Comparative genomic analysis of a naturally competent *Elizabethkingia anophelis* isolated from an eye infection

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*Elizabethkingia anophelis* has now emerged as an opportunistic human pathogen. However, its mechanisms of transmission remain unexplained. Comparative genomic (CG) analysis of *E. anopheles* endophthalmitis strain surprisingly found from an eye infection patient with twenty-five other *E. anophelis* genomes revealed its potential to participate in horizontal gene transfer. CG analysis revealed that the study isolate has an open pan genome and has undergone extensive gene rearrangements. We demonstrate that the strain is naturally competent, hitherto not reported in any members of *Elizabethkingia*. Presence of competence related genes, mobile genetic elements, Type IV, VI secretory systems and a unique virulence factor arylsulfatase suggests a different lineage of the strain. Deciphering the genome of *E. anophelis* having a reservoir of antibiotic resistance genes and virulence factors associated with diverse human infections may open up avenues to deal with the myriad of its human infections and devise strategies to combat the pathogen.

*Elizabethkingia* belonging to the family *Flavobacteriaceae* is a genus with 4 known species: *E. meningoseptica*, *E. anophelis*, *E. miricola* and *E. endophytica*. Since its discovery in 1959, *E. meningoseptica* has been described to be involved in diverse human infections such as meningitis, keratitis and sepsis among immunocompromised individuals. In 2003, *E. miricola* was reported for the first time from the condensed water samples of the Russian space station Mir. *E. miricola* was later found to be associated with sepsis, bacteraemia and pneumonia. *E. endophytica* isolated from corn has not yet been associated with any of the human infections. In 2011, *E. anophelis* was discovered from the midgut of *Anophelis gambiae* mosquito. It has been linked to neonatal meningitis, nosocomial outbreaks and catheter associated infections with high mortality rates.

*Elizabethkingia* strains have been found to encode metallo-beta-lactamases conferring carbapenem resistance. Because of the bacterium's innate resistance to several classes of antibiotics, treatment of *Elizabethkingia* infections is challenging. Several studies have reported that patients with severe underlying diseases and a history of antibiotic exposure are more susceptible to *Elizabethkingia* related nosocomial infections. Scant information is available on the potential modes of infection and their ability to adapt to diverse host environments. Hence, analysis of their genomes to study the potential for horizontal gene transfer leading to enhanced infection capabilities and survival abilities across various ecological niches assumes additional significance.

Here, we have investigated a new *E. anophelis* endophthalmitis strain, a multidrug resistant pathogen isolated from a post-operative endophthalmitis patient. After sequencing, CG analysis was performed with twenty-five other *E. anophelis* strains whose genomes were publicly available to get an insight into their phylogenetic position and also to understand the similarities and differences in their gene contents with the aim of determining the unique features of the organism isolated for the first time from an eye infection. Although the genome of the

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The organism exhibited intermediate resistance to tigecycline and ciprofloxacin (Table 1).

## Table 1. Antibiotic susceptibility profile of *E. anophelis* endophthalmitis. Antibiotic susceptibility test was performed using Vitek-2 as per clinical laboratory standards Institute (CLSI) guidelines. MIC – minimum inhibitory concentration of the antibiotic in µg/ml.

| S.no | Antibiotic Class | Antibiotic | MIC (µg/ml) | Resistance profile |
|------|------------------|------------|-------------|--------------------|
| 1    | Penicillins      | Piperacillin/tazobactam | ≥=128       | Resistant          |
| 2    | Cephalosporins   | Ceftazidime | ≥=64        | Resistant          |
| 3    |                  | Cefepime   | ≥=64        | Resistant          |
| 4    | Monobactum       | Aztreonam  | ≥=64        | Resistant          |
| 5    | Carbapenems      | Imipenem   | ≥=16        | Resistant          |
| 6    |                  | Meropenem  | ≥=16        | Resistant          |
| 7    | Aminoglycosides  | Amikacin   | ≥=64        | Resistant          |
| 8    |                  | Gentamicin | ≥=16        | Resistant          |
| 9    | Quinolones       | Ciprofloxacin | 2          | Intermediate Resistant |
| 10   |                  | Levofloxacin | 2          | Sensitive          |
| 11   | Tetracyclines    | Minocyclin | ≤=1         | Sensitive          |
| 12   |                  | Tigecycline | 4          | Intermediate Resistant |
| 13   | Trimethoprim     | Trimethoprim/Sulfamethoxazole | 80 | Resistant          |

organism has overall similarities with the genomes of other *E. anophelis* strains, presence of several unique genes and the ability of horizontal gene transfer indicate its distinct origin.

## Results

### Characteristics of the patient.

The patient was diagnosed with post-operative endophthalmitis (anterior chamber hypopyon, dense vitreous haze with yellow reflex). There was no view of the retina and his vision was only perception of light. Subsequently, vitrectomy was performed to debulk the infection and pus was removed from the infected eye. Empirical treatment for endophthalmitis was immediately initiated.

### Identification and antibiotic susceptibility profile.

The isolate was initially identified as *Elizabethkingia meningoseptica* by Vitek-2. Subsequently, with the availability of additional *Elizabethkingia* genomes and 16s rDNA sequence analysis, the genome of the study isolate was re-identified as *Elizabethkingia anophelis*. MIC analyses of 13 antibiotics revealed that the pathogen was resistant to penicillins, cephalosporins, monobactam, carbapenems, aminoglycosides and trimethoprim/sulfamethoxazole; and sensitive to levofloxacin and minocycline. The organism exhibited intermediate resistance to tigecycline and ciprofloxacin (Table 1).

### Comparative analysis of core and pan genome.

26 genomes of *E. anophelis* including the genome of the new *E. anophelis endophthalmitis* study strain were retrieved from NCBI database for CG analysis. The average size of the genomes is 4.03 Mb and average G+C% is 35.61%. The strain with the maximum size is NUHP1 (4.36 Mb) and the smallest genome is As1 (3.59 Mb) (Table 2). The predicted protein sequences of all the 26 *E. anophelis* genomes were used as input to conduct the core-pan genome analysis. CG analysis revealed that 1404 (40.79%) genes were shared between all the 26 strains, which may be considered to be “core genome”. The accessory genome varied from 656 to 2240 genes (avg. 59.2%) across the strains (Fig. 1). Most notably, exclusion of the study isolate’s genome from the analysis led to an inclusion of an additional 4 genes in the core genome of the remaining 25 genomes. The study strain was found to have maximum number of exclusively absent (846) and unique (156) genes (Table 3), revealing that it has an open pan genome. Annotation of the pan-genome of all the 26 genomes to understand the enrichment of the COG (Fig. 2) and KEGG pathways (Fig. 3) has been mapped.

### Core genome based phylogenetic analysis.

Phylogenetic tree was constructed to infer the relationship among these genomes using the concatenated sequences of 1,404 core proteins identified from the pan/core genome analysis. The tree separated the 26 *E. anophelis* genomes into two distinct branches. In the lower branch, genomes of 502, B2D, Endophthalmitis were found to be closely related. Genomes of *E. anophelis* from Singapore have clustered together (NUH6, NUH11, NUH1, NUHP1, NUHP2, NUHP3) excepting NUH4. Further, the genomes of the four Wisconsin outbreak isolates CSID300521207, CSID3015183678, CSID3015183681 and CSID3015183684 have been found to cluster together. The other branch was found to have two sub clusters. 12012-2PRCM, As1, Ag1, R26, 0422, PW2806 and PW2809 genomes grouped into one, while EM361-97, Po0527107, LDVH-AR107, V0378064, FMS007 and NUH4 genomes were part of the second sub cluster (Fig. 4).

### Resistance to antibiotics and toxic compounds.

Screening of all the study genomes revealed the presence of *bla*GOB, *bla*H and *bla*CMY (excepting strain AS1) beta-lactamase genes in most of the strains. Further, vancomycin (VanW) resistance was predicted among all the 26 strains. A gene encoding bile salt hydrolase - Choloylglycine hydrolase (EC 3.5.1.24) was found in all the analysed genomes excepting As1. This enzyme has been previously reported to protect *Brucella abortus* in the host gut from the toxic and antimicrobial activity of the...
bile salts\textsuperscript{14}. Genes associated with CzcCBA, a membrane bound protein complex aiding in heavy metal resistance\textsuperscript{15} have been identified in all the 26 genomes. Genes coding for proteins conferring resistance to several heavy metals (copper, Zinc, cadmium, cobalt) have been discovered among all the study isolates. In addition, genes coding for the proteins leading to arsenic resistance have been predicted in all the genomes. Further, some of the strains (NUH1, NUH11, NUH4, NUH6, NUHP1, NUHP2 and NUHP3) were found to possess arsenic resistance operon repressor suggesting the inducibility of the system’s resistance. Several genes encoding for the multidrug efflux pumps including RND (CmeB – 26 strains, CmeC – 14 strains), multi antimicrobial extrusion protein (Na\(+\)/drug antiporter – 26 strains)\textsuperscript{16} were found in the study genomes (Supplementary Table 2).

**Putative virulence and anti-virulence genes.** Many genes that may be associated with invasion and intracellular resistance in humans have been identified. We have identified homologs of the gene encoding for an agmatine deiminase in all the 26 isolates. Agmatine deiminase has been reported to aid growth at low pH and biofilm formation, confer acid tolerance in addition to being a potential adherence factor in the colonization of vagina\textsuperscript{17}. Putative hemolysin and a hemolysin secretion protein have been predicted among all the 26 genomes. Hemolysin has been implicated as a virulence factor among several gram-negative and gram-positive pathogens\textsuperscript{18}. It has also been reported that hemolysin could be a potential ocular virulence factor in *Bacillus cereus* and *Staphylococcus aureus* leading to endophthalmitis\textsuperscript{19,20} and keratitis\textsuperscript{21} respectively. However, among the 26 genomes included in this study, arylsulfatase has been identified only in the current study isolate. Arylsulfatase has been implicated previously in *E. coli* infection of the brain microvascular endothelial cells (BMEC) of the host. Presence of arylsulfatase may contribute to the ability of the pathogen to cross the blood-brain barrier leading to meningitis\textsuperscript{17,22}. An operon consisting of Quinolinate synthetase [EC 2.5.1.72], L-aspartate oxidase [EC 1.4.3.16], and quinolinate phosphoribosyl transferase [EC 2.3.2.19] involved in quinolinate biosynthesis has been identified in all the genomes being analysed (Supplementary Table 3). These genes are considered to have potential anti-virulence function as their activation was reported to inhibit invasion and intracellular spread among *Shigella* species\textsuperscript{23}.

**Analysis of Mobile Genetic Elements (MGEs).** MGEs are the major contributors for Horizontal Gene Transfer (HGT). A total of 59 prophage related regions have been identified across the study genomes. Of these, 56 of them appear to be incomplete prophage genomes in 23 study isolates. The other three strains (422, PW2806 and PW2809) were predicted to contain questionable prophage genomes. The size of the prophage genomes varied from 5.6 kb to 37.4 kb. Strains NUH1, NUH4, NUHP1, NUHP2 and NUHP3 were found to harbor maximum

| Strain Name      | Year of Collection | Accession id        | Genome size in Mb | GC %  | No. of Contigs | Genes | Proteins |
|------------------|--------------------|---------------------|-------------------|-------|----------------|-------|----------|
| 0422             | 1950               | NZ_LNOG01000011     | 3.9599            | 35.6  | 26             | 3634  | 3548     |
| 502              | 2012               | NZ_AVCO01000001     | 3.96066           | 35.5  | 21             | 3617  | 3520     |
| 12012-2PRCM      | 2009               | NZ_LPGX01000011     | 4.02331           | 35.6  | 83             | 3671  | 3554     |
| Ap1              | 2010               | NZ_AHHG01000001     | 4.04571           | 35.5  | 51             | 3723  | 3572     |
| As1              | 2013               | NZ_LKFO01000006     | 3.59087           | 35.5  | 12             | 3303  | 3229     |
| B2D              | 2013               | NZ_INCG01000007     | 3.93625           | 35.5  | 50             | 3553  | 3473     |
| CSD0_3000521207  | 2016               | NZ_CP015067         | 3.85345           | 35.7  | 1              | 3490  | 3390     |
| CSD0_3015183678  | 2016               | NZ_CP014805         | 3.93122           | 35.8  | 1              | 3562  | 3461     |
| CSD0_3015183681  | 2016               | NZ_CP015068         | 3.93122           | 35.8  | 1              | 3563  | 3461     |
| CSD0_3015183684  | 2016               | NZ_CP015066         | 3.93122           | 35.8  | 4              | 3562  | 3458     |
| EM361-97         | 2010               | NZ_KV757122         | 4.08405           | 35.7  | 27             | 3729  | 3614     |
| Endophthalmitis  | 2014               | JSA01000100         | 4.01982           | 35.5  | 167            | 3729  | 2302     |
| FMS-007          | Not available      | NZ_CP006576         | 3.93897           | 35.6  | 1              | 3578  | 3470     |
| LDVH-AR107       | 2004               | NZ_FTPG01000001     | 3.98893           | 35.7  | 108            | 3658  | 3538     |
| NUH1             | 2012               | NZ_ASYH01000001     | 3.43466           | 35.6  | 59             | 3993  | 3879     |
| NUH11            | 2012               | NZ_ASYK01000005     | 4.09148           | 35.6  | 59             | 3757  | 3642     |
| NUH4             | 2012               | NZ_ASYI01000001     | 4.23949           | 35.6  | 50             | 3912  | 3811     |
| NUH6             | 2012               | NZ_ASYJ01000011     | 4.1238            | 35.6  | 74             | 3812  | 3696     |
| NUHP1            | 2012               | NZ_CP007547         | 3.46983           | 35.6  | 1              | 4016  | 3898     |
| NUHP2            | 2012               | NZ_ASYF01000003     | 3.3465            | 35.5  | 59             | 3988  | 3882     |
| NUHP3            | 2012               | NZ_ASYG01000009     | 3.44311           | 35.5  | 71             | 3985  | 3871     |
| Po0327107        | 2006               | NZ_CCAC01000008     | 4.03206           | 35.5  | 89             | 3674  | 3573     |
| PW2806           | 2012               | NZ_CBVD01000003     | 3.91281           | 35.9  | 388            | 3612  | 3456     |
| PW2809           | 2012               | NZ_CBVE01000002     | 3.92215           | 35.8  | 278            | 3598  | 3452     |
| R26              | 2006               | NZ_ANIW01000066     | 4.03272           | 35.4  | 66             | 3726  | 3633     |
| V0378064         | 2011               | NZ_CCAB010000170    | 4.03675           | 35.7  | 214            | 3648  | 3556     |

Table 2. Comparison of genome characteristics of the 26 *E. anophelis* isolates used in analysis.
number of prophage regions (4 numbers) compared to the others. The average G + C% among these prophage genomes was 34.55% compared to the average G + C% of the study genomes (35.62%) (Supplementary Table 4).

A total of 107 Genomic Islands (GIs) have been identified in 25 of the study genomes excepting B2D. The smallest of the predicted GI was 8.2 kb in the strain EM361-97 and the largest GI was 31.8 kb in NUH1 and NUHP3 strains. NUH1, NUH4 and NUHP2 were found to harbor 8 GIs. Maximum number of coding DNAs (66 numbers) were found in the GI (region III) of the strain 422. Several virulence factors, antibiotic resistance genes, pathogenicity islands, insertion sequences, prophage related genes and genes of the secretion systems have been identified in these GIs (Supplementary Table 5). Further, remnants of several Integrative and Conjugative Elements (ICEs) have been found in many of the study genomes.

Type VI and Type IV Secretion systems. T6SS plays an important role in bacterial pathogenesis by allowing the transport of virulence factors, targeting the host cells as well as helping in competing with other bacteria in their niche24. They are widely distributed in the genomes of the phylum Bacteroidetes of which Flavobacteriaceae is a family. More specifically, T6SSiii has been reported to be prevalent among the members of the Flavobacteriaceae25. Consistent with earlier reports25-26, several of the study genomes were found to possess T6SSiii. Genes annotated to TssN, TssO and TssP proteins which are unique to only T6SSiii along with those coding for the other core components such as TssB, TssC and extracellular components VgrG and HcP have been identified in twenty-four genomes. Only strains NUH6 and As1 did not appear to harbor any genes coding for the T6SS components (Supplementary Table 6).

T4SS are established components of bacterial conjugation and virulence. T4SS genes are also acquired as part of the Integrative and Conjugative Elements (ICEs). Our analysis revealed that 23 of the genomes possessed the genes associated with T4SS but are absent in strains As1, B2D and R26 (Supplementary Table 7).

Defence and repair systems. Bacteria employ a host of mechanisms to protect themselves from the invading genetic elements. These include Restriction Modification systems (RMs) which are considered to be innate immune systems and Clustered Regularly Interspaced Short Palindromic repeat sequences (CRISPRs), considered to be adaptive immune systems27. Our analysis revealed the presence of Type I and Type II RM systems.
Unlike all other genomes analyzed in the study, *E. anophelis* endophthalmitis was found to possess maximum number of RM system associated genes. The genome was found to possess 12 RM genes, while the strains 502 and B2D did not possess any genes coding for RM systems (Supplementary Table 8). Analysis for CRISPRs indicated that only four of the study genomes (FMS007, LDVH-AR107, Po0527107 and V0378064) possess confirmed CRISPRs (Supplementary Table 9). Analysis for the presence of anti-restriction systems28 led to the detection of an anti-restriction gene ArdA among 19 of the study genomes. ArdA protein was reported to support the MGEs in evading the Type I RM systems and augment the spread of resistance determinants29 (Supplementary Table 10).

CG analysis indicated that most of the DNA repair pathways are represented in all the study genomes. Most of the repair pathways appear to be intact. In the majority of the genomes excepting those of the Wisconsin strains, there was no disruption in the *mutY* (Adenine DNA glycosylase) gene. In the genomes of the four Wisconsin strains, the 1,029 bp *mutY* gene was found to be disrupted by the insertion of a 62,212 bp ICE *Ea1* (Supplementary Fig. 1)13.

A total of 32 protein coding genes (excluding hypothetical genes) involved in transposition, excision of the conjugal transposon, heavy metal resistance and tetracycline resistance have been found inside the ICE. *E. anophelis* endophthalmitis is naturally competent.

Although a number of *Elizabethkingia* strains have been identified and characterized, natural transformation has not yet been reported in any of them. Given the presence of a considerable number of GIs and other gene clusters possibly acquired through HGT in the strain, we investigated the capability of the organism to carry out HGT. In the absence of any well characterized bacteriophages for these group of bacteria, we resorted to study natural transformation.

Natural transformation was observed after exposing plasmid DNA to exponentially growing cells. PCR analysis for the presence of BDNF gene in the plasmids isolated from the transformants confirmed that *E. anophelis* is naturally competent (Fig. 5). However, natural competence was observed only when OD₆₀₀ reached 0.84. Genome analysis revealed three genes – (a) DNA internalization-related competence protein ComEC/Rec2, (b) Competence protein F homolog and (c) Competence/damage-inducible protein CinA involved in DNA internalization and transformation. These genes are present in majority of the analyzed genomes indicating that natural transformation may be occurring in other *Elizabethkingia* (Supplementary Table 11).

Table 3. Core, accessory, unique and exclusively absent genes in the 26 *E. anophelis* genomes after pan-core genome analysis using BPGA pipeline. Core genes – number (No.) of genes that are shared by all the study genomes, Accessory genes – Genes that are not shared by all the genomes, Unique genes – genes that are found exclusively in a particular genome, exclusively absent genes – genes that are exclusively absent in a particular genome but are otherwise found in the other genomes.

| Strain Name       | No. of core genes | No. of accessory genes | No. of unique genes | No. of exclusively absent genes |
|-------------------|-------------------|------------------------|---------------------|--------------------------------|
| 0422              | 1404              | 1950                   | 94                  | 3                              |
| 502               | 1404              | 1888                   | 108                 | 2                              |
| 12012-2PRCM       | 1404              | 1930                   | 131                 | 10                             |
| Ag1               | 1404              | 2010                   | 10                  | 30                             |
| As1               | 1404              | 1693                   | 5                   | 149                            |
| B2D               | 1404              | 1868                   | 101                 | 5                              |
| CSID207           | 1404              | 1896                   | 0                   | 27                             |
| CSID678           | 1404              | 1958                   | 0                   | 0                              |
| CSID681           | 1404              | 1959                   | 0                   | 1                              |
| CSID684           | 1404              | 1956                   | 0                   | 0                              |
| EM361-97          | 1404              | 1964                   | 113                 | 3                              |
| Endophthalmitis   | 1404              | 656                    | 156                 | 846                            |
| FMS007            | 1404              | 1913                   | 56                  | 6                              |
| LDVH-AR107        | 1404              | 2006                   | 21                  | 16                             |
| NUH1              | 1404              | 2240                   | 0                   | 0                              |
| NUH4              | 1404              | 2185                   | 22                  | 0                              |
| NUH6              | 1404              | 2107                   | 33                  | 5                              |
| NUH11             | 1404              | 2097                   | 12                  | 1                              |
| NUH1P1            | 1404              | 2236                   | 0                   | 0                              |
| NUH1P2            | 1404              | 2236                   | 0                   | 0                              |
| NUH1P3            | 1404              | 2238                   | 2                   | 0                              |
| Po0527107         | 1404              | 2032                   | 45                  | 0                              |
| PW2806            | 1404              | 1956                   | 3                   | 12                             |
| PW2809            | 1404              | 1948                   | 5                   | 13                             |
| R26               | 1404              | 2052                   | 38                  | 0                              |
| V0378064          | 1404              | 2035                   | 20                  | 0                              |
Discussion
In this manuscript, we have described the uncommon features of a new strain of *E. anopheles* isolated from a post-operative endophthalmitis patient. This is the first ever isolate of the species from a patient suffering from this infection. Initial Vitek-2 analysis from the vitreous fluid of the patient led to the identification of the organism as *Elizabethkingia meningoseptica*. Availability of several *Elizabethkingia* genomes due to the advent of whole genome sequencing and 16srDNA analysis lead to the unambiguous identification of the pathogen as *Elizabethkingia anophelis*, a closely related species to *E. meningoseptica*. Due to the unusual nature of the pathogen, an investigation of the patient’s eye drops, sinks and cubicle in the ward where the patient stayed was conducted for the presence of the pathogen. However, culture results did not indicate the presence of *E. anophelis* ruling out the possibility of nosocomial acquisition. This raised the possibility that alternate modes have been employed in the transmission of the bacterium leading to postoperative endophthalmitis. Our findings suggest that *E. anophelis* is a slow growing bacterium compared to *E. coli* and is capable of natural transformation during a narrow window of exponential phase of growth.

CG analysis with 25 other sequenced strains of *E. anophelis* showed several differences. Pan and core genome analysis revealed that the strain *E. anophelis* endophthalmitis has undergone massive gene rearrangements indicated by the high number of unique and exclusively absent genes predicted in its genome. Exclusion of *E. anophelis* endophthalmitis genome from the pan-core genome analysis has led to an increase in the core genome size by 37% indicating the strain’s divergence. The presence of a large number of MGEs and the horizontally acquired genes would have contributed to the genome rearrangement in the organism, diverging it from other analyzed genomes. These data along with others described below suggest the possibility of the recent emergence of the strain.

Survival in diverse environments such as mosquito mid gut and human tissues necessitates the bacterium to adapt to the respective niche environments. In such a scenario, possession of MGEs would help adjust and thrive. Analysis for MGEs revealed the presence of GIs, remnants of phage genomes and ICEs amongst several of the study genomes, indicating that the organisms with incomplete prophages and ICEs have further undergone gene gain/loss which may be of an evolutionary requirement for the pathogen to be successful in diverse ecological niches. Interplay between the bacterial defence systems (RMs, CRISPRs) and anti-RM proteins are known to
decide the fate of the cell towards or against HGT (Supplementary Table 12). Taken together, these observations highlight the potential of the organism to be involved in more robust HGT. The presence of T6SS and T4SS secretome further confirms that HGT is a norm and not an exception in *E. anophelis*. Given this scenario, it is not surprising to find several genes associated with antibiotic resistance, efflux pumps, virulence factors and competence. The finding of an exclusive virulence gene, arylsulfatase in the organism along with other genes with potential for pathogenesis and disease in humans, may provide further insights into the adaptation mechanisms of the pathogen to thrive under diverse ecological niches.

Notably, the virulence factor arylsulfatase with potential to cause meningitis was not found in any of the other analysed genomes including those from the central african republic outbreak (Po0527107 and V0378064) associated with neonatal meningitis. This indicates that the current study strain may also have the potential to cause meningitis and there may be other meningital virulence factors that are yet to be identified and characterized.

From the features described, it is evident that *E. anophelis* endophthalmitis is a novel human pathogen with distinct characteristics. It is tempting to speculate how the organism could have been transmitted possibly by the mosquito vector to cause eye infection hitherto not associated with *E. anophelis*. Different mosquito species are highly prevalent in India, Africa and elsewhere. Thus, it is not unrealistic to suggest the transmission of the pathogen to human hosts by the mosquito vector in whose gut *E. anophelis* was first discovered. We speculate that the mosquito may have bitten near the eye after the patient has undergone surgery, thus transmitting the bacterium into the host's ocular system. However, there could be alternate explanations which could account for the transmission. Also, further investigations are needed to confirm the transmission routes including zoonotic transmission of this enigmatic pathogen.

Given the enhanced antimicrobial resistance observed in *E. anophelis*, and its capacity to thrive in many ecological habitats it is very critical to implement best healthcare practices when in contact with the pathogen and initiate appropriate surveillance measures before the pathogen gets involved in the next outbreak.

**Methods**

**Case presentation and strain characterization.** A 67-year-old patient with complaints of redness, pain and loss of vision in the right eye after cataract surgery was admitted to the Department of Ophthalmology, Sri Sathyai Sai Institute of Higher Medical Sciences, Prasanthigram, India. *E. anophelis* was isolated from the vitreous fluid of the infected eye. Identification, antibiotic susceptibility testing (AST N281 card) and determination of Minimum Inhibitory Concentration (MIC’s) were performed using Vitek 2 (BioMérieux, France) as per CLSI (Clinical and Laboratory Standards Institute) guidelines.
Natural transformation. *E. anophelis* endophthalmitis cultures were harvested at an OD600 of 0.37, 0.54, 0.61, 0.75 and 0.84. A plasmid (pCMV6-BDNF) containing eukaryotic Brain Derived Neurotrophic Factor (BDNF) with an ampicillin selection marker was added at 0 ng/mL, 28.2 ng/mL, 56.4 ng/mL and 84.6 ng/mL concentrations and

**Figure 4.** Core Phylogenetic tree: The core phylogenetic tree is based on the protein sequences of the concatenated core genes (1404) of the 26 *E. anophelis* genomes. The tree was generated by Neighbor Joining Method in MUSCLE using default parameters. The tree was plotted using the Perl package Bio::Phylo::Treedrawer available at http://search.cpan.org/~rvosa/Bio-Phylo-v2.0.1/lib/Bio/Phylo/Treedrawer.pm as implemented in BPGA pipeline. The scale bar presented at the foot of the tree indicates time period in millions of years (MYA).

**Figure 5.** PCR confirming the natural transformation in *E. anophelis* endophthalmitis. Lane 1: 1 kb DNA ladder. Lane 2: Positive control (pCMV-6 BDNF plasmid used as template). Expected band size of BDNF gene is 750 bp. Lane 3–5: PCR products from plasmid isolated from *E. anophelis* endophthalmitis after natural transformation. Lane 6–7: Negative controls.
incubated for 90 minutes at 30°C. Untransformed E. anophelis endophthalmitis was found to exhibit resistance to ampicillin (250μg/mL). Hence, transformation mixtures were plated on LB-amp agar plates at a higher concentration (600μg/mL) in three dilutions, i.e., 1:2, 1:20 and 1:100. PCR was performed for the presence of BDNF gene in the plasmids isolated from the transformed colonies using specific primers (forward Primer: 5'-GGATCCATG ACCATCCTTTTCTATGAGG-3'; reverse Primer: 5'-AAGCTTCTATCTTCCCTTTTAATGGTCAGT-3') in a final volume of 20μl containing 10μl of PCR Master Mix (Takara) which includes dNTPs, MgCl2, Tag DNA polymerase and PCR buffer, 0.5μM of forward and reverse primers, template DNA (2μl) and nuclease free Water (4μl). The PCR conditions employed for the amplification of BDNF gene were 94°C - 2 minutes followed by 30 cycles of 94°C - 15 seconds, 55°C - 30 seconds, 72°C - 47 seconds and a final elongation at 72°C - 7 minutes. PCR amplicons were analysed on 1.2% agarose gel. E.coli DH5α and untransformed E. anophelis endophthalmitis served as negative controls.

**Growth kinetics.** To determine the growth curve of E. anophelis endophthalmitis, 1% inoculum from overnight culture was added to 100 ml of fresh LB medium and incubated at 37°C. Absorbance at 600 nm was measured at every 30 minutes. Growth kinetics revealed that the generation time of the isolate is 78 minutes at 37°C (Supplementary Fig. 2).

**Genome characterization.** Draft genome sequencing and assembly of E. anophelis endophthalmitis genome was recently reported. The contigs from the draft assembly were subjected to gene prediction using PRODIGAL tool with default parameters as recommended. Predicted protein sequences were annotated using BLASTp against UNIPROT bacterial proteins database with evalue < 0.001, > 70% as query coverage and %Identity > 30. E. anophelis NUHPI (gi Number: 675102482) genome was used to construct the genome map of E. anophelis endophthalmitis. This led to successful organization of 111 contigs out of 167 contigs. A total of 3,729 ORFs encoding 2302 proteins were predicted from the assembled genome. All available (fifty-five other) E. anophelis genome sequences (as on February 02, 2017) were obtained from NCBI Genomes database for CG analysis and RAST (version 2.0) annotation has been repeated to obtain unambiguous results (Supplementary Table 1).

**Bacterial Pan Genome Analysis (BPGA)** was used for comprehensive pan/core genome analysis, functional annotation of the core, accessory and unique genes to Cluster of Orthologous groups (COG) categories and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways using default parameters. Phage genomes were identified by PHASTER. Antibiotic resistance genes were predicted by RAST (version 2.0), Resfinder (version 3.0) and VRprofile (version 2.0). Virulence factors, genomic islands, Insertion sequences and T6SS Secretory systems were analyzed by VRprofile (version 2.0). TASS genes were identified by SecReT4 (version 1.0). ICEberg (version 1.0) was used to screen for ICEs. Restriction Modification (RM) and anti-restriction systems were identified by RAST (version 2.0). CRISPRfinder was used to predict potential CRISPR gene clusters. Unless otherwise mentioned, all the above mentioned analyses were performed using default parameters.

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