Different Expression Patterns of Genes from the Exo-Xis Region of Bacteriophage λ and Shiga Toxin-Converting Bacteriophage Φ24b following Infection or Prophage Induction in Escherichia coli

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Abstract

Lambdoid bacteriophages serve as useful models in microbiological and molecular studies on basic biological process. Moreover, this family of viruses plays an important role in pathogenesis of enterohemorrhagic Escherichia coli (EHEC) strains, as they are carriers of genes coding for Shiga toxins. Efficient expression of these genes requires lambdoid prophage induction and multiplication of the phage genome. Therefore, understanding the mechanisms regulating these processes appears essential for both basic knowledge and potential anti-EHEC applications. The exo-xis region, present in genomes of lambdoid bacteriophages, contains highly conserved genes of largely unknown functions. Recent report indicated that the Ea8.5 protein, encoded in this region, contains a newly discovered fused homeodomain/zinc-finger fold, suggesting its plausible regulatory role. Moreover, subsequent studies demonstrated that overexpression of the exo-xis region from a multicopy plasmid resulted in impaired lysogenization of E. coli and more effective induction of λ and Φ24b prophages. In this report, we demonstrate that after prophage induction, the increase in phage DNA content in the host cells is more efficient in E. coli bearing additional copies of the exo-xis region, while survival rate of such bacteria is lower, which corroborated previous observations. Importantly, by using quantitative real-time reverse transcription PCR, we have determined patterns of expressions of particular genes from this region. Unexpectedly, in both phages λ and Φ24b, these patterns were significantly different not only between conditions of the host cells infection by bacteriophages and prophage induction, but also between induction of prophages with various agents (mitomycin C and hydrogen peroxide). This may shed a new light on our understanding of regulation of lambdoid phage development, depending on the mode of lytic cycle initiation.

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Introduction

Bacteriophage λ, which is a virus infecting Escherichia coli cells, has been used as a model in studies in the fields of microbiology and molecular biology for over 6 decades [for a review, see [1]]. Shortly after its discovery, it appeared that there are other bacteriophages, which genomes’ organization and developmental pathways are similar to those of λ, therefore, the family of lambdoid bacteriophages has been formally established. One of characteristic features of these viruses is their ability to follow two alternative developmental pathways. The lytic pathway includes phage genome replication and synthesis of phage-encoded regulatory and structural proteins, leading to production and liberation of progeny virions. The lysogenic pathway consists of integration of the phage genome into host chromosome, forming a prophage, and a passive replication of this form of the viral genome together with bacterial DNA (host cells bearing integrated phage genomes are called lysogens). However, the lysogenic stage is not permanent. A developmental switch, which consists of prophage induction, excision of phage DNA from the host chromosome, and initiation of the lytic mode of phage development, can occur under certain conditions causing a DNA damage in the host cell [for reviews, see [2,3]].

Genomes of some lambdoid bacteriophages, apart from genes characteristic for the whole family, contain also genes (abbreviated stx) encoding Shiga toxins. If E. coli is lysogenic with such a phage, it may be highly pathogenic to humans. Bacterial strains bearing such prophages are called Shiga toxin-producing E. coli (STEC), and this group includes enterohemorrhagic E. coli (EHEC) strains that are particularly dangerous pathogens [4–6]. Bacteriophages...
bearing stx genes in their genomes are known as Shiga toxin-converting bacteriophages or Stx phages [7–9]. The recent outbreak that occurred in Germany in 2011 resulted in over 4,000 symptomatic infections, including over 50 fatal cases. This can be an indication of severity of STEC-mediated infections and significance of medical problems caused by bacteria lysogenic with these phages [10–14].

In this light, detailed understanding the mechanisms of regulation of lambdoid phages’ development appears crucial, particularly because Stx prophage induction and effective replication of the phage genome are indispensable for efficient production of Shiga toxins [15–18]. This stems from the fact that expression of majority of genes of lambdoid prophages, including stx genes (for reviews, see [2,3]). In the case of Shiga toxin-converting prophages, among such factor and agents there are UV irradiation, antibiotics interfering with DNA metabolism (like mitomycin C), and hydrogen peroxide (which appears to be the most plausible compound causing induction of Stx prophages) [20,21]. However, despite determination of molecular principles of cl-mediated regulation of gene expression, we are still far from complete understanding of mechanisms influencing efficiency of lambdoid prophage induction and its further lytic development.

One mystery in lambdoid phage biology is the b region in the viral genome. It is dispensable for phage development under standard laboratory conditions (a phenomenon which is unusual among viruses), but contains an evolutionarily conserved fragment, located between exo and xis genes and transcribed from the p1 promoter. This fragment is called the exo-xis region, and consists of several open reading frames which functions in phage development are largely unknown. Previous studies demonstrated that overexpression of genes from the exo-xis region caused impairment lysogenization of E. coli by bacteriophage λ [22]. Subsequent report indicated that the presence of multiple copies of these genes on plasmids positively influenced efficiencies of induction of prophages λ and Φ24B, one of Shiga toxin-converting phages [23]. Interestingly, it was also found that two orthologs of the λ Ea8.5 protein, encoded by a gene located between exo and xis, contain a fused homeodomain/zinc-finger fold [24]. This strongly suggest a regulatory role of this protein.

In the light of the above facts, we aimed to investigate the exo-xis region in more details. Genetic maps of exo-xis regions from genomes of bacteriophages λ and Φ24B are shown in Fig. 1. We asked what are patterns of expression of genes from this region in E. coli cells either infected with bacteriophage λ or Φ24B, or lysogenic with these phages (after prophage induction). Unexpectedly, we found that in both phages λ and Φ24B, these patterns were significantly different not only between conditions of infection of the host cells by bacteriophages and prophage induction, but also between induction of prophages with various agents (mitomycin C and hydrogen peroxide). This may shed a new light on our understanding of regulation of lambdoid phage development, depending on the mode of lytic cycle initiation.

**Materials and Methods**

**Bacterial strains, bacteriophages, and plasmids**

Phages Φ24 (Δstx2::cat) [25] and λ papa (from our collection) were employed in this study. Bacteriophage suspensions were routinely stored in TM buffer (10 mM Tris-HCl, 10 mM MgSO4, pH 7.2) at 4°C. E. coli MG1655 strain [26] was the host of choice for bacteriophage infection and prophage induction experiments. Plasmids pGAW3775tet (bearing phage λ exo-xis region), pSBex-xis Φ24B (as pGAW3775tet but bearing the exo-xis region from phage Φ24B), and pJW0tet (pGAW3775tet with removed λ exo-xis region), used in this work, have been described [23]. Derivatives of pGAW3775, constructed previously [22] and bearing various fragments of the λ exo-xis region, are as follows: pJWexa8.5 (pJW0tet bearing the exa8.5 gene), pJWeca22 (pJW0tet bearing the ec22 gene), pJWorf (pJW0tet bearing orf61, orf60a and orf63 open reading frames), pJWeorfca22 (pJW0tet bearing orf61, orf60a and orf63 open reading frames and the ec22 gene), and pJWe22eca8.5 (pJW0tet bearing exa8.5 and ec22 genes). The frameshift mutations in each of the analyzed ORFs (orf60a, orf63, orf61, orf73) and genes (exa22, ec22, 8.5) within the λ exo-xis region of pGAW3775tet plasmid were introduced separately by deleting one base pair, to produce plasmids pGAWorf60a_mut, pGAWorf63_mut, pGAWorf61_mut, pGAWorf73_mut, pGAWeca22_mut, and pGAWeca8.5_mut, respectively. The site-directed mutagenesis was performed using GeneArt Site-Directed Mutagenesis PLUS System and AccuPrime Pfu polymerase, purchased from Life Technologies, and according to the manufacturer’s protocol. Primers used in the mutagenesis, with indication of the deleted nucleotide relative to the wild-type allele, are listed in Table 1.

**Prophage induction experiments**

Bacteria lysogenic with tested phages were cultured in Luria–Bertani (LB) medium at 30°C to a 0.8 of 0.1. Three induction conditions were tested: 0.2 μg/ml mitomycin C, 1 mM H2O2, and UV irradiation (50 J/m2). At indicated times after induction, samples of bacterial cultures were harvested, and 30 μl of chloroform were added to 0.5 ml of each sample. The mixture was vortexed and centrifuged for 5 min in a microcentrifuge. Bacteria were grown in LB medium supplemented with MgSO4 and CaCl2 (to a final concentration of 10 mM each). Plates were incubated at 37°C overnight. Analogous experiments but without induction agents were performed (control experiments) with each lysogenic strain. Presented values show phage titer (PFU/ml) normalized to results withdrawn and centrifuged (3,000 x g, 10 min). Each pellet was suspended in 1 ml (1/10 of initial volume) of 3 mM NaN3 in LB. Following 5-min incubation at 30°C, the phage was added to
multiplicity of infection (m.o.i.) of 0.05. Phage adsorption was carried out at 30°C for 10 min. The mixture was diluted ten-fold in warm (30°C) 3 mM NaN₃ in LB and centrifuged (3,000 x g, 10 min). Bacterial pellet was suspended in 1 ml of LB with 3 mM NaN₃ and centrifuged again (3,000 x g, 10 min). This procedure was repeated three times. The suspension was then diluted 1,000-fold with LB, prewarmed to 30°C (time 0), and aerated in a water bath shaker at this temperature. The number of infective centers was estimated from nine samples taken in the interval of 0–15 min after the dilution by plating under permissive conditions. The number of intracellular progeny phages (samples previously shaken vigorously for 1 min with equal volume of chloroform and cleared by centrifugation) was estimated by plating on appropriate indicator bacteria. Plates were incubated at 37°C overnight. Each experiment was repeated three times.

Table 1. Primers used for site-directed mutagenesis.

| Primer name                  | Sequence (5’→3’)                          | Deleted nucleotide |
|------------------------------|-------------------------------------------|--------------------|
| pF_pGAW3775tet_mut_orf60a   | CAATCACTTTCGTTCTCCCGTTACAAGCGAG [A]       |                    |
| pR_pGAW3775tet_mut_orf60a   | CTCGCCCTTTAAGCGAGAAGGGAAGTGATTG           |                    |
| pF_pGAW3775tet_mut_orf63    | CAAAGCATCTTCTGTGGATTAAAGAACGAGTATC [G]    |                    |
| pR_pGAW3775tet_mut_orf63    | GATACTCGTCTTTAACTAAAGAGATGCTTTT           |                    |
| pF_pGAW3775tet_mut_orf61    | CTTCTATCTTTGTTGCTTAGTGCTTGGGCGCATAG     | [C]                |
| pR_pGAW3775tet_mut_orf61    | ATGTGCGGATTACACACACAGAATAGAAG             |                    |
| pF_pGAW3775tet_mut_orf73    | GAAATAGAAGAATTACGCCACACAGAATGAA          | [A]                |
| pR_pGAW3775tet_mut_orf73    | TATTTGCTGTGTGTGCACCCAGAATCTCTCTCTATTTC   |                    |
| pF_pGAW3775tet_mut_ea22     | TGGGATTGTGACGGATCCCTTTCCATGAATTG         | [A]                |
| pR_pGAW3775tet_mut_ea22     | CAATTCATGAAAAGGCTTGCTCAATAATACCCCA       |                    |
| pF_pGAW3775tet_mut_ea8.5    | TTATCAATTGTCGTTGCGATGCAATCTACTACATGC    | [A]                |
| pR_pGAW3775tet_mut_ea8.5    | GAGACAAAGACCCGGACTGCACACATTTGAA          |                    |

The asterisk (*) indicate position of deleted nucleotide in forward primer. Analogous deletion was introduced in the reverse primer but is not shown for clarity of presentation.

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Figure 1. Maps of exo-xis regions and other genes of bacteriophages λ and Φ²48 analyzed in this work (accession numbers: GI:9626243 and GI:307604077 respectively). Dark arrows with continuous outside lines represent highly conserved genes and open reading frames (over 70% nucleotide as well as amino acid sequence identity). Dark arrows with punctuated outside lines represent highly conserved (above 70% sequence identity) open reading frames present in genomes of λ and 933W phages, available in the NCBI database but uncharacterized in annotations. The presence of orf73 in the λ exo-xis region was indicated by [48]. Light arrows represent genes and open reading frames with low level of sequence identity (<38%). Note the high homology between λ and Φ²48 exo-orf73 regions and low level of identity of other analyzed genes. Arrows indicate positions of promoters predicted with BPROM program. The localizations and −10 and −35 sequences of predicted promoters p₁, λ and p₁, Φ²48 are exactly the same (see Table 3). Schematic steam-loop structures (●) indicate localizations of predicted transcription terminators, found on the basis of nucleotide sequence analysis with ARNold software. The localizations and sequences of predicted terminators t₁ and t₂ are exactly the same in case of both phages λ and Φ²48 (see Table 4). Note that in the case of Φ²48 phage, some ORFs from the exo-xis region: vb_24B_9c, vb_24B_8c, vb_24B_7c, putative C4 zinc finger protein and vb_24B_6c are homologues of λ orf60a, orf63, orf61, orf73 and gene ea22 respectively. For clarity of this work, only the names of λ ORFs were used.

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Figure 1. Maps of exo-xis regions and other genes of bacteriophages λ and Φ²48 analyzed in this work (accession numbers: GI:9626243 and GI:307604077 respectively). Dark arrows with continuous outside lines represent highly conserved genes and open reading frames (over 70% nucleotide as well as amino acid sequence identity). Dark arrows with punctuated outside lines represent highly conserved (above 70% sequence identity) open reading frames present in genomes of λ and 933W phages, available in the NCBI database but uncharacterized in annotations. The presence of orf73 in the λ exo-xis region was indicated by [48]. Light arrows represent genes and open reading frames with low level of sequence identity (<38%). Note the high homology between λ and Φ²48 exo-orf73 regions and low level of identity of other analyzed genes. Arrows indicate positions of promoters predicted with BPROM program. The localizations and −10 and −35 sequences of predicted promoters p₁, λ and p₁, Φ²48 are exactly the same (see Table 3). Schematic steam-loop structures (●) indicate localizations of predicted transcription terminators, found on the basis of nucleotide sequence analysis with ARNold software. The localizations and sequences of predicted terminators t₁ and t₂ are exactly the same in case of both phages λ and Φ²48 (see Table 4). Note that in the case of Φ²48 phage, some ORFs from the exo-xis region: vb_24B_9c, vb_24B_8c, vb_24B_7c, putative C4 zinc finger protein and vb_24B_6c are homologues of λ orf60a, orf63, orf61, orf73 and gene ea22 respectively. For clarity of this work, only the names of λ ORFs were used.

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Table 2. Primers used in the real-time PCR assay.

| Primer name | Sequence (5′ → 3′) |
|-------------|---------------------|
| pF_0248_int | CAGTTGCCGGTATCCCTGT |
| pR_0248_int | TGAGGCTTCTCGTGGTCA |
| pF_0248_ea22 | TCGACGACTGACCATCAGC |
| pR_0248_ea22 | GGTGAAAGGCTAGAGTGT |
| pF_0248_orf73 | CGAACCTCTCTTTAATCTCT |
| pR_0248_orf73 | TCCAGGGTGGTCCGACCTT |
| pF_0248_orf61 | TTAGGCTGACGGGCAATG |
| pR_0248_orf61 | CCGACATGGAGACTTCA |
| pF_0248_orf63 | GGTGGCTCTCCTGTTGTC |
| pR_0248_orf63 | TAGGCAATCTCCCTTCAAA |
| pF_0248_orf60a | CATAAGGCTCTCCTGTTTAT |
| pR_0248_orf60a | CCGAATCCGGAAGAGCAC |
| pF_0248_N | AGGCCTTTGCTGAGACCTT |
| pR_0248_N | TACACCGGCCCTCTGCTAAAG |
| pF_0248_cro | CGAAGGGCTTGTGGGAGTTAG |
| pR_0248_cro | GTCTTGGGAGGACTGAGT |
| pF_0248_Q | GGGAGTGAGGCTTGAGATGG |
| pR_0248_Q | TACAGAGGTTCTCCCTCCCG |
| pF_0248_R | GGGTGATGTGTAAGCTCTG |
| pR_0248_R | TAACCCGGTCCGGATTTC |
| pF_l_int | TTTGATTCTCAATTTTGTCCCA |
| pR_l_int | ACCATGCGCACAAGATTCG |
| pF_l_ea8.5 | GGGCAAGTATCGTCCCTC |
| pR_l_ea8.5 | GCAATGTCGGAAGAAATGACTG |
| pF_l_ea22 | GCAGTCCAGCAACTGAT |
| pR_l_ea22 | AATGCAATGACGACTG |
| pF_l_orf73 | CACCTGAACCTCTCTGTTTACTG |
| pR_l_orf73 | CAGGGGTTACTGGACTTG |
| pF_l_orf61 | TTAGGCTGACGGGCAATG |
| pR_l_orf61 | CCGACATGGAGACTTCA |
| pF_l_orf63 | ACCTGTGCTCTCTACCTG |
| pR_l_orf63 | GATAGCCCAGATCTCCCTC |
| pF_l_orf60a | GCATACAGCCCCCTGTATT |
| pR_l_orf60a | CCGAAATCCATGGAAGAC |
| pF_l_N | CTCCTGATTCCTGTTGCG |
| pR_l_N | AAGCAGCATAAAAATCCCTTG |
| pF_l_cro | ATGCGGAGAGGTAAGGCCC |
| pR_l_cro | TGGAAATTGTAAGACGCC |
| pF_l_Q | TTCTGCGTCAAGACGAC |
| pR_l_Q | TGCATCAGATTTGAGTCCTTT |
| pF_l_R | ATGCACGATGAGAC |
| pR_l_R | GCTGCAACTGACCATACCA |
| pF_E.coli_icdA | CGAAGCGGCTGACCTT |
| pR_E.coli_icdA | GCTGCAACTGACCATACCA |
| pF_E.coli_16SrRNA | CCTTACGACCAGGGCTACAC |
| pR_E.coli_16SrRNA | TTTAGGGCCGCTTGCTC |

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Bacterial suspensions were stained with LIVE/DEAD supernatants were removed and pellets washed in 0.85% NaCl. Each experiment was repeated three times.

Presented values show percent of bacterial suspensions at emission = 530 nm by the fluorescence intensity at emission = 630 nm. SYTO 9 dye, and 630 nm for propidium iodide dye [28]. Data were analyzed by dividing the fluorescence intensity of stained bacteria with intact as well as damaged membranes (e.g. arising as a result caused by phage host-cell lysis). In contrast, propidium iodide penetrates only into bacteria with damaged membranes, causing a reduction in the green dye fluorescence when both stains are present. Measurements of fluorescence were performed in microplate reader using excitation wavelength = 485 nm and emission wavelengths = 530 nm for SYTO 9 dye, and 630 nm for propidium iodide dye [28]. 

Measurement of bacterial viability during prophage induction experiments

Bacteria lysogenic for tested phages were cultured in LB medium at 30°C to A_{600} = 0.1 with induction agent, either 0.2 μg/ml mitomycin C or 1mM H_2O_2, added at time zero. At indicated times after induction, samples equal to 2×10^9 cells/ml were withdrawn and centrifuged at 10,000 ×g for 10 min. The supernatants were removed and pellets washed in 0.85% NaCl. Bacterial suspensions were stained with LIVE/DEAD BacLight Bacterial Viability Kit (Molecular Probes), according to the manufacturer’s protocol. The mentioned kit utilizes mixtures of the green-fluorescent nucleic acid stain SYTO 9 and the red-fluorescent nucleic acid dye, propidium iodide. When used alone, the SYTO 9 dye stains bacteria with intact as well as damaged membranes (e.g. arising as a result caused by phage host-cell lysis). In contrast, propidium iodide penetrates only into bacteria with damaged membranes, causing a reduction in the green dye fluorescence when both stains are present. Measurements of fluorescence were performed in microplate reader using excitation wavelength = 485 nm and emission wavelengths = 530 nm for SYTO 9 dye, and 630 nm for propidium iodide dye [28].

Data were analyzed by dividing the fluorescence intensity of stained bacterial suspensions at emission = 530 nm by the fluorescence intensity at emission = 630 nm. Presented values show percent of live bacteria normalized to results of control experiment, non-induced cultures which at each time were assumed as 100% of live bacteria. Each experiment was repeated three times.

Estimation of relative phage DNA amounts

Bacteria lysogenic for tested phages were cultured in LB medium at 30°C to A_{600} = 0.1. Induction of prophages was provoked in lysogenic bacteria by addition of mitomycin C to a final concentration of 0.2 μg/ml or H_2O_2 to a final concentration 1 mM. At indicated times after induction, 2-ml samples with 120 μl of chloroform were vortexed for 10 s and centrifuged in a microcentrifuge for 5 min. The supernatants were collected and filtered through a membrane filter with pore size of 0.22 μm (Sigma-Aldrich) to remove bacterial cells. Filtered samples were first treated with DNase I (20 μg/ml; Sigma-Aldrich) for 30 min at 37°C to remove any free bacterial DNA. The viral DNA was then liberated from virions using the method described by [29], with a minor modification. All DNase-treated samples were heated at 37°C for 60 min in the presence of 20 mM EDTA (Sigma-Aldrich) and 50 μg/ml proteinase K (Sigma-Aldrich). DNA was quantified by staining with Qubit dsDNA BR Assay Kit (Invitrogen), according to the manufacturer’s instructions. Concentration of phage DNA (in μg/ml) were calculated relative to analogous experiments but without induction agents (control experiments) with each lysogenic strain. Presented values show phage DNA concentration (μg/ml) normalized to results of control experiments. Each experiment was repeated three times.

Bacteriophage infection

Host bacteria were grown to A_{600} of 0.3 at 30°C. Then, 120 ml volume was centrifuged and the pellet was washed with 30 ml of 0.85% NaCl. After centrifugation, the pellet was suspended in 36 ml of LB medium supplemented with MgSO_4 and CaCl_2 (to a final concentration of 10 mM each). The mixture was incubated for 30 min at 30°C and then chilled on ice. Bacteriophage lysate was added to m.o.i. of 5. Following 30 min incubation on ice, at indicated times, 1×10^9 bacterial cultures were treated with NaN_3 (Sigma-Aldrich) to a final concentration of 10 mM and harvested. The preparation of RNA and cDNA were performed as described in the subsequent subsection.

Preparation of RNA and cDNA from bacteria

For the preparation of RNA, the induction of temperate bacteriophages from E. coli strain MG1655 was performed with
Figure 3. Development of λ (panels A, B, C, D) and F24b (panels E, F) bacteriophages after prophage induction with 0.2 μg/ml mitomycin C (panels A, C, E) or 1 mM H2O2 (panels B, D, F) in E. coli MG1655 host at 30°C. The results with host cells containing the pJW0tet vector (open squares) or a plasmid bearing the whole exo-xis region from with λ (pGAW3773tet) or F24b (pSBe.x.r) (closed squares) are presented in each panel. In other experiments presented in panels A and B, bacteria contained plasmids bearing following parts of this region from λ: the ea8.5 gene (pJWea8.5; closed circles), the ea22 gene (pJWea22; open circles), orf61, orf60a and orf63 (pJWorf; open diamonds), orf61, orf60a, orf63 and ea22 (pJWorfea22; open triangles), ea22 and ea8.5 (pJWea22ea8.5; closed triangles). In other experiments presented in panels C and D, bacteria contained plasmids bearing the whole exo-xis region but with a frameshift mutation in one of following ORFs or genes: orf60a (pGAWorf60a_mut; open diamonds), orf63 (pGAWorf63_mut; closed diamonds), orf61 (pGAWorf61_mut; open triangles), orf73 (pGAWorf73_mut; closed triangles), ea22 (pGAWea22_mut; open circles) or ea8.5 (pGAWea8.5_mut, closed circles). The presented results are mean values from 3 experiments with error bars indicating SD (note that in the most cases the SD were smaller than sizes of symbols).

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mitomycin C (final concentration 0.2 mg/ml) or H2O2 (final concentration 1 mM) as described in previous subsections. To inhibit the growth of bacteria, all samples were treated with NaN3 (Sigma-Aldrich) to a final concentration of 10 mM. Total RNA was isolated from 1×10^9 bacterial cells with the High Pure RNA Isolation Kit (Roche Applied Science). RNA preparations were repeatedly digested with TURBO DNase from TURBO DNA-free Kit (Life Technologies) for 60 min at 37°C, as described by the manufacturer. To evaluate the quality and quantity of total isolated RNA, we used a NanoDrop spectrophotometer, considering the ideal absorbance ratio (1.8≤A260/A280≤2.0), and visualized the band patterns of total RNA by electrophoresis. The absence of DNA from RNA samples was controlled by PCR amplification, and by real-time PCR amplification of the all tested genes. RNA preparations were stored at −80°C for use. The preparation of cDNA from the total RNA samples (1.25 μg) was performed with Transcripter Reverse Transcriptase and random hexamer primers (Roche Applied Science), following the instructions supplied by the manufacturer. cDNA reaction mixtures were diluted 10-fold for use in real-time PCR.

Real-time PCR Assay

For transcriptional analysis of tested genes by quantitative real-time reverse transcription-PCR (qRT-PCR), the qRT-PCR was performed with the LightCycler 480 Real-Time PCR System (Roche Applied Science), with cDNA samples from lysogenic bacteria. Transcription rates of ϕ24B and λ genes were compared in parallel to those of the icdA (according to a procedure described by [30]) or 16S rRNA housekeeping genes. Primers were developed by Primer3web version 4.0.0 and produced by Sigma-Aldrich or GENOMED. The transcriptional analysis of ϕ24B and λ genes was performed with primers presented in Table 2. Real-time PCR amplifications were performed for 55 cycles in 20-μl reaction volumes by employing LightCycler 480 SYBR Green I

Figure 4. Relative level of DNA of λ (panels A and B) or ϕ24B (panels C and D) bacteriophages after prophage induction with 0.2 μg/ml mitomycin C (panels A and C) or 1 mM H2O2 (panels B and D) in E. coli MG1655 host at 30°C. Host cells contained either the pJW0tet vector (open squares) or a plasmid containing the exo-xis region from with λ (pGAW3773tet; panels A and B) or ϕ24B (pSBe.x.r; panels C and D) (closed squares). The presented results are mean values from 3 experiments with error bars indicating SD.

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Reactions were performed in Roche 96-well plates containing 10 µl 2x SYBR Green I Master Mix, 6.25 ng/µl cDNA and 200 nM of each gene-specific primer (Table 2). Relative quantification assays were performed with cDNA in an icdA or 16S rRNA and phage genes multiplex assay. All templates were amplified using the following program: 95°C for 5 min; 55 cycles of 95°C for 10 s; 60°C for 15 s and 72°C for 15 s. No template control was included with each run. Each

Table 3. Predicted promoters for the orf73 coding region of bacteriophages λ and Φ24b.

| Promoter name | Strand | −10 box | −35 box | Promoter score | Elements of predicted transcriptional factor binding sites |
|---------------|--------|---------|---------|----------------|--------------------------------------------------------|
| p1_λ          | Minus  | TTTTATTAT TCA TCA | 4.82 | rpoD17: CTCCTTT argR: TTT TTTT TTT |
|               |        |         |         |                | argR2: TTT TTTT TTT |
| p1_Φ24b       | Minus  | TTTTATTAT TCA TCA | 4.82 | rpoD17: CTCCTTT argR: TTT TTTT TTT |
|               |        |         |         |                | argR2: TTT TTTT TTT |
| p2_λ          | Minus  | TCA TCA TCA TCA | 0.59 | lrp: TGATTTTT |
|               |        |         |         |                | fadR: GGA TTG |

Figure 5. Survival of *E. coli* MG1655 cells lysogenic for λ (panels A and B) or Φ24b (panels C and D) bacteriophages after prophage induction with 0.2 µg/ml mitomycin C (panels A and C) or 1 mM H2O2 (panels B and D) at 30°C. Host cells contained either the pJW0tet vector (open squares) or a plasmid containing the exo-xis region from with λ (pGAW3773tet; panels A and B) or Φ24b (pSBe.x.r; panels C and D) (closed squares). The presented results are mean values from 3 experiments with error bars indicating SD.

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Table 4. Predicted terminators in the *exo-xis* regions of bacteriophages λ and Φ24B.

| Terminator name | Strand | Terminator sequence | Program that produced prediction | Score [Free energy of stem-loop region (kcal/mol)] |
|-----------------|--------|----------------------|----------------------------------|-----------------------------------------------|
| t₁,λ | Minus | TTACAAAGCGAGCTGGGAT7TCCGCGCTITTGTATCC | RNAmotif | –14.10 |
| t₂,λ | Minus | AAAATCTACAGAGCTAGGCTCCCTTTTTATATT | RNAmotif | –12.70 |
| t₃,λ | Minus | TTACAAAGCGAGCTAGGCTCCCTTTTTATATT | Erpin and RNAmotif | –14.10 |
| t₄,λ | Minus | TTATCTGAGCTAGGCGGTITTGTATCG | Erpin and RNAmotif | –14.10 |
| t₅,λ | Minus | AAGAACACCAAGCGCTGATGCGGTTTTCTGCGTG | Erpin and RNAmotif | –14.10 |
| t₆,λ | Minus | TCAACTAACCAACCGCCCTGGGCGGTTTATTATGGCTG | RNAmotif | –14.10 |

Secondary structures are indicated, where loops are in italic font and stems in bold underlined font. The sequences of predicted terminators t₁ and t₂ are exactly the same in the case of both phages λ and Φ24B.

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Figure 6. Expression patterns of genes from the *exo-xis* region, as well as *int*, *N*, *cro*, *Q* and *R* genes of bacteriophage λ (panel A) and Φ24B (panel B) infecting *E. coli* MG1655 host at 30°C. Levels of transcripts corresponding to particular genes or ORFs were determined at following times after infection: 2.5 (violet), 5 (green), 7.5 (red), 10 (white), 20 (black), and 30 (orange) minutes in case of phage λ and 10 (white), 20 (black), 30 (orange), 40 (gray), 50 (yellow), and 60 (blue) minutes in case of phage Φ24B. The presented results are mean values from 3 experiments with error bars indicating SD.

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Real-time PCR data analysis

To analyze the relative changes in gene expression revealed by quantitative Real-Time PCR experiments, the calibrator normalized relative quantification method with efficiency correction (so-called E-Method) was used. The E-Method provides an efficiency corrected calculation mode by using the determined PCR efficiency of target (Eₜ) as well as the efficiency of reference (Eᵣ). Relative fold change ratio was calculated by using the following formula, described in the application manual of Roche LightCycler Real-Time PCR Systems [31]:

\[ \text{Relative expression ratio} = \frac{E_\text{t} \times CT(\text{t})}{E_\text{r} \times CT(\text{r})} \]

(where “t” is target, and “r” is reference).

The sample at the time point “zero” was a calibrator. The raw run data for Φ24B and λ genes were transferred from the LightCycler 480 to the LinRegPCR 12.5 software using the “LC480 Conversion: conversion of raw LC480 data” software (available at http://www.hartfaalcentrum.nl/index.php?main=files&sub=0). PCR efficiency was determined for each gene by LinRegPCR program [32,33]. This software was successfully used previously to calculate PCR efficiency [34–38].

Prediction of the presence of transcription promoters and terminators in phage genomes

Prediction of the presence of promoters in the sequences of genomes of λ and Φ24B phages was performed using BPROM—the bacterial β’ promoter recognition program available at: http://linux1.softberry.com. Promoters were searched within ~400 bp long sequence fragment before the start of the *orf73* coding region, encompassing whole sequence of *orf61* and the region between *orf61* and *orf73*. BPROM has accuracy of 60–80%, and considers promoters with score above 0.20 [39].

Promoters were searched within ~400 bp long sequence fragment before the start of the *orf73* coding region, encompassing whole sequence of *orf61* and the region between *orf61* and *orf73*. BPROM has accuracy of *E. coli* promoter recognition about 80%, and considers promoters with score above 0.20 [39].

Predicted terminators were found within the *exo-xis* region of λ and Φ24B using ARNold, the online analysis tool which predicts the existence and location of rho-independent transcription terminators using RNAmotif and ERPIN complementary programs [40–43]. The ARNold program is available at: http://rna.igmors.u-psud.fr/toolbox/arnold/.
Effects of the multicopy exo-xis region on λ and ϕ24B development and host survival at 30°C

Previous studies demonstrated that the presence of the exo-xis region on a multicopy plasmid in the host cells caused enhanced efficiency of induction of prophages λ and ϕ24B under standard laboratory conditions (rich medium, 37°C). Since the main aim of this work was to determine detailed patterns of expression of genes from the exo-xis regions of the tested lambdoid phages, after preliminary experiments, we decided to culture host bacteria at 30°C, rather than at 37°C, to slow metabolic processes down which made the analyzes more precise. However, such a change in cultivation conditions relative to previously reported studies (temperature 30°C instead of 37°C) made it necessary to check whether effects of multiple copies of the exo-xis region on phage development are similar in both experimental systems.

We found that lytic developments of both tested phages, λ and ϕ24B, after infection of the E. coli cells growing at 30°C, were not significantly affected by the presence of corresponding exo-xis regions on multicopy plasmids (Fig. 2). In the case of phage λ, effects of the presence of particular fragments of this region were also tested, again with no considerable changes detected (Fig. 2). This corroborates results of analogous experiments performed at 37°C and reported previously [23].

Accordingly to previous studies [23], we have also tested development of both tested phages in the presence of the plasmid-borne exo-xis region after prophage induction with UV irradiation. Similarly to experiments with mitomycin C and hydrogen peroxide (Fig. 3), production of phage progeny of both λ and ϕ24B started earlier and was more efficient when additional copies of the exo-xis region were present in host cells (data not shown). Because of these similarities, in further experiments we have focused on prophage induction conditions which are more likely to occur in the intestine, a natural environment of the host bacterium [4–9], i.e., the presence of an antibiotic (mitomycin C) or H₂O₂, rather than UV irradiation.

Figure 7. Expression patterns of genes from the exo-xis region, as well as int, N, cro, Q and R genes of bacteriophage λ (panels A and B) and ϕ24B (panels C and D) after prophage induction with 0.2 μg/ml mitomycin C (panels A and C) or 1 mM H₂O₂ (panels B and D) in E. coli MG1655 host at 30°C. Levels of transcripts corresponding to particular genes or ORFs were determined at following times after induction: 64 (red), 128 (gray), 136 (orange), 144 (dark blue), 152 (yellow), 160 (green), 176 (maroon) and 192 (light blue) minutes. The presented results are mean values from 3 experiments with error bars indicating SD.

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Results

Effects of the multicopy exo-xis region on λ and ϕ24B development and host survival at 30°C

This corroborates results of analogous experiments performed at 37°C and reported previously [23].

Studies on λ and ϕ24B development after prophage induction were carried out in the host cells treated with either mitomycin C or hydrogen peroxide. Again, results obtained in experiments conducted at 30°C (Fig. 3) were generally similar to those at 37°C, described in the previous article [23]. However, positive effects of multiple copies of the exo-xis region in both λ and ϕ24B were even more pronounced at 30°C (compare Fig. 3 with [23]). These results also confirmed our presumption that performing the planned experiments at 30°C, rather than 37°C, may allow us to detect any putative differences more efficiently.
Finding experimental conditions (30°C rather than 37°C) causing more pronounced differences in development of bacteriophages between hosts containing and lacking additional copies of the exo-xis region allowed us to address the problem whether expression of genes and/or ORFs from this region is responsible for the observed effects. This was reasonable as one might assume that the tested fragment of phage genome might bind regulatory factor(s), titrating it/them out, and thus causing changes is the control of viral development.

To test such a possibility, we have employed the λ model with two experimental systems. First, instead of using the whole exo-xis region, host cells were transformed with plasmids bearing particular genes or ORFs or their combinations. Second, we have constructed a series of derivatives of pGAW3775tet (a plasmid bearing the whole exo-xis region) where each plasmid contains a frameshift mutation in particular gene or ORF. In the first experimental system, the absence of particular fragments of the exo-xis region relative to pGAW3775tet resulted is intermediate effects, i.e. the phage development was more effective than in the host bearing the vector, but less effective than in the pGAW3775tet-bearing host (Fig. 3 A and B). However, in some cases, specific effects were detected, namely, the presence of either pJWea22 (bearing ea22) or plWorfca22 (bearing orf61, orf60a, orf63 and ea22) did not result in more rapid induction or phage development in the mitomycin C-treated host (Fig. 3A), and the presence plWorfca22 (bearing ea22) caused a delay in the phage development in hydrogen peroxide-treated cells (Fig. 3B).

In the second experimental system, frameshift mutations in orf61 and ea22 (in the exo-xis fragment present in the plasmid) resulted in a delay in phage development after prophage induction with mitomycin C (Fig. 3C), while frameshift mutations in orf61, orf73 and ea8.5 caused similar effects in hydrogen peroxide-treated bacteria (Fig. 3D). In addition, we have measured the frequency of spontaneous (without addition of any specific agent) prophage induction to find similar values in cells bearing pGAW3775tet (with wild-type exo-xis region) and most of constructs with frameshift mutations in one of genes or ORFs (the frequencies were about 10^{-5} per cell), which were about 10 times higher than in bacteria bearing a control plasmid pJW0tet (about 10^{-6} per cell). However, a frameshift mutation in orf60a abolished the effect caused by the presence of a plasmid with the exo-xis region (the frequency was about 10^{-6} per cell). Although these results did not exclude a possibility for titrating out regulatory factors(s) by the exo-xis region, they suggested specific roles of expression of at least some genes and ORFs, particularly ea22, ea8.5, orf61, orf73, and orf60a.

Additional confirmation of the more efficient development of phages λ and φ24B in cells treated at the lysogenic stage with mitomycin C or hydrogen peroxide was provided by measurement of an increase in bacteriophage DNA amount. Again, more efficient increase in the level of DNA of both tested phages was observed after induction of corresponding prophages with both tested inducers in cultures of hosts bearing plasmids with appropriate exo-xis region relative to those containing plasmid vector (Fig. 4). As expected, in the same experimental system, survival rate of bacteria with the vector was always higher than cells bearing a plasmid with the exo-xis region (Fig. 5). Survival of a high percentage of bacterial cells after the induction might seem surprising. However, one should note that contrary to UV irradiation, the efficiency of prophage induction after treatment of the host (lysogenic with λ or φ24B) with mitomycin C or hydrogen peroxide may be moderate. In fact, experimental data indicated that in mitomycin C- or hydrogen peroxide-treated cultures of such bacteria, prophage induction occurred in less than 50% or even in only a few percent of cells, respectively [20,21]. Therefore, relatively large fraction of cells may survive, and due to lysogenic state, they are immune to superinfection by the same phage. This may allow these cells to grow and divide, resulting in values even over 100% in the employed experimental system (Fig. 5).

In summary, these results clearly indicated that the presence of the exo-xis region on a multicopy plasmid stimulates development of bacteriophages λ and φ24B after treatment of lysogenic cells with either mitomycin C or hydrogen peroxide at 30°C.

Expression of genes from the exo-xis regions of bacteriophages λ and φ24B in infected host cells

To assess patterns of expression of genes from the exo-xis region in phage-infected bacteria, E. coli wild-type (MG1655) cells were infected with either λ or φ24B, and at certain times after infection total RNA was isolated and levels of tested transcripts were determined by quantitative real-time reverse transcription PCR. The exo-xis region is believed to be transcribed from the leftward P1 promoter, one of two major lytic promoters of lambdoid phages [3,8]. Positions and sequences of predicted promoters and terminators located in the exo-xis regions of genomes of both tested phages were predicted, and they are indicated in Fig. 1 and summarized in Tables 3 and 4, respectively.

Lytic development of bacteriophage λ is quicker than that of φ24B, as demonstrated in Fig. 3, therefore, it is not a surprise that expression of the major genes of phage lytic development, N and cro, occurred earlier after the infection with the former phage than in the latter one (Fig. 6). The presence of low level signals, followed by a maximal amount of the transcript, and finally by decreased intensities of signals in the reactions of detection of N- and cro-specific mRNAs indicates that appropriate time-frames were chosen to assess the expression efficiency. Another proof of the properly chosen times of sample withdrawn are low levels of mRNAs for Q and R genes, coding for proteins synthesized at the late stage of bacteriophage development.

Interestingly, some of the genes from the exo-xis regions were expressed (at the RNA level) as efficiently as, or with a similar efficiency to, the N gene. This was true for orf60a, orf63, and orf61 in phage λ, and orf60a, orf63, orf73, and ea22 in phage φ24B (Fig. 6). Moreover, although levels of mRNA for orf73, ea22 and ea8.5, which are downstream of the predicted terminator (located between orf61 and orf73), were significantly lower than genes located upstream of the terminator in λ, considerably different expression pattern has been observed in φ24B. In host cells infected by the latter phage, orf61 was poorly expressed, while levels of mRNAs for orf73 and ea22 were as high as those for N, orf61a and orf63 (Fig. 6). Interestingly, upstream of orf73 and ea22, a promoter P1,φ24B was predicted by BPROM program (Fig. 1 and Table 3). The P1,φ24B promoter sequences −10 and −35 are identical with analogous sequences of the P1,λ promoter, localized upstream of the λ orf73.

Expression of genes from the exo-xis regions of bacteriophages λ and φ24B after prophage induction with various agents

Expression of phage genes after prophage induction was assessed in host cells treated with either mitomycin C or hydrogen peroxide. In both tested phages, λ and φ24B, characteristic time-course of mRNA amounts for N and cro genes, encompassing low level, maximum, and decreased levels, was achieved at significantly later times after induction than after infection (Fig. 7 and Fig. 6, respectively). In phage λ, levels of N and cro transcripts
were significantly lower than those of genes from the exo-xis region, which differs from the pattern determined in phage-infected cells. Moreover, N and evo were expressed at similar times to Q expression, which again indicates the difference between two variants of initiation of the phage lytic development (infection vs. induction). Additionally, in the case of mitomycin C-induced λ prophage, the level of expression of ea8.5, ea22 and orf73 was significantly decreased in comparison with other ORFs from the exo-xis region (Fig. 7A) what could be explained by the presence of predicted t2_1 and t2_2 transcription terminator, localized between orf61 and orf73 (Fig. 1 and Table 4).

In phage Φ24B, expression of all tested genes was delayed after the prophage induction relative to initiation of the lytic development by infection (Fig. 7 and 6, respectively). Nevertheless, expression patterns were quite similar between mitomycin C-treated lysogenic cells and bacteria infected with viruses. Surprisingly, when the prophage induction was caused by treatment of host cells with hydrogen peroxide, levels of mRNAs for orf73 and ea22 were significantly higher than other tested genes. This expression pattern differs considerably from that observed in mitomycin C-induced lysogens and might be explained by the presence of the p1_Φ24B promoter upstream of orf73 (Fig. 1 and Table 3). On the other hand, in both cases, hydrogen peroxide- and mitomycin C-induced Φ24B prophages, the change in the expression level was observed between orf61 and orf73 (Fig. 7C and D). At this point, it should be noted that localizations and sequences of predicted transcription promoters p1_1 and p1_Φ24B, as well as terminators t2_1 and t2_Φ24B are exactly the same (Fig. 1 and Table 4).

Discussion

Although bacteriophage λ and related phages have been used as models for genetic studies, including regulation of gene expression, reports on global analyses of gene expression (especially time course of the expression) of lambdoid viruses are rare in the literature. Recently, expression pattern of phage λ genes after thermal induction of the prophage bearing a temperature-sensitive mutation in the cl gene was investigated by ribosome profiling [44], and genes’ expression of phage Φ24B in the lysogenic host was studied using a proteomic approach [19]. In both cases, unexpected results were obtained. In bacteriophage λ developing after prophage induction, global gene expression analysis revealed the activity of various previously unappreciated open reading frames [44]. In both λ and Φ24B, apart from cl, rexA, rexB, tom and bor – genes previously known as those transcribed from a prophage, several other genes were found to be expressed in non-induced lysogens; those included ea8.5 in λ, and exo in Φ24B, genes studied also in our work. The recently published reports, mentioned above, indicated quite an unexpected complexity in the patterns of expression of genes of lambdoid phages, suggesting the existence of multiple regulatory systems, not yet identified in these viruses.

In this work, we have investigated expression patterns of genes from the exo-xis regions of phages mentioned above (λ and Φ24B) after infection of host cells or induction of corresponding prophages. We have chosen this genome region because recent studies indicated that genes included there had significant effects on efficiency of lysogenization and prophage induction in both tested phages [22,23]. Moreover, the Eal8.5 protein, encoded in this region, was found to contain a specific motif, strongly suggesting its regulatory role [24].

Similar to two other recent analyses [19,44], our studies led to unexpected results. First, time courses of expression of the investigated genes, including those coding for major regulatory proteins, N and evo, were significantly different in phage-infected cells and in induced lysogens. Second, despite homologous regulatory sequences (promoters and terminators), identified and predicted in genomes of λ and Φ24B, gene expression patterns were significantly different between these two tested phages. Third, even in the same phage, considerably different patterns of gene expression were detected, depending on the nature of agent (mitomycin C or hydrogen peroxide) used to induce the Φ24B prophage.

At the current stage of our knowledge, it is difficult to predict the mechanisms of the differential expression of phage genes during lytic development initiated by different ways, either infection or prophage induction. Even harder to understand is different expression of genes from the exo-xis region of phage Φ24B, when prophage induction is caused by either mitomycin C or hydrogen peroxide. One would expect that both agents should induce the S.O.S. response in the host cells, which should lead to degradation of the cl repressor and subsequent prophage excision, followed by expression of phage genes as in the lytic cycle initiated by infection. Definitely, regulations of these processes are significantly more complicated than assumed.

One example of unexpected specific regulation arises from analysis of the patterns of expression of genes from the exo-xis region of phage Φ24B. In silico analysis predicted the existence of both promoter and terminator between orf61 and orf73, homologous to those in λ. In bacteriophage λ, levels of transcripts for ORFs located upstream of the predicted terminator are lower than those located downstream of this terminator, irrespective of the way of initiation of the lytic development. However, orf73 and ea22 of Φ24B are efficiently expressed despite the presence of this terminator, which is especially well pronounced in cells treated with hydrogen peroxide, where orf73 and ea22 are expressed at significantly higher levels than the rest of the exo-xis region. This might suggest the presence of a promoter upstream of orf73, and it was confirmed by the analysis with the use of BPROM program which allowed us to localize a predicted σ70-dependent promoter upstream of orf73. Despite unknown mechanisms responsible for differential expression of genes from the exo-xis region of bacteriophages λ and Φ24B, the fact that transcripts of ORFs from this region occur at the significant levels during phages' development suggests that they can play important regulatory roles in development of these viruses, as suggested previously on the basis of biological experiments [22,23]. Such a proposal is corroborated by results presented in this report (Figs. 2–5), indicating more efficient development of both bacteriophages after prophage induction in cells bearing additional copies of the exo-xis region on plasmids, and by the recent finding that Eal8.5, encoded in this region, contains fused homeodomain/zinc-finger fold [24]. The experiments with plasmids bearing particular ORFs or genes, rather than the whole exo-xis region, as well as with plasmids bearing frameshift mutations in particular ORFs or genes, did not exclude a possibility that an excess of phage-derived sequences could cause effective binding of some regulatory factors and titrating them out. This could be responsible for observed effects on prophage development. However, such a scenario seems unlikely as a sole mechanism of this phenomenon, especially in the light of specific effects of particular mutations (Fig. 3). Therefore, we suggest that it is more probable that stimulation of development of λ and Φ24B in cells bearing additional copies of the exo-xis region arises from effects of expression of certain genes, and possible cooperative actions of at least some of their products. In fact, only two genes from this region, ea8.5 and ea22, were confirmed to date to encode proteins, thus others are named as
orfs. However, specific effects of frameshift mutations in orf61, orf73, and orf69a (Fig. 3C and D) suggest that they may code for biologically active polypeptides.

Recent bioinformatics and microarray analyses have indicated a growing number of genes encoding small proteins in the range of 20–130 amino acids [45–47]. Increasing amount of experimental data demonstrates that such small proteins have variety of roles and different mechanisms of action. They can regulate functions of larger proteins, act as signaling factors or structural proteins [47]. Expression patterns of the λ genome from ribosome profiling [44] as well as our results from qRT-PCR analyses show increased expression of ORFs of unknown function during λ phage lytic development (in between these localized in the exo-xis region as shown in Fig. 1 from work [44] and Fig. 7 in this work). Although expression is observed at different times after induction, it could be explained by application of different inducers, temperatures of cultivation and measurement methods. Additionally, in this work we first present the increased expression of ORFs from exo-xis region during lytic development of phage Φ2lp. It is important to note that the level of expression of some of the analyzed ORFs from the exo-xis region is comparable to that of known genes or even higher. As suggested previously [44] such observation allow to suppose that these ORFs might be translated into active polypeptide products. Therefore, our further research will focus on determination of biological and biochemical roles of products of genes included in the exo-xis region, as well as on determination of regulatory mechanisms operating in the process of the controlling of expression of these genes.

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Author Contributions

Conceived and designed the experiments: SB BN-F JML AW GW. Performed the experiments: SB BN-F AD AF. Analyzed the data: SB BN-F JML AF AW GW. Contributed reagents/materials/analysis tools: SB BN-F AW GW. Contributed to the writing of the manuscript: SB BN-F AW GW.

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