The paired-like homeo box gene MHox is required for early events of skeletogenesis in multiple lineages

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Formation of cartilage and bone involves sequential processes in which undifferentiated mesenchyme aggregates into primordial condensations that subsequently grow and differentiate, eventually forming the adult skeleton. Although much has been learned about the structural molecules that compose cartilage and bone, little is known about the nuclear factors that regulate chondrogenesis and osteogenesis. MHox is a homeo box-containing gene that is expressed in the mesenchyme of facial, limb, and vertebral skeletal precursors during mouse embryogenesis. MHox expression has been shown to require epithelial-derived signals, suggesting that MHox may regulate the epithelial–mesenchymal interactions required for skeletal organogenesis. To determine the functions of MHox, we generated a loss-of-function mutation in the MHox gene. Mice homozygous for a mutant MHox allele die soon after birth and exhibit defects of skeletogenesis, involving the loss or malformation of craniofacial, limb, and vertebral skeletal structures. The affected skeletal elements are derived from the cranial neural crest, as well as somitic and lateral mesoderm. Analysis of the mutant phenotype during ontogeny demonstrated a defect in the formation and growth of chondrogenic and osteogenic precursors. These findings provide evidence that MHox regulates the formation of preskeletal condensations from undifferentiated mesenchyme.

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tions, little is known about the nuclear factors that regulate these events. Homeo box-containing genes such as the msh-like genes msx-1 and msx-2 have been implicated as regulators of inductive events in the vertebrate limb and cranium (Davidson et al. 1991; Robert et al. 1991; Takahashi et al. 1991; Satokata and Maas 1994). In addition to the msx genes, the homeo box-containing gene MHox is expressed in the limb and facial primordia during embryogenesis (Cserjesi et al. 1992; Kuratani et al. 1994).

MHox was originally identified as a nuclear factor that bound an essential element in the muscle creatine kinase enhancer (Cserjesi et al. 1992). It was also isolated independently using a polymerase chain reaction (PCR)-based strategy (Kern et al. 1992). The human homolog of MHox, called Phox, was cloned in a yeast screen for proteins that interact with serum response factor (SRF) to activate transcription through the serum response element (Grueneberg et al. 1992). Within the homeo domain, MHox is most closely related to the paired family of homeo box-containing genes. However, MHox lacks a Paired domain and contains a glutamine in place of serine at position 9 of the recognition helix that determines DNA-binding specificity of the homeo domain (Desplan et al. 1998). During mouse and chick embryogenesis, MHox is expressed in the lateral mesoderm and in the mesenchyme of the limb buds and facial primordia, but its expression is excluded from overlying epithelial tissues. Removal of surface epithelium inhibits expression of MHox in the underlying mesenchyme, suggesting that MHox may respond to or mediate inductive signals emanating from the epithelium (Kuratani et al. 1994).

To investigate the role of MHox in mouse embryogenesis, we generated mice carrying a germ-line mutation in the MHox locus. The MHox mutant mice died neonatally and displayed multiple craniofacial defects resulting from the deletion or malformation of specific skeletal elements derived from the neural crest of the first branchial arch and cephalic mesoderm. In addition, MHox mutant mice had a limb defect in which the mutant zeugopod was bowed and foreshortened. However, proximal and distal limb structures were spared. A small percentage of MHox mutants also exhibited abnormalities of the thoracic and lumbar spine, in which the neural arches were flared and the cartilaginous precursors of the dorsal spinous process were truncated, resulting in spina bifida. Analysis during ontogeny revealed that the skeletogenic abnormalities identified in the MHox mutant mice arose as a consequence of a defect in growth of chondrogenic and osteogenic precursors. These findings reveal that MHox plays a central regulatory role in the tissue interactions that give rise to craniofacial and limb skeletal structures and, to a lesser extent, the axial skeleton.

Results

Generation of MHox mutant mice

To introduce a mutation in the MHox gene, we constructed a targeting vector containing 6.5 kb of genomic DNA encompassing the region of the MHox gene that encodes the homeo domain, and the neomycin resistance cassette was inserted into the 5' region of the homeo box (Fig. 1A, B). The linearized targeting vector was electroporated into AB-1 embryonic stem (ES) cells which were then subjected to positive-negative selection using G-418 and FIAU (see Materials and methods).

Insertion of neo into the MHox locus introduced novel EcoRI, Ncol, and ClaI sites that could be used to distinguish the targeted from the wild-type MHox allele by Southern analysis. From a total of 590 ES cell clones, 38 specific targeted events were identified by Southern analysis for a targeting frequency of ~1:15. Two independent ES clones containing an MHox mutant allele were injected into 3.5-day blastocysts derived from C57Bl/6 mice, and chimeras were obtained. Transmission of the mutation through the germ line yielded offspring heterozygous for the mutant MHox allele. Subsequent heterozygous intercrosses resulted in the generation of mice homozygous for the mutant allele. Southern blot analysis of tail DNA from mice of the three MHox genotypes (+/+, +/−, and −/−) is shown in Figure 1C. Genotyping of 253 neonatal and embryonic mice revealed that the neonatal MHox mutant mice were produced at a Mendelian frequency, demonstrating that the MHox mutant mice survived throughout gestation.

To confirm that the targeted gene expressed no functional MHox transcripts, MHox mRNA from wild-type (+/+) and mutant (−/−) animals was analyzed by reverse transcriptase PCR (RT-PCR) using primers surrounding the region of the MHox mRNA where neo was inserted. The MHox transcript from the wild-type mouse and the 10T1/2 cell line, which expresses high levels of MHox mRNA, gave rise to an amplified product of 350 bp (Fig. 1D). However, no PCR product was detected in the MHox mutant mouse, demonstrating that the MHox mRNA was disrupted by the mutation. Transcripts for the ubiquitously expressed ribosomal protein L7 were analyzed as an internal control to confirm the integrity of the RNA.

The MHox (−/−) mutation results in neonatal lethality and craniofacial abnormalities

MHox (−/−) mice were readily identifiable at birth because of cranial abnormalities, including microcephaly, low-set ears, and a pointed snout (Fig. 1E–H). The mutant neonates also demonstrated respiratory compromise as evidenced by gasping motions and cyanosis. Mutant mice never fed, and they developed abdominal distention due to inflation of the digestive tract with air.

Gross examination of MHox mutant neonates revealed that they had a large cleft of the secondary palate (Fig. 2A, B), which may have contributed to their difficulty in feeding and breathing. Coronal sections through the head clearly showed the unfused palatal shelves in the mutants (Fig. 2C and D).

Numerous cranial skeletal elements are altered or absent in MHox–mutant neonates

To identify the specific anatomic defects underlying the craniofacial abnormalities in MHox (−/−) mice, skele-
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Figure 1. Targeting of the mouse MHox locus. (A) Restriction map of the region of the MHox gene encoding the homeo domain (indicated in black). The restriction sites are EcoRI (R), StuI (S), BglII (B), XbaI (X), NcoI (N), and ClaI (C). The amino acid sequence of the homeo domain is shown. The position of the intron is indicated by the vertical arrow below the sequence, the insertion site of the neo cassette is indicated by the vertical arrow above the sequence. The positions of two alternative 3' exons are indicated downstream of the homeo domain (Cserjesi et al. 1992; Kern et al. 1992). Both potential AUG codons are also shown. The positions of PCR primers used for RT–PCR are indicated by horizontal arrows. (B) Strategy for targeting the MHox locus. The region of the MHox gene that was targeted is shown at the top. The targeting vector (middle), contained neo within the homeo domain at the position indicated in A, the predicted structure of the targeted MHox allele is shown at the bottom. The positions of the two probes used for Southern analyses are indicated at the top. (C) Southern blots of genomic DNA from mice of the three MHox genotypes (+/+, +/-, -/-) are shown. Probe A hybridizes to bands of 6.5 and 4.2 kb in genomic DNA from wild-type and MHox mutant mice, respectively, digested with EcoRI. Probe B hybridizes to bands of 9.1 and 5.7 kb in genomic DNA from wild-type and MHox mutant mice, respectively, digested with NcoI plus ClaI. (D) Transcripts for MHox and L7 were detected by RT–PCR as described in Materials and methods. RNA from 10T1/2 cells was included as a positive control. No RT–PCR products were detected when reverse transcriptase was omitted from the reaction (data not shown). (E,F) Lateral views of wild-type (E) and MHox(-/-) (F) neonates. The arrow in F denotes the low-set ear of the mutant. This mutant is also beginning to show abdominal distention and lacks milk in its stomach, which can be seen in the wild-type littermate. (G,H) Frontal views of wild-type (G) and MHox(-/-) (H) neonates. Note the microcephaly and pointed snout of the mutant.

tons of neonates were stained for cartilage and bone. Several major defects were readily apparent in the skulls of the mutant mice. The squamosal bone and the ascending lamina of the alisphenoid bone were absent, resulting in a large defect on the lateral aspect of the skull [Fig. 3, cf. A,C,E, with B,D,F]. There was also no trace of the zygomatic bone, gonial bone, tympanic ring [Fig. 3B,D,F], or supraoccipital bone [Fig. 4B]. At the base of the skull, the pterygoid bone was hypoplastic and the palatal processes of the palatal and maxillary bones were missing [Fig. 2E,F, data not shown]. Several defects were also observed in mandibles from MHox mutant mice. The coronoid, condylar, and angular processes of the mandible were hypoplastic, and the dentary component was shortened in all mutant animals [Fig. 3G,H]. The alveolar processes and tooth buds, however, were unaffected. These affected skeletal structures of the MHox mutant mice are derived primarily from neural crest of the first branchial
Figure 2. *MHox* mutant mice are born with bilateral cleft of the secondary palate. [A,B] Ventral view of the upper jaw of wild-type and *MHox*(−/−) neonates, respectively. The arrows point to the lateral edges of the secondary palate. Note there is no outgrowth of the palatal shelves in the mutant, leaving an open view of the nasal cavities, whereas the palatal shelves have fused at the midline in the wild type. [C,D] Coronal sections of the heads of wild-type and *MHox*(−/−) neonates, respectively, showing the palatal shelves. The palatal shelves in the mutant failed to extend toward the midline. Magnification, 108×. [E,F] Ventral view of the base of skulls of wild-type and *MHox*(−/−) neonates, respectively. Skulls were stained for bone (purple) and cartilage (blue). The palatal and pterygoid bones are hypoplastic in the mutant. (bo) Basioccipital bone; (bs) basisphenoid bone; (ns) nasal septum; (pl) palatal bone; (ps) palatal shelves; (pt) pterygoid bone; (t) tongue.

arch, except for the alisphenoid and supraoccipital bones, which are derived from cephalic mesoderm [Noden 1988; Le Douarin et al. 1993]. Both membranous bones, such as the squamosal, and enchondral bones, such as the supraoccipital, were affected by the mutation.

In addition to the cranial and jaw abnormalities, the *MHox* mutation resulted in several defects in the middle ear ossicles, which arise from first- and second-arch neural crest [Le Douarin et al. 1993]. In modern mammals the malleus, incus, and stapes form an ossicular chain that functions to transmit sound. The head of the malleus, which is derived from the proximal end of Meckel's cartilage [Novacek 1993], was the correct size in the mutant neonates. However, the manubrium of the malleus, which normally attaches to the tympanic membrane, was truncated and the processus brevis was displaced (Fig. 5, cf. A and B). Rostral to the mallear head, an additional cartilaginous process was identified in every mutant animal examined (Fig. 5B). During development, the incus, which is derived from the proximal end of the palatoquadrate cartilage [Novacek 1993], forms a separate cartilaginous body that articulates with both the stapes and the head of the malleus. In *MHox* mutants, the incudo-mallear joint was intact but the incus remained fused to a persistent quadrate cartilage on the lateral aspect of the skull and so did not articulate with the stapes (Fig. 5, cf. C and D).

The third middle ear ossicle, the stapes, is derived from the rostral portion of Reichert's cartilage in the second pharyngeal arch [Moore and Persaud 1993]. In mutant mice, the stapes was columniform and remained fused to Reichert's cartilage. In addition, the anterior and posterior crus were fused, obliterating the foramen through which the stapedial artery normally passes (Fig. 5, cf. E and F).

All of the above defects were fully penetrant. The head structures that were affected by the *MHox*-null mutation were highly specific, as many cranial bones were unaffected. Table 1 summarizes the cranial defects in *MHox* mutant mice, which were restricted primarily to skeletal elements derived from cranial neural crest within the first branchial arch and, to a lesser extent, to structures that arise from the second arch crest and the cephalic mesoderm. Transverse and coronal sections through the head of *MHox* mutant neonates revealed that the cranial musculature was intact [not shown]. Because cranial neural crest cells give rise to both neurogenic and skeletogenic components of the head and face [Noden 1983; 1988; Le Douarin et al. 1993], we analyzed the trigeminal nerve, which innervates the first branchial arch and its derivatives. Histologic analysis showed that the trigeminal [Gasserian] ganglion was normal in mutant animals [not shown]. Immunostaining of embryos 10.5 days post-coitum [dpc] with an anti-neurofilament antibody also showed that cranial nerves VII, IX, and X were unaffected by the *MHox* mutation [not shown].

Developmental time course of the cranial mutant phenotype

To begin to define which developmental processes were affected by the *MHox* mutation, we examined the forming cartilaginous and bony skeletons of mouse embryos.
Cartilage staining of mutant and wild-type embryos demonstrated that the \(\text{MHox}\) mutant phenotype was apparent at 13.5 dpc. The defects at this stage included the absence of the tectum posteriorus, the cartilaginous precursor of the supraoccipital bone [Fig. 6A,B]. Meckel’s cartilage remained unfused and had a sigmoidal morphology in mutant embryos, compared with the straight rod of cartilage seen in wild-type embryos. In addition, the ala temporalis, a cartilaginous structure that gives rise to the alisphenoid bone, was detectable in wild-type embryos at this stage but was not present in mutant embryos [Fig. 6C,D]. One day later, the mutant Meckel’s cartilage fused, although it remained morphologically abnormal [Fig. 6E–H]. The ala temporalis, though present, remained hypoplastic relative to wild type [not shown]. The forming occipital arch and parietal cartilage remained deficient in the \(\text{MHox}\) mutant mice at 14.5 dpc [Fig. 6E,F]. Fusion of the incus to the persistent quadrate cartilage on the lateral aspect of the skull was also obvious at this stage. The abnormal formation of cartilage in the developing skull of the \(\text{MHox}\) mutant results in a trabeculated appearance characteristic of phylogenetically more primitive animals [Fig. 6E,F; Novacek 1993]. Thus, many of the morphologic abnormalities observed in the mutant neonates were the end result of the failure of cartilaginous precursors to form or grow at normal rates. Similarly, ossification centers for the membranous squamosal and tympanic bones, which were deleted in mutant neonates, were never detected in 14.5 and 16.5 dpc mutant embryos [not shown].

**Figure 3.** Lateral views of the skulls of wild-type and \(\text{MHox}\) mutant neonates. Lateral views of the skulls and upper vertebral region of wild-type \(\{A,C,E\}\) and \(\text{MHox}^{-/-}\) \(\{B,D,F\}\) neonates stained for bone [red] and cartilage [blue]. \(\{C,D\}\) A diagram of the skull bones in wild-type and mutant neonates, respectively; \(\{E,F\}\) an enlargement of the region indicated by the boxes in \(\{C,D\}\). Skeletal structures that are altered in the mutant are indicated in pink in \(\{C,D\}\). \(\{G,H\}\) The isolated mandibles from wild-type and mutant neonates, respectively. In \(\{F\}\), note that only remnants of the squamosal and alisphenoid bones appear in the mutant, resulting in the absence of a temporomandibular joint. The coronoid, condylar, and angular processes of the mandible are also hypoplastic, and the tympanic, gonial, and zygomatic bones are absent. \(\{A\}\), Angular process; \(\{At\}\) atlas; \(\{AS\}\) alisphenoid bone; \(\{Ax\}\) axis; \(\{C\}\) coronoid process; \(\{CO\}\) condylar process; \(\{D\}\) dentary process; \(\{E\}\) exoccipital bone; \(\{F\}\) frontal bone; \(\{H\}\) hyoid bone; \(\{I\}\) interparietal bone; \(\{M\}\) malleus; \(\{Mn\}\) mandible; \(\{Mx\}\) maxillary bone; \(\{N\}\) nasal bone; \(\{OC\}\) otic capsule; \(\{P\}\) parietal bone; \(\{SQ\}\) squamosal bone; \(\{S\}\) supraoccipital bone; \(\{T\}\) tympanic bone; \(\{Z\}\) zygomatic bone.
Limb abnormalities in MHox mutant embryos and neonates

During embryogenesis, MHox is expressed in the lateral mesoderm prior to limb formation and later in the mesenchyme of the forming limb bud (Cserjesi et al. 1992; Kuratani et al. 1994). In MHox mutant neonates, the ossified shafts of the radius and ulna of the forelimb and the tibia and fibula of the hindlimb were abnormally shortened and broader than wild-type bones [Table 2]. In addition, the radius and tibia of mutant animals had a bowed shape when compared to wild-type structures [Fig. 7]. No defects were detected in the epiphyseal cartilage of the long bones of neonatal animals by whole-mount cartilage staining or by histologic analysis [Fig. 7; data not shown]. The proximal and distal limb structures were unaffected [Table 2].

The long bones of the extremities form by endochondral ossification in which a cartilaginous template is replaced by bone [Ham 1987]. Analysis of the cartilaginous precursors of the affected long bones at 13.5 dpc demonstrated that they were also bowed and foreshortened [Fig. 8A, B]. At 14.5 dpc, the long bones of the mutants revealed a delay in formation of the diaphyseal ossification center, and the bowing was more pronounced [Fig. 8C–F]. Thus, the abnormalities evident in the limbs of the mutant neonates can be traced to a defect in formation and subsequent ossification of specific cartilaginous precursors.

Vertebral defects in MHox mutant mice

Analysis of skeletons of MHox mutant neonates revealed that ~12% of the MHox mutant mice had abnormalities of the dorsal aspects of the thoracic and lumbar vertebrae. The neural arches of the affected vertebrae were abnormally shaped [Fig. 9]. In wild-type neonatal mice, the neural arches form a convex structure that surrounds the spinal cord laterally while the cartilaginous precursor of the spinous process forms the dorsal covering of the spinal cord. In MHox mutant mice, the neural arches failed to form a convex structure but were, instead, flared laterally. In addition, the dorsal cartilaginous structures were truncated. The result of these abnormalities was spina bifida.

Discussion

We have analyzed the functions of the MHox homeobox gene by gene targeting in mice. MHox mutant mice showed defects in numerous skeletal structures that are derived from the neural crest, the lateral mesoderm, and the somitic mesoderm. The observed defects can be classified primarily as deletion or hypoplasia of specific elements that can be traced to absent and growth-retarded precursors. The MHox mutation affected both endochondral and membranous bones, which form by distinct pathways. In the case of endochondral bone, mesenchyme condenses into cartilage, which in turn grows and differentiates and is eventually replaced by bone. The formation of membranous bone is not preceded by a cartilaginous intermediate, rather, mesenchymal condensations directly give rise to osteoblasts [Ham 1987].
Figure 5. Abnormalities of the middle ear of MHox(-/-) mice. Lateral views of the dissected middle ear ossicles of wild-type (A,C,E) and MHox(-/-) (B,D,F) mice. (A,B) Malleus; (C,D) incus; and (E,F) stapes; (G,H) Diagrams of the malleus-incus-stapes complexes as they appear in the skulls of wild-type and mutant neonates, respectively. Note the complete absence of the tympanic and gonial bones in the mutant (B). The manubrium of the malleus in the MHox mutant is truncated, and the processus brevis is posteriorly displaced (A,B). The arrows in B and H point to an ectopic process of Meckel’s cartilage. The incus from the mutant (D) is fused to a persistent quadrate cartilage on the lateral aspect of the skull. The wild-type stapes is stirrup-shaped (E) while the mutant stapes is columelliform (F). (c) Crus of the stapes; (E) exoccipital bone; (F) frontal bone; (fp) foot plate of the stapes; (G) gonial bone; (I) incus; (LH) lesser horn of the hyoid; (lp) long process of the incus; (M) malleus; (mb) manubrium of the malleus; (mc) Meckel’s cartilage; (pb) processus brevis of the malleus; (rc) Reichert’s cartilage; (S) stapes; (SO) supraoccipital bone; (St) stylloid bone; (SL) stylohyoid ligament; (T) tympanic bone.

arch structure to be affected by the MHox mutation was the stapes, a derivative of Reichert’s cartilage.

Most knowledge of the embryonic origins of the vertebrate skull is extrapolated from chick-quail grafting experiments. These studies have shown that the cranial and visceral arch skeletons are derived from the cranial neural crest, the cephalic mesoderm, or the occipital somites (Noden 1988; Lumsden et al. 1991; Le Douarin et al. 1993). Neural crest cells that contribute to the first and second arches originate from the dorsal neural tube at the level of rhombomeres 1–4 and undergo an epithelial-to-mesenchymal transition as they migrate (Lumsden et al. 1991; Sechrist et al. 1993). Migration of cranial neural crest cells occurs in two waves: the first wave gives rise to cartilage, bone, and connective tissues, and the second to neurogenic derivatives (Nichols 1986). The affected neural crest-derived bones of the skull, such as the squamosal and zygomatic bones, form by membranous ossification. Ossification centers for these bones were never detected in MHox mutant mice during embryogenesis, suggesting that the formation of the precursors of these bones had been perturbed. The abnormalities detected in endochondral bone of the skull of mutant mice can be related to defects in formation of
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Table 1. Defects in skeletal structures derived from the pharyngeal arches in MHOX mutant mice

| Derivatives | Defect |
|-------------|--------|
| First arch  |        |
| Meckel's cartilage | ectopic process |
| malleus | truncated manubrium |
| incus | fused to cartilaginous mass in lateral skull |
| tympanic bone | absent |
| maxillary bone | missing palatal process |
| dentary (mandibular) bone | hypoplastic angular, coronoid process and head of ramus |
| palatine bone | remnant |
| squamosal bone | remnant |
| pterygoid bone | remnant |
| zygomatic bone | absent |
| Second arch |        |
| stapes | fusion of anterior and posterior crus, fusion to Reichert’s cartilage |

Cartilaginous precursors. For example, the precursors of the supraoccipital and alisphenoid bones were either missing or growth delayed.

The absence of skeletal elements of MHOX mutants was not likely a result of a defect in neural crest cell migration or specification because MHOX expression is first detected after the onset of these events [Cserjesi et al. 1992]. Moreover, expression of the neural crest marker AP2 [Mitchell et al. 1991] in the first branchial arch was normal in MHOX mutant embryos (not shown). The MHOX mutation also produced no detectable effect on the cranial ganglia, which form from neurogenic neural crest cells in the first and second arch (not shown). These findings suggest that only a subset of the mesenchymal neural crest require MHOX to complete their developmental program.

The neural crest populates the branchial arches and developing facial processes and subsequently gives rise to cartilage and bone anlagen by aggregation of loose mesenchyme. These condensations differentiate into the specific cartilage and bone of the skull. The ultimate differentiation of cranial skeletal precursors has been shown to require epithelial–mesenchymal interactions [Hall and Miyake 1992]. Previous work suggesting that MHOX expression in embryonic mesenchyme requires a signal from the overlying epithelium implies that MHOX function is important in epithelial–mesenchymal interactions that eventually result in the differentiation of mature cartilage and bone [Kuratani et al. 1994]. Interruption of these intercellular signaling pathways could account for the loss of the cranial bones observed in MHOX mutant neonates.

The MHOX mutation results in defects in a subset of long bones and vertebrae

During the development of endochondral long bones, chondrocytes within the diaphysis, or shaft, of a cartilage body proliferate and increase the length of the structure. These chondrocytes then mature and calcify and eventually are replaced by bone forming the diaphysial ossification center. Late in development and postnatally, long bones lengthen as a result of growth in the cartilage of the epiphyseal growth plate. In addition, osteoblasts located within the periosteum encasing the bone lay down matrix and increase the width of the bone by appositional growth [Ham 1987]. In MHOX mutant neonates, the affected endochondral long bones were shorter and thicker than normal. A defect was observed in the cartilaginous templates of these long bones and later in the diaphysial ossification center. However, the epiphyseal growth plates were unaffected by the MHOX mutation. Appositional growth also seemed to be intact because the width of the bone was not diminished.

Vertebral abnormalities were also detected in a small percentage of MHOX–mutant neonatal mice. Although this defect was not analyzed during ontogeny, MHOX expression has been detected within the condensing vertebrae of developing mice [Cserjesi et al. 1992], suggesting that MHOX plays a role in the formation of vertebral precursors. These results suggest that the endochondral bone abnormalities observed in MHOX mutant mice are secondary to a defect in chondrogenesis of cartilaginous precursors.

Although the developmental processes that give rise to endochondral and membranous bone are distinct in many ways, the formation of both bone types is initiated by the aggregation of mesenchyme into prechondrogenic and preosteogenic condensations. This critical stage of skeletogenesis precedes the overt differentiation of cartilage and bone and is dependent on epithelial–mesenchymal tissue interactions for its completion [Hall and Miyake 1992]. Because the MHOX mutation affects both bone types, derived from all three embryologic sources, it is likely that the function of MHOX is exerted at the condensation stage of skeletal development. The primary mechanisms directing formation of condensations are cell proliferation and adhesion, which result in increased local cell density. Analysis of MHOX mutant embryos for abnormalities of growth rates within condensations by BrdU labeling has not revealed differences from wild-type embryos [J. Martin, unpubl.]. However, it is possible that these experiments would not reveal subtle proliferative changes within the mutant condensations. Aberrations in the adhesive properties of cells within condensing mesenchyme is also a conceivable mechanism responsible for defects of condensations within MHOX mutant mice. In this regard, it is interesting to note that the promoter of the cell adhesion molecule, NCAM, has a required recognition element that has been shown to bind a related factor, Phox2 [Valarché et al. 1993].

Only a subset of embryonic structures derived from MHOX-expressing cells are affected by the MHOX mutation

During mouse embryogenesis, MHOX is expressed in mesenchymal cells within the branchial arches begin-
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MHox expression is observed at high levels in the limb buds, lateral mesoderm, the maxillary processes, the frontonasal mass, and the prevertebrae. Not all regions of MHox expression show defects of equal severity. In the limb buds, for example, where MHox is expressed at high levels, we detected only subtle defects in MHox mutant mice. The dichotomy of observed phenotypes between the head and limb in MHox mutant mice suggests that there may be other genes that can substitute for MHox functions within the limb. The homeodomain protein S8 would be most likely to substitute for MHox because these two homeo domain proteins are highly related within their homeo domains, bind the same DNA target sequence, and show overlapping expression patterns during mouse embryogenesis (Opstelten et al. 1991; Cserjesi et al. 1992, 1994).

**Relationship of the MHox mutant phenotype to other homeo box gene mutations**

The phenotype of MHox mutant mice is similar to, but distinct from those mice bearing mutations in other ho-
Figure 7. Limb skeleton of wild-type and MHox mutant neonates. Bones from the forelimb (A,B) and hindlimb (C,D) of wild type (A,C) and MHox mutant (B,D) neonates were stained for bone (red) and cartilage (blue). The radius, ulna, tibia, and fibula are truncated in the mutants, whereas the humerus, femur, and distal elements are normal. (f) Fibula; (r) radius; (t) tibia; (u) ulna.

Little is known about the molecular mechanisms regulating the development of the skull. A priori, it would be expected that insight into these mechanisms should also contribute to an understanding of how cranial diversity between taxa is generated. Embryology and studies in comparative anatomy have suggested that a common mechanism to create cranial diversity is heterochrony—variation in the timing of developmental events. For example, the emergence of snakes from reptilian ancestors is thought to have occurred by modification in the timing of formation of cranial precursors, eventually altering the morphology of the adult skull (Rieppel 1993). The MHox mutation appears to result in a cranial atavism secondary to altered growth rates of cartilaginous precursors, supporting the notion that heterochronic change is an important evolutionary mechanism used to create diversity in the vertebrate skull. Similar notions have been postulated recently to explain the phenotype observed in Hoxa2 and Hoxd13 mutant mice. Hoxd13 mutant mice demonstrate altered growth rates of cartilage precursors in the limb, which lead to atavistic and neotenic changes in morphology (Dollé et al. 1993). In Hoxa2–mutant mice, similarities of the palatoquadrate cartilage were identified (Rijli et al. 1993). The MHox mutation provides further experimental support for these ideas and so provides insight into the mechanisms that control ontogeny and evolution of the skull.

Materials and methods

Construction of MHox targeting vector

MHox genomic clones were isolated by screening a mouse 129/Sv genomic library with the full-length MHox cDNA (Cserjesi et al. 1992). Positive clones were purified and subcloned into the pBSK2 plasmid (Stratagene). To construct the MHox targeting vector, a neomycin resistance cassette under control of the phosphoglycerol kinase (PGK) promoter was inserted into a unique StuI site near the amino terminal region of the homeo domain. A pMC1–HSV thymidine kinase gene (Mansour et al. 1988) was placed into a Xhol site at the 3’ end of the vector. Prior to electroporation, the MHox targeting vector was linearized with NotI.
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Figure 8. Limb skeleton of wild-type and MHox mutant embryos. Whole-mount cartilage staining of wild-type (A,C,E) and mutant (B,D,F) embryos at 13.5 dpc (A,B) and 14.5 dpc (C-F). At 13.5 dpc the cartilaginous precursors for the radius and ulna of the upper limb are bowed and shorter than normal (A,B). At 14.5 dpc the defect in the radius and ulna of the upper limb (D) and the tibia and fibula of the hindlimb (F) are more apparent. The arrow in B points to the misshapen radial precursor. The arrows in C–F point to the diaphyseal ossification center, which is delayed in the mutant limb. (f) Fibula; (r) radius; (t) tibia; (u) ulna.

Generation of MHox-null mice

The MHox targeting vector (50 µg) was electroporated into AB-1 ES cells (McMahon and Bradley 1990) using a Bio-Rad gene pulser (500 µF, 240 V), and cells were plated on SNL76/7 cells and cultured under the positive and negative selection of G-418 and FIAU (McMahon and Bradley 1990). Addition of FIAU to the medium resulted in an approximate eight-fold enrichment for targeting events. Following selection, surviving clones were isolated and replica plated onto SNL76/7 fibroblast feeder cells in 96-well plates. Southern analysis was performed on these colonies as described (Ramirez-Solis et al. 1992) using an EcoRI digest and probe A (see Fig. 1A). Ten ES cell clones that were found to contain targeted events within a single MHox allele were expanded, and their genomic DNA was analyzed further by Southern analysis using multiple restriction enzymes to confirm the targeting events. Rearrangements in the vicinity of the MHox locus were not detected. Two independent clones were injected into 3.5 dpc mouse embryos that were reimplanted into foster mothers to generate high percentage chimeras. One chimera transmitted the mutation through the germ line, giving rise to mice heterozygous for the MHox mutation.

Genotyping of progeny

To identify mice carrying the MHox mutation, Southern blot analysis was performed on genomic DNA that was obtained from tail biopsies of neonatal and 10-day-old mice and from the yolk sacs of mouse embryos. To isolate genomic DNA, tissue was incubated in lysis buffer (10 mM Tris at pH 8.0, 25 mM EDTA at pH 8.0, 100 mM NaCl, 1% SDS, 0.2 mg/ml of proteinase K) at 50°C for 3 hr, followed by phenol–chloroform extraction and ethanol precipitation. Genomic DNA was digested with the indicated restriction enzyme and fractionated on a 0.7% agarose gel. Digested DNA was transferred to Zeta-probe GT membranes and hybridized with probe A or B.

RT–PCR analysis of MHox mRNA

Total RNA was isolated from neonatal mice as described previously (Chomczynski and Sacchi 1987). To synthesize cDNA, 1 µg of total RNA was denatured in the presence of random hexamers, followed by incubation with reverse transcriptase (BRL) at 42°C for 1 hr. One-fifth of the cDNA reaction mixture was used for the PCR reaction, which was performed under the following conditions: Denature at 94°C for 3 min, 1 cycle, 94°C for 30 seconds, 60°C for 30 seconds, 72°C for 1 min, 28 cycles; 72°C for 15 min, 1 cycle. Each PCR reaction mixture contained 0.5 µCi of [32P]dCTP for detection of the amplified products. The sequence of L7 oligonucleotides has been described (Hollenberg et al. 1993). The sequence of the oligonucleotides used for amplifying the MHox transcript were as follows: MH-B, 5'-ACC ATG ACC TCC AGC TAC GG-3' and MH-O, 5'-TCC
The vertebral skeletons from wild-type (A,C) and mutant Dissected first lumbar vertebrae from wild type (C) and mutant and lumbar spine of the mutant was abnormally widened (B). Figure cartilage that is truncated in the mutant. Vertebral arch; (vb) vertebral body. The arrows denote the dorsal is abnormally flared and the dorsal cartilage is truncated. (a) (D) are representative of the defect in which the vertebral arch neonates were stained for cartilage and bone. The lower thoracic tenth of the reaction mixture was fractionated on a 6% acrylamide gel.

Skeletons were cleared in 2% KOH and stored in 100% glycerol. Neonatal mice were eviscerated and placed in water overnight. Skeletons were immersed in a 65°C water bath for 1 min, skinned, and fixed in 100% ethanol for 3 days followed by alcin blue stain (15 mg of alcian blue 8Gx (Sigma), 80 ml of 95% ethanol, 20 ml of glacial acetic acid) for 8–12 hr. Skeletons were rinsed in 100% ethanol overnight and cleared in 2% KOH for 6 hr. Counterstaining for bone was performed using alizarin Red (Sigma, 50 mg/liter of 2% KOH) for 3 hr. Skeletons were cleared in 2% KOH and stored in 100% glycerol.

Histology
Embryos and neonates were fixed in Bouin’s solution or 4% paraformaldehyde overnight and then dehydrated through graded alcohols and embedded in paraffin. Paraffin blocks were sectioned at 7–10 μm and stained with hematoxylin and eosin.

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