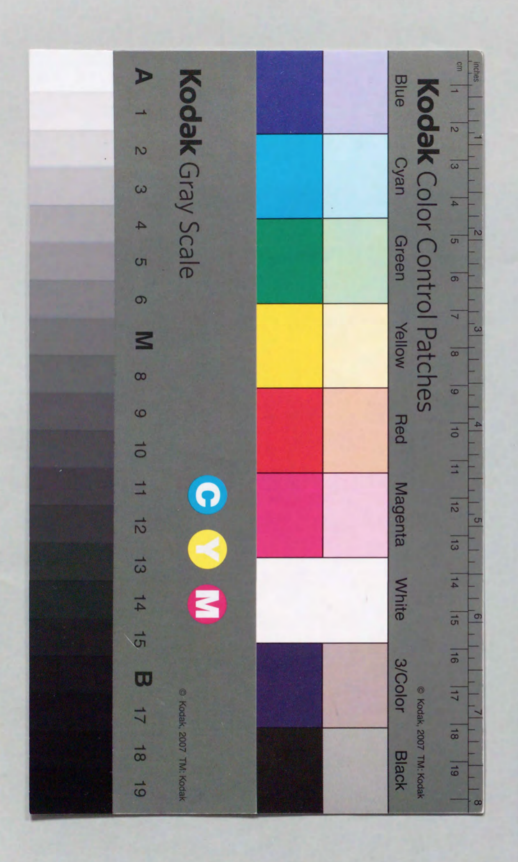


Title	Regionalized expression of the Dbx family homeobox genes in the embryonic CNS of the mouse
Author(s)	Shoji, Hiroki
Citation	大阪大学, 1996, 博士論文
Version Type	VoR
URL	https://doi.org/10.11501/3113111
rights	
Note	

Osaka University Knowledge Archive : OUKA

https://ir.library.osaka-u.ac.jp/

Osaka University



Regionalized expression of the Dbx family homeobox genes in the embryonic CNS of the mouse

HIROKI SHOJI

Regionalized expression of the Dbx family homeobox genes in the embryonic CNS of the mouse

マウス Dbx ホメオボックス遺伝子群の同定と 中枢神経系発生過程における領域特異的な発現

by

Hiroki Shoji

THESIS

Contents

Summary2	
Introduction	
Materials and Methods5	
Genomic and cDNA libraries5	ĺ
DNA sequencing5	
Chromosomal mapping5	
In situ hybridization6	,
Results8	
Genomic and cDNA clones of <i>Dbx</i> gene family	
Chromosomal location of Dbx210	
Expression of Dbx and $Dbx2$ in the embryonic CNS	
(1) Expression in the telencephalon	
(2) Expression in the diencephalon	
(3) Expression in the mesencephalon	
(4) Expression in the rhombencephalon and spinal cord	
Expression of $Dbx2$ outside of the CNS	1
(1) Expression in the limb buds	
(2) Expression in the tooth germs	
Discussion	,
Dbx gene family	,
Expression of Dbx and $Dbx2$ in the embryonic brain	
Expression in the spinal cord	
Expression of homologues in other animal species	
Complementary expression teritories with other homeobox genes 18	,
Expression in the limb buds and tooth germs	
Acknowledgements)
Acknowledgements	
References 2 1	
Figures	

Summary

Here I report the identification of a novel homeobox gene family Dbx in mouse, which consists of Dbx and Dbx2. The two genes share similar structural organization and are encoded by different chromosomes. The predicted Dbx and Dbx2 proteins share 85% identity in their homeodomain amino acid sequences, but otherwise showed no significant similarity. Characterization of the expression of these two genes in the embryos suggested their role in the development of the CNS. In the forebrain, Dbx is expressed in various regions, while Dbx2showed a more restricted pattern of expression. In the midbrain, the expression domains of Dbx and Dbx2 overlap along the dorso-lateral wall of the ventricle. In the hindbrain and spinal cord, both genes are expressed in the boundary separating the basal and alar plates, which seems to correspond to the sulcus limitans. Expression of the Dbx/Dbx2genes is restricted to the ventricular region of the embryonic CNS except for that of Dbx in the septum of the telencephalon. Together these observations indicate possible participation of the members of the Dbx family in regionalization of the CNS. While the expression of Dbxwas restricted to the CNS, Dbx2 was also expressed in some of the mesenchymal cells, such as limb buds and tooth germs.

commonutes, and are differentially expressed with individually among commonuted specificity in various embryonic anlages depending on the pastern within the clusters analogous to the Drasophilas Ham-C the reviews. Resent and Grant. 1990; Kenyon, 1994; Kramhaf, 1994). Other management and Grant. 1990; Kenyon, 1994; Kramhaf, 1994). Other management and Grant. 1995.

al. 1990).

Introduction

Our understanding of the molecular mechanisms underlying embryonic morphogenesis has been accelerated by genetic studies achieved in *Drosophila melanogaster*. Number of regulatory genes has been identified which have been integrated into a cascade of genetic networks (for review, Akam, 1987). Homeoboxes were discovered as highly conserved 180 bp sequence motifs shared by a number of genes involved in morphogenesis (McGinnis, et al., 1984; Scott and Weiner, 1984). Homeoboxes are translated into homeodomains which serve as DNA binding domains in transcriptional regulation (for reviews, Levine and Hoey, 1988; Scott et al., 1989; Hayashi and Scott; 1990; Gehring et al., 1994).

Homeoboxes are highly conserved among various animal species not only in terms of the basic feature of coding amino acid sequences but also in the genomic composition of the homeobox-containing genes. In several animal species, a family of homeobox-containing genes have been identified, and classified into subfamilies based upon the amino acid sequences of their homeodomains. In mouse, nearly a hundred homeobox genes have been identified (Kappen et al., 1993). Murine *Hox* genes are organized into four analogous gene clusters on different chromosomes, and are differentially expressed with individually unique rostro-caudal specificity in various embryonic anlages depending on the position within the clusters analogous to the *Drosophila Hom-C* (for reviews, Kessel and Gruss, 1990; Kenyon, 1994; Krumlauf, 1994). Other homeobox genes, such as *Pax* and *Evx1* are specifically expressed along the dorso-ventral axis of the CNS (Gruss and Walther, 1992; Bastian et al., 1990).

matrackettel.

In several unional species, a family of nomeobovecontaining peners have been streetled, and classified into contamilies based upon the amino Involvement of the homeobox genes in regional specification of the embryonic anlages, as suggested by analysis of their embryonic expression patterns, have been demonstrated by the phenotypes of the mutant animals lacking function of one of these genes (for reviews, Gruss and Walther, 1992; McGinnis and Krumlauf, 1992; Krumlauf, 1994).

The majority of vertebrate homeobox genes have been identified by their similarity to Drosophila homeobox genes. In order to identify homeobox genes including those which are highly divergent from Drosophila genes, I screened a mouse genomic library using a set of degenerated oligonucleotides corresponding to conserved motif in the homeodomain. This enabled me to isolate a new family of homeobox genes named Dbx (developing brain homeobox genes) which consists of Dbx (Lu et al. 1992) and Dbx2. These two genes are similar in their homeobox sequences and in gene organization, and are both expressed in the embryonic CNS with regional specificities.

the fragment of DhareDNA clone-tell-40 to Fig. 11 in the same buller of C. followed by two washes with Ly SSC, 0.18 SDS for 30 min at 1 9C. Screening of the width history ming in 0.75 an Eco R1 8 and in Fig. 11 was note in the same hybridization condition as described on the SA in the same hybridization condition as described on the SA followed by two washes with DawsSC, D 12 SASS for the same hybridization condition as described on the SA followed by two washes with DawsSC, D 12 SASS for the same hybridization condition as described on the same hybridization condition

DNA sugirating

ming Sequence Version 2.0 DNA Sequencing Elv (US Burchemical)

Circumsonal mapping

Materials and Methods

Genomic and cDNA libraries

BALB/c mouse liver DNA was partially digested with Sau3AI and fractionated by sucrose density gradient centrifugation. Fragments with the size of 15-20 kb were ligated to λEMBL3 vector arms (Clontech) and packaged in vitro (Stratagene). cDNAs for polyA+RNAs from 12.5 day ICR mouse embryos were methylated by EcoRI methylase, ligated to EcoRI linkers and digested with EcoRI. cDNAs larger than 500 bp were selected by Sepharose-CL4B column chromatography, and cloned into a \(\lambda \text{gt10} \) vector. Screening of the genomic library using the synthetic oligonucleotide probe was done in hybridization buffer of 6x SSC, 0.1% SDS, 0.2% Denhardt's solution, 100 μg/ml denatured salmon sperm DNA, at 55 °C overnight, followed by 3 washes with 2x SSC, 0.1% SDS for 15 min at 55 °C. Screening for isolation of Dbx2 genomic clone (gS4) was done using as probe the 0.56 kb fragment of Dbx cDNA clone (c164) (c in Fig. 1) in the same buffer at 65 °C, followed by two washes with 1x SSC, 0.1% SDS for 30 min at 65 °C. Screening of the cDNA library using a 0.75 kb EcoRI-BamHI fragment of g164 or a 0.2 kb EcoRI fragment of gS4 as probes (a and f in Fig. 1) was done in the same hybridization condition as described above for gS4, followed by two washes with 0.1× SSC, 0.1% SDS for 30 min at 65 °C.

DNA sequencing

DNA was sequenced by the dideoxy method (Sanger et al., 1977) using Sequenase Version 2.0 DNA Sequencing Kit (US Biochemical).

Chromosomal mapping

Intersubspecific backcross progeny were generated by mating (C57BL/6 \times MSM) F₁ females to MSM males (kindly provided by Drs. K. Moriwaki and N. Miyashita). DNA isolation, polymerase chain reaction (PCR) and polyacrylamide gel electrophoresis were performed as described (Dietrich et al, 1994). The primer sequences used for D15Mit15 and D15Mit40 were the same as described (Dietrich et al., 1994).

In situ hybridization

In situ hybridization with sections was done essentially as described previously (Wakamatsu and Kondoh, 1990). RNA probes were synthesized in vitro by transcription of a linear DNA template of 3' untranslated regions of c164 and gS4 (d and e in Fig. 1) using 200 μCi[35S]-UTP and T7 RNA polymerase (Toyobo). Paraffin sections of 6.5-15.5 day embryos of ICR mice were prepared on glass slides. The slides were treated with 0.2 N HCl and proteinase K, and then acetylated. Hybridization was done for 20 hours at 50 °C in 50% formamide, 0.2% Denhardt's solution, 2x SSC, 10 mM EDTA, 10% Dextransulphate, 500 μg/ml denatured salmon sperm DNA, 500 μg/ml yeast tRNA, 10 mM DTT, and with ³⁵S-labelled RNA probes. After hybridization, the slides were treated with 20 μg/ml RNase A and 1 unit/ml RNase T1 at 37 °C for 30 minutes, and washed in 0.1x SSC, 20% formamide, 10 mM DTT at 50 °C overnight. Autoradiography was done by exposing the slides to Kodak NTB-2 emulion for 1-2 weeks at 4 °C and by developing with Konikadol X (Konika).

Whole-mount in situ hybridization of 9.5 day embryos was performed as described by Sasaki and Hogan (1993) for Dbx. However, hybridization signal with Dbx2 was too faint to detect, so a procedure described by Kobayashi et al. (1994) was employed to improve signal

Intersubspecific threating property were generated by memory CS7BL76 x MSM1 39 tensine to MSM males throstly provided by Des K. Moriwaki and M. Missakhtat. 1376A declarion, polymerate chain reaction (PCR) and polymerylamids gut electrophorests were performed as described (Dietrich et al. 1994). The primer sequences used for DISMirtS and DISMirtS and DISMirtS and DISMirtS and DISMirtS and DISMirts were the same as described thetrich et al.

In the hybridization with authors was done excentally in the critical previously (Wakananes and Rondon, 1990). ANA probose were synthesized in other by quancilprion of a linear DNA tamplate of the minearistical argums at citif and xNd (d and e in Fig. 1) using 200 and (c) 25-1078 and 17 RMA polymerise (Toyolor). Partilia sections of 6.5-13.5 day embryos of ECR mice were proposed on glass slides. The slides were treated with 0.2 N IR 1 and probance K, and then nearlylaned likybridization was done to 20 hours at 50 °C in 50% formandes, 0.2% Deshardix volution. 2x SMC 10 mit EDTA, 10% Deshardix volution. 2x SMC 10 mit EDTA, 10% Deshardix to and with 25-1abellos KNA probes. After hybridization, the slides were treated with 20 agint RNAs probes. After hybridization, the slides were treated with 20 agint RNAs probes. After hybridization, the slides for 30 minutes and washed in 0 ix SSC: 20% formanida, 10 mid DFF at SOC overright. Automating capity was done by exposing the pitters to Kodak NTB-2 cintilion for 1-2 weeks at all 2 and by developing with

whole-mount in the hybridization of 955 day contract the performed sy described by Sasaki and Hogan (1993) for the Hoganises, by hybridization signal with their was not laint to detect, so a procedure hybridization signal with their state of the second so a procedure

detection. A digoxigenin labeled uridine for RNA probe was purchased from Boehringer Mannheim.

Genomic and cDNA clones of Dax gene family

of the homeoboxes

[(A/O)TCAAG(A/G)TCTGGTT(C/T)CA(A/G)AA(C/T)(C/A)GG(C/A)G(C/G)ATG

AAGGA] under mild stringency. From 8x10⁵ plagues, three positive closes, g161, g164 and g169, were obtained. Restriction analysis and nucleotide sequence determination indicated that g161 and g169 corresponded to HarB1 (Frotunan et al., 1990) and KarB3 (Lond et al., 1987), respectively, while g164 represented a novel gene (Figs. 1A and

mouse carbryonic cDNA library using a fragment of g164 as probe. The cDNA clone obtained, c164, was 1795 by long, contained a poly A tall which was accompanied by a polyadenylation afgoral (AATAAA) (Figs. 1A and 2A). The longest open reading frame (ORF) found in the c164 sequence began at the 5' end and learningted at nucleotide position 230 Comparison of the c164 and g164 sequences indicated that a putotive translational initiation site conforming to Kozak's consensus (Kozak, 1986) existed 171 by upstream of the 5' end of c164 (Fig. 2A). Translational initiation at this site is consistent with the size of the transcript estimated from Northern analysis (2 kb, data not shown).

gener exist in mouse and chicken genomes; I used a fregment of cities which included the homeobox is in Fig. 1) in Southern hybridization analysis. In addition to the predicted gift fregments, several different fragments of mouse DNA hybridized to the probe (data not shown)

Results

Genomic and cDNA clones of Dbx gene family

A λ EMBL3 mouse genomic library was screened by hybridization with synthetic oligonucleotides representing the most conserved region of the homeoboxes

[(A/G)TCAAG(A/G)TCTGGTT(C/T)CA(A/G)AA(C/T)(C/A)GG(C/A)G(C/G)ATG AAGGA] under mild stringency. From 8x10⁵ plaques, three positive clones, g161, g164 and g169, were obtained. Restriction analysis and nucleotide sequence determination indicated that g161 and g169 corresponded to *Hox*B1 (Frohman et al., 1990) and *Hox*D3 (Lonai et al., 1987), respectively, while g164 represented a novel gene (Figs. 1A and 2A).

To isolate the corresponding cDNA, I screened a λgt10 12.5 day mouse embryonic cDNA library using a fragment of g164 as probe. The cDNA clone obtained, c164, was 1795 bp long, contained a poly A tail which was accompanied by a polyadenylation signal (AATAAA) (Figs. 1A and 2A). The longest open reading frame (ORF) found in the c164 sequence began at the 5' end and terminated at nucleotide position 830. Comparison of the c164 and g164 sequences indicated that a putative translational initiation site conforming to Kozak's consensus (Kozak, 1986) existed 171 bp upstream of the 5' end of c164 (Fig. 2A). Translational initiation at this site is consistent with the size of the transcript estimated from Northern analysis (2 kb, data not shown).

To determine whether homeobox sequences related to the g164 gene exist in mouse and chicken genomes, I used a fragment of c164 which included the homeobox (b in Fig. 1) in Southern hybridization analysis. In addition to the predicted g164 fragments, several different fragments of mouse DNA hybridized to the probe (data not shown).

Companie and exists visual about the passes by total an animalian and alternation and alternation and alternation and animalian and animalian anim

of the homeohouse

Charles of the last was separated and gives seen addition of the seen of the s

And all respectively, while with represent a new particles, the sea

and A wheel a home-way a work of the state o

And in the party bearing and an analysis of the party of

acquerace began as the transferred as accounted as accounting parties will

The still - butto by June of Bell by superstant and Till Benezie Court.

Compared and the contract of t

lety in annually a deep I company neglects but a seen it when every notation and asked and

downship too with come are an excitation of the same and allegians

Multiple hybridizing bands were also obtained with chicken DNA. These results suggested that g164 and related genes form a gene family.

Further screening of the λ EMBL3 mouse genomic library with a c164 fragment as probe, resulted in two positive clones. One of them corresponded to Hlx (Allen et al., 1991), while the other (clone gS4) represented a novel gene (Figs. 1B and 2B).

I screened the 12.5 day mouse embryonic cDNA library using a restriction fragment of gS4 as probe. One clone, cS4 consisted of a 1035 bp insert with a 703 bp ORF which was missing an ATG at the 5' end but contained a poly A tail at the 3' end (Figs. 1B and 2B).

The nucleotide sequences and exon-intron boundaries of the genes represented by g164 and gS4 were reconstructed with the aid of their cDNA sequences, and are shown in Fig. 2. After these sequences were determined, Lu et al. (1992 and 1994) reported a new gene Dbx (developing brain homeobox gene) which was identical to the gene for g164/c164. There were several sequence disagreements in the 5' and 3' noncoding regions between the cDNA sequence reported by Lu et al. (1994) and g164 (Fig. 2A, nucleotides 799, 3027, 3096-8, 3409, 3415, 3517, 3565-6), which may reflect polymorphism or sequence error. gS4 which was entirely new in the mouse, I named Dbx2.

The amino acid sequence conservation between the homeodomains of Dbx and Dbx2 is 85% (Fig. 3). In addition, a significant similarity can be found in the 15 amino acids located C-terminal to the homeodomains. Comparison of the genomic and cDNA clones of Dbx indicated that the gene consists of at least 4 exons. The structure of Dbx2 has not been completely resolved but the position of the intron/exon junctions in the homeobox was identical to that of Dbx

which making the boundaries and once there are adjusted adjusted with the property of the party and the party and

Purchas superior of the Annual States of the States of the

contraponded in Mir (Afred at of, 1962), while on when fifteen past

Part a see instrument like reside and a mirror on hig to interment maintained

but contained a poly A salt or the 2' and Pape 4D and Sol

transmitted by all the same state or are at an area and area ar

determined, its of all cross and level has deep to all dominated to the continued of the co

the second well districted to the second to the second purpose pullbrance of

1817. 3565-0), which was reflect polymers of collection (\$601)

The matter are less to the latter of the lat

Additional allocations and in management of the present of the anisotropic management of the property of the p

closes of First indicated that the property products of the popular of

the introduction parameters of the horizontal was annihilated and

(Figs. 1 and 2). The structural similarity and expression specificity to be described below indicated that Dbx and Dbx2 compose a gene family.

Comparison of the amino acid sequences of Dbx and Dbx2 homeodomains with those of other proteins revealed that the zebrafish hlx-I (Fjose et al., 1994) and the chicken CHoxE (Rangini et al., 1991) appeared to be homologues of Dbx and Dbx2, respectively. hlx-I shared a 97% similarity with Dbx and CHoxE a 93% similarity with Dbx2 in the homeodomain (Fig. 3). I have isolated a chicken clone encoding a homeobox sequence that is more homologous to Dbx than Dbx2 or CHoxE (data not shown), suggesting that the Dbx gene family is conserved evolutionarily in both mouse and chicken genomes.

Chromosomal location of Dbx2

The mouse chromosomal location of Dbx2 was determined by intersubspecific backcross analysis. The 3' intron of the gene contained a $(CA)_n$ repeat (Fig. 2B) which in many cases gives rise to genetic variation. I synthesized two oligonucleotide primers,

5' CACCCTGCACATAAACATGTG 3', 5' CTAGTAGAGCACCTATGTGTG 3', that spanned a 164 bp region containing the CA repeat. DNA isolated from C57BL/6 and the MSM strain originating from a Japanese wild mouse, $\underline{Mus\ mus\ molossinus}$, were subjected to polymerase chain reaction (PCR). Analysis of the products by electrophoresis on a polyacriamide gel exhibited a simple sequence length variation between them. A segregation analysis was then carried out for 131 intersubspecific backcross mice obtained by mating (C57BL/6 × MSM) F₁ to MSM males. The strain distribution pattern was compared to that of microsatellite loci that had been determined by published markers (Dietrich et al., 1994). Fig. 4 summarizes the results of typing backcross mice. A clear linkage of the Dbx2 locus was found with two markers, D15Mit15

(*Hox*C8) and D15Mit40. From this haplotype analysis, I concluded that the most likely order of loci on chromosome 15 was cen. . .D15Mit15-0.8 cM-D15Mit40-6.1 cM-*Dbx2*. . .telo. Since *Dbx* has been mapped to mouse chromosome 7 (Johnson et al., 1992), *Dbx* and *Dbx2* do not form a gene cluster.

Expression of Dbx and Dbx2 in the embryonic CNS

The spatio-temporal distribution of Dbx and Dbx2 mRNAs was examined by in situ hybridization in embryos from stages older than 6.5 day using Dbx or Dbx2 antisense probes (d and e in Fig. 1). Expression of both genes was detected in the neural tube of embyros of stages later than 9.5 day as shown in embryos hybridized in whole mount (Fig. 5). Remarkably, both Dbx and Dbx2 were expressed in a zone separating alar and basal plates along the length of the spinal cord and rhombencephalon. Expression of Dbx2 occurred at a lower level and in a broader band than Dbx (Fig. 5). Dbx expression was also observed in the brain, most conspicuously in the mammillary areas, dosal thalamus to pretectum, and midbrain.

The regional specificity of Dbx/Dbx2 expression in the brain was more closely examined in 10.5-15.5 day embryos by in situ hybridization of serial sections through saggital, transverse and frontal planes. Representive examples of in situ hybridization data are shown in Figs. 6-8 and summarized as a scheme in Fig. 12. The pattern of expression of these genes did not substantially change during this period. The expression of Dbx and Dbx2 in the brain was restricted to the ventricular zone exept for that of Dbx in the septum of the telencephalon to be described below.

(1) Expression in the telencephalon

In the telencephalon, Dbx expression could be detected in the caudal ganglionic eminence (Fig. 7E) and septum (Fig. 6A), while Dbx2 was not detectable. Dbx epression in the septum was detected in cells out of the ventricular zone.

(2) Expression in the diencephalon

In the diencephalon, Dbx was expressed mainly in three domains rostral to the dorsal thalamus. One domain covered the anterior entopeduncular area (Fig. 6C, E) and the second domain covered the eminentia thalami (Fig. 7E). The third domain covered the optic chiasma, basal zone of the hypothalamus, mammillary areas (including the retromammillary area) and zona limitans intrathalamica (the boundary separating the dorsal and ventral thalamus) (Figs. 6A, C, E and 7A). Out of these domains, Dbx-positive cells scattered among negative cells in the alar zone of the hypothalamus and caudal region of the anterior preoptic area (Figs. 6C, 7A and 12A). In the same diencephalic subdivision Dbx2 was expressed in a short band located at the alar-basal boundary, from caudal region of hypothalamus to rostral region of the ventral thalamus (arrows in Figs. 6D, 7B, F and 8B, D-F).

Caudal to the zona limitans intrathalamica, Dbx expression in the alar zone started near the rostral boundary of the dorsal thalamus and epithalamus, and was extended through the middle of the pretectum (Fig. 6C). In the caudal half of the pretectum, Dbx expression was confined to the boundary between alar and basal zones (Figs. 6C). Dbx2 was expressed in a continuous band along the alar-basal boundary spanning the dorsal thalamus and the pretectum (Figs. 6D, 7B and 8A, C, E, F). It was noted that expression of both Dbx and Dbx2 increased in the dorsal thalamus away from the zona limitans intrathalamica.

(3) Expression in the mesencephalon

In the mesencephalon, Dbx was strongly expressed in the dorso-lateral wall, continuously from the pretectum and to the rhombencephalic isthmus (Fig. 6A, C, E). Very weak expression of Dbx2 could also been seen throughout the dorso-lateral wall overlapping with that of Dbx (Figs. 6B, D and 10A). In addition, Dbx2 was expressed at the boundary between the dorso-lateral wall and the basal zone in its rostral half (arrowhead in Fig. 6D).

In the rhombencephalic isthmus, Dbx was expressed at a high level while Dbx2 was not detected (Fig. 6C, D).

(4) Expression in the rhombencephalon and spinal cord

Along the length of the neural tube posterior to the rhombencephalic isthmus, the expression of Dbx and Dbx2 was detected as a bilateral pair of narrow bands at the boundary between the basal and alar plates (Figs. 5, 6, 7C-F, 8C, 9).

In the spinal cord, expression of Dbx was restricted to the boundary between the alar and basal plates (Fig. 5A and 9C, D). Expression of Dbx2 was similar to Dbx in its distribution, but broader, extending out from both sides of the Dbx-expressing band and with less defined borders (Fig. 5B and 9E, F). In the rhombencephalon, expression of both genes was restricted to the boundary of the dorso-ventral zones in a manner similar to that seen in the spinal cord (Fig. 5, 7C-F, and 8C).

At early stages when lateral plates of the spinal cord still consisted of neuroepithelial germinal cells (9.5-10.5 day), expression of Dbx and Dbx2 can be detected throughout the entire thickness of the plates (Fig. 9C, E). At later stages when neural differentiation initiates in the intermediate zone, expression of both genes become confined to the ventricular zone (Fig. 9D, F). But regardless of the developmental stage, the dorso-ventral width of the Dbx/Dbx2 expressing bands

remained fairly the same. Expression of Dbx2 appears to increase with the development of the embryo as judged by the increase in intensity of the hybridization signal (Fig. 9E, F).

Expression of Dbx2 outside of the CNS

Although expression of Dbx was confined to the CNS, Dbx2 was also expressed in mesenchymal components of the limb buds and tooth germs.

(1) Expression in the limb buds

In the limb bud mesenchyme, a significant level of Dbx2 expression occured after 11.5 day (Fig. 10A-C). High expression appears localized to the internal mesenchymal cells which later participate in chondrogenesis. After 13.5 day the expression was localized to the cells surrounding the precartilaginous condensation of the digits (Fig. 10D, E).

(2) Expression in the tooth germs

In the tooth germs, Dbx2 was expressed in the mesenchymal cells surrounding the enamel organ. Data of 13.5 and 15.5 day embryos are shown in Fig. 11. The Dbx2-positive cells included both dental papilla and follicle cells, but not the cells immediately adjacent to the internal enamel epithelia of the enamel organ and the cells surrounding the dental ledge. The expression pattern did not substantially change during this period.

Discussion

Dbx gene family

I have described the gene organization and expression of two new mouse homeobox genes which compose the Dbx family. Dbx and Dbx2 have similar homeodomain sequences and exon-intron organizations. Dbx cDNA was cloned independently by Lu et al. (1992) and its gene organization reported (Lu et al. 1994), which are in general agreement with my results.

The homeodomain and C-proximal flanking amino acid sequences are very similar between Dbx and Dbx2, but the sequences of other regions are quite divergent. Dbx and Dbx2 likely can recognize analogous nucleotide sequences, but may differ in their action as transcriptional regulators. Proximal to the C terminus of Dbx protein, there is a cluster of acidic residues often present in transcriptional activators (Sigler, 1988), but this motif is absent in Dbx2.

Comparison of the homeodomain amino acid sequences indicated that the Dbx family is related, but apparently distinct from the H2.0 family (Fig. 3). 63-68% similarity was found between the Dbx homeodomain and the homeodomains of the H2.0 family: H2.0 (Barad et al., 1988), H1x (Allen et al., 1991) and AHox1 (Saiga et al., 1991). Inverterate members for the Dbx family have not yet been reported.

Expression of Dbx and Dbx2 in the embryonic brain

Dbx and Dbx2 are expressed in the embryonic brain with regional specificities as summarized in Fig. 12A, B. Expression of Dbx is mapped in various regions of the diencephalon with boundaries largely consistent with the neuromeric model of the embryonic brain of the mouse proposed by Bulfone et al. (1993), by Puelles and Rubenstein

malangradid.

Tailed from a deligated and a supplied to the state of th

went of the case of the set of climber of the set of the second of the set of the second of the seco

with my viriality.

The furniture and a special theories will exposure.

and wanty election but mad (1902), but the organization of order and allege and allege was encountered.

The state of the second of the state of the

netwater 15gler, 1968, but her ment in nicht in Ding.

that the Day taken is settled, but applicably interest from the MC P

to be state and the property of the state of the state of the state of the state of

averterant mannings for the 11th States and values and resident

attent almost one and the same and the tentagement and the molecularity

designed at a set the residence in the last at the state of the set of the se

the warrous which the description of the same state of the same state of the

modern proposed by Dallins at al 11991, as Francisco and Polemnica

(1993) and by Rubenstein et al. (1994). Dbx expression in the brain had also been reported by Lu et al. (1992 and 1994), but not in great detail. Dbx2 expression in the diencephalon is limited to the alar-basal boundaries of the caudal hypothalamus to the region rostral to the ventral thalamus, dorsal thalamus and pretectum. In the midbrain, expression of Dbx and Dbx2 largely overlaps along the dorso-lateral wall.

It should be noted that expression of the Dbx genes emphasizes the separation of the alar and basal compartments. It is also remarkable that Dbx expression in the alar compartment of the pretectum continuous from the dorsal thalamus terminates in the middle of the pretectum with clear demarcation. This indicates that pretectum may be divided into rostral and caudal components as proposed by Figdor and Stern (1993). Thus, the Dbx genes provide two important genetic landmarks to the generic map of the brain.

Expression in the spinal cord

Along the length of the neural tube posterior to the rhombencephalic isthmus, both Dbx and Dbx2 are expressed at the boundary of the alar and basal plates which later develop to sensory and motor columns, respectively (Fig. 12C, D). It is important to note that these genes are expressed only in mitotically active cells of the ventricular zone in these regions and that these genes encode putative transcriptional regulators. It is possible that the Dbx family plays a role in specifying a small population of neural cells at the boundary between the columns, a region classically called the sulcus limitans. Expression of these genes overlaps, but the expression of Dbx2 is broader in a dorso-ventral direction. Therefore, the cells expressing both Dbx and Dbx2 are flanked by cells expressing only Dbx2. The overlapping

expression of the genes may indicate functional redundancy, but I prefer the view that the products of these genes have different transcriptional regulational activities, as suggested by the high divergence outside of the homeodomain.

It is interesting to speculate a possible function for the sulcus limitans cells in relation to expression of Dbx genes. It is known that the differentiation of motor neurons depends on inducing signals from the notochord and floor plate (Placzek et al., 1991; Yamada et al., 1991). Cells expressing Dbx genes may aquire a capacity to halt the signal from the ventral region which otherwise induces motor neurons throughout the spinal cord. Altman and Bayer (1984) have proposed that the region of the sulcus limitans (called the intermediate plate) is a source of contra- and ipsilaterally projecting relay neurons. The Dbx family may contribute to the determination of cell fates of these neurons. In either case, it is interesting to note that a certain class of cadherins is expressed in ventricular zone cells of the sulcus limitans similar to the Dbx family (M. Takeichi, personal communication). It is an intriguing possibility that the Dbx family of genes may regulate cadherin expression, which modifies cell interactions and consequently alters fate of the neuroepithelial cells.

Expression of homologues in other animal species

Zebrafish hlx-1, a putative homologue of Dbx, shows an expression pattern similar to Dbx in the CNS (Fjose et al., 1994). However, prior to expression in the CNS, hlx-1 is expressed in an area of primitive mesoderm (Fjose et al., 1994) which was not observed in the mouse embryo.

Chicken CHoxE, the putative chicken homologue of Dbx2, has been reported to be expressed in the developing neural tube posterior to the

rhombencephalic isthmus almost identically to Dbx2. But there is no description of more anterior expression or expression in the other organs (Rangini et al., 1991).

Complementary expression teritories with other homeobox genes

The proposed neuromere map of the CNS (Bulfone et al., 1993; Puelles and Rubenstein, 1993; Rubenstein et al., 1994) is based on the expression patterns of various homeobox genes. Examination of the expression of such genes indicated that some are expressed in regions of brain complementary to Dbx/Dbx2 expression. For instance, Pax-6 is expressed roughly complementary to Dbx in the diencephalon (Walther and Gruss, 1991; Puelles and Rubenstein, 1993). Caudal to rhombomere 6 of the rhombencephalon and spinal cord, Gbx-2 is expressed both in alar and basal plates, but not in the sulcus limitans (Bulfone et al., 1993), showing a complementarity to Dbx family expression. These observations may suggest that characteristics of individual neuromeres are defined by multiple superpositions of dichotomous choice of gene expression. It is interesting to see if complementary pairs of transcription factor genes are generally found with boundaries which conform to the model of the CNS.

Expression in the limb buds and tooth germs

Both in the limb buds and tooth germs, Dbx2 is expressed in undifferentiated mesenchymal cells. In the limb buds, Dbx2 seems to be expressed in the cells prior to paticipate in the precartilaginous condensation. In the tooth germs, Dbx2 is expressed in the mesenchymal cells surrounding the enamel organ, but not in the cells immediately adjacent to the internal enamel epithelia which may be

chombencephalic isthmas aimost identically in 1962. But their is not description of more actarios expression or camersion in the other organs (Rangini et al., 1991).

Complementary expression teritories with other nomentary genes.

The proposed neuromera map of the CNS thubbone of all 1993, problems and Rubensuela, 1993; Rubension of all 1993, is tased on the expression generally of various homeobox games. Hastminutes of the expression of such games indicated that some are expressed in regions of brain complementary to the that some are expressed in regions of brain complementary to the thought expression. For instance, Purch is and Gress, 1991; Puelles and Rubenstein, 1993; Candal at thembomeric of the thombomerically combenerability and Rubenstein, 1993; Candal at thembomeric of the thombomerically and basal plates, but and spinal cond, Ghz-2 is expressed both in also observations may suggest that characteristics of individual neuromeres observations may suggest that characteristics of individual neuromeres observations may suggest that characteristics of individual neuromeres.

Expression in the limb bads and toots germs. Dive? is expressed in undifferentiated mescapitymal cells. In the limb bads, 192.2 seems to be expressed in the cells prior to principate in the processification. In the tooth years, 192.2 is expressed in the most cold germs. Dive? its expressed in the most cold germs. Dive? its expressed in the mescapitymal cells surrounding the cuamel organ, but not in the cells formation in the cells.

undergoing differentiation into odontoblasts. Therefore, in the mesenchymal cells of these organs, Dbx2 seems to be expressed transiently immediately before the start of differentiation, which may suggest that Dbx2 is playing a role in determination of these cells. This is also consistent with the observation in the CNS that Dbx2 is exclusively expressed in the undifferentiated cells of the ventricular zone.

Professor Yusuke kamachi, Dr. Yoshio Wakamatsu, Dr. Daihachirch Yoshousane, Dr. Naoto Hayanaka and all the members of Kondoh's Inhoratory for their helpful advice, discussions and supports,

Richardsonal mapping of Dhr?

Melecular Biology, Nagoya University) for his guidance in neuroanatomy of the embryon, Professor Masatoshi Takeichi (Department of Biological Science and Technology, University of Tokushims) for advice in whole-mount in single-bybridization and Dr. Ruth Yu (Graduated school of Bolosciences, National of Science and Technology) for critical reading

Kanne Ohiski (Graduated school of Boioscionees, Nava Institute of Science and Technology) and Dr. Mitanto Oyanagi (First department of Biochemistry, Nicasa University School of Medicine) for their technical

Finally, I second like to thank Professor Hiromu Sugino and Assistant Professor Takamori Nakamura for their supports and giving me the time to finish this work here in Justitute for Enzyme Research, University of Tokushims.

Acknowledgements

I am indebted to Professor Naoki Takahashi (Graduated school of Boiosciences, Nara Institute of Science and Technology) and Professor Hisato Kondoh (Institute for Molecular and Cellular Biology, Osaka University) for their valuable guidance and innumerable helpful advice throughout my work.

I am grateful to Associated Professor Yujiroh Higashi, Assistant Professor Yusuke kamachi, Dr. Yoshio Wakamatsu, Dr. Daihachiroh Tomotsune, Dr. Naoto Hayasaka and all the members of Kondoh's laboratory for their helpful advice, discussions and supports.

I am also grateful to Professor Ryo Kominami (First department of Biochemistry, Niigata University School of Medicine) for his support in the chromosomal mapping of Dbx2.

I would like to thank Professor Hajime Fujisawa (Department of Molecular Biology, Nagoya University) for his guidance in neuroanatomy of the embryos, Professor Masatoshi Takeichi (Department of Biophysics, Kyoto University) for information of cadherin expression, Professor Sumihare Noji (Department of Biological Science and Technology, University of Tokushima) for advice in whole-mount *in situ* hybridization and Dr. Ruth Yu (Graduated school of Boiosciences, Nara Institute of Science and Technology) for critical reading.

I thank Dr. Toshiroh Ito (National Institute for Basic Biology), Ms. Kanae Ohsaki (Graduated school of Boiosciences, Nara Institute of Science and Technology) and Dr. Mitsuru Oyanagi (First department of Biochemistry, Niigata University School of Medicine) for their technical supports.

Finally, I would like to thank Professor Hiromu Sugino and Assistant Professor Takanori Nakamura for their supports and giving me the time to finish this work here in Institute for Enzyme Research, University of Tokushima.

References

- Akam, M. (1987) Development 101, 1-22.
- Allen, J. D., Lints, T., Jenkins, N. A., Copeland, N. G., Strasser, A., Harvey, R. P. and Adams, J. M. (1991) Genes Dev. 5, 509-520.
- Altman, J. and Bayer, S. A. (1984) In Beck, F., Hild, W., van Limborgh, J., Ortmann, R., Pauly, J. E., Schiebler, T. H. (eds), Advances in Anatomy, Embryology and Cell Biology vol 85, Springer Verlag.
- Barad, M., Jack, T., Chadwick, R. and McGinnis, W. (1988) EMBO J. 7, 2151-2161.
- Bastian, H. and Gruss, P. (1990) EMBO J. 9, 1839-1852.
- Bulfone, A., Puelles, L., Porteus, M. H., Frohman, M. A., Martin, G. R. and Rubenstein, J. A. R. (1993) J Neurosci. 13, 3155-3172.
- Dietrich, W. F., Miller, J. C., Steen, R. G., Merchant, M., Damron, D., Nahf, R., Gross, A., Joyce D. C., Wessel, M., Dredge, R. D., Marquis, A., Stein, L. D., Goodman, N., Page, D. C., and Lander, E. S. (1994) Nature Genetics 7, 220-225.
- Figdor, M. C. and Stern, C. D. (1993) Nature 363, 630-634.
- Fjose, A., Izpisúa-Belmonte, J. C., Fromental-Ramain, C. and Duboule, D. (1994) Development 120, 71-81.
- Frohman, M., Boyle, M. and Martin, G. R. (1990) Development 110, 589-607.
- Gehring, W. G., Qian, Y. Q., Billeter, M., Furukubo-Tokunaga, K., Schier, A. F., Resendez-Perez, D., Affolter, M., Otting, G. and Wüthrich, K. (1994) Cell 78, 211-223.
- Gruss, P. and Walther, C. (1992) Cell 69, 719-722.
- Hayashi, S., and Scott, M. P. (1990) Cell 63, 883-894.
- Johnson, K. R., Lu, S., Murtha, M. T., Ruddle, F. H. and Davisson, M. T. (1992) GENOMICS 14, 1107-1109.

Kappen, C., Schughary, K. and Ruddle, F. H. (1993) Genomics 18, 54 -70.

Kenyon, C. (1994) Cell 78, 175-180.

Kessel, M. and Gruss, P. (1990) Science 249, 374-379.

Kobayashi, S., Saito, H. and Okada, M. (1994) Develop. Growth & Differ. 36, 629-632.

Kozak, M. (1987) Nucl. Acids Res. 15, 8125-8146.

Krumlauf, R. (1994) Cell 78, 191-201.

Levine, M. and Hoey, T. (1988) Cell 55, 537-540.

Lonai, P., Arman, E., Czosnek, H., Ruddle, F. H. and Blatt, C. (1987) DNA 5, 409-418.

Lu, S., Bogarad, L. D., Murtha, M. T. and Ruddle, F. H. (1992) Proc. Natl. Acad. Sci. USA 89, 8053-8057.

Lu, S., Wise, T. L. and Ruddle, F. H. (1994) Mech. Dev. 47, 187-195.

McGinnis, W., Levine, M.S., Hafen, E., Kuroiwa, A. and Gehring, W. J. (1984) Nature 308, 428-433.

McGinnis, W. and Krumlauf, R. (1992) Cell 68, 283-302.

Placzek, M., Yamada, T., Tessier-Lavingne, M., Jessell, T. M. and Dodd, J. (1991) Development Supplment 2, 105-122.

Puelles, L. and Rubenstein, L. R. (1993) TINS 16, 472-479.

Rangini, Z., Ben-Yehuda, A., Shapira, E., Gruenbaum, Y. and Fainsod, A. (1991) Mech. Dev. 35, 13-24.

Rubenstein, J. L. R., Martinez, S., Shimamura, K. and Puelles, L. (1994) Science 266, 578-580.

Saiga, H., Mizokami, A., Makabe, K. W., Satoh, N. and Mita, T. (1991)

Development 111, 821-828.

Sanger, F., Nicklen, S. and Coulson, A. R. (1977) Proc. Natl. Acad. Sci. USA 74, 5463-5467.

Sasaki, H. and Hogan, B. L. M. (1993) Development 118, 47-59.

Appen, C., Schughmy, S., and Ruddle, F. H. (1993) Genomics 18, 54-70. Senyon, C. (1994) Cell 78, 175-180.

Kessel, M. and Gross, F. (1990) Science 249, 314-379, Kobayashi, S., Salto, H. and Okada, M. (1994) Develop, Growth &

Kozak, M. (1987) Nucl. Acids Res. 15, 8125-8146 Krumlauf, R. (1994) Cell 78, 191-201

Long, P., Arman, E., Crownell, H., Ruddle, E. H. and Blatt, C. (1987) DNA S.

Luc. S., Bogard, L. D., Mardee, M. T. and Ruddle, E. H. (1997) Proc. Nat. Acad. Sci. USA 87, 8053-8057.

McGinnis, W., Levine, M.S., Halan, E., Kunolwa, A. and Gebring, W. I.

McCinnia, W. and Krambarl, R. (1992) Call 68, 283-302.
Placack, M., Yamada, T., Texalor-Lavingna, M., Jassell, T. M. and Dodd, J.

Puelles, L. and Rubonstein, L. R. (1993) TIME 16, 472-479, Rangini, Z., Ben-Yetoda, A., Shapira, E., Groenburm, Y. and Frimsod, A.

Rubenstein, J. L. R., Martinux, S., Shmannur, K. and Poetics, L.

Saign, H., Mirokami, A., Makabe, E. W., Saigh, N. and Mile, T. (1991).

Sanger, F. Mickley, S. and Coulson, A. R. (1973) Proc. Natl Sci.

Sanki, H. and Hogen, B. L. M. (1993) Desciopment 118, 47-39.

Scott, M. P. and Weiner, A. (1984) Proc. Natl. Acad. Sci. U.S.A 81, 4115-4119.

Scott, M. P., Tamkun, J. W. and Hartzell, III, G. W. (1989) Biochem. Biophys. Acta. 989, 25-48.

Sigler, P. B. (1988) Nature 333, 210-212.

Suda, T., Oyanagi, M., Wakana, S., Takahashi, Y., Kanada, H., Yonekawa, H., Miyashita, N., Shiroishi, T., Moriwaki, K. and Kominami, R., (1994) DNA Research 1, 169-174.

Wakamatsu, Y. and Kondoh, H. (1990) Acta Histochem. Cytochem. 23, 367-374.

Walther, C. and Gruss, P. (1991) Development 113, 1435-1449.

Yamada, T., Placzek, M., Tanaka, H., Dodd, J. and Jessell, T. M. (1991) Cell 64, 635-647. Scott, M. P. and Weiner, A. (1984) Proc. Natl. Acad. Sci. U.S.A. 81,

Scott, M. P., Tamkun, J. W. and Hartzell, III, G. W. (1989) Biochem. Binobys, Acta, 989, 25-48.

Sigler, P. B. (1988) Nature 333, 210-212

Suda, T., Oyanagi, M., Wakana, S., Takahashi, Y., Kanada, H., Yonekawa, H.,

DNA Research 1, 169-174

Wakamatso, Y and Kondob, H. (1990) Acta Histochem. Cytochem. 23.

Watcher, C. and Grass, P. (1991) Development 113, 1435-1449.
Yamada, T., Placzek, M., Tanata, H., Godd, J. and Jonedi, T. M. (1991)

Figures

Fig. 1. Structure of Dbx (A) and Dbx2 (B) genes. Exon-intron organization maps of genomic clones of Dbx (g164) and Dbx2 (gS4), in comparison with cDNA clones shown below (c164, cS4). Wide boxes indicate the protein coding sequences, open boxes the homeoboxes, and narrow boxes the 3' untranslated regions. a to f indicate probes used for phage library screenig (a, c and f), genomic Southern hybridization (b) and in situ hybridization (d and e). BB in the fourth exon of Dbx indicates twice repeated BamHI sites separated by a nucleotide between their recognition sequences. Restriction sites, B, BamHI; E, EcoRI; H, HindIII.

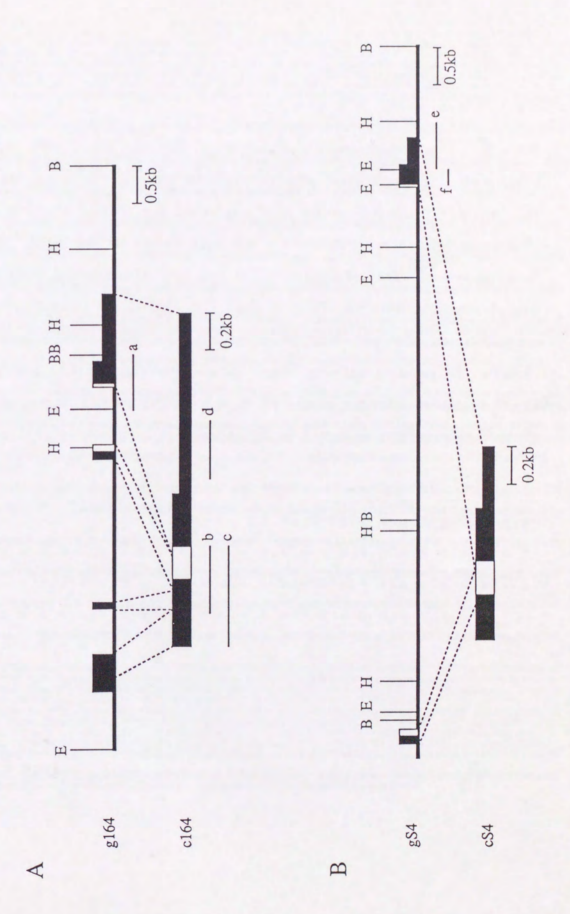


Fig. 2. Nucleotide sequences of Dbx (A) and Dbx2 (B) genes. Nucleotides are numbered arbitrary just to indicate the positions in the Figure. Predicted amino acid sequences are indicated below the nucleotide sequences with residue numbers shown to the right. The homeodomains are shaded. The (CA)n repeat, the regions complementary to the PCR primer set used for chromosomal mapping and the polyadenylation signal are underlined. Arrows indicate the putative splicing sites. Solid arrowhead in (A) indicates the 5' end of c164. Sequences of 5' to [in (B) are cDNA sequence. Open triangles indicate the poly A addition sites.

TTGTTGAACTTCATTGTCAGAGCTGTCAGTTTTCAAAGAGCTGGAGCGGGCGG	180 270 360 450 540 630 720 810
CCTGCGACCCACGCCCACCTTAACACTGCCCCAGTCCCTTCAATCCGCATTTTCCGGCCACTCTAGTTTCCTAGTAGAAGATTTGATCC L R P T P T L T L P Q S L Q S A F S G H S S F L V E D L I R	990 46
CATCAGCCGGCCTCCCACCTACCTGTCCCGCAGCATACCCCGCTGCCAGCCTGTCACCCCCCAGTCAAGAGGCCCCTGCAGCCCTCGCCGISRPPTYLSRSIPAASLSPPSQEAPAALAD	1180 76
CTCGGGGACCTCAGACCTGGGCTCTCCTGGGTCCGGGAGCCGGCGAGCCGAGCCAACAGACCGCCCTCTCCCCTGCCAGCGAGCCCA S G T S D L G S P G S G S R R G S S P Q T A L S P A S E P T	1270 106
GTTTCTGAAGTTTGGGGTGAATGCCATCCTTTCCTCGGCGCCTAGAAGAGGGTAAGATACAGCCGTGCCCTCGACGTGCGTTCCGCGG (F L K F G V N A I L S S A P R R E	1358 123
	1440 137
GCCTTTCCATACTTCGAAGGCTCGTTCCAACCCTTTATCAGATCCTCCTATTTCCCAGGTAGGT	1530 157
TGGACAGCCCGTTCCAGTCCCGGCCGGTCGCTCCCTACCCCGAAGTCGTATTTCTCCGTCCACCCC(-1.5kb-) AAAGGCAATTGT GCTGCTGTTTCGTGGGTTTTTCCTAATCCGGGTTTGCACGGTTGAAAGCGAAAGTTATTTGTGCGGAAAGCGCCTTTCACTCAGACAGG TGTAGGTTCTCCGTCTATTCTCTACGGGGAGAAAAAGGAGTGTTTTAAAGAGATGAAGGGACTCGAGGGGGAAAAAGGAGGGAG	1700
AGTGAAAACAGCTTGGGAGAGGCGGATGCTCGGCTCTGCAGGACACCCGGCTCATTCTCTACTTCCCTCACCCCGCAGCTTCCTCCAGC S S S	1880 160
PCGTGCCTATCCCTGGGACCTTCTCCTGGCCGCTGCTGCTCGCGGGCAAGCCTCGCCGGGGCATGCTGCGCAGAGCCGTGTTCTCTGAC V P I P G T F S W P L A A R G K P R R G M L R R A V F S D	1970 190
TGCAGCGGAAAGCCCTGGAGAAGACGTTCCAGAAGCAGAAGTACATCAGCAAGCCAGACCGGAAAAAGCTGGCCTCCAAGCTTGGCTTG Q R K A L E K T F Q K Q K Y I S K P D R K K L A S K L G L	2060 220
AGGACTCGCAGGTGCGAGCCGCCTCCACCG(-0.5kb-) CAAGGCTGGTGGGGAACAACACAGAAATCACCTTGAGGGCAGCGCCTTC D S Q	2140 224
TACATGTTCCCCAAGCCTGGCTTCTAAGGGTTGCCCCAACCACCACCACCACCACCACCACCACCACCACCA	2230
	2320 238
CCAAAGAGCGCGAGCTCCTGTCTAGCGGGGGCTGCCGAGAGCCAGACCCTTCCCACAAAACTAAACCCCCATCCAGACCTTAGTGATGTA K E R E L L S S G G C R E Q T L P T K L N P H P D L S D V	2410 268
CCCAGAAGGGACCTGGGGATGAGGAGGAAGACAATCCGGGTGCCCGCCTGGCCTACCACGCGCCCGCAGACCCGCGACATTTGCTGGAA Q K G P G D E E E D N P G A R L A Y H A P A D P R H L L E	2500 298
GGCCACTGCCTGCCGGCGCACTCGAGCAGCCCGGGTAAACCGTCAGACTTCTCTGATTCTGATGAGGATGAGGAGGGCGAGGAG P L P A S P A H S S S P G K P S D F S D S D E D E E G E E	2590 328
ATGAGGAAATCACGGTGTCCTAGGAGCCCCTCCAGCGGCTCTGGGGCTGTTGCAATTGGAAACTGGAGAGGGAACGCCCACGTGGCCAC	2680 335
CTGTTCAGGATCCTGGATCCTGCCTCTTCTCACAACAGACCCACCACCTTCTCAGGGGTGGCATCACCCTTCCTGTTGTCCCAGAGAC TTCCTTGCGCCGTGCTGCCACCTCTAAACAACCCTTCCCACGGGAGCGTGGGTCTTTTCTACAAGAGCAACATTGCATCTATTTATT	2860 2950 3040 3130 3220 3310 3400 3490
CACCATGTACAGACTTTTTATATATGTATATGTATCAAATGGACAAATGCCA <u>AATAAA</u> AAGTAGAATGAGTTTGATGGCTAAAGGGC CTGTTAAGTCACAAAAACGTGATGGCTGAAAGAACCAGGATTGGTCTTGTGGGTACCCAGAAGGCAAA	3647

В																																		
			GCA																												GCT		AGCT	90 30
							TGG																								CCA			180 60
	CCC										[TG	CAG	GTO	CAA	CGG	AT	CAT	CAT	TT	гтт	GGG	STT	TC	AAG	GA	AGI	cci	AGC	AC	TCA	GAT	AGC	TTA	248 63
AG	CA	ACA	GA	GAAG	CAG	CCT	GCC	ACC	GTO	GTG	AAC	TAT	GAC	SAT	ACC	CAC	TGT	TCI	GA	GAG	TCC	CGA	GG	GCT	TC	ccc	TC:	TCI	'AA	ccc	TGT	GCC	CCTT	338
гт	GT	ATI	TC	TG	гст	CTG	CCA				ACA H																						CGA R	428 85
																																	CTC	518 115
							AAG(GGT	ACA	GTO	GAG	GAG	AA	AGG	TGC	ATO	CCT	AGI	rcc	TAC	CTT	TC	AAC	AGO	CAG	TG	TGA	AAT	TTG	TTC	608 125
GA AC GT GA	CAC ACA CAA AA	GGA TACA ATT TACA	TCO ATA CAC TT' AGA	CACACACACACACACACACACACACACACACACACACA	ACACACACACACACACACACACACACACACACACACAC	CACA AATA ATCA CAC	TGAC CCAC ACAC AATC AGTC AGGC	GGT CAC GTA GTG GAG	GGC GTC ACA TCA CTA	CAA CA ACA ACA	AGC. TCT CAC. ATA CTC. AGT	AGA GCA ACA GAG ATC TAG	CCC TGT CTC CAP CTP	CCC AAT ACT	GAA ATG CGA ATA TAG CAG	TTG CA' TG ATG AA	C (- TGC) CAA' GATO AAG' TAGO	GCA TCC GTT TGC GAG	Kb- CAC TAA CTC	CAC TTA AAA GTG	CAC ACA ATC GTA GTC GCA	AAA AAA ATT GTC	TAR CCR TAR GGG	AAC CAT AAA AAA GTT	ACC GCI TCI CTI GAT	ACA AAC AAC TTT GTC	CAC ACAC TTAC CTC	ACA ACA AAT AGC	TAC TAC TTTC GTC	AACA GGT CAA GCT TTA	TAC GCT CAA TTT GCA	TGT ACA CTA GCC GAA GTT	ACA GGA GGG	
CG	GAZ	ATA	AGO	GACT	TTC	TTT	CCT:	TAT	TAT	TTT	TAT	CAT	TAC																		CCA			1408
																															CGC			1498 173
																															AGG:			1588 203
																															AGA			1678 233
Т	TGI	ACG	GGI	AACA	ATA	CTC	CGT	STA	CAT	CT	AAA	AGA	ACC	CT	TAG	CC	AGT	AAC	ACT	TG	GAT	TA	AAC	SCC	AG	TTA	GA	TTG	TG	CCT	CAA	AAC	TGC	1768
_																																		1858 1948
							TTTC																					ATA	TGO	GAA	CAA	ATT	CAT	2038 2111

Fig. 3. Amino acid sequences of the homeodomains of *Dbx* and *Dbx2* in comparison with those of *CHoxE* (Rangini et al., 1991), *hlx-1* (Fjose et al., 1994), *Hlx* (Allen et al.,), *H2.0* (Barad et al., 1988), AHox1 (Saiga et al., 1991) and *Antp* (Scott and Weiner, 1984). Similarity score of each homeodomain sequence relative to *Dbx* is shown on the right together with the animal species from which the gene was derived. Dashes indicate the residues identical to *Dbx*.

mouse chick zebrafish mouse fly ascidian	fly
0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	38%
GMLRRAVFSDVQRKALEKTFQKQKYISKPDRKKLASKLGLKDSQVKIWFQNRRMKWRNSK -IEEMTRVSEIED	-TLEE-HFNR-LTRRR-IEI-HA-C-TER-I
Dbx Dbx2 CHoxE hlx-1 Hlx H2.0	Antp

Fig. 4. Distribution of the haplotypes for 131 progeny from intersubspecific backcross mice. The loci followed in the cross are indicated with arrows to the partial chromosome 15 linkage map. The filled squares represent the C57BL/6 allele, and the open squares represent the MSM allele. Each column represents the chromosome identified in the progeny. The numbers of the progeny carrying each type of chromosome are listed at the bottom. In the linkage map, more proximal markers mapped previously are indicated in smaller type (Dietrich et al. 1994; Suda et al. 1994).

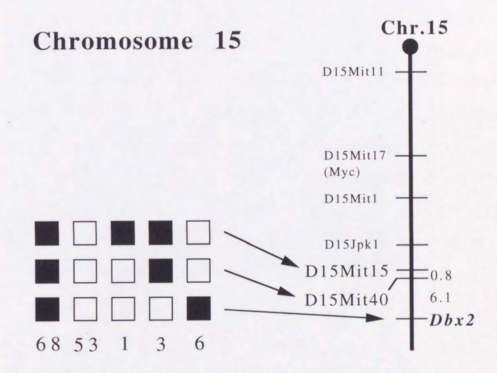


Fig. 5. Whole-mount in situ hybridization data of 9.5 day embryos probed with Dbx (A) and Dbx2 (B). The procedure for detection of Dbx and Dbx2 was slightly different as described in Experimental Procedures. The stripes of Dbx and Dbx2 expression along the length of rhombencephalon to spinal cord are indicated by arrowheads. In (A), three conspicuous sites of Dbx expression are marked: me, mesencephalon; d, a complex of areas from dorsal thalamus to rostral pretectum; ma, mammillary to retromammillary areas. In (B), deposit of the stain in structures other than the CNS was not reproducible. Scale bar, 0.5 mm.

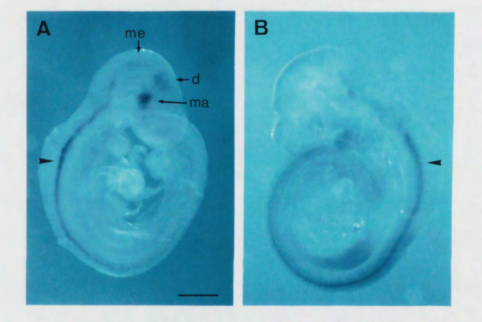


Fig. 6. Expression pattern of Dbx (A, C and E) and Dbx2 (B and D) in 11.5 day embryo in sagittal sections. In situ hybridization data are shown. (A) and (B), (C) and (D) are pairs of adjacent sagittal sections. (E) is a medial section parallel to (C). (F) shows hybridization with sense probe. An arrowhead in (D) emphasizes the hybridization signal at the boundary between the dorso-ventral wall and the basal zone of the mesencephalon which may escape from attention. An arrow in (D) indicates the expression of Dbx2 in the caudal region of the hypothalamus. Lines with labels AB, CD and EF indicate the positions and orientations of transverse and frontal sections in Fig. 7. Abbreviations: aep, anterior entopeduncular area; bh, basal zone of the hypothalamus; cpt, caudal pretectum; dt, dorsal thalamus; h, hypothalamus; is, rhombencephalic isthmus; m, mesencephalon; ma, mammillary area; oc, optic chiasma; poa, anterior preoptic area; pt, pretectum; rh, rhombencephalon; rpt, rostral pretectum; sc, spinal cord; se, septum; vt, ventral thalamus; zi, zona limitans intrathalamica. Scale bars, 1 mm.

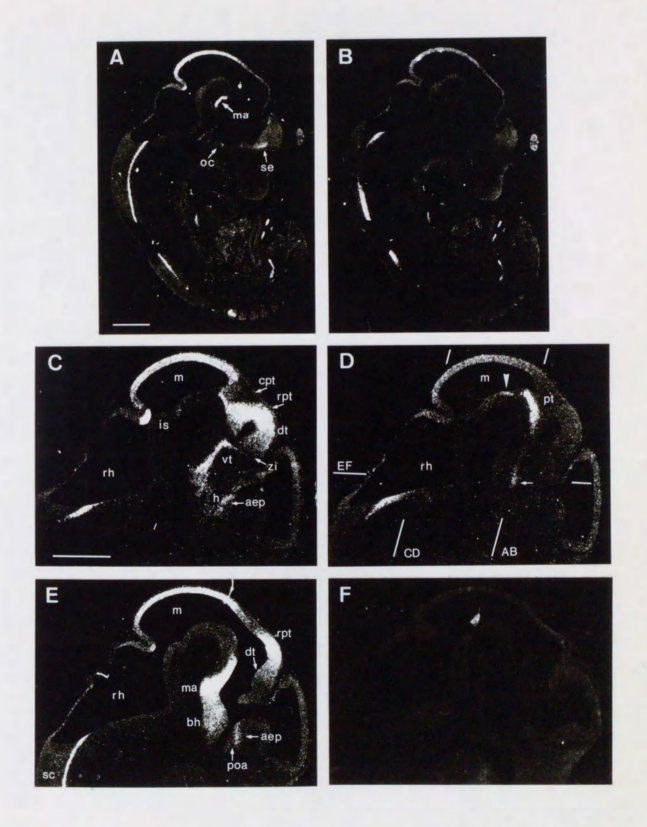


Fig. 7. Frontal (A-D) and transverse (E, F) sections showing expression of Dbx (A, C and E) and Dbx2 (B, D and F) in 11.5 day embryonic brain. (A) and (B), (C) and (D), (E) and (F) are adjacent pairs of sections. Arrows in (B) and (F) indicates the expression of Dbx2 in the caudal region of the hypothalamus. The positions and orientations of sections are indicated in Fig. 6. Abbreviations: cge, caudal ganglionic eminence; emt, eminentia thalami; otherwise the same as in Fig. 6. Scale bars, 1 mm. Scales for (E) and (F) are the same as (A).

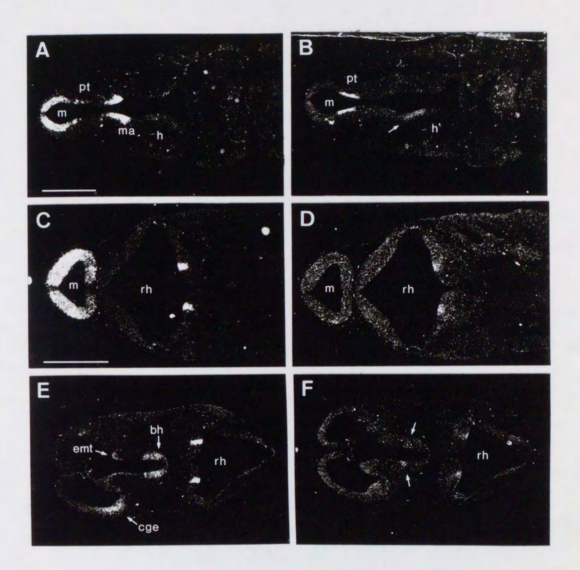


Fig. 8. Expression pattern of Dbx2 in the embryonic brain of the stages other than 11.5 day. (A) and (B) are serial frontal sections of 10.5 day embryonic brain. (C) and (D) are serial transverse sections of 12.5 day embryonic brain. (E) and (F) are frontal sections of 13.5 day and 15.5 day embryonic brains, respectively. Arrows in (B), (D), (E) and (F) indicates the expression of Dbx2 in the caudal region of the hypothalamus. Arrowheads in (E) and (F) indicates the expression of Dbx2 in the pretectal area. Signal in the telencephalon close to the bottom of the figure (D) is artifact of the section. An area of shining between the telencephalon and diencephalon in (E) is due to hemocytes rather than to a signal. Abberivations are the same as in Figs. 6 and 7. Scale bars: (A), 0.5 mm; (C) and (E), 1 mm. Scale for (F) is the same as (E).

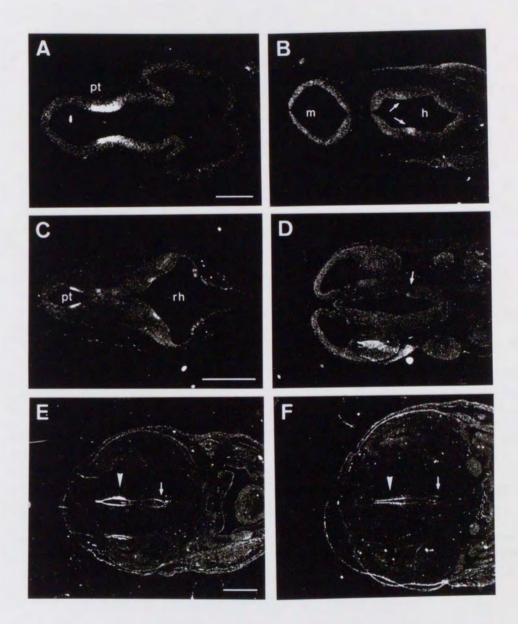


Fig. 9. Comparison of Dbx (C and D) and Dbx2 (E and F) expression in the spinal cord using adjacent transverse sections of 10.5 day embryos (A, C, E) and 12.5 day embryos (B, D, F). Sections of 10.5 day embryos are at the hind limb level and those of 12.5 day embryos are at a cervical level. (A) and (B) are phase contrast images of (E) and (F), respectively. Scale bar, 0.2 mm.

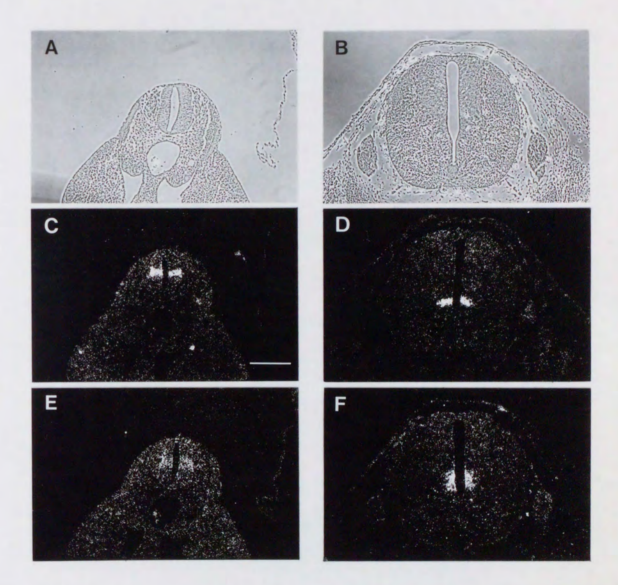


Fig. 10. Expression of Dbx2 in the limb bud mesnchyme. (A)-(C). A frontal section of 11.5 day embyro at different magnifications showing expression of Dbx2 in the hind limb bud. Arrows in (A) indicate the hybidization signal in the dorso-lateral wall of the mesencephalon (m), the hind limb bud (hl) and the spinal cord (sc). (B) is a magnification of the hind limb bud in (A). (D) and (E). A transverse section of 13.5 day embryonic fore limb bud at level of digits. Arrows in (D) indicate the hybridization signal. I and V in (F) indicate the digit number. (C) and (E) are phase contrast images of (B) and (D), respectively. Scale bars: (A), 1 mm; (B), 0.2 mm. Scales for (D) and (E) are the same as (B).

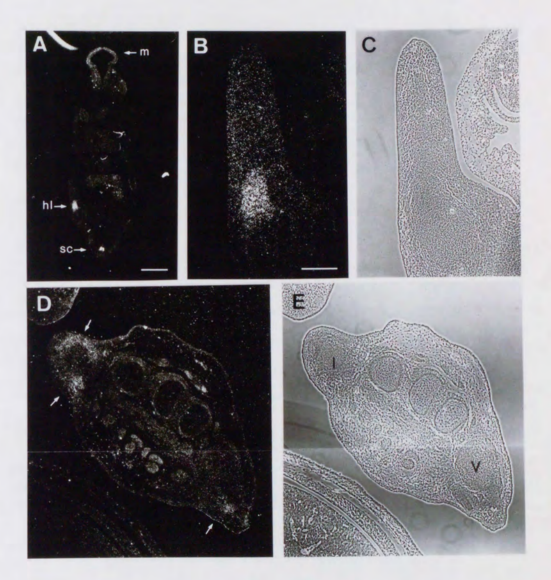


Fig. 11. Expression of Dbx2 in the tooth germ mesenchyme. (A)-(C) and (D)-(F) are frontal sections of 13.5 day and 15.5 day embryonic head, respectively. (B) and (E) are magnifications of the tooth germs indicated with arrowheads in (A) and (D), respectively. Arrows in (A) and (D) show expression of Dbx2 in the other tooth germs. (C) and (F) are phase contrast images of (B) and (E), respectively. The region between two arrowheads in (F) is dental ledge. Signal in lens seen in (A) is non-specific. Abbreviations: df, dental follicle; dp, dental papilla; ie, internal enamel epithelia. Scale bars: (A), 1 mm; (B), 0.2 mm.

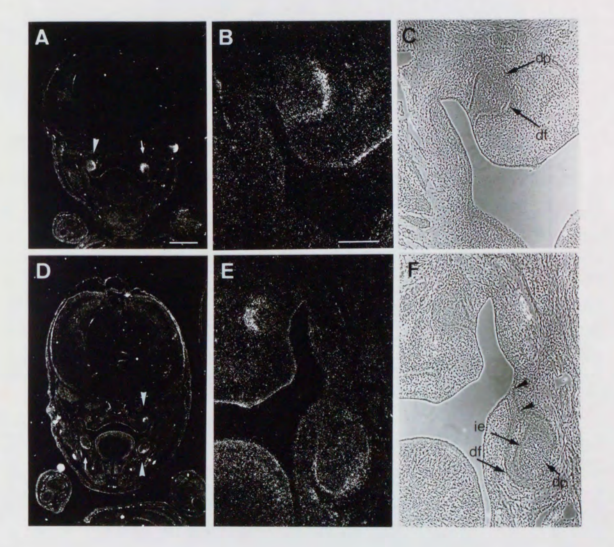


Fig. 12. Schematic illustrations of Dbx and Dbx2 expression patterns. (A) and (B). Expression of Dbx and Dbx2, respectively, in the brain of 11.5-12.5 day embryo. Gray areas indicate expression domains. The intensity of these paints reflect the relative intensity of hybridization signals in each illustration. Shaded area indicates scattering of Dbx-positive cells. (C) and (D). The spinal cord of 9.5-10.5 day embryos (C) and 11.5-12.5 day embryo (D). Filled areas are the regions expressing both Dbx and Dbx2, and shaded areas the regions expressing only Dbx2. Abbreviations: aep, anterior entopeduncular area; ap, alar plate; bh, basal zone of the hypothalamus; bp, basal plate; cge, caudal ganglionic eminence; cpt, caudal pretectum; dt, dorsal thalamus; emt, eminentia thalami; et, epithalamus; fp, floor plate; h, hypothalamus; is, rhombencephalic isthmus; lge, lateral ganglionic eminence; m, mesencephalon; ma, mammillary area; mge, medial ganglionic eminence; oc, optic chiasma; os, optic stalk; poa, anterior preoptic area; r1-r8, rhombomere 1-8; rm, retromammillary area; rp, roof plate; rpt, rostral pretectum; sc, spinal cord; se, septum; vt, ventral thalamus; vz, ventricular zone; zi, zona limitans intrathalamica.

