Destruction of a translationally controlled mRNA in *Xenopus* oocytes delays progesterone-induced maturation

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The maternal mRNA D7 is a moderately abundant transcript in *Xenopus laevis* whose expression is highest in, and perhaps restricted to, oogenesis and early embryogenesis. The nucleotide sequence of cloned D7 cDNA was determined and shown to have the capacity to code for a 31-kD protein. This amino acid sequence was searched against a protein data base, and no homologous proteins were found. Antibodies directed against D7 recognize in *Xenopus* embryos a soluble, cytoplasmic protein with an apparent molecular weight on SDS gels of 36,000. The D7 protein is absent from oocytes and first begins to accumulate during oocyte maturation. Its levels are highest during the first day of embryonic development and then decrease; D7 protein was not detected in adult tissues. D7 mRNA was selectively destroyed by injection into oocytes of antisense oligodeoxynucleotides. Analysis of injected oocytes by Northern and Western blotting showed site-specific cleavage and subsequent degradation of the D7 mRNA and the failure of the D7 protein to accumulate during progesterone-induced maturation. The loss of D7 protein affects the maturation process itself, significantly delaying the time course of germinal vesicle breakdown. Thus, D7 is a newly described protein involved in oocyte maturation.

[**Key Words:** *Xenopus laevis,* maternal mRNA, translational control, antisense oligodeoxynucleotide, oocyte maturation]

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was carried out. Subsequent in vitro maturation of such oocytes revealed interference in the maturation process, manifested as a pronounced delay in the maturation time course.

Results

Developmentally restricted expression of D7 mRNA and protein

A cDNA clone called D7.0 (originally D7, Dworkin and Dworkin-Rastl 1985) was derived from a moderately abundant X. laevis maternal mRNA (D7 mRNA) that is present throughout oogenesis but decreases in titer to very low levels within 2 days of development (Fig. 1). The amount of this mRNA in the adult tissues examined is at or below the level of detection (Fig. 1). Antibodies to the D7 protein, obtained from rabbits immunized with a fusion protein consisting of part of the putative D7 polypeptide fused to Escherichia coli β-galactosidase [Materials and methods], recognized on Western blots a single 36-kD protein that was expressed in a developmentally restricted pattern [Fig. 2]. D7 protein was absent (at this level of detection) from stage VI oocytes [as well as from stage I–IV oocytes; data not shown], consistent with the nonpolysomal fractionation of D7 mRNA in these cells [Dworkin et al. 1985]. Lysates from unfertilized eggs, however, reacted with the purified antibodies, demonstrating that translation of D7 mRNA and accumulation of D7 protein begin during oocyte maturation. The level of D7 protein was highest in embryos during the first day of development and then decreased, in accordance with the developmental profile of D7 mRNA. No D7 protein was detected in adult muscle or liver.

In situ immunofluorescence of paraffin sections of cleavage embryos with anti-D7 antibodies showed cytoplasmic labeling with no obvious localization within the embryo; nuclei were not stained (data not shown). Subcellular fractionation studies indicated that D7 is a soluble, cytoplasmic protein [data not shown].

Sequence of two D7 cDNA clones and the putative D7 protein

The size of D7 mRNA on Northern blots is ~1200 nucleotides. Because the clone initially isolated contained a cDNA insert of 898 bp [excluding poly(A) and cloning tails], it was not a full-length clone. Therefore, another cDNA library constructed from egg poly(A)+ RNA was screened for additional D7 cDNA sequences. A new clone, D7.1, was thus obtained, and it contained 153 additional nucleotides at the 5’ end but lacked 131 nucleotides at the 3’ end [Fig. 3A], compared with clone D7.0. The cDNA inserts of both clones were completely sequenced [Fig. 3]. The two sequences were identical in the 767 nucleotides of overlapping region, except for two single mismatches (positions 611 and 785; Fig. 3A). Whereas the mismatch at position 785 is a silent change, the mismatch at nucleotide position 611 causes the introduction of a stop codon in clone D7.0 into what is a continuous open reading frame (ORF) in clone D7.1. The shorter ORF of clone D7.0 has a coding capacity for a 22-kD protein, whereas the longer ORF of clone D7.1 has a coding capacity for a 31-kD protein, which is close to the size of the protein recognized by the anti-D7 antibodies [Fig. 2]. To test whether both types of sequences were represented in oocyte and egg mRNA, sequence-
Thus, we conclude that D7 mRNA predominantly (or much stronger with the D7.1-specific oligonucleotide. 

Materials and methods. Hybridization to the RNA was hybridized under differential hybridization conditions to each of the two oligonucleotide probes (Fig. 4; see also Materials and methods). Hybridization to the RNA was much stronger with the D7.1-specific oligonucleotide. Thus, we conclude that D7 mRNA predominantly (or exclusively) consists of molecules of the D7.1 type, lacking the stop codon at nucleotide positions 609–611.

The composite D7 mRNA sequence (Fig. 3B), consisting of the complete D7.1 sequence and the 3' extension derived from D7.0, is 1051 nucleotides long. Primer extension analysis indicated that the D7 transcript extends another 27 nucleotides beyond the 5' end of the cloned cDNA (data not shown). The first ATG codon in the cloned sequence is at nucleotides 27–29 and lies in a good sequence context for translation initiation [Kozak 1987]. The ORF has a coding capacity for 278 amino acids. Because no in-frame stop codon is present 5' to this ATG codon in the cloned sequence, a short extension of the ORF upstream cannot be excluded. The 3'-noncoding region is 191 nucleotides long, is very AT rich (73%), and contains several A[T][A] motifs [n = 3–5] (Fig. 3B). The presence of AU-rich sequences, including AU[A], motifs, in the 3'-noncoding region of eukaryotic mRNAs has been correlated with transcript instability [Shaw and Kamen 1986]. A potential polyadenylation signal [AATATA] is located 33 nucleotides from the 3' end. This suggests that the complete 3'-noncoding sequence is contained in the cDNA clone [D7.0], although this cannot be known with certainty because the cDNA library from which clone D7.0 was isolated had been constructed by AT tailing [Dworkin and Dworkin-Rastl 1985].

The putative polypeptide coded for by the ORF has a calculated pI of 7.00. It is mostly hydrophilic and lacks any long hydrophobic region, consistent with it being a soluble protein (data not shown). Its calculated molecular weight of 31,364 is lower than the apparent molecular weight on SDS gels [36,000] observed for the immunoreactive Xenopus protein. Because the D7 protein synthesized in vitro by translation of egg poly[A]+ RNA and isolated by immunoprecipitation also shows an apparent molecular weight on SDS gels of 36,000 (data not shown), the difference between the apparent and calculated molecular weights is most likely due to aberrant migration on SDS gels. The putative D7 amino acid sequence showed no significant homology to any sequence in the protein sequence data base of the Protein Identification Resource. Although we have not yet directly tested whether D7 is a phosphoprotein, it does not contain an obvious recognition site for protein kinase A [Edelman et al. 1987] and does not itself appear to be a protein kinase [Doolittle 1986; Brenner 1987; Bairach and Claverie 1988]. The D7 protein does contain a region rich in proline [P], glutamic acid [E], serine [S], and threonine [T] ('PEST' region), flanked with the positively charged residues lysine and arginine (amino acids 151–165, Fig. 3B).

Specific oligonucleotide probes were made to the nucleotide region in question for each of the two clones (Table 1). X. laevis oocyte and egg poly[A]+ RNA were hybridized under differential hybridization conditions to each of the two oligonucleotide probes (Fig. 4; see also Materials and methods). Hybridization to the RNA was much stronger with the D7.1-specific oligonucleotide. Thus, we conclude that D7 mRNA predominantly [or

Figure 3. Nucleotide sequence of D7 cDNA and predicted amino acid sequence of D7 protein. [A] Relationship between clones D7.0 and D7.1. The nucleotide positions at the beginning and end of each clone are indicated above the horizontal lines representing the clones in the 5' to 3' direction. The two nucleotide differences between the two clones are shown with an asterisk [*] (see text). The positions complementary to oligonucleotides D7/1–D7/4 are indicated by dashed arrows, A(T)[A] regions by wavy underlines, and the presumptive polyadenylation signal AATATA by a solid underline.

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Oligonucleotide-mediated degradation of D7 mRNA
To analyze the possible function of D7 protein, we attempted to construct phenocopies of null mutants for D7 by eliminating D7 mRNA from the oocyte. Toward
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Table 1. Synthetic oligodeoxynucleotides

| Designation | Sequence (5’ to 3’) | Complementarity |
|-------------|---------------------|-----------------|
| EBI1016     | TTGCCTGCATCATGGGTTT | nucleotides 602–620, specific for D7.0 |
| EBI1017     | TTGCCTGCACCATGGGTTT | nucleotides 602–620, specific for D7.1 |
| D7/1        | ATTACAGCTGCCCCATGTGC | nucleotides 200–219 |
| D7/2        | GGCTCCATGTTACCCCTGT | nucleotides 220–239 |
| D7/3        | TCACTAGTGTTGCTTTGG  | nucleotides 555–574 |
| D7/4        | GATCAAGCTCCTGTTAGGG | nucleotides 179–198 |
| EBI896      | GGTTTGCTAGTAGTGCTAAG | nucleotides 602–620, specific for D7.1 |
| EBI898      | CAGGAGATGGTAAAAGG   | nucleotides 555–574 |

Oligonucleotides EBI1016 and EBI1017 were used as hybridization probes specific for sequences D7.0 and D7.1, respectively; oligonucleotides D7/1–D7/4 were used for injection into oocytes to mediate D7 mRNA degradation; oligonucleotides EBI896 and EBI898 were used as controls in injection experiments (they have no known complementarity to *X. laevis* DNA). The nucleotide positions indicated refer to the D7 sequence given in Fig. 3B.

| Figure 4. | Differential hybridization of oligodeoxynucleotides to D7 SP6 transcripts and *Xenopus* RNA. SP6 transcripts (5 ng per lane) of D7.1 type and D7.0 type (A,C), or poly[A]+ RNA (3 μg per lane) from total ovary (ov) and unfertilized eggs (egg) [B,D] were hybridized under differential hybridization conditions (Materials and methods) with the D7.0-specific oligonucleotide EBI1016 [A,B] or the D7.1-specific oligonucleotide EBI1017 [C,D]. D7 mRNA [D7] is slightly larger in egg RNA than in ovary RNA, probably due to increased length of the poly[A] tail. rRNAs (28S and 18S) show nonspecific reaction with the probes. |

this end, four different 20-mer oligonucleotides [D7/1–D7/4], complementary to D7 mRNA, were synthesized (Table 1; Fig. 3B) and injected into stage VI oocytes. Hybridization of the oligonucleotides to D7 mRNA would likely be followed by cleavage of the RNA–DNA hybrid by an RNase H activity present in oocytes (Cazenave et al. 1987b; Dash et al. 1987; Shuttleworth and Colman 1988). The effect of the injection of D7 mRNA was assayed by extracting total RNA from the injected oocytes and probing for the integrity of D7 mRNA by Northern blot analysis. Injection of anti-D7 oligonucleotides resulted in a discrete shortening of D7 mRNA within 1 hr, the size shift in each case being consistent with an RNase H-mediated cleavage of the RNA–oligonucleotide hybrid (Fig. 5A; with D7/1, D7/2, and D7/4, only the larger 3’-cleavage products are seen on the gels). All oligonucleotides resulted in a reduction in the amount of full-length D7 mRNA. Oligonucleotides D7/2 and D7/3 were more efficient in mediating degradation of D7 mRNA than D7/1 and D7/4, reducing the amount of D7 mRNA to undetectable levels. The RNA cleavage product mediated by oligonucleotide D7/4 was the least stable compared with intermediates generated with D7/1, D7/2, and D7/3 (Fig. 5A). Upon continued incubation of the injected oocytes, these shortened transcripts were degraded (Fig. 5B). There was no evidence of new transcription of D7 mRNA during the incubation times. The same result was obtained when incubation was carried out in the presence of progesterone to induce maturation (see below; data not shown). However, we did observe some variation in the time courses of D7 mRNA cleavage among different batches of oocytes. To control for RNA yield and quality, the RNA blots were either cohybridized with, or stripped of the radioactive D7 probe and rehybridized with, a probe for a different maternal mRNA, A10 (Fig. 5; Dworkin et al. 1985). The observed persistence of A10 mRNA excludes a nonspecific effect of the anti-D7 oligonucleotide on mRNA in general. Also, injection of a control oligonucleotide (EBI898, Table 1), not complementary to D7 mRNA, did not affect D7 mRNA levels (Fig. 5A). Thus, the degradation of D7 mRNA is a sequence-specific event. In some batches of oocytes, two RNA bands hybridized with the D7 probe (Fig. 5B,C). Both transcripts were degraded upon injection of the antisense oligonucleotide.

The oligonucleotide-mediated degradation of D7 mRNA required considerable amounts of oligomer to be effective. In the experiments shown in Figure 5, A and B, 60 ng of oligonucleotide per oocyte was injected. When 20 ng per oocyte [oligonucleotide D7/1] was injected, a size shift occurred in only a fraction of the D7 mRNA.
molecules, and when 7 ng of oligonucleotide per oocyte was injected, no size shift was observed (Fig. 5C). The requirement for such high levels of oligonucleotide for successful RNA elimination could be due to the instability of oligonucleotides in oocytes. To test this, $^{32}$P-end-labeled oligonucleotide [D7/1, 60 ng per oocyte] was injected into oocytes, the oocytes were incubated up to 20 hr, and oocyte lysates were prepared and separated on a 20% DNA sequencing gel [Fig. 6]. Much of the $^{32}$P label was rapidly lost, probably due, in part, to phosphatase activity. However, intermediate-size oligonucleotides were also detected, and by 3 hr, the amount of intermediate-sized oligonucleotides exceeded the amount of remaining 20-mer. After 6 hr of incubation, only very short oligomers remained, as well as a small amount of 20-mer which did persist. Thus, during the course of incubation, the oligonucleotide was largely destroyed, possibly to deoxynucleotides. The gel also shows that initially some lengthening of the 20-mer [to a 21-mer] occurred in the oocytes. Instability of oligonucleotides in Xenopus oocytes has also been reported by Cazenave et al. [1987a].

In untreated oocytes, translation of D7 mRNA is induced during oocyte maturation (Fig. 2). Thus, in oocytes injected with anti-D7 oligonucleotides, D7 protein should not accumulate following induction of maturation. To test this, oocytes were injected with either the anti-D7 oligonucleotides or with a control oligonucleotide [EBI898, Table 1], incubated for 3 hr in progesterone-free medium, and then treated with progesterone. Progesterone induced maturation of the injected oocytes, albeit with a delay in the case of oocytes injected with the D7-specific oligonucleotides (see below). Protein of matured oocytes was assayed by Western blotting with anti-D7 antibodies [Fig. 7]. These blots were also coreacted with antiserum against Xenopus enolase, a protein whose level is unchanged during oocyte maturation [Segil et al. 1988], to demonstrate equal recovery of protein in the different samples. The results show that the level of D7 protein in matured oocytes was reduced with all experimental oligonucleotides; the D7 protein was reduced to background levels with oligonucleotides D7/1, D7/2, and D7/3. Injection of the control oligonucleotide did not affect D7 protein levels. Due to an abundant Coomassie blue-stainable protein at the position of D7 that gives a slight, nonspecific reaction to immunoglobulins, we cannot exclude the presence of low levels of D7 protein. By scanning autoradiographs of Western blots, we have determined that the background signal in stage VI oocytes at the D7 position is 20% the level of the D7 protein signal obtained from untreated in vitro-matured eggs.

Figure 5. Northern blots of total RNA from oocytes injected with antisense oligonucleotides. (A) Comparison of different oligonucleotides. [D7/1–D7/4] Oocytes injected with 60 ng antisense oligonucleotides against D7 mRNA; [898] oocytes injected with 60 ng control oligonucleotide EBI898; [O] uninjected stage VI oocytes. RNA was extracted 1 hr [D7/1–D7/4] or 3 hr [898] after injection. Blots were first hybridized with nick-translated plasmid D7.0, stripped of the probe, and rehybridized with nick-translated plasmid A10 [Dworkin et al. 1985] for the three right-most lanes, a shorter exposure of A10 is shown than for the other lanes. [B] Time course of degradation of D7 mRNA after injection of antisense oligonucleotide. RNA was extracted 3, 5, or 8 hr after injection of 60 ng oligonucleotide D7/1 per oocyte. [O] uninjected stage VI oocytes. [C] Dependence of D7 mRNA degradation on antisense oligonucleotide concentration. RNA was extracted 3 hr after injection of 20 ng or 7 ng oligonucleotide D7/1 per oocyte. The blots shown in B and C were cohybridized with nick-translated plasmids D7.0 and A10. A shorter exposure of the region of A10 hybridization is shown at bottom. [D7] Full-length D7 mRNA; [A10] A10 mRNA. In the oocytes used in experiments B and C, the D7 probe recognized a slightly larger RNA species in addition to the main D7 mRNA band, both RNAs were cleaved after oligonucleotide injection.
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Delayed maturation of oocytes with reduced D7 protein levels

Groups of stage VI oocytes were injected with 60 ng per oocyte of oligonucleotides D7/1–D7/4 or with the control oligonucleotides EBI896 or EBI898 (Table 1). After a minimum of 3 hr of incubation, the oocytes were transferred to medium containing progesterone. Maturation was scored visually under a dissecting microscope for formation of the white maturation spot. The results of six experiments (with oocytes from six different frogs) are shown in Table 2. In all experiments shown, uninjected oocytes as well as oocytes injected with the control oligonucleotides underwent 100% germinal vesicle breakdown (GVBD). Oocytes that had been injected with antisense oligonucleotides against D7 mRNA exhibited delays in the maturation time course. The extent of the delay depended on the oligonucleotide used (Fig. 8; Table 2), the most effective oligonucleotide being D7/2, followed by oligonucleotides D7/1 and D7/4. These oligonucleotides not only delayed GVBD but also caused a flattening of the slope in the curve of the maturation time course (Fig. 8). Although the delay of progesterone-

**Table 2. Delay of progesterone-induced maturation after injection of anti-D7 mRNA oligonucleotides**

| Experiment | Oligonucleotide | Time          | Percent mature |
|------------|-----------------|---------------|----------------|
| 1          | none            | 3 hr, 45 min  | 100            |
|            | EBI898          | 3 hr, 45 min  | 100            |
|            | D7/1            | 3 hr, 45 min  | 50             |
|            | D7/2            | 3 hr, 45 min  | 5              |
|            | D7/3            | 3 hr, 45 min  | 83             |
|            | D7/4            | 3 hr, 45 min  | 43             |
| 2          | none            | 5 hr          | 100            |
|            | EBI898          | 5 hr, 25 min  | 100            |
|            | D7/1            | 5 hr, 25 min  | 25             |
| 3          | none            | 10 hr, 30 min | 100            |
|            | EBI896          | 10 hr, 30 min | 100            |
|            | D7/2            | 10 hr, 30 min | 6              |
| 4          | none            | 6 hr, 30 min  | 100            |
|            | EBI896          | 5 hr          | 100            |
|            | D7/1            | 6 hr, 30 min  | 9              |
|            | D7/2            | 10 hr, 15 min | 9              |
| 5          | none            | 9 hr, 30 min  | 100            |
|            | EBI896          | 9 hr, 30 min  | 100            |
|            | D7/1            | 8 hr, 30 min  | 8              |
|            | D7/2            | 8 hr, 30 min  | 4              |
| 6          | none            | 8 hr, 30 min  | 100            |
|            | EBI896          | 8 hr, 30 min  | 100            |
|            | D7/1            | 8 hr, 30 min  | 10             |
|            | D7/2            | 8 hr, 30 min  | 4              |

Experiments 1 and 3 are identical to the experiments shown in Figs. 8 and 9, respectively. In experiments 1–5, 60 ng of oligonucleotide per oocyte was injected, and in experiment 6, 130 ng of oligonucleotide per oocyte was injected. GVBD was scored by visual inspection of maturation spots in experiments 1 and 2, by analysis of paraffin sections in experiments 3–5, and by dissection of trichloroacetic acid-fixed oocytes in experiment 6. The earliest time when control oocytes showed 100% maturation is given.

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induced maturation with D7/1- and D7/2-injected oocytes was very reproducible, the exact extent of the delay varied among different batches of oocytes. In some experiments, such as experiment 4 (Table 2) in which experimental oocytes (D7/2) showed <10% maturation after 10 hr in progesterone [control oligonucleotide-injected oocytes showed 100% GVBD by 5 hr], maturation may have been essentially inhibited rather than only delayed. Using higher amounts of oligonucleotide D7/2 [130 ng per oocyte], it was possible to achieve complete inhibition of maturation (experiment 6, Table 2). In this experiment, all but one of the oocytes injected with the oligonucleotide failed to mature within the time of the experiment [36 hr in progesterone], whereas uninjected oocytes or oocytes injected with control oligonucleotide [130 ng per oocyte] reached 100% GVBD after 8.5 hr in progesterone. It is likely, then, that D7 mRNA was completely eliminated in this experiment (see Discussion).

Oligonucleotide D7/3 affected the time course of oocyte maturation only slightly (Fig. 8; Table 2), even though this oligonucleotide was very effective in causing cleavage of D7 mRNA (Fig. 5A) and reducing D7 protein levels in the matured oocytes (Fig. 7). Because oligonucleotide D7/3 hybridizes to a region in D7 mRNA much farther 3' than the other oligonucleotides tested (Fig. 3B), it may be that the cleavage product of the D7 mRNA can still code for a functional [but truncated] polypeptide. Such a truncated polypeptide was not detected, however, in Western blot experiments (Fig. 7).

In addition to undergoing delayed maturation, oocytes injected with oligonucleotides D7/1, D7/2, or D7/4 ('experimental oocytes') displayed a matured phenotype distinct from that of uninjected oocytes or oocytes injected with oligonucleotides EB1896 or EB1898 ('control oocytes'), or oocytes injected with the ineffective [with respect to delaying maturation] oligonucleotide D7/3. In particular, whereas control oocytes usually developed obvious white maturation spots surrounded by a ring of darker pigment, the appearance of the maturation spot in experimental oocytes was varied, depending on the particular batch of oocytes. In general, white spot formation was less distinct in experimental oocytes than in control oocytes, and frequently the animal hemisphere developed a mottled or 'swirly' appearance. Because of these pigment effects, the scoring of maturation spots under the dissecting microscope was often difficult in experimental oocytes; therefore, in some experiments the percentage of GVBD in a particular population of oocytes was scored histologically, after embedding the oocytes in paraffin and sectioning, or after fixation of oocytes in trichloroacetic acid and subsequent dissection (Table 2). The histological determination of GVBD almost always confirmed the prior evaluation of oocytes made under the dissecting microscope.

The experiment shown in Figure 5A had demonstrated that different oligonucleotides were differentially effective in mediating D7 mRNA degradation. These oligonucleotides caused delays in progesterone-induced maturation to different extents (Fig. 8; Table 2), consistent with their respective efficiencies for mRNA degradation [except for D7/3, above]. This suggested that the extent of maturation delay was a function of residual D7 mRNA. We addressed this further by determining the relationship between the amount of a given oligonucleotide injected and the resultant delay in the maturation time course. As shown in Figure 5C, the efficiency of mRNA degradation depended very sensitively on the amount of oligonucleotide injected. Oligonucleotide D7/2 was injected into oocytes at three different concentrations (up to 60 ng per oocyte in this experiment), the oocytes were incubated overnight (>9 hr) in medium without progesterone, transferred to progesterone-containing medium, and the maturation time courses were followed by visual inspection under the dissecting microscope (Fig. 9). When control oocytes had reached 100% GVBD, all oocytes were fixed and processed for histological determination of GVBD. At this time, only 6% (1 of 18) of the oocytes that had been injected with 60 ng of oligonucleotide D7/2 had matured, whereas in the same time period, 50% (8 of 16) of the oocytes injected with 30 ng, and 100% (14) of the oocytes injected with 15 ng had matured. These results demonstrate a dependence of the delay in the maturation time course on the amount of oligonucleotide injected, which, in

Figure 8. Delay of in vitro maturation after injection into oocytes of antisense oligonucleotides against D7 mRNA. Stage VI oocytes were injected with 60 ng of antisense oligonucleotides against D7 mRNA (D7/1–D7/4) or 60 ng of the control oligonucleotide EB1898 (control), incubated for 3 hr, and induced to mature by the addition of progesterone. The number of oocytes injected per oligonucleotide was ≥21. The time courses of maturation were determined by scoring the formation of maturation spots under the dissecting microscope. Noninjected and control oligonucleotide-injected oocytes showed identical maturation time courses.
At this concentration, the oligonucleotide is present at a molar excess of about 10⁵ over a moderately abundant mRNA. At such high levels of oligonucleotide, the distribution of the oligonucleotide to its complementary region on the RNA which, in turn, suggests a dependence on the amount of D7 protein allowed to accumulate in these oocytes.

Discussion

These experiments demonstrate that in *X. laevis* stage VI oocytes, an endogenous mRNA that is translationally repressed at this stage can be destroyed by injection of an antisense oligodeoxynucleotide. Oligonucleotide-mediated mRNA destruction in *X. laevis* had been shown previously for exogenous RNAs, as well as recently for endogenous RNAs (see introductory section). The efficiency of mRNA destruction in these studies varied for different transcripts and oligonucleotides. It is likely to depend on the accessibility of a given oligonucleotide to its complementary region on the RNA which, in turn, is determined by the secondary structure of the RNA and its association with protein. Especially in the case of an mRNA that is translationally repressed in oocytes (like D7 mRNA), it is likely that the RNA is tightly complexed with protein, limiting the regions available for hybridization to the oligonucleotide. In this context, it is actually surprising that all four oligonucleotides tested for their ability to mediate degradation of D7 mRNA were effective, although their exact efficiencies varied.

We have also tested this approach with other mRNAs that are translationally repressed in oocytes and have encountered oligonucleotides completely inefficient in this assay [E. Dworkin-Rastl, unpubl.]. The extent of mRNA destruction depends very sensitively on the amount of oligonucleotide injected, with differences by factors of 2–4 having a strong effect (see also Dash et al. 1987; Jessus et al. 1988, Shuttleworth and Colman 1988). In our work, as well as in the studies cited above, oligonucleotide amounts in the range of 25–180 ng per oocyte were needed for successful mRNA destruction. At this concentration, the oligonucleotide is present at a molar excess of about 10⁶ over a moderately abundant RNA like D7 mRNA. The reason for the requirement for such high levels of oligonucleotide may be slow diffusion of the oligonucleotide, as well as a low rate of hybridization that would be expected in oocytes at ambient temperature. Furthermore, we and others (Cazenave et al. 1987a) have shown that oligonucleotides are very short-lived in *Xenopus* oocytes. The initial mRNA degradation products obtained with the different oligonucleotides are discrete and consistent with an RNase H-mediated cleavage mechanism (Dash et al. 1987, Shuttleworth and Colman 1988). RNase H cleavage products are unstable in the oocyte and eventually become undetectable by Northern blot analysis. We have observed some variability in the efficiency of mRNA degradation by a given oligonucleotide with different batches of oocytes, which could be a reflection of variation in RNase H levels.

The possibility to generate oocytes specifically depleted of D7 mRNA provides a basis for studying the function of the D7 protein product. In untreated oocytes, D7 protein is first detectable during oocyte maturation. In experimental oocytes injected with an effective anti-D7 mRNA oligonucleotide (e.g., D7/2) prior to progesterone treatment, the accumulation of D7 protein was reduced to background levels. These oocytes exhibited a significant increase in the time needed for oocyte maturation, or, in several experiments, never achieved a high percentage of maturation during the time the experiment was followed (see Table 2). It is possible that the extent of delay in maturation is a reflection of the efficiency of D7 mRNA destruction. This hypothesis is consistent with the observed correlation between the efficiencies of different anti-D7 mRNA oligonucleotides in mediating mRNA degradation and the different extents of delay in maturation resulting from these oligonucleotides (an exception to this correlation was oligonucleotide D7/3, as described in Results). Further evidence in support of this hypothesis is the observed dependence of the maturation delay on the amount of a given oligonucleotide injected. Thus, our current hypothesis is that D7 protein is required at a certain point in the maturation process and that maturation cannot proceed beyond this point unless a threshold amount of D7 protein is provided. This threshold amount must be <20% of the 'wild-type' D7 protein level, the background level of D7 protein in Western blots (see Results). However, on the...
basis of the current data, it cannot be excluded that D7 has only a nonessential role in maturation and represents a protein whose presence, not strictly required, only speeds up maturation. The increased accumulation of D7 protein after fertilization might indicate an additional function of D7 during early development. A more detailed analysis of the D7 null phenocopy is in preparation.

Materials and methods

RNA preparations and Northern analysis

Embryos were staged according to Nieuwkoop and Faber [1967] and oocytes according to Dumont [1972]. RNA was extracted from staged embryos, staged oocytes, or total ovary, and poly(A)+ RNA was prepared as described in Dworkin and Dawid [1980]. For analysis of oligonucleotide-mediated RNA degradation, total RNA was prepared from samples of three oocytes each. Northern analysis with nick-translated plasmid probes was carried out as described previously [Dworkin et al. 1984], except that hybridization and wash conditions were according to Church and Gilbert [1984].

Differential oligonucleotide hybridization

Synthetic RNA transcripts were made with SP6 RNA polymerase [Boehringer–Mannheim] according to Melton et al. [1984], using as templates [1] a plasmid containing the PstI–HindIII fragment of clone D7.0 [nucleotides 570–999, Fig. 3B] inserted into the respective sites of the vector pGEM4 (Promega Biotech) and linearized with PvuII to yield an SP6 transcript of ~600 nucleotides, and [2] plasmid D7.1, linearized with Scal to yield an SP6 transcript of ~2900 nucleotides. Differential hybridization conditions were determined using oligonucleotides EBI1016 and EBI1017 [specific for sequence D7.0 and D7.1, respectively, Table 1] end-labeled with T4 polynucleotide kinase, followed by extensive dialysis at 4°C against water systems DNA synthesizer, purified by reverse-phase high performance liquid chromatography (HPLC), following the protocol of Applied Biosystems, and dissolved in water. The radio-labeled oligonucleotide was prepared by end labeling trace amounts of D7/I oligonucleotide [Table 1] with T4 polynucleotide kinase, followed by extensive dialysis at 4°C against water (dialysis tubing: molecular weight cutoff of 3500) and lyophilization. The stability of oligodeoxynucleotides in stage VI oocytes was determined by injecting 60 ng [26,000 cpm] of D7/I oligonucleotide per oocyte, incubating oocytes for various times, and quick-freezing oocytes at ~70°C. Lysates from individual oocytes were prepared by crushing each oocyte in 12 µl sequencing gel sample buffer [Maxam and Gilbert 1980], boiling for 2 min, and centrifuging in the microfuge for 3 min. An aliquot of the supernatant containing 5000 cpm was analyzed by separation on a 20% DNA sequencing gel and subsequent autoradiography.

Screening of an egg cDNA library, DNA sequencing, and computer programs

An egg cDNA plasmid library, prepared from poly(A)+ RNA isolated from unfertilized eggs by GC tailing into the vector pGEM4 [Promega Biotech], was screened for D7 sequences using a nick-translated insert fragment of clone D7.0 [NcoI–EcoRI fragment, nucleotides 230–458, Fig. 3B]. DNA sequences were determined on both strands by either the method of Maxam and Gilbert [1980] or by chain-termination sequencing of double-stranded DNA templates [Chen and Seeburg 1985], using a Sequenase sequencing kit (USB) and oligodeoxynucleotide primers. Homology searches were performed of the Protein Identification Resource data bank [release 10.0], using the FASTP program [Lipman and Pearson 1985]. Calculations of pI value and hydrophobicity utilized the IBI DNA/protein sequence analysis software package.

Expression vector construction and fusion protein isolation

The BstEI–HindIII fragment of clone D7.0 [nucleotides 225–999, Fig. 3B], made blunt ended at the BstEI site by fill in with reverse transcriptase, was cloned between the BamHI site [filled in with reverse transcriptase] and the HindIII site of the β-galactosidase fusion protein vector pUR292 [Rüther and Müller-Hill 1983]. This resulted in an in-frame fusion of the D7 fragment to the carboxy-terminal end of β-galactosidase. The resulting plasmid was used to transform E. coli K-12 strain 71-18 [Messing et al. 1977]. Fusion protein was induced by the addition of isopropyl-β-thiogalactoside [Pharmacia] to a final concentration of 1.5 mM to log-phase bacteria [OD600 = 1.0] and grown for an additional 2 hr at 37°C. Fusion protein was purified from the supernatant fraction of the bacterial lysate by affinity chromatography on a p-aminophenyl-β-thiogalacto-pyranoside–agarose column [Sigma] according to Ullmann [1984]. Rabbits were immunized, and the resulting antisera was affinity-purified in two steps to obtain antibodies directed against D7: [1] by eliminating antibodies binding to β-galactosidase, and [2] by selecting for antibodies binding to the fusion protein. Details of the procedures used for fusion protein isolation are described elsewhere [Smith et al. 1988]. Serum from two rabbits immunized with the fusion protein gave indistinguishable results on Western blots.

Protein extracts and Western blots

Eggs and embryos were dejellied in 2% cysteine-HCl (pH 8). Larger-scale extracts of oocytes, eggs, embryos, and adult tissues were made as described previously [Segil et al. 1988]. For the analysis of oligonucleotide-injected oocytes, protein was prepared from samples of two to three oocytes by homogenizing the cells on ice in 25 µl full-strength modified amphibian Ringer’s solution [Vincent et al. 1986]. The homogenate was centrifuged for 30 sec at 4°C in a microfuge, and the supernatant analyzed by Western blotting. Western blots were done as described previously [Segil et al. 1988], using 7 µg/ml affinity column-purified anti-D7 antibodies and, where indicated, antisera against X. laevis enolase [Segil et al. 1988] at a dilution of 1:5000.

Oocyte injections, in vitro maturation, and histology

Oocytes were freed manually from the ovary in full-strength

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modified amphibian Ringer’s solution (Vincent et al. 1986), and stage VI oocytes were selected for injection. Up to 80 nl of oligonucleotide solution per oocyte was injected; the amounts of oligonucleotide injected are given in Results. Oocyte maturation was induced by the addition of 1-2 μg/ml progesterone in Ringer’s (above) containing (in some experiments) 1% calf serum or fetal calf serum. For histological evaluation of maturation, the oocytes were fixed overnight in 100% cold (4°C) methanol, embedded in paraplast, and sectioned; sections were stained with Giemsa solution (Merk). Alternatively, oocytes were fixed in 5% trichloroacetic acid for 5 min, bisected with a blade, and maturation evaluated under the dissecting microscope.

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Note

Sequence data described in this paper have been submitted to the EMBL/GenBank Data Libraries.

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