The 27 kDa human heat shock protein (hsp27) is encoded by a gene family of 4 members. Two genomic fragments hybridizing to cDNA encoding hsp27 have been isolated, characterized, and sequenced. One clone is a member of a cluster of three genes linked within a 14-18 kb region of the genome and encodes a transcript interrupted by two intervening sequences. A single open reading frame encodes a polypeptide of 22,300 deduced molecular weight. The 5' flanking region contains two transcription start sites and sequences homologous to the Drosophila consensus heat inducible control element. Induction of both potential transcripts follows heat shock in vivo. Accurate heat inducible transcription occurs at both start sites after injection into Xenopus oocytes. The second genomic clone is a processed pseudogene lacking promoter elements and is unlinked with the other members of the hsp27 gene family. The amino acid sequence of human hsp27 shows striking homology with mammalian α crystallin, and contains a region towards the carboxy terminus which shares homology with the small hsp of Drosophila and other organisms.

INTRODUCTION

The induction of transcription of the heat shock genes in response to elevated temperatures, first described in Drosophila (reviewed in 1), has now been demonstrated to be universal among prokaryotes and eukaryotes (2). The polypeptides (hsp) encoded by the heat shock genes are conserved to varying degrees. The nucleotide sequences encoding the 70 kDa hsp, the most conserved class, cross hybridize with the corresponding sequences of organisms as phylogenetically distant as insects and bacteria (2). The members of the 80-90 kDa class of hsp of diverse eukaryotes still share antigenic determinants (3). Similarities among the small, 20-30 kDa hsp of different organisms are only evident upon examination of the amino acid sequences (4,5). Key regulatory components of the heat shock response are also conserved. Heat shock genes from various organisms have been shown to be heat inducible following introduction into heterologous cells (6-8), and are activated at the temperature characteristic of the heat shock response of the host cell. Deletion studies have identified a consensus sequence (Pelham sequence) in the promoter region required for heat inducibility (6), suggesting conservation of control signals common to all heat shock genes. The heat shock proteins belong to a larger
group of polypeptides, the stress proteins, that are induced in various combinations in response to environmental challenges and developmental transitions. The heat shock genes therefore provide an excellent system in which to study the sequence requirements for coordinate regulation of overlapping subsets of a dispersed group of genes by different inducers of transcription.

The specific functions of the individual hsp have not yet been identified, but synthesis of the small hsp has been shown to be correlated with the acquisition of thermotolerance (2,9). In cultured Drosophila cells, ecdysone induces synthesis of only the low molecular weight class of hsp. Such induced cells are found to be thermotolerant without prior heat treatment (10). Mutants of Dictyostelium that are unable to acquire thermotolerance have been shown to lack the ability to synthesize the small hsp (11).

The mammalian small (27-28 kDa) heat shock protein (hsp27) labels very weakly with $^{35}$S methionine; it was therefore overlooked in early studies and has only recently been subjected to detailed analysis (12-15). Several isoelectric variants of mammalian hsp27 are found, which also differ slightly in size. The variants derive in part from differences in phosphorylation state (13,14), but the possibility that they also represent several unrelated polypeptides cannot yet be excluded. Both the induction of synthesis of hsp27, and the extent of phosphorylation, are responsive to different environmental stimuli. Synthesis of hsp27 is induced in mammalian cells in the presence of arsenite and amino acid analogs as well as during hyperthermia (13). Increased phosphorylation of mammalian hsp27, without increased synthesis of the proteins, follows serum stimulation of growth, or exposure to tumor promoters or calcium ionophores (13). This suggests that covalent modification may regulate the activity of these polypeptides.

We report here the characterization and sequence of two of the four human genomic sequences encoding hsp27, one of which is closely linked to two others. The linked gene encodes 2 overlapping transcripts that have different 5' transcription start sites and contain 2 intervening sequences. The promoter region of this gene contains sequences homologous to the Drosophila consensus heat inducible control element. The gene is accurately transcribed in vitro, and is heat inducible when introduced into Xenopus oocytes. The second of the genes characterized in this study is unlinked to the others, and is shown to be a processed pseudogene that contains no intervening sequences or transcription control signals. The deduced amino acid sequence of human hsp27 is homologous to the reported sequences of small hsp from a variety of organisms (4,7,16,17).
The amino acid sequence also shares with other small hsp a striking homology to the sequence of mammalian α crystallin (4,16).

MATERIALS AND METHODS

Materials: Restriction enzymes, DNA polymerase, and T4 polynucleotide kinase were from Bethesda Research Laboratories, reverse transcriptase from BioRad, calf intestine alkaline phosphatase from Boehringer, SI nuclease from P.L, RNasin from Promega Biotec, and T4 ribonuclease, protease K, and DNAase from Sigma. All sequencing reagents and ³²P dATP were obtained from Amersham, while ³²P-labelled ATP, UTP, and dCTP were from ICN. Electrophoresis supplies were DNA grade from BioRad and nitrocellulose filters were from Schleicher and Schuell.

Screening a human genomic library: An Alu I-Hae III human genomic library prepared by Maniatis (18) in phage Charon 4A was screened with the cDNA clone pHS208 (19) that encodes the human hsp27. From one million plaques, four hybridizing phage were isolated, plaque purified, and mapped by restriction digestion and Southern blotting using standard techniques (20). The gene containing fragments were identified by hybridization to the cloned cDNA probe pHS208, and by preferential hybridization to cDNA that was made to RNA from heat shocked cells using short oligonucleotide primers as described (20). Restriction fragments containing the mRNA coding region were subcloned into pUC12.

Southern analysis of genomic sequences: High molecular weight human placenta or sperm DNA was digested with restriction enzymes, and 10μg aliquots electrophoresed on agarose, and blotted onto nitrocellulose (21). The blots were hybridized to the nick-translated cloned pHS208 probe at 68° according to standard procedures, with a final stringent wash for 15 min at 68° in 0.1 x SSC, 0.1 % SDS.

SI nuclease analysis and primer extension: SI nuclease protection analysis was done with uniformly labelled (22) or end-labelled probes (23), according to the protocol described by Favaloro (24). Uniformly labelled probes were obtained by nick-translation, ligation and restriction of the subcloned plasmid DNA, followed by isolation of the insert fragments on gels (20). The 3' end-labelled probes were prepared with Klenow, and the 5' end-labelled probes were first treated with phosphatase and then labelled with T4 polynucleotide kinase. Probes were denatured at 90° for 10 min, and hybridization continued for 3 hours at 58°. The SI nuclease digestions were carried out at 22° for 30 min at 200 units/ml. Primer extension analysis was performed as described by Lee and Luse (25). The precise transcription start site was determined by comparison of
Figure 1 Characterization of human genomic clones encoding hsp27. The maps of the Charon 4A phage λhsB and λhsII are shown. The filled region indicates the restriction fragment hybridizing to the cloned cDNA and to 32P cDNA. The portion subcloned is shown expanded in the second line. The regions sequenced are marked with an open bar on the map of the subclone. Beneath the maps, the open arrows indicate the S1 nuclease protected regions and the direction of transcription. Restriction enzymes: B-Bam HI, Bg-Bgl II, E-Eco RI, H-Hind III, K-Kpn I, P-Pst I, Pv-Pvu II, S-Sma I, X-Xba I.

Expression in vitro and in vivo: In vitro transcription was assayed in extracts of HeLa cells prepared according to Manley (26). Linearized plasmid DNA (500 ng) was added to each 10 μl assay, and transcription was carried out in the presence of 32P labelled UTP. The labelled transcription products from assays incubated with or without α-amanitin were analyzed by electrophoresis and autoradiography.

In vivo transcription was assayed following injection of 5 ng of plasmid DNA into the germinal vesicle of Xenopus oocytes as described by Voellmy (8). Injected oocytes were held at 22°C 14–20 hours. For heat induction of transcription, oocytes were incubated at 36°C for 90 min. RNA was prepared from control oocytes kept at 22°C, or from induced oocytes immediately following heat shock (8). The presence of transcripts from the introduced plasmid was assayed by S1 nuclease protection analysis using 5' end-labelled probes, or by primer extension as described above.
**Sequencing:** Sequencing of restriction fragments subcloned into M13 was performed by the dideoxy termination method (27). For collation of the complete sequence, the programs of Schwindiger were used (28). Homologies among DNA and protein sequences were determined with the PCS program (29).

**RESULTS**

**Isolation and characterization of genomic clones:** Four phage were isolated from the Charon 4A library that hybridized to the cDNA clone pHS208. We have shown by hybridization selection and translation that this cDNA clone is complementary to a heat inducible RNA that specifically encodes human hsp27 (19). Restriction mapping revealed that three phage, represented by λhs11, contained DNA overlapping the same genomic region, while the fourth, λhs8, was quite different (Figure 1). Each phage contained only one region homologous to hsp27 mRNA (Fig 1, 2711 and 2708). A 2500 bp Eco RI fragment from λhs8 and a 3500 bp fragment from λhs11 were cloned into plasmid vectors to give subclones 2708 and 2711, respectively (Fig. 1). The subcloned segments were analyzed by SI protection experiments. The uniformly labelled Eco RI insert from subclone 2708 gave a single 750 bp protected fragment after hybridization with RNA from heat shocked cells. The Eco RI-Hind III segment from 2711, however, gave protected fragments of 450, 350, and 65 bp, indicating the presence of two intervening sequences. The boundaries of the transcription units and intervening sequences as depicted in Fig. 1 were localized using 5' and 3' end-labelled restriction fragments as hybridization probes. The precise boundaries of the intervening sequences and the 3' termini were verified by comparison with the sequence of the cloned cDNA (see below). Experiments using 5' end-labelled probe fragments indicated that the gene cloned in 2711 has two transcription start sites located approximately 50 bp apart (see Fig. 3) Both transcripts can be found at low levels in control cells, with the shorter transcript more abundant than the longer one. Levels of the two mRNAs are induced more than 20-fold during heat shock, without a significant change in proportions (data not shown).

The sequence of the human hsp27 genes: In Figure 2A the sequence of the entire transcribed region of 2711 plus 160 5' flanking nucleotides and 1000 3' flanking nucleotides, are shown aligned with the homologous sequences of 2708. The entire sequence of the cDNA clone pHS208 has been determined and it corresponds precisely to the sequence of 2711 presented in uppercase letters between the black triangles in Figure 2A. The sites at which transcription initiates in 2711 were determined to within one or two bases by the SI and primer extension analysis shown in Figure 3. Because the SI nuclease gave multiple bands at the
Figure 2A. The aligned DNA sequences of 2711 and the homologous regions of 2708. The numbered DNA sequence of 2711 is shown, with +1 being the start of the initiator ATG. Upper case letters indicate sequences present in the processed mRNA; the amino acid sequence is aligned below the DNA. The sequence of 2708 is aligned above that of 2711; dashes indicate identity of nucleotides; spaces have been introduced to maximize matching. Only the amino acids in 2708 which differ from 2711 are shown, above the sequences. The common control sequences are shaded. The Pelham sequence is underlined. Large arrows indicate transcription start sites. The black triangles indicate the boundaries of the cDNA sequence cloned in pHS208. The dotted underlines and overlines, and the horizontal arrow indicate recognized sequences described in the text. The small double brackets over the sequence in IVS I and in the 3' flanking region delineate regions of homology to the human Alu repeat.

B. The 5' flanking sequences of the pseudogene 2708: Sequences 5' to the 2708 regions in Fig 2A are shown. Upper case letters indicate homology with 2711, and nucleotide +1 is at the ATG. The double small brackets enclose the Alu repeat. Dotted underlines indicate the polymerase III promoter. The flanking direct repeat is marked with dotted overlines.

A+T rich first cap site at all digestion temperatures and nuclease concentrations assayed, the termini indicated by primer extension were used in assigning the cap sites (Fig. 2A, large arrows). The resulting mRNAs have 5' untranslated leaders of 40 (major transcript) and 90 (minor transcript) nucleotides. The transcribed region contains two intervening sequences of 723 and 120 nucleotides, bounded by appropriate splice signals. The precise boundaries of the intervening sequences were determined by comparison with the sequence of the cDNA clone pHS208. The first ATG in both transcripts is designated base +1 in the numbering of 2711 shown in Figure 2A. This ATG opens a reading frame that extends to a stop codon at nucleotide 1437, and predicts a polypeptide of 199 amino acids with one internal methionine, and a deduced molecular weight of 22,300. All other potential reading frames would encode polypeptides no longer than 47 amino acids. The 3' terminal nucleotide, corresponding to the polyadenylation site of the hsp27 cDNA, occurs 20 bases downstream from a consensus AATAAA polyadenylation signal. The two complete spliced transcripts are 825 and 775 nucleotides long and have 139 3' noncoding nucleotides. Particular G + T rich sequences twenty to thirty-eight bases 3' to the AATAAA have been reported.
Figure 3. Determination of the transcription start site in 2711: S1 nuclease protection and primer extension were used to identify the precise transcription start site as described. A: T lane of the sequence ladder; B: S1 nuclease protected fragments; C: primer extension products. The sequence of the region shown is aligned above the figure; arrows indicate the start sites. The source of the probes is shown below the figure. The map shows the portion of 2711 containing the complete first exon, and 250 flanking bases. E-Eco RI, P-Pst I, Pv-Pvu II, T-Taq I. Labelling was at the Taq I site as shown.

To be important for efficient processing of mRNAs (30). A similar sequence is found thirty-five nucleotides downstream from the AATAAA in 2711 (Fig. 2A, between the stars).

The 5′ flanking sequences of 2711 contain TATA boxes (CATAA) 22 nucleotides upstream from each transcription start site (Figure 2A, shaded). One hundred twenty-one nucleotides 5′ to the distal cap site is a sequence homologous to the Drosophila consensus heat control element (CTGGAAT[N]TTCTAGA) (6), with 8 nucleotides separating the initial CT from the remaining sequence (underlined). Seven bases downstream there is an abbreviated repeat of this element (also underlined). A dotted overline in the same region marks an inverted complement of the first consensus sequence. From nucleotides -190 to -180, underlined with an arrow, is an inverted copy of the consensus Spl binding site (31).
Both the first intervening sequence and the 3' flanking sequences contain regions of homology with the human Alu repeat family (32). The double brackets over the sequences in Figure 2A mark the boundaries of a sequence in IVS I which corresponds to one-half of the Alu repeat, and of a 3' flanking sequence which contains an entire copy of Alu but is not flanked by the usual short direct repeats, and contains no apparent polymerase III promoter sequences.

The hsp27 gene sequences cloned in 2708 clearly indicate a processed pseudogene. The intervening sequences present in 2711 are absent from 2708, and the closed splice sites correspond precisely to the corresponding sequences of the cDNA clone. Also, 2708 is flanked at the 5' and 3' boundaries with a poly A sequence and lacks the 5' leader sequences corresponding to the distal transcription start site in 2711. The homologous region of 2708 differs from the 2711 transcript by 14.3%, of which 10% consist of single base differences and small deletions and additions. The remainder is the result of one 20 base and one 12 base segment present in 2711 and absent in 2708. Of the single base changes, 24 are transversions, and 30 are transitions. A total of 6 base differences in 2708 are insertions, and in addition to the stretches mentioned above, 9 bases are deleted. The 3' terminus is also shortened by about 7 nucleotides with respect to 2711. The 2708 sequence encodes an open reading frame of 193 amino acids with 23% differing from 2711, 7% due to nucleotide changes in codons, and the rest due to frame shifts.

Figure 2B shows 820 5' flanking nucleotides of 2708, including the poly A and the 2711-homologous sequences up to the ATG (nucleotide +1). There are no 5' flanking sequences similar to polymerase II transcription control elements. However, between bases -497 and -190 there is a complete member of the Alu repeat family (double brackets), flanked by an 11 base direct repeat (dotted overline), and containing sequences corresponding to the split polymerase III promoter (dotted underline) (33).

Expression of the cloned genes in vitro and in vivo: Both forms of the hsp27 gene contain sequences that act as promoters in the in vitro transcription system described by Manley (26). DNA from 2711 linearized with Hind III produces a 2.2kb transcript in vitro (Figure 4A). This agrees with the start site predicted by the sequence, and by the results of SI analysis. Transcription from 2711 is completely inhibited by α-amanitin, confirming that the gene contains an active polymerase II promoter. In contrast, there is no evidence for polymerase II transcription of 2708. Transcription from Eco RI digested 2708 DNA is only slightly sensitive to 5 μg/ml of α-amanitin and labels an RNA of about 1.2 kb, 450 nucleotides longer than predicted by SI mapping. RNA of
The image shows a gel electrophoresis result with indications of amanitin concentration in μg/ml. There are markers for kb (kilobase) and bands at 2.3 and 2.0. The gel contains DNA fragments labeled with H and X restriction enzymes, with bands at 179 and 128. The gel is labeled with M, 2711, and 2708.
this size, however, would result from initiation at the polymerase III promoter identified in the 5' flanking sequences of 2708. Northern blot analysis of RNA from HeLa cells incubated at 42°C for 3 hr, hybridized with pHS208, shows one band of 960 nucleotides, appropriate for a polyadenylated transcript from 2711 (19, 20). No evidence of an 1.2 kb mRNA is found, indicating that transcription initiating at the polymerase III promoter in vivo does not continue through the regions of 2708 that are homologous to hsp27 mRNA.

The hsp 27 gene in 2711 also functions in vivo when injected into the Xenopus oocyte (Figure 3B). S1 nuclease protection studies of the RNA isolated from heat stressed Xenopus oocytes shows that transcription is initiated accurately at both start sites on the introduced gene, in approximately the same proportions as is seen in human cells. No protected fragments corresponding to the 2711 transcripts are found in oocytes held at the control temperature. Northern transfer analysis of RNA from injected oocytes shows that approximately 50% of the transcripts from 2711 are processed to a size which comigrates on agarose gels with authentic human hsp27 mRNA, while the remainder appears in a diffuse region of higher molecular weight (not shown).

Homologies between hsp27 and other low molecular weight heat shock proteins and mammalian α crystallins: The amino acid sequence of human hsp27 shows two large regions with more than 50% homology to mammalian α crystallins, separated by a non-homologous region 14 amino acids longer in hsp27 than in the crystallins (Fig. 5A). Fifty percent of the residues between amino acids 10 and 59 of hsp27 are found in the same positions in either mouse or bovine α crystallin. In a region between human hsp27 amino acid 84 and 178, there is 64% agreement with the sequence of the α crystallins. Included in this region is a stretch of 11 contiguous amino acids identical to the corresponding sequence of bovine α crystallin B.

Comparison of the sequences of human hsp27 and small hsp from Drosophila (16,17), Xenopus (5), Caenorhabditis (4), and soybean (34) indicates that the carboxy terminal half of the polypeptide is the most highly conserved. Within

Figure 4 A. In vitro expression of genomic sequences encoding hsp27: Autoradiograph of transcription products from Hind III cut 2711 and Pst I cut 2708 incubated in the Manley system in the presence of the indicated amounts of α amanitin. Arrows indicate the polymerase II transcript from 2711, and the polymerase III transcript from 2708. M- Hind III lambda marker, P- control transcription assay containing pBR322. B. Heat inducible transcription of 2711 in Xenopus oocytes: Transcription products from plasmid DNA introduced into oocytes as described in Methods were analysed by S1 nuclease protection. Fragments protected by RNA isolated from oocytes were compared with those protected by HeLa RNA. H - RNA from HeLa cells; X - oocyte RNA; c - incubation at control temperature; h - heat shocked cells or oocytes. Numbers indicate length in bases of protected fragments. Star indicates position of probe fragment.
Figure 5 A. Comparison of the amino acid sequences of human hsp27 and mammalian α crystallins: The amino acid sequence of human hsp27 (HS) has been aligned with that of murine α crystallin A (A), and bovine α crystallin B (B). Hyphens were introduced to maximize homologies between the crystallins; spaces were introduced to gain the best alignment with the heat shock protein sequence. Amino acid homologies between hsp27 and either crystallin are capitalized and shaded.

B. Comparison of the regions of homology of small hsp from several organisms: The region of 78 amino acids of human hsp27 showing greatest homology to the amino acid sequences of other heat shock proteins is shown aligned with the corresponding sequences from Drosophila (Dros), Xenopus, Caenorhabditis (Caeno), and α crystallin (Crys). Alignment was maximized by introduction of spaces (hyphens), and deletions (letters below the lines). Homologies are indicated by shaded upper case letters. Arrows indicate the boundaries of the second exon in human hsp27.
Figure 6 A. Southern analysis of genomic sequences encoding hsp27: Human sperm DNA (lanes 4, 7, 10) or placental DNA (lanes 5, 8, 11) was digested with the indicated restriction enzymes, and Southern blots were hybridized to nick-translated pH208. Appropriate digests of the genomic subclones were run in parallel: lane 3, Eco RI cut 2708, lane 6, Eco RI-Xba I cut 2711, lane 9, Pst I cut 2708. Lane 1 - lambda markers, lane 2 - linearized pH208. The stars indicate incomplete digestion products.

B. Map of the organization of human hsp27 genes: The heavy bars indicate the limits of the DNA pieces cloned in phage λhsB and λhs11, as indicated. Short arrows show the location and direction of transcription of the cloned regions hybridizing to pH208. Only restriction enzymes used in Fig. 6A are indicated; the numbers between the vertical lines give the fragment sizes in kb. The location of the four members of the gene family are indicated on the map by the labels they are given in the Eco-Xba digest in Fig 4A, Lanes 7&8.
the region between amino acids 100 and 178 of human hsp27, the small hsp presented in Figure 5B show from a minimum of 42% to a maximum of 50% homology. The soybean 17.5 kDa hsp shows only a short region of homology, with 8 of 14 amino acids identical between amino acid 158 and 171 of hsp27 (not shown). The 5 amino acid sequence, Asp-Gly-Val-Leu-Thr, conserved in all Drosophila small hsp, lies within this region of the soy protein, with Asn substituted for Asp (34). In human hsp27 the changes in this region have been conservative; the equivalent sequence reads Glu-Gly-Thr-Leu-Thr.

Organization of hsp27 genes within the genome: We have isolated a heat inducible gene and a pseudogene encoding human hsp27. To determine whether there are additional human genomic sequences encoding hsp27, human DNA from two sources was analyzed by Southern blotting. Human placental and sperm DNA were digested with specific restriction enzymes which did not cut the pHS208 cDNA clone, and which gave unique fragments of predicted sizes from the genomic subclones that hybridize with the cloned cDNA. Parallel digests of the subclones were electrophoresed for comparison beside the genomic DNA, and the blot was probed with nick-translated pHS208 (Figure 6).

An Eco RI digest of λhs8 yields a single internal 2.8 kb fragment that hybridizes with pHS208. In λhs11, however, the hybridizing Eco RI fragment contains a linker-generated site adjacent to a phage arm, and the actual size of the equivalent genomic Eco RI fragment can not be predicted. However, digestion of λhs11 with both Eco RI and Xba I gives an internal hybridizing fragment of 2.0 kb. Pst I digestion of λhs8 and λhs11 gives hybridizing fragments of 1.6 and 2.3 kb respectively. Figure 6A shows the presence of both gene arrangements in each preparation of human DNA. Hybridization of the hsp27 cDNA probe to Eco RI digested DNA reveals 3 hybridizing fragments, each containing no more than 1 or 2 gene copies (quantification not shown). The fragments are labelled a, b, and c in order of decreasing size. Fragment c comigrates with the 2.8 kb Eco RI fragment from λhs8. In the Eco RI-Xba I digest, the fragments are given the label of the Eco RI fragment to which they correspond. In this digest, a 2.0 kb hybridizing fragment is seen (ax2), that is derived from Eco RI fragment a, and comigrates with the Eco RI-Xba I fragment of λhs11. The predicted 2.3 and 1.6 kb fragments are also present in the Pst I digest, labelled a (containing the 2711 gene) and c (containing the 2708 gene). Thus, there does not appear to have been any gene rearrangement during cloning and amplification.

The data in Figure 6A also reveal two more members of the gene family homologous to pHS208, that are linked with the gene cloned in λhs11 in a
contiguous region of the genome of between 14 and 18 kb. The third form of the
gene appears as fragment b in the Eco RI digest, and reveals a restriction site
polymorphism between the two sources of human DNA. Fragment b is unchanged in
the Eco RI-Xba I digest, but Eco RI fragment a is cleaved by Xba I into two
hybridizing fragments. Only one hybridizing fragment, ax2, can be derived from
the DNA region cloned in λhs11 (see Figure 1). Therefore, the new fragment,
axl, represents a fourth separate sequence homologous to pHS208, that is linked
to the λhs11 sequence within an 8.6 kb Eco RI fragment in the genome. The axl
sequence and the Eco RI b sequence appear to be included together in the 9.2 kb
Pst I b fragment. Thus, the Eco RI b sequence is also linked to the λhs11 se-
quence in the genome within a fragment of at least 14 kb, and no more than 18
kb. The map of the gene organization derived from Southern analysis is shown in
Figure 6B. The λhsB pseudogene does not appear to be linked with any of the
other hsp27 genes.

DISCUSSION

The general picture of the structure and organization of heat shock genes,
as derived from the Drosophila system, is one of clustered families of related
genes lacking intervening sequences. The Drosophila hsp70 genes are found at
two loci on the same chromosome, one containing a pair of genes, the other a
cluster of four (35). The small hsp genes are clustered within a region of 11
kb (36-38). The clustering of small hsp genes within the genome now appears to
be common to many organisms. Four small hsp genes of Caenorhabditis are linked
in a remarkable head to head inverted repeat (4). Two Xenopus small hsp genes
have also been isolated as a tandem array (7). The unlinked member of the human
hsp27 gene family proved to be a processed pseudogene. In Drosophila, the only
hsp gene found to contain an intervening sequence is the single copy hsp83
gene. This gene is expressed in unstressed cells as well as during heat shock
(2). Intervening sequences are also present in the "cognate" genes that encode
proteins homologous to hsp70, but are not heat inducible (39). It initially
appeared likely, therefore, that intervening sequences in heat shock genes
characterized constitutively expressed homologues of the hsp, or heat inducible
genes with high levels of expression at control temperatures. As the heat shock
genes of a wider variety of organisms have been investigated, a more complex
picture has emerged. The small hsp genes of nematodes, which are under strict
temperature regulation, have been shown to be interrupted (4), and we present
here the sequence of a heat inducible human hsp27 gene which gives a spliced
transcript. The rapidity with which the hsp mRNAs reach the cytoplasm has been
attributed to the abbreviated processing required for unspliced mRNAs (40). There is evidence that processing and/or transport of most non heat shock mRNAs is inhibited during heat shock (40, our unpublished results). This suggests that there may be a specific mechanism by which the hsp27 transcript circumvents this inhibition.

The 5' regulatory sequences of the gene cloned in 2711 show characteristics common to most published heat shock gene sequences (2,4,6,7,15,17,35,41-43). Multiple copies of the Pelham consensus heat inducible control element are present: one quite exact, one inverted, and one abbreviated. This agrees with the complexity of this region found by Voellmy for the human hsp70 gene (41). The presence of more than one Pelham sequence has been reported to be required for efficient heat inducible expression of hsp genes introduced into homologous cells (44,45). The human hsp27 gene also contains an inverted copy of the sequence TGGCGGCCC, corresponding to the Sp1 protein binding site necessary for transcription of SV40 genes, that has also been found upstream from many eukaryotic genes (31). In SV40 this sequence functions in either orientation, since it is required both for early and late transcription. The key element of this sequence consists of GG6CGG, which is also found repeated in the 5' sequences flanking the human hsp70 gene (31,41). The TATAA sequences of 2711 correspond to the type (CATAAA) found in the majority of the globin genes (46).

At the 3' terminus the sequence GCCCT is found 10 nucleotides downstream from the polyadenylation site. This sequence is common to the 3' termini of many genes, however, no function has been determined for it (47). On the other hand, a sequence similar to GGTGTGG found 24-38 bases downstream from the AATAAA has been shown in deletion studies with the HSV thymidine kinase gene to be important for efficient processing and polyadenylation (30). This sequence has been identified in a number of viral and cellular genes (30), and is present in the human hsp27 gene at the appropriate site.

Pseudogenes, which have been reported in most gene families, appear to arise either as tandem repeats of a functional gene that are subsequently inactivated (48,49), or by reverse transcription of mRNA and insertion of the cDNA into the genome (50). The Xenopus small hsp pseudogene described by Bienz is of the first type; it is linked with functional hsp genes, but has a divergent Pelham sequence, encodes a polypeptide quite different from the normal small hsp, and lacks a functional promoter (7). In contrast, the human hsp27 pseudogene described here is of the second type, a processed pseudogene, characterized by precise excision of intervening sequences from the coding region, termination of the sequence at the polyadenylation site, and presence of a direct repeat.
flanking the pseudogene sequence. The presence of the long open reading frame encoded in 2708, unexpected in a pseudogene, is particularly unusual given the divergence between the 2708 and 2711 sequences. It is possible that the presence of the potential coding region in 2708 may reflect a sequence present in one of the hsp27 genes that have not yet been studied.

Hsp27 is a major cellular phosphoprotein, and has been reported to be phosphorylated primarily on serine residues (51). Interestingly, the small heat shock protein encoded in 2711 contains 21 serine residues (10%), as well as 13 threonine residues and 5 of tyrosine, giving a protein with almost 20% of its amino acids susceptible to phosphorylation. The presence of the single methionine residue in the polypeptide encoded by 2711 was surprising, since in our hands labelling of hsp27 with $^{35}$S methionine in vivo was not detected (12). Similar lack of labelling of human low molecular weight hsp with methionine has also been reported elsewhere (13). A recent report, however, shows that a low level of incorporation of methionine into a form of hsp27 can be seen following prolonged labelling of human lymphocytes at elevated temperature (52). The polypeptides encoded in the remaining two hsp27 genes may entirely lack methionine, which could mask the incorporation of label into the polypeptide encoded by the 2711 gene.

It is striking that the homology between human hsp27 and mammalian α crystallins is much more extensive than that between hsp27 and small hsp of lower organisms. The derivation of crystallins from the duplication of part of an ancestral small hsp has been suggested (53). Regions of amino acid alignment between hsp27 and α crystallin are found along almost the entire length of the crystallin. This suggests that the α crystallin gene may have been derived from a complete hsp gene that had an amino terminal region differing in sequence from earlier small hsp genes. The human small hsp would thus be closer evolutionarily to α crystallins than to invertebrate small hsp. Since crystallins have been shown to have highly conserved exon-intron distribution (53), the location of an intervening sequence at the same position in the Caenorhabditis hsp and the α crystallin gene is interesting. This position is quite different from that of either IVS in the human gene. The crystallin-like arrangement of exons may yet be found in one of the human hsp27 genes which has not been characterized. It is also possible that the Caenorhabditis gene is closer in structure, if not in sequence, to the ancestral form from which the crystallins are derived. Where present, the intervening sequences delineate protein regions which are most highly conserved (4, Fig 5). In Caenorhabditis the splice junction precedes the segment of greatest homology to the crystallins, and to
small hsp of other organisms. In the human hsp gene, the second exon corresponds to the region most similar to other small hsp and to the α crystallins. The amino acid sequence encoded in exon 2 forms a highly hydrophilic domain, which might be expected to be found on the exterior of the protein. The conservation of this domain may reflect its importance for the interaction of small hsp with other cellular components during stress. Analysis of the sequence and structure of the remaining human hsp27 genes, and those of other vertebrates, will clarify evolutionary relationships and identify the conserved domains that are likely to be of functional significance.

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