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The immune system and COVID-19: Friend or foe?

Fereshteh Yazdanpanah, Michael R. Hamblin, Nima Rezaei

Aim: Coronavirus disease 2019 (COVID-19) is a novel highly contagious infection caused by SARS-CoV-2, which has become a global public health challenge. The pathogenesis of this virus is not yet clearly understood, but there is evidence of a hyper-inflammatory immune response in critically ill patients, which leads to acute respiratory distress syndrome (ARDS) and multi-organ failure.

Material and methods: A literature review was performed to identify relevant articles on COVID-19 published up to April 30, 2020. The search resulted in 361 total articles. After reviewing the titles and abstracts for inclusion, some irrelevant papers were excluded. Additional relevant articles were identified from a review of citations referenced.

Key findings: SARS-CoV-2, directly and indirectly, affects the immune system and avoids being eliminated in early stages. On the other hand, the secretion of inflammatory cytokines creates critical conditions that lead to multi-organ failure.

Significance: The immune system which is affected by the virus tries to respond via a cytokine storm and hyperinflammation, which itself leads to further multi-organ damage and even death.

1. Introduction

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) causes the coronavirus disease 2019 (COVID-19), and has affected people's lives globally, since first observed in Wuhan, China in the last days of 2019 [1,2]. The main route of virus entry and transmission is respiratory droplets that are expelled and absorbed by the mucous membranes, especially the nasal and larynx mucosa. COVID-19 spreads readily via person-to-person contact [3]. The clinical spectrum of COVID-19 varies from an asymptomatic form to severe respiratory failure (SRF) that necessitates mechanical ventilation and support in an intensive care unit (ICU) and can lead to multi-organ failure. Pneumonia is the most frequent serious manifestation of COVID-19, characterized primarily by fever, dry cough, and dyspnea. Other less common symptoms are headaches, sore throat, and rhinorrhea. In addition to respiratory symptoms, gastrointestinal symptoms, myalgia, skin rashes, and neurological involvement have also been reported [1,3-6].

2. SARS-CoV-2 and the immune system

2.1. SARS-CoV-2 pathology

SARS-CoV-2 belongs to the coronavirus family, members of which have caused two previous epidemics at the beginning of the 21st century; one named SARS-CoV and the other Middle East Respiratory Syndrome (MERS). Coronavirus are large enveloped viruses with a positive sense RNA genome. The lipid bilayer envelope of the virus contains several proteins with different tasks. The spike or S glycoprotein (SP), has two domains of S1 and S2, is responsible for invasion, attachment, and entry into human cells. The receptor-binding domain (RBD) in S1 interacts with angiotensin-converting enzyme 2 (ACE2) on the human host cell surface, which is a similar entry mechanism to SARS-CoV; however, the S2 domain is responsible for virus-cell membrane fusion and viral entry with higher affinity [7]. Higher expression of the ACE2 receptor in adults compared to children may be a reason for the higher infection rate seen in adults [8,9]. Another noteworthy point

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is the increased level of enzymes in the liver, heart, and kidneys in COVID-19 patients with pneumonia, which is consistent with the tissue expression profile of the ACE2 receptor [10]; this could also explain the occurrence of multi-organ failure in some patients [11].

2.2. Effects of SARS-CoV-2 on the immune system

Since both SARS-CoV and SARS-CoV-2 have the same cell entry mechanism, the pathogenesis of both viruses could be the same, or at least very similar [12]. ACE2 is the common factor that binds to the superficial S glycoprotein on the envelope of the virus. It seems that this binding is sensed (essentially) by Toll-like receptor-7 (TLR-7), which is present in endosomes, and which then leads to the secretion of inflammatory cytokines [13,14]. ACE2 is highly expressed in some organs, like lung epithelial cells, especially type II pneumocytes, and in cells of the heart, kidneys, gastrointestinal tract, liver, and bladder [15,16]. Therefore these organs constitute the main target for the virus. Following entry of SARS-CoV-2 into the cell, the viral RNA genome is transferred from the envelope into the cytoplasm and the translation process begins. After replication of the RNA new viral particles are formed, by incorporating part of the host cell membrane in the new viral envelope. Although, SARS-CoV-2 buds from the infected cell, it does not lyse it directly [17]. Infected lung epithelial cells produce interleukin IL-8 which acts as a chemotactant for neutrophils and T lymphocytes [18]. The innate immune response is initially triggered by lung epithelial cells, alveolar macrophages and neutrophils. In the next stage, adaptive immune responses are triggered involving T and B lymphocytes to complete the complete immune response [19]. Virus particles containing single-stranded ssRNA, act as pathogen-associated molecular patterns (PAMPs), and provoke a strong innate immune response after recognition by Toll-like receptor 7 (TLR7), which is expressed on monocyte-macrophages and dendritic cells (DC). TLR7 can activate several signaling pathways and transcription factors, such as Janus kinase transducers (JAK/STAT), nuclear factor κB (NF-κB), activator protein 1 (AP-1), interferon response factor 3 (IRF3), and IRF7. This signaling cascade leads to increased secretion of pro-inflammatory cytokines, like IL-1, IL-6, monocyte chemo attractant protein-1 (MCP-1), MIP-1A, tumor necrosis factor α (TNF-α) and ultimately interferon 1 (IFN1) [20]. Furthermore, neutrophils are rapidly recruited to sites of infection, where they kill viruses by an oxidative burst, defensin secretion, and neutrophil extracellular traps (NETs) [21]. Along with these events, antigen presentation subsequently stimulates the body’s specific adaptive immunity (both humoral and cellular immunity) which culminates in approximately 7–14 days after infection. Following the representation of antigens by APCs to the CD4+ and CD8+ T-cells, pro-inflammatory cytokines are produced via the NF-κB signaling pathway. Activated B cells secrete virus-specific antibodies, while antigen-specific T cytotoxic cells kill virus-infected cells [17,22]. Additionally, Th17 cells, neutrophils, and granulocytes secrete IL-17, which in turn stimulates production of IL-1, IL-6, IL-8, MCP-1, Gro-α, G-CSF, GM-CSF, TNF-α, and PGE2. All these mediators can increase the recruitment of neutrophils, monocytes, and other immune cells. Besides, it has been reported that IL-17 expression is correlated with several inflammatory respiratory diseases [23]. All these immune signaling pathways are designed to create an inflammatory environment with the goal of eradicating SARS-CoV-2.

2.3. The immune system response

The pathology of SARS-CoV-2 is not yet completely understood; most of our knowledge has been based on research into SARS-CoV and MERS, which previously caused epidemics of acute respiratory syndromes. In this short duration of the present pandemic, studies have shown that SARS-CoV-2 has several defense mechanisms, which makes its eradication more difficult. The SARS-CoV-2 envelope includes attached proteins like M (membrane), S (spike), E (envelope), and N (nucleocapsid). Similar to other coronaviruses, the N protein of SARS-CoV-2 inhibits IFN1 by regulating IFN-β synthesis and signaling. On the other hand, the effectiveness of the innate immune response against viral infection depends mainly on IFN1 production and its downstream signaling that results in controlling viral replication and induction of an adequate adaptive immune response [7,20]. However the virus could avoid this attack due to the complex immune dysregulation caused by this infection. Chronic stimulation of T cells, resulting in a cytokine storm and T cell exhaustion, weakens the overall body defenses and puts the patient in a dangerous situation. High-grade chronic viral infections result in CD8+ T cell exhaustion (Exh) leading to a decreased effector function and lower proliferative capacity. Tex leads to over-expression of inhibitory receptors, including CD279 (PD-1), a lymphoid cell surface protein of the Ig superfamily, and a member of the extended CD28/CeCLA-4 family of T cell regulators, which act as a mature T cell checkpoint for the modulation of apoptosis. PD-1 can bind to either of its ligands (PD-L1[CD274] and PD-L2[CD273]) both members of B7 family of T cell co-receptors. This binding causes significant suppression of the immune system by affecting T cells, as well as B cells and NK cells [7,20,24,25]. Another important observation is the strong correlation between inflammatory markers, including ESR, CRP and IL-6, and the relevant subset of lymphocytes [26]. Overall, general lymphopenia is seen in COVID-19 patients, especially in severe cases [27,28].

2.4. Disease in some particular groups of patients

Similar to all infectious diseases, the immune system plays an essential role in virus suppression. Therefore, it can be assumed that suppression of the immune defense system will make the situation worse, but it is not as simple as it sounds [29]. There has not yet been enough scientific evidence to generally employ immunosuppressive drugs in autoimmune diseases like rheumatoid arthritis (RA) [30,31]. On the other hand, the hyper-inflammatory and cytokine release syndrome (CRS) typical of COVID-19 causes tissue damage to the lung epithelium and ARDS [32]; therefore, immunosuppressive drugs may be useful as there is some evidence that an anti-IL-6 approach is effective in critically ill patients in the ICU [33]. Another high-risk group for COVID-19 is cancer patients, who are administered cytotoxic drugs. Although their immune function may be at an acceptable level, some cytotoxic drugs such as 6-MP used in chemotherapy, have shown destructive effects on virus replication in vitro [31]. On the other hand, the high number of critically ill patients and increased mortality in patients with underlying diseases (such as hypertension and diabetes) has been proven [34]. Diabetes mellitus type II can include a hyperglycemic condition of diabetic patients can be worsened after COVID-19 infection. Chronic stimulation of T cells, resulting in a cytokine storm and T cell exhaustion, weakens the overall body defenses and puts the patient in a dangerous situation. High-grade chronic viral infections result in CD8+ T cell exhaustion (Exh) leading to a decreased effector function and lower proliferative capacity. Tex leads to over-expression of inhibitory receptors, including CD279 (PD-1), a lymphoid cell surface protein of the Ig superfamily, and a member of the extended CD28/CeCLA-4 family of T cell regulators, which act as a mature T cell checkpoint for the modulation of apoptosis. PD-1 can bind to either of its ligands (PD-L1[CD274] and PD-L2[CD273]) both members of B7 family of T cell co-receptors. This binding causes significant suppression of the immune system by affecting T cells, as well as B cells and NK cells [7,20,24,25]. Another important observation is the strong correlation between inflammatory markers, including ESR, CRP and IL-6, and the relevant subset of lymphocytes [26]. Overall, general lymphopenia is seen in COVID-19 patients, especially in severe cases [27,28].

3. The immune system as a foe

3.1. An entry route for the virus

CD147, also known as Basigin or extracellular matrix metallopeptinase inducer (EMMPRIN), is also recognized as a red blood cell (RBC) receptor for the parasite that causes malaria in humans, and is a transmembrane protein of the immunoglobulin family. Its expression is induced in several conditions such as asthma, cancer stem cells, high glucose concentration in monocytes, and inflammatory processes [39].
Wang et al. recently demonstrated that the SARS-CoV-2 SP also binds to CD147 [40]; so, the immune system itself could be an entryway for SARS-CoV-2.

### 3.2. Macrophage activation syndrome and the cytokine storm

Patients with COVID-19 have increased levels of inflammatory cytokines and chemokines, such as IL-1, IL-6, IL-8, IL-17, IL-17, CCL-2, TNF-α, G-CSF, IP-10, MCP-1, and MIP. The concentration of these markers fluctuates depending on the individual's condition; it seems that increased cytokine levels, especially IL-6, have a direct association with a worsened patient condition [41]. Furthermore, as higher levels of cytokines rapidly lead to deterioration of the patient's condition and death, they could be considered to be prognostic markers in the clinic [42]. The cytokine storm (CS) is typical of macrophage activation syndrome (MAS) or secondary hemophagocytic lymphohistiocytosis (sHLH). Consequently, tissue damage, lung injury and acute respiratory distress syndrome (ARDS) could be expected [43]. Furthermore, a study showed that the peripheral blood of a patient with severe COVID-19 had a strikingly high number of Th17 cells, which secrete IL-17, and are associated with autoimmune and inflammatory diseases. In the mucosal immune response, IL-22, IL-17, and TNF-α are known to induce antimicrobial peptides. Also, IL-22 upregulates mucins, fibrinogen, and anti-apoptotic proteins; therefore, IL-22 may contribute to the formation of life-threatening edema, and the lungs may become enriched with mucins and fibrin leading to the progression of ARDS, seen in COVID-19 patients [44]. All of the above-mentioned factors result in pneumonia and ARDS, besides hyper-ferritinemia, coagulopathy, and multi-organ failure in the liver, heart, and kidneys with elevated D-dimer, CRP, BUN, and Cr that are typical of MAS/sHLH [27,45]. However, NK cell reduction, suppression of antiviral defense, activation of aggressive tissue-damaging immune responses via increased IL-6 secretion, and secondary CS leads to a general picture of hyper-inflammation and severe respiratory failure [27,46]. Although the typical MAS/sHLH pathology that occurs in immunocompetent patients, may also be seen in COVID-19, but the differentiation between these two patterns is difficult.

### 3.3. Hyper-inflammation and severe respiratory failure

Although COVID-19 has numerous manifestations, lung injury and severe respiratory failure (SRF) are more common than others. The cornerstone of this condition is MAS/sHLH or immune dysregulation. Alveolar macrophages secret IL-6 that cause overproduction of pro-inflammatory cytokines by monocytes, and dysregulation of lymphocytes, characterized by CD4 lymphopenia and subsequently B cell lymphopenia. In parallel, the absolute natural killer (NK) cell count is depleted, probably as a result of the rapidly multiplying virus. Moreover, IL-6 decreases the expression of HLA-DR on the membrane and production of IFN-γ by CD4 cells [4]. In conclusion, it seems that innate immunity, characterized by neutrophils and proinflammatory cytokines in the format of the cytokine storm, is trying to limit the infection and overcome the virus; however, it leads to an excessive inflammatory response rather than harming the virus. This causes lung injury with tissue damage and life-threatening edema, enriched with mucins and fibrin, that unfortunately causes SRF and death in severely affected patients [22].

### 4. The immune system as a friend

#### 4.1. Role of immunoglobulin and immunomodulation in recovery

It seems that lymphopenia, hyper-inflammatory state, and cytokines all play a role in the pathology of COVID-19; hence the hypothesis was proposed that immunomodulatory drugs may be beneficial to reverse the condition and treat the disease [47]. Considering the pivotal role of IL-6 in CS, tocilizumab as an IL-6 inhibitor could be effective in COVID-19 patients with a severe condition [33]. There are limited data about the effects of other immunosuppressive drugs, such as glucocorticoids, IL-1 inhibitors, mycophenolate mofetil, anti-TNF-α agents, methotrexate, and NSAIDs in COVIS-19, and further in vitro and in vivo studies are recommended [31]. Due to the T cell lymphopenia and reduced function, immunoglobulins produced by B cells represent the main arm of the immune system to combat the virus. Serocconversion and antibody production occurs by the first two weeks after infection [48]. Like other infections, specific IgM is the first defense to appear, and disappears after a short time, but specific IgG remains a long-term defense against the virus. Therefore, neutralizing IgGs play a major role in the patient recovery and control of infection. IgGs reach their peak in the serum during the convalescent phase, and tend to wane after recovery, but memory B cells could still survive to offer long-term protection [22,49].

#### 4.2. Convalescent plasma therapy

There is a global effort to find an effective treatment for COVID-19. Although the benefits of some available drugs have been proven, they are not yet a certain specific cure for the disease, and controversies remain [50]. According to previous experience with other coronavirus family members, convalescent plasma therapy has been tested on some critically ill patients, with promising results. In this regard, several studies have confirmed the positive effects of this treatment, especially in critically ill patients in the ICU and with ARDS to shorten hospitalization time and reduce the mortality rate. The COVID-19 specific IgG antibodies act as passive immune therapy, and after administration by the transfused plasma might neutralize viral particles and activate the complement system, which could consequently lead to viral elimination. Despite its efficacy, this method has some difficulties such as concerns about causing an infection, and non-infectious transfusion reactions. Transfusion-induced infection is rare in industrial countries (unlike poor countries) while the latter is a global worry known as transfusion-related acute lung injury (TRALI) and allergic reactions. As an important point, IgG collected in one country may be different from another country, because lifestyle, diet, genetics, and the environment could play an important role in the development of specific antibodies against the virus. Furthermore, differences between strains of coronavirus should be considered in geographically distinct areas; so, treatment of infected cases with polyclonal IgG, collected from the same virus is much lower than found in previous pandemics. What makes COVID-19 so significant and dangerous, is the rapid acceleration of the virus transmission. Despite all efforts to control and treat this disease, public quarantine and lockdown measures cannot be implemented forever, and all tested treatment methods have not been sufficiently effective. Therefore, along with trying to find more effective specific drugs, the effort to find a vaccine should be redoubled in order to gain herd immunity and allow the world to return to normal life and resume its routine. Because the process of vaccine preparation and testing is complicated and lengthy, validating and marketing vaccines can be tedious, but given the body of work by various research groups, there is
a clear prospect of achieving this goal sooner rather than later [20, 58, 59].

5. Conclusion

The COVID-19 pandemic is an ongoing issue that affects the lives of most people around the world. Most countries are now semi-closed, strict travel regulations have been enacted, international relations have been affected, and humans are experiencing an unprecedented regime, which has changed ordinary life [60]. Therefore, it is of the utmost importance to understand the pathophysiology of disease and how the immune response to the pathogen affects the disease. Although the immune system plays an important role in fighting COVID-19, paradoxically it could also be harmful. Most critically ill patients in ICU, that develop ARDS, have high levels of inflammatory cytokines in their circulation, known as CRS. Considering all the reported data from observations and measurements, it seems that when the immune system is severely damaged and becomes inefficient by lymphopenia and Tcx, it tries to compensate by triggering the CRS, which could potentially lead to complications like ARDS and multi-organ failure. It is necessary to find efficient drugs and vaccines to return to the normal situation and reduce the mortality rate.

Author contribution

Both FY and NR had role in concept and design of the paper. FY drafted the manuscript, while NR and MH critically revised it. All authors approved the final draft of manuscript.

Declaration of competing interest

The authors declare that they have no conflicts of interest.

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