Computational characterization and analysis of molecular sequence data of *Elizabethkingia meningoseptica*

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**Abstract**

**Objective:** *Elizabethkingia meningoseptica* is a multidrug resistance strain which primarily causes meningitis in neonates and immunocompromised patients. Being a nosocomial infection causing agent, less information is available in literature, specifically, about its genomic makeup and associated features. An attempt is made to study them through bioinformatics tools with respect to compositions, embedded periodicities, open reading frames, origin of replication, phylogeny, orthologous gene clusters analysis and pathways.

**Results:** Complete DNA and protein sequence pertaining to *E. meningoseptica* were thoroughly analyzed as part of the study. *E. meningoseptica* G4076 genome showed 7593 ORFs it is GC rich. Fourier based analysis showed the presence of typical three base periodicity at the genome level. Putative origin of replication has been identified. Phylogenetically, *E. meningoseptica* is relatively closer to *E. anophelis* compared to other *Elizabethkingia* species. A total of 2606 COGs were shared by all five *Elizabethkingia* species. Out of 3391 annotated proteins, we could identify 18 unique ones involved in metabolic pathway of *E. meningoseptica* and this can be an initiation point for drug designing and development. Our study is novel in the aspect in characterizing and analyzing the whole genome data of *E. meningoseptica*.

**Keywords:** Bioinformatics, *Elizabethkingia meningoseptica*, Genome annotation, Pathway analysis, Subtractive genomics

**Introduction**

In 1959, Elizabeth O King, discovered *Elizabethkingia* (renamed in 2005) [1], earlier known as *Chryseobacterium*. It is a non-glucose fermenting, non-motile, catalase-oxidase positive gram negative bacteria belonging to *Flavobacteriaceae* family, ubiquitous in soil, fresh and salty water [2]. The genus comprises of six species [3] that is, *E. meningoseptica* associated with meningitis and sepsis in premature neonates, [4, 5] *E. anophelis* isolated from the midgut of *Anopheles gambiae* mosquitoes which causes respiratory tract illness in human [6], *E. miricola*, isolated from condensation water on the Mir space station of Russia collected in 1997 [7], and *E. brunniana*, *E. ursingii* and *E. occulta* (three CDC genomospecies) [8].

*Elizabethkingia meningoseptica* is causative agent of meningitis in neonates and sepsis in immunocompromised patients [9]. The occurrence of nosocomial infection has risen, mainly in patients, with prolonged hospitalization, treated with invasive procedures, subsequently on use of broad-spectrum antimicrobials as well as having concomitant infections [10]. The mortality rate in patients infected with *E. meningoseptica* is significantly higher due to its unusual resistance pattern and mechanism [11]. Further studies are needed to initiate the most effective therapeutic approach. One can follow the time
on previous studies, expectation value cut off of $10^{-4}$ and minimum bit score of 100 used as threshold to short-list non-homologous proteins [27]. Further, these non-homologous proteins were queried against Database of Essential Genes (DEG) server to get a list of essential genes for *E. meningoseptica* using e-value cut off $10^{-10}$ and bit score value of 100 as threshold [28]. These shortlisted essential genes that were non-homologous to host and essential for bacteria were studied further with respect to metabolic pathway.

**Metabolic pathway analysis and subcellular localization prediction**

Essential non-homologous proteins of *E. meningoseptica* were further analyzed using KAAS (KEGG Automated Annotation Server) in order to study metabolic pathways [29]. KEGG analysis performed BLAST comparison against available KEGG gene database and provide metabolic pathway maps including KO and EC number for a particular gene. To determine the location of proteins in a cell PSORTb version 3.0 server was used [30]. The essential gene subjected to BLASTP analysis against FDA approved drug targets from Drugbank to search novel drug targets. Targets with identification of more or equal 80% are druggable targets and others that show considerable low degree of matching with already approved drug target can be used as novel targets for new drug identification [31].

**Results**

**Genomic features of *E. meningoseptica* G4076 and its comparison with other species**

The whole genome data of *E. meningoseptica* G4076 having length of 3,873,125 bp showed a mean GC content of 36.5%, number of genes as per annotation is 3477 and the percentage base composition viz %A $\approx$ %T i.e., 31.76 and %G $\approx$ %C i.e., 18.23 calculated using ORIS software [14] (Additional file 1: Figure S1) which is in agreement with Chargaff’s parity rule [32]. Open reading frame is effective in identifying genes that encodes proteins. Total number of 7593 ORFs were found in whole genome. The products are of varying length and it shows that the number of ORFs found are actually slightly more than the annotated number of proteins (Additional file 1: Figure S2). To visualize sequence conservation, the circular genome plot was created using CG view Server (Additional file 1: Figure S3). Gene coding segments of *E. meningoseptica* genome does show the typical three-base periodicity indicating underlying codon structure that enables us to predict and identify all possible genes in majority of the bacterial genome with very high accuracy [17]. Additional file 1: Figure S4A shows all the bases considered for the fourier spectrum and indicates the

**Main text**

**Methods**

**Genome analysis of *E. meningoseptica* G4076 and its comparisons with Elizabethkingia family**

The whole genome (Accession Number NZ_CP016376) and protein sequences of *Elizabethkingia meningoseptica* G4076 were downloaded from NCBI (www.ncbi.nlm.nih.gov). Nucleotide composition of genome was obtained using ORIS software [14]. To find all open reading frames in the genome, ORF finder, a graphical tool was used (https://www.ncbi.nlm.nih.gov/orffinder/) [15]. CG-Viewer was used for plotting circular plot of genomes [16]. Discrete Fourier Transform based computational approach using customized python codes was carried out to see the typical three-base periodicity feature embedded in *E. meningoseptica* genomic sequence [17]. Rapid Annotation using Subsystem Technology (RAST) server was carried out for studying genome annotation [18, 19]. Ori-Finder [20] and ORISv1.0 [14] software tools were used to identify putative origin of replication (oriC) sites in the genome. MegaX software was utilized to carry out phylogenetic analysis for species within the same genus such as *E. miricola*, *E. meningoseptica*, *E. anophelis*, *E. bruuniana*, *E. ursingii* and *E. occulta* as well as *Flavobacterium columnare* ATCC49512, *Riemerella anatipes* G4076 and *E. meningoseptica* (Host) were downloaded from NCBI database [24, 25]. Out of the total 3406 proteins in *E. meningoseptica*, hypothetical proteins and proteins having length less than 100 amino acids were discarded. Remaining 2503 proteins were subjected to BLASTP against proteomes of *Homo sapiens* [26]. Based

**Subtractive genomics based computational analysis**

All protein sequences of *Elizabethkingia meningoseptica* G4076 and *Homo sapiens* (Host) were downloaded from NCBI database [24, 25]. Out of the total 3406 proteins in *E. meningoseptica*, hypothetical proteins and proteins having length less than 100 amino acids were discarded. Remaining 2503 proteins were subjected to BLASTP against proteomes of *Homo sapiens* [26]. Based consuming and labor-intensive experimental approach but advancement in bioinformatics field provided enormous software tools, that are used to analyze and extract information from the molecular sequence, structure, expression and pathway data [12, 13].

The current study focused on analyzing the whole genome data of *Elizabethkingia* to unravel the embedded features hitherto not reported, secondly to explore the possibility of getting some lead in the directions of possible novel therapeutic candidates. Accordingly, we have studied genomic features, origin of replication sites, phylogenetic relationships, comparative genomics among *E. meningoseptica* species and further explored subtractive genomics approach together with pathway analysis.

The whole genome (Accession Number NZ_CP016376) and protein sequences of *Elizabethkingia meningoseptica* G4076 were downloaded from NCBI (www.ncbi.nlm.nih.gov). Nucleotide composition of genome was obtained using ORIS software [14]. To find all open reading frames in the genome, ORF finder, a graphical tool was used (https://www.ncbi.nlm.nih.gov/orffinder/) [15]. CG-Viewer was used for plotting circular plot of genomes [16]. Discrete Fourier Transform based computational approach using customized python codes was carried out to see the typical three-base periodicity feature embedded in *E. meningoseptica* genomic sequence [17]. Rapid Annotation using Subsystem Technology (RAST) server was carried out for studying genome annotation [18, 19]. Ori-Finder [20] and ORISv1.0 [14] software tools were used to identify putative origin of replication (oriC) sites in the genome. MegaX software was utilized to carry out phylogenetic analysis for species within the same genus such as *E. miricola*, *E. meningoseptica*, *E. anophelis*, *E. bruuniana*, *E. ursingii* and *E. occulta* as well as *Flavobacterium columnare* ATCC49512, *Riemerella anatipes* G4076 and *E. meningoseptica* (Host) were downloaded from NCBI database [24, 25]. Out of the total 3406 proteins in *E. meningoseptica*, hypothetical proteins and proteins having length less than 100 amino acids were discarded. Remaining 2503 proteins were subjected to BLASTP against proteomes of *Homo sapiens* [26]. Based
presence of three base periodic signal as seen in most of bacterial genomes. Signal strength is prominent for purine-pyrimidine (Additional file 1: Figure S4B) whereas in the case of individual bases it is considerably low (Additional file 1: Figure S4C–F).

RAST server shows annotated data indicating 3477 putative genes, 61 RNAs which includes 4,4,4 (55, 165, 23S) ribosomal RNAs and 49 tRNAs and 335 subsystems (set of functional role) under 27 categories [18]. Sixty two coding sequences were related with antibiotics resistance and toxic compounds which suggests *E. meningoseptica* might be multiple drug resistant (Additional file 1: Figure S5).

Ori-Finder (a web based software tool for finding oriCs) predicted oriC region of 649 bp ranging from 740,720 bp to 741,368 bp having three DnaA box sequence motifs (TTATCCACA) with no more than one mismatch. Further, replication related gene, dnaA located from 2,613,273 to 2,614,727 bp which is followed by dnaN gene (Fig. 1A) [20]. A cluster of three DnaA boxes and two AT rich DNA unwinding elements (DUE) are indication of functional chromosomal origin (Fig. 1F). Similar kind of result was found with ORIS v1.0 software tool. DNA asymmetry, distribution of DnaA boxes as well as location of the dnaA gene help in predicting OriC regions [33–36]. Both graphs enable us to pin-point or identify ORI/TER site. The difference in the position (genome coordinates) of OriC predicted by Ori-Finder and ORIS are well within 1 kb and hence, close agreement.

Genomic comparison among *Elizabethkingia* species [*E. meningoseptica* G4076 (WP_016198861.1), *E. miricola* BM10 (WP_034866598.1), *E. ursingii* G4123 (WP_078402796.1), *E. anophelis* NUHP1 (WP_009086312.1) *E. brunniiana* G0146 (WP_034866598.1), *F. columnare* ATCC49512 (WP_014166114.1), *R. anatipestifer* ATCC11845 (WP_004918717.1)] has been done using MEGAX software. It depicts phylogenetic relatedness by comparing homology of protein sequence specifically by comparing homology of protein sequence specifically using e-value cut off 10–4 and bit score > 100, shortlisted 692 proteins that are essential for *E. meningoseptica* that further can be considered as potential therapeutic targets. Unique *E. meningoseptica* essential proteins non-homologous to host further subjected to BLASTP against proteins of *Homo sapiens* (host). Using e-value cut off 10–4 and bit score > 100, it was found total of 2052 proteins were non-homologous to host protein. Thereafter, these proteins were subjected to BLAST analysis using DEG server and using e-value cut off 10–10 and bit score > 100, shortlisted 692 proteins that are essential for *E. meningoseptica* but absent in host (Additional file 1: Table S1). DEG contains gene that plays important role in cell survival and can be novel targets for antibacterial drugs (Fig. 2).

**Prediction of essential genes in *Elizabethkingia meningoseptica***

Subtractive genomic analysis is unique, fast and efficient method for identifying essential genes in pathogenic species that are non-homologous to human (host). These non-homologous essential genes can be used as putative drug targets against pathogens [38]. The genome of *E. meningoseptica* G4076 has 3391 annotated proteins. After exclusion of protein which are <100 amino acids and hypothetical, remaining 2503 were subjected to BLASTP against proteins of *Homo sapiens* (host). Using e-value cut off 10–4 and bit score > 100, it was found total of 2052 proteins were non-homologous to host protein. Thereafter, these proteins were subjected to BLAST analysis using DEG server and using e-value cut off 10–10 and bit score > 100, shortlisted 692 proteins that are essential for *E. meningoseptica* G4076 but absent in host (Additional file 1: Table S1). DEG contains gene that plays important role in cell survival and can be novel targets for antibacterial drugs (Fig. 2).

**Metabolic pathway analysis of essential gene and subcellular localization prediction**

The shortlisted non-homologous essential genes were analyzed using KEGG database for metabolic pathway annotation. It was found, only 41 out of 692, are present in pathogen as unique pathways (Table 1). Majority of them were involved in DNA binding response regulator, ribosomal proteins, replication and repair, Glycan biosynthesis, protein folding and sorting, two-component system, biotin metabolism and ATP transporters. It is very important for drug designing to determine whether target protein resides on cell surface or in cytoplasm. Localization of proteins play important role in drug binding and action. Subcellular localization reveals, out of 41 target proteins, 80% of total are cytoplasmic, rest located in periplasm or cytoplasmic membrane and no extracellularly proteins were obtained (Additional file 1: Figure S8). Extracellularly secreted proteins may be better opted for vaccine development. Here, it is clear that majority of proteins resides in cytoplasm and cytoplasmic membrane that further can be considered as potential therapeutic targets. Unique *E. meningoseptica* essential proteins non-homologous to host further subjected to BLASTP against FDA approved drug targets from Drugbank which shortlisted to 18 target proteins. Out of which penicillin binding protein (2), ABC transporter ATP binding proteins (2) that targets for broad-spectrum antibiotics. The rest includes ribosomal proteins (rpsB, rpsl, rpsG, rpsJ, rpsE,
Fig. 1. Output results using Ori-Finder—A Z curve (AT, GC, RY and MK disparity) for *Elizabethkingia meningoseptica* G4076. Peaks with the diamonds show DnaA boxes, bold arrow indicates oriC location, and solid short black lines show replication marker genes i.e., dnaN, dnaA, gidA, hemE etc). B–E Cumulative GC, AT, MK, RY skew graph of *E. meningoseptica* G4076 using ORISv1.0 software tool having window size 40,000 with increment of 4000 bp. Bold solid arrow indicates putative ori site. F OriC sequence wherein showed DnaA boxes (capitalized and underlined) with not more than one mismatch to *E. coli* DnaA box. AT clusters, in oriC region are shown in bold.
rpsM, rpsK, rpsD, rplD, rplP), recombination protein (recR), DNA polymerase subunit III tau (dnaX), and signal peptidase which could be further explored as starting point for discovering novel drug candidate. Ribosomal proteins can be more suitable candidates for drug binding as it mainly involves in translation. Another work also
Table 1  Unique and novel essential genes in *E. meningoseptica* G4076

| Protein product                                   | Protein name                                      | KEGG orthology | EC       | Subcellular localization |
|--------------------------------------------------|---------------------------------------------------|----------------|----------|--------------------------|
| **Two component system**                         |                                                   |                |          |                          |
| WP_016198590.1                                    | Response regulator transcription factor           | K07665         | –        | Cytoplasmic              |
| WP_016199055.1                                    | LytTR family DNA-binding domain-containing protein | K07705         | –        | Cytoplasmic              |
| WP_016198441.1                                    | Response regulator transcription factor           | K07665         | –        | Cytoplasmic              |
| WP_016199099.1                                    | LytTR family DNA-binding domain-containing protein | K07705         | –        | Cytoplasmic              |
| WP_016200043.1                                    | VanW family protein                               | K18346         | –        | Unknown                  |
| WP_016199769.1                                    | Two-component sensor histidine kinase             | K07636         | 2.7.13.3 | Cytoplasmic membrane     |
| **Beta-lactam resistance**                       |                                                   |                |          |                          |
| WP_016170088.1                                    | Penicillin-binding protein 2a                      | K05515         | 3.4.16.4 | Cytoplasmic membrane     |
| WP_016170028.1                                    | Transglycosylase domain-containing protein         | K05366         | 2.4.1.129| Cytoplasmic membrane     |
| **DNA replication**                              |                                                   |                |          |                          |
| WP_016199473.1                                    | DNA primase                                       | K02316         | 2.7.7.101| Cytoplasmic              |
| WP_019051072.1                                    | DNA polymerase III subunit gamma/tau a            | K02343         | 2.7.7.7  | Cytoplasmic              |
| WP_016197797.1                                    | DNA polymerase III subunit delta a                | K02341         | 2.7.7.7  | Cytoplasmic              |
| **Homologous recombination**                     |                                                   |                |          |                          |
| WP_016198910.1                                    | Holliday junction branch migration protein RuvA   | K03550         | 3.6.4.12 | Cytoplasmic              |
| WP_016198560.1                                    | Holliday junction branch migration DNA helicase RuvB | K03551      | 3.6.4.12 | Cytoplasmic              |
| WP_016200627.1                                    | Recombination protein RecR                        | K06187         | –        | Cytoplasmic              |
| WP_016199534.1                                    | DNA replication and repair protein RecF           | K03629         | –        |                          |
| **Translation**                                   |                                                   |                |          |                          |
| WP_009085459.1                                    | 30S ribosomal protein S2 a                        | K02967         | –        | Cytoplasmic              |
| WP_016200426.1                                    | 30S ribosomal protein S9 a                        | K02996         | –        | Cytoplasmic              |
| WP_009087383.1                                    | 30S ribosomal protein S12 a                       | K02946         | –        | Cytoplasmic              |
| WP_009087378.1                                    | 30S ribosomal protein S7 a                        | K02992         | –        | Cytoplasmic              |
| WP_016197802.1                                    | 30S ribosomal protein S10 a                       | K02946         | –        | Cytoplasmic              |
| WP_009087341.1                                    | 50S ribosomal protein L4 a                        | K02926         | –        | Cytoplasmic              |
| WP_016197785.1                                    | 50S ribosomal protein L16 a                       | K02878         | –        | Cytoplasmic              |
| WP_009087327.1                                    | 50S ribosomal protein L14                         | K02874         | –        | Cytoplasmic              |
| WP_016197784.1                                    | 50S ribosomal protein L24                         | K02895         | –        | Cytoplasmic              |
| WP_009087314.1                                    | 30S ribosomal protein S5 a                        | K02988         | –        | Cytoplasmic              |
| WP_016197779.1                                    | 50S ribosomal protein L15                         | K02876         | –        | Cytoplasmic              |
| WP_016197776.1                                    | 30S ribosomal protein S13 a                       | K02892         | –        | Cytoplasmic              |
| WP_009087288.1                                    | 30S ribosomal protein S11 a                       | K02948         | –        | Cytoplasmic              |
| WP_016170211.1                                    | 30S ribosomal protein S4 a                        | K02986         | –        | Cytoplasmic              |
| WP_016170209.1                                    | 50S ribosomal protein L17                         | K02879         | –        | Cytoplasmic              |
| WP_016198662.1                                    | 30S ribosomal protein S16                         | K02959         | –        | Cytoplasmic              |
| WP_016200457.1                                    | 50S ribosomal protein L20                         | K02887         | –        | Cytoplasmic              |
| WP_016199408.1                                    | 30S ribosomal protein S1                          | K02945         | –        | Cytoplasmic              |
| WP_016200561.1                                    | 50S ribosomal protein L9                         | K02939         | –        | Cytoplasmic              |
| **ABC Transporters**                             |                                                   |                |          |                          |
| WP_016198610.1                                    | ATP-binding cassette domain-containing protein a   | K09812         | –        | Cytoplasmic membrane     |
| WP_016198126.1                                    | ABC transporter ATP-binding protein a             | K09810         | 7.6.2. - | Cytoplasmic membrane     |
| **Protein export**                               |                                                   |                |          |                          |
| WP_026149261.1                                    | Signal peptidase I b                              | K03100         | 3.4.21.89| Cytoplasmic membrane     |
| **Methane metabolism**                           |                                                   |                |          |                          |
| WP_016198341.1                                    | Phosphoenolpyruvate carboxylase a                 | K01595         | 4.1.1.31 | Cytoplasmic              |
| **Base excision repair**                         |                                                   |                |          |                          |
| WP_016170024.1                                    | Endonuclease III                                  | K10773         | 4.2.99.18| Cytoplasmic              |
| **Biotin metabolism**                            |                                                   |                |          |                          |
| WP_016199146.1                                    | Dethiobiotin synthase                            | K01935         | 6.3.3.3  | Cytoplasmic              |

* Potential therapeutic candidates as per FDA approved drugbank
lend support for choosing the specific drug target [39]. In that regard, computational analysis may include homology modelling and docking of selected candidate.

Discussions
Meningitis and sepsis is a major illness in newborn and immunocompromised patients caused by *Elizabethkingia meningoseptica*. Though typical clinical diagnostics are used to identify the illness but a greater understanding of molecular based diagnosis is desired and it is a long term goal. Increase in number of cases in Intensive care units (ICUs) makes it big challenge for clinicians to deal and manage. In this context, comprehensive analysis of whole genome data and pathway analysis were explored as we do not see much work related to computational analysis. Accordingly, bioinformatics approach was undertaken for characterizing molecular sequence data of *Elizabethkingia*. Our study identified 41 unique proteins in *Elizabethkingia* with respect to the host using subtractive genomics which further narrow down to 18 therapeutic target proteins using *in-silico* comparative genomics. The suitable shortlisted ribosomal proteins which are linked to translation may be useful for future treatment and management of the infection. We have studied in an integrated fashion of considering and analyzing sequence data of *E. meningoseptica* together with pathway analysis. Our study is small step in the direction of rapid diagnosis and possible drug development.

Limitations
The current investigation is limited to in silico study only.

Abbreviations
*E.*: Elizabethkingia; ORF: Open reading frame; GC: Guanine cytosine; RAST: Rapid Annotation using Subsystems Technology; COGs: Cluster of groups; CDC: Centre of Disease Control; DNA: Deoxyribonucleic acid; NCBI: National Center for Biotechnology Information; AT: Adenine thymine; MK: Amino-keto; RY: Purine pyrimidine; ORIS: ORI search; OrC: Origin or replication C; MEGA: Molecular evolutionary genetic analysis; ATCC: American type culture collection; bp: Base pair; kb: Kilobase pair; RNA: Ribonucleic acid; CDS: Coding sequences; DUE: DNA unwinding element; TER: Terminal; ICUs: Intensive Care Units; e-value: Expectation value; BLAST: Basic Local Alignment Search Tool; DEG: Database of essential genes; KEGG: Kyoto Encyclopedia of Genes and Genomes; FDA: Food and Drug Administration.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s13104-022-08011-5.

Additional file 1: Figure S1. Percentile distribution of DNA base composition in *E. meningoseptica* G4076 genome. **Figure S2.** Open reading frame viewer—a window showing ORFs on the interval from 1 to 50,000 nucleotides. **Figure S3.** Circular genomic plot of *E. meningoseptica* G4076 having Accession Number NZ_CP016376 was downloaded from NCBI site https://www.ncbi.nlm.nih.gov/genome/14625?genome_assembly_id=309079. All the protein sequences (numbering 3406) available in FASTA format were used for BLASTP analysis against human dataset option. Selected protein sequences (described in material method section) were further used as input for subtractive genomic analysis.

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Authors’ contributions
AK—conceptualization, methodology, formal analysis, writing—review and editing, visualization, supervision. NK—conceptualization, writing—review and editing, supervision, project administration. NG—formal analysis, investigation, data curation, writing—original draft. All authors read and approved the final manuscript.

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Availability of data and materials
The whole genome sequence of *Elizabethkingia meningoseptica* G4076 having Accession Number NZ_CP016376 was downloaded from NCBI site https://www.ncbi.nlm.nih.gov/genome/14625?genome_assembly_id=309079. All the protein sequences (numbering 3406) available in FASTA format were used for BLASTP analysis against human dataset option. Selected protein sequences (described in material method section) were further used as input for subtractive genomic analysis.

Declarations
Ethics approval and consent to participate
The authors declare that no ethical approval is required for current study.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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References
1. King EO. Studies on a group of previously unclassified bacteria associated with meningitis in infants. Am J Clin Pathol. 1959;31(3):241–7. https://doi.org/10.1093/ajcp/31.3.241.
2. Ceyhan M, Celik M. *Elizabethkingia meningoseptica* (Chryseobacterium meningosepticum) infections in children. Int J Pediatr. 2011. https://doi.org/10.1155/2011/215237.
3. Lin J, Lai C, Yang C, Huang Y. *Elizabethkingia* infections in humans: from genomics to clinics. Microorganisms. 2019;7:295. https://doi.org/10.3390/microorganisms7040295.
4. Hazuka BT, Dajani AS, Talbot K, Keen BM. Two outbreaks of *Flavobacterium meningosepticum* type *E* in neonatal intensive care unit. J Clin Microbiol. 1977;6(5):450–5.
5. Amer MZ, Bandey M, Bukhari A, Nemenquani D. Neonatal meningitis caused by *Elizabethkingia meningoseptica* in Saudi Arabia. J Infect Dev Ctries. 2011;5(10):745–7. https://doi.org/10.3855/jidc.1570.
6. Kempf P, Matthews H, Glaeser SP, Martin K, Lodders N, Faye I. *Elizabethkingia anaphelis* sp. nov., isolated from the midgut of the mosquito
10. Pereira GH, Garcia Dde O, Abboud CS, Barbosa VL, Silva PS. Nosocomial
9. Bloch KC, Nadarajah R, Jacobs R.
8. Nicholson AC, Gulvik CA, Whitney AM, et al. Revisiting the taxonomy
7. Kim KK, Kim MK, Lim JH, Park HY, Lee ST. Transfer of
20. Gao F, Zhang C. Ori-Finder: a web-based system for finding
17. Tiwari S, Ramachandran D, Bhattacharya A, Bhattacharya S, Ramaswamy R. Prediction of probable genes by Fourier analysis of genomic sequences. Bioinformatics. 1997;13(3):265–70.
15. ORFfinder. https://www.ncbi.nlm.nih.gov/orffinder/.
14. Singh, et al. ORIS: an interactive software tool for prediction of replication origins based on Z-curve analysis. J Phylogenetics. 2014;4(2):104–12. https://doi.org/10.12741/jphy.2014.2.104.
13. Alfat-Ul-Amin M, et al. Recent trends in computational biomediical research. Life. Basel. 2022. https://doi.org/10.3390/life10010027.
12. Hogeweg P. The roots of bioinformatics in theoretical biology. PloS Comput Biol. 2011. https://doi.org/10.1371/journal.pcbi.1002021.
11. Young SM, Lingam G, Tambyah PA.
10. 2174/1389202915999140328162938.
9. Zhiqiang CT, Zhang R, Ou HY. The Z curve database: a graphic representation of genome sequences. Bioinformatics. 2003;19(5):593–9. https://doi.org/10.1093/bioinformatics/btg041.
8. Roy S. Molecular markers in phylogenetic studies—a review. J Phylogenetics. 2014;4(2):2. https://doi.org/10.12741/jphy.2014.2.2.
7. Uddin R, Saeed K. Identification and characterization of potential drug targets by subtractive genome analyses of methicillin resistant Staphylococcus aureus. Front Mol Biosci. 2018;5:48. https://doi.org/10.3389/fmolb.2018.00048.
6. Zhiqiang CT, Zhang R, Ou HY. The Z curve database: a graphic representation of genome sequences. Bioinformatics. 2003;19(5):593–9. https://doi.org/10.1093/bioinformatics/btg041.
5. Mackiewicz P, Zakrzewska-Czerwińska J, Zawilak A, Dudek MR, Ćebrat S. Where does bacterial replication start? Rules for predicting the origin region. Nucleic Acids Res. 2004;32(13):3781–91. https://doi.org/10.1093/nar/gkh699.
4. Gao F. Recent advances in the identification of replication origins based on the Z-curve method. Curr Genom. 2014;15(2):104–12. https://doi.org/10.2174/1389202915999140328162938.
3. Zhang CT, Zhang R, Ou HY. The Z-curve database: a graphic representation of genome sequences. Bioinformatics. 2003;19(5):593–9. https://doi.org/10.1093/bioinformatics/btg041.
2. Hogeweg P. The roots of bioinformatics in theoretical biology. PloS Comput Biol. 2011. https://doi.org/10.1371/journal.pcbi.1002021.
1. Young SM, Lingam G, Tambyah PA.