Protamine 3'-untranslated sequences regulate temporal translational control and subcellular localization of growth hormone in spermatids of transgenic mice

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Although the mouse protamine 1 gene (mP1) is first transcribed in round spermatids, its mRNA is not translated until about 1 week later in elongating spermatids. To determine what mP1 sequences are important for its transcriptional and translational regulation, we have constructed fusions between mP1 and the human growth hormone (hGH) structural gene and analyzed their expression in transgenic mice. We show that mP1 sequences 5' to the start of transcription are sufficient to confer spermatid-specific expression on the hGH gene. We also show that 156 nucleotides of mP1 3'-untranslated sequence is sufficient to confer mP1-like translational regulation on the hGH mRNA. Interestingly, the subcellular localization of hGH was dependent on the time during spermiogenesis that it was made. Synthesis of hGH in early round spermatids resulted in localization in the acrosome, whereas synthesis in late elongating spermatids resulted in intracellular, but not acrosomal, localization.

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During the haploid stage of spermatogenesis, called spermiogenesis, round spermatids differentiate into spermatozoa (Bellve 1979; Hecht 1986). Among the many morphological changes that occur are the reshaping of the nucleus (Fawcett et al. 1971; Dooherr and Bennett 1973), the development of a flagellum (Fawcett and Phillips 1969), and the construction of the acrosome (Fawcett et al. 1971), a secretory vesicle located on the apical surface of the nucleus that is required for egg penetration during fertilization (Wassarman 1987). In the mouse, spermiogenesis lasts ~2 weeks and has been described as consisting of 16 stages based on various morphological criteria associated with changes in nuclear morphology and acrosomal development (Oakberg 1956).

The mouse protamine 1 (mP1) gene is one of two genes that encodes small basic proteins that replace the histones and testis basic proteins during sperm nuclear condensation (Bellve 1979). Protamine 1 expression is under both transcriptional and translational control. The gene is transcribed exclusively in haploid round spermatids (Hecht et al. 1986). mP1 mRNA is stored as a cytoplasmic ribonucleoprotein particle (mRNP) for up to 1 week before it is translated in elongating spermatids (Balhorn et al. 1984; Kleene et al. 1984).

Translational control is an important mechanism of regulation during spermatogenesis in many organisms. In the rainbow trout, protamine mRNA is stored for 15–30 days before it is translated (Iatrou and Dixon 1977). In Drosophila, where there is no postmeiotic transcription (Olivieri and Olivieri 1965), and in the mouse, where transcription ceases at about stage 9 (Monesi 1964; Kierszenbaum and Tres 1975), many proteins required for spermatozoon assembly are not synthesized until days after transcription ceases (O’Brien and Bellve 1980; Kuhn et al. 1988). Synthesis of these proteins is therefore dependent on mRNAs made either during meiosis or in early spermiogenesis. However, not all mRNAs are subject to translational regulation. Translation of other mRNAs occurs in early round spermatids, implying that a selective mechanism exists that ‘earmarks’ specific mRNAs for storage. In addition to mRNA storage, there may also be a defined temporal program for the translational recruitment of specific mRNAs encoding different components of the various spermatozoon structures. The mechanisms of selective mRNA storage and activation are poorly understood. The abundance of the mP1 mRNA in spermatids and the precision of its translational control make it an excellent system in which to begin investigating such mechanisms.

We have previously described the cloning of the mP1
gene and studied its transcriptional regulation in transgenic mice [Peschon et al. 1987]. Recently, we have shown that as little as 0.46 kb of mP1 5' sequence is sufficient to direct the expression of an oligonucleotide-tagged version of mP1 to haploid spermatids [Peschon et al. 1989]. In this study we have extended the analysis of the transcriptional control and have also investigated the cis-acting sequence elements required for the proper translational regulation of mP1. Our results demonstrate that the mP1 3'-untranslated region (UTR) contains all of the elements required for proper translational control and that they function even when transferred to a heterologous mRNA. Unexpectedly, we also discovered that the time during spermiogenesis at which different transgenic mRNAs are translated affects the subcellular localization of the product of the transgene within the developing spermatid.

**Results**

**Tissue- and stage-specific transcriptional control**

To study the transcriptional and translational regulation of the mP1 gene, two fusion genes were constructed between mP1 and the hGH gene [Fig. 1]. Construct mP1-hGH-mP1 3' contains 4.1 kb of the mP1 gene extending 5' to the transcription start site, 91 bp of mP1 5'-untranslated sequence, the hGH structural gene, and 156 bp of mP1 3'-untranslated sequence. Construct mP1-hGH-hGH 3' is identical to mP1-hGH-mP1 3', except that the mP1 3' UTR was replaced by the natural 112-bp hGH 3' UTR. Transgenic mice were generated with each of the gene fusions by microinjection into the pronuclei of fertilized mouse eggs [Brinster et al. 1985].

Northern blot analysis of RNA isolated from the testes and six somatic tissues of animals carrying the two transgenes shows that expression of the transgenes was restricted to the testis [Fig. 2]. The RNA blot was hybridized with a radiolabeled mP1 5' UTR probe that is complementary to the endogenous mP1 mRNA and to both transgene mRNAs. Figure 2 shows that the abundance of the two transgenic mRNAs and the endogenous mP1 transcript were similar.

The first round of spermatogenesis in the mouse is synchronous, starts at birth, and is completed in ~34 days [Nebel et al. 1961]. The time of appearance of an mRNA after birth can be used to deduce the cell type in which the gene is first expressed [Bellve 1979]. Northern blot analysis shows that mRNA from both transgenes, as well as the endogenous mP1 gene, was not detectable at day 19 after birth [Fig. 3, lanes a,f,k], a time at which cells at the leading edge of the first round of spermatogenesis are in the pachytene spermatocyte stage. However, at day 25 [lanes b,g,l], a time when cells in the first round are well into the haploid round spermatid stage, mRNA from both transgenes, as well as the endogenous gene, was detectable. The mRNAs continued to accumulate at day 28 and eventually reached a plateau in 32-day-old and adult mice. These results suggest that both transgenes are transcribed exclusively in round spermatids and that mP1 5' sequences extending from −4.1 kb to +91 bp are sufficient for appropriate mP1 transcriptional control.

**Figure 1.** Transgene constructs. Nontranscribed and intron DNA are represented as thin lines, protamine exons (non-coding) are represented as solid boxes, and growth hormone exons are represented as open boxes. The positions of the translational start and stop codons are designated with arrows above the genes. The break in the line that corresponds to mP1 sequence denotes that portion of the transgene that is not drawn to scale. Transgenes mP1-hGH-hGH 3' and mP1-hGH-mP1 3' both contain −4.1 kb of mP1 5'-untranslated sequence extending 5' from the transcriptional start site to a HindIII site at −4.1 kb. Both genes have chimeric 5' UTRs of 159 bp [91 bp of mP1 5' UTR, 7 bp of linker DNA, and 61 bp of hGH 5' UTR] and the complete hGH-coding sequence and introns [DeNoto et al. 1981]. mP1-hGH-hGH 3' contains the complete wild-type hGH 3' UTR [105 bp], and transgene mP1-hGH-mP1 3' contains a chimeric 3' UTR, which has the first 7 bp of the hGH 3' UTR and 156 bp of the mP1 3' UTR.

**Figure 2.** Tissue specificity Northern blot analysis. Total RNA (15 µg) extracted from various tissues was separated on 1.5% agarose–formaldehyde gel, transferred to nitrocellulose paper, and hybridized with a 32P-labeled 175-bp NcoI–NcoI fragment that contains 91 bp of mP1 5' UTR, which both transgenes and the endogenous mP1 gene contain. Arrows designate the positions of the transgenic mRNAs and the endogenous mP1 mRNA. Letters above each lane represent the tissue from which the RNA was extracted: [T] testis; [B] brain; [L] liver; [K] kidney; [S] spleen; [M] skeletal muscle. (Lane h) RNA isolated from line 65-4 (mP1-hGH-mP1 3'); (lane d) RNA isolated from line 314-4 (mP1-hGH-hGH 3'); (lanes a, e-i) RNA isolated from line 65-9 (mP1-hGH-mP1 3'); (lanes c, j-n) RNA isolated from line 315-6 (mP1-hGH-hGH 3').
Temporal translational regulation

To determine the stage during prepubertal testis development at which the two transgenic mRNAs were translated, immunocytochemistry was performed on testis sections from prepubertal animals. In mice containing the mPI-hGH-mP1 3’ construct [Fig. 4e], hGH was first detected in elongating spermatids within a small percentage of seminiferous tubules of 28-day-old animals. This corresponds to the same time that the endogenous mP1 protein is also first detected (Balhorn et al. 1984; Kleene et al. 1984). By day 32 [Fig. 4h], many of the tubules within the testis contained hGH immunopositive spermatids. Immunocytochemical analysis of adult testis sections showed that hGH was first detected in elongating spermatids at about stage 12 [Fig. 4k]. In contrast, in transgenic animals that contained the mPI-hGH-hGH 3’ gene, hGH protein was first detected in round spermatids in a few tubules at day 25 after birth [Fig. 4c]. By day 28 [Fig. 4f], more tubules contained immunopositive spermatids, and by day 32 [Fig. 4i], many of the tubules were immunopositive. Immunocytochemical analysis of adult testes first revealed hGH in round spermatids at about stage 3–4 [Fig. 4l]. This stage correlates with the most mature spermatids in day-22 prepubertal animals and is the earliest time at which mPI mRNA is detected (Kleene et al. 1983). Thus, despite the fact that the mPI-hGH-mP1 3’ and mPI-hGH-hGH 3’ genes are under the same transcriptional control, translation of the two messages occurs at different stages of spermiogenesis. These results show that as little as 91 nucleotides of mP1 5’-untranslated sequence and 156 nucleotides of mP1 3’-untranslated sequence are sufficient to confer mPI-like translational control on a heterologous mRNA. In addition, these results demonstrate that the mP1 3’ UTR is necessary for proper translational control. We also tested a mPI-hGH-mP1 3’ construct that contained only 7 bp of mP1 5’ UTR instead of 91 bp. Immunocytochemical analysis of testes from adult transgenic mice carrying this construct revealed that hGH was only synthesized in elongating spermatids, thus demonstrating that the mP1 3’ UTR is sufficient for mP1 translational control [data not shown].

Subcellular localization of hGH protein

Synthesis of hGH in early round spermatids versus late elongating spermatids resulted in different subcellular localization of the protein. In animals that contained the mPI-hGH-mP1 3’ gene, hGH protein accumulated intracellularly in elongating spermatids at about day 28 and continued to increase there through day 32 [Fig. 4]. In animals that contained the mPI-hGH-hGH 3’ gene, hGH protein was localized in the acrosome of spermatids. At day 25 [Fig. 4c], and in adult animals [Fig. 4l], this can be seen as punctate staining adjacent to the nucleus; at day 28 [Fig. 4f], it is seen as a half-moon; and at day 32 [Fig. 4i], it is seen as a crescent-shape stain corresponding to the shape of a mature acrosome. Thus, the time and, therefore, the stage at which hGH is synthesized during spermiogenesis determines its localization within the cell.

Immunocytochemistry of epididymal tissue from the mPI-hGH-mP1 3’ animals also revealed the presence of hGH protein in the cytoplasm of the epithelial cells that line the efferent ducts [data not shown]. These cells are known to resorb fluids and cellular debris generated from the residual bodies and cytoplasmic droplets released by the spermatids (Mason and Shaver 1952), and it is likely that the hGH staining seen in them reflects their endocytic role. hGH was not seen in the epithelial cells of the efferent ducts in the mPI-hGH-hGH 3’ animals. This observation is consistent with immunocytochemical data that revealed that hGH was stable in the acrosome at least through the caput section of the epididymis (Braun et al. 1989).

To confirm the differential translational regulation of the two transgenes, total SDS-soluble protein extracts were prepared from the testes of transgenic animals aged 19, 25, 28, and 32 days old and analyzed by Western blot analysis with hGH-specific antibodies [Fig. 5]. hGH was first detected at day 28 [lane e] in those animals that contained the mPI-hGH-hGH 3’ construct and at day 32 [lane h] in those animals that contained the mPI-hGH-mP1 3’ construct. The time during prepubertal development that hGH was first detected was delayed for both constructs due to the lower sensitivity of this method compared to immunocytochemistry. Figure 5 shows that the amount of protein that eventually accumulated [lanes g,h] was approximately equivalent. These results confirm the immunocytochemical analysis showing that the two transgenes are under different translational control.

hGH is normally a secreted protein and is synthesized with an amino-terminal signal peptide that is cleaved in the rough endoplasmic reticulum (RER) to generate the mature secreted protein. Figure 5 shows that a protein of
Prepubertal immunocytochemical analysis. Testes from prepubertal animals, aged 25 (a–c), 28 (d–f), or 32 days old (g–i), and adult animals (j–l) were fixed, sectioned, and treated with a rabbit anti-hGH primary antibody and a peroxidase-conjugated goat anti-rabbit secondary antibody, as described in Methods. (a,d,g,j) Sections were from animals of the 65-4 line (mP1-hGH-mP1 3') and were only incubated in the presence of the secondary antibody; (b,e,h,k) sections were also from the 65-4 line; (c,f,i,l) sections were from animals of the 314-4 line (mP1-hGH-hGH 3'),. The arrow in i shows the punctate hGH staining seen in the acrosome of stage 3–4 tubules. Additional negative controls included both the primary and secondary antibody on testes from nontransgenic animals.
Differential stability of transgene mRNAs

The immunocytochemical analyses described above revealed that if hGH was synthesized in elongating spermatids, it was intracellular but not acrosomal, whereas if it was synthesized in round spermatids, it was localized in the acrosome. In light of this, we wondered why mPI-hGH-mPI 3' mRNA was not translated in both round and in elongating spermatids, resulting in both acrosomal and nonacrosomal localization of hGH. One explanation could be that mPI-hGH-hGH 3' mRNA was only present in round spermatids. To determine whether the mPI-hGH-mPI 3' and mPI-hGH-hGH 3' mRNAs were present in both early- and late-stage spermatids, we performed in situ hybridization experiments with mPI and hGH-specific probes.

Hybridization of testis serial sections from a mPI-hGH-mPI 3' animal with 35S-labeled, mPI-specific (Fig. 6a,b) and hGH-specific (Fig. 6c,d) antisense RNA probes shows that the transgenic mRNA was present in the same staged tubules as was the endogenous mPI mRNA, suggesting that the stabilities of the endogenous and transgenic mRNA are similar. However, comparison of serial sections from a mPI-hGH-hGH 3' animal hybridized with the mPI-specific (Fig. 6e,f) and hGH-specific (Fig. 6g,h) antisense RNA probes shows that the overall pattern of hybridization with the two probes was different. Whereas those tubules that contained early-stage spermatids show the same hybridization signal with both probes, tubules with later stage spermatids only hybridized with the mPI riboprobe, suggesting that the endogenous mPI mRNA and the mPI-hGH-hGH 3' transgenic mRNA have different stabilities.

Discussion

Translational control

We have shown that chimeric transgenesc containing the promoter and transcriptional regulatory elements of the mPI gene, the hGH structural gene, and the mPI 3' UTR are under the same transcriptional and translational control as the endogenous mPI gene. These data and others (Peschon et al. 1989) demonstrate that all of the cis-acting sequence elements required for spermatid-specific transcription of mPI lie between -465 bp and the start of transcription. The mPI 3' UTR is critical for both delaying translation until the late stages of spermiogenesis and for mRNA stability. The importance of the mPI 3' UTR in translational control was predicted based on sequence conservation among the 3' UTRs of the mouse, human, and bovine protamine 1 genes (Krawetz et al. 1987, Lee et al. 1987a,b, Peschon et al. 1987). All three genes share at least 75% similarity in their 3' UTRs, with the most conserved regions mapping near the poly(A) site. In fact, the mouse and human genes are more similar in their 3' UTRs (89%) than they are in their protein-coding sequence (69%). Sequences in 3' UTRs have also been shown to be involved in iron-dependent regulation of transferrin receptor mRNA (Casey et al. 1988), plastid mRNA regulation (Gruissem et al. 1988), and t-PA translational control in mouse oocytes (Strickland et al. 1988).

In amphibians and in marine invertebrates, translational regulation is a common regulatory mechanism during oogenesis and early embryogenesis (Davidson 1986, Richter 1987; Rosenthal and Wilt 1987). In addition to there being a general overall inhibition of translation in unfertilized versus fertilized eggs, there are examples of both selective inhibition (Ford et al. 1977) and selective activation (Rosenthal et al. 1980) of translation. In mouse oogenesis, the tissue plasminogen activator gene (Huarte et al. 1987, Strickland et al. 1988) is

Figure 5. Prepubertal Western blot analysis. Total SDS-soluble protein extracts from testes were prepared from prepubertal animals of the 314-4 line (lanes a,c,e,g) and 65-4 line (lanes b,d,f,h). Equivalent amounts of protein from each prepubertal time point (60 µl, day 19, 40 µl, day 25, 30 µl, day 28, and 20 µl, day 32) were precipitated with 20% TCA, separated by SDS-PAGE, electroblotted, and treated with rabbit anti-hGH antibody and a peroxidase-conjugated goat anti-rabbit IgG secondary antibody, as described in Methods. (Lane j) Media harvested from baby hamster kidney (BHK) cells that had been transfected with a metallothionin-hGH gene (lane j) and corresponds to the mature form of hGH generated by cleavage of its amino-terminal signal peptide (DeNoto et al. 1981). Therefore, regardless of the subcellular localization of hGH, its signal peptide was properly cleaved. A second protein of ~22 kD was also observed and probably corresponds to a variant of the mature protein generated by alternative splicing that eliminates the coding sequence for the first 15 amino acids of the third exon of the hGH gene (Lewis et al. 1978, DeNoto et al. 1981). The ratio of the two proteins is different than that observed in BHK cells transfected with a metallothionin-hGH gene (cf. lanes h and j) and in the pituitary (Lewis et al. 1978, DeNoto et al. 1981; Cooke et al. 1988). We have not determined whether the altered ratio of the two proteins in the testis is due to differences in the frequency of the alternative splicing event or to differences in the stability of the two proteins.

Differential stability of transgene mRNAs

The immunocytochemical analyses described above revealed that if hGH was synthesized in elongating spermatids, it was intracellular but not acrosomal, whereas if it was synthesized in round spermatids, it was localized in the acrosome. In light of this, we wondered why mPI-hGH-hGH 3' mRNA was not translated in both round and in elongating spermatids, resulting in both acrosomal and nonacrosomal localization of hGH. One explanation could be that mPI-hGH-hGH 3' mRNA was only present in round spermatids. To determine whether...
Figure 6. In situ hybridization. Serial sections of testes from either the 65-4 line [mP1-hGH-mP1 3' (a-d)] or the 314-4 line [mP1-hGH-hGH 3'; (e-h)] were hybridized with a 35S-labeled mP1-specific antisense probe (a,b,e,f) or an hGH-specific antisense probe (c,d,g,h), as described in Methods. Negative controls included hybridization with a mP1-specific sense probe or an hGH-specific sense probe (data not shown). (E) Early-stage spermatids, (L) late-stage spermatids. (Left) Bright-field microscopy, (right) dark-field microscopy.
transcribed in primary oocytes but is not translated until their maturation. During recent years, the importance of translational control during spermatogenesis has also become apparent. Specific genes that are regulated post-transcriptionally include the mouse PGK-2 gene (Gold et al. 1983), the trout protamine genes (Iatrou and Dixon 1978), the rat TP1 gene (Heidaran et al. 1988), and the *Drosophila mst*(3)gl-9 gene (Kuhn et al. 1988). In oogenesis and early embryogenesis, the two mechanisms that are thought to play a role in inhibiting translation are limitations of specific translation factors (Richter and Smith 1981; Collin and Hille 1986) and masking of mRNAs by proteins (Rosenthal et al. 1980). In one special case—the histone mRNAs of sea urchins—translation is inhibited by sequestering the mRNA in the female pronucleus until nuclear membrane breakdown during the first mitotic division (Goustin 1981; Wells et al. 1981; Angerer et al. 1984). The association of mP1 mRNA with mRNPs in early spermatids (Kleene et al. 1984), where it is not translated, and with polysomes in elongating spermatids, where it is translated, suggests that masking of mP1 mRNA inhibits its translation in round spermatids. Because the mP1 mRNA is present at 10,000 copies per cell (K. Palmiter and D. Wolgemuth, unpubl.), it is likely that factors that bind to mP1 mRNA are also abundant. We are currently attempting to purify such factors.

The mRNA for mP1 is initially found in round spermatids as a 560-nucleotide transcript and later in elongating spermatids as a heterogeneous size class of molecules down to as small as 400 nucleotides (Fig. 3, lanes 1–o; Kleene et al. 1984). This decrease in mRNA size is due to a shortening of the poly(A) tail and occurs at the same time as translation of mP1 commences. It has yet to be resolved if deadenylation is a prerequisite for, or a consequence of, translation. Deadenylation also seems to be associated with translation of Xenopus oocyte histone mRNAs (Ruderman et al. 1979). In other systems it is usually polyadenylation that is correlated with translation (Wilt 1977; Rosenthal et al. 1983; Rosenthal and Ruderman 1987).

**Message stability**

Although we detected mP1-hGH-mP1 3' mRNA by in situ hybridization in both early- and late-stage spermatids, mP1-hGH-hGH 3' mRNA was clearly more abundant in early spermatids than in late-stage spermatids. Because both genes are first transcribed at about stage 3–4 of spermiogenesis and overall transcription ceases at about stage 9 (Monesi 1964; Kierszenbaum and Tres 1975), transcriptional differences are not likely to be involved. However, the half-lives of the two mRNAs could be different. Such a difference could be due to two, not mutually exclusive, possibilities. First, sequences present in hGH 3' UTR could be directly responsible for a change in half-life. In both prokaryotes and eukaryotes, 3' UTRs have been shown to affect mRNA stability (Shaw and Kamen 1986; Brawerman 1987). Second, premature translation of the mP1-hGH-hGH 3' mRNA relative to the mP1-hGH-mP1 3' mRNA could indirectly lead to a shorter half-life. Linkage between mRNA stability and ongoing protein synthesis is a common occurrence and may reflect the association of RNases with the translational apparatus (Graves et al. 1987; Yen et al. 1988).

**Protein sorting**

Most cells have either a constitutive secretory pathway, in which proteins are rapidly exported from the cell following their synthesis, or a regulated secretory pathway, in which proteins are stored in vesicles until their release following the appropriate signal (Tartakoff and Vassalli 1978; Burgess and Kelly 1987). Proteins in both pathways are synthesized on the rough endoplasmic reticulum (RER), pass through the cis-cisterna of the Golgi complex together, and are sorted in the trans-cisterna (Palade 1975; Orci et al. 1987; Tooze et al. 1987). hGH is normally expressed in the pituitary and is a member of the regulated class of secretory proteins (Burgess and Kelly 1987). The acrosome is a regulated secretory vesicle in that its contents are stored until their release following induction of the acrosome reaction during fertilization (Wassarman 1987). Translation of mP1-hGH-hGH 3' mRNA in early spermatids resulted in localization of hGH in the acrosomal compartment suggesting that the hGH protein is recognized as an acrosomal protein when synthesized in early spermatids. In some regulated secretory systems, for example, in lysosomal vesicles, proteins are recognized and sorted as a result of covalent modification (von Figura and Weber 1978; Darquar 1985). The sorting signals for hGH are currently unknown but do not involve protein modification (Guan and Rose 1984; Moore and Kelly 1986). The localization of hGH in the acrosome suggests that the protein-sorting machinery used in spermatids to construct the acrosome is functionally equivalent to that used in pituitary cells.

**Acrosome development**

Acrosome development begins early in spermiogenesis and consists of four basic steps: the Golgi phase, stages 1–3; the cap phase, stages 4–7; the acrosome phase, stages 8–11; and the maturation phase, stages 12–16 (Leblond and Clermont 1952; Oakberg 1956; Dooh and Bennett 1973). In mP1-hGH-hGH 3' animals, hGH first appeared in the developing acrosome of round spermatids at about stages 3–4, whereas in mP1-hGH-mP1 3' animals, hGH protein first appeared in elongating spermatids at about stage 12 and was not acrosomal. Because the signal peptide was removed, we presume that the mRNA was translated on RER and is sequestered in some, as yet undetermined, organelle. Nonetheless, the failure of hGH, synthesized during stages 12–15, to enter the acrosome suggests that the system targeting hGH to the acrosome in round spermatids is no longer functional in elongating spermatids, implying that the sorting of normal acrosomal proteins is completed before or soon after the beginning of the maturation phase. Electron microscopy has shown that the Golgi apparatus
moves caudally during the later stages of spermiogenesis and that an extensive network of various forms of RER extends throughout the cytoplasm [Susi et al. 1971; Clermont and Rambourg 1978]. The intracellular localization of hGH thus serves as a molecular marker for the cessation of acrosomal protein synthesis. Of interest is the signal that marks the completion of acrosomal assembly and the mechanism that dissociates the Golgi from the acrosome.

In cells that have both a regulated and constitutive secretory pathway, blockage of the regulated pathway results in rerouting of the proteins destined for the regulated pathway into the constitutive pathway, resulting in rapid secretion [Neufeld et al. 1975; Moore et al. 1983]. The fact that hGH does not appear to be secreted in elongating spermatids suggests that they do not have a constitutive secretory pathway. Alternatively, either another regulated pathway may be established late in spermiogenesis or remnants of the sorting signals for the acrosomal pathway, although disconnected from the acrosome, may still be present.

Methods

DNA and RNA analysis

DNA manipulations were performed using standard procedures. RNA used for Northern blot analysis was extracted by homogenization in guanidinium isothiocyanate, followed by precipitation with lithium chloride, as described by Cathala et al. [1983]. RNA samples were electrophoresed in 1.5% agarose-formaldehyde gels, transferred to nitrocellulose paper, and hybridized with radioactive DNA probes prepared by nick translation or random oligonucleotide-primed synthesis [Feinberg and Vogelstein 1983]. Single-stranded 32P-containing sense and antisense riboprobes were made in vitro with linear templates and either SP6 or T7 RNA polymerase [Melton et al. 1984]. The mPl template was made by cloning a 250-bp restriction fragment containing a portion of mPl exon 1, the intron, and a portion of mPl exon 2 into pGEM-2. The hGH template was constructed by cloning a 170-bp restriction fragment containing hGH exon 5 into pGEM-2.

Microinjection

The pronuclei of fertilized eggs derived from [C57BL/6 x SJL]F1 females mated with identical hybrid males were microinjected with -2 nl of DNA solution, as described by Brinster et al. [1985]. Eggs that survived microinjection were implanted into the oviducts of pseudopregnant Swiss foster females. Two lines of transgenic mice were generated and analyzed for each transgene described in Figure 1. Lines 314-4 and 315-6 carry the mPl-hGH-hGH 3' gene, and lines 65-4 and 65-9 carry the mPl-hGH-mPl 3' gene.

Protein extracts and Western blotting

Testes were decapsulated and homogenized in 0.8 ml of buffer containing 10 mM HEPES, 50 mM NaCl, 10 mM EDTA, and 0.2 mM PMSF. The proteins were solubilized in 10% SDS on ice for 15 min, sonicated until they were no longer viscous, and centrifuged in a microfuge for 5 min. Equivalent amounts of supernatants were precipitated with 20% TCA, resuspended in Laemmli buffer [Laemmli 1970], boiled, and electrophoresed in SDS-polyacrylamide gels. The proteins were electroblotted onto nitrocellulose [Towbin et al. 1979] for 3 hr at 200 mA at room temperature. Filters were washed at room temperature for 1 hr in 5% nonfat dry milk and PBS incubated with a 1/500 dilution of primary antibody overnight at 4°C, washed in 5% nonfat dry milk and PBS, incubated with a goat anti-rabbit IgG antibody conjugated to horseradish peroxidase [HRP] (Bio-Rad) for 3 hr at room temperature, and washed again in 5% nonfat dry milk and PBS. The HRP was detected with a diaminobenzidine substrate [Bio-Rad], using the manufacturer's instructions. Equivalent amounts of protein from all of the prepubertal time points were determined in pilot experiments by comparing the total amount of protein on Coomassie-stained SDS-PAGE gels.

Immunocytochemistry

Testes were fixed in Carnoy’s reagent, embedded in paraffin, and cut into 5-μm sections. Sections were deparaffinized using standard procedures. Indirect immunocytochemistry was performed using the reagents and suggested protocols of Vectastain ABC and Zymed SABC kits. The primary antibody, rabbit anti-hGH, was obtained from the National Hormone and Pituitary Program (Baltimore, Maryland), and used at a dilution of 1/500. Aminoethyl carbazole [Zymed Laboratories, Inc.] was used as the substrate chromogen. Sections were counterstained with hematoxylin. Additional negative controls included the use of both the primary and secondary antibodies on testes isolated from nontransgenic animals.

In situ hybridizations

Testes were dissected from adult animals, fixed in 4% paraformaldehyde, embedded in paraffin, and cut into 5-μm sections. The paraffin was extracted with xylene and the tissue was rehydrated using standard conditions. The sections were washed twice for 5 min in 0.1 M Na2HPO4/NaH2PO4 [pH 7.5], incubated with proteinase K at 1 μg/ml for 10 min at 37°C, washed twice for 2 min in H2O, washed once in 0.1 M triethanolamine [TEA] and 150 mM NaCl [pH 8.0] for 2 min, treated with 0.25% acetic anhydride in TEA/NaCl for 10 min, washed twice for 5 min in 2 x SSC, and hybridized for 6–12 hr at 45°C with radioactive probes. Hybridization solution contained 50% formamide, 0.6 M NaCl, 10 mM Tris [pH 7.5], 1 mM EDTA, 1 × Denhardt’s solution, 10% dextran sulfate, 10 mM DTT, and yeast RNA at 50 μg/ml. Probes were used at a concentration of -0.2 μg/ml, with an approximate sp. act. of 5 × 106 cpm/μg. Slides were washed twice in 4 x SSC and 5 mM DTT for 15 min, incubated for 30 min at 37°C in 10 mM Tris, 1 mM EDTA, 500 mM NaCl, and 25 μg/ml RNAse A, washed in 2 x SSC and 2 mM DTT for 60 min at 22°C, washed in 0.1 x SSC and 2 mM DTT for 30 min at 45°C, and washed in 0.1 x SSC for 5 min at 22°C. Slides were dehydrated with ethanol using standard procedures, dried, dipped in Kodak NTB emulation, stored at 4°C, developed after 1–5 days, and counterstained with hematoxylin.

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