



Genome Sequences of Four *Vibrio parahaemolyticus* Strains Isolated from the English Channel and the River Thames

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ABSTRACT *Vibrio parahaemolyticus* is the lead causative agent for seafood-borne human gastroenteritis. While its occurrence has traditionally been uncommon in Europe and the United Kingdom, rising sea surface temperatures have resulted in an increased prevalence. Here, we present the complete genome sequences of four novel *V. parahaemolyticus* strains isolated in the United Kingdom.

Vibrio parahaemolyticus is a ubiquitous marine bacterium and an important causative agent for human gastroenteritis (1). It is found in seawaters where temperatures exceed 15°C (2); thus, *V. parahaemolyticus* abundance and high rates of infection have traditionally been associated with outbreaks in Asia, Africa, South America, and the United States (3–5). However, rising sea surface temperatures have resulted in an increased prevalence in Europe and the United Kingdom (6, 7). In the midst of a warming climate, it is important to map the genetic profile of *V. parahaemolyticus* to better understand how to prevent and treat human infection. Here, we present the genome sequences for four UK environmental *V. parahaemolyticus* isolates, isolated as follows: (i) EXE V18/004 from *Ostrea edulis* at Chichester Harbor (2018), (ii) V12/024 from *Crassostrea gigas* at Weymouth (2012), (iii) V05/313 from *Eriocheir sinensis* in the River Thames (2005), and (iv) V05/027 from an unknown shellfish source in Southampton (2005). Strains EXE V18/004 and EXE V13/004 were isolated at the University of Exeter, while strains V12/024 and V05/027 were isolated at Cefas Weymouth Laboratories. All four of these strains were isolated directly from shellfish following previously described methods (2, 8). Strain V05/027 was donated to Exeter University from Cefas Weymouth laboratories.

V. parahaemolyticus was initially identified based on colony morphology on selective agar (marine agar and thiosulphate citrate bile sucrose agar) and by PCR targeting the *toxR* region (9). Isolates were grown in marine broth (10) at 37°C overnight, and 1 ml of each culture ($\sim 10^9$ cells · ml⁻¹) was added directly to a Qiagen DNeasy PowerWater kit (Germany) for DNA extraction, followed by library preparation for short-read (2 × 250 bp; Illumina, San Diego, CA) and long-read (MinION; Oxford Nanopore Technologies) sequencing. For short-read libraries, DNA was quantified in triplicate with the Quant-iT double-stranded DNA (dsDNA) high-sensitivity (HS) assay in an Eppendorf AF2200 plate reader. Genomic DNA libraries were prepared using a Nextera XT library prep kit (Illumina) following the manufacturer's protocol with the following modifications: 2 ng of DNA instead of 1 ng was used as the input, and PCR elongation time was increased from 30 s to 1 min. DNA quantification and library preparation were carried out on a Hamilton Microlab Star automated liquid handling system. Pooled libraries were quantified using the Kapa Biosystems library quantification kit for Illumina on a Roche LightCycler 96 quantitative PCR (qPCR) machine. Libraries were sequenced on an Illumina HiSeq instrument. Long-read sequencing was performed in-house using a multiplexed SQK-LSK108 library preparation and sequenced on a FLO-MIN106 flow cell, following Oxford Nanopore

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TABLE 1 Chromosome length, sequence run statistics, and assembly statistics for the *Vibrio parahaemolyticus* strains described in this study

<i>V. parahaemolyticus</i> strain	No. of contigs	Chromosome length (chromosome 1; chromosome 2 [bp])	No. of MinION reads	Median MinION read length (bp)	Median short-read coverage (chromosome 1; chromosome 2 [×])	Median long-read coverage (chromosome 1; chromosome 2 [×])	G+C content (%)	No. of protein-coding genes
EXE V18/004	5	3,263,543; 1,747,363	241,527	1,943	38; 30	187; 151	45	4,524
V12/024	2	3,315,999; 1,877,872	18,595	17,487	27; 22	61; 58	45	4,604
V05/313	3	3,319,614; 1,940,186	1,116,682	1,173	53; 42	784; 636	45	4,738
V05/027	2	3,438,892; 1,705,150	168,588	3,832	62; 51	135; 111	45	4,669

Technologies protocol (11). Short reads were adapter trimmed using Trimmomatic v3.0 (12) with a sliding window quality cutoff of Q15. Long-read sequences were base called using the Guppy v2.3.5 FlipFlop algorithm and then demultiplexed and adapter trimmed using Porechop (<https://github.com/rrwick/Porechop>) with the following settings: -require_two_barcode -discard_unassigned -discard_middle. Hybrid genome assembly was performed using Unicycler v0.4.7 (13), and each assembly was uploaded to the Integrated Microbial Genomes (IMG) platform (14), developed by the Joint Genome Institute (JGI; USA) for annotation. Short-read coverage was calculated using BMap v38.22 (<https://sourceforge.net/projects/bbmap/>) with the following settings: idfilter=0.95 covstats=covstats.txt. Long-read coverage was calculated by first using minimap2 v.2.17 (<https://github.com/lh3/minimap2>) and SAMtools v.1.9 (<http://samtools.sourceforge.net/>) to create a bam file (minimap2 -t 16 -ax map-ont <assembly> <long_reads> | samtools view -F 4 -buS | samtools sort -o long.sorted.bam) and then using BMap's pileup.sh to calculate coverage (pileup.sh in=long.sorted.bam ref=ref.fa out=long.covstats.txt). Default parameters for all software were used unless otherwise noted.

Each strain was found to have two chromosomes (Table 1), one with an average size of 3.3 Mbp and a smaller one with an average size of 1.8 Mbp, which is consistent with the literature (15). Two strains, EXE V18/004 and V05/313, were found to contain plasmids with sizes of 49,878 bp and 71,936 bp, respectively.

Data availability. Assembled and annotated genomes are publicly available from JGI IMG/M (<https://img.jgi.doe.gov/>) using the following taxon identifiers (IDs): [2816332655](https://img.jgi.doe.gov/taxonomy/term/2816332655) (*V. parahaemolyticus* EXE V18/004), [2816332656](https://img.jgi.doe.gov/taxonomy/term/2816332656) (*V. parahaemolyticus* V12/024), [2816332657](https://img.jgi.doe.gov/taxonomy/term/2816332657) (*V. parahaemolyticus* V05/313), and [2816332658](https://img.jgi.doe.gov/taxonomy/term/2816332658) (*V. parahaemolyticus* V05/027). Read data are available from the European Nucleotide Archive under the following strain names and accession numbers: *V. parahaemolyticus* EXE V18/004, [ERS3342146](https://www.ebi.ac.uk/ena/record/ERS3342146); *V. parahaemolyticus* V12/024, [ERS3342147](https://www.ebi.ac.uk/ena/record/ERS3342147); *V. parahaemolyticus* V05/313, [ERS3342148](https://www.ebi.ac.uk/ena/record/ERS3342148); and *V. parahaemolyticus* V05/027, [ERS3342149](https://www.ebi.ac.uk/ena/record/ERS3342149).

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