

NIH Public Access

Author Manuscript

J Mol Biol. Author manuscript; available in PMC 2012 January 7

Published in final edited form as:

J Mol Biol. 2011 January 7; 405(1): 125–142. doi:10.1016/j.jmb.2010.10.049.

Temporal regulation of gene expression of the *Thermus thermophilus* bacteriophage P23-45

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Abstract

Regulation of gene expression during infection of the thermophilic bacterium Thermus thermophilus (T. th.) HB8 with the bacteriophage P23-45 was investigated. Macroarray analysis revealed host transcription shut-off and identified three temporal classes of phage genes: early, middle, and late. Primer extension experiments revealed that the 5' ends of P23-45 early transcripts are preceded by a common sequence motif that likely defines early viral promoters. T. th. HB8 RNA polymerase (RNAP) recognizes middle and late phage promoters in vitro but does not recognize early promoters. In vivo experiments revealed the presence of rifampicin-resistant RNA polymerizing activity in infected cells responsible for early transcription. The product of the P23-45 early gene 64 shows a distant sequence similarity with the largest, catalytic subunits of multisubunit RNAPs and contains the conserved metal-binding motif that is diagnostic of these proteins. We hypothesize that ORF64 encodes rifampicin-resistant phage RNAP that recognizes early phage promoters. Affinity isolation of T. th. HB8 RNAP from P23-45-infected cells identified two phage-encoded proteins: gp39 and gp76, that bind the host RNAP and inhibit in vitro transcription from host promoters, but not from middle or late phage promoters, and may thus control the shift from host to viral gene expression during infection. To our knowledge, gp39 and gp76 are the first characterized bacterial RNAP-binding proteins encoded by a thermophilic phage.

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Keywords

Thermus thermophilus; thermophage; phage promoters; RNA polymerase; RNAP-binding proteins

Introduction

Transcription is the first step and primary regulatory determinant of gene expression. Multisubunit DNA-dependent RNA polymerases (RNAPs) are complex, highly regulated molecular machines. RNAP, alone or in complex with regulatory factors, is central to the transcription process. Every stage of transcription is regulated via RNAP interactions with various transcription factors. Bacteriophages (phages) have evolved highly effective mechanisms to modify bacterial RNAP to serve the needs of the phage (reviewed in 1). Recent studies indicate that phages are the most abundant life form in the Biosphere ^{2; 3}, and that the phage gene pool is the largest source of natural gene diversity. Consequently, while many phages use common strategies to subjugate their hosts, the number of phage-encoded regulatory mechanisms is virtually endless⁴. At the time of writing, more than 580 complete phage genome sequences (NCBI, last modified July 2010) had been determined. Comparative genomic analysis provides important insights into the diversity and evolution of phages and their hosts. However, our understanding of gene expression regulation strategies utilized by phages is relatively limited. Classical studies of gene regulation mechanisms in a handful of model *Escherichia coli* (E. coli) phages (e.g. γ , T4, and T7) have led to key discoveries in molecular biology. More recently, transcription profiling and bioinformatic predictions have been successfully applied to aid our understanding of the molecular features of phage regulatory networks 5; 6; 7; 8. The ultimate understanding of transcription regulatory mechanisms is based on structure-function analysis of RNAP alone, and RNAP-regulator complexes. However, a high resolution structure of E. coli RNAP is absent, therefore, almost no structural information on the action of transcription factors encoded by model E. coli phages in complex with E. coli RNAP has been obtained. Currently, only the structures of bacterial RNAP from the thermophilic eubacteria Thermus aquaticus (T. aq.)^{9; 10; 11} and T. th. ^{12; 13} have been determined. Thus, at present, the most reasonable way to structurally investigate phage-driven prokaryotic transcription regulation is to study phages that infect thermophilic eubacteria (i.e. thermophages). Despite the advances in our understanding of the structures of thermophilic eubacterial RNAPs, there is insufficient knowledge about thermophages, in particular their biology and the gene regulation mechanisms they employ during host bacterial infection. The genomes of several thermophages infecting *Thermus* species have been completely sequenced ^{14; 15; 16}, and the gene expression strategy of one of these phage, φ YS40, has been investigated in detail⁸.

In this work we studied the temporal regulation of transcription of another *T. th.* HB8 phage, P23-45, whose genome has recently been sequenced ¹⁵. We identified three temporal classes of P23-45 genes and their corresponding promoters by a combination of gene macroarray, *in vivo* primer extension, and *in vitro* transcription experiments. P23-45 middle and late promoters have consensus elements that differ from those of *T. th.* housekeeping promoters and φ YS40 thermophage promoters ⁸. Yet, P23-45 middle and late promoters are recognized by unmodified host *T. th.* RNAP *in vitro* or *in vivo*. In contrast, P23-45 early promoters are not recognized by host *T. th.* RNAP either *in vitro* or *in vivo*. The early promoters are defined by an unusual 11 bp conserved sequence motif, which is likely recognized by a unique phage-encoded RNAP. Affinity isolation of *T. th.* RNAP from P23-45-infected cells identified two phage-encoded *T. th.* RNAP-binding proteins: gp76 and gp39, the products of an early and a middle gene, respectively. These proteins bind the host RNAP *in vitro* and efficiently inhibit transcription from host bacterial promoters but not from middle or late phage promoters. Thus, these proteins may be responsible for the shut-off of host

transcription and therefore the switch from host to viral gene expression. To our knowledge, this is the first description of thermophage-encoded thermophilic bacterial RNAP-binding proteins.

Results

Macroarray analysis of gene expression during P23-45 infection

Almost half of the P23-45 genome (ORFs 1-78) is transcribed in the same direction (leftward in Fig. 1A). These genes form a single cluster in the left arm of the genome. The remaining P23-45 ORFs (ORFs 79–117) are transcribed in the rightward direction. These ORFs form a single cluster at the right arm of the genome. The only exception is ORF5, which is transcribed in the rightward direction, but is located in the left arm of the genome (Fig. 1A). To characterize the temporal profile of P23-45 gene expression, a macroarray of P23-45 phage genes was prepared. The array contained spots with equal amounts of PCRamplified DNA fragments of 20 P23-45 genes and one non-coding region of the P23-45 genome (marked by black dots in Fig. 1A and B). The genes chosen for the array encode proteins of different functional classes. One group of spots represented the abundance of mRNA from a cluster of small genes with unknown functions from the left arm (genes 57, 68, 69, 76, 78 and the non-coding region of triplex-forming mirror repeats ¹⁵). The second group of spots represented left arm genes involved in DNA replication, recombination, and nucleotide metabolism (genes 4, 5, 11, 14, 24, 39, and 46). The third group of spots on the array represented right arm genes that encode the P23-45 virion structural proteins (genes 82, 89, 94, 96, and 114), and predicted DNA packaging (gene 85) and lysis proteins (genes 108 and 112). Since closely spaced or partially overlapping genes are usually transcribed from the same promoter, some spots on the array may likely report the abundance of transcripts of multiple P23-45 genes.

In order to determine whether P23-45 shuts off host transcription, PCR-amplified DNA fragments of seven housekeeping *T. th.* HB8 genes: *rpoC* (encoding the RNAP β' subunit), *sigA* (encoding the primary sigma factor σ^A), *dnaK* (encoding a chaperone), *TTHA0466* (encoding alcohol dehydrogenase), *infB* and *infC* (encoding translation initiation factors IF2 and IF3, respectively), and *rpsA* (encoding the ribosomal protein S1) were spotted on the membrane. The array also contained spots with total *T. th.* HB8 genomic DNA, P23-45 genomic DNA, and a PCR-amplified DNA fragment of the *Drosophila melanogaster* (*D. me.*) *zfrp8* gene that was used as a normalizing and loading control.

T. th. HB8 cells were infected with P23-45 at a multiplicity of infection of 10 and total RNA was extracted 5, 20, and 40 minutes post-infection. As a control, RNA was extracted from T. th. HB8 cells immediately prior to P23-45 infection. Equal amounts of total RNA were used to generate radioactively labeled cDNA by random priming/reverse transcription followed by hybridization to the macroarray membrane. The amount of radioactivity hybridized to different spots of the macroarray reflected the abundance of transcripts of each corresponding gene. For each time point, three independent macroarray experiments were performed. To quantitatively analyze the macroarray data, the radioactive signal from each spot was corrected for background and normalized according to the relative strength of the signal from the D. me. zfrp8 spot. The mean amount of radioactivity for each macroarray spot was plotted as a function of time post-infection (Fig. 2). As expected, the total amount of P23-45 transcripts increased with time post-infection relative to the control zfrp8 spots (Fig. 2A). In contrast, the total amount of T. th. HB8 transcripts decreased throughout the same period (Fig. 2A), indicating that P23-45 either executes host transcription shut-off or increases the rate of host RNA decay. The behavior of individual host transcripts during P23-45 infection was complex; while the abundance of most transcripts decreased throughout the infection cycle, the rates at which the transcripts decreased varied and in one

To compare the behavior of individual P23-45 transcripts, plots of normalized spot signal intensities as a function of time post-infection were scaled to equalize the mean abundance of each transcript. The accumulation of individual transcripts peaked at different times during the infection cycle, indicating the presence of different temporal classes of phage genes (Fig. 2B). Phage genes were clustered based on the time when their transcripts became most abundant ¹⁷. Analysis of transcript abundance patterns revealed three distinct temporal classes of genes: early, middle, and late (Fig. 2B). The early class gene transcripts peaked 5 minutes post-infection, while middle class gene transcripts peaked 20 minutes post-infection. Finally, the abundance of transcripts of the late class of genes increased dramatically by the end of infection (in our conditions the eclipse period of P23-45 was 35 minutes and lysis of *T. th.* HB8 by the phage was complete ~ 60 minutes post-infection). The average value of the scaled abundances calculated for each of the three temporal classes are shown as separate panels in Figure 2C.

P23-45 genes used in the macroarray analysis and the temporal classes they belong to are indicated by black dots and different colors, respectively, in Figure 1. Most centrally located genes with unknown functions belong to the early class, however, gene 78 and the non-coding triplex-forming region located in this part of the P23-45 genome belong to the middle and the late classes, respectively. Some genes encoding DNA replication and recombination components also belong to the early temporal class. Other genes from this functional group, as well as genes encoding nucleotide metabolism enzymes, comprise the middle class. As expected, genes of the late class were found exclusively in the right arm of the genome. They encode the P23-45 virion structural proteins, DNA packaging proteins, and lysis proteins.

Mapping P23-45 promoters

The previous automated annotation of P23-45 genes suggested that P23-45 does not encode its own RNAP ¹⁵ and must therefore rely on host RNAP to transcribe its early, middle, and late genes throughout the infection. Such temporal regulation can be achieved either by using specific sequences defining promoters of different temporal classes and/or by modification of RNAP promoter specificity by phage-encoded transcription factors. In the following section, we report our analysis of the P23-45 middle and late promoters followed by a discussion of the analysis of the P23-45 early promoters.

P23-45 middle and late promoters—Non-coding P23-45 regions upstream of middle and late phage genes were examined by primer extension. Five primer extension products corresponding to the 5' RNA ends of three middle (P_4 , P_{35} , and P_{39}) and two late (P_{80} and P_{103}) putative promoters were detected. Additionally, one middle promoter, P_{68M} , located in the early gene cluster was also identified by primer extension. The kinetics of primer extension product accumulation for these promoters during P23-45 infection matched the macroarray data for middle and late genes. Representative primer extension experiments with primers specific to middle and late genes are shown in Figure 3A.

To confirm that the *in vivo* identified 5' RNA end points are transcription start points, we tested the ability of P23-45 genomic DNA fragments containing putative middle and late P23-45 promoters to serve as templates for *in vitro* abortive transcription initiation with purified *T. th.* HB8 σ^{A} -associated holoenzyme. For each promoter tested, combinations of nucleotide substrates that should have permitted transcription initiation (based on the results)

of *in vivo* primer extension analysis) were used. In all cases, robust transcription was detected (for example, see Fig. 5B, lanes 14 and 15). Therefore, we conclude that *in vivo* primer extension products correspond to transcription start points and that unmodified host RNAP- σ^A holoenzyme can recognize P23-45 middle and late promoters *in vitro*.

Comparisons of sequences upstream of the transcription start points of P23-45 middle and late transcripts revealed a motif that was common to promoters of both classes (Fig. 3B). The middle and late P23-45 promoters are characterized by a -10-like element (consensus sequence 5'-GTATanT-3') with the highest conservation at positions -11 (A) and -7 (T) relative to the experimentally determined transcription start point (Fig. 3B). In addition, an extended -10 "TG/TGTG" motif is present 0-2 bp upstream of the -10 element. In two cases, appropriately positioned motifs similar to the consensus *T. th.* -35 promoter element were identified in two middle phage promoters, P₃₅ and P₃₉ (Fig. 3B, for the consensus *T. th.* -35 promoter sequences may be due to the small number of promoter sequences examined. Alternatively, the distinct temporal patterns of activity from these promoters may be caused not by differences in basal promoter elements or by binding of transcription factors to specific regulatory sites, but by the phage-dependent modification of host *T. th.* HB8 RNAP and/or differences in intrinsic promoter strengths.

P23-45 early promoters—Based on previous studies of other phages, one would expect that transcription of early P23-45 genes should be driven by strong promoters recognized by the housekeeping form of the host RNAP holoenzyme: *T. th.* HB8 σ^A -associated holoenzyme. Early phage promoters need to be strong to efficiently compete with host promoters for host RNAP. Thus, early phage promoters would be expected to have a good match to the host σ^A -associated holoenzyme consensus promoter sequence. Indeed, such an expectation was fulfilled by the phage ϕ YS40, a *T. th.* HB8 phage previously studied in our laboratory ⁸.

We utilized a *T. th.* RNAP- σ^A holoenzyme promoter bioinformatic profile described in ⁸ to search P23-45 DNA upstream of early P23-45 genes. To our surprise, with the exception of a likely σ^A -dependent promoter upstream of gene 68 located in the middle of the early cluster (P_{68E}), no high-matching candidate sequences were found. Visual inspection of non-coding regions separating P23-45 early genes revealed the presence of a common 11 bp sequence motif (5'-TTATTCcTTTA-3') located immediately upstream of annotated start codons (Fig. 4). Copies of this motif were identified upstream of early genes 59–61, 63, 64, 67–69, and 71–77. No additional copies of the 11 bp motif are present in any other region of the P23-45 genome. The logo of the motif is shown below the alignment in Figure 4 and compared to the *T. th.* promoter consensus logo. As can be seen, the two logos are clearly distinct from each other.

We hypothesized that this 11 bp motif may define early P23-45 promoters. To test this hypothesis, RNA samples used in macroarray experiments were subjected to primer extension analysis using primers specific to eight genes from the early gene cluster (see the alignment in Fig. 4, the experimentally identified 5' ends are underlined). In all cases, an identical result was obtained: a primer extension product whose end corresponds to the last nucleotide of the 11 bp motif. Moreover, the kinetics of primer extension product accumulation for early P23-45 genes was in agreement with the macroarray data. Representative primer extension experiments with primers specific to genes 64 and 68 are shown in Figure 5A.

The results presented above suggest that the 11 bp motif may define the early P23-45 promoters that are obviously distinct from known host or phage promoters. Alternatively,

the 11 bp motif could be the site of post-transcriptional processing of an early P23-45 polycistronic transcript(s) that initiates elsewhere upstream. The following experiments were performed to distinguish between these possibilities. First, putative P23-45 early promoters whose transcription start sites had been identified in vivo were tested in an in vitro abortive transcription initiation assay using purified T. th. HB8 RNAP- σ^A holoenzyme. For each putative promoter tested, combinations of nucleotide substrates that should have permitted transcription initiation (based on the results of in vivo primer extension analysis) were used. Unexpectedly, T. th. RNAP- σ^{A} holoenzyme was either unable to transcribe or yielded only small amounts of product for almost every transcription template tested (Fig. 5B; lanes 1-3 and 6–13); the only exception was P_{68E} (lanes 4–5). However, robust transcription from this promoter could be explained by the fact that the 11 bp motif is embedded in a recognizable σ^{A} -dependent promoter (as revealed by bioinformatic analysis; Fig. 4 alignment, the -35like and -10-like elements are underlined). For comparison, T. th. HB8 RNAP- σ^A holoenzyme actively transcribed from P23-45 middle and late promoters (P_{68M} and P₁₀₃, see Fig. 5B, lanes 14 and 15). Therefore, we conclude that unmodified host T. th RNAP- σ^A holoenzyme is unable to initiate transcription from DNA fragments containing the 11 bp motif in vitro.

If processing of an early polycistronic precursor transcript were responsible for the appearance of primer extension products whose 5' ends are located just downstream of the 11 bp motif, then transcription initiation of this precursor transcript most likely originate in the non-coding region of the phage genome that separates the divergently transcribed early and late gene clusters (Fig. 1). This region, between genes 77 (an early gene) and 80 (a late gene) contains an 11 bp motif upstream of gene 77. In an effort to identify a hypothetical early P23-45 promoter located upstream of gene 77 and responsible for early viral transcription, the following experiment was performed. The intergenic region between P23-45 genes 77 and 80 was cloned into the T. th.-E. coli shuttle plasmid pMKE1¹⁸; to generate pMKE77-80, a plasmid containing the validated late promoter, P₈₀, and the putative divergent early promoter, P77 (Fig. 6A). T. th. HB8 was transformed with pMKE77-80 and the resulting strain was infected with P23-45 (Fig. 6A). Total RNA was extracted from infected cells at various times post-infection and in vivo primer extension analysis with primers complementary to plasmid sequences upstream and downstream of the P23-45 insert was performed. As a control, primer extension reactions with P23-45 phage genome-specific primers that reported the transcriptional activity of P_{77} and P_{80} located on the P23-45 genome were also performed (Fig. 6B). Control reactions revealed the expected late accumulation of the P80 transcript and the early accumulation of a transcript whose 5' end coincided with the last nucleotide of the 11 bp motif; i.e. P77 (Fig. 6B, two lower panels). A primer extension product corresponding to plasmid-located P₇₇ was absent from uninfected cells, peaked 5 minutes post-infection and steadily decreased afterwards (Fig. 6B, upper left panel). A primer extension product corresponding to plasmid-located late P_{80} was detectable in the absence of infection and continuously increased throughout the infection (Fig. 6B, right upper panel). The appearance of the P₈₀ transcript in the absence of the phage shows that this promoter is recognized by host RNAP in uninfected cells in vivo and is consistent with our findings that unmodified host T. th. RNAP initiates transcription from P₈₀ (and other P23-45 middle and late promoters) *in vitro*. The absence of this transcript immediately post-infection may be due to phage-dependent modification(s) of the host RNAP that regulates the coordinated temporal transcription program of P23-45 and prohibits premature recognition of late promoters via unknown mechanism. No primer extension product upstream of P77 was detected in uninfected cells. Our attempts to identify a hypothetical promoter located upstream of P₇₇ with several additional primers, both P23-45 phage genome-specific and pMKE77-80 plasmid-specific, were similarly unsuccessful (data not shown). These data make us conclude that it is very unlikely that there is a strong viral

promoter from which early P23-45 genes are transcribed by host RNAP to produce a single precursor transcript.

An "internal" putative early promoter P_{59} was also cloned into pMKE1 and the activity of this promoter in P23-45 infected and uninfected cells was monitored. An identical result to that described for P_{77} (no activity in uninfected cells and an "early" pattern of activity during infection) was observed (data not shown). Taken together, these data suggest that despite the lack of *in vitro* activity, P_{59} and P_{77} (and, likely, other genomic sites defined by the presence of the 11 bp motifs) may function as early phage promoters that are recognized immediately after infection by either a phage-encoded, but as yet unidentified RNAP, or a phage-modified host RNAP.

Transcription of P23-45 early genes is Rifampicin-resistant—Rifampicin (Rif), a strong inhibitor of bacterial RNAPs, including the *T. th.* RNAP, binds to a pocket formed by the RNAP β subunit and efficiently blocks the synthesis of transcripts longer than 2–3 nt ¹⁹. Conversely, all phage-encoded RNAPs studied thus far are resistant to Rif ^{6; 20}. Therefore, if early P23-45 genes were transcribed by host RNAP, then the appearance of these transcripts shall be suppressed by the addition of Rif. On the other hand, if a phage RNAP were responsible for early P23-45 transcription, the appearance of these transcripts shall not be affected by Rif.

We performed P23-45 infection followed by the addition of Rif at different time points of infection followed by additional 10 minutes incubation and primer extension analysis of selected phage transcripts. Since middle and late P23-45 mRNAs are transcribed by host RNAP, we expected that Rif will inhibit accumulation of these transcripts. This expectation was fulfilled (Fig. 7, panels B and C): the addition of Rif 5, 10, and 20 minutes postinfection inhibited accumulation of a middle (P₃₉) and late (P₈₀) promoter transcripts compared to a control infection that was not treated with Rif. The effect was less pronounced when Rif was added 20 minutes post-infection, since by this time P_{39} and P_{80-} originated transcripts started to accumulate prior to Rif addition. For 5 and 10 minutes time points, Rif addition led to complete disappearance of primer extension bands corresponding to either P₃₉ or P₈₀. In contrast, the addition of Rif had a completely different effect on the abundance of primer extension product corresponding to the early P_{64} phage transcript (Fig. 7, panel A). The addition of Rif 5 or 10 minutes post-infection led to increase of this transcript abundance compared to untreated control cells. Addition of Rif 20 minutes postinfection, when early transcription ceases based on the kinetics of early transcript accumulation, had no effect on the P64 transcript abundance. The difference in the abundances of the P64-originated transcript in total RNA extracted from Rif-treated and Rifuntreated cells may be explained by increased proportion of the P_{64} transcript in Rif-treated cells, where host RNAP-dependent transcripts are not accumulating. We conclude that the synthesis of early P23-45 transcripts is Rif-independent and may therefore be due to activity of Rif-resistant RNAP that is distinct from Rif-sensitive host RNAP.

Identification of P23-45-encoded T. th. RNAP-binding proteins

Based on our macroarray data, P23-45 executes shut-off of *T. th.* HB8 transcription, expresses its genes in a coordinated manner, and must rely on the host transcription machinery for the expression of middle and late genes. Therefore, we hypothesized that P23-45 may encode transcription factors to alter *T. th.* HB8 RNAP promoter specificity and activity. To identify such proteins, we affinity isolated *T. th.* HB8 RNAP genomically-tagged with a protein A (4PrA) tag from *T. th.* HB8 cells infected with P23-45 and identified RNAP co-isolating proteins in the sample using the MudPIT technique ²¹ (Fig. 8). As a control, analysis of proteins affinity isolated from P23-45-infected wild-type *T. th.* HB8 cells

(untagged RNAP) was performed. Proteins present in both the *T. th.* HB8/P23-45-infected (untagged) and the *T. th.* HB8 RNAP-4PrA/P23-45-infected (tagged) samples were filtered out of the data set obtained for the RNAP-4PrA tagged strain.

Analysis of the material affinity isolated from the T. th. HB8 RNAP-4PrA/P23-45-infected cells revealed the presence of the core RNAP subunits ($\alpha\beta\beta'\omega$) and the primary σ factor, σ^A . The RNAP subunits and σ^{A} were present at stoichiometric levels, as estimated by normalized spectral counts (Fig. 8), and another protein present at stoichiometric levels was CarD (TTHA0168). In a recent study, it was demonstrated that T. th. HB8 CarD could interact with the N-terminus of the RNAP β subunit in a bacterial two-hybrid assay ^{22, 23}. Several known transcription elongation and anti-termination factors (NusA, NusG, and GreA), the transcription-repair coupling factor (TRCF), the nucleoid associated protein HU and the exonuclease ABC subunit A (UvrA) were also detected, although these proteins were present at lower levels. An uncharacterized protein, TTHA1350, was also identified; although TTHA1350 was detected at low levels, this result indicates that TTHA1350 may play a role in the T. th. HB8 transcription cycle. In addition to the host bacterial proteins, two P23-45-encoded proteins: gp39 (16.2 kDa; detected at a level substoichiometric to the core RNAP subunits) and gp76 (5.8 kDa; detected at a level stoichiometric to the core RNAP subunits) were identified (Fig. 8). To corroborate the affinity isolation results, the DNA encoding these two proteins was cloned into an E. co. pET28-derived expression vector ²⁴ and recombinant gp39 and gp76 proteins were purified to homogeneity and assayed for their ability to bind to host T. th HB8 RNAP core and σ^{A} -associated holoenzymes (Fig. 9A). The phage proteins were incubated with T. th. RNAP core or σ^{A} holoenzymes and the mixtures were resolved by native gradient PAGE (Fig. 9A, left panel). The results of subsequent denaturing SDS-PAGE indicated that gp39 and gp76 can interact with both the core and the σ^A -holoenzymes (Fig. 9A, for gp39 lanes 2 and 6, and lanes 2' and 6'; for gp76 lanes 3 and 7, and lanes 3' and 7'). Thus, we conclude that gp39 and gp76 are able to bind T. th. HB8 RNAP in vivo and in vitro. As can be seen from Figure 9, they can interact with RNAP simultaneously (Fig. 9A, lanes 4 and 8, and lanes 4' and 8') but do not interact with each other (ZB and LM, unpublished results), suggesting that they bind to distinct sites on RNAP. To our knowledge, this is the first documentation of thermophageencoded thermophilic host bacterial RNAP-binding proteins. Gp39 (a middle phage protein) and gp76 (an early phage protein) have no recognizable conserved motifs or similarities with other proteins in the public databases ¹⁵. Thus, they are novel bacterial RNAP-binding proteins.

In order to elucidate the possible role(s) of gp39 and gp76 in P23-45 infection, we tested their ability to influence transcription by the host RNAP. In vitro abortive transcription initiation reactions using DNA fragments containing three different T. th. HB8 σ^{A} dependent promoters (P_{rpoB-1} , P_{rpoB-2} , and P_{infB}) and two P23-45 promoters, a middle promoter (P_{68M}) and a late promoter (P_{103}) , were performed in the presence or in the absence of either gp39 or gp76 (Fig. 9B). As can be seen, both proteins efficiently inhibited transcription from the host bacterial promoters that belong to the -10/-35 promoter class (Fig. 9B, lanes 1–9). When added together, gp39 and gp76 demonstrated a strong additive effect in transcription inhibition at these promoters (data not shown). In contrast, both gp39 and gp76 were much less efficient at inhibiting transcription from the phage middle and late promoters that belong to the extended -10 class of promoters that lack the -35 promoter element (Fig. 9B, lanes 10–15, see also P23-45 middle and late promoter alignment in Fig. 3B). Thus, the binding of gp39 and gp76 to T. th. HB8 RNAP leads to promoter-specific transcription inhibition. Both proteins were purified as polyhistidine-tagged versions. To check if the tags may introduce non-physiological activities, we compared his-tagged and untagged (with his tag removed by thrombin) proteins and found that the tag does not interfere with their RNAP binding and transcription inhibition activities in vitro. Our data

suggests that gp39 and gp76 may be responsible for host transcription shut-off during P23-45 infection and may likely act by interfering with -35 promoter element-RNAP interactions. However, the molecular mechanisms underlying host transcription inhibition remain to be fully elucidated.

Discussion

In this work, we investigated the transcription strategy of P23-45, a lytic thermophilic siphovirus infecting the thermophilic eubacterium T. th. HB8. To study the P23-45 gene expression pattern, we used a combination of bioinformatic and biochemical methods, an approach that was successfully used by us to study host and viral gene expression during infection of T. th. HB8 by an unrelated thermophage, a large myovirus φ YS40⁸. In the case of P23-45, macroarray analysis and *in vivo* primer extension revealed that *i*) host bacterial transcription is shut-off during the P23-45 infection cycle and *ii*) three temporal classes of viral genes: early, middle, and late, exist. Most of the known phages do not encode their own RNAP for expression of early genes and must therefore rely on host RNAP to transcribe their genes. Early promoters of such phages tend to be very strong, with a good match to the host promoter consensus, to successfully compete with host promoters for host RNAP at the onset of infection. Initially, automated bioinformatic analysis of the P23-45 genome did not reveal a recognizable RNAP gene ¹⁵; leading us to hypothesize that P23-45 early promoters are similar to host promoters and are recognized by the host T. th. RNAP- σ^A holoenzyme. Contrary to this expectation, no early P23-45 promoters with homology to host σ^{A} -dependent promoters were also identified. Instead, we determined that many early phage genes are preceded by a common sequence motif. This highly conserved 11 bp motif has a consensus sequence 5'-TTATTCcTTTA-3', with the highest conservation at positions -9 (T), -8 (A), -7 (T), -6 (T), and -2 (T) relative to the experimentally determined 5' end of the transcript. In 13 of the 15 early P23-45 promoters, the 11 bp motif is located immediately upstream of the annotated translation start codon. Thus, many early phage transcripts appear to be leaderless. A similar situation was observed during the analysis of late and middle transcripts of the φ YS40 thermophage⁸.

Comparison of putative P23-45 early promoters with the *T. th.* -10/-35 consensus promoter elements used in bioinformatic searches indicated that they are clearly different (Fig. 4, compare logos). Thus, it was not at all surprising that *T. th.* RNAP- σ^A holoenzyme did not recognize these sequences as promoters *in vitro*, or in the absence of P23-45 infection *in vivo*. Nevertheless, DNA fragments containing the 11 bp motifs, when positioned on a *T. th.* plasmid, led to the appearance of "correctly" initiated RNAs that behaved as early transcripts. We take these results as a strong indication that the 11 bp conserved sequences define early phage promoters. Experiments conducted in the presence of host RNAP inhibitor Rif clearly show that early P23-45 transcription is resistant to Rif, while middle and late transcription, which is carried out by *T. th.* RNAP, is Rif-sensitive. The data suggest that P23-45 encodes a Rif-resistant RNAP that transcribes its early genes from promoters defined by the 11 bp consensus motif.

Given that biochemical data strongly suggest the existence of P23-45-encoded RNAP, a more thorough bioinformatic analysis of the P23-45 genome was undertaken. All known RNAPs are divided into two unrelated families based on sequence, structure, and subunit composition. One family includes large multisubunit RNAPs of bacteria, archaea, and eukaryotes ²⁵; ²⁶; ²⁷. The other family consists of small single-subunit RNAPs related to phage T7 RNAP and found in some bacteriophages, in mitochondria, and in chloroplasts ²⁸; ²⁹. The principal enzymatic activities are performed in both families of RNAPs via the same two-metal catalytic mechanism. All multisubunit RNAPs share a universal metal-binding signature motif: NADFDGD, in their largest subunits that together with additional

conserved domains forms the active site. All T7-related RNAPs have other catalytic motifs and conserved domains/amino acids involved in the active center formation. In principle, it would seem more likely that P23-45 would encode a single-subunit RNAP that is present in many other bacteriophages. However, despite careful searches, we did not observe identify any sequence similarity to single-subunit RNAPs in the P23-45 genome. In contrast, a BLASTP search ³⁰ with the ORF64 sequence used as the query retrieved as the best hit (after the closely related ortholog from bacteriophage P74-26) the A subunit of RNAP I from the stramenopile *Thalassiosira pseudonana* (*T. ps.*)

The detected region of similarity included a 65 amino acid segment which aligned with 32% amino acid identity and 47% similarity; the similarity was not statistically significant (expect value of 1.4). However, it was notable that the alignment encompassed a portion of the catalytic double-psi beta-barrel (DPBB) domain of RNAP, the most conserved portion of both the β' and β subunits of all multisubunit RNAPs that is directly involved in nucleotide polymerization ²⁶; ²⁷; ³¹. Moreover, the RNAP amino acid signature that includes the three invariant and essential aspartates required for the coordination of the two Mg^{2+} ions that participate in catalysis was fully conserved in ORF64 ([NA]AD[FY]DGD, the magnesiumchelated aspartates are underlined). When the ORF64 sequence was combined with the T. ps. RNAP sequence to generate a position-specific scoring matrix (PSSM), the second iteration of the PSI-BLAST search ³⁰ readily retrieved numerous RNAP sequences. Using the PHI-BLAST program ³², we found that, when the ORF64 sequence was compared to the non-redundant protein sequence database at the NCBI under the additional requirement that the signature Mg²⁺-binding motif was matched, the only retrieved sequences were those of RNAP subunits. Using the HHPred program ³³, we found that the ORF64 sequence produced the best hit with the DPBB domain of the *T. th.* RNAP β' subunit, with the E-value 0.091 when the Interpro collection of HMMs was searched. When the HHPred search was initiated with the isolated sequence of the predicted DPBB domain of ORF64, the same best hit was obtained, with a statistically significant E-value of 0.0031.

A multiple alignment of the putative DPBB domain of ORF64 and a representative set of eukaryotic, archaeal and bacterial RNAPs is shown in Figure 10. The alignment suggests that ORF64 contains a short version of the DPBB domain similar to the highly diverged β' homologs of RNAPs from baculoviruses and fungal mitochondrial plasmids ³¹. The DPBB domain consists of six β -strands ^{27; 31}. In addition to the (NA)D(FY)DGD signature motif, ORF64 retains several other invariant and essential amino acid residues of the DPBB domain such as the arginine at the end of S3 (corresponding to β' R704 in *T. aq.* RNAP), and the proline and leucine in the loop downstream of S3 (β' P706 and L708, respectively, in *T. aq.* RNAP) (Fig. 10).

We propose that ORF64 is a distant member of the multisubunit family of RNAPs. However, the putative P23-45 RNAP contains no counterparts to several other essential residues of the β' RNAP subunit, suggesting that this enzyme could be mechanistically distinct from the known RNAPs. In our previous work highly sensitive mass spectrometric analysis of pure P23-45 virions has not detected ORF64¹⁵. More recently, Western blot analysis revealed no ORF64 traces among P23-45 virion proteins using anti ORF64 polyclonal antibodies (Minakhin *et al.*, unpublished data). It remains to be determined whether ORF64 functions on its own or requires additional host or phage-encoded factors for activity.

Macroarray and *in vivo* primer extension analyses also revealed P23-45 middle and late genes and the corresponding P23-45 promoters upstream of these genes; at present we cannot distinguish between the middle and the late promoters based on their consensus element sequences. Similarly to φ YS40, the P23-45 middle/late promoters are characterized

by a -10 consensus element supplemented with a "TG/TGTG" motif (Fig. 3B). Middle and late promoters of P23-45 are recognized by unmodified *T. th.* HB8 RNAP *in vitro*; as these promoters are inactive early in infection, it follows that their activity is somehow repressed until later stages of the infection cycle.

A combined view that emerges from our work thus indicates that the P23-45 transcription strategy is a simplified version of the transcription strategy employed *E. coli* phage N4. In the case of N4 infection, early genes are transcribed by the Rif-resistant phage-encoded RNAP that is encapsulated in the virion and injected into the infected cell with the N4 genome ^{20; 29}. The middle genes are transcribed by another phage-encoded RNAP, a product of early N4 genes ³⁴. Late N4 genes are transcribed by phage-modified host RNAP. In the case of P23-45, early genes are likely transcribed by the Rif-resistant RNAP encoded by ORF64, while the middle and late genes are transcribed by host RNAP. Previous analysis did not reveal the presence of ORF64 in P23-45 virions ¹⁵. This could have been caused by a low copy number of the putative phage RNAP in the virion. Alternatively, ORF64 may be initially transcribed by the host RNAP- σ^A promoter (see alignment in Fig. 4, the -10/-35 elements are underlined). These possibilities are currently being investigated in our laboratory.

Using one-step affinity isolation of host *T. th.* HB8 RNAP and subsequent mass spectrometric analysis (MudPIT) of the affinity isolated sample, we identified two P23-45-encoded *T. th.* HB8 RNAP-binding proteins: gp76 and gp39. To the best of our knowledge, this is the first documentation of thermophage-encoded thermophilic bacterial RNAP-binding proteins. Gp76 and gp39 are encoded by early and middle P23-45 genes, respectively, and both proteins efficiently inhibit *in vitro* transcription by host RNAP from host promoters, but are less effective at inhibiting transcription from P23-45 middle and late promoters (Fig. 9B). Taken together, these data suggest that gp76 and gp39 may be involved in the shut-off of host transcription during the P23-45 infection program. Further biochemical and structural analysis of these proteins, in complex with *T. th.* RNAP, should make it possible to obtain a structure-based model of the action of a phage-encoded transcription regulator.

Experimental Procedures

Bacterial growth and phage infection

Bacteriophage P23-45 was generously provided by Dr. Michael Slater, Promega Corporation (Madison, WI). To isolate individual P23-45 plaques, 150 μ L of a freshly grown *T. th.* HB8 culture (OD₆₀₀~0.4), grown in the TB medium (0.8% [w/v] tryptone, 0.4% [w/v] yeast extract, 0.3% [w/v] NaCl, 1 mM MgCl₂, and 0.5 mM CaCl₂), was combined with a 100 μ L dilution of phage stock, incubated for 10 minutes at 65°C, plated in soft TB agar (0.75 % [w/v]), and incubated overnight at 65°C. To prepare a phage stock suspension, an individual plaque was picked and subjected to two more rounds of plaque purification. The phage lysate was prepared using a previously described procedure ¹⁵.

E. coli strains XL-1Blue (New England Biolabs) and BL21(DE3) (Novagen) were used for molecular cloning and recombinant protein expression, respectively.

Total DNA purification and molecular cloning

P23-45 genomic DNA was extracted with a Lambda Midi kit (Qiagen) using the procedure recommended by the manufacturer. *T. th.* HB8 genomic DNA was purified by extraction with phenol-chloroform and subsequent precipitation with ethanol.

Plasmids encoding either polyhistidine-tagged gp39 or gp76 for recombinant protein production and purification were constructed as follows: the DNA encoding gp39 or gp76 was PCR-amplified using primers that appended *NdeI* and *Eco*RI sites at the 5' and 3' ends of each gene, respectively. The resultant PCR products were cleaved with *NdeI* and *Eco*RI and cloned between the *NdeI* and *Eco*RI sites of a pET28a-based plasmid, creating pSKB2-39_{HIS} and pSKB2-76_{HIS}. The plasmid pMKE77-80 was constructed as follows: a 504 bp DNA fragment of the P23-45 genome comprising the divergent P₇₇ and P₈₀ promoters together with the proximal parts of ORF77 and ORF80 coding regions was PCRamplified using primers that appended *NdeI* and *Hind*III sites at the 5' and 3' end, respectively. The resultant PCR fragment was digested with *NdeI* and *Hind*III and cloned between the *NdeI* and *Hind*III sites of the *E. coli* – *T. th.-E. coli* shuttle plasmid pMKE1¹⁸. The resultant plasmid, pMKE77-80, was used in *in vivo* primer extension experiments.

Strain construction and affinity isolation of Protein A-tagged T. th. HB8 RNAP

A similar strategy to that used to construct a *T. th.* HB8 strain encoding a polyhistidine tag ([His]₁₀) appended to the 3' end of the *rpoC* gene (encoding the RNAP β ' subunit) was used to construct a *T. th.* HB8 strain encoding a genomic β '-Protein A (4PrA) fusion protein ⁸. The resultant strain, *T. th.* HB8*rpoC::4PrA*, demonstrated a slightly slower growth rate compared to wild-type cells, but was infected with P23-45 as efficiently as wild-type.

To prepare P23-45-infected biomass, wild-type *T. th.* HB8 or *T. th.* HB8*rpoC::4PrA* cells were grown at 65 °C in 4 L of TB medium until OD₆₀₀ ~ 0.35 and were infected with P23-45 at a multiplicity of infection of 10. Infection was ceased 20 minutes post-infection by rapidly cooling the samples in an ice water bath. Cells were harvested by centrifugation and washed once with ice-cold 10% (v/v) glycerol. Next, 1 mL of lysis buffer (20 mM Hepes [pH 7.5], 0.2 mg/mL phenylmethylsulfonyl fluoride (PMSF), 4 mg/mL pepstatin) was added to every 10 g of cell pellet, and the cells were frozen in liquid nitrogen. Both tagged and untagged *T. th.* HB8 cells were cryogenically lysed using the MM 301 Mixer Mill (Retsch; ²⁴) and stored at -80° C until use. Affinity isolation of RNAP-4PrA and co-isolating proteins from P23-45 infected *T. th.* HB8 cells was performed as described for *E. coli* RNAP and co-isolating proteins ²⁴; ³⁵.

MudPIT analysis

Elution of *T.th.* HB8 RNAP-4PrA and co-isolating proteins from the IgG conjugated Dynabeads (Invitrogen) was achieved with 0.5 M ammonium hydroxide and 0.5 mM ethylenediaminetetraacetic acid (EDTA). The eluted proteins were frozen in liquid nitrogen and evaporated to dryness in a SpeedVac (Thermo Savant). The dried protein pellets were denatured, reduced, alkylated, and digested with endoproteinase LysC (Roche Applied Science) followed by digestion with trypsin (Promega). The peptide mixtures were pressure-loaded onto triphasic microcapillary columns, installed in-line with a Quaternary Agilent 1100 series HPLC pump coupled to a Deca-XP ion trap tandem mass spectrometer (ThermoElectron) and analyzed via ten-step chromatography ³⁶. The MS/MS data sets were searched using SEQUEST ³⁷ against a database of 117 P23-45 predicted gene products, combined with 2238 protein sequences from *T. th.* HB8 as described previously ¹⁵. Lists of detected proteins were established and compared using DTASelect/CONTRAST ³⁸ as described previously ¹⁵. Protein levels across different samples were compared using Normalized Spectral Abundance Factor (NSAF) values ^{39; 40}.

Proteins

Polyhistidine-tagged *T. th.* HB8 core RNAP and σ^A were used in native gel electrophoresis protein-protein interaction and *in vitro* transcription assays. The proteins were purified essentially as described in ⁸. Either recombinant polyhistidine-tagged gp39 or gp76 were

produced as follows; the expression plasmids either pSKB2-39_{HIS} or pSKB-76_{HIS} were transformed into E. coli BL21 (DE3) cells, and transformants were selected in the presence of 50 μ g/mL kanamycin. Cultures (4 L) were grown at 37 °C to an OD₆₀₀ ~ 0.8 and recombinant protein overexpression was induced with 1 mM IPTG for 4 hours at 37 °C. Cells containing overexpressed recombinant proteins were harvested by centrifugation, and disrupted by sonication in buffer A (10 mM Tris-HCl [pH 8.0], 500 mM NaCl, 5 mM imidazole [pH 8.0], 5% glycerol, 0.2 mg/mL PMSF). Inclusion bodies were dissolved in buffer B (10 mM Tris-HCl [pH 8.0], 500 mM NaCl, 2 mM imidazole [pH 8.0], 7 M urea); loaded onto a 5 mL nickel-chelated Hi-Trap sepharose column (GE Healthcare) equilibrated in buffer B, and the column was washed with buffer B supplemented with 25 mM imidazole. The bound proteins, either gp39 or gp76, were eluted from the column with buffer B supplemented with 200 mM imidazole, and dialyzed against buffer C (20 mM Tris-HCl [pH 8.0], 50 mM NaCl, 0.5 mM EDTA). The proteins were loaded onto a MonoQ column (GE Healthcare) equilibrated in TGE buffer (10 mM Tris-HCl [pH 8.0], 50 mM NaCl, 1 mM EDTA, 5% [v/v] glycerol), eluted with a linear gradient of NaCl from 150 mM to 450 mM, dialyzed against buffer D (10 mM Tris-HCl [pH 8.0], 100 mM NaCl, 1 mM EDTA, 50% [v/ v] glycerol) and stored at -80°C.

Macroarray membrane preparation and data analysis

DNA fragments corresponding to each of the selected P23-45 ORFs, *T. th.* HB8 housekeeping genes and the *D. me. zfrp8* gene (control) were PCR-amplified from the corresponding genomic DNA using gene-specific primer pairs (the sequences of the primers are available from the authors upon request). Membrane preparation, cDNA synthesis, and macroarray hybridization were performed as described ⁷. After hybridization, the amount of radioactivity from each spot was quantified using the ImageQuant software (Molecular Dynamics) and the background signal was subtracted from signals corresponding to every ORF spot. To allow comparison between the signals on different membranes, the background-corrected signals were normalized relative to the average of the two *D. me. zfrp8* spot signals; the normalized spot signals were used for data analysis.

Primer extension

Primer extension reactions were performed essentially as described in our previous work⁸. Exponential phase T. th. HB8 cells were infected with P23-45 and harvested at the same time points post-infection as for the macroarray experiments. In experiments utilizing Rif, Rif (Sigma-Aldrich) was added to T. th. HB8 cells infected with P23-45 at the designated time points to yield a final concentration of 2 mg/mL followed by 10 minutes incubation prior to RNA extraction. Total RNA was extracted using the RNeasy Mini Kit (Qiagen) according to the manufacturer's procedure. For each primer extension reaction, 10 µg of total RNA was reverse-transcribed with 100 units of SuperScript III enzyme from the First-Strand Synthesis kit for RT-PCR (Invitrogen) in the presence of 10 pmol of γ -³²P endlabeled primer. The reactions were treated with RNase H, precipitated with ethanol and dissolved in formamide loading buffer. To identify the 5' ends of the primer extension products, DNA sequencing reactions, accomplished using the *finol* DNA Cycle Sequencing kit (Promega), containing both the correspoding PCR-amplified P23-45 genome fragments and end-labeled primers used for the primer extension reaction were performed. The reaction products were resolved on 6-8 % (w/v) polyacrylamide sequencing gels and visualized using a PhosphorImager (Molecular Dynamics).

Protein complex analysis

Either *T. th.* HB8 core RNAP or σ^{A} -holoenzyme (reconstituted with 1 μ M core RNAP and 1 μ M σ^{A}) was incubated with either gp39 (~ 5 μ M) or gp76 (~ 5 μ M) in 10 μ L of transcription buffer (30 mM Tris-HCl [pH 7.9], 40 mM KCl, 10 mM MgCl₂, 2 mM β -mercaptoethanol)

for 10 minutes at 65°C. Subsequently, 4 μ L of the reaction mixture was resolved on a native 4–15% (w/v) Phast gradient polyacrylamide gel (GE Healthcare); bands due to proteins were visualized by Coomassie blue staining. To interrogate the protein composition of the bands resolved by native gel electrophoresis, the bands were excised from the native gel and placed into the wells of a gradient 12–16% (w/v) gradient polyacylamide denaturing SDS gel, followed by electrophoresis and staining with silver.

In vitro transcription

A typical abortive transcription reaction was perfomed in a final volume of 10 μ L and contained 200 nM of *T. th.* HB8 σ^{A} -holoenzyme and between 20–40 nM of a PCR-amplified DNA fragment containing either a *T. th.* HB8 or a P23-45 promoter in standard transcription buffer (30 mM Tris-HCl [pH 7.9], 40 mM KCl, 10 mM MgCl₂, 2 mM β –mercaptoethanol). Reactions (where indicated) were supplemented with either gp39 or gp76 (15 μ M), incubated for 10 minutes at 65 °C, followed by the addition of various RNA dinucleotides (100–500 μ M), [α -³²P] NTPs (3000 Ci/mmol) and the corresponding cold NTPs (100 μ M). The reactions were incubated for a further 10 minutes at 65 °C prior to being terminated by the addition of an equal volume of urea-formamide loading buffer. The reaction products were resolved on a 20% (w/v) polyacrylamide denaturing gel and visualized using a PhosphorImager.

Sequence analysis

The sequences of the predicted proteins encoded in the P23-45 genome were searched against the non-redundant protein sequence database at the NCBI using the iterative PSI-BLAST ³⁰ and the pattern-hit-initiated BLAST (PHI-BLAST) ³² programs searches with P23-45 deduced ORFs. Additional searches were performed using the HHPred program that implements pairwise comparison of hidden Markov models ³³. The multiple alignment of the putative DPBB domains of ORF64 from P23-45, ORF62 from P74-26, and a representative set of multisubunit RNAPs was constructed using the MUSCLE program ⁴¹ Results of secondary structure prediction made using the PredictProtein ⁴² and JPred ⁴³ programs were taken into consideration to manually refine the alignment.

Acknowledgments

Bacteriophage P23-45 was generously provided by Dr. Michael Slater from Promega Corporation. We thank Dr. E. Peter Geiduschek for critical reading of the manuscript and helpful advice. This work was supported by NIH grant R21 AI074769 (to L.M.), by NIH grant R01 GM61898 (to S.A.D.), by NIH grants RR00862 and RR022220 (to B.T.C.), by the Stowers Institute for Medical Research (L.F. and M.P.W.) by NIH grant R01 GM59295 (to K.S.), by Molecular and Cell Biology grant from the Presidium of Russian Academy of Sciences, Russian Foundation for Basis Research grant 07-04-00366-a, and by a National Center for Biotechnology of the Republic of Kazakhstan grant (to K.S.); E.V.K. is supported by the US Department of Health and Human Resources (National Library of Medicine, National Institutes of Health).

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Figure 1. Genome and transcription map of the *T. th.* **HB8 bacteriophage P23-45 A**. Functional annotation of the predicted P23-45 ORFs. The different colors of the arrows represent the different functions assigned for the proteins encoded by P23-45. Upper arrows denote a rightward orientation of the genes, while lower arrows denote a leftward orientation of the genes.

B. The different colors of the transcription map indicate different temporal classes of genes: blue, early; brown, middle; green, late. Promoters are colored corresponding to their temporal class and are depicted as bent arrows. A section of the early cluster is shown in more detail. In both **A** and **B** black dots indicate genes used in gene macroarray analysis of the temporal transcription program.



Figure 2. Macroarray data analysis

A. Normalized abundances of total *T. th.* HB8-encoded transcripts and total P23-45-encoded transcripts are shown as red and black lines, respectively.

B. Normalized abundances for individual P23-45 transcripts are presented. Transcripts that belong to different temporal classes are colored: early, blue; middle, brown; late, green. The expression curves represent average results of three independent experiments.

C. Average normalized abundances of P23-45 transcripts of different temporal classes.



Figure 3. Characterization of middle and late P23-45 promoters

A. The kinetics of accumulation of *in vivo* primer extension products obtained with P23-45 transcripts from representative middle (P_{39}) and late (P_{80}) promoters during infection; the DNA products corresponding to 5' RNA ends are labeled with arrows.

B. An alignment of P23-45 middle and late promoter sequences is shown. The putative -35 (where identified), -10 and TG/TGTG promoter elements are highlighted in bold uppercase text. Experimentally identified transcription start sites are highlighted in bold and underlined. The annotated translation initiation codons are shown in bold lowercase text. The corresponding sequence logos for P23-45 middle and late promoters are depicted below the alignment. The size of the letters indicates the degree of conservation; positions are defined with respect to identified transcription start sites.



Figure 4. Alignment of DNA sequences upstream of the annotated translation initiation start codons of P23-45 early genes

The novel promoter elements are highlighted in bold uppercase text. Experimentally identified 5' RNA ends are highlighted in bold text and underlined. The annotated translation initiation start codons are highlighted in bold lowercase text. The grey boxes indicate stem-forming regions of putative *rho*-independent terminators. The sequence logos for the early P23-45 promoters and the independently aligned -35 and -10 regions of *T. th.* HB8 -10/-35 promoters are depicted below the alignment. The size of the letters indicates the degree of conservation; positions are defined with respect to putative or identified transcription start sites.



Figure 5. Characterization of putative early P23-45 promoters

A. The kinetics of accumulation of representative *in vivo* primer extension products obtained with P23-45 transcripts from putative early promoters P_{64} and P_{68E} during infection; the DNA products corresponding to 5' RNA ends are labeled with arrows.

B. The results of abortive transcription initiation by *T. th.* HB8 RNAP- σ^A holoenzyme at representative P23-45 early promoters (lanes 1–13) are shown. Dinucleotide RNA substrates corresponding to the -2/-1 and/or -1/+1 positions with respect to the putative transcription start sites (see Fig. 4) were used as the primers. Radioactively labeled NTP corresponding to the putative +1 or +2 positions was used to form the trinucleotide RNA product. As a positive control, abortive transcription at representative P23-45 middle and late promoters is shown (lanes 14 and 15).



Figure 6. Putative P23-45 early promoters are functional in vivo

A. Schematic diagram illustrating an experiment to probe the functionality of P23-45 promoters *in vivo*. *T. th.* HB8 harboring the plasmid pMKE77-80 was infected with P23-45 and total RNA was extracted throughout the infection at the time points indicated. **B.** The results of *in vivo* primer extension analysis of RNA transcribed from the divergent plasmid-located P23-45 early (P₇₇) and late (P₈₀) promoters (upper panels) and from the genome-located P₇₇ and P₈₀ promoters (lower panels). The DNA products corresponding to 5' RNA ends are labeled with arrows.



Figure 7. Transcription from P23-45 early promoters is Rif-resistant and does not depend on host RNAP

The results of *in vivo* primer extension analysis of RNA extracted from phage-infected cells in the absence of Rif (-) or in the presence of Rif (+) added at different time points postinfection. The panels A, B, and C demonstrate the kinetics of accumulation of transcripts from representative early, middle, and late P23-45 promoters, respectively.



Figure 8. MudPIT analysis of proteins that co-isolate with *T. th.* HB8 RNAP affinity isolated from P23-45 infected cells

Normalized Spectral Abundance Factor (NSAF) values for RNAP subunits and co-isolated *T. th.* HB8- and P23-45-encoded proteins are shown. *T. th.* HB8-encoded proteins are shown as white colored bars and P23-45-encoded proteins are shown as black colored bars.



Figure 9. The P23-45 phage proteins gp39 and gp76 bind to *T. th.* HB8 RNAP and inhibit transcription initiation from host promoters but not from middle and late phage promoters A. Gp39 and gp76 bind to both core RNAP and RNAP- σ^A holoenzyme. The proteins alone or together were incubated with RNAP core (lanes 2–4) or with RNAP- σ^A holoenzyme (lanes 6–8) and analyzed by native gradient PAGE. Next, the native gels were stained with Coomassie stain, and the bands due to proteins (indicated by brackets in Fig. 9A) were excised from the gels and their composition were determined by denaturing SDS-PAGE (lanes 1'-8'). Gp39 and gp76 were loaded as markers (lanes 9 and 10, respectively). B. The results of abortive transcription initiation by *T. th.* HB8 RNAP- σ^A holoenzyme at several representative host *T. th.* HB8 (lanes 1–9) and middle/late phage (lanes 10–15) promoters in the presence or absence of either gp39 or gp76 are shown.

Secondary structure	ORF64		eee	eeeee	
P23 ORF64 YP 0014679	917.1	243-341	GPNTGFGG-ILLSPKILPF	LGLHGLEDGGVLAYTRRWKP	
P74 ORF62 YP 0014680	032.1	270-368	GPNTGFGG-ILLSPKILPF	LGLHGLEDGGLLAYTRRWRP	
RPAI XP 002290924.1	Thaps	466-631	GKRVNFACRSVISPDPY	IGTNEIGLPLYF	75
RPA1 EEY66296.1	Phyin	466-637	GKRVNYAARSVISPDPY	ISTSQIGVPLRF	84
RPA1 XP 794863.2	Strpu	430-597	GKRVNYAARSVISPDPY	INTDEIGIPQVI	82
RPA1 EAW99467.1	Homsa	434-604	GKRVDYAARSVICPDMY	INTNEIGIPMVF	82
RPA1 EEH52215.1	Micpu	443-631	GKRVNYAARSVIMPDPY	LKTSEIGVPPVF	100
RPA1 ACN85301.1	Oryco	430-605	GKRVNYACRSVISPDPY	LAVNEIGIPPVF	87
RPA' 2WAQ A	Sulsh	315-472	GKRVDFSSRTVISPDPN	IISIDEVGVPEII	79
RPA' ZP 06213371.1	Metsp	318-482	GKRVNFSSRTVISPDPC	CLSINEVGVPEVV	82
RPb' AAB59112.1	Bacsu	333-445	GKRVDYSGRSVIVVGPH	ILKMYQCGLPKEM	57
RPb' 2GHO D	Theaq	329-444	GKRVDYSGRSVIVVGPQ	LKLHQCGLPKRM	57
			* ::: ::	:	
Secondary structure	2WAQ		eeeeeeee	eeeee	
Secondary structure	2GHO		eeeeeeee	eeeee	
DPBB consensus struc	cture:		eeeSleee	eeS2ee	
Secondary structure	ORF64	eeeee	eeeee h	hhhhhhh	eeeeee
P23_ORF64_YP_0014679	917.1	GERVIFNR	RPDLPTGQSAVELTYVGLSF	PIADSVIAHEHDIAPTGADYDG	D-IGYVFPTPEMG
P74_ORF62_YP_0014680	032.1	GERVIFNR	RPDLPTGQSAVELTYLGLSF	PIADSVIAHEGDIASTGADYDG	D-IGYLFPTPEKG
RPA1_XP_002290924.1	Thaps	GDMVLMNR	QPTLHKPGIMAHRVRVLFS F	TQNTLRMHYANCNTYNADYDG	DEMNCHFPQSYLA
RPA1 EEY66296.1	Phyin	GDVVLMNR	PTLHKPSIMAHTARVLTNE	KMQTIRMHYANCNTFNADFD	DEMNMHFPQNELA
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RPA1_XP_794863.2	Strpu	GDIVLLNRQ	PTLHKE	GIMAHK	VRVL-	-PGEKTL	RLHYSACK	TYNADFDGI	EMNV	HFPQNELG	
RPA1 EAW99467.1	Homsa	GDILLLNRQ	PTLHRE	SIQAHR	ARIL-	-PEEKVL	RLHYANCK	AYNADFDGI	EMNA	HFPQSELG	
RPA1_EEH52215.1	Micpu	GDVLLVNRQ	PTLHKE	GIMAHT.	AKVL-	-PGQRTI	RMHYANCS	TYNADFDGI	EMNL	HFPQDHLA	
RPA1_ACN85301.1	Oryco	GDIVLVNRQ	PTLHKE	SMMAHF	VRVL-	-PGEKTI	RMHYANCS	TYNADFDGI	EMNV	HFPQDEIS	
RPA ' 2WAQ	Sulsh	GDVVLFNRQ	PSLHRI	SMMAHR	VRVL-	-KG-LTF	RLNLLVCF	PYNADFDGI	EMNL	HVPQSEEA	
RPA' ZP_06213371.1	Metsp	GDIVLYNRQ	PSLHRN	ISIMAHR	VRVL-	-PY-RTF	RHNLCVCF	PYNADFDGI	EMNL	HVPQSEEA	
RPb' AAB59112.1	Bacsu	EHPVLLNRA	PTLHRI	GIQAFE	PTLV-	-EG-RAI	RLHPLVCI	AYNADFDGI	QMAV	HVPLSAEA	
RPb ' 2GHO	Theaq	GKVVLLNRA	PTLHRI	GIQAFQ	PVLV-	-EG-QSI	QLHPLVCE	AFNADFDGI	QMAV	HVPLSSFA	
_		:::**	* *	:	:		:	:**:**	* :	:*	
Secondary structure	2WAQ	eeeeeee		eeeeee	eeee	ee	eeee		eeee	ee	
Secondary structure	2GHO	eeeee		eeeeee	eee	eee	e		eeee	ee	
DPBB consensus structure:		eeS3e		eeeS4ee		eS5			eeS6ee		

Figure 10.

Alignment of the double-psi beta-barrel (DPBB) domains of ORF64 from P23-45 and ORF62 from closely related P74-26 and β' orthologs from selected multisubunit RNAPs. The alignment was generated using the MUSCLE program ⁴¹, and the N-terminal part was manually modified on the basis of secondary structure predictions. The range of amino acid residues of the aligned segments in the respective protein precedes each sequence. Inserts in RNAP subunit sequences are shown as the numbers of amino acids. The residues conserved in all aligned sequences are denoted by asterisks; similar residues are denoted by colons. The most conserved segments of the alignment including the NADFDGD motif are shown as grey boxes. The predicted secondary structure of ORF64 (consensus of the PredictProtein and JPred predictions) is shown above the alignment. The secondary structure from two crystal structures, those of the RNAPs of the archaeon Sulfolobus shibatae (PDB code 2WAQ⁴⁴) and the bacterium *T. aq.* (2GHO⁴⁵) are shown underneath the alignment along with the consensus structure of the DPBB domain ³¹. In the secondary structure lines, e stands for extended conformation (β -strand) and h stands for α -helix; the S1–6 numbering is from ³¹. The organism name abbreviations: Thaps, *Thalassiosira pseudonana*; Phyin, Phytophthora infestans; Strpu, Strongylocentrotus purpuratus; Homsa, Homo sapiens; Micpu, Micromonas pusilla; Oryco, Oryza coarctata; Sulsh, Sulfolobus shibatae; Metsp, Methanocaldococcus sp.; Bacsu, Bacillus subtilis; Theaq, Thermus aquaticus.