Collagen is the building component of temporomandibular joint (TMJ) discs and is often affected by inflammation in temporomandibular disorders. The macromechanical properties of collagen are deteriorated by chronic inflammation. However, the mechanism by which inflammation influences disc function remains unknown. The relationship between the ultrastructure and nanomechanical properties of collagen in inflamed discs should be clarified. Seven-week-old female Sprague–Dawley rats were randomly divided into two groups. Chronic TMJ inflammation was induced by intra-articular injection of complete Freund’s adjuvant, and samples were harvested after 5 weeks. Picrosirius staining revealed multiple colours under polarized light, which represented alternative collagen bundles in inflamed discs. Using atomic force microscopy scanning, the magnitude of Young’s modulus was reduced significantly accompanied with disordered collagen fibril arrangement with porous architecture of inflamed discs. Transmission electron microscopy scanning revealed a non-uniform distribution of collagen fibres, and oversized collagen fibrils were observed in inflamed discs. Fourier transform infrared microspectroscopy revealed a decrease in 1 338 cm⁻¹/amide II area ratio of collagen in different regions. The peak positions of amide I and amide II bands were altered in inflamed discs, indicating collagen unfolding. Our results suggest that sustained inflammation deteriorates collagen structures, resulting in the deterioration of the ultrastructure and nanomechanical properties of rat TMJ discs.

**INTRODUCTION**

Temporomandibular disorders (TMDs) are a large family of diseases affecting the structure and function of the stomatognathic system. The temporomandibular joint (TMJ) disc absorbs peak stress during mandibular movement; thus, the disc effectively protects the TMJ. Abnormal temporomandibular disc function plays a significant role in pathogenesis of TMD. Disc displacement is one of the most common subtypes of TMD; however, its pathologic mechanism is unclear.

A normal disc has a concave configuration, which is consistent with the surface of condyle and temporal bone. Most studies have reported that disc deformation contributes to disc displacement and is often accompanied by inflammation and pain. However, different opinions exist. In several cases, TMJs exhibit inflammation and disc degeneration without disc displacement. Early studies on cadavers revealed that the deformation of the TMJ disc is associated with synovial inflammation without displacement; previous animal studies have reported that induced chronic inflammation of TMJ results in disc degeneration, including thickening and deformation. In addition, chronic inflammation of the TMJ could lead to the deterioration of macroscopic mechanical properties of discs, including a decrease in the tensile and compression modulus. Therefore, our group proposes an innovative theory: “inflammation may be the key factor leading to TMJ disc displacement”.

Stress distribution in the TMJ disc during mandible movement exhibits significant differences in normal and disc displacement groups; a disc with displacement endures more stress. Several findings have revealed the close relationship between disc structure and function. Studies have indicated that the arrangement of collagen fibrils influence tensile ability, and glycosaminoglycan influences compressive ability. However, how disc structural alteration results in biomechanical changes remains unclear. Moreover, little information is available regarding the relationship between inflammation and collagen deterioration in TMJ discs.

Complete Freund’s adjuvant (CFA) has been widely used for inducing acute or chronic TMJ inflammation by intra-articular injection. In our previous study, we found that the thickness, matrix content, biomechanical properties and surface structure of rat TMJ discs presented remarkable changes after CFA-induced chronic TMJ inflammation, and these changes result in the exposure and degradation of sub-superficial collagen fibrils. These results suggested that chronic inflammation in the TMJ could lead to the deterioration of mechanical properties and alteration of the disc surface ultrastructure, which might contribute to TMJ disc displacement. These previous studies prompted us to evaluate the pathologic alteration of internal collagen fibres to clarify the process of collagen degradation under sustained-inflammation conditions.

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1. Department of Orthodontics, Peking University School and Hospital of Stomatology, 22# Zhongguancun South Avenue, Haidian District, Beijing, China; 2. Center for Craniofacial Stem Cell Research and Regeneration, Peking University School and Hospital of Stomatology, 224 Zhongguancun South Avenue, Haidian District, Beijing, China; 3. Center for Temporomandibular Disorders and Orofacial Pain, Peking University School and Hospital of Stomatology, 22# Zhongguancun South Avenue, Haidian District, Beijing, China.

Correspondence: Yan-Heng Zhou (yanhengzhou@vip.163.com) or Xue-Dong Wang (xuedongwang2012@gmail.com)

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In the present study, we hypothesize that chronic inflammation contributes to the deterioration of the nanomechanical properties and ultrastructure of collagen fibres in TMJ discs. Considering that synovitis and internal derangement exhibit increased prevalence in females compared with males, we chose female rats to induce chronic TMJ inflammation similar to our previous studies. We applied picrosirius red staining, atomic force microscopy (AFM), transmission electron microscopy (TEM), Fourier-transformed infrared microspectroscopy (FTIRM) and circular dichroism to evaluate the changes in the arrangement, nanomechanical properties, ultrastructure and chemical structure of collagen fibrils in inflamed TMJ discs. An explanation of the mechanism of inflammation-induced degeneration of TMJ discs from the view of collagen fibres was provided.

**RESULTS**

Arrangement of different collagen fibres in picrosirius red staining. Picrosirius red staining is proportional to the content and orientation of collagen fibre in the extracellular matrix of TMJ discs and significantly facilitates the detection of collagen bundle arrangements. The inflamed and control discs exhibited considerable differences during polarizing microscopy. Haematoxylin and eosin (HE) staining for routine microscopy revealed the distinct locations of TMJ discs. Picrosirius red staining revealed the detailed morphology of the collagen fibres of discs at high magnification (Fig. 1). Although the orientation of collagen fibres was heterogeneous in the control group, the fibres were well organized. Compared with the control group, inflamed collagen fibres were disordered. The collagen fibres from the control group appeared greenish-yellow to yellow in most positions of the discs in the anterior band, intermediate zone and posterior band. The inflamed discs exhibited apparent red and yellow refraction when observed under polarized light. In the anterior band, the collagen fibres in the entire area exhibited a predominant red colour, and the green colour significantly diminished. In the intermediate zone and posterior band, the collagen fibres exhibited multiple colours, including yellow, orange and red. The greenish-yellow to red colour under polarized light indicates the changes of collagen arrangement and type as well as the transition from fine fibres to thick collagen fibre bundles. The statistical analysis of integrated optical density showed the significant differences of the two groups in the anterior and posterior bands. In addition, it showed the same tendency in the intermediate zone without significant differences of greenish colour.

Micro-morphology is disordered in inflamed discs by AFM imaging. Representative mapping images from various locations of the control and inflamed discs are presented in Fig. 2a–l. The arrangement of the collagen micro-fibrils in the control discs was regular and directional. The bundles consisted of distinctive collagen micro-fibrils, and the clearance between adjacent bundles was narrow in the anterior band and intermediate zone. In the posterior band, the clearance of collagen bundles was wider compared with the other two locations. Although the control samples exhibited various morphologies, tropocollagen molecules staggered from each other by approximately 67 nm, which is called D-period, exhibited a good performance in all of the mapping images from different positions. Both the arrangement and structure of collagen micro-fibrils in the inflamed samples were disordered and exhibited large interspace between adjacent bundles. The magnitude of irregular collagen bundles increased after induced inflammation.

The internal morphological changes in the collagen fibrils were analysed using AFM. The ultrastructure of collagen micro-fibrils is presented in Fig. 2m–p. In the control group, collagen micro-fibrils presented a clear D-period, and the repeated cycle was approximately 67 nm, which represented typical collagen ultrastructure.

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**Fig. 1** Histopathologic changes of TMJ discs. a–c Representative microscopic features of the discs from the control group subject to HE staining. The disc was thinner in the intermediate zone and thicker in the anterior and posterior bands. d–f Representative images of the disc from the control group assessed by picrosirius staining under polarized light. Collagen bundles presented yellowish to green reflection, and distribution of different colours was organized. The black dotted squares in a–c are magnified in d–f, respectively. g–l Representative microscopic features of the discs from the inflamed group subject to HE staining. The disc was thickened in all three regions. j–l Representative image of the disc from the inflamed group assessed by picrosirius staining under polarized light. Collagen bundles exhibited varied colours, but yellow to pink dominated. The distribution of different colours was disordered. The black dotted squares in g–l are magnified in j–l, respectively. m–o Inflammation decreased green intensity and increased red intensity in anterior and posterior bands significantly and exhibited the same tendency noted in the intermediate zone without significant differences. Y-axis represents IOD (integrated optimal density of positive pixel area). A, anterior band; In, intermediate zone; P posterior band. *P < 0.05, **P < 0.01.
However, in the inflammation group, the inflamed collagen micro-fibrils exhibited an attenuated D-period and disordered ultrastructure.

Collagen fibrils became hypertrophic in inflamed discs as assessed by TEM scanning. The collagen fibres in the control group were configured regularly with similar diameters and spaces. However, in the inflamed discs, the distribution of the diameters of collagen fibres manifested a broad range, and hypertrophy of collagen fibrils was observed in all the three bands. The oversized collagen fibrils, especially fibrils with a diameter of >120 nm, dramatically increased in most areas of the entire disc (Fig. 3). The distribution of the diameters of collagen fibrils revealed significant differences between the control and inflammation groups (P < 0.001) in all the three bands.

Elastic modulus of collagen decreased in inflamed discs as assessed by the AFM test. The Derjaguin–Muller–Toporov model was employed to calculate elastic modulus during AFM scanning. Representative colour-coded mapping images revealed a visible difference between the control and inflamed groups. The elastic modulus of a different location exhibited a small range in the control group (anterior band, 1.0–2.0 GPa; intermediate zone: 0.8–1.9 GPa; posterior band: 0.8–1.9 GPa, median: 1.3 GPa). Compared with the control group, the modulus of the elastic modulus decreased significantly in the anterior band (0.9–1.3 GPa, median: 1.1 GPa), intermediate zone (0.6–1.7 GPa, median: 1.2 GPa) and posterior band (0.6–1.6 GPa, median: 1.2 GPa) in the group with inflamed discs. The reduction in the modulus represents the capacity to resist extrinsic stress, which results in the deterioration of the nanomechanical property of collagen fibrils (Fig. 4).

Integrity damage detected through FTIRM and circular dichroism under inflammation conditions. FTIRM represented the spectra of the chemical group of collagen in the control and inflamed discs. Collagen absorbance at 1338 cm\(^{-1}\) is sensitive to a decrease in intensity when collagen degrades. The colour-coded mapping images present the different distributions and magnitudes of the 1338 cm\(^{-1}\) peak area. The peak area decreased significantly in the inflamed TMJ discs (Fig. 5a–i). The comparison of spectral parameters and the area ratio between the control and inflamed specimens revealed the differences between normal and inflamed TMJ discs. After the induction of chronic inflammation, the 1338 cm\(^{-1}\)/amide II area ratio decreased significantly in the anterior, intermediate and posterior bands. The representative spectra from all the FTIRM results are presented in Fig. 5j–l. Alterations in amide I in the posterior band (from 1638 to 1646 cm\(^{-1}\)) and amide II in the anterior band (from 1545 to 1533 cm\(^{-1}\)) were apparent and revealed deterioration of the primary collagen structure. Circular dichroism revealed alterations in the secondary structure of collagen as demonstrated by negative peaks at 208 and 222 nm in the control group and characterized by an α-helix. However, the peaks diminished in the inflamed sample (Fig. 5m), reflecting uncoiling of the α-helix.

**DISCUSSION**

The aim of the present study was to elucidate possible mechanisms of sustained inflammation resulting in degenerative changes in TMJ discs. As one of the main components of TMJ discs, collagen contributes to disc structure and function. In the present study, we discovered structural disorganization in inflamed collagen fibres through histological staining. Then, we detected the ultrastructure of collagen fibrils by TEM and AFM scanning. The results revealed that collagen fibril dimensions in inflamed discs were disordered and characterized by increasing hypertrophic and small fibrils. In addition, ultrastructure disorganization of collagen micro-fibrils and degradation of collagen single fibril were also typical in inflamed discs. Simultaneously, the elastic modulus of collagen fibrils decreased significantly in inflamed discs. To elaborate the mechanism involved in the changes in the ultrastructure and nanomechanical properties of
Fig. 3 TEM analysis of internal section of discs of TMJs with or without inflammation. a–c Representative cross-sections of discs from the control group. Collagen fibril diameter ranged from 20 to 120 nm with 40–70 nm noted most often. d–f Representative cross-sections of the discs from the inflamed group. Some large collagen fibrils (d > 120 nm) were observed, and the distribution was heterogeneous. Black arrows indicate abnormal hypertrophic collagen fibrils. g Nonparametric tests of collagen fibril diameter distributions based on three rats per group. The box presents 5%, lower quartile, median, higher quartile and 95%. The black dots represent fibrils count <5% or >95%. Large collagen fibrils appeared after inflammation was induced. The control and inflammation groups exhibited significant differences in collagen diameter. ***P < 0.001, scale bar: 200 nm

Inflamed discs, we analysed primary and secondary collagen structures using FTIRM and circular dichroism. We observed alterations in the primary and secondary structures based on absorbance at 1 338 cm⁻¹ and amides I and II, and collagen presented uncoiling during inflammation. We suggest that inflammation-induced collagen molecule structure alternations contribute to deterioration of collagen ultrastructure, further influence nanomechanical properties and finally result in TMJ disc dysfunction.

The results of structural testing demonstrated how inflammation deteriorates the collagen structure of TMJ discs. Picrosirius staining demonstrated the derangement and change in collagen fibre levels between the control and inflamed groups. AFM scanning revealed that the deterioration of collagen nanofibrils may result in histological changes in collagen bundles. In consideration of the partial disappearance of the staggered banding of collagen fibrils, AFM scanning images imply the disruption of the intermolecular connections in collagen nanofibrils. Given that the collagen fibrils were composed of tropocollagens connected to one another by covalent and hydrogen bonds,23 sustained inflammation may deteriorate the original tropocollagen structure. Once the collagen structure was imperfect, the D-period and nanomechanical indentation modulus exhibited significant reductions.24 The three-dimensional (3D) reconstruction results clearly revealed the disordered arrangement of inflamed collagen fibrils in 3D space. Collagen fibrils in adult rats varied widely in diameter (range: 10–120 nm; peak: 40–70 nm). TEM revealed that the inflamed discs exhibited significant non-uniformity in collagen micro-fibril dimensions that manifested as abnormal thick fibrils (d > 120 nm). The diameters and the arrangement of control collagen fibrils exhibited predominant middle-diameter fibrils with small-diameter fibrils wedged between the middle-diameter fibrils. This pattern matches the function of enduring mechanical stress.25 However, in inflamed discs, the enlargement of collagen fibrils, especially hypertrophy, disturbed the regular alignment of collagen fibrils. This feature may contribute to reductions in the nanomechanical property of collagen in the inflamed discs. This enlargement of collagen fibrils was similar to the age-related changes in rat TMJ discs.26 However, compared with the physiological ageing process, inflammatory degenerative alterations were attributed to the disordered repair of inflamed collagen.

AFM has been applied increasingly in recent years to detect the nanomechanical and ultrastructural changes in cartilage and fibrous tissues with inflammation.27–30 To the best of our knowledge, this study is the first to test the ultrastructural and nanomechanical properties of TMJ discs using AFM. The elastic moduli of the inflamed discs decreased, which is consistent with our previous macroscopic result demonstrating that the instantaneous Young's moduli of the inflamed discs decreased significantly compared with the control group.31 Given that the microarchitecture of collagen fibrils was destroyed simultaneously by sustained inflammation, we believe that the internal changes resulted in the deterioration of the mechanical properties of TMJ discs. Nevertheless, as AFM mapping could not separate single fibrils from the sample, further studies are needed to determine whether the elastic modulus of single collagen was influenced by inflammation.

FTIRM and circular dichroism analysis revealed the structural deterioration of inflamed collagen proteins. Several indexes can be used to evaluate collagen degradation by FTIRM; these indexes have been widely utilized to observe early cartilage degeneration in osteoarthritis samples.31,32 The absorbance at 1 338 cm⁻¹, which represents collagen C=O side chain vibrations, decreased when collagen is degraded.21 The changes in 1 338 cm⁻¹ normalized to the amide II area ratio in different regions represents collagen damage.33 The decreased absorbance at 1 338 cm⁻¹ indicates a change in collagen fibril structure and reveals the loosening bond between collagen molecules. This loosening may affect the biomechanical properties of collagen. Primary molecular vibrations related to these wave-number absorbances include amide I (C=O) amide II out of phase in-plane N-H deformation and C-N stretching.34 The shift of the amide I and amide II peak location was related to intramolecular and
intermolecular hydrogen deterioration, and the secondary \( \alpha \)-helix structure was also decreased. Sustained inflammation upregulated the expression of matrix metalloproteinases (MMPs), which interact with collagen peptide. These changes result in collagen unfolding, destroyed the hydrogen bond of the active site and formed the looped-out collagen chain. These protein structural changes partly explain why the collagen fibrils exhibited hypertrophy and a disordered arrangement. The collagen fibril structure is responsible for the elastic viscosity of TMJ discs. Although the structure of collagen fibrils in different regions is anisotropic, all three regions exhibited the same changing tendency of the elastic modulus in the present study. Deterioration of biomechanical properties possibly resulted from collagen degradation. Furthermore, collagen degradation formed a porous structure between collagen bundles that caused the elastic modulus to decrease in the entire region of inflamed discs.

Inflammation could also change the micro-environment of extracellular matrix, which directly influences collagen synthesis and assembly. Our previous study revealed that under the condition of chronic inflammation, the total content of collagen was dramatically increased, which indicates the “swelling” of inflamed disc ascribed to abnormal collagen synthesis. After procollagen is secreted out of the disc cell, an aberrant micro-environment may result in disturbed collagen fibril structure. Additionally, proteinases, such as a disintegrin and metalloproteinase with thrombospondin motif 2 (ADAMTS2) and MMPs, are involved in the process of collagen fibril maturing and assembly. Thus, both the synthesis and degradation of collagen fibrils were influenced during the sustained inflammation, which could explain the possible mechanism involved in the deterioration of collagen structure and function.

However, further studies are required to elucidate the intensive connection between structure and function and determine the possible signal pathway involved during the process of collagen metabolism in the inflammatory micro-environment. Intensive studies are needed to elucidate whether the changes in the structure and mechanical properties of collagen fibrils could be prevented by anti-inflammation treatment and whether disc displacement in clinical TMD patients can be induced by chronic inflammation in an animal model.

In conclusion, sustained inflammation influences collagen structures and results in the deterioration of the ultrastructure and nanomechanical properties of TMJ discs, which may change the stress distribution during mandible movement and could represent a key factor in disc displacement.

**METHODS**

Induction of inflammation

Seven-week-old female Sprague-Dawley rats (180–200 g) were randomly divided into inflammation and control groups. TMJ inflammation was induced by two intra-articular injections of CFA (Sigma) for 5 weeks as previously described. The inflammation group was injected with 25 \( \mu \)L CFA (emulsified in 25 \( \mu \)L saline) in the upper compartment of the TMJs on days 0 and 14. The control group was injected with equivalent saline.

All rats were sacrificed by pentobarbital overdose on day 35. The animal procedures were approved by the Peking University Animal Ethics Committee prior to the initiation of the study (Approval number: LA2014221). The methods employed were performed in accordance with approved guidelines.

Tissue preparation

For histological staining, entire TMJs were removed from three rats in each group, fixed in 4% paraformaldehyde in 0.1 molL\(^{-1}\) phosphate-buffered saline, and demineralized in 15% ethylene-diaminetetraacetic acid. The specimens were dehydrated in graded alcohol and xylene, embedded in paraffin, and sectioned to 4 \( \mu \)m thickness \((n = 3)\).

For AFM and FTIRM, we harvested TMJ discs from three rats per group. Each disc was cut into anterior, intermediated, and posterior segments; embedded by Tissue-Tek (Sakura Finetek, Torrance, CA, USA); and stored at \(-80^\circ\)C for cryopreservation. The specimens were sectioned to 40 \( \mu \)m thickness parallel to the disc surface using a cryostatic microtome at \(-40^\circ\)C. All the sections were stored at \(-40^\circ\)C before testing.

HE and picrosirius red staining

The tissue sections were stained with HE to locate the different segments of discs. The sections adjacent to the HE staining

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**Fig. 4** AFM mechanical scanning images of TMJ discs. **a-c** Representative nanomechanical mapping features of discs from the control group. Magnitude of Young’s modulus was consistent with the collagen D-period banding. **d-f** Representative nanomechanical mapping features of discs from the inflamed group. Compared with the control group, the inflamed samples exhibited a decreased magnitude and disordered distribution of Young’s modulus. **g** Nonparametric tests of collagen fibril diameter distribution based on three rats per group. The box presents 5%, lower quartile, median, higher quartile and 95%. The inflammation group exhibited significant differences in Young’s modulus compared with the control group. **\( ** \) \( P < 0.01 \), scale bar: 400 nm**

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**Fig. 5** FTIRM mapping images of TMJ discs. **a–f** Mapping images of the integration area of 1338 cm\(^{-1}\). The magnitude of the inflammation group was decreased compared with the control group, and the distribution was diverse in inflamed discs. **g–j** The 1338 cm\(^{-1}\)/amide II area ratio exhibited a significant decrease in the inflammation group in the anterior band, intermediate zone and posterior band, respectively. **j–l** Representative single spectra of the TMJ discs (from the anterior band). In the anterior band, intermediate zone and posterior band, the amide I peak position moved to a large wave-number, and the amide II peak position moved to a small wave-number. **m** Circular dichroism of the TMJ disc collagen solution. The negative absorbances of 208 and 222 nm were distinctive in the control group, but less pronounced in the inflammation group.
samples were subjected to picrosirius red staining by incubation in sirius red (0.1% in saturated picric acid; #43 665, Fluka) for 1 h at room temperature. Subsequently, the sections were examined through polarization microscopy. The green and red colours of the polarized images were analysed using Image-Pro Plus 6.0 according to the positive pixel per area ($n = 3$). Integrated optimal density of green and red colour was calculated in every image. To avoid subjective factors causing errors, the investigator was blind to the samples and groups.

AFM imaging
Imaging was performed using an Icon atomic force microscope as previously described.43 The results of AFM imaging and nanoin- dentation obtained from three rats were analysed with NanoScope Analysis 1.40. The samples were imaged in Peak-Force QNM mode. A TAP525A silicon cantilever was utilized, and deflection sensitivity was calibrated using a sapphire model. Calibration of the spring constant and tip radius was conducted using a standard model with a known elastic modulus of 2.7 GPa as a reference; the value of the spring constant was 2.5–3 N m$^{-1}$, and the value of the tip radius was 12–15 nm.

To calculate the elastic modulus, three scans of representative locations were performed for each specimen. Each scan generated a mapping image with 256 × 256 resolution for the elastic modulus. Ten 100 nm × 100 nm samples were randomly selected from the collagen fibril region in each mapping image.

TEM analysis
Three TMJ discs per group were fixed with 2.5% glutaralde- hyde solution (Sigma) at 4 °C for 12 h, and TEM was performed as described previously.44 The discs were double-fixed with 1% osmium tetroxide (Sigma) and stained with lead citrate and uranyl acetate. Then, the anterior band, posterior band and intermediate zone of the discs were embedded in epoxy resin. Transverse ultrathin sections (100 nm) were obtained vertical to the disc surface. Counting and measurement of the diameter of collagen from the TEM images were accomplished by marking the fibrils individually in Image-Pro Plus 6.0 software. Collagen fibrils with diameters >120 nm were regarded as irregular.

FTIRM and circular dichroism testing for chemical changes in collagen
The contiguous sections of the AFM specimens were assessed using FTIRM. The chemical changes in areas of the anterior band, intermediate zone and posterior band were examined. To obtain the requisite spatial resolution, measurements were performed on the infrared beam line in the laboratory of Bruker Corporation (Beijing, China). Interferograms were simulta-

Statistical analysis
Statistical analysis was performed using the SPSS 11.0 software. The nonparametric Kolmogorov–Smirnov test was implemented to compare the differences in collagen diameter and elastic modulus between the control and inflammation groups. Statistical comparisons of the picrosirius red staining and FTIRM results were performed using independent $t$ tests. $P < 0.05$ was considered significant.

Data availability
The authors declare that the data supporting the findings of this study are available within the paper.

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AUTHOR CONTRIBUTIONS
S.-J.C. and X.-D.W. performed animal and laboratory experiments. S.-J.C., Y.F., Y.L., X.-X. K., J.-N.Z., and Y.H.G. contributed to data analysis and interpretation. S.-J.C. wrote a draft of the manuscript. Y.-H.Z. and X.-D.W. contributed to the concept and design of the study, the coordination of all experiments and critical review of the manuscript.

ADDITIONAL INFORMATION
Competing interests: The authors declare no competing interests.

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