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The immune response and immune evasion characteristics in SARS-CoV, MERS-CoV, and SARS-CoV-2: Vaccine design strategies

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ABSTRACT

The worldwide outbreak of SARS-CoV-2, severe acute respiratory syndrome coronavirus 2 as a novel human coronavirus, was the worrying news at the beginning of 2020. Since its emergence complicated more than 870,000 individuals and led to more than 43,000 deaths worldwide. Considering to the potential threat of a pandemic and transmission severity of it, there is an urgent need to evaluate and realize this new virus’s structure and behavior and the immunopathology of this disease to find potential therapeutic protocols and to design and develop effective vaccines. This disease is able to agitate the response of the immune system in the infected patients, so ARDS, as a common consequence of immunopathological events for infections with Middle East respiratory syndrome coronavirus (MERS-CoV), SARS-CoV, and SARS-CoV-2, could be the main reason for death. Here, we summarized the immune response and immune evasion characteristics in SARS-CoV, MERS-CoV, and SARS-CoV-2 and therapeutic and prophylactic strategies with a focus on vaccine development and its challenges.

1. Introduction

The pandemic occurrences of the SARS-CoV infection were observed in November 2002 in Guangdong, Southern China, that infected over 8422 humans and caused 916 deaths worldwide during 2002–2003 [1,2]. The MERS-CoV was firstly reported in Saudi Arabia in the year of 2012 and led to 2494 laboratory-confirmed infected cases, 858 mortalities, and 38 deaths since September 2012 [3–7]. Coronavirus disease 2019 (COVID-19) as a novel coronavirus, SARS-CoV-2 that is first identified in a seafood market in Wuhan, Hubei Province, China, have been recent outbreaks in China and many other countries around the world [2,8]. Since February 23, 2020, approximately 76,936 cases have been reported in the Chinese Mainland and also 1,875 cases outside of the Mainland [9]. Moreover, over 9,700 cases were confirmed in China, and 106 cases were identified in 19 other countries on 30 January 2020 [9,10]. According to the last WHO reports, on 4 August 2020, there have been 18,142,718 confirmed cases of COVID-19, and 691,013 deaths, globally. Also, approximately 88 574 cases have been reported in China that accompanied with 4678 deaths [11].

CoVs virulent is mainly related to some defined virulence genes that are able to antagonize the responses of the cellular innate immune system in most cases. During virus infection, an immune response against the virus is triggered by host factors [12]. Nonetheless, it is valuable to consider that immunopathogenesis as a process of disease development is related to severe (out of control) immune response, which can lead to lung damage, malfunction, and probably the reduced capacity. Thus, viral interactions with the innate immune system play a primary role in displaying of infection outcome [13]. In other words, it can be referred that the insufficient or malfunction of the immune system could potentially lead to higher viral replication and consequence tissue damages.

Some structural and non-structural proteins have been involved in human CoV pathogenesis and could be considered for vaccine production from highly attenuated strains of common human CoVs [14–19]. Effective therapeutic and protective strategies like vaccine designing that can be promptly applied against new emergent strains is a research priority, as these viruses spread rapidly out of heterogeneous and zoonotic [20]. Virus vaccines must have the following characteristics to be
utilized in human cases: immunogenicity, safety, broad-spectrum stability, and suitability for long-lasting immunity induction. To achieve these factors, comprehensive knowledge of the humoral and cellular immune responses and their defensive roles, virus molecular structure, genetic configuration, antigens, biology, and pathogenesis could be helpful in vaccine investigations. Knowledge about the details and complexity of different serotypes related to CoV is an important step for designing vaccines or a human neutralizing monoclonal antibody, which is recognized as a promising therapeutic method against the infection linked CoV or possible prophylactic agents [21–23].

Along with MERS-CoV, the SARS-CoV spike (S) glycoprotein is known as the main target for defensive immunity in vivo condition; however, SARS-CoV has different immunological effects than other beta coronaviruses with limited antiviral cross-reactivity by antibodies. Indeed, the diversity of important immunogenic factors of the S protein should be addressed for the vaccine development to supply broad protection [24]. Actually, among all viral proteins, S and nucleocapsid (N) attract the most interest among scientists for vaccine production against MERS-CoV. In contrast, other functional proteins like envelope (E) protein and a non-structural protein (NSP)16 are chosen as potential immunogens in vaccine design [25–27].

Approximately 89% of the SARS-CoV-2 nucleotide sequence is approximately similar to that of SARS-like coronaviruses. Due to this, the early advancement of the potential SARS-CoV-2 vaccine is planned to manufacture based on those advanced earlier for SARS-CoV [28].

Researchers are working seriously to find possible cures for saving human lives and producing vaccines to prevent possible future infections. In this review, we summarized the immunopathology of COVID-19, SARS-CoV, MERS-CoVs, and vaccine development strategies, progresses, and challenges for this emerging virus.

2. Coronaviruses

Coronaviruses, as known as zoonotic pathogens, belong to the Coronaviridae family of the order Nidovirales. The Coronavirus genome consists of a single-stranded positive-sense RNA (+ssRNA) (~30 kb) that is linked with a 3′-poly-A tail (crown-shape peplomers with 80–160 nM in size) and a structure of 5′-terminal cap [29]. Coronaviruses have the largest genomes (26.4e31.7 kb) among all known RNA viruses. SARS-CoV as a single strand virus covered by a lipid membrane ((M), E, and S glycoproteins) [30] appears to mediate viral entry to the host target cells, and this entrance could be facilitated via ACE2 as the functional receptor [31].

The S protein is an essential stimulator for the induction of neutralizing antibodies. Moreover, the receptor-binding domain (RBD) in the S1 subunit of S protein is composed of multiple conformational structures of neutralizing epitopes that are significant markers for vaccine development (Fig. 1).

It is proposed that recombinant proteins that contain RBD directly or vectors encoding the RBD sequence may be used as an ideal safe vaccine.

Fig. 1. The genomic structure and phylogenetic tree of coronaviruses and coronavirus Spike Glycoprotein.
for the prevention of SARS-CoV infection. Therefore, the RBD fragment in the middle part of the S1 subunit is considered as a useful vaccine against the SARS-CoV challenge [32].

MERS-CoV is responsible for MERS infection, which belongs to the genus β-coronavirus, a positive sense, single-stranded RNA β-coronavirus. Since this virus is basically originated from bats as a zoonotic disease, it has been suggested that bats are the most natural reservoir of MERS-CoV [33]. MERS-CoV genome consists of at least 10 ORFs, encoding for 4 structural proteins like SARS-CoV, including S, E, M, five accessory proteins listed as ORF3, ORF4a, ORF4b, ORF5s, and ORF8b, N proteins and 16 NSP (NSP1-NSP16) [34]. SARS-CoV and newly appeared COVID-19 are the members of the same beta coronavirus subgroup, they only have 70% similarity at the genome level, and interestingly the novel group has been found to be genetically different from SARS-CoV [35]. This isolated novel β-coronavirus, like other typical coronaviruses, has different ORFs (at least ten). ORF1a/b, as the first one, can be recognized about 75% of viral RNA, which is used for the production of two large poly-proteins. The complex of viral replicate transcriptase is formed by pp1a and pp1ab poly-proteins in MERS-CoV and SARS-CoV due to processing into 16nsp-like [36].

The ns proteins mediate the rearrangement of membranes originated from the rough endoplasmic reticulum, resulting in vesicles with double-membrane formation where viral replication and transcription occur [37,38]. While SARS-CoV-2 binds to the ACE2 receptor, cellular receptor dipeptidyl peptidase 4 (DPP4 also called CD26) is the target of MERS-S via the RBD in the N-terminal surface subunit (S1). The molecular interaction of MERS-CoV with hCD26 has been delineated. The S1 domain is responsible for the detection of hCD26, situated in a C-terminal 240-residue RBD with a core and an external subdomain [39]. It was reported that the binding of the surface S1 unit facilitates the attachment of the virus to the surface of target host cells. Additionally, S protein cleavage at the S1/S2 and the S2 site is linked to priming processes through cellular proteases, which entails virus fusion and cellular membranes, driven via the S2 subunit. Engagement of ACE2 by SARS-S as the entry receptor [40] leads to the cellular serine protease TMPRSS2 employment for priming of S protein [41,42].

The functional characterization and defined antigenicity of S protein make this protein an important target for vaccine development [43]. The 3D structure of the RBD region in the S protein regulates the formation of the van der Waals forces [24]. It is reported that Lys 31 residue can be critically recognized on the human ACE2 receptor by 394 Glu residue in the RBD domain of SARS-CoV-2 [44]. Briefly, after receptor attachment, the S protein undergoes a conformation change, which facilitates the fusion of viral envelope with the cell membrane via the endosomal pathway, resulting in the RNA release of SARS-CoV-2 into the target cells. Genome RNA is initially translated into the polyproteins of viral replication (pp1a and pp1ab), then can be cleaved into small fragments by viral proteases. Discontinuous transcription of polymeerase activity could lead to the production of different subgenomic mRNAs that can be translated later to the viral proteins. Finally, viral proteins and RNAs assembling into virions in the Golgi and ER organelles stimulates the transportation of vesicles to release [45].

3. Immune response

3.1. Innate immunity

The launch response of the immune system to the invading of a microorganism such as a virus is directly related to the host sensing of the target organism and its linked constituents like uncapped viral RNA or the cellular stress response and consequent biological changes or damages due to infection [46]. This response could be primarily conducted by germline-encoded pattern recognition receptors (PRRs) such as Toll-Like Receptors (TLRs) or Retinoic acid-inducible gene I (RIG-I)-like from the components receptors (RLRs) that enable to detect PAMPs (Pathogen Associated Molecular Patterns) originated of a virus or its replication intermediates, promoting the initial antiviral signaling cascades in response to the infection (Fig. 2) [47]. RLRs has known RNA sensors that are localized in the cytosol include three main members: laboratory of genetics and physiology 2 (LGP2), RIG-I, and melanoma differentiation-associated protein 5 (MDA5). RLRs are the essential sensors for virus infections that mediate effectively the transcriptional induction of various genes associated with the antiviral response of the host and type I interferons [48]. More interestingly, these sensors not only able to recognize viral RNAs but also the mislocalized or misprocessed cellular RNAs (unusual forms). Such unusual cellular RNAs processing or localization could potentially indicate the infection or possible occurrence of sterile inflammatory pathways; therefore, the activation of RLRs is crucial in immunopathology concept [49]. MDA5, and RIG-I, as Interferon Stimulated Genes (ISGs), are mainly transcribed during the infection of the cells by SARS-CoV under in vitro condition. It is reported that Murine coronavirus (MHV) could be detected by MDA5 in microglial cells and brain macrophages, and in oligodendrocyte cells, by MDA5 and RIG-I. Though it has not been identified yet whether SARS-CoV infection can be recognized by MHV or RLRs, SARS-CoV is more likely to have similar intermediates for the replication process, as putative ligands for RLRs sensors; therefore, SARS-CoV is able to be potentially recognized by the same protocols [47].

Toll-like receptors (TLRs) as one of the PRRs, could be expressed by several types of immune cells. [50]. TLRs can be expressed either extracellular such as TLR1, TLR2, TLR4, TLR5, and TLR6 on the surface of the cells or intracellular like TLR3, TLR7, TLR8, and TLR9 in the cytosolic compartment, where they detect the genetic material of microorganisms [51].

Viruses are mainly characterized based on the complexity of their genomes and also divided according to their replication mechanism (Baltimore classification). The nucleic acid sequence of viral genomes could have consisted of either RNA or DNA, single-stranded or double-stranded, continuous or segmented, and negative or positive in polarity [52,53].

Though there is no proof for the direct implication of TLR in SARS-CoV detection, it was reported that TLR4 could potentially detect the viral related glycoproteins of RSV presented mainly on the lung epithelium’s surface cells as an essential cofactor for the entrance of respiratory viruses. It has also been identified that TLR4 plays as a protective factor against the activity of MHV-1 in an animal investigation model for SARS- respiratory disease. Further studies also reported that by the injection of MA15-SARS-CoV to a mouse model and human dendritic cells, the transcriptional level of TLRs increased. Additionally, TLR3 activation plays a protective effect in the infected mouse model by SARS-CoV.

RNA viruses can be internalized into the cells via different mechanisms, including binding to the surface receptors, such as ACE2 for SARS-CoV or CEACAM1 for MHV and plasma fusion. This internalization process could expose the viral genomic (RNA) to the dsRNA sensing system (MDA5, TLR3, and RIGI) in the cells. These proteins could induce IFNβ and the production of IFN (IFNβ/α) receptors on the surface of the surrounding or same cells, activating the signaling pathway of Stat1 that is associated with the activation of several anti-viral genes with ISRE promoter. This could raise a question that how coronaviruses, either human (OC43, NL63, SARS-CoV, and 229E) or animal (MHV, IBV, and TGEV), interact with the IFN system (type I) [54].

Antiviral cytokine IFN not only has a vital role in controlling many viral infections but also they could promote the process of viral clearance through programing of the adaptive immune response. However, the aberrant response of IFN, cytokine, and ISGs was recognized in those patients suffering from an advanced level of SARS disease compared to the healthy group with innate immune regulated SARS disease [54,55].

Several reports indicate that IFN (Type I) could inhibit the growth of SARS-CoV in the cell culture and also the viral replication process in
different animal models such as a mouse or cynomolgus macaques. Further investigations confirm that the SARS-CoV-infected animal model with a deficiency in IFN receptors such as Type I or Type III show more advanced stages in the replication process of viruses, particularly in the lungs. The expression of IFN receptor (Type III) was identified 24 h before the transcripts of Type I IFN in 2B4 cells, showing a delay in Type I IFN signaling and proving the protective function of Type III IFN in response to SARS-CoV infection [56]. Overall, studies in animal models of SARS-CoV-1 and MERS-CoV infection demonstrated that failure to induce IFN-I response in early stages correlates with the severity of the disease. Of note, these models may be indicated that timing is critical, as IFN-I response is protective in the beginning stage of the disease but is pathologic in the late stage [57].

Although SARS-CoV could effectively infect alveolar epithelial or airway cells, the infection of different hematopoietic cells like dendritic cells (DCs) and monocyte-macrophages is failed. The infection of DCs with SARS-CoV could also induce the lower levels of antiviral cytokines (IFN-α/β) expressions and control the pro-inflammatory cytokines and chemokines up-regulations such as TNF, IL-6, CCL2, CCL3, CCL5, and CXCL10 [58]. In the same way, macropathies infected with SARS-CoV indicate a considerable delay but a significant rise in pro-inflammatory cytokines levels, particularly IFN [59]. Also, airway epithelial cells that are infected by SARS-CoV represent a large production of CCL2, CCL3, CCL5, and CXCL10 [56]. It is thought that delayed but higher amounts of chemokines and cytokines productions could lead to the deregulation or malfunction of the innate immune response to the infection. It was found that the serum levels of chemokines (IL-8, CXCL10, CCL2, and CXCL9) and pro-inflammatory cytokines (IL-12, IFN-γ, IL-6, TGFβ, and IL-1) were increased in the patients with an advanced level of SARS infection compared to the patients with an uncomplicated level of SARS infection [60].

However, infected patients with SARS in advanced levels had a meager amount of anti-inflammatory cytokines such as IL-10. Despite the increased level of chemokines and pro-inflammatory cytokines in SARS-infected patients, the levels of ISGs such as CCL-2 and CXCL10 and IFNs like IFN-γ and IFN-α were enhanced in the infected patients compared to healthy control or the infected patients at a moderate level. These data indicate the possible role of ISGs and IFNs in the immunopathogenesis of infected patients with SARS. Therefore, it seems that dysregulation and exaggeration of chemokine or cytokine in response to SARS-CoV in macrophages, AECs, and DCs could play a considerable function in the pathogenesis of SARS [60].

MERS-CoV, similar to SARS-CoV, could infect airway epithelial cells, inducing the responses of pro-inflammatory cytokines like IL-1β, IL-6, IL-8, and IFNs significantly but delayed. Though MERS-CoV is able to be replicated in either naïve or activated DCs and monocyte-macrophages, activated T cells can only support the MERS-CoV replication [53]. This is a significant contrast that SARS-CoV could abortively infect T cells, monocyte-macrophages. The infection of THP-1 cells, dendritic cells, and DCs by MERS-CoV induce the higher production levels of chemokines and pro-inflammatory cytokines but in a delayed manner. However, IFN-α/β induction through DCs and monocyte-macrophages was not a significant except for the pDCs, which produce a remarkable amount of IFNs during the infection of MERS-CoV. It was documented that the serum levels of chemokines such as CCL5, IL-8, and CXCL-10 were increased considerably in individuals infected with an advanced level of MERS in comparison with those who are suffering from moderate disease.

**Fig. 2.** Innate and adaptive immunity in SARS-COV2 infection. Immune response primarily conducted by PRRs like TLRs or RLRs that enable the detection of PAMPs originated from a virus or its replication intermediates, promoting the initial antiviral signaling cascades in response to the infection, including inflammatory cytokine production. The presentation of antigen can consequently induce humoral and cellular immunity in the human body, which are enhanced by B and T cells (virus-specific cells).
The higher amount of these factors in the serum of MERS-infected patients is correlated with increased numbers of monocyte and neutrophil in the peripheral blood cells and lungs, showing the possible function of these cells in the pathology of the lungs [60].

The kappa light-chain-enhancer of activated B cells (NF-κB) as a well-known transcriptional nuclear factor is the most critical regulator for the induction of different pro-inflammatory cytokines such as IL-6 and IL-8, the early IFN-β expression upon viral infection and innate immunity system. More interestingly, it has been proved that the induction of IL-8, TNF-α, and IL-6 could have occurred in the cells with spike protein overexpression (a protein of SARS-CoV) through the NF-κB pathway [61,62].

Neutrophils, as well-known immune cells, are presented in several types of lung diseases related to ARDS (acute respiratory distress syndrome) and probably could contribute to the acute injury of lungs. Neutrophils have been poorly investigated in aspects of their response to viral infection, particularly viral disease in the respiratory system. To be able to reach the target site (the potential pathogenic site), neutrophils breakdown the collagen in pulmonary tissue (extracellular matrix (ECM)) by the production of MMP9 proteins. The production of TNF in the surrounding cells could increase the release of MMP9, leading to the absorption of more numbers of neutrophils towards the inflammation or infection. Subsequently, neutrophils start phagocytosis and degranulation of neutrophil myeloperoxidase, elastase, neutrophil extracellular traps [63], and reactive oxygen species (ROS), which contribute to the pathogenic microorganism clearance.

Neutrophils as a first line of immune defense in the beginning of a viral infection, increase in the number of these kinds of immune cells as well as their raised lifespans, composed the important phenomenon in the patients with COVID-19. Despite their protective effect against viral infections, neutrophils could injure in host tissues infected with a virus. Activation and degranulation of these cells in the microenvironment of infected cells can damage the host tissue and worsen the disease outcome. The neutrophils, via the induction of pro-inflammatory cytokines, as well as ROS production, could worsen the disease manifestation [64]. It is suggested that severe host response in the infected patients with COVID-19 is due to improper activation of leukocytes such as neutrophils in peripheral blood. These results indicate that when the normal signals to moderate inflammation process are lost, specifically under cytokine storm conditions or a signaling interruption between neutrophils and macrophages, can cause progressive and consequently uncontrollable inflammation [65].

Netosis is known as specific cell death of neutrophils whereby NETS (decondensed chromatin fibers, granule components, and histones) are deployed in order to kill and immobilize pathogens while accelerating its death process. Though its main aim is to restrict the pathogenic spread, some viruses or bacteria could lead to excessive formation of NETs [66]. In other words, severe production of NET can induce a cascade reaction in an inflammatory procedure that is related to the promotion of metastasis in cancer disease, micro thrombosis facilitation, and damaging of surrounding tissues, resulting in permanent destroy of organs such as renal, pulmonary, or probably cardiovascular systems [65].

The lung infection by HPAI, MERS-CoV, influenza A virus, and SARS-CoV is reported to cause a severe acute lung injury such as ARDS that is characterized by considerable deterioration to the alveolar epithelium and neutrophils infiltration [67]. A study conducted by Chen and et al. showed that 38% of the patients infected by COVID-19 had higher levels of neutrophils [68]. The critical question is that neutrophilia is driven whether by virus-induced cytopathy or viral infection response.

ARDS, as a common consequence of immunopathological events for infections with MERS-CoV, SARS-CoV, and SARS-CoV-2, could be the main reason for death. In a study was represented that 6 out of 41 cases that infected by SARS-CoV-2 died because of ARDS [69]. It has been reported that one of the critical detrimental mechanisms of ARDS is the uncontrollable inflammatory response that leads to the release of a huge concentration of chemokines and pro-inflammatory cytokines by the immune system in SARS-CoV infection. Similar to individuals infected by SARS-CoV, patients with an advanced level of MERS-CoV infection reveal increased amounts of pro-inflammatory cytokines and chemokines in serum samples compared to the patient with a moderate level of infection. The cytokine storm can induce a detrimental attack on the immune system, leading to ARDS or organ failures that can cause death or severe infection condition in the patients [70].

Histological tests of lungs from individuals infected by SARS indicated a remarkable cellular infiltrate in the alveoli and interstitium, which includes macrophages and neutrophils as predominant cells. These data are associated with the increment of monocyte and neutrophil numbers and decrement of CD8+ and CD4+ T cells in the patients infected with fatal SARS.

In addition to the immunohistochemical test indicated that MERS-CoV could mainly infect alveolar and airways epithelial cells, macrophages, and endothelial cells. The severity of lung damage is related to uncontrolled levels of macrophages and neutrophils infiltration into the lungs and higher counts of these cells in the sample of peripheral blood in the MERS-infected patients [60].

3.2. Adaptive immunity

Antigenic peptides are predominately expressed by the MHC (major histocompatibility complex) in humans and can be detected by CTLs (virus-specific cytotoxic T lymphocytes). Therefore, recognition of the fundamental mechanism of the SARS-CoV-2 antigen presentation could significantly contribute to comprehend the COVID-19 pathogenesis. Because there have not been any reports regarding this issue yet, some information, according to the previous studies on MERS-CoV and SARS-CoV were described in this part. The SARS-CoV antigen presentation is mainly dependent on MHC I and partially MHC II. The presentation of antigen can consequently induce humoral and cellular immunity in the human body, which are enhanced by B and T cells (virus-specific cells) (Fig. 2) [70,71].

3.2.1. Humoral immunity

Similar to other viral infections, the antibody profile has a common production pattern of IgG and IgM antibodies in the response of the SARS-CoV infection. The IgM antibodies specified for SARS can be detected up to 12 weeks, though the IgG antibody could be identified after a long time-frame, indicating that IgG antibody play probably the protective function and IgG antibody is mainly either S-specific and N specific [71].

The infection of the SARS-CoV induced the process of seroconversion at the beginning of 4 days after the onset of the disease and was recognized in most patients for 14 days. It was reported that long-lasting neutralizing and some specific IgG antibodies were even detectable after 2 years since the infection time. The seroconversion process is observed for the infection of MERS-CoV at the end of the second or third weeks of the onset. It has been studied that the response of weak and delayed antibody is related to the severe disease condition in both kinds of coronavirus infection. Moreover, all serum samples from infected patients could neutralize the detrimental function of the SARS-CoV-2 in vitro condition, showing the possible successful increment of humoral responses. However, the humoral responses of some specific antibodies are whether associated with the severity of the disease is not clear yet [72].

Although the response of successful neutralizing is indicated in most cases, higher titers are also linked to more intense clinical symptoms, proposing that the response of a robust antibody could not be sufficient alone to prevent severe diseases. In the previous epidemic of SARS-CoV-1, it was indicated that neutralizing titers were considerably lower in the recovered patients than decreased patients. This has brought a challenge that the responses off antibodies to such viruses could potentially lead to pulmonary pathology though the induction of antibody-dependent
enhancement (ADE) as an interesting phenomenon that concerns many researchers in this field.

This phenomenon can be seen when virus-specific IgG antibodies (non-neutralizing) contribute to the entrance of virus particles to several cells such as monocytes or macrophages via expressed Fc-receptor (FcR), inducing the activation of inflammatory procedure in these cells. Research on the infected rhesus macaques with SARS-CoV-1 showed that anti-S IgG is contributed in increasing the number of macrophages and monocytes in the lung and severe acute lung injury (ALI) [57]. Jaume et al. and Yip et al. also reported that though anti-S antibodies inhibit the entrance of viruses into permissive cells, they increase the infection by attaching to the IgG Fc-receptor-II positive (FcRII+) cells, such as macrophages or B cells [73]. Furthermore, ADE is recognized in the isolation of monoclonal antibodies from a patient infected with MERS-CoV. ADE of disease is also a common concern for the development of vaccines and antibody therapies because the mechanisms that motivate antibody protection against any virus have a theoretic potential to amplify the infection or eliciting harmful immunopathology. This phenomenon occurs in MERS, SARS-CoV-1, Zika, HIV, and dengue virus infection and vaccination [74]. Vaccine-specific variations in ADE could occur for many reasons, including modifications in vaccine protein glycosylation, vaccine adjuvant, and previous exposure to other CoV strains. Multiplex approaches developed for influenza can be rapidly adapted to this use, particularly to evaluate the balance between vaccine-induced protection from infection against the increased risk of severe disease with the following infection despite vaccination [75]. However, there is not available data that supports ADE contributed by sera in animal models such as vaccinated rats with SARS-CoV-2 BD and immunized macaques with the SARS-CoV-2 vaccine [76,77].

SARS-CoV-2, as a known mucosal targeted virus, is expected to produce secretory IgA (sIgA) and also promote the mucosal immunity strongly. It is valuable to refer that the function of sIgA in the infection of COVID-19 has studied limitedly, though SARS-CoV-2 could enter into the body via respiratory mucosa, and sIgA plays a crucial role in mucosal defenses. Moreover, different reports regarding COVID-19 infection have indicated that the presence of IgA antibody in infected patients with SARS-CoV-2 and vaccinated with anti-SARS (administered either sub-lingually or intranasally). These reports show the importance of studying the role of sIgA secrets in infected patients with COVID-19 and recognition of its antiviral function in respiratory tract mucosa, inflammation, and immune response [78].

Regarding the function of systemic and mucosal IgA in the infection of COVID-19, IgA induction by using lactoferrin to promote the signaling pathway of canonical TGF-β, or retinoic acid to increase the responses of lactoferrin-induced IgA could be a potential and novel method for COVID-19 therapy. Yu et al. reported, however, the enhanced response of IgA was detected in severe COVID-19, which may show its damaging effects. It is proposed that COVID-19 is possible a part of an IgA-mediated disease (associated with IgA vasculitis and deposition), which explains the related organ injuries in COVID-19 infection such as kidney injury or acute pulmonary embolism [79].

3.2.2. Cellular immunity

The migrated cells (DCs) from the lungs to the T-cell area can activate naïve CD8+ T cells over the viral infection, resulting in the differentiation and proliferation of them into CTLs [80]. CD8+ T and CD4+ T lymphocytes (mainly CD8+ T lymphocytes) employ different cell-associated mediators to transdifferentiate as FasL, perforin, or Granzyme, as a result of the apoptosis process in target cells. Because the induction of cytolytic needs to the engagement of antigen receptors in T lymphocytes by viral complexes of peptide/MHC molecules, the apoptosis mediated by T-cells mainly restricts the infected cells. As a significant exception, the mediated cytolytic by T lymphocytes plays a negligible function in the development of injured tissue generated by the adaptive immune system upon viral infection. However, the inflammatory mediators (soluble) derived from T cells such as MIP-1α TNF and IFNγ are able to deteriorate even uninfected cells and also increase the infiltration of injury-promoting innate immune cells [71].

CD4+ T cells are able to contribute to the production of antibodies, induce the activity of cytotoxic T lymphocyte (CTL) in CD8+ T cells, and act as functional memory cells. T helper cells have essential roles in the regulation of B cell proliferation, differentiation and switching of immunoglobulin class. In particularly Th2 cells also contribute to the somatic mutation of B cells in the germline centers. [81]. ICOS and CD40 ligand (CD40L) are abundantly expressed in Th2 cells, which its ligation with ICOSL and CD40 is fundamental for the B cell’s immune responses [82].

Cytotoxic CD4+ T cells (ThCTL) are mainly determined by their cytotoxic function and/or phenotype. ThCTL can also be detected in the peripheral blood mononuclear cells (PBMCs) (human species) in severe viral infections such as HCMV (human cytomegalovirus), HIV-1 (human immunodeficiency virus 1), and hepatitis viruses. A study that was conducted by Brown et al. indicated that, in vitro generated CD4+ cells could get cytolytic characteristic and have protective effect on influenza virus infection in combination with B cell help. In another study they indicated that, functional ThCTL could be found as a large effector population in lung (besides PBMC) as well following influenza infection in mouse model and play the cytotoxic role in the infection loci. It was reported that, effector profiles of CD4+ cells is distinct in the lung than draining lymph node (DLN) and that IFN-γ production via CD4+ cells at the site of infection could play a protective role. Moreover, CD4+ cells using perforin-mediated cytotoxicity that get in the lung, but not the DLN, are able to augment recovery from lethal influenza virus infection [83-85]. More studies are in need to evaluate these cells in COVID-19.

The most current reports reveal that the numbers of CD8+ T and CD4+ cells are considerably reduced in the peripheral blood of the patients infected with SARS-CoV-2; however, their over-activation is a proof for high positive fractions of HLA-DR (as described CD4+ 3.47% and CD38+ (CD8 39.4%). Likewise, the response of SARS-CoV-infected patients at the acute phase is related to the severe reduction of CD4+ T and CD8+ T cells. In 12 patients recovering from mild COVID-19, robust T cell responses specific for viral N, M, and S proteins were detected by IFN-γ ELISPOT. In the absence of antigen, CD8+ T and CD4+ T memory cells can even persist for several years in recovered patients with the SARS-CoV infection and capable of continuing the proliferation of T cells, IFN-γ production, and DTH response.

A study reported that the infection of SARS-CoV peptide is able to be recognized in response to specific memory T cells in 14 out of 23 recovered patients with SARS infection. It has also been reported that specific CD8+ or T cells can indicate a similar influence on MERS-CoV removal in mice. These investigations could probably provide useful data about rational vaccines designing for SARS-CoV-2 infection [15]. The rise of CD8+ T cells in MERS-CoV-infected patients at the early phase could be associated with the severity of the disease, and T helper cells (Th1 type) are recognized at the convalescent stage. The cells of airway memory CD4+ T are specified for conserved epitope play a protective role against lethal function in animal models and can also observe cross-reaction between the MERS-CoV and the SARS-CoV.

The strong response of T cells associated remarkably with higher amounts of neutralizing antibody; however, more Th2 cytokines in serum such as IL-4, IL-5, and IL-10 were identified in the fatal individuals. As neutrophils have a detrimental effect on all infections, the protective or possible destructive roles of Th17 against the infection of human coronavirus remains unclear [72].

4. Immune evasion by SARS-CoV and MERS-CoV

MERS-CoV and SARS-CoV can utilize different strategies to escape from the immune system to stay alive in host cells. Over the replication cycle of SARS-CoV, the dsRNA intermediates segregation of viruses in DMVs (Double Membrane Vesicles) is able to cover viral PAMPS from PRRs recognition in the cytosol. It has not been clarified whether the
products of viral dsRNA or ssRNA degradation could be sequestered in DMVs or detected by PRRs. Due to the lack of a cap (50 nucleotides) in viral mRNAs, the eukaryotic mRNAs can be distinguished, and many viruses such as SARS-CoV have a sophisticated mechanism for mimicking the host capping system [47]. Although IFN-γ has a significant protective effect on the infection of the SARS-CoV and MERS-CoV, the IFN-1 pathway is controlled in the infected animal model. It is indicated that accessory protein 4α belongs to MERS-CoV is able to inhibit IFN induction at the MDAS activation level via interaction directly with ds RNA. Moreover, ORF5, ORF6b, ORF6, and ORF7b, and membrane proteins of this virus could block IRF3 (nuclear transport of IFN regulatory factor 3) and subsequent IFN β promoter activation. Coronavirus can also possibly affect the presentation of antigen. As an example, it was shown that the expression of genes is linked to the presentation of antigens is remarkably down-regulated after MERS-CoV infection [70].

Reportedly, SARS-CoV, through encoding nsp3-deubiquitinase [18], nsp3-macrodomain, nsp1, ORF6, ORF9b, and ORF3b, could subvert the and also inhibition of SARS-CoV replication in the respiratory tract of SARS-CoV were considered as IFN antagonists; however, the functional addition, some structural proteins, such as nucleocapsid and membrane proteins of SARS-CoV [91] . Furthermore, cellular immunity activated utilized to completely inactivate the whole virus as a novel vaccine that and tested in several animal models. U.V and formaldehyde were uti

vaccines have been produced for both SARS-CoV and MERS-CoV virions not involving laborious genetic manipulation [90] . Several inactivated whole virus or whole killed virus vaccine (WKV) is

incomplete. In spite of their high level of efficacy, they led to eosino

entry when they combined with adjuvants like Freund

immunosenescence in older individuals impacts slightly on the innate immune system, but significantly on the responses of T cell-dependent adaptive immunity. More evidence shows that the level of proinflammatory cytokines is considerably increased in elderly mice, and alveolar macrophages are sensitive to IFN-γ activation. In other words, immunosenescence and probably comorbidity disorders are able to induce viral cytokine storm in older individuals, leading to the failure of life-threatening respiratory and multisystemic involvement. Therefore individual medicine can be developed in older people according to their personal medical history [89].

5. Vaccine development and its challenges

5.1. Inactivated whole virus

Inactivated Whole Virus or Whole killed Virus vaccine (WKV) is remarkable because of its safety, cost-effectiveness, easily prepared, and not involving laborious genetic manipulation [90]. Several inactivated vaccines have been produced for both SARS-CoV and MERS-CoV virions and tested in several animal models. U.V and formaldehyde were utilized to completely inactivate the whole virus as a novel vaccine that could significantly increase antibodies concentrations against the viral proteins of SARS-CoV [91]. Furthermore, cellular immunity activated has been shown by stimulation of IFNγ and interleukin-4 production, and also inhibition of SARS-CoV replication in the respiratory tract of mice vaccine by SARS-CoV (Utah) plus ALCOH3 as an adjuvant [92]. Qu et al. have shown that formaldehyde-inactivated-SARS-CoV strain G250 prompts neutralizing antibodies up to 1:640 when administered intranasally. Specific IgA was detected in the tracheal lung fluid of immunized mice when used either alone or with cholera toxin B (CTB) or polyethylene glycol (PEG) as an adjuvant [93]. In another study, formaldehyde-inactivated-SARS-CoV strain NS-1 represented an effective response, both humoral and mucosal immunity against SARS-CoV infection in monkeys [94]. The improved humoral immune response due to the application of the inactivated virus with adjuvants, either MF59 or alum associated with the stimulation of the CD4+ T cells, has been observed even after gene-based vaccination [95]. In other two studies conducted by Roberts and See, a whole-killed (deactivatend by β-propiolactone) SARS-CoV vaccine or β-propiolactone (BPL) inactivatend, and a mixture of two adeno-virus-based vectors, one expressing the N and other expressing the S protein (named Ad S/N vaccine), were examined in BALB/c mice, golden Syrian hamsters, and 129/Sv/Ev mice, respectively. The results of the first study showed the efficiency of the WKV associated with the higher neutralizing antibody titers compared to that of the Ad S/N vaccine. Nonetheless, intranasally administration of S/N limited noticeably the SARS-CoV replication in the lungs [96,97]. Similar results were reported for the neutralizing antibodies production with various protection levels using inactivated SARS-CoV virion vaccines with or without adjuvant in different animal models [63,96,98-105]. Moreover, SARS-CoV vaccine studies in mice lead to the observation of Th2-type immune responses, and also eosinophilia. It was indicated that oligomers of the SARS-CoV S protein act as an immunogen and cause eosinophilia in animals [98,99]. These observed side effects are significant and should be accurately evaluated before using this type of vaccine in humans. Also, if genetically attenuated viruses were utilized in the starting point of virus inactivation process, the vaccine-induced eosinophilia would be probably abated.

The application results of the inactivated SARS-CoV vaccine to 36 human cases revealed its high level of safety, well tolerating, and the ability to induce neutralizing antibodies. Nonetheless, the responses of SARS-CoV-specific IFN-γ-secreting T-cells have been identified in the mice vaccinated by a mixture of WKV and Ad S/N vaccine. Therefore, it has appeared that WKV vaccines are reliably safe and able to induce either SARS-CoV neutralizing antibodies or T lymphocytes, though the mechanism of action is still unknown [106].

Chemical-based inactivated MERS-CoV vaccine where the viruses are deactivated by formaldehyde, could trigger neutralizing antibodies in mice without the activation of any T-cell responses, as reported by Deng et al. [107]. However, introducing of adjuvant (alum and Cpg ODN) to these vaccines could improve the protective immunity in the mice received human dipetidyl peptidase4, hDPP4 [107]. Wirsbich et al. designed an inactivated whole-vaccines by mixture vectors encode a fusion protein such as the S1 domain of MERS-CoV linked to the C-terminus of G protein (RABV, rabies virus).

In MERS-CoV combined RAVB, the S1 domain was combined with the RAVB particles (BNSP333-S1). The responses of neutralizing antibodies were recognized after applying of the vaccines in mice with considerable safety [108]. Despite the advantages of IVW-based vaccine responses, such as neutralizing antibodies induction and reduction of the viral load in hDPP4 transgenic mice, unwanted responses like hyper-sensitivity lung immunopathologic reactions were observed [109,110]. Besides the investigation of inactivated vaccines in animals, phase I of human trials was performed with or without adjuvants. The vaccine caused strong responses against the RBD, and the RBD-specific antibodies in the antisera could efficiently block receptor binding and virus entry when they combined with adjuvants like Freund’s complete or incomplete. In spite of their high level of efficacy, they led to eosinophilia and other related symptoms of immunopathological disorders [101]. Therefore, it is necessary to develop novel vaccines with a high level of both antibodies and T-cells response; however, limited eosinophilic reactions. Although to the complex compositions of whole-cell antigens, the evaluation and quality control of WKV is complicated, several institutions started WKV development by different strains with confident low or no pathogenicity (Table 1) [111].

Although the Th2 immune responses, which induce production of IL-4, may be sufficient for protection against viral infections, defense
Table 1
Potential preclinical vaccine candidates against SARS-CoV, MERS-CoV, and SARS-CoV2.

| Vaccine type | Antigen target | Administration route | Reference |
|--------------|----------------|---------------------|-----------|
| **Inactivated Whole Virus** | SARS-CoV (UV-V)/TLR | SC, IP, IN | [63] |
| | SARS-CoV - BPL | IM/IN | [91] |
| | SARS-CoV (Utah)/S protein + Al(OH)3 | IN | [92] |
| | SARS-CoV | IM | [94] |
| | SARS-CoV | IN or IM | [97] |
| | SARS-CoV | IN | [98] |
| | SARS-CoV + Alum | IN | [104] |
| | SARS-CoV (Advax delta inulin adjuvant) | IM | [105] |
| **Live - attenuated vaccine** | SARS-CoV | IP | [208] |
| | SARS-CoV /NF-xB inhibitors | IP | [134] |
| | SARS-CoV (MA15) | IN | [209] |
| | SARS-CoV ΔNSP16 | IN | [19] |
| | SARS-CoV (MAwt)/MA-Exon | IN | [128] |
| **DNA vaccine** | N protein | IP | [193] |
| | S, M, N, or E protein | IM | [192] |
| | S protein | IM | [191] |
| | S protein | IM | [191] |
| **Recombinant vaccine** | RBD protein | – | [178] |
| **Subunit vaccines** | RBD-Fc protein ID, IM | [101] |
| | S318-510 protein SC | [155] |
| | S ectodomain IN | IN | [157] |
| | trimeric Spike protein | IP, SC | [158] |
| | S1-Fc domain protein | SC, IM | [160] |
| | M protein | ID | [166] |
| | N protein | SC | [167] |
| | S2 protein | IP | [168] |
| | SARS-CoVΔNSP16/ExoN | IN | [211] |
| | trimeric Spike protein | IP, SC | [212] |
| **Potential vaccine candidates against MERS-CoV** | S protein (alum + CpG) | IM | [107] |
| | Chimeric RABV/S1 protein | IM | [108] |
| | MERS-CoV | IM | [109] |
| | MERS-CoV | IM | [213] |
| **Live - attenuated vaccine** | MERS-CoV mutant rMERS-CoV-RFP, or rSARS-CoV-cDNA | In vitro | [126] |
| | Truncated, soluble variant of S protein | IP | [214] |
| | MERS-CoVΔNSP16 (mutant) | IN | [215] |
| **DNA vaccine** | N protein | IP | [216] |
| | DNA encoding S protein | IM followed by EP | [194] |
| | DNA encoding S1 protein | IN | [196] |
| | Full-length (pS) or S1-subunit (pS1) | IM | [189] |
| **Recombinant vaccine** | RBD protein | IM or SC | [143] |
| | S protein | IM and IN | [180] |
| | RBD-Fc protein | IM | [177] |
5.2. Live attenuated vaccines

Though live attenuated vaccines application has been limited due to the risk of reversion to a virulent type, they are mainly known as strong immunogenic, so a single administration in the absence of any adjuvant can provoke sufficient and effective protective immunity. Furthermore, they can stimulate strong cellular and antibody responses and often induce immunity persisting for several decades, by only one dose administration [123]. Live-attenuated vaccines either viruses with reduced fidelity capacity (mutated in nsp14), partial viruses (viruses eliminated in the envelope section (E) of protein), or the attenuated vector of viruses (AAV adenovirus, measles virus, parainfluenza, VSV, poxvirus, and rabies virus) are nontoxic and can trigger antibody and T-cell responses against SARS and MERS coronaviruses [124].

Based on the investigation, a MERS-CoV live attenuated vaccine in response to was designed by eliminating of the E gene from the MERS-CoV genome as a recombinant form (rMERS-CoV) by Almazan et al. This engineered virus could not cause infection and replicate in one cycle. Both safety problems related to the risk of virulence reversion could occur by vaccines based on the live attenuated viruses, while MERS-CoV lacking E gene is propagation defective and prevents directed reverse to virulence, allowing safer alternative approaches [125].

Awareness of MERS-CoV molecular clone and recombinant viruses presenting special genes such as red fluorescent protein (RFP), provides procedures for developing antiviral agents and designing of LAV. In a study conducted by Scooby et al., the transfected cDNA (full-length) recovered some recombinant viruses (rMERS-CoV) for inserting into the component clones. Furthermore, the accessory protein ORF5 was deleted and replaced with tomato red fluorescent protein (rMERS-RFP). These recombinants (rMERS-CoV-without ORF5, MERS-CoV-RFP, and rMERS-CoV) were replicated in the high titers, while MERS lacking ORF3-5 indicated decreased titer compared to rMERS-CoV [126]. A live-attenuated measles virus (MV) vaccine that encodes the MERS-CoV spike glycoprotein (MERS-S) and nucleocapsid protein (MERS-N) was characterized in a study by Bodmer et al. The results showed that replication-competent MV-MERS-S(H) vaccine could trigger strong neutralizing antibody titers and cellular immunity (IFN-γ and TNF-α) in adult mice by a fraction of MERS CoV-specific CD8+ T cells and MV-specific CD4+ T cells. Furthermore, the recombinant vaccine of MV expressing MERS-CoV-N induces N-specific T cell responses in vaccinated animals.

In general, live-attenuated vaccines can be recognized by the innate immune system like dendritic cells expressing pattern-recognition receptors, including the TLRs. However, many viruses can induce different signaling pathways of receptors and activation of immune cells [127].

Replication fidelity as a dominant virulence factor in coronaviruses is about 20-fold greater than other RNA viruses, which is associated with a 3′→5′ exonuclease (ExoN) activity. Considering this issue, a live-attenuated vaccine against SARS-CoV, according to mouse-adapted SARS-CoV (MAwt) was investigated and found that MA-ExoN as a stable mutator phenotype could profoundly reduce fidelity after regular administration and attenuate pathogenesis in young, old and immunocompromised BALB/c mice. The obtained results indicated that high amounts of neutralizing antibodies are generated by the vaccinated mice with MA-ExoN. Also, stable weakening of CoVs and, possibly, other RNA viruses can be performed by ExoN inactivation [128,129].

Another effective attenuated vaccine based on the deletion of the full-length gene SARSCoV (SARS-CoV-ΔE) or the carboxy-terminal region of nsp1 protein was developed in the cell culture or in vivo in BALB/c mice. The attenuation of these recombinant viruses by attenuating of E and nsp1 genes mutations showed protective effects via induction of both T cells and antibody responses in the vaccinated animal model against the lethal parental virus [130].

Ribose 2′-O-methylation of viral as a molecular pattern of mRNA, which is associated with the viral nonstructural protein nsp16, disrupts the induction of type I interferon production. On the other hand, type I interferon induction dependent on the cytoplasmic RNA sensor Mda5 (melanoma differentiation-associated protein 5) in viruses with deficient in 2′-O-methyltransferase. Therefore, the study by Züst et al. showed that mutants lacking 2′-O-methyltransferase activity-induced large production of type I interferon. The attenuation mixture of wild-type and mutant NSP16 (CoV 2 = O MTase) was examined in C57BL/6 mice, which indicated good efficiency in old mice and present rationale reasons for producing live attenuated coronavirus vaccines [131].

The efficiency of another live-attenuated SARS-CoV vaccine was evaluated in the golden Syrian hamster using a recombinant SARS-CoV without the E gene (rSARS-CoV-E). The high titers of serum-neutralizing antibodies were discovered. This antibody prevents the replication process of heterologous SARS-CoV and homologous (SARS-CoV Urbani) in the respiratory tract of animal models [132]. Also, a high neutralizing antibody titer was obtained by rVSV-S vaccination in a study done by Vogel et al. [133]. A live-attenuated vaccine comprising a recombinant SARS coronavirus without the E gene was also investigated, and it was found that the rSARS-CoV-E vaccine is immunogenic and effective in hamsters as an animal model, despite being attenuated in the replication process in the respiratory tract [132,134].

The IWV and LAV are the oldest vaccines developed with new technology and could be potentially known as the first vaccine for SARS-CoV-2 in clinical trial investigations. Different research departments in China have isolated the various strains of SARS-CoV-2 and begun to manufacture these kinds of vaccines. A new deoptimized live-attenuated vaccine against SARS-CoV-2 has been designed in India (Codagenix/Serum Institute) by the sequence of the viral genome as named “rationally designed.” It is in the pre-clinical stage, and the anticipated time is in summer 2020 (Table 2) [111]. Along with, recombinant SARS-CoV-2 vaccine that incorporates the adenovirus type 5 vector (Ad5-nCoV), designed by CanSino Biological Incorporation, Beijing Institute of Biotechnology, Canadian Center for Vaccinology comes into Phase I, Phase II, and Phase I/II respectively [Id. NCT04313127, Id. NCT04341389, Id.NCT04398147].

5.3. Subunit vaccines

To develop both MERS and SARS vaccines, the most focus of the subunit vaccine was on the recombinant S protein (RBD) synthesized in heterologous expression systems. The subunit vaccines can be designed by full-length S protein and its subunits (S1, S2), N-terminal domain (NTD) with or without adjuvants to improve their immunogenicity [135]. In fact, the RBD in the S1 subunit plays a critical role in the binding process of the virus to host cellular receptors [136]. High-titer neutralizing antibodies can be induced by the subunit vaccine based on both RBD and non-RBD S protein (full-length) like the S2 subunit. As an example, a recombinant S protein trimer of MERS-CoV (MERS-S-2P) can fuse to the DPP4 receptor, and RBD, S2-specific neutralizing antibodies produced. However, neutralizing antibodies induction was observed when this protein was utilized against divergent pseudotyped MERS-CoV in mice [137–142]. Contrary, partial effectiveness in the protection of immunized macaques and reduction of pneumonia and viral load as indicated by the RBD of S protein vaccination in Lan et al. study [143]. It was also reported that various fragments of the RBD, including 358–588 [144], 367–588 [145], 377–588, and 367–606 [39] could be used to vaccine development. The fragment of RBD containing residues 377–588 of MERS-CoV is a critical neutralizing domain and induces the highest humoral responses and neutralizing antibodies in immunized animals, especially when it fused with Fc of human IgG (S377–588–Fc) [144]. In another study, S377–588–Fc of MERS-CoV with MFS9 as an adjuvant evaluated in BALB/c mouse at different doses. The results showed strong humoral and cellular immune responses in the immunized mice at the lowest doses [136]. Similar results indicated that MERS-CoV neutralization could rise by the immune response against the 358–to-588, 1-to-357, and S1-Fc 1-to-747 variants [146]. It was proved
that the fragment-comprising residues 377–588 of MERS CoV could sufficiently protect hDPP4-Tg and Adenovirus (Ad) Ad/hDPP4-transduced mice against MERS-CoV without inducing of immunological
toxicitiy or eosinophilic immune development [140,144,146–148].
Also, the residues of 367–588 or 367–606 in the MERS-CoV S1 subunit
contain RBD [145,149,39]. In addition to being immunogenic, this
intranasal administration of the RBD domain of the S protein in MERS-
CoV can lead to more powerful response of local mucosal immune sys-
tem in the lung tissue in comparison with the subcutaneous immuniza-
tion [150].

Similar to MERS-CoV, several non-neutralizing immunodominant
domains are presented in the S protein of SARS-CoV (full-length) and
are responsible for attaching the receptor to cellular ACE2, that may
assist to the immunogenicity of main neutralizing domains or trigger
detrimental immune responses [151–154]. In addition to being immu-
ogenous, these subunit vaccines indicate a high safety level and repres-
ent considerable side effects. In several studies on SARS-CoV chal-
lenges, the role of the S protein in the induction of serum-
neutralizing antibodies and the production of protective immune re-
sponses by S protein in the animal models (mice, monkey) were
confirmed [154–160]. Also, it was observed that CD4+ and the responses
of CD8+ T-cells could be triggered by SARS-CoV S protein [161,162].
Other subunit vaccines that are able to induce protective responses in
mice and rabbits have been designed based on SARS-CoV RBD. It was
discovered in another report that a chimeric protein (RBD linked to RBD-
Fc) could provoke neutralizing antibodies in the different immunized
animals (mice and rabbits), which could last for months and protect most
of the vaccinated animal models from SARS-CoV infection [163].
Alongside RBD, several studies on the neutralizing activity of S1 protein
or non-neutralizing antibody responses of S2 protein have confirmed the
defensive efficiency of S protein fragments (S1 and S2) against SARS-
CoV infection [155,160,164]. It has also been found that other subunit
vaccines produced based on non-S structural proteins such as M proteins
can lead to the neutralizing activity in the animal model [165].
Moreover, the identified protective response of immunodominant
epitopes of M proteins (M1-31 and M132-161) that were obtained from
the serum of SARS recovered patients in the immunized mouse and
rabbit have been shown to cause immunogeneity and can trigger IgG
specific antibodies in rabbits [166]. Various investigations showed that
the CoV N protein could be considered as another antigen candidate in
SARS-CoV vaccine production [167,168]. Unlike M protein, the inter-
action of antibodies with CoV N proteins could provoke the activity of no
virus-neutralizing under in vitro condition [154]. However, in vivo
conditions, the induction of cell-mediated immunity by the protein can
cause the protection to be activated [106]. The defensive mechanism of
N- and M-based SARS subunit vaccines is still unidentified.

SARS-CoV-2 S protein subunits such as S1 subunit that consist of two
domains including the C-terminal domain (CTD) linked RBD domain
and NTD, and the S2 subunit consists of the membrane fusion peptide (FP), a
membrane-proximal external region (MPER), and a transmembrane
domain (TM) [169]. Several institutes have recently developed many
subunit vaccines against SARS-CoV-2. All of these vaccines are in the
pre-clinical stage. Drosophila S2 insect cell expression system VLPs
(Virus-like particles) is a developed strategy that has been currently used
by Expres2ion. Virus-like particles or multiple recombinant structural
proteins have enclosed a cell membrane lipid envelope or nano-
structures enclosing the capsid proteins within itself. Also, it can fold the
antigenic proteins of several microorganisms, such as bacteria, fungi,
sects, mammalian cell lines, and even in transgenic plants. VLPs can
act as non-replicating vectors for subunit and live-attenuated vaccine
production in the absence of its genome [170].

Other subunit candidate vaccines based on S protein have been
developed by WRAIR/USAMRIID, University of Epivax, Georgia, and
Queensland/GS, which are in pre-clinical stages. Adjuvanted S protein
trimer introduced by Clover Biopharma/Genentech and Sanofi Pasteur/Glaxo Smith Kline institutes. Also, II-Key peptide and plant-based subunit vaccines are produced by Bio/CC-Pharming and Genex/EpiVax, respectively. In addition, full-length S-trimer/nanoparticle plus Matrix M by Novavax and S1 or RBD protein by Bio/CC-Pharming were listed in the WHO list March 2020 (Table 1) [171,172].

5.4. Recombinant vaccines

For an extended period, recombinant vaccines were widely produced in yeast or mammalian cells, which both are expensive expression platforms. The main disadvantages of such systems include high costs for preparation culture media and the risk of contamination by human pathogens. Additionally, the production of recombinant vaccines in bacterial systems was not successful due to improper folding of eukaryotic peptides and the occurrence of inclusion bodies in bacterial hosts [173]. Most of the recombinant vaccines are designed based on highly purified recombinant proteins or subunits of pathogens (Table 1).

5.4.1. Recombinant vector vaccines (RVVs)

Recombinant vector vaccines (RVVs) are produced based on bacterial or live viral vectors that are designed to represent various exogenous antigens [174]. RVVs appear to be the most promising types of live vaccines for the development of a transmissible vaccine platform [175]. Overall, gene-based vaccines are able to stimulate the responses of both cellular and humoral immune systems strongly, and viral vectors could potentially be an effective approach for antigen-encoding genes delivery and antigen presentation. To be applied as an effective vaccine carrier, the provided viral vector should be firstly safe and then present pathogen-specific antigens efficiently to the immune system [176]. RVVs appears to remain effective, even when genetically are not in a stable form and prone to reversion to insert free vector. These types of vaccines seem to be the most promising transmissible vaccine platform [175].

It has been shown that although recombinant RBD vaccines were formulated with Freund’s adjuvant and Sigma adjuvant system® (monophosphoryl-lipid A and trehalose dicorymycololate adjuvant) to elicit neutralizing antibodies with strong protective immunity in the vaccinated animals, they significantly show reduced or eliminated antibody-dependent immune enhancement (ADE) and other related detrimental inflammatory or immune responses [177]. It has been also demonstrated that a recombinant SARS-CoV RBD protein fused with human Fc produces highly strong immune responses, leading to the completely protected vaccinated mice against SARS-CoV infection [163]. Considering the robust capacity of RBD for the induction of neutralizing antibody, the recombinant proteins linked RBD can be utilized for the development of SARS-CoV vaccines [178]. Another study showed that long-term protective immunity in animal models could induce highly robust neutralizing antibodies after the administration of SARS-CoV RBD proteins [179].

What makes S protein of MERS-CoV as a key target for vaccine development is the immunogenic property that can induce neutralizing antibodies. For this reason, a chimeric virus was constructed based on the vesicular stomatitis virus (VSV) through the replacement of the G gene by the MERS-CoV S gene (VSVΔG-MERS) to develop a vaccine against MERS-CoV disease [180]. Previous studies on MERS have revealed that subcutaneous (s.c.) administration of the recombinant protein (RBD-Fc) can promote the potent ability of the vaccine to induce systemic neutralizing antibody responses in the vaccinated mice. Furthermore, intranasal (i.n.) use of MERS-CoV RBD-Fc can be able to induce the responses of the humoral immune system compared to the one provoked by s.c. Uses, including the release of neutralizing antibodies, but more strong systemic responses of cellular and local mucosal immune responses in mouse lungs [150]. Other vaccines have been developed against the MERS-CoV infection highlights the function of RBD on the spike protein of MERS-CoV and the protective role of the recombinant NTD (rNTD) of spike proteins as an effective vaccine formulation in female BALB/c mice [38]. Studies on the development of rRBD vaccination against MERS-CoV infection have shown that while the RBD mainly induces most of the responses of the host immune system, this type of vaccine-induced only partial protection against the infection in both mice and non-human primates [143,181].

It has been demonstrated systemic responses of lung resident memory T-cells, secretory IgA, and IgG could be induced by the recombinant adeno-virus-based vaccine, which expresses MERS-CoV S protein when utilized intrasinally into BALB/c mice, providing a stable neutralizing immunity to MERS spike pseudotyped virus; therefore, it is suggested that this vaccine has probably protective role against MERS-CoV infection [182]. The responses of the immune system to these provided vaccines were assessed in BALB/c mice, and obtained results indicated that RV based vaccines are able to induce earlier antibody responses and higher amounts of cellular immunity compared to the GEM particle vector [40]. Based on an animal study, the induction of neutralizing antibodies and the consequence of protection from the virulent of SARS particles in mice emerged by the recombinant vaccine [183]. As discussed, another vaccine that is developed for clinical trial phase I is a DNA-based vaccine against SARS-COV [184]. Besides the vaccines undergoing phase I clinical trials, one of the notable COVID-19 vaccine candidates is a recombinant formulated vaccine in oral tablet form, which is used the genome sequence of SARS-CoV-2 and can induce strong systemic and mucosal immunity (Table 1) [185].

A clinical trial study conducted by Zhu et al. on recombinant adeno-virus type-5 (Ad5) vectored vaccine against COVID-19 expressing the S glycoprotein of SARS-CoV-2 virus is e is tolerable and also immunogenic in healthy adults 28 days after vaccination. The peaked of specific humoral responses against SARS-CoV-2 was shown at day 28 post-vaccination in healthy adult persons, and rapid, specific T-cell responses were peaked from day 14 post-vaccination suggesting the potential investigation of Ad5 vectored COVID-19 vaccine that makes it a strong warrant for further investigation. Additional suggestions on the safety and also immunogenicity of the Ad5 vectored COVID-19 vaccine will be provided via an ongoing phase 2 trial in China (NCT04341389).

Along with, recombinant SARS-CoV-2 vaccine that incorporates the adeno-virus type 5 vector (Ad5-nCoV), designed by CanSino Biological Incorporation, Beijing Institute of Biotechnology, Canadian Center for Vaccinology comes into Phase I, Phase II, and Phase I/II respectively [Id. NCT04313127, Id.NCT04341389, Id.NCT04398147] [186,187].

5.5. DNA vaccines

DNA vaccines represent two significant benefits compared to protein-based vaccines: (i) the easy DNA manipulation and (ii) low cost of production [188]. In addition, in contrast to the other protein-based subunit vaccines, DNA vaccines can lead to the Th1-biased immune response [189].

The success in the outcome of vaccination is linked to both antibody and T-cell-mediated immunity, and generally, among different vaccines, only live-recombinant vaccines can effectively induce cellular immunity. It has been demonstrated that DNA vaccines encode proteins from pathogens induce both humoral and cellular immune responses. What makes DNA vaccine to induce a cellular immune response is the mimicking of the live viruses effects, which produce antigenic proteins and efficiently represented by MHC class I; therefore, inducing the responses of CD8+ T-cells [190].

Several successful DNA vaccines have been developed for the functional proteins of SARS-CoV, such as the one designed for S, and also those identified for M and N proteins [191]. A DNA vaccine that is provided based on M protein expression has been proved that is able to provoke neutralizing antibody and the activity of cytotoxic T-lymphocytes in mice [192]. The N-protein peptide N220 of the SARS-CoV synthesized via bioinformatics assessment supplies essential information for designing of therapeutic vaccines against SARS-COV infection.
are associated with minimized risk of the immunopathologies, DNA vaccines expressing S1 subunit can be considered as a potential vaccine candidate. It needs to note that full-length DNA vaccines against MERS-CoV lead to considerable titers of IgG2a and IgG2b antibodies (Th1-skewed response) along with very subtle S1-specific CD8+ T cell responses. This result suggests that raised amounts of all IgG isotypes in a balanced Th1/Th2 response is generated by pS1-immunization with markedly increased CD8+ IFN-γ response compared to the pS group [189]. Another study supports the use of the plasmid encoding S1 protein for the development of DNA vaccines against MERS-CoV S1 protein. The obtained results confirmed the strong protective humoral and cellular immune responses of DNA based vaccines in mice and also antigen-specific CD4+ and CD8+ T cells secreting IFN-γ and other generated cytokines provoked by gene-based vaccines [196].

The previous reports on MERS have revealed that although subcutaneous vaccination with RBD of MERS-CoV S protein-induced systemic humoral immune responses, much stronger local mucosal immune responses, those induced by intranasal vaccination are strong and more robust [150]. Recently animal studies in rhesus macaques, mice, and camels demonstrated immunization with a MERS-CoV RBD vaccine, results in the activation of T-cell responses and increased production of Th1 and Th2 cytokines [194].

The combination vaccines (protein and DNA), encodings protein, which is in the preclinical phase, and DNA-based vaccines with moderate and mild symptoms in Phase I and II clinical trials are two other vaccines against MERS-CoV [139,189].

In a recent animal study on DNA vaccine, which encodes more than 700 residues of the S1 protein, the defined vaccine induces strong protective responses of the antigen-specific humoral and cellular immune responses in mice [196]. Based on the fact that SARS-CoV-2 shares approximately high nucleotide similarly (89%) to SARS-like coronaviruses, the strategies for designing of SARS-CoV-2 vaccines are built based on the previous advances for SARS. Furthermore, genomic analyses indicated that either SARS-CoV or SARS-CoV-2 exhibits genomic similarities in the receptor-binding domain that can directly bind to the human receptor ACE2, suggesting its essential implications for the designing of novel vaccines [197].

A DNA vaccine candidate targeting MERS have advanced into clinical trials [Id.NCT02670187]. Another vaccine candidate against MERS has been recently presented to phase I/II clinical trials as a DNA vaccine [198]. Furthermore, an efficient vaccine candidate named MERS001 for MERS-CoV with ChAdOx1 that encodes the S protein of the virus is developed at phase I clinical trial [Id. NCT03399578].

To date, some companies experience the designing of DNA-based vaccines that encode S protein against the COVID-19 challenge since March 13, 2020, according to the WHO report, including Inovio Pharmaceuticals (USA), Sanofi Pasteur/BARDa (France/USA), Sciences/Evvivax (USA/Italy), Takis Biotech/Applied DNA [199].
response. Therefore, to achieve more information about the COVID-19 and immune response interaction, further investigation is required.

There is limited information about SARS-CoV-2 infection and more questions than responses for the lately recognized virus, such as the structural features, etiology, or related epidemiology, pathogenesis, and immunopathogenesis, etc. Access to necessary information on highly pathogenic SARS-CoV2, like identifying of mediators that manage the immune response, the involved pathways, and the mechanism regulate virus behavior, will help the design and development of successful antiviral therapeutics and vaccine candidates. To stop disease progression and the immune response promotion, it is necessary to formulate an effective vaccine.

High mutation rates in SAR-CoV-2 are the critical subject in vaccine design, specifically when it can affect the binding affinity of the virus. A SARS-CoVirus -2 variant carrying the spike protein amino acid mutation D614G has grown into the most dominant form in the global pandemic. Active tracking of variant frequencies shown a recurrent mutation D614G has grown into the most dominant form in the global pandemic. To stop disease progression and the immune response promotion, it is necessary to formulate an effective vaccine.

Despite fast developing in vaccines, it will probably come too late to have an impact on the first wave of a potential pandemic. Critical lessons can be attained for the development of potential vaccines against rapidly emerging viruses. Notably, if the virus to be maintained within their host communities, SARS-CoV-2 vaccines will be critical to reducing morbidity and mortality. Moreover, as SARS-CoV-2 targets the mucosa of the respiratory tract, developing a vaccine that boosts the robust immunity through the in route as an efficient strategy to stop COVID-19 infection. The immunogenicity of vaccines is determined via numerous factors; the route of administration is one of the important factors. Therefore, the administration of vaccines in an optimal route and the appropriate adjuvant to ensure an optimal immune response could play a significant role in the development and successful application of SARS-CoV-2 vaccines.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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