SARS-CoV-2 therapeutics: how far do we stand from a remedy?

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Abstract
The SARS-CoV-2 has affected millions worldwide and has posed an immediate need for effective pharmacological interventions. Ever since the outbreak was declared, the medical fraternity across the world is facing a unique situation of offering assistance and simultaneously generating reliable data with high-quality evidence to extend the scope of finding a treatment. With no proven vaccine or other interventions available hitherto, there is a frenzied urgency of sharing preliminary data from laboratories and trials to shape a global response against the virus. Several clinical trials with investigational and approved repurposed therapeutics have shown promising results. This review aims to compile the information of the reported molecules approved for emergency use and those under clinical trials and still others with good results in the studies conducted so far. Being an RNA virus, SARS-CoV-2 is prone to mutation; thus, the possibility of gaining resistance to available drugs is high. Consequently, a cocktail therapy based on drug interaction with different stages of its replicative cycle is desirable to reduce the chances of evolving drug resistance. Since this virus encodes several proteins, including 16 nonstructural and 4 structural proteins, this review also offers an insight into potential drug targets within SARS-CoV-2.

Keywords Antiviral · COVID-19 · Drug repurposing · SARS-CoV-2 · Target proteins

Abbreviations
3CLpro  Chymotrypsin-like protease
ACE2  Angiotensin-converting enzyme 2
ARDS  Acute respiratory distress syndrome
COVID-19  Coronavirus Disease 2019
CQ  Chloroquine
HCQ  Hydroxychloroquine
IFN  Interferon
MERS-CoV  Middle East Respiratory Syndrome Coronavirus
Nsp  Nonstructural protein
ORF  Open reading frame
RCT  Randomised Controlled Trial
RdRp  RNA-dependent RNA polymerase
SARS-CoV  Severe Acute Respiratory Syndrome Coronavirus
SARS-CoV-2  Severe Acute Respiratory Syndrome Coronavirus 2
TMPRSS2  Transmembrane Serine Protease 2
RECOVERY  Randomised Evaluation of COVID-19 Therapy

Introduction
Very recently, in December 2019, a novel coronavirus, now designated as Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2), triggered a pneumonia outbreak in Wuhan, China [1], and since then has spread rapidly across the world. This outbreak was declared to be a “pandemic by the World Health Organisation (WHO) on March 11, 2020”. SARS-CoV-2 belongs to the same family of viruses as Severe Acute Respiratory Syndrome Coronavirus (SARS-CoV) and the Middle East Respiratory Syndrome Coronavirus (MERS-CoV), which caused outbreaks in 2002 and 2012, respectively. According to Johns Hopkins University and Medicine Coronavirus Resource Center (https://coronavirus.jhu.edu), there have been 56,901,880 laboratory-confirmed cases with over

We do not recommend these drugs to be taken without a valid prescription. Guidelines from regulating authority must be followed strictly.

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1,360,408 deaths in more than 191 countries/regions worldwide as on November 20, 2020. With approximately 100,000 new cases appearing every day since May 15 which has increased to approximately 150 thousand by June 19, 2020, we are still in the middle of a pandemic, heedless when the downside of the pandemic will begin. COVID-19 (Coronavirus Disease 2019) presents with symptoms like fever, cough, fatigue, phlegm production, hemoptysis, headache, diarrhoea, lymphopenia, and shortness of breath, appearing after a mean incubation period of about 5.2 days. These could be mild or asymptomatic to severe ones progressing to pneumonia and other complications like Acute Respiratory Distress Syndrome (ARDS) which could be fatal, especially in older patients with comorbid conditions [2–5]. ARDS might result from 'cytokine storm' which occurs due to the release of large amounts of proinflammatory cytokines, including interleukin-6 (IL-6) and tumour necrosis factor-alpha (TNFa). ARDS was also presented during the SARS outbreak, wherein during endocytosis of SARS-CoV, ACE2 receptor gets simultaneously endocytosed, leading to reduced ACE2 on the cell surface. ACE2 is an inactivator of angiotensin 2 (AngII) and the former’s reduced concentration leads to increased levels of AngII. AngII is not only a vasoconstrictor but also a pro-inflammatory cytokine [3, 6–9]. Thus, effective treatment of moderate cases is desirable to prevent progression into severe ones, to reduce the overall mortality rate.

In the absence of either the approved therapeutics or any effective vaccine against SARS-CoV-2, we completely rely on an individual’s immune system to clear the infection. Strict isolation norms and extended contact tracing can help in what we call ‘flattening the curve’ and, hence, prevent the healthcare system from being overwhelmed. As of November 20, 2020, the worldwide fatality rate stands at 2.39% (1,360,408/56,901,880) and varies greatly in different countries. China, where the outbreak was recorded first, has 5.15% (4742/91,935) fatalities; the USA has 2.15% (252,555/11,717,827), India records its fatality rate at 1.46% (132,162/9,004,365) and most of the European countries recorded higher fatalities. Rate of fatality varies widely in different territories across continents with lower than 1% in the middle-east to higher than 9% in Mexico (https://coronavirus.jhu.edu).

There could be many effective therapeutic targets within the life cycle of SARS-CoV-2 including:

1. Blocking the viral attachment to host cells by disrupting bonding between viral cell attachment proteins and the cellular receptor, and inhibiting Spike protein cleavage by the host TMPRSS2 which stimulates the formation of endosome facilitating virus internalisation.
2. Inhibition of transcription, translation, and replication of the viral genome.
3. Stalling the assembly process by targeting structural and nonstructural proteins critical in viral assembly.

Coming up with exclusive drugs for clinical presentations takes years of effort and billions of dollars (~$2.5 billion) [10] in investments. Time constraints posed by a sudden and novel pandemic makes it almost impossible to propose such novel molecules by conventional protocols. Thus, efforts to explore off-targeting of already approved molecules remains the only hope to possibly provide an answer. Drug repurposing saves the precious time and money in the development, formulation and the subsequent phases of safety trials. Synergistic use of both computational and experimental approach to delineate molecule(s) of interest and to validate them in cell lines and animal models can expedite subsequent efficacy trials in humans [11]. In this paper, we try to review several potential repositioned therapeutics that have emerged from imprints in different countries. A range of already in use drugs have been shown to bind to important targets of SARS-CoV-2. These repurposed medications could possibly be used in a cocktail therapy subject to validation of drug interaction on a case-to-case basis. The purpose of this review is to compile the knowledge on the therapeutic potential of the reported molecules. We made a modest attempt to aid scientific community to design newer molecules using the reported ones as HITS and to formulate novel drug combinations to achieve enhanced outcomes. Moreover, this review also gives an insight into the drug targets within SARS-CoV-2.

Proteins encoded by SARS-CoV-2 genome as potential therapeutic targets

SARS-CoV-2 contains a single-stranded positive-sense RNA (+ ssRNA) of 29,903 nucleotides in length with 5’-cap and 3’-polyA tail. Typically coronavirus genome transcribes into 10 ORFs (1 large genomic and at least 9 smaller sub-genomic) of which the first ORF (ORF 1a/b) encodes for the key polyproteins pp1a and pp1ab and covers nearly 60% of the entire genome. Other ORFs present in the downstream one-third region of the genome encode for four structural proteins along with special accessory proteins. These accessory proteins differ in different coronaviruses, with SARS-CoV-2 containing 6 of them [12, 13]. Further, viral 3 chymotrypsin-like protease (3CLpro) or main protease (Mpro) with papain-like protease (PLpro), process these polyproteins into 16 nonstructural proteins: nsp1–16. These nonstructural proteins perform all the vital functions in viral replication and infection and hence, many among them are important drug targets [13, 14].
Nonstructural proteins like Nsp1 induce host mRNA degradation and strongly inhibit host protein synthesis. Additionally, it is reported to block the host’s innate immune response, and hence plays an important role in the infection process [15, 16]. Nsp3 is a multi-domain protein with at least 8 conserved domains throughout the family Coronaviridae [17]. It is the largest protein and performs many functions to support the survival of the virus by interfering with the host’s antiviral response. In the early phase of infection, its binding to nucleocapsid protein is essential to establish infection and genome replication [18, 19]. The papain-like protease (PLpro), a subdomain of Nsp3, is responsible for cleaving Nsp1, 2, and 3 from the polyprotein [20]. Moreover, PLpro now is also known to demonstrate de-ubiquitinating and de-IGSylating (removal of ubiquitin-like protein ISG15) activities, justifying the role of Nsp3 in suppressing host’s immune response [21]. Nsp3b, also known as ‘X-domain’ (Macro-domain 1) has a predicted function of ADP-ribose-1″-phosphatases (ADRP). Thus, various domains of Nsp3 in themselves are attractive targets for novel therapeutics [22, 23]. Along with Nsp4, Nsp3 is also reported to play an important function in the rearrangement of host-derived membranes [24]. Nsp5, also known as 3CLpro, is auto-cleaved from polyprotein, and it then further cleaves downstream to ensure maturation of Nsp4–Nsp16 [25]. 3CLpro is highly conserved in SARS-CoV-2 and its function in processing key replicative proteins makes it an attractive anti SARS-CoV-2 target [26]. Nsp6 of SARS-CoV-2 generates auto-phagosomes. It is also reported to induce the formation of double-membrane vesicles (DMVs) along with Nsp3 and 4 [27, 28]. A complex of Nsp7–Nsp8 functions by enhancing the activity of RdRp and helps it bind to its substrate, i.e., RNA [29]. Nsp8 functions as a second RdRp and is involved in the synthesis of a primer, an oligonucleotide of fewer than 6 residues required by the primer-dependent main RdRp (Nsp12) [30]. Nsp9 in SARS-CoV is present as a dimer and binds RNA. It is known to mediate replication and virulence [31, 32]. Two critical proteins in a complex, nsp10/16, modify the viral genomic RNA to make it appear more like the host cell RNA and allow the virus to hide from the cells. Designing drugs to inhibit nsp10/nsp16 interaction would make it easier for the cells to detect the viral genome and clear it faster within the cell [33, 34]. Nsp12 or RNA-dependent RNA polymerase (RdRp) remains conserved in coronaviruses and synthesises the RNA strand in a primer-dependent fashion [30, 35], making it a very good target for novel therapeutics and some of the currently approved drugs such as remdesivir are directed against it. Helicase or Nsp13 works in 5′–3′ direction to separate double-stranded DNA and RNA. It is highly conserved in coronavirus and helps in its replication [36], making it a suitable potential target. Earlier recognised as an Mg2+-dependent 3′–5′ exonuclease, which played a crucial role in proof-reading and repair, Nsp14 is now also found to function as N7 Methyltransferase (N7-MTase) to methylate the RNA cap at the N7 position. Interestingly, its exonuclease activity also removes the incorporated nucleoside analogues from RNA and, thus, poses a challenge to Nucleoside analogue (NA) drugs [34, 37–39]. This makes Nsp14 a suitable target that should be simultaneously targeted, along with NAs targeting RdRp. While Nsp15 functions in viral replication and transcription as a Mn2+-dependent nidoviral uridylate-specific endoribonuclease (NendoU), present in Nidovirales order to which the family Coronaviridae belongs [40]. Nsp16 is a 2′-O-methyltransferase (2′-O-MTase) and in complex with Nsp10 helps to methylate cap-0 to form cap-1 structure [33, 41]. Interestingly, it was reported that NendoU fails to cleave RNAs with 2′-O-ribose methyl groups, which possibly explains the close association of Nsp15 and Nsp16 [42].

Like all other beta-coronaviruses, SARS-CoV-2 genome encodes four structural proteins, i.e., Spike (S), nucleocapsid (N), membrane (M), and envelope (E), which play a vital role in its attachment to host cell, replication, and assembly of the mature virus [43]. With around 75% homology to SARS-CoV, the spike protein of SARS-CoV-2 extends till an ecto-domain fragment via a trans-membrane moiety starting from a short cytoplasmic segment [44, 45]. S protein is present as a homo-trimer, with each monomer consisting of S1 and S2 subunit [45]. S1 helps in the attachment to host cells while the S2 subunit mediates membrane fusion during infection. S protein is required to be cleaved by host cell protease TMPRSS2 to facilitate the internalisation of the virus [46]. S1 subunit hosts a receptor-binding domain (RBD) that binds the human ACE2 (Angiotensin-Converting Enzyme 2) [46] with a better affinity than that in SARS-CoV [45]. A cascade of events then leads to disruption of the S1 subunit and formation of a more stable S2 subunit, thus facilitating the fusion of viral and cell membranes [46, 47]. Hence, spike protein determines the host range and primarily interacts with the host receptor to establish infection, making it an attractive target for therapeutics [43]. Envelope proteins are integral structural proteins that form viroporins and mediate pathogenesis [48]. While E protein (form E channel) plays an important role in virus assembly and release [49], the M or Membrane protein helps shape the virions and in the interaction of viral envelope proteins (S and E) with the nucleocapsid protein, thus playing a central role in mediating viral assembly [50]. Disrupting its interaction with either N or E or S protein can stall the assembly process. Nucleocapsid or N protein binds the viral RNA and helps in its encapsidation. Upon fusion of viral and host membranes, N protein interacts with cellular processes and helps to wind down the host cell replication and overall immune response [43, 51]. This makes all the structural proteins important for directing novel therapeutics.
Table 1  Structural and nonstructural proteins of coronavirus and their functions

| S no | PROTEIN/(references) | Function | PDB ID (SARS-CoV-2) |
|------|-----------------------|----------|----------------------|
| 1    | Spike Protein [43–47] | Attach to host cell (receptor); membrane fusion during infection | 6VXX: S protein (closed state) 6VYB: S protein (open state) 6VW1: RBD–ACE2 complex 6LXT, 6M1V: Post fusion core of S2 subunit |
| 2    | Envelope Protein [48, 49] | Helps in viral assembly and release; form viroporins | Unavailable |
| 3    | Membrane Protein [50] | Shape the virions; helps in mediating interactions of other structural proteins and therefore plays a central role in the assembly | Unavailable |
| 4    | Nucleocapsid Protein [43, 51] | Binds viral RNA; helps in encapsidation; post-fusion processing to enhance survival | 6M3M: N-terminal Domain (NTD) of N protein 6WJ7: C-terminal Domain (CTD) of N protein |
| 5    | Nsp1 [15, 16] | Critical virulence factor: Suppresses host innate immune response, degrades host mRNA, and inhibits host protein synthesis | Unavailable |
| 6    | Nsp2 [20] | Unclear; may function in the infection process by making virions more contagious | Unavailable |
| 7    | Nsp3 [17–23] | It has multiple subdomains: an acidic domain (Nsp3a); Nsp3b, also known as ‘X-domain’, function as ADP-ribose-1″-phosphatases (ADRP); Nsp3c is a SARS unique domain; Nsp3d functions as papain-like protease and also acts to suppress the interferon response | 6WEY: ‘X domain’ of Nsp3 6W02: ADRP complex with ADP ribose 6W6Y: ADRP complex with AMP 6W9C: PLpro 6WX4: PLpro in complex with peptide inhibitor VIR251 6WUU: PLpro in complex with peptide inhibitor VIR250 |
| 8    | Nsp4 [24] | Along with Nsp3 plays an important role in rearrangement of host-derived membranes; induces the formation of double-membrane vesicles (DMVs) | Unavailable |
| 9    | Nsp5 [25, 26] | Chymotrypsin-like protease (3CLpro) which is highly conserved in coronaviruses. It cleaves downstream and ensures maturation of Nsp4–Nsp16 | 6M2Q: 3CLpro apo structure 6LU7: Main protease in complex with an inhibitor N3 |
| 10   | Nsp6 [27, 28] | Generates autophagosome; induces the formation of double-membrane vesicles (DMVs) | Unavailable |
| 11   | Nsp7 [29] | Nsp7–Nsp8 enhances enzyme activity and thus processivity of RdRp and helps it bind to RNA | 6WQ: 6WQD: Complex of Nsp7 and C-terminal domain (CTD) of Nsp8 |
| 12   | Nsp8 [29, 30] | Primer-independent RdRp; synthesise primers to be used by Nsp12 RdRp | Nsp7–Nsp8 complex [Refer Nsp7] |
| 13   | Nsp9 [31, 32] | Binds RNA; mediates replication and virulence | 6W4B: Nsp9 |
| 14   | Nsp10 [33, 34] | Promotes binding of Nsp16 to both S-adenosyl-L-methionine (SAM) cofactor and RNA substrate | 6W75, 6W4H, 6WKS: Nsp10–Nsp16 Complex 7C2J: Nsp16–Nsp10 complexed with SAM |
| 15   | Nsp11 | Unclear | Unavailable |
| 16   | Nsp12 [30, 35] | Primer-dependent RdRp; conserved in coronaviruses | 7BW4: RdRp 6M71, 7BV1: nsp12(RdRp)–nsp7–nsp8 complex 7BV2: Nsp12–Nsp7–Nsp8 complexed with template-primer RNA and remdesivir triphosphate |
| 17   | Nsp13 [36] | Helicase; works in 5′–3′ direction | 6ZSL: Helicase (nsp13) 6XEZ: Helicase bound to replication–transcription complex |
| 18   | Nsp14 [34, 37–39] | N7 Methyltransferase (N7-MTase) [generate cap-0 structure]; Mg2+-dependent 3′-5′ exoribonuclease | Unavailable |
| 19   | Nsp15 [40, 42] | Mn2+ dependent nidosiral uridylic-specified endoribonuclease (NendoU) | 6VWW: Nsp15 6W01: Nsp15 complexed with a citrate |
| 20   | Nsp16 [33, 41, 42] | 2′-O-Methyltransferase (2′-O-MTase) [Nsp10/ Nsp16 methylates cap-0 to form cap-1structure]; help virus escape host immune response | Nsp10-Nsp16 Complex [Refer Nsp10] |
and MERS coronaviruses and also against SARS-CoV-2
and has shown in vitro therapeutic efficacy against SARS
metabolised to an analogue of adenosine triphosphate
It works by inhibiting viral RNA polymerases after being
initially designed against Ebola by Gilead Sciences, Inc.
Remdesivir is a prodrug of a nucleotide analogue and was
compiled promising candidates for drug repositioning and
focusing on drug repurposing to achieve a fast resolution.
While this approach could save pharmaceutical companies
in capitals, the odds of accomplishing a cure are of more
interest (Extensively reviewed by Pushpakom and cowork-
eries [11]). Multiple trials are being done across the world
to test several therapeutic options for COVID-19, mostly
focusing on drug repurposing to achieve a fast resolution.
These are enlisted in drugbank’s COVID-19 information
dashboard (https://www.drugbank.ca/covid-19) and Milken
Institute’s COVID-19 treatment and vaccine tracker (https :
://covid-19tracker.milkeninstitute.org/). In this review, we
compiled promising candidates for drug repositioning and
the results of their clinical trials as of now. Table 2 provides
a compiled list of all the described repurposed therapeutics
in various stages of clinical trials.

**Remdesivir**

Remdesivir is a prodrug of a nucleotide analogue and was
initially designed against Ebola by Gilead Sciences, Inc.
It works by inhibiting viral RNA polymerases after being
metabolised to an analogue of adenosine triphosphate
and has shown in vitro therapeutic efficacy against SARS
and MERS coronaviruses and also against SARS-CoV-2
with Nsp3b [have a predicted function of ADP-ribose-1″-
phosphatases (ADRP)], RdRp, E-channel (E protein), and
type-II transmembrane serine protease (TMPRSS2) enzymes
as potential binding targets [53–55]. With several studies
in moderate and severe COVID-19 patients still on, in one
of the trials in a small cohort, Grein and co-workers sug-
ject that remdesivir may have clinical benefits in severe
COVID-19 patients with improvement in oxygen-support
status. Though, the majority of them also experienced com-
mon side effects like increased hepatic enzymes, diarrhoea,
renal impairment, and hypertension. However, with a small
sample size, it was also reported that improvement was com-
paratively less frequent in cases with invasive ventilation
than those on non-invasive support and in patients above
70 years as compared to those younger than 50 years [56].
Intravenous remdesivir was administered to the first con-
firmed COVID-19 patient in the United States, and improved
clinical symptoms were observed [57].
However, findings of another randomised, double-blind,
placebo-controlled clinical trial in Hubei, China, found
no significant benefits of treatment of severe COVID-19
with remdesivir over normal supportive care. Wang and
co-workers reported that patients receiving remdesivir
showed faster clinical improvement, but this was statisti-
cally insignificant [58]. Another preliminary report from a
randomised, controlled trial [Adaptive COVID-19 Treatment
Trial 1 (ACTT-1)] comprising 1063 patients suggests that
treatment with remdesivir lead to 31 percent faster recov-
ery time, the median of which was 11 days as compared
to 15 days of the group, that received placebo [59]. These
results prompted successive trials ACTT-2 (Remdesivir plus
Baricitinib against Remdesivir; NCT04401579) and ACTT-3
(Remdesivir plus Interferon Beta-1a against Remdesivir;
NCT04492475) to formulate better treatment regimen (https :
c://clinicaltrials.gov/). Remdesivir is now cleared for emer-
gency use by the US FDA (Food and Drug Administration)
[60].

**Lopinavir and ritonavir**

Lopinavir is an antiviral used to treat Human Immuno-
deficiency Virus (HIV) type 1 infection. It inhibits HIV pro-
tease, and when used along with Ritonavir, the plasma half-
life of lopinavir increases. It has shown inhibitory activity
against SARS-CoV in vitro [61].
In a randomised, controlled, open-label trial in 199
patients, Cao and co-workers observed that there was no
significant improvement in clinical signs or mortality upon
treatment in 99 patients of the lopinavir–ritonavir group.
The authors found that though the treatment reduced the
patient’s stay in intensive care units (ICU), there were sig-
nificant gastrointestinal adverse events in this group against
those which did not [62]. These findings were substantiated.
### Table 2: List of repurposed therapeutics: with clinical trial result

| Sl. no | DRUG/references                                      | Primary use                                      | Potential target                                      | comments                                                                                                                                 |
|--------|------------------------------------------------------|--------------------------------------------------|-------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------|
| 1      | REMDESIVIR [53–60]                                   | Antiviral against EBOV (NA; targets RdRp)        | Nsp3b, RdRp, E-channel, and TMPRSS2                   | 1. Small cohort: effective<br>2. RCT: no significant benefits<br>3. RCT: faster recovery<br>4. Approved for emergency use by FDA |
| 2      | LOPINAVIR\(^A\) and RITONAVIR\(^B\) [61–65]          | Antiviral against HIV1 (\(^{\text{A,B}}\) protease inhibitor) | 3CL like protease                                      | 1. RCT: No significant improvement, major side effects<br>2. Intervention arm suspended in the SOLIDARITY trial (WHO) and RECOVERY (Randomised Evaluation of COVID-19 Therapy) trial |
| 3      | ARBIDOL\(^C\) and OSELTAMIVIR\(^D\) [55, 63, 64, 68, 69, 72] | Antiviral against Influenza virus (\(^{\text{C}}\)prevents fusion of viral and cell membrane; \(^{\text{D}}\)Neuraminidase inhibitor) | Nsp7/Nsp8 complex, Nsp14, Nsp15, E-channel, or Spike | 1. RCT: Less effective than favipiravir<br>2. Retrospective study: can drastically reduce the mortality rate<br>3. Retrospective study: early administration of arbidol (preferably with IFN) can reduce viral shedding |
| 4      | FAVIPIRA VIR [63, 65–68]                            | Antiviral against Influenza virus (targets viral RdRp) | RdRp                                                  | 1. Low binding score in docking studies<br>2. RCT: more effective than arbidol. It can prevent progression to ARDS |
| 5      | DANOPREVIR\(^E\) and RITONAVIR\(^F\) [70–73]         | Antiviral against HCV (\(^{\text{E}}\)protease inhibitor) (\(^{\text{F}}\)upregulate MHC I expression) | 3′-5′ exo-ribonuclease (Nsp14), endo-ribonuclease (Nsp15) | RCT: Can improve the clinical presentation |
| 6      | RIBAVIRIN\(^G\) and LOPINAVIR–RITONAVIR and IFN [53, 55, 73, 75, 77, 83] | Antiviral against HCV (\(^{\text{G}}\)guanosine analogue; targets RdRp and also interfere with RNA capping) | Plpro                                                 | 1. Open-label randomised trial: Triple combination of ribavirin, lopinavir–ritonavir, and IFN led to faster negative conversion, reduced hospital stay, and improved clinical presentations |
| 7      | DARUNAVIR\(^H\) and COBICISTAT [55, 80, 81]          | Antiviral against HIV-1 (\(^{\text{H}}\)protease inhibitor) | Nsp3c, Plpro, E-channel, or Spike proteins            | 1. No in vitro activity<br>2. RCT: Ineffective |
| 8      | FAMOTIDINE\(^I\) and HCQ [55, 82, 83, 88]           | Gastric acid suppression (\(^{\text{I}}\)histamine-2 receptor antagonist) | 3CL.pro                                               | 1. Retrospective study: Improve clinical symptoms<br>2. Proof-of-principle trial: combination of dual histamine blocker (cetirizine and famotidine) is effective in controlling disease progression |
| Sl. no | DRUG/references | Primary use | Potential target | comments |
|--------|-----------------|-------------|------------------|----------|
| 9      | HCQ\(^{1}\)     | Antimalarial drug; and \(^{2}\)a broad-spectrum antibacterial | RdRp      | 1. RCT: Effective, synergistic effect with azithromycin  
2. RCT: Shortened time to clinical recovery  
3. Prospective study: Inconclusive clinical evidence  
4. Randomised uncontrolled trial: Red flagged for safety concern with high dosage  
5. RCT: No significant benefit  
6. WHO temporarily paused trial for HCQ over safety concern but later resumed on June 03, 2020  
7. US FDA emergency use approval revoked on June 15, 2020 |
|        | HCQ and AZITHROMYCIN\(^{K}\) \[53, 67, 84–94\] |            |                  |          |
| 10     | TOCILIZUMAB     | Rheumatoid arthritis, and other inflammatory and autoimmune disorder | IL-6 receptor antagonist | 1. Retrospective study: Improve clinical presentations without significant side effects  
2. Significant reduction in the requirement of mechanical ventilation  
3. RCT: No mortality benefit by tocilizumab intervention |
| 11     | IVERMECTIN\(^{L}\) and DOXYCYCLINE\(^{M}\) \[101–107, 114, 115\] | Antiparasitic against worm infestations; and \(^{M}\)a broad spectrum antibacterial | 3CLpro, HR2 domain (spike protein) | 1. In vitro activity at IC\(_{50}\) of \(\sim 2 \mu M\)  
2. Retrospective study: effective in reducing mortality rate (even in subgroup of severe COVID-19 patients)  
3. Pilot-intervention trial: Ivermectin used as add-on therapy with HCQ and azithromycin showed better efficacy than control, with shorter duration of hospital stay and reduced time to turn negative in PCR test  
4. A medical team from Bangladesh claims to have found it effective |
| 12     | DEXAMETHASONE   | Endocrine, rheumatic and other conditions | Glucocorticoid receptor agonist | 1. The first drug to show a reduced mortality rate in severe cases  
2. Few studies also reported negative impact of corticosteroid |
| 13     | METHYLPREDNISOLONE | Dermatologic, allergic and other conditions | Glucocorticoid receptor agonist | 1. Shown to decrease the risk of ICU admission and the need for ventilation support  
2. Several randomised trials are ongoing |
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by other randomised controlled trial and docking studies [63]. Lopinavir–ritonavir SOLIDARITY trial (WHO) and RECOVERY (Randomised Evaluation of COVID-19 Therapy) trial were suspended after failing to find any clinical benefits [64, 65].

Arbidol/favipiravir/arbidol and oseltamivir

Arbidol, a broad-spectrum antiviral drug used against influenza and other respiratory viral infection in Russia and China, works by inhibiting the fusion of the virus’ lipid membrane with the host cells [55, 66]. Favipiravir, an antiviral used primarily against influenza infection, works by targeting the influenza viral RNA-dependent RNA polymerase (RdRp) [67]. It has a broad spectrum activity owing to the conserved nature of the catalytic domain of RdRp among various types of RNA viruses [68]. Both Influenza and SARS-CoV-2, being the RNA viruses, require RdRp for genome replication and have similar clinical presentations. Efficacy of both arbidol and favipiravir is now being investigated for the treatment of COVID-19. Docking studies predicted arbidol to potentially bind to Nsp7–Nsp8 complex, Nsp14 (exoribonuclease), Nsp15 (endoribonuclease), E protein, and/or Spike protein [55]; and favipiravir to bind RdRp in SARS-CoV-2 [69]. However, Chen and co-workers, in a prospective, multicenter, open-labeled, randomised superiority trial, found favipiravir to have a superior clinical recovery rate as compared to arbidol in moderate patients with COVID-19 infection [70] and indicated its use to prevent progression of disease into Acute Respiratory Distress Syndrome.

Another RCT supported this by concluding that treatment with arbidol may not improve clinical outcomes in mild/moderate COVID-19 patients [63]. However, in a retrospective study of 504 patients with confirmed COVID-19 infection in China, Liu and co-workers came to a different conclusion. On carefully studying data of these patients, the authors found that arbidol treatment led to slightly higher SpO2 levels and showed faster lesion absorption. This preprint report suggests that arbidol either alone or in combination with oseltamivir can drastically reduce mortality rate [71]. Further, in a retrospective study in 238 hospitalised COVID-19 patients in China, a prolonged viral shedding was correlated with delay in arbidol intervention (more than 7 days after illness onset) compared to those patients which received arbidol within 7 days of illness onset. Also the viral shedding was reduced in patient receiving combination of arbidol and interferon, compared to those receiving arbidol alone [72]. Oseltamivir as monotherapy or in combination with the other drugs is currently under multiple trials (NCT04303299, NCT04338698, NCT04457609, NCT04261270) to ascertain its efficacy in the treatment of the COVID-19 patients.
Danoprevir/danoprevir and ritonavir/danoprevir and ritonavir along with interferon

Approved and marketed in China since 2018, Danoprevir is an antiviral used to treat chronic hepatitis C patients by inhibiting HCV protease (NS3/4A) and is demonstrated to be safe and well tolerated in HCV patients. It is often boosted with ritonavir (an HIV protease inhibitor in itself at high dosage), to enhance its plasma concentration [73, 74]. Danoprevir is predicted to bind to 3′-5′ exoribonuclease (Nsp14), endoribonuclease (Nsp15) of SARS-CoV-2 [75].

A study suggests that in moderate COVID-19 patients, 4–12 days treatment of danoprevir/ritonavir in the presence or absence of interferon nebulisation can suppress viral replication and improve clinical conditions. In a trial with 11 patients, Chen and co-workers found that patients with hypertension or those who did not respond to treatment with lopinavir/ritonavir with or without worsening of symptoms were showing negative PCR test after treatment with danoprevir/ritonavir at a median of 2 days, ranging from 1 to 8 days with reduced ground-glass opacity [76]. This study also stresses the importance of containing the disease from progressing from a moderate to severe level. However, randomised controlled trials with a larger cohort are required to further substantiate the usefulness of danoprevir in the treatment of SARS-CoV-2 infection.

Ribavirin/ribavirin and interferon

Ribavirin is a guanosine analogue that, through multiple mechanisms, interferes with the replication of RNA and DNA viruses. Other than targeting polymerases, it is also found to interfere with RNA capping that is dependent on natural guanosine, by inhibiting the latter’s generation, and hence destabilises viral RNA. Viral replication in the presence of ribavirin occurs with reduced fidelity inducing random mutations, further reducing its viability [77, 78]. In safety trials of ribavirin for chronic HCV infection, the adverse effects were well established and were found to be well tolerated [79]. However, it is contraindicated during pregnancy, owing to its teratogenic properties [80].

Ribavirin was used during SARS infection in 2003, but findings remained unclear due to the deleterious effects of previous treatment and demanded more placebo-controlled trials. Likewise, during the MERS outbreak, the benefits of administering ribavirin were not sufficient to outplay its toxicity. With promising in vitro results, then again SARS-CoV and MERS-CoV and now against SARS-CoV-2, ribavirin is currently under clinical trial (NCT04356677) in combination with standard care therapy in Canada to check its efficacy against SARS-CoV-2 in humans [53, 61, 80, 81]. Chinese guidelines for the treatment of COVID-19 mention ribavirin in combination with IFN alpha as one of the therapeutic options [82]. Docking studies further suggest PLpro of SARS-CoV-2 as a potential target for ribavirin with low binding energy [55] in addition to RdRp. In an open-label randomised trial in 127 patient, a triple combination of ribavirin, interferon beta-1b and lopinavir–ritonavir when compared to lopinavir–ritonavir, led to a significantly shorter median time to convert nasopharyngeal swab negative for viral RNA, reduced duration of hospital stay, and a better clinical improvement with only mild side effects [83]. Data from other ongoing trials (NCT04392427, NCT04460443) would prove helpful in further establishing if it provides more benefits on early administration upon presentation with pneumonia than when the situation worsens.

Darunavir and cobicistat

Darunavir is an HIV-1 protease inhibitor which is used in combination with cobicistat as a pharmaco-enhancer. Docking results found it to have an affinity to SARS-CoV-2 Nsp3c, PLpro, E-channel, or Spike proteins, but the affinity for the main protease (3CL protease) was of most interest [55]. Closer visualisation showed that darunavir had very few interactions with the important residue of 3CL protease and was found to have no in vitro activity against SARS-CoV-2 [84, 85]. However, it still underwent a clinical trial on 30 COVID-19-positive patients at Shanghai Public Health Clinical Centre, and results showed that it was indeed ineffective [85]. Another clinical trial NCT04304053, dropped darunavir owing to in vitro data and other findings.

Famotidine/famotidine and hydroxychloroquine (HCQ)

Famotidine is a histamine-2 receptor antagonist and is commonly used for gastric acid suppression. It has shown in vitro antiviral properties against HIV replication [86]. Docking studies suggest that famotidine is likely to inhibit 3CLpro [55], which is essential for proteolytic processing of the SARS-CoV-2 polypeptide, and hence for viral replication. In a retrospective cohort study with large sample size, it was found that the use of famotidine in COVID-19 patients who were not in an intensive care setting, initially led to a twofold reduction in clinical deterioration. These patients were at reduced risk for intubation or death [87]. Further, a combination of dual histamine blocker (H1 antagonist Cetirizine, and H2 antagonist Famotidine) was used in a proof-of-principle, non-randomised, un-controlled trials and was found to be effective in a good prognosis and reduced ventilator support [88]. A randomised clinical trial (NCT04370262) is now being carried out to assess the efficacy of famotidine in combination with HCQ for the treatment of COVID-19.
**Hydroxychloroquine/HCQ and azithromycin**

HCQ is an antimalarial drug first approved by the FDA in 1955. It is a derivative of chloroquine (CQ) but has less toxicity. Apart from its use for the treatment of uncomplicated malaria, it is also prescribed for rheumatoid arthritis (RA) and systemic lupus erythematosus [89]. In the current crisis, it remained the most controversial molecule, and various studies reported it to be effective, while others found it ineffective and still others reported adverse effects of HCQ in COVID-19-affected individuals. Studies demonstrated that HCQ and CQ have a potent in vitro activity against SARS-CoV-2 [53], which is attributed to an increased endosomal pH, and its anti-inflammatory and immunomodulating effects [90]. On March 28, 2020, the FDA gave emergency use approval for the use of chloroquine phosphate and hydroxychloroquine sulphate to treat adult COVID-19 patients but later revoked it on June 15, 2020 considering its “known and potential risk” [91].

**Studies in favour:** An open-label, non-randomised trial was conducted in 36 patients in which 26 patients received hydroxychloroquine. 6 patients in the hydroxychloroquine group also received azithromycin. Despite a small sample size, Gautret and co-workers found that after 6 days of treatment, 100% of patients who received azithromycin and hydroxychloroquine together were negative for SARS-CoV-2; against 57.1% of those who received only hydroxychloroquine and 12.5% in the control group. Authors thus reported hydroxychloroquine as a potent drug to clear COVID-19 and found even better results when used synergistically with azithromycin [92]. In another randomised clinical trial in 62 COVID-19 patients, Chen and co-workers reported that 31 patients assigned to receive hydroxychloroquine (HCQ) showed shortened time to clinical recovery (TTCR) and improved absorption of pneumonia. Significantly reduced duration of high body temperature, recovery time, and cough remission time were reported in the HCQ group [93].

**Studies against:** This was contradicted when Molina and co-workers did not find any strong clinical evidence when in a prospective study, 11 severe COVID-19 patients with significant comorbidities were administered HCQ (600 mg/d for 10 days) and azithromycin (500 mg on day 1 and 250 mg on days 2–5) in the same dosage as used by Gautret and co-workers [92, 94]. Another uncontrolled double-blinded, randomised trial in 81 patients conducted to address different treatment dosage of chloroquine in severe COVID-19 patients red-flagged the use of high dosage (12 g of CQ in total, for 10 days) due to potential safety hazard [95]. No significant benefit of HCQ treatment was found by Tang and co-workers in an open-label, randomised, controlled trial with 75 patients in the HCQ group. In mild to moderate COVID-19 patients, the negative conversion probability in both the groups remains approximately similar. However, there were more adverse effects in the HCQ recipient group [96]. Even for COVID-19 patients with hypoxic pneumonia, there was no reduction in ICU admission or death after 7 days of hospital admission when treated with HCQ [97]. Thus, the efficacy of HCQ with or without azithromycin remains questionable and demands more randomised controlled trials. On May 25, 2020, WHO temporarily paused the trial of HCQ within its solidarity trial, owing to safety concerns [98] but later resumed it on the recommendation of the data safety and monitoring committee on June 03, 2020, only to again discontinue it on July 04, 2020 [64, 99].

**Tocilizumab**

Cytokine storm mediated by IL-6 plays a significant role in COVID-19 patients with severe symptoms (discussed above). Tocilizumab, an anti-human IL-6 receptor monoclonal antibody, works by binding to soluble and membrane-bound IL-6 receptors (sIL-6R and mIL-6R, respectively) preventing IL-6 binding and, thus, inhibits signal transduction pathway. It was found to be well tolerated and is currently approved for the treatment of severe rheumatoid arthritis and other inflammatory and autoimmune conditions [100, 101].

With randomised trials (NCT04310228, NCT04317092, NCT04339712, NCT04330638, NCT02735707, NCT04372186 [https://clinicaltrials.gov] still underway, there have been several retrospective studies. In one such study, it was reported that tocilizumab treatment in 21 severe and critical COVID-19 patients in China resulted in improved clinical presentations with no significant adverse effects. This study suggested that early treatment could also prevent deterioration of symptoms [102]. A different retrospective study at the Italian center found that bacterial superinfection in ICU settings could complicate treatment with tocilizumab [103]. 30 COVID-19 patients in France with severe and rapidly deteriorating pneumonia were treated with tocilizumab. When compared with the control group, Roumier and co-workers reported a significant reduction in mechanical ventilation requirements and reduced risk of ICU admission [104]. Similar improved survival was reported from a retrospective case-control study at a medical center in New York [105]. Further, reports from first randomised, placebo-controlled trial (COVACTA trial) of tocilizumab found no benefits in the death rates in the intervention arm. This study did not meet its primary endpoint (clinical status on a 7-category ordinal scale: 1 for discharge/readied to be discharged; to 7 for death) and the clinical presentations were not improved significantly as compared to control. However, reduced duration of hospital stay (median 8 days shorter than placebo) and ICU stay (median 5.8 days shorter than placebo)
were demonstrated in patients treated with tocilizumab [106]. These results contradict the earlier reports from various retrospective trials and indicate no effect of tocilizumab in managing mortality in COVID-19 patients. Results from further trials are awaited to support or refute the existing evidences.

### Ivermectin and doxycycline

Ivermectin is an FDA-approved broad-spectrum antiparasitic medication used against worm infestations. It has shown to inhibit several viruses, including HIV-1, Dengue, among others in vitro [107, 108]. This broad-spectrum activity of ivermectin is attributed to targeting of host importin (IMPa/B1) transport protein, which is used by many viruses during infection [109].

Caly and co-workers recently reported in vitro activity of ivermectin against SARS-CoV-2 with ~5000-fold reduction in the viral RNA load in 48 h at IC50 of ~2 μM [110]. However, it remains to be seen if such high plasma concentration could be safely attained, provided that it targets the host protein [111]. A bioinformatic study suggests that ivermectin could potentially bind to 3CL protease and the HR2 domain (spike protein) of SARS-CoV-2 synergistically [112]. Thus, a randomised control trial could shed more light on the efficacy of ivermectin in treating COVID-19, and multiple such studies are currently underway (https://clinicaltrials.gov/). However, Chaccour and coworkers pointed out that concurrent use with boosted antivirals like lopinavir/ritonavir and darunavir/ritonavir/cobicistat could increase systemic concentration (leading to neurotoxicity) of ivermectin [113]. A multicenter, retrospective cohort study at four hospitals in South Florida showed ivermectin to have significant effect in lowering mortality rate even in severe COVID-19 cases. However, no difference in duration of hospital stay or rate of extubation was observed in intervention and control arms [114]. In a pilot interventional trial in mild to moderate patients, a single dosage of ivermectin used as an add-on therapy over HCQ and azithromycin (ivermectin group) showed better efficacy when compared to synthetic controls (patients already treated and later used as an external control) receiving hydroxychloroquine and azithromycin alone [115]. Meanwhile, a medical team from Bangladesh claimed to have treated all 60 patients with a combination of ivermectin and doxycycline within 4 days without any adverse effect [116]. Retrospective and pilot trials are often bound by puzzling factors, albeit unmeasured at times. Larger placebo-controlled randomised trials are required to derive better evidence on effectiveness of ivermectin. However, such placebo-controlled trials tend to be unethical at times of pandemic.

### Dexamethasone

Corticosteroids have been reported in several retrospective studies to prevent the worsening of symptoms and reduce the risk of intubation [117–119]. Treatment with corticosteroid was associated with faster alleviation of high body temperature and improvement of SpO2 when compared with the group which did not receive them [120]. However, some other studies found increased mortality risks on the administration of corticosteroids with more cases of multiple organ dysfunctions [121, 122]. As described earlier, the deleterious effect of high levels of pro-inflammatory cytokines leads to extensive mortality in severe cases. Corticosteroid is an immunosuppressive drug that winds down the immune response and also helps to reduce inflammation. This anti-inflammatory response could thus prevent further alveolar damage [123]. However, the use of steroids was not advised by WHO for treatment of COVID-19 cases due to adverse effects and delayed rate of viral clearance as experienced in previous outbreaks during SARS-CoV, MERS-CoV, and H1N1 epidemics [124].

One such inexpensive steroid, dexamethasone is the first drug to have shown to reduce mortality in severe COVID-19 cases. In one of the arms of RECOVERY (Randomised Evaluation of COVID-19 Therapy) trial in the UK, dexamethasone was found to prevent one-third of the deaths of patients requiring mechanical ventilation. 2104 enrolled patients were administered a dose of 6 mg per day of the drug for 10 days while 4321 patients in the control group received only the standard care. It was preliminarily reported that dexamethasone does improve the chances of survivability in severe COVID-19 cases. The patients on oxygen therapy also reported 20 percent reduced mortality; however, there were no effects on less severe cases (those not requiring oxygen support). Following this, it was immediately authorised by the UK government for use in patients requiring oxygen support, including those who need mechanical ventilation [125–127].

Another FDA-approved glucocorticoid methylprednisolone has shown to benefit moderate to severe COVID-19 patients in a multicentric, partially randomised trial. This study reflects on the benefits of a short term course of methylprednisolone to decrease the risk of admission to ICU or need for ventilation support and posed no major side effect. However, hyperglycemia was frequently observed [128]. Another multicentre observational study concluded that early treatment of severe cases with methylprednisolone could prevent disease progression [129]. Results from several ongoing randomised trials (NCT04263402, NCT04345445, NCT04329650) could shed more light on its efficacy in the treatment of COVID-19 cases. In a meta-analysis by the ‘WHO Rapid Evidence Appraisal for COVID-19 Therapies (REACT) Working Group’, results
from 7 randomised clinical trials covering 1703 critical COVID-19 patients (678 to corticosteroid against 1025 in control group) from 5 continents suggested lowered all-cause-mortality after 28 days post randomisation. Overall 32% mortality was documented in corticosteroid intervention group against 40% in placebo. The analysis further recommended corticosteroid intervention along with standard care for critically ill COVID-19 patients, ‘in the absence of compelling contraindications’ [130].

**Convalescent plasma**

Convalescent plasma from recovered patient has been used historically to treat diseases in the events of lack of prophylactic or therapeutic interventions [131]. This mode of intervention is under trial for the treatment of COVID-19 patients. In a Cochrane review consisting a total of 5443 participants in 20 studies (16 non-controlled non-randomised studies of interventions (NRSIs), 3 controlled NRSIs, and 1 Randomised control trial), no certain evidence to substantiate the efficacy of convalescent plasma in reducing mortality, or improving clinical symptoms, or prolonging time-to-death was found [132]. Similarly, a randomised control trial in Netherlands did not conclude improvement in mortality and severity in the intervention arm. This study was however prematurely halted on recommendation of data safety monitoring board [133]. A more recent randomised control trial was reported from India, wherein 464 participants underwent randomisation (235 in intervention arm against 239 in control arm) at 39 trial sites. Authors found no significant difference in mortality or disease progression among moderate COVID-19 patients. They, however, reported improved resolution of indications (like fatigue and shortness of breath), and faster negative conversion of viral RNA in patients in the intervention arm but failed to demonstrate any anti-inflammatory property [134]. Undergoing clinical studies (clinicaltrials.gov: NCT04333251; NCT04344535; NCT04345523; NCT04332835; NCT04342182; NCT04345991; NCT04345289; NCT04346446) on convalescent plasma therapy could add to existing knowledge when completed.

Furthermore, other immunotherapies using monoclonal antibodies, cytokines, interferon, and mesenchymal stem cell therapy are also under clinical trials in an effort to modulate immune response in COVID-19 patients (reviewed by Mansourabadi and coworkers [135]).

**Under-trial molecules**

With such a large number of infected patients, the pandemic warrants the immediate need for treatment. Multiple trials are being conducted across the world to test several therapeutic options in an effort to generate strong evidences and build a uniform treatment protocol. Table 3 enlists some interventions (investigational and approved) that are under trial against COVID-19 infection with their status from [https://clinicaltrials.gov/](https://clinicaltrials.gov/) and Chinese Clinical Trial Register ([http://www.chictr.org.cn/](http://www.chictr.org.cn/)); and primary usage information from drugbank ([https://go.drugbank.com/](https://go.drugbank.com/)).

**Molecules with favourable in vitro/docking results but not under clinical trial yet**

Table 4 enlists some drugs with favourable docking for SARS-CoV-2 targets but which are not currently under clinical trial. Anti-HIV drugs like nelfinavir have shown affinity for 3CLpro [136, 137]. Ganciclovir was found to potentially bind RdRp, exonuclease, and helicase [136]. In vitro activity against SARS-CoV-2 was also exhibited by miglustat which is expected to work at the host cell level. In the postinfection process, it targets folding and glycosylation of viral protein in the endoplasmic reticulum. Miglustat is an approved drug to treat some genetic disorders and the in vitro activity was observed at a safe and achievable plasma concentration level [138–140]. Targeting multiple proteins simultaneously could reduce the chances of developing resistance. Elbasvir has been shown to target key SARS-CoV-2 proteins like RdRp, PLpro, helicase, and also the viral spike protein [141]. N4-deoxycytidine, a ribonucleoside analogue with broad antiviral properties, and its prodrug EIDD-2801, have demonstrated in vitro efficacy in Vero cells against SARS-CoV-2 at an IC50 of 0.3 µM. It has been shown to be potent even against coronaviruses with resistance for other nucleoside analogue drugs like remdesivir [142].

**Conclusion**

This inimitable illness has affected the entire world’s population, either directly or indirectly. With hundreds of thousands of deaths already recorded due to COVID-19, there are still no signs of infection trail slowing down. Many low-income countries faced an added challenge of preventing deaths due to starvation and, hence, made efforts to open the economies. With aerosol as one of the primary modes of transmission among others [143], proper face masks, eye protection along with physical distancing measures could significantly reduce the chance of infection and prevent the healthcare system from being overwhelmed [144]. With no specific antivirals available, patients have primarily been provided symptomatic treatment, and the effort has been broadly aimed at containing the spread of infection.

Remdesivir, an antiviral that was initially being developed for Ebola infection, gained approval by the US FDA for
emergency use after showing impressive results in a US clinical trial [59, 60, 145]. In contrast, earlier few trials showed little or no clinical benefit. Treatment with Lopinavir/Ritonavir had many adverse gastrointestinal events with no clear benefits. However, it did reduce the patient’s stay in the ICU [62]. This intervention regime has been dropped from both SOLIDARITY trial (WHO) and RECOVERY trial after failing to show clinical benefit [64, 65]. Chen and co-workers reported that patients who did not respond to treatment with lopinavir/ritonavir reported negative PCR tests after treatment with danoprevir/ritonavir [76]. Approved for the treatment of influenza infection, both arbidol and favipiravir indicated clinical recovery. Favipiravir was also suggested to prevent the progression of the disease to ARDS, which is one of the primary reasons for death in severe cases [70, 71]. There is a ‘cytokine storm’ in the latter phase of the

Table 3 Antivirals under trial against COVID-19 infection with no results reported yet

| Sl. no | Drug | Primary use | Trial id | Status |
|--------|------|-------------|---------|--------|
| 1      | CLEVUDINE | Investigational for use in HBV infection | NCT04347915 | RCT, recruiting, Phase 2 |
| 2      | ASC-09 | Investigational for use in HIV-1 infection | NCT04261270 | RCT, Recruiting, Phase 3 |
| 3      | GALIDESIVIR | Investigational for use in Ebola infection; broad-spectrum antiviral | NCT03891420 | RCT, Recruiting, Phase 1 |
| 4      | AZVUDINE | Investigational for use in HIV-1 infection | 1. ChiCTR2000030487  
2. ChiCTR2000030424  
3. ChiCTR2000029853 | 1. Non-randomised; recruitment status unknown  
2. Non-randomised; recruitment pending  
3. RCT, recruitment status unknown |
| 5      | BALOXAVIR MARBOXIL | Antiviral against Influenza | ChiCTR2000029548 | RCT, Not yet recruiting |
| 6      | EMTRICITABINE/ TENOFOVIR METENKEFALIN + TRIDECAC-TIDE | Antiviral against HIV | ChiCTR2000029468 | Non-randomised; Not yet recruiting |
| 7      | FINGOLIMOD | Treatment of multiple sclerosis | NCT04280588 | Non-randomised; Recruiting, Phase 2 |
| 8      | IBUPROFEN | NSAID used as an analgesic, anti-inflammatory and antipyretic | 1. NCT04382768  
2. NCT04334629 | 1. Compassionate Use Program; Recruiting  
2. RCT, Recruiting, Phase 4 |
| 9      | EIDD-2801 | Investigational for use in influenza, MERS-CoV and SARS-CoV-2 infection | NCT04405570  
NCT04405739 | 1. RCT, Recruiting, Phase 2  
2. RCT, Recruiting, Phase 2 |
| 10     | ATAZANAVIR | Treatment of HIV infection | 1. NCT04459286  
2. NCT04468087  
3. NCT04452565 | 1. RCT, Not yet Recruiting, Phase 2  
2. RCT, Not yet Recruiting, Phase 2/3  
3. RCT, Recruiting, Phase 2/3 |
| 11     | NAFAMOSTAT | Used as anti-coagulant in patients requiring renal replacement therapy | 1. NCT04352400  
2. NCT04473053 | 1. RCT, Not yet Recruiting, Phase 2  
2. RCT, Recruiting, Phase 2 |
| 12     | OPAGANIB | Investigational, Sphingosine kinase-2 (SK2) selective inhibitor | 1. NCT04414618  
2. NCT04467840 | 1. RCT, Recruiting, Phase 2  
2. RCT, Recruiting, Phase 2 |
| 13     | TELMISARTAN | Antihypertensive | 1. NCT04360551  
2. NCT04355936  
3. NCT04359953 | 1. RCT, Recruiting, Phase 2  
2. RCT, Recruiting, Phase 4  
3. RCT, Recruiting, Phase 3 |
| 14     | DIPYRIDAMOLE | Used as anticoagulant | 1. NCT04424901  
2. NCT04391179  
3. NCT04410328 | 1. RCT, Recruiting, Phase 2  
2. RCT, Recruiting, Phase 2  
3. RCT, Not yet Recruiting, Phase 3 |
| 15     | AT-001 | Investigational, aldose reductase inhibitor | NCT04365699 | Non-randomised controlled trial, Recruiting, Phase 2 |

[Status: ClinicalTrials.gov (https://clinicaltrials.gov/) and Chinese Clinical Trial Register (http://www.chictr.org.cn/); Primary usage information: Drugbank (https://go.drugbank.com/)]

NSAID Non-steroidal anti-inflammatory drug, RCT randomised controlled trial
disease, which may lead to ARDS. Tocilizumab, an IL-6 receptor inhibitor, was initially suggested to wind down this storm and prevent deterioration of symptoms [104, 105, 146]. However, tocilizumab failed to provide any mortality benefit when compared to control [106]. With increasing safety concerns, the role of HCQ largely remains unclear in tackling COVID-19 infection. The US FDA recently revoked the emergency use approval for HCQ over safety concerns [91]. Further, convalescent plasma therapy that initially looked promising failed to impart any significant benefit in reducing mortality or curtailing disease progression. It however, resulted in faster negative conversion and improved symptom resolution when compared to controls [134].

So far ivermectin and doxycycline appeared to be an effective therapeutic combination [116] supported by data from a retrospective study which found ivermectin to significantly lower mortality rate [114]. In a first result of its kind, steroid dexamethasone was reported from a RECOVERY (Randomised Evaluation of COVID-19 Therapy) trial in the UK to reduce mortality in severe COVID-19 cases [125, 126]. Being an RNA virus and under antiviral pressure, SARS-CoV-2 is prone to mutations. Combination therapy is thus desirable, which could address multiple targets of the virus and reduce the chance of developing drug resistance [76] that could potentially be used on a case-to-case basis subject to validation of drug interaction. Availability of a convenient animal model a “Syrian hamster” [147] may prove to be appropriate for studying the pathogenesis of SARS-CoV-2, and evaluating immunotherapies and antiviral drugs. It will overcome the shortcomings of the previously available animal models namely hACE2 transgenic mice and macaques (primate model) to a great extent [148].

There is a clear need to channel information from various trials to delineate a standard treatment protocol and guiding the future clinical trials. With hundreds of trial still in the pipeline and many already reporting conclusions, a compiled review serves as a quick source for reliable information. Till date, dexamethasone have shown proven efficacy in reducing mortality and remdesivir in improving clinical recovery, while many like hydroxychloroquine and convalescent plasma failed to meet expected efficacy. Given the severity of the disease, compassionate use of these drugs along with the symptomatic treatment and standard supportive care, is now being prescribed by the professionals. As it seems quite plausible that COVID-19 is here to stay for long, the efforts made towards finding an answer are still looking insufficient and require further research into greater depths [149]. Even if an effective solution is found, sooner or later emergence of resistance due to the highly mutable nature of the virus is inevitable. This warrants researchers to continue to race against time to formulate a better concoction and a vaccine to the SARS-CoV-2.

Again, we recommend against self-medication and reiterate the importance of seeking proper medical attention in line with local guidelines.

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#### Conflict of interest

The authors declare that they have no conflict of interest.

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#### Availability of data and material

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#### Consent to participate and publication

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