Dominant clade-featured SARS-CoV-2 co-occurring mutations reveal plausible epistasis: An in silico based hypothetical model

A. S. M. Rubayet Ul Alam1 | Ovinu Kibria Islam1 | Md. Shazid Hasan1 | Mir Raihanul Islam2 | Shafi Mahmud3 | Hassan M. Al-Emran4 | Iqbal Kabir Jahid1 | Keith A. Crandall5 | M. Anwar Hossain6,7

1Department of Microbiology, Jashore University of Science and Technology, Jashore, Bangladesh
2Division of Poverty, Health, and Nutrition, International Food Policy Research Institute, Bangladesh
3Department Genetic Engineering and Biotechnology, University of Rajshahi, Rajshahi, Bangladesh
4Department of Biomedical Engineering, Jashore University of Science and Technology, Jashore, Bangladesh
5Department of Biostatistics and Bioinformatics, Computational Biology Institute, Milken Institute School of Public Health, The George Washington University, Washington DC, USA
6Office of the Vice Chancellor, Jashore University of Science and Technology, Jashore, Bangladesh
7Department of Microbiology, University of Dhaka, Dhaka, Bangladesh

Abstract
Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has evolved into eight fundamental clades with four of these clades (G, GH, GR, and GV) globally prevalent in 2020. To explain plausible epistatic effects of the signature co-occurring mutations of these circulating clades on viral replication and transmission fitness, we proposed a hypothetical model using in silico approach. Molecular docking and dynamics analyses showed the higher infectiousness of a spike mutant through more favorable binding of G614 with the elastase-2. RdRp mutation p.P323L significantly increased genome-wide mutations (p < 0.0001), allowing for more flexible RdRp (mutated)–NSP8 interaction that may accelerate replication. Superior RNA stability and structural variation at NSP3:C241T might impact protein, RNA interactions, or both. Another silent 5′-UTR:C241T mutation might affect translational efficiency and viral packaging. These four G-clade featured co-occurring mutations might increase viral replication. Sentinel GH-clade ORF3a:p.Q57H variants constricted the ion-channel through intertransmembrane–domain interaction of cysteine(C81)-histidine(H57). The GR-clade N:p.RG203-204KR would stabilize RNA interaction by a more flexible and hypo-phosphorylated SR-rich region. GV-clade viruses seemingly gained the evolutionary advantage of the confounding factors; nevertheless, N:p.A220V might modulate RNA binding with no phenotypic effect. Our hypothetical model needs further retrospective and prospective studies to understand detailed molecular events and their relationship to the fitness of SARS-CoV-2.

KEYWORDS
clades, co-occurring mutations, COVID-19, fitness, infection paradox, SARS-CoV-2, virulence

1 | INTRODUCTION
Coronavirus disease (COVID-19) has caused 239,642,888 infection cases with 4,882,436 deaths worldwide until October 15, 2021 (https://coronavirus.jhu.edu/map.html). Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), the etiological agent of COVID-19 pandemic, has gained some extraordinary attributes that make it extremely infectious: High replication rate, large burst size, high stability in the environment, strong binding efficiency of spike glycoprotein (S) receptor-binding domain (RBD) with human angiotensin-converting
envelope 2 (ACE2) receptor, and additional furin cleavage site in S protein. In addition to those, it has proof-reading capability ensuring relatively high-fidelity replication. The virus contains four major structural proteins: spike glycoprotein (S), envelope (E), membrane (M), and nucleocapsid (N) protein, along with 16 nonstructural proteins (NSP1–NSP16) and seven accessory proteins (ORF3a, ORF6, ORF7a, ORF7b, ORF8a, ORF8b, and ORF10). Mutational spectra within the protein, which has direct roles in receptor binding and immunogenicity, thus viral immune escape, transmission, and replication fitness. Mutations in proteins other than spike could also affect viral pathogenicity and transmissibility, but the role of those occurring mutations, if not all, might have gained their relative evolutionary fitness through alterations of protein/RNA structure, function, and cross-talk (protein-protein or protein-RNA interaction). It is difficult to precisely determine how these co-occurring mutations, if not all, might have gained their relative evolutionary fitness through alterations of protein/RNA structure, function, and cross-talk (protein-protein or protein-RNA interaction). Overall, this in silico study aims to determine any plausible individual or epistatic impact of those mutants during replication in terms of viral entry and fusion, evasion of host cell lysis, replication rate, ribonucleoprotein stability, protein–protein interactions, translational capacity, and ultimately the probable combined effect on viral transmission and fitness.

2 | MATERIALS AND METHODS

2.1 | Retrieval of sequences and mutation analyses

This study analyzed 240,207 high-coverage (<1% Ns and <0.05% unique aa mutations) and complete (>29,000 nucleotides) genome sequences from a total of 32,142 sequences submitted to GISAID (https://www.epicov.org/epi3/frontend#18d55a) from January 1, 2020, to January 3, 2021 (Supplementary Material 1) for calculating the yearly percentage of the clades in 2020. We removed the non-human host-generated sequences during data set preparation. For the genome-wide mutation analysis, we initially selected 37,179 sequences from our whole data set before performing an alignment with MAFFT v7 (https://mafft.cbrc.jp/alignment/server/) against the Wuhan-Hu-1 (Accession ID: NC_045512.2) isolate as the reference genome. A python script (https://github.com/hrifdoy04/counting-mutations) was used to partition that data set into two subgroups (RdRp wild-type or “C” variant; 9,815; and mutant or “T” variant; 27,364) based on the presence of RdRp: C14408T mutation and then estimate the genome-wide single nucleotide polymorphisms (SNPs) for each strain (Supplementary Material 2). The SNP frequency was tested for significance with the Wilcoxon rank-sum test between the RdRp “C” and “T” variants implemented in IBM SPSS statistics 25. We chose this non-parametric test because a Kolmogorov-Smirnov test of mutational frequency showed these data do not fit a normal distribution (p < 0.001).

2.2 | Stability, secondary and three-dimensional structure prediction analyses of S, RdRp, ORF3a, and N proteins

We used DynaMut26 and FoldX 5.027,28 to determine the stability of both wild and mutated variants of N, RdRp, S, and ORF3a proteins. The following NCBI reference sequences were used as the wild and subsequently generated mutated aa sequence of N, RdRp, S, and ORF3a proteins, respectively, YP_009724397.2, YP_009725307.1, YP_009724390.1, and YP_009724391.1. We further used PredictProtein29 for analyzing and predicting the possible secondary structure and solvent accessibility of both wild and mutant variants of those proteins. The SWISS–MODEL homology modeling webtool30 was utilized for generating the three-dimensional (3D) structures of the RdRp, S, and ORF3a protein using 7c2k.1A, 6xr8.1A, and 6xdc.1A PDB structure as the template, respectively. We also used Modeller v9.2531 to generate the structures against the same templates to check the validity of SWISS–MODEL-derived structures. I-TASSER32 with default protein modeling mode was employed to construct the N protein 3D structure of wild and mutant type since there was no template structure available for the protein. The built-in structural assessment tools (Ramachandran plot, MolProbity, and Quality estimate) of SWISS–MODEL were used to check the quality of the generated structures.
Determination of the active sites affected by binding is a prerequisite for docking analysis. We chose aa residue 323 along with the surrounding residues (315–324) of RdRp and the residues 110–122 of NSP8 monomer as the active sites based on the previously reported structure.32 The passive residues were defined automatically, where all surface residues were selected within the 6.5Å radius around the active residues. The molecular docking of the wild and predicted mutated RdRp with the NSP8 monomer from the PDB structure 7C2K was performed using the HADDock2.4 To evaluate the interaction,34 The binding affinity of the docked RdRp-NSP8 complex was predicted using the PRODIGY.35 The number and specific interfacial contacts (IC) for each of the complexes were identified. The human neutrophil elastase (hNE) or elastase-2 (PDB id: 5A0C) was chosen for docking of the S protein, based on earlier reports.36 Here we employed CPORT37 to find out the active and passive protein-protein interaction residues of hNE. The S protein active sites were chosen based on the target region (594–638) interacting with the elastase-2. The passive residues of S protein were automatically defined as mentioned for RdRp-NSP8 docking analysis. Afterward, we individually docked wild (614D) and mutated (614G) S protein with the hNE using HADDock 2.4. The binding affinity of the docked complexes and the number and specific interfacial contacts (IC) for each of the complexes were identified. The structural stability of the protein complexes (RdRp-NSP8 and Spike-Elastase2) and their variants were assessed through the YASARA Dynamics software package. We used the AMBER14 force field for these four systems, and the cubic simulation cell was created with the TIP3P (at 0.997 g/L, 25°C, and 1 atm) water solvation model. The PME or particle mesh Ewald methods were applied to calculate the long-range electrostatic interaction by a cut-off radius of 8Å.39 We used the Berendsen thermostat to maintain the temperature of the simulation cell. The time step of the simulation was set as 1.25fs,40 and the simulation trajectories were saved after every 100 ps. Finally, we conducted the molecular dynamics simulation for 100 ns.41

Our analysis showed possible individual effect of a total of nine mutations in S, RdRp, ORF3a, N, 5'-UTR, leader protein (NSP1), and NSP3 found in the dominant clades G (15.2%), GH (20.8%), GR (32.6%), and GV (22.6%) of 2020 on viral replication cycle and transmission. We uniquely approached to dock spike with elastase-2 and RdRp with NSP8. Zeng et al.23 showed the links of these mutations toward possible epistatic effects on fitness using statistical analysis that duly suits our purpose of presenting how the mutations might play the combined roles. The overall epistatic interactions of the mutated proteins and/or RNA were depicted in Figure 1 as a hypothetical model.

3.1 Spike protein D614G mutation favors Elastase-2 binding

This study found interesting structural features of the S protein while comparing and superimposing the wild protein (D614) over mutated protein (G614). The secondary structure prediction and surface accessibility analyses showed a slight mismatch at the S1–S2 junction (681PRRAR*686), where serine at 686 (S686) was found covered in G614 and exposed to the surface in D614. However, S686 in both G614 and D614 were exposed to an open-loop region to have possible contact with the proteases (Figure S1). However, further investigation on the aligned 3D structures showed no conformational change at the S1–S2 cleavage site (Figure 2C). We also observed no structural variation in the surrounding residues of the protease-targeting
The predictive 3D models and structural assessment of D614 and G614 variants confirmed the cleavage site at 815–816 of the S2 subunit (812PSKR↓S816) or S2′ had no structural and surface topological variation (Figure 2D,E). Instead, the superimposed 3D structures suggested a conformational change in the immediate downstream region (618TEVPVAIHADQLTPT632) of the 614th position of mutated protein (G614) that was not observed in D614 variants (Figure 2A,B).

Several experiments suggested that mutated (G614) protein contains a novel serine protease cleavage site at 615–616 that is cleaved by host neutrophil elastase. The level of this elastase at the site of infection during inflammation will facilitate the host cell entry for G614.\(^{36,49,50}\) The elastase restrictively cuts valine at 615 due to its valine-dependent constriction of catalytic groove.\(^{51}\) The current sequence setting surrounding G614 (P6−610VLYQGVLNCTEv620,P′S) showed higher enzymatic activity on the spike,\(^{36}\) which cannot be entirely aligned with the previous works on the sequence-based substrate specificity of elastase-2.\(^{22}\)

However, the first non-aligned residue of the superimposed G614, located at the P′4 position (T618), may also be essential for binding with the elastase-2, and further down the threonine (T) at 618, the residues may affect the bonding with the respective aa of elastase-2 (Figure 2A,B). This changed conformation at the downstream binding site of G614 may help overcome unfavorable adjacent sequence motifs around G614 residue. Therefore, the simultaneous or sequential processing of the mutated S protein by TMPRSS2/Furin/Cathepsin and/or elastase-2 facilitates a more efficient SARS-CoV-2 entry into the host cells and cell–cell fusion.\(^{36,49,50}\)

This study further observed the possible association of the S protein with elastase-2 and found an increased binding affinity in the case of G614 (Table 1). Hence, the active sites of the mutated protein interacted efficiently with more aa of elastase-2 (Table 2), possibly providing a better catalytic activity, as shown by Hu et al.\(^{36}\) The mutation may induce a better structural configuration of the elastase-2 cleavage site of mutated spike protein for an easier and more accessible enzymatic cleavage (Figures 2A,B and 3). The efficient cleaving of this enzyme, although located in an upstream position of the S1–S2 junction, may assist in releasing S1 from S2 and change the conformation in a way that later helps in cleaving at the S2′ site by other protease(s) before fusion.\(^{53,54}\)

The complex of mutated spike protein and the elastase-2 was more flexible than the wild spike–elastase complex, and the interactions with enzyme were also different, as shown in root-mean-square deviation (RMSD) between the complexes (Figure 4).
FIGURE 2  (See caption on next page)
This G\textsuperscript{614} aa replacement may destabilize the overall protein structure (Table 2 and Figure 2A,B), and the deformed flexible region at or near G\textsuperscript{614} proves this destabilizing change (Figures 2F and S3). The S1 will release from S2 more effectively in G614 protein by increasing the chance of 1RBD-up conformation due to breakage of both intra-and interprotomer interactions of the spike trimer and symmetric conformation will increase binding potential with an ACE2 receptor and increase antibody-mediated neutralization.\textsuperscript{55,56} Our analyses provided the in silico proof of this fact by showing that the mutated protein was more flexible than the wild-type protein by missing a hydrophobic interaction between G\textsuperscript{614} and Phe\textsuperscript{592} (Figure 2G,H). The overall structural flexibility may assist the mutated S protein by providing elastase-2 a better binding space and attachment opportunity onto the cleavage site (Figure 3A,B), thus providing a more stable interaction that increases the credibility of an efficient infection (Figure 1).

### Table 1

| Variables                      | types |
|-------------------------------|-------|
|                               | RdRp/NSP8 | Spike–Elastase |
| HADDOCK score                 | Wild   | Mutant       |
|                               | -82.2 ± 7.8 | -118.3 ± 2.5 |
|                               | -43.0 ± 8.9 | -61.9 ± 4.5 |
| ΔG (kcal mol\textsuperscript{-1}) | Wild   | Mutant       |
|                               | -10.6  | -10.5        |
|                               | -13.3  | -13.7        |
| K\textsubscript{d} (M) at 37.0°C | Wild   | Mutant       |
|                               | 3.5E\textsuperscript{-08} | 3.9E\textsuperscript{-08} |
|                               | 4.5E\textsuperscript{-10} | 2.3E\textsuperscript{-10} |

| Number of interfacial contacts (ICs) per property | Wild                                      | Mutant                                      |
|--------------------------------------------------|-------------------------------------------|---------------------------------------------|
|                                                  | Charged-charged (5); charged-polar (9);   | Charged-charged (17); charged-polar (22);   |
|                                                  | charged-apol (15); polar-polar (2);      | charged-apol (32); polar-polar (5); polar-    |
|                                                  | polar-apol (16); and apolar-apol (21)    | apolar-apol (31); and apolar-apol (23)      |
|                                                  |                                           |                                             |
| Associated amino acids of Elastase-2 with possible docking interactions (for spike) or NSP8 (for RdRp) | Wild                                      | Mutant                                      |
|                                                  | P323: Asp (112), Cys (114), Val (115)    | P323: Asp (112), Cys (114), Val (115)       |
|                                                  | and Pro (116)                            | and Pro (116)                               |
|                                                  |                                           |                                             |
|                                                  | 605 (Ser) and 607 (Gln): 36 (Arg); 618    | 614 (Gly): 101 (Val); 618 (Thr): 181 (Arg),|
|                                                  | (Thr): 199 (Phe); 619 (Glu): 199 (Phe),  | 223–226 (Val, Arg, Gly, Gly); 619 (Glu): 103|
|                                                  | Cys (227); 620 (Val): 198 (Cys), 225:227  | (Leu), 181 (Arg), 222–225 (Phe, Val, Arg, Gly),|
|                                                  | (Gly, Gly, Cys)                          | 236 (Ala); 620 (Val): 223–227 (Val, Arg,|
|                                                  |                                           | Gly, Gly, Cys)                             |

### Figure 2

Diverse structural and stability comparisons of the wild and mutant spike protein. Structural superposition of wild and mutant spike proteins (A) and (B); conformation in S1–S2 (C) and S2’ sites (d) and (E); representation of vibrational entropy energy change on the mutant type structure (F); and interatomic interaction prediction of both wild (G) and mutant (H) types. For Figure (A)–(E), the gray and yellow colors represent the wild and mutant protein, respectively. (A) The downstream (617–636) of D614G in wild (green) and mutant (red) S protein was focused. Overlapping the wild (D\textsubscript{614G}) and mutant (G\textsubscript{614G}) S protein showed a conformational change in the 3D structures. (B) However, the conformational change is in the loop region (618–632) of the proteins thus may potentially play a role in interacting with other proteins or enzymes, such as elastase-2, as we focused on in this study. (C) No change was found in the S1–S2 cleavage site (685–686), depicted in blue color, of the wild and mutant protein. (D) Surface and (E) cartoon (2°) structure of the superimposed wild and mutant proteins where the S2’ (pink) is situated in the surface region and does not show any change in accessibility in the residual loop region. (F) The mutant (G\textsubscript{614G}) protein showed higher flexibility in the C\textsubscript{614} (sticks) and its surroundings (red). The intramolecular interaction determined the overall stability of the (G) wild and (H) mutant structure where C\textsubscript{p} of D\textsuperscript{614E} (aspartic acid at 614; green stick modeled) had two hydrophobic interactions with the benzene rings. These intramolecular contacts stabilize the S protein of wild-type. The missing of this bond destabilizes the mutant (G\textsubscript{614G}) protein. The mutant protein has glycine at 614, which has less chance of interacting with neighboring aa due to its shorter and nonpolar R-group. The color code representing the bond type is presented in each (G) and (H).

### 3.2 Increased flexibility of RdRp–NSP8 complex: Compromise proof-reading efficiency with replication speed

The binding free energy (ΔG) of the RdRp–NSP8 complexes have been predicted to be ~10.6 and ~10.5 kcal mol\textsuperscript{-1}, respectively, in wild
**Table 2** Assess the effect of mutations on structural dynamics of NSP-12/RDRP, Spike, NS3, and N Protein of SARS-CoV-2 using DynaMut

| Protein name | Mutation with position | ΔΔG DynaMut kcal/mol | ΔΔG ENCoM kcal/mol | ΔΔG mCSM kcal/mol | ΔΔG SDM kcal/mol | ΔΔG DUET kcal/mol | ΔΔGFoldX kcal/mol | Results* | ΔΔSVib ENCoM kcal/mol | ΔΔSVib ENCoM mol<sup>-1</sup>K<sup>-1</sup> |
|--------------|------------------------|----------------------|-------------------|------------------|-----------------|-----------------|-----------------|---------|----------------------|------------------|
| RdRp         | P323L                  | 1.054                | −0.441            | −0.264           | 0.700           | 0.118           | −0.733          | Stabilizing | −0.551               |                   |
| Spike        | D614G                  | −0.769               | +0.408            | −0.492           | 2.530           | 0.195           | +0.289          | Destabilizing | 0.510                |                   |
| ORF3a        | Q57H                   | 0.275                | −0.128            | 0.788            | 0.520           | −0.464          | −1.438          | Stabilizing | −0.160               |                   |
| N Protein    | RG203-04KR             | −                    | −                  | −                | −                | −3.4262         | Highly Destabilizing |                   | −                   |
| N protein    | A220V                  | 0.109                | 0.458             | −0.586           | −1.460          | −0.567          | +1.6            | Stabilizing | −0.572               |                   |

Note: The value of ΔΔG < 0 indicates that the mutation causes destabilization and ΔΔG > 0 represents protein stabilization. For ΔΔSVibENCoM, positive and negative value denotes the increase and decrease of molecular flexibility, respectively.

Abbreviation: SARS-CoV-2, severe acute respiratory syndrome coronavirus 2.

*The final result of the stability for each protein was determined based on the intramolecular interactome analysis.

**Figure 3** The molecular docking of wild and mutant with elastase-2. Both the (upper figure) wild (D<sub>614</sub>) and (lower figure) mutant (G<sub>614</sub>) version of S protein was shown in golden color, whereas the elastase-2 docked to D<sub>614</sub> and G<sub>614</sub> in blue and green color, respectively. The enlarged views of the docked site were shown in separate boxes. (A) The possible docked residues (stick model) on the wild S protein (warm pink) and elastase-2 (green) are 618(Thr)–619(Glu)–620(Val) and 198(Cys)–199(Phe)–225:227 (Gly, Gly, Cys), respectively. The aspartic acid at 614 is 17.3°A far away from the valine (101), apparently the nearest aa of elastase-2 to the cleavage site (615–616). (B) The possible interacting residues (stick model) on the mutant S protein (blue) and elastase-2 (warm pink) are 614(Gly)–618(Thr)–619(Glu)–620(Val) and 101(Val)–103(Leu)–181(Arg)–222:227(Phe, Val, Arg, Gly, Gly, Cys), and 236 (Ala), respectively. In this case, the glycine at 614 is 5.4°Afar from the valine (101), the nearest aa of elastase-2 to the cleavage site (615–616).
(P323) and mutated (L323) type, which suggests a more flexible interaction for the mutated protein (Table 2). The increased number of contacts found in the L323–NSP8 complex (Table 1) was possibly due to slightly more hydrogen bonds, which had no considerable impact on protein flexibility (Figure 4D). Our analyses identified that proline (P323) or leucine (L323) of RdRp can interact with the aspartic acid (D112), cysteine (C114), valine (V115), and proline (P116) of NSP8 (Table 1 and Figure 5). RdRp binds with NSP8 in its interface domain (from residues alanine: A250 to arginine: R365), forming positively charged or comparatively neutral "sliding poles" for RNA exit, and enhance the replication speed probably by extending the RNA-binding surface in that domain area.57,58 Molecular dynamics of the mutated RdRp–NSP8 complex supported this by showing a more expanded surface area in the interacting site (Figure 4B) and maintained integrity throughout the simulation (Figure 4C). Besides, we did not find any interaction of NSP8 with the zinc-binding residues (H295, C301, C306, and C310) of the RdRp protein (Table 1 and Figure 5).59,60 Therefore, the P323L mutation within the RdRp interface domain’s conserved site may only affect the RdRp–NSP8 interaction without changing metal binding affinity.

The results from six state-of-the-art tools of protein stability suggested that mutated (L323) protein cannot be concluded as "stable," only because of ambiguous ΔΔG estimates (Table 2); instead, the interaction with the adjacent aa mainly defined the stability.61 The superimposed 3D structures and secondary structure analyses showed no deviation in the loop/turn structure of the mutated protein, even though hydrophobic leucine (L323) was embedded (Figures S1 and S4). The mutation stabilized the L323 structure to some extent making the protein more rigid and bound less firmly with the NSP8 by expanding the interacting region.

Together, these structural variations may increase the replication speed by helping exit the processed RNA genome from the RdRp

**FIGURE 4** Molecular dynamics of spike-elastase2 and RdRp-NSP8 complex (A) Both the wild and mutated spike protein had lower RMSD profile till 60 ns, then it rose and maintained a steady state. Although the spike protein had a higher degree of deviation in the RMSD profile than RdRp, they did not exceed 3.0 Å. The RMSD demonstrated that mutant and wild RdRp protein complex has an initial rise in RMSD profile due to flexibility. Therefore, both RdRp complexes stabilized after 30 ns and maintained a steady peak. The wild-type RdRp complex had a slightly higher RMSD peak than mutant RdRp, indicating the more flexible nature of the wild-type. (B) The spike protein complex had a similar SASA profile, did not change its surface volume, and maintained a similar trend during the whole simulation time. The higher deviation of SASA indicates that mutant and wild-type RdRp had a straight line. Still, the mutant structure had a higher SASA profile, indicating the protein complex had enlarged its surface area. Therefore, the mutation in RdRp protein leads to more surface area expansion than wild types as their average SASA value had a significant difference. (C) Mutated spike exhibits a little more Rg profile than the wild-type, which correlates with the comparative labile nature of the mutant. The higher level of Rg value defines the loose packaging system and mobile nature of the protein systems. The mutant RdRp had a lower level of fluctuations and maintained its integrity during the whole simulation time. The wild-type RdRp complexes had higher deviations and more mobility than the mutant complex. (D) Any aberration in hydrogen bond numbers can lead to higher flexibility. Therefore, the mutant and wild spike proteins exhibit the same flexibility in terms of H-bonding. The mutant RdRp protein had more hydrogen bonding than the wild types, but they did not differentiate too much, and a relatively straight line was observed for the protein. RMSD, root-mean-square deviation
transmembrane helical configuration (Figure S1). Aligned 3D structures have also shown no variation of TM1 in the monomeric state (Figure 6A). The mutant (H$_{57}$) protein has a nonsignificant increase in structural stabilization and a minimal decrease in molecular flexibility (Table 2 and Figure S5). This higher stability is because of the weak ionic interaction of H$_{57}$C$_3$ with the sulfur atom of cysteine (C$_81$) that is present in TM2 and the hydrogen bond of terminal N$_2$ of lysine (K$_{61}$) with one of the endocyclic nitrogens of H$_{57}$ (Figure 6B). The Q$_{57}$ in wild-type protein forms the major hydrophilic constriction within the ORF3a channel pore. Thus, further favorably increasing constrictions within the H$_{57}$ protein channel pore due to diagonal H$_{57}$ (TM1)–C$_{81}$ (TM2) ionic interaction (Figure 6B) and the replacement of charge-neutral glutamine with a positively charged histidine in the selectivity filter may reduce the passing of positive ions, such as Ca$_{2+}$, Na$_{+}$, and K$^+$, by either electrostatic repulsion or blocking. This speculation for ORF3a mutated protein was supported by another study showing the reduction of ion permeability of Na$_{2+}$ and Ca$_{2+}$ through the H$_{57}$; however, that decrease was not found statistically significant ($p > 0.05$).

The decreased intracellular concentration of cytoplasmic Ca$_{2+}$ ions potentially reduces caspase-dependent apoptosis of the host cell, mainly supporting viral spread without affecting replication, as shown in Figure 1. Moreover, the ORF3a can drive necrotic cell death wherein the permeated ions into cytoplasm and the insertion of ORF3a as viroporin into lysosome play vital roles. The H$_{57}$ mutant may thus decrease pathogenicity and symptoms during the early stages of the infection, that is, reducing “cytokine storm” in the host. Besides, ORF3a was shown to affect inflammasome activation, virus release, and cell death, as detailed by Castaño-Rodriguez et al. That the deletion of ORF3a reduced viral load and morbidity in animal models.

Even though similar proteins of ORF3a have been identified in the sarbecovirus lineage infecting bats, pangolins, and humans, only one pangolin derived strain from 2017 in Guangxi, China contains H$_{57}$ residue as shown by mutation analyses (Figure S5), and also reported by Kern et al. A possible explanation behind that presence could be the more adaptive nature of the virus toward reverse transmission by being less virulent, that is, from human to other animals, as observed in recent reports.

### 3.4 | N protein mutation: Augmenting nucleocapsid stability and exerting miscellaneous effects

Our study has observed that the combined mutation (N: p.RG203→204KR) causes no conformational change in secondary and 3D structures (Figures S1 and Figure 7, respectively) of the conserved SR-rich site (184 → 204) in the linker region (LKR: 183 → 254) of the N protein (Figure S7). But there is a minor alteration among buried or exposed residues (Figures S7C and S8). The superimposed 3D structures showed structural deviation, rather at $^{235}$ESKMSGKGGQQQQQTTVT$^{247}$ of the LKR

### 3.3 | Q$_{57}$H substitution in ORF3a viroporin: The roles of decreased ion permeability

Our study found that the replacement of glutamine (Q$_{57}$) with positively charged histidine (H$_{57}$) at aa position 57 of ORF3a transmembrane region 1 (TM1) does not change secondary groove structure more swiftly (Figure 1). The increasing replication speed might be due to the perturbation of interaction between RdRp and NSP8, or less possibly, the complex tripartite interactions (RdRp, NSP8, and NSP14) responsible for the speculated decrease of proof-reading efficiency. Thus, RdRp mutants might increase the mutation rate by a trade-off between high replication speed and low fidelity of the mutated polymerase. The Wilcoxon rank-sum test revealed that the frequency of mutation (median = 8) in L$_{323}$ mutants ($n = 27,364$) is significantly higher ($p < 0.0001$) than the frequency (median = 6) of wild-type (P$_{323}$) strains ($n = 9815$). This increased mutation rate in L$_{323}$ mutants can surpass the constant proof-reading fidelity, which might help adapt more quickly in adverse climatic conditions, evade the immune response, and survive within different selective pressure.
Impedance to form particular SR-motif due to RG→KR mutation might disrupt the phosphorylation catalyzed by host glycogen synthase kinase-3. After the virus enters the cell, this synchronized hypo-phosphorylation of KR203-204 protein should make the viral ribonucleoprotein (RNP) unwind in a slower but more organized fashion that might impact translation and immune-modulation. In KR203-204, replacing glycine with arginine may increase the nucleocapsid (N protein–RNA complex) stability by forming stronger electrostatic and ionic interactions due to increased positive charge. Besides, the more disordered orientation of the downstream LKR site and the highly destabilizing property of KR203-204 may assist in packaging a stable RNP. N protein also utilizes the intrinsically disordered dynamic linker region (LKR) that controls its affinity toward M protein, self-monomer, 5′-UTR, and cellular proteins. The phosphorylation at the LKR site may play an essential role in regulating these interactions. These plausible interactions and impacts upon mutations are depicted in Figure 1.

3.5 Silent mutations may not be silent

The C241T of 5′-UTR, a single nucleotide "silent" mutation, is located at the UUCGU pentaloop part of the stem-loop region (SLR5B). This pentaloop of 5′-UTR remains unchanged and maintains a particular structure with a potential role in viral packaging. The RNA secondary structures predicted no change in the 241T structure (Figure S8A). However, C241T is present just upstream to the ORF1a start codon (266–268 position) and may be involved in differential RNA binding affinity to the ribosome and translational factors.

In the case of multi-domain NSP3 (papain-like protease), we have observed superior stability of the RNA after gaining the synonymous mutation 3037C>T (C318T) where wild and mutant RNA structure...
translational modifications as a part of protein regulation. This normality results in low translation efficiency affecting post-translational elongation that generally contributes to a range of abnormalities centered replication complex. The N protein can also affect the membrane stability through an uncharacterized interaction with the M protein, ultimately producing more stable virion particles.

A more robust N protein would help to modulate the replication and transcription. The mutant N protein may impact viral replication and transcription, like other coronaviruses, through binding with the NSP3 protein, which linked the RdRp-centered replication complex. The N protein can also affect the membrane stability through an uncharacterized interaction with the M protein, ultimately producing more stable virion particles.

A more robust N protein-RNA complex provokes a slower intracellular immune response. At the same time, the mutated virus can remain highly contagious and aggressive because of the simultaneous presence of G clade featured S protein and RdRp mutations (Figure 1). The rapid within-host replication and modified replication dynamics might be correlated with the fitness of G clade strains.

The epistatic effects of the co-occurring mutations, simultaneous multisite variations in the same or different proteins or genes, have provided new insights into the dynamic epistatic network by employing differential molecular interactions. The epistatic effects of the mutations were reported to control viral fitness and virulence through modulating the replication cycle and virus-host interactions, as observed before for Influenza and Ebola virus. The co-occurring mutations of the major SARS-CoV-2 clades discussed in this study contained epistatic links and positive selection pressure except for the synonymous mutations. Observable (https://observablehq.com/) also presents those mutated sites as positively selected.

We speculated no interlinked functional relationship between p:D614G of the S protein and p:P323L of the RdRp, two important G clade featured co-occurring mutations (Figure 1). The sequence-based prediction showed no potentially significant epistatic link as well. These seemingly unrelated mutations can cumulatively escalate the infectiousness of the virus because of a higher viral load and shorter burst time. The S:p.D614G might assist in rapid entry into the host cells followed by quick dissemination, and the RdRp:p:P323L may instead boost the replication by a faster RNA processing (exit). NSP3 is a scaffolding protein for the replication-transcription complex, and the possible change in its structure may affect the overall dynamics of viral replication.

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hiding the virus from host cellular immunity of the host and increasing stability in the environment.

Conversely, we have not found any literature for even other coronaviruses that showed ORF3a: p.Q57H correlating with the rest of the co-occurring mutations. The H57 mutant, possibly linked to the mild or asymptomatic cases, may allow the silent transmission and increase the chance of viral spread by lowering the activation of the inflammatory response (Figure 1), such as reduced viral particle release and cytokine storm.\(^{21,108,109}\)

According to our hypothesis and Wolf et al. (2021), the social interventions on movement could also play a role in disseminating these G, GH, and GR clades at the early and later pandemic phases well.\(^{110}\)

The GV strains featuring an A222V mutation in the S protein have probably no effect on the viral transmission, severity, and antibody escape due to its structural position; rather, super-spreading founder events might be the reason behind its faster spreading.\(^{11,111}\)

The A220V mutation stabilized the mutated N protein’s linker region (Table 2) might affect RNA binding affinity;\(^{20}\) however, different mutations at positions 220 in N found in other major lineages showed no phenotypic consequence.\(^{17}\) There was also no epistatically linked pairing between GV clade co-occurring mutations.\(^{22,23}\) Altogether, the co-occurring mutations of GV strains might not affect transmission fitness.

Vaccine inequity, immunocompromised patients, and a tremendous number of hosts are now frequently introducing variants with mutations in the RBD of the spike protein. The introduced mutations in a lineage (a variant of concern/interest) on top of the original clade-featured ones in the genome might play the most crucial role in increasing transmission fitness and a slightly reduced neutralization to antibodies by showing epistatic effects.\(^{112,113}\) Future studies are necessary to investigate the roles of the “mutation package” in each of these variants of concern/interest.

## 4 | CONCLUSION

In 2020, the course of the COVID-19 pandemic was dominated by the G, GH, GR, and GV clades. The G clade-featured co-occurring mutations might increase the viral load, alter host immune responses, and modulate intrahost virus genome plasticity that raises the speculation of their potential role in continuous transmission. The GR and GH clade mutant with the signature mutation, respectively, in nucleocapsid and ORF3a protein, might contribute to the host’s immune response and viral transmission. The GV strains, however, could have spread quickly by superspreading events with no apparent epistatic effect. Therefore, the fitness of SARS-CoV-2 may increase in terms of replication and transmission where viral strains are always giving their spread capacity within a population the top priority by calibrating the infection cycle. However, further in vivo and ex vivo studies and more investigations are required to prove and bolster this hypothesis.

## ACKNOWLEDGMENTS

We want to acknowledge the team at GISAID and all contributing authors, who generated and submitted sequence data, for creating the SARS-CoV-2 global database. Jashore University of Science and Technology provided the funding for the research. We want to thank the Microbial Genetics and Bioinformatics Laboratory of the University of Dhaka for supporting high-performance computer access. We especially appreciate the efforts of M. Shaminur Rahman and Spencer Mark Mondol in protein structure prediction and stability analyses.

## CONFLICT OF INTERESTS

The authors declare that there are no conflict of interests.

## AUTHOR CONTRIBUTIONS

Iqbal Kabir Jahid, Ovinu Kibria Islam, and A. S. M. Rubayet Ul Alam hypothesized about the work. A. S. M. Rubayet Ul Alam performed the sequence analysis part after Ovinu Kibria Islam and Md. Shazid Hasan compiled the data set. The Python coding, structural (both RNA and protein), and protein docking were done by A. S. M. Rubayet Ul Alam. MSH predicted protein structure in I-TASSER and performed stability analysis in DynaMut. Shafi Mahmud performed the molecular dynamics study. Hassan M. Al-Emran and Mir Raihanul Islam performed the statistical analysis. Hassan M. Al-Emran then reviewed and organized the manuscript expertly. Iqbal Kabir Jahid, Keith A. Crandall, and M. Anwar Hossain supervised, suggested, and revised the write-up to produce the final draft.

## DATA AVAILABILITY STATEMENT

All the sequence and structural data were taken from the GISAID (https://www.gisaid.org/) and RCSB PDB (https://www.rcsb.org/), as mentioned in the methodology section. We provide all the necessary information, such as accession numbers and the date-based data source for helping readers and reviewers to check the authenticity of the work. The data that support the findings of this study are available in GISAID at https://www.gisaid.org/. These data were derived from the resources available in the public domain: GISAID, https://www.gisaid.org/.

## ORCID

A. S. M. Rubayet Ul Alam \(\text{http://orcid.org/0000-0001-9295-9865}\)

Ovinu Kibria Islam \(\text{https://orcid.org/0000-0002-1114-3125}\)

Md. Shazid Hasan \(\text{https://orcid.org/0000-0002-2021-4693}\)

Iqbal Kabir Jahid \(\text{https://orcid.org/0000-0003-0717-0806}\)

M. Anwar Hossain \(\text{http://orcid.org/0000-0001-9777-0332}\)

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SUPPORTING INFORMATION
Additional supporting information may be found in the online version of the article at the publisher’s website.

How to cite this article: Alam ASM Rubayet UI, Islam OK, Hasan MS, et al. Dominant clade-featured SARS-CoV-2 co-occurring mutations reveal plausible epistasis: An in silico based hypothetical model. J Med Virol. 2022;94:1035-1049. doi:10.1002/jmv.27416