A variant limb deformity transcript expressed in the embryonic mouse limb defines a novel formin

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The formins constitute a set of protein isoforms encoded by the alternatively spliced transcripts arising from the limb deformity (ld) locus of the mouse. Mutations in this locus disrupt formation of the anteroposterior axis of the embryonic limb. Although ld transcripts are widely expressed during embryogenesis, we have identified a novel transcript that is expressed in the mesenchyme and apical ectodermal ridge of the developing limb. This pattern of expression coincides with the earliest morphological defects observed in ld mutant limb buds. Moreover, the formin encoded by this transcript bears a highly acidic amino terminus, as distinguished from the basic amino terminus encoded by other ld transcripts, suggesting that it may have a distinct biochemical function.

[Key Words: Alternative splicing, apical ectodermal ridge, embryogenesis, limb development, pattern formation]

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Classic embryological manipulations combined with recent molecular analyses have established the developing limb as an important model system for elucidating the mechanisms of vertebrate pattern formation (for review, see Tabin 1991). Embryological studies, in particular, implicate reciprocal inductive interactions between the ectoderm and mesenchyme in establishing the limb pattern (Zwilling 1956a). Morphogenetic structures in both the limb ectoderm and mesenchyme are required in this process, namely, the apical ectodermal ridge (AER) and the zone of polarizing activity (ZPA). The AER is a rim of columnar epithelial cells at the distal tip of the limb bud that maintains the underlying mesenchyme in a proliferative and undifferentiated state (Saunders 1977). Patternning of the mesenchyme, or establishing the positional identity of a cell along the anteroposterior axis, has been proposed to be regulated by a morphogen released from the ZPA (Wolpert 1969), a group of mesenchymal cells at the posterior margin of the limb (Saunders and Casseling 1968).

Loss-of-function mutations that disrupt normal limb development define a set of genes required for limb morphogenesis (Grueneberg 1963; Lyon and Searle 1989). The phenotypes of the mouse mutant limb deformity (ld) indicate that the ld gene is essential for proper limb pattern formation (Kleinebrecht et al. 1982; Woychik et al. 1985). Specifically, the earliest embryonic manifestations of the limb defect include incomplete AER differentiation and a marked reduction in the anteroposterior axis of ld mutant limb buds (Zeller et al. 1989). These ld mutations result in abnormal morphogenesis of both the limb and the kidney. The five known recessive ld alleles all display similar phenotypes, including fusion of distal bones of the limb, reduction in the number of digits, and a variably penetrant renal aplasia (Cupp 1960; Kleinebrecht et al. 1982; Messing et al. 1990, Woychik et al. 1985, 1990a). Cloning of the ld gene, which encodes a novel set of proteins, the formins, has identified one of the genes necessary for correct patterning of the vertebrate limb (Woychik et al. 1985, 1990b).

Because a complex array of transcripts arises from the ld gene, owing to alternative splicing and differential polyadenylation (Woychik et al. 1990b), we have sought to characterize the expression pattern of these transcripts in the developing limb. Expression of ld transcripts is observed in a wide array of adult tissues and throughout the embryo. Consistent with this RNA analysis, Trumpp et al. (this issue) have localized the chicken ld gene products by immunohistochemistry on chick embryos to the developing limb, kidney, and central nervous system. Considering the tissue-specific alternative splicing of ld transcripts observed in adult mice (Woychik et al. 1990b), it is possible that these splice forms may also be expressed differentially during embryogenesis. Using a probe common to all ld transcripts, we have demonstrated previously ld expression in both developing limb bud ectoderm and mesenchyme (Zeller et al. 1989). Identification of the particular splice form expressed in the developing limb would implicate its involvement in limb morphogenesis.

Here, we report the differential expression of several ld transcripts during mouse embryogenesis. As only one of these characterized splice forms is expressed in the developing limb, we have focused our further analysis on
this form. The sequence of this transcript predicts a protein with an amino-terminal domain distinct from the formin sequences reported previously [Woychik et al. 1990b]. In addition, in situ hybridization and ribonuclease protection analyses reveal expression of this transcript in both the AER and mesenchyme of the developing limb. Combined with the known alterations of \( ld \) transcripts in \( ld \) mutants [Maas et al. 1990], these data suggest a role for this particular \( ld \) gene product in normal limb development.

Results

Expression of one \( ld \) splice form in the developing limb

To determine which \( ld \) transcripts are expressed in the developing organs affected by \( ld \) mutations, we have generated transcript-specific probes from four different partial cDNA clones corresponding to splice variations [see Materials and methods; Woychik et al. 1990b]. Ribonuclease protection analysis of representative adult tissue RNAs (Fig. 1A) shows coordinate expression of splice forms I, II, and III, with highest levels in testis, brain, and kidney. The tissue distribution of splice form IV is distinct from the other three transcripts, displaying qualitative or quantitative differences in expression in most tissues assayed. Expression patterns during embryogenesis, and particularly in developing limbs, were analyzed similarly using embryonic RNAs from gestational day 8.5–13.5 postcoitum (p.c.) (Fig. 1B). Again, striking differences in expression of these two classes of transcripts were observed. Splice forms I, II, and III display coordinate expression (Fig. 1A, and data not shown) that is first detectable by day 9.5 p.c. and increases in level in pro-

![Figure 1. Ribonuclease protection analysis of the \( ld \) splice forms expressed in adult tissues and during mouse embryogenesis. Twenty micrograms of total RNA was hybridized to splice form-specific riboprobes (for probe details, see Materials and methods) and analyzed by ribonuclease protection. The different splice forms are illustrated to schematically represent both the unique and the common exons for each transcript. (A) Embryo and limb bud RNAs were from gestational day 11.5 p.c. The bottom row indicates hybridization to a 3' probe that detects all \( ld \) transcripts [Zeller et al. 1989]. (B) ES RNA was from mouse embryonic stem cells (ES-D3; Doetschman et al. 1985) grown on gelatinized plates in the presence of leukemia inhibitory factor [1000 U/ml; Amrad] (without feeder cells). Embryonic RNAs were from gestational days 8.5–13.5 p.c.; the actual stages are 0.5 days later than indicated by the labeling. The upper panel corresponds to protection of a probe that detects splice forms I, II, and III; the lower panel corresponds to protection of the splice form IV probe. (H) Head; (B) body without limbs.](image-url)
gressively older embryos. In contrast, splice form IV is present at moderate levels much earlier in development than the other three forms, for example, as seen in day 8.5 p.c. embryos. In addition, this transcript is present in undifferentiated embryonic stem cells, which may reflect the in vivo expression of this splice form at earlier stages. However, the most notable difference in expression is in the developing limb, where only splice form IV is detected in limb buds dissected at days 10.5 and 11.5 p.c. (Fig. 1A,B). These early stages of limb development precede the formation of the cartilaginous skeletal elements [detailed by Martin 1990] and correspond to the stages when morphological defects are apparent in Id mutants [Zeller et al. 1989]. The sole expression of splice form IV in the limb during these stages led us to focus on cloning the 5' end of this transcript and refining our analysis of its expression pattern.

Isolation of Id cDNA clones from embryonic limb buds

To complete the characterization of the splice form IV limb transcript, we prepared and screened a cDNA library from day 11.0 to day 11.5 p.c. limb buds. Random hexamer-primed first-strand synthesis was used to enrich for clones containing 5' ends of the large 13-, 7-, and 5-kb Id transcripts [Woychik et al. 1990b]. A library of 5 \times 10^6 primary clones was screened with a DNA fragment from the 5' end of clone 4 [for details, see Materials and methods, Woychik et al. 1990b]. This screen yielded four additional overlapping cDNA clones [for details, see legend to Fig. 2]. The sequence of these cDNAs encodes a novel isoform of the set of Id gene products described previously, the formins [Woychik et al. 1990b]. The entire novel portion of this 5' cDNA sequence is contained within one exon (boxed in Fig. 2) that encodes a unique 457-amino-acid amino-terminal domain, as determined by sequence comparison with a corresponding genomic clone [described below]. The composite 4241-bp sequence of the cDNA clones with the deduced 1206-amino-acid open reading frame (ORF) is presented in Figure 2. Apart from the carboxy-terminal region common to formins I, II, and IV, this sequence does not display any significant similarities to sequences entered in the GenBank data base as of this writing. An obvious biochemical function of this formin is not apparent from its primary sequence. However, there is one notable distinction between the two different amino-terminal formin sequences. The splice form IV amino-terminal domain is acidic [calculated pl 4.5, calculated charge at pH 7.0 = -30], whereas the amino-terminal domain common to formin isoforms I, II, and III is basic [calculated pl 9.8, calculated charge at pH 7.0 = +18]. Such differences suggest that these formins may have distinct functions.

Given the structural complexity of the Id transcripts, we wanted to determine whether the composite sequence shown in Figure 2 represents the only splice form containing this 5' sequence. Polymerase chain reaction (PCR) amplification of embryonic and adult tissue first-strand cDNA carried out using an oligonucleotide primer from the 3' end of this new exon and primers from progressively 3' exons did not identify any additional splice forms [data not shown]. Because major differences in the lengths of Id transcripts result from differential polyadenylation [Woychik et al. 1990b], more subtle size differences between potential splice forms could be detected by conferring a uniform 3' end on all Id transcripts. This was done with RNase H cleavage of RNA after hybridization to a DNA fragment near the 3' end of the coding region [represented by the solid box in the diagram in Fig. 3]. Northern analysis of this uniformly cleaved RNA from salivary gland was hybridized with probes specific for the splice form IV 5' exon [Fig. 3, probe A] or the common 3' region [Fig. 3, probe B]. Both probes detect a single RNA species of ~4 kb, in contrast to the uncleaved transcript sizes of 13, 7, and 5 kb (Fig. 3). Similar but less intense signals were observed from parallel samples from day 10.5 p.c. embryonic RNA [data not shown]. These data confirm that the size heterogeneity of Id transcripts is predominantly a result of differential polyadenylation [Woychik et al. 1990b] and demonstrate that the 5' and 3' regions are linked on a segment of RNA that is comparable in size to the full-length coding region. Taken together with the multiple stop codons found upstream of the initiation codon, these results indicate that the overlapping cDNAs represent the full-length ORF, although they are probably missing a portion of the 5'-untranslated region present in the mRNA.

Genomic organization of the alternative exons

To define the genomic organization of the alternatively spliced Id exons, we isolated cosmid clones containing some of these exons. Restriction mapping was used to define exon order, and comparative sequence analysis of the cDNAs with the genomic coding regions was used to define exon borders [Fig. 4, and data not shown]. The position of two known Id mutations that disrupt the 3'-coding region common to all of these Id transcripts is illustrated in Figure 4 [Maas et al. 1990; Woychik et al. 1990b]. Splice forms I, II, and III use the most 5' amino-terminal-coding exon (the basic amino acid sequence) but splice to three different downstream exons. Splice form IV has a unique 5' end that encodes the new acidic amino-terminal domain described above.

Limb ectoderm and mesenchyme show differential expression of Id transcripts

Using a probe common to all Id transcripts, Id expression has been shown previously to be fivefold higher in limb ectoderm than in limb mesenchyme [Zeller et al. 1989]. Our analysis of the different Id transcripts present in total limb bud RNA [Fig. 1] suggested that these results probably reflected splice form IV expression. To test this directly, we performed a similar analysis on RNAs isolated from day 10.5 [data not shown] and day 11.5 p.c. embryos, dissected limb buds, and dissociated limb ec-
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Figure 2. Nucleotide sequence and predicted amino acid sequence of the mouse $Id$ composite limb cDNA. Nucleotide and amino acid positions are labeled at left and right, respectively. The 5'-boxed sequence is distinct from that reported by Woychik et al. (1990b) and left right, Jackson-Grusby et al. 32 sequence corresponds to a 108-bp segment 4 (from Woychik et al. 1990b) 1175-4241. A 3' region of divergence from the previous Id genomic clone. The initiation codon at position 226 is preceded by termination codons in all three frames. The 5' and 3' borders of 56 include nucleotides 1-1595 (amino acids 1-457). This unique sequence corresponds to a single exon and was confirmed from a...
transcripts in the limb probe: A
RNase H: - + B - + f epidermal cells defining the AER in limb ectoderm and is not observed in adjacent nonridge ectoderm. Hybridization of a control sense probe to an adjacent section showed no signal in the AER, indicating the specificity of the anti-sense probe. Mesenchymal expression of this transcript, shown to be at least fivefold lower than ectodermal expression by ribonuclease protection (Fig. 5), cannot be distinguished from background hybridization in these studies (Fig. 6C). It remains possible that the mesenchymal expression of this transcript is localized, but at levels below our limit of detection.

Figure 3. RNase H mapping defines the transcript size for the coding region. Duplicate samples containing 5 μg of salivary gland poly(A) RNA were annealed to a 200-bp DNA fragment (illustrated by the solid box 3' of probe B). One sample was digested with RNase H; both were then analyzed by Northern blot hybridization sequentially with probes A and B.

Limb ectoderm expression is localized to the AER
To further refine the pattern of expression of splice form IV in the limb, we analyzed day 10.5 p.c. mouse limb buds by in situ hybridization with a probe specific for this transcript (Fig. 3, probe A). As shown in Figure 6, A and B, expression of this transcript is localized to the

Discussion
Reciprocal inductive interactions between the ectoderm and mesenchyme are essential for proper patterning of the vertebrate limb (Tickle 1980; Kelly and Fallon 1981; Fallon et al. 1983; see also introductory section). Inductive signals from the mesenchyme are required both for AER formation (Reuss and Saunders 1965) and maintenance (Zwilling 1956b; Zwilling and Hansborough 1956), whereas the AER provides factors that keep the underlying mesenchyme undifferentiated (Globus and Vethamany-Globus 1976) and promote mesenchymal cell pro-

Figure 4. Structure of the Id gene and its alternatively spliced transcripts. The genomic structure [top] of the 5'-alternatively spliced exons was determined by sequence comparison of genomic and cDNA clones. Cloned exons are all indicated by unique fill patterns; coding regions not defined in genomic DNA are indicated by open boxes [L. Jackson-Grusby and R.P. Woychik, unpubl.]. Uncloned intervening sequences are indicated by the broken line. Positions of the known Id mutations are marked by arrows [Maas et al. 1990; Woychik et al. 1990a,b]. The exon usage of splice forms I-IV is shown beneath the genomic structure. Splice forms I, II, and III represent composite structures of cDNA clones from Woychik et al. (1990b). The sequence of splice form IV is shown in Fig. 2. The size of the exons relative to the ~200-kb locus [L. Jackson-Grusby, unpubl.] is not to scale but does approximate the relative size of the exons.
This suggests that the deficient AER resulting from \( ld \) mutations is not the sole cause of the \( ld \) limb patterning defects, but rather, \( ld \) mutations have a more direct effect on the patterning of limb mesenchyme. Splice form IV is the best candidate \( ld \) gene product for functioning in anteroposterior limb patterning, as it is the only \( ld \) splice form that we have detected in limb mesenchyme (Fig. 5). Consistent with this, the expression of the chick \( ld \) gene product has been localized by immunohistochemistry to

**Figure 5.** Differential expression of \( ld \) splice forms within the limb. Ribonuclease protection analysis of RNAs isolated from embryos (Emb), limb buds (Lb), and dissociated limb ectodermal jackets (Ect) and mesenchymal cones (Mes) all dissected between gestational days 11.0 and 11.5 p.c. The 315-nucleotide probe contains the 3' 119 nucleotides from the 5' exon of splice form IV plus the two adjacent alternatively spliced exons (181 nucleotides) and 15 nucleotides of polylinker. The different amounts of RNA for each sample are indicated above their respective lanes. Exposure time was 5 days, using an intensifying screen.

liferation (Solursh et al. 1981). As best described for the chick limb, the regulation of anteroposterior axis patterning requires a direct interaction between the AER and the mesenchymal ZPA (Saunders and Gasseling 1968; Tickle 1980).

An analysis of the \( ld \) mutant phenotype (Zeller et al. 1989) and the molecular characterization of the \( ld \) gene products that we describe suggest roles for \( ld \) in both of these morphogenetic structures. The embryonic limb phenotypes for the known recessive \( ld \) mutations include a foreshortened anteroposterior limb axis, aberrant or incomplete AER differentiation, as well as a more subtle phenotype of mesenchymal necrosis (Zeller et al. 1989). These multiple defects are most consistent with the requirement of \( ld \) function in both the AER and limb mesenchyme. Both ZPA transplantation experiments and embryological analyses of other limb mutants indicate a primary role of the mesenchyme in anteroposterior axis determination (Tickle 1980). The dependence of anteroposterior limb patterning on the AER has been analyzed in studies by Rowe and Fallon (1981). By partial AER removal in chick embryos, anteroposterior limb truncations somewhat similar to those observed in \( ld \) mutants were generated. Although these operations resulted in limbs with missing and fused digits as observed in \( ld \) mutants, they failed to produce fusions of the distal bones of the limb that are well documented in \( ld \) mu-

**Figure 6.** Expression of \( ld \) splice form IV in the AER. In situ hybridization analysis of \( ^{35} \)S-labeled riboprobes from the unique 5' exon of splice form IV on a day 10.5 p.c. embryonic limb bud. The probes correspond to the sense and anti-sense strands of probe A (Fig. 3). (A) Bright-field illumination of a section hybridized to the anti-sense probe. (B) Dark-field illumination of the section shown in A. The arrowhead indicates hybridization to the AER. (C) Dark-field illumination of the control sense probe hybridized to an adjacent section. Exposures were for 14 days. Bar, 100 \( \mu \)m.
coincide with the ZPA region [Trumpp et al., this issue]. Furthermore, we observe splice form IV expression in embryos prior to limb bud formation [Fig. 1B], which is consistent with a role for this transcript in the early determinative events involved in limb patterning.

Expression of splice form IV is also observed at relatively high levels in the AER [Fig. 6], in addition to a new splice form that we have identified in limb ectoderm [Fig. 5]. The consistency of this expression in the AER with the observed AER defect in ld mutants suggests that one or both of these gene products are required for complete AER differentiation. This deficiency in the AER of ld mutants has no apparent effect on proximal-distal limb patterning [Zeller et al. 1989], a well-established consequence of AER removal [Saunders 1948]. Thus, the function of ld in the AER is specific for the complete morphological differentiation of these cells because ld mutations do not disrupt all AER functions.

In addition to its expression in the developing limb, splice form IV is also observed throughout the embryo together with splice forms I, II, and III [Fig. 1]. However, the known ld phenotypes indicate a functional requirement for the ld gene products only in the developing limb and kidney. Several possibilities could account for the discordance of the observed expression pattern of ld transcripts with the more restricted ld phenotypes. If the known ld alleles represent the null phenotype, it is possible that partially redundant functions exist between ld and other genetic loci in regions of ld expression apart from the limb and kidney [Maas et al. 1990 and references therein]. Alternatively, expression of ld in these other regions may not be essential, or phenotypic defects in these regions might be too subtle to have been noted.

It is not necessarily the case that the known ld alleles are null mutations. Nonetheless, two lines of evidence are consistent with this interpretation. First, the known ld mutations disrupt the coding region for most ld gene products [Maas et al. 1990], including splice forms I, II, and IV. The consequence of these mutations on the encoded formins is a predicted truncation of the carboxyl terminus involving a loss of ~10% in the structure of these relatively large proteins [illustrated in Fig. 4]. Second, all of the known ld alleles display fairly uniform limb and renal phenotypes, whereas variation in phenotype might have been expected from different partial loss-of-function mutations [Cupp 1960; Kleinebrecht et al. 1982; Woychik et al. 1985, 1990a; Zeller et al. 1989, Messing et al. 1990]. On the other hand, these observations are also consistent with the possibility that the known ld alleles are partial loss-of-function mutations and that null alleles result in embryonic lethality. This interpretation is supported by the fact that the known mutations in this widely expressed gene have such a tissue-specific phenotype. The known ld mutants may represent the strongest possible viable ld phenotype. Targeted mutagenesis in mouse embryonic stem cells [Gosslor et al. 1986; Thomas and Capecchi 1987] should allow us to produce unequivocal null mutations to decide this point and to discern the potentially distinct roles of the different ld gene products in vertebrate development.

Transcripts from the ld gene encode various distinct proteins, the formins [Woychik et al. 1990b]. The complex expression patterns of these transcripts (Fig. 1), along with the striking differences in their deduced primary sequences, suggest that the different formins may have distinct functions. We have identified a transcript [splice form IV] expressed in the limb that encodes a potentially new formin [Fig. 2]. The unique amino-terminal exon used in this transcript encodes an acidic domain, in contrast to the other formins, which have basic amino termini [Woychik et al. 1990b]. Interestingly, Trumpp et al. [this issue] have recently cloned the chick ld transcript homologous to splice form IV, which revealed the conservation of the acidity of this domain (calculated pt 4.5, calculated charge at pH 7.0 = —34), although the absolute sequence conservation of this domain (~40%) is much lower than that of the carboxy-terminal region (~80%). In addition, we have identified another splice form, encoding an as yet uncharacterized formin, which is expressed in limb ectoderm along with splice form IV [Fig. 5]. Although we do not know whether these gene products act independently or whether they interact, it is intriguing to speculate that expression of different combinations of ld gene products may result in distinct ld gene functions.

Molecular analysis of the ld transcripts presented in this work suggests an important role for the ld splice form that we have characterized in patterning of the anteroposterior axis of the limb and AER differentiation. Expression patterns of other candidate regulatory genes in the limb, such as homeo box, retinoic acid receptor, and growth factor genes, also suggest their involvement in limb patterning [for review, see Tabin 1991]. Our identification of the transcripts from the ld locus involved in limb morphogenesis will allow the continued analysis of the function of ld gene products and their interaction with other genes in regulating limb development.

Materials and methods

Probes

Splice form-specific probes for forms I, II, and III were generated by PCR using the different clones described by Woychik et al. (1990b) as templates. The probe structures with reference to the original cDNA clone number and the nucleotide sequence of the probe borders are as follows: splice form I (clone 9) 1980–2180; splice form II (clone 7) 1980–2042 + 2338–2460; splice form III (clone 5) 1980–2042 + 2398–2460; splice form IV (clone 4) 1476–1775 [Fig. 3]. Splice forms I, II, and III refer to the ld sequence with the Genbank accession number X53599 [Woychik et al. 1990b]. The six other types of cDNA clones described by Woychik et al. (1990b) represent either differences in the 3′ untranslated region as a result of differential polyadenylation, or testis-specific transcripts, and were not used in our analysis.

PCR reactions with Taq polymerase [Cetus] were done according to the manufacturer’s recommendations. The reaction products were cloned into Bluescript pBS(−) and sequenced to eliminate clones with PCR artifacts. The splice form IV probe is a subclone of the 612 bp EcoRI–PstI fragment from the 5′ end of clone 4 [Woychik et al. 1990b] into pGEM-1, cut with Pvull to
generate the 300-nucleotide-protected fragment shown above. A probe containing the 200-bp BamHI–HindIII fragment (1974–1983) of the 5’ exon common to splice forms I, II, and III detects the expression of all three of these splice forms. The probe specific for the 5’ exon of splice form IV is a 517-bp BamHI–PstI fragment (421–937, Fig. 2; probe A, Fig. 3).

Embryonic dissections
Dissected embryos were derived from FVB/N matings. Gestational age was defined as the number of days p.c., such that the morning of vaginal plugging was designated day 0.5 as described by Rugh (1990). Embryos were also staged according to Rugh to eliminate any embryos that did not correspond to the appropriate developmental age. Dissection of limb buds and dissociations of limb ectoderm and mesenchyme as were described in Zeller et al. (1989). Dissociated ectodermal jackets and mesenchymal cones were sorted using a dissecting microscope, transferred to fresh Hanks balanced salt solution (HBSS) containing 10% serum, and resorted twice to ensure clean separations. All partially dissociated limb buds were discarded.

Ribonuclease protection
Total RNA was isolated using the method of Chirgwin et al. (1979). Ribonuclease protection was performed essentially as described by Krieg and Melton (1987). Probes were purified using Millipore Ultra-free spin filters. Radioactive probe (1 x 10^6 cpm) was hybridized to 20 μg of total RNA except as noted for the limb ectoderm in Figure 5.

cDNA library construction and screening
Poly[A]^+ RNA was prepared from gestational day 11.0 to day 11.5 prechondrogenic mouse limb buds (inbred strain FVB/N), using the methods of Chirgwin et al. (1979) and Aviv and Leder (1972), with two rounds of purification on oligo(dT)–cellulose (Collaborative Research). Random hexamer-primed double-stranded cDNA was prepared (Boehringer Mannheim cDNA synthesis kit) and fractionated by agarose gel electrophoresis to select for cDNAs >600 bp. Limb cDNA was modified by the addition of EcoRI adapters (Promega), ligated to vector arms (Lambda ZapII, Stratagene), and packaged in vitro (Gigapack Gold II, Stratagene). Poly[A]^+ RNA (1.0 μg) yielded a library of 5 x 10^4 independent clones, which were plated without amplification and screened by filter hybridization following standard methods (Sambrook et al. 1989).

Sequence analysis
Sequence of both strands of the largest and 5’-most splice form IV cDNA, plus 250–300 bp of each end of the other cDNA clones, was determined by deoxyxide chain termination using T7 DNA polymerase (Pharmacia). A genomic subclone containing this 5’ exon was also sequenced to verify this unique region, which is not present in the previously published sequence (Woychik et al. 1990b). Computer-assisted sequence analysis was done with software from the Genetics Computer Group, University of Wisconsin (Madison). The sequence is deposited in GenBank under accession number X62379.

RNase H mapping
Five micrograms of poly[A]^+ RNA from salivary gland or whole day-10.5 p.c. embryos was hybridized to 20 ng of a 200-bp Id 3’ region PstI restriction fragment in 30 μl of 50% formamide, 0.4 M NaCl, 0.04 M Pipes (pH 6.4), and 1 mM EDTA at 55°C overnight, after heating for 10 min at 90°C. The samples were diluted to 200 μl with 0.3 M NaAc (pH 5.2) and precipitated with 2.5 volumes of ethanol. The washed and dried precipitates were resuspended in 20 mM Tris-HCl (pH 7.5), 10 mM MgCl₂, 100 mM KCl, 0.1 mM dithiothreitol, and 5% sucrose and incubated for 30 min with 1 unit of RNase H (BRL) at 37°C (Irminger et al. 1987). The digestion was terminated by the addition of EDTA to 10 mM, extracted with phenol–chloroform, and ethanol-precipitated. The digestion products were analyzed by Northern blot hybridization after electrophoresis in formaldehyde–agarose gels. Filters were hybridized to random hexamer-labeled restriction fragments as described (Sambrook et al. 1989).

Cosmid cloning
A cosmid library was constructed using 35- to 50-kb fragments of BALB/c liver DNA generated by partial digestion with Sau3A, phosphatase treatment, and fractionation through two rounds of salt gradient ultracentrifugation. The vector was modified from the double COS site vector C2RB (Bates and Swift 1983) to contain RNA polymerase promoters and NotI sites flanking the BamHI cloning site and internal to the flanking EcoRI sites. Vector was cut with Smal, phosphatased, and cut with BamHI. One microgram of vector was ligated to 0.5 μg of genomic insert DNA in 5 μl. In vitro packaging (Gigapack XL, Stratagene) of 1 μl of the ligations yielded 4 x 10^5 CFU, which were screened directly without amplification.

In situ hybridization
Day 10.5 p.c. embryos were fixed for 1–4 hr in 4% paraformaldehyde, embedded in paraffin, cut into 2-μm sections, and placed on gelatin-coated microscope slides prior to hybridization. Synthesis of 35S-labeled riboprobes was from templates linearized to yield sense or anti-sense versions of the 517-bp BamHI–PstI fragment [probe A in Fig. 3] subcloned into BlueScript II KS(–)]. Sections were pretreated, hybridized, and washed essentially as described by Wilkinson et al. (1987), with some modifications (Sassoon et al. 1988). Slides were dipped in emulsion (Kodak NTB-2) and exposed for 14–21 days before developing.

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References
Aviv H. and P. Leder. 1972. Purification of biologically active globin messenger RNA by chromatography on oligo-
thymidylic acid-cellulose. *Proc. Natl. Acad. Sci.* 69: 1408–1412.

Bates, P.F. and R.A. Swift. 1983. Double cos site vectors: Simplified cosmid cloning. *Gene* 26: 137–146.

Chirgwin, J.M., A.E. Pryzbyla, R.J. MacDonald, and W.J. Rutter. 1979. Isolation of biologically active ribonucleic acid from sources enriched in ribonucleic acid. *Biochemistry* 18: 5294–5299.

Cupp, M.B. 1960. *Mouse News* 22: 50.

Doetschman, T.C., H. Eistetter, M. Katz, W. Schmidt, and R. Kemler. 1985. The *in vitro* development of blastocyst-derived embryonic stem cell lines: Formation of visceral yolk sac, blood islands and myocardium. *J. Embryol. Exp. Morphol.* 87: 27–45.

Fallon, J.F., D.A. Rowe, J.M. Frederick, and B.K. Simandl. 1983. Studies on epithelial-mesenchymal interactions during limb development. In *Epithelial-mesenchymal interactions in development* [ed. R.H. Sawyer and J.F. Fallon], pp. 3–25. Praeger Scientific, New York.

Globus, M. and S. Vethamany-Globus. 1976. An *in vitro* analogue of early chick limb bud outgrowth. *Differentiation* 6: 91–96.

Gossler, A., T. Doetschman, R. Korn, E. Serfling, and R. Kemler. 1986. Transgenesis by means of blastocyst-derived embryonic stem cell lines. *Proc. Natl. Acad. Sci.* 83: 9065–9069.

Grueneberg, H. 1963. The pathology of development. *A study of inherited skeletal disorders in animals.* Blackwell Scientific Publications, Oxford, England.

Irmscher, J.-C., K.M. Rosen, R.E. Humbel, and L. Villa-Komaroff. 1987. Tissue-specific expression of insulin-like growth factor II mRNAs with distinct 5' untranslated regions. *Proc. Natl. Acad. Sci.* 84: 6330–6334.

Kemler. 1985. The *in vitro* development of blastocyst-derived embryonic stem cell lines: Formation of visceral yolk sac, blood islands and myocardium. *J. Embryol. Exp. Morphol.* 87: 27–45.

Cupp, M.B. 1960. *Mouse News* 22: 50.

Doetschman, T.C., H. Eistetter, M. Katz, W. Schmidt, and R. Kemler. 1985. The *in vitro* development of blastocyst-derived embryonic stem cell lines: Formation of visceral yolk sac, blood islands and myocardium. *J. Embryol. Exp. Morphol.* 87: 27–45.

Fallon, J.F., D.A. Rowe, J.M. Frederick, and B.K. Simandl. 1983. Studies on epithelial-mesenchymal interactions during limb development. In *Epithelial-mesenchymal interactions in development* [ed. R.H. Sawyer and J.F. Fallon], pp. 3–25. Praeger Scientific, New York.

Globus, M. and S. Vethamany-Globus. 1976. An *in vitro* analogue of early chick limb bud outgrowth. *Differentiation* 6: 91–96.

Gossler, A., T. Doetschman, R. Korn, E. Serfling, and R. Kemler. 1986. Transgenesis by means of blastocyst-derived embryonic stem cell lines. *Proc. Natl. Acad. Sci.* 83: 9065–9069.

Grueneberg, H. 1963. The pathology of development. *A study of inherited skeletal disorders in animals.* Blackwell Scientific Publications, Oxford, England.

Irmscher, J.-C., K.M. Rosen, R.E. Humbel, and L. Villa-Komaroff. 1987. Tissue-specific expression of insulin-like growth factor II mRNAs with distinct 5' untranslated regions. *Proc. Natl. Acad. Sci.* 84: 6330–6334.

Kelley, R.O. and J.F. Fallon. 1981. The developing limb: An analysis of interacting cells and tissues in a model morphogenetic system. In *Morphogenesis and pattern formation.* [ed. T.G. Connelly et al.], pp. 49–85. Raven Press, New York.

Kleinebrecht, J., J. Selow, and W. Winkler. 1982. The mouse mutant *limb-deformity* (ld). *Anat. Anz.* 152: 313–324.

Krieg, P.A. and D.A. Melton. 1987. *In vitro* synthesis with SP6 RNA polymerase. *Methods Enzymol.* 155: 397–415.

Lyon, M.F. and A.G. Searle. 1989. Genetic variants and strains of the laboratory mouse, 2nd ed. Oxford University Press, Oxford, England.

Maas, R.L., R. Zeller, R.P. Wojcik, T.F. Vogt, and P. Leder. 1990. Disruption of formin-encoding transcripts in two mutant *limb deformity* alleles. *Nature* 346: 853–855.

Martin, P. 1990. Tissue patterning in the developing mouse limb. *Int. J. Dev. Biol.* 34: 323–336.

Messing, A., R.R. Behringer, J.R. Slepak, G. Lemke, R.N. Palmer, and R.L. Brinster. 1990. Insertional mutation at the LD locus (again!) in a line of transgenic mice. *Mouse News* 87: 107.

Reuss, C. and J.W. Saunders Jr. 1965. Inductive and axial properties of prospective limb bud mesoderm in the early chick embryo. *Am. Zool.* 5: 214.

Rowe, D.A. and J.F. Fallon. 1981. The effect of removing posterior apical ectodermal ridge of the chick wing and leg on pattern formation. *J. Embryol. Exp. Morphol.* 65: 309–325.

Rugh, R. 1990. *The mouse. Its reproduction and development.* Oxford University Press, Oxford, England.

Sambrook, J., E.F. Fritsch, and T. Maniatis. 1989. *Molecular cloning: A laboratory manual*, 2nd ed. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York.

Sassoon, D.A., I. Garner, and M. Buckingham. 1988. Transcripts of alpha-cardiac and alpha-skeletal actins are early markers for myogenesis in the mouse embryo. *Development* 104: 155–164.

Saunders, J.W. Jr. 1948. The proximo-distal sequence of origin of the parts of the chick wing and the role of the ectoderm. *J. Exp. Zool.* 108: 363–403.

———. 1977. The experimental analysis of chick limb bud development. In *Vertebrate limb and somite morphogenesis* [ed. D.A. Ede, J.R. Hinchliffe, and M. Balls], pp. 1–24. Cambridge University Press, Cambridge, England.

Saunders, J.W. Jr. and M.T. Gasseling. 1968. Ectodermal-mesenchymal interactions in the origin of limb symmetry. In *Epithelial-mesenchymal interactions* [ed. R. Fleischmajer and R.E. Billingham], pp. 78–97. Williams and Wilkins, Baltimore, MD.

Solursh, M., C.T. Singley, and R.S. Reiter. 1981. The influence of epithelia on cartilage and loose connective tissue formation by limb mesenchyme cultures. *Dev. Biol.* 86: 471–482.

Tabin, C.J. 1991. Retinoids, homeoboxes and growth factors: Toward molecular models for limb development. *Cell* 66: 199–217.

Thomas, K.R. and M.R. Capecchi. 1987. Site-directed mutagenesis by gene targeting in mouse embryo-derived stem cells. *Cell* 51: 503–512.

Tickle, C. 1980. The polarizing region and limb development. In *Development in mammals* [ed. M.H. Johnson], vol. 4, pp. 101–136. Elsevier/North-Holland Biomedical Press, Amsterdam, The Netherlands.

Wilkinson, D.G., J.A. Bailes, and A.P. McMahon. 1987. A molecular analysis of mouse development from 8 to 10 days post coitum detects changes only in embryonic globin expression. *Development* 99: 493–500.

Wolpert, L. 1969. Positional information and the spatial pattern of cellular differentiation. *J. Theor. Biol.* 25: 1–47.

Woychik, R.P., T.A. Stewart, L.G. Davis, P. D'Eustachio, and P. Leder. 1985. An inherited limb deformity created by insertional mutagenesis in a transgenic mouse. *Nature* 318: 36–40.

Woychik, R.P., W.M. Generoso, L.B. Russell, K.T. Cain, N.L.A. Cacheiro, S.J. Bultman, P.B. Selby, M.E. Dickinson, B.L.M. Hogan, and J.C. Rutledge. 1990a. Molecular and genetic characterization of a radiation-induced structural rearrangement in mouse chromosome 2 causing mutations at the *limb deformity* and *agouti* loci. *Proc. Natl. Acad. Sci.* 87: 2588–2592.

Woychik, R.P., R.L. Maas, R. Zeller, T.F. Vogt, and P. Leder. 1990b. "Formins": Proteins deduced from the alternative transcripts of the *limb deformity* gene. *Nature* 346: 850–853.

Zeller, R., L. Jackson-Crusby, and P. Leder. 1989. The *limb deformity* gene is required for apical ectodermal ridge differentiation and anteroposterior limb pattern formation. *Genes & Dev.* 3: 1481–1492.

Zwilling, E. 1956a. Interactions between limb bud ectoderm and mesoderm in the chick embryo. I. Axis establishment. *J. Exp. Zool.* 132: 157–172.

Zwilling, E. 1956b. Interactions between limb bud ectoderm and mesoderm in the chick embryo. IV. Experiments with a wingless mutant. *J. Exp. Zool.* 132: 241–253.

Zwilling, E. and L. Hansborough. 1956. Interactions between limb bud ectoderm and mesoderm in the chick embryo. III. Experiments with polydactylyous limbs. *J. Exp. Zool.* 132: 219–239.
A variant limb deformity transcript expressed in the embryonic mouse limb defines a novel formin.

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