1	Parallel evolution of two clades of a major Atlantic endemic Vibrio parahaemolyticus pathogen
2	lineage by independent acquisition of related pathogenicity islands
3	
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7	Running Title: parallel evolution of ST631 Vibrio parahaemolyticus
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21 ABSTRACT

22 Shellfish-transmitted Vibrio parahaemolyticus infections have recently increased from 23 locations with historically low disease incidence, such as the Northeast United States (US). This 24 change coincided with a bacterial population shift towards human pathogenic variants occurring 25 in part through the introduction of several Pacific native strains (ST36, ST43 and ST636) to near-26 shore areas off the Atlantic coast of the Northeast US. Concomitantly, ST631 emerged as a 27 major endemic pathogen. Phylogenetic trees of clinical and environmental isolates indicated that 28 two clades diverged from a common ST631 ancestor, and in each of these clades, human 29 pathogenic variants evolved independently through acquisition of distinct *Vibrio* pathogenicity 30 islands (VPaI). These VPaI differ from each other and that bear little resemblance to hemolysin-31 containing VPaI from pandemic strains. Clade I ST631 isolates either harbored no hemolysins, 32 or contained a chromosome I-inserted island we call VPaI β that encodes a type three secretion 33 system (T3SS2B) typical of Trh hemolysin-producers. The more clinically prevalent and clonal 34 ST631 clade II had an island we call VPaly that encodes both *tdh* and *trh* and that was inserted in 35 chromosome II. VPaIy was derived from VPaIß but with some elements in common with 36 pandemic strains, exemplifying the mosaic nature of pathogenicity islands. Genomics 37 comparisons and amplicon assays identified VPaIy-type islands containing *tdh* inserted adjacent 38 to the *ure* cluster in the three introduced Pacific and most other lineages that collectively cause 39 67% of Northeast US infections as of 2016.

40

41 **IMPORTANCE**

42 The availability of three different hemolysin genotypes in the ST631 lineage provided a
43 unique opportunity to employ genome comparisons to further our understanding of the processes

44 underlying pathogen evolution. The fact that two different pathogenic clades arose in parallel 45 from the same potentially benign lineage by independent VPaI acquisition is surprising 46 considering the historically low prevalence of community members harboring VPaI in waters 47 along the Northeast US Coast that could serve as the source of this material. This illustrates a 48 possible predisposition of some lineages to not only acquire foreign DNA but also to become 49 human pathogens. Whereas the underlying cause for the expansion of V. parahaemolyticus 50 strains harboring VPaly along the US Atlantic coast and spread of this element to multiple 51 lineages that underlies disease emergence is not known, this work underscores the need to define 52 the environment factors that favor strains harboring VPaI in locations of emergent disease.

53

54 **INTRODUCTION**

55 *Vibrio parahaemolyticus* is an emergent pathogen capable of causing human gastric infections when consumed, most often with contaminated shellfish (1, 2). Some human 56 57 pathogenic V. parahaemolyticus strains evolve from diverse non-pathogenic communities 58 through horizontal acquisition of Vibrio pathogenicity islands (VPaI) (3-5). Gastric pathogenic V. 59 *parahaemolyticus* typically harbor islands with at least one of two types of horizontally acquired 60 hemolysin genes (*tdh* and *trh*) that are routinely used for pathogen discrimination even though 61 their role in disease appears modest (6-11). Most pathogenic V. parahaemolyticus isolates also produce accessory type three secretion systems (T3SS) that translocate effector proteins that 62 63 contribute to host interaction (12-14). Two evolutionarily divergent horizontally-acquired 64 accessory systems (T3SS2 α or T3SS2 β) contribute to human disease and are genetically linked 65 to hemolysin genes (two *tdh* genes with T3SS2 α , and *trh* with T3SS2 β) in contiguous but distinct 66 islands (4, 15-17). The first described *tdh*-harboring island [called by several different names

67	including Vp-PAI (15), VPaI-7 (4), and <i>tdh</i> VPA (17)] from an Asian pandemic strain called							
68	RIMD 2210366 is fairly well-characterized (4, 5, 13, 18, 19). In contrast, islands containing							
69	T3SS2 β linked to <i>trh</i> and a urease (<i>ure</i>) cluster, which confers a useful diagnostic phenotype,							
70	[where similar islands are described by others as Vp-PAI _{TH3966} (16), or <i>trh</i> VPA(17, 20)] have							
71	received only modest attention. Strains harboring both tdh and trh are increasingly associated							
72	with disease in North America (21-26), and yet, to our knowledge, the exact configuration of							
73	hemolysin-associated VPaI(s) in isolates that contain both <i>tdh</i> and <i>trh</i> have not yet been							
74	described [although see (20)]. Thus it is unclear how virulence loci and islands in these emergent							
75	pathogen lineages carrying both hemolysins evolved and spread.							
76	The expanding populations of V. parahaemolyticus have increased infections even in							
77	temperate regions previously only rarely impacted by this pathogen and where most							
78	environmental isolates harbor no known virulence determinants (27). A related complex of Asia-							
79	derived pandemic strains, most often identified as serotype O3:K6 and also known as sequence							
80	type (ST) 3 (based on allele combinations of seven housekeeping genes) causes the most disease							
81	globally (28). An unrelated Pacific native strain called ST36 (also described as serotype O4:K12)							
82	currently dominates infections in North America, including from the Northeast United States							
83	(US) (21, 26, 29). The introduction of ST36 into the Atlantic Ocean by an unknown route							
84	precipitated a series of outbreaks from Atlantic shellfish starting in 2012 (29, 30). Prior to 2012,							
85	local strains contributed to low but increasing sporadic infection rates on the Northeast US coast							
86	(https://www.cdc.gov/vibrio/surveillance.html, 2017) (21), with ST631 emerging as the major							
87	lineage that is endemic to near-shore areas of the Atlantic Ocean bordering North America (the							
88	northwest Atlantic Ocean) (31). However, we previously identified a single ST631 isolate							

lacking hemolysins (21, 27) suggesting this pathogen lineage may have recently evolved through
VPaI acquisition.

91 The goal of our study was to understand the genetic events and changing population 92 context for the evolution of the ST631 pathogenic lineage. We conducted whole and core 93 genome phylogenetic analysis of three environmental and 39 clinical ST631 isolates along with 94 isolates from other emergent lineages from the region, which revealed two ST631 clades of 95 common ancestry, from which human pathogens have evolved in parallel. The single clade I 96 clinical isolate acquired a *recA* gene insertion previously seen associated with Asian strains, and 97 had VPaI that is typical of isolates harboring *trh* in the absence of *tdh*. In contrast, the clonal 98 ST631 clade II that dominates Atlantic-derived ST631 infections (31) had a related but distinct 99 VPaI. This VPaI contained a *tdh* gene inserted within, not next to, an existing *ure-trh*-T3SS2β 100 island in close proximity to the *ure* cluster. Nearly all emergent resident lineages and invasive 101 strains, including all three Pacific lineages (ST36, ST636 and ST43) contained islands that 102 similarly had a *tdh* gene inserted within the VPaI in an identical location adjacent to the *ure* 103 cluster providing a mechanism for simultaneous acquisition of both hemolysins with T3SS2β. 104

105 **RESULTS**

106 Atlantic endemic ST631 and several invasive strains harboring both the *tdh* and *trh*

107 hemolysin genes are clinically prevalent in four reporting Northeast US States.

108 Ongoing analysis of clinical isolates revealed that even as the Pacific-derived ST36 109 lineage continued to dominate infections (50%), the endemic (autochthonous) ST631 lineage 110 accounted for 14% of infections (Table 1). Concurrently, a limited number of other lineages 111 contributed individually to fewer infections (\leq 3% each), among which were two lineages that

112 have caused infections in the Pacific Northwest in prior decades: ST43 and ST636 (22, 23). 113 ST43 and ST636 only recently (2013 and 2011 respectively) (21) have been linked to product 114 harvested from waters along the Northeast US coast, and also caused infections in subsequent 115 years. As is common among US clinical isolates, pathogenic strains of all the aforementioned 116 lineages harbor both the *tdh* and *trh* hemolysin genes (Table 1). Among environmental isolates, 117 ST34 and ST674 are the most frequently recovered pathogen lineages but these caused 118 comparatively few infections (Table 1). ST34 was first reported from the environment in 1998, 119 from both the Gulf of Mexico and near-shore areas of MA, and was also recovered in NH in 120 2012 (21) suggesting it is an established resident in the region. ST674 which was first reported 121 from an infection in Virginia in 2007 (32) was first recovered from the local environment in 122 2012 (www.pubmlst.org/vparahaemolyticus) (21). Notably even though all four ST674 123 environmental isolates, like ST34, harbored both hemolysin genes, the single ST674 clinical 124 isolate (MAVP-21) lacked hemolysins (Table 1) (21). The decrease in clinical prevalence of trh-125 harboring Atlantic endemic ST1127, which caused no infections in the last three years, coincided 126 with the increase in clinical prevalence of all three Pacific-derived lineages which harbor both 127 hemolysins. Notably, very few other clinical isolates harbored *trh* in the absence of *tdh* and 128 clinical isolates containing only *tdh* (i.e. ST1725) were extremely rare (Table 1). Concurrent with 129 this shift in composition of clinical lineages that includes multiple Pacific-derived strains, 130 hemolysin producers have increased in relative abundance in nearshore areas of the region, 131 where historically these represented $\sim 1\%$ of all isolates (27). Since 2012, hemolysin producers 132 have been recovered more frequently, and in the last two years their proportion has increased by 133 up to an order of magnitude (comprising as much as 10%) in some regional shellfish associated 134 populations (data not shown).

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136 A single clinical ST631 lineage isolate with an unusual *recA* allele harbors *trh* in the

- 137 absence of *tdh*
- 138 Employing ST631-specific marker-based assays (see methods), we identified two
- additional 2015 environmental isolates (one from NH and one from MA) and one additional
- 140 2011 local-source clinical isolate (MAVP-R) (21) with a hemolysin profile $(trh^+$ without tdh)
- 141 that is atypical of the ST631 lineage (Table 1). Although analysis of the seven-housekeeping
- 142 gene allele combination confirmed the environmental isolates were indeed ST631, MAVP-R was
- 143 not ST631 based on only one locus: recA. Examination of the recA locus of MAVP-R uncovered
- a large insertion within the ancestral ST631 *recA* gene (allele recA21;
- 145 www.pubmlst.org/vparahemolyticus) incorporating an intact but different *recA* gene into the
- 146 locus [allele recA107(33)] and fragmenting the ancestral gene (Fig. 1). The insertion in the
- 147 ancestral *recA* gene in MAVP-R is identical to one observed in the *recA* locus of two Hong Kong
- isolates (strains S130 and S134) and similar to the one in strain 090-96 (ST189a) isolated in Peru
- 149 but believed to have originated in Asia (33).
- 150
- 151 ST631 forms two divergent clades

The existence of three different hemolysin profiles (Table 1) among all available ST631 draft genomes suggested there could be more than one ST631 lineage. Therefore we evaluated whole genome maximum likelihood (ML) phylogenies of select ST631 isolates and all other lineages causing two or more infections reported in four Northeast US States to evaluate whether there was more than one ST631 lineage (Table 1) (Fig. 2). The phylogenetic tree showed that ST631 isolates, regardless of their hemolysin genotype, clustered together but they formed two

158 distinct clades, indicative of common ancestry (Fig. 2). Clade I harbored either *trh* or no 159 hemolysins and consisted of all three environmental isolates which were from MA and NH, and 160 the single clinical isolate MAVP-R, whereas clade II consisted of all other isolates all of which 161 harbor both hemolysins. The two distinct ST631 clades shared 85% of their DNA in common 162 and displayed polymorphisms in $\leq 12\%$ of the shared DNA content. The most closely related 163 sister lineage to ST631 was formed by *trh*-harboring ST1127 isolates that have been exclusively 164 reported from clinical sources in the Northeast US (21). 165 We next evaluated the relationships of all available ST631 isolate genomes at NCBI and 166 sequenced by us (Supplemental Table 1) using a custom core genome multi-locus sequence 167 typing (cgMLST) method as previously described (31). Minimum spanning trees built from core 168 genome loci from 42 ST631 isolates indicated that only 390 loci varied between the most closely 169 related isolate of clade I (MAVP-L) and clade II (G6928) (Fig. 3). The most distantly related 170 isolates within clade I (G149 and MAVP-R) exhibited 80 core genome loci differences whereas 171 clade II is clonal with only 51 variant loci between the most divergent isolates: clinical isolate 172 09-4436 and environmental isolate S487-4, both reported from PEI Canada (Fig. 3) (31). 173 174 Each ST631 clade independently acquired a distinct pathogenicity island positioned on 175 different chromosomes 176 Given the variation in ST631, comparisons between these isolates could elucidate the

events that led not only to the evolution of two pathogenic clades but also address unresolved questions about the unique configurations and contents of pathogenicity islands in western Atlantic Ocean emergent lineages. The physical proximity of *tdh* with the *ure* cluster and *trh*, and the co-occurrence of *tdh* with T3SS2β reported in many *tdh*⁺/*trh*⁺ clinical isolates suggested *tdh* could be harbored within or next to the same pathogenicity island harboring *trh* in at least
some lineages as was previously suggested (20, 24, 34).

183 To identify the location and determine the architecture of the pathogenicity elements 184 harboring hemolysin genes, we generated high quality annotated genomes for the clade I ST631 185 isolate MAVP-R and clade II ST631 isolate MAVP-Q (both reported in 2011 from MA) 186 employing PacBio sequencing. The pathogenicity island regions in these isolates genomes were 187 extracted, aligned, and the contents compared with pathogenicity island harboring two *tdh* genes 188 [previously called Vp-PAI (15), VPaI-7 (4) and tdhVPA(17)] from RIMD 2210366 and Vp-189 PAI_{TH3996} (16) [also called *trhVPI* (17)] harboring *trh* (Supplemental Table 2). This comparison 190 revealed that MAVP-R harbored a pathogenicity island typical of *trh*-containing strains that 191 includes a linked *ure* cluster and T3SS2 β that is orthologous, with the exception of few unique 192 regions, with Vp-PAI_{TH3996} (16) (Supplemental Table 2 and Fig. 4). Because the lack of 193 convention in uniformly naming syntenous islands that distinguish them from distinctive and yet 194 functionally analogous islands can impede communication, we hereafter will consistently 195 reference the same island by a common descriptive name regardless of strain lineage. Hereafter 196 we will refer to islands sharing the same general configuration to that in MAVP-R by the name 197 VPaI β , and refer to *tdh*-containing islands similar to that described in strain RIMD 2210366 by 198 the name VPaIa, regardless of strain background. We adopted this simplified nomenclature in 199 reference to the version of the key virulence determinant carried in the islands (T3SS2 α and 200 T3SS2 β) in the two already described island types. This scheme importantly accommodates 201 naming of additional uniquely-configured islands as they are identified. As noted previously (16, 202 17, 20), VPaI β is dissimilar to VPaI α in most gene content with ~ 78 ORFs unique to VPaI β 203 (where the number of identified ORFs used for comparison can differ slightly depending on

204 which annotation program is applied) (Supplemental Table 2, Fig. 4). Even so, VPaIB had many 205 homologous genes of varying sequence identity (n=~38 ORFs, excluding *tdh* homology with *trh*) 206 when compared to VPaI α (Supplemental Table 2, Fig. 4)(4, 5, 16). Identification of some 207 homologs required that we relax matching to 50% such as for the divergent, but homologous 208 T3SS2 α and T3SS2 β genes encoding the apparatus, chaperones, and some shared effectors 209 (Supplemental Table 2). VPaI β from strain TH3996 and VPaI α from pandemic strain RIMD 210 2210633 are inserted in an identical location in chromosome II adjacent to an Acyl-CoA 211 hydrolase-encoding gene. In contrast the VPaIßs in MAVP-R, ST1127 isolate MAVP-25, and 212 Asia-derived AQ4037 are in chromosome I, in each case in the same insertion location identified 213 for strain AQ4037 (17). 214 MAVP-Q contained both *tdh* and *trh* within the same contiguous unique VPaI (hereafter 215 called VPaI γ) that shared features with both VPaI α and VPaI β (Fig. 4, Supplemental Table 2). 216 Specifically, VPaIy had a core that with few exceptions was orthologous in content and 217 syntenous with VPaI β from MAVP-R (Fig. 4). VPaI γ displays high conservation with VPaI α 218 near its 3' end, as has been described in other draft tdh^+trh^+ harboring genomes (20) as well as in 219 the VPaI β island of strain TH3996, although the presence of this element may not be typical of 220 VPaIβ (e.g. it is absent in the islands from AQ4037 (17), MAVP-R and MAVP-25). The VPaIγ 221 also contained a *tdh* gene homologous to *tdh2* (also called *tdhA*) from VPaIa (98.6%) near its 5' 222 end but not at the 5' terminus of the island (Fig. 4). Rather, the DNA flanking both sides of the

tdh gene in VPaIγ was conserved in VPaIβ of MAVP-R and absent from VPaIα, (Fig. 4).

Analysis of 300 genomes of V. parahaemolyticus (representing a minimum of 28 distinct

sequence types) of sufficient quality for analysis confirmed that the module of four hypothetical

226 proteins preceding the *tdh2* homolog was present only in *trh*-harboring genomes, but not in

227	genomes harboring tdh in the absence of trh (i.e. VPaI α containing genomes), providing							
228	evidence that the <i>tdh</i> gene was acquired horizontally by insertion into, not next to, an existing							
229	VPaI β , perhaps through activity of the adjacent transposase gene (11) (Supplemental Table 3,							
230	Supplemental fig. 1, and data not shown). Like with VPaI α from RIMD 2210633, and VPaI β of							
231	TH3996, VPaIy of clade II ST631 is located in a conserved location of chromosome II, adjacent							
232	to an Acyl-CoA hydrolase-encoding gene.							
233	The final environmental ST631 clade I isolate that lacked hemolysins, G149, had no							
234	VPaI α , β or γ elements in its genome. Close examination of the DNA corresponding to the VPaI							
235	insertion sites in either chromosome revealed no remnants of these islands in either chromosomal							
236	location indicating this isolate likely never acquired a pathogenicity island (Supplemental Fig. 2							
237	and data not shown). Because clade I isolate G149 lacked these islands, this could be the							
238	ancestral state of the ST631 lineage (21).							
239								
240	Most clinically prevalent strains from the Northeast US harbor similar contiguous							
241	pathogenicity islands containing <i>tdh</i> inserted in the same location of their VPaI							
242	We next asked which other strains likely residing within the mixed population with							
243	ST631 in near-shore areas of the Northeast US harbored islands of similar structure to VPaIy that							
244	contain both hemolysin genes. Assembly of short-read sequences into contigs that cover the full							
245	length of VPaI which is necessary for comparative analysis of entire island configuration was							
246	impeded by the fact that homologous transposase sequences were repeated multiple times							

throughout the island. Therefore, we determine whether other lineages harboring both hemolysin

hypothetical proteins (to the left or 5' of *tdh*) and the *ure* cluster (to the right or 3' of *tdh*) (Fig. 4)

genes harbor *tdh* in the same island location, between the conserved VPaI β/γ module of four

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by combining bioinformatics analysis of sequenced genomes with amplicon assays 250 251 (Supplemental Fig. 1). First we analyzed assembled draft genomes for *tdh* co-occurrence and 252 proximity with the four adjacent hypothetical protein-encoding genes (See Methods). Every 253 emergent pathogenic lineage (Table 1) harboring both *tdh* and *trh* carried homologous DNA 254 corresponding to all four hypothetical proteins adjacent to the *tdh* gene in a contiguous segment 255 (Supplemental Table 3). To determine whether *tdh* was also adjacent to the *ure* cluster in these 256 same strains we next designed specific flanking primers and amplified the unique juncture 257 between the *tdh*-containing transposon associated module and the *ure* cluster for all clinical 258 isolates harboring both *tdh* and *trh* (See Methods) (Supplemental Fig. 1). The results were 259 congruent with our bioinformatics assessment (Supplemental Table 3), and demonstrated that 260 isolates from all emergent pathogenic lineages harboring both hemolysins have *tdh* inserted in 261 close proximity to an *ure* cluster in a configuration similar to VPaly from MAVP-Q (Fig. 5, 262 Table 1). This confirmed that these strains harboring both hemolysins harbor *tdh* within, and not 263 next to, the same VPaI thereby facilitating simultaneous acquisition of both hemolysin genes. 264

265 **DISCUSSION**

Even preceding the increased illnesses from Pacific-invasive lineages, two different clades of the predominant endemic Atlantic lineage of pathogenic *V. parahaemolyticus*, ST631 (31) evolved and contributed to a rise in sporadic illnesses in the four reporting Northeast US States (Table 1, Fig. 2 & 3). Several lines of evidence support the interpretation of parallel pathogen evolution. The two lineages exhibit differences in both clinical and environmental prevalence suggesting the pathogenic variants of each clade have not evolved the same degree of virulence (Table 1). Pathogenic members in each lineage also acquired different pathogenicity 273 islands with different hemolysin gene content (Fig. 2 & 3). Although it was a formal possibility 274 that ST631 clade II evolved from clade I by independent horizontal acquisition of *tdh* into its 275 existing VPaI β , it is notable that other resident and even invasive lineages now in the Atlantic 276 harbor VPaIy with *tdh* inserted into the same location, suggesting a common evolutionary origin 277 of this hybrid type island (Fig. 4 and Supplemental Fig. 1). Finally, each of the two clades harbor 278 VPaI insertions on different chromosomes: the less clinically prevalent ST631 clade I contains 279 three isolates that harbor VPaI β in chromosome I (Fig. 3) and a single environmental isolate 280 lacking any island (Table 1, supplemental Fig. 2), whereas the clonal ST631 clade II isolates all 281 harbor VPaIy on chromosome II. 282 Given that several other resident lineages harbor similar β and γ -type VPaI, pathogens in 283 each clade could have acquired their islands from the reservoir of strains already circulating in 284 the Atlantic even before the presume arrival of invasive Pacific lineages. Several well-285 documented members of the Gulf of Mexico V. parahaemolyticus population (35-37) may also 286 have expanded their range through movement of ocean currents and could be the source for these 287 VPaI (Table 1, Fig. 5). But historically, hemolysin producers were extremely rare in near shore 288 areas of the Atlantic US coast (25) and represented only about $\sim 1\%$ of isolates in an estuary of 289 NH as of a decade ago (27) limiting the potential for interacting partners or sources for acquired 290 VPaI. Given this historical context, it is remarkable that two different clades from the same 291 lineage independently acquired different VPaI-which for clade II ST631 occurred prior to 2007 -292 well before the recent shift in abundance of hemolysin producers. 293 The parallel evolution of two different lineages through lateral DNA acquisition alludes 294 to the possibility that as-yet-undefined attributes may increase the chances of acquisition or

295 prime some bacterial lineages (such as ST631) to more readily acquire and maintain genetic

296 material or become pathogenic upon island acquisition. Even though the ecological niche in 297 which horizontal island acquisition took place is unknown, it is conceivable that co-colonization 298 of hosts or substrates favorable to the growth of ST631 and hemolysin producers may have 299 facilitated island movement. Certainly, association of bacteria with specific marine substrates 300 such as chitinous surfaces of plankton that also induce a natural state of competence could 301 promote lateral transfer through close contact between the progenitors of the pathogenic 302 subpopulation of each clade and island donors (3, 38, 39). Alternatively, conjugative plasmids or 303 transducing phage could have been the agents of island delivery. The finding that the only 304 clinical clade I isolate, MAVP-R, also harbors a second horizontal insertion in its recA locus that 305 matched one previously found in Asia-derived strains (33) indicates it acquired more than one 306 segment of foreign DNA during its evolution as a pathogen (Fig. 1) further illustrating that 307 mechanisms that facilitate DNA transfer and acquisition may both have been at play. It also 308 suggests that horizontal transfer of DNA from introduced strains not yet detected in the Atlantic 309 could add to the genetic material available for pathogen evolution from Atlantic Ocean 310 populations. The more detailed molecular epidemiological, comparative genomic, and functional 311 analyses necessary to assess the impact of introduced pathogens on resident Atlantic lineages are 312 warranted given this evidence and the documented introduction of multiple Pacific-derived 313 strains in the region (Table 1).

There has been some consideration of the roles of human virulence determinants in ecological fitness, but the natural context of pathogenic *V. parahaemolyticus* evolution is still unknown (40-42). Whereas *tdh* and T3SS2 α each may promote growth when bacteria are under predation, isolates that carry *trh*-containing islands (which likely also have T3SS2 β) do not derive similar benefits from their islands (43). This is surprising considering the islands encode 319 several homologous effectors (Fig. 4 and Supplemental Table 2) that are not thought to 320 contribute to human disease but could mediate eukaryotic cell interactions with natural hosts 321 thereby promoting environmental fitness (13, 14). The general lack of knowledge of unique 322 T3SS2 β effectors and other gene function in these islands (Fig. 4 and Supplemental Table 2) 323 even with regard to human disease, limits comparative analysis with the well-studied and 324 functionally defined VPaI α which could elucidate the bases for pathogen evolution. The higher 325 clinical prevalence of clade II ST631 than clade I which has also been recovered on more than 326 one occasion from the environment (Table 1) could indicate that VPaly confers greater virulence 327 potential than VPaI β , perhaps owing to the presence of *tdh*, a known virulence factor (1, 7, 44). 328 However, the resident community members in both the Pacific and the Atlantic Ocean that 329 harbor *tdh* and T3SS2α comparatively rarely cause human infections (21-23). The unique 330 environmental conditions that underlie pathogen success from northern latitudes that favors 331 strains with VPaIß and VPaIy including two different ST631 lineages suggests the shared content 332 of these islands could confer abilities that are distinct from VPaIa which could underlie the 333 repeated acquisition and maintenance of these related islands by so many different lineages now 334 present in near-shore areas of the Northeast US.

335

336 MATERIALS AND METHODS

337 Bacteria isolates, media and growth conditions.

V. parahaemolyticus clinical isolates for this study were provided by cooperating public
health laboratories in Massachusetts, New Hampshire, Maine, and Connecticut whereas a select
number of environmental isolates were enriched from estuarine substrates as described (21).
Detailed information about these isolates was described previously (31) and listed in

342 Supplemental Table 1. Isolates were routinely cultured in Heart Infusion (HI) media

343 supplemented with NaCl at 37°C as described (21).

344

345 Whole genome sequencing, assembly, annotation and sequence type identification.

346 Genomic DNA was extracted using the Wizard Genomic DNA purification Kit (Promega,

347 Madison WI USA) or by organic extraction (21). The qualities of all the genomic DNA was

348 measured by NanoDrop (ThermalFisher, Waltham MA USA). Libraries for DNA sequencing

349 were prepared using a high-throughput Nextera DNA preparation protocol (45) using an optimal

350 DNA concentration of 2ng/µl. Genomic DNA was sequenced using an Illumina – HiSeq2500

device at the Hubbard Center for Genome Studies at the University of New Hampshire, using a

352 150bp paired-end library. De novo assembly was performed using the A5 pipeline (46), and the

assemblies annotated with Prokka1.9 using the "genus" option and selecting "Vibrio" for the

reference database (47). The sequence types were subsequently determined using the SRST2

355 pipeline (48). The sequence type of each genome was determined when using *V*.

356 *parahaemolyticus* as the database (https://pubmlst.org/vparahaemolyticus/). For most isolates

357 where the combination of each allele was not found in the database representing novel sequence

358 types, the genome was submitted for a new sequence type designation

359 (www.pubmlst.org/vparahaemolyticus).

360 Isolates MAVP-Q and MAVP-R were sequenced using the Pacific Biosciences RSII

technology. Using between 3.7-5.3 μg DNA, the library preparation and sequencing was

362 performed according to the manufacturer's instructions (Pacific Biosciences, Menlo Park CA,

363 USA) and reflects the P5-C3 sequencing enzyme and chemistry for MAVP-Q isolate and the P6-

364 C4 configuration for MAVP-R. The mass of double-stranded DNA was determined by Qubit

365	(Waltham, MA USA) and the sample diluted to a final concentration of 33 μg / μL in a volume
366	of 150 μ L elution buffer (Qiagen, Germantown MD USA). The DNA was sheared for 60
367	seconds at 4500 rpm in a G-tube spin column (Covaris, Wobrun MA USA) which was
368	subsequently flipped and re-spun for another 60 seconds at 4500 rpm resulting in a ~20,000 bp
369	DNA verified using a DNA 12000 Bioanalyzer gel chip (Agilent, Santa Clara, CA USA). The
370	sheared DNA isolate was then re-purified using a 0.45X AMPure XP purification step (Beckman
371	Coulter, Indianapolis IN USA). The DNA was repaired by incubation in DNA Damage Repair
372	solution. The library was again purified using 0.45X Ampure XP and SMRTbell adapters ligated
373	to the ends of the DNA at 25°C overnight. The library was treated with an exonuclease cocktail
374	(1.81 U/µL Exo III 18 and 0.18 U/µL Exo VII) at 37°C for 1 hour to remove un-ligated DNA
375	fragments. Two additional 0.45X Ampure XP purifications steps were performed to remove
376	<2000 bp molecular weight DNA and organic contaminant.
377	Upon completion of library construction, samples were validated using an Agilent
378	DNA 12000 gel chip. The isolate library was subjected to additional size selection to the range
379	of 7,000 bp – 50,000 bp to remove any SMRTbells < 5,000 bp using Sage Science Blue Pippin
380	0.75% agarose cassettes to maximize the SMRTbell sub-read length for optimal de
381	novo assembly. Size-selection was confirmed by Bio-Analysis and the mass was quantified using
382	the Qubit assay. Primer was then annealed to the library (80°C for 2 minute 30 followed by
383	decreasing the temperature by 0.1% to 25°C). The polymerase-template complex was then
384	bound to the P5 or P6 enzyme using a ratio of 10:1 polymerase to SMRTbell at 0.5 nM for 4
385	hours at 30°C and then held at 4°C until ready for magbead loading, prior to sequencing. The
386	magnetic bead-loading step was conducted at 4°C for 60-minutes per manufacturer's guidelines.
387	The magbead-loaded, polymerase-bound, SMRTbell libraries were placed onto the RSII machine

388	at a sequencing concentration of 110-150 pM and configured for a 180-minute continuous							
389	sequencing run. Long read assemblies were constructed using HGAP version 2.3.0 for de novo							
390	assembly generation. Further, hybrid assemblies were generated and error corrected with							
391	illumina raw reads using Pilon v1.20 (49).							
392								
393	Lineage-specific marker-based assays							
394	To more rapidly identify ST631 isolates from clinical and environmental collections we							
395	developed PCR-amplicon assays to unique gene content in ST631. Whole genome comparisons							
396	were performed on MAVP-Q (ST631 clinical strain), G149 (ST631 environmental strain),							
397	MAVP-26 (ST36), RIMD2210633 (ST3), and AQ4037 (ST96) (Supplemental Fig. 3). A total of							
398	26 distinct genomic regions, each greater than 1kb in size, were present in MAVP-Q but absent							
399	in other comparator genomes, including environmental ST631 that lacks hemolysins (G149)							
400	(Supplemental Fig. 3). Within a large genomic island ~37.6 Kb in length with an integrase at one							
401	terminus and an overall lower GC content (40.6% compared to 45.8% for the genome) a single							
402	ORF homologous to restriction endonucleases (AB831_06355) that was restricted to clinical							
403	ST631 isolates in our collection and publicly available draft genomes (n=693)							
404	(http://www.ncbi.nlm.nih.gov/genome/691, 2017) was selected as a suitable amplicon target. The							
405	distribution of this locus was further analyzed using the BLAST algorithm by a query against the							
406	nucleotide collection, the non-redundant protein sequences, and against the genus Vibrio (taxid:							
407	662), excluding V. parahaemolyticus (taxid: 691), using the default settings for BLASTn (50).							
408	Similar approaches were applied to identify ST631 diagnostic loci inclusive of the single							
409	environmental isolate (G149), which identified a hypothetical protein encoding region							
410	(AB831_06535) (ST631env). Oligonucleotide primers were designed to amplify the diagnostic							

411 regions including AB831_06355 using primers ST631end F

- 412 (5'AGTTCATCAGGTAGAGAGTTAGAGGA3') and ST631endR
- 413 (5'TCTTCGTTACCATAGTATGAGCCA3') which produces and amplicon of c.a. 494bp, and
- 414 AB831_06535 using primers ST631envF (5'TGGGCGTTAGGCTTTGC3') and ST631-envR
- 415 (5'GGGCTTCTACGACTTTCTGCT3') producing an amplicon of 497bp.

416 Amplification of diagnostic loci was evaluated in individual assays using genomic DNA

- 417 from positive and negative controls: MAVP-Q and G149 (ST631), G4186 (ST34), G3578
- 418 (ST674), and MAVP-M (ST1127), MAVP-26 (ST36) and G61 (ST1125). Amplification of
- 419 specific sequence types were performed with Accustart enzyme mix on purified DNA. Cycling
- 420 was performed with an initial denaturation at 94°C for 3 min., followed by 30 cycles of a
- 421 denaturation at 94°C for 1min, annealing at 55°C for 1 min, and amplification at 72°C for 30s
- 422 with a final elongation at 72°C for 5 min. The primer pairs only produced amplicons from
- 423 template DNA from ST631 and each was the expected size (data not shown, and Supplemental
- 424 Fig. 3). Amplicon assays were applied to 208 clinical isolates from the Northeast US States (ME,
- 425 NH, MA and CT) and 1140 environmental isolates collected from 2015-2016 from NH and MA.
- 426 These assays identified all known ST631 clinical isolates with 100% specificity and also
- 427 identified an additional 7 tdh^+trh^+ clinical isolates (ST631*end* and ST631env positive), and two
- 428 environmental (ST631end negative and ST631env positive) isolates from our archived collection.
- 429 Each, with the exception of MAVP-R, was subsequently confirmed to be ST631 by seven-locus
- 430 MLST (www.pubmlst.org).
- 431

432 Examination of *recA* allele and adjacent sequences

433	The PacBio sequenced genome of MAVP-R, contig 000001 (Accession No.
434	MPPP00000000) that contained the recA gene, was annotated using PROKKA1.9 (47). The
435	sequences of <i>recA</i> and its surrounding DNA was then compared to the contig containing <i>recA</i>
436	region from strain S130 (AWIW01000000), S134 (AWIS01000000), 090-96 (JFFP01000036)
437	(33) and MAVP-Q (Accession No. MDWT00000000). The map of <i>recA</i> region of the five
438	isolates was illustrated using Easyfig (51).
439	
440	Core genome SNP determination and phylogenetic analysis
441	Whole genome phylogenies were constructed with single nucleotide polymorphisms
442	(SNPs) identified from draft genomes using kSNP3 to produce aligned SNPs in FASTA format
443	(52). A maximum likelihood (ML) tree was then built from the FASTA file using raxMLHPC
444	with model GTRGAMMA, -f a and 100 bootstraps (53).
445	Minimum spanning tree (MST) analysis was built based on core gene SNPs produced
446	from a cluster analysis. The cluster analysis of ST631 was performed using a custom core
447	genome multi-locus sequence type (cgMLST) analysis using RidomSeqSphere+software v3.2.1
448	(http://www.ridom.de.seqsphere, Ridom GmbH, Münster, Germany) as previously described
449	(31). Briefly, the software first defines a cgMLST scheme using the target definer tool with
450	default settings using the PacBio generated MAVP-Q genome as the reference. Then, five other
451	V. parahaemolyticus genomes (BB22OP, CDC_K4557, FDA_R31, RIMD2210633, and UCM-
452	V493) were used for comparison with the reference genome to establish the core and accessory
453	genome genes. Genes that are repeated in more than one copy in any of the six genomes were
454	removed from the analysis. Subsequently, a task template was created that contains both core and
455	accessory genes. Each individual gene locus from MAVP-Q was assigned allele number 1. Then

456	each ST631 isolate genome assembly was queried against the task template, where any locus that
457	differed from the reference genome or any other queried genome was assigned a new allele
458	number. The cgMLST performed a gene-by-gene analysis of all core genes (excluding accessory
459	genes) and identified SNPs within different alleles to establish genetic distance calculations.
460	
461	Configuration and distribution of VPaIs
462	The VPaI sequence from the PacBio sequenced genomes of MAVP-Q and MAVP-R
463	were identified by comparison with the published RIMD2210633 VPaI-7 (NC_004605 region
464	between VPA1312 – VPA1395) and VPaI _{TH3996} (AB455531) (16). Identification of the complete
465	MAVP-Q VPaI γ and genomic junctures in chromosome II was done by comparison with the
466	same region of chromosome II in MAVP-R and G149 (which lack an island in this location)
467	using Mauve (54). In a reciprocal manner, the absence of an island in chromosome I in MAVP-Q
468	and G149 was assessed by comparison with chromosome I of MAVP-R. MAVP-Q VPaI γ
469	(MF066646) and MAVP-R VPaI β (MF066647) were then extracted as a single contiguous
470	sequence and annotated using Prokka 1.9. Gene content and order of the VPaI elements in
471	MAVP-Q, MAVP-R and RIMD2210633 were then illustrated by Easyfig (51). Roary (55) was
472	then employed to determine homologs among VPaIs based on each island's annotated sequences
473	with identity set at 50%. Identification of the genome locations of VPaI β in ST1127 isolate
474	MAVP-M (accession number GCA_001023155) and for VPaI γ in AQ4037 (accession number
475	GCA_000182365) (17) was also done using Mauve (54).
476	To examine the distribution of the VPaI γ in all publicly available draft genomes
477	(https://www.ncbi.nlm.nih.gov/genome/genomes/691, 2016) and genomes from archived
478	regional isolates, whole draft genome sequences were aligned to a 6,118 bp subsequence of the

479	MAVP-Q VPaI with NASP version 1.0.2 (56) (https://pypi.python.org/pypi/nasp/1.0.2, 2017).
480	This subsequence spanned the unique juncture of the four conserved hypothetical proteins
481	(AB831_22090, AB831_22095, AB831_22100, AB831_22105) with the adjacent inserted <i>tdh</i>
482	(AB831_22110, c.a. 2549 bp upstream of <i>ure</i> cluster)(Supplemental Fig. 1). Percent coverage of
483	the reference sequence was used to determine whether each genome harbored only the four
484	hypothetical proteins, only a <i>tdh</i> gene, or the entire module including the fusion of the four genes
485	with <i>tdh</i> (Supplemental Fig. 1 and Supplemental Table 3). The sequence type of each genome
486	harboring the fused element characteristic of VPaI γ was then determined using the SRST2
487	pipeline (48). Where sequencing reads were not available as the input for SRST2, they were
488	simulated from assemblies using an in-house Python script
489	(https://github.com/kpdrees/fasta2reads).
490	A PCR amplification approach was developed and applied to survey the presence of <i>tdh</i>

491 adjacent to the *ure* gene cluster. Primers were designed to conserved sequences of the 3' end of

- 492 *tdh* (PIHybF8: 5'GCCAACATGGATATAAAAAAATGA3') and the 5' end of *ureG*
- 493 (tdhUreGrev5: 5'GACAAAGGTATGCTGCCAAAAGTG3') as determined by gene alignments,
- 494 which when used together produced a 2631 bp amplicon of the insertion juncture when used with
- 495 MAVP-Q as a template (Supplemental Fig. 4). Amplification was performed on purified DNA
- 496 with Accustart enzyme mix, with an initial denaturation at 94°C for 3 min., followed by 30
- 497 cycles of a denaturation at 94°C for 1 min, annealing at 61°C for 1 min, and amplification at 72°C
- 498 for 2.5 min, with a final elongation at 72°C for 5 min. This amplification was performed in
- 499 parallel with a diagnostic multiplex PCR amplification of *tdh*, *trh* and *tlh* using published
- 500 methods (10, 57) to investigate the co-occurrence of VPaIγ with both hemolysin encoding genes

- 501 in representative isolates of various clinically prevalent sequence types. Amplicons were
- 502 visualized using a 1.2% agarose gel in TAE buffer (Supplemental Fig. 4).
- 503
- 504 Nucleotide sequence accession numbers.
- 505 The accession number of Pacific Biosciences sequenced genome for MAVP-Q is
- 506 MDWT00000000, and for MAVP-R is MPPP00000000. The accession number of Illumina
- sequenced draft genome for G6928 is MPPN00000000, for MA561 is MPPM00000000 and for
- 508 G149 is MPPO00000000. Detailed information about all other ST631 isolate draft genomes were
- described previously (31) and are listed in Supplemental Table 1. The accessions for the short
- 510 reads for the remaining sequenced genomes are listed in Supplemental Table 4. The accession
- 511 number of VPaIβ from MAVP-R is MF066647 and the accession number of VPaIγ from MAVP-
- 512 Q is MF066646.
- 513

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515

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536								
537	REFERENCES							
538	1.	Hiyoshi H, Kodama T, Iida T, Honda T. 2010. Contribution of Vibrio						
539		parahaemolyticus virulence factors to cytotoxicity, enterotoxicity, and lethality in mice.						
540		Infect Immun 78: 1772-1780.						
541	2.	Scallan E, Hoekstra RM, Angulo FJ, Tauxe RV, Widdowson M-A, Roy SL, Jones						
542		JL, Griffin PM. 2011. Foodborne illness acquired in the United States—major						
543		pathogens. Emerg Infect Dis 17(1):7-15.						
544	3.	Hazen TH, Pan L, Gu J-D, Sobecky PA. 2010. The contribution of mobile genetic						
545		elements to the evolution and ecology of Vibrios. FEMS Microbiol Ecol 74:485-499.						

546	4.	Hurley CC, Quirke A, Reen FJ, Boyd EF. 2006. Four genomic islands that mark post-
547		1995 pandemic Vibrio parahaemolyticus isolates. BMC Genomics 7:104
548		DOI:110.1186/1471-2164-1187-1104.
549	5.	Boyd EF, Cohen AL, Naughton LM, Ussery DW, Binnewies TT, Stine OC, Parent
550		MA. 2008. Molecular analysis of the emergence of pandemic Vibrio parahaemolyticus.
551		BMC Microbiol 8:110.
552	6.	Kishishita M, Matsuoka N, Kumagai K, Yamasaki S, Takeda Y, Nishibuchi M. 1992.
553		Sequence variation in the thermostable direct hemolysin-related hemolysin (trh) gene of
554		Vibrio parahaemolyticus. Appl Environ Microbiol 58:2449-2457.
555	7.	Honda T, Ni Y, Miwatani T, Adachi T, Kim J. 1992. The thermostable direct
556		hemolysin of Vibrio parahaemolyticus is a pore-forming toxin. Can J Microbiol 38:1175-
557		1180.
558	8.	Park K-S, Ono T, Rokuda M, Jang M-H, Iida T, Honda T. 2004. Cytotoxicity and
559		enterotoxicity of the thermostable direct hemolysin-deletion mutants of Vibrio
560		parahaemolyticus. Microbiol Immunol 48:313-318.
561	9.	Shirai H, Ito H, Hirayama T, Nakamoto Y, Nakabayashi N, Kumagai K, Takeda Y,
562		Nishibuchi M. 1990. Molecular epidemiologic evidence for association of thermostable
563		direct hemolysin (TDH) and TDH-related hemolysin of Vibrio parahaemolyticus with
564		gastroenteritis. Infect Immun 58:3568-3573.
565	10.	Panicker G, Call DR, Krug MJ, Bej AK. 2004. Detection of pathogenic Vibrio spp. in
566		shellfish by using multiplex PCR and DNA microarrays. Appl Environ Microbiol
567		70: 7436-7444.

568	11.	Nishibuchi M, Kaper JB. 1995. Thermostable direct hemolysin gene of Vibrio
569		parahaemolyticus: a virulence gene acquired by a marine bacterium. Infect Immun
570		63: 2093.
571	12.	Park K-S, Ono T, Rokuda M, Jang M-H, Okada K, Iida T, Honda T. 2004.
572		Functional characterization of two type III secretion systems of Vibrio parahaemolyticus.
573		Infect Immun 72: 6659-6665.
574	13.	Broberg CA, Calder TJ, Orth K. 2011. Vibrio parahaemolyticus cell biology and
575		pathogenicity determinants. Microb Infect 13:992-1001.
576	14.	Zhang L, Orth K. 2013. Virulence determinants for Vibrio parahaemolyticus infection.
577		Curr Opin Microbiol 16:70-77.
578	15.	Makino K, Oshima K, Kurokawa K, Yokoyama K, Uda T, Tagomori K, Iijima Y,
579		Najima M, Nakano M, Yamashita A. 2003. Genome sequence of Vibrio
580		parahaemolyticus: a pathogenic mechanism distinct from that of V. cholerae. The Lancet
581		361: 743-749.
582	16.	Okada N, Iida T, Park K-S, Goto N, Yasunaga T, Hiyoshi H, Matsuda S, Kodama T,
583		Honda T. 2009. Identification and characterization of a novel type III secretion system in
584		trh-positive Vibrio parahaemolyticus strain TH3996 reveal genetic lineage and diversity
585		of pathogenic machinery beyond the species level. Infect Immun 77:904-913.
586	17.	Chen Y, Stine OC, Badger JH, Gil AI, Nair GB, Nishibuchi M, Fouts DE. 2011.
587		Comparative genomic analysis of Vibrio parahaemolyticus: serotype conversion and
588		virulence. BMC Genomics 12:1.
589	18.	Zhou X, Gewurz BE, Ritchie JM, Takasaki K, Greenfeld H, Kieff E, Davis BM,
590		Waldor MK. 2013. vopZ A Vibrio parahaemolyticus T3SS effector mediates

591 pathogenesis by independently enabling intestinal colonization and inhibiting TAK1

592 activation. Cell Reports **3**:1690-1702.

- 593 19. Hubbard TP, Chao MC, Abel S, Blondel CJ, zur Wiesch PA, Zhou X, Davis BM,
- 594 Waldor MK. 2016. Genetic analysis of *Vibrio parahaemolyticus* intestinal colonization.
- 595 Proc Nat Acad Sci USA **113**:6283-6288.
- 596 20. Ronholm J, Petronella N, Leung CC, Pightling A, Banerjee S. 2016. Genomic
- 597 Features of Environmental and Clinical *Vibrio parahaemolyticus* Isolates Lacking
- 598 Recognized Virulence Factors Are Dissimilar. Appl Environ Microbiol **82:**1102-1113.
- 599 21. Xu F, Ilyas S, Hall JA, Jones SH, Cooper VS, Whistler CA. 2015. Genetic
- 600 characterization of clinical and environmental Vibrio parahaemolyticus from the
- 601 Northeast USA reveals emerging resident and non-indigenous pathogen lineages. Name:
- 602 Front Microbiol **6:**272.
- 603 22. Banerjee SK, Kearney AK, Nadon CA, Peterson C-L, Tyler K, Bakouche L, Clark
- 604 CG, Hoang L, Gilmour MW, Farber JM. 2014. Phenotypic and genotypic
- 605 characterization of Canadian clinical isolates of *Vibrio parahaemolyticus* collected from

606 2000 to 2009. J Clin Microbiol **52:**1081-1088.

- 607 23. Turner JW, Paranjpye RN, Landis ED, Biryukov SV, González-Escalona N, Nilsson
- 608 WB, Strom MS. 2013. Population structure of clinical and environmental *Vibrio*
- 609 *parahaemolyticus* from the Pacific Northwest coast of the United States. PLoS ONE
- 610 **8(2):**e55726
- 611 24. Jones JL, Lüdeke CH, Bowers JC, Garrett N, Fischer M, Parsons MB, Bopp CA,
- 612 **DePaola A.** 2012. Biochemical, serological, and virulence characterization of clinical and
- 613 oyster *Vibrio parahaemolyticus* isolates. J Clin Microbiol **50**(7):2343-2352.

614	25.	DePaola A	, Ulaszek J, Ka	vsner CA,	Tenge BJ.	, Nordstrom JL,	Wells J, Puhr	N,
· · ·			,	,,				• •

- 615 Gendel SM. 2003. Molecular, serological, and virulence characteristics of Vibrio
- 616 *parahaemolyticus* isolated from environmental, food, and clinical sources in North
- 617 America and Asia. Appl Environ Microbiol **69:**3999-4005.
- 618 26. Haendiges J, Timme R, Allard MW, Myers RA, Brown EW, Gonzalez-Escalona N.
- 619 2015. Characterization of *Vibrio parahaemolyticus* clinical strains from Maryland (2012–
- 620 2013) and comparisons to a locally and globally diverse *V. parahaemolyticus* strains by
- 621 whole-genome sequence analysis. Front Microbiol **6**.125
- 622 27. Ellis CN, Schuster BM, Striplin MJ, Jones SH, Whistler CA, Cooper VS. 2012.
- 623 Influence of seasonality on the genetic diversity of *Vibrio parahaemolyticus* in New
- Hampshire shellfish waters as determined by multilocus sequence analysis. Appl Environ
 Microbiol **78**:3778-3782.
- 626 28. Nair GB, Ramamurthy T, Bhattacharya SK, Dutta B, Takeda Y, Sack DA. 2007.
- 627 Global dissemination of *Vibrio parahaemolyticus* serotype O3: K6 and its serovariants.
- 628 Clin Microbiol Rev **20:**39-48.
- 629 29. Martinez-Urtaza J, Baker-Austin C, Jones JL, Newton AE, Gonzalez-Aviles GD,
- 630 DePaola A. 2013. Spread of Pacific Northwest *Vibrio parahaemolyticus* strain. N Engl J
 631 Med 369:1573-1574.
- 632 30. Newton AE, Garrett N, Stroika SG, Halpin JL, Turnsek M, Mody RK, Division of
- 633 Foodborne W, Environmental D. 2014. Notes from the field: Increase in *Vibrio*
- 634 *parahaemolyticus* infections associated with consumption of Atlantic coast shellfish—
- 635 2013. MMWR Morb Mortal Wkly Rep **63**:335-336.

636	31.	Xu F, Gonzalez-Escalona N, Haendiges J, Myers RA, Ferguson J, Stiles T, Hickey E,
-----	-----	---

- 637 Moore M, Hickey JM, Schillaci C. 2017. Sequence type 631 Vibrio parahaemolyticus,
- 638 an emerging foodborne pathogen in North America. J Clin Microbiol **55:**645-648.
- 639 32. Lüdeke CH, Gonzalez-Escalona N, Fischer M, Jones JL. 2015. Examination of
- 640 clinical and environmental *Vibrio parahaemolyticus* isolates by multi-locus sequence
- 641 typing (MLST) and multiple-locus variable-number tandem-repeat analysis (MLVA).
- 642 Frontiers in microbiology **6**.564
- 643 33. González-Escalona N, Gavilan RG, Brown EW, Martinez-Urtaza J. 2015.
- 644 Transoceanic spreading of pathogenic strains of *Vibrio parahaemolyticus* with distinctive
- 645 genetic signatures in the recA gene. PloS one **10**:e0117485.
- 646 34. Park K-S, Suthienkul O, Kozawa J, Yamaichi Y, Yamamoto K, Honda T. 1998.
- 647 Close proximity of the *tdh*, *trh* and *ure* genes on the chromosome of *Vibrio*
- 648 *parahaemolyticus*. Microbiology **144:**2517-2523.
- 649 35. Johnson C, Flowers A, Young V, Gonzalez-Escalona N, DePaola A, Noriea III N,
- 650 **Grimes D.** 2009. Genetic relatedness among *tdh*+ and *trh*+ *Vibrio parahaemolyticus*
- 651 cultured from Gulf of Mexico oysters (Crassostrea virginica) and surrounding water and
- 652 sediment. Microb Ecol **57:**437-443.
- 653 36. González-Escalona N, Martinez-Urtaza J, Romero J, Espejo RT, Jaykus L-A,
- 654 DePaola A. 2008. Determination of molecular phylogenetics of *Vibrio parahaemolyticus* 655 strains by multilocus sequence typing. J Bacteriol 190:2831-2840.
- 656 37. Ellingsen BA, Olsen JS, Granum PE, Rorvik LM, González-Escalona N. 2013.
- 657 Genetic characterization of trh positive Vibrio spp. isolated from Norway. Front Cell
- 658 Infect Microbiol **3:**107.

- 659 38. Chen Y, Dai J, Morris JG, Johnson JA. 2010. Genetic analysis of the capsule
- 660 polysaccharide (K antigen) and exopolysaccharide genes in pandemic Vibrio
- 661 *parahaemolyticus* O3: K6. BMC Microbiol **10:**1.
- 662 39. Meibom KL, Blokesch M, Dolganov NA, Wu C-Y, Schoolnik GK. 2005. Chitin
- 663 induces natural competence in *Vibrio cholerae*. Science **310**:1824-1827.
- 40. Takemura AF, Chien DM, Polz MF. 2014. Associations and dynamics of Vibrionaceae
- in the environment, from the genus to the population level. Front Microbiol 5:38.
- 41. **Lovell CR.** 2017. Ecological fitness and virulence features of *Vibrio parahaemolyticus* in
- 667 estuarine environments. Appl Microbiol Biotechnol 101:1781-1794.
- 42. **Johnson CN.** 2013. Fitness factors in vibrios: a mini-review. Microb Ecol **65**:826-851.
- 43. Matz C, Nouri B, McCarter L, Martinez-Urtaza J. 2011. Acquired type III secretion
- 670 system determines environmental fitness of epidemic *Vibrio parahaemolyticus* in the
- 671 interaction with bacterivorous protists. PloS one 6:e20275.
- 672 44. Nishibuchi M, Kaper JB. 1985. Nucleotide sequence of the thermostable direct
- hemolysin gene of *Vibrio parahaemolyticus*. J Bacteriol **162**:558-564.
- 45. Baym M, Kryazhimskiy S, Lieberman TD, Chung H, Desai MM, Kishony R. 2015.
- 675 Inexpensive multiplexed library preparation for megabase-sized genomes. PloS one
- 676 **10:**e0128036.
- 677 46. Tritt A, Eisen JA, Facciotti MT, Darling AE. 2012. A5. An integrated pipeline for *de* 678 *novo* assembly of microbial genomes. PLoS ONE 7:e42304.
- 679 47. Seemann T. 2014. Prokka: rapid prokaryotic genome annotation. Bioinformatics.
 680 30:2068-9

- 681 48. Inouve M, Conway TC, Zobel J, Holt KE. 2012. Short read sequence typing (SRST): 682 multi-locus sequence types from short reads. BMC Genomics 13:338.
- 683 49. Walker BJ, Abeel T, Shea T, Priest M, Abouelliel A, Sakthikumar S, Cuomo CA,
- 684 Zeng Q, Wortman J, Young SK. 2014. Pilon: an integrated tool for comprehensive
- 685 microbial variant detection and genome assembly improvement. PloS one 9:e112963.
- 686 50. Camacho C, Coulouris G, Avagyan V, Ma N, Papadopoulos J, Bealer K, Madden
- 687 TL. 2009. BLAST+: architecture and applications. BMC Bioinformatics 10:421.
- 688 51. Sullivan MJ, Petty NK, Beatson SA. 2011. Easyfig: a genome comparison visualizer. 689 Bioinformatics 27:1009-1010.
- 690 52. Gardner SN, Slezak T, Hall BG. 2015. kSNP3. 0: SNP detection and phylogenetic
- 691 analysis of genomes without genome alignment or reference genome. Bioinformatics 692 **31:**2877-8.
- 693 53. Stamatakis A. 2006. RAxML-VI-HPC: maximum likelihood-based phylogenetic 694
- analyses with thousands of taxa and mixed models. Bioinformatics **22**:2688-2690.
- 695 54. Darling AC, Mau B, Blattner FR, Perna NT. 2004. Mauve: multiple alignment of
- 696 conserved genomic sequence with rearrangements. Genome Res 14:1394-1403.
- 697 55. Page AJ, Cummins CA, Hunt M, Wong VK, Reuter S, Holden MT, Fookes M,
- 698 Falush D, Keane JA, Parkhill J. 2015. Roary: rapid large-scale prokaryote pan genome 699 analysis. Bioinformatics 31:3691-3693.
- 700 56. Sahl JW, Lemmer D, Travis J, Schupp J, Gillece J, Aziz M, Driebe E, Drees K,
- 701 Hicks N, Williamson C. 2016. The Northern Arizona SNP Pipeline (NASP): accurate,
- 702 flexible, and rapid identification of SNPs in WGS datasets. Microb Genom. 2:e000074

703	57.	Whistler CA, Hall JA, Xu F, Ilyas S, Siwakoti P, Cooper VS, Jones SH. 2015. Use of
704		Whole-Genome Phylogeny and Comparisons for Development of a Multiplex PCR Assay
705		To Identify Sequence Type 36 Vibrio parahaemolyticus. J Clin Microbiol 53:1864-1872.
706	58.	Jolley KA, Chan M-S, Maiden MC. 2004. mlstdbNet-distributed multi-locus sequence
707		typing (MLST) databases. BMC Bioinformatics 5:86.
708	59.	Alikhan N-F, Petty NK, Zakour NLB, Beatson SA. 2011. BLAST Ring Image
709		Generator (BRIG): simple prokaryote genome comparisons. BMC Genomics 12:402
710		

711

712	Table 1: Clinical and environmental	prevalence of emergent Northeast US V.	parahaemolvticus

713	lingage	with	associated	virulanca	fasturas
115	micages	with	associated	viruiciice	icatures.

Sequence		t US States ^b		Г Database ^с	Hemolysin	VPaI
type ^a	Clinical	Environmental	Clinical	Environmental	genotype	type ^d
3	2	0	217	33	tdh^+	α
36	91	1	58	5	tdh^+trh^+	γ
	24	0	12	0	tdh^+trh^+	γ
631	1^{e}	2	0	0	trh^+	β
	0	1	0	0	neither	absent
43	5	0	17	4	tdh^+trh^+	γ
636	4	0	2	0	tdh^+trh^+	γ
1127	4	0	0	0	trh^+	β
110	3	0	0	1	tdh^+trh^+	γ
34/324	2	2	4	19	tdh^+trh^+	γ
674	0	4	1	20	tdh^+trh^+	γ
0/4	1	0	0	0	neither	absent
308	2	0	0	2	tdh^+trh^+	γ
12	2	0	0	4	trh^+	β
162	2	0	1	1	neither	absent
194	2	0	1	0	neither	absent
809	2	0	0	1	trh^+	β
1716	2	0	0	0	trh^+	β
1123	1	1	0	0	trh^+	β
8	1	0	13	5	trh^+	β
23	1	0	0	3	tdh^+trh^+	γ
749	1	0	1	0	tdh^+trh^+	γ
1295	1	0	0	1	neither	absent
134	1	0	1	0	neither	absent
741	1	0	0	1	neither	absent
98	1	0	0	1	trh^+	β
1205	1	0	0	1	neither	absent
1561	1	0	0	0	neither	absent
1717	1	0	0	0	neither	absent
1725	1	0	0	0	tdh^+	α

714

^a Some clinical isolates had insufficient sequencing coverage to determine sequence type and included

715 eight tdh^+trh^+ isolates, one tdh^+ isolate, four trh^+ isolates, and 11 isolates without hemolysins, some of 716 which were from wound infections. Two wound infection isolates lacking hemolysins were of known

717 sequence types and are not listed above.

718 ^bData generated from all available gastric infection clinical and environmental isolates four reporting

719 Northeast US States including ME, NH, MA, and CT between 2010 and 2016.

720 ^chttp://pubmlst.org/vparahaemolyticus, 2017 (36, 58)

721 ^dPresence of the VPaly architecture was determined by PacBio genome sequencing of strain MAVP-Q

722 and MAVP-26, whereas for other strains, identification of VPaI type was determined through illumina 723 genome sequencing, PCR amplification and Sanger sequencing.

724 ^eThis single isolate harbors a *recA* allele (allele 21) typical of ST631 fused to allele 107 through an

725 insertion event, generating a hybrid allele previously described (33).

726

727

728 Figure 1. Schematic of a horizontally acquired insertion in the *recA*-encoding region of MAVP-R. 729 Sequences of the *recA* gene and flanking region from MAVP-Q (reference ST631 genome), 730 MAVP-R, Asia-derived isolates \$130/\$134 and Peru-derived isolate 090-96 were extracted and 731 aligned. Open reading frames designated with arrows and illustrated by representative colors to 732 highlight homologous and unique genes. The % similarity between homologs is illustrated by 733 grey bars. 734 735 Figure 2. Phylogenetic relationships of V. parahaemolyticus lineages and identification of 736 distinct ST631 clades. An ML phylogeny of representative V. parahaemolyticus genomes of 737 clinical strains causing two or more infections was built on whole genome SNPs identified by 738 reference-free comparisons as described in the methods. The branch length represents the 739 number of nucleotide substitutions per site. Numbers at nodes represent percent bootstrap 740 support where unlabeled nodes had bootstraps of less than 70. 741 742 Figure 3. Minimum spanning tree relationships among clade I and clade II ST631. A cgMLST 743 core gene-by-gene analysis (excluding accessory genes) was performed and SNPs were 744 identified within different alleles. The numbers above the connected lines (not to scale) represent 745 SNP differences. The isolates are colored based on different hemolysin genotypes as labeled. 746 747 Figure 4. Comparisons of the pathogenicity islands containing hemolysins and T3SS2. 748 Sequences of VPaI were extracted from select genomes and aligned. VPaIa was derived from 749 ST3 strain RIMD2210633, VPaly was derived from ST631 clade II isolate MAVP-Q, and VPalß 750 was derived from ST631 clade I isolate MAVP-R. ORFs are depicted in defined colors and

- similarities (\geq 75%) among ORFs are illustrated in grey blocks. Homologs between VPaIa and
- 752 VPaI β/γ (50>75% identity) are named and listed in Supplemental Table 2.
- 753
- 754 Figure 5. Distribution of VPaIy in emergent pathogen lineages. The presence of *tdh*, *trh* and
- 755 VPaIγ along with positive control *tlh* was determined by PCR amplification using gene-specific
- primers and visualized on a 1.2% agarose gel. The order from left to right is 1kb+ ladder, ST3
- 757 (MAVP-C), ST36 (MAVP-26), ST631 CII (clade II isolate MAVP-Q), ST631 CI (clade I
- 758 isolates MAVP-R and G149), ST43 (MAVP-71), ST636 (MAVP-50), ST1127 (MAVP-M),
- 759 ST110 (MAVP-46), ST34 (CTVP19C), ST324 (MAVP-14), and ST674 (CT4291, MAVP21).
- 760 The corresponding sizes of the ladder fragments are as labeled to the left and the identity of the
- amplicons listed to the right of the gel image.

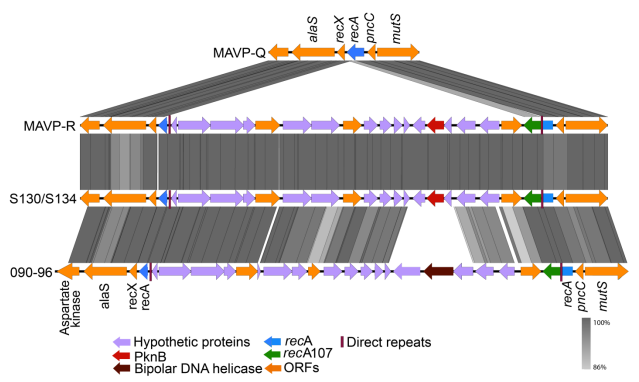


Figure 1. Schematic of a horizontally acquired insertion in the *recA*-encoding region of MAVP-R. Sequences of the *recA* gene and flanking region from MAVP-Q (reference ST631 genome), MAVP-R, Asia-derived isolates S130/S134 and Peru-derived isolate 090-96 were extracted and aligned. Open reading frames designated with arrows and illustrated by representative colors to highlight homologous and unique genes. The % similarity between homologs is illustrated by grey bars.

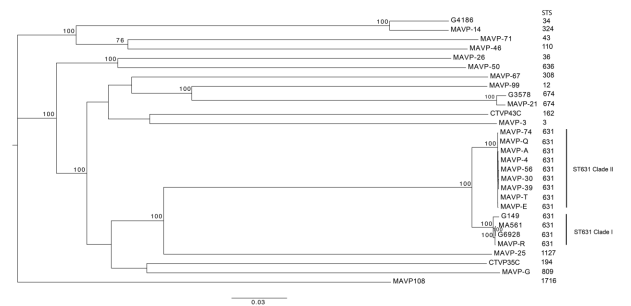


Figure 2. Phylogenetic relationships of *V. parahaemolyticus* lineages and identification of distinct ST631 clades. An ML phylogeny of representative *V. parahaemolyticus* genomes of clinical strains causing two or more infections was built on whole genome SNPs identified by reference-free comparisons as described in the methods, The branch length represents the number of nucleotide substitutions per site. Numbers at nodes represent percent bootstrap support where unlabeled nodes had bootstraps of less than 70.

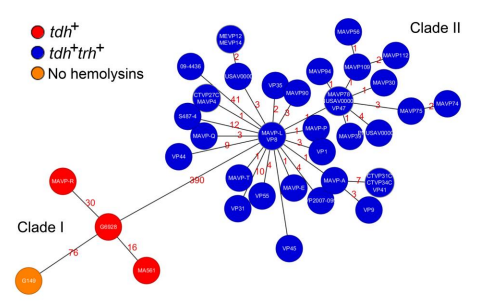


Figure 3. Minimum spanning tree relationships among clade I and clade II ST631. A core geneby-gene analysis (excluding accessory genes) was performed and SNPs were identified within different alleles. The numbers above the connected lines (not to scale) represent SNP differences. The isolates are colored based on different hemolysin genotypes as labeled.

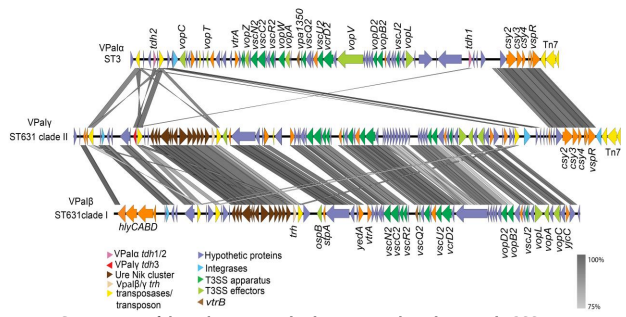


Figure 4. Comparisons of the pathogenicity islands containing hemolysins and T3SS2. Sequences of VPaI were extracted from select genomes and aligned. VPaI α was derived from ST3 strain RIMD2210633, VPaI γ was derived from ST631 clade II isolate MAVP-Q, and VPaI β was derived from ST631 clade I isolate MAVP-R. ORFs are depicted in defined colors and similarities (\geq 75%) among ORFs are illustrated in grey blocks. omologs between VPaI α and VPaI β/γ (50>75% identity) are named and listed in supplemental table 2.

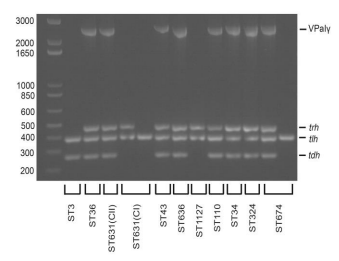


Figure 5. Distribution of VPaIy in emergent pathogen lineages. The presence of *tdh*, *trh* and VPaIy along with positive control *tlh* was determined by PCR amplification using gene-specific primers and visualized on a 1.2% agarose gel. The order from left to right is 1kb+ ladder, ST3 (MAVP-C), ST36 (MAVP-26), ST631 CII (clade II isolate MAVP-Q), ST631 CI (clade I isolates MAVP-R and G149), ST43 (MAVP-71), ST636 (MAVP-50), ST1127 (MAVP-M), ST110 (MAVP-46), ST34 (CTVP19C), ST324 (MAVP-14), and ST674 (CT4291, MAVP21). The corresponding sizes of the ladder fragments are as labeled to the left and the identity of the amplicons listed to the right of the gel image.