Redundant Cooperative Interactions for Assembly of a Human U6 Transcription Initiation Complex

Beicong Ma and Nouria Hernandez*

Howard Hughes Medical Institute, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York 11724

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The core human U6 promoter consists of a proximal sequence element (PSE) located upstream of a TATA box. The PSE is recognized by the snRNA-activating protein complex (SNAPc), which consists of five types of subunits, SNAP190, SNAP50, SNAP45, SNAP43, and SNAP19. The TATA box is recognized by TBP, which together recruit TBP to the U6 transcription initiation complex. The hnRNP A1 protein interacts with TBP to form the core U6 promoter complex, which is located about 200 bp upstream of the transcription start site. The PSE recruits the snRNA-activating protein complex (SNAPc), which contains an unusual Myb domain with a half Myb repeat (Rh) followed by four full-length Myb repeats (Ra, Rb, Rc, and Rd), and is recognized by TBP in the context of mini-SNAPc. The SNAPc complex is then assembled on the core promoter through cooperative interactions with Brf2. Our results identify complexes smaller than mini-SNAPc that are transcriptionally active and show that there are at least two redundant mechanisms to stably recruit TBP to the U6 transcription initiation complex.
a built-in damper that down-regulates its binding to DNA and that is deleted in mini-SNAP$_c$ (19).

In addition to the interaction between the activator Oct-1 and SNAP$_c$, protein-protein contacts are likely to occur among core promoter binding factors, because they also display cooperative binding. For example, TBP and Brf2 bind cooperatively to a TATA box (1), and SNAP$_c$ and TBP bind cooperatively to probes containing both a PSE and a TATA box (18). We have shown that mini-SNAP$_c$, although unable to bind cooperatively with the Oct-1 POU domain, can bind cooperatively with TBP (19). Because mini-SNAP$_c$ binds relatively efficiently to DNA on its own, cooperative binding in this case results, in effect, in recruitment of TBP to the DNA (19).

Here we have assembled complexes that were smaller than mini-SNAP$_c$ (stmSNAP$_c$) and tested their ability to recruit TBP to the DNA. The results identify a 50-amino-acid (aa) region within the N-terminal part of SNAP190 that is required for cooperative binding of mini-SNAP$_c$ with TBP. When fused to the ReRd repeats of SNAP190 or the GAL4 DNA binding domain, this region is sufficient to recruit TBP to a TATA box. By using stmSNAP$_c$s lacking the TBP-recruiting region, we show that there are at least two redundant mechanisms to recruit TBP to U6 transcription initiation complexes. Our results also identify a stmSNAP$_c$ that is 192 aa smaller than mini-SNAP$_c$ and yet is still capable of directing basal RNA polymerase III transcription.

FIG. 1. Composition of various SNAP$_c$ subcomplexes. (A) Map of the known subunit-subunit interactions within SNAP$_c$. Within SNAP190, aa 84 to 133 associate with SNAP19 and aa 164 to 268 of SNAP43, aa 263 to 503 correspond to the Myb domain with the Rh, Ra, Rb, Rc, and Rd repeats, aa 869 to 912 correspond to the Oct-1-interacting region (OIR), and aa 1281 to 1393 associate with SNAP45. Within SNAP43, aa 1 to 164 associate with SNAP50. Mini-SNAP$_c$ contains SNAP190 aa 1 to 514, SNAP43, and SNAP50, as outlined in red. (B) The composition of each stmSNAP$_c$ is indicated. All stmSNAP$_c$s including stmSNAP$_c$#23-26 contain a SNAP190 protein truncated after aa 505.
FIG. 2. A 50-aa region of SNAP190 is required for cooperative binding of the complex with TBP. (A) EMSA performed with the proteins indicated above the lanes and a probe containing the wild-type mouse U6 PSE and either the wild-type or a mutated human U6 TATA box as indicated above the lanes. In panels A to C, the upper complex in the doublet observed with stmSNAP_c#1 (#1) is labeled with an asterisk to the left of the complex, and the upper complex of the doublet observed with stmSNAP_c#1 and TBP is labeled with an arrowhead to the left of the complex. α-190, anti-SNAP190 antibody; −, only complex #1 was added. (B) EMSA performed with the proteins indicated above the lanes and a probe containing the wild-type mouse U6 PSE and the human U6 TATA box. The absence (−) or presence of the stmSNAP_c proteins (indicated by the stmSNAP_c number) is indicated over the lanes. The absence (−) and amount (indicated by the height of the triangle) of TBP is indicated over the gel. TBP was titrated over a threefold range. The protein-DNA complexes containing the stmSNAP_c only or both the stmSNAP_c and TBP are indicated. (C) EMSA performed with the proteins indicated above the lanes and a probe containing a wild-type mouse U6 PSE and a mutated TATA box. The presence or absence (−) and the amount (indicated by the height of the triangle over the lanes) of TBP are indicated over the lanes.
FIG. 3. When fused to the SNAP190 RcRd Myb repeats, SNAP190 aa 1 to 90 are sufficient to recruit TBP to a TATA box. (A) The structures of the various fusion proteins are illustrated. HT, His tag; G6, a run of six glycines. (B) EMSA performed with probes containing either wild-type or mutated mouse U6 PSE and wild-type or mutated human U6 TATA box, as indicated above the panels, and the proteins indicated above the lanes. The absence (−) or presence (+) of TBP is indicated over the gel. TBP was titrated over a threefold range. The proteins present in the various protein-DNA complexes are indicated to the left of the panels. (C) EMSA performed with the probes and proteins indicated above the lanes. The proteins present in the protein-DNA complex are indicated to the left.
**MATERIALS AND METHODS**

Protein expression and purification. The various versions of the SNAP190, SNAP43, and SNAP50 proteins were expressed in *Escherichia coli* with the T7 system of Studier and colleagues (27). In some cases, both SNAP43 and SNAP50 were expressed from a single plasmid. All SNAP190 derivatives were tagged at their C terminus with six histidines. The lysates from cells expressing SNAP43, SNAP50, and SNAP190 were then mixed and incubated with rotation for 2 h at 4°C, and the SNAP complexes were purified by affinity chromatography on nickel-nitrotriacetic acid (Ni-NTA) agarose beads (Qiagen). TBP and the RacRd portion of SNAP190 were expressed as glutathione S-transferase (GST) fusion proteins in *E. coli* and purified by affinity chromatography on glutathione agarose beads. The proteins were then either eluted with reduced glutathione (G-TBP) or cleaved from the GST moiety with thrombin. Brf2 carrying a His tag at its C terminus was expressed in *E. coli* and purified by affinity chromatography on Ni-NTA agarose beads. The GAL4 DNA binding domain (GAL4 DBD-HT) and the (1-90)-GAL4 DNA binding domain fusion protein [(1-90)GAL4 DBD-HT), both carrying a His tag at their C termini, were expressed in *E. coli* with the T7 system and purified by affinity chromatography on Ni-NTA agarose beads.

Electrophoretic mobility shift assays (EMSSAs). The binding reactions were performed in a total volume of 20 μl containing 80 mM KCl, 20 mM HEPEs (pH 7.9), 5 mM MgCl₂, 0.2 mM EDTA, 10% glycerol, 1 mM dithiothreitol (DTT), and were complemented with 12.5 ng of recombinant TBP and 2 μl of fetal bovine serum. The reaction mixtures were incubated for 20 min at 4°C before addition of the radiolabeled probe, followed by a 30-min incubation at 30°C. The binding reactions were fractionated on a 5% polyacrylamide gel (39:1 acrylamide-bisacrylamide) in TGE buffer (50 mM Tris base, 380 mM glycine, 0.07% Tween 20, 0.5 μg of poly(dG-dC)·(dG-dC), 0.25 μg of pUC118, and 20 μg of bovine serum albumin. The reaction mixtures were incubated for 20 min at 4°C before addition of the radiolabeled probe, followed by a 30-min incubation at 30°C.

**RESULTS**

The SNAP190 region from aa 34 to 83 is required for cooperative binding with TBP in the context of mini-SNAP<sub>E</sub>. To determine which region of SNAP<sub>E</sub> might be required for cooperative binding with TBP, we generated subcomplexes of SNAP<sub>E</sub> that can bind to the PSE on their own and monitored their ability to recruit TBP in gel shift assays. Figure 1 shows a map of subunit-subunit interactions within the full-length SNAP<sub>E</sub> that we established previously (13). Mini-SNAP<sub>E</sub>, which efficiently recruits TBP, consists of SNAP50, SNAP43, and the N-terminal 514 aa of SNAP190 including the SNAP190 Myb domain (19). As shown in Fig. 1A, within the mini-SNAP<sub>E</sub> subunits, neither the first 83 aa of SNAP190 nor the SNAP43 region extending from aa 269 to 368 are required for subunit-subunit interactions. Indeed, we previously showed that stm-SNAP<sub>s</sub> that bind to the PSE can be assembled (13). We assembled a series of such complexes, which are listed in Fig. 1B. StmSNAP<sub>S</sub>1 is identical to mini-SNAP<sub>E</sub> except that the SNAP190 fragment present is slightly shorter at the C terminus, ending after aa 505. All other stm-SNAP<sub>s</sub> contain a SNAP190 truncated after aa 505, and in addition stm-

![FIG. 4. The complex observed in the presence of TBP and the (1-90)RcRd-HT fusion protein contains both TBP and (1-90)RcRd-HT. EMSA performed with the proteins indicated above the lanes and a probe containing the wild-type mouse U6 PSE and the human U6 TATA box. G(1-90)RcRd and GTBP are GST fusion proteins. The proteins present in each protein-DNA complex are indicated to the right. In lane 1 no proteins were added to the probe.](http://mcb.asm.org/)
cause they were supershifted by addition of an anti-SNAP190 antibody (lane 5), and TBP, because their migration was retarded when TBP was replaced by a larger GST-TBP fusion protein (lane 6). As expected, none of the TBP-containing complexes were observed on probes containing a mutated TATA box (lanes 8 to 17). However, the complexes containing just stmSNAPc#1 were still formed on these probes, as expected (lanes 13 to 17) and could be supershifted by addition of the anti-SNAP190 antibody (lane 15). Thus, on probes containing both a PSE and a TATA box, TBP can be recruited to the TATA box by stmSNAPc#1.

To compare the abilities of the different stmSNAP's to recruit TBP, we first performed a titration experiment to determine the amount of each complex required to obtain equal binding to the high-affinity mouse U6 PSE. As shown in Fig. 2B, we were able to titrate each complex so as to obtain roughly equal amounts of protein-DNA complex (compare lanes 4, 7, 10, 13, 16, 19, 22, 25, 28, 31, 34, and 37). (Note that in this case, the stmSNAP's containing full-length SNAP43, such as stmSNAPc#1, gave rise only to the lower complex of the doublet observed in Fig. 2A.) Little or no detectable binding was obtained with increasing amounts of full-length TBP.

FIG. 5. SNAP190 aa 1 to 90 can recruit TBP to the DNA when fused to a heterologous DNA binding domain. (A) The structures of the various fusion proteins are illustrated. HT, His Tag. (B) EMSA performed with probes containing three tandem copies of the G17 GAL4 DNA binding site (3 X G17) (3) inserted 5 bp upstream of the wild-type (lanes 1 to 14) or mutated (lanes 15 to 20) HIV-1 TATA box (see Materials and Methods for the sequences of the probes), and the proteins indicated above the lanes. The presence (+) or absence (−) of TBP and the amounts (indicated by the thickness of the triangle) of the proteins containing the GAL4 DNA binding domain (DBD) are shown over the gel. DBD, GAL4 DBD-HT; (1-90)DBD, (1-90)GAL4 DBD-HT (see panel A). Both the GAL4 DBD-HT and (1-90)GAL4 DBD-HT proteins were titrated over a threefold range. In lanes 17 and 18, the same amount of GAL4 DBD-HT was used as in lane 5. In lanes 19 and 20, the same amount of (1-90)GAL4 DBD-HT was used as in lane 11. The proteins present in the various protein-DNA complexes are indicated to the left of the panels.
on its own (lanes 2 and 3). In sharp contrast, when both stmSNAPc#1 and TBP were incubated with the probe, the complex containing both stmSNAP c#1 and TBP appeared, as before (complex labeled stmSNAP cs/H11001 TBP [lanes 5 and 6]). (Note that at high concentrations of TBP, the upper band of the doublet appeared [lane 6, see also lanes 12, 18, 30, and 36, complex labeled with an asterisk], and as before, this complex was supershifted similarly to the corresponding main stmSNAPc-PSE complex with TBP [see lanes 6, 12, 18, 30, and 36, complex labeled with an arrowhead]).

Deletion of the N-terminal 32 aa of SNAP190 had no effect on TBP recruitment (stmSNAPc#3; Fig. 2B, lanes 11 and 12); however, deletion of the first 62 aa weakened TBP recruitment (stmSNAPc#5; lanes 17 and 18), and deletion of the first 83 aa resulted in a barely detectable stmSNAPc-TBP complex (stmSNAPc#7; lanes 23 and 24). This suggested that a region required for cooperative binding with TBP lay between SNAP190 aa 63 (or slightly N-terminal of aa 63) and aa 84. Surprisingly, however, an internal deletion encompassing aa 50 to 83 of SNAP190 had little effect (stmSNAPc#25; lanes 35 and 36), and only a larger internal deletion from aa 34 to 83 reduced cooperative binding significantly (stmSNAPc#23; lanes 29 and 30). Combination of the SNAP190 deletions with deletion of the last 100 aa of SNAP43 had little additional effect on cooperative binding with TBP (compare each odd-numbered complex with the next even-numbered complex). Together, these results indicate that in the context of mini-SNAPc, the SNAP190 region extending from aa 34 to 83 contains redundant sequences required for cooperative binding with TBP.

When we tested whether cooperative binding is dependent on a wild-type PSE and TATA box, we did not detect binding of the stmSNAPc to probes containing a mutant PSE, consistent with our previous results (13), even upon addition of TBP (data not shown). The results with a probe containing a wild-type mouse U6 PSE and a mutant TATA box for all of the stmSNAPc are shown in Fig. 2C. As before for stmSNAPc#1 (Fig. 2A), we did not observe any of the stmSNAPc-TBP complexes in the absence of the TATA box, indicating that cooperative binding is dependent on both an intact PSE and an intact TATA box, as observed previously (18). However, for all stmSNAPc containing full-length SNAP43, we did observe formation of the upper complex of the doublet upon addition of TBP (Fig. 2C, lanes 4, 8, 12, 16, 20, and 24, complex labeled with an asterisk). As shown above, the mobility of this complex was not affected by replacing TBP with a larger GST-TBP fusion protein or with Brf2 (Fig. 2A, compare lanes 16 and 17), again suggesting that it corresponds to another conformation of the stmSNAPc.

The N-terminal 90 aa of SNAP190 are sufficient for cooperative binding with TBP when fused to the SNAP190 RcRd Myb repeats. The above results suggest that in a complex consisting of SNAP43 (or the first 268 aa of SNAP43), SNAP50, and the N-terminal third of SNAP190, the N terminus of SNAP190 is required for cooperative binding with TBP. To determine whether this SNAP190 region might recruit TBP in the absence of other SNAPc subunits, we fused it via a six-glycine linker to either the N [(1-90)RcRd] or C [RcRd(1-90)] terminus of the RcRd repeats of SNAP190 (Fig. 3A). The SNAP190 RcRd repeats can bind to the PSE on their own, as visualized with a gel lacking MgCl2, although with reduced specificity (32; M. Wong and N. Hernandez, unpublished results). As illus-

FIG. 6. The transcription activities of different stmSNAPc do not correlate with their ability to recruit TBP. A whole-cell extract was either left untreated (lanes 1 and 2) or treated with preimmune (PRE-IMM.) antibody beads (lanes 3 and 4) or with a mixture of anti-SNAP190 (α-SNAP190) and anti-SNAP43 (α-SNAP43) antibody beads (lanes 5 to 20). The extracts were then complemented with the proteins indicated above the lanes. The presence (+) or absence (−) of TBP and the amount of the stmSNAPc protein (indicated by the height of the triangle) are indicated over the lanes. The extracts were programmed with the plasmid pU6/Hae/RA.2, which carries the human U6 promoter. The titrations of the stmSNAPc and the (1-90)RcRd protein were over a threefold range. The transcripts were analyzed by RNase T1 protection. The signals corresponding to correctly initiated U6 RNA (U6) or to a control RNA (IC) included in the reaction mixtures to monitor RNA handling and recovery are indicated to the left of the gel.
FIG. 7. TBP can be recruited to the U6 transcription initiation complex in the absence of SNAP190 aa 1 to 83. EMSA performed with the proteins indicated above the lanes and a probe containing the wild-type mouse U6 PSE and the human U6 TATA box. The amounts (indicated by the height of the triangle) of an anti-SNAP190 antibody (α-190) (CS696) and G-TBP, a GST-TBP fusion protein, are indicated over the lanes.
treated in Fig. 3A, we also generated a protein consisting of just the N-terminal 90 aa of SNAP190. All the constructs carried a His tag (HT) for easy purification from E. coli lysates.

The results are shown in Fig. 3B. As before, stmSNAPc#1, but not stmSNAPc#7, efficiently recruited TBP to the DNA (lanes 1 to 8), and this effect was dependent on both an intact TATA box (second panel) and an intact PSE (third panel). Neither the (1-90)RcRd nor the RcRd (1-90) fusion proteins bound to DNA on their own (lanes 9 and 12). However, upon addition of TBP, increasing amounts of both fusion proteins gave rise to increasing amounts of a protein-DNA complex [lanes 10, 11, 13, and 14, complexes labeled (1-90)RcRd or RcRd(1-90) + TBP]. These complexes were dependent on an intact TATA box (second panel) but surprisingly not on an intact PSE (third panel). We then checked whether the SNAP190 region from aa 1 to 90 was required for this effect. As shown in Fig. 3C, neither the (1-90)RcRd protein nor the RcRd protein bound efficiently to DNA on their own (lanes 2 and 3), although it should be noted that the same preparation of RcRd protein bound to DNA, as visualized with a gel lacking MgCl₂, consistent with our previous results (data not shown) (32). Importantly, in the presence of TBP, the (1-90)RcRd fusion protein, but not the RcRd protein, gave a prominent complex (compare lane 5 to lane 6), and this complex was dependent, as before, on an intact TATA box (lane 11) but not on an intact PSE (lane 17). Thus, the prominent complex is dependent on the presence of the first 90 aa of SNAP190 in the fusion protein.

Since the prominent complex observed with TBP and the (1-90)RcRd fusion protein did not depend on an intact PSE, we wondered whether it indeed contained both TBP and the SNAP190-derived fusion proteins. As shown in Fig. 4, (1-90)RcRd and TBP each bind poorly to the probe alone, as did GST fusion derivatives of each protein [G(1-90)RcRd and GTBP, lanes 2 to 5]. When (1-90)RcRd was combined with TBP, however, a protein-DNA complex was observed, as before (lane 6). The complex migrated with increasingly slower mobility upon replacement of (i) (1-90)RcRd with the larger protein G(1-90)RcRd (lane 7), (ii) TBP with the larger protein GTBP (lane 8), and (iii) both (1-90)RcRd and TBP with the larger GST fusion proteins (lane 9). It is not clear why fusion of the GST moiety to the (1-90)RcRd protein affected migration of the complex less than fusion of the GST moiety to TBP (lanes 7 and 8). Nevertheless, the observation that fusion of the GST moiety to either protein does retard migration of the complex confirms that the complex does indeed contain both components. Together, these results suggest that a fusion protein consisting of the first 90 aa of SNAP190 and the RcRd DNA binding domain is capable of recruiting TBP to the DNA. The observation that this recruitment is not dependent on an intact PSE suggests that the (1-90)RcRd and RcRd(1-90) proteins bind with low specificity to DNA.

The N-terminal 90 aa of SNAP190 are sufficient for cooperative binding with TBP when fused to the GAL4 DNA binding domain. We also expressed two proteins, shown in Fig. 5A, consisting of either the GAL4 DNA binding domain alone (GAL4 DBD-HT, aa 2 to 94 of GAL4) or the first 90 aa of SNAP190 fused to the GAL4 DNA binding domain [(1-90)GAL4 DBD-HT] with, in each case, a His tag (HT) at the C terminus. We tested the abilities of these two proteins to recruit TBP to probes carrying three tandem GAL4 DNA binding sites (G17 site [3]) upstream of either a wild-type or mutated TATA box derived from the human immunodeficiency virus type 1 (HIV-1) promoter (see Materials and Methods for the sequences of these probe). The presence of three GAL4 binding sites allowed us, in effect, to test three different spacings between the GAL4 site and the TATA box in a single probe. As shown in Fig. 5B, TBP on its own bound very weakly to the probe, as expected, such that the complex is not visible at this exposure of the gel (lane 2). In contrast, both the GAL4 DBD-HT and the (1-90)GAL4 DBD-HT proteins [labeled DBD and (1-90)DBD, respectively, in Fig. 5B] bound to DNA on their own (lanes 3 to 5 and 9 to 11). The GAL DBD-HT protein formed at least two complexes even at the lowest concentration used (lane 3, complexes labeled 3xDBD and 2xDBD), while the (1-90)GAL4 DBD protein formed at least two complexes only at the highest concentration used [lane 11, complexes labeled 3x (1-90)DBD and 2x (1-90)DBD]. These complexes probably correspond to occupancy of two and three of the three GAL4 DNA binding sites on the probe (a weak, smaller complex probably corresponding to occupancy of a single site can be seen just above the probe signal on low exposures of the gel). Very strikingly, in the presence of TBP, a new, prominent complex was obtained with the (1-90)GAL4 DBD-HT but not the DBD-HT protein [compare lanes 12 to 14 with lanes 6 to 8, complex labeled (1-90)DBD + TBP]. This complex migrated more slowly than either the TBP-TATA box complex (which is visible in lane 2 on very long exposures of the gel) or the 3x (1-90)DBD complex (lane 11), was dependent on the presence of a wild-type TATA box (compare lanes 12 to 14 with lane 20), and probably corresponds, therefore, to a complex containing both TBP and the (1-90)GAL4 DBD-HT protein. Thus, the first 90 aa of SNAP190 are capable of recruiting TBP to the DNA when fused to at least two types of DNA binding domain, the RcRd repeats from SNAP190 or the DNA binding domain of the Saccharomyces cerevisiae GAL4 protein. This indicates that the first 90 aa of SNAP190 are sufficient to recruit TBP to a TATA box when fused to a heterologous DNA binding domain.

The symbols to the right of different protein-DNA complexes indicate complexes containing proteins as follows: white circles, TBP and Brf2; blue circles, G-TBP and Brf2; white triangles, stmSNAPc#1 and TBP; blue triangles, stmSNAPc#1 and G-TBP; red triangle, stmSNAPc#1, TBP, and anti-SNAP190; white diamonds, stmSNAPc#1, TBP, and Brf2; blue diamonds, stmSNAPc#1, G-TBP, and Brf2; red diamonds, stmSNAPc#1, TBP, Brf2, and anti-SNAP190. (B) EMSA performed with probes containing the wild-type mouse U6 PSE and either a wild-type or mutant human U6 TATA box as indicated above the lanes, and the proteins indicated above the lanes. The Brf2 titrations were over a threefold range. The symbols at the right of different protein-DNA complexes indicate complexes containing proteins as follows: white circles, TBP and Brf2; white triangles, stmSNAPc#1 and TBP; white diamonds, stmSNAPc#1 or stmSNAPc#8, TBP, and Brf2.
FIG. 8. Known functional regions of SNAP190. The known functional regions of SNAP190 follow: SNAP190 aa 34 to 83, TBP recruitment region 1 (TRR1); SNAP190 aa 84 to 138, SNAP19- and SNAP43-interacting region; SNAP190 aa 263 to 503, Myb domain; SNAP190 aa 869 to 912, Oct-1-interacting region (OIR); and SNAP190 aa 1281 to 1393, SNAP45-interacting region.

The stmSNAP_s are competent for U6 transcription. We depleted a transcription extract of endogenous SNAP_c with a mixture of anti-SNAP190 and anti-SNAP43 antibodies and tested the abilities of the various stmSNAP_s to restore transcription in the depleted extract. As shown in Fig. 6, addition of TBP to the untreated extract stimulated transcription, indicating that TBP was limiting (lane 2). Upon treatment of the extract with preimmune antibody beads, transcription was diminished but less so than upon depletion with anti-SNAP190 and anti-SNAP43 beads (compare lanes 3 and 5). Upon addition of TBP, transcription from the extract treated with preimmune beads was restored to near-wild-type levels, whereas transcription from the extract depleted with anti-SNAP190 and anti-SNAP43 beads was stimulated only slightly (compare lanes 4 and 6). To test the stmSNAP_s, we complemented the depleted extract with a constant amount of TBP and increasing amounts of the various stmSNAP_s. The amounts of the various stmSNAP_s were normalized as in Fig. 2B to give rise to equal binding to the mouse U6 PSE. As shown in Fig. 6, lanes 7 to 18, the various stmSNAP_s restored transcription to different levels, but there was no correlation between the ability of an stmSNAP_c to restore transcription and its ability to recruit TBP. Thus, stmSNAP_c#7, stmSNAP_c#8, stmSNAP_c#23, and stmSNAP_c#24, which cannot recruit TBP efficiently (Fig. 2), were as active or more active than stmSNAP_c#1 and stmSNAP_c#2, which recruit TBP efficiently. Addition of the (1-90)RcRd protein did not stimulate transcription above the levels observed with addition of just TBP, indicating that this protein is inactive for transcription (compare lanes 19 and 20 to lane 6). Together, these results indicate that the ability of the stmSNAP_s to recruit TBP to the TATA box is not essential for transcription. Further, they show that stmSNAP_c#8, which contains SNAP190 aa 84 to 505, SNAP43 aa 1 to 268, and SNAP50, contains all the information required to direct basal levels of RNA polymerase III transcription. On the other hand, the (1-90)RcRd derivative of its own is not capable of directing transcription, indicating that it is lacking essential parts present in stmSNAP_c#8.

TBP can be recruited to the DNA through cooperative binding with Brf2. Cooperative binding of SNAP_c and TBP to the U6 promoter achieves both recruitment of TBP and recruitment of SNAP_c. The stmSNAP_s, however, lack the damper of DNA binding present in the full-length SNAP_c. In our experiments, we adjusted their concentrations so as to obtain in each case efficient binding to the PSE. Thus, under these conditions, cooperative binding mainly serves to recruit TBP. The observation that stmSNAP_s unable to recruit TBP efficiently are nevertheless active for transcription suggests, then, that there is another route to recruit TBP to the U6 promoter. Indeed, Brf2 has been shown to bind cooperatively with TBP (1). We asked, therefore, whether Brf2 could recruit TBP to the U6 promoter in the presence of stmSNAP_s. The results are shown in Fig. 7.

We first characterized the complexes obtained with various combinations of Brf2, stmSNAP_c#1, and TBP, as shown in Fig. 7A. As before, TBP alone did not bind efficiently to the probe, nor did a GST-TBP fusion protein, Brf2, and an antibody directed against SNAP190 (lanes 2 to 5). StmSNAP_c#1 alone bound to the probe (lane 6) and was able to recruit TBP, as expected (lane 7, complex labeled with a white triangle). The stmSNAP_c#1-TBP complex was retarded in its migration by an antibody directed against SNAP190 (lane 8, red triangle) or by replacement of TBP by the larger GST-TBP fusion protein (lane 9, blue triangle), consistent with it containing both stmSNAP_c#1 and TBP. Brf2 and TBP, each of which alone binds very poorly, if at all, to the probe (lanes 3 and 4) gave a strong complex when added together (lane 10, white circle). This complex migrated more slowly when TBP was replaced with the larger GST-TBP fusion protein (lane 11, blue circle), indicating that it does indeed contain TBP. Thus, as observed before (1), TBP and Brf2 bind cooperatively.

When Brf2 was added to the stmSNAP_c#1-TBP complex, a new complex of lower mobility was obtained (Fig. 7A, lane 13, white diamond). The mobility of this complex was retarded by increasing amounts of anti-SNAP190 antibody (lanes 14 and 15, red diamond) and by replacement of TBP with the larger GST-TBP fusion protein (lanes 18 and 19, blue diamond), consistent with it containing both stmSNAP_c#1 and TBP. Moreover, complexes formed in the presence of Brf2 consistently migrated slightly more slowly than the corresponding complexes lacking Brf2, confirming that these complexes indeed contain Brf2 [compare (i) the stmSNAP_c#1-TBP-Brf2 complex in lane 13 (white diamond) with the stmSNAP_c#1-TBP complex in lane 12 (white triangle); (ii) the stmSNAP_c#1-TBP-Brf2 complex in lane 14 and 15 (red diamond) to the stmSNAP_c#1-TBP complex in lane 14 and 15 (red triangle); (iii) the stmSNAP_c#1-G-TBP-Brf2 complex in lanes 18 and 19 (blue diamond) to the stmSNAP_c#1-G-TBP complex in lane 16 (blue triangle)]. Together, these results indicate that TBP binds cooperatively not only with stmSNAP_c#1 but also with Brf2. Further, a complex containing all three components is efficiently obtained.

We next compared stmSNAP_c#1, which can recruit TBP, with stmSNAP_c#8, which is deficient in TBP recruitment, for assembly into complexes containing TBP and Brf2. As shown in Fig. 7B, neither TBP nor Brf2 bound efficiently to DNA on their own, but together, they formed a strong complex, as before (lane 4, white circle). Both stmSNAP_c#1 and stm-
SNAPc#8 bound efficiently to the mouse U6 PSE (lanes 5 and 11) but only stmSNAPc#1 recruited TBP to the DNA (compare lanes 6 and 12, white triangle). Furthermore, we did not observe cooperative binding of Brf2 and any of the two stmSNAPc#s (lanes 10 and 16). Upon addition of increasing amounts of Brf2, increasing amounts of the stmSNAPc#1-TBP-Brf2 complex were obtained (lanes 7 to 9, white diamond). Remarkably, addition of Brf2 to the binding reaction mixture containing stmSNAPc#8 and TBP also resulted in efficient formation of a complex containing the three components, indicating that TBP can be recruited through Brf2 (lanes 13 to 15, white diamond). In addition, a complex containing just TBP and Brf2 was observed in the presence of both stmSNAPc#s (for example, lane 15, white circle), although with stmSNAPc#1 the complex was difficult to visualize because it comigrated with the complex that probably corresponds to an altered conformation of stmSNAPc#1 (compare lane 9, white circle, with the binding reaction mixture lacking Brf2 in lane 6). All the complexes containing TBP and Brf2 were dependent on the presence of a TATA box in the probe (lanes 18 to 28).

**DISCUSSION**

We have shown before that TBP and SNAPc bind cooperatively to the U6 promoter. Here we have identified a region in mini-SNAPc that is required for recruitment of TBP. Thus, as shown in Fig. 8, we now know of five functional regions within SNAP190. Two of these are involved in association with other SNAPc subunits; the region extending from aa 84 to 133 associates with SNAP43 and SNAP19, and the region extending from aa 1281 to 1393 associates with SNAP45 (13). The Myb domain extends from aa 263 to 503 and contains the two Myb repeats Rc and Rd that are required for DNA binding (19, 32). Two other regions are involved in cooperative binding with other members of the U6 initiation complex: (i) the Oct-1-interacting region (OIR) from aa 869 to 912, which allows cooperative binding with Oct-1 through direct protein-protein contacts (2, 17); and (ii) the TBP recruitment region (TRR1) from aa 34 to 83, which is required for cooperative binding with TBP in the context of mini-SNAPc. The TRR1 is also sufficient to recruit TBP when fused to a heterologous DNA binding domain.

With the full-length SNAPc and with probes containing the low-affinity human U6 PSE, cooperative binding of SNAPc and TBP effectively results in recruitment of both proteins to the DNA. In such a context, the cooperative interaction between the two proteins is necessary for efficient transcription from the human U6 core promoter (18). With mini-SNAPc on probes containing the high-affinity mouse U6 PSE, cooperative binding results mainly in the recruitment of TBP to the DNA, since it can be recruited by cooperative binding with TBP or with Oct-1. Such redundant mechanisms may ensure efficient formation of the transcription initiation complex on chromatin templates in vivo. Furthermore, they probably ensure that once the U6 transcription initiation complex is formed, it remains stable for several rounds of transcription.

It will be very interesting to determine whether SNAPc#s that still contain the damper of DNA binding but are unable to bind cooperatively with TBP are inactive for transcription from the core U6 promoter lacking the Oct-1 binding site. We do not know, however, whether deletion of the SNAP190 TRR1 will be sufficient to debilitate recruitment of TBP by the full-length SNAPc. Indeed, although this region is required for efficient TBP recruitment in the context of stmSNAPc#1 (and is sufficient to recruit TBP when fused to the RcRd repeats), it is quite possible that the full-length complex contains other regions, either in the C-terminal two thirds of SNAP190 or in SNAP45 or in SNAP19, that act redundantly with the SNAP190 TRR1 region.

The smallest stmSNAPc we tested in transcription is stmSNAPc#8, which lacks the last 100 aa of SNAP43 as well as the first 83 aa and the last 9 aa of the SNAP190 truncation present in mini-SNAPc. Remarkably, this complex is active for basal RNA polymerase III transcription from the human U6 promoter, indicating that it contains all the information required to recruit, ultimately, RNA polymerase III. We do not know whether RNA polymerase III directly contacts SNAPc, but our results suggest that any essential contact would involve subunits or part of subunits present in stmSNAPc#8, the smallest functional SNAPc subcomplex we have assembled so far.

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