Evaluation of Antibody-Dependent Fc-Mediated Viral Entry, as Compared With Neutralization, in SARS-CoV-2 Infection

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Fc-mediated virus entry has been observed for many viruses, but the characterization of this activity in convalescent plasma against SARS-CoV-2 Variants of Concern (VOC) is undefined. In this study, we evaluated Fc-mediated viral entry (FVE) on FcγRIIa-expressing HEK293 cells in the presence of SARS-CoV-2 convalescent plasma and compared it with SARS-CoV-2 pseudovirus neutralization using ACE2-expressing HEK293 cells. The plasma were collected early in the pandemic from 39 individuals. We observed both neutralization and FVE against the infecting Washington SARS-CoV-2 strain for 31% of plasmas, neutralization, but not FVE for 61% of plasmas, and no neutralization or FVE for 8% of plasmas. Neutralization titer correlated significantly with the plasma dilution at which maximum FVE was observed, indicating Fc-mediated uptake peaked as neutralization potency waned. While total Spike-specific plasma IgG levels were similar between plasma that mediated FVE and those that did not, Spike-specific plasma IgM levels were significantly higher in plasma that did not mediate FVE. Plasma neutralization titers against the Alpha (B.1.1.7), Beta (B.1.351), Gamma (P.1) and Delta (B.1.617.2) VOC were significantly lower than titers against the Washington strain, while plasma FVE activity against the VOC was either higher or similar. This is the first report to demonstrate a functional shift in convalescent plasma antibodies from neutralizing and FVE-mediating against the earlier Washington strain, to an activity mediating only FVE and no neutralization activity against the emerging VOC, specifically the Beta (B.1.351) and Gamma (P.1) VOC. It will be important to determine the in vivo relevance of these findings.

Keywords: SARS-CoV-2, coronavirus – COVID-19, variant of concern, antibody dependent enhancement, neutralization, pseudovirus, Fc-mediated virus entry, spike (S) protein
INTRODUCTION

SARS-CoV-2 is a Betacoronavirus that is the causative agent of coronavirus disease 2019 (COVID-19). COVID-19 has a broad spectrum of disease presentation; an estimated 80% of patients are asymptomatic or have mild flu-like symptoms, while 20% of patients develop severe respiratory illness (1, 2). Critical outcomes include respiratory failure, multiple organ dysfunction and shock. While determinants of disease severity and duration may be impacted by patient’s age and underlying health conditions, there is also growing evidence that the adaptive humoral immune response itself may correlate with COVID-19 disease severity. Patients with severe COVID-19 symptoms have been shown to develop earlier and higher concentrations of SARS-CoV-2 specific IgG as compared to patients with mild symptoms (3–5).

The SARS-CoV-2 spike (S) protein is the main viral protein exposed on the virion surface and is the primary target for the immune response. The S protein mediates virus entry through the human angiotensin converting enzyme 2 (ACE2) receptor and is therefore responsible for directing the host range and tissue tropism (6, 7). The coronavirus S protein is a class I fusion protein that is composed of two subunits; the S1 head subunit mediates cellular attachment, and the S2 stalk subunit mediates membrane fusion (8). Cleavage of the S protein by host proteases separates the S1 and S2 subunits, which remain non-covalently associated in a metastable, homotrimeric prefusion formation. The S1 subunit contains two major domains, the N-terminal domain (NTD) and the C-terminal domain (CTD). The CTD includes the receptor-binding domain (RBD) positioned at the top of the trimeric S (9). Binding of SARS-CoV-2 S RBD to cellular ACE2 results in infection of pneumocytes and other host cells that express ACE2.

Antibodies against the S protein have been shown to be protective against infection and disease severity (10–12). Antibodies are a critical feature of adaptive immunity that provide protection through several mechanisms, including viral neutralization or clearance (Ab dependent cellular phagocytosis, ADCP) and elimination of infected cells (Ab-dependent cellular cytotoxicity, ADCC). Potent neutralizing monoclonal antibodies (mAbs) that target the SARS-CoV-2 S RBD and NTD can inhibit viral infection by blocking receptor engagement or by interfering with the post-attachment fusion processes (10, 13, 14). Additionally, NTD neutralizing mAbs and vaccine-elicited antibodies have been shown to leverage Fc-mediated protective activities, including monocyte, neutrophil and complement engagement (10, 13, 15).

While Fc-mediated functions can be protective, plasma IgG can also facilitate infection by promoting antibody-dependent enhancement (ADE). ADE can occur when suboptimal neutralizing or non-neutralizing antibodies bind to the virus and facilitate entry in Fc receptor expressing cells (16). Viral-induced enhanced disease has been most notably reported for Dengue virus (DENV), where pre-existing, cross-reactive Abs from one DENV serotype increase infection and clinical severity of another DENV serotype or the related Zika virus (17). ADE expands the infectable host cell range and increases the cellular viral load. Binding of virus-antibody complexes to FcγRII on immune cells initiates receptor-mediated signaling for cellular activation and enhanced inflammatory cytokine expression. In vitro ADE, as defined by Fc-mediated viral entry (FVE), has been reported for many viruses, including SARS-CoV and MERS, and in some cases has been shown to impact disease severity in humans or animal models (16, 18–20).

Enhanced viral entry of infecting SARS-CoV-2 virions by convalescent plasma has been observed in vitro (21, 22) using the Washington or Wuhan viral strains from very early in the pandemic. In this study, we evaluated the impact of SARS-CoV-2 convalescent plasma antibodies on in vitro viral entry of SARS-CoV-2 pseudoviruses in ACE2 or FcγRIIa-expressing target cells. We determined plasma neutralization and FVE activity against the early SARS-CoV-2 Washington strain, as well as the variants of concern (VOC) Alpha (B.1.1.7), Beta (B.1.351), Gamma (P.1) and Delta (B.1.617.2). Additionally, we evaluated plasma antibody binding features that associate with in vitro plasma neutralization and/or FVE. Convalescent plasma FVE activity against the infecting strain was observed in 31% of participants and was significantly associated with lower S-specific IgM binding. The plasma dilution at which maximum FVE activity was observed correlated significantly with neutralization titer, indicating that as antibody concentration decreases and neutralization wanes, Fc-mediated uptake peaks. Plasma neutralization of VOC was significantly lower compared to neutralization of the Washington strain, while FVE against the VOC was higher or similar. These data indicate that increasing neutralization-resistance of VOC may facilitate an uptick in viral entry in the absence of neutralizing plasma activity.

RESULTS

Production and Characterization of FcγRIIa- and ACE2-Expressing Target Cell Lines

Given the previous observations of FcγRIIa-expressing entry of SARS-CoV (18), we transduced HEK293 cells with an FcγRIIa expression vector to produce a stable FcγRIIa expressing cell line, and then utilized it to evaluate FVE of SARS-CoV-2 pseudoviruses (PSVs). For comparison with FVE, we utilized a commercially available HEK293 ACE2-expressing cell line to evaluate ACE2-mediated PSV entry and neutralization. Cell surface expression of FcγRII and ACE2 receptors was evaluated by flow cytometry for both cell lines and the parental HEK293 cells (Figures 1A, B). We observed low ACE2 surface expression (3.4% positive) for the HEK293_ACE2 cell line, while the FcγRIIa-expressing cell line was about 65% positive for FcγRII.

We confirmed the former expresses ACE2 by western blot (Figure 1C). The 110 kDa ACE2 protein and an ~80 kDa cleavage product (lane 4, red infrared (IR) bands) were detectable for only the HEK293_ACE2 cell line. Clathrin, a housekeeping gene, was used as a control for each cell line (green IR bands).
We then evaluated susceptibility of the target cell lines to viral entry by five SARS-CoV-2 S variants, including the original US variant Washington, and the Alpha (B.1.1.7), Beta (B.1.351), Gamma (P.1) and Delta (B.1.617.2) VOC. To establish baseline viral entry and to compare infectivity of these PSV strains, we performed viral titrations on the HEK293_ACE2 and HEK293_FcγRIIa cell lines (Figures 1D, E). Viral input was quantified and standardized by determining the pseudotyped HIV p24 capsid concentration of each viral stock. For both target cell lines we observed greater infectivity with the VOC as compared to the Washington strain (grey), which was approximately one log less infectious than the Beta variant (red).

**FcγRIIa and ACE2-Mediated Entry of SARS-CoV-2 Pseudovirus in the Presence of Convalescent Plasma**

We performed viral entry assays in the presence of convalescent (n=39) or healthy donor (n=16) plasma to observe the impact of plasma antibody on FcγRIIa or ACE2-mediated entry of the SARS-CoV-2 Washington strain PSV. All convalescent patients reported mild symptoms; plasma was not available from severe infection or asymptomatic individuals. Fold increase in viral entry was quantified and compared to the Washington strain (grey), which was approximately one log less infectious than the Beta variant (red). The black dotted line indicates 50% neutralization, and the green dotted line indicates the baseline level of viral entry in the absence of plasma in the HEK293_FcγRIIa cells. Three representative samples showed a 5 to 10-fold maximum increase in entry in the presence of plasma (FVE+; PTs 2, 4, and 15), while two plasmas showed minimal FcγRIIa-mediated increase in viral entry (FVE-; PTs 10 and 12). All five represented samples showed >50% neutralization for at least 2 dilutions. Three endpoint characteristics were determined to define FVE and quantify FcR-mediated entry into HEK293_FcγRIIa cells in the presence of antibody. These endpoint characteristics included area under the curve >1 (FVE AUC), maximum fold increase in viral entry (max FVE), and the plasma dilution at which maximum viral entry was observed. Plasma-mediated neutralization on HEK293_ACE2 cells was reported as the 50% inhibitory dilution (ID50).

We observed a significant relationship between FVE AUC and max FVE for the convalescent plasma, as expected (Figure 2G, p<0.0001, r=0.9599). We compared convalescent plasma to healthy donor plasma activities to define viral entry greater than 4-fold above the level of viral entry seen without plasma, and an AUC greater than 3 (Figure 2G; FVE+ quadrant shown in light green). By these criteria, all healthy donor plasma were FVE+ and FVE was maintained over multiple dilutions in samples defined as FVE+. Using these FVE cutoff criteria, we observed that convalescent plasma from 12 individuals (31%, green) mediated FVE and neutralization, 24 individuals (61%, black) mediated neutralization but not FVE, and 3 individuals (8%, grey) didn’t mediate FVE or neutralization of the SARS-CoV-2 virus.
Washington strain (Figure 2H). Individuals in these functional plasma activity groups were not distinguished by available clinical or demographic data. Plasma from all healthy donors did not mediate FVE or neutralization (Figure 2F and data not shown).

No correlation was observed between FVE AUC or max FVE and the dilution at which the maximum viral entry was observed (p=0.2348, r=-0.1948, and p=0.3172, r=-0.1644 respectively; data not shown). However, we observed a significant correlation between the plasma dilution at which max FVE was observed and the plasma neutralization ID50 value (Figure 2I; p=0.0001, r=0.5712), indicating an intersection between the decline of plasma antibody neutralizing capacity and the peak FVE activity.

**FcγRIIa and ACE2-Mediated Entry of SARS-CoV-2 VOC Pseudoviruses in the Presence of Convalescent Plasma**

We next analyzed a subset of 12 convalescent plasma samples (3 plasmas mediated neutralization but not FVE, and 9 plasmas mediated both neutralization and FVE against the Washington strain) and 16 healthy plasma samples to evaluate the impact of plasma antibody on FcR or ACE2-mediated entry of the SARS-CoV-2 Alpha (B.1.1.7), Beta (B.1.351), Gamma (P.1) and Delta (B.1.617.2) VOC PSVs. The VOC are shown in consistent colors throughout the figures: Alpha = blue, Delta = orange, Beta = red, and Gamma = purple. The same definition of FVE was used to compare viral entry between convalescent and healthy donor plasma for each VOC (Figures 3A–D).

A significant reduction in plasma neutralization was observed against all VOC, as compared to the Washington strain; results are shown against each strain in aggregate by all plasmas in Figure 3E. While all 12 of the convalescent plasma samples neutralized the Washington strain, only 3 or 4 convalescent plasma samples neutralized the Gamma (P.1) or Beta (B.1.351) VOC, respectively. In the radar plot shown in Figure 3F, the color-coded lines indicate the ID_{50} for each convalescent participant (PT) plasma against a given strain. PT plasma number is indicated at the perimeter, and the closer the point is to the perimeter, the greater the plasma neutralization is for that PT. The grey line indicates the higher level of neutralization of the Washington strain by each PT, as compared to the lower ID_{50}s for all other...
variants, likely due to the fact these individuals were infected with the Washington strain. There was no significant correlation between plasma neutralization activity against the Washington strain and any of the VOC (data not shown). Conversely, we observed a significant increase in plasma-mediated viral entry against the Alpha (B.1.1.7) variant, while the FVE of the other VOC was comparable to that of the Washington strain (Figure 3G). Notably, Figure 3H demonstrates that two PT samples mediated higher FVE for the Alpha and/or Delta variants as compared to the Washington variant.

Within this subset of 12 convalescent plasma samples selected for analysis against the VOC, there was an expansion of the individuals that demonstrated FVE, but no neutralization, as can be seen in the lighter green bars in Figure 3I. A shift in plasma activity from antibody-dependent viral entry and neutralization against the Washington strain to antibody-dependent viral entry without neutralization against the Beta (B.1.351) and Gamma (P.1) VOC was specifically observed (Figure 3I).

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### Convalescent Plasma Antibody Specificity and FcR Binding

Plasma binding antibody responses in all 39 convalescent plasma were evaluated by Luminex multiplex assay using Washington strain antigens to determine if plasma antibody specificity, subclass or Fc reactivity impacted plasma neutralization and/or FVE. We evaluated the correlation between convalescent plasma neutralization (Figure 4A, left) or plasma FVE (Figure 4A, right) against the Washington strain, with the magnitude of antigen (shown in rows) and Fc-specific (shown in columns) binding antibody responses. Analysis of Fc-reactivity included characterization of Ig isotype, including IgM and IgG, or binding to common Fcγ receptors, including FcγRIa, FcγRIIb, FcγRIIa and FcγRIIb. The magnitude of binding antibodies directed to SARS-CoV-2 antigens correlated directly with plasma neutralization for all Fc detectors, except for FcγRIIb-specific binding to S NTD antigen (p=0.7385, r -0.058; peach box in Figure 4A left Neutralization panel). Interestingly, significant
inverse correlations were observed between plasma FVE AUC and the magnitude of IgM binding antibodies to S1, S ECD, S trimer and S RBD antigens only (p=0.014 to p=0.038, shown in bold text). The magnitude of antigen specific binding activity for total IgG or total IgM was compared between individuals that mediated neutralization and FVE versus individuals that mediated neutralization but not FVE against the Washington strain (Figures 4B, C, respectively). Total IgG binding was similar between groups. We observed a higher magnitude of S protein specific IgM responses for individuals that did not mediate FVE as compared to individuals with FVE; significant differences were observed for S1 (p=0.0219), S ECD (p=0.0317), S trimer (p=0.0411) and S RBD (p=0.0495) antigens, as indicated. We further compared time between participant’s onset of symptoms and plasma collection as it may impact antigen-specific IgG and IgM levels and observed no differences between groups (Neut+/FVE+: mean 32 days, range 22 – 41 days; Neut+/FVE-: mean 30 days, range 22 – 47 days).

Additionally, for the 12 convalescent plasma samples tested, we evaluated the correlation between neutralization or FVE against the Washington strain and antibody characteristics (binding to Washington strain antigens and Ig or Fc receptor engagement; Table S1 and Figures 5A–D). We observed significant direct correlations between plasma neutralization of all VOC and multiple S antigen binding antibody responses. Plasma FVE responses also correlated significantly with S antigen binding antibody responses for the Alpha (B.1.1.7) and Delta (B.1.617.2) VOC (Figures 5A, B, Table S1), but not for the Beta (B.1.351) and Gamma (P.1) VOC (Figures 5C, D, Table S1). The relationship between plasma FVE, neutralization, and IgG binding to the Washington strain S trimer, shown in Figure 5, demonstrates similar trends between the Alpha (B.1.1.7) and Delta (B.1.617.2) VOC, and separately the (B.1.351) and Gamma (P.1) VOC.

As we had previously observed an inverse correlation between plasma binding antibody titers and FVE against the Washington strain for all 39 patients, we re-evaluated this relationship for this subset of 12 individuals. For this subset of patients, which primarily included patients that mediated FVE, we observe a positive correlation between binding antibody titers and FVE of the Washington strain, like what was observed for the VOC (Table S1, bottom). Interestingly, for this subset of patients, we observe significant direct correlations between FVE and binding antibody responses to only the S trimer and S2 ECD. However, only 12 individuals were included in this analysis and further work is needed to clarify these trends.

**DISCUSSION**

Over 513 million COVID-19 cases have been confirmed worldwide, as of 1 May 2022. While pulmonary impairment is
the most common symptom of infection, concurrent extra-
pulmonary manifestations and post-COVID conditions are also prevalent and diverse (23). Due to the recency of the SARS-CoV-2 pandemic, the long-term consequences of COVID-19 infection won’t be fully characterized for some time.

Persistence of symptoms beyond the initial SARS-CoV-2 infection (4 or more weeks) implicates the adaptive immune response in propagating or exacerbating long-term sequelae (24). In this study, we evaluated humoral immune responses from convalescent individuals to determine the impact of their plasma antibodies on inhibiting or mediating SARS-CoV-2 entry in ACE2- or FcR-expressing cells, respectively. We determined that convalescent plasma antibodies could both inhibit and mediate viral entry of the infecting Washington SARS-CoV-2 strain. Plasma FVE against the Washington strain was observed for 31% of convalescent individuals and was detected only in the presence of neutralizing antibodies. For these individuals, viral entry and neutralization activities correlated inversely across multiple plasma dilutions, in that FVE activity peaked when neutralizing activity was declining. Additionally, significantly lower S protein specific plasma IgM was observed in individuals that mediated FVE, indicating IgM may potentially compete with FVE-mediated S-specific IgG for S antigen binding. Previous reports have shown the protective effect of S-specific IgM (25, 26). Infected individuals that developed earlier and higher IgM responses recovered more quickly, while higher IgG levels during recovery correlated with disease severity (27, 28).

Extrinsic ADE can occur through FcR endocytosis of IgG and viral immune complexes. In vitro ADE has been documented for multiple viruses, including HIV (29), influenza (30), Ebola (31) and flaviviruses (32–34). ADE was first and most notably described for DENV (35, 36); pre-existing antibodies can enhance infectivity and virulence of a secondary infection, increasing likelihood of Dengue hemorrhagic fever and dengue shock syndrome (37, 38). ADE occurs when non-neutralizing or sub-neutralizing concentrations of antibody bind to viral antigens without inhibiting or clearing infection. This cross-reactive but reduced-affinity antibody binding has been observed across DENV serotypes and between flaviviruses (39–41). In this study, we similarly observed Fc-mediated entry coinciding with declining or sub-neutralizing plasma antibody dilutions but have refrained from using the term ADE in reference to this activity as it is unclear whether true “enhancement” is occurring in this in vitro setting. Thus, we have preferred to define the activity simply as FcγR-mediated virus entry, or FVE. The role of IgG Fc receptors in potential ADE of SARS-CoV-2 infection is still unclear. However, as neutralization-resistant SARS-CoV-2 VOC emerge, further understanding of plasma antibody cross-reactivity and potential ADE are needed (42).

Several mechanisms of coronavirus FVE have been reported. RBD mAbs have been shown to trigger conformational changes in MERS or SARS-CoV S, facilitating entry into Fc receptor expressing cells through receptor mimicry (18, 43). Additionally, infection enhancing SARS-CoV-2 NTD mAbs, isolated from a patient with severe COVID-19, have been shown to induce conformational changes in S that promote ACE2 binding and fusion (44). In this study, we evaluated Fc-mediated viral entry utilizing FcγRIIa, as this Fc receptor was previously shown to mediate viral entry of SARS-CoV (18). Another study recently observed FcγRIIb-mediated enhancement of viral infection with RBD-specific SARS-CoV-2 neutralizing antibodies, while NTD-specific non-neutralizing antibodies were shown to mediate enhancement through a FcγR-independent mechanism (45). In this work, these infection-enhancing antibodies were protective in vivo when infused in monkeys and mice, however lung inflammation was observed in a small number of infused animals, as compared to untreated infected controls (45). Adding further layers of complexity to the potential FVE mechanisms, alternative receptors have been demonstrated to mediate or facilitate SARS-CoV-2 entry. Cellular molecules, including KREMEN1, ASGR1, and CD147 have been shown to mediate viral uptake (46, 47), while infection enhancing molecules, such as the phosphatidylycerine receptor, facilitate viral internalization in an ACE2-dependent manner (48, 49). These studies demonstrate the diverse and complex cellular entry mechanisms of SARS-CoV-2 (50). Further analysis is needed to explore the potential mechanism of convalescent plasma antibody FVE observed in this study.

Here we evaluated the impact of convalescent plasma from Washington strain SARS-CoV-2 infection on viral entry of SARS-CoV-2 VOC, including Alpha (B.1.1.7), Beta (B.1.351),
Gamma (P.1) and Delta (B.1.617.2). Convalescent plasma had reduced neutralization potency against all tested VOC; however, the reduction was most notable for the Beta (B.1.351) and Gamma (P.1) VOC. This is consistent with previous reports showing increased neutralization resistance of Beta (B.1.351) and Gamma (P.1) as compared to other variants, due to the shared E484K mutation in RBD (51–54). Amino acid mutations, S characteristics and monoclonal antibody neutralization sensitivity are also more similar between the Beta (B.1.351) and Gamma (P.1), as compared to Alpha (B.1.1.7) or Delta (B.1.617.2) VOC (51, 55). Similarly, we observe consistent patterns in plasma reactivity against the Beta (B.1.351) and Gamma (P.1) versus the Alpha (B.1.1.7) and Delta (B.1.617.2) VOC. FVE against the VOC was similar to or higher than the Washington strain. Thus, when plasma neutralization capacity declined significantly, plasma antibodies were found to mediate viral entry in the absence of neutralization activity. Further studies are needed to evaluate the impact of convalescent antibodies from currently circulating SARS-CoV-2 infections on FVE of emerging SARS-CoV-2 VOC that may be more neutralization-resistant.

The recently emerged Omicron (B.1.1.529) VOC contains a significantly larger number of mutations that result in neutralization resistance against many therapeutic and convalescent plasma antibodies (56, 57). Increased transmissibility of VOC has also been reported; Alpha (B.1.1.7) VOC was determined to be 43-90% more transmissible than the Washington strain, Delta (B.1.617.2) VOC was determined to be 60% more transmissible than Alpha (B.1.1.7) VOC, and Omicron (B.1.1.529) VOC was determined to be 100% more transmissible than Delta (B.1.617.2) VOC (57–59). Further, population-level analysis has identified a significant increase in reinfection risk associated with Omicron (B.1.1.529), but not Beta (B.1.351) or Delta (B.1.617.2) VOC (60). The implications of diminished reactivity of pre-existing antibodies in Omicron (B.1.1.529) reinfection have yet to be determined. However, while Omicron (B.1.1.529) is highly transmissible, disease severity of infection or reinfection appear to be diminished, with a significantly reduced risk of COVID-19 hospitalization as compared to Delta (B.1.617.2) VOC infection (61, 62).

The balance of antibody functions in vivo is complicated and cannot be reduced to functional activity observed in isolation in vitro. Neutralization and Fc-mediated antibody functions have a demonstrated role in prevention and control of SARS-CoV-2 (54, 63–65). A single antibody can have multifunctional capacity (66) by mediating both neutralization and ADCC, and similarly could mediate both ADCC and FVE, depending on environmental conditions. Passive immunotherapies have been developed for treatment and protection against emerging SARS-CoV-2 variants, recently reviewed in detail (67). Among those treatments, convalescent plasma therapies have been evaluated with mixed outcome, but are generally determined to be safe. Convalescent plasma therapy treatment effectiveness is impacted by the quality, titer and timing of administered antibodies (68–71). Qualities of the humoral immune response to SARS-CoV-2 have been shown to vary with disease severity (72–76). Patients with severe COVID-19 have unique serological signatures, including afucosylated Fc glycans which have enhanced interactions with FcγRIIa and result in proinflammatory activity (73, 75). Our study further demonstrates varying levels of convalescent plasma antibody Fc-engagement as determined by FVE. However, we have yet to explore the potential of this antibody Fc-engagement to result in protective Fc-mediated antibody functions, such as ADCC or ADCP. Further, convalescent antibody activities have been shown to differentially wane over time, with neutralization potency declining more rapidly than Fc-mediated ADCC and ADCP (77). The dynamic nature of the SARS-CoV-2 humoral immune response and antibody functions may shift over time in the context of this evolving COVID-19 pandemic.

These experiments employed cell line models transduced to express the ACE2 or FcγRIIa receptors, and therefore receptor expression may vary from in vivo target cells. The antibody functions against SARS-CoV-2 described herein have not yet been well characterized using primary viral isolates or using lung cells or macrophages. The impact of antibody on viral entry in primary cells co-expressing ACE2 and FcγRIIa receptors is unknown (as illustrated in Figure S1 as a graphical abstract). Viral entry and cytokine expression, which was not measured in this study, may vary between cell types and environmental conditions. The downstream cellular consequences of FVE observed in this study were thus not analyzed as they may not be physiologically relevant and could therefore lead to misleading conclusions. Studies using the challenging primary cell culture systems and wild type virus should shed further light on the in vivo importance of the functional activities observed in this study. While neither medical encounter, nor follow up data were available for the individual samples accessible for this study, analysis of longitudinal plasma samples from patients with varying clinical severity, and/or known reinfection status will help elucidate the relationship between in vitro Fc-mediated viral entry versus neutralization and in vivo disease progression. Despite these limitations, the data presented here suggest a relationship between S protein antibody binding titers and antibody functions that mediate virus neutralization or Fc-mediated virus entry. These observations warrant further investigation using well characterized, curated samples and primary viral and cell systems to further elucidate the impact of specific antibody functions on the pathogenesis of SARS-CoV-2 and COVID-19 disease.

MATERIALS AND METHODS

SARS-CoV-2 Convalescent Plasma

Convalescent plasma were purchased from Seracare Life Sciences (Milford, MA) and BEI Resources (Manassas, VA) and were collected from individuals infected between February and April 2020 in the SARS-CoV-2 pandemic. Healthy donor plasma samples were from the WRAIR IRB-approved RV229 protocol...
and were collected from individuals prior to the SARS-CoV-2 pandemic.

**Lentiviral Particle Transduction and Culture of Cell Lines**

Lentiviral particles were produced by co-transfecting HEK293T/17 cells (ATCC, Manassas, VA) with a Lenti-Pac HIV expression packaging kit (GeneCopoeia, Rockville, MD) and an ORF expression clone for FCGR2A (GenBank Accession: NM_021642.3) with neomycin resistance (GeneCopoeia, Rockville, MD) according to the manufacturer’s instructions. Lentiviral particle p24 concentration was determined by HIV-1 p24 antigen capture kit (Advanced Bioscience Laboratories, Rockville, MD) and HEK293 cells (ATCC, Manassas, VA) were transduced with an MOI of 10. FcγRIIa expressing cells were selected for using 1 mg/mL of G418 antibiotic in the growth medium. ACE2-expressing HEK293 cells were commercially obtained (Integral Molecular, Philadelphia, PA) and cultured with 1 µg/mL of puromycin in the growth medium.

**Characterization of Target Cells by Flow Cytometry**

Cells were detached using non-enzymatic cell dissociation buffer (Millipore Sigma, Burlington, MA) and 2.5 x 10^5 cells per well were added to a 96-well plate. Cells were washed with PBS and then stained with 50 µL of LIVE/DEAD fixable violet stain kit (Thermo Fisher Scientific, Waltham, MA) according to the manufacturer’s recommendations. After washing off excess viability stain, the cells were stained extracellularly using antibodies specific for ACE2 (R&D Systems, Minneapolis, MN) or FcγRIIa (BD Biosciences, Franklin Lakes, NJ) for 30 mins at 4°C. The cells were then washed and fixed with 2% formaldehyde (Tousimis, Rockville, MD). Data was recorded using the Becton FACSymphony (BD Biosciences, Franklin Lakes, NJ) and post-acquisition analysis was performed with FlowJo 10.7.1 (FlowJo, Ashland, OR).

**Characterization of Target Cells by Western Blot**

Cells were lysed, treated with Mammalian Protein Extraction Reagent (Thermo Fisher Scientific, Waltham, MA) containing protease inhibitors (Roche, Basel, Switzerland), including phenylmethylsulfonyl fluoride (Sigma-Aldrich, St. Louis, MO) and shaken at 4°C. Extracts were centrifuged at 14,000 G at 4°C for 10 minutes, then supernatants were collected, and protein concentrations were determined using Pierce bicinchoninic acid (BCA) protein assay (Thermo Fisher Scientific, Waltham, MA). The S expression plasmid sequence was derived from the Wuhan Hu-1 strain (GenBank # NC_045512), which is also identical to the IL1/2020 and WA1/2020 strains, and was codon optimized and modified to remove the last 18 amino acids of the cytoplasmic tail to improve S incorporation into the PSV, thereby enhancing infectivity. S expression plasmids for SARS-CoV-2 VOC were produced similarly and included S sequences for Alpha (B.1.1.7), Beta (B.1.351), Gamma (P.1), and Delta (B.1.617). Concentration of p24 in PSV stocks was determined by HIV-1 p24 antigen capture kit (Advanced Bioscience Laboratories, Rockville, MD) according to the manufacturer’s protocol. To determine infectivity, PSVs were serially diluted and 25 µL/well was added to a white 96-well plate with an equal volume of growth medium. Target cells were added to each well (40,000 cells/well) and incubated at 37°C for an additional 48 hr. Bright-Glo Luciferase Assay System substrate was added and Relative Light Units (RLUs) were measured using the EnVision Multimode Plate Reader (Perkin Elmer, Waltham, MA).

**SARS-CoV-2 Pseudovirus Neutralization and FVE Assays**

Test plasma were diluted 1:25 in growth medium, serially diluted, then 25 µL/well was added to a white 96-well plate. An equal volume of diluted SARS-CoV-2 PSV was added to each well and plates were incubated for 1 hr. at 37°C. HEK293 target cells expressing ACE2 or FcγRIIa were added to each well (40,000 cells/well) for neutralization or FVE assays, respectively. Plates were incubated for 48 hr. at 37°C. Then Bright-Glo Luciferase Assay System substrate (Promega, Madison, WI) was added and RLUs were measured using the EnVision Multimode Plate Reader (Perkin Elmer, Waltham, MA). Neutralization dose–response curves were fitted by nonlinear regression and final titers were reported as the reciprocal of the dilution of plasma necessary to inhibit 50% of viral entry (ID_{50}). FVE was calculated as the fold increase in viral entry in the presence of plasma; curves were analyzed using GraphPad Prism to determine the area under the curve >1 (FVE AUC), maximum fold increase in viral entry (max FVE), and the plasma dilution at which maximum viral entry was observed.
SARS-CoV-2 Antigens

Seven SARS-CoV-2 antigens were used to profile antibody responses, including S extra-cellular domain (S ECD), which comprises the subunit 1 (S1) and 2 (S2) (S1+S2 ECD), S1, S2, the nucleoprotein (NP) (Sino Biological; 40589-V08B1, 40591-V08B1, 40590-V08B and 40588-V08B), the S Receptor Binding Domain (RBD), S trimer (LakePharma; 46438 and 46328) and S N Terminal Domain [NTD; produced in house (78)]. All antigens corresponded to the SARS-CoV-2 Wuhan-Hu-1 strain.

Luminex Multiplex Binding Antibody Assay

Bead-coupling and multiplex Luminex assays were performed as previously described, with modifications (79). Thirty µg of each S antigen and 15µg of NP antigen were covalently coupled to six million fluorescently coded carboxylated magnetic MagPlex beads (Luminex Corporation, Austin, TX). Samples were tested in triplicate at a dilution of 1:400; 1,200 antigen-conjugated beads for each antigen were used per well. Total IgG and IgM antibody responses were detected with 20µl of phycoerythrin (PE)-labeled secondary antibodies specific for total IgG and IgM (Southern BioTech; Birmingham, AL) at a final concentration of 3µg/ml. Prior to use, biotinylated proteins corresponding to Fc gamma receptors FcyR2a, FcyR2b, FcyR3a and FcyR3b (produced under the direction of Dr. James Peacock in the Duke Human Vaccine Institute Research Protein Production Facility which received funding support from the Collaboration for AIDS Vaccine Research Bill and Melinda Gates Foundation (OPP1066832)) were mixed with a 1/4th molar ratio of Streptavidin-R-Phycocerythrin (SAPE; Prozyme) and incubated with rotation for 30 min at room temperature. One volume percent of 500µM free biotin (Thermo Fisher Scientific, Waltham, MA) was then added to block any free streptavidin binding sites and the SAPE-FcyRs were stored at 4°C and used within 24 hours. Binding to FcyRs was detected with 20µl of SAPE-FcyR2a, -FcyR2b, -FcyR3a and -FcyR3b at a final concentration of 1µg/ml.

Magnetic beads were acquired on a FlexMap 3D instrument (Luminex Corporation, Austin, TX) using the xPONENT software (version 4.2). A minimum of 100 beads per antigen and per well were collected. The output of the assay is the median fluorescence intensity (MFI) determined from the sampled beads. One single plate was run for each detection antibody and FcγR. Two pre-pandemic plasma samples from uninfected individuals were used as negative controls and two plasma samples from SARS-CoV-2 convalescent individuals were used as positive controls, each ran in triplicate at a 1:400 dilution. Each set of replicates was evaluated for outlier MFI values. The coefficient of variation (CV) of the triplicates was calculated; when the CV was higher than 20%, the replicate with the MFI signal furthest from the mean MFI was excluded. The mean MFI was then calculated for each sample.

Data Analysis and Statistics

Multiple variable correlation analyses were performed using nonparametric Spearman correlation. Statistical differences between groups were determined using Wilcoxon matched pairs signed rank tests and Mann Whitney U tests.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by Institutional Review Board at Walter Reed Army Institute of Research. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

LW, MZ, KP, JD, SP, and VP designed the studies. SP and VP provided project oversight. VD and SK contributed plasma and constructed SARS-CoV-2 S plasmids. MZ, SM, and EK developed and characterized the cell lines. MZ, EK, SM, and JH developed and performed SARS-CoV-2 infectivity, neutralization and FVE assays. MM, BB, SW-R, and MR performed SARS-CoV-2 Luminex antibody profiling. LW and MZ conducted data analysis. LW, MZ and VP wrote the manuscript. MM, WD, SW-R, KP, JD, SK, MR and SP reviewed and edited the manuscript. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2022.901217/full#supplementary-material
**Supplementary Figure 1** | Graphical summary of the route of viral entry (top) and the impact of antibody on viral entry (bottom) for (A) the HEK293, Fc-fills cell line, (B) the HEK293, ACE2 cell line and (C) primary cells, including monocytes and macrophages. The image was created using software available on BioRender.com.

**REFERENCES**

1. Guan WJ, Ni ZY, Hu Y, Liang WH, Ou CQ, He JX, et al. Clinical Characteristics of Coronavirus Disease 2019 in China. *N Engl J Med* (2020) 382:1708–20. doi: 10.1056/NEJMc2002302

2. Rieck DO, Malone RW. Medical Countermeasures Analysis of 2019-nCoV and Vaccine Risks for Antibody-Dependent Enhancement (ADE). *Preprints* (2020) 2020030138. doi: 10.20944/preprints202003.0138.v1

3. Marklund L, Nilsson J, Zurbuchen Y, Valaperti A, Schreiner J, Wolfensberger A, et al. Systemic and Mucosal Antibody Responses Specific to SARS-CoV-2 During Mild Versus Severe COVID-19. *J Allergy Clin Immunol* (2021) 147 (2):545–57.e549. doi: 10.1016/j.jac.2020.01.040

4. Liu W, Moore MF, Vasilieva N, Sui J, Wong SK, Berne MA, et al. Angiotensin-Converting Enzyme 2 is a Functional Receptor for the SARS Coronavirus. *Nature* (2003) 426(6965):450–4. doi: 10.1038/nature02145

5. Hamming I, Timens W, Bulbuis ML, Lely AT, Navis G, van Goor H. Tissue Distribution of ACE2 Protein, the Functional Receptor for SARS Coronavirus. A First Step in Understanding SARS Pathogenesis. *J Pathol* (2004) 203(2):631–7. doi: 10.1002/path.1570

6. Benton DJ, Wrobel AG, Xu P, Roustan C, Martin SR, Rosenthal PB, et al. Receptor Binding and Priming of the Spike Protein of SARS-CoV-2 for Membrane Fusion. *Nature* (2020) 588(7837):327–30. doi: 10.1038/s41586-020-2772-0

7. Yuan Y, Cao D, Zhang Y, Ma J, Qi J, Wang Q, et al. Cryo-EM Structures of Human Monoclonal Antibody. *Cell* (2021) 184(9):2316–31.e2315. doi: 10.1016/j.cell.2021.03.029

8. Roozendaal R, Solforosi L, Stieh DJ, Serroyen J, Straetemans R, Dari A, et al. SARS-CoV-2 Binding and Neutralizing Antibody Levels After Ad26.COV2.S Vaccination Predict Durable Protection in Rhesus Macaques. *Science* (2021) 371(6531):823–9. doi: 10.1126/science.abf4830

9. Gorman MJ, Patel N, Guebre-Xabier M, Zhu AL, Atyeo C, Pullen KM, et al. Fab and Fc Contribute to Maximal Protection Against SARS-CoV-2 Following NVX-CoV2373 Subunit Vaccine With Matrix-M Vaccination. *Cell Rep Med* (2021) 2(9):100405. doi: 10.1016/j.xcrm.2021.100405

10. Smatti MK, Al Thani AA, Yasmine HM. Viral-Induced Enhanced Disease Illnesses. *Front Microbiol* (2018) 9:2991. doi: 10.3389/fmicb.2018.02991

11. Khanda R, Munjal A, Dhamla K, Karthik K, Tiwari R, Malik YS, et al. Modulation of Dengue/Zika Virus Pathogenicity by Antibody-Dependent Enhancement and Strategies to Protect Against Enhancement in Zika Virus Infection. *Front Immunol* (2018) 9:597. doi: 10.3389/fimmu.2018.00597

12. Houser KV, Broadbent AJ, Gretebeck L, Vogel L, Lamirande EW, Sutton T, et al. Enhanced Inflammation in New Zealand White Rabbits When MERS-CoV Reinfection Occurs in the Absence of Neutralizing Antibody. *PLoS Pathog* (2017) 13(8):e1006565. doi: 10.1371/journal.ppat.1006565

13. Ye J, Yang JY, Liu WT, Huang JC, Chen YMA, Wang SF. SARS Coronavirus has Antibody-Dependent Enhancement (ADE) Effect Through the Autologous Antibodies Against Envelope Spikes on Fc Receptor Expressing Cells. *J Virus Erad* (2016) 2:48. doi: 10.1016/S2055-6640(20)31216-4

14. Hu T, Kuroda M, Armbrust T, Yamasaki S, Halfmann P, et al. Antibody-Dependent Enhancement of Viral Infectivity. *Cell Rep Med* (2021) 2(9):100405. doi: 10.1016/j.xcrm.2021.100405

15. Ochiai H, Kurokawa M, Matsu S, Yamamoto T, Kuroki Y, Kishimoto C, et al. Antibody-Dependent Enhancement of SARS-CoV-2 Virus Infection Is Mediated by the IgG Receptors FcγRIIa and FcγRIIb But Does Not Contribute to Aberrant Cytokine Production by Macrophages. *mBio* (2021) 2021:e01987-21

16. Wu F, Yan R, Liu M, Liu Z, Wang Y, Luan D, et al. Antibody-Dependent Enhancement (ADE) of SARS-CoV-2 Infection in Recovered COVID-19 Patients: Studies Based on Cellular and Structural Biology Analysis. *medRxiv* (2020). doi: 10.1101/2020.08.20.20209114

17. Higgins V, Sohaei D, Diamandis EP, Prassas I. COVID-19: From an Acute to Chronic Disease? Potential Long-Term Health Consequences. *Crit Rev Clin Lab Sci* (2021) 58(5):297–310. doi: 10.1080/01463670.2021.1860895

18. Roozendaal R, Solforosi L, Stieh DJ, Serroyen J, Straetemans R, Dari A, et al. SARS-CoV-2 Binding and Neutralizing Antibody Levels After Ad26.COV2.S Vaccination Predict Durable Protection in Rhesus Macaques. *Nat Commun* (2017) 8:15092. doi: 10.1038/ncomms15092
36. Halstead SB, Shotwell H, Casals J. Studies on the Pathogenesis of Dengue Infection in Monkeys. II. Clinical Laboratory Responses to Heterologous Infection. J Infect Dis (1973) 128(1):15–22. doi:10.1093/infdis/128.1.15

37. Halstead SB, Nirmannmiya S, Cohen SN. Observations Related to Pathogenesis of Dengue Hemorrhagic Fever. IV. Relation of Disease Severity to Antibody Response and Virus Recovered. Yale J Biol Med (1970) 42(5):311–28.

38. Bhatt P, Sabeena SP, Varma M, Arunkumar G. Current Understanding of the Pathogenesis of SARS-CoV-2 Virus Infection. Curr Microbiol (2021) 78(1):17–32. doi:10.1007/s00284-020-02284-w

39. Salje H, Cummings DAT, Rodrigues-Barraca I, Katzelnick LC, Lessler J, Klungthong C, et al. Reconstruction of Antibody Dynamics and Infection Histories to Evaluate Dengue Risk. Nature (2018) 557(7707):719–23. doi:10.1038/s41586-018-0157-4

40. Izmirly AM, Alturki SO, Connors J, Haddad EK. Challenges in Dengue Vaccines Development: Pre-Existing Infections and Cross-reactivity. Front Immunol (2020) 11:1055. doi:10.3389/fimmu.2020.01055.

41. Bonheur AN, Thomas S, Soshnick SH, McGibbon E, Dupuis AP 2nd, Hull R, et al. A Fatal Case Report of Antibody-Dependent Enhancement of Dengue Virus Type 1 Following Remote Zika Virus Infection. BMC Infect Dis (2021) 21(1):749. doi:10.1186/s12879-021-04682-0

42. Chen RE, Zhang X, Case JB, Winkler ES, Liu Y, VanBlargan LA, et al. Resistance of SARS-CoV-2 Variants to Neutralization by Monoclonal and Serum-Derived Polyclonal Antibodies. Nat Med (2021) 27(4):717–26. doi:10.1038/s41591-021-02944-w

43. Walls AC, Xiong X, Park YJ, Tortorici MA, Snijder J, Quiep J, et al. Unexpected Receptor Functional Mimicry Activates Activation of Coronavirus Fusion. Cell (2019) 176(5):1026–39.e1015. doi:10.1016/j.cell.2018.12.028

44. Liu Y, Soh WT, Kishikawa JI, Hirose M, Nakayama EE, Li S, et al. An Infectivity-Enhancing Site on the SARS-CoV-2 Spike Protein Targeted by Antibodies. Cell (2021) 184(13):3452–66.e3418. doi:10.1016/j.cell.2021.05.033.

45. Li D, Edwards RJ, Manne K, Martinez DR, Schäfer A, Alam SM, et al. The Functions of SARS-CoV-2 Neutralizing and Infection-Enhancing Antibodies In Vitro and in Mice and Nonhuman Primates. bioRxiv (2021). doi:10.1101/2021.12.31.424729

46. Gu Y, Cao J, Zhang X, Gao H, Wang Y, Wang J, et al. Receptor Profiling Identifies KREMEN1 and ASGR1 as Alternative Functional Receptors for SARS-CoV-2. Cell Res (2022) 2022:1–14. doi:10.1038/s41467-022-00654-6

47. Wang K, Chen W, Zhang Z, Deng Y, Lian JQ, Du P, et al. CD147-Spike Protein is a Novel Route for SARS-CoV-2 Infection to Host Cells. Signal Transduct Target Tther (2020) 5(1):283. doi:10.1186/s13239-020-00046-x

48. Bohan D, Van Ert H, Ruggio N, Rogers KJ, Badreddine M, Aguilar Briseño JA, et al. Phosphatidylinositol Receptors Enhance SARS-CoV-2 Infection. PLoS Pathog (2017) 13(11):e1006973. doi:10.1371/journal.ppat.1006973

49. Galadane LC, McSweeney KR, Qaradakhi T, Ali B, Zulli A, Apostolopoulos V. Live Imaging of SARS-CoV-2 Escape From a Highly Neutralizing COVID-19 Convalescent Plasma. Proc Natl Acad Sci USA (2021) 118(36). doi:10.1073/pnas.2103154118

50. Dussupt V, Sankhala RS, Mendez-Rivera L, Townsley SM, Schmidt F, Wiercokz L, et al. Low-Dose In Vivo Protection and Neutralization Across SARS-CoV-2 Variants by Monoclonal Antibody Combos. Nat Immunol (2021) 22(12):1503–14. doi:10.1038/s41590-021-01068-z

51. Hastie KM, Li H, Bedinger D, Schendel SL, Dennison SM, Li K, et al. Defining Variant-Resistant Epitopes Targeted by SARS-CoV-2 Antibodies: A Global Consortium Study. Sci (New York NY) (2021) 374(6566):472–8. doi:10.1126/science.abb2315

52. Weisblum Y, Schmidt F, Zhang F, DaSilva J, Keyt BA, Steholl LM. Passive Immunotherapy Against SARS-CoV-2: From Plasma-Based Therapy to Single Potent Antibodies in the Race to Stay Ahead of the Variants. BioDrugs (2022) 2022:1–93. doi:10.1007/s40259-022-00529-7

53. Marconato M, Abela IA, Hauser A, Schwarzmüller M, Katzeneiner R, Braun DL, et al. Antibodies From Convalescent Plasma Promote SARS-CoV-2 Clearance in Individuals With and Without Endogenous Antibody Response. J Clin Invest (2022). doi:10.1172/JCI158190

54. Orfutza MB, Yoon H, Goldfeld KS, Troxel AB, Daily JP, Wu Y, et al. Efﬁcacy and Safety of COVID-19 Convalescent Plasma in Adults With Severe COVID-19. J Clin Invest (2021) 131(13). doi:10.1172/JCI150646

55. Dispinseri S, Vecchi S, Pirillo MF, Tolazzi M, Borghi M, Eckhardt C, et al. Antibody Neutralizing Responses to SARS-CoV-2 in Symptomatic COVID-19 is Persistent and Critical for Survival. Nat Commun (2021) 12(1):2670. doi:10.1038/s41467-021-22958-8
Chakraborty S, Gonzalez J, Edwards K, Mallajosyula V, Buzzanco AS, Sherwood R, et al. Proinflammatory IgG Fc Structures in Patients With Severe COVID-19. Nat Immunol (2021) 22(1):67–73. doi: 10.1038/s41590-020-00828-7

Vigón L, García-Pérez I, Rodríguez-Mora S, Torres M, Mateos E, Castillo de la Osa M, et al. Impaired Antibody-Dependent Cellular Cytotoxicity in a Spanish Cohort of Patients With COVID-19 Admitted to the ICU. Front Immunol (2021) 12:742631. doi: 10.3389/fimmu.2021.742631

Larsen MD, de Graaf EL, Sonneveld ME, Plomp HR, Nouta J, Hoepel W, et al. Afucosylated IgG Characterizes Enveloped Viral Responses and Correlates With COVID-19 Severity. Sci (New York NY) (2021) 371(6532). doi: 10.1126/science.abc8378

Yates JL, Ehrbar DJ, Hunt DT, Girardin RC, Dupuis AP2nd, Payne AF, et al. Serological Analysis Reveals an Imbalanced IgG Subclass Composition Associated With COVID-19 Disease Severity. Cell Rep Med (2021) 2(7):100329. doi: 10.1016/j.xcrm.2021.100329

Lee WS, Selva KJ, Davis SK, Wines BD, Reynaldi A, Esterbauer R, et al. Decay of Fc-Dependent Antibody Functions After Mild to Moderate COVID-19. Cell Rep Med (2021) 2(6):100296. doi: 10.1016/j.xcrm.2021.100296

Dussupt V, Sankhala RS, Gromowski GD, Donofrio G, de la Barrera RA, Larocca RA, et al. Potent Zika and Dengue Cross-Neutralizing Antibodies Induced by Zika Vaccination in a Dengue-Experienced Donor. Nat Med (2020) 26(2):228–35. doi: 10.1038/s41591-019-0746-2

Merbah M, Wollen-Roberts S, Shubin Z, Li Y, Bai H, Dussupt V, et al. A High-Throughput Multiplex Assay to Characterize Flavivirus-Specific Immunoglobulins. J Immunol Methods (2020) 487:112874. doi: 10.1016/j.jim.2020.112874

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