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Surveillance report

Surveillance generated by nf-ncov-voc for Kappa variant

Date

This report is generated on 2022-01-18 using 162696 number of genomes collected between 2020-02-25 and 2021-12-20

Pango Lineages

Pango Lineages in this report ['B.1.617.1']

Indicator

This table contains key indicators identified

Indicator	Sub-categories from POKAY	Mutations
Transmissibility between hu-	transmissibility	p.D614G, p.E484Q, p.L452R,
mans		p.P681R
Infection Severity	ACE2 receptor binding affinity,	p.D614G, p.E154K, p.E484Q,
	viral load, outcome hazard ratio	p.G142D, p.L452R, p.P681R,
		p.Q1071H, p.T95I
Immunity after natural infection	convalescent plasma escape, rein-	p.D614G, p.E154K, p.E484Q,
	fection, humoral response dura-	p.G142D, p.L452R, p.P681R,
	bility	p.Q1071H
Vaccines	vaccine neutralization efficacy	p.D614G, p.E154K, p.E484Q,
		p.G142D, p.L452R, p.P681R,
		p.Q1071H, p.T95I
Monoclonal antibodies	monoclonal antibody serial pas-	p.E484Q, p.G142D, p.L452R
	sage escape, pharmaceutical ef-	
	fectiveness	
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 Depth	Alternate Allele	Alternate Frequency
p.Q1071H	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant combination (representing lineage B.1.617.1) showed a 1.12x increase in binding (KD) relative to D614G, but described as "not significant". [exact variant list not provided in manuscript, is inferred and represents minimal set agreed upon commonly]	B.1.617.1	Gong et al. (2021)	419	419	1.0
p.Q1071H	convalescent plasma escape	Relative to B.1, Kappa (B.1.617.1) shows 5.71x decrease in neutralization efficiency by convalescent plasma [no cohort details provided]	B.1.617.1	Wilhelm et al. (2021)	419	419	1.0
p.Q1071H	trafficking	extasciitilde3.5x cleavage of S2 relative to WA1 (D614G) wildtype by Kappa (B.1.617.1) variant as measured by mass spectrometry of Vero-TMPRSS culture (compare to extasciitilde4.5x for closely related Delta B.1.617.2 also with P681R at the cleavage site).	B.1.617.1	Esclera et al. (2021)	419	419	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.Q1071H	vaccine neutraliza- tion efficacy	Inferring from two B.1.617.1 variants tested, estimate baseline 3.3x reduction in ID50 relative to D614G wild-type using pseudotyped VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	419	419	1.0
p.Q1071H	vaccine neutralization efficacy	Neutralization efficiency (ID50) against B.1.617.1- v1 ("Kappa") reduced 3.4x relative to D614G wildtype using pseudo- typed VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	419	419	1.0
p.Q1071H	vaccine neutralization efficacy	1.5x reduction in neutralization (ID50) in sera 3 weeks after one dose of Pfizer/BioNTech BNT162b2 (up to 5 naive and post infection vaccinees) [exact set of variants used is not listed in the manuscript]	B.1.617.1	Gong et al. (2021)	419	419	1.0
p.Q1071H	vaccine neutraliza- tion efficacy	Relative to B.1, Kappa (B.1.617.1) shows mean 1.97x decrease in neutralization efficiency by 18 vaccinee sera (BNT162b2 and mRNA1273).	B.1.617.1	Wilhelm et al. (2021)	419	419	1.0
p.Q1071H	vaccine neutralization efficacy	Average 2x neutralization efficiency decrease against sera from 26 Phase II trial Covaxin (BBV152) vaccine recipents. Very similar to B.1.1.7 (extasciitilde2x) in the same study, and the neutralization from 17 sera of natural infections across B1 lineages.	B.1.617.1	Yadav et al. (2021)	419	419	1.0
p.E154K	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant combination (representing lineage B.1.617.1) showed a 1.12x increase in binding (KD) relative to D614G, but described as "not significant". [exact variant list not provided in manuscript, is inferred and represents minimal set agreed upon commonly]	B.1.617.1	Gong et al. (2021)	418	418	1.0
p.E154K	convalescent plasma escape	Escape mutant found after in passage in plasma pool of 26 convalescents mean 1.5 post symptom onset.	B.1.617.1	Schmidt et al. (2021)	418	418	1.0
p.E154K	convalescent plasma escape	Relative to B.1, Kappa (B.1.617.1) shows 5.71x decrease in neutralization efficiency by convalescent plasma [no cohort details provided]	B.1.617.1	Wilhelm et al. (2021)	418	418	1.0
p.E154K	trafficking	extasciitilde3.5x cleavage of S2 relative to WA1 (D614G) wildtype by Kappa (B.1.617.1) variant as measured by mass spectrometry of Vero-TMPRSS culture (compare to extasciitilde4.5x for closely related Delta B.1.617.2 also with P681R at the cleavage site).	B.1.617.1	Esclera et al. (2021)	418	418	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.E154K	vaccine neutraliza- tion efficacy	Inferring from two B.1.617.1 variants tested, estimate baseline 3.3x reduction in ID50 relative to D614G wild-type using pseudotyped VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	418	418	1.0
p.E154K	vaccine neutralization efficacy	Neutralization efficiency (ID50) against B.1.617.1-v1 ("Kappa") reduced 3.4x relative to D614G wildtype using pseudotyped VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	418	418	1.0
p.E154K	vaccine neutraliza- tion efficacy	1.5x reduction in neutralization (ID50) in sera 3 weeks after one dose of Pfizer/BioNTech BNT162b2 (up to 5 naive and post infection vaccinees) [exact set of variants used is not listed in the manuscript]	B.1.617.1	Gong et al. (2021)	418	418	1.0
p.E154K	vaccine neutralization efficacy	Relative to B.1, Kappa (B.1.617.1) shows mean 1.97x decrease in neutralization efficiency by 18 vaccinee sera (BNT162b2 and mRNA1273).	B.1.617.1	Wilhelm et al. (2021)	418	418	1.0
p.E154K	vaccine neutralization efficacy	Average 2x neutralization efficiency decrease against sera from 26 Phase II trial Covaxin (BBV152) vaccine recipents. Very similar to B.1.1.7 (extascitilde2x) in the same study, and the neutralization from 17 sera of natural infections across B1 lineages.	B.1.617.1	Yadav et al. (2021)	418	418	1.0
p.E484Q	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant combination (representing lineage B.1.617.1) showed a 1.12x increase in binding (KD) relative to D614G, but described as "not significant". [exact variant list not provided in manuscript, is inferred and represents minimal set agreed upon commonly]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.E484Q	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant combination (representing lineage B.1.617) showed a 1.85x increase in binding (KD) relative to D614G. [exact variant list not provided in manuscript, is inferred fro common knowledge]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.E484Q	antibody epitope effects	>20% (ELISA significance threshold) drop in antibody binding by this variant against monoclonal antibody VH-Fc ab8.	B.1.617.1	Sun et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate
p.E484Q	convalescent plasma binding	This variant combination (representing	B.1.617.1	Gong et al. (2021)	420	420	1.0
		lineage B.1.617) showed					
		a 1.22x decrease in Spike binding (relative					
		to D614G alone) by					
		5 plasma collected 8					
		months post-symptom- onset. [exact variant					
		list not provided in					
		manuscript, is inferred					
o.E484Q	convalescent plasma	fro common knowledge] In 3 of 11 subjects'	B.1.617.1	Greaney et al.	420	420	1.0
J.12404Q	escape	convalescent sera in an	B.1.017.1	(2021)	420	420	1.0
	_	early+late mutational					
		landscape analysis of the RBD, E484Q shows					
		a notably resistant					
		profile, comparable to					
		or even more resistant than E484K at later					
		time points (i.e. more					
		resistant to immune					
		cell somatic mutation evolution), see Figure					
		5a,b. Subject C 32 days					
		post-infection showed					
		»10 fold reduction in neutralization, reducing					
		to extasciitilde10-fold					
		by day 104. Subject B					
		26 days post-infection showed extasciitide10					
		fold reduction in neu-					
		tralization, reducing to					
		extasciitilde4x at day 113. Notably, Subject					
		B also showed smaller					
		than typical (extasci-					
		itilde10 vs 30+ fold) reduction in one-month					
		neutralization by E484K					
		at day 32, and no E484K					
		immune escape at day 104. Subject I 26 days					
		post-infection showed					
		extasciitilde10 fold					
		reduction in neutraliza- tion, with no reduction					
		in escape at day 102.					
		Notably, Subject I also					
		showed smaller than typical (extasciitilde10					
		vs 30+ fold) reduction					
		in one-month neutraliza-					
		tion by E484K at day 26, and no E484K immune					
		escape at day 102.					
.E484Q	convalescent plasma	Pseudotyped viruses	B.1.617.1	Tada et al.	420	420	1.0
	escape	for B.1.617 was 2.3-fold resistant to neutraliza-		(2021)			
		tion by convalescent					
		sera compared to wild					
		type - a finding that was similar to that of					
		the 3-fold resistance of					
		the South Africa B.1.351					
		variant using the same assay. The resistance of					
		B.1.617 was caused by					
		the L452R and E484Q					
		mutation, based on results from viruses					
		pseudotyped for indi-					
		vidual variants within					
		B.1.617. [details on the convalescent patient					
		sera collection are not					
		abundantly clear in the					
E4940	convolescent =lase=	preprint]	B 1 617 1	Wilhelm et al.	420	420	1.0
.E484Q	convalescent plasma escape	Relative to B.1, Kappa (B.1.617.1) shows 5.71x	B.1.617.1	(2021) Wilhelm et al.	420	420	1.0
		decrease in neutraliza-					
		tion efficiency by con- valescent plasma [no co-					



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.E484Q	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.1.617.1	Starr et al. (2021)	420	420	1.0
p.E484Q	pharmaceutical effectiveness	Bamlanivimab (LY-CoV555) lost extasci- itilde20x binding against this isolated mutation. Casirivimab lost extasci- itilde4x binding against this isolated mutation.	B.1.617.1	Engelhart et al. (2021)	420	420	1.0
p.E484Q	symptom prevalence	Gross examination of 3+3 hamster lung specimens showed pronounced congestion and hemorrhages on days 5 and 7 post-infection in the case of the B.1.617.1 as compared with the B.1. The lung lesions with the B.1 variant were minimal to mild whereas with B.1.617.1 they were moderate. For B.1 pneumonic changes were minimal to mild (inflammatory cell infiltration, focal consolidation and mild congestion). The pronounced changes (moderate to severe) with mononuclear infiltration in the alveolar interstitial septal thickening, consolidation and pneumocyte hyperplasia were observed with B.1.617.1 variant consistently.	B.1.617.1	Yadav et al. (2021)	420	420	1.0
p.E484Q	trafficking	extasciitilde3.5x cleavage of S2 relative to WA1 (D614G) wildtype by Kappa (B.1.617.1) variant as measured by mass spectrometry of Vero-TMPRSS culture (compare to extascitilde4.5x for closely related Delta B.1.617.2 also with P681R at the cleavage site).	B.1.617.1	Esclera et al. (2021)	420	420	1.0
p.E484Q	trafficking	This variant alone shows a 10x decrease in cell en- try efficiency (RLU mea- surement in 293T cells) compared to D614G.	B.1.617.1	Ferriera et al (2021)	420	420	1.0
p.E484Q	transmissibility	The combination caused a 3-fold increase in infec- tivity relative to D614G wild type. [compare to 3.5x for L452R alone]	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.E484Q	transmissibility	Normalized for particle number, on ACE2.293T cells showed that the B.1.617 spike protein was >2-fold increase in infectivity relative to D614G wild type.	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.E484Q	vaccine neutraliza- tion efficacy	Inferring from two B.1.617.1 variants tested, estimate baseline 3.3x reduction in ID50 relative to D614G wild-type using pseudotyped VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.E484Q	vaccine neutralization efficacy	Neutralization efficiency (ID50) against B.1.617.1- v1 ("Kappa") reduced 3.4x relative to D614G wildtype using pseudo- typed VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	420	420	1.0
p.E484Q	vaccine neutralization efficacy	Nine stored sera from Pfizer BNT162b2 vaccinees were tested against a range of spike mutation bearing PV. E484Q had a extasciitilde5x drop in neutralization (vs extasciitilde10x for E484K). When E484Q and L452R were combined, the fold change was significant, but similar to that of L452R alone (extasciitilde2x), suggesting no evidence for an additive effect [perhaps even E484Q effect dilution].	B.1.617.1	Ferreira et al. (2021)	420	420	1.0
p.E484Q	vaccine neutralization efficacy	1.5x reduction in neutralization (ID50) in sera 3 weeks after one dose of Pfizer/BioNTech BNT162b2 (up to 5 naive and post infection vaccinees) [exact set of variants used is not listed in the manuscript]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.E484Q	vaccine neutraliza- tion efficacy	1.4x reduction in neutralization (ID50) in sera 3 weeks after one dose of Pfizer/BioNTech BNT162b2 (up to 5 naive and post infection vaccinees)	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.E484Q	vaccine neutraliza- tion efficacy	Pseudotyped viruses for B.1.617 was 4-fold resistant to neutralization by 6 BNT162b2 vaccine sera 28 days post-booster compared to wild type - a finding that was similar to that of the 3.4-fold resistance of the South Africa B.1.351 variant using the same assay. Neutralization by 3 Moderna vaccine sera 28 days post-booster was 5-fold resistant (vs. 2.2-fold for B.1.351). The resistance of B.1.617 was caused by the L452R and E484Q mutation, based on results from viruses pseudotyped for individual variants within B.1.617.	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.E484Q	vaccine neutraliza- tion efficacy	Relative to B.1, Kappa (B.1.617.1) shows mean 1.97x decrease in neutralization efficiency by 18 vaccinee sera (BNT162b2 and mRNA1273).	B.1.617.1	Wilhelm et al. (2021)	420	420	1.0
p.E484Q	vaccine neutralization efficacy	Average 2x neutralization efficiency decrease against sera from 26 Phase II trial Covaxin (BBV152) vaccine recipents. Very similar to B.1.1.7 (extascitilde2x) in the same study, and the neutralization from 17 sera of natural infections across B1 lineages.	B.1.617.1	Yadav et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.E484Q	vaccinee plasma binding	This variant combination (representing lineage B.1.617) showed a 1.30x 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. 1.18x 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. [exact variant list not provided in manuscript, is inferred fro common knowledge]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.E484Q	viral load	In 9 infected ham- sters each for B.1 and B.1.617.1, no significant change in viral load or subgenomic RNA levels were detected.	B.1.617.1	Yadav et al. (2021)	420	420	1.0
p.P681R	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains- Fc portion IgG complex, this variant combination (representing lineage B.1.617.1) showed a 1.12x increase in binding (KD) relative to D614G, but described as "not significant". [exact variant list not provided in manuscript, is inferred and represents minimal set agreed upon commonly]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.P681R	convalescent plasma escape	Relative to B.1, Kappa (B.1.617.1) shows 5.71x decrease in neutralization efficiency by convalescent plasma [no cohort details provided]	B.1.617.1	Wilhelm et al. (2021)	420	420	1.0
p.P681R	symptom prevalence	Gross examination of 3+3 hamster lung specimens showed pronounced congestion and hemorrhages on days 5 and 7 post-infection in the case of the B.1.617.1 as compared with the B.1. The lung lesions with the B.1 variant were minimal to mild whereas with B.1.617.1 they were moderate. For B.1 pneumonic changes were minimal to mild (inflammatory cell infiltration, focal consolidation and mild congestion). The pronounced changes (moderate to severe) with mononuclear infiltration in the alveolar interstitial septal thickening, consolidation and pneumocyte hyperplasia were observed with B.1.617.1 variant consistently.	B.1.617.1	Yadav et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.P681R	trafficking	extasciitilde3.5x cleavage of S2 relative to WA1 (D614G) wildtype by Kappa (B.1.617.1) variant as measured by mass spectrometry of Vero-TMPRSS culture (compare to extasciitilde4.5x for closely related Delta B.1.617.2 also with P681R at the cleavage site).	B.1.617.1	Esclera et al. (2021)	420	420	1.0
p.P681R	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 \$680. In a homologuous site in Infectious Bronchitis Virus (IBV, Gammacoronaviruses), abolition of \$680 phosphorylation improves furin cleavage (and presumably cell entry). [Inference from similar positively charged substitution P681H actually described in the work]	B.1.617.1	Maaroufi (2021)	420	420	1.0
p.P681R	trafficking	Quantification of the band intensities showed that the P681R mutation, which lies near the proteolytic processing site, caused a small increase in proteolytic processing as measured by a 2-fold decrease in the ratio of S/S2.	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.P681R	transmissibility	Normalized for particle number, on ACE2.293T cells showed that the B.1.617 spike protein was >2-fold increase in infectivity relative to D614G wild type.	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.P681R	vaccine neutralization efficacy	Inferring from two B.1.617.1 variants tested, estimate baseline 3.3x reduction in ID50 relative to D614G wild-type using pseudotyped VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	420	420	1.0
p.P681R	vaccine neutralization efficacy	Neutralization efficiency (ID50) against B.1.617.1- v1 ("Kappa") reduced 3.4x relative to D614G wildtype using pseudo- typed VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	420	420	1.0
p.P681R	vaccine neutraliza- tion efficacy	1.5x reduction in neutralization (ID50) in sera 3 weeks after one dose of Pfizer/BioNTech BNT162b2 (up to 5 naive and post infection vaccinees) [exact set of variants used is not listed in the manuscript]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.P681R	vaccine neutraliza- tion efficacy	Relative to B.1, Kappa (B.1.617.1) shows mean 1.97x decrease in neutralization efficiency by 18 vaccinee sera (BNT162b2 and mRNA1273).	B.1.617.1	Wilhelm et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.P681R	vaccine neutraliza- tion efficacy	Average 2x neutralization efficiency decrease against sera from 26 Phase II trial Covaxin (BBV152) vaccine recipents. Very similar to B.1.1.7 (extasciitilde2x) in the same study, and the neutralization from 17 sera of natural infections across B1 lineages.	B.1.617.1	Yadav et al. (2021)	420	420	1.0
p.P681R	viral load	In 9 infected hamsters each for B.1 and B.1.617.1, no significant change in viral load or subgenomic RNA levels were detected.	B.1.617.1	Yadav et al. (2021)	420	420	1.0
p.P681R	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. [PG: Inferred by conservative AA substitution of described P681H]	B.1.617.1	Tada et al. (2021)	420	420	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.1.617.1	Gong et al. (2021)	418	389	0.93
p.T95I	convalescent plasma binding	No change in Spike binding (relative to D614G alone) by 5 plasma collected 8 months postsymptom-onset.	B.1.617.1	Gong et al. (2021)	418	389	0.93
p.T95I	vaccine neutraliza- tion efficacy	Neutralization efficiency (ID50) against B.1.617.1-v1 ("Kappa") reduced 3.4x relative to D614G wildtype using pseudo- typed VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	418	389	0.93
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 nonseroconverted 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.1.617.1	Gong et al. (2021)	418	389	0.93
p.L452R	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains- Fc portion IgG complex, this variant showed a 2.66x increase in binding (KD) relative to D614G.	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.L452R	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant combination (representing lineage B.1.617) showed a 1.85x increase in binding (KD) relative to D614G. [exact variant list not provided in manuscript, is inferred fro common knowledge]	B.1.617.1	Gong et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.L452R	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant combination (representing lineage B.1.617.1) showed a 1.12x increase in binding (KD) relative to D614G, but described as "not significant". [exact variant list not provided in manuscript, is inferred and represents minimal set agreed upon commonly]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.L452R	ACE2 receptor binding affinity	extasciitilde1.7-fold increase in binding affinity	B.1.617.1	Motozono et al. (2021)	420	420	1.0
p.L452R	T cell evasion	vs wild type. L452R derivative virus did not induce IFN- gamma expression even at the highest concen- tration tested (10 nM) in two different A*24:02 convalescent sera donor plasma (linear epitope NYNYLYRLF 448,456).	B.1.617.1	Motozono et al. (2021)	420	420	1.0
p.L452R	antibody epitope effects	Resistent to some neutralizing antibodies: mAbs X593 and P2B-	B.1.617.1	Li et al. (2020)	420	420	1.0
p.L452R	antibody epitope effects	Mutant screen in neutralization assay with a broad range of monoclonal antibodies shows resistence to more than one antibody.	B.1.617.1	Liu et al. (2021)	420	420	1.0
p.L452R	antibody epitope effects	10 of 14 RBD-specific mAbs that showed at least 10-fold reduced neutralization of B.1.427/B.1.429 variant pseudotype (S13I, W152C, and L452R) were also found to poorly bind to just a L452R RBD mutant, demonstrating a role for this mutation as an escape mechanism for certain RBD-targeting mAbs.	B.1.617.1	McCallum et al. (2021)	420	420	1.0
p.L452R	antibody epitope effects	extasciitilde20% (ELISA significance threshold) drop in antibody binding (ELISA) by this variant against monoclonal antibody VH ab6.	B.1.617.1	Sun et al. (2021)	420	420	1.0
p.L452R	convalescent plasma binding	2.15x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.L452R	convalescent plasma binding	This variant combination (representing lineage B.1.617) showed a 1.22x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptomonset. [exact variant list not provided in manuscript, is inferred fro common knowledge]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.L452R	convalescent plasma escape	Observed extasciitilde2x decrease on average in 16 health workers' convales- cent sera.	B.1.617.1	Alenquer et al. (2021)	420	420	1.0
p.L452R	convalescent plasma escape	Ablation of neutralization capability of 3 of 4 convalescent sera tested, the other is significantly hindered.	B.1.617.1	Liu et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.L452R	convalescent plasma escape	Pseudotyped viruses for B.1.617 was 2.3-fold resistant to neutralization by convalescent sera compared to wild type - a finding that was similar to that of the 3-fold resistance of the South Africa B.1.351 variant using the same assay. The resistance of B.1.617 was caused by the L452R and E484Q mutation, based on results from viruses pseudotyped for individual variants within B.1.617. [details on the convalescent patient sera collection are not abundantly clear in the preprint]	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.L452R	convalescent plasma escape	Relative to B.1, Epsilon (B.1.417/429) shows 1.74x-2.35x decrease in neutralization efficiency by convalescent plasma [no cohort details provided]	B.1.617.1	Wilhelm et al. (2021)	420	420	1.0
p.L452R	convalescent plasma escape	Relative to B.1, Kappa (B.1.617.1) shows 5.71x decrease in neutralization efficiency by convalescent plasma [no cohort details provided]	B.1.617.1	Wilhelm et al. (2021)	420	420	1.0
p.L452R	gene expression in- crease	Experimentally, Spike gene expression in- creased 0.32 fold	B.1.617.1	Starr et al. (2020)	420	420	1.0
p.L452R	monoclonal anti- body serial passage escape	Ranked effective mutant against this position in the RBD for highly neu- tralizing COV2-2096	B.1.617.1	Greaney et al. (2020)	420	420	1.0
p.L452R	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.1.617.1	Starr et al. (2021)	420	420	1.0
p.L452R	monoclonal anti- body serial passage escape	Class 2/3 antibody C628 and class 2 antibody C643 selected for the emergence of the L452R mutation in vitro.	B.1.617.1	Wang et al. (2021)	420	420	1.0
p.L452R	pharmaceutical effectiveness	Bamlanivimab (LY-CoV555) lost extasci- itilde5x binding against this isolated mutation. Cligavimab lost extasci- itilde4x binding against this isolated mutation. Regdanvimab lost ex- tasciitilde4x binding against this isolated mutation.	B.1.617.1	Engelhart et al. (2021)	420	420	1.0
p.L452R	pharmaceutical effectiveness	Bamlanivimab (LY-CoV555) entirely lost its neutralizing activity due to the central location of L452R in the epitopes recognized by this mAb. Regdanvimab (CT-P59), and to a smaller extent etesevimab, showed a reduction in neutralization potency.	B.1.617.1	McCallum et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.L452R	symptom prevalence	Gross examination of 3+3 hamster lung specimens showed pronounced congestion and hemorrhages on days 5 and 7 post-infection in the case of the B.1.617.1 as compared with the B.1. The lung lesions with the B.1 variant were minimal to mild whereas with B.1.617.1 they were moderate. For B.1 pneumonic changes were minimal to mild (inflammatory cell infiltration, focal consolidation and mild congestion). The pronounced changes (moderate to severe) with mononuclear infiltration in the alveolar interstitial space, interstitial septal thickening, consolidation and pneumocyte hyperplasia were observed with B.1.617.1	B.1.617.1	Yadav et al. (2021)	420	420	1.0
p.L452R	trafficking	variant consistently. We observed increased entry by pseudoviruses carrying the L452R mutation compared to D614G alone, with a 6.7 to 22.5-fold increase in 293T cells and a 5.8 to 14.7-fold increase in human airway organoids.	B.1.617.1	Deng et al. (2021)	420	420	1.0
p.L452R	trafficking	extasciitilde3.5x cleavage of S2 relative to WA1 (D614G) wildtype by Kappa (B.1.617.1) variant as measured by mass spectrometry of Vero-TMPRSS culture (compare to extasciitilde4.5x for closely related Delta B.1.617.2 also with P681R at the cleavage site).	B.1.617.1	Esclera et al. (2021)	420	420	1.0
p.L452R	trafficking	This variant alone shows a extasciitilde5x decrease in cell entry efficiency (RLU measurement in 293T cells) compared to D614G. [listed as L454R in Figure, but L452R in text, also text suggests not statistucally significant, but error bars say otherwise in Figure 4]	B.1.617.1	Ferriera et al (2021)	420	420	1.0
p.L452R	trafficking	Increased stability of RBD expression in yeast, suggesting in- creased Spike protein stability.	B.1.617.1	Motozono et al. (2021)	420	420	1.0
p.L452R	transmissibility	Increased infectivity of the B.1.617 spike was attributed to L452R, which itself caused a 3.5-fold increase in infec- tivity relative to D614G wild type. [In combina- tion with E484Q caused a lower 3-fold increase]	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.L452R	transmissibility	The combination caused a 3-fold increase in infec- tivity relative to D614G wild type. [compare to 3.5x for L452R alone]	B.1.617.1	Tada et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.L452R	transmissibility	Normalized for particle number, on ACE2.293T cells showed that the B.1.617 spike protein was >2-fold increase in infectivity relative to D614G wild type.	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.L452R	vaccine neutraliza- tion efficacy	Inferring from two B.1.617.1 variants tested, estimate baseline 3.3x reduction in ID50 relative to D614G wild-type using pseudotyped VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	420	420	1.0
p.L452R	vaccine neutralization efficacy	Neutralization efficiency (ID50) against B.1.617.1-v1 ("Kappa") reduced 3.4x relative to D614G wildtype using pseudotyped VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	420	420	1.0
p.L452R	vaccine neutraliza- tion efficacy	Nine stored sera from Pfizer BNT162b2 vacci- nees were tested against a range of spike muta- tion bearing PV. L452R conferred about a two- fold reduction in neutral- isation by vaccine sera, but was not statistically significant with this sam- ple size.	B.1.617.1	Ferreira et al. (2021)	420	420	1.0
p.L452R	vaccine neutralization efficacy	Nine stored sera from Pfizer BNT162b2 vaccinees were tested against a range of spike mutation bearing PV. E484Q had a extasciitilde5x drop in neutralization (vs extasciitilde10x for E484K). When E484Q and L452R were combined, the fold change was significant, but similar to that of L452R alone (extasciitilde2x), suggesting no evidence for an additive effect [perhaps even E484Q effect dilution].	B.1.617.1	Ferreira et al. (2021)	420	420	1.0
p.L452R	vaccine neutralization efficacy	1.4x reduction in neutralization (ID50) in sera 3 weeks after one dose of Pfizer/BioNTech BNT162b2 (up to 5 naive and post infection vaccinees)	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.L452R	vaccine neutraliza- tion efficacy	1.5x reduction in neutralization (ID50) in sera 3 weeks after one dose of Pfizer/BioNTech BNT162b2 (up to 5 naive and post infection vaccinees) [exact set of variants used is not listed in the manuscript]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.L452R	vaccine neutraliza- tion 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.1.617.1	Jacobson et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.L452R	vaccine neutralization efficacy	Pseudotyped viruses for B.1.617 was 4-fold resistant to neutralization by 6 BNT162b2 vaccine sera 28 days post-booster compared to wild type - a finding that was similar to that of the 3.4-fold resistance of the South Africa B.1.351 variant using the same assay. Neutralization by 3 Moderna vaccine sera 28 days post-booster was 5-fold resistant (vs. 2.2-fold for B.1.351). The resistance of B.1.617 was caused by the L452R and E484Q mutation, based on results from viruses pseudotyped for individual variants within B.1.617.	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.L452R	vaccine neutraliza- tion efficacy	Relative to B.1, Epsilon (B.1.417/429) shows 1.74x-2.35x decrease in neutralization efficiency by 18 vaccinee sera (BNT162b2 and mRNA1273).	B.1.617.1	Wilhelm et al. (2021)	420	420	1.0
p.L452R	vaccine neutralization efficacy	Relative to B.1, Kappa (B.1.617.1) shows mean 1.97x decrease in neutralization efficiency by 18 vaccinee sera (BNT162b2 and mRNA1273).	B.1.617.1	Wilhelm et al. (2021)	420	420	1.0
p.L452R	vaccine neutralization efficacy	Average 2x neutralization efficiency decrease against sera from 26 Phase II trial Covaxin (BBV152) vaccine recipents. Very similar to B.1.1.7 (extascitilde2x) in the same study, and the neutralization from 17 sera of natural infections across B1 lineages.	B.1.617.1	Yadav et al. (2021)	420	420	1.0
p.L452R	vaccinee plasma binding	1.05x 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. 1.16x 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.1.617.1	Gong et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.L452R	vaccinee plasma binding	This variant combination (representing lineage B.1.617) showed a 1.30x 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. 1.18x 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. [exact variant list not provided in manuscript, is inferred fro common knowledge]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.L452R	viral load	In 9 infected hamsters each for B.1 and B.1.617.1, no significant change in viral load or subgenomic RNA levels were detected.	B.1.617.1	Yadav et al. (2021)	420	420	1.0
p.L452R	virion structure	Estimated free energy change (ddG) for this variant is -0.67 kcal/mol (i.e. destabilizing relative to wild type)	B.1.617.1	Spratt et al. (2021)	420	420	1.0
p.G142D	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant combination (representing lineage B.1.617.1) showed a 1.12x increase in binding (KD) relative to D614G, but described as "not significant". [exact variant list not provided in manuscript, is inferred and represents minimal set agreed upon commonly]	B.1.617.1	Gong et al. (2021)	401	401	1.0
p.G142D	convalescent plasma escape	Relative to B.1, Kappa (B.1.617.1) shows 5.71x decrease in neutralization efficiency by convalescent plasma [no cohort details provided]	B.1.617.1	Wilhelm et al. (2021)	401	401	1.0
p.G142D	monoclonal anti- body serial passage escape	Escape mutation against Spike N terminal do- main antigenic supersite i mAbs S2M28, S2X28, S2X333	B.1.617.1	McCallum et al. (2021)	401	401	1.0
p.G142D	monoclonal anti- body serial passage escape	Selected twice in passage with mAb COV2-2489.	B.1.617.1	Suryadevara et al. (2021)	401	401	1.0
p.G142D	trafficking	extasciitilde3.5x cleavage of S2 relative to WA1 (D614G) wildtype by Kappa (B.1.617.1) variant as measured by mass spectrometry of Vero-TMPRSS culture (compare to extasciitilde4.5x for closely related Delta B.1.617.2 also with P681R at the cleavage site).	B.1.617.1	Esclera et al. (2021)	401	401	1.0
p.G142D	vaccine neutraliza- tion efficacy	Inferring from two B.1.617.1 variants tested, estimate baseline 3.3x reduction in ID50 relative to D614G wild-type using pseudotyped VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	401	401	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.G142D	vaccine neutraliza- tion efficacy	Neutralization efficiency (ID50) against B.1.617.1-v1 ("Kappa") reduced 3.4x relative to D614G wildtype using pseudotyped VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	401	401	1.0
p.G142D	vaccine neutraliza- tion efficacy	1.5x reduction in neutralization (ID50) in sera 3 weeks after one dose of Pfizer/BioNTech BNT162b2 (up to 5 naive and post infection vaccinees) [exact set of variants used is not listed in the manuscript]	B.1.617.1	Gong et al. (2021)	401	401	1.0
p.G142D	vaccine neutralization efficacy	Relative to B.1, Kappa (B.1.617.1) shows mean 1.97x decrease in neutralization efficiency by 18 vaccinee sera (BNT162b2 and mRNA1273).	B.1.617.1	Wilhelm et al. (2021)	401	401	1.0
p.G142D	vaccine neutraliza- tion efficacy	Average 2x neutralization efficiency decrease against sera from 26 Phase II trial Covaxin (BBV152) vaccine recipents. Very similar to B.1.1.7 (extascitilde2x) in the same study, and the neutralization from 17 sera of natural infections across	B.1.617.1	Yadav et al. (2021)	401	401	1.0
p.D614G	ACE2 receptor binding affinity	B1 lineages. Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant combination (representing lineage B.1.617.1) showed a 1.12x increase in binding (KD) relative to D614G, but described as "not significant". [exact variant list not provided in manuscript, is inferred and represents minimal set agreed upon commonly]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.D614G	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant showed a 2.66x increase in binding (KD) relative to D614G.	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.D614G	ACE2 receptor binding affinity	Using flow cytometry and ACE2 ectodomains-Fc portion IgG complex, this variant combination (representing lineage B.1.617) showed a 1.85x increase in binding (KD) relative to D614G. [exact variant list not provided in manuscript, is inferred fro common knowledge]	B.1.617.1	Gong et al. (2021)	420	420	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.1.617.1	Gong et al. (2021)	420	420	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.1.617.1	Li et al. (2020)	420	420	1.0
p.D614G	convalescent plasma binding	2.15x increase in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptom- onset.	B.1.617.1	Gong et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.D614G	convalescent plasma binding	This variant combination (representing lineage B.1.617) showed a 1.22x decrease in Spike binding (relative to D614G alone) by 5 plasma collected 8 months post-symptomonset. [exact variant list not provided in manuscript, is inferred fro common knowledge]	B.1.617.1	Gong et al. (2021)	420	420	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.1.617.1	Gong et al. (2021)	420	420	1.0
p.D614G	convalescent plasma escape	Pseudotyped viruses for B.1.617 was 2.3-fold resistant to neutralization by convalescent sera compared to wild type - a finding that was similar to that of the 3-fold resistance of the South Africa B.1.351 variant using the same assay. The resistance of B.1.617 was caused by the L452R and E484Q mutation, based on results from viruses pseudotyped for individual variants within B.1.617. [details on the convalescent patient sera collection are not abundantly clear in the preprint]	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.D614G	convalescent plasma escape	Relative to B.1, Kappa (B.1.617.1) shows 5.71x decrease in neutralization efficiency by convalescent plasma [no cohort details provided]	B.1.617.1	Wilhelm et al. (2021)	420	420	1.0
p.D614G	convalescent plasma escape	Relative to B.1, Epsilon (B.1.417/429) shows 1.74x-2.35x decrease in neutralization efficiency by convalescent plasma [no cohort details provided]	B.1.617.1	Wilhelm et al. (2021)	420	420	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.1.617.1	Landis et al. (2021)	420	420	1.0
p.D614G	syncytium formation	Slight increase in Vero cell-cell membrane fu- sion assay under infec- tion with VSV pseudo- typed virus.	B.1.617.1	Kim et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
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	B.1.617.1	Planas et al. (2021)	420	420	1.0
p.D614G	trafficking	circulating variant shown in vitro to not have major defects or enhancement of cell surface protein trafficking (i.e. Spike cleavage or fusion required for cell entry)	B.1.617.1	Barrett et al. (2021)	420	420	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.1.617.1	Daniloski et al. (2021)	420	420	1.0
p.D614G	trafficking	extasciitilde3.5x cleavage of S2 relative to WA1 (D614G) wildtype by Kappa (B.1.617.1) variant as measured by mass spectrometry of Vero-TMPRSS culture (compare to extasciitilde4.5x for closely related Delta B.1.617.2 also with P681R at the cleavage site).	B.1.617.1	Esclera et al. (2021)	420	420	1.0
p.D614G	trafficking	No change in infectivity (24h) relative to D614G alone in Caco-2 cells, Vero or Calu-3.	B.1.617.1	Kim et al. (2021)	420	420	1.0
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.1.617.1	Kim et al. (2021)	420	420	1.0
p.D614G	trafficking	Among S variants tested, the D614G mutant shows the highest cell entry (extasciitlde3.5x wild type), as supported by structural and binding analyses.	B.1.617.1	Ozono et al. (2020)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.D614G	trafficking	Quantification of the band intensities showed that the P681R mutation, which lies near the proteolytic processing site, caused a small increase in proteolytic processing as measured by a 2-fold decrease in the ratio of S/S2.	B.1.617.1	Tada et al. (2021)	420	420	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.1.617.1	Zhang et l. (2020)	420	420	1.0
p.D614G	transmissibility	Increased infectivity of the B.1.617 spike was attributed to L452R, which itself caused a 3.5-fold increase in infec- tivity relative to D614G wild type. [In combina- tion with E484Q caused a lower 3-fold increase]	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.D614G	transmissibility	The combination caused a 3-fold increase in infectivity relative to D614G wild type. [compare to 3.5x for L452R alone]	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.D614G	transmissibility	Normalized for particle number, on ACE2.293T cells showed that the B.1.617 spike protein was >2-fold increase in infectivity relative to D614G wild type.	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.D614G	vaccine neutralization efficacy	Inferring from two B.1.617.1 variants tested, estimate baseline 3.3x reduction in ID50 relative to D614G wild-type using pseudotyped VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	420	420	1.0
p.D614G	vaccine neutralization efficacy	Neutralization efficiency (ID50) against B.1.617.1-v1 ("Kappa") reduced 3.4x relative to D614G wildtype using pseudotyped VSV assay on 8 Moderna vaccinee sera one week post-booster.	B.1.617.1	Choi et al. (2021)	420	420	1.0
p.D614G	vaccine neutralization 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.1.617.1	Garcia- Beltran et al. (2021)	420	420	1.0
p.D614G	vaccine neutralization efficacy	1.5x reduction in neutralization (ID50) in sera 3 weeks after one dose of Pfizer/BioNTech BNT162b2 (up to 5 naive and post infection vaccinees) [exact set of variants used is not listed in the manuscript]	B.1.617.1	Gong et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.D614G	vaccine neutraliza- tion efficacy	1.4x reduction in neutralization (ID50) in sera 3 weeks after one dose of Pfizer/BioNTech BNT162b2 (up to 5 naive and post infection vaccinees)	B.1.617.1	Gong et al. (2021)	420	420	1.0
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.1.617.1	Kuzmina et al. (2021)	420	420	1.0
p.D614G	vaccine neutralization efficacy	Pseudotyped viruses for B.1.617 was 4-fold resistant to neutralization by 6 BNT162b2 vaccine sera 28 days post-booster compared to wild type - a finding that was similar to that of the 3.4-fold resistance of the South Africa B.1.351 variant using the same assay. Neutralization by 3 Moderna vaccine sera 28 days post-booster was 5-fold resistant (vs. 2.2-fold for B.1.351). The resistance of B.1.617 was caused by the L452R and E484Q mutation, based on results from viruses pseudotyped for individual variants within B.1.617.	B.1.617.1	Tada et al. (2021)	420	420	1.0
p.D614G	vaccine neutraliza- tion efficacy	Relative to B.1, Kappa (B.1.617.1) shows mean 1.97x decrease in neutralization efficiency by 18 vaccinee sera (BNT162b2 and mRNA1273).	B.1.617.1	Wilhelm et al. (2021)	420	420	1.0
p.D614G	vaccine neutraliza- tion efficacy	Relative to B.1, Epsilon (B.1.417/429) shows 1.74x-2.35x decrease in neutralization efficiency by 18 vaccinee sera (BNT162b2 and mRNA1273).	B.1.617.1	Wilhelm et al. (2021)	420	420	1.0
p.D614G	vaccine neutraliza- tion efficacy	Average 2x neutralization efficiency decrease against sera from 26 Phase II trial Covaxin (BBV152) vaccine recipents. Very similar to B.1.1.7 (extascitilde2x) in the same study, and the neutralization from 17 sera of natural infections across B1 lineages.	B.1.617.1	Yadav et al. (2021)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
p.D614G	vaccinee plasma binding	1.05x 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. 1.16x 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.1.617.1	Gong et al. (2021)	420	420	1.0
p.D614G	vaccinee plasma binding	This variant combination (representing lineage B.1.617) showed a 1.30x 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. 1.18x 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. [exact variant list not provided in manuscript, is inferred fro common knowledge]	B.1.617.1	Gong et al. (2021)	420	420	1.0
p.D614G	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.1.617.1	Gong et al. (2021)	420	420	1.0
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.1.617.1	Plante et al. (2020)	420	420	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.1.617.1	Spratt et al. (2021)	420	420	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.1.617.1	Weissman et al. (2020)	420	420	1.0



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Mutations	Sub-category	Function	Lineages	Citation	Sequence Depth	Alternate Allele	Alternate Frequency
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.1.617.1	Yurkovetskiy et al. (2020)	420	420	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.1.617.1	Zhang et al. (2020)	420	420	1.0



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The results here are in whole or part based upon data hosted at the Canadian VirusSeq Data Portal: https://virusseq-dataportal.ca/.We wish to acknowledge the following organisations/laboratories for contributing data to the Portal: Canadian Public Health Laboratory Network (CPHLN), CanCOGGeN VirusSeq and the list of labs available at https://virusseq-dataportal.ca/acknowledgements)

