Complete genome sequence of *Pseudoalteromonas* phage vB_PspS-H40/1 (formerly H40/1) that infects *Pseudoalteromonas* sp. strain H40 and is used as biological tracer in hydrological transport studies

René Kallies¹*, Bärbel Kiesel¹, Matthias Schmidt², Johannes Kacza³, Nawras Ghanem¹, Anja Narr¹, Jakob Zopfi⁴, Lukas Y. Wick¹, Jörg Hackermüller⁵, Hauke Harms¹,⁶ and Antonis Chatzinotas¹,⁶

**Abstract**

*Pseudoalteromonas* phage vB_PspS-H40/1 is a lytic phage that infects *Pseudoalteromonas* sp. strain H40. Both, the phage and its host were isolated in the 1970s from seawater samples collected from the North Sea near the island of Helgoland, Germany. The phage particle has an icosahedral capsid with a diameter of ~43 to 45 nm and a long non-contractile tail of ~68 nm in length, a typical morphology for members of the *Siphoviridae* family. The linear dsDNA genome of *Pseudoalteromonas* phage vB_PspS-H40/1 has a sequence length of 45,306 bp and a GC content of 40.6%. The genome has a modular structure and contains a high proportion of sequence information for hypothetical proteins, typically seen in phage genome sequences. This is the first report of the complete genome sequence of this lytic phage, which has been frequently used since the 1990s as biological tracer in hydrogeological transport studies.

**Keywords:** *Pseudoalteromonas* phage, *Siphoviridae*, AquaDiva, Marine phage, Virus, Genome, Bacteriophages as tracers

**Introduction**

*Pseudoalteromonas*, affiliated with the order Alteromonadales [1, 2] of the *Gammaproteobacteria* [2, 3], is a genus of heterotrophic, Gram-negative marine bacteria [4]. Members of this genus are widely distributed in marine ecosystems and have attracted interest due to their frequent association with eukaryotic hosts and their production of biologically active compounds [5–7]. Both inhibitory as well as synergistic chemical interactions between strains of *Pseudoalteromonas* and various marine eukaryotes have been described [8], indicating that members of this genus are potentially involved in complex ecological networks across trophic levels. Viruses, as the most abundant biological entity in the oceans, are a major cause of host mortality and thus key players within these ecological networks; they influence host community structures and thereby also influence global biogeochemical cycles and genetic landscapes [9].

As of April 2016, 14 complete *Pseudoalteromonas* phage genomes have been deposited at GenBank (10 of them unpublished). Ten representatives belong to the *Caudovirales* order (three siphoviruses, four podoviruses, two myoviruses and one unclassified caudovirus), one is a corticovirus and three are unclassified viruses. *Pseudoalteromonas* phages have been shown to represent a significant group of phages in the ocean [10, 11], making it likely that the number of yet unknown phage genomes is much higher. Characterization of additional *Pseudoalteromonas* phage genomes is a further step towards a better understanding of the diversity, the biology and the ecological impact of this group of phages.

* Correspondence: rene.kallies@ufz.de
*¹Department of Environmental Microbiology, Helmholtz Centre for Environmental Research - UFZ, 04318 Leipzig, Germany
Full list of author information is available at the end of the article

© The Author(s). 2017 Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.
and contributes to an improved interpretation of viral metagenome data and dynamics of viral populations in the environment [12–14]. Moreover, comparison of potentially closely related viral genomes is a prerequisite to understand virus evolution and intraspecies genomic variation [15, 16].

In this report we describe the genome of the Pseudoalteromonas phage vB_PspS-H40/1, isolated in 1978 from the North Sea near the island of Helgoland (Germany) [17]. Notably, this phage has been used as a non-reactive biological agent to trace the flow of water in surface and subsurface environments and promises utility in (geo-)hydrological transport studies [18–21]. According to the scheme for the nomenclature of viruses the phage was re-named from H40/1 to vB_PspS-H40/1 [22].

Organism Information
Classification and features
The bacterial host H40 was isolated from seawater samples collected between 1969 and 1978 near the island of Helgoland in the North Sea [17]. Sequence analysis of the 16S-rRNA gene revealed H40 as a member of the Pseudoalteromonas genus. The partial 16S-rRNA sequence was deposited at GenBank (acc. no. KX236488). Strain H40 was used as the bacterial host for screening of lytic marine bacteriophages from the same sampling site resulting in the isolation of phage vB_PspS-H40/1 [17].

Pseudoalteromonas phage vB_PspS-H40/1 is a lytic phage forming clear, well-contrasted plaques of four to five mm in diameter. Transmission electron microscopy showed that vB_PspS-H40/1 is a B1 morphotype with an icosahedral capsid of 42.7 nm in length (±1.7 nm) and 44.5 nm in width (± 2 nm). The long, non-contractile tail had a length of 67.5 nm (± 3.9 nm) and a diameter of 6.7 nm (± 0.6 nm) (Fig. 1). These morphological characteristics are typical for members belonging to the Siphoviridae family of the order Caudovirales [23].

The phage surface is moderately charged (zeta potential of −11 ± 3 mV (100 mM K2HPO4/KH2PO4, pH = 7)) and of moderate hydrophobicity (water contact angle = 53 ± 3°) as determined by standard physico-chemical characterization methods of bacterial surfaces (e.g. [24]).

Pseudoalteromonas phage vB_PspS-H40/1 has a linear dsDNA genome comprising 45,306 bp with a GC content of 40.6%. It showed the highest similarity (55.3% identity) over the whole genome to Pseudoalteromonas phage H103 (GenBank acc. no. KP994596), an unclassified representative of the Caudovirales order infecting the marine host Pseudoalteromonas marina [25] (Fig. 2). Phylogenetic analysis of the terminase large subunit (TerL) amino acid sequence grouped phage vB_PspS-H40/1 together with phage H103 in one clade (Fig. 3). Both phages shared a most recent common ancestor with TerL sequences found...
in unclassified members of the Caudovirales order and (probably) prophage sequences from members of the bacterial family Enterobacteriaceae [26, 27]. These unclassified phages belong to all three families of the Caudovirales order, i.e. Siphoviridae, Podoviridae and Myoviridae. Taken together, TerL-based phylogenetic analysis indicates phage vB_PspS-H40/1 occupies (perhaps together with phage H103) a phylogenetic position distinct from established genera of the Siphoviridae family.

Phylogenetic classification and general features of Pseudoalteromonas phage vB_PspS-H40/1 are summarized in Table 1.

---

**Genome sequencing information**

**Genome project history**

Pseudoalteromonas phage vB_PspS-H40/1 is one of the few known marine siphovirus isolates [28] and belongs to a group of important phages found in the ocean [10, 11]. Genome sequencing of this phage will increase available information and facilitate future studies on diversity, evolution and ecological impact of marine viruses. A second reason to select this phage for sequencing is its frequent application in biological tracing experiments [18–21]. Phage vB_PspS-H40/1 is one of the marine phages that are currently used in the frame of the Collaborative Research...
Centre AquaDiva to trace the hydrological flow and reactive transport of colloidal particles from the surface into the Earth's subsurface [29]. Environmental influences might inactivate a still to define percentage of transported phages. Knowledge of a phage genome will facilitate the detection of this phage using PCR and thus allow to (quantitatively) distinguish between biologically active (e.g. detected by plaque assay) from inactive phages and might hence help in the interpretation of findings from these transport experiments.

The dsDNA genome of phage vB_PspS-H40/1 was sequenced using the Illumina MiSeq system. Experiments, genome assembly, annotation and submission to GenBank were performed at the Department of Environmental Microbiology at the Helmholtz Centre for Environmental Research - UFZ, Leipzig, Germany. The sequencing project was started in December 2015 and finished in February 2016 and its outcome is available in the Genome Online Database under project number Gp0133998. The complete annotated genome sequence was submitted to Genbank (GenBank acc. no. KU747973). Information on the project is summarized in Table 2.

Growth conditions and genomic DNA preparation
The bacterial host *Pseudoalteromonas* sp., strain H40 was grown and maintained in 2216E medium [30] (containing nutrients at 50% of the original concentration) at 20 °C. The phage was propagated on its host in petri dishes with 2216E agar (with nutrients as above) using the double agar-layer technique. Five ml of SM buffer (100 mM NaCl, 8 mM MgSO₄·7H₂O, 50 mM Tris–HCl, pH 7.5) and a few drops of chloroform were added to the plates after confluent lysis of bacterial host cells. Plates were gently shaken for 2 h at room temperature, supernatant was collected and cell debris was removed by centrifugation at 10,000 × g for 15 min. One volume of chloroform was then added to the supernatant, gently mixed and centrifuged at 5,000 × g for 5 min. The phage particle-containing upper phase was

| Table 1 Classification and general features of *Pseudoalteromonas* phage vB_PspS-H40/1 infecting *Pseudoalteromonas* sp. strain H40 |
| MIGS ID | Property | Term |
|---------|----------|------|
| Classification | Domain: N/A | |
| | Genome group: dsDNA viruses, IDA | |
| | Phylum: unassigned | |
| | Class: unassigned | |
| | Order: Caudovirales | TAS [23] |
| | Family: Siphoviridae | TAS [23] |
| | Genus: unassigned | |
| | Species: unassigned | |
| | Strain: vB_PspS-H40/1 | |
| Particle shape | Isometric capsid, long non-contractile tail | IDA |
| Gram strain | N/A | |
| Cell shape | N/A | |
| Motility | N/A | |
| Sporulation | N/A | |
| Temperature range | N/A | |
| Optimum temperature | N/A | |
| pH range; optimum | N/A | |
| Carbon source | N/A | |
| MIGS-6 Habitat | Marine water column | TAS [17] |
| MIGS-6.3 Salinity | N/A | |
| MIGS-22 Oxygen requirement | N/A | |
| MIGS-15 Biotic relationship | Intracellular parasite of *Pseudoalteromonas* sp. strain H40 | TAS [17] |
| MIGS-14 Pathogenicity | Virulent phage of *Pseudoalteromonas* sp. strain H40 | IDA |
| MIGS-4 Geographic location | North Sea, Helgoland, Germany | TAS [17] |
| MIGS-5 Sample collection | 1978 | TAS [17] |
| MIGS-4.1 Latitude | 54°10’N | IDA |
| MIGS-4.2 Longitude | 7°52’E | IDA |
| MIGS-4.4 Altitude | N/A | |

| Evidence codes - IDA inferred from direct assay, TAS traceable author statement, N/A not applicable. These evidence codes are from the Gene Ontology project [46] |

| Table 2 Project information |
| MIGS ID | Property | Term |
|---------|----------|------|
| MIGS 31 | Finishing quality | Finished |
| MIGS-28 Libraries used | One paired-end Illumina library |
| MIGS 29 | Sequencing platforms | Illumina MiSeq |
| MIGS 31.2 Fold coverage | ~1200x |
| MIGS 30 | Assemblers | Geneious Assembler version R6 |
| MIGS 32 | Gene calling method | RAST, GLIMMER, GeneMark.hmm |
| Locus Tag | NA* |
| Genbank ID | KU747973 |
| GenBank Date of Release | Jun 07, 2016 |
| GOLD ID | Gp0133998 |
| BIOPROJECT | NA* |
| MIGS 13 Source Material Identifier | NA* |
| Project relevance | Diversity of marine bacteriophage, Hydrological transport studies |

*NA not available
passed through a 0.22 μm polyvinylidene fluoride CHROMAFIL® membrane filter to remove unlysed host cells and debris. The resulting phage suspension was stored at 4 °C. DNA from phage particles was extracted following the protocol of Thurber et al. [31].

**Genome sequencing and assembly**

The extracted phage DNA was sheared into ~300 to 500 bp fragments using the Covaris M220 Focused-ultrasonicator™ instrument and one paired-end library was prepared with the NEBNext® Ultra™ DNA Library Prep Kit for Illumina®. Sequencing was performed at the Helmholtz Centre for Environmental Research - UFZ on an Illumina MiSeq system (2 × 150 bp). In total, 418,468 paired-reads were obtained for *Pseudoalteromonas* phage vB_PspS-H40/1. Raw reads were split into 10 subsets (approximately 42,000 reads for each subset) to facilitate improved assembly [32]. Independent assemblies were performed for each subset by Geneious Assembler (version R6) resulting in nearly the same single contig for each of the subsets but with different starting points indicating a circularly permuted genome of phage vB_PspS-H40/1. For confirmation, PCR primers were designed matching the ends of the contigs with an outward orientation and used in PCR. The resulting amplicon was Sanger sequenced and used to close the contigs for *Pseudoalteromonas* phage vB_PspS-H40/1. The coverage was estimated by reference mapping of the raw reads to the contig resulting in an approximate 1200-fold coverage (~92% of all reads) of the 45,306 bp genome.

**Genome annotation**

Genes and ORFs in the phage genome were predicted using a combination of three gene calling methods: the RAST annotation server [33], GLIMMER3 [34] and GeneMark.hmm [35]. Only ORFs that were predicted by two of the three gene calling methods were included in the annotation. Functional annotation of translated ORFs was improved by BLASTp alignments against the NCBI non-redundant database [36]. In addition, RPS-BLAST searches against the Conserved Domain Database [37] and HMMER search [38] against the UniProtKB database were performed. Protein domains were predicted using the COG [39], Pfam [40], TIGRFAMs [41] and KEGG [42] databases. Phoebius [43] was used to predict signal peptides and transmembrane helices.

**Genome properties**

The complete genome of *Pseudoalteromonas* phage vB_PspS-H40/1 was assembled into one linear contig of 45,306 bp with a GC content of 40.6%. In total, 73 putative coding sequences were predicted in the phage genome (Fig. 2, Additional file 1: Table S1). Seventeen of these 73 protein coding genes were assigned to putative protein functions. The functions of the remaining 56 putative protein coding genes remained unknown and they were annotated as hypothetical proteins. One gene with a signal peptide was identified together with eight proteins containing transmembrane helices. *Pseudoalteromonas* phage vB_PspS-H40/1 genome properties are summarized in Table 3 and genes assigned to COG functional categories are listed in Table 4.

**Insights from the genome sequence**

When all 73 predicted CDSs were subjected to functional annotation only 17 CDSs could be assigned to a specific function. These functions were related to DNA packaging, head and tail assembly, DNA replication and metabolism (Fig. 2 and Additional file 1: Table S1). Twenty-nine of the predicted CDSs, including mainly hypothetical proteins but also TerL and structural proteins, showed highest similarity to the unclassified *Caudovirales* member *Pseudoalteromonas* phage H103 after blastp analysis (Fig. 2). Highest similarity of other CDSs was found to marine *Pseudoalteromonas* phages belonging to the *Siphoviridae* family, i.e. *Pseudoalteromonas* phage TW1 (GenBank acc. no. KC542353), *Pseudoalteromonas* phage Pq0 (GenBank acc. no. NC_029100) and *Pseudoalteromonas* phage H105/1 (GenBank acc. no. NC_015293). However, proteins involved in DNA replication (helicase, RecA-NTPase and methylase) were related to those found in *Vibrio* phage H188 (GenBank acc.no. KT160311) and *Escherichia* phage vB_EcoM-ep3 (GenBank acc. no. NC_025430), two members of the *Myoviridae* family, suggesting mosaicism of the genome.

Phylogenetic inferences deduced from the TerL amino acid sequence showed no close phylogenetic relationship to any of the established *Siphoviridae* genera (Fig. 3).

| Attribute                        | Value   | % of Total |
|----------------------------------|---------|------------|
| Genome size (bp)                 | 45,306  | 100.00     |
| DNA coding (bp)                  | 42,786  | 94.44      |
| DNA G + C (bp)                   | 17,376  | 40.60      |
| DNA scaffolds                     | 1       | 0.00       |
| Total genes                      | 73      | 100.00     |
| Protein coding genes             | 73      | 100.00     |
| RNA genes                        | 0       | 0.00       |
| Pseudo genes                     | 0       | 0.00       |
| Genes in internal clusters       | 0       | 0.00       |
| Genes with function prediction   | 17      | 23.29      |
| Genes assigned to COGs           | 6       | 8.22       |
| Genes with Pfam domains          | 18      | 24.66      |
| Genes with signal peptides       | 1       | 1.34       |
| Genes with transmembrane helices | 8       | 10.96      |
| CRISPR repeats                   | 0       | 0.00       |

Table 3 Genome statistics
The characterized complete genome of lytic Pseudoalteromonas phage vB_PspS-H40/1 that was isolated from seawater in the North Sea improves our knowledge of this significant group of phages. The linear dsDNA genome has a size of 45,306 bp and a GC content of 40.6%. The obtained sequencing data indicate phage vB_PspS-H40/1 uses headful packaging strategy and that the genome is circularly permuted. Among the 73 protein coding sequences only 17 were functionally annotated. Transmission electron microscopy and phylogenetic analysis of TerL sequences suggest this phage might belong to a genus of a yet unclassified group of Siphoviridae. Next to studies on specific phage-host interactions in marine systems, phage vB_PspS-H40/1 will be used in surface and groundwater tracer experiments and its genome sequence and morphological description will help interpreting results from these studies.

### Conclusions

The characterized complete genome of lytic Pseudoalteromonas phage vB_PspS-H40/1 that was isolated from seawater in the North Sea improves our knowledge of this significant group of phages. The linear dsDNA genome has a size of 45,306 bp and a GC content of 40.6%. The obtained sequencing data indicate phage vB_PspS-H40/1 uses headful packaging strategy and that the genome is circularly permuted. Among the 73 protein coding sequences only 17 were functionally annotated. Transmission electron microscopy and phylogenetic analysis of TerL sequences suggest this phage might belong to a genus of a yet unclassified group of Siphoviridae. Next to studies on specific phage-host interactions in marine systems, phage vB_PspS-H40/1 will be used in surface and groundwater tracer experiments and its genome sequence and morphological description will help interpreting results from these studies.

### Table 4

| Code | Value | Sargo | Description |
|------|-------|-------|-------------|
| J    | 0     | 0.00  | Translation, ribosomal structure and biogenesis |
| A    | 0     | 0.00  | RNA processing and modification |
| K    | 1     | 1.34  | Transcription |
| L    | 3     | 4.11  | Replication, recombination and repair |
| B    | 0     | 0.00  | Chromatin structure and dynamics |
| D    | 0     | 0.00  | Cell cycle control, Cell division, chromosome partitioning |
| V    | 0     | 0.00  | Defense mechanisms |
| T    | 0     | 0.00  | Signal transduction mechanisms |
| M    | 1     | 1.34  | Cell wall/membrane biogenesis |
| N    | 0     | 0.00  | Cell motility |
| U    | 0     | 0.00  | Intracellular trafficking and secretion |
| O    | 0     | 0.00  | Posttranslational modification, protein turnover, chaperones |
| C    | 0     | 0.00  | Energy production and conversion |
| G    | 0     | 0.00  | Carbohydrate transport and metabolism |
| E    | 0     | 0.00  | Amino acid transport and metabolism |
| F    | 0     | 0.00  | Nucleotide transport and metabolism |
| H    | 0     | 0.00  | Coenzyme transport and metabolism |
| I    | 0     | 0.00  | Lipid transport and metabolism |
| P    | 0     | 0.00  | Inorganic ion transport and metabolism |
| Q    | 0     | 0.00  | Secondary metabolites biosynthesis, transport and catabolism |
| R    | 1     | 1.34  | General function prediction only |
| S    | 1     | 1.34  | Function unknown |
| -    | 66    | 90.41 | Not in COGs |

The total is based on the total number of protein coding genes in the genome.
7. Skovhus TL, Holmström C, Kjelleberg S, Dahllof I. Molecular investigation of the distribution, abundance and diversity of the genus Pseudomonas in marine samples. FEMS Microbiol. Ecol. 2007;61(2):348–61.

8. Bowman JP. Bioactive compound synthetic capacity and ecological significance of marine bacterial genus Pseudomonas. Mar Drugs. 2007;5(4):20–41.

9. Suttle CA. Marine viruses - major players in the global ecosystem. Nat Rev Microbiol. 2007;5(10):801–12.

10. Wilches A, Bels ST, Gelderblom HR, Brinkhoff T, Muyzer G, Schütz C. Bacteriophage diversity in the North Sea. J Appl Environ Microbiol. 1998;64(12):38–33.

11. Duhaime MB, Wilches A, Waldmann J, Teeling H, Gökçener F. Ecogenomics and genome landscapes of marine Pseudomonas aeruginosa phage H105/1. ISME J. 2011;5:107–21.

12. Kang I, Oh HM, Kang D, Cho JC. Genome of a SAR116 bacteriophage shows the prevalence of this phage type in the oceans. Proc Natl Acad Sci U S A. 2013;110(3):12343–8.

13. Zhao Y, Temperton B, Thrash JC, Schwalbach MS, Vergin KL, Landry ZC, et al. Phenotypic and genotypic variations within a single bacteriophage species. Virology. 2011;13:179–84.

14. Moebus K, Nattkemper H. Bacteriophage sensitivity patterns among bacteria isolated from marine waters. Helgol Meeressunters. 1981;34:375–85.

15. Flynn R, Hunkeler B, Gerber C, Burn C, Rossi P, Aragno M. Bacteriophages as indicators and indicators of transport in a porous medium. Environ Sci Technol. 2016.

16. Koppich M, Prangishvili D, Hendrix RW, Barnford DH. Genomics of bacterial and archaeal viruses: dynamics within the prokaryotic virosphere. Microbiol Mol Biol Rev. 2011;75(4):610–35.

17. Pope WH, Jacobs-Sera D, Russell DA, Peebles CL, Al-Atrache Z, Alcoser TA, et al. Expanding the diversity of mycobacteriophages: insights into genome architecture and evolution. PLoS One. 2011;6(1):e16329.

18. Ceyssens PJ, Glonti T, Kropinski NM, Lavigne R, Chanishvili N, Kulakov L, et al. Phenotypic and genotypic variations within a single bacteriophage species. Virology. 2011;13:179–84.

19. Metcalf N, Kiesler B, Kallies R, Harms H, Chatzinotas A, Wick L. Marine phages as tracers: effects of size, physico-chemical and physico-chemical surface properties on transport in a porous medium. Environ Sci Technol. 2016. accepted for publication.

20. Kropinski AM, Prangishvili D, Lavigne R. Position paper: The creation of a rational scheme for the nomenclature of viruses and Archaea. Environ Microbiol. 2009;11(1):2775–7.

21. King AMQ, Adams MJ, Carstens EB, Lefkowitz EJ. Virus taxonomy: classification and nomenclature of viruses: Ninth Report of the International Committee on Taxonomy of Viruses. San Diego: Elsevier Academic Press; 2012.

22. Flynn R, Hunkeler D, Guerin C, Burn C, Rossi P, Aragno M. Geochemical and we will help you at every step:

23. Marchler-Bauer A, Derbyshire MK, Gonzales NR, Lu S, Chitsaz F, Gerbier LY, et al. CDD: NCBI’s conserved domain database. Nucleic Acids Res. 2013;41(Database issue):D222–6.

24. Finn RD, Clements J, Arndt W, Miller BL, Wheeler TJ, Schreiber F, et al. HHMWER web server: 2015 update. Nucleic Acids Res. 2015;43(W1):W30–8.

25. Pope WH, Jacobs-Sera D, Russell DA, Peebles CL, Al-Atrache Z, Alcoser TA, et al. Phenotypic and genotypic variations within a single bacteriophage species. Virology. 2011;13:179–84.

26. Goldscheider G, Haller L, Poté J, Wildi W, Zopfi J. Characterizing water circulation and contaminant transport in Lake Geneva using bacteriophage tracer experiments and limnological methods. Environ Sci Technol. 2007;41(15):5252–8.

27. Khanellia M, Sato Y, Kawashima M, Furumichi M, Tanabe M. KEGG as a reference resource for gene and protein annotation. Nucleic Acids Res. 2000;28(11):33–6.

28. Kall L, Krogh A, Sonnhammer EL. A combined transmembrane topology and signal peptide prediction method. J Mol Biol. 2005;348(5):1267–76.

29. Sullivan MJ, Petty NK, Beatson SA. Easyfig: a genome comparison visualizer. Bioinformatics. 2011;27(7):1009–10.

30. Haft DH, Delcher AL, Powers EC, Salzberg SL. Identifying bacterial genes and endosymbiont DNA with Glimmer. Bioinformatics. 2007;23(6):673–9.

31. Kurz A, Wiberg P, Jansson M, Schmitt J, Gyllensten U. The prevalence of this phage type in the oceans. Proc Natl Acad Sci U S A. 2013;110(3):12343–8.

32. Kall L, Krogh A, Sonnhammer EL. A combined transmembrane topology and signal peptide prediction method. J Mol Biol. 2005;348(5):1267–76.

33. Sullivan MJ, Petty NK, Beatson SA. Easyfig: a genome comparison visualizer. Bioinformatics. 2011;27(7):1009–10.

34. Guindon S, Gascuel O. A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. Syst Biol. 2003;52(5):696–704.

35. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. Basic local alignment search tool. J Mol Biol. 1990;215(3):403–10.

36. Marchler-Bauer A, Derbyshire MK, Gonzales NR, Lu S, Chitsaz F, Gerbier LY, et al. CDD: NCBI’s conserved domain database. Nucleic Acids Res. 2015;43(Database issue):D222–6. doi:10.1093/nar/gku1211.

37. Finn RD, Clements J, Arndt W, Miller BL, Wheeler TJ, Schreiber F, et al. HHMWER web server: 2015 update. Nucleic Acids Res. 2015;43(W1):W30–8. doi:10.1093/nar/gkv1070.

38. Punta M, Coggill PC, Eberhardt RY, Mistry J, Tate J, Boursnell C. The Pfam database: sequence similarity alignments to protein families. Nucleic Acids Res. 2012;40(Database issue):D290–301. doi:10.1093/nar/gkr1065.

39. Haft DH, Selengut JD, Richter RA, Harkins D, Basu MK, Beck E. TIGRFAMs and Genome Properties in 2013. Nucleic Acids Res. 2013;41(Database issue):D387–95. doi:10.1093/nar/gks97.

40. Kall L, Krogh A, Sonnhammer EL. A combined transmembrane topology and signal peptide prediction method. J Mol Biol. 2005;348(5):1267–76.