Consistent and High-Frequency Identification of an Intra-Sample Genetic Variant of SARS-CoV-2 with Elevated Fusogenic Properties

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Abstract

The severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has a genome comprised of a ~30K nucleotides non-segmented, positive single-stranded RNA. Although its RNA-dependent RNA polymerase exhibits exonuclease proofreading activity, viral sequence diversity can be induced by replication errors and host factors. These variations can be observed in the population of viral sequences isolated from infected host cells and are not necessarily reflected in the genome of transmitted founder viruses. We profiled intra-sample genetic diversity of SARS-CoV-2 variants using 15,289 high-throughput sequencing datasets from infected individuals and infected cell lines. Most of the genetic variations observed, including C->U and G->U, were consistent with errors due to heat-induced DNA damage during sample processing, and/or sequencing protocols. Despite high mutational background, we confidently identified intra-variable positions recurrent in the samples analyzed, including several positions at the end of the gene encoding the viral S protein. Notably, most of the samples possesses a C->A missense mutation resulting in the S protein lacking the last 20 amino acids (SΔ20). Here we demonstrate that SΔ20 exhibits increased cell-to-cell fusion and syncytia formations. Our findings are suggestive of the consistent emergence of high-frequency viral quasispecies that are not horizontally transmitted but involved in intra-host infection and spread.

Author summary

The severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) and its associated disease, COVID-19, has caused significant worldwide mortality and unprecedented economic burden. Here we studied the intra-host genetic diversity of SARS-CoV-2 genomes and identified a high-
frequency and recurrent non-sense mutation yielding a truncated form of the viral spike protein, in both human COVID-19 samples and in cell culture experiments. Through the use of a functional assay, we observed that this truncated spike protein displays an elevated fusogenic potential and forms syncytia. Given the high frequency at which this mutation independently arises across various samples, it can be hypothesized that this deletion mutation provides a selective advantage to viral replication and may also have a role in pathogenesis in humans.
Introduction

Observed for the first time in 2019, the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) and its associated disease, COVID-19, has caused significant worldwide mortality and unprecedented economic burden. SARS-CoV-2 is an enveloped virus with a genome comprised of a ~30K nucleotides non-segmented, positive-sense single-stranded RNA (vRNA) [1,2]. The virus is composed of four main structural proteins, encoded at the last 3’ terminal third of the viral genome: the spike glycoprotein (S), membrane (M), envelope (E) and the nucleocapsid (N) [3–5]. Attachment to the host receptor angiotensin-converting enzyme 2 (ACE2) is mediated by the S protein expressed on the surface of the virion [6]. Following its association, the S protein is cleaved into two separate polypeptides (S1 and S2), which triggers the fusion of the viral particle with the cellular membrane [7,8]. Once inside a cell, its RNA-dependent RNA polymerase (RdRp), which is encoded in the first open reading frame of the viral genome [9], carries out transcription and replication of the vRNA genome. In addition, mRNAs coding for the structural proteins (e.g., S, M, E and N) are expressed by subgenomic RNAs [9]. Once translated, the S, M and E proteins localize and accumulate in the endoplasmic reticulum–Golgi intermediate compartment (ERGIC) [10]. The S protein possesses an endoplasmic reticulum retrieval signal (ERRS) at its carboxy terminus, which is required for this localization [11]. At this location, the N protein associates with the viral genome and assembles into virions, which are transported along the endosomal network and released by exocytosis [9]. It was also observed for several coronaviruses that the S protein can localize to the cell surface and mediate cell fusion between adjacent cells, producing multinucleated cells or syncytia [8,12,13].

Genomic sequencing of SARS-CoV-2 vRNA from infected populations has demonstrated genetic heterogeneity [14–20]. Several recurrent mutations have been identified in consensus
sequences, and the geographical distribution of clades was established. Because they induce an abundance of missense rather than synonymous or non-sense mutations, it was suggested that regions of the SARS-CoV-2 genome were actively evolving and might contribute to pandemic spreading [20]. It was observed that variations are mainly comprised of transition mutations (purine->purine or pyrimidine->pyrimidine) with a prevalence of C->U transitions and might occur within a sequence context reminiscent of APOBEC-mediated deamination (i.e., [AU]C[AU]; [21,22]). Consequently, it was proposed that host editing enzymes might be involved in coronavirus genome editing [23,24].

Consensus mutations are only part of the genetic landscape in regard to RNA viruses. Replication of RNA viruses typically produces quasispecies in which the viral RNA genomes do not exist as single sequence entity but as a population of genetic variants [25]. These mutations are most frequently caused by the error-prone nature of each of their respective viral RdRps and by host RNA editing enzymes, such as APOBECs and ADARs [26]. However, the RdRp complex of large RNA viruses, such as coronaviruses, sometimes possess exonuclease proofreading activity, and consequently have lower error rates [25,27]. Quasispecies may sometimes exhibit diminished replicative fitness or deleterious mutations and exert different roles that are not directly linked to viral genomic propagation [28]. Mutations that form the intra-host genetic spectrum have been shown to help viruses evade cytotoxic T cell recognition and neutralizing antibodies and also render viruses more resistant to antiviral drugs [28]. These mutations can also be involved in modulating the virulence and transmissibility of the quasispecies [28].

In this study, we focussed on assessing intra-genetic variations of SARS-CoV-2. We analyzed high-throughput sequencing datasets to profile the sequence diversity of SARS-CoV-2 variants within distinct sample populations. We observed high genetic intra-variability of the viral
genome. By comparing variation profiles between samples from different donors and cell lines, we identified highly conserved subspecies that independently and recurrently arose in different datasets and, therefore, in different individuals. We further analyzed the dominant variant SΔ20 in a functional assay and demonstrate that this truncated spike protein enhances syncytium formation. Here we provide evidence for the existence of a consistently emerging variant identified across geographical regions that may influence intra-host SARS-CoV-2 infectivity and pathogenicity.

**Results**

**High intra-genetic variability of the SARS-CoV-2 genome in infected individuals.**

To assess the extent of SARS-CoV-2 sequence intra-genetic variability, we analyzed 15,224 publicly available high-throughput sequencing datasets from infected individuals. The raw sequencing reads were mapped to the SARS-CoV-2 isolate Wuhan-Hu-1 reference genome, and the composition of each nucleotide at each position on the viral genome was generated. Consensus sequences were produced for each dataset and the nucleotide composition for each position were compared to respective consensuses. To reduce the number of variations due to amplification bias and sequencing errors, duplicated reads were combined, and only positions mapped with a sequencing depth of 50 reads and having at least 5 reads with variations compared to the sample consensus were considered. Overall, we identified 301,742 variations from 11,362 samples located on 26,113 positions of the 29,903 nt SARS-CoV-2 genome. We observed an average of 26.6+/−132.0 variable nucleotides per sample (ranging from 1 to 5295 variations/sample; Fig 1A).
Analysis of the type of intra-genetic variations present in SARS-CoV-2 samples from infected individuals.

The analysis of the type of nucleotide changes within samples revealed that 52.2% were transitions (either purine->purine or pyrimidine->pyrimidine) and 47.8% were transversions (purine->pyrimidine or pyrimidine->purine). Notably, the highest nucleotide variations corresponded to C->U transitions (43.5%) followed by G->U (28.1%) transversion (Fig 1B), both types encompassing 71.6% of all variations. Since editing by host enzymes depends on the sequence context, we extracted two nucleotides upstream and downstream from each genomic position corresponding to variations and generated sequence logos. Our results indicated a high number of As and Us around all variation types and sites (62.1+/-3.4%; Fig 1B). Because SARS-CoV-2 is composed of 62% A/U, this suggests the observed number of As and Us around variation sites are mainly due to the A/U content of the viral genome, that no motifs are enriched around these sites and that these intra-genetic variations are likely not originating from host editing enzymes.

Identification of recurrent genetic variants of SARS-CoV-2 in samples from infected individuals.

To identify biologically relevant intra-genetic variations, we examined the variable positions that are recurrent in the samples analyzed. The variable positions were tabulated for each sample and then recurrent intra-genetic variations were calculated as percentages of samples containing variation at each position. Most variations are distributed homogeneously on the viral genome and most are poorly shared amongst samples (Fig 1C and 1D). However, our analysis
reveals 15 recurrent intra-variations shared by at least 5% of the samples analyzed (Fig 1C, above blue line; Table 1). Amongst these, four transversions (at nt 25,324, 25,334, 25,336 and 25,337) located at the 3’ end of the S gene are the most recurrent variations (inset of Fig 1C and Table 1). Three of these transversions (at nt 25,334, 25,336 and 25,337) correspond to missense mutations: E1258D (46.4%), E1258Q (27.6%) and D1259H (20.1%). Interestingly the most observed variation (at nt 25,324) is shared by 58.7% of the samples (6,668 of the 11,362 samples) and corresponds to a C->A transversion producing a nonsense mutation at amino acid 1,254 of the S protein (Fig 1C and 1D, red line; Fig 2B, red rectangle). The resulting S protein lacks the last 20 amino acids (SΔ20), which includes the ERRS motif at its carboxy terminus (Fig 2B, white letters on a black background). Amongst the sample with this intra-genetic variation, this C->A transversion represents from 2.9% to 42.4% of the subspecies identified (mean or 8.2+/-2.9%; Fig 2C and Table 1).

| Position (nt) | Proportion of Samples (%) | Type of Variation | Gene | Amino acid | Consensus codon | Variant codon | Consensus amino acid | Variant amino acid | Context (-2 to +2) | Frequency distribution (% of population) |
|--------------|---------------------------|------------------|------|------------|----------------|--------------|-------------------|-------------------|-----------------|---------------------------------------|
| 25324        | 58.69                     | C->A             | S    | 1254       | UGC            | UGA          | CYS (C)            | STOP             | UGCAA           | 8.19   2.89   2.86   42.37                                    |
| 25336        | 46.37                     | A->C             | S    | 1258       | GAA            | GAC          | GLU (E)            | ASP (D)          | GAAGA           | 6.38   2.10   2.42   29.09                                    |
| 25334        | 27.57                     | G->C             | S    | 1258       | GAA            | CAA          | GLU (E)            | HIS (H)          | AGAAC           | 4.79   1.63   2.03   22.81                                    |
| 25337        | 20.11                     | G->C             | S    | 1259       | GAC            | CAC          | ASP (D)            | HIS (H)          | AACAC           | 4.68   2.12   2.07   28.57                                    |
| 29187        | 10.95                     | C->U             | N    | 305        | GCA            | GLA          | ALA (A)            | VAL (V)          | UGAC            | 3.35   2.53   1.81   46.91                                     |
| 29188        | 10.68                     | A->G             | N    | 305        | GCA            | GCG          | ALA (A)            | VAL (V)          | ACAC            | 3.32   2.56   1.79   46.91                                     |
| 18591        | 10.21                     | C->G             | ORF1ab| 6108       | GUC            | GUG          | VAL (V)            | VAL (V)          | UGGUU           | 3.78   0.85   2.54   7.96                                    |
| 11874        | 10.02                     | U->C             | ORF1ab| 3870       | GUA            | GCA          | VAL (V)            | ALA (A)          | AEGAG           | 4.21   2.27   2.68   38.55                                    |
| 15965        | 8.12                      | G->U             | ORF1ab| 5233       | UGU            | ULU          | CYS (C)            | PHE (F)          | CUGUU           | 3.01   2.48   1.88   44.19                                    |
Analysis of intra-genetic variations present in SARS-CoV-2 samples from infected cells.

To further investigate variations in a more controlled system, and to determine whether host proteins are involved in SARS-CoV-2 genome editing, we used 65 high-throughput sequencing datasets generated in a recent transcription profiling study of several cell lines infected with SARS-CoV-2 [29]. Firstly, we mapped raw sequencing reads to the human genome to assess host modifying enzyme expression. For all cell lines, normalized counts for mRNAs corresponding to most host modifying enzymes were very low or non-detected (Fig 3), suggesting that these cell lines poorly expressed these host editing proteins. As above, the raw sequencing reads from infected cells were mapped to the SARS-CoV-2 genome sequence, the composition of each nucleotide at each position on the viral genome were generated, and nucleotide variations when compared to respective consensus sequences were calculated. Because the sequencing depths of the samples were low, we considered positions mapped by at least 20 reads and having at least 2 reads with variation compared to the sample consensus. In the samples derived from infected cells, we observed 29.7% and 70.3% of transitions and transversions, respectively. Similar to observations in samples from infected individuals, the highest nucleotide variations corresponded to G->U transversions (26.1%) and C->U (21.6%) transitions (Fig 4B). We then analyzed nucleotide compositions two nucleotides upstream and downstream of the intra-genetic variations.
As above, a high number of A/U (57.8+/-%7.7%) were present around sites showing variation (Fig 4B), consistent with the 62% A/U composition of the SARS-CoV-2 genome, indicating no enrichment of sequence motifs around these sites, except for the expected high number of As and Us.

We then examined the intra-genetic variable positions that are recurrent amongst the cell lines analyzed. We identified 29 positions within the viral populations showing intra-genetic variation enrichment in at least 10% of the cell cultures and most of them are located on structural genes, which are encoded at the last 3’ terminal third of the viral genome (Fig 4C and 4D). Similar to our observation from the samples from infected individuals, a cluster of recurrent variations is located at the 3’end of the S gene, including the C->A transversion at position 25,324 shared in 58.9% of the cell lines analyzed (Fig 4C and 4D, red line; Table 2). Overall, our results indicate consistent results between intra-genetic variations observed in infected cell lines and in samples from infected individuals, including the presence the viral subspecies resulting in a S protein truncated of its last 20 amino acids (SΔ20).

Table 2: Recurrent SARS-CoV-2 genome intra-variations shared by at least 10% infected cell cultures. Frequency distributions were calculated using data similar to Fig 2C. The variations are sorted by their recurrence, with the most shared variation at the top of the table.

| Position (nt) | Proportion of Samples (%) | Type of Variation | Gene | Amino acid | Consensus codon | Variant codon | Consensus amino acid | Variant amino acid | Context (-2 to +2) | Frequency distribution (% of population) |
|---------------|---------------------------|-------------------|------|------------|----------------|--------------|---------------------|------------------|------------------|----------------------------------------|
| 28853         | 82.14                    | U>A               | N    | 194        | UCA           | ACA          | SER (S)            | THR (T)           | GU>HC           | 8.16 7.99 1.52 35.42                      |
| 25336         | 58.93                    | A>C               | S    | 1258       | GAA           | GAC          | GLU (E)            | ASP (D)           | GA>GGA          | 21.77 5.81 12.00 42.22                     |
| 25324         | 58.93                    | C>A               | S    | 1254       | UGC           | UGA          | CYS (C)            | STOP             | UG>CAA          | 25.21 7.37 12.00 42.37                     |
| 23525         | 58.93                    | C>U               | S    | 655        | CAU           | UAU          | HIS (H)            | TYR (Y)           | AA>CAU          | 8.36 3.35 3.49 16.67                      |
| 25337         | 58.93                    | G>C               | S    | 1259       | GAC           | CAC          | ASP (D)            | HIS              | AA>GAA          | 20.43 4.65 12.86 35.56                     |
### SARS-CoV-2 Δ20 increases cell-to-cell fusion and the size of syncytia.

For several coronaviruses, the S protein mediates syncytia formation [8,12,13]. To investigate the effect of Δ20 on cell-to-cell fusion, HEK293T cells stably expressing human
ACE2 were co-transfected with a plasmid expressing GFP and plasmids expressing or not wild-type S or SΔ20 under a cytomegalovirus (CMV) major immediate early promoter. As expected, in the absence of the S protein (i.e., pCAGGS alone) syncytia formation was not observed (Fig 5A). Consistent with previous findings [8], we observed the presence of large cytoplasmic masses containing GFP in the presence of the wild-type S protein, indicating induction of cell-to-cell fusion (Fig 5A and 5B). Our results not only indicate that SΔ20 also induces fusion, but that the cytoplasmic masses are larger than the wild-type S protein (Fig 5A and 5B). To complement this approach, we quantified cell fusion using a bi-molecular fluorescence complementation (BiFc) assay composed of target cells containing GCN4 leucine zipper-Venus1 (ZipV1) with or without myc-ACE2, and effector cells containing GCN4 leucine zipper-Venus2 (ZipV2) with or without S/ SΔ20 (Fig 5C). Two hours after incubating the two cell populations, we observed an increase in fluorescence signal in the presence of wild-type S and ACE2, indicating fluorescence complementation and thus cell-to-cell fusion (Fig 5D). Consistent with our microscopy observations, fluorescence complementation was increased in the presence of SΔ20. Taken together, our results indicate that SΔ20 displays increased syncytia formation in HEK293T cells expressing ACE2 as compared to the wild-type S protein.

**Discussion**

Previous analyses of SARS-CoV-2 nucleotide variations indicated the prevalence of C->U transitions suggesting that the viral genome was actively evolving and those host editing enzymes, such as APOBECs and ADARs, might be involved in this process [23,24]. Although instructive on the role of host involvement in SARS-CoV-2 genome evolution, these studies were performed
on consensus sequences (*i.e.*, one per sample) and explore only part of the genetic landscape of this RNA virus. Here, we used a large number of high-throughput sequencing datasets to profile the intra-sample sequence diversity of SARS-CoV-2 variants, both in infected individuals and infected cell lines. We observed extensive genetic variability of the viral genome, including a high number of transversions, and identified several positions with recurrent intra-variability in the samples analyzed. Notably, most of the samples possessed a C->A missense mutation resulting in the S protein lacking the last 20 amino acids (SA20) that increases cell-to-cell fusion and syncytia formations.

Most intra-sample variations are distributed homogeneously across the viral genome, are not conserved or recurrent amongst samples, and a large number of them are C->U or G->U mutations. Previous analyses of SARS-CoV-2 sequence variations proposed that host editing enzymes might be involved in coronavirus transition editing based on results showing that C->U transitions occur within a sequence context reminiscent of APOBEC1-mediated deamination (*i.e.*, [AU]C[AU]) [21–24]. Here, we investigated nucleotide compositions at each variation site and observed a high number of As and Us around all variation types and sites. However, since the SARS-CoV-2 genome is 62% A/U-rich and similar percentages of As and Us were observed around all variations, we concluded that no motifs are enriched around these variations in the viral subspecies analyzed. Consequently, our results cannot support that host editing enzymes are a major source of these intra-sample variations.

Although it is possible that host RNA-editing enzyme are responsible for the occurrence of some variations, C->U transitions and G->U transversions are also generally associated with nucleotide deamination and oxidation, respectively [30–37]. It is common practice to thermally inactivate SARS-CoV-2 samples before performing RNA extractions followed by RT-PCR and
sequencing [38]. Heating samples can form free radicals, such as 8-hydroxy-20-deoxyguanine (8-
Oxo-dG), that could cause high levels of C->A and G ->U mutations and promote the hydrolytic
deamination of C->U [30–33,35,37,39,40]. It was previously reported that these types of mutations
occur at low frequency, that they are mostly detected when sequencing is performed on only one
DNA strand and that they are highly variable across independent experiments [32,34].
Consequently, most transversions observed in our analysis are likely due to heat-induced damage,
RNA extraction, storage, shearing and/or RT-PCR amplification errors. However, we identified
several positions with intra-sample variability recurrent in several independent samples, both from
infected individuals and infected cells. They were detected at moderate to high frequencies,
ranging from 2.5% to 39.3% per sample (Table 1 and 2), and most were derived from pair-end
sequencing (90.7%) in which the two strands of a DNA duplex were considered. Thus, it is likely
that these variations are genuine and represent hot spots for SARS-CoV-2 genome intra-sample
variability.

Amongst the variable positions identified in infected cells, most of them are located at the
last 3’ terminal third of the viral genome. These cells were infected with a high number of viruses
(i.e., high multiplicity of infection; MOI) for 24h [29]. The presence of several variations at
positions in the region coding for the main structural proteins likely reflects that this is a region
with increased transcriptional activity due to the requirement of producing their encoded mRNAs
from sub-genomic negative-sense RNAs [9].

Interestingly, a cluster of variations located at the 3’end of the S gene was observed for the
two datasets analyzed. They correspond to four transversions located at the 3’end of the S gene
and are shared by a large proportion of the samples. Three of these correspond to missense
mutations changing the charged side chains of two amino acids (E1258D, E1258Q and D1259H).
Notably, most of the samples possess a variability at position 25,324 producing a nonsense mutation at amino acid 1,254 of the S protein. The resulting protein lacks the last 20 amino acids (SΔ20) and thus does not include the ERRS motif at its carboxy terminus. For SARS-CoV-1, the ERRS domain localizes the S protein to the ERGIC and facilitates its incorporation into virions [11]. Deletion of this motif might cause the S protein of SARS-CoV-2 to accumulate to the plasma membrane and increase the formation of large multinucleated cells known as syncytia. Consistent with these observations, our results indicate an increase in syncytia formation frequency and size with SΔ20 as compared to the complete S protein. Although the biological function of SΔ20 is unknown, it is possible that the formation of syncytia facilitates the spread of the virus directly to neighboring cells. SΔ20 might also increase virus replication, as similar mutants (SΔ18, SΔ19 and SΔ21) were recently reported to increase both infectivity and replication of vesicular stomatitis virus (VSV) and human immunodeficiency virus (HIV) pseudotyped with SARS-CoV-2 S protein in cultured cells [41–44]. It is also tempting to suggest a link between SARS-CoV-2 pathogenesis and the presence of SΔ20, since severe cases of the disease were recently linked to considerable lung damage and the occurrence of syncytia [45,46].

Our findings indicate the presence of consistent intra-sample genetic variants of SARS-CoV-2, including a recurrent sub-population of SΔ20 variants with elevated fusogenic properties. Further investigation is required to better define the extent of SARS-CoV-2 variability in infected hosts and to assess the role of these subspecies in the life cycle of this virus. More importantly, further studies on the presence of SΔ20 and its link with viral pathogenicity could lead to better diagnostic strategies and designer treatments for COVID-19.
Methods

Analysis of intra-variability within SARS-CoV-2 samples.

15,289 publicly available high-throughput sequencing datasets were downloaded from the NCBI Sequence Read Archive (up to July 10, 2020). They comprise of 15,224 and 65 datasets from infected individuals and infected cell lines, respectively. The datasets from infected cells were generated by Blanco-Melo et al. [29]. Duplicated reads were combined to reduce amplification bias and mapped to the SARS-CoV-2 isolate Wuhan-Hu-1 reference genome (NC_045512v2) using hisat2 (v.2.1.0)[47]. For each dataset, the consensus sequences and the frequency of nucleotides at each position were extracted from files generated by bcftools (v.1.10.2) of the samtools package (v.1.1) with an in-house Perl script [48,49]. All further calculations were performed in R. To reduce the number of variations due to sequencing errors and/or protocol differences, only positions mapped with a sequencing depth of 50 reads and having at least 5 reads with variations compared to the sample consensus were considered. Sequence logos were generated with the ggseqlogo package (v.0.1) [50].

Differential expression analysis of transcript coding for APOBECs and ADARs.

High-throughput sequencing datasets generated in a recent transcription profiling study of several cell lines infected with SARS-CoV-2 were downloaded from SRA [29]. Duplicated reads were combined and mapped to the human reference genome (Homo_sapiens.GRCh38.83) using hisat2 (v.2.1.0)[47]. Transcript abundance was performed using HTSeq 0.12.4 [51] and normalized into Transcripts Per Million (TPM) in R.
Cell culture and plasmids.

Human embryonic kidney 293T (HEK293T) were obtained from the American Type Culture Collection (ATCC CRL-11268) and maintained in Dulbecco’s Modified Eagle’s Medium (DMEM) supplemented with 5% fetal bovine serum (Fisher Scientific), 5% bovine calf serum (Fisher Scientific) and 1x antibiotic-antimycotic (Fisher Scientific). HEK293T stably expressing human ACE2 (kind gifts of Dr. Hyeryun Choe, The Scripps Research Institute) were cultured and maintained in DMEM (Wisent) supplemented with 10% fetal bovine serum (Sigma), 1 U/mL penicillin, 1 µg/mL streptomycin, and 3 µg/mL glutamine (Corning). All cells were cultured at 37 °C in a humidified atmosphere containing 5% CO2. The original bi-molecular fluorescence complementation (BiFc) constructs GCN4 leucine zipper-Venus1 (ZipV1) and GCN4 leucine zipper-Venus2 (ZipV2) were sourced from Stephen W. Michnick (reviewed in [52]). pCEP4(myc-ACE2) was a gift from Erik Procko (Addgene plasmid #141185). pCAGGS expressing the SARS-CoV-2 S protein (pCAGGS(S)) was provided by Dr. Florian Krammer (Mount Sinai). pCAGGS(SΔ20) was generated using overlapping PCR to introduce a termination codon at residue 1254.

Syncytium Formation Assay.

HEK293T expressing ACE2 cells were seeded in 24-well plates in complete media to obtain an 85% confluence the following day. Cells were then placed in serum-free DMEM and transiently co-transfected using JetPRIME (Polyplus Transfection, France) with plasmids encoding GFP (MLV-GFP, a kind gift of Dr. James Cunningham, Brigham and Women’s Hospital), pCAGGS(S) or pCAGGS(SΔ20), and empty pCAGGS at a 0.15:0.2:0.65 ratio. 18 hours
post transfection, cells were imaged (ZOE Fluorescent Cell Imager, Bio-Rad) for syncytia formation using the Green channel to visualize fusion of GFP positive cells as performed previously [53]. GFP area was quantified on ImageJ [54].

Quantification of cell-to-cell fusion using bi-molecular fluorescence complementation (BiFc).

HEK293T cells were seeded in a 12-well microplate (500,000 cells/well) in complete media for 24h. Transient transfections were performed using JetPRIME (Polyplus transfection, France) according to the manufacturer's instructions. Target cells were transfected with ZipV1 (0.5 μg) or pCEP4(myc-ACE2) (0.05 μg) and ZipV1 (0.5 μg). Effector cells population were transfected with ZipV2 (0.5 μg) and pCAGGS(S) (0.125 μg) or pCAGGS(SΔ20) (0.125 μg). Total DNA was normalized using the empty pCAGGS vector DNA to 1 μg. Following transfection, cells were incubated at 37 °C for 24 h. Then, cells were rinsed with PBS and detached with versene (PBS, 0.53 mM EDTA) and counted. 35,000 cells/well of both populations were co-seeded in DMEM without serum in a 384-well black plate with optical clear bottom and incubated for 2 hours at 37 °C, 5% CO₂. BiFC signal was acquired using Biotek Synergy Neo2 plate reader (BioTek) using monochromator set to excitation/emission of 500 and 542 nm.
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Figure Legends

Fig 1: Intra-sample variability of the SARS-CoV-2 genome in infected individuals. (A) Number of intra- variations observed for each sample analyzed. The red dots represent the 11,362 samples analyzed and the blue violon shows the distribution of the data. (B) Type of variation and sequence context for each intra-sample variable position. Bars represent the percentage of each type. Sequence context is represented by logos comprised of the consensus nucleotides (center) with two nucleotides upstream and two downstream from each intra-sample variable position. (C) Recurrent intra-genetic variations are represented as percentages of samples containing variations at each position. The SARS-CoV-2 genome and its genes are represented by yellow boxes below the graph. The blue line indicates 5% shared variations and was used to extract the recurrent intra-sample variations listed in Table 1. The inset represents a magnification of the cluster identified at the end of the S gene. (D) One-dimension representation of the data shown in panel C for each type of variation individually. The location of the C->A variation at position 25,324 is indicated by a red line in (C) and (D).

Fig 2: Localization of the C->A missense mutation on the SARS-CoV-2 S protein. (A) Schematic representation of the functional domain of the SARS-CoV-2 S protein. (B) Localization of the C->A variation on the carboxy terminal domain (CTD) of the S protein. The mutation is colored and boxed in red. The carboxy terminal domain (CTD) and the ERRS are colored in yellow and black, respectfully in (A) and (B). (C) Distribution of the intra-sample proportion of the C->A transversion at position 25,324 in the 6,668 samples containing this subspecies. The inset
represents the distribution using red dots to represent the samples having this intra-genetic variation and a blue violin to show the distribution of the data.

Fig 3: Heatmap representation for the expression of genes coding for APOBEC and ADAR family members. Counts are represented as Transcripts Per Million (TPM). The blue scale also correlates with TPM values.

Fig 4: Intra-sample variability of the SARS-CoV-2 genome in infected cells. (A) Number of intra-variations observed for each sample analyzed. The red dots represent the 11,362 samples analyzed and the blue violin shows the distribution of the data. (B) Type of variation and sequence context for each intra-sample variable position. Bars represent the percentage of each type. Sequence context is represented by logos comprised of the consensus nucleotides (center) with two nucleotides upstream and two downstream from each intra-sample variable position. (C) Recurrent intra-genetic variations are represented as percentages of samples containing variation at each position. The SARS-CoV-2 genome and its genes are represented by yellow boxes below the graph. The blue line indicates 10% shared variations and was used to extract the intra-sample variation listed in Table 2. The inset represents a magnification of the cluster identified at the end of the S gene. (D) One-dimension representation of the data shown in panel C for each type of variation individually. The location of the C->A variation at position 25,324 is indicated by a red line in (C) and (D).
Fig 5: Effect of SARS-CoV-2 S protein and SΔ20 on cell-to-cell fusion. (A) Phase-contrast and fluorescence microscopy of HEK293T/ACE2 cells expressing GFP with empty vector (pCAGGS), or plasmid expressing wild-type S (pCAGGS(S WT)) or SΔ20 (pCAGGS(SΔ20)). (B) Quantification of syncytia expressing GFP. The area of 129 and 125 syncytia from wild-type S and SΔ20, respectfully, are represented by black dots. The p-value (pv) was calculated using unpaired t-test. (C) Schematic representation of the bi-molecular fluorescence complementation (BiFc) assay used to quantify cell fusion. (D) Quantification of cell fusion using the BiFc assay. Bars represent averages ± standard deviations of five independent experiments and the p-value (pv) was calculated using paired t-test. Schematic of BiFc created with BioRender.com.
Fig 1
Fig 2

A figure showing the structure of the S1 and S2 subunits, with regions labeled as SP, NTD, RBM, FP, HR1, HR2, TM, and CTD. The figure also includes a sequence alignment with highlighted amino acids and a histogram showing C→A variation at nt 25,324 (%/sample).
Fig 4
