| 1 | Assessment of serological assays for identifying high titer convalescent plasma |
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29 ABSTRACT

30 The COVID-19 pandemic has been accompanied by the largest mobilization of therapeutic convalescent plasma (CCP) in over a century. Initial identification of high titer units was based 31 32 on dose-response data using the Ortho VITROS IgG assay. The proliferation of SARS-CoV-2 33 serological assays and non-uniform application has led to uncertainty about their 34 interrelationships. The purpose of this study was to establish correlations and analogous cutoffs 35 between commercially available serological tests (Ortho, Abbott, Roche), a spike ELISA, and a virus neutralization assay using convalescent plasma from a cohort of 79 donors from April 36 37 2020. Relationships relative to FDA-approved cutoffs under the CCP EUA were identified by 38 linear regression and receiver operator characteristic curves. Relative to the Ortho VITROS assay, the r^2 of the Abbott, Roche, the anti-Spike ELISA and the neutralizing assay were 0.58, 39 40 0.5, 0.82, and 0.44, respectively. The best correlative index for establishing high-titer units was 41 3.82 S/C for the Abbott, 10.89 COI for the Roche, 1:1,202 for the anti-Spike ELISA, and 1:200 42 by the neutralization assay. The overall agreement using derived cutoffs compared to the CCP 43 EUA Ortho VITROS cutoff of 9.5 was 92.4% for Abbott, 84.8% for Roche, 87.3% for the anti-S 44 ELISA and 78.5% for the neutralization assay. Assays based on antibodies against the 45 nucleoprotein (Roche, Abbott) and neutralizing antibody tests were positively associated with the Ortho assay, although their ability to distinguish FDA high-titer specimens was imperfect. The 46 47 resulting relationships help reconcile results from the large body of serological data generated 48 during the COVID-19 pandemic.

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52 INTRODUCTION

53 COVID-19 convalescent plasma (CCP) has been one of the primary therapies deployed in the 54 COVID-19 pandemic. In this current iteration of a classic therapy, serological assays to quantify 55 antibodies to the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) spike (S) 56 protein play a critical role in characterizing human immune responses and identifying CCP 57 donors. Commercial SARS-CoV-2 serological assays have accordingly emerged at a rapid pace. 58 Within the first year of the pandemic, more serological assays were available for SARS-CoV-2 59 than for any other infectious disease, with over 65 emergency use authorizations (EUA) granted 60 for serological testing alone (1). The CDC and Infectious Diseases Society of America (IDSA) 61 have both defined relatively narrow and limited clinical applications for SARS-CoV-2 62 serological to include CCP donors identification, infection diagnosis in patients more than 14 63 days from symptom onset, and establishing seroprevalence in populations (2–4). However, the 64 clinical utility of these assays has been questioned (5, 6), in part, due to the challenge of 65 reconciling results from serological assays with clinical outcomes (7–9) and poor agreement 66 between commercial serological assays and virus neutralization assays (10-12).

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Identification of CCP with antibody content sufficient for therapeutic use of CCP has emerged as a key quantitative application for SARS-CoV-2 serological assays (2, 5). Anti-S IgG responses in particular were identified early as key correlates of SARS-CoV-2 immunity. At the pandemic's outset, the absence of an FDA-approved serological assay was a major obstacle to identifying CCP units with sufficient antibody content. A highly sensitive and specific laboratory developed S-based ELISA was quickly developed (13) and used to identify CCP donors with antibodies to SARS-CoV-2 following RT-PCR confirmed infection (14). The initial FDA

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75 recommendation was to use a minimum titer of 1:160, with an ideal titer \geq 1:320, as a criterion 76 for CCP donation (14). A subsequent study demonstrated that high-titer CCP, which was defined 77 as a signal of \geq 18.45 on the Ortho-Clinical Diagnostics VITROS IgG assay, was associated with 78 lower risk of mortality than those receiving low-titer units in a large retrospective analysis of 79 patients treated through an FDA expanded access protocol (15). A subsequent analysis of this 80 cohort through the Biomedical Advanced Research and Development Authority (BARDA) found 81 that patients receiving CCP with a neutralizing antibody titer > 1:250 experienced lower 82 mortality than those receiving units with titers < 1:250 (16, 17). Neutralizing antibody assays, 83 however, are highly laborious and require biosafety level 3 facilities if using live SARS-CoV-2, 84 limiting their use primarily to research laboratories. As a result, neutralizing assays were 85 correlated with the Ortho Clinical IgG assay, with a minimum signal of 12.0 distinguishing units 86 with high neutralizing titers (18).

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88 In February 2021, the FDA reissued a letter of authorization for CCP with several revisions to 89 the previous EUA (19). Importantly, this included a decision to release only high-titer CCP units 90 for patient use. Cutoffs were provided so that multiple serological assays could be used to define 91 high-titer CCP and previously established titers approved by the FDA were modified. The titer to 92 establish high-titer units with the Ortho Clinical assay was lowered from 12.0 to \geq 9.5 S/C, and 93 the original anti-S ELISA threshold was raised from 1:320 to \geq 1:2,880 in an ELISA performed 94 at Mt. Sinai Hospital. The revised EUA also established cutoffs for distinguishing high-titer units 95 using seven other commercial serological assays. For example, the Abbott SARS-CoV-2 IgG 96 assay and the Roche Elecsys anti-SARS-CoV-2 assay were approved for qualifying high-titer 97 units with results \geq 4.5 signal-to-cutoff (S/C) and \geq 109 cutoff index (COI), respectively.

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99 Little published literature is available to correlate neutralizing antibody titers, commercial 100 serological assays, and anti-S ELISAs. Several studies have assessed the positive percent 101 agreement and negative percent agreement (PPA and NPA) between assays (10, 11, 20). 102 However, the signal from other commercial, serological assays that best correlate to anti-S 103 ELISA titers of 1:320, neutralizing titers of 1:250, and the Ortho Clinical S/C of 12.0, have not 104 been determined. The purpose of our study was to establish correlations and analogous cutoffs between widely used commercial serological assays, anti-S ELISA, and neutralizing assays with 105 106 authentic SARS-CoV-2. The resulting relationships will help reconcile results from the large 107 body of serological studies and CCP trials results that continue to emerge during the COVID-19 108 pandemic.

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110 MATERIALS AND METHODS

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112 Human subjects

113 This study was approved by the Washington University Institutional Review Board. Serum 114 specimens were drawn on patients with RT-PCR-confirmed SARS-CoV-2 infection at least 14 115 days after infection and prior to donation of convalescent plasma. Patient reported demographic 116 information including age, gender, race, comorbidities, and duration of symptoms were collected 117 by survey on each patient. After collection, specimens were immediately frozen in 100 μ L 118 aliquots and stored at -80°C until further analysis.

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120 Assays

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121 Specimens were thawed at room temperature and analyzed within 3 days. Three commercial 122 serological assays and an anti-S ELISA granted EUA at Mt. Sinai Hospital, but used on a 123 research basis for this study, were used to directly measure antibody levels in serum specimens. 124 These assays detected antibodies to SARS-CoV-2 S or nucleocapsid proteins. The anti-S ELISA 125 was performed as previously described (13). In short, plasma specimens were diluted to 1:30 in 126 PBS, then serially diluted to 1:65,610 in a 96-well plate (Corning). Wells were washed, 127 incubated with a secondary anti-human IgG, followed by another wash step. Wells were then 128 incubated with o-Phenylenediamine dihydrochloride (Sigma-Aldrich) followed by a stop 129 solution (3M hydrochloric acid). The optical density was then measured at 490 nm and the cutoff 130 for a positive result was determined as an optical density that was three standard deviations 131 above the mean signal from a negative control specimen run with each plate. This signal was 132 extrapolated from the generated curves to quantify the titer (21).

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134 An authentic SARS-CoV-2 neutralization assay was used to measure neutralizing antibody titers. 135 Focus reduction neutralization assays were performed as previously described (10). SARS-CoV-136 2 strain n-CoV/USA_WA1/2020 was obtained from the Centers for Disease Control. Virus was 137 propagated in Vero E6 cells in Dulbecco's Modified Eagle Medium (DMEM, Corning) that was 138 supplemented with 10% FBS, glucose, L-glutamine, and sodium pyruvate. Patient sera were diluted and incubated with 1×10^2 focus forming units (FFU) of SARS-CoV-2 for 1 h at 37° C. 139 140 The plasma/virus complex was then added to Vero E6 monolayers at 37°C for 1 h. After 141 overlaying with methylcellulose, cells were harvested at 30 h, methylcellulose was removed, and 142 fixed with 4% paraformaldehyde (PFA) for 20 min. Plates were washed and incubated with 1 143 µg/mL anti-S antibody (CR3022) and HRP conjugated goat anti-human IgG. Infected cells were

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144 visualized with TrueBlue peroxidase substrate (KPL) and quantified using an ImmunoSpot 145 microanalyzer (Cellular Technologies). A minimum of eight dilutions was performed for each 146 specimen, a standard curve generated, and the 1/log10 plasma dilution (EC50) determined as the 147 dilution at which 50% of the cells were infected.

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149 All specimens were analyzed on three commercially available serological assays. The Ortho 150 Clinical VITROS SARS-CoV-2 IgG assay was performed on an Ortho Clinical VITROS 5600 151 Immunodiagnostic System and targets antibodies to the S protein. The Abbott SARS-CoV-2 IgG 152 assay was performed on an Abbott Architect i2000 and detects antibodies to the nucleocapsid 153 protein. The Roche Elecsys anti-SARS-CoV-2 assay was performed on a Cobas e601 and 154 identifies antibodies to the nucleocapsid protein. All commercial assays have FDA EUA as 155 qualitative methods and were performed and interpreted according to the manufacturer's 156 instructions. The positive cutoffs for each assay are 1.0 (S/C), 1.4 (S/C), and 1.0 (COI) for the 157 Ortho Clinical, the Abbott, and the Roche assays, respectively. All three assays report a numeric 158 signal to cut-off that is the amount of signal generated by the sample for each assay relative to 159 the signal from a single calibrator.

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161 Statistical analysis

Association between assays were compared with least squares regression to calculate intercept, slope and r^2 . Ideal cutoffs from linear regression were established by interpolating relative to each cutoff. Receiver operator curves (ROC) were also plotted to assess the ideal cutoffs using Youden Index to establish cutoff with maximum positive and negative percent agreement. Final

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166 cutoffs for distinguishing high- and low-titers units by each assay were established by averaging167 across all cutoffs.

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169 **RESULTS**

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171 COVID-19 convalescent plasma donors

Serum specimens were obtained from 79 adults at Washington University/Barnes-Jewish Hospital Medical Center in St. Louis, Missouri, U.S.A., with a history of positive SARS-CoV-2 RT-PCR testing who expressed interest in donating CCP between 4/6/2020-4/29/2020. The median age was 49 (range 20-69) (**Table 1**). 55.7% of patients were female, 91.1% were white. The most common comorbidity was asthma. Only 2 patients (2.5%) were hospitalized, and the median duration of symptoms was 12 days (range; 1-31). The median time from symptom onset to positive RT-PCR result was 4 days (range; 0-20).

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180 Anti-S ELISA IgG titers in this cohort spanned four orders of magnitude (1:21 - 1:17,278) with 181 a median titer of 1:2,315 (Fig 1A). A broad range of responses also was evident among 182 commercial serological assays. These results are consistent with substantial variability in 183 antibody responses to SARS-CoV-2 proteins among recovered adults. The median signal of the 184 Ortho Clinical assay was 15.4 S/C (95% CI: 12.7-18.0 S/C) (Fig 1B), the Abbott assay was 5.2 185 S/C (95% CI: 4.3-6.1 S/C) (Fig 1C), and the Roche assay was 23.94 COI (95% CI: 13.8-37.1 186 COI) (Fig 1D). As with the anti-S ELISA IgG, live virus neutralization titers spanned a broad 187 range (1:20 - 1:3,622), with a median titer of 1:316 (95% CI: 1:251-1:398) (Fig 1E). These

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results are consistent with substantial variability in neutralizing antibody responses to SARSCoV-2 proteins among recovered adults (12).

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191 Serological characteristics of donors

192 Linear relationships between each commercial assay were defined relative to the anti-S ELISA 193 IgG titer, Ortho-Clinical assay, and neutralization titer (Supplemental Figure 1). Slopes and 194 intercepts for each commercial assay relative to the anti-S ELISA, Ortho Clinical IgG, and 195 neutralization assay are found in **Table 2**. For the anti-S ELISA IgG titer, the initial FDA titer of 196 1:320 was used to establish high and low titers (Supplemental Fig. 1A). Interpolated signal by 197 the Ortho Clinical was 5.13 S/C (4.12-6.04), for the Abbott was 1.39 S/C (0.59 to 1.91) and for 198 the neutralization assay was 1.82 (1.68 to 1.93). Due to poor fit, an interpolated cutoff could not 199 be calculated for the Roche assay. For the Ortho Clinical assay, the cutoff of 12.0 S/C was used 200 to distinguish high- and low-titer units (Supplemental Fig 1B). The interpolated signal by the 201 Abbott assay was 3.62 S/C (3.1 to 4.07), for the Roche assay was 7.46 COI (4.78 to 10.73), for 202 the ELISA was 2.96 (2.87 to 3.04), and for the neutralization assay was 2.25 (2.09 to 2.37). Low-203 and high-titer cutoffs were also calculated at the Ortho Clinical cutoff of 4.62 S/C and 18.45 S/C, 204 respectively. For the neutralization assay, a single cutoff of 1:250 was used to determine low-205 and high-titer units (Supplemental Fig 1C). Relative to the neutralization assay, the interpolated 206 cutoffs for the Ortho Clinical was 12.39 S/C (10.38 to 14.16), for the Abbott was 3.92 S/C (3.14 207 to 4.57), for the Roche assay was 8.45 COI (3.48 to 15.89), and for the anti-S IgG ELISA was 208 3.05 (2.91 to 3.18).

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210 ROC curves were generated for each serological assay using the Ortho Clinical cutoff of 12 S/C 211 (Fig 2A) and the neutralizing cutoff of 1:250 (Fig 2B). Relative to the Ortho Clinical assays, all 212 assays had an AUC > 0.8, with the anti-S IgG ELISA having the best correlation (AUC = 0.927). 213 The assay with the greatest correlation with the neutralizing assay was the anti-S IgG ELISA, 214 with an AUC of 0.856 (0.771-0.941). Final cutoffs were established by averaging the ideal 215 cutoffs from linear regression in **Table 2** and the ideal cutoffs from the ROC curves using 216 Youden's Index. Using this approach, the average cutoff for distinguishing high- and low-titer 217 units by the Abbot assay was 3.82 S/C, the Roche assay was 10.89 COI, the anti-S IgG ELISA 218 was 3.08, 1:200 for the neutralization assay and 14.14 S/C for the Ortho-Clinical assay (Table 219 3). ROC curves were also generated relative to the low and high Ortho-Clinical cutoffs of 4.62 220 S/C and 18.45 S/C, respectively (Supplemental Fig 2) and for the low and high neutralizing 221 titers of 1:150 and 1:500, respectively (Supplemental Fig 3).

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223 Specimens were segregated as low- or high-titer using the Ortho Clinical cutoff of 9.5 S/C or 12 224 S/C and scatterplots generated (Fig. 3A and B). Using the cutoffs established in Table 3 (dotted 225 black lines), all four assays (Abbott, Roche, ELISA, and neutralization) demonstrated 226 comparable performance relative to the Ortho Clinical cutoffs of 9.5 S/C and 12 S/C for 227 identifying patients with high and low antibody titers. Decreasing the signal for identifying high-228 titer plasma on the Ortho Clinical assay led to improved NPA and PPA with the Abbott and 229 Roche assay and an improved NPA with a modest decrease in PPA with the anti-S IgG ELISA 230 and the neutralization titer (Supplemental Table 1). The overall agreement using the derived 231 cutoffs with the Ortho Clinical assay cutoff of 9.5 S/C was 92.4% for Abbott, 84.8% for Roche, 232 87.3% for the anti-S ELISA and 78.5% for the neutralization assay. Relative to the FDA Abbott

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233 cutoff of 4.5 S/C (dashed gray line) for identifying high-titer units, 5 additional specimens would 234 have been labeled as low-titer by the Abbott but high-titer by the Ortho Clinical assay. Using the 235 FDA cutoff of ≥ 109 COI for the Roche assay, all 79 specimens would have been qualified as 236 low-titer units. Specimens also were segregated as low- or high-titer using the neutralization 237 assay cutoff of 1:250 with similar results (Fig. 3C). The tiered Ortho Clinical and neutralizing 238 cutoffs used to identify patients with low medium and high titers are found in **Supplemental** 239 Figures 4 and 5, respectively. Patients with high ratios of nucleocapsid to S as measured by the 240 Abbott and Ortho assays were more likely to have low neutralizing antibody titers 241 (Supplemental Figure 6).

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243 **DISCUSSION**

Despite accumulating evidence of associations between commercial serological assay values and neutralizing antibody titers with human immunity and CCP efficacy, few published studies permit correlation between the assay formats in use. Here, we tested three widely used commercial serological assays, an EUA anti-S IgG ELISA, and neutralizing antibodies and correlated each assay with the ideal cutoffs for establishing high-titer plasma.

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An important finding from this study is that commercial assays and the anti-S ELISA performed similarly for identifying specimens with high neutralization titers. Our approach using linear regression for the Ortho Clinical assay with the FDA-established cutoff of 12 S/C and the neutralizing titer of 1:250, coupled with ROC curves that established maximal PPA and NPAidentified cutoffs made these assays largely interchangeable for identifying high-titer CCP. The antigenic target of the assay did not change the PPA and NPA, with assays measuring antibodies

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to the viral S protein performing similarly to those measuring antibodies to the nucleocapsid protein. This finding is similar to other studies from acutely infected patients with severe symptoms and patients with mild symptoms (10, 22).

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260 It is notable that FDA's reissued CCP authorization letter incorporated multiple EUA serological 261 assays, several of which are included in this report (16). The cutoff for the Abbott assay 262 established here (≥ 3.82 S/C) is similar to the cutoff of ≥ 4.5 S/C established by the FDA. This is 263 despite the lowered Ortho Clinical assay S/C cutoff (from 12 to 9.5) in the reissued CCP EUA 264 (16). Nonetheless, the Abbott assay cutoff of 3.82 S/C had better PPA with the Ortho Clinical 265 cutoffs of 9.5 S/C and 12.0 S/C, without sacrificing NPA. In contrast, the FDA-approved cutoff 266 of 109 COI for the Roche assay would have disqualified all units as low-titer, with a signal 267 approximately 10-fold higher than the ideal cutoff identified in this study. The derived cutoff 268 from this study with the anti-S IgG ELISA (1:1,202) also was lower than that established by the 269 FDA (1:2,880). This resulted in a considerable difference in PPA, with far more convalescent 270 donor units being excluded under the FDA cutoffs for ELISA than for the Ortho Clinical assay. 271 The cutoff identified in our study that best distinguishes neutralizing titers $\geq 1:250$ in the Ortho 272 Clinical assay was 14.14 S/C, higher than the original cutoff of 12.0 S/C from the FDA. An S/C 273 of 9 on the Ortho Clinical assay correlated to a neutralizing titer of \sim 1:100. This is notably similar to the neutralizing titer of 1:104 we found to be sufficient to reduce weight loss in mice 274 275 (23). Nevertheless, the neutralizing assay used in this study cannot be assumed to perform 276 similarly to the assay used in the BARDA study (14, 15) due to non-standardization of SARS-277 CoV-2 strains, cell lines, and reagents/procedures. These differences underlie the difficulties in

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| 278 | harmonizing SARS-CoV-2 | serological assay | results. These | discrepancies | should be | considered |
|-----|----------------------------|---------------------|------------------|----------------|-------------|------------|
| 279 | when attempting to use any | serological assay a | as a proxy for m | neasuring neut | ralizing an | tibodies. |

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281 Several studies demonstrate survival benefits with early, high-titer CCP administration and in 282 patients with hematological malignancies, implying a continued need to identify CCP or immune 283 globulin donors (15, 24–26). This study attempts to harmonize several commercially available 284 assays that have been extensively studied and published. While numerous studies have addressed 285 the analytical performance of available serological assays, little correlative information is 286 available in the published literature to relate multiple serological assay results. Many blood 287 centers in the US are currently using or switching to Ortho Clinical IgG assay for identification 288 of high-titer units. The data presented here suggest that multiple commercial assays could be 289 used to identify CCP donors with high levels of neutralizing antibodies. This study may also 290 provide useful information for contextualizing previous seroprevalence studies and multiple CCP 291 studies across the literature emerging from this pandemic.

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293 There are several limitations associated with this study. Among the greatest limitations is the 294 lack of standardization between assays, even among the same manufacturers. This was 295 previously noted with the neutralization assay, though the same is true among commercial 296 assays. Since several of the assays have been designated as qualitative (*i.e.* the Roche, Abbott, 297 and Ortho Clinical assays), there is limited evidence that semi-quantitative results are 298 comparable between different instruments by the same manufacturer above the cutoff. For 299 example, since there is no material to verify linearity at higher concentrations, a result of 15 S/C 300 at one institution using the Ortho Clinical assay may vary from the Ortho Clinical assay at

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301 another institution. This may underlie the differences between the established cutoff and FDA 302 cutoff for the Roche assay. In general, this problem will continue to plague the field until 303 quantitative assays are universally adopted and standardized to SARS-CoV-2 antibody reference 304 material, such as that recently released by the World Health Organization (27). This is further 305 complicated by unclear direction as to how to report a qualitative assay result as quantitative 306 under an EUA, which does not permit modification of the manufacturer's Instructions for Use. 307 Another limitation of the current study is that a limited number of assays were evaluated, 308 limiting the generalizability of results. It is also important to note that these specimens were 309 obtained early during the course of the pandemic, and that continued viral evolution (which may 310 lead to extensive antigenic changes in the S protein) means that the quantitative relationships in 311 this manuscript could become outdated. Ongoing studies are required to confirm the 312 relationships established here in patients infected with SARS-CoV-2 variants and using 313 neutralizing assays that utilize SARS-CoV-2 variants. Finally, while these results provide 314 evidence of that the cutoffs identified by the FDA are helpful for identifying high-titer CCP 315 units, there were several specimens by each assay that were not in agreement. These specimens 316 demonstrate the requirement for further study before the cutoffs proposed by the FDA are 317 modified.

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In conclusion, we demonstrate that assays based on nucleoprotein antibodies (Roche, Abbott) and neutralization were positively associated with Ortho assay results (anti-S), though their ability to distinguish FDA high-titer specimens was marginal. Association with a traditional ELISA serologic test was high. All assays were positively associated with neutralization titers, though associations were strongest with S-based assays.

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337 **REFERENCES**

- FDA. 2020. In Vitro Diagnostics EUAs. <u>https://www.fda.gov/medical-devices/coronavirus-</u>
 <u>disease-2019-covid-19-emergency-use-authorizations-medical-devices/in-vitro-diagnostics-</u>
 euas. Accessed March 2021.
- 2. Infectious Diseases Society of America. 2020. IDSA COVID-19 Antibody Testing Primer.
- 342 3. CDC. 2020. Interim Guidelines for COVID-19 Antibody Testing. Centers for Disease
- 343 Control and Prevention. <u>https://www.cdc.gov/coronavirus/2019-</u>
- 344 <u>ncov/lab/resources/antibody-tests-guidelines.html</u>. Accessed March 2021.

Farnsworth 17

| 345 | 4. | The Infectious | Diseases Society | y of America | Guidelines o | on the Diagnosis | of COVID-19: |
|-----|----|----------------|------------------|--------------|--------------|------------------|--------------|
| | | | - | | | 0 | |

- 346 Molecular Diagnostic Testing. <u>https://www.idsociety.org/practice-guideline/covid-19-</u>
- 347 guideline-diagnostics/. Accessed January 2021.
- 348 5. Farnsworth CW, Anderson NW. 2020. SARS-CoV-2 Serology: Much Hype, Little Data.
- 349 Clin Chem 66:875–877.
- 350 6. Theel ES, Slev P, Wheeler S, Couturier MR, Wong SJ, Kadkhoda K. 2020. The Role of
 351 Antibody Testing for SARS-CoV-2: Is There One? J Clin Microbiol 58:e00797-20.
- 352 7. Tang MS, Hock KG, Logsdon NM, Hayes JE, Gronowski AM, Anderson NW, Farnsworth
- 353 CW. 2020. Clinical Performance of Two SARS-CoV-2 Serologic Assays. Clin Chem
 354 66:1055–1062.
- Tang MS, Hock KG, Logsdon NM, Hayes JE, Gronowski AM, Anderson NW, Farnsworth
 CW. 2020. Clinical Performance of the Roche SARS-CoV-2 Serologic Assay. Clin Chem
 66:1107–1109.
- 358 9. Theel ES, Harring J, Hilgart H, Granger D. 2020. Performance Characteristics of Four

359 High-Throughput Immunoassays for Detection of IgG Antibodies against SARS-CoV-2. J
360 Clin Microbiol 58:e01243-20.

- 10. Tang MS, Case JB, Franks CE, Chen RE, Anderson NW, Henderson JP, Diamond MS,
- Gronowski AM, Farnsworth CW. 2020. Association between SARS-CoV-2 neutralizing
 antibodies and commercial serological assays. Clin Chem 66:1538-1547.

| 364 | 11. | Bal A, Pozzetto B, Trabaud M-A, Escuret V, Rabilloud M, Langlois-Jacques C, Paul A, |
|-----|-----|---|
| 365 | | Guibert N, D'Aubarede-Frieh C, Massardier-Pilonchery A, Fabien N, Goncalves D, |
| 366 | | Boibieux A, Morfin-Sherpa F, Pitiot V, Gueyffier F, Lina B, Fassier J-B, Trouillet-Assant |
| 367 | | S, COVID SER STUDY GROUP. 2021. Evaluation of high-throughput SARS-CoV-2 |
| 368 | | serological assays in a longitudinal cohort of patients with mild COVID-19: clinical |
| 369 | | sensitivity, specificity and association with virus neutralization test. Clin Chem |
| 370 | | 10.1093/clinchem/hvaa336. |
| 371 | 12. | Klein SL, Pekosz A, Park H-S, Ursin RL, Shapiro JR, Benner SE, Littlefield K, Kumar S, |
| 372 | | Naik HM, Betenbaugh M, Shrestha R, Wu AA, Hughes RM, Burgess I, Caturegli P, |
| 373 | | Laeyendecker O, Quinn TC, Sullivan DJ, Shoham S, Redd AD, Bloch EM, Casadevall A, |
| 374 | | Tobian AAR. 2020. Sex, age, and hospitalization drive antibody responses in a COVID-19 |
| 375 | | convalescent plasma donor population. J Clin Invest 130:6141-6150. |
| 376 | 13. | Amanat F, Stadlbauer D, Strohmeier S, Nguyen THO, Chromikova V, McMahon M, Jiang |
| 377 | | K, Arunkumar GA, Jurczyszak D, Polanco J, Bermudez-Gonzalez M, Kleiner G, Aydillo T, |
| 378 | | Miorin L, Fierer DS, Lugo LA, Kojic EM, Stoever J, Liu STH, Cunningham-Rundles C, |
| 379 | | Felgner PL, Moran T, García-Sastre A, Caplivski D, Cheng AC, Kedzierska K, Vapalahti |
| 380 | | O, Hepojoki JM, Simon V, Krammer F. 2020. A serological assay to detect SARS-CoV-2 |
| 381 | | seroconversion in humans. 7. Nature Medicine 26:1033–1036. |
| 382 | 14. | FDA. 2020. Recommendations for Investigational COVID-19 Convalescent Plasma. FDA. |
| 383 | | FDA. https://www.fda.gov/vaccines-blood-biologics/investigational-new-drug-ind-or- |
| 384 | | device-exemption-ide-process-cber/recommendations-investigational-covid-19- |
| 385 | | convalescent-plasma. Accessed June 2020. |

| 3 | 86 | 6 15. | Joyner MJ, 0 | Carter RE, S | Senefeld JW, | Klassen SA, | , Mills JR, | Johnson PW | , Theel ES, |
|---|----|-------|--------------|--------------|--------------|-------------|-------------|------------|-------------|
| | | | | , | , | , | | | , |

- 387 Wiggins CC, Bruno KA, Klompas AM, Lesser ER, Kunze KL, Sexton MA, Diaz Soto JC,
- 388 Baker SE, Shepherd JRA, van Helmond N, Verdun NC, Marks P, van Buskirk CM, Winters
- 389 JL, Stubbs JR, Rea RF, Hodge DO, Herasevich V, Whelan ER, Clayburn AJ, Larson KF,
- 390 Ripoll JG, Andersen KJ, Buras MR, Vogt MNP, Dennis JJ, Regimbal RJ, Bauer PR, Blair
- 391 JE, Paneth NS, Fairweather D, Wright RS, Casadevall A. 2021. Convalescent Plasma
- Antibody Levels and the Risk of Death from Covid-19. N Engl J Med 384:1015-1027.
- 393 16. FDA. 2020. COVID Convalescent Plasma EUA.
- 394 <u>https://www.fda.gov/media/141477/download</u>. Accessed September 2020.
- 395 17. Villa C. COVID-19 Convalescent Plasma EUA Decision.
- 396 https://www.fda.gov/media/141480/download . Accessed January 2021.
- 397 18. FDA. 2020. FDA Issues Emergency Use Authorization for Convalescent Plasma as
- 398 Potential Promising COVID–19 Treatment, Another Achievement in Administration's
- 399 Fight Against Pandemic. <u>https://www.fda.gov/news-events/press-announcements/fda-</u>
- 400 issues-emergency-use-authorization-convalescent-plasma-potential-promising-covid-19-
- 401 <u>treatment</u>. Accessed December 2020.
- 402 19. FDA. 2021. Revised COVID Convalescent Plasma EUA.
- 403 <u>https://www.fda.gov/media/141477/download</u>. Accessed March 2021.
- 404 20. Suhandynata RT, Hoffman MA, Huang D, Tran JT, Kelner MJ, Reed SL, McLawhon RW,
- 405 Voss JE, Nemazee D, Fitzgerald RL. 2021. Commercial Serology Assays Predict
- 406 Neutralization Activity against SARS-CoV-2. Clin Chem 67:404–414.

| 407 | 21. | FDA. Mount Sinai Laboratory COVID-19 ELISA IgG Antibody Test EUA Summary. |
|-----|-----|--|
| 408 | | https://www.fda.gov/media/137032/download. Accessed March 2021. |
| 409 | 22. | Long Q-X, Liu B-Z, Deng H-J, Wu G-C, Deng K, Chen Y-K, Liao P, Qiu J-F, Lin Y, Cai |
| 410 | | X-F, Wang D-Q, Hu Y, Ren J-H, Tang N, Xu Y-Y, Yu L-H, Mo Z, Gong F, Zhang X-L, |
| 411 | | Tian W-G, Hu L, Zhang X-X, Xiang J-L, Du H-X, Liu H-W, Lang C-H, Luo X-H, Wu S-B, |
| 412 | | Cui X-P, Zhou Z, Zhu M-M, Wang J, Xue C-J, Li X-F, Wang L, Li Z-J, Wang K, Niu C-C, |
| 413 | | Yang Q-J, Tang X-J, Zhang Y, Liu X-M, Li J-J, Zhang D-C, Zhang F, Liu P, Yuan J, Li Q, |
| 414 | | Hu J-L, Chen J, Huang A-L. 2020. Antibody responses to SARS-CoV-2 in patients with |
| 415 | | COVID-19. Nat Med 26:845–848. |
| 416 | 23. | Winkler ES, Gilchuk P, Yu J, Bailey AL, Chen RE, Chong Z, Zost SJ, Jang H, Huang Y, |
| 417 | | Allen JD, Case JB, Sutton RE, Carnahan RH, Darling TL, Boon ACM, Mack M, Head RD, |
| 418 | | Ross TM, Crowe JE, Diamond MS. 2021. Human neutralizing antibodies against SARS- |
| 419 | | CoV-2 require intact Fc effector functions for optimal therapeutic protection. Cell |
| 420 | | https://doi.org/10.1016/j.cell.2021.02.026. |
| 421 | 24. | Joyner MJ, Senefeld JW, Klassen SA, Mills JR, Johnson PW, Theel ES, Wiggins CC, |
| 422 | | Bruno KA, Klompas AM, Lesser ER, Kunze KL, Sexton MA, Diaz Soto JC, Baker SE, |
| 423 | | Shepherd JRA, van Helmond N, van Buskirk CM, Winters JL, Stubbs JR, Rea RF, Hodge |
| 424 | | DO, Herasevich V, Whelan ER, Clayburn AJ, Larson KF, Ripoll JG, Andersen KJ, Buras |
| 425 | | MR, Vogt MNP, Dennis JJ, Regimbal RJ, Bauer PR, Blair JE, Paneth NS, Fairweather D, |
| 426 | | Wright RS, Carter RE, Casadevall A. 2020. Effect of Convalescent Plasma on Mortality |
| 427 | | among Hospitalized Patients with COVID-19: Initial Three-Month Experience. medRxiv |
| 428 | | https://doi.org/10.1101/2020.08.12.20169359. |

| 429 | 25. | Duan K, Liu B, Li C, Zhang H, Yu T, Qu J, Zhou M, Chen L, Meng S, Hu Y, Peng C, Yuan |
|-----|-----|--|
| 430 | | M, Huang J, Wang Z, Yu J, Gao X, Wang D, Yu X, Li L, Zhang J, Wu X, Li B, Xu Y, |
| 431 | | Chen W, Peng Y, Hu Y, Lin L, Liu X, Huang S, Zhou Z, Zhang L, Wang Y, Zhang Z, Deng |
| 432 | | K, Xia Z, Gong Q, Zhang W, Zheng X, Liu Y, Yang H, Zhou D, Yu D, Hou J, Shi Z, Chen |
| 433 | | S, Chen Z, Zhang X, Yang X. 2020. Effectiveness of convalescent plasma therapy in severe |
| 434 | | COVID-19 patients. Proc Natl Acad Sci USA 117:9490-9496. |
| 435 | 26. | Thompson MA, Henderon JP, Shah PK, Rubinsteain SM, Joyner MJ, Choueiri TK, Flora |
| 436 | | DB, Griffiths EA, Gulati AP, Hwang C, Koshkin VS, Papadopoulos EB, Robilotti EV, Su |
| 437 | | CT, Wulff-Burchfield EM, Xie Z, Yu PP, Mishra S, Senefeld JW, Shah DP, Warner JL. |
| 438 | | Convalescent Plasma and Improved Survival in Patients with Hematologic Malignancies |
| 439 | | and COVID-19 medRxiv. |
| 440 | | https://www.medrxiv.org/content/10.1101/2021.02.05.21250953v1 |
| 441 | 27. | WHO. 2021. First WHO International Standard Anti-SARS-CoV-2 Immunoglobulin |
| 442 | | (Human). |
| 443 | | https://www.nibsc.org/products/brm_product_catalogue/detail_page.aspx?catid=20/136. |
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Farnsworth 22

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461 **Table 1.** Convalescent plasma donors' characteristics

| Variable | Total n=79 (%) |
|----------------------|----------------|
| Age (median [range]) | 49 (20-69) |
| Sex | |
| Female | 44 (55.7) |
| Male | 35 (44.3) |
| Race | |
| White | 72 (91.1) |
| Black | 4 (5.1) |
| Asian | 2 (2.5) |
| Other | 1 (1.3) |
| Comorbidities | |
| Asthma | 15 (19) |
| Lung disease | 0 (0) |
| Heart disease | 2 (2.5) |

Farnsworth 23

| Hypertension | 13 (16.5) |
|--|-----------|
| Diabetes mellitus | 3 (3.8) |
| Cancer | 6 (7.6) |
| Autoimmune disease | 5 (6.3) |
| Other | 28 (35.4) |
| Hospitalization | 2 (2.5) |
| Duration of symptoms in days (median | 12 (1-31) |
| [range]) | |
| Days from symptom onset to positive test | 4 (0-20) |
| (median [range]) | |

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464 **Table 2.** Linear fit and interpolated cutoffs for each assay

| | | Ortho Clinical | Abbott | Roche | anti-S ELISA | Neutralizing |
|----------------|---------------------------------|---------------------------|---------------------------|------------------------------|-----------------------------------|------------------------------|
| anti-S ELISA | Intercept | -632.5 | -359.5 | 467.9 | | -2875 |
| Titer | (95% CI) | (-1062 to - 203.1) | (-993.0 to 210.2) | (53.7 to 882.2) | | (-4281.0 to - 1468.0) |
| | Slope (95% CI) | 121.2 (96.05 to 146.4) | 337.3 (221.7 to 452.9) | 21.0 (11.8 to 30.2) | | 1625.0 (1070.0 to 2180.0) |
| | r^2 | 0.54 | 0.3 | 0.21 | | 0.31 |
| | FDA cutoff 1:320 (95% CI) | 5.13 (4.12- 6.04) | 1.39 (0.59 to 1.91) | NA* | | 1.82 (1.68 to 1.93) |
| | FDA cutoff 1:2,884 | 17.75 (16.8 to-18.75) | 5.64 (5.23 to 6.07) | 52.53 (43.05 to 65.31) | | 2.69 (2.62 to 2.774) |
| Ortho-Clinical | | | | | | |
| | Intercept (95% CI) | | 1.72 (-1.1 to 4.55) | 9.64 (7.29 to 11.99) | -24.549 (-28.74 to - 20.24) | -14.77 (-22.44 to -7.1) |
| | Slope (95% CI) | | 2.83 (2.29 to 3.38) | 0.16 (0.1 to 0.21) | 12.12 (10.83 to 13.4) | 11.91 (8.89 to 14.94) |
| | r^2 | | 0.58 | 0.5 | 0.82 | 0.44 |
| | FDA cutoff 12 (95% CI) | | 3.62 (3.1 to 4.07) | 7.46 (4.78 to 10.73) | 3.01 (2.87 to 3.04) | 2.25 (2.09 to 2.37) |
| | High Cutoff 18.45(95% CI) | | 5.91 (5.45 to 6.46) | 41.02 (2.73 to 84.33) | 3.54 (3.46 to 3.64) | 2.79 (2.66 to 2.96) |
| | Low Cutoff 4.62 (95%CI) | | 1.02 (0.03 to 1.73) | 1.06 (0.43 to 1.93) | 2.4 (2.28 to 2.51) | 1.63 (1.3 to 1.83) |
| Neutralizing | | | | | | |

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| (EC50) | Intercept (95% | 1.94 | 9.64 | 2.09 | 0.59 | |
|--------|-----------------|---------------|---------------|---------------|----------|--|
| | CI) | (1.77 to 2.1) | (7.29 to | (1.9 to 2.28) | (0.24 to | |
| | | | 11.99) | | 0.94) | |
| | Slope (95% CI) | 0.04 | 0.16 | 0.34 | 0.59 | |
| | _ | (0.03 to | (0.1 to 0.21) | (0.2 to | (0.48 to | |
| | | 0.05) | | 0.47) | 0.69) | |
| | r^2 | 0.44 | 0.5 | 0.24 | 0.61 | |
| | FDA cutoff 250 | 12.39 | 3.92 | 8.45 | 3.09 | |
| | (95% CI) | (10.38 to | (3.14 to | (3.48 to | (2.97 to | |
| | | 14.16) | 4.57) | 15.89) | 3.2) | |
| | High Cutoff 500 | 20.49 | 6.28 | 67.04 | 3.61 | |
| | (95% CI) | (18.56 to | (5.58 to | (33.4 to > | (3.49 to | |
| | | 22.94) | 7.23) | assay) | 3.75) | |
| | Low Cutoff 125 | 12.39 | 1.58 | 1.07 | 2.58 | |
| | (95%CI) | (10.38 to | (0.11 to | (0.01 to | (2.38 to | |
| | | 14.16) | 2.48) | 2.78) | 2.72) | |
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| 469 | Table 3. PPA and NPA for each a | assay at various | cutoffs relative to the | he Ortho | Clinical a | ssay and |
|-----|---------------------------------|------------------|-------------------------|----------|------------|----------|
|-----|---------------------------------|------------------|-------------------------|----------|------------|----------|

470 Neutralizing titers

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| | | Ortho-Clinical | | Neutral | | |
|--------|---------|----------------|---------------|----------------|---------------------|-------|
| | | FDA cutoff 12 | Youden Index | FDA cutoff 250 | Youden Index | Mean |
| Abbott | | | | | | |
| | Cutoff | 3.62 | 3.78 | 3.92 | 3.96 | 3.82 |
| | PPA | 92.3 (81.8 to | 92.3 (81.8 to | 87.2 (74.8 to | 87.2 (74.8 to 94.0) | |
| | (95%C) | 97.0) | 97.0) | 94.0) | | |
| | NPA | 74.1 (55.3 to | 81.5 (63.3 to | 68.8 (51.4 to | 68.8 (51.4 to 82.1) | |
| | (95%CI) | 86.8) | 91.8) | 82.1) | | |
| Roche | | | | | | |
| | Cutoff | 7.46 | 16.78 | 8.45 | 10.86 | 10.89 |
| | PPA | 90.4 (79.4 to | 82.7 (70.3 to | 87.2 (74.8 to | 81.1 (72.3 to 92.6) | |
| | (95%C) | 95.8) | 90.6) | 94.0) | | |
| | NPA | 62.96 (44.2 to | 85.2 (67.5 to | 59.4 (42.3 to | 56.3 (39.3 to 71.8) | |
| | (95%CI) | 78.5) | 94.1) | 74.5) | | |
| ELISA | | | | | | |
| | Cutoff | 2.96 | 3.04 | 3.05 | 3.25 | 3.08 |
| | PPA | 94.23 (84.4 to | 92.3 (81.8 to | 85.1 (72.3 to | 83.0 (69.9 to 91.1) | |
| | (95%C) | 98.4) | 97.0) | 92.6) | | |
| | NPA | 74.07 (55.3 to | 85.2 (67.5 to | 65.6 (48.3 to | 75.0 (57.9 to 86.8) | |
| | (95%CI) | 86.8) | 94.1) | 79.6) | | |

Farnsworth 25

| Neut/ Titer | | | | | | |
|----------------|----------------|------------------------|------------------------|------------------------|---------------------|-------|
| | Cutoff | 2.25 | 2.34 | | | 2.30 |
| | PPA (95%C) | 86.5 (74.7 to 93.3) | 84.6 (72.5 to 92.0) | | | |
| | NPA (95%CI) | 55.6 (37.3 to 72.4) | 66.7 (47.8 to 81.4) | | | |
| Ortho-C | linical | | | | | |
| | Cutoff | | | 12.38 | 15.9 | 14.14 |
| | PPA (95%C) | | | 80.9 (67.5 to 89.6) | 70.2 (56.0 to 81.4) | |
| | NPA (95%CI) | | | 59.4 (42.3 to 74.5) | 84.4 (66.3 to 93.1) | |

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477 **Figure Legends**

478 Figure 1. Histogram of each assay for 79 convalescent plasma donors with confirmed SARS-

479 CoV-2 infection. Dashed line is the median.

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481 **Figure 2.** ROC Curves for serological SARS-CoV-2 assays with **A**. the Ortho-Clinical IgG assay

482 using a cutoff of 12 to distinguish high titers and **B**. neutralizing assay using a cutoff of 250.

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484 Figure 3. Scatter plots of EUA serological SARS-CoV-2 assays using the ideal cutoffs identified

485 by linear regression and ROC curves relative to A. Ortho-Clinical cutoff of 9.5, B. Ortho Clinical

486 cutoff of 12.0 and **C.** a neutralizing titer cutoff of 1:250. Dotted line represents the mean cutoffs

487 identified in Table 3. Dashed lines represent the FDA approved cutoffs from EUA.











