Abstract. Epidemiological investigations indicate that certain ingredients in tea bricks can antagonize the adverse effects of fluoride. Tea polyphenols (TPs), the most bioactive ingredient in tea bricks, have been demonstrated to be potent bone-supporting agents. ClC-7 is known to be crucial for osteoclast (OC) bone resorption. Thus, in this study, we investigated the potential protective effects of TPs against fluorosis using a mouse model and explored the underlying mechanisms with particular focus on ClC-7. A total of 40, healthy, 3-week-old male C57BL/6 mice were randomly divided into 4 groups (n=10/group) by weight as follows: distilled water (control group), 100 mg/l fluoridated water (F group), water containing 10 g/l TPs (TP group) and water containing 100 mg/l fluoride and 10 g/l TPs (F + TP group). After 15 weeks, and after the mice were sacrificed, the long bones were removed and bone marrow-derived macrophages were cultured ex vivo in order to perform several experiments. OCs were identified and counted by tartrate-resistant acid phosphatase (TRAP) staining. The consumption of fluoride resulted in severe fluorosis and in an impaired OC function [impaired bone resorption, and a low mRNA expression of nuclear factor of activated T-cells 1 (NFATc1), ATPase H\(^{+}\) transporting V0 subunit D2 (ATP6v0d2) and osteopetrosis-associated transmembrane protein 1 (Ostm1)]. In the F + TP group, fluorosis was attenuated and OC function was restored, but not the high bone fluoride content. Compared with the F group, mature OCs in the F + TP group expressed higher mRNA levels of ClC-7 and Ostm1; the transportation and retaining of Cl\(^{-}\) was improved, as shown by the fluorescence intensity experiment. On the whole, our findings indicate that TPs mitigate fluorosis in C57BL/6 mice by regulating OC bone resorption. Fluoride inhibits OC resorption by inhibiting ClC-7 and Ostm1, whereas TPs attenuate this inhibitory effect of fluoride.

Introduction

Fluorosis, a disease caused by excessive fluoride intake through water and food, has been documented repeatedly in many field investigations and community surveys in a number of countries worldwide, such as Africa, China, Germany, India, United Republic of Tanzania and the US (1). In China, fluorosis includes 3 types: water-type fluorosis, coal-burning-type fluorosis and brick tea-type fluorosis. Excessive exposure to fluoride can lead to disruptions in bone metabolism and enamel development, causing skeletal fluorosis and dental fluorosis, respectively. Dental fluorosis is a less serious disease that causes a chalk-like discoloration of teeth with unsightly mottled spots or lines on tooth enamel. However, skeletal fluorosis is a crippling disease that causes pain and damage to bones and joints (2). Moreover, there is no established strategy to date for the treatment of patients with skeletal fluorosis. Thus, it is crucial to to obtain a better understanding of the mechanisms regulating the pathogenesis of fluorosis for the prevention of the potential health hazards of fluoride.

The characteristic pathological change associated with fluorosis is the acceleration of bone turnover, in which osteoclasts (OCs) play a critical role (3). An OC is a type of bone cell that disassembles and digests the composite of hydrated protein and mineral by secreting acid and collagenase, a process known as bone resorption. OCs play a critical role in the maintenance, repair and remodeling of bones. Monocytes/macrophages are recognized as OC precursors. Monocyte/macrophage precursors undergo 3 stages during their differentiation into functionally mature OCs. First, mononuclear pre-osteoclasts fuse together to become non-functional multi-nucleated OCs expressing tartrate-resistant

ClC-7/Ostm1 contribute to the ability of tea polyphenols to maintain bone homeostasis in C57BL/6 mice, protecting against fluorosis

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acid phosphatase (TRAP) and lacking ruffled membranes (RM); finally, non-functional multi-nucleated OCs are fully developed into functional OCs which secrete H+ and enzymes (4-8). This differentiation process is regulated by various factors. Nuclear factor of activated T-cells (NFATc1), a master regulator of osteoclastogenesis, regulates a number of OC-specific genes, such as TRAP, cathepsin K and ATPase H+ transporting V0 subunit D2 (ATP6v0d2), which are known to be involved in bone resorption. It has been demonstrated that NFATc1 binds directly to the promoter regions and induces the expression of fusion-mediating molecules, such as ATP6v0d2, thereby regulating OC multi-nucleation (9). In mature OCs, ATP6v0d2, as a component of V-type H+-ATPase, mediates the acidification of the resorption lacuna, which is a prerequisite for the secretion of enzymes (10,11). The efficient acidification of the resorption lacuna requires the neutralizing current through chloride channel-7 (CIC-7). CIC-7 is a Cl-/H+ antiporter, that resides mainly in late endosomes, lysosomes and RM of multi-nucleated OCs (12-14). The stability of CIC-7 depends on its association with osteopetostosis-associated transmembrane protein 1 (Ostm1), and Ostm1 serves as a β-subunit to support the function of CIC-7 (15,16). It has been demonstrated that the loss of function of CIC-7 or of its association with Ostm1 in humans and mice result in osteopetrosis (12). Thus, CIC-7 is crucial for OC bone resorption. Our previous study reported that fluoride decreased OC bone resorption via the inhibition of NFATc1 (45); however, to date, it has not been determined whether CIC-7 in OCs is involved in the pathogenesis of fluorosis.

The brick tea-type fluorosis caused by the consumption of fluoride from drinking tea, particularly brick tea compressed by green tea or black tea. It is a public health concern in the Northwest China, where minorities drink brick tea as drinking water. It is estimated that 31 million individuals in that region are affected (17). In addition, it is considered that the dental fluorosis detection rate in children and the skeletal fluorosis detection rate in areas affected by the brick tea-type fluorosis are higher than the rates in areas affected by water type-fluorosis (18); however, a lower number of patients with severe dental fluorosis and disabilities were found in areas affected by the brick tea-type fluorosis. Thus, it is worthy to explore whether some ingredients in brick tea can antagonize the adverse effects of fluoride. It is proposed that oxidative stress plays a critical role in the pathogenesis of fluorosis (19,20). Tea polyphenols (TPs), the most bio-active ingredient in brick tea (21), have received considerable attention due to their antioxidant effects and they have been shown to exert beneficial effects on health, including maintaining bone mass (22,23). Nevertheless, the potential effect of TPs on fluorosis and the underlying mechanisms have not yet been elucidated. The role of OCs in skeletal fluorosis has attracted increased attention (24,25,45). TPs have been demonstrated to be potent bone-supporting agents. Therefore, the present study aimed to investigate the potential protective effects of TPs against fluorosis using a mouse model and to explore the underlying mechanisms with particular focus on CIC-7.

Materials and methods

Animals and treatments. A total of 40 healthy, 3-week-old male C57BL/6 mice (Vital River Laboratories Animals Technology Co., Ltd., Beijing, China) were allowed to acclimatize to an environmentally controlled room with a 12-h light/dark cycle in an animal care facility for 7 days before beginning the experiment. Subsequently, these mice were randomly divided into 4 groups (n=10/group) by weight as follows: the distilled water (control group), 100 mg/l fluoridated water (F group), water containing 10 g/l TP (TP group) and water containing 100 mg/l fluoride and 10 g/l TP (F+ TP group). The animals were allowed to drink the experimental water for 15 weeks. Sodium fluoride (guaranteed reagent) was purchased from Tianjin Guangfu Fine Chemical Research Institute (Tianjin, China). TPs with a purity >98% were purchased from Zhengzhou Chengda Chemical Products Co., Ltd. (Zhengzhou, China), and the catechins account for approximately 70% of the extractable solids. Distilled water mixed with TPs and/or fluoride and was prepared fresh daily. Body weights were monitored weekly throughout the treatment period. At the end of the experiment, all mice were euthanized by ether asphyxiation. The 4 legs were aseptically removed and dissected free of adherent soft tissue without disarticulation at the joint. All animal procedures were approved by the Animal Care and Use Committee of Harbin Medical University, Harbin, China.

Dental fluorosis phenotyping. The dental fluorosis status of each mouse was evaluated and scored by 2 independent examiners based on the following modified Dean Index (26,27) (Fig. 1A).

Assessment of skeletal fluorosis. To measure bone mineral density, the humerus bones of the mice were excised and cleaned by removing the soft tissues, splitting open and removing all marrow with cold PBS. The specimens were dried for 4 h at 105°C and were then weighed. Prior to ashing, the bones were heated in a smoke-free environment in a fume hood. The bones were then ashed for 6 h at 550°C and weighed again, dissolved in 1N HCl, and neutralized by the addition of 1N NaOH. A standard method with the ion-specific F electrode (Shanghai Weiye Instrument Plant, Shanghai, China) was used to determine the F- concentration of each dissolvent in the Institute for Endemic Fluorosis Control, Center for Endemic Disease Control, Chinese Center for Disease Control and Prevention (Harbin Medical University).

Determination of bone fluoride content. Femur bone samples were collected and cleaned by removing the soft tissues, splitting open and removing all marrow with cold PBS. The specimens were dried for 4 h at 105°C and were then weighed. Prior to ashing, the bones were heated in a smoke-free environment in a fume hood. The bones were then ashed for 6 h at 550°C and weighed again, dissolved in 1N HCl, and neutralized by the addition of 1N NaOH. A standard method with the ion-specific F electrode (Shanghai Weiye Instrument Plant, Shanghai, China) was used to determine the F- concentration of each dissolvent in the Institute for Endemic Fluorosis Control, Center for Endemic Disease Control, Chinese Center for Disease Control and Prevention (Harbin Medical University).

In situ bone OC numbers. The long bones were transected at a distance from the knee joints to avoid damaging the joint. Following the removal of the soft tissues, the joints were fixed in 10% formaldehyde for 7 days. The formalin-fixed tissues were then decalcified in 5% EDTA for 1 month prior to paraffin-embedding. The paraffin-embedded specimens were then sectioned (6 µm). The slides were stained using the TRAP staining kit (Shanghai Sunbio Co., Shanghai, China) and counterstained with hematoxylin. The number of TRAP+ cells was counted across 4 fields of view just under the growth plate of each bone section by light microscopy at x20 magnification.
in an Olympus BX-51 microscope (Olympus Corp., Tokyo, Japan) and digital images were captured. Data were reported as the mean number of TRAP<sup>+</sup> cells within the 4 fields per bone section as the formation of OCs.

**Assessment of OC ultrastructure in the transmission electron microscope.** The metaphyses were fixed in 2.5% glutaraldehyde for 4 weeks at 4°C and the fixed tissues were decalcified in 10% EDTA for 4 weeks. The specimens were post-fixed for 2 h in 2% aqueous OsO<sub>4</sub>, dehydrated in acetone, and embedded in Epon (ElectroN Microscopy Sciences, Hatfield, PA, USA). Ultrathin sections were cut from blocks and then stained with uranyl acetate and lead citrate. The specimens were observed under an H-7650 transmission electron microscope (Hitachi, Ltd., Tokyo, Japan), as previously described (29,30). The cell surface structures, facing matrix, were defined as RM, an area with multiple folding of the cell membrane and clear zone (CZ), attachment of the cell to the bone surface.

**Immunohistochemistry.** The sections were treated with 0.3% H<sub>2</sub>O<sub>2</sub> in ethanol to inactivate endogenous peroxidase. The sections were digested with 0.2% Triton (Amresco, LLC, Solon, OH, USA), blocked with 5% BSA (SABC kit, Wuhan Boster Biological Technology, Ltd., Wuhan, China), incubated for 2 h with a 1:100 dilution of anti-rabbit ClC-7 antibody (ab86196; Abcam, Cambridge, UK) at 37°C, treated with a biotinylated anti-rat antibody, and DAB chromogenic liquid (SABC kit). The sections were counterstained with hematoxylin.

**Osteoclastogenesis.** Bone marrow-derived macrophages (BMMs) were prepared as described previously (4,31) and osteoclastogenesis was induced by receptor activator of NF-κB ligand (RANKL) and macrophage colony-stimulating factor (M-CSF). The femora and tibiae were aseptically removed and dissected free of adherent soft tissue. The bone ends were removed and the marrow cavity was flushed out into a Petri dish by slowly injecting α-MEM at one end of the bone using a sterile 5 ml syringe. The bone marrow suspension was passed repeatedly through a glass pipette to obtain a single cell suspension. The bone marrow cells were then resuspended in α-MEM, incubated for 24 h, and plated in 6-well plates with thye addition of 5 ng/ml M-CSF (Peprotech, Inc., Rocky Hill, NJ, USA) to deplete the stromal cells of cell preparations. Non-adherent cells were replated in 12-well plates at a density of 2x10<sup>6</sup> cells/well for extracting DNA and protein or in 96-well plates at a density of 1x10<sup>5</sup> cells/well in α-MEM for OC identification, containing 30 ng/ml M-CSF for 3 days to obtain the BMMs. Each group included 5 parallel wells. The BMMs were incubated with 30 ng/ml M-CSF and 100 ng/ml RANKL (Peprotech, Inc.) for another 4 days which were named day 1 to 4 accordingly. The medium was replaced every 2 days.

OCs were identified and stained using the TRAP staining kit (Shanghai Sunbio Co.) on days 3 and 4. TRAP<sup>+</sup> multinucleated cells were counted as total OCs (≥3 nuclei) or giant OCs (≥20 nuclei). Data are presented as the means ± SD.

**Determination of NFATc1 and ATP6v0d2 mRNA expression in OCs in vitro.** Gene expression in the cells was assessed by quantitative PCR (qPCR). The induced cells on day 4 were dissolved in TRIzol reagent and total RNA was extracted using the total isolation kit (Takara Bio, Dalian, China). First-strand cDNA was synthesized using 1 µg total RNA with the PrimeScript RT reagent kit with gDNA Eraser according to the manufacturer’s instructions (Takara Bio). qPCR was performed on the Real-Time Fluorescence PCR Instrument (Bio-Rad Laboratories, Inc., Hercules, CA, USA) using SYBR-Green chemistry. The following commercial available primer were used for PCR: GAPDH (forward, 5'-CTGACGTGCAGCCCT GGAGAAGC-3', and reverse, 5'-CCGGGCTCAAGGTGG AAGAGT-3'), NFATc1 (forward, 5'-CTGCAA CGGGAAACG GAAGAGAAG-3', and reverse, 5'-TATACCCCAAGA CGGCATAGC-3'), ATP6v0d2 (forward, 5'-ACAGACGCGC TTATATCACTACCTC-3', and reverse, 5'-ATCCCCCTTCC TTTCCTCAATAAC-3') and Ostn1 (forward, 5'-GGGGAG AAACAAGGAGCACAACAA-3'; reverse, 5'-ATACCCCAA CACTCTGTCTCCT-3'). The relative mRNA expression levels were calculated based on threshold cycle number (Ct) and normalized to GAPDH using the comparative Ct method.

**Western blot analysis.** The cells were lysed in RIPA buffer (Beyotime Institute of Biotechnology, Shanghai, China) and protein concentrations of cellular extract were measured using an enhanced BCA Protein Assay kit (Beyotime Institute of Biotechnology). An automated capillary-based simple western system (Simon; ProteinSimple, Santa Clara, CA, USA) was then used to quantify the protein levels of CIC-7. Compared to the traditional western blot analysis, automation allows for the more accurate and reproducible assessment of protein levels, as previously demonstrated (32). In brief, 16 ng of protein samples were used for analysis. The proteins were immunoprobed using anti-CIC-7 antibody (ab86196; Abcam) or anti-GAPDH antibody (G9545; Sigma, St. Louis, MO, USA). Other reagents were using Simon-Rabbit (15-150 kDa) Master kit provided by ProteinSimple (San Jose, CA, USA). For quantitative analysis, the CIC-7 signals were normalized against GAPDH.

**Chloride permeation.** Cl permeation was measured using the fluorescent Cl<sup>-</sup> indicator, N-ethoxycarbonylmethyl-6-methoxy quinolinium bromide (MQAE), as previously described (33). The mechanism of detection is diffusion-limited collisional quenching of MQAE fluorescence. The degree of fluorescence quenching in MQAE indicate an increase in the Cl<sup>-</sup> concentration. OCs were loaded with 6 mmol/l MQAE in a chloride-free buffer containing 140 mmol/l NaNO<sub>3</sub>, 5 mmol/l KNO<sub>3</sub>, 5 mmol/l 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), 1 mmol/l Mg(NO<sub>3</sub>)<sub>2</sub>, and 5 mmol/l glucose for 60 min at 37°C. Just before detection, the culture buffer was changed to a chloride buffer of similar composition with Cl<sup>-</sup> as the substituting anion. At the very moment, OCs were exposed to excitation light for 20 times at an interval of 5 sec on the stage of an inverted microscope (Nikon Diaphot; Nikon, Tokyo, Japan) and the digitized image was recorded. The original fluorescence intensity was measured by changing from a chloride buffer to chloride-free buffer at time zero. The mean relative intensity was calculated by dividing the fluorescence intensity by the original fluorescence, as previously described (34).

**Statistical analysis.** All experiments were repeated 5 times, and representative results are presented. Data are presented
as the means ± SD and analyzed using SPSS 16.0 software. A value of P<0.05 was considered to indicate a statistically significant difference. The normality of distribution and homogeneity of variance were tested. Dental fluorosis among the 4 groups was analyzed using the Chi-square test. The IOD of the humerus bone, TRAP⁺ cells in vivo and in vitro, the relative mRNA expression of 4 genes, and CIC-7 protein levels in vitro were analyzed by two-way ANOVA followed by Fisher’s LSD post hoc test to evaluate the effects of fluoride, TPs or their combination. Linear by linear association was used to evaluate the association between the fluorescence intensity and detection time.

Results

Body weight and bone fluoride. Neither fluoride nor TP significantly affected the body weights of the mice throughout the study period (data not shown). The fluoride content in long bone was detected to reflect the difference of fluoride accumulation among the groups. The data indicated that the bone fluoride content in the F group (4098.64±58.84 mg/kg) and F + TP group (3855.70±140.77 mg/kg) was significantly higher than that in the control group (876.26±14.46 mg/kg) (P<0.001 and P=0.002, respectively); however, there was no difference between the F group and F + TP group. The fluoride content in the TP group (970.27±17.05 mg/kg) was similar to that of the control group (876.26±14.46 mg/kg) (data not shown).

Dental fluorosis. Compared with the mice receiving distilled water or TPs, the mice receiving fluoride (F and F + TP groups) had a higher prevalence of dental fluorosis. From the 6th week, the teeth of the mice in the F group were characterized by a change in color and thin white horizontal lines. At the 15th week, half of the mice in the F group had severely fluorosed enamel with discrete or confluent pitting. It is worth noting that the prevalence of dental fluorosis in the F + TP group was lower than that in the F group (P<0.05; Fig. 1B).

Skeletal fluorosis. A radiographic examination of the excised humerus bones confirmed the status of long bone growth (Fig. 1C). Marrow space, growth plate and cortical margins were apparent in the control specimens. However, the specimens in the F group exhibited a thickened cortical margin and a brighter performance. However, there was no apparent change in intensity in the TP group and F + TP group by the naked eye. The IOD of the humerus (Fig. 1D) in the F group was significantly higher than that in the control group (P=0.024). Of note, the IOD of the humerus in the F + TP group was significantly lower than that in the F group (P=0.033); there was thus an interaction between fluoride and TPs (P=0.015).

In situ bone OC numbers. TRAP⁺ cells were detected as a measure of OC formation in situ and the majority of these cells appeared in the zone of cartilage calcification (Fig. 2A). The number of OCs was consistently elevated in all the treatment groups (Fig. 2B). Specifically, compared with the control group, the number of OCs increased 7-fold in the TP group, 5-fold in the F + TP group and 3-fold in the F group. Furthermore, the OCs in the TP group were more intense than those in the other groups by the naked eye.

OC ultrastructure. Ultrastructural analyses of the proximal metaphyses of femurs were performed using a transmission electron microscope (Fig. 2C). In the sections from the control group and TP group, the OCs with well-developed RM and CZ were frequently attached to the bone and these features
were considered to reflect capacity of bone resorption. Many vesicles containing electron-dense material (arrows) were also observed in the OCs. By contrast, a large proportion of OCs in the F group lacked the morphological signs of proper polarization, e.g., RM and CZ. Instead, the OCs exhibited a matrix-facing area with no characteristic cell surface structures, which can be considered as an immature developed cell. Moreover, the OCs in F + TP group exhibited vacuolation degeneration.

Immunohistochemistry of ClC-7 protein in long bone. Immunohistochemical staining of ClC-7 was performed on proximal metaphyses of femurs (Fig. 3). In the control group, intense immunostaining was observed in the connecting zone of cartilage calcification and ossification. The location of ClC-7 protein was similar among the 4 experimental groups.

OC potential. To examine the effects of fluoride and/or TPs on OC formation, TRAP-positive multi-nucleated OCs were observed in all groups (Fig. 4A). On day 3, the number of OCs (Fig. 4B) in the TP and F + TP groups was significantly higher than the control group (P=0.031 and P<0.001, respectively) and the number in the F + TP group was significantly higher than that in the F group (P=0.023). On day 3, the number of giant OCs (Fig. 4C) in the TP and F + TP groups was significantly higher than the control group (P=0.014 and P=0.002, respectively) and the number in the F + TP group was also significantly higher than that in the F group (P=0.030). These differences were evident until day 4. On day 4, the number of giant OCs in the 4 groups increased, compared with that at day 3. Of note, the number of giant OCs in the control group was significantly lower than the other 3 groups.

mRNA expression of NFATc1, ATP6v0d2 and Ostml in OCs in vitro. We examined the effect fluoride and/or TP administration on the mRNA expression of NFATc1 and ATP6v0d2 (Fig. 4D). The expression of NFATc1 and ATP6v0d2 in the F group was significantly lower than that in the control group (P=0.043 and P=0.039, respectively). The mRNA of both genes in the F + TP group was significantly higher than that in the F group (NFATc1, P=0.009; ATP6v0d2, P=0.002).

mRNA expression of Ostml in OCs in vitro. Ostml mRNA expression in the F group decreased almost 600-fold compared to the control group (Fig. 5A). The mRNA expression of Ostml in the F + TP group was significantly higher than that in the F group (Ostml, P=0.033).

ClC-7 expression in OCs in vitro. Giant OCs were regarded as having absorbing capacity, and thereafter the protein level of ClC-7 in the F group (0.08±0.02) was significantly lower than that in the control group (0.15±0.02, P=0.006). ClC-7 protein expression levels in the TP group (0.24±0.02) and F + TP group (0.16±0.05) were both significantly lower than the level in the F group (P=0.012 and P=0.03, respectively). There was thus an interaction between fluoride and TPs (P=0.029).

Fluorescence intensity in OCs in vitro. The ability of ClC in OCs to transport Cl⁻ in all groups was detected on day 4 (Fig. 5D).
Figure 3. Immunohistochemical staining of ClC-7. Representative images of osteoclasts under the growth plate of the (A) control, (B) fluoride (F), (C) tea polyphenol (TP); and (D) fluoride and tea polyphenol (F + TP) (FT) groups. Location of protein expression of ClC-7 was similar among the 4 experimental groups. Magnification, x200.

Figure 4. Osteoclast formation and function-related gene mRNA expression. (A) Formation of osteoclasts differentiated from BMMs isolated from mice. Morphology of osteoclasts was similarly among the 4 groups. (B and C) Osteoclasts were counted as total osteoclasts (≥3 nuclei) or giant osteoclasts (≥20 nuclei). Data are presented as the means ± SD. (D) Giant osteoclasts were generally considered as mature osteoclasts. NFATc1 and ATP6v0d2 mRNA expression on day 4 was determined. *P<0.05 vs. control; #P<0.05 vs. F group; &P<0.05 vs. the same group on day 3. F, fluoride group; TP, tea polyphenol group; FT, fluoride and tea polyphenol group; BMMs, bone marrow-derived macrophages.
Within 90 sec, the fluorescence intensity was continuously declining in the TP and F + TP groups (P<0.001 and P=0.024, respectively), suggesting that Cl\textsuperscript{-} could be accumulated in OCs in these groups; however, the fluorescence intensity in the F group exhibited an increasing trend (P<0.001), suggesting that OCs in the F group did not retain Cl\textsuperscript{-}.

**Discussion**

The skeletal fluorosis phenotype of the murine model and human disease is characterized by a combination of osteosclerosis, osteomalacia and osteoporosis in varying degrees. The low bone mass in C57BL/6 mice appears to be sensitive to the effects of fluoride on bone cells (35-39). Therefore, in the present study, C57BL/6 mice were utilized to investigate the potential protective effects of TPs against chronic fluorosis. In the F group, the mice had a higher prevalence of dental fluorosis and severe fluorosed enamel with discrete or confluent pitting. A radiographic examination of excised humerus bones revealed a thickened cortical margin and significantly higher IOD values. Such results were in agreement with those of another study using female C57BL/6 mice (35), indicating that fluorosis model was successfully constructed. The bone fluoride content remained consistent between the F group and F + TP group, suggesting that TP supplementation did not affect the accumulation of fluoride in the bone. However, compared to the F group, the F + TP group exhibited a significant mitigation of fluorosis, as shown by the prevalence of dental fluorosis and IOD value of the humerus, suggesting that TPs exerted a beneficial effect on bone homeostasis, protecting against the toxic effects of fluoride.

In our mouse model, the consumption of 100 mg/l fluoride stimulated bone formation. Bone remodeling is a complex, yet coordinated process that is responsible by two main types of cells, osteoblasts and OCs. Fluoride, as an anabolic agent, is capable of promoting osteoblast proliferation, whereas the influence of fluoride on OCs remains controversial. In our study, 100 mg/l fluoride increased the number of TRAP\textsuperscript{+} cells in the region below the growth plate, suggesting that fluoride stimulated the formation of OCs, which is consistent with a previous report using the C3H/HeJ (C3H) inbred mouse strain with 50 mg/l fluoride for 3 weeks (37). However, ultrastructural analysis revealed that the OCs in the F group exhibited an underdeveloped RM and CZ, and failed to attach to the bone matrix, which is consistent with previous animal and cell studies (40,41). It is known that OCs resorb bone by establishing a circle between themselves and the bone surface, and that the formation and development of specific membrane structures, such as RM and CZ, are important for bone resorption (42). Moreover, patients with defective acidification have increased numbers of OCs with decreased bone resorption (43,44). Therefore, our results suggested that fluoride increased the number of OCs; however, these OCs exhibited defective acidification and could not resorb bone matrix. Compared to the F group, mice in F + TP group had a greater number of TRAP\textsuperscript{+} cells in the region below the growth plate, and most OCs exhibited a well-developed RM and CZ, suggesting that TP alleviated defective acidification induced by fluoride. OCs are terminally differentiated multi-nucleated cells derived from mononuclear hematopoietic cells of the monocyte/macrophage lineage in bone marrow. RANKL and M-CSF are essential and sufficient for osteoclastogenesis from BMMs. Thus, primary BMMs isolated from mouse long bones are utilized as OC precursors to generate mature OCs *ex vivo*. In this study, we isolated primary BMMs from differently treated mice to observe the effects of fluoride and/or TPs on OC formation and
for ex vivo osteoclastogenesis assays. We noted that fluoride had little effect on osteoclastogenesis; this result was similar to that of our previous study (45); the number of OCs (≥3 nuclei) or the number of giant OCs (>20 nuclei) in mice in the F + TP group was greater than that in the F group on day 3, suggesting that TPs promoted osteoclastogenesis in association with exposure to high levels of fluoride.

Although the molecular mechanisms by which fluoride regulates bone resorption have not yet been illustrated, a previous in vitro study by our group indicated that fluoride inhibited the resorption activity of OCs by inhibiting NFATc1 and its downstream genes, such as Atpt6v0d2 (45). NFATc1 plays an important role in osteoclastogenesis by regulating the fusion of pre-osteoclasts and bone resorption. Thus, we investigated the effect of fluoride and/or TPs on the expression of NFATc1 and Atpt6v0d2 in new OCs at the mRNA level. Our results revealed that the expression of NFATc1 and Atpt6v0d2 in the F group was significantly lower than that in the control group, which is consistent with our previous study, whereas the mRNA expression of both genes in the F + TP group was significantly higher than that in the F group, suggesting that TP attenuated the inhibitory effect on these two genes induced by fluoride.

CIC-7, as one member of the CIC family, may support bone resorption by electrically shunting the H^+-ATPase that acidifies the resorption lacuna of OCs, or by facilitating the insertion of proton-pumping containing vesicles into the RM. It is known to be associated with osteopetrosis, that is characterized by OC dysfunction, impaired bone resorption and poor bone remodeling (48). In osteopetrosis, OCs are developed without an RM and fail to generate an acidic bone resorption compartment or resorb bone. In our model, OCs in the F group exhibited an underdeveloped RM and CZ, and failed to attach to the bone matrix; thus, we investigated the effect of fluoride and/or TPs on CIC-7. In the present study, the CIC-7 level in the F group was significantly lower than that in the control group; however, CIC-7 expression in the F + TP group was significantly higher than that in the F group, suggesting that TPs attenuated the inhibitory effect on CIC-7 protein induced by fluoride.

CIC-7, as a Cl^-/H^+ antiporter, is important for lysosomal acidification. It is the primary Cl^- permeation pathway in lysosomes (15). Thus, we further detected Cl^- permeation using fluorescent the Cl^- indicator, MQAE (33). The degree of fluorescence quenching in MQAE indicates an increase in Cl^- in cells. In the F group, new OCs did not retain Cl^-; however, new OCs in the F + TP group retained Cl^-: These results were consistent with the change in the protein expression of CIC-7. CIC-7 can reach the lysosome and can be degraded in the lysosome due to the lack of N-linked glycosylation sites. Ostml, a type I membrane protein with heavily glycosylated amino-terminal portion and a short cytosplasmic tail, shields CIC-7 from degradation by lysosomal proteases (15,16,49). Thus, we investigated the influence of fluoride and/or TPs on Ostml protein at the mRNA level. Our results indicated that in new OCs in the F group, Ostml mRNA expression was significantly lower than that in the control group; however, Ostml mRNA expression in the FT group was significantly higher than that in the F group, suggesting that fluoride decreased CIC-7 expression by inhibiting Ostml; TPs, however, attenuated this effect. This may be one of the reasons as to why the severity of fluorosis in areas affected by brick tea-type fluorosis is milder than that in areas affected by water-type fluorosis. Thus, perhaps the use of TPs could be considered for the prevention of fluorosis in areas affected by water-type fluorosis.

In conclusion, in the present study, we demonstrated that TPs exerted beneficial effects against fluorosis by regulating OC resorption ability. We reported that fluoride inhibited OC resorption by decreased CIC-7 and Ostml expression, whereas TPs attenuated these inhibitory effects of fluoride.

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