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Evaluation of transplacental transfer of mRNA vaccine products and functional antibodies during pregnancy and early infancy

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30

31 Abstract

32 Studies are needed to evaluate the safety and effectiveness of mRNA SARS-CoV-2 vaccination 33 during pregnancy, and the levels of protection provided to their newborns through placental 34 transfer of antibodies. We evaluated the transplacental transfer of mRNA vaccine products and 35 functional anti-SARS-CoV-2 antibodies during pregnancy and early infancy in a cohort of 20 36 individuals vaccinated during pregnancy. We found no evidence of mRNA vaccine products in 37 maternal blood, placenta tissue, or cord blood at delivery. However, we found time-dependent 38 efficient transfer of IgG and neutralizing antibodies to the neonate that persisted during early 39 infancy. Additionally, using phage immunoprecipitation sequencing, we found a vaccine-specific 40 signature of SARS-CoV-2 Spike protein epitope binding that is transplacentally transferred 41 during pregnancy. In conclusion, products of mRNA vaccines are not transferred to the fetus 42 during pregnancy, however timing of vaccination during pregnancy is critical to ensure 43 transplacental transfer of protective antibodies during early infancy.

44

45 Keywords

46 SARS-CoV-2, COVID-19, Pregnancy, Vaccine, Antibody, Neonatal Immunity, Neutralizing

47 Antibody, Phage Array, mRNA Vaccination, BNT-162b2, mRNA-1273, Placenta, Cord Blood

48

49 Introduction

50 Growing evidence has shown that pregnant individuals are at higher risk for SARS-CoV-51 2-related morbidity and mortality¹⁻⁴. Despite this, vaccination uptake by pregnant individuals has 52 been slower than the general population⁵, in part because of maternal concern of adverse

53 effects on the embryo or fetus, even with strong consensus recommendations for COVID-19 vaccination prior to or during pregnancy from several medical societies⁶. Pregnant individuals 54 55 were excluded from initial vaccine trials, and complete data on safety, efficacy, optimal timing of the vaccine in pregnancy, or its impact on the fetus has been delayed⁷, which may impact 56 57 individual medical decision making. Current COVID-19 vaccines fully approved and under emergency use in the United States include the mRNA vaccines BNT-162b2 and mRNA-1273, 58 which target the SARS-CoV-2 Spike protein and stimulate protective immune responses ^{8,9}. In 59 addition to protecting the mother against severe disease, vaccination during pregnancy may 60 61 protect the newborn through passive transfer of maternal immunoglobulin. SARS-CoV-2 62 infection and vaccination during pregnancy produces an IgG response that is transferred to the fetus¹⁰⁻¹⁶. Evidence of newborn protection might help address maternal concerns about adverse 63 effects. However, detailed studies of the transplacental transfer of vaccine products and 64 65 vaccine-related antibody dynamics, functional properties, and persistence during infancy of transferred SARS-CoV-2 antibodies are needed to provide such evidence. 66

We examined the transplacental transfer of mRNA vaccine products and humoral
responses using samples from pregnant individuals and their infants vaccinated with either
BNT-162b2 or mRNA-1273 mRNA vaccine during pregnancy.

70

71 Results:

Cohort: We evaluated 20 pregnant individuals who received COVID-19 mRNA vaccines during pregnancy and their infants. Participants were vaccinated between December 2020 and April 2021. Gestational age at first dose ranged from 13 weeks to 40 weeks (mean 31.2, SD 5.9 weeks). Nineteen participants delivered live, singleton infants between January 2021 through April 2021 at gestational ages ranging from 37.4 to 41.1 weeks (mean 39.2, SD 1.1 weeks). One participant who was vaccinated at 13 weeks had a termination of pregnancy due to a lethal skeletal dysplasia of genetic etiology at 20.4 weeks. Eight participants received BNT-162b2

79 (Pfizer-BioNTech) and twelve received mRNA-1273 (Moderna) vaccines. Eighteen participants 80 received both vaccine doses prior to delivery, and two participants received the second dose 81 after delivery. The time from first mRNA vaccine dose ranged from 6-97 (mean 51, SD 24.3) 82 days prior to delivery, time from the second dose ranged from 2-75 (mean 32, SD 21.3) days 83 prior to delivery, and in two participants 15 and 21 days after delivery. No participants received a 3rd dose prior to delivery. Infants born to vaccinated mothers were followed up at convenience 84 85 time points ranging from age 3 weeks to 15 weeks of life (mean 8.3, SD 3.2). Further demographic data is detailed in Table S1. 86

87

88

Vaccine mRNA products do not cross the placenta

89 To determine the transplacental transfer of mRNA vaccine derived products, we examined available maternal blood at delivery, placenta tissue, and cord blood for Spike protein 90 91 by Western blot and vaccine mRNA by PCR. All available delivery samples (maternal blood, 92 placental tissue, and cord blood) were negative for Spike protein by Western blot (Supp Figure 93 1, Supp Table 3) and did not have detectable levels of vaccine mRNA by PCR (Suppl Table 3). 94 Together, this indicates that products of mRNA vaccination do not reach the fetus after 95 vaccination during pregnancy at readily detectable levels.

96

97 mRNA vaccination in pregnancy leads to a robust antibody response

Similar to prior studies^{14,15,17}, we found that mRNA vaccination during pregnancy led to 98 an increase in anti-SARS-Cov-2 IgG following dose 1 (n=7, mean 388.6, SD 224.8 RFU) and an 99 100 even further robust increase after vaccination dose 2 (n=12, mean 3214, SD 1383 RFU). Anti-

101 SARS-CoV-2 IgM (n=7, mean 53.3, SD 50.2 RFU) was detected in two maternal participants

102 following dose 1, but only 1 participant following dose 2 (n=12, mean 23.8, SD 17 RFU, Fig 1).

104 Vaccine induced anti-SARS-CoV-2 IgG and neutralizing antibodies are transplacentally

105 transferred

106 We then evaluated the transplacental transfer of maternally derived anti-SARS-CoV-2 107 IgG antibodies to their infants. Maternal blood at delivery was available in 19/20 participants and 108 cord blood was available in 17/20 participants. Anti-SARS-CoV-2 IgG was detectable in 94.7% 109 (18/19) of maternal blood samples at delivery (mean 3235, range [10, 7811] RFU). Anti-SARS-110 CoV-2 lgG was detectable in 88.2% (15/17) cord blood samples (mean 2243, range [2, 4959] 111 RFU). One participant received one mRNA vaccine dose 9 days prior to delivery, and both the 112 maternal and cord blood were negative for IgG at the time of delivery. Another participant 113 received two doses of mRNA vaccine (23 and 2 days) prior to delivery and maternal blood was 114 positive at 55 RFU (positive cutoff >50 RFU), however cord blood IgG was negative (Figure 2A). 115 Maternal and cord blood anti-SARS-CoV-2 IgG levels were moderately correlated, but not 116 statistically significant (p=0.074, R_s=0.446, Fig 2A). All cord blood samples were anti-SARS-117 CoV-2 IgM negative.

118 We next evaluated the transplacental transfer of neutralizing antibody titers by a label-119 free surrogate neutralization assay (sVNT) from mother to cord blood. Maternal and cord blood 120 at delivery had robust neutralizing responses (maternal n=17, mean 220.2, range [0, 422]. Cord 121 blood n=16, mean 296.6, range [0, 485], Fig 2B). All mother-infant dyads with positive IgG 122 serology at delivery had detectable transplacental transfer of neutralizing antibodies with the 123 exception of one pair in which the mother was borderline IgG positive at delivery and cord blood 124 was negative, for which both maternal and cord blood were negative for neutralizing titers (Fig 125 2B). However, maternal and cord blood neutralizing titers were not significantly correlated 126 (p=0.361, $R_s=-0.244$, Fig 2B). Taken together, this indicates that maternal mRNA vaccination 127 induces functional neutralizing antibodies that are transferred to the infant.

128

129

Maternally-derived vaccine induced anti-SARS-CoV-2 IgG and neutralizing antibodies

130 persist through early infancy

131 A subset of infants was sampled at convenience timepoints during follow up (infant 132 n=11, weeks of life range [3,15] mean 8.3 weeks). Anti-SARS-CoV-2 lgG levels were positive in 81.8% of infants at follow-up (9/11 infants, mean 1290, range [1, 3225] RFU, Fig 2A), with one 133 134 infant still IgG positive at 12 weeks of age (Fig 2C). The two infants that were IgG negative at 135 follow up were both born to mothers who received only one vaccine dose prior to delivery (6 and 136 9 days, respectively). One of these infants did not have paired maternal or cord blood available 137 at the time of delivery for comparison, and the other was IgG negative in cord blood. Maternal and infant follow-up anti-SARS-CoV-2 IgG levels were not significantly correlated; however, 138 cord blood and infant follow-up IgG levels were significantly associated (p=0.492, R_s=0.249 and 139 140 p=0.021, R_s=0.76, respectively, Fig 2A). All infants were IgM negative at the time of follow up. 141 All infants with available IgG positive samples at follow up had detectable neutralizing 142 titers (n=8, mean 154, range [41-256], Fig 2B). Maternal and infant follow-up neutralizing titers 143 were not significantly correlated, as well as cord and infant follow up neutralizing titers (p=0.665, 144 R_s =-0.191 and p=0.662, R_s =0.214, respectively, Fig 2B).

To compare the rate of decay of IgG antibody levels in mothers and their infants, we evaluated 5 dyads with paired maternal and infant blood samples on the same day at the time of follow-up (range 3-9 weeks post-delivery). Maternal antibody IgG levels decreased faster in mothers than infants (mean delta -974 RFU and -670 RFU, respectively. Fig 2E) at the follow up timepoint. Taken together this indicates, maternally-derived functional vaccine induced antibodies persist at high levels in newborns through early infancy during a critical time of immune vulnerability and may decay slower than maternal IgG antibodies.

152

153 Vaccine induced antibody timing and transplacental facilitated transfer

We assessed the relationship of anti-SARS-CoV-2 IgG levels to neutralizing antibody levels. We found a strong correlation between IgG and neutralizing titers in maternal plasma at delivery (R_s =0.744, p=0.0012) and infant follow up (R_s =0.738, p=0.046) timepoints, but no significant association between IgG and neutralizing titers in cord blood (R_s =0.121, p=0.656, Figure 3).

159 We then evaluated the impact of timing of vaccination on maternal antibody levels at 160 delivery. We found no statistically significant correlation between maternal IgG levels at delivery 161 and time since dose 1 (R_s =-0.335, p=0.160) and gestational age at delivery (R_s =0.270, p=0.265, 162 Fig 4A,B). This lack of correlation appeared to be driven by two participants that had low or 163 absent levels of antibodies at delivery and received their first dose of vaccine within 30 days of 164 delivery. We then evaluated neutralizing titers in those participants with known detectable IgG 165 levels at delivery and found that maternal neutralizing titers at delivery trended with days since 166 vaccine dose 1 but was not statistically significantly (R_s=-0.422, p=0.093), and maternal neutralizing titers at delivery was not associated with gestational age at dose 1 (R_s=0.074. 167 168 p=0.780). One participant was borderline IgG positive at delivery (vaccinated within 30 days of 169 delivery) and did not have detectable neutralizing titers at delivery (Fig 4C,D).

170 To assess facilitated antibody transfer, we evaluated cord-to-maternal antibody IgG and 171 neutralization titer ratios by time since vaccination and gestational age. We found that IgG ratios 172 were highly correlated with both time since first maternal vaccination dose and gestational age 173 at first dose (R_s=0.917, p<0.0001 and R_s=-0.739, p=0.002, respectively. Fig 4E,F). However, 174 neutralization titer cord-to-maternal ratios by time since first vaccination dose and gestational age at first dose were not significantly associated (R_s =0.366, p=0.179 and R_s =-0.032, p=0.913, 175 176 respectively, Figure 4G,H). Together, this may indicate that timing of vaccination in pregnancy is 177 critical for maternal-fetal antibody transfer, and functional neutralizing antibodies are 178 differentially transferred to the fetus as compared to total anti-SARS-CoV-2 IgG during 179 gestation.

180

mRNA vaccination leads to a unique SARS-CoV2 Spike protein antibody epitope binding signature

We next investigated antibody linear epitope binding and transplacental transfer using the PhIP-seq/VirScan SARS-CoV-2 Spike protein phage display array in mother-infant dyads at the time of birth (Figure 5). We found that timing of vaccination was important for the transplacental transfer of Spike protein epitope binding antibodies. Two mother-infant dyads had minimal Spike protein specific epitope binding. The first dyad only received one dose of mRNA vaccine 9 days prior to delivery, and the other dyad received the second vaccine dose 2 days prior to delivery.

190 We found high levels of SARS-CoV-2 Spike protein epitope binding in 4 major peaks we 191 designate as regions 1-4 (Figure 5A). Region 1 overlays the carboxy terminal of the N-terminal 192 domain. Region 2 overlaps with key residues for the S1/S2 cleavage site. Regions 3 and 4 are 193 within S2, flanking the fusion loop and the transmembrane portion of the Spike protein, 194 respectively. However, we found minimal binding in the receptor binding domain (RBD) of Spike 195 protein. Prior evaluation using the PhIP-seg/VirScan SARS-CoV-2 epitope phage array during 196 SARS-CoV-2 infection demonstrated similar binding in regions 3 and 4, however in SARS-CoV-197 2 infection there was minimal binding in regions 1 and 2 demonstrating that antibody epitope binding in these regions may be unique to vaccination¹⁸. Additionally, there is proportional 198 199 representation of linear epitope binding across the SARS-CoV-2 Spike protein proteome 200 between mothers and infants (Figure 5B). Taken together, SARS-CoV-2 antibody linear epitope 201 binding after vaccination during pregnancy shows similar patterns, with multiple 202 immunodominant regions found in the majority of mothers and infants. Some of these regions are unique to vaccination and not observed during natural infection¹⁸⁻²⁰. 203 204

205 Discussion

206 Among twenty women who received the COVID-19 mRNA vaccine during pregnancy, 207 our study found no evidence of transplacental transfer of mRNA vaccine products but did find 208 high levels of functional vaccine-derived antibodies that transferred to the infant at delivery and 209 persisted during early infancy. Additionally, we identified high levels of epitope binding in two regions of Spike protein unique to SARS-CoV-2 vaccination¹⁸. These data may address some of 210 211 the many unanswered questions regarding COVID-19 vaccination in pregnancy: including the 212 dynamics of antibody production in the pregnant immune state, and the optimal timing of 213 immunization in pregnancy to impart passive immunity to the newborn during the vulnerable first 214 few weeks of infancy.

Uptake of COVID-19 vaccination in pregnancy has been slow⁵, and reasons for vaccine 215 216 hesitancy are likely multifactorial — but theoretical concerns that vaccine mRNA could cross the 217 placenta have been raised. We found no evidence of mRNA vaccine products in any of our 218 delivery samples. Additionally, no infants in our study had a fetal immune response to Spike 219 protein as demonstrated by a negative anti-SARS-CoV-2 IgM antibody in cord blood and infant 220 follow up samples. This further supports the lack of transfer of vaccine products, as only IgG is 221 transferred from the mother, and IqM production would indicate an endogenous fetal immune 222 response which has rarely been seen in natural infection with SARS-CoV-2 during pregnancy ^{16,21-23}. This provides additional reassurance that mRNA vaccination is safe during pregnancy. 223 224 We found that the timing of immunization during pregnancy is important to ensure trans-225 placental transfer of protective antibodies to the neonate, and during critical windows of immune 226 vulnerability during early infancy. Consistent with prior studies showing robust immune 227 responses to mRNA vaccination^{14,15,17}, we found high levels of IgG after two doses of mRNA 228 vaccine. However, completion of the vaccination series well before delivery was important to 229 ensure transfer of antibodies to the infant. Two mothers only received one vaccine dose prior to 230 delivery and did not transfer antibodies as demonstrated by the lack of antibodies in cord (in one 231 with available cord blood) and in both infants at follow-up. Additionally, neutralizing antibodies

were not transferred in a mother who received her second dose of vaccine 2 days prior to delivery. All evaluated mothers who received both doses during pregnancy and with the second dose greater than 9 days prior to delivery transferred IgG and neutralizing antibodies to their infants. Consistent with early studies of antibody transfer after COVID-19 vaccination in pregnancy, most of our participants were vaccinated in the third trimester of pregnancy. Larger studies of individuals vaccinated prior to pregnancy and in the first and second trimester are needed to understand persistence and waning of vaccine-induced immune responses.

239 Additionally, we believe we are the first to report that infants in the first few months of life 240 continued to have maternal vaccine-derived anti-SARS-CoV-2 antibodies that were functional 241 as demonstrated by high levels of neutralizing antibodies presenting infants up to 12 weeks of age. This is consistent with known persistence of maternally-derived antibodies from other 242 vaccinations including pertussis, rubella, varicella²⁴⁻²⁶. Additionally, we have previously found 243 244 persistence of anti-SARS-CoV-2 IgG antibodies in infants after natural infection up to 6 months¹⁶. However, the functional capability of these antibodies as compared to anti-SARS-245 246 CoV-2 vaccination-derived antibodies is unknown. Further evaluation of the longitudinal 247 persistence of maternal vaccine-derived antibodies during infancy will be critical to determine 248 optimal timing of COVID-19 vaccination in infancy.

Evaluation of paired maternal and baby samples at post-partum follow up timepoints showed a faster decline in maternal IgG antibody levels than infants, suggesting that persistence of maternally-derived antibody may be prolonged for infants. Differences in renal excretion and neonatal Fc receptor (FcRn) expression, which is involved in antibody degradation²⁷ in the infant as compared to adults, could underly these differences and should be explored further.

255 Consistent with observations in non-pregnant adults, we found that IgG levels in mothers 256 at delivery, and at infant follow-up were highly correlated with neutralizing titers²⁸. However, 257 cord blood IgG levels did not correlate with neutralizing titers. Moreover, IgG cord-to-maternal

258 ratios, which represent a proxy of maternal to fetal antibody transfer, were highly correlated with 259 timing of vaccination (gestational age and days since the first dose), but cord-to-maternal 260 neutralizing titer ratios were not significantly associated with time since vaccination nor 261 gestational age. During gestation there is facilitated transfer of maternally derived antibodies through the binding of the neonatal Fc receptor in the synctiotrophoblast layer²⁹. Differences in 262 glycosylation^{30,31}, FcR/FcRn binding affinity^{17,32}, preferential IgG subclass transfer^{33,34} may be 263 264 different in functional neutralizing antibodies as compared to total IgG antibody transfer. 265 However, a limitation of this study is the majority of participants were vaccinated in the third 266 trimester. Further investigations on factors that influence the transport of functional antibodies 267 across trimesters are needed to understand antibody dynamics and optimal transfer of 268 protective antibodies to infants.

269 Using a PhIP-seg/VirScan SARS-CoV-2 Spike protein phage array we were able to 270 compare linear epitope antibody binding in mothers and their infants. Consistent with IgG and neutralizing antibody evaluation, timing of vaccination was critical to ensure the transplacental 271 272 transfer of antibodies to the infant. Additionally, we identified unique regions of antibody epitope 273 binding in our vaccinated cohort that were not identified using the same phage library in a prior evaluation of a cohort of SARS-CoV-2 infected individuals¹⁸. One of these regions included the 274 275 carboxy terminal of the N-terminal domain, with other work having shown that the N-terminal domain is targeted by neutralizing antibodies against Spike protein³⁵. We did not see significant 276 277 binding in the receptor binding domain (RBD), which may be attributable to the fact that the 278 phage display library displayed short, linear peptides while antibodies targeting RBD are known 279 to target conformational epitopes. Lastly, we found that the same immunodominant regions 280 targeted by antibodies targeting the Spike protein in both mothers and infants.

In summary, this work provides further evidence that mRNA vaccination is safe in pregnancy and demonstrates that it generates time-dependent protective, functional antibody responses in mothers and infants that persist during early infancy.

284

285 <u>Methods</u>

286 Cohort and Sample collection: The University of California San Francisco (UCSF) institutional 287 review board approved the study (20-32077). Written informed consent was obtained from all 288 participants. We enrolled 20 pregnant individuals who were vaccinated with either BNT-162b2 289 or mRNA-1273 mRNA vaccines. Pregnant individuals were followed through delivery, and their 290 infants were followed up to 12 weeks of life. Maternal blood was collected during pregnancy 291 (pre-vaccine, 3-4 weeks post-dose 1, 4-8 weeks post-dose 2). During delivery, maternal blood, 292 placenta tissue, and cord blood was collected. Infant follow-up blood was collected at 293 convenience timepoints. Whole blood was immediately added to RNAlater in a 1:1.3 ratio. 294 Plasma was isolated from whole blood by centrifugation and immediately cryopreserved. Full-295 thickness placental biopsy was collected within 1 hour of delivery, washed three times with 296 phosphate buffered saline, and preserved in RNAlater.

297

298 SARS-CoV-2 plasma serology. Anti-SARS-CoV-2 plasma IgM and IgG antibodies were 299 measured using the Pylon 3D automated immunoassay system (ET Healthcare, Palo Alto, CA). 300 In brief, guartz glass probes are pre-coated with either affinity purified goat anti Human IgM (IgM 301 capture) or Protein G (IgG capture) are dipped into diluted patient sample. Samples are 302 washed, and then the probe is dipped into the assay reagent containing both biotinylated 303 recombinant spike protein receptor binding domain (RBD) and nucleocapsid protein (NP). After 304 a washing, the probe is incubated with a Cy®5-streptavidin (Cy5-SA) polysaccharide conjugate 305 reagent, allowing for cyclic amplification of the fluorescence signal. The background corrected 306 signal is reported as relative fluorescent units (RFU) which is proportional to the amount of 307 specific antibodies in the sample allowing for quantification. Levels of IgM and IgG were 308 considered positive if greater than 50 relative fluorescence units.

309

310 SARS-CoV-2 neutralizing assay

311 SARS-CoV-2 antibody neutralization titers were measured using a label-free surrogate 312 neutralization assay (LF-sVNT) previously described²⁸. Briefly, the method measures the 313 binding ability of recombinant RBD (Sino Biological, Wayne, PA) coated onto sensing probes 314 (Gator Bio, Palo Alto, CA) to recombinant ACE2 (Sino Biological, Wayne, PA) after neutralizing RBD with SARS-CoV-2 antibodies in serum. Measurements were done using a thin-film 315 316 interferometry (TFI) label-free immunoassay analyzer (Gator Bio, Palo Alto, CA). Each serum 317 sample was diluted in a series (1:25, 1:100, 1:250, 1:500, 1:1000, 1:2000) in running buffer 318 (PBS at pH 7.4 with 0.02% Tween 20, 0.2% BSA, and 0.05% NaN₃) for analysis. The first 319 testing cycle for each diluted sample measured the binding ability of RBD to ACE2 with 320 neutralization, and the second cycle provided the full binding ability of RBD without 321 neutralization. In each cycle, the recorded time course of signals, as known as the sensorgram, 322 was recorded. The readout measured the signal increase in RBD-ACE2 complex formation, 323 representing the quantity of RBD-ACE2 complex on the sensing probe. A neutralization rate 324 was calculated as the ratio of the readout in the first cycle to that in the second cycle, presented 325 as a percentage. To obtain the neutralizing antibody titer (IC_{50}) for each serum sample, the 326 neutralization rates were plotted against dilutions, and the points were fitted using a linear 327 interpolation model. The reciprocal of the dilution resulting in a 50% neutralization rate was 328 defined as the neutralizing antibody titer.

329

SARS-CoV-2 Spike protein Western blot. Maternal blood and cord blood were diluted in
RNAlater in 1:1.3 ratio, placenta was preserved in RNAlater. Protein lysates were obtained from
samples using RIPA buffer (150 mM NaCl, 25 mM Tris-HCl (pH 7.4), 1% NP-40, 0.5% sodium
deoxycholate, 0.1% sodium dodecyl sulfate) containing Halt[™] protease inhibitor cocktail
(ThermoScientific).Cell Lysates were resolved by SDS/PAGE on a Bis-Tris methane 4–12%
polyacrylamide gel and transferred to a nitrocellulose membrane, blocked with 5% skimmed

milk diluted in PBS, an incubated overnight at 4°C with anti-SARS-CoV-2 Spike mouse mAb
(1A9, GeneTex) or anti-GAPDH rabbit polyclonal antibody (GTX100118, GeneTex) respectively
diluted 1:1,000 or 1:5,000 in blocking buffer. The membrane was washed in PBS buffer
containing Tween-20 (0.1%) and then incubated for 1 h with horseradish peroxidase-conjugated
anti-mouse and anti-rabbit secondary antibody (Jackson ImmunoResearch) diluted respectively
1:5,000 and 1:10,000. The membrane was thoroughly washed, and proteins visualized using
Immobilon Forte Western HRP substrate (Millipore).

343

344 SARS-CoV-2 Spike mRNA PCR. Maternal blood and cord blood were diluted in RNAlater in 345 1:1.3 ratio, placenta was preserved in RNAlater. Tissues were kept at -80°C until analyzed. 346 RNA was isolated from samples using the RNeasy Micro or Mini Kit (Qiagen) according to 347 manufacturer's protocol. RNA concentration was measured using nanodrop and all samples had 348 >30 ng/ul total RNA. 500ng RNA was transcribed into cDNA using gScript cDNA synthesis kit 349 (Quantabio). Primers were design to detect the vaccines mRNA (mRNA-1273 Moderna and 350 BNT162b2 Pfizer-BioNtech) as previously described³⁶. Forward primer: 351 AACGCCACCAACGTGGTCATC. Reverse primer: GTTGTTGGCGCTGCTGTACAC. Primers 352 were shown to detect samples containing as low as 1.5 pg of vaccine using vaccine standard 353 curve (Table S2), QuantaStudio 6 Flex (Applied Biosystems) instrument and SsoFast EvaGreen 354 supermix (Bio-Rad) were used for PCR reaction: 30 second 95°C followed by 40 cycles of 5 355 second 95°C and 20 seconds 60°C. All samples were run in triplicate as 20 µL reactions, and Ct 356 values corresponding to <1.5 pg of vaccine based on standard curve (Table S2) were 357 interpreted as a negative result. For vaccines cDNA standard curves, 10000 pg/µL vaccine 358 mRNA (as cDNA) sample was used for serial dilution in 1:3 ratio, up to 0.06 pq/μ L. Two μ L of 359 these diluted samples were used in each well to create standard curves. 360

361 PhIP-Seq/VirScan Coronavirus phage display assay

362 Immunoprecipitation of phage-bound patient antibodies

Maternal plasma at delivery and cord plasma were evaluated by PhIP-Seq/Virscan Coronavirus phage display. Construction of the Coronavirus PhIP-Seq library and detailed methods for immunoprecipitation, sequencing and bioinformatic processing of data are identical to what has previously been described¹⁸. For the purposes of the analysis conducted in this study, analysis was restricted to sero-reactivity against the SARS-CoV-2 Spike protein. As previously described, a total of two rounds of amplification and selection were performed for all PhIP-Seq analyses.

370

371 Next Generation Sequencing library prep

Amplicon sequencing library preps were performed using the Labcyte Echo 525 and an Integra Via Flow 96 and were identical to what has previously been described¹⁸. All libraries were pooled by equal volume, cleaned and size selected using Ampure XP beads at 1.0X per manufacturer's protocol. Libraries were quantified by High Sensitivity DNA Qubit and qualitychecked by High Sensitivity DNA Bioanalyzer. Sequencing was then performed on a NovaSeq S1 (300 cycle kit with 1.3 billion clusters) aiming for sequencing depths of at least 1 million reads per sample.

379

380 Bioinformatic Analysis of PhIP-Seq Data

Sequencing reads were aligned to a reference database of the full viral peptide library using the Bowtie2 aligner. For all VirScan libraries, the null distribution of each peptide's log10(rpK) was modeled using a set of 95 pre-pandemic, healthy control sera. All counts were augmented by 1 to avoid zero counts in the healthy control sera samples. Multiple distribution fits were examined for these data, with the Normal distribution showing the best fit. These null distributions were used to calculate p values for the observed log10(rpK) of each peptide within a given sample. The calculated p values were corrected for multiple hypothesis using the Benjamini-Hochberg

method. Any peptide with a corrected p value of < 0.001 was considered significantly enriched
over the healthy background. To identify regions targeted by host antibodies, all library peptides
were aligned to the SARS-CoV-2 reference genome. Using the aligned position of the
significantly enriched peptides which aligned full-length against the reference, we determined
the proportion of individuals (mothers and infants) that were reactive at each residue of the
Spike protein. All plots were generated using the R ggplot2 package.

394

395 Statistical analysis:

396 Statistical analyses were performed using PRISM v9.2 (GraphPad), STATA 16 (StataCorp), and 397 R version 3.6.3 and R Studio version 1.1.447. Descriptive statistics include mean, standard 398 deviations, and ranges for continuous variables. The Wilcoxon rank-sum test was used for two-399 group comparisons of continuous variables including maternal pre- and post-vaccine antibody 400 responses. Associations between continuous variables were assessed using Spearman's rank 401 correlation (R_s) including comparisons between maternal, cord and infant follow-up antibody IgG 402 and neutralizing titer responses, and timing of vaccination. Two-sided p values were calculated 403 for all test statistics, and p<0.05 was considered significant. PhIP-Seg/VirScan bioinformatics as 404 detailed above.

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406 Data Availability

The data set generated during and/or analyzed during the current study are available from thecorresponding author on reasonable request.

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- 431 analyzed mRNA PCR experiments, performed data analysis.
- 432 A.G.C. Recruited and consented enrollees, oversaw sample collection, performed chart review,
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- 437 H.C. and U.J. performed and helped design critical experiments, and data collection.
- 438 C.Y.L., V.J.L., M.C., L.W., S.B. Performed and coordinated sample collection, and data collection.

- 439 V.J.F. Helped conceive and coordinate the project.
- 440 A.P.M. Provide funding.
- 441 W.C.G. Helped design western blot and oversaw data analysis.
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- 443 K.L.L. Designed and oversaw all neutralizing antibody experiments.
- 444 J.R. Designed, analyzed, and oversaw phage immunoprecipitation sequencing assays.
- 445 S.L.G. conceived and designed the project, oversaw recruitment, oversaw experiment design,
- 446 oversaw data analysis, provided funding, and helped draft the manuscript.
- 447 M.P., Y.G., Y.U., L.L., A.H.B.W, W.C.G, K.L.L., and S.L.G verified the underlying data.
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463 Figure 1. Anti-SARS-CoV-2 IgG and IgM antibody responses following vaccination

A. Maternal plasma anti-SARS-CoV-2 IgG antibody relative fluorescence units (RFU) levels prior to vaccination (n=4), 3-4 weeks post-dose 1 (n=7), and 4-8 weeks post-dose 2 (n=12). B. Maternal plasma anti-SARS-CoV-2 IgM (RFU) levels prior to vaccination (n=4), 3-4 weeks post-dose 1 (n=7), and 4-8 weeks post-dose 2 (n=12). Wilcoxon rank-sum testing. Data represent median \pm quartiles, two-sided *p* values were calculated for all test statistics.





480 Figure 2. Paired maternal, cord, and infant IgG and neutralization antibodies

481 A. Paired maternal plasma at delivery (n=19), cord plasma (n=17), and infant follow-up (n=10)

- 482 by anti-SARS-CoV-2 IgG antibody relative fluorescence units (RFU), (Spearman's rank
- 483 correlation, dotted line indicates positive cutoff value of 50 RFU). B. Paired maternal plasma at

484	delivery (n=17), cord plasma (n=16), and infant follow-up (n=8) by SARS-CoV-2 label-free
485	surrogate neutralization assay (sVNT), (Spearman's rank correlation, dotted line indicates
486	positive cutoff value of 25). C. Paired cord plasma (n=9) and infant follow-up plasma (n=11)
487	anti-SARS-CoV-2 IgG by weeks of life. D. Paired cord plasma (n=7) and infant follow-up plasma
488	(n=8) label-free surrogate neutralization assay (sVNT) by weeks of life. E. Paired maternal
489	plasma at delivery (n=5), cord plasma (n=5), and paired maternal follow-up (n=5) and infant
490	follow-up plasma (n=5) anti-SARS-CoV-2 IgG. Two-sided <i>p</i> values were calculated for all test
491	statistics.
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- 504 Figure 3. Neutralization to IgG antibody correlation
- A. Maternal plasma at delivery (n=17) B. Cord plasma (n=16) C. Infant follow-up plasma (n=8)
- 506 SARS-CoV-2 label-free surrogate neutralization assay (sVNT) by anti-SARS-CoV-2 IgG
- 507 correlation (Spearman's rank correlation). Two-sided *p* values were calculated for all test
- 508 statistics.







536	antibody transfer ra	atio by gestational age	at vaccine dose 1	l (n=15). ⁻	Two-sided <i>p</i> values were
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- 537 calculated for all test statistics.

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- 563 binding
- A. Heatmap displaying results of significant enriched (p<0.001) linear SARS-CoV-2 Spike
- 565 protein epitope binding from 15 paired mother-infant dyads in maternal plasma at delivery and
- 566 cord plasma by vaccine type and time since vaccine dose 1. Areas of high cumulative epitope

- 567 binding designated by regions 1-4. B. Cumulative fold enrichment of mothers and infants linear
- 568 SARS-CoV-2 Spike protein epitope binding.

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• Supplemental Appendix Evaluation of transplacental of mRNA vaccine products and functional antibodies during pregnancy and early infancy. pdf