Genomics of the Argentinian cholera epidemic elucidate the contrasting dynamics of epidemic and endemic *Vibrio cholerae*

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In order to control and eradicate epidemic cholera, we need to understand how epidemics begin, how they spread, and how they decline and eventually end. This requires extensive sampling of epidemic disease over time, alongside the background of endemic disease that may exist concurrently with the epidemic. The unique circumstances surrounding the Argentinian cholera epidemic of 1992–1998 presented an opportunity to do this. Here, we use 490 Argentinian *V. cholerae* genome sequences to characterise the variation within, and between, epidemic and endemic *V. cholerae*. We show that, during the 1992–1998 cholera epidemic, the invariant epidemic clone co-existed alongside highly diverse members of the *Vibrio cholerae* species in Argentina, and we contrast the clonality of epidemic *V. cholerae* with the background diversity of local endemic bacteria. Our findings refine and add nuance to our genomic definitions of epidemic and endemic cholera, and are of direct relevance to controlling current and future cholera epidemics.

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Latin America stands apart from the rest of the world in terms of its history of large-scale cholera epidemics. There have been seven global cholera pandemics in recorded history, between 1819 and the present day. Of these, the second pandemic (1829–1850) was the first to be seen in Latin America. Between the 1830s and 1895, the region suffered from repeated cholera outbreaks, linked to global pandemics 2 through 5 (refs. 4, 6). Importantly, although the sixth pandemic (1899–1923) affected most of Africa and Asia, Latin America did not experience cholera epidemics during this period, and thus was free of epidemic cholera for 96 years. In 1991, during the seventh pandemic (1961–present), cholera returned to Latin America. Phylogeographic data have shown that this was attributable to the simultaneous introduction of two distinct sub-lineages of a globally-circulating phylogenetic lineage of *Vibrio cholerae* serogroup O1 biotype El Tor, dubbed 7PET7–9. An antimicrobial-sensitive sub-lineage of 7PET (LAT-1) was introduced into Peru in January 1991 and spread rapidly across South America7,10,11. Concurrently, a separate and distinct drug-resistant 7PET sub-lineage (LAT-2) was introduced into Mexico7,12. The Haitian cholera epidemic in 2010 was also caused by 7PET, albeit by a third, independently-introduced sub-lineage, LAT-3 (refs. 7,8,12).

Since Latin America had been free of cholera epidemics for 96 years, and because the major epidemics seen in the 1990s and in 2010 are attributed to the introduction of epidemic 7PET sub-lineages originating in South Asia7, Latin America presented a unique opportunity to understand the longitudinal evolution of pandemic *V. cholerae* upon its introduction into a naïve population. Argentina is an ideal setting in which to study the evolution of the pandemic clone during the 1990s because, unlike some other countries in the region, the socioeconomic position of Argentina10,13, and its preparedness for epidemics by the time cholera reached the country, are thought to have enabled the monitoring and control of the epidemic. Argentina instituted the mandatory notification of cholera cases nationwide during 1991 after cholera broke out in Peru14, and developed public information campaigns which resulted in a concomitant increase in the rate of diarrhoeal disease reporting15. This included changes to the Argentinian national diarrhoeal surveillance system, and the creation of the National Diarrhoea Network (formerly the National Cholera Laboratories from the South).

Results
Cholera outbreaks occurred annually in Argentina (1992–1997). Between 1992 and 1998, WHO and PAHO data15 suggest that there were two peaks of cholera incidence in Argentina, one in 1993 (2,080 cases) and one in 1996/1997 (474 and 637 cases respectively) (Fig. 1a). However, there are discrepancies between these apparent maxima and other reports, which have suggested that there were seven epidemics of cholera in Argentina between 1992 and 1997 (refs. 14,12–25). To clarify this issue, we analysed the patterns of submission date for *V. cholerae* isolates sent to INEI. Between 10th February 1992 and 1st April 2002, INEI records show that the laboratory received at least 3,631 *V. cholerae* samples. Sixty-nine of these isolates were recorded as coming from countries other than Argentina: from Bolivia (*n* = 43), Chile (*n* = 16), Colombia (*n* = 1), Paraguay (*n* = 1), and Peru (*n* = 8). Four isolates have no recorded origin. The remaining 3,562 isolates originate from the North and Centre of Argentina (Fig. 1b), and may therefore represent over 82% of the PAHO/WHO-reported cholera cases for the whole country from this 1990s epidemic. Following the cholera outbreak in Peru, as part of epidemic preparedness, it was decided at the national level that every suspected cholera case reported in Argentina should be tested for *Vibrio cholerae* and the isolate sent to INEI from all microbiology laboratories from the network across the country, including the South region. *V. cholerae* was not reported by any of the laboratories from the South.

We used the metadata available for all *V. cholerae* samples received between 1992 and 2002, broken down by serogroup (O1/non-O1) and by serotype (Ogawa/Inaba). Of the isolates received, 2,189 were recorded as being of serogroup O1 (60.2%), and a total of 1,308 non-O1 *V. cholerae* were recorded. The vast majority (2,077, 94.8%) of *V. cholerae* O1 were of clinical origin, and 112 were either environmental isolates or their sources were not recorded. Of the non-O1 isolates, just 714 were of clinical origin (54.5%). From these data, it is clear that whilst there were periods during which no *V. cholerae* O1 were received (Fig. 1c), non-O1 *V. cholerae* were submitted to INEI more consistently during the 1990s, and their receipt rose coincidentally with peaks in over time after introduction to a region naïve to cholera, we sequenced nearly 500 *V. cholerae* isolates from the INEI collection. The collection is composed of all *V. cholerae* isolated from clinical suspected cholera cases as defined by the Ministry of Health between 1991 and 1998 (encompassing the epidemic period). During this time, all suspected cholera cases were tested in microbiology laboratories nationally, and putative *V. cholerae* were isolated and sent to INEI for further characterisation. After 1999, the National Diarrhoea and Foodborne Pathogens Network defined that one in every five stool samples should be tested for *Vibrio* spp., and all resultant *V. cholerae* isolates should be sent to INEI for further characterisation. The collection also includes *Vibrio* spp. from various environmental surveillance programmes, which had been collected to study the contribution of the environment to *Vibrio cholerae* dynamics in Argentina. Non-clinical environmental sources are defined by INEI as food, environmental and animal fodder; the majority of environmentally-isolated *V. cholerae* were derived from water sources, including drinking water.

We use these 490 clinical and environmental isolates and their associated records to link epidemiological reports of epidemic cholera in Argentina to genomic data. We also include non-O1 *V. cholerae* isolates of both clinical and environmental origins, to explore the underlying diversity of *V. cholerae* present in Argentina concurrent with the 1990s cholera pandemic, and to understand how these relate to those that caused the pandemic.
**V. cholerae** O1 receipt. In addition, there were 134 isolates for which there were no serogroup data recorded \((n = 129)\), that were autoagglutinable \((n = 4)\), or were recorded as being of serogroup O139 \((n = 1)\).

Figure 1c illustrates at least six peaks of **V. cholerae** O1 receipt within these data, occurring in early months of each year. This is consistent with previous reports, which allude to seven seasonal epidemics of cholera in Argentina during the 1990s\(^{14,23-25}\), and likely reflects the fact that WHO/PAHO data are only available as annual case/fatality numbers and are not broken down by month. **V. cholerae** serotype Ogawa dominated the number of received **V. cholerae** O1, with the exception of the incidence peak in early 1996. Ogawa isolates accounted for 1795 of all **V. cholerae** O1 (82.0%); 369 isolates were serotype Inaba (16.8%) (Fig. 1c). This is
in agreement with previous reports, which indicated that V. cholerae serotype Ogawa was predominant in Argentina during the epidemic, despite the initial cholera epidemic in Peru being ascribed to V. cholerae Inaba. Twenty-five V. cholerae O1 (1.1%) did not have a serotype assigned (Fig. 1c). The peak of V. cholerae in January 1993 for which there were no serogroup data recorded (n = 106; 79% of all no-data isolates) coincides with a peak in V. cholerae O1 receipt (n = 196), and describes an outbreak of V. cholerae O1.

These data, particularly the shifts between Inaba and Ogawa serotype, suggested subtleties in the dynamics of cholera epidemics in Argentina during the 1990s that could not be understood from epidemiological data alone. We revived and sequenced the genomes of 490 archived V. cholerae isolates from INEI archives. These were chosen principally to capture diversity of both O1 and non-O1 V. cholerae at the beginning (1992–1993) and the end (1996–1997) of the Argentinian epidemic (Fig. 1d). The sequenced isolates were a spatiotemporally-broad cross-section of cholera incidence, from all regions of Argentina that experienced cholera cases, and were chosen to capture apparent shifts between Inaba and Ogawa serotype (Fig. 1c, d).

The LAT-1 sub-lineage caused pandemic cholera in Argentina.

Most of the sequenced V. cholerae isolates were found to be members of the 7PET phylogenetic lineage (425/490, 86.7%)7. These sequences were placed into phylogenetic context with 518 additional 7PET genomes7 (Supplementary Data 1; Fig. 2). The vast majority of sequenced Argentinian 7PET V. cholerae were members of LAT-1 (421/425, 99.05%), the sub-lineage introduced into Peru in 1991 (ref. 7) (Fig. 2). No Argentinian isolates were members of the LAT-2 sub-lineage, which was introduced into Mexico in the early 1990s. In addition to the LAT-1 isolates, four isolates which lacked the genes encoding the cholera toxin (i.e. were non-toxigenic) clustered together with F99/W, a previously-described non-toxigenic 7PET genome, also from Formosa (Fig. 2; Supplementary Data 1; Supplementary Figs. 1–3).

In order to obtain a LAT-1-specific phylogeny, we mapped the reads for the genomes in this sub-lineage, as well as the direct ancestors of LAT-1 and related genomes from Angola, Côte d’Ivoire, and Sao Tome27,28, to a closed genome assembly of strain A1552 (ref. 29). This Inaba V. cholerae O1 was isolated in 1992 from a Peruvian traveller29,30 and harbours the WASA-1 genomic ancestors of LAT-1 and related genomes from Angola, Côte d’Ivoire, and Sao Tome (cluster 3, Fig. 3a).

Province-level geographic data were available for 475 of the 490 sequenced isolates, and for 416 of the 421 LAT-1 isolates (Fig. 3a; Supplementary Fig. 4). Sequences did not cluster by province or region. Strikingly, isolates from different Northern provinces were interspersed amongst one another, as were isolates from other countries in the region, such as Bolivia (Fig. 3a). However, we did observe limited clustering by date of isolation. Argentinian isolates from multiple provinces in 1996 and 1997 clustered phylogenetically, and isolates from 1997 occupied cluster 1 (Fig. 3a). Similarly, cluster 2 contains Argentinian genomes from 1992 to 1993 and one isolate from 1997, from multiple provinces, as well as one Bolivian and one Peruvian genome from 1992 to 1991 respectively (Fig. 3a; Supplementary Data 2, see Microreact project linked to this paper for an interactive map).

The LAT-1 sub-lineage has limited genetic variation.

Across all LAT-1 genomes and across the entire time period, we observed a striking lack of variation, at the levels of SNVs, gene gain/loss, and recombination. We found that just 0.03% of the A1552 genome was predicted to have undergone recombination (Supplementary Fig. 5), and a mean of only 26.05 non-recombining SNVs across both chromosomes separated the sequence of each LAT-1 isolate from that of the A1552 reference genome (Fig. 3b) (min 10, max 149, stdev 14.10). Of the 2,651 SNVs in the LAT-1 alignment, 72.6% were private to single genomes in the data set. This result contrasted with genomic studies of 7PET in other settings; in hyper-endemic settings, six co-circulating sub-lineages of 7PET could be identified over a 4-year period32, in Africa, multiple 7PET sub-lineages have co-existed and replaced one another since 1970 (refs. 28, 33), as has been observed in China34,35. This provided further evidence supporting the hypothesis that the Argentinian cholera epidemic was caused by one highly-clonal sub-lineage of 7PET.

We calculated a pangenome for LAT-1, and found that 3,368 genes were core to these isolates (present in ≥97% of isolates), suggesting that ~89% of the 3,776 annotated genes in the A1552 reference genome are core to LAT-1. Although gene gain and loss events were rare within LAT-1, there was evidence of the loss of the entire CTX bacteriophage within the data set, as it was absent from 51 of the isolates in the LAT-1 phylogeny (Fig. 3a, Supplementary Fig. 6A). It is possible that this loss was a result of long-term culture (some isolates were stored for 27 years before being sequenced) as has been noted previously32,36. We did not identify any candidate genes, which might have influenced the rapid spread of LAT-1. Rare and sporadic gene gains were also evident: four Argentinian LAT-1 V. cholerae carried extended-spectrum β-lactamase (ESBL) genes, blaCTX-M-3, blaOXA-8, and blaTEM (Fig. 3a; Supplementary Figs. 7, 8). By manual inspection of the genome assemblies, we confirmed that these three ESBL genes were carried on contigs that also included IncA/C plasmid replicons (Fig. 3a; Supplementary Figs. 8, 9; Supplementary
Data 2). Multidrug resistance plasmids have been seen in *V. cholerae* strains from Argentina previously, Algeria, and in Angolan isolates from 1988 and the early 1990s, two of which did harbour *bla*TEM (Supplementary Fig. 8; Supplementary Data 2). However, the complement of resistance determinants in these isolates does not match those found in Argentinian *V. cholerae* (Supplementary Fig. 8, Supplementary Data 2).

Serotype variation and multiple *wbeT* mutations in LAT-1. The Inaba and Ogawa serotypes of *V. cholerae* O1 are commonly differentiated by diagnostic laboratories and by epidemiologists, as well as in the formulation of cholera vaccines, such as Dukoral, because both serotypes elicit different immunological responses. Methylation of the terminal perosamine sugar on the O1 lipopolysaccharide chain by the WbeT enzyme confers an Ogawa phenotype; lack of methylation by disruption of *wbeT* confers an Inaba serotype. There is evidence that reversion from Inaba to Ogawa serotype can occur in vivo, albeit rarely.

In order to explain the apparent shifts between Inaba and Ogawa *V. cholerae* seen in Argentina (Fig. 1c), and more broadly in Latin America, we examined the sequence of *wbeT* in LAT-1. We identified nine distinct mutations in *wbeT* across LAT-1 which were predicted to disrupt the WbeT protein by non-synonymous, frameshift and truncating mutations (Fig. 3a; Supplementary Fig. 7, Supplementary Data 2, see Methods for full details). The genomic predictions correlated well with the longitudinal data detailed in Fig. 1c and with the phenotypic serotype assigned to each isolate; the *wbeT* genotype matched the phenotypic serotype for all but two of the 398 serotyped LAT-1 isolates sequenced in this study (99.4% concordance) (Fig. 3a; Supplementary Figs. 7, 8; Supplementary Data 2; Methods).

From previous reports, we know that the initial 1991 cholera epidemics in Peru and elsewhere in Latin America were associated with serotype Inaba *V. cholerae*, which became dominated by serotype Ogawa bacteria in 1992 and thereafter. Genome sequences show that the Peruvian Inaba isolates from 1991 harbour the N165fs mutation in *wbeT* (Fig. 3a; Supplementary Fig. 7, Supplementary Data 2). Since LAT-1 was introduced into Latin America from West Africa, we compared these data to West African Inaba isolates sharing a common ancestor with LAT-1, but collected between 1992 and 1995.
Angola, just after LAT-1 had been introduced into Peru\(^7,28\). These isolates were found to harbour a different mutation, N62fs (Fig. 3a; Supplementary Fig. 7). The N165fs mutation is present in 68 of the genomes of LAT-1 isolates collected since 1991. These include isolates from Brazil, Mexico, Chile, Argentina and Colombia, as well as isolates from Peru, all of which were originally serotyped as Inaba (Fig. 3a; Supplementary Figs. 7, 8; Supplementary Data 2). In addition, environmental isolates from Mexico, collected between 2004 and 2010, also harbour this mutation and are part of the same cluster of isolates (Fig. 3a; Supplementary Fig. 7). Hence, the N62fs and N165fs mutations are likely to have arisen independently, prior to spreading within West Africa and Latin America, respectively.

It has been hypothesised that cholera entered Argentina through the North of the country, which shares borders with Chile, Bolivia, Paraguay and Brazil\(^22\). Genomes from bacteria isolated in 1991 and 1992 from Chile, Bolivia and Brazil were included in our phylogeny\(^7,9\) (Fig. 3a). These were either Ogawa (Bolivia, \(n=7\); Brazil, \(n=1\)) or Inaba (N165fs; Brazil, \(n=6\); Chile, \(n=1\)) serotype. We found that these were interspersed amongst contemporaneous serotype Ogawa isolates, which were collected in Northern provinces of Argentina (Fig. 3a). All of these isolates were members of cluster 4, except for one Bolivian genome (1992) which was a member of cluster 2 (Fig. 3a). This observation, and the lack of genetic diversity within LAT-1, are consistent with the same \textit{V. cholerae} sub-lineage circulating within, and between, countries at the Northern border of Argentina.

In 1996, cholera cases resurged in Argentina\(^17\) following a relative lull in 1995. This was associated with serotype Inaba

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**Fig. 3 A phylogeny of the LAT-1 sub-lineage.** a A maximum-likelihood phylogeny was calculated from 2,651 non-recombinant SNVs determined relative to the A1552 reference sequence (see Methods). The tree is rooted on the sequence of CNRVC980048, an isolate lacking WASA-1 from Burkina Faso, 1984. Scale bar denotes the number of mutations per variable site. Clusters were defined using Fastbaps using an alignment of 725 parsimony-informative SNVs. Metadata are reported where available; gaps are presented where data were unavailable, not determined, or recorded as no data. Initial figure produced using iCANDY and re-coloured manually (Adobe Illustrator CC). b Density plot illustrating the number of SNVs separating each of the LAT-1 isolates from the A1552 reference sequence. The blue line indicates the mean (26.05 SNVs). Interactive phylogeny available in Microreact (https://microreact.org/project/VAZD_K0kZ).
**V. cholerae** (Fig. 1c). We found that 17 Argentinian Inaba **V. cholerae** isolates from 1996 formed a closely-related subclade within cluster 4 of the LAT-1 phylogeny (Fig. 3a), and harbour a unique mutation in wbeT, Q274trunc. This clade includes one 2010 Inaba isolate from Mexico (Fig. 3a; Supplementary Figs. 7 and 8). In addition, this subclade shares a common ancestor with the clade of isolates from 1997 which are serotype Ogawa and form a separate phylogenetic cluster (cluster 1, 48 isolates, Fig. 3a; Supplementary Figs. 7, 8). The 1996/1997 outbreak was not geographically-restricted; isolates from multiple provinces were part of this clade (Fig. 3a).

**Non-7PET diversity contrasts with LAT-1 clonality.** Sixty-five isolates sequenced in this study were not members of 7PET, but were obtained from the same regions and times as the LAT-1 isolates (Figs. 1d, 4a, b; Supplementary Data 3). Thus, we placed these into context with a more diverse collection of **V. cholerae** sequences2, together with genomes from a recently-published study of non-epidemic **V. cholerae** in China48, and then calculated a pangenome using these sequences (Fig. 4). The rate of gene discovery as sequences were added to the non-7PET pangenome was much greater than was observed in a LAT-1 pangenome, despite there being 38% more isolates in the LAT-1 pangenome (Fig. 4c, d). This indicates that genes are not being gained or lost by LAT-1. The non-7PET isolates were also extremely genetically diverse in comparison to the 7PET genomes, with a mean average nucleotide identity (ANI) relative to A1552 of 97.61 (min 95.90, max 99.65, stdev 0.960; Fig. 4e), in contrast to LAT-1 (mean ANI 99.99, min 99.96, max 99.98, stdev 0.0032; Fig. 4e). An ANI value of 95% is a common threshold for separating species49. The non-7PET isolates had a considerably expanded accessory genome when compared to LAT-1 (23,458 cloud genes in the collection of 383 diverse genomes compared to 3,313 in the 532 LAT-1 genomes) (Supplementary Fig. 6).

Of these 65 isolates, four were phenotypically and genotypically serogroup O1. Two were members of the previously-described Gulf Coast lineage of **V. cholerae** O1, including the single sequenced **V. cholerae** O1 from 1998. Both Gulf Coast isolates harboured CTX and were toxigenic, and the two remaining **V. cholerae** O1 isolates were members of ELA-3 (ref.7) (Supplementary Fig. 10a; Supplementary Data 3). All four isolates were of clinical origin (Supplementary Data 3). The remaining 61 isolates lacked the genes required to produce cholera toxin, and were confirmed in silico not to harbour genes encoding the O1 antigen, though 45 of these were of clinical origin. We identified four new lineages of non-O1 non-7PET **V. cholerae** amongst these isolates, defined as clades formed by three or more Argentinian non-7PET isolates in the phylogeny, and denoted as A1–A4, where A stands for ‘Argentina’ (Fig. 4f). These lineages contained isolates that were of clinical origin alone (A1, A3) or clinical and environmental origin (A2, A4), were acquired in different years (A3, A4), and from different regions (A2, A3, A4), suggesting that these represent populations of non-7PET **V. cholerae** local to Argentina (Fig. 4f, Supplementary Data 3).

Of the 61 non-O1 non-7PET isolates, 21 harboured one of three distinct Type III secretion systems (Fig. 4f; Supplementary Data 3). These included the T3SS-2a described in **V. cholerae** AM_19226 (refs. 50,51), the less-common T3SS-2β system described by Carpenter et al.31, and a third putative T3SS system which most closely resembles genes present in the genomes of two virulent Chilean *Vibrio anguillarum* isolates32 (Supplementary Fig. 10). This putative T3SS was found in lineage A2. The presence of T3SS-2β in lineage A3 is of particular interest — this lineage is composed of clinical isolates, contains the previously-described Argentinian isolate TUC_T2734 (ref. 7), and includes one isolate from Salta province in the year 2000. T3SS elements were mutually exclusive and were never detected in the same genome at the same time (the distribution of T3SS is described in Supplementary Fig. 10 and Supplementary Data 3). From these limited data it is clear that more T3SS-positive non-7PET were of clinical origin than environmental (T3SS-2α: 10 clinical, 2 environmental; T3SS-2β: 5 clinical, 0 environmental; **V. anguillarum** element; 1 clinical, 3 environmental). It is also important to note that none of these T3SS are present in 7PET.

**Discussion** We contend that the observations described here could only have been made in Latin America and in Argentina. This is because the limited introductions of 7PET sub-lineages into the region, and the consequentially-enhanced surveillance schemes, provided a unique opportunity to understand the long-term evolutionary dynamics of epidemic **V. cholerae** evolving from point-source introductions on a continent where there have been millions of cholera cases from 1991 to present. This is the largest genomic study to date that investigates pandemic cholera in a single country, and we believe that it is the largest sequencing project designed to investigate the genomics of a single bacterial pathogen in Argentina. These data have had direct impact on national health policies in Argentina by changing the national alert system to distinguish between pandemic 7PET lineage and local **V. cholerae** using whole-genome sequencing. This recognises the increased risk presented by an outbreak of 7PET relative to that of a non-7PET outbreak, even if toxigenic or serogroup O1. This is to ensure that an epidemic response focuses on high-risk 7PET clones, and that there is also efficient monitoring of the contributions of endemic non-7PET clones to public health, including via environmental surveillance using a One Health approach.

Our data show that Ogawa **V. cholerae** from Argentina in 1992 are closely related to the Inaba isolates sequenced from Peru (Fig. 3a), and show that the Inaba/Ogawa shift observed in Peru and elsewhere in Latin America represented variation within LAT-1, rather than a separate introduction of another strain (Fig. 3a; Supplementary Fig. 8). Likewise, the outbreak of Inaba **V. cholerae** in Argentina in 1996 arose by mutation of wbeT from wild-type to an Inaba genotype (Q274trunc), and this mutation may have occurred in Argentina. The Q274trunc mutation is distinct from others identified within the data set, particularly the mutation associated with the Inaba phenotype in contemporaneous Mexican isolates (N165fs). This indicated that Argentinian cholera in 1996 was not caused by an introduction of a new Inaba (sub)lineage from elsewhere in Latin America; rather, LAT-1 **V. cholerae** that had already been introduced into Northern Argentina or the neighbouring countries acquired a new Inaba genotype and phenotype. The 1997 Argentinian outbreak, in turn, was caused by a close relative of the 1996 Inaba clone, but the topology of our phylogeny suggests that this was not a result of reversion from the Inaba Q274trunc genotype to an Ogawa genotype (Fig. 3a), adding nuance that would have been useful for public health at the time of the epidemic. These data underline that Ogawa/Inaba phenotypic variation is not phylogenetically informative, and may not be appropriate to use as an epidemiological marker.

Perhaps surprisingly, in spite of the sustained circulation of LAT-1, which was disseminated across Northern Argentina, an area of ~1.2 million km² (Fig. 1b, Supplementary Fig. 4), these data suggest that very little genetic change, at the level of SNVs and gene gain/loss, occurred in this sub-lineage over a period of nearly 6 years. This lack of diversity is reminiscent of the clonality
in other bacterial pathogens, such as *Salmonella Typhi* H58 (ref. 53). Our data also show that LAT-1 circulated amongst the countries at the Northern borders of Argentina during the early 1990s — for instance, cholera was first reported in Bolivia in August 1991 (refs. 5,54), and Bolivian genomes from the early 1990s are intermixed amongst the Argentinian genomes from 1992 (Fig. 3a; Microreact). The lack of clustering by region is likely to reflect the rapid dissemination of LAT-1 across South America.

LAT-1 invariance is juxtaposed with the diversity observed in non-O1 *V. cholerae* in Argentina (Figs. 3 and 4; Supplementary Fig. 6). Although *V. cholerae* research has tended to focus on studying epidemics and outbreaks, by definition, this tends to describe epidemic lineages. Non-7PET *V. cholerae* are highly
variable, and as well as examples of local lineages of non-7PET V. cholerae, we also identified isolates which were confirmed microbiologically to be Vibrio cholerae, but were diverse phylogenetically and as measured by ANI values. Non-7PET V. cholerae continue to be associated with clinical cases of disease, but remain understudied. The disease associated with these isolates — and whether virulence determinants such as T3SS contribute to this disease — is the focus of future work, though there is evidence to suggest that T3SS contributes to diarrhoea caused by non-7PET V. cholerae. We also see here, with the caveat of a small sample size, that the clinical non-7PET isolates were enriched for the presence of T3SS (16/21 isolates).

We show that a single clone of V. cholerae O1 (Fig. 2), now known to be one sub-lineage of 7PET, was responsible for pandemic cholera in Argentina, in spite of the seasonal fluctuations and serotype variation observed (Fig. 1c). Our genomic data provide fine-scaled insight into the evolution of LAT-1 that would not have been captured by routine surveillance alone. However, it is also important to state that these data validate fundamental observations made by public health authorities during the cholera epidemics of the 1990s — that the outbreaks in Argentina were caused principally by Ogawa V. cholerae, which were closely related to the Peruvian strain as determined by PFGE. Furthermore, by including non-7PET V. cholerae in our study, we found a highly diverse population of this species in Argentina existing concurrently with the extremely invariant LAT-1 pandemic sub-lineage. We suggest that these non-7PET, which include O1 and non-O1 serogroup isolates, represent those V. cholerae that are truly endemic to Argentina, and are evolving locally but lack the propensity to cause epidemics and to spread in the same way as 7PET. Therefore, the fact that Latin America was cholera-free for 97 years was due solely to the absence of pandemic lineages, and it is a consequence of elevated surveillance due to the LAT-1 introduction that endemic V. cholerae were captured. In the absence of clinical data associated with these non-7PET isolates, we cannot determine whether they are aetiological agents of cholera, or of a cholera-like illness. However, it is clear that non-7PET V. cholerae were present, and associated with disease at a low level, in Argentina throughout the 1992–1998 cholera epidemic and thereafter (Fig. 1c).

It is vital to understand the diversity of the local, endemic V. cholerae that co-exist alongside 7PET during a cholera epidemic. This is because non-epidemic V. cholerae present in a country may contribute to cases of disease that are symptomatic of cholera, but do not pose the same relative risk to public health as 7PET. The case in the Chaco region during 2005, which was caused by non-epidemic V. cholerae O1 of the MX-2 lineage and failed to cause epidemic cholera, exemplifies this point. Similar observations have recently been made in China. Relative risk of V. cholerae lineages should be accounted for in the magnitude of epidemic preparedness responses to such outbreaks, as is now being done in Argentina. The Global Task Force on Cholera Control has committed to reducing deaths from cholera by 90% before the year 2030 (ref. 59). This campaign focuses on the control of cholera, the disease, rather than on 7PET, the aetiological agent of epidemic cholera. As cholera control is implemented in countries that currently experience a high incidence of cholera attributed to 7PET, cases of cholera will decline. We anticipate that as epidemic cholera is reduced in magnitude, disease caused by non-7PET V. cholerae will become more visible, just as has been observed in Argentina. By using genomic definitions to differentiate pandemic lineages for public health epidemic preparedness responses, as is being done in Argentina, we propose that concerted control efforts including epidemiologists, public health authorities and microbiology laboratories targeting 7PET specifically, and accounting for background levels of endemic non-7PET disease, could see epidemic cholera eliminated in Latin America.

**Methods**

### Bacterial strains and oligonucleotide primers

A list of bacterial isolates sequenced in this study is reported in Supplementary Data 1–3, together with relevant metadata and results from genomic analyses. Additional genome sequences used for contextualisation are listed in Supplementary Data 1–3. The sequences of oligonucleotides used for PCR (see Microbiology section below and Supplementary Methods) are detailed in Supplementary Table 1.

### Bacterial culture, genomic DNA isolation and sequencing

Vibrio cholerae isolates were revived from archived stocks on tryptic soy agar plates or in alkaline peptone water, and were subsequently passaged on thiosulphate-citrate-bile salts media to select for V. cholerae. Minimal biochemical tests (oxidase, Kligler tests) were performed to confirm Vibrio spp. identity. Genomic DNA extractions at INEI were carried out from 1 ml of liquid culture using the QiaCube Connect (Qiagen). Extractions carried out at the Sanger Institute were performed using the Epicentre Masterpure kit and a modified version of the manufacturer’s protocol — briefly, single purified colonies of V. cholerae were used to lawn an LB agar plate. Loopfuls of bacterial laws were resuspended in 300 μl Tissue & Cell Lysis Solution supplemented with Proteinase K, incubated at 65 °C for 20–25 min with intermittent vortexing to lyse cells, and then treated with RNase A at 30 °C to remove contaminating RNA. Thereafter, the manufacturer’s protocol was followed to remove protein contaminants and to purify genomic DNA. Approximately 0.5 μg of genomic DNA per isolate was used for sequencing with the Illumina HiSeq X10 platform at the Wellcome Sanger Institute.

### Microbiology

Isolates were received and subjected to biochemical and molecular testing by INEI at the time of their receipt, principally during the 1992–1998 epidemic period. During the cholera epidemic, minimal biochemical tests (Kligler, oxidase, haemagglutination and indole) and complementary tests (chloride and deoxyribonuclease) were performed to verify the identity of V. cholerae. The production of cholera toxin was assessed using enzyme-linked immunosorbent (ELISA) assays. Where required, PCR was performed to confirm an isolate as V. cholerae, as well as the presence of genes encoding tcpA and tcpB, and other virulence factors. The oligonucleotide primer sequences and reaction conditions for these PCR tests are detailed in Supplementary Methods and Supplementary Tables 1–5. All of the microbiological and molecular assays used at INEI-ANLIS for the characterisation of V. cholerae are also described in a publicly-accessible protocol manual.

### Sequence data quality control

A total of 21 sequenced isolates contained substantial amounts of contaminating sequences from non-Vibrio species, and were excluded from this study, for a total of 490 sequences used in this analysis. Contamination was assessed using Kraken, by examining the overall length of the SPAdes assembly (data were summarised using assembly-stats v1.0.1).
using SPAdes v3.8.2 (ref. 65) as part of a high-throughput analysis pipeline66, and annotated using Prokka v1.5 (ref. 67). External publicly-available sequences were similarly assembled from raw sequencing reads — where these were not available, assemblies were downloaded from Genbank and annotated using Prokka v1.5 for uniformity within the data set.

**Pangenome analysis**. Pangenomes were calculated using annotated genome sequences for a diverse collection of V. cholerae (Supplementary Data 3) as well as the LAT-1 subset of a 7PET-specific data set (Supplementary Data 2). Roary v3.12.0 (ref. 68) was used for these calculations, with options ‘–e-mafﬁ –s -cd 97’. For the non-7PET genome collection, an alignment of 2719 core genes was used for phylogenetic analysis (see below).

**SNV identiﬁcation and phylogenetic analysis.** For 7PET and LAT-1 phylogenetic analyses, sequencing reads were mapped to reference genomes (accession numbers LT907989/LT907990 for N16961; CP025936/CP025937 for AI552) using SMALT v0.74 (http://www.sanger.ac.uk/science/tools/smalt-0). The reference was indexed using SAMLT using a kmer size of 20 and a step size of 13 (k 20 + 13), and the reads were aligned using default parameters but with the maximum insert size (i) set as three times the mean fragment size of the sequencing library (target insert size 450 bp). PCR duplicate reads were identiﬁed using Picard v1.92 (https://broadinstitute.github.io/picard) and ﬂagged as duplicates in the BAM ﬁle. High-quality single nucleotide polymorphisms, including small indels, were identiﬁed as described by Harris et al.49 and reported previously for V. cholerae22,26, Brieﬂy, CBFs of variant sites were generated using samtools mpileup v0.1.19 (ref. 70) (parameters -d 1000 -Dhight) and bcftools v0.1.19 (http://samtools.github.io/ bcftools/). The bcftools option to call genotypes at variant sites was used. The following bcftools cut-off thresholds were applied: quality <50, map quality <30, af1 <0.95, ratio <0.75, depth <4, depth_strand <2, strand_bias <0.001, map_bias <0.001. If any of these ﬁlters were not met, the base called as uncertain. A pseudo-genome was constructed by substituting the base call at each site (variant or non-variant) in the BCF ﬁle into the reference genome. Uncertain sites were substituted with an N. Insertions with respect to the reference genome were ignored and deletions with respect to the reference genome were ﬁlled with Ns in the pseudo-genome to keep it aligned and the same length as the reference genome.

Regions of the genome which were predicted to be recombined, and which might therefore affect the topology of calculated phylogenies, were identiﬁed and removed from the pseudogenome alignment using Gubbins v4.1.10 (ref. 71). Alignments consisting entirely of variable nucleotides were produced from whole-genome alignments using SNP-sites v2.4.1 (ref. 72). The non-7PET core-gene alignment was trimmed using trimal v1.4.1 (ref. 73), and SNP-sites v2.5.1 was used to produce an alignment of 201,790 variable nucleotides. Maximum-likelihood phylogenetic trees were then calculated from SNV-only alignments using IQ-Tree v1.6.10 (ref. 74) under the general time reversible (GTR) and ascertainment bias correction (ASC) models75. Five thousand approximate likelihood ratio tests76 and ultrafast bootstrap approximations77 were performed to assess the robustness of the computed phylogenies.

LAT-1 genomes were clustered using Bayesian hierarchical clustering and partitioned using the Dirichlet Process Mixture model with Fastbaps v1.0.1 (ref. 31), run using default parameters. Fastbaps was similarly used to cluster sequences in the V. cholerae species phylogeny, using the Bayesian Hierarchical Clustering prior28 and excluding the three outgroup sequences from the alignment. Parsimony-informative SNVs were extracted from SNV-only alignments using extract_PL_SNPs.py (https://gist.github.com/asanashah9/396dcd014b33c3ce12514) and these alignments were used as the input for Fastbaps. SNV distance matrices were calculated from SNV-only alignments using snp-dists v0.4 (https://github.com/tseemann/snp-dists). Average nucleotide identity (ANI) values were calculated using FastANI v1.0 (ref. 49).

**Detection of plasmid replicons, antimicrobial resistance genes, ctxB variants and in silico serotype assignment.** wcbE and ctxB gene sequences were assigned using ARIBA v2.12.1 (ref. 78) and a custom database consisting of the ctxB nucleotide sequence from N16961 (LT907989/LT907990) and the intact wcbE sequence from NCTC 9420 (ref. 80) (CP013319/CP013320), which translates into a protein sequence which is 100% identical to the WcbE sequence from the Ogawa isolate VX44945 (AEAN0191.1)81. We assumed that an Inaba phenotype would be conferred on isolates in which ARIBA was unable to detect or assemble wcbE in its totality, and if a mutation in wcbE was detected that was predicted to frameshift or truncate translated wcbE (N62Ls, N165Fs, F244Fs, Q274trunc), this was associated with Inaba phenotypes (I206K), or was otherwise known to confer an Inaba phenotype (S158F)82. We assumed, since none of the isolates harbouring the E36D wcbE mutation was an Inaba phenotype, that this mutation does not result in abolition of the Ogawa serotype. The sequences of other virulence genes were taken from the sequence of N16961 (LT907989/LT907990) or of the Classical V. cholerae isolate O939 (CP000626/CP000627). We confirmed using BLASTn and the pangenome gene presence/absence matrix that the WASA-1 genomic island, a marker characteristic of the LAT-1 sub-lineage32, was present in the assemblies for these sequences. Plasmid replicons and antimicrobial resistance genes were detected using ARIBA, the ResFinder database83, and the PlasmidFinder database84 (both databases accessed on 23/06/2019). The presence and absence of the V. cholerae serogroup O1 biosynthesis operon in non-7PET genomes was confirmed using the pangenome gene presence/absence matrix, and by testing for the presence of the O1 biosynthesis operon sequence using BLASTn (co-ordinates 234,000–286,000 in the N16961 reference genome, accession LT907989).

**Data visualisation.** Data were visualised and maps were annotated using Tablet Desktop 2018.31. Maps in Tablet were produced using OpenStreetMap (7 OpenStreetMap contributors) which is licensed under a CC-BY-SA licence (https://www.openstreetmap.org/copyright). Phylogenetic trees were visualised using Figtree v1.4.3 (http://tree.bio.ed.ac.uk/software/ﬁgtree/) and iTOL v3 (ref. 85). Gene presence/absence matrices were visualised using roary_plots.py v0.10.0 (https://github.com/sanger-pathogens/Roary/tree/master/contrib/roary_plots).

Other ﬁgures were produced using R v3.5.1 with the ggplot2 v3.1.1 (ref. 86) and reshape v0.8.8 (ref. 87) packages, Artemis v16 (ref. 88), ACT v13 (ref. 88), DNA-plotter v1.11 (ref. 89), the Phandango web server90, Easyf v2.2.2 (ref. 90), and iCANDY (https://github.com/simonharris/iCANDY). Where ﬁgures were manually edited using Adobe Illustrator CC v23.0.4.

**Ethics.** Not applicable. This study uses archived bacterial samples processed by INEI. No identiﬁable data were available or used in this study.

**Reporting summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**

All next-generation sequencing data generated in this study have been deposited into the European Nucleotide Archive (ENA; http://www.ebi.ac.uk/ena) under accession number ERP118863 (https://www.ebi.ac.uk/ena/browser/view/ERP118863). The assemblies which were used to produce Supplementary Figs. 9, 10B have been deposited into the ENA as part of ERP118863 (https://www.ebi.ac.uk/ena/browser/view/ERP118863). The original data which underpin Fig. 1 are held and maintained by INEI-ANLIS, a National Reference Laboratory of Argentina. Therefore, in order to maintain conﬁdentiality and protect privacy, these raw data have not been published with the article, but managed access to these records can be facilitated on request (via Dr. J.C., senior author). All other data used to generate ﬁgures in this paper, including sequence alignments, phylogenetic trees, and data matrices, are available in Figs hare (https://doi.org/10.6084/m9. ﬁgshare.1130131) or in the Supplementary Data associated with this paper. An interactive LAT-1 phylogeny is available in Microreact93 (https://microreact.org/project/ VAZD_K0kZ).

**Code availability**

The R code used to produce the initial versions of Figs. 3b, and 4e is available in Figs hare (https://doi.org/10.6084/m9.ﬁgshare.1130131).

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**Author contributions**

N.R.T. and J.C. designed the study and supervised the work. S.C. supervised C.T., M.R.V., M.P., M.M. curated and cultured isolates. M.R.V., G.Z., L.K., D.D., C.T., J.C. and M.J.D. prepared gDNA for sequencing. M.J.D. and D.D. performed genomic analyses. T.P. digitised metadata. M.J.D. and T.P. analysed metadata. M.J.D., G.D., J.P., G.P.C.S., N.B., M.L.C., J.C. and N.R.T. analysed and interpreted data. M.J.D. wrote the paper, with major contributions from D.D., J.C. and N.R.T. All authors contributed to editing the paper.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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