Oocyte and somatic tyrosine tRNA genes in *Xenopus laevis*

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Over a period of many months, *Xenopus* oocytes stockpile large quantities of tRNA for use during the first few hours of embryogenesis. To test the idea that these tRNAs are transcribed from one set of genes and that another set is used by somatic cells, we used synthetic oligonucleotides to analyze the sequence and steady-state levels of unspliced tyrosine tRNA precursors in *Xenopus laevis* oocytes, embryos, and cultured kidney cells. These analyses identify four kinds of tyrosine tRNA genes, two oocyte-type and two somatic-type, whose unspliced transcripts are distinguishable from one another by their different 5' leader and intervening sequences. The oocyte-type tyrosine tRNA precursors are present in oocytes, very abundant in gastrula embryos, but absent from postembryonic somatic cells. The somatic-type precursors are undetectable in oocytes but are found in gastrula and later stage embryos and in somatic cells. The major switch from oocyte-type to somatic-type transcripts occurs early during embryogenesis, between the midblastula transition and the onset of neurulation, but some oocyte-type precursors are also detectable in tadpoles.

[Key Words: tRNA gene; tRNA precursors; intervening sequences; developmental regulation; embryogenesis; *Xenopus laevis*]

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*Xenopus laevis* contains an unusually large number of tRNA genes (~8000 per haploid genome; Clarkson et al. 1973), many of which are located in extensive multigene families (Müller and Clarkson 1980; Rosenthal and Doering 1983). One such family is composed of 3.18-kb DNA fragments, each of which contains eight genes for seven different tRNA species, that are repeated tandemly ~150 times at a single chromosomal locus (Fostel et al. 1984; Müller et al. 1987).

A potential explanation for these highly repeated genes is that they are needed to satisfy the unusual synthetic demands of oogenesis, during which single oocytes accumulate ~90 ng of tRNA (Dixon and Ford 1982). The tRNA requirements of somatic cells could be met by a smaller number of genes located in a different sequence context. A precedent for this type of developmental regulation is provided by the *Xenopus* oocyte-type and somatic-type 5S RNA genes (for review, see Wolfe and Brown 1988).

One of the eight genes on the 3.18-kb repeat contains an intervening sequence, it codes for tyrosine tRNA and is referred to as TyrC. A second kind of tRNA*^{Tyr}* gene has been isolated from *X. laevis* that is present in 1–3 copies per haploid genome; this gene, referred to as TyrD, differs from TyrC by a single purine transition in the coding region, by extensive differences in the intervening sequence and flanking regions, and by its ~6 times higher transcriptional activity in somatic S-100 extracts (Gouilloud and Clarkson 1986). Thus, it possesses many of the features that might be expected for a somatic-type gene.

In the work presented here we used synthetic oligonucleotides to analyze the sequence and steady-state levels of unspliced tyrosine tRNA precursors during early development. The results indicate that the TyrC and TyrD genes are indeed oocyte type and somatic type, respectively, but that there are additional genes of each class. We describe some sequence features of these genes and compare the developmental changes of tyrosine tRNA with the oocyte-somatic switch of 5S RNA.

**Results**

RNA sequences and oligonucleotide probes that distinguish the TyrC and TyrD transcripts

Figure 1 shows a comparison of the transcription products of the TyrC and TyrD genes in S-100 extracts of cultured *X. laevis* kidney cells (Gouilloud and Clarkson 1986). Transcription of TyrC starts with a pppAp located 5 nucleotides before the mature coding sequence, and it terminates at many of the T residues within a downstream T$_3$GT$_4$ stretch to yield primary transcripts of 97–102 nucleotides (Pre-tRNA 1, Fig. 1). These precursors are converted to a processing intermediate of 89 nucleotides that contains mature 5' and 3' ends (Pre-tRNA 2); the 13-nucleotide intervening sequence then is removed by splicing and ligation reactions to yield the mature length tRNA of 76 nucleotides. TyrD produces primary transcripts of ~101 nucleotides that contain a 5' leader of 7 nucleotides, an intervening sequence of 12 rather than 13 nucleotides, an A
rather than a G at position 57 of the coding region, and 3’ trailer sequences that end within the first 2–4 residues of a T15 stretch. A second initiation site, an A at -3, gives rise to a minor fraction of TyrD primary transcripts of ~97 nucleotides. These precursors are converted to a correspondingly shorter processing intermediate of 88 nucleotides and then to the mature tRNA.

Figure 1 also shows some of the oligonucleotide probes that were used to examine the kinds of tRNA\textsuperscript{tyr} gene transcripts made in vivo. Oligonucleotides C1 and D1 are specific for the intervening sequences of TyrC and TyrD, respectively. Under appropriate conditions, they should anneal only to the complementary unspliced precursors. The tyrosine coding sequence probe M1 should detect the same precursors, and any additional ones, together with the mature tRNA\textsuperscript{tyr}. These three probes range in length from 25–28 nucleotides and were designed to yield RNA–DNA hybrids of comparable stabilities, assuming that a G–C base pair is twice as stable as an A–T base pair [Suggs et al. 1981].

**TyrC is an oocyte-type gene and TyrD is a somatic-type gene**

When incubated with Northern blots of total ovary RNA, oligonucleotide C1 anneals to a heterogeneous set of RNAs of ~100 nucleotides and to a shorter, apparently unique RNA species of ~90 nucleotides (Fig. 2, lane c). Thus, these precursors correspond in both sequence and length to the TyrC primary transcripts and to their processing intermediates. Oligonucleotide M1 hybridizes to these same precursors and also yields the expected strong signal in the mature tRNA region [Fig. 2, lane b]; however, none of these bands is detected by the TyrC intron probe in RNA from cultured kidney cells [Fig. 2, lane f]. These results suggest that the highly repeated TyrC genes are active in oogenesis but that their expression is repressed in somatic cells.

Conversely, the TyrD intron probe hybridizes weakly to a single precursor band in the kidney cell RNA and fails to give a signal with ovarian RNA [cf. lane g with lane d, Fig. 2]. From the mobilities of internal labeled markers in the Northern blots (data not shown), this somatic precursor also has a length of ~100 nucleotides. Oligonucleotide M1 gives a much stronger signal with RNAs of this length, together with an intense band that corresponds to mature tRNA\textsuperscript{tyr} [Fig. 2, lane e]. This probe also hybridizes to a group of kidney cell RNAs of ~90 nucleotides; these molecules presumably represent tRNA\textsuperscript{tyr} precursors that either lack an intervening sequence or contain one that differs from the one within TyrD. Thus, these results indicate that TyrD is a minor somatic-type gene and that most of the kidney cell tRNA\textsuperscript{tyr} precursors are derived from a different set of somatic-type tyrosine tRNA genes. It is possible that TyrD also is expressed during oogenesis but that, because of its much lower copy number than TyrC, its transcripts are undetectable.

**TyrC is one of two tRNA\textsuperscript{tyr} genes expressed in the ovary**

A direct way to characterize these various transcripts is to anneal the oligonucleotides with the RNA samples in solution, to extend the primers with reverse transcriptase and then to sequence the extended products. This is not straightforward technically because the substrates are present in low amounts, and they contain a high degree of structure and some modified bases, both of which cause frequent premature termination. The cDNA yield can be maximized, however, by use of a primer such as M1 that is complementary to sequences near the 5’ end of the mature tRNA.

Figure 3 shows the reverse transcriptase products obtained from ovary tRNA precursors primed with oligonucleotide M1, together with sequence analysis of some of the longer cDNAs. Band D contains the 25-nucleotide primer and an additional 11 nucleotides that are complementary to the first 11 nucleotides of mature tyrosine

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**Figure 1.** Transcripts and oligonucleotides specific for the TyrC and TyrD genes. The unspliced precursors and oligonucleotide probes are aligned with the 5’ end of the mature tRNA\textsuperscript{tyr} produced from each gene. Nucleotides are numbered according to the convention for tRNAs and their genes [Sprinzl et al. 1987]. Transcripts are shown in their unmodified form and the anticodon is underlined. The 5’ leaders, intervening sequences, and the purines at position 57 are shown in boldface type.
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Figure 2. Northern blot analysis of X. laevis ovary and kidney cell RNA. Samples of total RNA from ovary tissue and from cultured kidney cells were fractionated over two wide slots of a thick sequencing gel and were transferred electrophoretically to a nylon membrane. The region of the filter that contained RNA sample was cut into strips, and pairs of strips, one from each sample, were hybridized with the 5'-labeled oligonucleotide probes (lanes b–d and e–g). Exposures were for 1 day at −70°C on preflashed film with an intensifying screen. The ethidium fluorescence of RNA samples electrophoresed under identical conditions is shown (lanes a and h). Some uncharacterized components, possibly polysaccharides, that migrate between ovary 5S and 4S RNA cause the anomalously fast mobilities of the tRNA precursors.

tRNA; presumably this cDNA is not derived from the mature tRNA\(^{\text{Tyr}}\), which was removed by prior gel electrophoresis, but rather from tRNA\(^{\text{Tyr}}\) precursors that have undergone 5'-end processing.

The longer cDNAs are extension products of tRNA\(^{\text{Tyr}}\) precursors with 5' leaders. Band B contains an extra 5 nucleotides (5'-CCGGGT-3') that are precisely complementary to the 5' leader sequence of the longest TyrC transcripts made in vitro. Thus band B is identified as the cDNA product of TyrC primary transcripts. Band C is a truncated version of B that contains 3 of these 5 nucleotides (5'-CCG-3'). This cDNA could arise through partial degradation of the TyrC primary transcripts or their incomplete copying; it seems unlikely that it represents a full-length copy of a primary transcript because this would imply initiation with a pppCp. Thus, these results provide direct evidence for the TyrC gene transcripts in the ovary.

Sequence analysis of band A (Fig. 3) has revealed the presence of another kind of tyrosine tRNA precursor in this tissue. This cDNA contains the 36 nucleotides characteristic of tRNA\(^{\text{Tyr}}\) plus an additional 7 nucleotides that are not complementary to the 5' leaders of either TyrC or TyrD primary transcripts. Apart from uncertainty about the last residue, this sequence appears to be unique (5'-CAGGAGN-3'). Thus, band A is derived from transcripts of a second type of tRNA\(^{\text{Tyr}}\) gene that is expressed in the ovary; moreover, the failure to detect longer cDNAs suggests that band A is the cDNA product of its primary transcripts.

The two tRNA\(^{\text{Tyr}}\) genes expressed in the ovary contain different intervening sequences

TyrC precursors contain a 13-nucleotide intervening sequence and a G at position 57 of the mature tRNA [Fig. 1]. To determine whether these features are found also in the other ovarian tyrosine tRNA precursors, an oligonucleotide complementary to the last 14 nucleotides of mature tRNA\(^{\text{Tyr}}\) (oligonucleotide M2, Fig. 1) was used for primer extension analysis. Reactions with this primer are particularly inefficient because the reverse transcriptase has to copy almost the entire highly structured tRNA moiety of the precursors. Nevertheless, some full-length cDNAs are obtained that, when sequenced (Fig. 4), exhibit a unique C that corresponds to position 57 of mature tRNA. Thus, the absence of a strong T reaction suggests that a G occupies this position in both kinds of ovarian tRNA\(^{\text{Tyr}}\) precursors.

The remainder of the cDNA sequence is unique also, and complementary to tyrosine tRNA, except for the presence of some double reactions within the region that corresponds to the intervening sequence of TyrC (Fig. 4). This implies that the two kinds of tRNA\(^{\text{Tyr}}\) precursors contain intervening sequences of the same length (13 nucleotides) but that they differ in some positions. The intervening sequence within the second ovarian tRNA\(^{\text{Tyr}}\) precursor may be deduced by subtracting that of TyrC; because of a suboptimal C reaction, the deduced sequence is not unambiguous but it is clear that the two intervening sequences differ in at least two positions (Figs. 4 and 5).

A fourth kind of tRNA\(^{\text{Tyr}}\) gene is expressed in somatic cells

Primer extension sequencing was used similarly to analyze the nature of the unspliced tRNA\(^{\text{Tyr}}\) precursors synthesized in cultured kidney cells. Oligonucleotide M1 yields a series of cDNAs that extend beyond the 5' end of the mature tRNA; unlike their ovary counterparts, these cDNAs appear to be derived from a major set of tRNA\(^{\text{Tyr}}\) precursors that contain just one kind of 5' leader sequence (data not shown). The longest cDNA contains an extra 7 nucleotides (5'-CTTGATAN-3') that is not complementary to the 5' leaders of TyrD, TyrC, or the other ovarian precursors. The sequences characteristic of this novel 5' leader also are found in full-length extension products of the M2 primer (Fig. 4). More clearly visible in this particular gel is the unique T reaction at a position complementary to R\(_5\) of the mature tRNA. The major somatic transcripts therefore contain an A residue at this position. They also possess a 13-nucleotide intervening sequence (Fig. 4) that is different from the 12-nucleotides intron within TyrD.

Hence cultured kidney cells transcribe a fourth type of tRNA\(^{\text{Tyr}}\) gene that is distinct from TyrD and from the two kinds of genes that are expressed in the ovary. The results imply further that both kinds of genes active in the ovary, and not merely TyrC, are repressed in somatic cells. The primer extension analyses are summarized in Figure 5, which aligns the noncoding DNA strands of the two oocyte-type and two somatic-type genes.
Developmental expression of oocyte-type and somatic-type genes

To determine when the TyrC, TyrD, and major somatic-type genes are expressed during development, Northern blots of total RNA from staged oocytes (Dumont 1972) and embryos (Nieuwkoop and Faber 1967) were annealed with oligonucleotides M1, C1, and D1 (Fig. 1) and a fourth probe, oligonucleotide S1 (Fig. 5). From the primer extension sequencing results, this new probe should be specific for the intervening sequence of the major somatic-type tRNA
precursors.

Tyrosine tRNA precursors are very difficult to detect in oocytes under conditions that readily reveal the precursors in gastrula and later stage embryos (Fig. 6). With increased RNA amounts and longer exposure times, however, tRNA
precursors of ~100 and ~90 nucleotides can be detected in stage I oocytes with oligonucleotides M1 (Fig. 6) and C1 (data not shown). Therefore, these precursors include unspliced transcripts of the TyrC genes. From the pattern of hybridization with oligonucleotide M1, their steady-state levels remain roughly constant from stages I–IV and then decrease in stages V and VI. These results suggest that at least the TyrC oocyte-type genes are expressed very early during oogenesis and that changes occur in the synthesis, processing, or degradation of their transcripts as the oocyte nears maturation. In contrast, the D1 and S1 probes give no detectable signal with RNA from staged oocytes or from ovary tissue under conditions that yield a strong ~100 nucleotide band in kidney cell RNA (Fig. 2 and data not shown). This implies either that somatic-type transcripts are not synthesized during oogenesis or, if they are made, that they account for a very minor fraction of the oocyte tRNA
precursor population.

No tRNA
precursors are found between fertilization and early blastula (stage 8) with any of the four probes, even after long exposures of blots that contain large amounts of RNA (data not shown). The M1 probe detects the mature tRNA
readily, however, which remains roughly constant per embryo during these early synchronous cleavage stages (Fig. 6). The hybridization patterns then change dramatically after the midblastula transition (MBT). Oligonucleotide M1 first detects tRNA
precursors at late blastula (stage 9); their steady-state levels are very high in gastrula embryos, with a peak at stage 10, and then drop sharply in neurula and tail-bud stages. A second phase of tRNA
precursor accumulation then commences at around the time of hatching (Fig. 6).

The TyrC precursors are detected first at stage 9, they reach a maximum at stage 10, decline at the end of gastrulation, and are not found in late neurula and tail-bud stages; they then reappear in low amounts after hatching (Fig. 6). The major somatic-type tRNA
precursors first appear at stage 10, and they peak at late gastrula (stage 12); they also decline to some extent at the end of gastrulation but are still present in neurula stages; they then exhibit their greatest accumulation in tadpoles (Fig. 6). The TyrD precursors follow the same pattern but they are present in at least 10-fold lower amounts (Fig. 6; note the 5 times longer exposure of the D1 blot).

To try to confirm and extend these Northern blot analyses, primer extension sequencing was applied to RNA from selected embryonic stages. With RNA from stage-10 embryos, the M1 primer generates cDNAs that...
are identical in length and sequence to those produced from ovary RNA (data not shown). Thus, both kinds of tRNA\textsuperscript{Tyr} genes expressed in the ovary, and not merely TyrC, are also active in gastrula stage embryos. Moreover, the two kinds of oocyte-type transcripts account for the vast majority of the tRNA\textsuperscript{Tyr} precursors present at this stage.

In contrast, the M1 primer yields cDNA sequences from embryonic stages 38, 24–26, 18–20, and 14–16 that are of the major somatic-type exclusively (data not shown). This implies that the very abundant oocyte-type transcripts found in stage-10 and stage-12 embryos are processed or degraded by the beginning of neurulation. The switch from the oocyte-type to somatic-type tyrosine tRNA is thus largely over by this point. Despite the apparent reactivation of some oocyte-type genes later in development (Fig. 6), most of the tRNA\textsuperscript{Tyr} precursors found in neurula and later-stage embryos exhibit the major somatic-type sequence.

**Discussion**

**Oocyte and somatic tRNA\textsuperscript{Tyr} genes**

We present evidence here that *X. laevis* contains at least two kinds of developmentally regulated tyrosine tRNA genes. These are the oocyte-type genes and are defined as such because their transcripts are found in oocytes; they appear transiently during early embryogenesis, but are absent from post-embryonic somatic cells [Figs. 2, 3, 4, and 6]. As long suspected, one of the oocyte-type genes is TyrC, a member of an unusual and complex multigene family. This tRNA\textsuperscript{Tyr} gene is located, along with seven other tRNA genes, on tandemly repeated 3.18-kb DNA fragments at a single chromosomal locus [Müller and Clarkson 1980; Fostel et al. 1984; Müller et al. 1987]. Although direct evidence is as yet lacking, it seems highly probable that all eight genes on these repeats also are regulated developmentally and are of the oocyte type.

Information on the second oocyte-type tRNA\textsuperscript{Tyr} gene is more limited but, from the sequence gel shown in Figure 4, its transcripts appear to be almost as abundant in ovary RNA as those of the TyrC genes. It seems reasonable to suppose that these transcripts also are derived from highly repeated genes. A possible candidate for their location is a second cluster of tandemly repeated DNA fragments that contain several tRNA genes [Rosenthal and Doering 1983]. Originally, these 3.1-kb repeats were thought to represent a minor tRNA gene cluster, but current estimates suggest that there are \( \sim 800 \) copies of the 3.1-kb repeats and \( \sim 200 \) copies of the 3.18-kb repeats per haploid *X. laevis* genome [J.L. Doering, pers. comm.]. Clearly, it will be of interest to determine whether the 3.1-kb repeats contain a tRNA\textsuperscript{Tyr} gene and, if so, whether this gene matches the second oocyte-type sequence shown in Figure 5.

The two other genes discussed here are designated somatic-type. Their transcripts are not found during oogenesis, perhaps because they are below the detection limit of the present methods, but they are seen clearly from the gastrula stage onward and in postembryonic somatic cells [Figs. 2, 4, and 6]. The TyrD gene, present in only 1–3 copies per haploid genome, is transcribed very actively in somatic S-100 extracts [Gouilloud and Clarkson 1986], yet in vivo it gives rise to only a minor fraction of somatic-type transcripts. The major fraction is derived from another kind of gene whose unspliced precursors exhibit unique 5' leader and intervening sequences (Figs. 4 and 5). Taken together, these results suggest that the major somatic-type transcripts are not derived from a single copy gene, but rather from a small family of \( \sim 20–50 \) highly conserved genes. Nothing is known yet of their organization, but they should be selectable with the oligonucleotide S1 probe (Fig. 5).
The sequences of the four known \textit{X. laevis} tRNA\(^\text{\textsuperscript{Tyr}}\) genes. The non-coding DNA strands of TyrC [Müller and Clarkson 1980] and TyrD (Gouilloud and Clarkson 1986) are aligned with those of the two new tRNA\(^\text{\textsuperscript{Tyr}}\) genes inferred from primer extension sequencing. The anticodon is underlined. The 5'-flanking regions, intervening sequences, and the purines at position 57 are shown in boldface type; vertical bars depict identical nucleotides within these three regions. The intron within the second oocyte-type gene contains several ambiguous positions; the sequence considered most likely is enclosed in brackets. Oligonucleotide S1 is the probe designed to be specific for the major somatic-type tRNA\(^\text{\textsuperscript{Tyr}}\) precursors.

The 5'-flanking sequences of the two oocyte-type genes are somewhat related, as are the comparable regions of the two somatic-type genes (Fig. 5). The intervening sequences within each pair of genes are even more similar, with an 11 out of 13 match for the somatic-type pair and perhaps as much as 10 out of 13 match for the oocyte-type pair. In addition, all four intervening sequences are identical in the last four positions (AGCA, Fig. 5). As noted previously for the TyrC and TyrD transcripts [Gouilloud and Clarkson 1986], these sequences would preclude base-pairing with the entire anticodon, thereby yielding a more open structure than that found in unspliced tRNA precursors from yeast. The presence of these same four nucleotides in two more \textit{X. laevis} tRNA\(^\text{\textsuperscript{Tyr}}\) genes indicates that these sequences are under strong selective pressure. Although recent evidence suggests that the intron sequence is unimportant for the splicing of yeast tRNA precursors by the \textit{Xenopus} endonuclease [Mattoccia et al. 1988], this enzyme may still have a preference for the sequence or for the more open structures of its homologous substrates.

There are indications, however, that the oocyte-type and somatic-type tRNA\(^\text{\textsuperscript{Tyr}}\) precursors are not spliced with equal efficiency. The oligonucleotide C1 probe detects precursors of ~100 nucleotides and ~90 nucleotides that are identified as the TyrC primary transcripts and their unspliced processing intermediates, respectively, whereas the somatic-specific D1 and S1 probes reveal only ~100 nucleotide precursors [Figs. 2 and 6]. Thus, splicing appears to be the rate-limiting processing step for oocyte-type transcripts but not for somatic-type precursors. This may reflect sequence differences in their introns or the purine transition at position 57 of the coding sequence (Fig. 5).

**Developmental changes in tRNA\(^\text{\textsuperscript{Tyr}}\) and 5S RNA**

Transcription is repressed after breakdown of the oocyte nuclear membrane at meiosis, and it remains inactive after fertilization and during the early synchronous cleavage stages of embryogenesis [Newport and Kirchner 1982, and references therein]. When embryos are labeled with \([\alpha-\text{\textsuperscript{32}}\text{P}]\text{GTP}, new tRNA synthesis is found first at the MBT, stage 8 [Wakefield and Gurdon 1983]. Northern blot analysis, which measures RNA steady-state levels rather than synthesis, first detects new tRNA\(^\text{\textsuperscript{Tyr}}\) precursors at late blastula, stage 9, and these are predominantly oocyte-type; the two kinds of somatic-type precursors appear somewhat later, at stage 10.
pressed in tadpoles (Fig. 6); oocyte-type 5S RNA has not been described in these late stages. Finally, and again in the oocyte-type (Fig. 6), which may suggest that all of the oocyte-type tRNA\(^{\text{yr}}\) genes are activated first and then are repressed preferentially by the midgastrula stage. Consistent with this suggestion, oocyte-type transcripts are most abundant at stage 10, in which they constitute the great majority of the tRNA\(^{\text{yr}}\) precursor population, whereas the somatic-type transcripts accumulate somewhat later, at stage 12 (Fig. 6). A second major transition then occurs at the hatching stage, which seems to involve a reactivation of some oocyte-type genes and an increased synthesis or stabilization of both kinds of somatic-type precursors (Fig. 6).

The pattern of expression of tRNA\(^{\text{yr}}\) genes resembles that of the X. laevis 5S RNA genes, in which the 5S RNA population changes from predominantly oocyte-type during oogenesis to somatic-type by the end of gastrulation. The two phenomena differ, however, in at least one important respect. The 5S RNA synthesized in late blastulae comprises a nearly equal mixture of oocyte-type and somatic-type transcripts, which indicates that most of the highly repeated oocyte-type 5S RNA genes are normally not transcribed after oocyte maturation (Wormington and Brown 1983). In contrast, the tRNA\(^{\text{yr}}\) precursors that are first detectable after the MBT are predominantly oocyte-type (Fig. 6), which may suggest that all of the oocyte-type tRNA\(^{\text{yr}}\) genes are activated first and then are repressed selectively. A second potential difference is that some oocyte-type tRNA\(^{\text{yr}}\) genes are expressed in tadpoles (Fig. 6); oocyte-type 5S RNA has not been described in these late stages. Finally, and again in apparent contrast to 5S RNA, the present work failed to detect somatic-type tRNA\(^{\text{yr}}\) precursors in oocytes.

Several mechanisms have been proposed to account for the selective inactivation of the oocyte-type 5S RNA genes. Early models emphasized the importance of the decreasing amounts of the 5S-specific factor TFIIIA during embryogenesis, the early replication of the somatic-type genes, and their greater affinity for TFIIIA (for review, see Brown 1984). All of these have been found wanting in some respects: Oocyte-type 5S RNA repression can occur in the presence of an excess of TFIIIA and in the absence of DNA synthesis (Andrews and Brown 1987), and the major oocyte and somatic genes have equal affinities for TFIIIA (McConkey and Bogenhagen 1988). More recent work suggests that oocyte-type transcription complexes are selectively destabilized in somatic cells, and that a limitation of common RNA polymerase III transcription factors can contribute to this destabilization (for review, see Wolffe and Brown 1988). This partly reflects sequence differences in the internal promoters, but sequences farther upstream also contribute to the differential transcription of oocyte and somatic 5S RNA genes in vitro (Reynolds and Azer 1988).

Some of these mechanisms may function in oocyte-type tRNA\(^{\text{yr}}\) gene repression but an involvement of TFIIIA seems unlikely. This factor is not needed for the transcription of tRNA genes by RNA polymerase III (Lassar et al. 1983), and elevated levels of TFIIIA in developing embryos stimulate 5S RNA synthesis at the MBT but are without effect on tRNA synthesis (Andrews and Brown 1987). These results imply further that two common factors required for 5S RNA and tRNA synthesis, TFIIIB and TFIIIC, are not limiting in early embryogenesis, which may help to explain the very high levels of the oocyte-type tRNA\(^{\text{yr}}\) precursors found in gastrulae (Fig. 6). Whether factor levels then decrease and contribute to the inactivation of the oocyte-type tRNA\(^{\text{yr}}\) genes, is not yet known. Also, it is not known yet whether the oocyte-somatic switch of tyrosine tRNA can be reproduced in embryonic extracts. However, the somatic-type gene TyrD is transcribed more actively than TyrC in somatic S-100 extracts; this differential activity is not the result of intragenic base changes but reflects sequence alterations in the 5' flanking DNA (Gouilhoud and Clarkson 1986). The oocyte-switch of tyrosine tRNA, and perhaps also of 5S RNA, may depend on some critical factor or polymerase interactions with upstream DNA sequences.

Materials and methods

Oocyte and embryonic RNA preparation

Total RNA was isolated from X. laevis staged oocytes (Dumont 1972) and embryos (Nieuwkoop and Faber 1967) by the method of Feramisco et al. (1982) with minor modifications. Frozen oocytes or embryos were homogenized in 1 volume [20 \(\mu\)l per oocyte or embryo] of 4 M guanidinium thiocyanate, 50 mM Tris-HCl (pH 7.6), 10 mM EDTA, 2% sarcosyl, and 140 mM 2-mercaptoethanol and were heated for 15 min at 60°C. Sequentially, 1 volume of 0.1 M sodium acetate [pH 5.2], 10 mM Tris-HCl [pH 7.4], 1 mM EDTA, 1 volume of phenol [pH 8.0], and 1 volume of chloroform-isoamyl alcohol [24 : 1] were added to the homogenate. Samples were left for another 10 min at 60°C with intermittent shaking and then were centrifuged. Each aqueous phase was reextracted once with 1 volume of phenol plus 1 volume of chloroform-isoamyl alcohol, and once with 2 volumes of chloroform-isoamyl alcohol. RNA samples were recovered by ethanol precipitation at \(-20°C\) and were resuspended in water.

Northern blot analyzes

Total RNA from 1–10 oocyte or embryo equivalents was dried down, resuspended in 7 M urea, 0.25 × TEB [1 × is 50 mM Tris-
HCl, 50 mM boric acid, 1 mM EDTA) and fractionated on 2-mm thick 8% polyacrylamide–7 M urea gels in 1 x TEB. Gels were stained for 20 min in transfer buffer [12 mM Tris, 6 mM sodium acetate, 0.3 mM EDTA (pH 7.5)] that contained 1 mg/ml of ethidium bromide, photographed, and then soaked for 25 min in fresh transfer buffer with slow agitation. The RNA was transferred to a nylon membrane [Zeta probe, Bio-Rad] by semi-dry blotting [Kyhse-Andersen 1984] by use of a Multiphor II Nova Blot System [LKB]. After a 1-hr transfer at 0.8 mA/cm², the membrane was rinsed for 30 min in transfer buffer, baked for 2 hr at 80°C, and then prehybridized for 4 hr at 39°C in 0.9 M NaCl, 50 mM Tris-HCl (pH 7.5), 5 mM EDTA, 0.1% NaDodSO₄, and 5 x Denhardt’s [1 x is 0.2% each of bovine serum albumin, Ficoll and polyvinylpyrrolidone] Hybridization was carried out overnight at 39°C after the addition of 5–10 pmol of 5'-end-labeled oligonucleotide probe [3–5 x 10⁶ Cerenkov cpm/pmol] to the prehybridization mix. Filters were rinsed 3 times in 6 x SSC, 0.1% NaDodSO₄ at room temperature, and were washed for 30 min at 39°C in 0.1 x SSC, 0.1% NaDodSO₄ then subjected to autoradiography at ~70°C in the presence of an intensifying screen.

Primer extension analyses

Tyrosine tRNA gene products present in X. laevis ovaries, cultured kidney cells, and staged oocytes or embryos were detected by primer extension analyses by use of a protocol adapted from Reeder et al. (1983) and Dingermann and Nerke (1987). In preliminary experiments with total RNA, the oligonucleotide M1 primer was found to generate a very strong cDNA from mature precursors. Consequently, the following gel purification step was employed prior to reactions with this primer. Total RNA (200 µg) was electrophoresed through a 2-mm thick 8% polyacrylamide–7 M urea gel, a slice corresponding to the tRNA precursor region was excised, and the RNA was recovered by electroelution and was purified further by ion-exchange chromatography [NACS Prepac columns, BRL]. RNA samples were precipitated with 5 pmol of oligonucleotide M1 that had been 5’-end-labeled with [γ-32P]ATP and T4 polynucleotide kinase [Boehringer] to 3–5 x 10⁶ Cerenkov cpm/pmol. The dry pellets were dissolved in 30 µl of 250 mM KCl, 10 mM Tris-HCl (pH 8.0) and were incubated for 2 hr at 25°C. Primer extension was initiated by addition of 50 µl of a buffer containing 24 mM Tris-HCl [pH 8.0], 16 mM MgCl₂, 16 mM DTT, 0.4 mM each of dATP, dCTP, dGTP, and dTTP, 0.16 mg/ml of actinomycin D, and 20 units of AMV reverse transcriptase [Boehringer]. After incubation for 30 min at 25°C and for 30 min at 42°C, the cDNAs were recovered by phenol extraction and ethanol precipitation. Reactions with the M2 primer were performed in the same way except that they contained 200 µg of unfractionated RNA in fourfold greater volumes. Primer extension products were fractionated on 0.5-mm thick 12% polyacrylamide-7 M urea sequencing gels, selected bands were excised and eluted by diffusion in 0.5 M ammonium acetate [pH 7.0], 1 mM EDTA. The cDNAs were ethanol-precipitated and then sequenced by the chemical method of Maxam and Gilbert (1980).

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