Severe Acute Respiratory Syndrome Coronavirus-2 (SARS-CoV-2), causative agent of Coronavirus Disease 19 (COVID-19), has spread throughout the world since its first emergence in Dec. 2019 in Wuhan, China. As anticipated, SARS-CoV-2 evolution is most likely through selection, perhaps conferring enhanced fitness and/or infectivity. Numerous reports detail the molecular identification of D614G mutation in Spike protein (S-protein) (Korber et al. 2020; Koyama et al. 2020; Maitra et al. 2020) and are experimentally characterizing the functional impact upon the viral life cycle (Li et al. 2020; Luthy and Kistler 1989). Regardless of how D614G arose, it is clear that this variant has a distinct phenotype. With the report of D614G, it was hypothesized that viruses containing G614 were more infectious than D614 viruses (Grubaugh et al. 2020; Korber et al. 2020). Experimental evidence quickly confirmed this hypothesis (Korber et al. 2020; Li et al. 2020). However, a report from COVID-19 Genomics UK Consortium (Report #9 - 25th June 2020) showed G614 virus has grown 1.22 times faster than the D614, but the statistical significance was low, indicating the role of other factors such as mutations in other genes.

To gain insight into the distribution of mutations in SARS-CoV-2 nonstructural proteins (nsps) and structural proteins, we analyzed protein sequences (n = 7232) from the United States (n = 6302), Europe (n = 420), China (n = 104), and India (n = 406), and determined the mutations with respect to Wuhan-Hu-1 isolate (NCBI Reference Sequence: NC_045512.2). A Circos diagram showing nonsynonymous mutations in select viral proteins (nsp8, nsp12, nsp13, nsp14, and S-protein) is shown in Fig. 1a. SARS-CoV-2 has evolved by developing mutations in different viral proteins. While mutations in 5 proteins is shown in Fig. 1a, mutations in other viral proteins were also identified in this analysis. The data also revealed that some mutations were widespread, while other mutations appeared to be restricted geographically. Most strikingly, mutation P323L in nsp12 (an RNA-dependent RNA polymerase or RdRp) and D614G in S-protein co-evolved throughout the world. Other mutations,
such as P504L and Y541C in nsp13 (helicase) were more prevalent in the USA (Fig. 1a).

To further analyze the coevolution of P323L and D614G, we determined the mutation frequency of P323L and D614G in SARS-CoV-2 nucleotide sequences from the United States (n = 7233) over a six-month period (January, 2020 - June, 2020). The results of the temporal analysis of the mutation frequency of D614G, C241U and P323L in the virus isolated from the USA patients. The frequency was calculated as the number of sequences containing mutation divided by total number of sequences, then multiplied by 100. For example, a T to C mutation frequency = (Number of T mutations / (Number of T mutations + number of regular C)) × 100. Panel c. This panel shows temporal change in frequency of I156V and M129I in nsp8 (orange filled triangle and red filled square, respectively), F233L in nsp14 (blue filled circle), P504L and Y541C in nsp13 (violet filled diamonds and orange filled hexagons, respectively) in addition to D641G, P323L and C241U from panel a. Plots in panel b and c were generated by Matplotlib Python script.

By the end of June 2020, 91.5% of patients from the U.S. had the D614G mutation (compared to Wuhan-Hu-1 isolate, GenBank ID NC_045512.2). Since SARS-CoV-2 emerged in
Asia, a significant increase in the D614G mutation in the U.S. indicates that this mutation may be associated with enhanced infectivity of the virus outside of China. The structure of G614 S-protein has not been reported yet. However, in the cryoEM structure of the full-length S-protein trimer, D614 is located at the interface of two monomers and forms inter- and intramonomer contacts (Fig. 1e) (Wrapp et al. 2020). D614 forms a salt-bridge with K854 and an H-bond with the backbone of V860 of the neighboring monomer. D614 also forms an intramonomer salt-bridge with R646. A mutation of D614 to G614 should result in the loss of these interactions, which could alter the dynamics of S-protein conformational changes during SARS-CoV-2 infection.

Reported experimental data show that D614G mutation enhances the infectivity of SARS-CoV-2 virus. However, it is highly unlikely that a seemingly high infection rate of SARS-CoV-2 is solely the result of this mutation. A near 100% coexistence of P323L (nsp12) and C241U (5′-UTR) with D614G likely contributes to the viral replication, infectivity, or a combination of attributes that are complex and interplay with the host machinery. P323 is located in the Interface domain (residues 251–398) of nsp12 (Hillen et al. 2020). However, in the cryoEM structure of replication-transcription complex (RTC), consisting of nsp7, nsp8, nsp12 and nsp13, the Interface domain is packed against nsp8 (Chen et al. 2020). The role of nsp7 and nsp8 in the context of the RTC is not known, but they can serve as the processivity factor using the replicating complex of nsp12/nsp7/nsp8/RNA (Hillen et al. 2020; Posthuma et al. 2017) similar to that of thioredoxin in the replicating structure of T7 DNA polymerase (Doublie et al. 1998). P323 is located at the interface of nsp12 and nsp8 (Hillen et al. 2020; Kirchdoerfer and Ward 2019) and mutation P323L is expected to position the leucine side chain within interacting distance to F396, which mediates hydrophobic interactions involving L122 an N118 (Cβ) of nsp8 and T323 (Cy2) and L270 of nsp12 (Fig. 1f). Enhanced interaction between nsp12 and nsp8 would further improve the processivity of nsp12 and thereby help in the viral replication. C241 is located in the 5′UTR of the viral RNA, 25 nucleotides upstream of the AUG start codon. SHAPE analysis of viral RNA predicted it in the loop of SLSb (Sun et al. 2020). The SHAPE activity of C241 and neighboring residues in the loop are low in vitro probing than in vitro probing, suggesting these residues may be bound by protein or engaged in long-range RNA interactions under in vivo conditions, and thus are protected from 2′-OH modification in SHAPE analysis. The C241U may change the structure of the SL5 loop or affect protein binding for easy ribosomal scanning, which is favored in translation initiation.

The nsp8 mutation M129I correlated to D614G at ~2.4%. M129 is also located at the nsp8/nsp12 interface (Fig. 1g). It is positioned such that mutation M129I would result in increased hydrophobic interactions with nsp12 residues L388, A400, and V405. The increased hydrophobic interactions between nsp12 and nsp8 could also provide enhanced processivity of RNA synthesis by nsp12. Two nsp13 mutations P504L and Y541C are correlated to D614G by ~15%. P504 is at the surface of nsp13, whereas Y541 is located at the putative RNA-binding channel. From the available structures, a reliable role of P504L and Y541C mutations cannot be deduced. Similarly, the two positions (P233 and T372) of nsp14 are away from the 5′-5′-exoribonuclease or methyl-transferase sites. Currently available structure of SARS-CoV nsp14 in complex (Ma et al. 2015) with nsp10 does not provide a conclusive role of mutations at these sites.

In summary, from currently available structural, genetic and biochemical data, the higher infectivity of SARS-CoV-2 D614G mutation is not fully understood. However, from the coexistence of several mutations with relatively high frequency suggest that the infectivity of G614 virus mutation is not solely dependent on the entry. Remarkably, COVID-19 presents with a wide variety of symptoms, including neuropathology, delayed onset symptoms, and anosmia. While these coevolving mutations could impact viral fitness, the breadth and complexity of the clinical symptoms may also be associated with new mutations and adaptations. While there has been an unprecedented eruption of COVID-19 publications, clearly we have much to learn going forward.

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Author Contributions KS and SNB conceptualized the study; KS wrote the first draft. SRK, ATA, ANS and KS conducted genetic analyses; XH performed the experiments; SNB, TPQ and CLL, and AS edited the manuscript and contributed to understanding the pathogenicity of CoVs. All authors approved the final manuscript.

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