Toxigenic *Vibrio cholerae* evolution and establishment of reservoirs in aquatic ecosystems

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The spread of cholera in the midst of an epidemic is largely driven by direct transmission from person to person, although it is well-recognized that *Vibrio cholerae* is also capable of growth and long-term survival in aquatic ecosystems. While prior studies have shown that aquatic reservoirs are important in the persistence of the disease on the Indian subcontinent, an epidemiological view postulating that locally evolving environmental *V. cholerae* contributes to outbreaks outside Asia remains debated. The single-source introduction of toxigenic *V. cholerae* O1 in Haiti, one of the largest outbreaks occurring this century, with 812,586 suspected cases and 9,606 deaths reported through July 2018, provided a unique opportunity to evaluate the role of aquatic reservoirs and assess bacterial transmission dynamics across environmental boundaries. To this end, we investigated the phylogeography of both clinical and aquatic toxigenic *V. cholerae* O1 isolates and show robust evidence of the establishment of aquatic reservoirs as well as ongoing evolution of *V. cholerae* isolates from aquatic sites. Novel environmental lineages emerged from sequential population bottlenecks, carrying mutations potentially involved in adaptation to the aquatic ecosystem. Based on such empirical data, we developed a mixed-transmission dynamic model of *V. cholerae*, where aquatic reservoirs actively contribute to genetic diversification and epidemic emergence, which underscores the complexity of transmission pathways in epidemics and endemic settings and the need for long-term investments in cholera control at both human and environmental levels.

Significance

Persistent aquatic environmental reservoirs for *Vibrio cholerae* O1 are present in Asia; however, their existence in other parts of the world remains controversial. The single-source introduction of toxigenic *V. cholerae* O1 in Haiti provides a unique opportunity to assess the potential role played by aquatic reservoirs in subsequent seasonal outbreaks. Whole-genome sequence Bayesian phylogeography showed robust evidence of *V. cholerae* O1 evolution in riverine sites, through the establishment of reservoirs, during lull periods of the Haitian epidemic. Novel lineages emerged in the environment from sequential population bottlenecks, characterized by mutations in genes potentially involved in adaptive response. The data highlight the need for long-term investments in cholera control at both human and environmental levels.

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The authors declare no competing interest.

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Data deposition: The genomic sequences reported in this paper have been deposited with the National Center for Biotechnology Information (NCBI) in the Sequence Read Archive, https://www.ncbi.nlm.nih.gov/nuccore (BioProject ID code PRJNA4510624). R scripts and xml files are available on GitHub (https://github.com/salemilab/environmental_cholera_haiti).

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aquatic ecosystems (2), elucidating the role of aquatic environmental reservoirs has become critical to the development of elimination strategies.

Results

Four departments—Artibonite, Centre, Nord-Ouest, and Ouest—account for 90% of reported cholera cases in Haiti (9). We focused our efforts on the Ouest Department (Fig. 1A), a coastal region rich in aquatic ecosystems where we have isolated toxicogenic V. cholerae O1 from fixed riverine and estuarine sites (Fig. 1A). The prevalence of toxicogenic aquatic V. cholerae O1 correlated with increased temperature and rainfall but not with fecal coliform counts, suggesting that the presence was not due to fecal contamination alone (10). Monthly case counts in Ouest, oscillating yearly but decreasing during the dry season and increasing during the rainy season (April to October), correlated with increased temperature and rainfall (Fig. 1B). This is in agreement with patterns of endemic V. cholerae O1 in Bangladesh and Peru, where seasonal cholera outbreaks correlate with increased V. cholerae O1 in the environment and increases in water temperature or rainfall (11, 12).

To test the hypothesis that V. cholerae established aquatic reservoirs in the Haitian river system, contributing to recurrent seasonal epidemics, we carried out whole-genome sequencing of 27 aquatic environmental toxigenic V. cholerae O1 strains, isolated in Gressier between 2012 and 2015, and 89 clinical strains collected in Ouest and Artibonite between 2010 and 2015 (Fig. 1B and Datasets S1 and S2). We investigated their phylogenetic relationship by maximum-likelihood (ML) (SI Appendix, Fig. S1C) and ML ancestral-state reconstruction (Fig. 1C) based on high-quality single-nucleotide polymorphisms (hqSNPs) (Dataset S3), after confirming the presence of a robust phylogenetic signal (SI Appendix, Fig. S1A and B). Environmental strains are intermixed with clinical strains in each major phylogenetic clade, including isolates from seasonal epidemic waves. Connection of epidemic waves through environmental lineages suggests that such lineages played a substantial role in promoting the epidemic, rather than simply providing an occasional contribution.

Sufficient temporal signal was present (SI Appendix, Fig. S1 D and E) to calibrate a reliable molecular clock (SI Appendix, Table S1) and infer time-scaled phylogenies. We reconstructed the spatiotemporal spread of the epidemic and quantified the potential contribution of environmental V. cholerae to this spread using Bayesian phylogeography (13). In agreement with previous findings (14), the maximum clade credibility (MCC) time-scaled tree obtained by discrete trait analysis (DTA) (Fig. 2A) and by Bayesian structured coalescent approximation (BASTA) (SI Appendix, Fig. S2A) showed a staircase phylogeny typical of evolution driven by repeated selective sweeps through sequential population bottlenecks with a major surviving lineage leading, over time, to each subsequent wave (15). The surviving lineage is represented by 4 preenvironmental contribution unambiguous and unique SNPs (uuSNPs): 98 C→T (gene Vch1786_I10360), 32 G→A (gene Vch1786_I1120), 51 T→G (gene exeA), and 53 T→C (upstream of gene rseA); there are 10 uuSNPs linked to environmental contribution: 1 C→T (gene Vch1786_I0012), 3 C→T (gene Vch1786_I0051), 30 G→A (gene Vch1786_I0998), 44 A→G (gene Vch1786_I1539), 45 T→G (gene Vch1786_I1601), 52 G→A (gene rseA), 61 A→C (gene epsG), 69 C→A (gene Vch1786_I2482), 105 G→A (upstream of gene Vch1786_I0538), and 107 C→A (gene Vch1786_I0794); and there is 1 postenvironmental contribution uuSNP: 110 C→T (gene Vch1786_I0977) (SI Appendix, Fig. S3). Eleven waning lineages defined by uuSNPs, shared in both inferences, are summarized in SI Appendix, Table S2 (SI Appendix, Fig. S3). The classic source–sink cholera population dynamic, where each epidemic wave is essentially propagated by ongoing transmissions within the human population (source) followed by spillover into the environment (sink), predicts that the backbone path (trunk) of the tree, that is, the surviving lineate successfully propagated through time (Fig. 2A), would be occupied only by lineages of clinical origin. Therefore, we used DTA and BASTA to assess the origin (clinical or environmental) of the internal branches (ancestral cholera lineages) in the tree through ancestral-state reconstruction. Both methods consistently inferred aquatic environmental V. cholerae lineages along the trunk of the tree giving rise, in turn, to clinical isolates circulating during different epidemic waves (Fig. 2A and SI Appendix, Fig. S2A). The presence of environmental ancestors directly connected along the trunk of the tree was highly supported (posterior probability > 0.9) between May 2012 (95% highest posterior density [HPD] November 2011 to August 2012) and January 2014 (95% HPD November 2013 to April 2014) (Fig. 2A and SI Appendix, Fig. S2A and B). This finding provides robust evidence

Fig. 1. Cholera epidemic in the Ouest Department, Haiti, 2010 to 2015. (A) Geographical distribution of environmental (n = 27) and clinical (n = 89) isolates sampled from the Haitian toxigenic V. cholerae O1 lineage in the Ouest Department between 2010 and 2015. (B, Upper) Monthly case counts of cholera infections in the Ouest Department and mean monthly precipitation (mm/mo) between 2010 and 2015. (B, Lower) Temporal distribution of V. cholerae genomes sampled in Haiti and of the number of cases reported monthly to the World Health Organization from October 2010, the beginning of the epidemic, until December 2015 (ticks correspond to the month of January for each year indicated on the x axis). The size of the circle is proportional to the number of environmental (green) and clinical (violet) genomes sampled in our study for each year. (C) Ancestral-state reconstruction using maximum likelihood inferred with PastML from a rooted ML phylogenetic tree inferred with IQ-TREE (SI Appendix, Fig. S1E). ML tree clades are merged vertically (clade circles contain the number of tips of the initial tree contained in them) and horizontally (branch size corresponds to the number of times its subtree is found in the initial tree) to cluster independent events of the same kind.
of sustained replication and evolution of *V. cholerae* in the aquatic environment followed by reintroduction into the human population, that is, environmental isolates playing an active role in epidemic spread rather than contributing solely to dead-end transmission chains. Moreover, we observed the presence of a monophyletic clade in the phylogeny, which arose in 2012, dominated by environmental isolates with few intermixed clinical ones (Fig. 2A). Although we cannot exclude that the clade is an artifact of sampling bias, its presence in the tree suggests a concurrent scenario where independent evolution of *V. cholerae* in aquatic reservoirs can occasionally spill over to humans without necessarily contributing to an epidemic wave. Previous genetic studies suggest that the first cholera outbreak in the Port-au-Prince area, after the full period in 2014, was propagated by autochthonous local transmission in the south rather than introduction from the north (16). Such observations, together with our findings, re-inforce a scenario of undetected environmental reservoirs serving as a source for a later wave of clinical cases.

To investigate further, we quantified the contribution of *V. cholerae* environmental ancestors to the backbone path of the tree by calculating the proportion of time spent by isolates in environmental (river/estuarine) or clinical (human host) periods (Fig. 2B). The estimates support the concept that aquatic reservoirs were a main source of *V. cholerae* from June 2012 to May 2014 (60 to 80% of the trunk, \(P < 0.0001\)) (Fig. 2B and SI Appendix, Fig. S2C). Since Bayesian phylogeography reconstruction can be sensitive to unequal sampling size, all analyses were repeated on 10 additional datasets, each one including all 27 aquatic isolates as well as a random subsample of 27 clinical isolates. Results were in agreement with the findings for the full dataset (SI Appendix, Fig. S4). The majority of the transitions (connections between two neighboring nodes) in the MCC tree (76.6%) occurred between isolates of clinical origin, representing a large number of human-to-human transmission events throughout the epidemic (Fig. 2C). Yet, a proportion of all transitions (19.3%) occurred between aquatic isolates, with about half of these (83%) located in internal branches (i.e., connections between ancestral neighboring nodes). These data support, again, a scenario of independent replication/evolution in the aquatic environment. Clinical-to-environmental and environmental-to-clinical transitions were also present at a low proportion in the tree, indicating spillover between the environment and humans (Fig. 2C). Spillover between host and environment, as well as host colonization (17) or host-to-host transmission cycles (14), can cause bottlenecks that dramatically reduce the bacterial (population genetic measure) effective population size (\(N_e\)). Indeed, *V. cholerae* demographic history, inferred from the Bayesian time-scaled phylogeny (18), showed two population bottlenecks in mid-2012 and 2014 (SI Appendix, Fig. S2D). The demographic history also showed that, despite the low prevalence of clinical cases observed in 2014, bacterial diversification continued in the aquatic environment, as indicated by the \(N_e\) increase in 2012 to 2013 after the first bottleneck. *V. cholerae* is capable of surviving in aquatic environments due to its varied adaptive responses to stressors, including nutrient deprivation (19), changes in salinity, and temperature and predation by bacteriophages and protists (20). Persistence in the environment has been linked to a variety of phenotypes and mechanisms (21, 22), including various independent biofilm formation (23). To explore the rate at which toxigenic *V. cholerae* isolates replicated, we calculated the weighted average of synonymous substitution rates (\(K_{SS}\)) as a proxy for bacterial replication rate. To control for the potential bias of transient polymorphisms, \(K_{SS}\) was separately estimated along internal and external branches, of either environmental or clinical isolates,
of Bayesian phylogenies, after testing for phylogenetic quality and structure of the datasets and calibrating the best evolutionary model (SI Appendix, Figs. S5 and Tables S3 and S4). The environmental $K_S$ estimate for internal branches was about twice higher than the clinical ($P < 0.001$) (Fig. 2D and SI Appendix, Figs. S6 and Table S5), evidence of faster replication in isolates circulating in aquatic environmental sources. While not completely in keeping with prior findings that the human gut is the “optimal” environment for *V. cholerae* (24) replication, it is known that stress-induced mutations contribute to adaptive evolution (25) and bacteria increase their mutation rates in response to environmental stressors such as starvation (25). A faster replication of environmental isolates is expected to lead to accumulation of deleterious mutations in the population, with a large number of transient polymorphisms segregating on external branches of the phylogeny. Indeed, while $K_S$ in external branches of environmental phylogenies, that is, along lineages that have not propagated successfully, is significantly higher than in internal branches (Fig. 2D and SI Appendix, Table S5), the contrary is observed for clinical isolates (Fig. 2D and SI Appendix, Figs. S6 and Table S5).

To distinguish between evolution under selective pressure (adaption) or genetic drift, we calculated the nonsynonymous ($dN$) and synonymous substitution ($dS$) rate ratio ($dN/dS$) (SI Appendix, Figs. S6 and Table S5). The evolutionary dynamics of the clinical isolates was mainly driven by purifying selection ($dN$/

$dS = 0.6, P < 0.001$), consistent with a negative Tajima’s $D$ ($-2.3, P < 0.005$) and a population experiencing bottlenecks and/or selective sweeps. Negative selection is expected to drive the already-established epidemic in humans that has reached a peak in the adaptive landscape during an early epidemic stage driven by diversifying selection (14). In contrast, environmental isolates showed greater nonsynonymous rates ($dN/dS = 2.1, P < 0.001$), characteristic of a heterogeneous microbial population recently introduced to a new environment and undergoing fixation of new variant-driven diversifying selection. In fact, we found environmental-specific molecular imprints in the genomes of *V. cholerae* that successfully replicated and persisted in the aquatic environment. A total of seven nonsynonymous mutations in coding regions, potentially granting a selective advantage to the environmental population of *V. cholerae*, emerged along the backbone of the environmental lineage during 2012–2013 (Fig. 3 and SI Appendix, Table S7). Three environmental-specific mutations, two early in the backbone bifurcation (Fig. 3) and one in an internal branch (SI Appendix, Table S7), affected genes in the type II secretion system (TSSII): general secretion pathway proteins A and G, and K (SI Appendix, Table S7). The TSSII plays a pivotal role in the virulence and survival of *V. cholerae* in different niches, such aquatic reservoirs or human hosts (26). Other mutations found along internal branches affected genes related to environmental stress response, chemotaxis/motility of *V. cholerae* in response to fluctuating environmental cues, flagellum hook-length control, biofilm formation in the extraintestinal environment, and exponential growth in response to available nutrients (SI Appendix, Figs. S7 and Table S7). Environmental isolates that contributed successfully to the evolving environmental lineage were collected not only near hospitals and cholera treatment centers located in Gressier but also in the nearby Cape and Carrefour regions (Fig. 3).

The evidence of epidemiological and environmental evolutionary forces driving the Haitian epidemic motivated us to consider a dynamic model of cholera tracking transmission and population genetics through aquatic environments and hosts. As in prior models of cholera from our group (27), our underlying epidemiological model is based on the susceptible–infected–recovered with reservoir (SIRW) framework, an extension of the classic susceptible–infected–recovered (SIR) framework with an added compartment for pathogen concentration in an aquatic reservoir (W) (Fig. 4 and SI Appendix, Figs. S8 and S9). In keeping with Kirpich et al. (27), we allowed for replication outside of the host, along with seasonality in environmental transmission, shedding, and decay, as suggested by the significant correlation in case counts with precipitation. In addition to this epidemiological formulation, we considered an evolutionary component in our model. Both neutral drift and selection pressures are accounted for in our multifocus framework, where we measure $N_e$ by calculating genetic diversity and utilizing coalescent approaches (SI Appendix, Text). Consistent with other studies, we assumed a tradeoff between environmental survivability and fitness in host transmission for loci under selection. The results from model simulations recapitulate qualitative features of $N_e$ and monthly case counts (Fig. 4A and SI Appendix, Fig. S8).

First, the model output and data both display a rise and drop in $N_e$ initially reflecting rapid expansion followed by negative selection after the initial outbreak (Fig. 4A). Then came the observation of a second growth in $N_e$ in 2013, larger in extent than would be expected by host transmission alone, produced by a preceding seasonal flare-up and extending into a period of reducing incidence. Our hypothesis of diversification within the aquatic reservoir offers an explanation for this relative upsurge in $N_e$, and is supported by reasonable model simulations where inclusion of environmental replication allows for this evolutionary exploration in contrast to cases without environmental replication (Fig. 4A and SI Appendix, Fig. S8). In particular, during the late 2012- to mid-2013 peak, uuSNPs were detected in the following genes of the surviving environmental lineage: Vch1786_10012 (1 C→T), Vch1786_10051 gene (3 C→T), Vch1786_10098 (30 G→A), Vch1786_11539 (44 A→G), rscA (52 G→A), epsG (61 A→C), and Vch1786_100794 (107 C→A) (SI Appendix, Fig. S3). Mutations in these genes may have contributed to increased adaptation of environmental strains in the aquatic ecosystem, although future in vitro studies will be needed to investigate their contribution to *V. cholerae* fitness. The peak in environmental replication is followed by a substantial decline in $N_e$ beginning prior to the major lull period in 2014. Again, the case incidence does not fully explain the magnitude of this drop. While the population bottleneck induces a decrease in $N_e$, simulations suggest the environmental replication is necessary to accelerate fixation of genes adapted to persisting in the aquatic reservoir. At the end of 2014, environmental reservoirs spill over to hosts with a spike in clinical cases, possibly boosted by a beneficial host adaptation and/or loss of immunity in the host population. Extensive simulations calibrated to the case data suggest that in order to additionally fit the observed $N_e$, environmental replication should be included in the model. A major ramification of our findings is that the replication and adaptation within the aquatic reservoir may make control strategies targeting host transmission more difficult. Indeed, projecting forward from 2015 in simulations, we observed that vaccination can readily clear the pathogen in the absence of environmental replication (SI Appendix, Fig. S10). However, for environmental adaptation and replication consistent with the observed $N_e$, vaccinating a much larger portion of the population or adding an intervention affecting the aquatic reservoir is necessary to eliminate cholera in our dynamic mixed-transmission model (Fig. 4B).

**Discussion**

Our findings have direct applicability to the development of control and elimination strategies for this ancient and devastating disease. Assumptions that transmission outside of the Indian subcontinent occurs only at a direct person-to-person level have resulted in a major focus on “quick response” teams that go to case households to provide disinfection combined with a local emphasis on water, sanitation, and hygiene. Using Haiti as an example, our data underscore a greater complexity of the transmission process, illustrating the key role that aquatic reservoirs play in evolution, dissemination, and maintenance of the disease in an area well-removed from its Asian homeland. This translates into
a recognition that elimination of toxigenic \textit{V. cholerae} may be difficult, if not impossible, so long as environmental reservoirs are present. This, in turn, emphasizes the need for careful and ongoing environmental studies in areas where cholera cases have occurred and the need for a long-term control strategy, including long-term investments in water, sanitation, and hygiene infrastructure, even if cholera cases are not being identified. It also plays into the question of vaccination. There has been a recent focus on local, “ring” vaccination for cholera to try to limit spread from identified human cases. Our data, in keeping with earlier models published by our group, suggest that population-based vaccination may be more effective, recognizing the potential for broad distribution of the disease through environmental sources.

There has been a well-recognized link between water and cholera epidemics (28). However, recent genetic studies based on clinical isolates from global sources hypothesized that environmental reservoirs, while important in the persistence of the disease on the Indian subcontinent, have been playing a minimal role in cholera outbreaks in Africa and the Americas (5, 6, 29). Such studies lacked an analysis of aquatic environmental toxigenic \textit{V. cholerae} O1 isolates. Our findings show the critical role played by aquatic isolates in the transmission and evolution of cholera in Haiti. We propose a dynamic mixed-transmission model that assumes bacterial dissemination into an aquatic reservoir, evolution and adaptation to the new condition, and spillover to humans with subsequent transmission within human populations. Ultimate control of cholera is possible, but requires a recognition of the complexity of the transmission process and the need for long-term investments in cholera control at both the human and environmental levels.

Materials and Methods

**Sample Collection.** Between 2010 and 2017, we have isolated and characterized ∼800 toxigenic \textit{V. cholerae} O1 strains from cholera patients attended in diverse cholera treatment centers and clinics run by nongovernmental agencies and Haitian national clinics. For environmental monitoring of toxigenic \textit{V. cholerae} O1 strains, we collected surface water samples monthly from 17 sentinel sites in the Gressier/Leogane regions in Haiti effective May 2012. Environmental survey resulted in the isolation of 27 toxigenic \textit{V. cholerae} O1 strains between 2012 and 2015. Both clinical and environmental \textit{V. cholerae} O1 isolates were confirmed by standard microbiological, biochemical, serological, and genetic analysis as described previously (10, 30). Of 800 clinical isolates, 205 strains (116 from the Ouest Department) as well as all 27 environmental O1 strains were analyzed in this study. Information about the samples is available in SI Appendix, Tables S1 and S2.

Monthly case counts in the Ouest Department were obtained from the Pan American Health Organization (https://new.paho.org/hq/images/Atlas_IHR/CholeraHispaniola/atlas.html) (SI Appendix, Table S2), and mean monthly precipitation and average water surface temperature at 10 m above displacement height in Haiti were obtained from the NASA Goddard Distributed Active Archive Center (https://disc.gsfc.nasa.gov/). We tested if the variables used in a model to predict the case precipitations and temperatures were relevant to the outcome (cases) using a stepwise selection using the Akaike information criterion. The F test was used to compare this model with the one with the intercept only (no variables).

**Whole-Genome Mapping and HqSNP Calling.** Our dataset was composed of 116 toxigenic \textit{V. cholerae} strains: 27 environmental isolates were obtained in the Ouest Department, 23 isolates were sequenced in this study, and 4 were previously sequenced by our group (14); and 89 clinical isolates, of which one is the reference strain EL-1786 for the Haitian epidemic (31): 37 isolates were sequenced in this study, 32 isolates were previously sequenced by our group and collected between 2010 and 2012 (14), and 20 isolates also added to ensure correct calibration of the root of the tree were collected in the Artibonite and Ouest departments at the beginning of the epidemic between 2010 and 2011 (32). The newly sequenced toxigenic \textit{V. cholerae} strains were confirmed by serology and PCR (33). After subculture, genomic DNA extraction was performed using the Qiagen DNeasy Blood and Tissue Kit. Genomic DNA from all isolates was cultured and extracted from bacterial pellets. Sample library construction using the Nextera XT DNA Library Preparation Kit was performed (Illumina). Whole-genome sequencing on all isolates was executed on the Illumina MiSeq for 500 cycles (Illumina). Adapter and raw sequence reads were filtered by length and quality by using the program Trimmomatic (34). After quality filtering, Bowtie 2 (35) was used to map the sequence reads to the reference genome, \textit{V. cholerae} O1 strain 2010EL-1786 (GenBank accession nos. NC_016445.1 and NC_016446.1) (31). This reference sequence was isolated in Haiti and is generally used for
reference mapping of the samples that were collected in Haiti. After the reads were mapped to the reference genome, duplicate reads were marked and the reads were realigned using the program Picard (http://broadinstitute.github.io/picard). The reference-based mapping alignment was then verified and fixed accordingly, if needed. FreeBayes (36) was used to create a custom genome-wide SNP calling database (dbSNP) from all isolates in the dataset to perform base quality score recalibration (BQSR), outlined in GATK’s best practice guidelines for germline variation (https://gatk.broadinstitute.org/hc/en-us). When the variant call format (VCF) file was obtained, hard filtering of called SNPs was performed and the subsequent VCF file was used as a dbSNP for BQSR. When the BQSR step was completed, the alignment files were recalibrated and FreeBayes (36) was then used again to call variants on the recalibrated alignment files. Afterward, the newly created VCF file was filtered only for SNPs. After filtering, the VCF file was normalized using BCFtools (http://www.htslib.org/doc/bcftools.html). Normalization simplifies the represented variants in the VCF file by showing as few bases as possible at a particular SNP site in the genome. SNPs were then filtered by depth of coverage, quality, and genotype likelihood, as described in Azarian et al. (14). A SNP alignment in FASTA format was extracted from the VCF file from a custom python script. The SNP alignment was then filtered by site, leaving only sites with greater than 75% SNPs at that particular site, making a high-quality SNP alignment (SI Appendix, Table S3). The hqSNP alignment was annotated using the program Snpeff (37). HqSNPs found in protein-coding regions of the V. cholerae O1 genome that were identified by annotation were used to produce a codon alignment that was extracted in-house scripts (available upon request). All bioinformatic analysis was performed using this optimized pipeline implemented on the University of Florida’s HiPerGator cluster (https://www.vcr.ufl.edu/services/hipergator).

**Fig. 4.** Mixed-transmission model of cholera and vaccination prediction. (A) \( N_e \) (effective population size) observed from data (gray) and in simulation with environmental replication (green) and \( N_e \) in simulation without environmental replication (violet). (B) Distinct scenarios of vaccination and control for the model simulation with environmental replication, in particular vaccination coverages of 64% (rate of 0.01 d\(^{-1}\)), 88% (rate 0.04 d\(^{-1}\)), and 64% with a 10% increase in environmental decay rate. Here we define vaccination coverage as the percent reduction in susceptible individuals (upon reaching an approximate steady state after 2 to 3 mo of vaccination). As opposed to Kirpich et al. (27) and simulations without environmental replication (SI Appendix, Fig. S10) in which 64% vaccination coverage eradicates the pathogen within a year, here either more vaccination (88% coverage) or increased environmental decay (10% increase) is needed to control cholera in the presence of environmental replication.

Phylogenetic Quality Assessments and Maximum-Likelihood Analysis. All datasets used in this study passed phylogenetic quality checks as follows: in order to evaluate the presence of sufficient phylogenetic signal to resolve the phylogenetic relationship among V. cholerae O1 isolates, we performed likelihood mapping analysis with IQ-TREE (http://www.iqtree.org/) that calculates and maps the likelihood of all possible sequence quartets using the best-fitting nucleotide substitution model (38). Absence of substitution saturation was assessed by plotting pairwise transition/transversion vs. genetic distance with DAMBE6 (http://dambe.bio.uottawa.ca/DAMBE/) (39) (SI Appendix, Fig. S2). For each dataset, the presence of temporal signal was assessed with TempEst version 1.5 (http://tree.bio.ed.ac.uk/software/tempest/) (40) on ML phylogenies inferred by IQ-TREE (41, 42) (http://iqtree.cibiv.univie.ac.at) using the best-fitting nucleotide substitution model according to the Bayesian information criterion (41, 42) and ultrafast bootstrap approximation (1,000 replicates) to assess robustness of the phylogeny internal branches (43) (SI Appendix, Fig. S2). In order to test whether or not the spread of sampling times was sufficient to allow the substitution rate to be estimated accurately, that is, the presence of temporal structure within each of the time-structured datasets, we also performed the date randomization test on all phylogeographic datasets using TipDatingBeast implemented in the R package (44) (SI Appendix, Fig. S2). Ancestral-state reconstruction using maximum likelihood inferred with PastML (45) from a rooted maximum likelihood phylogenetic tree inferred with IQ-TREE, and the tree was compressed vertically and horizontally: clades have been vertically (clade circles contain the number of tips of the initial tree contained in them) and horizontally (the branch size corresponds to the number of times its subtree is found in the initial tree) merged to cluster independent events of the same kind.

**Bayesian Coalescent Inference: Discrete Phylogeographic Reconstruction, Markov Jumps, and Markov Rewards.** To test our hypothesis of long-term aquatic reservoirs of toxigenic V. cholerae O1 being established in Haiti, we used the Bayesian phylogeographic (46) coalescent-based method (15) implemented in
the BEAST version 1.8.4 (47) software package. The reconstruction of V. cholerae spatiotemporal spread in different environments through Bayesian phylogenetics requires calibration of a molecular clock. Evolutionary rates were estimated implementing the HKY nucleotide substitution model (48) with empirical base frequencies, gamma distribution of site-specific rate heterogeneity, and ascertainment bias correction (49, 50), testing a constant demographic prior against nonparametric demographic models, Gaussian Markov random-field Skyride (51), and Bayesian Skyline (52) to rule out spurious changes in effective population size inferred by a nonparametric model that is not fit to empirical sampling time (53). Additionally, for each demographic model, we compared strict and relaxed uncorrelated (lognormal distribution among branches) molecular clocks (18, 54). The best molecular clock and demographic model were chosen by estimating the marginal likelihood of each model, with path sampling and stepping-stone methods, followed by Bayes factor comparisons (47, 55) (SI Appendix, Tables S1 and S3). Each chain Monte Carlo samplers were run for a number of generations (50 to 200 million) sufficient to achieve proper mixing of the Markov chain, which was evaluated by calculating the effective sampling size (ESS) of each parameter estimate under a given model. ESS values >200 for all parameter estimates were considered as evidence of proper mixing. The origin of each isolate, environmental or clinical, was used as a trait for “location” in order to reconstruct the evolution of V. cholerae O1 and bacterial flow (migration) events between environmental and/or clinical sources. Transitions between discrete states (environmental and clinical) were estimated using the continuous-time Markov chain model using the asymmetric migration model with Bayesian stochastic search variable selection (46). In our reconstruction of ancestral states, we assume migration occurs at the tree nodes. Bayesian phylogenetic reconstructions are sensitive to disproportional sampling across subpopulations. We tested for sampling bias using two approaches: 1) using the less sensitive to sampling biases Bayesian structured coalescent approximation (56) implemented in BEAST2 version 2.5.2 for the Oustet dataset (SI Appendix, Fig. S5); and 2) using 10 resampled datasets on clinical (187) and environmental strains (27) isolated in 2012 to 2015, since no environmental strains were collected in 2016 to 2017, and all clinical strains sampled during these 2 yr were monophyletic (i.e., representing a new epidemic wave) not connected to any previously sampled environmental strain. Each dataset was generated by randomly subsampling clinical strains to match the same number of sequences available for environmental isolates collected between 2012 and 2015, and by adding the isolates collected at the beginning of the epidemic (2010) to allow correct rooting of the tree (31). The analyses described below were carried out in parallel on each of the resampled datasets and are shown in SI Appendix, Fig. S4.

The maximum clade credibility trees were obtained from the posterior distribution of trees with TreeAnnotator version 1.8.4 after 10% burn-in. The MCC phylogeny was manipulated in R using the package GGTREE (57) for publishing purposes.

Within the same phylogeographic inference, we also calculated the number of transitions between environmental and clinical states (Markov jump) (58) and the time between state changes (Markov reward) (58, 59). Markov jumps (or transitions) were plotted as the total number of state counts for the migration in and out of each location (environment, clinical) among all internal and external branches. Markov rewards can be estimated along the phylogenetic tree trunk to calculate the length of time that an ancestral population occupies a particular location (environmental or clinical) (60–62). We defined the backbone path, or trunk of the tree, following Lema et al. (63), as the internal subset of branches that connects the root node of a time-scaled phylogeny to the most recent common ancestor of the sequences sampled at the latest time point. In other words, the trunk represents the successful lineage that persists through time. Trunk proportions for time and ancestral locations were calculated after 10% burn-in, slicing time in 0.5-yr sections. If the tree trunk is occupied by multiple locations, cholera would exhibit source-sink population dynamics (60), and if the trunk is occupied by multiple locations, cholera would emerge from a population migrating between states (61, 62). Markov rewards were compared against a null distribution of 10 replicates obtained by randomization of the tip states performing binomial tests (64) to assess whether the mean of the posterior probability at every time point differed. Markov jumps and rewards were plotted using the R package ggplot2 (65).

Calculation of the Weighted Average of Synonymous Substitution Rates and Selection Analysis. In order to investigate replication rates among environmental versus clinical strains and to assess the selective pressure driving their evolution, we filtered the genome-wide hsSNP data to hsSNPs located in protein-coding regions. Using hsSNP in codon format, a subset of 200 Bayesian MCC genealogies was randomly obtained from the posterior distribution of trees for each subsampled data set and used for selection analysis. The weighted average of synonymous substitution rates (κS) and nonsynonymous substitution rates (κA) in the protein-coding regions of the V. cholerae O1 genome for all, internal and external, branches were obtained from a subset of 200 Bayesian MCC trees randomly obtained from the posterior distribution of trees, as described by Lema et al. (63). The ratio of nonsynonymous and synonymous substitution rates (dN/dS)—providing insight on whether or whether not positive selection is occurring mainly through random genetic drift (dN/dS < 1) or positive (dN/dS > 1) or negative (dN/dS < 1) selection—was obtained from a subset of 200 Bayesian MCC trees randomly obtained from the posterior distribution of trees, as described by Lema et al. (63). For the calculation of Tajima’s D (66) and P values, we used the tajima.test function in the R package pegas.

Statistical Analyses for dN and dS Substitution Accumulation. The null hypothesis of equality of two means, H0:p1=p2, against the alternative H1:p1≠p2, was tested through a Welch’s two-sample location test (67). It is an adaptation of the Student’s t-test and is applicable when the two samples have unequal variances. The resulting P value is used as evidence against the hypothesis that the two population means are equal. Testing the null hypothesis that the ratio of the two means is equal to 1 can be stated as H0:μ1=μ2, against the alternative H1:μ1≠μ2, which means that μ1=μ2, or μ1≠μ2, where μ1=μ2, and the null hypothesis on the ratio can be restated in terms of equality in means, that is, H0:μ1=μ2, against H1:μ1≠μ2. CIs are computed using the bias-corrected and accelerated bootstrap method. It adjusts for both bias and skewness in the bootstrap distribution (68).

SIRW Mathematical Model. See SI Appendix for a mathematical model for the eco-evolutionary dynamics of cholera.

Additional Data. R scripts and xml files are available at the GitHub page: https://github.com/salem1ab/environmental_cholera_haiti. The genomic sequences have been deposited with the National Center for Biotechnology Information (NCBI) in the Sequence Read Archive under BioProject ID code PRJNA510624. All other data are available in SI Appendix or upon request.

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