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Ecology of *Vibrio parahaemolyticus* and *Vibrio vulnificus* in the Coastal and Estuarine Waters of Louisiana, Maryland, Mississippi, and Washington (United States)

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Johnson, C. N., Bowers, J. C., Griffitt, K. J., Molina, V., Clostio, R. W., Pei, S., Laws, E., Paranjpye, R. N., Strom, M. S., Chen, A., Hasan, N. A., Huq, A., Noriea, N. F., Grimes, D. J., Colwell, R. R. (2012). Ecology of *Vibrio parahaemolyticus* and *Vibrio vulnificus* in the Coastal and Estuarine Waters of Louisiana, Maryland, Mississippi, and Washington (United States). *Applied and Environmental Microbiology*, 78(20), 7249-7257.

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- 1 Ecology of Vibrio parahaemolyticus and Vibrio vulnificus in the coastal and estuarine
- 2 waters of Louisiana, Maryland, Mississippi, and Washington, United States
- 3
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- 16 Running Title: Coastal ecology of the vibrios
- 17 Key Words: Vibrio parahaemolyticus, Vibrio vulnificus, Crassostrea virginica, water,
- 18 sediment, thermostable direct hemolysin, *tdh*-related hemolysin, temperature, turbidity,
- 19 salinity, plankton
- 20
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- 24 Submitted to: Applied and Environmental Microbiology
- 25

26 ABSTRACT

27

28 Vibrio parahaemolyticus (Vp) and Vibrio vulnificus (Vv), native to estuaries globally, are 29 agents of seafood-borne or wound infections, both potentially fatal. Like all vibrios 30 autochthonous to coastal regions, their abundance varies with changes in 31 environmental parameters. Sea surface temperature (SST), sea surface height (SSH), 32 and chlorophyll have been shown to be predictors of zooplankton and thus factors 33 linked to vibrio populations. The contribution of salinity, conductivity, turbidity, and 34 dissolved organic carbon to the incidence and distribution of Vibrio spp. has also been reported. Here, a multi-coastal, 21-month study was conducted to determine 35 36 relationships between environmental parameters and V_P and V_V populations in water, 37 oysters, and sediment in three coastal areas of the United States. Because ecologically 38 unique sites were included in the study, it was possible to analyze individual parameters 39 over wide ranges. Molecular methods were used to detect thermolabile hemolysin *tlh*, 40 thermostable direct hemolysin tdh, and tdh-related hemolysin trh, as indicators for Vp 41 and hemolysin vvhA for Vv. SST and suspended particulate matter were found to be 42 strong predictors of total and potentially pathogenic Vp and Vv. Other predictors included chlorophyll-a, salinity, and dissolved organic carbon. For the ecologically 43 44 unique sites included in the study, SST was confirmed as an effective predictor of 45 annual variation in vibrio abundance, with other parameters explaining a portion of the 46 variation not attributable to SST.

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49 INTRODUCTION

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51 It has long been established that *Vibrio* spp. are autochthonous to the marine, 52 estuarine, and riverine environment. Vibrios cultured from environmental samples 53 commonly lack genes coding for functions associated with pathogenicity for humans 54 and marine animals, e.g., the thermostable direct hemolysin (tdh) in Vibrio 55 parahaemolyticus. Yet, pathogenic subpopulations of vibrios are potential agents of 56 disease outbreaks and pandemics (7, 19, 23, 37, 44, 50, 65), notably in developing 57 countries where access to safe drinking water is limited (26, 56) and/or in countries where consumption of raw or undercooked shellfish is common (11, 80). Vibrio 58 59 parahaemolyticus is most frequently associated with gastroenteritis and has been linked 60 to annual outbreaks (7, 8, 44). Vibrio vulnificus is more frequently associated with 61 wound infections, with a case fatality rate as high as 50% (5, 10, 27). The abundance 62 and distribution of these three human pathogens have been linked to environmental 63 factors most notably temperature and salinity, depending on the pathogen and its 64 habitat, and the geographic location (4, 13, 14, 18, 24, 29, 31, 35, 39, 70, 72, 83). Dissolved oxygen (30, 54, 58), chlorophyll (6, 20, 31, 33), and plankton (2, 31, 41, 59, 65 66 74) have also been found to be important in describing the ecology of vibrios. 67 Regulatory authorities responsible for oversight of recreational waters and shellfish 68 harvesting areas employ rainfall, fecal coliform counts, river stages, and, more recently, 69 enterococcus counts to determine opening and closing of specific areas to protect 70 public health (21, 25, 62, 76). Standard microbiological approaches to classification and opening/closing of oyster harvest areas, unfortunately not useful for control of exposure 71

72 to pathogenic Vibrio, spp., continue to be used and are generally accepted for

regulating exposure to other pathogens in the U.S. (36).

74 Naturally occurring pathogens, notably vibrios, are ubiquitous in the aquatic 75 environment and contribute to carbon and other nutrient cycling (24, 61). Clearly, 76 human exposure to these pathogens cannot be completely eliminated, but incidence of 77 illness can be reduced if environmental conditions that significantly elevate risk could be 78 identified and monitored. Communication of such conditions to stakeholders (regulatory 79 agencies, the shellfish industry, public health officials, at-risk consumers, etc.) would 80 reduce exposure and subsequent disease. An informative, robust system of 81 identification of conditions associated with high risk requires quantifying the association 82 of environmental factors with abundance of total vibrio populations and potentially 83 pathogenic vibrios. Given proven associations as predictors of vibrio abundance the 84 relevant environmental data can be collected by satellite remote sensing (13, 39, 83). 85 Development of models to predict presence of vibrio populations is facilitated by 86 collecting observations over a range of environmental parameters and recognition that 87 predictive relationships may vary across regions due to differences in ecology (for 88 example, models developed for the Gulf of Mexico may not be applicable to the Pacific 89 Northwest). Furthermore, potentially pathogenic subpopulations of environmental 90 vibrios are not necessarily a constant proportion of the total vibrio population (17, 18, 91 31, 32, 55, 83). Here we describe an analysis of environmental factors providing the 92 potential for improving upon existing predictive models for V. parahaemolyticus and V. 93 vulnificus. Specifically, we determined densities of total V. parahaemolyticus (tlh), and 94 potentially pathogenic V. parahaemolyticus, as indicated by the presence of the

thermostable direct hemolysin (*tdh*) and *tdh*-related hemolysin genes (*trh*) (48, 64), and total *V. vulnificus* as indicated by the presence of the *V. vulnificus* hemolysin gene (*vvhA*) that has been used as a marker for the species (45, 81, 82). These were determined for water, oyster, and sediment samples collected at sampling stations located in the Pacific Northwest, northern Gulf of Mexico, and Chesapeake Bay.

100

101 MATERIALS AND METHODS

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103 Sample collection and processing. From December, 2008 to August, 2010, 104 water, oyster, and sediment samples were collected in the Pacific Northwest in Hood 105 Canal in Washington (WA), in the northern Gulf of Mexico spanning Louisiana (LA) and 106 Mississippi (MS), and in the Chester River and Tangier Sound of the Chesapeake Bay 107 in Maryland (MD). All samples were collected concurrently in LA, MS, and MD, but in 108 WA, because of logistical problems, oyster and sediment samples were collected inter-109 tidally and relayed to a dock where the water samples were collected sub-tidally. At all 110 sampling stations, 6 – 12 L of water, 20 – 25 oysters, and 100 g of sediment were 111 collected and transported to the laboratory in coolers containing ice or ice packs. 112 Water, oyster, and sediment samples were processed as described elsewhere (31). 113 Specifically, water samples were shaken as previously described (1), oysters were 114 scrubbed, shucked, and homogenized, and pore water was decanted from sediment 115 then diluted 1:1 and shaken as previously described (31). 116 V. parahaemolyticus and V. vulnificus were enumerated as follows: 1 mL water,

- 117 0.1g and 0.01g oyster, and varying wet weights of sediment (0.0005 0.1g) were
- 5

spread plated on T₁N₃ agar (1% tryptone, 3% NaCl, pH 7.2) and VVA agar (2% 119 peptone, 3% NaCl, 1% cellobiose, 0.06% bromthymol blue, pH 8.2). Detection of tdh+ 120 and trh+ V. parahaemolyticus was accomplished using 1L, 100mL, and 10mL water 121 enrichments and 10g and 1g oyster enrichments in 10X alkaline peptone water (10X 122 APW, 10% peptone, 1% NaCl, pH 8.5). All samples were incubated at 33 °C for 16 – 18 123 hours, as described previously (31).

124 Enumeration of vibrios. To enumerate V. parahaemolyticus and V. vulnificus 125 by direct plating / colony hybridization (DP/CH), Whatman 541 ashless filters (Whatman, 126 Kent, ME) were used to lift bacterial colonies from plates, as described elsewhere (31). 127 The filters were probed using alkaline phosphatase-conjugated oligonucleotide probes 128 (DNA Technology A/S, Risskov, Denmark) specific for vvhA, tdh, trh, and tlh (31). The 129 DP/CH method is most effective for directly enumerating *tlh* and *vvh*, but *tdh* and *trh* 130 populations are often too sparse to rely on DP/CH for enumeration. Therefore, samples 131 were enriched in APW as described above to increase the tdh and trh populations to 132 higher levels that could be detected and enumerated using the most probable number 133 (MPN) method (49). For both total (th+) and potentially pathogenic (tdh+ and trh+) V. 134 parahaemolyticus, serial MPN dilutions were determined in triplicate for water and 135 oyster enrichments using real-time PCR, as previously described (31, 49). For samples 136 collected in Louisiana and in Mississippi, probes and equipment were used that have 137 been described elsewhere (31). For MD samples, probes were used as described 138 previously (49) and reactions were carried out using an AB 7500 thermal cycler (Applied 139 Biosystems, Carlsbad, CA). For WA samples, a Stratagene Mx300Sp Real-Time PCR 140 System (Agilent Technologies, Santa Clara, CA) was used for real-time PCR analysis.

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| | 141 | |
|------------------|-----|---------------------------|
| | 142 | DNA Technologies (Cora |
| | 143 | Applied Biosystems (Fos |
| ini ⁻ | 144 | Brilliant Multiplex QPCR |
| lo I | 145 | following reaction compo |
| Õ | 146 | primers at 75 nM, and B |
| e d c | 147 | remainder of the reactior |
| ahe | 148 | described above, nuclea |
| e (| 149 | profile employed through |
| nlin | 150 | 95°C, followed by 40 cyc |
| Ō | 151 | annealing/extension step |
| Jec | 152 | amplification cycle. The |
| lis | 153 | described previously (49 |
| duc | 154 | All sediment samp |
| S | 155 | pathogenic vibrios in sed |
| 0 | 156 | sediment has previously |
| cc | 157 | and they could therefore |
| \triangleleft | 158 | analysis of sediment san |
| \mathbb{N} | 159 | (data not shown). V. vul |
| \mathbb{Z} | 160 | among the 174 water, oy |
| | 161 | V. vulnificus was detecte |
| | | |

| 141 | The <i>tlh</i> and internal amplification control (IAC) probes were purchased from Integrated |
|-----|---|
| 142 | DNA Technologies (Coralville, IA), and the tdh and trh probes were obtained from |
| 143 | Applied Biosystems (Foster City, CA). Each 25- μ l reaction consisted of 12.5 μ l of 2X |
| 144 | Brilliant Multiplex QPCR Master Mix (Agilent Technologies, Santa Clara, CA) and the |
| 145 | following reaction components (final concentrations): all three probes at 150 nM, all six |
| 146 | primers at 75 nM, and BSA (New England Biolabs, Beverly, MA) at 400 ng/µl. The |
| 147 | remainder of the reactions consisted of 1 μI of the IAC template at the concentration |
| 148 | described above, nuclease-free water, and 5 μI of template. The two-step thermal |
| 149 | profile employed throughout the study consisted of an initial 9.5 min denaturation step at |
| 150 | 95°C, followed by 40 cycles of 30s denaturation at 95°C and a 45s combined |
| 151 | annealing/extension step at 58°C. Fluorescence data were collected at the end of each |
| 152 | amplification cycle. The primer and probe sequences employed were the same as |
| 153 | described previously (49). |
| 154 | All sediment samples were analyzed by DP/CH alone to enumerate total and |
| 155 | pathogenic vibrios in sediment. The PCR/MPN method was not used because |
| 156 | sediment has previously been found to contain very high levels of all four gene targets, |
| 157 | and they could therefore be effectively enumerated using DP/CH; in addition, PCR |

nples has proven unfruitful based on our previous experiences

nificus levels were consistently low in the Pacific Northwest;

vster, and sediment samples collected there during this study,

ed in only one water and in one sediment sample. Thus,

162 determinations of vvhA densities in WA samples were excluded from statistical

analyses. In addition, during analyses of samples from the Pacific Northwest using the 163

DP/CH method, cross-reactivity was noted with either another *Vibrio* species or an unknown bacterium. *V. parahaemolyticus tdh* and *trh* data from oyster and water analyses were therefore excluded and only PCR/MPN data from WA were used for water and oyster samples. To correct for possible cross-reactivity at other sites, the DP/CH data were excluded from the analyses for the GC and MD sites when calculating *tdh* and *trh* densities. The resulting data pools are listed in Table 1.

170 Environmental parameters. At each collection site, water temperature and 171 salinity were measured at the surface and bottom using a digital handheld conductivity 172 meter (model 30-25FT, Yellow Springs Instruments, Yellow Springs, OH). Chlorophyll-a 173 was measured by high-performance liquid chromatography at the University of Hawaii 174 as follows. Triplicate volumes of up to 200 mL were filtered using 25 to 47mm diameter 175 GF/F filters (Whatman, Kent, ME). Filters were stored at -20 °C until shipped overnight 176 on dry ice to the University of Hawaii where concentrations of chlorophyll-a (chl-a) were 177 measured in methanol extracts on a Cary model 50 UV-visible spectrophotometer, as 178 described previously (38). Suspended particulate matter (SPM) was measured by 179 weighing pre-dried GF/F filters using a high-precision scale and filtering up to 200 mL 180 water; the filters were dried overnight at 65 °C and re-weighed. 181 To determine dissolved organic carbon (DOC) concentration, triplicate water 182 samples were pre-filtered using an acrodisc (Pall Acrodisc PSF GHP 0.45 µm) and

183 treated with HCl to convert inorganic carbon to CO₂. The samples were stored at -20 °C

184 until analysis using the method described as follows. CO₂ from inorganic carbon was

185 manually purged from samples by adding additional HCI followed by overnight

186 incubation at room temperature. Total organic carbon was measured using a Shimadzu

TOC-V CSN carbon analyzer equipped with an ASI-V autosampler (Shimadzu Scientific
 Instruments, Columbia, MD).

189 Statistical analyses. Multi-level generalized linear mixed models (GLMM) were 190 used to estimate the distribution of vibrio abundance in ovster, sediment, and water and 191 the relationship between abundance and environmental predictors. Underlying (latent) 192 distributions of vibrio abundance were assumed to be lognormal with mean log₁₀ 193 densities generally presumed to be linearly related to environmental parameters being 194 considered as predictors of abundance. However, given the wide range of salinities 195 observed across sampling locations and consequent likelihood of a nonlinear 196 dependence, a quadratic polynomial was used to model the effect of salinity. Estimates 197 of location and scale of latent distributions of abundance for each combination of gene 198 target, sample type, and sampling location were obtained by fitting null (intercept-only) 199 models with no predictor variables. To facilitate identification of associations between 200 abundance and environmental predictors weakly identified when considering each 201 sampling location separately, data were pooled across sampling locations for each 202 combination of gene target and sample type. Raw plate count and real-time PCR-MPN 203 observations for multiple aliquots and dilutions of the same sample were treated as 204 repeated and discrete-valued measurements of the same underlying abundance in the 205 given sample. Raw observations comprise the response variables of GLMM regression 206 with plate counts and PCR-MPN outcomes at each dilution level treated as independent 207 Poisson and binomial outcomes, respectively, conditional on latent distribution of 208 abundance and volume of sample examined in each aliquot or dilution. Given apparent 209 inhibition of the PCR reaction at low dilutions in some samples, the PCR-MPN data

| | 210 | were truncated to one |
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| | 211 | temperature and salini |
| | 212 | parts per thousand, re |
| rìni | 213 | 10 logarithms of their r |
| o J | 214 | estimates were determ |
| io To | 215 | posterior Bayesian dis |
| edd | 216 | prior distributions. Ass |
| ahe | 217 | parameters were sum |
| Je | 218 | measure of the propor |
| nlìr | 219 | variation in the enviror |
| 0 | 220 | WinBUGS (40) and the |
| hec | 221 | associations was asse |
| lis | 222 | regression parameters |
| ouk | 223 | For graphical p |
| ts | 224 | vibrios was determined |
| 0 | 225 | one or more plates by |
| CC | 226 | sediment examined. |
| \triangleleft | 227 | Therefore, the limit of |
| \mathbf{N} | 228 | sediment were 1-250 |
| A | 229 | respectively, because |
| | 230 | were tested per V. vuli |
| | | |

dilution, as described elsewhere (31). In regression analyses, ity parameters were expressed in units of degrees Celsius and spectively, while chl-a, DOC and SPM where expressed in base respective measurement units. GLMM regression parameter nined by Markov Chain Monte Carlo (MCMC) sampling of stributions, conditional on the observed data and noninformative sociations between vibrio abundance and environmental marized using McKelvey and Zavoina's pseudo-R² (28, 67), as a tion of variation in latent distributions of abundance attributable to nmental parameter. Statistical analyses were conducted using e R2WinBUGS package of R (57, 71). Statistical significance of essed by identifying Bayesian 95% credible intervals for s that were exclusive of zero (51). resentation of data, the number of vvhA+, tlh+, tdh+, and trh+ d by dividing the total number of colony-forming units (CFU) on the corresponding total volume of water or weight of oyster and Only CFU counts between 1 and 250 CFU per plate were plotted. detection (LOD) ranges for V. vulnificus in water, oysters, and CFU/mL, 10-25,000 CFU/g, and 100-83,333 CFU/g,

1 mL water, 0.1 - 0.01 g oyster, and 0.01 to 0.003 g sediment nificus plate and only 1-250 CFU were counted per plate. The 231 LODs for V. parahaemolyticus in water, oysters, and sediment were 1-250 CFU/mL,

10–25,000 CFU/g, and 20–83,333 CFU/g, respectively, because 1 mL water, 0.1 - 0.01
g oyster, and 0.05 to 0.003 g sediment were tested per *V. parahaemolyticus* plate.

235 **RESULTS**

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234

237 Environmental parameters. Sea surface temperature ranges across the four 238 sample sites were relatively similar (Fig. 1). The lowest temperatures were measured in 239 WA, with LA and MS highest, and MD samples showing the widest temperature range. 240 Based on their similar geography and climate, the two Gulf Coast sites, LA and MS, 241 were combined (GC) for analytical and reporting purposes. Salinities were highest for 242 WA samples and lowest for MD samples; GC samples had the widest salinity range. 243 WA samples contained the lowest median chl-a concentration, median DOC, and SPM, 244 and GC samples had the highest medians for the three parameters. 245 Model-based estimates of abundance. A large number of negative results 246 were obtained using the DP/CH method for guantitation of tdh and trh in water samples 247 and in oysters. The DP/CH approach, therefore, was concluded not to be as

informative as the MPN approach for *tdh* and *trh* enumeration, and these data were

249 excluded for this reason. Results for the WA samples showed *vvhA* to be very low for

250 all three sample types, an indication that V. vulnificus, if present, was below the limit of

251 detection levels in agreement with previous studies (34). At all sampling sites, the

252 largest numbers of vibrios (*tlh*, *tdh*, *trh*, and *vvhA*) were determined when the water

temperature was high (Fig. 2). All GC samples had high *tlh* and *vvhA* numbers and

temperatures. The GC samples also had the highest *tdh* and *trh* densities in oysters,

but the WA samples had the highest *tdh* and *trh* densities in sediment. Overall, trends in vibrio population numbers in water were similar for the all sample types, but WA samples exhibited low *V. vulnificus* densities. MD samples exhibited the lowest mean vibrio densities, with respect to the *tdh* and *trh* gene targets, with relatively large standard deviations.

260 Intra-sample comparisons of V. parahaemolyticus and V. vulnificus. 261 Densities of V. parahaemolyticus (tlh) were compared to V. vulnificus (vvhA) on a 262 sample-by-sample basis, i.e., each *tlh* abundance was compared to *vvh* abundance in 263 the same sample. Comparisons summarizing relative abundance in CFU/mL water or 264 CFU/g of oyster or sediment were used to infer prevalence of one species over the 265 other across sample type and temperature range (Fig. 3). Data are presented in this 266 fashion due to the relatively high rate of non-detection, making calculation of 267 percentages problematic on a sample-by-sample basis. For LA samples, when *tlh* and 268 vvhA were detectable by DP/CH, th outnumbered vvhA in most samples (Fig. 3). 269 Specifically, *tlh* outnumbered *vvhA* about 2/3 of the time in water and sediment and 270 about 4/5 of the time in oysters. The reverse was observed for MS and MD samples, 271 where vvhA typically outnumbered tlh. Thus, overall, in LA samples, V. 272 parahaemolyticus was dominant more frequently in all sample types than V. vulnificus, 273 whereas MS and MD samples were more often dominated by V. vulnificus than by V. 274 parahaemolyticus. 275 Environmental predictors of abundance. Multi-level (GLMM) regression 276 models and associated measures of relative importance of predictor variables (pseudo-

277 R² values) were applied to data pooled across sampling locations to assess the

278 proportion of variation in vibrio abundance attributable to variation in each 279 environmental parameter. Where identified as statistically significant, DOC accounted 280 for 13% of *tlh* variability in oysters (Table 2), 15% of *tdh* variability in water, and 12% of 281 trh variability in sediment; its impact on the other factors were insignificant. Where 282 identified as statistically significant, chl-a accounted for 5% of th variability in sediment, 283 22% of tdh variability in sediment, 13% of trh variability in oysters, and 9.8% of trh 284 variability in sediment. Similarly, where identified as statistically significant, SPM 285 accounted for 6 - 29% of variability in vibrio abundance, depending on sample type, with 286 tdh in oysters being the highest. Salinity accounted for 9% of tlh variability in oysters and 3.7% of *tlh* variability in sediment. Although the pseudo-R² value for salinity in 287 288 Table 2 was high for *tdh* in the water column (31%), this value was not statistically 289 significant and most probably due to chance. SST accounted for 7.1 – 34% of V. 290 vulnificus and V. parahaemolyticus and was a strong predictor in all samples except for 291 tdh in water and oysters and trh in oysters. In most sample types for which SST was a 292 significant predictor, SST explained a larger percentage of variability than any of the 293 other parameters measured. In instances where SST was not significant, SPM was the 294 strongest predictor (Table 2). DP/CH detection rates were highest in sediment samples, 295 followed by oysters and water, and the highest *tdh* or *trh* non-detect rates were in water 296 and in oysters (Table 3).

Estimates determined by analysis of data pooled across sampling locations were further evaluated by comparing to results of analyses of un-pooled data, considering each site separately. Analysis of un-pooled data was conducted to assess consistency of identified associations in the pooled analyses across each sampling location. Similar

patterns of association were observed in analysis of un-pooled data as in the analysis of pooled data, but uncertainty of the identified relationships was much greater. For parameters SST and SPM, which exhibited a relatively strong association with vibrio abundance, the estimated effect size (magnitude and sign of regression coefficients) across sampling locations was more consistent with that of the pooled analysis than was the case with other parameters.

307

308 DISCUSSION

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310 A major goal of this group is to develop ecological models that can be used in 311 conjunction with remotely sensed data collected from and applicable to different 312 geographic regions of the world (i.e., algorithms such as 313 http://www.eol.ucar.edu/projects/ohhi/vibrio/). Development of ecological models for 314 bacteria is strengthened by collection and analysis of samples from diverse geographic 315 locations. Inclusion of geographically distinct study sites to maximize understanding of 316 the role of environmental parameters is a unique contribution of this study. An 317 additional strength is the length and intensity of sampling, which included 594 water, 318 oyster, and sediment samples collected weekly to biweekly over 21 months and a range 319 of environmental parameters measured; this was one of the longest and most intensive 320 sampling programs associated with vibrio abundance and distribution (15, 18, 22, 43, 321 46, 52, 54, 73, 78, 83). Furthermore, our sampling was carried out year-round to 322 examine seasonal variations in vibrio densities. Environmental factors associated with 323 incidence and geographic distribution of V. parahaemolyticus, potentially pathogenic V.

325 areas were analyzed. 326 Ranges in vibrio densities were wider and detection rates were higher in this 327 study than in our previous study, as were ranges of environmental parameters (31). 328 The current study identified highest SPM levels on the Gulf Coast, a result that was not 329 surprising since the Mississippi River plume contributes to turbidity and eutrophy 330 (http://earthobservatory.nasa.gov/IOTD/view.php?id=4982, accessed July 4, 2012), and 331 southerly wind events frequently resuspend sediment in the shallow waters of the 332 northern Gulf of Mexico (75). There appeared to be a degree of niche-specific 333 sequestering, as evidenced by the fact that vvhA exhibited the highest detection rates in 334 oysters while *tlh* exhibited the highest detection rates in sediment. Thus, V. 335 parahaemolyticus and V. vulnificus differed in their niches. In addition, the intra-sample 336 dominance of *tlh* in LA as compared to that of *vvhA* in MS and MD samples indicated 337 some state-to-state variability that may merit consideration as model-based risk 338 assessments are further developed; i.e., vvhA did not consistently outnumber th on a 339 within-sample basis, and in the current study this ratio varied by geographic location. 340 By extending previous work on both vibrio ecology and ecological models for 341 prediction of Vibrio spp. abundance in the aquatic environment, it was reaffirmed that 342 temperature is a strong predictor of abundance and distribution of total vibrios (3, 4, 30, 343 31, 33, 42, 53, 54, 63, 74, 77), and this is particularly useful in the warmer Gulf Coast 344 states included in this study (LA and MS). Though it is clear that temperature is 345 dominant, there is no specific hierarchy among the parameters; environmental factors

parahaemolyticus, and V. vulnificus at four sampling locations in three U.S. coastal

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interact to influence vibrio abundance, but precise details of all such variables and howthey interact have yet to be fully described.

Despite its dominance with respect to *tlh* and *vvhA*, SST was not a strong predictor for densities of vibrios with the pathogenicity genes *tdh* and *trh* in this study. This finding suggests that environmental factors may differentially affect the abundance of pathogenic subpopulations. This is particularly relevant given previous observations that the percentage of total *V. parahaemolyticus* containing these *tdh* and/or *trh* genes appears to be variable and inversely related to temperature (18, 19, 83).

354 Observed associations between abundance and salinity were minimal despite 355 the relatively wide salinity range of this study. Salinity correlated significantly only with 356 tlh in oysters and sediment but with no other measurements in the analysis of data 357 pooled over sampling locations. This was unexpected, given previous observations of 358 significant correlations between salinity and vibrios in samples from Mississippi and 359 Alabama (31). This finding did not appear to an artifact of the pooled data analysis, as 360 analyses of un-pooled data, by sampling location separately, were generally consistent. 361 Specifically, although effects of salinity did not follow the same nonlinear (quadratic) 362 relationship at each sampling location (e.g., due to narrow range of salinity above or 363 below an optimum), the apparent effects at each sampling location were consistent with 364 that of the pooled data, even when the effect overall was identified as not statistically 365 significant. Regression models incorporating an interaction between temperature and 366 salinity were explored but did not significantly improve overall goodness-of-fit or 367 otherwise provide an interpretation for the unexpected findings.

| 368 | Some studies have identified a significant relationship between vibrios and |
|-----|--|
| 369 | salinity (6, 12, 30, 31, 60, 69, 79), while others did not (47, 60, 66, 68), so the |
| 370 | relationship with salinity may be variable and complex. For example, Griffitt and Grimes |
| 371 | (manuscript in preparation) report that large salinity shifts, as seen during the opening of |
| 372 | the Louisiana Bonnet Carré Spillway following the Mississippi River floods of 2011, can |
| 373 | cause detectible and significant change in the relative numbers of pathogenic vibrios. In |
| 374 | vitro growth rates of V. vulnificus biotypes 1, 2, and 3 (9) have been related to distance |
| 375 | from shore (47) with respect to salinity and analyzed to determine the salinity of the |
| 376 | coastal ocean and estuaries with respect to vibrio abundance and distribution. |
| 377 | Observed differences between studies may also be attributable to different salinity |
| 378 | ranges or other factors such as the nutrients sparing the salinity requirement for growth |
| 379 | (66). |
| 380 | The statistically significant contributions of chlorophyll and DOC to the vibrios in |
| 381 | this study were minimal, but findings for SPM were suggestive of a stronger effect. This |
| 382 | is consistent with the significant relationships previously identified between turbidity and |
| 383 | V. parahaemolyticus abundance (30, 31, 33, 47, 54). A positive association with |
| 384 | turbidity is consistent with expectations because vibrios, like many bacteria, are |
| 385 | frequently attached when in the aquatic environment (16). A higher density of |
| 386 | particulate matter suspended in the water column logically provides habitat for a greater |
| 387 | density of vibrios. The current study represents initial efforts to quantify that |
| 388 | relationship. |
| 389 | It was surprising that SST was the only factor that was a statistically significant |
| 390 | predictor of vvhA density in any sample type, even when accounting for the paucity of |

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391 vvhA in Washington. We and others have previously demonstrated relationships 392 between vvhA and environmental parameters including temperature, salinity, and 393 chlorophyll (31). Also interesting, WA samples exhibited the lowest median chl-a, DOC, 394 and SPM levels, as well as the highest salinities.

395 The proportions of *tdh* and *trh* in the Pacific Northwest as measured by DP/CH 396 were high compared to total V. parahaemolyticus (i.e., tlh+), suggesting that tdh+ and/or 397 trh+ V. parahaemolyticus are present in very large numbers. The relatively high 398 salinities in the Pacific Northwest were concluded to be unrelated to the high tdh/trh 399 rates because similar salinities observed at the other sampling sites in this study were 400 not associated with high rates of tdh/trh detection (data not shown). To investigate the 401 possibility of cross-reactivity with other vibrios in the Pacific Northwest, a small subset of 402 vibrios in the GC collection was queried. Of the 23 vibrios containing trh, only two were 403 identified as V. alginolyticus, with the remaining 21 identified as V. parahaemolyticus; 404 tdh was only found in V. parahaemolyticus (Rachel Clostio, personal communication). 405 Other studies of WA tdh+ and trh+ strains, including strain genotyping 406 (Paranjpye, et al., manuscript in preparation) and both multilocus sequence typing and 407 complete genomic sequencing (Turner, et al., manuscripts in preparation), demonstrate 408 that strains in the Pacific Northwest carrying *tlh*, *tdh*, and *trh* are indeed V. 409 parahaemolyticus. Thus, the explanation for the high tdh/trh rates in the Pacific 410 Northwest as measured by DP/CH remains unknown. 411 The unforeseen need for the asymmetrical treatment of DP/CH results from 412 Washington due to unexpectedly high rates of tdh and trh DP/CH results was deemed 413

acceptable and as contributing minimal artifacts because both DP/CH and PCR/MPN

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| 414 | methods target the same genes (49). The PCR/MPN method does include additional |
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| 415 | regions of specificity by its nature because it includes two oligonucleotide primers and a |
| 416 | fluorescent probe while DP/CH only includes an alkaline phosphatase-conjugated probe |
| 417 | that binds to the region targeted by the forward PCR primer. However, potential |
| 418 | variability and artifacts were minimized by treating all three sites in the same manner |
| 419 | where possible, i.e., only including PCR/MPN results for <i>tdh</i> and <i>trh</i> gene targets. |
| 420 | Future studies will address the impact of individual parameters on vibrio |
| 421 | abundance, for which microcosm studies have been initiated at the University of |
| 422 | Maryland that address molecular genetic determination of the vibrios indigenous to the |
| 423 | respective geographic regions of this study, and these results will be presented |
| 424 | elsewhere. Additional data will also be analyzed as a result of a recently concluded |
| 425 | concurrent sampling regime in the four sampling states. A focus of analysis of these |
| 426 | data will be exploring possible differences in relationships between vibrio abundance |
| 427 | and predictor variables across sampling locations, and this will provide further insight |
| 428 | about the initial assessment based on pooling of data. Findings from microcosm study |
| 429 | will be evaluated to better inform model selection in the analysis of field study |
| 430 | observations. Zooplankton and phytoplankton densities and relationships with |
| 431 | additional pigments indicative of phytoplankton will also be analyzed. A sufficiently |
| 432 | large complement of data will facilitate identification of statistical models that are both |
| 433 | interpretable and provide for the best possible predictive value. |
| 434 | In conclusion, the microbial ecology of selected Vibrio spp. has been extensively |
| 435 | studied to determine the importance of specific environmental parameters influencing |
| 436 | the incidence, distribution, and abundance of total and pathogenic vibrios. This study |

438 geographic regions, vibrio densities, seasons, and environmental parameters not 439 studied previously. Maximizing the size of the study made it possible to study 440 parameter ranges that cannot be investigated by studying only a single study site. This 441 study confirmed some previously reported findings (e.g., the impact of temperature) but 442 also identified some new findings (e.g., the differences in the strength of correlation of 443 V. parahaemolyticus and V. vulnificus densities to environmental parameters). 444 Diversifying the geographic niches included in this study improves the chances of 445 identifying environmental signatures that can be used to predict and possibly prevent 446 vibrio outbreaks in a wide and possibly global range of geographic locations. 447 448 Acknowledgments 449 450 This work was supported by NSF RAPID grant #1043126 and NSF grant EF-451 0813285/EF-0813066 as part of the joint NSF-NIH Ecology of Infectious Diseases 452 program and by NASA grant NNX09AR57G. Additional support was provided by the 453 National Oceanic and Atmospheric Administration's Oceans and Human Health Initiative 454 (NA04-OAR-4600214) and NOAA's National Marine Fisheries Service. The authors are 455 indebted to Dr. Robert Gambrell, Dr. Bob Bidigaire, Stephanie Christensen, Drusilla 456 Cowan, Erica Stephens, Pete Cable, Atticus Finger, Conor Smith, William Nilsson, 457 Gladys Yanagida, Asta Stojanowski, Andy DePaola, Adrienne Flowers, Halley Murray, 458 The Olympia Oyster Company in Shelton, WA, and the Port Gamble S'Klallam Tribe in 459

Kingston, WA for sample analyses, technical support, intellectual assistance, and

builds upon existing data sets and findings by including an exceptionally wide range of

437

| 460 | logis | tical assistance, to the anonymous reviewers for a thorough review of the |
|------------|-------|--|
| 461 | man | uscript, and to Dr. Estelle Russek-Cohen for reviewing the manuscript and |
| 462 | provi | ding statistical advice. |
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| 745 | | |

- 746
- 747 Tables.

| target. | | | | | | |
|--|--|--|--|--|--|--|
| Gene | Sample | PCR/MPN method | DP/CH method | | | |
| | water | WA, GC, MD ¹ | GC, MD | | | |
| tlh | oyster | WA, GC, MD | GC, MD | | | |
| | sediment | | WA, GC, MD | | | |
| | water | WA, GC, MD | | | | |
| tdh | oyster | WA, GC, MD | | | | |
| | sediment | | WA, GC, MD | | | |
| | water | WA, GC, MD | | | | |
| trh | oyster | WA, GC, MD | | | | |
| | sediment | | WA, GC, MD | | | |
| | water | | GC, MD | | | |
| vvhA ² | oyster | | GC, MD | | | |
| | sediment | | GC, MD | | | |
| ¹ WA = W probable for all W | Vashington; e number; D A sample s | GC = Gulf Coast; MD = Ma P/CH = direct plating/colony ources for WA were omitted | ryland; MPN = most y hybridization; ² <i>vvhA</i> data I. | | | |

Table 1. Pooled data as pooled for WA, GC, and MD according to gene target.

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| Acce |
| M A |
| AE/ |

| | | tlh | | | vvhA | | | tdh | | | trh | |
|------------|--------|---------|--------|--------|----------|--------|-------|--------|-------|----------------------|----------|------|
| | wat | oys | sed | wat | oys | sed | wat | oys | sed | wat | oys | sed |
| log | | - | | | | | | | | | | |
| DOC | 1.88 | 13.1 | 0.96 | 1.56 | 2.05 | 4.36 | 15.3 | 4.17 | 5.05 | 7.77 | 5.89 | 12.4 |
| log | | | | | | | | | | | | |
| Chla | 1.98 | 0.63 | 5.01 | 1.29 | 2.55 | 1.38 | 12.1 | 6.31 | 22.0 | 3.11 | 12.9 | 9.80 |
| log | | | | | | | | | | | | |
| SPM | 15.6 | 5.99 | 7.40 | 2.38 | 2.80 | 3.80 | 22.9 | 28.9 | 1.91 | 8.64 | 17.4 | 10.1 |
| SalBoth | 2.45 | 8.98 | 3.70 | 3.56 | 3.72 | 2.60 | 30.7 | 5.24 | 4.17 | 5.78 | 4.72 | 4.79 |
| SST | 11.0 | 18.0 | 34.3 | 11.5 | 27.6 | 9.00 | 4.65 | 3.65 | 11.2 | 14.2 | 7.72 | 7.05 |
| Relative i | mporta | ince of | enviro | nmenta | al parar | neters | based | on pse | udo-R | ² statist | tics der | ived |

from GLMM analysis. SalBoth = both salinity and quadratic salinity combined; bold = P < 0.05.

| Table 3. Summary statistics for V. parahaemolyticus and V. vulnificus | | | | | | | | | |
|---|--------------------------|----------------------------|-------------------|--|--|--|--|--|--|
| densities | | | | | | | | | |
| | Range in CFU/mL or CFU/g | | | | | | | | |
| | Probe | for respective DP/CH probe | DP/CH Detects (%) | | | | | | |
| | | (Median) | | | | | | | |
| | vvh | <1 - >250 (6.0) | 79.2 | | | | | | |
| Iter | tlh | <1 - 204 (1.5) | 69.5 | | | | | | |
| ≥ S | tdh | <1 - 66 (<1) | 18.1 | | | | | | |
| _ | trh | <1 - 39 (<1) | 19.7 | | | | | | |
| s | vvh | <10 - >2.5E4 (673.9) | 86.3 | | | | | | |
| ter | tlh | <10 - 2.2E4 (186) | 81.5 | | | | | | |
| Oys | tdh | <10 - 241 (<10) | 24.8 | | | | | | |
| | trh | <10 - 982 (<10) | 34.9 | | | | | | |
| nt | vvh | <100 - >8.3E4 (525) | 61 | | | | | | |
| me | tlh | <20 - >8.3E4 (715) | 89.7 | | | | | | |
| edi | tdh | <20 - 2.4E3 (25) | 61.3 | | | | | | |
| Š | trh | <20 - 3.5E3 (50) | 64.2 | | | | | | |
| <i>tlh,</i> thermolabile hemolysin; <i>tdh,</i> thermostable direct hemolysin; <i>trh, tdh-</i> | | | | | | | | | |
| related hemolysin; vvh, V. vulnificus hemolysin; DP/CH, direct | | | | | | | | | |
| plating/colony hybridization. CFU/mL and CFU/g data are from DP/CH. | | | | | | | | | |

Table 2. Relative importance of environmental parameters for all sampling locations combined.



Figure 1.

| 1 | Figure 1. Box plots of sea surface temperature (SST), salinity (Sal), chlorophyll-a |
|---|--|
| 2 | (Chl-a), dissolved organic carbon (DOC), and SPM (suspended particulate |
| 3 | matter) in Washington (WA), Mississippi + Louisiana (Gulf Coast, GC), and |
| 4 | Maryland (MD). Box plots summarize distribution by indication of the maximum, |
| 5 | 75th percentile, median, 25th percentile, and minimum values. Additional circles |
| 6 | indicate outlier values identified by the statistical package R. Points more than |
| 7 | 1.5 times the interquartile range above the third quartile or below the first quartile |
| 8 | were plotted individually as outliers. Median values are indicated below the |
| 9 | graphs. |
| | |



Figure 2.

| 1 | Figure 2. Model-based estimates of mean and standard deviation (SD) of log |
|---|--|
| 2 | abundance by location and gene target. Estimates of mean and SD of log |
| 3 | CFUs/mL (water) and log CFUs/g (oysters and sediment) are based on |
| 4 | measurements from DP/CH (all vvhA and sediment data points), from real-time |
| 5 | PCR/MPN (tdh and trh in all water/oyster, and tlh in WA water/oyster), or from |
| 6 | both (tlh in GC water/oyster and MD water/oyster). Means are presented with |
| 7 | standard deviation of the distributions and not standard error of means. |
| 8 | |



Figure 3A. The *tlh:vvhA* densities for water samples.









- 1 Figure 3. The *tlh:vvhA* relative densities by habitat and location. The DP/CH-
- 2 derived *tlh* densities were compared to DP/CH-derived *vvhA* densities on a
- 3 sample-to-sample basis for water (A), oysters (B), and sediment (C). Black bars
- 4 = *tlh* densities; gray bars = *vvhA* densities; diamond lines = sea surface
- 5 temperature in °C plotted on secondary (right) y-axis. WA data were excluded
- 6 from these graphs because of the lack of *vvhA* counts.
- 7