Perspective on the Role of Antibodies and Potential Therapeutic Drugs to Combat COVID-19

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Abstract
The sudden emergence of the novel severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) causing the coronavirus disease of 2019 (COVID-19) has brought the world to a standstill. Thousands of people across the globe are biting the dust with every passing day and yet more are being tested positive for the SARS-CoV-2 infection. In order to dispense this current crisis, numerous treatment options have been tried and tested and many more are still under scrutiny. The development of vaccines may help in the prevention of the global pandemic, however, there is still a need for the development of alternate approaches to combat the disease. In this review we highlight the new discoveries and furtherance in the antibody based therapeutic options and the potent drugs, with special emphasis on the development of the monoclonal and polyclonal antibodies and the repurposed drugs, which may prove to be of significant importance for the treatment of COVID-19, in the days to come. It is an attempt to evaluate the currently presented challenges so as to provide a scope for the ongoing research and assistance in the development of the effective therapeutic options against SARS-CoV-2.

Keywords Antibody · SARS-CoV-2 · Convalescent plasma · COVID-19 · Cytokine · Therapeutic drug

Abbreviations
IL Interleukin
GM-CSF Granulocyte–macrophage colony-stimulating factor
IFN Interferon
TNF Tumor necrosis factor
ARDS Acute respiratory distress syndrome
APC Antigen presenting cells
MHC Major Histocompatibility Complex
TCR T-cell receptor
CTLs Cytotoxic T lymphocyte

1 Introduction
The SARS-CoV-2 (severe acute respiratory syndrome coronavirus 2) or the 2019-nCoV, responsible for the COVID-19, is a member of the ‘Coronaviridae’ family of viruses which are known to mainly cause respiratory tract illnesses ranging from common cold to SARS [1]. The first occurrence of the disease was observed during early December in Wuhan, China and it was reported to the WHO (World Health Organisation) on 31st December 2019. Thereafter, it was soon declared as a global pandemic on March 11, 2020, by the WHO. As on 18th August, 2020, there are more than 21 million confirmed active cases of COVID-19 and more than 7 lakh deaths globally [2]. The 2019-nCoV are enveloped viruses containing single-stranded positive-sense RNA. This RNA is used as a template that encodes for both non-structural (NSP) and structural proteins. There are 16 NSPs encoded by the viral genome, which play a specific role in the viral replication and transcription, 4 major structural proteins and 5–8 accessory proteins [3]. The 4 structural proteins of the virus include the membrane (M), spike (S), envelope (E) and nucleocapsid (N) proteins, that are essentially required for the viral attachment, infection and assembly. The Spike protein [S], which is a structural protein, is one of the major drug targets and is known to be composed

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of two subunits, S1 and S2. The S1 unit contains the RBD (receptor binding domain) which mediates the attachment of the virus to the ACE2 (angiotensin converting enzyme 2) receptor of the host (humans), whereas the S2 unit is responsible for the fusion of the host and viral cell membranes [4].

It was also confirmed that the genomic sequence of the SARS-CoV-2 was ~79.6% and 96% identical to the SARS-CoV and bat coronavirus, respectively [5]. In spite of being similar to the SARS-CoV (Severe acute respiratory syndrome coronavirus) and MERS-CoV (Middle East respiratory syndrome coronavirus) viruses, which had led to a similar outbreak in the years 2002 and 2012, respectively, the SARS-CoV-2 is considered to be more dangerous, since its rate of human to human transmission is much more as compared to the aforesaid viruses. It was also observed that the binding affinity of the SARS-CoV-2 to the ACE2 receptor is much higher as compared to the SARS-CoV virus, making it even riskier [6].

The very fact that the discovery and large-scale production of a SARS-CoV-2 targeted vaccine would take a very long time to get into market, makes it important to analyse and research about the existing medicinal drugs which can be utilised for the treatment. We may also develop new therapeutic options that may take a shorter time to be produced and are effective in treating the COVID-19 patients [7]. This review hence consists of a comprehensive study of some of the re-purposed drugs and antibody based therapeutic options, that have been developed or are in pipeline, for controlling the COVID-19 pandemic outcomes in the future. It provides insights into the recent discoveries and advancements related to them.

2 Antibodies for the Expeditious Rescue

Today the major research focuses on the development of antibody (Ab) molecules against the SARS-CoV-2. The administration of passive Abs either help to neutralize the antigen (Ag), which is expected to be present in smaller quantities during the initial phase of infection or mediate the modification of the inflammatory and cellular immune response, which is also easy to manage before the actual symptoms start to appear. The efficiency and effectiveness of the therapy is largely dependent on the composition and amount of Abs conferred [8].

As the technology has advanced, it has become possible over the last few decades, to purify and manufacture immunoglobulins (IGs) and hyperimmune globulins (H-IGs). The H-IGs preparations are often reported to be more effective as compared to the IVIgs (intravenous immunoglobulins), since these preparations contain known titers of Ab against a specific Ag, derived from the plasma of sero positive donors with sufficient titers of NAbs (Neutralizing antibodies). The H-IGs also present an advantage over the CP (convalescent plasma), because it prevents the transfer of any harmful coagulation factors present in the serum and its composition and preparation is more consistent as compared to CP [9]. Recently, a Japanese drug making company called Takeda Pharmaceutical Co., was reported to experimentally develop anti-SARS-CoV-2 polyclonal hyperimmune globulins (H-IGs)) that was derived from the blood plasma of the recovered COVID-19 individuals [10].

The onset of the coronavirus infection begins when the RBD in the spike protein of the virus binds to the host cell surface receptor, ACE2, in case of both SARS-CoV and SARS-CoV2. Hence an effective anti-SARS-CoV2 mAb (monoclonal antibody) or an anti-ACE2 mAb might prove to be an effective method of fighting against CoV-19. NAbs like 80R, CR3014, F26G18, F26G19, etc. binding to the spike receptor binding protein in SARS-CoV were discovered earlier [11]. It was observed that the NAbs that targeted the ACE2 binding site of SARS-CoV, did not bind to the SARS-CoV2, implying that there was a difference between the RBDs of the two strains, that have an important role to play. Thus, the development of novel NAbs specific for SARS-CoV-2 RBD is the need of the hour [12]. An interesting hypothesis which gathered attention was that the rabbit sera, derived from rabbits that are immune to SARS and contained Abs raised against its S1 subunit, was observed to reduce the entry of both SARS-CoV and SARS-CoV-2 efficiently. Thus, Abs raised in response to SARS-CoV were observed to offer some level of protection against SARS-CoV-2, however the level of cross reactivity and efficiency of these Abs to neutralize the novel virus has not yet been determined. Two available IVIgs preparations: Gamunex®-C (Grifols Therapeutics, Inc., NC, USA) and Flebogamma® DIF (Instituto Grifols S.A., Barcelona, Spain), initially developed using plasma from donor patients of USA, were also observed to contain Abs that cross reacted positively with the SARS-CoV-2, at a concentration of 100 μg/ml and 1 mg/ml, respectively. These can thus be used as a potent therapeutic option; however further validation of these IVIgs is still under process [13, 14]. In a recent study a group of scientists characterized 206 RBD specific mAbs from single cell B cells of eight COVID-19 infected patients by using the FACS (fluorescence activated cell sorting) technique. The effectiveness of binding and neutralization of these Abs were tested using surface plasmon resonance assay and neutralization assay on both the pseudotyped and original SARS-CoV-2 virus. The crystal structure analysis of the RBD bound Ab revealed 2 potential human monoclonal NAbs named P2C-1F11 and P2B-2F6 [15].

Another study based on ELISA-cross reactivity of Ab containing supernatants, of SARS-S hybridomas (spike protein antibodies obtained using hybridoma technology)
obtained from immunized mice encoding chimeric immunoglobulins (Human- Heavy and Light chains; H2L2 and constant region of rat origin), was also conducted recently. The hybridomas were made by the fusion of the spleen cells of the immunized mouse and the SP2/0 myeloma cells. 4 of these Abs showed cross reactivity with the S1 subunit and 1 out of these even showed cross neutralizing activity towards both SARS-CoV and SARS-CoV-2. This chimeric Ab was named as 47D11 H2L2, which was fully humanized by cloning the V_{H} and V_{L} human chain regions into a human IgG1 isotype backbone. This finally led to the creation of the first recombinantly expressed human 47D11 mAb, against the SARS-CoV-2. The Ab was also successful in preventing the VeroE6 cells from the infection of SARS-CoV and SARS-CoV-2 pseudo typed viruses [16]. Attempts have also been made to develop a recombinant Ab by the fusion of the extracellular domain of the ACE2 to the constant region of the human IgG1 immunoglobulin. The recombinant Ab showed high affinity towards the RBD and was suggested and tested to neutralize both SARS-CoV and SARS-CoV-2 pseudotyped viruses, in mice. However, for high scale development for therapeutic purposes, this rAb was limited by the short half-life and was also not tested in-vivo on the live viruses [17].

In the U.S., many teams are independently running various studies to discover mAbs, for the emerging threat of COVID-19. The research groups are funded by the Pentagon’s defense advanced research projects agency (DARPA) and even by the Bill and Melinda Gates foundation. AbCellera Biologics, Canada, is currently working on an engineered version of spike protein as a bait to collect Abs against SARS-CoV-2. They collected 500 Abs against the spike protein from a single patient of Seattle, out of which 24 were the most potent ones, since they retained their shape on mass production and were bound to the S protein for the longest time. Another US based company, Regeneron, is also trying to develop a pair of Ab formulation or an Ab cocktail preparation (human and mouse derived Abs) which targets or binds to different sites of the spike protein itself, ensuring a better neutralizing ability, taking into account the mutations that may occur in the SARS-CoV-2. Vir Biotechnology, has also come up with an Ab that was obtained from a SARS-CoV recovered patients of 2003, which cross neutralizes the SARS-CoV-2 by binding to the highly conserved region of its RBD. They are now trying to make further modifications to the Ab, to increase its effective half-life. The modifications being made are of two kinds: one which extends the viable life of the antibody by preventing it from degradation while the other improves the supposed vaccinal impact, which signals T cells to help decimate the infected cells.

In his recent research, Brouwer et al. also isolated mAbs from the convalescent plasma of 3 COVID-19 patients. In order to capture the S protein specific Abs, present in the sera of the patients, stabilization techniques were used to generate a soluble and stabilized S protein in its prefusion conformation. These S proteins were further fluorescently labelled and used to analyse the S-protein specific B cells using flow cytometry technique. These cells were then sorted out from the entire pool of B cells, for the isolation of S protein specific mAbs. The Abs were then expressed in the HEK293T cells and screened for the binding to the S protein using ELISA technique. On the other hand, Surface Plasmon Resonance based competition assays and electron microscopy methods were used to demonstrate that the S protein of SARS-CoV-2 contained various antigenic sites (RBD or non-RBD epitopes). Further, 84 mAbs that showed affinity towards the S protein were selected and these were further narrowed down to 19 Abs that efficiently inhibited the pseudo SARS-CoV-2 virus by binding to various different antigenic sites on the spike protein. 14 out of these 19 mAbs were found to bind to the receptor binding domain and 2 were expected to be highly potent NAbs which showed a 47 and 60 pM (picomolar) neutralizing activity [18].

Many other groups like the collaborative team of AstraZeneca and Vanderbilt University, are creating a huge library of random Abs, using phages that are targeting regions other than the RBD of the spike protein of the novel virus. Ever since the search for the Ab against the SARS-CoV-2 began, many attempts have been made to develop these Abs using the phage display technology. In this technique a gene that encodes for the protein (antibody) of interest, is integrated into a phage (virus that infects bacteria) genome, such that it allows the expression of this gene on the surface or capsid of the phage, thus, displaying it on the exterior. This phage can then keep producing similar new phages that are displaying desired peptides, once they infect a bacteria (commonly *E.Coli*) and uses its replication machinery for this purpose. A few of these developments include: a single-domain Ab from llama which neutralizes the S antigen of SARS-CoV and pseudotyped SARS-CoV-2 as a bivalent human IgG Fc-fusion (fragment crystallizable) protein and a synthetic human Fab (fragment antigen binding) library screened against the RBD domain of the SARS-CoV-2 [19, 20]. Computer techniques like molecular modelling softwares are also being applied, by scientist like, Jacob Glanville, Distributed Bio, to develop a huge library of variants of Abs of SARS, which could be further sorted to find Abs that binds to the SARS-CoV-2 and neutralize a broad range of other coronaviruses in future [21].

GigaGen Inc., has also initiated development of recombinant anti-coronavirus (COVID-19) hyperimmune gamma globulin (rCIG) polyclonal antibody (pAbs) after agreement from the FDA (Food and Drug Administration). The company uses single cell technology to recreate the library of COVID-19 Abs. rCIG helps in reproduction of the complete
Ab reserves (B cells reserve) from recovered patients and this will be used to intravenously inject the COVID-19 patients. It can also be used as a prophylaxis in case of high-risk individuals. The recombinant pAbs are expected to have a higher potency as compared to plasma derived antibodies [22]. Another pAb called COVID 19 Spike Protein Coronavirus Polyclonal Antibody or SARS-CoV-2 Spike Antibody, is also under investigation. It is an IgG isotype derived from rabbit (the host system) [23]. Attempts are also being made to design pAbs which target different regions including the S1 RBD, Nucleocapsid protein, S2 subunit, etc. [24].

2.1 Therapeutic Aspects of Convalescent Plasma in COVID-19

The convalescent plasma (CP) therapy is a very old approach which involves the transfusion of blood serum of a person, who has just recovered from a viral infection and hence contains the Abs raised against the infecting antigen, into another infected person. This helps the infected person in fighting the viral antigen, due to the presence of Abs that either bind to it or activate the immune response (cellular cytotoxicity or phagocytosis) against it [25]. Several cases and studies based on treating various viral infections by the early administration of the CP have been reported earlier [Supplementary Material Table 1].

These findings helped hypothesize that the use CP therapy may also prove to be beneficial as an immediate solution for the patients of CoV-19. A study was conducted in the Third People’s Hospital in Shenzhen, China, between 20th January and 25th March, 2020, where 5 patients that were critically ill with the CoV-19 infection, were treated with CP containing NAbs [26, 27]. Similar studies were also carried out in different parts of China and South Korea, for different durations between January-March, 2020, on critically ill CoV-19 patients [28–30]. All 27 patients, however, showed no adverse effects, indicating that the CP therapy was well tolerated among patients. The clinical status of all the patients improved and an increase in the Ab titre and decrease in the viral load within 12–45 days of transfusion were observed. These patients were however, supported by antiviral drugs or hydroxychloroquine, which made it difficult to conclude the exact consequences of the therapy alone and the results were not completely confirmative, since these were tested on a very small number of patients, that included no control. All the studies, though in favour of the therapy, suggested that an optimal dose, timing and procedure, is still required for further investigation via controlled clinical trials on a larger scale [31, 32].

CP therapy was proposed by US FDA for treating severely ill CoV-19 patients on 24th March 2020, whereas the requirements to be fulfilled by the donor were established by the FDA on 11th April, 2020 [33]. The ICMR (Indian Council of Medical Research) has also approved to conduct the clinical trials for the CP therapy at the SCTIMST (Sree Chitra Tirunal Institute for Medical Sciences and Technology), India [34]. CP therapy is also currently in clinical trials under the COVID-19 Convalescent Plasma Project, a program in the U.S. In France, Germany, Italy, Netherlands and Belgium as well, the blood transfusion centers are aiming at collection of plasma from donor patients [35].

Even though CP therapy has shown positive outcomes in numerous cases, it also has some risks associated with it. These risks broadly fall into two categories: serum sickness and Ab dependent enhancement of infection (ADE). The serum disease is associated with the risk related to the transfer of components like other infectious agents from the blood serum of the donor, to the patient being treated whereas the ADE occurs when the Abs contrarily mediate the viral entry and eventually infection in host cells [36]. There is no evidence of occurrence of ADE in COVID-19 patients, until date. However, the ADE was observed to occur in the infection of SARS-CoV. Also, studies in macaque models (primates) suggested an enhanced pro-inflammatory response in alveolar macrophage, which lead to acute lung injuries, when treated with anti-SARS-CoV Spike IgG [37]. CP also has other issues associated with its administration, which include the variable NAb titres, infectious agents getting transferred during transfusion, difficulty in collecting the plasma from suitable donors, etc.[38]. To avoid such risks to occur while treating the SARS-CoV-2 infection, various conditions have to be met before actually implementing the administration of the convalescent serum to the CoV-19 patients.

2.2 Controlling the Cytokine Storm Syndrome

The cytokine storm has been observed to occur during infectious diseases including the ones caused by influenza viruses and coronaviruses. It is an exuberant immune response which leads to an elevated level of cytokines/chemokines, causing acute lung injuries, ARDS (acute respiratory distress syndrome), multiple organ failure and even death, in many infected patients. A report from China showed that there is an elevated concentration of all the anti and pro-inflammatory response factors, that is, ILs (Interleukins: IL-2ra, IL-6, IL-10, IL-18), IF γ (Interferon), monocyte chemotactic protein 3, granulocyte colony stimulating factor, interferon gamma induced protein 10, macrophage inflammatory protein 1 alpha, tumor necrosis factors (TNF), etc., in the critically ill COVID-19 patients [39, 40]. These studies helped understand that the cytokines may be utilized as biomarkers to predict the progression and severity of the infection. This leads to the implication that cytokine storm should also be considered of significant importance while looking for therapeutic opportunities, for COVID-19 [41].
A good therapeutic option is Tocilizumab, which is a recombinant and humanized mAb that specifically binds and blocks the IL-6 receptors [42, 43]. It has the ability to bind to both the mIL6R (transmembrane IL-6 receptor) and sIL-6R (soluble IL-6 receptors) receptors of IL-6. It is sold under the name Actemra, which was earlier approved by FDA for use in treatment of diseases like rheumatoid arthritis, etc. [44]. The clinical examinations with this therapeutic mAb have shown to reduce fever and C-proteins (complement proteins) within the patients of COVID-19 worldwide [45–48]. Recently, Roche’s Genentech unit received US FDA approval after clearing the phase 3 trial of Actemra, in severely ill COVID-19 patients [49]. Several other Ab preparations like the completely humanized IgG1 mAb, called Emapalumab, which earlier got an FDA approval for the treatment of haemophagocytic disorder and Infliximab and Etanercept, known for their use in treating autoimmune diseases, could also be suggestive therapeutic options. Emapalumab is known to bind to both free and receptor bound IF γ whereas Infliximab and Etanercept target the TNF α, reducing the hyper immune response. Although more studies, research and clinical trials need to be done on the safety and efficacy of these drugs [50]. Recently, IL-1 targeting mAb called Canakinumab, a humanized IgG1 mAb called Gimsilumab targeting the granulocyte–macrophage colony-stimulating factor (GM-CSF) and Rituximab-CD20, an anti-CD-20 mAb, have also entered into clinical trials for the treatment of patients with COVID-19 [50–52]. Lenzilumab, a GM-CSF targeting drug has also received an approval by the FDA for the phase three trials, considering the significant role GM-CSF may play in the hyperinflammatory response in the COVID-19 patients [53].

The immunomodulatory agents may also be used in combination with the antiviral drugs, to reduce the cytokine storm effects in the COVID-19 patients. These agents include: corticosteroids, interferon gamma, IVIg, IL-1 and IL-6 antagonists (like Anakinra, etc.), TNF blockers, chloroquine, hydroxychloroquine, ulinastatin, etc. [54–56]. All these agents may directly or indirectly help in suppressing the effects of the cytokine storm. Some other drugs that are currently under Phase 2 clinical trials include Sarilumab, an IL-6 receptor inhibitor; Baricitinib and Ruxolitinib, JAK-STAT signalling inhibitors; Fingolimod, which inactivates IL-6 and also STAT3 pathway and statins which show anti-inflammatory effects by blocking NF-κB and NLRP3, are being explored [57, 58]. Minocycline, another drug with potential re-purposing capabilities, has also been recently proposed for the management of the cytokine storm effects in the COVID-19 patients due to its potent immunomodulatory effects [59].

A recent study suggested the use of ursodeoxycholic acid or UDCA, in the treatment of COVID-19, may help in preventing inflammation and apoptosis, thereby, projecting new opportunities which still require to be evaluated. This study was in support of China promoting the use of Bear Bile, which is rich in the UDCA [60]. Hypothesis of usage of drugs like Pirfenidone and Thalidomide, that had initially been developed for the treatment of Idiopathic Pulmonary Fibrosis and morning sickness, respectively, have also gained much attention. Both of the drugs have been observed to decrease the effects of a number of released cytokines. Pirfenidone is also known to reduce the expression of the ACE2 host receptor and shows antioxidiant and antiapoptotic effects. On the other hand, Thalidomide shows anti-inflammatory effects, thus, making both these drugs potent fighters in the battle against COVID-19 [61, 62]. Another therapeutic option, which has also been observed to down-regulate the ACE2 and TMPRSS2 (activates spike protein in SARS-CoV-2) expression levels in the human liver derived cells, is Nrf2-activator PB125® [63].

### 3 Other Pharmacotherapies to Counter COVID-19

Apart from the various attempts in developing the monoclonal and polyclonal antibodies targeting either the human receptor, viral proteins or the immune response elements, the other major focus is towards the medications/drugs that have been repurposed, in order to treat the COVID-19 patients (Fig. 1). Some of these majorly used medications, such as Remdesivir, Lopinavir-Ritonavir, Hydroxychloroquine (HCQ), Favipiravir, etc. are extensively under clinical trials. Each of these drugs have their own mechanism of action, advantages and side effects of administration.

#### 3.1 Remdesivir

Remdesivir is a well-known antiviral drug. It is metabolized into its active adenine analogue that interferes with the activity of RNA dependent RNA polymerase (RdRp), further promoting the viral exoribonuclease activity of proofreading and thereby inhibit the synthesis of viral RNA. The drug is responsible for lowering the RNA levels of the virus by acting at an early stage of viral infection [64]. It had previously shown its efficacy while treating the Ebola virus infection in Rhesus monkeys and was also approved for use in treating infections caused by MERS and SARS-CoV. The potential of Remdesivir for the treatment of SARS-CoV-2 was proven in the clinical trials that were held in various parts of the world. The drug effectively treated 14 American cruise passengers in Japan who were infected by the novel virus and even patients in the USA who administered this drug after showing pneumonia symptoms [65, 66]. The importance of this drug was also stated in the guidelines issued by American Society
Fig. 1 Inhibitory functions and roles of different antibodies and repurposed drugs for battling COVID-19. The COVID-19 (coronavirus disease of 2019) infection begins with the attachment of the spike protein of the SARS-CoV-2 (severe acute respiratory syndrome coronavirus 2) to the ACE2 (angiotensin converting enzyme 2) human receptor. Various monoclonal (mAb) and polyclonal antibodies are being designed and convalescent plasma collected from the donor patients are being used to inhibit the attachment of the novel virus. This is followed by the virus and host membrane fusion and entry of the virus which is inhibited by drugs like Hydroxychloroquine (HCQ) and Chloroquine (CQ). Once the uncoating occurs and the viral RNA is released, it starts replicating with the help of the enzyme RNA dependent RNA polymerase (RdRp). Drugs like Remdesivir, Favi-piravir and Ribavirin, interfere with the activity of this enzyme. In order to further generate viral proteins, the processing (capping) of the viral RNA is required for maintaining its stability. This processing is inhibited by drugs like Ribavirin. On the other hand, combination of Lopinavir and Ritonavir is used for inhibiting the proteases like 3CL<sup>pro</sup> (chymotrypsin like cysteine protease) which are mainly responsible for the proteolytic cleavage and generation of non-structural proteins involved in viral replication. Once the viral assembly is complete and the new virions are ready, the exocytosis of these virions occur. This is inhibited by drugs like CQ and HCQ. The infection by the novel virus and release of the new virions which infect the neighbouring cells of the body lead to an exaggerated cytokine release which can be managed using drugs like corticosteroids, various mAbs targeting different cytokines and cytokine receptors, etc.

Post the approval by FDA on 1st of May 2020, the drug was also announced to be used as an emergency treatment option for COVID-19 treatment by the Drug Controller General of India on 1st of June, 2020. Companies such as Cipla, Jubilant Life Sciences, Dr. Reddy’s Laboratory, Ferozsons Laboratory and Zydus Cadila were approached to produce the drug in India. Hetero and Cipla, being the only approved manufactures priced the drug for Rs.5400 and less than Rs. 5000, respectively [71, 72].

3.2 Lopinavir-Ritonavir

Lopinavir-ritonavir act as inhibitors of proteases such as 3C like and papain-like proteases, which are important targets of the Ortho Coronaviridae family. In India, this drug has been previously used to prevent the transmission of HIV from infected mother to child. Amongst several other anti-HIV drugs which possess abilities of efficient polyprotein processing, it has been observed that Lopinavir showed the highest inhibitory activity against SARS-CoV-2. The recommended drug dosage includes 200 mg/50 mg. In patients having trouble taking oral dosage, 400 mg/100 mg...
Diseases, as a potential treatment for COVID-19 [74]. In of America and American Association for study of Liver therapy was promoted by both Infectious Disease Society its antiviral activity [65]. Ritonavir assisted combination vir, thereby increasing the half-life of the drug and hence ritonavir acts as an obstruction in the metabolism of Lopina-
This combination is sold under the brand Kaletra. Rito-
ritonavir is efficient in inhibiting the antiretroviral protease.

However, the clinical trials for the drug, conducted on 199 patients in Wuhan, that were further divided into two groups, one group getting the lopinavir -ritonavir combination while other treated with standard care showed no difference in the combination therapy as compared to the standard care. Though the clinical improvements were seen in both the groups. The adverse effects were less in the patients treated with the combination of both drugs, but no reduction in the levels of viral load was observed when compared with the standard care [76]. Even according to the British researchers, the combination of lopinavir and ritonavir showed no efficacy during the UK recovery trials of COVID 19 patients. The chief investigator of the trials conducted by the Oxford, stated that the drug had no clinical benefits in treating the COVID-19 patients [77] and hence more studies are required to be conducted in this regard.

3.3 Favipiravir

This antiviral drug was used to treat the influenza virus infection in 2014 after being approved in Japan. The drug hinders the viral replication, by binding to the conserved domain of polymerase which further puts a stop to the addition of nucleotides into the growing viral RNA, during replication [78]. The drug gets converted into an active phosphoribosyl form which is recognized by the viral RdRp (RNA dependent RNA polymerase) as a substrate thus, inhibiting its polymerase activity [79].

On 17th March, 2020, the Director of China National Centre for Biotechnology announced that Favipiravir was successfully tested during a clinical trial held on 80 patients infected by SARS-CoV-2 [80]. Studies were also conducted in other parts of China and Japan on a larger section of patients. It was observed in all these studies, that the patients administered with Favipiravir, showed to be negative for the infection caused by the virus, in a shorter period as compared to some other anti-viral drugs. Researchers from Stanford are also planning to conduct clinical trials for further checking the effectiveness of Favipiravir on 120 COVID-19 patients [81]. Recently the Glenmark Pharmaceuticals was given the approval for manufacturing the drug in India, on 19th June 2020. The recommended dosage of the drug is 1800 mg twice on the first day followed by 800 mg twice for the next 14 days. The drug is currently being sold at a price of Rs 103 per tablet for mild to moderate cases of COVID-19 [82].

3.4 Hydroxycloroquine in Combination with Azithromycin

Towards the end of World War II, the Germans found the substitute of quinacrine (compound derived from Cinchona bark) called Chloroquine (CQ), which was explored to be effective not only as an antimalarial drug but also against arthritis, lupus erythematosus rashes, prophylaxis and other diseases with symptoms of fever. Later, an even safer derivative of CQ, with an OH group added to its composition was developed, called HCQ. Though both these drugs have commonly been used for many years now, but its usual symptoms also include dermatological changes, pruritus, nausea, cardiovascular issues, headaches, abdominal pain, etc. [83]. The mechanism of action of both the drugs in treating COVID-19, is not very well studied. However, it is suggested that the intracellular phagolysosome is alkalised (rise in endosomal pH) by these drugs, thus preventing the fusion and uncoating of the viral membrane and hence the spread of the virus [84]. As per the observation made after the clinical trials of CQ/ HCQ, it was concluded that these drugs inhibit the entry of both SARS-CoV and SARS-CoV-2, by impeding the terminal glycosylation of ACE2 host receptors [85]. Studies show that HCQ had the better potential as a drug against COVID-19 as compared to CQ [86]. In view of a paper published in Lancet describing the concerns and increase in the mortality rate due to use of HCQ for the treatment of COVID-19, the WHO officially halted the ongoing clinical trials of the drug on 25th May, 2020. A week later, after further clarification and better understanding these trials were resumed [87, 88]. However, soon studies revealed that the benefits of the drug were not able to outnumber the risks that came along with the usage of this drug in the treatment of COVID-19. Thus on 15th of June, 2020, FDA in consultation with BARDA, announced the cancellation of its usage as an emergency authorized drug [89].

Azithromycin belonging to the macrolide class of antibiotics, is another drug used in combination with HCQ, for the treatment of COVID-19 patients. This drug is commonly used for the treatment of MAC (Mycobacterium avium complex), bronchitis, or pneumonia infections. A group of researchers from the University of New Mexico demonstrated that Azithromycin behaves like a weak base having acidotropic lyophilic properties and regulates the pH of the Golgi transport network as well as endosomes. Its
effects on various organelles were similar to that caused by HCQ [90]. In a controlled study on 86 infected patients, treated with a combination of HCQ and Azithromycin, negative results of the viral infection by 97.5%, were observed on day 5 after administration, implying it to be an effective therapeutic option against CoV-2 infection [91]. A comparative study also revealed that the combination of these two drugs helped cured 100% of the patients (under study) after 5–6 days while the recovery was observed in only 57.1% patients when treated with HCQ alone [92].

3.5 Ribavirin

Ribavirin is a drug that is not only known to interfere with the DNA and RNA polymerisation process in viruses, but also with the capping process of the RNA. It also inhibits the inosine monophosphate dehydrogenase, which is an essential enzyme for the production of guanosine [93]. The usage of this drug and its benefits were also analysed in earlier outbreaks of SARS-CoV and MERS-CoV, where it was used in combination with several other drugs like corticosteroids, lopinavir/ritonavir, oseltamivir, prednisolone, interferons, etc. However, no previously conducted studies gave any clear answers regarding its advantages [83, 94–96]. These previously conducted studies of the use of ribavirin in treating the viral infections caused by viruses similar to SARS-CoV-2, encouraged the use of this drug in treating patients of COVID-19 as well. Many researchers are still trying to evaluate the role of this drug in fighting the novel coronavirus, around the world [97–101]. However, many studies have yet proven its efficiency and even the optimum dosage prescriptions are yet to be finalised for the combinational or non-combinational use of this drug.

4 Discussion

The seamless spread of the novel coronavirus and the augmented rise in the number of infected patients, indicate that quarantine alone cannot help in controlling the devastating risks and threats that the SARS-CoV-2 infection is imposing on the world. As the spread of the virus and consequently the mortality rate of the infected individuals is increasing at a rapid pace, even the research and health/clinical care community are facing several challenges, globally, to find potential therapies or drugs to fight the novel virus. Studies related to the genome and structure of the novel coronavirus, has immensely boosted the process of designing rapid test kits, vaccines, antibodies and drugs to combat the viral infection. Many structural proteins of the SARS-CoV-2, host receptor: ACE2 and cytokines/chemokines have been projected to be significant drug targets. Although, it is important to emphasize that there exist many non-structural and accessory proteins of the virus, which are yet to be characterized and well-understood. These might also help in developing suitable therapeutic targets once their roles are well defined.

Several antiviral and immunomodulatory drugs, vaccines, antibodies, protease inhibitors and IFN and stem cell-based therapies are also under extensive study while some are under clinical trials. Very recently, India’s first COVID-19 “inactivated” vaccine, “COVAXIN”, developed by Bharat Biotech, had entered the first phase of the human clinical trials, in collaboration with ICMR and NIV, Pune [102]. Another Indian, plasmid DNA based vaccine, called ZyCoV-D was developed by Zydus Cadila, Ahmedabad. On 15th July, 2020, this vaccine also entered the Phase I/II human clinical trials [103]. Another adeno viral vector-based vaccine, called ChAdOx1 nCoV-19 or AZD1222, was helpful in generating a strong innate, humoral (development of neutralizing Abs within 28 days of vaccination) and cellular (development of T-cell to attack the virus within 14 days of vaccination) immune response to the RBD of the S protein of SARS-CoV-2. It was reported to be developed at the Oxford University in collaboration with Astra Zeneca. The vaccine had successfully completed the Phase I/II trials and has recently entered into the Phase III trials [104, 105]. Some mRNA-based vaccines have also gathered much attention in the recent times. These include the mRNA-1273, developed by Moderna in consortium with NIAID (National Institute of Allergy and Infectious Diseases), U.S. The vaccine utilizes the ability of the mRNA to encode for a full-length S protein, which then commences an immune response allowing generation of Abs against SARS-CoV-2. This vaccine candidate entered into the Phase 3 clinical trials on 27th July, 2020 [106, 107]. Pfizer in association with BioNTech have also developed a similar mRNA vaccine called BNT162b2, with a single nucleoside modification in the mRNA sequence of the S protein. The Phase 2/3 trials have also begun for this vaccine [108].

Even though the vaccine development is occurring at an unprecedented rate, it will take a long time to manufacture and vaccinate a large number of people worldwide. In order to bridge this gap, the focus shifted towards the development of antibody-based therapy and repurposing drugs in order to meet the need of the hour (Table 1). Multiple other drugs apart from the repurposed drugs discussed earlier are being evaluated for their potential use in combating COVID-19. These include Ivermectin, Nitazoxanide, Lianhuaoqingwen capsules, Dipyridamole, etc. [109].

Though such extensive research work is being carried out all over the world, no specific treatment/therapy/drug for COVID-19 is yet available. There are several challenges to overcome, which not only includes the development of effective therapeutic options and diagnosis kits, but also, select, evaluate and mass produce them. There is also a need to conduct further studies to explore and perform in depth
analysis about the pathogenesis of the novel virus, which could possibly lead us to identify new drug targets. Thus, the greatest challenge today undoubtedly requires the entire research community to work with cooperation towards the common goal and the day is not far off, when a therapeutic agent will prove to be effective against COVID-19.

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Compliance with Ethical Standards

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Table 1 A summary of currently available therapeutic drugs

| Therapeutic drugs       | Mechanism of action                                                                 | Currently under usage/trial | References |
|-------------------------|-------------------------------------------------------------------------------------|-----------------------------|------------|
| Remdesivir              | Blocks the viral synthesis by interfering with activity of RdRp (RNA dependent RNA polymerase) | +                           | [64]       |
| Lopinavir-ritonavir     | Inhibits proteases such as Papain and 3C                                            | −                           | [65]       |
| Favipiravir             | Blocks the addition of nucleotide into the viral RNA, by binding to conserved domains of RdRp | +                           | [78]       |
| Azithromycin and Hydroxychloroquine combination | Viral membrane fusion (endocytosis) and exocytosis (viral release) is blocked, inhibition of Interleukins (IL) like IL-6 | +                           | [90]       |
| Chloroquine             | Increase the pH and thus decrease the viral membrane fusion and inhibit ACE2 (Angiotensin converting enzyme 2) glycosylation thus preventing virus binding | −                           | [84, 85]  |
| Ivermectin              | Blocks the terminal glycosylation of ACE2 receptor                                  | −                           | [110]      |
| Nitazoxanide            | Blocks IMPα/β1 and thus prevent nuclear-cytoplasmic translocation                  | −                           | [111]      |
| Tocilizumab (Actemra)   | Inhibit the glycosylation of spike protein                                          | +                           | [46, 47]  |
| Dexamethasone           | Anti-CD-6 IgG1 mAb (monoclonal antibody)                                           | +                           | [113]      |
| Octagam                 | Immunoglobulin mixture                                                               | +                           | [114]      |
| Emapalumab              | Anti-inflammatory response                                                           | −                           | [50]       |
| Infliximab and Eternacept| Binds to both free and receptor bound IF γ (interferon gamma)                      | +                           | [50]       |
| Lenzilumab              | Target TNF α (Tumor Necrosis Factor alpha)                                          | +                           | [53]       |
| Baricitinib and Ruxolitinib| GM-CSF (Granulocyte–macrophage colony-stimulating factor) targeting drug            | −                           | [57]       |
| Canakinumab             | Controls the elevated interleukin (IL)-1β levels                                    | +                           | [115]      |
| Sarilumab               | IL-6 inhibitor                                                                      | +                           | [114]      |
| Pirfenidone and Thalidomide| JAK-STAT (Janus Kinase/Signal Transducer and Activator of Transcription) signaling inhibitors | −                           | [61, 62]  |
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