A Kayvirus Distant Homolog of Staphylococcal Virulence Determinants and VISA Biomarker is a Phage Lytic Enzyme That Decreases S. aureus Tolerance to Vancomycin

Aleksandra Glowacka-Rutkowska¹, Magdalena Ulatowska¹, Joanna Empel², Magdalena Kowalczyk¹, Jakub Boreczek¹, Małgorzata Łobocka*¹

¹ Institute of Biochemistry and Biophysics of the Polish Academy of Sciences, Warsaw, Poland; glowacka@ibb.waw.pl (A.G.-R.); miulatowska@gmail.com (M.U.); mk@ibb.waw.pl (M.K.); jakub.boreczek@ibb.waw.pl (J.B.); lobocka@ibb.waw.pl (M. Ł.)
² Department of Epidemiology and Clinical Microbiology, National Medicines Institute, Warsaw, Poland; j.empel@nil.gov.pl (J.E.)
* Correspondence: lobocka@ibb.waw.pl; Tel.: +48-22-592-1300

Abstract: Staphylococcal bacteriophages of Kayvirus genus are candidates for therapeutic applications. One of their proteins, Tgl, is slightly similar to staphylococcal virulence factors, secreted autolysins of lytic transglycosylase motifs, IsaA and SceD. We show that Tgl is also a lytic enzyme secreted by bacterial transport system and localizes to cell peripheries, like IsaA and SceD. It caused lysis of E. coli cells expressing the cloned tgl gene, but could be overproduced when depleted of signal peptide. S. aureus cells producing Tgl lysed in the presence of nisin, which mimics the action of phage holin. In vitro, Tgl protein was able to destruct S. aureus cell walls. The production of Tgl decreased S. aureus tolerance to vancomycin, unlike the production of SceD, which is associated with the decreased sensitivity to vancomycin. In the genomes of kayviruses, the tgl gene is located a few genes away from gene lysK, encoding the major endolysin. While lysK is a late phage gene, tgl can be transcribed by a host RNA polymerase, as are phage early genes. Taken together our data indicate that tgl is a part of kayviruses lytic module and encodes an additional endolysin which can act in concert with LysK in cell lysis.

Keywords: Staphylococcus aureus; bacteriophage; Kayvirus; endolysin; virulence determinants; vancomycin

1. Introduction

Staphylococcus aureus is one of the most challenging bacterial pathogens, due to the increasing number and spread of antibiotic resistant strains that are a serious threat to health and life [1]. Thus, bacteriophage therapy may become a future option of choice to overcome the S. aureus antibiotic-resistance problem [2]. As a rule, phages target bacteria of certain strains or species independently of their resistance to antibiotics. They are harmless to eukaryotic cells and propagate in a self-limiting manner, which is controlled by the availability of a sensitive host [3]. Most importantly bacteriophages specific for certain bacterial pathogens do not destruct bacteria of natural human or animal microflora and do not select for antibiotic resistance [4].

A dominant group of staphylococcal phages in the therapeutic phage collections are tailed phages of Kayvirus genus of the Twortvirinae subfamily, family Herelleviridae [5]. They are obligatorily lytic, and infect a broad spectrum of S. aureus strains [6, 7]. Several Kayvirus genus representatives have been successfully used in experimental antistaphylococcal therapies in humans and animals [2, 8, 9, 10, 11, 12, 13, 14]. The genomic sequences of most of them have been determined [6, 15]. They vary between 127-152 kb and do not transfer bacterial DNA by transduction. Core genome regions of kayviruses and other phages of Twortvirinae subfamily are conserved and resemble in organization and coding properties the core genome regions of other phages of Herelleviridae family [5, 6]. Nearly half of about 200 kayviruses genes have been assigned function or putative function based on homologies or, to a lesser extent, on results of experimental studies.
It is commonly accepted that in addition to obligatorily lytic propagation strategy and inability to transfer bacterial DNA by transduction a feature that qualifies phages for the potential use in antibacterial therapies is the lack of genes encoding toxins or other virulence determinants, as well as antibiotic resistance markers [16, 17]. In our previous work we identified in the genomes of Kayvirus genus representatives, a conserved gene (designated by us as tgl) whose product is a distant homolog of staphylococcal virulence determinants, proteins IsaA and SceD [6]. Additional analysis of known or predicted proteins encoded by 22 kayviruses revealed that Tgl is the only gene product of these phages which has slight similarities at the amino acid sequence level to proteins associated with bacterial virulence [17].

IsaA and SceD are paralogous staphylococcal autolysins and surface antigens [18,19]. They are similar in size (231 and 233 amino acid residues, respectively), share 33% of amino acid sequence, and have a lytic transglycosylase domain motif (pfam 06737) in their C-terminal part. They both contribute to cell wall remodeling, are required for normal growth of S. aureus and are associated with S. aureus virulence [18, 19, 20]. Each of them contributes differently to biofilm formation and beta-lactam resistance, and they have opposite effects on S. aureus cells clumping and septation [18, 21, 22, 23, 24].

In the genome of S. aureus the isaA and sceD genes are located in unrelated, monocistronic operons that are far away from each other [see e.g. 25] (GB acc. no. NC_007795.1). They both belong to the essential WalKR two-component system regulon (formerly YycFG) controlling cell wall metabolism and are positively regulated by WalR [26, 27]. Additionally, they are oppositely controlled by two global regulators of virulence genes, SarA and agr [18, 28, 29, 30]. Besides the common factors participating in the control of both genes expression, sceD is positively regulated by sigma B, and by two-component regulatory systems LytSR and SaeRS [30]. While the inactivation of isaA leads to the elevated level of sceD expression, the reciprocal effect was not observed, indicating the overlapping as well as distinct physiological roles of IsaA and SceD [18, 21].

IsaA is a major S. aureus immunodominant antigen, which is surface-exposed and has been identified in exoproteins of all tested S. aureus clinical isolates studied [19, 31, 32, 33, 34]. It is bound to S. aureus cell wall by non-covalent interactions [35]. Diseases associated with S. aureus colonization, including sepsis caused by methicillin resistant S. aureus (MRSA), are associated with the increased IgG response against IsaA [36, 37, 38, 39]. Monoclonal anti-IsaA antibodies were effective in the treatment of staphylococcal infections [40, 41, 42].

SceD is mostly secreted to the growth medium [35]. Its increased production was essential for the establishment of S. aureus nasal colonization in rats [18, 43]. Moreover, SceD is overproduced in most of the MRSA strains with decreased sensitivity to vancomycin, namely VISA (vancomycin-intermediate S. aureus) or hVISA (heterogeneous vancomycin-intermediate S. aureus). Hence it has been proposed to be a potential biomarker for the detection of VISA and hVISA strains [44, 45, 46, 47, 48]. Pieper et al. suggested that its action on the cell wall is required to increase the tolerance to vancomycin [46].

Similarity of Tgl to IsaA and SceD, albeit limited, poses a question about the role of Tgl protein in the development of kayviruses or their interaction with a bacterial host. In a search for the answer to these questions we analyze here the extent and significance of Tgl homologies to IsaA and SceD, and show that Tgl is an additional kayviruses endolysin that may possibly contribute to the high lytic efficacy these phages.

2. Materials and Methods

2.1. Bacterial Strains, Bacteriophages and Growth Conditions of Bacteria

The bacterial strains used in this study were E. coli DH5α (F− λ80lacZAM15 Δ(lacZYA-argF) U169 recA1 endA1 hisD1717(kr−, mcr−) phoA supE44 λ− thi-1 gyrA96 relA1) [49, 50], E. coli BL21 (F−ompT hsdS8 (rB−, mcr−) gal dcm (DE3)) [51] and S. aureus RN4220 - a restriction-deficient derivative of NCTC8325 [52, 53]. Phage MSA6, a representative member of Kayvirus genus phages and a source of tgl gene, was described previously [6, 54]. Bacteria were grown in Luria-Bertani broth (LB; Difco), or in trypticase soy broth (TSB; Difco) with constant shaking (200 rpm) or on agar-solidified LB medium at 30, 37 or 42°C, where indicated. When required, antibiotics were added to the media at the following concentrations: 100 µg/ml ampicillin (E. coli) or 20 µg/ml chloramphenicol (S. aureus). LB solid medium supplemented with 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (x-gal; 200 µg/ml) served to detect the production of β-galactosidase by S. aureus and E. coli cells.
2.2. Plasmids

Shuttle E. coli - S. aureus expression vectors used in this study are based on a hybrid plasmid, pMLE2, derived from the E. coli - S. aureus shuttle pSK5630 vector kindly provided by Ron Skurray [55], and the expression cassette of pDAS122 plasmid kindly provided by David A. Schofield [56], (Figure S1). Briefly, to construct pMLE2, the expression cassette cut out of the pDAS122 plasmid using Styl and BamHI and blunt-ended at StylI end with Klenow fragment of DNA polymerase I was inserted to the SalII and BamHI-digested pSK5630 vector blunt-ended at the SalII end. The cloning site for genes of interest in the pMLE2 vector and its derivatives is preceded by the synthetic promoter (Pro3), whose transcription is repressed by the temperature-sensitive C1 repressor of phage P1 (C1-100) and the P1 Bof (Lxc) protein, which enhances the repression [56]. Genes encoding both regulators are present in pMLE2. At the permissive temperature (30°C), repressor binds to its operator sites and blocks transcription, while at the non permissive temperature (42°C), C1-100 is unstable and transcription is proceeding. The SalII restriction site of pMLE2 was replaced by the PmaCI restriction site, by amplification of the whole pMLE2 sequence with primers OMSL001F and OMSL002R, digestion of the amplicon obtained with PmaCI, and circularization by ligation to make the pMLE3 vector. The hybridization product of pMLE1F and pMLE2R oligonucleotides, was inserted in place of the Spe1-SphI fragment of pMLE3 to generate the pMLE4 plasmid, which contains an additional Clal restriction site between the sequences recognized by PmaCI and SphI and is devoid of the lacZ gene. The hybridization product of two oligonucleotides pMLE3F and pMLE4R, which contained the recognition sites for SacI, AgeI and XbaI restriction endonucleases was inserted in place of the Clal-SphI fragment of pMLE4 to generate the pMLE5 vector. The pMLE5-derived plasmid, encoding Egfp fluorescent protein was constructed by the replacement of AgeI-XbaI fragment of pMLE5 downstream of the Pro3 promoter, with the AgeI-XbaI fragment of the pEGFP plasmid (Clontech; GenBank acc. no. U76561) containing the egfp gene. Constitutive expression vector derivatives of the pMLE3 and pMLE5 were constructed by the introduction of a frameshift mutation into the C1-100 encoding sequence. Briefly, the pMLE3 and pMLE5 vector was digested using BglII. The BglII-generated sticky ends were blunt with Klenow fragment of E. coli DNA polymerase I and religated to generate the pAGL1 and pAGL2 plasmid, respectively. The pMLE5-derived plasmid, pAGL3 containing the tgl gene of MSA6, was constructed by the amplification of tgl, using MSA6 DNA as a template, and primers OAGL40 and OAGL39, digestion of the amplicon with Clal and SphI and insertion of the fragment obtained to the Clal- and SphI-digested pMLE5 plasmid. The pMLE6-derived plasmid pAGL4 containing the translational fusion of full length tgl gene with the egfp gene, was constructed by the amplification of tgl, using MSA6 DNA as a template, and primers OAGL40 and OAGL49, digestion of the amplicon with Clal and AgeI and insertion of the fragment obtained to the Clal and AgeI-digested pMLE6 plasmid. To construct a pMLE3-derived plasmid carrying the promoter-operator region of tgl gene inserted just upstream of the promoterless lacZ gen in pMLE3 (pAGL5), the sequence containing the putative tgl promoter was amplified using bacteriophage MSA6 DNA as a template and primers OAGL57 and OAGL78. The PCR product obtained was digested with NaiI and PmaCI and inserted in place of NaiI-PmaCI fragment of pMLE3. To construct a pMLE5-derived plasmid containing the tgl gene truncated with 28 codons at its 5' end the truncated version of tgl was amplified with MSA6 phage DNA as a template and primers OAGL108 and OAGL109. The amplicon obtained was digested with Clal and SphI and inserted in place of Clal-SphI fragment of pMLE5 plasmid, to generate a plasmid designated as pAGL6. All DNA manipulation procedures were performed according to standard protocols [57] or as recommended by enzyme suppliers. The sequence correctness of all plasmid fragments obtained by PCR was verified by DNA sequencing, with primers OAGL41 and OAGL42. The sequencing was performed at the Laboratory of DNA Sequencing and Oligonucleotides Synthesis of the Institute of Biochemistry and Biophysics, Polish Academy of Sciences. Oligonucleotide sequences used for the DNA amplification or construction of plasmids are listed in Table S1.

2.3. Bioinformatic analysis

Comparative sequence analysis of Tgl protein and gene products of the tgl-flanking regions in the genomes of Kayvirus genus phages, with proteins in databases was performed using Blastp [58], PSI-Blast [59], and Tblastn [58] at NCBI, and HHpred [60]. The N-terminal sequence of the Tgl protein was searched for putative transmembrane domains and signal peptides as described elsewhere [61].
2.4. Testing the influence of intracellular Tgl protein on the growth and survivability of E. coli and S. aureus cells

*E. coli* DH5α or *S. aureus* RN4220 cells, bearing the pAGL3 plasmid with the *tgl* gene under the control of thermoinducible Pro3 promoter were grown overnight in LB supplemented with ampicillin (100 µg/ml, *E. coli*) or chloramphenicol (20 µg/ml, *S. aureus*) at 30°C. Overnight cultures were diluted 1:100 in similar medium (OD₆₀₀ of about 0.001) and incubated with shaking (200 rpm) at 42°C to derepress the expression of the cloned *tgl* gene. Bacterial growth was monitored spectrophotometrically (OD₆₀₀). The number of viable cells in each culture was determined based on the number of colony forming units (CFU/ml) obtained after plating of diluted culture samples and the overnight incubation of plates at 30°C. Cells harboring the empty pMLE5 plasmid were used as a control. Each experiment was performed in triplicate.

2.5. Determination of the intracellular localization of Tgl-EGFP protein in *S. aureus* cells

*S. aureus* RN4220 cells bearing the pAGL6 plasmid, which carries the *tgl*-egfp fusion under the control of thermoinducible promoter, were grown overnight in LB supplemented with chloramphenicol at 30°C. Overnight cultures were diluted 1:100 in similar medium and incubated at 30°C with shaking (200 rpm) until the optical density (OD₆₀₀) of about 0.3. Cultures were then transferred to 42°C to derepress the expression of cloned *tgl* gene. To visualize the Tgl-Egfp protein in individual cells, bacteria were immobilized on an agarose-padded microscope slide. Briefly, 100 µl of molten (75°C) agarose solution (1.5% Agarose SeaKem GTG in 0.7% NCI) was dropped on a prewarmed microscope slide, covered with a cover glass and left to solidify. Next, the cover glass was removed. Cell culture (1 ml) was harvested by centrifugation (10,000 × g for 5 min) and resuspended in 100 µl of 1.5% NaCl. Aliquots of cell suspension (10 µl) were dropped on the agarose and coated with a new cover glass. Cells were viewed with an Eclipse TE2000-E confocal microscope equipped with color camera DS-5Mc (Nikon) and the CFI Plan Apochromat oil immersion objective (100x, numerical aperture = 1.4) with an additional 10x zoom option. A Nikon module for fluorescence equipped with an argon laser (Melles Griot) and BA515-555 Nikon filter set was used to excite the fluorescence of Egfp with blue light (488 nm wavelength) and to observe the Egfp emission light within the wavelength range of 500-530 nm. Confocal microscopy observations were performed at the Laboratory of Confocal and Fluorescence Microscopy of the Institute of Biochemistry and Biophysics of Polish Academy of Sciences, Warsaw, Poland.

2.6. Assays of β-galactosidase activity

Bacteria for the assays of β-galactosidase activity were grown overnight in LB supplemented with ampicillin (*E. coli*) or chloramphenicol (*S. aureus*) at 30°C. Overnight cultures were diluted 1:100 in similar medium, incubated at 30 or 42°C with shaking (200 rpm) until the optical density (OD₆₀₀) of about 0.4, and chilled in ice for 15 min to stop growth. *S. aureus* cells for the assays (0.5 ml) were harvested by centrifugation, resuspended in the same volume of lysis buffer (0.01 M potassium phosphate buffer [pH 7.8], 0.015 M EDTA, 1% Triton X-100, 100 mg of lysostaphin per ml) and incubated at 37°C for 30 min with vigorous shaking. The portions of lysed *S. aureus* cells or *E. coli* cultures were mixed with 0.5 ml of chilled Z buffer (60 mM Na₂HPO₄, 7H₂O; 40 mM NaH₂PO₄, H₂O; 10 mM KCl; 1 mM MgSO₄, 7H₂O; 50 mM β-mercaptoethanol; pH 7.0) to obtain the final volume of 1 ml. Lysis of *E. coli* cells was performed in the reaction mixtures by adding chloroform (50 µl) and 0.1% SDS (25 µl). Next, the probes were vigorously vortexed for 10 s and incubated for 20 min, at 28°C. Reactions were started by adding 0.2 ml of o-nitrophenyl-β-D-galactopyranoside (ONPG, Sigma; freshly prepared solution in H₂O; 4 mg/ml), incubated until the appearance of yellow coloration and stopped by the addition of 1 M Na₂CO₃ (0.5 ml). The enzyme activity was calculated according to Miller's method [62].

2.7. Preparation of *S. aureus* cell walls for zymography

*S. aureus* cells walls were prepared as described previously [63]. Briefly, overnight culture of *S. aureus* was diluted 1:100 in 250 ml of TSB and incubated at 37°C with shaking (200 rpm) until the optical density (OD₆₀₀) of about 1.0. Next, bacteria were harvested by centrifugation (10,000 × g for 15 min, room temperature), washed with 250 ml of Milli-Q water, centrifuged as above and resuspended in 30 ml of Milli-Q water. The cell suspension was autoclaved (15 min, 121°C), centrifuged as above and the pellet obtained was stored overnight at -20°C. Next, the pellet was resuspended in 3 ml of Milli-Q water and...
distributed into pre-weighted tubes. The suspension was dried overnight in a speed vac. The dried pellet was weighted, resuspended in Milli-Q water to reach the final concentration 50 mg/ml and stored at -20°C.

2.8. Overproduction of the TglASP protein and zymography

Overnight culture of *E. coli* BL21 cells harboring the pAGL6 plasmid, which carries the truncated version of tgl gene (tglASP), was diluted 1:100 in 100 ml of fresh LB supplemented with ampicillin and incubated at 30°C with shaking (200 rpm) until the optical density (OD$_{600}$) of about 0.4. Cells harboring pMLE5 plasmid (empty vector) were used as controls. Cultures were then transferred to 42°C to derepress the transcription from the Pro3 promoter. After 5 hours of growth with shaking the cells were harvested by centrifugation (10,000 × g for 15 min), resuspended in 1 ml of lysis buffer (50 mM NaH$_2$PO$_4$, 300 mM NaCl, 10 mM imidazole, 1 mM PMSF, pH 8.0) containing a protease inhibitor cocktail (complete EDTA-free Protease Inhibitor Cocktail; Roche), and disrupted by sonication (Sonication system- Bioruptor UCD-200, Diagenode, Belgium; operating conditions: power setting- high; sonication cycle- 30 sec ON, 30 sec OFF; total sonication time - 10 min). The cell extracts were centrifuged (10,000 × g for 30 min, 4°C) to remove insoluble cell debris. The supernatant obtained was left for further analysis. The pellet was dissolved in 1 ml of lysis buffer with 8 M urea. Next, the 20 µl samples of supernatant or dissolved pellet were mixed with Laemmli buffer (Bio-Rad), heated at 95°C for 5 min and separated electrophoretically in 10% SDS-PAGE gel. Gels were kept in staining solution (InstantBlue, Expedeon) for 1 h at room temperature with gentle shaking, submersed in Milli-Q water and incubated until the protein bands became visible. For zymography, 20 µl portions of the supernatant or dissolved cell pellet were mixed with Laemmli buffer (Bio-Rad) and loaded (without prior boiling) to a 10% SDS-PAGE gel (0.75 mm thickness) containing *S. aureus* cells walls (prepared as above) at a final concentration of 2 mg/ml [63]. Following electrophoresis, the gel was rinsed briefly with Milli-Q water and washed three times with Milli-Q water with gentle agitation for 15 min at room temperature. Then, the gel was incubated overnight at 37°C with gentle agitation in freshly prepared renaturation buffer (50 mM Tris-HCl [pH 7.5], 0.1% [v/v] Triton X-100, 10 mM CaCl$_2$, 10 mM MgCl$_2$). Zymograms were stained in methylene blue solution (0.1% [w/v]) in 0.01% potassium hydroxide) for 1 h and destained in water until the bands formed by transparent gel fragments became clearly visible.

2.9. Preparation of nisin stock solution and nisin activity assay

Brain Heart Infusion growth medium (110 ml; BHI; Oxoid) supplemented with glucose (0.8% w/v) was inoculated with 100 µl of fresh o/n culture of the nisin producing *Lactococcus lactis* IBB 1339 strain from the IBB PAS laboratory culture collection. After incubation at 30°C for 20 h, the culture was centrifuged at 6 800 × g for 15 min to remove cells. Ammonium sulfate (30 g) was added to 100 ml of supernatant, dissolved at room temperature by gentle mixing, the samples were kept in ice for 60 min to allow protein precipitation, and centrifuged at 12 850 × g for 40 min. The pellet containing nisin was dissolved in 1 ml of sterile Milli-Q water, and incubated at 100°C for 2 min to kill contaminating cells. Nisin concentrate was stored at -20°C. The activity of nisin in the concentrate was assayed according to the critical dilution method [64] as previously described [65, 66]. The *L. lactis* subsp. *lactis* IL1403 strain, which is sensitive to nisin, served as an indicator. Briefly, fresh GM17 medium (Oxoid; composition per 1 l: pancreatic digest of casein 5.0 g, soy peptone 5.0 g, beef extract 5.0 g, yeast extract 2.5 g, ascorbic acid 0.5 g, magnesium sulfate 0.25 g, disodium-$\beta$-glycerophosphate 19.0 g) was used to prepare two fold serial dilutions of the nisin concentrate. Aliquots (10 µl) of each dilutions were dropped onto the layer of approximately 10$^7$ indicator cells embedded in GM17 soft agar (0.7% w/v) on the surface of GM17 solid medium in Petro dish. The nisin activity, expressed in arbitrary units (AU) per ml was calculated as follows: the highest dilution factor yielding a clear zone of inhibition on the indicator lawn after 18 h incubation was multiplied with 100 to obtain AU/ml. Each assay was performed in triplicate.

2.10. Testing the influence of Tgl protein production on the sensitivity of S. aureus to vancomycin

The sensitivity of *S. aureus* cells producing Tgl to vancomycin was assayed by the determination of minimal inhibitory concentration (MIC) for vancomycin, and by monitoring the growth of bacteria in cultures supplemented with vancomycin. The MIC for vancomycin was determined in Muller Hinton broth (MHB; Difco) using two-fold dilution plate method, as described previously [67], with some modifications. Briefly, overnight cultures of *S. aureus* cells bearing pAGL3 (tgl+) or pMLE5 plasmid (empty control vector)
were diluted 1:100 in MHB supplemented with chloramphenicol (20 µg/ml) and incubated with shaking for 1 h at 30°C and for additional 1 h at 42°C (to derepress the expression of cloned tgl gene). At zero time 100 µl portions of cell cultures were transferred to wells of prewarmed (42°C) honeycomb plates containing 100 µl MHB with or without vancomycin. MHB medium (200 µl) without antibiotic served as a control for medium sterility. The plates were incubated for 5 h at 42°C. After incubation, 66 µl of dimethyl thiazolyl diphenyltetrazolium bromide (MTT) (0.3 mg/ml) or triphenyl tetrazolium chloride (TTC) (0.1%) per well were added and the plates were incubated for 90 min at 37°C with gentle shaking (110 rpm). The MICs were read manually, based on the changes of natural colors of MTT and TTC diluted in MHB to the colors of their reduced forms.

To monitor the growth of \textit{S. aureus} producing Tgl in the presence of vancomycin, overnight cultures of \textit{S. aureus} RN4220 cells bearing pAGL3 (tgl+) or pMLE5 plasmid (empty control vector) were diluted 1:100 in MHB supplemented with chloramphenicol (20 µg/ml) and incubated with shaking for 1 h at 30°C and for additional 1 h at 42°C (to derepress the expression of cloned tgl gene). At time zero 100 µl portions of the cultures were transferred to wells of prewarmed (42°C) honeycomb plates containing 100 µl MHB supplemented with vancomycin at various concentrations. The plates were incubated in the Bioscreen C Microbiology Plate Reader (Growth Curves USA, New Jersey, USA) for 6 h at 42°C with medium intensity shaking. The optical density of samples (OD\textsubscript{600}) was measured during the whole experiment in 15 min intervals.

3. Results

3.1. Analysis of the amino acid sequence of Tgl protein

Analysis of the 230-aa Tgl protein sequence reveals that it contains a core lysozyme-like domain motif at its C terminal end (cd13925, pos. 165-230, E=3.76e-4) with a conserved glutamate, which is involved in catalytic activity of core-lysozyme like domain-containing proteins, a soluble \textit{E. coli} lytic transglycosylase Slt among them (Figure 1) \cite{68}. Additionally, Tgl contains a predicted signal peptide at its N-terminal end, and the central part of Tgl sequence contains a motif characteristic of essential cell division protein FtsN (PRK10927, pos. 86-143, E=2.47e-03).

Regions of homology between Tgl and staphylococcal autolysins SceD and IsaA span nearly the whole Tgl sequence (Figure 1). One of them (pos. 1-27) overlaps with the sequences of SceD and IsaA preprotein signal peptides. Preproteins of IsaA and SceD are cleaved by staphylococcal type I signal peptidase SpsB, at pos. 29/30 and 27/28, respectively, after the conserved AXA motif essential for the recognition and cleavage \cite{69, 70}. This motif is conserved in Tgl. The remaining regions encompass the Tgl fragment between aa residues 28 and 89, the region of PRK10927 motif and the core lysis-enzyme domain motif. The first of these regions corresponds to the region of IsaA essential for the binding to cell wall \cite{35}. The second region, homologous to the PRK10927 motif in Tgl is the least conserved. Its significance is unclear. No homolog of \textit{E. coli} FtsN protein was described in staphylococci. Additionally, the similarities between Tgl and PRK10927 motif do not include amino acid residues of FtsN essential for its functions or protein-protein interactions (data not shown). The corresponding IsaA fragment cannot bind to the cell wall by itself but binds the cell wall, when associated with the preceding fragment, indicating that it may facilitate the binding \cite{35}. 
Figure 1. Alignment of the amino acid sequences of MSA6 Tgl and its homologs encoded by type phages of Baoshanvirus, Twortivirus and Sepunavirus genus [5, 112, 113] with S. aureus SceD and IsaA. The Clustal Omega program (CLUSTAL O [1.2.4]) at its site (https://www.ebi.ac.uk/Tools/msa/clustalo/ [71]) served to make the alignment. Protein designations are the following: Tgl - AFN38714, phage phiSA_BS2 - phiSABS2_87 - AVR55531, phage phiIBB-SEP1 gp165 - YP_009601090, phage Twort ORF051 - YP_238708.1, IsaA - WP_130826635, SceD - WP_000752008. Signal peptides of IsaA and SceD preproteins are bolded, according to [18, 69]. AXA sequences of known or predicted recognition sites for type I signal peptide are underlined. Regions of known or predicted core lysozyme-like domains are indicated by the line above the alignment, those of predicted PRK motif by the dotted line above the alignment. Amino acid residues that are identical in all proteins are marked with an asterisk, those of similar properties are marked with double or single dots. The conserved glutamate known as essential for catalytic activity of lytic proteins with core lysozyme-like domains is highlighted in grey. The region of IsaA sequence essential for the binding to staphylococcal cell wall is highlighted in grey.

In addition to S. aureus IsaA and SceD and their homologs in S. aureus-related bacterial species, proteins that are similar to Tgl over its entire length are encoded only by S. aureus phages of Kayvirinae, Twortvirinae, Sepunivirus [5], and Baoshanvirus [112] genus of Twortvirinae subfamily, as verified by Tblastn [58]. Similarities between Tgl and its closest homologs in these phages are: 100/99-100; 89/44; 82-89/33-82; 56/33 as indicated by %coverage/%identity. Twortvirinae of Silviavirus [5] and Sciuriunavirus [113] genus encode proteins of short regions slightly similar to the core-lysozyme like domain-containing region of Tgl (19-30 coverage/43-44 identity) but they do not contain signal sequences, are products of...
genes encoded by structural genome modules and are annotated as tail-tip proteins or tail lysins (data not shown).

SceD and IsaA have been classified to lytic transglycosylases based on the presence in their C-terminal parts a motif characteristic of lytic transglycosylase domain (pfam06737) [18]. This motif overlaps with the motif of core lysozyme-like domain of resuscitation-promoting factor proteins, which is common for IsaA, SceD and Tgl, and in Tgl is in the region between aa 165 and 207 amino acid residues (3.66e-03), and with the motif of lysozyme-like domains (ci00222; 3.66e-03 in Tgl, 2.60e-14 in SceD and 1.83e-05 in IsaA). The latter is common for a large family of enzymes involved in the cleavage of beta-1,4-linked polysaccharides, including soluble lytic transglycosylases (SLT), goose egg-white lysozyme (GEWL) and bacteriophage lambda endolysin. When searched with HHpred against protein structure database, the Tgl region encompassing the motif of lysozyme-like domains appeared to be significantly similar to active enzyme domains of Pseudomonas aeruginosa MltF (4P0G; 1.1e-8; 5A5X_B, 4.8e-8), Escherichia coli MltC (4CFP_B, 4.7e-8), Ralstonia sp. GH23 family chitinase A-471, Neisseria meningitidis LtgA (5O29_A, 9.1e-8), E. coli Slt70 (1QSA_A, 1.1e-7), henn egg-white lysozyme (2VB1_A, 5.2e-8), P. aeruginosa Slt (5OHU_A, 3.1e-7), Campylobacter jejuni Cj0943 (6CF8_A, 1.5e-7), goose-type lysozyme (4G9S_A, 9.2e-7), E. coli EtgA (4XP8_A, 1.1e-6) and E. coli MltE (6GL4_B, 2.1e-6). All of the aforementioned enzymes, but lysozymes and Ralstonia sp. GH23 family chitinase, which has an atypical structure [72] belong to LTs of Gram-negative bacteria with LT domains classified to family 1 [73, 74]. All these LTs as well as SceD, IsaA and phage homologs of Tgl have been classified as LTs of GH23 family of glycoside hydrolyses according to CAZy (Carbohydrate Active enZYmes) database classification (http://www.cazy.org/) [75]. Consistently, although in general the predicted secondary structure and the region of catalytic glutamic acid residue (at pos. 165 of Tgl sequence) aligns well with the corresponding regions of cod, goose and swan lysozyme, the region containing two aspartic acid residues (at pos. 90 and 101 of cod lysozyme) critical for the catalytic activity of lysozymes and absent in LTs [76, 77] is not conserved in Tgl (data not shown).

3.2. Localization of Tgl protein in S. aureus

To test whether the homology of Tgl to SceD and IsaA in the region of SceD and IsaA signal peptide sequences reflects the ability of Tgl to be transported to or through the cytoplasmic membrane, we constructed a translational fusion of tgl gene to the gene encoding a fluorescent protein, EGFP. In the S. aureus cells producing Tgl-EGFP fusion protein from the resultant plasmid (pAGL4), the concentration of fluorescence was detected in cell peripheries, indicating the functionality of the predicted Tgl signal peptide in the transport of Tgl (Figure 2).
Figure 2. Localization of Tgl-EGFP protein in S. aureus cells. (A) Schematic picture of tgl-egfp expression cassette in pAGL4 plasmid. (B) Confocal microscope images (left panel) and differential interference contrast (DIC) images (right panel) of S. aureus cells producing Tgl-EGFP protein. The central panel shows the merge of both images.

3.3 Influence of Tgl on the growth and survivability of E. coli and S. aureus

Attempts to clone the tgl gene in E. coli in a shuttle E. coli-S. aureus vector, pAGL2, under the control of constitutive promoter were unsuccessful. Thus, to analyze the tgl function we cloned tgl in a shuttle pMLE5 vector under the control of thermoinducible promoter Pro3 (Figure 3A, Figure S1).
Figure 3. Influence of the intracellular Tgl protein on the growth and survivability of *E. coli* DH5α and *S. aureus* RN4220 cells. (A) Schematic picture of *tgl* expression cassette in the pAGL3 plasmid. (B,C) Changes in the optical density (dotted lines) and the number of colony forming units (solid lines) in cultures of *E. coli* DH5α (B) and *S. aureus* RN4220 (C) cells harboring the pAGL3 or pMLE5 plasmid (empty control vector). At zero time the diluted overnight culture of each strain (1:100; OD$_{600}$ of about 0.001) was transferred to 42°C to derepress the transcription from the Pro3 promoter. The left Y-axis at each graph represents the log of CFU/ml, the right Y-axis represents OD$_{600}$. Each curve shows the representative results of one of three independent experiments.

Induction of the *tgl* transcription in *E. coli* cells carrying the resultant plasmid, pAGL3, caused cell lysis, as indicated by a decrease in the optical density of cell culture as well as by a decrease in the number of viable cells (CFU/ml, Figure 3B). Clearly, Tgl can cause the lysis of *E. coli* cells from within, indicating that it has a mурalytic activity and can pass the cytoplasmic membrane to get access to the *E. coli* cell wall. The induction of *tgl* expression from the same plasmid in *S. aureus* cells also caused a decrease in the number of viable cells but to a much lesser extent and with a long delay as compared to the decrease observed in the case of *E. coli* (Figure 3C). Moreover, the optical density of *S. aureus* cells expressing *tgl* grew much slower than that of cells with the empty vector, but no decrease of optical density was observed even in 10-th hour upon the induction of *tgl* expression. The pMLE5 plasmid whose backbone is derived from the shuttle pSK5630 vector (Figure S1) is driven by a high-copy-number ColE1 plasmid replicon in *E. coli* cells, but by the low-copy-number replicon of pSK1 plasmid in *S. aureus* cells [55]. Despite the copy-number differences the retention of pSK5630 backbone-based plasmids in *E. coli* and in *S. aureus* is similar (Figure S2). However, we observed that the level of β-galactosidase measured in *S. aureus* and *E. coli* cells harboring the pMLE3 plasmid, parental for pMLE5 and containing the *lacZ* gene under the control of Pro3 promoter, is nearly three times higher in *E. coli* than in *S. aureus* (Figure S3). Thus, the observed differences between the response of *E. coli* and *S. aureus* to the induction of *tgl* expression may be
due, at least in part, by the differences between the level of \( tgl \) transcription in \( E. coli \) and \( S. aureus \) carrying pAGL3.

3.4. Muralytic activity of Tgl protein against \( S. aureus \) cell walls

To overproduce the Tgl protein in \( E. coli \) cells we attempted to eliminate the problem of Tgl lethality for \( E. coli \), by creating a version of Tgl (TglΔSP) depleted of the N-terminal 28 amino acid residues comprising the signal peptide (Figure 4A). \( E. coli \) cells expressing the recombinant \( tgl \) gene from the pAGL6 plasmid, in contrast to control cells with an empty vector, produced a protein of predicted molecular mass of TglΔSP (24 kDa), indicating that it is TglΔSP (Figure S4). The majority of the TglΔSP protein was in the cell pellet.

![Figure 4. \( S. aureus \) cell degrading activity of the Tgl protein. (A) Schematic picture of the expression cassette of pAGL6 plasmid encoding the recombinant version of Tgl (TglΔSP) protein deprived of the signal peptide. (B) Zymograms obtained upon separation of pellet proteins of \( E. coli \) BL21 cells harboring plasmid pAGL6 (Ba) or pMLE5 (empty control vector) (Bb), by SDS-PAGE in a gel containing dead \( S. aureus \) cells. The band of clearing was detected upon protein renaturation in the gel (see Materials and Methods). It is indicated by the arrowhead.]

To test whether the TglΔSP can act as \( S. aureus \) cell wall degrading protein, extracts from crude preparations of induced \( E. coli \) BL21 strain cells, harboring plasmid pAGL6 were subjected to denaturing polyacrylamide gel electrophoresis in gels containing \( S. aureus \) cell walls. Protein extracts of cells harboring pMLE5 plasmid (empty vector) served as a negative control in this experiment. After the renaturation of proteins in the gel and gel staining with Coomassie blue a clear lysis zone on a blue background was observed in the region of predicted TglΔSP localization, in lanes containing pellet proteins of the lysed \( E. coli \) BL21/pAGL6 cells. No similar zone was present in lanes with control (Figures 4B and Figure S5). Clearly, the Tgl protein can act as an \( S. aureus \) cell wall hydrolase. Additionally, our results indicate that the N-terminal 28-aa region of Tgl is dispensable for the ability of Tgl to degrade the \( S. aureus \) cell wall.

3.5. The influence of nisin on the growth and survivability of Tgl producing \( S. aureus \) cells

The hydrolytic activity of TglΔSP against \( S. aureus \) cell wall suggested that the intact Tgl protein can function as an additional endolysin of Twortvirus genus phages that encode it. To verify, whether Tgl can lyse \( S. aureus \) cells from within, we supplemented a culture of \( S. aureus \) producing Tgl, with nisin. Nisin is a membrane-permeabilizing bacteriocin produced by certain \( Lactococcus \) strains. It forms pores in a bacterial cell membrane and dissipates the proton motif force (pmf) \([78]\), thus mimicking the action of phage holins. While nisin at the concentration used did not inhibit the growth and did not decrease the survivability of \( S. aureus \) cells with pMLE5 plasmid (empty vector), it caused the complete cessation of increase in the optical density of cells carrying the \( tgl \) encoding plasmid and a decrease in the number of viable cells in their culture (Figure 5). Taken together our results indicate that Tgl can function as an endolysin in \( S. aureus \) cells, but requires the help of pore-forming protein, to get access to \( S. aureus \) cell wall. In cells infected with a Tgl-encoding Kayvirus genus phage this help is likely to be provided by a phage-encoded holin.
Figure 5. Effect of intracellular Tgl protein on growth (A) and survivability (B) of nisin-treated S. aureus cells. Overnight cultures of S. aureus cells bearing pAGL3 (tgl+) or pMLE5 plasmid (empty control vector) were diluted 1:100 in LB and incubated at 30°C until the optical density (OD_600) of about 0.1. At time zero cultures were transferred to 42°C, supplemented with nisin concentrate (50 µl per 25 ml S. aureus culture) and incubated further. Parallel samples without nisin served as controls. Each curve shows the representative result of one of three independent experiments.

3.6. Analysis of the transcriptional activity of the putative early tgl promoter

In the genomes of Twortvirinae that encode Tgl protein, the tgl gene is preceded by a sequence that is similar to promoters for housekeeping RNA polymerase of S. aureus cells (Figure S6) [6, 79].

Figure 6. Transcriptional activity in E. coli and S. aureus of the region preceding the MSA6 tgl gene. (A) Schematic picture of transcriptional fusion in the pAGL5 plasmid. The coordinates, in the genome of MSA6,
of promoter region cloned are: 32540 – 33289 bp. (B) Colonies formed on LB solid medium with x-gal at 30°C by *E. coli* DH5α and *S. aureus* RN4220 cells that were transformed with pAGL5 or pAGL7 plasmid (control). (C) The activity of β-galactosidase in cells of *E. coli* DH5α and *S. aureus* RN4220 bearing the pAGL5 or pAGL7 plasmid. The activity of β-galactosidase in *S. aureus* cells with plasmids was normalized by the subtraction of background that has been observed in *S. aureus* RN4220 and results from yellow coloration of 2-aminophenoxazin-3-one that is produced by the majority of *S. aureus* strains [114]. Each value is the average of results of at least three independent assays; vertical bars show average deviations.

To verify whether this sequence can function as an early promoter of *tgl*, we inserted it upstream of the promoterless reporter gene, *lacZ*, in the pAGL7 plasmid (Figure 6A). *E. coli* as well as *S. aureus* cells with the resultant plasmid (pAGL5) formed blue colonies on the solid LB medium supplemented with x-gal, indicating that the inserted DNA fragment can drive the transcription of reporter gene in both these bacterial species in the absence of any phage proteins (Figure 6B). Consistently, the activity of β-galactosidase in cells of *E. coli* and *S. aureus* containing the pAGL5 plasmid reached a few hundred Miller Units, while in cells with the empty vector no β-galactosidase activity was detected (Figure 6C). Clearly, the *tgl* gene can be transcribed from an early phage promoter, which is dependent on the host RNA polymerase with a housekeeping sigma factor. Although the activity of β-galactosidase in *E. coli* cells with pAGL5 appeared to be twice as high as in *S. aureus* cells with this plasmid, the difference between the copy number of pAGL5 in *E. coli* as compared to *S. aureus* cells is likely to contribute to this difference.

3.7. Susceptibility of Tgl-producing *S. aureus* to vancomycin

The increased production of *S. aureus* SceD protein was shown to correlate with hVISA or VISA phenotype [48, 80, 81] which in turn is associated with vancomycin treatment failure [82, 83]. Thus, Cui et al. [17] speculated that the expression of staphylococcal phages homologs of SceD during *S. aureus* infection may increase the minimum inhibitory concentration (MIC) of vancomycin required to treat infections by MRSA. To verify this possibility we compared the minimal inhibitory concentrations of vancomycin required to inhibit growth of *S. aureus* RN4220 expressing the *tgl* gene from a plasmid with that of RN4220 carrying an empty vector. We did not observe any decrease in the susceptibility of *S. aureus* producing Tgl to vancomycin, as compared to the control strain (Figure 7). To the contrary, we reproducibly
Figure 7. Influence of intracellular Tgl protein on the susceptibility of S. aureus to vancomycin. (A) Growth of Tgl producing S. aureus RN4220 cells in the presence of vancomycin at various concentrations. Overnight cultures of S. aureus cells bearing pAGL3 (tgl+) or pMLE5 plasmid (empty control vector) were diluted 1:100 in MHB supplemented with chloramphenicol (20 µg/ml) and incubated with shaking for 1 h at 30°C and for additional 1 h at 42°C (to derepress the expression of cloned tgl gene). At time zero 100 µl portions of the cultures were transferred to wells of prewarmed (42°C) honeycomb plates containing 100 µl MHB supplemented with vancomycin at concentrations indicated. The plates were incubated in the Bioscreen C Microbiology Plate Reader (Growth Curves USA, New Jersey, USA) for 6 h at 42°C with medium intensity shaking. The optical density of samples (OD<sub>600</sub>) was measured during the whole experiment in 15 min intervals. Each curve shows the representative result of one of three independent experiments. (B) Determination of MICs for vancomycin for S. aureus cells producing Tgl protein. MICs for vancomycin were determined in MHB using two-fold dilution plate method, as described previously [67], with some modifications. Briefly, 100 µl portions of RN4220 cell cultures containing pAGL3 or pMLE5 (as indicated) and prepared like in (A) were transferred to wells (columns A-K) of prewarmed (42°C) honeycomb plates containing 100 µl MHB with (column A-J) or without vancomycin (column K). Column L contained 200 µl of MHB without antibiotic as a control for medium sterility. Plates were incubated for 5 h at 42°C. After incubation, 66 µl of MTT (0.3 mg/ml) or TTC (0.1%) per well were added (as indicated) and the plates were incubated for 90 min at 37°C with gentle shaking (110 rpm). The MICs were read manually. Comparison between columns L and K shows a change of natural color of MTT and TTC diluted in MHB (respectively yellow and colorless - column L) to the reduced form (respectively violet and red - column K).

observed a slightly increased susceptibility to vancomycin of S. aureus producing Tgl as compared to that with an empty vector (MIC change from 4 to 2), indicating that Tgl can decrease the S. aureus tolerance to vancomycin.

4. Discussion
Phage therapy is a promising alternative to antibiotic therapy in curing infections caused by the drug resistant *S. aureus* strains [84]. However, each potential therapeutic phage should be well characterized at the structural and functional level, before its introduction to a wide use, to ensure the safety of therapy [16, 17]. Although representatives of *Twortvirinae* subfamily phages that proved to be effective in curing infections with *S. aureus*, have been characterized at the level of genomic sequence, the function of over half of their genes is unknown [6, 17, 85, 86]. Of special interest is the *tgl* gene, whose predicted product is slightly homologous to major surface antigens and virulence factors of *S. aureus* cells, SceD and IsaA [6]. Both of them act as autolysins, and have been classified to lytic transglycosylases based on the presence in their C-terminal parts a motif characteristic of lytic transglycosylase domain (pfam06737) [18]. Here we show, that homologies of Tgl with SceD and IsaA, which span the region of lytic transglycosylase motif at protein C-terminal domains and overlap the region of pre-SceD and pre-IsaA signal peptide sequences correlate with functional similarities. Like SceD and IsaA [18], Tgl protein can digest peptidoglycan of *S. aureus* cell wall. Like SceD and IsaA [18, 87], Tgl can be transported through the cytoplasmic membrane by a bacterial transport system, as indicated by the localization of Tgl in the *S. aureus* cell peripheries and by its ability to cause lysis of *E. coli* cells from within. High homologies of the predicted Tgl signal peptide to the signal peptides of SceD and IsaA, including the conserved AXA motif essential for the cleavage by type I signal peptidases (SPases), suggests that Tgl can be also translocated through the cytoplasmic membrane by these SPases with the removal of signal peptide. Consistently, we show that the N-terminal fragment of Tgl, corresponding to the predicted signal peptide is dispensable for the muralytic activity of Tgl.

The physiological significance of PRK10927 motif, characteristic for FtsN proteins, in the amino acid sequences of Tgl is unclear. *S. aureus* does not encode any obvious homolog of FtsN or proteins that directly interact with FtsN in Gram-negative bacteria. Additionally, the amino acid residues that are essential for the function of FtsN are absent from Tgl, SceD and IsaA (data not shown). Possibly, this region can facilitate the interaction of Tgl with cell wall as it has been proposed for the corresponding region of IsaA [35].

In the genomes of staphylococcal kayviruses the *tgl* gene is located in the cluster of leftward transcribed genes, three genes downstream of *lysK*, which encodes the major endolysin of these phages (Figure S6 and 8) [86]. LysK protein contains two domains associated with its muralytic activity: amidase-2 family domain (N-acetylmuramoyl-L-alanine amidase) and CHAP domain (cysteine, histidine-dependent amidohydrolase/peptidase) [88]. They cleave the *S. aureus* peptidoglycan at the amide bond between the N-acetylmuramic acid (MurNAc) of glycan moiety and the L-alanine of peptide side chain, and at the peptide bond between D-alanine and the first glycine of the pentaglycine cross-bridge, respectively [89]. As a canonical endolysin, which relays on the function of pore-forming holin to get access to a cell wall, LysK does not contain any signal peptide or transmembrane domain, and its gene is immediately preceded by a holin encoding gene (*holA*). Based on the close neighborhood of *tgl* gene to *lysK* and *hol*, and on the confirmed mureinolytic activity of Tgl protein, we postulate that *tgl* is an additional component of kayviruses lytic module, and encodes a second endolysin of these phages. The conserved structural and amino acid sequence motifs of family 1 lytic transglycosylases in the Tgl protein indicate that Tgl cleaves the glycan component of cell wall peptidoglycan, and that the cleavage occurs on the reducing side of N-acetylmuramic acid. In this respect Tgl resembles the lambda phage endolysin (product of gene R) and the *Pseudomonas* phiKZ phage endolysin, Gp144, which are also lytic transglycosylases [73, 90, 91 and references therein].

Several endolysins containing two domains of muralytic activity have been identified experimentally or by the bioinformatic analysis of predicted products of phage genes [reviewed by 61; 92 and references therein]. However, fewer examples of phages are known that encode two endolysins targeting different bonds in peptidoglycan. In general, combinations of lysins of different cleavage specificity as well as lysins of multiple muralytic domains of different cleavage specificity perform better in the disruption of bacterial cell walls than enzymes of single muralytic domain when acting alone [reviewed by 92]. Consistently, a cocktail of LysK and lysostaphin as well as engineered LysK enriched with the glycyl-glycine M23 endopeptidase domain of lysostaphin outperformed either of these combination components in the ability to lyse *S. aureus* [93, 94]. Two endolysins of different cleavage specificity may also extend the phage host range, as it was shown for *Bacillus thuringiensis* phage GIL01 [95]. The production by kayviruses of two different endolysins, LysK and Tgl, targeting three kinds of bonds in peptidoglycan may be a feature of these phages contributing to their high lytic efficacy and wide host range.
Proteins of phage lytic modules fulfill their function at the last step of the phage development. However, typically, the expression of endolysin genes and, in certain cases, the secretion of endolysins to the cell wall environment starts at an early stage of infection and reaches its maximum at late stages [96 and references therein, 97]. The tgl gene is preceded by a sequence that is similar to promoters for host RNA polymerase with a housekeeping sigma factor [6, 79]. Indeed, we show here, that it is transcribed by a host RNA polymerase in the absence of any phage proteins – a hallmark of early phage genes. Meanwhile, the transcripts of lysK were detected in phage infected cells between 10 and 20 min after phage infection, as shown in studies with type kayvirus K [unpublished data referred to in 86]. Consistently, neither lysK nor holA genes are preceded by any sequence resembling promoters for bacterial RNA polymerase holoenzyme [6]. We conclude that the transcription of tgl gene is at least in part independent of the transcription of lysK and starts early in phage development.

Despite the early production of Tgl protein and its likely host SpsB-dependent transport through the cytoplasmic membrane, kayvirus-infected cells do not lyse prematurely. This is consistent with the proposed dependence of endolysin-mediated cell lysis on prior dissipation of membrane proton motif force (pmf) and cell death, which is normally triggered by holins and was shown to be a holin function additional to the formation of membrane pores [98]. According to this lysis model even an endolysin accumulated in the extracytoplasmic environment acts ineffectively until holin-mediated dissipation of the pmf at a defined time can cause cell death and prevent the repair of endolysin-mediated lesions by cellular proteins. In support of that, the genomes of phages predicted to encode only secretory endolysins harbor a holin gene [97, 99]. Our results fully conform to this model. Although S. aureus cells producing Tgl from the cloned tgl gene stopped growing, they did not lyse unless their culture was supplemented with nisin - a peptide antibiotic, which forms pores in the cytoplasmic membrane and dissipates the pmf, thus mimicking the function of holins. Similar phenomena were observed in the case of the other secreted endolysins: Lys44 of Oenococcus oeni bacteriophage fOg44 which contains SP, and LysPP1 of Bacillus subtilis which was artificially enriched with SP [97, 98, 99]. Exponentially growing cells of Gram-positive hosts (O. oeni or L. lactis, and B. subtilis, respectively) producing either of these endolysins from a cloned gene or treated with externally added endolysin at physiological concentrations, lysed only if their culture was supplemented with nisin or exposed to other factors or conditions dissipating the membrane pmf.

The dependence of intracellular Tgl on nisin in the lysis of S. aureus cells that was observed in this work contrasts with the ability of Tgl to cause rapid lysis of E. coli cells in the absence of nisin. This difference is likely to result from a lower level of tgl gene expression in S. aureus as compared to E. coli in our experimental system and from the requirement of more enzyme for the disruption of S. aureus peptidoglycan, which is much thicker than that of E. coli. In the case of SP1 bacteriophage endolysin enriched with SP, the requirement of nisin for cell lysis also depended on the concentration of endolysin [100]. While at physiological endolysin concentration lysis required nisin, it could occur in the absence of nisin when the endolysin concentration was several times higher.

The tgl, lysK and holA genes are in the region encompassing 8 genes between the functionally different genome modules of which one encodes tRNAs and the other ribonuclease and structural virion components (Figure 8) [6]. Of three genes between lysK and tgl, two, mbpV and mbpS, encode predicted membrane proteins. MbpV is a 102-aa protein with a transmembrane domain in its N-terminal part and numerous positively charged amino acid residues in its C-terminal moiety (data not shown). In this respect it resembles holins. The amino acid sequence of MbpS contains a conserved SPFH domain motif (PF01145) characteristic for stomatin, prohibitin, flotillin, and HflK/C proteins [101]. Additionally, the predicted secondary structure of MbpS in the region between amino acid residues 57 and 244 is highly similar to that of archael stomatin of Pyrococcus horikoshi (3BK6_C; 4e-19) as indicated by results of HHpred analysis. In bacterial cells stomatins, flotillins and related proteins are parts of so called functional membrane microdomains (FMM), which are similar to the lipid rafts of eukaryotic cells [102, 103, 104]. FMMs allow for the compartmentalization of different processes within the membrane despite their close proximity [103]. The main component of lipid rafts is flotillin, a scaffold protein which recruits other proteins and stabilizes their multimeric complexes [102]. One of the most represented FMM-associated protein is SecA, as it was shown for S. aureus, B. subtilis and Borrelia burgdorferi [103]. The interaction of Sec secretion machinery with flotillins is important for the Sec-associated protease and protein secretion activities, as it was shown in the case of B. subtilis [103, 105]. One cannot exclude that in Staphylococcus cells infected with a Kayvirus, MbpS protein facilitates the SpsB(Sec)-mediated transport of Tgl through the membrane.
The organization of tgl and lysK gene region appears to be similar to that in MSA6 in all Kayvirus genus phages genomes (Figure 8). In the genomes of Twortvirus, Baoshanvirus and Sepunavirus genus phages, which also encode a homolog of Tgl, this organization is slightly different indicating genomic rearrangements. Genomes of Silviavirus and Sciuriunavirus genus do not encode a homolog of Tgl. Conceivably the tgl gene was acquired by phages of certain Twortvirinae genera from their bacterial host after their separation from a common ancestor, and evolved to serve bactericidal functions. Production of two endolysins targeting different bonds in peptidoglycan may be advantageous for Twortvirinae. Lytic modules of Silviavirus and Sciuriunavirus genus phages, which do not encode a homolog of Tgl protein, also encode two proteins of predicted muralytic activity [79, 85, 106, 107].

The homology of Tgl protein to S. aureus autolysins, SceD and IsaA, has been a major concern. Cui et al., [17] wandered whether the S. aureus cells infected with a Kayvirus genus phage and producing Tgl could be less sensitive to vancomycin than non-infected cells. We do not confirm such possibility and demonstrate that S. aureus cells producing Tgl are less tolerant to vancomycin than their Tgl-free counterparts. The amino acid sequence of Tgl is only in 36% identical to that of SceD over 82% of the entire protein length. Conceivably, the influence of Tgl on S. aureus cells is limited to mureinolysis.

Certain natural or engineered phage endolysins of amidase or endopeptidases activity were shown to act synergistically with vancomycin to kill S. aureus cells [108, 109, 110, 111]. The influence of endolysins of lytic transglycosylase activity on the sensitivity to vancomycin has not been studied previously. Our results show for the first time that endolysins of this activity can also act as allies in the fight with pathogens of reduced vancomycin sensitivity.

5. Conclusions

Enzymes of phage and bacterial origin, responsible for the cleavage of the same kinds of bonds in peptidoglycan, have enzymatic domains of similar sequence or structure, indicating the past transfer of their ancestral genes between bacteria and phages. However, further fates of these genes in evolution are tightly associated with the needs of genomes that carry them. Staphylococcal phages of certain Twortvirinae genera, kayviruses among them, encode a distant homolog (Tgl) of two secreted autolysins and virulence factors of S. aureus (IsaA and SceD), which was apparently acquired by them after the differentiation of Twortvirinae. Here we show, that like IsaA and SceD, Tgl has features of lytic transglycosylase and can disrupt S. aureus and E. coli cell wall. Like IsaA and SceD it is secreted through the cytoplasmic membrane.
with the help of its signal peptide and host signal peptidases of type I. However, the location of tgl gene in a genome lytic module of phages that have this gene indicates that Tgl protein functions as a phage endolysin. Like other secreted endolysins Tgl, when not overproduced, requires a dissipation of pmf to cause cell lysis. Early expression of tgl gene in phage development, and lytic transglycosylase properties of Tgl protein imply that Tgl can support the function of major kayviral endolysin, LysK, which is a late phase protein and has the amidase and cysteine, histidine-dependent amidohydrolase/peptidase activity. Possibly, the combined ability of Tgl and LysK to target three kinds of bonds in cell wall peptidoglycan contributes to the wide host range and high lytic efficacy of kayviruses and related Twortvirinae. Most importantly we show here, that the production of Tgl by S. aureus, unlike the production of SceD, is not associated with the decreased sensitivity to vancomycin but causes the decrease in tolerance to vancomycin.

Supplementary Materials: The following are available online at www.mdpi.com/xxx/s1, Supplementary experimental procedure: Plasmid retention test; Table S1: Oligonucleotides used in this study; Figure S1: Organization of genes and regulatory sites in the S. aureus - E. coli shuttle expression vector, pMLE2, and its derivatives; Figure S2: Retention of the pSK5630, pDAS122, pMLE3 or pAGL1 plasmid in E. coli DH5α and S. aureus RN4220 cells grown in the absence of selection for the resident plasmids; Figure S3: The activity of β-galactosidase in E. coli DH5α and S. aureus RN4220 cells containing pDAS122, pMLE3 or pAGL1 plasmid; Figure S4: Overproduction of the recombinant version of the Tgl protein (TglΔSP) deprived of the N-terminal fragment corresponding to the signal peptide; Figure S5: S. aureus cell wall destructing activity of the TglΔSP protein (full image); Figure S6: Genetic organization of the tgl gene region of Kayvirus genome [based on Lobocka 6].

Author Contributions: Conceptualization, Aleksandra Glowacka-Rutkowska and Malgorzata Lobocka; Formal analysis, Malgorzata Lobocka; Funding acquisition, Aleksandra Glowacka-Rutkowska and Malgorzata Lobocka; Investigation, Aleksandra Glowacka-Rutkowska, Magdalena Ulatowska and Malgorzata Lobocka; Methodology, Aleksandra Glowacka-Rutkowska, Magdalena Ulatowska, Joanna Empel and Malgorzata Lobocka; Resources, Magdalena Kowalczyk and Jakub Boreczek; Supervision, Malgorzata Lobocka; Validation, Aleksandra Glowacka-Rutkowska and Malgorzata Lobocka; Visualization, Aleksandra Glowacka-Rutkowska and Magdalena Ulatowska; Writing – original draft, Aleksandra Glowacka-Rutkowska and Malgorzata Lobocka; Writing – review & editing, Aleksandra Glowacka-Rutkowska and Malgorzata Lobocka.

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