Abstract: The purpose of this review is to address some of the latest aspects regarding molecular features, pathogenic mechanisms, and immune system response to severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), based on recent publications in this field from March 2020 to May 2021. Interpretation keys for periodic re-emergence of coronavirus infections and other lethal viral pandemics are suggested. Antibody-dependent enhancement (ADE) and other potential mechanisms of immune system deception are put forward. Therefore, vaccine development must take into account ADE and other unwanted side effects of immune-based medical intervention. Features reported in our review will allow both clinicians and basic science researchers to take home ideas to improve their knowledge about SARS-CoV-2.

Keywords: SARS-CoV-2; cytokine storm; immune system; antibody-dependent enhancement (ADE)

1. The New Coronavirus: Molecular Proofs

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is the third documented example of animal coronavirus transmission to humans in the last two decades, following the severe acute respiratory syndrome coronavirus (SARS-CoV) in 2002 and the Middle East respiratory syndrome coronavirus (MERS-CoV in 2012) [1]. All three coronaviruses (CoVs) can spread from person to person [2]. SARS-CoV-2 is associated with a severe disease called coronavirus disease 19 (COVID-19) [3]. SARS-CoV-2 is an enveloped virus with positive single-stranded RNA, coding for four structural proteins: spike (S), envelope (E), membrane (M), and nucleocapsid (N). The S protein comprises two functional subunits. The S1 subunit is responsible for binding to the host cell receptor, while the S2 subunit determines fusion of the viral and cellular membranes [4].

Circulation of SARS-CoV-2 infection has had a variable impact worldwide, determined by implemented lockdown measures (i.e., social distancing and travel restrictions), public health system, socioeconomic status and lessons learnt from previous outbreaks [5]. Prevalence and confirmed cases of SARS-CoV-2 have been continuously changing in different regions around the world. However, the present health emergency has been strictly monitored and it is possible to follow the progression of the pandemic in real-time. In the globalization era, the immediate availability of epidemiological data and genomic sequences of the new virus, collected worldwide, help to monitor the diversity of emerging virus isolates and to assist public health policy [6].
Molecular and phylogenetic analyses can estimate genetic variability in real-time, eventually identifying specific genetic variants associated with different clinical outcomes and disease prognoses. When the number of infected people increases, viruses become more genetically distinct, and natural selection pressure makes them more likely to fit with the host [7]. SARS-CoV-2 intra/inter-host genetic variation analysis detected the same minority variants by deep-sequencing, suggesting a predisposition to specific mutations and a subsequent revival of the epidemic [7]. If the evolutionary rate of SARS-CoV-2 is slower than its transmission rate, then several identical genomes are rapidly spreading. Accordingly, genetic similarity places limitations on the ratio of imported cases to local transmission calculation. However, due to this slow rate of virus mutation, even one difference can provide enough information if appropriate molecular methods are used [8].

At present SARS-CoV-2 is classified into viral variants, such as Variant of Concern (VOC) and Variant of Interest (VOI). The WHO has provided working definitions of VOCs and VOIs, considering changes that lead to an altered phenotype, which are harder to reflect in genotype-based classifications [9]. The VOCs have been associated with one or more of the following: in vitro evidence for an increase in transmissibility and virulence, change in epidemiology and clinical disease presentation, decrease in effectiveness of public health measures, vaccines and therapeutics. On the other hand, VOIs show genetic changes to receptor binding sites, reducing neutralization by antibodies (from previous infection or vaccination), and efficacy of treatments thereby increasing transmissibility and disease severity [9,10].

During the first pandemic wave, SARS-CoV-2 strains were classified into two subtypes (a and b) according to synonymous mutations (D614G) in the spike gene by phylogenetic analysis. Since this amino acid change is located in the S1 domain (B-cell epitope), it may play a crucial role in virus evasion of the immune system. Indeed, SARS-CoV-2b strains (exhibiting D614G) reduce antigenic indices compared to the SARS-CoV-2a subtype [11]. More than one year after the first confirmed case of COVID-19, SARS-CoV-2 infection around the world is maintained by several lineages, such as B.1, B.2, B.3, B.4, and VOCs (i.e., B.1.1.7, B.1.351, P1, B.1.617.2, and so on) (Table 1).

Table 1. SARS-CoV-2 variants and related substitutions.

| WHO Label | First Identified | Pango Lineage | Spike Protein | ORF1ab Protein |
|-----------|-----------------|---------------|---------------|----------------|
| **VOCs (Variants of Concern)** | | | | |
| Alpha | United Kingdom | B.1.1.7 | 69del, 70del, 144del, E484K, S494P, N501Y, A570D, D614G, P681H, T716I, S982A, D1118H K1191N | None |
| Beta | South Africa | B.1.351 | K417N, E484K, N501Y | None |
| Gamma | Japan/Brazil | P1 | K417T, E484K, N501Y | None |
| Delta | India | B.1.617.2 | 19R, G142D, 156del, 157del, R158G, L452R, T478K, D614G, P681R, D950N | None |
| Epsilon | California | B.1.427, B.1.429 | S13I, W152C, L452R, Q677H | I4205V (ORF1a); D1183Y (ORF1b) |
| **VOIs (Variants of Interest)** | | | | |
| Eta | United Kingdom/Nigeria | B.1.525 | E484K, D614G, Q677H | None |
| Theta | The Philippines | P3 | E484K, N501Y, D614G, P681H | None |
| Iota | United States (New York) | B.1.526 | L5F, D80G, T95I, Y144L, F157S, D253G, L452R, S477N, E484K, D614G, A701V, T859N, D950H, Q957R | None |
| Kappa | India | B.1.617.1 | T95I, G142D, E154K, L452R, E484Q, D614G, P681R, Q1071H | None |
| Lambda | Peru | C.37 | L452Q, F490S, D614G | None |
| Mu | Colombia | B.1.621 | T95I, Y144S, Y145N, 146N insertion, R346K, E484K, N501Y, D614G, P681H, D950N | None |
VOCs harbor substitutions at the receptor-binding site (RBD) of the S protein that impact on viral fitness, transmissibility, and immunity [12]. According to the latest European Centre for Disease Prevention and Control (ECDC) report released on 3 June 2021, VOCs are all more transmissible than the wild-type strain. The emergence of novel variants and their spread into new countries must be constantly monitored. The B.1.1.7 (alpha), B.1.351 (beta), and P.1 (gamma) variants possess substitutions, such as N501Y, E484K, and K417N/T capable of increasing transmissibility, pathogenicity, and ability to escape immunity [13]. B.1.617.1 (kappa) and B.1.617.2 (delta), first detected in India, carry the main substitutions, L452R, E484Q, D614G, and P681R [14] (Table 1). In particular, P681R, which facilitates spike protein cleavage and enhances viral fusogenicity, was reported as responsible for increased spread and pathogenicity [15]. However, the B.1.617.1 (kappa) variant is less efficient at infecting host cells [16].

Adaptive evolution of SARS-CoV-2 suggests that specific patterns of changes in the RBD domain, N and S proteins confer a selective advantage within a broader epistatic context [17]. In particular, B.1.351 (beta) and B.1.617.2 (delta) variants clearly escape neutralization by convalescent plasma and vaccines are likely to be less effective against them [18]. B.1.1.7 (alpha) and B.1.617.2 (delta) variants previously linked to limited cases or outbreaks have become more widespread due to genomic substitutions influencing cellular binding and viral replication [12]. Currently, according to epidemiological data, B.1.617.2 (delta) is the most prevalent variant worldwide and at least 40% more transmissible than B.1.1.7 (alpha) [9]. Very recently, the B.1.621 (mu) variant was identified for the first time in Colombia, South America. On 30 August 2021, it was classified as a VOI by the WHO. Similar to beta and delta, the mu variant is associated with increased transmissibility and decreased antibody responses, following both recovery from COVID-19 disease and vaccine administration [9].

SARS-CoV-2 appears to be less deadly but more transmissible than other coronaviruses. This virus does not kill the host quickly, therefore allowing its dissemination. It is possible that genomic mutations selected during this outbreak might reduce SARS-CoV-2 virulence and transmissibility as previously reported for SARS-CoV [19]. In any case, increasing the number of sequences available in public databases, will allow researchers to better understand the evolution and direction of the new coronavirus infection.

2. Pathogenesis and Different Organ Involvement

Unlike other respiratory viruses, CoVs are well-known to attach to the parenchymal region of the lung, besides the perihilar and stromal areas. This feature is very particular and should draw the attention of researchers to explain this using cellular and molecular mechanisms. The study by Zhou and co-workers has clarified this aspect by demonstrating that SARS-CoV-2 uses the angiotensin-converting enzyme 2 (ACE2) as a specific receptor among the alveolar cells, namely type II alveolar cells (AT2) [20]. Likewise, the pathway followed by SARS-CoV-2 to arrive in the lung alveoli has been elucidated in a recent paper evaluating a particular type of SARS-CoV-2 trans-membrane protease on bronchial cells, TMPRSS2 [21].

In its more severe forms, the clinical course of infection due to the new coronavirus shares many features with sepsis [22]. Among patients who display symptoms, the disease is mild in about 80% of cases, while it is severe or critical in 14% and 5% of patients, respectively [23,24]. Mild disease is characterized by the presence of a flu-like syndrome with fever, fatigue, dry cough, and myalgia. When dyspnea and hypoxia occur, the disease can be considered severe. It becomes critical when acute respiratory distress syndrome (ARDS), shock, or multiple organ dysfunction syndrome (MODS) are presented. Although the three most common symptoms of COVID-19 are fever (99%), fatigue (70%), and dry cough (59%) [25–27], it has also been reported that the presence of fever varies according to patient characteristics. Indeed, fever is often more frequent during hospitalization compared to actual hospital admission (89% vs. 44%) [28]. Other symptoms, such as choriza, headache, and enteric manifestations (nausea, vomiting, and diarrhea) may be
present, even if in a minimal proportion of cases [27,29]. The occurrence of olfactory and gustatory dysfunctions has been recently reported as frequent symptoms in COVID-19 patients. In a study of 417 patients diagnosed with COVID-19, olfactory and gustatory dysfunctions were reported in 85.6% and 88.0% of patients respectively, with a significant association between both disorders [30]. Finally, neurological manifestations, such as acute cerebrovascular diseases, impaired consciousness, and skeletal muscle injury, have also been reported, especially in patients with a more severe clinical course of infection [31,32].

Aside from this clinical scenario, the association between the many sepsis-like mediators and the severe/late course of SARS-CoV-2 infections is intriguing. Such severe cases have been reported to be related to the enhancement of leucocyte and granulocyte numbers and increased concentrations of D-dimer [33].

It has been demonstrated that the expression of human ACE2 in different organs and tissues would be of relevance regarding clinical susceptibility and prognosis during SARS-CoV-2 infection. The single-cell RNA-seq analysis data suggest that Asian people showed a much higher ACE2 expression cell ratio than Caucasian and African-Americans. This experimental finding supports the different SARS-CoV-2 susceptibility among various human races and may account for the higher sensitivity of Asian people to the new coronavirus [34].

Although the apparent target of SARS-CoV-2 is the lung, research, from the beginning, has made evident the role played by different extra-respiratory tissues and organs, particularly in cases with a more severe clinical course of infection. The digestive tract and liver of humans may be involved in the spreading of this virus within the human body. Recently the putative receptor (ACE2) for SARS-CoV-2 has been found on bile duct epithelial cells. Therefore, an inflammatory response of cholangiocytes and the compensatory proliferation of cholangiocyte-derived hepatocytes may underline the potential mechanism of liver injury by SARS-CoV-2 [35,36]. Accordingly, the presence of SARS-CoV-2 in fecal samples has been recently reported [37].

Such findings have different implications. First, the fecal-oral route of viral transmission could be as crucial as the aerial transmission. Second, the delayed colonization of the digestive tract may suggest a novel pathogenic mechanism and finally, novel clinical and virological assessment (RT-PCR on fecal samples) might be needed in order to follow the natural history of human infection by SARS-CoV-2 [37].

Central nervous system (CNS) involvement has also been recently reported in SARS-CoV-2 infection. Neurological symptoms and signs such as: nausea, vomiting, headache, and reduced consciousness without apparent alterations in head CT scans suggested a diagnosis of encephalitis in some SARS-CoV-2 patients. Molecular tests were carried out on cerebrospinal fluid of some such patients, and SARS-CoV-2 was identified. Coronavirus has been found in the brainstem of both humans and experimental animals. It has also been reported that coronavirus can spread to the medullary cardiorespiratory center from peripheral chemoreceptors and from the lungs through both the blood and synapse routes. This feature might contribute to acute respiratory failure in the most severe cases of SARS-CoV-2 infection [11,38].

The spread of COVID-19 via blood circulation or across the cribriform plate of the ethmoid bone can lead to brain involvement. Such a feature has been reported for SARS-CoV patients. Subsequent budding of the virus from the capillary endothelium and slight injury to the endothelial lining can allow access to the brain. Subsequently, virus interaction with ACE2 receptors found in neurons and glial cells, may then lead to a cycle of viral budding with neuronal damage in the absence of an apparent inflammatory reaction.

Two significant clinical features of the early stage of COVID-19 are hyposmia and ageusia. There is a sharp contradiction regarding the mechanisms of these two symptoms. Some investigators have reported that a change in the mucosal cells of the nasal district might account for such a feature. Indeed, it is very unusual for such a change of feeling, without any overt inflammation and rhinorrhea.
The other possible explanation includes a primary effect of SARS-CoV-2 on the nerve fibers and cells. Many nervous regions, like the raphe nucleus (mesencephalon) and some nuclei of the lobus limbicus, as well as basal nuclei, are significantly involved in olfactory sensitivity. Similarly, ageusia might be explained by an effect on the tongue mucosal cells. Indeed, saliva is a matrix rich in SARS-CoV-2. Moreover, the nervous fiber of peripheral nerves involved in gustatory sensation could be substantially impaired [30,39]. Cerebral involvement alone, with the potential for causing cerebral edema, or small bleeding along with the cerebral microcirculation during COVID-19, may lead to death even before systemic homeostatic deregulation or lung diffuse consolidation. Moreover, entry of SARS-CoV-2 to the central nervous system via the transcriptional route has been reported for other CNS targeting pathogens. Such a route may explain a clinical COVID-19 with hyposmia and the cases of acute respiratory failure in COVID-19. Moriguchi described the first case of meningitis/encephalitis associated with SARS-CoV-2 not detected by nasopharyngeal swab, but confirmed in cerebral-spinal fluid [40].

There is no data which confirms that pregnancy increases the susceptibility of women to COVID-19, and it is still unknown if pregnancy could predispose to a more severe clinical course [41]. Clinical presentations in pregnant women are very similar to non-pregnant ones, and a small case series of women in their third trimester of pregnancy reported no adverse outcomes after delivery [42]. However, some authors suggest performing bimonthly fetal growth monitoring for the possibility of intrauterine growth restriction [43]. One explanation of such findings may be the lack of ACE2 receptors in placental tissue. This is consistent with the lack of SARS-CoV-2 in amniotic fluid, cord blood, and neonatal throat swabs [44].

In children and new-borns, COVID-19 seems to be mild and complications rarely occur [45]. Most children diagnosed with COVID-19 had a benign clinical course, with fewer and mild symptoms [46–48]. Reasons why children are not likely to present severe disease are still unknown, although the particular characteristics of the immune system and lower maturity and function of ACE2 receptors in children compared to adults have been suggested as possible underlying mechanisms [49].

3. Immune System Deception by SARS-CoV-2

The S protein has a well-defined composition comprising 14 binding residues that directly interact with human ACE2 receptors. Of these amino acids, eight are conserved in SARS-CoV-2. However, particular changes in some amino acids between S proteins of SARS-CoV and SARS-CoV-2 have been correlated with the enhanced binding affinity of SARS-CoV-2 and ACE2 receptor vs. SARS-CoV [46,50]. The mutations responsible for the greater transmissibility and the corresponding VOCs/VOIs are shown in Table 1.

The S protein binds to CD147, a trans-membrane protein of the immunoglobulin family, known as an entry route in red blood cells for the Malaria parasite. Such a protein could also be used by the new coronavirus [51]. These features may explain the increased human-to-human transmission exhibited by SARS-CoV-2 [46,50].

It has been demonstrated that increased levels of serum IgG antibodies could be proportional to the viral load and enhanced disease severity [52]. SARS-CoV-2 viral load, innate immunity and the several viral replication steps of COVID-19 are of vital importance to better understand the disease scenario. This could help to clarify why some patients remain asymptomatic, while other patients (where SARS-CoV-2 invades the alveoli and replicates in type II pneumocytes) may exhibit pneumonia, often with a fatal outcome [33]. Innate immunity is crucial to control early viral replication before the immune system can generate an effective adaptive immune response and also to establish this adaptive immunity. Innate immunity against SARS-CoV-2 is similar to that against other pathogens and is based on the release of some humoral components and the action of different cellular elements. It would be intriguing to investigate the possible role of endogenous antibiotics, such as magainins, against SARS-CoV-2 in both in vitro and in vivo models [53]. The presence of such innate immunity mediators could account
for the significant and timely effect of convalescent plasma transfusions on the severe complications of COVID-19, even in the presence of non-neutralizing antibodies in the transfused plasma [54]. Even Toll-like receptors (TLRs) play a role in pathogenesis of SARS-CoV-2 infection. This feature has been associated with a signal mediated by TLR3, TLR7, or TLR8 leading to immunopathology [55].

Coronaviruses causing severe epidemics may disappear several months after causing deadly pandemics, only to re-emerge after some years to generate another outbreak [56]. Antibodies are primary tools in the protection against respiratory viruses [57]. Historical data regarding periodical re-emergence of the most critical viral respiratory epidemics show undoubtedly that the most studied are the flu epidemics [58]. The study of flu epidemics began with the 1889–1892 Russian Flu, followed by the Spanish Flu (1918–1920). The latter deadly epidemic has recently been analyzed based on the evaluation of antibody titer in survivors and on the direct demonstration of the strong neutralizing effect of such antibodies in an in vivo flu murine model [57]. Strong systemic cytokine reactions of remarkable pathogenic significance followed in this elegant in vivo experimental model of Spanish Flu, reminding us of the immunopathogenesis associated with the most severe COVID-19 clinical cases.

Similarly, in 2002–2003 the first relevant pandemic by a coronavirus causing respiratory pathology, was able to spread over 26 countries with 8096 human cases and 774 deaths [59]. A similar respiratory epidemic emerged again in 2012 spreading over 25 countries [59]. These pandemics have been followed by the present COVID-19 pandemic, which has so far caused more than 170,000,000 cases and more than 3,000,000 fatal outcomes. One traditional interpretation of such periodic behavior of these pandemics is reported as cyclical theory. The lack of population immunity to the previous pathogen would account for the periodicity of such epidemics. Indeed, such immunity would only last while most of the survivors are still alive [60].

However, the reappearance of an epidemic caused by respiratory viruses after only 8–10 years since the previous outbreak, is in contrast with cyclical theory. In this case, a different interpretation must be taken into account to explain such a re-emergence of the same epidemic after such a short period. Dealing with the historical behavior of several flu pandemics, some authors state that previous exposure to the influenza virus has paradoxically enhanced, instead of decreasing, susceptibility to such viral disease [60].

Two possible explanations are reported to account for a second more severe infection based on the same viral etiology of the first one (Figure 1).

The first is explained by the antibody-dependent enhancement (ADE) theory, where antibodies specific to the etiological agent, causing the first epidemic are released. ADE is the phenomenon in which preexisting, non-neutralizing, or poorly neutralizing antibodies lead to enhanced infection, increasing viral entry into cells. The mechanism of ADE can also occur due to a lower concentration or lesser affinity of neutralizing antibodies. Several factors are involved in ADE such as affinity, concentration, specificity, and isotype of antibodies. It is interesting how high-affinity neutralizing antibodies, even at minimal concentrations, can avoid ADE. In clinical settings, ADE has been reported in several human viruses (HIV, Dengue virus, RSV, Ebola virus, Zika virus, Influenza virus) and veterinary pathogens [55]. On the contrary, according to our knowledge, direct scientific evidence of a clinical impact of ADE during human SARS-CoV-2 infection is lacking. Similarly, ADE emerging after SARS-CoV-2 vaccine administration in humans has not yet been published.

According to Tillett et al., previous exposure to SARS-CoV-2 might not guarantee total immunity. Their findings led to the hypothesis that a mechanism of antibody-dependent enhancement might be the cause of a symptomatic reinfection with SARS-CoV-2 in a 25-year-old man [61].
Figure 1. Schematic view of Immune System role during SARS-CoV-2 infection. The antibody-dependent enhancement (ADE) theory is shown in blue, the antigenic imprinting in orange.

Focosi et al. collected a number of demographic and clinical data of their patients and evaluated the level of antibodies against antigens of both minor coronaviruses and SARS-CoV-2. Statistical association between the levels of anti-minor coronavirus antibodies and worse clinical outcome was demonstrated. They suggested a possible ADE during SARS-CoV-2 infection due to previous alphacoronavirus (minor coronavirus) immunity and they found that previous antibodies to the seasonal endemic coronaviruses could be associated with a worse clinical outcome in COVID-19 patients [62].

Dengue virus and other viral organisms have been demonstrated to cause ADE. It is a general view that host antibodies play a beneficial role during all viral diseases. Previous studies on SARS-CoV-2 infection showed the peculiar mechanism used by the virus to convert immune cells, such as macrophages and B cells, into weapons against the host. Such a mechanism envisages that SARS-CoV-2-specific antibodies, after engagement with coronavirus or its product, could link receptors specific for the Fc fragment of the immunoglobulin molecule. Such interaction could produce both inflammatory reactions and persistent viral replication in patient tissues [63–65]. The formation of immune complexes might lead to the cytokine storm particularly in severe COVID-19 [66]. Some cytokines (IL-1β, IFNγ, IL-8, and IL-4) have been investigated concerning such an atypical viral pathogenic mechanism. Indeed, those patients infected with SARS-CoV-2 were found to exhibit a high concentration of IL-1β, IFNγ, and IP10 (CXCL10), leading to a T-helper-1 (Th1) cascade. This unusual viral pathogenic mechanism has also been reported as a “cy-
"cytokine storm" [29]. Two relevant aspects should be underlined regarding the cytokine role in SARS-CoV-2 infections. High levels of chemokines like IP10, MCP1, MIP1α, and relevant concentration of the pro-inflammatory cytokine TNFα found in the severe cases, might explain the intense tissue inflammation and massive consolidation of large lung areas. Moreover, Th17 cells, neutrophils, and granulocytes all produce IL-17, which stimulates production of several pro-inflammatory cytokines [51].

On the contrary, the release of Th2 and Treg cytokines (IL-4 and IL-10), with an anti-inflammatory potential, might account for the many mild cases of SARS-CoV-2 infections [29,67]. Regarding the early release pro-inflammatory cytokines, IL-6 has been reported to be among the main components of the cytokine storm [67].

It is a relevant issue for viral infections that the internalized immune complexes modulate host immune response enhancing viral replication and aggravating disease severity [68]. In the context of SARS-CoV-2, ADE may enhance viral replication in immune cells since the virus might productively infect immune cells including monocytes and B cells both in vivo and ex vivo. Wu et al. studied in vitro ADE of SARS-CoV-2 entry into FcγRII receptor-bearing cells (Raji cells are lymphoma cells derived from human B lymphocytes) by plasma and antibodies from patients who recovered from COVID-19 [69]. The enhancement of SARS-CoV-2 infection was mediated by IgG and it was more commonly observed in plasma from elderly patients with critical conditions and prolonged disease duration, suggesting a possible association of ADE with worse clinical outcomes in COVID-19 patients [69].

Vaccine development needs to take into account the serious possibility of any potentially dangerous effect which triggers an ADE phenomenon. Due to the very large population that must be vaccinated, the very frequent spread of minor coronavirus infections (HCoV-229E, -NL63, -OC43, -HKU1) throughout the general population should be considered. Such spread probably allows several populations of anti-coronavirus antibodies to be maintained at very low levels for a long time. Thus, the potent trigger of a SARS-CoV-2 vaccine might provoke an uncontrolled ADE with hazardous side effects for the host. Any vaccine must therefore avoid ADE and other unwanted side effects.

A vaccine inducing high titers of neutralizing antibodies is considered safer than one inducing low titers because the new virions are neutralized before the ADE occurs. Furthermore, neutralizing antibodies mediate ADE only at the suboptimal neutralizing concentration. Plasma and antibodies from patients who have recovered from COVID-19 should be tested for potential ADE effects before clinical usage [69]. The development of a vaccine or immunotherapeutic agent targeting only the neutralizing epitopes represents an optimal approach to induce a better immune response reducing the possibility of ADE and hence protecting the host from COVID-19 [55].

Alongside the complex ADE phenomenon, another interpretation which may account for a second more severe infection based on the same viral etiology of the first one is antigenic imprinting. This hypothesis proposes a reduced host immune response against a respiratory virus when such a pathogen comes into contact with the host for the first time early in life. The trend of human immune system B cells and T cells to overly trust on previous/first contact with any given pathogen, leads to ineffective short-cuts of immune cells functions which eventually would not be able to eradicate the pathogen [58,70].

From the ideas reported above, one may hypothesize that the first time a coronavirus or another primary respiratory virus spreads in a human population, a small and limited epidemic would result. However, a second hit, one or two decades later, would cause a deadly pandemic.

The risks due to the ADE phenomenon associated with SARS-CoV-2 have important implications for COVID-19, MIS-C treatment, B-cell vaccines, SARS-CoV-2 antibody therapy, and convalescent plasma therapy for patients. ADE risk for SARS-CoV-2 B-cell vaccines for different subsets of populations, can be associated with various clinical and demographic characteristics such as age, cross-reactive antibodies, variability of antibody...
levels over time and pregnancy. These new studies place greater emphasis on the need to develop safe SARS-CoV-2 T-cell vaccines that will not be dependent on antibodies [71].

4. Conclusions

The COVID-19 deadly pandemic offers researchers the chance to shed light on many aspects of SARS-CoV-2 and possibly of similar viruses. This virus took advantage of some out-of-the-ordinary social and behavioral aspects within cities with a very large population, during a period of the year when the majority of the community was in transit due to the beginning of the Chinese lunar calendar. In modern times, infectious organisms can exploit the possibility of infecting a large part of the world population due to year-round travel, by air and by cruise ship. On the contrary, until 30 or 40 years ago, very few people could have afforded such frequent trips for either work or leisure. SARS-CoV-2 exploited the quickest and simplest transmission pathway, airborne, instead of slower and less efficient diffusion mechanisms (by contaminated food, by a parenteral route, by specific vectors). Such different and highly efficient transmission patterns of viruses must be thoroughly investigated to avoid other infectious agents from replicating the COVID-19 pandemic. In Table 2 we suggest the main points of SARS-CoV-2 infection for attention.

Table 2. Take home messages.

| Number | Text                                                                 | Related References       |
|--------|----------------------------------------------------------------------|--------------------------|
| First  | Mutations in SARS-CoV-2 strains could help the virus elude the immune system response | Cherian S. et al., 2021 [13] |
| Second | The cytokine storm during SARS-CoV-2 infection is closely related to expression of ACE2 in different organs and tissues, shaping clinical susceptibility and prognosis. | Cao Y. et al., 2020 [34] |
| Third  | The ADE phenomenon and/or antigenic imprinting could explain the immune system deception by SARS-CoV-2. | Fu Y. et al., 2020 [65], Roncati L. et al., 2020 [58] |
| Fourth | Development of safe SARS-CoV-2 T-cell vaccines, that are not dependent on antibodies, is necessary | Ricke D.O. et al., 2021 [71] |

Researchers should address the life cycle and biology of SARS-CoV-2, and the molecular mechanisms used to initiate and develop infection in the human host. Such illness begins in the upper airways, but may rapidly progress to the lung tissue and might even spread to several different vital organs, including the brain, liver, heart, kidney, and possibly others. The efficacy of currently available therapeutic strategies against COVID-19 is not yet established. However, different clinical trials are ongoing and preliminary data may be useful in guiding clinicians toward the best management of patients suffering from COVID-19.

Finally, even if the ADE phenomenon is not well addressed in SARS-CoV-2, it is necessary to take into account the side effects of uncontrolled ADE. Hence, investigators should identify effective and safe vaccine strategies to prevent the further spread of this infection and hopefully stop this pandemic.

Future studies dealing with the COVID-19 immune response should address in depth the cellular response of host immunity. Such a response has not been fully addressed to date and would reveal many extremely interesting pathogenic features, as well as suggesting both vaccine and therapeutic improvements.
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