In eukaryotes RNA polymerase II (Pol II) is responsible for transcribing nuclear genes encoding the mRNAs and several small nuclear RNAs. Like RNA Pol I and Pol III, Pol II cannot recognize its target promoter directly and initiate transcription without accessory proteins. Instead, this large multisubunit enzyme relies on both general transcription factors (GTFs) and transcriptional activators and coactivators (both positive and negative) to regulate transcription from class-II nuclear gene promoters [for review, see Roeder 1996]. The primary DNA anchor of this complicated macromolecular machine is transcription factor IID (TFIID), a 700-kD complex composed of the TATA box-binding protein (TBP) and a set of phylogenetically conserved, Pol II-specific TBP-associated factors [for review, see Burley and Roeder 1996]. DNA binding by human TFIID was first demonstrated with the adenovirus major-late promoter (AdMLP). DNase I footprinting studies of the AdMLP and selected human gene promoters revealed sequence-specific interactions with the TATA element, which are primarily mediated by TBP. Protection outside of the TATA box displays a nucleosome-like pattern of DNase I hypersensitivity, varies radically among promoters, and can be induced by some activators [for review, see Burley and Roeder 1996].

Genes encoding TBPs have been cloned from organisms ranging from archaea to human. The molecules share a phylogenetically conserved 180-residue carboxy-terminal or core segment, which contains two imperfect direct repeats and supports all of the protein’s biochemically important functions in Pol II transcription [for review, see Burley and Roeder 1996]. Because of the original purification and characterization of Saccharomyces cerevisiae TBP, there has been considerable progress toward understanding its mechanisms of action. Specific nanomolar-affinity [Hahn et al. 1989] binding to a TATA element entails DNA bending [Horikoshi et al. 1992] and occurs exclusively via minor groove interactions [Lee et al. 1991; Starr and Hawley 1991]. Three-dimensional structures of a full-length TBP from Arabidopsis thaliana [Nikolov et al. 1992; Nikolov and Burley 1994], yeast core TBP [Chasman et al. 1993], and full-length TBP from the archaeon Pyrococcus woesei [DeDecker et
al. 1996) have been determined. The monomeric protein consists of two nearly identical domains and adopts a quasisymmetric α/β structure, resembling a molecular saddle complete with stirrups (Fig. 1). The concave underside of the saddle is a highly curved, 10-stranded, antiparallel β-sheet, containing the amino acids involved in DNA binding (Fig. 2). The convex upper surface of the saddle consists of four α-helices, which interact with other transcription factors (for review, see Nikolov and Burley 1994). Cocrystal structure determinations of *A. thaliana*, yeast core, and human core TBPs interacting with similar TATA elements (Kim et al. 1993a,b, Kim and Burley 1994; Juo et al. 1996; Nikolov et al. 1996) revealed an unusual protein–DNA complex, characterized by extensive hydrophobic interactions with the minor groove, severely distorted DNA, and phenylalanine side chains kinking DNA by insertion between base pairs at the 5′ and 3′ ends of the TATA box. The same conformational change has been observed in triple-complex cocrystal structures of TBP plus DNA with human TFIIIB [Nikolov et al. 1995], an archaeal homolog of TFIIIB [Kosa et al. 1997], and yeast TFIIA [Geiger et al. 1996; Tan et al. 1996]. Further relevant work on TBP includes examination of the kinetics and thermodynamics of TATA element binding [Hoopes et al. 1992; Petri et al. 1995, 1998; Parkhurst et al. 1996], and studies of the effects of prebending promoter DNA (Parvin et al. 1995) and TBP-induced DNA deformation [Sun and Hurley 1995; L. Hurley, unpubl.).

Computational studies of the eukaryotic promoter database (EPD) have also yielded important insights into the function of TBP (see Table 1 for a summary of results available on the internet via http://www.epd.isb-sib.ch/promoter_elements). An exhaustive statistical survey documented that the TATA box is an A/T-rich 8-bp segment often flanked by G/C-rich sequences (Bucher 1990). Despite the marked preference for A:T and T:A base pairs, C:G and/or G:C base pairs occur frequently at five of the eight positions. Thus, TBP can bind productively to a large number of diverse TATA elements, some of which bear little resemblance to the optimal TATA sequence (5′-TATATAAG-3′), identified by an in vitro binding-site selection experiment with *Acanthamoeba* TBP [Wong and Bateman 1994].

In an effort to understand how TBP can support Pol II transcription initiation from many different TATA boxes in class-II nuclear gene promoters, we have made a systematic X-ray crystallographic study of a canonical wild-type TBP (*A. thaliana* TBP isoform 2) recognizing 10 naturally occurring variants of the AdMLP TATA element. The structure of the TBP–DNA complex is essentially independent of TATA element sequence. Three distinct molecular mechanisms allow TBP to induce the same DNA deformation, one of which uses Hoogsteen base pairs adjacent to the 3′ kink site. Detailed analyses of these high-resolution cocrystal structures document that TATA element recognition has remained constant over the course of evolution.

Results and Discussion

**Study design and functional characterization of TATA element variants**

Table 1 illustrates the oligonucleotides used for crystallization with wild-type TBP. The 10 sequences represent naturally occurring single-base variants of the AdMLP TATA element (5′′-TATATAAAAG-3′′), including at least one substitution at each position. Diffraction-quality cocrystals were obtained with C:G or G:C base pairs at four of the five TATA box positions, where they occur with non-zero estimated frequency in the EPD (ranging from 2.9% to 38.4%). Defining the intrinsic strength of the AdMLP TATA box to be 100% (see Materials and Methods), the variants show transcription activity levels ranging between 5% and 110%. (Similar cocrystalliza-
tion trials were unsuccessful with oligonucleotides that do not support transcription initiation or TBP binding, including 5'-TGTAAAAG-3', 5'-TCTAAAAG-3', 5'-TA-
AAAAG-3', 5'-TATGAAG-3', and 5'-TATACAG-3'.) Table 2 provides a summary of the crystallographic statistics, showing that all 10 newly determined cocrystal structures are of the highest quality. The presence of two protein–DNA complexes/asymmetric units in each case permits an estimate of the precision of each set of atomic coordinates. Root-mean-square deviations (rmsds) between α-carbons range between 0.2 and 0.4 Å, which is comparable to the precision of the X-ray method at resolution limits between 1.8 and 2.3 Å (Brünger and Rice 1997).

**TBP structure is not affected by TATA element sequence variation**

Figure 1B illustrates the structures of TBP extracted from each cocrystal structure and overlaid by least-squares superposition on our original AdMLP cocrystal structure (Kim et al. 1993a; Kim and Burley 1994). With the exception of the short segment connecting α-helix H1 to β-strand S2, TBP structural variation as a function of

---

**Figure 2.** TBP sequence alignments. Sequences of 21 TBPs and *D. melanogaster* TBP-related factor aligned by use of the three-dimensional structure of *A. thaliana* TBP2. (A1) *A. thaliana* TBP isoform 1; (A2) *A. thaliana* TBP isoform 2; (M1) maize TBP isoform 1; (M2) maize TBP isoform 2; (PO) potato TBP; (W1) wheat TBP isoform 1; (W2) wheat TBP isoform 2; (SP) S. pombe TBP; (SC) *S. cerevisiae* TBP; (DM) *D. melanogaster* TBP; (HS) human TBP; (XL) *Xenopus laevis* TBP; (DD) *Dictyostelium discoideum* TBP; (AC) Acanthamoeba castellanii TBP; (OV) Onchocerca volvulus TBP; (CE) Caenorhabditis elegans TBP; (TT) *Tetrahymena thermophila* TBP; (SS) *Sulfolobus shibatae* TBP; (PW) *P. woesei* TBP; (TC) *T. celer* TBP; (PF) *P. falciparum* TBP; (TRF) *D. melanogaster* TRF. The secondary structure is denoted with arrows (β-strands) and solid bars (α-helices). Scheme for identifying protein–DNA contacts is as follows: (r) ribose; (p) phosphate; (w) water-mediated backbone; (b) base; (i) intercalating phenylalanine. Color coding of residues interacting with bases is identical to that used in Fig. 4.
TATA element sequence is comparable to the precision of the atomic coordinates (α-carbon rmsds = 0.2–0.5 Å).

All subsequent structural comparisons are based on the superpositions in Figure 1B. The phenylalanine pairs (Phe-148 and Phe-165, Phe-57, and Phe-74) responsible for kinking the TATA element at its 5' and 3' ends also show no significant structural variation (Fig. 1C,D).

Trajectory of the DNA double-helix axis is not affected by TATA element sequence variation

Figure 3 illustrates only minimal variation in the trajectory of the DNA double-helix axis as a function of TATA element sequence. This similarity documents that TBP binding induces the same deformation in the core promoter, independent of the precise sequence and transcription activity of the TATA element.

Detailed analyses of the DNA structural parameters (Lavery and Sklenar 1989; Stofer and Lavery 1993) for each TATA element (data not shown) reveal only minor differences when compared with the AdMLP TATA box bound to the same protein (Kim and Burley 1994).

Table 1. TATA element variants

<table>
<thead>
<tr>
<th>Oligonucleotide</th>
<th>cUAS (%)</th>
<th>MEL1 (%)</th>
<th>Observed frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>GC T A TAAA A GGGCA CGA T A TTTT T CCCGT</td>
<td>100</td>
<td>100</td>
<td>A/C/G/T</td>
</tr>
<tr>
<td>A(−31) A ATAAAAAG</td>
<td>16</td>
<td>11</td>
<td>5.0/11.0/4.5/79.5</td>
</tr>
<tr>
<td>T(−30) T TAAAAAG</td>
<td>29</td>
<td>20</td>
<td>83.5/1.3/1.3/13.9</td>
</tr>
<tr>
<td>C(−29) T ACAAAAAG</td>
<td>24</td>
<td>15</td>
<td>4.4/3.3/0.9/91.4</td>
</tr>
<tr>
<td>T(−28) T ATTAAAG</td>
<td>N.D.</td>
<td>14</td>
<td>89.2/0.8/1.7/8.4</td>
</tr>
<tr>
<td>T(−27) T ATATAAG</td>
<td>43</td>
<td>25</td>
<td>71.0/0.8/0.5/27.7</td>
</tr>
<tr>
<td>T(−26) T ATATAAG</td>
<td>6</td>
<td>6</td>
<td>84.8/2.9/9.5/2.8</td>
</tr>
<tr>
<td>G(−26) T ATAAAG</td>
<td>N.D.</td>
<td>18</td>
<td>84.8/2.9/9.5/2.8</td>
</tr>
<tr>
<td>T(−25) T ATAAAAAG</td>
<td>110</td>
<td>90</td>
<td>45.0/3.4/16.4/35.2</td>
</tr>
<tr>
<td>C(−25) T ATAAAACG</td>
<td>5</td>
<td>6</td>
<td>45.0/3.4/16.4/35.2</td>
</tr>
<tr>
<td>T(−24) T ATAAAAAG</td>
<td>N.D.</td>
<td>N.D.</td>
<td>35.8/14.0/38.4/11.8</td>
</tr>
</tbody>
</table>

Crystallization oligonucleotide and two assays of transcriptional efficiency [(N.D.) not done] are given for each cocrystal structure of wild-type TBP bound to a variant of the AdMLP TATA element. Eukaryotic Promoter Database estimated frequency statistics are given for each of the eight positions in the TATA element (http://www.epd.isb-sib.ch/promoter_elements). The estimated frequencies corresponding to the AdMLP and the TATA variant base pairs are denoted with boldface type and underlining, respectively. Values of ~1% or less are incompatible with the structural data and are thought to reflect noise in the computational process used to identify and align TATA elements in the EPD (P. Bucher, pers. comm.).

TATA element sequence is comparable to the precision of the atomic coordinates (α-carbon rmsds = 0.2–0.5 Å). All subsequent structural comparisons are based on the superpositions in Figure 1B. The phenylalanine pairs (Phe-148 and Phe-165, Phe-57, and Phe-74) responsible for kinking the TATA element at its 5' and 3' ends also show no significant structural variation (Fig. 1C,D).

Trajectory of the DNA double-helix axis is not affected by TATA element sequence variation

Figure 3 illustrates only minimal variation in the trajectory of the DNA double-helix axis as a function of TATA element sequence. This similarity documents that TBP binding induces the same deformation in the core promoter, independent of the precise sequence and transcription activity of the TATA element. Detailed analyses of the DNA structural parameters (Lavery and Sklenar 1989; Stofer and Lavery 1993) for each TATA element (data not shown) reveal only minor differences when compared with the AdMLP TATA box bound to the same protein (Kim and Burley 1994).

Structure of the TBP–DNA complex is not affected by TATA element sequence variation

Pairwise comparisons of our TBP–DNA cocrystal structures document that even the detailed structure of the protein–DNA complex is not significantly affected by variations in the TATA element (rmsds for core α-carbon and TATA element C1’ atoms = 0.2–0.5 Å). A comparison of plant and archaeal TBP–DNA complexes reveals a very similar protein–DNA interface (rmsds for TATA element C1’ atoms = 1.1 Å), despite the fact that the upper surfaces of the two molecular saddles are somewhat different [data not shown]. We conclude that the structure of the TBP–TATA element complex is independent of the sequence of the TATA element.

Figure 4 provides a schematic view of the protein surface responsible for TATA element recognition. In all available crystal structures containing TBP and DNA, only 15 highly conserved residues contribute to interactions with the minor-groove edges of the bases. Excluding Plasmodium falciparum TBP and Drosophila melanogaster TBP-related factor (TRF) from the sequence comparison, these 15 residues are almost absolutely invariant (Fig. 2). The only exceptions are a Thr-82 → Leu substitution in Schizosaccharomyces pombe TBP, and Thr-173 → Ser substitutions in P. woesei and Thermococcus celer TBPs. The sequence of P. falciparum TBP displays six conservative substitutions, including Val-29 → Thr, Phe-57 → Ile, Val-80 → Met [which is an Ile in Drosophila TRF], Val-119 → Ile, Pro-149 → Ala, and Val-171 → Ile. The residues responsible for direct and water-mediated interactions with the DNA back-
bone are also highly conserved, showing minimal variation particularly in the carboxy-terminal half of the protein [Fig. 2]. Thus, it is not surprising that the three-dimensional structure of TBP and its interactions with DNA have remained unchanged throughout evolution.

Three mechanisms explain how TBP exploits the same induced fit strategy to recognize all 10 variants of the AdMLP TATA element

The major-groove face of B-DNA is varied chemically as a function of sequence, which explains why most sequence-specific DNA-binding proteins interact with the major-groove edges of base pairs in their recognition elements (for review, see Patikoglou and Burley 1997). In contrast, the minor-groove face is chemically monotonous as a function of sequence [Seeman et al. 1976]. T → A and A → T transversions leave the structure of the minor-groove face essentially unchanged, because both A:T and T:A display similarly positioned pairs of hydrogen bond acceptors on their minor-groove edges. G:C and C:G, on the other hand, provide some relative chemical variation. Minor-groove differences between G:C and A:T or C:G and T:A arise from an exocyclic NH$_2$ protruding from G. It is not surprising that DNA-binding proteins exhibiting little or no sequence specificity interact primarily with the minor groove (for review, see Patikoglou and Burley 1997). TBP exploits an induced-fit mechanism that relies, in part, on the chemical monotony of the minor groove to recognize many different core promoters during Pol II transcription initiation.

Minor-groove structural degeneracy of A:T and T:A permits TBP-DNA complex formation on many A/T-rich promoter sequences, albeit with dramatically reduced transcriptional efficiency in some cases (Table 1). Figure 5 depicts the consequences of changing the first base pair of the TATA element from T:A to A:T in our A(−31) cocrystal structure of an AATA box (5 base pair of the TATA element from T:A to A:T in our A(−31) cocrystal structure of an AATA box (5

Table 2. Summary of crystallographic statistics

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Resolution [Å]</td>
<td>6–1.9</td>
<td>6–2.3</td>
<td>6–1.8</td>
<td>6–1.9</td>
<td>6–2.1</td>
<td>6–2.3</td>
<td>6–2.1</td>
<td>6–1.93</td>
<td>6–2.23</td>
<td>6–1.95</td>
<td>6–1.86</td>
</tr>
<tr>
<td>Rmerge [I]</td>
<td>0.052</td>
<td>0.053</td>
<td>0.051</td>
<td>0.037</td>
<td>0.045</td>
<td>0.058</td>
<td>0.060</td>
<td>0.050</td>
<td>0.072</td>
<td>0.054</td>
<td>0.042</td>
</tr>
<tr>
<td>Completeness (%)</td>
<td>93.2</td>
<td>90.9</td>
<td>98.1</td>
<td>86.8</td>
<td>97.0</td>
<td>96.2</td>
<td>89.5</td>
<td>98.7</td>
<td>98.5</td>
<td>99.5</td>
<td>96.5</td>
</tr>
<tr>
<td>Refined statistics</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Resolution [Å]</td>
<td>6–1.9</td>
<td>6–2.3</td>
<td>6–1.8</td>
<td>6–1.9</td>
<td>6–2.1</td>
<td>6–2.3</td>
<td>6–2.1</td>
<td>6–1.93</td>
<td>6–2.23</td>
<td>6–1.95</td>
<td>6–1.86</td>
</tr>
<tr>
<td>Rmerge [I]</td>
<td>0.197</td>
<td>0.182</td>
<td>0.193</td>
<td>0.209</td>
<td>0.194</td>
<td>0.194</td>
<td>0.193</td>
<td>0.196</td>
<td>0.191</td>
<td>0.199</td>
<td>0.210</td>
</tr>
<tr>
<td>Rfree [I]</td>
<td>0.265</td>
<td>0.257</td>
<td>0.239</td>
<td>0.275</td>
<td>0.266</td>
<td>0.263</td>
<td>0.251</td>
<td>0.252</td>
<td>0.267</td>
<td>0.255</td>
<td>0.264</td>
</tr>
<tr>
<td>ΔBonds (Å)</td>
<td>0.013</td>
<td>0.009</td>
<td>0.008</td>
<td>0.011</td>
<td>0.009</td>
<td>0.009</td>
<td>0.009</td>
<td>0.008</td>
<td>0.009</td>
<td>0.009</td>
<td>0.010</td>
</tr>
<tr>
<td>ΔAngles (°)</td>
<td>1.7</td>
<td>1.5</td>
<td>1.4</td>
<td>1.6</td>
<td>1.5</td>
<td>1.4</td>
<td>1.4</td>
<td>1.5</td>
<td>1.5</td>
<td>1.5</td>
<td></td>
</tr>
<tr>
<td>Δβ-factors (Å$^2$)</td>
<td>1.8</td>
<td>2.0</td>
<td>1.9</td>
<td>2.0</td>
<td>1.9</td>
<td>2.0</td>
<td>1.9</td>
<td>2.0</td>
<td>1.9</td>
<td>1.9</td>
<td>1.9</td>
</tr>
<tr>
<td>No. of water molecules</td>
<td>520</td>
<td>269</td>
<td>478</td>
<td>350</td>
<td>350</td>
<td>263</td>
<td>378</td>
<td>502</td>
<td>329</td>
<td>472</td>
<td>439</td>
</tr>
<tr>
<td>(B) protein (Å$^2$)</td>
<td>26</td>
<td>26</td>
<td>25</td>
<td>31</td>
<td>28</td>
<td>29</td>
<td>26</td>
<td>26</td>
<td>30</td>
<td>25</td>
<td>26</td>
</tr>
<tr>
<td>(B) DNA (Å$^2$)</td>
<td>31</td>
<td>31</td>
<td>28</td>
<td>36</td>
<td>31</td>
<td>36</td>
<td>32</td>
<td>31</td>
<td>36</td>
<td>30</td>
<td>32</td>
</tr>
<tr>
<td>(B) water (Å$^2$)</td>
<td>40</td>
<td>39</td>
<td>39</td>
<td>44</td>
<td>40</td>
<td>41</td>
<td>39</td>
<td>41</td>
<td>42</td>
<td>37</td>
<td>40</td>
</tr>
<tr>
<td>PDB code</td>
<td>1QNE</td>
<td>1QNC</td>
<td>1QNA</td>
<td>1QNA</td>
<td>1QNA</td>
<td>1QNA</td>
<td>1QNA</td>
<td>1QNA</td>
<td>1QNA</td>
<td>1QNA</td>
<td>1QNA</td>
</tr>
</tbody>
</table>

Rmerge [I] = $\Sigma I - $ (I) | $\Sigma I$, where $I$ = observed intensity and (I) = average intensity obtained from multiple observations of symmetry related reflections. ΔBonds and ΔAngles are the respective rmsds from ideal values. Δβ-factors is the rmsd between the $B$ values of covalently bonded atomic pairs. Free R-factor was calculated with 10% of the data omitted from the structure refinement. Source/detector denotes beamline (X25 at National Synchrotron Light Source, A1 and F1 at Cornell High Energy Synchrotron Source) and detector ([IP] image plate, [CCD] charge-coupled device).
structures only.

Fig. 1. (A) Color coding of each double-helix axis corresponds to those used calculated by use of CURVES (Stofer and Lavery 1993). The trajectory of the DNA double-helix axis for each cocrystal structure, representation of TBP with a line representation of the traje-

cctory of the DNA double-helix axis for each cocrystal structure, calculated by use of CURVES (Stofer and Lavery 1993). The color coding of each double-helix axis corresponds to those used in Fig. 1. [A] Viewed along the twofold quasisymmetry axis; [B] viewed through the stirrups of the molecular saddle. Overlay was performed by least-squares superposition of the protein structures only.

The second mechanism underlying relaxed DNA-binding specificity allows TBP to accommodate the exocyclic NH$_2$ of G in the hypothetical TAGA box, which almost never appears in the EPD (frequency <1%). The same constraints may not apply to a hypothetical TATAAAC box (frequency = 2.9%), because the cavity at position 6 is somewhat elongated and may receive the protruding amino group from the bottom strand of the promoter. Further analyses of all available TBP–TATA element interfaces do not reveal the existence of any other sizable cavities [data not shown], and we believe that the cavity mechanism of specificity broadening is restricted to positions 3 and 6.

The final contributor to relaxed DNA-binding specificity involves TBP-induced formation of Hoogsteen base pairs (Hoogsteen 1963), which have not been described previously in protein–DNA complexes. Inspection of our AdMLP cocrystal structure suggests that TBP should not tolerate C:G or G:C base pairs at position 7, because the exocyclic NH$_2$ would clash with the side chain of Leu-72 (see the structure of TBP bound to 5' TATAAATG-3', illustrated in Fig. 7C). Bucher’s studies of the EPD, however, yielded non-zero estimates for the frequencies of C:G and G:C base pairs at position 7 [3.4% and 16.4%, respectively]. Cocrystallization of TBP with the oligonucleotide corresponding to a TATAAAC box [C(−25) 5'-TATAAACG-3', frequency = 3.4%, activity = 5%–6%] allowed us to uncover a remarkable explanation for broadened DNA-binding specificity at position 7. A 180° torsion angle change about the C1’–N9 bond [Fig. 8], giving a syn instead of the normal anti conformation, creates a C:G Hoogsteen base pair that is stabilized by interstrand hydrogen bonding [C N4–G O6 = 2.9 Å and possibly C N3–G N7 = 3.0 Å] plus an intrastrand hydrogen bond with the backbone (G N2–phosphate O = 2.9 Å). This additional DNA deformation prevents a steric clash between the exocyclic NH$_2$ of G and Leu-72 that would be produced by the corresponding Watson–Crick base pair [data not shown]. At the same time, the Hoogsteen base pair preserves many of the van der Waals interactions with Phe-57 and Phe-74, which are largely responsible for the DNA kink at the 3’ end of the TATA box. Similar steric arguments apply to the problem of accommodating a G:C base pair at this position, and model building suggests that TATA elements bearing a G at position 7 exploit an analogous G:C Hoogsteen base pair [data not shown]. The corresponding TATAAAG box occurs often in eukaryotic promoters [frequency = 16.4%] and is capable of supporting Pol II transcription initiation both in vitro and in vivo (Wobbe and Struhl 1990).

We believe that Hoogsteen base pairs can form at position 7 because the energy barrier for the anti to syn glycosidic torsion angle change is very low for unstacked DNA bases [Ornstein et al. 1978], which is precisely the case in the vicinity of the 3’ kink. Steric clashes between TBP and Watson–Crick C:G or G:C base pairs at position 7 preclude assembly of stable protein–DNA complexes in the absence of further conformational changes in the nucleic acid. TBP has solved this problem by exploiting the phenylalanine-induced kink between positions 7 and
8 to allow rotation of the position 7 G base about the C1'–N9 bond. Noncrystallographic symmetry allows us to rule out some lattice packing artifact in C(−25), because we see the same behavior in both crystallographically independent TBP–TATAAAC box cocrystal structures comprising the asymmetric unit.

Figure 4. Complementary protein and DNA-binding surfaces. (A) MOLSCRIPT drawing of the DNA-binding surface of TBP viewed along the twofold quasisymmetry axis. Amino acid side chains making base contacts are drawn as stick figures. Atom color codes: yellow, carbon; blue, nitrogen; red, oxygen. Side chain color codes: red, Pro149; green, intercalating phenylalanines Phe-148, Phe-165, Phe-57, Phe-74; magenta, Leu-163, Leu-72; dark blue, Val-171, Val-119, Val-80, Val-29, light blue, Thr-173, Thr-82; yellow, Asn-27, Asn-117. Color coding of residues interacting with bases is identical to that used in Fig. 2. (B) Schematic drawing of the minor-groove surface of the crystallization oligonucleotide showing all direct and water-mediated contacts. The location of the twofold quasisymmetry axis is denoted by a solid, vertical oval. A and B have been prepared as if the image of the underside of TBP was on one page of an open book facing an image of the minor groove of the AdMLP TATA element.

Figure 5. A(−31), T(−30), and C(−29) TATA variants bound to wild-type TBP. Overlays of the structure of TBP bound to the AdMLP (green) and variant TATA elements (yellow), calculated with least-squares superposition of the protein structures. Each composite view shows only the base pair differing between the two crystallization oligonucleotides, surrounding protein residues and bridging water molecules. Protein and DNA are illustrated as atomic stick figures, with color coding for atom type. (C) Yellow; (O) red; (N) blue; (P) pink. Prime denotes bases of the noncoding strand (color coded red in Fig. 1A). Electron density difference \(|F_{\text{observed}}| - |F_{\text{calculated}}|\) maps were calculated with simulated annealing (maximum temperature = 1000 K) after omitting the illustrated base pair from the phase calculation and fixing surrounding atoms with a 3 Å shell. This procedure eliminates model bias, yielding experimental electron density for the structural differences observed between AdMLP and the variant TATA element. The same image format and color-coding scheme is used for Figs. 6–8. (A) A(−31) variant [5'-AATAAAAAG-3', frequency = 5.0%, activity = 11%–16%]. Substitution of A:T for T:A yields similar contacts between Pro-149 and the minor-groove edge of the first base pair. Broken lines denote van der Waals contacts between Pro-149 and O2 of T(−31) (yellow) and N3 of A(−31) (green) with distances of 3.1 and 3.3 Å, respectively. (B) T(−30) variant [5'-TTTAAAAG-3', frequency = 13.9%, activity = 20%–29%]. The C2 atoms of A(−30) (yellow) and A(−30) (green) are similarly close to Leu-163 (3.7 and 3.6 Å, respectively). (C) C(−29) variant [5'-TACAAAG-3', frequency = 3.3%, activity = 15%–24%]. Exocyclic NH2 of G(−29) is accommodated in a pocket, formed by the side chains of Val-119, Val-171, Thr-173, and Leu-163. The closest G(−29) NH2 contact occurs with Val-171 (3.6 Å). Inspection of our original AdMLP cocrystal structure reveals a cavity (denoted by a meshwork polygon) between A(29') and the residues forming the exocyclic NH2 pocket. (D) Simulated annealing electron density map corresponding to the view illustrated in C, contoured at 2.5σ. The map shows clear density for the exocyclic NH2 of G(−29').
Although Hoogsteen base pairs have not been described in protein–DNA complexes, there is extensive literature on Hoogsteen base pairing in drug–DNA complexes (for review, see Chen and Patel 1995). C:G Hoogsteen base pairs were detected in a cocrystal structure of Triostin A bound to a self-complementary duplex oligonucleotide with sequence 5'–GCGTACGC–3' (Wang et al. 1986). Like our observation at position 7 of the TATA sequence, this substitution yields subtle changes in the structure of the protein–DNA interface, permitting entry of a water molecule (yellow sphere) bridging between the minor groove edge of the A and a threonine side chain [A(−26) N3–H2O = 2.9 Å, H2O–Thr-82 Oγ1 = 2.6 Å]. (D) Simulated annealing electron density map corresponding to the view illustrated in C, contoured at 2.5σ. The map shows clear electron density for the variant T:A base pair and interfacial water molecule.

Figure 7. G(−26), T(−25), and T(−24) TATA variants bound to wild-type TBP. (A) G(−26) variant [5’–TATAAGAG–3’, frequency = 9.5%, activity = 18%]. Exocyclic NH2 of G(−26) is accommodated in a pocket, formed by the side chains of Val-29, Val-80, Thr-82, and Leu-72 (omitted for clarity). The closest G(−26) NH2 contact occurs with Val-80 (3.7 Å). The map shows clear density for the exocyclic NH2 pocket. (B) Simulated annealing electron density map corresponding to the view illustrated in A, contoured at 2.5σ. The map shows clear density for the exocyclic NH2 of G(−26). (C) T(−25) variant [5’–TATAATAG–3’, frequency = 35.2%, activity = 90%–110%]. The van der Waals contact between A C2 and Leu-72 is preserved [A(−25) C2–Leu-72 CD1 = 3.6 Å vs. A(−25) C2–Leu-72 CD1 = 3.7 Å in the AdMLP cocrystal structure]. This panel is very similar to the view illustrated in Fig. 5B, in which A:T and T:A base pairs make van der Waals contacts with another leucine side chain related by twofold quasisymmetry [Leu-163]. (D) T(−24) variant [5’–TATAAAAT–3’, frequency = 11.8%, activity = ND]. Like position 1 (Fig. 5A), all four base pairs are tolerated at this position. In the AdMLP cocrystal structure, the closest contact between protein and DNA involves CE1 of Phe-57 and the exocyclic NH2 of G(−24) [3.6 Å]. Substitution of G:C with T:A yields a similar van der Waals contact [A(−24) C2–Phe-57 CE1 = 3.4 Å].
TATA element recognition by TBP is conserved

Figure 8. Hoogsteen base pairs in two different TBP–DNA complexes. (A) C(−25) variant (5′-TATAAACG-3′, frequency = 3.4%, activity = 5%–6%). A C:G Watson–Crick base pair at this position would create a steric clash between the exocyclic NH₂ of G(−25) and the side chain of Leu-72. The C(−25) variant of the AdMLP TATA element avoids this steric clash by undergoing a 180° rotation about the C1′-N9 glycosidic bond, going from the normal anti to the syn conformation. The resulting Hoogsteen base pair [hydrogen bonds denoted with white broken lines] is stabilized by a G(−25) NH₂–C(−24) O1P hydrogen bond (2.9 Å) between the base and the DNA backbone. (B) Simulated annealing electron density map corresponding to the view illustrated in A, contoured at 2.5σ. The map shows excellent electron density for C in the Hoogsteen base pair. The electron density for G is not as well defined. (C) G:C Hoogsteen base pair at position 8 in the cocrystal structure of the T(−30) variant, superimposed on the corresponding base pair of the AdMLP cocrystal structure. Unlike the Hoogsteen base pair depicted in A and B, this unusual DNA structure is the result of a stabilizing interaction with a neighboring protein–DNA complex in the crystal lattice. (D) Simulated annealing electron density map corresponding to the view illustrated in C, contoured at 2.5σ. The map shows excellent electron density for the entire Hoogsteen base pair.

TATAAAC box, these Hoogsteen base pairs flank drug intercalation sites where unstacking occurs. Formation of a C:G (or G:C) Hoogsteen base pair stabilized by two hydrogen bonds [Fig. 8] requires protonation of C at position N3 [pKa = 4.6 in the absence of environmental effects]. X-ray crystallography at 1.95 Å resolution cannot reveal the protonation state of a particular titratable group, but we presume that the pKa of N3 is shifted toward neutral in our Hoogsteen base pairs, as observed by NMR in drug–DNA complexes [for review, see Escude et al. 1996].

We also detected a G:C Hoogsteen base pair on the other side of the 3′ kink, created by insertion of Phe-57 and Phe-74. Figure 8, C and D, illustrates a portion of the TTATA box cocrystal structure [T(−30) 5′-TTTAAAAG-3′, frequency = 13.9%, activity = 20%–29%]. Bucher’s [1990] studies of the EPD reveal no substantial base pair preference [Table 1], and our model building results suggest that all four Watson–Crick base pairs can be accommodated at position 8 [data not shown]. In this case, we are confident that the observed Hoogsteen base pair actually is an artifact of lattice packing within this particular protein–DNA crystal (each of the 10 newly determined cocrystal structures displays a unique lattice packing arrangement). Arg-65, protruding from a nearby TBP–DNA complex in the crystal lattice, makes two coplanar hydrogen bonds with G(−23) O6 and N7 [both 2.9 Å] and stacks on the π electron cloud of G(−24) [3.4 Å interplanar distance], thereby stabilizing the observed Hoogsteen base pair [Fig. 8C,D].

Detection of Hoogsteen base pairs adjacent to the 3′ kink site in the TATA element suggests that a similar situation might be obtained at the 5′ end of the recognition element, where the DNA is equally kinked. Many TATA boxes are quasisymmetric and the underside of the TBP saddle displays approximate twofold symmetry about an axis passing between β-strands S1 and S1′ and running perpendicular to the β-sheet [Fig. 4]. Leu-72 even has a counterpart, Leu-163, which is related by twofold quasisymmetry. On this side of the saddle, however, Leu-163 really does preclude G:C or C:G base pairs at position 2 [frequencies = 1%], and wild-type TBP does not make productive complexes with either TCTA or TGTA boxes [Wobbe and Struhl 1990]. Site-directed mutagenesis of Leu-163 → Val [with an accompanying Ile → Phe substitution] yielded a mutant TBP capable of directing Pol II transcription initiation in vivo from both TATA and TGTA boxes [Strubin and Struhl 1992]. Thus, the TGTA box must be able to undergo the obligate TBP-induced DNA deformation when it is bound to the mutant protein. Our attempts to cocrystallize wild-type TBP with oligonucleotides containing TCTA or TGTA boxes were unsuccessful [data not shown], and we believe that Hoogsteen base pair formation does not occur in the vicinity of the 5′ kink site.

This functional asymmetry between the 5′ and 3′ ends of the TATA element has been detected independently by Hurley and coworkers [Sun and Hurley 1995; L. Hurley, unpubl.]. Their chemical modification studies with pluramycin, a probe for deformed DNA, demonstrated that TBP-dependent reactivity extends beyond the confines of the 3′ end of 5′-TATAAAA-3′ but never 5′ of the T:A base pair at position 1. In contrast, a perfectly symmetric TATA element (5′-TATATAA-3′) shows equal TBP-dependent pluramycin reactivity at both ends of the oligonucleotide. Thus, subtle asymmetry in the deformability of various TATA element sequences may contribute to asymmetric be-
behavior during complex formation. If true, this phenomenon could help dictate the polarity of TBP binding, which continues to elude definitive explanation. This intrinsic effect of the TATA element would be independent of any polarity-determining role that TFIIIB may play, via interactions with the promoter upstream of the TATA box (Cox et al. 1997; Lagrange et al. 1998; Qureshi and Jackson 1998). It is remarkable that the TBP-dependent effects of pluramycin reactivity are not influenced by addition of TFIIIB [L. Hurley, unpublished].

Not surprisingly, all 10 newly determined cocrystal structures display subtle changes at the level of both TBP and DNA, when compared with our AdMLP cocrystal structure. The most remarkable difference was observed in our TATAAT box cocrystal structure [T(−26)5′-TATAATTAG3′, frequency = 2.8%, activity = 6%]. Modest structural rearrangements created a void in the protein–DNA interface that is occupied by a water molecule (Fig. 6C,D) in both halves of the asymmetric unit. This finding was somewhat unexpected, because there is no precedent for an ordered water molecule being found between TBP and DNA. It is made even more interesting by the fact that this particular TATA box is almost 20-fold weaker than its AdMLP counterpart, which differs by only 1 bp [A:T → T:A].

Functional definition of the TATA element

The wealth of information available on TBP–DNA complexes makes a functional definition of the TATA element possible. Inspection of extant cocrystal structures plus the results of model building exercises allow us to predict which of the four possible base pairs are tolerated at each of the eight positions comprising a TATA box. Specifically, we can predict which octameric sequences can present a minor groove surface that is complementary to the underside of the molecular saddle. Regrettably, we cannot make quantitative judgments regarding whether or not a particular combination of allowed base pairs will yield a sequence capable of undergoing the obligate conformational change on binding to TBP. We have already discussed the example of an A tract, but there are bound to be other sequence-specific effects that preclude certain combinations of allowed base pairs. For reviews on sequence-dependent effects in protein–DNA complexes see Dickerson (1998) and Olson et al. (1998).

Assuming that all functional TATA elements must present a complementary minor-groove surface to TBP after they undergo the same DNA deformation and taking each of the eight positions in turn, we derived a structure-based definition for a TATA element that is also consistent with the results of Bucher’s (1990) studies of the EPD.

TATA definition

\[
\begin{align*}
T_3 \times c > a \equiv g/A & >> t/T >> a \equiv c/A >> t/T >> a/ \\
A >> g > c \equiv t/A & \equiv T > g > c/G \equiv A > c \equiv t.
\end{align*}
\]

Position 1 \((T >> c > a \equiv g)\)

All four base pairs are observed in nature and are compatible with our structural insights. The EPD survey identifies a marked preference for T (frequency = 79.5%), which may be related to the fact that the energy of T:A stacking on the adjacent base pair is relatively small [except for T:A stacking on C:G] (Ornstein et al. 1978) and easy to overcome by intercalation of Phe-148 and Phe-165.

Position 2 \((A >> t)\)

A:T and T:A appear to be equally acceptable from the structural standpoint. Both G:C and C:G are forbidden by steric clashes with Leu-163. The EPD analysis confirms that G:C and C:G are vanishingly rare [frequencies = 1%]. There is a marked preference for A:T over T:A [frequencies = 83.5% and 13.9%, respectively], which is probably correlated with the frequency of T:A in position 1. T:A on A:T stacking energy is the lowest of all possible combinations (Ornstein et al. 1978), again favoring unstacking of the first two base pairs by Phe-148 and Phe-165.

Position 3 \((T >> a \equiv c)\)

T:A, A:T and C:G are all structurally compatible at this position, whereas the exocyclic NH2 of G:C would clash with Val-119. T:A dominates (frequency = 91.4%), which may reflect the preference for T:A in position 1 followed by A:T in position 2. An A:T in position 3 [frequency = 4.4%] would create an A tract.

Position 4 \((A >> t)\)

Our structures suggest that only A:T and T:A are permitted, which is consistent with database findings [frequencies = 89.2% and 8.4%, respectively]. Val-119 precludes G:C and C:G [both frequencies = 1%].

Position 5 \((T >> a)\)

Like position 4, A:T and T:A are structurally permitted and observed in nature [frequencies = 71.0% and 27.7%, respectively], whereas G:C and C:G are precluded by Val-29 [both frequencies <1%]. We infer that mutation of either of these critical valine residues [29 or 119] to alanine would greatly broaden the specificity of TBP binding.

Position 6 \((A >> g > c \equiv t)\)

Our collection of cocrystal structures includes A:T, T:A and G:C, all of which function in transcription (Wobbe and Struhl 1990). The remaining possibility, C:G, is present at a low but statistically significant level [frequency = 2.9%] and is active in transcription (Wobbe and Struhl 1990).

Position 7 \((A \equiv T > g > c)\)

The EPD contains examples of all four base pairs at this position. Our structural study focused on C:G [frequency = 3.4%] revealed the most significant structural
ing with G in the syn conformation, thereby avoiding a sterical clash with Leu-72.

Position 8 (G ≡ A > c ≡ t)

All four base pairs are found in the EPD and are compatible with the structural data.

Implications for promoter sequence conservation and regulation of Pol II transcription initiation

Our functional definition of the TATA element implies that 6144 of the 65,536 possible octameric sequences could present complementary minor-groove surfaces to the underside of the molecular saddle. The actual number must be lower, because some combinations of individually allowed base pairs cannot undergo the required conformational change during TBP binding. In advance of a systematic study, it is impossible to know how many putative TATA elements bind TBP productively, but it probably numbers in the thousands. Typical sequence-specific DNA-binding proteins could not bind such large ensembles of sequences with high affinity, because they interact with the major groove. Minor-groove recognition of TATA elements by TBP, on the other hand, represents an architecturally elegant solution to two potential problems arising from errors occurring during DNA replication. First, random point mutants in TATA boxes controlling expression of essential genes are not invariably lethal. Second, variations in TATA box sequence do not significantly perturb later steps in PIC assembly. TATA element deformation by TBP creates a structurally invariant nucleoprotein complex that serves as the receptor for TFIIB and TFIIA. Subsequent entrants to the PIC (Pol II/TFIIF, TFIIE, TFIIH) would also see the same multiprotein–DNA complex no matter what TATA box was actually present in the promoter. It is, of course, possible that one or more factors could gain access to the major-groove face of the TATA box while it is bound to TBP and affect transcription initiation from specific promoters, as suggested by the results of Lee and Roeder [1997].

Transcription activation in vivo is usually thought of as being composed of two distinct subprocesses, antirepression and true activation. We have already discussed the connection between TBP binding to the minor-groove face of the TATA box and the repressive effects of chromatin [Kim et al. 1993a], and now go on to consider true activation. Our crystallographic analysis of DNA recognition by TBP does not specifically address the issue of how transcriptional activators work, but the structural and functional data do provide some mechanistic insights. The fact that the cocrystal structures are essentially identical, despite 20-fold differences in transcription activity [all other things being equal in two different in vivo assays], immediately tells us that thermodynamic, kinetic, or dynamic differences must be responsible for the observed variations in transcriptional efficiency.

Preliminary biophysical observations demonstrate that our transcriptionally weaker TATA element variants bind to TBP with lower affinities when compared with the AdMLP [A.K. Mollah, B. Gilden, E. Jamison, M.D. Librizzi, S.K. Burley, I.M. Willis, and M. Brenowitz, in prep.]. Variations in association ($k_{on}$) and/or dissociation ($k_{off}$) rates could reduce both binding affinity ($K_D = k_{off}/k_{on}$) and the efficiency of transcription initiation. If the obligate, TBP-induced conformational change takes the same amount of time (i.e., $k_{on}$ remains unchanged) for some TATA elements, the half-lives of the corresponding TBP–DNA complexes must be decreased in the weaker TATA boxes (i.e., $k_{off}$ increases). Alternatively, changes in TATA-box sequence could also slow the rate of TBP–DNA complex formation. During Hoogsteen base pair formation at position 7, we know that $k_{on}$ is reduced [Mollah, S.K. Burley, and M. Brenowitz, in prep.], presumably because it takes longer to form a biochemically productive TBP–DNA complex dependent on rotation about the glycosidic bond. We suggest that some transcriptional activators will exert their positive effects on mRNA production by increasing the half-life of the foundation on which the PIC is assembled on a specific promoter. Presumably, this strategy allows regulatory proteins bound upstream to overcome the effects of an intrinsically weak TATA element, as seen for the Zta trans-activator protein [Lieberman and Berk 1991]. A transcriptional activator could also up-regulate gene expression by increasing TBP (or TFIID) recruitment and increasing $k_{on}$, which may well be the case with the artificial lex–TBP fusion transcription system [Chatterjee and Struhl 1995], the Zta trans-activator [Lieberman and Berk 1994], and other experimental systems cited in Chi et al. [1995].

Regrettably, our study does not provide any direct insight into the precise molecular mechanisms responsible for transcription initiation from the so-called TATA-less promoters [for review, see Smale et al. 1998]. However, we do believe that the functional definition of the TATA element discussed earlier will serve as a useful tool for identifying bona fide TATA-less promoters (i.e., sequences that are not capable of forming the protein–DNA complex illustrated in Fig. 1A). This definition does not preclude TBP binding to G/C-rich sequences upstream of transcription start sites with a different molecular recognition strategy. Although such a scenario represents a formal possibility, it seems more likely that other components of TFIID interact with the core promoters of class-II nuclear genes in the absence of a functional TATA element. Alternatively, initiator element-binding proteins could recruit components of the Pol II transcription machinery to a TATA-less promoter.

Conclusion

This paper presents an atomic resolution analysis of the molecular mechanisms responsible for TATA element recognition by TBP during Pol II transcription initiation. Our work provides a detailed picture of how TBP exploits minor-groove interactions and formation of Hoogsteen base pairs to recognize thousands of octameric sequences while inducing a dramatic distortion in
the DNA double helix. The structure of TBP bound to the deformed core promoter is independent of the origin of TBP and of the sequence of the TATA box, demonstrating that the Pol II PIC is assembled on a nucleoprotein foundation that has remained unchanged throughout evolution.

Materials and methods

Reagent preparation and crystallization

Wild-type TBP isoform 2 from Arabidopsis thaliana was overexpressed in Escherichia coli and purified to homogeneity (Nikolov et al. 1992). The 14-bp oligonucleotides containing variants of the AdMLP TATA box (Table 1) were prepared as described previously (Kim et al. 1993a). Cocrystals were obtained by mixing an equimolar ratio of DNA and TBP to form the complex at a final concentration of ~0.5 mM in 40 mM 2-(N-morpholino)ethane sulfonic acid (MES) at pH 5.9, 60 mM or 100 mM KCl [depending on the oligonucleotide], 4 mM MgCl₂, 14% [vol/vol] glycerol, 300 mM ammonium acetate, and 10 mM DTT and equilibrating against a reservoir containing 12% [vol/vol] glycerol, 25 mM MES (pH 5.9), and 10 mM DTT with sitting drop vapor diffusion at 4°C. Following seeding, plate-like cocrystals (0.8 × 0.8 × 0.1 mm) grew in weeks.

X-ray data collection, structure determination, and refinement

Diffraction data were obtained from flash-frozen cocrystals via X-ray data collection, structure determination, and refinement. The publication costs of this article were defrayed in part by payment of page charges. This article must therefore be hereby marked “advertisement” in accordance with 18 USC section 1734 solely to indicate this fact.

Acknowledgments

We thank Drs. L. Berman, M. Capel, and R.M. Sweet for help at beamline X25 at the National Synchrotron Light Source; Drs. S. Ealick and D. Thiel and the MacCHESS staff for help at beamlines A1 and F1 at the Cornell High Energy Synchrotron Source; E. Halay for help with DNA production; Drs. J. Bonanno, D. Jouzalim, J. Marcotrigiano, D.B. Nikolov, S.K. Nair, and X. Xie for useful discussions and much help with computing and figure preparation. For their many useful suggestions, we are grateful to Drs. K. Arndt, M. Brenowitz, P. Bucher, R.E. Dickerson, J. Kahn, L. Hurley, J. Kuriyan, W.K. Olson, G.A. Petsko, R.G. Roeder, P.B. Sigler, and M. Vasseur. This work was supported by the Howard Hughes Medical Institute (S.K.B.).

The reporter gene experiments

The reporter plasmids described above were transformed into yeast strain Sc18 (GAL4, gal80, ura3-52, leu2-3,112, his3, trp, MEL1). Transformed cells were grown on minimal media lacking uracil to mid-log phase and then harvested as described previously (Vashee and Kodadek 1995). The carbon source was 2% galactose, 3% glycerol, and 2% lactic acid. Reported β-galactosidase values are accurate to ±15% and are the result of at least three independent, replicate measurements. For AdMLP, the absolute levels of β-galactosidase activity supported by the consensus and MEL1 UAS Gal4-binding sites differed by an order of magnitude. In Table 1, each is listed as 100% and the activity levels obtained with the other TATA elements are appropriately normalized.

Construction of reporter gene plasmids with variant TATA elements

A synthetic promoter was constructed by introducing the AdMLP TATA element into the yeast Gal1 promoter with a Gal4-binding site [consensus UAS (5′-CGGACCAGTCGTCGTTCCG-3′) located 200 bp upstream. The synthetic promoters were placed immediately upstream of a lacZ reporter gene, yielding two reporter plasmids (pLS1 and pLS2), which differ only in the sequence of the Gal4-binding site. By use of pLS1 and pLS2 as parent plasmids, other constructs containing nine variant TATA elements were made by site-directed mutagenesis.

References


Patikoglou et al.
TATA element recognition by TBP is conserved


Roeder, R.G. 1996. The role of general initiation factors in tran-


Starr, D., B. Hoopes, and D. Hawley. 1995. DNA bending is an important component of site-specific recognition by the TATA binding protein. *J. Mol. Biol.* 250:434–446.


TATA element recognition by the TATA box-binding protein has been conserved throughout evolution

Georgia A. Patikoglou, Joseph L. Kim, Liping Sun, et al.

*Genes Dev.* 1999, 13:

References

This article cites 53 articles, 14 of which can be accessed free at:

http://genesdev.cshlp.org/content/13/24/3217.full.html#ref-list-1

License

Email Alerting Service

Receive free email alerts when new articles cite this article - sign up in the box at the top right corner of the article or click here.