O-antigen diversity and lateral transfer of the wbe region among Vibrio splendidus isolates

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Title: O-Antigen Diversity and Lateral Transfer of the \textit{wbe} Region Among \textit{Vibrio splendidus} Isolates

Running Title: O-antigen Diversity Among \textit{Vibrio splendidus}

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Summary

The O-antigen is a highly diverse structure expressed on the outer surface of Gram-negative bacteria. The products responsible for O-antigen synthesis are encoded in the wbe region, which exhibits extensive genetic diversity. While heterogeneous O-antigens are observed within Vibrio species, characterization of these structures has been devoted almost exclusively to pathogens. Here, we investigate O-antigen diversity among coastal marine Vibrio splendidus-like isolates. The wbe region was first identified and characterized using the sequenced genomes of strains LGP32, 12B01, and Med222. These regions were genetically diverse, reflective of their expressed O-antigen. Additional isolates from physically distinct habitats in Plum Island Estuary (MA, USA), including within animal hosts and on suspended particles, were further characterized based on multilocus sequence analysis (MLSA) and O-antigen profiles. Results showed serotype diversity within an ecological setting. Among 48 isolates which were identical in three MLSA genes, 41 showed gpm genetic diversity, a gene closely linked to the wbe locus, and at least 12 expressed different O-antigen profiles further suggesting wbe genetic diversity. Our results demonstrate O-antigen hyper-variability among these environmental strains and suggest that frequent lateral gene transfer generates wbe extensive diversity among V. splendidus and its close relatives.

Introduction

The O-antigen, a polysaccharide chain composed of repeated units of 2-6 sugars, protrudes from the surface of Gram-negative bacteria as the outermost portion of lipopolysaccharide (LPS). This outer membrane structure is in direct physical interaction
with the surrounding substrates and thus subject to environmental selective pressures. Consequently, O-antigens exhibit high diversity in basic composition and shape, largely due to the variation of monosaccharide building blocks, their linkage into repeat units, and the number of units (Reeves et al., 1996; Chatterjee and Chaudhuri, 2004). For example, hundreds of serotypes, or conspecific strains which encode and express distinct O-antigens, have been observed for *Escherichia coli* (Samuel and Reeves, 2003), *Salmonella enterica* (Popoff, 2001), and *Vibrio cholerae* (Chatterjee and Chaudhuri, 2004). This phenotypic diversity manifests in the *wbe* chromosomal region which ranges in size from ~40 to 70 kilobases (kb) reflecting differences in both shared and non-homologous gene content located within *wbe* regions. While shared *wbe* genes differ based on mutations, non-homologous genes result from lateral gene transfer (LGT) (Reeves et al., 1996; Stroeher et al., 1998).

Historically, O-antigen diversity among pathogenic bacteria was proposed to be influenced by selective pressure exerted by the host immune system in which strains expressing rare or novel structures evade immune detection and cause disease (Reeves, 1995). This hypothesis explains O-antigen diversity among pathogens that undergo phase variation which increases bacterial fitness by evasion within a host (Maskell et al., 1991; Meyer, 1991; Lukácová et al., 2008), but fails to explain serotype diversity among other pathogens that express stable O-antigens, such as *E. coli* O157, *S. enterica* serovar Typhi, and *V. cholerae* O1 and O139 which cause bacteremia, typhoid fever and cholera, respectively. Although conspecific strains may carry virulence genes, these serotypes are thought to be non-pathogenic (Guhathakurta et al., 1999; Bakhshi et al., 2008; Rahman et al., 2008; Ottaviani et al., 2009). Moreover, most isolates, including pathogenic ones,
spend the majority of their lifecycle in an environment not attributed to causing disease suggesting that other ecological selective pressures influence O-antigen diversity. For instance, O-antigen diversity among *S. enterica*, which spend most of its time as a gut commensal, may be maintained by intestinal amoeboid predation (Wildschutte et al., 2004; Wildschutte and Lawrence, 2007). Vibrios are marine microbes that have multiple lifestyles and survive either free-living, particle associated, or within animal hosts. Selective pressures may exist such as phage and protist predation, competition for attachment to particulate carbon sources in nutrient deprived waters, or from habitat differences encountered when traveling from hosts to the water column. Thus, knowledge of ecology may be necessary to understand bacterial genetic and structural diversity.

While O-antigen characterization has been well documented among individual pathogenic *Vibrio* strains including *V. cholera* O1 and O139 (Stroeher et al., 1998; Chatterjee and Chaudhuri, 2004), serotype diversity at the population level remains less studied. The *Vibrio splendidus* clade represents the dominant vibrioplankton group in the temperate coastal ocean (Thompson et al., 2004a; Thompson et al., 2004b; Thompson et al., 2005) and has been found free-living and associated with numerous marine substrates including suspended organic particles, zooplankton, mussels and crabs [Preheim et al., submitted; (Thompson et al., 2005; Hunt et al., 2008)]. Since isolates survive in various habitats, O-antigen diversity may persist among strains because certain structures provide fitness benefits against different selective pressures. To initially characterize O-antigen diversity and establish that different serotypes occur among *V. splendidus*-like isolates, we used the published genomes of LGP32, 12B01, and Med222 to identify and define the *wbe* region and show that its genetic diversity reflects O-antigen differences. These
environmental strains were isolated from different geographic locations; LGP32 was isolated from an oyster pond in France, 12B01 from Plum Island Estuary (PIE) of coastal Massachusetts, and Med222 from the Mediterranean Sea (Le Roux et al., 2009). We extended this study to *V. splendidus*-like environmental isolates within the PIE to determine if O-antigen diversity persists among strains within a geographical area but from diverse marine habitats including different body regions of crabs and mussels, and zooplankton. Combined methods of multilocus sequencing analysis (MLSA) and O-antigen profiling were used to show that O-antigen hyper-variability exists among *V. splendidus*-like isolates. Sequence analysis of the *gpm* gene, a housekeeping gene closely linked to the *wbe* locus, was used to investigate LGT about the *wbe* region. Extensive *gpm* genetic divergence as well as phylogenetic incongruencies between MLSA and *gpm* tree topologies, suggest a more frequent transfer of the *wbe* region compared to MLSA housekeeping genes among our environmental isolates and with LGP32, 12B01, and Med222. Together, these methods provide an excellent means for discriminating between closely related isolates and may prove useful in linking bacterial diversity to ecological parameters.

**Results**

**Genetic Diversity of the *V. splendidus wbe Locus.***

The *wbe* loci of the *V. splendidus*-like strains LGP32, 12B01, and Med222 were identified and determined to be bounded by the *gmhD* and *gpm* genes (Figure 1a). The *gmhD* gene product (also referred to as *rfaD*) encodes an epimerase involved in heptose synthesis and is required for core LPS in many Gram-negative bacteria (Coleman, 1985;
Stroeher et al., 1998). Among annotated vibrios, the $gmhD$ ORF has been shown to have strong linkage to the $wbe$ region (Stroeher et al., 1998). Initially using $gmhD$ as a guide, we identified the $wbe$ regions in LGP32, 12B01, and Med222. For each strain, this locus was found on the larger of two chromosomes, which contains core loci involved in cellular processing, signaling, and metabolism (Le Roux et al., 2009). The $wbe$ regions differ in size between strains by almost 20 kb: the 12B01 $wbe$ is the largest at 54.4 kb, Med222 is 43 kb, and LGP32 is 37 kb. Although the ORFs within these regions have predicted functions in the synthesis, linkage, and modification of sugars, the wide range in size is largely due to non-homologous $wbe$ gene content between strains (Figure 1a). While pairwise comparisons of ORFs flanking the $wbe$ region were highly conserved, many ORFs within our predicted $wbe$ region were non-homologous with respect to each region suggesting gain and/or loss through lateral gene transfer and further supporting our identification of each $wbe$ coding region.

Homologous ORFs were identified within the $wbe$ region between strains with LGP32 as a reference (Table 1). Separate analyses were conducted using 12B01 or Med222 as the reference (Tables S1 and S2). Three gene groups show similarity among the strains (indicated by gray shading in Figure 1a; also refer to Table 1). The first group (I) is represented in LGP32 as ORFs labeled 1-7. Group I ORFs, which include the $gmhD$ gene required in LPS synthesis (see above) were found in all three strains, suggesting conserved functions among these strains. Other predicted Group I gene products include a regulator and a transferase. Given their conserved location relative to $gmdH$, these may be involved in assembling heptose into core, which was found in LPS from all three strains (Table 2). Group II (LGP32 ORFs 12-14), is shared between 12B01 and LGP32.
Gene products in this group have proposed functions in polysaccharide export. Interestingly, these ORFs were not identified in Med222, suggesting this strain uses a different system for O-antigen export. Finally, Group III (LGP32 ORFs 26-29), has homologues in both 12B01 and Med222; however, in these strains the ORFs are not adjacent to one another. Within Med222 Group III ORFs are represented as ORFs 8, 9, 13, and 14. In 12B01, these ORFs are observed twice, at ORFs 9-12 and 45-48, suggesting a duplication event or two independent transfers. The predicted functions of these genes are involved in the glucose and rhamnose synthesis pathways, which we verified to be incorporated into the O-antigen of each strain (Table 2). Besides these similarities, most ORFs among LGP32, 12B01 and Med222 are non-homologous genes with respect to each wbe region, and likely encode different proteins that help assemble diverse O-antigens. Taken together, our results indicate the overall wbe composition is diverse among these closely related strains.

The wbe loci of Gram-negative bacteria are typically marked by JUMP (Just Upstream of Many Polysaccharide regions) sites, which include a short conserved signal sequence for DNA uptake and are thought to be involved in LGT during transformation of competent cells (Hobbs and Reeves, 1994; Snyder et al., 2007). These short conserved sequences reside just prior to wbe regions of other vibrios (González-Fraga et al., 2008). Genome searching revealed JUMP sites to be exclusively located within our defined wbe region of LGP32, 12B01 and Med222, just prior to a series of ORFs transcribed in one direction (Figure 1b). The LGP32 JUMP site is located downstream of putative O-antigen transporter genes. In 12B01 and Med222, this sequence is immediately upstream of ORFs 8 and 9, respectively. Interestingly, 12B01 has another very similar JUMP sequence just
upstream of ORFs 45-48 which is homologous to the ORFs 9-12 (Figure 1a). The conserved JUMP site sequence and its location just prior to wbe gene clusters transcribed in the same direction suggest that these sites are involved in the transfer of multiple wbe encoded genes during a single LGT event.

**O-Antigen Structural Variability Reflects wbe Genetic Diversity.**

In other vibrios, the wbe gene region has been shown to encode proteins responsible for O-antigen synthesis (Stroeher et al., 1998; Chatterjee and Chaudhuri, 2004). Different structures are phenotypically manifested through the incorporation of dissimilar monosaccharides and their linkage into polysaccharide units. Thus, variation in wbe gene content (i.e., ORFs encoding monosaccharide synthesis, transferases, and transporters) is likely to influence the O-antigen expressed by a strain. Given the observed wbe genetic diversity between LGP32, 12B01, and Med222 (Figure1a and Tables 1, S1 and S2), we next analyzed the LPS core and O-antigen expressed by each strain through silver staining. This method allows visualization of differences in O-antigen repeat units through differential banding patterns, such that different profiles represent dissimilar O-antigens. Different O-antigen profiles were observed among LGP32, 12B01 and Med222 indicating each strain is of a distinct serotype (Figure 1c).

To address whether the differences in O-antigen profiles could be attributed to the inclusion of monosaccharides unique to each strain, the glycosyl residues belonging to the LGP32, 12B01 and Med222 O-antigens were determined through combined gas chromatography and mass spectrometry (Merkle and Poppe, 1994). For all strains, we were able to identify monosaccharides common to the LPS core (heptose and glucose), and those typically included in the O-antigen (galactose, rhamnose and ribose) (Table 2)
Overall, these shared residues represent most of the conserved regions among LGP32, 12B01, and Med222 (Figure 1a and Table 1). We also detected residues not shared by all strains. For example, glucuronic acid, which has been shown to be included in the O-antigen of other Gram-negatives (Samuel and Reeves, 2003; Chatterjee and Chaudhuri, 2004), was detected in 12B01, and an unidentified amino sugar was unique to Med222 (Table 2). These residues are likely to contribute at least partially to the observed differences in O-antigen structures (Figure 1c). Together, these results support that wbe genotypic diversity contributes to phenotypic diversity between serotypes.

Serotype Diversity Among Closely Related V. splendidus-like Isolates.

O-antigen diversity was observed among V. splendidus-like strains LGP32, 12B01 and Med222 (Figure 1) which were originally isolated from diverse geographical regions (Table S3) (Le Roux et al., 2009). Our recent study of population-level diversity among vibrios in the PIE affords the opportunity to determine O-antigen diversity among closely related, co-existing strains (Preheim et al., submitted). We chose 114 representatives within the V. splendidus clade from several marine habitats (Table S3), including zooplankton, crabs, and mussels (Pacocha, et al. 2010) to investigate serotype diversity.

As an estimate of overall relatedness of these 114 strains, concatenated nucleotide sequences of the adk, hsp60, and mdh housekeeping genes were used for MLSA and a maximum likelihood tree was generated (Figure 2a). Isolates had either different sequence types (ST) (n=37) meaning they were closely related based on nucleotide changes within the genes used for MLSA or they shared a ST with another strain (n=77) suggesting genetically identical or clonal isolates. Overall, we observed relatively little
genetic divergence among all these strains, as inferred from branch lengths and position
relative to LGP32, 12B01 and Med222, thus limiting the time-scale for genome and O-
antigen variation to accrue. Serotype diversity was characterized by visualizing O-antigen
profiles for a total of 53 PIE isolates consisting of 37 with different STs as well as 16 that
shared STs (Figure 2a); another 61 isolates that shared either ST 3, 12, or 243 were
characterized in separate analyses (see below). Silver staining revealed at least 9 different
O-antigen structures from these isolates (Figures S1), and in some cases isolates of the
same ST were found to express different structures (for example, 9ZC32 and 9ZC73;
9CH134 and 9CHC140), thus confirming multiple serotypes of closely related V.
splendidus-like strains are present within PIE.

Given the number of O-antigen structures observed within the closely related
isolates, and that some strains with the same ST showed different O-antigen profiles
(Figures S1 and 2a), we next examined serotype diversity among strains having the same
ST to further constrain the time scale of O-antigen variation. Strains with ST 3 (n=25)
and ST 12 (n=23) were isolated from multiple habitats including crabs, mussels, and
zooplankton while ST 243 (n=13) originated from one individual crab. Silver staining
was used to examine serotype diversity among isolates belonging to each ST (Figure 3).
For ST 243 we observed no differences in O-antigen banding patterns (Figure 3a)
possibly due to clonal expansion within a single host specimen. Surprisingly, a variety of
O-antigen profiles were observed for ST 3 (Figure 3b and c) and 12 (Figure 3d and e),
and overall we estimate at least 12 different serotypes within these groups alone. This is
interpreted as a conservative estimate since profiles that appear similar may not
absolutely represent the same O-antigens. These results demonstrate that diverse V.
splendidus-like serotypes occur within PIE, and further suggest O-antigen hyper-
variability among strains, as isolates of identical ST can express distinct structures.

Genetic Diversity of the gpm Gene and wbe LGT.

Our sequence analysis of the LGP32, 12B01 and Med222 genomes led to the
identification of putative JUMP sites within the wbe regions (Figure 1a and b), which
have been implicated in LGT between other bacteria (Hobbs and Reeves, 1994). As a
means to investigate if LGT is a possible mechanism of wbe diversity and to discriminate
between serotypes having the same ST, we performed a phylogenetic analysis of gpm, a
housekeeping gene in close linkage to wbe and required in glycolysis (Figure 1a). Overall,
we observed a ~6-fold increase in divergence of gpm coding sequence (4.6%), compared
to MLSA divergence (0.81%) among our environmental strains (Table S4). Furthermore,
81 unique STs were observed among the sample set using gpm sequence analysis while
only 37 unique STs were identified based on adk, hsp60 and mdh alone.

Using an approach similar to our initial MSLA with the adk, hsp60 and mdh
genes, we generated a gpm-based maximum likelihood tree as a means to infer strain
relatedness with respect to wbe (Figure 2b). Compared to the MLSA generated tree, the
gpm gene tree exhibits longer branch lengths as a result of increased gpm nucleotide
change. More importantly, the topology of the gpm tree (Figure 2b) differs from the
MLSA tree (Figure 2a). For instance, strains with identical sequences based on MLSA
genes (ST 3 and 12) are scattered throughout the gpm based tree, showing they are not
genetically identical even if they have similar O-antigen profiles; and LGP32, 12B01 and
Med222 appear more closely related to environmental isolates in the gpm-based tree as
opposed to MLSA. In addition, strain 9CSC152 is more closely related to the SWAT3
outgroup based on the gpm gene (Figure 2b) than to the PIE environmental isolates (Figure 2a). These results suggest that wbe transfer occurs frequently across the V. splendidus clade. Of the 61 strains belonging to STs 3, 12, and 243, a total of 41 unique gpm sequences were observed: 24 of 25 isolates for ST 3, and 17 of 23 for ST 12 (Figure 2b). Identical strains based on gpm were mostly of ST 243, again suggesting clonality. In combination with the identification of wbe JUMP sites within the available V. splendidus-like genomes, O-antigen hyper-variability among PIE isolates with the same STs, and gpm gene diversity along with tree incongruencies between MLSA and gpm sequences, these data indicate frequent LGT of wbe loci within the PIE marine column resulting in multiple V. splendidus-like serotypes.

Discussion

Marine bacteria constantly encounter diverse habitats while carried through the water column. Ecological selective pressures ranging from predation to surface adherence likely exist on spatial scales and may influence O-antigen diversity among serotypes. The ability to change an O-antigen through LGT of the wbe region may offer advantages in fitness across diverse environments. Using the sequenced genomes of LGP32, 12B01, and Med222, we showed characteristics of LGT such as non-homologous genetic differences between strains and the presence of JUMP sites, which are believed to facilitate wbe gene transfer (Figure 1). With the acquisition of wbe regions, entire functional pathways involving the synthesis of different O-antigen structures can be gained with the potential result of serotype conversion. We have previously shown that V. splendidus is found in different marine environments such as free-living within the water
column, attached to suspended particles, and on marine hosts [Preheim et al., submitted, (Thompson et al., 2005; Hunt et al., 2008)]. We suggest that the acquisition and expression of different wbe regions among V. splendidus and its close relatives could influence bacterial fitness through environmental interactions by the O-antigen resulting in the maintenance of O-antigen diversity.

To investigate LGT among environmental V. splendidus-like strains, we chose closely related and even identical strains based on MLSA to constrain O-antigen variability. Related strains were on average 0.81% divergent based on the concatenated adk, hsp60, and mdh sequences consisting of 1254 base pairs (Table S4), while strains having the same sequences (such as ST 3 and 12 strains) were devoid of mutations. Even with this mutational constraint, extensive genetic diversity was observed in the gpm gene amongst ST 3 and ST 12 isolates—an average and maximum nucleotide divergence of gpm was 5.25% and 13.5%, respectively. Because gpm is closely linked to the wbe region (Figure 1), selective sweeps are precluded and gpm diversity is likely maintained through hitchhiking with the wbe locus. Furthermore, incongruencies between MLSA and gpm phylogenies (Figure 2) and the presence of disparate O-antigens within a ST (Figure 3d and e) suggest that LGT occur at the wbe chromosomal location.

High rates of transfer within the wbe region, as suggested by extensive genetic diversity of the gpm gene, provide a means for serotype selection. We did not observe a predominant serotype among or within hosts (except for one crab where clonal expansion of strains with ST 243 is evident) which supports our recent study that V. splendidus are generalists among invertebrate hosts (Preheim et al., submitted). However, we did observe closely related serotypes (expressing the same O-antigen) among different hosts.
For instance, strains 9CS34 (ST 12), 9CG23 (ST 3), and 9CSC94 (ST 3) from crab specimens 2, 2, and 5, respectively, were of the same serotype; and 9CG33 (ST 12), 9MHC17 (ST 12), and 9MHC23 (ST 12) from different mussels and a crab were of another serotype. If O-antigen selection occurs in the water column, prior to association within a host, then closely related serotypes could be found dispersed among different marine invertebrates. Continued studies to identify possible selective pressures influencing O-antigen and wbe diversity in the marine environment are being investigated.

Serotypes expressing the same O-antigen usually have different gpm sequences resulting from mutations within gpm or because of its close linkage to wbe making it susceptible to lateral transfer while preventing gpm selective sweeps. However, it is possible for strains to have the same gpm sequences yet dissimilar O-antigens. We amplified 483 base pairs of the gpm gene starting 168 base pairs downstream from the start site; if recombination occurs before the amplified region or within the wbe locus, then gpm gene sequences may be identical. For instance, a group of six strains from crab specimen 7 were identical based on gpm gene analysis (Figure 2b). It was expected that these were clonal isolates because all were of ST 12 based on MLSA; however, the O-antigen profiles among these strains differ. For example, 9CHC127, 9CHC133, and 9CS146 show one profile, while 9CSC139, 9CSC158, 9CS151 show another (Figure 3d and e). This is also seen with (1) 9CS134 and CS126 and (2) 9CS24 and 9MG29 which have the same gpm sequence but dissimilar O-antigen profiles. These results suggest that LGT occurred within the wbe region without involving the gpm gene. Furthermore, we would predict that genetic diversity of genes surrounding the wbe region would decrease with distance from the wbe locus if this region is under strong selection.
Our results suggest that the O-antigen hyper-variability observed among environmental *V. splendidus*-like serotypes reflects LGT-driven diversity of the *wbe* region. Frequent *wbe* transfer is evident among these strains and as well as the more distantly related LGP32, 12B01, Med222, all within the *V. splendidus* clade. The selective pressures that maintain O-antigen diversity remain unknown but may be related to phage infection, protist predation, or ecological interactions during life history in the water column. MLSA approaches that include loci with hyper-variable outer membrane structures have improved capacity to discriminate among otherwise identical STs and can provide greater insight into ecologically relevant differentiation among closely related strains.

**Experimental Procedures**

**Strain Isolation and Growth Media.**

Water samples and invertebrates were collected from Plum Island Sound Estuary, Ipswich, MA in the spring and fall of 2008 as described in (Preheim et al., submitted). Briefly, seawater samples were collected at high tide in 4 L bottles from the shore. Zooplankton was isolated by filtering 100 L of seawater through a 64 µm mesh net. Samples were rinsed three times with sterile seawater, washed into a 50 ml conical tube and kept at ambient temperature in the dark until processing ~2 hours later. Living and dead zooplankton were differentiated by eye under a dissecting microscope based on movement and 10-140 individuals of each category were picked from each 100 L concentrate. Collections also included four male green crabs (*Carcinus maenas*); eight male and four female shore crabs (*Hemigrapsus sanguineus*; and sixteen blue mussels
(Mytilus edulis). All animals were washed with sterile seawater and placed in a whirl
pack and cooler until processing. For crabs, gill (one brachia), stomach (entire tissue)
and hindgut (~4 cm beginning with anus) were collected following stunning prior to
dissection (no anesthesia). For mussels, approximately 1.5 cm² of gill and hindgut
(including the anus) tissue was collected. For both crabs and mussels, gastrointestinal (GI)
contents were collected by flushing tissue with 4 ml sterile seawater with a syringe.
Tissues were washed 3x with sterile seawater to ensure only attached bacteria were
collected. Crab and mussel tissue and GI tract contents samples were homogenized in a
tissue grinder, serially diluted (10- to 10,000-fold) in sterile seawater, and plated for
isolation on Vibrio-selective marine TCBS media (BD Difco TCBS + 1% NaCl). A total
of 160 isolates were picked from each sample type per season (20 per specimen) using
the most dilute samples with sufficient growth. Isolated colonies were re-streaked 3x
alternating 1% TSB media (BD Bacto + 2% NaCl) and marine TCBS media to ensure
purity of isolates.

**PCR Amplification for MLSA and gpm Analysis.**

Partial amplification of the heat shock protein (hsp60), adenylate kinase (adk), and malate
dehydrogenase (mdh) genes were performed with all isolates for MLSA. Primers were as
follows:  

**adk**, 5’GTATTCCACAAATYTCTACTGG3’ and 5’GCTTCTTTACCGTAGTA3;  

**hsp60**, 5’GAATTCGAIIIIICIGGIGAYGGIACIACIAC3’ and 5’CGCGGGATCCYKIYKITCICCRAAICCIIGGICYTT;  

**mdh**, 5’GATCTCGAGYCATATCCWCAC3’ and 5’GCTTCWACMACYTCRGTACCCG3’.

PCR amplification was carried out as previously described with annealing temperature at
41°C for *adk* and *hsp60* and 60°C for *mdh* and sequences were submitted to GenBank (Preheim *et al.*, submitted). For *gpm* gene analysis, partial gene amplification was performed using primers 5’GATGGYCAAATGGGTAACTC3’ and 5’CAGCACGGTAGTTCATGAAG3’. PCR amplification was carried out for 30 cycles with an annealing temperature of 60°C. Amplicons were sequenced bidirectionally at the Bay Paul Center at the Marine Biological Laboratory, Woods Hole, MA with the same primers for each respective gene. *gpm* sequences were submitted to GenBank under accession numbers GU990234-GU990351.

**Phylogenetic Tree Construction and Gene Divergence.**

Concatenated *adk*, *hsp60*, and *mdh* for MLSA and single *gpm* gene sequences were used to generate sequence alignments and gene divergence matrices using default parameters in ClustalX. The Vibionales bacterium SWAT-3 was used as the outgroup. Maximum likelihood trees were constructed from the alignment using PhyML set with HKY85 substitution parameters (Guindon and Gascuel, 2003). Bootstrapping was performed in 100 replicates and values >70% are shown.

**Whole Cell Lysates and Silver Stain for Estimation of O-antigen Diversity.**

Strains were grown overnight in 5 mL of TSB at room temperature (RT). When cultures reached an OD$_{600}$ of 1.0, 1 mL of cells were aliquoted and spun at 13,000 rpm for 3 min. Cells were resuspended in 100 μl lysis buffer (1M Tris HCl; pH 6.8, 2% SDS; 4% β-mercaptoethanol; 10% glycerol), incubated at 100°C for 10 minutes, and then cooled to below 60°C. Lysates were treated with 1.3 μl of 20 mg/ml of proteinase K and incubated for 1 hr at 55°C. Bromophenol blue was added to each lysate for visualization, and 14 μl each loaded to a precast 10-20% tricine Novex gel. Following electrophoresis, gels were
silver stained as previously described to visualize the O-antigen (Hitchcock and Brown, 1983). Briefly, each gel was fixed in 40% ethanol and 5% acetic acid for 1 hr, oxidized in the fixative with 0.7% periodic acid, and then incubated in silver stain (0.6% silver nitrate; 0.14 M NaOH; 1 ml 37% ammonium hydroxide) for 10 min. Gels were developed by incubation for 2 min in developing buffer (50 μM citric acid and 0.7% formaldehyde in 200 ml volume) at 40°C. All gels were repeatedly washed after each step as described.

**O-Antigen Glycosyl Composition Analysis.**

Procedures were carried out as previously described (Merkle and Poppe, 1994). Single colonies of LGP32, 12B01, and Med222 were picked and grown overnight at RT in 75 ml of Difco TSB media. Cultures were pelleted by centrifugation for 10 min at 10,000 rpm and resuspended in 1 ml of water. Five 5 ml of 95% ethanol was added and cells were incubated at room temperature for an hour. Each cell suspension was pelleted and the supernatant removed. The Complex Carbohydrate Research Center at the University of Georgia determined core and O-antigen glycosyl residues after acid hydrolysis of purified LPS.

**Analysis of the wbe region.**

The gmhD gene from *V. cholera* strain N16961 (locus tag VC0240) was used to determine the presence and location of gmhD gene in the sequenced genomes of *V. splendidus*-like strains LGP32, 12B01, and Med222. We identified the gmhD gene in each genome screened (LGP32, YP_002415885; 12B01, ZP_00989916; and Med222, ZP_01065583) and used its location as a reference point to manually analyze adjacent open reading frames (ORFs) for predicted functions involved in synthesis, linkage, and
modification of sugars. ORFs bounded by \textit{gmhD} and \textit{gpm} represented the \textit{wbe} region of each strain.

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\section*{References}


Figure Legends

Figure 1. The *wbe* genotypic and O-antigen phenotypic diversity of *V. splendidus* strains 12B01, Med222, and LGP32. (A) The regions exhibit extensive genetic diversity between the *gmhD* and *gpm* flanking genes. Each *wbe* region encodes similar and different genes whose putative functions are O-antigen construction. Rectangular boxes represent ORFs. ORFs depicted above and below the respective genome baseline indicated forward and reverse transcription, respectively. Grey lines between genomes indicate homology between those genes. Black bars above LGP32 ORFs identified as I, II, and III indicated regions of shared homology with other *wbe* loci. Open and closed circles represent JUMP sites. (B) JUMP sites shown for *V. cholera* 01, 12B01, Med222, and LGP32 which contains the conserved DNA uptake signal sequenced (USS). Circles represent respective JUMP sequence locations in the *wbe* region. Bold and shaded sequences represent the conserved USS in *V. cholera* and *V. splendidus*, respectively. (C) Silver stain showing different O-antigen profiles; lanes 1, molecular marker; 2, *Salmonella enterica* LT2; 3, *Escherichia coli* K12; 4, 12B01; 5, Med222; and 6, LGP32.

Figure 2. Maximum likelihood trees of *V. splendidus*-like strains isolated from different marine habitats. (A) Phylogenetic relatedness based on MLSA of concatenated *adk, mdh* and *hsp60* partial gene sequences consisting of 1254 base pairs. The strains with ST 3, 12 or 243 are boxed grey and their ST# is present after their strain name. Sequenced genomes are bolded and marked with a *. (B) Phylogenetic relatedness based on *gpm* partial gene sequence consisting of 483 base pairs. Strains are labeled
according to the season and animal sample of isolation: Fall and spring is designated by 9 or 4, respectively, and the specific animal sample is identified by CG, crab gills; CH, crab intestines; CHC, crab intestinal lining; CS, crab stomach; CSC, crab stomach lining; MHC, mussel intestinal lining; ZC, zooplankton. Colors represent the individual host or zooplankton sample they were isolated from.

Figure 3. Silver stains showing the O-antigen profiles of V. splendidus-like environmental isolates. (A) Strains isolated from an individual crab host having MLSA ST 243 express the same O-antigen. (B and C) ST 3 and (D and E) ST 12 strains isolated from either crabs, mussels, or zooplankton show similar and different O-antigen profiles. Strains were isolated from different individual hosts as indicated by numbers and strain nomenclature, as described in Figure 2.