Genomic epidemiology and transmission dynamics of SARS-CoV-2 in congregate healthcare facilities in Santa Clara County, California

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Summary: Genomic sequences from 196 staff and residents of LTCFs in Santa Clara County, California characterized transmission of SARS-CoV-2 in congregate settings. Timely pairing of epidemiologic and clinical data with genomic sequencing is an important tool to refine SARS-CoV-2 mitigation strategies.
**Background:** Outbreaks of SARS-CoV-2 in long term care facilities (LTCFs) cause significant morbidity and mortality. Mapping viral transmission within and between by combining genomic sequencing with epidemiologic investigations enables targeting infection control interventions.

**Methods:** We conducted weekly surveillance of residents and staff in LTCFs in Santa Clara County, CA with at least one confirmed COVID-19 case between March and July 2020. Positive samples were referred for whole genome sequencing. Epidemiological investigations and phylogenetic analyses of the largest outbreaks (>30 cases) were carried out in six LTCFs (Facilities A through F).

**Results:** Among the 61 LTCFs in the county, 41 had at least one confirmed case during the study period, triggering weekly SARS-CoV-2 testing. The six largest outbreaks accounted for 60% of cases and 90% of deaths in LTCFs, though the bed capacity of these facilities represents only 11% of the LTCF beds in the county. Phylogenetic analysis of 196 whole genome sequences recovered from those facilities showed that each outbreak was monophyletic, with staff and residents sharing a common viral lineage. Outbreak investigations revealed that infected staff members often worked at multiple facilities, and in one instance, a staff member infected while working in one facility was the likely index case in another.

**Conclusions:** We detected a pattern of rapid and sustained transmission after a single introduction of SARS-CoV-2 in six large LTCF outbreaks, with staff playing a key role in transmission within and between facilities. Infection control, testing, and occupational policies to reduce exposure and transmission risk for staff are essential components to keeping facility residents safe.

Keywords: genomics; epidemiology; elderly; healthcare; Providers
Introduction

Santa Clara County, an ethnically and economically diverse region of California also known as “Silicon Valley,” had an early introduction of the SARS-CoV-2 novel coronavirus\(^1\). Its first case of COVID-19 was confirmed in a traveler on January 31, 2020, and subsequent cases were introduced by multiple routes including travel from Wuhan, China, passengers from the Diamond and Grand Princess cruises, and travel from Washington state\(^2,3\). The first community-acquired case of SARS-CoV-2 in Santa Clara County (SCC) was announced on February 28, and within a week the virus was found to have spread from the community into long-term care facilities (LTCF), including skilled nursing facilities (SNFs), resulting in 4 deaths among 419 cases by April 30\(^4\). Residents of these congregate healthcare facilities are particularly prone to poor outcomes including mortality because of advanced age and comorbidities such as hypertension, diabetes, and chronic respiratory conditions. LTCFs also have the potential for rapid and sustained viral transmission due to shared living quarters and close contact between residents and care providers.

To reduce the potential for outbreaks of COVID-19 in LTCFs, the County of Santa Clara Public Health Department (SCCPHD), with assistance from the Centers for Disease Control and Prevention (CDC) provided facilities with infection control guidance and checklists, increased access to personal protective equipment, and established policies prohibiting family visitation and group activities. Beginning April 1, the County introduced response-driven testing in LTCFs, where a suspected cluster of COVID-19 cases triggered, at a minimum, weekly facility-wide testing of residents and staff. Positive PCR specimens were submitted for whole genome sequencing.
The use of genomics as a complement to traditional epidemiologic investigations and surveillance has been well established in global health contexts, with successful investigations and responses to outbreaks caused by Ebola, Zika, and seasonal influenza A/B viruses \(^5\text{--}^8\). Several studies have successfully used genomic epidemiology to describe SARS-CoV-2 transmission in hospitals and nursing homes in the United Kingdom and the US \(^9^,\text{10}\). Viral genome sequencing can provide independent and complementary evidence to contact tracing, which is especially useful in congregate healthcare settings where complex patterns of staff and resident movement within and between facilities create competing hypotheses for transmission routes.

Here, we characterize the outbreak transmission patterns of SARS-CoV-2 between LTCF residents and staff across LTCFs in Santa Clara County from March 18 to July 31, 2020. We demonstrate how pairing genomic sequencing of SARS-CoV-2 samples with contact tracing and epidemiologic data revealed relationships between individuals and facilities, and how this additional evidence informed infection control recommendations and public health decision making.

**Methods**

**Testing and Screening Strategy**

Beginning April 1, 2020, facility-wide testing of staff and residents was coordinated by the SCCPHD and LTCF management when clusters of three or more suspected cases or one confirmed case of COVID-19 among staff or residents were detected in any LTCF, which included licensed skilled nursing, intermediate care, and assisted living facilities. As first cases were detected in each facility, the health department engaged in site visits and facility-wide PCR screening (i.e., Response-Driven testing). Nasopharyngeal swabs for all employees and residents were collected weekly for SARS-
CoV-2 testing by the SCCPHD laboratory (with a transition to nasal swabs on June 18). At a minimum, weekly screening was sustained until no new cases were detected. Visitors and non-essential staff were excluded from LTCFs and were not included in testing. Outbreaks were declared contained in the absence of new suspected or confirmed cases after two consecutive weeks. Standardized case line lists contained demographics, clinical data and outcomes, and resident room assignment.

We analyzed all confirmed cases of SARS-CoV-2 detected at 61 LTCFs from March 18 – July 31, 2020. Based on the CDC definition, cases of COVD-19 were confirmed by the presence of SARS-CoV-2 RNA as detected by a quantitative molecular amplification detection test (RT-PCR). Whole genome sequencing on PCR positive samples from SCC started April 1, 2020. We focused our detailed genomic analysis to six LTCF outbreaks (Facilities A to F), each with a minimum of 30 cases of COVID-19.

**Sequencing and Bioinformatics**

All respiratory samples from LTCFs were processed for RT-PCR testing at the SCCPHD Laboratory using the CDC protocol for the detection and amplification of SARS-CoV-2. Positive samples with a cycle threshold (Ct) ≤32 were forwarded to the Chan Zuckerberg (CZ) Biohub for genomic sequencing via a modified version of the Primal-Seq Nextera XT version 2.0 protocol, using ARTIC Network V3 primers, and paired-end 2x150bp sequencing on Illumina NovaSeq. Consensus sequences were obtained using MN908947.3 as reference, using minimap2, samtools, and ivar in the CZ Biohub consensus genome pipeline (https://github.com/czbiohub/sc2-illumina-pipeline). Viral genomes with at least 90% coverage were uploaded to GISAID.
Phylogenetic assemblies from SCCPHD were constructed using iqtree in the augur pipeline with default settings. Phylogenetic results, paired with demographic and epidemiologic data, were visualized in NextStrain and displayed using baltic (https://github.com/evogytis/baltic). Non-study samples were downsampled by a factor of 3 after tree construction for ease of display.

**Communication to Facilities**

Positive PCR results were communicated to the facility, with an average turnaround time of 48 hours. Efforts to minimize the turnaround time for PCR results and genomic data translated into actionable information. Where applicable, phylogenetic relationships were confidentially communicated to SCCPHD and LTCF management to educate and reinforce infection control messaging.

**Non-Research Public Health Activity Adjudication**

The representative for the Department of Public Health of the Health Services Institutional Review Board (IRB) of the County of Santa Clara reviewed this activity and deemed it to be a public health surveillance activity, granting it a non-research activity determination that did not require IRB review.

**Results**

**Outbreak Characteristics**

The first COVID-19 LTCF outbreaks occurred in mid-March 2020, prompting weekly testing in 41 sites by the end of July 2020. We focused our analysis on the largest outbreaks which occurred at six LTCFs. Facilities A through F comprised 64.5% of all cases (491/761) and 90.0% (72/80) of all deaths.
among all LTCFs in the county, despite comprising just 11.3% (761/6722) of the total bed capacity. These facilities included four large SNFs and two assisted living facilities with memory care services. Table 1 describes the characteristics of their staff (median age 47 years) and residents (median age 81 years). Their demographics were comparable to other facilities in the county (data not shown). Moreover, among the Centers for Medicare and Medicaid Services (CMS)-certified SNFs in the county, outbreak size was not statistically associated with facility quality rating or minutes of care per day by nurses or by aides (Spearman correlations ρ=-0.12-0.15, p=0.38-0.76). Likewise, quality ratings among the four SNFs with the largest outbreaks included the highest (n=1), lowest (n=1) and average (n=2) facility quality scores.

The large outbreaks came in two waves. The initial cases from the four SNF outbreaks in March 2020 (Table 1, Facilities A-D) were detected in symptomatic staff. These outbreaks arose early in the pandemic when treatment, diagnostic, and PPE constraints were significant. The next wave included two large outbreaks, both detected in June, in assisted living/memory care centers, with the first cases also identified in staff. Table 2 outlines the timeline and scale of the outbreaks with the recovery of viral genomes from staff and residents. On average, 40.7% of residents as a percent of facility capacity (Range: 12.5%-79.6%) tested positive across the six facilities, while outbreaks in other county facilities were limited to less than 10% of their capacity (data not shown). All 239 PCR-positive samples for SARS-CoV-2 with a Ct value of no greater than 32 were submitted for whole-genome sequencing (WGS), and near-complete (>90%) viral genomes were recovered for 196 samples, representing 50 sequences from staff (27.9% of known staff cases) and 146 from residents (46.8% of known resident cases).
Phylogenetic Analysis

We constructed a phylogenetic tree consisting of the 196 SARS-CoV-2 genomes from the LTCF facilities investigated together with 509 additional genomes from residents of Santa Clara County diagnosed with COVID-19, to understand the relationship between the outbreaks and other circulating genotypes (Figure 1). The additional genomes include COVID-19 cases from hospitals, community clinics, public testing sites, and jails. We found that Santa Clara County contained all the major global clades circulating at the time, as defined by NextStrain (19A, 19B, 20A, 20B, 20C). Each SNF outbreak (Facilities A-D) formed a separate, monophyletic cluster on the tree. Within each outbreak, a plurality of samples (42.3% on average) shared a single genotype ancestral to all other genotypes recovered from the outbreak, consistent with an introduction by an index case followed by rapid spread (Table 2). Each outbreak lineage is defined by a small number (1-4) of single nucleotide mutations (denoted as single nucleotide variants or SNV) relative to the root Wuhan-Hu-1 genotype, which is their earliest common ancestor (Facility A: C10277T, Facility B: C25692T, Facility C: A14940G, Facility D: C25916T). Interestingly, all sequences belong to the 19A clade, and none of the genomic lineages found at Facilities A through D were of the SCC1 lineage (defined by SNV, G29711T) which defined early community spread within SCCPHD, nor the WA1 lineages characterizing the Grand Princess cruises. The outbreaks in both assisted living facility outbreaks were also monophyletic (Facilities E and F). Notably, the Facility F outbreak lineage was a descendent of the Facility E outbreak lineage: Facility F genomes contained the G7960T mutation associated with the Facility E outbreak as well as an additional A14827G mutation. This genetic connection between the outbreaks was further supported by epidemiological data described below.

Genetic analysis uncovered further instances of staff sharing between facilities. While 37 of the 50 staff genomes (74%) recovered matched the lineage associated with the facility in which the staff
member was tested, 13 of the genomes (26%) were found during response driven testing at a different facility. Further case investigation revealed that each of the 13 staff also worked at the facility where their matching genotype was in circulation, and those cases are colored in Figure 1 according to the latter facility. Even in this small sample, those links suggest transmission opportunities between facilities by infected staff.

Transmission Kinetics of SARS-CoV-2

To better understand the kinetics of transmission in LTCFs we studied the genetic and epidemiologic relationships between residents and staff. Figure 2 shows the viral genomes of cases among residents (orange) and staff (blue) from Facilities A and B. Samples from other LTCFs (pink) and the community (dark grey) are also included as reference. Thirteen genomes, from the earliest cases at Facility A, have an identical genotype. Subsequent mutations define limited onward transmission within the facility. One such transmission chain (Figure 2A, red box) contains a cluster of four residents who were roommates or across a corridor from one another. They were transferred to a COVID-19 isolation wing after receiving positive PCR results. Two weeks later, a downstream genotype containing one additional SNV was detected in a staff member.

The red box in Figure 2b contains 10 identical genotypes from 9 residents and 1 staff who were tested at Facility B. All of these residents were originally co-located along one wing (e.g., shared rooms and restrooms, across the hallway). This one geographic hotspot within the facility appeared to give rise to genotypically identical cases for six weeks from late March to early May.
We also investigated the association between Facilities E and F suggested by the genomic data (Figure 3). One asymptomatic staff member working at both Facilities E and F had a plausible role in introducing COVID-19 to Facility F (red arrow). This individual had a positive PCR result at Facility E in late June, and was also the earliest identified case from the outbreak at Facility F. The genome recovered from this case had one SNV distinguishing it from the genotype circulating in Facility E and that was subsequently present in all viral genomes recovered from the 59 cases at Facility F. Two asymptomatic certified nursing assistants (CNAs) who shared a household, worked in at least three LTCFs including Facilities E and F. The viral genotypes sequenced from these individuals were identical and matched the dominant genotype associated with Facility E. In this circumstance, transmission was likely associated with the shared household. There was no evidence to indicate that either staff transmitted the virus to others at any of the multiple sites they were employed. Rapid intervention, exclusion from work, and home isolation may have prevented further transmission into the multiple facilities where these providers worked.

**Staff-Resident Interactions**

We also sought to use the genomic data to clarify whether transmission was staff-to-staff, staff-to-resident, or resident-to-resident. Across 146 recovered resident genomes, we identified 10 pairs of resident roommates with identical genomes, and 11 pairs of roommates with unmatched genomes differing by at least 1 SNV (consistent with unidentified intermediaries). While roommates share close proximity and an increased likelihood of receiving care from the same staff, other breaches in infection control could present opportunities for transmission. Of the 50 genomes obtained from healthcare workers across the six outbreaks, 26 (54.2%) were from CNAs, who perform duties such
as feeding, bathing, and toileting activities, involving close or face-to-face contact increasing the opportunity for person to person transmission.

With shelter-in-place orders imposed since March 13, 2020, non-essential care or elective wellness activities were all but discontinued in SNFs, and residents were unable to socialize or circulate as they may have chosen to do normally. While the direction of transmission between residents and staff can be challenging to ascertain, with residents largely confined to their living quarters, transmission was likely amplified by healthcare worker intermediaries.

**Discussion**

Our investigation, spanning months of active, response-driven PCR testing for SARS-CoV-2, revealed a consistent pattern of transmission driving large LTCF outbreaks in Santa Clara County. In each case, a single introduction was followed by rapid proliferation, with the same SARS-CoV-2 lineage persisting for weeks despite infection prevention efforts. Healthcare providers were infected by the same viral lineage circulating among their patients. While the direction of transmission cannot be established from this analysis, it reinforces the need for setting-specific and real-time infection control guidance for staff, visitors, and residents.

CNAs represent a majority of caregiver roles in LTCFs who provide essential face-to-face patient care services. CNAs often care for multiple residents, sometimes exceeding the recommended ratios, and are less likely to receive comprehensive and refresher infection control training, compared with licensed nursing staff. These factors can elevate transmission risks among CNAs and similar caregiver types. Other investigations have identified comparably credentialed personnel in the UK
and US to be similarly vulnerable\textsuperscript{20,21}. It is prudent to enhance the infection control training offered to these provider groups.

In the United States, it is common for staff to work in multiple LTCFs, and a worker exposed in one facility may bring the virus to another.\textsuperscript{20,22} In March 2020, the CDC identified staff members working in multiple nursing homes as a likely source of early spread in Washington state\textsuperscript{2}, and recent work of Chen et al. used smartphone data to show a correlation in cross-staff movement and outbreak size across US nursing homes\textsuperscript{20}. Genomics can validate such observation and statistical data, and our study provides evidence of staff acting as a transmission link between facilities.

To curb the likelihood of onward transmission within and across healthcare facilities, in-depth contact tracing for healthcare providers may be beneficial. Screening all household and close contacts of positive healthcare providers could provide insight into the intermediaries between community- and healthcare-associated transmission.

Transitioning WGS from research into applied public health requires rapid return of phylogenetic results from clinical samples, ideally within 3 to 5 days\textsuperscript{9}. In our experience, creating an end-to-end process from clinical sample to an assembled phylogenetic tree within one business week required multiple stakeholder commitments, including: standardized processes moving samples between testing and sequencing labs, batched sequencing workflows to control costs, agile analysis methods, and frequent communication. Providing visualizations of phylogenetic data to public health and facility management became an infection control tool, illustrating, in real-time, targets to break transmission.
While it is challenging to balance the social and behavioral needs of SNF residents under pandemic restrictions, the exclusion of community, family, visitors, and non-essential support services may have prevented multiple introductions into facilities, which could amplify already escalating case counts and resource constraints. However, the necessity of staff entry and interaction with residents creates a baseline level of transmission risk, making appropriate training, testing, cohorting, and support for staff essential components of infection control. Going forward, it is important to continue genomic surveillance in light of SARS-CoV-2 antigenic evolution, waning natural immunity, and shifts in transmission patterns.
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Figure Legends

FIGURE 1. Six major SARS-CoV-2 outbreaks in congregate senior healthcare facilities in Santa Clara County are monophyletic. Shown is a maximum-likelihood phylogenetic tree built from 706 viral genomes from Santa Clara County samples collected between February and July 2020. Resident and staff samples from each facility are drawn as dots colored by the facility at which the individual was tested or worked (n=196). Dashed boxes contain all nodes descended from the index genotype of each facility, defined as the common ancestor of all genotypes sampled from residents at that facility (incoming branches are labelled with the defining SNV). Nextstrain clades are labelled 19A-20C. All staff and resident genomes from each facility descend from the index genotype, consistent with a single introduction. Grey nodes (n=16) indicate contextual samples which are also descended from facility index genotypes, representing potential onward transmission.

FIGURE 2. Genetic linkages between resident and staff within each facility. Insets of the tree in Figure 1 showing staff and residents from Facilities A (panel A) and B (panel B) together with samples from elsewhere descended from the index genotype. Instances where clusters of staff and residents share additional mutations beyond the index genotype of each outbreak indicate repeated transmission between those groups, and instances where external samples also share those mutations indicate likely onward transmission from the facility back into the community. Red boxes indicate clusters with epidemiological relationships discussed in the text.

FIGURE 3. Shared staff seed outbreak. The index case of the outbreak at Facility F (red arrow) was a staff member also working at Facility E. The genomic diversity of Facility F is nested entirely inside that of Facility E. Red asterisks indicate two additional CNAs who worked at both facilities and share a household.
## Tables

### Table 1: Demographic characteristics across six congregate settings, March 2020-July 2020

|                      | Facility A | Facility B | Facility C | Facility D | Facility E | Facility F | Totals |
|----------------------|------------|------------|------------|------------|------------|------------|--------|
| **LTCF Capacity**    | 199        | 99         | 201        | 59         | 104        | 104        | 766    |
| **LTCF Type**        | SNF        | SNF        | SNF        | SNF        | AL/MC      | AL/MC      | N/A    |
| **Confirmed Cases**  | 151        | 64         | 126        | 61         | 30         | 59         | 491    |
| **Sex, female (% all cases)** | 85 (56.3)  | 40 (62.5)  | 77 (61.1)  | 46 (75.4)  | 25 (83.3)  | 44 (74.6)  | 317 (64.6) |
| **Resident Age (median, yrs)** | 75.5       | 79         | 76         | 87         | 84         | 90         | 81     |
| **Staff Age (median, yrs)** | 48         | 53         | 40.5       | 52         | 48         | 47.5       | 47     |
| **COVID Severity**   |            |            |            |            |            |            |        |
| **Hospitalization**  | 44         | 5          | 11         | 10         | 0          | 1          | 71     |
| ICU Admission | 8 | 1 | 23 | 3 | 6 | 5 | 46 |
|---------------|---|---|----|---|---|---|----|
| Deaths        | 27| 10| 11 | 11| 5 | 8 | 72 |
Table 2: SNF Outbreak Characteristics and Outcomes, March 2020-July 2020

| Outbreak Characteristics | Facility A | Facility B | Facility C | Facility D | Facility E | Facility F | Total |
|--------------------------|------------|------------|------------|------------|------------|------------|-------|
| Outbreak onset (earliest confirmed positive PCR) | March Week 3 | March, Week 1 | March, Week 2 | June, Week 1 | June, Week 3 | N/A |       |
| Days to Outbreak peak | 22 | 19 | 36 | 27 | 13 | 5 | 20.3* |
| Days to Last Case | 54 | 38 | 109 | 47 | 18 | 30 | 49.3* |
| COVID-19 positive staff | 49 | 17 | 54 | 18 | 17 | 24 | 179 |
| COVID-19 positive residents | 102 | 47 | 72 | 43 | 13 | 35 | 312 |
| Residents % positive | 51.3 | 47.5 | 79.6 | 35.8 | 12.5 | 33.7 | 40.7 |
| Sequences Recovered | 47 | 33 | 32 | 25 | 19 | 40 | 196 |
| Staff Sequences (%) known cases | 9 (18.4) | 8 (47.1) | 8 (14.8) | 1 (5.6) | 10 (58.8) | 14 (58.3) | 50 (27.9) |
| Residents Sequences | (% known cases) | 38 (37.3) | 25 (53.2) | 24 (33.3) | 24 (55.8) | 9 (69.2) | 26 (74.3) | 146 (46.8) |
|---------------------|----------------|-----------|-----------|-----------|-----------|----------|----------|-----------|
| Sequences with      |                |           |           |           |           |          |          |           |
| index genotype (% of WGS) |            | 15 (31.9) | 4 (12.1)  | 19 (59.4) | 8 (32.0)  | 13 (68.4) | 26 (65.0) | 83 (42.3) |

Note: For SNFs (Facilities A-D), average bed capacity based on Centers for Medicare and Medicaid Services, 2019 licensing data (medicare.gov/nursinghomecompare), For Assisted Living Facilities, total bed capacity was used as the denominator

*Average time to outbreak peak/resolution in days
