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Pbx1 Represses Osteoblastogenesis by Blocking Hoxa10-Mediated Recruitment of Chromatin Remodeling Factors

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Abdominal-class homeodomain-containing (Hox) factors form multimeric complexes with TALE-class homeodomain proteins (Pbx, Meis) to regulate tissue morphogenesis and skeletal development. Here we have established that Pbx1 negatively regulates Hoxa10-mediated gene transcription in mesenchymal cells and identified components of a Pbx1 complex associated with genes in osteoblasts. Expression of Pbx1 impaired osteogenic commitment of C3H10T1/2 multipotent cells and differentiation of MC3T3-E1 preosteoblasts. Conversely, targeted depletion of Pbx1 by short hairpin RNA (shRNA) increased expression of osteoblast-related genes. Studies using wild-type and mutated osteocalcin and Bsp promoters revealed that Pbx1 acts through a Pbx-binding site that is required to attenuate gene activation by Hoxa10. Chromatin-associated Pbx1 and Hoxa10 were present at osteoblast-related gene promoters preceding gene expression, but only Hoxa10 was associated with these promoters during transcription. Our results show that Pbx1 is associated with histone deacetylases normally linked with chromatin inactivation. Loss of Pbx1 from osteoblast promoters in differentiated osteoblasts was associated with increased histone acetylation and CBP/p300 recruitment, as well as decreased H3K9 methylation. We propose that Pbx1 plays a central role in attenuating the ability of Hoxa10 to activate osteoblast-related genes in order to establish temporal regulation of gene expression during osteogenesis.

Embryonic patterning of the skeleton is a complex process responsible for regulating the positional identity, shape, and size of skeletal primordial elements by an intricate orchestration of signaling molecules and transcriptional regulators. The most widely studied transcriptional regulators governing embryonic patterning are the Hox homeodomain DNA-binding proteins. Several emerging lines of evidence have demonstrated that these Hox proteins play a wider role, extending their functions from patterning the developing embryo to regulating formation and maintenance of mature tissues such as bone (13, 18, 19, 21). In adults, bone turnover is a continual process of resorption and renewal that depends on the commitment of mesenchymal progenitors to the osteoblast lineage. This process is mediated by a variety of extracellular signals that are also necessary for embryonic skeletal development, including canonical Wnt signals and bone morphogenetic proteins (BMPs) (5, 12, 27). Homeodomain proteins, including the abdominal-class Hox family, have been identified as downstream targets and regulators of osteogenic BMP signaling (23, 25, 26). One of the more compelling questions regarding Hox function is the mechanism(s) by which these proteins regulate transcription in a cell-specific manner to promote mature tissue regeneration.

Several abdominal-class Hox proteins are upregulated in bone repair, including Hoxa2, Hoxd9, Hoxa11, and Hoxa13 (15, 24), suggesting that the embryonic program is reactivated during new bone formation. For example, Hoxa10 is strongly induced in osteoblasts immediately following BMP2 treatment, coincident with the robust expression of Runx2, the master regulator of osteoblast commitment (2). Hoxa10 has been shown to bind to several phenotypic bone promoters at Hox consensus sequences and to enhance osteocalcin (bglap2; Ocn) transcriptional activity in reporter assays in vitro (19). Although the exogenous expression of Hoxa10 can induce the transcription of several bone genes in normal osteoblasts, Hoxa10 binds to chromatin of endogenous bone genes preceding the binding of Runx2 (19). Hoxa10 has been postulated to mark chromatin for activation by allowing Runx2 binding and subsequent gene expression. It is important to note that during osteoblast differentiation phenotypic genes are temporally expressed, and this sequential pattern of expression suggests that several chromatin remodeling factors are required for osteoblastic gene activation. Perhaps due to the critical but varied roles of Hox proteins, a large complement of coregulatory molecules serve as coactivators or corepressors of Hox-mediated transcriptional activity, functioning by direct protein-protein interactions or by modulating Hox DNA-binding activity (31).

One of the best characterized groups of Hox-interacting proteins is the three-amino-acid loop extension (TALE) family of homeodomain-containing transcription factors, which includes the pre-B-cell leukemia homeobox protein 1 (Pbx1). Like Hox proteins, Pbx1 plays a critical role in patterning of the axial skeleton, with deletion of the gene resulting in severe...
skeletal malformations due to defects in chondrocyte matura-
tion (40). Pbx1 regulation of skeletal development has been
shown to be both dependent and independent of Hox protein
involvement (6, 31). Several TALE proteins are closely related, but loss of Pbx2 or Pbx3 does not confer a skeletal phenotype
(36, 41); thus, Pbx1 may have a distinct role during skeletal
development. Furthermore, mapping of polymorphisms re-
lated to bone mineral density in Chinese populations revealed
a strong correlation between nucleotide variation in the
Pbx1 gene and decreased bone mass (10). It has been suggested
that during mesenchymal cell commitment to the myogenic lineage,
Pbx proteins may act as “pioneer” transcription factors that
penetrate repressive chromatin and mark specific genes for
activation by MyoD (38). In this study we addressed the func-
tional role of Pbx1 in osteogenesis and the Hoxa10-dependent
activation of the osteocalcin and bone sialoprotein (Bsp) genes
as classical models for osteoblast-specific gene expression.

MATERIALS AND METHODS
Cell culture. Human embryonic kidney cells (HEK-293T), mouse preoste-
blasts (MC3T3-E1), and mesenchymal cells (CH110T1/2) were obtained from
ATCC (Manassas, VA). HEK-293T and MC3T3-E1 cells were maintained in
minimum essential medium alpha medium (αMEM) (Invitrogen/Gibco Carls-
bad, CA) supplemented with 10% fetal bovine serum (FBS) (HyClone, Logan,
Utah, UT). MC3T3-E1 and mesenchymal cells (CH110T1/2) were maintained in
Dulbecco’s modified Eagle’s medium (DMEM) (Invitrogen/Gibco) containing
2 mM l-glutamine, and Pen/Strep. C3H10T1/2 cells were maintained in Dulbecco’s
modified Eagle’s medium (DMEM) (Invitrogen/Gibco) containing 10% FBS, 2 mM l-glutamine,
and Pen/Strep. Primary rat osteoprogenitors were isolated from embryonic (day
21) rat calvaria as previously described (16). Osteoprogenitor cells were main-
tained in Dulbecco’s minimal essential medium (MEM) (Invitrogen/Gibco) con-
taining 10% FBS, 2 mM l-glutamine, and Pen/Strep. To induce osteogenic
differentiation, media were further supplemented with BMP2 (100 ng/ml) gener-
ously provided by John Wozney [Wyeth Research, Women’s Health and Mus-
culoskeletal Biology, Cambridge, MA)] and/or 280 μM ascorbic acid and 10 mM
β-glycerophosphate (BGP) (Sigma Aldrich, St. Louis, MO). Media were re-
placed every 2 days for the duration of all experiments. All cells were maintained
at 37°C in a humidified 5% CO2 environment.

Expression constructs and short hairpin RNA (shRNA) virus generation.
The Xpress-Pbx1 plasmid used in this study was previously described (19). The
Flag-Pbx1 construct was generated by excising the Pbx1 fragment from pCMV-
Pbx1 (generously provided by Corey Largman, University of California, San
Francisco) by BamHI digestion followed by ligation into linearized pCDNA3
(Invitrogen). To create the Pbx1-pSuper construct used for lentiviral packaging,
a 938-bp region of the rat bone sialoprotein proximal promoter region was
previously described (34) (generously provided by the late Jaro Sodek, University
of Toronto). Alteration of the Pbx or Hox DNA consensus sequence was per-
formed by using a QuikChange site-directed mutagenesis kit (Stratagene, Cedar
Creek, TX). Cells were transfected with 500 ng of reporter plasmid and the
indicated concentrations of Hoxa10, Pbx1, or pCDNA 3.1 expression vector using
FuGene6 transfection reagent. Following the harvest of DNA from the
lentiviral packaging cell line (Roche), At 16 to 24 h posttransfection, cells were scraped into PBS, collected by centrifugation, and lysed with 5X passive lysis buffer (Promega, Madison, WI).

ChIP assays.

Immunoblotting.

RNA isolation and RT-qPCR. Total RNA was isolated using Trizol reagent
(Invitrogen) according to the manufacturer’s specifications. DNase I-treated
total cellular RNA was primed with oligo(dT) and reverse transcribed to cDNA
during mesenchymal cell commitment to the myogenic lineage,
using the Superscript first-strand cDNA synthesis kit (Invitrogen) accord-
ing to the manufacturer’s instructions. Gene expression was determined by quan-
titative real-time reverse transcription-PCR (RT-qPCR) using Power SYBR
green PCR master mix (Applied Biosystems Inc., Foster City, CA) and gene-
specific primers in an ABI Prism 7000 thermocycler. For each gene, expression
levels were normalized to GAPDH (glyceraldehyde-3-phosphate dehydro-
genase). Experiments were performed in triplicate and results displayed
as mean values ± standard error of the mean (SEM).

Immunoblotting. Cells were lysed in direct lysis buffer (2% SDS, 10 mM
dithiothreitol [DTT], 10% glycerol, 12% urea, 10 mM Tris-HCl [pH 7.5], 1 mM
phenylmethylsulfonyl fluoride [PMSF], 1X protease inhibitor cocktail [Roche],
25 μM MG132 [proteasome inhibitor]) and boiled for 5 min. Samples were
quantified, and equal amounts of protein were resolved by SDS-PAGE. Proteins
were transferred to a polyvinylidene difluoride membrane and were subjected to
immunoblotting with the appropriate antibodies. Immune-reactive proteins
were detected using Western Lightning chemiluminescence reagent (Perkin-Elmer,
Boston, MA).

ChIP assays. Chromatin immunoprecipitation (ChIP) assays were performed as
previously described (17). Briefly, formaldehyde cross-linking was performed for
15 min, and cells were washed once in 1X PBS and then lysed in lysis buffer (25
mM Tris-HCl [pH 8.0], 5 mM MgCl2, 10 mM EDTA, 1% SDS, 1% Triton X-100,
162.5 mM NaCl, 25 μM MG132, and 1X Complete protease inhibitor). Lysates
were sonicated to obtain DNA fragments with an average size of 0.2 kbp to 0.6
kbp. In some experiments, chromatin was treated with PstI restriction endonu-
clease (NEB, Danvers, MA) for 2 h at 37°C. Immunoprecipitations were per-
formed with the appropriate antibodies or immunoglobulin G (IgG) as a control.
Immunocomplexes were collected, followed by recovery of DNA using the
ChIPplex Kit (Upstate/Millipore, Billerica, MA).

Sequential ChIP studies were performed using the primary pulldown from
one antibody, which was divided into equal aliquots for the second pulldown with
antibodies specific for coregulatory molecules. Instead of the elution step (1%
formic acid), antibodies specific for coregulatory molecules were used to

acetylated histone H3 dimethyl lysine 9 (ab9050; Abcam), anti-histone H4

acetylated lysine 16 (ab1762; Abcam, Cambridge, MA), anti-HDAC2 (C-19,
sc-6296; Santa Cruz), anti-HDAC6 (L-18, sc-5258; Santa Cruz), and anti-histone

H3 histone (ab9505; Abcam).

Immunohistochemistry. Long bones of 2-day-old mice were fixed with PLP
fixative (2% paraformaldehyde, 0.075 M lysine, 0.037 M sodium phosphate,
1.0 M bis (H13003)

mM, pH 7.4) for 48 h. The fixed long bones were then demineralized by
sequential treatment with 18% EDTA (disodium salt dihydrate, pH 7.4) for 14
days, dehydrated, and embedded in paraffin using standard procedures. Paraffin-
embeded tissues were sectioned (5 μm), adhered to glass slides, and rehydrated,
and antigens were recovered by treatment with retrieval buffer (1 mM Tris, 0.5
mM EGTA, pH 9.0). Sections were blocked with phosphate-buffered saline
supplemented with 1% bovine serum albumin, 0.05% saponin, and 0.2% gelatin.
Serial sections were incubated overnight at 4°C with anti-Pbx1 (Cell Signaling),
anti-Hoxa10 (N-20), or anti-Runx2 (M-70) diluted (1:1000 in phosphate-buffered
saline (PBS) supplemented with 0.1% bovine serum albumin and 0.3% Triton
X-100. Labeling was visualized with species-specific horseradish peroxidase-
detected secondary antibody (P448, 1:200; Dako). Sections were counter-
stained with methyl green (0.5% methyl green, 0.1 M sodium acetate trihy-
drate, pH 4.2) or toluidine blue (0.1% toluidine blue, 7% ethanol, 150 mM
sodium chloride, pH 2.3).

Transfection and reporter assays. A 1.1-kbp DNA fragment encompassing the
proximal, tissue-specific promoter of the rat osteocalcin gene was inserted into
pGL3 basic 3 vector containing a 938-bp region of the rat bone sialoprotein proximal promoter region was
previously described (34) (generously provided by the late Jaro Sodek, University
of Toronto). Alteration of the Pbx or Hox DNA consensus sequence was per-
formed by using a QuikChange site-directed mutagenesis kit (Stratagene, Cedar
Creek, TX). Cells were transfected with 500 ng of reporter plasmid and the
indicated concentrations of Pbx1, Pbx2 or pCDNA 3.1 expression vector using
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ChIP assays. Chromatin immunoprecipitation (ChIP) assays were performed as
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mM Tris-HCl [pH 8.0], 5 mM MgCl2, 10 mM EDTA, 1% SDS, 1% Triton X-100,
162.5 mM NaCl, 25 μM MG132, and 1X Complete protease inhibitor). Lysates
were sonicated to obtain DNA fragments with an average size of 0.2 kbp to 0.6
kbp. In some experiments, chromatin was treated with PstI restriction endonu-
clease (NEB, Danvers, MA) for 2 h at 37°C. Immunoprecipitations were per-
formed with the appropriate antibodies or immunoglobulin G (IgG) as a control.
Immunocomplexes were collected, followed by recovery of DNA using the
ChIPplex Kit (Upstate/Millipore, Billerica, MA).
SDS, 1.1% Triton X-100, 1.2 mM EDTA, 167 mM Tris-HCl [pH 8.1], 167 mM NaCl and used for the secondary immunoprecipitations.

Aliquots of each recovered DNA sample were assayed by quantitative PCR to detect the proximal Ocn promoter region spanning bp –198 to –28 upstream of the transcription start site. As a control for nonspecific binding/proper DNA fragmentation, a region of the Ocn distal promoter from bp –1377 to –1184 was also analyzed. The oligonucleotides used were as follows: proximal promoter, 5’-GCG AGC CTC TGA TTG TGT CC-3’ (–198 to –179) and 5’-ATT ATC CAC TCG CTG AGG GG-3’ (–47 to –28); Pbx site, 5’-GTC TCT AGG GCC AGC TAC GGC TAC TGG GTG CGT CCT GAC AT-3’ (–112 to –91) and 5’-GCC CCG AGT TGG TCT GTG GGA-3’ (–5 to +15); and distal promoter, 5’-TAG TGT CTC CAG GCC AGC AGA-3’ (–1377 to –1356) and 5’-AAA TCT GCA GCC GTT CCC CCA GT-3’ (–1207 to –1184). Samples were normalized to the initial input and expressed as percent chromatin pulldown (compared to input).

**Immunoprecipitations.** Initially, protein A/G plus agarose beads (Santa Cruz) (100 µl; 1:1 beads/PBS) were noncovalently complexed with 20 to 100 µg of specific antibody. Unbound antibodies were washed away with phosphate-buffered saline (PBS), and antibody-bound protein A/G agarose was resuspended in dimethyl sulfoxide (DMSO) and added to the antibody-protein A/G solution at a final concentration of 1 mM. The cross-linking reaction was performed at room temperature for 1 h. Excess DSS was removed by washing the resin four times with 400 µl of Tris-buffered saline (TBS) (50 mM Tris, 150 mM NaCl, pH 7.2), four times with 0.1 M glycine (pH 2.8) to remove free antibody, and finally three times with TBS. The cross-linking efficiency was evaluated by A280. Approximately 10⁷ cells per immunoprecipitation were lysed in 500 µl of Nonidet P-40 (NP-40) lysis buffer (150 mM NaCl, 50 mM Tris [pH 8.0], 1% NP-40, 1× Complete protease inhibitor [Roche], 25 µM MG132 [Sigma Aldrich]) for 15 min at 4°C. Cell lysates were sonicated, followed by centrifugation at 16,000 × g for 15 min at 4°C. The supernatant was transferred to a clean microcentrifuge tube and precleared with 40 µl of antibody-protein A/G plus agarose complex was added and incubated at 4°C with agitation for 1 to 4 h. Beads were washed three times with 1× PBS containing 1× protease inhibitors and eluted in 50 µl 0.1 M glycine (pH 2.8). Samples were neutralized by addition of 50 µl Tris-HCl (pH 9.2), denatured by addition of 6× sample buffer followed by boiling for 5 min, and then analyzed by Western blotting.

**Statistical analysis.** Statistical analysis was performed using Prism software (Graphpad Software, La Jolla, CA). Gene expression data were repeated at least three times and analyzed by one or two-way analysis of variance (ANOVA) followed by the Bonferroni posttest. Chromatin immunoprecipitation data were repeated at least two times and analyzed by Student’s t test.

**RESULTS**

**Pbx1 is highly expressed in osteoprogenitors and decreases during osteoblast differentiation.** Pbx1 is strongly associated with skeletal patterning during embryonic development (40); however, the expression profile of Pbx1 from mesenchymal progenitors through stages of osteoblast differentiation is not clear. The commitment to the osteoblast phenotype is normally characterized by expression of specific genes (e.g., Runx2, the alkaline phosphatase gene [Alp], Bsp, and Ocn) at defined time points (Fig. 1A), and we sought to determine the expression of Pbx1 through these defined stages. We used two different progenitor cell lines representing distinct stages of commitment to the osteoblast lineage. When treated with BMP2 or dexamethasone to induce osteoblast differentiation, C3H10T1/2 cells, a model of multipotential mesenchymal progenitors, exhibited decreased Pbx1 mRNA expression (Fig. 1B) and protein levels (Fig. 1C). Analysis of lineage-committed MC3T3-E1 preosteoblastic cells revealed that Pbx1 expression was relatively high during the proliferative stage (day 4) but decreased consistently after day 15 with maturation of the osteoblast precursors (Fig. 1D). Correspondingly, Pbx1 protein levels were reduced at later stages of the differentiation time course (Fig. 1E). In contrast, mRNAs of other TALE family proteins (Pbx2, Pbx3, and Meis1) were detectable at low levels and did not decline during osteoblast differentiation (Fig. 1D). These findings suggest that unlike other TALE family members, Pbx1 is specifically downregulated during differentiation to a mature osteoblast.

**Pbx1 is expressed in the bone environment.** We sought to determine if the pattern of Pbx1 expression observed in maturing osteoblasts was indicative of protein expression in native bone. Immunohistochemical staining of mouse long bones revealed that Pbx1 was expressed in bone marrow cells and proliferating chondrocytes but could not be observed in hypertrophic chondrocytes (Fig. 2A). Pbx1 was also expressed in preosteoblasts and active osteoblasts on newly forming trabecular bone (Fig. 2A, G, and J). On the periosteal bone surface, robust Pbx1 staining could be observed in muscle cells and osteoprogenitors but not in mature osteoblasts (cells directly

**FIG. 1.** Pbx1 expression decreases during osteoblast differentiation. (A) Schematic illustrating temporal gene expression during osteoblast differentiation. Osteoblast-like differentiation was induced in C3H10T1/2 cells by treatment with either 2% FBS (control), 100 ng/ml BMP2, or 10 nM dexamethasone for 7 days. (B) Relative mRNA expression of Pbx1 in C3H10T1/2 cells was determined by RT-qPCR. (C) Protein expression was determined by Western blotting using an anti-Pbx1 antibody. MC3T-3E1 cells were treated with 280 µM ascorbic acid–5 mM β-glycerophosphate to induce osteogenic differentiation for a total period of 28 days. (D) Total RNA was isolated, and relative expression of TALE family protein genes was determined by real-time quantitative PCR using gene-specific primers. (E) Relative protein expression of Pbx1 over the 28-day time course was determined by Western blotting using an anti-Pbx1 antibody. Error bars indicate SEM.
adjacent to new bone) (Fig. 2D and J). In contrast, Hoxa10 could be observed in most cells within cartilage, bone, and muscle (Fig. 2B, E, H, and K), while Runx2 was restricted primarily to hypertrophic chondrocytes and osteoblasts and was not expressed in bone marrow or muscle (Fig. 2C, F, I, and L). These findings suggest that in bone Pbx1 is present in osteoprogenitor cells but is reduced in mature osteoblasts, supporting the notion that Pbx1 is downregulated during osteoblast maturation.

**Pbx1 functionally impairs osteoblast differentiation.** To understand the contribution of Pbx1 in commitment and maintenance of the osteoblast lineage, Pbx1 was exogenously expressed in C3H10T1/2 cells under the control of a CMV promoter using recombinant lentivirus. Increased virus titer demonstrated a dose-dependent correlation with Pbx1 protein levels (Fig. 3A), and high expression of Pbx1 was maintained over the 12-day experimental period (data not shown). C3H10T1/2 cells were infected with Pbx1 or control (green fluorescent protein [GFP]) lentivirus and treated with BMP2 to induce osteoblast differentiation. Pbx1 overexpression decreased alkaline phosphatase activity by 50% compared to controls (Fig. 3B). Examination of a panel of osteoblast markers (Alp, Runx2, Ocn, and Bsp) by RT-qPCR also showed significant ($P < 0.01$) reductions (10 to 25% of control values) in expression of these endogenous genes in response to Pbx1 (Fig. 3C). Similarly, C2C12 cells (a mesenchymal/myogenic progenitor with osteoblastic potential) examined under the
same conditions demonstrated a significant ($P < 0.01$) decrease in BMP2-induced osteoblastogenesis in response to Pbx1 (see Fig. S1 at http://labs.umassmed.edu/steinlab/). Our studies with the committed preosteoblast MC3T3-E1 cells also echoed a reduction in osteoblast-specific differentiation by Pbx1 expression from the onset of alkaline phosphatase activity (day 6) through the matrix maturation stage (day 12) (Fig. 3D). Due to the committed osteogenic phenotype of MC3T3-E1 cells, Pbx1 had no effect on expression of the early osteogenic marker Runx2 (day 12), while mature osteoblast genes Bsp and Ocn were significantly ($P < 0.01$) decreased, by 50% (Fig. 3E). Taken together, these findings indicate that Pbx1 is a negative regulator of mesenchymal progenitor commitment to the osteoblast phenotype as well as the maturation of osteoprogenitors.

**Pbx1 negatively regulates osteogenic genes: depletion of Pbx1 by shRNA increases osteoblast differentiation.** The reduction of osteoblast markers by Pbx1 overexpression in differentiated cells suggested that Pbx1 may have a biological role in attenuating osteoblast-specific gene expression. To address this question, several Pbx1 shRNAs were generated and exogenously expressed in C3H10T1/2 cells using recombinant lentivirus. All three Pbx1 shRNA sequences (T1, T2, and T3) successfully decreased Pbx1 protein levels (Fig. 4A). This decrease in Pbx1 levels was maintained for the period of the experiments (>20 days) (data not shown), and the shRNA construct generating the most efficient knockdown (T2) was used for subsequent studies. C3H10T1/2 cells with reduced Pbx1 expression exhibited increased (3-fold) alkaline phosphatase activity and expression compared to cells infected with nonsilencing (NS) control (Fig. 4B). Additionally in Pbx1-shRNA-infected cells, Ocn and Bsp mRNA levels, markers of the mature osteoblast, were upregulated 5- to 6-fold compared to NS control cells (Fig. 4C), while Runx2 mRNA levels did not change significantly. Pbx1-shRNA resulted in a similar enhancement of osteoblast differentiation in MC3T3-E1 cells, with significant ($P < 0.01$) increases in Alp activity and Alp, Ocn, and Bsp mRNA levels compared to those in NS-infected controls but no change in Runx2 mRNA (Fig. 4D and E).

These findings are consistent with the Pbx1 expression profile in osteoblasts (Fig. 1 and 2) and forced-expression studies (Fig. 3) and establish that Pbx1 is a potent negative regulator of osteoblastic genes expressed during early and late stages of osteoblast differentiation.

A Pbx1 regulatory element is required to inhibit Hoxa10-mediated activation of osteocalcin and bone sialoprotein promoters. The potent effects on osteoblast gene expression mediated by alteration of Pbx1 expression levels suggested that Pbx1 may directly regulate bone-related gene promoter activity. Pbx1 has been demonstrated to bind directly to DNA and modulate the binding of Hox proteins for several promoters (22). We sought to determine if the negative regulation of the osteoblast phenotype by Pbx1 was dependent on the putative Pbx regulatory elements (33) located in the proximal promoters of several bone-related genes, including Bsp and Ocn. To address this question, Pbx1 and Hoxa10 expression plasmids were cotransfected with luciferase reporter constructs containing approximately 1 kbp of the Ocn ($\sim 1.1$ kbp Ocn-Luc) (Fig. 5A) or Bsp ($\sim 988$ bp Bsp-Luc) promoter region (see Fig. S2 at http://labs.umassmed.edu/steinlab/). MC3T3-E1 preosteoblasts transfected with Hoxa10 in addition to the Ocn or Bsp luciferase reporter exhibited significantly ($P < 0.01$) increased luciferase activity (5- to 6-fold) (Fig. 5A and data not shown). However, cotransfection of cells with Hoxa10 and Pbx1 along with either reporter construct (Ocn or Bsp) resulted in a 75% decrease in luciferase activity compared to that in cells transfected with Hoxa10 alone (Fig. 5A; also see Fig. S2 at http://labs.umassmed.edu/steinlab/). There was no change in luciferase activity in cells transfected with Pbx1 alone. These results indicate that Pbx1 acts to reduce Hoxa10-mediated transcriptional activation at bone-specific promoters.

We next identified that the Pbx1-binding site is functionally active by determining that in vitro-translated (IVTT) recombinant Pbx1 and MC3T3-E1 nuclear protein extracts could bind to the radioactively labeled probe (Ocn promoter sequence GCCAGCTCTGGTGTCC) by electrophoretic mobility shift assay (see Fig. S3 at http://labs.umassmed.edu/steinlab/).
Ocn

porter construct (pGL3-Basic) containing the erase activity was assessed with a luminometer. (A) Luciferase re-

pCDNA3, pCDNA-Hoxa10, and/or pCDNA-Pbx1, and relative lucif-

luciferase reporter plasmids as well as the indicated amounts (ng) of MC3T3-E1 preosteoblasts were transfected with 500 ng of the indicated promoter results in decreased attenuation of Hoxa10 activity.

sioned to loss of Pbx1 repression of Hoxa10-mediated Ocn/H11002

1.1 kb mutHox-Ocn-Luc) or the Hox DNA consensus sequence (1.1 kb mutPbx-Ocn-Luc). Hoxa10-mediated luciferase reporter activity of the mutHox-Ocn promoter (Fig. 5C) could be due to the contribution from increased Runx2-mediated transcriptional activity, as Hoxa10 can directly upregulate Runx2 expression (19). However, all Hox-mediated activation is significantly (P < 0.05) decreased by cotransfection of Pbx1 (Fig. 5C). Taken together, these results provide direct evidence that Pbx1-DNA interactions are required to block Hoxa10-mediated transcriptional activation of osteoblast-related genes.

Pbx1 is associated with repression of chromatin remodeling on the osteocalcin promoter. Our previous findings identified Hoxa10 as an early activator of bone gene expression and demonstrated that Hoxa10 functions by facilitating the remodeling of chromatin (18). Given the role of Pbx1 as a mediator of Hox function, we sought to establish if Pbx1 is associated with chromatin remodeling of the temporally expressed osteocalcin gene during osteoblast differentiation. The Ocn gene promoter (Fig. 6A) and the Bsp promoter (see Fig. S4A and B at http://labs.umassmed.edu/steinlab/) are strongly regulated by chromatin structure and are tightly packaged as heterochromatin in proliferating cells (day 4) (32). During transcriptional activation, the proximal promoter must adopt an open conformation to allow binding of transcriptional regulators, resulting in an exponential increase in gene expression (day 12) (Fig. 6B). By chromatin immunoprecipitation analysis, Pbx1 is strongly associated with the Ocn and Bsp promoters prior to expression (day 4) but not during exponential gene expression (day 12) (Fig. 6C; also see Fig. S4C at http://labs.umassmed.edu/steinlab/). This result was in contrast to the pattern observed with RNA polymerase II and known transcriptional activators (e.g., Runx2 and Hoxa10), all of which demonstrated increased Ocn promoter occupancy with increased Ocn gene expression (Fig. 6C).

Several mediators and covalent modifications of histone proteins associated with chromatin remodeling, including CBP/p300, acetylated histone H4, and acetylated H4 lysine 16 (H4K16), were found to increase with osteoblast differentiation (Fig. 6D; also see Fig. S4D at http://labs.umassmed.edu/steinlab/), although acetylation of H3 lysine 9 (H3K9) was unchanged. In contrast repressive marks on chromatin, including histone deacetylases (HDAC) 1, 2, and 3, as well as methylation of histone H3K9, demonstrated a decrease upon osteoblast
differentiation (Fig. 6E; also see Fig. S4E at http://labs.umassmed.edu/steinlab/). These results strongly suggest that Pbx1 is associated with negative regulation of chromatin remodeling at the osteocalcin and bone sialoprotein promoters.

To address whether there is a direct association of Hoxa10 and Pbx1 with regulators of chromatin remodeling at the osteocalcin promoter in differentiated osteoblasts, two-step chromatin immunoprecipitation (ChIP-reChIP) experiments were performed using a primary pulldown with an anti-Hoxa10 or anti-Pbx1 antibody. The chromatin fraction associated with Hoxa10 at the $Ocn$ promoter demonstrated a high level of association with Pbx1, as the secondary pulldown with anti-Pbx1 antibody resulted in a signal that was approximately 60% of the input (Hoxa10) (Fig. 7A). Consistent with the primary ChIP analysis (Fig. 6C), the association of Pbx1 with the Hoxa10 chromatin fraction was greatly reduced at day 12. In comparison, Runx2 was weakly associated with the Hoxa10 chromatin fraction at day 4 and subsequently increased at day 12. This increased binding of Runx2 in association with Hoxa10 on the $Ocn$ promoter at day 12 reflects upregulation of the $Ocn$ gene, which is also demonstrated by the 4- to 5-fold-increased association of the transcriptional coactivator p300.

In contrast to Hoxa10, the Pbx1- $Ocn$ chromatin fraction was only weakly associated with Runx2, CBP, or p300 (by secondary pulldown) at either day 4 or day 12 (Fig. 7B), reflecting the role of Pbx1 as a repressor of $Ocn$ gene expression. However, the Pbx1 chromatin fraction was strongly associated with HDAC1 at the osteocalcin promoter when the gene was silent during proliferation (day 4) and during bone-specific gene expression (day 12), and neither HDAC1 nor Pbx1 was strongly associated with Hoxa10 at day 12. Due to the cooccupancy of Pbx1 with negative regulators of chromatin remodeling at the osteocalcin promoter during proliferation and differentiation, Pbx1 displays a pattern of gene-repressive functionality on the osteocalcin promoter. (A) Diagram of rat osteocalcin promoter displaying relative binding sites and primer sites used for chromatin immunoprecipitation analysis. Rat calvarial osteoblasts were isolated from embryonic day 18.5 rat pups and collected during the proliferative stage (day 4) or cultured in differentiating conditions and collected during exponential increase in osteocalcin expression (day 12). (B) Calvarial osteoblasts were analyzed at the indicated differentiation stages by RT-qPCR to determine levels of osteocalcin gene expression. (C to E) The ChIP analysis was performed on cleared lysates from primary osteoblasts using $\sim 5 \mu g$ of the indicated antibody. Recovered DNA was then quantified by qPCR using primers specific for the proximal promoter region of the osteocalcin gene to determine relative occupancy of transcriptional activators (C), activation markers (D), or repressive markers (E). Statistical significance was determined by Student's t test ($\ast$, $P < 0.05$ versus matched control). ChIP experiments were repeated at least two times with similar results, and one representative experiment is presented ($\pm$ standard deviation [SD]).
we sought to determine if association of Pbx1 with these factors (including Hoxa10) was by direct protein-protein interactions. Chromatin digested with PstI restriction endonuclease to separate the Pbx and Hox binding sites was immunoprecipitated with Hoxa10, Pbx1, or control (IgG) antibodies (Fig. 7C). DNA containing the Hoxa10-binding site could be detected in both Hoxa10 and Pbx1 pulldowns by qPCR. Similar results were observed for the Pbx1-binding site fragment, suggesting that Hoxa10 and Pbx1 may interact through direct protein-protein interactions on the \textit{Ocn} promoter. To detect other protein-protein interactions, whole-cell lysates from confluent MC3T3-E1 cells were immunoprecipitated with Hoxa10, Runx2, Pbx1, HDAC1, or control (IgG) antibodies. Proteins were resolved by SDS-PAGE, and Hoxa10 and Pbx1 were detected by Western blotting with specific antibodies. (A) Whole-cell lysates from confluent MC3T3-E1 cells were immunoprecipitated with Hoxa10, Runx2, Pbx1, HDAC1, or control (IgG) antibodies and quantified by qPCR. (B) Lysates from MC3T3-E1 were used for immunoprecipitation using anti-HDAC1 followed by Western blotting with anti-Pbx1 or specific HDAC antibody.
was decreased compared to that in NS-infected cells, as would be predicted based on reduced cellular levels of Pbx1 (Fig. 9D). While there was no effect on Hoxa10 binding, a large increase in both Runx2 and p300 occupancy at the osteocalcin promoter in cells deficient in Pbx1 as well as a reduction in occupancy of HDAC1 and HDAC2 was found, consistent with findings from ChIP-reChIP studies (Fig. 7). These results indicate that Pbx1 may be important for recruiting these HDAC factors to the osteocalcin promoter to mark the gene for expression while maintaining a repressive state during proliferation. The data suggest that Pbx1 contributes to the switching of chromatin remodeling factors from repression to open chromatin for activation, as illustrated in Fig. 9E.

DISCUSSION

Here we have identified an integral role for the Hox-interacting protein Pbx1 as an important regulator of the osteoblast phenotype. Pbx1 acts to coordinate the temporal expression of the Ocn and Bsp genes by Hoxa10 during osteogenesis. It is apparent that Hox and TALE genes have defined roles in embryonic development; however, our results indicate that these proteins can have analogous or concordant roles in the maintenance of adult tissues. It has been suggested that Pbx1, as well as other TALE family proteins, can act in concert with (1, 35) or independent of (4, 8) Hox proteins in its role of regulating gene expression through direct binding to cognate regulatory elements. We have provided several lines of evidence that Pbx1 is a critical negative regulator of osteoblast differentiation by demonstrating that (i) Pbx1 is highly expressed in osteoprogenitors and its expression at the transcript and protein levels decreases upon commitment to the osteoblast phenotype and again during late stages of differentiation; (ii) exogenous expression of Pbx1 or depletion by targeted-shRNA expression inhibits or increases, respectively, osteoblastogenesis and the expression of osteoblast-related genes; (iii) Pbx1 acts directly on the Ocn and Bsp promoters to repress Hoxa10-mediated expression of the gene by a mechanism in which Pbx1 recruits negative regulators of chromatin remodeling; and (iv) Pbx1 has the ability to act as a molecular switch functioning with Hoxa10 in proliferating cells to maintain osteoblast-specific genes in a repressed state and facilitate Hoxa10 activation of gene promoters by dissociating from the chromatin in differentiated cells. These key findings highlight a novel regulatory role for Pbx1 to repress and mark osteoblast genes for expression and limit Hoxa10-mediated gene transcription for regulating the temporal activation of postproliferative gene expression.

Our data suggest that Pbx1 has a role in attenuating osteogenesis by repressing osteoblast gene expression at several stages. Increased Pbx1 expression in mesenchymal precursors decreased expression of early osteoblast genes (Runx2 and Alp) and late-stage genes (Bsp and Ocn), whereas decreased Pbx1

|FIG. 9. Modification of Pbx1 levels in osteoblasts results in alteration of histone-modifying enzymes at the osteocalcin promoter. (A and B) Rat calvarial osteoblasts were isolated from embryonic day 18.5 rat pups and infected with Pbx1 or empty (EV) lentiviral constructs, and cells were collected at 48 h after infection after just reaching confluence (day 6). (A) Relative expression of the Pbx1 and Ocn genes was monitored by RT-qPCR. (B) ChIP analysis was performed on cleared lysates from primary osteoblasts using ~5 μg of the indicated antibody. Recovered DNA was then quantified by qPCR using primers specific for the proximal promoter region of the osteocalcin gene to determine the relative occupancy of the indicated proteins. (C) Isolated osteoblasts (as described above) were infected with Pbx1-shRNA or nonsilencing shRNA (NS) lentiviral constructs, and relative expression of Pbx1 and Ocn was determined by RT-qPCR. (D) ChIP analysis was performed on cleared lysates from treated primary osteoblasts (as described above) to determine the relative occupancy of the indicated proteins on the osteocalcin promoter. Statistical significance was determined by Student’s t test (*, P < 0.05 versus matched control). ChIP experiments were repeated two times with similar results, and the data presented are representative of one experiment (± SD). (E) Schematic model of Pbx-mediated repression of the osteocalcin gene.
levels resulted in increased osteoblast gene expression. The observation of enhanced commitment to the osteoblast phenotype in the absence of Pbx1 is concordant with reports of premature mineralization of skeletal elements, particularly at the hypertrophic chondrocyte zone, in growth plates in Pbx1-null mice (40). In contrast to Pbx1-null mice, which die at embryonic day 15, mice with knockouts of other TALE family genes (Pbx2 and Pbx3) survive after birth and show only slight skeletal defects, with no apparent effects on osteoblast differentiation (7, 41). Meis1-null mice die due to defects in fetal hematopoiesis and do not develop the skeletal abnormalities observed in Pbx1<sup>−/−</sup> mice (20), together suggesting that Pbx1 may have a specific role in the commitment of precursors to the osteoblast lineage. It is interesting to note that conditional Pbx1 deletion in hematopoietic stem cells results in loss of self-renewal and the premature initiation of transcriptional mechanisms, resulting in maturation of progenitor cells to a differentiated state (14). Likewise, Pbx1 depletion from the osteoprogenitor stage to the osteoblast appears to have a temporal role in promoting differentiation. Thus, Pbx function is to limit differentiation by maintaining a mesenchymal phenotype, similar to the role of the protein in maintaining the embryonic mesoderm (7).

Our previous studies demonstrated that Hoxa10 functions as an immediate-early response to the osteogenic BMP2 signal and is capable of inducing osteogenic gene expression. Therefore, Hoxa10 is an important cofactor to establish the osteoblast phenotype (19). In this study, our findings show that Pbx1 occupies gene promoter coincident with Hoxa10, resulting in tempered Hoxa10-mediated gene expression. In committed osteoprogenitors where Pbx1 expression is robust, it is clear that Pbx1 and Hoxa10 both bind to the osteocalcin promoter preceding gene expression. These findings are consistent with our earlier studies that showed a significant increase in Hoxa10 occupancy of bone promoters that are expressed in mature osteoblasts (19). Here we show that Hoxa10 is associated with chromatin of the actively transcribed genes (day 12).

The regulation of osteoblast genes by Pbx1 and Hoxa10 appears to be facilitated by the conserved organization of regulatory elements in the promoters of osteoblast-related genes. The proximal promoter regions of several genes generally associated with bone formation feature multiple homeodomain-binding DNA motifs, including some that preferentially bind Pbx1 (tTGAC/T) (33). Strikingly in the Bsp and Ocn promoters, the Pbx1 consensus sequence and the Hoxa10-binding site flank functional Runx2-binding elements (16, 37) (Fig. 6A; also see Fig. S4A at http://labs.umassmed.edu/steinlab/). The regulation of the Ocn promoter by Hoxa10 involves the accumulation of Hoxa10 on the Ocn promoter, peaking with active gene transcription. This finding establishes a key event for Hoxa10-mediated gene activation by dissociation of Pbx1 from the Ocn and Bsp genes when they are actively transcribed. Coincident with the dissociation of Pbx1, there is an increase in Runx2 and RNA Pol II, which marks the onset of active gene transcription. Thus, we conclude from our studies of primary osteoblasts progressing through stages of differentiation that dissociation of Pbx1 may be an important step (or switching mechanism) in allowing RNA Pol II and/or Runx2 recruitment to these promoters, resulting in gene transcription.

Chromatin remodeling through the recruitment of histone-modifying enzymes that regulate the methylation or acetylation of lysine and arginine residues is a fundamental underlying mechanism regulating transcription (29, 30). Hoxa10 as well as Pbx1 has a defined role in the patterning of the axial skeleton (6, 43, 44) by regulating chromatin remodeling that leads to phenotypic gene expression in several tissues (4, 28). Pbx1 has been demonstrated to be involved in chromatin remodeling events leading to the activation of MyoD-dependent promoters (such as myogenin), and the protein is constitutively bound to silent chromatin prior to initiation of muscle differentiation (4, 11). Supporting the concept that Pbx1 exerts functional activity through epigenetic control, the pattern of histone modifications and histone-modifying enzymes at the osteocalcin promoter indicates that Pbx1 is associated with a repressed chromatin state and that the chromatin adopts an “active” conformation marked by higher levels of H3K4 methylation and acetylation upon Pbx1 dissociation. Several studies have reported that HOX proteins (including Hoxa10) can mediate a modified chromatin acetylation status leading to transcriptional activation or repression by interacting with p300/CPB (3, 9, 42). The ChIP-reChIP experiments presented in this study demonstrate that Hoxa10 is strongly associated with CBP at the osteocalcin promoter during both proliferation and differentiation and with p300 during differentiation, while Pbx1 is localized on the osteocalcin promoter coincident with histone deacetylases and Hoxa10 in proliferating cells. This result would further imply that Pbx1 is a repressor of chromatin remodeling and gene transcription in osteoblasts. Supporting this model, it has been previously demonstrated that Hox-Pbx complex functions can promote histone deacetylation by recruiting class I histone deacetylases and Hoxa10 in proliferating cells. This result would further imply that Pbx1 is a repressor of chromatin remodeling and gene transcription in osteoblasts. Supporting this model, it has been previously demonstrated that Hox-Pbx complexes can promote histone deacetylation by recruiting class I histone deacetylases to repress gene transcription (39). It is therefore intriguing that when colocated at the osteocalcin promoter, Hoxa10 and Pbx1 have the characteristics of both repressional and activational regulators, containing both HDACs and histone acetyltransferases (CBP/p300). Our findings suggest that even though Pbx1 is a strong inhibitor of osteoblastogenesis, it may have a temporal role in assisting in the activation of chromatin remodeling and support a switching mechanism from repression to activation. In conclusion, in this study we have demonstrated functional interactions between Pbx1 and Hoxa10 in progenitor cells and osteoblasts for control of the temporal expression of bone-specific genes at multiple levels, including transcriptional regulation at the promoter level and initiation of transcriptional activity through chromatin remodeling. This characterization of the Pbx1 and Hoxa10 interactions on the osteocalcin promoter has provided a clearer understanding of the commitment to an osteogenic lineage preceding Runx2 recruitment in mesenchymal progenitors. This concept has implications for bone repair and molecular intervention to correct bone disorders.

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