Extensive genetic diversity with novel mutations in spike glycoprotein of severe acute respiratory syndrome coronavirus 2, Bangladesh in late 2020

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Abstract

In Bangladesh, coronavirus disease 2019 (COVID-19) has been highly prevalent during late 2020, with nearly 500,000 confirmed cases. In the present study, the spike (S) protein of severe acute respiratory coronavirus 2 (SARS-CoV-2) circulating in Bangladesh was genetically investigated to elucidate the diversity of mutations and their prevalence. The nucleotide sequence of the S protein gene was determined for 15 SARS-CoV-2 samples collected from eight divisions in Bangladesh, and analysed for mutations compared with the reference strain (hCoV-19/Wuhan/WIV04/2019). All the SARS-CoV-2 S genes were assigned to B.1 lineage in G clade, and individual S proteins had 1–25 mutations causing amino acid substitution/deletion. A total of 133 mutations were detected in 15 samples, with D614G being present in all the samples; 53 were novel mutations as of January 2021. On the receptor-binding domain, 21 substitutions including ten novel mutations were identified. Other novel mutations were located on the N-terminal domain (S1 subunit) and dispersed sites in the S2 subunit, including two substitutions that remove potential N-glycosylation sites. A P681R substitution adjacent to the furin cleavage site was detected in one sample. All the mutations detected were located on positions that are functionally linked to host transition, antigenic drift, host surface receptor binding or antibody recognition sites, and viral oligomerization interfaces, which presumably related to viral transmission and pathogenic capacity.

Keywords: Bangladesh, mutation, severe acute respiratory syndrome coronavirus 2, spike glycoprotein

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Introduction

Coronavirus disease 2019 (COVID-19) pandemic is caused by a novel species of coronavirus, i.e. severe acute respiratory syndrome coronavirus type 2 (SARS-CoV-2) [1]. The virus was first recognized in Wuhan, China, in December 2019 and since then its worldwide spread has had a detrimental effect on global health and economy [2]. In Bangladesh, the first case of COVID-19 was identified on 8 March 2020 (https://corona.gov.bd/). Then, to prevent further human-to-human transmission, measures were scheduled quickly by the Government of Bangladesh including initial lockdown, social distancing, movement restrictions, closure of public and private facilities, and reduction of domestic and international air transport. Nevertheless, despite these strict measures, Bangladesh has been vulnerable to the current COVID-19 pandemic because it is one of the world’s most densely populated countries. As a consequence, until the end of January 2021, a total of 535,230 cases with 8,102 deaths have been recorded (https://corona.gov.bd/) officially in the country.
Given the evolving nature of the SARS-CoV-2 genome, drug and vaccine developers continue to be vigilant for the emergence of new variants or sub-strains of the virus [3]. Knowledge of the molecular background of circulating strains is vital for the appropriate selection of protective vaccine targets and to maintain molecular diagnostic tools for Bangladesh. Although structural proteins of SARS-CoV-2 show extremely high conservation, mutations have been identified, most frequently in the spike (S) protein as well as in the nucleocapsid protein, among global genomic data of this virus [4]. The S protein of coronaviruses is a trimeric glycoprotein that contains two subunits (S1 and S2) that mediate attachment and fusion of viral and cellular membranes, respectively [5,6]. In the process of viral entry into a cell, the receptor-binding domain (RBD) of the S1 subunit attaches to the host cell’s receptor angiotensin-converting enzyme 2, while the S2 subunit mediates fusion of viral and cellular membranes [7]. For the process of virus–cell fusion, the S protein requires priming by host proteases in cleavage sites with a polybasic (furin) cleavage motif (RRAR) at the S1/S2 boundary [7,8]. Mutations of the S protein have been considered important because they are associated with alteration of infectivity and antigenicity of SARS-CoV-2 [9,10].

In Bangladesh, the SARS-CoV-2 genome has been analysed in several reports, revealing variation and frequency of mutations in all the viral proteins [11–17]. However, mutations on the S protein have not yet been well investigated and sufficient information is not available. In the present study, we analysed variation and frequency of missense mutations in the S protein of SARS-CoV-2 from geographically wide areas in Bangladesh. The results revealed extensive diversification of the S protein associated with a wide spectrum of mutations that may affect viral transmission and pathogenicity.

Materials and methods

Viral RNA samples from COVID-19 patients were collected from laboratories in eight divisions of Bangladesh (Barisal, Chittagong, Dhaka, Khulna, Mymensingh, Rajshahi, Rangpur, Sylhet) between July and December 2020. Initial detection of SARS-CoV-2 in nasopharyngeal swab samples was performed by quantitative RT-PCR targeting the ORF1ab and N genes using a commercial kit (Novel Coronavirus Nucleic Acid Diagnostic Kit, Sansure Biotech, Changsha, China). Among 229 samples collected, 15 samples with higher viral load, presumed by the Cq value of the ORF1ab and N genes, were selected for further analysis (see Supplementary material, Table S1).

Full-length of S gene was amplified by conventional RT-PCR with newly designed primer sets (see Supplementary material, Table S2). PCR products were purified and were subjected to Sanger-sequencing. The obtained sequences were assembled, edited and aligned using BioEdit software. Sequence data obtained in this study were deposited in the GISAID database, from which accession numbers were assigned (see Supplementary material, Table S3). A phylogenetic tree was generated using MEGA X software [18] applying the maximum likelihood method and the Tamura–Nei model [19]. Deduced amino acid sequences of S proteins were aligned and compared with the reference strain (hCoV-19/Wuhan/WIV04/2019). The major amino acid changes at the receptor binding site, antigenic site and cleavage site of the S protein were represented in a three-dimensional model using Swiss-Model (https://swissmodel.expasy.org/) software. This study was approved by the institutional review board of Mymensingh Medical College, Bangladesh (MMC/IRB/2020/290).

Results

Nucleotide sequences of 15 SARS-CoV-2 S genes were assigned to clade G (GISAID nomenclature) and further clustered with the B.1 lineage [20] with 98.1%–99.9% identity to the reference strain hCoV-19/Wuhan/WIV04/2019. Phylogenetically, these 15 S protein genes were scattered over several sub-branches together with viruses of this group distributed globally (see Supplementary material, Fig. S1), showing the highest nucleotide sequence identity (99.21%–100%) to strains from the USA, India, Bangladesh, the Netherlands, Colombia, Egypt, Australia, South Africa and Canada (see Supplementary material, Table S4).

A total of 189 single nucleotide polymorphisms by comparison with the S gene sequence of the reference strain were detected in 15 SARS-CoV-2 in Bangladesh (Table 1). Deduced amino acid sequences of 15 samples exhibited a total of 133 replacements (substitution, deletion) with 110 different types of amino acid substitution (see Supplementary material, Table S5). Individual S proteins had 1–25 amino acid replacements (average 8.9 mutations). Among the amino acid replacements identified in the present study, 80 were previously known and 53 were novel mutations as of January 2021.

The S1 subunit contained more amino acid substitutions than S2, almost equally distributed in the N-terminal domain and RBD, and others in the subdomains and furin cleavage site (FCS). Among the 15 Bangladesh SARS-CoV-2 samples, previously reported substitutions in the S1 subunit were located mostly in the N-terminal domain and in the vicinity of the FCS (48 among 58 substitutions). In contrast, novel mutations in the S1 subunit (total 26 substitutions) were identified mainly in the RBD (ten substitutions) and subdomain–FCS (nine substitutions). On the S protein RBD, 21 mutations were...
identified. Among the ten novel substitutions on RBD, three substitutions (P491H, G496W, P499A) were located on the receptor-binding motif along with two known substitutions. Substitution D614G was detected in all samples, among which two samples had mutation Q675H. Substitution P681R, located adjacent to the FCS motif, was detected in one sample. Two novel substitutions, N282D and N1098D, were presumed to remove a potential N-glycosylation site. As depicted in Fig. 1, the positions of representative novel substitutions in RBD are mapped near the outermost edge of S protein, whereas N1098D is located in the root of the S2 subunit close to the viral envelope. The missense mutations identified in S proteins were predicted to affect antigenicity, receptor binding ability, and viral oligomerization depending on the functional domains in S protein (see Supplementary material, Table S6).

### Discussion

Mutation rate of SARS-CoV-2 has been estimated to be $6 \times 10^{-4}$ nucleotide/genome/year, which is lower than other RNA viruses [21,22]. However, minimal mutations in the genome, represented by a single D614G mutation in the S protein, have the capacity to alter the traits of a protein, affecting virus infectivity, clinical outcomes, and also epidemiology [23,24]. Accordingly, it is important to monitor genetic diversity and mutations of SARS-CoV-2 on a global scale during the ongoing pandemic.

In Bangladesh, almost solely the D614G substitution in SARS-CoV-2 S protein was noted and its predominance was revealed by genomic analysis [11,13–15], as observed in the present study. However, other mutations in the S protein have not yet been analysed in detail, and only a few have been described [11,16,17]. In a recent study by Hasan et al. [17], a missense mutation in the S protein was reported to be the third most frequent among viral proteins, following ORF1ab and N protein, by analysis of 371 viral genomes. In contrast, lower mutation rates of the S protein have been described elsewhere [14]. Although mutations in other viral proteins were not analysed in the present study, we identified high genetic diversity in the S protein, with missense mutation ranging from only one to 25 per sample. Therefore, associated with a nationwide pandemic in Bangladesh since late 2020, SARS-CoV-2 is suggested to have undergone rapid genetic evolution, acquiring various mutations. The presence of functional domains in the S protein underscores the importance of analysing mutations in this protein.

| Sample ID | No. of single nucleotide polymorphisms | No. of amino acid replacements | Amino acid replacement (substitution/deletion) | Known | Novel* |
|-----------|----------------------------------------|---------------------------------|-----------------------------------------------|--------|--------|
| D3        | 22                                     | 7                               | D614G, P681R, I1081V                         |        |        |
| D8        | 10                                     | 25                              | R158I, S162T, Q173H, V213L, Q218H, L244V, Q257C, A260T, R273T, D614G, I664K, T676N, A685S, Q1201R |        |        |
| C2        | 21                                     | 12                              | D614G, Q1180V, E1182D, N1187                  |        |        |
| C3        | 30                                     | 20                              | D614G, Q779H, V785F, T883A, F888S, G889A, Q891V, A892V, Q895R, G908C, S974L, D1260N |        |        |
| R2        | 12                                     | 9                               | L517P, A522S, P527L, T531N, D614G             |        |        |
| R5        | 1                                      | 1                               | D614G                                         |        |        |
| B4        | 27                                     | 18                              | V13D, L18S, P272S, K378R, Y380F, V382L, S477I, T500L, S157I, D614G, K1073N |        |        |
| B9        | 5                                      | 4                               | K1048R, T307S, E309Q, D614G                   |        |        |
| S3        | 12                                     | 5                               | R158K, E298K, D614G, V213L                    |        |        |
| S7        | 9                                      | 4                               | V13D, D614G, V620                               |        |        |
| T16       | 2                                      | 1                               | D614G                                         |        |        |
| T17       | 6                                      | 6                               | P541L, D614G, L578I, N574S, A771S             |        |        |
| M1        | 17                                     | 2                               | D614G                                         |        |        |
| M77       | 3                                      | 17                              | L141I, M153, A163V, N544Y, D574Y, R575S, V579L, D614G, Q675H, P809S |        |        |
| K6        | 2                                      | 2                               | D614G, Q675H                                  |        |        |
| Total     | 189                                    | 133                             | 80                                            |        |        |

del, deletion.

*Analysis includes the sequences available up to 15 January 2021.
infectivity of this variant [24,25]. A number of characteristic variants combined with the D614G substitution in the S glycoprotein have been identified previously and described as modifying the pattern of infection [9]. Among them, a D614G+Q675H variant was found in two samples in the present study.

In the present study, three novel substitutions were identified on the receptor-binding motif in RBD, suggesting their effect on alteration of viral infectivity. Although more evidence remains to be determined, the novel S protein mutations in SARS-CoV-2 in Bangladesh give rise to the possibility of a fitness advantage to target cells, which is likely to allow more successful person-to-person transmission. Another notable substitution was P681R, which is located adjacent to the FCS motif, which is highly conserved and important for pathogenesis of the virus [26,27]. On the other hand, loss of the FCS motif is related to attenuation of replication in cells and disease in vivo [26]. The P681R mutation was reported in only one study in Bangladesh [16], so appears to be still rare. Further surveillance may be necessary to monitor this mutation regarding its epidemiological and clinical impact. Mutations at N-glycosylation sites (amino acid 282 and 1098) were identified in the present study. Such glycosylation mutants had been reported previously and considered to be implicated in alteration of antigenic epitopes [9].

In conclusion, the present study revealed diverse nucleotide polymorphisms and amino acid substitutions in the S glycoprotein of SARS-CoV-2 in Bangladesh, suggesting rapid genetic evolution during the pandemic phase in late 2020. The epidemic of COVID-19 is still ongoing in Bangladesh, and the UK variant B.1.1.7 of SARS-CoV-2 was first identified in January 2021 [28], so continuous monitoring of genetic variation is necessary.

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Conflict of interest

The authors declare no conflicts of interest.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.nmni.2021.100889.

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