

# Complete Genome Sequence of the Marine Fish Pathogen *Vibrio anguillarum* Harboring the pJM1 Virulence Plasmid and Genomic Comparison with Other Virulent Strains of *V. anguillarum* and *V. ordalii*<sup>∇†</sup>

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Received 25 March 2011/Returned for modification 21 April 2011/Accepted 4 May 2011

**We dissected the complete genome sequence of the O1 serotype strain *Vibrio anguillarum* 775(pJM1) and determined the draft genomic sequences of plasmidless strains of serotype O1 (strain 96F) and O2β (strain RV22) and *V. ordalii*. All strains harbor two chromosomes, but 775 also harbors the virulence plasmid pJM1, which carries the anguibactin-producing and cognate transport genes, one of the main virulence factors of *V. anguillarum*. Genomic analysis identified eight genomic islands in chromosome 1 of *V. anguillarum* 775(pJM1) and two in chromosome 2. Some of them carried potential virulence genes for the biosynthesis of O antigens, hemolysins, and exonucleases as well as others for sugar transport and metabolism. The majority of genes for essential cell functions and pathogenicity are located on chromosome 1. In contrast, chromosome 2 contains a larger fraction (59%) of hypothetical genes than does chromosome 1 (42%). Chromosome 2 also harbors a superintegron, as well as host “addiction” genes that are typically found on plasmids. Unique distinctive properties include homologues of type III secretion system genes in 96F, homologues of *V. cholerae* zot and ace toxin genes in RV22, and the biofilm formation *syp* genes in *V. ordalii*. Mobile genetic elements, some of them possibly originated in the pJM1 plasmid, were very abundant in 775, resulting in the silencing of specific genes, with only few insertions in the 96F and RV22 chromosomes.**

*Vibrio anguillarum* is a marine pathogen that causes vibriosis in close to 50 species of fish, including cultured and wild fish, mollusks, and crustaceans, in marine, brackish, and fresh water (1). Vibriosis is a hemorrhagic septicemia with dire consequences for fish rearing, especially in countries that depend heavily on fish for their food consumption. Despite the fact that *V. anguillarum* is a dramatic cause of vibriosis in fish, little is known about the genomic composition of this important pathogen. Although 23 serotypes have been reported in *V. anguillarum*, the O1 and O2 serotypes are the major causative agent of fish vibriosis (32, 66, 75). Many O1 serotype strains harbor 65-kb pJM1-type plasmids, which carry the siderophore anguibactin biosynthesis and transport genes, a main virulence factor of *V. anguillarum*, while one of the O1 serotype strains and other serotypes, such as all of the O2 strains, are plasmidless (1, 13, 33, 76). Many virulence factors have been characterized, but we are still far from getting the whole picture of the virulence mechanisms of this pathogen (1, 50). The fact that the pJM1 plasmid is an important component of virulence for the 775 strain but that the other three strains examined do not harbor this plasmid and are still virulent indicates that they must have different mechanisms to cause disease. Moreover,

O1 serotype strains cause disease in salmonid fish, whereas O2β strains are usually isolated from cod and other nonsalmonids (1, 32, 43). We have previously performed random genome sequencing of *V. anguillarum* 775 genomic DNA and identified potential virulence factors (61). Some of the genes identified in that study, such as hemolysins and the Rtx toxin, were later confirmed as virulence factors of this bacterium (34, 60).

In this study, we have determined the whole genome sequence of *V. anguillarum* 775(pJM1) and have done comparative genomics with virulent plasmidless strains of serotype O1 and O2β and with *V. ordalii*. The last is highly related to *V. anguillarum* and was previously recognized as *V. anguillarum* biotype 2 (64). Schiewe et al. (63) proposed the name *V. ordalii* based on cultural and biochemical characteristics, as well as DNA homology with biotype 1. These experiments demonstrated that *V. anguillarum* and *V. ordalii* are 58 to 69% related based on DNA-DNA hybridization (64). Histopathological studies have shown that vibrioses caused by *V. anguillarum* and *V. ordalii* are dramatically different (59). In early stages of infection, *V. anguillarum* strains cause histopathological changes in blood, loose connective tissue, kidney, spleen, gills, and posterior gastrointestinal tract, and these bacteria are most abundant in the blood, although they appear uniformly dispersed throughout the affected tissues. Conversely, *V. ordalii* develops the bacteremia only at late stages of disease, and the concentration of bacterial cells in blood is normally less than in the *V. anguillarum* infection. Tissues displaying histopathological changes after infection with *V. ordalii* are skeletal and cardiac muscles, anterior and posterior gastrointestinal

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† Supplemental material for this article may be found at <http://iai.asm.org/>.

∇ Published ahead of print on 16 May 2011.

tract, and gills, and the bacteria are not evenly dispersed but are present within tissues as colonies or aggregates of cells. In this work, we identified common and strain-specific virulence genes from *V. anguillarum* and *V. ordalii* that could explain the virulence mechanisms of these bacteria and that would be useful for future development of vaccines.

#### MATERIALS AND METHODS

**Genome sequence data.** We sequenced the genomes of *Vibrio anguillarum* 775 (ATCC 68554), 96F, and RV22 and *V. ordalii* ATCC 33509. The *V. anguillarum* 775 strain was isolated from Coho salmon (*Oncorhynchus kisutch*) in the U.S. Pacific Ocean coast (14), strain 96F was isolated from striped bass (*Morone saxatilis*) in the United States (76), and strain RV22 was isolated from turbot (*Scophthalmus maximus*) in Spain (33). In addition, we sequenced a strain of *V. ordalii* ATCC 33509 isolated from Coho salmon (*Oncorhynchus kisutch*) kidney in Puget Sound, WA (63). There is some confusion in the literature as to the genus name for *Vibrio anguillarum*, because it was suggested that it should be renamed *Listonella* (37). However, workers in the vibrio field have been using the classical *Vibrio* name for the genus. Recently, we demonstrated that *Vibrio anguillarum* is a *bona fide* *Vibrio* species, and we proposed the use of *V. anguillarum* instead of the more recent heterotype *Listonella* for this bacterium (69).

**Library, sequencing, gap closure, and assembly.** To complete the genome, *V. anguillarum* 775 was sequenced using Roche 454 technology. Whole genomic DNA was subjected to 454 Titanium sequencing with 32-fold coverage. For gap closure, we generated amplicons using long-range PCR and sequence assembly by primer walking and prepared cosmid and fosmid libraries for fosmid (cosmid) end sequencing, scaffold assembly on gapped genome, and primer walking sequence assembly. For homopolymers, we searched for regions with >5 identical bases in a row and then cloned the region and confirmed by Sanger capillary sequencing using an AB 3130XL instrument in the MMI Core Facility.

The reads were assembled using Newbler software version 2.3 (Roche/454 Life Sciences, Branford, CT) combined with the generation of cosmid and fosmid libraries using SuperCos1 (Stratagene, La Jolla, CA) and a CopyControl fosmid library production kit (Epicentre, Madison WI) using total DNA from *V. anguillarum* 775. The cosmid and fosmid DNA was purified and subjected to end sequencing by the Sanger method. For gap closure of the initial 71 contigs, comparative genomics with closely related *Vibrio* genomes was used to design amplicons using primers from the contig ends. The amplicons were amplified using PCR methods. Draft genome sequencing of *V. anguillarum* 96F and RV22 and *V. ordalii* ATCC 33509 was also performed using the Roche 454 technology, and the reads were also assembled using Newbler software version 2.3 (Roche/454 Life Sciences) of the 454 Titanium pyrosequencing approach, resulting in the following contigs: 220 for 96F, 362 for RV22, and 281 for *V. ordalii*. The average coverage for all four genomes was 32-fold.

**Annotation and comparative genomics.** The complete genome sequence of *V. anguillarum* 775 and the draft genomic sequences of 96F, RV22, and *V. ordalii* ATCC 33509 were annotated by the Rapid Annotation System Technology (RAST) server (4). The annotation is based on subsystems, i.e., a set of functional roles that an annotator has decided should be thought of as related. Frequently, subsystems represent the collection of functional roles that make up a metabolic pathway, a complex (e.g., the ribosome), or a class of proteins (e.g., two-component signal transduction proteins). The steps in annotation by RAST are as follows. First, the call of tRNAs and rRNAs, then an initial call of open reading frames (ORFs), is performed using GLIMMER2 (17). After an initial prediction, a set of representative sequences universal or nearly universal in prokaryotes, for example, the tRNA synthetases, is used to search protein-encoding genes from the new genome. The outcome can be used to estimate the closest phylogenetic neighbors of the newly sequenced genome and to confirm the putative gene prediction by GLIMMER2. The next step is an initial metabolic reconstruction, reflecting basic divisions of function, producing detailed estimates of the genome contents that were successfully connected to the subsystems. To perform the comparative analysis, we use the RAST Compare Metabolic Reconstruction tool. This tool utilizes the BLAST algorithm to perform the analysis. In addition, we used the GeneWiz browser 0.94 server to visualize all of the genomes (24).

**Prediction of replication origins.** To predict the origins of replication of the chromosomes, we used the DoriC tool, which utilizes z-curve analysis for nucleotide distribution asymmetry, DnaA box distribution, genes adjacent to candidate *oriCs*, and phylogenetic relationships (89).

**GIs and determination of dinucleotide relative abundance values.** To predict the genomic islands (GIs), we used the IslandViewer tool (31). The tool utilizes

TABLE 1. General features of the *V. anguillarum* 775 genome

Feature	Result for:		
	Chromosome 1	Chromosome 2	pJM1
Size (bp)	3,063,912	988,135	65,009
No. of CDS <sup>b</sup>	2,864	951	65
GC content (%)	44.65	44.09	42.57
No. of tRNAs	82	4	0
No. of rRNAs	22	0	0
<i>oriC</i> localization (nt) <sup>c</sup>	219251...219628	535047...535332	Ori1
<i>oriC</i> GC content (%)	39.95	30.42	
Length of <i>oriC</i> (bp)	377	285	
No. of <i>dnaA</i> boxes	5	4	
Genbank accession no. for:			
ParA	VAA_00834	VAA_00096	AY312585.1 <sup>a</sup>
ParB	VAA_00833	VAA_00097	AY312585.1 <sup>a</sup>
Soj-like protein	VAA_03384		
MukBEF	VAA_02164- VAA_02166		
StbDE toxin-antitoxin		VAA_00010- VAA_00011	

<sup>a</sup> Accession number of plasmid pJM1.

<sup>b</sup> CDS, coding sequences.

<sup>c</sup> nt, nucleotides.

SIGI-HMM and IslandPath-DIMOB methods (26, 79). The SIGI-HMM method uses a hidden Markov model (HMM) and measures codon usage to identify possible genomic islands. The IslandPath method identifies genomic islands by visualizing several common characteristics of GIs, such as abnormal sequence composition or the presence of genes that are functionally related to mobile elements. We determined the dinucleotide relative abundance value for each island predicted by IslandViewer. Mononucleotide and dinucleotide frequencies were calculated using COMPSEQ (EMBOSS). Dinucleotide relative abundances ( $\rho^*XY$ ) were calculated using the equation  $\rho^*XY = f_{XY}/f_Xf_Y$  where  $f_{XY}$  denotes the frequency of dinucleotide XY, and  $f_X$  and  $f_Y$  denote the frequencies of X and Y, respectively (28). Statistical theory and data from previous studies indicate that the normal range of  $\rho^*XY$  is between 0.78 and 1.23 (28). The difference in genome signature between two sequences is expressed by the genomic dissimilarity ( $\delta^*$ ), which is the average of the relative abundance in the dinucleotide difference between two sequences. The dissimilarities in relative abundance of dinucleotides between both sequences were calculated using the equation described by Karlin et al. (28):  $\delta^*(f,g) = 1/16 \sum |\rho^*XY(f) - \rho^*XY(g)|$  (multiplied by 1,000 for convenience), where g indicates the island genome and the sum extends over all dinucleotides.

**Nucleotide sequence accession numbers.** The complete genome of *V. anguillarum* 775 has been deposited at DDBJ/EMBL/GenBank under the accession numbers CP002284 to CP002285. The sequences of the genomes of *V. anguillarum* 96F and RV22 and *V. ordalii* ATCC 33509 have been deposited at DDBJ/EMBL/GenBank under the accession numbers AEZA00000000, AEZB00000000, and AEZC00000000, respectively.

## RESULTS

***Vibrio anguillarum* 775 genome.** Table 1 and Fig. 1 show that the complete genome sequence of *V. anguillarum* 775 consists of two circular chromosomes, chromosome 1 (Chr1) of 3,063,912 bp and chromosome 2 (Chr2) of 988,135 bp, in addition to the 65,009-bp pJM1 plasmid, which carries anguibactin biosynthesis and ferric anguibactin transport genes (19). *V. anguillarum* 775 contains 2,093 (54%) products (protein) identified in subsystems and 1,783 (46%) not included in subsystems, indicating a large portion of proteins with unknown function according to the RAST method. The GC contents of Chr1 and Chr2 are 45% and 44%, respectively, and that of the pJM1 plasmid is 43%. The total number of subsystems found in the 775 genome is 396. In *V. cholerae* El Tor N16691, the total number of subsystems is 423. The lower number in *V. anguil-*

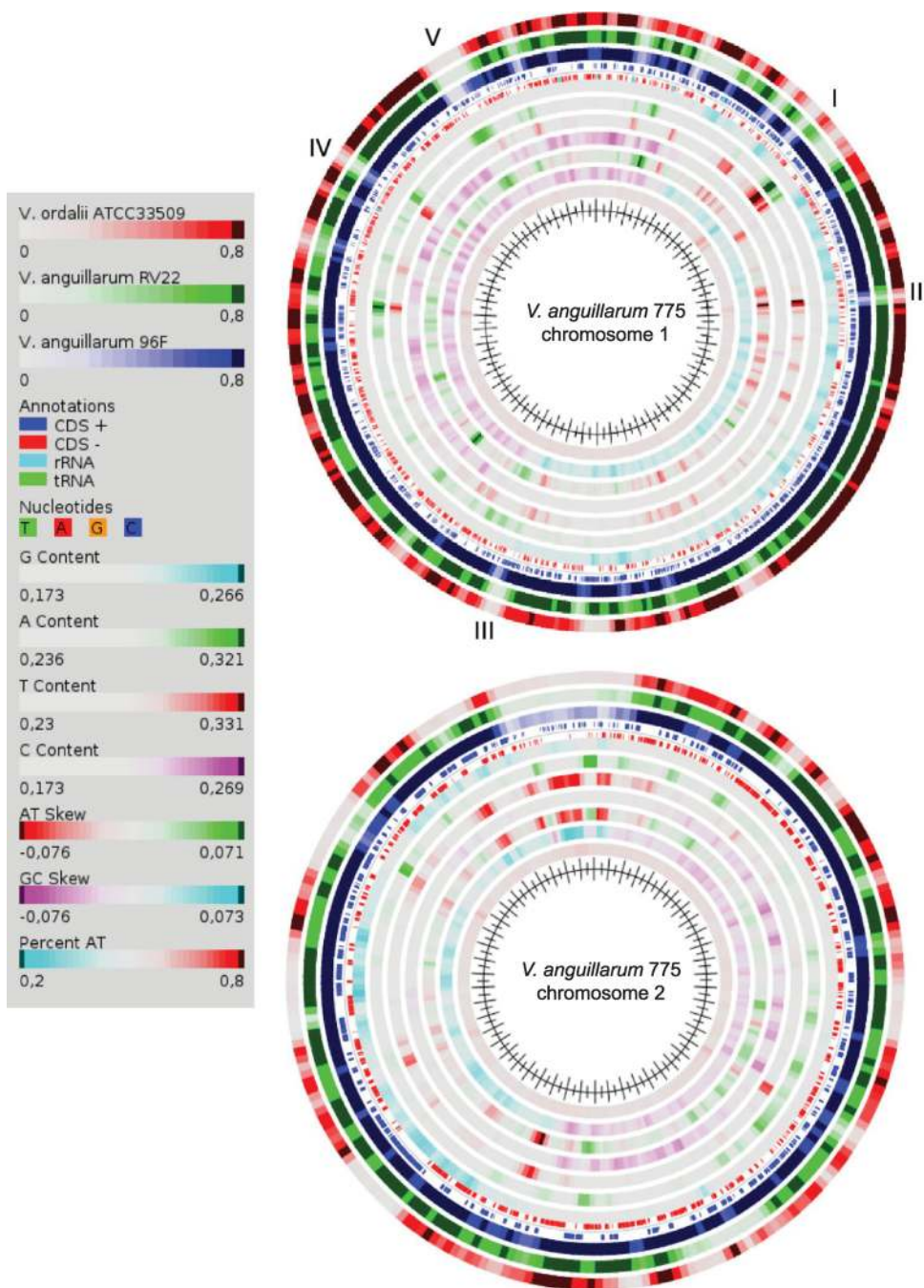


FIG. 1. Atlas visualization of chromosomes 1 and 2 of strain 775. The GeneWiz browser 0.94 server was used to visualize all of the genomes (24). The reference genome is *V. anguillarum* 775, and the four outer circles represent the genomes from the other *Vibrio* strains. The Roman numerals represent genes that are absent from the 96F and RV22 strains and *V. ordalii*. The numbers in the table at left correspond to the intensity of the colors.

larum could be due to the fact that this is the first complete sequencing of the *V. anguillarum* genome and that, therefore, knowledge about the subsystems needs to be enhanced. This total genome was predicted to have 3,876 ORFs in the two chromosomes and 65 ORFs on the pJM1 plasmid. Approximately 1,007 (26%) of the total number of chromosomal and plasmid genes represent hypothetical proteins. Figure 2 shows that the majority of gene products were classified into catego-

ries such as production and/or utilization of carbohydrates, amino acids, and derivatives, as well as protein metabolism. Table S1 in the supplemental material shows the comparison of the molecular parameters of the genomes of *V. anguillarum* 775 (4.1 Mb), 96F (4.0 Mb), and RV22 (4.0 Mb) and *V. ordalii* (3.4 Mb).

**Distribution of origins of chromosomes, plasmids, and partition genes in the 775 strain.** Table 1 shows the origins of

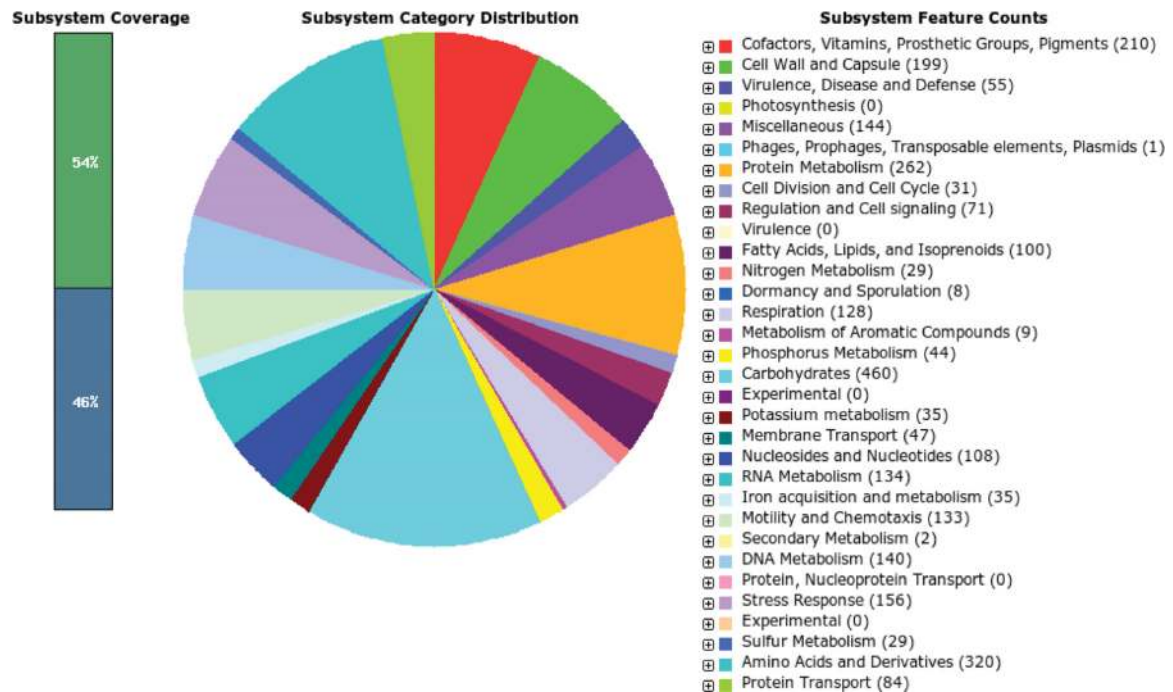


FIG. 2. Distribution of subsystems found in the *V. anguillarum* 775 genome. Subsystem coverage is indicated by green and blue bars. The green bar corresponds to the percentage of the proteins included in the subsystems, and the blue bar corresponds to the percentage of the proteins that are not included in the subsystems. The complete genome sequence of *V. anguillarum* 775 and the draft genomic sequences of 96F, RV22, and *V. ordalii* ATCC 33509 were annotated using the Rapid Annotation System Technology (RAST) server (4).

replication identified as follows. *V. anguillarum ori1* (*ori1<sub>va</sub>*) in Chr1 is approximately 39.95% GC, with 5 *dnaA* boxes, and is close to the *dnaA* gene. *ori2<sub>va</sub>* in Chr2 is approximately 30.42% GC, with 4 *dnaA* boxes, and is close to the *parA* gene. We also identified *parA* and *parB* homologues in both chromosomes and in the pJM1 plasmid, although Chr1 encodes exclusively partition proteins, such as Soj-like protein and MukBEF, and Chr2 has StbDE as a toxin-antitoxin system. The origin of pJM1 DNA replication has been located adjacent to ORF50 (9).

**Genes of interest in *V. anguillarum* 775.** We first investigated the location of housekeeping and other essential genes and found the majority located on Chr1, as is the case in other vibrios (data not shown). We also analyzed proven and potential virulence-associated and environmentally associated genes and their distribution in the 775 chromosomes and the plasmid pJM1. Table 2 shows that the majority of the ferric anguibactin biosynthesis and transport genes are present in the pJM1 plasmid, with the exception of *vabA*, which is essential for anguibactin biosynthesis but resides on Chr1 (3). The TonB1 and TtpC-TonB2 clusters, which are essential for iron transport, are encoded on Chr1 (67, 68). The *fur* gene is located on Chr1, which also harbors the heme determinants, although we have also identified a heme-related *V. cholerae hutR* homologue on Chr2 (42, 48, 49, 70). Genes involved in ferric vanchrobactin and/or ferric enterobactin transport are located on Chr1, although vanchrobactin biosynthesis in 775 is abrogated by a mutation in *vabF* and there are no biosynthesis genes for enterobactin (51). It is of interest that potential ABC transporters FvtB-FvtE for vanchrobactin and enterobactin uptake are found on Chr2. Following the trend of other genes essential

for iron transport, Chr1 also harbors the genes *feoABC* for ferrous iron, *fbpABC* for ferric iron, and *fhuABCDE* for ferrichrome transport, which are homologues to those of *V. cholerae* (62, 85).

The genes related to the repeat-in-toxin gene *rtx* are located on Chr1. This toxin was also implicated in *V. anguillarum* virulence (34). In addition, we identified two more *rtx* toxin genes, one on Chr1 and the other on Chr2. As in *V. vulnificus*, only one RTX cluster of a total of three contains a complete set of *rtx* genes (8, 34). The hemolysin genes are located mainly on Chr1, except for *vah1*, which was located on Chr2 (25, 60, 86). The metalloprotease genes are located on Chr2 (44, 46). Genes for other secreted enzymes are located on Chr2, except for lysophospholipase L2, which is on Chr1. We also identified flagellar and chemotaxis genes previously demonstrated to be associated with virulence in immersion challenges (41, 45). These genes were identified on Chr1, whereas chemotaxis genes were found on both chromosomes. We identified *V. cholerae* type 4 pilus and mannose-sensitive hemagglutinin (MSHA) pilus homologues (39). These genes were located on Chr1, which also harbored the type VI secretion system locus containing the *vstA-vstH* genes (81). In addition, we found another novel locus harboring a second type VI secretion system in Chr2. These two systems are highly related; however, the genes in the two systems are organized differently.

Quorum sensing-related genes are distributed in both chromosomes, although Chr2 carries most of the genes described in the literature (16). Chr2 also harbored genes for homologues of regulators, such as *slyA* and *virK*, as adjacent genes transcribing in opposite directions. These genes were demon-

TABLE 2. *V. anguillarum* and *V. ordalii* genes proven to be or potentially associated with virulence or the environment

Gene category and name	Presence/absence of gene <sup>b</sup> in <i>V. anguillarum</i> 775:			GenBank accession no.	Presence/absence of gene <sup>b</sup> in:		
	Chr1	Chr2	pJM1		96F	RV22	<i>V. ordalii</i>
<b>Anguibactin biosynthesis and iron transport</b>							
<i>angA-angE, angGHMNRU</i>	—	—	x	AY312585.1 <sup>a</sup>	—	—	—
<i>vabA-vabE</i>	x	—	—	VAA_02392-VAA_02395, VAA_02076	x	x	x
<i>fatA-fatD</i>	—	—	x	AY312585.1 <sup>a</sup>	—	x	—
<i>tipC, tonB2, exbB2, exbD2</i>	x	—	—	VAA_01800-VAA_01804	x	x	—
<i>tonB1, exbB, exbD1</i>	x	—	—	VAA_01389-VAA_01387	x	x	x
Utilization protein for unknown catechol siderophore	—	x	—	VAA_01367	x	x	x
<i>fur</i>	x	—	—	VAA_03429	x	x	x
<b>Heme transport</b>							
TonB-dependent heme receptor HutR gene	—	x	—	VAA_02922	x	x	—
<i>huvA-huvD, huvSXZ</i>	x	—	—	VAA_01392, VAA_01385-VAA_01383, VAA_01390-VAA_01391	x	x	x (except <i>huvXZ</i> )
<b>Ferric enterobactin and ferric vanchrobactin transport</b>							
<i>fvtA</i>	x	—	—	VAA_02079	x	x	x
<i>fvtB-fvtE</i>	—	x	—	VAA_03000-VAA_03003	x	x	—
<i>fetAB</i>	x	—	—	VAA_00188-VAA_00189	x	x	—
<b>Ferrous and ferric transport</b>							
<i>feoABC</i>	x	—	—	VAA_03398-VAA_03400	x	x	x
<i>fbpCBA</i>	x	—	—	VAA_01820-VAA_01823, VAA_00769-VAA_00767	x	x	x
<b>Ferrichrome</b>							
<i>fhuACBD</i>	x	—	—	VAA_02423-VAA_02426	x	x	Only FhuA
<b>Hemolysins</b>							
Hemolysin genes	x	—	—	VAA_00718	x	x	x
Putative hemolysin genes	x	—	—	VAA_02701	x (4)	x (4)	x (1)
21-kDa hemolysin precursor gene	x	—	—	VAA_03731	x	x	x
<i>vah1</i>	—	x	—	VAA_02992	x	x	x
<i>vah2-vah5</i>	x	—	—	VAA_01090, VAA_03209, VAA_03298, VAA_00294	x	x (except <i>vah4</i> )	x (except <i>vah4</i> )
<i>mliD</i>	x	—	—	VAA_03554	x	x	x
<b>RTX toxins</b>							
<i>rtxEDBHCGA</i>	x	—	—	VAA_03091-VAA_03085	x	x	x
RTX toxin and related Ca <sup>2+</sup> -binding protein genes	—	x	—	VAA_02949	x	x	x
RTX toxin gene	x	—	—	VAA_02070	x	—	—
<b>Metalloproteases</b>							
<i>empA</i>	—	x	—	VAA_01306	x	x	x
<i>priV</i>	—	x	—	VAA_02995	x	x	x
Zinc protease, insulinase family gene	—	x	—	VAA_01003	x	x	—
<b>Secreted enzymes</b>							
Phospholipase C gene	—	x	—	VAA_01330	x	—	—
Nonhemolytic enterotoxin lytic component L1 gene <sup>c</sup>	—	x	—	VAA_00241-VAA_00240	x	x	x
Microbial collagenase gene, secreted	—	x	—	VAA_02998	x	x	—
Lysophospholipase L2 gene	x	—	—	VAA_01963	x	x	x
Lipase family protein gene	—	x	—	VAA_01024	x	x	x
<b>Motility and chemotaxis</b>							
<i>fliEDBGI</i>	x	—	—	VAA_03469-VAA_03465	x	x	x
<i>fliS, fleSQ</i>	x	—	—	VAA_03463-VAA_03459	x	x	x
<i>fliE-fliR, fliB</i>	x	—	—	VAA_03459-VAA_03443	x	x	x

Continued on following page

TABLE 2—Continued

Gene category and name	Presence/absence of gene <sup>b</sup> in <i>V. anguillarum</i> 775:			GenBank accession no.	Presence/absence of gene <sup>b</sup> in:		
	Chr1	Chr2	pJM1		96F	RV22	<i>V. ordalii</i>
<i>flgTOPNMA</i>	x	—	—	VAA_03532-VAA_03526	x	x	x
<i>cheVR</i>	x	—	—	VAA_03525-VAA_03524	x	x	x
<i>flgB-flgL</i>	x	—	—	VAA_03522-VAA_03512	x	x	x
<i>flaAC</i>	x	—	—	VAA_03511-VAA_03510	x	x	x
<i>flhA</i>	x	—	—	VAA_03392	x	x	x
<i>flhL</i>	x	—	—	VAA_00455	x	x	x
<i>rpoHN</i>	x	—	—	VAA_00451, VAA_02739	x	x	x
<i>motYBA</i>	x	—	—	VAA_02089, VAA_01161-VAA_01162	x	x	x
Methyl-accepting chemotaxis gene	x (23)	x (19)	—		x (40)	x (45)	x (28)
Type IV pilus							
<i>pilABCD</i>	x	—	—	VAA_00732-VAA_00728	x	x	x
<i>pilVWE</i>	x	—	—	VAA_01191, VAA_01193, VAA_01195	x	x	x
Putative type IV pilin gene	x	—	—	VAA_01194	x	x	x
<i>pilUT</i>	x	—	—	VAA_01655-VAA_01656	x	x	x
<i>pilF</i>	x	—	—	VAA_03067	x	x	x
Multimodular transpeptidase-transglycosylase gene	x	—	—	VAA_02627, VAA_00759	x	x	x
<i>pilMNOPQ</i>	x	—	—	VAA_02628-VAA_02632	x	x	x
MSHA type 4							
<i>mshQPODCABF</i>	x	—	—	VAA_01721-VAA_01728	x	x	x
<i>mshGENMLKJIH</i>	x	—	—	VAA_01729-VAA_01738	x	x	x
Type VI secretion system							
<i>vtxA-vtsH</i> system I	x	—	—	VAA_03042-VAA_03021	x	x	—
<i>vtxA-vtsH</i> system II	—	x	—	VAA_02833-VAA_02849	x	x	—
Quorum sensing							
<i>luxS</i>	x	—	—	VAA_00716	x	x	x
<i>luxR</i>	x	—	—	VAA_01741	x	x	x
<i>vanMN</i>	—	x	—	VAA_00956-VAA_00957	x	x	x
<i>vanUO</i>	x	—	—	VAA_02384-VAA_02385	x	x	x
<i>vanRI</i>	—	x	—	VAA_01245-VAA_01246	x	x	x
<i>vanQP</i>	—	x	—	VAA_01548-VAA_01547	x	x	x
<i>vanT</i>	x	x	—	VAA_00743, VAA_00909	x	x	x
Regulators							
<i>virK, slyA</i>	—	x	—	VAA_02801-VAA_02800	x	x	x
DNA-binding protein H-NS gene	x	—	—	VAA_02203	x	x	x
Transcriptional activator HlyU gene	—	x	—	VAA_00407	x	x	x
Transcriptional regulator VpsT gene	—	x	—	VAA_01253	x	x	x
<i>hfq</i>	x	—	—	VAA_00118	x	x	x

<sup>a</sup> GenBank accession number of plasmid pJM1.

<sup>b</sup> “x”, present genes; “—”, absent genes. The numbers in parentheses indicate the number of copies in the genome.

<sup>c</sup> Similar to VCA0884.

strated to be important for virulence in different bacteria, such as *Salmonella enterica* serovar Typhimurium for *slyA* and *Campylobacter jejuni*, *Salmonella* Typhimurium, and *Shigella flexneri* for *virK* (18, 21, 52, 54). Regulators *hlyU* and *hns* were located on Chr2 and Chr1, respectively. We recently reported that homologues of these genes in *V. vulnificus* operated in silencing-antisilencing regulation of the *rxnA1* gene, which we proved to be an important virulence factor in this bacterium (35, 36). Chr2 also harbors a homologue of the *V. cholerae* *vpsT* gene, which regulates matrix production and motility in that bacterium (30). The RNA chaperone Hfq has previously been shown to be necessary for *V. anguillarum* NB10 virulence (80). This gene was located on Chr1.

**GIs.** Ten putative GIs were identified in *V. anguillarum* 775, with the majority (eight) on chromosome 1 and two on chromosome 2, totaling 199 genes (Fig. 3; also see Table S2 in the supplemental material). To predict the islands, we utilized the IslandViewer tool, which uses a hidden Markov model and measures codon usage. The 10 islands ranged in size from 4 kb to 140 kb, and the largest genomic island (GI2) had 74 genes, whereas the smallest (GI7) had 5 genes (Fig. 3; Table S2). The largest island had a GC content of approximately 39%, indicating a region of easy transcription but difficult translation due to codon usage bias (84). The largest island comprised genes for transposase, dTDP-rhamnose synthesis (lipopolysaccharide [LPS] component), glycosyltransferase, aminotrans-

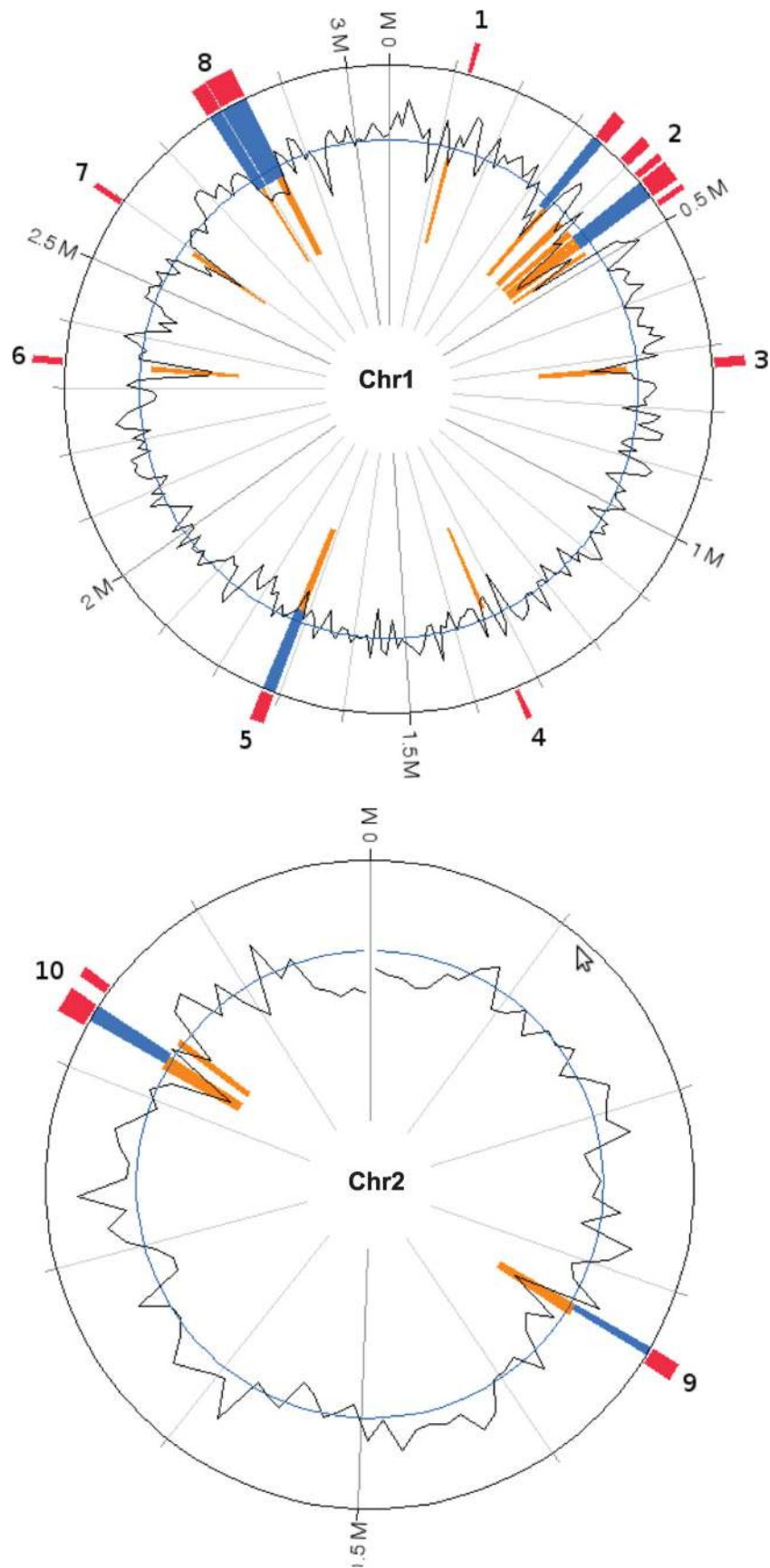


FIG. 3. Genomic islands of strain 775 predicted by IslandViewer. The colors red, blue, and orange represent the three methods for island detection (integrated detection, SIGI-HMM, and IslandPath-DIMOB, respectively). The black line represents the GC content (%).

ferases, and acetoin and butanediol metabolism. Three islands, GI3, GI6, and GI8, are inserted adjacent to tRNA-Leu, tRNA-Tyr, and tRNA-Met loci, respectively. Six of the islands contain integrase and transposase genes, which may indicate a possible mechanism of insertion. The genomic signature between the genomes of *V. anguillarum* 775 and 96F strains is 1.5, whereas the genomic signature for each GI compared with the chromosomes ranged from 37 to 116, showing the high genomic dissimilarity between the islands and the genomes.

***Vibrio anguillarum* 775 superintegron.** The superintegron region of strain 775 was approximately 53 kb, with a GC content of 41%, and contained homologues of the *V. cholerae* *infC* gene (translation initiation factor 3), *rpmL* and *rplT* (ribosomal proteins L20 and L35), the integrase gene, and the *attI* and *attC* recombination sites as found in the *V. cholerae* superintegron (40). The superintegron has 96 genes, of which 72 (75%) were identified as hypothetical proteins (see Table S3 in the supplemental material). The *V. anguillarum* strains 96F and RV22 and *V. ordalii* have the same genes as the 775 superintegron.

**Comparative analysis of *V. anguillarum* strains 775, 96F, and RV22 and *V. ordalii*.** Table S1 shows the comparison of the molecular parameters of the genomes of *V. anguillarum* 775 (4.1 Mb), 96F (4.0 Mb), and RV22 (4.0 Mb) and *V. ordalii* (3.4 Mb). What leaps to the eye is the reduced size of the *V. ordalii* genome. The difference in genome size is similar to the value we obtained using DNA-DNA hybridization analysis, which demonstrated that the *V. ordalii* genome is about 70% of the genome size of the O1 and O2 strains (64). In order to determine the similarity between the strains and strain-specific genes, a comparison was performed using BLAST Atlas for both chromosomes (Fig. 1). In Fig. 1, the outermost circle (red) represents the *V. ordalii* ATCC 33509 genome, which is the most distinct in relation to strain 775, followed by the plasmidless strains of serotype O2 $\beta$  RV22 (green circle) and serotype O1 96F (blue circle). The genomes of *V. anguillarum* strains 775 and 96F had higher numbers of genes in common. Chr1 shows higher gene conservation among these strains than does Chr2, underscoring the higher plasticity of Chr2. There are 5 specific regions in Chr1 of strain 775 in comparison with the other strains. The Roman numerals in Fig. 1 represent genes that are absent from the 96F and RV22 strains and *V. ordalii* (also see Table S4 in the supplemental material). In total, these regions corresponded to 173 unique genes for *V. anguillarum* 775. All five specific regions of strain 775 have integrases and transposases. The regions I, II, III, and V include the islands GI2, GI3, GI5, and GI8, respectively (Fig. 3). The GC content of region II, at 39%, is lower than that for the total chromosome. This region contains several tRNA and rRNA genes. The genes in each of these regions are shown in Table S4. There are 2,681 genes common to all of the strains, using a cutoff of 60% identity. The majority of the 775 chromosome-specific genes are in the five major regions. There are 65 genes carried on the pJM1 plasmid that are also unique for 775, and because we did not close the gaps on 96F, RV22 and *V. ordalii*, we did not search for smaller clusters or individual chromosomal genes, since they could have been missed in the draft sequence of the latter genomes.

**Distribution of virulence-related and environmentally related genes in the chromosomes of 775, 96F, RV22, and *V. ordalii*.** Table 2 shows the comparison of putative virulence

gene distributions in the three *V. anguillarum* strains and *V. ordalii*. The majority of the genes were conserved throughout the strains; however, the anguibactin biosynthesis and transport genes were found only on the pJM1 plasmid in the 775 strain, although the RV22 strain had homologues of the *fatABCD* genes. These genes appeared not to be expressed (5, 51). The *tonB1* and *tonB2-tpc* clusters were found in all of the *V. anguillarum* strains, but only the *tonB1* system was identified in *V. ordalii*. The vanchrobactin system was found in all strains, but *V. ordalii* lacks the ABC transporters *fvbB-fvbE* and *fetAB*. HutR homologues were identified only in the *V. anguillarum* strains. All hemolysin genes were conserved except for *vah4*, and the third copy of the *rtx* toxin gene and the phospholipase C gene were identified only in O1 serotype strains. Type VI secretion systems and the gene for microbial collagenase are present in all of the *V. anguillarum* strains but absent from *V. ordalii*, while the chemotaxis genes are distributed in the two chromosomes in all strains.

**Distribution of transposase and phage proteins in *V. anguillarum* and *V. ordalii*.** We compared the chromosome distributions of transposase and phage proteins in the three *V. anguillarum* strains and *V. ordalii* (see Table S5a in the supplemental material). *V. anguillarum* 775 contains more transposase genes (about 53) than 96F (about 23), RV22 (about 42), and *V. ordalii* (about 18). In pJM1, there are two insertion sequences, ISV-A1 (transposase, ORF2) and ISV-A2 (transposase, ORF11), and many transposases for which insertion sequences have not been identified (73). Table S5b in the supplemental material shows transposases that possibly originated in the pJM1 plasmid in many locations of the chromosomes of *V. anguillarum* 775. One of these insertions took place in the *vabF* gene, resulting in the abrogation of vanchrobactin biosynthesis. Two of the pJM1 transposases were not found in either of the two chromosomes. An important and exciting finding was the presence of only one pJM1 transposase on one site of the chromosomes of 96F and RV22, whereas no pJM1 transposase was found in *V. ordalii*.

***Vibrio anguillarum* genes specific to 96F compared to 775.** Table S6 in the supplemental material shows that strain 96F had genes not present in strain 775. These included several genes involved in stress response (catalase and peroxidase) and the toxin-antitoxin system (YoeB-YefM), a prophage, RNase H1, Tra system genes, OmpA/MotB, and tRNAs.

We also identified in *V. anguillarum* 96F a type III secretion system (T3SS) cluster (Fig. 4; also see Table S7 in the supplemental material) that is highly related to the *V. parahaemolyticus* type III secretion system 2 (T3SS2), which was also found in *V. cholerae* and *V. mimicus* (2, 6, 20, 38, 55). The T3SS has been experimentally shown to be important for virulence in *V. parahaemolyticus* and *V. cholerae* (20, 57). The 96F T3SS locus contains genes that are predicted to encode putative T3SS apparatus proteins and putative T3SS regulatory and effector proteins. Those genes are highly conserved with other T3SS2 genes in different *Vibrio* species. In strain 775, this cluster is missing from both chromosomes, although we find a partial sequence in the same place, indicating that possibly this strain previously carried this locus. Interestingly, this position in 775, corresponding to the 96F T3SS locus, instead contains three sets of transposase sequences, one of which appears to have its origin in pJM1, suggesting that the deletion of the T3SS locus



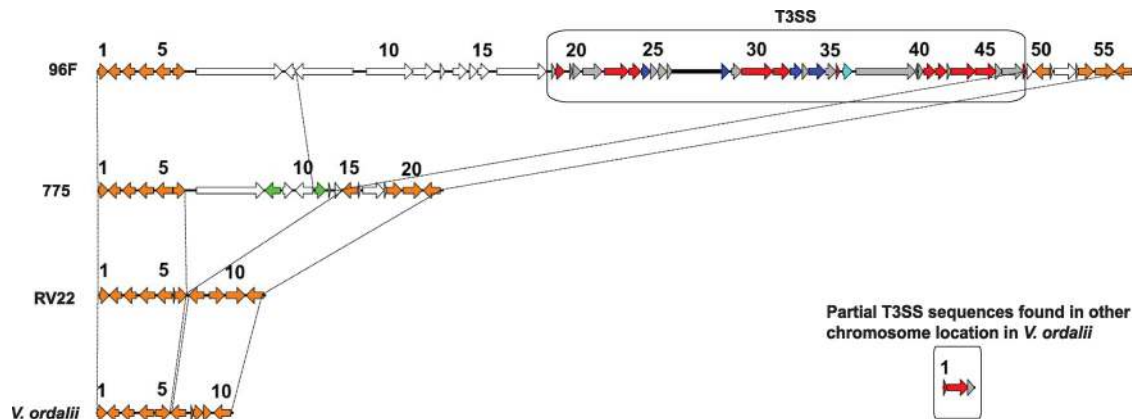


FIG. 4. T3SS of *V. anguillarum* 96F. The box shows the location of the T3SS in 96F: apparatus (red), effectors (blue), and regulator (cyan). Genes conserved among all of the strains are shown in orange. Transposase sequences are shown in green. The numbers correspond to the ORF numbers in Table S7 in the supplemental material, which shows the complete list of genes.

could have been caused by a transposition, deletion, and inversion event. The T3SS cluster was not found in RV22 or in *V. ordalii* in the 96F location, although *V. ordalii* has three T3SS homologous genes in a different location.

***V. anguillarum* genes specific to RV22 compared to 775.** The strain RV22 harbors the toxin-antitoxin systems YoeB-YefM, ParE-ParD, and HigB-HigA and has other genes not present in strain 775 (see Table S8 in the supplemental material). These genes comprised a variety of functions, including amino acid and carbohydrate metabolism, 2-keto-3-deoxyoctulosonic acid (KDO)-lipid A biosynthesis (modification and synthesis of lipid A [77]), and cell wall and capsule synthesis (e.g., sialic acid metabolism). In RV22, we also identified a TniA putative transposase and a TniB NTP-binding protein and genes that could be associated with virulence, such as those encoding the accessory *V. cholerae* enterotoxin (Ace) and the *Zonula occludens* toxin (Zot). In *V. cholerae*, Zot decreases ileal tissue resistance by modulating intercellular tight junctions, while Ace increases short-circuit current in Ussing chambers and causes fluid secretion in a ligated rabbit ileal model (22, 78). We have found that *zot* and *ace* homologues of *Vibrio* species exist only in *V. anguillarum* RV22. Although there is a *zot*-like gene in the list of 96F-specific genes, our detailed BLAST analysis indicated that it does not encode Zot.

Strain RV22 has genes specifying biosynthetic proteins and transport for a yersiniabactin-like siderophore. This cluster was also found in *Vibrio ordalii* (see Table S9 in the supplemental material). This cluster is highly conserved in many *Vibrio* species, and *Photobacterium damsela* subspecies *piscicida* carries a highly related cluster (56).

***Vibrio ordalii* genes specific to ATCC 33509 compared to 775.** Strain ATCC 33509 has genes not present in strain 775, such as those encoding production of catalase, peroxidase, and superoxide dismutase. In addition, we find genes for a ribosyl nicotinamide transporter and metal transport and toxin-antitoxin systems YefM-YoeB, ParE-ParD, and HigB-HigA. As stated above, the yersiniabactin-like siderophore cluster was also found in *V. ordalii* (see Table S9 in the supplemental material). We have also identified an entire biofilm formation cluster, the Syp system, in the *V. ordalii* genome, except for SypE, a negative regulator (Fig. 5; also see Table S10a in the supplemental

material). The Syp system has mainly been characterized as an essential factor of *V. fischeri* to colonize squid organs (47, 87, 88). It can also be found in other pathogenic *Vibrio* species, such as *V. vulnificus* and *V. parahaemolyticus*, and it contributes to biofilm formation in *V. vulnificus* (29, 88). Although the genes surrounding the Syp system are conserved, the Syp cluster was absent from the 775, 96F, and RV22 genomes (Fig. 5; Table S10a).

**Comparison of the O antigen genes in the *V. anguillarum* strains 775, 96F, and RV22 and *V. ordalii*.** It has been shown that the regions for O-antigen biosynthesis in *V. cholerae* O1, O22, O31, O37, and O139 are surrounded by *gmhD* and *rjg* genes and that this region is also important for the production of capsule of non-O1 *V. cholerae* O31 and O139 (7, 10, 11, 12). In the *V. anguillarum* serotype O1 strain, *virA* (*wbhS*) and *virB* (*wbhR*) are involved in LPS biosynthesis and have been shown to be important for virulence (53). Those genes are located in the cluster adjacent to *gmhD* (27). Furthermore, LPS genes such as *rmlD* (*wbhC*) and *rmlC* (*wbhD*) are essential for stability of the outer membrane ferric anguibactin-specific receptor FatA in the 531A strain of *V. anguillarum*, which carries the pJM1-like plasmid pJHC1. Thus, the LPS genes are necessary for ferric anguibactin transport and virulence (82). Until now, the *rjg* gene has not been described in *V. anguillarum*; however, in this work we located the entire locus in Chr1 of *V. anguillarum* 775, which included the *gmhD* and *rjg* genes surrounding the O-antigen cluster genes (Fig. 6; also see Table S10b in the supplemental material). In addition to genes that are responsible for LPS biosynthesis, we found genes such as *wza-wzc*,

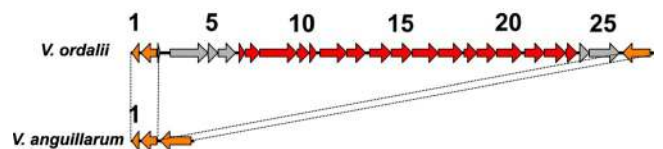


FIG. 5. Syp system in the *V. ordalii* genome. Shown are *syp* genes (red), genes encoding hypothetical proteins (gray), and genes found outside the *syp* genes in *V. ordalii* and present in *V. anguillarum* (orange). The numbers correspond to the ORF numbers in Table S10a in the supplemental material, which shows the complete list of *syp* genes.

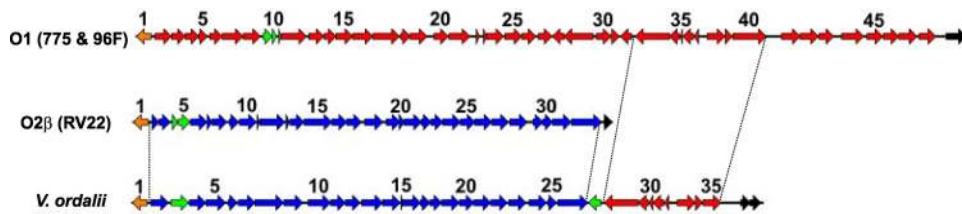


FIG. 6. Comparison of the O-antigen genes in the *V. anguillarum* strains 775, 96F, and RV22 and *V. ordalii*, which are flanked by *gmhD* (orange) and *rjg* (black) genes. The figure shows the genes for antigens O1 (red) and O2 $\beta$  (blue). Transposase sequences are shown in green. The *V. ordalii* O-antigen cluster is a hybrid of *V. anguillarum* O1 and O2 $\beta$  antigen genes. ORF2 to ORF26 of the *V. ordalii* cluster are highly related to ORF2 to ORF32 of the O2 $\beta$  cluster, and ORF28 to ORF35 of the *V. ordalii* cluster are highly related to ORF33 to ORF40 of the O1 cluster. The numbers correspond to the ORF numbers in Table S10b in the supplemental material, which shows the complete list of genes.

*wbfB-wbfD*, and *orf1*, shown to be important in exopolysaccharide biosynthesis (15). These genes are also located within this cluster. Very related *gmhD* and *rjg* genes are also found in 96F, RV22, and *V. ordalii*. It is clear from Table S10b that the O-antigen clusters for 775 and 96F are almost identical, as could be expected of these two O1 serotype strains. In RV22, the O2 $\beta$  serotype cluster between *gmhD* and *rjg* had genes different from those of 775 and 96F. *V. ordalii* presents a mosaic situation in which about 25 genes from the *gmhD* locus are highly homologous to those of the O2 $\beta$  serotype strain RV22. The rest of the genes up to *rjg* are O1 cluster genes, suggesting that *V. ordalii*, which can react with O2 $\alpha$  serum, is actually a hybrid serotype.

## DISCUSSION

The main serotypes of *V. anguillarum* associated with virulence are O1 and O2. O1 serotype strains have been reported to be the most important in causing disease in salmonid fish (32). Serotype O2 can be divided into O2 $\alpha$  and O2 $\beta$ , and O2 $\alpha$  strains have been isolated from both salmonid and marine fish, while O2 $\beta$  strains are usually isolated from cod and other nonsalmonids (1, 32, 43).

The majority of serotype O1 strains of *V. anguillarum* harbor pJM1-type plasmids carrying the siderophore anguibactin biosynthesis and transport genes and are responsible for epizootics in many parts of the world (13, 58, 65). These virulent strains are classified into two groups based on the geographic locations where they were isolated and special features of the pJM1-like plasmid. Strains from the Pacific coast of the United States and Japan harbor a pJM1 plasmid very similar to pJM1 in 775 and produce similar levels of anguibactin, whereas strains originated in the U.S. Atlantic coast and in Spain, Scotland, and the Scandinavian countries carry pJM1-like plasmids that possess several extra insertion elements and a mutation in the *angR* gene (71, 74). This mutation results in higher anguibactin production levels than those in Pacific coast strains, although this difference was not translated into an enhanced virulence phenotype and no major differences were found by DNA hybridization and biochemical studies (72, 82). These findings suggest that all O1 strains carrying a pJM1-like plasmid, independent of plasmid type and geographical isolation, are very similar. It is then clear that 775 isolated from Coho salmon in the Pacific Northwest coast of the United States can be used as the paradigm for the plasmid-carrying strains.

Therefore, to understand the genomics of virulence in *V. anguillarum*, we completely sequenced the genome of the paradigm 775 harboring the virulence plasmid pJM1 and obtained the draft genome sequence of two plasmidless strains, one of serotype O1, 96F isolated from striped bass, and another an O2 $\beta$  serotype, RV22, isolated from turbot in Spain. We also sequenced a *V. anguillarum* strain of biotype 2 ATCC 3350, now named *V. ordalii*.

We determined that each of the three *V. anguillarum* strains and the *V. ordalii* strain possess two chromosomes, whereas the 775 strain also harbors the plasmid pJM1, carrying the genes to produce anguibactin and cognate transport genes to internalize ferric anguibactin, which requires the Chr1 TonB2-TtpC system. Ferric anguibactin production and transport are the most important virulence factors once the bacterium invades the host fish. Although it does not produce these virulence factors, the 775 strain can transport vancomycin, enterobactin, ferrichrome, and heme as well. In the O2 $\beta$  serotype strain RV22 and in *V. ordalii*, we also identified biosynthesis and transport genes for a yersiniabactin-like siderophore, which potentially could play a role in virulence.

Genomic analysis using Karlin's genomic signature dissimilarity identified the presence of eight genomic islands (GIs) in Chr1 of *V. anguillarum* 775(pJM1) and two GIs in Chr2 where novel DNA was present. The 10 islands ranged in size from 4 kb to 140 kb. The genomic signature between *V. anguillarum* 775 and 96F strains is 1.5, whereas the genomic dissimilarity for each island ranged from 37 to 116, showing the high genomic dissimilarity between the islands and the genomes. Three islands, GI3, GI6, and GI8, with the highest coefficient of dissimilarity were inserted adjacent to tRNA-Leu, tRNA-Tyr, and tRNA-Met loci, respectively. The islands contain integrase and transposase genes, which may indicate a possible mechanism of insertion. Another DNA sequence that can operate in the acquisition of foreign DNA, superintegron, in the strain 775 contains integrase genes and *attI* and *attC* recombination site. The evolutionary genomic plasticity conferred by these large chromosomal regions unique to *V. anguillarum* suggests that they may play an essential role in the dynamic adaptation of this pathogen to specific ecological niches. The 775 strain carrying the pJM1 plasmid also shows a genomic variability due to the insertion of phages and transposases, many of them with a likely origin in the pJM1 plasmid.

The 775 strain harbors, in addition to many other potential virulence factors, flagellar and chemotaxis genes that have

been demonstrated to be important in virulence for fish by immersion challenge and two extra copies of the *rtx* gene that has previously been shown to be part of the *V. anguillarum* virulence repertoire (34, 41, 45). Only the O1 serotype 96F strain possesses an entire complement of type III secretion system genes, whereas only in the O2β serotype RV22 strain are the genes for the *V. cholerae* Ace and the *V. cholerae* Zot toxins found. From these results, it appears that in addition to the carriage of the pJM1 virulence plasmid, different serotypes have evolved different potential virulence mechanisms.

*V. anguillarum* causes acute hemorrhagic septicemia, but *V. ordalii* causes a very slow infection whose outcome is uneven dispersion within tissues and possibly the formation of biofilms. This result is in accordance with our finding that this bacterium carries the *syp* genes involved in biofilm formation. *V. ordalii* has suffered a large deletion of its chromosome (about 600 kb) compared to 775 and is attenuated in virulence in the trout model (59). The process of genome reduction that appears to have occurred in the transition of *V. anguillarum* to *V. ordalii* can be encountered in other bacteria, such as *Mycobacterium leprae* and *Mycobacterium tuberculosis*, the first being reduced from the second (23). Another example is that of the endosymbiont *Wolbachia* (83). It is possible that *V. ordalii* is going through a reductive evolution and could be on its way to becoming an endosymbiont, since it has the smallest genome of the vibrios. Comparisons of known virulence genes and of surface-exposed genes and colonization determinants across all sequenced *V. anguillarum* and *V. ordalii* genomes identified not only common genes but also genomic variability, because some genes are restricted to specific serotypes. Our studies will result in an increase of our understanding of the evolution of the *V. anguillarum* genome and will contribute new insights into virulence factors and conserved proteins, which could lead to new avenues in the exploration of targets for vaccine development.

ACKNOWLEDGMENTS

J.H.C. was supported by USDA grant 2007-35600-185-27 and NIH grant RO1-19018, whereas F.L.T., C.C.T., and G.M.D. were supported by grants from CNPq, CAPES, and FAPERJ.

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