Clinical characteristics of 10 Chinese patients with melorheostosis and identification of a somatic MAP2K1 variant in one case

Xiaojun Han | Yang Xu | Zhanying Wei | Chun Wang | Hua Yue | Zhenlin Zhang

1Department of Orthopedics, Qingpu Branch of Zhongshan Hospital Affiliated to Fudan University, Shanghai, China
2Shanghai Clinical Research Center of Bone Disease, Department of Osteoporosis and Bone Disease, Shanghai Jiao Tong University Affiliated Sixth People's Hospital, Shanghai, China

Correspondence
Hua Yue and Zhenlin Zhang, Shanghai Clinical Research Center of Bone Disease, Department of Osteoporosis and Bone Disease, Shanghai Jiao Tong University Affiliated Sixth People’s Hospital, Shanghai 200233, China. Email: yueyinglonghua@163.com (H. Y.) and zhangzl@sjtu.edu.cn (Z. Z.)

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Abstract
Background: Melorheostosis (MEL) is an exceptionally rare sclerosing bone dysplasia with asymmetrically exuberant bone formation and soft tissue lesions in a segmental distribution. We aimed to summarize the clinical characteristics of Chinese MEL patients and identify their pathogenic cause.

Methods: In total, 10 Chinese MEL patients were recruited, and clinical manifestations and radiological characteristics were recorded. Sanger sequencing of the LEMD3 gene was performed on peripheral blood samples of all patients, while the exome sequencing of matched peripheral blood, melorheostotic bone, and skin lesion samples was conducted on one patient who provided affected bone and skin tissues. Micro-computed tomography (micro-CT) was also used to scan the melorheostotic bone tissue.

Results: We found the average age of the 10 MEL patients was 29.5 years (range 11–40 years), and the major symptoms were bone pain, restricted movement, and bone deformity. The lesions sites were mainly located in femur (8/10), tibia (8/10), fibula (6/10), and foot (7/10), the next was pelvis (4/10), and the last were patella (1/10), hand (1/10) and spine (1/10). Radiological examinations showed a mixture of hyperostosis consisting of classic “dripping candle wax,” “osteoma-like,” or “myositis ossificans-like” patterns in most patients. No germline pathogenic variants in the LEMD3 gene were found in all patients, but a disease-causing somatic variant of MAP2K1 (c.167A>C, p.Gln56Pro) was detected in melorheostotic bone from one patient. Moreover, the micro-CT analysis showed increased porosity in the melorheostotic bone with somatic MAP2K1 variant.
Melorheostosis (MEL, OMIM 155950) is a sclerosing bone disease characterized by hyperostotic lesions in a “dripping candle wax” pattern on the cortex of the long tubular bones (Freyschmidt, 2001). It is a sporadic disease occurring in males and females equally, and the low prevalence of 1/1,000,000 makes MEL rare (Kotwal & Clarke, 2017; Wynne-Davies & Gormley, 1985). MEL lesions are asymmetrically distributed, often with multiple contiguous bones involved. Although the involvement of one limb is typical, more widespread involvement is not uncommon (Kotwal & Clarke, 2017; Smith et al., 2017; Wordsworth & Chan, 2019). Although the overgrowth lesions of MEL do not metastasize, the malignant change to osteosarcoma has been reported (Bostman et al., 1987; Murphy et al., 2003). Abnormalities of the adjacent soft tissues may result in sensory deficits, linear epidermal nevus (LEN), scleroderma-like skin (SLS), subcutaneous fibrosis, ectopic bone formation, and fibrolipmas accompany (Hellemans et al., 2004). Despite the absence of definitive diagnostic criteria, patients with MEL can usually be recognized from a combination of clinical findings, radiographs, and bone scans (Kotwal & Clarke, 2017).

The majority of MEL cases are isolated but a minority occur against a background of other hyperostotic bone diseases, osteopoikilosis (OPK, OMIM 166700) and Buschke-Ollendorff syndrome (BOS, OMIM 166700), two benign disorders caused by germline variants in the LEMD3 gene (OMIM 607844) (Hellemans et al., 2004). LEMD3 encodes the inner nuclear membrane protein MAN1, an antagonist in the TGF-β/BMP signaling pathway (Lin et al., 2005). The germline variant of LEMD3 accompanied by heterozygous somatic KRAS (OMIM 190070) variant was identified as the cause of the SLS lesion in a patient with MEL and familial OPK (Whyte et al., 2017). In patients with isolated MEL, no pathogenic germine LEMD3 variants were identified, but somatic activating variants in MAP2K1 (OMIM 176872), encoding MEK1 kinase, were found by comparing the exome sequences of affected and unaffected tissues (Mortier et al., 2019; Mumm et al., 2007). In addition to disease-causing variants in the negative regulatory domain of MEK1 kinase (p.Gln56Pro, p.Lys57Glu, and p.Lys57Asn), which were proved to increase enzymatic activity and phosphorylation of extracellular signal-regulated kinase (ERK), the variant in the catalytic domain (p.Cys121Ser) was also identified as associated with MEL (De Ridder et al., 2020; Kang et al., 2018). Furthermore, somatic activating variants in SMAD3 (OMIM 603109) were also discovered to cause MEL by up-regulating the TGF-β/SMAD pathway (Kang et al., 2020). So far, the etiology and molecular pathogenesis of MEL have been preliminarily elucidated. However, no Chinese MEL patients with definite genetic diagnoses were reported and no clear summary of the clinical characteristics of Chinese MEL patients was made.

In this study, we investigated 10 Chinese MEL patients diagnosed with clinical manifestations and typical radiological findings. Through Sanger sequencing on the LEMD3 gene of all the patient’s peripheral blood samples and exome sequencing on the biopsied melorheostotic bone and skin lesion tissues of one case, we identified a pathogenic somatic variant in MAP2K1 for the first time in a Chinese MEL patient.

### Materials and Methods

#### 2.1 Ethical compliance

The present study was reviewed and approved by the Ethics Committee of the Shanghai Jiao Tong University Affiliated Sixth People’s Hospital, and written informed consent were obtained from all the patients and the healthy volunteer.

#### 2.2 Subjects

Ten sporadic MEL patients of Han ethnicity were recruited in our outpatient from August 2009 to June 2018. The patients were diagnosed with MEL according to their clinical manifestations, plain radiography, and some even radionuclide bone scan. The principal characteristics of these patients were listed in Table 1.
Among the patients, patient 10 first came to our department in February 2013. She was born to a healthy non-consanguineous couple. From 30 years old, restricted motion range of the left hip, knee, and ankle made the patient limp, and at 37 years old, she came to our department for severe pain in the above joints. Physical examination revealed mild scoliosis, bilateral lower extremities asymmetry, and hard lumps around the left popliteal fossa and ankle. A large cutaneous lesion with prominent brown macules and erythema extended from her posterior waist to left thigh and leg following Blaschko lines.

2.3 | Blood and tissue sampling

Peripheral blood (2 ml) was obtained from all 10 patients for genomic DNA extraction. Patient 10, underwent open surgery to remove hyperplastic and sclerosing bone tissues, which restricted her motion, in our hospital. The excised bone and lesional skin tissue were immediately placed in liquid nitrogen for DNA extraction or 4% paraformaldehyde for micro-CT. The control bone sample with no radiographic evidence of MEL was obtained from the proximal tibia of a middle-aged female patient who underwent amputation surgery after a car accident. Unfortunately, the remaining 9 patients with MEL were not operated on at our hospital, so we were unable to obtain their affected bone or skin tissues.

2.4 | Exome sequencing and Sanger sequencing

Genomic DNA was extracted from 10 blood, two bone samples, and one skin lesion tissue using QuickGene DNA whole-blood Kit (Kurabo Industries Ltd., Osaka, Japan) and Qiagen DNeasy tissue Kit (Qiagen, Venlo, The Netherlands).

First, Sanger sequencing on the *LEMD3* gene was performed on the 10 patients. All exons and the exon-intron boundaries of the *LEMD3* gene (GenBank accession No. NM_014319.4) were amplified via polymerase chain reaction (PCR) using the primers designed by Primer 3 software (http://frodo.wi.mit.edu/primer3/). Direct nucleotide sequencing of the amplicons was performed using the BigDye Terminator Cycle Sequencing Ready Reaction Kit (version 3.1; Thermo Fisher Scientific, Inc., Waltham, USA) and analyzed by an ABI 3130 automated sequencer (Thermo Fisher Scientific, Inc., Waltham, USA).

Exome sequencing was applied to genomic DNA isolated from matched blood, bone, and skin samples of Patient 10. All samples were fragmented into 100–800bp pieces with a peak size of ~250bp via NEBNext dsDNA Fragmentase (New England Biolabs, Ipswich, UK). And then, end-repairing, dA-Tailing, and adaptor ligation were conducted using the NEBNext DNA Library Prep Reagent Set from Illumina (New England Biolabs, Ipswich, UK). 2% agarose gel electrophoresis was used to fraction adaptor-ligated DNA fragments and fragments of the desired size (300–400bp) were excised. 10 PCR cycles were adopted to amplify the extracted DNA with PE primers (Illumina, San Diego, USA) and Phusion DNA polymerase (New England Biolabs, Ipswich, UK). The PCR products were then subjected to exome sequence capture using the SureSelect Human All Exon V6 (Agilent, Santa Clara, USA). The amplicons were size-checked and quantitated using a BioAnalyzer 2100 (Agilent, Santa Clara, USA), and then subjected to 2 × 150bp paired-end massively parallel sequencing using a HiSeq2000 Sequencing System (Illumina, San Diego, USA). After filtering out the candidate pathogenic variants by comparing the exome sequencing data of blood, bone, and skin tissue, validation was conducted with Sanger sequencing again.

2.5 | Micro-CT of bone tissues

The biopsied bone from patient 10 with MEL and corresponding control bone tissue from the volunteer were analyzed by SkyScan 1176 (Bruker MicroCT, Kontich, Belgium) operated at 65 kV with a 0.5 mm Al filter and 520 ms exposure time per projection. The scanning was captured at 0.35 degrees over a total of 180 degrees. The 1.6 version of NRecon software was used for 3D reconstruction and image viewing. Then, the segmentation was performed with the range from 20/130 to 120/255, and after segmentation, the inverted images were used to obtain the bone mass and pores. The parameters of bone mass include BMD (mg/cm³) and percent bone volume (BV/TV, %), and properties of pores including percent pore volume (Po.V/TV, %), pore diameters (Po.Dm, mm) and pore number (Po.N, 1/mm), were calculated in a cubic (1.2 mm*1.2 mm*1.2 mm) region of interesting (ROI) by CTan software (Bruker MicroCT, Kontich, Belgium). Three cubic regions in normal bone tissue and six cubic regions in biopsied osteophytes were selected for statistical analysis. The 3D rendering of the bones and pores was performed using CTvox software (Bruker MicroCT, Kontich, Belgium).

3 | RESULTS

3.1 | Clinical features

Table 1 listed the clinical features of the 10 sporadic MEL patients. There were 4 men and 6 women with an average age of 29.5 years (range 11–40 years), and the average age at symptoms onset was 21.2 years (range 10–38 years).
| Patient | Gender | Age (years) | Affected side | Affected sites | Age at symptom onset (years) | Symptoms and signs | Radiographic pattern | Skin lesions | Bone scan |
|---------|--------|-------------|---------------|---------------|-----------------------------|-------------------|---------------------|-------------|----------|
| 1       | Male   | 27          | Right         | Pelvis, femur, tibia, fibula, foot | 12 | Enlarged right lower extremity, bone pain | Mixed (classic+ osteoma-like+ myositis ossificans-like) | − | + |
| 2       | Female | 40          | Left          | Tibia, fibula  | 38 | Tibia protrusion, bone pain | Classic | − | + |
| 3       | Male   | 32          | Right         | Femur, patella, tibia | 16 | Tibia protrusion, bone pain, restricted movement | Mixed (classic+ osteoma-like) | − | NT |
| 4       | Female | 22          | Left          | Pelvis, femur, tibia, fibula, foot | 12 | Lower extremity deformity, restricted movement | Classic | − | NT |
| 5       | Female | 36          | Right         | Femur, tibia, foot | 33 | Bone pain, restricted movement | Mixed (classic+ osteoma-like) | − | NT |
| 6       | Male   | 32          | Right         | Hand          | 22 | Bone pain | Mixed (classic+ osteoma-like) | − | NT |
| 7       | Male   | 11          | Right         | Femur, tibia, fibula, foot | 10 | Bone pain | Mixed (classic+ osteoma-like) | − | NT |
| 8       | Female | 27          | Left          | Pelvis, femur, tibia, fibula, foot | 17 | Bone pain | Mixed (classic+ osteoma-like) | Scleroderma-like skin | + |
| 9       | Female | 31          | Left          | Femur, foot | 22 | Bone pain | Mixed (classic+ myositis ossificans-like) | − | NT |
| 10*     | Female | 37          | Left + spine  | Spine, pelvis, femur, tibia, fibula, foot | 30 | Bone pain, restricted movement | Mixed (classic+ osteoma-like+ myositis ossificans-like) | Scleroderma-like skin, linear epidermal nevus | + |

Abbreviations: NT, not tested; +, present; −, not present.

*A 37-year-old female who carries a somatic p.Gln56Pro variant of the MAP2K1 gene in affected bone tissue.
FIGURE 1 Radiographs, bone scintigraphy, and photographs of patient 10. (a) X-rays of the left knee joint show a “dripping candle-wax” appearance of hyperostosis in the left fibula, “myositis ossificans-like” pattern in the vicinity of the joint, “osteoma-like” hyperostosis in the endosteal surface of the left femur. (b) Radiographs of the left ankle joint reveal melorheostotic lesions in the left fibula, calcaneus, talus, navicular, cuboid, and cuneiforms. (c) X-rays of the pelvis demonstrated a “dripping candle-wax appearance” of MEL in L4, L5, left ilium, and periosteal and endosteal surfaces of the cortex of the left femur. (d) Bone scintigraphy displays larger areas than that shown by X-rays. (e–h) a classic linear epidermal nevus and scleroderma-like skin lesions extend from the patient’s posterior waist to the left thigh and leg following Blaschko lines
The disease manifested clinically with bone pain, restricted movement, and bone deformities. In the 10 patients, MEL lesions were involved with multiple contiguous bones unilaterally: the lesion sites were mainly located in the lower extremity, including femur (8/10), tibia (8/10), fibula (6/10), and foot (7/10), the next was pelvis (4/10), and the last were patella (1/10), hand (1/10), and spine (1/10). X-ray imaging revealed bony changes consistent with MEL, in which, 2 patients showed the classic MEL feature, namely “dripping candle wax” appearance of hyperostosis in periosteal or endosteal bone surfaces, 5 displayed a mixed pattern of “classic” and “osteoma-like” hyperostosis, and the remaining 3 manifested with a more complicated mixed pattern of “classic”, “osteoma-like” and “myositis ossificans-like” hyperostosis. 4 patients underwent a radionuclide bone scan, and all skeletal melorheostotic abnormalities demonstrated intensely elevated 99mTc-MDP uptake. Among all the patients, patient 10 had the most extensive bone involvement and relatively heavy disease burden (Figure 1). In addition to bone lesions, 2 patients also presented skin lesions. Patient 8 displayed a small SLS lesion in the anterior tibia, while patient 10 had a large cutaneous lesion, including LEN and SLS lesion, extended from her posterior waist to left thigh and leg following Blaschko lines (Figure 1).

### 3.2 Identification of somatic variant in MAP2K1

After Sanger sequencing on the LEMD3 gene, no pathogenic variants were found in the genomic DNA from peripheral blood samples of the 10 MEL patients. Then, exome sequencing was conducted on genomic DNA isolated from matched blood, biopsied bone, and skin samples of patient 10. Sequence alignment files were generated to duplicate removal, local realignment, and base quality recalibration with the Genome Analysis Toolkit (GATK). Variations including single-nucleotide variants (SNVs) and small insertions or deletions (Indels) were identified with variant quality score recalibration (VQSR). Then, SNVs with a minor allele frequency of < 1% in the Chinese 1000 genome database and SIFT score of < 0.05, Polyphen-2 score of > 0.85, or MutationTaster score of > 0.85 were considered as significant of not being benign. According to the above restrictions and comparison among blood, bone, and skin tissues, somatic variants were filtered from those that were present in the affected bone or skin tissue, but not present in the blood, and caused protein changes or splicing changes. Finally, the somatic missense variant in exon 2, the coding region of MAP2K1 (GenBank accession No. NM_002755.3), c.167A>C (p.Gln56Pro), causing amino acid substitution of glutamine to proline in the protein MEK1, was identified in the bone tissue of Patient 10 (Figure 2). However, no pathogenic somatic variants in MAP2K1 or other genes were found in the skin lesion tissue. The MAP2K1 variant, p.Gln56Pro, occurred at a conserved position in eight vertebrates based on Universal Protein Resource (Figure 2). The above MAP2K1 variant was then verified by Sanger sequencing to be absent in the control bone sample from the volunteer. Since we were unable to obtain the affected bone or skin tissues of the remaining 9 patients, we could not determine whether they carried pathogenic somatic variants in the MAP2K1 gene or other genes through exome sequencing.

### 3.3 Micro-CT of bone tissues

To explore the microstructures of bone with melorheostotic lesions, micro-CT with a resolution of 8.96 um was adopted to scan the osteophytes biopsied from Patient 10 and unaffected bone samples obtained from the volunteer, respectively. The 2D radiographies of micro-CT showed homogeneously dense cortical tissue with consistently sized pores in unaffected bone tissue, but increased pores with larger size were found in biopsied affected osteophytes with MAP2K1 variant (Figure 3). The 3D renderings segmented the calcified tissue and vascular pores, and clearly showed decreased bone volume and elevated porosity in the biopsied osteophytes (Figure 3). Quantitatively, BMD and percent bone volume (BV/TV) was decreased in affected bone versus unaffected bone. In addition, the increased porosity in affected bone was manifested with elevated percent pore volume (Po.V/TV), pore...
diameters \((P_{o.Dm})\), and pore numbers \((P_{o.N})\) versus unaffected bone (Table 2). The pore diameters also showed significantly different distributions (Figure 3). In unaffected bone, the maximum pore diameter was 0.134 mm, while the range of pore diameters was notably wider with the maximum pore diameter of 0.368 in affected bone tissue. The results showed significantly increased porosity due to increased tissue vascularity in the affected bone tissue.

4 | DISCUSSION

With the progress of research, the molecular pathogenesis of MEL is gradually revealed. In the majority of sporadic MEL cases, the causal somatic variants in \(MAP2K1\) and \(SMAD3\) are identified from melorheostotic bone (De Ridder et al., 2020; Kang et al., 2018). Most variants in the \(MAP2K1\), cluster in the negative regulatory domain of

TABLE 2  Comparison of microstructure characteristics between melorheostotic and unaffected bone tissues

|                | Unaffected | Affected | \(p\)-value |
|----------------|------------|----------|-------------|
| **BMD (g/cm\(^2\))** | 0.740 ± 0.001 | 0.673 ± 0.022 | .002** |
| **BV/TV (%)**    | 94.44 ± 2.25 | 76.41 ± 8.80 | .012* |
| **Po.V/TV (%)**  | 5.72 ± 2.27  | 23.91 ± 8.91 | .012* |
| **Po.Dm (mm)**   | 0.071 ± 0.013 | 0.141 ± 0.012 | <.001** |
| **Po.N (/mm)**   | 0.781 ± 0.185 | 1.676 ± 0.573 | .037* |

Abbreviations: BMD, bone mineral density; BV/TV, percent bone volume; Po.Dm, pore diameters; Po.N, pore numbers; Po.V/TV, percent pore volume. *\(p < .05\); **\(p < .01\).
MEK1 (p.Gln56Pro, p.Lys57Glu, and p.Lys57Asn), causing loss of inhibitory effect, activating MEK1 in the RAS signaling pathway and following increasing phosphorylation of ERK, which promote osteoblast proliferation and reduce their differentiation. A novel variant (p.Cys121Ser) in the catalytic domain is also identified as associated with MEL (De Ridder et al., 2020). In some MAP2K1-negative cases, somatic SMAD3 variants are identified, which up-regulate the TGF-β/SMAD pathway (Kang et al., 2018; Kang et al., 2020). In MEL cases that appeared based on OPK or BOS, germline deactivating variants in LEMD3 influence the negative regulation of MAN1 on SMAD transcription factors in TGF-β/BMP superfamily signaling (Couto et al., 2007; Hellemans et al., 2004). Somatic variant in KRAS located upstream of the RAS signaling pathway was also detected in the SLS of a case with polyostotic MEL and familiar OPK caused by germline LEMD3 variant (Whyte et al., 2017). This may implicate variants in the RAS signaling pathway can be the “second hit” upon the LEMD3 haploinsufficiency, and abnormalities in RAS and TGF-β signaling pathways lie at the pathogenic heart of MEL. Considerable crosstalk between RAS and TGF-β signaling and RUNX2 activation via MAPK and PI3K/AKT pathway made the mechanism of MEL more complicated and attractive (Cohen-Solal et al., 2015; Grusch et al., 2010; Vivekanandhan & Mukhopadhyay, 2019). In this study, we failed to find pathogenic germline variants in LEMD3 in any of the ten MEL patients, but a somatic MAP2K1 (p.Gln56Pro) variant was identified in the bone lesion of one patient. However, no pathogenic somatic variants in MAP2K1 or other genes were found in the skin lesion tissue of the same patient. Unfortunately, we were unable to collect the affected bone and skin tissues of the remaining 9 patients, since they did not undergo relevant surgery in our hospital. The difficulty of obtaining bone and skin samples may be a big obstacle to detect somatic variants. To the best of our knowledge, this is the first report of a Chinese MEL patient carrying a somatic MAP2K1 variant, which is a rare but apparently specific invariant for MEL. The finding may further confirm the pathogenicity of somatic MAP2K1 variants and validate the molecular mechanism of MEL in Chinese patients.

In Table 3, the clinical features of ten Chinese MEL patients were summarized and compared with those in the largest published clinical review about MEL in Mayo Clinic (Smith et al., 2017). The average age of our patients seems to be younger, and the male-to-female ratio seems to be higher. Concordant with Mayo Clinic’s findings, we found recurrent pain, movement limitation, and limb deformity were the major presenting concerns, but numbness and weakness were not observed in our patients (Smith et al., 2017). MEL is typically limited to one limb but more widespread involvement is not unusual (Wordsworth & Chan, 2019). Unilateral limb involvement was present in 90% of our cases, and the affected lesions were contiguous and often crossed joint lines. Among all sites, appendicular skeleton especially lower extremities including femur, tibia, fibula, and foot were more commonly affected, while axial skeleton including skull and vertebrae was rare. The average number of affected sites per patient of 3.6 was similar to that of 4.4 in the Mayo cohort. In addition to classic “dripping candle wax” radiological features, three other radiologic presentations have also been described about MEL: (1) “osteoma-like” hyperostosis, which involved the endosteal surface and orientated in the long axis of the affected bone, (2) “osteopathy striata-like” hyperostosis, showing unilateral, long and dense striations near the inner surface of the cortex, (3) “myositis ossificans-like” pattern, with or without intraosseous hyperostosis.
(Freysschmidt, 2001). In our study, most cases displayed a mixture of one or more of the above patterns, which may confuse the diagnosis of MEL with OPK, BOS, and osteopathia striata (Ethunandan et al., 2004; Gnoli et al., 2019; Ihde et al., 2011; Mumm et al., 2007; Zhang et al., 2009). The microstructure of melorheostotic lesion showed two strikingly different regions including the outer periosteal apposition (PA) tissue with little porosity and deeper osteonal remodeling (OR) bone with more porosity according to Fratzl-Zelman et al. (2019). In this study, Micro-CT of “myositis ossificans-like” bone biopsied from the patient showed significantly increased porosity with increased tissue vascularity in OR bone, however, no PA region was observed, which may be due to differences in hyperostosis types and biopsy locations. Moreover, consistent with the literature, a large area of LEN and SLS were observed overlying the affected bones in the female patient (Jha et al., 2019; Rhys et al., 1998; Woolridge et al., 2005).

5 | CONCLUSION

To conclude, this is a summary of clinical features of Chinese MEL patients and the first study to identify somatic MAP2K1 variant in Chinese MEL patients. Our findings further validate the molecular genetic mechanism of MEL and broaden its phenotype spectrum in different populations.

AUTHOR CONTRIBUTIONS
Xiaojun Han, Yang Xu: Investigation, Formal analysis, Data collection, Writing-original draft; Zhiyang Wei: Data collection; Formal analysis; Chun Wang: Funding acquisition, Supervision; Hua Yue, Zhenlin Zhang: Conceptualization, Funding acquisition, Supervision, Writing-review&editing. All authors read and approved the final version of the manuscript.

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CONFLICT OF INTEREST
The authors declare no conflicts of interest.

DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available from the corresponding author upon reasonable request.

ETHICS STATEMENT
This study was approved by the Ethics Committee of the Shanghai Jiao Tong University Affiliated Sixth People’s Hospital, and written informed consent was obtained from all the patients and healthy volunteers. All methods were performed in accordance with the Declaration of Helsinki.

ORCID
Hua Yue © https://orcid.org/0000-0002-9179-5894
Zhenlin Zhang © https://orcid.org/0000-0002-4339-1617

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