Transcription factor AP-2 is expressed in neural crest cell lineages during mouse embryogenesis

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We have analyzed the expression pattern of transcription factor AP-2 in mouse embryos to evaluate the potential of AP-2 as a regulator during vertebrate development. A partial cDNA encoding AP-2 was isolated from a mouse embryo cDNA library and used to prepare probes to measure AP-2 mRNA levels by RNase protection and RNA in situ hybridization. Between 10.5 and 15.5 days of embryogenesis, the relative abundance of AP-2 mRNA is greatest at 11.5 days and declines steadily thereafter. RNA in situ hybridization analysis of embryos between 8.5 and 12.5 days of gestation identified a novel expression pattern for AP-2. The principle part of this expression occurs in neural crest cells and their major derivatives, including cranial and spinal sensory ganglia and facial mesenchyme. AP-2 is also expressed in surface ectoderm and in a longitudinal column of the spinal cord and hindbrain that is contacted by neural crest-derived sensory ganglia. Additional expression of AP-2 occurs in limb bud mesenchyme and in meso-metanephric regions. This embryonic expression pattern is spatially and temporally consistent with a role for AP-2 in regulating transcription of genes involved in the morphogenesis of the peripheral nervous system, face, limbs, skin, and nephric tissues.

[Key Words: AP-2; transcription factor; mouse embryogenesis; neural crest cells; vertebrate development]

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Insights from invertebrate developmental genetics and from studies in mammalian transcription regulation have facilitated the identification of several gene families that may play important regulatory roles during vertebrate embryogenesis. Foremost among these are the homeo domain and paired-box gene families that encode Hox, POU, and Pax proteins suspected of specifying positional regulatory information [Akam 1989; He et al. 1989; Affolter et al. 1990; Kessel and Gruss 1990]; the retinoic acid (RA) receptor genes that encode proteins believed to mediate morphogenetic effects of RA [Evans 1988, Green and Chambon 1988], and the HLH genes (with helix–loop–helix domains) whose products may affect differentiation of cell lineages [Tapscott et al. 1988; Murre et al. 1989]. These diverse proteins share the common feature that they bind specific DNA sequences and regulate expression of particular target genes.

Potential regulatory roles in embryogenesis remain to be assessed for many other sequence-specific DNA-binding proteins that have been newly identified in the context of mammalian transcription regulation studies or by low-stringency cloning strategies using probe sequences from previously cloned DNA-binding regulatory proteins [for review of mammalian DNA-binding transcription factors, see Mitchell and Tjian 1989]. Some of these candidate developmental regulators belong to multigene families, while others have little or no primary sequence similarity to previously identified vertebrate or invertebrate proteins. To sort out the roles of these factors it is necessary to define their individual fields of action in the organism and the target genes that they regulate within those fields. A first step toward this goal is to analyze the distribution of the factors in adult organisms and in developing embryos. Comparing the expression patterns of many different transcription factors during embryogenesis may suggest possible regulatory cascades and factor interactions that are involved in transcriptional control of vertebrate development (Sassoon et al. 1989; Gaunt and Singh 1990; Kessel and Gruss 1990; Lonai and Orr-Urtreger 1990).

Transcription factor AP-2 was first identified and purified from HeLa cell nuclear extracts by DNA affinity chromatography using an oligonucleotide sequence from the transcriptional enhancer region of the SV40 virus [Mitchell et al. 1987]. Purified HeLa AP-2 stimulates RNA polymerase II transcription of test promoters in vitro in an AP-2-binding site-dependent manner. The protein binds sequences with the consensus 5'-GCCCGAGGC-3'; some cellular gene promoters with AP-2-binding sites include human metallothionein-IIA, proenkephalin, collagenase, and murine major histocompatibility complex (MHC) H-2Kb. Isolation of a
human AP-2 cDNA clone from a HeLa cell cDNA library was accomplished using oligonucleotide probes based on partial peptide sequences of HeLa AP-2 tryptic fragments (Williams et al. 1988). We showed recently that levels of AP-2 mRNA and protein are transiently induced during RA-dependent differentiation of human NT2 teratocarcinoma stem cells into cell types that include neurons (Lüscher et al. 1989). This finding suggested that AP-2 might participate in an RA-activated regulatory cascade during NT2 cell differentiation. A role for RA as a naturally occurring developmental morphogen has been most clearly supported by studies of chick limb skeletal development (Tickle et al. 1985; Eichele 1989); other vertebrate structures found to be developmentally sensitive to retinoid compounds include the nervous system, skin, and skeletal elements of the face (Cohlan 1953; Koplan et al. 1987; Sulik et al. 1988; Sive et al. 1990).

The results presented here suggest that AP-2 has a role in transcriptional regulation during some retinoid-affected morphogenetic processes, as AP-2 mRNA is found in several embryonic tissues at times when they are known to be developmentally retinoid sensitive. The neural crest, a major source of early AP-2-expressing cell types, has been the subject of a rich tradition of vertebrate experimental embryology (for review, see Le Douarin 1982, Hall 1988). Neural crest cells participate in the morphogenesis of a variety of structures that distinguish vertebrates from invertebrates, and it is believed that the migratory, pluripotent, and inductive features of neural crest cells have played an important part in vertebrate evolution (Maderson 1987). Fate mapping experiments have shown that neural crest-derived cell lineages give rise to much of the peripheral nervous system (sensory and autonomic neurons, glia, and Schwann cells), melanocytes, adrenal medullary cells, several other paracrine cell types, and connective tissues, bone, some muscle, and some adipose tissues in the face and ventral neck. There is evidence that cranial neural crest cells are the targets for teratogenic effects of excess RA leading to craniofacial abnormalities; however, molecular mechanisms that direct facial morphogenesis have been elusive (Wedden 1987; Sulik et al. 1988). Analyses of multiple RA receptors (RARs), retinoid-binding proteins, and RA-inducible genes are beginning to shed light on ways that complex patterns of retinoid activities may direct changes in gene transcription that underlie vertebrate morphogenetic processes (Dollé et al. 1989a,b; Eichele 1989; Perez-Castro et al. 1989; Ruberte et al. 1990, Summerbell and Maden 1990). Further investigation of the role of transcription factor AP-2 in this regard may also contribute useful information concerning retinoid-sensitive regulation of vertebrate development.

Results

Isolation of a murine AP-2 cDNA

To facilitate analysis of the embryonic expression of AP-2, a murine AP-2 cDNA clone, X22, was isolated from a 12.5-day mouse embryo cDNA library. The partial amino acid sequence of murine AP-2 deduced from the DNA sequence of the EcoRI insert in X22 is compared with the homologous carboxy-terminal region of human (hu) AP-2 (cDNA from HeLa cells, amino acids 259–437, Williams et al. 1988). Amino acid differences are noted. [B] Three segments of hu AP-2 were subcloned into plasmid vectors and used to synthesize RNA probes for RNase protection and RNA in situ hybridization experiments. The X22 insert consists of 57 bp of 3′-untranslated sequence (hatched region) adjacent to 534 bp encoding 178 amino acids of murine AP-2 (open region). A 40-bp segment (boxed region) at the 5′ end of the X22 insert is not homologous to human AP-2 (see Materials and methods, DNA sequencing). The amino acid sequence in A does not include this region. Sequence data have been submitted to the EMBL/GenBank Data Libraries.

Figure 1. (A) Murine AP-2 partial cDNA [mu AP-2 λ22] cloned from a 12.5-day mouse embryo cDNA library. The partial amino acid sequence of murine AP-2 deduced from the DNA sequence of the EcoRI insert in λ22 is compared with the homologous carboxy-terminal region of human [hu] AP-2 (cDNA from HeLa cells, amino acids 259–437, Williams et al. 1988). Amino acid differences are noted. [B] Three segments of mu AP-2 were subcloned into plasmid vectors and used for synthesis of RNA probes used in RNase protection and RNA in situ hybridization experiments. The λ22 insert consists of 57 bp of 3′-untranslated sequence [hatched region] adjacent to 534 bp encoding 178 amino acids of mu AP-2 (open region). A 40-bp segment [boxed region] at the 5′ end of the λ22 insert is not homologous to human AP-2 (see Materials and methods, DNA sequencing). The amino acid sequence in A does not include this region. Sequence data have been submitted to the EMBL/GenBank Data Libraries.

AP-2 mRNA levels in adult mouse tissues and cultured cell lines

Amounts of AP-2 mRNA in several cultured cell lines and adult mouse tissues were compared by RNase protection analyses of cytoplasmic RNA samples using an antisense probe for a 145-nucleotide segment of mouse AP-2 (R2 region; Fig. 1B). For reference, antisense probes were also used to measure RNA from Hox-2.1 (Hox-2 gene family member; Hauser et al. 1985) and Dhfr (dihydrofolate reductase, a cell proliferation-linked house-keeping enzyme, Chang et al. 1978). In these experiments, AP-2 mRNA was most abundant in fibroblastic cell lines 3T3, 3T6 [Fig. 2, lanes 1 and 2], and C3H10T1/2 (not shown). AP-2 was expressed at very low levels in undifferentiated F9 teratocarcinoma stem cells (20-fold less than in 3T3 cells, data not shown) and was not sig-
AP-2 mRNA in the mouse embryo

Figure 2. RNase protection analysis to measure AP-2 mRNA levels in several murine cell lines and adult mouse tissues. Each lane represents protected RNA segments obtained after hybridizing 5 μg cytoplasmic RNA with AP-2, Hox2.1, and Dhfr antisense RNA probes simultaneously for 16 hr at 63°C and digesting with RNase to remove unhybridized RNA (R2 probe for AP-2 was used; Fig. 1B). The gel autoradiograph bands from one experiment have been cut into strips and arranged for easy reference.

Figure 3. RNase protection analysis to assess relative abundance and region-specific expression of AP-2 mRNA during mid-embryogenesis in the mouse. R2 region antisense probe for AP-2 was used (Fig. 1B).

Temporal and spatial distribution of AP-2 mRNA during mouse embryogenesis

Poly[A]⁺ RNA samples from whole mouse embryos between 10.5 and 15.5 days of embryogenesis [or days postcoitus (dpc)] were analyzed by RNase protection. During this period, the relative level of AP-2 mRNA was greatest at 11.5 dpc and gradually declined thereafter (Fig. 3A). The period between 11.5 and 13.5 dpc represents a dramatic phase of organ and tissue morphogenesis in the mouse. To determine whether AP-2 was region-specifically expressed during embryogenesis, RNA samples prepared from dissected parts of 12.5- and 13.5-dpc embryos were analyzed by RNase protection analysis using antisense RNA probes for AP-2 and four reference genes: Hox-2.1, MyoD (Tapscott et al. 1988), Sp1 (Kadonaga et al. 1987), and Dhfr (Fig. 3). AP-2 mRNA was distributed in all three parts of dissected 12.5-dpc embryos [central nervous system, face, and body; the CNS is defined as the brain and spinal cord]. The 13.5-dpc embryos were dissected into nine parts; here, region-specific differences in AP-2 expression were observed. AP-2 mRNA was most abundant in the hindbrain, spinal cord, face, and limbs; lower levels were seen in the body [carcass]. Very low expression of AP-2 was observed in yolk sac, heart/lung, and forebrain.
while no expression was detected in the liver/gut compartment. Three of the reference genes {Hox-2.1, MyoD, and Dhfr} also exhibited region-specific differences in expression; their individual patterns were different from each other and from AP-2. In marked contrast, mRNA of transcription factor Sp1 was present at very similar levels in all nine parts of 13.5-dpc embryos. These experiments encouraged us to analyze the distribution of AP-2 mRNA during embryogenesis in more detail.

**RNA in situ hybridization to embryo sections**

AP-2 expression was analyzed by RNA in situ hybridization to paraffin serial sections of mouse embryos between 7.5 and 12.5 dpc. At 7.5 dpc (presomite gastrula), AP-2 mRNA was not detected in the embryo proper, but was expressed extraembryonically in trophoblastic tissue (not shown). At 8.5 dpc {8–12 somite stage}, AP-2 mRNA was abundant in the anterior portion of the embryo and also persisted in trophoblastic cells [Fig. 4A,B]. Transverse sections through the head revealed strong signal in lateral head mesenchyme and surface ectoderm but little or no expression in the brain plate or the sparse mesenchyme medial to it (Fig. 4C,D). In the posterior body, AP-2 mRNA was detected in surface ectoderm and neural folds but not in the neural plate, somites, or presomitic mesoderm [Fig. 4C,E]. The pattern of AP-2 mRNA in neural folds and head mesenchyme at this time closely approximates the spatial distribution of the earliest identified neural crest cells in the mouse embryo [Chan and Tam 1988]. Neural crest cells originate in a craniocaudal wave from neural fold neuroectoderm along the lateral margins of the vertebrate neural plate before and during closure of the neural tube. Migrating away from the dorsal axis [LeDouarin 1982; Tan and Morriss-Kay 1986, Chan and Tam 1988], neural crest cells proliferate and collect in multiple locations where they form or contribute to a variety of structures [neural crest-derived cell types were mentioned earlier]. The location of AP-2 signal in neural folds at 8.5 days [Fig. 4C,D] suggests that expression in neural crest cells begins prior to or coincident with their emigration.

In frontal sections of 9.5-dpc embryos, AP-2 mRNA continued to persist in surface ectoderm (pre-epidermis) and was pronounced in cephalic mesenchyme (Fig. 5A). The latter consists predominantly of midbrain- and hindbrain-originating neural crest cells that are forming rostral and branchial arch mesenchymal populations and doroasal ganglioblast aggregations [cranial ganglia primordia] adjacent to the hindbrain. The prominent lateral regions of AP-2 expression seen in Figure 5A are consistent with the trigeminal and facial–auditory ganglia primordia (left) and branchial arch mesenchyme slightly ventral to ganglia primordia (right) [Altman and
Sensory neurons that arise in the trigeminal and facial—acoustic ganglia and motor neurons that pass through the ganglia will innervate mandibular and hyoid arch-derived facial structures via cranial nerves V, VII, and VIII. At this time, AP-2 was also expressed at low levels in mesonephric regions, in pericardial tissues, and diffusely around the dorsal aorta, gut, and lateral body and in limb bud mesenchyme (sections not shown).

At 10.5 dpc, AP-2 signal was evident in the dorsal root ganglia (DRG) [Fig. 5D,G]. DRG develop from neural crest cells that aggregate segmentally on either side of the spinal cord in register with the somites; these spinal sensory ganglia are related to the more specialized cranial ganglia to which neural crest cells contribute [Le Douarin and Smith 1988]. Frontal sections passing through the head identified continued strong AP-2 expression in the epidermis, trigeminal ganglia, and regions of facial mesenchyme heavily populated by neural crest cells [Fig. 5F]. AP-2 signal was also evident in the 10.5-dpc embryo spinal cord [Fig. 5G].

In 11.5-dpc embryos, AP-2 mRNA persisted in the same structures as at 10.5 dpc, and several new sites of expression were also observed [Fig. 6]. High AP-2 expression was seen in a continuous longitudinal region of the CNS extending at least as far anteriorly as the hindbrain—midbrain junction [Fig. 6A,B] and posteriorly in the spinal cord to at least the level of the hind limbs [Fig. 6C,D]. Transverse views of the spinal cord and hindbrain showed that this AP-2-expressing column was mediolateral and mainly dorsal to the sulcus limitans [Fig. 6C,D]. The dorsal-most part of the expressing region coincided with the dorsal entry zone where developing axons of DRG sensory neurons enter the spinal cord. Similarly, in the hindbrain, the AP-2 column coincided with the area where boundary caps of cranial ganglia contacted the CNS [Fig. 6C, facial—acoustic ganglia; others not shown]. The disposition of this CNS column suggests that it includes not only the discrete regions where developing axons of sensory neurons enter the CNS but also sensory nuclei [synaptic centers] and longitudinal tracts along which sensory fibers travel to reach these nuclei. Sensory nuclei and tracts typically develop in columns dorsal and lateral to the sulcus limitans, in contrast with motor nuclei, which are found in ventral—medial columns [Arey 1974; Hopper and Hart 1984]. Our present data suggest that AP-2 expression in the lateral CNS may mature craniocaudally, as expression seems to be restricted to the hindbrain and anterior spinal cord at 10.5 dpc [Fig. 5A,C, and sections not shown]. At 11.5 dpc, AP-2 expression was also detected in sympathetic ganglia [SG, all neural crest cell derived], along spinal nerve paths between DRG and SG, and at low levels in cranial ganglia developing posterior to the otic vesicle [including the superior, petrosal, and nodose ganglia, Fig. 6B and sections not shown].

In 12.5-dpc embryos, AP-2 signal continued to be found in peripheral nervous system [PNS] ganglia, epidermis, and lateral CNS [Fig. 7]. Expression in limb mesenchyme was concentrated distally (in the progress zone) and in a small medial patch [Fig. 7D]. Strong AP-2 signal was evident in facial mesenchyme adjacent to nasal epithelia and along the nasal lacrimal groove [Fig. 7A—D], and in corneal layers of the eye [Fig. 7B] [the contribution of neural crest cells to the cornea has been noted in chick embryos; Le Douarin 1982]. The transverse section in Figure 7B shows AP-2 mRNA on the right side of the head in the trigeminal ganglion [ophthalmic branch] and anterior metencephalon. Serial reconstruction showed that this slightly skewed section passes through the bottom of the right eye and middle of the left eye. The section most likely marks the anterior boundary of the continuous column of AP-2 expression in the lateral CNS [expression in the left trigeminal ganglion and metencephalon was observed several sections more posteriorly]. The arc of AP-2 signal in the metencephalon extends from the dorsal alar region into the pons and is similar to the known migration path of alar plate neuroblasts and glio blasts that form the pontine nuclei. These become synaptic centers connecting the phylogenetically newest part of the cerebellar cortex with the newest part of the cerebral cortex for the coordination of muscular activity [Hopper and Hart 1984].

Sections passing more caudally through 12.5-dpc metencephalon and myelencephalon regions showed AP-2 expression in alar regions [future cerebellum], in patches near the sulcus limitans [possibly corresponding to sensory nuclei and visceral efferent nuclei], and along pathways characteristic of alar cells migrating to form the olivary nuclei in the ventral myelencephalon [data not shown]. The pattern of AP-2 expression in the hindbrain and spinal cord between 11.5 and 12.5 dpc is consistent with expression in sensory nuclei and sensory tracts which, when taken together, constitute a more or less continuous longitudinal column caudal to the hindbrain—midbrain junction. Autonomic efferent neurons also develop in intermediolateral regions of the CNS (in general, visceral efferent nuclei), and it is possible that the AP-2-expressing zone in the lateral CNS may include some of these autonomic nuclei as well. In addition to the continuous lateral column of AP-2 expression in the CNS, AP-2 mRNA was also seen in limited portions of the mesencephalon and diencephalon at 12.5 dpc [Fig. 6D and sections not shown].

Discussion

Transcription factor AP-2 exhibits a novel expression pattern between 8.5 and 13.5 days of mouse embryogenesis and is restricted to a subset of ectodermal and mesodermal tissues. The principle part of this pattern is a remarkable summary of the movements and aggregations of the major neural crest cell lineages; AP-2 mRNA is present in neural folds and head mesenchyme of neural crest cell origin at 8.5 dpc and subsequently in neural crest cell-populated tissues, which include PNS ganglia and facial and branchial arch mesenchyme where neural crest cells form connective tissue and bones. Appreciable AP-2 expression also occurs in epi-
dermal ectoderm, limb bud mesenchyme, mesometanephric regions, and a lateral portion of the CNS. AP-2 is not significantly expressed in mesoderm of the notochord, somites, or heart, which lie medial to the neural axis, or in a variety of endodermally derived organs and tissues (liver, lungs, gut, and yolk sac). This endodermal exclusion is reflected in vitro in a similar lack of appreciable AP-2 expression in endodermally fated F9 teratocarcinoma and HepG2 hepatoma cells [Lüscher et al. 1989].

Figure 5. (See facing page for legend.)
A potentially significant link between AP-2-expressing regions in the embryo is that they are all within tissues known to be developmentally sensitive to retinoid compounds. High concentrations of RA (acidic vitamin A) cause craniofacial, limb, nervous system, and primary body axis defects in vertebrate embryos (Wedden 1987; Wilde et al. 1987; Sive et al. 1990). The nature and severity of these defects are stage- and dose-dependent and generally affect both ectodermal and mesodermal tissues. In young animals, lack of dietary vitamin A (retinol) or retinoid derivatives leads to skeletal and nervous system deficiencies, dry and thickened skin, kidney degeneration, and sterility in both sexes (Wolbach and Howe 1925). The pattern of expression of transcription factor AP-2 in the embryo is highly suggestive of a role for AP-2 in regulatory programs during retinoid-sensitive morphogenetic periods in many of the above-mentioned tissues and structures.

Expression in neural crest cells and other ectodermally derived tissues

AP-2 expression in DRG is unambiguously neural crest lineage-specific as DRG contain cell types (sensory neurons, support cells, and Schwann cells) descended entirely from neural crest cells [LeDouarin 1982]. In rostral and ventral craniofacial regions where AP-2 mRNA is found, mesectodermal neural crest cells constitute the predominant mesenchymal cell type (especially between 8.5 and 10.5 dpc). These sites, together with the DRG, provide strong evidence that AP-2 is expressed in both neural and mesectodermal lineages of neural crest cells. In cranial ganglia (CG), support cells are commonly held to be all neural crest-derived; however, sensory neurons in CG can originate from neural crest cells, placodal ectoderm, or both depending on the particular ganglia [D'Amico-Martel and Noden 1983; LeDouarin and Smith 1988]. Thus, AP-2-expressing cells in CG are not necessarily of neural crest descent. However, although AP-2 is expressed in surface ectoderm as early as 8.5 dpc, it is not appreciably expressed in otic, olfactory, or epicranial placodes (data not shown). This suggests that AP-2 is turned off during differentiation of placodal ectoderm from surface ectoderm. In view of this, it seems more likely that AP-2-expressing cells in all CG are neural crest-derived neurons and/or support cells as in DRG.

High AP-2 expression occurs in the epidermis and the lateral CNS. Melanoblasts originate from neural crest cells and begin to migrate into future dermal regions as early as 8.5 dpc in the mouse but are not believed to invade the epidermis until several days later. Thus, AP-2 signal in surface ectoderm at 8.5 dpc is not likely to be contributed by melanoblasts. As persistent AP-2 expression in multiple neural crest cell lineages is already evident, it is reasonable to ask whether there is a fundamental ectodermal lineage relationship that links AP-2-expressing cell types in surface ectoderm, neural fold neuroepithelia, and the lateral CNS. Single cell in situ labeling has identified individual precursor cells in the neural plate of early somite chick embryos that generate both neural crest cells and a sister lineage of neural cells that remain in the CNS [Bronner-Fraser and Fraser 1989]. Whether the latter cells proliferate and come to populate a specific region of the CNS after neural tube closure (such as the lateral CNS region where AP-2 is expressed by 10.5 dpc) is not known. Another fate mapping experiment in the chick has indicated that, even as late as the 3-somite stage [8-dpc mouse embryo], neural fold epithelia in the anterior head still harbors some cells destined to form surface ectoderm in the face (Couly and LeDouarin 1990). These experiments underscore potential lineage relationships between epidermal, neural crest, and CNS progenitor cells within or near the margins of the neural plate at this time.

Expression in mesodermal tissues

AP-2 transcripts are detected in a subset of mesodermal tissues that include facial and limb bud mesenchyme and mesometanephric regions. In keeping with the observed expression of AP-2 in embryonic nephric tissues,
Figure 6. RNA in situ hybridization to 11.5-dpc mouse embryo sections. AP-2 expression in DRG [1], lateral CNS [2], and sympathetic ganglia [3] is noted. Approximate planes for sections in C and D are shown in A. [A] Sagittal section showing expression in DRG, frontonasal-prominence (FN), and lateral portions of the myelencephalon (My, posterior hindbrain) and spinal cord (sc). [See note in Fig. 6D regarding artifact signal from blood cells (bc)]. (Tel) Telencephalon; (Di) diencephalon; (Met) metencephalon (anterior hindbrain), (Mes) mesencephalon, (Ph) pharynx; (Mn) mandibular arch/tongue. (B) Parasagittal section through the lateral head and upper body, passing through the eye, mandibular (Mn) and hyoid (Hy) arches, pharynx, otic vesicle (ov, auditory vesicle), and heart (Ht). Dark-field views shows AP-2 signal in the lateral spinal cord and hindbrain (extending anteriorly to Met), in DRG (drg), sympathetic ganglia [3] near the heart, and facial–acoustic cranial ganglia [gVII–VIII] [part of signal here is due to extraneous material.
AP-2 mRNA in the mouse embryo

limited to this section [possibly torn from the adjacent hindbrain]. Low, diffuse expression is seen in petrosal [gXi] and nodose [gXii] [i, inferior] ganglia and along spinal and cranial nerve paths. Bright- and dark-field views in B are adjacent serial sections. [C] Frontal section showing expression in myelencephalon, facial—acoustic ganglia and their boundary caps apposing the myelencephalon (My), facial mesenchyme lateral to the pharynx, epidermis, spinal cord, DRG, mesonephric regions (mnp), and forelimb buds (Lf). The lateral patches of AP-2 expression in the My are mediolateral and mainly dorsal to the sulcus limitans (sl), the slight indentation that marks the boundary between dorsal/sensory/alar plate and the ventral/motor/basal plate (in transverse views of the spinal cord in C and D, the sulcus limitans is seen on the outer walls). [D] Frontal section showing AP-2 signal in facial mesenchyme, epidermis, DRG, spinal cord, mesonephric duct (md), and diencephalon. [Ns] Nostrils; [csc] spinal cord caudal to hindlimbs; [Lh] hindlimb bud.
Figure 7. (See facing page for legend.)
AP-2 mRNA is easily detected in cell lines derived from kidney (CV-1 and BSC-40 monkey kidney cells; P.J. Mitchell, unpubl.). HeLa cells, the human cell line in which AP-2 was first identified, were originally cultured from an epithelial carcinoma of the cervix. In the embryo, nephric and genital tract tissues develop in intimate association from urogenital ridge mesoderm. The mesonephros, which expresses AP-2, directly contributes tissue to the developing genital tracts of both males and females during its regression and replacement by the metanephros [adult kidney]. Analysis of later embryonic stages and adult tissues to look for AP-2-expressing cells in developing or adult genital tracts may suggest whether AP-2 expression in HeLa cells is consistent with their tissue origin.

In the embryonic head, AP-2-expressing cells are nearly continuous from the lateral CNS, to CG, and out to facial mesenchyme and epidermis. Perhaps concerted AP-2 expression in these different ectodermal and mesenchymal cell lineages reflects the fact that they have inherited gene expression programs that are fundamentally coordinated for interactive morphogenetic processes during facial development and innervation. In facial mesenchyme, the distribution of AP-2 mRNA is most consistent with expression in cranial neural crest-derived mesectoderm, which will form skeletal elements and connective tissues in the face. In early limb buds, AP-2 mRNA is diffusely distributed throughout the mesenchyme, but by 12.5 dpc it is concentrated at the distal end [progress zone] and in at least one small medial patch. Expression in limb epidermis is evident at all stages analyzed. A recent fate mapping experiment demonstrated that trunk level neural crest cells normally migrate into chick limb bud mesenchyme in detectable numbers [Serbedzija et al. 1990]. These cells were considered likely to be Schwann cell precursors that associate with incoming neurons, which will innervate the limb. Although pre-Schwann cells could conceivably contribute part of the AP-2 signal in limbs, the pattern of AP-2 expression in limb bud mesenchyme, as in facial mesenchyme, is most consistent with expression in cells that are precursors to skeletal element- and connective tissue-forming cells.

A prevailing model of limb development asserts that RA is distributed in a morphogenetic gradient emanating from a posterior part of the limb bud, the zone of polarizing activity (ZPA). Undifferentiated and actively dividing mesenchymal cells in the distal progress zone receive cues from the limb apical ectodermal ridge (AER) which, when interpreted in the context of the RA gradient, render them competent to differentiate into a particular skeletal structure on being displaced proximally out of the progress zone [Eichele 1989; Summerbell and Maden 1990]. As an outcome of this patterning process, appropriate skeletal elements of the limb develop in a proximodistal order. Morphogenesis of the limbs and the largely neural crest-derived frontonasal mass of the face are both notably sensitive to RA [Wedden 1987, Wilde et al. 1987, Eichele 1989]. Exposure to high concentrations of RA alters the normal patterning of skeletal elements in these sites causing truncation or duplication depending on the mode and time of exposure. Spatially and temporally, the expression pattern of AP-2 in the face and limbs is consistent with a role for AP-2 in regulatory programs that bring about patterning of skeletal elements in these regions.

The basic molecular principles that regulate retinoid-sensitive morphogenetic processes are probably not far from being understood in rough outline [Eichele 1989; Summerbell and Maden 1990]. An important part of this control is mediated by intracellular RARs, which are sequence-specific DNA-binding transcriptional regulators [Evans 1988; Green and Chambon 1988]. RARs and other retinoid-binding cellular proteins that may be involved in mediating morphogenetic effects of retinoids are distributed in complex patterns in vertebrate embryos [Dollé et al. 1989a; Perez-Castro et al. 1989, Ruberte et al. 1990]. Additional regulators of gene expression that have been implicated in retinoid-affected processes include Hox genes [Dollé et al. 1989b; Kessel and Gruss 1990; Wilkinson and Krumlauf 1990]. Like AP-2, many of the murine Hox genes are first expressed between 7 and 8.5 dpc, reach peak levels in mid-embryogenesis, and are expressed region specifically in, among other places, continuous longitudinal compartments of the CNS with discrete anterior boundaries [Kessel and Gruss 1990]. Also like AP-2, many of the Hox genes are demonstrably induced by RA treatment of NT2 teratocarcinoma cells, and the varying kinetics of RA induction for different Hox family members supports the idea that they could be differentially regulated by endogenous retinoids in the embryo [Simeone et al. 1990; Stornaiuolo et al. 1990]. Other indirect evidence suggesting that Hox genes may be involved in regulating retinoid-
sensitive aspects of morphogenesis is found in transgenic mice that express Hox-1.1 ectopically (Balling et al. 1989). Nearly ubiquitous expression of Hox-1.1, normally not found in head regions, results in a spectrum of craniofacial defects resembling the teratogenic effects of RA.

Target genes regulated by transcription factor AP-2

The pattern of AP-2 expression in neural crest cells and the embryonic nervous system suggests that a major role for AP-2 may be in transcriptional regulation involved in establishing the PNS and its connections with the CNS. As the molecular mechanisms that generate positional regulatory information may be fundamentally similar for different morphogenetic or patterning events in the embryo, it is possible that the role of AP-2 in the nervous system is related to its role in the face, limbs, epidermis, and nephric system.

On the basis of current understanding of transcriptional activation, it is likely that target genes for AP-2 will vary in different parts of the embryo as a result of synergistic or antagonistic interactions between AP-2 and other region-restricted transcription factors or modifers of AP-2 activity. By analogy to invertebrate systems where developmental regulatory hierarchies are at least superficially outlined, we suspect that AP-2 acts in regulatory cascades, that at least some of its target genes are other regulatory proteins, and that it may have negative as well as positive transcriptional effects on target genes. A location for AP-2 generally in middle stages rather than very early or terminal differentiation stages of developmental regulatory cascades is suggested by several observations. AP-2 appears to be dramatically activated between 7.5 and 8.5 dpc of mouse embryogenesis, well beyond the initiation of gastrulation at 6.5 dpc. Expression peaks between 10.5 and 12.5 dpc, while many primordial organs and structures are being established, and then declines steadily thereafter. During induced differentiation of several types of cell lines in vitro, we find that AP-2 mRNA levels actually decline or are only transiently increased [unpublished observations with 3T3-L1 adipocytes and G7 and G8 myoblasts (P.J. Mitchell, unpubl.) and previous data on NT2 teratocarcinoma cells (Lüscher et al. 1989)]. Finally, expression levels in adult organs that we have tested are consistently lower than expression in embryonic or less differentiated cultured versions of these tissues. These findings suggest that in the ectodermal and mesodermal lineages where AP-2 is used, it may be more frequently involved in stem cell or precursor integrity and act during proliferative and determinative phases, rather than during terminal phases of differentiation.

Regulators of AP-2 expression

The anterior location of AP-2 mRNA in 8.5-dpc embryos discounts most of the known Hox genes as positive activators of very early AP-2 expression; an exception may be Hox 7.1, which exhibits some temporal and spatial overlap with AP-2 in the anterior head (Hill et al. 1989; Robert et al. 1989). The complex overlap between domains of expression for AP-2 and different Hox genes in more posterior regions of the embryo does not suggest any simple regulatory relationships between these factors. Similarly, the embryonic expression patterns of the known RARs and retinoid-binding proteins (Dollé et al. 1989b; Perez-Castro et al. 1989; Ruberte et al. 1990) overlap with that of AP-2 to different extents, and no straightforward regulatory relationships are immediately evident. Our previous analysis of NT2 teratocarcinoma cells suggested that the AP-2 gene was not a primary target for activation by RARs in these cells, as the kinetics of AP-2 mRNA appearance were more consistent with AP-2 being secondarily induced in an RA-activated regulatory cascade [Lüscher et al. 1989]. The fact that AP-2 mRNA levels remained low throughout a 7-day RA treatment of F9 teratocarcinoma cells, which initially express RAR α, β, and γ [Hu and Gudas 1990], also suggests that the AP-2 promoter may not be directly regulated by RARs. However, as new RAR types and isoforms are still being characterized (Kastner et al. 1990; Mangelsdorf et al. 1990) and RARs may activate their target genes by synergistic interaction with other transcription factors of restricted distribution, the possibility that AP-2 is directly regulated by RARs in certain cell types cannot be excluded. A better understanding of the regulation of AP-2 awaits more thorough analyses of the AP-2 gene and promoter.

Future prospects

In our present study we have identified regions where AP-2 transcripts are expressed in the mouse embryo between 7.5 and 13.5 dpc, a period that covers many of the major morphogenetic events of vertebrate embryogenesis. AP-2 protein and transcriptional activity in the embryo may actually be more restricted than the RNA distribution pattern as a result of RNA processing, translational, or post-translational regulations operating in different cell types. Future experiments will be directed toward localizing AP-2 protein at the cellular level, identifying important target genes, dissecting the AP-2 gene promoter, and determining the effects of blocking AP-2 function in different regions of the embryo. Additional in situ studies with appropriate markers should enable a more thorough characterization of cell types expressing AP-2, including cells not specifically addressed in our present study such as neural crest derivatives in the adrenal medulla and other paraendocrine tissues. These and experiments to examine the relationships between AP-2 and other suspected developmental regulators may ultimately contribute to a better understanding of transcriptional regulation during vertebrate embryogenesis.

Materials and methods

Screening a murine igt10 cDNA library for AP-2 clones

A 45-nucleotide oligonucleotide 5’-GTCTGTGAACCTCCT-CGATAGCCG-3’
TGGCAATCTGTTGTTGGCIAIGACA TTCT-3’ deduced from a 15-amino acid peptide sequence of human AP-2 (Williams et al. 1988) was end-labeled with [32P]ATP using T4 polynucleotide kinase. This probe was used to screen a λgt10 cDNA library prepared from 12.5-day mouse embryo cDNA (Hauser et al. 1985). Procedures were essentially according to Benton and Davis (1977) with minor modifications; the final wash conditions were 48°C with 2 x SST + 0.5% SDS. A screen of 106 plaques yielded a single murine AP-2 cDNA clone, λ22, containing a 635-bp EcoRI insert that hybridized strongly to a full-length human AP-2 probe in a moderately stringent Southern blot analysis.

DNA sequencing
The 635-bp EcoRI insert from λ22 and RsaI, Alul, and HaeIII fragments derived from it were sequenced after cloning into EcoRI-cut plasmid KS/SK (Stratagene) or SmaI-cut M13mp10 using the dideoxy-nucleotide method (Biggin et al. 1983). A 40-bp segment at one end of the 635-bp EcoRI insert of λ22 is not homologous to sequences in human AP-2. This short segment, which is adjacent to the 5'-most portion of the mouse AP-2 sequence, most likely was appended as a cloning artifact during the construction of the λgt10 library because it is not protected in RNase protection experiments using adult or embryo mouse mRNAs (including RNA from 12.5-day embryos, which the λ22 clone represents).

Cell culture
F9 cells were grown on gelatinized tissue culture plates. The confluent plates were rinsed twice with cold PBS; cells were harvested in 0.5% SDS. A screen of 106 plaques yielded a single murine AP-2 cDNA clone, λ22, containing a 635-bp EcoRI insert that hybridized strongly to a full-length human AP-2 probe in a moderately stringent Southern blot analysis.

RNA preparation
RNA from cultured cells was prepared as follows: Near-confluent plates were rinsed twice with cold PBS; cells were scraped with a rubber policeman into cold PBS, transferred to a microfuge tube containing 220 μl Bonner buffer [7 M urea, 350 mM NaCl, 10 mM Tris HCl (pH 7.5), 10 mM EDTA, 1.4% SDS] and vortexed. RNA was prepared by two extractions with dH2O-saturated phenol, one extraction each with phenol/urea, 350 mM NaCl, 10 mM Tris HCl (pH 7.5), 10 mM EDTA; and proteinase K (freshly prepared buffer was preincubated for 10 min at 37°C), incubated for 30 min at 37°C, phenol/chloroform-extracted until the interface was clear, chloroform-extracted for a second time, and precipitated with ethanol. Poly(A)- RNA was isolated from total embryo RNA by oligo(dT)-cellulose chromatography.

Plasmids for RNA probe synthesis
Fragments cloned into pBluescript KS/SK vectors for radiolabeled RNA probe synthesis using T7 and T3 RNA polymerases were as follows. AP-2: The 165-bp RsaI fragment (R2) containing the 5'-most 143 bp of mouse AP-2 sequences in the λ22 insert was taken as a BamHl–EcoRI fragment from an M13mp10 clone used for DNA sequencing (see above), this was recloned into EcoRI + BamHl-cut pBS-KS. The resulting plasmid pKS-R2 was linearized with EcoRI for AP-2 antisense probe synthesis with T7 RNA polymerase. The 22-bp non-AP-2 segment in the R2 probe does not interfere with the quantitation of AP-2 by RNase protection. However, in RNA in situ hybridization experiments, we observed that the R2 probe gave substantially higher background signal compared with similar exposure times with two other AP-2 antisense probes. Other AP-2 fragments: The 240-bp BglII fragment of mouse AP-2 was cloned into the BamHl-cleaved pBS-SK to make pB240, the 340-bp BglII–EcoRI fragment was cloned into BamHl–EcoRI-cleaved pBS-KS to make pB340. After appropriate linearization within polymer linker regions, these plasmids were used to prepare B240 and BR340 AP-2 antisense and sense probes using T3 and T7 RNA polymerases, respectively. Sp1: Approximately 220-bp PstI–BamHl fragment from p11muSp1 was subcloned into PstI–BamHl-cleaved pBS-KS; the resulting plasmid, pKS-Sp1Pb, was linearized with PstI for antisense probe synthesis with T7 RNA polymerase. Hox-2.1: A 390-bp PvuII–Xbal fragment from the EcoRI insert of λ6.8 (Hauser et al. 1985) was cloned into HindIII–XbaI-cleaved pBS-KS; the resulting plasmid, pKS-A6XP was linearized with XhoI for antisense probe synthesis with T7 polymerase. Dfr: A 450-bp PstI–BamHl fragment from pDHFR21 (Chang et al. 1978) was cloned into PstI–BamHl-cleaved pBS-KS; the resulting plasmid KS-DHFR.PB was cut with HpaII for antisense probe synthesis with T7 polymerase.

Preparation of RNA probes and RNase protection analysis
[a-32P]UTP RNA probes were prepared and RNase protection analysis was performed essentially as described by Lüscher et al. (1989).

RNA in situ hybridization to mouse embryo sections
CBA/Ca mice were mated, and embryos were obtained from pregnant females by standard techniques (Hogan et al. 1986). Paraffin-embedded sections were prepared from paraformaldehyde-fixed embryos, and in situ hybridization was performed essentially as described by Wilkinson et al. (1987). α-[32P]UTP-labeled antisense and sense RNA probes were prepared for T7 and T3 polymerase-directed synthesis using linearized pBluescript plasmids. When hybridized to adjacent serial sections, B240 and BR340 AP-2 antisense probes (nonoverlapping probes, Fig. 1B) identified similar region-specific AP-2 expression; however, the B240 probe consistently yielded the lowest background [this could be related to the presence of a GC-rich tract in the 3'-untranslated portion of AP-2 in BR340]. Results shown here were obtained with the B240 probe. The mouse cardiac actin probe (Sassoon et al. 1988) protects a 50-bp segment in the 5'-untranslated portion of cardiac actin. Hybridizations were done at 55°C for 16 hr, and washes were performed at high stringency. Slides were coated with Ilford Nuclear Research Emulsion, and autoradiographic exposures were done for 5–7 days. Negative results with AP-2 sense control probes demonstrated the specificity of our in situ probes; and RNase protection analysis of RNA purified from 13.5-dpc embryo parts pro-
vided a stringent complementary assay that was consistent overall with the region-specific pattern of AP-2 expression documented by our RNA in situ analysis.

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