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Quantifying how single dose Ad26.COV2.S vaccine efficacy depends on Spike sequence features

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In the ENSEMBLE randomized, placebo-controlled phase 3 trial (NCT04505722), estimated single-dose Ad26.COV2.S vaccine efficacy (VE) was 56% against moderate to severe–critical COVID-19. SARS-CoV-2 Spike sequences were determined from 484 vaccine and 1,067 placebo recipients who acquired COVID-19. In this set of prespecified analyses, we show that in Latin America, VE was significantly lower against Lambda vs. Reference and against Lambda vs. non-Lambda [family-wise error rate (FWER) $p < 0.05$]. VE differed by residue match vs. mismatch to the vaccine-insert at 16 amino acid positions (4 FWER $p < 0.05$; 12 q -value ≤ 0.20); significantly decreased with physicochemical-weighted Hamming distance to the vaccine-strain sequence for Spike, receptor-binding domain, N-terminal domain, and S1 (FWER $p < 0.001$); differed (FWER ≤ 0.05) by distance to the vaccine strain measured by 9 antibody-epitope escape scores and 4 NTD neutralization-impacting features; and decreased ($p = 0.011$) with neutralization resistance level to vaccinee sera. VE against severe–critical COVID-19 was stable across most sequence features but lower against the most distant viruses.

Initial SARS-CoV-2 vaccine candidates were based on the virus's original lineage, as represented by the index strain with Spike D614 (NC_045512; <https://www.ncbi.nlm.nih.gov/nucore/1798174254>). As the virus has evolved, the efficacy of these vaccines against symptomatic infection has waned^{1,2}, and new vaccine inserts have been developed.

Based on data from a randomized, placebo-controlled vaccine efficacy (VE) trial on clinical outcomes and pathogen sequences isolated from participants experiencing clinical outcomes, sieve analysis assesses how VE depends on pathogen sequence features^{3,4}. Pajon et al.⁵ and Sadoff et al.⁶ showed how the VE against symptomatic COVID-19 was lower against certain variants than against the Reference strain in the phase 3 COVE trial of two doses of Moderna's mRNA-1273 vaccine and the phase 3 ENSEMBLE trial of a single dose of Janssen's Ad26.COV2.S vaccine, respectively. [As in ref. 6, Reference is defined as the basal outbreak lineage B.1, which bears the D614G mutation.] Cao et al. showed that VE was higher in COVID-19 VE trials where circulating viruses had shorter Spike sequence

Hamming distances to the vaccine strain⁷. These sieve analyses only considered Spike viral variation defined by the WHO-defined variant category or the unweighted Spike protein distance. They did not assess how VE depends on other Spike sequence features, such as at the level of individual mutations or features that impact immunological functions such as anti-SARS-CoV-2 neutralization^{8–13}, relevant given the strong evidence of neutralizing antibodies (nAbs) as a cross-platform correlate of protection^{14–16}.

In this work, we report the results of a sieve analysis of the ENSEMBLE trial (NCT04505722), which enrolled over 40,000 participants and was conducted in Argentina, Brazil, Chile, Colombia, Mexico, Peru, South Africa, and the United States^{6,17}. The sieve analysis considered baseline SARS-CoV-2 seronegative per-protocol participants and the primary endpoint (moderate to severe–critical COVID-19), as well as the severe–critical COVID-19 endpoint, during the double-blinded period of follow-up. The major conclusions of the current work are that in Latin America, where Spike diversity was greatest, VE differed by multiple Spike, receptor-binding domain

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(RBD), N-terminal domain (NTD), and S1 sequence features, as well as by distance to the vaccine strain as measured by multiple antibody-escape scores and neutralization-impacting features. Most of these significant sieve effects are linked to the Lambda lineage, implicating Lambda as a likely escape variant. Moreover, VE against severe-critical COVID-19 was generally stable across most sequence features, although it was lower against the most distant viruses.

Results

SARS-CoV-2 sequence data

A total of 1345 SARS-CoV-2 Spike amino acid sequences were obtained from 1224 participants experiencing the moderate to severe-critical primary endpoint. All sequences were variant-typed to either the Reference lineage or to one of nine different WHO-defined variants (Table 1, Fig. 1a, and Supplementary Table 5). In Latin America, lineages that circulated at the beginning of the study period, e.g., Reference, were closer to the sequence from the vaccine insert than later emerging lineages,

with Lambda the most distant (Fig. 1b and Supplementary Fig. 1). Similar results were obtained in South Africa and the United States (Supplementary Figs. 2, 3, respectively).

Greatest SARS-CoV-2 Spike diversity in Latin America

Most sequences were obtained from participants in Latin America ($n = 776$), with additional sequences from the US ($n = 323$) and South Africa ($n = 125$) (Supplementary Table 6). Participant demographics of the sieve analysis cohort are shown in Table 2 (Latin America) and Supplementary Tables 7–9 (US, South Africa, all regions pooled, respectively). Five main variants circulated in Latin America (Reference, Zeta, Gamma, Lambda, Mu), while the South African sequences were 76% Beta and 17% Delta, and the US sequences were 85% Reference (Fig. 1a). There was greater Spike AA sequence diversity in Latin America compared to South Africa and the US (Rao's $Q = 10.1$ vs. 7.7 vs. 3.3, respectively; Supplementary Fig. 4).

The succession of distinct co-circulating variants in Latin America and the resulting broadest dynamic range of inter-individual sequence

Table 1 | Numbers of primary endpoint COVID-19 cases with Spike amino acid sequence data by treatment arm, geographic region, and primary endpoint case lineage

Geographic Region	Latin America		South Africa		United States		Pooled	
	Vaccine (329) ^a	Placebo (634)	Vaccine (62)	Placebo (110)	Vaccine (93)	Placebo (323)	Vaccine (484)	Placebo (1067)
Primary endpoint case lineage								
Reference	72	196	1	4	52	221	125	421
Alpha	4	10	1	2	4	16	9	28
Beta	-	-	36	59	-	-	36	59
Delta	-	-	11	10	-	-	11	10
Epsilon	-	2	-	-	8	15	8	17
Gamma	73	111	-	-	1	-	74	111
Iota	-	-	-	-	-	4	0	4
Lambda	43	45	-	1	-	-	43	46
Mu	38	57	-	-	-	-	38	57
Zeta	33	92	-	-	1	1	34	93
No Sequence Obtained	66	121	13	34	27	66	106	221

A primary endpoint case is defined as the moderate to severe-critical primary COVID-19 endpoint in the per-protocol baseline seronegative cohort, with disease onset starting 14 days post-vaccination through to a participant's unblinding date.

^aNumbers in parentheses are numbers of moderate to severe-critical COVID-19 primary endpoints caused by the listed SARS-CoV-2 lineage, regardless of the availability of SARS-CoV-2 sequence data

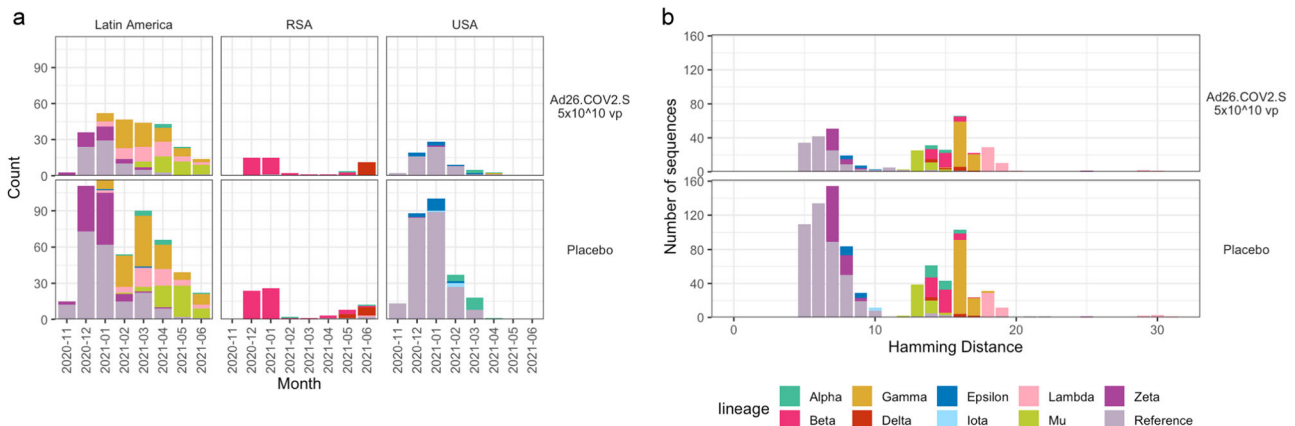


Fig. 1 | Diversity of circulating SARS-CoV-2 lineages. **a** The distribution of SARS-CoV-2 lineages of COVID-19 primary endpoints. The number of lineage sequences identified each month is shown for vaccine and placebo participants. **b** The distribution of variant sequences identified in Latin America as a function of their

Spike Hamming distance from the vaccine insert. Lineages are color-coded as follows: Alpha, teal; Beta, pink; Delta, red; Epsilon, dark blue; Gamma, mustard; Iota, light blue; Lambda, pink; Mu, light green; Zeta, purple; and Reference, lavender. vp viral particles.

Table 2 | Demographics of participants in Latin America in the sieve analysis cohort

Characteristics	Vaccine (N = 329)	Placebo (N = 634)	Total (N = 963)
Age			
Age 18–59	261 (79.3%)	523 (82.5%)	784 (81.4%)
Age ≥60	68 (20.7%)	111 (17.5%)	179 (18.6%)
Mean (Range)	43.4 (18, 83)	44.3 (18, 83)	44.0 (18, 83)
Sex assigned at birth			
Female	139 (42.2%)	270 (42.6%)	409 (42.5%)
Male	190 (57.8%)	364 (57.4%)	554 (57.5%)
BMI			
Underweight BMI <18.5	5 (1.5%)	3 (0.5%)	8 (0.8%)
Normal 18.5 ≤ BMI <25	103 (31.3%)	194 (30.6%)	297 (30.8%)
Overweight 25 ≤ BMI <30	72 (21.9%)	154 (24.3%)	226 (23.5%)
Obese BMI ≥30	149 (45.3%)	282 (44.5%)	431 (44.8%)
-	-	1 (0.2%)	1 (0.1%)
Ethnicity			
Hispanic or Latino	319 (97.0%)	611 (96.4%)	930 (96.6%)
Not Hispanic or Latino	6 (1.8%)	19 (3.0%)	25 (2.6%)
Unknown	4 (1.2%)	4 (0.6%)	8 (0.8%)
Race			
American Indian Or Alaska Native	101 (30.7%)	187 (29.5%)	288 (29.9%)
Asian	5 (1.5%)	2 (0.3%)	7 (0.7%)
Black Or African American	12 (3.6%)	22 (3.5%)	34 (3.5%)
White	146 (44.4%)	299 (47.2%)	445 (46.2%)
Multiple	50 (15.2%)	104 (16.4%)	154 (16.0%)
Other	15 (4.6%)	20 (3.2%)	35 (3.6%)
Country			
Argentina	45 (13.7%)	102 (16.1%)	147 (15.3%)
Brazil	109 (33.1%)	212 (33.4%)	321 (33.3%)
Chile	6 (1.8%)	9 (1.4%)	15 (1.6%)
Colombia	105 (31.9%)	209 (33.0%)	314 (32.6%)
Mexico	3 (0.9%)	9 (1.4%)	12 (1.2%)
Peru	61 (18.5%)	93 (14.7%)	154 (16.0%)
Risk for severe COVID-19^a			
At-risk	114 (34.7%)	211 (33.3%)	325 (33.7%)
Not at-risk	215 (65.3%)	423 (66.7%)	638 (66.3%)
Age, risk for severe COVID-19			
Age 18–59 at-risk	75 (22.8%)	152 (24.0%)	227 (23.6%)
Age 18–59 not at-risk	186 (56.5%)	371 (58.5%)	557 (57.8%)
Age ≥60 at-risk	39 (11.9%)	59 (9.3%)	98 (10.2%)
Age ≥60 not at-risk	29 (8.8%)	52 (8.2%)	81 (8.4%)
HIV status			
Negative	324 (98.5%)	623 (98.3%)	947 (98.3%)
Living with HIV	5 (1.5%)	11 (1.7%)	16 (1.7%)

^a “At-risk” is defined as having one or more comorbidities [listed in ref. 17] associated with elevated risk of severe COVID-19.

diversity, and the greatest number of COVID-19 endpoints, implies that sieve analyses of the Latin America region have the greatest statistical power. In contrast, the domination of the Reference lineage in the United States and the Beta and Delta lineages in South Africa constrained the sequence diversity’s dynamic range and limited the power of these sieve analyses. Therefore, we focus on the results from Latin America, with the United States and South Africa results reported in Supplementary Note 1.

Differential vaccine efficacy by SARS-CoV-2 lineage

All reported results on VE by SARS-CoV-2 features are based on feature-specific proportional-hazards models^{18,19} [see the Statistical Analysis Plan (SAP), provided in ref. 20 and at the end of the Supplementary Information]. Figure 2a shows VE against the primary COVID-19 endpoint caused by the Reference, Gamma, Zeta, Lambda, and Mu lineages, and Fig. 2b shows VE against the primary COVID-19 endpoint caused by the groupings of all other lineages excluding each individual lineage (referred to as not-lineage). Figure 2c shows differential VE against pairs of lineages or against pairs of lineage vs. not-lineage. VE was significantly higher against Reference than against Lambda and against not-reference lineages [family-wise error rate (FWER) $p < 0.05$]. It was also significantly higher against not-Lambda vs. Lambda and against Zeta vs. Lambda (FWER $p \leq 0.05$), and higher against Reference vs. Gamma, Reference vs. Mu, Zeta vs. Gamma, and Zeta vs. Mu (q value ≤ 0.20). Country-specific results for the same analyses are shown in Supplementary Figs. 5–7, where due to reduced sample size, the confidence intervals are generally much wider, and fewer pairwise comparisons of VE against lineages could be performed.

Higher vaccine efficacy against vaccine-matched AA residues

We scanned across all Spike AA positions with sufficient residue variability (at least 20 endpoints with a vaccine-mismatched residue: $n = 37$ positions). VE significantly differed (q value ≤ 0.20) by residue match vs. mismatch to the vaccine strain residue at 16 positions (Fig. 2d; four positions with FWER $p \leq 0.05$: 75, 76, 253, 490). Similarly, when assessing the presence or absence of specific residues at each AA position, VE significantly differed (q value ≤ 0.20) for 38 residues/residue features (75V vs. not-75V and 76I vs. not-76I with FWER $p \leq 0.05$) distributed across these 16 positions (Fig. 2e). Thirteen of these 16 positions harbored characteristic mutations of the Lambda variant and not for any other variants, and very highly covaried with Lambda vs. not-Lambda (Supplementary Fig. 8, $M_{\text{star}}^{21} > 0.85$), thereby providing nearly equivalent signatures of differential VE captured by Lambda vs. not-Lambda. The full results of the covariability analysis are in Supplementary Note 1. Four of the 1277 analyzed AA positions (417, 452, 484, 490) were prespecified as being hypothesized to impact neutralization based on an association with a reduced nAb response in mRNA vaccine recipients^{22–24}, or evidence for increased transmissibility (452)²² or increased infectivity in vitro (452, 490)^{22,24,25}. Of these, positions 452 and 490 were found to significantly impact VE (FWER $p \leq 0.05$). Latin America country-specific results for the same analyses are shown in Supplementary Figs. 9–12, where due to reduced sample size fewer AA positions were screened in for assessment of differential VE.

Supplementary Figs. 13–15 provide complete results, including by geographic region.

Vaccine efficacy decreases with vaccine-insert distance

VE significantly decreased with physicochemical-weighted Hamming distance (between the observed vs. vaccine insert sequence) for Spike, RBD, NTD, and S1 (Fig. 3, FWER $p < 0.001$) but not for S2 ($p = 0.78$). Against viruses with shortest Spike distances (average six residue mismatches), VE was 69% (95% CI: 60 to 76%), and against viruses with 25th, 50th, 75th, and 95th percentile Spike distances (average 8.1, 12.9, 17.8, 18.6 residue mismatches), VE was 64% (56, 71%), 52% (44, 58%), 34% (19, 46%), and 30% (13, 44%), respectively. The median distances of sequences for vaccine:placebo were 15.0:9.5 for Spike, 2.6:1.0 for RBD, 4.0:1.6 for NTD, 11.7:6.2 for S1, and 3.1:3.2 for S2. A sensitivity analysis of VE by physicochemical-weighted Hamming distance performed after removing the Lambda and Zeta outliers (defined by distance > 20) in Fig. 3 yielded similar results (Supplementary Fig. 16). Supplementary Tables 10 and 11 show inferences about differences in

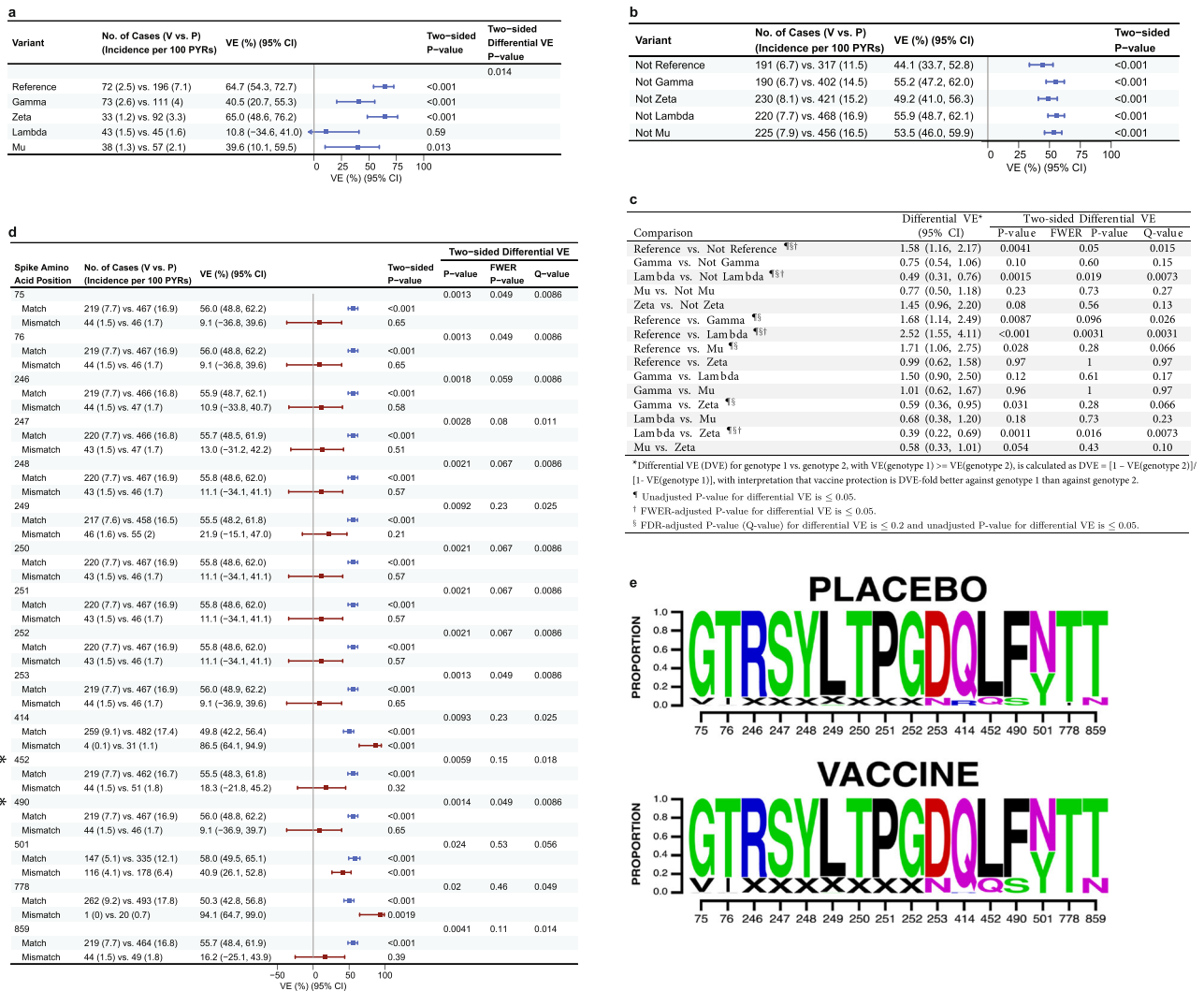
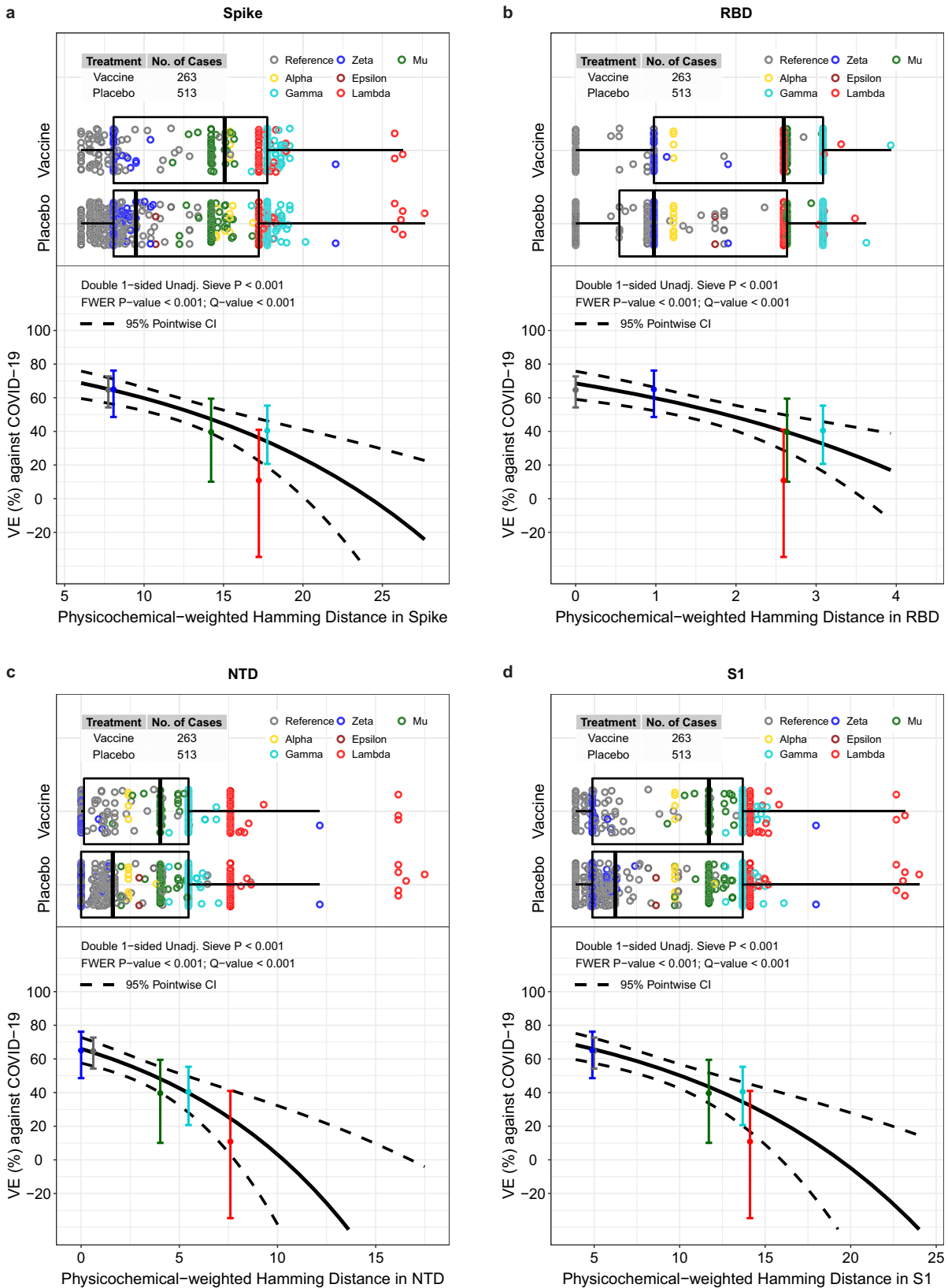


Fig. 2 | For the Latin America cohort, differential vaccine efficacy (VE) by lineage or by residue feature. a VE estimates against the primary COVID-19 endpoint caused by SARS-CoV-2 lineages (lineage “X”). **b** VE estimates against the primary COVID-19 endpoint caused by all other lineages combined (“Not X”). **c** Differential VE estimates against the primary COVID-19 endpoint across pairs of lineages or across a lineage (“X”) vs. all other lineages (“Not X”). **d** VE estimates against the primary COVID-19 endpoint caused by SARS-CoV-2 with a vaccine-matched or vaccine-mismatched residue at each of the 16 Spike amino acid (AA) residues with differential VE (q value <0.2 and unadjusted $p \leq 0.05$). Results for VE against matched residue genotypes are shown in blue and for mismatched residue genotypes in maroon. The two amino acid positions hypothesized to impact VE (452 and 490)^{22,24,25} are identified with an asterisk. In the forest plots in **a**, **b**, and **d**, the solid squares represent the VE point estimates, and the error bars display the 95% confidence intervals. In panels **a**, **b**, and **d**, the “Two-sided P value” is from testing the null hypothesis H_{A10} vs. H_{A12} using the test statistic U_{2j} (pp 17–18 of Heng et al.¹⁸). In panels **a**, **c**, and **d**, the “two-sided differential VE P value” is from testing the null hypothesis H_{B0} vs. H_{B2} using the test statistic T_2 (pp 18–19 of Heng et al.¹⁸). **e** Logo plots showing the distributions of AA residues and gaps/deletions (represented by “X”) by treatment arm at the 16 positions in (d). AA residues are color-coded by their chemical properties: green, polar; red, acidic; blue, basic; black, hydrophobic. CI confidence interval, DVE differential vaccine efficacy, FDR false discovery rate, FWER family-wise error rate, PYRs person-years.

mean distances of vaccine vs. placebo sequences. Supplementary Figs. 17–21 and Supplementary Table 12 provide complete results, including by geographic region, where Supplementary Table 12 shows that VE decreased with weighted Hamming distance for RBD, NTD, and S1 in the US (q value ≤ 0.20).

By lineage, ordered by placebo arm COVID-19 endpoint Spike distance to the vaccine strain, Reference viruses had 6.0–17.7 residue mismatches, Zeta 8.1–22.1 mismatches, Epsilon 10.7 mismatches, Mu 12.2–16.8 mismatches, Alpha 14.5–16.8 mismatches, Gamma 16.7–20.2 mismatches, and Lambda 17.2–27.7 mismatches. This ordering of lineages by protein distance matches the ordering of the VE estimates by lineage category, suggesting that overall Spike evolution is a reasonable metric capturing VE decline with a variant. The results are generally similarly ordered for the RBD, NTD, and S1 distances (Supplementary Fig. 22).

Neutralization-relevant RBD features were defined where mutations impact binding in deep mutational scanning (DMS) experiments²⁶ (see Supplementary Methods). Escape scores were defined for all antibodies and for each of 10 epitope-specific clusters of AA sites (see Methods), labeled DMS (all antibodies) and DMS1 through DMS10. Vaccine efficacy significantly decreased (q value ≤ 0.20) with each of the DMS, DMS2, DMS6, DMS7, and DMS8 escape scores (FWER $p \leq 0.05$) as well as for DMS1, DMS5, DMS9 (q value ≤ 0.20 and FWER >0.05) (Supplementary Table 13). Supplementary Tables 14 and 15 show the mean DMS escape scores in the vaccine arm and in the placebo arm of the disease-causing SARS-CoV-2 isolates, as well as the difference in mean DMS escape score between the two arms. To accommodate for missing sequences, in the analysis whose results are shown in Supplementary Table 14, doubly robust targeted minimum loss-based estimation was used; in



Supplementary Table 15, inverse probability weighting was used. Results were generally similar using the two different statistical methods, with greater mean DMS escape scores in the vaccine arm than the placebo arm for all of the clusters and the lower limit of the 95% CI usually greater than zero. The greatest difference in mean DMS escape score (vaccine – placebo) was seen for DMS5 [0.051 (0.0032, 0.098) in Supplementary

Table 14, 0.13 (0.073, 0.19) in Supplementary Table 15. Geographic region-specific results are also shown in Supplementary Tables 14 and 15.

Alternatively, we defined putative antibody footprint site sets (including whole Spike) based on structures of SARS-CoV-2 in complex with antibodies available from the Protein Data Bank (PDB). Each sequence was assigned an escape score based on a class of epitopes

Fig. 3 | For the Latin America cohort, vaccine efficacy (VE) against the primary COVID-19 endpoint by different physicochemical-weighted (PCW) Hamming distances. The Hamming distances were from the disease-causing SARS-CoV-2 isolate to that of the vaccine-insert sequence, for the following regions: **a** Spike, **b** the RBD domain, **c** the NTD domain, or **d** the S1 region. The top plot in each panel shows the distributions of distances by treatment arm, color-coded by lineage: Alpha, yellow; Epsilon, brown; Gamma, turquoise; Lambda, red; Mu, green; Reference, gray; Zeta, blue. The left and right edges of the box plots represent the 25th and 75th percentiles of PCW-Hamming distance in the designated region, and the vertical middle line represents the 50th percentile. The horizontal bars extend from the 25th (or 75th) percentile of PCW-Hamming distance to the minimum (or maximum) PCW-Hamming distance within the 25th (or 75th) percentile of Hamming distance minus (or plus) 1.5 times the interquartile range. The bottom plot in each panel shows the estimated VE by SARS-CoV-2 sequence distance. The dotted lines

are pointwise 95% confidence intervals. The dots are overall VE estimates for the given lineage placed at the lineage-specific median distance of placebo arm end-points, with vertical bars indicating their pointwise 95% confidence intervals. In each panel, the “Double I-sided unadjusted sieve p value” doubles the p value from a one-sided Wald test of the null hypothesis of constant VE vs. the alternative hypothesis of a decreasing VE with an increasing value of the feature on the x-axis (Juraska and Gilbert¹⁹, Section 5). Two Zeta sequences are visible outliers from other Zeta sequences; both sequences have two large deletions (9AA and 7AA in length) in the N-terminal domain. The plots reveal that Lambda has two sub-lineages, one ($n = 79$) with a range of distances 17.2–18.9 and a second ($n = 9$) with a range of distances 25.8–27.7, due to a 13-AA deletion between sites 64 and 76. CI confidence interval, FWER family-wise error rate, NTD N-terminal domain, RBD receptor-binding domain, Unadj. unadjusted.

(see Supplementary Methods). These features are referred to as PDB1 through PDB14, with the first 12 clusters in the RBD and PDB13 and PDB14 in the NTD. Vaccine efficacy significantly decreased (q value ≤ 0.20) with the escape scores for PDB4, PDB7, PDB8, and PDB13 (FWER $p \leq 0.05$) as well as for PDB1 and PDB3 (q value ≤ 0.20 and FWER > 0.05) (Supplementary Table 16). Supplementary Tables 17 and 18 show the mean PDB escape scores in the vaccine arm and in the placebo arm of the disease-causing SARS-CoV-2 isolates, as well as the difference in mean PDB escape score between the two arms. Analyses were performed the same as for Supplementary Tables 14 and 15. For each cluster, the mean PDB escape score was generally higher in the vaccine arm than in the placebo arm. The greatest difference in mean PDB escape score (vaccine – placebo) was seen for PDB13 [0.27 (0.043, 0.5) in Supplementary Table 17 and 0.47 (0.24, 0.7) in Supplementary Table 18. Geographic region-specific results are also shown in Supplementary Tables 17 and 18.

To interpret the DMS and PDB results, we focus on the epitope-specific features with FWER $p \leq 0.05$ that carry the greatest amount of independent information based on inter-correlation and hierarchical clustering analysis (Supplementary Note 1 and Supplementary Figs. 23, 24): DMS2, PDB7, PDB8, and PDB13. The sieve analysis results are similar across these four features, with estimated VE at 60–70% against viruses with an escape score of zero and decreasing to 0%–20% against viruses with a maximum escape score. PDB8 and PDB13 rank highest for discriminating VE with a slightly greater span of VE point estimates over the range of escape scores (spans 20–60%, 16–60%, 21–69%, and 1–57% for DMS2, PDB7, PDB8, and PDB13, respectively) (Fig. 4a–d).

Figure 5a lists the Spike AA residues in each epitope footprint, and the visualizations in Fig. 5b–e show the positions comprising the four antibody-epitope footprints on a Spike monomer structure. Supplementary Figs. 25–33, 34–40 provide complete results for DMS and PDB features, respectively. Another reason PDB8 was highlighted is its balanced contacts across the whole receptor-binding motif (RBM) whereas the other RBM-specific clusters (PDB1–PDB6) are more tightly grouped within a region of the RBM (Fig. 5f). The epitope of PDB8 is better to monitor broadly the mutations in the RBM than PDB1–6, which have more limited targets. Among the non-RBM focusing antibodies (PDB7, PDB9–PDB14), PDB7 (which includes variable sites) and PDB13 correspond to the most accessible sites on Spike in a closed prefusion trimer (Fig. 5g). These sites are relatively variable among SARS-CoV-2 sequences. The PDB9, PDB10, and PDB11 epitopes are cryptic and conserved, whereas the epitope for PDB12 appears accessible and conserved. Among NTD antibodies, the Fc domain may be more accessible for PDB13 than for PDB14 (Fig. 5h), which may be linked to improved Fc-dependent virus clearance.

Vaccine efficacy decreases with NTD-linked reduced neutralization

Seven dichotomous NTD features (Methods) were assessed for a sieve effect as for vaccine-match vs. vaccine-mismatch binary features. Six of

the 7 NTD features significantly impacted VE (q value ≤ 0.20): NTD4, NTD6, NTD1, NTD3, NTD5, and NTD7 (where the last four also had FWER $p \leq 0.05$) (Fig. 6a). Figure 6b–e show the spatial locations in the NTD of the four features that impacted VE (FWER $p \leq 0.05$).

Vaccine efficacy decreases by neutralization resistance score

All of the sieve analyses study how VE depends on Spike AA features except one: a neutralization sieve analysis that scores each virus's lineage by its experimentally measured sensitivity to neutralization by Ad26.COV2.S vaccinee sera^{27,28}. VE decreased with this variant-neutralization resistance score ($p = 0.011$) (Fig. 7). Under one model for the neutralization assay being a perfect correlate of protection, the estimates of VE for each of the five lineages would fall on the curve of VE by variant-neutralization resistance score. Lambda had evidence of deviating from the curve, with VE 55% (48, 62%) when estimated based on measured neutralization sensitivity compared to VE 11% (–35, 41%) when estimated based on direct analysis of Lambda ignoring neutralization data. In contrast, the weighted Hamming distance analyses yielded VE estimates at Lambda-variant distance values that are closer to the VE 11% figure.

Supplementary Fig. S41 provides complete results by geographic region.

Multivariable virus features as predictors of treatment arm

A variable importance measure (VIM) analysis by ensemble machine learning²⁹ was conducted to assess how well different groups of virus features in COVID-19 endpoint cases predicted treatment arm beyond that provided by baseline risk factors (whether the COVID-19 endpoint was from Colombia and enrollment periods). Virus features defined based on neutralization data were the top-performing predictors of the treatment arm, with the DMS RBD antibody-escape features having the highest estimated VIM (0.073, p value for a test of the null hypothesis of zero VIM = 0.043; Supplementary Fig. 42).

The second-most important classifying variables were the set of all nAb correlate of protection (CoP) hypothesis features, defined as DMS RBD and PDB antibody-escape features, NTD neutralization-relevant features, and the variant-neutralization sensitivity score (VIM = 0.051); PDB antibody-escape features (VIM = 0.049); and the variant-neutralization sensitivity score (VIM = 0.048) (Supplementary Fig. 42). The unbiased features specified to include all Spike AA variation ignoring neutralization hypotheses had the lowest estimated variable importance (VIM = 0.036 to 0.046 for the weighted Hamming distances and Spike AAs).

Generally stable vaccine efficacy against severe-critical COVID-19

Differential VE against severe-critical COVID-19 by lineage could only be assessed for Latin America, with VE of 83% (64, 92%) against Reference, 64% (26, 83%) against Gamma, 94% (–27, 100%) against Zeta, 62% (–31%, 89%) against Lambda, and 84% (42, 96%) against Mu

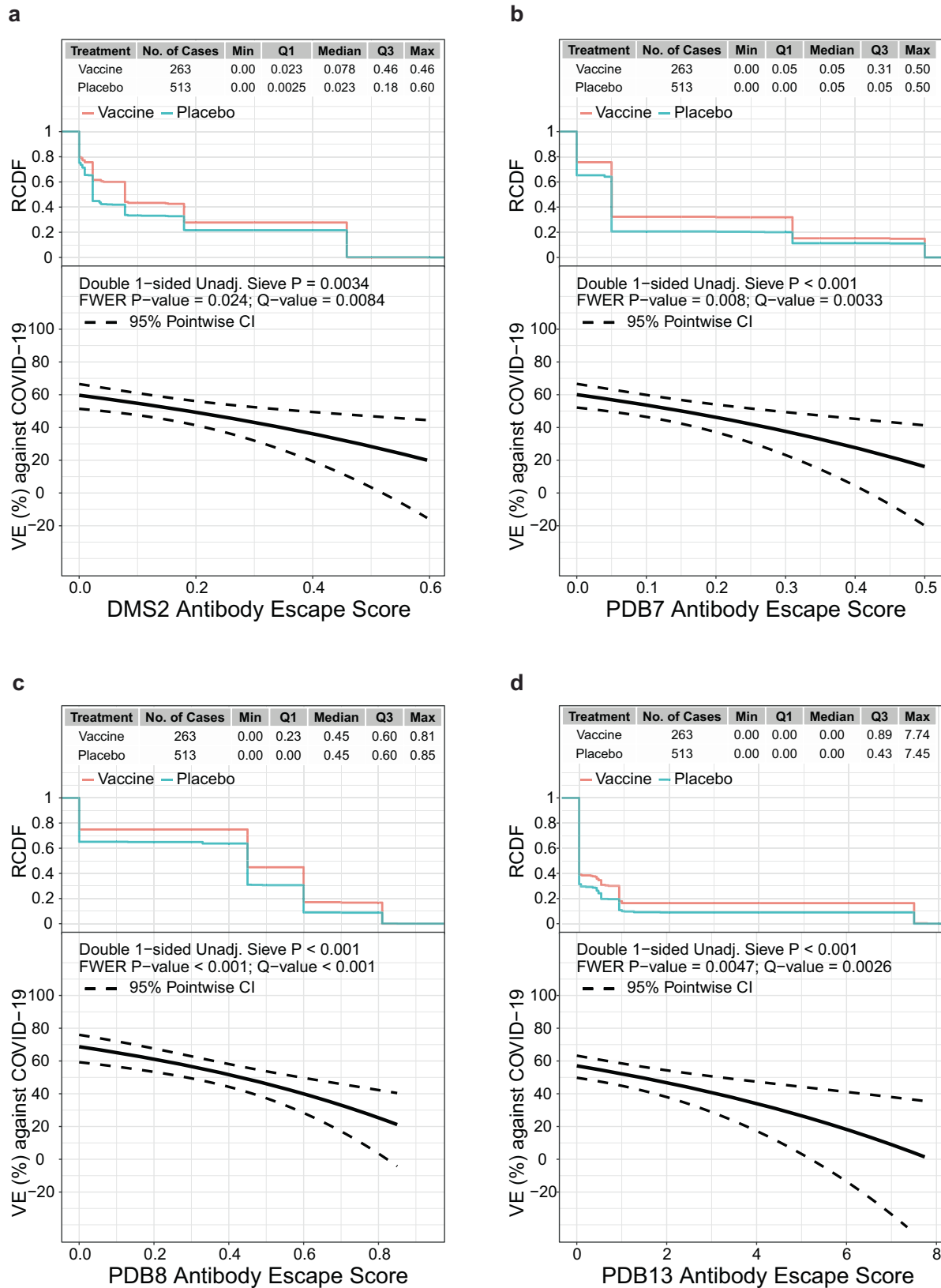


Fig. 4 | In the Latin America cohort, vaccine efficacy (VE) against the primary COVID-19 endpoint by the SARS-CoV-2 antibody-escape score. VE (point estimates as solid line, 95% confidence intervals as dashed lines) is shown by the antibody-escape scores for **a** DMS2, **b** PDB7, **c** PDB8, and **d** PDB13. In each panel, the “Double 1-sided unadjusted sieve *p* value” doubles the *p* value from a one-sided Wald test of the null hypothesis of constant VE vs. the alternative hypothesis of a

decreasing VE with an increasing value of the feature on the x-axis (Juraska and Gilbert¹⁹, Section 5). The plot at the top of each panel shows the reverse cumulative distribution function (RCDF) of the relevant antibody-binding escape score across SARS-CoV-2 isolates by treatment arm: Vaccine, pink; Placebo, turquoise. CI confidence interval, DMS deep mutational scanning, FWER family-wise error rate, PDB Protein Data Bank, Unadj. unadjusted.

a Antibody Escape Score-Based Putative Epitope Footprints (Spike AA Residues)

Antibody Escape Score	Spike AA Residues (Location)
A. DMS2	• 346, 348, 352, 417, 420, 455, 456, 468, 473, 475, 493, 504 (RBD)
B. PDB7	• 339, 340, 341, 344, 345, 346, 347, 348, 349, 351, 352, 354, 356, 399, 449, 450, 466, 468, 469, 470, 472, 481, 482, 483, 484, 490, 492 (RBD)
C. PDB8	• 351, 403, 417, 444, 445, 446, 447, 449, 450, 452, 453, 455, 456, 470, 472, 475, 483, 484, 485, 486, 487, 488, 489, 490, 492, 493, 494 (RBD)
D. PDB13	• 144, 145, 146, 147, 148, 150, 152, 246, 247, 248, 249, 250, 251, 252, 253, 254

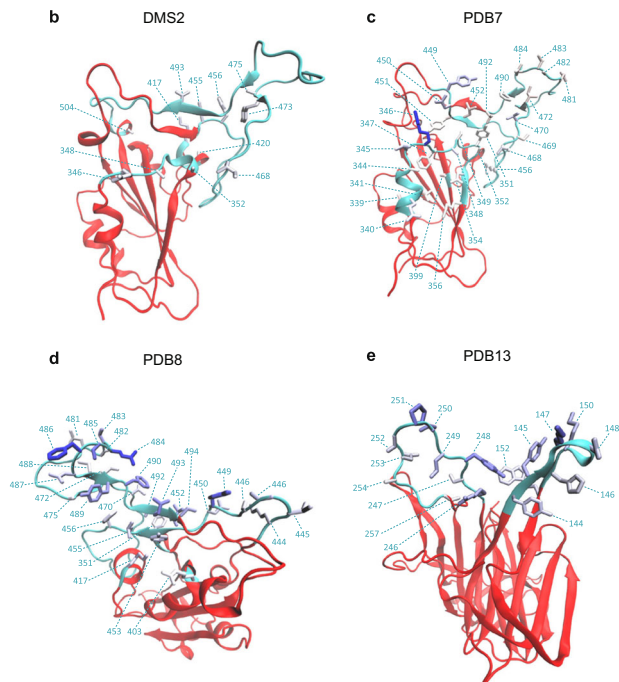
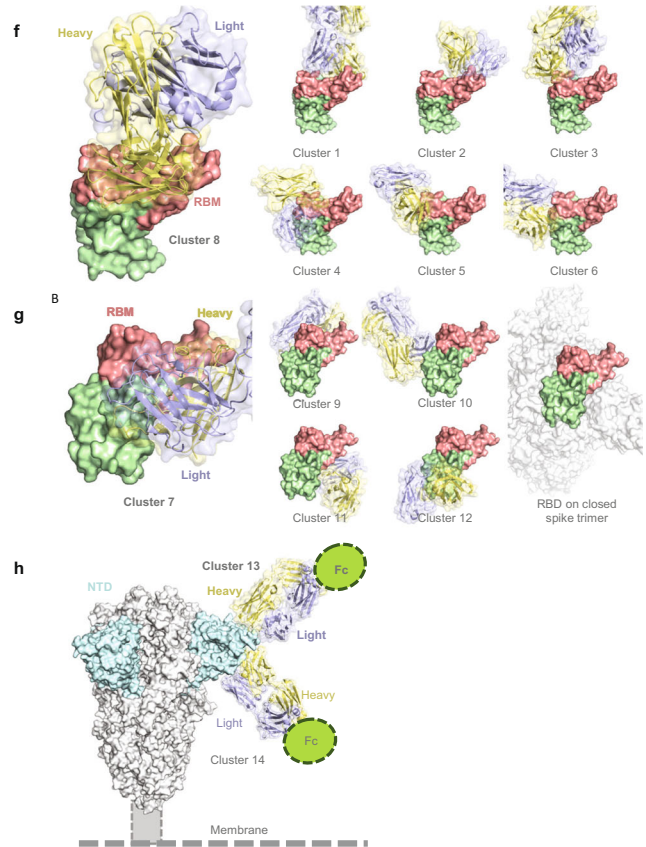


Fig. 5 | Structural visualizations of antibody footprints in Spike on antibody-escape score clusters and neutralization features where a sieve effect was seen. **a** Spike amino acid (AA) residues constituting each antibody-escape score-based putative epitope footprint. **b–e** For each set of residues constituting an antibody-epitope footprint for DMS2, PDB7, PDB8, and PDB13, the image shows the set of AA positions comprising the footprint on a Spike monomer NTD or RBD structure. Cyan ribbons highlight epitope footprint residues, while red ribbons make up the rest of RBD [**b** DMS2, **c** PDB7, and **d** PDB8]) or NTD (**e** PDB13). Residue numbers and



cyan dashed lines are used to label footprint residues. Each structure's orientation was chosen to best visualize all residues of a footprint. Residues are colored based on their cluster weights going from white to blue with increasing weight. **f–h** Epitope location of representative clusters 8, 7, and 13. Color coding denotes the following: Receptor-binding motif, red; heavy chain, yellow; light chain, lavender; N-terminal domain, turquoise; clusters, green. DMS deep mutational scanning, NTD N-terminal domain, PDB Protein Data Bank, RBD receptor-binding domain.

(Supplementary Table 19). There was no evidence of variation in VE across the lineages ($p = 0.50$) (Supplementary Tables 19, 20). The estimates of VE were similar/stable across AA positions with vaccine-matched vs. vaccine-mismatched residue, with all unadjusted p values for differential VE above 0.05 (Supplementary Fig. 43). For the key positions 452 and 490 found to show sieve effects for the primary COVID-19 endpoint, the results for the severe-critical COVID-19 endpoint were VE 79% (68, 87%) against 452-matched virus compared to VE 70% (3, 91%) against 452-mismatched virus ($p = 0.58$ for difference), and VE 80% (68, 87%) against 490-matched virus compared to VE 62% (–31, 89%) against 490-mismatched virus ($p = 0.34$ for differential VE). For the DMS antibody-escape score distances, the data support stable VE across the distances (Supplementary Table 21). Similarly, the data support stable VE across RBD and PDB Spike-antibody-escape scores (Supplementary Table 22). VE was stable by variant-neutralization resistance score, with VE = 84% (67%, 92%) for the most sensitive lineage (ancestral) and VE = 73% (50, 85%) for the least sensitive lineage (Mu) ($p = 0.33$, Supplementary Fig. 44).

There was a trend of VE against severe–critical COVID-19 decreasing with the weighted Hamming distance for the Spike, NTD, and S1 regions (q values = 0.20) (Supplementary Table 23 and Supplementary Figs. 45, 47, 48). The point estimates of VE suggested moderate declines of VE with distances. For example, the VE for Spike was 87% (71%, 94%) against viruses with a shortest distance of 6 and

66% (34%, 83%) against viruses with a long distance of 20 ($p = 0.12$). Supplementary Figs. 45–49 and Supplementary Table 24 provide complete information by geographic region. In addition, while VE was stable across levels of NTD1 through NTD4 ($p > 0.20$), it differed by levels of NTD5, NTD6, and NTD7, with VE of 61% (31, 78%) vs. 88% (76, 94%) for the two NTD5 genotypes ($q = 0.10$ for difference), VE of 60% (20, 80%) vs. 84% (72, 91%) for the two NTD6 genotypes ($q = 0.12$ for difference), and VE of 64% (32, 80%) vs. 85% (73, 92%) for the two NTD7 genotypes ($q = 0.12$ for difference) (Supplementary Table 24).

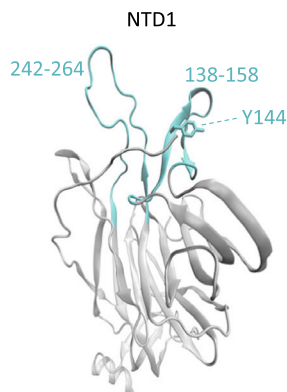
Exploratory analyses of Lambda's escape and driver mutations

A reviewer suggested additional analyses in pursuit of understanding how the level of VE against Lambda was lower than expected based on its level of neutralization resistance to vaccinee sera (Fig. 7) and the potential impact of driver vs. bystander mutations. Based on the reviewer's suggestions, we conducted two post-hoc exploratory analyses of Latin American data. First, we repeated the AA position site-scanning analysis, except removing all Lambda lineage viruses. Of the 16 AA positions with sieve effect evidence (Fig. 2d), three qualified for the sensitivity analysis based on sufficient residue variability (sites 414, 501, 778). The results of this sensitivity analysis also supported a sieve effect for all three positions (Supplementary Fig. 50), with VE estimates against vaccine-matched COVID-19 and against vaccine-mismatched COVID-19 in the Lambda-excluded analysis very similar to the all-

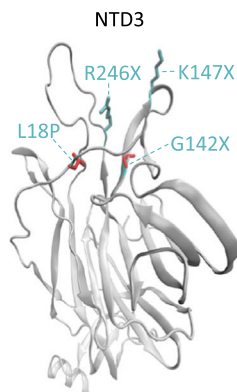
a

NTD Features	No. of Cases (V vs. P) (Incidence per 100 PYRs)	VE (%) (95% CI)	Two-sided P-value	Two-sided Differential VE		
				P-value	FWER P-value	Q-value
NTD1				0.0016	0.0065	0.0025
Mark Value = 1	93 (3.3) vs. 126 (4.6)	31.0 (10.3, 46.8)	0.0055			
Mark Value = 0	170 (6) vs. 387 (14)	58.7 (51.0, 65.3)	<0.001			
NTD3				0.0017	0.0065	0.0025
Mark Value = 1	46 (1.6) vs. 50 (1.8)	12.5 (-30.1, 41.2)	0.51			
Mark Value = 0	217 (7.6) vs. 463 (16.7)	56.1 (48.9, 62.2)	<0.001			
NTD4				0.097	0.19	0.12
Mark Value = 1	133 (4.7) vs. 223 (8.1)	45.3 (33.0, 55.4)	<0.001			
Mark Value = 0	130 (4.6) vs. 290 (10.5)	57.2 (47.9, 64.9)	<0.001			
NTD5				<0.001	0.0021	0.0012
Mark Value = 1	132 (4.6) vs. 188 (6.8)	35.2 (19.8, 47.6)	<0.001			
Mark Value = 0	131 (4.6) vs. 325 (11.7)	61.8 (53.7, 68.6)	<0.001			
NTD6				0.12	0.19	0.12
Mark Value = 1	77 (2.7) vs. 120 (4.3)	41.5 (22.7, 55.7)	<0.001			
Mark Value = 0	186 (6.5) vs. 393 (14.2)	55.2 (47.1, 62.0)	<0.001			
NTD7				<0.001	0.0013	0.0012
Mark Value = 1	117 (4.1) vs. 157 (5.7)	31.7 (14.2, 45.7)	0.0011			
Mark Value = 0	146 (5.1) vs. 356 (12.9)	61.0 (53.2, 67.5)	<0.001			

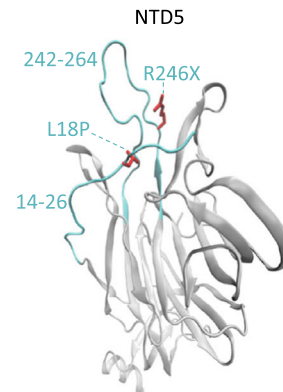
b



c



d



e

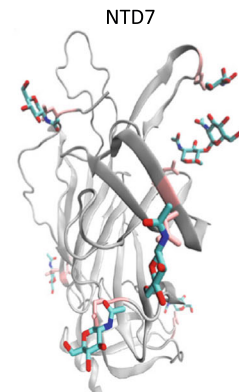


Fig. 6 | In the Latin America cohort, N-terminal domain (NTD) sequence feature sieve analysis and visualization of defining sites in the NTD features that impacted vaccine efficacy (VE). **a** VE estimates against the primary COVID-19 endpoint caused by SARS-CoV-2 with (vs. without) a NTD feature value, screened in as a specific hypothesis-driven neutralizing antibody (nAb) correlate of protection. VE estimates against SARS-CoV-2 harboring the NTD feature value are shown in blue; those against SARS-CoV-2 without the NTD feature value are shown in maroon. The solid squares represent the VE point estimates, and the error bars display the 95% confidence intervals. The “two-sided P value” is from testing the null hypothesis H_{A10} vs. H_{A12} using the test statistic U_{2j} (pp 17–18 of Heng et al.¹⁸) and the “two-sided differential VE P value” is from testing the null hypothesis H_{B0} vs. H_{B2} using the test statistic T_2 (pp 18–19 of Heng et al.¹⁸). **b–e** Visualization of the

constituent mutation sites that define the NTD feature, for each NTD features that significantly impacted VE with an FWER $p < 0.05$ (NTD1, NTD3, NTD5, and NTD7) (PDB 7L2C). **b** NTD1: Cyan colors the two insertions or deletions regions with one common residue for deletion (Y144) shown. **c** NTD3: The four specific mutations are labeled and colored cyan or red based on being mutations of interest or not respectively. **d** NTD5: Cyan colors the two substitution regions highlighting the two exclusions that are both found in this structure. **e** NTD7: Pink colors all asparagine residues that form N-linked glycans with the corresponding carbohydrate colored by atom (C: cyan, N: blue, O: red). Each structure shows the entire NTD region and colors all non-highlighted portions gray. CI confidence interval, FWER family-wise error rate, PYRs person-years.

lineage results shown in Fig. 2d. Therefore, the association of AAs at these positions with VE did not depend on the lineage being Lambda.

Secondly, as not accounting for ancestral relationships among sequences can potentially detect associations of AA changes with VE that are due to bystander mutations within distinct lineages, we conducted phylogenetically corrected GenSig analysis^{30–34} of the 37 screened-in sites to better understand vaccine associations with residues (Supplementary Tables 25–27). This method, based on

phylogenetic reconstruction from the nucleotide sequences (as both silent and non-silent mutations are accounted for in the imputed ancestral evolution of the lineages), scans all amino acid positions throughout the full Spike protein to assess whether ancestral changes to or away from each residue are associated with vaccination status. Of the 16 AA positions significantly associated with VE in Latin America (Fig. 2d), the Lambda variant mutation RSYLTTPGD246-253N, RSYLTTPGD with the N246 glycan in the NTD supersite epitope, was

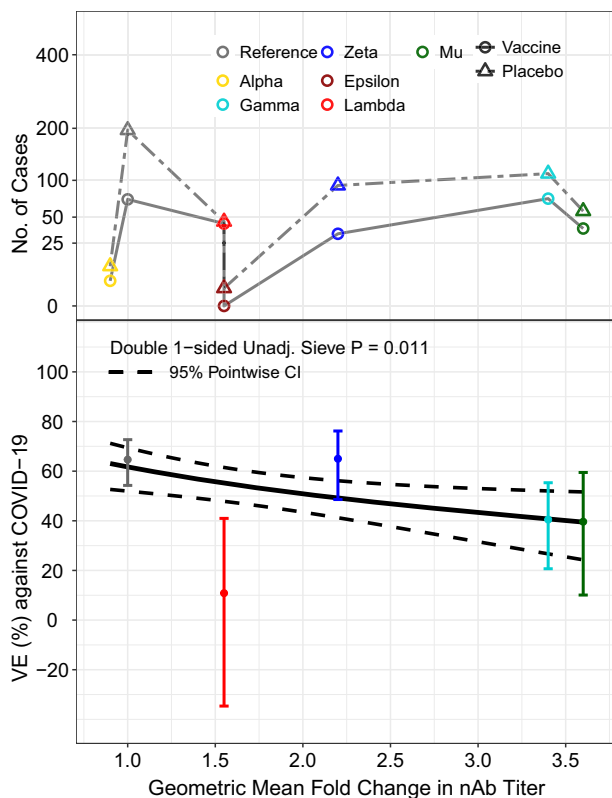


Fig. 7 | In the Latin America cohort, neutralization phenotype sieve analysis. Vaccine efficacy (VE) against the primary COVID-19 endpoint is shown by geometric fold change in neutralizing antibody titer against the disease-causing SARS-CoV-2 variant vs. against the D614G Reference strain. The top plot shows the numbers of cases by treatment arm (Vaccine: open circle; Placebo: open triangle) and color-coded by lineage: Alpha, yellow; Epsilon, brown; Gamma, turquoise; Lambda, red; Mu, green; Reference, gray; Zeta, blue. The bottom plot shows the estimated vaccine efficacy by geometric fold change in nAb titer against the disease-causing SARS-CoV-2 variant vs. against the D614G Reference strain. The dashed lines are pointwise 95% confidence intervals. The dots are VE point estimates against the given lineage, with the vertical bars showing 95% confidence intervals. The “double one-sided unadjusted sieve p value” doubles the p value from a one-sided Wald test of the null hypothesis of constant VE vs. the alternative hypothesis of a decreasing VE with an increasing value of the feature on the x-axis (Juraska and Gilbert¹⁹, Section 5). CI confidence interval, nAb neutralizing antibody, Unadj. unadjusted.

associated with decreased VE after phylogenetic correction (FDR-adjusted $p < 0.2$). Because the GenSig tool runs a test at each AA position, to make sure it was indeed the full deletion to glycan mutation that yielded significance after phylogenetic correction, we reran after excluding 6 sequences with ambiguous calls in the deletion, and forced the 246–253 motifs into a single mutation. The N246 glycan change was significant after phylogenetic and multiple testing correction (FDR-adjusted $p = 0.002$). While no other AA position variables were significantly associated with VE after phylogenetic and multiple testing correction, other mutations characteristic of the Lambda variant (G75V, T76I, L452Q, F490S) and/or associated with changes in RBD class I and II bnAb binding (L452R, E484K, F490S) or neutralization potency (L18F, T20N, K417T/N, L452R) were significantly enriched in the breakthrough sequences from the vaccinated group (FDR-adjusted $p < 0.10$).

Structure and immune evasion of the Lambda variant glycoprotein

To gain insight into the potential mechanisms underlying these observed Lambda variant sieve effects, we determined the structure of the Lambda-variant glycoprotein spike trimer bound to S309 (an RBD-

targeting nAb³⁵), S2L20 (an NTD-targeting non-nAb⁴³), and S2X303 (an NTD-supersite targeted nAb³⁶) using cryoEM (Fig. 8a). S383C/D985C mutations³⁷ were used to staple the RBDs closed, thereby enabling the use of C3 symmetry to aid structural determination.

Local classification and refinement of the RBD bound to S309 improved the local resolution of this region, permitting atomic model building in the cryoEM density, including the L452Q mutation and the main chain of F490S (Fig. 8b). Both mutations at positions L452 and F490 are associated with escape from site Ib RBD-targeted antibodies. ELISA experiments confirmed that Lambda escapes three of a panel of 12 RBD-targeting antibodies (Fig. 8d). Yeast-display experiments showed that the F490S mutation does not improve binding to ACE2, while the L452Q mutation hardly has an effect on ACE2 binding³⁸. Consistent with this study, binding experiments confirmed the Lambda RBD binds with approximately equivalent affinity to ACE2 as the index strain RBD (Fig. 8e).

Local classification and refinement of the NTD bound to S2L20 and S2X303 improved local resolution of this region, permitting model building, including the R246N mutation and the 247–253 deletion (commonly referred to as the “RSYLTPGD246-253N” mutation) (Fig. 8c). This deletion introduces a new glycan sequon and the linked glycan could be visualized and built in the cryoEM map. These mutations are in the NTD antigenic supersite loop and are therefore predicted to permit escape from NTD-targeted nAbs. ELISA experiments confirmed that Lambda escapes 10 of a panel of 11 NTD-targeted nAbs (Fig. 8d).

Discussion

Sieve analysis compares genotype-specific or immunophenotype-specific COVID-19 incidence between randomized study groups, therefore directly assessing the causal effects of vaccination and providing inferences for how vaccine efficacy depends on SARS-CoV-2 features. In addition to the strength of a randomized, double-blinded, placebo-controlled phase 3 trial, the present sieve analysis of ENSEMBLE had ample statistical precision due to the large number of SARS-CoV-2 Spike sequences (measured from more than 1,200 participants) and the broad proteomic variability of the SARS-CoV-2 Spike sequences causing these endpoints. Consequently, the sieve analysis could provide many insights into how the efficacy of the Ad26.COVS.2 vaccine, evaluated in baseline SARS-CoV-2 negative individuals, depended on virus features.

In the Latin American cohort, VE against the moderate to severe-critical COVID-19 primary endpoint significantly declined with Spike sequence distance as measured in myriad ways, including lineage, weighted Hamming distances calculated for Spike, RBD, NTD, and SI, scores reflecting degree of escape from epitope-specific antibodies computed using deep mutational scanning or based on crystal structures in the PDB, and NTD features previously shown to impact neutralization. Estimates of VE by lineage were consistently ordered by the distances of the different lineages to the vaccine strain. VE declined similarly with Spike, RBD, NTD, and SI distances (VE about 70% against viruses closest to the vaccine and 20% against viruses beyond the 90–95th percentile of distances) but did not depend on S2 distances. This may be explained by S2’s relative conservation when compared to SI. As such, almost all variant-characteristic mutations are not in S2, and none of the prescribed antibody-epitope footprint clusters included S2 positions (only rare epitopes in PDB mapped to S2), reflecting S2’s stalk location and relative lack of exposure to the immune system.

VE significantly declined with 14 of the 20 evaluable antibody-epitope escape scores. Six antibody-epitope clusters had no evidence of impacting VE: DMS3, PDB2, PDB5, PDB6, PDB9, and PDB14. Of the 14 clusters with a sieve effect, nine include at least one site that harbors a characteristic mutation of Lambda, whereas three include site 417 which is a characteristic mutation of Mu and Gamma, one includes site 501 that harbors a characteristic mutation of Gamma, Alpha, and Mu,

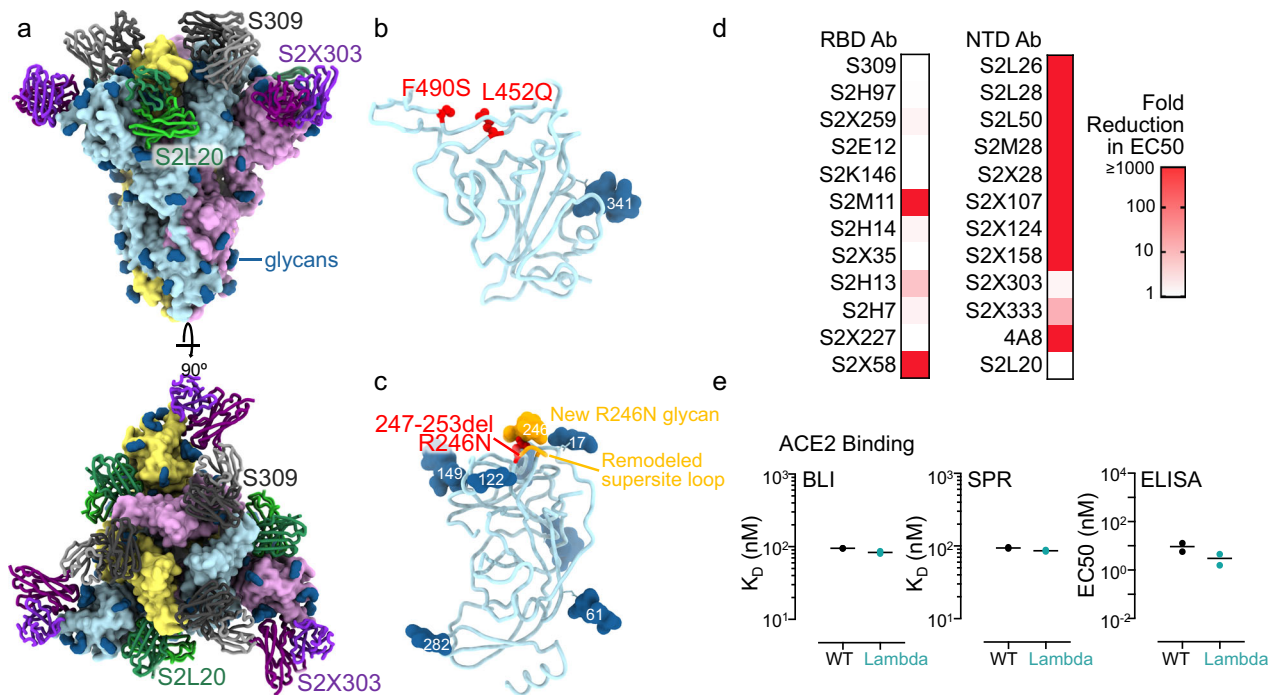


Fig. 8 | CryoEM structure of the SARS-CoV-2 Lambda Spike ectodomain trimers and analysis of monoclonal antibody (mAb) and ACE2 binding. **a** Structure of the Lambda Spike trimer (surface rendering) bound to the S2L20, S2X303, and S309 Fabs (ribbons). SARS-CoV-2 Spike protomers are colored pink, cyan, and gold, whereas the S2L20 Fab heavy and light chains are colored dark and light green, respectively. The S2X303 Fab heavy and light chains are colored dark and light purple, respectively. The S309 Fab heavy and light chains are colored dark and light orange, respectively. Only the Fab variable domains are resolved and therefore modeled in the map. N-linked glycans are rendered as dark blue spheres. **b** Zoomed-in view of the S309-bound Lambda RBD with L452Q and F490S shown as red spheres. **c** Zoomed-in view of the S2L20- and S2X303-bound Lambda NTD with the R246N mutation shown as red spheres; the remodeled loop caused by the 247–253 deletion as well as the new R246N glycan are shown in orange. **d** Binding of a panel of 12 neutralizing RBD-specific mAbs, 11 neutralizing (antigenic site i) NTD-specific mAbs, and one non-neutralizing (S2L20, antigenic site iv) NTD-specific

mAbs to recombinant SARS-CoV-2 Spike variants analyzed by ELISA displayed as a heat map (relative to SARS-CoV-2 index strain Spike binding). **e** (Left) Biolayer interferometry (BLI) binding analysis of the human ACE2 ectodomain (residues 1–615) to immobilized biotinylated SARS-CoV-2 index strain and Lambda RBDs. Data from one biological replicate is shown with 1–2 technical replicates each. (Center) Surface plasmon resonance (SPR) binding affinity analysis of the human ACE2 ectodomain (residues 1–615) for immobilized biotinylated SARS-CoV-2 index strain and Lambda RBDs. Data from two biological replicates are shown with two technical replicates each. (Right) ELISA binding analysis of the SARS-CoV-2 index strain and Lambda RBDs to immobilized human ACE2 ectodomain (residues 1–615) shown as 50% effective concentrations (EC50). Data from two biological replicates are shown with two technical replicates each. CryoEM cryo-electron microscopy, NTD N-terminal domain, RBD receptor-binding domain. Source data are provided as a Source Data file.

and one includes both sites 417 and 501. Thus the nine sieve-effect clusters appear to be driven by the differential VE by Lambda vs. not-Lambda, whereas the other five appear to be driven by mutations at the important sieve-effect sites 417 and 501 that impact neutralization. Of the six non-sieve-effect clusters, only one (PDB14) included a site harboring a characteristic mutation of Lambda, site 75, which was a sieve-effect site with FWER $p \leq 0.05$. The potential for sieve effects in different epitope sets depends on many factors, including the level of accessibility to nAbs, conservation, and the narrowness of the footprints on the tridimensional structure they target.

Given that most of the sieve effects found in this study are linked with Lambda, it is not surprising that prior work suggests that Lambda's mutations enable it to function as an immune escape variant. Kimura et al.³⁹, Wang et al.⁴⁰, and Acevedo et al.²⁴ showed that Lambda exhibited reduced neutralization sensitivity to antibodies induced by both vaccination and prior infection, with evidence suggesting that these effects were caused by the L452Q, F490S, and RSYLTPGD246–253N mutations. Kimura et al. and Acevedo et al. also considered other variants, including Gamma, and found that Lambda exhibited the least neutralization sensitivity among all those studied, and Kimura et al. found that Lambda was more infectious than the other variants due to its T761 and L452Q mutations. Additionally, ref. 41 directly compared Lambda with Delta and demonstrated that Lambda was more successful at evading vaccine-induced humoral

immunity, while showing similar binding affinity to ACE2 as the ancestral strain.

nAb assays have performed well at predicting vaccine efficacy against COVID-19 and severe-critical COVID-19 across SARS-CoV-2 lineages^{15,16,42}. Importantly, one of the sieve analyses in the present work scored viruses by their lineage's directly measured resistance to neutralization by sera from ENSEMBLE Ad26.COVID.S vaccine recipients, providing a way to study a neutralization correlate of protection (CoP) in a complementary way to individual-level (e.g., refs. 43–46) and population-level immune correlates analyses (e.g., ref. 47). VE significantly declined against lineages with greater neutralization resistance scores, providing validation of pseudovirus neutralization titer as a CoP. However, while refs. 24,39–41 demonstrated that Lambda was more resistant to nAbs than other variants, Lambda was less resistant to neutralization by Ad26.COVID.S vaccinee sera than Mu, Gamma, and Zeta, such that the ordering of variants by this neutralization scoring was discordant with the level of vaccine efficacy against the variants, indicating that vaccine efficacy against Lambda was lower than predicted from a serum nAb-CoP model. While, to our knowledge, no studies have directly compared the immune response susceptibility of Lambda vs. Zeta and of Lambda vs. Mu, two of the studies above^{24,39} demonstrated that Lambda showed a lower neutralization sensitivity than Gamma, suggesting that neutralization readouts from vaccinee sera cannot be used as the sole

basis of extrapolating predictions of VE, where other immune functions (e.g., cellular, Fc effector) could be important to include in such predictions. It is interesting to note that ref. 48 showed that Spike-specific CD4 + T-cell and CD8 + T-cell responses in Ad26.COV2.S vaccine recipients ($N=28$) were generally maintained across the Alpha, Beta, Gamma, Delta, B.1.1.519, Kappa, Lambda, and R.1 variants, although at the individual-donor level, one Ad26.COV2.S vaccine recipient showed reduced Lambda-Spike CD4 + T-cell responses and one showed reduced Lambda-Spike CD8 + T-cell responses (both vs. Ancestral-Spike responses). Additionally, the association of weighted Hamming distances with vaccine efficacy (Fig. 3) were concordant with vaccine efficacy by variant, especially for NTD, with Lambda being most distant from the vaccine strain (with notable outlying viruses with 25 residue mismatches to the vaccine strain) and against which vaccine efficacy was lowest. This suggests that amino acid sequence distances may have advantages as a biomarker for reliably predicting vaccine efficacy, a result also supported by ref. 7. A caveat of the neutralization sieve analysis is that the lineage scores were estimated from vaccinee sera from only eight ENSEMBLE participants, although the scores were supported by additional data from 17 Ad26.COV2.S vaccine recipients in the COV2001 phase 1/2a study²⁷.

The relative prevalence of SARS-CoV-2 lineages changed over time (Fig. 1a and Fig. 1 of ref. 6) where in Latin America the median (range) number of days from enrollment until the COVID-19 endpoint among placebo recipients was 48 (15, 197) for Reference.45, (15, 141) for Zeta, 114 (42, 220) for Gamma, 126 (57, 204) for Lambda, and 170 (109, 219) for Mu. If newer variants tended to expose participants later in follow-up than older variants, it could cause spurious genotypic sieve effects that are instead due to waning vaccine efficacy. This potential bias was mitigated by controlling for the calendar time of enrollment in the sieve analyses.

Given that the sieve effects identified in this analysis have strong linkages with viral variants (especially Lambda), a natural consideration is to identify which of these features are drivers of vaccine escape versus lineage-linked “bystander” features that are merely along for the ride. Among the four identified sieve sites in the RBD (414, 452, 490, 501), the deep mutational scanning work by ref. 9 identified sites 452 and 490 with significant antibody-escape in some variants, and that site 490 is in a Class Ib epitope targeted by antibodies such as LY-CoV555. Of the ten identified sieve sites in the NTD (75, 76, 246, 247, 248, 249, 250, 251, 252, and 253), eight of them fall in the NTD supersite identified by ref. 13, by way of Lambda’s RSYLTPGD246-253N mutation. The end result of this mutation is an N-linked glycosylation motif starting at position 246, which is a novel motif only found in Lambda. The glycan at position 246 and the deletion remodel the supersite and promote NTD-targeted nAb escape, as confirmed by ELISA, showing that binding of 10 out of 11 tested NTD-targeted mAbs was abrogated. As R246A or R246Q point mutations in the Lambda spike abolish this glycan sequon and do not affect neutralization of BNT162b2 vaccine recipient sera³⁹, the 247–253 deletion rather than the new glycan may be solely responsible for the association of the RSYLTPGD246-253N mutation with decreased VE. These findings provide a potential mechanism by which the Lambda variant deletion RSYLTPGD246-253N in the NTD supersite epitope was associated with decreased VE after phylogenetic correction. The other two NTD sieve sites (75 and 76) are the trailing residues in an N-linked glycosylation motif beginning at position N74; similarly, as a glycan at this site may promote binding, mutations in these downstream positions disrupt the motif and may facilitate immune escape.

A phylogenetically corrected analysis of a subset of 1159 codon-aligned nucleotide Spike sequences confirmed a significantly lower VE against the Lambda variant. This analysis also found additional sites implicated in higher bnAb neutralization resistance to be enriched in the vaccine group breakthrough sequences. While these additional associations were not significant after the phylogenetic

correction, one needs to take into account the low diversity and recent evolutionary history of the Spike protein at the time of this analysis compared to HIV-1, for which the GenSig tool was originally designed, thus underpowering the overall analysis. Interestingly, many of these sites are enriched in the Lambda variant, against which VE was found to be significantly lower after phylogenetic correction. At the time it emerged, many studies indeed predicted higher resistance to RBD bnAbs, infectivity, and higher likelihood to escape vaccines in the Lambda variant compared to contemporary variants^{25,41}.

Another important consideration in the relevance of these results is how they are reflected in the current epidemic. The Ad26.COV2.S vaccine sieve effects observed here, based on data collected prior to July 10, 2021, revealed broader vaccine adaptation features as several sieve signature sites showed mutations in subsequent variant waves, including many that are still circulating today. At the time of this writing (October 2023), mutations at sites 252, 484, 490, and 501 are dominant in currently circulating Omicron sub-lineages and recombinants, including BA.2.86, EG.5.1, FL.1.5.1, GL.1, and HK.3. (Global proportion between June 27, 2023 and September 24, 2023: G252V = 75.2%; E484A = 96.5%; F490S = 93.2%; N501Y = 96.9%⁴⁹.) Of note is the sieve signature site F490S. While rare until the end of 2022, this mutation became dominant in early 2023 with the rapid global spread of the XBB.1.5 recombinant lineage⁵⁰.

Other mutations correlating with vaccine efficacy in this analysis have shown to be characteristic of other transient lineages, although they did not persist as long-term circulating features: e.g., G75V with DZ.2 and XBB.1.14; T76I with BA.5.5; S247N with BF.31.1; Y248D with BQ.1.145; T250I with AY.33; and P251H with BQ.1. L452R exhibited a limited show of prominence up through the start of 2022 with the Delta variant AY.4, and again in the latter half of 2022 with the Omicron sub-lineage BA.5.2.1, but has since faded from circulation. Even Q414R and T859N, the two VE-associated mutations that were not characteristic of any of the WHO-labeled variants observed in this study, emerged briefly as characteristic mutations in future minor lineages (AY.29.2 and B.1.637, respectively).

The fact that sieve analysis predicted currently relevant mutations could be expected, since SARS-CoV-2 has shown remarkable patterns of convergent evolution since the initial appearance of variants, with numerous recurrent mutations, especially in the RBD, shared across lineages over time⁵¹. Conversely, some of the sites found to be associated with Ad26.COV2.S efficacy in this analysis (R246 and L249) were not strongly characteristic of any future lineages of note. The R246N mutation in Lambda, as mentioned above, results in a novel N-linked glycosylation motif, and its apparent absence in future lineages seemingly validates Kimura et al.’s finding that a glycan at this position does not impact the virus’s fitness. Site 249, while also a part of the NTD supersite, seemingly does not tolerate substitutions very well, as its wild-type residue (L) is currently present in over 99% of Spike sequences, with the next-most prevalent mutation being a deletion. As such, its identification as a sieve site in this analysis may be due to the happenstance of being part of Lambda’s RSYLTPGD246-253N mutation.

A strength of this study was that it was conducted in three separate geographic regions with different circulating lineages, which contribute insights based on these lineages and their characteristic signature mutations, and different distributions of genetic distances of circulating sequences to the vaccine strain. The analyses of Latin American study sites provided the greatest insights given that 63% of primary COVID-19 endpoints with sequence data were in Latin America, where the circulating SARS-CoV-2 sequences were the most diversified. All features showing sieve effects in the US also showed sieve effects in Latin America, constituting independent replication of results. The result of no sieve effects in South African study sites can likely be explained by the vast majority of circulating sequences being

Beta or Delta variants with limited dynamic range of genetic distances within each variant and a lack of Reference viruses that are close to the vaccine strain.

Another strength of this study was that VE against severe-critical COVID-19 could be assessed. The results support that VE against this endpoint also declines with Spike sequence distance as measured in multiple ways, yet with VE starting higher against viruses closest to the vaccine strain and diminishing less rapidly with increasing degrees of sequence mismatch. Thus, the results generally endorse the hypothesis that a single dose of Ad26.COVS vaccine will durably protect against severe outcomes. However, it is unlikely to protect against reinfection, as might be expected of any member of the coronavirus family⁵². It is difficult to deconvolute the effects of antigenic change on protection from severe outcomes and protection from reinfection from the natural decay of infection-blocking immunity as compared to the near permanence of immunity against severe outcomes.

Overall, the finding that protection against severe-critical COVID-19 is more invariant to sequence changes than against less-symptomatic COVID-19 may have clinical implications for planning updates of vaccines with new variants. The severe-critical classification covers a broad spectrum of clinical phenotypes ranging from individuals with only repeated low partial pressure of oxygen to severe pneumonia requiring respiratory support. Protection against hospitalization with severe consequences is clinically most important but sieve analysis specific to this outcome could not be performed given the small number of cases. Yet, ENSEMBLE and post-approval trials have shown high Ad26.COVS efficacy against this outcome, especially in South Africa after a 6-month boost, suggesting that neutralization resistance and sequence variation may be playing a less-dominant role in vaccine-induced protection against the most serious disease, perhaps due to CD8 + T cells⁵³.

Methods

Trial design, study cohort, and COVID-19 endpoints

Trial enrollment began on September 21, 2020. Participants were not compensated for their participation. The end of the double-blind period varied by country; the data cutoff for this analysis was July 9, 2021. The main endpoint for sieve analysis is the same COVID-19 primary endpoint (moderate to severe-critical) as in the primary analyses^{6,17}, restricting to endpoints starting 14 days post-vaccination. Moderate COVID-19 was defined by a positive RT-PCR test for SARS-CoV-2 as well as two or more of the following symptoms (new or worsening): fever or chills, cough, heart rate ≥ 90 beats/minute, muscle or body pain, headache, new loss of taste or smell, sore throat, red or bruised-looking feet or toes, nausea, vomiting, or diarrhea; or one or more of the following signs or symptoms: shortness of breath, respiratory rate > 20 breaths/minute, clinical or radiologic evidence of pneumonia, deep vein thrombosis, or abnormal oxygen saturation (but above 93%)^{6,17}.

Severe-critical COVID-19 was defined by a positive RT-PCR test for SARS-CoV-2 with one of the following features: respiratory failure; evidence of shock (systolic blood pressure < 90 mm Hg, diastolic blood pressure < 60 mm Hg, or requiring vasopressors); respiratory rate > 30 breaths/min; heart rate ≥ 125 beats/min; oxygen saturation of 93% or less (ambient air at sea level), or a ratio of the partial pressure of oxygen to the fraction of inspired oxygen < 300 mm

Hg; intensive care unit admission; significant acute renal, hepatic, or neurologic dysfunction, or death^{6,17}.

Sieve analyses were also conducted for severe-critical COVID-19, again using the same definition as used in the primary papers^{6,17}. Analyses were conducted in the per-protocol baseline seronegative cohort¹⁷. See Section 1 of the SAP (provided in ref. 20 and at the end of the Supplementary Information) and Supplementary Methods for further details.

SARS-CoV-2 sequencing and sequence data

The Virology Laboratory at the University of Washington, Department of Laboratory Medicine and Pathology (“UW Virology”) conducted next-generation sequencing of SARS-CoV-2 Spike sequences with the Swift Biosciences SNAP workflow version 2.0 on Illumina platforms^{17,54}. Only Spike gene sequence information was obtained, and the assignment of WHO-labeled variants was based on profiles of predefined and characteristic amino acid substitutions in the Spike protein relative to the reference sequence [GenBank accession number NC_045512 (<https://www.ncbi.nlm.nih.gov/nuccore/1798174254>)]. For SARS-CoV-2 sequences, Nextclade (<https://clades.nextstrain.org/>)⁵⁵ and Pangolin (<https://cov-lineages.org/resources/pangolin.html>)⁵⁶ were used for lineage assignments. Sequenced samples were obtained as close as possible to the start of the symptoms from individuals acquiring a COVID-19 primary endpoint, where typically but not exclusively, samples with SARS-CoV-2 viral load above 1,000 copies/mL were shipped for sequencing.

When the sequencing was successful, the consensus sequence from the set of sequencing reads from a single run on a single clinical sample was used as the sequence in the analysis. For the 109 participants with sequences from more than one sample (97 with sequences from two samples and 12 with sequences from three samples), the consensus sequence from the chronologically earliest timepoint was used in the analysis.

Sequences were selected for analysis if they were obtained within 36 days following the first RNA-positive timepoint associated with the first moderate to severe-critical COVID-19 primary endpoint. See Supplementary Methods for further details.

In our analyses, we compared the observed sequences with the sequence to the insert of the Ad26.COVS vaccine. The vaccine insert strain/sequence is from the index strain (prior to the D614G mutation) with two stabilizing mutations at K986P and V987P, and two additional mutations (R682S and R685G) in the furin cleavage site. All site positions mentioned are relative to those in the index sequence.

nAb titers

nAb titers were measured to a panel of Spike antigens representing the Reference strain B.1.D614G and several variants^{27,28}. The nAb assay used a commercial cell line consisting of HEK293T target cells stably expressing the human ACE2 and human TMPRSS2 genes [CoronaAssay-293T(hACE2-hTMPRSS2), procured from VectorBuilder; Cat. CL0015]. R (version 3.4.3) was used to calculate SARS-CoV-2 neutralizing titers using a four-parameter curve fit as the sample dilution at which a 50% reduction (IC50) of luciferase readout was observed compared with luciferase readout in the absence of serum (High Control). Relative light units were measured using an EnSight Multimode Plate Reader (Perkin Elmer) running Kaleido software (version Kaleido 3.0.3067.117x). Each variant was assigned a score defined as the log₁₀-transformed ratio of the geometric mean titer of vaccinee sera against the variant and the geometric mean titer of vaccinee sera against the Reference strain.

Sieve analysis

This analysis was prespecified and documented in the SAP. The sequences and clinical data were pre-processed into an analysis dataset as specified by the SAP, using R⁵⁷ (version 4.3.1) with the seqinr⁵⁸ package (version 4.2–30). The sieve analyses were conducted for each of the four geographic regions: Latin America, South Africa, the United States, and the three geographic regions pooled (hereafter, “geographic-region analyses”). For each geographic-region analysis, lineages with at least 20 COVID-19 endpoints were included. For amino acid (AA) position scanning sieve analysis that considered residue match-vs.-mismatch to the vaccine-strain residue, positions with at least 20 COVID-19 endpoints with a residue match and at least 20 COVID-19 endpoints

with a residue (or gap/insert/deletion) mismatching the vaccine-strain residue, were included. Similar AA position scanning sieve analyses were done that focused on specific residues at given positions, where residues at positions with at least 20 COVID-19 endpoints with the residue and at least 20 COVID-19 endpoints without the residue, were included. The same screening rule was used for the Latin America country-specific analyses. For the severe-critical COVID-19 endpoint, ten severe-critical COVID-19 endpoints were used to down-select the lineages and AA position features for each geographic-region analysis. Further details are given in Section 1.6 of the SAP.

While the primary analyses^{6,17} both counted endpoints starting 14 or 28 days post-vaccination, all of the sieve analyses restrict to starting 14 days post-vaccination, given that similar results are expected and more COVID-19 endpoints could be included in the analysis.

Specification of Spike AA sequence features for sieve analysis

We performed unsupervised learning of the treatment-blinded trial sequence data to fully specify and down-select the set of AA sequence features that were studied for sieve effects. All statistical inferences were prespecified in the SAP before treatment unblinding. Then, inferential statistical analysis (supervised learning that produces VE estimates, differential VE estimates, confidence intervals, and p values) was conducted in an automated/press-button fashion, with the inferences valid based on the pre-specification of inferences and the reproducibility of the computer code.

Features were classified into two types: (1) all Spike AA sequence features with sufficient variability to study for potential sieve effects (“All” features) and (2) the subset of All features

that are directly connected to a hypothesis that nAbs are a correlate of protection, and were selected based on knowledge/data or hypotheses that different levels of the feature affect neutralization (nAb-CoP-hypothesis features). See Section 1.3 of the SAP for further details.

Handling of missing sequences

For primary endpoint COVID-19 cases, either all Spike sequence features are observed, or no Spike sequence features are observed. Thus, the structure of the missing data pattern for primary endpoint cases is simple, with complete sequence data or no sequence data. See Section 2.4 of the SAP for details on how each specific sieve analysis method handles the missing sequence data.

Quantification of viral diversity

Spike sequence diversity within each geographical location (Latin America, South Africa, United States of America) was quantified using Rao’s Q , which measures the average phylogenetic distance between any two Spike AA sequences randomly chosen from any given region⁵⁹.

Structural modeling

All protein structures were generated using VMD⁶⁰ (version 1.9.4). PDB files were obtained from the RSCB Protein Data Bank (PDB) and are specified as described in Supplementary Methods. Using Python (version 3.9), NumPy⁶¹ (version 1.20.3) was used for data processing and MDAnalysis^{62,63} (version 2.0.0) was used for processing, editing, and generating PDB structure files. All protein backbones were drawn in the NewCartoon style to keep focus on the residues and regions of interest. In order to apply a weighted color to the residues, a Python script (publicly available at ref. 64) was used to translate cluster weights into a VMD readable scale metric. This metric, the beta factor, can be used as a dummy field in PDB files and applies a red-to-blue gradient at values of 0 and 200, respectively. Cluster weights were set to a baseline of 100 before adding the cluster weight scaled by a factor of 20 in order to highlight cluster variation.

AA sequence sieve analysis methods: prospective VE sieve analysis

For sieve analyses that answer the questions of whether and how VE depends on AA sequence features of exposing SARS-CoV-2 viruses [prospective VE sieve analysis⁶⁵], the Spike AA sequence feature-specific VE estimands for measuring sieve effects are defined in the SAP. The main estimand used is genotype-specific hazard-ratio VE, which for a given genotype is defined as 100% times one minus the genotype-specific hazard ratio (vaccine/placebo) of COVID-19 over the follow-up period 14 days post-vaccine or placebo administration until unblinding. An AA sequence sieve effect is defined as statistically significant evidence for differential VE across multiple levels of a given AA sequence feature. Details on the sieve analysis methods are given in Section 2 of the SAP.

For these hazard ratio-based prospective VE sieve analyses, the following software was used: R 4.2.3⁵⁷ and Rstudio⁶⁶ 2023.03.0 + 386. For hazard ratio-based sieve model fitting, the R packages sievePH^{19,67} (version 1.0.4) and cmprskPH⁶⁸ were used. Information on additional R packages used is provided in the Supplementary Methods.

Multiple hypothesis testing adjustment for AA sequence sieve analysis

For the AA sequence sieve analysis, the following plan was implemented for multiple testing adjustment, separately for each geographic region-specific analysis:

Family-wise error rate (FWER) adjusted p values are Holm–Bonferroni and FDR-adjusted p values (q values) are Benjamini–Hochberg, computed separately for the two per-protocol baseline seronegative cohort analyses defined by All features and nAb-CoP-hypothesis features, and within each of these analysis types separately for each of the classes of defined sequence feature sets given in Section 2.11 of the SAP. Significant results are marked at two levels of evidence, the higher evidence being FWER $p \leq 0.05$ and the lesser evidence being all three outcomes of unadjusted p value ≤ 0.05 , q value ≤ 0.20 , and FWER $p > 0.05$. Results in the text with q value ≤ 0.20 are only reported as such if also the unadjusted p value ≤ 0.05 .

Classification sieve analysis

For the classification sieve analyses assessing multivariable viral predictors of the treatment arm, we estimated multivariable prediction functions using a Super Learner ensemble⁶⁹ and performed a variable importance analysis, defining variable importance as the difference in area under the receiver operating characteristic curve between including versus excluding a group of virus features in COVID-19 endpoint cases. We computed point estimates and p values for each group. R⁵⁷ (version 4.3.1) was used. Information on the R packages used is provided in the Supplementary Methods.

Deep mutational scanning (DMS) antibody-escape scores

Various antibody-escape scores representing how mutations in RBD impact antibody binding²⁶ (and hence likely relevant to neutralization) were calculated. This involved two steps: identification of epitope-specific escape scores based on putative antibody-epitope footprints, and subsequent calculation of the antibody-escape scores for each sequence. These steps are detailed in Supplementary Methods.

Protein Data Bank (PDB) antibody-escape scores

The PDB antibody-escape scores are based on sets of putative antibody footprint sites, which are associated with Spike-antibody structural interactions. These site sets were determined from unique SARS-CoV-2 Spike and human anti-Spike antibody complexes that were downloaded from the PDB database (<https://www.rcsb.org/>) ($n = 274$ on May 4, 2022). For each PDB complex, epitope sites were defined as antigen sites that are in contact with the antibody in the antigen-antibody

complex (i.e., all sites that have non-hydrogen atoms within 4 Å of the antibody). Quantitation of the interaction between an epitope site and the antibody is detailed in Section 2.1.5 of the SAP and Supplementary Methods.

Neutralization hypothesis-driven sieve analysis

Mutations, substitutions, insertions and/or deletions at the following locations were identified as impacting neutralization in *in vitro* experiments: residues at AA positions 14–20, 140–158, and 245–264 that encompass the supersite epitope; mutations in the NTD signal peptide that may impact where the signal peptide is cleaved off (e.g., S12P or S13I); residues at positions 12–13, 14–26, 138–158, 242–264; D80 mutations; deletions at positions 69 and 70; mutation T95I; and R190 mutations. Based on these observations, we defined the following seven dichotomous NTD features:

- NTD1: One or more deletions or insertions in positions 138–158 and 242–264;
- NTD2: Mutations at positions 12 or 13, which may delay cleavage of the signal peptide;
- NTD3: One or more of the following mutations, R246X, G142X, K147X, or L18P;
- NTD4: One or more substitutions in 138–158 excluding positions 142 and 147;
- NTD5: One or more substitutions in 14–26 and 242–264 excluding substitutions R246X and L18P;
- NTD6: One or more substitutions D80X or L18F;
- NTD7: One or more substitutions that add or remove a glycan sequon within the NTD.

Further details are given in Section 2.1.6 of the SAP.

Covariability analysis

The covariability of two AA positions is quantified by the normalized mutational information M_{star} , which has a range from 0 to 1²¹. M_{star} equals the likelihood ratio statistic for testing the independence of the AA changes at the two AA positions, normalized to weight the positions equally regardless of diversity. For calculating the covariability of an AA position with a lineage, M_{star} is calculated in the same way as for a pair of AA positions, using two levels of the lineage variable present vs. absent.

Phylogenetic trees

The overall phylogeny was reconstructed by approximate maximum likelihood with FastTree v2.1.11 compiled with double precision⁷⁰, under the Jones–Taylor–Thornton (JTT) substitution model and CAT approximation with 20 rate categories⁷¹. This tree was rooted by the Reference group, and the sequences for the vaccine insert and the reference sequence [GenBank accession number NC_045512 (<https://www.ncbi.nlm.nih.gov/nuccore/1798174254>)] were included for comparison. The trees for each region were extracted and then visualized using the *ggtree*⁷² and *patchwork*⁷³ packages in R⁵⁷ (version 4.1.1.4).

GenSig

We used the LANL tool GenSig to identify signature sites associated with vaccine status after a phylogenetic correction that accounts for potentially spurious associations due to lineage effects^{30,31}. Further details are in the Supplementary Methods.

Production of recombinant glycoproteins

The SARS-CoV-2 S Lambda ectodomain contains Lambda mutations G75V, T76I, R246N, 247–253del, L452Q, F490S, D614G, and T859N, in addition to Hexapro stabilizing mutations⁷⁴, and DS RBD stapling mutations³⁷. S trimer was produced in 100 mL cultures of Expi293F Cells (Thermo Fisher Scientific, #A14527) grown in suspension using Expi293 Expression Medium (Thermo Fisher Scientific) at 37 °C in a humidified 8% CO₂ incubator rotating at 130 rpm. Cells grown to a

density of 3 million cells per mL were transfected using the ExpiFectamine 293 Transfection Kit (Thermo Fisher Scientific) and cultivated for 4 days, at which point the supernatant was harvested. S ectodomains were purified from clarified supernatants using a Cobalt affinity column (Cytiva, HiTrap TALON crude), washing with 20 column volumes of 20 mM Tris-HCl pH 8.0 and 150 mM NaCl, and eluted with 20 mM Tris-HCl pH 8.0, 150 mM NaCl, and 600 mM imidazole. The S ectodomain was then concentrated using a 100 kDa centrifugal filter (Amicon Ultra 0.5 mL centrifugal filters, MilliporeSigma), residual imidazole was washed away by consecutive dilutions in the centrifugal filter unit with 20 mM Tris-HCl pH 8.0 and 150 mM NaCl, and finally concentrated to 1 mg/mL before use immediately after purification. The RBD construct was based on reported constructs (with the exception of Lambda L452Q and F490S mutations in the RBD) and were produced and biotinylated as previously⁷⁵. The hACE2 construct was synthesized by GenScript into pCMV (residues 19–615 from UniProt Q9BYF1 with a C-terminal AviTag-10xHis-GGG-tag, and N-terminal signal peptide) and was expressed in HEK293.sus using standard methods (ATUM Bio). The hACE2 protein was purified via Ni Sepharose resin followed by isolation of the monomeric hACE2 by size exclusion chromatography using a Superdex 200 Increase 10/300 GL column pre-equilibrated with PBS.

CryoEM sample preparation and data collection

Fabs were generated by LysC digestion [1:3000 (w/w) antibody:LysC] for 16 h at 37 °C. Next, 50 µL of 2 mg/mL SARS-CoV-2 S Lambda ectodomain was incubated with 40 µL 3.4 mg/mL S309 Fab, 3.6 µL 28 mg/mL S2X303 Fab, and 2.2 µL of 67 mg/mL S2L20 Fab in 150 mM NaCl and 20 mM Tris-HCl pH 8 for 15 min at 37 °C. Unbound Fab was then washed away with six consecutive dilutions in 400 µL of 20 mM Tris-HCl pH 8.0 and 150 mM NaCl over a 100 kDa centrifugal filter (Amicon Ultra 0.5 mL centrifugal filters, MilliporeSigma). The complex was concentrated to 3.5 mg/mL and 3 µL was immediately applied onto a freshly glow discharged 2.0/2.0 UltraFoil grid (84) (200 mesh), plunge frozen using a Vitrobot Mark IV (Thermo Fisher Scientific) using a blot force of –1 and 6.0 s blot time at 100% humidity and 23 °C. Data were acquired using the Legion software⁷⁶ to control a FEI Titan Krios transmission electron microscope equipped with a Gatan K3 direct detector and operated at 300 kV with a Gatan Quantum GIF energy filter. The dose rate was adjusted to 3.75 counts/super-resolution pixel/s, and each movie was acquired in 75 frames of 40 ms with a pixel size of 0.843 Å and a defocus range comprised between –0.4 and –2.0 µm.

CryoEM data processing

Movie frame alignment, estimation of the microscope contrast-transfer function parameters, particle picking, and extraction (with a down-sampled pixel size of 1.686 Å and box size of 256 pixels²) were carried out using Warp⁷⁷. Reference-free 2D classification was performed using cryoSPARC⁷⁸ to select well-defined particle images. 3D classification with 50 iterations each (angular sampling 7.5° for 25 iterations and 1.8° with local search for 25 iterations) were carried out using Relion without imposing symmetry. 3D refinements were carried out using non-uniform refinement in cryoSPARC (62) before particle images were subjected to Bayesian polishing using Relion⁷⁹, during which particles were re-extracted with a box size of 512 Å at a pixel size of 0.843 Å. Next, 86 optics groups were defined based on the beam tilt angle used for data collection. Another round of non-uniform refinement in cryoSPARC was then performed concurrently with global and per-particle defocus refinement. For focused classification of the NTD and RBD, particles were symmetry-expanded and 3D classified in Relion without alignment, and then particles in well-formed 3D classes were then used for local refinement in cryoSPARC. For the NTD, the mask that encompasses the NTD, the S2L20 VH/VL region, and the S2X303 VH/VL region. For the RBD, the mask encompasses the RBD and the S309 VH/VL region. Reported resolutions are based on the

gold-standard Fourier shell correlation of 0.143 criterion and Fourier shell correlation curves were corrected for the effects of soft masking by high-resolution noise substitution^{80,81}.

Supplementary Table 28 provides information on cryoEM data collection, refinement, and validation statistics.

CryoEM model building and analysis

UCSF Chimera⁸² and Coot⁸³ were used to fit atomic models of S2L20, S309, and SARS-CoV-2 S (PDB 7SOB) into the cryoEM maps. The model was then refined and rebuilt into the map using Coot, Rosetta^{84,85}, and ISOLDE⁸⁶. Model validation and analysis used Phenix⁸⁷. Figures were generated using UCSF ChimeraX⁸⁸.

ACE2 binding measurements using Biolayer interferometry

Lambda RBD was biotinylated and immobilized at 5 ng/μL in undiluted 10X kinetics buffer (Pall) to SA sensors that were pre-hydrated in water for 10 min and then equilibrated into 10X Kinetics Buffer (Pall). The RBDs were loaded to a level of 1 nm total shift. The loaded tips were then dipped into a dilution series of monomeric ACE2-his in 10X Kinetics Buffer (Pall) starting at 1000 or 5000 nM for 300 s prior to 300 s dissociation in 10X Kinetics buffer for kinetics determination. The data were baseline subtracted, and the plots were fitted using the Pall FortéBio/Sartorius analysis software (v.12.0). Data were plotted in GraphPad Prism (v.9.0.2).

ACE2 binding measurements using surface plasmon resonance

RBD:ACE2 affinity measurements were performed using a Biacore T200 instrument. The Cytiva Biotin CAPture Kit, Series S, was used for surface capture of biotinylated RBD. The running buffer was HBS-EP+ pH 7.4 (Cytiva) and measurements were performed at 25 °C. Experiments were performed with a threefold dilution series of monomeric hACE2: 300, 100, 33, 11, 3.7 nM. The association was 300 s and dissociation was 450 s. Data were double reference-subtracted and fit to a 1:1 binding model using Biacore Evaluation software.

ELISA

For ELISA experiments with NTD-targeted mAbs, 384-well Maxisorp plates (Thermo Fisher Scientific) were coated overnight at 4 °C with 2 μg/mL of S glycoprotein in 20 mM HEPES pH 8 and 150 mM NaCl. For ELISA experiments with RBD-targeted mAbs, 384-well Maxisorp plates (Thermo Fisher Scientific) were coated overnight at 4 °C with 4 μg/mL of hACE2-His in 20 mM Sodium Phosphate pH 8 and 100 mM NaCl. The antibodies that were used were purified previously^{13,36}. Plates were slapped dry and blocked with Blocker Casein in TBS (Thermo Fisher Scientific 37532) for one hour at 37 °C. Plates were slapped dry and mAbs were serially diluted in TBST with an initial concentration of 50 μg/ml. Plates were left for one hour at 37 °C and washed 4X with TBST, then 1:5000 Goat anti-Human (Thermo Fisher Scientific A18817) was added. Plates were left for 1 h at 37 °C and washed 4x with TBST, and then TMB Microwell Peroxidase (Seracare 5120-0083) was added. The reaction was quenched after 4 min with 1 N HCl and the A450 of each well was read using a BioTek plate reader.

Sex and gender in reporting

Information on participant sex was self-reported, solicited, and collected by four predefined options (female, male, unknown, and intersex).

Sadoff et al.¹⁷ determined that sex had no meaningful impact on vaccine efficacy. As such, we scoped our sieve analyses accordingly, as any finding of viral features impacting vaccine efficacy by sex would likely be a false discovery or would need to be interpreted as a qualitative interaction.

Inclusion and ethics

The COV3001 (ENSEMBLE) study was reviewed and approved by all relevant local ethics committees and Institutional Review Boards. All

participants provided written informed consent. All experiments were performed in accordance with the relevant guidelines and regulations.

Site PIs were invited as co-authors according to the enrollments performed in the study, and were given the opportunity for intellectual contribution.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

The sequence data used in this study are available in two groups: Information pertaining to the SARS-CoV-2 sequences obtained from study participants, including their GISAID accession numbers, is provided in the Supplementary Data 1 file. The sequences curated by LANL to define the canonical variant sequences are available on GISAID through identifier EPI_SET_221208syn (<https://doi.org/10.55876/gis8.221208syn>)⁸⁹. Available information includes contributors' details, such as accession number, virus name, collection date, originating lab, submitting lab, and the list of authors.

The deep mutational scanning (DMS) data used to identify the DMS antibody-escape scores are available at https://raw.githubusercontent.com/jbloomlab/SARS2_RBD_Ab_escape_maps/651fe6fa5a7fccec2b662ddb45b6d2c7421ae74/processed_data/escape_calculator_data.csv. The representative Protein Data Bank (PDB) complexes for the PDB escape scores (Supplementary Table 2) are available from the PDB (<https://www.rcsb.org/>).

The cryoEM structures have been deposited at the PDB (<https://www.rcsb.org/>) and at the EMDB under the following accession numbers: D_1000281320 = Lambda global refinement, PDB: 8VYE, EMDB: EMD-43658; D_1000281321 = Lambda NTD local refinement, PDB: 8VYF, EMDB: EMD-43659; D_1000281322 = Lambda RBD local refinement, PDB: 8VYG, EMDB: EMD-43660.

Source data for Fig. 8 and Supplementary Figs. 53, 54 are provided with this paper.

The data sharing policy of Janssen Pharmaceutical Companies of Johnson & Johnson is available at <https://www.janssen.com/clinical-trials/transparency>. The data needed to execute the custom code for the sieve analysis as well as the neutralizing antibody data supporting the findings of this study, are proprietary to Janssen and may be obtained from the authors upon reasonable request as determined by an agreement with Yale Open Data Access (YODA) Project to serve as the independent review panel for evaluation of data requests. Project metrics for past data requests via YODA are available at <https://yoda.yale.edu/metrics/>. Source data are provided with this paper.

Code availability

All custom code for the sieve analysis, including code for: the unsupervised learning of the treatment-blinded trial sequence data to fully specify and down-select the set of amino acid (AA) sequence features that were studied for sieve effects, implementing hazard-based sieve analysis, covariability analysis of any pairs of AA positions, the SuperLearner-based supervised learning sieve analysis, reproducing the figures in the supplemental material, calculating the epitope distance analyses, and generating the structural visualizations in the manuscript is publicly available at Figshare (<https://doi.org/10.6084/m9.figshare.24911373.v1>)⁶⁴.

References

1. Feikin, D. R. et al. Duration of effectiveness of vaccines against SARS-CoV-2 infection and COVID-19 disease: results of a systematic review and meta-regression. *Lancet* **399**, 924–944 (2022).
2. Ssentongo, P. et al. SARS-CoV-2 vaccine effectiveness against infection, symptomatic and severe COVID-19: a systematic review and meta-analysis. *BMC Infect. Dis.* **22**, 439 (2022).

3. Gilbert, P., Self, S., Rao, M., Naficy, A. & Clemens, J. Sieve analysis: methods for assessing from vaccine trial data how vaccine efficacy varies with genotypic and phenotypic pathogen variation. *J. Clin. Epidemiol.* **54**, 68–85 (2001).
4. Rolland, M. & Gilbert, P. B. Sieve analysis to understand how SARS-CoV-2 diversity can impact vaccine protection. *PLoS Pathog.* **17**, e1009406 (2021).
5. Pajon, R. et al. Initial analysis of viral dynamics and circulating viral variants during the mRNA-1273 Phase 3 COVE trial. *Nat. Med.* **28**, 823–830 (2022).
6. Sadoff, J. et al. Final analysis of efficacy and safety of single-dose Ad26.COV2.S. *N. Engl. J. Med.* **386**, 847–860 (2022).
7. Cao, L. et al. Rapid evaluation of COVID-19 vaccine effectiveness against symptomatic infection with SARS-CoV-2 variants by analysis of genetic distance. *Nat. Med.* **28**, 1715–1722 (2022).
8. Greaney, A. J. et al. The SARS-CoV-2 Delta variant induces an antibody response largely focused on class 1 and 2 antibody epitopes. *PLoS Pathog.* **18**, e1010592 (2022).
9. Greaney, A. J., Starr, T. N. & Bloom, J. D. An antibody-escape estimator for mutations to the SARS-CoV-2 receptor-binding domain. *Virus Evol.* **8**, veac021 (2022).
10. Greaney, A. J. et al. A SARS-CoV-2 variant elicits an antibody response with a shifted immunodominance hierarchy. *PLoS Pathog.* **18**, e1010248 (2022).
11. Alenquer, M. et al. Signatures in SARS-CoV-2 spike protein conferring escape to neutralizing antibodies. *PLoS Pathog.* **17**, e1009772 (2021).
12. Greaney, A. J. et al. Comprehensive mapping of mutations in the SARS-CoV-2 receptor-binding domain that affect recognition by polyclonal human plasma antibodies. *Cell Host Microbe* **29**, 463–476 e466 (2021).
13. McCallum, M. et al. N-terminal domain antigenic mapping reveals a site of vulnerability for SARS-CoV-2. *Cell* **184**, 2332–2347 e2316 (2021).
14. Gilbert, P. B. et al. A Covid-19 milestone attained — a correlate of protection for vaccines. *N. Engl. J. Med.* **387**, 2203–2206 (2022).
15. Cromer, D. et al. Neutralising antibody titres as predictors of protection against SARS-CoV-2 variants and the impact of boosting: a meta-analysis. *Lancet Microbe* **3**, e52–e61 (2022).
16. Cromer, D. et al. Predicting vaccine effectiveness against severe COVID-19 over time and against variants: a meta-analysis. *Nat. Commun.* **14**, 1633 (2023).
17. Sadoff, J. et al. Safety and efficacy of single-dose Ad26.COV2.S vaccine against Covid-19. *N. Engl. J. Med.* **384**, 2187–2201 (2021).
18. Heng, F., Sun, Y. & Gilbert, P. B. Estimation and hypothesis testing of strain-specific vaccine efficacy with missing strain types, with applications to a COVID-19 vaccine trial. Preprint at [arXiv:2201.08946 \[stat.ME\]](https://arxiv.org/abs/2201.08946) (2022).
19. Juraska, M. & Gilbert, P. B. Mark-specific hazard ratio model with missing multivariate marks. *Lifetime Data Anal.* **22**, 606–625 (2016).
20. Gilbert, P. B. et al. ENSEMBLE phase 3 trial sieve analysis statistical analysis plan. [figshare <https://doi.org/10.6084/m9.figshare.21920652.v1>](https://doi.org/10.6084/m9.figshare.21920652.v1) (2023).
21. Gilbert, P. B., Novitsky, V. & Essex, M. Covariability of selected amino acid positions for HIV type 1 subtypes C and B. *AIDS Res. Hum. Retroviruses* **21**, 1016–1030 (2005).
22. Deng, X. et al. Transmission, infectivity, and neutralization of a spike L452R SARS-CoV-2 variant. *Cell* **184**, 3426–3437 e3428 (2021).
23. Liu, Y. et al. Neutralizing activity of BNT162b2-elicited serum. *N. Engl. J. Med.* **384**, 1466–1468 (2021).
24. Acevedo, M. L. et al. Differential neutralizing antibody responses elicited by CoronaVac and BNT162b2 against SARS-CoV-2 Lambda in Chile. *Nat. Microbiol.* **7**, 524–529 (2022).
25. Motozono, C. et al. SARS-CoV-2 spike L452R variant evades cellular immunity and increases infectivity. *Cell Host Microbe* **29**, 1124–1136 e1111 (2021).
26. Starr, T. N. et al. Deep mutational scanning of SARS-CoV-2 receptor binding domain reveals constraints on folding and ACE2 binding. *Cell* **182**, 1295–1310 e1220 (2020).
27. Sadoff, J. et al. Durable antibody responses elicited by 1 dose of Ad26.COV2.S and substantial increase after boosting: 2 randomized clinical trials. *Vaccine* **40**, 4403–4411 (2022).
28. Jongeneelen, M. et al. Ad26.COV2.S elicited neutralizing activity against Delta and other SARS-CoV-2 variants of concern. Preprint at [bioRxiv <https://doi.org/10.1101/2021.07.01.450707>](https://doi.org/10.1101/2021.07.01.450707) (2021).
29. Williamson, B. D., Gilbert, P. B., Simon, N. R. & Carone, M. A general framework for inference on algorithm-agnostic variable importance. *J. Am. Stat. Assoc.* **118**, 1645–1658 (2023).
30. Bricault, C. A. et al. HIV-1 neutralizing antibody signatures and application to epitope-targeted vaccine design. *Cell Host Microbe* **25**, 59–72 e58 (2019).
31. Bhattacharya, T. et al. Founder effects in the assessment of HIV polymorphisms and HLA allele associations. *Science* **315**, 1583–1586 (2007).
32. He, P. et al. SARS-CoV-2 Delta and Omicron variants evade population antibody response by mutations in a single spike epitope. *Nat. Microbiol.* **7**, 1635–1649 (2022).
33. Cao, Y. et al. Omicron escapes the majority of existing SARS-CoV-2 neutralizing antibodies. *Nature* **602**, 657–663 (2022).
34. Zhou, B. et al. A broadly neutralizing antibody protects Syrian hamsters against SARS-CoV-2 Omicron challenge. *Nat. Commun.* **13**, 3589 (2022).
35. Pinto, D. et al. Cross-neutralization of SARS-CoV-2 by a human monoclonal SARS-CoV antibody. *Nature* **583**, 290–295 (2020).
36. McCallum, M. et al. Molecular basis of immune evasion by the Delta and Kappa SARS-CoV-2 variants. *Science* **374**, 1621–1626 (2021).
37. McCallum, M., Walls, A. C., Bowen, J. E., Corti, D. & Velesler, D. Structure-guided covalent stabilization of coronavirus spike glycoprotein trimers in the closed conformation. *Nat. Struct. Mol. Biol.* **27**, 942–949 (2020).
38. Greaney, A. J. et al. Complete mapping of mutations to the SARS-CoV-2 spike receptor-binding domain that escape antibody recognition. *Cell Host Microbe* **29**, 44–57 e49 (2021).
39. Kimura, I. et al. The SARS-CoV-2 Lambda variant exhibits enhanced infectivity and immune resistance. *Cell Rep.* **38**, 110218 (2022).
40. Wang, M. et al. Reduced sensitivity of the SARS-CoV-2 Lambda variant to monoclonal antibodies and neutralizing antibodies induced by infection and vaccination. *Emerg. Microbes Infect.* **11**, 18–29 (2022).
41. Liu, H. et al. The receptor binding domain of SARS-CoV-2 lambda variant has a better chance than the delta variant in evading BNT162b2 COVID-19 mRNA vaccine-induced humoral immunity. *Int. J. Mol. Sci.* <https://doi.org/10.3390/ijms231911325> (2022).
42. Khoury, D. S. et al. Correlates of protection, thresholds of protection, and immunobridging among persons with SARS-CoV-2 infection. *Emerg. Infect. Dis.* **29**, 381–388 (2023).
43. Gilbert, P. B. et al. Immune correlates analysis of the mRNA-1273 COVID-19 vaccine efficacy clinical trial. *Science* **375**, 43–50 (2022).
44. Fong, Y. et al. Immune correlates analysis of the ENSEMBLE single Ad26.COV2.S dose vaccine efficacy clinical trial. *Nat. Microbiol.* **7**, 1996–2010 (2022).
45. Fong, Y. et al. Immune correlates analysis of the PREVENT-19 COVID-19 vaccine efficacy clinical trial. *Nat. Commun.* **14**, 331 (2023).
46. Benkeser, D. et al. Immune correlates analysis of a phase 3 trial of the AZD1222 (ChAdOx1 nCoV-19) vaccine. *NPJ Vaccines* **8**, 36 (2023).

47. Khoury, D. S. et al. Neutralizing antibody levels are highly predictive of immune protection from symptomatic SARS-CoV-2 infection. *Nat. Med.* **27**, 1205–1211 (2021).
48. Tarke, A. et al. SARS-CoV-2 vaccination induces immunological T cell memory able to cross-recognize variants from Alpha to Omicron. *Cell* **185**, 847–859 e811 (2022).
49. Korber, B. et al. Tracking changes in SARS-CoV-2 spike: evidence that D614G increases infectivity of the COVID-19 virus. *Cell* **182**, 812–827 e819 (2020).
50. Wang, Q. et al. Alarming antibody evasion properties of rising SARS-CoV-2 BQ and XBB subvariants. *Cell* **186**, 279–286 e278 (2023).
51. Corey, L. et al. SARS-CoV-2 variants in patients with immunosuppression. *N. Engl. J. Med.* **385**, 562–566 (2021).
52. Milne, G. et al. Does infection with or vaccination against SARS-CoV-2 lead to lasting immunity? *Lancet Respir. Med.* **9**, 1450–1466 (2021).
53. Alter, G. et al. Immunogenicity of Ad26.COVS vaccine against SARS-CoV-2 variants in humans. *Nature* **596**, 268–272 (2021).
54. Yu, X. et al. Convergence of immune escape strategies highlights plasticity of SARS-CoV-2 spike. *PLoS Pathog.* **19**, e1011308 (2023).
55. Aksamentov, I., Roemer, C., Hodcroft, E. B. & Neher, R. A. Nextclade: clade assignment, mutation calling and quality control for viral genomes. *J. Open Source Softw.* **6**, 3773 (2021).
56. O’Toole, A. et al. Assignment of epidemiological lineages in an emerging pandemic using the pangolin tool. *Virus Evol.* **7**, veab064 (2021).
57. R Core Team. R: a language and environment for statistical computing. R Foundation for Statistical Computing. <https://www.R-project.org/> (2022).
58. Charif, D. & Lobry, J. in *Structural Approaches to Sequence Evolution: Molecules, Networks, Populations* (eds Bastolla, U., Porto, M., Roman, H. & Vendruscolo, M.) Ch. 10 (Springer, 2007).
59. Rao, C. R. Diversity and dissimilarity coefficients: a unified approach. *Theor. Popul. Biol.* **21**, 24–43 (1982).
60. Humphrey, W., Dalke, A. & Schulten, K. VMD: visual molecular dynamics. *J. Mol. Graph.* **14**, 33–38 (1996). 27–38.
61. Harris, C. R. et al. Array programming with NumPy. *Nature* **585**, 357–362 (2020).
62. Michaud-Agrawal, N., Denning, E. J., Woolf, T. B. & Beckstein, O. MDAnalysis: a toolkit for the analysis of molecular dynamics simulations. *J. Comput. Chem.* **32**, 2319–2327 (2011).
63. Gowers, R. J. et al. MDAnalysis: a Python package for the rapid analysis of molecular dynamics simulations. In *Proc. 15th Python in Science Conference* (eds Benthall, S. & Rostrup, S.) 98–105 (SciPy, 2016).
64. Magaret, C. A. et al. Code used to execute the SARS-CoV-2 sieve analysis for the ENSEMBLE (CoVNP 3003) efficacy trial of the Janssen Ad26.COVS vaccine. figshare (software). <https://doi.org/10.6084/m9.figshare.24911373.v1> (2024).
65. Gilbert, P. B., Self, S. G. & Ashby, M. A. Statistical methods for assessing differential vaccine protection against human immunodeficiency virus types. *Biometrics* **54**, 799–814 (1998).
66. RStudio Team. RStudio: integrated development for R. RStudio. <http://www.rstudio.com/> (2023).
67. Juraska, M. & Gilbert, P. B. Mark-specific hazard ratio model with multivariate continuous marks: an application to vaccine efficacy. *Biometrics* **69**, 328–337 (2013).
68. Gray, B. cmprsk: subdistribution analysis of competing risks. R package. <https://cran.r-project.org/package=cmprsk> (2022).
69. van der Laan, M. J., Polley, E. C. & Hubbard, A. E. Super learner. *Stat. Appl. Genet. Mol. Biol.* <https://doi.org/10.2202/1544-6115.1309> (2007).
70. Price, M. N., Dehal, P. S. & Arkin, A. P. FastTree 2—approximately maximum-likelihood trees for large alignments. *PLoS ONE* **5**, e9490 (2010).
71. Jones, D. T., Taylor, W. R. & Thornton, J. M. The rapid generation of mutation data matrices from protein sequences. *Comput. Appl. Biosci.* **8**, 275–282 (1992).
72. Yu, G. C., Smith, D. K., Zhu, H. C., Guan, Y. & Lam, T. T. Y. GGTREE: an R package for visualization and annotation of phylogenetic trees with their covariates and other associated data. *Methods Ecol. Evol.* **8**, 28–36 (2017).
73. Pedersen, T. patchwork: the composer of plots. <https://patchwork.data-imaginist.com>, <https://github.com/thomasp85/patchwork> (2023).
74. Hsieh, C. L. et al. Structure-based design of prefusion-stabilized SARS-CoV-2 spikes. *Science* **369**, 1501–1505 (2020).
75. Walls, A. C. et al. Structure, function, and antigenicity of the SARS-CoV-2 spike glycoprotein. *Cell* **181**, 281–292 e286 (2020).
76. Suloway, C. et al. Automated molecular microscopy: the new Legimon system. *J. Struct. Biol.* **151**, 41–60 (2005).
77. Tegunov, D. & Cramer, P. Real-time cryo-electron microscopy data preprocessing with Warp. *Nat. Methods* **16**, 1146–1152 (2019).
78. Punjani, A., Rubinstein, J. L., Fleet, D. J. & Brubaker, M. A. cryoSPARC: algorithms for rapid unsupervised cryo-EM structure determination. *Nat. Methods* **14**, 290–296 (2017).
79. Zivanov, J., Nakane, T. & Scheres, S. H. W. A Bayesian approach to beam-induced motion correction in cryo-EM single-particle analysis. *IUCrJ* **6**, 5–17 (2019).
80. Rosenthal, P. B. & Henderson, R. Optimal determination of particle orientation, absolute hand, and contrast loss in single-particle electron cryomicroscopy. *J. Mol. Biol.* **333**, 721–745 (2003).
81. Chen, S. et al. High-resolution noise substitution to measure overfitting and validate resolution in 3D structure determination by single particle electron cryomicroscopy. *Ultramicroscopy* **135**, 24–35 (2013).
82. Pettersen, E. F. et al. UCSF Chimera—a visualization system for exploratory research and analysis. *J. Comput. Chem.* **25**, 1605–1612 (2004).
83. Emsley, P., Lohkamp, B., Scott, W. G. & Cowtan, K. Features and development of Coot. *Acta Crystallogr. D. Biol. Crystallogr.* **66**, 486–501 (2010).
84. Frenz, B. et al. Automatically fixing errors in glycoprotein structures with Rosetta. *Structure* **27**, 134–139 e133 (2019).
85. Wang, R. Y. et al. Automated structure refinement of macromolecular assemblies from cryo-EM maps using Rosetta. *Elife* <https://doi.org/10.7554/eLife.17219> (2016).
86. Croll, T. I. ISOLDE: a physically realistic environment for model building into low-resolution electron-density maps. *Acta Crystallogr. D. Struct. Biol.* **74**, 519–530 (2018).
87. Liebschner, D. et al. Macromolecular structure determination using X-rays, neutrons and electrons: recent developments in Phenix. *Acta Crystallogr. D. Struct. Biol.* **75**, 861–877 (2019).
88. Goddard, T. D. et al. UCSF ChimeraX: meeting modern challenges in visualization and analysis. *Protein Sci.* **27**, 14–25 (2018).
89. GISAIID Identifier: EPI_SET_221208yn. <https://doi.org/10.55876/gis8.221208yn> (2022).

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Competing interests

ALG reports contract testing from Abbott, Cepheid, Novavax, Pfizer, Janssen, and Hologic and research support from Gilead and Merck. JS declares support for the submitted work from the Janssen Pharmaceutical Companies of Johnson & Johnson and partial support (in the form of funding to his institution) from BARDA for the submitted work, declares support within the past 36 months from the Janssen Pharmaceutical Companies of Johnson & Johnson and BARDA funding for part of this work, has patents (US 11,384,122 B2) on invention of the Janssen COVID-19 vaccine, and has Johnson & Johnson stock and stock options. SR had

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Ethics committees

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Additional information

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