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***Basic Study***

**Transcriptomic alterations underline aging of osteogenic bone marrow stromal cells**

Cheng YH *et al.* Transcriptomic alterations underline BMSC aging

Yu-Hao Cheng, Shu-Fen Liu, Jing-Cheng Dong, Qin Bian

**Yu-Hao Cheng,** Institute for Cell Engineering, Johns Hopkins University School of Medicine, Baltimore, MD 21205, United States

**Shu-Fen Liu,** Institute of Spine, Shanghai University of Traditional Chinese Medicine, Shanghai 200030, China

**Jing-Cheng Dong, Qin Bian,** Department of Integrative Medicine, Huashan Hospital, Fudan University, Shanghai 200040, China

**Author contributions:** Liu SF and Bian Q performed the *in vivo* and *ex vivo* experiments, respectively; Cheng YH wrote the manuscript and analyzed the data; Dong JC and Bian Q edited the manuscript and provided feedback; Bian Q designed the project.

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**Corresponding author: Qin Bian, MD, PhD, Professor,** Department of Integrative Medicine, Huashan Hospital, Fudan University, Middle Wulumuqi Road, Shanghai 200040, China. bianqin213@126.com

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**Abstract**

BACKGROUND

Multipotent bone marrow stromal cells (BMSCs) are adult stem cells that form functional osteoblasts and play a critical role in bone remodeling. During aging, an increase in bone loss and reduction in structural integrity lead to osteoporosis and result in an increased risk of fracture. We examined age-dependent histological changes in murine vertebrae and uncovered that bone loss begins as early as the age of 1 mo.

AIM

To identify the functional alterations and transcriptomic dynamics of BMSCs during early bone loss.

METHODS

We collected BMSCs from mice at early to middle ages and compared their self-renewal and differentiation potential. Subsequently, we obtained the transcriptomic profiles of BMSCs at 1 mo, 3 mo, and 7 mo.

RESULTS

The colony-forming and osteogenic commitment capacity showed a comparable finding that decreased at the age of 1 mo. The transcriptomic analysis showed the enrichment of osteoblastic regulation genes at 1 mo and loss of osteogenic features at 3 mo. The BMSCs at 7 mo showed enrichment of adipogenic and DNA repair features. Moreover, we demonstrated that the WNT and MAPK signaling pathways were upregulated at 1 mo, followed by increased pro-inflammatory and apoptotic features.

CONCLUSION

Our study uncovered the cellular and molecular dynamics of bone aging in mice and demonstrated the contribution of BMSCs to the early stage of age-related bone loss.

**Key Words:** Bone marrow stromal cell; Mesenchymal stem cell; Mesenchymal stromal cell; Aging; Bone modeling and remodeling; Transcriptome

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**Core Tip:** Multipotent bone marrow stromal cells (BMSCs) are adult stem cells that form functional osteoblasts and play a critical role in bone remodeling. During aging, an increase in bone loss and reduction in structural integrity lead to osteoporosis and result in an increased risk of fracture. In this study, we examined age-dependent histological changes in murine vertebrae and uncovered that bone loss begins as early as the age of 1 mo. The BMSCs isolated at different ages revealed a consistent decreasing trend in both colony-forming and osteogenic commitment capacity. Moreover, we obtained the transcriptomic profiles of BMSCs at 1 mo, 3 mo, and 7 mo to investigate the distinct molecular and regulatory features that underpin the early loss of osteogenic potential. We showed the enrichment of osteoblastic regulation genes at 1 mo and loss of osteogenic features at 3 mo. The adipogenic and DNA repair features were enriched in the later age at 7 mo. Moreover, we demonstrated that the WNT and MAPK signaling pathways were upregulated at 1 mo, followed by increased pro-inflammatory and apoptotic features.

**INTRODUCTION**

Multipotent bone marrow stromal cells (BMSCs) are adult stem cells that can be isolated through plastic adherence and differentiate into three distinct lineages, including adipocytes, osteoblasts, and chondrocytes *in vitro*. BMSCs are responsible for constant renewal and generate functional osteoblasts *in vivo* that work with bone-resorbing osteoclasts to remodel bone structure[1]. It is well-known that an increase in bone loss with age leads to a more fragile skeletal integrity and causes osteoporosis[2,3]. In mice, the size of cortical bone, trabecular bone volume, and bone strength reach the peak at 3 mo followed by a constant decline[4,5]. In humans, bone mineral density peaks at the age of early 20 s and declines with age advancing[6,7]. Despite much scrutiny in age-dependent bone loss, the causal relationship between BMSCs and age-dependent bone loss is inferred mainly from the osteogenic features of BMSCs. There are limited studies about the role of multipotent BMSCs in bone aging. A study examined age-dependent changes in the number and lineage potential of multipotent BMSCs between the age of 3 mo and 24 mo and revealed that the self-renewal and osteogenic potential of BMSCs reached peaks at 3 mo and then decreased, which is consistent with the phenotypic features observed[8].

With the recent advances in microarray and sequencing technology, identification of the molecular features and dynamics of multipotent BMSCs in an unbiased manner has become possible. From the global transcriptome analysis, BMSCs showed an upregulation of genes that encode extracellular matrix components compared with other stem cells and mature cell types. Besides, BMSCs also present distinct transcriptomic signatures of mobility, proliferation, and oxidative stress response[9]. Aside from functional molecular features, the transcriptome comparison across time of human BMSCs identified 155 genes associated with BMSCs aging *in vitro*[10]. A recent study isolated two distinct BMSC populations, the CXCL12-abundant reticular cells and the platelet-derived growth factor receptor-a+Sca1+ cells. The comparison of transcriptomes between BMSCs isolated at 2 wk, 2 mo, and 2 years revealed the upregulation of pro-inflammatory gene expression that is associated with aging *in vivo*[11].

To identify the detailed dynamics of bone loss, we conducted a temporal observation of murine vertebrae at different time points, focusing on the early to middle age, and subsequently uncovered the molecular dynamics that underpin BMSC aging by global transcriptome analysis.

**MATERIALS AND METHODS**

***Animals***

Male imprinting control region mice (*n* = 10 per time point) were obtained from the Shanghai Laboratory Animal Center (SCXK 2007-0005, Science and Technology Commission of Shanghai Municipality). The study was approved by the Shanghai Animal Ethics Committee.

***Micro-computed tomography***

Lumbar spine specimens were fixed in 4% paraformaldehyde for 24 h, washed for 2 h, and examined. L4 vertebrae in each group were subjected to a 3D model without adnexa, the transverse, and the spinous processes. The images were captured using µCT 80 radiograph microtomography (Scanco Medical AG, Switzerland), and further processed with 3DCalc, cone reconstruction, and AVG model building software (HP, Japan). The bitmap data set was employed to reconstruct the 3D model. Scores for the bone mass density (BMD), the ratio of bone volume to tissue volume (BV/TV), the connectivity density of trabeculae (Conn.D.), the trabecular number (Tb.N), the trabecular thickness (Tb.Th), and the trabecular spaces (Tb.Sp) were measured from the 3D model.

***Histological and histomorphometric analyses***

Lumbar spines from the mice were fixed in 4% paraformaldehyde for 24 h, decalcified in 10% EDTA for 4 wk, and embedded in paraffin wax. The sections were stained with hematoxylin and eosin or underwent the following tartrate-resistant acid phosphatase staining (Sigma-Aldrich). The morphometric analysis was performed with an image auto-analysis system (Olympus BX50; Japan). The static parameters, including trabecular bone area (T.Ar) and the bone perimeter (B.Pm), were collected from the L4 lumbar spine and applied to compute the bone remodeling parameters, which include the osteoblast formation (N.ob/T.Ar, N.ob/B.pm) and the number of the osteoclast (N.oc/B.pm).

***BMSC culture and treatment and*** ***colony-forming unit-fibroblast assay***

BMSCs were obtained from the bone marrow aspiration of the bilateral tibia and femur. The cells were further subjected to either colony-forming assay, differentiation assay, or microarray detection. The marrow cavity was flushed with α-MEM (Gibco, United States) containing 10% fetal bovine serum (FBS, Gibco, United States) and 1% penicillin-streptomycin (Gibco, United States). The cells were cultured and expanded in 10 cm dishes (2 mice per dish) for 7 d for microarray detection. The number of colony-forming unit-fibroblasts (CFU-F) was counted under an inverted light microscope on the 3rd day of culture. For alkaline phosphatase (ALP) staining, the cells were fixed with 4% paraformaldehyde and stained with nitro blue tetrazolium and 5-bromo-4-chloro-3-indolyl phosphate (Pierce, United States) for 30 min.

***NimbleGen gene expression profiling***

The transcriptomic profiles were captured using NimbleGen gene expression profiling (No. PXH100525) containing 26991 genes (3 pieces per group, *n* = 9). Total RNA was isolated using TRIzol (Invitrogen) and the RNeasy kit (Qiagen) per the manufacturer’s instructions, including the DNase digestion step. After quality control *via* RNA measurement on the Nanodrop ND-1000 and denaturing gel electrophoresis, the samples were amplified and labeled using a NimbleGen one-color DNA labeling kit and hybridized in the NimbleGen hybridization system. Subsequently, the chip underwent steps of washing and scanned with the Axon GenePix 4000B microarray scanner. Raw data were obtained with NimbleScan software (version 2.5). NimbleScan software’s implementation of RMA offers quantile normalization and background correction. The gene summary files were imported into Agilent GeneSpring software (version11.0) for further analysis. Differentially expressed genes were identified through fold-change and *t*-test screening and visualized *via* the heatmap function of the Seurat package in R[12].

***Gene set enrichment analysis***

We used the R package enrichR to complete the gene set enrichment analysis (GSEA). The top enriched markers of each group were extracted and subjected to enrichR function. The mouse cell atlas database was used to characterize the cell typing features of samples, and the Wikipathway 2019 Mouse database was used to analyze the most enriched pathways features.

***Statistical analysis***

The data are expressed as the mean ± SE, and statistical significance was calculated using one-way ANOVA followed by the Tukey’s test (heterogeneity of variance) using aov and TukeyHSD function in R. The significance level was defined at *P* < 0.05.

**RESULTS**

We applied µCT imaging to quantitatively measure the densitometry and capture the structural dynamics in lumbar spines at multiple time points. We combined the imaging findings with histological features to quantitatively analyze the dynamic changes of osteoblasts and osteoclasts in bone tissue. Subsequently, we isolated BMSCs and characterized their self-renewal and osteogenic potential to support the imaging and histological findings. Finally, we identified the molecular signatures that underpin the bone loss by obtaining the gene expression profile near the initiation time point when BMSCs lose self-renewal and differentiation potential.

***Skeletal remodeling dynamics during aging***

We obtained the µCT three-dimensional images of the L4 lumbar spine from mice at the age of 1 mo, 3 mo, 7 mo, 12 mo, 15 mo, and 18 mo. The loss of bone volume and decrease in connectedness in the cancellous compartment with age were apparent (Figure 1A). The quantitative analysis revealed the changes in the cancellous bone integrity with age. The BMD continuously increased and plateaued at 15 mo with a 45% increase, and later remained constant. The BV/TV varied across the time and peaked at 12 mo with a 75% increase, followed by a dramatic decrease at 18 mo. For trabecular bone, the Tb.N decreased over time, but the difference between 7 and 18 mo was not significant. Between 1 and 18 mo, Tb.N decreased by 36%. The Tb.Th continuously increased and reached the plateau at 12 mo. The Tb.Sp increased from 1 to 18 mo by 60%. The Conn.D. decreased from 1 to 12 mo, while beyond the age of 12 mo, the Conn.D. remained constant. The degree of anisotropy in trabecular bone orientation stayed the same with age (Figure 1B). The finding that the trabeculae number decreased along with the increase in BMD with age indicated the enhancement of mineral deposit and active bone remodeling. The decrease in cancellous bone volume and trabecular thickness from 12 mo suggested that bone remodeling reached the equilibrium of formation and resorption at 12 mo and later skewed to the resorption.

***Bone remodeling and BMSC osteogenic capacity during aging***

To investigate the cellular composition with age, we performed histological analysis on the L4 lumbar trabecular bone at 1 mo, 3 mo, 7 mo, 12 mo, 15 mo, and 18 mo. In the aged 18-mo-old mice, we observed bone structure reduction and a lower density of osteoblasts. Across the age followed, the number of osteoblasts per trabecular area (T.Ar) maintained at a relatively high level until 3 mo, followed by a drop, and then remained constant with advancing age. The difference was similar to the results normalized by bone perimeter (B.pm). On the bone resorption side, the osteoclast number per vertebra peaked at 1 mo with a sharp drop and gradually returned to a similar level at the age of 18 mo (Figure 2A).

To determine the self-renewal and differentiation features of BMSCs, we flushed bone marrow cells from 1 mo, 3 mo, 7 mo, 12 mo, 15 mo, and 18-mo-old mice and performed CFU-F and ALP staining assays. The number of colonies decreased at 3 mo and began to increase from the age of 12 mo. The self-renewal capacity reached a peak at 15 mo, followed by a sharp decrease at 18 mo. To study whether aging alters the osteogenic potential of BMSCs, we examined the ALP activity *via* staining. The results showed that ALP activity reached two peaks at 1 and 15 mo, which represented the modeling and the remodeling phase during development and aging (Figure 2B). The results suggested that the self-renewal and differentiation potential of BMSCs dropped during the modeling phase, followed by an increase at the middle age between 12 and 15 mo. After that, the capacity again dropped sharply at 18 mo.

***Transcriptome profile of BMSCs at modeling phase***

We collected the transcriptome profiles of BMSCs *via* microarray at the age of 1 mo, 3 mo, and 7 mo to uncover the molecular regulation contributing to the drop of self-renewal and differentiation potential observed in 1-mo-old mice. The feature signatures at 1 mo included genes associated with mesenchymal migration, such as *Coro1c*, and telomere regulation, such as *Terf1*. Some critical osteoblastic regulation genes, such as *Tnpo1, Dock7,* and *Apoa2* were also enriched at an early age. Multiple Hox genes were relatively enriched at the age of 1 mo, suggesting the involvement of patterning of the bone tissue. At 3 mo, the BMSCs did not show the osteogenic regulation and distinct aging-preventing features that enriched in 1 mo but showed enrichment of chondrogenic regulators Sox9 and Snai1. BMSCs also gained the osteoblastic inhibition and osteoclast promoting feature gene *Efna1*. The BMSCs at 7 mo showed enrichment of the bone remodeling regulators Hey1 and Hey2. We also observed enhanced expression of DNA repair gene *Neil3* (Figure 3A).

Subsequently, we subjected the differential genes to the GSEA analysis. Using the cell atlas database, we uncovered that 1-mo and 7-mo samples were relatively enriched in osteoblasts and bone features, but lost of the osteogenic features at 3 mo (Figure 3B). When we applied the pathway analysis database as a reference, we uncovered the underlying signaling pathways that were involved in gaining distinct features. At 1 mo, the MAPK and WNT signaling pathways were highly enriched. The 3-mo sample presented pathways related to apoptosis, stress features, and more pro-inflammatory signaling pathways, including IL-1 and IFNg signaling pathways. At the age of 7 mo, the pathway related to adipogenesis began to appear, and also present features of DNA repair. High enrichment of both ossification and matrix metalloproteinase resorption pathways suggested active bone remodeling (Figure 3C).

**DISCUSSION**

Bone loss is a specific feature during aging that is partially caused by the impaired osteogenic capacity of BMSCs. In this study, we characterized bone and BMSC features in mice at different ages and uncovered the underlying transcriptomic changes. The µCT results revealed the dynamic alteration in murine vertebrae during aging. The finding was similar to the previous study that observed the age-related changes in the long bone tibia[13]. Although both vertebrae and long bone showed a constantly decreased trabecular number, the bone volume and trabecular thickness in vertebrae fluctuated, which increased at an early age followed by constant downward trend. The pattern of changes in mouse vertebrae was also similar to that observed in humans. In both species, the BV/TV and Tb.N in lumbar vertebrae increased at an early age during growth and development followed by a downward trend with age[14,15]. However, there was a difference between the trabecular bone of humans and mice. In humans, Tb.Th presents finer alterations that showed both thinner trabecula and thickening remaining trabecula, which did lead to overall significant changes with age[16]. In mice, the thinness of trabecular bones was significantly increased at an early age and remained constant after the age of 7 mo. The temporal changes in bone with age were surprisingly similar between mice and humans. The total bone mass peaks at midlife around 30 and 40 years of age in humans while the bone mass in mice also peaks at the middle age of 15 mo[17].

The cellular components in the bone remodeling of mouse vertebrae changed with age. The decreasing density of bone-forming osteoblasts and the increasing number of osteoclasts were consistent with the previous reports that the activity of bone formation decreased while bone resorption increased with age[18,19]. However, we noticed a peak in the number of osteoclasts at an early age, which suggested the early activation of bone modeling. Unlike osteoblasts, the stem and progenitor populations BMSCs that are responsible for bone repopulation showed a different pattern of changes with age. Considering both self-renewal colony-forming assay and osteogenic assessment, a similar peak presented at the early life at 1 mo, which was consistent with the high osteoblast density as we observed; however, we observed a significant increase in the amount of colony formation with high osteoblastic features at 15 mo. The previous study assessed the number and differentiation potential of BMSCs between the ages of 3 mo and 18 mo also reported a similar pattern[8]. The discrepancy between a high osteogenic capacity and a low number of osteoblasts could be related to the high bone remodeling at the age of 15 mo and exertion of the BMSC populations. However, the other study isolated BMSCs from rats of 1 mo and 16 mo of age did not uncover a similar pattern. The BMSCs from the 16-mo-old rats had significantly decreased colony number, size, and ALP expression compared with the 3-wk-old rats[20]. The difference suggested that the transient recovering capacity of BMSCs might vary across species and require a finer follow-up interval to uncover.

We obtained the transcriptomic profile of BMSCs at the early time points to investigate the underlying molecular regulation that underpins the early decrease in BMSC capacity. At the age of 1 mo, the BMSCs showed multiple feature genes related to osteogenic regulation, including *Tnpo1, Dock7,* and *Apoa2*. A previous study showed that knockdown of *Tnpo1* *via* siRNA abrogated the osteoblast differentiation of BMSCs[21]. The other study reported that Dock7 was responsible for trabecular maintenance. Loss of Dock7 in the mice resulted in an impairment of periosteal and endocortical envelope expansion and lower trabecular bone mass[22]. A recent study uncovered that *Apoa2* knockout mice derived BMSCs had a less osteogenic commitment and an increased tendency to form adipocytes[23]. Aside from osteogenic regulation, the top differential genes also uncovered general features that were distinct in early life. *Terf1* is the gene that was correlated to telomere maintenance in stem cell populations and was highly associated with aging[24,25]. *Terf1* was also highly enriched at the age of 1 mo compared with the other two time points. The finding was consistent with the fact that the sample was the youngest among the others. Unlike the osteogenic related features, the 3-mo data revealed more differential genes that are related to chondrogenic commitment including *Snai1* and *Sox9*. Sox9 is a well-known master regulator that governs chondrogenic commitment[26]. A previous study of *Snai1* knockout showed a substantial defect in the long bones[27]. The chondrogenic features along with the endochondral ossification features in the following time point suggested a temporal progression of bone growth and development. The BMSCs of the later age of 7 mo revealed features of aging and bone resorption. *Neil3*, one of the top differential genes, is involved in DNA repair, which is one of the critical factors related to genome stability and cellular aging[28,29]. The other feature gene *Efna1* was reported to promote osteoclastogenesis and inhibit osteoblast formation, suggesting the tendency of enhanced bone resorption[30,31].

The following GSEA of the differential genes allowed us to have a broader picture of the feature genes at different ages. When we applied the cell atlas database to be the reference, BMSCs from mice aged 1 mo or 7 mo revealed the enrichment of osteoblast features, while BMSCs from mice at the age of 3 mo showed a more diverse cell type feature. The loss of the osteogenic features at 3 mo was temporally consistent with our findings of a drastic decrease in the osteogenic capacity of BMSCs. The GSEA using the signaling-pathway database as a reference also supported the findings. The pathways involved at 1 mo were largely related to osteogenic regulation, including WNT and MAPK pathways[32,33]. At 3 mo, the signaling pathways revealed more pro-inflammatory features, including IL-1 and IFNg signaling pathways. The most significant apoptotic pathways suggested the bone loss as we observed at an early stage, which was probably caused by the activation of apoptosis in BMSCs, and ultimately contributed to the remodeling of the bone architecture. The pathways involved in BMSCs at the later age of 7 mo were related to DNA mismatch repair and bone remodeling, which included ossification and MMP enzymes. The finding was consistent with the top enriched genes discovered at a later age.

**CONCLUSION**

In summary, our study showed the histological and cellular dynamics of bone aging in mice and demonstrated the temporal changes of the osteogenic BMSCs. Moreover, we uncovered the temporal features *via* transcriptomic analysis, which suggested the contribution of BMSCs to the early stage of age-related bone loss.

**ARTICLE HIGHLIGHTS**

***Research background***

Multipotent bone marrow stromal cells (BMSCs) form functional osteoblasts and are involved in bone formation. During aging, significant bone loss leads to osteoporosis and results in an increased risk of fracture.

***Research motivation***

We discovered that an early bone loss occurs as early as 1 mo in mice, and we would like to investigate the role of BMSCs during early bone loss.

***Research objectives***

To understand the functional alterations of BMSCs during the early bone loss and uncover the transcriptomic dynamics that underpin the early loss of osteogenic potential.

***Research methods***

We collected BMSCs from mice at early to middle ages and assessed their self-renewal and differentiation potential. Subsequently, we obtained the transcriptomic profiles at a young age to reveal the features of BMSCs during early bone loss.

***Research results***

The colony-forming and osteogenic commitment capacity decreased at the age of 1 mo. At 3 mo, BMSCs were enriched in osteoblastic regulation genes, and at 7 mo, the transcriptomic features shifted toward adipogenic and DNA repair. The gene set enrichment analysis suggested the involvement of WNT and MAPK signaling pathways at the osteogenic phase and increased pro-inflammatory and apoptotic features at the latter phase.

***Research conclusions***

We demonstrated the contribution of BMSCs to the early stage of age-related bone loss and uncovered the underlying transcriptomic dynamics.

***Research perspectives***

Resolving the detailed cellular and molecular mechanism underlying bone aging is crucial. In this study, we demonstrated the role of BMSCs in early bone loss and revealed the transcriptomic dynamics to better understand the underlying molecular mechanism.

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**Footnotes**

**Institutional animal care and use committee statement:** The study was approved by the Shanghai Animal Ethics Committee.

**Conflict-of-interest statement:** Qin Bian is currently a research associate at Johns Hopkins University. The work has been done prior to her entrance to Johns Hopkins University.

**Data sharing statement**: No additional data are available.

**ARRIVE guidelines statement:** The authors have read the ARRIVE guidelines, and the manuscript was prepared and revised according to the ARRIVE guidelines.

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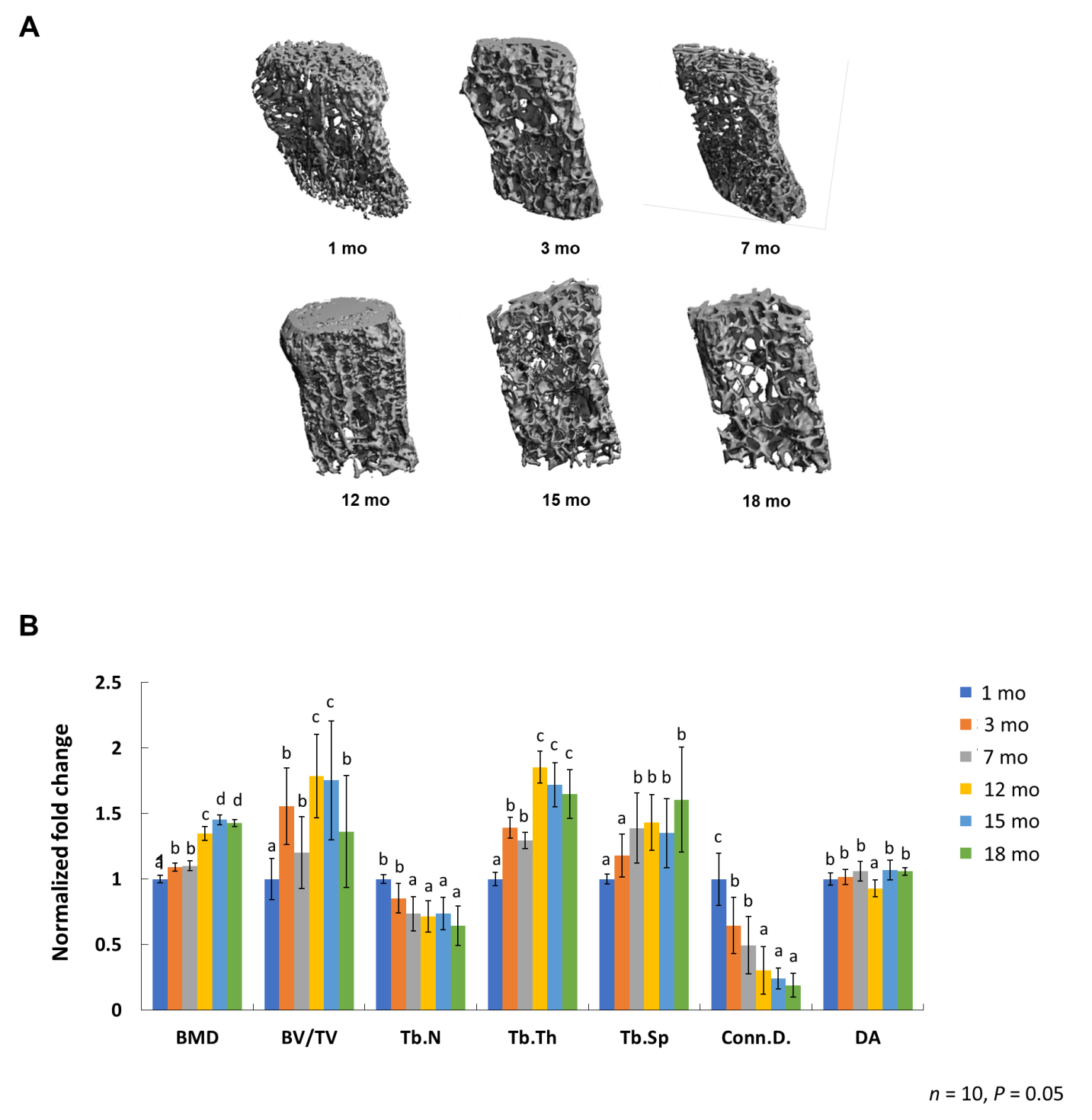
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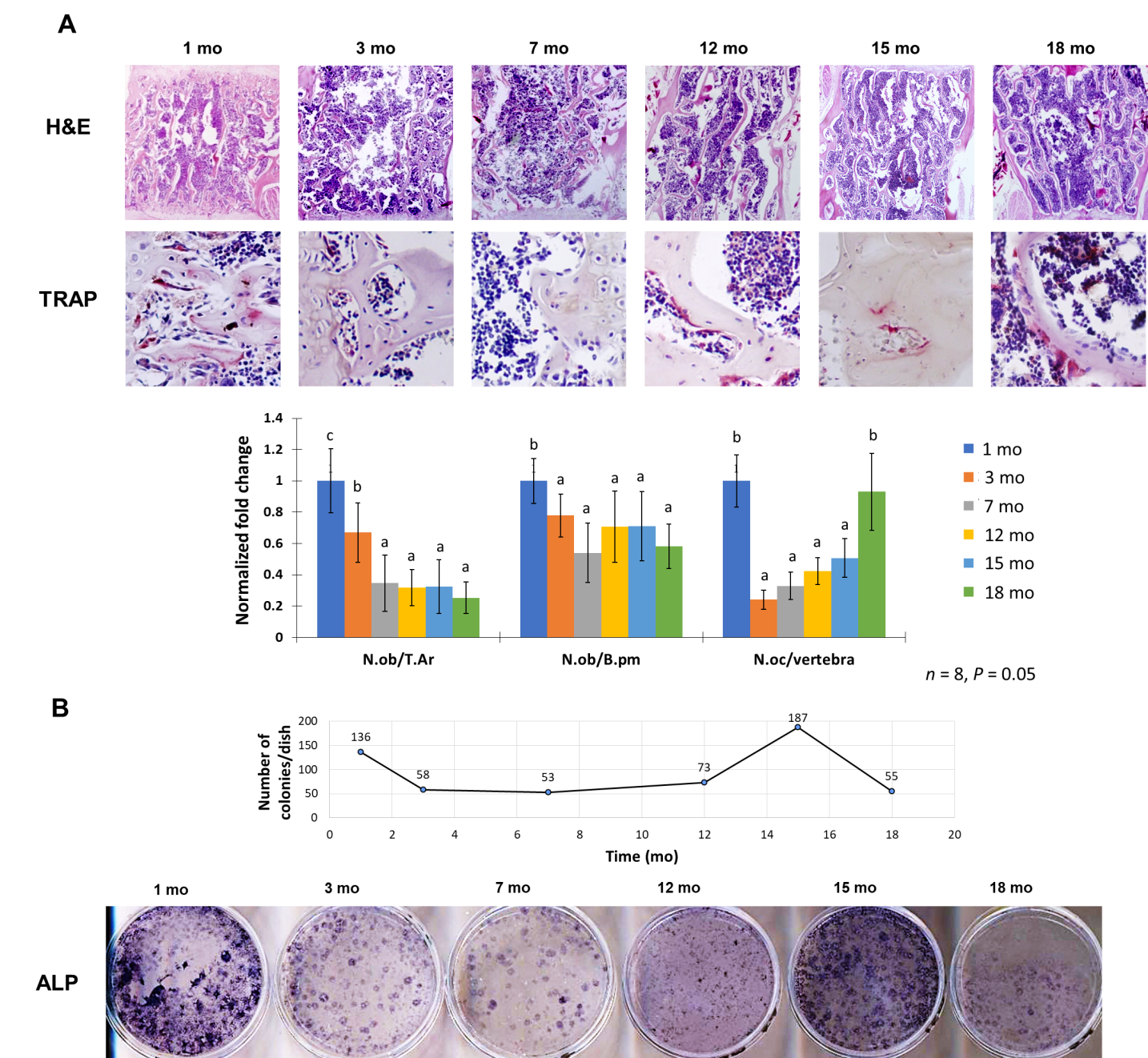
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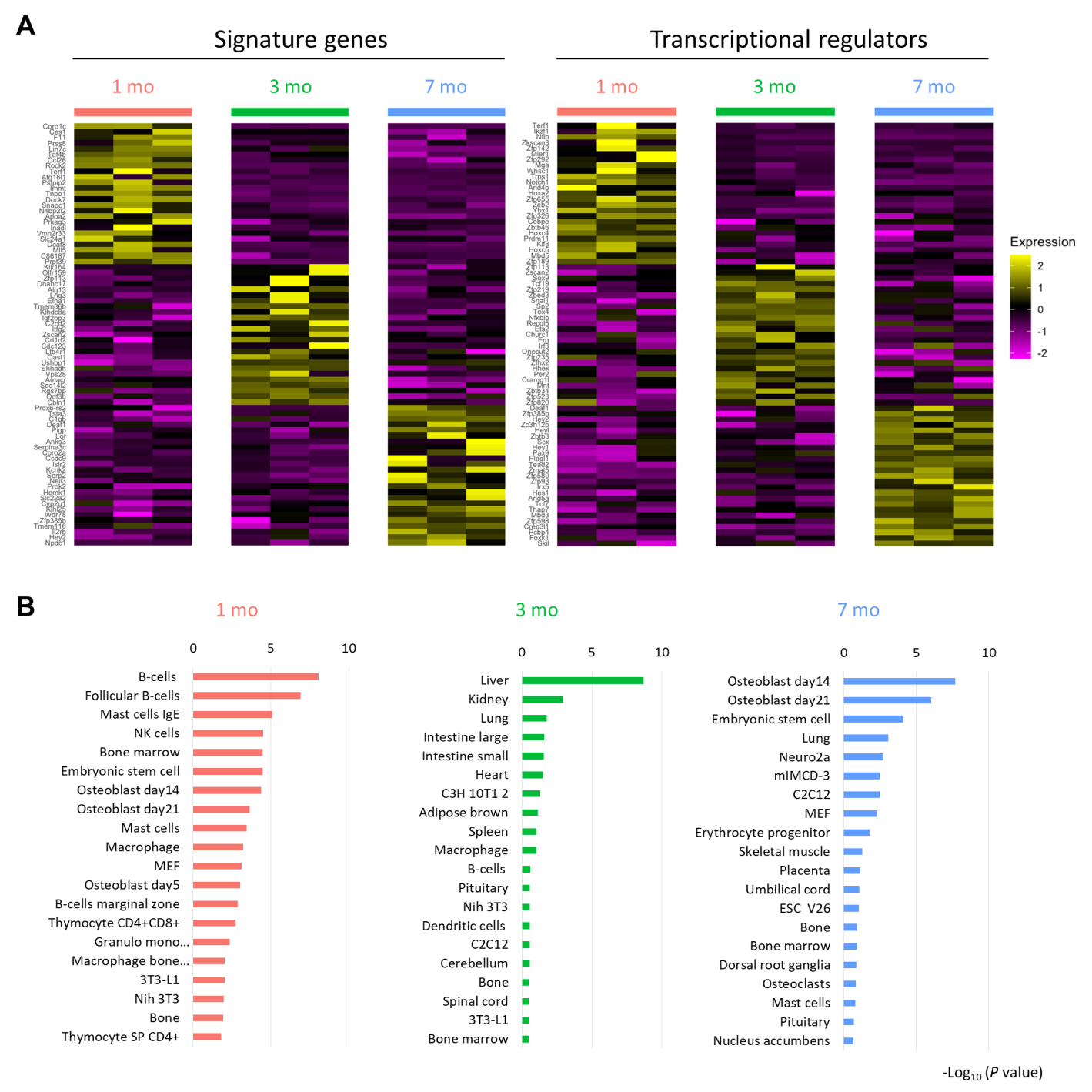
**Figure Legends**

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**Figure 1 Quantitative measurement of skeletal features.** A: Micro-computed tomography images of L4 lumbar spine at 1 mo, 3 mo, 7 mo, 12 mo, 15 mo, and 18 mo; B: Quantitative measurement of the densitometry and structural parameters of cancellous bone, including the ratio of bone volume to tissue volume (BV/TV), the connectivity density of trabeculae (Conn.D.), the trabecular number (Tb.N), the trabecular thickness (Tb.Th), and the trabecular spaces (Tb.Sp). Bone mass density, Tb.Th, and Tb.Sp increased with age while Tb.N and Conn.D. decreased constantly. The BV/TV increased and reached the plateau at the age of 12 mo. The degree of anisotropy did not change over time between the window of 1 mo and 18 mo. BMD: Bone mass density; BV/TV: Bone volume to tissue volume; Tb.N: Trabecular number; Tb.Th: Trabecular thickness; Tb.Sp: Trabecular spaces; Conn.D.: Connectivity density of trabeculae; DA: Degree of anisotropy.

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**Figure 2 Histological assessment of bone properties and osteogenic bone marrow stromal cells during bone aging.** A: Mouse lumbar spine at different ages was sectioned and stained with hematoxylin and eosin for osteoblast detection, or underwent tartrate-resistant acid phosphatase staining for quantifying the number of osteoclasts. The quantitative analysis measured the osteoblast number per trabecular surface area or perimeter, and osteoclast number per vertebra; B: Self-renewal and differentiation capacity of bone marrow stromal cells assessed *via* colony-forming assay and alkaline phosphatase staining. Two peaks at 1 mo and 15 mo were detected, suggested the modeling and remodeling phase during development and aging. H&E: Hematoxylin and eosin; TRAP: Tartrate-resistant acid phosphatase; ALP: Alkaline phosphatase; N.ob: Number of osteoblasts; T.Ar: Trabecular bone area; B.pm: Bone perimeter; N.oc: Number of the osteoclast.





**Figure 3 Transcriptomic profile of bone marrow stromal cells in young mice.** A: The heatmap demonstrated the signature genes and the transcription regulators differentially expressed at different ages that were computed from the microarray data; B: The gene set enrichment analysis taking mouse cell atlas as reference revealed the fate tendency of bone marrow stromal cells (BMSCs); C: The signaling pathway analysis uncovered the associated signaling pathways enriched at different ages in BMSCs.



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