Events during Initiation of Archaeal Transcription: Open Complex Formation and DNA-Protein Interactions

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Transcription in Archaea is initiated by association of a TATA box binding protein (TBP) with a TATA box. This interaction is stabilized by the binding of the transcription factor IIB (TFIIB) orthologue TFB. We show here that the RNA polymerase of the archaean Methanococcus, in contrast to polymerase II, does not require hydrolysis of the β-γ bond of ATP for initiation of transcription and open complex formation on linearized DNA. Permanganate probing revealed that the archaeal open complex spanned at least the DNA region from −11 to −1 at a tRNA\textsuperscript{Val} promoter. The Methanococcus TBP-TFB promoter complex protected the DNA region from −40 to −14 on the noncoding DNA strand and the DNA segment from −36 to −17 on the coding DNA strand from DNase I digestion. This DNase I footprint was extended only to the downstream end by the addition of the RNA polymerase to position +17 on the noncoding strand and to position +13 on the coding DNA strand.

Initiation of transcription in archaea is mediated by orthologues of the eucaryotic transcription factors TATA box binding protein (TBP) and transcription factor IIB (TFIIB) (11, 26, 30). These two factors, archaeal TBP (αTBP) and transcription factor B (TFB), along with RNA polymerase, are sufficient for initiation of cell-free transcription at some promoters (12, 23). This finding is in line with the data derived from analyses of archaean genomes that indicate the absence of additional eucaryotic-like initiation factors like TFIIB, TFIIF, and TFIIH in archaea. Competition experiments using templates of different eucaryotic-like initiation factors like TFIIA, TFIIF, and TFIIH in archaea. Competition experiments using templates of different length and gel shift and footprinting experiments have shown that αTBP binds to the archaeal TATA box and that TFB stabilizes binding of αTBP to the promoter (8, 12, 23, 26, 30). This preinitiation complex seems to recruit the archaeal RNA polymerase to the promoter. The pathway of assembly of transcriptional components at the promoter, the existence of a TATA box at position −25 as a major signal directing initiation of transcription and start site selection (10, 25, 27), and the subunit structure and sequence of RNA polymerase (20) indicate a specific evolutionary relationship of the archaeal and eucaryotic RNA polymerase II (Pol II) transcriptional machinery.

In Pol II cell-free transcription systems on linear or relaxed templates, the DNA helicase activity of TFIIF is necessary for open complex formation (14). This activity requires hydrolysis of the β-γ phosphohydride bond of ATP in a step prior to initiation of transcription (15, 29, 31). This TFIIF requirement is bypassed by a supercoiled template at some basal Pol II promoters. ATP hydrolysis is required to drive at least two steps in the Pol II system, open complex formation and promoter clearance (25, 31). The striking similarity of the archaeal and eucaryotic Pol II systems posed the question of whether the archaeal RNA polymerase uses a similar mechanism for promoter opening.

In this study, we have used a highly purified Methanococcus cell-free system to investigate the early phase of transcription initiation. Promoter opening was analyzed by potassium permanganate footprinting, and DNA-protein interactions of the preinitiation complex were studied by DNase I footprinting.

MATERIALS AND METHODS

Reagents and enzymes. [γ\textsuperscript{−32}P]ATP and [α\textsuperscript{−32}P]UTP were purchased from Hartmann Bioanalytics (Braunschweig, Germany). The modified nucleotides and the dinucleotide UpG were from Sigma (St. Louis, Mo.). Restriction endonucleases and other DNA-modifying enzymes were purchased from Fermentas (Vilnius, Lithuania) or New England Biolabs.

Templates for in vitro transcription. The plasmid pIC-31/2 containing the promoter of the tRNA\textsuperscript{Val} gene was used in standard transcription reactions (10). The plasmid pIC-31/30PRO-C25 was used in addition. It contains the wild-type tRNA\textsuperscript{Val} promoter region, but all cytosine residues from positions +2 to +25 were replaced by other nucleotides (13). Templates were linearized with Aval, which cleaves the DNA in the vector region.

Purification of RNA polymerase. RNA polymerase from Methanococcus thermaUtolithrophicus was purified as described previously (7).

Expression and purification of recombinant αTBP. The coding region of αTBP (EMBL accession no. AJ271331) was subcloned using PCR amplification to generate the coding region with an NdeI restriction site at the 5’ end of the sequence and a BamHI site at the 3’ end. The amplified insert was then cloned between the NdeI and the BamHI restriction sites of the pET14b expression vector to generate pET14b-αTBP. After expression of αTBP with an N-terminal hexahistidine tag, BL21(DE3) cells containing the pET14b-αTBP plasmid were grown to an A\textsubscript{600} of 0.8 at 25°C. Expression of the protein was induced by addition of 1 mM isopropyl-1-β-D-thiogalactopyranoside (IPTG). Cells were harvested by centrifugation 3 h after induction, resuspended in buffer (50 mM Tris, pH 8.0, 50 mM NaCl, 20% glycerol), and disrupted by passage through a French press. The lysate was clarified by centrifugation at 4°C (100,000 × g for 20 min), and αTBP was purified by Ni-nitrilotriacetic acid agarose (Qiagen), MonoQ (Pharmacia), and Superdex 200 (Pharmacia) chromatography. The purified proteins were analyzed by sodium dodecyl sulfate-polycrylamide gel electrophoresis and stored at −70°C.

Expression and purification of recombinant TFB. The coding region of TFB (EMBL accession no. AJ271467) was subcloned using PCR amplification to generate the coding region with an NdeI restriction site at the 5’ end of the sequence.
The amplified insert was then cloned between the NdeI and the EcoRV restriction sites of the pET17b expression vector to generate pTFFBm17. BL21(DE3) cells containing the pTFFBm17 plasmid were grown to an A600 of 0.8 at 37°C. Protein expression and preparation of crude extract with buffer (50 mM Tris, pH 7.5, 300 mM NaCl) was done as described for aTBP. TFB was purified by HiTrap heparin (Pharmacia) and Superdex 200 chromatography.

In vitro transcription reactions. In vitro transcription experiments were performed in 25-μl mixture reactions that contained 190 fmol of template, 0.8 pmol of purified RNA polymerase, 1.7 pmol of recombinant aTBP, 1.7 pmol of recombinant TFB, and 4 μM [α-32P]UTP (370 Bq/pmol) in transcription buffer (20 mM Tris, pH 8.5, 2 mM MgCl2, 0.1 mM EDTA, 40 mM KCl, 3 mM dithiothreitol). Other ribonucleotides (Roche Diagnostics, Mannheim, Germany) used for in vitro transcription are indicated in the figure legends. After incubation for 20 min at 37°C, the reaction was terminated by adding 12.5 μl of loading buffer containing 98% formamide, 10 mM EDTA, 0.1% bromophenol blue, and 0.1% xylene cyanol. RNA products were analyzed by electrophoresis on denaturing 20% polyacrylamide gels.

Potassium permanganate footprinting. Potassium permanganate footprinting was performed with minor modifications of the procedure described by Jiang and Graff (18). Reaction mixtures were assembled as described above, with 20 fmol of negatively supercoiled or linearized DNA template in one-step transcription buffer. After 20 min of incubation, potassium permanganate was added to 6 mM for 3 min, followed by β-mercaptoethanol (final concentration, 700 mM) to stop the potassium permanganate reaction. EDTA to 10 mM and SDS to 0.5% were added, followed by phenol-chloroform extraction. The DNA was passed through a Sephadex G50 spin column for desalting. Potassium permanganate-sensitive sites were detected by asymmetric PCR using end-labeled M13 primers (located 139 bp downstream of the promoter) or M13 reverse primers (located 29 bp upstream of the promoter), 2.4 fmol of the desalted DNA, and a modified Taq DNA polymerase (Fermentas). PCR products were identified on 6% sequencing gels by using sequencing reactions with the same primer as size markers. For nonradioactive detection, a fluorescent-dye-labeled primer (DYEnamic ET primer; Amersham Pharmacia Biotech) was used instead of a radiolabeled primer. For nonradioactive detection, a fluorescent-dye-labeled primer (DYEnamic ET primer; Amersham Pharmacia Biotech) was used instead of a radiolabeled primer, and analysis was performed on an ABI 373 or ABI PRISM 310 automated sequencer.

RESULTS

Initiation of archael transcription does not require hydrolysis of the β-γ phosphoanhydride bond of ATP and GTP. To address the question as to whether ATP is required for activation of Methanococcus transcription, a reconstituted cell-free system consisting of bacterially produced Methanococcus TBP and TFB and highly purified RNA polymerase was used. The first 6 nucleotides (nt) of RNA initiated at the Methanococcus tRNAVal wild-type promoter did not contain an A residue (Fig. 1A). Therefore, provided that hydrolysis of the β-γ bond of ATP is not required for initiation of transcription, synthesis of a hexanucleotide was expected to occur in cell-free transcription reaction mixtures containing only GTP, CTP, and UTP. Analysis of the RNA products revealed that on both supercoiled and linearized templates, a transcript of the expected size was synthesized (Fig. 1B, lanes 9 and 10). When the ATP analogue cordycepin-5'- triphosphate (3'-dATP) was added at a concentration of 50 μM in addition to GTP, CTP, and UTP, synthesis of the same product and of a slightly longer RNA product was observed (Fig. 1B, lane C). With increasing concentrations of 3'-dATP, the intensity of the upper band increased (data not shown), suggesting that the longer RNA product is a heptanucleotide carrying 3'-dAMP at its 3' terminus. These findings supported the conclusion that the observed 6-nt transcript initiated accurately and exclusively at the G residue located 22 nt downstream of the TATA box that had been identified previously as the initiator nucleotide at this promoter (7). Control reactions showed that the synthesis of this RNA product was dependent upon the presence of two archael transcription factors and on the initiator nucleotide of GTP (Fig. 1B, lanes 1 to 4 and 6 to 9). These results demonstrated that a hydrolyzable β-γ phosphoanhydride bond of ATP was not necessary for initiation of transcription in Methanococcus.

To exclude the possibility that hydrolysis of the β-γ phosphoanhydride bond of GTP catalyzed open complex formation, GTP was replaced in cell-free transcription reaction mixtures by the analogs GTPγS and GMP-PNP containing nonhydrolyzable β-γ phosphoanhydride bonds. On both linearized and supercoiled DNA as a template, GTPγS or GMP-PNP was able to replace GTP (Fig. 2A, compare lanes 1, 4, and 7). Due to nonhydrolyzable β-γ phosphoanhydride bond to transcription reaction mixtures did not increase the rate of hexanucleotide synthesis occurring in the presence of either GTP or of analogs of GTP (Fig. 2A, lanes 2, 3, 5, 6, 8, and 9). These findings suggested that initi-
investigated DNA melting of the tRNAVal promoter in the archaeal promoter.

or GTP was not required for initiation of transcription at this position, and that hydrolysis of the GTP-initiated hexanucleotide (Fig. 2B, lane C). These data provide evidence that archaeal transcription could be initiated by a dinucleotide and that hydrolysis of the GTP-initiated transcript (labeled by an arrow) was reduced compared to that of the dATP-initiated transcript (lane C).

To provide conclusive evidence that hydrolysis of the β-γ phosphoanhydride bond of ATP or GTP for synthesis of the first five phosphodiester bonds (Fig. 2A, compare lanes 1 to 3, 4 to 6, and 7 to 9). In the presence of dATP, synthesis of longer RNA products occurred due to trace contaminations of dATP with ATP or to misincorporation of dAMP instead of AMP into RNA. Therefore, the electrophoretic mobility of the dinucleotide-initiated transcript was reduced in the presence of dATP (Fig. 2, lanes 3, 6, and 7), although dATP itself did not inhibit hexanucleotide synthesis.

To provide conclusive evidence that hydrolysis of the β-γ bond of a purine nucleotide was not required for initiation of transcription, GTP was replaced in transcription assays by the chemical probe specific for thymidines in single-stranded DNA. First, we have analyzed open complex formation on the non-template (lanes 1 to 6) and the template strand (lanes 6’ to 11) were analyzed by asymmetric PCR using radioactive labeled primers. Positions of reactive thymidine residues were determined by comigration of a sequence ladder terminated with ddATP and are indicated in relation to the transcription start site. Compare the following: lanes 1 and 6’, without protein and nucleotides as a control; lanes 2, 3, 7, and 8, without nucleotides; lanes 4 to 6 and 9 to 11, in the presence of 20 μM ATP (A), GTP (G), or CTP (C). (B) DNA sequence of plasmid pIC-31/2 containing the promoter and the region of the transcription start site. The positions of the modified thymidine residues are boxed. Position numbers refer to the transcription start site.

Potassium permanganate (KMnO₄) was used as a chemical probe specific for thymidines in single-stranded DNA. First, we have analyzed open complex formation on linearized wild-type DNA. Transcriptional components were incubated with DNA at 55°C. After treatment with KMnO₄, the hyperreactive sites on the non-template (lanes 1 to 6) and the template strand (lanes 6’ to 11) were analyzed by asymmetric PCR using radioactive labeled primers. Positions of reactive thymidine residues were determined by comigration of a sequence ladder terminated with ddATP and are indicated in relation to the transcription start site. The positions of the modified thymidine residues are boxed. Position numbers refer to the transcription start site.

Analysis of open complex formation. Since ATP is required for open complex formation in the Pol II system, we have investigated DNA melting of the tRNAVal promoter in the archaeon M. thermolithotrophicus in the presence and absence of ATP. Potassium permanganate (KMnO₄) was used as a chemical probe specific for thymidines in single-stranded DNA. First, we have analyzed open complex formation on linearized wild-type DNA. Transcriptional components were incubated with DNA at 55°C. Modified thymidine residues in single-stranded DNA regions were identified by an asymmetric PCR using a 32P-end-labeled primer and Taq DNA polymerase (see Materials and Methods).

Figure 3A shows the KMnO₄-detectable opening of the non-template and template DNA strands. Incomplete preinitiation complexes containing only aTBP and TFB showed no significant KMnO₄ sensitivity on both DNA strands (lanes 2 and 7) compared with the control lanes (1 and 6). When RNA polymerase was added to the aTBP-TFB-promoter complexes, modification of thymidine residues occurred at positions −3 to −1 and −6 to −8 and also at −10 (Fig. 3A, lane 2). These findings indicate that the open region extended at least from positions −1 to −10 at the noncoding DNA strand. Addition of either ATP, GTP, or CTP had no detectable effect on the modification patterns (Fig. 3A, lanes 4 to 6 and 9 to 11). These findings provide evidence that promoter opening was catalyzed...
by the archaeal RNA polymerase and that this step did not require energy provided by nucleotide hydrolysis.

When KMnO₄-detectable opening of the template strand was analyzed in reaction mixtures containing aTBP, TFB, and RNA polymerase, two prominent signals corresponding to T residues at positions −5 and −11 were observed (Fig. 3A, lane 5). The analysis of the KMnO₄ footprinting patterns on both DNA strands revealed that the open complex extended at this promoter at least from positions −11 to −1 (Fig. 3B, boxed T residues).

**Analysis of temperature dependence of open complex formation.** *M. thermolithotrophicus* is a moderate thermophile that grows between 30 and 70°C with an optimum at 65°C (16), and the cell-free transcription reactions and analyses of open complex formation shown here were carried out at 55°C. To investigate the ability of the *Methanococcus* RNA polymerase to form open complexes at temperatures comparable to conditions allowing open complex formation in enteric bacteria and eucaryotes and to analyze the temperature dependence of open complex formation, KMnO₄ sensitivity was assayed on both linearized and supercoiled DNA of the nontemplate strand containing a mutated tRNAVal promoter (pIC31/30PRO-C25; see Materials and Methods) at 40, 25, and 10°C. This template, which contains its first C residue at position 26, allows the synthesis of a 25-nt RNA product in transcription reactions carried out in the absence of CTP (13). Permangenate sensitivity was also detected by a PCR-based primer extension reaction, but a nonradioactive fluorescence-labeled primer was used and the modified thymidine residues causing termination of the primer extension reaction were detected by an ABI automated DNA sequencer (see Materials and Methods).

Analysis of KMnO₄-detectable opening of linear DNA revealed that at 40 and 25°C, the open complex extended from position −10 to −1 (Fig. 4A and data not shown). Even at 25°C, ATP had no effect on open complex formation (Fig. 4A, compare third and fourth panels from the top).

With negatively supercoiled DNA as template, promoter opening was shown to occur even at 10°C. The T residues corresponding to position −1 to −3, −6 to −8 and −10 were clearly modified in both the presence and absence of ATP (Fig. 4B, second and third panels from the top). When ATP, GTP, and UTP were added to this reaction, strong additional permanganate signals corresponding to positions +22, +20, +17/ +16, and +5 were detected and upstream signals disappeared, indicating bubble extension to the downstream DNA region (data not shown). This finding suggests that at 10°C, synthesis of a 25-nt transcript from a supercoiled template was still possible and an RNA product of this size could be also detected in cell-free transcription assays carried out at 10°C (Fig. 61x132)
Analysis of DNA protein contacts in preinitiation complexes. To couple the permanganate sensitivity assays of open regions in the DNA template to direct analysis of the RNA polymerase interaction with the promoter, DNase I footprinting experiments were carried out. The DNase I footprinting patterns of the αTBP promoter complex, the αTBP-TFB promoter complex and the αTBP-TFB-RNA polymerase promoter complex were analyzed. Footprinting reactions were carried out with DNase I by using a fluorescence-labeled DNA fragment. The nicked fragments were analyzed by an automated DNA sequencer.

On the nontemplate strand, αTBP protected the DNA region from −29 to −20 from DNase I digestion (Fig. 5A, TFB). TFB did not produce a footprint, and the αTBP-TFB footprint extended from position −40 to −14 (Fig. 5A, TFB and αTBP/TFB). DNase I hypersensitivity sites were observed at positions −5 to −7 and at positions −10 and −11 (αTBP/TFB). When complexes containing αTBP, TFB, and RNA polymerase were analyzed, a considerable extension of the footprint to the downstream DNA region was observed. This complex protected the DNA region from −40 to +17 from DNase I digestion (Fig. 5A, bottom panel). The hypersensitivity sites disappeared in this complex.

On the template strand, a protection of the BRE element and the TATA box region by αTBP, from positions −36 to −20, was found (Fig. 5B, αTBP). Binding of TFB to DNA was not detected (Fig. 5B, TFB), and the αTBP-TFB footprint extended from positions −36 to −17 (Fig. 5B, αTBP/TFB). DNase I hypersensitivity sites were generated by αTBP and TFB at positions −13/−14 and −37. The complex containing RNA polymerase in addition protected the DNA region from −35 to +13 from DNase I digestion (Fig. 5B, bottom panel). The hypersensitivity sites were also on the DNA strand not observed in the complex containing the RNA polymerase in addition. The results of the footprinting experiments are summarized in Fig. 6.

DISCUSSION

The archaeal RNA polymerase does not require hydrolysis of the β-γ bond of ATP for initiation of transcription. Transcription initiation by Pol II on a linearized template requires hydrolysis of the β-γ bond of ATP. Two steps during early transcription of Pol II promoters are dependent upon energy provided by ATP hydrolysis, open complex formation, and extension of the transcription bubble during RNA synthesis (31). One major finding described here is that although the archaeal transcriptional machinery strikingly resembles the Pol II machinery (see the introduction), the archaeal RNA polymerase did not require hydrolysis of the β-γ bond of ATP or GTP for initiation and thus uses a different mechanism for the early steps of transcription initiation.

Characteristics of the archaeal open complex. Permanganate footprinting experiments have shown that αTBP- and TFB-bound promoter DNA is in the closed conformation, whereas addition of RNA polymerase results in open complex formation.

FIG. 5. Interaction of the transcriptional components with the tRNA\(^{\text{Val}}\) gene. The DNA fragment was incubated with the components (25 pmol of αTBP, 8 pmol of TFB, 0.8 pmol of RNA polymerase) as indicated at the bottom of each chromatogram for 20 min at 37°C, followed by DNase I treatment. Further treatment was as described in Materials and Methods. For size calibration of the peaks of the template strand, size markers with a different fluorescent dye were added to each probe and size calling was done by using the global Southern method according to the instructions of the supplier. The calculation of the fragment length was calibrated using sequencing reactions generated with the fluorescence-labeled primer. For the analysis of the nontemplate strand, the 220-bp DNA fragment was also generated by PCR using a biotinylated M13 reverse primer and a fluorescent-dye (ABI JOE)-labeled M13 primer. The DNA sequences of the plasmid pIC-31/30PRO-C25 are shown on the tops of the chromatograms in such a way that the peaks of the chromatograms can be directly correlated to the individual base positions within the DNA sequence. The promoter and the transcription start site are underlined. * DNase I hypersensitive sites.
Similar to Pol II (31) and E. coli RNA polymerase (6), the archaeal enzyme is able to melt a DNA segment of 10 nt (Fig. 3) located between positions −11 and −1. In striking contrast to the Pol II system, open complex formation occurred on linear DNA in the absence of nucleotides and was not enhanced in the presence of ATP (Fig. 3 and 4). ATP-independent promoter opening of the DNA region between −1 and −12 of a Sulfolobus tRNA promoter has been shown (1). However, in this case, the template was in negatively supercoiled conformation. DNA in this conformation can be also melted by Pol II in the absence of ATP (9, 31). In the Methanococcus system, the extent of the promoter opening is independent of template topology, but negative supercoiling of DNA clearly energetically favors promoter opening as indicated by the lower temperature limit for promoter opening and RNA synthesis with negatively supercoiled DNA compared to linear DNA (Fig. 5).

**DNase I footprinting analyses of an archaeal preinitiation complex.** The αTBP-TFB promoter complex has been studied in the Pyrococcus and Sulfolobus systems (12, 24). In addition, the crystal structure of the Pyrococcus preinitiation complex containing N-terminally-truncated TFB and αTBP in complex with promoter DNA was recently determined (22), but the interaction of archaeal RNA polymerase with these αTBP-TFB promoter complexes has not yet been described. Evidence for semispecific initiation of purified Sulfolobus RNA polymerase independent of a TATA box and the presence of transcription factors has been obtained (2, 17). A DNase I footprint of purified Methanococcus RNA polymerase on a homologous hisΔ promoter (3) and exonuclease III footprints of this enzyme on several promoters (27, 28) were reported, but Methanococcus transcription factors were not included in these studies since they were discovered later (7, 11).

A sequence of 6 nt immediately upstream of the TATA box was identified recently as the recognition element for TFB (BRE) and found to direct the polarity of archaeal transcription (2, 22). This element was also contained within the DNA region protected after addition of Methanococcus TFB. The GC base pair at position −1 of the tRNAVal promoter (relative to the TATA box) deviates from the consensus for BRE sequences RNWAAW (R = purine, W = A or T, N = any base). However, earlier mutational analyses of the tRNAVal promoter provide evidence that the major structural determinants are conserved between the BRE element of the Sulfolobus T6 and Methanococcus tRNAVal promoters. A single-point mutation of the G residue at position −1 (relative to the TATA box) to a T did not affect transcriptional activity of the tRNAVal promoter (10), suggesting that this deviation does not influence the interaction of Methanococcus TFB with BRE. By contrast, when the A residue at position −3 of the tRNAVal BRE element that has been identified as a key determinant of Sulfolobus BRE (2) was replaced by a G, the rate of transcription was reduced to 45% (10).

Similar to the Sulfolobus system (2), a DNase I hypersensitivity site 8 bp upstream of the TATA box was observed. This site was shown to correspond to a bend of the DNA in the Sulfolobus system that brings the major groove of the DNA in the BRE in close contact with a helix-turn-helix motif of TFB (22). In contrast, the DNase I hypersensitivity site downstream of the TATA box has not been observed in the Sulfolobus TBP-TFB promoter complex. This finding provides additional evidence for a significant difference in the interaction of Methanococcus and Sulfolobus TFB with promoter DNA.

Addition of RNA polymerase resulted in a large extension of the footprint to the 3’ end on both DNA strands (Fig. 5 and 6). As the footprint from position −40 to −14 can be attributed to interaction of transcription factors with DNA, the DNA region that was in contact with RNA polymerase extended at least from position −15 to +17 on the nontemplate strand and from −15 to +14 on the template strand. Direct contacts of the enzyme with the DNA region protected by the transcription factors could not be demonstrated even though the enzyme may also bind to this DNA region.

**Comparison with Pol II and bacterial RNA polymerase holoenzyme.** The archaeal system shows similarities to both Pol II and the bacterial RNA polymerase. Like the ββ′σ70 system, the archaeal RNA polymerase was able to melt promoter DNA without additional factors and ATP hydrolysis. By contrast, open complex formation in the Pol II system requires an additional enzymatic activity, the ATP-dependent DNA helicase of TFIIH.

The RNA polymerase of E. coli protects about 70 bp of DNA from DNase I digestion in the preinitiation complex (5, 19). In contrast to the E. coli enzyme and similar to Pol II, the archaeal enzyme binds to a preformed ternary complex of αTBP and TFB with promoter DNA, and like Pol II (4), the Methanococcus RNA polymerase extends the footprint to the DNA region downstream of the TATA box. In contrast to the Pol II system, no RNA polymerase-induced extension of the DNase I footprint to the region upstream of the TFB binding site was observed in the Methanococcus system. A further difference from the Pol II system was that archaeal
TFB extended the TBP footprint on either site of the TATA box, whereas TFIIIB causes an extension of the TBP footprint only to the downstream site of the TATA box (4).

Our definition of the open DNA region and of DNA protein interactions in a preinitiated complex allows an examination of the mechanism of transcription bubble and RNA polymerase translocation in isolated complexes stalled in various registers.

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