Aspirin modulates the inflammatory response in a thrombus-stimulated LMVEC model

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Abstract. The purpose of the present study was to examine whether aspirin interferes with the inflammatory response in a thrombus-stimulated lung microvascular endothelial cell (LMVEC) model. The LMVECs were randomly divided into eight groups: Normal group (group N), model group (group M), model + ASP group (group M+A), model+CX3CL1-short hairpin (sh)RNA group (group M+SH), model + CX3CL1-overexpression vector group (group M+CX3), model + ASP + shRNA group (group M+A+SH), model + ASP + CX3CL1-overexpression vector group (group M+A+CX3), and normal + virus control group (group N+V). The endothelial cells were cultured, and a thrombus was added to the cells. Briefly, 12 h following the precipitation of the thrombus, data from ELISA, reverse transcription-quantitative polymerase chain reaction analysis and confocal microscopy revealed that the levels of tumor necrosis factor (TNF)-α, interleukin (IL)-6, CX3C chemokine ligand 1 (CX3CL1), CX3C chemokine receptor 1 (CX3CR1) and nuclear factor-κB (NF-κB) in group M were increased, compared with those in group N (P<0.01). These levels, with the exception of TNF-α, were significantly lower in group M+SH, compared with those in group M (P<0.01). Furthermore, the levels of IL-6 in groups M+A, M+CX3 and M+A+CX3 were decreased, compared with those in group M (P<0.01); the level of TNF-α in group M+A+SH was decreased, compared with that in group M (P<0.01); the level of CX3CR1 was lower in groups M+A and M+A+SH, compared with that in group M (P<0.01), and the level of NF-κB in group M+SH was decreased, compared with the level in group M (P<0.05). In conclusion, the thrombus-stimulated LMVEC model exhibited induced production of TNF-α, IL-6, CX3CL1, CX3CR1, NF-κB and intercellular adhesion molecule-1. Furthermore, it was confirmed that the signaling pathways involving CX3CL1-NF-κB, IL-6 and TNF-α were partly inhibited by aspirin.

Introduction

Acute pulmonary embolism (APE) refers to the blockage of at least one vessel in the lungs, which occurs due to thrombus lodging and leads to disrupted pulmonary circulation. The pathological mechanisms of APE include endothelial injury, activation of the coagulation cascade, inflammation, and the promotion of thrombus formation via inflammatory cell recruitment (1,2). APE-associated inflammation has been somewhat neglected in investigations over previous years, and there are limited reports on the condition. Our previous studies demonstrated that a wide range of inflammatory responses, including the expression of tumor necrosis factor-α (TNF-α), interleukin (IL)-1β, IL-8, the irregular chemokine CX3C chemokine ligand 1 (CX3CL1) and its receptor CX3C chemokine receptor 1 (CX3CR1) were found in an APE rat model. In addition, the level of CX3CL1 was shown to positively correlate with the level of TNF-α (3,4). It was also found that lipopolysaccharide (LPS) can induce the expression of nuclear factor (NF)-κB and CX3CL1 in human bronchial epithelial cells (5), and it was found that aspirin improved the pathological changes in rats with APE via the CX3CL1/CX3CR1 signaling pathway (3,4). Based on these findings, adenovirus vectors carrying CX3CL1-short hairpin (sh)RNA and
Materials and methods

Materials. Aspirin enteric-coated tablets were purchased from Nanjing Baijingyu Pharmaceutical Co., Ltd. (Nanjing, China; specifications: 25 mg x 100 tablets/bottle, batch no. 141201).

Western blot analysis. Western blot analysis was used to detect NF-κB. A PVDF membrane was obtained from EMD Millipore (Billerica, MA, USA); cat. no. IPVH00010; batch no. K5JA5013L. Anti-NF-κB p65 antibody was obtained from Abcam (Cambridge, MA, USA; cat. no. ab7970). β-actin was purchased from Multi Sciences (Lianke) Biotech, Co., Ltd. (cat. no. ab008-100). Goat anti-rabbit IgG was bought from Multi Sciences (Lianke) Biotech, Co., Ltd. (cat. no. GAR0072). Goat anti-mouse IgG was obtained from Bioworld Technology, Inc. (St. Louis Park, MN, USA; cat. no. BS12478). An electrophoresis system (Mini-Protein Tetra system) was purchased from Bio-Rad Laboratories, Inc. (Hercules, CA, USA). A gel imager was obtained from Bio-Rad Laboratories, Inc. (Chemidoc XRS+ system).

Reverse transcription-quantitative polymerase chain reaction (RT-qPCR) analysis. RT-qPCR analysis was performed to detect the expression levels of CX3CL1, CX3CR1, IL-6 and TNF-α. A highly purified total RNA was extracted in the pre-experimental data. For each group, experimentation was run in triplicate. ELISA was used to detect IL-6, TNF-α and IFN-γ. The expression levels of cytokines were analyzed using the following forward and reverse primers as previously described (8). The mRNA expression of CX3CL1, CX3CR1, IL-6 and TNF-α was determined using qPCR using IQ SYBR Green SuperMix PCR array kit was purchased from Bio-Rad.

Methods

Establishment of an LMVEC model induced by thrombus stimulation (6,7). All animal experiments, including the method of sacrifice, were performed in compliance with the institutional animal care regulations and guidelines of Zhejiang Chinese Medical University (Hangzhou, China) and performed according to the Association for Assessment and Accreditation of Laboratory Animal Care International (AAALAC) and the Institutional Animal Care and Use Committee (IACUC) guidelines. The present study was approved by the Laboratory Animal Management and Ethical Review Committee (no. ZSLL-2015-1015) of Zhejiang Chinese Medical University (Hangzhou, China).

The rat LMVECs were purchased from Pri Cells Biomedical Technology Co., Ltd. (Wuhan, China). Frozen LMVECs were recovered and cultured in LHC-8 serum-free medium ( Gibco; Thermo Fisher Scientific, Inc., Waltham, MA, USA) at 37°C, in a 5% CO2 incubator. For subculture, the cells were washed with PBS, digested with 0.125% trypsin + 0.01% EDTA, and maintained in logarithmic phase culture. For treatment, the cells were digested, counted and seeded in 6-well plates at a density of 2x10^5 per well. Male Sprague-Dawley (SD) rats (n=3; 8 weeks old, 200 g) were purchased from Shanghai SLA Laboratory Animal Co., Ltd. (Shanghai, China). The animals were housed in 12 h dark/12 h lights, temperature of 22-26°C and atmosphere of ~0.03-0.04% CO2. Animals were given access to food and water ad libitum. Following a 6-day acclimation, the experiment was initiated. The rats were anesthetized by intraperitoneal injection of 2% sodium pentobarbital (0.25 ml/100 g); all surgery was performed under sodium pentobarbital anesthesia and all efforts were made to minimize suffering. Rat venous blood was collected, and a thrombus 5 mm in size was prepared. The LMVECs were randomly divided into two groups (n=3): Normal group (group N) and model group (group M); the cells in group N did not receive thrombus stimulation. Thrombosis precipitated naturally (a 2/3 thrombus pavement was considered a base). Time points of 0, 0.5, 1, 4, 8 and 12 h were set prior to experiment commencement; the 12-h time point was selected as the appropriate time to wash away the thrombus for measurement based on the pre-experimental data. For each group, experimentation was run in triplicate. ELISA was used to detect IL-6, TNF-α and IFN-γ. RT-qPCR analysis was performed to measure the expression levels of CX3CL1, CX3CR1, IL-6 and TNF-α; the primer sequences are listed in Table I. Total RNA was extracted using TRIzol reagent (Thermo Fisher Scientific, Inc.), according to the manufacturer's protocol. A total of 2 µg of extracted RNA was converted to cDNA by MMLV-reverse transcriptase (Fermentas; Thermo Fisher Scientific, Inc.), which was used according to its manufacturer's protocol. The cDNA was amplified using the following forward and reverse primers as previously described (8). The mRNA expression of CX3CL1, CX3CR1, IL-6 and TNF-α was determined with qPCR using IQ SYBR Green SuperMix PCR Array kit was purchased from Bio-Rad.
The sequence of primers was as follows: Rat CX3CL1 forward, GCC ATG AAG AGG GAc; rat CX3CR1 reverse, GGG GTT GAC TCC. The reaction was designed and synthesized at Shanghai Generay Biotech (Shanghai, China). The reaction was performed and analysed by a CFX Connect Real-Time PCR System (Bio-Rad Laboratories, Inc.). The relative expression levels of the mRNA in each sample were calculated using the 2^ΔΔCt method (9). These levels were expressed in arbitrary units.

Western blot analysis was used to evaluate the expression of NF-κB. Protein samples were extracted from the rat LMVECs using Protein Extraction kit (Beyotime Institute of Biotechnology, Jiangsu, China). The protein concentration was determined using bicinchoninic acid method. Protein samples (30 mg) were separated using 4% to 12% Bis-Tris SDS-PAGE gels and then transferred to polyvinylidene fluoride membranes. Membranes were blocked with 5% skim milk in PBS/1% Tween-20 for 1 h at room temperature. Following incubation PVDF membrane was washed with 1X TBST solution for 5 min. The membranes were then incubated with primary antibodies as described above overnight at 4°C. Subsequently, the membranes were incubated with secondary antibodies as described above for 2 h at room temperature. ECL chemiluminescence reagent (Beyotime Institute of Biotechnology) was used for visualization.

Chemiluminescent images of the blots were finally captured using a ChemiDoc System. ImageJ software was used to calculate the integrated absorbance of identified bands (IA), and the expression of protein was calculated using the following formula: 

\[ \text{Relative expression of protein} = \frac{\text{IA}_{\text{protein}}}{\text{IA}_{\text{actin}}} \]

Laser confocal microscopy was used to evaluate the expression levels of CX3CL1/CX3CR1 and CX3CL1/NF-κB. Fractalkine (CX3CL1) antibody (Santa Cruz Biotechnology, Inc.; 1:10), anti-NF-κB p65 antibody (1:50), anti-CX3CR1 antibody (1:100) (both from Abcam), anti-rabbit IgG secondary antibody (488 conjugate; Thermo Fisher Scientific, Inc.; excitation wavelength/emission wave length: 488/520 nm; 1:200), anti-goat IgG secondary antibody (555 conjugate; Thermo Fisher Scientific, Inc., excitation wavelength/emission wave length: 555/562 nm; 1:200), and DAPI (Sigma; EMD Millipore; excitation wavelength/emission wavelength: 358/461 nm) were used.

A fluorescein isothiocyanate (FITC) labeling kit, which was purchased from Sangon Biotech Co., Ltd. (Shanghai, China), with excitation and emission wavelengths of 490 and 525 nm respectively, was used to label recombinant protein. The freeze-dried powder of the recombinant protein was dissolved in PBS and mixed with FITC. The sample was then incubated in the dark at room temperature for 5 h. Subsequently, the sample was purified in a gel column, and unbound FITC was removed. A cell climbing slice was prepared, and FITC-labeled CX3CL1/CX3CR1 and CX3CL1/NF-κB were added to the slice and incubated at room temperature for 10 min. Following incubation, the slice was washed with PBS four times. Following mounting of the slice, laser confocal microscopy was used to evaluate the cellular localization of CX3CL1/CX3CR1 and CX3CL1/NF-κB. For staining, CX3CL1 (goat anti human antibodies) was combined with either CX3CR1 (rabbit anti human antibodies) or NF-κB p65 (rabbit anti human antibodies) for double staining. The fluorescent secondary antibody applied to CX3CL1 was donkey anti-goat IgG (H+L), secondary antibody® 647 (near-infrared; pseudocolor green). The fluorescent secondary antibody applied to CX3CR1 and NF-κB p65 was goat anti-rabbit IgG (H+L), secondary antibody® 594 (red). ImageJ 4.1 software (National Institutes of Health, Bethesda, MD, USA) was used to calculate the fluorescence value (unit: pixels): Cell fluorescence intensity = fluorescence value/mean value of group N.

**Vector construction.** The construction of vectors carrying CX3CL1-overexpression shRNA was performed as an in vitro experiment. Vector pHBAD-MCMV-GFP (Hanbio, Shanghai, China), vector pHBAD-U6-GFP (Hanbio), Escherichia coli strain DH5α (Tiangen Biotech Co., Ltd.), restriction enzyme, T4 ligase (both from Fermentas Thermo Fisher Scientific, Inc.) and a plasmid DNA extraction kit (CWBio, Beijing, China) were used. The TCID50 method was applied to detect the titer (10). The viral concentration was MIO=10.

**Verification of the effect of CX3CL1 on the model.** The adenovirus vectors of CX3CL1-overexpression were constructed. The LMVECs were randomly divided into four groups (n=3): Normal group (group N), model group (group M), model + CX3CL1-overexpression vector group (group M+CX3), and normal + virus control group (group N+V). ELISA was used to determine the levels of IL-6, TNF-α and ICAM-1. RT-qPCR analysis was performed to detect the expression levels of CX3CL1, CX3CR1, IL-6 and TNF-α, whereas the cell culture supernatant was separated for ELISA.

**Verification of the effects of CX3CL1-shRNA on the model.** The adenovirus vectors carrying CX3CL1-shRNA were constructed. The LMVECs were randomly divided into four groups (n=3): Group N, group M, model+CX3CL1-shRNA group (group M+SH), and group N+V.

**Verification of the effects of aspirin combined with CX3CL1 on the model.** The LMVECs were randomly divided into eight groups (n=3): Group N, group M, model + aspirin group (group M+A), model + SH, group M+CX3, model + shRNA + aspirin group (group M+A+SH), model + aspirin + CX3CL1-overexpression vector (group M+A+CX3), and group N+V (Table II).

**Statistical analysis.** IBM SPSS 21.0 (IBM SPSS, Armonk, NY, USA) was used for statistical analysis. The experimental results are expressed as the mean ± standard deviation, and two independent samples were compared using the paired t-tests. Single factor analysis of variance was used to compare
Results

Thrombus-stimulated model

ELISA. ELISA was used to detect the levels of IL-6, TNF-α and ICAM-1. The levels of IL-6 (207.90±16.69) and TNF-α (239.60±15.27) in group M was significantly increased, compared with those in group N (85.93±20.31 and 101.23±11.91, respectively) (all P<0.01); whereas there was no difference in the level of ICAM-1 between group N (1.44±0.15) and group M (1.79±0.35) (P>0.05; Fig. 1A).

RT-qPCR detection. The levels of CX3CL1 (0.00033±0.000006) and CX3CR1 (0.27±0.03) in group N were similar to those in group M (0.00020±0.000000 and 0.35±0.04) (P>0.05).

multiple samples, and the LSD test was used for comparison among groups. P<0.05 was considered to indicate a statistically significant difference.

Table I. Primer sequences for reverse transcription-quantitative polymerase chain reaction detection of CX3CL1, CX3CR1, IL-6 and TNF-α.

| Target gene | Primer sequence (5’-3’) | Product size (bp) |
|-------------|--------------------------|-------------------|
| Genbank ID: 89808 | | |
| RatCX3CL1 Forward | GCAGTGACTGGATCGTCTCC | 117 |
| RatCX3CL1 Reverse | ACTCGGCCCATGGTGTTG | |
| Genbank ID: 171056 | | |
| RatCX3CR1 Forward | GAGTTGGCCAGAAGGGAAG | 97 |
| RatCX3CR1 Reverse | GGGTTGAGGAGCA | |
| Genbank ID: 25464 | | |
| RatICAM1 Forward | ATCATGTGCGCTCGTG | 127 |
| RatICAM1 Reverse | AGGGAGCGGGGCTGTAC | |
| Genbank ID: 24835 | | |
| RatTNF-α Forward | CCACACGCTCTTCAGTCTA | 150 |
| RatTNF-α Reverse | GGGGATGAGGGCTTGCAC | |
| Genbank ID: 24498 | | |
| RatIL6 Forward | TCCAGGCTGCTTGCTTCTT | 103 |
| RatIL6 Reverse | TGTGTGGGGTGTATTCCTG | |

CX3CL1, CX3C chemokine ligand 1; CX3CR1, CX3C chemokine receptor 1; ICAM-1, intercellular adhesion molecule-1; IL-6, interleukin-6; TNF-α, tumor necrosis factor-α.

Table II. Experimental grouping.

| Group | Treatment |
|-------|-----------|
| Normal group (group N) | No treatment, cultured for 72 h. |
| Model group (group M) | No treatment, cultured for 72 h, with re-prepared thrombus added in last 12 h. |
| Model + ASP group (group M+A), Model + shRNA group (group M+SH) | Aspirin added at a concentration of 5 mM. Pre-prepared thrombus added in last 12 h. Treated with MIO=10; knockdown virus added for 72 h, with final thrombus added in last 12 h. |
| Model + overexpression group (group M+CX3) | Treated with MIO=10⁴, overexpression virus added for 72 h, final thrombus added in last 12 h. |
| Model + ASP + shRNA group (group M+A+SH) | Treated with MIO=10, knockdown virus and 5 nM aspirin added for 72 h, with final thrombus added in the last 12 h. |
| Model + ASP + CX3CL1-overexpression vector (group M+CX3) | Treated with MIO=10, overexpression virus and 5 nM aspirin were added for 72 h, with final thrombus added in last 12 h. |
| Normal + virus control group (group N+V) | Adenovirus was added for 72 h. |

⁴MIO is a viral unit, 10 refers to one cell with 10 viruses.
There was a significant difference in the level of IL-6 between groups N (0.11±0.02) and M (0.66±0.05) (P<0.01). In addition, the level of TNF-α in group N (0.02±0.01) differed significantly from that in group M (1.06±0.04) (P<0.01; Fig. 1B).

**Western blot analysis detection of NF-κB.** No difference in the level of NF-κB was found between groups N (0.66±0.62) and M (0.56±0.25) (P>0.05; Fig. 1C and D).

**Laser confocal microscopy.** Laser confocal microscopy was used to evaluate the co-expression of CX3CL1/CX3CR1 and CX3CL1/NF-κB, and to compare the cell fluorescence intensities. The level of NF-κB in group N (1.00±0.09) was decreased compared with that in group M (1.21±0.07) (P<0.05). Compared with group N (1.00±0.31), the level of CX3CR1 in group M (3.94±0.58) was significantly increased (P<0.01; Fig. 1E).

**Verification of the effects of CX3CL1 on the model**

**ELISA detection of IL-6, TNF-α and ICAM-1.** The level of IL-6 in group N was significantly decreased compared with the levels in groups M and M+CX3 (159.27±15.02) (P<0.01; Fig. 2A). Compared with group M, the levels of IL-6 in groups N, M+CX3 and N+V (107.40±12.26) were decreased (P<0.01). The level of TNF-α was significantly higher in groups M, M+CX3 (216.97±9.20) and N+V (158.00±13.95), compared with that in group N (P<0.01; Fig. 2A). Compared with group M, the levels of TNF-α were significantly decreased in groups N and N+V (P<0.01). Compared with the group N, there was no difference in the level of ICAM-1 in the other groups (P>0.05; Fig. 2A), nor was there a statistically significant difference in the level in group M+CX3 (1.37±0.23), compared with that in group M (P>0.05).

**RT-qPCR detection of CX3CL1, CX3CR1, IL-6 and TNF-α.** Compared with groups N and M, the level of CX3CL1
in group M+CX3 (0.55±0.01) was significantly increased (P<0.01; Fig. 2B). Compared with groups N and M, the level of CX3CR1 in group N+V (2.09±0.36) was significantly increased (P<0.01). Compared with group N, the levels of IL-6 in groups M and M+CX3 (0.29±0.03) were significantly increased (P<0.01; Fig. 2B). Compared with group M, the levels of IL-6 in groups N and N+V (0.15±0.02) were significantly decreased (P<0.01). Compared with the group N, the levels of TNF-α in groups M and M+CX3 (0.88±0.07) were significantly elevated (P<0.01). Compared with group M, the level of TNF-α in groups N, M+CX3, and N+V (0.05±0.01) were also significantly decreased (P<0.01; Fig. 2B).

Western blot analysis detection of NF-κB. Compared with groups N (0.66±0.62) and M (0.56±0.25), there was no significant difference between the levels of NF-κB in groups M+CX3 (1.14±0.74) and N+V (0.29±0.11) (P>0.05; Fig. 2C and D).

Laser confocal microscopy evaluation of co-expression of CX3CL1/CX3CR1 and CX3CL1/NF-κB. CX3CL1 fluorescence intensity was significantly decreased in group M+CX3 (0.59±0.09) compared with that in group M (P<0.01; Fig. 2E). The intensity of CX3CL1 fluorescence was elevated significantly in groups M and N+V (3.47±0.16), compared with that in group N (P<0.01). Compared with group M, the intensity of CX3CL1 fluorescence was decreased significantly in groups N and M+CX3 (1.10±0.16) (P<0.01). No significant differences in the intensity of NF-κB fluorescence was observed between groups (P<0.05). Compared with group M, the
level of NF-κB fluorescence was significantly decreased in groups N, M+CX3 (0.41±0.08) and N+V (0.82±0.05) (P<0.05; Fig. 2E).

Verification of the effect of CX3CL1-shRNA on the model ELISA detection of IL-6, TNF-α and ICAM-1.

Compared with group N, the level of IL-6 was significantly elevated in groups M, M+SH (183.60±11.52) and N+V (P<0.01; Fig. 3A). Compared with group M, the level of IL-6 was significantly decreased in groups N, M+SH, and N+V (P<0.01). Compared with group N, the level of TNF-α was significantly increased in the other four groups (P<0.01; Fig. 3A). Compared with group M, no significant difference in the level of TNF-α was found between groups N, M+SH (282.00±5.63) and N+V (P>0.05). The level of ICAM-1 in group M+SH (2.39±0.13) was higher compared with that in group N (P<0.01; Fig. 3A). The level in group M+SH differed significantly compared with that in group M (P<0.05).

RT-qPCR detection of CX3CL1, CX3CR1, IL-6, and TNF-α.

The level of CX3CL1 was significantly different in groups M and N+V (P<0.05; Fig. 3B). Compared with group M, the level of CX3CL1 was significantly different in groups N, M+SH, and N+V (P<0.01). Compared with group N, there was no significant difference in the level of CX3CR1 in group M+SH (0.33±0.05) (P>0.05; Fig. 3B). Compared with group M, the level of CX3CR1 was significantly increased in group N+V (P<0.01). The level of IL-6 was significantly elevated in groups M and M+SH (0.82±0.17) compared with that in group N (P<0.01; Fig. 3B). Compared
with group M, the level of IL-6 was decreased significantly in groups N and N+V (P<0.01). The level of TNF-α was increased significantly in groups M and M+SH (0.41±0.04) compared with that in group N (P<0.01; Fig. 3B). Compared with group M, the level of TNF-α different significantly in groups N, M+SH and N+V (P<0.01).

**Western blot analysis for the detection of NF-κB to verify the effect of CX3CL1-shRNA on the model.** Compared with groups N (0.66±0.62) and M (0.56±0.25), no significant differences were found in groups M+SH (0.79±0.19) and N+V (0.29±0.11) (P>0.05; Fig. 3C and D).

**Evaluation of the co-expression of CX3CL1/CX3CR1 and CX3CL1/NF-κB by laser confocal microscopy.** Compared with groups N and M, the intensity of CX3CL1 fluorescence was significantly decreased in group M+SH (0.41±0.07) (P<0.01; Fig. 3E). Compared with group N, the intensity of CX3CR1 fluorescence was significantly elevated in groups M and N+V (P<0.01; Fig. 3E). Compared with group M, CX3CR1 fluorescence was significantly decreased in groups N and M+SH (0.85±0.18) (P<0.01; Fig. 3E). NF-κB fluorescence intensity was significantly different in groups M, M+SH (0.33±0.7) and N+V, compared with that in group N (P<0.05). Compared with group M, NF-κB fluorescence intensity was significantly decreased in the other three groups (P<0.05).

**ELISA detection of IL-6, TNF-α and ICAM-1** (Fig. 4). Compared with group N, the level of IL-6 was significantly increased in groups M (P<0.01), M+A (118.50±13.56) (P<0.05), M+SH (P<0.01), M+A+SH (193.53±28.86; P<0.01) and M+CX3 (P<0.01). Compared with group M, the level of IL-6 was decreased in groups N, M+A, M+CX3, M+A+CX3 (97.63±13.57) and N+V (P<0.01). Compared with group M+A, a significant difference in the level of IL-6 was found in groups N (P<0.05), M (P<0.01), M+SH (P<0.01), M+CX3 (P<0.05) and M+A+SH (P<0.01). Compared with group N+V, the level of IL-6 differed significantly in groups M, M+SH, M+A+SH and M+CX3 (P<0.01). Compared with group N, the level of TNF-α was significantly increased in the other seven groups (P<0.01). Compared with group M, there was significant difference in groups N (P<0.01), M+SH (P<0.01), M+A+SH (197.90±14.19; P<0.01), M+A+CX3 (205.07±25.81; P<0.05) and N+V (P<0.01). Compared with group M+A (216.67±21.60), there was a significant difference in groups N, M+SH and N+V (P<0.01). Compared with group N+V, the level of TNF-α was significantly increased in groups M, M+A, M+SH, M+CX3, M+A+SH and M+A+CX3 (P<0.01).
Compared with group N, the level of ICAM-1 was significantly elevated in groups M+SH and M+A+SH (3.00±0.15) (P<0.01). Compared with group M, there was a significant difference in groups M+SH (P<0.01), M+A+SH (P<0.01), M+cX3 (P<0.05) and M+A+cX3 (1.32±0.32; P<0.05). Compared with group M+A (1.74±0.18), there was a significant difference in the level of ICAM-1 in groups M+SH (P<0.01), M+A+SH (P<0.01) and M+cX3 (P<0.05). Compared with group N+V, the level of ICAM-1 was increased significantly in groups M+SH and M+A+SH (P<0.01).

RT-qPCR detection of CX3CL1, CX3CR1, IL-6 and TNF-α (Fig. 5). The level of CX3CL1 was found to differ significantly in the other seven groups, compared with that in group M+CX3 (P<0.01). Compared with group M+CX3 (1.30±0.08), the level was decreased significantly in the other seven groups (P<0.01). The level of CX3CR1 was decreased significantly in the other seven groups, compared with that in group N+V (P<0.01). Compared with group N, the level of IL-6 was significantly different in groups M, M+A (0.27±0.02), M+SH, M+CX3 and M+A+SH (0.86±0.05) (P<0.01). Compared with group M, the level was increased significantly in groups M+SH (P<0.05) and M+A+SH (P<0.01), but was decreased in groups N (P<0.01), M+SH (P<0.01), M+CX3 (P<0.01), M+A+CX3 (P<0.01) and N+V (P<0.01). Compared with group M+A, the level differed significantly in groups N, M, M+SH, M+A+SH, M+CX3 (0.09±0.01) and N+V (P<0.05). Compared with group N+V, the level differed significantly in groups M (P<0.01), M+CX3 (P<0.05), M+SH (P<0.01), M+A+SH (P<0.01) and M+CX3 (P<0.05). For TNF-α, compared with group N, the level of was increased significantly in all groups, with the exception of group N+V (P<0.01). Compared with group M, there was a significant difference in levels between groups N (P<0.01), M+SH (P<0.01), M+CX3 (P<0.05) and M+A+SH (0.58±0.10; P<0.01), N+V (P<0.01). Compared with group M+A (1.01±0.04), there was a significant difference in levels between groups N, M+SH, M+A+SH and N+V (P<0.01). Compared with group N+V, the level of TNF-α was increased significantly in all groups, with the exception of group N (P<0.01).

Western blot analysis for the detection of NF-κB (Figs. 6 and 7). Compared with group M+CX3 (1.14±0.74), the level of NF-κB was decreased significantly in group M+CX3 (0.47±0.30) (P<0.05). Compared with group N+V
Compared with group N, the intensity of CX3CR1 fluorescence was significantly elevated in groups M, M+A (2.64±0.21), M+A+SH (1.99±0.47), M+A+CX3 (2.42±0.32) and N+V (P<0.01). Compared with group M, fluorescence intensity was decreased significantly in groups M+SH, M+CX3 and M+A+CX3 (P<0.01). Compared with group M+A, fluorescence was significantly decreased in groups N (P<0.01), M+SH (P<0.01), M+CX3 (P<0.01) and M+A+SH (P<0.05). Compared with group N+V, a significant difference was found between all groups, with the exception of group M (P<0.01).

Compared with group N, NF-κB fluorescence intensity was significantly different in groups M (P<0.05), M+A (1.21±0.07; P<0.01), M+SH (P<0.01), M+CX3 (P<0.01), M+A+SH (0.80±0.09; P<0.05), M+A+CX3 (0.68±0.06) (P<0.01) and N+V (P<0.05). Compared with group M, fluorescence intensity was significantly different in group N (P<0.05), M+SH (P<0.01), M+CX3 (P<0.01), M+A+SH (P<0.01), M+A+CX3 (P<0.01) and N+V (P<0.01). Compared with group M+A, fluorescence intensity was significantly different in all groups but group M (P<0.01). Compared with group N+V, there was a significant difference in groups N (P<0.05), M (P<0.01), M+A (P<0.01), M+SH (P<0.01) and M+CX3 (P<0.01).

**Discussion**

In order to elucidate the potential roles of aspirin in the inflammatory responses involving the LMVEC model, the present study investigated gene-protein and receptor-nuclear interactions. A total of four experimental stages were performed: Validation of the successful construction of a thrombus-stimulated cell model; validation of the effect of the overexpression of CX3CL1 in the model; validation of the effect of CX3CL1-inhibition in the model; and examination of the effects of aspirin combined with CX3CL1 on the model. The data from the first three experiments were derived from the fourth experiment, and the experiments were performed in triplicate.

When thrombosis occurs, damage in endothelial cells leads to an inflammatory response (2). In the initial pre-experimental stage of the present study, a thrombus-stimulated LMVEC model and a cell scratch test were used, which revealed that the inflammatory effects were more pronounced, in the former and a previous report argued that the model could be successfully manufactured (6). Accordingly, the model was improved in the present study using the instrument-specialized flow of blood to an advantage, with the experiment involving dynamic change. Subsequently, the changes in NF-κB B, IL-6, TNF-α, ICAM-1, CX3CL1 and CX3CR1, generated by the stimulatory effect of the thrombus in LMVECs, were investigated, and the effects of the overexpression and inhibition of CX3CL1, or the effects of aspirin application were examined. ELISA, RT-qPCR analysis and cell fluorescence revealed that the model group secreted significantly higher levels of TNF-α, IL-6, ICAM-1, CX3CL1, CX3CR1 and NF-κB, compared with the control group, suggesting that the inflammatory response model of the thrombus-stimulated LMVECs had been constructed successfully.

Subsequently, the present study investigated the changes generated by the stimulatory effects of the thrombus on (0.29±0.11), the level was significantly elevated in group M+CX3 (1.14±0.74) (P<0.05). There was no significant difference between groups N (0.66±0.62), M (0.56±0.25) and N+V (0.29±0.11) (P<0.05).

**Laser confocal microscopic evaluation of the co-expression of CX3CL1/CX3CR1 and NF-κB (Figs. 8-10).** Compared with group N, the intensity of CX3CL1 fluorescence was decreased significantly in groups M+SH and M+CX3 (P<0.01). Compared with group M, fluorescence was significantly decreased in groups M+SH, M+CX3 and M+A+CX3 (0.80±0.17) (P<0.01). Compared with group M+A (1.49±0.19), fluorescence was significantly decreased in groups N (P<0.01), M+SH (P<0.01), M+CX3 (P<0.01), M+A+SH (0.98±0.26) (P<0.05), M+A+CX3 and N+V (P<0.05). Compared with group N+V, there was a significant difference in fluorescence between groups M+A (P<0.05), M+SH (P<0.01) and M+CX3 (P<0.01).
LMVECs via the overexpression of CX3CL1. It was found that, compared with the control group, the overexpression of CX3CL1 in the intervention model significantly stimulated the secretion of IL-6, TNF-α and CX3CR1, but had minimal effect on the expression of NF-κB, ICAM-1 and CX3CR1.

In the third step, it was noted that the levels of IL-6, TNF-α and ICAM-1 were significantly elevated in the CX3CL1-shRNA intervention model, compared with those in the control group. However, the expression levels of IL-6, ICAM-1, CX3CL1, CX3CR1 and NF-κB expression were significantly decreased in the model group.

Finally, the present study found that aspirin inhibited the expression of IL-6 in the model group; aspirin combined with shRNA or the overexpression of CX3CL1 inhibited the secretion of IL-6 in the model group, whereas aspirin combined with shRNA inhibited TNF-α in the model group. The inhibition of CX3CL1 by shRNA in the model suggested that the construction of a CX3CL1-shRNA adenovirus vector had been successful. Aspirin, either alone or combined with shRNA, inhibited CX3CR1 in the model, with shRNA being more effective than aspirin at inhibiting CX3CR1. shRNA inhibited NF-κB, and was more effective than aspirin at inhibiting NF-κB in the model.

Fractalkine (CX3CL1) is the only member of the CX3C family that can bind to its specific receptor (CX3CR1) and mediate tight adhesion between inflammatory cells and vascular endothelial cells (11). CX3CL1 is important in the recruitment of inflammatory cells into the vascular wall in endothelial cell injury (12, 13). Previous studies have reported that TNF-α stimulates the CX3CL1/CX3CR1 inflammatory...
signaling pathway, whereas our previous studies revealed that the expression levels of TNF-α, CX3CL1 and CX3CR1 were significantly increased in rats with pulmonary embolism (3,14). Certain studies have reported that the presence and the stimulation of the CX3CL1 receptor, CX3CR1, is correlated with the reduced release of IL-1β and TNF-α (15,16). Sukkar et al (17) reported that chemokines are important in bronchial asthma and chronic obstructive pulmonary disease, with CX3CL1 acting as a chemo attractant for cell adhesion molecules, monocytes and T cells. The combination of interferon-γ and TNF-α stimulated the mRNA and protein secretion of CX3CL1 dependent on time and concentration, whereas dexamethasone significantly promoted the secretion of CX3CL1. Studies have also reported that CX3CL1 is involved in a variety of diseases, including acute severe pancreatitis, cerebral ischemia and acute hypoxia, breast cancer, rheumatoid arthritis, and increased pulmonary hypertension (18-22). The regulatory mechanisms associated with CX3CL1 may also involve NF-κB. Yang et al (23) suggested that TNF-α promotes the adhesion of monocyte THP-1 to human umbilical vein endothelial cells via the regulation of adhesion molecules by the NF-κB signaling pathway, including the expression of chemokine CX3CL1/monocyte chemotactic protein-1 of ICAM/vascular cell adhesion molecule/S-select in. Cimato et al (24) found that inflammatory cytokines, rather than cholesterol, regulated the expression of CX3CL1 in atherosclerosis. Cao et al suggested that human dermal microvascular endothelial cell-1 in patients with inflammatory skin disease induced chemokines, including CXCL8, CX3CL1 and CXCL16, by activating the NF-κB signaling pathway when stimulated by TNF-α, which is in agreement with our previous finding that LPS-NF-κB-CX3CL1 exists in human bronchial epithelial cells (5,25). The present study revealed that CX3CL1-shRNA inhibited the secretion of NF-κB in the thrombus-stimulated LMVEC model, which was also consistent with a previous report (26). In addition, aspirin combined with CX3CL1-shRNA inhibited the secretion of NF-κB, further supporting our previous finding that aspirin decreased the levels of NF-κB in rats with pulmonary embolism (4). Our previous study found that aspirin significantly decreased pulmonary arterial pressure, improved pathological changes in embolism, and decreased the expression of CX3CL1/CX3CR1 and CX3CL1/NF-κB, whereas the adenovirus CX3CL1-overexpression vector aggravated the inflammatory changes in APE improved
by aspirin. In addition, the intervention of the adenovirus CX3CL1 vector reduced the changes, and its combination with aspirin significantly improved the changes (4). These in vivo experimental results were further validated in the cell experiments in the present study.

ICAM-1 belongs to an immunoglobulin superfamily, and is a transmembrane, single chain glycoprotein expressed in leukocytes, fibroblasts and epithelial cells (27). ICAM-1 can induce the expression of inflammatory factors, including IL-1α, IL-1β, IL-6 and TNF-α, via signaling pathways involving NF-κB, thereby promoting the development of inflammation (28). In the present study, it was found that a significantly higher level of ICAM-1 was expressed in the model group, compared with that in the control group, and that CX3CL1-shRNA decreased the expression of ICAM-1. Aspirin did not effectively inhibit the secretion of ICAM-1 in the model group, which was consistent with a previous finding that CX3CL1 induced cell migration by upregulating the expression of ICAM-1 via the CX3CR1/Pi3K/Akt/NF-κB pathway in human osteosarcoma cells (29).

Aspirin irreversibly inhibits cyclooxygenase, preventing the formation of thromboxane A2 in platelets (30). The substance can also have anti-inflammatory effects by preventing the activation of NF-κB (31), associated with the inhibited activation of NF-κB kinase β by aspirin, decreasing the binding activity of NF-κB DNA (32). Studies have reported that aspirin exerts certain effects on NF-κB, IL-6, TNF-α, ICAM-1, CX3CL1 and CX3CR1, which is consistent with the findings of the present study (33,34).

As the primary mechanism underlying pulmonary embolism formation and propagation is not caused by the inflammatory response itself, future investigations are required to focus on the association between inflammation and blood coagulation following endothelial cell damage in the pulmonary vasculature.

A major limitation of the present study was that the Virchow’s triad mechanism of venous thrombosis involves vascular endothelial cell injury, circulatory stasis and a hypercoagulable state, therefore, the association between inflammation and coagulation requires further investigation.

In conclusion, the present study showed that TNF-α, IL-6, CX3CL, CX3CR1, NF-κB and ICAM-1 were secreted in the thrombus-stimulated LMVEC model; the signaling pathways of CX3CL1-NF-κB, IL-6 and TNF-α were important in the disease process and were partly inhibited by aspirin.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions

LW proposed the study and wrote the paper. RY analyzed and interpreted the data from Thrombus-stimulated model, HJ analyzed and interpreted the data from verification of the effects of CX3CL1 on the model, QJ analyzed and interpreted the data from verification of the effect of CX3CL1-shRNA on the model, JK was involved in the ELISA and western blotting, ZZ was involved in the confocal microscopy, YS in was involved in reverse transcription-quantitative polymerase chain reaction, DC in was involved in the confocal microscopy and RW was involved in vector construction. All authors read and approved the final manuscript.

Ethics approval and consent to participate

All animal experiments, including the method of sacrifice, were performed in compliance with the institutional animal care regulations and guidelines of Zhejiang Chinese Medical University, and performed according to the AAALAC and the IACUC guidelines. The present study was approved by the Laboratory Animal Management and Ethical Review Committee (approval no. ZSLL-2015-1015) of Zhejiang Chinese Medical University (Hangzhou, China).

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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