Positive and negative regulation of the gene for transcription factor IIIA in *Xenopus laevis* oocytes

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Expression of the positively acting 5S gene-specific transcription factor, TFIIIA, is regulated during development, with highest levels of mRNA and protein occurring during oogenesis. By analysis of TFIIIA promoter mutants microinjected into late stage *Xenopus* oocytes, we have determined DNA sequences required for the transcription of this gene and we have identified proteins that bind to these regulatory sequences. A negative element lies between positions -306 and -289. Three positive-acting sequences are located between positions -289 and -253, -250 and -173, and -144 and -101. Gel shift analyses of TFIIIA promoter fragments incubated with *Xenopus* oocyte extracts have identified two DNA-protein complexes. One complex, designated B1, requires sequences within the promoter region extending from -271 to -253 while the second complex, designated B2, involves promoter sequences from -235 to -221. The protein involved in formation of the B1 complex has been found to be related to the human adenovirus major late transcription factor, USF.

[Key Words: TFIIIA; transcription; cis-acting elements; trans-acting factors; developmental control; *Xenopus laevis*]

Received February 3, 1989; revised version accepted March 8, 1989.

Regulation of eukaryotic gene expression frequently is achieved at the level of transcription via the interaction of specific proteins with sequence elements located within the promoter region [Maniatis et al. 1987]. Quantitative and/or qualitative changes in specific DNA-binding proteins can positively or negatively modulate the temporal and spatial pattern of gene expression during maturation of the organism. The expression of the 5S ribosomal RNA genes during the development of the South African clawed toad, *Xenopus laevis*, has been characterized extensively [Engelke et al. 1980; Korn 1982; Shastry et al. 1982; Wormington et al. 1982] and provides a model system for the analysis of developmental control at the level of transcription. The primary step required for expression of the 5S genes in *Xenopus* is the binding of a positively acting gene-specific transcription factor, TFIIIA, to the internal control region of the 5S gene [Bogenhagen et al. 1980; Engelke et al. 1980; Bieker and Roeder 1984]. TFIIIA binding permits the subsequent formation of a transcription complex involving at least two other transcription factors, TFIIB and TFIIC, and RNA polymerase III [Lassar et al. 1983; Bieker et al. 1985]. TFIIIA was the first eukaryotic transcription factor that was purified to homogeneity and for which a cDNA was isolated and the corresponding primary amino acid sequence obtained [Ginsberg et al. 1984]. On the basis of this sequence, Miller et al. [1985] proposed a TFIIIA structure with several zinc-binding ‘fingers’ which are involved in DNA binding. This structure has since been identified in other DNA binding proteins [Kadonaga et al. 1988].

The TFIIIA protein serves a dual role in 5S gene expression, acting both as a 5S gene-specific transcriptional activator [Engelke et al. 1980] and as a specific RNA-binding protein that interacts with 5S RNA to form the 7S ribonucleoprotein storage particle [Honda and Roeder 1980; Pelham and Brown 1980]. During the development of *X. laevis*, a distinct pattern of expression of two classes of 5S RNA genes is observed: whereas the somatic 5S gene family is expressed in both oocytes and somatic cells, expression of the oocyte 5S genes decreases dramatically following oogenesis. Although the differential regulation of the two 5S gene classes involves several control mechanisms [Andrews and Brown 1987; Wolffe and Brown 1987, 1988], the demonstration that TFIIIA is required for transcription of both 5S gene families [Engelke et al. 1980] suggests that an alteration in the level or the activity of TFIIIA may play an important role in the regulation of 5S gene expression during development.

The expression of the TFIIIA gene in *Xenopus* has also been shown to be under developmental control. Both TFIIIA protein and steady-state mRNA levels are highest in early oogenesis, decrease significantly in later-stage oocytes, and are reduced drastically during embryogenesis and in somatic cells. Thus, TFIIIA protein levels are reduced from 10^{12} to 10^{9} molecules per cell, whereas RNA levels are reduced from 10^{6} to <1 molecules per...
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cell during development [Ginsberg et al. 1984; Shastry et al. 1984, Taylor et al. 1986]. The high levels of TFIIIA protein in oogenesis appear necessary to ensure full activation of both the oocyte type and the somatic type 5S genes as well as stabilization of the RNA product, whereas the low level in somatic cells may be in part responsible for the restriction of activation to somatic genes. An understanding of the mechanism by which TFIIIA gene expression is regulated during Xenopus development would provide insight into an apparent hierarchical network of developmental control in which a gene that encodes a transcriptional regulatory factor is itself regulated in a developmental fashion.

To investigate the process of TFIIIA gene regulation during Xenopus development, we have analyzed the transcription of this gene in oocytes. In this paper we describe the isolation and characterization of a TFIIIA genomic DNA clone and the identification of cis-acting promoter sequences that act both positively and negatively to affect expression of the TFIIIA gene in Xenopus oocytes. Furthermore, using gel retardation assays, we identify DNA-binding proteins that interact with these cis-acting sequences, and we discuss how they may interact to modulate the expression of TFIIIA during development.

Results
Isolation and characterization of TFIIIA genomic clones
A X. laevis genomic library was screened using the TFIIIA cDNA clone, 3a1.b, as probe [Ginsberg et al. 1984]. A homologous clone containing the 5′ end of the cDNA probe was isolated and characterized further. A gross restriction map of this clone is depicted in Figure 1A. While generally in agreement with the TFIIIA allele described by Tso et al. [1986], restriction analysis of this gene indicates the presence of at least one additional EcoRI site. The 5′ end of the mRNA was mapped within a 2.2-kb genomic EcoRI–PstI fragment by primer extension analysis and the region from position −425 [relative to the transcription start site] through the first exon was sequenced [Fig. 1B]. This gene lacks the PstI site within the first exon, which is present in the allele described previously by Taylor et al. [1986] and displays heterogeneity in the 5′ leader sequence when compared to the allele described by Tso et al. [1986]. A consensus TATA sequence is found at position −32 and a putative CAAT box homology begins at position −94.

Positive and negative cis-acting elements are involved in TFIIIA gene expression in oocytes
We have begun an analysis of the nature and location of the 5′-flanking sequences that are involved in the transcription of this gene following microinjection into X. laevis oocytes [stages V and VI]. To facilitate later analysis, we replaced most of the body of the TFIIIA gene with a fragment derived from the human β-globin gene [Poncz et al. 1983]. The chimeric constructs contain a transcription start site derived from the TFIIIA gene in addition to a part of the second and the third exon of the β-globin gene, including an intron and a functional polyadenylation signal [Fig. 1C].

A set of sequential exonuclease Bal31-generated 5′ deletions was constructed within the TFIIIA promoter region; the exact end point of each deletion was determined by DNA sequencing [Fig. 1D]. Deletion mutants, along with a marked X. laevis histone H4 gene, were analyzed following coinjection into stage V and VI oocytes. Primer extension of a properly initiated TFIIIA transcript generates a fragment 109 nucleotides in length while the extension product for the H4 control gene is 181–187 nucleotides in length. The result of a typical analysis is shown in Figure 2A. Figure 2B shows the level of transcription of each deletion mutant relative to the level observed for the H4 control gene. Each number represents the average of a minimum of four independent experiments. In general, the same pattern of expression was seen whether low [400 pg] or high [2 ng] concentrations of DNA were injected.

Comparison of the level of expression directed by the −1500 and −306 constructs indicates that sequences downstream of −306 are adequate for efficient expression of the TFIIIA gene, although the possibility of other functional elements occurring upstream of −306 is not necessarily excluded. Deletion of sequences downstream from −306 revealed several major cis-acting elements in addition to the TATA box. Thus, removal of sequences between position −306 and −300 resulted in a substantial increase in transcription from the mutant promoter, indicating the presence of an element which has a negative influence on transcription. Deletion to −289, −267, and −265 progressively decreased transcription to the level observed with the wild-type (−1500) promoter. Further deletion through −253 to −235 decreased transcription still further [to about 25% of control levels]. These results indicate the presence of one or more positive cis-acting elements within these regions. It should be noted that in several independent experiments, the −175 deletion mutant promoted a somewhat higher level of transcription than did the −235 mutant, suggesting the presence of another negative element. The −148, −76, and −49 deletion mutants directed approximately the same low level of expression; therefore, in the absence of upstream sequence elements, elimination of the putative CAAT box at position −94 by 5′ deletion had no significant effect on transcription. Deletion of the TATA box at position −32 abolishes transcription from the TFIIIA promoter.

To investigate further the region downstream of the cis-acting elements between −306 and −235, as well as to identify potential cis-acting elements whose function is dependent on the presence of upstream sequences, several internal deletion mutants were constructed using the −306 TFIIIA promoter mutant as a template. Some portion of the deleted sequence was replaced with linker sequence. Analysis of these mutants by microinjection both confirmed and extended the conclusions drawn from the 5′ deletion studies [Fig. 3]. Of the nine
Figure 1. (A) A gross restriction map of the TFIIIA gene showing orientation of the EcoRI fragments within the λ genomic clone. Symbols for restriction sites are: (RI) EcoRI; (H) HindIII; (P) PstI; (B) BglII; (R) RsaI; (Sc) SacI; (HIII) HindIIIII; (S) SalI. The two EcoRI sites flanking the insert represent artificial linkers added during cloning. The 3' end of the gene, as defined by the cDNA clone 3αl.b, lies in a 3.5-kb EcoRI fragment contained within a second nonoverlapping genomic clone. (B) Nucleotide sequence of the BglII-RsaI fragment. The CAAT and TATA boxes are marked. The stars indicate the sequence heterogeneity in the 5' leader as compared with Tso et al. (1986). (C) Structure of the chimeric TFIIIA/β-globin gene. The region of the TFIIIA gene extending from −306 with respect to the transcription start site (+1) [single line] through +88 within the first exon [stippled box] was fused to a 1689-bp segment of the human β-globin gene, which extended from a BamHI site within the second exon to a PstI site within the 3'-flanking sequence [striped boxes designate β-globin exon sequence; the intron sequence appears as a triple line]. The TFIIIA ATG and the β-globin polyadenylation signal are indicated. (D) 5' End points of various TFIIIA promoter deletion mutants. Positions of the CAAT and TATA boxes are indicated.
internal deletion mutants [IDM] examined, IDM 4 (-181/-173 deletion), IDM 7 (-75/-58 deletion), and IDM 8 (-64/-48 deletion) promoted the same level of expression as the -306 deletion mutant, which will be considered wild type in describing the following analyses. Therefore, internal deletion of sequences between -181 and -173 and between -75 and -48 did not affect transcription of the microinjected gene. However, IDM 2 (-223/-173 deletion) and IDM 3 (-214/-173 deletion) were transcribed at only 49% and 53% of the wild-type levels, respectively, whereas IDM 1 (-250/-226 deletion) reduced transcription to 17%. Taken together, these data define at least one cis-acting element lying between -250 and -173 within the TFIIIA promoter. Deletion of the region from -144/-101 [IDM 5] reduced transcription to 42%, suggesting the presence of yet another positive-acting element within this region. Removal of internal sequences containing the CAAT box [IDM 6, -93/-78] reduced transcription approximately twofold, whereas deletion of the region containing the TATA box [IDM 9, -38/-4] abolished transcription.

Identification of protein factors that bind within the TFIIIA promoter

Because much of the expression of the TFIIIA gene in stage V and VI oocytes was dependent on sequences within the region spanning -306 to -175, we attempted to identify by gel retardation assays any oocyte proteins that bind within this region. A double-stranded promoter fragment that included sequences from -306 to -180 was labeled and incubated with a crude cellular extract prepared from a whole Xenopus ovary containing
Figure 3. (A) Primer extension analysis of total RNA from oocytes microinjected with mutants containing internal deletions in the TFIIIA promoter. (B) Quantitation of primer extension analyses of IDM. Values represent the percent of transcription of each mutant relative to the wild-type -306 construct and normalized to the expression of the H4 control gene. Sequences deleted and spacing changes generated are shown below the graph. Details on the internal deletion mutants are given in Materials and methods.

Oocytes representing all six stages of oogenesis. Following electrophoresis on a 6% native acrylamide gel, two retarded complexes, designated B1 and B2, could be identified [Fig. 4]. To prove that the association of these proteins within the TFIIIA promoter was sequence specific and to delimit the regions in which we bind, we tested the ability of various DNA fragments derived from deletion mutants of the TFIIIA promoter to compete for these interactions. A fragment containing promoter sequences extending from -289 to +88 was able to compete for binding of the proteins complexed in both B1 and B2. However, similar fragments with 5' end points at -265, -253, and -235 were able to compete only for the B2 complex. The -175 and -148 deletion fragments were unable to compete for either complex. Additional assays using double-stranded oligonucleotides as competitors [Fig. 5A, B] showed strong competition for the B1 complex with sequences spanning positions -289 to -252 and -271 to -252 and less competition with an oligonucleotide spanning -306 to -265. The B2 complex was competed with an oligonucleotide containing sequences from -252 to -221. An oligonucleotide spanning positions -220 to -180 did not compete for either complex. These results indicate that the binding site for the protein involved in the formation of the B1 complex includes the region between -271 and -265, and that sequences between -265 and -252 stabilize this interaction. The binding site for the B2 complex lies downstream of -235 and is disrupted or eliminated by removal of sequences between -235 and -221.

**The B1 binding site shares homology with the consensus sequence for the adenovirus major late transcription factor, USF.**

Analysis of the nucleotide sequence in the region involved in the formation of the B1 complex revealed a partial homology (-269 5'-CACGTG-3' -264) to the binding sequence for the adenovirus major late transcription factor, USF [Carthew, et al. 1985; Sawadogo and Roeder 1985]. Preincubation with an oligonucleotide containing the human USF consensus binding site, 5'-GGCCACGTGACC-3', inhibited formation of the B1 complex, indicating that this sequence is indeed involved in the binding of the B1 protein[s]. Competition with a nonspecific oligonucleotide containing an octamer transcription factor -1 binding site [Fletcher et al. 1987] had no effect on the formation of the complex.

Figure 4. Detection of two prominent DNA binding complexes in *X. laevis* whole ovary extracts. A 32P-labeled probe spanning the TFIIIA promoter region from -306 to -180 was incubated with 30 μg of whole ovary extract in the standard gel electrophoresis DNA binding assay. Two complexes, designated B1 and B2, could be resolved on a 6% polyacrylamide gel. Competitor fragments (100 ng, 25- to 50-fold molar excess) were isolated from TFIIIA 5' deletion mutants and included sequences from the position indicated at the top of each lane through +88.
Figure 5. (A) Double-stranded oligonucleotides used as competitors in gel shift analysis. Position within the TFIIIA promoter is shown above each sequence. BamHI restriction sites flank each oligonucleotide. (B) The TFIIIA promoter fragment from -306 to -180 was incubated with 30 μg of whole ovary extract under standard gel shift conditions. Competitor oligonucleotides (200 ng) [500- to 1500-fold molar excess) were added as designated above each lane.

(Fig. 6A). To substantiate further the relationship between the Xenopus B1 complex protein(s) and human USF, a partially purified preparation of USF (the kind gift from M. Sawadogo, Rockefeller University, New York) was substituted for the oocyte extract in a gel shift assay (Fig. 6B). Clearly, the human USF bound to the TFIIIA promoter, generating the two retarded complexes characteristic of the binding of USF to the major late promoter (Sawadogo 1988; Sawadogo et al. 1988). The shifted bands have approximately the same Rf value as does the B1 complex [data not shown].

The region between -235 and -221 (5'-AGGG-GAGTGTCCAGA-3'), within which the B2 protein binds, contains a 9- out of 12-bp homology [bases underlined] to the consensus binding sequence for the NFκB protein [5'-GGGGACTTTCCG-3'] (Sen and Baltimore 1986). However, no competition for B2 complex formation was observed in gel shift analyses with oligonucleotides containing the NFκB consensus sequence, the related binding sequence for the ubiquitous transcription factor H2TF1 [5'-GGGGAGTTCCC-3'] (Baldwin and Sharp 1987), or the immunoglobulin heavy-chain NTF protein [5'-GGGGAGCTTCCC-3'] (B. Yoza and R.G. Roeder, in prep.). Furthermore, purified NFκB protein [the kind gift of K. Kawakami, Rockefeller University, New York; Kawakami et al. 1988] did not bind within the B2 region [data not shown].

Point mutations affect binding of B1 and B2 proteins as well as efficiency of transcription of microinjected constructs

To delineate further the sequences required for formation of the B1 and B2 complexes, as well as to verify the requirement for these sequences in the efficient expression of the TFIIIA promoter in vivo, point mutations were made within the binding regions delimited by the gel shift analyses. In the two mutants analyzed, the 5' end point was at position -306. The B1 promoter mutation is shown in Figure 7A. Microinjection analyses showed that this mutant promoted 45% of the expression of the wild-type -306 promoter [Fig. 7B]. Gel shift analyses using this mutant promoter as competitor showed that although it can compete for binding of the oocyte protein(s) to the wild-type promoter fragment, it does so with a five- to sevenfold reduced efficiency [Fig. 8A].

The B2 box point mutant (Fig. 7A) was transcribed in oocytes with approximately twofold lower efficiency than the -306 promoter [Fig. 7B]. When used as a competitor in gel shift analyses, the B2 mutant promoter was found to compete three- to fivefold less well than the wild-type promoter for the B2 complex [Fig. 8B].

Discussion

The regulation of transcription in eukaryotic cells involves the interaction of protein factors with specific DNA promoter sequences. To understand the role that these protein–DNA interactions play in the modulation of gene expression, it is necessary to determine how the expression of the genes encoding transcription factors is regulated. Because the 5S gene-specific transcription factor TFIIIA is both regulated developmentally and one of the first eukaryotic factors for which the corre-
Figure 6. Detection of a USF-related protein in Xenopus whole ovary extract. (A) A USF oligonucleotide competes specifically for the B1 complex. The standard binding reaction using 30 μg of whole ovary extract and 32P-labeled probe spanning from −306 to −180 is shown. Competitors included in the reaction mixture are increasing concentrations of a USF oligonucleotide (lanes 2-4) or of a nonspecific oligonucleotide containing an octamer (OTF1) binding site (lanes 5-7) [see Materials and methods for oligonucleotide sequences]. (B) Human USF protein binds the TFIIIA promoter sequence. A 32P-labeled probe containing sequences from −425 to −253 and various dilutions of a partially purified USF fraction were used in the binding reaction. The shifted complexes appear as a doublet (arrow).

TFIIIA gene expression

A functional analysis of the region between −306 and −173 has implicated at least three distinct and closely spaced sequence elements in the expression of the TFIIIA gene in X. laevis oocytes. A negative element lies between −306 and −289, whereas a positive element lies directly downstream of the negative element and involves sequences from position −289 to −253. Internal deletions and point mutations in the regions between −250 and −173 and between −144 and −101 reduce transcriptional efficiency, indicating the presence of additional positive-acting elements within these regions. That these latter elements were only detected in analysis of internal deletions or point mutants suggests that upstream promoter regions not present in the 5’ deletion mutants may be required for their activity. We cannot ignore the possibility, however, that effects of internal deletions that have not yet been confirmed by point mutations may result from changes in spacing generated by the deletions. Although removal of the putative CAAT box by 5’ deletion had no appreciable effect on transcription, an internal deletion mutant in which sequences between −93 and −78, including the CAAT sequence, were removed supported 49% of wild-type expression; whether this decrease is due to removal of the CAAT box or an adjacent element is presently under investigation. Removal of sequences between −47 and −11 abolishes transcription from the TFIIIA promoter. Because the TATA box is found within this region, we assume that most if not all of this effect results from its removal. Recently, Matsumoto and Korn (1988) reported the identification of two positive elements within the

TFIIIA alleles exhibit polymorphism, yet are highly conserved throughout the promoter region

Tso et al. (1986) have described previously allelic differences among cloned TFIIIA cDNA and genomic DNA sequences based in part on the presence or absence of a PstI site at position 139 within the protein coding sequence. On the basis of this criterion, the gene described in this report most closely resembles the allele described by Taylor et al. (1986), which lacks a PstI site at position 139. On a gross level, our data further substantiate the proposed allelic heterogeneity, as we have observed differences in at least one additional restriction site between the two genes described. Nucleotide sequence analysis revealed further heterogeneity within the 5’ leader sequence; within 50 bp there are five nucleotide differences between the two alleles.

One negative and three positive cis-acting elements have been defined within the TFIIIA promoter

A functional analysis of the region between −306 and
Two oocyte proteins bind within elements required for the efficient expression of the TFIIIA promoter in oocytes

To define the nature and function of the proteins interacting within the TFIIIA promoter, we have used gel shift assays to identify sequence-specific DNA binding proteins present in X. laevis oocyte extracts. To date, we have resolved two DNA–protein complexes, designated B1 and B2, which involve a promoter fragment containing sequences from −306 to −180. We have shown by competition analysis that the binding of the protein(s) involved in the B1 complex requires sequences between −289 and −252, a region that is shown to be involved in the activation of this gene in oocytes. Similar analysis of the B2 complex has allowed us to map the binding site for this protein between −235 and −221 within the TFIIIA promoter, a region contained within a second activator element located between −250 and −173. Therefore, sequences shown to be required for the in vivo expression of the TFIIIA gene bind proteins, which we assume act as regulators of TFIIIA gene expression.

The protein(s) involved in the formation of the B1 complex is related to the major late transcription factor, USF

It has recently become apparent that, in addition to eukaryotic RNA polymerases (Sentenac 1985), several eukaryotic transcription factors have been highly conserved during evolution (Buratowski et al. 1988). Several mammalian factors have been shown to have homologous counterparts in Xenopus. The Xenopus cytoskeletal actin gene contains a 20-bp homology to the human c-fos serum response element that binds a Xenopus factor homologous to the human serum response factor (Mohun et al. 1987). Doebbeling et al. (1988) have identified a cell-specific activator region within the Xenopus A2 vitellogenin gene that is active transcriptionally in rat liver nuclear extracts. A Xenopus histone H4 transcription factor, XF4TF1, is most likely homologous to the human histone H4 transcription factor H4TF2 (G.H. Thomsen and R.G. Roeder, in prep.). Emerson and Roeder (1984) have shown that Xenopus and human RNA polymerase III factors are interchangeable in functional assays. The relationship we have identified between the Xenopus B1 protein and the human major late transcription factor extends these observations.

The major late transcription factor, USF (Sawadogo and Roeder 1985) or MLTF (Carthew et al. 1985) was identified initially by its ability to bind selectively to and activate transcription from the adenovirus major late promoter. More recently, it has been found to bind to and activate cellular genes, such as the rat γ-fibrinogen promoter (Chodosh et al. 1987) and the mouse metallothionein I gene promoter (Carthew et al. 1987). USF-related proteins also have been reported in yeast (Bram and Kornberg 1987). We have found that the X. laevis oocyte protein involved in the formation of the B1 complex is similar in size and binding properties to the human USF protein, binding to a 6-bp core CACGTG of the 12-bp adenovirus major late sequence; the resulting complex migrates with a similar Rf value as does the complex formed between the Xenopus–DNA fragment and the human USF protein [data not shown]. Recently, it has been demonstrated that polyclonal mouse antibodies raised against human USF (Sawadogo et al. 1988) interact with the B1 complex in a gel shift assay, resulting in a second shift with lower electrophoretic mobility [H. Kaulen, unpubl.]. Whether the Xenopus protein is functionally similar to the human protein awaits purification and in vitro analysis. However, the location of its binding site over 260 bp upstream of the start site of transcription suggests that if its function involves interaction with TFIIID, as has been suggested for the major late USF (Sawadogo and Roeder 1985), bending of the DNA molecule is likely involved in the juxtaposition of these two proteins.
Developmental regulation of the TFIIIA gene—a perspective

In *X. laevis*, TFIIIA gene expression is regulated developmentally, with highest levels occurring during the maturation of the oocyte; the regulation of this gene-specific transcription factor has been implicated in the regulation of the *Xenopus* 5S gene family. Regulation of TFIIIA gene expression may itself be modulated by changes in the interplay of various constitutive and regulatory transcription factors within the promoter elements throughout development. It is interesting to speculate how the closely spaced positive and negative cis-acting elements within the TFIIIA promoter may be occupied differentially by factors during *Xenopus* development. A model to explain the regulation of the TFIIIA gene would involve a decrease in the binding of activators and an increase in the binding of repressor/silencers during development, because of either quantitative or qualitative changes in the transcription factors. This model is similar to the mechanism described for regulation of the β-interferon gene, in which a negative regulatory element in proximity to a constitutive transcriptional element binds a factor that is released on induction (Goodbourn et al. 1986; Zinn and Maniatis 1986). We are continuing our search for trans-acting factors involved in the expression of this gene and will determine whether the modulation of these proteins throughout oogenesis, embryogenesis, and in somatic cells is involved in the regulation of the TFIIIA gene.

Materials and methods

Cloning and characterization of the TFIIIA gene

A *Xenopus* genomic library constructed in EMBL4 using partial Sau3A-digested high-molecular-weight DNA was the kind gift of D. Melton (Harvard University). The library was screened with the TFIIIA cDNA 3a1.b (Ginsberg et al. 1984) using standard techniques (Maniatis et al. 1982). Two clones were analyzed further and the TFIIIA transcription unit contained
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therein was oriented relative to subclones of the 3a1.3 cDNA. Previous primer extension analysis (Ginsberg et al. 1984) allowed us to map the 5' end of the cDNA within a genomic 2.2 kb EcoRI–PstI fragment, which was subcloned into pUC 12 or 13, or M13mp8 or M13mp9, prior to nucleotide sequencing. Unless otherwise noted, all restriction enzymes used for these and subsequent analyses were obtained from either New England Biolabs, Bethesda Research Laboratories, or Boehringer–Mannheim Biochemicals, and reactions were carried out according to suppliers’ specifications.

Construction of 5’ deletion mutant chimeric genes

The pUC 13 plasmid containing the 2.2-kb EcoRI–PstI fragment was linearized by digestion with BgIII. Sequential deletions were generated using the exonuclease Bal31 and conditions recommended by the vendor [IBI]. Following digestion, the ends were repaired with the Klenow fragment of DNA polymerase I and EcoRI linkers were added at the deletion end points. EcoRI–Rsal fragments extending from the 5’ deletion end point through 88 bp of the first TFIIIA exon were sized by gel electrophoresis, isolated, and subcloned into EcoRI–Smal-digested pUC 13. The end points of the 5’ deletion mutants were determined in a sequencing reaction using the chain-termination method and Sequenase reagents (U.S. Biochemical Corp.).

TFIIIA/β-globin chimeric genes were constructed by joining each promoter deletion mutant to a 1689-bp BamHI–PstI fragment derived from the human β-globin gene (Poncz et al. 1983) containing part of the second and all of the third exon of the β-globin gene, including the second intervening sequence and a polyadenylation signal.

Cloning of the internal deletion mutants

The 3’ promoter deletion mutants were constructed following the same general strategy as described above. A pUC 13 plasmid containing 306 bp of the TFIIIA promoter region was digested with BamHI and treated with Bal31 exonuclease [IBI]. BamHI linkers were ligated to the truncated 3’ ends of the TFIIIA promoter. EcoRI–BamHI fragments of varying size were isolated and subcloned into EcoRI–BamHI-digested pUC 13.

Suitable 5’ and 3’ promoter mutants in pUC 13 were chosen for cloning of the internal deletion mutants. The 3’ mutants were digested with BamHI, and the 5’ overhanging ends were filled in with DNA polymerase I Klenow fragment and recut with HindIII, a polynuker restriction site close to the BamHI site. The 5’ deletions were digested with EcoRI, filled in, and redigested with HindIII. The EcoRI–HindIII fragments were gel-purified and ligated to the 3’ mutants prepared as described.

All internal deletion mutants were confirmed by sequence analysis. For IDM 2, IDM 3, and IDM 4, the deletion end points were separated by a 14-bp synthetic linker, 5’-GGGATC-CAATTCTC-3’). IDM 5 contains a 19-bp insertion 5’-GGGGATC-CAATTCTCAGG-3’. 13 bp were provided by a synthetic linker and 6 bp are due to a rearrangement at the target sequence during cloning. IDM 6, IDM 7, IDM 8, and IDM 9 contain a 13-bp synthetic linker sequence 5’-GGGGATC-CAATTCTC-3’. In some cases, the inserted sequence restored the original promoter sequence, which was taken into account when calculating the spacing change.

Site-directed mutagenesis

The point mutants in the B1 and B2 boxes and IDM 1 were constructed using the site-directed mutagenesis protocol of Kunkel et al. (1987), with minor modifications. A TFIIIA promoter fragment including sequences from −306 to +88 was subcloned into the vector pEMBL19, which contains an ori origin of replication and can be used to generate single-stranded templates [Dente et al. 1985]. The recombinant plasmid was transformed into the Escherichia coli dut− ung− strain BW313 (Kunkel 1985). Following superinfection with 11 phage, single-stranded DNA was purified for use in subsequent mutagenesis procedures. The oligonucleotides used to produce the B1 box mutation contained the sequence −280 5’-TCATGTAT-TATTCGACGCTCCACTAGGC-3’ −252; the B2 box mutant oligonucleotide included the sequence −253 5’-ACT-CAAAGCTAAGGAGATCATGCTCAAAAACC-3’ −214 (altered bases are underlined). The internal deletion IDM 1 (−250 to −226) was created using oligonucleotides flanking the deletion site. Sequenase (U.S. Biochemical Corp.) replaced T4 DNA polymerase in the in vitro synthesis reactions. Mutations were verified by nucleotide sequencing.

Microinjection experiments in stage V and VI X. laevis oocytes

Ovaries were obtained from large adult female X. laevis purchased from either Nasco (Fort Atkinson, Wisconsin) or Xenopus (Michigan). Individual stage V and VI oocytes (staged according to Dumont (1972)) were teased from the ovary and stored in MSB-H buffer [88 mM NaCl, 1 mM KCl, 2.4 mM NaHCO3, 0.82 mM MgSO4, 0.33 mM Ca(NO3)2, 0.41 mM CaCl2, 10 mM HEPES (pH 7.6), 10 mg/ml of penicillin, 10 mg/ml of streptomycin] for several hours up to 4 days prior to use. Subsequent microinjections and incubations were performed in MSB-H buffer. For each TFIIIA promoter mutant analyzed, a X. laevis histone H4 gene, into which a 9-bp HindIII linker was inserted for identification [G. Thompsen, Rockefeller University, New York] was coinjected as a control; 400 pg or 2 ng of both the control and mutant plasmid DNA in 10 nl or 40 nl of H2O, respectively, were injected into the germline vesicles of 25–50 oocytes. Injected oocytes were incubated for 20 hr at 23°C, then harvested for further analysis.

Isolation of RNA from microinjected oocytes

Surviving oocytes (75–100%) were collected, pooled, and washed several times in TE buffer [10 mM Tris (pH 8.0), 1 mM EDTA], then homogenized in an extraction buffer containing 10 mM Tris (pH 7.6), 0.5 mM dithiothreitol, 10 mM EDTA, 0.5% SDS, and 0.14 mg/ml Proteinase K and incubated for 1 hr at 37°C. Following incubations, samples were brought to 0.3 M sodium acetate and extracted several times with phenol, phenol/chloroform, and chloroform. The RNA was precipitated, recovered by centrifugation, and resuspended in 10 μl of H2O per oocyte equivalent prior to quantitation. Between 3 and 5 μg of total RNA was recovered per oocyte. RNA integrity was monitored by agarose gel electrophoresis prior to analysis.

Primer extension experiments

Prime extension analyses were performed in primer excess, using 1.5 oocyte equivalents of total RNA for analysis of TFIIIA gene expression (5–7 μg), and 0.5 oocyte equivalents of RNA for analysis of transcription of the control H4 gene (1.5–2.5 μg), and 3 ng (1 × 106 to 2 × 106 cpm) of the appropriate primers. Synthetic oligonucleotides corresponding to a region of exon 2 of the human β-globin gene and spanning a synthetic modification introduced in the histone H4 gene were radioactively labeled with T4 polynucleotide kinase and γ-32P[ATP]. Neither primer was found to cross-hybridize to any endogenous oocyte.
transcript. Extension of the 26-base globin primer on a properly initiated transcript generates a 109-base fragment. The H4 primer is 34 bases and the resulting extension product is 181–187 bases long; this heterogeneity is due to multiple start sites (G. Thompson, pers. comm.). RNA and primer were heated to 80°C for 2 min and incubated for 2 hr at 65°C (globin primer) or 55°C (histone H4 primer) in a hybridization buffer containing 0.4 M NaCl, 40 mM PIPES (pH 6.5), and 1 mM EDTA. Specific RNA was eluted in 50 mM Tris (pH 8.3), 10 mM MgCl₂, 40 mM KCl, 5 mM DTT, 0.2 µg of actinomycin D, 0.2 mM dNTPs, 45 units RNase inhibitor (RNasin, Boehringer–Mannheim), and 5–10 units AMV reverse transcriptase (Boehringer–Mannheim) at 42°C. The reaction was stopped by heating to 70°C for 10 min. The RNA was degraded with DNase I-free RNase A and the elongation product was precipitated, re-suspended in sequencing dye, and electrophoresed through an 8% sequencing gel. Signals were quantitated by scintillation counting.

Preparation of total ovary extract

Oocytes were released from whole ovaries by gentle rotation in a solution of 0.3% collagenase IA (Sigma), prepared in MSB-H, for 12 hr at 23°C. Following extensive washing with MSB-H, the oocytes were washed in TE (pH 8.0), and lysed by 4 strokes of a Kontes all-glass Dounce homogenizer (B-type pestle). The lysis buffer contained 50 mM Tris (pH 8.4), 170 mM KCl, 0.2 mM EDTA, 20% glycerol, 0.5 mM DTT, and 0.5 mM PMSF. The extract was centrifuged at 100,000g at 4°C for 1 hr. Aliquots of the supernatant were quick-frozen and stored at -70°C prior to use. Protein concentrations were determined according to Bradford (1976).

Gel shift assays

Gel shift assays were performed according to Garner and Revzin (1981) with some modifications: 1 ng of a 32P-labeled DNA fragment was incubated with 4 µg of poly[d(1-C)], 30 µg of oocyte protein, and competitor DNA where indicated in a buffer containing 25 mM HEPES (pH 7.9), 10 mM MgCl₂, 1 mM DTT, 0.1 mM CaCl₂, 60 mM KCl, and 10% glycerol at room temperature for 15 min. The reaction mixture was electrophoresed through a 6% polyacrylamide gel in 1 x TBE (89 mM Tris, 89 mM boric acid, 2 mM Na₂EDTA). The five TFIIA promoter oligonucleotides used as competitors contained sequences between -26 to -306, -289 to -252, -271 to -252, -252 to -221 and -220 to -180; each oligonucleotide was flanked with a BamHI restriction site. The USF oligonucleotide used as competitor contained the sequence 5’-CCGAATTCCTGGCATTACACTG-3’. The non-specific oligonucleotide competitor, which included a binding site for the human histone H2B octamer protein, contained the sequence 5’-AATTCACCACCTGGATAATGGCTTCG-3’ (Fletcher et al. 1987). Assays in which the human USF protein was substituted for the oocyte extracts were performed as described by Sawadogo et al. (1988). The human USF protein, kindly provided by M. Sawadogo, was a partially purified fraction following chromatography on Mono Q, as described by Sawadogo et al. (1988).

DNA techniques

All DNA manipulations were done using standard techniques (Maniatis et al. 1982) and according to the National Institutes of Health [NIH] guidelines.

Acknowledgments

We thank Dr. D.A. Melton for providing the X. laevis genomic DNA library, and Dr. A.M. Ginsberg for isolation of the genomic clone. We are grateful to Drs. Kevin C. Gorman and Eckart Bartsch for assistance in the preparation of the figures, to M.C. Gleich for technical assistance, and to Dr. Gerald H. Thomsen and the members of the Roeder laboratory for helpful scientific discussion. This work was supported by Public Health Service grant CA-2567 from the NIH to R.G.R. and by general support from the Pew Charitable Trusts. K.W.S. was supported in part by an American Cancer Society fellowship [PF-2648] and by a National Research Service Award from the NIH (GM-10777). K.K. was supported by a postdoctoral fellowship from the German Research Foundation.

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*Genes Dev.* 1989, 3:
Access the most recent version at doi:10.1101/gad.3.5.651

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