Role of ACE2 receptor and the landscape of treatment options from convalescent plasma therapy to the drug repurposing in COVID-19

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Abstract

Since the first case reports in Wuhan, China, the SARS-CoV-2 has caused a pandemic and took lives of > 8,35,000 people globally. This single-stranded RNA virus uses Angiotensin-converting enzyme 2 (ACE2) as a receptor for entry into the host cell. Overexpression of ACE2 is mainly observed in hypertensive, diabetic and heart patients that make them prone to SARS-CoV-2 infection. Mitigations strategies were opted globally by the governments to minimize transmission of SARS-CoV-2 via the implementation of social distancing norms, wearing the facemasks, and spreading awareness using digital platforms. The lack of an approved drug treatment regimen, and non-availability of a vaccine, collectively posed a challenge for mankind to fight against the SARS-CoV-2 pandemic. In this scenario, repurposing of existing drugs and old treatment options like convalescent plasma therapy can be one of the potential alternatives to treat the disease. The drug repurposing provides a selection of drugs based on the scientific rationale and with a shorter cycle of clinical trials, while plasma isolated from COVID-19 recovered patients can be a good source of neutralizing antibody to provide passive immunity. In this review, we provide in-depth analysis on these two approaches currently opted all around the world to treat COVID-19 patients. For this, we used “Boolean Operators” such as AND, OR & NOT to search relevant research articles/reviews from the PUBMED for the repurposed drugs and the convalescent plasma in the COVID-19 treatment. The repurposed drugs like Chloroquine and Hydroxychloroquine, Tenofovir, Remdesivir, Ribavirin, Darunavir, Oseltamivir, Arbidol (Umifenovir), Favipiravir, Anakinra, and Baricitinib are already being used in clinical trials to treat the COVID-19 patients. These drugs have been approved for a different indication and belong to a diverse category such as anti-malarial/anti-parasitic, anti-retroviral/anti-viral, anti-cancer, or against rheumatoid arthritis. Although, the vaccine would be an ideal option for providing active immunity against the SARS-CoV-2, but considering the current situation, drug repurposing and convalescent plasma therapy and repurposed drugs are the most viable option against SARS-CoV-2.

Keywords Cytokine storm syndrome · ARDS · Rheumatoid arthritis · Herd immunity · Vertical transmission · Vaccine nationalism

COVID-19 and SARS-CoV-2 virus

An infectious disease COVID-19 caused by a virus belongs to the Coronaviridae family was first reported in December 2019 in the Wuhan city of China. Several other lethal viruses such as Severe Acute Respiratory Syndrome-related Coronavirus (SARS-CoV) and Middle Eastern Respiratory Syndrome coronavirus (MERS-CoV) also belong to this family. The SARS-CoV is a single-stranded, enveloped positive-sense strand RNA virus with a genome size between 27 and 34 kilobases that is comparatively larger than other RNA viruses. SARS-CoV-2 driven endemic unfurled into
a pandemic on 11th of March 2020 by the World Health Organization (WHO). So far, a total of seven human coronaviruses (hCoVs) types have been identified as shown in Fig. 1. The newest coronavirus strain that caused the current pandemic is known as severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). The latest member SARS-CoV-2 has similarity close to 70% with SARS novel coronavirus. The SARS-CoV-2 infection is being characterized by severe clinical manifestations of the respiratory tract with a highly complex pathogenesis. Among targets of the virus are the epithelial cells of respiratory tract, which upon infection result to diffuse alveolar damage and severe lung injury. After entry into the cells, the virus propagates in the cytoplasm, which is also the site for formation and budding of the virus containing vesicles. The destruction of cells occurs upon release of vesicles [1].

**Structure of SARS-CoV-2 virus**

SARS-CoV-2 RNA codes for four types of significant proteins: specific spike (S), membrane (M), nucleocapsid (N), and envelope (E) [2]. The detailed structure of the virus has been shown in the Fig. 2. The S protein is a transmembrane glycoprotein, which facilitates the virus entry into the host cell by using the signal sequence of N-terminal to have access to the endoplasmic reticulum [3]. Due to its gigantic size, it creates distinct spikes on the viral surface. The N protein helps in viral RNA synthesis, while E and M proteins are instrumental in viral assembly.

**Mechanism of entry into the host cells, RAAS, and replication of SARS-CoV-2**

The S protein consists of two subunits: S1 & S2 by a protease Transmembrane Serine Protease 2 (TMPRSS2). TMPRSS2 is a furin-like protease that contains a single transmembrane domain and single domain for SR TRYPsin and LDLA domains as well [4]. The gene encoding for this protein is located on 21q22.2. The primary localization of TMPRSS2 is restricted to the plasma membrane. TMPRSS2 is a secretory protein as its presence has been reported in biological fluids like semen [5] and urine [6]. The protein architecture of TMPRSS2 is shown in Fig. 3.

The priming of the S protein for pathogenicity is carried out by TMPRSS2 in coronaviruses like SARS-CoV, and MERS-CoV [7]. S protein consists of two subunits: S1 and S2. Among these, S1 subunit contains the receptor-binding

![Fig. 1 Classification of RNA-based viruses and flow-chart showing the belongingness of Coronavirus and other closely related RNA viruses. This schematic classification of the Coronaviridae family shows how the members are divided based on sense and anti-sense strands. SARS-CoV-2 falls in category of single stranded sense strand RNA virus that is enveloped and possesses helical capsid. The α-coronaviruses are: 229E and NL63. Except SARS-CoV-2, there are other members of the β-coronavirus types are: OC43, HKU1, SARS-CoV, and MERS-CoV](image-url)
domain (RBD), which binds to the SARS-CoV-2 viral receptor angiotensin-converting enzyme II (ACE2). ACE2 is a carboxypeptidase that contains one transmembrane domain and also one signal peptide [4, 8]. The gene encoding for this protein is located on Xp22. The primary localization of ACE2 is membranous and the secondary localization is extracellular. ACE2 is a secretory protein as its presence has been reported in biological fluids like plasma [5], and urine [6]. The protein architecture of ACE2 protein has been shown in Fig. 3. Further, mRNA expression levels of ACE2 in normal tissues are quite heterogeneous [9] as shown in RNAseq derived data in Fig. 4. The descending order of mRNA expression level was as follows: small intestine (93.7 ± 16.1), duodenum (69 ± 6.29), gall bladder (32.6 ± 14.37), kidney (30.8 ± 17.14), testis (26.9 ± 8.99), heart (12.3 ± 10.95), thyroid (1.39 ± 0.928), liver (1.29 ± 0.38), stomach (1.18 ± 0.937), and lung (0.345 ± 0.3).

SARS-CoV-2 exploits S protein for binding to its receptors ACE2 or DPP4 (dipeptidyl peptidase 4, in bronchial epithelial cells) [10]. The mechanisms of entry and replication of SARS-CoV-2 have been shown in Fig. 5. The gene encoding for this protein is located on 2q24.3. The primary localization of CD26 is a plasma membrane and the secondary localization is extracellular. CD26 is a secretory protein as its presence has been reported in biological fluids like plasma [11], serum [12], semen [5], tears [13], and urine [14]. The protein architecture of CD26 protein has been shown in Fig. 3.

The N protein, which is phosphorylated, binds to SARS-CoV-2 genome are like a bead on a string fashion. The E protein is a transmembrane protein found in lower concentration and play an important role in assembly & releasing of the virus, and therefore crucial for pathogenesis. The M protein is a dimer and most abundant one among M, N, S and E protein.

Hemagglutinin-esterase (HE) exists as a dimer protein present in some beta coronaviruses. It binds to sialic acids on the surface of glycoproteins, and increase S protein-mediated viral entry into the cells, and eventually the virus spread through the mucosa. Unlike other β-coronaviruses, SARS-CoV-2 infection occurs not only in the mucosal epithelium (nasal depression and pharynx) of the upper respiratory tract.
but also in other organs such as of the gastric tract. Other cells types that are infected during the pathogenesis may include the neurons in brain, tubular epithelial cells of the kidneys, and intestinal mucosa cells. There have been reports of infection in sites that may lead to heart injury, failure of organs such as liver, intestine, and kidney [15].

SARS-CoV-2, cytokine storm syndrome, and organ failure

One major issue in COVID-19 cases is the blood upregulation of pro-inflammatory cytokines such as IL-1, IL6, TNF, and interferon γ. The major source of cytokine production are macrophages, as upon activation they can produce cytokines like TNF-a, interleukins including IL6, IL1, IL4, IL13, and IL18. Those further activate the cascade reaction of inflammatory factors that eventually lead to the cytokine storm syndrome (CSS), an uncontrolled response of cytokines. In CSS, an increased and uncontrolled secretion of pro-inflammatory cytokines give rises to acute respiratory distress syndrome (ARDS). It is characterized with progressive arterial hypoxemia, and breathing difficulties [16]. Respiratory failure due to ARDS is a major cause of death in COVID-19 patients [17]. CSS has been reported not only in avian H5N1 influenza virus, SARS and Middle East Respiratory Syndrome (MERS), but also in other diseases like multiple sclerosis and pancreatitis. Role of different cytokines in relation to COVID-19 has been well documented [18]. Dust cells, which are present in the alveolar region of lungs, are macrophages that play an important role in CSS. Type I IFN low levels are common to COVID-19, MERS, and SARS which could suppress Th1, but favor Th2 responses [19].

Transmission of SARS-CoV-2

Transmission of the virus can happen even from a person who shows no symptoms for COVID-19 (asymptomatic) [20]. The COVID-19 patients starts developing symptoms such as mild respiratory issues, and fever with an incubation period between 5 and 6 days that can get extended up to 1–14 days [21]. Mode of COVID-19 transmission can be through different routes including contact, saliva, droplet, faecal and aerosol transmission [22]. Possibility of vertical transmission of COVID-19 has been also suspected, where the virus can be transmitted from parents to offspring’s via placental barrier, transcytosis of the cell-associated virus, during delivery, or through breast-feeding, but vertical transmission in case of COVID-19 was not reported until recently [23]. The first case of vertical
transmission of SARS-CoV-2 in India was reported from Sassoon General Hospital, Pune, Maharashtra (India) [24].

SARS-CoV-2 transmission can happen by touching contaminated surfaces followed by nose, eyes, or mouth. To stop the transmission of SARS-CoV-2 a number of Do’s and Don’ts have been recommended by the WHO as well as by agencies like NIH and ICMR. The lists of Do’s and don’ts required to mitigate COVID-19 transmission have been mentioned in the Table 1.

**COVID-19 and herd immunity**

When a higher percentage of the community becomes immune to a disease (that could be due to prior illness or vaccination) and makes spreading of the disease improbable is known as *herd immunity*. Even non-vaccinated (such as newborns and the immunocompromised one) but susceptible individuals offer immune-protection because the disease has hardly any possibility to spread within the
| S. no. | Parameter                          | Don’ts | Do’s | Mechanism                                                                                           | References |
|-------|-----------------------------------|--------|-----|-----------------------------------------------------------------------------------------------------|------------|
| 1     | Smoking                           | ✔      |     | Smoking have increased expression of ACE2 receptor in the host cells infected by the SARS-CoV-2 virus | Brake et al. [25] |
| 2     | PPE (medical professional)        | ✔      |     | Personal protection equipment kit is very essential for health care personnel to wear specially those dealing with nCoV-19 patients/suspects | Giwa et al. [26] |
| 3     | Face mask                         | ✔      |     | SARS-CoV-2 spreads through droplets or aerosol so a proper mask is required to stop spreading transmission of SARS-CoV-2 virus. N95 masks have penetration size from 0.1 to 0.3 micron. Facemasks prevent spreading of droplets coming in cough and sneezing | Tirupathi et al. [27], Ozma et al. [28] |
|       |                                   |        |     |                                                                                                      | [https://www.who.int/emergencies/diseases/novel-coronavirus-2019/technical-guidance/infection-prevention-and-control](https://www.who.int/emergencies/diseases/novel-coronavirus-2019/technical-guidance/infection-prevention-and-control) |
| 4     | Hand sanitization                 | ✔      |     | The cleaning and washing hands with alcohol or with soap and water must be done or 20–30 s. Methanol, isopropyl alcohol and ethanol are major disinfectant agents. Ethanol concentration between 60 and 95% (v/v) is safe and effective for disinfection | Berardi et al. [29] |
|       |                                   |        |     |                                                                                                      | [https://www.who.int/emergencies/diseases/novel-coronavirus-2019/advice-for-public](https://www.who.int/emergencies/diseases/novel-coronavirus-2019/advice-for-public) |
| 5     | Social/physical distancing        | ✔      |     | Maintain a social distance of at least 2 m or ~6 feet is required to avoid infection                   | MacIntyre et al. [30] |
| 6     | Visit of Crowded places           | ✔      |     | Gathering is a good source to spread or getting infection with SARS-CoV-2 virus because at crowded places it is hard to maintain a social or physical distance of 2 m (6 feet). Social distancing is crucial in preventing community transmission | Wilder-Smith et al. [31] |
| 7     | Touching eyes, nose, and mouth    | ✔      |     | The virus transmission can be through infected person’s airways/droplets (Aerosol), nose or mouth to recipient’s eyes, mouth or nose. Touching surface with bare hands can caught viral infection | West et al. [32] |
| 8     | Updating of information           | ✔      |     | It is of utmost importance to keep up to date on the latest information from trusted sources, such as WHO or your local and national health authorities as they provides the most updated information/advice that need to follow in the area they are residing during the pandemic | Santos et al. [33] |
| 9     | Alcoholic sanitizers to be out of reach | ✔      |     | It is poisonous and must be out of children’s reach of children as they have more probability of accidental swallowing and ingestion. Reported health effects after swallowing are drowsiness, eye irritation, nausea, vomiting, etc. Methanol is very toxic and may prove life threatening sometimes. Alcohol vapor is also harmful | Santos et al. [33] |
| 10    | Sharing of cup, utensils, food, or drink items | ✔      |     | It transmits infection                                                                                     | Müller et al. [34] |
| S. no. | Parameter | Don’ts | Do’s | Mechanism | References |
|-------|-----------|--------|------|-----------|------------|
| 11    | Stay home | ✔      |      | The measures by which transmission can be reduced are; rules on distance and hygiene to prohibitions on meetings and extension of school and university holidays to the closure of all non-system-relevant facilities in a country in connection with the regulation of individual freedom of movement i.e. lockdown | Adhikari et al. [35] |
| 12    | Namaste   | ✔      |      | This way of greeting avoids physical touch as observed in hand shaking | Kulkarni et al. [36] |
| 13    | Hugs, handshake, high five, fist bump, & | ✔      |      | If individuals will go for these types of greetings or mode of celebration, transmission of virus will happen | Adhikari et al. [35] |
| 14    | COVID-19 associated app provided by the government | ✔      |      | It’s always good to upload COVID-19 app on your mobile to know latest update as well as COVID-19 positive patient in the vicinity | Ming et al. [37, 38] |
| 15    | Make a note of helpline number of emergency medical facility | ✔      |      | In case if you submit the symptoms such as fever, difficulty in breathing and cough. This put stats accordingly if you are at high or low risk of getting the disease, and further in tracing and monitoring of the patient | Collado-Borrell et al. [38] |
community [39]. This is associated with the R0 (R Zero or R naught or basic reproductive number), which represents the infectivity of an infectious agent like SARS-CoV-2. The R0 value has been estimated in different studies on the SARS-CoV-2 virus ranging from 2 to 6. Between the two cohorts, the R0 observed was 2.2 [40], and 5.7, respectively [41]. In a recent study, with an R0 value of 3 for SARS-CoV-2, the threshold for herd immunity was ~67% which means that the decline in the incidence of SARS-CoV-2 infection will begin in the population when it surpasses 0.67 [42]. To develop herd immunity against SARS-CoV-2 there are two ways i.e. first, we vaccinate at a massive scale, but it’s not possible without the availability of a safe and efficacious vaccine. The second option is via natural immunization of the world population with the virus but has seriously implication, as a large proportion of the population must be infected with SARS-CoV-2.

Here, we present a systematic review cum meta-analysis conducted to evaluate the significance of currently used treatment options for COVID-19, associated issues, and future challenges in dealing with infection and management of SARS-CoV-2. To achieve this goal, we carried out this study to evaluate the repurposing drug agents so far used for the treatment of COVID-19. For this, we used “Boolean Operators” search criteria in PUBMED to get relevant search outcome [43]. The schema for fetching the data and further filtering of the articles has been shown in Fig. 6.

We used keywords such as:

I. COVID-19 OR coronavirus = 48,139
II. COVID-19 AND repurposing drugs = 234
III. COVID-19 AND repurposing drugs = 232
IV. COVID-19 AND repurposing drugs = 02

We have searched the literature and screened published research articles to further dig-down to list which molecule the repurposed drug targets and their route of administration whether oral, cutaneous, subcutaneous or in the form of injection, through which those have been given to the patients. Next, we corroborated the additional information by visiting the https://clinicaltrials.gov/ to get additional information on the clinical trials where the repurposed drugs have been used. The protein architecture of some of the important proteins such as ACE2, TMPRSS2, and DPP4 was extracted from the human protein reference database (HPRD) freely accessible at https://hprd.org [4]. Further, the structures of the repurposed drugs were drawn using ChemDraw Professional Version 16.0 software. The rest of the figures were made using Adobe Illustrator CS5 version 15.0.0.

There had been many treatment options adopted worldwide to treat COVID-19 patients. Among those: convalescent plasma therapy, and repurposing the drugs are taking the lead in the absence of a vaccine or unavailability of a neutralizing antibody for coronavirus.

**Convalescent plasma as a potential therapy for COVID-19**

The plasma derived from COVID-19 patients those successfully overcome its infection is referred to as *convalescent plasma* (CP). CP had been used in the past for treatment of deadly viral diseases such as Severe Acute Respiratory Syndrome (SARS), H1N1, Spanish flu, Ebola, and the MERS.

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**Fig. 6** Schema for screening of the articles reporting drugs repurposed for COVID-19 The NCBI search engine was searched using Boolean operators such as AND, NOT, & OR. The articles were fetched for repurposing drugs, synergism or convalescent plasma in combination with COVID-19. The articles were further segregated based on the agent used for drug repurposing.
The German scientist Emil von Behring got the noble prize in 1901 for the usage of CP in the treatment of diphtheria. The CP provides neutralizing antibodies to the patient against infectious agents [39]. A must to do task is to get a measurement of titer of the neutralizing antibody in advance prior to giving the plasma to the COVID-19 patients. A titer of > 1:320 must be there for the neutralizing antibody [40].

The patients who recovered from COVID-19 can be identified as potential donors if they have: (i) prior diagnosis of COVID-19, (ii) complete resolution of symptoms at least 14 days prior to donation, (iii) a negative RT-PCR result for COVID-19, and (iv) desired SARS-CoV-2 neutralizing antibody titers (optimally > 1:320). Although, the donor titer varies in current CP based trials from > 1:40 (NCT04374487 from India) to > 1:320 (USA:NCT04377672, NCT04373460, NCT04344535, Hungary:NCT04345679), the higher the better. There are more specifics coming up on the criteria for exclusion and inclusion for a donor as well as for a recipient in CP therapy. These are listed based on different clinical trials from NIH, USA as well as trials from other countries in Supplementary Table 1. Though only a handful of studies are there on the usage of CP therapy on COVID-19 patients, those have been summarized in Table 2. The exclusion and inclusion criteria for donors and acceptor have been mentioned in Supplementary Table 1.

In a small study on COVID-19 patients in Guangdong (China), after the 12th day of hospitalization of patients with severe condition CP was given. Three out of four patients discharged, and the last one was found negative using RT-PCR, two out of four patients were able to produce anti-SARS-CoV-2 IgG ~ 14 days post-transfusion [41]. A high titer antibody present in the recovered COVID-19 patients must be sufficient enough to bind SARS-CoV-2 and neutralize it to avoid access to normal cells. One of the major challenges is that CP is not used alone but in combination with other agents like corticosteroids. The neutralizing antibodies present in the CP are capable to accelerate the clearance of infected cells as well. CP constituents are capable of activating the effector mechanisms such as complement activation and phagocytosis [49]. A combination of CP and corticosteroids can reduce the viral load as well as reduce the excess of inflammatory response [47]. The initial findings from all around the world are encouraging from CP therapy supporting the evidence that the human anti-SARS-CoV-2 plasma could be able to modulate the virulence exerted by the SARS-CoV-2 via neutralization [50].

**Drug repurposing as an alternative therapy for COVID-19**

Drug repurposing or drug repositioning (which is sometimes also defined as drug re-profiling or drug re-tasking) is an approach for exploring the new maneuver of already approved drugs, which have been used for the treatment of other diseases [51]. In contrast, synergism is an interaction between two or more drugs that leads to overall effect to be cumulatively more than the sum of individual effect. Drug synergism is measured by calculating the combination index (CI) using freely available software CompuSyn [52]. The CI value > 1, = 1 and < 1 represents antagonistic, additive, and synergistic interaction between two or more drugs [53]. Based on the literature survey, we divided the repurposed drugs that can be used for the trials to treat COVID-19 into

| Table 2 | Titer ratio among different studies where Convalescent plasma has been used as a treatment option for COVID-19 patients |
| --- | --- | --- | --- | --- |
| Region/country | Titer | Patient received CP | Patient outcome | References |
| Dongguan, Xiangtan, Xiaolan cities of China | > 1:320 | 06 | The patients treated with CP did not require mechanical ventilation and 11 days post-CP treatment were transferred to a general ward | Zhang et al. [41] |
| Shenzhen, China | Antibody against anti-SARS-CoV-2 AB > 1:1000, and neutralization titer > 40 | 05 | Anti-SARS-CoV-2 antibody titers ranged between 1:800 and 16:200, NAb titers from 80 to 480, reduced the viral load | Shen et al. [44] |
| Wuhan, China | Neutralizing Anti-SARS-CoV-2 AB > ~1:640 | 10 | Reduced the viral load | Duan et al. [45] |
| Wuhan, China | Not mentioned | 06 | An instant accretion in titer of anti-SARS-CoV-2 AB titers in patients #2 and #3 | Ye et al. [46] |
| South Korea | Not mentioned | 02 | Convalescent plasma therapy was given to two COVID-19 patients. Both showed a favorable outcome | Ahn et al. [47] |
| Italy | ≥ 1:160 | 46 | Primary outcome was 7-days hospital mortality and 6.5% patients died within 7 days | Perotti et al. [48] |
five categories: (I) Anti-malarial drugs (II) Drugs used for Rheumatoid arthritis (III) Cytokine modulators (IV) Protease Inhibitors (V) Others. Additionally, the details of the currently going on clinical trials are summarized in Supplementary Table 2.

The drug repurposing offers benefits in terms of time and costs required as compared to the development of a new drug from the beginning. The repurposed drugs are approved by the Food and Drug Administration (FDA), for their pharmacological properties, safety, and clinical efficacy for a different indication [54]. Therefore when used for the COVID-19 treatments, the toxicity or safety profiles of the repurposed drugs are already known. Therefore, a number of drugs have been proposed for the repurposing to treat the COVID-19 patients across the world. We are presenting basic properties of these drugs, their target and study outcome or at least observations reported in studies globally. We have summarized the information on repurposed drugs, their target, and diseases information for which those were made in Table 3.

**Anti-malarial/anti/protozoan drugs**

The structures of anti-malarial or anti-parasitic drugs have been shown in Fig. 7.

**Chloroquine phosphate**

Primarily chloroquine phosphate had been used for the treatment of malaria. It is a quinolone that possesses anti-inflammatory properties, and some time for amoebiasis as well. It is also known as chloroquine (CQ). CQ offers an advantage, as it does not pose complications associated with infectious complications exerted by drugs like methotrexate and leflu-nomide. Evidence-based on different studies showed that CQ possesses broad-spectrum anti-viral activities [80, 81]. Both anti-viral as well as anti-inflammatory activities of CQ possibly responsible for CQ’s efficacy in treating the pneumonia of COVID-19 patients [82].

**Hydroxychloroquine (HCQ)**

HCQ has other names/synonyms such as Oxychlorochin, Plaqueril, and Oxichloroquine. An in vitro activity against anti-SARS-CoV of HCQ was found to be superior as compared with CQ [83], and the HCQ clinical profile is also superior to CQ [84]. In terms of side effects when compared CQ with HCQ, it was found that CQ treated patients have some side effects like circular defects or Bull’s eye maculopathy, retinopathy, diametric retina defects and cardiomyopathy, but patient treated with HCQ have reduced tissue accumulation that could be responsible for lesser adverse events of HCQ as compared with CQ. A high dose for > 5 years of HCQ led to retinopathy development, which is in concordance with the HCQ as a therapy [85, 86].

**Emetine**

Emetine is an alkaloid isolated from the flowering plant *Carapichea ipecacuanha* a member of the family Rubiaceae. Emetine has been used against protozoan infections and also to induce vomiting. Emetine is a translation-inhibiting drug that had been used in amoebiasis treatment. It is capable of inhibiting translation machinery of the malaria parasite (*Plasmodium falciparum*) by binding to the E ribosomal site. This shows anti-viral activity against a wide range of viruses (both DNA and RNA based) including Zika, rabies, cytomegalovirus, Ebola, and HIV-1 virus. Emetine also showed anti-viral activity against hCoV-OC43, SARS-CoV, hCoV-NL43, MHV-A59, and MERS-CoV in an in vitro condition. The viral polymerase enzyme and some host proteins are the targets of Emetine [18]. It has been recently reported that emetine inhibits the replication of SARS-CoV-2 at ~ 0.5 µM concentration. The in vivo achievable concentration of emetine in plasma is 0.075 µg/mL (0.156 µM), lower than the in vitro EC₅₀ against SARS-CoV-2 [19].

**Anti-viral drugs**

These drugs work against different viruses including retro-viruses like HIV-1 and have been proposed to use for the treatment of COVID-19. The structures of selected anti-retroviral/anti-viral drugs have been shown in Fig. 7.

**Favipiravir**

Favipiravir is an oral anti-viral drug used for the treatment of influenza. It came into limelight for Ebola treatment during the 2014 epidemic in West Africa as there was no standard of care (SOC) was available. Effectiveness of favipiravir was also observed for prophylaxis and infectious animal models of lethal Ebola virus [87]. It is a purine analogue and also known as 6-fluoro-3-hydroxy-2-pyrazinecarboxamide or T-705 or Avigan as a brand name, which targets viral RdRp (RNA dependent RNA Polymerase). On an urgent basis, favipiravir had been approved for the clinical trial in adult COVID-19 patient’s treatment (2020L00005). SARS-CoV-2 also possess RdRp gene similar to other members of the family (SARS-CoV and MERS-CoV), which makes favipiravir eligible to be tested against SARS-CoV-2 virus. It is a pro-drug which upon ribosylation and phosphorylation form an active metabolite intracellularly called T-705RTP or favipiravir ibofuranosyl-5′-triphosphate (T-705RTP), which interfere with the replication of the virus by competing with the naturally occurring purine nucleosides and inhibits the viral RdRp of SARS-CoV-2.
| Name of the agent (mode of giving to the patients) | Type | Originally used for the diseases | Target & mode/mechanism of action | References |
|--------------------------------------------------|------|---------------------------------|----------------------------------|------------|
| Baricitinib (oral)                               | Active ingredient of Olumiant | Rheumatoid arthritis             | Reversible JAK inhibitor         | Kuriya et al. [55] |
| Favipiravir (an oral anti-viral drug)             | Analogue of purine origin    | Influenza                        | RdRp catalytic site preventing    | Furata et al. [56] |
|                                                  |                                               |                                 | virus replication, It is erroneously |            |
|                                                  |                                               |                                 | concede as purine nucleotide by   |            |
|                                                  |                                               |                                 | the RdRp                          |            |
| EIDD-2801 (an oral anti-viral drug)               | Ribonucleotide analog, it is isopropyl ester prodrug of [N4-hydroxy-cytidine] | Tested in influenza, MERS-CoV, SARSCoV-2 | Same enzyme targeted by remdesivir. | Sheahan et al. [57] |
| Oseltamivir (orally administered drug)            | Sialidase inhibitor               | Influenza A and B               | Neuraminidase                      | Oliveira et al. [58] |
| Remdesivir (intravenously)                        | An adenosine analogue             | Flu virus (influenza)           | Targets viral RNA polymerase       | Eastman et al. [59] |
|                                                  |                                               |                                 | via incorporation of adenosine     |            |
|                                                  |                                               |                                 | analogue in the nascent RNA chain  |            |
|                                                  |                                               |                                 | using viral RdRp                   |            |
| Methylprednisolone (oral)                         | A corticosteroid                  | Diminish pro-inflammatory        | Binds to nuclear receptor         | Lu et al. [60] |
| Tofacitinib (oral)                                | An-acylpiperidines                | Psoriatic arthritis and rheumatoid arthritis | Selective JAK1 & JAK3 inhibitor, inhibits CYP3A4 | Emori et al. [61], Guo et al. [62] |
| Ruxolitinib (oral)                                | Apyrrolo [2,3-d]pyrimidines       | High-risk myelofibrosis         | Inhibitor of JAK1 & JAK2           | Elli et al. [63] |
| Chloroquine                                       | Anaminoquinolone derivative       | Malaria, HIV, Q fever, Whipple’s disease | Inhibits caspase-1 (CASP1), NLRP3. | Oscanoa et al. [64] |
|                                                  |                                               |                                 | CQ escalates the pH of the        |            |
|                                                  |                                               |                                 | endosomes as well as impede the    |            |
|                                                  |                                               |                                 | glycosylation of some receptors   |            |
| Hydroxychloroquine (oral)                         | 4-Aminoquinoline                  | Malaria, Rheumatoid arthritis and Lupus | ALDH1 and QR2. The mechanism of action is very similar to CQ | Graves et al. [65] |
| Oseltamivir (oral)                                | Anti-viral neuraminidase inhibitor  | Influenza A & B                  | Neuraminidases, Ion channels of nicotinic acetylcholine receptors | O’Hanlon et al. [66], Ono et al. [67] |
| Lopinavir (oral)                                  | Anti-retroviral protease inhibitor  | HIV                              | Mimetic Protease that impede viral protease activity | De Clercq [68], Paskas et al. [69] |
| Ruxolitinib (oral)                                | Cancer growth blocker             | Myelofibrosis                    | Janus Kinase inhibitor (JAK inhibitor) with selectivity for subtypes JAK1 and JAK2 | Mesa [70] |
| Carolacton                                        | Macrolide keto-carboxilic acid (A natural product Bio-film inhibitor Isolated from Sorangiumcellulosum) | Antibacterial                      | FoID/MTHFD                         | Fu et al. [71] |
| Ivermectin (oral and tropical)                   | Macrocyclic lactones              | Parasite infestations (Head lice, Scabies, Onchocerciasis, Strongyloidesis, trichuriasis, ascariasis and lymphatic filariasis) | Activates glutamate—gated Cl- channels Cys -loop receptor, P2X4 receptor, fernesoid X receptors | Chen et al. [72] |
In Shenzhen (China), a clinical trial of favipiravir on COVID-19 patients was conducted for evaluation of safety and efficacy (ChiCTR2000029600). A total of 35 patients in the favipiravir arm showed significantly a shorter viral clearance duration in contrast with the control arm containing 45 patients. Further, these findings were corroborated with chest X-rays showing improvement in the favipiravir arm (91.43% vs 62%) [88]. In another multi-centric randomized study (ChiCTR200030254), favipiravir treatment of COVID-19 patients led to an improved recovery at 7th day from 55.86 to 71.43% [89].

**Remdesivir**

Remdesivir (also known as GS-5734) is a 1’-cyano-substituted adenosine analogue. It is a pro-drug that inhibits viral RNA polymerases, has shown in vitro activity against coronaviruses like SARS-CoV-2, CoV-229E, SARS-CoV, CoV-OC43, and MERS-CoV [90]. It is a mono phosphoramide pro-drug possessing wide anti-viral spectrum covering filoviruses, coronaviruses, pneumoviruses, and paramyxoviruses. It has been observed that remdesivir inhibits humans as well as animal coronaviruses in vitro, including SARS-CoV-2. Remdesivir proved to be a superior drug in a lethal murine MERS model as compared with a regimen of IFN-b, and lopinavir-ritonavir combination. An EC50 of remdesivir was 0.77 μM against SARS-CoV-2 virus [91]. It has been documented mutations such as F476L and V553L in the nsp12 polymerase gene of murine hepatitis virus confer remdesivir resistance [92].

**Alovudine**

Alovudine (also known as fluorothymidine) a DNA polymerase inhibitor developed by Medivir is an anti-viral agent. Due to toxicity issues, in 2005 after phase II clinical trial, it was discontinued. Alovudine is a nucleoside reverse transcriptase inhibitor analog of thymidine [93]. Alovudine is able to terminate the RNA synthesis SARS-CoV-2 virus, but more work is required before it makes an entry into a clinical trial.

**Drug used for rheumatoid arthritis**

The structures of drugs used for rheumatoid arthritis, but now repurposed for treating the COVID-19 patients are shown in Fig. 7.

**Baricitinib**

Baricitinib is an orally available agent used for rheumatoid arthritis. It is sold with the brand name Olumiant. It inhibits the response of inflammatory molecules and
cytokine production via modulation of JAK-STAT pathway [94]. It is an active ingredient of olumiant. Baricitinib is a reversible inhibitor of JAK1/JAK2. According to the EU Clinical Trials Register there are already phase-II, and III (2020–001854-23), and phase-IV (2020–001354-22) clinical trials using Baricitinib on COVID-19 patients. Modulation of cytokine dysregulation could affect the host inflammatory response and entry of viruses into the cells.
This makes it an ideal agent to be tested in COVID-19 patients [95, 96].

**Tofacitinib (oral)**

Baricitinib and tofacitinib are first-generation JAK inhibitors. Tofacitinib is a small molecule inhibitor of Janus Kinases particularly JAK1/JAK3 [97]. It is sold with the brand name Xeljanz. It had been used for the treatment of RA (moderate to severe form). Tofacitinib is N-acylpiperidine compound. Tofacitinib inhibits STAT also, but in a reversible manner. Tofacitinib subjects to hepatic metabolism through cytochrome CYP3A4 mainly which means a combination of CYP3A4 inhibitors could be tested first in vitro to see if there is a synergistic impact [98].

**Ruxolitinib (oral)**

It has been used for the treatment of moderate to high-risk myelofibrosis. It is sold in the market with the trade name Jakafi or Jakavi. Ruxolitinib is an oral kinase inhibitor that inhibits JAK1 and JAK2. It is also known as INC801842. Metabolism of ruxolitinib is facilitated by CYP3A4. The chemical constituent of Ruxolitinib belongs to the pyrrolo [2, 3-d] pyrimidines class of organic compound.

**Cytokine modulators**

**Tocilizumab**

An anti-IL6 blocker targets the IL6 receptor proved to be effective in rheumatoid arthritis treatment [99], and later for juvenile idiopathic arthritis [100], giant cell arteritis [101]. IL6 is a bonafide marker for inflammation. It is also known by another famous name Actemra and recently approved by the FDA for testing in a clinical trial for COVID-19 patients. It is a recombinant antibody humanized and of IgG1 class. Actemra is capable of disrupting inflammatory response exerted by IL6 is known as cytokine release syndrome (CRS). The efficacy of Actemra was tested on COVID-19 patients at The First Affiliated Hospital of the University of Science and Technology, China. Among 21 patients tested, the body temperature returns to normal in all the cases. An improvement in respiratory function was seen in 100% of the patients and the recovery rate was ~ 95% as seen in CT scan reports, and the patients were discharged within 14 days of post-tocilizumab treatment. The findings extrapolated on 500 severe or critical patients enrolled in a clinical trial (ChiCTR2000029765) [102]. In contrast, the Italian guidelines suggest that tocilizumab use is suitable in patients with major symptoms including when high viral load is over, and patients don’t have any fever (Apyretic) for > 72 h or 7 days post-onset of symptoms, and increased IL6 levels [94].

**Anakinra**

Anakinra is a recombinant human antagonist of IL1R that has been used in rheumatoid arthritis. These drugs also play an important role in the management of CRS. Due to the release of IL1R SARS-CoV-2 causes an advanced form of cell death occurs due to inflammation (pyroptosis) and mediated by caspase-1. The repurposed drug anakinra in case of COVID-19 patients in phase III randomized clinical trial able to reduce both requirement of invasive mechanical ventilation in ICU as well as the mortality rate in severe COVID-19 cases without serious side-effects [79].

**Adalimumab (anti-TNF-α agent)**

It has been earlier used for treatment of Rheumatoid arthritis. The mode of adalimumab for patients is subcutaneous. FDA approved it long back in 2002 for treatment of RH. Biosimilar of adalimumab (Hyrimoz) is also available which is available by the name adalimumab-adaz approved in Oct 2018 by the FDA. Adalimumab binds to TNFα and leads to inhibition of interaction with the receptor of TNF by binding with p55 & p75. There is a trial going on (ChiCTR2000030089) where one arm includes conventional treatment along with adalimumab [103]. An interesting observation has been mentioned regarding levels TNF-α that it was moderately high in SARS but significantly higher levels in COVID-19 patients.

**Protease inhibitors anti-retroviral/anti-viral drugs**

**Lopinavir**

Lopinavir is an anti-retroviral drug used for the treatment of HIV patients. It is a protease inhibitor, which has been used for the treatment of SARS-CoV infected patients in combination with ritonavir & ribavirin in a non-randomized clinical trial. Only few SARS patients progressed to ARDS with this treatment as compared with patients receiving only ribavirin and corticosteroids. It was sold by the brand name Kaletra. It is interesting to note that Lopinavir is exclusively given along with ritonavir because lopinavir possesses poor oral bioavailability and extensive biotransformation. On the other hand, Ritonavir is an inhibitor of the enzymes related to lopinavir metabolism, therefore a co-administration boosts the lopinavir exposure and significantly improves the anti-viral activity [104]. Finding of a randomized control trial (ChiCTR2000029308) on SARS-CoV-2 showed no significant benefit of lopinavir-ritonavir combination in SARS-CoV-2 patients as compared with SOC [105]. Combining lopinavir with other agents to treat SARS-CoV-2 virus not only increased synergism but also decreased the lopinavir inhibitory concentration.
Ritonavir

It is another anti-retroviral drug used against HIV. It’s a protease inhibitor that inhibits the productive cycle of the HIV virus. It is sold with the trade name Norvir. Ritonavir inhibits HIV-1 protease as well as host’s cytochrome P450 3A4 enzyme that helps in metabolizing lopinavir. The NIH panel recommended not using the combination of lopinavir/ritonavir or other HIV protease inhibitors due to unfavorable outcome post-treatment of these agents.

Ribavirin

Ribavirin or tribavirin is an anti-viral drug used for Rous sarcoma virus infection, viral hemorrhagic fevers and hepatitis C. Ribavirin is synthetic guanosine nucleoside that interferes with the viral mRNA synthesis. Ribavirin has been used in combination with interferon beta-1b, lopinavir-ritonavir in a randomized phase-II clinical trial in COVID-19 patients and the early results showed that it was superior to alone lopinavir-ritonavir combination in reducing the symptoms exerted by the virus and shortening shedding of the virus [106].

Camostat mesilate

Camostat mesilate (CM) is an inhibitor of TMPRSS2. It inhibits the serine proteases like TMPRSS2 [7]. CM proved to be effective in blocking the spreading and virulence of SARS-CoV in a lethal mouse model [107]. It was observed that CM is capable of blocking the entry of SARS-CoV-2 into the lung cells. Camostat inhibits diverse range of proteases including plasmin, trypsin, kallikrein and thrombin [108]. University of Tokyo, Japan planned to conduct a clinical trial on the combination of CM and nafamostat on COVID-19 patients. CM was approved in Japan for treatment of pancreatic inflammation [109].

Homoharringtonine

Homoharringtonine is also known as omacetaxine meptesuccinate or HHT. HHT is a cephalotaxine ester. It was isolated from the leave of Cephalotaxus fortunei of family Taxaceae. Omacetaxine received orphan drug status from FDA in March 2006 (according to FDA an orphan drug is the one which is intended to treat a rare disease which affect <200,000 persons in the United States) for treatment of chronic myeloid leukemia patients particularly those who found to be resistant to > 2 tyrosine kinase inhibitors. HHT shows anti-cancer activity via inhibition of translation by binding to ribosomal site-A. This forces the cells to lose proteins like MCL1 and c-MYC (both with short half-life), crucial for leukemia cell’s survival. HHT showed activity against a large number of viruses including pseudorabies virus, rabies virus, hepatitis B virus, Newcastle disease virus, and echovirus 1 [110]. In an in vitro screening in Vero E6 cells, HHT inhibited SARS-CoV-2 replication at an EC₅₀ of < 100 μM [111].

An urgent requirement and challenges for more treatment options for COVID-19

Availability of safe and efficacious vaccine against COVID-19

Multiple pharmaceutical companies and academic institutions are joining the hands for the collaborative efforts to develop a vaccine against SARS-CoV-2. There are multiple candidates proposed for the potential vaccine against SARS-CoV-2 that include mRNA vaccine, inactivated virus vaccine, DNA vaccine recombinant protein vaccine, and viral vector-based vaccine. Globally, the experts in vaccine development think that it will take around 18 months to develop a SARS-CoV-2 vaccine, although it is very optimistic even if we consider this to be the fastest created new vaccine in the history. In traditional settings, it takes 5 to 6 years to develop a vaccine, but the high mutation rate of RNA viruses and therefore changing the specific immune response make it even harder to develop an efficacious vaccine. Every vaccine in human clinical trials goes through three phases: Phase-I is the safety trial in a small group of healthy volunteers, where a vaccine is tried out with different dosages to find out the strongest immune response without serious side effects. The phase-II vaccine trials test how well the vaccine works in hundreds of people of diverse age and health status. Next, in phase three, the vaccine is given to thousands of people who are already at the risk of infection, and then wait to see if the vaccine reduces the number of people getting sick. Since phase-III is tried out in natural disease condition and larger population size, it is usually the longest phase. A number of potential candidates for vaccines are in pre-clinical studies and a few in clinical trials in different parts of the world including the USA, Russia, China, UK, and India. Regardless which of these countries get success and finally a vaccine gets approved, the first challenge is whether the country is willing to share it with the rest of the world, and a proper storage and distribution system. So, overcoming these challenges requires close collaboration between pharma giants, regulatory bodies like FDA, active involvement and cooperation of the scientific community, and healthcare systems.

Development of antibodies neutralizing the virus

Most of the anti-SARS-CoV novel antibodies (nAbs) have been targeted against S protein, RBD [112], S2 subunit, and S1/S2 proteolytic cleavage sites. Some nAbs like S230.15,
found to be negative for increased, but TNF-α level was decreased. The MSCs disappeared. The Dendritic cell population and IL10 were increased, and on day 6 cytokine secreting cells (CXCR3+ cells) were increased. CT scans showed that the pneumonia of clinical trials are time consuming processes and require target to testing in the animals and then different phases frame may vary because from the selection of the suitable vaccine to deliver the vaccine in next 12 to 18 months but the time clinic nationalism is against the principles of global public health, unfortunately, there is no law to prohibit the pre-purchase in pandemic like COVID-19.

Conclusion

The existence of SARS-CoV-2 was reported in 2019. Since then it posed a threat to mankind around the world. Hastening of treatment options for COVID-19 brought nothing so far but we have to look at old treatment option as a savior because we know convalescent plasma therapy and repurposing drugs approaches had been used in the past in crisis period. While massive-scale efforts to make a suitable vaccine are on the way, time being number of drugs used for other diseases have been currently repurposed to tackle the COVID-19 pandemic.

Globally, the experts in vaccine development are optimistic to deliver the vaccine in next 12 to 18 months but the time frame may vary because from the selection of the suitable target to testing in the animals and then different phases of clinical trials are time consuming processes and require

An unobvious challenge of vaccine nationalism

During COVID-19 pandemic one issue emerged where different countries particularly the USA, Russia, and China are in the race of making a vaccine against SARS-CoV-2. At the same time, countries like USA and Russia are securing a large and sufficient number of vaccine doses against COVID-19 for their own people and prioritizing their own market rather than making available to other countries. This is popularly known as ‘vaccine nationalism’. This can be executed through pre-purchase agreements between a vaccine manufacturer and the government. WHO issued a warning regarding vaccine nationalism as instead of helping mankind, it’s going to help the virus. It’s not a new challenge, as a similar scenario was observed during The H1N1 flu pandemic in 2009. At that time, among vaccine producers for H1N1 flu Australia was the leader and the government blocked the exports, but at the same time, the rich countries went for the pre-purchase agreements with some pharma giants. In the case of COVID-19, US government already has shown interest to secure 600,000 doses. Though vaccine nationalism is against the principles of global public health, unfortunately, there is no law to prohibit the pre-purchase agreements with some pharma giants. In the case of COVID-19, US government already has shown interest to secure 600,000 doses. Though vaccine nationalism is against the principles of global public health, unfortunately, there is no law to prohibit the pre-purchase agreements with some pharma giants. In the case of COVID-19, US government already has shown interest to secure 600,000 doses. Though vaccine nationalism is against the principles of global public health, unfortunately, there is no law to prohibit the pre-purchase agreements with some pharma giants. In the case of COVID-19, US government already has shown interest to secure 600,000 doses.

Mesenchymal stem cell therapy for COVID-19

Stem cell therapy proved to be very useful in treating a number of diseases including cancer [117], and diabetes [118]. Mesenchymal stem cells (MSC) are characterized by low invasive nature and high proliferation rate, and additionally devoid of ethical & social issues that makes it as the preferred therapeutic option over others [119]. MSCs play an important role in immunomodulatory effects via secreting many types of cytokines by paracrine secretion or make direct interactions with immune cells. The source of MSCs can be peripheral blood (PB), bone marrow (BM), adipose tissues [(AT), buccal fat pad, abdominal fat, & infrapatellar fat pad], placenta, umbilical cord, Warton jelly, amniotic fluid, and blood cord. Therefore, it seems MSCs-based therapy may possibly be an ideal candidate for clinical trials or at least the combination of treatment to treat COVID-19 patients.

MSC therapy was applied in COVID-19 patients on seven patients. The levels of peripheral lymphocyte were increased, and on day 6 cytokine secreting cells (CXCR3+ CD4+ T, CXCR3+ CD8+ T, and NK CXCR3+ cells) were disappeared. The Dendritic cell population and IL10 were increased, but TNF-α level was decreased. The MSCs were found to be negative for ACE2 and TMPRSS2 suggesting there was no SARS-CoV-2 infection [120].

In addition, recently a case study was reported in China on a female patient with an acute COVID-19 syndrome that the results of laboratory tests and CT images provided extremely effective results after 21 days of treatment with umbilical cord MSCs. Upon treatment with MSCs, an increase in lymphocyte and a decrease both in WBCs and neutrophils was observed. Some T cell surface markers like CD3, CD4, and CD8 were increased. CT scans showed that the pneumonia was cleared [121]. Including the ground-glass opacity in the lung, the other typical diagnosis characteristic of critically ill patients was a significant decrease in lymphocytes along with the increase of neutrophils. However, only a handful studies on MSCs show promising results in the treatment of COVID-19 patients. Though only on a limited number of patients, but these studies showed that MSC therapy alone or in combination with other drugs can be used to treat SARS-CoV-2 infected patients.

m396, S109.8 and S227.14 showed neutralizing activity against human, raccoon dog, and palm civet but none of these have been evaluated in clinical studies. Antibodies like 80R (scFv or mAb) is capable of neutralizing the infection of SARS-CoV via blockage of RBD-ACE2 interaction [113]. There is a high sequence similarity for S protein between SARS-CoV-2 and SARS-CoV [114]. This suggests a cross neutralizing/cross-reactivity of nAbs between SARS-CoV and SARS-CoV-2 infection. The SARS-CoV mAb CR3022 (RBD-specific) could possibly bind to RBD of SARS-CoV-2 because RBD of SARS-CoV & SARS-CoV-2 are very similar [115]. Different fragments like S1-NTD, S2, and RBD have been used as a target for the development of nAbs. A similar strategy could be adopted for SARS-CoV-2. CP is currently in use for the treatment of COVID-19 patients, but non-nAbs targeting other regions than RBD of S-protein can create antibody-dependent enhancement (ADE) effect on virulence as well as on the disease [116].

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quality controls. Though, FDA has put vaccine development and their approval on the fast track, but there are no short cuts for different stages of developments and quality controls directly associated with the safety and efficacy. There are already concerns raised by different scientists including Dr Anthony Fauchi about the COVID-19 vaccine from Russia. With all the hopes on the potential vaccine’s progress, the strategies are also required to build the infrastructure for equitable distribution of the vaccine and to avoid bottleneck on the availability as soon as the vaccine successfully completes the clinical trials. While we wait for a vaccine to come into the picture, convalescent plasma therapy, and Repurposing the drugs treatment options proved to be suitable (if not perfect). We need to have suitable vaccine development, neutralizing nABs antibody as prophylactic and therapeutic, and mesenchymal stem cell-based treatment options for effectively dealing with COVID-19. Until an ideal treatment comes, people must follow proper caution such as wearing masks, follow social distancing, and as much as possible do activities, which could be afforded through online mode/ route.

Author contributions MKK and AH conceived and guided the research. MKK, AKS, PK, AK, GT, & AT carried out the exhaustive search on the data. MKK, AH, AKS, PK, AK, GT, AT, KM, and PCM analyzed and interpreted the data. MKK wrote the manuscripts. All the authors read, critically evaluated, gave their feedback, and edited the manuscript.

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Data availability All data generated and analyzed during our study are included in the published article and also supplied in two supplementary tables.

Compliance with ethical standards

Conflict of interest Authors declare no potential conflicts of interest and also no competing financial interest associated with this manuscript.

Ethical approval The data included in the study was collected form

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