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Postnatal Osterix but not DMP1 lineage cells significantly contribute to intramembranous ossification in three preclinical models of bone injury

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Murine models of long-bone fracture, stress fracture, and cortical defect are used to discern the cellular and molecular mediators of intramembranous and endochondral bone healing. Previous work has shown that Osterix (Osx+) and Dentin Matrix Protein-1 (DMP1+) lineage cells and their progeny contribute to injury-induced woven bone formation during femoral fracture, ulnar stress fracture, and tibial cortical defect repair. However, the contribution of pre-existing versus newly-derived Osx+ and DMP1+ lineage cells in these murine models of bone injury is unclear. We addressed this knowledge gap by using male and female 12-week-old, tamoxifen-inducible Osx Cre_ERT2 and DMP1 Cre_ERT2 mice harboring the Ai9 TdTomato reporter allele. To trace pre-existing Osx+ and DMP1+ lineage cells, tamoxifen (TMX: 100 mg/kg gavage) was given in a pulse manner (three doses, 4 weeks before injury), while to label pre-existing and newly-derived lineage Osx+ and DMP1+ cells, TMX was first given 2 weeks before injury and continuously (twice weekly) throughout healing. TdTomato positive (TdT+) cell area and cell fraction were quantified from frozen histological sections of injured and uninjured contralateral samples at times corresponding with active woven bone formation in each model. We found that in uninjured cortical bone tissue, Osx Cre_ERT2 was more efficient than DMP1 Cre_ERT2 at labeling the periosteal and endosteal surfaces, as well as intracortical osteocytes. Pulse-labeling revealed that pre-existing Osx+ lineage and their progeny, but not pre-existing DMP1+ lineage cells and their progeny, significantly contributed to woven bone formation in all three injury models. In particular, these pre-existing Osx+ lineage cells mainly lined new woven bone surfaces and became embedded as osteocytes. In contrast, with continuous dosing, both Osx+ and DMP1+ lineage cells and their progeny contributed to intramembranous woven bone formation, with higher TdT+ tissue area and cell fraction in Osx+ lineage versus DMP1+ lineage calluses (femoral fracture and ulnar stress fracture). Similarly, Osx+ and DMP1+ lineage cells and their progeny significantly contributed to endochondral callus regions with continuous dosing only, with higher TdT+ chondrocyte fraction in Osx+ versus DMP1+ cell lineages. In summary, pre-existing Osx+ but not DMP1+...
Introduction

Bone is one of the only tissues in the body that can heal with scarless tissue regeneration. This remarkable capacity for self-repair requires a complex, multi-faceted process that involves growth factors, mechanical cues, and unique populations of cells. Based on these environmental factors, bone healing occurs either via endochondral or intramembranous ossification. In endochondral ossification, progenitor cells first differentiate and form a cartilage callus that is later replaced by bone. In contrast, intramembranous ossification results in direct bone formation from progenitor cells, bypassing the cartilage intermediate. Although still unclear, studies indicate that endochondral processes are favored in environments with low oxygen tension, vascular disruption, and some micromotion (non-rigid fixation) (Toyosawa et al., 2004; Marsell and Einhorn, 2011; Bahney et al., 2015; Miller et al., 2015). With nearly 5–10% of fractures progressing to delayed healing or non-union (Wooll and Pfleger, 2003; Einhorn and Gerstenfeld, 2015) and resulting in increased medical cost and loss of productivity (Bonafede et al., 2013), understanding the cellular and molecular mediators of both endochondral and intramembranous ossification following bone injury is paramount.

Preclinical models of bone injury are critical for dissecting the cellular and molecular processes controlling endochondral and intramembranous vascularization. The most common injury model used is the transverse, full fracture (“Einhorn model”) first developed by Bonnarens and Einhorn (Bonnarens and Einhorn, 1984). This model has been adapted for use in both the tibia and femur of rats and mice (Bonnarens and Einhorn, 1984; An et al., 1994; Zondervan et al., 2018; Buettmann et al., 2019), and utilizes blunt trauma to induce a mid-diaphyseal fracture that is stabilized with an intramedullary rod. Due to the semi-stable nature of fixation, this model heals by periosteal intramembranous woven bone formation near the callus periphery and endochondral ossification near the fracture site, with both woven bone tissue and cartilage visible by day 14 post-injury (Colnot, 2009; Buettmann et al., 2019). Tissue transplantation studies have determined that cells from the periosteum are the primary contributors to callus formation in this model, with smaller contributions from the adjacent skeletal muscle and marrow (Colnot, 2009; Julien et al., 2022). In contrast, the rodent stress fracture model, developed and characterized in our lab, utilizes forelimb cyclic fatigue loading to create a non-displaced ulnar fracture that heals predominantly by periosteal intramembranous woven bone formation 10–14 days post-injury (Hsieh and Silva, 2002; Uthgenannt et al., 2007; Wohl et al., 2009; Martinez et al., 2010). Bulk RNAseq analysis comparing the transverse, full fracture model versus stress fracture model in mice indicates that the stress fracture model has a shorter, less pronounced inflammatory phase and a more enriched osteogenic signature (Coates et al., 2019). Another widely used bone repair model is the monocortical defect injury. In this model, a small monocortical defect (0.4–0.8 mm in diameter) is drilled in the mid-diaphysis of the long-bone (Liu et al., 2018; Buettmann et al., 2019; Li and Helms, 2021). Healing progresses after injury with inflammation followed by small amounts of periosteal cartilage callus formation between days 3 and 7 (Hu and Olsen, 2016; Liu et al., 2018). By days 5–10 after injury, intramedullary intramembranous hard callus formation occurs, followed by resolution at days 14–21. Due to the differing healing modalities among these three bone injury models, their simultaneous utilization can provide insights into the unique cellular and molecular mediators of bone healing (Supplementary Figure S1).

Tracking the cellular mediators of bone healing has been aided by the recent development of many tamoxifen-inducible Cre constructs (Cre_ERT2) that can be crossed with fluorescent transgenic reporters (Ai9, Ai14, mTmG, YFP, etc.), allowing for longitudinal tracking of targeted cell populations that contribute to fracture healing in vivo (Feil et al., 2009; Madisen et al., 2010; Abe and Fujimori, 2013; Seine et al., 2015). The emerging role of different skeletal stem cells in fracture repair has been reported by numerous groups and was reviewed recently (Serowoky et al., 2020). We have focused on cells at the later stage of the osteoblast lineage (Osx and later), and used continuous tamoxifen dosing to demonstrate that Osx+ lineage cells and their progeny (labeled in OsterixCre_ERT2 (Maes et al., 2010) Ai9 (Madisen et al., 2010) mice) contributed greater cell numbers than DMP1+ lineage cells (labeled in Dentin Matrix-Protein 1 Cre_ERT2 (Powell et al., 2011) Ai9 (Madisen et al., 2010) mice) and their progeny to woven bone formation in femoral transverse, ulnar stress fracture, and tibial cortical defects (Buettmann et al., 2019). However, because cells were labeled before and during healing by continuous tamoxifen, we could not determine the contribution of pre-existing versus newly differentiated Osx+ and DMP1+ lineage cells and their progeny (herein labeled Osx+ or DMP1+ lineage cells) to fracture callus tissues. More
recent work by our lab group used pulse-chase labeling strategies and demonstrated that pre-existing Osx+ and DMP1+ lineage cells and their progeny contribute significantly to early lamellar bone formation following anabolic (non-damaging) skeletal loading (Zannit and Silva, 2019; Harris and Silva, 2022). Interestingly, we observed that these lineage-labeled cells, especially DMP1+ lineage cells, are rapidly depleted from the periosteal bone surface when a higher loading stimulus induces woven bone formation (Zannit and Silva, 2019; Harris and Silva, 2022). Together these data indicate that Osx+ and DMP1+ lineage cells play a role in load-induced bone formation and bone healing, however the relative contributions of pre-existing versus newly-derived Osx+ and DMP1+ lineage cells and their progeny across various bone injury types remains poorly defined.

Using both continuous and pulse-chase tamoxifen dosing strategies, we sought to determine the role of pre-existing and/or newly-differentiated Osx+ and DMP1+ lineage cells and their progeny in three pre-clinical models of bone repair: transverse femoral fracture, ulnar stress fracture, and tibial cortical defect (Supplementary Figure S1). Due to the wider resident bone cell population reported to be targeted with Osx Cre_ERT2 construct (Maes et al., 2010), we hypothesized that pre-existing Osx+ lineage cells target a greater portion of woven bone regions versus pre-existing DMP1+ lineage across all three injury models. Furthermore, we hypothesized that lineage-labeled cells in woven bone callus would be significantly increased with continuous dosing compared to pulse dosing in both Osx Cre_ERT2 Ai9 and DMP1 Cre_ERT2 Ai9 mice.

Methods

Mouse lines

All mouse breeding and experimental protocols were approved by Washington University in St. Louis IACUC. Mouse lines including Osx Cre_ERT2 (Maes et al., 2007), DMP1 Cre_ERT2 (Powell et al., 2011), and Ai9 (RCL-tdTomato) (Madisen et al., 2010) were previously generated and described. Osx Cre_ERT2 and DMP1 Cre_ERT2 breeders were shared from the laboratories of Drs. Henry Kronenberg and Paola Pajevic, respectively. Ai9 (RCL-tdT; Catalog #007909) breeders were purchased from Jackson Laboratories. All mice were obtained from a previously backcrossed C57BL/6J line. To generate inducible Cre reporter mice, male mice hemizygous for Cre were crossed to female mice containing homozygous Ai9 alleles (Figure 1).

Experimental overview and tamoxifen dosing timeline

Tamoxifen (TMX) was mainly administered by oral gavage dissolved in corn oil (Sigma-Aldrich, CAS #10540-29-1; 100 mg/kg). In an initial cohort of mice (~10% of study), TMX was given by chow diet (ENVIGO TD. 130859; ~40 mg/kg daily) for continuous dosing strategies but later discontinued in favor of gavage dosing. We did not observe differences in TdTomato expression during bone healing between tamoxifen administration methods when used continuously (data not shown). Experimental mice harboring Osx Cre_ERT2+/−; Ai9+/− (+= presence of transgene; -= absence of transgene) or DMP1 Cre_ERT2+/−; Ai9+/− and given TMX served as Cre reporter mice and are labeled as OsxTMX and DMP1TMX, respectively (Table 1). Mice harboring Cre and Ai9 alleles and only given the vehicle corn oil or chow without tamoxifen were used to assess Cre non-inducible recombination (i.e. “leakiness”) and are labeled as OsxVEH or DMP1VEH, respectively. To label pre-existing Osx+ and DMP1+ lineage cells (and their progeny) after bone injury, mice were given TMX continuously (2x weekly; 100 mg/kg) starting at 2 weeks before injury and throughout healing (Figure 2). These mice are referred to as OsxTMX;Continuous and DMP1TMX;Continuous groups, respectively. To label only pre-existing Osx+ and DMP1+ lineage cells (and their progeny) as well as newly-derived Osx+ and DMP1+ lineage cells and their progeny after bone injury, mice were given TMX continuously (2x weekly; 100 mg/kg) starting at 2 weeks before injury and throughout healing (Figure 2). These mice are referred to as OsxTMX;Continuous and DMP1TMX;Continuous groups, respectively.

To label only pre-existing Osx+ and DMP1+ lineage cells (and their progeny), mice were given three TMX doses 4 weeks before bone injury at 8 weeks of age (Figure 2). These mice are referred to as OsxTMX;Pulse and DMP1TMX;Pulse groups, respectively. We have previously reported residual tamoxifen effects on bone formation are negligible following a 4-week clearance time (Zannit and Silva, 2019). Male and female mice were used as available and in approximately equal numbers among experimental groups. We utilized both males and females.
in this study since both mouse sexes have been readily utilized in these inducible Cre lines in previous literature (Buettmann et al., 2019; McKenzie et al., 2019; Harris and Silva, 2022). Mice were group housed under a standard 12-h light/dark cycle and given access to food and water ad libitum.

Models of bone repair

For each bone repair model, the right limb was injured whereas the left contralateral limb served as the uninjured control. Mice were given buprenorphine SR LAB (1 mg/kg, s. c.) one hour before injury, and anesthetized during all procedures with isoflurane (1–3% v/v). The right limb was shaved and sterilized with betadine and alcohol (70%) before surgery. Following all bone injury procedures, mice were returned to their cage and placed on electronic heating pads (BeanFarm; Ultratherm) until awake and sternal. Mice were monitored daily for signs of pain and distress and open wounds were quickly resutured and treated with topical triple antibiotic ointment.

Femoral semi-stabilized transverse fracture

Right femurs were prepared for fracture as previously described (McBride-Gagyi et al., 2015; McKenzie et al., 2018). Briefly, a complete (full) transverse bone fracture was made in the femoral mid-diaphysis via three-point bending using a custom designed fixture on a materials testing machine (Instron, DynaMight 8841). The fracture was stabilized with a 24-gauge stainless steel pin (Microgroup, #304 H24RW) and the wound sutured with 3–0 nylon sutures in a simple interrupted pattern (Ethicon, #1669H). Immediately after fracture, lateral radiographs at ×3 magnification (Faxitron, Ultrafocus 100) were taken to verify proper fixation of the fracture site. Mice were allowed to heal for 14 days post-injury (PID 14), when the intramembranous woven bone on the callus periphery and cartilage undergoing endochondral ossification near the

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<th>Mouse genotype</th>
<th>Treatment (100 mg/kg)</th>
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<tr>
<td>Osx Cre&lt;sup&gt;ERT2&lt;/sup&gt;−/−; A&lt;sup&gt;9&lt;/sup&gt;−/−</td>
<td>Tamoxifen in Corn Oil</td>
<td>Osx&lt;sup&gt;TMX&lt;/sup&gt;</td>
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<td>Osx Cre&lt;sup&gt;ERT2&lt;/sup&gt;−/−; A&lt;sup&gt;9&lt;/sup&gt;−/−</td>
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<td>DMP1 Cre&lt;sup&gt;ERT2&lt;/sup&gt;++/−; A&lt;sup&gt;9&lt;/sup&gt;−/−</td>
<td>Tamoxifen in Corn Oil</td>
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fracture line are both visible (Supplementary Figure S1) (Einhorn and Gerstenfeld, 2015; Buettmann et al., 2019).

Ulnar stress fracture

Right ulnas had a stress fracture generated as previously described (Martinez et al., 2010; Buettmann et al., 2019). Briefly, a non-displaced (partial) stress fracture was made in the ulnar mid-diaphysis via fatigue loading by cyclic compression on a material testing machine (Instron, DynaMight 8841). Right forelimbs were loaded at a calibrated peak force of 3.1 N to a 50% increase in cyclic displacement from the 10th cycle of loading. Previous work has shown that loading to this average cyclic displacement level in similarly aged wildtype C57BL/6 mice produces a reproducible non-displaced crack on the compressive surface (Buettmann et al., 2019). Mice were allowed to heal for 10 days post-injury (PID 10), when the woven bone response, predominantly formed via periosteal intramembranous ossification, is maximal (Supplementary Figure S1) (Uthgenannt et al., 2007; Martinez et al., 2010).

Tibial cortical defect

The right tibia was prepared as previously described (Kim et al., 2007; Liu et al., 2018). Briefly, a 0.78 mm monocortical circular defect was made using a #68 sterilized drill bit attached to a Dremel tool (Bosch Tool Group, Model 395). It was centered on the anterior medial cortex of the tibia and was located 4.3 mm from the tibial plateau. Following drilling, the cortical defect was irrigated with sterile saline with the wound closed using 5-0 nylon sutures (McKesson, #1034511). Mice were allowed to heal for 7 days post-injury (PID 7), when the woven bone response, formed via intramembranous ossification, encompasses the entire localized marrow space (Supplementary Figure S1) (Uthgenannt et al., 2007; Martinez et al., 2010).

Frozen histology

Injured and contralateral uninjured limbs were harvested at previously mentioned timepoints (transverse femoral fracture–PID14, ulnar stress fracture–PID10, tibial cortical defect–PID7) and immediately fixed in 4% paraformaldehyde (Electron Microscopy Sciences; #15710) for 24 h. A small subset of transverse femoral fracture femurs (injured + contralateral) were also harvested at days 5 (pulse TMX dosing strategy) and day 7 (continuous TMX dosing strategy) to investigate Osx+ and DMP1+ lineage cells in the rapidly expanding periosteum and mesenchyme before robust woven bone formation. All specimens underwent standard decalcification for 14 days (14% EDTA, pH 7.0) and subsequent tissue processing (30% sucrose infiltration) followed by embedding and freezing in O.C.T. Compound (Tissue-TekTM; #25608-930). Sections were cut longitudinally at a thickness of 5 μm using the Leica CryoJane Tape-Transfer System and stored at −80°C until use.

Imaging and TdTTomato quantification

Slides were rehydrated in deionized water, counterstained using DAPI (Sigma-Aldrich, #D9542, 1:1,000 in DiH2O), and mounted with Fluoromount aqueous mounting media (Thermo Fisher Scientific, #0-4958-02). Sections were subsequently imaged under consistent exposure settings for DAPI and TRITC signal at 20–40× magnification by the Nanozoomer Digital Slide Scanning System (Hamamatsu, S360 System). Images containing both channels (DAPI; TdTTomato) were exported using NDP viewer2 (Hamamatsu, #U12388-01) software with consistent image settings (Contrast = 200%; γ = 1.8).

Contralateral uninjured femur analysis

40X images were randomly taken from each cortical diaphyseal quadrant (ROIs: anterior-proximal; anterior-distal; posterior-proximal; posterior-distal) from uninjured D7 and D14 continuous TMX and vehicle mice from each Cre_ ERT2 line. We did not see any differences in TdTTomato expression (Cre activation) between uninjured D7 and D14 images. Images were blinded and manually counted for TdTTomato positive (TdT+) osteocytes, periosteal labeled surface and endosteal labeled surface using the FIJI (Schindelin et al., 2012) ROI manager and cell counter plug-in. TdTTomato positive (TdT+) cells were normalized to total number of osteocytes or endosteal/periosteal bone surface length for their respective indices. Indices for all four cortical ROIs were averaged on each specimen for final data statistical analysis. 20x images from the femoral mid-diaphysis and distal femoral growth plate were also captured to qualitatively determine relative targeting of skeletal muscle, marrow cells, and chondrocytes based on cellular morphology and anatomical location (Supplementary Figure S2).

PID14 femur fracture analysis

Woven Bone (Intramembranous Region): Any tissue between the skeletal muscle and cortical bone was considered callus tissue. 40X images were randomly taken from two woven bone regions in the callus (~2.5–3 mm peripheral to the fracture site), one on the anterior side of the bone and the other on the posterior side. Images were blinded and manually counted for TdTTomato positive (TdT+) osteocytes within woven bone (Wo.B). Osteocytes were counted as any cell within the woven bone (Wo.B) tissue, excluding the bone surface and adjacent marrow spaces (marked by clusters of overlapping cells). The multi-layered outline of cells encompassing the perimeter of the woven bone (i.e. expanded periosteal perimeter) was also manually counted for TdT+ cells using the FIJI (Schindelin et al., 2012) ROI manager and cell counter plug-in. Both
indices were normalized to total Wo_B osteocytes and callus perimeter length, respectively. TdT+ cellular area was also computed automatically by FIJI as per previous methods and normalized to total callus area (Wang et al., 2019; Shihan et al., 2021). In brief, TdT+ cell area was counted automatically by collecting data only on the red channel (split channel function), thresholding to make the image binary (threshold 190), and calculating the thresholded area (particle analysis–no restrictions on size/circularity).

Cartilage (Endochondral Region): 40X images were randomly taken from two cartilage regions anterior and posterior to the fracture site away from the mineralizing woven bone front. TdT+ cartilage cellular area was also computed automatically by FIJI as per exact methods listed for the woven bone region and normalized to total cartilage area. TdT+ cartilage cells were calculated for the same images by splitting the red and blue channels, and using particle analysis to automatically count the ratio of TdT+ to DAPI+ cells. In brief, TdT+ cells were counted by binary thresholding (threshold 190), discretizing overlapping cells by watershed analysis, and running particle analysis (size: 20–200 microns; circularity: 0.2-1.0). DAPI+ cells were counted by binary thresholding (threshold 150), discretizing overlapping cells by watershed analysis, and running particle analysis (size: 20–200 microns; circularity: 0.2-1).

**PID10 stress fracture analysis**

To complement the woven bone analysis for femoral fracture mice, the periosteal stress fracture callus was also analyzed. For this, 10X images were taken that were centered at the stress fracture crack line of the compressive region of the callus (this ROI encompassed the majority of the callus). Images were blinded and manually counted for TdT+ positive osteocytes within woven bone (TdT+ Wo_B Osteocytes) and TdT+ callus perimeter similar to methods used for the femoral woven bone analysis. TdT+ Wo_B cells were calculated for the same images by splitting the red and blue channels and using particle analysis to automatically count the ratio of TdT+ to DAPI+ cells within the 10X callus region, regardless of location. This callus population included the total number of TdT+ woven bone lining cells, woven bone marrow cells and osteocytes. In brief, TdT+ cells were counted by binary thresholding (threshold 150), discretizing overlapping cells by watershed analysis, and running particle analysis (size: 20–200 microns; circularity: 0.2-1). DAPI+ cells were counted by binary thresholding (threshold 20), discretizing overlapping cells by watershed analysis, and running particle analysis (size: 20–200 microns; circularity: 0.2-1).

**PID7 tibial cortical defect analysis**

To determine if contributions of pre-existing Osx+ and DMP1+ lineage cells differ following marrow-derived intramembranous bone repair, a small number of mice were given cortical defect injuries following pulse TMX regimens. (Cortical defect experiments were not performed under the continuous TMX protocol.) 10X images were taken centered at the PID7 cortical defect site around the anterior medial surface of the tibia and used to investigate TdT+ cells expression within the intramedullary woven bone.

**Statistics**

Quantitative outcomes of TdTomato cellular expression per tissue area (woven bone or cartilage), per perimeter (callus or bone), or per cell number were analyzed within each inducible Cre line (Osx or DMP1). Due to the smaller sample sizes used (n = 2-4), data normality was first assessed by Q-Q plots and assumed to be normal if not deviating significantly from a straight diagonal line. Depending on outcome, data was compared by unpaired t-test or ANOVA (normally distributed) or Mann-Whitney or Kruskal-Wallis to test for the significant effects of tamoxifen dosing (continuous, pulse, vehicle) in GraphPad Prism Pro Version 9 (La Jolla, CA). The type of statistical test for each figure is noted in the legend. Direct statistical comparisons between Cre lines were avoided due to potential confounding technical differences in A9 recombination efficiency between Osx Cre_ERT2 and DMP1 Cre_ERT2 constructs, which may not reflect accurate changes in biology. Mouse sex was not tested as an independent variable because our study wasn’t adequately powered to compute male and female differences (so they were pooled for analysis). For added clarity, data points from male and female mice are represented on graphs as diamonds and circles, respectively as noted in each figure’s caption. Post-hoc Tukey’s (parametric - ANOVA) or Dunn’s (non-parametric - Kruskal–Wallis) were used to determine significance differences between individual groups after accounting for multiple comparisons corrections. Statistical significance was defined as p < 0.05 and trending values were denoted as p < 0.10. Data are presented as mean ± SD with individual sample sizes for each outcome denoted as data points in each graph and in the figure caption.

**Results**

**Inducible Osx Cre_ERT2 under continuous dosing targets a higher percentage of femoral cortical bone cells in uninjured bones compared to inducible DMP1 Cre_ERT2**

Uninjured contralateral femurs were first assessed for TdT+ cells in intracortical osteocytes and bone surfaces at the mid-diaphysis following continuous TMX administration for 4 weeks. Overall, OsxTMS;Continuous femurs showed greater targeting of cells compared to DMP1TMS;Continuous femurs in
each bone component analyzed (Figure 3). For example, Osx<sup>TMX;Continuous</sup> femurs had 98% of osteocytes labeled TdT<sup>+</sup> compared to 72% in DMP1<sup>TMX;Continuous</sup> femurs. In addition, Osx<sup>TMX;Continuous</sup> femurs had 91% and 85% of the periosteal and endosteal surfaces labeled, whereas DMP1<sup>TMX;Continuous</sup> femurs had 66% and 77% of the periosteal and endosteal surfaces labeled, respectively (Figure 3). The majority of TdT<sup>+</sup> labeling was attributed to tamoxifen induction as expected, as both Osx<sup>TMX;Continuous</sup> and DMP1<sup>TMX;Continuous</sup> femurs had significantly increased TdT<sup>+</sup> labeling in all investigated cortical compartments versus respective vehicle-treated controls (p < 0.05). In the absence of TMX, the periosteal surface and endosteal surface had negligible non-inducible recombination (“leakiness”) in either Cre<sub>ERT2</sub> line, however leakiness was readily apparent in intracortical osteocytes. For example, 9.4% of osteocytes were TdT<sup>+</sup> in Osx<sup>VEH</sup> femurs while in DMP1<sup>VEH</sup> femurs 21% of osteocytes were TdT<sup>+</sup> (Figure 3).

Qualitative assessment of TdT<sup>+</sup> cell labeling outside the cortical diaphyseal bone in the marrow, skeletal muscle and primary spongiosa was also performed (Supplementary Figure S2). Osx<sup>TMX;Continuous</sup> and DMP1<sup>TMX;Continuous</sup> femurs both showed minimal TdT<sup>+</sup> expression in marrow cells (Supplementary Figure S2; Panel 1). Notably, both DMP1<sup>TMX;Continuous</sup> and DMP1<sup>VEH</sup> femurs showed robust TdT<sup>+</sup> expression in skeletal muscle cells, indicative of non-inducible recombination at this site (Supplementary Figure S2; Panel 2). Looking at the distal femoral growth plate, a place undergoing endochondral ossification similar to the fracture callus, Osx<sup>TMX;Continuous</sup> femurs showed greater targeting of growth plate chondrocytes (white arrows) and trabecular bone within the primary spongiosa compared to DMP1<sup>TMX;Continuous</sup> femurs (Supplementary Figure S2; Panel 3). In summary, these results demonstrated that DMP1<sup>VEH</sup> femurs had greater non-inducible TdTomato expression and hence leakiness in multiple tissue compartments, notably intracortical osteocytes and skeletal muscle, compared to Osx<sup>VEH</sup>. However following a 4-week period of TMX dosing, Osx<sub>Cre<sub>ERT2</sub></sub> caused Cre activation in a greater number of bone cells compared to DMP1<sup>TMX;Continuous</sup> femurs, such as intracortical osteocytes, periosteal and endosteal lining cells, and growth plate chondrocytes.
Pulse-chase labeling reveals that pre-existing Osx⁺ but not DMP1⁺ lineage cells and their progeny give rise to most intramembranous woven bone osteocytes following femoral fracture.

The callus from the fractured femurs was next analyzed for TdTomato expression in woven bone regions at the callus periphery, known to primarily undergo intramembranous ossification, and revealed a large contribution of pre-existing Osx⁺ but not DMP1⁺ lineage cells. With pulse dosing, Osx⁺璀璨 callus had noticeably increased TdT⁺ stained woven bone area compared to OsxVEH control (16% vs. 0.02%; Figure 4A) but still nearly 3-fold less staining than OsxTMX;Continuous femurs (16% vs. 44%; Figure 4A). Notably, TdT⁺ osteocytes were significantly more abundant in OsxTMX;Pulse calluses (74%) than OsxVEH controls (0.47%) although less abundant than OsxTMX;Continuous femurs (99%, p < 0.05; Figure 4A). Lastly, OsxTMX, Pulse femurs also had more TdT⁺ labeled cells lining the perimeter of the intramembranous woven bone callus compared to OsxVEH (24% vs 0%) but this only reached significance versus vehicle in the OsxTMX;Continuous group (66% vs 0%, p < 0.05; Figure 4A). Taken together,
these results indicate that pre-existing Osx+ lineage cells and their progeny (identified by pulse-chase labeling) make up about a fifth of intramembranous callus tissue (16%), which is two to three-fold less than the amount labeled by continuous dosing (44%), which captures both pre-existing and newly differentiated Osx+ lineage cells and their progeny. Notably, the majority of intramembranous woven bone osteocytes are derived from pre-existing Osx+ lineage cells and their progeny (74%).

In contrast, DMP1\textsuperscript{TMX}; Pulse calluses demonstrated minimal increases in TdT+ woven bone area that was not significantly different from DMP1\textsuperscript{VHS}; controls (2.6% vs. 0.3%; Figure 4B). Moreover, DMP1\textsuperscript{TMX}; Pulse calluses had significantly less TdT+ intramembranous area compared to DMP1\textsuperscript{TMX};Continuous Calluses (2.6% vs 19%; p < 0.05; Figure 4B). Differential TdT+ labeling between DMP1\textsuperscript{TMX}; Continuous and DMP1\textsuperscript{TMX}; Pulse was even more apparent in woven bone intracortical osteocytes and the callus periphery. For instance, DMP1\textsuperscript{TMX}; Pulse intramembranous calluses showed a trending but non-significant increase in TdT+ osteocytes compared to DMP1\textsuperscript{VHS}; controls (26% vs. 7.2%, p < 0.10; Figure 4B) but was significantly less compared to DMP1\textsuperscript{TMX};Continuous calluses (26% vs. 87%, p < 0.05; Figure 4B). In addition, only DMP1\textsuperscript{TMX};Continuous calluses had significantly more TdT+ cells lining the periphery of the intramembranous woven bone compared to DMP1\textsuperscript{VHS}; (36% vs. 1.3%, p < 0.05; Figure 4B). These DMP1+ lineage results indicate that pre-existing DMP1+ lineage cells contribute minimally to callus formation and only become a small fraction of the total DMP1+ cell lineage population (pre-existing and newly-derived). These pre-existing DMP1 lineage cells only contribute to the initial woven bone osteoblasts and osteocytes (marked by proximity to original cortical bone) during fracture healing.

Comparing Osx and DMP1 Cre\_ERT2 models, it appears that most osteocytes within woven bone come from pre-existing Osx+ lineage cells and their progeny and will acquire DMP1+ expression as evidenced by the similar labeling of osteocytes between Osx\textsuperscript{TMX}; Pulse and DMP1\textsuperscript{TMX}; Continuous calluses (74% versus 87%). This is further supported when looking at early timepoints of fracture healing such as PID5 and PID7 in pulsed and continuous fracture calluses, respectively (Supplementary Figures S3, S4). For example, by PID5 a greater extent (i.e. longitudinal length) of the expanded periosteum is labeled by pre-existing Osx+ lineage cells than DMP1+ lineage cells and their progeny within woven bone tissue at PID7 (Supplementary Figure S4). Overall, our results indicate that pre-existing Osx+ lineage cells and their progeny, but not DMP1+ lineage cells (and their progeny), contribute to early

FIGURE 5
Newly-derived but not pre-existing Osx+ (panel (A)) and DMP1+ (panel (B)) lineage cells and their progeny make up cartilage callus following femoral fracture (endochondral region). 2.5X (Overview; scale bar 1 mm) and 40X images (cartilage - scale bar 50 μm; black/white ROI boxes) were randomly taken and averaged from two regions of interest centered in cartilage regions adjacent to the PID14 fracture site (Fx). TdTomato positive callus area were calculated from automated thresholding for TdT+ area within each field of view. Cartilage TdTTomato positive cells (TdT+ Cg cells - white arrow) were thresholded, counted and normalized to total number of DAPI+ chondrocytes (TdT− and TdT+ cells - orange arrow). Data presented as mean ± SD with n = 2 – 4 per group. Mouse sex of each data point is represented by shape (circle = female; diamond = male). * Significant Tamoxifen Effect by 1-WAY ANOVA. &p < 0.10, *p < 0.05; **p < 0.005; Significantly Different by Tukey Post-Hoc.
woven bone formation in the fracture callus both by lining new woven bone surfaces and becoming embedded osteocytes.

Newly-derived but not pre-existing Osx+ and DMP1+ lineage cells and their progeny make up cartilage callus following femoral fracture

TdTomato expression in multiple cartilage regions immediately adjacent to the femoral fracture site was averaged to evaluate the role of pre-existing versus newly-derived Osx+ and DMP1+ lineage cells and their progeny in endochondral ossification at 14 days post-fracture. Overall, we saw little evidence of pre-existing Osx+ or DMP1+ lineage cells contributing to cartilage formation. For example, with pulse dosing, OsxTmX; Pulse calluses had non-significant TdT+ stained cartilage callus tissue (0.3% versus 0.0%) and cartilage cells (2.3% versus 0.1%) compared to OsxVEH control (Figure 5A). However, with continuous tamoxifen dosing there were trending increases in OsxTmX;Continuous TdT+ stained callus tissue (6.2 versus 0.3%) and cells (48% versus 2.3%; p < 0.10)

FIGURE 6
Pre-existing Osx+ (panel A) but not DMP1+ (panel B) lineage cells and their progeny significantly contribute to woven bone cells during stress fracture. 10X and 20X images (S_Fx insert - scale bar 100 µm) centered around the PID10 stress fracture line (Fx). TdT+ positive cells within the callus were thresholded, counted and normalized to total number DAPI+ callus cells (TdTomato positive Wo. B Cells). TdT+ positive Wo. B osteocytes (TdTomato positive Wo. B osteocytes - white arrow) were normalized to total number of osteocytes (TdTomato and TdT- osteocytes - orange arrow). TdT+ periosteal callus perimeter was normalized to total callus perimeter length (C.P.). Data presented as mean ± SD with n = 1–4 per group. Mouse sex of each data point is represented by shape (circle—female; diamond—male). * Significant Tamoxifen Effect by 1-WAY ANOVA. **p < 0.005; ****p < 0.00005 Significantly Different by Tukey Post-Hoc.
compared to OsxVEH control (Figure 5A) indicating the majority of Osx+ lineage cartilage cells are newly-derived following fracture. Similarly, with pulse dosing, DMP1VEH; Pulse calluses had minimal cartilage callus area (0.1% versus 0.0%) and cartilage cells (0.7% versus 0.0%) targeted compared to DMP1VEH but this was significantly enhanced with continuous tamoxifen expression (Figure 5B). While continuous TMX dosing resulted in significant cartilage labeling compared to vehicle controls in both Cre lines, OsxTMX; Continuous femurs, on average, targeted approximately 10-fold more chondrocytes compared to DMP1TMX. Continuous femurs (48% versus 6.0%). These data indicate that pre-existing Osx+ and DMP1+ lineage cells and their progeny give rise to minimal chondrocytes in the fracture callus. However, it appears that a large portion of total chondrocytes become Osx+ lineage cells once formed in the fracture callus between PID7 and PID14, with an even smaller population of chondrocytes becoming DMP1+ lineage cells near sites of endochondral ossification.

Pre-existing Osx+ but not DMP1+ lineage cells and their progeny contribute a significant but small portion of periosteal woven bone osteocytes following ulnar stress fracture

The ulnar stress fracture model was utilized in each Cre_ERT2 line (Osx and DMP1) with pulse and continuous TMX dosing to further assess the role of each osteoblast cell lineages’ contribution to periosteal woven bone intramembranous repair. These results partially mirrored the findings in the intramembranous region of the femoral fracture callus and suggest that pre-existing Osx+ but not DMP1+ lineage cells and their progeny contribute significantly more to woven bone formation following stress fracture. For example, OsxTMX; Pulse stress fracture calluses had significantly increased TdT+ cells within the woven bone regions of the stress fracture callus compared to OsxVEH control (20% versus 0%; p < 0.05; Figure 6A). However, the overall TdT+ cell population was significantly less in OsxTMX; Pulse calluses compared to OsxTMX; Continuous Calluses (20% vs 82%; p < 0.05; Figure 6A). Stratifying TdT+ cells based on location, the majority of TdT+ cells in OsxTMX; Pulse stress fracture calluses were embedded woven bone osteocytes (24% of Wo.B osteocytes TdT+; p < 0.05 compared to 0% in OsxVEH) but not callus peripheral cells in the expanded periosteum (4.6% peripheral cells TdT+; p > 0.05 compared to 0% OsxVEH). With continuous TMX dosing, OsxTMX; Continuous calluses showed a significant elevation in TdT+ targeting of woven bone osteocytes (95% of Wo.B osteocytes TdT+; p < 0.05) and expanded callus periosteum (72% of peripheral cells TdT+; p < 0.05) compared to OsxTMX; Pulse and OsxVEH groups (Figure 6A). These data indicate that pre-existing Osx+ lineage cells and their progeny make up a small but significant portion of total Osx+ lineage cells in intramembranous callus tissue following stress fracture, mainly in the form of woven bone osteocytes. In addition, the majority of intramembranous callus cells acquire Osx+ lineage cell specification after injury.

In contrast, DMP1TMX; Pulse stress fracture calluses showed no significant elevation of TdTomato expression in callus cells (including total cells, osteocytes, or callus perimeter) versus DMP1VEH controls (Figure 6B). However, with continuous TMX dosing, DMP1TMX; Continuous stress fracture calluses had significantly elevated TdTomato
expression in the callus (41% cells TdT +), osteocytes (67% TdT +), and callus perimeter cells (14% TdT +) compared to DMP1TMX; Pulse and DMP1VEH controls (Figure 6B; p < 0.05). Comparing Cre lines with continuous dosing, OsxTMX; Continuous had 2-fold greater TdT + labeling in total stress fracture callus cells versus DMP1TMX. Continuous callus (82% vs 41%). Moreover, OsxTMX; Continuous had approximately 1.5-fold greater osteocyte labeling (95% vs. 67%) and nearly 5-fold greater callus peripheral labeling (72% vs. 14%) compared to DMP1TMX; Continuous calluses. Collectively, these data indicate that pre-existing Osx + but not DMP1 + lineage cells and their progeny contribute a significant number of cells to stress fracture calluses. In addition, although newly-derived Osx + and DMP1 + lineage cells and their progeny make up the majority of cells in the stress fracture callus, newly-derived Osx + lineage cells contribute significantly more than newly-derived DMP1 + lineage cells to non-osteocytic populations.

Pre-existing Osx + but not DMP1 + lineage cells and their progeny significantly contribute to cells in the intramedullary woven bone following tibial cortical defect

To investigate the contribution of Osx + and DMP1 + lineage cells and their progeny in another widely used model of bone repair, a tibial cortical defect was created in OsxTMX; Pulse and DMP1TMX; Pulse mice. With TMX pulsing, TdT + signal was strongly present in the majority of woven bone cells, including woven bone osteocytes, woven bone lining cells and injured marrow surrounding the defect in OsxTMX; Pulse but not DMP1TMX; Pulse mice (Figure 7). OsxVEH and DMP1VEH defects showed minimal non-inducible expression. In all, this suggests that the majority of woven bone cells following cortical defect arises from pre-existing Osx + and their progeny but not pre-existing DMP1 + lineage cells and their progeny.
Discussion

We investigated the contributions of pre-existing versus newly-derived Osx+ and Dmp1+ lineage cells and their progeny to regenerated tissues in three preclinical models of bone injury using inducible Osx CreERT2 A9 and DMP1 Cre_ERT2 A9 mice. Using two different tamoxifen dosing regimens: 1) pulse-labeling with washout (4 weeks) before injury or 2) biweekly dosing before (2 weeks) and during bone injury healing, we found across injury models (femoral fracture, unlar stress fracture, tibial cortical defect) that pre-existing Osx+ lineage cells and their progeny, but not pre-existing DMP1+ lineage cells and their progeny, contributed a significant amount of total TdT+ labeled tissue area and cells versus respective vehicle controls (Figure 8). These results support our first hypothesis and demonstrate that pre-existing Osx+ lineage cells and their progeny but not DMP1+ lineage cells and their progeny are a significant source of woven bone forming osteoblasts and osteocytes following bone injury. In addition, continuous tamoxifen administration significantly increased labeling within each inducible Cre line. For example, Osx Cre_ERT2 showing significantly higher targeting of callus tissue with continuous TMX dosing across all scenarios compared to DMP1 Cre_ERT2, supporting our second hypothesis (Figure 8). Importantly, these results suggest that pre-existing Osx+ lineage cells and their progeny are likely critical for postnatal injury-induced bone formation, although their contribution varies based on skeletal site and the type of bone injury.

In the femoral fracture callus at day 14 post-injury, the specific Cre model and tamoxifen dosing regimen led to differential targeting of cells based on callus region. Comparing pulse to continuous dosing allowed us to see the maximum contribution of Osx+ or DMP1+ lineage cells and their progeny to bone healing (pre-existing and newly-derived) within each Cre line. Using this methodology, we saw when comparing Osx+; Pulse to Osx+; Continuous, that the majority of Osx+ lineage osteocytes (74% versus 99%) and some callus border cells (24% versus 66%) and virtually no chondrocytes (2.3% versus 48%) were derived from pre-existing Osx+ lineage cells and their progeny (Figure 4A and Figure 5A). Comparing DMP1+; Pulse to DMP1+; Continuous, we saw that out of all DMP1+ lineage cells involved in femoral fracture, only a small percent of pre-existing DMP1+ lineage cells become osteocytes (26% versus 87%), and virtually none become callus border cells (0.5% versus 36%) or chondrocytes (0.7% versus 6.0%; Figure 4B and Figure 5B). Our DMP1+; Pulse results are similar to previous results by Root et al., whom utilized the DMP1 CreERT2; A9 mouse crossed to the 2.3 kb Col1 Cre thymidine kinase (tk) mouse (Visnjic et al., 2001) during transcortical fracture healing in 8 week old mice (Root et al., 2020). In this study, the authors used ganciclovir administration (GCV) for 16 days prior to fracture to eliminate the proliferating 2.3Col1 tk+ osteoblast lineage cells, which overlap significantly with pulse-labeled DMP1+ lineage cells (Matic et al., 2016), leaving only transcartical DMP1+ lineage cells prior to injury (Root et al., 2020). Tracing of these pre-labeled transcartical DMP1+ lineage cells following fracture for 7 days revealed minimal contribution of DMP1+ lineage cells to periosteal woven bone, although all DMP1+ lineage cells lining woven bone were 2.3Col1 GFP+, suggesting that they were bone-forming osteoblasts (Root et al., 2020). Similarly, our data also suggest minimal contributions of pre-existing DMP1+ lineage cells (even those that may not be targeted by 2.3Col1 tk) and their progeny to woven bone following transverse fracture.

Looking at Osx and DMP1 CreERT2 lines collectively in our results indicate that pre-existing Osx+ osteoprogenitor lineage cells and their progeny at the time of fracture readily become woven bone callus lining cells, woven bone forming osteoblasts and the majority of embedded osteocytes, whereas pre-existing DMP1+ lineage cells and their progeny are mainly absent from fracture callus tissues (a few become woven bone lining osteoblasts and osteocytes) (Figure 8). Furthermore, the higher percentage of overall Osx+ lineage cells compared to DMP1+ lineage cells in intramembranous callus tissue area (44% versus 19%) and chondrocytes (48% versus 6.0%), but similar overlap in the percentage of osteocytes (99% versus 87%) targeted under continuous dosing regimen suggests that Osx Cre_ERT2 targets a wider population of bone cells that eventually go on to become DMP1+ lineage concurrent with woven bone formation and matrix embedding (i.e. osteocytogenesis in woven bone) (Figure 8). This wider targeting of osteoblast lineage cells using Osx Cre_ERT2 over DMP1 Cre_ERT2 is supported by previously published works that DMP1 is expressed at the mature osteoblast and osteocyte stages of differentiation during matrix mineralization and osteocyte cell embedding (Maes et al., 2007; Lu et al., 2011; Powell et al., 2011; Kim et al., 2012; Kalajzic et al., 2013; Matic et al., 2016; Shiflett et al., 2019).

Based on the continuous dosing regimen labeling pre-existing and newly-derived cells and their progeny, we also found that cells will acquire Osx+ lineage specification and to a lesser degree DMP1+ lineage specification within sites of endochondral ossification at day 14 in the femoral fracture callus (Figure 8). A limitation of this work is that we did not use co-staining to better characterize the identity of these newly-derived Osx+ or DMP1+ lineage cells observed near the cartilage to bone transition zone (e.g., Collagen type II or Collagen X staining). Another limitation is that our study was underpowered to detect differences in these TdT+ cell populations between mouse sexes. As emerging data suggests that mouse sex may differentially regulate the response to tamoxifen (Ceasrine et al., 2019) and lead to biological changes in fracture healing.
(particularly cartilage formation) (Haffner-Luntzer et al., 2021), future research is needed to determine if mouse sex significantly alters Cre specificity during bone healing. The similar trends in Cre specificity seen between males and females in our data suggest that mouse sex effects are subtle compared to the tamoxifen dosing regimen and Cre construct used for inducible cell targeting. Non-ethereal, the anatomic location of these newly-derived Osx+ and DMP1+ lineage cells and their progeny in both mouse sexes, within the chondrocyte transition zone (near vasculature), are in line with other reports showing that Osx+ lineage cells labeled continuously during fracture healing can demonstrate a hypertrophic chondrocyte phenotype (labeled by Collagen X) (Hu et al., 2017; Buettmann et al., 2019). In addition, DMP1 mRNA has previously been shown via in situ hybridization to be weakly expressed in a small number of hypertrophic chondrocytes in the growth plate (Lu et al., 2011) and during fracture repair (Toyosawa et al., 2004). These results, along with our own, are consistent with the trans-differentiation of chondrocytes to osteoblast lineage cells as proposed by others (Bahney et al., 2015; Hu et al., 2017). Our use of pulse-labeling strategies extends these prior results and indicates that chondrocytes likely do not arise from pre-existing Osx+ or DMP1+ lineage cells following bone injury. Therefore, researchers studying conditional gene deletion postnatally during transverse fracture repair would minimize targeting of cartilage cells with Osx Cre_ERT2 or DMP1 Cre_ERT2 mice by using a similar pulse dosing strategy. However, our results differ from Mizoguchi et al., which showed that Osx+ lineage cells labeled at postnatal day 5 (P5), can become fracture callus chondrocytes following bone injury nearly 15 weeks later (Mizoguchi et al., 2014). Overall, this suggests that there is a critical time-window between birth and 8 weeks postnatally in which pre-existing Osx+ osteoprogenitor cells are bipotent in vivo.

In order to complement our femoral fracture results, we tested the requirement of pre-existing and newly-derived Osx+ and DMP1+ lineage cells and their progeny to contribute to stress fracture repair. This model heals predominantly by intramembranous ossification (Martinez et al., 2010) and has not been extensively explored in the literature using Cre reporter mice. Our results largely mirror the woven bone results seen at day 14 of healing in the intramembranous region of the femoral fracture callus, with pre-existing Osx+ but not DMP1+ lineage cells and their progeny significantly contributing to callus woven bone cells based on changes from each Cre lines respective vehicle controls (Figure 6). These results reinforce that the stress fracture model largely mirrors the intramembranous processes in the femoral fracture model as we previously reported (Wohl et al., 2009). However, what was striking is that the overall percentage of total Osx+ lineage Wo.B osteocytes labeled in pulse versus continuous dosing was much lower in the stress fracture (~20% total Osx+ lineage Wo.B osteocytes came from pre-existing Osx+ lineage cells) compared to femoral fracture (~75% total Osx+ lineage Wo.B osteocytes came from pre-existing Osx+ lineage cells). In contrast, osteocytes expressing DMP1+ cell lineage between pulse vs. continuous labeling were relatively unchanged (~20–25% total DMP1+ lineage Wo.B osteocytes came from pre-existing DMP1+ lineage cells) between full fracture and stress fracture repair. These findings suggest, that pre-existing Osx+ lineage osteoprogenitors and their progeny contribute less to total callus area and cellularity in the less traumatic ulnar stress fracture than the femoral fracture model. The overall result that pre-existing Osx+ lineage cells contribute more to callus cells than DMP1+ lineage cells with higher degrees of bone damage are consistent with previous reports using anabolic tibial loading at graded force levels (Harris and Silva, 2022), and may potentially reflect the smaller overall cellularity and decreased proliferative processes in stress fracture versus full fracture injuries as previously shown (Coates et al., 2019). However, it may also reflect changes in anatomic location (ulna versus femur) or slight differences in analysis regions between the two fracture models used in the current study (i.e. majority of callus used to analyze stress fracture vs. callus periphery in transverse fracture).

Despite these differences in pre-existing Osx+ lineage cell recruitment, the full fracture and stress fracture model also show some striking similarities in the types of cells targeted between both inducible Cre drivers. For example, in both models, Osx\textsuperscript{TMX}; Continuous but not DMP1\textsuperscript{TMX}; Continuous labeling results in a strong TdT+ signal within woven bone marrow spaces known as sites of progenitor cell and blood vessel invasion (Figure 4 and Figure 6) that support bone healing (Hausman et al., 2001; Lu et al., 2006; Tomlinson et al., 2013). Furthermore, Osx\textsuperscript{TMX}; Pulse labeling shows much weaker TdT+ signal compared to Osx\textsuperscript{TMX}; Continuous calluses at these woven bone marrow sites. Maes et al. has demonstrated previously that osteoblast precursors labeled instantaneously by Osx Cre (but not Collagen 1 Cre), can take on a pericyte-like profile and co-invade woven bone spaces in the fracture callus, thereby supporting angiogenesis and subsequent bone formation (Maes et al., 2010). This concept was further supported by Buettmann et al., where using continuous dosing in Osx Cre_ERT2 VEGFA\textsuperscript{fl/fl} mice led to decreased femoral fracture and stress fracture angiogenesis and subsequent woven bone formation (Buettmann et al., 2019). Our pulse labeling strategy expands upon these results and suggests that pre-existing Osx+ lineage cells and their progeny, due to reduced TdT+ targeting of Wo.B marrow cells in femoral fracture and ulnar stress fracture, likely do not co-invade with vasculature (neither do more mature DMP1+ lineage cells). Thus, if Osx Cre_ERT2 VEGFA\textsuperscript{fl/fl} mice were pulse-dosed with TMX (rather than continuously dosed as previously performed in Buettmann et al., 2019), we hypothesize that femoral fracture and ulnar stress fracture healing would not be impaired.

Lastly, we showed that pre-existing Osx+ but not DMP1+ lineage cells and their progeny make up a majority of intramedullary woven bone tissue following monocortical defect. In particular, Osx\textsuperscript{TMX}; Pulse showed TdT+ cells
encompassing the majority of woven bone surfaces, osteocytes and even adjacent marrow, whereas these sites were largely void of TdTomato expression in DMP1\textsuperscript{TMX}; Pulse defects (Figure 7). These results indicate that, at 8 weeks age, pre-existing Osx\textsuperscript{+} lineage cells but not pre-existing DMP1\textsuperscript{+} lineage cells and their progeny significantly contribute to intramedullary bone formation following cortical defect. Although the exact bone compartment contributing to this differential TdTomato expression is unknown, work by Colnot suggests that both endosteal and marrow derived cellular niches act locally to play a large role in monocortical defect healing (Colnot, 2009). Therefore, it is likely that the pre-existing Osx\textsuperscript{+} but not pre-existing DMP1\textsuperscript{+} lineage cells contributing to defect labeling are derived from the endosteurum or marrow niche. Although continuous labeling revealed similar endosteal (Figure 3) and minimal marrow (Supplementary Figure S2; Panel 1) targeting in Osx Cre\_ERT2 and DMP1 Cre\_ERT2 mice, pre-existing lineage cells at these sites were not quantitated in pulse-labeled uninjured specimens, which is a limitation of the current work. Other reports indicate that later pulse labeling (14 days postnatal or after) in Osx Cre\_ERT2 and DMP1 Cre\_ERT2 labels vascular associated reticular marrow cells and endosteal bone-lining cells that decrease in number over time (Powell et al., 2011; Kim et al., 2012; Park et al., 2012). For example, Matic et al. demonstrated that DMP1\textsuperscript{+} lineage endosteal bone lining cells decrease by 50–75\% 3 weeks following tamoxifen induction (Matic et al., 2016). Therefore, it is possible that the differential Osx\textsuperscript{+} and DMP1\textsuperscript{+} lineage cell labeling in the intramedullary woven bone seen in our study is due to a preferential decline in DMP1\textsuperscript{+} over Osx\textsuperscript{+} lineage endosteal cells during the 4 weeks between pulse labeling and the cortical defect creation. Another possibility is that Osx Cre\_ERT2 targets a marrow or endosteal lineage cell population with higher regenerative capacity overall compared to DMP1 Cre\_ERT2. This differential Cre specificity would be in line with previous reports showing that peri-vascular stromal Osx\textsuperscript{+} lineage cells in the marrow have high regenerative capacity following bone injury (Park et al., 2012; Mizoguchi et al., 2014). Future studies, using dual-labeling strategies, to determine the instantaneous degree of overlap between Osx\textsuperscript{+} and DMP1\textsuperscript{+} lineage cells in various bone compartments, would be particularly informative.

In all, we have shown in the current study that pre-existing postnatal Osx\textsuperscript{+} lineage cells and not pre-existing DMP1\textsuperscript{+} lineage cells and their progeny contribute significantly to cells populating woven bone in multiple widely used preclinical models of bone injury. This study underscores the importance that pre-existing Osx\textsuperscript{+} lineage cells play in bone regeneration, especially for early woven bone formation, and suggest that bone targeting therapies to improve healing might target this particular cellular subset. Furthermore, this work provides a tissue and cellular atlas for inducible Cre targeting using the Osx Cre\_ERT2 and DMP1 Cre\_ERT2 models during bone healing, thereby providing a framework for researchers using these widely available tools in future studies.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

Ethics statement

All animal procedures have been approved by the staff veterinarians of the Washington University in St. Louis Department of Comparative Medicine (DCM) and the Washington University in St. Louis School of Medicine Institutional Animal Care and Use Committee (i.e. WUSTL IACUC).

Author contributions

Study design: MS, JM, EB. Study conduct: EB, JM, PH, SY. Data collection: EB, JM, PH, SY. Data analysis: EB, JM. Data interpretation: EB, JM, SY, PH, MS. Drafting manuscript: EB. Revising manuscript: EB, JM, MS. All authors revised the manuscript critically and approve of the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fphys.2022.1083301/full#supplementary-material


