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A Novel SARS-CoV-2 Variant of Concern, B.1.526, Identified in New York

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



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Emergence and Expansion of the SARS-CoV-2 Variant B.1.526 Identified in New York

Short title: Emergence of SARS-CoV-2 variant B.1.526

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1 **Recent months have seen surges of SARS-CoV-2 infection across the globe with considerable**
2 **viral evolution¹⁻³. Extensive mutations in the spike protein may threaten efficacy of vaccines**
3 **and therapeutic monoclonal antibodies⁴. Two signature mutations of concern are E484K,**
4 **which plays a crucial role in the loss of neutralizing activity of antibodies, and N501Y, a**
5 **driver of rapid worldwide transmission of the B.1.1.7 lineage. Here, we report the emergence**
6 **of variant lineage B.1.526 that contains E484K and its alarming rise to dominance in New**
7 **York City in early 2021. This variant is partially or completely resistant to two therapeutic**
8 **monoclonal antibodies in clinical use and less susceptible to neutralization by convalescent**
9 **plasma or vaccinee sera, posing a modest antigenic challenge. The B.1.526 lineage has now**
10 **been reported from all 50 states in the US and numerous other countries. B.1.526 rapidly**
11 **replaced earlier lineages in New York upon its emergence, with an estimated transmission**
12 **advantage of 35%. Such transmission dynamics, together with the relative antibody**
13 **resistance of its E484K sub-lineage, likely contributed to the sharp rise and rapid spread of**
14 **B.1.526. Although SARS-CoV-2 B.1.526 initially outpaced B.1.1.7 in the region, its growth**
15 **subsequently slowed concurrent with the rise of B.1.1.7 and ensuing variants.**

16
17

18 **Main**

19 While evolution of SARS-CoV-2 was deemed to be slow at the beginning of the global pandemic⁵,
20 multiple major variants of concern have emerged over the past year^{1-3,6}. These lineages are each
21 characterized by numerous mutations in the spike protein, raising concerns that they may escape
22 from therapeutic monoclonals and vaccine-induced antibodies. The hallmark mutation of B.1.1.7,
23 a SARS-CoV-2 variant of concern that emerged in the UK, is N501Y located in the receptor-
24 binding domain (RBD) of spike¹. This variant is seemingly more transmissible and virulent⁷⁻⁹,
25 perhaps due to a higher binding affinity of N501Y for ACE2¹⁰ or a greater propensity to evade
26 host innate immune responses¹¹. Two other variants of concern, B.1.351² and P.1¹², share the
27 N501Y mutation with B.1.1.7 but also contain an E484K substitution in RBD^{2,3}. P.1 emerged as
28 part of a second surge in Manaus, Brazil despite a high pre-existing SARS-CoV-2 seroprevalence
29 in the population¹³. Reinfections with P.1 and another related Brazilian variant P.2 harboring
30 E484K, have been documented^{14,15}. Our previous study on B.1.351 demonstrated that this variant
31 is refractory to neutralization by a number of monoclonal antibodies directed to the top of RBD,
32 including several that have received emergency use authorization⁴. B.1.351 was markedly more
33 resistant to neutralization by convalescent plasma and vaccinee sera. Importantly, these effects
34 were in part mediated by the E484K mutation. These findings are worrisome in light of recent
35 reports that three vaccine trials showed a substantial drop in efficacy in South Africa^{16,17}. Likewise,
36 P.1 was also relatively resistant to antibody neutralization, although not as severely¹⁸. We therefore
37 implemented rapid molecular screening for signature mutations implicated in the success of these
38 early variants of concern.

39

40 **Rapid screening for SARS-CoV-2 mutations**

41 We first developed rapid PCR-based single-nucleotide-polymorphism (SNP) assays (Extended
42 Data Fig. 1) to search for N501Y and E484K mutations in SARS-CoV-2 positive clinical samples
43 stored in the Columbia University Biobank. Between November 1, 2020 and May 1, 2021, 1,602
44 samples were successfully genotyped by PCR. We identified 182/1,602 (11%) samples with
45 E484K and 63/1,602 (3.9%) with N501Y. Eight samples contained both mutations. The earliest
46 case with E484K was collected in mid-November 2020. The proportion of E484K PCR-screened
47 cases substantially increased from 2.0% at the end of 2020 to 24.3% between February 21st and
48 March 5th, 2021 (Fig. 1a), when targeted PCR genotyping was replaced by whole-genome

49 sequencing. Viruses harboring N501Y also increased over time, from the earliest detection in mid-
50 January to 5.3% of screened isolates by the beginning of March.

51

52 **Genomic surveillance of SARS-CoV-2**

53 We next performed untargeted whole genome nanopore sequencing of nasopharyngeal samples
54 collected throughout the study period with cycle threshold (Ct) ≤ 35 . We successfully obtained
55 1,507 SARS-CoV-2 whole genomes (59% of samples with $Ct \leq 35$; Extended Data Fig. 2).
56 Sequencing results verified the E484K and N501Y substitutions in all samples identified by PCR
57 screening. Of sequenced N501Y isolates, 31/41 (76%) were consistent with the B.1.1.7 lineage.
58 Samples which harbored both N501Y and E484K were genotyped as P.1 (n=6), B.1.351 (n=1),
59 and B.1.623 (n=1). However, quite unexpectedly, the large majority of PCR-screened cases with
60 E484K (n=98/128, 77%) fell within a single lineage, B.1.526,¹⁹ recently labeled the Iota variant
61 by the WHO²⁰.

62

63 Analysis of the entire collection of CUIMC genomic sequences (Fig. 1b) showed that by May
64 2021, SARS-CoV-2 variants (including B.1.526, B.1.1.7, and more recently P.1) comprised two-
65 thirds of all sequenced isolates, replacing the vast majority of earlier lineages (Fig. 1b). The
66 proportion of cases caused by B.1.526 rose rapidly from late 2020 through February 2021, and
67 remained at approximately 40-50% of all sequenced cases from March to May 2021, despite a
68 concurrent increase in B.1.1.7. In fact, during the months of December and January when the
69 prevalence of B.1.1.7 was still negligible (Fig. 1b, marking under horizontal axis), the frequency
70 of all viruses in the B.1.526 lineage rose from <5% to 50% while the frequency of other lineages
71 declined from >95% to 50% (Fig. 1b, where white blank space represents other lineages).
72 Calculations using these numbers in a head-to-head comparison and an established mathematical
73 method²¹ indicate that B.1.526 has a growth advantage of ~5% per day. Likewise, fitting a logistic
74 regression model to 478 individual observations from the extended timeframe of November 2020
75 through January 2021 shows that B.1.526 had a similar growth advantage of 4.6% per day (95%
76 CI 2.8–6.5% per day). Given that the serial interval for SARS-CoV-2 transmission is about 7
77 days²² in the absence of any intervention, these results suggest that B.1.526 is ~35% more
78 transmissible than non-variant viruses.

79

80 Demographic and clinical features, including clinical outcomes, were largely comparable in
81 patients with E484K versus those without the signature E484K or N501Y mutations, and between
82 patients with B.1.526-E484K versus those with non-variant lineages²³ (Extended Data Table 1).
83 However, significantly lower Ct values were associated with both E484K (29.49 vs 30.71,
84 $p=0.013$) and B.1.526-E484K (27.65 vs 28.81 in non-variant lineages, $p=0.015$), indicating a
85 modestly higher viral load in these variant samples. A significantly higher proportion of patients
86 B.1.526-E484K were also admitted to the hospital or presented to the emergency department
87 ($p=0.037$).

88

89 **Signature B.1.526 lineage mutations**

90 We identified signature spike-protein mutations in the B.1.526 lineage by comparing all genomes
91 generated in this study (Fig. 1c). Phylogenetic examination showed that the B.1.526 lineage is
92 comprised of two closely related sub-lineages harboring either E484K (B.1.526-E484K; defined
93 as Pangolin lineage B.1.526) or S477N (B.1.526-S477N; Pangolin lineage B.1.526.2), and the
94 additional sub-lineage B.1.526.1, harboring the L452R substitution (B.1.526-L452R). Both
95 B.1.526-E484K and B.1.526-S477N share characteristic spike-protein mutations L5F, T95I,
96 D253G, D614G, and either A701V or Q957R along with either E484K or S477N. Non-spike
97 mutations widely shared by B.1.526-E484K and B.1.526-S477N isolates include: T85I in ORF1a-
98 nsp2; L438P in ORF1a-nsp4, a 9bp deletion $\Delta 106-108$ in ORF1a-nsp6; P323L in ORF1b-nsp12;
99 Q88H in ORF1b-nsp13; Q57H in ORF3a; and P199L and M234I in the N gene. While B.1.526-
100 L452R isolates shared a number of mutations across the genome in ORF-1ab, ORF-3ab, ORF-8,
101 and N, it does not share characteristic spike mutations with B.1.526-E484K and B.1.526-S477N.

102

103 To further investigate the evolutionary history of B.1.526, we performed phylogenetic analyses on
104 genomes in this collection and in GISAID harboring the ORF1a-nsp6 deletion $\Delta 106-108$, along
105 with mutation A20262G that uniquely defines the parent clade containing B.1.526 and related
106 viruses (Fig. 2a). We observed a stepwise emergence of the key lineage-defining mutations, with
107 T95I, D253G, and L5F appearing in the earliest phylogenetic nodes. Isolates subsequently
108 branched into four sub-lineages, with two major groups B.1.526-E484K and B.1.526-S477N
109 containing A701V, with a smaller sub-lineage B.1.526-S477N containing Q957R. The B.1.526-

110 L452R lineage, which also emerged in parallel, is related to B.1.526-E484K and B.1.526-S477N
111 yet forms a distinct phylogenetic branch (Fig. 1c).

112
113 Fig. 2b displays the localization of signature B.1.526-E484K and B.1.526-S477N mutations within
114 the S protein. D253G resides in the antigenic supersite within the N-terminal domain²⁴, which is a
115 target for neutralizing antibodies²⁵, whereas E484K is situated at the RBD interface with the
116 cellular receptor ACE2. The A701V mutation near the furin cleavage site is also shared with
117 variant B.1.351.

118
119 **Antibody neutralization of B.1.526**

120 The impact of the signature S protein mutations in B.1.526 on antibody neutralization was first
121 assessed using vesicular stomatitis virus (VSV)-based pseudoviruses as previously described^{4,25}.
122 Pseudoviruses containing S477N or E484K alone and all five signature mutations (L5F, T95I,
123 D253G, A701V, and E484K or S477N), termed NYΔ5(E484K) or NYΔ5(S477N), were
124 constructed and subjected to neutralization by 12 monoclonal antibodies including 5 with
125 emergency use authorization, 20 convalescent plasma, and 22 vaccinee sera. The specifics of these
126 monoclonal antibodies and clinical specimens were previously reported⁴. The neutralizing activity
127 of 12 monoclonal antibodies covering a range of epitopes on RBD was essentially unaltered
128 against the S477N and NYΔ5(S477N) pseudoviruses (Extended Data Fig. 3a) showing that this
129 mutation has no discernible antigenic impact, as was confirmed using convalescent plasma and
130 vaccinee sera (Extended Data Fig. 3b). However, against E484K and NYΔ5(E484K)
131 pseudoviruses, the activities of several antibodies were either impaired or lost, including
132 REGN10933 and LY-CoV555 that are already in clinical use (Fig. 3a). Likewise, neutralizing
133 activities of convalescent plasma or vaccinee sera were lowered by 4.1-fold or 3.3-3.6-fold,
134 respectively, against NYΔ5(E484K) (Fig. 3b). Neutralization studies of the authentic B.1.526-
135 E484K virus yielded similar results, although the magnitude of resistance to convalescent plasma
136 or vaccinee sera was slightly lower at 2.6-fold or 1.8-2.0-fold, respectively (Fig. 3b). A
137 comparative analysis with other variants of concern (Fig. 3c) showed that such risks are likely
138 lower than B.1.351 and closer to P.1. Overall, these results demonstrate the need to modify our
139 antibody therapy and to monitor the efficacy of current vaccines in regions where B.1.526-E484K
140 is prevalent.

141

142 **B.1.526 surge across New York and the US**

143 Prevalence of the novel variant B.1.526 surged alarmingly in our hospital catchment area (Fig. 4a)
144 and throughout New York State (Fig. 4b) after its emergence in late 2020, replacing other lineages
145 and initially outpacing B.1.1.7. A multinomial logistic regression model describing the concurrent
146 growth rates of these two lineages shows that starting in mid-April 2021, B.1.1.7 surpassed
147 B.1.526 due to a slightly higher fitness, with estimated growth rates in New York State of 5.3%
148 per day for B.1.1.7 (95% CI 5.0–5.7%) and 3.4% per day for B.1.526 (3.2–3.6%) (Fig. 4b). These
149 estimates suggest a fitness advantage of B.1.526 over existing non-variant lineages of 22–25%
150 over a serial interval of 7 days^{21,22} during a period when multiple variants are competing
151 simultaneously. Furthermore, the estimates also suggest a fitness advantage of B.1.1.7 over
152 existing non-variant lineages of 35–40%, as well as a fitness advantage of B.1.1.7 over B.1.526 of
153 12–15%. Both lineages grew quickly (Fig. 4a,b), but once they reached a high frequency of
154 circulating viruses, the competition between them caused the growth of B.1.1.7 to slow and
155 B.1.526 to decline.

156

157 Frequency trajectories of B.1.1.7 and B.1.526 across states (Fig. 4c, Extended Data Fig. 4) show
158 two general patterns: (1) initial rapid increase of both lineages until the proportion of other lineages
159 had been eclipsed, followed by decline of B.1.526 seen in New York and in several neighboring
160 states; and (2) rapid growth and resulting dominance of B.1.1.7 preventing the further rise of
161 B.1.526. The dynamics between these two lineages is further shown in Fig. 4d, which plots the
162 logistic growth rate of B.1.526 against the frequency of B.1.1.7, again at the state-level. At lower
163 frequencies of B.1.1.7, all states have a similarly rapid growth of B.1.526 as it replaces non-variant
164 lineages. As B.1.1.7 increases in frequency, however, it slows the growth of B.1.526, again
165 indicative of a slightly higher fitness for B.1.1.7. At a minimum, B.1.526 rose rapidly where
166 B.1.1.7 was not already dominant and, in several states, continued to grow at a similar pace as
167 B.1.1.7 (Extended Data Fig. 4).

168

169 Phylogeographic analysis of the B.1.526 lineage revealed ancestral viruses originating in New
170 York in August 2020, diversifying within the state, and then dispersing to other states (Figs. 4e
171 and 4f). State-level genomic data showed that B.1.526 was concentrated primarily in New York

172 and surrounding states, including New Jersey and Rhode Island (Extended Data Fig. 4). This
173 suggests that B.1.526, and B.1.526-E484K in particular, became widespread in the region, the
174 original epicenter of COVID-19 in the US^{26,27}, although the lineage has also grown in states outside
175 the Northeastern US (e.g., North Carolina). By the end of April 2021, the geographic makeup of
176 B.1.526 within the US was quite diverse, and the lineage has emerged and expanded in multiple
177 states across the country (Fig. 4f). The rise of B.1.526 over a short timeframe across the United
178 States (Extended Data Fig. 4), as well as its international spread, are notable.

179

180 **Discussion**

181 Here we report the emergence of the SARS-CoV-2 lineage B.1.526, and its surge in New York
182 during the second wave of the COVID-19 pandemic. Neutralization studies on B.1.526-E484K
183 demonstrate that the activities of several antibodies were either impaired or lost, including two
184 (Ly-CoV555 and REGN10933) already in clinical use. Furthermore, neutralizing activities of
185 convalescent plasma or vaccinee sera were lower against B.1.526 harboring E484K (Fig. 3b). The
186 S477N mutation, a key signature of another B.1.526 sub-lineage, on the other hand, did not have
187 an impact on antibody neutralizing.

188

189 Several limitations of our study need to be considered. This was a single-center genomic survey
190 representing patients presenting to a hospital system and may not have fully captured patients with
191 milder disease. However, our results are comparable to genomic data released by public health
192 laboratories in the region and incorporate all publicly available data for phylogeographic context
193 and growth rate calculations. As in all genomic surveillance studies, we predominantly sequenced
194 samples with a Ct<30 but covered a high proportion of samples throughout the study period. In
195 addition, our PCR screen allowed us to obtain unbiased estimates of E484K and N501Y prevalence
196 early on in the study. PCR approaches may be increasingly warranted for continued surveillance
197 during non-surge periods, during which Ct values trend higher. Lastly, transmissibility estimates
198 based on observed prevalence are imperfect as they reflect observed growth rates rather than
199 intrinsic transmissibility of the virus.

200

201 Taken together, our findings underscore the importance of the E484K mutation, which has
202 emerged in at least 246 different lineages of SARS-CoV-2²⁸, a real testament to convergent

203 evolution. This highlights that E484K can rapidly emerge in multiple clonal backgrounds and may
204 warrant targeted screening for this key mutation in addition to robust genomic surveillance
205 programs. However, B.1.526 is one of the few lineages with E484K that has risen to prominence.
206 The greatest threat of B.1.526 appears to be its ease of spread, with an estimated transmissibility
207 of ~35% greater than non-variant viruses when competing head-to-head. Despite the notable
208 transmissibility of B.1.1.7, B.1.526 was able to spread rapidly in the US to replace other lineages
209 and continued to increase in frequency in several states where both B.1.526 and B.1.1.7 were
210 predominant. Similarly, while B.1.351 may pose the greatest antigenic challenge to antibodies and
211 vaccines, the B.1.526-E484K sub-lineage also exhibits resistance to antibody neutralization. The
212 findings herein present a clear-cut example of SARS-CoV-2 evolution in real time. B.1.526, with
213 its higher transmissibility, appeared suddenly and rose to dominance, only to wane as variants
214 (B.1.1.7, and B.1.617.2 more recently) with even greater fitness emerged. These observations are
215 a stark reminder that if SARS-CoV-2 is allowed to continue its spread, increasingly worrisome
216 variants are to be expected in the future.

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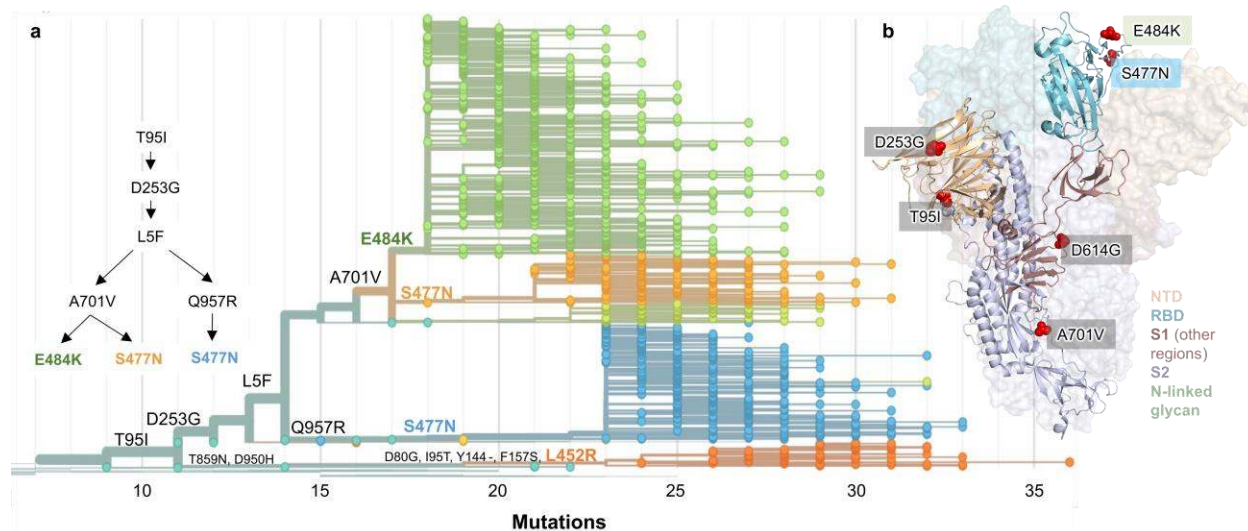
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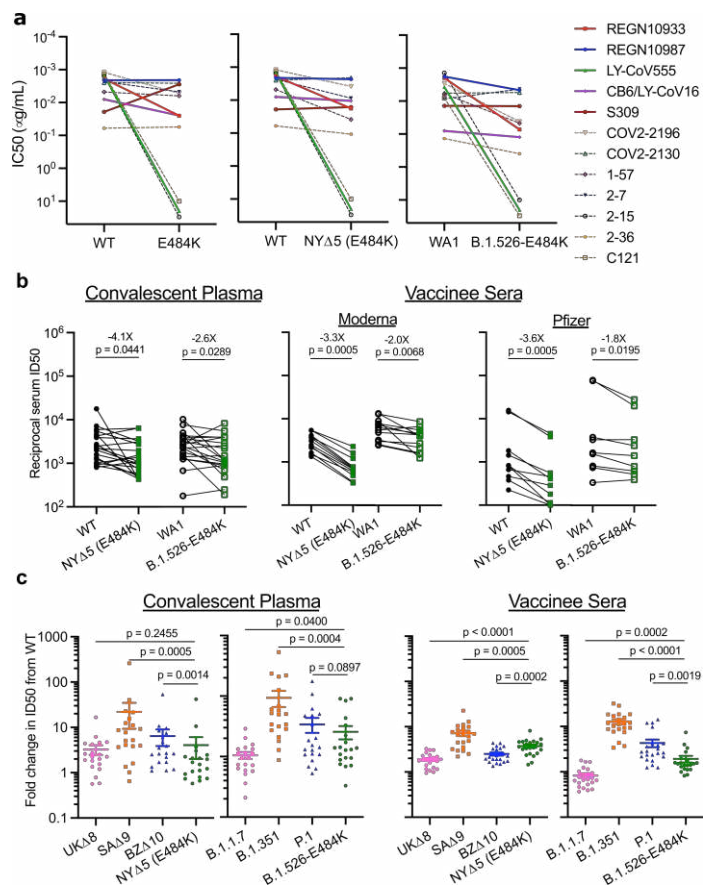
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- 324
- 325

327 **Figure 1. Prevalence of E484K-harboring SARS-CoV-2 and B.1.526.** (a) Detection of viruses
328 with key signature mutations in spike over time. The earliest detected E484K-harboring variant
329 was collected in mid-November 2020. The prevalence of E484K (samples with E484K/total PCR-
330 genotyped samples) subsequently increased over time, from 4.8% in early December 2020 up to
331 24.3% in early March 2021. Throughout late 2020 and early 2021, we identified fewer N501Y-
332 than E484K-harboring isolates, with a maximum of 5.9% of N501Y during mid-February 2021.
333 (b) Distribution of different viral lineages identified by whole genome sequencing. Within our
334 genomic collection (n=1,507), the B.1.526 lineage rose rapidly in early 2021, replacing the
335 majority of other lineages (shown as the white blank space) present during this timeframe. This
336 was followed by a steady rise in B.1.1.7 by mid-2021. The marking below the X axis denotes the
337 time-period used to calculate the growth advantage of B.1.526 over other earlier viruses. (c)
338 Phylogenetic tree of SARS-CoV-2 variants identified by sequencing and alignment of key spike
339 mutations. Unique patterns of spike protein mutations present in genomes sequenced from our
340 hospital center with at least one mutation of interest or concern (E484K, N501Y, S477N, or
341 L452R; n=64) are shown. Residues at which at least one sample harbored a mutation are displayed
342 above the S-protein schematic.
343
344

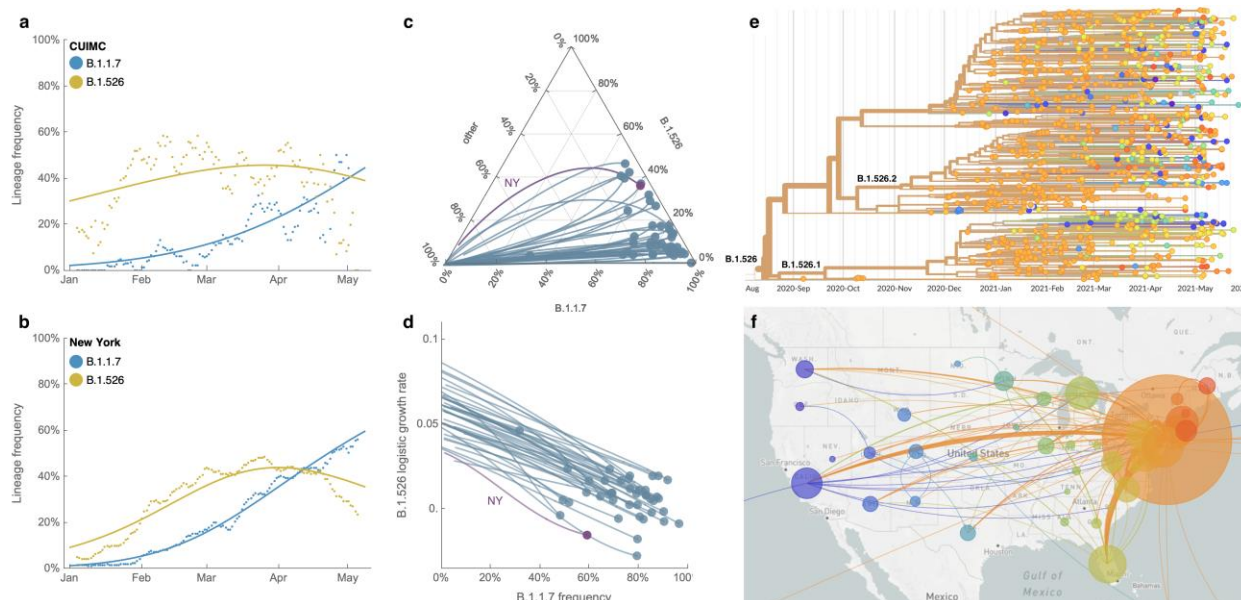


345
346 **Figure 2. Spike protein amino acid substitutions and structural changes represented in**
347 **sequenced isolates.** (a) Maximum-likelihood phylogenetic tree of 2,309 SARS-CoV-2 viruses
348 colored according to spike protein haplotype. Spike protein mutations are labeled on the tree
349 showing the stepwise accumulation of signature B.1.526 mutations T95I, D253G, and L5F, and
350 branching of B.1.526-E484K (green) and two B.1.526-S477N sub-lineages (light orange, blue).
351 The B.1.526-L452R sub-lineage (dark orange) emerged in parallel. An interactive version of this
352 figure is available at <https://nextstrain.org/groups/blab/ncov/ny/B.1.526>. (b) Key mutations of
353 B.1.526 displayed on the spike trimer. The D253G mutation resides in the antigenic supersite
354 within the N-terminal domain (NTD), a target for neutralizing antibodies, E484K and S477N at
355 the receptor binding domain (RBD) interface with the cellular receptor ACE2, and A701V near
356 the furin cleavage site.



357
358 **Figure 3. Neutralization studies of B.1.526-E484K and comparative analyses.** (a) Neutralizing
359 activities of 12 monoclonal antibodies against pseudoviruses containing E484K alone or all five
360 signature B.1.526 mutations (L5F, T95I, D253G, A701V, and E484K), termed NYΔ5(E484K) as
361 well as against the authentic B.1.526-E484K. Antibodies with emergency use authorization are
362 shown in bold solid lines. Data are represented as mean ± SEM of technical triplicates and
363 represent one of two independent experiments. (b) Neutralizing activities of convalescent plasma
364 (n=20) and vaccinee sera (n=22) against the NYΔ5(E484K) pseudovirus compared to wildtype
365 pseudovirus as well as against authentic B.1.526-E484K and wildtype virus (WA1). (c) Fold
366 change in convalescent plasma and vaccinee sera neutralization ID50 of different variant
367 pseudoviruses and live viruses compared to wildtype counterparts. The data on B.1.1.7, B.1.351
368 and P.1 were derived from our prior publications^{4,18}. Data from 20 convalescent patients or 22
369 vaccinated individuals were averaged and are represented as arithmetic mean ± SEM (individual
370 data points also shown). Statistical comparisons were made using the Wilcoxon matched-pairs
371 signed rank test; two-tailed p-values are reported.
372

373



374

375 **Figure 4. Spread of lineages B.1.1.7 and B.1.526 in New York and the USA.** (a, b) Frequencies
376 of lineages B.1.1.7 (blue) and B.1.526 (yellow) in the CUIMC catchment area (in panel a) and
377 New York State (in panel b) with dots representing daily 7-day sliding window averages and lines
378 representing fit to a multinomial logistic regression model. (c) Ternary plot of state-level frequency
379 trajectories for 42 states separating frequencies of B.1.1.7, B.1.526 and other lineages. Each state-
380 level trajectory is a line in this plot moving from lower left in January 2021 when both B.1.1.7 and
381 B.1.526 were rare, rightward as B.1.1.7 and B.1.526 increase in frequency. The trajectory of New
382 York State is highlighted in purple. (d) The same data as in panel c, except plotting frequency of
383 B.1.1.7 against logistic growth rate of B.1.526. (e) Phylogenetic tree of 933 B.1.526 samples from
384 across the US where branch tips are colored based on location of sampling and branches are
385 colored by inferred ancestral location. (f) Phylogeographic view of data from panel e, where each
386 sampling location is represented as a circle with area proportional to sample count and each
387 inferred transition event across the phylogeny is drawn as an arc connecting inferred origin and
388 destination. Most migration events are inferred to be direct dispersals from New York State.

389 **Methods**

390 **Clinical cohort.** This observational study took place at an academic quaternary care center in New
391 York City. Nasopharyngeal swabs obtained as part of routine clinical care were tested by the
392 Clinical Microbiology laboratory, and positive specimens were transferred to the Columbia
393 University Biobank for inactivation and storage. Electronic health records data extracted for this
394 analysis included demographics, laboratory results, admission, discharge, and transfer dates,
395 current and historical international classification of disease (ICD 9 and 10) codes extracted from
396 the clinical data warehouse. This study was reviewed and approved by the Columbia University
397 Institutional Review Board (protocol number AAAT0123).

398
399 **PCR screening.** Extended Data Figure 1 describes our overall protocol for variant screening. To
400 enable rapid PCR-based screening, we prepared RNA using the heat inactivation method in place
401 of RNA isolation methods²⁹. First, 50 μ l of nasal swab sample in VTM solution was transferred
402 into 96-well PCR plates, covered with an adhesive aluminum foil (VWR 60941-076) and
403 incubated at 95°C for 5 min using the PCR instrument. After the centrifugation of the plate at
404 $>2,100 \times g$ for 5 min, 5 μ l of the supernatant from each sample, which contains viral RNA, was
405 used for the SNP assay.

406
407 The SNP assay consists of four steps as follows: reverse transcription (RT) of viral RNA, pre-read
408 of the SNP assay, real-time PCR and post-read of the SNP assay. 5 μ l of RNA from the supernatant
409 was added to 15 μ l of the single step RT-qPCR reaction mix, which consists of 5 μ l of TaqPath 1-
410 step RT-qPCR Master Mix, CG (4x) (ThermoFisher Scientific), 500 nM of forward and reverse
411 primers, 120 nM of VIC-MGB probe, 50 nM of FAM-MGB probe, 1/2000 volume of ROX
412 Reference Dye (Invitrogen) as the final concentration, and nuclease-free water to adjust the total
413 reaction volume of 20 μ l. Each reaction plate included 8 control wells, 5×10^6 and 5×10^3 copies of
414 WA-1 (wild type), UK variant and South African variant, which were generated by PCR to match
415 the variant sequences, and 2 wells with water as no template controls (NTC).

416
417 The primer pairs and probes used are as follows. For the SNP assay for position **501**, a primer
418 pair of 501.F: 5'-GGT TTT AAT TGT TAC TTT CCT TTA CA-3' and 501.R: 5'-AGT TCA
419 AAA GAA AGT ACT ACT ACT CTG TAT G-3' were used with two TaqMan probes

420 (ThermoFisher Scientific), one for wild type, VIC.N501MGB: [VIC]-AA CCC ACT AAT
421 GGT-MGBNFQ and the other for variant type, FAM.Y501MGB: [FAM]-AAC CCA CTT ATG
422 GT-MGBNFQ. For position **484**, a primer pair of 484.F: 5'-AGA GAG ATA TTT CAA CTG
423 AAA TCT ATCAGG-3' and 484.R: 5'-GAA ACC ATA TGA TTG TAA AGG AAA GTA AC-
424 3' were used with two probes, one for wild type, VIC.E484MGB: [VIC]-ATG GTG TTG AAG
425 GT-MGBNFQ and the other for variant type, FAM.K484MGB: [FAM]-ATG GTG TTA AAG
426 GT-MGBNFQ.

427
428 The reaction plate was subjected to 1) reverse-transcription reaction (RT) at the condition at 25°C
429 for 2 min, at 50°C for 15 min and a hold at 4°C; 2) SNP assay (pre-read) at 60°C for 30 sec; 3)
430 real-time PCR at 95°C for 20 sec followed by 50 cycles of two-step PCR, at 95°C for 3 sec and at
431 60°C for 30 sec with the fast 7500 mode; followed by 4) SNP assay (post-read) at 60°C for 30 sec
432 using ABI 7500 Fast Dx Real-Time PCR Instrument with SDS Software (ThermoFisher
433 Scientific). The genotype at each key position for each sample was determined by reading the
434 component signal of the amplification and the allelic discrimination analysis software in the
435 program.

436
437 **Whole genome sequencing.** Extended Data Fig. 2 displays a flowchart outlining samples available
438 for this study. Isolates with cycle threshold (Ct) values below 35 were selected for sequencing
439 using the ARTIC v3 low-cost protocol targeting 400bp amplicons³⁰ or Rapid Barcoding kit
440 protocol targeting 1,200bp amplicons³¹. Briefly, RNA was extracted using the Qiagen RNeasy
441 Mini kit or Zymo DNA/RNA Mini kit. Reverse transcription was performed using LunaScript RT
442 SuperMix (NEB). Tiling PCR was performed on the cDNA, and amplicons were barcoded using
443 the Oxford Nanopore Native Barcoding Expansion 96 kit. Pooled barcoded libraries were then
444 sequenced on an Oxford Nanopore MinION sequencer using R9.4.1 flow cells. Basecalling was
445 performed in the MinKNOW software v21.02.1. Sequencing runs were monitored in real-time
446 using RAMPART (<https://artic-network.github.io/rampart/>) to ensure sufficient genomic coverage
447 with minimal runtime. Consensus sequence generation was performed using the ARTIC
448 bioinformatics pipeline (<https://github.com/artic-network/artic-ncov2019>). Genomes were
449 manually curated by visually inspecting sequencing alignment files for verification of key residues
450 in Geneious v10.2.6.

451
452 **Phylogenetic analysis.** Phylogenetic reconstruction of amino acid changes (Fig. 2A) was
453 conducted using the Nextstrain³² workflow at <https://github.com/nextstrain/ncov> which aligns
454 sequences against the Wuhan-Hu-1 reference via nextalign
455 (<https://github.com/nextstrain/nextclade>), constructs a maximum-likelihood phylogenetic tree via
456 IQ-TREE³³, estimates molecular clock branch lengths via TreeTime³⁴ and reconstructs nucleotide
457 and amino acid changes also via TreeTime. This workflow was applied to 2309 SARS-CoV-2
458 genomes possessing the 9bp deletion Δ 106-108 in ORF1a-nsp6 along with mutation A20262G
459 which demarcates the parent clade to lineage B.1.526 alongside 688 global reference viruses. This
460 analysis was conducted on data downloaded from gisaid.org³⁵ on April 5, 2021. Phylogeographic
461 reconstruction of spread from New York state (Fig. 4E-F) was similarly conducted using the same
462 Nextstrain workflow with the addition of performing ancestral trait reconstruction of the
463 geographic “division” attribute of 933 SARS-CoV-2 genomes downloaded from gisaid.org on Jun
464 6, 2021.

465
466 **Neutralization studies of pseudoviruses.** We assayed the neutralizing activity of monoclonal
467 antibodies (mAbs), convalescent plasma, and vaccinee sera against E484K, S477N, and WT
468 (D614G) pseudoviruses, as well as pseudovirus NY Δ 5 containing all five signature mutations of
469 B.1.526-E484K (L5F, T95I, D253G, E484K, D614G, A701V), as previously described²⁵. We
470 examined four mAbs with emergency use authorization (CB6, REGN10987, REGN10933 and
471 LY-CoV555), plus eight additional RBD mAbs, including ones from our own collection (2-15, 2-
472 7, 1-57, & 2-36)²⁵ as well as S309³⁶, COV2-2196 & COV2-2130³⁷, and C121³⁸. We also examined
473 convalescent plasma collected in Spring of 2020 (n=20 patients), and Moderna and Pfizer vaccinee
474 sera (n=22)⁴. Briefly, Vero E6 cells (ATCC) were seeded in 96-well plates (2×10^4 cells per well).
475 Pseudoviruses were incubated with serial dilutions of the test samples in triplicate for 30 min at
476 37 °C. The mixture was added to cultured cells and incubated for an additional 24 h. Luminescence
477 was measured using a Britelite plus Reporter Gene Assay System (PerkinElmer), and IC₅₀ was
478 defined as the dilution at which the relative light units were reduced by 50% compared with the
479 virus control wells (virus + cells) after subtraction of the background in the control groups with
480 cells only. The IC₅₀ values were calculated using nonlinear regression in GraphPad Prism 8.0.

481 Statistical analysis was performed using a Wilcoxon matched-pairs signed rank test. Two-tailed p-
482 values are reported.

483
484 **Neutralization of infectious SARS-CoV-2.** Infectious SARS-CoV-2 isolate hCoV-19/USA/NY-
485 NP-DOH1/2021 was isolated at the Aaron Diamond AIDS Center (Columbia University Medical
486 Ctr) from nasopharyngeal swab and propagated for one passage in Vero E6 cells (ATCC).
487 Infectious titer of the resulting virus was determined by an end-point dilution and cytopathic effect
488 (CPE) assay on Vero-E6 cells as described previously²⁵. The virus has since been deposited at BEI
489 Resources (Cat#NR-55359). SARS-CoV-2 virus USA-WA1/2020 (WA1) obtained from BEI
490 Resources (Cat# NR-52281) served as the control in experiments.

491
492 An end-point dilution microplate neutralization assay was performed to measure the neutralization
493 activity of twenty patient convalescent plasma samples and twelve purified monoclonal antibodies.
494 In brief, plasma samples were subjected to successive 5-fold dilutions starting from 1:100.
495 Similarly, antibodies were serially diluted (5-fold dilutions) starting at 50 µg/ml. Triplicates of
496 each dilution were incubated with SARS-CoV-2 at an MOI of 0.1 in EMEM with 7.5% inactivated
497 fetal calf serum (FCS) for 1 hour at 37°C. Post incubation, the virus-antibody mixture was
498 transferred onto a monolayer of Vero-E6 cells grown overnight. The cells were incubated with
499 the mixture for ~70 hours. Cytopathic effect (CPE) of viral infection was visually scored for each
500 well in a blinded fashion by two independent observers. The results were then converted into
501 percentage neutralization at a given sample dilution or antibody concentration, and the averages ±
502 SEM were plotted using a five-parameter dose-response curve in GraphPad Prism v8.4.

503
504 **Growth dynamics.** Growth dynamics of B.1.1.7 and B.1.526 were obtained through by
505 downloading “metadata” from gisaid.org on June 6, 2021 for all 422,760 viruses sampled from the
506 USA collected after January 1, 2021. This metadata has PANGO lineages³⁹ already assigned to
507 each genome sequence. Daily state-level frequencies (and frequencies for CUIMC) were extracted
508 for plotting via 7-day sliding window averages of the prevalence of B.1.1.7 and B.1.526, calculated
509 as the number of sequence-verified samples from each strain divided by the total number of
510 positive samples with cycle threshold (Ct) values below 35, as this threshold value was used for
511 sequencing. Separately, a multinomial logistic regression model was fit directly to the observation

512 data consisting of individual genomes, their dates of sampling (independent variable X in days
513 since January 1, 2021) and their categorical labels (dependent variable Y , “B.1.1.7”, “B.1.526” and
514 “other”). This results in a 4-parameter model where both B.1.1.7 and B.1.526 have parameters
515 specified for frequency at day 0 (January 1, 2021) and logistic growth rate. This model was fit to
516 the data using the Classify package of Mathematica v12.2.

517
518 **Data availability.** All genomes and associated metadata generated as a part of this study have been
519 uploaded to GISAID (gisaid.org) and NCBI GenBank (BioProject Accession PRJNA751551).
520 Biological materials (i.e. variant pseudoviruses) generated as a part of this study will be made
521 available but may require execution of a materials transfer agreement.

522
523 **Code availability.** Data processing and visualization was performed using publicly available
524 software and packages, primarily RStudio v1.2.5033, GraphPad Prism v8.4, and iTOL
525 (<https://itol.embl.de/>). The exact workflow used for phylogenetic (Fig. 2A) and phylogeographic
526 analysis of public GISAID data (Fig. 4E-F) is available at <https://github.com/blab/ncov-ny>.
527 Frequency dynamics were modeled using Mathematica in notebooks also available at
528 <https://github.com/blab/ncov-ny>.

529
530 **Methods References**

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555 genomic epidemiology. *Nat Microbiol* 5, 1403-1407, doi:10.1038/s41564-020-0770-5
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557
558
559

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575
576 **Competing Interests:** P.W., M.S.N., Y.H., and D.D.H. are inventors on a provisional patent
577 application on monoclonal antibodies against SARS-CoV-2. D.D.H. is a member of the scientific
578 advisory board of Bria Biosciences, which has provided a grant to Columbia University to support
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580 Co. unrelated to this study.

581
582 **Author Contributions: Conceptualization** – A.-C.U., D.D.H., M.K.A., H.M.; **Data curation** –
583 M.K.A., H.M., J.E.Z., P.W., M.S.N., Z.S., T.B., A.G.-S., Y.H., A.L.K., M.T., A.-C.U.; **Formal**
584 **analysis** – M.K.A., P.W., J.E.Z., T.B., A.G.-S.; **Funding acquisition** – A.-C.U., D.D.H., M.K.A.;
585 **Investigation** – M.K.A., H.M., J.E.Z., P.W., M.S.N., A.L.K., M.T., T.B., Y.H.; **Methodology** –
586 M.K.A., H.M., P.W., M.S.N., T.B., Y.H.; **Supervision** – A.-C.U., D.D.H.; **Visualization** –
587 M.K.A., P.W., T.B.; **Writing – original draft** – A.-C.U., M.K.A., H.M., D.D.H.; **Writing –**
588 **review and editing** – all authors

589
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591 (au2110@cumc.columbia.edu) or David D. Ho (dh2994@cumc.columbia.edu).

592

593 **Extended Data**

Extended Data Table 1. Clinical characteristics of patients infected with SARS-CoV-2 based on viral genotype

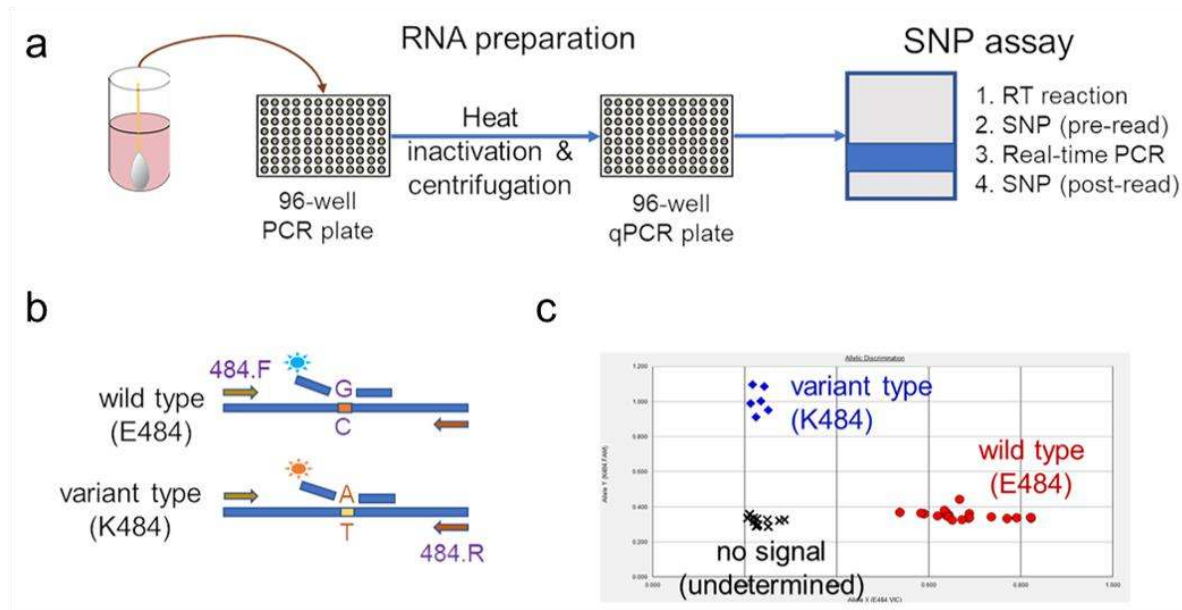
Clinical Characteristic	E484K (n=170) ¹	Wildtype ¹ (n=1,180)	P ²	Non-VOI/VOC lineages ¹		P ²
				B.1.526-E484K ¹ (n=232)	(n=790)	
Demographics						
Male sex, n (%)	80 (47.1)	539 (45.7)	0.806	97 (41.8)	373 (47.3)	0.159
Age, years (median [IQR])	56 [36, 68]	55 [33, 70]	0.965 ³	51 [32, 69]	51 [30, 68]	0.743 ³
Race and ethnicity, n (%)			0.288			0.117
Hispanic/Latino	87 (51.2)	548 (46.5)		122 (52.6)	357 (45.3)	
Black	11 (6.5)	131 (11.1)		27 (11.6)	89 (11.3)	
White	24 (14.1)	172 (14.6)		23 (9.9)	119 (15.1)	
Other	48 (28.2)	328 (27.8)		60 (25.9)	223 (28.3)	
Place of residence, n (%)			0.008			0.002
NYC	151 (88.8)	959 (81.3)		204 (87.9)	627 (79.4)	
Yonkers	7 (4.1)	34 (2.9)		9 (3.9)	23 (2.9)	
Outside NYC and Yonkers	12 (7.1)	187 (15.8)		19 (8.2)	140 (17.7)	
Comorbidities						
BMI, kg/m ² (median [IQR])	28.9 [25.0, 33.1]	27.1 [23.5, 31.3]	0.01 ³	28.6 [24.7, 33.1]	26.6 [23.1, 30.5]	0.001 ³
Hypertension, n (%)	70 (41.7)	444 (41.2)	0.981	86 (38.4)	291 (40.6)	0.602
Diabetes mellitus, n (%)	51 (30.4)	265 (24.6)	0.134	63 (28.1)	166 (23.2)	0.157
Chronic kidney disease, n (%)	21 (12.5)	126 (11.7)	0.864	18 (8.0)	93 (13.0)	0.059
Coronary artery disease, n (%)	13 (7.7)	108 (10.0)	0.428	16 (7.1)	72 (10.1)	0.240
Solid organ transplant, n (%)	7 (4.2)	42 (3.9)	1.00	7 (3.1)	41 (5.7)	0.171
Cycle threshold value (mean (SD))⁴	29.49 (5.64)	30.71 (5.66)	0.013	27.65 (4.88)	28.81 (4.93)	0.015
Severity of care and outcomes						
Highest level of care, n (%)			0.175			0.037
Admitted	50 (29.6)	417 (35.4)		81 (35.1)	228 (29.0)	
Emergency Department	64 (37.9)	339 (28.8)		83 (35.9)	243 (31.0)	
ICU	13 (7.7)	86 (7.3)		15 (6.5)	54 (6.9)	
Outpatient	42 (24.9)	335 (28.4)		52 (22.5)	259 (33.0)	
Supplemental oxygen, n (%)	48 (85.7)	348 (77.5)	0.217	67 (81.7)	187 (73.9)	0.199
Outcome, n (%)			0.454			0.565
Deceased or discharged to hospice	8 (4.7)	87 (7.4)		10 (4.4)	45 (5.7)	
Further care at external facility	16 (9.5)	85 (7.2)		16 (7.0)	44 (5.6)	
Discharged to home	145 (85.8)	1003 (85.4)		202 (88.2)	693 (88.5)	

¹ Wildtype isolates are defined as those without E484K or N501Y mutations. For comparisons based on lineage, B.1.526-E484K was compared with non-variant of interest (VOI) and non-variant of concern (VOC) lineages (i.e. B.1.526.1, B.1.526.2, B.1.1.7, P.1, P.2, and B.1.351 were excluded). Comparisons were made using the first genotyped sample per patient to exclude multiple measures.

² T-tests (two-sided) were performed for continuous variables and chi-squared tests for categorical variables, unless otherwise indicated as below

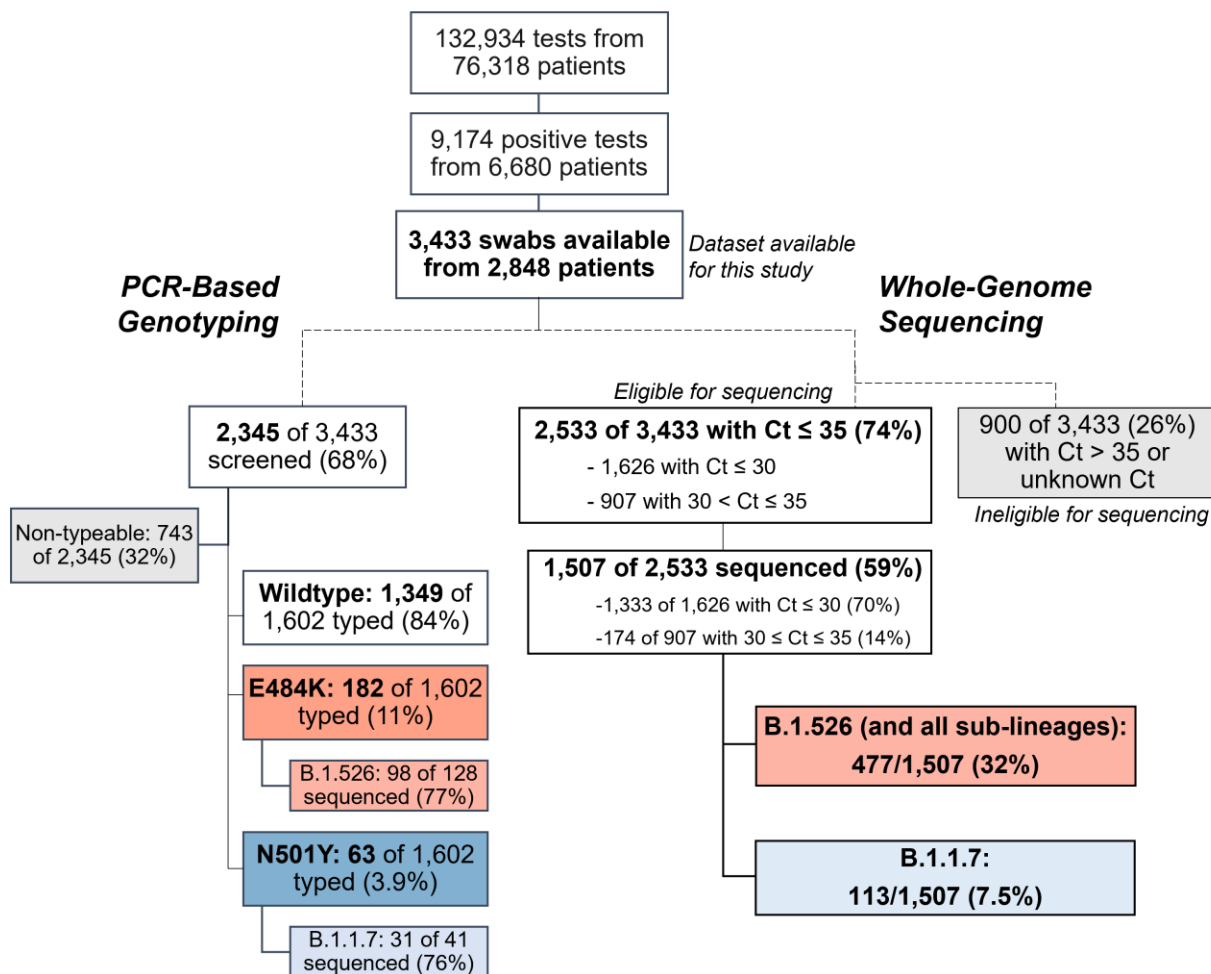
³ Due to non-normal distribution, Kruskal-Wallis non-parametric test was used

⁴ Cycle threshold value as determined through our rapid qPCR-based screening assay on heat-inactivated nasopharyngeal swab samples



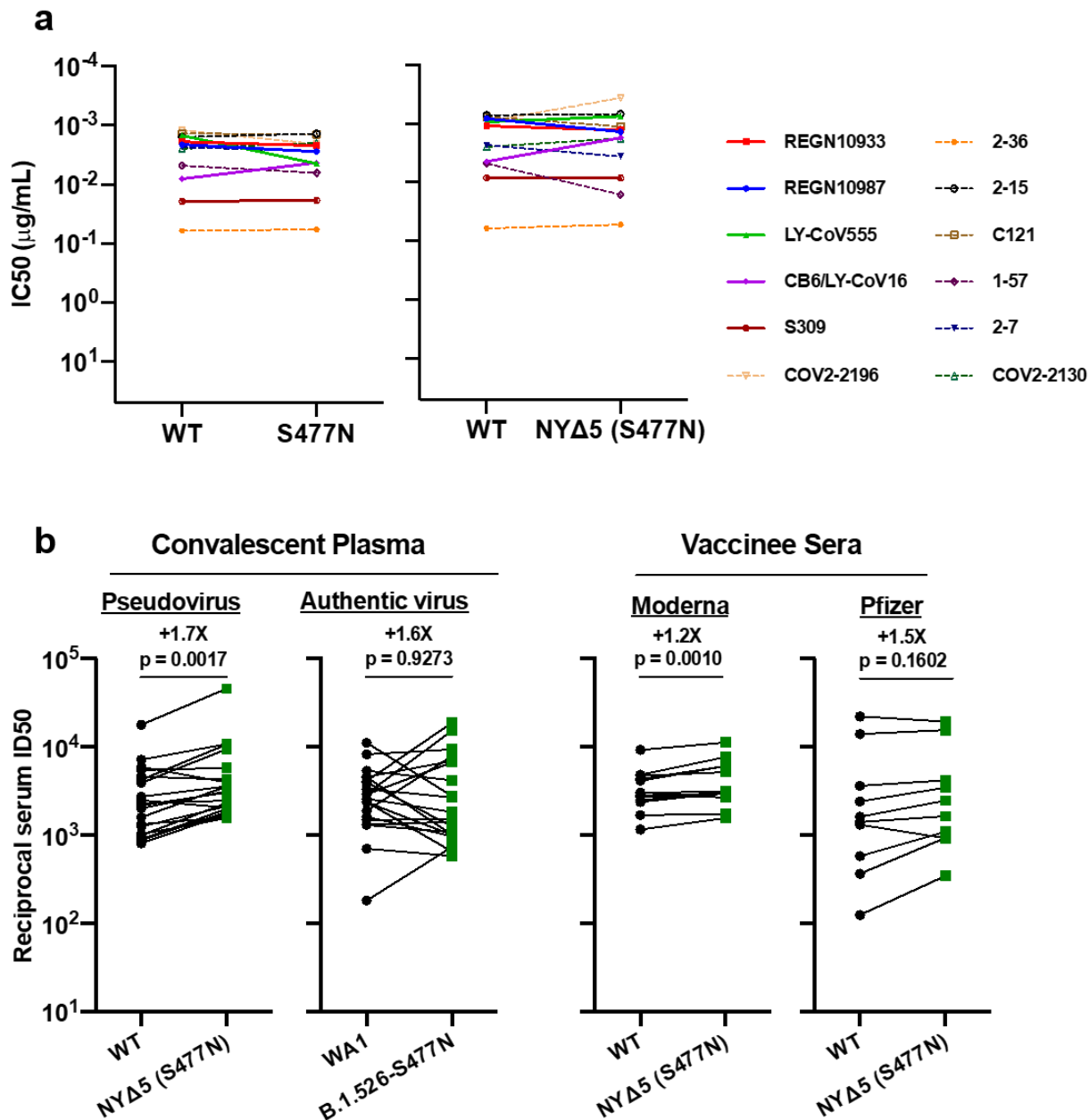
Extended Data Figure 1. Rapid PCR-based screening assay protocol to identify samples harboring key substitutions. (a) Viral RNA is prepared by heat inactivation and centrifugation. The supernatant is then used for the SNP assay, which entails four steps: the reverse transcription (RT) reaction, pre-PCR reading of the plate to assess background fluorescence (SNP pre-read), real-time PCR, and post-PCR reading of the plate to measure fluorescence (SNP post-read). The runtime for this entire protocol is approximately two hours. (b) Genotype at targeted sites in COVID-19 viral RNA can be determined with two MGB probes, one for wild type (conjugated with VIC) and the other for variant type (conjugated with FAM). (c) Example signals for the variant type (K484; blue), the wild type (E484; red) and samples with no signal (black) are shown.

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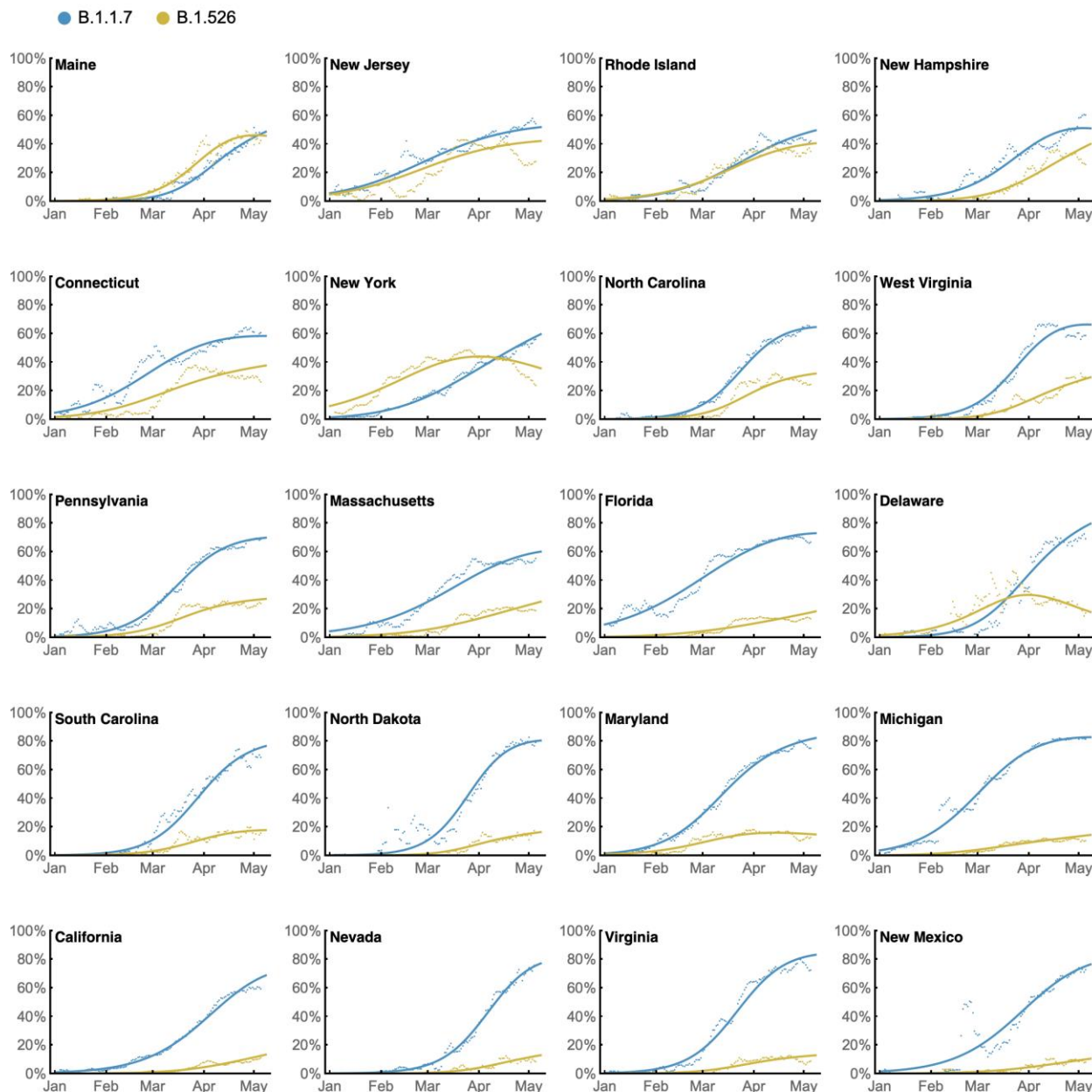
Note: 8 swabs identified with both E484K and N501Y were sequence-verified as P.1 (n=6); B.1.351 (n=1); and B.1.623 (n=1)

Extended Data Figure 2. Flowchart for SARS-CoV-2-positive nasopharyngeal swabs included in this study. (top) During the study period of November 1, 2020 to May 1, 2021, 6,680 patients tested positive for SARS-CoV-2 at our hospital center and affiliated hospitals. From these 9,174 positive nasopharyngeal swabs, 3,433 swabs were stored as part of the Columbia University Biobank COVID-19 sample repository and available for this study. (left) PCR-based genotyping assays for E484K and N501Y (see Extended Data Fig. 1) were performed on 2,345 samples. We identified a significant proportion of samples with E484K (11%), later confirmed through sequencing to primarily fall within the B.1.526 lineage, and a number of samples with N501Y (3.9%), primarily within the B.1.1.7 lineage. (right) We performed whole-genome sequencing on 1,507 samples. Of these, 32% belonged to B.1.526 and the sublineages B.1.526.1 and B.1.526.2, while B.1.1.7 constituted a much smaller proportion of samples at our center (7.5%).



Extended Data Figure 3. Neutralization studies of B.1.526-S477N. (a) Neutralizing activities of 12 monoclonal antibodies against pseudoviruses containing S477N alone or all five signature B.1.526-S477N mutations (L5F, T95I, D253G, A701V, and S477N), termed NY Δ 5(S477N). Antibodies with emergency use authorization are shown in bold solid lines. Data are represented as mean \pm SEM. of technical triplicates and represent one of two independent experiments. (b) Neutralizing activities of convalescent plasma (n=20) against NY Δ 5(S477N) as well as against the authentic B.1.526 virus with S477N, and neutralizing activities of vaccinee sera (n=22) against the NY Δ 5(S477N) pseudovirus, compared to wildtype counterparts. Statistical comparisons were made using the Wilcoxon matched-pairs signed rank test; two-tailed p-values are reported.

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599 **Extended Data Figure 4. State-level growth dynamics of B.1.526 and B.1.1.7.** Daily state-level
600 frequencies of B.1.526 (in yellow) and B.1.1.7 (in blue), based on GISAID data downloaded on
601 June 6, 2021, were used to plot 7-day sliding window averages of the prevalence of each lineage
602 (shown as dots in the figure). A 4-parameter multinomial logistic regression model was fit directly
603 to the observation data, in which both B.1.1.7 and B.1.526 have parameters specified for frequency
604 at day 0 (January 1, 2021) and logistic growth rate (shown as lines in the figure). States are ordered
605 according to frequency of B.1.526 at the final timepoint of May 8, 2021.