Original Article

Unexpected details regarding nosocomial transmission revealed by whole-genome sequencing of severe acute respiratory coronavirus virus 2 (SARS-CoV-2)

Sofia Myhrman MD1,a, Josefin Olausson PhD2,a, Johan Ringlander MD2, Linéa Gustavsson MD3, Hedvig E. Jakobsson PhD2, Martina Sansone PhD1,4 and Johan Westin PhD2,4

1Department of Clinical Microbiology, Infection Control Unit, Region Vastra Gotaland, Sahlgrenska University Hospital, Gothenburg, Sweden, 2Department of Infectious Diseases, Institute of Biomedicine, Sahlgrenska Academy, University of Gothenburg, Sweden, 3Department of Geriatric Medicine, Region Vastra Gotaland, Sahlgrenska University Hospital, Gothenburg, Sweden and 4Department of Infectious Diseases, Region Vastra Gotaland, Sahlgrenska University Hospital, Gothenburg, Sweden

Abstract

Objective: Effective infection prevention and control (IPC) measures are key for protecting patients from nosocomial infections and require knowledge of transmission mechanisms in different settings. We performed a detailed outbreak analysis of the transmission and outcome of coronavirus disease 2019 (COVID-19) in a geriatric ward by combining whole-genome sequencing (WGS) with epidemiological data.

Design: Retrospective cohort study.

Setting: Tertiary-care hospital.

Participants: Patients and healthcare workers (HCWs) from the ward with a nasopharyngeal sample (NPS) positive for severe acute respiratory coronavirus virus 2 (SARS-CoV-2) RNA during the outbreak period.

Methods: Patient data regarding clinical characteristics, exposure and outcome were collected retrospectively from medical records. Stored NPSs from 32 patients and 15 HCWs were selected for WGS and phylogenetic analysis.

Results: The median patient age was 84 years and 17 (53%) of 32 were male. Also, 14 patients (44%) died within 30 days of sampling. Viral loads were significantly higher among the deceased. WGS was successful in 28 (88%) of 32 patient samples and 14 (93%) of 15 HCW samples. Moreover, 3 separate viral clades were identified: 1 clade and 2 subclades among both patient and HCW samples. Integrated epidemiological and genetic analyses revealed 6 probable transmission events between patients and supported hospital-acquired COVID-19 among 25 of 32 patients.

Conclusions: WGS provided an insight into the outbreak dynamics and true extent of nosocomial COVID-19. The extensive transmission between patients and HCWs indicated that current IPC measures were insufficient. We recommend increased use of WGS in outbreak investigations to identify otherwise unknown transmission links and to evaluate IPC measures.

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WGS and phylogeny for a detailed outbreak analysis of COVID-19 in a geriatric hospital setting.

**Methods**

**Setting**

The outbreak occurred during the first wave of the pandemic in a 2,000-bed tertiary-care hospital in western Sweden that serve a population of ~700,000. The affected 30-bed unit was assigned for orthogeriatric patients without COVID-19 infection and comprised 6 single-bed rooms, 6 two-bed rooms, and 3 four-bed rooms. During the outbreak period, 166 patients received inpatient care at the ward.

**Definitions and data collection**

The outbreak period was set from the sampling day of the index case until 14 days had passed without any newly discovered cases, considering an incubation period of 2–14 days. A patient or HCW from the ward with a nasopharyngeal sample (NPS) positive for SARS-CoV-2 RNA was considered an outbreak case. Patient data were retrospectively collected from medical records, and information regarding contact tracing and ward occupancy from the IPC team and the hospital administrative unit. No individual data were available for HCWs.

HCAIs were classified according to Meredith et al. as true when confirmed >14 days after admission, days 7–14 after admission (suspected), days 5–6 after admission (indeterminate), and ≤2 days after admission (community associated).

**Infection prevention and control measures**

In accordance with the Swedish National Health Authority, personal protective equipment (PPE) was recommended only when within 1 m from a suspected or confirmed case of COVID-19 and included a plastic apron and a full-face visor (stretching below the chin) or a surgical mask (IIR) and face shield or goggles. A respirator (FFP2-3) was added if aerosol-generating procedures were performed in the room. Gloves and a long-sleeve apron were recommended for those at risk of contact with bodily fluids.

Patients were triaged for COVID-19–associated symptoms upon arrival at the emergency department and were considered suspected cases if they presented at least 2 of the following symptoms: cough, sore throat, fever, and shortness of breath (or upon judgment by the treating physician). Suspected cases were isolated at a quarantine ward until the diagnosis was confirmed or averted. Confirmed cases were transferred to assigned COVID-19 wards. Visitor restrictions were enforced throughout the hospital and admitted patients were restricted to their rooms.

Contact tracing was performed around all confirmed cases. Patients sharing a room with a case during their infectious phase were considered close contacts and were isolated and monitored for symptoms for 14 days. HCWs were considered close contacts when exposed to an infectious case without PPE and continued to work if asymptomatic during the incubation period. HCWs with symptoms of COVID-19 self-quarantined at home for at least 7 days unless they tested negative. Testing resources were limited and prioritized for suspected patient cases requiring in-hospital care.

Laboratory methods and details on bioinformatics and phylogenetic analysis are provided in the Supplementary Material (online).

**Ethical statement**

Approval for this study was granted by the Swedish Ethical Review Authority (protocol no. 2020-03276).

**Results**

**Outbreak description**

In total, 32 patients and 15 HCWs were included in this study (Fig. 1). The index case (patient 1) developed COVID-19 symptoms and was sampled 8 days after admission (outbreak day 0). The first secondary case (patient 2) tested positive on outbreak day 5, 11 days after admission. Close contact between them was excluded. Several staff members reported illness during this period, and HCW–patient transmission was suspected. An outbreak investigation initiated by staff management and the IPC team identified several possible factors contributing to transmission: difficulties in symptom interpretation, crowding in workspaces, PPE shortage and insufficient IPC training. Testing of HCWs was available from outbreak day 14. IPC training and PPE resource allocation was initiated in week 3. By week 5, HCWs used full-face visors in all patient care activities and social gatherings were limited during breaks. The ward closed for new admissions on day 30, and screening of the remaining patients (n = 17) identified 8 cases, of whom 5 were asymptomatic. Repeated screening (n = 9) on day 32 identified 1 additional case.
Case characteristics

Patient characteristics are shown in Supplementary Table 1 (online). The median age was 84 years and 17 (53%) of 32 patients were male. The overall 30-day mortality was 44% (death occurring in median 8 days after sampling). No additional mortality was observed within 90 days, and no cases were lost to follow-up.

Viral load was significantly higher among the deceased. Also, 5 asymptomatic patients were identified, of whom 4 developed symptoms within the following 4 days (Supplementary Fig. 1 online). One patient remained asymptomatic within 5 days of follow-up and had a possibly false-positive test due to a very low viral load (Ct value, 39).

Outbreak analysis

WGS was successful in 28 (88%) of 32 patient samples and 14 (93%) of 15 HCW samples. Patient 31 was excluded from phylogenetic analysis due to low viral load. Patients 15 and 23 and HCW 7 were excluded due to lack of material and patient 27 was excluded due to low genomic coverage. Viral strains from 3 genetically distinct clades (20A–C) were found among both outbreak and community sequences, although outbreak sequences showed greater internal genetic similarity (Supplementary Fig. 2a online).

Two outliers among patient sequences were found in clade 20B and clade 20C. Based on the phylogenetic clustering, clade 20A was separated into subclades I and II, which also appeared during different phases of the outbreak (Fig. 1). Clade 20A-I and 20B were dispersed among both patient and HCW sequences (Supplemental Fig. 2b online). These phylogenetic analyses suggest 4 separate introductions, of which 3 (20A-I and 20B) resulted in secondary transmission.

Contact tracing revealed an epidemiological link (close contact) between 22 of 32 patient cases (Fig. 2). The phylogenetic analysis did not support transmission in 5 of these cases due to clade differences (patients 4, 18, and 21) or sequence differences (patients 9 and 13) (Supplementary Fig. 2b online and Fig. 2). In contrast, for 6 of the 17 remaining cases (patients 11, 17, 20, 25, 27, and 28), a patient–patient transmission event between close contacts was supported by a positive nasopharyngeal sample (NPS) or symptom onset occurring within 2–14 days of each other (Fig. 2).

Based on the case definitions, 12 of 32 patients had true HCAIs. The phylogenetic analysis revealed a close relationship to other outbreak sequences for 27 of 28 patients (Supplemental Fig. 2a online). The single finding of clade 20C (patient 18, NPS 4 days after admission) supported community transmission. Patient 13

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**Fig. 2.** Patient–patient transmission of SARS-CoV-2 in a hospital ward outbreak. The panel display 22 patient cases defined as close contacts due to sharing a room with another case. Individual case numbers and letters indicating shared room are seen on the y-axis, and timeline of the outbreak period (days) seen on the x-axis. Bars show day of admission until discharge from the affected ward. Colors represent duration of shared room with another case and viral clade. Dots indicate time point for sampling and stars represent symptom onset. A patient–patient transmission event was probable if: sequence differences did not exclude a genetic relationship, and day of sampling or symptom onset for 2 close contacts occurred within 2–14 days of each other.
(NPS 6 days after admission, day 20) and HCW 9 (day 30) were genetically similar although more closely related to community sequences than other outbreak sequences. However, the direction of transmission between them is unclear due to lack of clinical data for HCWs. Subclone 20AI was first identified in HCW 3, and patient 1 may have introduced 20AI (Fig. 1). In contrast, clade 20B was unlikely introduced by either patient 4 or 5 because they had no close contact and were sampled the same day (Supplementary Fig. 1 online). Altogether, strong support of true HCAI was found for 25 (78%) of 32 patients.

Discussion

We present an integrated epidemiological and genetic analysis of a COVID-19 hospital outbreak resulting in the discovery of not 1 but multiple separate viral clusters. We also found that instead of 4 patients, 25 patients likely had true HCAI, highlighting both the uncertainties of nosocomial COVID-19 case definitions and the power of WGS.

The mortality in our study was in line with the national mortality rate for COVID-19 in the group aged 80–89 years during this period,25 although it was higher than previous reports of ~30% among elderly hospitalized COVID-19 patients.3,4,28,29 COVID-19 has been reported as a significant risk factor of death within 30 days for patients with hip fractures,11,28,29 which may have influenced our results. The short median survival time (8 days) corresponds with previous findings,2,27 supporting the finding that death was caused by acute infection. Significantly higher viral load was seen among the deceased patients, previously reported in geriatric patients.30 The severe outcome stresses the importance to protect this patient group from COVID-19.

Asymptomatic transmission has been suggested a key factor in hospital outbreaks.7,8 Interpreting symptoms in elderly patients with COVID-19 may be difficult6,31 and asymptomatic cases are more common among patients aged >80 years.32 The asymptomatic cases identified in our study support the insufficiency of a strictly symptom-based testing strategy. We recommend screening combined with serial testing, especially during significant community transmission or when hospital outbreaks are suspected.9,33

We identified only 6 events of probable patient–patient transmission. This finding suggests that transmission from HCWs might have occurred, which has been reported previously.4,7,9,12,20,33 However, the direction is often unclear, and exposure from HCWs seldom appears to result in infection.34 The close genetic relationship between sequences from HCWs and patients support the hypothesis that transmission occurred between them; hence, breaches in IPC measures were identified. The symptom-based recommendations overlooked silent transmission from pre- or asymptomatic individuals, and PPE recommendations may have been insufficient. Adherence to IPC measures was unknown. Therefore, the cause of infection could have been inadequate IPC or PPE recommendations, insufficient adherence, or all of these.

The main limitation of this study was the restricted testing policy, which resulted in unrecognized cases among patients and HCWs that might have influenced the course of the outbreak. Establishing the direction of transmission was complicated due to lack of clinical information for HCWs. Patients and HCWs had multiple contacts in other units and transmission outside of the ward may have been overlooked.

The details provided by WGS and phylogeny emphasize the limitations of classic outbreak investigations and the potential of molecular characterization. We recommend increasing the use of WGS for outbreak investigations to clarify transmission links, to identify nosocomial infections, and to evaluate IPC measures.

Supplementary material. To view supplementary material for this article, please visit https://doi.org/10.1017/ice.2021.374

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Conflicts of interest. All authors report no conflicts of interest relevant to this article.

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