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Surveillance report

Surveillance generated by nf-ncov-voc for Omicron variant

Date

This report is generated on 2022-01-21 using 164953 number of genomes collected between 2020-02-25 and 2021-12-29

Pango Lineages

Pango Lineages in this report ['B.1.1.529', 'BA.1', 'BA.2']

Indicator

This table contains key indicators identified

Indicator	Sub-categories from POKAY	Mutations
Transmissibility between hu-	transmissibility	p.N440K
mans		
Infection Severity	ACE2 receptor binding affinity,	p.A701V, p.D614G, p.G339D,
	viral load, outcome hazard ratio	p.H655Y, p.H69del, p.K417N,
		p.N440K, p.N501Y, p.P681H,
		p.T95I, p.V70del
Immunity after natural infection	convalescent plasma escape, rein-	p.E484A, p.H69del, p.K417N,
	fection, humoral response dura-	p.N501Y, p.P681H, p.Q493R,
	bility	p.V70del
Vaccines	vaccine neutralization efficacy	p.D614G, p.K417N, p.N501Y,
		p.P681H
Monoclonal antibodies	monoclonal antibody serial pas-	p.E484A, p.G142D, p.K417N,
	sage escape, pharmaceutical ef-	p.N440K, p.N501Y, p.Q493R,
	fectiveness	p.R158G, p.R346K, p.S373P
Diagnostics	clinical indicators, antigenic test	
	failure, symptom prevalence	

Mutation Significance

This table contains key functional impacts of mutations identified

Mutations	Sub-category	Function	Lineages	Citation	Sequence	Alternate	Alternate
					Depth	Allele	Frequency
p.K417N	ACE2 receptor binding affinity	The K417N mutation decreased the affinity extasciitilde4 fold, mainly by decreasing the k(on) but also by increasing the k(off) as measured by surface plasmon resonance.	B.1.1.529, BA.1, BA.2	Barton et al. (2021)	750	T	1.0
p.K417N	ACE2 receptor binding affinity	This combination showed extasciitilde3x increase binding to ACE2 vs wild type, about half that of the B.1.1.7 lineage, suggesting that the K417N mutation is slightly detrimental to ACE2 binding, probably as a result of disrupting the salt bridge formed with ACE2 residue D30	B.1.1.529, BA.1, BA.2	Collier et al. (2021)	750	Т	1.0
p.K417N	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant showed a 1.5x decrease in binding (KD) relative to D614G.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	750	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.K417N	ACE2 receptor binding affinity	RBD containing the N501Y mutation results in 9-fold stronger binding to the hACE2 receptor than wild type RBD. The E484K mutation does not significantly influence the affinity for the receptor, while K417N attenuates affinity. As a result, RBD from B.1.351 containing all three mutations binds 3-fold stronger to hACE2 than wild type RBD but 3-fold weaker than N501Y.	B.1.1.529, BA.1, BA.2	Laffeber et al. (2021)	750	T	1.0
p.K417N	ACE2 receptor binding affinity	Studying the key covariants in lineage of concern 501Y.V2, observed about 2-fold increase in ACE2 binding vs wildtype, but greatly decreased mAb binding, suggesting evolutionary optimum tension between immune evasion and ACE2 binding affinity as the N501Y variant alone has 10x increase in affinity but no effect on tested mAb binding.	B.1.1.529, BA.1, BA.2	Liu et al. (2021)	750	T	1.0
p.K417N	ACE2 receptor binding affinity	Using Mircoscale Thermopheresis, the B.1.351 variant harboring three mutations, binds ACE2 at nearly five-fold greater affinity than the original SARS-COV-2 RBD (Kd 87.6, vs 402.5 nM).	B.1.1.529, BA.1, BA.2	Ramanathan et al. (2021)	750	Т	1.0
p.K417N	ACE2 receptor binding affinity	Reported 3-fold decrease in affinity compared to wild-type RBD on the cell surface (Kd	B.1.1.529, BA.1, BA.2	Tian et al. (2021)	750	Т	1.0
p.K417N	ACE2 receptor binding affinity	Reported slight increase in affinity compared to wild-type RBD on the cell surface (Kd	B.1.1.529, BA.1, BA.2	Tian et al. (2021)	750	Т	1.0
p.K417N	ACE2 receptor binding affinity	The affinity of ACE2 for this mutation combination was twice as high as for wild type. Having in mind that the affinity of SARS-CoV-2 for ACE2 is only 4-fold higher compared to SARS-CoV-1, this factor of 2 is expected to be biologically significant.	B.1.1.529, BA.1, BA.2	Vogel et al. (2021)	750	Т	1.0
p.K417N	antibody epitope effects	>20% (ELISA significance threshold) drop in antibody binding (ELISA) by this variant against IgG1 monoclonal antibody ab1.	B.1.1.529, BA.1, BA.2	Sun et al. (2021)	750	Т	1.0
p.K417N	antibody epitope effects	5 antibodies tested were less potent against K417N by ten-fold or more (class 1 mAbs)	B.1.1.529, BA.1, BA.2	Wang et al. (2021)	750	Т	1.0
p.K417N	antibody epitope effects	Pseudotyped virus model ablates binding by RBD-directed mAbs CB6 and 910-30 (targeting the inner side of the RBD). Pseudotyped virus model impairs binding by RBD-directed mAbs 4-20 and REGN10933.	B.1.1.529, BA.1, BA.2	Wang et al. (2021)	750	T	1.0
p.K417N	convalescent plasma binding	2.16x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	750	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.K417N	convalescent plasma escape	The 501Y.V2 to first wave IC50 ratio ranged from 6 to 200-fold. Averaging across all 7 participant convalescent sera highlighted the dramatic decrease in sensitivity to neutralization of authentic 501Y.V2 variants. PG: I'm purposefully ignoring D614G and A701V as contributors	B.1.1.529, BA.1, BA.2	Cele et al. (2021)	750	Т	1.0
p.K417N	convalescent plasma escape	In 19 convalescent human sera extasci- itilde1mo post infection, Two-tailed Wilcoxon matched-pairs signed- rank test shows mild resistence P	B.1.1.529, BA.1, BA.2	Chen et al. (2021)	750	Т	1.0
p.K417N	convalescent plasma escape	27% of 44 early pandemic exposure convalescent plasma/sera lose all activity against a RBD triple mutant pseudovirus (RBD mutatants of the 501Y.V2 "South African" lineage), while only 23% retained high titres	B.1.1.529, BA.1, BA.2	Wibmer et al. (2021)	750	Т	1.0
p.K417N	convalescent plasma escape	Nearly half (21 of 44, 48%) of early pandemic exposure convalescent plasma/sera failed to neutralize the 501Y.V2 ("South African") lineage pseudovirus construct Only 3 of 44 convascent sera (those with the highest titer, which correlated directly with initial infection severity) had high neutralization against this 501Y.V2 PG: note that lineage variant R246I was excluded from the text in reference to these sera assays, not sure if that was an oversight.	B.1.1.529, BA.1, BA.2	Wibmer et al. (2021)	750	T	1.0
p.K417N	gene expression increase	Experimentally, Spike gene expression in- creased 0.1 fold	B.1.1.529, BA.1, BA.2	Starr et al. (2020)	750	Т	1.0
p.K417N	monoclonal anti- body serial passage escape	Escape mutation against monoclonal antibody LY-CoV016	B.1.1.529, BA.1, BA.2	Starr et al. (2021)	750	Т	1.0
p.K417N	monoclonal anti- body serial passage escape	In vitro selection against class 1 (Spike 'up' conformation) monoclonal antibody C682, and to a lesser extent C614 and C660	B.1.1.529, BA.1, BA.2	Wang et al. (2021)	750	Т	1.0
p.K417N	pharmaceutical effectiveness	COR-101 lost extasci- itilde6x binding against this isolated mutation. Estesevimab lost ex- tasciitilde100x binding against this isolated mutation.	B.1.1.529, BA.1, BA.2	Engelhart et al. (2021)	750	Т	1.0
p.K417N	pharmaceutical effectiveness	Tixagevimab, Regdan- vimab and COR-101 display reduced binding affinity to virus pseu- dotyped as RBD from B.1.351.	B.1.1.529, BA.1, BA.2	Engelhart et al. (2021)	750	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.K417N	pharmaceutical effectiveness	COR-101 lost extasci- itilde20x binding against this double mutation. Estesevimab lost ex- tasciitilde16x binding against this double mu- tation. Regdanvimab lost extasciitilde6x bind- ing against this double mutation. M396 lost extasciitilde10x bind- ing against this double mutation.	B.1.1.529, BA.1, BA.2	Engelhart et al. (2021)	750	Т	1.0
p.K417N	pharmaceutical effectiveness	This mutated version of RBD completely abol- ishes the binding to a therapeutic antibody, Bamlanivimab, in vitro.	B.1.1.529, BA.1, BA.2	Liu et al. (2021)	750	Т	1.0
p.K417N	trafficking	extasciitilde2x more infectivity than D614G alone in HEK293T- ACE2 cells 48h post- transduction.	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	750	Т	1.0
p.K417N	trafficking	extasciitilde9x more infectivity than D614G alone in HEK293T- ACE2 cells 48h post- transduction (no syn- ergy as level approx. that of N501Y alone).	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	750	Т	1.0
p.K417N	trafficking	Lentiviral pseudotyped with this individual mutation from B.1.351 was tested on ACE2.293T cells. Luciferase activity was measured two days postinfection, showing mild decrease in infection rate amongst the cells, suggesting that this mutation does not contributing to cell entry fitness.	B.1.1.529, BA.1, BA.2	Tada et al. (2021)	750	Т	1.0
p.K417N	vaccine neutralization efficacy	This variant showed only minor in Pfizer sera (one or two dose) neutralization efficiency vs D614G (using lentivirus pseudotype).	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	750	Т	1.0
p.K417N	vaccine neutraliza- tion efficacy	This variant showed >5x decrease in Pfizer sera (3 weeks post-first dose: n	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	750	Т	1.0
p.K417N	vaccinee plasma binding	1.76x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously nonseroconverted subjects. 1.75x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	750	Т	1.0
p.K417N	virion structure	Estimated free energy change (ddG) for this variant is -0.86 kcal/mol (i.e. destabilizing relative to wild type)	B.1.1.529, BA.1, BA.2	Spratt et al. (2021)	750	Т	1.0
p.S373P	monoclonal anti- body serial passage escape	Reduce affinity for mildly cross-reactive CR3022 (2003 pandemic SARS monoclonal anti- body cross-reactive to SARS-CoV-2)	B.1.1.529, BA.1, BA.2	Long et al. (2020)	862	С	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.Q493R	antibody epitope effects	Ablates binding by class 2 mAbs such as C144 that directly interfere with ACE2 binding, but clonal somatic mutations of memory B cells at 6.2 months (evolving humoral immune response) show pronounced increase in binding to the variant.	B.1.1.529, BA.1, BA.2	Gaebler et al. (2021)	862	G	1.0
p.Q493R	antibody epitope effects	Massive reduction in binding efficiency vs wild type for mAbs CB6/LY-CoV16 and LY-CoV555.	B.1.1.529, BA.1, BA.2	Rappazzo et al. (2021)	862	G	1.0
p.Q493R	antibody epitope effects	Somewhat resistent to some class 1 (Spike 'up') antibodies tested. Mix of non- to strongly resistent in class 2 antibodies tested.	B.1.1.529, BA.1, BA.2	Wang et al. (2021)	862	G	1.0
p.Q493R	convalescent plasma escape	Escape mutant found after in passage in plasma pool of 26 convalescents mean 1.5 post symptom onset.	B.1.1.529, BA.1, BA.2	Schmidt et al. (2021)	862	G	1.0
p.Q493R	monoclonal anti- body serial passage escape	The engineered mutation cause 10-fold or more increase in the disassociation constant with C144, C002 and C121 monoclonal antibodies vs. wild type Spike protein RBD domain AAs.	B.1.1.529, BA.1, BA.2	Barnes et al. (2020)	862	G	1.0
p.Q493R	monoclonal anti- body serial passage escape	Escape mutation against monoclonal antibody LY-CoV555 (antibody that forms the basis for Eli Lilly's bam- lanivimab)	B.1.1.529, BA.1, BA.2	Starr et al. (2021)	862	G	1.0
p.Q493R	monoclonal anti- body serial passage escape	Class 2 mAb C627 modestly selected for the emergence of this mutation in vitro.	B.1.1.529, BA.1, BA.2	Wang et al. (2021)	862	G	1.0
p.Q493R	monoclonal anti- body serial passage escape	Strong positive selection (up to 37% of super- natant sequences) after two rounds of C135 mon- oclonal antibody pas- sage, overall 76% switch away from Q493 to K or R	B.1.1.529, BA.1, BA.2	Weisblum et al. (2020)	862	G	1.0
p.R158G	monoclonal anti- body serial passage escape	Escape mutation against Spike N terminal do- main antigenic supersite i mAb S2X28	B.1.1.529	McCallum et al. (2021)	1	G	1.0
p.D614G	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains- Fc portion IgG complex, this variant showed es- sentially no change in binding (KD) relative to D614G.	BA.1	Gong et al. (2021)	860	G	1.0
p.D614G	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains- Fc portion IgG complex, this variant showed a 1.21x increase in binding (KD) relative to D614G.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	G	1.0
p.D614G	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant showed a 1.51x increase in binding (KD) relative to D614G, mostly due to decreased in "off-rate" a.k.a. dissociation rate (Kdis).	BA.1	Gong et al. (2021)	860	G	1.0
p.D614G	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains- Fc portion IgG complex, this variant showed a 1.5x decrease in binding (KD) relative to D614G.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	G	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.D614G	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant showed a 2.52x increase in binding (KD) relative to D614G, mostly due to decreased in "off-rate" a.k.a. dissociation rate (Kdis). Compare to full Spike variant complements for major lineages containing this variant subset: 5.43x (B.1.1.7 aka Alpha), 3.56x (B.1.351 aka Beta), 4.24x (P.1 aka Gamma).	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	G	1.0
p.D614G	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains- Fc portion IgG complex, this variant showed a 1.23x decrease in binding (KD) relative to D614G.	B.1.1.529, BA.1	Gong et al. (2021)	861	G	1.0
p.D614G	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains- Fc portion IgG complex, this variant showed a 1.33x decrease in binding (KD) relative to D614G.	B.1.1.529, BA.1	Gong et al. (2021)	861	G	1.0
p.D614G	ACE2 receptor binding affinity	In four cell lines (including 293T-hACE2 cells), this mutation combination increases infectivity vs D614G alone	B.1.1.529, BA.1, BA.2	Li et al. (2020)	862	G	1.0
p.D614G	convalescent plasma binding	1.11x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	BA.1	Gong et al. (2021)	860	G	1.0
p.D614G	convalescent plasma binding	1.48x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	G	1.0
p.D614G	convalescent plasma binding	1.33x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	BA.1	Gong et al. (2021)	860	G	1.0
p.D614G	convalescent plasma binding	2.16x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	G	1.0
p.D614G	convalescent plasma binding	1.65x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	G	1.0
p.D614G	convalescent plasma binding	1.26x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	B.1.1.529, BA.1	Gong et al. (2021)	861	G	1.0
p.D614G	convalescent plasma binding	No change in Spike binding (relative to D614G alone) by 5 plasma collected 8 months postsymptom-onset.	B.1.1.529, BA.1	Gong et al. (2021)	861	G	1.0
p.D614G	immunosuppression variant emergence	Studying 94 COVID-19 extended infection cases with genomics April 1 to October 17, 2020, one case developed 23 mutations in a 19 day period, including this combination in Spike.	B.1.1.529, BA.1, BA.2	Landis et al. (2021)	862	G	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.D614G	reinfection	A 47yo Indian male was reinfected with B.1.36 lineage virus in September 2020 after infection with genetically distinct B.1.36 virus in July, with negative PCR tests in between. While the forst episode was asymptomatic, the second included fever, cough, and malaise. The second case additionally contained stopgain ORF3a:E261*	B.1.1.529, BA.1, BA.2	Rani et al. (2021)	862	G	1.0
p.D614G	syncytium forma- tion	Slight increase in Vero cell-cell membrane fu- sion assay under infec- tion with VSV pseudo- typed virus.	B.1.1.529, BA.1, BA.2	Kim et al. (2021)	862	G	1.0
p.D614G	syncytium formation	Slight increase in Vero cell-cell membrane fusion assay under infection with VSV pseudotyped virus relative to wild type, no change relative to D614G.	B.1.1.529, BA.1, BA.2	Kim et al. (2021)	862	G	1.0
p.D614G	tissue specific neutralization	The nasal mucosa of Pfizer vaccinees with time course collection was evaluated against VSV pseudotypes: results (only one nasal swab from different previously infected vacinee neutralizing at weeks 3 and 6 against B.1.1.7 and D614G) suggest that vaccinees probably do not elicit an early humoral response detectable at mucosal surfaces even though sera neutralization was observed. They strengthen the hypothesis that some vaccines may not protect against viral acquisition and infection of the oral-nasal region, but may prevent severe disease associated with viral dissemination in the lower respiratory tract.	B.1.1.529, BA.1, BA.2	Planas et al. (2021)	862	G	1.0
p.D614G	trafficking	Circulating variant shown in vitro to not have major defects or enhancement of cell sur- face protein trafficking (i.e. Spike cleavage or fusion required for cell entry)	B.1.1.529, BA.1, BA.2	Barrett et al. (2021)	862	G	1.0
p.D614G	trafficking	The increased transduction with Spike D614G ranged from 1.3- to 2.4-fold in Caco-2 and Calu-3 cells expressing endogenous ACE2 and from 1.5-to 7.7-fold in A549ACE2 and Huh7.5ACE2 over-expressing ACE2. Although there is minimal difference in ACE2 receptor binding between the D614 and G614 Spike variants, the G614 variant is more resistant to proteolytic cleavage, suggesting a possible mechanism for the increased transduction.	B.1.1.529, BA.1, BA.2	Daniloski et al. (2021)	862	G	1.0
p.D614G	trafficking	No change in infectivity (24h) relative to D614G	B.1.1.529, BA 1 BA 2	Kim et al. (2021)	862	G	1.0
		alone in Caco-2 cells, Vero or Calu-3.	BA.1, BA.2	(2021)			



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.D614G	trafficking	extasciitilde4x more efficient S2 domain cleavage compared to wild type in Caco-2 cells, mid-range of three cell line tested (Vero and Calu-3).	B.1.1.529, BA.1, BA.2	Kim et al. (2021)	862	G	1.0
p.D614G	trafficking	More efficient infectivity (24h) compared to wild type, in Caco-2 cells extasciitilde9x, Vero extasciitilde8x, and Calu-3 extasciitilde8x. Compare to wild type at extasciitilde5x across cell types.	B.1.1.529, BA.1, BA.2	Kim et al. (2021)	862	G	1.0
p.D614G	trafficking	extasciitilde4x more efficient S2 domain cleavage compared to wild type, no change relative to D614G alone in Caco-2 cells, mid-range of three cell line tested (Vero and Calu-3).	B.1.1.529, BA.1, BA.2	Kim et al. (2021)	862	G	1.0
p.D614G	trafficking	extasciitilde2x more infectivity than D614G alone in HEK293T- ACE2 cells 48h post- transduction.	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	862	G	1.0
p.D614G	trafficking	extasciitilde9x more infectivity than D614G alone in HEK293T-ACE2 cells 48h post-transduction (no synergy as level approx. that of N501Y alone).	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	862	G	1.0
p.D614G	trafficking	9x more infectivity than D614G alone in HEK293T-ACE2 cells 48h post-transduction.	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	863	G	1.0
p.D614G	trafficking	Among S variants tested, the D614G mutant shows the highest cell entry (extascitilde3.5x wild type), as supported by structural and binding analyses.	B.1.1.529, BA.1, BA.2	Ozono et al. (2020)	862	G	1.0
p.D614G	trafficking	We report here pseudoviruses carrying SG614 enter ACE2-expressing cells more efficiently than wild type (extasciitilde9-fold). This increased entry correlates with less S1-domain shedding and higher S-protein incorporation into the virion. D614G does not alter S-protein binding to ACE2 or neutralization sensitivity of pseudoviruses. Thus, D614G may increase infectivity by assembling more functional S protein into the virion.	B.1.1.529, BA.1, BA.2	Zhang et l. (2020)	862	G	1.0
p.D614G	vaccine neutraliza- tion efficacy	Pseudotyped D614G virus has reduced neutralization activity vs wild type: 1.2x (37 sera Pfizer median 9 days post 2nd dose, 37 sera Moderna median 18 days post 2nd dose). This was NOT significant by ANOVA.	B.1.1.529, BA.1, BA.2	Garcia- Beltran et al. (2021)	862	G	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.D614G	vaccine neutralization efficacy	Using a lentivirus virus pseudotyped with D614G Spike, sera from vaccinated individuals who received the second dose (9–11 days post-second dose of Pfizer) exhibited a robust neutralizing potential, with a mean NT50 value of 99,000. This was an average of a 2-fold increase, relative to sera drawn from the individuals who received one dose of vaccination—mean NT50 dilution of 51,300. Importantly, a 6-fold increase in mean NT50 dilution was obtained when sera from the first vaccination dose was compared to convalescent sera from cohort with severe disease (NT50 51,000 vs 8,700) 21 to 63 days post-onset.	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	862	G	1.0
p.D614G	vaccine neutralization efficacy	This variant showed only minor in Pfizer sera (one or two dose) neutralization efficiency vs D614G (using lentivirus pseudotype).	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	862	G	1.0
p.D614G	vaccine neutraliza- tion efficacy	This variant showed >5x decrease in Pfizer sera (3 weeks post-first dose: n	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	862	G	1.0
p.D614G	vaccine neutralization efficacy	This variant showed no change in Pfizer sera (one or two dose) neutralization efficiency vs D614G (using lentivirus pseudotype).	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	862	G	1.0
p.D614G	vaccine neutralization efficacy	No significant change in virus neutralzation by 18 Pfizer two dose vaccinee sera compared to B.1.1.7. [results without including the used mutation A27S likely generalizable, as this is not a lineage defining mutation]	B.1.1.529, BA.1	Zuckerman et al. (2021)	861	G	1.0
p.D614G	vaccinee plasma binding	1.19x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously nonseroconverted subjects. 1.1x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	BA.1	Gong et al. (2021)	860	G	1.0
p.D614G	vaccinee plasma binding	1.04x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously nonseroconverted subjects. 1.09x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	G	1.0



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Mutations	Sub-catego	ory	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.D614G	vaccinee binding	plasma	1.14x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously non-seroconverted subjects. 1.09x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	BA.1	Gong et al. (2021)	860	G	1.0
p.D614G	vaccinee binding	plasma	1.76x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously nonseroconverted subjects. 1.75x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	G	1.0
p.D614G	vaccinee binding	plasma	1.17x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously nonseroconverted subjects. 1.09x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	G	1.0
p.D614G	vaccinee binding	plasma	1.14x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously non-seroconverted subjects. 1.11x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	B.1.1.529, BA.1	Gong et al. (2021)	861	G	1.0
p.D614G	vaccinee binding	plasma	1.16x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously non-seroconverted subjects. 1.02x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	B.1.1.529, BA.1	Gong et al. (2021)	861	G	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.D614G	viral load	Hamsters infected with SARS-CoV-2 expressing spike(D614G) (G614 virus) produced higher infectious titres in nasal washes and the trachea, but not in the lungs, supporting clinical evidence showing that the mutation enhances viral loads in the upper respiratory tract of COVID-19 patients and may increase transmission.	B.1.1.529, BA.1, BA.2	Plante et al. (2020)	862	G	1.0
p.D614G	virion structure	Estimated free energy change (ddG) for this variant is 2.5 kcal/mol (i.e. stabilizing relative to wild type)	B.1.1.529, BA.1, BA.2	Spratt et al. (2021)	862	G	1.0
p.D614G	virion structure	Negative stain EM shows increased proportion of "one-up" trimer conformation of Spike proteins on the surface of virions, where the up conformation is presumed to be more likely to bind ACE2.	B.1.1.529, BA.1, BA.2	Weissman et al. (2020)	862	G	1.0
p.D614G	virion structure	CryoEM shows increased proportion of "one-up" trimer conformation of Spike proteins on the surface of virions, where the up conformation is presumed to be more likely to bind ACE2.	B.1.1.529, BA.1, BA.2	Yurkovetskiy et al. (2020)	862	G	1.0
p.D614G	virion structure	Based on pseudotyped virus experiments, D614G may increase infectivity by assembling more functional S protein into the virion.	B.1.1.529, BA.1, BA.2	Zhang et al. (2020)	862	G	1.0
p.N501Y	ACE2 receptor binding affinity	The N501Y mutation had the biggest effect on ACE2 affinity of any VOC mutation tested, increasing the affinity extasciitilde10 fold to KD extasciitilde7 nM, by increasing the k(on) extasciitilde1.8 fold and decreasing the k(off) by extasciitilde 7 fold as measured by surface plasmon resonance.	B.1.1.529, BA.1, BA.2	Barton et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	This combination showed extasciitilde3x increase binding to ACE2 vs wild type, about half that of the B.1.1.7 lineage, suggesting that the K417N mutation is slightly detrimental to ACE2 binding, probably as a result of disrupting the salt bridge formed with ACE2 residue D30	B.1.1.529, BA.1, BA.2	Collier et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	The most frequent RBM mutation N501Y (165,519 instances) makes defective the atypical N-glycosylation sequon NGV 501-503, becoming a key RBM position for the interaction with hACE2-binding hotspot 353.	B.1.1.529, BA.1, BA.2	Gamez et al. (2021)	862	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.N501Y	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant showed a 2.52x increase in binding (KD) relative to D614G, mostly due to decreased in "off-rate" a.k.a. dissociation rate (Kdis). Compare to full Spike variant complements for major lineages containing this variant subset: 5.43x (B.1.1.7 aka Alpha), 3.56x (B.1.351 aka Beta), 4.24x (P.1 aka Gamma).	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	RBD containing the N501Y mutation results in 9-fold stronger binding to the hACE2 receptor than wild type RBD. The E484K mutation does not significantly influence the affinity for the receptor, while K417N attenuates affinity. As a result, RBD from B.1.351 containing all three mutations binds 3-fold stronger to hACE2 than wild type RBD but 3-fold weaker than N501Y.	B.1.1.529, BA.1, BA.2	Laffeber et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	Reported 10-fold increase in ACE2 binding vs wildtype (Kd	B.1.1.529, BA.1, BA.2	Liu et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	Studying the key covariants in lineage of concern 501Y.V2, observed about 2-fold increase in ACE2 binding vs wildtype, but greatly decreased mAb binding, suggesting evolutionary optimum tension between immune evasion and ACE2 binding affinity as the N501Y variant alone has 10x increase in affinity but no effect on tested mAb binding.	B.1.1.529, BA.1, BA.2	Liu et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	extasciitilde4-fold in- crease in binding affinity vs wild type.	B.1.1.529, BA.1, BA.2	Motozono et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	Using Microscale Thermopheresis, this variant binds ACE2 at nearly two-fold greater affinity than the original SARS-COV-2 RBD (203.7 nM vs 402.5 nM).	B.1.1.529, BA.1, BA.2	Ramanathan et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	Using Mircoscale Thermopheresis, the B.1.351 variant harboring three mutations, binds ACE2 at nearly five-fold greater affinity than the original SARS-COV-2 RBD (Kd 87.6, vs 402.5 nM).	B.1.1.529, BA.1, BA.2	Ramanathan et al. (2021)	862	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.N501Y	ACE2 receptor binding affinity	In silico methods (Py-MOL and PDBePISA) involving mutagenesis (N501Y mutation) and interface analysis focusing on the Spike RDB-ACE2 interaction showed that the SARS-CoV-2 N501Y mutant (lineage B.1.1.7) establishes a more significant number of interactions relating to the mutant residue Y501 (Spike RDB) with residues Y41 and K353 (ACE2). This finding shows that the increased infectivity of SARS-CoV-2 lineage B.1.1.7 is associated with the interaction force between the Spike RBD Y501 mutant residue with the ACE2 receptor, which in this strain is increased.	B.1.1.529, BA.1, BA.2	Santos and Passos (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	Experimentally, ACE2 binding affinity in- creased 0.24 fold	B.1.1.529, BA.1, BA.2	Starr et al. (2020)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	This single mutation causes major increase in binding affinity vs. wild type as measured by IC50 vs pseudotyped lentivirus, but combined with the complete set of B.1.1.7 lineage variants no major change vs wild type affnity is observed.	B.1.1.529, BA.1, BA.2	Tada et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	This mutation combination causes major increase in binding affinity vs. wild type as measured by IC50 vs pseudotyped lentivirus, but combined with the complete set of B.1.1.7 lineage variants no major change vs wild type affnity is observed.	BA.1	Tada et al. (2021)	860	Т	1.0
p.N501Y	ACE2 receptor binding affinity	Reported 4-fold increase in affinity compared to wild-type RBD on the cell surface (Kd	B.1.1.529, BA.1, BA.2	Tian et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	Reported slight increase in affinity compared to wild-type RBD on the cell surface (Kd	B.1.1.529, BA.1, BA.2	Tian et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	The affinity of ACE2 for this mutation combination was twice as high as for wild type. Having in mind that the affinity of SARS-CoV-2 for ACE2 is only 4-fold higher compared to SARS-CoV-1, this factor of 2 is expected to be biologically significant.	B.1.1.529, BA.1, BA.2	Vogel et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	Among the first selected and fixed variants in an in vitro evolution exper- iment for ACE2 bind- ing. Calculated disasso- ciation constant for this variant is nearly four fold lower than wild type (Kd	B.1.1.529, BA.1, BA.2	Zahradnik et al. (2021)	862	Т	1.0
p.N501Y	ACE2 receptor binding affinity	N501Y residue inserts into a cavity at the binding interface near Y41 of ACE2. The additional interactions result in increased affinity of ACE2 for the N501Y mutant, accounting for its increased infectivity.	B.1.1.529, BA.1, BA.2	Zhu et al. (2021)	862	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.N501Y	T cell evasion	Vaccinated, but not post-infection sera, show decreased average T cell response to an N501Y peptide. When we primed transgenic mice expressing human HLA-DRB1*0401 with the Wuhan Hu-1 peptide pool, T cell responses to the B.1.1.7 variant peptide pool were significantly reduced (p	B.1.1.529, BA.1, BA.2	Reynolds et al. (2021)	862	Т	1.0
p.N501Y	antibody epitope effects	Ablates Class 3 N- terminal domain tar- geting antibody COV2- 2489, diminishes COV2- 2676.	B.1.1.529, BA.1, BA.2	Chen et al. (2021)	862	Т	1.0
p.N501Y	antibody epitope effects	Ablates Class 3 N- terminal domain tar- geting antibody COV2- 2489, diminishes COV2- 2676.	BA.1	Chen et al. (2021)	860	Т	1.0
p.N501Y	antibody epitope effects	Of 50 mAbs tested, major loss of neutralization observed for S2X128, S2D8, S2X192, S2D19, S2H14, S2H19.	B.1.1.529, BA.1, BA.2	Collier et al. (2021)	862	Т	1.0
p.N501Y	antibody epitope effects	Wildtype elicits immune response, COVID-19 cohort epitope score > 99th percentile of the 497 pre-pandemic controls, mutant drops PIWAS epitope score from 3% to 1.2% (poorer immune recognition) Together with other B1.1.7 lineage mutational changes (Spike: Y144del, A570D, P681H, Nucleoprotein: D3L, S235F) resulted in only 2 or 579 individuals (0.3% of the population) having a dramatic reduction in PIWAS antigen scores, which reflects the peak epitope signal along the entire antigen.	B.1.1.529, BA.1, BA.2	Haynes et al. (2021)	862	Т	1.0
p.N501Y	antibody epitope effects	Contrary to other reports on N501Y containing lineages (i.e. with additional mutations), N501Y alone may have an even greater affinity for a human monoclonal antibody specific for wild type. These results suggest that the individual N501Y mutation does not contribute to altered viral properties by itself, but may contribute to a collective conformational shift produced by multiple mutations.	B.1.1.529, BA.1, BA.2	Klegerman et al. (2021)	862	Т	1.0
p.N501Y	antibody epitope effects	mutations. Lowered the neutralization potency of mAb COVA1-12 to the limit of the assay. Decrease in potency was observed against the N501Y pseudotype for the cluster IX mAb COVA2-17.	B.1.1.529, BA.1, BA.2	Rees-Spear et al. (2021)	862	Т	1.0
p.N501Y	antibody epitope effects	Reduction in neutralization by mAbs COVA1-18 (extasci- itilde4x), COVA2-15 (extasciitilde9x), S309 (extasciitilde3x)	B.1.1.529, BA.1, BA.2	Shen et al. (2021)	862	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.N501Y	antibody epitope effects	Reduction in neutralization by mAbs COVA1-18 (extasciitilde4x), COVA2-15 (extasciitilde9x). PG: these effects are laregly missing in the deletion-alone data	BA.1	Shen et al. (2021)	860	Т	1.0
p.N501Y	antibody epitope effects	4 antibodies tested were less potent against K417N by ten-fold or more, in both mAb classes 1 and 3	B.1.1.529, BA.1, BA.2	Wang et al. (2021)	862	Т	1.0
p.N501Y	convalescent plasma binding	1.65x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	Т	1.0
p.N501Y	convalescent plasma escape	One convalescent sera tested showed 4-fold or greater reduction in neu- tralization efficiency.	BA.1	Alenquer et al. (2021)	860	Т	1.0
p.N501Y	convalescent plasma escape	Slight neutralization improvement on average in 16 health workers' convalescent sera.	BA.1	Alenquer et al. (2021)	860	Т	1.0
p.N501Y	convalescent plasma escape	The 501Y.V2 to first wave IC50 ratio ranged from 6 to 200-fold. Averaging across all 7 participant convalescent sera highlighted the dramatic decrease in sensitivity to neutralization of authentic 501Y.V2 variants. PG: I'm purposefully ignoring D614G and A701V as contributors	B.1.1.529, BA.1, BA.2	Cele et al. (2021)	862	Т	1.0
p.N501Y	convalescent plasma escape	0.7x reduction in neutralization by key variant in several variants of concern in sera collected from cohort of 10 with severe disease 21 to 63 days post-onset.	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	862	Т	1.0
p.N501Y	convalescent plasma escape	In 30 samples collected 111 to 260 days post onset of symptoms, the covalescent plasma can neutralize both the reference USA-WA1/2020 strain and the mouse adapted strain that contains the N501Y spike mutation with similar efficiency.	B.1.1.529, BA.1, BA.2	Rathnasinghe et al. (2021)	862	Т	1.0
p.N501Y	convalescent plasma escape	Neutralization activity of convalescent sera tested decreased extasci- itilde2x with this B.1.1.7 pseudotyped virus.	B.1.1.529, BA.1, BA.2	Shen et al. (2021)	862	Т	1.0
p.N501Y	convalescent plasma escape	Viruses containing the point mutations of B.1.1.7 showed that the single point mutations (\Delta 69-70 and N501Y) were neutralized as efficiently as D614G across 10 convalescent sera from April 2020 infectees.	B.1.1.529, BA.1, BA.2	Tada et al. (2021)	862	Т	1.0
p.N501Y	convalescent plasma escape	These key B.1.1.7 mutations as a combination neutralized slightly less well than D614G and this was noticeable in the lack of sera with high neutralizing titer for the viruses across 10 convalescent sera from April 2020 infectees.	BA.1	Tada et al. (2021)	860	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.N501Y	convalescent plasma escape	As measured by surface plasmon resonance, RBD with the N501Y mutation alone showed a mean 2.1x decrease in binding affinity for six batches of hyperimmune immunoglobulin (hCoV-2IG) preparations generated from SARS-CoV-2 convalescent plasma.	B.1.1.529, BA.1, BA.2	Tang et al. (2021)	862	Т	1.0
p.N501Y	convalescent plasma escape	27% of 44 early pandemic exposure convalescent plasma/sera lose all activity against a RBD triple mutant pseudovirus (RBD mutatants of the 501Y.V2 "South African" lineage), while only 23% retained high titres	B.1.1.529, BA.1, BA.2	Wibmer et al. (2021)	862	Т	1.0
p.N501Y	convalescent plasma escape	Nearly half (21 of 44, 48%) of early pandemic exposure convalescent plasma/sera failed to neutralize the 501Y.V2 ("South African") lineage pseudovirus construct Only 3 of 44 convascent sera (those with the highest titer, which correlated directly with initial infection severity) had high neutralization against this 501Y.V2 PG: note that lineage variant R246I was excluded from the text in reference to these sera assays, not sure if that was an oversight.	B.1.1.529, BA.1, BA.2	Wibmer et al. (2021)	862	T	1.0
p.N501Y	environmental condition stability	Relative to D614G, this mutation demonstrated significant increase in infectivity (i.e. heat stability) after incubation at 50C after 30 minutes or 1 hour	B.1.1.529, BA.1, BA.2	Tada et al. (2021)	862	Т	1.0
p.N501Y	environmental condition stability	Relative to D614G, this mutation demonstrated significant increase in infectivity (i.e. heat stability) after incubation at 50C after 1 hour.	BA.1	Tada et al. (2021)	860	Т	1.0
p.N501Y	homoplasy	Variant within the six key residues in the receptor binding domain (RBD). Independently reported in UK, Australia (same origin as UK), and South Africa (independent origin).	B.1.1.529, BA.1, BA.2	Flores-Alanis et al. (2021)	862	T	1.0
p.N501Y	immunosuppression variant emergence	Appeared (day 128) and persisted in chronic (152 day) SARS-CoV-2 in- fection of immunocom- promised patient with severe antiphospholipid syndrome	B.1.1.529, BA.1, BA.2	Choi et al. (2020)	862	Т	1.0
p.N501Y	monoclonal anti- body serial passage escape	In vitro selection against class 1 (Spike 'up' conformation) monoclonal antibody C663, and to a lesser extent C613.	B.1.1.529, BA.1, BA.2	Wang et al. (2021)	862	Т	1.0
p.N501Y	pharmaceutical effectiveness	COR-101 lost extasci- itilde8x binding against this isolated mutation. Regdanvimab lost ex- tasciitilde6x binding against this isolated mutation.	B.1.1.529, BA.1, BA.2	Engelhart et al. (2021)	862	Т	1.0
p.N501Y	pharmaceutical effectiveness	Tixagevimab, Regdan- vimab and COR-101 display reduced binding affinity to virus pseu- dotyped as RBD from B.1.351.	B.1.1.529, BA.1, BA.2	Engelhart et al. (2021)	862	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.N501Y	pharmaceutical effectiveness	COR-101 lost extasci- itilde20x binding against this double mutation. Estesevimab lost ex- tasciitilde16x binding against this double mu- tation. Regdanvimab lost extasciitilde6x bind- ing against this double mutation. M396 lost extasciitilde10x bind- ing against this double mutation.	B.1.1.529, BA.1, BA.2	Engelhart et al. (2021)	862	Т	1.0
p.N501Y	pharmaceutical effectiveness	This mutated version of RBD completely abol- ishes the binding to a therapeutic antibody, Bamlanivimab, in vitro.	B.1.1.529, BA.1, BA.2	Liu et al. (2021)	862	Т	1.0
p.N501Y	symptom prevalence	A higher proportion of cases infected with the B.1.1.7 variant were hypoxic on admission compared to other variants (70.0% vs 62.5%, p	BA.1	Snell et al. (2021)	860	Т	1.0
p.N501Y	symptom prevalence	In comparison of B.1.1.7 lineage (193 cases) vs. "wildtype" (125) in Berlin Jan 18 to March 29 2021, significant symptom changes are ab- sent loss of smell/taste (P	BA.1	van Loon et al. (2021)	860	Т	1.0
p.N501Y	syncytium formation	Slight increase in Vero cell-cell membrane fusion assay under infection with VSV pseudotyped virus relative to wild type, no change relative to D614G.	B.1.1.529, BA.1, BA.2	Kim et al. (2021)	862	Т	1.0
p.N501Y	trafficking	More efficient infectivity (24h) compared to wild type, in Caco-2 cells extasciitilde9x, Vero extasciitilde8x, and Calu-3 extasciitilde8x. Compare to wild type at extasciitilde5x across cell types.	B.1.1.529, BA.1, BA.2	Kim et al. (2021)	862	Т	1.0
p.N501Y	trafficking	extasciitilde4x more efficient S2 domain cleavage compared to wild type, no change relative to D614G alone in Caco-2 cells, mid-range of three cell line tested (Vero and Calu-3).	B.1.1.529, BA.1, BA.2	Kim et al. (2021)	862	Т	1.0
p.N501Y	trafficking	9x more infectivity than D614G alone in HEK293T-ACE2 cells 48h post-transduction.	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	863	Т	1.0
p.N501Y	trafficking	extasciitilde9x more infectivity than D614G alone in HEK293T-ACE2 cells 48h post-transduction (no synergy as level approx. that of N501Y alone).	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	862	Т	1.0
p.N501Y	trafficking	Decreased stability of RBD expression in yeast, suggesting de- creased Spike protein stability.	B.1.1.529, BA.1, BA.2	Motozono et al. (2021)	862	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.N501Y	trafficking	Lentiviral pseudotyped with this individual mutation from B.1.1.7 was tested on ACE2.293T cells. Luciferase activity was measured two days postinfection, showing slightly increased infection rate amongst the cells. [in what is essentially a replicate experiment in the same paper, because each B.1.351 lineage variant was independetly evaluated and N501 is in both lineages, a significant decrease was observed, therefore the error bars described in this paper should be interpreted carefully	B.1.1.529, BA.1, BA.2	Tada et al. (2021)	862	Т	1.0
p.N501Y	trafficking	Lentiviral pseudotyped with this mutation set from B.1.1.7 was tested on ACE2.293T cells. Luciferase activity was measured two days postinfection, showing significant (40%) increase in infection rate amongst the cells, much more than the effect of either the deletion or the point mutation alone, suggetsing that this combination has a synergistic effect contributing to cell entry fitness, moreso than this combination with the addition of P681H.	BA.1	Tada et al. (2021)	860	Т	1.0
p.N501Y	trafficking	Lentiviral pseudotyped with this mutation set from B.1.1.7 was tested on ACE2.293T cells. Luciferase activity was measured two days postinfection, showing significant (40%) increase in infection rate amongst the cells, much more than the effect of either the deletion or the point mutation alone, suggetsing that this combination has a synergistic effect contributing to cell entry fitness, but to a smaller extent than N501Y and the deletion alone.	BA.1	Tada et al. (2021)	860	T	1.0
p.N501Y	vaccine neutraliza- tion efficacy	Observed 1.3-fold reduction in neutralization efficiency of Pfizer vaccinee sera (collected 14 days after second dose) against pseudotype B.1.1.7 key variant lentivirus. Compare to 2.6-fold reduction against cultured B.1.1.7 virus.	B.1.1.529, BA.1, BA.2	Bates et al. (2021)	862	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.N501Y	vaccine neutralization efficacy	1.2x drop in neutralization using sera collected from 14 healthy adult participants that received two injections of the mRNA-1273 (Moderna) vaccine at a dose of 100 µg (18-55 years: day 1 and day 14 post-2nd dose) against a recombinant single variant virus (modified replicating WA-1 cDNA clone) relative to contemporary circulating D614G variant (USA/GA-EHC-083E/2020) using a live-virus Focus Reduction Neutralization Test (FRNT) assay.	B.1.1.529, BA.1, BA.2	Edara et al. (2021)	862	Т	1.0
p.N501Y	vaccine neutralization efficacy	The presence of this variant in 189 post-mRNA-vaccination COVID-19 cases was proportionally in line with lineage prevalence in Northen California during the study period, suggesting no effect of these variants on immune escape.	B.1.1.529, BA.1, BA.2	Jacobson et al. (2021)	862	Т	1.0
p.N501Y	vaccine neutraliza- tion efficacy	This variant showed no change in Pfizer sera (one or two dose) neutralization efficiency vs D614G (using lentivirus pseudotype).	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	862	Т	1.0
p.N501Y	vaccine neutraliza- tion efficacy	This variant showed >5x decrease in Pfizer sera (3 weeks post-first dose: n	B.1.1.529, BA.1, BA.2	Kuzmina et al. (2021)	862	Т	1.0
p.N501Y	vaccine neutralization efficacy	Human sera from 6 two-dose Pfizer vaccinated individuals (47-68 days post 1st-dose) can neutralize both the reference USA-WA1/2020 strain and the mouse adapted SARS-CoV-2 strain that contains the N501Y spike mutation with similar efficiency.	B.1.1.529, BA.1, BA.2	Rathnasinghe et al. (2021)	862	Т	1.0
p.N501Y	vaccine neutraliza- tion efficacy	In a cohort of 20 patients 8+ weeks after second vaccine dose of Moderna (mRNA-1273) or Pfizer- BioNTech (BNT162b2) vaccines, a modest de- crease in neutralization by vaccine plasma was observed.	B.1.1.529, BA.1, BA.2	Wang et al. (2021)	862	Т	1.0
p.N501Y	vaccinee plasma binding	1.17x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously non-seroconverted subjects. 1.09x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	Т	1.0
p.N501Y	virion structure	Estimated free energy change (ddG) for this variant is 0.69 kcal/mol (i.e. stabilizing relative to wild type)	B.1.1.529, BA.1, BA.2	Spratt et al. (2021)	862	Т	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.N501Y	virion structure	The Ratio of S2 (processed Spike) to full length Spike is higher for this mutation combination, due to a drop in the full length Spike measured, suggesting that this mutation compensates for decreased Spike production by improved proteolytic processing.	BA.1	Tada et al. (2021)	860	Т	1.0
p.N501Y	virion structure	The Ratio of S2 (processed Spike) to full length Spike is higher for this mutation combination, due to a drop in the full length Spike measured, suggesting that this mutation compensates for decreased Spike production by improved proteolytic processing.	BA.1	Tada et al. (2021)	860	Т	1.0
p.T95I	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant showed a 1.33x decrease in binding (KD) relative to D614G.	B.1.1.529, BA.1	Gong et al. (2021)	858	Т	1.0
p.T95I	convalescent plasma binding	No change in Spike binding (relative to D614G alone) by 5 plasma collected 8 months postsymptom-onset.	B.1.1.529, BA.1	Gong et al. (2021)	858	Т	1.0
p.T95I	vaccinee plasma binding	1.16x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously non-seroconverted subjects. 1.02x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	B.1.1.529, BA.1	Gong et al. (2021)	858	T	1.0
p.E484A	antibody epitope effects	Mutant screen in neutralization assay with a broad range of monoclonal antibodies shows high resistence to 4 antibodies, and broad low level resistence against much of the rest of the panel.	B.1.1.529, BA.1, BA.2	Liu et al. (2020)	750	C	1.0
p.E484A	convalescent plasma escape	Remarkably, several of the E484 escape mutants were resistant to neutralization at the highest concentration (1:80 initial dilution) of all 4 convalescent sera tested. Against a wider panel of 16 convalescent plasma (no replicates), all but one show major resistance.	B.1.1.529, BA.1, BA.2	Liu et al. (2021)	750	С	1.0
p.E484A	monoclonal anti- body serial passage escape	Ranked moderately effective mutant against this position in the RBD for highly neutralizing COV2-2050 monoclonal antibody	B.1.1.529, BA.1, BA.2	Greaney et al. (2020)	750	С	1.0
p.E484A	monoclonal anti- body serial passage escape	"E484A rose in frequency in linkage with F486I, but since E484A is not an escape mutation in our maps it is not	B.1.1.529, BA.1, BA.2	Starr et al. (2020)	750	С	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.E484A	monoclonal anti- body serial passage escape	Escape mutation against monoclonal antibody LY-CoV555 (antibody that forms the basis for Eli Lilly's bamlanivimab)	B.1.1.529, BA.1, BA.2	Starr et al. (2021)	75Ô	C	1.0
p.E484A	pharmaceutical effectiveness	Bamlanivimab (LY- CoV555) lost extasci- itilde8x binding against this isolated mutation.	B.1.1.529, BA.1, BA.2	Engelhart et al. (2021)	750	С	1.0
p.V70del	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant showed a 1.51x increase in binding (KD) relative to D614G, mostly due to decreased in "off-rate" a.k.a. dissociation rate (Kdis).	BA.1	Gong et al. (2021)	855	ТТА	1.0
p.V70del	ACE2 receptor binding affinity	This mutation combination causes major increase in binding affinity vs. wild type as measured by IC50 vs pseudotyped lentivirus, but combined with the complete set of B.1.1.7 lineage variants no major change vs wild type affnity is observed.	BA.1	Tada et al. (2021)	855	TTA	1.0
p.V70del	antibody epitope effects	Ablates Class 3 N- terminal domain tar- geting antibody COV2- 2489, diminishes COV2- 2676.	BA.1	Chen et al. (2021)	855	TTA	1.0
p.V70del	antibody epitope effects	Reduces neutralization by structurally un- mapped mAb COVA1-21 (cluster XI).	BA.1	Rees-Spear et al. (2021)	855	TTA	1.0
p.V70del	antibody epitope effects	Reduction in neutralization by mAbs COVA1-18 (extasciitilde4x), COVA2-15 (extasciitilde9x). PG: these effects are laregly missing in the deletion-alone data	BA.1	Shen et al. (2021)	855	TTA	1.0
p.V70del	convalescent plasma binding	1.33x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	BA.1	Gong et al. (2021)	855	TTA	1.0
p.V70del	convalescent plasma escape	One convalescent sera tested showed 4-fold or greater reduction in neu- tralization efficiency.	BA.1	Alenquer et al. (2021)	855	TTA	1.0
p.V70del	convalescent plasma escape	Slight neutralization improvement on average in 16 health workers' convalescent sera.	BA.1	Alenquer et al. (2021)	855	TTA	1.0
p.V70del	convalescent plasma escape	Fatal COVID-19 compli- cations in immunocom- primised patient after immune escape from con- valescent plasma	BA.1	Kemp et al. (2020)	855	TTA	1.0
p.V70del	convalescent plasma escape	Neutralization activity of almost all Moderna Phase 1 sera tested actually *increased*.	BA.1	Shen et al. (2021)	855	TTA	1.0
p.V70del	convalescent plasma escape	Viruses containing the point mutations of B.1.1.7 showed that the single point mutations $(\Delta 69\text{-}70 \text{ and } \text{N}501\text{Y})$ were neutralized as efficiently as D614G across 10 convalescent sera from April 2020 infectees.	BA.1	Tada et al. (2021)	855	ТТА	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.V70del	convalescent plasma escape	These key B.1.1.7 mutations as a combination neutralized slightly less well than D614G and this was noticeable in the lack of sera with high neutralizing titer for the viruses across 10 convalescent sera from April 2020 infectees.	BA.1	Tada et al. (2021)	855	TTA	1.0
p.V70del	environmental condition stability	Relative to D614G, this mutation demonstrated significant increase in infectivity (i.e. heat stability) after incubation at 50C after 1 hour.	BA.1	Tada et al. (2021)	855	TTA	1.0
p.V70del	immunosuppression variant emergence	The delH69/V70 enhances viral infectivity, indicating its effect on virus fitness is independent to the N501Y RBM change [with which it is found in lineage B.1.1.7] Possibly arisen as a result of the virus evolving from immune selection pressure in infected individuals and possibly only one chronic infection in the case of lineage B.1.1.7.	BA.1	Kemp et al. (2020)	855	TTA	1.0
p.V70del	symptom prevalence	A higher proportion of cases infected with the B.1.1.7 variant were hypoxic on admission compared to other variants (70.0% vs 62.5%, p	BA.1	Snell et al. (2021)	855	TTA	1.0
p.V70del	symptom prevalence	In comparison of B.1.1.7 lineage (193 cases) vs. "wildtype" (125) in Berlin Jan 18 to March 29 2021, significant symptom changes are ab- sent loss of smell/taste (P	BA.1	van Loon et al. (2021)	855	TTA	1.0
p.V70del	trafficking	Lentiviral pseudotyped with this mutation set from B.1.1.7 was tested on ACE2.293T cells. Luciferase activity was measured two days postinfection, showing significant (40%) increase in infection rate amongst the cells, much more than the effect of either the deletion or the point mutation alone, suggetsing that this combination has a synergistic effect contributing to cell entry fitness, moreso than this combination with the addition of P681H.	BA.1	Tada et al. (2021)	855	TTA	1.0
p.V70del	trafficking	Lentiviral pseudotyped with this mutation set from B.1.1.7 was tested on ACE2.293T cells. Luciferase activity was measured two days postinfection, showing significant (40%) increase in infection rate amongst the cells, much more than the effect of either the deletion or the point mutation alone, suggetsing that this combination has a synergistic effect contributing to cell entry fitness, but to a smaller extent than N501Y and the deletion alone.	BA.1	Tada et al. (2021)	855	TTA	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.V70del	vaccinee plasma binding	1.14x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously non-seroconverted subjects. 1.09x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	BA.1	Gong et al. (2021)	855	TTA	1.0
p.V70del	virion structure	The Ratio of S2 (processed Spike) to full length Spike is higher for this mutation combination, due to a drop in the full length Spike measured, suggesting that this mutation compensates for decreased Spike production by improved proteolytic processing.	BA.1	Tada et al. (2021)	855	TTA	1.0
p.V70del	virion structure	The Ratio of S2 (processed Spike) to full length Spike is higher for this mutation combination, due to a drop in the full length Spike measured, suggesting that this mutation compensates for decreased Spike production by improved proteolytic processing.	BA.1	Tada et al. (2021)	855	TTA	1.0
p.H69del	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant showed a 1.51x increase in binding (KD) relative to D614G, mostly due to decreased in "off-rate" a.k.a. disso-	BA.1	Gong et al. (2021)	1710	TTA	1.0
p.H69del	ACE2 receptor binding affinity	ciation rate (Kdis). This mutation combination causes major increase in binding affinity vs. wild type as measured by IC50 vs pseudotyped lentivirus, but combined with the complete set of B.1.1.7 lineage variants no major change vs wild type affnity is observed.	BA.1	Tada et al. (2021)	1710	TTA	1.0
p.H69del	antibody epitope effects	Ablates Class 3 N- terminal domain tar- geting antibody COV2- 2489, diminishes COV2- 2676.	BA.1	Chen et al. (2021)	1710	TTA	1.0
p.H69del	antibody epitope effects	Reduces neutralization by structurally un- mapped mAb COVA1-21 (cluster XI).	BA.1	Rees-Spear et al. (2021)	1710	TTA	1.0
p.H69del	antibody epitope effects	Reduction in neutralization by mAbs COVA1- 18 (extasciitilde4x), COVA2-15 (extasciitilde9x). PG: these effects are laregly missing in the deletion-alone data	BA.1	Shen et al. (2021)	1710	TTA	1.0
p.H69del	convalescent plasma binding	1.33x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	BA.1	Gong et al. (2021)	1710	TTA	1.0
p.H69del	convalescent plasma escape	One convalescent sera tested showed 4-fold or greater reduction in neu- tralization efficiency.	BA.1	Alenquer et al. (2021)	1710	TTA	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.H69del	convalescent plasma escape	Slight neutralization improvement on average in 16 health workers' convalescent sera.	BA.1	Alenquer et al. (2021)	1710	TTA	1.0
p.H69del	convalescent plasma escape	Fatal COVID-19 compli- cations in immunocom- primised patient after immune escape from con- valescent plasma	BA.1	Kemp et al. (2020)	1710	TTA	1.0
p.H69del	convalescent plasma escape	Neutralization activity of almost all Moderna Phase 1 sera tested actually *increased*.	BA.1	Shen et al. (2021)	1710	TTA	1.0
p.H69del	convalescent plasma escape	Viruses containing the point mutations of B.1.1.7 showed that the single point mutations $(\Delta 69-70 \text{ and } N501Y)$ were neutralized as efficiently as D614G across 10 convalescent sera from April 2020 infectees.	BA.1	Tada et al. (2021)	1710	TTA	1.0
p.H69del	convalescent plasma escape	These key B.1.1.7 mutations as a combination neutralized slightly less well than D614G and this was noticeable in the lack of sera with high neutralizing titer for the viruses across 10 convalescent sera from April 2020 infectees.	BA.1	Tada et al. (2021)	1710	TTA	1.0
p.H69del	environmental condition stability	Relative to D614G, this mutation demonstrated significant increase in infectivity (i.e. heat stability) after incubation at 50C after 1 hour.	BA.1	Tada et al. (2021)	1710	TTA	1.0
p.H69del	immunosuppression variant emergence	The delH69/V70 enhances viral infectivity, indicating its effect on virus fitness is independent to the N501Y RBM change [with which it is found in lineage B.1.1.7] Possibly arisen as a result of the virus evolving from immune selection pressure in infected individuals and possibly only one chronic infection in the case of lineage B.1.1.7.	BA.1	Kemp et al. (2020)	1710	TTA	1.0
p.H69del	symptom prevalence	A higher proportion of cases infected with the B.1.1.7 variant were hypoxic on admission compared to other variants (70.0% vs 62.5%, p	BA.1	Snell et al. (2021)	1710	TTA	1.0
p.H69del	symptom prevalence	In comparison of B.1.1.7 lineage (193 cases) vs. "wildtype" (125) in Berlin Jan 18 to March 29 2021, significant symptom changes are ab- sent loss of smell/taste (P	BA.1	van Loon et al. (2021)	1710	TTA	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.H69del	trafficking	Lentiviral pseudotyped with this mutation set from B.1.1.7 was tested on ACE2.293T cells. Luciferase activity was measured two days postinfection, showing significant (40%) increase in infection rate amongst the cells, much more than the effect of either the deletion or the point mutation alone, suggetsing that this combination has a synergistic effect contributing to cell entry fitness, moreso than this combination with the	BA.1	Tada et al. (2021)	1710	TTA	1.0
p.H69del	trafficking	addition of P681H. Lentiviral pseudotyped with this mutation set from B.1.1.7 was tested on ACE2.293T cells. Luciferase activity was measured two days postinfection, showing significant (40%) increase in infection rate amongst the cells, much more than the effect of either the deletion or the point mutation alone, suggetsing that this combination has a synergistic effect contributing to cell entry fitness, but to a smaller extent than N501Y and the deletion alone.	BA.1	Tada et al. (2021)	1710	TTA	1.0
p.H69del	vaccinee plasma binding	1.14x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously nonseroconverted subjects. 1.09x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in	BA.1	Gong et al. (2021)	1710	ТТА	1.0
p.H69del	virion structure	post-infection vaccinees. The Ratio of S2 (processed Spike) to full length Spike is higher for this mutation combination, due to a drop in the full length Spike measured, suggesting that this mutation compensates for decreased Spike production by improved proteolytic processing.	BA.1	Tada et al. (2021)	1710	TTA	1.0
p.H69del	virion structure	The Ratio of S2 (processed Spike) to full length Spike is higher for this mutation combination, due to a drop in the full length Spike measured, suggesting that this mutation compensates for decreased Spike production by improved proteolytic processing.	BA.1	Tada et al. (2021)	1710	TTA	1.0
p.P681H	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant showed a 1.23x decrease in binding (KD) relative to D614G.	B.1.1.529, BA.1	Gong et al. (2021)	858	A	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.P681H	antibody epitope effects	Ablates Class 3 N- terminal domain tar- geting antibody COV2- 2489, diminishes COV2- 2676.	B.1.1.529, BA.1	Chen et al. (2021)	858	A	1.0
p.P681H	antibody epitope effects	Wildtype elicits immune response, COVID-19 cohort epitope score > 99th percentile of the 497 pre-pandemic controls, mutant drops PIWAS epitope score from 7.8% to 1.2% (significantly poorer immune recognition) Together with other B1.1.7 lineage mutational changes (Spike: Y144del,N501Y, A570D Nucleoprotein: D3L, S235F) resulted in only 2 of 579 individuals (0.3% of the population) having a dramatic reduction in PIWAS antigen scores, which reflects the peak epitope signal along the entire antigen.	B.1.1.529, BA.1	Haynes et al. (2021)	858	A	1.0
р.Р681Н	antibody epitope effects	This variant is adjacent to the Spike protein furin cleavage site (cleavage of S into S1 and S2 subunits is required for viral membrane fusion and subsequent entry into host cells), a site shown to be highly immunogenic.	B.1.1.529, BA.1	Johnson et al. (2020)	858	A	1.0
р.Р681Н	convalescent plasma binding	1.26x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	B.1.1.529, BA.1	Gong et al. (2021)	858	A	1.0
p.P681H	convalescent plasma escape	One convalescent sera tested showed 4-fold or greater reduction in neu- tralization efficiency.	BA.1	Alenquer et al. (2021)	857	A	1.0
p.P681H	convalescent plasma escape	Slight neutralization improvement on average in 16 health workers' convalescent sera.	BA.1	Alenquer et al. (2021)	857	A	1.0
p.P681H	convalescent plasma escape	These key B.1.1.7 mutations as a combination neutralized slightly less well than D614G and this was noticeable in the lack of sera with high neutralizing titer for the viruses across 10 convalescent sera from April 2020 infectees.	BA.1	Tada et al. (2021)	857	A	1.0
р.Р681Н	environmental con- dition stability	Relative to D614G, this mutation demonstrated significant increase in infectivity (i.e. heat stability) after incubation at 50C after 1 hour.	BA.1	Tada et al. (2021)	857	A	1.0
р.Р681Н	trafficking	While the introduction of P681H in the SARS-CoV-2 B.1.1.7 variant may increase spike cleavage by furin-like proteases, this does not significantly impact viral entry or cell-cell spread. We consider that other factors are at play to account for the increased in transmission and disease severity attributed to this variant of concern (VOC).	B.1.1.529, BA.1	Lubinski et al. (2021)	858	A	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
р.Р681Н	trafficking	This mutation in the first base of the furin clevage site maintains the RXXR recognition motif, and is presumed to enhance cleavage based on the removal of a proline-directed phosphotase recognition site at S680. In a homologuous site in Infectious Bronchitis Virus (IBV, Gammacoronaviruses), abolition of S680 phosphorylation improves furin cleavage (and presumably cell entry).	B.1.1.529, BA.1	Maaroufi (2021)	858	A	1.0
p.P681H	trafficking	Lentiviral pseudotyped with this individual mutation from B.1.1.7 was tested on ACE2.293T cells. Luciferase activity was measured two days postinfection, showing NO statistically significant infection rate change amongst the cells, suggesting that furin cleavage typically used for cell entry is not affected by this change one amino acid upstream of the RXXR recognition pattern.	B.1.1.529, BA.1	Tada et al. (2021)	858	A	1.0
p.P681H	trafficking	Lentiviral pseudotyped with this mutation set from B.1.1.7 was tested on ACE2.293T cells. Luciferase activity was measured two days postinfection, showing significant (40%) increase in infection rate amongst the cells, much more than the effect of either the deletion or the point mutation alone, suggetsing that this combination has a synergistic effect contributing to cell entry fitness, but to a smaller extent than N501Y and the deletion alone.	BA.1	Tada et al. (2021)	857	A	1.0
p.P681H	vaccine neutraliza- tion efficacy	No significant change in virus neutralzation by 18 Pfizer two dose vaccinee sera compared to B.1.1.7. [results without including the used mutation A27S likely generalizable, as this is not a lineage defining mutation]	B.1.1.529, BA.1	Zuckerman et al. (2021)	858	A	1.0
p.P681H	vaccinee plasma binding	1.14x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously non-seroconverted subjects. 1.11x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	B.1.1.529, BA.1	Gong et al. (2021)	858	A	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
р.Р681Н	virion structure	The Ratio of S2 (processed Spike) to full length Spike is higher for this mutation, due to a drop in the full length Spike measured, suggesting that this mutation compensates for decreased Spike production by improved proteolytic processing.	B.1.1.529, BA.1	Tada et al. (2021)	858	A	1.0
р.Р681Н	virion structure	The Ratio of S2 (processed Spike) to full length Spike is higher for this mutation combination, due to a drop in the full length Spike measured, suggesting that this mutation compensates for decreased Spike production by improved proteolytic processing.	BA.1	Tada et al. (2021)	857	A	1.0
p.G339D	ACE2 receptor binding affinity	Experimentally, ACE2 binding affinity in- creased 0.06 fold	B.1.1.529, BA.1, BA.2	Starr et al. (2020)	863	A	1.0
p.G339D	gene expression in- crease	Experimentally, Spike gene expression in- creased 0.3 fold	B.1.1.529, BA.1, BA.2	Starr et al. (2020)	863	A	1.0
p.H655Y	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant showed a 1.21x increase in binding (KD) relative to D614G.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	Т	1.0
p.H655Y	anthropozoonotic events	Six minks were in- tranasally infected with WA1 isolate, all de- veloped this mutation during infection.	B.1.1.529, BA.1, BA.2	Esclera et al. (2021)	862	Т	1.0
p.H655Y	convalescent plasma binding	1.48x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	Т	1.0
p.H655Y	homoplasy	In experimental models of SARS-CoV-2 mutational evolution (without immune pressure), this mutation in the N terminal domain appears convergent.	B.1.1.529, BA.1, BA.2	Borges et al. (2021)	862	T	1.0
р.Н655Ү	vaccinee plasma binding	1.04x increase in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in previously non-seroconverted subjects. 1.09x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 3 weeks after one dose of Pfizer/BioNtech BNT162b2 vaccine in post-infection vaccinees.	B.1.1.529, BA.1, BA.2	Gong et al. (2021)	862	Т	1.0
p.H655Y	virion structure	Estimated free energy change (ddG) for this variant is 0.87 kcal/mol (i.e. stabilizing relative to wild type)	B.1.1.529, BA.1, BA.2	Spratt et al. (2021)	862	Т	1.0
p.N440K	ACE2 receptor binding affinity	Experimentally, ACE2 binding affinity in- creased 0.07 fold	B.1.1.529, BA.1, BA.2	Starr et al. (2020)	749	G	1.0
p.N440K	antibody epitope effects	N501Y substitution decreased the neutralizing and binding activities of CB6 and increased that of BD-23	B.1.1.529, BA.1, BA.2	Cheng et al. (2021)	749	G	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.N440K	antibody epitope effects	Ablates binding by class 3 mAbs such as C135 that do not directly interfere with ACE2 binding, but clonal somatic mutations of memory B cells at 6.2 months (evolving humoral immune response) show pronounced increase in binding to the variant.	B.1.1.529, BA.1, BA.2	Gaebler et al. (2021)	749	G	1.0
p.N440K	antibody epitope effects	Greater than 10-fold reduction of binding effeiency vs wild type for mAb LY-CoV555. Abolishes binding of mAb ADG-1.	B.1.1.529, BA.1, BA.2	Rappazzo et al. (2021)	749	G	1.0
p.N440K	antibody epitope effects	Resistent to class 3 antibodies (i.e. Abs that do not directly interfere with ACE2 binding).	B.1.1.529, BA.1, BA.2	Wang et al. (2021)	749	G	1.0
p.N440K	monoclonal anti- body serial passage escape	Class 3 antibody C669 mildly selected for the emergence of the N440K mutation in vitro (in contrast to N440H which caused mild escape in Class 1/2 mAb C653).	B.1.1.529, BA.1, BA.2	Wang et al. (2021)	749	G	1.0
p.N440K	monoclonal anti- body serial passage escape	Positive selection (up to 45% of supernatant sequences) under two rounds of C135 monoclonal antibody passage, eliminated in subsequent passages	B.1.1.529, BA.1, BA.2	Weisblum et al. (2020)	749	G	1.0
p.N440K	reinfection	A 47yo Indian male was reinfected with B.1.36 lineage virus in September 2020 after infection with genetically distinct B.1.36 virus in July, with negative PCR tests in between. While the forst episode was asymptomatic, the second included fever, cough, and malaise. The second case additionally contained stopgain ORF3a:E261*	B.1.1.529, BA.1, BA.2	Rani et al. (2021)	749	G	1.0
p.N440K	transmissibility	The N440K variant produced ten times higher infectious viral titers than a prevalent A2a strain, and over 1000 folds higher titers than a much less prevalent A3i strain prototype in Caco2 cells. Interestingly, A3i strain showed the highest viral RNA levels, but the lowest infectious titers in the culture supernatants, indicating the absence of correlation between the RNA content and the infectivity of the sample.	B.1.1.529, BA.1, BA.2	Tandel et al. (2021)	749	G	1.0
p.R346K	gene expression increase	Experimentally, Spike gene expression increased 0.12 fold	B.1.1.529, BA.1	Starr et al. (2020)	860	A	0.29
p.R346K	monoclonal anti- body serial passage escape	Strong positive selection (up to 53% of super- natant sequences) under two rounds of C135 mon- oclonal antibody pas- sage, overall 70% switch away from R346 to S, K or M	B.1.1.529, BA.1	Weisblum et al. (2020)	860	A	0.29
p.G142D	monoclonal anti- body serial passage escape	Escape mutation against Spike N terminal do- main antigenic supersite i mAbs S2M28, S2X28, S2X333	B.1.1.529, BA.2	McCallum et al. (2021)	2	A	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence	Alternate	Alternate
					Depth	Allele	Frequency
p.G142D	monoclonal anti-	Selected twice in passage	B.1.1.529,	Suryadevara et	2	A	1.0
	body serial passage	with mAb COV2-2489.	BA.2	al. (2021)			
	escape						



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