Complete Genome Sequences of Seven *Vibrio anguillarum* Strains as Derived from PacBio Sequencing

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Abstract

We report here the complete genome sequences of seven *Vibrio anguillarum* strains isolated from multiple geographic locations, thus increasing the total number of genomes of finished quality to 11. The genomes were de novo assembled from long-sequence PacBio reads. Including draft genomes, a total of 44 *V. anguillarum* genomes are currently available in the genome databases. They represent an important resource in the study of, for example, genetic variations and for identifying virulence determinants. In this article, we present the genomes and basic genome comparisons of the 11 complete genomes, including a BRIG analysis, and pan genome calculation. We also describe some structural features of superintegrons on chromosome 2s, and associated insertion sequence (IS) elements, including 18 new ISs (ISVα3 – ISVα20), both of importance in the complement of *V. anguillarum* genomes.

Key words: *Vibrio anguillarum*, chromosomal integrons, integrases, insertion sequences, IS-elements, PacBio sequencing.

Introduction

*Vibrio (Listonella) anguillarum* is a marine bacterium and the causative agent of hemorrhagic septicemia (or vibriosis), in fish, molluscs, and crustaceans (Frans et al. 2011). The pathogenic nature of *V. anguillarum* and its global impact on the aquaculture industry continues to keep this bacterium in the spotlight. In efforts to elucidate virulence determinants and/or to analyze genetic variations among strains, 44 genome sequences have been determined (Agarwala et al. 2018).

Recently, Holm et al. (2015) reported the complete genome of the virulent strain NB10, originally isolated from diseased rainbow trout (*Oncorhynchus mykiss*) on the Swedish coast of the Gulf of Bothnia. Its genome, which is typical in size (average 4.31 Mb), is 4,373,835 bp in total, and consists of two circular chromosomes and a pJM1-like plasmid named p67 (66.8 kb). This is 255 kb larger than the genomes of strains 775 and M3, which were published in 2011 and 2013, respectively (Naka et al. 2011; Li et al. 2013). The majority of the 255-kb DNA represent prophages, genomic islands, and genes of unknown function/hypothetical protein genes (Holm et al. 2015). Strain 775 was isolated from Coho salmon (*Oncorhynchus kisutch*) on the United States Pacific coast, and strain M3 was isolated from Japanese flounder (*Paralichthys olivaceus*) off the coast of China. Another previously available complete genome includes that of strain 90-11-286 (Castillo et al. 2017). The initially complete strains NB10, 775, and M3 all harbor a pJM1-like plasmid, strain 90-11-286 has no plasmid.

Materials and Methods

Bacterial Isolates

*Vibrio anguillarum* strains 87-9-116, JLL237, S3 4/9, CNEVA NB11008, VI843, and VI812 were kindly provided by Prof. Hans Rediers (KU Leuven Association, Sint-Katelijne-Waver, Belgium). Strain ATCC-68544 (synonym 775) was acquired from the ATCC Bacteriology Collection. Bacteria were routinely grown at 22°C on BD Difco Marine Agar 2216 (Fisher Scientific) and in liquid cultures in BD Bacto Tryptic Soy Broth (Fisher Scientific).

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DNA Isolation and DNA Sequencing

Total DNA was isolated from 6 ml overnight cultures at stationary phase using Genomic-tip 100/g (Qiagen) according to the manufacturer protocol. The final DNA concentration and quality were measured using a Nanodrop 2000c (Thermo Scientific) instrument. Integrity of high-molecular weight DNA was examined on a 1% agarose gel. DNA samples were sequenced at the Norwegian Sequencing Centre (NSC: a national sequencing core facility located in Oslo).

Genome Analysis

SMRT sequencing was performed at NSC. Libraries were constructed using Pacific Biosciences 20-kb library preparation protocol. Size selection of the final library was performed using BluePippin with a 7-kb cut-off. Libraries were sequenced on Pacific Biosciences RS II instrument using P6-C4 chemistry with 360-min movie time. Reads were assembled using HGAP v3 (Pacific Biosciences, SMRT Analysis Software v2.3.0). Contigs were circularized using Minimus2 software of Amos package (Schatz et al. 2013).

For CNEVA NB11008 and JLL237, the number of circular contigs were bioinformatically corrected. The BRIG software (Alikhan et al. 2011) was used to compare the 11 complete genomes (with the NB10 strain chromosomes as a reference). The genomes of ATCC-68544 and 775 were globally compared using the Artemis Comparison Tool (ACT), and the comparison file was produced with the DOUBLE ACT v.2 server (Carver et al. 2005).

Results and Discussion

As of March 2018, 11 V. anguillarum genomes of finished quality are available in the genome databases (table 1). A rough overview of the location at which these strains originate is shown in supplementary figure S1, Supplementary Material online (although exact geographical positions for most of them are unavailable). All strains originate from Europe, except 775/ATCC-68544 (from the United States Pacific coast) and M3 (from China).

The sequencing statistics for strains sequenced in this study are shown in supplementary table S1, Supplementary Material online. The complete genomes were assembled from Pacific Biosciences (PacBio) sequence reads (32,981–139,094) produced at the Norwegian Sequencing Centre (NSC). These sequences produced 84.4–207.8 genome coverage, and assembled into circular contigs (i.e., chromosomes and plasmids). The total genome sizes ranged from 4.14 to 4.89 Mb, with an average GC content of 44.4%.

Figure 1 shows a global BLAST comparison of the 11 complete genomes generated using the BRIG software (Alikhan et al. 2011). BLAST matches of sequences from each strain were mapped onto Chromosome 1 and Chromosome 2 of NB10 (i.e., the reference). Overall, the figure shows that the majority of sequences present in NB10 are also present in all others strains. However, the BRIG analysis does not display sequences not found in NB10. To add more information we therefore calculated the pan genome using the GET_HOMOLOGUES tool (Contreras-Moreira and Vinuesa 2013). The orthoMCL algorithm was used with default
parameters and the 11 complete genomes as input. In brief, the numbers from GET_HOMOLOGUES suggest that the pan genome include 7,667 gene clusters in total, 2,574 core clusters, 2,183 accessory clusters, and finally 2,910 unique clusters. This clearly demonstrates that the total number of genes greatly exceeds the number of genes in each genome (complete genomes contain between 3,426 and 4,127 genes).

Moreover, figure 1 shows that the genome gaps (GGs) B, C, and E–J, that have previously been described as sequences present in NB10, but not in 775 and M3 (Holm et al. 2015), are also missing from the majority of strains in the current analysis. However, NB10 sequences in GGs A and D are present in most strains. In general, sequences located in the GGs represent hypothetical CDSs, genomic islands, or prophages. Other GGs are also present, but will not be described in further detail in this work. Finally, based on the BRIG analysis, and as the only complete strain, 87-9-116 appears to contain all (or close to all) CDSs that are present in NB10. A likely explanation relies on the fact that both strains have been sampled from relatively close geographical locations in the Gulf of Bothnia, well adapted to the environment and local biotic factors, even though from different salmonid species.

Notably, according to the ATCC Bacteriology Collection, the strain names ATCC-68554 and 775 are synonymous. To verify this relationship ATCC-68554 was acquired directly from the ATCC Bacteriology Collection, cultured under standard conditions, and finally sequenced. A global pairwise genome comparison of the two genomes was done using the Artemis Comparison Tool (ACT) and the DOUBLE ACT v.2 server (Carver et al. 2005; see supplementary fig. S2, Supplementary Material online). This shows that both Chromosome 1 and Chromosome 2 sequences are highly similar, except for a region located approximately between positions 470,000–597,000 on Chromosome 2 (according to the ATCC-68554 sequence). This region represents a so-called “superintegron” (SI), which normally begins with an integron integrase, but in ATCC-68554 it is truncated and thus nonfunctional (locus tag CLI14_17245). AttC-containing regions (i.e., the SIs) are marked in yellow in supplementary figure S2C, Supplementary Material online. In ATCC-68554 this sequence is 54 kb longer than that of 775. In 775, the SI is followed by a 26-kb region, which is not present in ATCC-68554. The latter 775-specific sequence contains CDSs of various functions, and one tRNA-Gly gene. These discrepancies may be explained by technical artefacts during sequencing and assembly, or by real differences in the genomes, perhaps as a result of subculturing of the bacterium in different laboratories. The SIs and associated insertion sequences (ISs) are described in more detail below.

SIs are subsets of chromosomal integrons (CI) found in vibrios and a wide range of other gram negative bacterial species (for review, see, Cambray et al. 2010). Integrons contain a functional platform (i.e., the integrase encoding gene, intI, a primary integration site, attI, and a primary promoter,
administering integrated gene cassettes [i.e., ORF(s)] followed by a recombination site attC. Superintegron attC sites are species specific (Mazel 2006; Cambray et al. 2010), with a high degree of identity and a common set of characteristics that enable them to be identified, which is the reason why we focused on these cassette components as SI-markers. The supplementary figure S3, Supplementary Material online, shows an alignment of attC-sites from the serovar O1 NB10 strain, with a consensus 31-nt “cassette-identifier”: 5′-TAACAAACGnnTCAAGAGGGAnGnCACC-3′. This cassette-identifier constitutes part of the quality assurance system in the final assembly of the genomes, enabling calculations of the number of cassettes within the SI-part of chromosome 2 s. The SI gene content (attC-span) of finished genomes varies relative to chromosome 2-sizes, between 6.9% in strains 775 and M3 (harboring equally sized and the smallest chromosome 2 s, both with 64 attC-sites) and 28.9% in strain S3 4/9 (containing 147 attC-sites/cassettes). Worth mentioning in this context is the low number of attC-sites in the published partially complete genomes (span: 1–46; average: 22 cassette identifiers; see supplementary table S2, Supplementary Material online), most likely due to their missing genes (cassettes).

Vibrio anguillarum CIs harbor clusters of highly diverse gene cassettes (VAR; Vibrio anguillarum repeats), mostly of unknown function, but among others toxin/antitoxin cassettes and genes involved in substrate modification or interactions with virulence factors and DNA modification, similar to in Vibrio cholerae (Rowe-Magnus et al. 2003).

Also embedded in V. anguillarum genomes, and especially within SIs, are numerous insertion sequences (IS: i.e., transposases, and sometimes one or two accessory genes). The V. anguillarum SIs encode a specific integrase denoted VangIntIA (based on the naming of VchIntIA, a specific integrase in V. anguillarum [O1] El Tor strain N16961; Mazel et al. 1998).

A striking observation is that the VangIntIA gene is truncated in many strains, nearly always due to the insertion of an ISVα5-element (see supplementary fig. S4, Supplementary Material online). It is also worth mentioning that this truncation apparently co-occurs with the presence of a plM1-like plasmid (carrying two ISVα5 elements, see supplementary table S2, Supplementary Material online; bungled only by strain 87-9-116). Whether there is a functional link between these two genetic coincidences is unknown. A further exhaustive scrutiny of V. anguillarum CIsI genes is not within the scope of this study. However, the completion of seven additional genomes means that we are nevertheless able to present the scientific community with significant new knowledge.

The presence of repetitive IS-elements may present major technical challenges during sequencing and assembly of microbial genomes, especially when using short read methods, and the majority of genomes in the archives are therefore frequently found in a large number of contigs (Busschaert et al. 2015). Our work revealed 18 new IS-elements (ISVα3-ISVα20), which are available in the “ISfinder” database (Sigier et al. 2006) (see supplementary table S3 and data file S1, Supplementary Material online). Resolving the order of a high number of contigs by using, for example, long-range PCRs is very time-consuming and costly. As an alternative, we used PacBio sequencing, which offers long-sequence reads, and is therefore excellent for resolving regions with repetitive DNA. The resulting sequences were therefore de novo assembled into circular, gap free contigs without ambiguous bases. For strains CNEVA NB11008 and JLL237, discrepancies after PacBio sequencing and assembly were bioinformatically resolved. Two of the three PacBio circular contigs in strain CNEVA NB11008 were found to contain subsets of a super-integron located on Chromosome 2 (based on the presence and distribution of attC sites). Regarding the three PacBio circular contigs from strain JLL237, their size distribution clearly suggested a merger of the two smallest into a complete circularized Chromosome 1; a reassembly of the two was also supported by their lack of attC-sites. The Artemis Comparison Tool (ACT) was used to make comparisons between the respective PacBio contigs and the NB10 genome (LK021130/LK021129), forming the basis of the bioinformatic correction of their final chromosome sequences.

In summary, we have in this work sequenced seven strains of V. anguillarum to completion using the PacBio method, thus bringing the total number of finished genomes to 11 (as of March 2018). A pan genome based on the 11 genomes was calculated, and includes 7,667 gene clusters in total; 2,574 core clusters, 2,183 accessory clusters, and 2,910 unique clusters. These numbers show that the total number of genes among the strains is much greater than those found in each individual strain, which suggests considerable variation among strains, and that more genomes should be sequenced to completion in order to perform detailed genome comparisons, thus significantly further increasing the supply of resources for future studies of this important fish pathogen.

Supplementary Material
Supplementary data are available at Genome Biology and Evolution online.

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