEDURES**

**Cell Culture, Generation of Cell Senescence, Cell Growth Rate**—Chang cells were purchased from the American Type Culture Collection and were cultured in DMEM supplemented with 10% fetal bovine serum (FBS; Invitrogen) and antibiotics at 37 °C in a humidified incubator with 5% CO₂. Chang cells, immortalized normal human hepatocytes, were chosen to understand metabolic backgrounds related to stress-induced senescence due to their homogeneous replicative capacity and strong metabolic properties. To generate stress-induced senescence, Chang cells were exposed to 200 µM deferoxamine (DFO) or H₂O₂ for the indicated periods. For replicative senescence, the primary human diploid fibroblasts (HDFs) were generously provided by Dr. Lim (Ajou University, South Korea) (20). Confluent cells were evenly transferred into two new dishes, and the cells were cultured until getting confluent again to generate one population doubling. The numbers of population doublings (PD, number) as well as the doubling time (DT, days) were continuously counted. Young and old cells used in the present experiments were defined as the HDF with a doubling time of around 24 h and >1 week, respectively.

To evaluate cellular growth rate, cells were harvested by trypsinization and counted with a hemocytometer after staining with 0.4% (w/v) trypan blue (Invitrogen) to exclude dead cells. No significant dead cells were observed during both stress-induced and replicative senescence. The total cell number of trypan blue-negative live cells was obtained upon completion of the experiments.

**Estimation and Visualization of Subcellular Organellar Mass**—To estimate cellular organellar mass, cells were stained with organelle-specific fluorescence dyes according to the manufacturer’s instruction. Briefly, proper concentrations and incubation times for each organelle were first determined. Mitotracker Red CMXRos (100 nM, M7512; Molecular Probes), Lyso Tracker Green DND-26 (1 µM, L7526; Molecular Probes), and ER Tracker Blue-White DPX (1 µM, E-12353; Molecular Probes) were applied for 30 min at 37 °C. Especially to stain the Golgi body, cells were incubated for 30 min at 4 °C with 5 µM BODIPY C5-ceramide (D3521; Molecular Probes) after pre-conjugating the dye with BSA and further incubated in fresh medium at 37 °C for 30 min. After staining cells with the dyes, cells were harvested in phosphate-buffered saline (PBS), and their fluorescence intensities were analyzed by flow cytometry (FACSCanto™ II; Becton Dickinson Corp.). Subcellular organellar masses were represented as mean values of an arbitrary fluorescence unit of 10,000 cells.

To confirm specific staining of subcellular organelles with the dyes, fluorescence images of the stained cells were visualized by Axiovert 200M with a cooled AxioCam HRm operated by the Axiovision 4.8 software (Carl Zeiss AG, Göttingen, Germany). Nuclear membrane mass was represented by nucleus area (square micrometers) measured from the nuclear fluorescence images of the Hoechst 33258-stained cells. Cytoplasmic membrane mass was represented by cell area (square micrometers) measured from phase contrast images of the cells. The nuclear or cytoplasmic area was calculated automatically by Axiovision 4.8 software (Carl Zeiss AG).

**Determination of Intracellular ROS, and Cell Size and Granularity**—To estimate intracellular ROS level, cells were stained with 10 µM 2',7'-dichlorodihydrofluorescein diacetate (DCFH-DA; Molecular Probes) for 15 min at 37 °C before performing flow cytometric analysis with FACSCanto™ II. Cell size and granularity were evaluated by analyzing forward scatter (FSC) and side scatter (SSC) of the stained cells, respectively, as described previously (3).

**Thin Layer Chromatographic Analysis of Cellular Lipids**—To analyze cellular lipid profiles, cellular lipids were extracted as described by Folch et al. (22) with slight modification. Briefly, the indicated numbers of cells were harvested into glass tubes and mixed with 1 ml of chloroform:methanol (2:1). After vortexing, H₂O (500 µl) was added into the tubes and centrifuged at 1,500 rpm for 15 min. The lower chloroform phase was collected and backwashed by adding methanol (300 µl) and H₂O (300 µl). After removing the top layer, chloroform phases were concentrated by evaporating chloroform under N₂ gas and loaded onto a TLC plate (1.0572.0001, 20 × 20, silica-coated; Merck). To separate the cellular lipids, we used the Kupke and Zeugner method (23) with a slight modification. Briefly, the chromatogram was first developed in a saturated chamber with...
solvent I (chloroform:methanol:H₂O = 65:30:5) to allow polar lipids to be separated. The solvent front was allowed to migrate ~10 cm from the bottom of the plate. After evaporating the solvents, the nonpolar lipids were separated by developing the chromatogram with solvent II (n-hexane:diethyl ether:acetic acid = 80:20:1.5) up to 1 cm from the top of the plate. The plates were dried under a stream of air and sprayed with primulin (Sigma) solution (50 μg/ml in acetone:water (80:20)). The plates were dried again, and fluorescent lipid spots were scanned on a UV transilluminator (SL-20; Seoulin Scientific Co., Seoul, Korea) equipped with a camera (PC1210; Canon Inc., Japan). The densities of lipid spots were estimated by using Gel-Pro Analyzer 4.0 software (Media Cybernetics, Silver Spring, MD). Standard lipids were obtained from Sigma as follows: cholesteryl palmitate (C6072), palmitate (P5586), cholesterol (C8667), cardiolipin (C0563), 1,2-diacyl-sn-glycerol-3-phosphatidylserine (P7769), L-α-lysophosphatidylcholine (L4129), and 3-sn-lysophosphatidylethanolamine (L4754).

Western Blot Analysis and Antibodies—Cells were washed twice with PBS and lysed with lysis buffer (50 mM Tris-Cl, pH 7.5, 0.1 mM NaCl, 1 mM EDTA, 1% (v/v) Triton X-100, 10 μg/ml each aprotinin and leupeptin, and 1 mM PMSF). A portion (30 μg) of the lysate was applied to Western blot analysis as described previously (7). Antibodies against SREBP-1 (catalog 557036) and SREBP-2 (catalog 557037) were purchased from BD Biosciences, and antibody for β-actin (catalog 8457) was from Cell Signaling Technology. Antibodies against SREBP-1 (catalog 2217) and SREBP-2 (catalog 2216) were purchased from Santa Cruz Biotechnology. Antibodies for tubulin (T-9026) and anti-LC3 (G3526), were kindly provided by Prof. Kyung-Sup Kim (Yonsei University, Seoul, Korea). siRNA duplexes were introduced into cells using Oligofectamine (Invitrogen) according to the manufacturer’s instructions.

Senescence-associated β-Galactosidase (SA-β-galactosidase) Assay—SA-β-galactosidase activity was assayed at pH 6.0 as described by Dimri et al. (25) with a slight modification. Briefly, cells were washed twice with PBS, fixed to plates by 3.5% formaldehyde for 5 min, and then incubated overnight in freshly prepared staining solution (40 mM citrate-phosphate buffer, pH 6.0, containing 1 mg/ml 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (Sigma), 5 mM potassium ferrocyanide, 5 mM potassium ferricyanide, 150 mM NaCl, and 2 mM MgCl₂). The stain was visible 12 h after incubation at 37 °C. By counting the numbers of the blue and total cells using ImageJ software (National Institutes of Health), the percentage of the SA-β-galactosidase-positive cells was obtained to estimate the degree of senescence-associated cells.

RESULTS

Overall Mass Increases in Membranous Subcellular Organelles of Senescent Cells—First, we examined changes in mass of all cellular organelles, especially membranous ones, in stress-induced senescence triggered by DFO and H₂O₂ as described previously (7, 26, 27). Subcellular organelles were stained with organelle-specific fluorescent dyes, and their masses were estimated by flow cytometric analysis (Fig. 1A) after confirming their specificities by visualizing with fluorescence microscopy (supplemental Fig. S1). Unexpectedly, increased organelle mass was observed not only in lysosomes and mitochondria, but also in other organelles, such as the endoplasmic reticulum and Golgi. Whether the number of organelles was increased or whether their individual volumes enlarged without changes in number was difficult to clarify, one clear finding was that the total membranous portion of the specified organelle within a single cell was increased. In the case of the nucleus, senescent cells developed either a single giant nucleus or highly conjugated multinuclei. To estimate the membranous portion of the nucleus, we measured its single-sectioned area after staining with Hoechst 33258. Regardless of the morphological diversity, the nuclear area of most senescent cells was significantly increased compared with control cells, suggesting a marked increase in nuclear membrane (Fig. 1B). Increased cytoplasmic membrane was also identified using an analogous approach (Fig. 1C). Moreover, similar results of overall mass increases in membranous organelles were obtained during the replicative senescence of primary HDFs, which is the best representative model of cell senescence (Fig. 1, D and E). These results indicate that the mass of all subcellular membranous organelles is commonly increased in senescence, contributing to the increase in cell mass. The results also suggest that membrane lipids, major components of the organelles, may increase for this phenomenon.

Increase in Membrane Lipid Content and Lipogenesis in Senescence—Biological membranes are complex structures with up to 70% of their composition being membrane lipids. To measure membrane lipid content, total cellular lipids were extracted and separated by TLC (Fig. 2Aa). Total lipids (sum of the lipids indicated as 1–5 in Fig. 2Aa) increased roughly 3-fold.
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Potential membrane lipids (sum of the lipids indicated as 2–5 in Fig. 2A) positioned with similar retardation as the standard membrane lipids, such as phosphatidylethanolamine, phosphatidylserine, cardiolipin, and cholesterol, amounted to ∼89% of the total lipids in control cells and obviously increased to 95.5% in DFO-induced senescent cells, whereas the potential storage lipid (indicated as 1 in Fig. 2A) with retardation similar to that cholesteryl palmitate decreased (Fig. 2Ab). Increases in total lipid content and the membrane lipid portion were also observed in H$_2$O$_2$-induced senescent cells (supplemental Fig. S2A). Unquestionably, old HDFs were found to have increased total lipids (2.4-fold) and an increased portion of membrane lipids (96.11% in young, 98.4% in old; Fig. 3A). These results indicate that an increase in cellular lipids, especially membrane lipids, can also be recognized as a common characteristic of senescence, supporting the increase in organellar mass.

Next, we investigated whether the increase in membrane lipids results from the activation of lipid biosynthesis. We monitored the expression level of the mature form of SREBP-1, which is known to be a master regulator of the transcription of diverse lipogenic enzymes (28). As expected, the active mature form of SREBP-1 was clearly increased in DFO-induced senescence (Fig. 2B). The protein levels of SREBP-1 target enzymes FAS, ACC, and ACL were also monitored. When the enzyme levels were normalized to tubulin, they reached a maximum at 24 h (Fig. 2Bb) and then slightly decreased, which does not inhibit further studies.

### FIGURE 1.
Overall increases of membranous subcellular organelles in cell senescence. A–C, Chang cells were treated with 200 μM DFO or H$_2$O$_2$ for 3 days. A, subcellular organellar masses were estimated by comparing the fluorescence intensities with flow cytometry after staining cells with organelle-specific dyes as described under "Experimental Procedures." ER, endoplasmic reticulum. B, nuclear membrane mass of a cell was represented by single-sectioned area (square micrometers) of nucleus of each cell. Upper panels are the representative images for both single giant nucleus (SN) and multinuclei (MN) of H$_2$O$_2$-treated single senescent cell. Scale bar, 5 μm. C, cellular cytoplasmic membrane mass was represented by single-sectioned area (square micrometers) of each cell. Upper panels are representative cell phase images of H$_2$O$_2$-treated senescent cell. Scale bar, 10 μm. D, subcellular organelar masses of young (Y, PD 16 or 19) and old (O, PD 79 or 80) HDFs. E, nuclear membrane mass (a) and cytoplasmic membrane mass (b) of young (PD 16), mid-old (MO, PD 65) and old (PD 83) HDFs were also estimated as described in B. **p < 0.01. Error bars, S.D.

### FIGURE 2.
Increased lipogenesis in DFO-induced senescence. Chang cells were treated with 200 μM DFO for the indicated periods. A, cellular lipid profile of senescent cells obtained by TLC as described under "Experimental Procedures." Representative TLC image (a) and quantitative estimation (b) of cellular lipids extracted from different numbers of cells are shown. Standard lipid mixture (Std) containing 10 μg each was used. Cholesteryl palmitate (CP, 1) belongs to nonpolar storage lipid, and cholesterol (CL, 2), cardiolipin (CA, 3), phosphatidylcholine (PC, 4) and phosphatidylserine (PS, 5) belong to membrane lipids. d in the x axis stands for day. B, Western blot for SREBP-1. Full-length and mature forms of SREBP-1 are indicated by arrows. C, Western blot analyses for lipogenic enzymes. Expression ratios of ACC (f), ACL (Œ), and FAS (Œ) to tubulin (b) or lamin (c) are shown.
correlate with the pattern of total cellular lipid content. We confirmed that the increase in cytoplasmic proteins is much greater than the increase in nuclear proteins (data not shown), although nuclear mass also increases (Fig. 1B) during cell senescence, implying that overall cytoplasmic mass is the combined result of all of the other cytoplasmic organelles except the nucleus. Therefore, we normalized the enzyme expression levels to lamin (nuclear protein) to represent the protein level per cell because cellular lipid contents were analyzed based on cell number. In this case, progressive and dramatic increases in the lipogenic enzymes were clearly observed (Fig. 2Bc). Similar results were obtained with H2O2-induced senescence (supplemental Fig. S2C). This increased expression of lipogenic enzymes and the active form of their master key regulator, SREBP-1c, was further observed in replicative senescence (Fig. 3, B and C) and the aging F344 rat model (Fig. 4), even when normalized to tubulin. Taken together, these results suggest that enhanced lipogenesis via SREBP-1 activation may contribute to the increased cellular lipids of senescence and aging.

**Enhanced Lipogenesis Is Enough to Induce Cellular Senescence**—To determine whether enhanced lipogenesis is genuinely involved in the development of cell senescence, we examined the effect of enhanced lipogenesis using the ectopic expression of active SREBP-1. We introduced adenovirus harboring the N-terminal fragment (mature form) of rat SREBP-1 into Chang cells. As shown in Fig. 5A, overexpressed mature SREBP-1 (mSREBP-1) was sufficient to increase the expression of FAS and ACL without affecting endogenous SREBP-1 or SREBP-2 expression. mRNA levels of all of the lipogenic enzymes (FAS, ACC, and ACL) were significantly induced by mSREBP-1 at the transcriptional level, implying its transcriptional activation (supplemental Fig. S4A). This finding implies that mSREBP-1 overexpression can effectively enhance cellular lipogenesis in a cell culture system. Excitedly, this enhanced lipogenesis by mSREBP-1 overexpression alone clearly induced senescence, as shown by an increase in SA-β-galactosidase activity, arrest of cellular growth, increased cell granularity represented by SSC, and increased cell size represented by FSC (Fig. 5, B and C). The induction of senescence was further confirmed by the induction of the p21 and p16 proteins, the inhibitory cell cycle regulators (Fig. 5A). Similar and more remarkable results by overexpressing SREBP-1 on gain of senescence phenotypes, including SA-β-galactosidase, granularity (SSC), and delayed cell growth in addition to induction of lipogenic enzymes were obtained in young HDFs (Fig. 6 and supplemental Fig. S4B). These data provide strong evidence that enhanced lipogenesis, at least by SREBP-1 activation, is sufficient for inducing senescence.

**Blocking Lipogenesis Significantly Attenuates Cell Senescence**—Next, we investigated the effect of blocking lipogenesis on H2O2-induced senescence using siRNA-mediated knockdown of SREBP-1 or ACL expression and pharmacological FAS inhibitors. SREBP-1 knockdown effectively attenuated all senescence phenotypes induced by H2O2 (Fig. 7, A and B). Attenuation of the senescence phenotypes correlated well with intracellular ROS levels, suggesting that the enhanced lipogenesis may be tightly linked to cellular oxidative stress (Fig. 7D). The H2O2-induced senescence was also diminished by siRNA-mediated ACL knockdown or FAS-specific inhibitors cerulenin and C75 (29, 30) as shown in Fig. 7, C and D.

Finally, we investigated whether the inhibition of lipogenesis can reverse senescent phenotypes of old HDFs, which are developed spontaneously by a series of cell passage rather than by any
exogenous insult, to young-like phenotypes. For this purpose, we first determined optimal concentrations of FAS inhibitors (data not shown). Surprisingly, most senescent phenotypes of old HDFs were effectively reversed by repeatedly exposing the cells to a determined concentration of FAS inhibitors (cerulenin and C75), although the effect on cell granularity (estimated by SSC) was minor (Fig. 8). Higher concentrations of FAS inhibitors were toxic to cells, and single treatment of the low dose was not enough to display the inhibitory effects (data not shown). These data imply that lipogenesis is importantly involved in cellular senescence, and pharmacologically blocking of lipogenesis at fatty acid synthesis is an effective approach for attenuating cellular senescence.

DISCUSSION

Increases in cell mass during senescence and the aging process have often been documented (3). Although the detailed components of the mass have not been clearly understood, increasing numbers of reports on the mass of lysosomes and mitochondria in senescence indicate that particular organelles seem to be associated with the mass of the senescent cell (3–5, 7–9). However, in the present study, we emphasize that, rather than certain organelles, all cellular organelles, particularly membranous organelles, increase during the development of cell senescence, as shown in both stress-induced and replicative senescence. These organellar mass increases correspond well with the increase in membrane lipid content and induction of lipogenic enzymes (FAS, ACC, and ACL) and SREBP-1. This finding implies that enhanced lipogenesis may contribute to the overall organelle increase in cell senescence. Therefore, in preference to investigating the changes or importance of individual organelles, we focused here on exploring the involvement of lipogenesis in the development of senescence. Indeed, enhancing lipogenesis by overexpressing mature SREBP-1 was enough to induce senescent phenotypes in both immortalized Chang cells and primary HDFs. Furthermore, blocking lipogenesis using siRNA-mediated knockdown of SREBP-1 or ACL significantly attenuated H2O2-induced senescence. More interestingly, the inhibition of lipogenesis by FAS-specific inhibitors reversed the senescent phenotypes, not only of the stress-induced senescence, but also of replicative senescence. All of
these results demonstrate that enhanced lipogenesis is critically involved in the senescence process, possibly contributing to the membranous organellar mass and, consequently, cell mass (Fig. 9).

The relationship between phospholipid synthesis and senescence or aging has been discussed in only a few reports, although reports on storage fat or oxidatively damaged lipids with aging are increasing (31–33). Inhibitors of HMG-CoA reductase, one of the targets of SREBP, reduce cellular senescence (34, 35), supporting our hypothesis. However, contradictory results have also been reported. Maeda et al. (36) observed that de novo phospholipid synthesis declines in senescent human fibroblasts, although the formation of cholesterol was increased compared with young cells. In that same study, senes-

FIGURE 7. SREBP-1 knockdown attenuates H2O2-induced senescence of Chang cells. A–C, Chang cells (1 × 10^4) seeded on 6-well plates were transfected with siRNAs for SREBP-1 and then treated with 200 μM H2O2 for 4 days. Aa, knockdown of SREBP-1 expression was confirmed by Western blotting. Ab, SA-β-galactosidase activities are shown. B, cell size and cell granularity were estimated by comparing FSC (a) and SSC (b) of cell distribution with flow cytometry, respectively. Ba, cell growth rate was compared by counting trypan blue-negative live cells. Dead cells were hardly observed. Bd, intracellular ROS levels were estimated by flow cytometric analysis of DCF fluorescence after staining cells with DCFH-DA. C, Chang cells (1 × 10^4) were treated with siRNAs for ACL and then treated with 200 μM H2O2 for 4 days. SA-β-galactosidase activities (upper panel) and Western blot analysis of ACL (lower panel) are shown. D, Chang cells were pretreated with the indicated concentrations of cerulenin (Cer; Sigma C2389) or C75 (Sigma C5490) and then treated with 200 μM H2O2 for 4 days. SA-β-galactosidase activities (upper panel) and Western blotting were performed. Two different siRNAs for each protein (SREBP-1 or ACL) were used. siRNA of random sequence was used as negative control (NC) and DMSO was used as vehicle (V). ##, p < 0.01 versus no H2O2. **, p < 0.01 versus NC or V.

FIGURE 8. Reversal of senescent phenotypes of old HDFs by pharmacological inhibition of lipogenesis. Old HDFs (PD 73, DT 8) were treated with the indicated concentrations of FAS inhibitors and replenished with new medium containing the inhibitors every other day for 2 weeks. A, cell growth rate was estimated by counting trypan blue-negative live cells. Dead cells were hardly detected. B, SA-β-galactosidase activity is shown. C, cell size was estimated by FSC analysis with flow cytometry. D, cell granularity was estimated by SSC analysis with flow cytometry. *, p < 0.05; **, p < 0.01 versus vehicle; error bars, S.D.

FIGURE 9. Schematic diagram for involvement of lipid biosynthesis in cellular senescence.
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cent cells developed only after the 25th passage of cell culture, whereas the young HDFs obtained from a 4-year-old boy in our study reached senescence after 75 doublings, implying that the reported cell condition was somewhat different from ours. We also observed that cells in the later stage of senescence (just before reaching cell death) contain much lower levels of lipogenic activity than well developed senescent cells, with maximum SA-β-galactosidase activity in both stress-induced and replicative senescence (data not shown). Interestingly, increased expression of lipogenesis-associated proteins was further found in the aging F344 rats, presenting the potential involvement of lipogenesis in the aging process. All of these reports and findings further support our hypothesis of the important contribution of enhanced lipogenesis to senescence and aging.

All mammalian cells require the biosynthesis of membrane lipids for the duplication of membranous organelles undergoing cell division. Therefore, further activation of lipogenesis is expected in cancer development, in which the strength of cell growth and proliferation is increased. Indeed, an increasing number of studies on the importance of lipogenesis in carcinogenesis have been reported, and lipid synthesis is regarded as a new target for the development of cancer therapeutics (37–39). However, in terms of cell proliferation, cancer development is the opposite situation to senescence. Therefore, the question of why enhanced lipogenesis leads cells to senescence and not to cell proliferation is important. When we overexpressed mature SREBP-1 protein, we clearly observed p21 and p16 proteins, the negative cell cycle regulators, in addition to the increased expression of lipogenic enzymes. Although protein induction of ACC, the rate-limiting enzyme of fatty acid synthesis, by SREBP-1 was minor, overall flow of fatty acid synthesis may be enhanced by modulation of substrate (acetyl-CoA) and product (malonyl-CoA) levels via the increased ACC and FAS proteins. In addition, the result of the inhibitory cell cycle regulators corresponds well to the previous report that mature SREBP-1 causes cell cycle arrest through the induction of the cyclin-dependent kinase inhibitors, such as p21, p27, and p16 (40, 41). Therefore, enhanced lipogenesis via SREBP-1 activation alone may act as an inducer of senescence. In other words, cancer cells should acquire any additional strategy to suppress the action of the cyclin-dependent kinase inhibitors that may be induced by SREBP-1 to use the lipogenic power for cell proliferation.

The questions then are why the senescent cells need an increased organellar mass and whether the organelles are functional. The simple explanations previously applied to lysosomes and mitochondria are possibly applicable to these questions. Senescent cells regenerate organelles to compensate for the nonfunctional organelles damaged by senescence-associated ROS as a defensive response, but the organelles that are supposed to remove the damaged organelles (components), such as lysosomes and proteasomes, have also lost their function, resulting in an accumulation of nonfunctional organelles (14, 42, 43). The worse scenario is that even newly synthesized organelles may be nonfunctional due to an oxidatively stressed cellular environment, aggravating or maintaining senescence. However, in question is what happens to the newly generated organelles in the senescence induced by overexpressed mSREBP-1 protein alone, in which no additional stress exists. So far, whether the organelles have normal function, structure, and well balanced composition, including membrane lipids and proteins, is not clear. Interestingly, knocking down SREBP-1 reduced the senescent intracellular ROS level, implying that lipogenesis is probably linked to oxidative stress. Therefore, we can hypothesize that enhancement of lipid biosynthesis alone, without supporting other components (for example, proteins) for organelles, may act as a stress by forming only lipid-enriched nonfunctional organelles. For example, in lipid-enriched mitochondria, respiratory complexes may be dispersed, making their electron transfer reaction defective and increasing ROS generation. However, a detailed understanding of the functional and compositional status of individual organelles in senescence remains unexplored.

SREBP-1 is known to regulate the supply of membrane lipids in response to cell growth and is highly expressed in cells that are actively growing (44). However, SREBP-1-expressing livers of transgenic mice exhibit impaired regeneration after partial hepatectomy (41), indirectly supporting our results that mSREBP-1 overexpression sufficiently induces senescence-associated growth arrest. We also showed that knocking down the up-regulated SREBP-1 in the stress-induced senescent cells effectively attenuated the senescence phenotypes, indicating the potential role of SREBP-1 as a novel effective target for modulating senescence. However, we also observed that knocking down SREBP-1 below the level of control cells by using high dose siRNA also induced growth arrest (data not shown), suggesting that the well balanced regulation of lipogenesis is required to maintain normal cellular function, and the fine control of SREBP-1 activation is essential to control senescence. In addition, our previous study proved that glycogen synthase kinase 3 (GSK3) is commonly inactivated by being phosphorylated and is also importantly involved in all the cell senescence systems (DFO- and H2O2-induced, and replicative senescence) employed in this study (27). GSK3 is known to modulate SREBP1 activity through phosphorylation-mediated ubiquitination (21). Therefore, it is suggested that GSK3 may play a critical role in activating SREBP-1 as an upstream modulator in senescence induced by stresses. A detailed link between GSK3 and SREBP-1 in senescence is under investigation. Collectively, our results provide new insight into the metabolic background of cellular senescence and indicate SREBP-1-induced lipogenesis as a potentially effective novel target for senescence and aging-related diseases.

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Enhanced Lipogenesis via SREBP-1 in Cell Senescence
Supplementary Figure S1.

A

MitoTracker
Red CMXRsos

Con | DFO | H₂O₂

Mean value
44.1

Mean value
110.6

Mean value
131.0

B

Lyso Tracker
Green DND-26

C

ER Tracker
Blue-White

D

BODIPY
C₅-Ceramide
Supplementary Figure S2.

A

B

C (a)

Cellular lipids (% control)

SREBP1

Full length

Mature form

Tubulin

H2O2: C 1 2 3 (d)

FAS

ACC

ACL

Tubulin

Lamin

H2O2: C 6 12 24 48 (h)

Expression ratio (Enzymes/tubulin)

Expression ratio (Enzymes/lamin)
Supplementary Figure S3.

A

CP
PA
CL
CA
PE

Std  PS  PC  5  10  20  40  5  10  20  40 (x10^4 cells)

HDF:  PD 15  PD 82

1
2
3
4
5
Supplementary Figure S4.

A

Chang Lamin FAS

Lamin ACC

Lamin ACL

100 25 50 100 (MOI)

GFP ad-mSREBP1

mRNA expression

B

HDF Lamin FAS

Lamin ACC

Lamin ACL

100 25 50 100 (MOI)

GFP ad-mSREBP1

mRNA expression
Supplementary figure legends

Supplementary Fig. S1. Representative fluorescence images for subcellular organelles of stress-induced senescent Chang cells. Chang cells were exposed to 200 µM DFO or H2O2 for 3 days. A. Fluorescence images (upper panel), phase images (middle) and flow cytometric analysis (lower panel) of MitoTracker-stained cells. Fluorescence images of LysoTracker-stained cells (B), ER Tracker-stained cells (C) and BODIPY C5-Ceramide-stained cells (D) are presented.

Supplementary Fig. S2. Increased lipogenesis in H2O2-induced senescent Chang cells. Chang cells were exposed to 200 µM H2O2 for 3 days. A. Cellular lipid profile of senescent cells was obtained by TLC as described in Experimental procedures. Quantitative estimation of cellular lipids is shown. Cholesteryl palmitate (CP, 1) belongs to non-polar storage lipid, and cholesterol (CL, 2), cardiolipin (CA, 3), phosphatidyl choline (PC, 4) and phosphatidyl serine (PS, 5) belong to membrane lipids. ‘d’ in the x-axis stands for day. B. Western blot for SREBP-1. Full length and mature form of SREBP1 were indicated by arrows. * is unidentified non-specific proteins. C. Western blot analyses for lipogenic enzymes (a). Expression ratios of ACC (■), ACL (▲), and FAS (◆) to tubulin (b) or lamin (c) are shown.

Supplementary Fig. S3. Representative TLC image for cellular lipid profiles of young and old HDFs. Cellular lipid profiles of young (PD 15, DT 2) and old (PD 82; DT 9) HDFs were compared by TLC as described in Experimental procedures and figure 2A.

Supplementary Fig. S4. Ectopic expression of mature SREBP-1 induces mRNAs of lipogenic enzymes. Chang cells or young HDFs (PD 24, DT 2) were infected with recombinant adenovirus encoding mature form of SREBP-1 (ad-mSREBP1) for 4 days. Recombinant adenovirus encoding GFP (ad-GFP) was used as control. Total RNA was isolated using Trizol (Invitrogen) and cDNA was prepared using avian myeloblastosis virus reverse transcriptase (Promega, Madison, WI, USA). PCR involved 25-30 cycles of 94°C for 30 seconds, 52–55°C for 30 seconds, and 72°C for 60 seconds. Oligonucleotide primers were produced by Bioneer (Seoul, Korea). The PCR primer sequences were: 5’-CCTCAACTATGCTAAGAAGGAATC-3’ and 5’-TTGGCATGTAAGTCTGCTC-3’ for lamin; 5’-TCAACCCAGATTCACTCCGAG-3’ and 5’-ATGGCCCTGGTAGGCTTC-3’ for FAS; 5’-CACGCTCAAGTCACAAGAA-3’ and 5’-CTGCAG GTTCTCAATGCAA-3’ for ACC; 5’-ATCTCCGGCCTCTTACTTT-3’ and 5’-GGTTCAAGGTCAGCTTC-3’ for FAS. A. mRNA levels by RT-PCR of Chang cell expressing mSREBP-1. B. mRNA levels by RT-PCR in HDFs expressing mSREBP-1.
Sterol Regulatory Element-binding Protein (SREBP)-1-mediated Lipogenesis Is Involved in Cell Senescence
You-Mie Kim, Hyun-Taek Shin, Yong-Hak Seo, Hae-Ok Byun, Soo-Han Yoon, In-Kyu Lee, Dong-Hoon Hyun, Hae-Young Chung and Gyesoon Yoon

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