Rapid Genomic Characterization of SARS-CoV-2 Viruses From Clinical Specimens Using Nanopore Sequencing

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Abstract

The outbreak of the novel SARS-CoV-2 has swiftly spread worldwide. Rapid genome sequencing of the SARS-CoV-2 strains has become a helpful tool for better understanding of virus genomic characteristics and the origin. To obtain the virus whole-genome sequence directly from the clinical specimens, we performed the nanopore sequencing using a modified ARTIC protocol on portable nanopore sequencer, and validated the routine 8 hours workflow and 5 hours rapid pipeline. We had made some optimizations to improve the genome sequencing workflow. The sensitivity of the workflow was also tested by serially diluting RNA from clinical samples. The optimized pipeline was finally applied to obtain the whole genomes from 17 clinical specimens in Hangzhou from January 2020 to March 2020. In the obtained 17 complete genomes of SARS-CoV-2, 12 variations were found and analyzed. The genomic variations and phylogenetic analysis hinted the multiple sources and different transmission pattern during the COVID-19 epidemic in Hangzhou, China. In conclusion, the genomic characteristics and the origin of the virus could be quickly determined by nanopore sequencing following our workflows.

Introduction

In December 2019, an outbreak of atypical pneumonia with unclear etiology began in Wuhan, a major transportation hub in the center of China. A novel coronavirus similar to severe acute respiratory syndrome coronavirus (SARS-CoV) was then identified as the causative pathogen, which was officially named as SARS-CoV-2 by the International Committee on Taxonomy of Viruses (ICTV). Before that, most of human infected coronaviruses (HCoV) can only cause mild upper respiratory infections (HCoV-229E, HCoV-NL63, HCoV-OC43 and HCoV-HKU1), but sometimes they cross species and cause fatal respiratory disease and outbreaks, as experienced in cases of SARS-CoV or Middle East respiratory syndrome coronavirus (MERS-CoV). The seventh HCoV SARS-CoV-2 made a spillover event in late December 2019 and is swiftly spreading crossed the borders of city and province in the mainland of China, and soon became an emergency of major international concern. As of 24 April 2020, the cumulative number of confirmed human infections rapidly increased to 2,549,632 (175,825 deaths) reported by WHO (https://who.sprinklr.com/), almost 300 times more than the total number of SARS-CoV infected cases.

Hangzhou is a national tourism city with a registered population of 10.36 million, and located at the south wing of the Yangtze River Delta with a humid, subtropical climate facilitating the airborne survival and transmission of viruses associated with respiratory infections. The first case was found from a Wuhan returnee in January 19, 2020. As of March 2020, 186 infections were confirmed by viral RNA detection. As the virus genome could be sequenced rapidly on the portable MinION sequencer, the accurate genomic sequencing data generated could be used to trace back to the origin during the viruses spreading, which could bring the molecular epidemiology analysis close to the aim of front-line application.

Therefore, we conduct a modified ARTIC protocol for SARS-CoV-2 genome sequencing on the MinION platform. Two workflows were applied and validated by amplifying and sequencing the genome from SARS-CoV-2 infected clinical samples, and the 17 genomes characteristics of SARS-CoV-2 viruses in Hangzhou were analyzed to study the origin and transmission history of these viruses.

Results

8h and 5h Workflow for SARS-CoV-2 Nanopore Sequencing

In order to acquire the whole genome sequence of SARS-CoV-2 more efficiently, 8 hours workflow was designed for its sequencing throughput and speed after loading library into the flow cell, and 5 hours workflow was designed for rapid library building need (Figure 1). These two work flows were tested on the sample of HZCDC0001 with Ct value 26.51/27.03 (Orf1ab/N), the first sample appeared in Hangzhou, Zhejiang province. In the 8h workflow, nanopore ligation sequencing kit was applied because this protocol can maximize the sequencing throughput and length of reads, in figure 2c, the total bases and coverage of SARS-CoV-2 increased much faster than the 5h workflow, in only 10 minutes after loading library to the flow cell, the coverage reached to almost 100%, here we defined a base had a coverage with at least 15x depth. Meanwhile, 5h workflow took more than 1.5 hours to approach the 100% coverage. The 5h workflow advantaged at the rapid 15 minutes library preparation time, especially for the extreme condition, this work ow can hugely shorten the library preparation time compared to the 2 hours of ligation protocol (figure 1). However, as the nanopore rapid protocol would cleavage DNA to quickly add transposase adapters, sequencing throughput and speed behaved much poorly comparing with the 8h workflow (figure 2a-c). In both workflows, two regions (5231 - 5644 bp and 22798 – 23214 bp, primer pairs #18 and #76) appeared to be the short boards in genome mapping, which needs the further optimization.

Meanwhile, we had made some optimizations to improve the workflow. Since the step of RNA extraction is vital for the follow-up sequencing, we compared the magnetic beads extraction on NP968 (Tianlong, China) with column RNA extraction using RNeasy Mini Kit (QIAGEN, Germany) (Figure 2d). The latter approach seems to yield higher quality RNA, making the depth of coverage nearly doubled (Figure 2f). What's more, PCR procedure took more than 40% time of total workflow, so we tried to decrease the annealing and extension time from 5 minutes to 3 minutes and 1 minute, corresponding to the total time of PCR from about 3 hours to 1 hour. Even if the annealing time reduced to 1 minute, the whole genome sequence still could be obtained from the products of 1h PCR procedure (Figure 2e, g). On account of the different viral titers of SARS-CoV-2 in clinical samples, 3-min annealing time could be considered as the equilibrium point.
asymptomatic or oligo-symptomatic transmission during the first week of symptoms and their role in the global spread of the COVID-19 pandemic is still unclear.

The outbreak of COVID-19 caused by SARS-CoV-2 has swiftly spread worldwide. However, probable origin of SARS-CoV-2 associated with the COVID-19 pandemic is still unclear. Recently reports about COVID-19 cases with none or mild upper respiratory tract symptoms suggest the potential for asymptomatic or oligo-symptomatic transmission during the first week of symptoms. Hence, there is an urgent need for rapid identification and traceability of pathogens for the diseases control and prevention. Deep understanding of the novel virus first comes from the analysis of the genome sequence. In this study, we demonstrate the utility of nanopore sequencing for SARS-CoV-2 genomes from clinical specimens based on modified ARTIC protocol. It allowed confirmation of SARS-CoV-2 infection at genomic-level within a few minutes by sequencing and simultaneously introducing the errors of nanopore sequencing, we first filtered the low quality reads and only homozygous SNP with high quality (phred value > 20) and high site depth (> 50) were considered for downstream analysis. Besides, we also perform illumina sequencing in all samples including HZCDC6706 and HZCDC6789 imported from abroad, which provided the evidence that SNPs from our standard was 100% consistent to illumina data and SNPs can be called on only nanopore sequencing in SARS-CoV-2 genome.

The length of reference SARS-CoV-2 genome (MN908947.3) is 29,903 bp. However, a considerable part of the submitted SARS-CoV-2 genomes are uncompleted. Despite the strategy of building phylogenetic tree based on SNPs was applied to investigate the traceability of interested samples. To avoid introducing the errors of nanopore sequencing, we first filtered the low quality reads and only homozygous SNP with high quality (phred value > 20) and high site depth (> 50) were considered for downstream analysis. Besides, we also perform illumina sequencing in all samples including HZCDC6706 and HZCDC6789 imported from abroad, which provided the evidence that SNPs from our standard was 100% consistent to illumina data and SNPs can be called on only nanopore sequencing in SARS-CoV-2 genome.

In all obtained 17 complete genomes of SARS-CoV-2, 12 substitutions distributed in three coding sequences (CDS) and 5'UTR were identified based on sequences alignment (Figure 3), including C241T in 5'UTR, 5 synonymous variations and 6 missense variations (Table 2). Nine variations were found in orf1ab gene, the longest gene in SARS-CoV-2 genome, and only one variation was detected in S and ORF3a gene, respectively. The couple infected in Wuhan, HZCDC0012 and HZCDC0013 owned the same two variations in 11083 and 21282 site. Nine variations were detected in the samples HZCDC6789, which was nearest to now in time and imported from abroad, and few variations were detected in the early samples.

Based on marker variants from 507 full genomes shared (GISAID), three major subclades were named as S (Orf8: L84S), G (S: D614G) and V (Orf3: G251V). The phylogenetic relationship reveals that most of our genomes from early COVID-19 patients linked to Wuhan are in the main clade, except HZCDC0025 and HZCDC0167 belonging to subclade V (Figure 3). Two genomes from the international imported cases are in subclade G and V respectively (Figure 3), which hinted multiple sources of transmission from oversea. However, the clade classification will need to be adjusted when the marker is not stable occurring in the other newly sequenced genomes.

Discussion

The outbreak of COVID-19 caused by SARS-CoV-2 has swiftly spread worldwide. However, probable origin of SARS-CoV-2 associated with the COVID-19 pandemic is still unclear. Recently reports about COVID-19 cases with none or mild upper respiratory tract symptoms suggest the potential for asymptomatic or oligo-symptomatic transmission during the first week of symptoms. Hence, there is an urgent need for rapid identification and traceability of pathogens for the diseases control and prevention. Deep understanding of the novel virus first comes from the analysis of the genome sequence. In this study, we demonstrate the utility of nanopore sequencing for SARS-CoV-2 genomes from clinical specimens based on modified ARTIC protocol. It allowed confirmation of SARS-CoV-2 infection at genomic-level within a few minutes by sequencing and simultaneously mapping the reads to reference genome and analyzing the output data in real-time.

Comparing with nasal/oropharyngeal swabs, the lower respiratory tract specimens from COVID-19 could be detected more readily. Our data shows that the virus genomes from different parts of respiratory tract are consistent. However, the difference of viral loads in samples will affect the stability of the average depth and genome-wide coverage, and increase the difficulty of whole genome mapping, suggesting the importance of sample collection for the later genome sequencing.

To characterize the genomic variations, we found 12 different substitution sites distributed in four coding regions among 17 genomes of SARS-CoV-2, without any recombination event. Genomic evidence supported that most of the first early 15 infections were directly or indirectly linked to Wuhan, and
the genomes from two imported infections occur in March 2020 have specific variation comparing with domestic strains. The phylogenetic analysis indicated the potential for human to human transmissions as previously reported elsewhere. For example, the identical substitutions (G11083T: Orf1ab Leu3606Phe and A21282G: Orf1ab Asp7006Gly) in the virus genomes from a couple hinted the family-cluster transmission history of SARS-CoV-2 from a Wuhan traveler to her husband.

In summary, we performed the SARS-CoV-2 genome sequencing on portable nanopore sequencer. Combined with 8 hours workflow, the genomic characteristics and the origin of the virus could be quickly determined. The rapid 5 hours workflow with 15 minutes fast library preparation could be applied on trace-back task out of lab, bringing the genome-level molecular epidemiology analysis to the front-line of the outbreaks. Therefore, based on prompt diagnosis and rapid whole-genome analysis, the swift and decisive response to the SARS-CoV-2 outbreak will be benefit to the diseases control and prevention.

**Methods**

**Ethics statement**

This study and all experimental protocols were approved by the Institutional Review Board of Hangzhou Center for Diseases Control and Prevention. We confirm that all methods were carried out in accordance with relevant guidelines and regulations. Signed informed consents were obtained from the patients or their spouses and personally identifiable information was anonymized.

**Viral Infections Diagnose**

Upper and/or lower respiratory tract samples, including nasal, oropharyngeal swabs, sputum, tracheal aspirate samples and bronchoalveolar-lavage fluid, were collected from suspected cases with informed consent from patients or their spouses, and were sent to Hangzhou Center for Diseases control and prevention on ice for diagnosis within six hours. The viral RNA was extracted directly from 200 μL supernatant of clinical sample using the RNeasy Mini Kit (QIAGEN, Germany) according to the manufacturer's instruction in a biosafety cabinet in Biosafety Level 2 Laboratory, and tested for the presence of SARS-CoV-2 using the diagnostic real-time reverse transcription polymerase chain reaction (qRT-PCR) on ABI7500 (ABI, USA) following the diagnostic kit manual.

**Workflows of Virus Genome Sequencing**

Viral RNA extracted from clinical samples was used as a template to amplify and sequence the SARS-CoV-2 genome. Briefly, cDNA was synthesized from 11 μL viral RNA using SuperScript™ IV First-Strand Synthesis System (Invitrogen) with random hexamers. PCR was performed using Q5 Hot Start High-Fidelity DNA Polymerase (NEB) and a set of primers targeting regions of the SARS-CoV-2 genome designed by ARTIC network (https://artic.network/ncov-2019). The PCR mixture was initially incubated 2 minutes at 98 °C for the denaturation, followed by 35 cycles at 98 °C for 15 seconds, 65 °C for 1, 3 or 5 minutes (depending on the workflows). The amplified products were purified by equal volume of AMPure XP beads (Beckman Coulter) to exclude small nonspecific fragments.

According to eight-hour routine workflow (Figure 1), the purified DNA was repaired with NEBNext FFPE Repair Mix (NEB), followed by the DNA ends preparation using NEBNext End repair / dA-tailing Module (NEB) and the successively attachment of native barcodes and sequencing adapters supplied in the EXP-NBD104/114 kit (Nanopore) to the DNA ends. The DNA concentration was determined by a Qubit 3.0 using dsDNA HS Assay Kit (Thermo Fisher). After priming the flow cell, 60 ng DNA per sample of products were pooled in the DNA library with final volume of 65 μL. Following the ligation sequencing kit (SQK-LSK109) protocol, MinION Mk1B was used to perform the genome sequencing on an R9.4.1 Flow Cell for 1 hour per sample. For rapid barcoding workflow, a fragmentation mix in the SQK-RBK004 kit (Nanopore) was used to attach the barcodes to the DNA ends, followed by the attachment of sequencing adapters.

**Reads Preprocessing and Consensus Building for Nanopore Sequencing**

Base calling was performed by guppy (https://community.nanoporetech.com) with the parameter “-c dna_r9.4.1_450bps_hac.cfg -x auto” and different samples were separated and adapters were trimmed with the additional parameter “-trim_barcodes -barcodes EXP-NBD104/EXP-NBD114/SQK-RBK004”. FASTQ reads were filtered for quality control with the cutoff “length >= 200 and Phred value >= 7” using program “fitlong v0.2.0” (https://github.com/rrwick/Filtlong).

After quality control, artic-ncov2019 pipeline (https://artic.network/ncov-2019) was applied to perform the sequences mapping, primers trimming, variations calling and consensus assembly building. Variations were called using Medaka 0.11.1 (https://github.com/nanoporetech/medaka). In the stage of consensus assembly building, the site with depth lower than 50 x will be masked by N, and reference will be substituted by the homozygous variations with phred quality >= 20. "Samtools depth" was used to calculate the depth of each site and "Samtools bedcov" was used to calculate the window depth scanning in the genome.
Reads Preprocessing and Variations Calling for Illumina Sequencing

Raw illumina PE reads were trimmed and quality controlled by the software fastap 0.20.0 with default parameters. Bwa 0.7.17-r1188 was used to mapping the clean reads to SARS-CoV-2 reference genome and sam/bam files were manipulated by samtools 1.9. Variations were detected with the program "mpileup and calling" from bcftools 1.9, variations will be considered as positive when the homozygous variations has a phred quality value >= 20 and a depth >=50.

Phylogeny and Variations Analysis

In order to remove the bias from the gaps in uncompleted genome, sequence alignment from all SNPs site were chosen to build the phylogenetic tree. First, all SNPs were called aligned to SARS-CoV-2 reference sequences using “nucmer” and “dnadiff” programs from MUMmer 3.23, the effect of SNPs were estimated using SnpEff 4.3t. Second, all SNPs site were connected to a single sequence for every sample based on the variations calling results from the last step. Then, these sequences were combined to perform phylogeny analysis, maximum likelihood phylogenies were estimated by FastTree 2.1.10 with default parameters. The phylogenetic tree with variations heatmap matrix was drew by phyD3. The group and clade numbers were assigned to achieve consistency with the earlier studies.

Declarations

Data Availability

All genome sequences in this study are available in GISAID (http://gisaid.org) (the full list of the accession numbers are available from table 1).

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Competing Interests:

The authors declare no competing interests.

Authors’ Contributions:

W.X.C., L.J., W.H.Q., M.L.F., P.J.C. and S.J.L. designed the study and wrote the manuscript. C.J.F., S.Z. involved in the sample collection. C.S.C., L.J., Y.X.F. and Q.X. did the viral detection. L.J., Y.H. and C.S. participated in the genomes sequencing on nanopore MinION sequencer and illumina Miseq platform. M.L.F., L.J. performed the data analysis. All authors reviewed the manuscript.

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**Tables**

**Table 1** The information of seventeen Hangzhou SARS-CoV-2 viruses from COVID-19 patients in this study.

| Strain number | Source              | Gender | Age (yr) | Date of onset | Collection date | History                          | GISAID accession number | Average depth | Cover-age(%) |
|---------------|---------------------|--------|----------|---------------|-----------------|----------------------------------|-------------------------|---------------|--------------|
| HZCDC0001     | Tracheal aspirate   | Male   | 31       | 2020-01-18    | 2020-01-19      | Infected in Wuhan Spouse of HZCDC0013 | EPI_ISL_407313          | 754.88        | 100.00       |
| HZCDC0012     | Nasal, oropharyngeal swab | Male   | 47       | 2020-01-18    | 2020-01-20      | Infected in Wuhan                | EPI_ISL_421236          | 326.64        | 100.00       |
| HZCDC0013     | Nasal, oropharyngeal swab | Female | 45       | 2020-01-15    | 2020-01-20      | Infected in Wuhan                | EPI_ISL_421235          | 418.66        | 100.00       |
| HZCDC0025     | Nasal, oropharyngeal swab | Male   | 51       | 2020-01-21    | 2020-01-21      | Infected in Wuhan                | EPI_ISL_421234          | 509.46        | 99.99        |
| HZCDC0048     | Nasal, oropharyngeal swab | Male   | 35       | 2020-01-16    | 2020-01-21      | Contact to patient from Wuhan     | EPI_ISL_421233          | 506.14        | 100.00       |
| HZCDC0048L    | Tracheal aspirate   | Female | 40       | 2020-01-17    | 2020-01-21      | Contact to patient from Wuhan     | EPI_ISL_421232          | 385.17        | 100.00       |
| HZCDC0090     | Nasal, oropharyngeal swab | Female | 34       | 2020-01-17    | 2020-01-21      | Contact to patient from Wuhan     | EPI_ISL_421230          | 233.75        | 99.15        |
| HZCDC0090L    | Bronchoalveolar-lavage fluid | Female | 30       | 2020-01-17    | 2020-01-21      | Contact to patient from Wuhan     | EPI_ISL_421229          | 555.60        | 100.00       |
| HZCDC0091     | Nasal, oropharyngeal swab | Male   | 31       | 2020-01-17    | 2020-01-21      | Contact to patient from Wuhan     | EPI_ISL_421228          | 264.45        | 99.16        |
| HZCDC0091L    | Bronchoalveolar-lavage fluid | Male   | 33       | 2020-01-17    | 2020-01-21      | Contact to patient from Wuhan     | EPI_ISL_421227          | 535.46        | 99.97        |
| HZCDC0119     | Nasal, oropharyngeal swab | Female | 41       | 2020-01-21    | 2020-01-22      | Contact to patient from Wuhan     | EPI_ISL_421226          | 464.11        | 100.00       |
| HZCDC0135     | Nasal, oropharyngeal swab | Male   | 62       | 2020-01-15    | 2020-01-22      | Infected in Wuhan                | EPI_ISL_421225          | 509.26        | 99.95        |
| HZCDC0162     | Nasal, oropharyngeal swab | Male   | 46       | 2020-01-21    | 2020-01-23      | /                                 | EPI_ISL_421224          | 488.18        | 99.61        |
| HZCDC0167     | Nasal, oropharyngeal swab | Female | 53       | 2020-01-21    | 2020-01-23      | /                                 | EPI_ISL_421223          | 464.55        | 98.08        |
| HZCDC6706     | Nasal, oropharyngeal swab | Female | 37       | 2020-03-12    | 2020-03-14      | Infected in U.S.A.                | EPI_ISL_421222          | 507.37        | 100.00       |
| HZCDC6789     | Nasal, oropharyngeal swab | Male   | 21       | 2020-03-13    | 2020-03-15      | Infected in U.K.                  | EPI_ISL_421221          | 365.88        | 100.00       |

**Table 2** The list of substitutions in all obtained 17 complete genomes of SARS-CoV-2.
## Variations

| Site | Variations | Nanopore SNP quality value | Nanopore depth | Illumina SNP quality value | Illumina depth | Variation type | Gene | Amino acid change | Samples |
|------|------------|---------------------------|----------------|---------------------------|----------------|----------------|------|------------------|---------|
| 241  | CàT        | 30.9                      | 447            | 225.0                     | 7421           | Intergenic region SNP | SUTR | None             | HZCDC6706 |
| 6187 | CàT        | 58.8                      | 412            | 228.0                     | 2645           | Synonymous variant    | orf1ab | His1974His       | HZCDC0025 |
| 9190 | GàT        | 56.0                      | 299            | 225.0                     | 7145           | Synonymous variant    | orf1ab | Val2975Val       | HZCDC6789 |
| 11083| GàT        | 31.4/22.8/33.5/32.2       | 183/178/329/162| 228.0/225.0/225.0/225.0 | 2909/4821/6846/6139 | Missense variant | orf1ab | Leu3606Phe | HZCDC0012, HZCDC0013, HZCDC00167, HZCDC6789 |
| 14034| TàC        | 59.2                      | 345            | 225.0                     | 7981           | Missense variant    | orf1ab | Met4590Thr       | HZCDC6789 |
| 14805| CàT        | 63.2                      | 368            | 228.0                     | 7689           | Missense variant    | orf1ab | Thr4847Ile       | HZCDC6789 |
| 16468| CàT        | 54.2                      | 647            | 228.0                     | 1577           | Synonymous variant    | orf1ab | His5401His       | HZCDC6789 |
| 17247| TàC        | 46.1                      | 319            | 228.0                     | 7958           | Missense variant    | orf1ab | Val5661Ala       | HZCDC6789 |
| 19646| TàC        | 58.1                      | 265            | 228.0                     | 3637           | Synonymous variant    | orf1ab | Leu6461Leu       | HZCDC6789 |
| 21282| AàG        | 56.2/44.0                 | 221/274        | 225.0/225.0               | 1391/2248      | Missense variant    | orf1ab | Asp7006Gly       | HZCDC0012, HZCDC0013 |
| 24133| CàT        | 61.5                      | 691            | 228.0                     | 6456           | Missense variant    | orf1ab | Gly657Val        | HZCDC6789 |
| 26144| GàT        | 35.4/24.9/58.8            | 412/407/324    | 228.0/228.0/228.0         | 7846/7748/7739 | Missense variant    | ORF3a | Gly251Val        | HZCDC0025, HZCDC0167, HZCDC6789 |

**Figures**

**Figure 1**

Overview of two nanopore sequencing workflows. The white boxes represent a series of tasks as the components of 8 hours routine workflow and 5 hours rapid workflow respectively. The numbers in the colorful boxes indicate the time required to complete the tasks.
Figure 2

Analysis of genome sequencing data of SARS-CoV-2 viruses by Oxford Nanopore Technologies using MinION sequencer. (a) Trends of the data of depth appearing among 16 hours sequencing using Ligation Sequencing Kit 109. (b) The genome-wide depth of coverage while using Rapid Barcoding Kit 004. (c) Comparative analysis of the average depth and genome-wide coverage between routine workflow and rapid workflow. (d-g) Method optimization of nucleic acid extraction (magnetic beads or spin column method) and PCR amplification procedure. (h-j) Repeated assays of nanopore sequencing of viral RNA with 10 times dilution. (k) The depth of genome-wide coverage appears when applying the optimized methods in clinical specimens.

Figure 3
Phylogenetic analysis of seventeen Hangzhou SARS-CoV-2 strains with reference genome sequences. The histogram in the upper panel refer to the frequency of occurrence of these variations. In the lower panel, the missense and synonymous variants were marked as dark blue and light blue respectively. ORF: open reading frame; S: spike protein gene; E: envelope protein gene; M: membrane protein gene; N: nucleocapsid protein gene. Numbers above ORFs are genome positions according to the reference SARS-CoV-2 genome (Genbank accession number MN908947).