Radical mutations reveal TATA-box binding protein surfaces required for activated transcription in vivo

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Regions on the surface of human TATA-box binding protein (TBP) required for activated transcription in vivo were defined by construction of a library of 89 surface residue mutants with radical substitutions that were assayed for their ability to support activated transcription in vivo, basal transcription in vitro, and TFIIA and TFIIIB binding in vitro. Four epitopes were identified in which substitutions in two to four neighboring surface residues greatly inhibited activated transcription in vivo. One epitope in which substitutions inhibited both basal and activated transcription (E284, L287) is the interface between TBP and TFIIIB. Another (A184, N189, E191, R205) is the recently determined interface between TBP and TFIIA. Mutations in residues in this TFIIA interface greatly inhibit activated, but not basal transcription, demonstrating a requirement for the TFIIA-TBP interaction for activated transcription in vivo in mammalian cells. The remaining two activation epitopes (TBP helix 2 residues R231, R235, R239, plus F250; and G175, C176, P247) are probably interfaces with other proteins required for activated transcription. The library of mutants responded virtually identically to two different types of activators, GL4-E1A and GAL4-VP16, indicating that transcriptional activation by different classes of activators requires common interactions with TBP.

[Key Words: TBP; TFIIID; transcription; activation; TFIIA]

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The TATA-box binding protein (TBP) is an essential transcription factor for each of the three eukaryotic nuclear RNA polymerases (Hernandez 1993; Burley and Roeder 1996). In mammalian and Drosophila systems for transcription of protein-coding genes by RNA polymerase II (Pol II), TBP is associated with ~12 (human) or 9 (Drosophila) additional polypeptides called TBP-associated factors (TAFs) in the general transcription factor TFIIID (Dynlacht et al. 1991; Goodrich and Tjian 1994, Burley and Roeder 1996). On promoters with a TATA box (~80% of protein-coding genes, Bucher 1990), TFIIID initiates assembly of an approximately ribosome-size preinitiation complex (~4 x 10^3 kD) by binding to TATA-box DNA (Davison et al. 1983; Burley and Roeder 1996). The TBP polypeptide binds to general transcription factors TFIIA (Geiger et al. 1996, Tan et al. 1996) and TFIIIB (Nikolov et al. 1995) and can participate in an in vitro basal transcription reaction along with TFIIB, TFIIIF, Pol II, TFIIIE, and TFIIFH (Burley and Roeder 1996). Activation of transcription in vitro, however, requires TFIIID TAFs (Dynlacht et al. 1991; Goodrich and Tjian 1994, Burley and Roeder 1996) and other coactivators (Ge and Roeder 1994, Kaiser et al. 1995; Luo and Roeder 1995, Shykind et al. 1995), or, in Saccharomyces cerevisiae, polypeptides associated with a Pol II holoenzyme complex (Kim et al. 1994; Koleske and Young 1994).

In vitro TBP binds multiple polypeptides in addition to TFIIA and TFIIIB. Among these are activation domains (Stringer et al. 1990; Horikoshi et al. 1991; Lee et al. 1991), the Pol II large subunit carboxy-terminal repeat domain (CTD) (Usheva et al. 1992), repression domains (Um et al. 1995), transcriptional inhibitors (Yeung et al. 1994; Kim et al. 1995), and TFIIF (Tang et al. 1996). To test the significance of these interactions observed in vitro for activated transcription in vivo, we constructed a library of human TBP mutants, each with a single amino acid residue substitution within the 180-residue essential carboxy-terminal core domain at a site where the side chain of the wild-type residue is exposed to solvent in the crystal structure of the TBP–TATA box DNA complex (Kim et al. 1993, 1995). Only the TBP core domain was analyzed, as the amino-terminal domain of human TBP is not required for activated transcription in vivo (Q. Zhou et al. 1993) or in vitro (Keveaney et al. 1993, Tansey et al. 1994). We expected that substitutions in a subset of surface residues would weaken or block TBP binding to proteins with which it must normally interact.
during regulated transcription. By correlating the effects of mutations on TBP binding in vitro with effects on transcription in vivo, we hoped to distinguish which interactions observed in vitro make important contributions to transcription in vivo.

Earlier studies of TBP mutants have already revealed a great deal about how the protein functions in Pol II transcription. Residues E284, E286, and L287 at the tip of the second stirrup of the saddle-shaped molecule [Kim and Burley 1994] have been shown to be critical to TFII B binding and in vitro transcription [Kim et al. 1994, 1995; Tang et al. 1996]. An analysis of human TBP mutants with multiple simultaneous substitutions in surface residues suggested that the TBP interaction with TAFu250 and several have been described recently that have substitutions on the upper surface of the TBP molecule [Stargell and Struhl 1996]. Further genetic analyses support the model that activator DNA is sufficient to activate transcription [Stargell and Struhl 1995], and several have been described recently that have substitutions in the upper surface of the TBP molecule [Stargell and Struhl 1996].

Results

In most cases, we introduced radical amino acid substitutions onto the surface of TBP to increase the likelihood that a single mutation would disrupt TBP's interaction with another protein in vitro and produce a clear phenotype in vivo. Lysine and arginine residues were substituted with glutamic acid, and aspartic and glutamic acid residues were substituted with arginine to introduce a bulky, charged side chain that might sterically block interactions with neighboring regions of the TBP surface. Because we were interested in identifying sites of protein-protein interaction, we did not mutate TBP residues that interact directly with DNA, except in a few cases. Eighty-nine single-residue substitution mutants [Table 1] were constructed and tested for their abilities to support activated transcription in vivo in a transient transfection assay in COS cells, to interact with TFII A and TFII B in gel mobility shift assays, and to support basal in vitro transcription.

Mutant TBPs expressed in and purified from Escherichia coli [Fig. 1A] were assayed for DNA binding alone and in the presence of recombinant TFII A and/or TFII B by gel mobility shift assay [Buratowski et al. 1989; Table 1; representative results shown in Fig. 1B]. These results will be discussed below together with results on in vivo function of these mutants. But in analyzing the in vivo data, it is essential to eliminate the class of mutations that inhibit TBP DNA-binding activity. Considering the radical nature of the substitutions introduced, we anticipated that some of the mutations might significantly distort TBP's conformation or even interfere with protein folding. Such mutants were recognized by their inability to bind DNA, because the structure of the TBP DNA-binding surface, that is, the bottom of the saddle-shaped molecule [see Fig. 5, below], depends on the overall conformation of the 180 residue carboxy-terminal core domain. Fourteen mutants failed to form DNA-protein complexes with TATA-box DNA in the gel mobility shift assay in the presence of either TFII A or TFII B [Table 1; e.g., Fig. 1B, A190E]. These mutants may be improperly folded, and any reduction in function they may exhibit in vivo [see below] may be explained by their inability to bind TATA-box DNA.

To assay the influence of these surface residue substitutions on activated transcription in vivo, surface mutations were constructed on the background of the m3 triple mutation [Strubin and Struhl 1992]. The three m3 mutations in the DNA-binding surface of TBP relax its binding specificity, allowing it, unlike wild-type TBP, to bind to a TG TAAA sequence with an affinity similar to that of wild-type TBP for a TATAAA box. This experimental method allows one to assay the influence of additional mutations in TBP on its ability to support Pol II transcription in vivo by assaying expression from a reporter gene with a TG TAAA box [Strubin and Struhl 1992].

We called the human TBP mutant with the three m3 mutations in the DNA-binding surface, but otherwise wild-type surface residues, wt hm3. Wt hm3 and all of the TBP mutants were epitope-tagged at the amino terminus so that expression levels could be assayed by Western blotting. We used two different TG TAAA-box reporter constructs with upstream GAL4-binding sites to test the generality of the mutant TBP phenotypes on different promoters. One used the promoter region from c-fos [Tansey et al. 1994], and the other, the β-retinoic acid receptor gene promoter region [Keaveney et al. 1993; Fig. 2].

To determine whether different surfaces of TBP are required for activation by different classes of activators, we analyzed activation by two different activation domains, one in GAL4–VP16 [Sadowski et al. 1988] and the other in GAL4–E1A conserved region 3 (CR3) [Martin et al. 1990]. These two activation domains probably function by distinct mechanisms because VP16 is a prototype acidic activation domain [Cress and Triezenberg 1991], see also Leuther et al. 1993], whereas E1A CR3 is not an acidic activator [Martin et al. 1990; Webster and Ricciardi 1991]. Also, GAL4–VP16 is a potent activator in S. cerevisiae [Sadowski et al. 1988], whereas GAL4–E1A is...
Table 1. Phenotypes of TBP mutants

| Position Mutation | c-fos E1A | EIA VP16 | β-ret E1A | VP16 | in vitro TnX | T | TA | TB | TAB column |
|-------------------|-----------|----------|-----------|------|-------------|---|----|----|-----------|
| S156E             | 69±18     | 78±17    | 85±15     | 48±4 | 29±4        | 3±3| 66±14| 11±19      | 70±4       |
| E197R             | 20±5      | 35±5     | 60±5      | 11±5 | 1±1         | 0±0| 2±2 | 0±0         | 1±1        |
| S158E             | 65±15     | 73±15    | 111±15    | 78±7 | 55±5        | 37±3| 60±11| 60±11       | 3±3        |
| S159E             | 111±17    | 61±17    | 51±17     | 60±6 | 41±4        | 3±3| 67±11| 67±11       | 3±3        |
| E161R             | 29±12     | 22±12    | 39±12     | 47±4 | 16±4        | 3±3| 66±11| 66±11       | 3±3        |
| L165E             | 7±4       | 18±4     | 18±4      | 5±5  | 22±2        | 3±3| 67±11| 67±11       | 3±3        |
| N173E             | 52±7      | 58±7     | 54±7      | 44±4 | 44±4        | 3±3| 66±11| 66±11       | 3±3        |
| G175R             | 18±2      | 22±2     | 41±2      | 47±4 | 6±6         | 3±3| 67±11| 67±11       | 3±3        |
| C176R             | 9±3       | 6±3      | 11±3      | 11±3 | 11±3        | 3±3| 67±11| 67±11       | 3±3        |
| K177E             | 72±12     | 121±12   | 63±12     | 70±7 | 8±8         | 3±3| 67±11| 67±11       | 3±3        |
| L244R             | 39±6      | 56±6     | 57±6      | 11±3 | 3±3         | 0±0| 2±2 | 0±0         | 1±1        |
| K181E             | 59±11     | 59±11    | 11±11     | 11±11| 11±11       | 3±3| 67±11| 67±11       | 3±3        |
| A184E             | 1±1       | 1±1      | 1±1       | 1±1  | 1±1         | 0±0| 2±2 | 0±0         | 1±1        |
| R188E             | 160±18    | 125±18   | 105±18    | 11±3 | 49±4        | 3±3| 66±11| 66±11       | 3±3        |
| A190E             | 1±1       | 1±1      | 1±1       | 1±1  | 1±1         | 0±0| 2±2 | 0±0         | 1±1        |
| N193R             | 6±4       | 14±4     | 9±4       | 14±4 | 14±4        | 3±3| 66±11| 66±11       | 3±3        |
| E191R             | 20±2      | 20±2     | 11±2      | 7±7  | 7±7         | 3±3| 66±11| 66±11       | 3±3        |
| N199E             | 26±3      | 53±3     | 50±3      | 44±4 | 44±4        | 3±3| 66±11| 66±11       | 3±3        |
| V194E             | 30±4      | 48±4     | 69±4      | 69±4 | 69±4        | 3±3| 66±11| 66±11       | 3±3        |
| R203E             | 20±2      | 20±2     | 11±2      | 7±7  | 7±7         | 3±3| 66±11| 66±11       | 3±3        |
| R205E             | 1±1       | 1±1      | 1±1       | 1±1  | 1±1         | 0±0| 2±2 | 0±0         | 1±1        |
| F215E             | 6±6       | 215±6    | 6±6       | 215±6| 6±6         | 3±3| 66±11| 66±11       | 3±3        |
| G223R             | 6±6       | 6±6      | 6±6       | 6±6  | 6±6         | 3±3| 66±11| 66±11       | 3±3        |
| E227R             | 6±6       | 6±6      | 6±6       | 6±6  | 6±6         | 3±3| 66±11| 66±11       | 3±3        |
| E228R             | 6±6       | 6±6      | 6±6       | 6±6  | 6±6         | 3±3| 66±11| 66±11       | 3±3        |
| R231E             | 6±6       | 6±6      | 6±6       | 6±6  | 6±6         | 3±3| 66±11| 66±11       | 3±3        |
| L232E             | 6±6       | 6±6      | 6±6       | 6±6  | 6±6         | 3±3| 66±11| 66±11       | 3±3        |
| R235E             | 6±6       | 6±6      | 6±6       | 6±6  | 6±6         | 3±3| 66±11| 66±11       | 3±3        |
| V240E             | 6±6       | 6±6      | 6±6       | 6±6  | 6±6         | 3±3| 66±11| 66±11       | 3±3        |
| K249E             | 6±6       | 6±6      | 6±6       | 6±6  | 6±6         | 3±3| 66±11| 66±11       | 3±3        |

Diagrams at the left indicate the position of the mutation on the surface of TBP (highlighted in red and enlarged) with the TBP molecule shown in one of the three positions presented in Fig. 5. The first four columns show results of in vivo function assays. Luciferase activity is shown as a percentage of that observed with wt hm3 for activation by Gal4-E1A (E1A) and Gal4-VP16 (VP16) from TGTAAA-box reporter genes using the c-fos and β-retinoic acid receptor gene (β-ret) promoter regions. Values are means of three independent transfection assays ± S.D. “in vitro TnX” shows activity in a basal transcription reaction (see Materials and Methods) as a percentage of the transcription observed with wt hm3. Values shown ± S.D. are means of 2–3 assays. Values without a S.D. are from a single assay. Columns 6–9 summarize gel shift assay results for reactions with mutant TBP alone (T), plus TFIIA (TA), plus TFIIB (TB), plus TFIIA and TFIIB (TAB). (+) Addition of protein produced a new complex; (+) addition of protein produced a new complex but with a different mobility from wt hm3; (-) no new complex observed compared with mutant TBP alone. The right column shows the percent binding to a Gsti–TFIIAαβγ column relative to 100% for wt hm3. Values showing S.D. are averages of 2–4 determinations.

an extremely weak activator in this organism (T.G. Boyer and A.J. Berk, unpubl.).

Of the 75 TBP mutants that bind TATA-box DNA together with either TFIIA or TFIIB, 13 were severely reduced (<20% of the activity of wt hm3) in their ability to support transcriptional activation by GAL4-E1A (Table 1, Fig. 3). Similar results were observed when each mutant was tested for its ability to support activation by GAL4-VP16 (Fig. 3). Results with both activators were similar on both the c-fos and β-retinoic acid receptor TGTAAA-box reporter genes (Table 1). Because similar results were observed for each TBP mutant on two different promoter regions and in response to two different activators, these results probably reflect general aspects of TBP function in Pol II transcription from promoters with a TATA box. We were not able to detect significant expression from these reporter genes without cotransfecting an expression vector for a GAL4 DNA-binding domain fused to an activation domain. Consequently, in vivo basal activity (if such a thing occurs in mammalian cells) could not be assayed.

To determine whether the in vivo activity of these mutants was affected by their expression levels, mutant TBPs were analyzed by Western blotting of whole-cell extracts from transfected cells with a monoclonal antibody specific for the amino-terminal epitope tag [repre-
Figure 1. Gel mobility-shift assays. (A) Representative Coomassie blue-stained SDS–PAGE gel of purified mutant hTBPs expressed in E. coli. Mutants E228A and R188E were expressed at lower levels than the other mutants. (B) Representative gel mobility-shift assays. Letters above a lane indicate the proteins added to the binding reaction: (T) mutant TBP; (A) TFIIA; (B) TFIIB.

Figure 2. Plasmids used in the in vivo function assays. The reporter plasmid was derived either from the c-fos promoter region (Tansey et al. 1994) or the β-retinoic acid receptor gene promoter region (Keaveney et al. 1993). Mutant TBPs with an amino-terminal epitope-tag were expressed from an SRα promoter/enhancer. GAL4 DNA-binding domain fusions to E1A conserved region 3 or the VP16 activation domain were expressed from the early SV40 promoter/enhancer, as was β-galactosidase as an internal control for transfection efficiency.

sentative data shown in Fig. 4]. Expression levels varied, but did not correlate with activity in the transfection assay. For example, mutants L185E and F280E had in vivo activities similar to that of wt hm3, yet were expressed at lower levels. This indicates that the high level of altered-binding-site TBP expressed from these replicating vectors saturates the transient transfection assay. Several mutants that were expressed at levels higher than wt hm3 had activities similar to wt hm3 [E228A, S215E, P267E, and V317E], indicating that squelching does not appear to be an important factor at these levels of expression in this assay.

In four successive transfection/Western blot experiments like that shown in Figure 4, for the mutants with <20% wt hm3 activity, N189E, R205E, R231E, R235E, R239S, E284R, and L287E were expressed at 50–100% the level of wt hm3. Mutants C176R, E191R, and R239S were expressed at 25–50% of wt hm3 concentration. Mutants G175R, A184E, and F250E, however, were consistently expressed at about 25% or less of the level of wt hm3. Consequently, the low in vivo activities of mutants N189E, E191R, R205E, R231E, R235E, R239S, E284R, and L287E result from defects in the function of these mutant TBPs and not from low expression levels. On the other hand, the low expression levels of mutants G175R, A184E, and F250E may have contributed to their low in vivo function. Also, the high expression level of some mutants in the transfection assay might mask the effects of mutations by driving weakened associations with other transcription factors through mass action.

The regions on TBP’s surface in which mutations significantly blocked in vivo function fall into four epitopes of two to four neighboring residues [Fig. 5]: the tip of the second repeat stirrup [E284R, L287E], the upstream surface of the first repeat [A184E, N189E, E191R, R205E], the top of the first repeat [H2 helix residues R231E, R235E, R239S, and F250E], and a region on the downstream surface of the first repeat, near the top [G175R, C176R, P247E]. The phenotypes produced by substitutions in two of these regions can be explained by the disruption of presently understood TBP interactions with two general transcription factors.
Mutations at E284, E286, and L287 inhibited TFIIB binding to TBP–TATA box DNA in an earlier alanine scanning analysis of the human TBP surface (Tang et al. 1996). Our substitutions in the same residues also inhibited TFIIB binding (Table 1; Fig. 1B). These three residues are deeply buried in the TFIIB core domain in the TFIIB–TBP–TATA box DNA crystal structure, and each of these three residues directly contacts TFIIB residues [Nikolov et al. 1995; see Fig. 6]. Consequently, the finding that our substitutions E284R, E286R, and L287E inhibit the TFIIB–TBP interaction is readily explained. The crystal structure of the complex also shows contacts between TFIIB and TBP residues corresponding to human residues Y283 and P285. Our mutations constructed at these positions yielded mutant TBPs that do not bind DNA (Table 1). Consequently, their impact on TFIIB binding could not be assessed in the gel mobility shift assay.

These three mutants [E284R, E286R, and L287E] were also defective for basal in vitro transcription (Table 1; Fig. 7), consistent with biochemical studies showing that TFIIB is required for both basal and activated in vitro transcription (Buratowski and Zhou 1993; Ha et al. 1993; Hisatake et al. 1993). Consequently, the failure of mutants E284R and L287E to support activated transcription in vivo is likely caused by a block in TFIIB binding.

Surprisingly, mutant E286R was reduced by only a factor of two in the in vivo assay (Table 1; Fig. 3), even though its ability to form a gel mobility shift complex with TBP–DNA was extremely reduced, even at high TFIIB concentration (Fig. 8B), and even though it had no in vitro basal transcription activity (Fig. 7).

Substitutions in a second epitope on the upstream surface of the first TBP repeat (A184E, N189E, E191R, R205E; Fig. 5) inhibited formation of the TFIIA–TBP–DNA (TA) complex in the gel mobility shift assay (Table...
Figure 4. In vivo expression of mutant TBPs in transfected cells. Mutants with <50% of the activity of wt hm3 and eight mutants with >50% of this in vivo activity were analyzed. COS cells were transfected with the standard amount of mutant TBP expression vector used in the in vivo assays of Table 1 and Figure 3. TBPs with two copies of the amino-terminal epitope were expressed. Total cell extracts were analyzed by Western blotting with monoclonal antibody 12CA5 against the amino-terminal epitope-tag on the mutant TBPs. The region of the gel with TBP is shown. Results for C176R and P247E were from a separate transfection experiment, with the wt hm3 control shown immediately to the left.

These results are perfectly consistent with the recent crystal structures of yeast TFIIA deletion mutant-core TBP–TATA box DNA complexes (Geiger et al. 1996; Tan et al. 1996, Fig. 6). Residues A184, N189, E191, and R205 are conserved between the yeast and human proteins, and the equivalent yeast amino acid side chains each make contacts with TFIIA in the yeast protein crystal structures. Substitutions A184E, N189E, and E191R block formation of the TA complex even at high TFIIA concentration (Fig. 8A), although these mutant TBPs can form DNA complexes with TFIIB (Fig. 1B). Mutant R205E formed detectable TA complex, although to a significantly lesser extent than wt hm3 (Fig. 8A). Substitutions in surface residues at the top of the first TBP repeat (TBP helix 2, R231E, R235E, R239S), in which earlier reports indicated possible contacts with TFIIA (Buratowski and Zhou 1992, Lee et al. 1992, Tang et al. 1996), did not affect TFIIA binding in the gel mobility shift assay (Fig. 1B, 8A). [Note that the lowest concentration of TFIIA used in the gel mobility shift assay with these mutants was just sufficient to shift wt hm3 (see Fig. 8A, lower right). This indicates that the affinity of these mutants for TFIIA was within twofold of wt hm3 as measured by the gel mobility shift assay.] However, several of these mutations did reduce TBP binding in a GST–TFIIA column-binding assay [Table 1]. Consequently, mutations in this region of the molecule may have small effects on the affinity for TFIIA, but we find that they have much less significant effects than mutations on the upstream surface of the first repeat (Fig. 5), where TBP contacts TFIIA in the crystal structures [Fig. 6].

Mutants R205E, E191R, and N189E had basal in vitro transcription activities similar to wt hm3 (Table 1; Fig. 7). These results are consistent with earlier biochemical studies showing that TFIIA is not required for basal in vitro transcription in reactions with TBP and purified transcription factors (Sayre et al. 1992; Sun et al. 1994). These mutants, however, had ≈5% of the activity of wt hm3 for transcription activated by GAL4–VP16 or GAL4–E1A in vivo. This phenotype is similar to the yeast TBP double mutant N2-1, characterized by Stargell and Struhl (1995), which is defective for binding to TFIIA in vitro and for supporting activated transcription in vivo, but which does support in vivo transcription from unregulated yeast promoters. Mutants R231E, R235E, R239S, and P247E on the upper surface of the first repeat [Fig. 4] were also extremely defective for supporting activated transcription in vivo (Table 1, Fig. 2), yet had high activity for basal in vitro transcription (Fig. 7).

Discussion

Our systematic mutational analysis of the TBP surface revealed four epitopes in which single substitutions in any of two to four neighboring surface residues significantly impair the ability of TBP to support activated transcription in vivo. Two of these epitopes represent surfaces of TBP that bind the general transcription factors TFIIA and TFIIIB, respectively. It is likely that the remaining two epitopes also represent TBP surfaces that interact with proteins required for activated transcription in vivo. The TFIIA and TFIIIB interaction surfaces are of similar size to the surface of E. coli CAP protein that interacts with the α subunit of E. coli RNA polymerase at the lac promoter (Y. Zhou et al. 1993; Busby and Ebright 1994). As such, these interaction surfaces are large enough to determine specific interactions in the preinitiation complex, but small enough to be sufficiently weak to allow the preinitiation complex to dissociate following initiation.

The TFIIA interaction epitope

Our identification of TBP mutations that inhibit TFIIA binding are perfectly consistent with the recent crystal structures of complexes of the core of yeast TBP, TATA-box DNA, and viable deletion mutants of yeast TFIIA (Geiger et al. 1996; Tan et al. 1996). The yeast TBP residues equivalent to the critical human TBP residues R205, N189, A184, and E191, in which substitutions inhibited TFIIA binding, all contact TFIIA in the crystal structure models. The crystal structure models also suggest contacts between TFIIA and side chains of TBP residues equivalent to human residues K181, L185, R188, A190, Y192, N193, R203, and I204. Our results with substitutions at K181, L185, and N193 [Table 1, Fig. 8A] suggest that these residues make less significant contributions to the energetics of the interaction than R205.
Figure 5. Positions of critical residues on the surface of TBP. Residues where mutations severely depress function are highlighted in red on space filling models of *Arabidopsis thaliana* TBP. The upstream surface of TBP is shown at the bottom; the molecule is rotated 90° in the middle of the figure to show the top, and then another 90° at the top to show the downstream surface. The first column highlights mutations that reduce in vivo transcription activated by Gal4-E1A to <20% of wt hm3. The middle column highlights residues in which mutations severely inhibit binding to TFIIB and the right column mutations that severely inhibit binding to TFIIA.

N189, A184, and E191. Comparisons between these radical mutations, however, are complicated because they may both eliminate interactions of the wild-type side chains and introduce repulsive forces into a protein–protein interface. We cannot draw conclusions about the significance of contacts with R188, A190, or Y192 because our substitutions at these positions prevented DNA binding even in the presence of TFIIA and TFIIB, probably by causing significant changes in the conformations of the mutant TBPs. We did not mutagenize I204 because the side chain is largely buried. Interpretation of the reduced gel mobility shift activity of mutant R203E is complicated by the fact that the normal arginine at this position also contacts a phosphate in TATA-box DNA and, therefore, contributes to the stability of the TBP–DNA interaction (Kim and Burley 1995). However, this mutant did form a gel mobility shift complex with both TFIIA and TFIIB (Table 1), suggesting that it is properly folded, and it was reduced in binding to a GST–TFIIA column (Table 1), as predicted by the crystal structure models.

In vitro studies have been somewhat equivocal about the function of TFIIA. TFIIA had no affect on basal transcription in reactions with TBP (Sun et al. 1994), but was

Figure 6. Structures of the TBP–TFIIB–DNA and TBP–TFIIA–DNA complexes with TBP residues highlighted where mutations inhibit complex formation. *Arabidopsis* TBP is present in the TFIIB complex (Nikolov et al. 1995), and *S. cerevisiae* TBP is in the TFIIA complex (Geiger et al. 1996). The highlighted TBP residues are visible in the upper portion of the figure where TFIIB and TFIIA are shown with wire diagrams. The space-filling models of TFIIB and TFIIA in the lower portion of the figure almost completely obscure the highlighted residues that are buried in the protein–protein interfaces.
either required for (DeJong et al. 1994; Ozer et al. 1994), or stimulated (Sun et al. 1994; Yokomori et al. 1994; Kang et al. 1995) basal transcription reactions with TFIIID. In some studies TFIIA was required for activated transcription (DeJong et al. 1994; Ozer et al. 1994), whereas in others it merely stimulated activated transcription a fewfold (Sun et al. 1994; Yokomori et al. 1994; Kang et al. 1995). Like TFIIID TAFs, TFIIA was not required for activated transcription in reactions with the yeast holoenzyme (Koleske and Young 1994; T. Kim et al. 1994). Yet, the genes encoding the subunits of yeast TFIIA are required for viability (Ranish et al. 1992), and shifting yeast cells with temperature-sensitive mutations in TFIIA to the nonpermissive temperature causes a drop in Pol II transcription (Kang et al. 1995). Stargell and Struhl (1995) isolated a yeast TBP mutant defective in supporting transcriptional activation in vivo by acidic activators but not constitutive transcription and found that it was also defective in binding TFIIA. These, plus other studies, led them to conclude that the interaction between TBP and TFIIA is required for activation by acidic activators in vivo (Stargell and Struhl 1995). Our results with human TBP mutants R205E, E191R, and N189E amply confirm this conclusion. As discussed above, these mutations occur in the TBP–TFIIA interface and inhibit TFIIA binding. They are able to form complexes with TFIIB and support near wild-type levels of in vitro basal transcription, indicating that they make all general transcription factor interactions required for this reaction. Nevertheless, they are severely defective for activated transcription in vivo. Moreover, these mutants are defective for activation by both an acidic [VP16] and a nonacidic [E1A CR3] activation domain. These results imply that the TBP–TFIIA interaction is generally required for transcriptional activation in vivo in mammalian cells.

Although these experiments show that the TBP–TFIIA interaction is required for activated transcription in vivo, they do not reveal how this interaction affects transcription. Earlier experiments indicated that a class of activation domains interact with TFIIA (Ozer et al. 1994, Kobayashi et al. 1996) to stimulate the extent, rate of formation, and stability of TFIIA–TFIID complex assembly on promoter DNA (Wang et al. 1992, Lieberman and Berk 1994; Ozer et al. 1994; Chi et al. 1995). The greatly increased rate of assembly may enhance transcription by protecting assembling complexes from inhibitory competing reactions in vivo, such as ATP-dependent dissociation by Mot1 (Auble et al. 1994) or formation of inactive complexes with TBP-binding inhibitors such as NC2/Drl (Yeung et al. 1994; Kim et al. 1995). TFIIA–TBP complexes are resistant to both of these inhibitory mechanisms. Activators also induce a conformational change in TFIIA–TFIID-promoter complexes (Lieberman and Berk 1994; Chi et al. 1995, Shykind et al. 1995), visualized by strong nuclease hypersensitivity near the transcription start site of some promoters (Lieberman and Berk 1994; Chi et al. 1995). This appears to be an activated conformation of the complex because it correlates with high initiation rates when the remaining transcription factors and Pol II are added. Consequently, the TFIIA–TBP interaction may be required for activators to stimulate the assembly of preinitiation complexes and to induce structural changes in the complex that increase the rate of initiation.

**Figure 7.** Basal in vitro transcription. Transcription was performed as described in Materials and Methods. For each mutant, control reactions without added TBP and with wt hm3 from the same gel and same exposure time are shown to the left.

**Figure 8.** Gel mobility-shift assay titrations. (A) For wt hm3 and each mutant [except L185E] five lanes show (from left to right) binding with TBP alone (5 ng), and plus 15, 30, 60, and 120 ng of recombinant TFIIA, respectively. For A184E, N189E, and the mutants shown in the top row, the assay performed with wt hm3 is shown at the top left. For F250E and L185E, the wt hm3 assay performed at the same time is shown to the left. For the L185E titration, 0, 4, 8, 15, 30, and 60 ng of rTFIIA were added from left to right. (B) Binding reactions contained mutant or wt hm3 TBP alone (5 ng), and plus 20, 60, 180, and 540 ng recombinant TFIIB, respectively, in lanes from left to right.
**Additional activation epitopes**

Mutations in two other regions on the TBP surface also inhibited activated transcription in vivo (Fig. 5). In interpreting the in vivo results for these mutants, it is critical to determine whether or not these mutations significantly alter the structure of TBP. One test for native protein structure was DNA-binding activity. All of the mutants in these two epitopes (R231E, R235E, R239S, F250E, and G175E, C176R, and P247E) formed gel mobility shift complexes with TFIIA and TFIIIB (Fig. 1B, 8A). Of particular significance regarding the question of mutant protein structure, mutants R231E, R235E, R239S, and P247E were functional for basal in vitro transcription (Fig. 7). This important result indicates that these mutants make all the general transcription factor interactions required for this complex reaction. Furthermore, mutants R231E, R235E, R239S, and P247E were expressed at higher level in the transfection assay than mutants F280E and L185E that have >70% wt hm3 activity (Fig. 3, 4). We conclude that mutants R231E, R235E, R239S, and P247E are severely defective for in vivo transcription because they are defective in making required TBP interactions, not because they are improperly folded or because they are not expressed at sufficient level. Consequently, as for the TFIIA-interaction surface, the TBP surface in which these substitutions occur is required specifically for the response to transcriptional activators. Because mutants G175R and C176R were defective for basal in vitro transcription yet bound TATA-box DNA and formed TA and TB gel mobility shift complexes, they may be defective for TBP interactions with other basal transcription factors.

Based on the examples of the TBP interfaces with TFIIA and TFIIA, these two epitopes probably indicate surfaces that interact with proteins yet to be identified. Obvious candidates include TFIID TAFs that are required for activated transcription in mammalian systems in vitro. Because the TFIID TAFs are stably bound to TBP in TFIIID, it is likely that the interaction surface that binds TBP to the TAF complex is larger than the TFIIIB or TFIIA interaction surfaces. Consequently, the TAF complex might interact with both of the remaining, closely spaced epitopes. Consistent with this possibility, Tansey et al. (1994) reported that a triple alanine substitution, R231A, R235A, R239A was reduced for binding to isolated TAF250 in an in vitro binding assay. Further studies will be required to determine whether TBP mutations in these regions interfere with assembly of the mutant TBPs into the TFIID complex in vivo.

**Implications for the significance of other TBP interactions**

Both the VP16 activation domain and E1A CR3 bind to the isolated human TBP protein in vitro (Stringer et al. 1990; Horikoshi et al. 1991; Lee et al. 1991) As discussed in Results, E1A CR3 and VP16 appear to be different types of activation domains. Because they are chemically distinct, it seems unlikely that they would interact with the identical surface of TBP. Consequently, we anticipated that if the activation-domain-TBP interactions observed in vitro contribute to the mechanism of activation in vivo, we would observe some mutations of the TBP surface that would influence E1A and VP16 activation differently, by interfering with the TBP interaction with one of these activation domains and not the other. In an analogous situation, different substitutions in the E. coli RNA polymerase α subunit carboxy-terminal domain have different effects on the transcriptional response to different activators (Busby and Ebright 1994). A striking result of our studies, however, was that the library of TBP mutants responded virtually identically to activation by GAL4–E1A and GAL4–VP16 (Fig. 3). This finding shows that transcriptional activation by both E1A and VP16 requires common interactions with TBP. Although we cannot rule it out, it seems unlikely that the E1A and VP16 activation domains would interact with the identical surface residues of TBP. It seems more likely that the TBP functional epitopes identified in this study are interaction surfaces for a common set of proteins that participate in transcription activated by both activators: TFIIA, TFIIIB, and, as hypothesized above, TFIID TAF complexes. In an earlier analysis of TBP mutants, Tansey and Herr (1995) also concluded that the in vitro interaction between VP16 and TBP does not contribute to the mechanism of VP16 activation in vivo.

Alanine substitution at TBP residue E320 decreased the binding of TFIIA to a TFIIA–TBP–TFIIB–DNA complex [TAB complex] in the presence of Pol II, and alanine
substitutions at E206 and L232 decreased the binding of Pol II to the same TAB complex in the presence of TFIIH [Tang et al. 1996]. Examining the effects of substitutions at these residues on in vivo transcription, we found that E206R and L232E were nearly as active as wt hm3 (Table 1). Consequently, interactions at these residues probably do not contribute greatly to in vivo transcription. Our arginine substitution at E320 produced a TBP molecule that did not bind to DNA in the gel mobility shift assay. Consequently, we cannot draw conclusions about the significance of interactions at this residue.

Fourteen of the 89 mutants we constructed did not form gel mobility shift complexes on TATA-box DNA. As expected from this low DNA-binding activity, none of these mutants were active for in vitro basal transcription (Table 1). Eight of these mutants had very low in vivo activity, as anticipated. However, six of these mutant TBPs that did not bind to a TATA box in vitro had in vivo activities of >20% wt hm3 (Table 1). Tansey et al. [1994] also observed TBP mutants with greatly decreased TAT-box-binding activity that, nonetheless, were active in vivo. These results suggest that compared with basal in vitro transcription, additional interactions occur during in vivo activated transcription that stabilize the incorporation of these mutant TBPs into a preinitiation complex. In related results Arndt et al. [1995] and Lee and Struhl [1995] found that a class of yeast TBP mutants selected to be deficient in activated, but not basal, transcription have mutations that decrease the affinity of TBP for TATA-box DNA. All these results are consistent with the model that activators function in part by promoting assembly of the preinitiation complex [Gaynor and Berk 1983].

Stargell and Struhl [1996] recently reported the characterization of four yeast TBP mutants [F148H, T153I, E236P, and F237D] specifically defective for the ability to support activated transcription in vivo, but wild-type for the ability to support transcription from unregulated promoters and for transcription by Pol I and Pol III. We mutated human TBP residues at three of the positions equivalent to these yeast residues in the TBP core domain. Human TBP mutants F246E and L251E (equivalent to yeast F148 and T153, respectively) had only modest effects on in vivo function in our assays, whereas mutant G334R [equivalent to yeast E236] was fully functional. Our failure to observe significant phenotypes for these human TBP mutations may be because our assays measured transcription from potently activated promoters with four or five binding sites for powerful viral activators, potentially masking the effects of all but the most deleterious TBP mutations. The yeast mutants fell into two classes as determined by whether their defects could be suppressed by recruiting the mutant TBP to a promoter through fusion to a LexA DNA-binding domain. Mutant T153I was fully suppressed and F148H was partially suppressed, whereas the mutations near the carboxyl terminus of TBP were not suppressed by fusion to the LexA DNA-binding domain. The yeast residues T153 and F148 lie near the epitopes defined by our mutations on the upper surface of the first TBP repeat (Fig. 5) and may influence interactions with yeast factors homologous to the human factors that we postulate interact with these epitopes. Because the yeast mutations can be suppressed by recruiting them to a promoter via the LexA DNA-binding domain fusion, the factors that interact with this region of TBP may function by helping to bind wild-type TBP to activated promoters, as suggested by Stargell and Struhl [1996].

Our strategy to identify functionally important regions on TBP’s surface was to substitute wild-type surface residues with amino acids having radically different side chains. Our reasoning was that such mutations would be more likely to severely impair protein–protein interactions and result in significant phenotypes than the frequently used approach of alanine substitution. Our strategy was effective in identifying a surface of TBP that interacts with THIA. In contrast, alanine substitutions of single residues in the TFIIA interaction surface did not significantly alter the affinity of TBP for THIA as measured in gel mobility shift assays [Tang et al. 1996]. This may be because there are a large number of interactions in the TFIIA–TBP interface, many of them with polypeptide backbone atoms that are not altered by amino acid substitutions [Geiger et al. 1996; Tan et al. 1996]. Substitution by alanine eliminates interactions beyond the β-carbon of the side chain substituted, and this apparently did not reduce the energy of binding sufficiently to observe a significant decrease in affinity. In contrast, substitution of a charged residue with a large residue of opposite charge, and substitution of short or hydrophobic side chains with a large charged side chain both eliminates interactions made by the wild-type side chain and may introduce repulsive forces into a protein–protein interface.

In the case of the TBP-TFIIA interaction, the resulting dramatic decreases in affinity were readily detected. In a second contrasting situation, whereas we observed strong phenotypes for individual mutations of basic residues in TBP helix 2 [R231E, R235E, and R239S], Tansey et al. [1994] found that a triple alanine substitution of these same residues in a single mutant [mutant H2, R231A+R235A+R239A] was reduced to only 50% of the activity of wt hm3 in their in vivo assay.

Surprisingly, alternative substitutions in one of the epitopes we defined as being required for Pol II transcription [R231, R235, and F250] specifically affect Pol III transcription in vivo in yeast [mutants in S. cerevisiae TBP K133T, K133S, R137W, R137C, F152G, F152E, Cormack and Struhl 1993]. These mutations in yeast TBP were selected to generate temperature-sensitive growth of yeast cells, and can be suppressed by overexpression of the Pol III TAF, Brl1 [Cormack and Struhl 1993]. It seems likely that the TAFs and general transcription factors specific for each of the three eukaryotic nuclear RNA polymerases evolved from proteins that functioned with a single complex RNA polymerase in a common ancestor of all eukaryotes, and as presently represented by organisms in the kingdom Archaea [Baumann et al. 1995; Langer et al. 1995; Thomm 1996]. The protein–protein interfaces with TBP in these initiation com-
plexes probably were conserved as the distinct Pol I, Pol II, and Pol III systems evolved. Based on this evolutionary argument and the results of Cormack and Struhl (1993) we suggest that residues R231, R235, and F250 contribute to the TBP surface that interacts with a domain of Brf1 in the Pol III factor TFIIIB and an evolutionarily related domain of a Pol II TAF in TFIIID. We suggest that selection of temperature-sensitive TBP mutants in S. cerevisiae yielded substitutions in this interface that weaken the TBP–Brf1 interaction just to the extent that it is broken at the nonpermissive temperature, but not at the lower permissive temperature. We propose that these substitutions are also in the interface with a yeast Pol II TAF, but that they do not weaken the interaction sufficiently to produce a rapid shut-off phenotype for Pol II functions following a shift to the nonpermissive temperature. In contrast, we propose that our (for the most part) more radical substitutions prevent the binding of mammalian Pol II TAFs, and we would predict the binding of the Pol III TAF, Brf1 as well. Further studies are required to test these ideas.

The drawback of radical substitutions is that they may alter the overall conformation of the protein domain being analyzed. For TBP, 14/89 (~15%) of the radical mutations introduced into surface residues inhibited TBP–DNA binding. For mutations in TBP residues that do not interact directly with DNA, we interpret the loss of DNA-binding activity to indicate a significant change in protein conformation. However, if mutants that distort overall protein conformation can be distinguished from those that do not, in this case by assaying TBP DNA-binding activity, then such mutants do not confuse the interpretation of the remaining, more meaningful mutants. Radical mutations in protein surface residues have the potential of introducing repulsive forces into protein–protein interfaces. This strategy may be a generally effective method to probe structure–function relationships of polypeptides incorporated into multiprotein complexes, such as the transcription preinitiation complex, in which eliminating a single side chain’s interactions through alanine substitution may not decrease the stability of the overall complex sufficiently to produce a clear phenotype.

Materials and methods

Mutagenesis of TBP

Altered binding site human TBP [hTBPm3] (Strubin and Struhl 1992) was modified by PCR to include one or two copies of the 9-amino-acid HA1 flu epitope at the amino terminus [hTBPm3e]. This hTBPm3e coding region was cloned into pBluescript (Stratagene), sequenced, and used for oligonucleotide-directed mutagenesis by use of Amersham Sculptor. Mutant plasmids were sequenced and restriction mapped.

Purification of TBP mutants

To facilitate mutant TBP purification, each mutant coding region was cloned into pQE-30 (Qiagen) expression vector so that a 6 His peptide was inserted before the first methionine of human TBP. To accomplish this, the nucleotide sequence encoding the first 184 amino acids of wild-type TBP was generated by PCR to contain a BamHI site at the amino terminus plus SstI and HindIII sites next to the native DraI site of TBP. This fragment was cloned into the BamHI–HindIII site of pQE-30 to generate pQE-30TBPA. The DraI–SstI fragment encoding most of the carboxy-terminal domain of each mutant TBP was then cloned between the DraI and SstI site of pQE-30TBPΔ to regenerate full-length TBP mutants with a 6× His tag. For mutants at amino acid positions 156–184, which are amino-terminal to the DraI site, transfer was accomplished in a two-step cloning strategy. First, a PsI–HindIII fragment from TBP was cloned into the PsI–HindIII site of pQE-30TBPΔ. Then the PsI–PsI fragments of the mutant TBP plasmids were cloned into the PsI site of this vector and the orientation was confirmed by restriction analysis.

Mutant TBP’s were then purified by heparin–Sepharose followed by Ni⁺2-NTA-chromatography. The pQE-30 vectors encoding each mutant TBP were transformed into E. coli strain M15. Bacteria were grown in 500 ml of 2× Tryptone Yeast Extract media (TYE) (DIFCO) to an OD₆₀₀ of ~0.7 at 37°C. Cells are then induced with 1mM isopropyl-β-D-thiogalactoside (IPTG) and grown for 45 min at 30°C. Cells were then pelleted and resuspended in 30 ml of D buffer [20 mM HEPES at pH 7.9, 20% glycerol, 0.2 mM EDTA, 10 mM β-mercaptoethanol, and 0.5 mM phenylmethylsulfonyl fluoride (PMSF)] containing 100 mM KC1 (D₁₀₀). Cells were sonicated, debris was removed by centrifugation, and His-tagged TBP mutants were purified from the soluble extract by binding the supernatant in batch to heparin–Sepharose (Pharmacia; 1 ml of heparin per 5–10 mg of protein) at 4°C for 1 hr. The resin was washed once with D₁₀₀ and loaded into a column that was washed with D buffer containing 300 mM KC1 (D₃₀₀) and eluted with D buffer containing 1 M KC1. The eluted protein was diluted with D buffer to a final concentration of 300 mM KC1. This protein fraction was then bound in batch for 1 hr at 4°C to 1 ml of Ni⁺²-NTA-Agarose (Qiagen) that was pre-equilibrated in D₃₀₀. The beads were then washed once with D₃₀₀ and loaded into a column that was washed with D₃₀₀ containing 5 mM imidazole. Finally, the columns were eluted with 400 μl D₃₀₀ containing 100 mM imidazole. The purified protein was then dialyzed for 4 hr at 4°C in D₃₀₀. Protein concentrations were determined by Bradford assay. Equivalent masses of different mutant TBP preparations were used in gel mobility shift assays and in vitro transcription assays as indicated below.

Gel mobility shift assays

Gel mobility shift assays contained 5 ng of mutant TBP or wt hTBPm3, 0.4 ng of [³²P]labeled double-stranded DNA (37 mer) with the adenovirus 2 major late promoter TATA box and, where indicated, 15 ng of recombinant TFIIIA or/and 15 ng of recombinant TFIIIB in 12.5 μl 40 mM KC1, 5 mM MgCl₂, 10 mM HEPES at pH 7.9, 10 mM β-mercaptoethanol, 20 μg/ml poly[d(G-C)], 500 μg/ml BSA, 10% glycerol, 0.1 mM EDTA. Reactions were incubated for 30 min at 37°C and products were resolved on a 5% polyacrylamide gel in 45 mM Tris, 45 mM boric acid at pH 8.3, and 1 mM EDTA.

In vivo function assays

In vivo function of each TBP mutant was assayed during transient transfection of COS cells.

TGTTAA-box reporter plasmids

The reporter derived from the β-retinoic acid receptor promoter region was pGal4-M1-Luc (Keveaney et al. 1993). The second reporter plasmid, p4xGal-fosTGTLuc, was derived from
Bryant et al.
c-fos (−56) [4xGAL] (Tansey et al. 1994) by replacing the CAT gene with luciferase.

**Mutant TBP mammalian expression vector**

Wt hm3 and each mutant TBP were expressed by use of the pSreMSVtkneo vector (Muller et al. 1991) that replicates from c-los (-

**pSR (~MSVtkneo vector** (Analytical Luminescence Laboratory) with 5 sec measurements with the Promega Luciferase Assay Substrate. β-Galactosidase was assayed by use of ONPG hydrolysis. Luciferase assays were normalized by dividing luciferase units by β-galactosidase units. The background control was from a transfection in which the TBP expression vector was replaced by the same vector expressing an inactive, truncated form of hm3 (deleted at the Stud site). Reported units are normalized luciferase units minus background as a percentage of the activity observed with wt hm3.

**Transient transfection assays**

Three micrograms of mutant TBP expression vector, 5 µg of reporter plasmid (p4xGALC-fos-TGTLuc or pGAL4-M1-Luc), 1 µg of pGAL4-E1A or 0.1 µg of pSGVP16, 1 µg of pCH110 (Hall et al. 1983) was assayed to normalize transient transfection assays.

**In vitro basal transcription**

For in vitro basal transcription, 20 ng of mutant TBP (or wt hm3) and 300 ng of yeast GAL4 1–147 fused to Ad2 E1A 121–223 (conserved region 3) from the SV40 early promoter. GAL4–VP16 was expressed from the SV40 early promoter by use of pSGVP16 (Sadowski et al. 1988). β-galactosidase expressed from pCH110 (Hall et al. 1983) was assayed to normalize transient transcription assays.

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