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Newly emerging viruses, such as severe acute respiratory syndrome coronavirus (SARS-CoV), Middle Eastern respiratory syndrome CoVs (MERS-CoV), and H7N9, cause fatal acute lung injury (ALI) by driving hypercytokinemia and aggressive inflammation through mechanisms that remain elusive. In SARS-CoV/macaque models, we determined that anti–spike IgG (S-IgG), in productively infected lungs, causes severe ALI by skewing inflammation-resolving response. Alveolar macrophages underwent functional polarization in acutely infected macaques, demonstrating simultaneously both proinflammatory and wound-healing characteristics. The presence of S-IgG prior to viral clearance, however, abrogated wound-healing responses and promoted MCP1 and IL-8 production and proinflammatory monocyte/macrophage recruitment and accumulation. Critically, patients who eventually died of SARS (hereafter referred to as deceased patients) displayed similarly accumulated pulmonary proinflammatory, absence of wound-healing macrophages, and faster neutralizing antibody responses. Their sera enhanced SARS-CoV–induced MCP1 and IL-8 production by human monocyte–derived wound-healing macrophages, whereas blockade of FcγR reduced such effects. Our findings reveal a mechanism responsible for virus-mediated ALI, define a pathological consequence of viral specific antibody response, and provide a potential target for treatment of SARS-CoV or other virus-mediated lung injury.

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Anti–spike IgG causes severe acute lung injury by skewing macrophage responses during acute SARS-CoV infection

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Introduction

Severe acute respiratory syndrome coronavirus (SARS-CoV), Middle Eastern respiratory syndrome CoVs (MERS-CoV), and H7N9, cause fatal acute lung injury (ALI) by driving hypercytokinemia and aggressive inflammation through mechanisms that remain elusive. In SARS-CoV/macaque models, we determined that anti–spike IgG (S-IgG), in productively infected lungs, causes severe ALI by skewing inflammation-resolving response. Alveolar macrophages underwent functional polarization in acutely infected macaques, demonstrating simultaneously both proinflammatory and wound-healing characteristics. The presence of S-IgG prior to viral clearance, however, abrogated wound-healing responses and promoted MCP1 and IL-8 production and proinflammatory macrophage/macrophage recruitment and accumulation. Critically, patients who eventually died of SARS (hereafter referred to as deceased patients) displayed similarly accumulated pulmonary proinflammatory, absence of wound-healing macrophages, and faster neutralizing antibody responses. Their sera enhanced SARS-CoV–induced MCP1 and IL-8 production by human monocyte–derived wound-healing macrophages, whereas blockade of FcγR reduced such effects. Our findings reveal a mechanism responsible for virus-mediated ALI, define a pathological consequence of viral specific antibody response, and provide a potential target for treatment of SARS-CoV or other virus-mediated lung injury.
during the acute stage of ARDS, as well as in the resolution phases of inflammation and recovery from ARDS. At steady state, resident macrophages are normally quiescent to prevent damaging the alveoli and are critically involved in normal tissue homeostasis. After tissue injury or during infection, resident macrophages become activated. Circulating monocytes can be efficiently recruited to the site of injury. Inflammatory monocytes/macrophages (IMMs) and resident macrophages undergo marked phenotypic and functional changes, and they can be classified into proinflammatory (M1 or classically activated) and inflammatory-resolving (M2, alternatively activated, wound-healing, or antiinflammatory) macrophages, with a continuum of macrophage polarization existing beyond these discrete categories (9). During acute infection, monocytes/macrophages often display a phenotype of classically activated macrophages. These cells mediate host defenses against viruses and also promote lung injury by producing nitric oxide (NO); ROS; IL-1, IL-6, and IL-8; and TNF. Simultaneously, some macrophages may become alternatively activated, exerting antiinflammatory function and regulating wound healing by producing matrix metalloproteinases (MMPs), growth factors, and antiinflammatory cytokines, particularly TGF-β. When the pathogen or inflammatory stimulus is eliminated, proinflammatory macrophages diminish. The predominant macrophage population assumes a wound-healing phenotype. At the final recovery stage, macrophages show a regulatory/suppressive phenotype, secreting increased levels of IL-10, which facilitates the resolution of wound healing and restores homeostasis. When the wound-healing response is well organized and controlled, the inflammatory response resolves quickly, and normal tissue architecture is restored. Disturbances in wound-healing response can lead to uncontrolled production of inflammatory mediators, contributing to a state of persistent injury (9–11). In patients who eventually died of SARS (hereafter referred to as deceased patients) and animal models, extensive lung damage is associated with high initial viral loads, increased IMM accumulation in the lungs, and elevated serum proinflammatory cytokine (IL-1, IL-6, IL-8, CXCL-10, and MCP1) levels (4, 5, 12, 13). While much is known about the terminal phase of SARS, little is known about the early immune events during the acute phase of infection. Studies defining macrophage heterogeneity and function during acute infection and ALI using nonhuman primate and patient specimens are limited, and the factors driving the hypercytokinemia and aggressive inflammation remain elusive.

During the SARS outbreak in Hong Kong, most patients (70%–80%) presented with abnormal chest radiographs, with approximately one-quarter of these individuals progressing to ALI. After 12 days, 80% of these SARS patients developed ARDS, coincident with IgG seroconversion (6). In a detailed analysis of antibody responses against SARS-CoV spike (S) glycoprotein, we reported that the anti-S neutralizing antibody (NAb) response developed significantly faster in deceased patients compared with recovered patients after the onset of clinical symptoms (14). It took an average of 20 days for the recovered patients to reach their peak of NAb activities, as opposed to only 14.7 days for the deceased patients. Moreover, the actual NAb titer is significantly higher in deceased patients compared with that in the recovered patients during the same time period (14). These findings suggest a role of anti-S antibodies in SARS-CoV–mediated ALI during acute infection. Consistently, preexisting serum antibodies against influenza antigens were found to associate with worse clinical severity and poor outcomes in patients during the 2009 influenza pandemic (15, 16). Moreover, multiple vaccine platforms and viral infection appeared to induce SARS-CoV–specific immune memory that enhanced lung inflammation following homologous challenge in mice and African green monkeys (17–19). The mechanism responsible for the immunopathologic reaction remains elusive. Recent studies suggested that T cells play a crucial role in protection of mice against lethal SARS-CoV infection (20–22). Enhanced pulmonary immunopathology in vaccinated and challenged animals reflects an inadequate Th1 response (22, 23). The role of virus-specific antibody response in SARS-CoV–induced lung injury has yet to be clearly defined. Therefore, we used vaccination and anti–S-IgG passive immunization strategies to evaluate the effects of anti-S antibodies on SARS-CoV–induced ALI in Chinese rhesus monkeys (Macaca mulatta). Our results now show that, in productively infected lungs, anti–S–IgG causes severe ALI by skewing inflammatory resolving responses during acute infection.

**Results**

*Vaccine-induced S-specific immunity resulted in severe ALI in SARS-CoV infected Chinese macaques.* We initially compared the pathological changes in the lungs of rhesus macaques vaccinated with a modified vaccinia Ankara (MVA) virus encoding full-length SARS-CoV S glycoprotein (ADS-MVA) or control at 7 and 35 days after pathogenic SARS-CoV PUMC (1 × 10⁵ tissue culture infectious dose of 50% [TCID₅₀]) challenge (24). Three healthy macaques were included as controls. Sixteen macaques were i.m. immunized twice: 8 animals with
ADS-MVA and another 8 animals with a control MVA (ADC-MVA) (Figure 1A) (24). Vaccination with ADS-MVA induced high levels of anti–SARS-CoV NAbs in all 8 macaques, with sera IC_{50} values ranging from 10,232- to 28,703-fold dilutions (Figure 1B). None of the macaques developed fever during or after vaccination (normal body temperature 37.8°C–38.1°C). The IC_{50} values of these sera were maintained, with an over-2,000-fold reciprocal serum dilution at 4 weeks after the secondary immunization, a time when all of the animals were i.n. challenged (Figure 1B). After viral challenge, all macaques developed fever between 1 and 5 days after inoculation (dpi) ranging from 38.6°C–39.2°C, and sera from control vaccinated macaques showed increased neutralizing activity after 14 dpi, indicating establishment of infection (Figure 1B). SARS-CoV RNA was readily detected on oral swabs from all control macaques examined at multiple time points using real-time reverse transcription PCR (real-time RT-PCR) but only in 3 of the 8 ADS-MVA–vaccinated macaques with lower NAb titers at 2 dpi (SL11, SE13, and SL14) (Figure 1C), suggesting reduced productive viral infection in the immunized macaques through S-specific immunity.

However, histological examination revealed acute DAD with various degrees of severity in 6 ADS-MVA–vaccinated macaques at 7 and 35 dpi, whereas most control macaques in the ADC-MVA group showed minor to moderate inflammation (Figure 1D). To better characterize the pathological changes, we adopted a 6-grade scoring system to describe the severity of the lung damage from least severe to most severe: 0, –; 1, +; 2, ++; 3, +++; 4, ++++; and 5, +++++ (Supplemental Figure 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.123158DS1). In the control vaccinated group, 5 macaques showed minor inflammation with slight septa broadening and inflammatory cells infiltration that we scored as + (CL23, CE25, CE20, CE21, and CL19; Figure 1D, Supplemental Figure 1B, and Supplemental Figure 2). Two macaques showed moderate inflammation with more interstitial mononuclear inflammatory infiltration and were scored as ++ (CL26, CE24; Supplemental Figure 1C and Supplemental Figure 2). Only 1 macaque showed typical symptoms of acute DAD with extensive exudation and cell infiltration at 35 dpi (CL22, scored as ++++; Supplemental Figure 2). In the ADS-MVA–vaccinated group, 4 macaques showed typical symptom of acute DAD with extensive exudation and many-cell infiltration in alveolar cavities (SL11, SE12, SL16, and SL18, scored as ++++; Supplemental Figure 1E and Supplemental Figure 2). One macaque showed severe acute DAD with exudation, hyaline membrane formation along the alveoli, pneumocyte desquamation, and damaged alveolus filled with hemorrhage and inflammatory cells (SL15, scored as +++++; Figure 1D). One macaque showed early symptom of acute DAD (SL14, scored as ++++; Supplemental Figure 1D), and the remaining 2 macaques showed moderate inflammation (SE13 and SE17, scored as ++; Supplemental Figure 2). The comparison of lung histopathological scores of 2 groups showed significantly enhanced lung injury in the ADS-MVA–vaccinated group at both 7 and 35 dpi compared with the control ADC-MVA group, suggesting that S-specific — but not MVA-specific — immunity promotes ALI during SARS-CoV infection (Figure 1E). Moreover, there is a moderate correlation between lung pathological scores and sera NAb titers at 0 dpi (Figure 1F), suggesting a role of S-specific antibody in enhancing SARS-CoV–mediated lung injury.

Anti–S-IgG induced severe lung injury during acute SARS-CoV infection. In our previous study, i.n. inoculation of Chinese macaques with SARS-CoV\_PUMC (1 × 10^5 TCID\_50) led to lower respiratory tract infection in all animals within 2 days (25). However, most of them rapidly cleared infection in the lungs, and all of the animals (n = 8) exhibited mild lung lesions before 3 dpi (25). Severe lung injury in SARS-CoV–infected Chinese macaques has not been detected until 7 dpi (26, 27). To determine the effect of anti-S antibody on the extent of SARS-CoV–mediated lung injury, we adoptively transferred 5 mg (low dose) and 200 mg (high dose) purified anti–S-IgG from ADS-MVA–vaccinated but unchallenged macaques into 2 groups of unvaccinated macaques (n = 6/group) via i.v. injection, and then i.n. challenged recipients with SARS-CoV\_PUMC (1 × 10^5 TCID\_50) (Figure 2A). As controls, another 2 macaques were administered 200 mg of control IgG (C-IgG) derived from ADC-MVA–vaccinated macaques. We sacrificed half of the macaques in each group at 2 dpi to avoid potential disruption by virus infection–induced antibodies against nucleocapsid protein (N) and other viral proteins — and prior to when induction of ALI was typically observed. The remaining half of the macaques in each group was sacrificed at 21 dpi to evaluate long-term impact (Figure 2A). At day 2, sera from macaques that received low- or high-dose S-IgG showed neutralizing activity, with IC_{50} values ranging from 10–260 and 1,000–10,000, corresponding to the dose of sera transfer; macaques that received C-IgG failed to show neutralizing activity (Figure 2B and Supplemental Table 1). While NAb titers in the sera of macaques in the high-dose group declined steadily between 2 and 21 dpi, macaques in the low-dose and control groups displayed increased NAb titers over time (Figure 2B), suggesting active viral replication.
Indeed, SARS-CoV RNA was detected using real-time RT-PCR on oral swabs, and/or virus was recovered from lung tissue in both control animals and 5 of 6 animals (83%) in the low-dose group. Only 2 of 6 animals (33%) in the high-dose group were viral RNA+ (Figure 2C and Supplemental Table 1), suggesting reduced viral production by high-dose S-IgG.

Consistent with the findings in the control ADC-MVA–vaccinated group, histopathological examination of the lungs indicated minor and moderate inflammation at day 2 and 21 in C-IgG recipients, respectively. In contrast, all S-IgG recipients exhibited symptoms of acute DAD with various degrees of exudation, hyaline membrane formation, and hemorrhage and inflammatory cells within alveoli at both day 2 and 21 (Figure 2, D and E, and Supplemental Table 2). Of note, despite the presence of high titer sera NAb titers at 0 dpi, only 1 of 5 C-IgG and control
ADC-MVA vaccine recipients showed acute DAD (Figure 1E and Figure 2E), indicating a time-restricted role for S-IgG, which mostly causes severe ALI in acutely infected macaques. Therefore, we conclude that, despite viral suppression, the presence of S-IgG at the acute stage of SARS-CoV infection caused severe ALI that persists until late stages.

Figure 2. Anti-spike antibodies induced ALI in SARS-CoV–infected Chinese rhesus macaques. (A) Experimental design used to investigate the influence of S-IgG on SARS-CoV–induced lung injury. Two groups of Chinese rhesus macaques (n = 6/group) were subjected to i.v. injection of high-dose (200 mg) or low-dose (5 mg) purified S-IgG from ADS-MVA–vaccinated but unchallenged macaques. As controls, another 2 macaques were administered 200 mg of C-IgG derived from ADC-MVA–vaccinated macaques. After 2 days, 3 groups of macaques were challenged i.n. with SARS-CoV PUMC (1 × 10^5 TCID₅₀). Half of the animals from each group were sacrificed at 2 and 21 dpi. (B) Sera from macaques at the indicated time points were tested for the capacity to neutralize SARS-CoV pseudotype virus. (C) Detection of viral RNA in oral swabs. SARS-CoV RNA was detected by nested RT-PCR in the swabs from one of the high-dose S-IgG–treated macaques (n = 6), 3 low-dose S-IgG–treated macaques (n = 6), and 2 C-IgG–treated macaques (n = 2) at the indicated time points. Left y axis shows the viral RNA copy number per milliliter swab. Right y axis shows the serum NAb titers of each macaque at 2 dpi, which are highlighted by shaded area. (D) Pathology changes of the lung tissue (200×). Sections from macaques were stained with H&E. Images show symptom of acute DAD exhibited in macaques received high-dose and low-dose S-IgG (n = 12) with extensive exudation (red arrows), hyaline membranes (black arrows), and massive cell infiltration (yellow arrows) at 2 and 21 dpi. Right panel shows minor and moderate inflammation in the macaques received C-IgG with slight alveolar septa broadening and sparse monocyte infiltration at 2 and 21 dpi. (E) Histopathological scores of the high-dose and low-dose S-IgG groups, including lung samples collected at both 2 and 21 dpi, were compared against the C-IgG group. See Supplemental Figure 1 for the scoring index based on severity of lung histopathology.
S-IgG failed to prevent SARS-CoV lower respiratory tract infection and amplified IMM infiltration and accumulation in the lungs. To determine the potential cause of S-IgG–enhanced ALI, we measured viral infection and monocytes/macrophages infiltration in the lungs and serum cytokine profile at 2 dpi. We previously found that pulmonary infection of Chinese macaques could be rapidly controlled. Viral RNA+ cells were detectable in the lungs of only 2 of 4 animals at 2 dpi, although all of them were viral nucleoprotein-positive (NP+) in the lungs and the hilar lymph nodes (LNs), where the lymphatics of the lungs drain (25). Therefore, we measured viral RNA+ cells by in situ hybridization (ISH), as well as NP signals in the lungs and the hilar LNs by IHC staining to evaluate the actual pulmonary infection of each macaque. Consistent with the results by viral isolation (Supplemental Table 1), viral RNA was detected in pneumocytes of 1 macaque in the high-dose (HS02, 33%), 2 macaques in the low-dose (LS01, LS02; 66%), and the C-IgG–treated macaque (Figure 3A and Supplemental Table 3), suggesting reduced productive infection by S-IgG. However, an NP antibody identified positive signals in the lungs and hilar LNs of all S-IgG and C-IgG recipients (7 of 7 macaques) but not healthy controls (Figure 3B and Supplemental Table 3), suggesting that S-IgG failed to prevent SARS-CoV entry, and SARS-CoV established lower respiratory tract infection in all of the challenged animals.

Consistent with previous findings in SARS-CoV–inoculated naive Chinese macaques (n = 4) (25), double-immunostaining with the NP antibody and antibodies specific to the stages of macrophage differentiation and inflammation (MAC387, CD68, CD163) revealed significant infiltration of IMMs in the lungs of C-IgG recipients, consisting of newly infiltrated (MAC387+) and inflammatory (CD163+) monocytes/macrophages (Figure 3C and Supplemental Figure 3). The CD163+ cells greatly exceeded the CD68+ population and typically gathered around NP+ cells to form cell clusters (Figure 3C), showing an association between viral infection and IMM infiltration in these macaques.

Compared with C-IgG recipients, more robust signals for MAC387 and CD163 were identified in the lungs of S-IgG recipients (Figure 3C and Supplemental Figure 3), suggesting enhanced IMM infiltration by S-IgG. MAC387+ and CD163+ monocytes/macrophages not only gathered around NP+ cells, but also formed clusters in tissues with little NP signal (Figure 3C), suggesting that infection-induced recruitment from circulation continued after viral clearance and accumulated at the inflamed site. Moreover, more-concentrated CD68+ signals were found, indicating altered macrophages functional response by S-IgG treatment (Figure 3C). Consistently, elevated serum levels of(100,452),(520,498) were observed in macaques treated with high-dose S-IgG at 2 dpi (Supplemental Figure 3), and the IL-8 levels are strongly correlated with anti–SARS-CoV NAb titers (Supplemental Figure 3). IL-8 is a macrophage-derived inflammatory cytokine important in initiating ALI. These results, therefore, suggest that S-IgG likely altered macrophage functional response in the lungs during acute infection, resulted in increase in IL-8 production, and enhanced ALI and IMM infiltration and accumulation.

Alveolar monocytes/macrophages assumed a wound-healing function as early as 2 dpi in macaques not treated with S-IgG. Macrophage activation is dynamic and plastic. Different monocyte/macrophage activation statuses have different specialized and critically timed roles, taking part in either the initiation and maintenance or resolution of inflammation (9). A well-organized wound-healing response leads to timely resolution of inflammation, whereas disturbance in wound-healing response results in persistent inflammation and injury. To understand the S-IgG–caused excessive inflammation, we further defined macrophage heterogeneity and activation statuses in the lung lesions of C-IgG (n = 1) and S-IgG recipients (n = 3) at 2 dpi. Three healthy macaques and 4 naive Chinese macaques challenged with the same dose of SARS-CoV PUMC and scarified at 2 dpi from a previous study were included as controls (25).

By combining 5 macrophages markers (MAC387, CD68, CD163, HAM56, and CD206) (28–30), 2 subpopulations of resident macrophages were shown in the lungs of Chinese rhesus macaques at steady state as previously described (29). They are interstitial macrophages (IMs; MAC387+HAM56 CD206-) (Figure 4A) and resident alveolar macrophages (AMs; CD68+ HAM56+ expressing low levels of CD163 and CD206, a marker specific to the macrophage wound-healing response, corresponding to their intrinsic antiinflammatory function (Figure 4B) (29). In lung lesions of both C-IgG–treated and naive macaque groups challenged with SARS-CoV PUMC, IMMs were further characterized and divided into 2 subpopulations. One subpopulation was the recently recruited MAC387+HAM56 CD68+ monocytes (Figure 4C) (29), which is like IM but is larger in size and primarily coexpressed CD163 (Supplemental Figure 4). The other subpopulation was single CD163+ inflammatory macrophages that had differentiated from newly recruited MAC387+ monocytes/macrophages (Figure 4D and Supplemental Figure 4) (29). Importantly, resident
AMs (CD68+) differentiated into HAM56+ cells, with a slight increase in numbers and increased size, with stronger CD163 and — more importantly — CD206 staining, suggesting that they are alternatively activated (Figure 4D). Strikingly, elevated levels of TGF-β expression were detected in all MAC387+ — and most enlarged CD163+ — as well as CD68+ macrophages in the lungs of both C-IgG–treated and unvaccinated macaques compared with healthy controls (Figure 4, G–I), indicating that many IMMs and activated resident AMs have assumed a wound-healing function in infected lungs within 2 days of infection.

S-IgG treatment skewed wound-healing macrophage response in the lungs during acute SARS-CoV infection. Compared with the macaques not treated with S-IgG, the number of IMMs (MAC387+ and CD163+HAM56+) significantly increased in S-IgG recipients (Figure 4, E and F). Moreover, many recently recruited MAC387+HAM56+ monocytes/macrophages did not coexpress CD163 (Supplemental Figure 4), representing the newest recruited blood monocytes, as a result of enhanced monocytes infiltration by S-IgG. Importantly, the majority of CD163+HAM56+ IMMs coexpressed CD68 (Supplemental Figure 4), suggesting a change of function of this subpopulation by S-IgG. Moreover, resident macrophages (CD68+HAM56+) in the lungs of S-IgG recipients no longer express CD206, which — being surrounded by many CD163+ IMMs — did not coexpress CD163 (Supplemental Figure 4), indicating that many IMMs and activated resident AMs have assumed a wound-healing function in infected lungs within 2 days of infection.

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Onset of an antibody response prior to viral clearance is associated with abrogated wound healing responses and increased IMM lung infiltration in unvaccinated Chinese rhesus macaques. To understand the mechanism underlying the time-restricted role of S-IgG in promoting ALI, we conducted a temporal analysis of the productive viral infection, antibody response, and macrophage functional changes in unvaccinated Chinese macaques during the first week of infection. Naive macaques were challenged i.n. with 1 × 10^5 TCID50 SARS-CoV PUMC and sacrificed at 2, 3, and 7 dpi (4 macaques/group) (Figure 5A) (25). Successful lower respiratory tract infection was confirmed by detection of viral NP using IHC staining in the lungs of all macaques (25). Viral RNA+ alveolar pneumocytes were primarily found at 2 dpi (2 of 4 macaques), and infrequently at 3 dpi (0 of 4 macaques; Figure 5B and Supplemental Table 4) (25). At 7 dpi, viral RNA was detectable in the lungs of 2 macaques, AD0516 and AD0518, suggesting productive infection there (Figure 5, B and C). Only few scattered NP+ cells were found in AD0517 (Figure 5D). Neither NP nor viral RNA was found in the lungs or swabs of AD0515, suggesting clearance of productive viral pulmonary infection in this macaque (Figure 5D and Supplemental Table 4). Serum NAb was not observed at 2 and 3 dpi until 7 dpi in 2 of 4 macaques, AD0515 and AD0516 (Supplemental Table 4 and Figure 5E).

In agreement with the findings in the C-IgG group, temporal analysis of macrophage heterogeneity revealed significantly increased number of IMMs (CD163+CD206+) in viral RNA+ lungs at 2 dpi (Figure 5F, right image). The IMMs diminished at 3 dpi (4 of 4 macaques) (Figure 5G) and became nearly completely absent at 7 dpi in macaques without productive pulmonary viral infection (2 of 4 macaques, AD0515 and AD0517) (Figure 5H), indicating that inflammatory response resolved with 7 days following the control of productive infection. Critically, wound-healing macrophages (CD163+CD206+ and CD163+TGF-β+) were observed in the lungs of all macaques at 2 dpi, despite productive pulmonary viral infection in macaques AD0506 and AD0508 (Figure 5F). In contrast, macroce AD0516, with serum NAb and productive virus infection in the lungs, displayed a significantly enhanced proinflammatory macrophage response and reduced inflammatory-resolving macrophages (Figure 5I), which was not observed in macaques with productive viral infection but undetectable NAb responses (AD0518) or in macaques with detectable NAb but undetectable viral replication (AD0515) (Figure 5D and Supplemental Table 4). Therefore, persistent viral replication and concomitant NAb detection is associated with increased IMM lung infiltration and reduced wound-healing macrophage numbers.

S-IgG modifies functional response of alternatively but not classically activated macrophages to SARS-CoV infection. To determine whether S-IgG skews wound-healing response through modifying wound-healing macrophage functional response during SARS-CoV infection, we treated in vitro–polarized alternatively activated human monocyte–derived macrophages (MDM) for 20 hours with SARS-CoV pseudovirus and heat-inactivated sera of high-dose S-IgG–treated macaques (S-IgG sera) collected at 2 dpi. We
then measured the concentrations of 13 inflammatory cytokines in cell culture supernatants, including IL-1β, IFN-α, IFN-γ, TNF-α, MCP1, IL-6, IL-8, IL-10, IL-12p70, IL-17A, IL-18, IL-23, and IL-33. Sera from C-IgG recipients or healthy macaques were included as controls. In vitro–polarized classically activated MDM was also studied and compared with alternatively activated MDM in parallel. Consistent with previous report, unstimulated classically activated MDM produced high levels of IL-8 and IL-6; alternatively activated MDM only produced minimal levels of those cytokines (Figure 6A). Addition of SARS-CoV pseudovirus induced IL-8 production in alternatively activated MDM to levels comparable with those found in unstimulated, classically activated MDM, as well as in low-level MCP1 and IL-6 production (Figure 6B). By contrast, SARS-CoV pseudovirus caused less than a 1-fold increase in IL-8 secretion, but not other cytokine secretion, by classically activated MDM (Figure 6C). Strikingly, while none of the sera samples induced cytokine production when they were administered into MDMs alone,
**Figure 4. Comparison of monocyte/macrophage phenotype and function in the lungs of S-IgG– and C-IgG–treated macaques.** (A–F) Phenotype of monocytes/macrophages subpopulations in the lungs of healthy, C-IgG–, and S-IgG–treated macaques (original magnification, 200×). These sections were triple-immunostained with antibodies for CD206 (cyan), HAM56 (FITC), and MAC387 (TRITC) (A, C, and E); CD206 (cyan), HAM56 (FITC) (B, D, and F), and CD68 (TRITC) (upper panel in B, D, and F); or CD163 (TRITC) (lower panel in B, D, and F). A and B show 2 subpopulations at steady states, including interstitial macrophages (IM) (MAC387+HAM56–CD206–) (yellow arrow), and resident alveolar macrophages (AM) (CD68+CD163loCD206loHAM56–) (white arrows). C and D show the presence of alternatively activated resident AM and accumulated IMMs in the C-IgG group. Alternatively activated AMs are CD206hiCD68+CD163hiHAM56+ (D, white arrows). IM are MAC387+HAM56–CD206– (C, yellow arrow). IMMs include newly infiltrating monocytes/macrophages (MAC387+) (C, green arrow); and inflammatory macrophages (CD163+CD68–CD206–HAM56–) (D, green arrow). E and F show significantly increased number of IMMs and decreased number of alternatively activated macrophages in the S-IgG group. IMMs include newly infiltrating monocytes/macrophages (MAC387+) (E, green arrow) and inflammatory macrophages that are CD68+HAM56+ (green arrow, F, upper panel) and CD163+CD206–HAM56– (green arrow, F, lower panel). Resident AMs are no longer expressing CD206 (CD68+HAM56+CD163–CD206+) (blue arrow, F, upper panel). (G–J) TGF-β and IL-6 expression in macrophages in the lungs. These sections were double-immunostained with antibodies for TGF-β or IL-6 (TRITC) and MAC387, CD68, or CD163 (FITC). G shows low expression levels of TGF-β and IL-6 in IMs (MAC387+) and AMs (CD68+) in healthy macaques. H shows increased TGF-β expression in the C-IgG group (white arrows) but increased IL-6 in the S-IgG group (blue arrows). I and J show the numbers of TGF-β+ and IL-6+ monocytes/macrophages in 3 groups in a 200× field.
administration of S-IgG sera caused a dose-dependent increase in the production of all 3 inflammatory cytokines in SARS-CoV–stimulated alternatively activated MDM, resulted in a greater than 10-fold increase in IL-6, an 8-fold increase in MCP1, and a 5-fold increase in IL-8 (Figure 6, D and E). This enhancement, however, was not observed when S-IgG sera were replaced with sera from C-IgG recipients or healthy macaques (Figure 6, F, G, and H), suggesting that S-IgG, but not other sera components, enhanced SARS-CoV–stimulated cytokine production by alternatively activated MDM. Administration of S-IgG sera had no effect on cytokine production by SARS-CoV–treated classically activated MDM (Figure 6, I, J, and K), indicating that S-IgG do not modulate virus-mediated classically activated macrophage functional response. MCP1 and IL-8 are key cytokines that regulate monocytes/macrophages migration and promote neutrophils infiltration, while IL-6 promotes persistent inflammation and injury. These results suggest that, in response to SARS-CoV infection, wound-healing macrophages produce IL-8 and low amounts of MCP1 to recruit neutrophils and blood monocytes, whereas the presence
of S-IgG can result in 5- to 10-fold increases in IL-8, MCP1, and IL-6 production by wound-healing macrophages, leading to significantly enhanced and persistent monocytes infiltration and tissue injury. Critically, in the presence of Fcγ receptor (FcγR) blocking antibody, IL-8 was significantly reduced when cells were treated with SARS-CoV pseudovirus and sera from high-dose S-IgG–treated macaques (diluted 1:4,000), indicating that anti–SARS-CoV antibody modifies wound-healing response partially through FcγRs (Figure 6L). Therefore, we conclude that S-IgG directly modifies functional response of alternatively activated macrophages during acute infection, thus skewing wound-healing response, leading to hypercytokinemia, aggressive inflammation and severe lung injury.
Deceased SARS patients displayed higher levels of serum NAbs, accumulated proinflammatory, macrophages infiltration, and absence of wound-healing macrophages in the lungs. To examine the role of S-IgG in promoting ALI in a clinical setting, we investigated antibody responses during the early stage of infection in 2 groups of deceased patients or individuals recovered from SARS and characterized the heterogeneity of monocytes/macrophages in the lungs of 3 deceased SARS patients. H&E staining of lung specimens derived from 3 deceased SARS patients revealed severe ALI (Figure 7A). Consistent with the macaque findings, IHC analysis revealed significant infiltration of CD163+ monocytes/macrophages into the lungs of patients, with little TGF-β expression and loss of the wound-healing marker CD206+, indicating predominantly proinflammatory macrophage activation and reduced inflammatory-resolving response (Figure 7B). Moreover, significantly higher levels of anti-S NAbs were detected in the sera of deceased patients (n = 6) during acute infection compared with recovered patients (n = 8) (Figure 7C). These results, therefore, are consistent with findings in experimental macaques.

Antisera from deceased SARS patients skewed wound-healing response during SARS-CoV infection partially through the FcγR. To examine whether human anti–SARS-CoV antibody modifies wound-healing response during SARS-CoV infection, we treated in vitro–polarized alternatively activated MDM with SARS-CoV pseudovirus for 20 hours in the presence of sera collected from SARS patients during acute infection. Macrophages treated with SARS-CoV pseudovirus or sera alone were included as controls. We then examined the protein levels of cytokines/chemokines in the cell culture supernatant. Consistent with findings in macaques, administration of sera from a deceased SARS patient (D1; Supplemental Table 5) into SARS-CoV–treated alternatively activated MDM caused a dose-dependent increase in the production of cytokines, leading to a 2- to 3-fold increase in MCP1 and IL-8 after 20 hours (Figure 8, A and B). This enhancement, however, was not observed in cells treated with sera alone (Figure 8, A–D). Administration of sera from recovered patients (n = 8; Supplemental Table 5) had no effect on cytokine production except 1 sample (Figure 8, C and D). These differences may be explained, at least in part, by their lower NAb titers, although it is also possible that sera from recovered patients contain different antibody populations. Consistently, the only serum sample — R2, which enhanced chemokine production — has a relatively higher NAb titer among the samples tested (Supplemental Table 5). Moreover, statistical analysis shows that IL-8 production strongly correlates with NAb titer of sera from deceased patients (Figure 8E). Furthermore, IL-8 production was significantly reduced when cells were treated with SARS-CoV pseudovirus and sera from deceased SARS patient (diluted 1:4,000) in the presence of FcγR blocking antibody, indicating that anti–SARS-CoV antibody modifies wound-healing response in humans partially through FcγRs (Figure 8F).

Discussion

Here, we present evidence of a detrimental role of anti–S-IgG in ALI during SARS-CoV infection. Respiratory CoVs infection poses a unique challenge to the immune system: not only must the virus be rapidly eliminated, but lung inflammation must also be controlled to prevent acute respiratory failure (33). In the present study, we show that, despite markedly reducing virus titers, anti–S-IgG caused lung injury during the early stages of infection by abrogating a wound-healing macrophage response and TGF-β production, while promoting proinflammatory cytokine IL-8 and MCP1 production and inflammatory macrophage accumulation. To our knowledge, our data demonstrate a previously unrecognized mechanism underlying virus-mediated ALI and suggest that modulation of the anti-S antibody response or blockage of Fcγ receptors during acute infection might be needed for effective treatment for respiratory CoV infection.

Many studies have demonstrated the role of NAbs induced by the S glycoproteins in protecting viral replication in susceptible hosts, including mice, ferrets, hamsters, and macaques (24, 34, 35). However, the effect of S-specific immunity in protecting against pulmonary immunopathology has been controversial. In some cases, vaccination-induced immunity assists the viral clearance and protects mice or ferrets against lethal challenge (36–39), whereas in many other situations, multiple vaccine platforms appear to induce increased eosinophilic proinflammatory pulmonary response upon challenge (18, 23, 40). These differences may be explained, in part, by the characteristics of the vaccine being studied, the infectious dose, and the viral strain employed, as well as by the type or quality of immune response elicited. Recent studies suggest that CD8+ T cell response plays a crucial role in viral clearance and protection of mice against eosinophilic immunopathology and lethal SARS-CoV infection, whereas immunopathology predominantly reflects an inadequate vaccine-induced Th1 response (20–23).
In fact, a protective effect against pulmonary immunopathology mediated by full length S protein–based vaccine in SARS-CoV–infected nonhuman primate models has not been described, to our knowledge. Although several vaccine candidates protected macaques against viral replication in the lungs, pulmonary immunopathology was not evaluated in these studies (24, 41, 42). By contrast, it was shown that SARS-CoV–specific immune memory induced by prior infection enhanced lung inflammation following homologous challenge in African green monkeys (17). Similarly, augmentation of disease by vaccination has also been described in studies of atypical measles and dengue hemorrhagic fever, as well as several respiratory diseases, including respiratory syncytial virus (RSV) and pandemic influenza (16, 43). In the study of RSV vaccine conducted in 1966 and 1967, 80% of RSV vaccinees needed hospitalization, whereas only 5% of RSV-infected children in the control vaccine group required admission, although the underlying mechanisms remain incompletely understood. Using Chinese rhesus macaques, we show that an ADS-MVA–based protected macaques against viral replication but significantly enhanced acute DAD at both 7 and 35 dpi compared with the control ADC-MVA group, suggesting that S-specific but not MVA-specific immunity promotes ALI (Figure 1). Until now, it remains unknown if SARS vaccine candidates with a focus on the receptor-binding domain (RBD) of S protein can avoid the induction of ALI in nonhuman primates. These vaccines have been shown to induce highly potent NAb responses and elicit long-term protective immunity in immunized small animals (38).

SARS-CoV infection of Chinese macaques is often characterized by the rapid control of viral replication with mild lung lesions in most macaques (25). The mechanisms underlying why Chinese macaques do not often develop ALI, as observed in most SARS patients, are largely unknown. Our data suggest that these effects, in part, reflect the rapid control of viral replication in the lungs, which peaked between 24 and 48 hours after inoculation (hpi) and diminished within 7 days, thus creating an interval between lung inflammation and high titer antibody production in most macaques. Indeed, although SARS-CoV infection resulted in significant infiltration of macrophages in the lungs, the number of macrophages was rapidly reduced following virus clearance at 3 dpi (Figure 5). At 7 dpi, inflammation appeared to be mostly resolved (Figure 5). By contrast, control of virus replication is less efficient in SARS patients, and the peak in viral load coincides with the first appearance of an antibody response, approximately 10 days after the onset of symptoms (6).

The low rate of ALI and interval between lung inflammation resolution and antibody production, however, makes Chinese macaques an ideal animal model for the present study. Using vaccination and passive immunization, we conditioned anti-S antibody titers in macaques during the early stage of infection when ALI was not typically observed. We found that prior administration of anti–S-IgG led to massive accumulation of monocytes/macrophages in the lungs within 2 dpi. This result is consistent with previous reports, in which mice given the SARS vaccine exhibited an immunopathologic lung reaction after the subsequent challenge with SARS-CoV (18, 19). Moreover, in SARS patients, clinical course and outcome are more favorable in children younger than 12 years of age compared with adolescents and adults (44), indicating that prior alternative CoV infection might
A recent study reported that pathogenic immune complexes promoted lung injury during the 2009 H1N1 pandemic (16). It remains unknown whether immune complexes may also play a role in driving ALI during SARS-CoV infection.

Detailed analyses of macrophage heterogeneity at different stages of infection and in lungs with severe injury or mild lesion revealed distinct roles for proinflammatory and wound-healing macrophages in SARS disease progression. We observed that the early transition of macrophage responses from proinflammatory...
to wound healing with increased TGF-β expression is associated with inflammation resolution and mild lung injury. By contrast, abrogated wound healing by S-IgG resulted in decreased TGF-β production and prolonged classical macrophage activation, and it promoted severe lung injury. Our findings are primarily based on imaging of formalin-fixed tissues of infected macaques, which may have technical limitations. A detailed flow cytometric analysis would be helpful to further characterize macrophage subpopulations in infected lungs. Unfortunately, we were not allowed to obtain any live cells, including macrophages, for analysis by flow cytometry outside the BSL-3 laboratory in China, which is in line with “WHO biosafety guidelines for handling of SARS specimens” (https://www.who.int/csr/sars/biosafety2003_04_25/en/). Having said this, TGF-β is mainly derived from wound-healing macrophages but not from classically activated macrophages. Therefore, by staining both TGF-β and CD206 (another biomarker of wound-healing macrophages), we believe that the finding of reduction of this type of macrophages in the S-IgG–treated lungs is convincing.

Patients who received convalescent plasma from recovered SARS patients had shorter hospital stays and lower mortality rates without evident adverse effects (45–47). These results might reflect a different quality of antibodies in recovered patients from deceased patients during early infection. Indeed, we previously observed that deceased patients had significantly higher NAb’s with a very low level of anti-N antibodies in serum during the early stage of infection (14). In contrast, recovered patients had more rapid anti-N antibody and slower NAb development (14). Consistently, serum from deceased patients — but not SARS survivors during early stages of infection — enhanced IL-6, IL-8, and MCP1 production by wound-healing macrophages (Figure 8). We previously identified epitopes in S protein that elicited NAb’s and antibodies that enhanced SARS-CoV infection (48, 49). Further studies that will attempt to identify antibodies and epitopes that mediate prevention or enhancement of ALI might be needed for future vaccine design and immunotherapy.

Lastly, blockade of FcγRs reduced proinflammatory cytokine production, suggesting a potential role of FcγRs for the postulated reprogramming of alternatively activated macrophage. FcγRs are functionally divided into activating and inhibitory receptors. Activating FcγRs, such as FcγRI, FcγRIIA, and FcγRIII, triggers production of proinflammatory chemokines and cytokines through an immunoreceptor tyrosine–based activation motif (ITAM) in their intracytoplasmic domain and SRC family kinases and spleen tyrosine kinase (SYK), whereas the inhibitory FcγR, FcγRIIB, has an immunoreceptor tyrosine–based inhibition motif (ITIM) in its intracytoplasmic domain and counteracts the signals that are mediated by activating FcγRs. Based on literature and our experiments, human monocyte–derived wound-healing macrophages express higher levels of CD16 (FcγRIII) and CD32A (FcγRIIA) (50). Therefore, it is likely that S-IgG promoted proinflammatory cytokine production through FcγRI and/or FcγRIIA. S-IgG did not affect classically activated macrophage function in vitro. This difference can be possibly explained by the high baseline level of proinflammatory cytokines before infection and the downregulated expression of FcγR on classically activated MDM by SARS-CoV treatment.

**Methods**

**Immunization of animals.** We vaccinated Chinese rhesus macaques as previously described (51). Briefly, 16 Chinese rhesus monkeys were immunized twice on days 0 and 28. Eight of these animals received ADS-MVA, whereas the other 8 animals received ADC-MVA. ADS-MVA and ADC-MVA vectors encoding SARS-CoV S protein or HCV E1E2 were constructed and produced as previously described (25). The vaccine strain SARS-CoV HKU39849 (AY278491) shares 100% sequence homology of the S gene with the challenge strain SARS-CoV PUMC (AY350750).

**Animals, tissues, and SARS-CoV infection.** SARS-CoV–infected animals and control animals were all adult female Chinese rhesus macaques. A total of 42 Chinese rhesus macaques were challenged i.n. with a live pathogenic SARS-CoV (PUM, TCID50 = 50) as previously described (25). An additional 3 uninfected macaques were inoculated with PBS as negative controls. Animals were subjected to daily measurement of anal temperature, routine blood assays, and chest radiography. Animals were euthanized at 2, 3, 7, 21, and 35 days after inoculation (Figure 1, Figure 2, and Figure 6). All animals were humanely euthanized with an i.v. overdose of pentobarbital and were immediately necropsied. Lung tissues (left and right cranial, middle and caudal lobes, and right accessory lobe) were collected and fixed in 10% neutral-buffered formalin, embedded in paraffin, and sectioned at 5 μm.

**Passive transfer of IgG.** SARS-CoV S protein–immune serum was generated by vaccination with ADS-MVA vectors. IgG was purified from pooled sera by protein G affinity chromatography to >95% purity. IgG was depyrogenated, concentrated to 20–30 mg protein ml⁻¹, and administered i.v. Recipients received 200 mg or 5 mg of purified IgG per kg body weight at 48 hours before the viral challenge.
Neutralization. A neutralization assay was established to determine the humoral immune responses generated by the vaccine as previously described (51). The neutralizing activity of heat-inactivated animal sera was determined using a pseudotype viral entry assay. The pseudotype virus was generated through cotransfection of 293T cells with 2 plasmids, pcDNA-Sop9 and pNL4-3Luc_Env_Vpr_, carrying the optimized S gene and a human immunodeficiency virus type 1 backbone, respectively. The serial diluted serum samples were incubated with equal amounts of pseudotype virus at 37°C for 1 hour. The serum-virus mixtures were subsequently added into preseeded HEK 293T-ACE2 cells. After 56 hours, infected cells were lysed to measure luciferase activity.

Nested RT-PCR and isolation of SARS-CoV. Pharyngeal swab samples were collected from infected monkeys on the first day after inoculation and subsequently tested for SARS-CoV using nested RT-PCR as previously described (25). Briefly, RNA was isolated using TRIZOL (Invitrogen). RT-PCR was performed in a 50 μl reaction volume with outer primer pair (5′-GCTGCATTGTTGTTATGCCTA-TGC-3′) and (5′-ATACAGATACAT-AGATTGCTGTATCC-3′), inner primer pair (5′-TCACTTGCTTC-CGTTAGGTCAGC-CAGCGTGGTGTTGCATACA-3′) and (5′-GGTTCGGAATGTATACACGC-GTCTCAGG-CAGAAAGCTGTAAGCT-3′). All PCR products were verified by nucleotide sequencing. SARS-CoV was isolated as previously described. Briefly, pharyngeal swab samples from the infected macaques were inoculated onto Vero-E6 cells and cultured in DMEM (Thermo Fisher Scientific). Both cytopathic effect (CPE) and immunofluorescence assay (IFA) (incubation with 1:10 dilution of SARS patient serum) were used to determine the infection status of the animals.

Double and triple immunofluorescence staining on paraffin sections. The samples were deparaffinized and rehydrated. After blocking with normal goat serum for 30 minutes at room temperature, a rabbit anti-SARS-CoV nucleocapsid antibody (ss-006-0100, eENZYME) was applied at 4°C overnight, followed by an Alexa Fluor 448 conjugated goat anti-rabbit IgG antibody for 1 hour at room temperature. Additional immunofluorescence staining was subsequently performed by overnight incubation with primary antibodies, followed by the detection of signals using appropriate fluorescent secondary antibodies (MilliporeSigma) for 1 hour at room temperature.

Primary antibodies for immunofluorescence staining. The following primary antibodies were used in this study: SARS-CoV NP (ss-006-0100, eENZYME); MAC387 (clone MAC387, AbD, Serotec); CD68 (clone KPI, DAKO; ab125047, Abcam); CD163 (clone EDHu1, AbD and ab183476, Abcam); HAM56 (clone HAM56, DAKO); CD206 (HPA004114, MilliporeSigma); TGF-β (sc-146, Santa Cruz Biotechnology Inc.); IL-6 (sc-1265, Santa Cruz Biotechnology Inc.); AE1/AE3 (clone AE1/AE3, DAKO).

ISH studies. ISH was developed for the detection of SARS-CoV. DNA fragments of N and S proteins from the SARS-CoV strain from Hong Kong (AY278491) were used as templates to generate RNA probes. The probes were labeled with digoxigenin (GENEWIZ Inc.). After deparaffinization and rehydration, the sections were treated with microwave heating in an 800 W domestic microwave oven (NN-ST556M, Panasonic) at medium power for 10 minutes in 0.01 M sodium citrate buffer. Hybridization was performed at high stringency with 150 ng/ml of denatured probe at 45°C for 16 hours in 50 μl of hybridization mix (50% formamide, 10% dextran sulfate in 2× saline sodium citrate [SSC]) in a moist chamber. Excess probe was removed by washing twice in 2× SSC for 20 minutes at 45°C, followed by 2 washes in 1× SSC and 0.1× SSC for 20 minutes at 45°C. Immunodetection was conducted using the DIG Nuclear Acid Detection Kit (Roche Diagnostics) according to the manufacturer’s instructions. Briefly, after incubation with 1% blocking reagent for 30 minutes, the sections were incubated with sheep anti-digoxigenin alkaline phosphatase conjugate (11093274910, Roche Diagnostics) diluted at 1: 500 for 2 hours at room temperature. After 3 separate washes with washing buffer (0.1 M maleic acid, 0.15 M NaCl [pH 7.5], 0.3% Tween 20), the signal was developed using 200 μl of a HNPP/Fast Red TR mix (1758888, Roche Diagnostics) in the dark for 45 minutes. The sections were counterstained with DAPI. Tissue sections incubated with sense RNA probe were used as a negative control.

Confocal microscopy. Confocal microscopy was performed using a Carl Zeiss LSM 700 confocal microscope equipped with 4 lasers (Zeiss). ZEN (Zeiss) was used to assign colors to the 4 channels collected: Alexa Fluor 568 and HNPP/Fast Red, which fluoresces when exposed to a 568-nm wavelength laser and appears red; Alexa Fluor 488 (Molecular Probes), which appears green; Alexa Fluor 647 (Molecular Probes), which appears cyan; and the differential interference contrast (DIC) image, which is in gray scale. The 4 channels were simultaneously collected. To differentiate between individual cells, Hoechst 33258 (nuclear marker; Molecular Probes) was used at 1 μg/ml and incubated for 5 minutes, followed by rinsing with water. The colocalization of antigens was demonstrated by the addition of colors as indicated in the figure legend.
Monocyte isolation, differentiation, and polarization of macrophages. Monocytes were purified from peripheral blood mononuclear cells obtained from blood buffy coats (provided by Red Cross, Hong Kong, China) or monkey peripheral blood, as we previously described (52). Monkey peripheral blood was obtained in heparinized vacutainer collection tubes from healthy female rhesus monkeys. To generate M1 macrophages, monocytes were differentiated in RPMI 1640 medium (Thermo Fisher Scientific) supplemented with GM-CSF (400 IU/ml), 2 mM glutamine, 10% decomplemented FCS, 100 IU/ml penicillin, and 100 μg/ml streptomycin (Thermo Fisher Scientific) for 4 days, followed by exposure to fresh medium supplemented with 5% FCS and containing GM-CSF and LPS (100 ng/ml) + IFN-γ (20 ng/ml) for an additional 72 hours. To generate M2 macrophages, monocytes were differentiated in RPMI 1640 medium (Thermo Fisher Scientific) supplemented with M-CSF (50 ng/ml), 2 mM glutamine, 10% decomplemented FCS, 100 IU/ml penicillin, and 100 μg/ml streptomycin for 4 days, followed by exposure to fresh medium supplemented with 5% FCS and containing M-CSF (50 ng/ml) + IL-4 (20 ng/ml) + IL-10 (100 pg/ml) for an additional 72 hours.

Quantification of cytokine levels. A panel of 13 cytokines, including IL-1β, IFN-α, IFN-γ, TNF-α, MCP1, IL-6, IL-8, IL-10, IL-12p70, IL-17A, IL-18, IL-23, and IL-33 in monkey sera and in culture medium were quantified using multiplexing laser bead technology (BioLegend). To assess the cytokine production of macrophages, in vitro–polarized cells were washed 3 times with PBS and were subsequently activated with SARS-CoV pseudotype virus in RPMI 1640 medium (Thermo Fisher Scientific) with or without administration of highly diluted and heat-inactivated (56°C) sera from macaques or SARS patients. FcγR-specific mouse monoclonal antibody (5 μg, clone 3G8, anti-hCD16, 16016682; clone FLI8.26, anti-hCD32, 16032891; and clone 10.1, anti-hCD64, 16064981 [BD Pharmingen]) was used for blockade of human FcγR. The supernatant of these cultures was recovered after 20 hours of stimulation, and the cytokine levels were measured and analyzed.

Statistics. All statistical analyses were performed using the 2-tailed Student’s t test. P values less than 0.05 were considered statistically significant. Data are presented as the mean values ± SEM of at least 3 independent experiments unless otherwise indicated.

Study approval. The experiment was conducted in a Biosafety Level 3 animal facility. Handling and experimental procedures, including injections, blood collection, virus challenge, and sacrificing were approved and performed according to the animal welfare committee on the Use of Live Animals in Teaching and Research of Institute of Laboratory Animal Science, Beijing, China. All study subjects gave informed consent, and procedures were reviewed and approved by the University of Hong Kong Research Ethics Committee.

Author contributions
ZC, LL, and QW designed the study. LL and QW analyzed the data. LL, ZC, KYY, SP, and AL contributed to drafting the manuscript. ZC is the principal investigator, with overall responsibility for the design of the study. QW and CQ coordinated the autopsy organization and collection of autopsy materials of Chinese macaques. QW was involved in the development and performance of the RT-PCR assay. LL, HT, TW, and KWC performed the neutralization assay. LL, XA, and AL were involved in the development of the H&E, SARS-CoV IHC, and ISH assays. HW performed the ISH assay and the IHC staining of NP and calculation of the mean fluorescence intensity in the lungs. LL, HK, and KN performed the H&E and double- or triple-IHC staining. JF and HT performed culture of monocyte-derived macrophages. QL, HT, JP, and ZT performed analysis of cytokine production in cell culture supernatant and monkey sera. KHC and KYY coordinated the collection of sera and tissue sample from SARS patients. All authors were involved in the correlative interpretation of the pathological and molecular data.

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