Cross-talk between primary osteocytes and bone marrow macrophages for osteoclastogenesis upon collagen treatment

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Homeostasis of osteoclast formation from bone marrow macrophages (BMM) is regulated by paracrine signals of the neighbourhood bone cells particularly mesenchymal stem cells (MSC), osteoblasts and osteocytes (OC). Besides paracrine cues, collagen and glycosaminoglycan are involved in controlling bone homeostasis. Towards this approach, different molecular weight collagens were reacted with MSC, OC and BMM to understand the bone homeostasis activity of collagen. The up-regulating effect of collagens on osteogenic cell growth was confirmed by the presence of mineralized nodules in the osteoblastogenic lineage cells and increased osteogenic stimulatory gene expression. The decreased BMM-derived TRAP+ osteoclasts number and osteoclastogenic regulatory gene expression of OC could demonstrate the exploitive osteoclastogenic activity of collagens. Osteoclastogenesis from BMM was triggered by paracrine cues of OC in some extend, but it was down-regulated by collagen. Overall, the effect of collagen on osteoclastogenesis and osteoblastogenesis may depend on the molecular weight of collagens, and collagen suppresses osteoclastogenesis, at least in part by downregulating the secretion of cytokines in OC.

Bone is connective tissue that continuously undergoes remodeling through the precise and localized coupling of resorption (removal of aged material) with replacement by newly formed bone. This process (bone homeostasis) requires regulated interactions between different cell types such as osteoblasts, osteoclasts (large plurinucleate cells), bone marrow macrophages and osteocytes. The failure of this system may help explain osteoporotic fractures in post-menopausal women and the elderly, a problem that has been increasing.

Osteoclast differentiation is triggered by cell-cell contact between osteoclast-precursor cells and other osteogenic progenitor cells in the bone including osteoblasts, osteocytes and bone marrow stromal cells1. The receptor activator of nuclear factor-kB ligands (RANKL) has been identified as the membrane-bound osteoclast differentiation factor of osteoclastogenesis-supporting cells that form stromal osteoclasts. Osteoblasts are believed to be the major cell type that expresses RANKL2. Nakashima et al.3 on the other hand stated that osteocytes have a greater capacity to support osteoclastogenesis in vitro than osteoblasts and bone marrow stromal cells. However, the major source of osteoclastogenesis remains unclear, as RANKL is expressed by several cell types in bone and

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bone marrow, including osteoblasts, osteocytes, mouse bone marrow mesenchymal stem cells (MMSC-bm) and lymphocytes. Therefore, it is important to know the actual cellular mechanisms of osteoclast formation to support work on bone remodeling.

Bone strength depends not only on the quantity of bone tissue but also on the quality, which is characterized by the geometry and the shape of bones, its mineral content and composition, and the fibrillar collagen microstructure. Collagen is a potential candidate for bone tissue engineering and may provide innate biological guidance to cells that favor cell attachment and promote chemotactic responses. As an extracellular matrix protein, collagen plays an important role in bone strength and remodeling. During bone homeostasis, osteoblast cells undergo mineralization mainly within the bone collagen to form mature bone cells (osteocytes). It was reported that administration of porcine skin collagen hydrolysates enhanced bone growth. In addition, oral administration of hydrolyzed type I collagen with calcitonin might inhibit bone resorption better than calcitonin alone. Guillermin et al. reported that specific collagen amino acid sequences such as Asp-Gly-Glu-Ala, Ala, Hyp-, Gly-Pro-Val and Gly-Pro-Hyp are involved in bone metabolism and bone growth. Unlike type I collagen, the osteogenic mechanism of type II collagen (CII) has not been fully evaluated. Our previous study claimed the excellent biocompatible properties of shark type II collagen. However, the influence of different molecular weight of type II collagens from shark cartilage with respect to the paracrine signaling mechanism of osteocytes for bone marrow macrophages derived osteoclast formation is needed to better understand for bone homeostasis and remodeling.

**Results and Discussion**

**Effect of collagen on osteocytes.** The result of proliferation assay is given in Fig. 1. On day 4, type II collagen (CII) and 57 kDa type II collagen polypeptide (57 kDa) treated groups (50 μg/ml) had a greater number of osteocytes than control, however, there were no significant differences observed between treated and control group on day 7. Similarly, rat tail type I collagen (150 μg/ml) coated osteocyte-like cell lines (MLO-Y4 cells) found to be an effective way to maintain the growth of osteocyte-like phenotype. In another study, primary osteocytes culture condition was optimized using type I collagen matrix with 0.2% FBS.

In general, osteocytes would not produce a high level of ALP, therefore, as an exploratory study, we estimated ALP activity to understand the regulatory mechanism of collagen and polypeptides in osteocytes. The results showed that the ALP level was elevated more in CII (50 μg/ml), 40 kDa type II collagen polypeptide (40 kDa) (1 μg/ml) and 57 kDa polypeptide treated groups than control on day 7 (p < 0.05) (Fig. 1). On day 4, there were no changes on ALP level in the treated groups. ALP activity is considered as an important factor in determining bone cell differentiation and mineralization and is used as a biochemical marker for determining mature bone cell phenotype. Therefore, an increased level of ALP in osteogenic cells would indicate increased osteocytes activity. As shown in Fig. 1, ALP activity was elevated with increasing concentrations of collagen and its polypeptides from 1 to 50 μg/ml compared to control. Similar to the present findings, ALP activity of murine osteoblasts was

![Figure 1. Proliferation effect (A) and ALP level (B) of pOC cells upon collagen and its polypeptides treatment.](image)
significantly increased in the presence of 1 mg/mL porcine collagen hydrolysates. Collectively, the above findings support an osteogenic capability of type-II collagens and also osteocytes might secrete ALP in some extent.

**mRNA expression.** mRNA levels of genes of interest in osteocytes treated with collagen and polypeptides are shown in Fig. 2. Col2a1 mRNA levels were not significantly altered by either collagen or polypeptides treatment, except high dose (50 μg/ml) of polypeptides at 1 h treatment (p < 0.05) and, low dose of CII (1 μg/ml) and 57 kDa (1 and 10 μg/ml) had increased the level of Col2a1 mRNA at 6 h. Il6ra mRNA levels were increased between 1 h and 6 h in untreated wells (p < 0.05). At 1 h and 6 h, Il6ra mRNA levels decreased with increasing CII and 57 kDa (50 μg/ml) (p < 0.05). It was proved that local stimulatory factors IL-6 and IL-6ra had predictive for osteoclast maturation, activation, and recruitment. Several studies demonstrated the higher level of IL-6ra responsible for bone loss in early post-menopause.

Tnfrsf11b mRNA levels were not significantly altered by either collagen or polypeptide treatment in 1 h. At 6 h, Tnfrsf11b mRNA levels were increased with increasing collagen (10 and 50 μg/ml) and 57 kDa (50 μg/ml) (p < 0.05). There was no significant difference in Tnfrsf11b mRNA levels between controls and collagen polypeptides (1 μg/ml and 10 μg/ml)-treated cells. Tnfrsf11b gene provides instruction for making a protein osteoprotegerin, which plays a vital role in bone remodelling and regulation of osteoclast cells. Tnfrsf11b acts as a decoy receptor for the receptor activator of nuclear factor kappa B ligand (RANKL), a major cytokine for osteoclastogenesis. Therefore, higher expression of Tnfrsf11b mRNA level by collagen and 57 kDa should reduce the osteoclast formation through osteocytes signaling mechanism.

Tumor necrosis factor ligand superfamily member 11 (Tnfsf11), also known as RANKL is a type-II membrane protein and has been identified to affect the immune system and control bone regeneration and remodeling. At 1 h and 6 h, Tnfsf11 mRNA levels were decreased with increasing collagen and polypeptides (57 kDa and 40 kDa) concentration (50 μg/ml) (p < 0.05). There is strong evidence that osteocytes produce paracrine signals and regulate osteoclast formation through RANKL production. The present study clearly depicted that treatment of collagen had obstructed the mRNA expression of RANKL in osteocytes.

Sost mRNA provides an instruction for producing a protein called sclerostin. It is produced in osteocytes and acts on the bone to stop bone formation. At 1 h, Sost mRNA levels were decreased with increasing collagen and
polypeptides concentration (50 μg/ml) when compared to control cells (p < 0.05). There was no significant difference in Sost mRNA levels by collagen and polypeptides in 6 h. From the data, collagen had down-regulate bone formation inhibitor gene, i.e., sost mRNA expression, which further supported the proliferation effect of collagen and polypeptides on osteocytes (Fig. 1). The recent study reported the decreased formation of the mineralized matrix and an increased mRNA expression of Sost in osteocytes, indicating a reduction in bone formation via the NF-κB pathway14.

Alpl is the gene for the alkaline phosphatase (ALP) enzyme, which helps in the growth and development of bones through bone mineralization. This enzyme helps for the deposition of calcium and phosphorus in developing bones15. Alpl mRNA levels were increased significantly between 1 h and 6 h in untreated cells (p < 0.05). Except for CII, 57 kDa and 40 kDa (10 μg/ml)-treated cells, there was no significant difference in Alpl mRNA levels between controls and collagen/polypeptides-treated cells in 1 h. At 6 h, Alpl mRNA levels decreased in collagen/polypeptides-treated cells (p < 0.05). Though, the proliferation rate and cellular alkaline phosphatase level of collagen treated osteocytes increased, the level of Alpl mRNA expression was not triggered at 6 h treatment.

Several hypotheses could be possible for the above result because in general, osteocytes are not the major sources for ALP productions for bone formation. The earlier study could demonstrate that an induction of Alpl mRNA during the in vitro osteogenic differentiation process expected for bone forming capacity of the bone marrow stromal cells in vivo16.

Taken together, these findings suggest that the expression of osteogenic regulatory genes such as Col2a1 and Alpl, was triggered initially at 1 h and declined later at 6 h in collagens treated osteocytes; and osteoclasts regulatory genes, such as Il6ra, and Tnfsf11 were suppressed in collagen and polypeptides treated osteocytes that could predict to down-regulate osteocyes cues for osteoclast formation. It was earlier confirmed by few authors that certain specific amino acid residues (asparagine, glutamine, glycine, and alanine) of collagen interact with integrin α2β1 on the mesenchymal stem cell membrane and can lead to up-regulate bone matrix synthesis through inhibition of TGF-beta signaling and activation of RUNX2 through FAK-JNK signaling17,18.

Effect of collagen on MMSC-bm. MMSC-bm activity contains three important phases including proliferation, matrix protein synthesis and mineralization of the bone matrix during new bone formation. The result of proliferation assay of MMSC-bm cells treated with different concentration of collagen and polypeptides is given in Fig. 3. In general, the proliferation rate of MMSC-bm cells was accelerated with increasing concentration of collagen and polypeptides. Notably, the rate was significantly higher on day 7 at 50 μg/ml of CII than other treatments. Similarly, Hennessy et al. and Gao et al. disclosed the osteogenic stimulatory activities of type I collagen
peptides and rat tail type I collagen on mesenchymal stem cell. In contrary, Song et al. observed decreasing trend of human bone marrow mesenchymal stem cells with coating and/or direct treatment of human type I collagen (5 μg/ml). Vleggeert-Lankamp et al. reported that bovine dermis type I collagen did not promote/inhibit the proliferation of human Schwann cells. In some studies, type I collagen was shown having the effect to increase cell proliferation, whereas, in some others, it did not have such effect, even decreased the proliferation of MSC cells. We speculated that this could be due to different molecular composition of collagens from different sources. Recently, Chiu et al. reported that the level of integrin α2β1 complex (VLA-2) expression on MSC-bm surface increased by type-II collagen treatment on day 4 and then gradually reduced from day 6 to a lower level than that of the control on day 10.

Collagens and polypeptides treated MMSC-bm cells cultured in osteogenic media were able to differentiate into the osteoblastogenic lineage cells. This was confirmed by the presence of mineralized nodules in collagens untreated cultures. There was no nodular aggregate deposited in negative control cultures (cells cultured without osteoblast supplements). This was clearly demonstrating that collagens and polypeptides accelerated osteoblast lineage cells differentiation from MMSC-bm and nodular aggregates present in collagens untreated cultures were calcium deposits. This is consistent with previous work showing that bone marrow-derived mesenchymal stem cells (BM-MSC) cultured on mammalian type-II collagen coated plates exhibited significant higher calcium deposition on day 12 and day 16, which suggests osteogenic induction properties of type-II collagen. Collagen treated cultures had larger alizarin red positive aggregates and stained more intensively, indicating that a more extensive calcium deposition had occurred. The presence of alizarin red stain with nodular cell aggregates observed in collagen and polypeptides treated cultures establishes that these amorphous deposits contain calcium and suggests that calcium deposits are made up of MMSC-bm cells differentiated to the osteoblastic lineage. Numerous possible mechanisms have been proposed for stimulating MMSC-bm osteogenic differentiation. Actual signaling mechanism of type II collagen during early osteogenic differentiation of MMSC-bm occurred by facilitating RUNX2 phosphorylation activation through integrin α2β1-FAK-JNK signaling axis, thus enhanced bone defect repair through an endochondral ossification-like process.

**Effect of collagen on osteoclast differentiation.** Macrophages harvested from mouse bone marrow were treated with RANKL and mCSF in the presence and absence of different concentration of collagen and polypeptides to assess the effects of the collagens on osteoclastogenesis. The result shows that increasing concentration of collagen and its polypeptides reduced osteoclast numbers considerably (p < 0.01) (Fig. 4). In contrast, the
murine oncostatin M (m-OSM) treated group increased osteoclast cell numbers (3+ nuclei) compared to control. But, the number of double nuclei osteoclast cells was decreased in m-OSM treated group than control (p < 0.01). The higher concentration of collagen and polypeptides treated groups had considerably decreased both double nuclei and 3+ nuclei TRAP+ cells compared to control (p < 0.01). Similarly, Guillermin et al. reported that murine osteoclasts differentiation reduced by the treatment of porcine collagen hydrolysates (2 or 5 kDa) with 1 mg/ml concentration, which might be due to inhibition of Transforming growth factor beta (TGF-β) through interaction of collagen-derived peptides (asparagine, glycine, glutamine and alanine).

The micrograph clearly depicted the lower number of osteoclast cells in the presence of higher concentration (50 μg/ml) of collagen and its polypeptides (Fig. 5). Among the treatment, 1 μg of 57 kDa collagen polypeptide-treated group had a high number of osteoclast cells than other treatment groups. In higher magnification (10×), “ghost” outlines remaining after osteoclast apoptosis cells were found in wells treated with the higher doses of CII but not in lower dose treatment. This observation indicated that increasing concentration of collagen increased apoptosis in osteoclasts.

**Co-culture model.** It was reported that differentiated primary osteoblasts express cytokines, RANKL and mCSF in presence of stimulators (IL-6, soluble IL-6 Receptor, prostaglandin E2 (PGE2), 1,25-Dihydroxy vitamin D3 (1,25 (OH)2D3) and oncostatin M), which bind with mouse BMM and support osteoclast differentiation15. In contrast, Nakashima et al. identified that purified osteocytes also express a much higher amount of RANKL and have a greater capacity to support osteoclastogenesis. Therefore, in the present study, we used a co-culture model with different stimulators to understand the effect of inducers such as m-OSM, IL-6, sIL-6R, 1,25 (OH)2D3 and PGE2 on osteoclast formation. The primary co-culture of BMM with pOC was grown in presence of inducers for 7 days. m-OSM significantly increased the number of single and double nuclei TRAP-positive cells, but few cells with more than 2 nuclei were observed. Compared to m-OSM, less number of TRAP-positive cells found in 1,25 (OH)2D3 and PGE2 group. In contrast, IL-6 in the presence of sIL6R supported only the production of TRAP-positive single and double nuclei cells to some extent (Fig. 6). No TRAP-positive cells were observed in wells treated with IL-6 or sIL-6R alone; this can also be seen in the micrographs (Fig. 6).
Our above study confirmed that osteocytes stimulated with m-OSM might support osteoclast differentiation to the point of a multinucleated cell, as previously observed with other stimuli in freshly isolated osteocytes. Further, we aimed to understand whether collagen and its polypeptide might play a similar role in the presence and absence of inducer, OSM in the co-culture of BMM and pOC. TRAP-positive single and double nuclei cells were formed in the collagens-OSM treated group (Fig. 7), but it was lower than m-OSM alone treated group (Fig. 6), which confirmed that collagen and collagen polypeptides might suppress m-OSM-stimulation of osteocytes towards osteoclast formation to some extent. In contrast, others reported no effect of hydrolyzed collagen on primary co-culture of osteoblast and osteoclasts growth. The down-regulation of osteoclastogenesis by collagen might be justified by the higher level of Tnfrsf11b mRNA (a decoy receptor for RANKL) that coupled with m-OSM-triggered-RANKL and leads to inhibition of RANK-RANKL ligand binding towards osteoclast formation.

The level of inhibition was higher (p < 0.05) in a collagen-OSM group than polypeptides treated groups, which might be due to the suppressive mechanism of osteoclastogenic regulatory genes such as Il6ra and Tnfsf11 expression of pOC by collagen (Fig. 2). Thus, pOC could not secrete necessary cytokines in order to stimulate osteoclast formation from collagen treated BMM-pOC co-culture, though the presence of positive inducer, m-OSM.

**Methods**

**Materials.** Type II collagen from whale shark cartilage was extracted with 0.5 M acetic acid containing 1% pepsin (Sigma-Aldrich Co., Ltd, Shanghai, China) and purified by gel-filtration column chromatography using a Sephadex G-100 (Sigma-Aldrich) column (25 × 3 cm). Collagen polypeptides were obtained by enzyme (thermolysin from Bacillus thermoproteolyticus rokko, EC number: 3.4.24.27, Sigma-Aldrich) hydrolysis of collagen. Briefly, 100 mg/mL collagen in 50 mM sodium phosphate buffer (pH 7.0) was hydrolyzed with thermolysin at 0.1% (w/w) of the protein content, 70 °C for 60 min and terminated with the addition of EDTA (20 mM) and two types of (57 and 40 kDa) collagen polypeptides were obtained using molecular weight cut-off filters (Millipore, Shanghai, China).

**Effect of collagen and collagen polypeptides on MMSC-bm and osteocytes.** Cell culture. Primary osteocytes (pOC) cells were harvested from mice (Sino-British Sippr/BK Lab Animal Co., Ltd, Shanghai, China) long bones (femora, and tibia) using 300 U/mL collagenase (Sigma-Aldrich) dissolved in α-minimal essential medium (α-MEM) (Gibco, Shanghai, China) and were cultured in α-MEM medium supplemented with 10% fetal bovine serum (Gibco) at 37 °C in a CO₂ incubator (Shanghai Hengyue Medical Instruments Co., Ltd, Shanghai, China), respectively. Animal study protocols and procedures were approved by the Shanghai Ocean University institutional animal care and use committee (Permit Number: 13-0012). All methods were employed in accordance with the relevant guidelines and regulations of Scientific and Ethical Care and Use of Laboratory Animals of Shanghai Ocean University.
Mouse bone marrow-mesenchymal stem cells (ZQ0465) (MMSC-bm) were purchased from the Shanghai Zhong Qiao Xin Zhou Biotechnology Co., Ltd, Shanghai, China and were cultured in mesenchymal stem cell culture medium (brand: Sciencell, Cat. No. 7501) containing 10% fetal bovine serum (FBS), (brand: Sciencell, Cat. No. 0025), mesenchymal stem cell growth supplement (1% MSCGS, Cat. No. 7552) and 1% penicillin/streptomycin (10,000 units/ml of Penicillin and 10,000 μg/ml of Streptomycin in a saline solution) (Cat. No. 0503) at 37 °C in a CO₂ incubator.

**Proliferation assay.** After confluence, cells pOC and MMSC-bm at passage 5 were seeded (5 × 10⁵ cells/well/48 well plate) in microtiter plates (Costar, Shanghai) along with the different concentration of collagen and polypeptides (1, 10 and 50 μg/ml). Controls consisted of uncoated (without collagen) wells. The total number of viable cells was counted using an Invitrogen cell counter (Countess II Automated Cell Counter, ThermoFisher Scientific, Shanghai, China) at 0, 4 and 7 days after seeding.

**Alkaline phosphatase (ALP) assay.** pOC cells were generated as described above. At each time point, cells were washed twice with 1 × phosphate buffered saline (PBS), harvested with lysis buffer (10 mM Tris buffer, pH 7.4) and sonicated (40 w, Shanghai Kedao Ultrasonic Instrument Co, Shanghai, China) with an ice water-bath for 30 sec to release cellular components. After a brief centrifugation (13,000 × g for 1 min at 4 °C, Universal 320 R, Andreas-Hettich, Buckinghamshire, Germany), the supernatant was used for the ALP assay. In brief, 25 μl per sample was added to 100 μl substrate (10 mM p-nitrophenyl phosphate (Sigma-Aldrich) in assay buffer (0.1 M Na₂CO₃ buffer, pH 10.0)). The reaction proceeded for 30 min at 37 °C; the reaction was terminated by addition of 50 μl NaOH (1 M) and absorbance measured at 410 nm using a plate reader (Bio-Rad Model 550, Shanghai). ALP activity was determined from a standard curve using 10 mM p-nitrophenol from 4000 to 62.5 nmoles and without substrate maintained as a control. The same volume of sample was used to determine protein content using bicinchoninic acid (BCA) as per the manufacturer’s instructions (Pierce, IL, USA). ALP activity was expressed as nmoles/min/mg protein determined as follows:

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\text{Alkaline Phosphatase level} = \frac{\text{ALP activity}}{\text{PC} \times \text{Time}}
\]

where, ALP level- Alkaline phosphatase level (nmoles/ml), PC- Protein concentration (μg/ml),Time-30 min

**mRNA expression.** pOC cells were seeded (250,000 cells/well) in 6 well microtiter plates (Costar) along with the different concentration of collagen and polypeptides (1, 10 and 50 μg/ml). Controls consisted of uncoated (without samples) wells. The percentage of mRNA expression (Sost, Alpl, Ilora, Tnfsf11, Tnfsf11b and Col2a1) was determined at a different time interval (0, 1 and 6 h) as follows.
and Molecular grade water to 20 μl FS master mix (Invitrogen, Shanghai, China), 3 μl cDNA synthesis.

Nanodrop ND-1000 spectrophotometer (ThermoFisher Scientific).

Briefly, cells were washed with PBS and fixed in 4% paraformaldehyde for 30 min and the effect of collagen on osteogenesis was confirmed by observing Ca deposition in cultured cells using the Alizarin red staining method. Cells were grown in the culture medium without sample and ObGS, respectively. The control and negative control cells were grown in the culture medium without sample and ObGS, respectively. The culture medium was removed from the cells by pipetting after an incubation period and washed twice with ice-cold PBS. The cell monolayer was lysed in TRizol (Invitrogen Life Technologies, Shanghai, China) (0.5 ml/well). The TRizol supernatant was passed through a 21 G needle using Luer-lock syringes (Zogear Industries Co., Ltd, Shanghai, China) to shear the RNA. Phase separation was obtained by addition of 0.2 ml chloroform for each 1 ml of TRizol used and shaken vigorously by hand until the sample was opaque. Then, cells were left on ice for a few min and spun at 12,000 × g for 15 min at 4 °C. The top clear phase containing RNA was collected and added to a fresh tube containing 0.5 ml isopropanol for RNA precipitation. The mixture was mixed thoroughly and centrifuged as previously. The pellet was washed with 0.5 ml 70% ethanol and re-centrifuged. The RNA pellet was suspended in an appropriate volume (10 μl) of RT-PCR water (Shanghai Biocolor BioScience & Technology Company, Shanghai, China). Dnase treatment was done using a Turbo Dnase kit (Shanghai Biocolor BioScience & Technology Company, Shanghai, China) as per the manufacturer's instructions. After centrifuging at 1000 × g for 2 min at 4 °C, RNA was quantified at an absorbance of 260 nm using a Nanodrop ND-1000 spectrophotometer (ThermoFisher Scientific).

cDNA synthesis. First strand cDNA synthesis was done in a 20 μl reaction mixture containing 1.0 μg RNA, 10 μl FS master mix (Invitrogen, Shanghai, China), 3 μl random primer (Invitrogen), 1 μl affinity script (Invitrogen) and Molecular grade water to 20 μl. The reaction mixture was incubated at 42 °C for 30 min and 85 °C for 5 min, followed by 5 min at 90 °C to inactivate the RT in a Biometra T3000 thermocycler (Biometra Gmbh, Gottingen, Germany).

Real-time polymerase chain reaction (RT-PCR) was done using a 96-well plate using an ABI 7500 Fast Real-Time PCR System (Applied Biosystems, Shanghai, China) using SYBR Green Fast qPCR Master Mix (Invitrogen, Shanghai, China). pOc regulatory primers designed using Primer-BLAST (Shanghai Biotechnology Co., Ltd, Shanghai, China) were listed in Table 1. The total volume of each PCR reaction was 10 μl, containing 5 μl SYBR Green Fast qPCR Master mix, 1.5 μl cDNA template sample, 0.5 μl of forward and reverse primers and 2.5 μl water. The PCR reaction was carried out at 95 °C for 30 min, 40 cycles at 95 °C for 5 min, 60 °C for 30 min, one cycle of 95 °C for 1 h, 55 °C for 30 min and 95 °C for 30 min.

Osteogenic differentiation and quantification. MMSC-bm cells at passage 5 were initially seeded at a density of 5 × 10^4 cells/well in microtiter 6 well plates. After 2 h, 50 μg/ml collagen or collagen polypeptides were added into each well. To induce osteogenesis, the cells were grown in osteoblast medium (Shanghai Zhong Qiao Xin Zhou Biotechnology Co., Ltd, Cat. No. 4601) with the addition of osteoblast growth supplement (ObGS) (Shanghai Zhong Qiao Xin Zhou Biotechnology Co., Ltd, Cat. No. 4652) composed of 100 nM dexamethasone, 10 mM β-glycerol phosphate, and 0.05 mM 2-phosphate-ascorbic acid for 21 days with media changes every 3 days. Control and negative control cells were grown in the culture medium without sample and ObGS, respectively. The effect of collagen on osteogenesis was confirmed by observing alizarin red staining method. Briefly, cells were washed with PBS and fixed in 4% paraformaldehyde for 30 min and stained with 1% alizarin red (Sinopharm Chemical Reagent Co., Ltd, Shanghai, China) containing 1% NH₄OH.

Effect of collagens on osteoclastogenesis. Bone marrow macrophage (BMM) preparation and RANKL-induced osteoclastogenesis. Mouse bone marrow macrophages (BMM) from 8 wk old wild mice (Sino-British) were cultured in a T75 medium flask (Costar) (20 ml final volume) containing 30 ng/ml mCSF (R&D Industries, Minneapolis, MN, USA) in endotoxin-free RPMI-10% heat-inactivated fetal bovine serum (FBS) (Gibco). The cells were cultured for 3 days and non-adherent BMM precursors were collected by centrifugation at 1,500 g for 4 min at 4 °C. The cells were re-suspended in αMEM medium (Gibco) and cell counts were adjusted to 50 × 10^5 BMM/ml. Then, they were seeded at 10^5 cells/well/0.2 ml medium in 48 well plates and treated with RANKL (100 ng/ml)/mCSF (30 ng/ml) (R&D Industries) (control group), mouse osteoclast M (m-OSM)-RANKL-mCSF (positive control), collagen and polypeptides (1, 10 and 50 μg/ml)-RANKL-mCSF.

| Primers | Sequence |
|---------|----------|
| Hprt1   | F'-TGATTAGCGATGATGAAACCAG<br> (housekeeping gene) R'-AGAGGGCCACAATGTGATG |
| Sost    | F'-GAGAACACACCACTATGAC<br> R'-GCTCGGCGCACTGTCATCT |
| Alpl    | F'-AAACCCCGACAAGCCTTC<br> R'-TCACACGGAGAAGGG |
| Trfsl1  | F'-TCCAGCTATGAGGAAGCT<br> R'-GTACCAAGGAGGAGG |
| Trfsl1b | F'-CCCTAATCAACAGCACTGC<br> R'-TAAGGCCCTTCCTCACACTC |
| Il6ra   | F'-TGCAACGCGGATCTGTGAGT<br> R'-CTGGACCTGCTTCCACACTC |
| Col2a1  | F'-ATCTGTAAGACCCAGACTGC<br> R'-GTTCTCCTTCTGCCCTTGG |

Table 1. List of primer sequence used in this study.
The plates were incubated at 37°C in a humidified atmosphere with 5% CO₂ for 7 days. Cells were fixed in 4% formaldehyde (Sinopharm) for 3 min followed by ethanol (1:1) for 1 min. After brief air dry, the cells were incubated with tartrate-resistant acid phosphatase (TRAP) stain (Sigma) for 8 min¹⁸, and the number of double and triple plus nucleated TRAP-positive osteoclasts were counted under a light microscope (Olympus Inc., Center Valley, PA, USA).

Co-culture model. BMM cultured as above were co-cultured with pOC (2 x 10⁴ cells/ml) in the presence of inducers such as oncostatin-M (m-OSM, 50 ng/ml), 10 nM 1α,25-Dihydroxyvitamin D₃ (1, 25 (OH)₂ D₃), 100 nM prostaglandin E₂, IL-6 (40 ng/ml) and soluble IL-6 receptor (100 ng/ml) (R&D Industries). BMM was also co-cultured with osteocytes in the presence of m-OSM (50 ng/ml) with or without collagen and polypeptides (50 μg/ml). From the previous results, m-OSM was selected as the potential inducer. Plates were incubated at 37°C in a humidified atmosphere with 5% CO₂ for 7 days, with a media change at day 3. Cells were then fixed with 4% paraformaldehyde (Sigma-Aldrich) for 10 min on ice, single and double plus nucleated osteoclasts were counted after TRAP stain.

Conclusion

In summary, the in-vitro results obtained with pOC, MMSC-bm and osteoclast cells demonstrated that type-II collagen and its polypeptides were able to stimulate osteogenesis and suppress osteoclastogenesis. The effect of collagen and its polypeptides in osteocytes and MMSC-bm cell growth is dose-dependent, which ultimately maintained the bone formation. Osteoclast numbers were reduced by the concentration of collagen and its polypeptides. Intriguingly, higher concentration of type-II collagen and polypeptides treatment had up-regulated the osteoblast lineage cell differentiation from MMSC-bm and had down-regulated the BMM-derived osteoclast cell formation, suggesting that collagen or collagen hydrolysate could be helpful in the management of bone diseases like osteoporosis. There are numerous reports disclosing the chondrogenic properties of type II collagen for cartilage repair, but here we are reporting that not only chondrogenesis, type II collagen might also play some role in ossification through modulating osteocytes paracrine signals for osteoclastogenesis. Therefore, the present work concluded that type II collagen and collagen polypeptides isolated from shark cartilages might be conceivably novel biomaterials to modulate bone formation and resorption activity; and could be of potential curiosity as a nutritional supplement in the preclusion of bone loss in osteoporosis. This work also highlighted that in some parameters native collagen was more efficient than that observed in collagen polypeptides. However, one important issue that needs to be addressed by further studies is the concern on the molecular interaction of collagen with osteocytes in order to maintain bone homeostasis.

Data Availability. The datasets generated and analyzed during the present study are available from the corresponding author upon reasonable request.

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Acknowledgements
This work received financial support from the National High Technology Research and Development Program of China (Project Grant No. 2011AA09070109), the National Natural Science Foundation of China (Project Grant No. 81750110548), an International Young Scientist Research Fellowship, and the Plan of Innovation Action in Shanghai (Project Grant No. 15410722500 and 17490742500).

Author Contributions
W.W.H. and B.B. designed the study, interpreted the data, and helped write the manuscript. J.E. conducted experiments, carried out laboratory-based analyses, analyzed data, prepared figures, and helped write the manuscript. S.W., R.G., and J.R., contributed to the interpretation of the data and preparation of figures. C.S., J.E.M.S.V., Y.H., C.W., and K.S.C.M.M., revised the manuscript. J.M.R. provided valuable advice on interpretation of data, revising figures and revised the manuscript. All authors reviewed the final manuscript and approved the final version.

Additional Information
Competing Interests: The authors declare no competing interests.

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