Autophagy enhanced by curcumin ameliorates inflammation in atherogenesis via the TFEB–P300–BRD4 axis

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Abstract
Disturbance of macrophage-associated lipid metabolism plays a key role in atherosclerosis. Crosstalk between autophagy deficiency and inflammation response in foam cells (FCs) through epigenetic regulation is still poorly understood. Here, we demonstrate that in macrophages, oxidized low-density lipoprotein (ox-LDL) leads to abnormal crosstalk between autophagy and inflammation, thereby causing aberrant lipid metabolism mediated through a dysfunctional transcription factor EB

Acetyl-H3, acetyl-histone 3; ATG5, autophagy-related 5; BET, bromodomain and extra-terminal; BRD4, bromodomain protein 4; ChIP, chromatin immunoprecipitation; CQ, chloroquine; Cur, curcumin; CVDs, cardiovascular diseases; Dil-ox-LDL, 1,10-dioctadecyl-3,3,30,30-tetramethyl-indocarbocyanine perchlorate labeled oxidized low-density lipoprotein; FCs, foam cells; HFD, high-fat diet; IL-1β, interleukin 1β; LIR, LC3-interacting region; MCP-1, monocyte chemotactic protein 1; mTORC1, mammalian target of rapamycin complex 1; NAC, N-acetyl-L-cysteine; ORO, Oil red O; ox-LDL, oxidized low-density lipoprotein; qRT-PCR, quantitative real-time polymerase chain reaction; Re-ChIP, re-chromatin immunoprecipitation; ROS, reactive oxygen species; SE, super-enhancer; siRNAs, small interference RNAs; TFEB, transcription factor EB; TNF-α, tumor necrosis factor α.

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1. Introduction

As an underlying pathological process of life-threatening cardiovascular diseases (CVDs), the incidence of atherosclerosis is rising globally. Atherosclerosis has become a costly public health issue. Dysregulated of lipid catabolism and a maladaptive inflammatory response driven by the intramural retention of lipid-laden macrophages in susceptible areas of the arterial wall are the major contributors to atherosclerosis. Initially, plasma-derived lipoproteins that are sequestered in the subendothelial space following their modification are often engulfed by macrophages and balanced by reverse cholesterol transporters. Excess lipid forces the formation of foam cells (FCs) that contribute to the atherosclerotic lesion. Secretion of proinflammatory cytokines by FCs causes the plaque inflammatory state and defective effec-ctocytosis. FCs also exacerbate the formation of the vulnerable advanced lesion with a necrotic lipid core, whose rupture results in serious clinical complications. Thus, macrophages as the key integrators of inflammatory and metabolic signals have a pivotal role in atherogenesis. Efficiently ameliorating dysfunctional lipid catabolism and FCs inflammation may be a promising strategy for the regression of atherosclerosis.

Autophagy is an evolutionarily conserved cellular self-renewal mechanism that is critical in bulk degradation of aging or damaged cytoplasmic material via double-membrane autophagosomes with ensuing lysosomal fusion. Accumulating evidence indicates that adequate induction of autophagy delays atherogenesis through multiple mechanisms, including promoting lipid breakdown, inhibiting inflammation and oxidative stress. Impairment of autophagy usually occurs in advanced plaque. Injury of the lysosome by the lethal accumulation of cholesterol crystals contributes to plaque development and vulnerability. The resulting release of lysosomal hydrolyses aggravates apoptosis, leading to a vicious cycle. So far, in medications against atherosclerosis, only rapamycin or everolimus are working as the promotors of autophagy. Unfortunately, they have many side effects including cytokine production, dyslipidemia, and hyperglycemia. Transcription factor EB (TFEB) is an emerging regulator of the autophagy—lysosome pathway via binding to E-box sequences (CANNTG). The protective effects of TFEB have been widely confirmed on a variety of diseases, such as Alzheimer’s, diabetes, ischemic injury, as well as atherosclerosis from in vivo and in vitro models. No effective drug targeting TFEB in CVDs has been described.

Many natural products with excellent pharmacological properties are reported to be good candidates for the prevention of atherosclerosis. Curcumin (Cur) is a natural polyphenol extracted from turmeric curry spice. Cur has multiple biological properties and health benefits without corresponding toxicity. Numerous studies have confirmed the favorable therapeutic effects of Cur on a broad array of diseases. Although its role in the prevention of atherosclerosis has been shown, Cur’s anti-atherogenic mechanism is still unclear. Recently, Cur is reported to regulate autophagy by inhibiting the mammalian target of rapamycin (mTOR) activity in endothelial cells undergoing oxidative stress. Phosphorylation that is dependent on mammalian target of rapamycin complex 1 (mTORC1) controls TFEB nuclear export and inhibits autophagy in normal conditions. Whether Cur can delay atherosclerosis by activating autophagy via TFEB and overcome the side effects of traditional autophagy modulating medicines has yet not been investigated.

Atherosclerosis is characterized by abnormal autophagy and chronic inflammation of the arterial wall. However, the mechanism involved in the crosstalk between autophagy and inflammation is still unclear. An imbalance between the two processes may trigger pathologic atherogenesis. Bromodomain protein 4 (BRD4), a member of the bromodomain and extra-terminal (BET) family proteins, usually localizes at gene promoters and enhancers. BRD4 acts as a scaffold for transcription factors. BRD4 recognizing acetylated histones through bromodomains, regulates the chromatin landscape, and promotes the transactivation of pathologic genes. Interestingly, BRD4 has also been implicated in CVDs, including pulmonary arterial hypertension, heart failure and cardiac hypertrophy, due to the epigenetic activation of proinflammatory genes. Whether BRD4 is involved in the atherogenesis and whether Cur can delay atherosclerosis by a BRD4-dependent epigenetic regulatory mechanism has not been reported. Given the inhibitory effect of autophagy on inflammation, whether there is a link between the crucial autophagy regulator TFEB and the inflammation regulator BRD4 is also unclear.

The present study examined two closely related issues. First, the novel epigenetic mechanism was investigated to understand the abnormal crosstalk between autophagy and inflammation, which will cause aberrant lipid metabolism in FCs. Second, we sought to...
Figure 1 Curcumin (Cur) restores foam cells (FCs) autophagy and attenuates lipid metabolic dysfunction by promoting TFEB nuclear translocation. (A, B) Cell viability of FCs was measured by the cell counting kit 8 (CCK-8) assay. (C–E) Immunoblot analysis of LC3 II and P62.
identify an effective drug to treat and prevent atherosclerosis. Our experiments demonstrated that Cur, a natural P300-specific histone acetyltransferases inhibitor, regulates the novel epigenetic TFEB–P300–BRD4 axis. The findings explain the pro-autophagy-induced anti-inflammation behavior of Cur in the formation of FCs.

2. Materials and methods

2.1. Cell culture

Human monocytic THP-1 cells (American Type Culture Collection, Manassas, VA, USA) were cultured in RPMI 1640 medium with 10% fetal bovine serum supplemented with 20 μg/mL penicillin–streptomycin at 37 °C in a 5% CO₂ humidified atmosphere. To establish the FCs models, THP-1 cells were firstly differentiated into macrophages by 100 ng/mL phorbol-12-myristate-13-acetate (La Jolla, CA, USA) for 72 h at 1.0 × 10⁶ cells/mL, which were subsequently transformed to FCs induced by 50 μg/mL oxidized low-density lipoprotein (ox-LDL, Peking Union-Biology Co., Ltd., China) for 24 h. For the auto-phagy inhibition experiment, the cells were pre-incubated with 1 mM 3-MA in serum-free medium for 2 h and then incubated with ox-LDL.

2.2. Atherosclerosis model

Male Apoe knock-out mice (Apoe−/−, 8 weeks) on the C57BL/6 background were purchased from Sippr-BK Laboratory Animal Co., Ltd. (Shanghai, China). The mice were housed in a specific pathogen-free facility under controlled conditions (temperature, 22 ± 2 °C; relative humidity, 55 ± 15%; noise, < 60 dB; light/dark cycle, 12/12 h). Male Apoe−/− mice were fed with a high-fat diet (HFD, 21% fat and 0.15% cholesterol; Xietong Organism Inc., China) for 16 weeks to induce atherosclerosis. Mice in the Cur group received daily Cur (20 mg/kg body weight) by gastric gavage for 16 weeks together with HFD. All of the Apoe−/− mice consumed the HFD throughout the experiment. All experimental procedures in animals were conducted by the National Institute of Health Guide for the Care and Use of Laboratory Animals and approved by the Animal Care and Use Committee of Nanjing Medical University (IACUC-2103042), Nanjing, China.

2.3. Construct generation and lentiviral particle production

The recombinant lentiviral construct specifically expressing Brd4 or deleting Tfeb in macrophages was produced by molecular cloning methods. Firstly, the full-length Brd4 and Tfeb cDNA was obtained from murine peripheral blood mononuclear cells and amplified by reverse transcription-polymerase chain reaction (RT-PCR). Then, the PCR product was inserted into the self-inactivating lentiviral vector pLVX-Puro (Clontech, Mountain View, CA, USA). The macrophage-specific human CD68 promoter (2.9 kb) along with an 89 bp intronic enhancer were amplified from human genomic DNA by PCR. After deleting the cytomegalovirus promoter from the vector, the CD68 promoter was incorporated upstream of the coding sequence to generate the final Brd4-expressing lentiviral construct pLVCD68-Brd4 overexpression and Tfeb-deleting lentiviral construct pLVCD68-Tfeb knock-out (Tfeb KO). Then, the viruses were harvested from the medium of 293T cells transient co-transfected with pLVCD68-Brd4, pLVCD68-Tfeb or Lenti-X HTX Packaging Mix (Clontech, USA) according to the manufacturer’s instructions. Similarly, a recombinant lentiviral vector pLVCD68 that contains the CD68 promoter and lacks the Brd4 coding sequence or the Tfeb coding sequence was also generated and used as a control.

2.4. Bone marrow transplantation

Bone marrow cells (BMCs) from the femurs and tibias of donor male Apoe−/− mice were filtered through 30 μm nylon gauze to obtain the single-cell suspensions. Then, they were transfected by recombinant lentiviral pLVCD68-Brd4 oe or pLVCD68-Tfeb KO supernatant for 24 h at an MOI of 20 with 1 × 10⁶ TU/mL titers containing 4 μg/mL polybrene and 2000 U/mL recombinant human macrophage-colony stimulating factor (rh-CSF, R&D Systems, Minneapolis, MN, USA). Next, the cells were cultured for another 4 days in fresh medium with 2000 U/mL rh-CSF and harvested before injection. Lethally irradiated (9 Gy) the male recipient Apoe−/− mice transplanted with 5 × 10⁶ lentiviral transduced BMCs via tail vein injection within 6 h of irradiation. After 4 weeks, mice were fed with a HFD for an additional 16 weeks. The mice received BMCs transduced with pLVCD68 lentivirus used as control.

2.5. Atherosclerotic lesion analysis

The aortas were perfused with PBS and excised from the proximal aortic arch to the common iliac artery. Then, the connective and adipose tissues were removed from the aorta and the section of aortas was cut longitudinally stained with Oil red O (ORO) solution. Finally, the specimens were digitally photographed.

(F) Immunofluorescence analysis of LC3 (green) puncta formation in macrophages treated by oxidized low-density lipoprotein (ox-LDL) combined with Cur. Scale bar = 20 μm. (G) Immuno blot analysis of the autophagic flux in Cur-treated FCs pretreated with chloroquine (CQ). (H) Immunoblot analysis of Tfeb subcellular distribution in FCs. (I) Immunoblot of mTOR/p-mTOR, AKT/p-AKT, calcineurin, and Tfeb/p-Tfeb in Cur-treated FCs. (J) Immunoblot analysis of the effect of mTORC1-specific agonist L-leucine (0.5 mmol/L) on the activity of mTOR and Tfeb in Cur-treated FCs. (K, L) Immunoblot (K) and immunofluorescence (L) analysis of autophagy related proteins in Cur-treated FCs after siTfeb. Scale bar = 20 μm. (M, N) The subcellular distribution of Tfeb in Cur-treated FCs combined with 3-MA and siATG5 was detected by immunoblot (M) and immunofluorescence (N). Scale bar = 20 μm. (O, P) The lipid accumulation (O) and the lipid uptake capacity (P) of Cur-treated FCs combined with siTfeb or siATG5 was evaluated by Oil red O (ORO) staining and 1,1'-dioctadecyl-3,3',3'-tetramethyl-indocarbocyanine perchlorate labeled oxidized low-density lipoprotein (DiI-ox-LDL), separately. Scale bar = 100 μm. (Q) Immunofluorescence analysis of lipid catabolism in FCs through double fluorescence labeling with Bodipy (red) and LC3 (green). Scale bar = 20 μm. (R) Immunofluorescence analysis of lipid catabolism in Cur-treated FCs combined with siTfeb or siATG5 through double fluorescence labeling with Bodipy (red) and LC3 (green) at 24 h. Scale bar = 20 μm. Data are expressed as mean ± SEM, n = 5; *P < 0.05, **P < 0.001; ***P < 0.001.
Cur reduces inflammation in FCs by inhibiting P300 and BRD4 to alter the chromatin environment for inflammatory gene expression. (A, B) Relative mRNA levels of inflammatory genes, including IL-1β, IL-6, IL-8, TNF-α, and MCP-1 in ox-LDL-treated macrophages combined with Cur or DMSO.
For histological analysis, the section of aortas was fixed in 4% paraformaldehyde at 4 °C for 24 h, dehydrated in a graded series of alcohol (70%, 80%, 90%, 95% and 100%, each 90 min), embedded in 5 μm-thick paraffin. Then, they were stained with hematoxylin and eosin, ORO, Sirius red and Masson. Images were captured using a light microscope (Olympus BX53, Olympus Corp.) at magnification, 10×, 20× and 40×.

2.6. Cell viability analysis

The cell viability was evaluated by the cell counting kit 8 (CCK-8) assay (Beyotime, Beijing, China) according to the procedure. After different treatments, 100 μL 1640 medium containing CCK-8 at 10:1 dilution ratio was added to the cells incubating for 1 h in the dark. The survival cells were reflected by the absorbance of each well at 450 nm.

2.7. Cell transfection

Small interference RNAs (siRNAs) used to knock down human TFEB, autophagy-related 5 (ATG5) and BRD4 (GenePharma, Shanghai, China), as well as the plasmid used to overexpress BRD4, was imported into the macrophages in 35 mm cultures. Firstly, lipofectamine™ 3000 (Thermo Fisher Scientific, Madison, USA) were diluted in Opti-MEM medium (Thermo Fisher Scientific, Madison, USA). Then, the master mix of siRNA/DNA was prepared by diluting siRNAs/DNA in the Opti-MEM medium (Supporting Information Table S1). After that, the diluted siRNAs/DNA was added into diluted lipofectamine™ 3000 at 1:1 ratio and incubated for 15 min at room temperature (RT). Finally, the siRNA/DNA–lipid complex was added into cells incubating for 1–3 days. Immunoblotting was used to analyze the transfection efficiency of different siRNAs and plasmids.

2.8. Immunofluorescence

After different treatments, the cells seeded in glass-bottom cell culture dishes were rinsed in PBS followed by fixed in 4% paraformaldehyde and soaked in 0.1% Triton X-100 for 30 min, respectively. After blocked with 3% bovine serum albumin for 1 h, the cells were incubated with specific primary antibodies overnight at 4 °C. The cells were then incubated with appropriate TRITC/FITC-conjugated secondary antibodies (diluted in 1% bovine serum albumin) or Bodipy (Invitrogen, Carlsbad, CA, USA) for neutral lipid staining for 1 h at 37 °C. Finally, the cell nuclei were counterstained with DAPI. The sections were viewed under a confocal laser scanning microscope (LSM800, Zeiss, Oberkochen, Germany).

2.9. Immunoblot analysis

The whole-cell extracts were obtained in RIPA buffer containing protease and phosphatase inhibitors on ice. The nuclear and cytosolic fractions were obtained by the nuclear/cytosol fractionation kit (Beyotime, Beijing, China). Equal amounts of protein samples were separated in 6%–15% SDS-PAGE and transferred onto PVDF membranes (Millipore, Billerica, MA, USA) which was blocked in 5% skimmed milk for 2 h at RT. Then, the membranes were immunoblotted with specific primary antibodies at 4 °C overnight with slight agitation followed by HRP-conjugated secondary antibodies (Jackson Labs, USA) for 1 h at RT (Supporting Information Table S2). Finally, the immunoreactive bands were detected by the enhanced chemiluminescence method (Amersham Imager 600, GE, USA). According to the Guidelines for the use and interpretation of assays for monitoring autophagy, we evaluated the expression of autophagy by comparing LC3 II with actin.

2.10. CO-immunoprecipitation

After different treatments, the cells were lysed in lysis buffer (Beyotime, Beijing, China) and placed on a low-speed rotating shaker for 30 min at 4 °C. After centrifuged at 12,000 × g for 10 min, 50 μL of the whole cell lysates was loaded as input sample and the remaining cell lysates were incubated with the indicated antibody with protein A/G agarose beads at 4 °C overnight. The next day, beads were washed with lysis buffer for 5 times and boiled at 95 °C for 10 min. All samples were subjected to SDS-PAGE separation and immunoblotted with the indicated antibodies.

2.11. Quantitative real-time PCR (qRT-PCR)

Total cellular RNAs were isolated using Trizol (Takara, Japan) according to the manufacturer’s instructions and quantified by NanoDrop™ OneC (Thermo Fisher Scientific, Madison, USA), followed by reverse-transcribed into cDNA using HiScript® II Q RT SuperMix for qPCR (Vazyme, Jiangsu, China). qRT-PCR was carried out with cDNA in triplicate using AceQ® qPCR SYBR® Green Master Mix (Vazyme, China) on QuanStudio 5 Real-Time PCR System (Thermo Fisher Scientific, USA). Gene expression values were normalized against that of Actin. Data were analyzed with the 2^ΔΔCt method. Structures of all primers used are listed in Supporting Information Table S3.

2.12. Oil Red O lipid staining

The lipid accumulation in cells was analyzed by ORO (Sigma, St. Louis, MO, USA) staining. Firstly, the FCs were fixed in 4% paraformaldehyde followed by dehydrated with 60% isopropanol for 2 min at RT. Next, filtered 0.3% ORO solution was used to stain the lipid in FCs for 10 min at RT. Finally, DAPI

with Cur in the presence of siTFEB, siATG5, 3-MA, or CQ. (C, D) The expressions of BRD2, BRD3, BRD4, and actin in ox-LDL-treated macrophages combined with Cur were measured by immunoblot (C) and immunofluorescence (D). Scale bar = 100 μm. (E, F) Relative mRNA levels of inflammatory genes in ox-LDL-treated macrophages combined with Cur were measured by qRT-PCR (E). (F) BRD4 overexpression (oeBRD4) plasmid or P300 activator CTB (F). (G) Chromatin immunoprecipitation (ChIP) analysis of the enrichment of no-an antibody (NA), P300, BRD2, BRD3, BRD4 and H3 at −1 kb from the promoter, at the promoter, and at +1 kb from the promoter of inflammatory genes (IL-8, IL-1β, TNF-α) in Cur-treated FCs. Data are expressed as mean ± SEM, n = 5. *P < 0.05, **P < 0.01, ***P < 0.001; #P < 0.01, ##P < 0.001; $P < 0.05, $$$P < 0.01, $$$$P < 0.001.
Figure 3  Epigenetic landscape of P300-BRD4 dependent inflammatory genes in Cur treated FCs. (A) ChIP analysis of the enrichment of P300, BRD2, BRD3, BRD4 and H3 at −1 kb from the promoter, at the promoter and at +1 kb from the promoter of inflammatory genes (IL-8, IL-1β) in
was used to counterstain cell nuclei. After washing, they were photographed under an optical microscope and quantified by Image-pro plus.

2.13. Dil-ox-LDL uptake assay

The uptake capacity of ox-LDL in FCs after different treatments were tracked by 1,1'-dioctadecyl-3,3',3',3'-tetramethylyclorcobucyanine perchlorate labeled ox-LDL (Dil-ox-LDL, Yiyuan Biotech, Guangzhou, China). The FCs were loaded with 20 μg/mL Dil-ox-LDL for 6 h at 37 °C in the dark and then counterstained with DAPI to track the nuclei. Finally, the FCs were detected under a fluorescence microscope at 514/565 nm.

2.14. Reactive oxygen species (ROS) assay

The generation of intracellular ROS was detected by DCFH-DA staining. Briefly, the cells after different treatments were firstly washed with RPMI 1640 and then incubated with DCFH-DA (Beyotime, Beijing, China) at 1:1000 ratio for 30 min at 37 °C in the dark. After washing, the fluorescent signal was measured by fluorescence microscope at 488/525 nm.

2.15. Chromatin immunoprecipitation (ChIP) and re-chromatin immunoprecipitation (Re-ChIP)

ChIP was performed as previously described19. Briefly, the cells grown in 15-cm dishes were cross-linked in 1% formaldehyde at RT for 10 min after different treatments, quenched in 0.125 mol/L glycine for 5 min and collected by centrifugation followed by sonication in SDS lysis buffer to generate ~500 bp chromatin fragments. The lysates were centrifuged at 4 °C for 10 min and 20 μL supernatant was used as input for quantitation. The left supernatant was diluted 10 times with dilution buffer and precleared in 20 μL protein A–agarose beads at 4 °C for 3 h. Subsequently, the supernatant was incubated with indicated antibodies at 4 °C with rotation overnight followed by incubating with 40 μL protein A–agarose beads at 4 °C for 2 h. The Txn stop buffer with 0.4 mg/mL glycogen and proteinase K (0.45 mg/mL, Roche, USA) was used to digest protein in immunoprecipitated DNA at 37 °C for 1 h. After that, DNA was isolated in phenol/chloroform/isoamyl alcohol (25:24:1) and precipitated in ethanol. For Re-ChIP, immunoprecipitated complexes were eluted with the elution buffer (1% SDS, 100 mmol/L NaCO3), diluted with the Re-ChIP buffer (1% Triton X-100, 2 mmol/L EDTA, 150 mmol/L NaCl, 20 mmol/L Tris pH 8.1), which was subjected to immunoprecipitation with BRD4 and MED1 antibody. Finally, the enrichment of immunoprecipitated material relative to input with gene-specific primers to the specified regions was determined by qRT-PCR (Supporting Information Table S4).

2.16. Chromatin conformation capture assay-qPCR (3C-qPCR)

After different treatments, the cells were cross-linked in 1% formaldehyde at RT for 10 min, quenched in 0.125 mol/L glycine for 5 min and centrifuged at 1000 × g for 10 min at 4 °C. Then, the pellets were lysed in lysis buffer for 10 min on ice and centrifuged to get nuclei, which were digested in restriction enzyme buffer containing 0.3% SDS for 1 h at 37 °C. Triton X-100 was added to the buffer for 1 h at 37 °C followed by 400 U XbaI incubation overnight to digest DNA. Next, the nuclei were incubated in 1.6% SDS for 25 min at 65 °C, suspended in ligation buffer containing 1% Triton X-100 for 1 h at 37 °C, then ligated by 100 U ligase for 4 h at 16 °C followed by 30 min at RT. Reaction was terminated, and de-crosslinking was performed with PK buffer (5 mmol/L EDTA, pH 8.0; 10 mmol/L Tris—HCl, pH 8.0; 0.5% SDS) at 65 °C overnight. After digested in 300 μg RNase for 45 min at 37 °C, DNA was isolated in phenol/chloroform/isoamyl alcohol (25:24:1) and precipitated in ethanol. 3C-DNA was amplified by qPCR. BAC 3C DNA was used to determine the PCR efficiency for each set of primers. The actin promoter was used as a negative control.

2.17. 1,6-Hexanediol disrupts BRD4 and MED1 phase separation

The experiments were performed as previously described20. THP-1 cells seeded in glass-bottom cell culture dishes and treated with 3% hexanediol for 15 s after ox-LDL stimulation for 24 h. Immunofluorescence staining with BRD4 and MED1 and the cell nuclei were counterstained with DAPI. The sections were viewed under a confocal laser scanning microscope (LSM800, Zeiss, Oberkochen, Germany).

2.18. Statistical analysis

Results obtained are presented as mean ± standard error of mean (SEM) based on five independent experiments. Differences among the treatment groups were assessed with one-way ANOVA followed by Tukey’s test for comparison when appropriate. Unpaired two-tailed Student’s t test was used for comparisons between two groups. Statistical significance was defined as *P < 0.05. Statistical analyses were performed with GraphPad Prism 8.0 software.

3. Results

3.1. Cur restores FCs autophagy and attenuates lipid metabolic dysfunction by promoting TFEB nuclear translocation

To determine whether Cur protects macrophage-derived FCs against ox-LDL injury, we initially used the CCK-8 assay to detect cell viability. As shown in Fig. 1A, the cell viability of macrophages decreased significantly in the process of differentiation

ox-LDL-treated macrophages combined with JQ-1 or siBRD4. (B, C) ChIP analysis of the enrichment of P300, BRD2, BRD3, BRD4 and H3 at −1 kb from the promoter, at the promoter and at +1 kb from the promoter of inflammatory genes (IL-8, IL-1β) in ox-LDL-treated macrophages combined with Cur in the presence of oeBRD4 (B) or CTB (C). (D) Immunoblot analysis of the expressions of P300, BRD4, actin, acetyl-H3 and H3 in Cur-treated FCs in the presence of oeBRD4 and CTB. Data are expressed as mean ± SEM, n = 5. *P < 0.05, **P < 0.01, ***P < 0.001; *P < 0.05, ***P < 0.001; ns, no significance.
Figure 4  Cur inhibits the P300–BRD4 pathway by attenuating ROS generation through TFEB-activated autophagy. (A) ChIP analysis of the enrichment of P300, BRD2, BRD3, BRD4 and H3 at −1 kb from the promoter, at the promoter and at +1 kb from the promoter regions of...
into FCs and was restored by Cur in a dose-dependent manner. At the same time, there was no obvious difference in the cell viability of FCs treated with 20 μmol/L Cur for 0, 6, 12, or 24 h (Fig. 1B). The cell viability of FCs decreased when the cells were treated with 20 μmol/L Cur for 48 h, which was consistent with previous report. Therefore, FCs treated with 20 μmol/L Cur for 24 h were subsequently used.

Overloaded lipid accumulation usually causes autophagy deficiency, which exacerbates the pathogenesis of atherosclerosis. To evaluate the effect of Cur on the autophagy of FCs, immunofluorescence and immunoblot analyses were performed to detect the expression levels of autophagy-specific markers. Upon autophagy induction, microtubule-associated protein 1 light chain 3-I (MAP1LC3/LC3 I) is converted to LC3 II through lipidation, promoting the formation of autophagosomes. Meanwhile, sequestosome-1 (SQSTM1)/P62 targets damaged cytoplasmic material for lysosomal degradation. The results reveal that macrophage autophagy was downregulated by ox-LDL as evidenced by the reduced expression of LC3 II along with the increased P62 expression (Fig. 1E and Supporting Information Fig. S1C). However, they were reversed by Cur in concentration- and time-dependent manners (Fig. 1C and Fig. S1A, Fig. 1D and S1B). Similarly, compared with untreated cells, immunofluorescence examination revealed that the number of LC3 puncta was decreased in FCs which was restored by Cur (Fig. 1F and Fig. S1D). Therefore, FCs incubated with 20 μmol/L Cur for 24 h is chosen as the optimal condition.

During the process of autophagy, the encapsulated contents as well as LC3 II and P62 in autophagosomes are usually degraded by fusing with lysosomes, which is called autophagic flux. To examine the Cur-induced autophagy, FCs were pretreated with a lysosomal inhibitor (chloroquine, CQ, 20 μmol/L, 1 h) to prevent the fusion and subsequent proteolysis. Compared with Cur-treated FCs, the expression of LC3 II and P62 were further increased in Cur combined with CQ (Fig. 1G and Fig. S1E). There was no further significant increase of LC3 II and P62 in FCs in response to CQ, suggesting a defect in autophagic flux in FCs. The collective findings indicate that Cur restores the autophagy level of FCs by promoting the formation of both autophagosomes and autophagic flux.

To detect whether TFEB was involved in Cur-induced autophagy in FCs, we assessed the cellular distribution of TFEB. Cur did not affect total TFEB expression. While the nuclear levels of TFEB gradually increased and reached the most at 24 h after Cur, the cytosol levels of TFEB presented the opposite trend (Fig. 1H and Fig. S1F). In most cases, mTORC1-dependent phosphorylation controls TFEB nuclear export and inhibits autophagy in the normal condition. TFEB stability can also be regulated by other kinases, such as AKT and calcineurin. We confirmed that Cur did not affect the expression of calcineurin and phosphorylated AKT (p-AKT) in FCs. Nevertheless, phosphorylated mTOR (p-mTOR; Ser2448), as the indicator of mTORC1 activation, and its downstream effector p-S6K1 were downregulated in Cur-treated FCs along with p-TFEB (Fig. 1I and Fig. S1G, Fig. 1K and Fig. S1I). The decreased expression of p-TFEB caused by Cur was largely recovered when co-treated with the mTOR specific activator L-leucine (Fig. 1J and Fig. S1H), indicating that Cur effectively promoted TFEB nuclear translocation in FCs by inhibiting mTOR activity. Furthermore, TFEB deficiency resulting from treatment with small interfering RNA (siRNA) markedly inhibited the autophagy activation in Cur-treated FCs (Fig. 1K, Fig. S1I and Supporting Information Fig. S2A, Fig. 1L and Fig. S1J). However, autophagy inhibitor 3-methyladenine (3-MA) or autophagy deficiency via siATG5 had no effect on Cur-induced TFEB nuclear translocation in immunoblot analysis (Fig. 1M, Fig. S1K and Fig. S2B) and immunofluorescence (Fig. 1N). These data demonstrate that Cur restores FCs autophagy mainly by activating TFEB nuclear translocation.

Cur could inhibit the decrease of ATG5 mRNA in macrophages caused by ox-LDL, which was reversed by siTFEB (Fig. 2C). Moreover, Cur increased the binding of TFEB to the ATG5 promoter regions as observed by ChIP (Fig. 2D). However, there was no protein interaction between TFEB and ATG5 (Fig. 2E). To further confirm whether TFEB influences the open state of chromosomes, ChIP analysis was performed to detect the open (active) and condensed chromatin landmarks. Open chromatin is generally marked by H3K4me1 and H3K27ac. Transcriptionally repressed (condensed) chromatin is generally marked by H3K9me3 and H3K27me3. Cur inhibited the binding of H3K9me3 and H3K27me3 to the ATG5 promoter regions and increased the binding of ATG5 promoter regions, which could be reversed by siTFEB (Fig. 2F). These results suggest that Cur could promote the open state of ATG5 gene located chromatin by increasing TFEB nuclear translocation.

Cur reportedly enhances lipid degradation in FCs. The underlying mechanism is unclear. Whether the effect of Cur on lipid degradation depends on TFEB-related autophagy was assessed. As shown in Fig. 1O and Fig. S1L, Cur apparently eliminated the lipid accumulation in FCs as stained by ORO. A binding assay using Dil-ox-LDL shows that Cur diminished the lipid uptake of FCs as evidenced by decreased red fluorescence intensity (Fig. 1P and Fig. S1M). At 24 h of treatment, Cur also resulted in an increased number of LC3-positive autophagosomes along with a decreased number of Bodipy-positive lipid. The findings suggest that Cur promoted the lipid breakdown of FCs (Fig. 1Q). All of the above changes induced by Cur in FCs were inhibited by siTFEB and siATG5 (Fig. 1O, P, and R). Therefore, Cur decreased the lipid accumulation in FCs by decreasing lipid uptake and increasing lipid catabolism.

3.2. Cur reduces inflammation in FCs by inhibiting P300 and BRD4 to alter the chromatin microenvironment for inflammatory gene expression

Proinflammatory cytokines participate in immune-mediated plaque formation and vulnerability. As shown in Fig. 2A and B, the increased mRNA levels of inflammatory cytokines, including
Figure 5  Cur regulates the chromatin open state in FCs in a BRD4-dependent manner. (A) ChIP-seq peaks (H3K27ac levels) and putative super-enhancer (SE) location (highlighted by the red box) around IL-8. Cell lines are marked with different colors, with K562 in blue, CD14-
interleukin 1β (IL-1β), interleukin 6 (IL-6), interleukin 8 (IL-8), tumor necrosis factor α (TNF-α), and monocyte chemotactic protein 1 (MCP-1) induced by ox-LDL were markedly suppressed by Cur. The anti-inflammatory effects of Cur on FCs were also eliminated by TFEB and autophagy deficiency. These results indicate that Cur attenuated inflammation in FCs by enhancing TFEB-related autophagy.

The epigenetic reader protein BRD4 functions as a critical coactivator of the transactivation of inflammatory-genes during aerol photographic-induced inflammation and heart failure<sup>7,28</sup>. To detect whether BRD4 is involved in the activation of inflammation in FCs, we first examined the expressions of the BET family genes, including BRD2, BRD3, and BRD4. Immunoblot and immunofluorescence analyses revealed that BRD4 was increased in FCs, which could be markedly inhibited by Cur, while BRD2 and BRD3 did not change (Fig. 2C, D and Supporting Information Fig. S3A). Furthermore, ox-LDL-induced inflammation in macrophages was substantially diminished when BRD4 was suppressed by siRNA or the inhibitor JQ-1 (Fig. 2E and Fig. S2G). The findings indicate the involvement of BRD4 in the activation of inflammation in FCs. The histone acetyltransferases CBP/P300 can facilitate the recruitment of BRD4 to activate enhancers and to control cell identity gene induction in adipogenesis and myogenesis<sup>27</sup>. In the present study, the activity of P300 reflected by the expression of acetyl-histone 3 (acetyl-H3) was also increased by ox-LDL, which was inhibited by Cur (Fig. 2C). ox-LDL also effectively increased the acetylated levels of H3K9 and H3K27, rather than those of H3K14, H3K18, and H3K23. The increases were reversed by the P300 inhibitor C646 (Fig. S3B). These results suggest that the specific site of P300 acetylation on FC histones is H3K9 and H3K27. Furthermore, plasmid-mediated enforced overexpression of BRD4 (oeBRD4, Fig. S2H) and the P300 activator CTB markedly repressed the anti-inflammatory effect of Cur on FCs (Fig. 2F). The collective results indicate that BRD4 and P300 are positive regulators involved in the inflammation of FCs, whose activities could be inhibited by Cur.

ChIP of the promoter elements (~1 to +1 kb) of inflammatory genes was used to confirm the mechanism underlying the regulation of P300 and BRD4 on FC-mediated inflammation. We systematically used ChIP to probe for BRD2, BRD3, BRD4, P300, and H3 (positive control) and no-antibody control (NA). No significant changes were evident in NA, BRD2, and BRD3 concerning IL-8, IL-1β, TNF-α, IL-6, and MCP-1 in each treatment group (Fig. 2G and Fig. S3C). Strikingly, the promoter regions of inflammatory genes were enriched for BRD4 and P300 in ox-LDL-treated macrophages, which was inhibited by Cur. The control marker H3 was pulled down evenly. The positive effects of Cur on lipid catabolism in vitro were all reversed by BRD4 overexpression (Fig. S3D–S3F). The collective results implicate BRD4 and P300 were involved in the control of the chromatin status to transactivate inflammatory genes, thereby promoting lipid accumulation. The decreased expression of LC3 II in FCs was rescued by siBRD4, consistent with the previous observation that BRD4 is also a transcriptional repressor of autophagy (Fig. S3G).

3.3. Epigenetic landscape of P300—BRD4-dependent inflammatory genes in Cur treated FCs

We next analyzed whether the effect of BRD4 on inflammation in Cur-treated FCs relies on P300 activity. BRD4 deficiency in FCs reduced the enrichment of BRD4 at the promoter regions of these inflammatory genes, while having no effect on P300 (Fig. 3A and Supporting Information Fig. S4A). ChIP data of IL-8, IL-1β, IL-6, TNF-α and MCP-1 revealed that the inhibitory effect of Cur on the binding of BRD4 and P300 to the promoter regions of inflammatory genes in FCs was reversed by oeBRD4 and the P300 activator CTB (Fig. 3B and Fig. S4B, Fig. S4C). However, CTB and oeBRD4 did not affect the expression or the activity of each other (Fig. 3D). CTB promoted increased binding of BRD4 to the promoter regions of the inflammatory genes in the presence of ox-LDL (Fig. 3C). In contrast, oeBRD4 had no effect on the binding of P300 (Fig. 3B). Consistently, P300 cooperates with the downstream histone acetylation reader BRD4 to achieve transcriptional activation<sup>30</sup>. Thus, Cur could reduce inflammation in FCs by inhibiting the P300—BRD4 pathway.

3.4. Cur inhibits the P300—BRD4 pathway by attenuating ROS generation through TFEB-activated autophagy

Cur is reportedly an inhibitor of P300. However, the detailed mechanism is unclear. Interestingly, we observed that Cur reduced BRD4 and P300 enrichment in TFEB oriented chromatin, which was abrogated by TFEB deficiency (Fig. 4A and Supporting Information Fig. S5A), as are the expressions of acetyl-H3 and BRD4 (Fig. 4B). The observations prompted us to consider how Cur decreases BRD4 expression in FCs and whether the regulation of TFEB on P300 activity relies on the activation of autophagy.

Two protein degradation systems, including ubiquitin–proteasome and autophagy systems, have been described<sup>31</sup>. We first used the proteasome inhibitor MG132 to explore the degradation pathway of BRD4 in Cur-treated FCs. The decreased expression of BRD4 in Cur-treated FCs was not changed by MG132 (Fig. S5B). This result prompted us to explore whether Cur decreased BRD4 expression via autophagy. Scrutiny of the iLIR autophagy database revealed an LC3-interacting region (LIR) motif in BRD4, which ensures targeting of autophagy receptors to LC3 anchored in the phagophore membrane (Fig. S5C). The LIR sequence is presented in Fig. S5D. Use of the autophagy flux inhibitor CQ revealed that the decrease of BRD4 in Cur-treated FCs was restored by CQ (Fig. S5E), suggesting that the BRD4 was degraded by autophagy in Cur-treated FCs. The interaction between BRD4 and LC3 in Cur-treated FCs, which was further increased in the presence of CQ (Fig. S5F),
Figure 6   Cur regulates BRD4-dependent SE associated with phase separation on inflammatory genes. (A) A color-coded schematic representation of the aligned amino acid sequences and corresponding prion-like domain disorder propensity plots for BRD4 and MED1 generated
Collectively, these findings confirm that Cur could degrade BRD4 in FCs by autophagy activation.

The inhibitory effect of Cur on P300 activity was attenuated in the presence of siATG5 (Fig. 4C), indicating that Cur can decrease P300 activity via TFEB-mediated autophagy activation. ROS could upregulate P300 activity-dependent CD36 expression and promote FCs formation \(^7\). Increased activation of autophagy can effectively decrease ROS generation through mitophagy \(^6\). We observed that the increased P300 activity mediated by ox-LDL was blocked by the ROS scavenger N-acetyl-l-cysteine (NAC) (Fig. 4D). This finding suggests the involvement of ROS in P300 activation during FC formation, which is consistent with previous studies. Moreover, Cur substantially inhibited the generation of ROS as indicated by DCFH-DA fluorescence staining, which was reversed by siTFEB or siATG5 (Fig. 4E). Over the course of exploring the basis of the ROS-mediated increase of P300 activity in FCs, we observed that phosphorylation of non-receptor tyrosine kinase BTK increased in ox-LDL-treated macrophages, which was inhibited by the ROS scavenger NAC (Fig. S5G). Furthermore, the increased acetylation of total histone in ox-LDL-treated macrophages was inhibited by BTK antisense oligonucleotides (ASO, Fig. S5H). These results confirm that ox-LDL resulted in ROS production, which in turn activated BTK, followed by increased P300 activity in macrophages.

The collective data suggest that ROS is an activator of P300 that promotes the binding of BRD4 to the promoters of inflammatory genes. This binding can be inhibited by TFEB-induced autophagy upon treatment with Cur.

### 3.5. Cur regulates chromatin open state in FCs in a BRD4-dependent manner

For the well-known proinflammatory cytokines IL-8 and IL-1β, the putative super-enhancer (SE) in blood predicted by the comprehensive human SE database was denoted as a horizontal line in blue in Fig. 5A and Supporting Information Fig. S6A \(^4\). In this putative SE regions strong signals of active SE marked by high levels of H3K27ac were found to be located in proximal to the IL-8 and IL-1β promoters. The corresponding Chip-seq data from three human blood cell lines (K562 \(^3\), CD14-positive monocyte \(^6\) and THP-1 \(^5\)) were displayed using the Integrative Genomics Viewer \(^8\) (Fig. 5A and Fig. S6A). To confirm the interaction between the promoters of IL-8 and these potential enhancer clusters as well as IL-1β, we performed a quantitative analysis of chromosome conformation capture assays (3C-qPCR). The primers for promoter and enhancer locus 2 in IL-8 and locus 4 in IL-1β showed the greatest interaction strength and strongest response to Cur, JQ-1, siBRD4, and P300 inhibitor C646 (Fig. 5B and Fig. S6B). An SE that strongly enriches for the H3K27 acetylation (H3K27ac), BRD4, and the mediator complex 1 (MED1) is comprised of clusters of transcriptional enhancers \(^9\). The IL-8 and IL-1β promoters were strongly enriched for BRD4, MED1, and H3K27ac in FCs, which were inhibited by JQ-1, siBRD4, or Cur (Fig. 5C and Fig. S6C). Re-ChIP assay results demonstrate that BRD4 and MED1 occupied the same DNA element of IL-8 and IL-1β in FCs, which was inhibited by Cur or siBRD4 (Fig. 5D and Fig. S6D). ox-LDL upregulated the expressions of BRD4, MED1, H3K27ac and H3K4me1 and decreased H3K27me3 expression. The findings suggest that ox-LDL contributed to the opening state of chromatin, which was reversed by Cur or siBRD4 (Fig. 5E). The collective findings indicate that ox-LDL promotes the enrichments of BRD4 and MED1 in the promoter regions of inflammation genes to form SE and thereby enhance the transactivation of inflammation, which can be inhibited by Cur.

### 3.6. Cur regulates BRD4-dependent SE associated with phase separation on inflammatory genes

Based on the latest research on phase separation published in Science \(^20\), we investigated the sensitivity of BRD4 and MED1 to 1,6-hexanediol, a compound known to disrupt liquid-like condensates. The architectural features of BRD4 and MED1 that are prone to phase separation are shown in Fig. 6A. Compared with macrophages, a dramatic increase was evident in the number of BRD4 and MED1 puncta in the nuclei of FCs. The increase was reversed by 1,6-hexanediol (Fig. 6B). 1,6-Hexanediol inhibited the greatest interaction strength between promoter and enhancer locus 2 in IL-8 and locus 4 in IL-1β (Fig. 6C and Fig. S6E), the binding of BRD4 and MED1 to the SE region of IL-8 and IL-1β (Fig. 6D and Fig. S6F), and the upregulation of IL-8 and IL-1β mRNA (Fig. 6E and Fig. S6G) in FCs. The increased BRD4 and MED1 puncta on macrophages treated with the phase separation activator PEG were also inhibited by Cur or the P300 inhibitor C646 (Fig. 6F and J). PEG increased the interaction between the IL-8 promoter and the enhancer locus 2, as well as the IL-1β promoter and the enhancer locus 4 (Fig. 6G and Fig. S6H). PEG also increased BRD4 and MED1 binding to the SE region of IL-8 and IL-1β (Fig. 6H and K, Fig. S6I and S6K). These reactions were inhibited by both Cur and C646, which were evident at the mRNA level for IL-8 and IL-1β (Fig. 6I and L, Fig. S6J and S6L). The increased BRD4 and MED1 puncta induced by ox-
Figure 7  Specific ectopic expression of BRD4 and TFEB in macrophages reduces the protective effect of Cur in Apoe<sup>−/−</sup> mice. (A) Immuno blot analysis of BRD4 in aortas from Apoe<sup>−/−</sup> mice fed with the normal chow (NC) and high-fat diet (HFD). All of the Apoe<sup>−/−</sup> mice were fed
Curcumin prevents atherosclerosis via inhibiting inflammation through TFEB—P300—BRD4 axis

**Figure 8** Proposed model of the autophagy enhanced by Cur ameliorates inflammation in atherogenesis via the TFEB—P300—BRD4 axis. ox-LDL leads to the autophagy deficiency and ROS production in macrophages. This resulted in the activation of P300 which promotes increased binding of BRD4 to the promoter regions of inflammatory genes. This consequently contributes to inflammation which leads to the formation of FCs. Cur can significantly increase autophagy activity in FCs by promoting TFEB nuclear translocation through the inhibition of mTOR. This optimizes lipid catabolism. Autophagy can effectively attenuate the ROS level and P300 activity. The decreased acetylated histone resulting from the suppression of P300 lessens the attraction of BRD4 to the promoter regions of inflammatory genes in FCs and finally attenuates the inflammation. Especially, Cur inhibits the formation of BRD4-dependent SE of the inflammatory genes associated with phase separation.

LDL was significantly inhibited by Cur (Fig. 6M). The collective findings indicate that Cur regulates BRD4-dependent SE associated with phase separation on inflammatory genes.

3.7. Specific ectopic expression of BRD4 and TFEB in macrophages abolishes the protective effect of Cur in Apoe<sup>−/−</sup> mice

Compared with Apoe<sup>−/−</sup> mice fed with normal chow (NC), the expression of BRD4 of aortas was increased in Apoe<sup>−/−</sup> mice fed with a HFD (Fig. 7A). Macrophage-specific Brd4 over-expressing Apoe<sup>−/−</sup> mice were constructed through bone marrow transplantation. C57BL/6 bone marrow cells transduced with either pLVCD68-Brd4 overexpression (pLVCD68-Brd4 oe) or pLVCD68 vector were transplanted into the Apoe<sup>−/−</sup> mice. After consuming the HFD for 16 weeks, atherosclerotic plaques were assessed. There was no difference in survival rate among different groups of mice (Supporting Information Fig. S7A). We confirmed that the mRNA level of Brd4 increased markedly in mice transplanted with pLVCD68-Brd4 oe-transduced BMCs and was partially decreased in the presence of Cur (Fig. 7B). Compared with the mice that received pLVCD68 + DMSO, the lesion area and lipid content of aortas from the mice that received pLVCD68 + Cur significantly decreased as assessed by morphometric analyses of lesion area in the aorta en face (Fig. 7C, Fig. S7B) and cross-sections of the aortic arch (Fig. 7D, Fig. S7C). Furthermore, compared with mice treated with pLVCD68 + DMSO, there was a trend toward increased plaque collagen content measured by Sirius red and Masson staining in plaques (Fig. 7D, Fig. S7C), and decreased inflammation of arteries from mice treated with pLVCD68 + Cur (Fig. 7E). No difference in blood pressure was evident in all groups of mice (Fig. S7D). However, the protective effect of Cur on atherosclerosis was significantly inhibited in mice transplanted with pLVCD68-Brd4 oe-transduced BMCs (Fig. 7C–E). Cur effectively decreased the plasma lipids in HFD-fed Apoe<sup>−/−</sup> mice that had received a bone marrow transplant from pLVCD68 macrophages. The decrease was reversed by macrophage-specific Brd4 over-expression (Fig. S7E).

To further demonstrate the role of Tfeb in the protective effect of Cur on atherosclerosis, we first assessed the Tfeb activity. In aortas of HFD-fed Apoe<sup>−/−</sup> mice, the phosphorylation level of TFEB was decreased by Cur treatment, while the total TFEB protein expression did not change between groups (Fig. 7F and Fig. S7F). We constructed the macrophage-specific Tfeb knock-out (KO) Apoe<sup>−/−</sup> mice by bone marrow transplantation. qRT-PCR confirmed the marked decrease in the mRNA level of Tfeb in mice transplanted with pLVCD68-Tfeb-KO-transduced BMCs (Fig. 7G). There was no difference in survival rate in all groups (Fig. S7G). The protective effects of Cur on the lesion area, lipid content, plaque collagen, inflammation, and plasma lipids were all inhibited in mice transplanted with pLVCD68-Tfeb-KO-transduced BMCs (Fig. 7H–J, Fig. S7H, S7I and S7K). No difference was evident in blood pressure in all groups of mice (Fig. S7J). The increased ROS generation and LC3 expression, and decreased expression of BRD4 in Cur-treated Apoe<sup>−/−</sup> mice were also reversed in macrophage-specific Tfeb KO mice (Fig. 7K and L, Fig. S7L). We also confirmed that the decreased P300 activity in the presence of Cur was also reversed in mice with macrophage-specific Tfeb KO (Fig. 7L and Fig. S7L). The
collective findings demonstrate that Cur could effectively enhance autophagy in FCs via TFEB and decrease P300 activity and BRD4 expression.

4. Discussion

In this study, we observed that ox-LDL led to abnormal crosstalk between autophagy and inflammation through the TFEB–P300–BRD4 axis. A proposed model is presented in Fig. 8. We also shed light on the mechanism of how Cur benefits atherosclerosis by promoting autophagy and thereby inhibiting inflammation in the FCs. These findings implicate Cur as a potential therapeutic agent for atherosclerosis with several novel drug targets identified.

Our results show that Cur markedly ameliorated autophagy deficiency in FCs and increased the autophagic flux. Therefore, Cur has a protective role in atherosclerosis. TFEB is a novel master molecule of the autophagy–lysosome pathway. Normally, TFEB is localized on the lysosome membrane in an inactivated form via phosphorylation by mTORC1. Interestingly, Cur has been reported to ameliorate intestinal barrier injury via AMPK–TFEB-dependent mitophagy. The Cur derivative Cur20 attenuates cerebral ischemic injury via HIF-1alpha/VEGF/TFEB-dependent angiogenesis. Another Cur analog could induce autophagy by directly binding to and activating TFEB in an mTOR-independent manner to treat neurodegenerative diseases. However, the TFEB response upon Cur treatment in FCs has never been reported. Here, we demonstrate that Cur induced the time-dependent redistribution of TFEB in the nuclei of FCs and corresponding activation of autophagy. These events were completely inhibited by silencing TFEB. Pharmacological inhibition of autophagy had little effect on TFEB nuclear translocation, implicating TFEB as an upstream regulator of autophagy. The expression of p-mTOR (Ser2448) in FCs was downregulated by Cur and was partly reversed by TFEB silence, while other previously reported TFEB activity regulators like AKT and calcineurin were not affected in FCs. The collective findings support the viewpoint that Cur significantly improves the autophagy of FCs by promoting mTORC1-dependent TFEB nuclear translocation. Furthermore, Cur could inhibit the binding of H3K9me3 and H3K27me3 to the TFEB promoter region and increase the H3K4me1 and H3K27ac binding of to the ATG5 promoter region. These events can be reversed by siTFEB. These results suggest that Cur could influence the open state of chromosomes by increasing TFEB nuclear translocation. In addition, Cur led to an increase in the content of autophagosomes and decreased Bodipy fluorescence in FCs, as well as the reduced lipid uptake and accumulation. All of the above events were reversed by TFEB or ATG5 deficiency, suggesting that TFEB nuclear translocation and subsequently autophagy activation are crucial in Cur-mediated promotion of lipid degradation.

Inflammation is causally related to atherogenesis. Overexpression of inflammatory cytokines may trigger activation of nuclear transcription factors like TFEB, which can translocate into the nucleus to regulate gene expression. As a potential natural anti-inflammation product, Cur can inhibit the inflammation by fine-tuning the TFEB- and ATG5-related inflammatory responses.

Recently, epigenetic regulation has been found to play a critical role in gene expression via post-translational modification of core histones in the genome, resulting in alternative chromatin configuration and altered gene transcription. The expression of inflammatory genes is tightly controlled by chromatin “readers” that specifically bind acetylated histones and provide a scaffold for sequence-specific transcription factors that comprise the transcriptional activation complex. Moreover, BET proteins as “readers” of histone acetylation have been demonstrated to mediate the inflammatory response by remodeling the communication between enhancer and RNA polymerase in heart failure and pulmonary artery hypertension. Whether BET families take part in the epigenetic regulation of inflammation in FCs has never been reported. Amazingly, we found that the expression of BRD4, but not BRD2 and BRD3, is dramatically increased in FCs. These events can be prevented by Cur. Consistent with the report that BRD4 acts as a transcriptional repressor of autophagy, the inhibition of autophagy in FCs was partially rescued by siBRD4. The anti-inflammation effect of Cur in FCs could be reversed by BRD4 overexpression. Additionally, we confirmed that either the overexpression of Brd4 or the knock-out of Tfeb in macrophage exacerbated atherosclerosis in Apoe−/− mice and abolished the anti-atherosclerotic effects of Cur. The decreased ROS generation, increased expression of LC3, and decreased H3 phosphorylation by mTORC1 were all inhibited by macrophage-specific Tfeb KO. Mechanically, ChIP analysis showed that under ox-LDL stimulation, the aberrant occupancy of the promoters of inflammatory genes by BRD4 was reversed by Cur and attenuated by Tfeb silencing. The collective findings indicate that activated TFEB is a negative regulator in the upstream of BRD4 in Cur-treated FCs. In assessing how Cur decreases BRD4 expression in FCs, we found that the Cur mediated decreases of BRD4 was reversed by inhibition of autophagic flux, rather than by a proteasome inhibitor. The finding was supported by the prediction that BRD4 harbors the iLIR autophagy database. Interaction between BRD4 and LC3 was demonstrated in Cur-treated FCs. The interaction was further increased in the presence of CQ. The collective findings confirmed that Cur can degrade BRD4 in FCs by activating autophagy.

Histone acetylation is usually achieved by histone acetyltransferases including P300. The latter is a major enzyme that mediates the function of BRD4. The role of P300 in atherosclerosis is controversial. One study reported that P300 promotes the atherogenesis by activating vascular smooth muscle cells proinflammatory genes. However, another study showed that P300 rescues the base excision repair enzyme activity to delay the plaque development by decreasing oxidative damage. The results of our present study show that P300 activity was upregulated in FCs as evidenced by the increase of acetylated histone 3. This event was reversed by Cur, which is consistent with the reported that Cur is an inhibitor of P300. Cur was reported to inhibit the binding efficiency of histones and acetyl CoA to P300 via binding with P300/MBD at a specific site. However, we proposed that Cur could inhibit P300 in another way. Here, P300 activity in FCs was downregulated in the presence of Cur, the same as the effect of the ROS broad scavenger NAC, in accordance with the report that...
ROS can increase P300 activity. This reminds us that ROS is the upstream activator of P300. In exploring whether TFEB activation caused by Cur regulated P300 through ROS, we observed that ROS generation in FCs was inhibited by Cur. This event was reversed by TFEB or ATG5 deficiency. These findings indicate that Cur can inhibit ROS generation by activating autophagy due to TFEB nuclear translocation, which in turn reduces P300 activity. We confirmed that ROS increases the P300 activity by increasing BTK phosphorylation. Although P300 is widely reported to be a negative regulator of autophagy, we confirmed that autophagy can also negatively regulate the activity of P300. This finding broadens the understanding of the interplay between P300 and autophagy.

The SE is composed of clusters of transcriptional enhancers, which are strongly enriched for the binding of MED1, BRD4, H3K27ac and other transcriptional coactivators to genes essential for certain cellular function. Data from bioinformation databases reveal an SE region at the promoter of inflammatory genes, which was further enhanced with ox-LDL treatment and inhibited by Cur. Thus, SE could be a new target for Cur. Phase separation is a physicochemical process by which a homogenous liquid solution of macromolecular components separates into a dense phase and a dilute phase. Biomolecular condensates produced by LLPS allow rapid movement of components into and within the dense phase and exhibit properties of liquid droplets, such as fusion and fission. The aberrant forms of phase separation are now thought to mediate many complex human diseases, such as neurodegenerative diseases, cancers, and infectious diseases. Recently, BRD4 has been reported to be involved in the formation of LLPS at SEs with coactivator MED1 to control gene expression. In the present study, BRD4 was upregulated in macrophages by treatment with the classical stimulator ox-LDL, which promoted the expression of inflammatory genes. Notably, LLPS in FCs was associated with BRD4. All these findings implicate FCs as a new model to develop new drugs targeting phase separation and BRD4, with the aim of controlling gene expression. Data from bioinformation databases reveal an SE region at the promoter of inflammatory genes. The collective evidence in the present study implicates that Cur can restore the autophagy and inflammatory genes. Notably, LLPS in FCs was associated with BRD4. The SE is composed of clusters of transcriptional enhancers, which are strongly enriched for the binding of MED1, BRD4, H3K27ac and other transcriptional coactivators to genes essential for certain cellular function. 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