



The biological and clinical significance of emerging SARS-CoV-2 variants

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Abstract | The past several months have witnessed the emergence of SARS-CoV-2 variants with novel spike protein mutations that are influencing the epidemiological and clinical aspects of the COVID-19 pandemic. These variants can increase rates of virus transmission and/or increase the risk of reinfection and reduce the protection afforded by neutralizing monoclonal antibodies and vaccination. These variants can therefore enable SARS-CoV-2 to continue its spread in the face of rising population immunity while maintaining or increasing its replication fitness. The identification of four rapidly expanding virus lineages since December 2020, designated variants of concern, has ushered in a new stage of the pandemic. The four variants of concern, the Alpha variant (originally identified in the UK), the Beta variant (originally identified in South Africa), the Gamma variant (originally identified in Brazil) and the Delta variant (originally identified in India), share several mutations with one another as well as with an increasing number of other recently identified SARS-CoV-2 variants. Collectively, these SARS-CoV-2 variants complicate the COVID-19 research agenda and necessitate additional avenues of laboratory, epidemiological and clinical research.

Humoral immunity

Immunity mediated via host antibodies including those that directly neutralize virus as well as those that recruit other host immune functions.

Viral mutation rate

The rate of mutation calculated in vitro as the number of nucleotide incorporation errors per round of replication or in vivo as the number of nucleotide changes observed during a fixed time such as 1 year. Although the two rates are related, the in vivo rate is also influenced by the number of replication cycles that occur over time and the frequency with which multiple mutations occur at the same position.

Among the many unprecedented aspects of the SARS-CoV-2 pandemic is the intense virological monitoring that has occurred, with more than two million virus isolates having undergone partial or complete genomic sequencing. Initially, genetic sequencing suggested that SARS-CoV-2 was exceptionally well adapted to humans, spreading rapidly with little evidence for natural selection among circulating viruses. This changed during the later months of 2020, with the first reports of emergent SARS-CoV-2 variants associated with increased transmissibility, disease severity and escape from humoral immunity.

In this Review, we create a framework for understanding SARS-CoV-2 variants by describing fundamental aspects of SARS-CoV-2 evolution, the structure and function of the SARS-CoV-2 spike protein and the laboratory methods used to characterize spike variants. We then describe the biological properties and epidemiological characteristics of these variants and their associated mutations. Lastly, we describe the types of study required for the research, clinical and public health communities to respond to the new threat posed by emerging SARS-CoV-2 variants. Given the wide public interest in this topic, we provide a box of key points. We also provide a repository of the SARS-CoV-2 variant neutralization data discussed in this Review ([Stanford University Coronavirus Antiviral & Resistance Database — Susceptibility Data](#)).

SARS-CoV-2 evolution

Coronaviruses contain an exonuclease enzyme that reduces their replication error rate by about 15-fold to 20-fold in vitro, resulting in an in vivo viral mutation rate about 10-fold lower than that of influenza^{1–3}. Nonetheless, they accumulate mutations and generate further diversity through the process of recombination when variants with different mutations infect the same host^{4–6}. Recombination between different SARS-related coronaviruses is likely to have led to the emergence of SARS-CoV-2 (REF.⁷) and, although it can be difficult to detect owing to the similarity of most sequences, recombination is occurring to some extent among circulating SARS-CoV-2 variants^{6,8}. Additionally, host-mediated RNA editing by APOBEC and ADAR enzymes, as evidenced by the dominance of C to U changes in specific dinucleotide contexts, contributes to SARS-CoV-2 diversity^{9,10}.

Although it had been previously assumed that waning immunity explained the observation that people are commonly reinfected with endemic common-cold coronaviruses¹¹, recent studies suggest that antigenic drift also contributes to the lack of long-lasting protection following coronavirus infections^{12,13}. HCoV-229E and HCoV-OC43 sequences over a 30-year period demonstrate a ladder-like phylogenetic tree topology consistent with the emergence of novel variants sweeping

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Key points

- The past several months have witnessed the emergence of four SARS-CoV-2 variants of concern (Alpha, Beta, Gamma and Delta) associated with increased transmissibility, increased risk of reinfection and/or reduced vaccine efficacy.
- Many additional SARS-CoV-2 variants sharing mutations and biological features with these variants are also increasingly being identified.
- The increasing number of SARS-CoV-2 variants share a repertoire of mutations that is enabling the virus to spread despite rising population immunity while maintaining or increasing its replication fitness.
- Whereas most emerging mutations reduce the protective effects of neutralizing antibodies generated by infection and vaccination, several recently identified mutations appear to antagonize the innate immune response to initial infection.
- The emergence of SARS-CoV-2 variants requires an expanded research agenda to improve our understanding of emerging SARS-CoV-2 mutations and the correlates of protective immunity against variants with these mutations.

APOBEC and ADAR enzymes

Apolipoprotein B mRNA-editing enzyme catalytic polypeptide-like (APOBEC) and adenosine deaminase acting on RNA (ADAR) are host enzymes that edit viral genomes. Although these enzymes represent an antiviral defence mechanism, it is often hypothesized that these enzymes can contribute to viral evolution.

through the human population similar to seasonal influenza, albeit at a slower rate, with virus isolates from one time point often evading neutralization by plasma from persons infected several years earlier^{12,13}.

The evolutionary rate of SARS-CoV-2 has been estimated to be between 0.0004 and 0.002 mutations per nucleotide per year^{14–19}. Although the possibility that synonymous mutations may influence SARS-CoV-2 phenotypic properties should not be discounted, there have been no reports of this phenomenon occurring within the SARS-CoV-2 spike gene. Therefore, in this Review, we use the term mutation to indicate an amino acid change from the Wuhan-Hu-1 reference sequence (GenBank accession: NC_045512.2).

The phylogenetic classification of emergent SARS-CoV-2 lineages has been difficult because new lineages often differ from one another by just a few nucleotides^{20,21}. Geographical classification has been challenging because most variants have been detected in multiple countries and there are marked disparities in the proportion of viruses undergoing sequencing in different countries. Two commonly used systems have been developed for epidemiological surveillance: the Phylogenetic Assignment of Named Global Outbreak (PANGO) lineage²² and NextStrain²³ systems. The PANGO lineage system provides greater specificity and is used more frequently. It contains an alphabetical prefix and a suffix containing up to three numbers separated by periods indicating sub-lineages (such as B.1.1.7). However, as the system allows for only three hierarchical levels, the introduction of a new lineage suffix can make it difficult to identify the ancestral lineage of a variant.

In addition, the lineage of a virus does not always correspond to its component mutations, as a virus can acquire additional biologically relevant mutations without being assigned to a new lineage.

The first indication of SARS-CoV-2 genetic evolutionary selection pressure became evident as a novel virus variant containing the spike mutation D614G emerged in early 2020 and rose to a prevalence of nearly 100% by June 2020 (REFS^{8,24–26}). By the end of 2020 and in early 2021, several variants with recurrent mutations (in addition to D614G) occurring primarily, but not exclusively, in the spike protein were also reported. In December 2020, B.1.1.7, a rapidly growing lineage in the UK associated with an unexpectedly large number of genetic changes, was reported on the virological.org discussion forum²⁷. Retrospective analyses determined that the earliest clinical sample of this variant had been obtained in the UK in late September 2020.

Within 1 month, two additional rapidly growing lineages with large numbers of genetic changes were reported from South Africa¹⁶ and Brazil¹⁹. The B.1.351 variant rose in prevalence in South Africa from 11% in October to 87% by December²⁸. The P.1 variant emerged in Manaus, Brazil, a region that was estimated to have achieved an infection rate approaching 75% by October 2020, but which experienced a surge in new cases beginning in November 2020 (REFS^{19,29,30}). Subsequently, a novel variant (B.1.617.2) increased in prevalence from 2% in February 2021 to 87% in May 2021 in Maharashtra, India, as India experienced a dramatic surge in cases³¹. Since then, the B.1.617.2 variant has spread widely in multiple countries^{32–34} and displayed evidence of being even more transmissible than the B.1.1.7 variant, and is likely to cause more severe disease than earlier virus variants^{35,36}.

Variants that have spread widely and displayed evidence for being more transmissible, causing more severe disease and/or reducing neutralization by antibodies generated during previous infection or vaccination have been classified as variants of concern (VOCs) by the World Health Organization (WHO)³⁷, US Centers for Disease Control and Prevention (CDC)³⁸ and COVID-19 Genomics UK Consortium (COG-UK)³⁹. Variants that have spread less widely but contain mutations similar to those present within VOCs have been classified as variants of interest (VOIs). On 31 May 2021, the WHO labelled VOCs and VOIs using the Greek alphabet, with the current VOCs designated as Alpha (B.1.1.7), Beta (B.1.351), Gamma (P.1) and Delta (B.1.617.2) (FIG. 1).

Spike structure and immune epitopes

The SARS-CoV-2 spike protein is a 1,273-amino acid trimeric glycoprotein responsible for virus entry into host cells (FIG. 1). Each spike monomer has a largely exposed S1 attachment domain (residues 1–686) and a partially buried S2 fusion domain (residues 687–1,273)^{40,41}. Part of S1, called the receptor-binding domain (RBD; residues 306–534), alternates between a closed/down position and an open/up position. When in the up position, it binds to the human angiotensin-converting enzyme 2 (ACE2) receptor, the necessary first step for entry into most, if not all, cells^{42–45}. Approximately 20 RBD residues form

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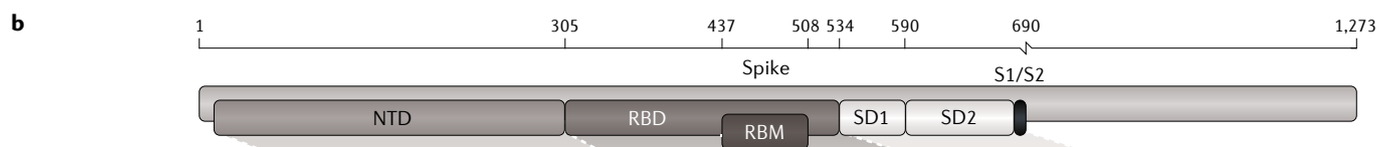
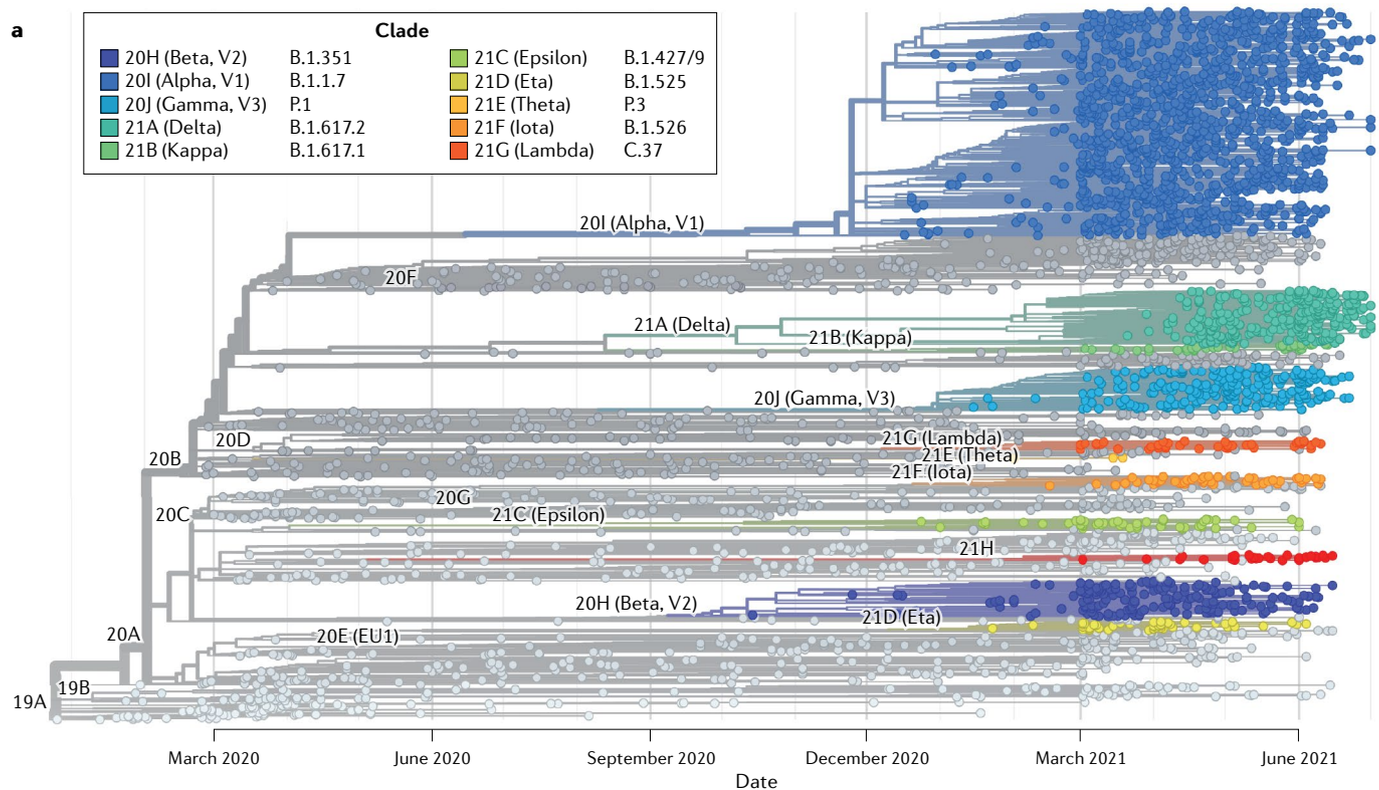
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WHO	PANGO lineage	Country of first identification	Spike														nsp6: 106	N:203/ 204
			18	69	95	144	242	417	452	478	484	490	501	614	681	701		
			L	H	T	Y	L	K	L	T	E	F	N	D	P	A	S	R/G
Alpha	B.1.1.7	UK		del		del							Y	G	H		del	K/R
Beta	B.1.351	South Africa	F				del	N				K	Y	G		V	del	
Gamma	P.1	Brazil	F					T				K	Y	G			del	K/R
Delta	B.1.617.2	India			I				R	K				G	R			M
Epsilon	B.1.427/9	USA							R					G				
Eta	B.1.525	Nigeria		del		del						K		G			del	
Theta	P.3	Philippines										K	Y	G	H			K/R
Iota	B.1.526	USA			I	del						K		G		V	del	
Kappa	B.1.617.1	India			I				R			Q		G	R			M
Zeta	P.2	Brazil										K		G				K/R
Lambda	C.37	Peru							Q				S	G			del	K/R

Fig. 1 | SARS-CoV-2 variants: evolution and constituent mutations. **a** | Phylogenetic tree based on subsampling of globally circulating sequences created by NextStrain (CC BY 4.0). The tree shows that nearly all variants of concern (VOCs; Alpha, Beta, Gamma and Delta) and variants of interest (VOIs; Kappa, Epsilon, Eta, Theta, Iota and Lambda) emerged independently beginning in late 2020. **b** | The most common mutations present in multiple VOCs and VOIs. Numeric column headers indicate spike protein positions except for two non-spike mutations in the nsp6 and nucleocapsid (N) genes. The second row indicates the residue found in the reference sequence.

Spike protein residues are mapped to their associated domain within the spike protein, as shown in various shades of grey above the table. Deletions are indicated 'del'. Several additional mutations in other viral proteins also appear to have arisen more than once, including orf3a:Q57H and nsp2:T85I. NTD, amino-terminal domain; PANGO, Phylogenetic Assignment of Named Global Outbreak; RBD, receptor-binding domain; RBM, receptor-binding motif; SD, subdomain; S1/S2, junction between the exposed S1 attachment domain and the partially buried S2 fusion domain; WHO, World Health Organization.

Neutralizing antibodies

Antibodies that alone can prevent virus infection of cells in vitro. Neutralization is determined almost entirely by an antibody fragment antigen-binding (Fab) region.

Epitope

An antigenic determinant of a protein. B lymphocyte antibody epitopes are often formed by amino acids from different parts of a protein that are brought together during protein folding. T lymphocyte epitopes are linear peptides recognized by T lymphocyte receptors when bound to a human leukocyte antigen (HLA) protein on a cell surface. As human HLA proteins are heterogenous, different people recognize different epitopes of the same protein.

contacts with the human ACE2 receptor. The part of the RBD containing these residues encompasses residues 438–506 and is called the receptor-binding motif (RBM), whereas the remainder of the RBD is called the RBD core.

Similar to the RBD, much of the S1 amino-terminal domain (NTD) is also exposed on the S trimer surface. The remainder of S1 contains two subdomains downstream of the RBD traditionally referred to as subdomain 1 (SD1) and subdomain 2 (SD2) that we refer to in this Review as the S1 carboxy-terminal domain (CTD). The spike protein also has 22 glycosylation sites, distributed among both the S1 and S2 domains. Within S1, eight of the glycans are found in the NTD, two are in the RBD core, three are in the CTD and nine are in S2 (REF. 46).

S1 displays more amino acid variability than S2 among SARS-related coronaviruses (FIG. 2). Within S1, the RBD and the NTD are more variable than the CTD. Within the RBD, the RBM is more variable than the RBD core. As of June 2021, 42 spike mutations have a global sampled prevalence $\geq 1.0\%$ including 15 in the NTD, 6 in the RBD, 5 in the CTD and 9 in S2 (Supplementary Table 1). Most of the 32 S1 mutations with a prevalence $\geq 1.0\%$ arose in multiple SARS-CoV-2 lineages.

The spike RBD is the main target of neutralizing antibodies^{47–52}. The presence of neutralizing monoclonal antibodies (mAbs) targeting the RBD correlates with protection in animal models and in previously infected and vaccinated persons, although cellular immune responses and potentially non-neutralizing antibodies are likely to have contributed to protection in these studies^{53–59}. The development of neutralizing antibodies early in the course of infection has been associated with lower virus levels and greater protection from severe infection^{58,60–63}. Finally, the passive administration of neutralizing mAbs reduces the severity of infection when administered early^{62,64–66}.

High-resolution X-ray crystallography and cryo-electron microscopy structures have been published for more than 100 mAbs, including 5 with US Food and Drug Administration (FDA) Emergency Use Authorizations (EUAs) and several additional mAbs in phase III clinical trials⁶⁷. Most mAbs target either the RBD RBM or the RBD core; several target the NTD. Those targeting the RBM compete with RBD binding to ACE2. Those targeting the RBD core often cross-neutralize other SARS-related coronaviruses^{49,68,69}. The NTD-targeting neutralizing antibodies primarily bind a single epitope comprising the largest glycan-free surface facing away from the viral membrane referred to as the NTD supersite^{51,52,70}.

Several classification schemes have been developed to describe RBD-binding mAbs based upon whether they bind the RBM or RBD core, whether they bind the RBD in its up and/or down configuration and the extent to which they compete with other mAbs^{71–74}. Among those mAbs targeting the RBM, one group binds epitopes that overlap extensively with the ACE2 binding site and, as a result, binds solely when the RBD is in the open state. This group, which is referred to as class 1 mAbs, is typically encoded by the closely related *IGHV3-53* and *IGHV3-66* heavy chain genes and has short complementarity-determining region H3 loops. The second main group of RBM-binding mAbs (class 2) has a smaller ACE2 binding footprint and, as a result, can often bind the RBD in the closed state. Several other RBM-binding mAbs are more difficult to classify, including a class that binds a quaternary epitope involving more than one RBD^{71,72}.

Antibodies that target the RBD core also form two major clusters, one on the surface-accessible face of the RBD and another whose epitope is buried in the closed state^{71–74}. Antibodies that bind to the surface-accessible face of the RBD core can bind in either

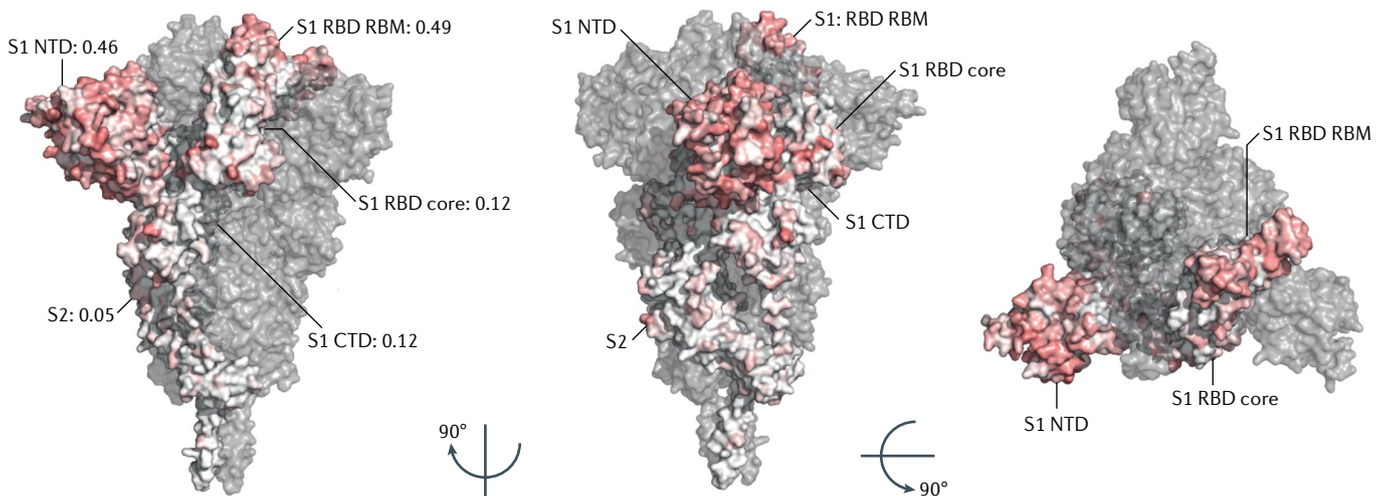


Fig. 2 | Genetic variability of the SARS spike proteins. Position-specific sequence variability and median domain-specific pairwise distances among SARS-related coronaviruses. Results were derived from an alignment of 24 representative sarbecovirus spike sequences having a nucleotide genetic distance (TN93 model) of ≥ 0.01 . Position-specific entropy is superimposed for one of three monomers on a surface representation of trimeric SARS-CoV-2 spike (Protein Databank (PDB) code: 6XR8), with white indicating conserved residues and the shade of red

indicating the extent of sequence variability. Two 90° rotated side views (left and middle panels) and one top view (right panel) of the spike trimer are shown. The median pairwise distance among SARS-related coronaviruses is greatest for the S1 amino-terminal domain (NTD) and receptor-binding domain (RBD). Within the RBD, the median pairwise distance is greater for the receptor-binding motif (RBM) than for the core region. CTD, carboxy-terminal domain; S1, exposed attachment domain; S2, partially buried fusion domain.

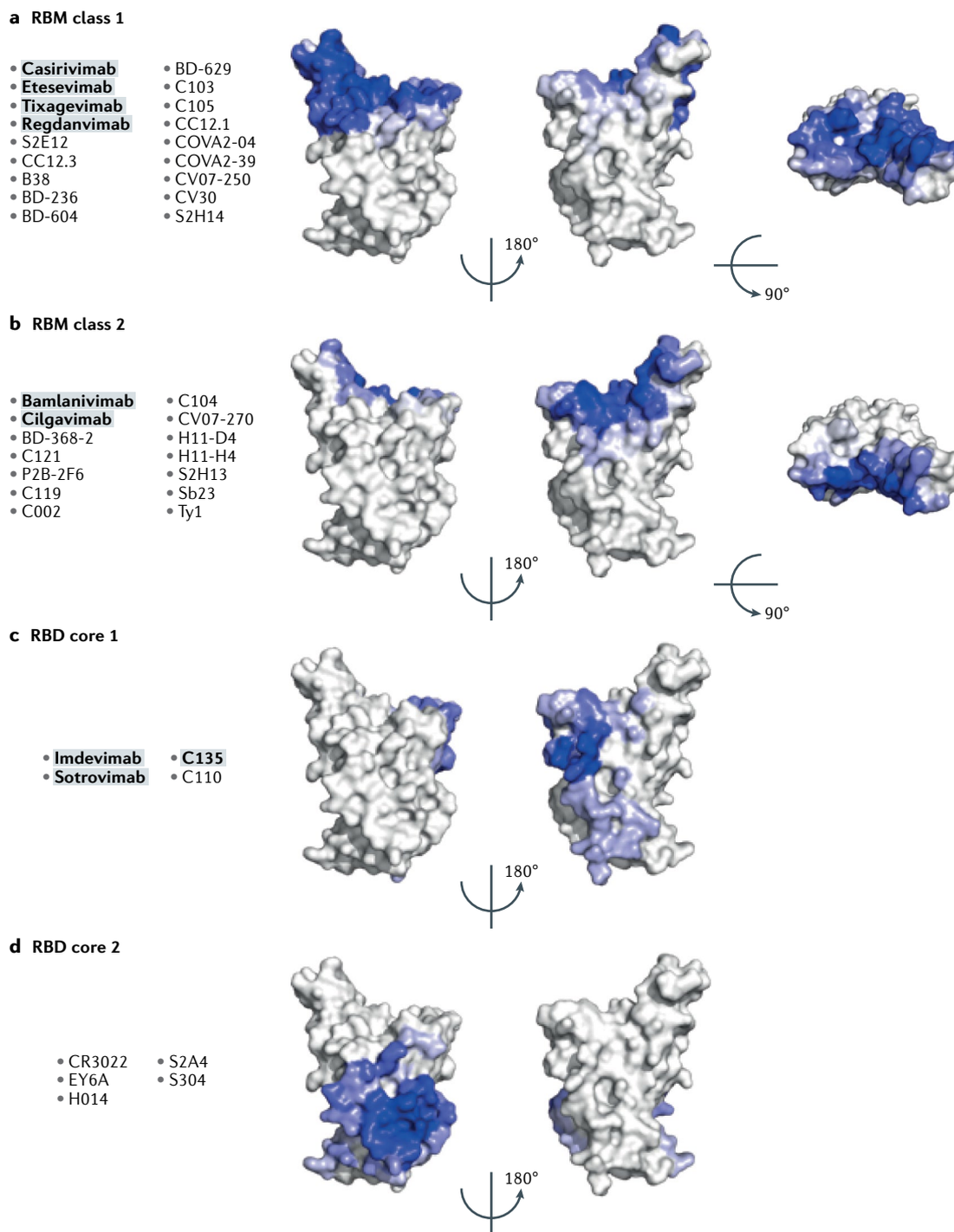


Fig. 3 | **SARS-CoV-2 spike-targeted antibody classifications.** Classification of monoclonal antibodies (mAbs) targeting the SARS-CoV-2 spike receptor-binding domain (RBD) epitopes. For the two classes of mAbs binding the receptor-binding motif (RBM), 90° rotated side views and one top view of a surface RBD representation are shown (parts **a,b**). For the two classes of RBD core-binding mAbs, just the 90° rotated side views are shown (parts **c,d**). Each image derived from coordinates of the Protein Databank (PDB) structure 6M0J. Bold highlighted mAbs are in phase III clinical trials (as of July 2021). Blue intensity is proportional to the number of mAbs binding to the underlying amino acid residues. RBM refers to the region of the RBD containing the angiotensin-converting enzyme 2 (ACE2)-binding residues. RBM class 1 mAbs (part **a**) bind the RBD only in its up position, whereas RBM class 2 mAbs (part **b**) can bind the RBD in its up or down position. A third RBM mAb class binds to a quaternary epitope comprising more than one RBD but is not shown as it would require the trimeric spike. Epitopes for amino-terminal domain-binding mAbs are not shown.

the open or closed state and those that target the RBD core epitope bind only in the open state. FIGURE 3 displays the epitopes of those mAbs with high-resolution

structures that are either in advanced clinical trials or have been assessed for their activity against viruses with mutant spike proteins. Supplementary Table 2

Fc-effector functions

Antibody functions mediated by their fragment crystallizable (Fc) region, including complement activation, antibody-dependent cellular cytotoxicity and antibody-dependent cellular phagocytosis. These functions may be particularly important for eliminating virally infected cells. They are more difficult to study *in vitro* than *in vivo*.

Convalescent plasma

Plasma samples from persons previously infected with SARS-CoV-2 usually contain neutralizing antibodies that bind to the SARS-CoV-2 spike protein and prevent it from infecting cells *in vitro*. Convalescent plasma samples are characterized by the number of months since recovery from infection, the severity of illness associated with infection, and the extent to which the plasma can be diluted and still retain the ability to prevent infection *in vitro* against different SARS-CoV-2 variants.

Deep mutational scanning

A method that makes use of next-generation sequencing to measure in a single experiment the activity of many unique variants of a protein.

describes the epitopes of those mAbs being studied in clinical trials.

In addition to RBD-targeting and NTD-targeting antibodies, there may be a protective role for neutralizing antibodies targeting other parts of the spike and for non-neutralizing antibodies. Several neutralizing antibodies bind the S1 CTD and S2 of multiple coronaviruses, although these have been much less potent than RBD-targeting and NTD-targeting mAbs^{75–77}. Non-neutralizing antibody Fc-effector functions such as complement activation, cellular cytotoxicity and phagocytosis have also been shown to afford protection in animal models^{78,79} and in the initial weeks following vaccine administration⁸⁰. However, the protection afforded by non-neutralizing antibodies in the absence of neutralizing antibodies is difficult to quantify^{81,82}.

The early development of cytotoxic T lymphocyte (CTL) responsiveness in persons infected with SARS-CoV-2 correlates with less severe COVID-19 illness^{83,84}. CTL responses also contribute to protection from severe infection in non-human primates in the presence of low titres of neutralizing antibodies^{53,85} and in persons with impaired humoral immunity⁸⁶. SARS-CoV-2 infection and immunization with the spike protein elicits helper T lymphocyte and CTL responses^{58,87,88}. Indeed, many studies have identified specific human leukocyte antigen (HLA)-restricted helper T cell and CTL spike epitopes^{58,89,90}. Analyses of peptide libraries from several VOCs have shown that, with a few exceptions, spike CTL epitopes either remain unchanged or able to bind most HLA molecules^{91,92}. Nonetheless, the number of T cell spike epitopes recognized by a person varies according to their HLA profile and is much lower than the total number of SARS-CoV-2 T cell epitopes^{93,94}. Mutations at T cell epitopes have also been observed within several cohorts with the common HLA type A*02 (REF.⁹⁵) and at low levels within the circulating viruses of individual patients⁹⁶.

In vitro selection and neutralization experiments

An increasing number of studies have described either the *in vitro* selection of SARS-CoV-2 immune escape spike mutations or the impact of mutations on the neutralizing activity of mAbs, convalescent plasma or plasma from vaccinated persons^{52,97,98}. Studies of plasma from patients who were previously infected provide insight into the risk of reinfection with a SARS-CoV-2 variant, whereas those from immunized persons are relevant to vaccine efficacy. Although most convalescent plasma samples studied so far were obtained prior to the emergence of immune escape variants ('pre-variant isolates'), more recent studies have studied plasma from patients infected with different VOCs^{99–104}.

In vitro selection experiments have been performed most commonly using non-replicative pseudo-typed or replication-competent chimeric viruses^{105–108}. SARS-CoV-2 pseudo-typed viruses are produced by co-transfecting a SARS-CoV-2 protein expression vector and a construct encoding the components required for replication of a different virus that lacks the coding sequence for their own surface protein — most

commonly vesicular stomatitis virus (VSV), HIV-1 or murine leukaemia virus. These constructs also encode a reporter gene such as luciferase or green fluorescent protein. Chimeric viruses contain the SARS-CoV-2 spike sequence in a VSV genome lacking the sequence for the VSV surface protein^{107,108}. VSV-based chimeric viruses are particularly useful for mutation selection studies because they can undergo multiple rounds of replication. There have also been reports describing the *in vivo* selection of spike mutations in animal models¹⁰⁹ and in persons infected with SARS-CoV-2 with prolonged infections or receiving mAbs^{109–114}.

Neutralization studies compare the ability of a mAb or plasma sample to inhibit the cellular entry of a virus containing one or more spike mutations with viruses lacking these mutations. Neutralization studies have been performed using either pseudo-typed viruses, VSV-based chimeric viruses, full-length cloned recombinant SARS-CoV-2 viruses^{115–120} or low-passage or plaque-purified cultured isolates^{121–124}. Pseudo-typed and chimeric viruses have been used most frequently because it is simpler to introduce mutations into a plasmid encoding just the spike gene compared with using a clinical virus isolate or a recombinant virus generated using a system that requires either multiple plasmids or bacterial or yeast artificial chromosomes^{119,120}. Although the results of neutralization experiments using pseudo-typed viruses, VSV-based chimeric viruses and full-length SARS-CoV-2 usually yield similar results^{97,107,125,126}, full-length viruses are expected to be more reliable and can be studied in animal models¹²⁷.

The effects of nearly all individual RBD mutations on protein expression in yeast (a correlate of protein folding stability), ACE2 binding and binding to a wide variety of mAbs and plasma samples have also been assessed using deep mutational scanning in which each yeast cell producing a different RBD mutation is labelled with a distinct genomic sequence^{50,128–131}. Although this approach does not quantify the effect of RBD mutations on mAb neutralization, it has proved useful as a screening assay to identify mutations that require further study in cell culture. High-throughput biochemical assays such as enzyme-linked immunosorbent assays (ELISAs) are also being developed to allow clinical laboratories to measure the ability of plasma to inhibit ACE2 binding to RBDs belonging to different SARS-CoV-2 variants^{132–134}.

In vitro neutralization experiments of SARS-CoV-2 variants are usually reported as a fold reduction in susceptibility compared with a control virus, such as the reference Wuhan-Hu-1 virus, variants containing just the D614G mutation or other pre-variant isolates⁹⁷. In the sections that follow, we summarize the neutralization susceptibility of different viruses as being <3-fold, 3–10-fold and >10-fold reductions compared with the control virus as these thresholds represent approximately one-half-log and one-log reductions in susceptibility. Nonetheless, the absolute level of neutralization is often more clinically relevant than the fold change compared with a control virus as the same reduction in susceptibility to a mAb or to vaccinee plasma will be more consequential for mAbs with low intrinsic activity or vaccines that do not elicit high levels of neutralizing

antibodies. In summarizing the results of neutralizing experiments, we have also pooled results obtained using pseudo-typed, chimeric and infectious viruses as the results of these assays are usually concordant^{97,107,125,126}.

Of the more than 6,000 plasma samples from persons receiving a complete course of vaccination, 76% were obtained from persons receiving an authorized mRNA vaccine (Pfizer/BioNTech BNT162b (54%) and Moderna mRNA-1273 (22%)), 9% from recipients of the adenovirus-vectored AstraZeneca AZD1222 vaccine, 6% from recipients of the inactivated Sinovac CoronaVac vaccine, 3% from recipients of the adenovirus-vectored Janssen Ad26.CoV2.s vaccine, 3% from recipients of the inactivated Bharat Biotech Covaxin (BBV152) vaccine and 1% each from recipients of the protein subunit Novavax NVX-CoV2373, the adenovirus-vectored Gamaleya Research Institute Sputnik V and the inactivated Sinopharm BBIBP-CorV vaccines.

SARS-CoV-2 mutations

Currently circulating SARS-CoV-2 VOCs and VOIs share several mutations that enable them to spread in the face of rising population immunity while maintaining or increasing their replication fitness. These mutations belong to a repertoire of recurrent mutations, most of which are in the spike gene. FIGURE 4 illustrates the most biologically and clinically significant spike mutations and their change in prevalence since the early stages of the pandemic. To understand the biological properties and epidemiological characteristics of the increasing number of SARS-CoV-2 VOCs and VOIs, it is necessary to understand their component mutations. Here, we divide these mutations into seven categories: D614G; the RBD mutation N501Y; the RBD mutation E484K; other RBD mutations; NTD mutations; mutations proximal to the S1/S2 furin cleavage site; and non-spike mutations.

D614G

The prevalence of the D614G mutation began increasing in late February 2020, and within several months it outcompeted all ancestral viruses and rose to a global prevalence approximating 100%⁸. Infectious virus clones with D614G replicated to higher levels in primary human airway cells and in the upper respiratory tracts of hamsters^{135–137}. D614G-containing virus clones were also associated with increased transmission between hamsters¹³⁶. Cryo-electron microscopy studies have shown that D614G disrupts one or more interprotomer contacts, resulting in a greater likelihood that one or more of the three RBDs are in an open versus closed position and, hence, compatible with ACE2 receptor binding^{138,139}. Subsequently, additional mutations in the spike CTD and in S2 have also been reported to possibly increase SARS-CoV-2 replication by a similar mechanism¹⁴⁰. D614G may also be responsible for increasing the number of spike proteins per virion^{141,142} and the rate of S1/S2 cleavage¹⁴³. Viruses with D614G have been slightly more susceptible to neutralization by mAbs, convalescent plasma and plasma from vaccinated individuals in some studies^{138,144} and slightly more resistant to neutralization in other studies^{136,145}.

Furin cleavage site

A short positively charged amino acid sequence at a specific location in a viral surface protein that is recognized by host furin protease enzymes. A furin cleavage site at the border of the SARS-CoV-2 spike S1 and S2 domains must be cleaved to enable viral cell fusion.

N501Y

N501Y is present in the Alpha, Beta and Gamma VOCs. N501Y increases ACE2 affinity^{128,146,147} and increases virus replication in human upper-airway cells and in the upper respiratory tracts of hamsters¹²⁷. N501Y does not influence the binding and neutralization of most mAbs^{48,109,117,121,148,149} (TABLE 1). Alone, it is also rarely associated with reduced susceptibility to convalescent plasma^{121,148–150} or plasma from persons receiving one of the two authorized mRNA vaccines (Pfizer/BioNTech BNT162b2 or Moderna mRNA-1273) or the Novavax NVX-CoV2373 protein subunit vaccine^{115,121,148,150–153} (FIG. 5).

E484K

E484 is recognized by a high proportion of the polyclonal antibodies developing within persons infected with SARS-CoV-2 (REF.⁵⁰). E484K is present in the Beta and Gamma VOCs^{16,19} and in the VOIs Eta (B.1.525), Iota (B.1.526)¹⁵⁴, Theta (P.3)^{155,156} and Zeta (p.2)¹⁵⁷. E484K has also been reported within several Alpha variant sub-lineages^{158,159}. E484Q has been reported in the Kappa VOI (B.1.617.1).

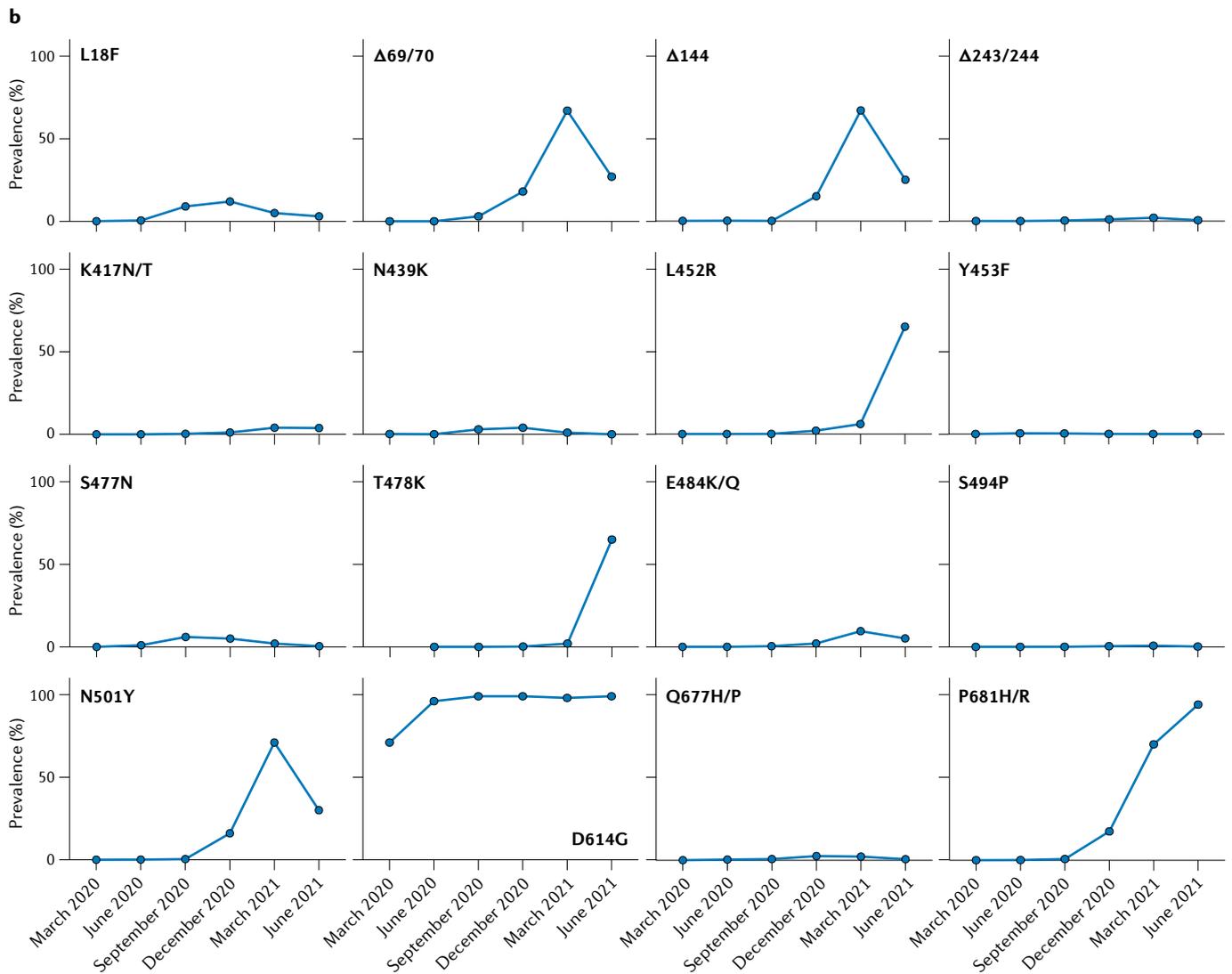
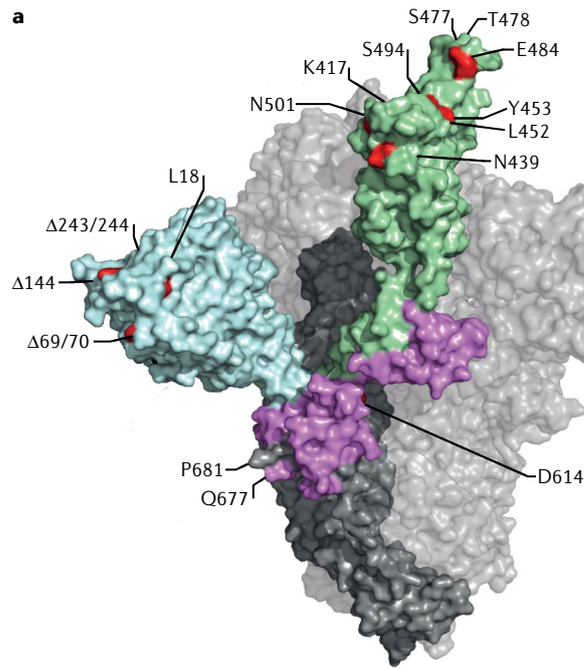
E484K has been selected *in vitro* by casirivimab and bamlanivimab and several other RBM class 1 and 2 mAbs^{48,105,110,130,160} and it reduces susceptibility to these mAbs^{64,105,109,117,121} (TABLE 1). E484K has resulted in 3-fold to 10-fold reduced susceptibility to about 30% and >10-fold reduced susceptibility to about 10% of convalescent plasma samples^{121,130,160–162} (FIG. 5). E484K has also resulted in 3-fold to 10-fold reduced susceptibility to about 30% of plasma samples from persons immunized with one of the authorized mRNA vaccines^{121,160–162} (FIG. 5).

Other RBD mutations

L452R is present in the Delta VOC, as well as the Kappa (B.1.617.1) and Epsilon (B.1.427/9) VOIs^{34,123}. It reduces susceptibility to several RBM class 2 mAbs, including bamlanivimab, but not to the other FDA EUA-approved mAbs^{109,110,160,163} (TABLE 1). L452R has resulted in 3-fold to 10-fold reduced susceptibility to about one third of convalescent and vaccinee plasma samples^{164–166} (FIG. 5). Pseudo-typed viruses containing L452R were associated with higher levels of cell entry in lung organoids compared with pseudo-typed viruses containing D614G alone but lower levels compared with pseudo-typed viruses containing N501Y (REF.¹²³).

K417N/T are present in the Beta (as K417N) and Gamma (as K417T) VOCs. K417N/T rarely occur in the absence of other RBM mutations, possibly because K417 mutations appear to reduce ACE2 binding^{130,159}. K417N confers >100-fold reduced susceptibility to etesevimab¹²⁹ and about 10-fold reduced susceptibility to casirivimab¹²¹ but retains susceptibility to bamlanivimab, imdevimab and sotrovimab¹²¹. K417N/T retain full susceptibility to plasma samples from persons previously infected with SARS-CoV-2 or immunized with one of the authorized mRNA vaccines^{117,121,167}.

N439K increases ACE2 affinity^{128,168,169} and reduces imdevimab susceptibility¹²⁹ (TABLE 1). Viruses containing N439K usually retain full susceptibility to convalescent



◀ Fig. 4 | **Locations and prevalence of key SARS-CoV-2 spike mutations.** **a** | Sites of 16 key S1 (exposed attachment domain) mutations on the SARS-CoV-2 spike trimer, including 9 in the receptor-binding domain (RBD; green), four in the amino-terminal domain (NTD; cyan) and three in the carboxy-terminal domain (CTD; purple). Spike trimer figure derived from a cryo-electron microscopy structure (Protein Databank (PDB) code 7BNN). S2 (partially buried fusion domain) shown in dark grey. Six of the RBD mutations (K417N/T, L452R, T478K, E484K and N501Y) are present in one or more of the World Health Organization (WHO) variants of concern (VOCs): Alpha (B.1.1.7), Beta (B.1.351), Gamma (P.1), and Delta (B.1.617.2). NTD mutations include three deletions and L18F, mutations present in two or more VOCs. CTD mutations include D614G, which became the consensus amino acid at this position prior to the emergence of the VOCs, and P681H and P681R, which are present in Alpha and Delta VOCs, respectively. **b** | Other than D614G, which has a prevalence close to 100%, the most prevalent mutations as of June 2021 are those present in Alpha (Δ 69/70, Δ 144, N501Y and P681H) and Delta (L452R, T478K, and P681R) VOCs. Prevalence data obtained from outbreak.info¹⁹⁶.

plasma^{50,148,169}. Increases in the prevalence of two lineages containing N439K were reported in the UK in September 2020, but their prevalence declined with the emergence of the Alpha variant (REF.¹⁶⁹).

Y453F emerged independently several times in mink lineages, including one that subsequently spread among humans but is no longer active¹⁷⁰. Y453F increases ACE2 binding but, nonetheless, remains rare^{128,140,171}. Y453F markedly reduces susceptibility to casirivimab but not to the other FDA EUA-approved mAbs^{105,167} (TABLE 1).

S477N was present in a variant that spread widely in Europe in the summer of 2020 (REF.¹⁷²). It increases the strength of ACE2 binding¹²⁸ but has since circulated at a low level. It has not been shown to reduce susceptibility to any of the FDA EUA-approved mAbs^{109,167}.

T478K is present in the Delta variant and a common variant in Mexico^{34,173}. By itself, it retains susceptibility to all but a few mAbs and to most convalescent and vaccinee plasma samples^{50,129,130,174}.

F490S and S494P are uncommon RBM mutations that have arisen independently within several Alpha variant sub-lineages¹⁵⁸. F490S is associated with highly reduced susceptibility to bamlanivimab but retains susceptibility to the other FDA EUA-approved mAbs^{64,105,175}. S494P is associated with >10-fold reduced susceptibility to bamlanivimab and about 5-fold reduced susceptibility to casirivimab^{64,109,175}.

NTD mutations

NTD deletions are present in several VOCs and VOIs, and have also been reported commonly in persons with prolonged SARS-CoV-2 infections^{112–114,176}. Deletions at positions 69–70 appear to be associated primarily with increased virus replication^{113,177} whereas those between positions 141–146 and 242–244 interfere with the neutralizing activity of NTD-binding antibodies^{51,121,178,179}. Other NTD mutations including L18F and D253Y also reduce susceptibility to NTD-neutralizing antibodies^{51,103,158}.

Mutations close to the S1/S2 furin cleavage site

Mutations just upstream of the polybasic S1/S2 furin cleavage — including Q675H/R, Q677H/P, N679K and P681H/R — have occurred independently in many SARS-CoV-2 variants¹⁸⁰. P681H is present in the Alpha VOC and the Theta VOI, and in several additional SARS-CoV-2 lineages¹⁵⁶. P681R is found in the Delta VOC and the Kappa VOI. The increased positive

charge associated with both P681H and P681R appears to influence virus tropism by increasing S1/S2 cleavage in human airway epithelial cells^{181–183}.

Non-spike mutations

Mutations outside the spike protein have been reported to increase SARS-CoV-2 transmissibility by antagonizing the host response to type I interferons. In one study, the Alpha and Beta variants displayed a mean 112-fold and 8-fold reduced susceptibility to several type I interferons, respectively, compared with an early pandemic virus¹⁸⁴. In a second study, the Alpha variant was found to cause lower levels of interferon- β (IFN β) expression and to be less sensitive to IFN β pretreatment compared with two early pandemic viruses¹⁸⁵. In this and in a third study, a D3L mutation in the Alpha variant nucleocapsid gene was found to introduce an enhanced transcription regulatory sequence (TRS) upstream of *Orf9b*, an interferon antagonist gene expressed as an alternative reading frame within the nucleocapsid coding region^{185–187}.

Although the nucleocapsid D3L mutation is not found in other VOCs or VOIs, most have mutations in genes associated with interferon antagonism. One example is a recurrent deletion (Δ 106–108) of unknown phenotypic consequence in *nsp6*, a component of the SARS-CoV-2 membrane-tethered complex that also antagonizes interferon¹⁸⁸ and is present in the Alpha, Beta and Gamma VOCs and the Eta, Iota and Lambda VOIs²⁵ (FIG. 1b). Another example is a recurrent adjacent three-nucleotide change in the nucleocapsid gene that probably arose by homologous recombination of the core sequence of the leader TRS and that results in the double amino acid change R203K/G204R and novel sub-genomic transcripts of unknown consequences^{189,190}.

SARS-CoV-2 variants

SARS-CoV-2 variants are classified according to their lineage and component mutations. As a result, viruses belonging to the same lineage but containing different subsets of mutations can be classified as different variants. Variants are characterized by their transmissibility, disease severity and ability to evade humoral immunity. Increased transmissibility is demonstrated by the ability of a variant to outcompete other variants and to display a higher effective reproduction rate and/or secondary attack rate compared with other circulating variants^{8,191–193}. Disease severity has been assessed using mortality data and rates of hospitalization^{194,195}. Variants associated with higher virus levels may be more transmissible and/or cause more severe disease. Evasion of humoral immunity has been assessed by comparing a variant's susceptibility to mAbs, convalescent plasma and vaccinee plasma with that of other variants^{52,97}. In the following sections, we summarize the biological, epidemiological and clinical characteristics of the WHO-defined VOCs and VOIs as of June 2021.

Alpha variant (B.1.1.7)

The Alpha variant spike mutations include the RBD mutation N501Y, P681H and NTD deletions at positions 69–70 and 144 (FIG. 1b). The position 69–70 deletion prevents the amplification of one of three genomic segments

Transcription regulatory sequence
(TRS). In coronavirus genomes, a short sequence located at several genomic locations that is responsible for producing a set of nested 3' and 5' co-terminal sub-genomic RNA molecules.

Effective reproduction rate
The average number of new infections caused by an infectious individual in a population where some individuals may no longer be susceptible. The effective reproduction rate of different variants is compared to control for factors other than transmissibility that might account for changes in the prevalence of different variants.

Secondary attack rate
The transmission of infection in a circumscribed group. The secondary attack of different variants is compared to control for factors other than transmissibility that might account for changes in the prevalence of different variants.

Table 1 | SARS-CoV-2 variants and their fold reduction in neutralization susceptibility to monoclonal antibodies in advanced clinical trials

Variant	BAM	ETE	BAM+ETE	CAS	IMD	CAS+IMD	SOT	REG	TIX	CIL	C144	C135	BRII-196	BRII-198	ADG20
Alpha (B.1.1.7)	1 ₁₂	3.6 ₉	1.3 ₃	1 ₁₄	0.8 ₁₅	1 ₈	2.7 ₁₂ ^a	1.4	1.5 ₅	0.7 ₅	–	0.9 ₂	0.5 ₃	0.2 ₃	1.4
Beta (B.1.351)	>100 ₉	>100 ₇	>100 ₄	59 ₁₁	0.8 ₁₂	1.2 ₈	1 ₉	27 ₂	4.9 ₄	0.7 ₄	–	0.9 ₂	0.6 ₅	6 ₃	2.5
Gamma (P.1)	>100 ₂	59 ₂	>100	>100 ₆	0.6 ₆	1 ₄	1 ₆	–	6.2 ₂	0.5 ₂	–	–	0.3	0.7	2.2
Delta (B.1.617.2)	>100 ₂	0.7 ₂	–	0.8 ₂	0.8 ₂	1	–	–	0.5	2.9	–	–	–	–	1.4
Kappa (B.1.617.1)	>100 ₂	0.6 ₂	5	4.6 ₂	1.3 ₂	1 ₂	0.7	–	0.7	5.1	–	–	–	–	2.5
Epsilon (B.1.427/9)	>100 ₂	3.3 ₂	7.7	1.3	2.1	1	0.8 ₂	14	–	–	–	–	–	–	–
N501Y	1 ₃	2.9 ₇	1	1 ₉	0.8 ₉	1 ₃	1.6 ₅ ^a	–	1.1 ₃	1 ₃	1.4 ₃	1.4 ₃	1 ₃	1.8 ₂	–
E484K	>100 ₄	2.7 ₆	17	13 ₁₂	1 ₁₂	1.6 ₆	0.6 ₅	–	6.4 ₄	1.5 ₄	>100 ₄	0.4 ₃	1.4 ₃	2.4 ₂	–
K417N	0.2 ₂	>100 ₅	1	8.9 ₇	0.9 ₆	0.8	0.6 ₄	–	0.3 ₃	0.6 ₃	0.7 ₂	0.3 ₂	1.8 ₃	0.3 ₂	–
L452R	>100 ₂	0.9 ₄	7.4	1.2 ₄	2 ₄	1.2 ₃	0.6	–	–	–	–	–	1.4	–	–
T478K	–	–	–	–	–	–	–	–	–	–	1.5	–	–	–	–
N439K	1.3	0.4 ₃	–	0.8 ₄	28 ₅	1.7	0.9 ₃	–	–	–	0.9 ₂	>100	1 ₂	1	–
Y453F	1.8	1.4 ₃	–	74 ₆	1.6 ₆	3.5	1.1	–	–	–	1.1	–	1	1.5	–
F490S	>100 ₂	1.1 ₂	1	1 ₂	1.4 ₂	1	0.8 ₂	–	–	–	4 ₂	1.2	–	–	–
S494P	86 ₂	0.6 ₂	1	4.5 ₃	0.9 ₂	1.1	2 ₂	–	–	–	73 ₂	0.8	0.7	1.6	–

Table shows fold reductions in neutralization (relative to control virus) for pseudo-typed and infectious viruses with combinations of spike mutations present in four World Health Organization (WHO)-defined variants of concern (VOCs; Alpha, Beta, Gamma and Delta), two WHO-defined variants of interest (VOIs; Kappa and Epsilon) and viruses containing individual spike mutations. Fold change is the median value of results, subscript is the number of results. ‘–’ indicates absence of susceptibility data. ^aLevels of fold reduction for SOT for N501Y and B.1.1.7 were much higher for IC₅₀ values than for IC₉₀ values. BAM, bamlanivimab (LY-CoV555); CAS, casirivimab (REGN10933); CIL, cilgavimab (AZD1061); ETE, etesevimab (LY-CoV016); IMD, imdevimab (REGN10987); REG, regdanvimab (CT-P59); SOT, sotrovimab (Vir-7831); TIX, Tixagevimab (AZD8895).

in a commonly used diagnostic PCR assay, resulting in a phenomenon referred to as S-gene target failure (SGTF), which has been used as a proxy for this variant¹⁹¹. The Alpha variant also contains several non-spike mutations including nsp6:Δ106–108 and the nucleocapsid mutations D3L, R203K and G204R, which may increase transmission by antagonizing innate immunity^{25,185–187}. By the second quarter of 2021, the Alpha variant accounted for the majority of infections in the USA and many European countries^{23,196}. Epidemiological studies suggest that it was approximately 50% more transmissible than previously circulating UK variants^{191,197,198}. It was also associated with threefold to eightfold higher upper-airway levels^{199–201} and an estimated 50% increased mortality^{194,195,202}.

The Alpha variant is susceptible to neutralization by most neutralizing mAbs as well as by most plasma samples from previously infected persons^{97,102,121,149,159,203–206} (FIG. 5). The fact that the Alpha variant is rarely associated with reduced susceptibility to convalescent plasma is consistent with it not being associated with an increased risk of reinfection²⁰⁷.

The Alpha variant has displayed 3-fold to 10-fold reduced susceptibility to approximately 15% of plasma samples from recipients of an authorized mRNA vaccine^{97,102,121,148,149,152,159,204,208,209}. In cohort studies from Israel and Qatar, the BNT162b vaccine also retained greater than 90% efficacy against this variant^{210,211}

(TABLE 2). In a post-hoc analysis of a NVX-CoV2373 clinical trial, vaccine efficacy was 86.3% against Alpha variants compared with 96.4% against non-Alpha variants²¹² (TABLE 2). Similar data for the AZD1222 vaccine have been inconsistent. In a study of vaccine trial participants, plasma samples were associated with a median reduction in neutralizing activity of ninefold to the Alpha variant compared with an earlier UK variant²¹³. In this trial, AZD1222 displayed a non-statistically significant reduction in efficacy against the Alpha variant (70%; 95% confidence interval (CI) 44–85%) compared with earlier variants (82%; 95% CI 68–89%)²¹³. However, in three other studies, the median reduction in neutralizing activity was between onefold and threefold^{102,214,215}.

Several Alpha variant sub-lineages have acquired additional mutations that might increase the risk of reinfection and vaccine failure, including the RBD mutations E484K, F490S and S494P (REF. 158).

Beta variant (B.1.351)

Between October 2020 and January 2021, daily cases in South Africa increased from approximately 2,000 to more than 20,000 reported cases per day. This increase occurred in a setting in which more than 30% of the population was estimated to have already been infected and was associated with the emergence of the Beta variant, which contains three RBD mutations (K417N, E484K and N501Y) and five NTD mutations,

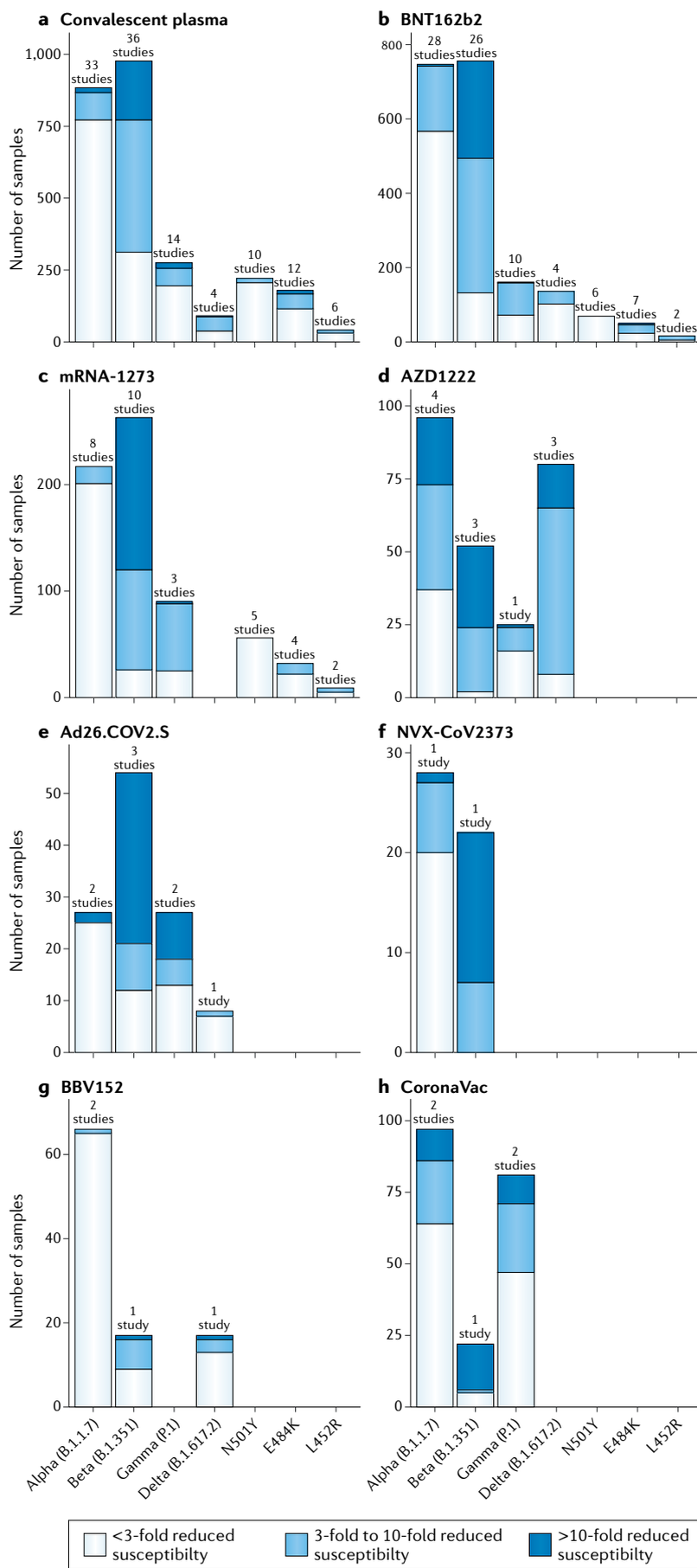


Fig. 5 | Effects of SARS-CoV-2 spike variants on susceptibility to neutralization. Fold-reduced susceptibility of the four variants of concern (VOCs; Alpha, Beta, Gamma and Delta) and three common spike receptor-binding domain (RBD) mutations (N501Y, E484K and L452R) to in vitro neutralization by plasma from previously infected persons (part a) and from persons vaccinated with the Pfizer/BioNTech BNT162b2 (part b), Moderna mRNA-1273 (part c), AstraZeneca AZD1222 (part d), Janssen Ad26.COVS.2.S (part e), Novavax NVX-CoV2373 (part f), Bharat Biotech BBV152 (part g) and Sinovac CoronaVac (part h) vaccines. y axes indicate number of plasma units tested. Colour scheme indicates fold reduction in neutralization. Only those data from plasma samples from persons receiving a full immunization schedule were included. Data obtained from <https://covdb.stanford.edu/search-drdb/> on 1 July 2021. In some plots, distributions are approximate as they include results reported only in aggregate as a mean fold reduction in susceptibility.

including a deletion within the NTD supersite at positions 242–244 (REF.¹⁶). The Beta variant was estimated to be 50% more transmissible than the lineages that preceded it¹⁶.

Reinfections with the Beta variant occurred commonly during a phase IIb trial of the NVX-CoV2373 vaccine performed in South Africa as approximately one third of infections in both the vaccine and placebo arms were reinfections²¹⁶. It is not known whether the Beta variant is associated with higher virus levels or disease severity because once detected it was no longer co-circulating with other lineages. As of June 2021, the Beta variant accounts for more than 50% of infections in many countries in sub-Saharan Africa^{23,196}.

The Beta variant is associated with reduced susceptibility to many mAbs because E484K interferes with binding of several RBM class 1 and 2 mAbs and K417N interferes with the binding of several RBM class 1 mAbs^{104,117,121,163,217}. Of the five FDA EUA-approved mAbs, bamlanivimab, etesevimab and casirivimab are largely inactive against B.1.351 whereas imdevimab and sotrovimab, which bind to the RBD core, retain neutralizing activity^{104,109,117,175,204} (TABLE 1).

The Beta variant has displayed 3-fold to 10-fold reduced susceptibility to 46% and >10-fold reduced susceptibility to 22% of convalescent plasma samples compared with early pandemic variants^{97,101,104,121,124,203,204,218–220} (FIG. 5). Of 34 convalescent plasma samples from persons infected with the Alpha variant, 59% had 3-fold to 10-fold reduced neutralizing activity and 18% had >10-fold reduced neutralizing activity to the Beta variant^{99,104,221}. Conversely, plasma from 22 persons infected with the Beta variant retained partial or full neutralizing activity against variants from earlier waves of the pandemic in all but three persons¹⁰¹.

Of plasma samples obtained from persons receiving one of the mRNA vaccines, 45% had 3-fold to 10-fold and 30% had >10-fold reduced Beta variant neutralizing activity^{97,104,117,121,151,204,208,209,222,223} (FIG. 5). Of plasma samples from persons receiving the AZD1222 vaccine, 42% had 3-fold to 10-fold and 54% had >10-fold reduced Beta variant neutralizing activity^{104,214,224} (FIG. 5).

Table 2 | Vaccine efficacy at preventing symptomatic infection for SARS-CoV-2 variants of concern

Variant	Pfizer/BioNTech BNT162b	AstraZeneca AZD1222 ^a	Novavax NVX-2373	Janssen AD26.CoV2 ^b	Sinovac CoronaVac ^c
Pre-variant	95% (trial) ^{56,59}	70% (trial) ^{59,248}	96% (trial) ^{59,212}	72% (trial) ^{59,225}	51% (trial) ⁵⁹
Alpha	90% (TNCC) ²¹¹	70% (trial) ²¹³	86% (trial) ²¹²	–	–
	92% (TNCC) ²³³	66% (TNCC) ²³²			
	93% (TNCC) ²³²	81% (TNCC) ²³³			
	97% (cohort) ²¹⁰				
Beta	75% (TNCC) ²¹¹	10% (trial) ²²⁴	51% (trial) ²¹⁶	64% (trial) ²²⁵	–
Gamma	–	–	–	–	42% (TNCC) ²⁴⁹
Delta	83% (TNCC) ²³³	60% (TNCC) ²³²	–	–	–
	88% (TNCC) ²³²	61% (TNCC) ²³³			

Table shows vaccine efficacy at preventing symptomatic SARS-CoV-2 infection with the Alpha, Beta, Gamma and Delta variants of concern (VOCs). Comparison is with efficacy against ‘pre-variant’ isolates. Data include persons receiving a full course of vaccination: 1–2 weeks following two vaccine doses for BNT162b, AZD1222, NVX-2373 and CoronaVac; 4 weeks after one vaccine dose for AD26.CoV2. ‘Trial’ refers to post-hoc analysis of clinical trial data. TNCC, test-negative case–control study; ‘–’, Not available. ^aAZD1222 efficacy data include some patients receiving the standard vaccine dose at both time points and others receiving a lower vaccine dose at the first time point. ^bAD26.CoV2 efficacy data based on preventing moderate to severe disease; pre-variant efficacy based on US data. ^cCoronaVac efficacy data against pre-variant isolates may include some persons infected with the Gamma VOC and the Zeta variant of interest (VOI). The TNCC study of vaccine efficacy against the Gamma VOC was performed in persons aged 70 years or older.

The Beta variant has also been associated with reduced vaccine efficacy (TABLE 2). In a phase II trial in South Africa of AZD1222 in which 39 HIV-negative patients became infected with the Beta variant, the estimated vaccine efficacy was just 10%²²⁴. In a phase II trial of NVX-CoV2373 in South Africa in which 38 of 41 infections were caused by the Beta variant, the estimated vaccine efficacy was 49%²¹⁶. In a phase III trial of the Janssen Ad26.COVS vaccine, vaccine efficacy in South Africa was 52% and 64% against moderate to severe COVID-19 with onset at least 14 and 28 days after administration, respectively²²⁵. In the previously cited Qatar cohort study, the efficacy of BNT162b against the Beta variant was estimated to be 72%, which was 15% lower than that for the Alpha variant²¹¹. Among nine BNT162b vaccine breakthrough infections in an Israeli case–control study, eight were with the Beta variant even though the Alpha variant predominated during the study period²²⁶.

Gamma variant (P.1)

The Gamma variant contains the RBD mutations N501Y, E484K and K417T (REF.¹⁹). It also contains five NTD mutations, of which L18F has been shown to interfere with the binding of NTD-targeting neutralizing antibodies⁵¹. As the Gamma variant was associated with a surge of infections in a region of Brazil estimated to have achieved a high infection rate, it is suspected of being able to infect and cause illness in persons previously infected with other variants^{19,29,30}. It was estimated to result in virus levels 3–4 times higher than earlier variants and to be responsible for an estimated 1.1-fold to 1.8-fold higher mortality¹⁹. By June 2021, the Gamma variant accounted for a high proportion of infections in several South American and Caribbean countries¹⁹⁶ and 10% of US infections³³. The resistance profile of the Gamma variant to the FDA EUA-approved mAbs is similar to that of the Beta variant^{103,109,217,227,228} (TABLE 1). Of convalescent

plasma samples obtained from persons infected with early pandemic variants or with the Alpha variant, about 20% had 3-fold to 10-fold and 10% had >10-fold reduced neutralizing activity^{97,103,117,203,217,228,229} (FIG. 5). Of plasma samples from persons receiving one of the two authorized mRNA vaccines, about 60% had 3-fold to 10-fold and 5% had >10-fold reduced neutralizing activity^{97,103,117,167,217,230} (FIG. 5). A similar distribution in the reduction in neutralizing activity was observed in plasma samples from recipients of the AZD1222 vaccine¹⁰³.

Delta variant (B.1.617.2)

As the pandemic surged in India in early 2021, two variants sharing a common ancestor, Delta (B.1.617.2) and Kappa (B.1.617.1), accounted for a high proportion of infections. The two variants probably diverged from a common ancestor between August and October 2020 (FIG. 1). Both variants contain the RBD mutation L452R, the proximal furin cleavage site mutation P681R and several mutations within *orf3*, *orf7a* and the nucleocapsid gene. The Kappa variant contained the RBD mutation E484Q whereas the Delta variant contained the RBD mutation T478K. The two variants also contain different mutations within *orf1a/b* and the spike NTD and S2 domains. Even though E484Q is more likely than T478K to evade antibody neutralization^{48,100,165}, only the Delta variant has demonstrated increased transmissibility, spreading to 54 countries and rapidly replacing the Alpha variant in the UK^{32,35} and the USA³³.

Among the FDA EUA-approved mAbs, the Delta variant is associated with high-level reduced bamlanivimab susceptibility^{100,214}. It results in approximately 3-fold to 10-fold reduced susceptibility to 45% and >10-fold reduced susceptibility to 5% of convalescent plasma samples^{100,118,214,215,231} (FIG. 5). Among plasma samples obtained from recipients of the BNT162b vaccine, approximately 15% displayed 3-fold to 10-fold reduced neutralizing activity against the Delta variant^{100,118,214} (FIG. 5).

Test-negative case–control study

(TNCC). A study design in which cases are patients presenting with a clinical syndrome who test positive for SARS-CoV-2 whereas controls are patients presenting with the same syndrome who test negative for SARS-CoV-2. The proportions of cases and controls who have been vaccinated are compared.

By contrast, among plasma samples obtained from recipients of the AZD1222 vaccine, higher proportions displayed reduced neutralizing activity against this variant^{100,209,214}. In case-control studies from the UK, the BNT162b vaccine was approximately 85% effective for the Delta variant whereas the AZD1222 vaccine was approximately 60% effective^{232,233}.

AY.1 and AY.2 are recently reported sub-lineages of the Delta variant that have developed additional mutations including the RBD mutation K417N (REF.¹⁹⁶).

Variants of interest

The Epsilon variant comprises two closely related lineages B.1.427 and B.1.429, first detected in California, USA. This variant was the first reported to contain the RBD mutation L452R. It was estimated to be about 20% more transmissible than co-circulating lineages and to be associated with twofold higher upper-airway virus levels^{123,192}. By February 2021, the Epsilon variant accounted for 15% of US infections, but by June 2021 its prevalence decreased to below 1%.

In contrast to the Delta VOC, the Kappa VOI has not demonstrated increased transmissibility. However, because of the presence of the RBD mutations L452R and E484Q, the Kappa variant has a somewhat greater ability to evade humoral immunity than the Delta variant. It displays reduced susceptibility to casirivimab and to bamlanivimab. Of convalescent plasma samples, about 40% had 3-fold to 10-fold and 15% had >10-fold reduced neutralizing activity^{100,122,234}. Of plasma samples from recipients of an mRNA vaccine, approximately 55% had 3-fold to 10-fold and 5% had >10-fold reduced neutralizing activity^{100,118,122,234}.

The Iota (B.1.526), Eta (B.1.525) and Zeta (P.2) variants are each characterized primarily by the RBD mutation E484K. The Iota variant was first identified in New York state. In June 2021, it had a prevalence of 5–10% in the USA but remained rare outside the USA¹⁵⁴. It contains the same nsp6 deletion as in the Alpha, Beta and Gamma variants. Of convalescent plasma samples, about 40% display 3-fold to 10-fold and 10% display >10-fold reduced Iota variant neutralizing activity^{166,235}. Of plasma samples from recipients of an mRNA vaccine, about 30% display 3-fold to 10-fold reduced neutralizing activity^{230,235}. The Eta variant is present at low levels in many countries, with Nigeria having the highest proportion of infections²³⁶. The ability of mAb, convalescent plasma and vaccinee plasma to neutralize the Eta variant has been infrequently studied^{118,166}. The Zeta variant was common in Brazil in late 2020 and early 2021 (REF.¹⁵⁷), but appears to be decreasing in prevalence.

The Theta variant was first reported in March 2021 in the Philippines. It contains 13 lineage-defining mutations including N501Y, E484K, P681H and the NTD deletion at positions 141–143 (REFS^{155,156}). However, it remains rare, accounting for a small proportion of infections even in the Philippines¹⁹⁶. The Lambda variant (C.37) has a unique set of spike mutations including L452Q and F490S within the RBD and the NTD deletion Δ 246–252 (REF.²³⁷). It is highly prevalent in several South American countries and is associated with reduced susceptibility to the locally used CoronaVac vaccine²³⁸.

Conclusions and implications for COVID-19

SARS-CoV-2 variants are characterized by their transmissibility, disease severity and ability to evade humoral immunity. The Alpha and Delta variants are each associated with increased transmissibility and greater disease severity because of immune evasion, and potentially because of higher virus levels resulting from the antagonism of innate immunity. The Beta and Gamma variants are each associated with increased transmissibility because of their ability to evade humoral immunity and cause reinfections. Although it is not surprising that mutations associated with reduced humoral immunity have recently emerged in association with rising population immunity, the concurrent emergence of mutations that intrinsically increase SARS-CoV-2 replication is more difficult to explain. The timing of this second category of mutations raises the possibility that they only emerged after a critical number of global infections or as compensation for subtle reductions in replication fitness associated with developing immune escape mutations.

As of June 2021, it is uncertain whether the current approaches to classifying variants will be sustainable. Should the current VOCs and VOIs develop multiple sub-lineages with additional biologically relevant mutations, it may become necessary to classify variants according to their component mutations rather than their ancestral lineages. However, despite the phenomenal progress in studying the impact of individual mutations in vitro and in animal models, classifying variants according to their component mutations will also prove challenging should the number of recurrent mutations also increase.

Although SARS-CoV-2 variants differ in their transmission rates, disease severity and risk of reinfection, there is no evidence that they are differentially affected by non-pharmaceutical public health measures such as social distancing and the use of personal protective equipment, or that they will respond differently to most antiviral therapies. Except for several mAb preparations (for example, bamlanivimab/etesevimab), most antiviral compounds are likely to retain activity against each of the current VOCs and VOIs. The two most recently approved mAb preparations, casirivimab/imdevimab and sotrovimab, retain activity against all VOCs and VOIs, in part because imdevimab and sotrovimab target the RBD core whereas the most prevalent immune escape mutations are in the RBD RBM. There have also been no reports of variants with mutations in the enzymatic targets of therapy that would reduce susceptibility to remdesivir or to the RNA-dependent RNA polymerase and protease inhibitors in clinical development.

The most important consequence of emergent SARS-CoV-2 variants, therefore, is their impact on vaccine efficacy. The levels of neutralizing mAbs elicited by the mRNA vaccines has been high and in most persons are likely to be maintained for many months above the levels required for protection against even the most resistant circulating variants^{164,239,240}. In addition, the mRNA vaccines appear to often elicit memory B cells that undergo somatic hypermutation, which should broaden the response to viruses with variant spike

proteins^{239,240}. Nonetheless, in vitro neutralization studies and epidemiological vaccine efficacy studies indicate weaker protection against emerging variants for most of the non-mRNA vaccines. Moreover, regardless of the vaccine received, reductions in vaccine-elicited humoral immunity is likely to be clinically significant for persons with impaired immunity as a result of underlying disease, immunosuppressive drugs or older age^{241–244}.

As the spectrum of SARS-CoV-2 variants is expanding and shifting faster than epidemiological studies can be conducted, laboratory correlates of protection against SARS-CoV-2 variants have become a high priority. As summarized in this Review, neutralizing antibody titres against pre-variant and variant SARS-CoV-2 isolates correlate with protection from infection in epidemiological studies. Whether this is because neutralizing antibodies are the most important means of viral protection and/or because these titres correlate with other aspects of protective immunity, including memory B cells, antibody-mediated effector functions and T cell immunity is not known. Therefore, a strategy that combines genomic surveillance, in vitro

neutralization studies and vaccine efficacy studies should be maintained to identify those variants that pose the greatest threat to current vaccines and to guide the development of immunogens for second-generation vaccines²⁴⁵.

An mRNA vaccine that incorporated most of the spike mutations present in the Beta variant (mRNA-1273.351; Moderna) was recently reported to increase both pre-variant and Beta variant-specific neutralizing antibody titres when administered as a booster to mice that had previously been immunized with mRNA-1273 (REF.²⁴⁶). Continued studies in animal species that are more predictive of responses in humans will be necessary to determine whether updated immunogens can broaden the response to multiple variants rather than just boost existing antibody responses generated by previous infections or vaccinations²⁴⁷. Moreover, as the spike protein has been found to contain multiple cytotoxic and helper T cell epitopes, it will be important that as many of these as possible are included in future vaccine preparations⁸⁹.

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Author contributions

K.T., P.L.T. and R.W.S. conceived of the project. K.T., P.L.T., J.N. and R.W.S. reviewed the primary literature. K.T. and P.L.T. created the database software necessary for the project. K.T. reviewed the primary data presented in the Review. K.T., P.L.T. and R.W.S. analysed the primary data presented in the manuscript. R.W.S. wrote the manuscript. D.F. assisted with creating manuscript figures. R.K.G., T.d.O., S.L.K.-P. and D.F. reviewed multiple drafts of the manuscript.

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