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Research note

SARS Coronavirus-2 variant tracing within the first Coronavirus Disease 19 clusters in northern Germany

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Objectives: Investigation whether in depth characterization of virus variant patterns can be used for epidemiological analysis of the first severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection clusters in Hamburg, Germany.

Methods: Metagenomic RNA-sequencing and amplicon-sequencing and subsequent variant calling in 25 respiratory samples from SARS-CoV-2 infected patients involved in the earliest infection clusters in Hamburg.

Results: Amplikon sequencing and cluster analyses of these SARS-CoV-2 sequences allowed the identification of the first infection cluster and five non-related infection clusters occurring at the beginning of the viral entry of SARS-CoV-2 in the Hamburg metropolitan region. Viral genomics together with epidemiological analyses revealed that the index patient acquired the infection in northern Italy and transmitted it to two out of 134 contacts. Single nucleotide polymorphisms clearly distinguished the virus variants of the index and other clusters and allowed us to track in which sequences worldwide these mutations were first described. Minor variant analyses identified the transmission of intra-host variants in the index cluster and household clusters.

Conclusions: SARS-CoV-2 variant tracing allows the identification of infection clusters and the follow up of infection chains occurring in the population. Furthermore, the follow up of minor viral variants in infection clusters can provide further resolution on transmission events indistinguishable at a consensus sequence level.

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Introduction

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) first emerged in late 2019 in Wuhan, China [1]. Sequences of SARS-CoV-2 were immediately publicly available and online tools, such as GISAID (Global initiative on sharing all influenza data), allowed phylogenetic network analyses [2–4]. Genomic epidemiology of SARS-CoV-2 in different countries allowed conclusions to be drawn on major viral lineages, transmission chains and routes of infection [5–7]. Genome surveillance has been highly useful to monitor distinct geographical lineages. However, no notable association of genome sequence variation with transmissibility and pathogenicity has been reported with the exception of one mutation in the spike protein, D614G, contributing to higher transmissibility [8] and becoming a major global variant.

We used viral genomics and variant calling to describe the initial entry of SARS-CoV-2 into the metropolitan region of Hamburg by proven and suspected travel returnees in February and March 2020. We were able to trace back the putative geographical infection sites and characterized viral transmission chains in the earliest documented infection clusters in Hamburg.

Methods

Patients and SARS-CoV-2 diagnostics

The index patient (P0), the first case of coronavirus disease 2019 (COVID-19) in northern Germany, tested SARS-CoV-2-positive on 27 February 2020 after returning from a vacation in northern Italy. His spouse (P02) and a co-worker (P01) tested positive on 4 March 2020 after returning from a vacation in northern Italy. Sequences from P01 and P02 acquired SARS-CoV-2 from P0, who was initially infected in northern Italy (see Supplementary material, Fig. S1). Sequences from P39 and P40, a couple returning from Italy, which cluster in II.2, do not show sequence variation at position 160.

Pattern II.3 and II.4, each separate from II.2 by one additional SNP. Pattern II.3 represents sequences from two families returning from a joint vacation in Italy. SNPs in II.3 were first described in sequences simultaneously sampled in the Netherlands, Italy and Austria. Sequences in pattern II.4 are derived from related travel returnees; characteristic SNPs in II.4 have been first reported in a sequence from France.

Pattern III, representing a family cluster, is defined by SNPs first reported in a sequence from North-Rhine Westphalia, Germany. Finally, pattern I, a sequence from a travel returnee from the Middle East shows 12 SNPs with six of them being previously reported in a sequence sampled in Oman.

Based on amplicon-seqencing, we defined minority variants, sub-consensus viral populations, at 14 positions with ten resulting in synonymous mutations.

Interestingly, in two family clusters, we identified minority variants allowing us to follow transmission events. In cluster II.4, sequences of P22 and P48 display multiple variants at nt 199 while P21 showed only one variant. The observed variant fractions are suggestive of transmission from P22 or P48 to P21 or alternatively infection from a common source and P21 has lost one variant. The transmission of intra-host variants with variant frequencies of the founding population in the recipient closely matching the donor suggests a high viral load exposure or a loose bottleneck in transmission events.

Similarly, in cluster II.3 with two families returning from a joint vacation, sequences from P06, P09 and P10 from the same household show intra-host variants at position 11,438 indicative of high infection dose exposure during viral transmission from a common source to all three patients or between these individuals. With sampling 4–5 days apart (Figs. 1a), P06 and P10 might have had the same infection source, and one of them transmitted the virus to P09.

Discussion

We here report the genomic analysis of the index cluster and five unrelated family/household clusters occurring at the beginning of the pandemic in the metropolitan region of Hamburg, Germany. Our analysis created important insights into transmission events and molecular epidemiology of SARS-CoV-2. Based on comparative
Fig. 1. (a) Clustering of viral variants of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) sequences recovered from the index patient, patient 1, patient 2 and 19 SARS-CoV-2 sequences from respiratory swabs collected in the same time period in comparison to the reference sequence, NC_045,521. Nucleotide positions are indicated at the bottom. Only variants with sufficient coverage (>10) and single nucleotide polymorphism (SNP) present in more than 33% of all reads in at least one sample are included. I–III summarizes sequence patterns as defined by SNPs. The frequency of variants is indicated by the heat map ranging from grey (reference), yellow to dark blue (variant). The quality score per individual site is indicated at the top. * indicates members within one family. Sampling dates are indicated on the right with the sampling date of the cases in the index cluster labelled in red. (b) Overview of individual SNPs defining pattern I, II (II.1–II.4) and III with sequence variation in comparison to the reference sequence.

Listed are nucleotides identified in all sequences of one cluster; nucleotide positions in bold indicate mutations identified in GISAID. REF: NC_045512.2. Rare: <10%; low 10-25%; medium 25-49%; high ≥50%; n.a.: not assigned; * Yin; C. Genomics 2020.
SNP analysis from our study and additional SARS-CoV-2 sequences from GISAID, we identified different virus clusters and variant patterns, which allowed us to discern the first infection clusters in Hamburg and to propose intra-cluster transmission routes. Transmission of SARS-CoV-2 from the index patient occurred to only two out of 132 contacts with whom prolonged and unprotected interaction took place.

The overall mutation frequency was relatively low in our analysis when compared with a previous report [12]. These differences may be explained by our variant calling approach, employing >33% variant frequencies as cut-off compared with 5% [12].

By following the transmission of intra-host variants, we propose high-dose transmission of SARS-CoV-2 occurring in these clusters. In contrast to our study, no evidence of transmission of intra-host variants has been described in a previous study [12], whereas in another study identical minor variants were identified in household and workplace clusters [13].

Overall, our study describes the initial SARS-CoV-2 clusters occurring in Hamburg and emphasizes that virus variant tracing can provide further resolution for the identification of transmission events, for example in clusters that are indistinguishable at the consensus sequence level.

Author information

SP, RK, TG, AG, MA, JK, ML and NF designed the study. SP, TG, RS, MA, JK, ML and NF performed the literature search. SP, RK, TG, RS, MA, JK, ML and NF wrote the manuscript. SP, RK, MC, DI, ML and NF performed the bioinformatics analysis. SP, TG, MC, DI, ML and NF generated the figures and tables.

Transparency declaration

We declare that there are no competing interests.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.cmi.2020.09.034.

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