Evolutionary Sweeps of Subviral Parasites and Their Phage Host Bring Unique Parasite Variants and Disappearance of a Phage CRISPR-Cas System

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ABSTRACT Vibrio cholerae is a significant threat to global public health in part due to its propensity for large-scale evolutionary sweeps where lineages emerge and are replaced. These sweeps may originate from the Bay of Bengal, where bacteriophage predation and the evolution of antiphage counterdefenses is a recurring theme. The bacteriophage ICP1 is a key predator of epidemic V. cholerae and is notable for acquiring a CRISPR-Cas system to combat PLE, a defensive subviral parasite encoded by its V. cholerae host. Here, we describe the discovery of four previously unknown PLE variants through a retrospective analysis of >3,000 publicly available sequences as well as one additional variant (PLE10) from recent surveillance of cholera patients in Bangladesh. In recent sampling we also observed a lineage sweep of PLE-negative V. cholerae occurring within the patient population in under a year. This shift coincided with a loss of ICP1’s CRISPR-Cas system in favor of a previously prevalent PLE-targeting endonuclease called Odn. Interestingly, PLE10 was resistant to ICP1-encoded Odn, yet it was not found in any recent V. cholerae strains. We also identified isolates from within individual patient samples that revealed both mixed PLE(+)/PLE(−) V. cholerae populations and ICP1 strains possessing CRISPR-Cas or Odn with evidence of in situ recombination. These findings reinforce our understanding of the successive nature of V. cholerae evolution and suggest that ongoing surveillance of V. cholerae, ICP1, and PLE in Bangladesh is important for tracking genetic developments relevant to pandemic cholera that can occur over relatively short timescales.

IMPORTANCE With 1 to 4 million estimated cases annually, cholera is a disease of serious global concern in regions where access to safe drinking water is limited by inadequate infrastructure, inequity, or natural disaster. The Global Task Force on Cholera Control (GTFCC.org) considers outbreak surveillance to be a primary pillar in the strategy to reduce mortality from cholera worldwide. Therefore, developing a better understanding of temporal evolutionary changes in the causative agent of cholera, Vibrio cholerae, could help in those efforts. The significance of our research is in tracking the genomic shifts that distinguish V. cholerae outbreaks, with specific attention paid to current and historical trends in the arms race between V. cholerae and a cooccurring viral (bacteriophage) predator. Here, we discover additional diversity of a specific phage defense system in epidemic V. cholerae and document the loss of a phage-encoded CRISPR-Cas system, underscoring the dynamic nature of microbial populations across cholera outbreaks.

Editor Graham F. Hatfull, University of Pittsburgh
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The authors declare a conflict of interest. K.D.S. is a scientific adviser for Nextbiotics Inc. All other authors declare no competing interests.
The project described was supported by Grant Numbers R01AI127652 and R01AI153303 (K.D.S.) from the National Institute of Allergy and Infectious Diseases and its contents are solely the responsibility of the authors and do not necessarily represent the official views of the National Institute of Allergy and Infectious Diseases or NIH. K.D.S. is a Chan Zuckerberg Biohub Investigator and holds an Investigators in the Pathogenesis of Infectious Disease Award from the Burroughs Wellcome Fund. icddr,b gratefully acknowledges the following donors, which provide unrestricted support: Government of the People’s Republic of Bangladesh, Global Affairs Canada, Swedish International Development Cooperation Agency (SIDA), and the Department for International Development, UK Aid.
Received 13 October 2021
Accepted 26 January 2022
Published 15 February 2022

January/February 2022 Volume 13 Issue 1 e03088-21
Pandemics of the severe intestinal disease cholera have plagued humanity on a global scale for over two centuries. Caused by the bacterium *Vibrio cholerae*, the first pandemic is thought to have occurred in the early to mid-19th century, originating near the Bay of Bengal and sweeping across the Indian subcontinent, Asia, and parts of east Africa (1). There have been six subsequent pandemics, costing millions of lives and culminating in the current 7th cholera pandemic, which accounts for ~120,000 deaths per year globally (2) and has spread heavily throughout Africa and Asia. Devastating outbreaks have also occurred in the Americas, Europe, Oceania, and the Middle East. The genetic history of the 7th pandemic is marked by several temporally overlapping waves of distinct *V. cholerae* lineages, which themselves are comprised of numerous inferred transmission events (3). A hallmark of *V. cholerae*’s evolution is that not only do new lineages frequently arise but also that prior lineages simultaneously disappear. Pandemic *V. cholerae* is principally the O1 serogroup that is divided into classical and El Tor biotypes (4). A particularly dramatic example of lineage replacement in *V. cholerae* is that the first six pandemics are thought to have been entirely O1/classical strains, while the 7th pandemic is predominantly O1/EL Tor (5).

Isolates from within the waves of the current pandemic are highly homogenous, and each can be traced back to the Bay of Bengal as the original source (6). Cholera outbreaks are endemic to communities surrounding this area, where it is thought that *V. cholerae* is in continuous circulation between the human population and estuarian environments that act as the natural reservoir (7, 8). *V. cholerae* is thought to undergo the majority of its diversification and selection in the aquatic reservoir, giving rise to novel lineages that can infect local communities, spread to cities, and, from there, spread globally (9). Due to this single-point origin of epidemic *V. cholerae*, surveillance of cholera along the coast of the Bay of Bengal, specifically in Bangladesh, has previously proven very important in identifying not only novel bacterial genotypes (10) but also factors in *V. cholerae*’s ecosystem that influence the competition between lineages and help shape the selection of genotypes that cause outbreaks (11, 12). Of these factors, bacteriophage predation is likely of particular importance given that phages specific to *V. cholerae* are found in water sources in regions where the disease is endemic as well as in cholera patient stool (12–14). With phage predation threatening *V. cholerae* throughout its life cycle, the fitness of epidemic strains is likely contingent on the evolution of phage defense mechanisms.

Of the bacteriophages known to target *V. cholerae*, ICP1 stands out for its persistence, having first been detected in 1992 in India (15) and consistently recovered in stool samples from Bangladesh since then (14–17). However, *V. cholerae* can defend against ICP1 through the acquisition of parasitic antiphage islands called PLEs (phage-inducible chromosomal island-like elements) (18). During the course of ICP1 infection, the integrated PLE excises from the chromosome (19), replicates (20), hijacks ICP1 machinery to package its own genome (21), and can accelerate lysis of the host *V. cholerae* host (22), halting spread of ICP1 to neighboring cells. Five distinct yet highly similar PLEs have been detected in *V. cholerae* isolates going back to 1949, and over time each variant rose to prominence before being replaced by another (18). This succession is likely reflective of the fact that ICP1 has also evolved counterdefenses against PLEs, including an endonuclease called Odn (23) and, remarkably, a fully functional CRISPR-Cas system (10). However, the presence of these two anti-PLE elements also fluctuates over time, as they target different PLEs and do not cooccur in the same genome (23). The earliest ICP1 isolates from Bangladesh in 2001 possess Odn, while CRISPR-Cas was first was detected in an ICP1 isolate from 2003 (15). Between 2006 and 2017, all ICP1 isolates from Bangladesh were CRISPR(+) (16). However, contemporary coevolutionary trends of ICP1 and PLE are currently unknown.

A recent study from our lab of *V. cholerae* isolates from cholera patient stool samples in Bangladesh between 2016 and 2019 found evidence of temporal fluctuations in ICP1’s antagonism with other mobile genetic elements and SXT integrative and
conjugative elements (24). Considering these findings, we were interested in examining how the dynamics between PLE and ICP1 have progressed during this time period. Surprisingly, we found that after dominating the epidemic landscape for nearly a decade (16), PLE disappeared almost completely in 2018. This disappearance coincided with a switch from CRISPR-Cas back to odn among the temporally concurrent ICP1. A phylogenetic comparison of all 148 *V. cholerae* isolates during the surveillance period revealed an abrupt and nearly total shift from one *V. cholerae* lineage to a new lineage lacking PLE. Remarkably, this transition occurred over the course of only a few months. We briefly detected a novel PLE, which responds to ICP1 infection and is resistant to the anti-PLE nuclease Odn. An expanded analysis of >3,000 *V. cholerae* genomes revealed the existence of additional novel PLEs and provides a clearer picture of PLE succession both geographically and temporally. We also compared *V. cholerae* isolates from within individual patients, revealing heterogenous intrapatient PLE(+)PLE(−) populations. Similarly, heterogenous ICP1 phages possessing CRISPR-Cas or Odn were isolated from a single patient and exhibited evidence of *in situ* recombination, underscoring the potential for rapid phage evolution in the human gut.

**RESULTS**

Clinical surveillance of *Vibrio cholerae* and a predatory bacteriophage uncovers regressive shifts in the arms race. We analyzed 239 stool samples from patients in Bangladesh with suspected cholera infections, as determined by a rapid diagnostic test commonly referred to as a dipstick test. These samples were recently interrogated to study the temporal dynamics of SXT integrative and conjugative elements as well as the relevant mechanisms of phage counteradaptation (24). Samples were collected between November 2016 and September 2019, with 119 originating from the capital city Dhaka and 121 from a small city, Mathbaria, located on the Bay of Bengal coast (Fig. 1A; see also Table S1 in the supplemental material).

Interestingly, ICP1 phages, which previously encoded only CRISPR-Cas in Bangladeshi stool samples after 2006 (16, 17), transitioned back to encoding odn around May 2018. Bulk metagenomic genomic DNA was extracted from 197 of the stool samples and used as a template for PCR to detect the presence of ICP1 by screening for odn and CRISPR. Notably, all ICP1 isolates sequenced thus far encode either odn or CRISPR-Cas, suggesting this PCR screen is a sufficient indicator for the presence of ICP1 in patient samples. This screen revealed that 71 of these 197 samples were positive for ICP1, with 36 (50.7%) samples possessing CRISPR, 32 (45.1%) possessing odn, and three (4.2%) having a mix of both anti-PLE loci (Fig. 1B). A sampling period over ~11 months followed where no ICP1 was detected, and then in April-May of 2019, both loci were detected again. In the final 4 months of surveillance (June to September 2019), only odn was detected. To further investigate this transition from CRISPR dominance to apparent odn dominance, we examined whole-genome sequences of 44 individual ICP1 isolates from these stool samples (Table S1). These genomes corroborated the PCR results that CRISPR-Cas dominance declined while the odn locus began to dominate in mid-2018 (Fig. 1C).

Since both ICP1-encoded loci defend against *V. cholerae*-encoded PLEs (23), we examined whole-genome sequences of *V. cholerae* (*n* = 148) from 110 stool samples spanning this period. From the genomes we determined whether each isolate contained PLE and, if so, which of the five known PLEs was present (Fig. 1D). This analysis led to two unexpected results. First, PLE1, which was observed to be the dominant PLE in Bangladesh during previous surveillance (16), appears to have disappeared almost entirely in favor of PLE(−) strains. Interestingly, the disappearance of PLE1 roughly correlated with the period of ICP1’s transition from CRISPR-Cas to odn, with only a single isolate possessing PLE1 after February 2018. Second, we observed that four *V. cholerae* isolates from one stool sample collected in May 2017 possessed a novel PLE variant, which we have designated PLE10. This new PLE was only observed once and was not detected in any later isolates. From these data it is difficult to say whether PLE10 provided inadequate defense against ICP1 and was selected against in the population or if
increased surveillance and isolate sequencing would reveal additional instances of PLE10 in other samples.

**PLEs have evolved and diversified both globally and temporally.** The discovery of a novel PLE and the surprising observation that PLEs in Bangladeshi *V. cholerae* isolates have diminished in recent years prompted us to look more broadly, both geographically and temporally, at other available *V. cholerae* genomes. Previous surveillance studies tracked the occurrence of PLEs to a limited extent (~200 genomes) (16, 18); however, this was likely insufficient to fully elucidate the flux of PLEs over time and space. To build on these analyses, we constructed a database of 3,363 sequenced *V. cholerae* genomes, including our new surveillance isolates and raw reads or fully assembled genomes from public repositories. This collection spans over a century (1916 to 2019) and includes strains collected across 25 different countries (Data Set S2).

Using the previously discovered five PLEs as queries (18), we performed BLASTn searches against each genome and identified four additional PLEs that, along with PLE10, double the total number of known PLE variants from 5 to 10. The abundance of these 10 PLEs has shifted over time, with PLE5 being the only PLE detected before 1987, followed by the other PLEs, culminating in the more recent dominance of PLE1 (Fig. 2A). Previously, PLE5 had been observed as early as 1949, but here we detected PLE5 in an isolate from 1931, increasing the known time range of PLEs circulating in *V. cholerae*. Generally speaking, the previous pattern of temporal succession, where one PLE dominates for a time...
Evolutionary Sweeps Drive *V. cholerae* Lineage Shifts

FIG 2 Genetic, temporal, and spatial variability of PLEs in >3,000 *V. cholerae* genomes collected between 1916 and 2019. (A) Timeline of PLE-positive strains colored by PLE variant type from 3,363 *V. cholerae* whole-genome sequences. The lower histogram provides the total number of strains per year, while the upper histogram represents a stacked percentage of those that are positive for each PLE. Because of the small number of sequenced genomes from isolates prior to 1960, those genomes were collapsed into a single column. Metadata for all strains analyzed can be found in Data Set S2. (B) Overview of the geographical occurrences of PLE(+) isolates by country of isolation. Each individual isolate is indicated with a colored dot corresponding to the PLE type, as in panel A, and grouped with all others for a given country. The relatively large numbers of isolates from Bangladesh are represented in the inset. (C) Global alignment of all 10 PLEs using progressiveMauve. The order was determined by phylogenetic distance between PLEs (Fig. S1). The mauve-colored regions are conserved across all PLE sequences, while the light gray regions are unique to that specific variant. All other regions vary in conservation between two or more PLEs and were assigned random colors from the variable palette. Annotated genes are shown to scale as black arrows, and several genes with known functions are labeled (§, lidI).
before being supplanted by another PLE (16, 18), was observed in this larger collection of isolates, corroborating previous analyses. However, we observed more temporal overlap of PLEs than has previously been appreciated. Furthermore, we observed that while PLE5 had appeared to go extinct among clinical isolates in 1991, three additional PLE5-positive isolates were detected in 2016 to 2017 after a nearly 30-year absence, pointing to unsampled reservoirs in nature where genotypes may persist for some time. Some of the newly discovered PLE variants, however, only appeared for a short time, such as PLE6, which was detected once in Bangladesh in 1987, and PLE8 in Vietnam in 1995 and 2004. We also detected one possible phage satellite with partial genetic relationship to PLEs, as has been described in non-cholera vibrios recently (25); however, its sequence could not be fully resolved from available data. It is highly likely that we still have not captured the full extent of PLE diversity and variant flux over time, as *V. cholerae* isolate sampling was sporadic and limited until recent decades (with over half of all isolates collected in only the last 9 years). For example, in this current surveillance effort we observed PLE10 in only a single patient sample, which likely would not have been observed at all with slightly less sampling depth. Whether these short-lived PLEs (and divergent satellites) are somehow less adept at parasitizing ICP1, are selected against by specific anti-PLY counterdefenses, or are victims of simple stochastic fluctuations is unknown.

Additionally, transmission bottlenecks that likely occur in the course of regional outbreaks may constrain PLEs geographically (Fig. 2B), such as in the two most recent outbreaks in Haiti (26) and Yemen (27), where no PLEs were detected. The most notable exception is Bangladesh, where 7 of the 10 PLE variants have been detected. While this may be partially due to the sampling focus on this region, as a significant fraction (~20%) of the *V. cholerae* genomes analyzed were from there, it also may be due to its juxtaposition with the Bay of Bengal, where the majority of *V. cholerae*’s diversification and evolution are predicted to occur.

Having found additional PLEs, we then performed a whole-genome alignment of all 10 PLE variants (Fig. 2C), which was used to generate a phylogenetic tree (Fig. S1). All PLEs are broadly syntenic and share numerous regions of nucleotide conservation. Annotation and sequence comparison of open reading frames confirmed that the newly detected PLEs 6 to 10 possess genes key to PLE function that have been characterized in the previously studied PLEs 1 to 5. These include *nixI*, which encodes a nickase that interferes with ICP1’s ability to replicate (25), *lidI*, which can accelerate lysis of the *V. cholerae* host cell (22), and *capR*, which acts to repress ICP1’s capsid morphogenesis operon (21). The conservation of these genes indicates that the newly identified PLEs play roles similar to those of PLEs 1 to 5 as parasites of ICP1. Two additional genes indicate that these PLEs are also still functional as mobilizable genetic elements: an integrase that catalyzes PLE excision from the host genome and integration following transduction, and a replication initiation factor that controls PLE replication.

PLEs encode one of two integrases (Fig. S2), the PLE1-type (found in PLEs 1, 3, 4, 5, 6, and 10), which directs integration into a *Vibrio cholerae* repeat (VCR) sequence in the superintegron and recognizes ICP1-encoded PexA as a recombination directionality factor to catalyze PLE excision (19), and the PLE2-type (found in PLEs 2, 7, 8, and 9), which directs PLE integration into the M48 family metallopeptidase gene (*vca0581*) and does not interact with PexA. VCRs are ~124-bp repeat elements that are interspersed between gene cassettes within the superintegron (28). VCRs are extremely conserved, which could allow PLEs with the PLE1-type integrase to integrate into any single VCR following horizontal transfer, resulting in isogenic PLEs flanked by different genes in the superintegron. Previous analyses showed that PLE1 integrates into various distinct VCRs under laboratory conditions, but such variability has not been observed among PLE(+) *V. cholerae* isolates from clinical specimens, suggesting that PLE transmission is largely vertical in nature (16, 18). To determine if this finding held true, we extracted the genomic regions flanking PLE for every PLE-positive *V. cholerae* isolate in our data set. While no flanking differences were observed among the PLEs with PLE2-type integrases, we identified 14 unique VCR integration sites for the PLE1-type PLEs.
integrase-positive *V. cholerae* isolates in our database (Fig. S3 and Data Set S3). While most PLE variants appeared to have a single integration site, either through preferential horizontal integration or vertical transmission, we did detect multiple sites for PLE1, PLE4, and PLE5, indicating horizontal acquisition events in nature.

The PLE-encoded replication initiation protein RepA has a conserved C terminus across all PLEs but varies at the N terminus, which directs DNA binding to the PLE origin of replication (ori) (20). Previous work found that RepA’s N terminus for PLEs 1 to 5 fell into two groups, one with PLEs 1, 4, and 5 and the second with PLEs 2 and 3. Diversification of the PLE replication module (comprised of a compatible RepA and ori) is hypothesized to be driven by ICP1’s nuclease Odn, which mimics the PLE1,4,5-type RepA protein to bind to the cognate ori and then cleave it. For PLE to escape Odn by modifying its ori, it must possess a compatible RepA N terminus as well as the conserved C terminus, which is hypothesized to facilitate recruitment of the replisome machinery from ICP1 (20). Including the five new PLEs described here, phylogenetic analysis revealed the existence of a novel third RepA N-terminal variant found in PLEs 6, 8, 9, and 10 (Fig. S4). In support of the designation of a third type of PLE replication module, previously validated ori sequences in PLE1-5 were not observed in PLEs with the PLE10-type RepA protein, indicating that these PLEs harbor a compatible unique ori sequence that would be resistant to Odn-mediated cleavage.

**PLE10 responds to ICP1 infection and is resistant to Odn.** All previously studied PLEs (PLEs 1 to 5) are triggered by ICP1 infection to excise, replicate, and package themselves into transducing particles using ICP1’s machinery (Fig. 3A) (18). Based on genetic similarity to PLEs 1 to 5, we hypothesize that the five new PLEs described here respond similarly to ICP1 infection. Unfortunately, of the new PLE variants we only had access to *V. cholerae* strains possessing PLE10 and therefore experimentally assessed how this variant responds to ICP1 infection.

To determine if this novel PLE10 variant indeed functions in a way similar to that of known PLEs, we performed several assays in tandem with PLE1. First, a circularization PCR showed that PLE10 excises and circularizes during ICP1 infection (Fig. 3B), as was expected due to the similarity of the PLE1 and PLE10 integrases (Fig. S2). Second, we observed that PLE10 replicates in response to ICP1 infection (Fig. 3C). This confirms that the divergent PLE10 replication module is functional and likely exploits ICP1 machinery to promote PLE replication. Interestingly, although PLEs 1 to 5 do not replicate during infection by ICP1 mutants lacking an accessory SF1B-type helicase (*helA* or *helB*) (29), PLE10 does replicate to low levels in the absence of this ICP1-encoded helicase (Fig. 3C), suggesting that the divergent replication module allows for additional flexibility in PLE’s reliance on ICP1’s replication machinery. Third, we observed that PLE10, which possesses a *lidI* homologue (22), also exhibited an accelerated lysis phenotype, like PLE1, following ICP1 infection (Fig. 3D). Fourth, PLE10 mobilized and transduced its genome to naive *V. cholerae* hosts during infection at rates similar to those of PLE1 (Fig. 3E). These results demonstrate that PLE10 exhibits the anticipated response to ICP1 infection, namely, excision, replication, and transduction.

As ICP1 is known to possess two anti-PLE counterdefense loci (*odn* and CRISPR-Cas), we tested the ability of ICP1 with either loci to plaque on *V. cholerae* having either PLE1 or PLE10. PLE1 is susceptible to ICP1 possessing CRISPR-Cas (those with a spacer specific to PLE1) (10), and one hypothesis is that this new PLE10 variant has an advantage over PLE1 through the acquisition of an anti-CRISPR, although none have been found previously in PLEs. However, we found that the CRISPR-Cas(+) phage, isolated from the same stool sample as the PLE10 *V. cholerae* isolates and naturally possessing spacers specific to both PLE1 and PLE10, could overcome PLE10 during infection and that this activity was dependent on ICP1’s CRISPR-Cas system (Fig. 3F). This was an identical result to that for the PLE1 control, which was overcome by the same CRISPR-Cas(+) phage. The inability to block plaque formation by CRISPR-Cas(+) phage indicates that PLE10 has no anti-CRISPR protein that might provide an evolutionary advantage over PLE1.
In contrast, an ICP1 phage with odn was unable to form plaques on a PLE10(+) host but was able to form plaques on an otherwise isogenic host harboring PLE1 (Fig. 3F). Purified Odn also did not exhibit nucleolytic activity against a probe amplified from the ORF-less region (expected to harbor the ori) of PLE10 compared to a probe from PLE1 that was cleaved as expected (Fig. S5). Collectively these data are consistent with
the specificity of the origin-targeting behavior of Odn (23) and the divergent PLE10 replication module conferring protection from Odn-mediated cleavage. The Odn(+)
phage, tested for its capacity to form plaques on PLE10, was from a more recent 2019 stool sample, a time period in which Odn-positive ICP1 phages were dominant in our surveillance (Fig. 1C). However, despite the observation that PLE10 is resistant to this Odn(+)
phage, it did not have an apparent selective advantage over PLE1 during this surveillance period (Fig. 1D).

Colony and plaque screens reveal intrapatient heterogeneity. Preliminary PCR screening of metagenomic DNA from stool samples revealed some samples with a positive signal for PLE but from which we later recovered V. cholerae isolates that were PLE(-). This finding prompted us to interrogate several samples further to better understand the ratios of PLE(+) to PLE(-) strains within intrapatient populations. We selected three samples from May 2017 (D19, D28, and D35) and one from September 2017 (D55), as both time points also possessed a mix of PLE(+) and PLE(-) isolates between patient samples (Fig. 1D). We performed PCR on a large number of colonies (n = 61 to 130) picked from each of these samples and observed that all samples were indeed heterogenous for the presence of PLE [ranging from 6 to 97% PLE(+)] (Fig. 4A). We additionally chose one sample from May 2017 from which all (n = 4) sequenced isolates were PLE(-) to see if PLE heterogeneity could be caused by stochastic loss under laboratory conditions. PCR of 218 colonies revealed all to be PLE(+), indicating that PLE loss was highly unlikely to occur as a result of the outgrowth and isolation procedures used. These results reinforce previous discoveries that V. cholerae can diversify to some extent during human infection (30), although whether this finding indicates a loss or gain of PLE is unclear.

Similarly, we performed PCR on a large number of ICP1 plaques from sample M76 (May 2018), in which the initial screen of metagenomic DNA extracted from stool revealed a mixed signal of both ICP1-encoded odn and CRISPR-Cas anti-PLE loci. This was also the first instance of odn reappearance (Fig. 1B) in this surveillance period. All previously discovered ICP1 isolates have either odn or CRISPR-Cas but never both (15, 17, 23), and these counter-defense mechanisms have always been in the same location within the genome. It is an intriguing possibility that some rare isolates could carry both odn and CRISPR-Cas, but PCR results from purified plaques (n = 56) from this sample showed that members of this intrapatient ICP1 population indeed only had CRISPR-Cas or odn (Fig. 4B).

In addition to the 56 screened plaques, we also sequenced the genomes of six isolates of ICP1 from this patient sample, two odn(+) isolates and four CRISPR(+) isolates. Whole-
genome sequencing revealed that both odn(+) phage had a canonical downstream region typically associated with odn, and three of the CRISPR(+) phage had the expected distinct conserved set of associated genes. However, one isolate (2018_Mat_164) harboring the CRISPR-Cas anti-PLE locus possessed the downstream region typically associated with odn (Fig. 4C). This appears to be an example of within-patient recombination between two ICP1 genotypes and is strong additional evidence that coinfection is a driver of ICP1 evolution, as has been demonstrated previously under laboratory conditions (22).

To the best of our knowledge, this is the first attempt to document intrapatient diversity of vibriophages and, as such, the rates of in situ recombination are unknown. Nonetheless, these data highlight that ICP1 evolution occurs in the human gut, giving rise to distinct ICP1 genotypes with distinct host ranges (23) that could differentially impact the fitness of cocirculating V. cholerae.

V. cholerae isolates from 2016 to 2019 in Bangladesh sort into two distinct phylogenetic lineages. The transmission and evolution of V. cholerae is characterized by successive waves of clonally related strains that are replaced over time (3), and the phylogenetic relationships between these waves generally have been determined by differences in single point mutations. We observed that PLE disappeared from V. cholerae isolates during the course of our surveillance (Fig. 1D); however, we did not know if this is the result of PLE being lost from a clonal population or if the PLE(+) lineage was replaced by a distinct PLE(−) lineage. To address this question, we identified 532 sites with single nucleotide variations (SNVs) (Data Set S4) common across all 148 isolates and used this information to construct a core genome phylogenetic tree (Fig. 5).

It was immediately clear that the V. cholerae isolates cluster into two highly distinct phylogenetic groups that largely coincide with the presence of PLE, suggesting that we captured a replacement event where the PLE(−) lineage emerged to dominate as opposed to witnessing the loss of this mobile genetic element (MGE) in an otherwise persistent lineage. This was further reinforced by a previous observation in this surveillance data set that another MGE that plays a role in phage defense, integrative and conjugative elements (ICEs) of the SXT/R391 family, exhibited a transition from a mix of VchInd6 and SXT(−) to VchInd5 over the same time period (24). Additionally, sequences of the wbeT gene, which is part of the biosynthetic pathway for the O1 antigen, also revealed a switch from the Inaba (nonfunctional wbeT) to Ogawa (functional wbeT) serotype that perfectly tracked with the lineage replacement. In the earlier Inaba lineage, wbeT was disrupted by a transposon with an identical sequence and integration site across all isolates. Remarkably, this replacement sweep appears to have occurred over months instead of years, with only a few isolates from the earlier PLE(+/VchInd6(+))/Inaba lineage persisting into 2018 and only a single one in 2019. This indicates that the later PLE(−)/VchInd5 (+)/Ogawa lineage has some competitive advantage over the earlier lineage driving its sudden and dominating emergence. We also saw no patterns separating the sampling locations of Dhaka and Mathbaria, indicating that the replacement sweep occurred rapidly geographically as well as temporally.

Finally, in addition to the samples examined by colony PCR (Fig. 4A), multiple isolates were sequenced from several other samples when recoverability permitted. While significantly more isolate sequencing would be required to draw meaningful conclusions about the full extent of intrapatient diversity, we did observe both SNV and MGE variability between isolates in some patient samples (Data Sets S1 and S4). However, in no sample did we detect isolates from both lineages among the intrapatient isolates, perhaps suggesting that although rapid, lineage sweeps are unlikely to occur in the context of a single patient’s infection.

DISCUSSION

The trajectories of global cholera outbreaks are generally defined by successive waves of V. cholerae in which a new lineage replaces the previous lineage (3). While it is unclear exactly what allows successive lineages to outcompete their predecessors, it is broadly theorized that each new lineage in the current 7th pandemic originated from the areas
Phylogeny of *V. cholerae* isolates during the surveillance period in Bangladesh reveals the replacement of a previously prevalent PLE(+) lineage with a PLE(−) lineage in 2018 to 2019. (Continued on next page)

**FIG 5** Phylogeny of *V. cholerae* isolates during the surveillance period in Bangladesh reveals the replacement of a previously prevalent PLE(+) lineage with a PLE(−) lineage in 2018 to 2019. A
surrounding the Bay of Bengal (6). Therefore, developing a better understanding of the
genetic variability of *V. cholerae* in this region is likely key to unlocking the mystery sur-
rrounding the cycle of global selective sweeps. Furthermore, predatory bacteriophages
like ICP1 and antiphage defenses such as PLE may play an important role in determining
which lineages become pathogenically relevant in human populations. In this study, we
describe findings from continued surveillance of *V. cholerae* in Bangladesh and build
upon previous work to further reinforce the importance of monitoring these microbial
populations in communities proximal to the Bay of Bengal.

The antagonism between ICP1 and PLE results in an ongoing arms race in which ICP1
evolves means to target PLE and inhibit PLE’s anti-ICP1 activity. The earliest known ICP1
isolates have the Odn nuclease, which cleaves PLE’s origin of replication (23), while PLEs
likely diversified to escape Odn targeting. It was assumed, however, that the extreme
flexibility and rapid adaptation provided by the later acquisition of a novel CRISPR-Cas system
had given ICP1 the upper hand in this battle (16), or at least that CRISPR-Cas would be so
beneficial that it would remain fixed in the ICP1 population. Remarkably, though, we
observed that despite the assumed superiority of a CRISPR-Cas counterdefense mechani-
ism, a reversion occurred in the ICP1 population back to carrying the *odn* locus instead
of CRISPR-Cas. During this same time frame before reversion to *odn*, we also found that
nearly all *V. cholerae* isolates lacked PLE. Whether or not these two nearly simultaneous
changes are functionally connected is unclear. Perhaps the much larger CRISPR-Cas locus
is too burdensome for the phage to encode when PLE no longer threatens its viability,
maybe *odn* is more effective at fully restricting PLE activity for the PLEs that it can target,
or possibly both (23). These hypotheses suggest that PLE was lost from the *V. cholerae*
population first, and in the absence of that selective pressure, ICP1 quickly reverted to a
more favorable genotype.

A similar reversion event was previously observed in these same surveillance samples
in which the promoter for *orbA* that inhibits antiphage components of the SXT ICE
*VchInd5* was missing from ICP1 isolates while *VchInd5* was absent from the *V. cholerae*
population and a different SXT ICE dominated. However, when *VchInd5* became domi-
nant in the population around mid-2018 (Fig. 5), phage isolates with *orbA* and its func-
tional wild-type promoter also reemerged as the dominant ICP1 genotype (24). It is possi-
bile that ICP1’s robust accessory genome allows it to keep pace with emergent *V. cholerae*
lineages, and reemergence of historical genotypes suggests unsampled reservoirs of ICP1
diversity, much like we expect for *V. cholerae*. If future lineage shifts cause PLE to re-
merge in the *V. cholerae* population, it may also mean that CRISPR-Cas, or perhaps yet
another anti-PLE mechanism, will again supplant Odn in the ICP1 population. However,
this hypothesis is potentially weakened by the discovery of a new PLE in one patient stool
sample approximately a year before the shift in ICP1’s anti-PLE loci. The new PLE (PLE10)
was shown to be functional as a phage parasite in response to ICP1 infection but was
also resistant to Odn targeting because it harbors a novel origin of replication. In this
case, why did PLE10 not proliferate in the *V. cholerae* population after ICP1 had switched
to *odn*? It is impossible to say for certain, but it may have some unknown competitive dis-
advantage or was simply unable to spread through the population while CRISPR-Cas with
PLE10 spacers was still cooccurring. It is further possible that the pressures inherent in the
dynamics of ICP1-PLE interactions are not responsible for driving *V. cholerae* lineage selec-
tion and PLE10 was simply not present in the lineage that swept to dominance in 2018.

In further investigating the genomics of this lineage switch among the *V. cholerae sur-
vellance isolates, we observed two very distinct clades based on SNVs. While the SNV
analysis did not reveal any candidate mutations that would indicate an obvious competi-

![FIG 5 Legend (Continued)](cladogram of 148 *V. cholerae* isolates based on 532 variable nucleotide genomic positions (Data Set 54). Columns next to the tree contain metadata for (left to right) the PLE variant present, the SXT ICE present, city of isolation, year isolated, and serotype (based on whole-genome sequencing and analysis of *wbeT*). Column coloring and order is shown in the legend from top to bottom. Strains lacking MGEs (PLE or SXT ICE) are indicated in light gray.)
Evolutionary Sweeps Drive V. cholerae Lineage Shifts

tive advantage of the new lineage, a sizable number of the nonsynonymous variations were perfectly separated between the two lineages (see Data Set S4 in the supplemental material). This indicates that the lineage shift reflects an emergence of a distinct V. cholerae variant that already had these mutations in its genome. Additionally, the serotype switch from Inaba to Ogawa also tracked precisely with the two lineages. The tendency of V. cholerae to change serotype between Inaba and Ogawa, or vice versa, has been well documented. This is in large part due to the ease with which serotype agglutination tests can be performed and has revealed these shifts to be very common (31). This phenomenon has been documented globally (32, 33) as well as in Bangladesh for almost five decades (34), including a time range that overlaps in part with our surveillance period and agrees with the observation made here (35). Despite this long history, it is still unknown what drives the Inaba/Ogawa serotype switching in V. cholerae. As the function of the wbeT gene is to modify the O1 antigen, it has been hypothesized that phage predation is a selective pressure (36); however, ICP1 (which uses the O1 antigen as its receptor) is insensitive to this modification (14), and there is no evidence to indicate that the other two known vibriophages frequently isolated from patient stool samples (ICP2 and ICP3) are impacted by such a modification. An alternative hypothesis is that O1 antibodies created as an immune response to V. cholerae in patient populations could select against whichever serogroup is currently circulating. While this has not been conclusively tested, a recent study found that patients challenged with an Inaba strain developed antibodies that cross-reacted effectively with the Ogawa serotype (37), suggesting a limited selective advantage to serotype switching in evading human immunity.

We did find some evidence of possible diversification within patients by detecting mixed populations of PLE(+) and PLE(−) colonies from several patient samples as well as in sequenced isolates from one patient that varied by several SNVs. These observations are reinforced by previous discoveries that found genetic variation of V. cholerae during human infection (30) and even the emergence of hypermutator phenotypes (38). Whether this indicates an in situ loss, possibly due to an incurred fitness cost of harboring the MGE, gain of PLE through lateral transfer, or coinfection with distinct strains is unknown. We also found evidence of bacteriophage heterogeneity leading to recombination within a single patient. We are not aware of this being observed in nature before, and there is experimental evidence of ICP1 recombination under laboratory conditions (22). The evolutionary variation in the V. cholerae/ICP1/PLE system is likely apparent, and perhaps more so, in the aquatic reservoir. A deeper investigation of individual patient samples as well as longitudinal surveillance and environmental samples will be important to begin to fully understand the dynamics at play in driving V. cholerae evolution and lineage selection.

**MATERIALS AND METHODS**

**Bacterial growth conditions.** The bacterial strains and plasmids used in this study are listed in Table S1 in the supplemental material. All bacterial strains were grown at 37°C in LB (Fisher) with aeration or on LB agar plates. Antibiotics were used when appropriate (100 μg/mL streptomycin, 75 μg/mL kanamycin, 2.5 μg/mL chloramphenicol for V. cholerae and 25 μg/mL for E. coli). Ectopic expression constructs in V. cholerae were induced 20 min prior to ICP1 infection with 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) and 1.5 mM theophylline.

**Phage growth conditions.** The phages in this study are listed in Table S1. Bacteriophages were propagated on V. cholerae, harvested via polyethylene glycol precipitation (39) or medium exchange on Millipore's Amicon Ultra centrifugal filters (40), and quantified via the soft-agar overlay method (39).

**Isolation of bacteria, phage, and total DNA from stool.** The collection of stool samples and isolation of V. cholerae and phages interrogated in this study was recently described (24). Briefly, deidentified rice water stool (RWS) from patients presenting with diarrheal disease were tested for the presence of V. cholerae via a crystal V. cholerae rapid dipstick test (RDT; Span Diagnostics, Surat, India) at icddr,b Dhaka Hospital and Government Health Complex of Mathbaria, Pirojpur, Bangladesh. RDT-positive stool samples were mixed with glycerol and frozen for transport to the University of California, Berkeley. V. cholerae was isolated from RDT-positive stool samples onsite in Bangladesh following enrichment in alkaline peptone water (APW) and plating on taurocholate tellurite gelatin agar (TTGA) (Difco). Additional isolates of V. cholerae from these RWS samples were obtained following APW outgrowth and culturing thioulate-citrate-bile salts-sulfate agar (Fish) and Vibrio ChromoSelect agar (Sigma) at the University of California, Berkeley. To isolate phages, dilutions of RWS and APW outgrowths were mixed with log-phase V. cholerae and plated in 0.3 to 0.5% LB top agar. Total DNA was extracted from stool samples using the DNeasy PowerSoil kit (Qiagen) with
Whole-genome sequencing. Whole-genome sequencing of V. cholerae and ICP1 isolates during the surveillance period was performed in our previous study (24). Briefly, DNA from V. cholerae and phages was extracted using commercially available kits (Qiagen DNeasy blood and tissue or monarch genomic DNA purification kit from New England Biolabs). For preparation of phage DNA, phage stocks were treated with DNase at 37°C for 30 min before heat inactivation prior to DNA extraction by following the manufacturer’s instructions. Libraries for Illumina sequencing were prepared with New England Biolabs’ Ultra II DNA or Ultra II FS DNA prep kits or performed by the Microbial Genome Sequencing Center. Sequencing (150- by 150-bp paired end) was performed by the QB3 Genomics Core at the University of California, Berkeley, or by the Microbial Genome Sequencing Center.

CRISPR/Odn and PLE PCR screening. To detect ICP1-encoded anti-PLE loci from patient stool, total DNA from stool was used as a template for PCR using the primers listed in Table S1. Colony PCR using the primers listed in Table S1 was performed on individual V. cholerae colonies to screen for the presence of PLE1 among isolates recovered from the same patient sample.

Genome database and PLE discovery. A database of publicly available genomes was generated by downloading all deduplicated assemblies from the NCBI V. cholerae genome directory (https://www.ncbi.nlm.nih.gov/genome/browse/#/prokaryotes/505/) and by a literature search for raw sequence reads available from the NCBI Sequence Read Archive (https://www.ncbi.nlm.nih.gov/sra). These raw reads were subsequently downloaded in fastq format and assembled with spades v3.14.0 using default settings. This data set is similar to but expanded from that of a previous work (24). Each of the five previously known PLEs were used as BLASTn queries against the completed V. cholerae database, and the resulting nonoverlapping hit lengths (E value of <0.001) were summed for each PLE. Summed hit lengths that exceeded 10% of that PLE’s total nucleotide length were considered putative PLEs and were manually curated with CLC Main Workbench 7 to ensure similar length, synteny, and sequence similarity to known PLEs. This BLASTn approach was also used to identify genomes with copies of the known PLEs.

PLE alignment and phylogeny. PLE fasta files were aligned using progressiveMauve (v. 2015-02-25) using default parameters as well as the assume colinear genomes setting. Region color combinations were randomly chosen from the Viridis color palette for each combination of PLEs, and the phylogenetic tree was generated from the guide tree file.

PLE integration sites. The first 100 bp of the genomic regions immediately flanking every PLE-positive V. cholerae isolate in the V. cholerae genome database were extracted and grouped by those that matched exactly. Only PLEs with a uniquely matched up to left and right regions flanking the integration site were considered (i.e., no left flank sequence occurred with more than one right flank sequence). This ensured that integration site differences were not due to genomic variability or shuffling within the superintegron of each PLE-positive V. cholerae isolate. Those PLEs that satisfied these requirements were grouped by identical flank pairs (i.e., integration sites) and plotted.

A cladogram of 148 Vibrio cholerae isolates was constructed based on 532 single nucleotide variations (SNVs). These variable locations were determined by aligning the assembled sequence of each isolate to the reference strain N16961 using MUMmer (v. 3.0). Only nucleotide locations that were present in all 148 isolates as well as the reference were considered for analysis.

333 μL of stool by following a modified extraction protocol. Briefly, 200 μL of bead solution was removed and replaced with 200 μL of phenol-Chloroform-isoamyl alcohol (pH 7 to 8; Sigma) before the stool was added. High liquid content of stool resulted in increased sample volume after bead beating, resulting in our scaling up of reagent volumes, after which the manufacturer’s instructions were followed.

Phage spot assays. Mid-log-phase V. cholerae was added to 0.5% molten LB agar, poured on a solid agar plate, and allowed to solidify. Tenfold dilutions of phage were overlaid in 3-μL spots. After spots were dry, plates were incubated at 37°C for ~6 h prior to imaging. Images are representative of three independent experiments.

V. cholerae phylogeny. A cladogram of 148 Vibrio cholerae isolates was constructed based on 532 single nucleotide variations (SNVs). These variable locations were determined by aligning the assembled sequence of each isolate to the reference strain N16961 using MUMmer (v. 3.0). Only nucleotide locations that were present in all 148 isolates as well as the reference were considered for analysis. The
Supplemental material is available online only.

**DATA SET S1**, XLSX file, 0.02 MB.

**DATA SET S2**, XLSX file, 0.1 MB.

**DATA SET S3**, XLSX file, 0.02 MB.

**DATA SET S4**, XLSX file, 0.2 MB.

**FIG S1**, EPS file, 2 MB.

**FIG S2**, JPG file, 2 MB.

**FIG S3**, EPS file, 1.5 MB.

**FIG S4**, JPG file, 1 MB.

**FIG S5**, EPS file, 1.9 MB.

**TABLE S1**, DOCX file, 0.02 MB.

**Acknowledgments**

We are especially thankful to icddr,b hospital and lab staff for support. We thank members of the Seed lab for critical feedback and thoughtful discussion regarding the manuscript.

The project described was supported by grant numbers R01AI127652 and R01AI153303 (K.D.S.) from the National Institute of Allergy and Infectious Diseases, and its contents are solely the responsibility of the authors and do not necessarily represent the official views of the National Institute of Allergy and Infectious Diseases or the NIH. K.D.S. is a Chan Zuckerberg Biohub Investigator and holds an Investigators in the Pathogenesis of Infectious Disease Award from the Burroughs Wellcome Fund. icddr,b gratefully acknowledges the following donors, which provide unrestricted support: Government of the People’s Republic of Bangladesh, Global Affairs Canada, Swedish International Development Cooperation Agency (SIDA), and the Department for International Development, UK Aid.

A.A., S.G.H., and K.D.S. designed the study. A.A. wrote the manuscript, which was edited by K.D.S. A.A. performed the sequence and bioinformatic analyses. S.G.H. performed PCR screens. S.G.H., M.H.T., and K.D.S. performed experiments to characterize PLE activity. F.J., M.S., and M.A. provided essential resources. M.A. and K.D.S. acquired funding. All authors reviewed and edited the manuscript.

K.D.S. is a scientific adviser for Nextbiotics, Inc. All other authors declare no competing interests.

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