Novel Giant Siphovirus from Bacillus anthracis Features Unusual Genome Characteristics

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Abstract

Here we present vB_BanS-Tsamsa, a novel temperate phage isolated from Bacillus anthracis, the agent responsible for anthrax infections in wildlife, livestock and humans. Tsamsa phage is a giant siphovirus (order Caudovirales), featuring a long, flexible and non-contractile tail of 440 nm (not including baseplate structure) and an isometric head of 82 nm in diameter. We induced Tsamsa phage in samples from two different carcass sites in Etosha National Park, Namibia. The Tsamsa phage genome is the largest sequenced Bacillus siphovirus, containing 168,876 bp and 272 ORFs. The genome features an integrase/recombinase enzyme, indicative of a temperate lifestyle. Among bacterial strains tested, the phage infected only certain members of the Bacillus cereus sensu lato group (B. anthracis, B. cereus and B. thuringiensis) and exhibited moderate specificity for B. anthracis. Tsamsa lysed seven out of 25 B. cereus strains, two out of five B. thuringiensis strains and six out of seven B. anthracis strains tested. It did not lyse B. anthracis PAK-1, an atypical strain that is also resistant to both gamma phage and cherry phage. The Tsamsa endolysin features a broader lytic spectrum than the phage host range, indicating possible use of the enzyme in Bacillus biocontrol.

Introduction

We present a novel temperate phage isolated from Bacillus anthracis, the causative agent of anthrax infections in wildlife, livestock and humans [1]. Bacillus anthracis is a member of the Bacillus cereus sensu lato group of six closely related species: B. cereus, B. anthracis, B. thuringiensis, B. mycoides, B. pseudomycoideae and B. weihenstephanensis [2]. This group contains both obligate and opportunistic animal pathogens, including B. anthracis, B. cereus, strains of which cause food poisoning and opportunistic infections in humans [3], and B. thuringiensis, an insect pathogen commonly used as a pesticide in agriculture [4]. Genomic studies have identified a number of putative prophages in the Bacillus cereus sensu lato group (e.g., [5], [6]), including four unique prophage elements in B. anthracis [7].

Lysogeny occurs commonly in B. anthracis [8] and may play an essential role in its life cycle [9]. Isolates of B. anthracis from soil frequently exhibit phage-derived plaques upon subculture [10]. Schuch et al. [9] showed that temperate phage infections of B. anthracis can affect sporulation, induce biofilm formation and promote colonization of earthworms and environmental reservoirs. Furthermore, the lytic activity and high specificity of bacteriophages provide a promising resource for the development of innovative treatments for human pathogens, including B. anthracis.

In this study, we describe the genome and host range of vB_BanS-Tsamsa, a novel temperate phage obtained from B. anthracis isolated in Etosha National Park (Etosha), Namibia. We named the phage Tsamsa, which in Hai|om means “place where the winds blow closed” referring to the endless vista of the Etosha pan and the dust devils that form there. Tsamsa phage is a giant siphovirus capable of infecting some members of Bacillus cereus sensu lato.

Materials and Methods

Phage Isolation and Preparation

We obtained isolates of the siphovirus from two carcass sites in Etosha, a 22,915 km² national park in northern Namibia with abundant wildlife populations that exhibit regular occurrences of anthrax infections (reviewed in [11]). Field sampling was authorized by the Namibian Ministry of Environment and Tourism under permit number 1448/2009 to HHG. Bacteriophages were isolated from a B. anthracis isolate obtained from a
swab of a plains zebra (Equus quagga) carcass from 2006 (Etosha Ecological Institute (EEI) carcass number: EB060318-01NVY; GPS coordinates: $-18.99376$, $15.01504$) and from soil collected near another plains zebra carcass in 2010 (EEI carcass number: EB100228-01MK; GPS coordinates: $-19.1731$, $15.92603$). Prior diagnostic testing showed that both carcasses were positive for *B. anthracis* isolates from the two carcasses were identified as genotype 6 and genotype 4 in the A cluster of *B. anthracis*, respectively [12]. Genotypes 6 and 4 are closely related members of a dominant *B. anthracis* strain that has been causing outbreaks in Etosha for a very long time [12]. Phages were obtained from the two samples by culturing to enrich for bacteria and by exposure to mitomycin C to induce prophages in the host genome to transition into a vegetative state. Methods for enrichment culture and induction are described by Sambrook [13] and Van Twest and Kropinski [14]. We did not obtain phages from either sample without induction. We used two approaches, one for the swab isolate and another for the soil sample, as follows:

1. For the swab isolate, we inoculated 3 ml of Bovine Heart Infusion medium (BHI, BD Difco, Sparks, MD, USA) with a single *B. anthracis* colony isolated from the swab and incubated the culture overnight at 37°C with aeration. We diluted the overnight culture 100-fold in 3 ml of BHI and incubated it at 37°C with aeration for one hour. To induce the release of prophages from the genome, mitomycin C was added to achieve a final concentration of 2.5 μg ml⁻¹. The culture was incubated at 37°C with aeration for 20 hr and pelleted for 15 min at 3000xg. The supernatant was filtered through a 0.22 μm filter unit and stored at 4°C.

2. For the soil sample, 20 g of soil were added to 15 ml of 1% Nutrient Broth medium (BD Difco, Sparks, MD, USA). After vortexing briefly, the sample was incubated at 35°C with aeration overnight. The resulting culture was centrifuged and the supernatant collected. Then we added 1% Nutrient Broth medium to attain a final volume of 7.5 ml. The sample was incubated in 1 μg ml⁻¹ mitomycin C for 30 minutes with gentle aeration at 30°C. Then the sample was filtered (0.22 μm) and concentrated using a Vivaspin 20 concentrator (Sartorius Stedim, Bohemia, NY, USA) by adding 10 ml of 1% nutrient broth medium and centrifuging the sample at 3000xg for 10 min. The resulting phage extract was stored at 4°C.

**Preparation of Plate Stocks of the Two Tsamsa Phage Isolates**

Phage preparations were purified and concentrated using standard techniques [13,15]. Preliminary plaque assays were performed with phage extracts from the two carcass site samples (swab isolate and soil sample) to harvest concentrated plate stocks. Soft agar overlays were performed as described previously by Adams [16]. Briefly, five microliters of a spore preparation of an avirulent pXO1 + pXO2 — *B. anthracis* strain (6602 R1 [17]) were spotted on top of plates seeded with a lawn of bacteria. Each plate was tilted to allow the phage solution to run down the plate. If phage titers were higher than $10^8$ pfu/ml, more dilutions were made. A phage was considered positive for infection of a certain strain if single plaques could be observed on one of the spots. We tested a set of 55 strains for susceptibility to Tsamsa phage (Table 1).

**Phage Host Range**

Tsamsa phage was originally propagated on *B. anthracis* 6602 R1 (an avirulent strain that lacks both pXO1 and pXO2 virulence plasmids, [17]). Using the spot-on-the-lawn method and a 100-fold and a 10,000-fold dilution of the phage stock, 10 μl were spotted on top of plates seeded with a lawn of bacteria. Each plate was tilted to allow the phage solution to run down the plate. If phage titers were higher than $10^8$ pfu/ml, more dilutions were made. A phage was considered positive for infection of a certain strain if single plaques could be observed on one of the spots. We tested a set of 55 strains for susceptibility to Tsamsa phage (Table 1).

**Phage Morphology**

The Tsamsa phage was negatively stained with 2% uranyl acetate on carbon-coated copper grids (Carbolung, Quantifoil, Jena, Germany) and observed in a Philips CM12 TEM microscope at 120 kV acceleration voltage with a Gatan Orius 1 k camera.

**Genome Sequencing and Genetic Characteristics**

We performed a standard DNA extraction using phenol-chloroform-isooamyl-alcohol of the cesium chloride purified stock [13]. After washing with 70% ethanol and drying, the DNA pellet was resuspended in sterile ultrapure water. We sequenced DNA from the two isolates using a SMRT sequencing approach (Pacific Biosciences RS) with 10 kb and 800 bp insert libraries (C2 chemistry) and one SMRT sequencing cell for each library. We used the standard error-correction workflow and SMRT portal software 1.3.1 for assembly of 36166 post-filter reads (with 2582 bp average read length). Open reading frames were predicted by RAST [20] and edited manually. The genome sequence was deposited at GenBank under accession number KC481682. The unassembled reads for both sequencing runs are available in the DNA Databank of Japan Sequence Read Archive under accession number DRA001229.

**Phylogenetic Analysis**

Comparisons were made between sequences of the large terminase subunit of Tsamsa phage and 17 other *Bacillus* phages (GenBank accession numbers: 955214, 955254, 7070024, 12980149, 13164871, 14697218, 14697335, 14697413, 14697831, NC_001884, NC_006557, NC_007457, NC_007458, NC_007734, NC_007814, NC_011167, NC_011421). Alignments and phylogenetic tree construction were performed in Geneious version 6.1 ( Biomatters Ltd., http://www.geneious.com). Muscle [21] was used to align the terminase gene (with 16 iterations). Then MrBayes 3.1.2 [22,23] was used to build the tree and determine Bayesian posterior probabilities (with the Monte Carlo Markov Chain run for 1.1 x 10⁶ generations).
Table 1. Host range of Tsamsa phage on 43 different *Bacillus* strains and 12 non-*Bacillus* strains and lytic spectrum of Tsamsa endolysin for a subset of the *Bacillus* strains. Presence of lysis is indicated by + and absence of lysis is indicated by -. n.d.: not determined.

| Strain name | Organism | Notes | Source | Phage lysis | Endolysin lysis |
|-------------|----------|-------|--------|-------------|----------------|
| 6602 R1     | *Bacillus anthracis* | pXO1-pXO2 negative | [39] | + | + |
| Sterne      | *Bacillus anthracis* | pXO2 negative | Institut Pasteur #7702 | + | + |
| Weybridge UM44 | *Bacillus anthracis* | pXO2 negative | [39] | + | + |
| Ames-non reverting | *Bacillus anthracis* | pXO2 negative | U.S. Dept. of Agriculture, Ames, Iowa | + | n.d. |
| Vollum 1b   | *Bacillus anthracis* | Laboratory Strain | + | n.d. |
| PAK-1       | *Bacillus anthracis* | Pakistan isolate, M. Hugh-Jones collection | - | n.d. |
| S69         | *Bacillus cereus* | [17] | + | n.d. |
| LA 925      | *Bacillus cereus* | CHUV | - | + |
| ATCC 14579  | *Bacillus cereus* | ATCC | - | + |
| ATCC 11778  | *Bacillus cereus* | ATCC | + | + |
| ATCC 10702  | *Bacillus cereus* | ATCC | - | + |
| ATCC 10876  | *Bacillus cereus* | ATCC | + | n.d. |
| DSM 2302    | *Bacillus cereus* | DSM | - | + |
| BO 366      | *Bacillus cereus* | This study | - | - |
| BO 372      | *Bacillus cereus* | This study | - | - |
| BO 493      | *Bacillus cereus* | This study | - | - |
| DSM 4218    | *Bacillus cereus* | DSM | - | + |
| ATCC 33019  | *Bacillus cereus* | ATCC | + | + |
| ATCC 14737  | *Bacillus cereus* | ATCC | + | + |
| DSM1274     | *Bacillus cereus* | DSM | - | + |
| ATCC 27522  | *Bacillus cereus* | ATCC | - | + |
| NCTC 11143  | *Bacillus cereus* | NCTC | - | + |
| NCIMB 8705  | *Bacillus cereus* | NCIMB | + | + |
| ATCC 6464   | *Bacillus cereus* | ATCC | - | + |
| B346        | *Bacillus cereus* | Mouse isolate | - | + |
| DSM360      | *Bacillus cereus* | DSM | - | - |
| HER1399     | *Bacillus cereus* | HER | + | + |
| WSB 10550   | *Bacillus cereus* | WSB | - | - |
| WSB 10556   | *Bacillus cereus* | WSB | - | - |
| WSB 10566   | *Bacillus cereus* | WSB | - | - |
| WSB 10583   | *Bacillus cereus* | WSB | - | - |
| DSM4421     | *Bacillus thuringiensis* | DSM | - | + |
| WSB 10204   | *Bacillus thuringiensis* | WSB | - | - |
| HER1211     | *Bacillus thuringiensis* | HER | + | + |
| Kurstaki    | *Bacillus thuringiensis* | Industry isolate | - | - |
| ATCC 10792  | *Bacillus thuringiensis* | ATCC | + | + |
| DSM168      | *Bacillus subtilis* | DSM | - | - |
| DSM675      | *Bacillus subtilis* | DSM | - | - |
| ATCC 23059  | *Bacillus subtilis* | ATCC | - | n.d. |
| DSM395      | *Bacillus sphaericus* | DSM | - | - |
| DSM90       | *Bacillus megaterium* | DSM | - | - |
| WSB 10550   | *Bacillus weihenstephanensis* | WSB | - | + |
| WSLC 3009   | *Listeria ivanovii* | WSLC | - | - |
| ATCC BAA-679| *Listeria monocytogenes* | ATCC | - | - |
| PSK         | *Staphylococcus aureus* | Laboratory Stock | - | - |
| Twort       | *Staphylococcus aureus* | Laboratory Stock | - | - |
Endolysin Production and Determination of Lytic Spectrum

Purified Tsamsa phage endolysin was recombinantly produced [24] and characterized. Lysis activity against different bacterial strains was tested with a turbidity reduction assay or plate lysis. Turbidity reduction experiments were performed by harvesting and washing an overnight bacterial culture (grown in half-strength BHI medium) in PBS (120 mM NaCl, 50 mM Na₂HPO₄, pH 8.0) buffer. Cell density was adjusted to OD₆₀₀ₙₐₜ of 1±0.05. 100 μM of lysin were added to the wells of a 96-well plate and 200 OD₆₀₀ₙₐₜ of 0.4–0.6 and diluted 1/100 in PBS immediately before bacterial suspension was added. Wells were measured at a wavelength of 600 nm in 10 second intervals until clear turbidity reduction was observed relative to the control.

Lysis activity by plate lysis assay was tested as follows. The Bacillus strains were grown in half-strength BHI medium to an OD₆₀₀ₙₐₜ of 0.4–0.6 and diluted 1/100 in PBS immediately before plating. The freshly spread lawns of Bacillus cells on ½ BHI agar plates were air dried for 30 min. Ten microliters of a 10-fold serial dilution (50, 5 and 0.5 μM) of the purified endolysin were spotted onto the plates. Spots were air-dried and plates incubated at 30°C for 16 hours. Cleared spots indicating cell lysis were assessed visually.

Results

We obtained two phage isolates from two different carcass sites (one from soil and one carcass swab). Both phages were morphologically very similar and unusually large compared with previously described Siphoviridae of Bacillus. We found by sequencing the genomes of these two isolates that both phages were 100% identical. The representative phage was named Tsamsa and its characteristics were investigated further.

Tsamsa Features a Broad Host Range in the B. anthracis Subgroup

A set of 55 bacterial strains was analyzed for susceptibility to Tsamsa. The phage lysed seven out of 25 B. cereus strains as well as two out of five B. thuringensis strains and six out of seven B. anthracis strains (6602 R1, Sterne, Weybridge UM44, Ames-non-reverting and Vollum 1b but not PAK-1) (Table 1). It should be noted that PAK-1 is an atypical member of B. anthracis, belonging to the A2 branch that contains very few isolates [25], and is resistant to both Cherry phage and Gamma phage [26]. Tsamsa phage is unable to infect B. subtilis 168 and DSM675, B. megaterium DSM90, B. sphaericus DSM395, B. weihenstephanensis WSBC10550 as well as all tested strains from other bacterial genera (Listeria ivanovii and L. monocytogenes, Staphylococcus aureus and S. epidermidis, Salmonella Typhimurium, Escherichia coli, Streptococcus salivarius, Lactococcus lactis or Enterococcus faecalis) (Table 1). Thus it is a narrow host-range virus that infects some members of the Bacillus cereus sensu lato group and exhibits moderate specificity for B. anthracis.

Tsamsa is a Giant Siphovirus

Most bacteriophages belong to the Order Caudovirales, which contains three families: Myoviridae with a contractile tail, Siphoviridae with a non-contractile flexible tail and Podoviridae with a short, non-contractile tail. Tsamsa exhibits typical siphovirus morphology, featuring a long, flexible and non-contractile tail of 440 nm (not including baseplate structure) and an isometric head of 82 nm in diameter (Figure 1A and B). Individual tail striations (disk-like structure) and a baseplate structure with appendages are visible (Figure 1C). The head features visible individual capsomers (Figure 1D), an observation previously made for a different class of large virulent phages belonging to the Spounavirinae subfamily within the family Mphoviridae [27]. Because of the large head dimensions and our experience from similarly sized myoviruses, we anticipate a triangulation number of 16 or higher but experimental proof is lacking.

The Phage Tsamsa Genome is Large and Unique

Genome sequencing and assembly resulted in a single large contig with an average coverage of 530-fold of error-corrected SMRT reads (Figure 2). Both phage isolates were identical. A repeat structure of 284 bp at both genome ends was identified during assembly and confirmed in restriction profiles (Figure 3). Methylome analysis revealed no base modifications in the genome. The genome sequence is 168,876 bp in length. Tsamsa features 272 open reading frames, 17 tRNA and 2 pseudo-tRNA genes. Database matches of predicted proteins encoded by Bacillus anthracis phage Tsamsa are provided in Table S1. The GC content is 34%, similar to published genome sequences of B. anthracis.

Virus Proteome

Virion proteins of Tsamsa were separated on a 10–20% SDS gradient PAGE. Resulting bands were extracted and protein content identified by mass spectrometry [28]. Six bands were...
allocated to gene products. The tape measure protein is present in two protein bands of 280 and 100 kDa in size, presumably because of instability of the large protein or post-translational modification. gp206 was identified in a band with an estimated mass of 38 kDa and gp199 and 207 were identified in bands of 26.5 and 19 kDa, respectively.

Figure 1. Electron microscopy of Tsamsa phage. TEM images were acquired from a preparation of pure phage particles negatively stained with 2% uranyl acetate on carbon-coated copper grids (Quantifoil, Jena, Germany) and observed using a Philips CM12 microscope at 120 kV acceleration voltage with a Gatan Orius digital camera. A, Preparation overview. B, Close-up of single phage particle. C, Details of the phage tail distal end. D, Details of the phage head structure. Individual capsomers are visible, an observation previously made for SPO1-related phages ([27]). Scale bars represent 100 nm. doi:10.1371/journal.pone.0085972.g001

Figure 2. Genome map of Tsamsa phage. Open reading frames are drawn to scale and transcription direction is indicated by arrows. Selected proteins with putative function are labeled. Genetic modules (i.e. structural genes, early genes) are indicated by coloring. doi:10.1371/journal.pone.0085972.g002
Figure 3. Restriction profile of Tsamsa phage. 500 ng DNA were digested with different restriction enzymes for 2 h at 37°C and electrophoresed. Clear band separation up to 24 kb in size was achieved and the restriction profiles matched with the sequenced genome size. The terminal redundancy location and size was determined from the fragment sizes as previously described [13,15]. Enzymes used: 1: Alw44I (NEB); 2: Eco91I (Fermentas); 3: NheI (NEB); 4: PacI (NEB); 5: SwaI (NEB); 6: Van91I (Fermentas); 7: XcmI (NEB). M1: Lambda 19 Mix Size standard (Fermentas); M2: 1 kb size standard (Fermentas). Numbers to the left indicate band size in kb.
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Comparison with Other Sequenced Bacillus Phages

The sequence of the large terminase subunit of Tsamsa phage was compared with 17 previously sequenced Bacillus phages: four B. anthracis phages (Fah, Cherry, WBeta, Gamma), two B. thuringiensis phages (IEBH, BTCS33), four B. subtilis phages (SPBeta, SPO1, SPP1, phi105), one B. clarkii phage (BCJA1c), two B. cereus phages (PBC1, TP21-L) and four B. pumilus phages (Andromeda, Curly, Eoghan, Finn). The Tsamsa phage terminase clearly differs from previously described phages isolated from B. anthracis (Figure 4), which were shown to be derived from a single B. anthracis prophage named W [29,30].

The Tsamsa Endolysin Shows Broad Activity

The endolysin of phage Tsamsa (gp217) was cloned and recombinantly produced in E. coli. The 35.78 kDa protein features an isoelectric point of 9.01 and contains three domains, one Amidase_2 enzymatic active domain (PF01510) and two SH3_3 (PF08239) cell-wall binding domains. Lytic activity of the protein was assessed with either turbidity reduction or plate lysis assay and activity against a set of Bacillus strains is outlined in Table 1. The Tsamsa endolysin features a broad spectrum of lytic activity and is able to lyse more Bacillus cereus sensu lato strains than the phage can infect.

Figure 4. Phylogenetic relationship of the large terminase subunit gene in 18 Bacillus phages. Phages are color coded according to their host bacterium: B. anthracis in red, B. cereus in blue, B. clarkii in orange, B. pumilus in green, B. subtilis in purple and B. thuringiensis in black. The tree and posterior probabilities were determined from 1.1x10⁶ generations with MrBayes 3.1.2 [22,23].
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Discussion

We obtained identical phage isolates of an unusually large siphovirus induced from *B. anthracis* in two different environmental samples. Such a large siphovirus is rarely isolated; only nine of the 539 Siphovirus genomes in the NCBI Genome database have a larger genome than Tsamsa phage (six *Caulobacter* Siphovirus genomes in the NCBI Genome database have a larger genome than Tsamsa phage). To our knowledge, Tsamsa is the largest sequenced siphovirus infecting *Bacillus*. Two larger *Bacillus* siphoviruses are known but not characterized: *B. mycoides* phage N5 and *B. thuringiensis* phage II, both of which feature approximately 20% larger dimensions than Tsamsa and are speculated to be identical (H.-W. Ackermann, personal communication). Large siphoviruses may be isolated infrequently due to a bias in phage isolation procedures towards smaller phages. Consequently, the distribution and abundance of large siphoviruses are poorly understood. Such a sampling bias has been reported for the so-called Jumbo-Phages [31].

Like many siphoviruses, the genome is structured in functional modules. The early gene cluster (genes for DNA replication, modification and repair, host takeover and nucleotide metabolism) spans roughly 70% of the genome, indicating active participation of virus-encoded genes in the metabolic processes associated with replication in the host cell. It is notable that the Tsamsa genome encodes three tyrosine integrase/recombinase enzymes of the Crc/XERD type (gp94, gp227, gp253; [32,33]), which exhibit no homology to each other. Tsamsa features a temperate lifestyle and these three enzymes may serve as means to integrate into different *attB* sites and ensure a large host range for lysogeny. Further work will elucidate the specificity and activity of the three recombinases. We also note the presence of two *ig*-domain containing proteins, gp233 and gp213 [33,34], which may play accessory roles during infection [35]. Ig-like domains in structural proteins were recently shown to play a potential role in phage attachment to mucosa of humans and animals by interaction of the Ig-like domain with the mucous glycan residues, providing a non-host derived immunity [36].

Tsamsa features distributed homologies in its structural proteins to SPO1-related phages: A511, A9, LP65 and SPO1 [27]. This finding is very unusual because SPO1-related phages belong to the *Spounavirinae* subfamily within an unrelated family of bacteriophages (Myoviridae) [27]. In addition, Tsamsa is a temperate phage and phages within the *Spounavirinae* are strictly virulent. Tsamsa also displays individual capsomers thought to be a hallmark of the *Spounavirinae* (Figure 1) [27]. An assessment of the virus particle proteome identified six structural protein bands in Tsamsa, namely the tape measure protein in two bands, and one band each for gp199, 206, 207 and 222. Tsamsa features an unusually long tail of 440 nm, which corresponds with the large size of the tape measure protein (3123 aa) [37,38] and the protein is disproportionately large in comparison to other sequenced bacteriophages. The large unknown gene 221 likely encodes for a tail fiber component, with the C-terminus featuring significant homologies to Celllobiosidase, S-layer associated endoglucanase or glycoside hydrolase domains. The Tsamsa endolysin (gp217) is accompanied by a holin (gp215) and is active when produced recombinantly. The enzyme shows a broad lytic spectrum and lysed more *B. cereus* and *B. thuringiensis* strains than the phage infects. Thus the Tsamsa lysis might be useful as an antimicrobial agent against some *Bacillus cereus* sensu lato organisms.

In conclusion, we present vB_BanS-Tsamsa, a novel temperate phage obtained from *B. anthracis* that is specific to some members of the *Bacillus cereus* sensu lato group. To our knowledge Tsamsa is the largest sequenced siphovirus infecting *Bacillus* organisms.

Supporting Information

Table S1 Database matches of predicted proteins encoded by *Bacillus anthracis* phage Tsamsa. (DOCX)

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Author Contributions

Conceived and designed the experiments: HHG CL MS FE MJL WMG RC J. Korlach WB J. Klumpp. Performed the experiments: HHG CL MS FE MJL WMG RC J. Korlach WB J. Klumpp. Analyzed the data: HHG CL FE RC J. Korlach J. Klumpp. Contributed reagents/materials/analysis tools: HHG CL MS FE MJL WMG RC J. Korlach WB J. Klumpp. Wrote the paper: HHG CL MS FE MJL WMG RC J. Korlach WB J. Klumpp.

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