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1 Title: Protein Vaccine Induces a Durable, More Broadly Neutralizing Antibody Response 2 in Macagues than Natural Infection with SARS-CoV-2 P.1 3 4 Authors: Albert To¹, Teri Ann S. Wong¹, Michael M. Lieberman¹, Karen Thompson², Laurent 5 Pessaint³, Jack Greenhouse³, Nisrine Daham³, Anthony Cook³, Brandon Narvaez³, Zack 6 Flinchbaugh³, Alex Van Ry³, Jake Yalley-Ogunro³, Hanne Andersen Elyard³, Chih-Yun Lai¹, 7 Oreola Donini⁴, Axel T. Lehrer^{1*} 8 9 Affiliations: 10 ¹Department of Tropical Medicine, Medical Microbiology, and Pharmacology, John A. Burns 11 School of Medicine, University of Hawai'i at Mānoa, Honolulu, HI, USA 12 ²Department of Pathology, John A. Burns School of Medicine, University of Hawai'i at Mānoa, 13 Honolulu, HI, USA 14 ³Bioqual Inc., Rockville, MD, USA 15 ⁴Soligenix, Inc, Princeton, NJ, USA 16 17 *Corresponding author at: Axel T. Lehrer: (lehrer@hawaii.edu). University of Hawai'i at Mānoa, 18 John A. Burns School of Medicine, Department of Tropical Medicine, 651 Ilalo Street, Honolulu, 19 HI 96813, United States 20 21 One Sentence Summary: A recombinant subunit protein formulated with CoVaccine HT[™] 22 adjuvant induces superior immunity than natural infection and reduces viral load while protecting 23 cynomolgus macaques from COVID-19-like disease caused by late SARS-CoV-2 P.1 (Gamma) 24 challenge.

25

26 Abstract: FDA-approved and Emergency Use Authorized (EUA) vaccines using new mRNA and viral-vector technology are highly effective in preventing moderate to severe disease, 27 28 however, information on their long-term efficacy and protective breadth against SARS-CoV-2 29 Variants of Concern (VOCs) is currently scarce. Here we describe the durability and broad-30 spectrum VOC immunity of a prefusion-stabilized spike (S) protein adjuvanted with liquid or 31 lyophilized CoVaccine HT[™] in cynomolgus macagues. This recombinant subunit vaccine is 32 highly immunogenic and induces robust spike-specific and broadly neutralizing antibody 33 responses effective against circulating VOCs (B.1.351 [Beta], P.1 [Gamma], B.1.617 [Delta]) for 34 at least 3 months after the final boost. Protective efficacy and post-exposure immunity were 35 evaluated using a heterologous P.1 challenge nearly 3 months after the last immunization. Our 36 results indicate that while immunization with both high and low S doses shorten and reduce viral 37 loads in the upper and lower respiratory tract, a higher antigen dose is required to provide 38 durable protection against disease as vaccine immunity wanes. Histologically, P.1 infection 39 causes similar COVID-19-like lung pathology as seen with early pandemic isolates. Post-40 challenge IgG concentrations were restored to peak immunity levels and vaccine-matched and 41 cross-variant neutralizing antibodies were significantly elevated in immunized macaques 42 indicating an efficient anamnestic response. Only low levels of P.1-specific neutralizing 43 antibodies with limited breadth were observed in control (non-vaccinated but challenged) 44 macaques suggesting that natural infection may not prevent reinfection by other VOCs. Overall, 45 these results demonstrate that a properly dosed and adjuvanted recombinant subunit vaccine 46 can provide long-lasting and protective immunity against circulating VOCs.

47

48 Main Text:

49

50 INTRODUCTION

51 COVID-19 is a respiratory disease caused by the Severe Acute Respiratory Syndrome
52 Coronavirus 2 (SARS-CoV-2) (*1*, *2*) and is commonly characterized as an asymptomatic

53 infection or a self-limiting, febrile illness co-presenting with cough, shortness of breath, and 54 fatigue (3, 4). Severe complications can result in approximately 6-10% of infected patients (5) 55 developing pneumonia, acute respiratory distress syndrome, multi-organ dysfunction and 56 arterial thromboembolic events that can result in hospitalization and/or death (6-9). 57 Transmission occurs through respiratory droplets during close contact with asymptomatic or 58 presymptomatic infected individuals (10-12). The synergistic relationship between the infectivity 59 and high transmissibility with low virulence has contributed to the ongoing global public health 60 crisis, and the emergence of more transmissible and potentially more pathogenic variants of 61 concern (VOCs). Longitudinal cross-sectional studies have indicated only a fraction of 62 asymptomatic to mild and moderate SARS-CoV-2 natural infections of humans results in 63 detectable neutralizing antibodies months after recovery (13-15) and current evidence suggests 64 that convalescent immunity may only provide transient protection (16-18). Furthermore, the 65 degree of immune evasion witnessed in vitro by spike (S) protein variants and the increasing 66 occurrence of reinfection (19-23) has raised the question whether previous infection or vaccine-67 derived immunity can provide heterologous protection against further transmission and limit 68 additional mutation of circulating VOC, even while protection against severe disease and 69 mortality has been sustained to date.

70

71 Some VOCs show increased transmissibility, cause more severe disease than original strains 72 and demonstrate greater immune evasion to neutralizing antibodies as they harbor mutations at 73 sites facilitating viral fusion (24, 25) and at critical epitopes in the N-terminal (NTD) and receptor 74 binding domains (RBD). The B.1.1.7 (Alpha), B.1.351 (Beta) and P.1 (Gamma) variants contain 75 the same set of mutations, K417T, E484K, and N501Y, while the B.1.617 (Delta) carries L452R 76 and T478K mutations, all of which are associated with enhanced infectivity and reduced serum 77 neutralization (26-28). These antigenic changes have abolished potent neutralizing epitopes 78 targeted by monoclonal antibody therapeutics (29) and antibodies elicited by mRNA vaccination

(30). Natural reinfection with the VOC strains after original strain infection, and *vice versa*, has
been documented (*20, 31-34*). Despite this, cross-neutralization of convalescent sera from
patients infected with non-variants and mRNA immunized individuals suggests that the parental

82 spike (S) protein in current vaccines affords some degree of protection (35-37).

83

84 Phase 1-3 clinical trials of novel mRNA (Moderna mRNA-1273 and Pfizer-BioNTech BNT162b2) 85 and viral-vector (Johnson & Johnson-Janssen Ad26.COV2.S) platform, utilizing the original 86 Wuhan-Hu-1 S protein (38-40), have shown these vaccines to be highly protective with 94-95% 87 and 67% efficacy against COVID-19 disease, respectively (41-43), and they all elicit uniformly 88 high protection against severe disease/death (i.e., >85%). Post-hoc sequencing during clinical 89 trials of these and other vaccine platforms in variant-dominant regions have revealed a slightly 90 lower efficacy against B.1.1.7 and B.1.351 variants, albeit usually remaining above the 50% 91 protection threshold (44, 45). Attempts at predicting protection based on immune correlates, 92 such as cross-neutralizing antibodies, have shown that vaccinated individuals had a 2 to 5-fold 93 reduction in titers against current VOCs as IgG levels wane, which are nonetheless believed to 94 confer a degree of resistance against infection (46-49). However, these samples were taken 95 during the peak period of the humoral response and may not be accurate predictors of 96 protection months onwards when the likelihood of infection is greater. Thus, understanding the 97 decay in antibody responses months after the final boost is an essential endpoint in vaccine 98 development.

99

Evaluating a delayed vaccine response in non-human primate (NHP) models can provide
valuable foresight into the relationship between waning immunity and protection, especially
against VOCs. In this study, we describe the immunogenicity and protective efficacy of a
recombinant pre-fusion spike subunit (based on the reference SARS-CoV-2 strain, Wuhan-Huadjuvanted with liquid or lyophilized (dry) CoVaccine HT[™], a squalane-in-water

105 nanoemulsion adjuvant containing immunostimulatory sucrose fatty acid sulfate esters (50), in 106 cynomolgus macaques. Other investigators have shown that infection with SARS-CoV-2 107 (prototype strain) produced COVID-19-like disease in this species and reflects similar viral 108 shedding kinetics and lung pathology as human infections (51, 52). We have previously 109 demonstrated that our vaccine candidate elicits a broad-spectrum IgG response including high 110 neutralizing antibody (NtAb) titers against the prototypic SARS-CoV-2 and VOCs, specifically 111 B.1.351 and P.1, and an in vitro antigen-specific IFN-y secreting response from immune 112 splenocytes taken from Swiss Webster mice (53). To elaborate on our understanding of the 113 elicited immunological response, we assessed the vaccine efficacy using a delayed challenge 114 scheme with the P.1 VOC to delineate how the maturation and/or decay of the humoral 115 response affects protection from a heterologous SARS-CoV-2 strain. Neutralizing antibody 116 responses to the vaccine-matched WA1/2020 strain, and VOCs, B.1.351, P.1 and B.1.617 were 117 also determined just prior to challenge and 14 days post-challenge (to assess anamnestic 118 responses). Furthermore, we characterize the viral kinetics and histopathological changes post-119 challenge in the lower and upper respiratory tracts of control (non-vaccinated, challenged), and 120 protected NHPs. To our knowledge, this is the first published account of P.1 VOC challenge in 121 NHPs.

122

123 **RESULTS**

124 Spike antigen with either liquid or lyophilized CoVaccine HT[™] elicits a durable humoral

125 response

Twelve Cynomolgus macaques were assigned into four groups and received two immunizations with 5 (Group A) or 25 (Group B and C) µg liquid prefusion spike trimer (S) antigen formulated with either lyophilized (Group A and C) or liquid (Group B) CoVaccine HT[™] adjuvant, or one dose of a co-lyophilized CoVaccine HT[™]-adjuvanted control containing an unrelated viral glycoprotein antigen (Group D) (Fig. 1A). The two doses were administered within a three-week 131 interval and all NHPs were challenged with the P.1 isolate 12 weeks post-boost (~ 3 months, 132 study week 15). Wuhan-Hu-1 S- and RBD-specific IgG titers were measured by a multiplexed 133 microsphere immunoassay (MIA) using insect cell expressed antigens coupled onto spectrally 134 distinct, magnetic beads as described previously (53). Serum S-specific IgG concentrations 135 were interpolated using a standard curve generated from S-specific human IgG purified from 136 vaccinated individuals (Fig. 1B). RBD-specific IgG titers were read out as median fluorescence 137 intensity (MFI) (Fig. S1). All NHP immunized with the adjuvanted S at both antigen doses 138 seroconverted after the prime (week 3) with S-specific antibodies in the range of 20 to 70 139 µg/mL, and peak serum IgG concentrations detected two weeks after the boost (week 5) in the 140 range of 70 to 753 µg/mL. Macaques given a 25 µg dose of S demonstrated a greater IgG 141 response to the antigen compared to those receiving 5 µg. RBD-specific IgG titers followed a 142 similar trend. As expected, animals in group D receiving an unrelated antigen did not develop 143 any detectable S-specific IgG during this phase of the study. S-specific IgG remained detectable 144 12 weeks after the boost (week 15) although IgG concentrations dropped 3.0 to 9.9-fold relative 145 to the prior peak titer.

146

147 The titer of NtAb against the prototype WA1/2020 strain was determined using a standard 148 PRNT with wild-type virus (Fig. 1C). NtAb from one animal in Group C was detectable after the 149 prime (week 3). All S-immunized NHPs developed a potent neutralizing response, peaking two 150 weeks after the boost (week 5) and generally remaining stable one week later (week 6). The 151 group receiving 5 µg of the antigen showed the greatest variability. Nearly all immunized NHPs 152 maintained neutralization capacity, with PRNT₅₀ greater than 1:150 dilution, 12 weeks after the 153 final immunization (week 15). NtAbs from groups receiving 25 µg S either remained stable or 154 decreased up to 3.0-fold, while the group receiving 5 µg S declined ~2.9-9.2-fold. A similar 155 reduction in serum neutralization was also verified using a surrogate rVSV-SARS-CoV-2 S 156 PRNT assay (Fig. S2).

157

158	A recombinant subunit vaccine induces stable neutralizing antibodies against VOCs
159	Circulating SARS-CoV-2 VOCs can evade vaccine-induced antibody responses and are
160	associated with breakthrough infections in those fully immunized, especially as antibody titers
161	wane (<i>54-60</i>). To determine whether an adjuvanted, prototypic Wuhan-Hu-1 S subunit can
162	generate durable cross-variant neutralizing antibodies, PRNTs using WT B.1.351, P.1, and
163	B.1.617 isolates were determined with sera collected at the time of peak neutralization and 12
164	months after the final immunization, a timepoint at which antibody titers are expected to have
165	waned. At week 6, potent neutralization of the B.1.351, P.1 and B.1.617 VOC was detected,
166	although 10.7-, 10.7- and 5.7-fold lower, respectively, compared to WA1/2020 in all vaccine
167	groups (Fig. 2A). As antibody titers waned 10 weeks later (study week 15) the gap between
168	
	neutralization titers against the WA1/2020 strain to the VOC isolates narrowed to a 4.5-, 6.3-
169	and 5.1-fold difference (Fig. 2B) driven by a proportionally larger decline in WA1/2020-specific
170	neutralizing activity. At this later time point, all S-immunized macaques, except for a single
171	macaque in Group A, maintained a detectable $PRNT_{50}$ titer greater than 1:40 dilution. A few
172	macaques in Groups B and C demonstrated greater VOC neutralization at week 15 compared
173	to week 6, suggesting a refinement or "maturation" of the humoral response towards
174	neutralizing epitopes during this interval.
175	
176	Immunization with recombinant subunits reduces viral burden from delayed P.1 (Gamma)
177	challenge
178	The characterization of the prototype strain and B.1.351 variant infections in NHP models has
179	been described previously (51, 52, 61-63). Here we characterize the clinical signs and
180	histological events of a P.1 VOC infection in cynomolgus macaques and determine whether a
181	prefusion spike subunit formulated with CoVaccine HT™ is effective at reducing viral load
182	against a circulating VOC after a 12-week interval following the final immunization. All NHPs

183 were challenged with a total of 1×10^6 TCID₅₀ of the P.1 isolate using simultaneous intranasal and intratracheal inoculation routes. None of the NHPs challenged developed visible clinical 184 185 signs of respiratory disease throughout the study, consistent with WA1/2020 strain infected 186 cynomolgus macagues (51). Bronchioalveolar lavages (BAL) from the lower respiratory tract, 187 and nasal and oral swabs (NS and OS respectively) from the upper respiratory tract were 188 collected at days 2, 4, 7, 10 and 14 after challenge to detect infectious virus, genomic RNA or 189 signs of viral replication. High levels of infectious virus, GMT Log₁₀ TCID₅₀ titer of 5.8 and 4.8, 190 were recovered two days post-challenge from the NS and BAL, respectively, from control 191 macaques (Group D, Fig. 3A & 3D and S3A-B). Infectious virus continued to be detectable in 192 both anatomic sites until at least day 7 post-challenge before becoming undetectable on day 10. 193 All S-immunized NHPs presented TCID₅₀ titers at least 1-2 Log₁₀ lower than control macaques 194 in both locations at all timepoints throughout the study, indicating protection conferred by both 5 195 and 25 µg S formulations of the subunit vaccine. No infectious virus could be cultured from the 196 BAL in S-immunized macaques by day 7. Infectious virus from the NS was recoverable from 197 immunized macagues throughout the study period and was variable between groups, however 198 these macaques on average had TCID₅₀ at least 1-2 Log₁₀ below those of control animals.

199

200 Quantitative RT-PCR was used as an additional measurement for viral load. We detected the 201 presence of viral RNA and viral replication throughout the 14-day study period in both the lower 202 (Fig 3B-C, S4A-C and S5A-C) and upper (Fig 3E-F, S4D-F and S5D-F) respiratory airways and 203 from the oral cavity (Fig 3G-H, S4G-I and S5G-I). Quantification of viral RNA showed that both 5 204 and 25 µg immunization had reduced the viral load by approximately 1-2 Log₁₀ RNA copies/mL 205 (or swab) over Group D macaques, corroborating the TCID₅₀ results. Virus replication in the oral 206 cavity was undetectable at least 3 days earlier in most S-immunized macaques than in the 207 control group, suggesting a reduced potential for transmission. High levels of viral RNA and 208 replication, between 5-6 Log₁₀ vgRNA and sgRNA copies/mL, respectively, were observed in

these control macagues before gradually resolving at day 10. Altogether, reduced viral loads at

209

all tested anatomic locations in Groups A, B and C macaques indicate a degree of durable 210 211 protection from P.1 VOC infection provided by an adjuvanted protein subunit vaccine. 212 213 High dose spike antigen reduces lung histopathology caused by P.1 (Gamma) challenge 214 Also consistent with previous prototype virus infection studies in cynomolgus macaques (51, 215 52), P.1 challenge did not cause changes in weight, temperature nor demonstrate any 216 observable signs of respiratory disease. To evaluate vaccine efficacy against COVID-19-like 217 pathology, a section slide from each lung lobe and the bronchi was prepared from each 218 macague and stained with hematoxylin and eosin to survey for histopathological changes 219 indicative of acute lung injury. The histopathological scoring system based on the 220 presence/severity of edema, intraalveolar and interstitial inflammation, perivascular lymphocytic 221 cuffing, and increased bronchiolar-associated lymphoid tissue (BALT) is outlined in Table S1. A 222 cumulative average score was determined for each macaque based on the evaluation of these 223 five characteristics per section for a total of 30 scores (Fig 4A). Significant differences in the 224 cumulative average scores for each vaccine formulation vs. controls were determined using a 225 one-way ANOVA followed by a Dunnett's multiple comparison test. Macagues in the control 226 Group D appeared to have developed mild to moderate respiratory disease. The lower 227 cumulative GMT scores of Groups B and C, immunized with 25 µg S, suggests that these 228 animals, except for a single macague, were completely protected from lung pathology while all 229 Group A (immunized with 5 µg S) macagues appeared to have developed mild disease despite 230 exhibiting lower viral loads. Week 15 WA1/2020 NtAb levels were determined to be inversely 231 correlated, although weakly, with histopathology scores using a Spearman's correlation test (r=-232 0.682, p=0.0178) (Fig. 4B). Week 15 pre-challenge neutralizing antibody levels were lower in 233 animals with higher histopathology scores, a trend seen with breakthrough infections in fully

immunized individuals (64). The moderate disease seen in the control animals is demonstrated

235 in Fig. 5A, where a representative, low-power magnification shows a distribution of lymphocytic 236 perivascular cuffing and clusters of intraalveolar macrophages. In contrast, immunized animals 237 (especially those receiving the higher antigen dose), appeared to have milder disease without 238 these pathological changes (Fig. 5B). Macagues developing mild to moderate disease exhibited 239 absent to moderate, focal and multifocal edema and perivascular cuffing with marked 240 lymphocytic infiltration (Figs. 6A-D). Syncytia of intraalveolar multinucleated giant cells 241 surrounded by acute and eosinophilic inflammatory infiltrate (Fig. 6A.) or increased intraalveolar 242 macrophages with interstitial lymphocytic inflammation, in addition to alveolar septal thickening 243 and complete perivascular cuffing can be seen in a moderately diseased, control macaque (Fig. 244 6B-C). Proteinaceous edema fluid filling alveolar spaces was also observed in unprotected 245 macagues (Fig. 6D). While these histopathological changes were also noticed in protected 246 macaques, the findings appeared less severe (Fig. 6E-F). A section from a protected macaque 247 revealed a few intraalveolar multinucleated giant cells without an increase in intraalveolar 248 macrophages, lymphocytic vascular cuffing, or interstitial inflammation (Fig. 6E). Similarly, only 249 partial lymphocytic perivascular cuffing with adjacent alveolar epithelial hyperplasia was noticed 250 without an increased intraepithelial macrophage infiltrate and normal alveolar septal thickness in 251 immunized macaques receiving 25 µg of S (Fig. 6F).

252

P.1 challenge recalls a broadly neutralizing anamnestic response and a variant-specific primary neutralizing response in naïve macaques

255 To better understand the immunological basis of events happening after late challenge, and to

- determine if P.1 challenge triggered an anamnestic response in immunized NHPs, post-
- 257 challenge S-specific IgG concentrations and PRNT₅₀ titers were measured from sera collected

258 upon necropsy at day 14. Late P.1 challenge significantly boosted serum IgG concentrations to

- levels attained at the post-immunization peak at Week 6, with less variance between animals,
- suggesting that a recall response was triggered by the heterologous infection (Fig. 7A).

261 Vaccine-matched (WA1/2020) and VOC neutralizing (i.e., B.1.351, P.1 and B.1.617) antibody 262 levels were also all significantly boosted to levels attaining or surpassing the peak levels 263 generated through initial immunization by 5-fold and upwards of 20-fold respectively (Fig. 7B 264 and S2). Unsurprisingly, the greatest increase from pre-challenge neutralization was against the 265 P.1 challenge isolate. Group A showed slightly higher P.1 NtAbs compared to Group B and C 266 macagues and was the only group to have significantly higher titers across different variants. 267 The lower degree of histopathology seen in Groups B and C may indicate that 25 µg S with 268 either form of CoVaccine HT[™] provides greater long-term protection. In Group D control 269 macaques, P.1 challenge generated low levels of NtAbs compared to immunization alone, and it 270 was only cross-reactive with the B.1.351 isolate and not to WA1/2020 nor B.1.617 isolates. 271 This suggests that primary infection with one isolate of SARS-CoV-2 does not generate broad 272 immunity against other variants in contrast to vaccine derived immunity, which is more broadly 273 cross-reactive with VOCs, even months after peak immunity is observed.

274

275 **DISCUSSION**

276 Understanding the durability and breadth of vaccine-generated immunity is critical for predicting 277 long-term protection during the COVID-19 pandemic and for informing policy on strategic 278 resource deployment to facilitate equitable vaccine access. Diversifying the type of vaccines 279 currently used beyond mRNA and viral vectors to include other vaccine platforms, such as 280 protein subunit vaccines, can bolster global availability by mobilizing more thermostable 281 vaccines to resource-poor areas, or overcoming anti-vector immunity and adverse effects, by 282 using this platform as a booster to restore pre-existing natural or vaccine-induced immunity (65-283 67).

284

We have demonstrated that two doses of a subunit vaccine consisting of a prefusion S trimer
(Wuhan-Hu-1) formulated with liquid or lyophilized CoVaccine HT[™] reduces viral load and

287 provides sufficient, durable, cross-variant protection from mild to moderate disease lasting at least 3 months after the final boost. This supports previous findings in rhesus macaques that S 288 289 trimer-based subunit protein vaccines are highly immunogenic and generate long-lasting, robust 290 antibody responses (68). Furthermore, using a two-step purification method of immunoaffinity 291 chromatography followed by a size-exclusion polishing step results in a homogenous antigen 292 composition and reduces host-cell carryover plaguing conventional purification strategies for 293 protein subunits (69). Also promising is the prospect that the adjuvant, CoVaccine HT[™], can be 294 lyophilized, and reconstituted simply with water for injection and still retain functionality. This 295 technology has been previously used to develop mono- and multivalent filovirus vaccine 296 formulations. Those studies demonstrated that single-vial, lyophilized formulations 297 preserved the higher order antigen structure and the biophysical properties of the adjuvant (70, 298 71). Robust levels of S-specific IgG and homologous, as well as B.1.351, P.1 and B.1.617 299 cross-neutralizing antibodies, were detected throughout the extended study period and inversely 300 correlate with viral load and lung damage. This report supports growing evidence that the 301 Wuhan-Hu-1 S peptide sequence encoded by highly effective mRNA, viral vectored, and 302 subunit protein vaccines, generates immunity that affords protection against circulating VOCs 303 (45, 46, 70-73), even months after the final vaccine dose when antibody titers are waning. 304 Furthermore, we noticed a few macaques receiving the higher antigen dose had developed 305 increasingly potent cross-variant neutralizing titers immediately prior to viral challenge which 306 were greater than titers observed during the peak humoral response. This suggests that a 307 potentially remaining antigen depot or persistence of APC's may foster continued accumulation 308 of somatic mutations and affinity maturation in memory B cells (74, 75) beyond the initial 309 vaccination phase.

310

This study is also the first to describe the course of infection and histopathology of a SARSCoV-2 P.1 infection in a NHP model. Consistent with what was described in WA1/2020 infected

313 cynomolgus macagues (51, 52), macagues challenged with the P.1 isolate did not develop elevated temperature or decreased weight, nor did they show observable signs of respiratory 314 315 disease. Viral load in the upper and lower respiratory tract peaked early during infection two 316 days after challenge and gradually decreased to undetectable sgRNA levels by day 14 in 317 unprotected animals, similar to viral kinetics described in another study using the same 318 inoculating dose $(1 \times 10^6 \text{ TCID}_{50})$, but with an early pandemic isolate (52). Although P.1 is 319 estimated to be upwards of 2.4-fold more transmissible (76), it is unclear whether the higher 320 viral load observed during peak P.1 replication in this study compared to time-matched reports 321 for a non-variant isolate is a characteristic of P.1 replication or a discrepancy between readouts 322 from different RT-PCR procedures. However, we show that immunization with both 5 and 25 μ g 323 doses of S effectively reduces viral load in both upper and lower respiratory tracts implying a 324 lower likelihood of viral transmission even in mildly symptomatic macaques. Histopathological 325 examination of lung and bronchial sections in unprotected macagues confirmed mild to 326 moderate disease, which was abrogated in immunized individuals. COVID-19 like disease 327 observed in this model consisted of increased intraalveolar and interstitial infiltration, as well as 328 giant syncytial cells, similar to previous observations (51, 77, 78). While both high and low 329 doses of S antigen generate statistically negligible differences in humoral responses during 330 peak immunity, it is clear from our findings that higher antigen doses, at least with CoVaccine 331 HT[™] produces a more durable and protective response at this later timepoint.

332

Late P.1 challenge induced an anamnestic response in all immunized macaques that restored
S-specific antibody titers to peak serum concentrations seen shortly after the final boost, and
significantly enhanced vaccine-matched and VOC-neutralizing antibody titers. In most
macaques receiving 25 µg S, the immune recall occurred with little to no disease. The durability
and cross-variant neutralizing nature of immune responses generated by mRNA and viral
vectors has been documented up to six months or later (*79-82*). However, breakthrough

339 infections, particularly with variant strains, have been reported even in the presence of 340 detectable and high-level NtAbs (54, 56, 59, 83). Anamnestic responses in fully immunized 341 NHPs have been characterized previously with homologous and B.1.351 variant challenge and 342 agree with our observations that viral challenge boosts functional antibody levels. Our analysis 343 shows that viral challenge with a heterologous strain not only rapidly boosts homologous and 344 challenge strain specific NtAbs, but also NtAbs against the unencountered variants, B.1.351 and 345 B.1.617, underpinning the broad-spectrum potential of subunit protein vaccines. In contrast, 346 control macagues, receiving no S antigen, developed moderate levels of P.1 NtAbs that are only 347 cross-neutralizing with B.1.351, and barely neutralize WA1/2020 and B.1.617, suggesting that 348 natural infection with one strain of SARS-CoV-2 may confer only limited protection against other 349 VOCs. Of course, further affinity maturation could not be observed here as the study ended 14 350 days after challenge.

351

352 The small size of our treatment groups (n=3) is a limitation of our study and may therefore not 353 provide enough statistical power to strongly correlate antibody concentrations to viral load or 354 histopathology score. Furthermore, we cannot directly compare the outcome of our study to 355 other studies that evaluate vaccine efficacy soon after the final booster when immunity is 356 greatest, as vaccine efficacy is known to decline over time. Likewise, we could not benchmark 357 the efficacy of our vaccine formulation when the immunity is greatest at week 5, however, our 358 late challenge scheme accurately reflects the current urgent need for decisions about timing and 359 candidates for possible booster vaccinations during the pandemic situation as it provides useful 360 information regarding the real-life durability and breadth of vaccine protection. It furthermore 361 may shed light into the utility of protein vaccines to serve as prime or boost in combination with 362 other vaccines.

363

364	In conclusion, we show that a two-dose regimen of a prefusion-stabilized trimeric S subunit
365	protein vaccine formulated with lyophilizable CoVaccine HT [™] adjuvant reduces viral burden and
366	high antigen doses can confer durable cross-variant immunity. Future efforts will therefore focus
367	on developing a thermostabilized vaccine formulation in a single-vial presentation, potentially
368	enabling facile worldwide distribution.
369	
370	MATERIALS AND METHODS
371	Ethical Statement
372	The investigators adhered fully to the "Guide for the Care and Use of Laboratory Animals" by
373	the Committee on Care of Laboratory Animal Resources Commission on Life Sciences, National
374	Research Council. Cynomolgus macaques (Macaca fascicularis) were housed at BIOQUAL Inc.
375	(Rockville, MD). All macaque experiments were reviewed and approved by BIOQUAL's Animal
376	Care and Use Committee. BIOQUAL Inc. is accredited by the American Association for
377	Accreditation of Laboratory Animal Care (AAALAC).
378	
379	Study Design Cynomolgus macaque studies were performed using three adjuvanted vaccine
380	formulations and an adjuvanted control using unrelated antigens: A) 5 μ g of SARS-CoV-2 S
381	protein with 10 mg lyophilized CoVaccine HT™, B) 25 µg S protein with 10 mg liquid CoVaccine
382	HT™, C) 25 μg S protein with 10 mg lyophilized CoVaccine HT™, and D) 25 μg of Ebola Virus
383	(EBOV) glycoprotein with 10 mg CoVaccine HT™, co-lyophilized in one vial. CoVaccine HT™
384	(BTG International Ltd, London, United Kingdom) was lyophilized as previously described (71)
385	and reconstituted with sterile PBS before mixing with the antigen. Each group consisted of both
386	male and female cynomolgus macaques ($n = 3$ for each formulation) weighing between 2.8 and
387	8.2 kg. Cynomolgus macaques of groups A-C were immunized intramuscularly (IM) in both

- deltoids (split dose) at weeks 0 and 3, control animals were immunized only once at week 0.
- 389 Pre-challenge sera were collected at weeks 0, 3, 5, 6, 15. Challenge at week 15 with SARS-

390 CoV-2 P.1 variant, hCoV-19/Japan/TY7-501/2021, TY7-501 (BIOQUAL-generated stock [lot no. 391 031921-1215] in Calu-3 cells from seed stock no.TY7-501 was performed with an inoculum 392 dose of 5x10⁵ TCID₅₀/mL administered to each animal in volumes of 1 mL by intratracheal and 393 intranasal injection at each site. Nasal and oral swabs (NS and OS respectively) were collected 394 on days 2, 4, 7, 10 and 14 after challenge. Bronchoalveolar lavage (BAL) samples were 395 collected on days 2, 4, 7 and 10 after challenge. Necropsies and lung tissue collection were 396 performed at the endpoint of the study at day 14 post-challenge. Samples were immediately 397 processed and subsequently stored at -80°C prior to analysis. S-specific serum IgG 398 concentrations were determined using a microsphere immunoassay (MIA; see below). NtAbs 399 were measured using wild-type and rVSV-SARS-CoV-2 S PRNTs (see below). Viral load in the 400 NS, OS and BAL was measured using plaque assays and guantification of genomic and 401 subgenomic N transcript RNA (see below). Lung and bronchi tissues were processed, and H&E 402 stained to delineate histopathological signs of disease in each animal. 403 404 Recombinant protein expression and purification 405 Plasmids were generated to express the pre-fusion, protease-resistant, trimeric transmembrane 406 (TM)-deleted spike (S) glycoprotein from SARS-CoV-2 as described previously (53). 407 Modifications to the gene include the removal of the furin and S2' cleavage site, the addition of 2 408 prolines between the heptad repeat 1 and central helix region, and a foldon trimerization 409 domain. A stably transformed cell line was created by hygromycin B selection at 300 µg/mL.

410 The cell line was scaled up to 1L using a WAVE bioreactor (Cytiva, Marlborough, MA) and

411 induced with 200 μM CuSO₄.

412

413 Recombinant S protein was purified from clarified cell culture supernatants by immunoaffinity

414 chromatography (IAC) using the SARS-CoV-2 cross-reactive mAb CR3022 (provided by Mapp

Biopharmaceutical) coupled to NHS-activated Sepharose at a concentration of 10 mg/mL. The

416 antigen was eluted with a glycine buffer (pH 2) in tandem into a HiPrep 26/10 desalting column 417 (Cytiva, Marlborough, MA) equilibrated with PBS. The oligomeric content was evaluated by size-418 exclusion chromatography using a HiLoad 16/600 column (GE Healthcare, Piscataway, NJ) 419 equilibrated with PBS. The S protein eluted as a single peak and the final product migrated as 420 two bands, corresponding to the monomer and trimer on SDS-PAGE, under denaturing 421 conditions, and was reactive to CR3022 mAb on a western blot. Antigens were sterile filtered 422 with a 0.22 µm syringe filter (Cytiva, Marlborough, MA) and stored at -80°C until use. 423 424 Analysis of antibodies by multiplex microsphere immunoassay (MIA) 425 The IgG antibody titers in sera were measured using a multiplex microsphere-based 426 immunoassay as described previously (53, 84-86). Spectrally distinct, magnetic MagPlex® 427 microspheres (Luminex Corporation, Austin, TX) were coupled to purified S, RBD or bovine 428 serum albumin (BSA). A mixture of the antigen-coupled beads was incubated with sera diluted 429 with PBS+ 1% BSA and 0.02% Tween 20 (PBS-BT) at 1:5,000 or 1:10,000 in black-sided 96-430 well plates for 3 hours at 37°C with agitation. Bound IgG was detected using 1µg/mL red 431 phycoerythrin (R-PE)-conjugated goat anti-human IgG antibodies (Jackson ImmunoResearch, 432 Inc., West Grove, PA) and resuspended in MAGPIX® drive fluid before being analyzed on a 433 MAGPIX® Instrument (Luminex Corporation, Austin, TX).

434

To determine S-specific IgG concentrations in the sera, the median fluorescence intensity (MFI)
readouts of each sample was interpolated against a standard curve generated using purified
human IgG at concentrations in the range of 7.44 to 1000 ng/mL. To produce the antibody
standard, IgG was purified from pooled sera of COVID-19 vaccinated human volunteers using
protein A affinity chromatography, followed by immunoaffinity chromatography (IAC) using NHSSepharose (Cytiva, Marlborough, MA) coupled with recombinant S to select for S-specific IgG.
Purity was assessed using SDS-PAGE and antibody concentration was quantified using UV₂₈₀

442 absorbance. The resulting MFI values were plotted against the Log₁₀-transformed

443 concentrations and fitted using a sigmoidal dose-response, variable slope model (GraphPad

444 Prism, San Diego, CA). The resulting curves yielded r^2 values > 0.99 with a well-defined top and

bottom and the linear range of the curve. The experimental S-specific IgG concentrations in

446 experimental samples were determined by interpolation on the standard curves, multiplied by

the dilution factors and plotted as antibody concentrations (ng/mL).

448

449 Recombinant vesicular stomatitis virus (rVSV) neutralization assay

450 Replication-competent rVSV expressing SARS-CoV-2 S protein (Wuhan-Hu-1) was generated

451 as described previously (87) and the virus stocks were amplified in Vero E6 cells. For the

452 plaque reduction neutralization test (PRNT), individual NHP serum samples were heat-

453 inactivated at 56°C for 30 minutes. Six 3-fold serial dilutions of serum samples, starting at 1:40

dilution, were prepared and incubated with 100 plaque-forming units (PFU) of rVSV-SARS-CoV-

455 2-S at 37°C for 1 hour. Antibody-virus complexes were added to Vero cell monolayers in 6-well

456 plates and incubated at 37°C for another hour followed by addition of overlay media mixed with

457 1% agarose. 72 hours later, cells were fixed and stained with a solution containing 1%

458 formaldehyde, 1% methanol, and 0.05% crystal violet overnight for plaque enumeration.

459 Neutralization titers (PRNT₅₀) were generated using a variable slope, nonlinear regression, with

460 upper and lower constraints (100% and 0% neutralization, respectively), using Prism 9

461 (GraphPad Software, San Diego, CA).

462

463 TCID₅₀ and Wild-type SARS-CoV-2 virus PRNT₅₀ assay

464 TCID₅₀ and PRNT₅₀ assays were performed in a biosafety level 3 facility at BIOQUAL, Inc.

465 (Rockville, MD). The TCID₅₀ assay was conducted by addition of 10-fold graded dilutions of

samples to Vero TMPRSS2 cell monolayers. Serial dilutions were performed in the cell culture

467 wells in quadruplicates. Positive (virus stock of known infectious titer in the assay) and negative

468 (medium only) control wells were included in each assay set-up. The plates were incubated at 469 37°C, 5.0% CO2 for 4 days. The cell monolayers were visually inspected for CPE, i.e. complete 470 destruction of the monolayer with cellular agglutination. The TCID₅₀ value was calculated using 471 the Read-Muench formula (88). For samples which had less than 3 CPE positive wells, the 472 TCID₅₀ could not be calculated using the Reed-Muench formula, and these samples were 473 assigned a titer of below the limit of detection (i.e., <2.7 log10 TCID₅₀/mL). For optimal assay 474 performance, the TCID50 value of the positive control should test within 2-fold of the expected 475 value.

476

477 To measure neutralization, sera from each NHP were diluted to 1:10 followed by a 3 fold- serial 478 dilution. Diluted samples were then incubated with 30 plague-forming units of wild-type SARS-479 CoV-2 USA-WA1/2020 (BEI-NR-52281), B.1.351 (BEI NR-55282), or P.1. (BEI NR-54982) 480 variants, in an equal volume of culture medium for 1 hour at 37°C. The serum-virus mixtures 481 were added to a monolayer of confluent Vero E6 cells and incubated for one hour at 37°C in 5% 482 CO₂. Each well was then overlaid with culture medium containing 0.5% methylcellulose and 483 incubated for 3 days at 37°C in 5% CO₂. The plates were then fixed with methanol at -20°C for 484 30 minutes and stained with 0.2% crystal violet for 30 min at room temperature. PRNT₅₀ titers 485 were calculated using a variable slope, nonlinear regression, with upper and lower constraints 486 (100% and 0% neutralization, respectively), on Prism 9 (Graphpad Software, San Diego, CA).

488 Histopathology

487

NHP Lung tissue specimens from each lung lobe and bronchi were harvested at time of
necropsy and preserved in 10% formalin before processing and parafilm embedded, fixed and
stained with hematoxylin & eosin. For each NHP, one section from the bronchi and one section
from each lobe on the right and left lung were selected for scoring, for a total of six sections.
Pathologic findings on each slide were scored on a scale of 0 – 2 for intraalveolar edema, the

amount of BALT (bronchiolar-associated lymphoid tissue), and the presence of Interstitial
inflammation; and 0 – 3 for perivascular inflammatory infiltrates (cuffing), and intraalveolar
inflammation. (See Table S1.) Scores for the six slides from each macaque were tabulated, and
a cumulative average score was calculated for each NHP using a total of 30 scores as
replicates.

499

500 SARS-CoV-2 Viral Genomic and subgenomic RNA quantitative RT-PCR

501 The presence of viral RNA and viral replication in the BAL, NS, and OS after SARS-CoV-2 P.1 502 strain challenge was determined by quantitative RT-PCR. RNA was isolated from 200 µL 503 sample using the QIAamp MinElute Virus spin kit (Qiagen, Frederick, MD). For the gRT-PCR 504 assay, viral RNA was first isolated from BAL, NS and OS using the Qiagen MinElute virus spin 505 kit. To generate a control for the amplification reaction, RNA was isolated from the applicable 506 virus stock using the same procedure. The number of copies for the control were calculated 507 using known RNA weights per mol. A master mix was prepared containing Tag-polymerase, 508 obtained from the TagMan RT-PCR kit (Bioline cat# BIO-78005), RT, RNAse inhibitor, a primer 509 pair at 2 µM concentration (2019-nCoV N1-F: 5'-GAC CCC AAA ATC AGC GAA AT-3' and 510 2019-nCoV N1-R: 5'-TCT GGT TAC TGC CAG TTG AAT CTG-3') and probe (2019-nCoV N1-511 P: 5'-FAM-ACC CCG CAT TAC GTT TGG TGG ACC-BHQ1-3') at a concentration of 2 µM. For 512 the reactions, 45 μ L of the master mix and 5 μ L of the sample RNA were added to the wells of a 513 96-well plate. All samples were tested in triplicate. Control RNA was prepared to contain 10⁶ to 514 10^7 copies per 3 µL. Eight (8) 10-fold serial dilutions of control RNA were prepared and 515 produced a standard curve with a range of 1 to 10⁷ copies/reaction. For amplification, the plate 516 was placed in an Applied Biosystems 7500 Sequence detector and amplified using the following 517 program: 48°C for 30 minutes, 95°C for 10 minutes followed by 40 cycles of 95°C for 15 518 seconds, and 1 minute at 55°C. Duplicate samples of each dilution were prepared as described 519 above. If the copy number exceeded the upper detection limit, the sample was diluted as

needed. The number of copies of RNA per mL was calculated by interpolation from the standard
curve and multiplying by the reciprocal of 0.2 mL extraction volume. This gave a practical range
of 50 to 5 x 10⁸ RNA copies per mL for BAL samples and for nasal and oral swabs the viral
loads were given per swab.

524

525 The RT-PCR assay for the sgRNA utilizes primers and a probe specifically designed to amplify 526 and bind to a region of the N gene mRNA from the Coronavirus, which is not packaged into the 527 virion. The signal was compared to a known standard curve of plasmid containing the sequence 528 of part of the messenger RNA and calculated to give copies per ml. The control DNA was 529 prepared to contain 10⁷ copies. Seven 10-fold serial dilutions of control RNA were prepared 530 using Buffer AVE and generated a standard curve with a range of 1 to 10⁶ copies/reaction. 531 Duplicate samples of each dilution were prepared as described above with the primer pair (SG-532 N-F: CGATCTCTTGTAGATCTGTTCTC and SG-N-R: GGTGAACCAAGACG CAGTAT) and 533 probe (FAM- TAACCAGAATGGAGAACGCAGTGGG -BHQ). If the copy number exceeded the 534 upper detection limit, the sample was diluted as needed. For amplification, the plate was placed 535 in an Applied Biosystems 7500 Sequence detector and amplified using the following program: 536 48°C for 30 minutes, 95°C for 10 minutes followed by 40 cycles of 95°C for 15 seconds, and 1 537 minute at 55°C. The number of copies of RNA per ml was calculated by interpolation from the 538 standard curve and multiplying by the reciprocal of 0.2 ml extraction volume. This gave a 539 practical range of 50 to 5×10^7 RNA copies per mL for harvested samples.

540

541 Statistical analysis

542 Statistically significant differences between the geometric mean of IgG concentrations or MFI in 543 groups given different vaccine formulations at each timepoint were determined using a two-way 544 ANOVA followed by Dunnett's Multiple Comparison. Comparisons of PRNT₅₀ values between 545 vaccine formulations was done using a Kruskal-Wallis Test followed by a Dunn's multiple

546	comparisons test. Correlations between IgG or PRNT titers to viral load in the BAL and NS was		
547	examined using the non-parametric Spearman's correlation test. Differences between		
548	histopathological scores in different treatment groups was calculated using a one-way ANOVA		
549	followed by Dunnett's multiple comparison test with the cumulative scores of all slides per		
550	macaque as replicates. All statistical analysis was completed using Graphpad Prism 9 software		
551	(San I	Diego, CA).	
552			
553	Suppl	ementary Materials	
554	Fig. S1. Serum Wuhan-Hu-1 RBD-Specific IgG Kinetics		
555	Fig S2. Pre-and Post-Challenge rVSV-SARS-CoV2-S Neutralizing Antibody Response		
556	Fig S3. Individual TCID ₅₀ Viral load from the BAL and NS		
557	Fig S4. Individual Genomic Viral RNA Load		
558	Fig S5. Individual Subgenomic N transcript RNA Load		
559	Table S1 Detailed Histopathology Scoring System		
560			
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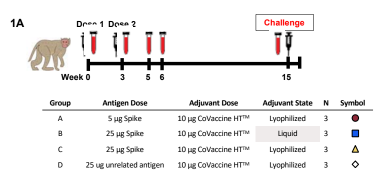
780 Author Contribution:

- 781 Conceptualization: AT, MML, HA, OD, ATL
- 782 Methodology: AT, TSW, KT, LP, HA, CYL, ATL
- 783 Investigation: AT, TSW, KT, LP, JG, ND, AC, BN, ZF, AVR, JYO
- 784 Formal Analysis: AT, MML, KT, ATL
- 785 Visualization: AT, KT
- 786 Funding acquisition: OD, ATL
- 787 Project administration: HA, OD, ATL
- 788 Supervision: HA, OD, ATL
- 789 Writing original draft: AT, KT, ATL
- 790 Writing review & editing: AT, TSW, MML, KT, HA, OD, ATL
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792 Competing Interests:

- ATL and OD are named inventors on a patent application covering a recombinant subunit
- vaccine for SARS-CoV-2. LP, JG, ND, AC, BN, ZF, AVR, JYO and HA are current employees of
- 795 BIOQUAL, Inc. OD is a current employee of Soligenix Inc. All other authors declare no
- competing interests.
- 797
- 798 Data and material availability: All data are available in the main text or the supplementary
- 799 materials

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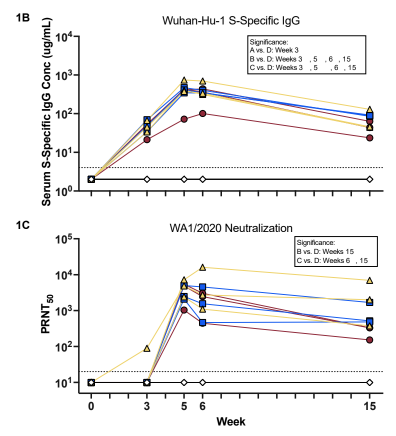


Figure 1. Vaccine Scheme, IgG and Neutralizing Antibody Kinetics (A) 12 cynomolgus macaques were separated into 4 groups and given either 5 or 25 μ g S protein formulated with either liquid or reconstituted, lyophilized CoVaccine HTTM adjuvant. Two doses were administered IM 3 weeks apart and sera was collected at indicated time points. Macaques were challenged IN and IT with a total of 1x10⁶ TCID₅₀ of the SARS-CoV-2 P.1 strain. (B) Serum Wuhan-Hu-1 S-specific IgG kinetics measured using a MIA with purified, human S-specific IgG standards to estimate serum concentration. Dashed line indicates the limit of quantification (LOQ, $\leq 4 \mu$ g/mL). Individual values falling below the LOQ were set to $\frac{1}{2}$ LOQ. Significance was calculated using a one-way ANOVA followed by a Dunnett's Multiple Comparison to Group D at each time point. (C) WA1/2020 neutralizing antibody kinetics were measured using a sigmoidal dose response curve. Dashed line indicates the limit of detection (LOD, $\leq 1:20$). Individual values falling below the LOD were set to $\frac{1}{2}$ LOD were set to $\frac{1}{2}$ LOD. Significance was calculated using a Kruskal-Wallis Test followed by a Dunn's Multiple Comparison. (*p ≤ 0.05 , **p ≤ 0.01).

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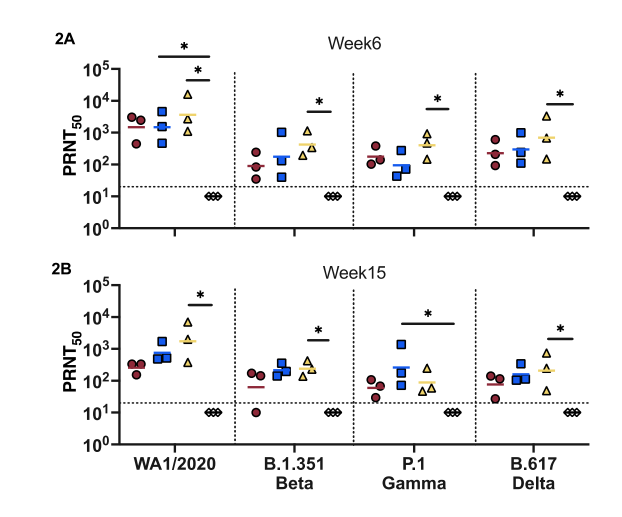


Figure 2. Cross-Neutralizing Antibody Titers Neutralizing antibody titers at (A) week 6 and (B) week 15 were measured using a WT SARS-CoV-2 PRNT assay with the WA1/2020, B.1.351 (Beta), P.1 (Gamma), B.1.617 (Delta) VOC. Curve-fitted PRNT₅₀ titers were calculated using a sigmoidal dose response curve. Dashed line indicates the LOD (\leq 1:20). Individual values falling below the LOD were set to ½ LOD. Significance was calculated using a Kruskal-Wallis Test followed by a Dunn's Multiple Comparison to Group D at each time point. Horizontal bars mark the GMT of each group (*p \leq 0.05).

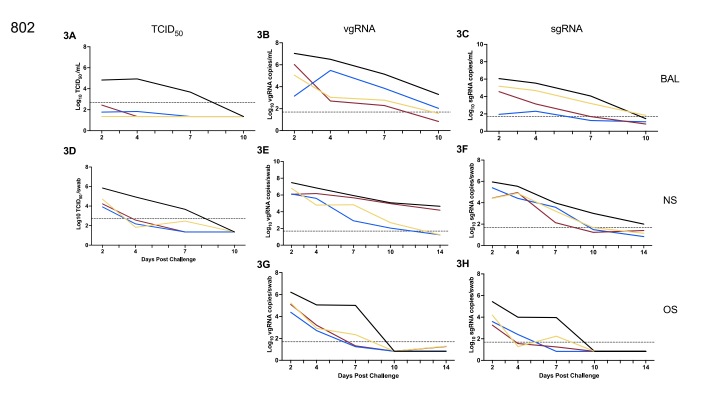


Figure 3. Viral Load Kinetics after P.1 Challenge GMT of the (A & D) $TCID_{50}$ titers; (B, E, G) viral genomic; and (C, F, H) subgenomic N RNA copies from the (A, B, C) bronchoalveolar lavage (BAL); (D, E, F) nasal swab (NS); and (G, H) oral swab (OS) collected at the time points indicated. Dashed line indicates the assay-specific LOD. Individual values falling below the LOD were set to $\frac{1}{2}$ LOD.

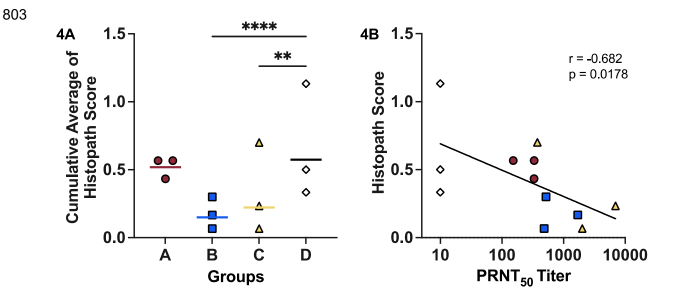


Figure 4. Histopathology Score and Correlation with Neutralizing Antibodies (A) Cumulative average histopathology score was determined by averaging the scores of six sections cut from each lobe and the bronchi. The presence and severity of edema, intra-alveolar inflammation, perivascular cuffing, increased BALT and interstitial inflammation was determined and assigned scores ranging from 0-3. Significant differences in lung histopathology were calculated using a one-way ANOVA followed by a Dunnett's multiple comparison to Group D with the scores from each section (30 scores per macaque) as replicates. Horizontal bars mark the GMT of each group. (** $p \le 0.01$, **** $p \le 0.0001$) (B) Correlation between histopathology scores and WA1/2020-specific PRNT₅₀ titers were determine using a Spearman's correlation test. bioRxiv preprint doi: https://doi.org/10.1101/2021.09.24.461759; this version posted September 25, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.



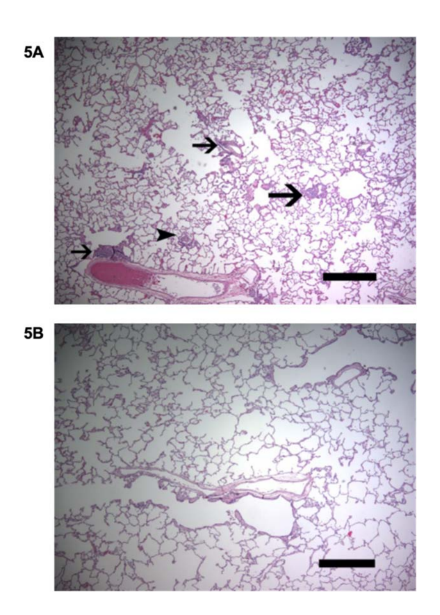


Figure 5 Histopathology observed in Hematoxylin and Eosin-Stained Lung Sections of Control Cynomolgus Macaques Compared to Immunized Macaques after P.1 challenge (A) Representative low power view of a lung section showing distribution of lymphocytic perivascular cuffing (arrows) and clusters of intraalveolar macrophages (arrowhead) in an unprotected macaque. (B) Representative low power view of a lung section from an immunized macaque which did not exhibit pathologic changes. Scale Bar = 500 microns.

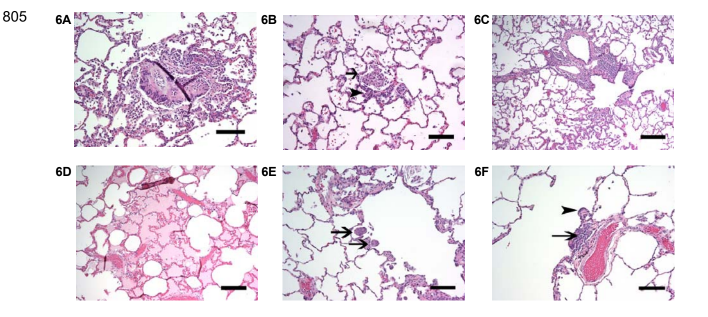


Figure 6 Histopathology Observed in Control Cynomolgus Macaques Compared to that Observed in Immunized Macaques after P.1 Challenge (A-D) unprotected macaques. (A) A syncytium of intraalveolar multinucleated giant cells is seen surrounded by an acute and eosinophilic inflammatory infiltrate. (B) Increased intraalveolar macrophages and an intraalveolar multinucleated giant cell accompanied by interstitial lymphocytic inflammation and thickening. (C) Complete lymphocytic perivascular cuffing with increased intra-alveolar macrophages. (D) Proteinaceous edema fluid in alveolar spaces. (E-F) Protected macaques. (E) Intraalveolar multinucleated giant cells (arrows) without increased intraalveolar macrophages, lymphocytic vascular cuffing or interstitial inflammation. (F) Partial lymphocytic perivascular cuff (arrow) with adjacent alveolar epithelial hyperplasia (arrowhead) without alveolar septal thickening. Scale bar = 10 microns.

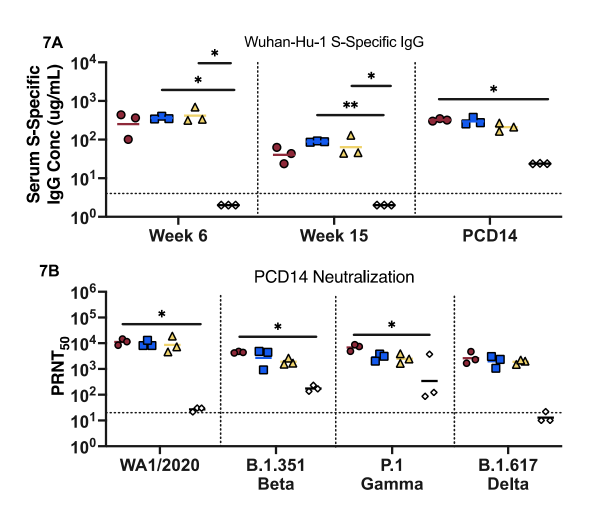


Figure 7. Post-Challenge Anamnestic Response (A) Vaccine-matched S-specific IgG conc. from sera collected at week 6, week 15 and at 14-days post-challenge, measured by MIA. IgG concentrations were estimated using a human S-specific IgG standard as described previously. Horizonal bars mark the GMT for each group. Dashed line indicates the LOQ ($\leq 4 \mu g/mL$) Individual values falling below the LOQ were set to ½ LOQ (B) Neutralizing antibodies 14 days after P.1 challenge were measured using a WT SARS-CoV-2 PRNT assay with the WA1/2020, B.1.351 (Beta), P.1 (Gamma), B.1.617 (Delta) VOC. Curve-fitted PRNT₅₀ titers were calculated using a sigmoidal dose response curve. Dashed line indicates the LOD ($\leq 1:20$). Individual values falling below the LOD were set to ½ LOD. Significance was calculated using a Kruskal-Wallis Test followed by a Dunn's Multiple Comparison to Group D at each time point. (*p ≤ 0.05 , **p ≤ 0.01).

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