Spec3: embryonic expression of a sea urchin gene whose product is involved in ectodermal ciliogenesis

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We have characterized the temporal and spatial expression of Spec3 mRNA in embryos of the sea urchin, Strongylocentrotus purpuratus. This mRNA, 2.0 kb in length, is present at low levels in unfertilized eggs but accumulates rapidly during cleavage, increasing 50-fold by hatching blastula stage. Message levels then decline abruptly, remain constant during mesenchyme blastula and gastrula stages, and increase again during prism and pluteus stages. This accumulation pattern is quite similar to that of the ectodermally expressed β-tubulin mRNAs described recently by Harlow and Nemer (1987a). In situ hybridization shows that although Spec3 message accumulates in all blastomeres at early blastula stages, it later becomes restricted to ectoderm. By late blastula stage, hybridization is strongest in the animal hemisphere. At gastrula, signals are variable over ectoderm, and by pluteus, grains are concentrated in the ciliary band, though present in other ectodermal cells as well. Deciliation and regeneration of cilia in gastrula-stage embryos results in a four- to fivefold increase in Spec3 mRNA levels, implying that the Spec3 gene product is associated with ciliogenesis. Spec3 mRNA is encoded by a single gene in the haploid genome, and characterization of the gene shows that it contains three exons that encode an open reading frame for a hydrophobic protein of 21.6 kD. The reading frame reveals that the carboxy-terminal part of the protein contains two long hydrophobic stretches, 31 and 37 residues long, separated by short hydrophilic regions of six to eight residues. The presence of these two distinct hydrophobic stretches suggests that the Spec3 protein contains two α-helical domains that either span the lipid bilayer or are associated with some other hydrophobic environment.

[Key Words: Sea urchin embryo, embryonic ectoderm, ciliogenesis, in situ hybridization, hydrophobic protein]

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in embryogenesis than had been reported for any other cloned gene, with the exception of the α-histone genes [Maxson and Wilt 1982]. Although Spec3 was selected because it is enriched in ectoderm cells of plutei, maximal accumulation of Spec3 message occurs long before the morphological differentiation of any ectodermal cell types.

To obtain a detailed picture of Spec3 transcript accumulation, we isolated total embryo RNA either at 4-hr intervals (unfertilized egg through prism stage) or 6-hr intervals (prism through pluteus stage) and analyzed it by RNA blotting. A portion of the Spec3 cDNA containing the entire open reading frame and small regions of 5' and 3' untranslated sequences was subcloned and used as a single-stranded hybridization probe. Multiple exposures of the autoradiogram were scanned densitometrically to quantify relative mRNA abundance (Fig. 1). As demonstrated in our earlier study, Spec3 transcripts are present at very low levels in the egg [Bruskin et al. 1981] and do not increase in abundance through the first four or five cleavages (6–7 hr). Levels then increase dramatically: threefold by morula stage (11 hr) and 50-fold by hatched blastula stage (20 hr). The transcript levels then decrease rapidly by gastrula stage to about one-fourth of their maximal levels. Spec3 mRNA accumulates again in prism and pluteus stages, though never to hatched blastula levels. These data indicate that Spec3 mRNA accumulates during embryogenesis in a biphasic manner, with a major peak of accumulation around hatched blastula stage and a minor peak around pluteus stage.

We found evidence for variation in the level of Spec3 in different egg batches. The experiment shown in Figure 1 was carried out with a pool of eggs from several females, but similar analyses were carried out to 50 hr of development on two other occasions with different pools of eggs. In each case, hybridization signals were normalized for RNA loads using a probe for mitochondrial 16S rRNA, whose concentration does not change during embryogenesis [Wells et al. 1982]. Although all three experiments showed the same rapid accumulation of blastula stage, there were significant differences, ranging from 10% to 50% of the maximal level, in the amount of message present between 25 and 50 hr.

Alexandraki and Ruderman [1985] showed that in *Lytechinus pictus*, RNA blots utilizing probes for both α- and β-tubulin mRNAs produced accumulation patterns similar to that seen for Spec3 transcripts in *S. purpuratus*. Recently, β-tubulin messages from *S. purpuratus* have been characterized by Harlow and Nemer [1987a]. Using a probe from the 3' untranslated region specific for the β1-tubulin RNA, they found that this message was highly enriched in ectodermal tissue. The inset in Figure 1 shows the temporal pattern of β1-tubulin mRNA expression redrawn from the data of Harlow and Nemer [1987a]. The accumulation of this message and of two other moderately ectoderm-specific tubulins, β2 and β3, bears a striking resemblance to Spec3 message accumulation.

Several lines of evidence indicate that the Spec3 transcript encodes a protein product. First, a nearly full-length cDNA clone was isolated from a library constructed from oligo(dT)-primed polysomal poly(A) RNA. Second, we have identified a 208-codon open reading frame that can be translated in vitro (see below and Eldon 1988). Third, hybridization analysis shows that
the sequence of the open reading frame is conserved between *S. purpuratus* and the distantly related species *L. pictus*, whereas the 3' untranslated region is not: a Spec3 probe containing the open reading frame detected three bands in a blot of genomic DNA when hybridized at a moderately high stringency of 0.18 M Na⁺ at 68°C, whereas a probe containing only 3' untranslated sequence did not (Eldon 1987).

**Spatial distribution of the Spec3 transcript during embryogenesis**

In situ hybridization shows that Spec3 mRNA is transiently synthesized in all regions of the very early blastula (12 hr), but rapidly becomes restricted to ectoderm lineages. In agreement with conclusions from the RNA blot analysis (Fig. 1), grain densities over sections of unfertilized eggs were not detectably higher than background (Fig. 2a). Two separate measurements indicated signals were three- and fourfold higher over sections of late cleavage-stage embryos (12 hr, or about 160–180 cells; Cox et al. 1984) than over sections of eggs. These new transcripts accumulate in all regions of the 12-hr embryo (Fig. 2b). In approximately 5% of randomly selected sections, we observed very small regions that appeared unlabeled, but corresponding areas in adjacent sections were labeled at the general level of the section. Thus, any regions free of Spec3 mRNA are either small (<5 μm in diameter) or artifically unlabeled. We conclude that all major lineages of the embryo (presumptive oral ectoderm, aboral ectoderm, primary mesenchyme, secondary mesenchyme, and endoderm) accumulate Spec3 mRNA during cleavage.

At early blastula stage before hatching (17 hr, Fig. 2c), the distribution of Spec3 mRNA is no longer uniform. In individual sections the majority of cells are heavily labeled, whereas one small region of contiguous cells is essentially unlabeled. The lack of distinguishing morphological characteristics prevents positive identification of the unlabeled cells at this stage, but they likely correspond to precursors of the thickened vegetal plate, which is the only consistently unlabeled region of the embryo.

**Figure 2.** Distribution of Spec3 mRNA in sea urchin embryos. Sections of *S. purpuratus* embryos were prepared and hybridized with [3H]-labeled antisense RNA probes as described in Methods. Shown are pairs of phase-contrast and dark-field photographs of egg (a), 12-hr early blastula (b), 17-hr blastula (c), 20-hr hatched blastula (d), 27-hr mesenchyme blastula (e), perpendicular sections through 55-hr gastrula (f,g), 73-hr pluteus (h), and 84-hr pluteus (i). All sections except that shown in c were taken from the same slide; therefore, grain densities can be compared. The exposure time was 4 weeks. The section shown in c was taken from another experiment in which the specific activity of the probe was lower (82%) and the exposure time was 31 days. (▼) Animal hemisphere; (▼▼) margins of the ciliary band; (cb) presumptive ciliary hand; (aoe) presumptive aboral ectoderm; (oe) presumptive oral ectoderm. Bar in a represents 10 μm.
20-hr hatched blastula (Fig. 2d). In the mesenchyme blastula (27 hr, Fig. 2e), the unlabeled region includes all primary mesenchyme cells within the blastocoel as well as the vegetal plate, which includes presumptive endoderm, secondary mesenchyme, and derivatives of the small micromeres. Cells of these lineages remain unlabeled throughout the remainder of embryonic development. The labeled region includes the majority of the surface of the blastula, which gives rise to both oral and aboral ectoderm. In many sections of blastulae, grain densities are highest at the animal pole and gradually diminish from animal to vegetal pole (Fig. 2e). The animal pole includes cells that produce the apical tuft, a structure of long rigid cilia first observed after hatching.

At gastrula stage, labeling is restricted again to ectodermal lineages, and grain densities over ectoderm are clearly heterogeneous. Figure 2, f and g, shows two sections, both cut through the animal–vegetal axis but perpendicular to each other, which illustrate that signals are highest in the animal hemisphere and in the region of the presumptive ciliary band. Signals are intermediate in presumptive aboral ectoderm and undetectable in portions of presumptive oral ectoderm, endoderm, and mesenchyme. Analysis of serial sections leads us to believe that the section shown in Figure 2g passes through the border of presumptive aboral and oral ectoderm. These are regions of contiguous cells, each comprising approximately one hemisphere of differentiating ectoderm (for example, see Lynn et al. 1983). The border between these regions contains cells of the presumptive ciliary band, and the uniformly high labeling intensities observed here over ectoderm are consistent with this interpretation.

The relative distribution of Spec3 mRNA established in various lineages at gastrula is maintained in the pluteus-stage larva. Grain densities observed in different sections vary somewhat, which appears to reflect actual differences in Spec3 mRNA content among different embryos, since quite similar levels of labeling are observed over corresponding regions of adjacent 5-μm sections (data not shown). Signal intensity over the perimeter of oral ectoderm corresponding to ciliary band cells is always equal to, and often significantly higher than, that over aboral ectoderm. Labeling of ciliary band cells increases at late pluteus stage. For example, Figure 2i shows a section cut close and parallel to the anal side of the pluteus in which a strip of oral ectoderm lying between the postoral arms is heavily labeled. Little or no signal is observed over a different cell type comprising the face of the pluteus with the border of the ciliary band (Fig. 2h). Thus, at these late stages strong signals correlate again with the presence of ciliated cells.

We have analyzed the spatial pattern of accumulation of β-tubulin mRNAs in embryos of two urchin species using probes containing coding sequence. In both *L. pictus* (B2 probe, Alexandraki and Ruderman 1983) and, as illustrated in Figure 3, in *S. purpuratus* ("blast 1" probe, Shepherd et al. 1983), the distribution of β-tubulin mRNA is very similar to that of Spec3 throughout embryogenesis. In particular, grain densities are lowest at the vegetal pole of the blastula and in the gastrula archenteron, intermediate in aboral ectoderm and its precursors, and highest at the animal pole of blastula and gastrula, especially in the presumptive ciliated band (20 hr: cf. Figs. 2d and 3a; 44 hr: cf. Figs. 2f and 3b). Figure 3c shows that tubulin mRNAs are present throughout the pluteus at rather uniform concentration except for two- to threefold higher concentration in the ciliated band (see arrows in Fig. 3c). With respect to ectodermal lineages, the patterns are identical to those of Spec3 mRNA (73 hr: cf. Figs. 2h and 3c). The only noticeable difference in *S. purpuratus* is that the tubulin probe gives some signal in endoderm and mesenchyme (see, e.g., Fig. 3c), because it cross-reacts with tubulin mRNAs that are not ectodermally restricted (Harlow and Nemer 1987a). The levels of signal obtained with Spec3 and β-tubulin probes are in close agreement with their relative abundances as determined by RNA blotting and solution titration (Nemer 1986; Harlow and Nemer 1987a).

**Accumulation of Spec3 message in deciliated and cilia-regenerating embryos**

The congruence of temporal and spatial patterns of Spec3 and β-tubulin mRNAs suggested that Spec3 protein might be involved in ciliogenesis. To test this possibility, we measured the accumulation of Spec3 mRNA during regeneration of cilia after deciliation induced by hypertonic shock (Burns 1973). This approach has been used recently by Harlow and Nemer (1987b) and Gong and Brandhorst (1987) to monitor β-tubulin mRNA levels after deciliation in *S. purpuratus* and *L. pictus*, respectively. Deciliation causes a transient increase in the synthesis of tubulin and other proteins associated with cilia (Stephens 1977) which, in the case of tubulin, has been shown to be regulated mainly at the level of transcription (Gong and Brandhorst 1987).Replicate dot blots were prepared with total RNA isolated from gastrula regenerating cilia. We compared the response of Spec3 with that of tubulin using a β-tubulin-coding sequence that hybridizes to all four known β-tubulin messages (Harlow and Nemer 1987a,b). As a control for nonspecific effects of deciliation, we used a probe for Spec1, an mRNA encoding a protein expressed in aboral ectoderm whose function is not expected to be related to ciliogenesis (Carpenter et al. 1984). The mitochondrial 16S rRNA probe was used to control for RNA loading and hybridization efficiency. Quantification of the results shows that both Spec3 (Fig. 4a) and β-tubulin messages (Fig. 4b) increase coordinately three- to fourfold in the first 90 min after deciliation, then decrease. A second round of deciliation produces a more rapid four- to fivefold accumulation of both messages with nearly identical kinetics. In contrast, Spec1 mRNA levels are unaffected by either round of deciliation. The decrease in Spec1 mRNA abundance over the 2-hr course of the experiment seen in both control and experimental cultures probably reflects the developmental decrease in message abundance seen for Spec1 at the late gastrula.
Figure 3. Distribution of β-tubulin mRNAs in sea urchin embryos. These in situ hybridizations were carried out in the same experiment as those shown in Fig. 2a, b, and d–i, using probes of the same specific activity. The sections in a, b, and c are from the same stages and correspond to the same sectioning planes as those shown in Fig. 2d (20 hr), 2f (44 hr), and 2h (73 hr), respectively. Signal intensities closely reflect relative concentrations of β-tubulin and Spec3 mRNAs since the concentration of the former is three times higher, the complexity of the tubulin probe is 75% of that for Spec3, and the exposure was one-half as long (14 days). Taking these parameters into account, the signals shown above should be 1.1 times those shown in Spec3 mRNAs in Fig. 2. Arrows indicate margins of the ciliary band. Bar in a represents 10 μm.

stage (Bruskin et al. 1981). We have performed identical experiments at blastula stage with similar results (data not shown; Eldon 1987), providing strong evidence that Spec3 is associated with ciliogenesis. Further strengthening this assertion are the results of Nemer (1986) and Harlow and Nemer (1987b), showing that animalization of embryos with Zn2+, which causes increased ciliation, results in increased levels of both Spec3 and β-tubulin mRNAs.

The Spec3 open reading frame

In our original report on Spec3 (Bruskin et al. 1981), we made use of a cDNA clone, pSpec3, which had a 1.15-kb insert and contained no significant open reading frames. With it, we isolated a longer Spec3 cDNA clone with a 1.95-kb insert only slightly smaller than the 2.0-kb Spec3 transcript (Fig. 5). The two EcoRI fragments from the longer Spec3 clone were subcloned into either pUC or M13 vectors and designated Spec3-kappa and Spec3-zeta (Fig. 5). Hybridizing single-stranded probes derived from Spec3-kappa to embryo RNA and sequencing Spec3-kappa defined the orientation of the Spec3 message and revealed an open reading frame beginning at the site of the EcoRI linker (added during the cloning procedure) and continuing 740 bases to the first translational stop codon, TAA.

To map the start of transcription, primer extension analysis was carried out. A synthetic primer complementary to the first 20 nucleotides of the putative protein coding sequence was hybridized to RNA isolated from blastula-stage embryos and extended using reverse transcriptase. As shown in Figure 6, the product of this reaction is a single fragment 152 bases long, 24 bases longer than the cDNA. Chain-termination sequencing of the genomic phage encoding the transcript was primed with the synthetic oligomer and run on the same denaturing acrylamide gel. The alignment of the extension product with the sequencing lanes reveals the nucleotide initiating transcription to be G, as indicated in Figure 7a. The first methionine in the only open reading frame occurs at position 133 and is assumed to be the start of translation of the Spec3 protein. The protein translated from this reading frame would be 208 amino acids long.

Several features of the predicted amino acid composition (Table 1) and sequence (Fig. 7a) are unusual. Acidic and basic amino acids are very rare; there are only five aspartic acids, three lysines, three histidines, and one arginine; glutamic acid is absent. Most of the nonpolar amino acids are arranged in a highly ordered pattern in the carboxy-terminal half of the protein (see Fig. 7b). The proline content is extremely high (14%) and is distributed asymmetrically in the protein sequence; all 29 prolines occur within the amino-terminal 138 residues, and 23 prolines occur within the first 63 residues. The sequence is highly reiterated: in 19 places a single amino acid is repeated twice, in eight places a single amino acid is repeated three times, and in three places a single amino acid is repeated four times. In addition, a six-amino-acid sequence is repeated three times, and an overlapping region of 15 amino acids is repeated twice. The placement of these reiterations is detailed in the legend to Figure 7a. There is also a single site for amino-linked glycosylation (Asn-Asn-Thr) and a single Lys-Lys
pair, which could act as a peptide cleavage site (Fig. 7a).

A Kyte-Doolittle hydrophathy plot (Kyte and Doolittle 1982) of the putative Spec3 protein displays several distinctive features (Fig. 7b). Despite high overall hydrophobicity there is no amino-terminal signal peptide. Rather, near the amino-terminus is a long hydrophilic region (residues 10–70). In contrast, the carboxy-terminal half of the molecule contains two strongly hydrophobic regions, spanning residues 105–136 and residues 146–183, bounded by short stretches of hydrophilic residues. Nine of the 12 charged amino acids found in the protein sequence occur in these hydrophilic stretches (small arrowheads, Fig. 7b). The nonrandom distribution of acidic and basic residues and the sharp boundaries between hydrophilic and hydrophobic domains argue that these regions have structural significance.

**Figure 4.** Quantity of Spec3, β-tubulin, and Specl RNA synthesized following deciliation of gastrula-stage embryos. (a) Spec3; (b) β-tubulin; (c) Spec1. ■ First deciliation; ● second deciliation; □ first control; ○ second control.

**Figure 5.** Restriction maps of Spec3 cDNA clones. The upper map is the nearly full-length clone isolated from a λgt10 library. The bars indicate the two EcoRI fragments, kappa and zeta, that were subcloned. The open reading frame is indicated by the box, the dark portion of which indicates the putative protein coding region. The lower map is the original Spec3 isolate (Bruskin et al. 1981). Restriction sites: [A] XbaI; [B] BamHI; [C] HindIII; [F] HindIII; [K] KpnI; [P] PvuII; [R] EcoRI; [T] PstI.

**A single Spec3 gene**

To determine the number of genes contributing to the complex temporal and spatial patterns of Spec3 expression, we used the 1.34-kb EcoRI–BamHI fragment of the Spec3-kappa subclone (Fig. 5) to isolate the number of recombinant λ phages from a λEMBL3 genomic DNA library made from a Sau3A partial digestion of S. purpuratus sperm DNA. All of the recombinant inserts derived from a single overlapping region of the genome as judged by their restriction maps. Southern hybridization analysis of one genomic recombinant, phage 39, using the Spec3-kappa and -zeta cDNA clones as probes, defined restriction fragments containing Spec3 sequences. These restriction fragments were partially sequenced to determine their similarities with the Spec3 cDNA clones and to identify the exon–intron boundaries. The results of this analysis (summarized in Fig. 8) indicate that phage 39 contains an entire Spec3 gene and that, where sequences could be compared, the match with the cloned cDNA is greater than 95% (data not shown; Eldon 1988). The Spec3 gene is 4 kb long and is composed of 3 exons, 0.51 kb, 0.18 kb, and approximately 1.4 kb long. The introns are clearly bounded by consensus splice sequences and are 1.4 kb and 0.32 kb in length (Fig. 8). The message size as measured by RNA gel blots corresponds closely to the sum of the exon sizes.

Though the genomic library screen yielded only one region of the S. purpuratus genome and sequencing showed a high degree of match with the cDNA clones, it was still possible that other Spec3 genes existed that were not represented in the λEMBL3 library. That this is not the case is shown by gel blots of sperm DNA from
Table 1. Amino acid composition of the Spec3 translation product

| Amino acid | Number | Percent |
|------------|--------|---------|
| Nonpolar   |        |         |
| Ala        | 22     | 10.6    |
| Val        | 11     | 5.3     |
| Leu        | 8      | 3.8     |
| Ile        | 18     | 8.7     |
| Pro        | 29     | 13.9    |
| Met        | 7      | 3.7     |
| Phe        | 10     | 4.8     |
| Trp        | 5      | 2.4     |
| Gly        | 28     | 13.5    |
| Ser        | 4      | 1.9     |
| Thr        | 12     | 5.8     |
| Cys        | 6      | 3.4     |
| Tyr        | 7      | 3.4     |
| Asn        | 14     | 6.7     |
| Gln        | 15     | 7.2     |
| Polar      |        | 41.9    |
| Acidic     |        |         |
| Asp        | 5      | 2.4     |
| Glu        | 0      | 0.0     |
| Basic      |        | 2.4     |
| Lys        | 3      | 1.4     |
| Arg        | 1      | 0.5     |
| His        | 3      | 1.4     |

several individuals. In the experiment shown in Figure 9, DNA from four individuals was digested with EcoRI and XbaI and the blots were probed with the 1.0-kb EcoRI-XbaI fragment of the Spec3-kappa clone [cf. Fig. 5]. Also included in this experiment was an EcoRI, XbaI digestion of phage 39 [Fig. 9, lane 4]. As predicted from the restriction map, the probe hybridized to two phase 39 fragments, a 3.4-kb fragment containing exon 1 and a 1.6-kb fragment containing exon 2 and part of exon 3. Individual 3 [Fig. 9, lane 2] shows the identical hybridization pattern as phage 39, indicating that under these hybridization conditions phage 39 represents the only Spec3 gene in this apparently homozygous animal. Individual 2 [Fig. 9, lane 1] shows a composite pattern: Two bands match the phage 39 pattern, and two bands at 4.1 kb and 1.8 kb appear in the same intensity ratio as the 3.4- and 1.6-kb bands. Presumably, this individual is heterozygous for the Spec3 EcoRI-XbaI fragments due to restriction fragment length polymorphisms at the Spec3 locus. DNA of individual 11 [Fig. 9, lane 4] was used to construct the EMBL3 genomic library; this individual is also heterozygous, with phage 39 representing one of the alleles. DNA from individual 7 [Fig. 9, lane 3] shows four bands, one that is novel and three that correspond to bands seen in individuals 2, 3, and 11. No additional bands appear when the stringency of the final wash is reduced to 0.15 M Na+, 55°C [data not shown]. It is clear from the genomic DNA blot analysis that the Spec3 gene is single copy and polymorphic with respect to the length of the EcoRI-XbaI fragment. Direct comparisons between different cDNA clones and genomic clones show minor variation in sequence, demonstrating that single base pair polymorphisms also occur throughout the length of the gene [data not shown; Eldon 1988].

Discussion

Correlation of the Spec3 message with ciliogenesis

Our results provide strong circumstantial evidence that the Spec3 message encodes a protein associated with ectodermal ciliogenesis, though it may have other tu-
Gene expression in embryonic ectoderm

...as well. The temporal pattern of accumulation of Spec3 mRNA is quite similar to that of the ectoderm-enriched β-tubulin mRNAs previously identified [Nemer 1986; Harlow and Nemer 1987a]. The rapid accumulation of these transcripts begins at late cleavage stage and coincides roughly with the initial elaboration of cilia; the second peak of accumulation coincides with the appearance of the ciliary band on the...
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Figure 8. Spec3 gene structure. A partial restriction map of 11 kb at the 5' end of the phage 39 insert is shown. A portion of this region encodes the Spec3 transcript. The boxes indicate the three exons of the mature mRNA transcript connected by lines indicating the two introns. Shaded areas indicate the presumed protein coding region. The bars below the map indicate the two EcoRI-XbaI fragments of genomic DNA detected by the probe used for genomic DNA blots (Fig. 9). Restriction sites are: (A) XbaI; (B) BamHI; (G) BglII; (H) HindIII; (K) KpnI; (P) PvuII; (R) EcoRI; (T) PstI; (X) XhoI.

Figure 9. Representation of the Spec3 gene in the genome. A Southern blot of EcoRI–XbaI digests of DNA from the genomes of four individuals was hybridized with the EcoRI–XbaI fragment of the Spec3-kappa cDNA subcloned into M13mp19. (Lanes 1–4) Individuals 2, 3, 7, and 11, respectively; (lane 5) phage 39 digested with EcoRI–XbaI (Fig. 8). Fragment sizes are indicated. The stronger hybridization intensity of the 1.2- to 2.0-kb bands are due to asymmetric labeling of the probe because the phage 39 bands (lane 5) show the same asymmetric hybridization pattern.

prism-stage embryo. In situ hybridization shows highest accumulation of Spec3 and β-tubulin mRNAs in regions thought to be actively involved in producing cilia. The gradation in grain intensities seen with the Spec3 and β-tubulin probes at blastula stages reflects the apparent length gradient of cilia on the embryo. Cilia, including the apical tuft cilia, have been found to be longest at the animal pole and shortest at the vegetal pole (Riederer-Henderson and Rosenbaum 1979; Harlow and Nemer 1987b). High grain densities are clearly associated with the ciliary band, a distinct cluster of ciliated columnar or pear-shaped cells that extends around the circumference of the oral surface of prism- and pluteus-stage embryos. Deciliation and subsequent cilia regeneration results in a transient four- to fivefold increase in the levels of the Spec3 and β-tubulin mRNAs, but not of Spec1 mRNA, whose expression is probably unrelated to ciliogenesis. In addition, embryos that have been animalized and induced to grow excessive cilia by treatment with Zn²⁺ synthesize increased levels of both Spec3 and β-tubulin mRNAs (Nemer 1986; Harlow and Nemer 1987b). Stephens (1977) has shown that the synthesis of several ciliary proteins is increased during cilia regeneration, and several studies demonstrate that β-tubulin mRNA levels are increased during cilia regeneration [Merlino et al. 1978; Gong and Brandhorst 1987; Harlow and Nemer 1987b]. In Chlamydomonas, regeneration of flagella results in a similar reduction of α- and β-tubulin mRNAs and mRNAs encoding other cilia-associated proteins (Lefebvre et al. 1980; Baker et al. 1984; Schloss et al. 1984; Williams et al. 1986). In the case of α- and β-tubulin, this induction involves new transcription as well as increased stability of the mRNA (Baker et al. 1984).

In S. purpuratus, cilia appear during the early blastula stages on the apical surface of all blastomeres, with the possible exception of mesenchyme cell precursors. Our in situ hybridization data show no grains above background in differentiated endoderm cells, yet at least some of these cells do have cilia. The cells of the mid- and hindgut are ciliated on the luminal side, consistent with the presence of cilia on their apical surface before invagination, though it is not clear whether or not those of the foregut are ciliated [G. Wessel, pers. comm.]. In L. variegatus, cilia on invaginating endodermal cells are shorter and stubbier than ectodermal cilia, suggesting structural differences [Morrill and Santos 1985]. Because we detect no Spec3 message in endoderm tissue, we assume that the Spec3 protein is not associated with endodermal cilia.

The quantity of the Spec3 transcript can be roughly estimated by comparison with β-tubulin message, which has been quantified by Harlow and Nemer [1987a]. They found that there are 3.5 × 10⁶ β-tubulin mRNA molecules per embryo at hatching blastula stage, and that about one-third of these are the ectoderm-specific β₁-tubulin mRNA. The abundance of Spec3 transcripts is also one-third that of total β-tubulin mRNA at this stage [Nemer 1986]. This correspondence suggests a developmental coupling of the Spec3 and β₁-tubulin mRNAs that is quantitative as well as qualitative. If the translation rate and stability of the Spec3 protein were similar to that of β₁-tubulin, the new embryonic Spec3 protein would be quite abundant and possibly stoichiometric with β₁-tubulin. Whether or not there are maternal stockpiles of Spec3 protein similar to the large quantities of tubulin subunits stored in the egg is unknown, so the relative abundance of tubulin and Spec3 protein in the embryo cannot yet be determined.
The fact that the Spec3 message accumulates to one-third the level of total β-tubulin message during development suggests that the Spec3 product is not a minor cellular protein. It may be that the Spec3 protein is not specific to cilia but is also associated with other microtubule structures such as the mitotic spindle or the cytoskeleton, much as the testes-specific β-tubulin in *Drosophila* is found in the meiotic spindles and cytoplasmic microtubules as well as in the sperm axoneme (Kemphues et al. 1982).

**Temporal and spatial expression of Spec3 mRNA**

The Spec3 mRNA is one of the earliest known developmentally regulated transcripts, accumulating in embryos as early as 12 hr after fertilization. The initial accumulation clearly takes place in all blastomeres, and the pattern of differential expression is established by sequential disappearance of Spec3 mRNA from different lineages. The transient decrease in Spec3 mRNA content between blastula and gastrula stages reflects both its loss from some lineages and its decreased levels in others. The mRNA disappears from endoderm and mesenchyme cells after 12 and probably closer to 17 hr, at about the time when cell cycle synchrony breaks down and cell division slows. The rapid disappearance of Spec3 mRNA implies a short half-life in these cells. Thus, the expression and decay of the Spec3 message is one of the earliest events of gene regulation identified for mesenchyme and endoderm cells. Spec3 mRNA is present in the entire ectoderm at high concentrations in the early blastula stages and persists at diminishing concentrations until pluteus. This implies either that there is continuing synthesis of Spec3 mRNA at low levels or that the mRNA synthesized earlier is rather stable later in development.

**The Spec3 protein**

We used the Bionet Computer Resource to search for protein sequences with similarities to the Spec3 open reading frame and found no significant matches. This is not surprising even if the Spec3 protein is a component of the cilium per se. This structure contains a complex assortment of more than 150 proteins (Piperno et al. 1977), and although several of the accessory proteins of cilia and flagella axonemes have been identified, including dynein arm components (Mitchell and Rosenbaum 1985) and radial spoke proteins (Piperno et al. 1981), there is little sequence information available for any of them. Recently cDNA clones specifying different radial spoke proteins and dynein from *Chlamydomonas* have been isolated (Williams et al. 1986), and the messages are all induced to varying extents upon deflagellation and regeneration. Whether the Spec3 protein has sequence similarities with any of these proteins remains to be seen.

We have been unable to identify the Spec3 proteins in vivo using several approaches [Eldon 1988]. Brandhorst and his colleagues have performed an extensive analysis of the temporal and spatial patterns of protein synthesis in *S. purpuratus* using two-dimensional polyacrylamide gels [Bruskin et al. 1982; Bedard and Brandhorst 1983], and close inspection of these gels shows no protein with the early temporal appearance and ectodermal specificity expected for the Spec3 protein. Given the long hydrophobic stretches and high proline content present in the Spec3 protein, it is possible that it has unusual physical properties. It is also possible that the Spec3 protein is modified post-translationally by proteolytic processing, glycosylation, or other modifications, and is not resolvable on two-dimensional gels.

The sequence and hydrophobicity analysis of the Spec3 open reading frame indicates that the putative protein has two distinct regions. The amino-terminal portion [residues 1–91] is hydrophilic, contains 26 of the 29 prolines, and has a site for amino-linked glycosylation. The carboxy-terminal portion [residues 92–208] is highly hydrophobic and contains two long hydrophobic stretches bounded by short hydrophilic linkers, where nine of the 12 charged amino acids reside. The presence of the hydrophobic regions suggests to us that the Spec3 protein may be associated with the plasma membrane of cilia via two potential α-helical membrane-spanning regions. The Spec3 translation product appears to lack an amino-terminal signal sequence, but several membrane proteins have been identified that use regions other than their amino terminus for membrane insertion [Wickner and Lodish 1985]. It is possible that the extensive hydrophobic regions found in the Spec3 translation product allow its incorporation into other hydrophobic environments within the cell, such as nonmembranous lipids or other proteins of a hydrophobic nature.

**The Spec3 gene**

The complex temporal and spatial expression of the Spec3 mRNA was monitored with probes containing Spec3 coding sequences. It is conceivable that this complexity could be the result of cross-hybridization of multiple Spec3 genes, each with its own pattern of expression. For example, the observation that the Spec3 transcripts are present in all cells at 12 hr after fertilization but restricted to ectodermal cells by 17 hr after fertilization could be explained by the existence of two Spec3 genes: one expressed initially in all cells and ceasing expression by 17 hr postfertilization, and the other expressed only in ectodermal cells beginning at about 17 hr after fertilization. Differential expression of individual members of multiple gene families has been reported in sea urchin embryos for genes encoding actins [Cox et al. 1986; Lee et al. 1986], histones [Mohun et al. 1985], β-tubulins [Alexandraki and Ruderman 1985; Harlow and Nemer 1987], and metallothioneins [Wilkinson and Nemer 1987]. However, our data demonstrate that multiple Spec3 genes do not exist and that expression from a single Spec3 gene is responsible for the patterns we observe. We isolated only a single region of the *S. purpuratus* genome that contained Spec3 sequences, and hybridization of Spec3 coding sequences with genomic DNA, even under low stringencies, yielded patterns indicating a single polymorphic gene.

Rather than representing a gene whose expression is
associated with the differentiation of a distinct cell lineage of cell type, as is the case for several genes recently described in sea urchins (Davidson 1986), expression of the Spec3 appears, at least in part, to be associated with developmental and physiological process—that of ectodermal ciliogenesis. The strong coupling of the Spec3 gene with the β-tubulin genes suggests that identical regulatory mechanisms control their expression. In particular, the stability of β-tubulin mRNA in most systems is controlled by the pool of β-tubulin monomer (Cleveland et al. 1981). It will be of interest to ask whether the Spec3 message stability is under similar control.

Methods

Embryo culture

Embryos were cultured at a density of 10^3–10^4/ml in artificial seawater (McClay 1986) at 14–15°C with constant stirring.

Hybridization probes from single-stranded M13 templates

Single-stranded M13 templates were hybridized to a 16-base hybridization probe primer [New England Biolabs] and elongated using the Klenow fragment of *E. coli* polymerase I (BRL) in the presence of a mix of dATP, dCTP, TTP (Pharmacia), and [a-32P]dCTP (Amersham) at 30°C for 30 min. Reaction products were precipitated with ethanol and washed to remove the bulk of the unincorporated nucleotides. Specific activities of 25 × 10^6 to 45 × 10^6 cpm/μg were obtained routinely. The probes labeled were: *Spec1-M13mp19* containing the 1.05-kb Styl–XbaI fragment of Spec1 cDNA (Carpenter 1984), used as a negative control in deciliation experiments; *Spec1-M13mp19* containing the 1.03-kb *EcoRI–XbaI* fragment of the kapa cDNA clone (Fig. 5), and *mitochondrial 16S rRNA-M13mp8* containing a 550-bp *EcoRI* fragment of the *S. purpuratus* mitochondrial 16S rRNA (Wells et al. 1982), used to standardize the amount of hybridizable RNA on the filters.

Random oligonucleotide-primer labeling of β-tubulin cDNA

The method of Feinberg and Vogelstein (1983) was used to label a gel-purified 365-bp BamHI–BglII fragment from the coding region of the β-tubulin cDNA clone 1*-pUC19-365S* [a gift from P. Harlow and M. Nemer] to a specific activity of 1.6 × 10^8 cpm/μg. Labeled fragments were ethanol-precipitated and washed to remove the bulk of the unincorporated nucleotides.

RNA accumulation during normal development

RNA was isolated from eggs and embryos every 4 hr from 3 to 59 hr and every 6 hr from 65 to 107 hr from synchronously growing cultures of *S. purpuratus* embryos by the guanidine-HCl method described by Bruskin et al. (1981). Ten-microgram samples were electrophoresed on formaldehyde agarose gels (Bruskin et al. 1981) and blotted to nitrocellulose. Blots were hybridized with an M13 single-stranded probe of the *EcoRI–XbaI* fragment of the Spec3 cDNA probe that included the entire open reading frame and 108 bp of 5' untranslated sequence as described (Bruskin et al. 1981). Final wash conditions were 68°C in 0.066 M NaCl for 1–2 hr, unless otherwise indicated. Signals were quantified by densitometric scanning of several autoradiographic exposures of the RNA blot to insure linearity. To standardize the amount of hybridizable RNA in each lane, the filter was probed with the 550-bp *EcoRI* fragment of the mitochondrial 16S rRNA sequence described above. Accumulation was plotted as a proportion of maximal level reached during development.

In situ hybridization

A 1.74-kb Spec3 cDNA fragment bounded by *EcoRI* restriction sites (Fig. 5) was inserted into two different vectors, each containing an Sp6 promoter. The insert in *pSpZ1* (Angerer et al. 1985) contains two of the Spec3 sequences in the same orientation, whereas, that in *pSp65* is a monomer. Run-off transcripts [3900 nucleotides (pSpZ1) or 1790 nucleotides (pSp65)] were synthesized from HindIII-digested templates as described previously (Lynn et al. 1983). Sister constructions provided comparable sense strand transcripts for use as controls for nonspecific background binding of probe. Similarly, a 1.3-kb *PstI* cDNA fragment containing β-tubulin sequence [blast 1, kindly provided by M. Nemer] was transferred to *pSp64* and *pSp65* for preparation of sense and antisense RNA probes with 36 RNA polymerase. The specific activity of the run-off transcripts was adjusted to 0.91 × 10^8 to 1.1 × 10^8 dpm/μg using [3H]CTP and [3H]UTP (Amersham). Methods for purification and reduction of the fragment length of probes to 150–200 bases, for glutaraldehyde fixation and prehybridization treatments of tissue and for in situ hybridization have been described previously (Angerer and Angerer 1981, Cox et al. 1984). Hybridizations were done at 45°C using saturating probe concentrations of 0.5–1 μg/ml, and autoradiographic exposure times were 28–31 days.

RNA accumulation following deciliation

Gastrula-stage embryos were deciliated by stirring 2 min in hypertonic seawater, following the protocol of Stephens (1986). Following deciliation, the embryos were washed once in artificial seawater and cultured at 14°C with constant stirring. A portion of the culture was harvested every 15 min for 2 hr and RNA was isolated by rapid lysis in a buffer containing 50 mM HEPES (pH 8.0), 0.5 mM sodium acetate (pH 6.0), 2% sodium deoxycholate, and 1% deuterium carbonate. The preparation was extracted once with phenol and once with chloroform and then precipitated with ethanol. The resulting pellet was resuspended in 3.0 mM sodium acetate (pH 6.0), and RNA was allowed to precipitate at −20°C overnight (Palmiter 1974). Equal amounts of total RNA were denatured for electrophoresis and loaded onto nylon membrane filters (Hybond-N, Amersham) using a dot-blot apparatus (BioRad). Quadruplicate filters were made and hybridized as described in Results, using the conditions described above for filter hybridization. The amount of probe hybridized to each filter was determined in a scintillation counter. The amount of total RNA on each dot was determined using the mitochondrial 16S rRNA probe as a standard. Accumulation was plotted as a fraction of the maximum level reached over the course of the experiment.

Mapping the initiation of transcription

One nanogram of a synthetic 20-base oligonucleotide complementary to the first 20 nucleotides of the putative protein coding sequence was labeled at the 5' end using polynucleotide kinase and hybridized with 50 μg of total blastula RNA for 2 hr at 15°C below the calculated *Tm*. The oligonucleotide primer was extended using AMV reverse transcriptase (Life Sciences) in the presence of 8 μM dNTPs. To determine the sequence of the extended product and identify the nucleotide with which transcription initiates, the synthetic 20-base oligonucleotide was used to prime chain-termination sequencing reactions from the appropriate genomic template. The products of the extension and sequencing reactions were resolved on the same 6% denaturing acrylamide gel cast in a buffer gradient.
Isolation of genomic clones

A λEML3 library, made with one male sea urchin’s genomic DNA partially digested with Sau3A, was screened with the 1.4-kb EcoRI–BamHI fragment of the Spec3-kappa cDNA clone, which had been radioactively labeled by nick translation. Phages producing positive plaques were purified, and DNA was isolated using standard procedures outlined by Maniatis et al. (1982).

DNA sequencing reactions

Restriction fragments labeled at a unique end were chemically degraded by the method of Maxam and Gilbert (1980) as modified by Bencini et al. (1984). Reaction products were electrophoresed on 8% and 20% denaturing polyacrylamide gels (Sanger and Coulson 1978).

Fragments subcloned into M13 were sequenced by the chain-termination method of Sanger et al. (1977). The chain-termination method as modified by Chen and Seeburg (1985) was employed to sequence double-stranded plasmids. Reaction products were separated on denaturing polyacrylamide gels cast in a buffer gradient (Biggin et al. 1983).

Genomic DNA blots

Ten micrograms of DNA isolated from the sperm of a single urchin was digested with restriction enzymes and electrophoresed on 0.8% neutral agarose gels and blotted onto nitrocellulose (Southern 1975). Blots were hybridized as described by Bruskin et al. (1981). Final wash conditions were 0.06 M Na+ at 68°C, unless otherwise indicated.

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Note added in proof

The blast 11 β-tubulin clone has been sequenced recently by P. Harlow, S. Litwan, and M. Nemer [J. Mol. Evol. 26: in press]. It contains open reading frame sequences, which hybridize to all β-tubulin messages and 3′ untranslated sequences, which hybridize only to β1-tubulin transcripts.

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