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# A whole-genome approach to identifying protein binding sites: promoters in *Methanocaldococcus* (*Methanococcus*) *jannaschii*

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## ABSTRACT

We have adapted an electrophoretic mobility shift assay (EMSA) to isolate genomic DNA fragments that bind the archaeal transcription initiation factors TATA-binding protein (TBP) and transcription factor B (TFB) to perform a genome-wide search for promoters. Mobility-shifted fragments were cloned, tested for their ability to compete with known promoter-containing fragments for a limited concentration of transcription factors, and sequenced. We applied the method to search for promoters in the genome of Methanocaldococcus jannaschii. Selection was most efficient for promoters of tRNA genes and genes for several presumed small noncoding RNAs (ncRNA). Protein-coding gene promoters were dramatically underrepresented relative to their frequency in the genome. The repeated isolation of these genomic regions was partially rectified by including a hybridization-based screening. Sequence alignment of the affinity-selected promoters revealed previously identified TATA box, BRE, and the putative initiator element. In addition, the conserved bases immediately upstream and downstream of the BRE and TATA box suggest that the composition and structure of archaeal natural promoters are more complicated.

#### INTRODUCTION

Relative to the explosive growth in DNA sequence data, our understanding of genomes, their dynamics and the organisms that are hosts to them seems to grow slowly. Although several sophisticated algorithms exist to predict protein-coding regions (e.g. 1–4), the identification of DNA elements that play crucial roles in their expression lags far behind. This is especially notable in the case of transcriptional promoters. The identification of promoters has been limited, by and large, to studies of individual genes. Traditional genetic, biochemical and structural studies of promoters, with their emphasis on identifying and characterizing individual promoters, cannot meet the challenge of the genomic era. These approaches—be they based on RNA analysis [e.g. nuclease protection (5) and reverse transcriptase run-off (6)] or based on DNA analysis [e.g. transcription factor or polymerase footprinting (7) and analysis of *in vitro* transcript run-offs (8)]—are not readily scalable to probe whole genomes.

There are few genomes with systematically annotated promoter features. Computational approaches to promoter identification are unreliable when the analysis is carried out a genome at a time (9). Comparative analysis of appropriately related genomes can identify important DNA sequences (10,11), but these methods are not specific to promoters. Even in organisms where promoters have been extensively studied, the available tools for in silico identification remain wanting. The situation is even more critical in the case of the Archaea, where there is little experimental data to provide the bases for the calibration of computational tools. Other genome-wide probing methods, such as RNA analyses with oligonucleotide arrays, can identify transcribed regions (under specific growth conditions), but do not locate the start site of the transcripts, nor do they provide information on cis-acting regulatory regions.

Archaeal transcription shares similarity with the eukaryal pol II transcription system, in terms of the enzyme and factors involved, as well as promoter architecture (12–15). Core components include the general transcription factors—TATA-binding protein (TBP) and transcription factor B (TFB), which are homologs of eukaryal TBP and TFIIB—and one multi-subunit RNA polymerase, similar to RNA pol II in overall architecture and subunit composition (16,17). Transcription initiation involves the recognition of promoter sequences by TBP, the rate-limiting step. A correctly oriented TBP–promoter

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complex is stabilized by binding of TFB, which is also responsible for recruiting RNA polymerase (14,15,18).

Mirroring the similarities in the transcription machineries, the available information supports the notion that core elements of archaeal promoters are similar to those of pol II promoters. A TATA box and an upstream purine-rich element (BRE) are key players in the assembly of the pre-initiation complex. The TATA box provides the initial recognition element for transcription factor binding, with the BRE stabilizing appropriate binding of TBP and being instrumental in defining transcription orientation. The formation of the tripartite archaeal preinitiation complex has been confirmed experimentally using footprinting assays, electrophoretic mobility shift assays (EMSA) and base-specific cross-linking (19-23). Crystallography of the ternary complex has revealed base-specific contacts between promoter elements and transcription factors TBP and TFB (24,25). A third element, the initiator (Inr), located at the transcription start site, is similar to its eukaryal counterpart (26,27). In Archaea, it appears to be less important, since in vitro transcription initiates readily from new sites in artificial promoters recovered by *in vitro* evolution (20), or in promoters with insertions/deletions between the TATA box and Inr (28). Moreover, while this element is conserved in Methanococcales and Sulfolobales, it is not evident in Haloarchaea (29).

Of the elements comprising the pre-initiation complex in Archaea (and in Eucarva), the promoter is the least understood. To facilitate systematic analyses, we set out to identify and characterize archaeal promoters on a genome-wide basis. We devised an efficient method for isolating and identifying genomic sequences that specifically bind to the general transcription factors TBP and TFB. We chose to carry out the analysis in the archaeon Methanocaldococcus jannaschii, the first member of the Archaea to have a fully sequenced genome (30). Since binding of transcription initiation factors to promoter sequences can be readily assayed by EMSA (31,32), we postulated that this could be modified into a method to select protein binding sequences from a pool of random genomic DNA fragments. To identify the desired fragments, we use the polymerase chain reaction (PCR) to amplify DNA sequences with increased apparent molecular weight, followed by cloning and sequencing of random and/or screened clones. The procedure is very similar in spirit to in vitro evolution and 'systematic evolution of ligands by exponential enrichment' (SELEX) (33,34) or, more closely, to genomic SELEX (35-40) since we did not wish to alter the sequences or binding ability of the natural DNAs.

## MATERIALS AND METHODS

Unless stated otherwise, all kits and enzymes were used as recommended by their manufacturers.

#### **Genomic DNA library**

Methanocaldococcus jannaschii cells were grown in a reactor vessel as previously described (41). Late

exponential cells from 11 of culture were lyzed by three cycles of freezing and thawing in the presence of 0.5% SDS and their genomic DNA was purified (42). One hundred micrograms of DNA, re-suspended in 1 ml 25% glycerol/1 M NaOAc (pH 5.5), was exhaustively fragmented using a Branson Sonifier Cell Disruptor by repeated 15 s pulses to yield fragments of 400-500-bp average size. Following ethanol precipitation, 45µg of fragmented genomic DNA was treated with 1U BAL31 nuclease (New England Biolabs, NEB) for 7 min at 30°C in a 100 µl volume. After extraction with phenol/chloroform and ethanol precipitation, the fragments were resolved in a 1.5% SeaPlaque (FMC Corp.) agarose gel. The region of the gel containing 200-300-bp fragments was excised and the DNA extracted (Qiagen Gel Extraction kit). DNA fragments measuring 320 ng were treated with 1 U T4 DNA polymerase (Invitrogen) for 30 min at 37°C and the DNA was ligated overnight at 16°C to 600 ng of SmaIcleaved and dephosphorylated pUC18 vector (Invitrogen). The ligation product was treated with 1U DNA polymerase I (NEB) for 1 h at 16°C to move (by nicktranslation) the nick between the insert and the (initially dephosphorylated) plasmid, thereby expanding the region of intact duplex DNA. Based on the frequency of colonies recovered following electrotransformation into Escherichia coli strain XL1Blue-MRF', we estimate that this primary library (20  $\mu$ l total volume) contains 4  $\times$  10<sup>6</sup> recombinant molecules/µl. This is likely an underestimate of the effective complexity in the PCR-based procedures described subsequently.

### DNA fragments for the selection

Aliquots of the primary library were used as template for generating a collection of random DNA fragments by PCR using M13 universal primers. Each 100 µl reaction contained 20 mM Tris–HCl (pH 8.4), 50 mM KCl, 2 mM MgCl<sub>2</sub>, 100 pmol of each primer, 40 nmol dNTPs, 1 µl library DNA and 3 U *Taq* DNA polymerase (Invitrogen). Following an initial denaturation at 94°C for 90 s, the library DNA was amplified for 25 cycles with denaturation at 95°C for 30 s, annealing at 55°C for 30 s and extension at 72°C for 30 s. The final elongation was 72°C for 10 min. The products of the amplification include 134 bp of plasmid DNA flanking the inserts. The amplified DNA was resolved in low-melting temperature agarose and the 300–400-bp DNA fragments were recovered as above.

For EMSA, DNA was end-labeled using  $[\gamma^{-32}P]ATP$ and T4 polynucleotide kinase (Invitrogen). Labeled fragments were purified using a QIAquick Nucleotide Removal Kit (Qiagen).

## Expression and purification of transcription factors

TBP from *M. jannaschii* was cloned as an N-terminal  $His_6$ -tagged recombinant protein in the expression vector **pQE31** (Qiagen). TFB, without intein, was cloned as a C-terminal  $His_6$ -tagged recombinant protein in the vector **pET29b** (Novagen). The recombinant proteins were expressed in *E. coli* BL2-CodonPlus<sup>®</sup>(DE3)-RIPL.  $His_6$ -tagged proteins were affinity-purified with

QIAexpressionist<sup>TM</sup> (Qiagen). Cells were grown to OD<sub>600</sub> of 0.5. Following a 1.5-h induction with 1 mM isopropyl- $\beta$ -D-thiogalactopyronoside (IPTG), cells were collected by centrifugation and re-suspended in lysis buffer [50 mM Na·PO<sub>4</sub> (pH 8.0), 300 mM NaCl and 10 mM imidazole], followed by two passages through a French pressure cell. The lysate was heat-treated at 70°C for 8 min, and spun at 15000 g for 15 min to remove denatured host proteins. The cleared cell lysate was mixed with 1/6th volume of Ni<sup>2+</sup>-NTA agarose equilibrated with lysis buffer, loaded onto the column and eluted with an increasing concentration of imidazole. Fractions containing the target protein were pooled and dialyzed against 20 mM Tris-HCl (pH 7.8), 500 mM NaCl and 40% glycerol. Purified proteins were concentrated by ultrafiltration using an Amicon Centriprep YM10.

To increase the stability of TFB, a C-terminal, DNAbinding fragment (TFBc) was created by removing the first 137 amino acids. The resulting sequence was cloned into pET29b expression vector and expressed as a C-terminal His<sub>6</sub>-tagged recombinant protein. The purification procedure was slightly modified. Briefly, host cells were induced with 0.4 mM IPTG when OD<sub>600</sub> reached 0.5, and harvested after 2h. The cell pellet was re-suspended in lysis buffer [20 mM Tris-HCl (pH 8.3), 50 mM NaOAc, 180 mM KCl, 10 mM β-mercaptoethanol, 20 mM imidazole and 10% glycerol], and lyzed by sonication using a Branson sonifier fitted with a microtip (10 times for 30s, with 30s breaks). The dialysis step was omitted. The pooled fractions were concentrated in an Amicon-stirred ultrafiltration cell, and then passed through a Pharmacia PD10 column to remove imidazole.

## **Electrophoretic Mobility Shift Assays (EMSA)**

For assaying the formation of the DNA/TFB/TBP complex, 20 µl reactions containing binding buffer [60 mM KCl, 20 mM Tris-HCl (pH 7.5), 10 mM MgCl<sub>2</sub>, 9 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 0.05 mM EDTA, 0.5 mM DTT, 0.05 mM phenylmethylsulfonylfluoride (PMSF), 5% glycerol, 2.5% PEG8000], 1µg poly(dI-dC), 200 ng TBP, 400 ng TFB and 1 ng labeled DNA were incubated at 65°C for 20 min. Binding assays for DNA/TFBc/TBP contained TFBc binding buffer [150 mM KCl, 20 mM Tris-HCl (pH 7.5), 10 mM MgCl<sub>2</sub>, 0.05 mM EDTA, 0.5 mM DTT, 0.1 mM PMSF, 5% glycerol], 20 ng TFBc, 50 ng TBP, 1 µg poly(dI-dC) and 1 ng labeled DNA; incubation was at 75°C for 30 min. After incubation, reactions were loaded on a 15cm long 5% polyacrylamide gel containing  $0.5 \times TBE$  (45 mM Tris-borate, 1 mM EDTA, pH 8.3), 1 mM MgCl<sub>2</sub> and 1% glycerol. Gels were run in  $0.5 \times TBE$ at 200 V for 2–3 h, exposed to X-ray film, or visualized by phosphorimaging, and quantitated with Image Gauge software.

Regions of the gel above the free probe containing the fraction of bound DNA (as inferred from their migration relative to that of tRNA<sup>Val</sup> promoter and TBP/TFB) were excised and the DNA extracted. Gel slices were soaked overnight at  $37^{\circ}$ C in buffer [0.5 M NH<sub>4</sub>OAc, 10 mM Mg(OAc)<sub>2</sub>, 1 mM EDTA (pH 8.0) and 0.1% SDS], followed by phenol and chloroform extractions and

recovered by ethanol precipitation in the presence of carrier yeast tRNA. This material was amplified using M13 universal primers. Fifty microlitre PCR reactions were assembled containing 1/10th of gel-recovered DNA, 20 mM Tris–HCl (pH 8.4), 50 mM KCl, 2 mM MgCl<sub>2</sub>, 50 pmol primers, 20 nmol dNTPs and 3 U *Taq* DNA polymerase. The program used was denaturation at 94°C for 2.5 min, 26 cycles (95°C for 30 s, 55°C for 30 s, 72°C for 30 s) and a final extension at 72°C for 10 min. The resulting products were digested with KpnI and SalI, and cloned into correspondingly cut pUC18 plasmid.

# Competition assay of selected DNAs versus a known promoter

The ability of the recovered fragments to compete effectively with the labeled tRNA<sup>Val</sup> reference promoter was tested in binding reactions as previously described, except for the addition of 200-ng unlabeled recombinant plasmid DNA. When binding reactions included TFBc (not TFB), the amount of unlabeled recombinant plasmid was raised to 400 ng. Gels were run as described and visualized by phosphorimaging. Labeled bound and free DNA were measured using Image Gauge, and the ratio of  $f_{b/f,i} = \text{DNA}_{\text{bound},i}/\text{DNA}_{\text{free},i}$  was calculated. The  $f_{b/f}$ was first normalized using the ratio from a reference binding reaction without competitor DNA  $(f'_{b/f,i} = f_{b/f}/f_{b/f,0})$ , and then referenced to the average  $f_{b/f, \text{ average}}^{(j)}$  for the gel  $(f_{b/f,i}^{\prime}n = f_{b/f, \text{ average}}^{\prime}/f_{b/f,i}^{\prime})$ . The resulting  $f_{b/f,i}^{\prime}n$  was used as a measure of the relative affinity to the transcription factors. The cut-off normalized score for promoters with 'high affinity' was set at 0.9.

## Hybridization screening of selected clones

For dot hybridizations, 30 ng plasmid DNA were denatured in 0.2 M NaOH at room temperature for 15 min, and transferred to Hybond-N+ nylon membranes. The membrane was rinsed briefly in  $2 \times$  SSC (0.3 M NaCl and 30 mM sodium citrate), and the DNA fixed by vacuum drying at 80°C for 2 h. Pre-hybridization was at 68°C for 30 min in 10 ml buffer containing 0.25 M Na·PO4 (pH 7.2), 1.0 mM EDTA, 7% SDS and 1% BSA. Hybridizations were to a mixed probe-containing promoter DNA from 15 tRNA genes and one non-coding RNA gene that had been repeatedly selected in the initial screening. Promotercontaining DNA fragments were generated by PCR and cut with KpnI and BamHI; 50 ng of each were pooled. The fragment mix was labeled with  $\left[\alpha^{-32}P\right] dCTP$  in a fill-in reaction using Klenow fragment of E. coli DNA polymerase (Invitrogen), and purified with QIAquick Nucleotide removal kit (Qiagen). The probe was denatured at 95°C for 4 min before use. After overnight hybridization at 68°C in the same buffer, the membrane was washed with  $2 \times SSC/0.1\%$  SDS for 20 min, then  $0.2 \times$  SSC/0.1% SDS for 20 min, followed by a final wash with  $0.1 \times SSC/0.5\% SDS$  for 20 min. Hybridization signal was measured by phosphorimaging, and a normalized signal intensity for each plasmid was calculated  $(S_i^n = S_i / S_{\text{average}}, \text{ where } S_{\text{average}} \text{ is the average signal}$ strength from plasmids from the same preparation and

the same membrane). The cut-off normalized score for 'strong hybridization' was set at 0.9.

#### Alignment of promoter sequence and Logos

Collected sequences were inspected for the presence of conserved elements either manually or using MEME (43). Sequences were aligned by centering the conserved elements. Matrices of the base composition were derived at every position, and information content was calculated as:

$$I = \sum_{i \in \{A, C, G, T\}} \frac{n_i + p_i}{N + 1} \log_2\left(\frac{n_i + p_i}{(N + 1)p_i}\right)$$

where, *I* is information content of the position in bits,  $p_i$  is random frequency of residue type *i*,  $n_i$  is the number of instances of residue type *i* at the position and *N* is total number of sequences analyzed. This formula includes a pseudocount of  $p_i$  for residue type *i* at each position, as small count correction. Sequence logos (44) were generated based on the local alignment. At each position, the height of the character for residue type *i* was determined by  $h_i = I n_i/N$ .

## RESULTS

#### EMSA of a known promoter

It has been previously demonstrated that the binding of transcription initiation factors TBP and TFB to a DNA fragment containing an archaeal promoter can substantially reduce the mobility of the DNA on a gel (31,32). Typically, the mobility shift of a labeled DNA fragment is used as an assay for transcription factor binding. We reasoned that we could select promoter-containing DNA regions in the *M. jannaschii* genome by using EMSA to separate fragments of genomic DNA that bound transcription initiation factors TBP and TFB (and thus presumably contain a promoter) away from those that do not.

We first optimized conditions for transcription factor binding and gel shift assays using a 254-bp radiolabeled DNA fragment containing the tRNA<sup>Val</sup> gene promoter (P<sub>tRNAVal</sub>) from Methanococcus vannielii (28), a sequence that we had previously shown to be efficiently bound by the M. jannaschii transcription factors (32). Recombinant M. jannaschii TBP and TFB genes were expressed in E. coli, and the His<sub>6</sub>-tagged proteins purified. Efficient formation of the TBP/TFB/DNA complex was limited to the temperature range 55-80°C (data not shown). Only trace amounts of shifted DNA were detected following incubation at lower temperature, and no shifted band was observed following incubation at higher temperature. The low temperature limit agrees with the fact that M. jannaschii is a thermophile. The reason for the high temperature limit was not pursued (possibilities include protein instability, complex instability or denaturation of the high-A + T DNA fragment). Incubation pH has little effect on formation of the ternary complex in the range from 6.8 to 8.3 (data not shown). The condition most critical for efficient complex formation was the monovalent cation concentration; the optimal binding was

found at 60–90 mM  $K^+$  (data not shown). Binding was specific, as demonstrated by the fact that it could be competed by addition of unlabeled cognate fragment, but not by addition of a DNA fragment of similar size containing the plasmid pUC18 polylinker region.

In the process of optimizing binding conditions, we noticed that upon incubation, purified TFB lost its binding ability precipitously. When the protein was preincubated in binding reaction conditions for 12 min at  $60^{\circ}$ C, only half of its binding ability was retained. When pre-incubation was carried out at  $65^{\circ}$ C for 15 min, the loss of binding activity was almost  $80^{\circ}$ . This loss of activity was observed over a wide range of pH values and ionic strengths. Instability of the *M. jannaschii* pre-initiation complex (consisting of TBP, TFB and RNA polymerase assembled on a suitable DNA substrate) has been reported (45), and our results suggest that it may be due to instability of TFB.

Archaeal TFB, like its eukaryal homologs, contains a C-terminal core domain responsible for interactions with TBP and DNA binding, and an N-terminal domain responsible for recruiting RNA polymerase. We evaluated the stability and binding ability of a truncated version of the protein lacking the N-terminal domain (referred to as TFBc). To construct TFBc, we aligned M. jannaschii TFB with homologs for which a crystal structure was available. The alignment revealed a conserved C-terminal domain of about 200 amino acids, corresponding to the core domain from Pyrococcus woesei (P. furiosus subsp. woesei) TFB used in crystallographic analyses (25). Based on these data, the first 137 amino acids of the protein were removed, and the C-terminal domain was expressed as a His<sub>6</sub>-tagged recombinant protein. TFBc formed the ternary DNA/TBP/TFBc complex with 5- to 10-fold greater efficiency than the full-length protein. Although a modest amount of non-specific binding was observed when reactions were carried out at 65°C, this was eliminated by raising the incubation temperature to 75°C and increasing the concentration of  $K^+$  to 150 mM. At least two factors are likely to contribute to the greater activity: TFBc is much more stable upon incubation than the full-length protein (only marginal loss was observed upon preincubation at  $75^{\circ}$ C for 30 min); and, it is possible that the N-terminal domain of the protein hinders the formation of the ternary complex, as has been observed in eukaryotes (46). Therefore, except for the first round of selection, all experiments below were conducted with TFBc, instead of the full-length protein. In contrast, TBP was stable after purification and did not lose activity measurably upon incubation in binding reaction conditions.

## Selecting promoter-containing genomic DNAs by transcription initiation factor binding

We constructed a library of *M. jannaschii* DNA by ligating  $\sim$ 250-bp random fragments into the SmaI restriction site of the plasmid pUC18. Vector sequences flanking the cloning site provided anchors for 25 cycles of PCR amplification of the DNA inserts to yield a population of  $\sim$ 385-bp linear DNA molecules for the selection for TBP/TFB-binding sequences. To ensure a thorough

sampling of genomic sequences in the selection, an aliquot of the primary library representing  $\sim 30$  genome equivalents was used as substrate for amplification. Because the inserts had been size-selected, they migrated as a well-defined band on an agarose gel (Figure 1A). For promoter selection, 12 ng (0.05 pmol) of labeled DNA fragments were incubated with 28 pmol each of TBP and TFB in a 60 µl reaction volume, and then resolved on a polyacrylamide gel (Figure 1B). No shifted band was directly observed, presumably because the amount of promoter-containing DNA constitutes only a small part of the entire population of fragments. Gel regions above the free DNA fragments were excised, the DNA extracted from them, and the resulting material subjected to 26 cycles of PCR amplification. Although the procedure was expected to recover the desired DNAs (those shifted to a lower mobility by the binding of transcription factors), it was also possible that the amplification products were due to a small amount of larger DNA contaminating the original library. Electrophoretic analysis of the amplified recovered DNAs verified that they were 350-400 bp, consistent with the original library (Figure 1C).

Having demonstrated that the recovered DNA was of the desired size, we needed to verify the ability of individual DNA fragments to bind transcription factors. The PCR products were cut with the restriction enzymes KpnI and SalI and ligated into similarly cleaved pUC18 plasmid. Following transformation and growth, recombinant plasmids were tested for their ability to compete with a labeled fragment containing the tRNA<sup>Val</sup> promoter ( $P_{tRNAVal}$ ) for a limited supply of transcription factors (Figure 1D). Not only did this approach avoid the need to label the fragment from each clone, but it also provided a measure of each fragment's affinity for TBP and TFB relative to that of  $P_{tRNAVal}$ .

Of 215 clone inserts tested in this way, 121 were judged to compete significantly with the  $P_{tRNAVal}$  (Figure 2A). Of those judged to compete, 70 were sequenced and mapped onto the annotated *M. jannaschii* genome (30). Of these, 55 (almost 80%) included potential promoters upstream of tRNA-encoding genes (Figure 2A). These encompass 15 of the 23 tRNA genes that we predict to have their own promoter (E.L., unpublished data). Individual promoter regions were isolated 1–8 times (Tables S1 and S2). All sequenced fragments were different from one another, so multiple isolates of a given genomic region are independent events, not artifacts of duplicate clones. Five (7%) of the 70 sequences included potential promoters adjacent to G + C-rich islands with no known function or identified



**Figure 1.** Isolation of *M. jannaschii* genomic fragments with affinity to the transcription factors TBP and TFB. (A) PCR amplification of a small insert (200–300 bp) *M. jannaschii* genomic library. Left lane, 1-kb DNA ladder (Invitrogen); right lane, PCR product. (B) EMSA of TBP/TFB and labeled DNA. Fractions 1–4 contain transcription factor-bound DNA. (C) Agarose gel analysis of the re-amplification of transcription factor-bound DNA. Left lane, 1-kb DNA ladder; rightmost 4 lanes, product amplified from recovered DNA fractions 1 to 4 (Figure 1B). (D) Competition assay of recovered DNA fragments with the reference  $P_{tRNAVal}$  promoter. Lane SP, specific competitor (plasmid containing the reference  $P_{tRNAVal}$  promoter); lane NSP, non-specific promoter (empty vector); lanes C1 to C19, various recombinant plasmids harboring recovered genomic fragments.



**Figure 2.** Flow chart of screening strategies to characterize putative promoters isolated as transcription factor-bound genomic DNA fragments. (A) Initial selection without hybridization-based screening. (B) Selection with hybridization-based screening against most commonly isolated genome regions.

homologs in other organisms. We presume that these G + C-rich regions are small non-coding RNAs (ncRNAs), as other workers have suggested for similar regions (47,48). Three distinct putative ncRNAs are represented among the five clones. The presence of appropriately oriented transcription factor-binding sequences adjacent to these islands lends credence to their identification as transcribed genes (Table S1). Among the 70 clones, we uncovered only 3(4%) putative promoters for protein-coding genes. The final eight fragments (11% of the 70) mapped to intragenic regions in the chromosome. Although it is tempting to regard these as false positives, this would be a potentially misleading designation in that they compete effectively with a tRNA gene promoter for transcription factors.

It appears that the competition screening introduced a heavy bias towards the isolation of strong promoters (and specifically tRNA promoters). For comparison, we performed additional competition assays against a protein-coding gene promoter. We used the promoter region of MJ1335, a gene annotated as encoding phosphoheptoisomerase. We had isolated this promoter region in the original selection and qualitatively characterized it as competing, but only weakly, with  $P_{tRNAVal}$ . We tested 47 of the DNA fragments that did not compete effectively with  $P_{tRNAVal}$ . Of these, 21 successfully competed with  $P_{MJ1335}$  and 18 of these were subsequently sequenced. Of these, five (28%) contained tRNA promoters, five (28%) contained presumptive ncRNA promoters and two (11%) contained a putative protein promoter. Finally, the number of fragments mapping to intragenic regions rose to 39%.

Our double competition criterion (competes with a protein gene promoter, but not a tRNA gene promoter)

decreased substantially the number of tRNA promoters (from 80% to 28%). Unexpectedly, the recovery of promoters for putative ncRNAs increased dramatically.

To assess the bias introduced by our competition-based screens, we sequenced 24 random recovered DNA fragments that did not compete efficiently with  $P_{tRNAVal}$ . Of these, 21% contained putative promoters (three for tRNA genes and two for protein-coding genes). One of the tRNA gene promoters found was not among those found previously, bringing the total to 16 of the expected 23 promoter regions. The remaining 79% mapped to intragenic regions. Because we have no confirmation of their binding of transcription initiation factors, some of them might be *bona fide* false positives from the selection.

## Screening clones by hybridization

The recurring isolation of tRNA gene promoters led us to explore the use of a hybridization-based screening step against the tRNA promoter regions that we had already isolated. A preliminary screen using colony hybridization (49) was judged to be insufficiently reproducible for reliable results (data not shown). Clearer results were obtained when we used dot-blot hybridization to survey clones isolated by EMSA (Figure 2B). We analyzed 996 of these to determine how well they hybridized to a pool of already recovered and labeled tRNA promoters. For each clone, we also assessed the relative affinity for the transcription factors, as inferred from its ability to exclude the reference labeled tRNA<sup>Val</sup> promoter from the ternary complex in a competition assay (TFBc was used in these assays since it is more stable in solution than the fulllength protein and yielded more reproducible data).

Therefore, each clone is identified by two coordinatesstrength of hybridization to the pool of tRNA promoters recovered in the initial study, and relative affinity to the transcription factors (see Materials and Methods section for details). A double logarithmic scatter plot  $[\log(S_i^n) \text{ versus } \log(f'_{b/f,i}n)]$  was generated (Figure 3). Clones were divided into four categories. Those that hybridized the mixed promoter probe and competed with  $P_{tRNAVal}$  (34% of the total) were expected to be repeat isolations of tRNA promoters, and sequenced instances indicated this fraction was enriched with tRNA gene promoters (68% of total). Those that did not hybridize the mixed promoter probe and did not compete with P<sub>tRNAVal</sub> (25% of the total) were candidates for false positives (even though they were isolated by the gel shift). Eleven out of 12 sampled instances were found to be intergenic regions or be downstream of genes. The 297 clones that did not hybridize the mixed promoter, but did compete with PtRNAVal (30% of the total) were considered the best candidates for potentially unidentified promoters, and they were extensively sampled and sequenced. The composition of 167 sequenced clones is summarized in Figure 2B. Of particular note is the increase in the proportion of protein-coding gene promoters, showing that they are being gel shifted by the transcription factors, but much less efficiently than the tRNA and ncRNA promoters. Additional evidence that the screening helped to survey new promoters is that four additional tRNA genes were represented among the upstream regions of 37 tRNA genes, bringing up the total of recovered tRNA promoters to 20, out of 23 predicted.

Although the introduction of screening schemes (both double competition assays, and prescreening of clones by hybridization) improved the recovery of novel putative promoter sequences, it is clear that the DNAs isolated in our selection are a non-random sample of the genome, being highly enriched in potential promoters for tRNAs and (probably) also for ncRNAs. These results suggest that—not surprisingly—promoters in *M. jannaschii* have different levels of affinity to the transcription factors TBP and TFB, and that the combination of the experimental conditions employed in the selection and subsequent screening are sensitive to these differences.

## Core promoter elements in M. jannaschii

The sampling of promoters found is far from random, so it is premature to attempt an unbiased analysis of the M. jannaschii promoter sequences. However, analyses of the promoters from the 70 clones sequenced in the first round of selection and screening [representing 20 distinct intergenic regions (Table S1)] were interesting. BRE and TATA box sequences that could be recognized by eye or by computer-aided search with MEME (43) were aligned (Table S2). A sequence Logo (44) was generated in each of 2 ways. First, the individual sequences were weighted by the number of times they were isolated, giving emphasis to the strongest binding sequences (Figure 4B). Second, the consensus was calculated with each sequence counted equally (Figure 4A). The combined height of all letters stacked at a position is the information content at the position in bits (see Materials and Methods section). The relative heights of the letters stacked at a position reflect the relative frequency of the corresponding nucleotides.

The BRE and TATA box are obvious. The observed TATA box consensus of TWTATATA (W = A or T) differs little from the previously proposed TTTATATA consensus for methanogens (29). However, the Logos show conservation amongst these promoters that continues at least four, and perhaps seven or nine, residues beyond the 3' end of the canonical TATA box. This is most obvious in Figure 4B, but the heavy emphasis on the 15 tRNA promoter sequences makes interpretation beyond four positions questionable.

These promoters also display an unusually long BRE consensus, spanning 9–10 nucleotides (MRCCGAAAAG, where M = A or C, and R = A or G), rather than the more usual 6–7 (18,20, but compare 26). Because of the A + T-richness of the *M. jannaschii* genome, the random chance of a C or a G is lower than that of an A or a T, so a conserved C or G can be more informative. In Figure 4, this is reflected in the greater heights of some conserved C's and G's than of conserved A's and T's.

Another feature of some promoters in Archaea and Eucarya is a short initiator element (Inr), which includes the start site of transcription (26,27). Previous *in vitro* studies of the  $P_{tRNAVal}$  showed that changes in this



**Figure 3.** Scatter plot of recombinant plasmids harboring transcription factor-binding genomic fragments. The axes represent relative affinity and hybridization strength. (I) High affinity and high hybridization; (II) Low affinity and high hybridization; (III) Low affinity and low hybridization; (IV) High affinity and low hybridization.

element, especially the start site residue, impaired promoter function (19). When aligned on the Inr (Table S3), our clones display a TGC consensus, with greatest conservation of the G, which is the start of the transcript (Figure S1). There are 19-24 bp of A + T-rich sequence between the canonical TATA box and the Inr (right side of Figure 4 and left side of Figure S1).

#### DISCUSSION

## Genome-wide selection of archaeal promoters

We developed a genome-wide method to experimentally identify promoter sequences in *M. jannaschii*, based on the fact that archaeal promoter DNA is readily complexed with general transcription factors *in vitro* to form stable complexes that can be separated from unbound DNA by gel electrophoresis. The procedure was inspired by *in vitro* selection and evolution protocols (33–40), and it provided a simple and quick approach to isolate protein-bound DNA since no cross-linking or protein-specific antibody was required.

As in previous genomic SELEX work (35–40), we used a small-insert (200–300 bp) genomic DNA library; therefore, our pool of potential transcription factor-binding sequences was derived from an organism. This allowed the selection of only natural sequences, as opposed to 'optimally interacting sequences'. The pitfalls inherent in searching for optimal binding sequences are apparent in previous studies. For example, 'best' –10 and –35 motifs for *E. coli* sigma factor  $\sigma^{S}$  were derived by SELEX using randomized DNA oligonucleotides; however,



Figure 4. Sequence Logo generated from aligned promoter sequences. (A) Logo of promoter sequences from the initial selection. (B) Same as A, but weighted for the frequency at which each sequence was selected. (C) Logo based on all alignable promoter sequences in this study.

transcription from an artificial promoter containing them was weaker than from a promoter with the -35consensus only (50). Similarly, in studies using SELEX to identify Lrp protein binding sites in *M. jannaschii*, the derived 'optimal' binding sequences performed poorly in identifying the natural binding sites for the proteins in the genome (51). Clearly, natural binding sites are not selected for maximal binding.

Some important modifications were adopted in our version of the affinity-based selection to improve efficiency. We used a simplified protocol to construct the genomic library (40). Genomic DNA fragments were size-fractioned and blunt-end ligated into the vector. The quality of the library was demonstrated by the independent isolation of multiple genomic DNA fragments covering the same region (Table S1). In addition, we performed only one round of binding, separation and recovery. Typically, in SELEX and genomic SELEX, multiple rounds are performed, resulting in the selection of sequences with progressively higher affinity, while lower-binding affinity sequences are effectively eliminated. In contrast, we reasoned that omitting repeated rounds of selection would allow us to recover a broader spectrum of promoter sequences, with varying affinities to the transcription factors; one potential problem associated with this strategy is that the background of (presumptive) false positives is increased. Subsequent screening was performed to isolate fragments with high affinity to the transcription factors, i.e. potential promoters from this archaeon. Criteria for the identification of putative promoters were based on genome context, as well as the frequency of recovered genomic DNA fragments (Figure S2).

Conserved motifs from the natural promoters were readily identified when recovered DNA sequences were aligned (Figure 4). In *M. jannaschii*, the TATA box (the binding site for TBP) is highly conserved, and nearly symmetrical as well, in agreement with the fact that the DNA-binding domain of archaeal TBP contains more nearly perfect repeats than does eukaryal TBP (29,52). This symmetry renders the TATA box alone incapable of determining the orientation of the transcription complex. Essential information regarding orientation is provided by the BRE (25), which also shows an extended region of conservation vis-a-vis its eukarval element. The extended BRE indicates that the core promoter is more complex than expected. Basespecific cross-linking in the pre-initiation complex revealed close contact between this region and the C-terminal region of TFB (23). Cytosine at -7 (relative to TATA box), the most conserved position in the BRE extension, was identified as a preferred base in the selection of randomized DNA oligonucleotides using archaeal TBP and TFB (20). Interestingly, similar results (though with a conserved G instead of C at -7) were obtained when human TBP and TFIIB were used, indicating that this base may contribute to the interaction with TFIIB in eukaryal promoters as well (53). Moreover, a 3D structure of the human ternary TBP/TFB/DNA complex confirmed that G at position -7 makes watermediated contacts with TFIIB (54). The conservation of position -7 across the archaeal-eukarval lineage strongly suggests the relevance of this position, likely in mediating contact with TFB/TFIIB in both domains. Although we cannot assign a functional role to the other three bases in the extended BRE, their conservation suggests the extended BRE is an integral part of archaeal promoters. Interestingly, the extended BRE is most conserved in promoters with particularly high affinity for transcription factors; this may be relevant in recognizing functional distinctions among promoters. The extended consensus does not necessarily mean an extended binding site for these two factors; other molecules could bind the region.

Given the compact nature of the M. jannaschii genome it was not surprising that, in most cases, promoters were found close to the start site of a downstream ORF or RNA gene. We used this feature to define putative small ncRNAs. If the gap between a promoter and the downstream gene was large, and the G + C content in the region was higher than the average for the genome, these promoters were tentatively assigned to previously unidentified ncRNA genes. In thermophiles, high G + Ccontent is a significant feature of structured RNAs, such as tRNAs and rRNAs. In silico studies have utilized the presence of high G + C islands to identify ncRNA genes in organisms with overall low genomic G + C content (47,48). We identified 10 presumptive ncRNA genes and their associated promoters from 27 gel shifted genomic DNA fragments. Four of these RNAs had previously been identified experimentally (47). Our results suggest that a systematic search for promoters is an effective strategy to identify novel transcripts. Further characterization of these ncRNA genes and their promoters is in progress.

# Experimental concerns of *in vitro* selection of natural promoters

The dramatic difference between the isolation frequency of tRNA gene promoters and protein gene promoters makes it important to consider the sources of bias in the procedures employed, and how they might be circumvented. Foremost among our concerns is the effective concentration of the transcription initiation factors. The DNA-binding reactions prior to electrophoretic separation were  $\sim 0.5 \,\mu\text{M}$  in each of the proteins. With the relative instability of TFB, we estimate that its effective concentration by the end of the incubation is  $<0.1 \,\mu\text{M}$ , about 20-fold lower than estimates of its in vivo concentration (Ying Jiang, C.I.R. and G.J.O., unpublished data). Given the demonstration that different promoters have different intrinsic affinities for the transcription factors (based on the competition assays), we presume that we are far below saturation binding of transcription factors to DNA. Although other factors could influence the distribution of promoters recovered, this effect is consistent with all of our observations. The most obvious solutions are to increase the stability of TFB by use of the TFBc fragment (as was done in later selections, although this could potentially introduce other biases), and to use higher concentrations of TFB (increasing solubility and other logistical problems).

A related issue is the possibility that transcription factor-DNA complexes formed, but that they dissociated during the electrophoretic separation. In particular, this could be aggravated by the use of a gel with (nominally) equal concentrations of  $Mg^{2+}$  and EDTA, in contrast to the binding reaction conditions in which  $Mg^{2+}$  is in excess. Although we have not observed decreases in shifted DNA with changes in gel conditions, nearly all of these observations are based upon the shifting of the strongest-binding promoters. Thus, it is possible that some classes of promoters (presumably those with lower affinity of the transcription factors) dissociate in the gel, and hence are not shifted. We have no direct measure of this, but we do observe a strong correlation between the efficiency with which weakly binding DNA fragments are mobility-shifted (as a function of transcription factor concentration) and the efficiency of the same fragment in competing for transcription factors in the binding reaction (measured by the gel shifting of a stronglybinding promoter).

Several of the isolated DNA regions were intragenic. We have already cautioned against classifying them as false positives. In organisms with compact genomes it is not unusual to find promoters inside genes. A comprehensive survey of 791 E. coli mRNA promoters indicated that 18% of them were positioned inside the preceding genes (55). Protein ORFs can also host, or overlap with, independently transcribed genes (particularly ncRNAs) as well. A variety of small non-messenger RNAs have been identified in a specialized cDNA library of the archaeon Archaeoglobus fulgidus (56); some of these RNAs are complementary to an ORF, some overlap an ORF and some are even internal to an ORF. In vitro, most of our 'false positives' were demonstrated to compete with bona fide promoters for transcription factors. While suggestive of in vivo binding and potential promoter activity, other explanations are possible, and our data are unable to distinguish among them.

Our protocol involves two PCR amplification steps, one before the promoter selection step and one after. At the end of the procedure, the combined errors of the two amplifications and the single-strand sequencing of recovered clones was  $\sim 2.5$  errors per 1000 bp DNA relative to the published genome sequence. Of these steps, only PCR errors introduced before the gel selection (at most 1.2 per 1000 bp, or about 1 per 4 genomic DNA fragments) could bias the selection. Only three selected clones had a sequence error in the presumed TATA or BRE, and these were excluded from the analyses reported above. Overall, there is no evidence that the PCR errors were sufficiently numerous to corrupt authentic promoters (false negatives) or create spurious promoterlike sequences (false positives).

A more problematic artifact was the creation of chimeric sequences. Overall, 12 (3.7%) of the sequenced clones included DNA from non-adjacent portions of the genome (again, they were excluded from the results reported). This artifact can be introduced either during the ligation step of library construction, or during the PCR amplification steps. Kanagawa (57) proposed two modes of chimera formation during multi-template PCR, either by template switching or premature termination of extension. Consistent with premature termination of extension, examination revealed homologous overlapping sequences at the junction point. Although this artifact was annoying, we cannot see any mechanism by which it could have altered our major observations. This said, PCR sequence errors could be reduced by decreasing the PCR cycles and using a proofreading polymerase; chimeras could be reduced by decreasing the PCR cycles and optimizing efficiency to minimize premature termination.

## SUMMARY AND IMPLICATIONS

In conclusion, we report an experimental genome-wide approach to identify promoters; the procedure is based on the fact that promoter sequences are substrates for transcription factor binding, and thus could be selected in a manner reminiscent of SELEX. Our efforts resulted in the selection of a subset of promoter regions in the genome, most of them for genes that are highly expressed. Information extracted from these sequences allowed us to refine our knowledge of the composition and architecture of promoters in this organism, and to infer that strict adherence to canonical promoter elements (and their extensions) is a key feature of highly transcribed genes. We view this as a test case for the application of EMSA to a whole-genome screen of binding sites for a variety of proteins. The approach can be applied to any protein-DNA complex that is sufficiently stable to gel-shift the binding-site-containing DNA fragment.

## SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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