AML-induced osteogenic differentiation in mesenchymal stromal cells supports leukemia growth

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Introduction

Acute myeloid leukemia (AML) is characterized by the accumulation of immature myeloid blasts in the bone marrow (BM) and peripheral circulation. This abnormal growth of AML cells disrupts normal hematopoiesis in the BM microenvironment (1). However, it is unclear how different the composition of the AML-BM microenvironment is compared to its normal counterpart. It is also not clear whether changes in the BM microenvironment are induced directly by AML cells or indirectly through other factors.

The BM microenvironment contains endosteal and vascular niches. The vascular niches, which consist mostly of sinusoidal endothelial cells and pericytes, have been reported to support AML cell survival and BM engraftment and promote drug resistance (4–6). MSCs exhibit an extensive self-renewal capacity and are able to differentiate into three major mesodermal lineages: osteoblasts, adipocytes, and chondrocytes (7, 8). Osteogenic differentiation of MSCs results in expression of osteogenic proteins, including osterix, osteopontin, RUNX2, and tissue nonspecific alkaline phosphatase (TNAP, also known as ALP) (9, 10). It is not clear if and how AML cells induce changes in differentiation potential of MSCs.

BMPs are members of the transforming growth factor-β superfamily that are able to induce osteogenic differentiation during embryonic development (11–13). BMPs exert their biological effects by signaling through two type I serine/threonine kinase receptors, RIA and RIB, which are transduced by Smad1/5 signaling to induce osteogenic differentiation in MSCs. Gene expression array analysis revealed that AML cells induce connective tissue growth factor (CTGF) expression in BM-MSCs irrespective of AML type. Overexpression of CTGF in a transgenic mouse model greatly enhanced leukemia engraftment in vivo. Together, our data suggest that AML cells induce a preosteoblast-rich niche in the BM that in turn enhances AML expansion.
reported to induce connective tissue growth factor (CTGF) expression in MSCs (16). CTGF is a member of the CCN family of proteins and has been shown to regulate MSC differentiation (17). CTGF is highly expressed and prognostic in acute lymphoid leukemias (18). Inhibition of CTGF activity by anti-CTGF monoclonal antibody increased the survival of leukemia-bearing mice (19).

Here, we hypothesized that AML cells mediate functional alterations in stromal cells to induce preosteoblastic-rich niches in the BM microenvironment. To address this, we examined AML-MSCs and discovered that they are induced to undergo osteogenic differentiation. Human AML xenograft and syngenic mouse AML models were used to study AML-stroma interactions in vivo and confirm that AML cells alter the differentiation potential of MSCs by inducing CTGF to create specialized hospitable niches that support their growth. Investigations of primary patient-derived BM biopsies confirmed these findings.

Results

AML-MSCs are phenotypically distinct from normal MSCs. We isolated MSCs from the BM of AML patients (AML-MSCs; n = 24) aged 40–70 years and age-matched normal healthy donors (N-MSCs; n = 11). Patient characteristics, including AML subtype, are summarized in Supplemental Table 1 (supplemental material available online with this article; https://doi.org/10.1172/jci.insight.90036DS1). Morphologically, AML-MSCs are polygonal or irregularly shaped and are much larger than spindle-shaped N-MSCs (diameter, 100–150 μM versus 40–60 μM; P < 0.01) (Supplemental Figure 1A). Growth analysis of AML-MSCs and N-MSCs showed that AML cells grow 2- to 3-fold more slowly (P < 0.01) than N-MSCs (Supplemental Figure 1B). Furthermore, BrdU pulse and propidium iodide (PI) labeling assay revealed that 9.6% ± 4.1% of N-MSCs in S-phase were positive for BrdU uptake, versus only 2.59% ± 0.38% of AML-MSCs (P < 0.001; Supplemental Figure 2), indicating a slower proliferation rate for AML-MSCs. The cell surface phenotypes of AML-MSCs and N-MSCs revealed that BM-MSC–associated markers, including CD44, CD51, CD73, CD90, CD105, CD106, CD140b, CD146, and SUSD2, were expressed on both cell types at equal intensities (Supplemental Figure 3). Neither CD45 nor CD31 was expressed on either AML-MSCs or N-MSCs (Supplemental Figure 3).

Flow cytometry revealed that TNAP (clone W8B2), known to be expressed on osteoprogenitor cells (20), mature osteoblasts, and naive MSCs (21), was significantly upregulated in AML-MSCs compared with N-MSCs (Figure 1A). In the cohort of primary MSC samples isolated from AML patients with different disease status (newly diagnosed or in remission or relapsed; n = 29), the average mean fluorescence intensity (MFI) for TNAP was approximately 10-fold higher than that in N-MSCs (n = 11; Figure 1B, P < 0.01). The median MFI for N-MSCs was 146, versus 1,033 for AML-MSCs. Only ≤10% of AML-MSCs showed TNAP MFI values <500, suggesting that most AML subtypes overexpress TNAP (Supplemental Table 1). However, MFI of other cell surface markers analyzed was not significantly changed between AML- and N-MSCs types (Supplemental Figure 3 and 4).

AML-MSCs are primed for osteogenic differentiation. Because the osteogenic differentiation marker TNAP was upregulated in AML-MSCs compared with N-MSCs, we determined whether other osteogenic lineage–associated genes were also upregulated in AML-MSCs. mRNA expression of several genes associated with osteogenic differentiation determined by qRT-PCR was upregulated by 3- to 10-fold in AML-MSCs compared with N-MSCs (Figure 1C, n = 3), including transcription factors RUNX2 and osterix and the cell surface or extracellular matrix–associated genes osteopontin and TNAP. Interestingly, AML-MSCs but not N-MSCs stained positive for ALP enzyme activity by the BCIP/NBT substrate assay (days 0, Figure 1D). However, AML-MSCs on day 0 (before induced osteogenic differentiation) were not positive for Alizarin Red S (Supplemental Figure 1B). Therefore, we found that TNAP expression was upregulated by more than 10- to 15-fold in N-MSCs cocultured with OCI-AML3 cells compared with CB-derived CD34+ or CB-derived CD33+ cells or cells cultured alone.
Figure 1. Acute myeloid bone marrow–derived mesenchymal stromal cells are primed to differentiate into osteoblasts. (A) Tissue nonspecific alkaline phosphatase (TNAP) expression was analyzed by flow cytometry on normal donor–derived (Normal-MSCs) (green) or acute myeloid bone marrow–derived mesenchymal stromal cells (AML-MSCs) (red) over unstained cells (gray). Cells were incubated with anti-TNAP antibody (clone W8B2) conjugated with phycoerythrin (PE). The TNAP-stained cells were overlaid on unstained cells; representative histograms (n = 3 for each) are shown. Data were analyzed by FlowJo software. (B) MFI of normal MSCs (N-MSCs) (n = 10) or AML-MSCs (n = 23) stained with TNAP antibody were determined. AML samples were compared with control medium (Figure 2C). OCI-AML3-CM–pretreated MSCs exhibited increased TNAP activity and Alizarin Red S staining as MSCs pretreated with control medium in a time-dependent manner (Figure 2D). In addition, osterix, RUNX2, osteopontin, and TNAP were upregulated by 2- to 10-fold in N-MSCs cocultured with OCI-AML3 compared with N-MSCs cultured alone (Figure 2E). (C) Alkaline phosphatase activity was determined by an enzyme assay in N- and AML-MSCs was measured by qRT-PCR. GAPDH served as an equal loading control. (D) Alkaline phosphatase enzyme activity and absorbance at 405 nm for Alizarin Red S staining were quantitated as described in the methods section. Statistical data were analyzed by GraphPad Prism software. One-way ANOVA was used for comparison of 3 or more groups and unpaired Student’s t test was used for comparisons of 2 groups. (*) P < 0.05, ** P < 0.01, *** P < 0.001 versus control. Dunnett’s multiple comparison test was used to check the statistical significance in difference between multiple groups.
Figure 2. AML cells induce osteogenic differentiation in N-MSCs. (A) N-MSCs were cocultured with OCI-AML3 cells (red) or with cord blood–derived CD34+ cells (green) or N-MSCs cultured alone (blue) for 3, 5, or 7 days, and TNAP expression was analyzed by LSR-II flow cytometer (n = 3). Data were analyzed and histograms were generated by FlowJo software. (B) MFI of TNAP expression was quantified in N-MSCs cocultured with cord blood–derived CD34+ cells or OCI-AML3 for 3 or 5 days. (C and D) N-MSCs were cultured with or without OCI-AML3 cell–derived conditioned medium (OCI-AML3-CM) for 5 days before
bone fragment implantation. As expected, the leukemia cells strongly engrafted within the regions of HBMI implantation, as revealed by in vivo bioluminescence imaging (Supplemental Figure 9D) and H&E staining (Supplemental Figure 9E) as early as 10 days after transplantation. Phenotypic characterization of HBMI-derived MSCs revealed that these cells expressed MSC-associated proteins, including CD44, CD90, CD140b, CD73, CD105, CD106, CD166, CD51, and SUSD2, and were negative for CD45 and CD31 (Supplemental Figure 10). When cultured in osteogenic or adipogenic differentiation medium, these MSCs developed into Alizarin Red S-positive osteoblasts and Oil Red O-positive adipocytes (Supplemental Figure 11), suggesting that components of the BM microenvironment in HBMI mice are intact and functional. These data indicate that the HBMI model is useful for studies related to factors regulating leukemia-stroma interactions.

To determine whether leukemia cells induce osteogenic differentiation in BM stromal cells in vivo, we injected Molm13 AML cells intravenously into the HBMI mouse model. After HBMI implantation, one group of mice received PBS solution (PBS; n = 10) and another group received Molm13 cells (1 × 10^6 cells/mouse; n = 10) in 100 μl PBS. After 4 weeks, the mice were sacrificed and the bone fragments from each mouse were examined by immunohistochemistry (IHC) with the Opal multiplex tissue staining approach. Sections were costained with osteogenic markers, including osterix and RUNX2, and the hematopoietic marker CD45. All antibodies were human specific and did not recognize mouse antigens.

We found that, in bone fragments derived from control mice, RUNX2 and osterix expression was restricted to the endosteal region of the bone (Figure 4B). However, in bone fragments from the Molm13 group, RUNX2 and osterix expression was upregulated in both the endosteal surface and medullar cavity, indicating greater osteogenic activity in HBMI mice harboring leukemia cells (Figure 4B). We also assessed the absolute number of osterix+CD45− or RUNX2+CD45− stromal cells in bone fragments from Molm13 and control groups using inForm image analysis software and determined that both osteogenic stromal types were present at a 5- to 7-fold higher rate in bone fragments from mice treated with Molm13 cells as compared with controls (Figure 4C), indicating that the increased osteogenic activity in leukemic BM is due to osteogenic differentiation of stromal cells.

**AML-BM displays higher osterix and RUNX2 expression than normal BM.** Next, we examined AML patient BM biopsies and normal BM specimens (n = 5 each) for osteogenic expression by IHC staining. In the AML-BM biopsy specimens, osterix and RUNX2 were expressed not only in the bone-lining cells, but also in the BM cavity, matching the HBMI findings. In contrast, these proteins were not expressed in the marrow regions of the normal BM (Figure 4, D and E). Overall, osterix and RUNX2 were upregulated 3- and 6-fold, respectively, in AML-BM specimens compared with normal BM specimens (P < 0.01), further suggesting that AML-BM specimens exhibit increased osteogenic activity.

**AML cells induce osteogenic differentiation in MSCs through BMP-mediated signaling.** To investigate the mechanisms involved in AML-induced osteogenic differentiation in MSCs, we focused on the role of BMP-mediated signaling, as the induction of osteogenic differentiation by BMPs is well established (23). N-MSCs were cultured in the presence or absence of OCI-AML3-CM, and the activation of Smad1/5 proteins was determined by Western blotting. Interestingly, the Smad1/5 proteins were phosphorylated in N-MSCs cultured with OCI-AML3-CM in a time-dependent manner, with the signal peaking at 60 minutes after treatment (Figure 5A). To determine whether Smad1/5 phosphorylation occurs though BMP ligands, but not through cross-talk with other signaling pathways, we cultured N-MSCs with OCI-AML3-CM for 60 minutes in the presence or absence of the BMP type-1 receptor–specific inhibitor LDN-212854 (24). Interestingly, we observed a progressive decrease in OCI-AML3-CM–induced Smad1/5 phosphorylation in MSCs with increasing concentrations of LDN-212854 (10–1,000 ng/ml), suggesting that Smad1/5 activation in MSCs is mediated though a BMP ligand secreted by AML cells (Figure 5B).

To determine the role of BMP signaling in AML-induced osteogenic differentiation in MSCs, N-MSCs were cultured in the presence or absence of OCI-AML3-CM for 5 days with or without LDN-212854 and analyzed for osteogenic markers by qRT-PCR. Expression of osterix, RUNX2, and osteopontin was 2- to 3-fold higher in N-MSCs cultured in OCI-AML3-CM (Figure 5C) compared with the control medium. In
Figure 3. AML cells inhibit adipogenic differentiation in MSCs. (A) N- and AML-MSCs were cultured in adipogenic differentiation medium for 3 weeks. mRNA expression of adipocyte-associated genes, aP2, lipoprotein lipase, and PPARγ, in N- and AML-MSCs before and after induction of differentiation was analyzed weekly by qRT-PCR. (B) N- or AML-MSCs (n = 5 each group) were cultured in adipogenic differentiation medium for 28 days. On day 28, the adipocytes were stained with Oil Red O. Scale bar: 100 μm. (C) Oil Red O–positive cells (i.e., adipocytes) were counted in 10 microscopic fields per sample. (D) N-MSCs were pretreated with OCI-AML3–conditioned medium for 5 days before they were cultured in adipogenic differentiation.
medium for 7 days. Expression of indicated adipogenic lineage–associated genes before induction of differentiation and on day 7 was analyzed by qRT-PCR. GAPDH served as an equal loading control. Two-way ANOVA was used for comparisons of 3 or more groups and unpaired Student’s t-test was used for comparisons of 2 groups (**P < 0.01, ***P < 0.001, ****P < 0.0001 versus control). In addition, Tukey’s multiple comparison test was also performed for multiple data sets.

In this report, we demonstrated that AML-MSCs are phenotypically and functionally different from normal donor-derived MSCs. We found that osteogenic lineage–associated genes were upregulated in AML-MSCs compared with N-MSCs. However, AML-MSCs were unable to differentiate into adipocytes. Coculture of AML cells with N-MSCs upregulated osteogenic marker expression in MSCs, suggesting that AML
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Figure 4. Human bone marrow implant model and patient AML-BM biopsy specimens validate AML-induced osteogenic differentiation in vivo. (A) Human femur-derived bone fragments were transplanted with Matrigel into the right flanks of NSG mice. Four weeks after implantation, vascularization of the HBMI bone fragments was confirmed by X-RAD 225 Cx cone-beam micro-CT scan using the contrast agent AuroVist 15 nm injected intravenously before image acquisition. Images were imported to MicroView software for analysis and segmentation. This representative image shows neovascularization of the bone fragment. Arrow points towards a mouse blood vessel leading to HBMI. (B) Bone fragments from HBMI mice transplanted (right) or not transplanted (left) with Molm13 cells were subjected to IHC staining for CD45 and osterix or CD45 and RUNX2 4 weeks after transplantation. The Opal multiplex system was used for signal amplification. CD45 was labeled with fluorescein (green), and RUNX2 and osterix were developed with Cy5 label (red). The nuclei were stained with DAPI (blue). Arrow points towards cells expressing osterix or RUNX2. Scale bar: 100 μm. (C) Quantification of osterix + or RUNX2 + stromal cells from HBMI mice transplanted or not transplanted with Molm13 cells. The absolute numbers of osterix +CD45– or RUNX2 +CD45– stromal cells from HBMI mice transplanted or not transplanted with Molm13 cells were quantitated by inForm image analysis software. Immunohistochemical analysis was performed for osterix (D) and RUNX2 (E) in normal (left) or AML patient-derived (right) BM specimens. The Opal multiplex system was used for signal amplification. CD45 staining was developed with fluorescein (green), osterix staining was developed with Cy3 (yellow), and RUNX2 staining was developed with Cy5 (red). Scale bar: 100 μm. Unpaired Student’s t test was used to test the statistical difference between the 2 groups (*P < 0.05). Arrow points towards cells expressing osterix or RUNX2.
Figure 5. BMP signaling mediates AML-induced osteogenic differentiation and induces CTGF expression in MSCs. (A) Immunoblotting was used to detect total and phosphorylated Smad1/5 in MSCs cultured with or without OCI-AML3–conditioned medium at indicated time points. (B) Immunoblotting also was used to detect total and pSmad1/5 in MSCs cultured with or without AML-CM in the presence or absence of type-1 BMP inhibitor LDN-212854 at indicated concentrations. β-Tubulin and β-actin served as loading controls. Data shown are representative of 3 independent experiments. (C) N-MSCs (2 × 10^5) were cultured in OCI-AML3–conditioned medium (OCI-AML3) with or without LDN-212854 for 5 days. mRNA expression of osteoprogenitor-associated genes was analyzed by qPCR and normalized to GAPDH. **p < 0.01. (D) Heatmap showing expression of various genes in AML-CM and AML-CM + LDN-212854 conditions. (E) mRNA expression of CTGF in N-MSCs cultured with or without LDN-212854. **p < 0.01. (F) mRNA expression of CTGF in OCI-AML3 cultured with or without LDN-212854. **p < 0.01.
cells induce osteogenic differentiation in MSCs. These findings were validated in vivo using an HBMI bone model. Osteogenic markers, including osterix and RUNX2, were upregulated in stromal cells in HBMI with leukemia compared to stromal cells in HBMI without leukemia. AML patient-derived BM biopsy samples also revealed higher osterix and RUNX2 expression compared with healthy BM. Mechanistic studies identified that AML cells secret BMPs and activate Smad1/5 phosphorylation in MSCs to induce osteogenic differentiation and CTGF expression in MSCs. Overexpressing CTGF in transgenic mice enhanced AML engraftment. These data demonstrate that AML cells induce osteogenic differentiation in BM-MSCs to gain growth advantage through growth factors secreted by osteogenic cells.

We previously demonstrated that AML-MSCs lack the ability to differentiate into adipocytes (22). However, their ability to differentiate into other lineages, including osteogenic or chondrogenic lineages, is not known. Here, we show that leukemia cells alter the BM microenvironment by inducing osteogenic differentiation in MSCs. In our previous report, we studied the effect of CTGF on MSCs but not on leukemia (22). Downregulation of CTGF induces adipogenic differentiation upon stimulation in vitro and spontaneously in vivo. These data support our current report that upregulation of CTGF in MSCs favors osteogenic differentiation but downregulation favors adipogenic differentiation. In our previous paper, we reported that AML cells home to BM with preexisting adipocytes compared with BM with fewer adipocytes (22). These data suggest that adipocytes could influence in AML homing to BM and engraftment. However, herein we show that AML cells influence MSCs in the BM to differentiate into osteoprogenitor cells. BMPs derived from AML cells activate Smad1/5 in MSCs and induce osteogenic differentiation. Once the cells become osteoprogenitor cells, they secret several factors, including CTGF, which supports leukemia growth. Several recent reports also suggest that the osteogenic BM niche affects AML growth. A report from Scadden’s lab showed that deletion of Dicer1 from osterix-expressing osteoprogenitor cells induced leukemia in mice (28). In another report, Kousteni’s group suggests that activating β-catenin mutations in osteoblasts induce AML in mice (29).

Very recently, Dong and Yu et al., reported that Ptpn11 mutations in osteoprogenitor and mesenchymal stem cells induced myeloproliferative neoplasm but that the same mutations in endothelial cells or mature osteoblasts did not induce this effect on myeloid cell compartment (30). These reports from different labs suggest that osteogenic niches support leukemia growth and disease progression.

Hanoun et al. reported that AML-BM contained higher numbers of osterix-expressing osteogenic cells than healthy BM in a syngeneic mouse model (31). Recent data also suggest osteoblast-mediated inhibition of AML growth (32). However, the authors used osteocalcin-positive mature osteoblasts in their model, which are different from osterix+RUNX2+ osteoprogenitor cells. These reports and our data demonstrate that AML cells induce and require an osteoprogenitor-rich niche, but not mature osteoblasts, for their expansion. In fact, mature osteoblasts are essential for normal hematopoiesis (33). In primary AML, the expression of both osterix and RUNX2 was substantially upregulated (Figure 4D). Taken together, our data and that of published reports demonstrate that AML cells prefer a preosteoblast/osteoprogenitor-rich niche for their growth and that they generate this niche in the BM microenvironment by inducing osteogenic differentiation in MSCs.

BMPs play a critical role in the induction of osteogenic differentiation during embryonic development (23). We found that AML-conditioned medium mediates Smad1 and -5 activation in MSCs. Recent reports indicate that BMPs are highly expressed in plasma samples from AML patients compared with those from healthy individuals (14, 15, 34), supporting our findings. Although it is not entirely clear which BMP proteins are involved, BMP6 and BMP9 have been shown to have the highest osteogenic activity among their family members (BMP1–15) (11) because of their association with Wnt signaling (12). In addition, BMP9 has been shown to directly regulate CTGF expression in MSCs (16), suggesting that it plays a functional role in AML-induced osteogenic differentiation in MSCs.
Figure 6. CTGF overexpression facilitates leukemia engraftment. (A) MLL-ENL-GFP-Nras-FFluci-p53−/− syngeneic mouse AML cells (1 × 10^5 per mouse) were transplanted intravenously into control (WT C57/B6; n = 3) or Colla2-CTGF–transgenic mice (CTGF tg; n = 3). Leukemic growth was measured by IVIS bioluminescence imaging at 8, 12, and 16 days after AML cell injection. (B) Total photon flux, representing the mean leukemia burden in the mice, was determined. (C) On day 16, peripheral blood mononuclear cells remaining after lysis of red blood cells were analyzed for GFP+ mouse leukemia cells by flow cytometry. (D) Sections of mouse spleen and liver from WT (left) or Colla2-CTGF–transgenic mice were subjected to H&E staining on day 16 after AML cell transplantation. Scale bar: 200 μm. (E) GFP immunofluorescence staining of spleens from WT mice (left) and Colla2-CTGF–transgenic mice (right) by the anti-GFP antibody on day 16 after AML cell transplantation. Scale bar: 200 μm. (F) Schematic representation of leukemia-stroma interaction in the BM microenvironment. Two-way ANOVA was used to test the data set in B and unpaired Student’s t test was used for the data set in C (** P < 0.01, ***P < 0.001 versus control). In addition, Tukey’s multiple comparison test was also performed for the data set in B.
CTGF expression has been reported to be upregulated during osteogenic differentiation of mesenchymal cells (26). CTGF has been shown to be a prognostic factor in acute lymphoid leukemia, and inhibition of its expression by an anti-CTGF antibody (FG-3019) led to increased survival of leukemia-bearing mice (19). In addition, CTGF has been shown as a critical factor for normal hematopoietic stem cell (HSC) maintenance and it has been shown that HSCs can upregulate CTGF expression in stromal cells (35). However, the role of CTGF in AML has not been elucidated. Our findings, for the first time to our knowledge, define the role of CTGF in AML cell growth. Our data indicate that AML cells, but not normal CD34+ cells, induce osteogenic differentiation through BMP/osterix/RUNX2-mediated signaling, leading to CTGF expression in MSCs, and that CTGF overexpression by stromal cells enhances leukemia engraftment in mice. Inhibition of BMP/CTGF-mediated signaling (Figure 6F) may inhibit AML growth and BM engraftment and could be developed into novel therapeutic concept in AML.

Methods

Isolation and culture of cells. MSCs were isolated from the BM of normal healthy donors undergoing BM harvest for use in allogeneic transplantation or AML patients. Each BM specimen was subjected to centrifugation (700 g for 20 minutes at 4°C) over a Ficoll-Hypaque gradient (Sigma-Aldrich) to separate mononuclear cells. After centrifugation, the buffy coat layer was carefully extracted; resuspended in α-MEM containing 10% FBS, L-glutamine, 4% pooled human platelet lysate (36), and 1% penicillin-streptomycin (MSC growth medium); and plated at an initial density of 5 × 10^4 cells/cm². After 3 days, the cultures were washed with PBS (Media Tech Inc.), and the remaining adherent cells were cultured until approximately 80% confluent. The cells were then subcultured at densities of 5,000–6,000 cells/cm². Only third or fourth passage cells were used for the experiments. AML cell lines, OCI-AML3 (gift from Mark Minden, Ontario Cancer Institute, Toronto, Canada) and Molm13 (Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH), were cultured in RPMI medium (Media Tech Inc.) with 10% FBS and 1% penicillin-streptomycin. CD34+ cells were isolated from human CB samples using magnet-assisted cell sorting (Miltenyi Biotech) as per the manufacturer’s instructions.

Cell growth/cell cycle analysis. Cell growth was assessed by counting live cells on daily basis. AML- or N-MSCs (n = 5 each; 3 × 10⁴ cells/well) were plated in 6-well culture dishes in MSC growth medium. Cells were trypsinized and counted daily for 6 days on an automated live-cell counter (Vi-CELL, Beckman Coulter). The data were analyzed in an Excel spreadsheet.

BrdU pulse labeling assay. AML- and N-MSCs (n = 3 each) were pulsed with 10 μM BrdU solution for 3 hours in MSC growth medium. The cells were washed, subjected to trypsinization, and fixed in ice-cold ethanol overnight. Following fixation, DNA was denatured by incubating cells in 2 N HCl/0.5% Triton X-100 for 30 minutes at room temperature and then neutralized with 0.1 M Na₂B₄O₇. The cells were then incubated with N-MSCs (n = 5 each; 3 × 10⁴ cells/well) were plated in 6-well cell culture dishes in MSC growth medium.

Flow cytometry. Flow cytometry analysis of MSCs isolated from normal donor–derived and AML patient–derived BM specimens was performed as described before (22). MSCs isolated from normal donor–derived and AML patient–derived BM specimens were subjected to trypsinization and washed once with PBS. The cells were then incubated for 20 minutes with 10 μl of fluorochrome-conjugated antibodies. The following antibody conjugates were used: anti-CD45, anti-CD105, anti-CD140b, anti-CD146, anti-CD73, anti-TNAP (clone W8B2), and anti-CD140a were conjugated with phycoerythrin (PE); anti-CD271 and anti-CD51 were conjugated with FITC; anti-CD44, anti-SUSD2, and anti-CD106 were conjugated with APC; and anti-CD31 was conjugated with APC/cyanine dye Cy7 (all purchased from BioLegend). Anti-CD90 antibody conjugated with APC–Alexa fluor 700 was purchased from Beckman Coulter. After incubation, the cells were washed once with PBS containing 0.5 μg/ml DAPI (to exclude dead cells) and analyzed on an LSR-II Flow Cytometer. Ten thousand events were acquired for each sample. The flow cytometric data were analyzed by FlowJo software.

Multilineage differentiation. For experiments related to osteogenic and adipogenic differentiation, we randomly selected AML and normal MSCs. Samples 5, 6, 11, and 22 from AML-MSCs and samples 31, 34, and 36 from normal MSCs (Supplemental Table 1) were used for differentiation induction or coculture experiments. Multilineage differentiation potential of the AML- and N-MSCs was investigated as described before (22, 37).
To induce osteogenic differentiation, 10,000 MSCs isolated from normal donor– or AML patient–derived BM were cultured with MSC growth medium (control) or with NH OsteoDiff medium (Miltenyi Biotec) for 21 days in 6-well dishes. The medium was replaced every 3 days. After 21 days, the cells were washed twice with PBS and fixed with 4% paraformaldehyde (PFA). To determine ALP activity, the cells were incubated with FAST BCIP/NBT substrate (Sigma-Aldrich) for 20 minutes at room temperature. Calcium deposition was analyzed by staining with 1% Alizarin Red S (Spectrum) for 20 minutes at room temperature. After the cells were washed, they were photographed by a Nikon Coolpix-950 camera attached to a Nikon-TMS light microscope.

To measure adipocyte differentiation, MSCs isolated from normal donor– or AML patient–derived BM were cultured in MSC growth medium (control) or adipocyte differentiation medium containing DMEM, 1 μM dexamethasone, 50 μM indomethacin, 500 nM IBMX, 5 μg/ml insulin, and 10% fetal calf serum (FCS) in a 12-well cell culture dish. The medium was replaced every 3 days. After 21–28 days, the formation of adipocytes was evaluated by fixing cells with 4% PFA and staining with Oil Red O dye (Sigma-Aldrich) for 15 minutes at room temperature.

To induce differentiation into the chondrocyte lineage, the cells were incubated in ChondroDiff medium (Miltenyi Biotec). Briefly, cells were washed once with PBS and plated into a u-bottom 96-well plate with 250,000 MSC cells/well. The medium was replaced every 3 days. After incubation for 28 days, the resulting cell pellets were fixed with 10% formalin, embedded in paraffin, and cut into 5-μm sections. Following deparaffinization and hydration, the sections were incubated with 1% Alcian Blue 8GX (w/v) solution (Sigma-Aldrich) in 3% acetic acid for 30 minutes at room temperature. The slides were washed in 3% acetic acid and then in distilled water. Photographs were taken by an Olympus DP72 digital camera attached to a BX43 microscope (Olympus America Inc).

**Alkaline phosphatase and mineralization quantification.** ALP activity was quantified by using the colorimetric Alkaline Phosphatase Assay Kit (Abcam) as per the manufacturer’s instructions. Mineral deposition by cultured cells was quantified directly from Alizarin Red S staining as previously described (38). Briefly, cells were trypsinized, washed once with PBS, and subjected to lysis in assay buffer. The assay was performed by adding 5 mM p-nitrophenol phosphate substrate and sample lysis buffer to cells in a 96-well flat-bottom plate. A standard curve was prepared by diluting 1 mM p-nitrophenol phosphate stock with appropriate amounts of assay buffer. The standards ranged from 0.015 to 0.15 μmol/ml. After incubation for 1 hour at room temperature, absorbance was measured at 405 nm by using Infinite 200 Pro (Tecan). A standard curve of absorbance versus p-nitrophenol concentration was plotted and used to determine the amount of p-nitrophenol generated by each sample, representing ALP activity.

The dye was extracted by incubating cells with 10% (v/v) acetic acid for 30 minutes at room temperature with shaking. The monolayer was scraped with a cell scraper and transferred to a 1.5-ml microcentrifuge tube. After vortexing for 30 seconds, the lystate was sealed with parafilm, heated to 85°C for 10 minutes, and immediately cooled on ice for at least 5 minutes. The lysate was then subjected to centrifugation at 20,000 g for 15 minutes. The supernatant was transferred to a new tube and neutralized with 10% ammonium hydroxide to reach pH between 4.1 and 4.5 before aliquoting to a 96-well plate. The standards ranged from 0.05 to 0.4 μmol/ml. Absorbance was read at 405 nm with an Infinite 200 Pro.

**Quantitative RT-PCR.** Unless otherwise mentioned, all qRT-PCR analyses were performed as SYBR-green PCR analysis. Total RNA was extracted by the standard Trizol-based method described elsewhere (22). The cDNA was prepared by using the Improm-II reverse transcriptase system (Promega). The SYBR-green PCR primer sequences and taqman assays used for experiments in this report are listed in Supplemental Table 2 and Supplemental Table 3, respectively. The Fast-SYBR-green or Fast-taqman master consisting of DNA polymerase and a nucleotide mix was purchased from Applied Biosystems (now ThermoFisher Scientific). An HT900 real-time PCR instrument (ThermoFisher Scientific) was used to perform the PCR reaction and acquisition of signal amplification.

**Human BM implant mouse model.** Human femur heads were obtained from leukemia-free patients undergoing hip replacement surgery at St. Luke’s Hospital (Houston, Texas, USA). NOD/SCID/IL-2rnull (NSG) mice were purchased from the Jackson Laboratory. Eight- to 10-week-old NSG mice were surgically implanted with the fresh human femur fragments as a model of normal human BM. The mice were irradiated (150 Gy) the day before the surgery, and the fresh bone fragments were irradiated (250 Gy) the day of the surgery to avoid graft-versus-host disease in the recipient mice. The implantation was done under sterile conditions. Before the surgery, animals were anesthetized with isoflurane (2.5%) and the mice were shaved...
on the dorsal side until the skin was clearly visible. Two 1-cm incisions were made in the right dorsal flank of each mouse using a sterile sharp scalpel, and human bone fragments immersed in Matrigel (ECMatrix, Millipore) were carefully implanted subcutaneously into these incisions. The incisions were then closed with surgical clips. The animals were maintained on a Bactrim diet (Harlan Laboratories Inc.) for up to 2 weeks following surgery. The bone implants were allowed to engraft in the mice for 4 to 12 weeks. For the leukemia engraftment assays, leukemia cells were introduced through the tail vein 4 weeks after bone implantation. The experiment was repeated 3 times using 5 mice per group.

**CT imaging.** The vascularization of the newly implanted human bone fragments in the HBMI mice was assessed by imaging on a X-RAD 225 Cx cone-beam micro-CT (Precision X-Ray) using the commercially available contrast agent AuroVist 15 nm (Nanoprobes) as described elsewhere (39). The injected doses of blood-pool agents were selected following the instructions provided by the manufacturer. Both before and after contrast scans were taken. The contrast material was injected continuously via a polyethylene catheter though the tail vein in a 200-μl solution per 25 g mouse (40 μg/mouse), with an injection time of 2 minutes. Scan tube voltage was 40 kVp and current was 5 mA. For each scan, 360 projections were acquired at 1-degree increments and reconstructed using a filtered back-projection algorithm, resulting in an image volume with isotropic 150-μm voxels. Images were imported into the MicroView system (Parallax Innovations) for analysis and segmentation.

**Bone generation by human bone implants.** New bone generation was assessed as described elsewhere (40). Briefly, the animals were injected intravenously with the targeted fluorescent in vivo bisphosphonate imaging agent OsteoSense 750EX (PerkinElmer; 100 μl/mouse), which images bone growth, reabsorption, remodeling, and microlcifications (41). As a control, mice without human bone implants were also injected with OsteoSense 750. The mice were imaged by using the IVIS (PerkinElmer) system, and the imaging data were analyzed by Living Image software (PerkinElmer).

**MSC generation from human bone implants.** The HBMI mice were killed 4 weeks after bone implantation, and the bone implants were removed and cut into 0.5-cm³ cubes. These pieces of recovered bone were washed once with PBS and incubated in a combination of dispase (0.5 mg/ml; Stem Cell Technologies) and collagenase (1 mg/ml; Sigma-Aldrich) at 37°C for 60 minutes in a shaking incubator. After incubation, the bone pieces were washed twice with DMEM containing 10% FBS. The HBMI-derived cells were subjected to red cell lysis buffer (Sigma-Aldrich) at 10 minutes to remove any residual erythrocytes. After incubation, the cells were washed twice with PBS and plated in MSC growth medium. The MSCs generated were isolated by FACS using anti-human CD140b and anti-human CD90 markers to exclude any mouse-derived MSCs.

**Conditioned medium incubation and Western blotting.** To investigate the induction of BMP signaling in MSCs by leukemic cells, AML-conditioned medium was prepared by seeding OCI-AML3 cells (0.3 × 10⁶ cells/ml) in RPMI medium with 2% FCS. The cells were harvested on day 4, and the supernatants were collected and filtered through 0.2-μm membranes (Millipore). N-MSCs, pretreated for 1 hour with BMP type-1 receptor inhibitor LDN-212854 (Sigma-Aldrich) at different concentrations were incubated in the N-MSC membranes (Millipore). N-MSCs, pretreated for 1 hour with BMP type-1 receptor inhibitor LDN-212854 (Sigma-Aldrich) at different concentrations were incubated in the conditioned medium for 1-2 hours. The conditioned medium was collected and used for Western blotting. The conditioned medium was collected and used for Western blotting. The proteins were then transferred to Immobilon-FL membranes (Millipore). The membranes were incubated with rabbit anti-human phospho (p) Smad1/5 or anti-Smad1 or anti-Smad5 antibodies (all from Cell Signaling Technologies) and mouse anti-human β-tubulin antibody (Sigma-Aldrich) overnight at 4°C. They were then washed and incubated with donkey anti-rabbit or anti-mouse IgG secondary antibody conjugated with Alexa fluor 680 or Alexa fluor 800 (both from LI-COR Biosciences). The membranes were washed again and scanned using the Odyssey fluorescence imaging system (LI-COR Biosciences).

**Leukemia engraftment.** To investigate leukemia engraftment in human bone implants, we transplanted Molm13 cells (110⁶) expressing GFP and firefly luciferase through the tail vein in a group of HBMI mice (n = 5). Another group of HBMI mice (n = 5) that did not receive leukemia cells served as controls. The leukemia engraftment was measured weekly by IVIS whole-body imaging (Caliper Life Sciences) and peripheral blood analysis by flow cytometry. Human leukemia engraftment in the peripheral blood was assessed by positive staining for anti-human CD45-PE antibody (Biolegend) and negative staining for anti-mouse CD45-APC. Leukemia engraftment in HBMI mice was also assessed by H&E staining (Sigma-Aldrich) and by IHC staining for anti-human CD45 and firefly luciferase.
The Col1a2-CTGF–transgenic mice used in this study have been described previously (42). Briefly, the murine homolog of CTGF (Fisp-12) was cloned into a vector containing the 6-kb enhancer and minimal promoter of the murine Col1a2 gene, an IRES-lacZ reporter, and the murine protamine polyA signal. Transgenic mice were produced by the standard method of pronuclear injection of linear DNA into a fertilized mouse egg. Genotyping was performed by PCR for the lacZ reporter on DNA extracted from ear biopsies. Mouse hematopoietic cells transduced with MLL-ENL-GFP Nras-FFLuci p53− (gift of Scott Lowe) were introduced into a group of CTGF-transgenic and WT (C57/B6) mice via tail vein. Bioluminescent imaging was performed at different time points by injecting D-luciferin (4 mg/mouse in 100 μl). The leukemia engraftment was measured weakly by whole-body imaging using IVIS (Caliper Life Sciences) and peripheral blood by flow cytometry analysis. Leukemia engraftment in the peripheral blood was assessed by GFP+ leukemia cells. Leukemia engraftment was also assessed by immunohistochemical staining using anti-GFP antibody.

Immunohistochemical analysis. After substantial human leukemia engraftment (>30% circulating human CD45+ cells), mice were killed and the human bone implants were removed and fixed in 4% PFA. The bone fragments were decalcified and the remaining tissues were embedded in paraffin and cut into thin sections (5 μm) onto glass slides using a microtome. The slides were stained with H&E and analyzed by light microscopy. For IHC staining, the tissue sections were deparaffinized and gently boiled in 1′ antigen retrieval buffer (DAKO) in a microwave oven for 15 minutes at 20% power. The slides were then washed and incubated for 10 minutes in serum-free blocking solution (DAKO) and then for 1 hour with the primary antibody. The tissue sections were washed 3 times with TBST and stained with horseradish peroxidase–conjugated species-specific secondary antibodies. The slides were then washed again and developed with Opal fluorescein and Cy5 reagents (Perkin Elmer) per the manufacturer's guidelines. The slides were imaged with a Vectra multispectral imaging system (Perkin Elmer), and the data were processed by inForm image analysis software (Perkin Elmer). Leukemia engraftment in CTGF-transgenic mice was analyzed by using Alexa fluor 488–conjugated anti-mouse secondary antibody (Life Technologies) instead of Opal.

Gene expression profiling. To investigate differences in gene expression among AML-associated MSCs and healthy MSCs, murine AML cells carrying different gene deletions/rearrangements were transplanted in C57/B6 mice. These primary leukemia cells were generated by introducing the desired oncogenic translocations, including p53-knockout C57BL/6 mice (AML1/ETO9a, MLL/ENL, MLL/ENL+FLT3-ITD) or p53 wt C57BL/6 mice (MLL/ENL+FLT3-ITD p53 wt). In vivo expanded genetically modified leukemia cells were transplanted into 10 mice (6 to 8 week old) by tail vein injection of 1 × 106 cells/recipient. PBS was injected in the control group. Mice were sublethally irradiated prior to the injections to favor a uniform disease onset in recipient animals. Leukemia engraftment and burden were subsequently monitored by bioluminescence imaging (IVIS system; Xenogen/Caliper Life Sciences).

Transplanted leukemia was allowed to progress for 2 (MLL/ENL and MLL/ENL+FLT3-ITD), 3 (AML1/ETO9a), or 4 weeks (MLL/ENL+FLT3-ITD p53 wt) according to disease burden. At this time, mice were sacrificed and BM-MSCs from mouse BM were isolated by FACS using a combination of the following cell surface markers: Ter119−, CD31−, CD45−, CD105+, Sca1+, CD106+, and PDGF-Rα+ α mice were sacrificed and BM-MSCs from mouse BM were isolated by FACS using a combination of the following cell surface markers: Ter119−, CD31−, CD45−, CD105+, Sca1+, CD106+, and PDGF-Rα+. The bone fragments were decalcified and the remaining tissues were embedded in paraffin and cut into thin sections (5 μm) onto glass slides using a microtome. The slides were stained with H&E and analyzed by light microscopy. For IHC staining, the tissue sections were deparaffinized and gently boiled in 1′ antigen retrieval buffer (DAKO) in a microwave oven for 15 minutes at 20% power. The slides were then washed and incubated for 10 minutes in serum-free blocking solution (DAKO) and then for 1 hour with the primary antibody. The tissue sections were washed 3 times with TBST and stained with horseradish peroxidase–conjugated species-specific secondary antibodies. The slides were then washed again and developed with Opal fluorescein and Cy5 reagents (Perkin Elmer) per the manufacturer's guidelines. The slides were imaged with a Vectra multispectral imaging system (Perkin Elmer), and the data were processed by inForm image analysis software (Perkin Elmer). Leukemia engraftment in CTGF-transgenic mice was analyzed by using Alexa fluor 488–conjugated anti-mouse secondary antibody (Life Technologies) instead of Opal.

Gene expression analysis. Differences in TNAP expression were compared by subjecting MFI ± SEM values from N- or AML-MSCs to ANOVA test using the Prism software. This method was also used to compare data from RT-PCR gene expression studies and enzyme activity assays. P values of less than 0.05 were considered statistically significant. One-way or two-way ANOVA was used for comparisons of 3 or more groups, and unpaired 2-tailed Student’s t test was used for comparisons of 2 groups. In addition, Dunnett’s or Tukey’s multiple comparison test was used for multiple groups. For comparison of osterix+CD45− or RUNX2+CD45− stromal cells in HBMI mice stimulated with leukemia cells or not stimulated, cell-level marker intensity was dichotomized relative to the median intensity in the population. Thus, every cell was assigned high or low marker status based on its intensity relative to the median. For each of the markers (osterix, CD45, and Runx2), the number of cells that were in each
subgroup (Runx2\(^{-}\)CD45\(^{+}\); Runx2\(^{-}\)CD45\(^{-}\); Runx2\(^{+}\)CD45\(^{-}\); and Runx2 CD45\(^{-}\)) were counted. Similarly, for the Osterix\(^{-}\) and CD45\(^{-}\) populations, a test of proportions was used to measure the difference in the population of Runx2\(^{-}\)CD45\(^{+}\) cells between the 3 groups relative to the control (without Molm13). This procedure was repeated for the osterix\(^{-}\)CD45\(^{-}\) comparison as well.

**Study approval.** All experiments performed with the mice were in compliance with the standards of care of the MD Anderson Cancer Center Institutional Animal Care and Use Committee as well as in accordance with a protocol approved by the MD Anderson Cancer Center Institutional Animal Care and Use Committee. All BM and bone donors provided written informed consent. Studies involving human materials were conducted according to a protocol approved by the Institutional Review Boards of St. Luke’s and MD Anderson Cancer Center.

**Author contributions**

VLB conceptualized and planned the study, conducted experiments, interpreted the data, and wrote the manuscript. PML and JCS performed the experiments and analyzed the data. KN, BY, XZ, SS, TM, VR, KAM, and XL performed the experiments. ES provided patient samples. ZW performed gene expression array experiments and the data was analyzed by RJ and RED. AR analyzed statistical data. GA provided patient samples. MK interpreted the data. MAH and CWC provided human bone samples. CBR provided samples and interpreted the data. MA planned the study and interpreted the data.

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