Expression, modification, and localization of the fushi tarazu protein in Drosophila embryos

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The fushi tarazu (ftz) protein of Drosophila is required during embryogenesis for the process of body segmentation. To study the biochemical properties of the ftz protein, ftz cDNA was expressed in Escherichia coli and the protein was purified to homogeneity. Polyclonal antibodies raised against the purified protein were used to localize and quantitate the protein during embryogenesis. Three temporally and spatially distinct phases of expression were observed, which include a previously undetected period later in embryogenesis. During this last phase, the protein is localized predominantly in the developing hindgut. Analysis of embryonic ftz protein on Western blots permitted us to approximate the number of protein molecules per nucleus. During the blastoderm phase of development, when ftz protein is most abundant, we estimate that there are ~20,000 molecules of protein per ftz-expressing nucleus. The embryonic ftz protein migrates more slowly on SDS-polyacrylamide gels than protein made either in E. coli or in a reticulocyte lysate system in vitro, indicating that it is modified in the embryo. To facilitate characterization of ftz protein made in embryos, an ftz overexpression system functional in Drosophila was developed. When fused to an hsp70 heat shock promoter and introduced into the germ line by P-element-mediated transformation, ftz could be overexpressed at all stages of development by heat shock. This protein is localized in the nucleus and comigrates on SDS-polyacrylamide gels with endogenous ftz protein. Two-dimensional gel electrophoresis followed by Western blotting resolves the overexpressed protein into a series of isoforms that differ in charge and electrophoretic mobility. Post-translational modification may influence the biochemical properties and functions of the ftz protein during embryogenesis.

[Key Words: ftz protein, expression, localization, modification, quantitation]

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An intriguing aspect of Drosophila embryogenesis is the process of body segmentation. In this process, the embryo is divided into repeating units that eventually adopt unique identities and functions [for reviews, see Gehring and Hiromi 1986; Scott and O'Farrell 1986]. The fushi tarazu (ftz) gene was first identified through its requirement in this metamerization process [Wakimoto and Kaufman 1981; Wakimoto et al. 1984]. Embryos that contain a homozygous deletion of the ftz gene produce only half the normal number of body segments and die before hatching as larvae. Because only certain regions of every other segment are deleted, ftz was classified as one of the 'pair-rule' class of segmentation genes. The gene is located within a cluster of homeotic genes that comprise the Antennapedia Complex [Wakimoto and Kaufman 1981]. The gene has since been cloned [Kuroiwa et al. 1984; Weiner et al. 1984] and sequenced [Laughon and Scott 1984; A. Kuroiwa, S. Baumgartner, and W.J. Gehring, unpubl.].

Transcription of the ftz gene is detectable only during embryogenesis [Kuroiwa et al. 1984; Weiner et al. 1984]. The message can be detected on Northern blots between the second and tenth hours of embryogenesis. During the first of two peaks in expression, the transcripts have been shown by in situ hybridization to accumulate in seven transverse bands that encircle the embryo [Hafen et al. 1984]. These bands correspond in location to the regions that are deleted in ftz~ embryos. Immunolocalization of the protein product shows that it accumulates in the nucleus [Carroll and Scott 1985]. Immunostaining of older embryos also demonstrated the occurrence of a second phase of ftz expression localized to the developing nervous system [Carroll and Scott 1985]. The use of ftz promoter sequence fusions to the reporter gene β-galactosidase also shows that the gene is expressed in both locations and that the temporal and spatial regulation lies predominantly at the level of transcript initiation [Hiromi et al. 1985; Hiromi and Gehring 1987]. It has been demonstrated recently that ftz expression in the nervous system is required for the expression of...
other genes and for the specification of certain cell neuronal lineages (Doe et al. 1988).

The first insight into the molecular mechanisms by which homeotic and segmentation genes control aspects of pattern formation came when the sequences of these genes were compared. It was found that the ftz gene has a 180-bp region of homology to homeotic genes found in the Antennapedia and Bithorax complexes (McGinnis et al. 1984a,b; Scott and Weiner 1984; Shepherd et al. 1984), which determines the identities of the anterior and posterior segments of the fly, respectively. This sequence, now known as the homeo box, encodes a 60-amino-acid peptide called the homeo domain. A portion of the homeo domain bears homology to the yeast developmental control genes Mata1 and Mata2 and the helix-turn-helix motif of a number of prokaryotic regulatory proteins, which enables them to bind DNA in a sequence-specific manner (Laughon and Scott 1984; Shepherd et al. 1984). A few homeo-domain-containing proteins have been shown to bind DNA with some sequence specificity (Desplan et al. 1985; Fainsod et al. 1986). Recently, Hall and Johnson (1987) have also demonstrated that the yeast protein Mata2 binds to specific DNA sequences in a homeo-domain-dependent manner. These results and the nuclear localization of these proteins suggest that they control aspects of pattern formation in Drosophila through an ability to bind to specific DNA sequences.

Although much is known about the temporal and spatial expression of the ftz gene, little is known about the actual properties and roles of the protein. Some of those properties are described here. Full-length protein expressed in Escherichia coli was purified and used to generate antibodies. Immunostaining of embryos using these antibodies confirms the data of Carroll and Scott (1985) and, in addition, shows that ftz protein is expressed a third time during a later stage of embryogenesis. It is also shown that ftz protein produced in Drosophila has a number of properties that are different from protein synthesized in two heterologous systems. Overexpression of ftz protein in Drosophila allowed us to characterize the protein and show that it is heavily modified.

**Results**

**Expression of ftz in heterologous systems**

To facilitate our biochemical characterization of the ftz protein, we first attempted to express the protein in bacteria as a means of producing large quantities of protein rapidly and economically. A cloning scheme was chosen that would produce the same protein as expected in Drosophila embryos, except for two additional amino acids encoded at the amino-terminal end. Two well-characterized expression systems were used initially in our attempt to express the gene in bacteria, and both failed to produce plasmid-specific expression of protein. In a third expression system, which has been described recently by Studier and Moffatt (1986), expression of a protein with an apparent molecular weight of 63,000 was induced (Fig. 1B, cf. lanes 1 and 2). The expression plasmid that we constructed, pGEMF1, was made by subcloning a ftz cDNA restriction fragment from one of the nonexpressing vectors into the plasmid pGEM-1 (Promega Biotec) such that the ftz gene is under the control of pGEM-1 T7 promoter (see Fig. 1A). Induction is achieved either by infecting cells that contain the plasmid with λ phages, which carry T7 RNA polymerase gene, or, alternatively, by using a lysogenic cell line that harbors a T7 RNA polymerase gene under the control of an inducible lac promoter.

The 63-kD protein induced by isopropanol-β-D-thiogalactopyranoside (IPTG) in pGEMF 1-transformed cells comigrated with protein translated in vitro using RNA transcribed from any of the ftz expression vectors constructed previously (Fig. 2, lanes 1 and 2). The reason that all expression systems were functional in vitro and that only the T7 system produced detectable levels of protein in bacterial cells (by either Coomassie blue staining or [35S]methionine labeling was not pursued; however, a number of potential advantages have been described by the developers of the system [see Studier and Moffatt 1986]. Even with these advantages, however, ftz protein only accumulated to a maximum abundance of 3–5% of total cellular protein, a value significantly lower than values of 30–50% reported for other proteins in this system [Studier and Moffatt 1986]. Inspection of Western blots shows that induced cells contain much degraded ftz protein, suggesting that protein synthesis may play a major role in limiting the abundance of the protein in the cell (data not shown).

Because the molecular weight of the ftz protein predicted by the sequence of the coding region is 47.5 kD, and the induced protein migrates on SDS–polyacrylamide gels as a 63-kD protein, we examined the pGEMF1 plasmid carefully for cloning errors. Restriction analysis, DNA sequencing, and in vitro translation of runoff transcripts (data not shown) showed that the mobility of the protein as a 63-kD polypeptide on SDS–PAGE appears to be an inherent property of the protein. Indeed, several homeotic and segmentation proteins have also been observed to migrate anomalously on SDS–polyacrylamide gels [Carroll and Scott 1985; Ollo and Maniatis 1987; U. Walldorf, P. LeMotte, and B. Dalle Carbonare, unpubl.].

Having achieved overexpression of the ftz protein in E. coli, we proceeded to work out a purification procedure. Lysis of the cells by sonication, French pressing, or treatment with detergent, followed by centrifugation consistently pelleted essentially 100% of the induced ftz protein (Fig. 1B, lane 3). The protein could not be solubilized by decreasing the amount of protein made in the cell or the rate of protein synthesis. Adding nonionic detergents and varying the pH and ion composition of the lysate also failed to solubilize the protein. Only treatment with denaturing levels of urea or guanidium hydrochloride proved successful, although complete removal of these reagents by dialysis again resulted in precipitation of the protein. Similarly, >50% of the protein produced in a reticulocyte lysate in vitro was also pre-
Figure 1. Expression and purification of ftz protein in E. coli. (A) The ftz expression plasmid pGEMFl contains ftz cDNA under the transcriptional control of the T7 phage promoter [arrow]. Transcription is achieved in an E. coli lysogen by inducing expression of an endogenous T7 polymerase gene. A short sequence between the T7 promoter and the ftz-coding region contains a strong T5 phage Shine–Dalgarno sequence (S.D.) and three codons for Met, Arg, and Asp, which replace the Met codon of the normal ftz gene. Thus, the encoded protein differs from the actual ftz protein by only two amino acids. (B) SDS–gel electrophoresis of samples prepared during ftz protein purification show uninduced cells [1], cells induced with IPTG for 2 hr [2], the cellular lysate pellet [3], protein after washes and solubilization [4], and purified protein eluted from a double-stranded DNA cellulose column [5]. Molecular weight marker sizes are indicated to the left [M].

Detection and quantitation of ftz protein synthesized in Drosophila

The production of antibodies directed against ftz protein was a prerequisite in performing the experiments that follow. Antibody production was made possible by the expression of ftz in E. coli. The ftz protein, purified as described, was injected into rabbits in the presence of 2 M urea. As shown in Figure 2, Western blot analysis of ftz protein expressed in E. coli shows that the antibodies bind specifically to the 63-kD ftz protein band [Fig. 2,
Protein by immunoblotting proved to be crucial.

Figure 2. Comparison of protein synthesized in Drosophila and in heterologous systems. Samples containing ftz protein were resolved on a 12% SDS–polyacrylamide gel and represent 35S-labeled ftz protein transcribed in vitro from the pGEMFl plasmid and translated in a reticulocyte lysate [1]. Coomassie-stained ftz protein purified from E. coli [2] ftz protein purified from E. coli detected by immunoblotting [3], ftz protein from 3- to 4-hr-old embryos detected by immunoblotting [4], and ftz protein overexpressed by heat-shocking AAl embryos carrying pHS/tz, also detected by immunoblotting [5]. Protein samples in lanes M, 1, and 2 were run on the same gel; those in lanes 3, 4, and 5 were run on a different gel.

To see whether the antibodies were also capable of recognizing the Drosophila protein, embryos that were 3–4 hr of age were homogenized in loading buffer and loaded directly on an SDS–polyacrylamide gel. After transfer to nitrocellulose, ftz protein was detected by immunostaining. A weak fuzzy band that migrates somewhat slower than the purified E. coli protein was detected [Fig. 2, lane 4]. The ability to detect the Drosophila ftz protein by immunoblotting proved to be crucial in permitting the comparisons of protein synthesized in different systems. That this band corresponds to the embryonic ftz protein will be shown more conclusively later in this paper.

One of the advantages of immunoblotting is that it allows quantitation of the protein that is detected. The ftz protein identified in lane 4 of Figure 2 was generated during the period of embryogenesis when it is most abundant [see below]. By cutting out and counting the 125I-labeled band [see Methods] and comparing the value to those generated by known amounts of purified E. coli protein, a rough estimation of its abundance could be made. Because the transfer of proteins to nitrocellulose was complete [as determined by Coomassie blue staining], polyclonal antibodies were added in excess, and the antigens detected were denatured, this comparison should be valid. It is possible, however, that proteins present in the extracts or that modifications covalently bound to the ftz protein might reduce the number of antibody molecules bound per molecule of antigen. If we assume, however, that both the E. coli and Drosophila proteins are recognized with equal affinity, the amounts of Drosophila ftz protein present in lane 4 of Figure 2 is ~1.1 ng. Knowing that this amount was derived from ~4 mg [wet weight], or 320 embryos, with each embryo possessing ~2100 nuclei that express ftz protein (~35% of 6000 blastoderm nuclei), we estimate the number of ftz protein molecules per nucleus to be ~20,000. Assuming that the volume of a blastoderm nucleus is ~1.5 X 10^-13 liters [using the dimensions 4.5 X 12.5 μm and averaging the calculated volumes for a rotational ellipsoid and a cylinder] and provided that the distribution of protein within a nucleus is uniform, 20,000 molecules would correspond to a concentration of ~2 X 10^-7 M.

The Western blots show that the ftz antibodies have a high specificity and affinity for the ftz protein (0.1 ng of ftz protein is detectable). However, the levels of protein expressed during embryogenesis are very close to the threshold of detectability. The quantitation described above also indicates that 4 kg, or ~3 X 10^8, of 3- to 4-hr-old embryos would be required to yield a maximum of 1 mg of ftz protein. Considering the short half-life of the protein in intact cells (~6 min, Edgar et al. 1986), this makes purification to homogeneity a daunting task.

Overexpression of ftz in Drosophila embryos

To facilitate the isolation and characterization of ftz protein made in Drosophila embryos, we attempted to construct an expression vector that would make it possible to overexpress the protein in all tissues and at all times during embryogenesis. This was accomplished by placing a genomic DNA fragment containing the ftz gene under the control of an hsp70 heat shock promoter (see Fig. 3A) in the P-element vector pHT4 [Schneuwly et al. 1987]. The resulting plasmid pHS/tz could then be transformed stably into flies using P-element-mediated germ-line transformation (Rubin and Spradling 1982). Eight stable fly lines were derived from injections with the pHS/tz P-element construct: AA1, AA9, AA12, AA17, and AA22 are third chromosome insertions, whereas lines AA3, AA19, and AA24 are second chromosome insertions. In the stocks AA3, AA12, and AA19, pHS/tz insertions are homozygous lethal and are therefore maintained as balanced stocks.

Transformants were tested for heat shock-dependent ftz expression by three different assays. In the first assay, SDS–gel electrophoresis of extracts from heat-shocked embryos representing each transformant line was performed and analyzed by immunoblotting to detect ftz protein. In each case, a band was observed corresponding to the band detected in 3- to 4- or 8- to 10-hr-old wild-type embryos [see, e.g. Fig. 2, lane 5]. Only background levels of protein were detected in non-heat-shocked embryos and in heat-shocked wild-type embryos [data not shown]. The amount of ftz protein produced upon heat
Figure 3. \textit{ftz} overexpression in \textit{Drosophila}. [A] The P-element vector, pHS/ftz, contains a promotor fusion of the hsp70 promoter, and a genomic \textit{ftz} DNA fragment that includes \textasciitilde 100 bp of leader and 420 bp of \textit{ftz} trailer DNA [from the \textit{KpnI} to \textit{EcoRI} (RI) sites]. \textit{ftz} DNA is flanked by hsp70 leader and trailer DNA [hatched in opposite direction], and hsp70 DNA, in turn, is flanked by P-element vector DNA (solid black regions). The \textit{rosy} gene (\textit{ry}) serves as a positive marker for fly transformation. [B] Expression of pHS/ftz in a blastoderm AA1 embryo. Anterior is to the left and dorsal is up. Two- to three-hour-old embryos were heat-shocked for 15 min at 37°C and fixed after a 15-min recovery at room temperature. \textit{ftz} protein was detected in whole embryos using \textit{ftz} antibodies and fluorescein-conjugated anti-rabbit secondary antibodies. Staining is uniform and predominantly nuclear. [C] pHS/ftz induction in larvae. Because immunolocalization is not possible on whole larvae, a salivary gland dissected from a heat-shocked third instar larva is shown, \textit{ftz} protein detected by immunofluorescence is also nuclear. [D] Immunofluorescent staining of a salivary gland from a heat-shocked AA1 larva (as in C) using preimmune serum at the same dilution as in C (1:50). The weak immunofluorescent signal seen in cellular membranes in C is also visible here. The magnification used is two times that of C.

shock varied approximately twofold between different transformant lines. Maximal expression of \textit{ftz} protein was achieved by heat-shocking at 37°C for 3 hr. It was also found that, by body weight, embryos produced about twice as much \textit{ftz} protein as third instar larvae and contained fewer proteolytic degradation products. Under optimal conditions, AA1 produced \textit{ftz} protein in a 20- to 40-fold excess over levels produced in wild-type 3- to 4-hr-old embryos (cf. lanes 4 and 5). Similar values were also found using embryos derived from the heat shock \textit{ftz} transformant \textit{hsf2} [Struhl 1985]. These levels of accumulation are far below those of heat shock proteins produced under similar conditions and suggest that there are processes that limit the expression or accumulation of \textit{ftz} protein and/or transcripts.

To examine the localization of protein expression, heat-shocked embryos and larvae were also analyzed by immunofluorescence. Usually, protein was detected uniformly in all nuclei and in all tissues, although the intensity of staining sometimes varied within and between individuals after longer heat shocks. This may have been due to the method of heat shock and/or the detrimental effects of heat or excess \textit{ftz} protein on different individuals. Figure 3B shows the typical uniform distribution of \textit{ftz} protein in a blastoderm embryo. Uniform distribution was also observed in heat-shocked \textit{hsf2} embryos obtained from Gary Struhl [Struhl 1985]. The embryo shown in Figure 3B was heat-shocked for 15 min at 37°C and fixed after a 15-min recovery; it should therefore be representative of embryos that produced the reciprocal \textit{ftz} phenotype described by Struhl [1985]. Figure 3C shows the nuclear staining of \textit{ftz} protein in a salivary gland dissected from a heat-shocked third instar larva. Because some background staining of the salivary gland cellular membranes was detected, immunostaining was also performed with preimmune serum (Fig. 3D). This also shows some staining of the cellular membranes, indicating that the background detected is recognized by antibodies that were already present in the rabbit before it was injected with \textit{ftz} protein. This staining was not detected with the higher dilutions of serum that were used in subsequent experiments.

Having ascertained that all pHS/ftz-transformed lines are capable of synthesizing \textit{ftz} protein and that the pro-
tein is transported into the nucleus, we looked for an indication that protein synthesized in this manner is functional. This was tested by examining the cuticular phenotype of pharate first instar larvae that were heat-shocked at 37°C between 2.5 and 3.5 hr of development. As stated earlier, such treatment has been shown previously to produce a near-reciprocal deletion of segments in a complementary fashion to those deleted in $ftz^{-}$ embryos [Struhl 1985]. We also obtained similar results using our pHStz transformant embryos [data not shown]. Although this does not show whether or not the protein expressed from the hsp70 promoter can perform the same functions as the endogenous protein, it indicates that it somehow interferes either with the distribution of other segmentation proteins or their ability to direct normal segment boundary formation in regions where $ftz$ is not normally expressed.

In summary, these three tests show that the $ftz$ protein can be overproduced under the control of a heat shock promoter, that it is a nuclear protein, and that it is probably capable of some, if not most, of the endogenous $ftz$ protein functions.

**Post-translational modification of $ftz$ protein**

As pointed out earlier, $ftz$ protein made in *Drosophila* embryos migrates more slowly on SDS–polyacrylamide gels than protein expressed in vitro or in *E. coli*. This suggests that the protein contains covalently bound modifications. To obtain more information about the nature of these modifications, protein samples were resolved further using two-dimensional PAGE [Fig. 4]. Samples of $ftz$ protein made either in *E. coli* or in embryos were separated first by charge on isoelectric focusing gels and then by molecular weight on SDS–polyacrylamide gels. The $ftz$ protein was then detected by Western blotting. Figure 4A shows that $ftz$ protein expressed in *E. coli* predominantly forms a single spot. In addition, there is a weak spot shifted slightly to the acidic side of the gel and, on blots that have been developed longer, a third very faint spot can be detected even farther to the right. The pattern generated by purified protein stained with Coomassie blue is identical, indicating that all of these spots represent forms of the $ftz$ protein and that some modification occurs in bacterial cells.

In contrast, when $ftz$ protein was expressed uniformly throughout the embryo by heat-shocking pHStz embryos, the protein was resolved into a series of at least 11 distinct isoforms [Fig. 4B]. These increase in negative charge and apparent molecular weight with respect to the major species made in *E. coli*. The same pattern was observed using other pHStz transformant embryos and is also essentially the same in heat-shocked pHStz third instar larvae [data not shown]. The pattern seen in Figure 4B was generated from embryos collected overnight, aged for 2 hr, and then heat-shocked for 3 hr. Thus, the mixture of isoforms represented by this pattern is the average of all isoforms produced throughout the embryo between 5 and 18 hr of embryogenesis.

**Temporal and spatial expression of $ftz$ protein during embryogenesis**

Earlier, it was shown that the $ftz$ antibodies detect a weak band on Western blots of total embryonic extracts [Fig. 2, lane 4]. To demonstrate more conclusively that the protein detected is $ftz$ and that the antibody is specific for this protein, we examined variations in its abundance during different periods of embryogenesis using a ‘developmental’ Western blot [Fig. 5]. In addition, we used immunological localization of $ftz$ protein on whole embryos to determine whether or not changes in abundance of the protein correlate with changes in
expression of protein in wild-type embryos. Embryos collected for 1-hr (0–10 hr) or 2-hr (10–16 hr) intervals were weighed and lysed in 4 μl/mg SDS–lysis buffer. The equivalent of 4 mg of embryos was loaded in each lane, separated by SDS–PAGE, and blotted to nitrocellulose. ftz protein was detected using ftz antibodies and 125I-labeled protein A. Time intervals indicating the ages of collected embryos are given, as are the amounts of ftz protein purified from E. coli used as size and quantitation standards.

The developmental Western blot was made using equivalent numbers of embryos collected and aged so that 1- or 2-hr intervals of embryogenesis are represented. Examination of Figure 5 shows that the band detected previously in Figure 2 (lane 4) first appears during the 3- to 4-hr interval of embryogenesis. It then subsides and reaches a second peak during the 8- to 9-hr interval. After 10 hr of development, the protein abundance drops substantially. Upon overexpression of the blot, a band can still be detected during the 12- to 14-hr interval (<0.1 ng protein). No protein is detectable after 14 hr. Hence, the expression of ftz protein is in agreement with the fluctuations of ftz mRNA reported previously, except that the protein is in higher relative abundance during the 8- to 10-hr interval than the message from which it is derived (see Kuroiwa et al. 1984; Weiner et al. 1984). The overall abundance of protein at 8–9 hr is ~75–80% of the amount present during the 3- to 4-hr interval (based on several repetitions of the experiment and quantitation of bands). In contrast, the amount of message at 8–10 hr is ~10–20% of the abundance at 2–4 hr (Kuroiwa et al. 1984; Weiner et al. 1984). The nuclear concentration of ftz protein during the 8- to 10-hr interval was not determined because both the numbers and sizes of nuclei expressing ftz in the developing nervous system vary rapidly with time (see Fig. 6D–G).

Immunofluorescence on whole embryos shows that the two peaks of ftz protein expression take place in different locations (as described previously by Carroll and Scott 1985). The embryos shown in Figure 6, A–C, demonstrate that ftz protein present between 3 and 5 hr of development is localized in seven zebra-like bands that encircle the embryo. We noted consistently that the bands appear in the following order: 2, 1 + 3, 5, 6, 7, and, finally, 4. This order of appearance is reflected by the relative intensity of each stripe, as seen in the embryo in Figure 6A (stripes are arbitrarily numbered from anterior to posterior). At this point, it can also be seen that there is a gradient of protein abundance across the stripes; nuclei at the edges of the bands stain very weakly, whereas those in the center produce the strongest signal. This bell-shaped distribution is resolved quickly into stripes with well-defined borders in which all nuclei apparently possess the same level of ftz protein and are flanked by nuclei that express no detectable protein (Fig. 6B).

The bands are dynamic in width and position, as discussed previously by Carroll and Scott (1985). In Figure 7B, nuclei on the posterior edges of most stripes that have intermediate levels of ftz protein abundance can be seen. Carroll et al. (1988) have interpreted this to mean that subsequent stripe narrowing occurs predominantly by loss of ftz protein in the cells representing the posterior edges of each stripe. Lawrence et al. (1987) arrived at the same conclusion previously by showing that the anterior edges of each ftz stripe remain lined up with engrailed stripes during the period of time that engrailed and ftz are coexpressed. We must point out, however, that this only appears to be true for stripes 1–5, because stripe 6 remains more or less constant at three to four cells in width, and stripe 7 becomes wider rather than narrower (see Fig. 6C). Because the cells are dividing during this time, the maintenance of stripe 6 at three to four cells in width is likely to involve a balance between the process of stripe narrowing due to loss of ftz expression in certain cells and widening due to the division of other ftz-expressing cells.

After completion of germ-band extension, expression of ftz protein becomes difficult to detect. Then, beginning at ~5–6 hr of development, when the stripes are no longer detectable, staining of distinct neuronal precursor cells in the developing nervous system becomes visible in each segment (Fig. 6D). These increase in number and eventually give rise to a large number of specifically labeled neurons. The pattern of expression in specific cells and the role of ftz protein in these cells is being studied elsewhere (see Doe et al. 1988). Localization of ftz protein in the blastoderm and nervous system confirms data reported previously (Carroll and Scott 1985).

Upon examination of older embryos, a third stage of expression was detected that has not been described previously (Fig. 6H; Fig. 7). This expression appeared dorsally at about 40% egg length as a small ring or tube in 12- to 15-hr-old embryos and was most prominent be-
Figure 6. Immunofluorescence localization of ftz protein expressed during embryogenesis. Staged embryos were incubated with anti-ftz antibodies, and a swine, anti-rabbit secondary antibody coupled to fluorescein. Embryos are oriented with the anterior end to the left and ventral surface down, except for D–F, which are viewed from the ventral side. Based on a 21-hr period of embryogenesis (at 25°C) embryos are estimated to be ~2 3/4 hr [A], 3 hr [B], 4 hr [C], 6 hr [D], 7 hr [E], 9 hr [F], 10 hr [G], and 14 hr old [H]. [A–C] Stages of pair-rule stripe expression; [D–G] nervous system expression; [H] hindgut expression. After 9 hr of development, autofluorescence of the yolk proteins present in the developing midgut can be detected and differs from the fluorescein signal in that it is yellow rather than green.
Figure 7. \(ftz\) protein expressed in the hindgut. \(ftz\) protein was detected using either fluorescein-coupled \(F,G\) or peroxidase-coupled \(A-E, H\) secondary antibodies. Lateral views are shown, with dorsal at the top and anterior to the left. (A) A 12-hr embryo, hindgut expression is just beginning. (B) A 13-hr-old embryo. In addition to the hindgut staining, regions of expression are indicated in the anterior and posterior regions of the embryo (arrows). (C) A 14-hr-old embryo. (D) A 15-hr-old embryo, showing widening and weakening of the hindgut site of expression. (E) A higher magnification of the hindgut expression site, showing nuclear localization. (F) Fluorescent staining of a 14-hr-old wild-type embryo. (G) A 14-hr-old \(ftz^{H154}\) embryo, which does not express \(ftz\) protein [Carroll and Scott 1985]. (H) A diagrammatic interpretation of a 14-hr-old embryo showing relevant features. The site of \(ftz\) expression in the hindgut is indicated as a solid black ring. (M.G.) Midgut; (M.T.) malpighian tubules; (H.G.) hindgut.
stain light yellow. The major site of expression, seen as a stained ring when viewed in cross section, coincides with a section of the hindgut just posterior to the site of malpighian tubule attachment (see Fig. 7H). At this point, the hindgut makes a major bend, with one end looping back to the posterior end of the embryo and the other extending past the malpighian tubules to the point where it fuses with the midgut. Expression begins as a short tube (Fig. 7A) that narrows during the peak of expression (Fig. 7B,C) and then widens once more as expression drops off (Fig. 7D). As seen in the previous two phases of ftz expression, protein localization also appears to be nuclear (Fig. 7E). Similar staining patterns were obtained using antibodies from two different rabbits. Staining was not observed when preimmune serum or protein-negative ftz- embryos were used (Fig. 7F,G).

Discussion

Synthesis of ftz protein in heterologous systems

Expression of ftz protein was achieved in an in vitro transcription–translation system and in E. coli using the T7 expression system. In both cases, the protein migrated anomalously on SDS–gels to approximately the same position, corresponding to an apparent molecular weight of 63,000. Protein produced in either system is insoluble and can only be solubilized and maintained in solution in the presence of urea or guanidium hydrochloride. This tendency of ftz protein to aggregate is probably an inherent property of the protein’s amino acid composition and sequence and suggests that in Drosophila, the protein must be folded differently, modified, or maintained in solution by associating with other factors. Indeed, no indication of insolubility has been encountered during partial purification of ftz protein from Drosophila embryos (H.M. Krause and W.J. Gehring, in prep.).

Analysis of the ftz DNA sequence predicts an unusually high abundance of proline, serine, threonine, and tyrosine residues (10%, 10%, 10%, and 9%, respectively). The first three are characteristic components of ‘PEST’ sequences [P = proline, E = lysine, S = serine, T = threonine], which have been found in a large number of eukaryotic proteins that are degraded rapidly (see Rogers et al. 1986). One could also imagine that repetition of these amino acids, the prolines in particular, could create defined kinks or bends in the secondary structure of the protein. These might facilitate protein aggregation and possibly inhibit quantitative binding and denaturation by SDS, thus affecting mobility in SDS–polyacrylamide gels as well.

Rapid degradation of the ftz protein in E. coli may have been the decisive factor that resulted in the failure of all but the T7 expression system to produce detectable levels of protein. The developers of the system claim that transcription and translation of the T7 polymerase transcribed gene in bacterial cells monopolizes all of the available resources. Thus, in the case of a protein that is extremely labile, this may preclude a host response sufficient to remove the levels of protein being synthesized. In addition, the T7 RNA polymerase is present in higher abundance, elongates more rapidly, and is less sensitive to termination than the E. coli RNA polymerase. Even so, ftz protein accumulation was far below the levels that have been reported for other proteins expressed in this system.

ftz antibodies are specific for ftz protein

ftz protein purified from E. coli, as described, was used to generate antibodies in rabbits. That the protein used to generate antibodies is ftz and that the antibodies recognize this protein as well as the Drosophila-made ftz protein, is shown by mutually agreeing experiments. On Western blots, the antibodies recognize only the 63-kD band of an induced E. coli lysate and a slightly larger, fuzzy band of higher apparent molecular weight in extracts from wild-type embryos. Peaks in the abundance of this band correlate in time with the results of immunological staining of embryos. The antibody detects seven stripes of protein located in the ectodermal nuclei of 3- to 5-hr-old embryos and the nuclei of neuronal precursors in the developing nervous system of 6- to 12-hr-old embryos. These results are also consistent with previously reported Northern blots on which ftz mRNA can be detected at similar times. A comparison of Northern and Western blots, however, shows that ftz protein is higher in relative abundance during nervous system expression than the mRNA from which it is derived. This, taken together with the fact that mRNA expression in the nervous system was not detected initially by in situ hybridization and that the protein is easily detectable by immunofluorescence, suggests that the efficiency of ftz mRNA translation or protein stability must be significantly higher in the nervous system.

It was found that rabbits injected with ftz protein consistently produced antibodies with high specific affinity for native, as well as denatured, ftz protein and generally showed very low background. Immunological staining of whole embryos produced essentially the same results using serum dilutions ranging from 1 : 100 to 1 : 1000, indicating a high titer of ftz-specific antibodies. No difference in staining patterns was detected using ftz antibodies that were purified over a ftz protein affinity column. It is possible that the high titer and specificity
of the rabbit sera may be partially attributable to the use of a purified and full-length, nonfusion protein that was injected in a seminative state to generate these antibodies.

The heat shock–ftz promoter fusion

Although engrailed and ftz proteins synthesized in E. coli bind DNA in a sequence-specific manner, the binding sites are numerous and variable in sequence (Desplan et al. 1985; Krause and Gehring 1988, and unpub.). Because problems in the solubility of these proteins were also observed, we wished to identify and compare the ftz protein made in Drosophila embryos with protein made in heterologous systems to see whether biochemical differences could be detected. It became obvious that normal expression of the protein in embryos was at the threshold of detection on immunoblots. A ftz—heat shock promoter fusion was therefore constructed so that elevated levels of ftz protein could be synthesized in Drosophila embryos. Use of the hsp70 heat shock promoter has the additional advantage of allowing temporal control of induction. By heat shocking pHS/ftz transformants, it was found that synthesis of ftz protein could be induced at all times during development and in all tissues examined, except perhaps prior to 2 hr of embryogenesis. Furthermore, the protein migrates to the nuclei of these cells as it does during normal expression in wild-type embryos and comigrates with the endogenous protein on SDS–polyacrylamide gels. In previous experiments, Struhl (1985) has shown that ftz protein expressed from the hsp70 promoter by heat shock during the blastoderm period of development disrupts the normal pattern of segmentation. Here, we have shown that heat shock does indeed produce ftz protein in his and our transformants and that the protein is distributed uniformly throughout the blastoderm nuclei as expected. Maximal levels of ftz protein produced by these transformants were less than expected, being only ~20–40 times higher than in 3- to 4-hr-old wild-type embryos. This is not much more than the value that might be expected if one considers that the whole volume of 5- to 18-hr-old embryos (the usual age after overnight collection, staging, and heat shock) is capable of expressing ftz protein rather than just the 2100 ftz-expressing nuclei that encircle the periphery of a normal blastoderm embryo. We have found that the ftz protein is extremely labile, both in E. coli and Drosophila, and this may quite possibly be the major factor that inhibits further accumulation in these cells. Edgar et al. (1986) have estimated that the half-lives of ftz mRNA and protein are ~10 and 6 min, respectively, during the blastoderm stage of expression. In contrast, under similar heat shock conditions, the various heat shock proteins can each accumulate to values approaching 1% of total cellular protein (Ashburner and Bonner 1979; Dura 1981). We estimate that the amount of ftz protein maximally produced is about 2 orders of magnitude lower. That proteins involved in embryonic development have a rapid turnover rate is perhaps not surprising as changes in developmental states must occur very rapidly to complete embryogenesis in a mere 21 hr.

ftz protein is covalently modified in Drosophila

On two-dimensional gels, ftz protein made in E. coli migrates predominantly as a single spot. The pH of the isoelectric focusing gels at this position is ~7. This value is in agreement with the pl value of 6.8 calculated for the protein based on its amino acid sequence. A very weak spot that comigrates with this one can also be detected in extracts from heat-shocked pHS/ftz transformant embryos. Because all other spots detected are increasingly negatively charged, this spot probably represents the unmodified form of the protein. Interestingly, two weak spots were also seen on two-dimensional gels, both before and after purification of the bacterially expressed protein. Thus, it appears that E. coli possesses the ability to modify the ftz protein covalently, although the percentage of modified protein and the number of modifications on each protein molecule were minimal under these conditions of expression. Because these isoforms of the protein also run more slowly in the SDS–PAGE dimension, it is unlikely that they are generated by the removal of amino acids due to proteolytic degradation. Protein translated in vitro in a reticulocyte lysate comigrated with the protein synthesized in E. coli, suggesting that it is also not modified extensively. In vitro-translated protein was not examined on two-dimensional gels.

ftz protein made in either wild-type Drosophila embryos or by heat shock in pHS/ftz-transformed embryos comigrated on SDS–polyacrylamide gels as broad bands which run more slowly than protein made in either of the two heterologous systems. When the protein overexpressed by heat shock was resolved on two-dimensional gels, at least 11 distinct spots could be detected. These appear to represent isoforms of the ftz protein which possess increasing numbers of covalently bound negative modifications as they become shifted from left to right on the two-dimensional gels. We have also detected complex patterns of spots in wild-type embryo extracts at stages when the protein is expressed maximally and also when expressed under the control of a heterologous promoter in the flight muscles of adult flies (H.M. Krause and W.J. Gehring, in prep.). Thus, it is unlikely that the process of heat shock is responsible for the creation of these isoforms. However, it may have an influence on the number and nature of modifications made to the protein. Several types of post-translational modifications could produce the type of pattern observed. The results of further experiments indicate that the process giving rise to these multiple isoforms is protein phosphorylation (H.M. Krause and W.J. Gehring, in prep.).

Post-translational modification of the ftz protein during embryogenesis could conceivably alter the regulatory potential of the protein by altering various biochemical properties. For example, the tendency of the unmodified protein made in heterologous systems to ag-
aggregate could be modulated by the addition of negatively charged groups to hydrophobic regions of the protein. In key positions, they could also alter the ability of the protein to bind DNA sequence specifically or to interact with other proteins such as transcription factors. Because the embryonic pattern of ftz protein produced on two-dimensional gels represents protein expressed during a great portion of embryogenesis, and throughout the embryo, it will be interesting to see whether distinct subpopulations of the observed isoforms are localized to specific regions during different stages of ftz expression.

Three phases of expression during embryogenesis

The high affinity of our ftz antibodies permitted the observation of very early and late phases of the segmentation and nervous system periods of ftz expression. In situ hybridization to ftz mRNA shows that the message is first detectable in a fairly uniform distribution over the seven-stripe domain (Hafen et al. 1984). This pattern then becomes resolved into the seven stripes reflected by the RNA (Hafen et al. 1984; Weir and Kornberg 1985; Edgar et al. 1986) and protein distribution (Carroll and Scott 1985). The earliest unambiguous expression of ftz protein that we detected is in a striped pattern at late blastoderm stages. The appearance of each stripe occurs in a consistent, defined order, as represented by the different stripe intensities of the fixed embryo in Figure 6A. This intermediate pattern can also be seen in the intermediate patterns of ftz transcript distribution (Hafen et al. 1984; Weir and Kornberg 1985; Edgar et al. 1986). Embryos that exhibited a uniform or double-wide intermediate stripe distribution were not observed.

Antibody detection at these initial stages of expression demonstrates the occurrence of rapid changes in stripe intensity and definition. At the onset of detection, nuclei at the edges of the stripes contain lower levels of ftz protein than those in the center. The quick transition to a state where nuclei at the edges of the bands either do or do not contain equivalent amounts of ftz protein is compatible with the models proposed by Meinhardt and Gierer (1980, Meinhardt 1986). To fit these models, the ftz protein must be capable of autoregulating its own expression positively, whereas segmentation gene products expressed in adjacent and initially overlapping stripes must repress ftz expression. The ratio of ftz protein concentration to other competing segmentation proteins in a given nucleus, therefore, determines whether or not ftz becomes maximally expressed or is turned off. Only one of the competing genes would eventually be expressed in each nucleus, and the stripe borders would thus become sharply defined. Because the sharpening of the initial striped pattern occurs after cellularization, it appears that this resolution process occurs cell autonomously. Evidence for positive autoregulation by ftz protein and repression of ftz expression by other segmentation genes has been obtained through the use of ftz promoter fusions to the β-galactosidase gene (Hiromi et al. 1985; Hiromi and Gehring 1987) and immunodetection of the protein expressed in various mutant backgrounds (Carroll and Scott 1986). After the initial resolution of the stripes, stripes 1–5 become narrower and stripe 7 becomes wider. This appears to occur by the removal or addition of ftz-expressing nuclei at the posterior edges of the stripes, as demonstrated previously by Lawrence et al. (1987) and Carroll et al. (1988).

Comparisons of the abundance of ftz protein during this first phase of expression with known amounts of E. coli protein on immunoblots permitted us to make a rough estimate of the concentration of ftz protein in the nuclei in which the gene is expressed. This abundance is comparable to levels of other eukaryotic regulatory proteins. For example, the enhancer-activating proteins AP-1 and AP-2 also have an estimated abundance of 20,000 molecules per cell (Lee et al. 1987; Mitchell et al. 1987). These numbers can also be compared with levels of prokaryotic regulatory proteins. The λ repressor, for example is present at ~100–200 copies per lysogen (Ptashne 1986). Because Drosophila blastoderm nuclei are ~150 times the volume of a bacterial cell (150 μm^3 vs. 1 μm^3), this abundance would also be equal to ~20,000 molecules per nucleus. Indeed, the estimated concentration of λ repressor in a lysogen is the same as our estimated value for the concentration of ftz protein in blastoderm nuclei (~2 × 10^-7 M).

Nervous system expression of ftz follows soon after the stripes have disappeared. The protein is first detectable between 5 and 6 hr of development in the midline precursor-2 [MP-2] cells (Doe et al. 1988) in a segmentally repeating pattern. Expression in many other identified neuronal precursors follows and reaches a second peak of abundance at ~9 hr of development. At the end of germ-band retraction, the protein disappears. Although it is not yet clear what the role of the ftz protein is during the development of the central nervous system, it has now been shown that failure to express ftz leads to a change in cell identity in the case of at least one pair of segmentally repeated neurons (Doe et al. 1988).

Following nervous system expression of the protein, a previously undetected phase of expression was observed. This phase begins at ~11.5 hr of development and persists until approximately the fifteenth hour of development. This is the only nonsegmental expression of ftz protein during embryogenesis and is predominantly localized to a section of the hindgut just posterior to the attachment site of the malpighian tubules. After the peak of expression at 13 hr, the ring seems to widen into a tube that partially overlaps the major bend in the hindgut (diagramed in Fig. 7H). As the embryo develops further, the signal disperses and weakens. In addition to this site of expression, protein was usually detectable at the posterior, dorsal surface of the embryo and in a few cells in the foregut. Again, as in the previous phases of expression, the protein localization is nuclear. Although no protein can be detected on the developmental Western blot at this time in development, examination of the Northern blots of Weiner et al. (1984) shows that some ftz mRNA is still present at low levels in 12- to
14-hr-old embryos. The band corresponding to ftz protein in the developmental Western blot was also detectable at this stage of development.

Several other lines of evidence are consistent with the assumption that the protein detected at this stage is ftz. Hindgut expression was detected with ftz antibodies produced in two different rabbits, either before or after affinity purification of ftz antibodies and not with preimmune serum from these same rabbits. Also, in homozygous ftz29H64 embryos, which are ftz protein negative (Carroll and Scott 1985), the hindgut was usually present, but immunostaining was absent. The possibility that the signal is due to cross-reactivity with another protein bearing similar epitopes is also unlikely because the antibodies appear to recognize only the ftz protein on immunoblots.

Further evidence that ftz is expressed and has a function in the hindgut is given in a study by Wakimoto et al. (1984). Flies carrying a temperature-sensitive ftz allele were heat-shocked at different times during embryonic development. Decreased viability was observed when heat shocks were administered before 4 hr, ~8 hr, and again at 12 hr of development. It might also be mentioned that expression of the genes hairy, engrailed, Ultrabithorax, and caudal have been detected in parts of the midgut or hindgut during similar periods of embryogenesis (Akam and Martinez-Arias 1985; DiNardo et al. 1985; Fjose et al. 1985; Ingham et al. 1985; Kornberg et al. 1985; Mlodzik et al. 1985).

In light of these data, it seems that certain homeotic and segmentation genes not only play a role in the process of segmentation but in nervous system and gut development as well. It is not immediately obvious what the common requirement for ftz protein might be in such functionally diverse tissues. It is possible that each tissue responds to ftz expression in a similar way or that each tissue or cell interprets the presence of the protein in an individual fashion. For example, during different phases of expression, different subsets of genes might be regulated by the ftz protein due to such variations as the presence or absence of different coregulatory factors or alterations in chromatin structure. Already, there is evidence that at least some of the genes that regulate ftz and others that are regulated by ftz vary during the blastoderm and nervous system periods of expression (Hiromi et al. 1987; Dodge et al. 1988). In this respect, it will be interesting to see whether protein modification may also play a role in determining which genes are controlled by ftz during different phases of development.

**Methods**

**Plasmids**

The expression of ftz protein in E. coli necessitated the construction of the plasmid pGEMF1 [Fig. 1A]. This was made by subcloning an EcoRI fragment from a previously constructed plasmid into the pGEM-1 plasmid (Promega Biotec), using standard cloning techniques (Maniatis et al. 1982). The donor plasmid, pDSBH2, was constructed and kindly provided by Johannes Wirz. The construction of the pDSBH2 plasmid involved a series of cloning steps in which ftz genomic and cDNA restriction fragments isolated from various plasmids were ligated together to generate the complete reading frame of the protein with the intron deleted. In the final step, a 1590-bp Ball–EcoRI ftz restriction fragment was inserted into the expression plasmid pDS6 (Stuber et al. 1984), using the BamHI and EcoRI sites of that plasmid. To make pGEMF1, an EcoRI fragment that includes 30 bp of the original pDS6 plasmid was subcloned from pDSBH2 into pGEM-1 such that the T7 phase Shine–Dalgarno sequence present in that plasmid is also utilized in the new construct. In the pGEMF1 plasmid, the orientation of the EcoRI fragment places the inserted ftz gene under the control of the T7 promoter, whereas it can be transcribed from the SP6 promoter in a second control plasmid, pGEMF2.

The pHStz construct was made by inserting a 1.8-kb KpnI–EcoRI restriction fragment from the plasmid pHyHF (obtained from Y. Hiromi) into the modified P-element vector pH74 (for a description of pH74, see Schneuwly et al. 1987). In addition to the complete ftz-coding region, the KpnI–EcoRI ftz fragment contains ~100 bp of 5′ ftz leader (to the Sau3A site) and 400 bp of 3′ trailer sequences. Sticky ends were filled in with T4 DNA polymerase [Boehringer] and blunt-end-ligated into the unique KpnI site of pH74.

**Purification of ftz protein from E. coli**

The plasmid pGEMF1 was first transformed, characterized, and maintained in the E. coli strain ED8739 in which T7 RNA polymerase-directed ftz expression can be induced by infection with genetically engineered λ phage (Studier and Moffatt 1986). For maximal expression, the plasmid was transformed into the BL21 : DE3 lysogen, in which induction is achieved by addition of IPTG. Bacterial growth and vector induction were performed essentially as described by Studier and Moffatt (1986). Cells were grown for 2 hr after the addition of 2 mM IPTG at a cell density corresponding to an OD600 reading of 1.0.

Cells from 1 liter of culture were pelleted by a low-speed spin and resuspended in 5 ml of 25% sucrose, 50 mM Tris–HCl (pH 7.5). Cells were then frozen overnight at −20°C. The following steps were carried out on ice or at 4°C and in the presence of the following protease inhibitors: 0.1 mM phenylmethylsulfonyl fluoride [PMSF] [Sigma], 5 μg/ml leupeptin [Fluka], 0.2 μM/ml kalikrein [CalBiochem], and 2 μg/ml pepstatin [Fluka]. Cells were thawed and resuspended in an equal volume of lysis buffer [100 mM Tris–HCl (pH 7.5), 4 mM EDTA, 2 mM dithiothreitol [DTT], and 2 mg/ml lysozyme] and incubated for 15 min. This was followed by sonication in four 15-ml Falcon tubes for 15 sec, repeated four times with not less than 1 min between repetitions. The lysate was spun at 14,000 rpm for 10 min in a Sorvall SS-34 rotor, and the pellet was washed twice by resuspending in 20 ml resuspension [R] buffer 10 mM Tris–HCl at pH 7.5, 100 mM NaCl, 1 mM EDTA, 10% glycerol], first containing 1% Triton X-100 and then 3 mM potassium thiocyanate. Each time, the mixture was resuspended using a Dounce homogenizer, and repelleted as above. Finally, ftz protein was solubilized by resuspending in 10 ml R buffer containing denatured 8 M urea, and after adding an equal volume of R buffer-urea insoluble material was removed by centrifugation for 10 min at 14,000 rpm in a Sorvall SS-34 rotor. The supernatant was then passed through a 1.5 × 10-cm DEAE–cellulose [Cellex D, Bio-Rad] column and dialyzed against R buffer containing 50 mM NaCl and 2 mM urea. Precipitated protein was again removed by centrifugation, and the supernatant was loaded on a 1 × 10-cm double-stranded DNA cellulose [Sigma] column. Bound protein was eluted with a 50–500 mM NaCl linear gradient in R buffer containing 2 mM urea. The protein was identified by SDS–
PAGE and eluted between 175 and 225 mM NaCl. Fractions containing ftz protein were pooled, and the majority were stored at 4°C for immediate studies. Aliquots were also prepared and stored at −70°C.

**In vitro transcription–translation**

RNA was transcribed in vitro from the plasmids pDSBH2 and pASF1 (in this plasmid, ftz is under the control of the a P1 promoter in the vector pA5-1; Mott et al. 1985), using *E. coli* RNA polymerase, or from the plasmid pGEMF1 using T7 RNA polymerase. Both enzymes were obtained from Boehringer Mannheim. In vitro transcription was carried out according to the procedure of Hurt et al. [1984]. Proteins labeled with [35S]methionine (Amersham) were translated in a rabbit reticulocyte lysate (kindly provided by G. Schatz, Biozentrum, University of Basel) were separated by SDS–PAGE and detected by autoradiography following fluorography and Dupont intensifying screens.

**Antibody preparation, purification, and manipulations**

Antibodies were generated in rabbits against full-length, native ftz protein purified as described above. ftz protein (250 µg) in 250 µl R buffer containing 2 M urea was mixed with an equal volume of complete Freund's adjuvant and injected subcutaneously at multiple sites. Secondary boosts were performed similarly with 50 µg of protein every month for 4 months. Rabbits were bled every 2–4 weeks until final exsanguination. Western blot analysis and immunolocalization of ftz protein in embryos was generally performed with crude serum. As a control, antibody was purified further either by adsorption against 16- to 20-h-old embryos or by affinity purification on a ftz protein–Affi-Gel column. Preadsorption was achieved by incubating 200 µl of crude serum with 0.5 ml of fixed 16- to 20-h-old embryos in a total of 1 ml of whole-mount solution A [1× PBS, 0.05% Saponin, 0.05% Triton X-100, 0.05% NP-40, 0.05% deoxycholate (DOC), 2 mg/ml bovine serum albumin (BSA), and 0.02% NaN3] overnight at 4°C with end-over-end rotation. Embryos were subsequently removed by a 1-min spin in an Eppendorf centrifuge.

For affinity purification, 2 mg of ftz protein purified from *E. coli* was dialyzed against 10 mM morpholinopropanesulfonic acid (MOPS) pH 7.0, 100 mM NaCl, 1 mM DTT, and 2 M urea. Preparation of Affi-Gel 10 [Bio-Rad] and coupling of protein were performed as described by the manufacturer. Coupling was allowed to proceed for 4 hr at 4°C with a 1 : 1 vol/vol ratio of ligand to Affi-Gel. Crude serum (20 ml) was passed over a 1-ml ftz protein–Affi-Gel column, and bound antibody was eluted with 0.2 N glycine–HCl (pH 2.8). The eluate was neutralized immediately with Tris–HCl (pH 8.0) and dialyzed against PBS.

Embryos were prepared for immunolocalization using a variation of the protocol described by Mitchison and Sedat [1983]. Embryos were dechorionated for 90 sec in a 3% Clorox solution and washed extensively. Fixation was carried out for 30 min at room temperature with moderate shaking in equal parts of heptane and a solution containing 100 mM PIPES (pH 6.9), 2 mM EGTA, 1 mM MgSO4, and 5% formaldehyde. The lower aqueous phase was then removed and replaced by an equal volume of methanol. Vitelline membranes were removed by shaking vigorously for 3–5 min. Intact embryos were allowed to settle, and all liquid was removed. This was followed by three methanol washes and resuspension in whole-mount solution A (described above). Embryos were stored for no longer than 2 days before use. If used immediately, embryos were incubated for 30 min in whole-mount solution A with end-over-end rotation at room temperature. Antibody binding was carried out in whole-mount solution B [1× PBS, 0.1% Triton X-1200, 2 mg/ml BSA, 5% swine serum, 0.02% NaN3] for 2 hr under the same conditions. ftz antibodies were used at a dilution of 1 : 500. Swine anti-rabbit antibody coupled to fluorescein (Dako) was generally used as the secondary antibody at a 1 : 100 dilution. When peroxidase coupled to swine anti-rabbit serum (Dako) was used as the secondary antibody, washed embryos were stained in PBS containing 20 µg/ml diaminobenzidine (Sigma) and 0.003% hydrogen peroxide. The reaction was stopped by washing in PBS and followed by dehydration in washes of 70%, 90%, and 100% ethanol. The embryos were then resuspended in DePeX mounting medium (Gurr) and mounted on a glass slide under a coverslip.

**Characterization of ftz protein modifications**

Samples of ftz protein made in *Drosophila* embryos were prepared for one- and two-dimensional polyacrylamide gel analysis simply by homogenizing embryos directly in the appropriate loading buffer. The PHF7ftz transformant lines AA1–AA24 were generated by P-element-mediated germ-line transformation using standard techniques (Rubin and Spradling 1982), with the recipient fly line R-3 [ry606, st, th, e]. Embryos or larvae were collected and cleaned before heat shock in a 37°C incubator in a moist container. Embryos were dechorionated, blotted dry, and weighed. For one-dimensional gels, a minimum of 50 mg of staged embryos were homogenized and boiled for 5 min in 4 µl/mg SDS–lysis buffer and spun for 2 min in an Eppendorf centrifuge to remove insoluble material. The equivalent of 4-mg embryos (wet weight) were loaded on SDS–polyacrylamide gels. Western blots were made by transferring proteins separated by SDS–PAGE to nitrocellulose, as described by Towbin et al. [1979]. Nonspecific binding of antibodies was blocked with BLOTTO (Johnson et al. 1984). ftz antibodies were bound at room temperature over a 2-hr period at a dilution of 1 : 1000 and visualized by labeling with 35S-labeled protein A and autoradiography using prefished film and intensifying screens. Sections of the blot containing ftz protein were then cut out and counted in scintillation fluid. For two-dimensional analysis of proteins (O'Farrell 1985), embryos were lyzed directly in 4 µl/mg isoelectric focusing lysis buffer. After SDS–PAGE in the second dimension, immunoblotting was carried out as described above, except that the second antibody used was swine anti-rabbit coupled to peroxidase (Dako). Enzymatic development was performed as described above for the whole-mount staining of embryos.

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