

# Negligible impact of SARS-CoV-2 variants on CD4<sup>+</sup> and CD8<sup>+</sup> T cell reactivity in COVID-19 exposed donors and vaccinees.

## AUTHORS

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## SUMMARY

The emergence of SARS-CoV-2 variants highlighted the need to better understand adaptive immune responses to this virus. It is important to address whether also CD4<sup>+</sup> and CD8<sup>+</sup> T cell responses are affected, because of the role they play in disease resolution and modulation of COVID-19 disease severity. Here we performed a comprehensive analysis of SARS-CoV-2-specific CD4<sup>+</sup> and CD8<sup>+</sup> T cell responses from COVID-19 convalescent subjects recognizing the ancestral strain, compared to variant lineages B.1.1.7, B.1.351, P.1, and CAL.20C as well as recipients of the Moderna (mRNA-1273) or Pfizer/BioNTech (BNT162b2) COVID-19 vaccines. Similarly, we demonstrate that the sequences of the vast majority of SARS-CoV-2 T cell epitopes are not affected by the mutations found in the variants analyzed. Overall, the results demonstrate that CD4<sup>+</sup> and CD8<sup>+</sup> T cell responses in convalescent COVID-19 subjects or COVID-19 mRNA vaccinees are not substantially affected by mutations found in the SARS-CoV-2 variants.

## INTRODUCTION

The emergence of several SARS-CoV-2 variants of concern (VOC) with multiple amino acid replacements has implications for the future control of the COVID-19 pandemic (Davies et al., 2020; Kirby, 2021; Tegally et al., 2020; Volz et al., 2021). Variants of concern include the UK (United Kingdom) variant 501Y.V1 lineage B.1.1.7 (Davies et al., 2020), the SA (South Africa) variant 501Y.V2 lineage B.1.351 (Tegally et al., 2020), the BR (Brazilian) variant 501Y.V3 lineage P.1 (Voloch et al., 2020) and the CA (California) variant CAL.20C lineage B.1.427 (Zhang et al., 2021). The B.1.1.7 variant is associated with increased transmissibility (Rambaut et al., 2020; Washington et al., 2021), and similar epidemiological observations have been reported for the SA and BR variants (Tegally et al., 2020; Voloch et al., 2020).

Mutations of greatest concern are present in the viral Spike (S) protein, and include notable mutations in the receptor binding domain (RBD), N-terminal domain (NTD), and furin cleavage site region. Several of these mutations directly affect ACE2 receptor binding affinity, which may impact infectivity, viral load, or transmissibility (Greaney et al., 2021; Starr et al., 2021; Wang et al., 2021a; Zahradnik et al., 2021). Several of the mutations were also noted to be in regions bound by neutralizing antibodies, so it is crucial to address to what extent the mutations associated with the variants impact immunity induced by either SARS-CoV-2 infection or vaccination.

Several reports address the effect of these mutations on antibody binding and function, by either monoclonal or polyclonal antibody responses, and considering both natural infection or vaccination (Edara et al., 2021; Greaney et al., 2021; Muik et al., 2021; Shen et al., 2021; Skelly et al., 2020; Stamatatos et al., 2021; Supasa et al., 2021; Wang et al., 2021a; Wang et al., 2021b; Wibmer et al., 2021; Wu et al., 2021). In general, the impact of the B.1.1.7 variant mutations on neutralizing antibody titers is moderate (Emary et al., 2021; Muik et al., 2021; Shen et al., 2021; Skelly et al., 2020; Supasa et al., 2021; Wu et al., 2021). In contrast, the mutations associated with the B.1.351 and P.1. variants are associated with more pronounced loss of neutralizing capacity (Cele et al., 2021; Skelly et al., 2020; Wang et al., 2021a; Wibmer et al., 2021; Wu et al., 2021). Concerning vaccination responses, the AstraZeneca ChAdOx1 vaccine has been associated with a partial loss of neutralizing antibody activity against B.1.1.7 (Skelly et al., 2020), and a large loss of neutralizing activity against B.1.351 (Voysey et al., 2021). Consistent with these reports, ChAdOx1 maintains efficacy against B.1.1.7 (Emary et al., 2021; Hall et al., 2021), but has a major loss in efficacy against mild COVID-19 with the B.1.351 variant (Voysey et al., 2021). Current epidemiological evidence is that the BNT162b2 Pfizer/BioNTech COVID-19 vaccine retains its efficacy against B.1.1.7 in the UK and in reports from Israel (Amit et al., 2021). Novavax (NVX-CoV2373) has reported differential protective immunity against the parental strain, B.1.1.7, and B.1.351 in vaccine clinical trials (96%, 86%, and 60%) (Novavax Inc., 2021), whereas the Janssen Ad26.COVS.1 1-dose COVID-19 vaccine, which elicits lower neutralizing antibody titers (Sadoff et al., 2021), has relatively similar protection for moderate COVID-19 against both the ancestral strain and B.1.351 (72% and 64%)(FDA, 2021a, b).

Several lines of evidence suggest that CD4<sup>+</sup> and CD8<sup>+</sup> T cell responses play important roles in resolution of SARS-CoV-2 infection and COVID-19 (Sette and Crotty, 2021), including modulating disease severity in humans (Rydyznski Moderbacher et al., 2020; Tan et al., 2021) and reducing viral loads in non-human primates (Munoz-Fontela et al., 2020). Further, persons with agammaglobulinemia or pharmaceutical depletion of B cells generally experience an uncomplicated COVID-19 disease course (Sette and Crotty, 2021; Soresina et al., 2020). Robust CD4<sup>+</sup> and CD8<sup>+</sup> T cell memory is induced after COVID-19 (Breton et al., 2021; Dan et al., 2021; Peng et al., 2020; Wang et al., 2021b), and multiple COVID-19 vaccines elicit CD4<sup>+</sup> and CD8<sup>+</sup> T cell responses (Baden et al., 2021; Dowd et al., 2020; Keech et al., 2020; Sadoff et al., 2021; Voysey et al., 2021). It is therefore key to address the potential impact of SARS-CoV-2 variants mutations on T cell reactivity; however, little data is currently available on this topic (Skelly et al., 2020).

Here, we take a combined experimental and bioinformatics approach to address how SARS-CoV-2 variants of concern impact T cell reactivity. We directly assess T cell responses from persons recovered from COVID-19 obtained before the emergence of the variants, and from recent Moderna mRNA-1273 or Pfizer/BioNTech BNT162b2 vaccinees, for their capacity to recognize peptides derived from the ancestral reference sequence and the B.1.1.7, B.1.351, P.1 and the CAL.20C variants. Bioinformatic analyses were used to predicted the impact of mutations in the various variants with sets of previously

reported CD4<sup>+</sup> and CD8<sup>+</sup> T cell epitopes derived from the ancestral reference sequence (Tarke et al., 2021).

## RESULTS

### Sequence analysis, peptide pool generation and selection of cohorts of COVID-19 convalescent and recent vaccinees

As a first step, we mapped the specific mutations (amino acid replacements and deletions) associated with several of the current variants of concern, including the SARS-CoV-2 B.1.1.7, B1.351, P.1 and the CAL.20C variants, as compared to the SARS-CoV-2 Wuhan ancestral sequence (NCBI acc. no. NC\_045512.2). Briefly, the genomic sequences were downloaded from GISAID, translated using the VIGOR4 tool available on the Virus Pathogen Resource (ViPR) (Pickett et al., 2012), and then compared with the protein sequence of the Wuhan ancestral strain to identify all the possible amino acid changes, as listed in **Table S1**.

Next, we synthesized the corresponding peptides associated with the different variants and generated new peptide pools spanning the full genome sequences of the ancestral Wuhan strain and the respective B.1.1.7, B1.351, P.1 and the CAL20C variants (**Table S2**). As described below, the resulting peptide pools were assessed for their capacity to be recognized by memory T cells responses derived from natural infection in convalescents and vaccinees, and responses to the variant and ancestral genome antigen-specific pools were compared.

Our convalescent donors were adults with ages ranging from 21 to 57 years of age (median 39); 27% were male and 73% female (**Table 1**). SARS-CoV-2 infection in these donors was determined by PCR-based testing during the acute phase of their infection, if available (55% of the cases), and/or seropositivity determined by plasma SARS-CoV-2 S protein RBD IgG ELISA (Stadlbauer et al., 2020)(**Fig. S1**). From these donors, PBMC samples were collected between July to October 2020 period, when the dominant local strain was the ancestral reference virus.

From vaccinated donors, we collected PBMC after recent vaccination with the Moderna mRNA-1273 or the Pfizer/BioNTech BNT162b2 vaccines, approximately 14 days following the second dose administration (**Table 1**). These donors ranged in age from 22 to 67 years (median 43) and 26% were male and 74% were female. All vaccinees had significant RBD IgG titers in the 1843 to 16365 range, consistent with recent vaccination (**Fig. S1**).

### CD4<sup>+</sup> and CD8<sup>+</sup> T cell antigenicity against Spike variant sequences in convalescent samples

We previously described the use of Activation Induced Marker (AIM) assays (Dan et al., 2021; Grifoni et al., 2020b; Mateus et al., 2020; Rydyznski Moderbacher et al., 2020; Tarke et al., 2021) to measure CD4<sup>+</sup> and CD8<sup>+</sup> T cell responses to pools of overlapping peptides spanning the entire sequence of the SARS-CoV-2 antigens. Here, we utilized the same AIM assays using OX40<sup>+</sup>CD137<sup>+</sup> and CD69<sup>+</sup>CD137<sup>+</sup> markers for CD4<sup>+</sup> and CD8<sup>+</sup> T cells reactivity, respectively (Grifoni et al., 2020b; Mateus et al., 2020). As shown in **Fig. 1A-B**, good CD4<sup>+</sup> and CD8<sup>+</sup> T cell reactivity was observed in convalescent donors with pools of overlapping peptides spanning the S protein of the ancestral Wuhan sequence, but also for each of the corresponding variant S pools. Geomean reactivity ranged from 0.09 to the 0.10 for CD4<sup>+</sup> T cells, and 0.08 to the 0.11 for CD8<sup>+</sup> T cells; No significant difference was observed between the pool of S peptides corresponding to the ancestral sequence and those corresponding to the different variants (CD4: UK p=0.90; SA p=0.50; BR p=0.49; CA p=0.85 and CD8: UK p=0.16; SA p=0.07; BR p=0.18; CA p=0.20 by the Wilcoxon test). These values (here and in subsequent graphs) are not corrected for multiple comparisons, as the correction would only decrease the statistical power for detecting a significant difference; therefore, not performing multiple comparison corrections is the more conservative and stringent test.

These T cell analyses were extended using a FluoroSPOT assay system, to measure the capacity of the various pools to elicit functional responses in terms of secretion of IFN $\gamma$  and IL-5 cytokines (**Fig. 1C-D**). As shown in **Fig. 1C**, reactivity was observed for the S pools in convalescent donors in terms of IFN $\gamma$  with geomean reactivity ranging from 32 to 45 Spot Forming Cells (SFC) per million PBMCs. Compared to the ancestral strain, mild decreases in the 24–30% range were noted for B.1.1.7, P.1 and CAL.20C variant pools (UK p=0.01; SA p=0.48; BR p=0.05 and CA p=0.01 by the Wilcoxon test), while

no difference was observed for B.1.351. As expected, no IL-5 reactivity was observed for any of the pools (Fig. 1D).

To further expand these findings, we considered the dose response of the various S pools in terms of stimulation of CD4<sup>+</sup> and CD8<sup>+</sup> T cell specific responses. As shown in Fig. 1E, CD4<sup>+</sup> T cell dose dependent responses for the Wuhan and four variant pools were similar. The same pattern was also observed for CD8<sup>+</sup> T cell responses (Fig. 1F).

### **CD4<sup>+</sup> and CD8<sup>+</sup> T cell antigenicity of proteome-wide SARS-CoV-2 variant sequences in convalescent samples**

As shown in Table S1, mutations found in the variants studied herein were not limited to the Spike protein, but occurred in several additional antigens encoded in the SARS-CoV-2 genome. To address their potential impact on the overall proteome-wide CD4<sup>+</sup> and CD8<sup>+</sup> T cell reactivity, we tested overlapping peptide pools spanning the entire proteome of the ancestral Wuhan sequence in comparison with corresponding pools representing the different variants.

Overall, reactivity to the peptide pools spanning the variant genomes was found to be similar to that against the ancestral Wuhan strain (Fig. 2). When the sum total of reactivity throughout the genome was considered, no differences or decreases in reactivity compared to the ancestral were noted for the variant pools (CD4: UK p=0.58; SA p= 0.46; BR p= 0.27; CA p= 0.08 and CD8: UK p= 0.25; SA p= 0.15; BR p= 0.02; CA p= 0.30 by the Wilcoxon test uncorrected p values) (Fig. 2A-B).

We previously showed that in COVID-19 convalescent subjects a set of 10 different antigens (nsp3, nsp4, nsp6, nsp12, nsp13, S, ORF3a, M, ORF8 and N) account for 83 and 81% of the total CD4<sup>+</sup> and CD8<sup>+</sup> T cell responses, respectively (Tarke et al., 2021). Here a similar overall pattern of dominant antigens was observed. When single proteins are considered, no variant pool showed a decrease in reactivity when a multi-hypothesis testing correction was applied (Fig. 2C-D). It is worth noting that this specific comparison is for illustration purposes only, as this study is not fully powered to rule out minor differences that could be observed in the individual antigens.

In conclusion, these experiments suggest that memory CD4<sup>+</sup> or CD8<sup>+</sup> T cells from individuals that have been infected with the ancestral SARS-CoV-2 strain recognize the ancestral reference strain and the variant genome-wide sequences with similar efficiency.

### **CD4<sup>+</sup> and CD8<sup>+</sup> T cell antigenicity against Spike variant sequences in recent vaccines samples**

We also studied T cell responses by individuals who received authorized mRNA COVID-19 vaccines. We focused our analysis on T cell reactivity to peptide pools spanning the Spike antigen of the ancestral strain, which is the basis of the presently used vaccines. For both CD4<sup>+</sup> and CD8<sup>+</sup> T cell reactivity, the magnitude of responses to pools encompassing the sequences from the ancestral Wuhan genome and the different variants considered range from a geomean of 0.15 to 0.19 for CD4<sup>+</sup> T cells and a geomean of 0.16-0.24 for CD8<sup>+</sup> T cells. Comparison of the variant pools to the ancestral sequence showed no significant difference for CD4<sup>+</sup> T and CD8<sup>+</sup> T cells reactivity, with the exception of the B.1.351 pools, where mild decreases of 29% and 33%, respectively, were observed (CD4: UK p=0.47; SA p=0.01; BR p=0.91; CA p=0.41; CD8: UK p=0.03; SA p=0.001; BR p=0.15 and CA p=0.02 by the Wilcoxon test) (Fig. 3A-B).

The results from the FluoroSPOT assay system (Fig. 3C-E) showed good reactivity in terms of IFN $\gamma$ , with geomean reactivity ranging from 54 to the 70 SFC per million PBMCs (Fig. 3C). Minimal IL-5 responses were observed, with geomean reactivity ranging from 22 to 25 SFC/10<sup>6</sup>, which is slightly above the limit of detection (Fig. 3D). On a per donor basis, the IFN $\gamma$  response was found to account for more than 80% of the total response, on average (range 81% to 87%), irrespective of whether the ancestral strain or any of the variants was considered (Fig. 3E).

Similar to the experiments in convalescent donors, we also considered the dose response of the various Spike pools in terms of stimulation of CD4<sup>+</sup> or CD8<sup>+</sup> T cell specific responses for vaccinees. As shown in Fig. 3F-G, CD4<sup>+</sup> and CD8<sup>+</sup> T cell dose responses for the ancestral pools and the four variant pools were similar. Taken together these results indicate that the responses to the ancestral and variant Spike pools are similar for both CD4<sup>+</sup> and CD8<sup>+</sup> T cells in mRNA vaccinees.

### **Conservation analysis of sets of defined CD4<sup>+</sup> and CD8<sup>+</sup> T cell epitopes**

We recently reported a comprehensive study of epitopes recognized in convalescent subjects, leading to the identification of 280 different CD4<sup>+</sup> T cell epitopes (Tarke et al., 2021). Here, we analyzed how many of those epitopes would be impacted by mutations in the different variants. As shown in **Fig. 4A**, we found that 89.6%, 90%, 94.3% and 97.1% (average 93%) of the CD4<sup>+</sup> T cell epitopes identified by Tarke et al. are conserved in the B.1.1.7, B.1.351, P.1 and the CAL20C variants. A similar pattern is observed when the magnitude of responses associated with the various epitopes is considered, rather than the simple number of epitopes (**Fig. 4B**). The fully conserved CD4<sup>+</sup> T cell epitopes account for 84.4%, 88.1%, 95.7% and 97.8% (average 91.5%) of the total response, when comparing the B.1.1.7, B.1.351, P.1 and the CAL.20C variants, respectively.

The study of Tarke et al. also reported the identification of 523 CD8<sup>+</sup> T cell epitope associated with unique HLA restrictions (Tarke et al., 2021). Performing a similar analysis as above, (**Fig. 4C**) we found that 508 (97.1%) of these 523 CD8<sup>+</sup> T cell epitopes are totally conserved within the B.1.1.7 variant, 509 (97.3%) are conserved within the B.1.351 variant, 509 (97.3%) are conserved within the P.1 variant and 512 (97.9%) within the CAL.20C variant. Similarly, in terms of magnitude of CD8<sup>+</sup> T cell responses associated with the various epitopes, totally conserved CD8<sup>+</sup> T cell epitopes account for 98.3%, 98.4%, 97.9% and 97.8% of the total responses (**Fig. 4D**) for the B.1.1.7, B.1.351, P.1 and the CAL.20C variants respectively, with an average of 98.1%.

Finally, we analyzed the degree of CD4<sup>+</sup> and CD8<sup>+</sup> T cell epitope conservation if we restricted our analysis only to epitopes contained in the Spike antigen. The number of S-derived epitopes conserved at 100% sequence identity was, on average, 84.5% for the CD4<sup>+</sup> T cell epitopes (**Fig. 4E**), and 95.3% for the CD8<sup>+</sup> T cell epitopes (**Fig. 4G**). Similarly, in terms of magnitude of CD4<sup>+</sup> T cell responses associated with the various S epitopes, totally conserved CD4<sup>+</sup> T cell epitopes account for 95.5%, 75.3%, 89.8% and 98.3% of the total responses (**Fig. 4F**) for the B.1.1.7, B.1.351, P.1 and the CAL.20C variants respectively, with an average of 89.7%. In terms of the magnitude of CD8<sup>+</sup> T cell responses, totally conserved epitopes account for 95.2%, 97.6%, 95.4% and 97.3% of the total responses (**Fig. 4H**) for the B.1.1.7, B.1.351, P.1 and the CAL.20C variants respectively, with an average of 96.4%.

While the restriction of the HLA class II epitopes in the Tarke et al. (Tarke et al., 2021) study was not unequivocally assigned, the restriction of the class I epitopes is implicitly inferred based on HLA allele specific predictions and testing in HLA matched donors. Accordingly, we further analyzed the extent to which the affected epitopes would be impacted by their respective associated mutation by determining, for each epitope/matching epitope variant, their predicted binding affinity for the corresponding putative HLA class I restriction element. Predicted binding capacity was determined using the NetMHCpan BA 4.1 tool provided by the IEDB's analysis resource (Dhanda et al., 2019; Reynisson et al., 2020).

In the case of the B.1.1.7, B.1.351, P.1 and the CAL.20C variants, the % of mutations associated with no decrease in binding capacity, conservatively defined as a 2-fold reduction, was 73.3%, 78.6%, 78.6% and 45.5%, respectively (with the CAL.20C variant having a smallest number of total mutations, noted above) (**Fig. S2** and **Table S3**). In conclusion, the analyses suggest that the vast majority of CD8<sup>+</sup> T cell epitopes are unaffected by mutations found in all the different variants. The corresponding mutations are also predicted to have minor effects on the total T cell response, thus providing a molecular basis for the marginal impact on T cell reactivity by COVID-19 convalescent subjects and recipients of COVID-19 mRNA vaccines.

## DISCUSSION

The present study addresses a key knowledge gap pertaining to the potential of emergent SARS-CoV-2 variants to evade recognition by human immune responses. We focused on T cell responses elicited by either natural infection or vaccination with the Pfizer/BioNTech and Moderna COVID-19 mRNA vaccines. We found negligible effects on both CD4<sup>+</sup> or CD8<sup>+</sup> T cell responses to all four variants investigated, to include the B.1.1.7, B.1.351, P.1 and CAL.20C variants found in the UK, South Africa, Brazil and California, respectively. To more comprehensively assess T cell functionalities, the comparison between the original Wuhan isolate and the variants was performed utilizing different T cell methodologies, such as the AIM assay (quantifying T cells with a range of functionalities), and the FluoroSPOT assay (quantifying cells with specific cytokine-secreting activity). We also tested whether any of the variant sequences might be associated with an altered cytokine polarization; marginal IL-5

production was detected in any of the conditions tested. This is relevant, since it was reported that single amino acid replacements in an epitope sequence can lead to a change in the cytokines produced (Evavold and Allen, 1991; Sloan-Lancaster and Allen, 1996), and a Th2-like response pattern was initially hypothesized to be linked to adverse outcomes in SARS-specific responses (Peeples, 2020).

The data provide some positive news in light of justified concern over the impact of SARS-CoV-2 variants of concern on efforts to control and eliminate the present pandemic. Undoubtedly, several of the variants are associated with increased transmissibility, and also have been associated with decreased susceptibility to neutralizing antibodies from infected or vaccinated individuals. In contrast, the data presented here suggests that T cell responses are largely unaffected by the variants. While it is not anticipated that circulating memory T cells would be effective in preventing SARS-CoV-2 infection, it is plausible that they can reduce COVID-19 severity (Lipsitch et al., 2020; Sette and Crotty, 2021). Several lines of evidence support this notion, such as observations that early SARS-CoV-2 T cell responses are associated with milder COVID-19 (Rydyznski Moderbacher et al., 2020; Tan et al., 2021). Thus, the T cell response may contribute to limiting COVID-19 severity induced by variants that partially or largely escape neutralizing antibodies. This is consistent with T cell mediated immunity observed in humans against a different respiratory pathogen, influenza, for which heterologous immunity against diverse influenza strains is associated with memory T cells to conserved epitopes (Greenbaum et al., 2009; Sridhar et al., 2013; Wilkinson et al., 2012).

Our data also provide a molecular basis for the lack of impact of the mutations associated with the variants analyzed on T cell responses. Prior reports have identified a large number of T cell epitopes recognized throughout the SARS-CoV-2 proteome, including Spike (Ferretti et al., 2020; Keller et al., 2020; Le Bert et al., 2020; Nelde et al., 2020; Peng et al., 2020; Snyder et al., 2020). We furthered this point by an analysis of the Tarke et al. data set, showing that 93% of CD4<sup>+</sup> T cell, and 97% of CD8<sup>+</sup> T cell, epitopes are completely conserved in the variants. Further, we found that even in the epitopes affected by single mutations, no negative affect HLA binding capacity in the majority of cases is expected. The apparent higher conservation of CD8<sup>+</sup> T cell epitopes is to be expected based on the shorter length of HLA class I binding peptides (usually 9-10 amino acids) as compared to their class II counterparts (13-17). This effect is counterbalanced by CD8<sup>+</sup> T cells being generally less tolerant of amino substitutions as compared to CD4<sup>+</sup> T cells (Grifoni et al., 2020a; Weiskopf et al., 2014). Overall, we observed that the effect of the variant mutations on the global CD4<sup>+</sup> and CD8<sup>+</sup> T cell responses was negligible.

Mutations associated with the variants could be reflective of adaptation in terms of optimizing replication or binding to ACE2, but also reflective of adaptation to escape immune recognition by antibodies (Andreano et al., 2020; Starr et al., 2020; Wang et al., 2021a; Wang et al., 2021b; Zahradnik et al., 2021). Indeed, higher viral binding to a cellular receptor can be a mechanism of compensatory viral evolution in the presence of neutralizing antibodies (Hensley et al., 2009). In that respect, while mutation to escape antibody binding has been well documented for influenza (Andrews et al., 2015; Doud et al., 2018; Krammer et al., 2018) and SARS-CoV-2, immune escape at the level of T cell responses in human populations has not been reported for other acute respiratory infections. Because of HLA polymorphism, the epitope repertoire recognized is likely to be substantially different from one individual to the next, greatly decreasing the likelihood of immune escape by an acute virus. An advantage conferred to the virus by a mutation in a person would not be linked to an immune response escape advantage in a non-HLA matched individual. At the same time, our data does not rule out that each person could be strongly affected by the mutations of specific variants. For SARS-CoV-2, this property of T cell recognition is further enhanced by the fact that the T cell responses against SARS-CoV-2 are highly multi-antigenic and multi-specific, with tens of different epitopes recognized by CD4<sup>+</sup> and CD8<sup>+</sup> T cells in a given individual (Braun et al., 2020; Ferretti et al., 2020; Nelde et al., 2020; Tarke et al., 2021).

The results here have potential implications for engineering coronavirus vaccines with broader protective immunity against variants of concern. Clearly the most straightforward path is to update the current vaccines to target a variant Spike, given how highly successful several COVID-19 vaccines have proven to be against the parental SARS-CoV-2 strain. Our results suggest that a parallel alternative approach could involve inclusion of additional antigens and epitopes, perhaps selected on the basis of low mutational propensity (Gaiha et al., 2019), to ensure that neutralizing antibodies are complemented with T cell responses to minimize COVID-19 morbidity and mortality.

### ***Limitations and future directions.***

The present study did not assess decreases in antibody reactivity, as several other studies have already investigated this matter (Edara et al., 2021; Greaney et al., 2021; Muik et al., 2021; Shen et al., 2021; Skelly et al., 2020; Stamatatos et al., 2021; Supasa et al., 2021; Wang et al., 2021a; Wang et al., 2021b; Wibmer et al., 2021; Wu et al., 2021). Further, our studies utilized overlapping peptide pools, and as such we could not exclude that some of the mutations might involve alterations in terms of antigen processing for either class I or class II, which would be undetected by using pools of “preprocessed” peptides. The number of donors studied was also limited, although variability in T cell reactivity suggestive of large variant-associated effects were not observed. While we have no reason to suspect that substantial differences might exist between the epitope specificity of responses elicited by different vaccines, our study did not address this point. Our study was designed to test for differences in overall response to the different variants and was not powered or designed to investigate differences between the mRNA vaccines. Furthermore, in our study we tested samples from convalescent donors infected before October 2020; thus, it is unlikely that any of the donors would have been infected by any of the variants, as this date precedes their diffusion to appreciable degree in the US and California. Finally, we have only investigated whether responses induced by the ancestral reference sequence are able to cross-recognize variant sequences, as this is relevant to the current situation. We have not examined whether responses induced by an infection with a variant sequence will be able to cross-recognize the ancestral reference sequence present in the currently approved vaccines.

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## **AUTHOR CONTRIBUTIONS**

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## FIGURE LEGENDS

**Figure 1. T cell responses of COVID-19 convalescent individuals against SARS-CoV-2 Spike for the different variants.** PBMCs of COVID-19 convalescent individuals (n=11) were stimulated with the Spike MPs corresponding to the ancestral reference strain (Wu, black) and the B.1.1.7 (UK, grey), B.1.351 (SA, red), P.1 (BR, orange) and CAL.20C (CA, light blue) SARS-CoV-2 variants. **A)** Percentages of AIM<sup>+</sup> (OX40<sup>+</sup>CD137<sup>+</sup>) CD4<sup>+</sup>T cells. **B)** Percentages of AIM<sup>+</sup> (CD69<sup>+</sup>CD137<sup>+</sup>) CD8<sup>+</sup> T cells. **C)** IFN $\gamma$  spot forming cells (SFC) per million PBMCs **D)** IL-5 SFC per million PBMCs. Paired comparisons of Wuhan S MP versus each of the variants were performed by Wilcoxon test and are indicated by the p values in panels **A-C**. The data shown in panels **A** and **B** are plotted to show the Spike MPs titration (1  $\mu$ g/mL, 0.1  $\mu$ g/mL, 0.01  $\mu$ g/mL) for CD4<sup>+</sup> (**E**) and CD8<sup>+</sup> (**F**) T cells in each SARS-CoV-2 variant and the geometric mean of the 0.1 $\mu$ g/mL condition is listed above each titration. In all panels, the bars represent the geometric mean.

**Figure 2. T cell responses of COVID-19 convalescent individuals against SARS-CoV-2 proteome for the different variants.** PBMCs of the COVID-19 convalescent individuals (n=11) were stimulated with the MPs for the entire viral proteome corresponding to the ancestral reference strain (Wu, black) and the B.1.1.7 (UK, grey), B.1.351 (SA, red), P.1 (BR, orange) and CAL.20C (CA, light blue) SARS-CoV-2 variants. **A)** Percentages of AIM<sup>+</sup> (OX40<sup>+</sup>CD137<sup>+</sup>) CD4<sup>+</sup> T cells for the total reactivity. **B)** Percentages of AIM<sup>+</sup> (CD69<sup>+</sup>CD137<sup>+</sup>) CD8<sup>+</sup> T cells for the total reactivity. Bars represent the geometric mean. Paired comparisons of Wuhan versus each of the variants were performed by Wilcoxon tests. **C)** Percentages of AIM<sup>+</sup> (OX40<sup>+</sup>CD137<sup>+</sup>) CD4<sup>+</sup> T cells for each MP **D)** Percentages of AIM<sup>+</sup> (OX40<sup>+</sup>CD137<sup>+</sup>) CD4<sup>+</sup> T cells for each MP.

**Figure 3. T cell responses of COVID-19 vaccinee individuals against SARS-CoV-2 Spike for the different variants.** PBMCs of Pfizer/BioNTech BNT162b2 (n=8, triangles) and Moderna mRNA-1273 COVID-19 vaccines (n=11, circles) were stimulated with the Spike MPs corresponding to the ancestral reference strain (Wu, black) and the B.1.1.7 (UK, grey), B.1.351 (SA, red), P.1 (BR, orange) and CAL.20C (CA, light blue) SARS-CoV-2 variants. **A)** Percentages of AIM<sup>+</sup> (OX40<sup>+</sup>CD137<sup>+</sup>) CD4<sup>+</sup> T cells. **B)** Percentages of AIM<sup>+</sup> (CD69<sup>+</sup>CD137<sup>+</sup>) CD8<sup>+</sup> T cells. **C)** IFN $\gamma$  spot forming cells (SFC) per million PBMCs **D)** IL-5 Spot forming cells (SFC) per million PBMCs. **E)** Percentages of IFN $\gamma$  were calculated from the total IFN $\gamma$  and IL-5 SFC per million PBMCs. Paired comparisons of the ancestral reference strain-based S MP versus each of the variants were performed by Wilcoxon test and are indicated by the p values in panels **A-E**. The data shown in panels **A** and **B** are also plotted showing the spike MPs titration (1  $\mu$ g/mL, 0.1  $\mu$ g/mL, 0.01  $\mu$ g/mL) for CD4<sup>+</sup> (**F**) and CD8<sup>+</sup> (**G**) T cells in each SARS-CoV-2 variant. The geometric mean of the 0.1 $\mu$ g/mL condition is listed above each titration. In all panels, the bars represent the geometric mean.

**Figure 4. SARS-CoV-2 T cell epitope sequences affected by the variants.** CD4<sup>+</sup> and CD8<sup>+</sup> T cell epitopes of the ancestral strain identified in a previous study (Tarke et al.) are analyzed as a function of the number and percentage of response that are or are not conserved across the B.1.1.7 (UK, grey), B.1.351 (SA, red), P.1 (BR, orange) and CAL.20C (CA, light blue) SARS-CoV-2 variants. The SARS-CoV-2 epitopes for the most immunodominant SARS-CoV-2 proteins in terms of numbers and percentage of response are shown for CD4<sup>+</sup> (**A-B**) and CD8<sup>+</sup> (**C-D**) T cells. The SARS-CoV-2 epitopes for the Spike protein only in terms of numbers and percentage of response are shown for CD4<sup>+</sup> (**E-F**) and CD8<sup>+</sup> (**G-H**) T cells.

## TABLES

**Table 1.** Characteristics of donor cohorts.

	<b>COVID-19 (n = 11)</b>	<b>Vaccinees (n = 19)</b>
<b>Age (years)</b>	21-57 [Median = 39, IQR = 33]	22-67 [Median = 43, IQR = 36]
<b>Gender</b>		
Male (%)	27% (3/11)	26% (5/19)
Female (%)	73% (8/11)	74% (14/19)
<b>Sample Collection Date</b>	July–Oct 2020	Jan–Feb 2021
<b>SARS-CoV-2 PCR</b>	Positive = 83% (5/6) Not tested = 45% (5/11)	N/A
<b>S RBD IgG Positive</b>	100% (11/11)	100% (19/19)
<b>Peak disease Severity<sup>a</sup></b>		
Mild	100% (11/11)	
Moderate	0% (0/11)	N/A
Severe	0% (0/11)	
Critical	0% (0/11)	
<b>Race-Ethnicity</b>		
White- not Hispanic or Latino	82% (9/11)	42% (8/19)
Hispanic or Latino	9% (1/11)	16% (3/19)
Asian	9% (1/11)	42% (8/19)
American Indian/Alaska Native	0% (0/11)	0% (0/19)
Not reported	0% (0/11)	0% (0/19)
<b>Days at Collection</b>	38-80 (11/11) [Median =50, IQR = 45] <sup>b</sup>	13-30 (8/19) Pfizer 12-15 (11/19) Moderna [Median = 14, IQR = 14] <sup>c</sup>

<sup>a</sup> According to WHO criteria.

<sup>b</sup> Post Symptom Onset

<sup>c</sup> 2<sup>nd</sup> dose of vaccination

## SUPPLEMENTAL MATERIALS

### SUPPLEMENTAL FIGURE LEGENDS

**Figure S1. SARS-COV-2 serology of the all the cohorts analyzed in this study.** Related to Figures 1, 2 and 3 and Table 1. Spike RBD serology in COVID-19 convalescents (n=11) and COVID-19 vaccines (n=19).

**Figure S2. Effect of mutations on CD8 epitope predicted HLA class I binding capacity.** Related to Figure 4. For each CD8<sup>+</sup> T cell epitope associated with a mutation found in the respective variants, the predicted HLA binding capacity of original sequence and the mutated sequence was calculated. Based on the results, each instance was categorized as a function of whether the binding capability of the mutated peptide is increased (>2-fold), neutral or decreased (<2-fold). Each analysis is done separately for the B.1.1.7 (UK, grey), B.1.351 (SA, red), P.1. (BR, orange) and CAL.20C (CA, light blue) SARS-CoV-2 variants.

**Figure S3. Gating strategy.** Related to Figures 1, 2, 3 and 4. Representative graphs illustrating the gating strategy used in the flow cytometry AIM assays in order to define antigen-specific CD4<sup>+</sup> (outlined in blue) and CD8<sup>+</sup> (outlined in red) T cells by the expression of OX40<sup>+</sup>CD137<sup>+</sup> and CD69<sup>+</sup> CD137<sup>+</sup>, respectively. These graphs depict one of the COVID-19 convalescent donors from this study and are representative of the gating strategy utilized with all donors tested.

## SUPPLEMENTAL TABLE LEGENDS

**Table S1.** Related to Figures 1, 2, 3 and 4. List of amino acid positions and relative amino acid changes in the different variants studied with respect to the ancestral Wuhan strain.

**Table S2.** Related to Figures 1, 2, 3 and 4. List of mutated peptides with respect to the ancestral Wuhan strain in the different variants studied.

**Table S3.** Related to Figure 4. Effect of mutations on CD8 epitope HLA class I binding capacity.

## STAR METHODS

### RESOURCE AVAILABILITY

#### Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Dr. Alessandro Sette ([alex@lji.org](mailto:alex@lji.org)).

#### Materials Availability

Aliquots of synthesized sets of peptides utilized in this study will be made available upon request. There are restrictions to the availability of the peptide reagents due to cost and limited quantity.

#### Data and Code Availability

The published article includes all data generated or analyzed during this study, and summarized in the accompanying tables, figures and supplemental materials.

### EXPERIMENTAL MODEL AND SUBJECT DETAILS

#### Human Subjects

*Convalescent COVID-19 Donors.* Convalescent donors were enrolled at either a UC San Diego Health clinic under the approved IRB protocols of the University of California, San Diego (UCSD; 200236X), or at the La Jolla Institute (LJI; VD-214). All donors were California residents and samples were collected from August to October 2020, before any of the SARS-CoV-2 variants described herein had been detected in California. These donors were referred to the study by a health care provider or were self-referred. The CRO BioIVT provided additional cohorts of COVID-19 convalescent donors who had been confirmed positive for COVID-19 by PCR following the resolution of symptoms. The total cohort of convalescent donors represented both sexes (27% male, 73% female) and ranged from 21 to 57 years of age (median 39 years). All samples were confirmed seropositive against SARS-CoV-2 by ELISA, as described below. Details of this convalescent COVID-19 cohort are listed in Table 1. All convalescent COVID-19 donors provided informed consent to participate in the present and future studies at the time of enrollment.

*COVID-19 vaccinees.* The La Jolla Institute recruited 19 healthy adults who had received the first and second dose of the Pfizer/BioNTech BNT162b2 (n=8) or Moderna mRNA-1273 COVID-19 vaccines (n=11). Blood draws took place under IRB approved protocols two to four weeks after the second dose of the vaccine was administered. All donors had their SARS-CoV-2 antibody titers measured by ELISA, as described below. The cohort of vaccinees represented ranged from 22 to 67 years of age (median 43 years) and represented both sexes (26% male, 74% female). At the time of enrollment in the study, all donors gave informed consent.

### METHOD DETAILS

#### Isolation of peripheral blood mononuclear cells (PBMCs) and plasma

Collection and processing of blood samples was performed as previously described (Dan et al., 2021; Tarke et al., 2021). Briefly, whole blood was collected in heparin coated blood bags or in ACD tubes and centrifuged for 15 minutes at 1850 rpm to separate the cellular fraction from the plasma. The plasma was then removed and stored at -20°C. The cellular fraction next underwent density-gradient sedimentation using Ficoll-Paque (Lymphoprep, Nycomed Pharma, Oslo, Norway) to separate the PBMCs as previously described (Weiskopf 2013). Isolated PBMCs were cryopreserved in cell recovery media containing 10% DMSO (Gibco), supplemented with 90% heat inactivated fetal bovine serum (FBS; Hyclone Laboratories, Logan UT) and stored in liquid nitrogen until used in the assays.

#### SARS-CoV-2 RBD ELISA

Serology to SARS-CoV-2 was determined for all donor cohorts as previously described (Rydzynski Moderbacher et al., 2020). Briefly, 96-well half-area plates (ThermoFisher 3690) were coated with 1 ug/mL SARS-CoV-2 Spike (S) Receptor Binding Domain (RBD) and incubated at 4°C overnight. The next day plates were blocked at room temperature for 2 hours with 3% milk in phosphate buffered saline (PBS)

containing 0.05% Tween-20. Heat-inactivated plasma was added to the plates for an additional 90-minute incubation at room temperature followed by incubation with the conjugated secondary antibody, detection, and subsequent data analysis by reading the plates on Spectramax Plate Reader at 450 nm using the SoftMax Pro. The limit of detection (LOD) was defined as 1:3. Limit of sensitivity (LOS) for SARS-CoV-2 infected individuals was established based on uninfected subjects, using plasma from normal healthy donors not exposed to SARS-CoV-2.

### **Mutation analysis of SARS-CoV-2 UK, California, South Africa and Brazil variants**

Genome sequences for the variant viruses were downloaded from GISAID. These sequences were screened to select those without ambiguous residues and generated from Illumina sequencing technologies using an in-house sequence QC script. The selected genomic sequences were then translated into protein amino acid sequences using the VIGOR4 tool available on the Virus Pathogen Resource (ViPR)(Pickett et al., 2012). Sequence variations in the variant viruses were derived by comparison with Wuhan-1 (NC\_045512.2). One or more representative sequences were considered for the UK (EPI\_ISL\_601443), Brazilian (EPI\_ISL\_804823), Californian (EPI\_ISL\_847619; EPI\_ISL\_847621; EPI\_ISL\_847643) and South Africa (EPI\_ISL\_660629; EPI\_ISL\_736930; EPI\_ISL\_736932; EPI\_ISL\_736944; EPI\_ISL\_736966; EPI\_ISL\_736971; EPI\_ISL\_736973; EPI\_ISL\_825104; EPI\_ISL\_825120; EPI\_ISL\_825131) variants. A summary of all the amino acids mutated in the different variants respect to the Wuhan sequence and considered in this study is available in **Table S1**.

### **SARS-CoV-2 Wuhan and variant peptide synthesis and pooling**

Peptides were synthesized that spanned entire SARS-CoV-2 proteins and corresponded to the ancestral Wuhan sequence or the B.1.1.7 (UK), B.1.351 (SA), P.1 (BR) and CAL.20C (CA) SARS-CoV-2 variants. Peptides were 15-mers overlapping by 10 amino acids and were synthesized as crude material (TC Peptide Lab, San Diego, CA). All peptides were individually resuspended in dimethyl sulfoxide (DMSO) at a concentration of 10–20 mg/mL. Megapools (MP) for each antigen were created by pooling aliquots of these individual peptides, undergoing another lyophilization, and resuspending in DMSO at 1 mg/mL.

### **Bioinformatic analysis of T cell epitopes**

The binding capacity of SARS-CoV-2 T cell epitopes, and their corresponding variant-derived peptides, for their putative HLA class I restricting allele(s) was determined utilizing the NetMHCpan BA 4.1 algorithm (Reynisson et al., 2020), as implemented by the IEDB's analysis resource (Dhanda et al., 2019; Vita et al., 2019). Predicted binding is expressed in terms of IC<sub>50</sub> nM. For each epitope-variant pair a ratio of affinities (WT/variant) was determined. Ratios >2, indicating a 2-fold or greater increase in affinity due to the mutation, were categorized as an increase in binding capacity, and <0.5 as a decrease; ratios between 0.5 and 2 were designated as neutral.

### **Flow cytometry-based AIM assay**

Activation induced cell marker (AIM) assay has previously been described in detail elsewhere (da Silva Antunes et al., 2021; Dan et al., 2021; Reiss et al., 2017). In summary, PBMCs were cultured for in the presence of SARS-CoV-2 specific (Wuhan or variant) MPs [1 µg/ml] in 96-well U-bottom plates at a concentration of 1x10<sup>6</sup> PBMC per well. As a negative control, an equimolar amount of DMSO was used to stimulate the cells in triplicate wells and as positive controls phytohemagglutinin (PHA, Roche, 1µg/ml) and a cytomegalovirus MP (CMV, combining CD4 and CD8 MPs, 1µg/ml) were also included. After incubation for 20–24 hours at 37°C, 5% CO<sub>2</sub>, the cells were stained with CD3 BUV805 or CD3 AF700 (4:100 or 4:100; BD Biosciences Cat# 612895 or Life Technologies Cat# 56-0038-42, respectively), CD4 BV605 (4:100; BD Biosciences Cat# 562658), CD8 BUV496 or BV650 (2:100 or 4:100; BD Biosciences Cat# 612942 or Biolegend Cat# 301042), and Live/Dead eFluor506 (5:1000; eBioscience Cat# 65-0866-14). Cells were also stained to measure activation with the following markers: CD137 APC (4:100; Biolegend Cat# 309810), OX40 PE-Cy7 (2:100; Biolegend Cat#350012), and CD69 PE (10:100; BD Biosciences Cat# 555531). All samples were acquired on a ZE5 5-laser or 4-laser cell analyzer (Bio-rad laboratories) and analyzed with FlowJo software (Tree Star). In the resulting data generated from the AIM assays, the background was removed from the data by subtracting the average of the % of AIM<sup>+</sup>

cells plated in triplicate wells stimulated with DMSO. The Stimulation Index (SI) was calculated by dividing the % of AIM<sup>+</sup> cells after SARS-CoV-2 stimulation with the average % of AIM<sup>+</sup> cells in the negative DMSO control. An SI greater than 2 and a minimum of 0.02 % or 0.03 % AIM<sup>+</sup> CD4<sup>+</sup> or CD8<sup>+</sup> cells, respectively, after background subtraction was considered to be a positive response. The gates for AIM<sup>+</sup> cells were drawn relative to the negative and positive controls for each donor. A representative example of the gating strategy is depicted in **Fig. S3**.

### **FluoroSPOT assays**

96-well FluoroSpot plates were coated with anti-cytokine antibodies for IFN $\gamma$  and IL-5 (mAbs 1-D1K and TRFK5, respectively; Mabtech, Stockholm, Sweden) at a concentration of 10 $\mu$ g/mL. PBMCs were stimulated in triplicate at a density of 200x10<sup>3</sup> cells/well with S MPs corresponding to each of the SARS-CoV-2 variants analyzed (1 $\mu$ g/mL), PHA (1 $\mu$ g/mL), and DMSO (0.1%), as positive and negative controls respectively. After 20 hours of incubation at 37°C, 5% CO<sub>2</sub>, cells were discarded and plates were washed before the addition of cytokine antibodies (mAbs 7-B6-1-BAM and 5A10-WASP; Mabtech, Stockholm, Sweden). After a 2-hour incubation, plates were washed again with PBS/0.05% Tween20 and incubated for 1 hour with fluorophore-conjugated antibodies (Anti-BAM-490 and Anti-WASP-640). An AID iSPOT FluoroSpot reader (AIS-diagnostika, Germany) was used to count the fluorescent spots that resulted from cells secreting IFN $\gamma$  and IL-5. Each peptide MP was considered positive compared to the DMSO negative control based on the following criteria: 20 or more spot forming cells (SFC) per 10<sup>6</sup> PBMC after subtraction, a stimulation index (S.I.) greater than 2, and a p value <0.05 by either a Poisson or T test calculated between the triplicates of the MP and the relative negative control.

### **QUANTIFICATION AND STATISTICAL ANALYSIS**

Data and statistical analyses were performed in FlowJo 10 and GraphPad Prism 8.4, unless otherwise stated. Statistical details of the experiments are provided in the respective figure legends and in each method section pertaining the specific technique applied. Data plotted in logarithmic scales are expressed as geometric mean. Statistical analyses were performed using Wilcoxon matched-pairs signed rank test for paired comparisons. Multi-hypothesis testing corrections (MHTC) have not been applied in the study by design. The study is not designed or powered to address differences across different proteins. The primary hypothesis is that no significant differences are observed across the different variants, and this is more stringently addressed avoiding to correct for MHTC, since a difference that is not significant would remain so even after corrections. Therefore, reporting the data without applying MHTC is a more stringent criterion which is appropriately being applied in this case to avoid false negatives. Details pertaining to significance are also noted in the respective figure legends.

### **DECLARATION OF INTEREST**

A.S. is a consultant for Gritstone, Flow Pharma, Oxford Immunotech, Caprion, Arcturus and Avalia. S.C. is a consultant for Avalia. All other authors declare no conflict of interest. LJI has filed for patent protection for various aspects of vaccine design and identification of specific epitopes.

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Fig. 1

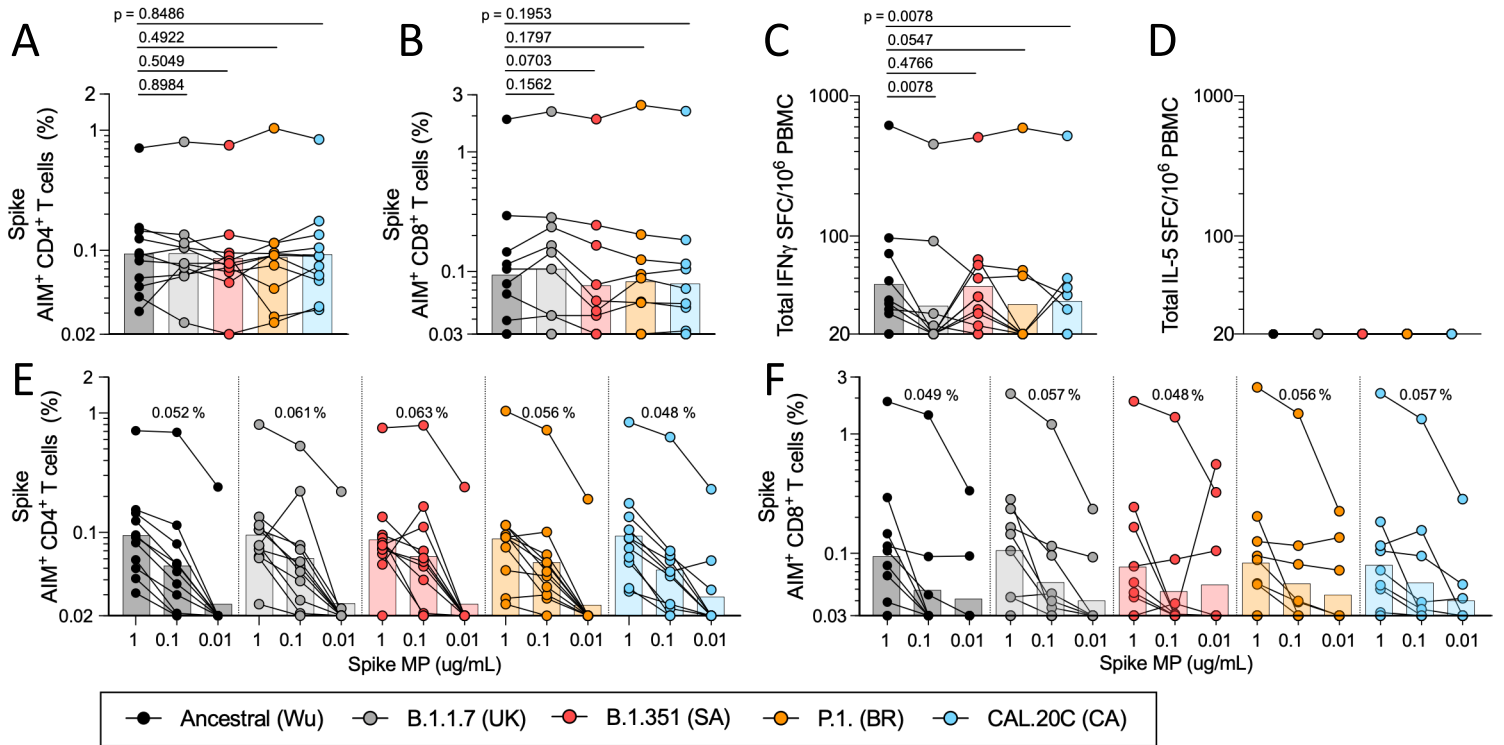
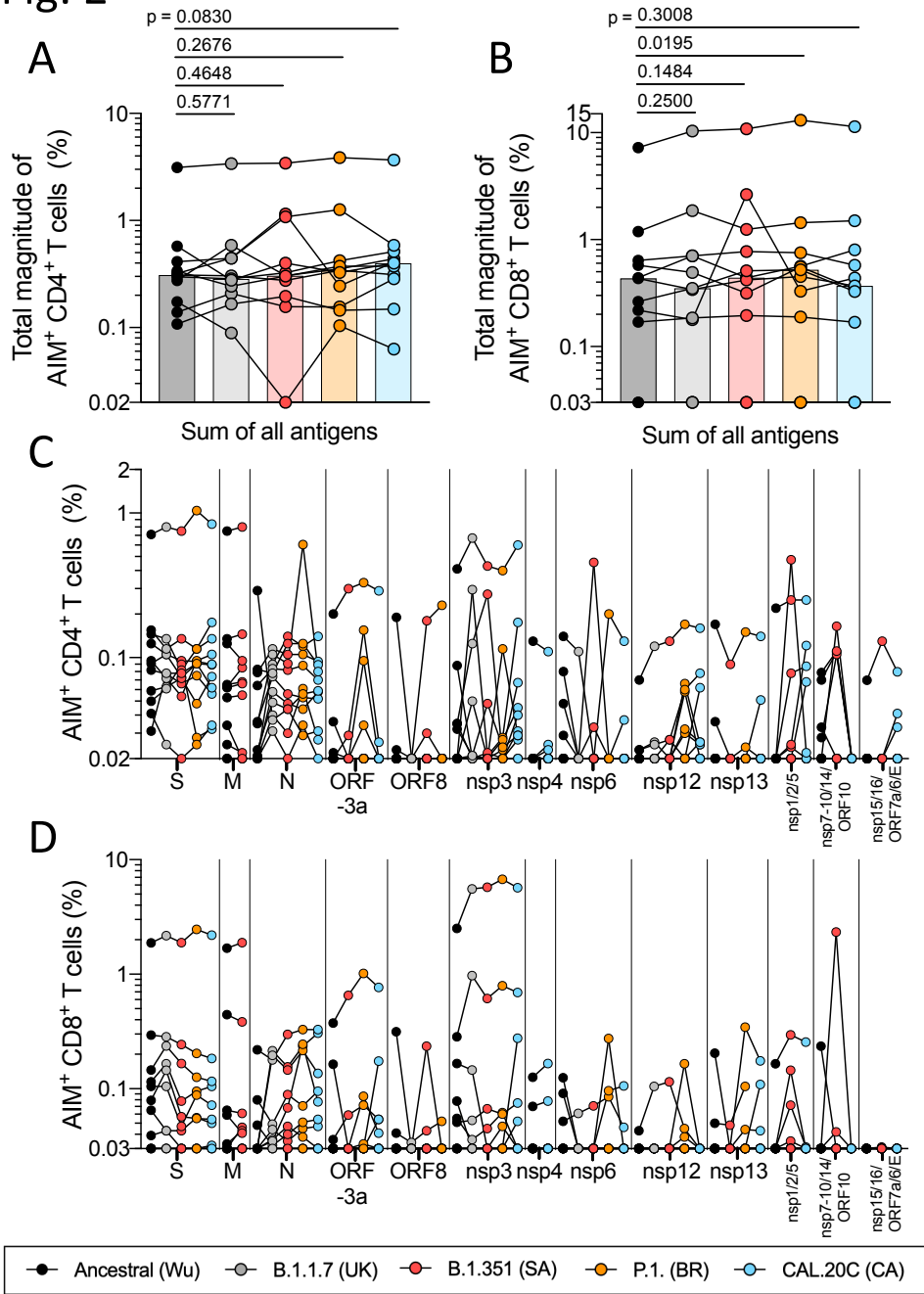


Fig. 2





### Fig. 3

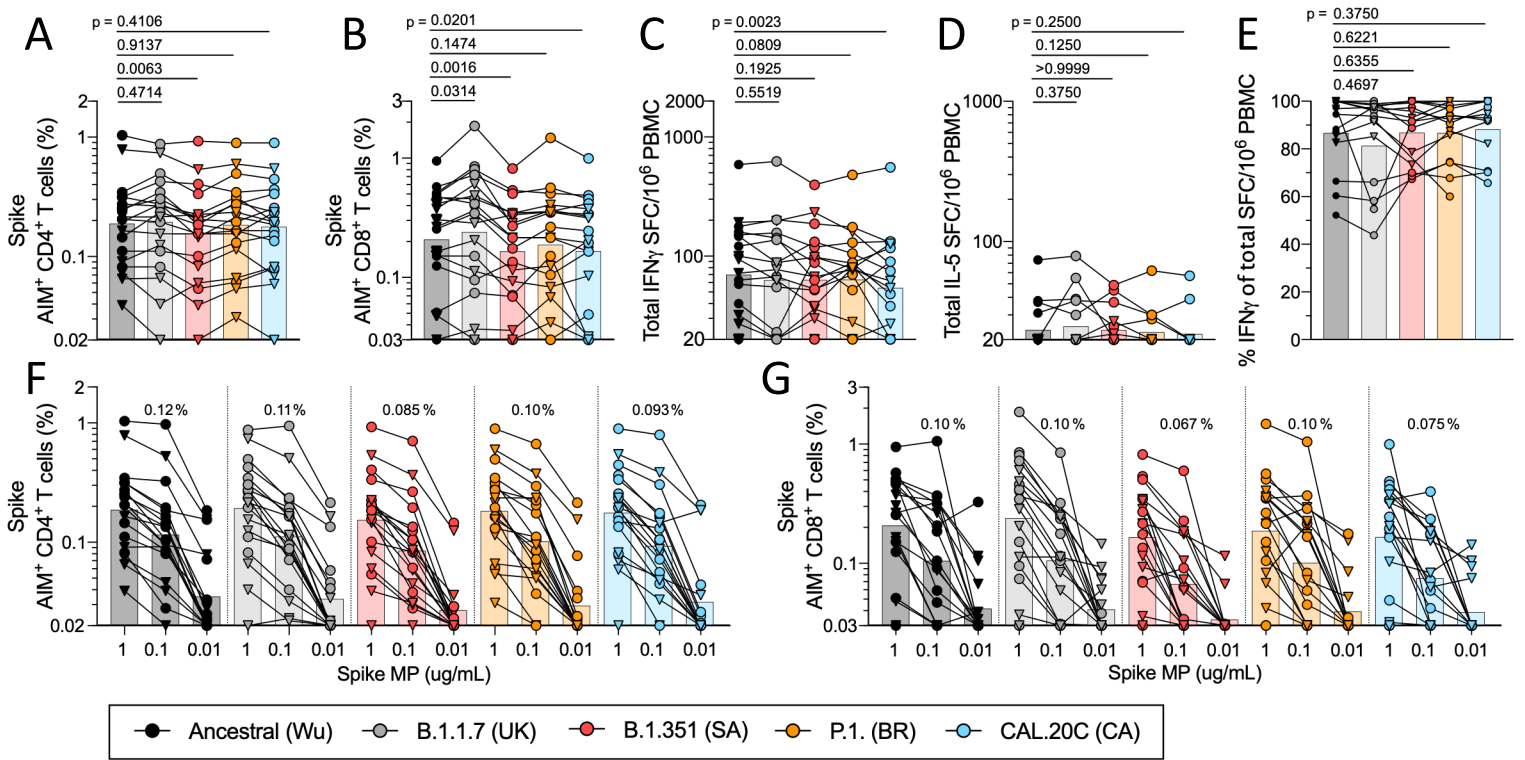


Fig. 4

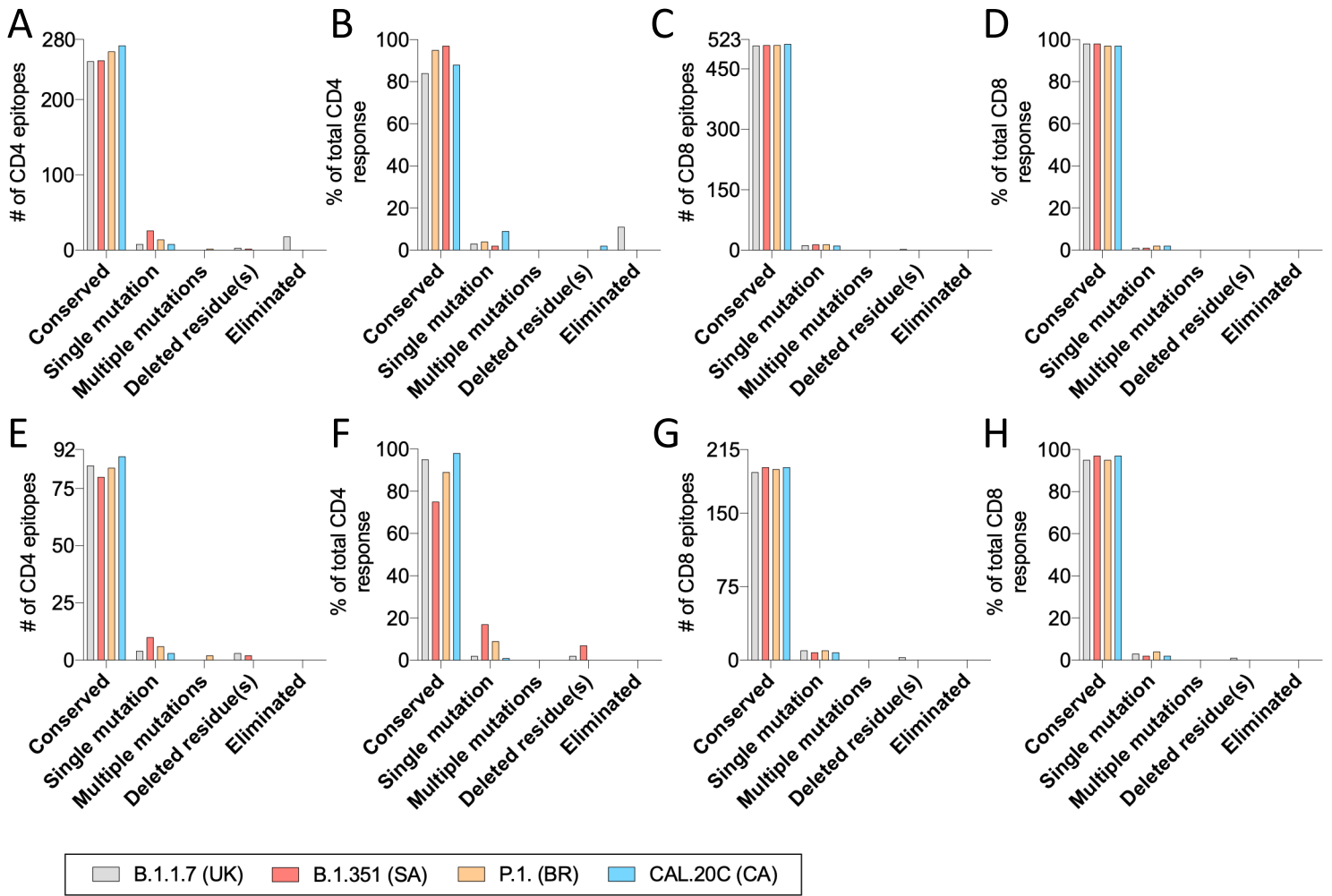


Fig. S1

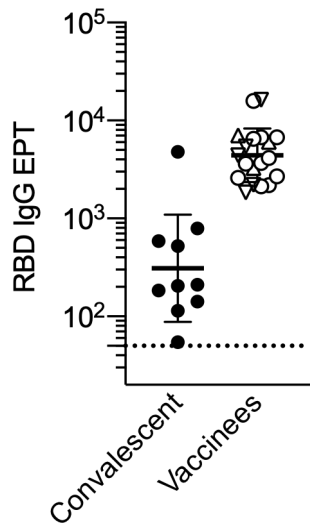


Fig. S2

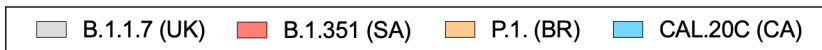
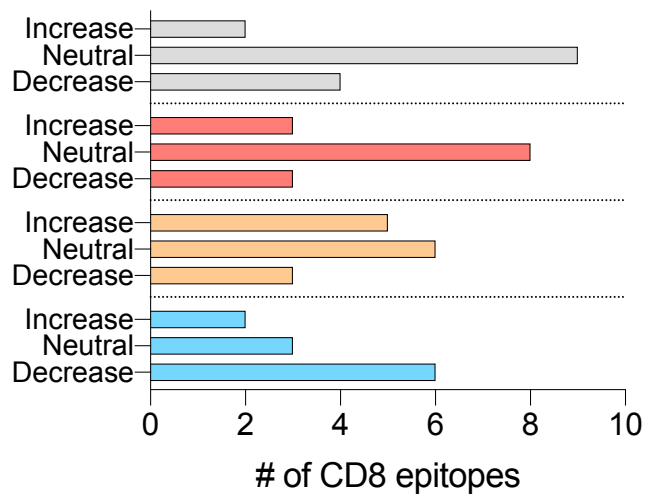
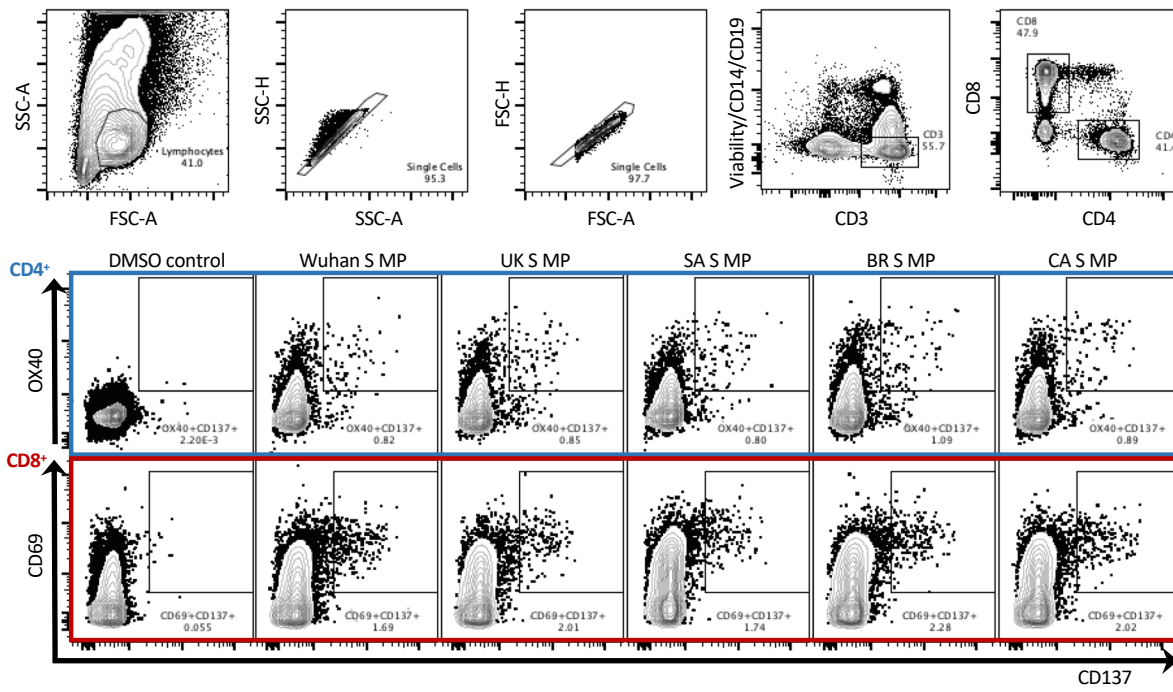


Fig. S3



**Table S1.** Related to Figures 1, 2, 3 and 4. List of amino acid positions and relative amino acid changes in the different variants studied with respect to the ancestral Wuhan strain.

Protein	Amino acid position	Ancestral (Wu)	B.1.1.7 (UK)	B.1.351 (SA)	P.1. (BR)	CAL.20C (CA)
S	13	S				I
S	18	L		F	F	
S	20	T			N	
S	26	P			S	
S	69	H	Del			
S	70	V	Del			
S	80	D		A		
S	138	D			Y	
S	145	Y	Del			
S	152	W				C
S	190	R			S	
S	215	D		G/H		
S	241	L		Del		
S	242	L		Del		
S	243	A		Del		
S	417	K		N	T	
S	452	L				R
S	484	E		K	K	
S	501	N	Y	Y	Y	
S	570	A	D			
S	614	D	G	G	G	G
S	655	H			Y	
S	681	P	H			
S	701	A		V		
S	716	T	I			
S	938	L				F
S	982	S	A			
S	1027	T			I	
S	1118	D	H			
S	1176	V			F	
S	1191	K				N
M	162	K		N		
N	3	D	L			
N	13	P		S		
N	32	R		H		
N	80	P			R	
N	203	R	K		K	K
N	204	G	R		R	R
N	205	T		I		I
N	212	G		C		
N	234	M				I
N	235	S	F			
E	71	P		L		

ORF3a	57	Q		H		H
ORF3a	131	W		L		
ORF3a	171	S		L		
ORF3a	253	S			P	
ORF7a	93	V		F		
ORF8	27	-	Stop			
ORF8	92	E			K	
ORF8	121	I		L		
nsp1	109	P		S		
nsp2	85	T		I		I
nsp2	339	G		S		
nsp2	366	S		T		
nsp2	427	Q		H		
nsp2	563	E		D		
nsp3	183	T	I			
nsp3	186	T			A	
nsp3	370	S			L	
nsp3	778	P				S
nsp3	837	K		N		
nsp3	890	A	D			
nsp3	926	C		S		
nsp3	977	K			Q	
nsp3	1180	T		I		
nsp3	1412	I	T			
nsp3	1778	N		S		
nsp4	395	S				T
nsp5	90	K		R		
nsp5	193	A		V		
nsp6	106	S	Del	Del	Del	
nsp6	107	G	Del	Del	Del	
nsp6	108	F	Del	Del	Del	
nsp6	125	L				F
nsp6	135	G		S		
nsp6	149	V		F		
nsp6	167	L				F
nsp9	65	I				V
nsp10	105	N		K		
nsp12	323	P	L	L	L	L
nsp13	53	P				L
nsp13	209	V				F
nsp13	260	D				Y
nsp13	341	E			D	
nsp13	588	T		I		
nsp14	177	L		F		
nsp14	326	F				L
nsp14	328	V		F		
nsp15	91	D				Y

Table S2. Related to figures 1, 2, 3 and 4. Lists of mutated peptides with respect to the amino acid position in the different variants studied.

Protein	Aminoacid position (Start)	Aminoacid position (End)	SARS-CoV-2 strain	Sequence
nsp3	173	187	B.1.1.7	QDGEDNQTTIIQTI
nsp3	178	192	B.1.1.7	DNQTTIIQTIVEVQP
nsp3	183	197	B.1.1.7	IIQTIVEVQPQLEME
nsp3	878	892	B.1.1.7	QDAYRRARAGEADNF
nsp3	883	897	B.1.1.7	RARAGEADNFCALIL
nsp3	888	902	B.1.1.7	EADNFCALILAYCNK
nsp3	1398	1412	B.1.1.7	NYLKSFPNFKSLINIT
nsp3	1403	1417	B.1.1.7	PNFSKLINITWIFLL
nsp3	1408	1422	B.1.1.7	LINITIWFLLSVCL
nsp6	92	106	B.1.1.7	MRIMTWLDMVDTSLK
nsp6	97	111	B.1.1.7	WLDMDVTSLLKDCV
nsp6	102	116	B.1.1.7	DTSLKDKDCVMYASA
nsp6	107	121	B.1.1.7	KLKDCVMYASAVLL
nsp12	309	323	B.1.1.7	HCANFNVLFTVFPFL
nsp12	314	328	B.1.1.7	NVLFSTVFPLTSFGP
nsp12	319	333	B.1.1.7	TVFPPLTSFGPLVRKI
N	1	15	B.1.1.7	MSLNGPQNQRNAPRI
N	191	205	B.1.1.7	RNSSRNPSTGSSKRT
N	196	210	B.1.1.7	NSTPGSSKRTSPARM
N	201	215	B.1.1.7	SSKRTSPARMAGNGG
N	221	235	B.1.1.7	LLLLDRNLQLESKMF
N	226	240	B.1.1.7	RLNQLLESKMFKGQQ
N	231	245	B.1.1.7	ESKMFKGQQQQGQT
ORF8	41	55	B.1.1.7	FYSKWYIRVGAIKSA
ORF8	46	60	B.1.1.7	YIRVGAIKSAPLIEL
ORF8	51	65	B.1.1.7	AIKSAPLIELCVDEA
ORF8	61	75	B.1.1.7	CVDEAGSKSPIQCID
ORF8	66	80	B.1.1.7	GSKSPIQCIDIGNYT
ORF8	71	85	B.1.1.7	IQCIDIGNYTVSCLP
S	56	70	B.1.1.7	LPFFSNVTWFHAISG
S	61	75	B.1.1.7	NVTWFHAISGTNGTK
S	66	80	B.1.1.7	HAISGTNGTKRFDNP
S	131	145	B.1.1.7	CEFQFCNDPFLGVYH
S	136	150	B.1.1.7	CNDPFLGVYHKNNKS
S	141	155	B.1.1.7	LGVYHKNNKSWMESE
S	491	505	B.1.1.7	PLQSYGFQPTYGVGY
S	496	510	B.1.1.7	GFQPTYGVGYQPYRV
S	501	515	B.1.1.7	YGVGYQPYRVVLSF
S	556	570	B.1.1.7	NKKFLPFQGFGRDID
S	561	575	B.1.1.7	PFQGFGRDIDDTDA
S	566	580	B.1.1.7	GRDIDDTDAVRDPQ
S	601	615	B.1.1.7	GTNTSNQVAVLYQGV
S	606	620	B.1.1.7	NQVAVLYQGVNCTEV
S	611	625	B.1.1.7	LYQGVNCTEVPVAIH
S	671	685	B.1.1.7	CASYQTQTNSHRRAR
S	676	690	B.1.1.7	TQTNSHRRARVASQ
S	681	695	B.1.1.7	HRRARSVASQSIIAY
S	706	720	B.1.1.7	AYSNNSIAIPINFTI
S	711	725	B.1.1.7	SIaipINFTISVTTTE
S	716	730	B.1.1.7	INFTISVTTTEILPVS
S	971	985	B.1.1.7	GAISSVLDILARLD
S	976	990	B.1.1.7	VLNDILARLDKVEAE
S	981	995	B.1.1.7	LARLDKVEAEVQIDR
S	1106	1120	B.1.1.7	QRNFYEPQIITHTNT
S	1111	1125	B.1.1.7	EPQIITHTNTFVSGN
S	1116	1130	B.1.1.7	TTHNTFVSGNCDVVI
N	1	15	B.1.351	MSDNGPQNQRNASRI
N	6	20	B.1.351	PQNQRNASRITFGGP
N	11	25	B.1.351	NASRITFGGSPDSTG
N	21	35	B.1.351	SDSTGSNQNGEHSGA
N	26	40	B.1.351	SNQNGEHSGARSKQR
N	31	45	B.1.351	EHSGARSKQRRPQGL
N	191	205	B.1.351	RNSSRNPSTGSSRGI
N	196	210	B.1.351	NSTPGSSRGISPARM
N	201	215	B.1.351	SSRGISPARMACNGG
N	206	220	B.1.351	SPARMACNGGDAALA
N	211	225	B.1.351	ACNGGDAALALLLLD
ORF8	107	121	B.1.351	DFLEYHDVVRVLDL
ORF7a	81	95	B.1.351	SVSPKLFIRQEEFQE
ORF7a	86	100	B.1.351	LFIRQEEFQELYSPI
ORF7a	91	105	B.1.351	EEFQELYSPIFLIVA
M	151	165	B.1.351	IAGHHLGRCDINDLP
M	156	170	B.1.351	LGRCDINDLPKEITV
M	161	175	B.1.351	INDLPKEITVATSR
ORF3a	46	60	B.1.351	LIVGALLAVFHSAS
ORF3a	51	65	B.1.351	ALLAVFHSASKIITL
ORF3a	56	70	B.1.351	FHSASKIITLKKRWQ
ORF3a	121	135	B.1.351	VRIIMRLWLCLKCRS
ORF3a	126	140	B.1.351	RLWLCLKCRSKNPLL
ORF3a	131	145	B.1.351	LKCRSKNPLLDANY
ORF3a	161	175	B.1.351	NSVTSSIVITLGDGT
ORF3a	166	180	B.1.351	SIVITLGDGTTSPIS
ORF3a	171	185	B.1.351	LGDGTTSPISEHDYQ
nsp1	96	110	B.1.351	QYGRSGETLGLVLSH
nsp1	101	115	B.1.351	GETLGLVLSHVGEIP
nsp1	106	120	B.1.351	VLVSHVGEIPVAYRK
nsp5	183	197	B.1.351	GPFVDRQTAQVAGTD
nsp5	188	202	B.1.351	RQTAQVAGTDTTITV
nsp5	193	207	B.1.351	VAGTDTTITVNLAW
nsp10	93	107	B.1.351	KGKYVQIPTTCAKDP
nsp10	98	112	B.1.351	QIPTTCAKDPVGFLL
nsp10	103	117	B.1.351	CAKDPVGFLLKNTVC
nsp12	309	323	B.1.351	HCANFNVLFTVFPFL
nsp12	314	328	B.1.351	NVLFSTVFPLTSFGP
nsp12	319	333	B.1.351	TVFPPLTSFGPLVRKI



nsp14	326	340	B.1.351	VLLPLVSSQCVNFTT
S	6	20	B.1.351	VSSQCVNFTTRTQLP
S	11	25	B.1.351	VNFTTRTQLPPAYTN
S	16	30	B.1.351	HAIHVSGTNGTKRFA
S	66	80	B.1.351	SGTNGTKRFANPVL
S	71	85	B.1.351	TKRFANPVLPFNDGV
S	76	90	B.1.351	FKIYSKHTPINLVRH
S	201	215	B.1.351	FKIYSKHTPINLVRG
S	201	215	B.1.351	KHTPINLVRHLPQGF
S	206	220	B.1.351	KHTPINLVRGLPQGF
S	206	220	B.1.351	NLVRHLPQGFSALEP
S	211	225	B.1.351	NLVRGLPQGFSALEP
E	61	75	B.1.351	RVKNLNSSRVLDLLV
nsp2	71	85	B.1.351	LQTPFEIKLAKKFDI
nsp2	76	90	B.1.351	EIKLAKKFDIFNGEC
nsp2	81	95	B.1.351	KKFDIFNGECPNFVF
nsp2	326	340	B.1.351	CGNFKVTKGKAKKSA
nsp2	331	345	B.1.351	VTKGAKKSAWNIGE
nsp2	336	350	B.1.351	AKKSAWNIGEQKSIL
nsp2	356	370	B.1.351	FASEAARVVRTIFSR
nsp2	361	375	B.1.351	ARVVRTIFSRLETA
nsp2	366	380	B.1.351	TIFSRLETAQNSVR
nsp2	416	430	B.1.351	VVMAYITGGVVHLTS
nsp2	421	435	B.1.351	ITGGVVHLTSQWLTN
nsp2	426	440	B.1.351	VHLTSQWLTNIFGTV
nsp2	551	565	B.1.351	MPLKAPKEIIFLDGE
nsp2	556	570	B.1.351	PKEIIFLDGETLPTE
nsp2	561	575	B.1.351	FLDGETLPTEVLTEE
nsp3	823	837	B.1.351	SFLGRYMSALNHTKN
nsp3	828	842	B.1.351	YMSALNHTKNWKYPQ
nsp3	833	847	B.1.351	NHTKNWKYPQVNGLT
nsp3	1168	1182	B.1.351	LHKPIVWHVNNAINK
nsp3	1173	1187	B.1.351	VWHVNNAINKATYKP
nsp3	1178	1192	B.1.351	NAINKATYKPNTWCI
nsp3	1768	1782	B.1.351	VAVKMFDAYVSTFSS
nsp3	1773	1787	B.1.351	FDAYVSTFSSFNVP
nsp3	1778	1792	B.1.351	STFSSFNVPMEKLLK
nsp13	577	591	B.1.351	SDRDLYDKLQFISLE
nsp13	582	596	B.1.351	YDKLQFISLEIPRRN
nsp13	587	601	B.1.351	FISLEIPRRNVATLQ
nsp6	92	106	B.1.351	MRIMTWLDMVDTSLK
nsp6	97	111	B.1.351	WLDVMVDTSLKLDKCV
nsp6	102	116	B.1.351	DTSLKLDKCVMYASA
nsp6	107	121	B.1.351	LKDCVMYASAVVLLI
nsp6	122	136	B.1.351	LLILMTARTVYDDSA
nsp6	127	141	B.1.351	TARTVYDDSARRVWT
nsp6	132	146	B.1.351	YDDSARRVWTLMNVL
nsp6	137	151	B.1.351	RRVWTLMNVLTLFYK
nsp6	142	156	B.1.351	LMNVLTLFYKVYYGN
nsp6	147	161	B.1.351	TLFYKVYYGNALDQA
S	231	245	B.1.351	IGINITRFQTLHRSY
S	236	250	B.1.351	TRFQTLHRSYLPDGD
S	241	255	B.1.351	LHRSYLPDGDSSSGW
S	406	420	B.1.351	EVRQIAPGQTGNIAI
S	411	425	B.1.351	APGQTGNIAIDYNYKL
S	416	430	B.1.351	GNIADYNYKLDDFT
S	476	490	B.1.351	EIQAGSTPCNGVKG
S	481	495	B.1.351	GSTPCNGVKGFNCYF
S	486	500	B.1.351	NGVKGFNCYFPLQSY
S	491	505	B.1.351	PLQSYGFQPTYGVGY
S	496	510	B.1.351	GFQPTYGVGYQPYRV
S	501	515	B.1.351	YGVGYQPYRVVLSF
S	606	620	B.1.351	GTNTSNQVAVLYQGV
S	611	625	B.1.351	NQVAVLYQGVNCTEV
S	616	630	B.1.351	LYQGVNCTEVPVAIH
S	691	705	B.1.351	SHIAYTMSLGVENSV
S	696	710	B.1.351	TMSLGVENSVAYSNN
S	701	715	B.1.351	VENSVAYSNNIAIP
N	66	80	P.1.	FPRGGQVPINTNSSR
N	71	85	P.1.	GVPINTNSSRDDQIG
N	76	90	P.1.	TNSSRDDQIGYYRRA
N	191	205	P.1.	RNSSRNSTPGSSKRT
N	196	210	P.1.	NSTPGSSKRTSPARM
N	201	215	P.1.	SSKRTSPARMAGNGG
ORF8	81	95	P.1.	VSLPFTINCQKPKL
ORF8	86	100	P.1.	FTINCQKPKLGLSLV
ORF8	91	105	P.1.	QKPKLGLSLVVRCSFY
ORF3a	241	255	P.1.	EEHVQIHTIDGSPGV
ORF3a	246	260	P.1.	IHTIDGSPGVVNPVM
ORF3a	251	265	P.1.	GSPGVVNPVMEPIYD
nsp12	309	323	P.1.	HCANFNVLFSVFPL
nsp12	314	328	P.1.	NVLFSTVFPLTSFGP
nsp12	319	333	P.1.	TVFPLTSFGPLVRKI
nsp13	327	341	P.1.	IDKCSRIIPARARVD
nsp13	332	346	P.1.	RIIPARARVDCFDKF
nsp13	337	351	P.1.	RARVDCFDKFKVNST
S	6	20	P.1.	VLLPLVSSQCVNFTN
S	11	25	P.1.	VSSQCVNFTNRTQLP
S	16	30	P.1.	VNFTNRTQLPSAYTN
S	21	35	P.1.	RTQLPSAYTNSFTRG
S	26	40	P.1.	SAYTNSFTRGVYYPD
S	126	140	P.1.	VVIKVEFCQFCNYPF
S	131	145	P.1.	CEFCQFCNYPFLGVYY
S	136	150	P.1.	CNYPFLGVYYHKNNK
S	176	190	P.1.	LMDLEGKQGNFKNLS
S	181	195	P.1.	GKQGNFKNLSEFVK
S	186	200	P.1.	FKNLSEFVKNIDGY
S	406	420	P.1.	EVRQIAPGQTGTIAD

S	471	485	P.1.	AGGQETADVYKPL
S	476	490	P.1.	GSTPCNGVKGFNCYF
S	481	495	P.1.	NGVKGFNCYFPLQSY
S	491	505	P.1.	PLQSYGFQPTYGVGY
S	496	510	P.1.	GFQPTYGVGYQPYRV
S	501	515	P.1.	YGVGYQPYRVVLSF
S	601	615	P.1.	GTNTSNQVAVLYQGV
S	606	620	P.1.	NQVAVLYQGVNCTEV
S	611	625	P.1.	LYQGVNCTEVPVAIH
S	641	655	P.1.	NVFQTRAGCLIGAEY
S	646	660	P.1.	RAGCLIGAEYVNNNSY
S	651	665	P.1.	IGAEYVNNSYECDIP
S	1016	1030	P.1.	AEIRASANLAAIKMS
S	1021	1035	P.1.	SANLAAIKMSECVLG
S	1026	1040	P.1.	AIKMSECVLGQSKRV
S	1166	1180	P.1.	LGDISGINASFVNIQ
S	1171	1185	P.1.	GINASFVNIQKEIDR
S	1176	1190	P.1.	FVNIQKEIDRLNEVA
nsp3	173	187	P.1.	QDGSEDNQTTTIQAI
nsp3	178	192	P.1.	DNQTTTIQAIIVEVQP
nsp3	183	197	P.1.	TIQAIIVEVQPQLEME
nsp3	358	372	P.1.	AVFDKNLYDKLVLSF
nsp3	363	377	P.1.	NLYDKLVLSFLEMKS
nsp3	368	382	P.1.	LVLFSLEMKSEKQVE
nsp3	963	977	P.1.	KGVQIPCTCGKQATQ
nsp3	968	982	P.1.	PCTCGKQATQYLQQ
nsp3	973	987	P.1.	KQATQYLQQESPFV
nsp6	92	106	P.1.	MRIMTWLDMVDTSLK
nsp6	97	111	P.1.	WLDVMVDTSLKDKCV
nsp6	102	116	P.1.	DTSLKDKCVMYASA
nsp6	107	121	P.1.	LKDCVMYASAVVLLI
N	191	205	CAL.20C	RNSSRNSTPGSSKRI
N	196	210	CAL.20C	NSTPGSSKRISPARM
N	201	215	CAL.20C	SSKRISPARMAGNGG
N	221	235	CAL.20C	LLLLDRLNQLESKIS
N	226	240	CAL.20C	RLNQLESKISGKQQ
N	231	245	CAL.20C	ESKISGKQQQQGQT
nsp2	71	85	CAL.20C	LQTPFEIKLAKKFDI
nsp2	76	90	CAL.20C	EIKLAKKFDIFNGEC
nsp2	81	95	CAL.20C	KKFDIFNGECPNFV
nsp3	768	782	CAL.20C	MSMTYGGQFGSTYLD
nsp3	773	787	CAL.20C	GQQFGSTYLDGADVT
nsp3	778	792	CAL.20C	STYLDGADVTKIKPH
nsp3	383	397	CAL.20C	ICISTKHFYWFFTN
nsp4	388	402	CAL.20C	KHFYWFFTNYLKRRV
nsp4	393	407	CAL.20C	FFTNYLKRRVVFNGV
nsp6	112	126	CAL.20C	DCVMYASAVVLLIFM
nsp6	117	131	CAL.20C	ASAVVLLIFMTARTV
nsp6	122	136	CAL.20C	LLIFMTARTVYDDGA
nsp6	157	171	CAL.20C	ALDQAISMWAFIISV
nsp6	162	176	CAL.20C	ISMWAFIISVTSNYS
nsp6	167	181	CAL.20C	FIIISVTSNYSVVT
nsp9	51	65	CAL.20C	LKWARFPKSDGTGT
nsp9	56	70	CAL.20C	FPKSDGTGTVYTELE
nsp9	61	75	CAL.20C	GTGTVYTELEPPCRF
nsp12	309	323	CAL.20C	HCANFNVLFTVFP
nsp12	314	328	CAL.20C	NVLFSTVFPVPLTSFGP
nsp12	319	333	CAL.20C	TVFPVPLTSFGPLVRKI
nsp13	42	56	CAL.20C	VLSVNPVVCNAPGCD
nsp13	47	61	CAL.20C	PYVCNAPGCDVTDVT
nsp13	52	66	CAL.20C	APGCDVTDVTLQYLG
nsp13	197	211	CAL.20C	EYTFEKGDYDAFVY
nsp13	202	216	CAL.20C	KGDYDAFVYRGTTT
nsp13	207	221	CAL.20C	DAFVYRGTTTYKLN
nsp13	247	261	CAL.20C	VRITGLYPTLNISYE
nsp13	252	266	CAL.20C	LYPTLNISYEFSSNV
nsp13	257	271	CAL.20C	NISYEFSSNVANYQK
nsp14	316	330	CAL.20C	VVKAALLADKLPVLH
nsp14	321	335	CAL.20C	LLADKLPVLHDIGNP
nsp14	326	340	CAL.20C	LPVLHDIGNPKAIKC
nsp15	79	93	CAL.20C	IAANTVIWDYKRYAP
nsp15	84	98	CAL.20C	VIWDYKRYAPAHIST
nsp15	89	103	CAL.20C	KRYAPAHISTIGVCS
ORF3a	46	60	CAL.20C	LIVGVALAVFHSAS
ORF3a	51	65	CAL.20C	ALLAVFHSASKIITL
ORF3a	56	70	CAL.20C	FHSASKIITLKKRWQ
S	1	15	CAL.20C	MFVFLVLLPLVSIQC
S	6	20	CAL.20C	VLLPLVSIQCVNLT
S	11	25	CAL.20C	VSIQCVNLTTRTQLP
S	141	155	CAL.20C	LGVYHKNKSCMES
S	146	160	CAL.20C	HKNKSCMESEFRVY
S	151	165	CAL.20C	SCMESEFRVYSSANN
S	441	455	CAL.20C	LDSKVGGNRYRRL
S	446	460	CAL.20C	GGNRYRRLFRKSN
S	451	465	CAL.20C	YRRLFRKSNLKPFE
S	601	615	CAL.20C	GTNTSNQVAVLYQGV
S	606	620	CAL.20C	NQVAVLYQGVNCTEV
S	611	625	CAL.20C	LYQGVNCTEVPVAIH
S	926	940	CAL.20C	QFNSAIGKIQDSFSS
S	931	945	CAL.20C	IGKIQDSFSSSTASAL
S	936	950	CAL.20C	DSFSSSTASALGKLQD
S	1181	1195	CAL.20C	KEIDRLNEVANLNE
S	1186	1200	CAL.20C	LNEVANLNEVSLIDL
S	1191	1205	CAL.20C	NNLNEVSLIDLQELGK

**Table S3.** Related to Figure 4. Effect of mutations on CD8 epitope HLA class I binding capacity

Origin	Ancestral reference sequence	Protein	Start	Mutation <sup>a</sup>	Mutated sequence <sup>b</sup>	HLA restriction	WT (IC <sub>50</sub> nM) <sup>c</sup>	Mutant (IC <sub>50</sub> nM)	Fold difference	Effect <sup>d</sup>
B.1.1.7	HVSGTNGTK	S	69	HV69-70 del	HAISGTNGTK	A*68:01	55	44	0.8	Neutral
B.1.1.7	GVYHKNNK	S	142	Y145 del	FLGVYHKNNK	A*03:01	28	1078	39	Decrease
B.1.1.7	YYHKNNKSW	S	144	Y145 del	VYHKNNKSW	A*24:02	117	308	2.6	Decrease
B.1.1.7	YGFQPTNGV	S	495	N501Y	YGFQPTYGV	B*51:01	3488	3541	1.0	Neutral
B.1.1.7	YQDVNCTEV	S	612	D614G	YQGVNCTEV	A*02:06	18	57	3.2	Decrease
B.1.1.7	QTNSPRRAR	S	677	P681H	QTNSHRRAR	A*31:01	35	33	0.94	Neutral
B.1.1.7	SPRRARSV	S	680	P681H	SHRRARSV	B*08:01	429	2449	5.7	Decrease
B.1.1.7	NSAIPTNF	S	710	T716I	NSAIPINF	B*57:01	1335	968	0.73	Neutral
B.1.1.7	IAIPTNFTI	S	712	T716I	IAIPINFTI	B*51:01	209	189	0.90	Neutral
B.1.1.7	IAIPTNFTI	S	712	T716I	IAIPINFTI	B*53:01	396	266	0.67	Neutral
B.1.1.7	IPTNFTISV	S	714	T716I	IPINFTISV	B*07:02	188	168	0.89	Neutral
B.1.1.7	IPTNFTISV	S	714	T716I	IPINFTISV	B*51:01	156	94	0.60	Neutral
B.1.1.7	SVLNDILSR	S	975	S982A	SVLNDILAR	A*68:01	109	92	0.84	Neutral
B.1.1.7	KLINIIWIF	nsp3	1407	I1412T	KLINITIWF	A*32:01	161	48	0.30	Increase
B.1.1.7	STVFPPTSF	nsp12	318	P323L	STVFPPTSF	B*57:01	1583	637	0.40	Increase
B.1.351	RFDNPVLPF	S	78	D80A	RFANPVLFPF	A*24:02	458	34	0.075	Increase
B.1.351	FDNPVLPFDGVYF	S	79	D80A	FANPVLFPFDGVYF	B*35:01	65	65	1.0	Neutral
B.1.351	TPINLVRDL	S	208	D215G	TPINLVRGL	B*07:02	213	119	0.56	Neutral
B.1.351	TPINLVRDL	S	208	D215H	TPINLVRHL	B*07:02	213	199	0.93	Neutral
B.1.351	QIAPGQTGK	S	409	K417N	QIAPGQTGN	A*68:01	137	27998	204	Decrease
B.1.351	YGFQPTNGV	S	495	N501Y	YGFQPTYGV	B*51:01	3488	3541	1.0	Neutral
B.1.351	YQDVNCTEV	S	612	D614G	YQGVNCTEV	A*02:06	18	57	3.2	Decrease
B.1.351	YTMSLGAENSVAY	S	695	A701V	YTMSLGVENSVAY	A*26:01	184	253	1.4	Neutral
B.1.351	LGAENSVAY	S	699	A701V	LGVENSVAY	B*35:01	19	21	1.1	Neutral
B.1.351	GPQNQRNAPRITF	N	5	K17N	GPQNQRNASRITF	B*07:02	640	696	1.1	Neutral
B.1.351	QSASKIITL	ORF3a	57	Q57H	HSASKIITL	B*08:01	1788	573	0.32	Increase
B.1.351	MSALNHTKK	nsp3	829	K837N	MSALNHTKN	A*30:01	102	7035	69	Decrease
B.1.351	MSALNHTKKW	nsp3	829	K837N	MSALNHTKNW	B*57:01	16	14	0.88	Neutral
B.1.351	SALNHTKKW	nsp3	830	K837N	SALNHTKNW	B*57:01	111	93	0.84	Neutral
B.1.351	STVFPPTSF	nsp12	318	P323L	STVFPPTSF	B*57:01	1583	637	0.40	Increase
P.1.	LPPAYTNSF	S	24	P26S	LPSAYTNSF	B*07:02	294	51	0.17	Increase
P.1.	LPPAYTNSF	S	24	P26S	LPSAYTNSF	B*35:01	44	4.1	0.093	Increase
P.1.	LPPAYTNSF	S	24	P26S	LPSAYTNSF	B*53:01	366	18	0.049	Increase
P.1.	QIAPGQTGK	S	409	K417T	QIAPGQTGT	A*68:01	137	20478	149	Decrease
P.1.	YGFQPTNGV	S	495	N501Y	YGFQPTYGV	B*51:01	3488	3541	1.0	Neutral
P.1.	YQDVNCTEV	S	612	D614G	YQGVNCTEV	A*02:06	18	57	3.2	Decrease
P.1.	AEHVNNYSY	S	653	H655Y	AEYVNNYSY	B*44:02	1038	904	0.87	Neutral
P.1.	AEHVNNYSY	S	653	H655Y	AEYVNNYSY	B*44:03	1020	577	0.57	Neutral
P.1.	RASANLAATK	S	1019	L1027I	RASANLAAIK	A*03:01	85	99	1.2	Neutral
P.1.	NASVVNIQK	S	1173	V1176F	NASFVNIQK	A*68:01	13	6.2	0.48	Increase
P.1.	NTNSSPDDQIGYY	N	75	P80R	NTNSSRDDQIGYY	A*01:01	44	44	1.0	Neutral
P.1.	SPDDQIGYY	N	79	P80R	SRDDQIGYY	B*35:01	101	17781	175	Decrease
P.1.	LYDKLVSSF	nsp3	364	S370L	LYDKLVLSF	A*24:02	77	70	0.91	Neutral
P.1.	STVFPPTSF	nsp12	318	P323L	STVFPPTSF	B*57:01	1583	637	0.40	Increase
CAL.20C	LPLVSSQCV	S	8	S13I	LPLVSIQCV	B*51:01	402	272	0.68	Neutral
CAL.20C	YYHKNNKSW	S	144	W152C	YYHKNNKSC	A*24:02	117	11134	95	Decrease
CAL.20C	SWMESEFRVY	S	151	W152C	SCMESEFRVY	A*29:02	49	980	20	Decrease
CAL.20C	KVGGNYYNYLY	S	444	L452R	KVGGNYYNYRY	A*29:02	101	505	5.0	Decrease
CAL.20C	VGGNYYNYLY	S	445	L452R	VGGNYYNYRY	A*29:02	94	519	5.5	Decrease
CAL.20C	NYYNYRRLF	S	448	L452R	NYYNYRRLF	A*24:02	21	108	5.1	Decrease
CAL.20C	YNYLYRLEFR	S	449	L452R	YNYRYRLEFR	A*31:01	16	12	0.75	Neutral
CAL.20C	YQDVNCTEV	S	612	D614G	YQGVNCTEV	A*02:06	18	57	3.2	Decrease
CAL.20C	QSASKIITL	ORF3a	57	Q57H	HSASKIITL	B*08:01	1788	573	0.32	Increase
CAL.20C	WFFSNYLKR	nsp4	392	S395T	WFFTNYLKR	A*31:01	70	98	1.4	Neutral
CAL.20C	STVFPPTSF	nsp12	318	P323L	STVFPPTSF	B*57:01	1583	637	0.40	Increase

<sup>a</sup>Mutation noted as ancestral residue-position-variant residue. Del refers to deletion of the corresponding residue.

<sup>b</sup>For deletion mutants, the peptide sequence shown represents the variant encompassing the same region that has the highest predicted binding affinity for the corresponding restricting allele.

<sup>c</sup>Indicates predicted IC<sub>50</sub> for the corresponding reported restricting allele. Predictions were performed using the NetMHCpan BA 4.1 algorithm, hosted by the IEDB.

<sup>d</sup>Increase/decrease in affinity defined by a two-fold difference in predicted IC<sub>50</sub> nM.