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Thermus thermophilus bacteriophage φYS40 genome and proteomic characterization of virions

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Abstract

We determined the sequence of the 152,372-bp genome of φ YS40, a lytic tailed bacteriophage of *Thermus thermophilus*. The genome contains 170 putative open reading frames and three tRNA genes. Functions for 25% of φ YS40 gene products were predicted on the basis of similarity to proteins of known function from diverse phages and bacteria. φ YS40 encodes a cluster of proteins involved in nucleotide salvage, such as flavin-dependent thymidylate synthase, thymidylate kinase, ribonucleotide reductase, and deoxycytidylate deaminase, and in DNA replication, such as DNA primase, helicase, type A DNA polymerase, and predicted terminal protein involved in initiation of DNA synthesis. The structural genes of φ YS40, most of which have no similarity to sequences in public databases, were identified by mass-spectrometric analysis of purified virions. Various φ YS40 proteins have different phylogenetic neighbors, including Myovirus, Podovirus, and Siphovirus gene products, bacterial genes, and in one case, a dUTPase from a eukaryotic virus. φ YS40 has apparently arisen through multiple acts of recombination between different phage genomes as well as through acquisition of bacterial genes.

Keywords

Thermus thermophilus; bacteriophage; genome; virion; proteomics; bioinformatics; DNA polymerase

Introduction

In the last decade, the genomes of several hundred phages have been completely sequenced (282 complete dsDNA phage genomes in the Genome Division of GenBank as of July 2006). While bacterial hosts of these phages are phylogenetically diverse, only ten of those completely sequenced phages are known to infect thermophilic microorganisms. Most of 'thermophilic'

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phages were isolated from a small number of archaeal species ¹⁻³. Sequence analysis revealed that archaeophages encode mostly uncharacterized proteins with no similarities to sequences in public databases, though more detailed examination revealed a limited number of recognizable ATPases, nucleotide salvage enzymes, and putative transcription factors ⁴. As of the time of this writing, the only sequenced genome of a phage from a thermophilic eubacterium is RM 378 that infects *Rhodothermus marinus*⁵.

During their development in a bacterial host, phages are known to regulate host macromolecular synthesis by modifying host transcription and translation machinery and making it serve the needs of the virus. Proteins from thermophilic bacteria are particularly amenable to structural studies of large complexes involved in DNA replication, DNA transcription, and RNA translation. Thus, structural and functional analysis of thermophilic phage-encoded regulators and their complexes with RNA polymerases, ribosomes, and other components of thermophilic bacteria can provide insights into molecular mechanisms of regulation of transcription, translation, and other cellular processes. With these ideas in mind, we determined the genomic sequence of φ YS40, a large myophage hosted by the thermophilic bacterium *Thermus thermophilus* (temperature range from 56 to 78°C)⁶. Here, we present the results of a preliminary study of the φ YS40 genome and the proteome of φ YS40 virions.

Results

Overview of the ϕ YS40 genome

The sequence of the φ YS40 genome was determined using the fimer technology and assembled into a single 152,372 bp contig using the phredPhrap package (see Materials and Methods). The G + C content of the φ YS40 genome is 32.59%, which is significantly lower than that of its host (69.4%). Though the GC-content of φ YS40 is close to values typical of the low-GC Gram-positive bacteria, there is no specific evolutionary affinity between sequences of φ YS40 and these bacteria, and the GC-content of the phage may instead reflect specific aspects of phage molecular biology, for example distinct mutational bias of its DNA polymerase. φ YS40 DNA appears to be unmodified as it is susceptible to digestion with all common methylationsensitive restriction endonucleases tested (data not shown).

A total of 170 ORFs were predicted in the φ YS40 genome (Table 1, Fig. 1). The intergenic regions were screened for additional genes by searching GenBank, GenPept, and the database of unfinished microbial genomes at NCBI, but no additional conserved ORFs were found. The predicted φ YS40 ORFs are between 43 and 1744 codons in length. As with most other phages, the genome of φ YS40 is tightly packed: coding sequences occupy 95% of the φ YS40 genome. There are 46 cases of overlaps (from 1 to 40 bases long) between neighboring ORFs. The longest non-coding region (390 bp) lies between ORF138 and ORF139. Most of the 170 predicted ORFs start at the AUG codon, 22 ORFs use GUG codon, and three use UUG. At the ends of φ YS40 genes, there are 90 TAA stop codons, 66 TGA codons, and 16 TAG codons.

Two-thirds of the φ YS40 genes (114 genes) are transcribed in one direction, designated as leftward in the genome map (Fig. 1), and 56 genes are transcribed in the rightward direction. The G+C content is approximately the same for both sets of ORFs. Taking a set of genes transcribed in the same direction and having no more than three consecutive intruders (*i.e.*, genes transcribed in a different direction) as a cluster, we find four gene clusters in the φ YS40 genome. The ORF1-ORF36 and ORF62-ORF146 clusters are transcribed in the rightwards direction, and ORF37-ORF61 and ORF147-ORF170 clusters are transcribed in the rightwards direction (Fig. 1). The probability of obtaining each of the four clusters by chance, calculated using equation 2 from Durand and Sankoff ⁷ is less than 0.1, indicating that at least part of the clustering may be due to evolutionary or functional constraints.

tRNA genes

Using the tRNA scan-SE program, we identified three tRNA genes in the φ YS40 genome. The tRNA1 gene overlaps with ORF61, whereas the tRNA2 and tRNA3 genes are both located between ORF139 and ORF140. Other large tailed dsDNA bacteriophages, such as coliphage T4⁸, vibriophage KVP40⁹, and phage phiKZ of *P. aeruginosa*¹⁰ also encode several tRNAs.

The φ YS40 tRNA1 and tRNA3 recognize ACA (threonine) and AGA (arginine) codons, respectively. These codons, while overrepresented in the φ YS40 genome, are the rarest threonine and arginine codons in *T. thermophilus* genes. tRNA2 has a CAU anticodon, which would correspond to methionine codon AUG if C34 in the wobble position is unmodified. In homologous tRNAs from a number of bacteria and bacteriophages, the corresponding cytidine is converted to lysidine, which results in the AUA (Ile) decoding ¹¹⁻¹³. Determinants for tRNA^{Ile} identity are thought to consist of anticodon loop bases A37 and A38, the discriminator base A73, and conserved base pairs in the D-arm (U12·A23), the anticodon arm (C29·G41), and the acceptor arm (C4·G69)¹⁴. All these characteristics are present in φ YS40 tRNA2, which therefore may decode the isoleucine codon AUA, another rare *T. thermophilus* codon that is much more frequent in φ YS40 ORFs. Thus, φ YS40-encoded tRNAs may ensure efficient decoding of codons that are overrepresented in the phage genome relative to its host.

Sequence analysis of predicted ϕ YS40 proteins

Analysis of intrinsic features of protein sequences indicates that seven φ YS40 ORFs encode proteins with putative transmembrane domains (from one to three) and four φ YS40 proteins are predicted to have coiled-coil regions. Only one protein, gp107, is predicted to be strongly non-globular, and only one protein, gp35, contains an N-terminal secretion signal peptide. All deduced amino acid sequences were compared to proteins in the non-redundant database at NCBI using the PSI-BLAST program with a slightly relaxed cutoff for profile inclusion (-h parameter). The comparison showed that ~25% of φ YS40 proteins display sequence similarity to proteins of known function (Table 1).

φYS40 proteins involved in nucleotide metabolism—Like other large phage genomes, **φ**YS40 encodes a number of enzymes involved in nucleotide metabolism. They are gp8, a homolog of mammalian/viral UTPase (EC 3.6.1.23); gp9, related to a predicted flavin-dependent thymidylate synthase (EC 2.1.1.148); GMP reductase gp17 (EC 1.7.1.7); thymidine kinase gp24 (EC 2.7.1.21); deoxycytidylate deaminase gp38 (EC 3.5.4.12); dNMP kinase gp60 (EC 2.7.4.-); and the catalytic α subunit of ribonucleotide reductase encoded by two adjoining ORFs, gp41 and gp42 (EC 1.17.4.1). Except for dUTPase gp8, all these gene products show stronger sequence similarity to prokaryotic or phage enzymes than to their eukaryotic or archaeal counterparts. The best database match and closest phylogenetic neighbor for dUTPase gp8 is dUTPase from *Lymantria dispar* nucleopolyhedrosis virus. Gene exchange between phages and bacteria has been suggested to account for odd gene phylogenies that are sometimes observed in the components of bacterial replication and transcription machinery ¹⁵. Our observation indicates that eukaryotic viruses, and perhaps their hosts, may also be involved in such exchange.

 φ YS40 proteins involved in DNA replication and recombination— φ YS40 encodes most of the proteins required for replisome formation, namely gp14, a replication initiation helicase DnaB; gp23, a bacterial DnaG-family DNA primase; gp26, a RecB family exonuclease; gp33, a type A DNA polymerase, and gp27, a DEAD box helicase. Another predicted DEAD-box helicase is encoded by gp79. Based on the fact that gp79 is a part of the φ YS40 virion, we suspect that it is involved in viral DNA packaging. φ YS40 also encodes two recombination proteins, gp12, a RecA/RadA recombinase, and gp114, an ssDNA-

annealing protein of the ERF family. There are no gene products with detectable sequence similarity to known ssDNA-binding proteins 16 or DNA ligases.

The product of gene 65 is of particular interest for understanding the replication mechanism of φ YS40. It shows a striking sequence similarity to a portion of the terminal protein (TP) of *B. subtilis* phage φ 29. The Ser²³² residue of the TP protein forms a phosphoester bond with the 5'-terminal dAMP of the phage genome, and is essential for protein-primed replication of linear dsDNA genome of φ 29¹⁷⁻¹⁹. This serine is conserved in φ YS40 gp65 (Fig. 2). Thus, it is likely that gp65 primes the replication of φ YS40 genomic DNA. It should be noted that the ends of the φ YS40 genome as presented in Fig. 1 are arbitrary, since no defined ends were revealed during genome sequencing and assembly, indicating that the φ YS40 genome may be circularly permuted or may have direct terminal repeats. This matter requires further investigation.

Properties of the \varphiYS40 DNA polymerase—The φ YS40 gp33 is a type A DNA polymerase, which contains a conserved nucleotidyltransferase domain and a 3'-5' exonuclease domain, but lacks the 5' \rightarrow 3' exonuclease domain. Since gp33 is the first known example of a type A DNA polymerase from a thermophilic phage, we expressed recombinant gp33 in *E. coli* and studied its properties *in vitro*. At 60-65 °C, recombinant gp33 exhibited moderate polymerization activity and very strong 3' \rightarrow 5'exonuclease activity toward both single-stranded DNA and double-stranded DNA substrates, even in the presence of 1 mM dNTP. As a result, at pH > 8.0 and low salt concentrations, the enzyme mostly hydrolyzed the primer. The increase of salt concentration partially inhibited the polymerase activity as well. The decay of primer-template substrate by gp33 exonuclease was abolished when primers were protected with thiolate modification, but the interference of the exonucleolytic activity during elongation resulted in poor DNA yield.

Gp33 was moderately thermostable. Both polymerase and exonuclease functions were lost after a 3-min incubation at 85 °C. At 75 °C, the polymerase activity decreased faster than the exonuclease activity; as a result, the enzyme produced shorter elongation products after heating. Similarly low thermostability has been reported for type B DNA polymerase from the *Rhodothermus marinus* phage (a half-life of 2 min at 90 °C⁵). These observations indicate that both processivity of φ YS40 DNA polymerase and its stability at elevated temperatures must be conferred by its interactions with other components of the replicative complex, in marked contrast with other DNA polymerases of bacteria and archaea, such as *Taq* or *Pfu*, which are processive and thermostable in the absence of cofactors.

Protein composition of \varphiYS40 virions—To identify φ YS40 structural proteins, φ YS40 virions were purified by double sedimentation in CsCl gradients. The results of SDS-PAGE analysis of purified φ YS40 virions are shown in Fig. 3. The two major protein components of the virion were identified by mass spectrometry as gp73 and gp19 (Fig. 3). These proteins may correspond to major head and tail proteins, but their function could not have been predicted by sequence comparison because of lack of database homologs.

Three independent φ YS40 lysates of increasing titer (from 2×10⁷ to 2×10⁹ pfu/ml) were also directly examined by multidimensional protein identification technology, MudPIT ²⁰ a shotgun proteomics approach where proteolytic peptides of a protein complex under study (in our case, phage virions) are generated, loaded onto triphasic microcapillary columns, eluted over several chromatography steps and analyzed directly by tandem mass spectrometry. Peptides matching 33 φ YS40 proteins were detected in one or more of these samples. There were also 79 host proteins, all of which decreased in abundance when the lysates of higher titer were used as a starting material for CsCl purification (Supplementary Table A). In contrast,

the NSAF (Normalized Spectral Abundance Factor, see Materials and Methods) values for φ YS40 proteins increased with the titer of phage in the starting sample (Supplementary Table B). Gp73 and gp19 were detected at the highest levels in all three analyses in agreement with these being major structural proteins. With the exception of gp52 (annotated as a UDP-3-O-[3-hydroxymyristory] glucosamine N-acyltransferase), gp69 (tail sheath protein), gp79 (DEAD-Box helicase), gp150 (putative baseplate assembly protein), and gp152 (fibritin neck whisker), most φ YS40 virion proteins identified in this analysis are novel proteins without any detectable database homologs. Interestingly, all multiply detected φ YS40 virion proteins are the products of adjacent co-transcribed genes, except for ORF19 (Fig. 4C). In particular, a group of 13 proteins detected at high levels are the products of genes at the end of the largest cluster of φ YS40 genes (ORF62-ORF146, above) that therefore may correspond to the late gene cluster.

DISCUSSION

Bacteriophages may be the most abundant living entities on Earth. It has been proposed that the origin of dsDNA bacteriophages is as ancient as DNA replication itself and that the analysis of the currently known bacteriophages may provide clues to early evolution of cellular and viral genomes ¹⁵.

Here, we report a preliminary analysis of *Thermus thermophilus* bacteriophage φ YS40 genome. The analysis shows that φ YS40 does not easily fit into previously established groups of dsDNA bacterial viruses and may represent a distinct branch of the *Myoviridae* family. A substantial fraction of φ YS40 genes codes for predicted proteins to which no function can be assigned; however, 25% of the φ YS40-encoded proteins show detectable homology to their counterparts in a broad phylogenetic range of microorganisms, and some proteins are homologous to proteins found in other dsDNA bacteriophages infecting diverse hosts, such as *Staphylococcus*, *Rhodothermus marinus*, and *Vibrio parahaemolyticus*. In agreement with morphological data, predicted tail genes are mostly *Myoviridae*-related. Most of other φ YS40 genes that have database homologs are, however, closer to either podoviral or siphoviral gene products: for instance, gp26 (RecB family exonuclease) and gp60 (dNMP kinase) are most closely related to homologs from a podovirus SIO1 and a λ -like siphovirus phi-BT1, respectively. Yet other genes are phylogenetically close to bacterial genes, and, in one case, to a homolog from a eukaryotic baculovirus. φ YS40 has apparently arisen through multiple acts of recombination between different groups of phages and perhaps even their hosts.

Molecular adaptations to thermophily in various species are of great interest. Comparative studies of the genomes of thermophilic, hyperthermophilic, and mesophilic prokaryotes have suggested several attributes of thermostability at the levels of amino acid sequence, properties of folded proteins, and gene content. The proposed sequence level predictors of thermostability, such as large charged-versus-polar (CvP) amino acid ratio or (E + K)/(Q + H) ratio, are not conclusive in the case of φ YS40, and genes that are indicative of the host ability to survive at extreme temperatures ²¹ are missing from the φ YS40 genome. Moreover, only seven φ YS40 gene products have closest phylogenetic neighbors in thermophilic microorganisms.

In its genome size, φ YS40 is similar to T4, an *E. coli* phage that is known to rely on host RNA polymerase for expression of its genes. During its development, T4 sequentially modifies host RNA polymerase to shut off transcription of host genes and to ensure correct expression of several classes of its own genes (reviewed in Ref. 22). Like T4, φ YS40 does not encode its own RNA polymerase and therefore has to rely on the host enzyme for transcription of its DNA. The early genes of φ YS40 should therefore be transcribed by the *T. thermophilus* RNA polymerase holoenzyme, most likely containing general initiation factor σ^A . Preliminary analysis reveals the presence of sequences with strong similarities to bacterial housekeeping

sigma promoters in front of many ϕ YS40 genes, but no such sequences are found in front of genes coding for ϕ YS40 structural proteins (A. Sevostyanova, M. Gelfand and KS, unpublished observations). Structural genes, which should be expressed late in infection, must be therefore transcribed by a modified form of host RNA polymerase. Further biochemical studies may reveal ϕ YS40 proteins that are required for these modifications.

MATERIALS AND METHODS

Cell growth and phage infection

The bacterial strain *Thermus thermophilus* HB8 and φ YS40 were generously provided by Dr. Tairo Oshima, Tokyo University of Pharmacy & Life Science. The cells and phage were grown overnight in the Tth medium (0.8% polypeptone, 0.4% yeast extract, 0.2% NaCl, and 0.35 M CaCl₂ and 0.4 M MgSO₄) at 65°C with vigorous agitation.

To isolate individual φ YS40 plaques, 1 ml of overnight HB8 culture (**OD600~1.6**) was centrifuged and resuspended in 100 µl of the Tth medium and combined with 5 µl dilutions of φ YS40 stock, incubated for 15 min at 65°C, plated in soft Tth agar (0.7 %), and incubated overnight at 65°C. An individual plaque was picked up and subjected to two more rounds of plaque purification, before making a phage lysate stock solution. To this end, a single plaque was resuspended in a small volume of the Tth medium and mixed with 0.1 ml of overnight HB8 culture. The mixture was incubated for 15 minutes at 65 °C to allow phage absorption, 5 ml of fresh Tth medium was added and the culture was incubated on a rotary shaker at 65 °C until complete lysis occurred (usually overnight). Cell debris was removed from the lysate by centrifugation at 12,000g for 15 minutes. The resultant phage stock (6×10⁹ pfu/ml) was saturated with chloroform and stored at 4 °C. The φ YS40 stock was used to prepare larger amounts of phage lysate using a scale-up of the procedure described above.

Purification of **\phyS40** virions

DNase I and RNase A (each to a final concentration of 1 μ g/ml) were added to ϕ YS40 lysed T. thermophilus culture followed by a 30-min incubation at 30°C. Solid NaCl was added a final concentration of 1 M and dissolved by swirling. The lysed culture was left on ice for 1 h and centrifuged at 11,000 g for 10 min at 4°C. To precipitate φ YS40, PEG 8000 was added to the supernatant to the final concentration of 10% (w/v) followed by a 1-h incubation on ice. Precipitated φ YS40 particles were recovered by centrifugation at 11,000g for 10 min at 4 °C. The phage pellet was resuspended in 2 ml of SM buffer (NaCl, MgSO₄, Tris-HCl, pH7.5, 2% gelatin). The PEG 8000 and cell debris were extracted from the phage suspension by adding an equal volume of chloroform and centrifuged at 3,000g for 15 min at 4°C. 0.5 g of solid CsCl per milliliter of bacteriophage suspension was added to the aqueous phase, which contained the bacteriophage particles, and dissolved by gentle mixing. CsCl step gradients (three steps with 1.45, 1.50, and 1.70 g/l density) were performed in Beckman SW41 polypropylene centrifuge tubes at 22,000 rpm for 2 hrs at 4 °C and at 38.000 rpm for 24 hrs at 4 °C (Beckman SW50.1 rotor, Beckman Coulter, Fullerton, CA). Purified bacteriophage suspension was dialyzed twice at room temperature for 1 h against a 1000-fold volume of 10 mM NaCl, 50 mM Tris-HCl pH 8.0, 10 mM MgCl₂.

Extraction of phage DNA

EDTA (to a final concentration of 20 mM), proteinase K (to a final concentration of 50 μ g/ml), SDS (to a final concentration of 0.5%) were added to bacteriophage solution and incubated at 56°C for 1 h. An equal volume of phenol was added to chilled bacteriophage suspension, mixed, and centrifuged at 3000 g for 5 min at room temperature. The aqueous phase was extracted with a 50:50 mixture of equilibrated phenol and chloroform, and equal volume of chloroform. DNA was precipitated with ethanol.

Genome sequencing

Initial sequence data were obtained using mini shotgun library of phage DNA. Several rounds of sequencing reactions were performed directly on phage DNA using ThermoFidelase and Fimer technology²³, ²⁴. Trace assembly was done with phredPhrap package (http://www.phrap.com/)²⁵. The final round of sequencing resulted in one pseudocircular contig with a no- errors quality level.

Sequence analysis

ORFs of φ YS40 were predicted using the GeneMark server (http://opal.biology.gatech.edu/ GeneMark/heuristic_hmm2.cgi, Ref. 26). The PSI-BLAST program ²⁷ was used to detect the homologs of φ YS40 genes in the DNA and protein databases, with profile inclusion cutoff *E*-value in PSI-BLAST (-h parameter) set at 0.02. Both options for low-complexity filtering (-F parameter) and composition-based statistics (-t parameter) were sometime adjusted for better detection in sequence similarities. Phylogenetic analysis was performed using the programs in the PHYLIP package.²⁸

tRNA genes were searched by using the tRNAscan-SE program. ²⁹ Searches for the presence of the transmembrane helices and coiled coil regions were done with the aid of the SEALS package. ³⁰

MudPIT

Three independent virion lysates were prepared by double sedimentation in CsCl gradients and had phage titers of 2×10^7 pfu/ml, 4.2×10^8 pfu/ml and 2×10^9 pfu/ml. These lysates were treated with for 30 minutes at 37°C with 0.1U of benzonase (Sigma, St. Louis, MO), then precipitated in 20% trichloroacetic acid, 100mM Tris-HCl, ph 8.5, overnight at 4°C. The dried protein pellets were denatured, reduced, alkylated and digested with endoproteinase LysC and trypsin (both from Roche Applied Science, Indianapolis, IN) as described previously. ³¹ Peptide mixtures were pressure-loaded onto split-triphasic microcapillary columns, installed in-line with a Quaternary Agilent 1100 series HPLC pump coupled to Deca-XP ion trap tandem mass spectrometer (ThermoElectron, San Jose, CA) and analyzed via seven-step chromatography as described in Ref. 31.

The MS/MS datasets were searched using SEQUEST ³² against a database of 171 YS40 predicted gene products, combined with 2224 protein sequences from *Thermus thermophilus*, strain HB8 (chromosome and large plasmid) downloaded from NCBI on 2005-08-01, as well as usual contaminants such as human keratins, IgGs, and proteases. In addition, to estimate background correlations, each sequence in the database was randomized (keeping the same amino acid composition and length) and the resulting "shuffled" sequences were concatenated to the "normal" sequences and searched at the same time (the total number of sequences searched was 5144).

DTASelect/CONTRAST program³³ was used to select spectra/peptide matches with normalized difference in cross-correlation score (DeltCn) of at least 0.11, a minimum cross-correlation score (XCorr) of 1.8 for singly-, 2.5 for doubly-, and 3.5 for triply-charged spectra, a maximum Sp rank of 10, and a minimal length of 7 amino acids. In addition, the peptides had to be fully tryptic. No peptides matching shuffled protein sequences passed this criteria set. Spectral counts are considered to be a good estimation of absolute protein abundance³⁴. To account for the fact that larger proteins tend to contribute more peptide/spectra, spectral counts are divided by protein length defining a Spectral Abundance Factor (SAF).³⁵ SAF values are normalized against the sum of all SAFs for each run (removing redundant proteins) allowing us to compare protein levels across different runs using the Normalized Spectral Abundance Factor (NSAF) value.

φYS40 DNA polymerase

The gene encoding φ YS40 DNA polymerase was PCR amplified using appropriate primers annealing at the beginning and the end of φ YS40 gene 33 and containing engineered *NdeI* site CATATG overlapping with the initiating ATG codon of gene 33 and a *Hind*III site downstream of the termination codon (primer sequences are available from the authors upon request). The amplified fragment with treated with *NdeI* and *Hind*III and cloned into appropriately digested pet21d plasmid and transformed into the *E. coli* expression strain BL-21 pLysS. Cells were grown in 1 L of LB medium and induced with 1 mM IPTG. Cell pellet was dissolved in 15 ml of lysis buffer and centrifuged at 17000 rpm for 30 min (no heat treatment). Lysate was diluted to 0.25M NaCl, and applied on a Heparin Sepharose High-Trap column (GE Healthcare, Newark, NJ), equilibrated with 50 mM Tris pH 7.5, containing 0.25 M NaCl and 2 mM mercaptoethanol. After washing with the same buffer, φ YS40 DNA polymerase was eluted in about 0.3-0.35 M NaCl and appeared to be over 80% pure by SDS-PAGE. Assays of its enzymatic activities were done essentially as described by Pavlov et al., Ref. 36.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.

The ϕ YS40 genome.

Bacteriophage φ YS40 genome is schematically presented with predicted ORFs indicated by arrows. Arrow direction indicates the direction of transcription. Several ORFs with clear functional predictions for their products are color-coded (see also Table 1 for more details).

Nf phi29 GA-1 B103	(74) VEPYKKKY V VOLKA NA ANA KAO NI DOCHAL SURVEY ISBORODYOOMAL LISBORVETSREED TO VERVAL HE LIGHMARKA ED (74) VEPKKKY VOLKA NA SURVEY VOLKA NA KAO NI DOCHAL SURVEY ISBORODYOOMAL LISBORVETSREED TO VERVAL HE LIGHMARKA ED (74) VEPKKYKY VOLKA NA SURVEY VOLKA NI NA KAI LISBORVEY ISBORODYOVERKELLAVIONARE EDITE VID VERVAL HE ALS MARKA ED (74) VEPKKYKY VOLKA NA SURVEY VOLKA NI NA KAI LISBORVEY ISBORODYOVERKELLAVIONARE EDITE VID VERVAL HE ALS MARKA ED (74) VEPKKYKY VOLKA NA SURVEY VOLKA NI NA KAI LISBORODYOVERKELLAVIONARE EDITE VID VERVAL HE ALS MARKA ED (74) VEPKKYKY VOLKA NA SURVEY VOLKA NI NA KAI LISBORODYOVERKELLAVIONARE EDITE VID VERVAL HE ALS MARKA ED (74) VEPKKYKY VOLKA NA SURVEY VOLKA NI NA KAI LISBORODYOVERKELLAVIONARE EDITE VID VERVAL HE ALS MARKA ED (74) VEPKKYKY VOLKA NA SURVEY VOLKA NI NA SURVEY I SORODYOVERKELLAVIONARE EDITE VID VERVAL HE ALS MARKA ED (74) VEPKKYKY VOLKA NA SURVEY VOLKA NI NA SURVEY I SORODYOVERKELLAVIONARE EDITE VID VERVAL HE ALS MARKA ED (74) VEPKKYKYKY VOLKA NA SURVEY VOLKA NI NA SURVEY I SORODYOVERKELLAVIONARE EDITE VID VERVAL HE ALS MARKA ED (74) VEPKKYKYKYKYKYKYKYKYKYKYKYKYKYKYKYKYKYKYK
PZA YS40	(74) YOPEKNAYOWAGKAKLARIERNTKENORLVDEKINAKDKEYYAGGKPOGTIEORIAMTSPARVTGINRPRDTDEKAYYSEKTLEES-MEMRTDPO MYOLIKOLPLIPPTIOTISLANLIKAIGESNOKN 20000000000000000000000000000000
Nf phi29 GA-1 B103 P2A YS40	

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Figure 2.

Sequence alignment of the TP proteins.

Multiple alignment of terminal proteins (TP) from φ 29 family phages and phage φ YS40 gp65. The stretch of * indicates a region of a predicted amphipathic alpha-helix in TP. Distances, in amino acid residues, from the ends of each sequence and between blocks, are shown in parentheses. A white font in blue indicates the residue identical in all sequences compared, yellow shading indicates the conservation of hydrophobic residues, grey shading indicates the conservation of polar and charged residues. The white font in red indicates the Ser₂₃₂ that is essential for TP priming activity.



Figure 3.

SDS-PAGE analysis of the φ YS40 virion proteins.

The SDS gel shows the protein composition of purified ϕ YS40 virions. The two major bands identified by mass-spectrometry are indicated.



Figure 4. MudPIT analysis of ϕ YS40 lysates.

- A. Normalized Spectral Abundance Factor (NSAF) values measured for ϕ YS40 proteins detected in at least two of the three runs.
- **B.** NSAFs for contaminating *T. thermophilus* proteins detected in at least two of the three runs.
- C. All 33 φ YS40 genes for which products were detected are plotted along the genome as a function of the measured NSAF values (when proteins were identified in several runs, maximal NSAF values are reported). The arrows under the x axis represent the position of the leftward and rightward predicted transcription clusters.

	Function and other properties b	distal tail fiber protein	unknown portal protein TM, unknown	S-adenosylmethionine decarboxylase (adoMetDC) unknown	unknown dUTPase	flavin-dependent thymidylate synthase unknown	gp18, unknown function	RecA/RadA recombinase P.od 20 strond - avolvance - modein	DNA helicase DnaB	unknown unknown IMP dehydrogenase / GMP reductase	DNA binding HTH-domain protein, transcription regulator	Major structural protein unknown	unknown unknown	DNA primase bacterial DnaG type thymidine kinase	ATP-dependent ClpP protease RecR family exonuclease	DEAD domain helicase	unknown sugar-phosphate nucleotidyltransferas e	unknown unknown	unknown DNA polymerase, without N-terminal 5-3 exonuclease domain	3 TMs, unknown 2 TMs, unknown	unknown unknown	deoxycytidylate deaminase unknown	unknown ribonucleotide reductase, alpha subunit, the N-terminus ribonucleotide reductase alpha subunit the C-terminus		unknown unknown unknown unknown	4 TMs, unknown
lable 1	Taxonomic origin of the best match	Vibrio phage KVP40	Staphylococcus phage K Flavobacterium johnsoniae 11W101	Methanocaldococcus jannaschii	Lymantria dispar micleonolyhedrovinis	Thermotoga maritima	Burkholderia cepacia phage Bcen22	Microbulbifer degradans Thermus thermoshilus HR27	Ralstonia metallidurans	Microbulbifer degradans	Novosphingobium aromaticivorans			Heliobacillus mobilis Photorhabdus luminescens	Borrelia burgdorferi Rosenhacter nhaoe SIO1	Streptococcus pneumoniae	Bacteroides fragilis YCH46		Bacteroides thetaiotaomicron			Xanthomonas campestris	Azotobacter vinelandii Thermonnaroharter	tengcongensis		
	The best The best database match with validated	sunuaruy 34419532	48696430 90591438	19924248	9631083	33357605	33860394	23029929 16200225	22978288	23029305	23110678			27262500 37526389	15595102 9964675	15900485	52216967		29348669			21229604	23104360 20808702			
ومارمته لمرمية بلامسم سنمنا	neur predicted inloted ORF length (amino acids)	643 001	945 218	144 74 202	203 180	275 171	157	339 101	447	97 369	136	411 145	126	558 202	237 308	449	466 169	585 237	157 703	96 88	92 121	142 142	174 797 199		68 150 71	147
2	1 DIIAge 7 1040 Allu I ORF strand/position ^a	-/(71938)	- / (1741+200) - / (45737410) - / (74128068)	- / (80968530) - / (85648788)	- / (88019412) - / (93999941)	- / (995510782) - / (1081611331)	-/(1131011783)	-/(1177612795)	- / (1341314756)	-/(14/4515036) 1512415453 1546716576	1664017050	-/(1710818343) -/(1840018837)	-/(1883419214) -/(1918719960)	- / (1994421620) - / (2166922277)	- / (2230223015) - / (22975_2301)	- / (2389825247)	2539626796 2682227331	- / (2732829085) - / (2909029803)	- / (2981830291) 3038732498	- / (3249132781) - / (3276833034)	- / (3303133309) 3338133746	3373034158 3418834616	3463135155 3520137594 37607 38206		3824038446 3845938911 3889839227 3922439439	3944139884
	Gene products of ORF name	c	1 m 4	v v	~ 8	9 10	11	12	41	21 81 71	18	19 20	21	23 24	25 26	27	28	30 31	32 33 3	35 35	36 37	38 39	40 41 42	j (44 45 46	47

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NIH-PA Author Manuscript	Function and other properties b	unknown unknown unknown	UDP-3-0-[3-hydroxy-myristory] glucosamine N-acyltransferase unknown unknown	unknown conserved bacterial protein, unknown	spore concex syntresis protein apovic unknown putative serine motein kinase	putative dNMP kinase	terminase large subunit unknown	unknown terminal protein	unknown unknown	unknown tail sheath protein	unknown Zn ribbon, similar to archaeal transcription factor IIB	unknown Major structural protein	unknown, 3 coiled coil regions	unknown unknown unknown. 3 coiled coil regions	unknown	helicase (DEAD motif replaced by DDAE) unknown	unknown iniknown	unknown	unknown	unknown	unknown	unknown 2 TMs, unknown	unknown	unknown	unknown	unknown unknown	unknown	3 TMs, unknown unknown	unknown	unknown	unknown
NIH-PA Auth	Taxonomic origin of the best match		Mesorhizobium sp. BNC1	Geobacter metallireducens Symbiobacterium thermophilum	baenoviorio pacieriovorus Desulfitohacterium hafniense	Streptomyces phage phi-BT1	Methanocaldococcus jannaschii	Bacillus phage B103		Chloroflexus aurantiacus	Staphylococcus phage K		Staphylococcus phage K			Aspergillus nidulans						Deinococcus radiodurans									
or Manusc	The best database match with validated similarity		45914890	23055325 51891857 12521855	42521050 23112542	29366771	15668504	22855150		22973075	48696435		48696431			40/44644						15805515									
ript	ORF length (amino acids)	102 115 151 490	290 175 161	233 340 466	400 616 619	195 149	622 262	126 183	88 246	216 648	220	171 470	544 86	1525 1744	121	1524 287	650 612	75	76 50	83	78 81	61 161	151	202 144	141	140 58	43	135 181	180	196 170	178
NIH-PA Author	ORF strand/position ^a	3987740185 4020140548 4055841013 41010_47482	4253643408 4341143938 4394044425	- / (4442645127) 4518746209 46100 47526	4019947330 4756449414 4945351312	5141051997 5203552484	- / (5247754345) - / (5432055108)	- / (5510555485) - / (5546656017)	5604956315 - / (5636257102)	- / (5710457754) - / (5777559721)	-/(59/8260492) -/(6049561157)	- / (6116761682) - / (6175663168)	-/(6320464838) -/(64838_65008)	- / (0508569662) - / (6508474918) - / (6968474918)	- / (7493175296)	- / (7530979883) - / (7988080743	- / (8078882740) - / (82771 84609)	- / (8486785094)	-/(8532885558) -/(85767_85019)	- / (8602286273)	-/(8638286618)	- / (8750587990)	- / (8807488529)	- / (8934989783)	- / (8979690221)	- / (9103691212)	9123191359	-/(9141791824) -/(91835_92380)	- / (9250393045)	-/(9304593635) -//93610_04131)	- / (9433794873)
Manuscript	ORF name	48 50 51	52 53 54 54	55 56 57	58 58 59	60 61	63 63	64 65	66 67	69 69	0/	72 73	74 75	91 77	78	6/ 08	81 82	83	84 85	86	87 88	89	90 10	92	93 04	95 95	96 52	97 89	66	100	102

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Function and other properties ^b	unknown unknown unknown unknown	ATPase unknown unknown unknown	unknown unknown glycosyltransferase unknown unknown	unknown unknown unknown unknown unknown unknown	unknown unknown unknown unknown	unknown unknown unknown unknown coiled coil, unknown unknown	unknown unknown unknown unknown unknown	unknown unknown coiled coil, unknown unknown M27/M37 peptidase	unknown 3 TMs, unknown unknown unknown	putauve baseptate assembly protent unknown unknown unknown unknown unknown unknown unknown
Taxonomic origin of the best match		Thermotoga maritima	Escherichia coli		Shigella flexneri		Corvnebacterium olutamicum	Rhodococcus equi		r usobacter um nucleatum Escherichia coli phage K3
The best database match with validated similarity	•	15643692	11992695		18462664		19557983	10956653		903862
ORF length (amino acids)	162 171 176 173	202 347 231 248	130 181 179 163 80 158	185 183 188 164 167 172	105 171 194 165	150 168 168 178 215	189 190 219 287 534	183 58 1014 174 346	316 86 131 162	631 796 703 116 287 266
ORF strand/position ^a	- (9488595373) - (9551096025) - (96809696626) - (9683397354) . (07575 00053)	- / (99280100323) - / (100462101157) - / (101227101973)	- / (102138102530) - / (102531103076) - / (103077103616) - / (103616104107) - / (104503105279)	- ((105422105979) - / (105969105520) - / (106510107076) - / (107090107539) - / (107552108046) - / (108772108729.10 - / (108772109290)	-/(109528.110513) -/(109998.1110513) -/(111157.1111654)	-/(111605112153) -/(112165112677) -/(1126891131955) -/(113202113630) 113852114388 -/(114385115032)	- ((115155115724) - ((115727116299) - ((116271116693) 116815117474 - ((117442118005) - /(118365119999)	- / (120226120777) 120821120994 - / (120953123997) - / (124553125593) - / (124553125593)	- / (125998126948) - / (126553126813) 126870127055 127065127460 12771127959	12/9/9/12/12/9/9/ 12/9/64131859 131870134260 134253136364 136588137644 137634138497 137634139269
lame									149	
ORF n	103 104 105 106	108 108 110	111 112 113 114 115	117 118 120 121 122 123	124 125 126	128 130 131 132 133 133 133	134 135 136 137 138	140 141 143 143	(45) 146 147 148	151 152 153 154 155 156 157

F name	OqtnV Vd-HIN ORF strand/position ^a 139253143296 143322143846 144155143846 144155143846 144155145396 144756145390147022) 147094147639 1477094147639 147767188689 147767188689 147767188689 147756151907 1510264.15157 151024.15157 151057 151057 1510557 1510557 1510557 1510557 15105	pdij ORF length (amino acids) acids) acids acids) acids acids) acids acid acids acids acid	SENUEV JOU The best database match with validated similarity 15674141	HIN VA-HIN Taxonomic origin of the best match Lactococcus lactis	Function and other properties ^b Emotion and other properties ^b unknown unkno
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a position of the ORFs in the phage YS40 genome; "-" indicates a leftwards transcription orientation.

 \boldsymbol{b} presence of transmembrane domains (TM) and coiled coil regions are indicated.