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Authors

Chan, Charles KF
Gulati, Gunsagar S
Sinha, Rahul
et al.

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Identification of the Human Skeletal Stem Cell

Charles K. F. Chan^{#a,b,*,‡}, Gunsagar S. Gulati^{#a,b}, Rahul Sinha^{#b}, Justin Vincent Tompkins^{#a}, Michael Lopez^{#a}, Ava C. Carter^{#c}, Ryan C. Ransom^{#a,b}, Andreas Reinisch^{#b}, Taylor Wearda^{#a}, Matthew Murphy^{#a}, Rachel E. Brewer^a, Lauren S. Koepke^a, Owen Marcic^a, Anoop Manjunath^{a,b}, Eun Young Seo^a, Tripp Leavitt^a, Wan-Jin Lu^b, Allison Nguyen^b, Stephanie D. Conley^b, Ankit Salhotra^a, Thomas H. Ambrosi^a, Mimi R. Borrelli^a, Taylor Siebel^b, Karen Chan^b, Katharina Schallmoser^d, Jun Seita^e, Debashis Sahoo^f, Henry Goodnough^g, Julius Bishop^g, Michael Gardner^g, Ravindra Majeti^h, Derrick C. Wan^a, Stuart Goodman[#], Irving L. Weissman^{#b}, Howard Y. Chang^{#c}, and Michael T. Longaker^{#a,b,*}

^aDepartment of Surgery, Stanford Medicine, CA 94305, USA

^bInstitute for Stem Cell Biology and Regenerative Medicine, Stanford Medicine, CA 94305, USA

^cCenter for Personal Dynamic Regulomes, Stanford University, CA 94305, USA

^dDepartment of Blood Group Serology and Transfusion Medicine, Medical University of Graz, Austria

^eMedical Sciences Innovation Hub Program, RIKEN, Tokyo, 103-0027, Japan

^fDepartments of Pediatrics and Computer Science and Engineering, University of California San Diego, CA 92093, USA

^gDepartment of Orthopedic Surgery, Stanford Medicine, CA 94305, USA

^hDepartment of Medicine, Division of Hematology, Stanford Medicine, CA 94305, USA

These authors contributed equally to this work.

Summary

Stem cell regulation and hierarchical organization of human skeletal progenitors remain largely unexplored. Here, we report the isolation of a self-renewing and multipotent human skeletal stem cell (hSSC) that generates progenitors of bone, cartilage, and stroma, but not fat. Self-renewing and multipotent hSSCs are present in fetal and adult bones and can also be derived from BMP2-treated human adipose stroma (B-HAS) and induced pluripotent stem cells (iPSCs). Gene expression analysis of individual hSSCs reveals overall similarity between hSSCs obtained from different sources and partially explains skewed differentiation towards cartilage in fetal and iPSC-

[‡] Lead contact for material and resource requests: chazchan@stanford.edu. *Correspondence: chazchan@stanford.edu (C.K.F.C.), longaker@stanford.edu (M.T.L.).

Author contributions

C.K.F.C., G.S.G., R.S., J.V.T., designed and performed experiments, and wrote the manuscript. M.L., A.C.C., R.C.R., A.R., T.W., M.M., R.E.B., L.S.K., O.M., A.M., E.Y.S., T.L., W.J.L., A.N., S.D.C., A.S., T.H.A., M.R.B., T.S., K.C., H.G., K.S., J.S., D.S. performed experiments, J.B., M.G., D.W., S.G., provided clinical skeletal specimens and edited the manuscript. R.M., I.L.W., H.Y.C., M.T.L. supervised the project.

Declaration of Interests

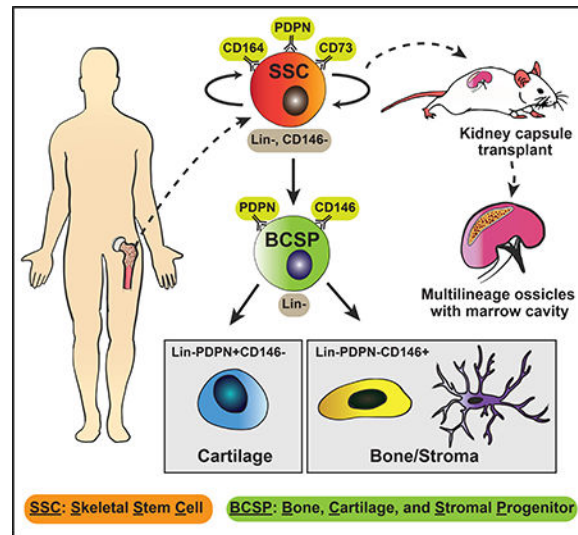
Patent is pending for the isolation, derivation, and use of human skeletal stem cells and downstream skeletal progenitors.

derived hSSCs. hSSCs undergo local expansion in response to acute skeletal injury. In addition, hSSC-derived stroma can maintain human hematopoietic stem cells (hHSCs) in serum-free culture conditions. Finally, we combine gene expression and epigenetic data of mouse skeletal stem cells (mSSCs) and hSSCs to identify evolutionarily conserved and divergent pathways driving SSC-mediated skeletogenesis.

In Brief

Identification of a human skeletal stem cell reveals conserved and species-specific pathways in skeletal development and response to injury and will guide future regenerative approaches.

Graphic Abstract:



Introduction

The human skeleton is a composite of diverse tissue types including bone, cartilage, fat, fibroblasts, nerves, blood vessels, and hematopoietic cells. Each tissue type in the skeleton is generated and maintained by the precise regulation of distinct stem cells that possess the ability to replenish themselves and give rise to multiple, nonoverlapping cell-lineages. Among the most well-characterized stem cells in the skeleton is the hematopoietic stem cell (HSC), marked by a combination of cell-surface proteins and defined by its functional ability to form colonies in vitro and serially reconstitute long-term hematopoiesis in conditioned hosts (Baum et al., 1992; Spangrude et al., 1988). Similarly, a significant amount of effort has been devoted towards identifying the cellular origins of the non-hematopoietic lineages in skeletal tissue, such as bone, cartilage, vascular endothelium, and stroma (Chan et al., 2015a; Marecic et al., 2015; Mendez-Ferrer et al., 2010; Worthley et al., 2015; Zhou et al., 2014). However, the absence of reliable cell-surface markers, tissue dissociation protocols, and functional stemness assays has hampered the discovery of putative stem cells in the human skeleton.

Early efforts to isolate non-hematopoietic stem cells in the bone relied on the ability of bone marrow cells to adhere to plastic plates. However, these cultures of “mesenchymal stem cells” contain heterogeneous mixtures of cells with indeterminate potencies and promiscuous contribution to many overlapping lineages, such as bone, cartilage, fat, muscle, fibroblast, endothelial cells, and stroma. Likely, these cells represent a population comprised of multiple types of distinct stem cells rather than a uniform purified skeletal stem cell (SSC). Identification of several specific cell-surface proteins has improved the enrichment for skeletogenic activity in mesenchymal cultures (Lv et al., 2014). THY1 (CD90), ENG (CD105), NT5E (CD73), and CD44 broadly label bone marrow stromal cells with variable colony-formation ability and differentiation to osteo-, chondro-, and adipo-lineages (Lange et al., 2005). NGFR (CD271) and MCAM (CD146) additionally select for cells with higher colony-formation ability and multilineage differentiation (Quirici et al., 2002; Sorrentino et al., 2008). CD146+ cells can also recreate a hematopoiesis-supportive human ossicle in a mouse (Sacchetti et al., 2007). Similarly, STRO-1 and STRO-3 are expressed in cells capable of supporting human hematopoietic stem cells for an extended time in the presence of serum (Bensidhoum et al., 2004; Simmons and Torok-Storb, 1991). Still, cells expressing these markers are heterogeneous in their function, and the true nature of a functional human skeletal stem cell (hSSC) hierarchy in the bone marrow is missing.

Our and others’ evidence from lineage-tracing and clonal analysis in mice supports the existence of mouse SSCs (mSSCs) that exclusively give rise to bone, cartilage, and stroma, but not fat, muscle, fibroblasts, or hematopoietic tissue (Chan et al., 2015a; Marecic et al., 2015; Worthley et al., 2015). We hypothesized that similar SSCs may exist in humans. In this study, we report the isolation of a human SSC (hSSC) capable of self-renewal and multilineage differentiation to bone, cartilage, and stroma. We demonstrate that these cells can be isolated from various human tissue sources and quantify their response to fracture. Cross-species comparison of hSSCs with our mSSCs additionally provides insights into convergent and divergent skeletal evolution.

Results

Single-cell transcriptome analysis of growth plate and diaphysis reveals markers for hSSC isolation

As described above, our findings in mice (Chan et al., 2015a; Marecic et al., 2015) suggested that similar types of lineally related skeletal stem and progenitor cells may also be prospectively isolated by fluorescence-activated cell sorting (FACS) from human skeletal tissue. We reported a high frequency of clonal mSSC-derived cell expansion in the growth plate of the mouse bone by lineage-tracing using Actin-Cre^{ER} x Rainbow mice (Chan et al., 2015a). Therefore, we performed single-cell RNA-sequencing (scRNA-seq) of the growth plate and diaphysis of a 17-week human fetal femur to identify individual cells with the greatest homology in terms of gene expression to the mSSC and the mouse bone, cartilage, stromal progenitor (mBCSP), which is immediately downstream of the mSSCs in the differentiation hierarchy (Figure 1, Figure 4 right panel) (Chan et al., 2013; Chan et al., 2015a; Fleming et al., 2015). We first microdissected the fetal bone into seven regions based on morphological differences exhibited by the cells in each region after Movat’s

pentachrome (MP) staining (Figure 1A, B) and intravital staining with Hoechst and Calcein-AM (Figure 1C). These regions consisted of the two halves of the proliferative zone (P1 and P2), two halves of the pre-hypertrophic zone (p-H1 and p-H2), two halves of the hypertrophic zone (H1 and H2), and the single diaphyseal zone (D) (Figure 1C). We then mechanically and enzymatically dissociated these tissue regions separately into single-cell suspensions and sorted non-hematopoietic CD45-CD235- cells from each region into 96-well plates for scRNA-seq using Smart-seq2 (Figure 1D–E) (Picelli et al., 2014). To query our scRNA-seq data and identify rare cells with stem cell-like mRNA expression, we used the Gene Expression Commons (GEXC) (Seita et al., 2012) to search for genes specifically expressed in mSSCs and mBCSPs (Chan et al., 2015a), but not their downstream progeny, and created a gene set of 76 mSSC/mBCSP-specific genes and their human orthologs (Supplemental Table 1). We then calculated the percentage of transcriptionally active human orthologs to mSSC/mBCSP-specific genes for each sequenced single cell in each region. In contrast to all other regions of the bone, the cells within the second half of the pre-hypertrophic zone (p-H2) and the first half of the hypertrophic zone of the growth plate (H1) showed significantly higher expression of human orthologs to mSSC/mBCSP-specific genes (Figure 1F; Supplemental Figure 1A). A comparative analysis of the most variable cell-surface proteins across the different regions, particularly those that were significantly enriched or depleted within the hypertrophic zone (H1) of the growth plate compared to the diaphysis (D), unveiled a list of candidate markers that could provide a basis for antibody-mediated isolation of human growth plate cells by FACS (Supplemental Figure 1B).

Through flow-cytometric analysis of those candidate cell-surface markers against which antibodies were available, we identified a panel including PTPRC (CD45), GYPA (CD235a), TEK (TIE2), PECAM1 (CD31), PDPN, MCAM (CD146), NT5E (CD73), CD164, and THY1 (CD90) that separate growth plate zone cells into distinct subpopulations (Supplemental Figure 1C). Immunofluorescent (IF) staining against PDPN, CD146, CD73, and CD164 confirmed heterogeneity in the expression of these markers among the cells within the growth plate and the diaphysis. PDPN was primarily expressed in the growth plate and hypertrophic zone (p-H2, H1/H2), while CD146 was mainly expressed in the diaphyseal zone (D) (Supplemental Figure 1D, middle panel). In addition, a PDPN+CD146+ cell population was also present in the growth plate (p-H2, H1/H2) (Supplemental Figure 1D, middle panel). CD73 and CD164 were both enriched in the growth plate (p-H2, H1/H2) and further divided cells from this region into subpopulations based on the presence or absence of these two cell-surface markers (Supplemental Figure 1D, right panel). Revisiting the scRNA-seq data, we found that *PDPN*, *NT5E*, and *CD164* transcripts were enriched and *MCAM* transcripts were depleted in the p-H2/H1 zones of the growth plate—a region with high homology to the mSSC/mBCSP in terms of mRNA expression. This contrasted with the regions that showed low homology to the mSSC/mBCSP, such as the diaphysis (D) (Figure 1F; Supplemental Figure 1E–F). Moreover, by grouping cells based on expression of specific genes, we observed that *PDPN+MCAM-NT5E+CD164+* single cells were enriched in the growth plate (p-H2, H1/H2), depleted in the diaphysis (D), and comprised a larger percentage of the cells that showed high degree of homology to mSSC/mBCSP (Supplemental Figure 1E). Consistent with these findings, the p-H2 and H1 zones of the growth plate had a high frequency of non-hematopoietic (CD45-CD235a-), non-endothelial

(TIE2-CD31-) PDPN+CD146-CD73+CD164+ cells compared to other zones by flow cytometry (Supplemental Figure 1C). Therefore, we hypothesized that PDPN+CD146-CD73+CD164+ cells in the bone are likely enriched in hSSCs.

Prospective isolation and characterization of PDPN+CD146-CD73+CD164+ hSSCs

To assess the intrinsic skeletogenic potential of distinct human fetal growth plate cell-subsets *in vivo*, we used FACS to isolate seven distinct non-endothelial, non-hematopoietic cell populations based on their differential surface expression of PDPN, CD146, CD73, CD164, and THY1 and transplanted each population individually under the renal capsule of 8–12-week-old NOD scid gamma (NSG) immunodeficient mice (Figure 2A–E, Supplemental Figure 2) as described earlier (Chan et al., 2015a). The xenografts in the transplanted kidneys were explanted and sectioned for MP staining after 4 weeks of development. The staining revealed whether the transplanted cell-subset was unipotent (capable of forming only bone or only cartilage) or multipotent (capable of forming bone, cartilage and stroma). Two of the seven cell-subsets were multipotent and capable of endochondral (bone and cartilage) ossification—a PDPN+CD146-CD73+CD164+ and a PDPN+CD146^{lo} population (Figure 2C-c–d, 2E; Supplemental Figure 2). The rest of the five cell-subsets were unipotent and could be divided into two categories: 1) two osteogenic subsets, a PDPN-CD146+THY1^{hi} and a PDPN-CD146+THY1^{lo} population—hereafter collectively referred to as the human osteoprogenitor (hOP) (Figure 2C-a–b, 2E), and 2) three chondrogenic subsets, a PDPN+CD146-CD73-CD164+, a PDPN+CD146-CD73+CD164-, and a PDPN+CD146-CD73-CD164- population—hereafter collectively referred to as the human chondroprogenitors (hCPs) (Figure 2C-e–g, 2E). Immunostaining with human nuclear antigen confirmed that human skeletal progenitor-derived ossicles in the renal capsule were indeed of human origin (Figure 2D; Supplemental Figure 2B). We next measured the ability of these distinct populations to form colonies *in vitro*. Out of the two multipotent subsets (see above), the PDPN+CD146-CD73+CD164+ subset gave rise to a higher frequency of colony forming units (CFUs). We next asked whether the different cell-subsets were capable of *in vitro* self-renewal by testing their ability to serially generate CFUs. Strikingly, the multipotent PDPN+CD146-CD73+CD164+ cell subset was capable of CFU formation after secondary and tertiary serial transfer indicating self-renewal capacity (Figure 2F, G). In addition, colonies derived from the PDPN+CD146-CD73+CD164+ cell subset were composed of cells of both cartilage and bone lineages as shown by positive immunostaining for cartilage (Collagen type II) and osteoblast markers (Osteocalcin) (Figure 2H). On the contrary, CD146+ cells, which are commonly associated with mesenchymal stem cells (Lv et al., 2014; Sacchetti et al., 2007), displayed reduced colony size and frequency compared to PDPN+CD146-CD73+CD164+ cells as observed by light microscopy and flow cytometry (Supplemental Figure 2F-G).

To demonstrate self-renewing ability and multipotency in an *in vivo* serial transplantation setting, we labeled the purified PDPN+CD146-CD73+CD164+ cells from the fetal growth plate with DiD lipophilic dye and transplanted >1.5 million labeled cells under the renal capsule. The ossicles were explanted 2 weeks later, and cells were isolated after mild digest (Supplemental Figure 2A). The PDPN+CD146-CD73+CD164+ cells had undergone multiple cycles of division as evidenced by log-fold reduced DiD-label on flow cytometry

(Supplemental Figure 2B). We then sorted DiD^{lo} cells from the primary graft for an in vitro serial colony formation assay and a serial transplantation assay under the renal capsule of a secondary NSG mouse as above (Supplemental Figure 2A). Sorted DiD^{lo} cells from the primary graft were capable of at least two rounds of successive (tertiary) colony formation and exhibited similar morphology and distribution of hSSCs and lineage cells as control hSSCs that were freshly sorted from fetal growth plates (Supplemental Figure 2C). Secondary renal capsule grafts were explanted 4 weeks after transplant, grossly examined by imaging, and stained with MP. The secondary ossicle was comparable to the ossicle generated after primary transplant (Supplemental Figure 2D) and revealed regions of Alcian blue-labeled cartilage and Saffron yellow-labeled bone by MP staining (Supplemental Figure 2E). Therefore, the combined in vitro and in vivo results showing serial colony formation and serial reconstitution of multilineage grafts, respectively, supports the identity of PDPN⁺CD146⁻CD73⁺CD164⁺ fetal growth plate cells as self-renewing, multipotent skeletal stem cells—hereafter referred to as putative hSSCs.

Determination of human skeletal lineage hierarchy

To determine whether individual chondrogenic, osteogenic, and the multipotent skeletogenic populations are lineally related or whether they each represent independent, self-renewing stem and progenitor fractions, we evaluated the capacity of the seven flow-sorted skeletal populations (see above) to differentiate into other subsets in vitro and in vivo. We cultured each of the purified cell populations in serum-rich base media. At weekly intervals, adherent cells were lifted from the culture dish by collagenase digestion and then analyzed by FACS (Figure 3A). We found that only the putative hSSC fraction (PDPN⁺CD146⁻CD73⁺CD164⁺) had the capacity to give rise to all other fractions—PDPN⁺CD146^{lo}, PDPN⁻CD146⁺, and PDPN⁺CD146⁻CD73^{+/-}CD164^{+/-} subsets (Figure 3B). Alizarin Red S staining showed that these putative-hSSC-derived cells comprised distinct osteogenic and non-osteogenic cells (Figure 3C). We also measured the adipogenic differentiation potential of the sorted putative hSSCs by either subjecting them to in vitro differentiation or by analyzing the ossicles formed after transplanting them in RFP-mice and did not observe any evidence of adipogenesis by either fetal or adult bone marrow-derived cell subsets (Supplemental Figure 3).

To validate our in vitro results, we transplanted 100,000 freshly isolated putative hSSCs under the renal capsule of immunodeficient NSG mice. We isolated the grafts 4 weeks after transplant and analyzed the cell suspension derived from the graft by FACS. As seen in vitro, the putative hSSCs gave rise to the other skeletal subsets after transplant (Figure 3D). To demonstrate *in situ* clonal contribution to multiple skeletal lineages, we labeled the putative hSSCs randomly with multiple distinct fluorescent proteins (RFP, EGFP, and CFP) using lentiviral transduction prior to transplant under the mouse renal capsule (Weber et al., 2011). We found that clonally marked putative hSSCs generated both cartilage and bone staining regions in the subcapsular ossicles (Figure 3E; Supplemental Figure 2E). Collectively, these data show that the putative hSSC fraction (PDPN⁺CD146⁻CD73⁺CD164⁺) is enriched in hSSCs that are capable of lineal generation of chondrogenic and osteogenic subsets and are at the top of the skeletogenic differentiation tree (Figure 3F).

hSSCs are locally amplified in response to skeletal injury

Stem cells determine the regenerative potential of adult tissues by constantly replenishing cells that have been damaged by aging, injury, or disease. As we observed earlier in mice (Marecic et al., 2015), hSSCs are significantly amplified in soft-callus fracture specimens when compared to their frequency in uninjured limb skeletal tissues that were obtained from patients undergoing bone graft procedures (Figure 4A, D upper panel; Supplemental Table 2). It is possible that variability among patients could confound analysis of fracture specimens obtained from different patients. Therefore, we established a new human bone xenograft mouse model to evaluate the effects of skeletal injury in human limb bones in a more controlled setting. We transplanted carefully dissected human fetal phalangeal grafts with intact periosteum obtained from 18-week fetuses into 5-day-old immunodeficient NSG mice (Figure 4B). The human fetal phalanges efficiently engraft and develop into growing bones on the flank of a mouse (Figure 4B). The phalangeal grafts grew much larger in size than recipient host mouse limbs. Moreover, we found that localized injury induced by unicortical fractures on the xenografted phalangeal bones resulted in significant localized expansion of hSSCs relative to unfractured areas in the same bone (Figure 4C–D, lower panel). Therefore, consistent with the fracture-induced expansion of mSSCs in mice (Marecic et al., 2015), the injury-induced hSSC (PDPN+146-CD73+CD164+) expansion in clinical specimens and in xenografts represents a regenerative response to skeletal injury.

hSSCs generate hematopoiesis-supportive stroma

As observed with mSSCs (Chan et al., 2009; Chan et al., 2013), the hSSCs transplanted under the renal capsule generate ossicles with a marrow cavity, indicating that they may give rise to skeletal lineage-derived, hematopoiesis-supportive stroma in addition to bone and cartilage. To determine whether hSSCs and hSSC-derived subsets can functionally maintain hHSCs, we established monolayer cultures with 100,000 purified hSSCs (PDPN+CD146-CD73+CD164+), hBCSPs (PDPN+CD146+), or hOPs (PDPN-CD146+THY1+/-) freshly isolated from 17–19-week-old fetal limbs. After cultures reached confluency, 5,000 Lin-CD34+CD38-THY1^{lo}CD45RA-cord blood hHSCs (Baum et al., 1992) were co-cultured with hSSCs, hBCSPs, hOPs, or no stroma in serum-free conditions supplemented with stem cell factor (SCF) and thrombopoietin (TPO). After 14 days of co-culture, the cells were lifted with EDTA and transplanted into recipient immunodeficient NSG mice that were conditioned using sublethal irradiation as shown earlier (Reinisch et al., 2015). Freshly isolated cord-blood hHSCs were also transplanted as a positive control. Eight weeks after transplant, bone marrow aspirates were collected from recipient mice and analyzed for the presence of human hematopoietic reconstitution (Supplemental Figure 4A). While there was little engraftment from transplants of hHSCs cultured without stroma, all hHSCs co-cultured with skeletal progenitor populations gave rise to some levels of engraftment. hHSCs co-cultured with hBCSPs generated the highest level of multilineage myeloid and lymphoid engraftment (Supplemental Figure 4B–C). We found that hSSCs and hSSC-derived subsets, express a variety of potential hematopoiesis-supportive cytokines with matching cognate receptors expressed on hHSCs and downstream hematopoietic progenitors, including *ANGPT1*, *CSF1*, *SDF*, *IL27*, *IL7*, and *SCF* (Supplemental Figure 4D–G). Consistently, we also found that hHSCs express a variety of factors that could have a potential role in supporting the hSSC lineage, including *BMP2*, *BMP8A*, *DHH*, *FGF3*, *WNT1*, and *WNT8*

(Supplemental Figure 4F-i), supporting a model of stem-cell crosstalk within the bone marrow.

PDPN+CD146-CD73+CD164+ expression identifies hSSCs from other skeletal tissue sources

We had observed in mice that mSSCs in fetal, newborn, and adults have similar surface marker profiles (Chan et al., 2015a), suggesting that hSSCs may also be present in a variety of different skeletal tissues and may be isolated by the same set of surface markers. Indeed, we found that cells isolated from freshly resected adult human femoral head specimens also contain significant populations of non-hematopoietic, non-endothelial (CD45-CD235-TIE2-CD31-) cells with distinct patterns of staining for PDPN, CD146, CD73, and CD164 (Figure 5A-i). As in fetal tissues, adult PDPN+CD146-CD73+CD164+ cells readily differentiate into osteoblasts and cartilage in vitro (Figure 5A-ii) and form skeletogenic ossicles when transplanted under the renal capsule or subcutaneously in vivo (Figure 5A-iii-v, 7D). We could also isolate hSSCs from bone marrow aspirates, although the percent of skeletal lineage cells, including hSSCs, was contingent on the presence of bone spicules in the sample (data not shown). PDPN+CD146-CD73+CD164+ expression also enriches for hSSCs formed by skeletal induction of human monocyte-derived iPSCs (Karp et al., 2006) (Figure 5B-i). Cultured iPSC-derived hSSCs form monolayers of mineralizing osteoblasts and micromasses of chondrocytes (Figure 5B-ii). In addition, subcapsular transplants of these cells form multilineage ossicles containing marrow cavity in vivo (Figure 5B-iii-v, 7D). We previously observed that high concentrations of BMP2 triggers skeletal reprogramming of adipose stromal cells in mice (Chan et al., 2015a; Lo et al., 2012). Therefore, we tested whether subcutaneous injection of human adipose stroma with high levels of recombinant BMP2 in matrigel can give rise to heterotopic ossicles enriched for PDPN+CD146-CD73+CD164+ hSSCs (Figure 5C-i). We observed that while PDPN+CD146-CD73+CD164+ hSSCs are not present in freshly isolated human adipose stroma (HAS) (Figure 5C-ii left), the BMP2-treated human adipose stroma (B-HAS) form vascularized heterotopic ossicles that harbor PDPN+CD146-CD73+CD164+ hSSCs and downstream PDPN-CD146+ hOPs (Figure 5C-ii middle, C-iii, 5D). In contrast, when VEGF signaling is antagonized by co-delivery of excess soluble VEGF-receptor during BMP2-induced ossification, bone formation is repressed in favor of chondrogenesis as PDPN+CD146-CD73-CD164+ hCPs increase in frequency in the induced ossicles that lack blood vessels (Figure 5C-ii right, C-iv, 5D).

Single-cell transcriptomic comparison of fetal-, adult-, B-HAS-, and iPSC-derived hSSCs

We observed differences in the skeletogenic potential of hSSCs isolated from fetal, adult, B-HAS, and iPSC tissues (Figure 5D). For example, fetal and iPSC-derived SSCs have a higher tendency to form cartilage than adult SSCs. To investigate these phenotypic differences between the various sources of hSSCs, we compared single hSSCs from fetal, adult, B-HAS, and iPSC-derived tissues using scRNA-seq (Figure 6A). Principal component analysis (PCA) of 42 fetal, 36 adult, 77 B-HAS-derived, and 41 iPSC-derived hSSCs demonstrates that hSSCs isolated from different sources cluster together (Figure 6B). However, the second principal component (PC2) separates a few iPSC-derived hSSCs from this cluster as outliers. The gene loadings that most contribute to PC2 include important

early skeletogenic factors such as *SPARC*, *COL1A1*, *SFRP2*, *LUM*, and *FOS* (Figure 6B)—suggesting that the outliers may comprise hSSC precursors or poorly differentiated cells during skeletal induction of iPSCs. To specifically study single hSSCs from a skeletal and developmental perspective, we curated a set of signaling pathways and bone development genes from the GEXC and the Molecular Signatures Database (Supplemental Table 3) (Liberzon et al., 2015; Seita et al., 2012). Using these 296 genes, we performed supervised hierarchical clustering of 196 single hSSCs and identified ten gene clusters (Figure 6C; 1–10). Five clusters demonstrate transcriptional similarity in expression of key skeletogenic genes, including *SPARC*, *COL1A1*, *ITGB1BPI*, and *RHOA* (Figure 6C; 1–4, 9). Four clusters recapitulate the phenotypic similarity of adult to B-HAS-derived hSSCs, and fetal to iPSC-derived hSSCs based on key genes such as *OSR2*, *SPG20*, *FOS*, and *WLS* (Figure 6C; 5, 6, 8, 10). The last cluster captures transcriptional differences in expression levels between primary isolates and in vitro derived hSSCs based on genes such as *TCF7L2*, *FST*, and *CHRDL1* (Figure 6C; 7). To assess the degree of similarity between individual hSSCs and groups of hSSCs, we calculated the Pearson correlation coefficients for each pair of single and groups of hSSCs (Figure 6D, E). When analyzed separately, we found that adult hSSCs were more heterogeneous compared to fetal, B-HAS-derived, or iPSC-derived hSSCs. The greater degree of heterogeneity among adult hSSCs may be a result of diverse specialization within the stem cell pool with age (Figure 6D). Upon aggregating cells within each group, we found that fetal hSSCs most closely correlated with iPSC-derived hSSCs and B-HAS-derived hSSCs most closely correlated with adult hSSCs. When we removed the iPSC-derived hSSCs from the comparison, we found that fetal hSSCs correlated more with adult hSSCs than with B-HAS-derived hSSCs (Figure 6E). The correlation data (Figure 6E) is largely consistent with our functional data (Figure 5D) except for B-HAS- and iPSC-derived hSSCs, both of which are functionally similar with respect to their differentiation potential but are less similar by single-cell mRNA expression.

Convergent and divergent skeletal regulatory programs between mouse and human SSCs

Humans and mice diversified from a common ancestor nearly 100 million years ago (Nei and Glazko, 2002). While many aspects of their physiology are similar, mouse and humans also differ in several ways related to their skeletal systems. We observe that sub-renally transplanted human fetal bones grow dramatically larger after 2 months of growth compared to their mouse counterparts (Figure 7A). In addition, hSSCs produce up to 100-fold larger CFUs compared to mSSCs (Figure 7B). These results suggest some aspects of skeletal size control are cell intrinsic and could reflect differences in genetic and epigenetic regulation at the level of individual SSCs.

To address this, we purified human and mouse fetal SSCs at similar developmental stages (e17.5 mouse and 19-week-old human fetal bones) for a comparative gene expression analysis by microarray. We normalized expression values using the GEXC (Seita et al., 2012) that assigns a threshold value to each probeset using StepMiner (Sahoo et al., 2007) and calculates a percentile value between –100% and +100% for each available gene, allowing us to compare mouse and human gene expression on a normalized, continuous scale. Focusing specifically on signaling pathways and genes associated with bone development (Supplemental Table 3), a pairwise comparison between fetal e17.5 mSSC and

19-week fetal hSSC revealed that over 66.5% of transcription within the selected signaling pathways and bone development genes are expressed in both mSSCs and hSSCs. We analyzed the genes that constitute the WNT, BMP, Hedgehog, FGF, and Notch pathways as well as additional genes that constitute the gene ontology (GO) gene set for bone development. As expected, a few active genes in mSSCs were turned off in hSSCs, while other genes that were not expressed in mSSCs were found to be active in hSSCs (Figure 7C, Supplemental Figure 5). Notable among divergent genes are key members of the WNT signaling pathways, *SOST*, *CXXC4*, and *DNAJB6*, all of which have been implicated earlier in human bone development but are not expressed in mSSCs (Figure 7C) (Kojima et al., 2009; Menezes et al., 2012; Moester et al., 2010). Among the genes that are expressed by both mouse and human SSCs, we found some well-known master transcriptional regulators of skeletogenesis, including *RUNX2*, *SOX9* (Figure 7C)—in addition to some lesser known transcription factors (TFs) such as *OSR2* and *TWIST1* (Supplemental Table 3).

To better understand epigenetic regulation of skeletal development, we performed ATAC-seq on mouse and human SSCs isolated from bones at similar developmental stages (18-week-old human fetal bone and e17.5 mouse fetal bone) as above to identify sites of accessible chromatin. This analysis revealed various open chromatin-sites comprising active promoters as well as gene-distal enhancers that are conserved between mouse and human (Supplemental Figure 6). To drill-down to the species-specific epigenetic differences, we focused our analysis on genes that are expressed in the hSSC but not in the mSSC (see above). For these gene-loci, we found that in human (but not always in mouse), the accessible open chromatin sites were enriched in TF-binding-motifs recognized by the core SSC-specific-TFs that are conserved between human and mouse in terms of their expression. For example, even though *RUNX2* is expressed in both human and mouse (Figure 7C), the predicted *RUNX2* binding-sites are enriched in the open chromatin landscape of human *SOST* but not mouse *Sost* locus and therefore, *SOST* is expressed only in hSSCs but not mSSCs (Figure 7D). The same is true for another human-specific gene, *DNAJB6* (Supplemental Figure 6I). Interestingly, overexpression of either human *SOST* or *DNAJB6*—both of which are inhibitors of WNT signaling—by lentiviral transduction in mSSCs led to significantly larger subrenal ossicle formation relative to GFP virus control (Supplemental Figure 7A–E). This is also consistent with the delayed osteogenesis in *SOST* or *DNAJB6* transduced mSSC-derived ossicles—as exemplified by the reduced bone staining (yellow) and enhanced cartilage staining (blue) of cross-sections of the ossicles harvested 16-days after transplant (Supplemental Figure 7C-iv lower panel, 7D-iv lower panel). These data suggest that functional genes that are expressed in a species-specific manner may be a result of divergent evolution of epigenetic regulation.

Genome-wide association studies (GWAS) have identified several non-coding genomic polymorphisms that are associated with skeletal traits in humans such as bone mineral density and height (Chan et al., 2015b; Estrada et al., 2012). We analyzed 56 such single nucleotide polymorphisms (SNPs) that are strongly associated with variation in bone mineral density in humans and found that one of the SNPs changes the predicted *RUNX* binding site within the *SOST* locus (Figure 7D) while another was within the accessible non-coding regions surrounding the *WNT4* promoter (Supplemental Figure 6J). These

observations further exemplify the key role played by the SSC-specific genes both during development and homeostasis.

Discussion

Our data support a developmental hierarchy of skeletal progenitors in humans organized by cells with differential expression of PDPN, CD146, CD73, and CD164. We find that a purified population of PDPN+CD146-CD73+CD164+ cells serially generate CFUs from single cells in vitro and multilineage ossicles containing bone, cartilage, and stroma upon subrenal transplantation in mouse. Notably, these stem cells do not differentiate into adipocytes, suggesting that adipocytes may be replenished by other sources of resident bone marrow stem cells. Instead, hSSCs lineally transition into an early bone, cartilage, and stroma progenitor (hBCSP), which then give rise to osteoprogenitors (hOPs) and chondroprogenitors (hCPs) prior to terminally differentiating into bone, cartilage, and stroma.

hSSCs from different skeletal sources are similar yet distinct

We show that hSSCs and their downstream progeny can also be isolated from skeletally-differentiated iPSCs and B-HAS. When compared to fetal and adult hSSCs, we observe that there is variation in the proportion of cartilage and bone in the ossicles from different hSSC sources, which correlates to gene expression differences between individual hSSCs as quantified by scRNA-seq. The gene sets that segregate different hSSC populations provide molecular landmarks for modifying skeletogenic conditions to endow B-HAS- or iPSC-derived hSSCs with enhanced self-renewal or increased chondrogenic and/or osteogenic activity.

A molecular glimpse of skeletal speciation

Finally, we compared mouse and human SSCs to gain insight into the evolutionary basis of stem cell-mediated skeletogenesis. We found that when equally sized human and mouse fetal bones were transplanted in mice, the human fetal bones far outgrew their mouse counterparts. Similarly, after equal time in culture, CFUs from hSSCs had over 100-fold more cells than CFUs from mSSCs. These initial observations along with molecular comparisons of SSCs from mouse and humans using microarray, and ATAC-seq analysis led us to suspect that there are intrinsic mechanisms that regulate skeletogenesis and that molecular comparison of mouse and human SSCs could explain divergences in skeletal phenotype.

Overexpression of two genes (*SOST* and *DNAJB6*) in mSSCs that are expressed in hSSCs but not mSSCs, lead to significantly larger mSSC-derived ossicles, suggesting that *SOST* and *DNAJB6* expression in hSSCs may be important for skeletal size control at the stem cell level. This was unexpected as *SOST* is a known negative regulator of WNT signaling and bone formation in osteoblasts (Compton and Lee, 2014). We hypothesize that expression of *SOST* by hSSC could be important for extending hSSC activity during fetal development, perhaps by delaying premature WNT-dependent osteoblast differentiation and/or osteoblast-mediated feedback signaling.

Our analysis also revealed differential expression of various genes involved in the synthesis of connective proteins (Supplemental Table 4, 5), such as types of collagen that could underscore differences in biomechanical properties between mouse and human skeletal tissues. With the identification of the SSC in both mouse and human, and the delineation of their downstream skeletal lineages, we provide a cell atlas to enable a series of new investigations into the genetic and molecular determinants of skeletal formation and development. Moreover, given the tremendous medical burden imposed by degenerative, neoplastic, post-traumatic and post-surgical skeletal disorders, we believe that identifying the hSSC and elucidating its lineage map will enable the molecular diagnosis and treatment of skeletal diseases (Beyth et al., 2011). Using this approach, we recently showed that even complex skeletal pathologies that arise from systemic diseases like diabetes can be precisely diagnosed and remedied by comparing the molecular phenotype of normal and diseased SSCs (Tevlin et al., 2017). Therefore, we anticipate that further characterization of the hSSC, and the ontogeny of hSSC-derived skeletal progenitors as described here will facilitate the development of patient-specific approaches to diagnosing and reversing diverse types of skeletal disorders.

STAR METHODS

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to, and will be fulfilled by, the Lead Contact, C.K.F.C.: chazchan@stanford.edu

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice—BA mice (C57BL/Ka-Thy1.1-CD45.1), purchased from the Jackson laboratories (Mouse strain datasheet JAX:000406), and used for isolation of mouse SSC and skeletal progenitors for gene expression and ATAC-sequencing experiments as described below. NSG (NOD.Cg-Prkdc^{scid} Il2rg^{tm1Wjl}/SzJ, JAX: 005557) immunodeficient mouse strains were used to establishing the human xenograft model and as recipients for either subcutaneous or renal capsule transplantation of human skeletal and hematopoietic stem and progenitor cells. RFP (B6.Cg-Tg(CAG-mRFP1)1F1Hadj/J, JAX: 005884) mice were used for hSSC and HAS/B-HAS transplantation to assess adipogenic potential in vivo. All mice were male and 8–12-weeks-old unless otherwise stated in the results section. Animals were maintained at Stanford University animal core facility in accordance with Stanford and National Institutes of Health guidelines.

We do not have any information regarding the sex of fetal or cord blood specimens utilized during this study. The gestation ages of fetal specimens are listed in Supplemental Table 2 and described in the respective results sections. Human adipose stroma (HAS) were obtained from female patients ranging from 30–50 years old for both in vitro and in vivo adipogenesis experiments as well as skeletal induction and scRNA-seq experiments. Adult hSSCs, and corresponding lineage populations, were obtained from femoral head bone marrow from a 58-year old male, a 71-year old female, and a 58-year-old female donor. Age and sex of donors that contributed fractured bone specimens is listed in Supplemental Table 2. The iPSC line, SCVI 113, used in this study was derived from peripheral blood monocytes of a

51-year-old male. We do not have age or sex information regarding primary cells used in the chondrogenic differentiation studies.

Fetal and adult human samples—Fourteen human fetal samples were obtained from Stemexpress (Folsom, CA) and shipped overnight. Samples ranged in age from 10 to 20 weeks of gestation with no restrictions on race or gender. Fetal sample procurement and handling was in accordance with the guidelines set by the Institutional Review Board (IRB-35711). 22 adult femoral heads (58–74 years old), 3 uninjured human bones (16–30 years old), and 10 fractured long bone specimens (20–90 years old) were obtained from Stanford Hospital in accordance with guidelines set by the Institutional Review Board (IRB-35711). On this IRB, informed consent was not required as samples were considered biological waste. No restrictions were made regarding the race, gender, or age of the specimen's donor. Following excision, all femoral head specimens were placed on ice, and skeletal stem and progenitor cells were isolated from the bone marrow cavity and articular cartilage, as described below. Adipose stromal cells were harvested from lipoaspirate human tissue (n=5) from female donors, 30–50 years old. Informed consent was obtained in all cases, and samples were obtained from Stanford Hospital in accordance with guidelines set by the Institutional Review Board (IRB-25954). Human core-blood samples were obtained according to the Administrative Panel on Human Subjects Research Institutional Review Board (IRB)-approved protocols (Stanford IRB no. 18329, no. 6453, and no. 5637) with informed consent. Cord blood was collected with written informed consent from the mother, which was obtained before delivery of full-term pregnancies at the Lucile Packard Children's Hospital, according to IRB-approved protocols (Stanford IRB no. 5637) or purchased from the New York Blood Center (NYBC). All cord-blood samples were processed within 24 h after delivery and used fresh.

Cell lines—Human monocyte derived induced Pluripotent Stem Cell (iPSC) line, SCVI 113, was a kind gift from the laboratory of Joseph C. Wu (joewu@stanford.edu). Donor was male 51-year-old.

Bacteria and Virus strains—Lentivirus expressing GFP, RFP, and CFP were subcloned in the Weissman laboratory from pLL3.7 deposited in Addgene (<https://www.addgene.org/11795/>). Validated lentiviral expression plasmids containing cDNA expression cassettes for *SOST* (Cat#Mm11378) and *DNAJB6* (Cat#Mm21356) were acquired from Genecopeia.

METHOD DETAILS

Human Fetal and Adult Bone dissociation—Human fetal long bones were separated, gently denuded of soft tissue with paper towels, and dissected for the growth plate and its constitutive zones. The distinct areas of the growth plate were distinguished based on differences in the size and shape of the chondrocytes and osteoblasts that are representative of each growth plate zones. These differences can be visualized in fixed tissue by histology with Movat Pentachrome (Saffron [Sigma-Aldrich, Cat#S8381–5G], Acid Red 73 [Sigma-Aldrich, cat#49823–25MG], Alcian Blue 8GX [Sigma-Aldrich, Cat#A3157–25G], Acid Fuchsin [Sigma-Aldrich, Cat#F8129–25G], Phosphotungstic Acid [Sigma-Aldrich, Cat#P4006], Hematoxylin [Sigma-Aldrich, Cat#MHS32–1L]) or in living tissue by vital dye

staining with Calcein-AM (Invitrogen, Cat#C1430) and Hoechst (Thermo Fisher Scientific™, Cat# 62249) (Figure 1B–C). The proliferative zone (P) is identified by the relatively small size of the chondrocytes, the pre-hypertrophic zone chondrocytes (p-H) are medium-sized, and the hypertrophic chondrocytes (H) are among some of the largest chondrogenic cells in the growth plate. Finally, the diaphyseal region (D) includes trabecular bone and marrow. For scRNA-seq analysis, we further split each of the chondrocyte zones (P, p-H, and H) into equivalently sized halves (1 & 2)—thus, a total of 7 distinct regions were used for analysis including the diaphyseal zone.

From adult human femoral head specimens, the articular cartilage was dissected out with scalpels and the bone marrow was scraped out with a spatula from the marrow cavity. The marrow and the cartilage tissue were then minced separately with razor blades, resuspended in 3000 U/mL type II collagenase (Sigma-Aldrich, Cat#C6885) digestion buffer supplemented with 100 U/mL DNase I (Worthington, Cat#NC9199796) and incubated at 37 °C for 40 minutes under constant agitation. The supernatant was filtered through a 70 µm nylon mesh and quenched with staining media (2% fetal bovine serum (FBS), in phosphate-buffered saline (PBS), GIBCO, Cat#C14190500BT). Digestion was repeated twice more prior to centrifugation at 200g at 4 °C followed by resuspension in staining media.

Fluorescence-activated cell sorting (FACS)—Human skeletal cells were separated from red blood cells (RBCs) and bone dust by density gradient separation using 1:1 Histopaque of density 1.119 g/mL (Sigma-Aldrich, Cat#11191). The buffy coat was collected, washed with staining media, and the resulting cell suspension was depleted of CD45+ cells by magnetic-activated cell sorting (MACS) (Miltenyi, Cat#130–045-801). Cells were blocked with mouse IgG and stained with fluorochrome-conjugated antibodies against CD45 (BioLegend, Cat#304029-BL), CD235a (BioLegend, Cat#306612-BL), CD31 (Thermo Fisher Scientific™, Cat#13–0319-82), CD202b (Tie-2) (BioLegend, Cat#334204), CD146 (BioLegend, Cat#342010), PDPN (Thermo Fisher Scientific™, Cat#17–9381-42), CD90 (THY1; BioLegend, Cat#328110), CD164 (BioLegend, Cat#324808), and CD73 (BioLegend, Cat#344016). Flow cytometry was performed on FACS Aria II (BD Biosciences). Gating schemes were established with fluorescence-minus-one (FMO: staining with all fluorophores except one) controls and negative propidium iodide (PI) (Sigma-Aldrich, Cat#P4170) staining (1 µg/ml) was used as a measure for cell viability.

In vivo transplantation of cells—SSCs or other skeletal cells were purified using FACS and resuspended in 2 µL of Matrigel (Corning, Cat#CB40234A) and then injected subcutaneously or underneath the renal capsule of 8–12-week-old immunodeficient NSG mice. Injected cells developed into a graft after 4 weeks. The grafts were surgically removed for analysis. For serial transplantation, SSCs from the graft were isolated by FACS, labeled with DiD lipophilic dye-AF647 (Cat#V22887), and >1.5 million cells were transplanted under the renal capsule of 8–12-week-old NSG mice. Ossicles were explanted 2 weeks later for analysis.

In vivo transplantation of whole bones—Fresh 10-week-old human fetal and e14.5 mouse fetal femurs were transplanted underneath the renal capsule of 8–12-week-old

anesthetized immunodeficient NSG mice. Bones were incubated for 2 months prior to surgical excision and analysis of graft.

Human xenograft Model—Ten 18-week-old fetal phalanges (proximal, middle and distal phalanges) were transplanted subcutaneously into postnatal day 5 NSG mouse pups. The pups reached skeletal maturity at 4 weeks and the xenograft was opened. Unicortical fractures were performed using a drill as shown in Figure 4B. The fractures were allowed to heal for 2 weeks and then harvested and dissociated as described above. Cells were analyzed by FACS and underwent osteogenic differentiation and Alizarin Red staining (Sigma-Aldrich, Cat#A5533) to assess osteogenic potential.

Immunofluorescence—Immunofluorescence (IF) on cryopreserved ectopic bone specimens were performed using M.O.M. immunodetection kit (Vector Laboratories, Cat#BMK-2202) according to manufacturer's instructions. Briefly, specimens were treated with a blocking reagent 5–2% donkey (Sigma-Aldrich Cat#D9663) or goat serum (Thermo Fisher Scientific™, Cat#G9023) supplemented with 1% Bovine Serum Albumin (BSA; Sigma-Aldrich, Cat#A9647) in Tris Buffered Saline (TBS, Sigma-Aldrich, Cat#T5912), then probed with monoclonal primary antibody at 4 °C overnight. Primary antibodies: CD164 (Sigma-Aldrich, Cat#HPA010636), Perilipin A (Chemicon, Cat#AB10200), Adiponectin (Invitrogen, Cat#MA1-054), Human Nuclear Antigen (Abcam, Cat#ab191181), Collagen II (Abcam, Cat#ab34712), Osteocalcin (Abcam, Cat#ab13418). Specimens were next washed with PBS, probed with secondary antibodies conjugated to Alexafluor-488 (AF488; Invitrogen, Cat#A11006) or AF647 (Invitrogen, Cat#A21236) as needed. Specimens were incubated with 1 ug/ml of DAPI (Biolegend, Cat#422801) for 10 mins and then washed twice. The specimens were then mounted with a coverslip using Fluoromount-G (Invitrogen, Cat#00-4958-02), and imaged with a Leica DMI6000B inverted microscope system.

IF on tissue-cultured cell specimens were performed similarly to cryopreserved specimens. Briefly, cultured cells in 6-well to 96-well culture plates were washed with PBS and fixed in 2% paraformaldehyde (PFA, Electron Microscopy Sciences, Cat#15710) at 4°C overnight. Specimens were treated with a blocking reagent (see above), then probed with monoclonal antibody at 4°C overnight. Next, specimens were washed with PBS, probed with secondary antibodies conjugated to Alexafluor-488 (AF488; Invitrogen, Cat#A11006) or AF647 (Invitrogen, Cat#A21236) as needed, washed, immersed in PBS, and imaged with a Leica DMI6000B inverted microscope system (LAS X software, Leica Microsystems). Monoclonal antibodies: human Collagen II (Abcam, Cat#ab34712), Osteocalcin (Abcam, Cat#ab13418), AF-conjugated secondary antibodies (AF455; Molecular Probes Cat#A11034, and AF647; Molecular Probes, Cat#A31573).

In vivo skeletal-induction with local factors—Adipose stromal cells were harvested from lipoaspirate human tissue, collagenase (Type II) digested, and then depleted of CD45-expressing cells by magnetic separation using anti-human CD45 microbeads (Miltenyi, Cat#130-045-801). 10 million CD45- lipoaspirate cells were then resuspended in 50 µL of Matrigel containing 10 µg rhBMP2 (recombinant human BMP2; R&D Systems, Cat#355-BM/CF). The cell solution was then injected subcutaneously into the inguinal fat pads of 8–

12-week-old adult NSG immunodeficient mice. One month after transplant, grafts were harvested for histological or FACS analysis as described.

Human adipose stromal cells were harvested and transplanted as above. Local induction with 10 ug rhBMP2 (recombinant human BMP2; R&D Systems, Cat#355-BM/CF) alone or rhBMP2 with rhVEGFR1 (recombinant human VEGFR1; R&D Systems, Cat#321-FL/CF) at a ratio of 3:50 μg allowed for osteo and chondro-induction, respectively. The recombinant proteins were mixed directly with the matrigel and the cell suspension.

Single Cell RNA sequencing (scRNA-seq)

Single cell capture and RNA extraction: Cells were dissociated into a single cell suspension and prepared for FACS as described under **Human Fetal and Adult Bone dissociation and Fluorescence-activated cell sorting (FACS)** above. Single cells were sorted into 96-well plates of lysis buffer (1 U/ μL RNase inhibitor (Clontech, Cat#2313B), 0.1% Triton (Thermo Fisher Scientific™, Cat#85111), 2.5 mM dNTP (Invitrogen, Cat#10297-018), and 2.5 μM oligo dT30VN (IDT, special order) in nuclease-free water (Thermo Fisher Scientific™, Cat#10977023)) as described (Picelli et al., 2014), centrifuged at maximum speed for 30 sec at 4 °C, snap frozen on dry ice, and then stored at -80 °C. ERCC (External RNA Controls Consortium) ExFold RNA Spike-In Mixes (Invitrogen, Cat#4456739) was added at 1:600,000 dilution to each lysis reaction to be processed in parallel to cellular mRNA.

Whole-transcriptome amplification, library preparation, and sequencing: Single cell cDNA libraries were constructed using the Smart-seq2 protocol (Picelli et al., 2014) with minor modifications. Briefly, cDNA from single cell RNA was generated by oligo-dT primed reverse transcription with MMLV SMARTScribe reverse transcriptase (Clontech, Cat#639538) and a locked template-switching oligonucleotide (TSO; Exiqon, special order). This was followed by 25 cycles of PCR amplification, which was the optimal number of cycles determined to obtain cDNA concentration of at least 0.05–0.32 ng/ μL from each human skeletal lineage cell, using KAPA HiFi hotStart ReadyMix (Kapa Biosystems, Cat#KK2602) and ISPCR primers (IDT; IDT, special order). Amplified cDNA was then purified using 0.6 \times volume Agencourt AMPure XP beads (BEECKMAN COULTER, Cat#A63882) to remove oligos smaller than 500 base pairs (bp). The resulting cDNA concentration and size distribution for each well was determined on a capillary electrophoresis-based fragment analyzer (Advanced Analytical). Based on this analysis, the cDNA concentration was then normalized to a desirable concentration range of 0.05–0.32 ng/ μL by diluting the amplified cDNA from each well. The cDNA was tagged, uniquely barcoded, and PCR enriched using the Nextera XT DNA Library Prep kit (Illumina, Cat#FC-131-1096). Libraries were then pooled and purified to remove fragments smaller than ~400 bp using 0.6 \times volume Agencourt AMPure XP beads. Pooled libraries were quantified by Bioanalyzer (Agilent), using High Sensitivity DNA analysis kit, and fluorometrically, using Qubit dsDNA HS Assay kits and a Qubit 2.0 Fluorometer (Invitrogen), and the final concentration was adjusted to 2 nM. The pooled libraries were then sequenced on NextSeq 500** (Illumina) to obtain 2 \times 151 bp paired-end reads at a depth of 1–4.5 \times 10⁶ reads/cell.

**Pooled libraries were initially sequenced on HiSeq 4000 (Illumina); however, due to presence of significant amount of index-switching (Sinha et al., 2017) in our data, library pools were rerun on the NextSeq 500.

Transcriptional Expression Profiling by microarray

mRNA isolation and microarray: We performed microarray analyses on highly-purified, double-sorted populations of e17.5 mouse and 19-week human SSCs. 10–30,000 cells were directly sorted into 1 mL TRIzol LS (Invitrogen, Cat#10296028). RNA was isolated with RNeasy Micro Kit (Qiagen, Cat#74004) as per manufacturer's instructions. RNA was twice amplified with an Arcturus™ RiboAmp™ PLUS Kit (Applied Biosystems™, Cat#KIT0521). Amplified cRNA was streptavidin-labeled, fragmented, and hybridized to Affymetrix arrays 430–2.0 (for mouse genome; Applied Biosystems™, Cat#900495), and HG-U133+ (for human genome; Applied Biosystems™, Cat#901569). Arrays were scanned with a Gene Chip Scanner 3000 (Affymetrix) running GCOS 1.1.1 software.

Histology—Dissected specimens were fixed in 2% PFA at 4 °C overnight. Specimens were decalcified in 400 mM Ethylenediaminetetraacetic acid (EDTA; Invitrogen, Cat#15573–038) in PBS (pH 7.2) at 4 °C for 2 weeks with a change of EDTA once at 1 week. The specimens were then dehydrated in 30% sucrose at 4 °C overnight. Specimens were embedded in Optimal Cutting Temperature compound (OCT) and sectioned at 5 μm. Representative sections were stained with freshly prepared Hematoxylin (Sigma-Aldrich, Cat#MHS32–1L) and Eosin (Eosin Y; Thermo Fisher Scientific™ Shandon™, Cat#6766009), Movat pentachrome, or Alizarin Red (Sigma-Aldrich, Cat#A5533) stains depending on the individual experiment. For zone demarcation in the growth plate and diaphysis, live bones were bisected along the longitudinal axis of the bone and stained directly with Hoechst (Thermo Fisher Scientific™, Cat#62249) and Calcein-AM (Invitrogen, Cat#C1430) intravital dyes as per standard protocols.

Cell culture—Skeletal progenitors were cultured in MEM-alpha medium with 10% human platelet-derived lysate (Stem Cell Technologies, Cat#06960), 1% Penicillin-Streptomycin solution (Pen-Strep; Thermo Fisher Scientific™, Cat#15140–122) in an incubator maintained at 37 °C with 5% CO₂. Culture vessels were first coated with 0.1% Gelatin (Millipore, Cat#ES-006-B). Cultured cells were lifted for analysis or passaging by incubating with 3000 U/mL type II collagenase in M199 (Sigma-Aldrich, #C6885).

For human colony forming assays, single cells were sorted into each well of a 96-well plate and cultured for 2 weeks. At this time, specimens were examined under a phase microscopy and a cloning ring was used for quantification. The cells were subsequently lifted for staining and analysis by FACS.

Adipogenic Differentiation—Populations of hSSC, hBCSP, hCP and hOP were prospectively isolated using FACS. Cells were seeded at a predetermined uniform density depending on typical cell yield from the specimen. The cell populations were subjected to adipogenic differentiation as described earlier (Ge et al., 2016). Briefly, cells cultured in proliferation media (DMEM/F12 (Thermo Fisher Scientific™, Cat#11320–033), 20% FBS,

1% Pen-Strep) for a couple of passages. After expansion, the cells were plated to confluency and changed to induction media (DMEM/F12, 10% FBS, 1% Pen-Strep, 1 μ M dexamethasone (MP Biomedicals, Cat#194561), 58 μ g/mL insulin (Human Insulin Recombinant Protein; Invitrogen, Cat#RP-10908), 0.5 mM 3-isobutyl-1-methylxanthine (IBMX; Sigma-Aldrich, Cat#I7018), 200 μ M indomethacin (Sigma-Aldrich, Cat#57413), and fed with fresh induction media every third day. Cells were then stained with Oil Red O (Sigma-Aldrich, Cat#O0625) to assess adipogenic potential after 1 week of differentiation.

Osteogenic Differentiation—Populations isolated by FACS were seeded and grown in vitro until 100% confluency, at which point the cell populations underwent osteogenic differentiation. Cells were fed with fresh osteogenic differentiation media (ODM) (MEM-alpha medium [Thermo Fisher, Scientific™, Cat#12561–056], 10% FBS, 1% Pen-Strep, 100 nM dexamethasone, 10 mM sodium β -glycerophosphate [Sigma-Aldrich, Cat#G9891], 2.5 mM ascorbic acid 2-phosphate [Sigma-Aldrich, Cat#A8960]) every 3 days for 10 days. Cells were then stained with Alizarin Red (Sigma-Aldrich, Cat#A5533) to assess osteogenic potential.

Chondrogenic differentiation—Cells were cultured as a micro-mass and differentiated using StemPro chondrogenesis differentiation kit (Thermo Fisher Scientific™, Cat#A1007101). Cells were resuspended at a cell-density of 1.6×10^7 cells/mL. A 5 μ L droplet of the cell suspension was seeded under high humidity conditions in a 24-well plate for 2 hours. After 2 hours, warmed chondrogenic differentiation media was added to the culture vessel. The growing micro-mass was fed with fresh chondrogenesis media every other day in a 37 °C incubator with 5% CO₂. At day 14 the micro mass was fixed, sectioned, and stained with Alcian blue (Sigma-Aldrich, Cat#A5268).

hSSC differentiation from induced pluripotent stem cell (iPSC)—Human iPSCs were plated onto Geltrex (Gibco, Cat#A1569601)-coated dishes in Essential 8 Medium (Gibco, Cat#A1007101) and cultured in an incubator maintained at 37 °C with 5% CO₂. iPSCs were fed daily with fresh media and maintained at 80% confluency. Cells were passaged 1:10 using 0.05 mM EDTA. To induce bone formation, iPSCs were differentiated when they reached 80% confluency by switching the cells over to ODM composed of MEM-alpha medium supplemented with 100 nM of dexamethasone, 10 mM β -Glycerophosphate, 2.5 mM ascorbic acid 2-phosphate, and 50 ng/mL of rhBMP2. Differentiating cells were fed with fresh ODM every other day and harvested on week 2 for FACS analysis.

HSC-stromal co-culture and transplant—To establish stromal monolayers for hHSC co-culture experiments, human stromal subsets, hSSCs, hBCSPs, and hOPs, were isolated by FACS and cultured as described under **Fluorescence-activated cell sorting (FACS)** and **Cell Culture** above. Human HSCs were enriched from freshly processed cord-blood samples by magnetic separation using CD34 microbeads (Miltenyi Biotec, Cat#130–056-701), according to the manufacturer's instructions. CD34+CD38-Lin-HSPC subpopulations (purity >95%) were isolated for co-culture with SSC-derived stromal populations. CD34+CD38-Lin-HSPC were sorted using a previously established panel that included anti-human lineage markers [CD2 (Cat#555328), CD3 (Cat#555341), CD4

(Cat#555348), CD7 (Cat#555362), CD8 (Cat#555368), CD11b (Cat#555389), CD14 (Cat#562692), CD16 (Cat#555408), CD19 (Cat#555414), CD20 (Cat#561761), CD56 (Cat#555517), all PE-Cy5; all dilution 1:50], CD235a(GPA)-PE-Cy5 (dilution 1:100; BD Biosciences, Cat#559944). Additionally, the following antibodies were used: CD38-PE-Cy7 (dilution 1:50; Cat#33579), CD90-FITC (dilution 1:25; Cat#555595), CD123-PE (dilution 1:10, Cat#554529), CD34-APC (dilution 1:25; Cat#659123)—all from BD Biosciences, CD10-APC-Cy7 (dilution 1:10; Cat#312212), and CD45RA-BV605 (dilution 1:20; Cat#304133)—both from Biolegend. Live cells assessed by negative staining for propidium iodide (PI, Sigma-Aldrich, Cat#P4170; final concentration: 1 μ g/ml) were double sorted. 5000 Lin-CD34+CD38-THY1^{lo}CD45RA- human cord blood HSCs were then co-cultured with or without confluent human stromal subsets for 14 days in Stemline serum-free hematopoietic stem cell expansion media (Sigma-Aldrich, Cat#S0192) supplemented with 10 ng/mL human recombinant stem cell factor (Peprotech, Cat#300-07), 5 ng/mL human recombinant thrombopoietin (Peprotech, Cat#300-18), 20 ng/mL human basic fibroblast growth factor (R&D Systems, Cat#P09038), and 25 ng/mL human insulin growth factor (R&D Systems, Cat#P05019). After 14-day co-culture, hHSCs were retro-orbitally injected into 6–8-week-old immunodeficient NSG mice conditioned using sub-lethal irradiation (250 rad in single dose) as described earlier (Reinisch et al., 2015). 8 weeks after transplant, bone marrow aspirates were obtained from recipient mice and analyzed by FACS for expression of human hematopoietic markers, HLA-ABC (pan-human; BioLegend, Cat#311404), hCD45 (pan-hematopoietic; BioLegend, Cat#103258), hCD19 (B lymphoid cells; BioLegend, Cat#302226), and hCD33 (myeloid cells; BioLegend, Cat#366626).

Lentiviral transduction of mSSCs—Fetal mSSCs were resuspended in MEM-alpha medium with 15% Fetal Calf Serum (FCS) and transduced individually with lentivirus carrying GFP, RFP, CFP, GFP/SOST, or GFP/DNAJB6 cDNA expression-cassettes. 48 hours after transduction, cells were sorted for GFP expression and transplanted as described above.

ATAC-seq—Cells for ATAC-seq were sorted as described above. The Fast-ATAC version of the ATAC-seq protocol was performed as described (Corces et al., 2016). Briefly, 10,000 cells per replicate (2 replicates per cell type) were resuspended in transposase reaction mixture (25 μ L of 2X Tagmentation DNA Buffer [TD buffer; Illumina, Cat#FC-121-1030], 2.5 μ L Tn5 enzyme [Illumina, Cat#FC-121-1030], 0.5 μ L 1% digitonin [Promega Cat#G9441], and 22 μ L water), incubated for 30 minutes at 37 °C. The tagged and transposed DNA fragments were purified using MinElute Reaction Cleanup kit (QIAGEN, Cat#28204) and eluted in 11 μ L elution buffer (from the MinElute kit) and amplified as described (Buenrostro et al., 2013). Libraries were sequenced to a depth of 12–23 million reads per replicate.

QUANTIFICATION AND STATISTICAL ANALYSIS

Fluorescence-activated cell sorting (FACS)—Flow cytometry was performed on FACS Aria II (BD Biosciences). Gating schemes were established with fluorescence-minus-one (FMO: staining with all fluorophores except one) controls and negative PI staining (1 μ g/ml) was used as a measure for cell viability. Cells from the human xenograft model and

from the in vivo skeletal-induction assays were also analyzed by FACS. FlowJo was used to analyze FACS data.

Transcriptional Expression Profiling—Raw microarray data were submitted to Gene Expression Commons (<http://gexc.riken.jp>) (Seita et al., 2012), where data normalization was computed against the Common Reference, which is a large collection of more than 11,939 mouse and 25,229 human array experiments deposited to the National Institutes of Health Gene Expression Omnibus (NIH GEO) database. GEXC assigns a threshold value to each probeset using the StepMiner algorithm (Sahoo et al., 2007) and calculates a percentile value between -100% and +100% for each available gene, allowing us to compare mouse and human gene expression on a normalized, continuous scale. Meta-analysis of the Common Reference also provides the dynamic range of each probe set on the array, and, in situations where there are multiple probe sets for the same gene, the probe set with the widest dynamic range was used for analysis. The Affymetrix Mouse Genome 430 2.0 Array includes 45,101 probe sets, of which 17,872 annotated genes are measurable. A list of mSSC/mBCSP-specific genes was generated by using the pattern search feature of the Gene Expression Commons (GEXC) database (Seita et al., 2012) for transcripts expressed in SSCs and BCSPs but not in their downstream progeny (**Supplemental Table 1**). The biomaRt R package (Durinck et al., 2009) was used to determine human orthologs to the mSSC/mBCSP-specific genes. Single cell RNA-seq data from each growth plate zone and diaphyseal region were then queried for expression of these human orthologs and zero-mean normalized by subtracting the mean gene expression. Percent positive gene enrichment was calculated by dividing the total number of positively-expressed genes by the total number of genes queried for each cell. Heatmap in Figure 1E was generated by plotting the most variable genes across single cells in the growth plate zones and diaphyseal region.

Scatter plots comparing mouse and human gene expression were plotted on ggplot2 platform in R (The R Foundation). Normalized gene expression values from probe sets with the widest dynamic range were plotted for a curated list of canonical signaling and bone development genes. Genes were broken into four quadrant groups: Q1: ↓Mouse ↑Human, Q2: ↑Mouse ↑Human, Q3: ↓Mouse ↓Human, Q4: ↑Mouse ↓Human; percent of each pathway represented in calculated as: $\frac{\text{No of pathway genes in quadrant}}{\text{Total number of pathway genes}}$. Heatmaps were generated using normalized expression values from GEXC (Seita et al., 2012).

Single-cell RNA-seq

Data processing: bcl2fastq2 2.18 (Illumina) was used to extract the data and generate FASTQ files for each single cell by using unique barcode combinations from the Nextera XT preparation. Raw reads were trimmed for base call quality (PHRED score ≥ 21) and for adapter sequences using Skewer (Jiang et al., 2014). Trimmed reads were then aligned to hg38 using STAR 2.4 (Dobin et al., 2013), and transcripts per million (TPM) was calculated using RSEM 1.2.21 (Li and Dewey, 2011). Raw TPM values were mean- and log₂-normalized. Cells with less than 200,000 uniquely mapped reads or fewer than 1,000 genes detected were excluded, leaving 1308 out of 1508 total cells for downstream analysis. Genes that were undetectable across all cells were also removed, leaving 21,414 genes out of total 26,856. Additional filtering criteria are described here for each figure.

Data analysis: Unsupervised principal component analysis (PCA) of single cells from fetal, adult, B-HAS-derived, and iPSC-derived hSSCs was completed using the `prcomp()` R function (R-Core-Team, 2017). Quality filtering prior to entering the matrix for PCA included (1) removal of cells with less than 2400 gene features, (2) removal of unexpressed genes and genes in which a single value occupied greater than 80% of the total sum. A total of 196 cells, including fetal (n=42), adult (n=36), B-HAS-derived (n=77), and iPSC-derived hSSCs (n=41) and 15,386 genes were analyzed.

Supervised analysis of our scRNA-seq data was guided by a gene set of canonical signaling pathways and bone development pathways created using Gene Expression Commons (GEXC) database (Seita et al., 2012) and the Molecular Signatures Database (MSigDB) (Liberzon et al., 2015). Quality filtering steps included (1) removal of cells with fewer than ~33% of gene features expressed and (2) removal of genes expressed in fewer than ~5% of cells. Gene expression values were \log_2 normalized. Total of 114 genes across 196 cells were analyzed. The same [cell x gene] matrix was used to generate a Pearson correlation matrix using the `ggcorrplot` package in R (Kassambara, 2016). For the bulk correlation matrix, mean gene expression was calculated for each gene per group (i.e. fetal, adult, B-HAS, iPSC) and analyzed using the `ggcorrplot` package in R by Pearson's correlation.

ATAC-seq analysis: ATAC-seq reads were trimmed using CutAdapt (Martin, 2011) and aligned to the human (hg19) or mouse (mm9) genome using Bowtie2 (Langmead and Salzberg, 2012). After filtering and removing duplicate reads, peaks were called using MACS2 as previously described, without a shifting model (Buenrostro et al., 2013). HOMER was used for motif analysis using default settings (Heinz et al., 2010). Promoter-proximal and promoter-distal peaks were separated based on distance to the nearest TSS (>2kb, distal. 2kb, proximal) using HOMER peak annotation. To identify peaks near SSC-specific, human-specific, or mouse-specific genes, we used the HOMER `annotatePeaks` tool to identify the nearest gene to a given peak. We then took the set of all peaks closest to an SSC-specific, human-specific, or mouse-specific gene and performed motif enrichment analysis. To find individual instances of RUNX2 motifs, we used the HOMER `annotatePeaks` tool with the "--motif" option specifying the RUNX2 consensus motif.

Statistical analysis—Bar chart data illustrating the analyses comparing SSCs in human fracture and xenografts are presented as the mean \pm SEM. Analyses were performed on GraphPad Prism. Unpaired, two-tailed Student's t-tests were used, with the assumption of unequal variances (Figure 5). Reproducibility was confirmed by repeating the analysis on multiple patients, n represents the number of patients.

DATA AND SOFTWARE AVAILABILITY

The ATAC-seq, RNA-seq, and Microarray data described in this study was deposited with the gene expression omnibus with the accession number SRA: PRJNA478935.

RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Mouse anti-human CD45-PB, Clone HI30 (1:50)	BioLegend	Cat#304029
Mouse anti-human CD235a-PB, Clone HIR2 (1:50)	BioLegend,	Cat#306612
Mouse anti-human CD31-biotin, Clone WM-59 (1:50)	Thermo Fisher Scientific™	Cat#13-0319-82
Mouse anti-human CD202b, biotin-linked, Clone 33.1 (1:50)	BioLegend	Cat#334204
Streptavidin, APC-AF750 (1:100)	Thermo Fisher Scientific™	Cat#SA1027
Mouse anti-human CD146-PE-Cy7, Clone SHM-57 (1:50)	BioLegend	Cat#342010
Mouse anti-human PDPN-APC, Clone NZ-1.3 (1:50)	Thermo Fisher Scientific™	Cat#17-9381-42
Mouse anti-human THY1-PE, Clone 5E10, (1:50)	BioLegend	Cat#328110
Mouse anti-human CD164-PE, Clone X, (1:50)	BioLegend	Cat#324808
Mouse anti-human CD73-FITC, Clone Ad2, (1:50)	BioLegend	Cat#344016
Mouse anti-human CD2-PE-Cy5, Clone RPA-2.10, (1:50)	BD Biosciences	Cat#555328
Mouse anti-human CD3-PE-Cy5, Clone HIT3a, (1:50)	BD Biosciences	Cat#555341
Mouse anti-human CD4-PE-Cy5, Clone RPA-T4, (1:50)	BD Biosciences	Cat#555348
Mouse anti-human CD7-PE-Cy5, Clone M-T701, (1:50)	BD Biosciences	Cat#555362
Mouse anti-human CD8-PE-Cy5, Clone RPA-T8, (1:50)	BD Biosciences	Cat#555368
Mouse anti-human CD11b-PE-Cy5, Clone ICRF44, (d1:50)	BD Biosciences	Cat#555389
Mouse anti-human CD14-PE-Cy5, clone M ϕ P9, (1:50)	BD Biosciences	Cat#562692
Mouse anti-human CD16-PE-Cy5, Clone 3G8, (1:50)	BD Biosciences	Cat#555408
Mouse anti-human CD19-PE-Cy5, Clone HIB19, (1:50)	BD Biosciences	Cat#555414
Mouse anti-human CD20-PE-Cy5, Clone 2H7, (1:50)	BD Biosciences	Cat#561761
Mouse anti-human CD56-PE-Cy5, Clone B159, (1:50)	BD Biosciences	Cat#555517
Mouse anti-human CD38-PE-Cy7, Clone HB7, (1:50)	BD Biosciences	Cat#33579
Mouse anti-human CD90-FITC, Clone 5E10, (1:25)	BD Biosciences	Cat#555595
Mouse anti-human CD123-PE, Clone 7G3, (1:10)	BD Biosciences	Cat#554529
Mouse anti-human CD235a (GPA)-PE-Cy5, clone GAR2 (1:100)	BD Biosciences	Cat#559944
Mouse anti-human CD34-APC-R700, Clone 8G12, (1:25)	BD Biosciences	Cat#659123
Mouse anti-human CD10-APC-Cy7, Clone HI10a, (1:25)	BioLegend	Cat#312212
Mouse anti-human CD45RA-BV605, Clone HI100, (1:20)	BioLegend	Cat#304133
Mouse anti-human CD45R-PE/DAZZLE 594, Clone RA3-6B2, (1:100)	BioLegend	Cat#103258
Mouse anti-human HLA-A,B,C FITC, Clone W6/32(1:100)	BioLegend	Cat #311404
Mouse anti-human CD19-AF700, Clone HIB19, (1:100)	BioLegend	Cat#302226
Mouse anti-human CD33-AF647, Clone P67.6 (1:100)	BioLegend	Cat#366626
Purified rabbit anti-human CD164, (1:100)	Sigma-Aldrich	Cat#HPA010636
Purified anti-Perilipin A, (1:100)	Chemicon	Cat#AB10200

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Purified Adiponectin, (1:100)	Invitrogen	Cat#MA1-054
Purified mouse anti-Human Nuclear Antigen, [235-1] (1:100)	Abcam	Cat#ab191181
Purified rabbit polyclonal Collagen II, (1:100)	Abcam	Cat#ab34712
Purified mouse anti-Osteocalcin [OC4-30], (1:100)	Abcam	Cat#ab13418
Secondary goat anti-rabbit-AF-488 (1:500)	Molecular Probes	Cat#A11034
Secondary goat anti-rat AF488 (1:500)	Invitrogen	Cat#A11006
Secondary goat anti-mouse AF647 (1:500)	Invitrogen	Cat#A21236
Secondary donkey anti-rabbit AF647 (1:500)	Molecular Probes	Cat#A31573
Anti-human CD45 microbeads (1:100)	Miltenyi Biotec	Cat#130-045-801
Anti-human CD34 microbeads (1:100)	Miltenyi Biotec	Cat#130-056-701
Bacterial and Virus Strains		
Lentivirus carrying GFP, RFP, and CFP.	Subcloned in the Weissman laboratory from pLL3.7 (Rubinson et al., 2003)	Addgene#11795
Chemicals, Peptides, and Recombinant Proteins		
DAPI	Biologend	Cat#422801
DiD lipophilic dye-AF647	Life Technologies	Cat#V22887
propidium iodide (PI)	Sigma-Aldrich	Cat#P4170
Matrigel	Corning	Cat#CB40234A
paraformaldehyde (PFA)	Electron Microscopy Sciences	Cat#15710
Bovine serum albumin (BSA)	Sigma-Aldrich	Cat#A9647
Donkey serum	Sigma-Aldrich	Cat#D9663
Goat Serum	Thermo Fisher Scientific™	Cat#G9023
Tris Buffered Saline (TBS), 10× solution	Sigma-Aldrich	Cat#T5912
Phosphate Buffered Saline (PBS)	GIBCO	Cat#C14190500BT
Fluoromount-G	Invitrogen	Cat#00-4958-02
rhBMP2 (recombinant human BMP2)	R&D Systems	Cat#355-BM/CF
rhVEGFR1 (recombinant human VEGFR1)	R&D Systems	Cat#321-FL/CF
Recombinant RNase inhibitor	Clontech	Cat#2313B
Triton X-100 10%	Thermo Fisher Scientific™	Cat#85111
MMLV SMARTScribe reverse transcriptase	Clontech	Cat#639538
KAPA HiFi hotStart ReadyMix	Kapa Biosystems	Cat#KK2602
Saffron	Sigma-Aldrich	Cat#S8381-5G
Acid Red 73	Sigma-Aldrich	Cat#49823-25MG
Alcian Blue 8GX	Sigma-Aldrich	Cat#A3157-25G
Acid Fuchsin	Sigma-Aldrich	Cat#F8129-25G
Phosphotungstic Acid	Sigma-Aldrich	Cat#P4006
Hematoxylin	Sigma-Aldrich	Cat#MHS32-1L
Eosin Y	Thermo Fisher Scientific™ Shandon™	Cat#6766009

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Calcein-AM	Invitrogen	Cat#C1430
Hoescht	Thermo Fisher Scientific™	Cat#62249
Type II collagenase	Sigma-Aldrich	Cat#C6885
100 U/mL DNase I	Worthington	Cat#NC9199796
Fetal bovine serum (FBS)	Fisher Scientific	Cat#16000-069
Histopaque of density 1.119 g/mL	Sigma-Aldrich	Cat#11191
Alizarin Red stain	Sigma-Aldrich	Cat#A5533
TRIzol LS	Invitrogen	Cat#10296028
Ethylenediaminetetraacetic acid (EDTA)	Invitrogen	Cat#15573-038
Optimal Cutting Temperature compound (OCT)	Fisher Scientific	Cat#23-730-571
Human platelet derived lysate	Stem Cell Technologies	Cat#06960
MEM alpha medium	Thermo Fisher Scientific™	Cat#12561-056
Penicillin-Streptomycin Solution	Thermo Fisher Scientific™	Cat#15140-122
0.1% Gelatin Solution	Millipore	Cat#ES-006-B
Media 199 (M199)	Sigma-Aldrich	Cat#C6885
Dulbecco's Modified Eagle Medium (DMEM)/F12	Thermo Fisher Scientific™	Cat#11320-033
Dexamethasone	MP Biomedicals	Cat#194561
Sodium β-glycerophosphate	Sigma-Aldrich	Cat#G9891
Ascorbic acid 2-phosphate	Sigma-Aldrich	Cat#A8960
Human Insulin Recombinant Protein	Invitrogen	Cat#RP-10908
3-isobutyl-1-methylxanthine (IBMX)	Sigma-Aldrich	Cat#I7018
Indomethacin	Sigma-Aldrich	Cat#57413
Oil Red O	Sigma-Aldrich	Cat#O0625
Alcian blue stain	Sigma-Aldrich	Cat#A5268
Geltrex	Gibco	Cat#A1569601
Essential 8 Medium	Gibco	Cat#A1007101
Human recombinant stem cell factor	Peptotech	Cat#300-07
Human recombinant thrombopoietin	Peptotech	Cat#300-18
Human basic fibroblast growth factor	R&D Systems	Cat#P09038
Human ng/mL insulin growth factor	R&D Systems	Cat#P05019
Stemline serum-free hematopoietic stem cell expansion media	Sigma-Aldrich	Cat#S0192
2× TD Buffer (Tagmentation DNA Buffer)	Illumina	Cat#FC-121-1030
Tn5 enzyme	Illumina	Cat#FC-121-1030
Digitonin	Promega	Cat#G9441
Agencourt AMPure XP beads	BEECKMAN COULTER	Cat#A63882
UltraPure™ DNase/RNase-Free Distilled Water	Thermo Fisher Scientific™	Cat#10977023
Critical Commercial Assays		
Nextera XT DNA Library Preparation kit	Illumina	Cat#FC-131-1096

REAGENT or RESOURCE	SOURCE	IDENTIFIER
High Sensitivity DNA analysis kit	Advanced Analytics, Stanford SFGF facility	N/A
Qubit dsDNA HS Assay kits	Thermo Fisher Scientific™	Cat#Q32851
M.O.M. immunodetection kit	Vector Laboratories	Cat#BMK-2202
RNeasy Micro Kit	Qiagen	Cat#74004
Arcturus™ RiboAmp™ PLUS Kit	Applied Biosystems™	Cat#KIT0521
StemPro chondrogenesis differentiation kit	Thermo Fisher Scientific™	Cat#A1007101
QIAGEN MinElute Reaction Cleanup kit	QIAGEN	Cat#28204
Deposited Data		
Microarray data, ATAC-seq data, sc-RNA-seq data	This paper	PRJNA478935
Experimental Models: Cell Lines		
Human: Human monocyte derived iPSC line SCVI 113	A gift from the laboratory of Joseph C. Wu	N/A
Experimental Models: Organisms/Strains		
Mouse: BA (C57BL/Ka-Thy1.1-CD45.1)	The Jackson Laboratory	JAX: 000406
Mouse: NSG (NOD.Cg-Prkdc ^{scid} Il2rg ^{tm1Wjl} /SzJ)	The Jackson Laboratory	JAX: 005557
Mouse: RFP (B6.Cg-Tg(CAG-mRFP1)1F1Hadj/J)	The Jackson Laboratory	JAX: 005884
Human: human fetal samples	Stemexpress	N/A
Oligonucleotides		
Oligo-dT30VN (5′ – AAGCAGTGGTATCAACGCAGAGTACATG-3′)	Invitrogen	N/A
template-switching oligonucleotide (TSO) (5′-AAGCAGTGGTATCAACGCAGAGTACATrGrG+G-3′)	Exiqon	N/A
ISPCR primers (5′-AAGCAGTGGTATCAACGCAGAGT-3′)	IDT	N/A
dNTP	Invitrogen	Cat#10297–018
ERCC (External RNA Controls Consortium) ExFold RNA Spike-In Mixes	Invitrogen	Cat#4456739
Recombinant DNA		
cDNA expression cassettes for <i>SOST</i>	Genecopeia	Cat#Mm11378
cDNA expression cassettes for <i>DNAJB6</i>	Genecopeia	Cat#Mm21356
Software and Algorithms		
Leica DMI6000B inverted microscope system running Leica Application Suite (LAS)-X software	Leica Microsystems	NA
ImageJ	NIH	http://rsb.info.nih.gov/ij/
GraphPad Prism 6.0	GraphPad Software	https://www.graphpad.com/scientific-software/prism/
Bioanalyzer	Agilent	https://www.genomics.agilent.com/en/Bioanalyzer-
Qubit 2.0 Fluorometer runs specific Invitrogen Qubit quantitation assays	Invitrogen	REF. Q32866
NextSeq 500; The NextSeq System Suite software; NCS v2.0	Illumina	ftp://webdata2@ussdftp.illumina.com/
HiSeq 4000	Illumina	ftp://webdata2@ussd-ftp.illumina.com/
FlowJo	FLOWJO LLC	https://www.flowjo.com/
bcl2fastq2 2.18	Illumina	https://support.illumina.com/sequencing/sequencing-software/bcl2fastq2-2.18.html
Skewer	Jiang et al., 2014	https://sourceforge.net/projects/skewer

REAGENT or RESOURCE	SOURCE	IDENTIFIER
STAR 2.4	Dobin et al. 2013	https://github.com/alexdobin/STAR
RSEM 1.2.21	Li and Dewey, 2011	https://deweylab.github.io/RSEM/
Gene Expression Commons (GEXC) database	Seita et al. 2012	http://gexc.riken.jp
The R package for Statistical Computing	The R Foundation	https://www.rproject.org/
BD FACSAria II	BD Biosciences	http://www.bdbiosciences.com/cn/home
Molecular Signatures Database	MSigDB	http://software.broadinstitute.org/gsea/msigdb
Gene Chip Scanner 3000 running GCOS 1.1.1 software	Affymetrix	http://www.affymetrix.com/products/software/speci
CutAdapt (Martin, 2011)	Martin, 2011	http://code.google.com/p/cutadapt/
Bowtie2	Langmead and Salzberg, 2012	http://bowtie-bio.sourceforge.net/bowtie2/index.shtml
MACS2 v. 2.0.10, pileup	Buenrostro et al 2013	http://liulab.dfci.harvard.edu/MACS/
HOMER	Heinz et al., 2010	Homer.ucsd.edu/homer/
Other		
Mouse Affymetrix 430–2.0 array	Affymetrix, Applied Biosystems™	Cat#900495
Human Affymetrix HG-U133+ array	Affymetrix, Applied Biosystems™	Cat#901569

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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- PDPN+CD146-CD73+CD164+ marks a self-renewing, multipotent human skeletal stem cell
- hSSCs can be isolated from fetal, adult, BMP2-treated human adipose stroma, and iPSCs
- hSSCs undergo local expansion in response to acute skeletal injury
- Comparison of mouse and human SSCs reveals evolutionary differences in skeletogenesis

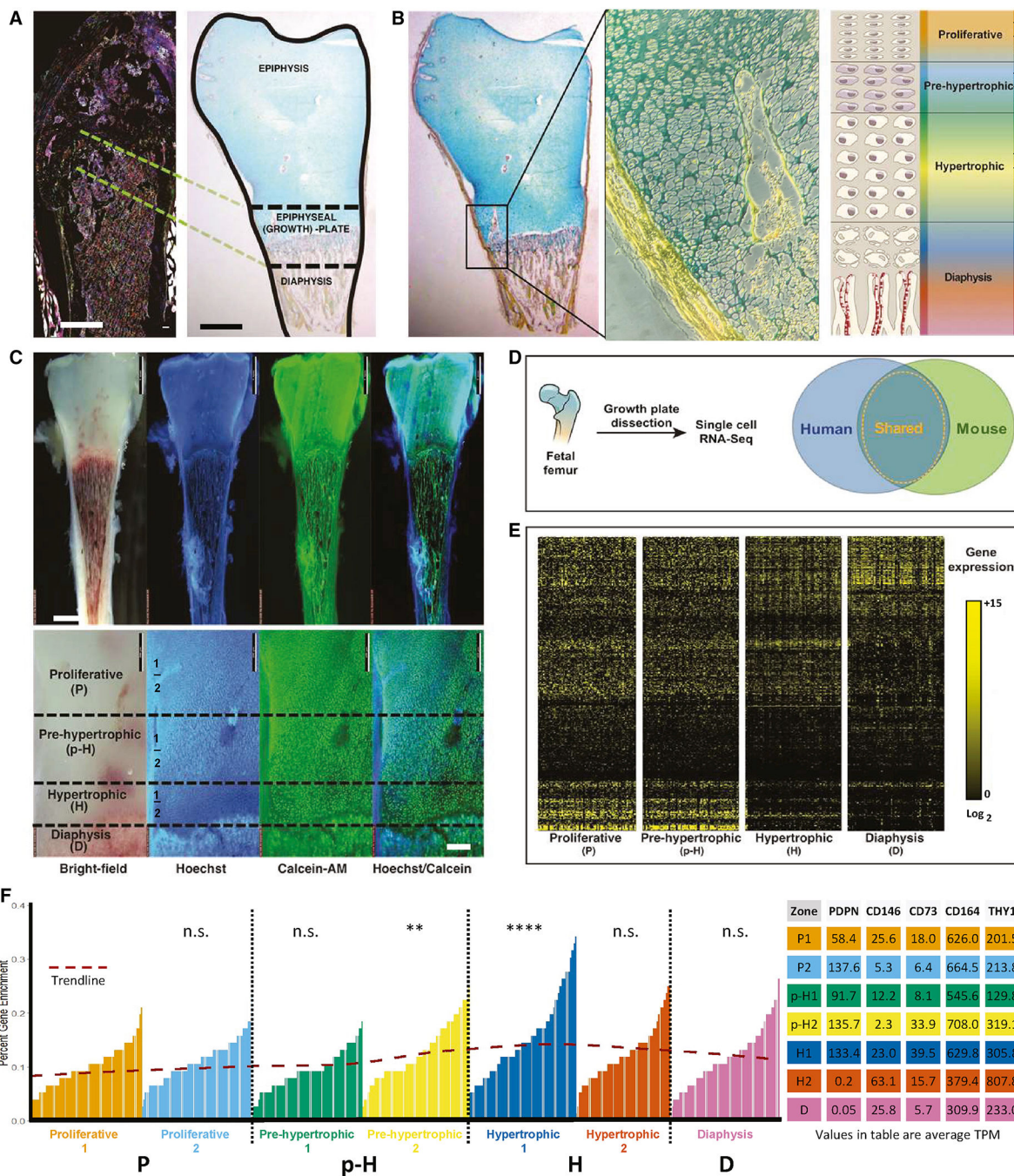


Figure 1. Single-cell analysis of skeletal growth plate zones.

A) A representative cross-section of a femur from an adult (8–12 weeks-old) Actin-Cre^{ER} X Rainbow mouse showing sites of clonal skeletogenesis in the growth plate (left). Scale: 500 μm. Homologous regions are shown in cross-section of a femoral head from a 17-week-old human fetus stained with Movat’s pentachrome (MP) (right). Scale: 2 mm.

B) Longitudinal cross-section of a femurhead from a 17-week-old human fetus stained with MP (left) alongside a magnified view of the growth plate (middle); yellow: bone, blue: cartilage, purple: marrow. Diagram of morphologically distinct zones in the growth plate of

17-week-old human fetal bone; P: proliferative zone, p-H: pre-hypertrophic zone, H: hypertrophic zone, D: diaphyseal zone (right). Scale: 100 μm .

C) Whole tissue stain of a viable 17-week-old human fetal femur with Hoechst and Calcein-AM intravital dyes (top; scale: 5 mm) revealing morphologically distinct regions (bottom; scale: 500 μm) corresponding to the six growth plate zones (P1, P2, p-H1, p-H2, H1, and H2) and the diaphyseal zone (D).

D) Experimental strategy for identifying human SSCs using scRNA-seq and a database of conserved SSC-specific genes.

E) Heatmap of most variable genes detected by scRNA-seq analysis of cells isolated from dissected human growth plate zones as shown in C.

F) Bar chart showing percent of human orthologs to mSSC/mBCSP-specific genes expressed in human CD45-CD235a- single cells for each zone described in C. Red trendline shows percent enrichment. Statistical significance for zones with highest enrichment for human orthologs to mSSC/BCSP-specific genes was calculated by one-way ANOVA with post-hoc Tukey HSD test with proliferative 1 (P1) as reference at an α -level of 0.01 (left). **p-value<0.005, ****p-value<0.0005; ns: not significant. Average transcripts per million (TPM) for five surface markers identified by scRNA-seq-enrichment-screen followed by empirical testing by FACS: PDPN, CD146 (MCAM), CD73 (NT5E), CD164, and THY1 (right). n=4.

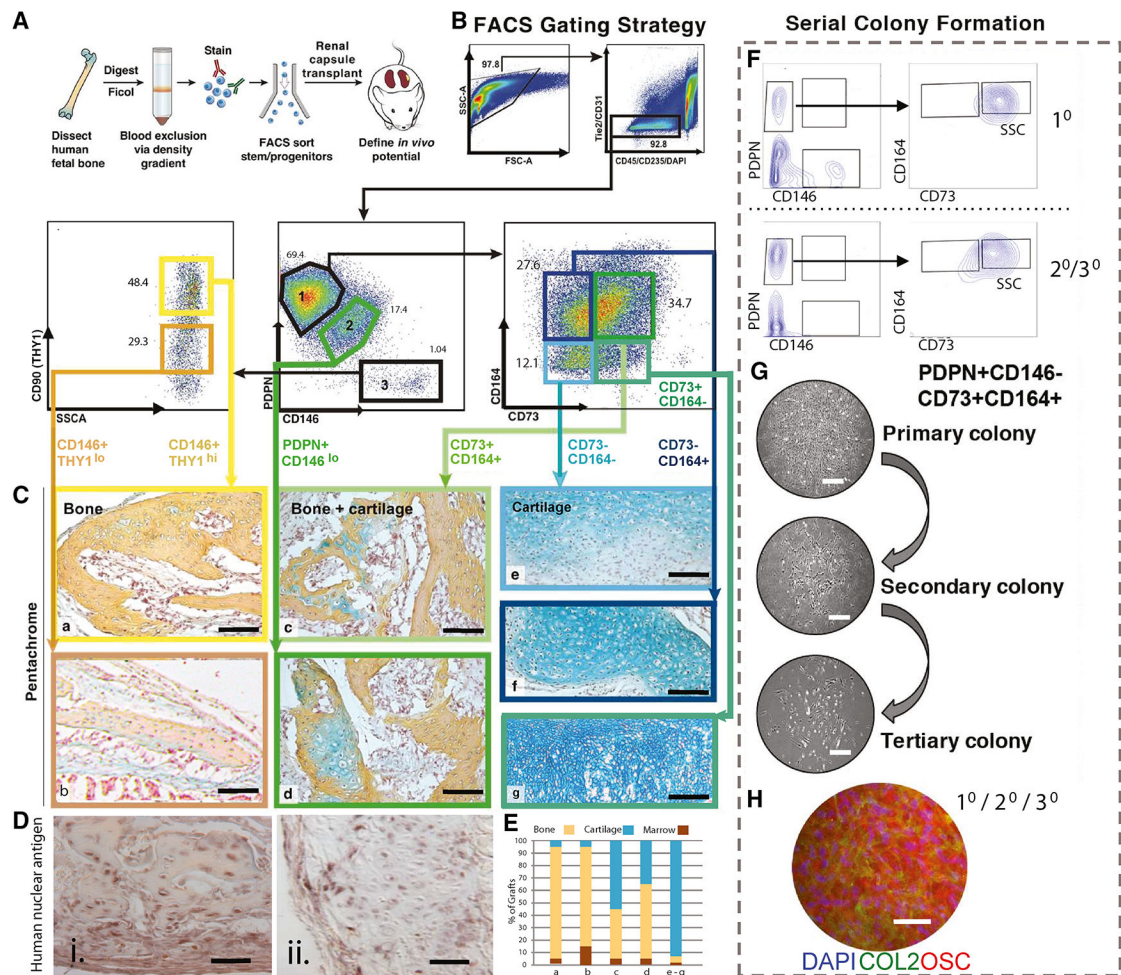


Figure 2. Prospective isolation of PDPN+CD146-CD73+CD164+ hSSCs.

A) Experimental strategy for isolation and in vivo functional characterization of hSSCs and progenitors by transplantation beneath the renal capsules of immunodeficient NSG mice. n=12.

B) Gating scheme for the isolation of distinct skeletal populations based on the varying expression of CD45, CD235, TIE2, CD31, PDPN, CD146, CD73, and CD164 using FACS.

C) MP stain of explanted human skeletal progenitor-derived grafts one month after subcapsular transplant into NSG mice; yellow: bone, blue: cartilage, purple: marrow. Stained cross-sections of the grafts that were derived from (a) PDPN-CD146+THY1+, (b) PDPN-CD146+THY1-, (c) PDPN+CD146-CD73+CD164+, (d) PDPN+CD146lo, (e) PDPN+CD146-CD73-CD164-, (f) PDPN+CD146-CD73-CD164+, and (g) PDPN+CD146-CD73+CD164- cells, respectively. Scale: 200 μ m.

D) Images showing immunohistochemical (IHC) staining for human nuclear antigen in (i) adjacent tissue cross-section as seen in C-a, and (ii) adjacent tissue cross-section as seen in C-c. Scale: 200 μ m.

E) Quantification of skeletal fates derived from populations C-a-f by morphometric analysis of histological specimens.

F) Representative FACS plots of primary colonies derived from a single PDPN+CD146-CD73+CD164+ cell (top), and secondary and tertiary colonies derived from a re-isolated single PDPN+CD146-CD73+CD164+ cell from the primary colony (bottom).

G) Phase contrast images showing primary, secondary, and tertiary PDPN+CD146-CD73+CD164+ colonies. Scale: 100 μ m.

H) IHC staining for nuclei with DAPI and cartilage and bone with anti-Collagen II (COL2), and anti-Osteocalcin (OSC) antibodies, respectively, in secondary colonies derived from a single, primary PDPN+CD146-CD73+CD164+ cell. Scale: 100 μ m; n=3.

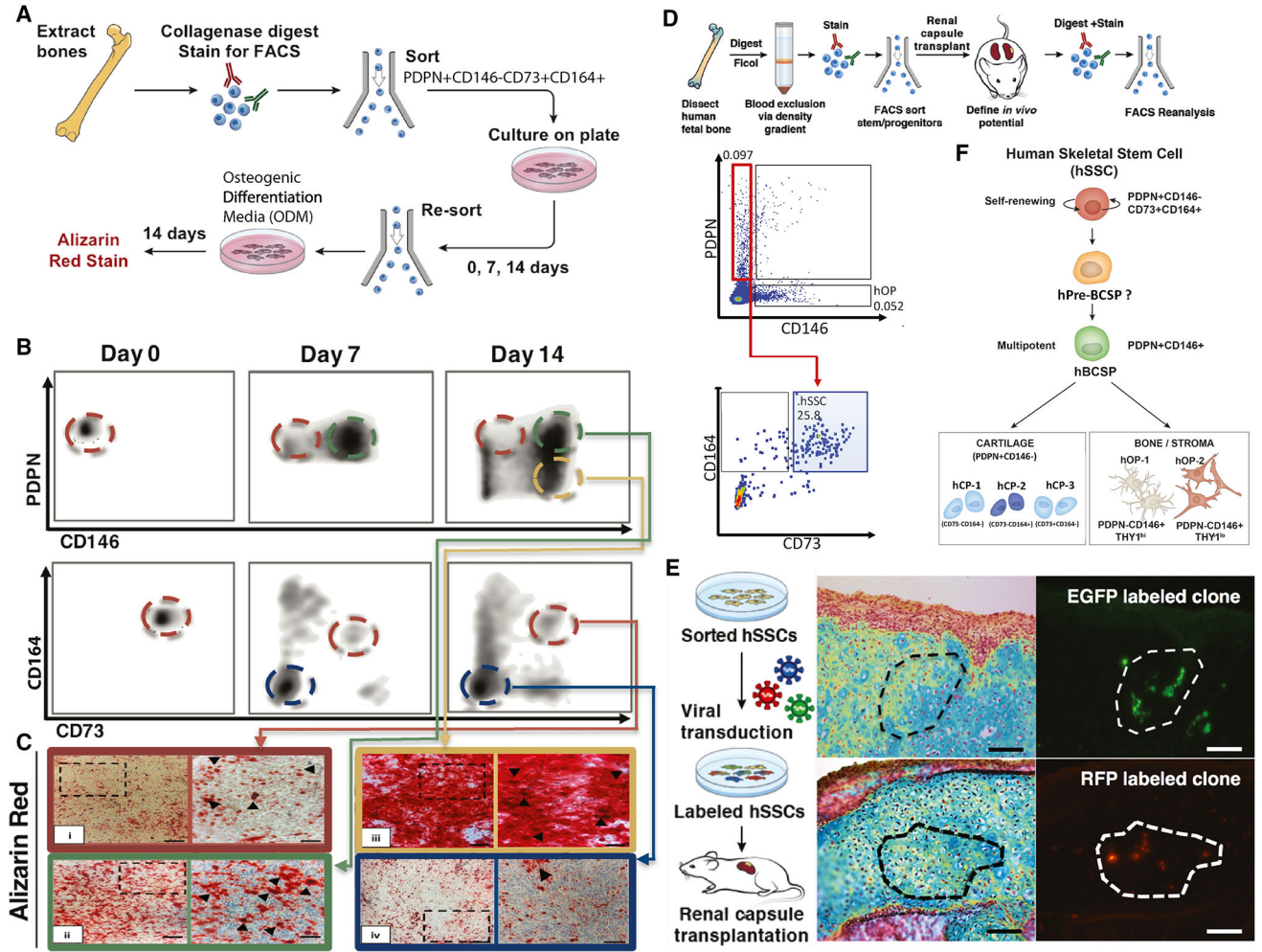


Figure 3. Determination of hSSC lineage hierarchy.

A) Experimental strategy for in vitro characterization of lineage progression of putative hSSCs (PDPN+CD146-CD73+CD164+) and other human skeletal progenitor subsets. $n=5$.

B) Representative FACS plots showing differentiation of cultured PDPN+CD146-CD73+CD164+hSSCs into lineage-restricted subsets from day 0, 7, and 14.

C) Alizarin Red S staining of individual human skeletal progenitor subsets to measure osteogenicity [i) hSSC: human skeletal stem cell; ii) hBCSP: human bone, cartilage, and stromal progenitor cell; iii) hOP: human osteogenic progenitor; iv) hCP: human chondrogenic progenitor]. Scale: 200 μm .

D) Experimental strategy for in vivo determination of hSSC lineage potential (top). Freshly isolated hSSCs were transplanted under the renal capsules of NSG mice, explanted after 4 weeks of development, dissociated to single-cell suspension and analyzed. FACS plot showing different skeletal populations within the dissociated primary graft including hSSC (blue box), hBCSP, hOP, and hCP all derived from originally transplanted fetal hSSC (bottom).

E) Experimental strategy for random red-green-blue (RFP, EGFP, CFP) lentiviral labeling and in vivo clonal expansion of the labeled hSSCs (left). MP-stained sections of kidney grafts derived from the labeled-hSSCs 4 weeks after transplantation (middle). Fluorescent

images of the adjacent sections of the same grafts showing clonal expansion of single color skeletal cells derived from a single-color-labeled hSSC (right). Individual EGFP and RFP labeled clones are shown. Scale: 100 μm .

F) Lineage map of hSSC and downstream skeletal progenitors. h-pre-BCSP: pre-bone, cartilage and stromal progenitor cell.

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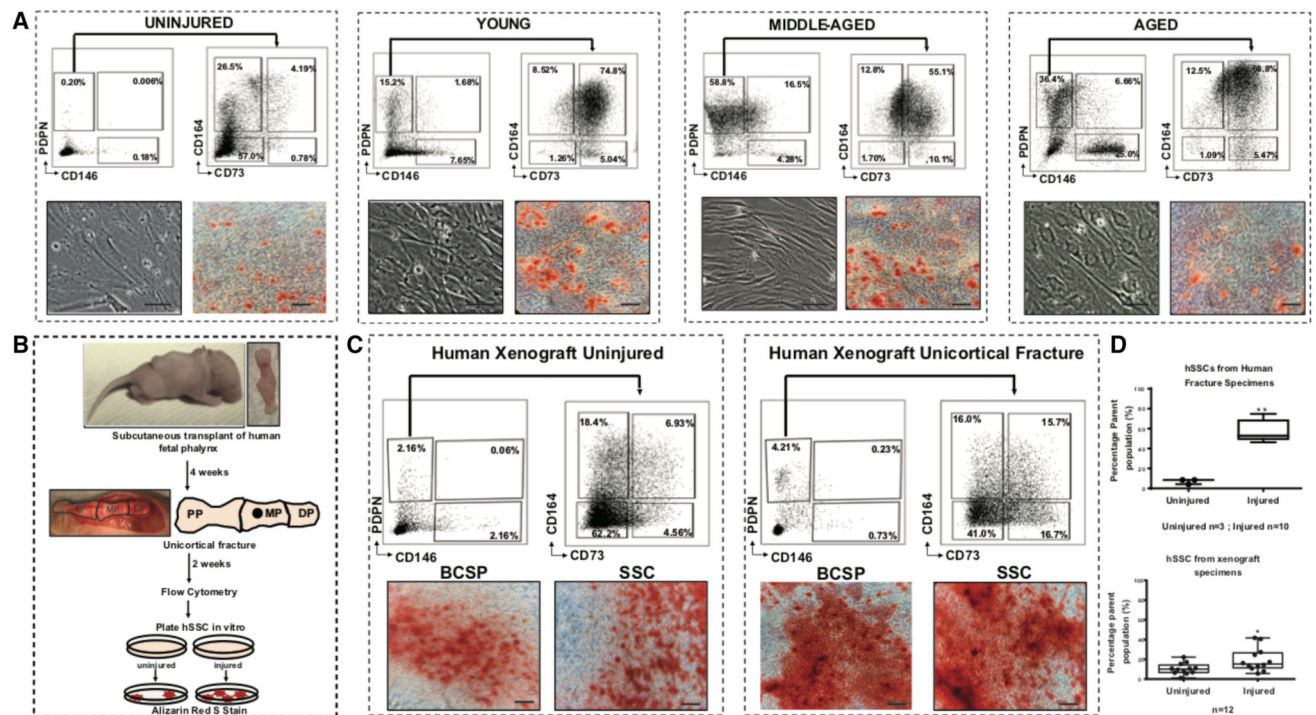


Figure 4. Osteogenic hSSCs are locally amplified in response to skeletal injury.

A) Injured (n=10) and uninjured (n=3) human bone-specimens were obtained and hSSCs were isolated. Uninjured age group: 16–30 years-old (yo); injured age groups: young: 20–40 yo (n=3), middle-aged: 41–70 yo (n=3), and aged: 71–90 yo (n=4) (left-right). Cells were isolated by FACS (top left and right), plated, differentiated, and stained using Alizarin Red. Brightfield and corresponding Alizarin Red stained images are shown in bottom left and bottom right panels, respectively. Scale: 100 μ m.

B) Postnatal day 5 NSG mouse pup host (top left). Phalanx of 18-week-old human fetus (top right). Human fetal phalanges from 18-week-old fetuses were transplanted subcutaneously under the dorsum of day 5 NSG mouse pups. Fractures were introduced at 4 weeks post-transplantation (middle left and right panels) within the engrafted human phalanx (PP: proximal phalanx, MP: middle phalanx, DP: distal phalanx; defect indicated by black circle in MP region). After a further 2 weeks post-injury the phalanges were explanted for analysis (n=12). For both uninjured and injured samples, explants were digested, hSSC were purified by FACS, plated, differentiated for 2 weeks, and then stained using Alizarin Red (bottom).

C) Comparison of uninjured (left) and injured (right) human xenografts. Each panel shows the corresponding FACS plots with gating strategy to isolate hBCSPs and hSSCs (top). Purified hBCSPs and hSSCs were plated, differentiated for 2 weeks, and stained using Alizarin Red (bottom). Scale: 100 μ m.

D) Bar chart showing the effect of injury on the hSSC populations in both the primary human samples (top) (uninjured, n=3; injured, n=10) and the human xenograft model (n=12) (bottom).

<0.05, **p<0.01.

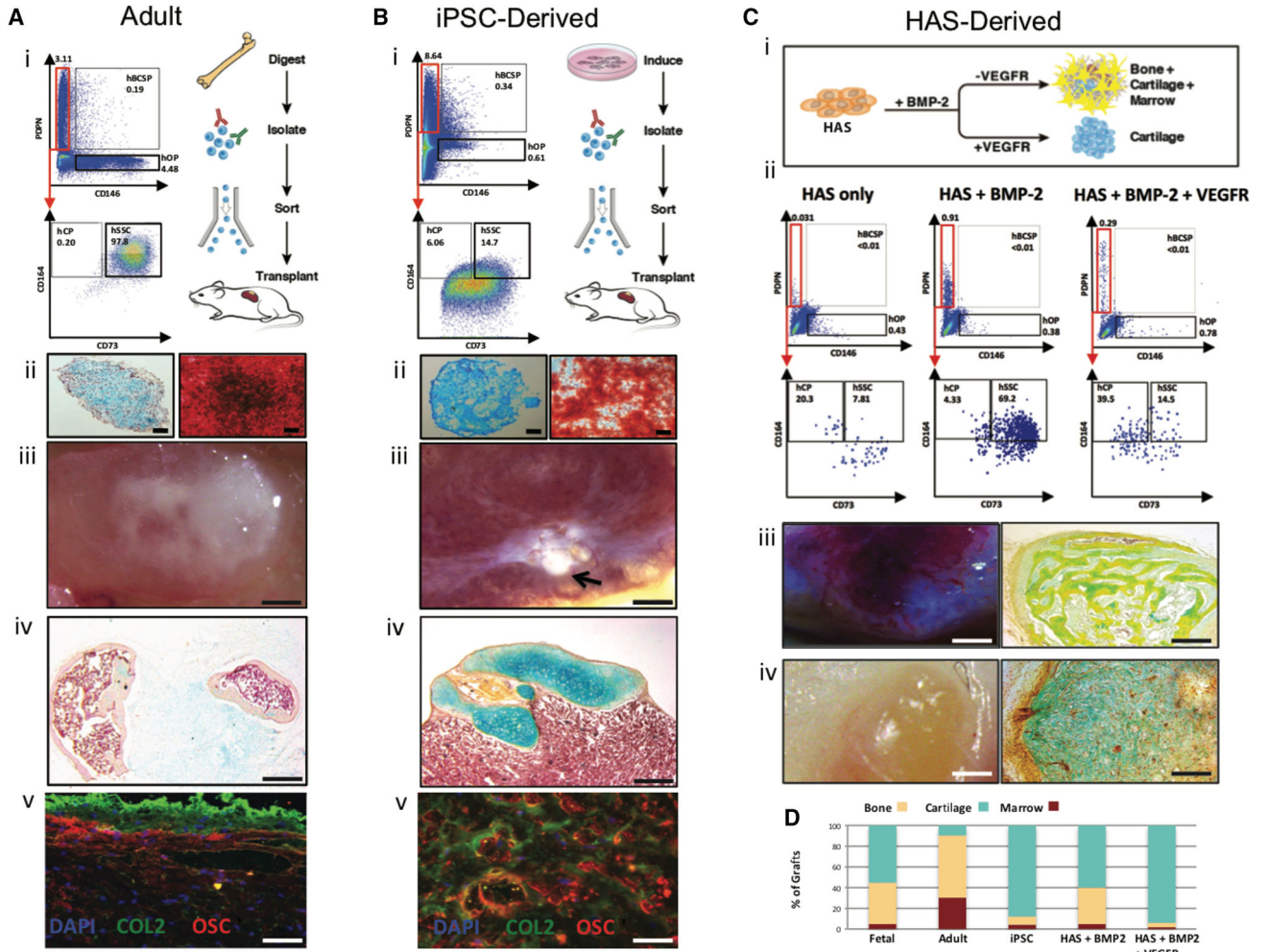


Figure 5. PDPN+146-CD73+CD164+ expression identifies hSSC from other tissue sources.

A) Isolation of hSSCs from adult human femoral tissue.

(i) Experimental strategy (right) and representative FACS plots (left) showing gating scheme for isolation of hSSCs from adult human femurhead tissue (n=6).

(ii) Alcian blue stain showing cartilage tissue in a cross-section of micro-mass generated by adult hSSCs after differentiation in vitro (left). Scale: 500 μ m. Alizarin Red stain showing bone tissue (osteoblasts) generated by adult hSSCs after differentiation in vitro (right). Scale: 100 μ m.

(iii) Image of ossicle formed 4 weeks after transplant of 100,000 hSSCs isolated in A-(i) under the renal capsule of NSG mice. Scale: 2 mm.

(iv) MP stain of a cross-section of the ossicle shown in A-(iii). Scale: 500 μ m.

(v) IHC staining of a cross-section of the ossicle shown in A-(iii) for nuclei with DAPI and cartilage, and bone with anti-Collagen II (COL2), and anti-Osteocalcin (OSC) antibodies, respectively. Scale: 100 μ m; n=6.

B) Isolation of hSSCs after skeletal induction of human monocyte-derived iPSCs.

(i) Experimental strategy (right) and representative FACS plots (left) showing gating strategy for the isolation of hSSCs 2 weeks after skeletal induction of the iPSCs.

(ii) Alcian blue stain showing cartilage in a cross-section of micro-mass generated 2 weeks after differentiation of iPSC-derived hSSCs in vitro (left). Scale: 500 μ m. Alizarin Red S stain showing bone (osteoblasts) generated 2 weeks after differentiation of iPSC-derived hSSCs in vitro (right). Scale: 100 μ m.

(iii) Image of an ossicle formed 4 weeks after transplant of 100,000 hSSCs purified from skeletal induced iPSC cultures and then transplanted under the renal capsule of NSG mice. Scale: 2 mm.

(iv) MP stain of a cross-section of the ossicle shown in B-(iii). Scale: 200 μ m.

(v) IHC stained image of a cross-section of the ossicle in B-(iii) for nuclei with DAPI and cartilage, and bone with anti-Collagen II (COL2), and anti-Osteocalcin (OSC) antibodies, respectively. Scale: 100 μ m; n=6.

C) Isolation of hSSCs from BMP2-treated human adipose stroma (B-HAS).

(i) Experimental strategy for human adipose stroma (HAS) induction with either BMP2 alone or with co-delivery of BMP2 and sVEGFR.

(ii) FACS analysis of HAS (left), HAS-derived ossicle after induction with BMP2 (middle), and HAS-derived cartilage after induction with BMP2 and sVEGFR (right).

(iii) Image of a vascularized ossicle generated 4 weeks after subcutaneous transplantation of HAS treated with BMP2 in NSG mice (left). Scale: 2 mm. MP stain of a cross-section of the same ossicle (right). Scale: 200 μ m.

(iv) Image of cartilage mass formation lacking vascularization generated 4 weeks after subcutaneous transplantation of HAS treated with BMP2 and sVEGFR in NSG mice (left). Scale: 2 mm. MP stain of a cross-section of the same subcutaneous cartilaginous mass (right). Scale: 200 μ m; n=6.

D) Bar chart showing quantification of percent contribution by the different skeletal fates to total graft mass for grafts derived from fetal hSSCs, adult hSSCs, skeletal-induced-iPSC, B-HAS, or BMP2+sVEGFR-treated HAS after morphometric analysis.

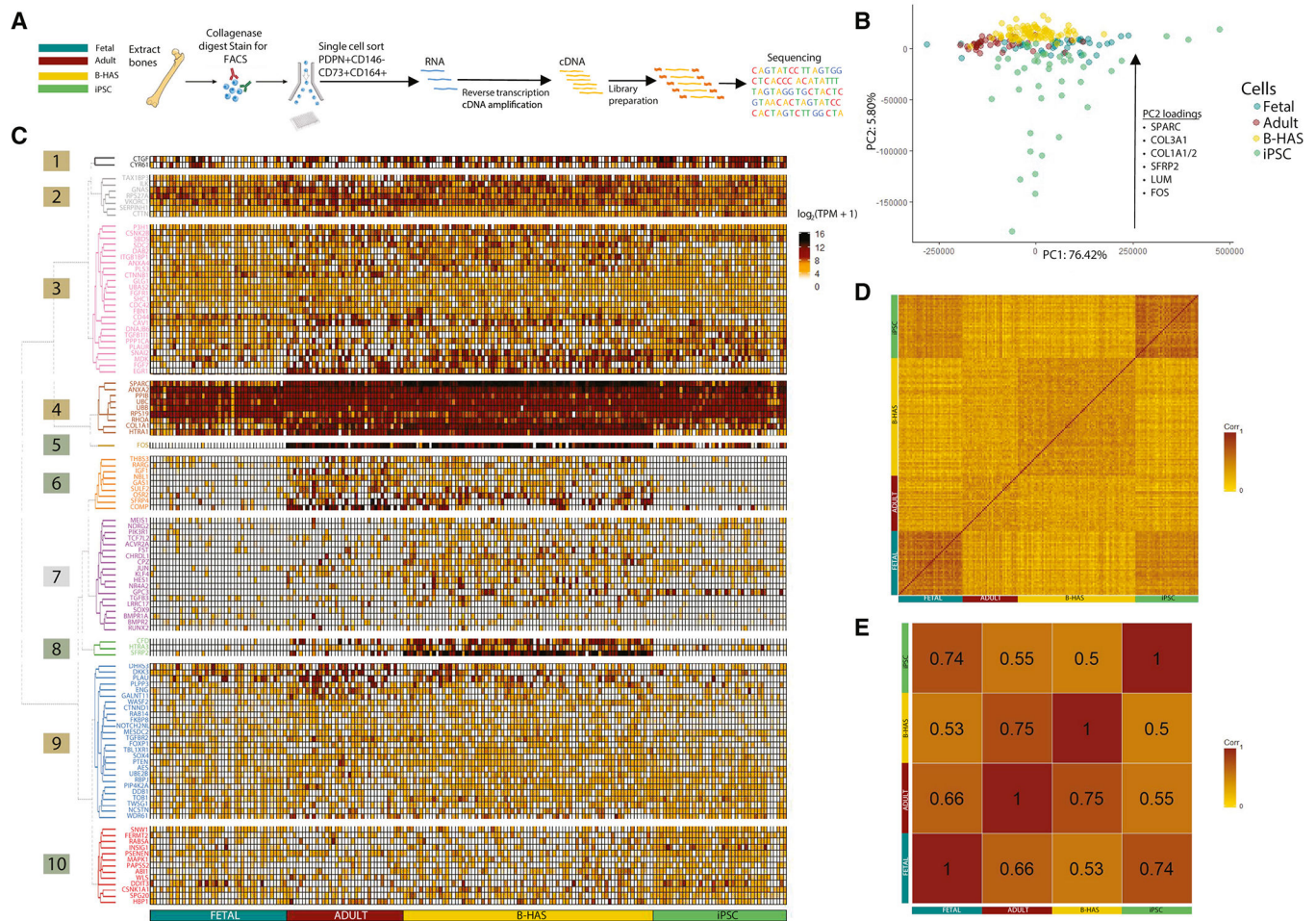


Figure 6. Comparative scRNA-seq analysis of hSSCs derived from different sources.

A) Experimental strategy for scRNA-seq of fetal (blue), adult (brown), B-HAS-derived (yellow), and iPSC-derived (green) hSSCs.

B) Unsupervised PCA of single hSSCs based on their expressed transcriptomes across principal component (PC) 1 and 2. Top gene loading contributors to the PC2 are listed along with the general direction of their eigenvectors. Colors of dots correspond to hSSC-source as in A.

C) Heatmap showing gene expression ($\log_2(\text{TPM}+1)$) of known genes involved in canonical signaling and/or bone development pathways (Supplemental Table 3) in single fetal (n=42), adult (n=36), B-HAS-derived (n=77), and iPSC-derived hSSCs (n=41). The heatmap is grouped into 10 clusters obtained based on the expression patterns of the selected genes using Ward's method.

D) Pearson correlation matrix showing similarities between single fetal, adult, B-HAS-derived, and iPSC-derived hSSCs based on gene expression values as in B. Scale shows color gradient for R^2 value from 0 to 1.

E) Pearson correlation matrix with numerical coefficients showing degree of similarity between aggregate fetal, adult, B-HAS-derived, and iPSC-derived hSSC populations. Mean $\log_2(\text{TPM}+1)$ was calculated separately for each group by calculating the sum of the

$\log_2(\text{TPM}+1)$ values for each gene in B and dividing the sum by the total number of cells in each group.

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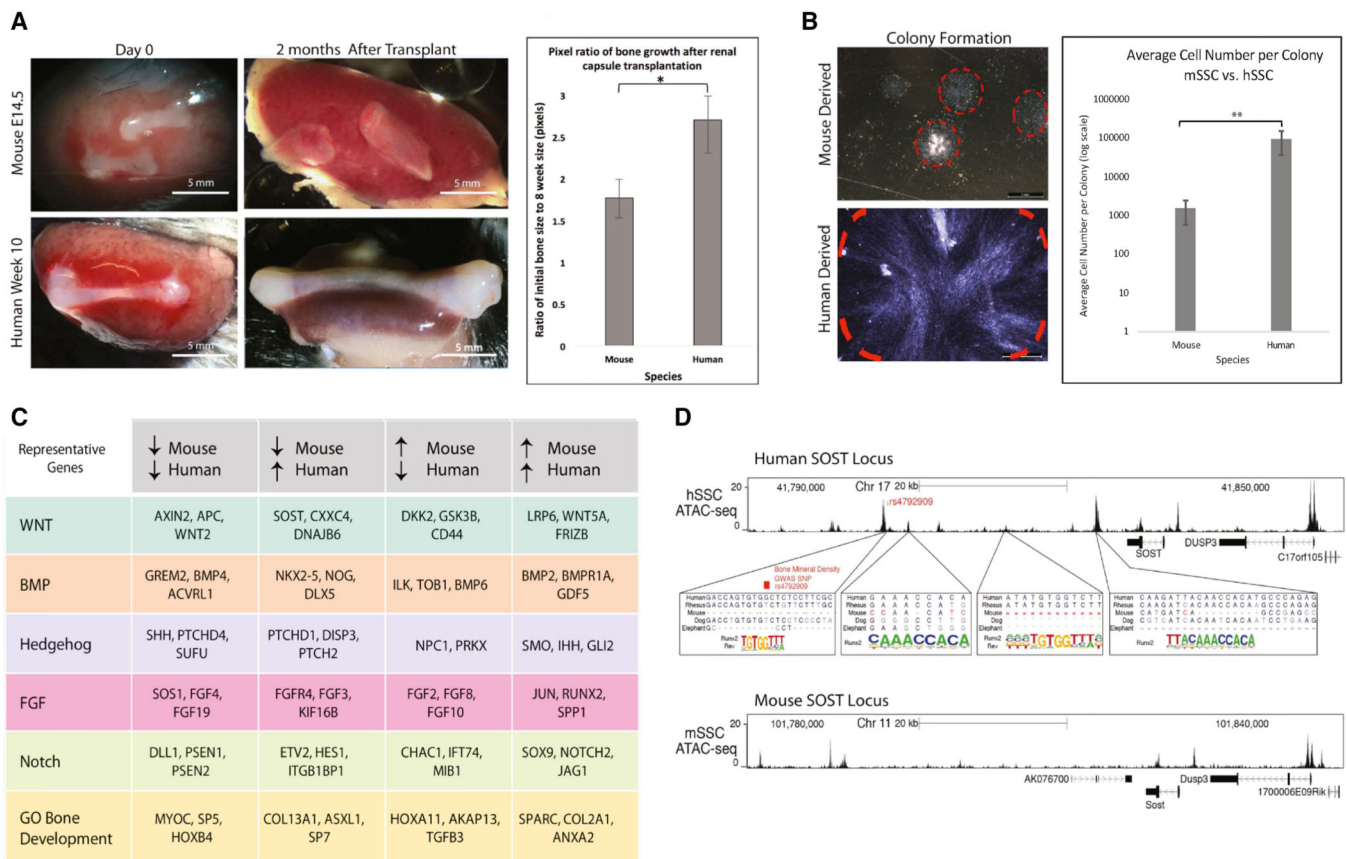


Figure 7. Convergent and divergent skeletal regulatory programs between mouse and human SSCs.

A) Renal capsule transplant of e14.5 mouse bones at day 0 (top left) and 10-week human fetal bones at day 0 (bottom left) in NSG mice. Images highlighting the resulting difference in size between the mouse (top middle) and human (bottom middle) bones after 2 months of growth in vivo. Scale: 5 mm; n=3. Pixels corresponding to the bone tissue in the images were measured and normalized to their respective pixel-counts at day 0 to calculate growth ratios, which are represented as bar chart (right). *p<0.05.

B) Brightfield image of CFUs (red dashed line) that were formed after 2 weeks of culture of either e14.5 fetal mSSCs (top) or 10-week fetal hSSCs (bottom) highlighting the difference in their sizes. Scale: 1 mm. Bar chart showing the average number of cells per hSSC-derived and mSSC-derived colonies (right). Error bars: standard deviation; **p<0.01; n=5.

C) Table listing shared and unshared representative genes and their expression trends (↑=upregulated, ↓=downregulated) between mouse and human focusing specifically on major signaling pathways and gene ontology (GO) gene set for bone development based on comparative microarray analysis of mSSCs (e17.5) and hSSCs (19-week) using the GEXC (n=3).

D) UCSC genome browser track showing the ATAC-seq signal at the *SOST* locus in 18-week fetal hSSCs (top). Regions within four ATAC-seq peaks that contain RUNX-binding motifs are magnified. Mammalian conservation track from UCSC genome browser is shown for human, rhesus, mouse, dog, and elephant. RUNX2-binding motif (forward or reverse)

found at each site is also shown (middle). Genome browser track showing ATAC-seq signal at the mouse *Sost* locus in e17.5 fetal mSSCs (bottom).

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