## NOTES

## Molecular and Developmental Characterization of the Heat Shock Cognate 4 Gene of *Drosophila melanogaster*

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The Drosophila heat shock cognate gene 4 (hsc4), a member of the hsp70 gene family, encodes an abundant protein, hsc70, that is more similar to the constitutively expressed human protein than the Drosophila heat-inducible hsp70. Developmental expression revealed that hsc4 transcripts are enriched in cells active in endocytosis and those undergoing rapid growth and changes in shape.

The cells of nearly all organisms have a conserved response to environmental stresses, consisting of the synthesis of several heat shock proteins (hsps) (for review, see reference 15). The most prominent stress protein in the majority of species, hsp70, is highly conserved among procaryotic and eucaryotic species. The hsp70 gene is a member of a family of closely related genes that includes both stressinducible genes (hsp's) and genes expressed constitutively during normal development, the heat shock cognate genes (hsc's).

The heat shock cognate proteins appear to be important for normal cellular function (reviewed in references 5, 15, 21, and 27). A number of recent results indicate that the hsp70like proteins act in an ATP-dependent manner in several cellular compartments. They may function to alter the conformations of proteins or affect protein-protein interactions (9, 18, 25). Additionally, they may play a role in the translocation of polypeptides across specific membranes (4, 7). The abundant cytoplasmic heat shock cognate protein in mammalian cells, hsc70, is involved in the ATP-dependent uncoating of clathrin from endocytotic vesicles (26, 31).

In Drosophila melanogaster, the hsp70 multigene family includes five copies of the heat-inducible hsp70 gene, one copy of the heat-inducible hsp68 gene, and seven heat shock cognate genes, hsc1 through hsc7, that are expressed during normal growth (6, 13, 15, 19). The Drosophila hsc70 protein, encoded by hsc4, is a very abundant polypeptide in all tissues and cells and is localized to a meshwork of cytoplasmic fibers concentrated around the nucleus (19).

Sequence and structure of the Drosophila hsc4 gene. The hsc4 gene of D. melanogaster was originally isolated on a recombinant plasmid, pMG34 (Fig. 1A and 2), following hybridization with a Drosophila hsp70 gene (6). The DNA sequence of hsc4 revealed a single open reading frame of 1,953 base pairs (bp) that potentially encodes a 651-amino-acid polypeptide with an estimated molecular weight of 71,108. S1 nuclease analysis indicated that the protein coding and the 5' untranslated regions were not contiguous and

that an intron was located 5' of the initiating ATG (data not shown). To confirm the position of this intron, a cDNA, cD12, was isolated and the sequence of the 5' end was determined. The cDNA sequence diverged from the genomic DNA sequence at the ATG (Fig. 1B). The protein coding and 5' untranslated regions of the *hsc4* gene were interrupted by a 1.6-kilobase (kb) intron.

The deduced amino acid sequence of the hsc4 gene (Fig. 3) was 73% identical to that of the *Drosophila* heat-inducible hsp70 (12) and 85% identical to that of the human hsc70 polypeptides (8). Furthermore, *Drosophila* hsc70 was 80% identical to *Caenorhabditis elegans* hsp70A, which is abundant throughout development and only marginally heat inducible (29). An unresolved question is whether constitutively expressed *hsp70*-related genes, such as *hsc4*, and those induced by stress, e.g., *hsp70*, have identical or different functions. The fact that *Drosophila* hsc4 is more closely related to vertebrate *hsc70*-like genes than an inducible gene from *D. melanogaster* suggests that the constitutively expressed proteins may be functionally distinct from the stress-induced proteins.

In situ hybridization to embryos reveals stage- and tissuespecific enrichment of hsc4 transcripts. Northern (RNA blot) analysis demonstrates that the major 2.3-kb hsc4 transcript is expressed throughout embryonic, larval, pupal, and adult development at relatively constant levels (6) (data not shown). In situ hybridization to wild-type embryos was performed as described by Hafen and Levine (11) or Tautz and Pfeifle (30). Radioactive DNA probes were labeled by nick translation with [<sup>35</sup>S]dCTP (New England Nuclear Corp.) to a specific activity of approximately  $5 \times 10^7$ cpm/µg, and the autoradiograms were developed after 2 to 3 days. Nonradioactive probes were prepared essentially by the protocol provided with the nonradioactive labeling and detecting kit (Boehringer Mannheim, catalog no. 1093657). hsc4 transcripts were localized in a complex spatial and temporal pattern during embryogenesis (Fig. 4 and 5), which was superimposed onto a basal level of expression apparent in virtually all cells of the developing embryo.

Enrichment of *hsc4* transcripts was first observed during late syncytial and cellular blastoderm stages and during early gastrulation in the cytoplasmic compartment between the

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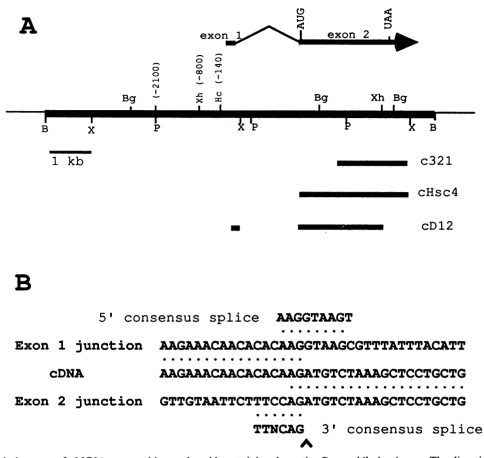


FIG. 1. (A) Restriction map of pMG34, a recombinant plasmid containing the entire *Drosophila hsc4* gene. The direction of transcription, positions of the intron and exons 1 and 2, start (ATG) and stop (TAA) codons of translation, and relevant restriction sites are included (B, *Bam*HI; Bg, *Bg*/II; Hc, *Hinc*II; P, *Pst*I; Xb, *Xba*I; Xh, *Xho*I). Below the restriction map, the approximate extents of three *hsc4* cDNAs are depicted. cDNA clone c321 was isolated from a 3- to 12-h embryonic cDNA library (23) in a differential screen designed to identify genes preferentially expressed in neuroblasts rather than differentiated neurons (L. A. Perkins, A. P. Mahowald, and N. Perrimon, submitted for publication) and was determined to encode *hsc4* sequence based on its localization to 88E on the salivary gland polytene chromosomes, Southern blot analysis with pMG34 as a probe, and partial DNA sequence analysis. cDNA clone cHsc4 was isolated at high stringency from a size-selected 9- to 12-h embryonic library with c321 as a probe (35). cDNA clone cD12 was isolated from an embryonic cDNA library provided by M. Goldschmidt-Clermont with pMG34 as a probe. (B) Location of the intron/exon boundaries in the *Drosophila hsc4* gene. This comparison shows the nucleotide sequence from cDNA cD12 aligned with the genomic DNA sequence from pMG34. This alignment does not permit the unambiguous determination of the precise boundaries of exons 1 and 2, but the predicted splice site (  $\bigstar$  ), based on the eucaryotic consensus (17), is shown.

blastoderm nuclei and the volk (Fig. 4A and B). These stages are characterized by the rapid assembly of cellular membranes to compartmentalize the nuclei. Tissue enrichment was next observed in neuroblasts of both the head and extending germ band (Fig. 4C, D, and E). Unlike transcripts from *Delta* and members of the *achaete-scute* gene complex, which are enriched in subsets of neuroblasts enlarging within the neurogenic ectoderm (2, 24, 32), hsc4 transcripts were only observed in newly segregated neuroblasts internal to the ectoderm. Enrichment was clearly observed in neuroblasts from the procephalic neurogenic ectoderm (Fig. 4C and D) and continued to be enriched in specific derivatives of this region (Fig. 4H). Enrichment of hsc4 transcripts was observed in cells of the embryonic gut from anterior and posterior midgut invagination to hatching (Fig. 4F to H) and transiently in developing mesodermal cells (Fig. 4F and G). Enrichment in the gut occurred while the cells were undergoing numerous cellular processes: mitoses, expansions, stretching, and volumetric growth (3). Enrichment in the mesoderm occurred as the somatic and splanchnic mesoderms became separate layers (Fig. 4F and G) and was readily apparent as the somatic muscles formed and single cells fused into syncytial myotubes and differentiated into somatic muscles (3).

During late embryogenesis, *hsc4* transcripts were most abundant in the garland gland (Fig. 4G and H), an organ postulated to segregate and store waste products (34). Cells from the garland gland are very active in endocytosis via coated vesicles. In fact, electron microscopy reveals the cortex of these cells to be a labyrinth of endocytotic pits or channels that "pinch off" to form clathrin coated vesicles (14; C. Poodry, personal communication). Since a clathrin "uncoating ATPase" activity has been detected in *Drosophila* cells (28), we propose that hsc70 in the garland gland functions in the uncoating of clathrin triskelions.

In conclusion, *hsc4* transcripts are present in most if not all cells during embryonic development but are enriched in cells active in endocytosis and those undergoing rapid growth and changes in shape. Studies in other organisms have demonstrated high levels of hsc70 in rapidly growing

- CCGAG CGCCAMAAAA TACCACGATC AATAAGAACT GCACTGTTGT TAAATGGCTG GGCAGCCGTG TGCGTCAAAT AAGTGCCGAT GGAGAACTAG AATAACCTTA -343
- TATAACGAAA GATTTATAAA TAAAAAAAATC CCATGTTCCA TATTCCACTG TTCCTTCATT AATTATTCCT ATATTATGAA TTATATTCAT TAAGAAAGGA ATAGGAAATG TGGTTTTTAA -223
- ATGTCTAAAG CTCCTGCTGT TGGTATTGAT TTGGGCACCA CCTACTCGTG CGTGGGCGTG TTCCAGCATG GCAAGGTCGA GATCATCGCC AACGACCAGG GTAATCGTAC CACTCCATCC 231 TATGTTGCCT TCACCGATAC GGAGCGTCTG ATCGGAGATG CCGCCAAGAA CCAGGTGGCG ATGAACCCGA CCCAGACGAT CTTCGACGCC AAGCGCTTGA TTGGTCGCAA GTTCGATGAT 351 GCCGCCGTGC AGTCTGACAT GAAGCACTGG CCCTTCGAGG TGGTCAGCGC CGATGGCAAG CCCAAGATCG AGGTGACCTA CAAGGACGAG AAGAAGACCT TCTTCCCCGA GGAGATCTCT 471 TCGATGGTGC TTACCAAGAT GAAGGAGACC GCCGAGGCCT ATCTGGGCAA GACTGTGACC AACGCGGTCA TCACCGTGCC GGCCTACTTC AACGACTCTC AGCGTCAGGC GACCAAGGAC 591 GCCGGCACCA TCGCCGGTCC GAACGTGCCG CGTATCATCA ACGAGCCCAC TGCCGCTGCT ATCGCTTACG GTCTGGACAA GAAGGCTGTT GGAGAGCGCA ACGTGCTCAT CTTCGATCTG 711 GECGGCGGCA CCTTCGATGT GTCCATCCTG TCGATCGATG ACGGTATCTT TGAGGTCAAG TCCACGGCCG GAGATACGCA TCTGGGTGGT GAGGACTTCG ACAACCGTCT GGTCACCCAC 831 TTCGTGCAGG AGTTCAAGCG CAAGCACAAG AAGGATCTGA CCACCAACAA GCGTGCTCTG CGTCGTCTGC GCACCGCTTG CGAGCGTGCA AAGCGTACCC TGTCGTCCTC CACCCAGGCC 951 AGCATTGAGA TCGACTCTCT GTTCGAGGGT ACCGACTTCT ACACCTCGAT TACTCGTGCC CGTTTCGAGG AGTTGAACGC TGATCTGTTC CGCAGCACCA TGGACCCCGT GGAGAAGGCT 1071 CTGCGTGACG CCAAGCTGGA CAAGTCGGTC ATCCACGACA TTGTGCTGGT CGGTGGCTCC ACCCGTATCC CCAAGGTGCA GCGCCTGCTG CAGGATCTGT TCAATGGCAA GGAGCTGAAC 1191 AAGTCGATCA ATCCCGATGA GGCTGTGGCC TACGGTGCTG CCGTCCAGGC GGCCATTCTG CACGGCGACA AGTCGCAGGA GGTGCAGGAT CTGCTGCTGC TCGATGTCAC TCCTCTGTCC 1311 CTGGGTATCG AAACCGCTGG CGGTGTGATG AGCGTGTTGA TCAAGCGCAA CACCACCATT CCGACCAAGC AGACCCAGAC CTTCACCACC TACTCGGACA ACCAGCCCGG TGTGCTGATC 1431 CAGGTGTACG AGGGAGAGCG TGCCATGACC AAGGACAACA ACCTGCTCGG CAAGTTCGAG CTGTCGGGCA TCCCCCCGC ACCACGTGGT GTGCCCCAGA TCGAGGTCAC CTTCGATATC 1551 GATGCCAACG GTATCCTCAA CGTGACTGCC CTGGAGCGTT CGACCAACAA GGAGAACAAG ATCACCATTA CCAACGACAA GGGTCGTCTC TCCAAGGAGG ACATCGAGCG CATGGTCAAC 1671 GAGGCCGAGA AGTACCGCAA CGAGGATGAG AAGCAGAAGG AGACCATTGC CGCCAAGAAC GGCCTCGAGT CGTACTGCTT CAACATGAAG GCCACCCTCG ACGAGGATAA CCTGAAGACC 1791 AAGATCTCGG ACTCTGACCG CACCACAATC CTGGACAAGT GCAACGAGAC CATCAAGTGG CTGGATGCCA ACCAGCTGGC TGACAAGGAG GAGTACGAGC ACCGCCAGAA GGAACTGGAG 1911 GGTGTGTGCA ACCCGATCAT TACCAAGCTA TACCAGGGCG CCGGTTTCCC ACCCGGTGGC ATGCCCGGCG GTGGTGGAGG TATGCCCGGA GCGGCTGGTG CCGCTGGCGC TGCCGGAGCC 2031 GGCGGTGCTG GCCCCACCAT CGAGGAGGTC GACTAAACCA TTCACCCCCCA CACCTCAATG CAACCATACA GTAACAGTTC TCCAAACAAT TTACCAACCA AACACAGTAG AAGAGTTGCT 2151 2168
- TAAACAAACT TGGATTC

FIG. 2. Complete nucleotide sequence of the *Drosophila hsc4* gene contained on plasmid pMG34. The entire protein-coding region, delimited by start (ATG) and stop (TAA) codons, is 1,953 bp and contains no introns. The predicted 5' and 3' splice sites of the intron separating exon 1 and exon 2 and the predicted start of transcription (5' END) are indicated by arrows. Nucleotides are numbered with reference to the predicted start of transcription (+1). The 5' untranslated region of the *hsc4* gene is approximately 120 bp (6); intronic sequences are not included in this numbering scheme. A consensus TATA box at -23 to -31, and two regions (-91 to -104 and -144 to -157) with sequence similarity to the consensus heat shock element (20) are observed upstream of the start of transcription. Stars indicate identical matches to this heat shock element, T—GAA—TAA—G. We have marked the approximate 5' end of the *hsc4* transcript as +1.

MPAIGIDLGTTYSCVGVYQHGKVEINAYDQGNRTTPSYVAFTDSERLNGEPAKNQVAMNPRNTVFDAKRLIGRKYDDPKIAEDMKHWPFKVVSD30	•	Dros hsp70
MSKAPAVGIDLGTTYSCVGVFQHGKVEIIANDQGNRTTPSYVAFTDTERLIGDAAKNQVAMNPTQTIFDAKRLIGRKFDDAAVQSDMKHWPFEVVSAD	K 100	Dros hsc70
MSKGPAVGIDLGTTYSCVGVFQHGKVEIIANDQGNRTTPSYVAFTDTERLIGDAAKNQVAMNPTNTVFDAKRLIGRRFDDAVVQSDMKHWPFMVVNDA	Ř 100	Hum hsc70
PKIGVEYKGESKRFAPEEISSMVLTKMKETAEAYLGESITDAVITVPAYFNDSQRQATKDAGHIAGLNVLRIINEPTAAALAYGLDKNLKGERNVLIFI	L 197	Dros hsp70
PKIEVTYKDEKKTFFPEEISSMVLTKMKETAEAYLGKTVTNAVITVPAYFNDSQRQATKDAGTIAGPNVPRIINEPTAAAIAYGLDKKAVGERNVLIFI	L 200	Dros hsc70
OCO PKVQVEYKGETKSFYPEEVSSMVLTKMKEIAEAYLGKTVTNAVVTVPAYFNDSQRQATKDAGTIAGLNVLRIINEPTAAAIAYGLDKKVGAERNVLIFI	L 200	Hum hsc70
${\tt GGGTFDVSILTIDEGSLFEVRSTAGDTHLGGEDFDNRLVTHLAEEFKRKYKKDLRSNPRALRRLRTAAERAKRTLSSSTEATIEIDALFEGQDFYTKVSON CONTRACT CONTRACTACT CONTRACTACT CONTRACT CONTRACT CONTRACTACT CONTRACTACT CONTRACTAC$	R 297	Dros hsp70
$\tt GGGTFDVSILSIDDG. IF \tt EVKSTAGDTH \tt LGGEDFDNRLVTH \tt FVQEFKRKHKKD \tt LTNKRALRRLRTACERAKRTLSSSTQASIEIDSLFEGTDFYTSITE STATE STA$	R 299	Dros hsc70
GGGTFDVSILTIEDG.IFEVKSTAGDTHLGGEDFDNRMVNHFIAEFKRKHKKDISENKRAVRRLRTACERAKRTLSSSTQASIEIDSLYEGIDFYTSI	R 299	Hum hsc70
ARFEELCANLFRNTLOPVEKALNDAKMDKGQIHDIVLVGGSTRIPKVQSLLOEFFHGKNLNLSINPDEAVAYGAAVQAAILSGDQSGKIQDVLLVDVA	L 397	Dros hsp70
ARFEELNADLFRSTMDPVEKALRDAKLDKŠVIHDIVLVGGSTRIPKVQRLLQDLFNGKELNKSINPDEAVAYGAAVQAAILHGDKSQEVQDLLLDVT	L 399	Dros hsc70
ARFEELNADLFRGTLDPVEKALRDAKLDKSQIHDIVLVGGSTRIPKIQKLLQDFFNGKELNKSINPDEAVAYGAAVQAAILSGDKSENVQDLLLLDVT	L 399	Hum hsc70
${\tt Slgietaggvmtklierncripckotktfstysdnopgvsiovyegeramtkdnnalgtfdlsgippaprgvpoievtfdldangilnvsakemstgki$	к 497	Dros hsp70
SLGIETAGGVMSVLIKRNTTIPTKQTQTFTTYSDNQPGVLIQVYEGERAMTKDNNLLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNNLGKFELSGIPPAPRGVPQIEVTFDIDANGILNVTALERSTNKINGSULVYEGERAMTKDNTALERSTNKGNTANGILNVTALERSTNKGTNANGILNVTALERSTNKGTNANGANTANGANGANTANGANGANTANGANGANTANGANTANGANTANGANTA	N 499	Dros hsc70
SLGIETAGGVMTVLIKRNTTIPTKQTQTFTTYSDNQPGVLIQVYEGERAMTKDNNLLGKFELTGIPPAPRGVPQIEVTFDIDANGILNVSAVDKSTGK	N 499	Hum hsc70
NITIKNDKGRLSQAEIDRMVNEAEKYADEDEKHRQRITSRNALESYVFNVKQSV.EQAPAGKLDEADKNSVLDKCNETIRWLDSNTTAEKEEFDHKME	L 596	Dros hsp70
KITITNDKGRLSKEDIERMVNEAEKYRNEDEKOKETIAAKNGLESYCFNMKATLDEDNLKTKISDSDRTTILDKCNETIKWLDANQLADKEEYEHROKI	Ľ 599	Dros hsc70
KITITNDKGRLSKEDIERMVQEAEKYKAEDEKQRDKVSSKNSLESYAFNMKATVEDEKLQGKINDEDKQKILDKCNEIINWLDKNQTAEKEEFEHQQKH	L 599	Hum hsc70
TRHCSPIMTKMHQQGAGAAGGPGANCGQQAGGFGGYSGPTVEEVD 641 Dros hsp70		
EGVCNPIITKLYQGAGFPPGGMPGGGGGGMPGAAGAAGAAGAAGAGGAGPTIEEVD 651 Dros hsc70		
exvcnpiitklygsaggmpggmpggfpgggappsggassgptieevd 646 Hum hsc70		

FIG. 3. Comparison of *Drosophila* hsc70, human hsc70, and *Drosophila* hsp70 amino acid sequences. The predicted amino acid sequence encoded by the *Drosophila* hsc4 gene (*Dros* hsc70) is compared with the predicted amino acid sequences of human hsc70 (Hum hsc70 [8]) and *Drosophila* hsp70 (*Dros* hsp70 [12]). The sequences were aligned by using the GAP program of the University of Wisconsin Genetics Computer Group with a gap weight of 5.00 and a length weight of 0.30. Identical amino acids ( $\bullet$ ) and conservative differences ( $\bigcirc$ ) are indicated.

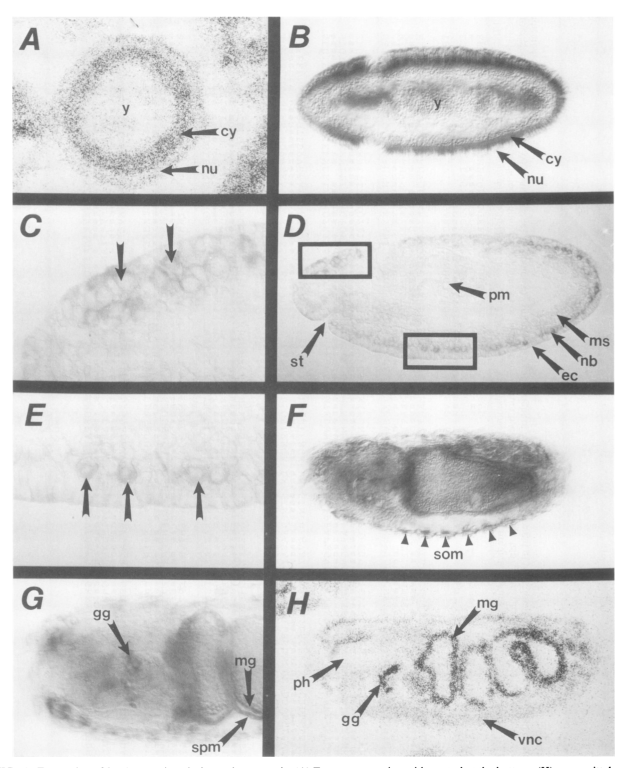


FIG. 4. Expression of hsc4 transcripts during embryogenesis. (A) Transverse section with ventral at the bottom; (H) parasagittal section with anterior to the left and ventral at the bottom; (B to G) whole-mount embryos labeled with nonradioactive probes (30). From fertilization through the early stages of cleavage, expression of hsc4 transcripts is uniformly distributed in the embryo (not shown). During the late syncytial and cellular blastoderm stages through early gastrulation, most of the hsc4 transcript is observed between the peripherally positioned nuclei (nu) and the central yolk (y), i.e., in the cytoplasmic compartment (cy) (A and B). hsc4 transcripts remain essentially uniform in distribution at the basal level in all embryonic tissues until the germ band is almost fully extended. At this time a punctate band of more intense hybridization internal to the region of the ectoderm (ec) and exterior to the mesoderm (ms), where neuroblasts (nb) have segregated (D, enlarged in E), is detected. With development the intensity of the band increases, presumably due to either increased numbers of cells becoming enriched for the hsc4 transcript or increased expression of the transcript in the enriched cells. Enrichment is also observed in the procephalic neurogenic regions (D, enlarged in C). Throughout the remaining stages of embryogenesis, the lining of the developing gut is

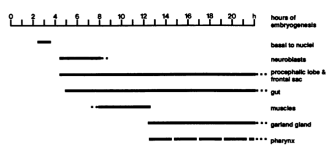


FIG. 5. Schematic summary of tissue-specific enrichment for hsc4 transcripts during embryonic development. The top scale represents hours of embryogenésis with hatching (h) occurring at 22 h of embryonic development. The solid lines indicate the times at which enrichment was observed in the tissues indicated at the right. The dashed line for the pharynx indicates that hsc4 transcription was below the basal level of transcription observed in other nonenriched tissues.

embryonic and transformed cells and in some secretory cells (1, 10, 16, 22), suggesting that hsc4 is a homolog of the mammalian hsc70 gene. Consistent with this interpretation is the fact that hsc4 is more closely related to the mammalian hsc70 than to the heat-inducible *Drosophila* hsp70 protein. In addition, like the mammalian hsc70 protein, the *Drosophila* hsc4 protein product translocates to the nucleus after thermal stress (19, 33).

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highly enriched. By germ band shortening and extending through dorsal closure, the developing somatic (som) and splanchnic (spm) musculature become enriched (F and G). Finally, late in embryogenesis hsc4 tissue enrichment is observed in the proventriculus, gut (mg), the garland gland (gg), and that region of the frontal sac dorsal to the pharynx which is derived from the procephalic lobe (H). Note that at this stage the lining of the pharynx (ph) shows hybridization below background. This is the only tissue observed during embryogenesis to have less than basal-level hybridization. Other abbreviation: vnc, ventral nerve cord. That the hybridization observed is specific for hsc4 and not other hsp70-related genes was supported by the fact that a 0.8-kb PstI-SaII fragment taken from the highly divergent 3' end of hsc4 (K. Palter, unpublished observations) (panels B to G) showed patterns of hybridization identical to that with cDNAs c321 and cHsc4, which extend into the less divergent 5' end (panels A and H).

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