Concurrent mutations in RNA-dependent RNA polymerase and spike protein emerged as the epidemiologically most successful SARS-CoV-2 variant

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The D614G mutation in the Spike protein of the SARS-CoV-2 has effectively replaced the early pandemic-causing variant. Using pseudotyped lentivectors, we confirmed that the aspartate replacement by glycine in position 614 is markedly more infectious. Molecular modelling suggests that the G614 mutation facilitates transition towards an open state of the Spike protein. To explain the epidemiological success of D614G, we analysed the evolution of 27,086 high-quality SARS-CoV-2 genome sequences from GISAID. We observed striking coevolution of D614G with the P323L mutation in the viral polymerase. Importantly, the exclusive presence of G614 or L323 did not become epidemiologically relevant. In contrast, the combination of the two mutations gave rise to a viral G/L variant that has all but replaced the initial D/P variant. Our results suggest that the P323L mutation, located in the interface domain of the RNA-dependent RNA polymerase, is a necessary alteration that led to the epidemiological success of the present variant of SARS-CoV-2. However, we did not observe a significant correlation between reported COVID-19 mortality in different countries and the prevalence of the Wuhan versus G/L variant. Nevertheless, when comparing the speed of emergence and the ultimate predominance in individual countries, it is clear that the G/L variant displays major epidemiological supremacy over the original variant.

Common coronaviruses differ from a pandemic causing SARS-CoV-2 coronavirus that emerged in late 2019. In addition, as opposed to previous coronavirus outbreaks such as SARS-CoV-1 and MERS, the new viral strain is not characterized by a particularly high case fatality rate, but rather by unprecedented infectivity and epidemiological success. Indeed, the virus has so far largely resisted all attempts of eradication, despite stringent measures taken in many countries around the globe.

SARS-CoV-2 is an enveloped RNA virus. They have a large genome and a sophisticated life cycle which is still poorly understood. Coronavirus have a proofreading machinery, which confers a relatively stable genome, as
compared to other types of RNA\textsuperscript{2–6}. Given its proofreading capability of SARS-CoV-2, prevalent SARS-CoV-2 mutations leading to significant amino acid changes should be relatively rare\textsuperscript{7}. The SARS-CoV-2 transmembrane Spike glycoprotein (S-protein) is the most abundant protein of the viral envelope. It mediates attachment of the virus to the host cell surface receptors and a subsequent fusion between the viral and the host cell membranes to facilitate viral entry\textsuperscript{8}. The S-protein is a trimeric class I fusion protein, which exists in a metastable pre-fusion conformation, divided into two main functional subunits responsible for binding to the host cell receptor (S1 subunit) and fusion of the viral and cellular membranes (S2 subunit, Fig. 1a). The trimer exists mainly in two distinct conformational states resulting from an opening of the receptor binding domain (RBD) in the S1 subunit: ‘up’ or ‘down’ (Fig. 1a). The ‘up’ conformation exposes the RBD to cell surface receptors of the host, which would otherwise be buried (‘down’ conformation) at the interface of each monomer\textsuperscript{9}. This conformational change is required for proteolytic processing and/or fusion of the S2 domain with the host cell membrane. It is now understood that a variety of proteases, including the PC family of proteases, trypsin-like proteases and cathepsins are able to cleave the spike protein\textsuperscript{10}.

The S-protein D614G mutation has received attention given that it could potentially alter the viral attachment, fusogenicity and/or immunogenicity. The mutation has been suggested to increase infectivity\textsuperscript{11} and/or the case fatality rate\textsuperscript{12}. More recent studies have demonstrated that the G614 variant indeed enhances viral entry into cells\textsuperscript{13–17}.

SARS-CoV-2 employs a multi-subunit replication/transcription machinery\textsuperscript{18}. A set of non-structural proteins (nsp) produced as cleavage products of the ORF1a and ORF1b viral polyproteins assemble to facilitate viral replication and transcription, making this machinery a potential target for therapeutic intervention\textsuperscript{19}. RNA-dependent RNA polymerase (RdRp or nsp12) is a key player in the synthesis of viral RNA. The crystal structure of SARS-CoV-2 nsp12 in complex with its nsp7 and nsp8 co-factors underlines the central role these non-structural proteins have in the replication and transcription of the virus\textsuperscript{20,21}. The P323L mutation of the RdRp has received less attention; however, it has been suggested that this mutation might alter viral proofreading and thereby lead to an increased down-stream mutation rate\textsuperscript{22}.

In this study, we investigated the emergence of a SARS-CoV-2 variant with concomitant mutations in the S-protein and RdRP-polymerase. Both mutations are in strategically relevant sites of the respective proteins and hence candidates to alter the biology of the virus. We show that the G614/L323 viral variant, but not the individual G614 or L323 mutants, is epidemiologically highly successful and replaced the original D614/P323 variant.

Results

S-protein D614G increases cell transduction of pseudotyped lentivectors. Cell entry of SARS-CoV-2 depends on the binding of viral S-proteins to cellular receptors and subsequent priming by proteases of the host cell\textsuperscript{23}. Our first aim was to establish cell lines that consistently express the angiotensin-converting enzyme-2 (ACE2) receptor, which is recognized by SARS-CoV-2; and relevant proteases, such as transmembrane serine protease 2 (TMPRSS2), which cleaves the S-protein. As seen in Fig. 2a,b, neither HeLa cells nor A549 cells expressed relevant amounts of ACE2 mRNA. Similarly, mRNA levels of TMPRSS2 were almost undetectable. However, both cell lines express low to intermediate gene levels of furin and moderately high levels of cathepsin L (Fig. 2a,b). Furin is able to cleave the spike protein of SARS-CoV-2, while cathepsin L is able to cleave S-proteins from both SARS-CoV-1 and SARS-CoV-2\textsuperscript{11}. In order to establish cell lines permissive for S-protein pseudotyped lentivectors, we transduced both HeLa and A549 cells with either ACE2 alone, TMPRSS2 alone or in combination.

To understand the impact of the S-protein D614G mutation on infectivity, we produced S-protein pseudotyped lentivectors. Specifically, we used full length and C-terminally truncated D614 protein, as well as full length and C-terminally truncated G614 S-protein. We included the C-terminal truncation, as it has been shown previously to increase transduction with spike pseudotyped subviral particles, probably due to the removal of a retention sequence\textsuperscript{23}. Our pseudotyped lentivectors expressed both a secreted luciferase as well as GFP; therefore, measurements were performed by luminescence detection and flow cytometry.

As expected, exposure of wild type (mock-transduced) HeLa or A549 cells to the lentivectors described above did not lead to GFP or luciferase expression, which suggests that these cells were not permissive to transduction by our pseudotyped lentivectors (Fig. 2c–f). Similarly, overexpression of TMPRSS2 by itself did not lead to detectable transduction. Using the D614 pseudotyped lentivector, low-level transduction of ACE2 and ACE2/TMPRSS2-expressing HeLa cells were observed, but which was hardly detectable with A549 cells. This changed markedly when the G614 pseudotyped vector was used, with there being significant transduction of both ACE2-expressing cell lines. Once again, transduction of ACE2-expressing HeLa cells was significantly higher than for ACE2-expressing A549 cells. Interestingly, overexpression of TMPRSS2 in combination with ACE2 expression led to an additive enhancement of transduction in A549 cells, which was not the case for HeLa cells. We hypothesized that our HeLa cell line most likely had high-level expression of a yet to be defined S-protein cleaving protease, and that overexpression of TMPRSS2 is not required for efficient transduction. In contrast, in A549 cells, proteolytic cleavage appears to be rate-limiting, and overexpression of TMPRSS2 is required to enhance transduction.

D614G mutation alters the interactions around the point of the mutation. Several trajectories from long molecular dynamics (MD) simulations of the S-protein of the SARS-CoV-2 virus have been released to the public domain (CC BY 4.0)\textsuperscript{24}. These long simulations (10 μs) were obtained in specialized supercomputers\textsuperscript{25}. For the present analysis, we used two trajectories of the trimeric SARS-CoV-2 spike glycoprotein\textsuperscript{24}. The first simulation was initiated from the closed, ‘down’ conformation (PDB entry 6VXX), and the second one from a partially open, ‘up’, conformation (PDB entry 6VYB)\textsuperscript{25}. The simulated structures contain modelled missing loops from the crystal structures and glycans\textsuperscript{27}. Although no conformational transitions between the two states
Figure 1. Structural implications of the D614G mutation in the S-protein. (a) Top and side views of the S-protein in the closed and open conformations. D614 in the three protomers is depicted in orange van der Waals spheres. The RBD of protomer A in the down and up state is coloured in light green and dark pink, respectively. Graphical representation of the domain organisation of the S-protein: signal peptide (SP), receptor binding domain (RBD), N-terminal domain (NTD), subdomains 1 and 2 (SD1 and SD2), fusion peptide (FP), heptad repeat 1 (HR1), central helix (CH), connector domain (CD), heptad repeat 2 (HR2), transmembrane domain (TM). (b) Time series of D614 with K835 and P826 at the A–C interface over the course of the simulations initiated from the closed (green) and partially open (pink) conformations. The equivalent interactions of D614 at the B–C interface are given in grey. The probability distribution function (PDF) of the two interactions are displayed on the top of each timeseries. (c) Upper panels: interactions involving D614 for the closed (left) and open (right) conformation at the end of each simulation. The D614 residue of chain A is depicted in pink and green spheres and the interacting residues of chain C are represented in white spheres. Bottom-panels: interactions involving D614, when mutated to glycine, for the closed (left) and open (right) conformation at the end of each simulation. The Cα atom of G614 of chain A is depicted in pink and green spheres and the interacting residues of chain C are represented in white spheres.
Figure 2. SARS-CoV-2 S-protein D614G variant shows increased infectivity. (a,b) Real-time RT-PCR for ACE2, TMPRSS2, furin and cathepsin L mRNA expression profiles in HeLa (a) and A549 (b) cells. (c–f) Spike pseudotyped lentiviral infectivity assays. Pseudotyped lentiviruses expressing Luciferase and GFP and pseudotyped with SARS-CoV-2-D614 or G614 full length or truncated version were used to transduce HeLa and A549 overexpressing or not either ACE2 and/or TMPRSS2. (c,d) Luciferase assay of HeLa (c) and A549 (d) at 5 days post transduction with 80 μL of SARS-CoV-2 spike pseudotyped lentivirus. (e,f) Percent of GFP + assay of HeLa (e) and A549 (f) cells at 5 days post transduction with 80 μL of SARS-CoV-2 S-protein pseudotyped lentivirus analysed by Flow cytometry. The experiments were done in triplicates and repeated three times with three independent lentiviral stocks that was resulting from independent transfection. One representative is shown with error bars indicating SEM of replicates.
of the D614(A)-K835(C) salt bridge in the closed conformation, but very little will change in the open conformation. The D614G mutation will disrupt the formation (Fig. 1c). The D614G mutation stabilizes the open conformation of the S-protein as the pairwise RMSD in the wild type. Another molecular dynamics study published during the revision of the present manuscript form of the S-protein shows that the mutation favours the 'up' conformation, 82% population compared to 42% recent article points in the same direction where negative stain electron microscopy reconstructions of the D614G correlation matrix is very stable (Supplementary Figure 2) even for the RDB (A) domain exposed to the solvent. A

The interactions involving D614, once the structures are fully relaxed after MD simulations, clearly reveal a different behavior between the closed and open conformations. The D614G mutation will disrupt the formation of the D614(A)-K835(C) salt bridge in the closed conformation, but very little will change in the open conformation (Fig. 1c). The D614G mutation stabilizes the open conformation of the S-protein as the pairwise RMSD correlation matrix is very stable (Supplementary Figure 2) even for the RDB (A) domain exposed to the solvent. A recent article points in the same direction where negative stain electron microscopy reconstructions of the D614G form of the S-protein shows that the mutation favours the 'up' conformation, 82% population compared to 42% in the wild type. Another molecular dynamics study published during the revision of the present manuscript also concludes that the D614G mutation favors the open conformation of the RDB.

Two abundant amino acid changes in SARS-CoV-2: S protein D614G and RNA-dependent RNA polymerase (RdRp) P323L. Available high-quality sequences of SARS-CoV-2 (27,084) were analysed for the D614 and G614 variants of the S-protein and the P323 and L323 of the RdR polymerase. Data are shown as total number of sequences per week (a, note the logarithmic scale) or percent of sequences per week (b).

Two major protein variants of SARS-CoV-2: S protein D614G and RNA-dependent polymerase (RdRp) P323L. 27,084 genomic sequences from GISAID (www.gisaid.org) representing 87 countries over the time period of 24 December 2019 to 19 June 2020, were used to perform the epidemiological study. Using the original D614/P323 variant (NC_045512.2) as a reference, we identified two highly abundant SARS-CoV-2 mutations leading to amino acid changes: the D614G mutation in the S-protein, and the P323L mutation in the RNA-dependent RNA polymerase (RdRp). Figure 3a,b shows the weekly frequency of these mutations in the viral sequences over time (log scale), while Fig. 3c,d shows the percentage of the respective mutants within a given week. Until mid-January 2020, both mutations were virtually absent in the available sequences. Thereafter, we can observe a notable increase in the frequency of this mutant, with a virtually complete replacement of the original sequences by May 2020. A similar striking increase in the frequency of both the D614G and P323L mutations should be noted.

Co-emergence and concurrence of the D614G and the P323L mutations. Similar kinetics in the appearance of D614G and P323L led us to further analyse the relationship between the two mutations. Figure 4a shows the occurrence of the four possible variants: D614/P323 (i.e., the original D614/P323 variant), G614/P323 (mutation only in the S-protein), D614/L323 (mutation only in RdRp), and G614/L323 (mutations in both the S-protein and RdR). In our dataset, these variants were first reported as follows: D614/P323 on 2019-12-24; G614/P323 on 2020-01-24, D614/L323 on 2020-03-10, and G614/L323 on 2020-01-24. Interestingly, only the double mutation G614/L323 was epidemiologically successful and at the time of writing accounted for ~70% of all available SARS-CoV-2 sequences; while the original Wuhan sequence, accounted for ~30%. The single mutants accounted for only 0.3% (G614/P323) and 0.1% (D614/L323) of all sequences. To investigate the emergence and spread of the two strains, we analysed the weekly numbers of sequences (Fig. 4b) and the cumulative number of sequences (Fig. 4c) of D614/P323 and G614/L323 variants. There was an evident first wave from December 2019 to February 2020, which was strongly dominated by the D614/P323 variant. From March 2020,
a second wave with a predominance of the G614/L323 variant can be observed. We also observed major differences between the variants in different countries (Fig. 4d). For example, while D614/P323 was the largely dominant variant in Hong Kong, South Korea and China; G614/L323 was almost exclusively observed in countries such as Switzerland, Czechia and Argentina. We also note that the occurrence of single mutants (G614/P323 and D614/L323) continued to be rare.

To better understand the relationship between the dynamics and geographic aspects of the pandemic, we analysed the weekly and cumulative numbers of sequences (Fig. 5) in different countries (only countries with at least 20 sequences were included in this analysis). At first glance, the D614/P323 variant can be observed as the dominant strain in the initial wave in Asian countries, while the G614/L323 variant was the main driver of the pandemic in other continents. However, in several Asian countries, including Taiwan, Thailand, Japan; the emergence of G614/L323 can be observed at later time points. Also, in many non-Asian countries, the pandemic started with the D614/P323 variant, followed by a strong predominance of the G614/L323 variant (e.g. Italy, UK, USA). Thus, country-by-country data do not suggest a simple geographic predominance or a bottleneck in the transmission that led to a random emergence of G614/L323 in non-Asian countries. Rather, it appears that G614/L323 is epidemiologically more successful than the original D614/P323 variant.

We also investigated the case fatality rate in different countries as a function of the percentage of D614/P323 vs. G614/L323. We found no significant correlation between strain predominance and either deaths per million population (Fig. 6a) or case fatality rate (Fig. 6b). However, we cannot entirely exclude such a correlation, as there was a weak trend towards a higher case fatality rate in countries with a high percentage of the G614/L323 variant.

We next investigated the epidemiological parameters in more detail. In 66 countries, sequences of both the D614/P323 variant and the G614/L323 were reported. In 37 countries, the D614/P323 variant was reported before the G614/L323 (Fig. 7a, light blue bars); while in 24 countries, the double mutation was reported first (Fig. 7a, dark blue bars). In five countries, both variants were reported for the first time on the same date. Note that even in countries where the D614/P323 variant was reported more than 20 days before the G614/L323, ultimately a clear predominance of the G614/L323 variant was observed (e.g. Belgium, USA, France). We, therefore, decided
Figure 5. Temporal evolution of the four possible D614G and P323L variants in different countries. Available high-quality sequences of SARS-CoV-2 (27 075) were analysed for the different possible combinations of the D614 and G614 variants of the spike protein and the P323 and L323 of the RdR polymerase based on country of origin of the sequences. Only countries with at least 20 sequences were included in this analysis. Results are shown as cumulative number of sequences over time (the weekly number of sequences is provided as a Supplementary Figure 4).
to compare the spread of the two dominant strains and the single mutant variants in different countries with respect to the total number of sequences collected within 14 days following the emergence of each respective strain (only countries with at least five sequences at day 14 were included). As shown in Fig. 7b for individual countries and summarized in Fig. 7c, the emergence of the single mutant strain was slow and of low efficiency. The D614/P323 variant shows a relatively slow emergence, which increases mostly after seven days. In contrast, the G614/L323 strain emerged very rapidly and efficiently. When comparing the rate at which the strains emerge within the given countries, the difference between the D614/P323 variant and the G614/L323 strain was statistically significant ($p$ value $< 2.2e^{-16}$; Wilcoxon signed rank test with continuity correction).

P323L mutation of the SARS-CoV-2 RdRp polymerase. The SARS-CoV-2 polymerase RdRp has an overall architecture that resembles the one seen in SARS-CoV-1 and consists of a nsp12-nsp7-nsp8 complex. The structure of the SARS-CoV-2 nsp12 contains a cupped, right-handed RdRp domain linked to a nidovirus RdRp-associated nucleotidyl-transferase domain (NiRAN) via an interface domain. The RdRp domain adopts the conserved architecture of the viral polymerase family 4 and is composed of three domains: a fingers domain, a palm domain and a thumb domain (Fig. 8). The nsp12 is stabilized by a nsp7–nsp8 heterodimer, which is packed against the thumb-finger interface, and a single nsp8 monomer interacting with the finger and interface domains. The P323L mutation is located on the interface domain of nsp12, and specifically on the loop that connects the three helices of the interface domain to the three β-strands of the same domain.

Although P323 engages only in a single interdomain interaction with F396 of the finger domain (Fig. 8), L323 is expected to affect the local dynamics of the regions around the point of the mutation. The conformational freedom of the proline-rich $\text{321FPPTS325}$ stretch, where the P323L mutation lies, is expected to be limited due to the pyrrolidine ring of the two Pro residues, which provide a degree of local structural rigidity. Phe has been known to be over-represented in sequence positions immediately following Pro–Pro motifs in protein turns and NMR data have shown a strong Pro–Phe, $\text{C}_\text{H}^\text{π}$ interaction that restricts the rotation of the Phe side chain$^{31}$. Therefore, although L323 will still in principle be able to interact with F396, the mutation is expected to increase the flexibility of this region.

Figure 6. SARS-CoV-2 variants: mortality, case fatality rate, and patient status. Relationship between the percentage of the two main SARS-CoV-2 variants (D614/P323 and G614/L323) in different countries and (a) number of deaths per 100,000 population and (b) case fatality rate in the respective countries. Numbers of deaths per 100,000 population and case fatality rates were taken from the Johns Hopkins Coronavirus Resource Center (coronavirus.jhu.edu/data/mortality) at 4th of August 2020.
A mutagenesis study has confirmed the functional role of nsp7 and nsp8 in SARS-CoV-1 replication. Specifically, Subissi et al. identified three nsp8 residues (P116, P183, and R190) that are critical for the nsp8/nsp12 interaction in vitro and in vivo, and the reported impaired SARS-CoV-1 polymerase activity in mutant variants underlines the key role of these residues in the nsp8/nsp12 association and subsequent SARS-CoV-1 RNA synthesis. Although similar experiments have to be carried out for the case of SARS-CoV-2, R190 and P116 are very close in space to the point of the P323L mutation, and P116 interacts directly with S325 of the 321FPPTS325 stretch (Fig. 8). Therefore, we anticipate the P323L mutation to have an impact on the nsp8 association. The level of association of nsp8 has been shown to be an important factor for obtaining a high RNA polymerase activity in SARS-CoV-1, and therefore enhanced interaction between the nsp12 and nsp8 upon the P323L mutation may give an evolutionary advantage to the virus.

Discussion

Compared to DNA-based organisms, RNA viruses are generally believed to replicate their genetic material with lower accuracy, as they lack the necessary proofreading and correction mechanisms. Given the high mutation rate seen in viruses, the mutations that give an advantage to the virus and, therefore, are successful, are fine-tuned through evolutionary pressure. Here, we have analysed the structural and functional impact of the widely discussed S-protein mutation D614G and investigated whether it provides by itself an epidemiological advantage to SARS-CoV-2. We demonstrate that indeed, this single amino acid substitution from aspartate to glycine at position 614 significantly enhances the infectivity of pseudotyped lentivectors. Through molecular modelling, we suggest that the G614 amino acid substitution destabilizes the closed conformation, while it does not alter the stability of the open one. The transition towards the open state is thought to be required for viral fusion, and thus, a structurally favoured open state is in line with the higher infectivity demonstrated by our biochemical assays.

However, epidemiological data suggests that the D614G mutation is not exclusively responsible as a driver mutation of SARS-CoV-2. The time series for the emergence of the D614G and P323L mutations indicates that the S-protein mutation (D614G) evolved simultaneously with P323L in RdRp. We observed small clusters of the
SARS-CoV-2 in general, possibly by providing a selective advantage as hypothesized by Korber et al.34. Although more experiments are needed to understand the underlying biological relevance of the two mutations, we speculate that the P323L mutation attenuates the clinical presentation of the SARS-CoV-2. Firstly, based on its location, we expect the P323L mutation to negatively impact the nsp8 association. The level of association of nsp8 has been shown to be an important factor for obtaining high RNA polymerase activity in SARS-CoV-132, and thus P323L is likely to attenuate polymerase activity in SARS-CoV-2. Secondly, it becomes increasingly obvious that asymptomatic infections are a key factor in the epidemiological success of SARS-CoV-2. We, therefore, speculate that the P323L RdRp mutation might decrease the production of viral RNA and hence increase the likelihood of asymptomatic infections and/or the time between infection and appearance of symptoms. The infectivity-enhancing D614G S-protein mutation would then counterbalance this initial decrease in virion production and allow transmission even by asymptomatic or presymptomatic individual.

What do we know about the underlying molecular mechanisms? Regarding the S-protein, the analysis of publicly available simulations reveals an interesting interaction network around the D614 residue. The main interactions change between the two molecular dynamics datasets, observing a cooperative effect among protomers following the symmetry of the oligomer. We believe that the mutation of D614 will disrupt the interactions that couple the AC, CB and BA interfaces. In particular, none of the salt bridges related to D614 could be formed, shifting the conformational population to a more open-like situation (Fig. 1a,b). Therefore, the D614G mutation observed more prevalently in specific regions might help to reduce the energetic cost of the conformational transition needed to attach to the cell receptors. In the case of the RdRp, the P323L mutation is located far from the RNA binding domain, which lies within the finger-palm-thumb domains and, therefore, any direct effect of the mutation on the ability of the mutated RdRp to bind RNA should be excluded. However, the mutation could exert its effect allosterically. Since allosteric effects are impossible to be inferred by static crystal structures, simulation techniques can be employed to further elucidate the effect of the mutation on the dynamics of the interface domain and the NiNAR, fingers, and palm domains that are in direct contact with the domain where the mutation is found. Moreover, experiments on the stability of the P323L variant are needed, since changes in the stability of the mutated nsp12 could alter its cellular concentration, which would subsequently affect the replication rate of the virus within the host cell.

Figure 8. Structural implications of the D614G mutation in S-protein. (a) Domain organisation of SARS-CoV-2 nsp12 in complex with nsp7 and nsp8 cofactors. The P323 is depicted in beige spheres. (b) Close up around P323. The residues of the 321FPPTS325 stretch are depicted in sticks, while F396 that P323 interacts with is depicted in spheres. (c) Residues P116, P183 and R190 of the nsp8 that has been reported to be crucial for the nsp8/nsp12 interaction in SAR-CoV-1 are depicted in red spheres.
Taken together, our study demonstrates that the mutation D614G in S-protein has high infectivity in cellular systems and provides a structural explanation of this effect through molecular modelling. Our data, however, are not compatible with the concept that the D614G S-protein mutation by itself is sufficient to explain the epidemiological success of the presently predominant SARS-CoV-2 strain. The coevolution of the S-protein mutation with the P323L mutation of the RdRp is striking and appears to be crucial for the success of the viral strain. We provide a working hypothesis where the polymerase mutation attenuates the symptoms and severity of the infection, while the spike mutation improves the ability of the D614G spike variant to infect cells.

Methods

**Cell lines and culture.** Cells were maintained in Dulbecco modified Eagle medium (DMEM) supplemented with 10% (v/v) fetal bovine serum (FBS), 100 μg/ml of penicillin and streptomycin, 2 mM l-glutamine, 1 mM sodium pyruvate and 1% non-essential amino acids. Cultures were maintained at 37 °C in a 5% CO2 atmosphere. Human cell lines used in the study were HEK 293T/17 (human kidney, ATC C#CRL-11268), A549 cells and HeLa cells.

**Luciferase reporter assay.** Cells were transduced with SARS-CoV-2 S-protein lentiviral vectors encoding GFP and Gaussia luciferase (GluC). Four to five days later, Gluc activity was measured by adding 5 μL of the cell supernatant sample to 50 μL of room temperature assay buffer (100 mM NaCl, 35 mM EDTA, 0.1% Tween 20, 300 mM sodium ascorbate, 4 μM coelenterazine in 1 × phosphate-buffered saline [PBS]) and immediately measuring luminescence in a luminometer (Glomax; Promega).

**Generation of cell lines expressing ACE2 and TMPRSS2.** HeLa and A549 cells were plated at a density of 2E05 cells per well in a 12 well plate. The day after cells were co-transduced either with ACE2/puromycin and/or TMPRSS2/blasticidin lentiviruses. Four days after transduction, cells were selected with blasticidin and puromycin at 5 μg/mL. Cells were maintained as polyclonal population.

**Plasmids.** ACE2 and TMPRSS2 cDNA ORFs were purchased from GenScript and were cloned into pCDH-CMV-MCS-EF1α-puro and pCDH-CMV-MCS-EF1α-Blast lentivectors respectively using standard cloning methods. pCG1_ScOV-2 plasmid encoding SARS-CoV-2 S-protein was provided by Prof. Dr. Stefan Pöhlmann (Deutsches Primatenzentrum, Leibniz-Institute for Primate research, Göttingen, Germany). G614 S-protein was cloned using site-directed mutagenesis with the following primers: 5′-GCG TGA ACT GTA CCG AAG TG-3′ and 5′-CAA TCT AGA TTA GCA GCA GCT GCC ACAG-3′. The pGAG plasmid was kindly provided by Didier Trono (Addgene plasmid 12259). The medium was changed 8 h post-transfection. After 48 h, the viral supernatants were collected and filtered using 45 μm PVDF filters and stored at −80 °C.

**Lentivirus production.** For recombinant-lentivirus production, plasmids were transfected in 293 T cells using the calcium phosphate method. Briefly, 4.5 × 10⁶ cells were plated in a 10-cm dish and transfected 16 h later with 15 μg either ACE2, TMPRSS2, or empty-expressing lentiviral vectors, 10 μg of packaging plasmid [psPAX2, gift from Didier Trono (Addgene plasmid 12260)], and 5 μg of envelope [pMD2G, gift from Didier Trono (Addgene plasmid 12259)]. The medium was changed 8 h post-transfection. After 48 h, the viral supernatants were collected and filtered using 45 μm PVD filters and stored at −80 °C.

**SARS-CoV-2 S-protein pseudotyped lentiviral production.** Spike-pseudotyped lentivirus was produced by co-transfection 293 T cells with psPAX2, pCDH-CMV-GluC-EF1α-GFP, and plasmids encoding either SARS-CoV-2 S-protein full length (FL), S-protein DeltaCter (DL), S-protein DeltaCter (DL) and GFP, or empty vector by using the calcium phosphate method as described above.

**SARS-CoV-2 S-protein pseudotyped lentiviral infectivity assays.** HeLa and A549 cells were seeded into 96-well plates at a density of 1000 cells per well and inoculated with 50 μL media containing S-protein pseudovirions. After 2-day incubation, cells were washed with PBS and the media was refreshed. Two days later, supernatants were collected to measure Gaussia luciferase activity. The cells were trypsynized and resuspended in PBS and were analysed by flow cytometry using Attune NxT Flow Cytometer (Thermo Fisher Scientific). Data were analysed using FlowJo 11 (FlowJo, LLC, Ashland, OR).

**Real-time quantitative RT-PCR.** Total RNA was extracted from A549 and HeLa cell lines using RNasy Micro Kit (Qiagen). Complementary DNA was synthesized from total RNA using PrimeScript RT Reagent kit (Takara). The real-time PCR measurement of cDNAs was performed using PowerUp SYBR Green Master Mix (Applied Biosystems) and normalized to the expression of GAPDH as control the housekeeping gene. The primers were: Furin forward: ggaacatgacagctgcaact; Furin reverse: tcgtcacgatctgcttctca; Cathepsin L forward: gaccatgt; GAPDH forward: gcacaagaggaagagagacc; GAPDH reverse: aggggagattcagtgtggtg. The real-time PCR measurement of cDNAs was performed using PowerUp SYBR Green Master.

**Epidemiological data.** Metadata and 36,105 sequences specified as ‘complete’ and ‘high coverage’ were retrieved from GISAID (www.gisaid.org/) as of 30 June 2020. Only sequences that had metadata information were filtered further to contain human as host, a full date including month of the day that were collected before 22 June 2020, leaving a total of 34,841 sequences. The latter, together with the reference sequence (NC_045512.2),
were used to perform multiple sequence alignment (MSA) using MAFFT\(^4\) (v7.453) with default parameters. We retained only sequences where the first amino acid in S and RdRp protein were identical to the reference proteins, respectively. Next, we filtered out sequences that contained non-nucleotide bases e.g. ‘N’ and whose sequence length was different from the respective reference gene. We kept only sequences that corresponded to the criteria for both S protein and RdRp, leaving a total of 27,084 sequences. Among these, there were two variants for both S-protein and RdRp protein representing the vast majority of sequences (Supplementary Figure 3). Due to the very low number of other sequence variants (N614 in the S-protein and F323, H323, S323 in the RdRp protein), we removed sequences containing other variants, leaving us with 27,075 sequenced genomes for further analysis.

Unless stated otherwise, only countries that had sequenced a minimum of 20 genomes were used in the analysis. However, for Worldwide analysis, all countries were included. All plots were done using ggplot2 in R. The correlation plots were done using ggscatter function from the ‘ggpubr’ library. The statistical interference between the two dominant variants was performed using pairwise ‘wilcox.test’ function.

**S-protein modelling.** The simulations of S-protein used the Amber f99SB-ILDN force field for proteins\(^5\), the CHARMM TIP3P model\(^6\) for water, and the generalised Amber force field\(^7\) for glycosylated asparagine. The system was neutralized and salted with 0.15 M concentration of NaCl. The interval between frames of the analysed trajectories is 1.2 ns, and the simulations were run at 310 K in the NPT ensemble.

**References**

1. Petersen, E. et al. Comparing SARS-CoV-2 with SARS-CoV and influenza pandemics. *Lancet Infect. Dis.* https://doi.org/10.1016/1473-3099(20)30484-9 (2020).
2. Smith, E. C., Blanc, H., Sureda, M. C., Vignuzzi, M. & Denison, M. R. Coronavirus lacking exoribonuclease activity are susceptible to lethal mutagenesis: evidence for proofreading and potential therapeutics. *PLoS Pathog.* 9, e1003565. https://doi.org/10.1371/journal.ppat.1003565 (2013).
3. Ferron, F. et al. Structural and molecular basis of mismatch correction and ribavirin excision from coronavirus RNA. *Proc. Natl. Acad. Sci. USA* 115, E162–E171. https://doi.org/10.1073/pnas.1718806115 (2018).
4. Denison, M. R., Graham, R. L., Donaldson, E. F., Eckerle, L. D. & Baric, R. S. Coronavirus: an RNA proofreading machine regulates replication fidelity and diversity. *RNA Biol.* 8, 270–279. https://doi.org/10.4161/rna.8.2.15013 (2011).
5. Bar-On, Y. M., Flammholz, A., Phillips, R. & Milo, R. SARS-CoV-2 (COVID-19) by the numbers. *eLife* https://doi.org/10.7554/elife.57309 (2020).
6. SanjuÁEAc, N., Nebot, M. R., Chirico, N., Mansky, L. M. & Belshaw, R. Viral mutation rates. *J. Virol.* 84, 9733–9748. https://doi.org/10.1128/jvi.00694-10 (2010).
7. Pathan, R. K., Biswas, M. & Khandaker, M. U. Time series prediction of COVID-19 by mutation rate analysis using recurrent neural network-based LSTM model. *Chaos Solitons Fractals* 138, 110018. https://doi.org/10.1016/j.chaos.2020.110018 (2020).
8. Alkhansa, A., Lakkis, G. & El Zein, I. Mutational analysis of SARS-CoV-2 ORF8 during six months of COVID-19 pandemic. *Gene Rep.* 23, 101024. https://doi.org/10.1016/j.genrep.2021.101024 (2021).
9. Yan, R. et al. Structural basis for the recognition of SARS-CoV-2 by full-length human ACE2. *Science (New York, N.Y.)* 367, 1444–1448. https://doi.org/10.1126/science.abb2762 (2020).
10. Wrapp, D. et al. Cryo-EM structure of the 2019-nCoV spike in the prefusion conformation. *Science (New York, N.Y.)* 367, 1260–1263. https://doi.org/10.1126/science.abb2507 (2020).
11. Jaimies, J. A., Millot, J. K. & Whitaker, G. R. Proteolytic cleavage of the SARS-CoV-2 spike protein and the role of the novel S1/S2 site. *Sci. Adv.* 23, 101212. https://doi.org/10.1126/sciadv.101212 (2020).
12. Korber, B. et al. Spike mutation pipeline reveals the emergence of a more transmissible form of SARS-CoV-2. *bioRxiv*, 2020.04.20.069054. https://doi.org/10.1101/2020.04.29.069054 (2020).
13. Becerra-Flores, M. & Cardozo, T. SARS-CoV-2 viral spike G614 mutation exhibits higher case fatality rate. *Int. J. Clin. Pract.* https://doi.org/10.1111/ijcp.13525 (2020).
14. Li, Q. et al. The impact of mutations in SARS-CoV-2 spike on viral infectivity and antigenicity. *Cell*. https://doi.org/10.1016/j.cell.2020.07.012 (2020).
15. Ogawa, J. et al. The D614G mutation in the SARS-CoV-2 Spike protein increases infectivity in an ACE2 receptor dependent manner. *bioRxiv*. https://doi.org/10.1101/2020.07.21.214932 (2020).
16. Daniloiski, Z. et al. The Spike D614G mutation increases SARS-CoV-2 infection of multiple human cell types. *eLife*. https://doi.org/10.7554/eLife.65365 (2021).
17. Bhattacharyya, C. et al. Global Spread of SARS-CoV-2 Subtype with Spike Protein Mutation D614G is Shaped by Human Genomic Variations that Regulate Expression of TMPRSS2 and MX1 Genes (Cold Spring Harbor Laboratory, 2020).
18. Goodfellow, Goodfellow, I. & Taube, S. I. Calicivirus replication and reverse genetics. In *Viral Gastroenteritis. Molecular Epidemiology and Pathogenesis*, 355–378 (2016).
19. Yin, W. et al. Structural basis for inhibition of the RNA-dependent RNA polymerase from SARS-CoV-2 by remdesivir. *Science (New York, N.Y.)* https://doi.org/10.1126/science.abf1560 (2020).
20. Gao, Y. et al. Structure of the RNA-dependent RNA polymerase from COVID-19 virus. *Science (New York, N.Y.)* 368, 779–782. https://doi.org/10.1126/science.abb7498 (2020).
21. Pachetti, M. et al. Emerging SARS-CoV-2 mutation hot spots include a novel RNA-dependent RNA polymerase variant. *J. Transl. Med.* 18, 179. https://doi.org/10.1186/s12967-020-02344-6 (2020).
22. Hoffmann, M. et al. SARS-CoV-2 cell entry depends on ACE2 and TMPRSS2 and is blocked by a clinically proven protonate inhibitor. *Cell* 181, 271–280.e278. https://doi.org/10.1016/j.cell.2020.02.052 (2020).
23. Fukushima, S. et al. Vesicular stomatitis virus pseudotyped with severe acute respiratory syndrome coronavirus spike protein. *J. Gen. Virol.* 86, 2269–2274. https://doi.org/10.1099/vir.0.09055-0 (2005).
24. Research, D. E. S. Molecular Dynamics Simulations Related to SARS-CoV-2. *D. E. Shaw Research Technical Data* (http://www.deshawresearch.com/resources_sarscovid2.html) (2020).

Received: 14 October 2020; Accepted: 25 May 2021
Published online: 01 July 2021
were involved in the study. SARS-CoV-2 and to the many participants who kindly provide the data. None of the participants in GISAIDous allocation of supercomputer time. We also are grateful to GISAID and its team for creating a resource for institutional affiliations.

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