Coding-Complete Genome Sequences of Three SARS-CoV-2 Strains from Bangladesh

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ABSTRACT We report the sequencing of three severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) genomes from Bangladesh. We have identified a unique mutation (NSP2_V480I) in one of the sequenced genomes (isolate hCoV-19/Bangladesh/BCSIR-NILMRC-006/2020) compared to the sequences available in the Global Initiative on Sharing All Influenza Data (GISAID) database. The data from this analysis will contribute to advancing our understanding of the epidemiology of SARS-CoV-2 in Bangladesh as well as worldwide at the molecular level and will identify potential new targets for interventions.

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) belongs to the genus Betacoronavirus, family Coronaviridae, and is the causative agent of the ongoing coronavirus disease 2019 (COVID-19) pandemic (https://coronavirus.jhu.edu/map.html). SARS-CoV-2 was first reported in Bangladesh in March 2020, and since then the country has experienced a steady rise in infections, resulting in ~115,786 cases and 1,502 deaths as of 22 June 2020 (https://corona.gov.bd/?gclid). Here, we report the complete sequences of three SARS-CoV-2 isolates from patients who tested positive using quantitative PCR (qPCR) in the National Institute of Laboratory Medicine and Referral Center (NILMRC). qPCR was performed using a Sansure Biotech novel coronavirus (2019-nCoV) nucleic acid diagnostic kit according to the manufacturer’s instructions. The threshold cycle (Ct) values of the N gene of the three positive samples were 24.3, 22.6, and 23.4, and the Ct values of the open reading frame (ORF) genes were 24.9, 27.5, and 22.6, respectively.

The study was approved by the ethics committee of NILMRC (Bangladesh). The samples were taken with the consent of all patients, and they provided informed written consent consistent with the experiment. To understand the molecular epidemiology of SARS-CoV-2 viruses in Bangladesh, we carried out whole-genome sequencing of three isolates (hCoV-19/Bangladesh/BCSIR_NILMRC_006/2020 [BCSIR_NILMRC_006], hCoV-19/Bangladesh/BCSIR_NILMRC_007/2020 [BCSIR_NILMRC_007], and hCoV-19/Bangladesh/BCSIR_NILMRC_008/2020 [BCSIR_NILMRC_008]) collected directly from nasopharyngeal swabs from three patients in Bangladesh. Viral RNA was extracted using a PureLink viral RNA/DNA minikit (catalog no. 12280050; Thermo Fisher Scientific, USA). The cDNA of all three samples was used to prepare paired-end libraries with the Nextera DNA Flex library preparation kit according to the manufacturer’s instructions (Illumina, Inc., San Diego, CA). The extracted viral RNA was converted into cDNA using the GoScript reverse transcription system protocol according to the manufacturer’s
The library pool of three samples was sequenced using a NextSeq high-output kit with an Illumina NextSeq 550 instrument in paired-end format (read length, 151 bp).

The library generated 73,164,498, 109,775,424, and 70,072,652 reads, of which 72,000,769, 106,842,322, and 68,549,83 unique reads, respectively, were found after excluding duplicate marked reads using BaseSpace DRAGEN RNA Pathogen Detection software version 3.5.1 (1). Sequence trimming and quality control were done using the BaseSpace DRAGEN app. Quality control examination of the sequencing reads revealed that >99% of the sequencing data yielded a Phred score of 30 or above. We did not identify any overrepresented adapter sequences in the sequencing libraries. After generating a FASTA file from the FASTQ files using the DRAGEN software, it was found that the complete genome sequences of the Bangladeshi SARS-CoV-2 strains (BCSIR_NILMRC_006, BCSIR_NILMRC_007, and BCSIR_NILMRC_008) have linear RNAs of 29,892 bp, 29,823 bp, and 29,758 bp, respectively, with an average GC content of 39%. Data for the three samples were uploaded to the Global Initiative on Sharing All Influenza Data (GISAID) database on 30 May 2020 (2). Phylogenetic analysis of these three virus genomes grouped them in SARS-CoV-2 clade 20B (Fig. 1).

This phylogeny shows the evolutionary relationships of human coronavirus 2019 (hCoV-19; or SARS-CoV-2) viruses from the ongoing COVID-19 pandemic. The results show an initial emergence in Bangladesh in May and June 2020, followed by sustained human-to-human transmission, leading to the sampled infections.

Compared with hCoV-19/Wuhan/WIV04/2019, for strain BCSIR-NILMRC-006, we found eight mutations, including NSP2_G339S, N_R203K, N_G204R, NSP3_Q172R, Spike_D614G, NSP2_I120F, NSP12_P323L, and NSP2_V480L. Six mutations were found in BCSIR-NILMRC-007, Spike_D614G, N_R203K, N_G204R, NSP12_K59N, NSP2_I120F, and NSP12_P323L. In BCSIR-NILMRC-008, the genome mutations Spike_D614G, N_R203K, N_G204R, NSP2_I120F, NSP12_P323L, and NSP3_P822S were observed (Table 1). A unique mutation (NSP2_V480I) was observed in the BCSIR-NILMRC-006
Data availability. The sequences of these SARS-CoV-2 genomes from Bangladesh were submitted to the GISAID database (accession no. EPI_ISL_455458, EPI_ISL_455420, and EPI_ISL_455459) and GenBank (accession no. MT539159, MT539158, and MT539160). The raw reads have been deposited in the BaseSpace cloud and were also submitted to the NCBI SRA under accession no. PRJNA658211, PRJNA657985, and PRJNA657938 for strains BCSIR_NILMRC_008, BCSIR_NILMRC_007, and BCSIR_NILMRC_006, respectively.

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