A novel role for Livin in the response to ultraviolet B radiation and pterygium development

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Abstract. A pterygium is an inflammatory, invasive and proliferative lesion on the ocular surface, which can decrease visual acuity, damage the ocular surface and affect the appearance of the eye. However, the underlying molecular mechanisms of the pathogenesis remain unclear. In the present study, the role of apoptosis-associated protein Livin in the occurrence and development of pterygium was investigated. Primary samples from quiescent or advanced clinical stages of pterygium and normal human conjunctival tissues were used to assess mRNA and protein expression levels of Livin using reverse transcription-quantitative PCR and immunohistochemistry, respectively. Livin was knocked down in pterygium epithelial cells (PECs) using small interfering RNA (siRNA), to investigate the role of Livin in PEC viability, migration, invasion ability and apoptosis. The cell viability, invasion ability and apoptosis of PECs following ultraviolet B (UVB) radiation alone or in combination with Livin silencing were also analyzed. Expression levels of Livin increased in the pterygium tissues compared with those in the normal conjunctiva at both the mRNA and protein levels. Livin expression levels in advanced pterygium were significantly higher compared with those in quiescent pterygium samples. Knockdown of Livin expression levels significantly reduced cell migration, invasion ability and cell viability, and induced apoptosis of PECs. Inhibition of Livin expression in PECs increased the expression levels of caspase-7, caspase-3 and E-cadherin, whereas expression levels of Snail were downregulated. Cell viability and invasion ability in PECs was enhanced following UVB radiation and Livin expression upregulated. UVB irradiation induced cell invasion ability of PECs and this was attenuated by Livin-silencing. Transfection with Livin siRNA also partially recovered the apoptosis rate of PECs, which was reduced by UVB irradiation. In conclusion, Livin was upregulated in pterygium, and UVB radiation functions in the development of pterygium by inducing Livin expression.

Introduction

Pterygium is a common disease of the ocular surface, characterized by invasion of triangular inflammatory fibrovascular lesions into the cornea, which can cause inflammation, irregular corneal astigmatism and stromal opacity (1). Currently, the underlying mechanisms of pterygium have not been well elucidated and surgical excision is the standard treatment; however, pterygium recurrence is common following surgery (2). In order to devise strategies for the non-surgical treatment of pterygium or the prevention of recurrence after resection, it is important to understand the underlying molecular mechanisms of the pathogenesis of pterygium.

Pterygium is more common in tropical and subtropical zones, and epidemiological evidence indicates that the pathogenesis of pterygium is strongly associated with high exposure to ultraviolet (UV) radiation (3). UV radiation induces expression of several proinflammatory cytokines, including transforming growth factor-β, interleukin (IL)-1, IL-6, and IL-8, and growth factors, including fibroblast growth factor and vascular endothelial growth factor, which promote the progression of pterygium (3-6); however, the exact mechanism by which UV leads to the onset of pterygium remains unclear.

Pterygium exhibits tumor-like features, including propensity to invade normal tissue, recurrence after excision and coexistence with premalignant lesions (4,7,8). Previous studies have demonstrated that pterygium epithelial cells (PECs) are highly proliferative, have features of epithelial-mesenchymal transition (EMT) and overexpress anti-apoptotic proteins (9-13). Livin is a member of the inhibitors of apoptosis protein (IAP) family (14), which function in tumor initiation, progression and resistance to chemotherapy (15). Livin has been reported to serve a prominent and specific role in certain malignancies and is associated with diverse cellular behaviors, including proliferation, invasiveness and motility (14). However, the role of Livin in ocular surface diseases has not been previously reported, and to the best of our knowledge, there is no research investigating the expression and molecular mechanisms of Livin in the development of pterygium.
In the present study, the expression levels and distribution of Livin in pterygium was investigated and the association between UVB radiation and Livin expression levels was explored. Knockdown of Livin expression was used to investigate the function of Livin in the cellular behavior of pterygium epithelium and the potential underlying molecular mechanisms.

Materials and methods

Patients and samples. The present study was in compliance with The Declaration of Helsinki and was approved by the Institutional Review Board of Hangzhou Red-Cross Hospital (Hangzhou, China). All patients provided written informed consent. A total of 19 male and 16 female patients with primary pterygium (age range, 52-79 years; mean age, 60.3±7.7 years), treated at Hangzhou Red-Cross Hospital between November 2018 and March 2019, were included in the present study. Exclusion criteria included a history of any other ocular surface disorders, previous ocular surgery, ocular trauma, systemic autoimmune disease or inflammation and use of any topical steroids or non-steroidal anti-inflammatory drugs. Normal conjunctival tissue segments included a history of any other ocular surface disorders, previous ocular surgery, ocular trauma, systemic autoimmune disease or inflammation and use of any topical steroids or non-steroidal anti-inflammatory drugs. Normal conjunctival tissue and pterygium were cut and ground, then fixed in 10% buffered formaldehyde solution and embedded in paraffin before subsequent experiments (<3 months).

Excised tissues were divided laterally into two symmetrical pieces of identical size. One piece (for measurement of Livin mRNA) was frozen and stored at -80˚C and the other piece was divided into two pieces of identical size. One piece (for measurement of Livin mRNA) was frozen and stored at -80˚C and the other piece was fixed in 10% buffered formaldehyde solution and embedded in paraffin before subsequent experiments (<3 months).

Reverse transcription-quantitative (RT-q)PCR. Normal conjunctival tissue and pterygium were cut and ground, then 1 ml TRIzol® (EZBioscience) was used to extract total cellular RNA according to the manufacturer's protocol. Subsequently, RNA purity and concentration were measured using a Nanodrop 2000 (Thermo Fisher Scientific, Inc.). Then RNA was reverse transcribed to cDNA using a Revert Aid First Strand cDNA Synthesis kit (Thermo Fisher Scientific, Inc.). qPCR was performed using SsoAdvance Universal SYBR-Green Supermix (Bio-Rad Laboratories, Inc.). PCR was performed at 95˚C for 3 min, then for 40 cycles at 95˚C for 10 sec, 60˚C for 30 sec and 72˚C for 30 sec. The primer sequences of the target genes were as follows: Livin forward, 5'-ACACACGAGCAGAGAGGAGGAGG-3' and reverse 5'-GGCAGTCAGCAGGCCAGT CAT-3'; and β-actin forward, 5'-GAACCCACTAAAGGCTACACAG AGAAAGG-3' and reverse 5'-CCACTGACATAAGGAGGAGGAAC-3'; β-actin was used as an internal control. The experimental results were automatically calculated using a CFX96™ Real-Time PCR Detection system (Bio-Rad Laboratories, Inc.). The relative expression of Livin gene was expressed as a fold-increase of β-actin calculated using the 2-ΔΔCq method (17).

Immunohistochemistry analysis. The tissues were fixed in a 10% formalin solution and embedded in paraffin. Samples were cut into 4-µm sections, mounted on glass and dried overnight at 37˚C. All sections were then deparaffinized in xylene, rehydrated with alcohol (100% ethanol for 2x3 min; 95% ethanol for 3 min; 70% ethanol for 3 min; 50% ethanol for 3 min) and washed in PBS. Sections were then inactivated using 3% H2O2 for 10 min. Sections were heated to 100˚C in EDTA (pH 9.0) for 10 min and then naturally cooled and soaked in PBS three times for 3 min each. The sections were blocked in a 37˚C oven for 45 min using 5% BSA (Sigma-Aldrich; Merck KGaA). The immunohistochemistry experiment was conducted using the streptavidin-biotin-peroxidase method (18). Anti-Livin primary antibody (cat. no. ab93750; 1:50; Abcam) was applied to the sections and incubated overnight at 4˚C. 3,3'-Diaminobenzidine was used as a chromogen, followed by rinsing with water for 5 min and counterstaining with hematoxylin for 2 min at room temperature.

The numbers of epithelial cells and Livin-positive cells of the pterygium or normal conjunctiva were counted in three independent fields using a high-power field under a light microscope (magnification, x40). Cells positively stained for the anti-Livin antibody were scored according to the percentage extent of staining and the measurements were averaged. Scores were as follows: 0, no positive staining; +, 1-10%; ++, 11-50%; and +++, >50% positive cells. Scores of +, ++ and +++ were considered to represent positive immunostaining, whereas a score of 0 indicated negative immunostaining. All evaluations of the staining were conducted by an observer blinded to the origins of the tissue samples.

Western blot analysis. Total protein was extracted from patient tissues using RIPA lysis buffer (Beyotime Institute of Biotechnology) and 20 µg protein/lane was loaded onto a 12% gel, resolved using SDS-PAGE and subsequently transferred to PVDF membranes (EMD Millipore). PVDF membranes were immersed in a blocking solution containing 5% skimmed milk powder in PBS or 5% BSA and blocked at room temperature for 2 h. Subsequently, the membranes were incubated with primary antibodies at 4˚C overnight, followed by secondary antibodies at room temperature for 2 h. The proteins were visualized using a Clarity™ Western ECL Substrate (Bio-Rad Laboratories, Inc.) and detected using a chemiluminescence detection system (Chemidoc MP; Bio-Rad Laboratories, Inc.) and the bands were analyzed using Image software (v1.46; National Institutes of Health). Anti-GAPDH antibodies were used as an internal control (cat. no. cst5174; 1:1,000; Cell Signaling Technology, Inc.). The primary antibodies included anti-Livin (cat. no. NB100-56548; 1:500-1:2,000; Novus Biologicals, Ltd.), anti-caspase-7 (cat. no. ab32522, 1:1,000; Abcam), anti-caspase-3 (cat. no. cst9662; 1:1,000; Cell Signaling Technology, Inc.), anti-E-cadherin (cat. no. 610405; 1:1,000; BD Biosciences), anti-Snail (cat. no. cst3879; 1:1,000; Cell Signaling Technology, Inc.) and anti-poly AD-ribose polymerase (cat. no. ab74290; PARP; 1:4,000; Abcam). The secondary antibodies included goat anti-rabbitIgG (cat. no. BL003A; 1:5,000; Biosharp Life Sciences) and goat anti-mouse IgG (cat. no. BL001A; 1:5,000; Biosharp Life Sciences).
Culture of PECs. PECs were isolated and cultured using previously established methods (10,16). Fresh primary pterygium samples at the advanced stage were washed with PBS with 1% penicillin (100 units/ml) and streptomycin (100 µg/ml) (Gibco; Thermo Fisher Scientific, Inc.). After the fibrovascular tissue was removed as much as possible, the remaining epithelial tissues were cut into small pieces of 1x1 mm and then cultured in DMEM supplemented with 10% FBS (Gibco; Thermo Fisher Scientific, Inc.), 100 U/ml penicillin and 100 mg/ml streptomycin at 37°C with 5% CO₂ in a humidified incubator. The solution was changed once every 2-3 days, and after the cells covered the bottom of the bottle, the cells were digested with 0.25% trypsin (Gibco; Thermo Fisher Scientific, Inc.) at 37°C for 2 min. Under a phase-contrast microscope, once cytoplasmic retraction and enhanced refractive index had been observed, 2 ml medium with 10% FBS was added to terminate digestion. The cell suspension was prepared, counted and inoculated. Finally, PECs were seeded in serum-free DMEM at 100 µl/well with 5% CO₂, in a humidified incubator. The cells from primary culture using tissue were defined as primary passage cells (P0) and the cells from the first passage after the primary culture were defined as first passage PECs (P1). The morphology of cells was relatively uniform, showing a fibrous/fusiform shape. Immunofluorescence results confirmed that almost all cells (P2) were positively stained with anti-cytokeratin 13 (cat. no. ab32522; 1:200; Abcam) at 4°C overnight, so the cell purity was considered to be very high, close to 100% (Fig. 1).

UVB irradiation of PECs. The cultured PECs were incubated in a 96-well culture plate with 4x10⁴ cells/well for 24 h. Culture medium was removed and replaced with PBS. The PECs were irradiated with UVB at a dose of 0, 0.5, 1 and 2.0 J/cm² using an UV irradiation chamber BS-02 (Opsyte Dr. Gröbel GmbH). PECs were collected for analysis of Livin expression, and analysis of cell viability and invasion ability following UVB irradiation.

CCK-8 assay for cell viability. Following UVB irradiation, PECs were incubated in 96-well plates at 100 µl/well with 5 replicate wells for each group for 24 h. Subsequently, 10 µl CCK8 was added to each well for analysis of cell viability according to the manufacturer’s protocol (Shanghai Yeasen Biotech Co., Ltd.). Blank control wells were set and incubated at 37°C in the dark. The optical density (OD) value was read at a wavelength of 450 nm using a microplate reader Plus848 ( Molecular Devices LLC) and the cell survival rate was calculated using the following formula: Cell viability (%%)=(experimental group OD_sixe0;/normal group OD_sixe0;) x100. The experiment was repeated three times.

PEC transfection. Small interfering RNA (siRNA) was used to knockdown endogenous Livin gene expression in PECs. Cells were transfected with Livin-specific siRNA (GE Healthcare Dharamcon, Inc.) and negative control scrambled siRNA using Lipofectamine® 2000 (Thermo Fisher Scientific, Inc.) according to the manufacturer's protocol. The PECs were seeded at a density of 1x10⁶ cells/well in 6-well plates and transfected with 30 nM Livin-siRNA using Lipofectamine 2000. After 48 h of incubation at 37°C, the cells were harvested and subjected to cell viability, migration and invasion assays, and flow cytometry analysis.

Cell apoptosis assay. Apoptosis was determined using an Annexin V-FITC/7-AAD assay. After transfection with Livin-siRNA, PECs were incubated at 37°C for 48 h. A total of 5x10⁴ cells were collected by centrifugation at 500 x g for 5 min at room temperature, washed twice with PBS and centrifuged again at 500 x g for 10 min at 37°C. A total of 100 µl binding buffer was used to re-suspend the cells. After adding 5 µl Annexin V-FITC and 5 µl 7-AAD (both Beijing Jiamay Biotech), the cells were incubated for 15 min at room temperature in the dark. After adding 400 µl binding buffer, cell apoptosis was detected using a NovoCyte flow cytometer (ACEA Biosciences, Inc.). The cells that were FITC-positive and 7-AAD-negative were indicated as apoptotic cells.

Cell migration assay. A total of 1x10⁵ cultured PECs/well were seeded in a 6-well culture plate. After incubation for 24 h at 37°C, a straight line was scratched across the cells using a P200 pipette tip to create a wound. The plates were then rinsed with PBS to remove the cells in suspension. Identical sites were observed at x40 magnification at 0 and 24 h post-scratching using a light microscope. The migration of cells toward the wounds was expressed as percentage of wound closure: % of wound closure=[(At=0 h−At=h)/At=0 h] x100%. At=0 h is the area of wound measured immediately after scratching; At=h is the area of wound measured 24 h after scratching.

Cell invasion assay. Cell invasion was measured in a Transwell cell culture chamber. Cultured PECs were seeded in the upper chambers with 25 µg Matrigel (8-12 mg/ml). A total of 100 µl cell suspension using serum-free medium was added to the upper chamber (9x10⁶ cells/chamber) and 600 µl DMEM containing 10% serum as a chemokine was added to bottom chamber. Incubation was performed at 37°C with 5% CO₂ for 48 h. The chamber was then removed, washed three times with PBS, fixed with immobilization solution for 20 min and washed three times with PBS. After staining with crystal violet for 20 min at room temperature, the cells were counted
Livin expression levels are upregulated in pterygium compared with those in normal conjunctiva.

The Livin expression levels were analyzed in normal human conjunctiva and pterygium tissues using RT-qPCR. Livin mRNA expression levels were increased 5-fold in quiescent stage pterygium and 5.3-fold in the pterygium at the advanced stage compared with that in the normal conjunctiva (Fig. 2A). Advanced stage pterygium tissues exhibited significantly higher Livin mRNA expression levels compared with quiescent stage tissues (Fig. 2A). Immunohistochemistry demonstrated that Livin protein was expressed in pterygium tissues primarily located in the nuclei of epithelial cells, whereas it was seldom expressed in the normal conjunctiva (Fig. 2B).

In the pterygium group, 28/35 samples (80.0%) stained positively for Livin, whereas in the normal conjunctiva group only 1/10 samples (10.0%) demonstrated Livin-positive staining. The difference between the pterygium and normal conjunctiva groups was significant (80.0 vs. 10%; P<0.001). Staining scores were higher in advanced-stage compared with quiescent-stage samples (Table I).

**Results**

*Livin expression levels are upregulated in pterygium compared with those in normal conjunctiva.* The Livin expression levels were analyzed in normal human conjunctiva and pterygium tissues using RT-qPCR. Livin mRNA expression levels were increased 5-fold in quiescent stage pterygium and 5.3-fold in the pterygium at the advanced stage compared with that in the normal conjunctiva (Fig. 2A). Advanced stage pterygium tissues exhibited significantly higher Livin mRNA expression levels compared with quiescent stage tissues (Fig. 2A). Immunohistochemistry demonstrated that Livin protein was expressed in pterygium tissues primarily located in the nuclei of epithelial cells, whereas it was seldom expressed in the normal conjunctiva (Fig. 2B).

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**Knockdown of Livin expression impairs cell migration, invasion and viability, and induces apoptosis in PECs.** To
further evaluate the role of Livin in pterygium development and progression, \textit{ex vivo} cultured PECs were transfected with Livin-siRNA or control siRNA. Western blot analysis showed that the expression levels of Livin protein and mRNA were significantly downregulated in Livin-siRNA-transfected group compared with those in the control group, suggesting successful knockdown of Livin in Livin-siRNA-transfected PECs (Fig. 3A-C). Knockdown of Livin expression significantly reduced invasion, migration and viability of PECs (Fig. 3D-H) and the proportion of apoptotic cells was increased in Livin-siRNA in PECs (Fig. 3I and J).

Livin regulates EMT and apoptosis via E-cadherin, Snail, caspases and PARP. Previous studies have demonstrated that EMT is an initial step required for cell metastasis and that it has a role in apoptotic resistance (19). Therefore, the role of Livin in regulating the expression of EMT-associated proteins was also investigated. The data indicate that knockdown of Livin expression significantly increased E-cadherin and reduced Snail expression levels (Fig. 4A and B), indicating Livin is involved in the regulation of EMT. The protein expression levels of apoptosis-associated enzymes, caspase-3 and 7, were significantly increased in the Livin-knockdown PECs. PARP protein expression levels were also increased in Livin-knockdown PECs, but this difference was not significant (Fig. 4B).

\textbf{Effect of UVB irradiation on Livin expression, cell viability and invasion ability in PECs.} UVB is a major risk factor for pterygium development (3). To further investigate the association between Livin expression levels and cell viability and invasion following UVB irradiation, \textit{ex vivo} PECs in culture were treated with UVB radiation. Compared with the non-radiation treatment group, Livin protein expression levels were significantly increased in UVB irradiation-treated PECs (Fig. 5A) and PEC viability was significantly increased with increasing doses of UVB radiation (Fig. 5B). In addition, cell invasion was promoted by UVB irradiation (Fig. 5C).

UVB irradiation may promote the development and progression of pterygium by inducing the expression of Livin. Western blot analysis was used to evaluate Livin protein expression following UVB irradiation with or without Livin-knockdown. Livin-knockdown followed by UVB irradiation resulted in partial inhibition of UVB-induced Livin protein expression (Fig. 6A and B). In the invasion experiments, UVB induced an increase in cell invasion ability, which was attenuated by Livin silencing (Fig. 6C and D). Cell apoptosis analysis revealed that UVB
irradiation reduced apoptosis of PECs, and transfection with Livin-siRNA prior to UVB irradiation increased the apoptosis of PECs compared with the control siRNA group (Fig. 6E and F).

Discussion

Pterygium is a fibrovascular neoformation composed of epithelium and highly vascular loose subepithelial connective tissue (20). Previous studies have demonstrated that apoptotic and oncogenic proteins, microsatellite instability, inflammatory mediators, extracellular matrix modulators, EMT and other factors are involved in the development of pterygium (4,9‑11,13). However, to the best of our knowledge, the precise molecular mechanisms of pterygium pathogenesis have not been resolved. It is hypothesized that epithelial cells function in the pathogenesis of pterygium, with EMT and secretion of matrix metalloproteinases from epithelial cells associated with the invasive and recurrent behavior of pterygium (21‑23).

Previous studies have demonstrated that a number of apoptosis-associated genes and proteins are differentially expressed in pterygium compared with normal conjunctiva tissues, including p53, Survivin and Bcl-2, indicating the role of apoptosis in the pathogenesis of pterygium (12,16,24). Pterygium may be associated with disrupted control of cellular apoptosis rather than an increase in proliferative capacity, therefore, the regulation of PEC apoptosis may have a role in the pathogenesis of pterygium (25). As a member of the IAP family, Livin participates in apoptosis, proliferation and the cell cycle by binding to caspases and regulation of EMT (14,26). Livin is not expressed in the majority of normal adult tissues; however, it is present in developmental tissues and highly expressed in most tumor tissues (27). In the present study, the molecular mechanism underlying the function of Livin in the pathogenesis of pterygium was investigated. Livin mRNA and protein expression levels were significantly upregulated in pterygium tissues compared with those in the normal conjunctiva. Moreover, Livin expression levels were higher in advanced pterygium compared with those in quiescent pterygium tissues. This difference may be due to the higher cell proliferation pattern in advanced pterygium compared with that in quiescent pterygium. Therefore, upregulated Livin may be involved in the occurrence and development of pterygium, which requires further research. In the present study, cultured PECs were
Livin expression in advanced pterygium may have a role in promoting epithelial proliferation, resulting in progression of the pterygium. To explore whether Livin expression is involved in the occurrence and development of pterygium, PECs were transfected with Livin-siRNA in the present study. Knockdown of Livin expression levels increased apoptosis of PECs and decreased their proliferation, migration and invasion abilities, which is consistent with the results of a previous study investigating other IAPs (16). It is hypothesized that EMT occurs during the pathogenesis of pterygium (13). Snail is a key transcription factor involved in EMT regulation (28). E-cadherin is downregulated in during EMT, associated with the detachment and invasion of cells from their primary site, and results in metastasis and progression (13,28,29). In the present study, Livin-knockdown reduced Snail expression and increased E-cadherin expression, consistent with the reduced...
proliferation, migration and invasion of PECs following Livin-knockdown, indicating that Livin may be associated with EMT. Therefore, Livin may be involved in the pathogenesis of pterygium via a function in EMT.

IAPs suppress apoptosis induced by a range of stimuli, primarily via direct binding of the BIR domain to specific intracellular proteases, including caspase-3, -7 and -9, inhibiting their activity (24). PARP is an important downstream substrate of caspase-3 (30). In the present study, Livin-knockdown increased the expression of caspase-3, -7 and PARP, consistent with results of a previous study investigating the function of Livin in oral squamous cells (27,31). Therefore, the anti-apoptotic function of Livin may be mediated by suppressing the activity of caspases in PECs.

A number of studies have demonstrated that UV exposure is a factor precipitating the development of pterygium (3,21,32,33). Histological examination has confirmed that elastic connective tissue lesions and severely damaged areas of the Bowman membrane indicate direct damage caused by solar radiation or indirect damage due to excessive proteolytic activity (3,21). The present study showed that Livin expression levels were increased by UVB irradiation and contributed to the increased cell viability and invasion ability in PECs. Meanwhile, knockdown of Livin expression attenuated the effects of UVB-induced Livin expression level on cell viability and cell invasion in PECs. This decreased cell viability and invasion ability in Livin-knockdown PECs following UVB exposure suggests that Livin may have a role in inducing pterygium formation by increasing cell viability and invasion. PEC apoptosis was reduced by low-dose UVB irradiation, and knockdown of Livin prior to UVB irradiation increased the apoptosis of PECs. Therefore, the increase in cell viability and invasion induced by UVB irradiation may be due to upregulated Livin expression, and silencing of Livin expression may protect against pterygium development.

Overall, the present study suggests that increased Livin expression levels may be involved in the pathogenesis and development of pterygium. Knockdown of Livin reduced the proliferation, migration and invasion of PECs, and induced apoptosis by regulating EMT and apoptosis-associated proteins. UVB radiation increased the expression of Livin and cell invasion; therefore, UVB is a risk factor for pterygium. The present study provides novel insight into the pathogenesis of pterygium and may contribute to the development of novel therapeutic agents for treatment. However, the interaction between Livin and other cytokines and growth factors in pterygium development still requires further investigation.

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Availability of data and materials

The datasets during the current study are available from the corresponding author on reasonable request.

Authors' contributions

SQW, QBX, WYS and LWZ designed the study. SQW, QBX and LWZ drafted the original manuscript, collected the data and reviewed the literature. QBX, WYS and LYS interpreted the data and critically reviewed the manuscript. All authors have read and approved the final manuscript.

Ethics approval and consent to participate

The present study was approved by The Institutional Review Board at Hangzhou Red-Cross Hospital (Hangzhou, China) and written informed consent was provided by all patients.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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