Staf, a promiscuous activator for enhanced transcription by RNA polymerases II and III

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Staf is a zinc finger protein that we recently identified as the transcriptional activator of the RNA polymerase III-transcribed selenocysteine tRNA gene. In this work we demonstrate that enhanced transcription of the majority of vertebrate snRNA and snRNA-type genes, transcribed by RNA polymerases II and III, also requires Staf. DNA binding assays and microinjection of mutant genes into Xenopus oocytes showed the presence of Staf-responsive elements in the genes for human U4C, U6, Y4 and 7SK, Xenopus U1b1, U2, U5 and MRP and mouse U6 RNAs. Using recombinant Staf, we established that it mediates the activating properties of Staf-responsive elements on RNA polymerase II and III snRNA promoters in vivo. Lastly a 19 bp consensus sequence for the Staf binding site, YY(A/T)CCC(A/G)N(A/C)AT(G/C)C(A/C)YY-RCR, was derived by binding site selection. It enabled us to identify 23 other snRNA and snRNA-type genes carrying potential Staf binding sites. Altogether, our results emphasize the prime importance of Staf as a novel activator for enhanced transcription of snRNA and snRNA-type genes.

Keywords: RNA polymerase II/RNA polymerase III/snRNA genes/transcriptional activator/zinc finger

Introduction

Genes for vertebrate small nuclear RNAs (snRNAs) are transcribed by either RNA polymerase II (Pol II) or RNA polymerase III (Pol III), depending on the type of promoters they harbor. The basal promoters of both types include an essential proximal sequence element (PSE) located at approximately ~59 upstream of the transcription start site. The Pol III-dependent genes also possess a TATA box at ~30 which acts as a major determinant of RNA Pol III specificity (Lobo and Hernandez, 1989; Mattaj et al., 1988; see Hernandez, 1992 for a review). A number of other short transcription units, such as the 7SK RNA, Y RNA, MRP RNA and H1 RNA genes have similar basal promoter elements and can be classified as snRNA-type genes. snRNA and snRNA-type genes contain, in addition to the cis elements described above, a distal sequence element (DSE). The DSE plays a major role in transcription efficiency, accounting for a 5- to 100-fold level of activation of Pol II or Pol III basal transcription in transfected cells or injected Xenopus oocytes.

Numerous Pol II and Pol III DSEs have been dissected and found to be composed of an octamer motif and another, usually close, element (for a review see Hernandez, 1992). Among the latter Sp1 binding sites in the human U2, Xenopus U2 and U6 genes (Ares et al., 1987; Janson et al., 1987; Tebb and Mattaj, 1989; Lesecre et al., 1992), an AP-2 binding site and a CRE motif in the human U4C gene (Weller et al., 1988), SPH motifs in the chicken U1 and U4B genes (Roebuck et al., 1990; Zamrod and Stumph, 1990; Cheung et al., 1993), CAAT motifs in human and Xenopus U3 genes (Ach and Weiner, 1991; Savino et al., 1992), a NONOCT motif in the human U6 gene (Danzeiser et al., 1993), a D2 motif in the Xenopus U2 gene (Tebb and Mattaj, 1989) and a CACCC box and octamer-like motifs in the human 7SK gene (Murphy et al., 1989, 1992; Kleinert et al., 1990; Boyd et al., 1995) have been identified.

The Sp1 and octamer motifs contain the recognition sites on the DNA for the well-characterized transcriptional activators Sp1 and Oct-1 respectively (Courey and Tjian, 1988; Sturm et al., 1988; for reviews see Herr, 1992; Hernandez, 1992). However, transcription factors interacting with the other elements described above have not been purified to homogeneity or cloned. Furthermore, owing to the occurrence of octamer or octamer-like sequences in a number of DSEs, it has been tacitly admitted that the activation function of the DSE is mediated essentially by Oct-1 binding at the octamer motif.

The basal promoter of the atypical selenocysteine tRNA gene is principally external to the coding region and comprises a PSE and a TATA motif functionally equivalent to those of vertebrate U6 snRNA genes (Carbon and Krol, 1991; Myslinski et al., 1993a). Additionally, its basal promoter is activated by the activator element (AE), an element which functions without assistance of the octamer (Myslinski et al., 1992, 1993b). Instead, the activation properties of the AE are mediated by Staf, a sequence-specific zinc finger protein that we recently characterized (Schuster et al., 1995). Experimental evidence provided in this work shows that Staf is also involved in transcriptional activation of a large variety of snRNA and snRNA-type genes transcribed by RNA Pol II and Pol III. Our results indicate that AP-2, D2, NONOCT, octamer-like and SPH motifs previously described as being involved in transcriptional activation of a number of these genes are in fact Staf-responsive elements. Staf is thus a key factor for transcriptional activation of snRNA and snRNA-type genes by RNA Pol II and Pol III.

Results

Staf binds specifically to the majority of snRNA and snRNA-type genes

To determine whether Staf is involved in transcriptional activation of snRNA and snRNA-type genes, gel retard-
Table I. snRNA and snRNA-type genes in this study

<table>
<thead>
<tr>
<th>Gene</th>
<th>Abbreviation</th>
<th>Transcribed by</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human U1</td>
<td>hU1</td>
<td>Pol II</td>
<td>Lund and Dalhberg (1984)</td>
</tr>
<tr>
<td>X.laevis U1b1</td>
<td>xU1b1</td>
<td>Pol II</td>
<td>Krol et al. (1985)</td>
</tr>
<tr>
<td>X.laevis U1b2</td>
<td>xU1b2</td>
<td>Pol II</td>
<td>Krol et al. (1985)</td>
</tr>
<tr>
<td>X.laevis U2</td>
<td>xU2</td>
<td>Pol II</td>
<td>Mattaj and Zeller (1983)</td>
</tr>
<tr>
<td>X.laevis U3A</td>
<td>xU3A</td>
<td>Pol II</td>
<td>Savino et al. (1992)</td>
</tr>
<tr>
<td>Human U4C</td>
<td>hU4C</td>
<td>Pol II</td>
<td>Bark et al. (1986)</td>
</tr>
<tr>
<td>X.laevis U5</td>
<td>xU5</td>
<td>Pol II</td>
<td>Kazmaier et al. (1987)</td>
</tr>
<tr>
<td>Human U6</td>
<td>hU6</td>
<td>Pol III</td>
<td>Kunkel et al. (1986)</td>
</tr>
<tr>
<td>Mouse U6</td>
<td>mU6</td>
<td>Pol III</td>
<td>Oshima et al. (1981)</td>
</tr>
<tr>
<td>X.tropicalis U6</td>
<td>xU6</td>
<td>Pol III</td>
<td>Krol et al. (1987)</td>
</tr>
<tr>
<td>Human H1 RNA</td>
<td>hH1 RNA</td>
<td>Pol III</td>
<td>Bae et al. (1990)</td>
</tr>
<tr>
<td>Human Y4</td>
<td>hY4</td>
<td>Pol III</td>
<td>Mariaia et al. (1994)</td>
</tr>
<tr>
<td>Human 7SK</td>
<td>h7SK</td>
<td>Pol III</td>
<td>Murphy et al. (1986)</td>
</tr>
<tr>
<td>X.laevis MRP RNA</td>
<td>xMRP RNA</td>
<td>Pol III</td>
<td>Bennett et al. (1992)</td>
</tr>
</tbody>
</table>

Determination assays were used in the first place to examine the ability of Staf to bind the DSEs arising from 14 genes transcribed by RNA Pol II and Pol III (see Table I). Labeled DNA fragments encompassing the various DSEs (see Materials and methods) were incubated with the purified Staf DNA binding domain and then analyzed on non-denaturing polyacrylamide gels. Figure 1 shows that Staf bound to the majority of the 14 DSEs tested. A high yield of binding was detected with the DSEs of xU1b1 (Figure 1, lane 6), xU2 (lane 12), hU4C (lane 20), xU5 (lane 24), hU6 (lane 28), mU6 (lane 32), hY4 (lane 42), h7SK (lane 46) and xMRP RNA (lane 50). The intensities of the retarded complexes observed in these lanes are comparable with that obtained with the tRNA\textsuperscript{sec} gene (Figure 1, lane 54). In contrast, a very low binding was observed with hU1 (lane 2) and hH1 RNA (lane 38). Lastly, no gel shift at all could be obtained with xU1b2 (lane 10), xU3A (lane 16) and xU6 (lane 36) DSEs. To demonstrate that these retarded complexes were caused by the specific binding of Staf, gel retardation assays were performed in the presence of an excess of two different double-stranded oligodeoxynucleotides acting as competitors. The first contains the AE of the Xenopus laevis tRNA\textsuperscript{sec} gene, which is specifically recognized by Staf, the other carries a mutant AE unable to bind Staf (Schuster et al., 1995). Band shifts were abolished in the presence of the specific competitor (Figure 1, lanes 3, 7, 13, 21, 25, 29, 33, 39, 43, 47, 51 and 55) but unaltered when the mutant AE was used instead (lanes 4, 8, 14, 22, 26, 30, 34, 40, 44, 48, 52 and 56). These results are consistent with a specific binding of Staf to xU1b1, xU2, hU4C, xU5, hU6, mU6, hY4, h7SK and xMRP RNA DSEs.

To localize the Staf binding sites, DNase I footprint analysis was carried out with labeled DNA probes harboring the various DSEs. Those DSEs binding Staf with high yield produced a clear footprint over at least 21 bp (Figure 2A). The protected regions are shown schematically in Figure 2B, together with that obtained on the AE of the tRNA\textsuperscript{sec} gene (Schuster et al., 1995). Sequence comparisons between the various binding sites revealed homologous sequences, on one strand or the other, allowing derivation of a 20 bp consensus sequence for the Staf binding site, YYTCCCANNRTNCCNYYGCRR (Figure 2B).
**snRNA-type promoter transcriptional activation by Staf**

**Fig. 2.** Identification of Staf binding sites in the DSEs of xU1b1, xU2, hU4C, xU5, hU6, mU6, hY4, h7SK and xMRP RNA genes.

(A) Footprint analysis of Staf–DNA complexes. DNase I digestion of the xU1b1, xU2, hU4C, xU5, hU6, mU6, hY4, h7SK and xMRP RNA probes in the absence (lanes 2, 6, 10, 14, 18, 22, 26, 30 and 34) or presence of 3 (lanes 3, 7, 11, 15, 19, 23, 27, 31 and 35) or 4 μl (lanes 4, 8, 12, 16, 20, 24, 28, 32 and 36) of Staf DNA binding domain. G+A chemical cleavage is indicated above the lanes. (B) Sequence comparisons of the different footprinted Staf DNA binding sites. DNase I-protected regions are diagramed above each sequence by a solid bar. For xU2, hU6 and h7SK, the reported protected regions correspond to the footprint obtained on the opposite strand. The sequence protected by Staf in the \textit{X.laevis} tRNASec gene (Schuster et al., 1995) is added at the bottom of the figure and the activator element of this gene (Myslinski et al., 1992) is indicated in white on a dark background. The numbers on the right and left sides indicate the distance from the transcription initiation site. Residues conserved in at least seven of the 10 mapped Staf binding sites are indicated on a shaded background. The 20 bp consensus sequence is derived at the bottom of the figure. N, R and Y stand for any nucleotide, purine and pyrimidine respectively. Pol II and Pol III, genes transcribed by RNA polymerases II and III.

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**Functional relevance of the mapped Staf binding sites**

We next analyzed the functional relevance of the mapped binding sites by: (i) creating substitution mutants either unable or showing severely reduced abilities to bind Staf (Figure 3A); (ii) assaying their transcription abilities by injection into Xenopus oocyte nuclei (Figure 3B). The substitution mutants changed positions 4–7 of the consensus sequence. The conserved CCCA (positions 4–7) in the xU1b1, hU4C, xU5, hU6, mU6, hY4 and xMRP RNA genes was substituted by AAAC. In xU2 and h7SK, CCCG and TCCA (at the same positions) were substituted.
By AAAT and GAAC respectively. In this injection assay, the transcription activities of seven of the nine mutants dropped considerably (Figure 3B). Normalized residual values, expressed relative to the corresponding wild-type levels, ranged between 2% (hU6 and hY4, lanes 9 and 10, and 13 and 14 respectively), 5% (xU1b1, lanes 1 and 2), 15% (xMRP RNA, lanes 17 and 18), 30% (h7SK, lanes 15 and 16) and 40% (hU4C and mU6, lanes 7 and 8, and 11 and 12 respectively).

Transactivating properties of Staf on Pol II and Pol III snRNA promoters

In order to show that Staf is actually responsible for this activation function, we used the X.laevis oocyte expression assay previously developed to establish that Staf mediated transcriptional activation of the tRNA	extsuperscript{sec} gene (Schuster et al., 1995). In this assay, the endogenous Staf background of the oocyte, which would interfere in the experiment, was eliminated by replacing the Staf DNA binding domain with that of Krox-20 (Krox-20 DBD; Figure 4A). The transcription ability of this chimeric protein, termed Staf–Krox-20, was assayed with wild-type Xenopus Pol II U1b2 (Krol et al., 1985) and Xenopus Pol III U6 (Krol et al., 1987) reporters (Figure 4B) and mutant versions thereof lacking the DSE (U1·ΔDSE and U6·ΔDSE) or containing instead the Krox-20 binding site E element (U1·3E and U6·3E). The mRNAs of the effectors Staf–Krox-20 and Krox-20 DBD were transcribed in vitro, capped and injected separately into oocyte cytoplasm (Schuster et al., 1995). After 20 h incubation, the various U1 and U6 reporters were injected into oocyte nuclei, along with [α-32P]GTP. After a second incubation, labeled RNAs were extracted, the levels of which measure the transactivation properties of the protein tested. In the presence of Staf–Krox-20, the transcription levels of U1wt, U1·ΔDSE, U6wt and U6·ΔDSE (Figure 4C, lanes 7, 8, 16 and 17 respectively) were identical to those observed in the absence of effector (lanes 1, 2, 10 and 11 respectively) or in the presence of Krox-20 DBD only (lanes 4, 5, 6 and 14 respectively). Remarkably, however, comparison of lanes 3 with 9 and 12 with 18 revealed that Staf–Krox-20 could significantly stimulate transcription of U1·3E and U6·3E. Of note, transcription levels varied from 10% (U1·3E) to 50% (U1·ΔDSE) and 0% (U6·3E) of the corresponding wild-type promoter level.

Transcription was not mediated by Krox-20 DBD, since transcription of U1·3E and U6·3E was unaffected by its presence (lanes 4, 5, 13 and 14 respectively). Remarkably, however, comparison of lanes 3 with 9 and 12 with 18 revealed that Staf–Krox-20 could significantly stimulate transcription of U1·3E and U6·3E. Of note, transcription levels varied from 10% (U1·ΔDSE) to 50% (U1·3E) and 0% (U6·ΔDSE) to 10% (U6·3E) of the corresponding wild-type promoter level. Transactivation was not mediated by Krox-20 DBD, since transcription of U1·3E and U6·3E was unaffected by its presence (lanes 6 and 15). These results demonstrate unambiguously the transactivating properties of Staf on Pol II and Pol III snRNA promoters.

Selection of DNA binding sites for Staf

To extend our knowledge of the Staf DNA binding sites, we employed the technique of PCR-mediated amplification of protein-selected random oligonucleotides (Blackwell
snRNA-type promoter transcriptional activation by Staf

C) YYRCR (Figure 5). Within the consensus, position 8 is degenerate and positions 4–7, 10, 11 and 13 are more highly constrained than bases at positions 1–3, 8, 9, 12 and 14–18. Positions 9, 12, 14 and 15, considered as fully degenerate in the first consensus derived from sequence comparisons of the different footprints (Figure 2B), in fact match the consensus derived from binding site selection. From the selection data it is obvious that position 20 is fully degenerate and not occupied by R, as deduced from Figure 2B.

Twenty three genes with potential Staf binding sites

Lastly, in addition to the 14 genes tested above, we have used the consensus binding site of Figure 5 to search for the presence of potential Staf binding sites in the other 34 vertebrate snRNA and snRNA-type genes found in the database (Gu and Reddy, 1996). Sequences with a high match (at least 14 out of 19) to the Staf consensus sequence occur in 23 Pol II or Pol III genes (Figure 6), residing between –245 and –185, similarly to the positions for the sites characterized experimentally (Figure 2B). In the light of these findings, we consider that the additional 23 sequences also constitute Staf binding sites. Together with the 10 genes for which we provided experimental evidence, our data strongly suggest that Staf is involved in transcriptional activation of at least 70% of the Pol II and Pol III snRNA and snRNA-type genes available up to now in the databases.

Discussion

Staf is a zinc finger protein that was recently identified as the transcriptional activator of the Pol III selenocysteine tRNA gene (Schuster et al., 1995). In the present work, we have demonstrated that enhanced transcription activity provided by Staf is not devoted to the selenocysteine tRNA promoters in X.laevis oocytes. (A) Schematic organization of the effector mRNAs synthesized in vitro. 5’ UTR and 3’ UTR are the corresponding untranslated regions of the X.laevis β-globin mRNA.

(B) Schematic organization of the Xenopus U1b2 and U6 snRNA reporter genes. PSE and TATA represent the basal promoter elements of the U1 and U6 genes, DSE their distal sequence elements. 3E indicates three E binding sites of the Krox-20 protein substituting for the wild-type DSE. The bent arrow indicates the start of transcription. (C) Enhanced transcription of U1 and U6 snRNAs. Positions of the SS maxi, U1 and U6 are indicated.

et al., 1990; Chittenden et al., 1991; Delwel et al., 1993). To this end, a chimeric protein was used which consisted of glutathione S-transferase fused to residues 257–475 of the Staf DNA binding domain. The fusion protein was purified by affinity binding to glutathione–Sepharose and the Sepharose-bound protein was used for binding and amplification reactions with a 57 bp oligonucleotide duplex that contained a core of 17 random nucleotides. Seventy three clones chosen from the final pool of selected DNAs were sequenced. Of the 22 positions tabulated, 18 positions (1–7 and 9–19) displayed a significantly higher degree of constraint with respect to base preference (Figure 5). Eleven out of 17 display strong secondary preferences (positions 1–3, 7, 9, 12, 14–17 and 19) when the base of first preference is lacking. The 19 bp consensus sequence thus derived is YY(A/T)CCC(A/G)N(A/C)AT(G/C)C(A/ C)YYRCR (Figure 5). Within the consensus, position 8 is degenerate and positions 4–7, 10, 11 and 13 are more highly constrained than bases at positions 1–3, 8, 9, 12 and 14–18. Positions 9, 12, 14 and 15, considered as fully degenerate in the first consensus derived from sequence comparisons of the different footprints (Figure 2B), in fact match the consensus derived from binding site selection. From the selection data it is obvious that position 20 is fully degenerate and not occupied by R, as deduced from Figure 2B.

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Discussion

Staf is a zinc finger protein that was recently identified as the transcriptional activator of the Pol III selenocysteine tRNA gene (Schuster et al., 1995). In the present work, we have demonstrated that enhanced transcription activity provided by Staf is not devoted to the selenocysteine tRNA promoter alone. We have presented several lines of evidence strongly suggesting that Staf is also involved in transcriptional activation of at least 70% of vertebrate snRNA and snRNA-type genes transcribed by RNA Pol II and Pol III. These include the chicken U1 52A and U4B, human U4C, U6, Y4 and 7SK and X.laevis U2 and MRP RNA genes, for which various motifs have been attributed a function by others based on sequences which we have here demonstrated to represent in fact Staf binding sites (Figure 7). In chicken U1 52A and U4B, Staf binding sites match perfectly the SPH motifs previously attributed a function by others (Schuster et al., 1995; Schuster et al., 1990; Schuster et al., 1988; Tebb and Mattaj, 1989; Murphy et al., 1989; Bennett et al., 1992; Danzeiser et al., 1993; Maraia et al., 1994; Boyd et al., 1995). In contrast, our data clearly demonstrate that a Staf-responsive element overlaps these motifs (Figure 7).

The high affinity Staf binding site generated by in vitro selection is a 19 bp consensus sequence which tolerates a high degree of degeneracy in 12 out of 19 positions
Fig. 5. Derivation of the Staf consensus binding site. Analysis of the Staf binding sequences occurring within affinity-selected oligonucleotides. The frequencies with which the four bases A, C, G and T were selected at positions 1–20 (numbering as in Figure 2B) are listed. Positions flanking the 20 bp oligonucleotide were also subjected to selection, labeled –1 at the 5′-end and +1 at the 3′-end. The consensus nucleotide(s) for each position appears below Number of sequences, with lower case letters indicating bases selected less frequently.

(Figure 5). Such a particularly extended binding site may explain the ability of Staf-responsive elements to accept the substantial number of base changes that occur in the different genes tested, without altering the binding of Staf. This is well illustrated by the example of the Staf-responsive elements in the human U6 and Y4 genes, which lack the 3′-part of the consensus Staf binding site (positions RCR in Figure 7) and yet are recognized efficiently by Staf.

In previous reports, we have shown that Staf possesses the capacity to stimulate CAT expression from a Pol II promoter (Myslinski et al., 1992; Schuster et al., 1995). Therefore, our data collectively demonstrate the particular ability of Staf to activate both snRNA-type and mRNA promoters and thus the whole diversity of Pol II and Pol III promoters. Comparison between Staf and its human homolog ZNF 76 revealed the presence, in addition to the central zinc finger domain, of six conserved motifs (Schuster et al., 1995). We hypothesize that some of these conserved motifs represent promoter-selective activation domains directing the differential activation of snRNA and mRNA promoters. This is currently under investigation.

Although the octamer sequence has been recognized for quite some time as a universal motif in the DSEs of vertebrate snRNA and snRNA-type genes, one major finding of our work is the high prevalence of Staf-responsive elements in the DSEs of these genes. About 70% of the DSEs contain both an octamer motif and a Staf binding site associated or not with a third element. The other DSEs contain either octamer or Staf motifs with or without a second element, depending on the DSE. For example, optimal transcription of the Xenopus and human U2 genes is dependent on the three octamer, Staf and Sp1 motifs (Ares et al., 1987; Tebb and Mattaj, 1989; this work). On the other hand, transactivation of the X.laevis selenocysteine tRNA promoter, and probably that of the human Y4 and X.laevis MRP RNA genes, is dependent on a Staf motif only (Myslinski et al., 1992, 1993b; this work). What might be the reason for the variability in the identity and number of motifs constituting the DSE? The answer(s) may reside in the arrangement and strength of the basal promoter elements, which are known to exert a marked effect on motif composition of the DSE and transcriptional activator function (Myslinski et al., 1993b; Das et al., 1995).

The combined presence of the octamer and Staf motifs in a number of genes indicates that enhanced transcription necessitates the simultaneous presence of Oct-1 and Staf transcription factors. These two motifs are always found in close proximity, separated by a maximum of 28 bp. In this regard, we have previously shown that addition of an octamer element in the vicinity of a Staf binding site in the Xenopus Pol II U1b2 and Pol III U6 genes produced a synergistic effect on transcriptional activation, with a marked dependence on the spacing between the two motifs (Myslinski et al., 1993b; our unpublished results). Similar results were obtained with chicken U1 52A and U4B (Roebuck et al., 1990; Zamrod and Stumph, 1990). This suggests a functional cooperativity between the two DNA-bound factors, the basis of which is unknown at the present time. Several possibilities can be invoked: (i) Staf and Oct-1 bind cooperatively to the DNA to activate transcription; (ii) the simultaneous presence of Oct-1 and Staf creates a unique surface for interaction with a co-
snRNA-type promoter transcriptional activation by Staf

Pederson, 1988), mouse U6 (~315–220) (Oshima et al., 1981) and *X.tropicalis* U6 (~335–178) (Krol et al., 1987) were 5′-end-labeled by PCR amplification of the corresponding genes using the proximal 3′-end-labeled primer. Human U1 (positions ~300–134) (Lund and Dahlberg, 1984), *X.laevis* U2 (~310–160) (Matta and Zeller, 1983), human U4C (~257–96) (Bark et al., 1986), *X.laevis* U5 (~260–111) (Kazmaier et al., 1987), human H1 RNA (~279–130) (Baer et al., 1990), human Y4 (~264–101) (Maraya et al., 1994), human 7SK (~243–143) (Murphy et al., 1986), *X.laevis* MRP RNA (~261–100) (Bennett et al., 1992) and *X.laevis* tRNA~Phe~ (~280–102) (Lee et al., 1990) were 5′-end-labeled on the non-coding strand by PCR amplification of the corresponding genes using the distal 3′-end-labeled primer.

**Plasmid constructions**

**Reporter constructs:***U1wt, U1-ΔDSE, U6wt and U6-ΔDSE** correspond to *X.laevis* U1b2 (Krol et al., 1985), *X.laevis* U1b2-ΔDSE (Murgu et al., 1991), *X.tropicalis* U6 (Krol et al., 1987) and C115 genes constructs (Myslinski et al., 1992) respectively. The U1-3E and U6-3E reporters were obtained by ligating in the inverted orientation the BglII fragment of *PvV-3E* (Chavrier et al., 1990) to the BamHI/BglII-cut *X.laevis* U1b2-ΔDSE and C115 constructs respectively. The E sites map at positions ~205–196, ~235–226, ~265–256 in U1-3E and ~219–210, ~249–240, ~270–270 in U6-3E.

**Effectors constructions.** Construction of pBR3-Staf/Krox-20 and pBR3/Krox-20 DBD was as described in Schuster et al. (1995).

**Oocyte microinjections**

In the experiments shown in Figure 3B, *X.laevis* oocytes were co-injected with 4 ng wild-type or mutant templates, 0.2 μCi [α-32P]GTP (800 Ci/mmol) and the 5S RNA maxigene (25 pg for Pol II genes and 100 pg for Pol III genes) as an internal control for oocyte injection and RNA recovery, except for hU4C, where the RNA~Q~ (100 pg) was used instead. For competition experiments, oocyte nuclei were co-injected with 8 ng each template and 25 pg 5S RNA maxigene. Oocytes were incubated at 19°C for 5 (Pol III transcription) or 16 h (Pol II transcription). RNAs were extracted from batches of 10 oocytes and analyzed as described in Schuster et al. (1995). Transcription efficiencies were quantitated with a Fuji Bioimage Analyzer Bas 2000 and normalized relatively to 5S RNA maxigene and RNA~Q~ transcription levels.

In the experiments shown in Figure 4, capped mRNAs (20 nl, 10 ng) were injected into the cytoplasm 20 h before nuclear injection of 20 nl containing the reporter DNA (50 μg/ml), the 5S maxigene (5 μg/ml) as an internal control and [α-32P]GTP (800 Ci/mmol, 0.2 μCi/oocyte). Incubation was for 16 h (U1 reporters) or 5 h (U6 reporters). Transcription of the reporter genes was analyzed as described in Schuster et al. (1995).

**Binding site selection**

The 57 bp oligonucleotide used in the binding selection, 5′-CTGAG-TCTGAGATCCCCTG(N)17AGGCTCAAAGCTGAATTCC-3′, contained an internal region of 17 degenerate bp flank on each side by a 20 bp sequence containing BamHI (S′) and EcoRI (3′) restriction sites. For PCR amplification, the oligonucleotides 5′-CTGATCTGAGATTCCCTG-3′ and 5′-AGGAATTCGACCCTGCTTACC-3′ served as forward and reverse primers respectively. Selection was performed essentially as described in Delwel et al. (1993). After six rounds of binding and amplification by PCR, an additional step was performed to ensure that the majority of the amplified 57 bp oligonucleotides represented perfect duplexes lacking mismatches (Chittenden et al., 1991). To do this, 200 pmol of each primer were added to the reaction and the mixture subjected to an additional PCR cycle. The final oligonucleotide amplification product was purified, BamHI/EcoRI digested and ligated to pBS (+). Isolated clones were sequenced by standard methods (Sambrook et al., 1989). In binding site comparisons, to avoid biasing the data, nucleotides recognized by the PCR primers within the defined sequence were excluded.

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