The coronavirus disease 2019 (COVID-19) pandemic caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has spread globally, with >365,000 cases in California as of 17 July 2020. We investigated the genomic epidemiology of SARS-CoV-2 in Northern California from late January to mid-March 2020, using samples from 36 patients spanning nine counties and the Grand Princess cruise ship. Phylogenetic analyses revealed the cryptic introduction of at least seven different SARS-CoV-2 lineages into California, including epidemic WA1 strains associated with Washington state, with lack of a predominant lineage and limited transmission among communities. Lineages associated with outbreak clusters in two counties were defined by a single base substitution in the viral genome. These findings support contact tracing, social distancing, and travel restrictions to contain the spread of SARS-CoV-2 in California and other states.

The novel severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), which causes coronavirus disease 2019 (COVID-19), is a pandemic that has infected more than 13.8 million people around the world and caused more than 591,000 deaths as of 17 July 2020 (1), including >3.5 million cases in the United States and >365,000 in California. An exponential growth in the number of cases has overburdened clinical care facilities and threatened to overwhelm the medical workforce. The reported case numbers also underestimate the true number of infections because of the presence of asymptomatic or mild cases who do not get tested (2–4). As a result, California, along with many other states and countries, has issued a “shelter-in-place” policy for all residents, effective 20 March 2020 and ongoing at the time of this report. These unprecedented measures have disrupted daily life for ~40 million inhabitants of the state and have incurred profound economic losses (5).

Until late February 2020, the majority of infections identified in the United States were related to travelers returning from high-risk countries, repatriated citizens under quarantine, or close contacts of infected patients. Community spread, in which the source of the infection is unknown, has since been documented in multiple states. In particular, Washington state reported a series of COVID-19 cases from 21 January to 18 March, following the identification of the earliest case reported in the United States, WA1, on 19 January; this suggests that a persistent WA1 lineage transmission chain was present in the community during that time period (6, 7).

Genomic epidemiology of emerging viruses has proven to be a useful tool for outbreak investigation and for tracking virus evolution and spread (7–9). During the Ebola virus disease epidemic of 2013–2016 in West Africa, genomic analyses established that the outbreak had a single zoonotic origin (9), that two major viral lineages were circulating (10), and that sexual transmission played a role in maintaining some transmission chains (11). Viral genome sequencing also uncovered the route that Zika virus traveled from northern Brazil to other regions (12), including Central America and Mexico (13) and the Caribbean and United States (14). However, real-time genomic epidemiology data for COVID-19 to inform public health interventions in California have been limited to date.

We recently developed a method called MSSPE (metagenomic sequencing with spiked primer enrichment) to rapidly enrich and assemble viral genomes directly from clinical samples (15). Here, we used this method and/or tiling multiplex polymerase chain reaction (PCR) to recover viral genomes from COVID-19 patients in Northern California and to perform phylogenetic analyses, with the goal of better understanding the genetic diversity of SARS-CoV-2 in the United States and the nature of transmission of virus lineages in the community.

We screened a total of 62 respiratory swab samples from 54 COVID-19 patients available from hospitals and clinics at the University of California, San Francisco (UCSF), the California Department of Public Health (CDPH), and eight county public health departments in Northern California (table S1). Presumptive positive cases were confirmed to be SARS-CoV-2-infected by a U.S. Centers for Disease Control and Prevention (CDC) assay approved by a Food and Drug Administration (FDA) Emergency Use Authorization (EUA) on 4 February 2020 (16). SARS-CoV-2 genomes (>62% coverage) were recovered from 36 patients (Fig. 1A and table S2). The 36 infected patients for whom viral genomes were obtained were collected from 29 January to 20 March 2020 and spanned nine counties in Northern California (Fig. 1B and table S2). The 36 infected patients for whom viral genomes were obtained were collected from 29 January to 20 March 2020 and spanned nine counties in Northern California (Fig. 1B and table S2). The 36 infected patients for whom viral genomes were obtained were collected from 29 January to 20 March 2020 and spanned nine counties in Northern California (Fig. 1B and table S2).
SARS-CoV-2 RNA genome, followed by metagenomic next-generation sequencing (mNGS) of pooled and indexed samples on Illumina NextSeq, HiSeq, or MiSeq instruments (18, 19). The PCR cycle thresholds ranged from 15.3 to 33.4, corresponding to virus loads of $1.1 \times 10^4$ to $2.7 \times 10^8$ copies/ml (fig. S1 and table S2). An average of 31 million [interquartile range (IQR), 23 million to 57 million] and 2.2 ± 0.2 million reads were generated per sample for use in M Snape and tiling multiplex PCR, respectively, and virus genomes were assembled by mapping to reference genome NC_045512 (Wuhan-Hu-1). The assembly yielded 34 SARS-CoV-2 genomes with genome coverage exceeding 65%, and these were included in the study. An additional two genomes sequenced from samples of a returning traveler from Wuhan, China and a household contact collected on 29 January by the CDC (CA3 and CA4) were also included in the analysis. The median coverage achieved across all samples was 97.7% (IQR, 90.4% to 99.7%).

Phylogenetic analysis revealed that the 36 SARS-CoV-2 genomes from California generated in this study were dispersed across the evolutionary tree of SARS-CoV-2 that was constructed from 789 worldwide genomes deposited into GISAID as of 20 March 2020 (Fig. 2A). The 36 genomes included 14 in the Washington state (WA1) lineage, 10 in a lineage associated with the Santa Clara County outbreak cluster (henceforth the SCC1 lineage), three from a Solano County cluster of three individuals, five related to lineages circulating in Europe and New York, and four related to early lineages from Wuhan or other regions of China (including two patients from San Benito County with identical genomes) (Figs. 1, 2A, and 3 and table S2).

A large outbreak was associated with travel on the Grand Princess cruise ship (with at least 78 confirmed positive cases out of 469 tested) as of 26 March (20). The Grand Princess undertook two consecutive voyages from San Francisco (voyage A to Mexico, 11 to 21 February; voyage B to Hawaii, 22 February to 4 March), with much of the same crew and a shared subset of passengers. Samples from 11 infected patients were sequenced, three of whom had been on voyage A and became sick after returning to their home county, and eight from crew members and passengers aboard the cruise ship on voyage B. Note that all 11 available sequenced genomes from the Grand Princess were part of the WA1 lineage (Fig. 2, A and B, and Fig. 3). In addition to sharing three single-nucleotide variants (SNVs) that define WA1 (C8782T, C18060T, and T26144C), the sequences from cruise ship passengers and crew also shared two additional SNVs, C17747T and A17858G, common to nearly all WA1 sequences sampled from Washington and California but not the basal WA1 case (Figs. 2B and 3).

The WA1 case was reported on 19 January (6) and thus substantially predated the voyages of the Grand Princess cruise ship (7, 20). In addition, six of eight passengers on voyage B (UC7 to UC11 and UC30) each carried at least two new mutations (G16975T and C23185T) not observed in UC1, UC19, and UC20, who were all on voyage A (Fig. 3). This suggested that the virus from UC19 could be basally positioned relative to the cruise ship strains from voyage B, and that COVID-19 infections associated with voyage A may have been passed on to passengers and crew on voyage B. However, because of sequencing artifacts from areas of low coverage, the initial WA1 subtree extracted from the global maximum likelihood phylogenetic tree did not place UC19 basal to...
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A

WA1 lineage
Santa Clara County cluster
Solano County cluster
San Benito County
Northern California, European / New York lineages
Northern California, other lineage
Other California (outside of this study)
Other US
European / New York lineages
G clade

B

WA1 Lineage
Santa Clara County (SCC1) Lineage

G clade (European / New York lineages with the D614G spike mutation)

D

Solano County Cluster

C

Santa Clara County (SCC1) Lineage
sequences from voyage B passengers (fig. S2). To establish a more accurate tree topology, we therefore reconstructed a new phylogenetic subtree of the WA1 lineage after excluding all ambiguous sites. In this new subtree (Fig. 2B), UC19 is basal to all other California genomes within the WA1 lineage. In addition, among the sequences from patients on voyage B, UC5 and UC6 group together, whereas UC7 to UC11 and UC30 group together with a sequence sampled in Minnesota.

The chronology and phylogeny of the cruise ship outbreak, and the predominance of the WA1 lineage in Washington state (7), together suggest that the virus on the Grand Princess likely came from Washington, although the cases may also have originated from a different region in which the WA1 strain is circulating. In addition to passengers and crew members aboard the Grand Princess, virus genomes sampled from three cases of community transmission in different counties of the San Francisco Bay Area (UC22, UC23, and UC28) were also of the WA1 lineage. UC22 was the son of an infected Grand Princess passenger (UC20) on voyage A and most likely contracted the virus from household contact. The UC23 and UC28 cases may also reflect transmission from disembarking Grand Princess passengers on voyage A, or preexisting circulation of the WA1 strain in the community.

Three patients examined in this study (CA3, CA4, and UC12) had COVID-19 infections associated with international travel or exposure to international travelers. CA3 corresponds to a resident of San Benito County who became sick shortly after returning from Wuhan, China in late January. The sequence of his SARS-CoV-2 genome is identical to that of CA4, a household contact who was also infected with the virus. Their viral genomes were found to be closely related to early lineages from China (Fig. 2A and data S1). UC12 had a prolonged exposure to a known positive traveler from Switzerland while attending a conference. The genome from UC12 fell within a lineage containing many sequences from European residents or travelers from Europe (Fig. 2A). Interestingly, four additional genomes (UC24, UC26, UC27, and UC36) were also grouped within the European lineage. UC27 and UC36 were both diagnosed shortly after returning to California from New York, consistent with reports that the New York outbreak that began in March 2020 originated with travelers coming from Europe (21, 22). UC26 also reported domestic travel from Los Angeles, whereas UC24 had no known travel history.

In Santa Clara County, we sequenced seven genomes from individuals who were part of a local outbreak of COVID-19 at a large workplace facility with multiple employers, large areas of shared space, and heavy pedestrian traffic. The genomes all shared the G2971T SNV that defines the SCC1 lineage (Figs. 2C and 3). Four employees (UC13, UC14, UC15, and UC34) had dates of symptom onset within 2 weeks of each other, although they did not know each other. UC6 and UC17 were family members of UC13 and lived in the same residence, while UC35 transported UC14 to the hospital via emergency medical services. Notably, the genomes from a Solano County resident (UC21) and a San Mateo County couple (UC18 and UC25) were also placed in the SCC1 lineage, suggesting possible spread to different counties. Further epidemiological investigation found that UC21 had visited a merchant in Santa Clara County, during which he likely became infected.

In Solano County, a small cluster of three cases included the first reported instance of community transmission in the United States on 26 February (UC4) (Figs. 2D and 3). The two other cases (UC2 and UC3) were health care workers who were taking care of patient UC4 and likely contracted the disease in the hospital, consistent with transmission of the disease from patients to health care providers (23). The genomic epidemiology of the COVID-19 cases associated with community spread studied here do not show any predominant SARS-CoV-2 lineage circulating in Northern California. In California, multiple recent and unrelated introductions of SARS-CoV-2 into the state via different routes appear to have given rise to the diversity of virus lineages reported in this study, with no single predominant lineage observed.

We note that this does not exclude the possibility of cryptic transmission of multiple lineages in California simultaneously, as the current level of sampling is not dense enough to confidently estimate the dates of the seeding events, nor the subsequent periods of cryptic transmission before a lineage was identified. There is growing evidence that WA1 is now an established lineage of SARS-CoV-2 in the United States. Here, we found WA1-lineage viruses from Grand Princess cruise ship passengers as well as from residents of several Northern California counties. In addition, WA1-lineage viruses have been identified in COVID-19 cases from many states including Minnesota, Connecticut, Utah, Virginia, and New York (24, 25). The early date and basal phylogenetic position of the WA1 virus make it likely that the direction of dissemination was from Washington state to California and other states. Notably, SARS-CoV-2 sequences from Connecticut (25) and British Columbia, Canada (Fig. 2B) are positioned close to the root of the subtree containing the WA1 sequences, raising the possibility that the virus may not have been first introduced into the United States via Washington state.

SARS-CoV-2, like other coronaviruses, contains a nonstructural gene with proofreading activity (26). Consequently, the virus evolves more slowly than many other human RNA viruses, on the order of one to two DNA base substitutions per month across its ∼29-kb genome (27). Thus, only one to three SNVs in general are needed to define a distinct lineage. The WA1 lineage consists of three key SNVs (G2972T, C18060T, and T28144C), whereas the SCC1 lineage associated with the Santa Clara County cluster and the Solano County cluster are each defined by only one SNV (G2971T and C9924T), respectively (Figs. 2 and 3).

Our epidemiological and genomic survey of SARS-CoV-2 has several limitations. First, this initial analysis represents a relatively sparse sampling of cases. Undersampling of virus genomes is due in part to the high proportion of cases (~80%) with asymptomatic or mild disease who do not get tested (2–4). Second, the majority of samples analyzed were obtained from public health laboratories and thus may not be representative of the general population. Finally, phylogenetic grouping of viruses from different locations, such as Washington state and California in the same WA1 lineage, does not prove the directionality of spread. Despite this, our study shows that robust insights into COVID-19 transmission are achievable if virus genomic diversity is combined and jointly interpreted with detailed epidemiological
Fig. 3. Multiple sequence alignment of all SARS-CoV-2 genomes reported across nine Northern California counties and the Grand Princess cruise ship. SNVs with respect to the reference genome (NC_045512) are shown as vertical red and black lines for lineage-defining SNVs and other SNVs, respectively. Cases that are part of the WA1 lineage include the first case of COVID-19 infection (WA1) in the United States, eight passengers and crew members aboard the Grand Princess cruise ship during its second trip (voyage B), and three individuals surveyed from three Northern California counties as passengers on the ship’s first trip (voyage A). The three SNVs C8782T, C18060T, and T28144C define the WA1 lineage, and the two SNVs C17747T and A17858G are common to Grand Princess passengers and crew. Viruses from voyage B passengers and crew share SNVs G16975T and C23185T that are lacking in viruses from voyage A passengers. Single SNV variants C9924T and G29711T define the lineages from Solano County and Santa Clara County, respectively. European lineages share SNV A23403G. The putative epidemiological link and sample collection date are shown beside each sequence alignment.
case data. In particular, we found that a returning traveler from New York was infected with a lineage circulating widely in Europe, thus suggesting an association between the New York outbreak and intercontinental travel to and from Europe before this was widely recognized (21, 22).

Public health containment measures such as isolation and contact tracing, as performed in the Solano County and Santa Clara County outbreak clusters, become more difficult to maintain once a lineage becomes established in the community. Our data suggest trends in this direction, such as the association between the WA1 lineage and community-acquired COVID-19 cases in several counties of Northern California, and the detection of a virus from the SCC1 lineage in residents of neighboring San Mateo and Solano counties. Social distancing interventions, such as the “shelter-in-place” directive that was issued by the governor of California on 20 March 2020, may have assisted in stemming spread from community to community. Inter-state dissemination of SARS-CoV-2 lineages has also been demonstrated coast-to-coast between Washington state and Connecticut (25), and from domestic and international travel into the Bay Area in the current study. Suspension of non-essential travel may help to prevent importation of new cases in California and other states.

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SUPPLEMENTARY MATERIALS

science.sciencemag.org/content/369/6503/582/suppl/DC1

Materials and Methods

Fig. S1 and S2

Tables S1 to S5

Materials and Methods

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(31–39)

View/request a protocol for this paper from Bio-protocol.

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Genomic surveillance reveals multiple introductions of SARS-CoV-2 into Northern California


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**Epidemic in Northern California**

Genome sequencing of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) outbreaks is valuable for tracing the sources and perhaps for drawing lessons about preventing future outbreaks. Genomic analysis by Deng *et al.* revealed that Northern California experienced a complex series of introductions of the virus, deriving not only from state-to-state transmission but also from international travel by air and ship. The study highlights the importance of being able to rapidly test and trace contacts of positive cases to enable swift control.

*Science*, this issue p. 582
Supplementary Materials for

Genomic surveillance reveals multiple introductions of SARS-CoV-2 into Northern California


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This PDF file includes:
Materials and Methods
Figs. S1 and S2
Captions for tables S1 to S5
Caption for data S1
References

Other supplementary material for this manuscript includes:
Tables S1 to S5 (Excel format)
Data S1 (.tree format)
Materials and Methods

Quantitative RT-PCR testing

Quantitative reverse transcription-polymerase chain reaction (RT-PCR) testing for SARS-CoV2 was performed by the US Centers for Disease Control and Prevention (CDC), California Department of Public Health (CDPH), University of California, San Francisco Clinical Microbiology Laboratory (UCSF), or Santa Clara County Public Health Department (SCCPHD). At UCSF, samples were extracted on a Magna Pure 24 Viral Kit (Roche Diagnostics, Indianapolis, USA) or an EZ1 Viral Mini Kit (Qiagen, Hilden, Germany). At the CDC, CDPH, or SCCPHD, samples were extracted using Qiagen DSP Viral RNA Mini Kit with carrier RNA added (Qiagen). Samples were tested for SARS-CoV-2 using the FDA EUA-approved 2019-nCoV CDC Real-Time RT-PCR Diagnostic Panel assay (16), which targets the N1 and N2 regions of the nucleoprotein gene. The cutoff for a confirmed positive sample was determined to be a cycle threshold (Ct) value of 40 cycles, with a positive result requiring detection of both the N1 and N2 targeted regions of the nucleoprotein gene.

Clinical SARS-CoV-2 samples

Nasopharyngeal and/or oropharyngeal swabs in universal transport media (Copan Diagnostics, Murrieta, CA, USA) from RT-PCR positive COVID-19 patients were obtained from the UCSF Clinical Microbiology Laboratory, CDPH, and SCCPHD. Patient samples were randomly selected for broad representation of the 9 counties in Northern California and the Grand Princess cruise ship.
Next-generation sequencing library preparation and PCR confirmation of the G29711T SNV

Extracted RNA from nasopharyngeal (NP) swab samples in universal transport medium was reverse transcribed to complementary DNA (cDNA) using the MSSPE method as previously described (15). The custom-designed 13-nucleotide (nt) SARS-CoV-2 primers (IDT Technologies) were constructed using an alignment of 30 SARS-CoV-2 genome references available in the NCBI GenBank database as of end of February 2020 (Table S3). Barcoded sequencing libraries were constructed from cDNA by Nextera tagmentation (Ilumina), followed by 12 cycles of amplification per the manufacturer’s protocol. A second round of amplification (14 cycles) was used to boost the yield of cDNA library due to low RNA yield in general from low-input samples such as NP swabs (33). The final libraries were pooled and sequenced on the MiSeq, NextSeq, or HiSeq 1500 (Illumina Inc., San Diego, USA) as 1x150 single-end or 2x150 paired-end reads. For some genomes with low coverage (<50%), tiling amplicon PCR using SARS-CoV-2 primers generated from the Primal algorithm (Table S4) was used to increase genome coverage according to the published protocol (17). To confirm that UC15 and UC25 belong to the SCC1 lineage, we performed specific PCR using designed primers to amplify a region containing the lineage defining G29711T SNV, followed by Sanger sequencing to confirm the presence of this SNV (Table S5).

Phylogenetic analysis

Raw reads were first screened via BLASTn (BLAST+ package 2.9.0) (34) for alignment to SARS-CoV-2 reference genome NC_045512. They were then aligned to the reference genome with LASTZ version 1.04.03. For libraries generated using the MSSPE protocol (15), single-end
reads were trimmed using Geneious version 11.1.3 by removal of 13 nucleotides (nt) (the length of the MSSPE primer) and low-quality reads from the ends, followed by removal of duplicate reads. Trimmed reads were then mapped to reference genome NC_045512 in Geneious with no gaps allowed and a maximum of 5% mismatches per read. The assembled contig was then manually annotated, and a consensus genome was generated using a majority threshold criterion. For libraries generated using tiling multiplex PCR (Primal) protocol (17), paired-end reads were processed using the BBTools suite, version 38.82 (https://sourceforge.net/projects/bbmap/). Quality scores were recalibrated, and reads were trimmed for removal of adapters and low-quality sequences using BBDuk. Subsequently, mate pairs were aligned with BBMap, and duplicate pairs removed on the basis of mapping to the same nucleotide positions. Mapped reads containing deletion events or multiple (>2) substitution events unsupported by any other read were removed to reduce overall noise, and soft clipping was applied to reduce the impact specifically of artifactual noise near read ends. Variants were called with CallVariants and applied to the genome using a depth cutoff of 3 to yield the final assembly. Regions with depth of less than 3 were represented by N.

We downloaded all 762 complete (>29,000 bp), high-coverage SARS-CoV-2 genomes that had been deposited into GISAID as of March 20, 2020 (31, 32), and then added the 36 genomes in the current study to generate an initial dataset of 798 genomes. This dataset was then trimmed by removal of low-quality genomes and known duplicate genomes from the same patient, yielding a final dataset of 789 genomes. Sequences were aligned using MAFFT v7.427 (35) using default settings and multiple sequence alignments were manually curated for accuracy. Phylogenetic maximum likelihood trees were constructed in PhyML v3.3 (36) under an HKY+Γ₄ substitution model (37, 38), after trimming the ends of the alignment. To compute
branch support values we used the aLRT (approximate likelihood ratio test) method, implemented in PhyML (39), which is better suited than bootstrapping to data sets with few informative sites per taxon.

For the separate WA1 lineage phylogenetic analysis in Figure 2B, we discarded all genomes of the WA1 lineage, leaving 88 genomes. We further removed 4773 nucleotide sites from the alignment that were either ambiguous or unknown in more than one genome. As above, the phylogenetic WA1 lineage subtree was constructed in PhyML v3.3 under an HKY+$\Gamma_4$ substitution model. The locations of SNVs and gaps shown in Figure 3 were extracted from an alignment of 154 sequences from the US, aligned to the reference sequence (NC_045512), using custom scripts (30).
Fig. S1. SARS-CoV2 genome coverage versus cycle threshold for respiratory swab samples from COVID-19 patients (n=34) in this study. The x-axis designates the virus titer based on the reported real-time RT-PCR Ct value (higher values correspond to lower viral loads), while the y-axis designates genome coverage in percentage. The data points are color coded to designate either the patient’s county of residence in Northern California or whether the patient was a passenger or crew member aboard the Grand Princess cruise ship.
Fig. S2. Initial phylogeny of the WA1 lineage pruned from the global phylogenetic tree of 789 SARS-CoV-2 genomes. Light blue circles denote the genome sequences obtained in this study, while gray circles denote sequences from Washington State (WA). The 3 key SNVs defining the WA1 lineage are highlighted in red, while an additional 2 SNVs found in viral genomes from the Grand Princess passengers and crew as well as the majority of the WA1 lineage viruses are highlighted in black. The WA1 virus from Washington State (first reported COVID-19 case in the US) is positioned at the root of the tree and is closely related to Fujian and Chongqing viruses from China. Bootstrap values (converted from the approximate likelihood ratio test, or aLRT score) are displayed at each node, with a value of 1 indicating 100% support.
Table S1. (separate file in Excel format) List of all 62 SARS-CoV-2 samples analyzed in the study along with sequencing controls.

Table S2. (separate file in Excel format) Patient epidemiologic metadata and sequencing metrics for 36 SARS-CoV-2 genomes recovered from COVID-19 infected patients.

Table S3. (separate file in Excel format) Custom primer sequences for the MSSPE method.

Table S4. (separate file in Excel format) Tiling multiplex PCR primers.

Table S5. (separate file in Excel format) PCR primers and Sanger sequencing reads obtained for confirmation of SNV G29711T from SARS-CoV-2 respiratory nasal swab samples.

Data S1. (separate file in Nexus tree format) Tree file for the global phylogeny of 789 SARS-CoV-2 genomes.
References


