nCOVID-19: Its diagnosis, possible preventive measures, therapeutic interventions and management

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Abstract

Novel coronavirus (nCOVID-19) is a current global threat causing severe infections to millions of people throughout the world. WHO has reported in their case-based surveillance that the aged and male populations are most vulnerable to this recent coronavirus infection. This disease-causing virus, named as SARS-CoV-2, is a respiratory coronavirus that probably originated from the bat with a substantially different genetic makeup concerning the other known coronaviruses. Structurally SARS-CoV-2 is a diverse single-stranded RNA virus having spike (S) glycoprotein in its envelop that is associated with the transmission of the virus by binding with the host target receptor. The principal target site of nCOVID-19 is angiotensin-converting enzyme 2 (ACE2) in the host receptor. The expression of ACE2 target site varies according to the genetic susceptibility and ABO blood group locus causing variation in the severity of the infection. The virus usually affects the inflammatory pathways, responsible for white blood cell activation and clot formation after interacting with pathogen pattern recognition receptors. The major pathological symptoms of severe nCOVID-19 are mild to severe respiratory and bowel syndromes. Prevention of infections has been achieved by social distancing depending on the concept of aerosolized/droplet transmission of virus and by maintaining personal hygiene. Remedies like ayurvedic, homeopathic, micronutrients such as vitamins and minerals along with regular physical exercises like yoga and meditation are also found to be helpful in disease prevention. The treatments that have been applied so far with some positive responses are antimalarial chloroquine and hydroxychloroquine, antivirals, plasma therapy, steroids, omega-3-fatty acid derivatives, vitamin C infusion, etc. Some global healthcare bodies are also in action for the development of vaccines therapy against nCOVID-19. Apart from the infection control, some other associated issues like long term social distancing, constant stress, anxiety, change in lifestyle are affecting adversely the aged populations and also the other part of the society. There is no doubt that government bodies from various countries throughout the world are taking extreme measures to control and mitigate the pandemic, but this untoward situation is still far from the control of human races and supposes to take more time and further scientific interventions and management are needed to be continued.

Abbreviation

ACE2: Angiotensin-Converting Enzyme 2; ADCC: Antibody-Dependent Cellular Cytotoxicity; ARDS: Acute Respiratory Distress Syndrome; B cells: Bursa cells; CD4+ T cells: Cluster of Differentiation 4+ Thymus cells; CD8+ T cells: Cluster of Differentiation 8+ Thymus Cells; CDC: Complement Dependent Cytotoxicity; CFT: Complement Fixation Test; CI: Confidence Interval; CNS: Central Nervous System; CoV HKU4: Tylonycteris bat coronavirus HKU4; CRS: Cytokine Release Syndrome; cryo-EM: cryo-Electron Microscopy; CSF: Cerebro Spinal Fluid; CT: Computerized Tomography; DFA: Direct Fluorescent Assay; DIC: Disseminated Intravascular Coagulation; E: Envelope protein; EC50: half maximal Effective Concentration; EIA: Enzyme-Immunonassay; ELISA: Enzyme-Linked Immunosorbent Assay; FP: Fusion Peptide; GS-5734: Remdesivir; GSK: GlaxoSmithKline; HCoV-229E: Human Coronavirus 229E; HCoV-HKU1: Human Coronavirus HKU1; HCoV-NL63: Human Coronavirus NL63; HCoVOC43: Human Coronavirus OC43; HI: Hemagglutination Inhibition; HR: Heptad Repeat 1; HR2: Heptad Repeat 2; HyFc: Hybridoma Fusion; IBD: Inflammatory Bowel Diseases; IC: Intracellular domain; IFA: Immuno-Fluorescence Assay; IFN-γ: Interferon gamma; IgD: Immunoglobulin D; IgG4: Immunoglobulin G4; IL-6: Interleukin-6; IL-7: Interleukin-7; INO-4800: DNA vaccine for SARS-COV-2; IQR: Interquartile Range; kb: kilo byte; LF: Loop Mediated Isothermal Amplification; LAT: Lateral-Flow Test; LFIA:
Lateral-Flow Immune chromatographic assay; mAbs: monoclonal antibodies; MERS-CoV: Middle East Respiratory Syndrome Coronavirus; mRNA-1273: Lipid Nanoparticle (LNP)-encapsulated mRNA-based vaccine; N: Nucleocapsid protein; nCOVID-19: novel Corona Virus Disease-2019; NIAID: National Institute of Allergy and Infectious Diseases; NK cells: Natural Killer; NKG2A: Natural Killer cell G2A; nM: nano Molar; np14: nonstructural protein 14; NTD: N-Terminal Domain; Orf1a,b: Open reading frame 1a and b 226; PCR: Polymerase Chain Reaction; PD: Peptidase Domain; RA: Rheumatoid Arthritis; RBD: Receptor-Binding Domain; RBM: Receptor-Binding Motif; RdRP: RNA-dependent RNA Polymerase; RNA: Ribonucleic Acid; rRT-PCR: Real-Time reverse transcription-Polymerase Chain Reaction; RSV: Respiratory Syncytial Virus; RTLAMP: Reverse Transcription Loop-Mediated Isothermal Amplification; RT-PCR: Reverse Transcription Polymerase Chain Reaction; SARS-CoV-2: Severe Acute Respiratory Syndrome Coronavirus 2; SD1: Subdomain 1; SD2: Subdomain 2; SLE: Systemic Lupus Erythematosus; T-705: Favipiravir; T-cell: Thymus cell; TM: Transmembrane region; TMPRSS2: Transmembrane Protease Serine 2; TNF-alpha: Tumor Necrosis Factor-alpha; WHO: World Health Organization

Introduction

The novel coronavirus, nCOVID-19, also named as severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), is a zoonotic virus causing mild respiratory tract infections to unusual fatal pneumonia in humans upon exposure [1]. In the recent past, the deadly infection due to nCOVID-19 has been first identified in Wuhan City, China on 29th December 2019 and rapidly takes a shape of a global pandemic by increasing exponentially during the last 7 months [2]. WHO has reported (as mentioned in the report released on 28th May 2020) that there are more than 10.1 million nCOVID-19 positive cases as the development of nCOVID-19 infections globally with almost 0.5 million deaths [3]. WHO has also claimed, based on their case-based surveillance that the occurrence of infection is age and gender-based (Figure 1). As per their report, the ratio of the rate of infection in males and females is 1.03:1 with a median age of 51 years (i.e. interquartile range, IQR: 36–65 years) [4]. The most common symptoms of the nCOVID-19 positive patients are mild to moderate in nature like nasal secretions, cough, fever, dyspnea, myalgia, and occasionally diarrhea [5]. Around 17% of the cases are developing acute respiratory distress syndrome (ARDS), among which 65% cases are deteriorated to the extent of septic shock and/or multiple organ failure [5]. It is also evident that patients with underlying medical conditions such as cardiovascular diseases, diabetes mellitus, renal failure, respiratory diseases are more susceptible to nCOVID-19 infections [6,7]. So far, symptomatic therapies along with some nonspecific antiviral therapies have been tried as a therapeutic regimen of nCOVID-19 positive cases as the development of nCOVID-19 specific therapies (including vaccines) are still under clinical trials and yet to be available for the treatment purpose.

The ongoing worldwide investigations on nCOVID-19 have highlighted many insights into this infection like viral structure, immunopathology, associated symptoms, diagnostic approaches, possible preventive measures, and control, etc. The impacts of this pandemic over the global socio-economic condition, mental health of the society, etc., have also been studied [8]. In this context, it may be mentioned that social distancing, long term lockdown, discontinuation of personal work routine, the constant fear of the infection, concern for family and friends, fear of losing financial independence are collectively affecting badly the mental health of a large section of the population [8]. This leads to an increase in the level of stress followed by anxiety, depression, sleeplessness, and more chances to expose in the diseased conditions [9]. Many misconceptions and unauthentic information related to nCOVID-19 are circulating in the social media platform which causing more panic among the masses [10]. In this review, the characteristics of nCOVID-19, its possible diagnosis, preventions, therapeutic interventions, and management have been discussed in detail based on the recent investigations going on worldwide. This summarized yet informative content of the review article will help to get an overall impact on the idea of the entire status of this ongoing nCOVID-19 pandemic at a glance which will help the mass to get a clear cut knowledge about this current pandemic in our belief.

Respiratory viruses

Some major viral pathogens such as influenza virus, rhinovirus, adenovirus, respiratory syncytial virus (RSV), and coronavirus are responsible for maximum cases of morbidity and mortality due to the development of acute respiratory diseases (ARD) [11,12]. The clinical manifestations of respiratory infections generally help to characterize the pathogens such as rhinovirus can be characterized by typical common cold and RVS by bronchiolitis [13]. But the common viral respiratory syndromes are often associated with more than one viral infection which is presented in Table 1.

Coronavirus

As described in Table 1, Coronavirus is a class of respiratory viruses belongs to the coronaviridae family. The main vector of this virus is the bat and the infections occur to the human...
The new coronavirus isolated in January 2020 is SARS-CoV-2, a diverse single-stranded RNA virus, probably originated from the bat with a substantially different genetic sequence concerning the other known coronaviruses [44,45]. Unlike SARS which has a shorter incubation period of 1-4 days, nCOVID-19 has a latency of up to 14 days under which the virus can infect an average of 3.77 numbers of other people [44,46]. The infection mechanism is dependent on the envelope spike (S) glycoprotein which facilitates receptor binding, membrane fusion, and host tropism as well as transmission power of the virus [45]. This trimeric S protein cleaved in S1 and S2 subunit during viral infections [47]. S1 consists of the receptor-binding domain (RBD) and S2 is responsible for membrane fusion. S1 directly binds to the peptidase domain (PD) of angiotensin–converting enzyme 2 (ACE2) in the host receptor and host

by various other cross-species such as avians, rodents, and chiropets [32,33]. Earlier six subtypes of human coronaviruses have been identified as follows: HCoV-HKU1, HCoV229E, HCoVOC43, HCoV-NL63, severe acute respiratory syndrome coronavirus (SARS-CoV), and middle east respiratory syndrome coronavirus (MERS-CoV) [34].

In 2002, SARS-CoV, the novel human Coronavirus, has been first recognized as a cause of atypical pneumonia mostly in older adults and a probable cause of epidemic which affects around 8000 humans worldwide after originating from Guangdong Province, China [35]. SARS-CoV is a subclass of Zoonotic virus having horseshoe bats as reservoir hosts [36]. The important traits of SARS-CoV are (a) potentially high RNA mutation rates (2×10^{-6}) in contrast to other RNA viruses and also being able to encode 3′→5′ exonuclease RNA proof-reading activity within nonstructural protein 14 (nsp14) [37,38], (b) coronavirus family frequently recombinant between genomes and subgenomic replication complexes during mixed infections [39] and (c) being a larger sized RNA virus (27–31 kb), coronavirus can change or modify as they have numerous open reading frames towards the 3′ end of the genome and shows the characteristics of rapid adaptation to the novel hosts [37].

MERS-CoV, causing respiratory disease in humans, has been reported first in June 2012 [40]. Human MERS-CoV is a 30 kb positive single standard RNA genome that closely resembles to lineage C β-coronaviruses of Tylonycteris bat CoV HKU4 and also with the MERS-CoV derived from camels with whom it has >99.5% identical nucleotides [41,42]. MERS-CoV infection has been reported to cause around 851 deaths in 27 countries due to a lack of commercial vaccines and therapeutic interventions as reported by WHO until September 2019 [43].

**The new virus SARS-CoV-2**

The new coronavirus isolated in January 2020 is SARS-CoV-2, a diverse single-stranded RNA virus, probably originated from the bat with a substantially different genetic sequence concerning the other known coronaviruses [44,45]. Unlike SARS which has a shorter incubation period of 1-4 days, nCOVID-19 has a latency of up to 14 days under which the virus can infect an average of 3.77 numbers of other people [44,46]. The infection mechanism is dependent on the envelope spike (S) glycoprotein which facilitates receptor binding, membrane fusion, and host tropism as well as transmission power of the virus [45]. This trimeric S protein cleaved in S1 and S2 subunit during viral infections [47]. S1 consists of the receptor–binding domain (RBD) and S2 is responsible for membrane fusion. S1 directly binds to the peptidase domain (PD) of angiotensin–converting enzyme 2 (ACE2) in the host receptor and host

### Table 1: Respiratory virus and their associated syndromes.

<table>
<thead>
<tr>
<th>Type of Respiratory viruses</th>
<th>Genus</th>
<th>Symptoms</th>
<th>Diagnostic tools</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Influenza virus</td>
<td>Influenza virus A, B, C, and D &amp; Orthomyxoviridae family, Isavirus, Quanjanvirus and Thogotovirus genus</td>
<td>Major symptoms: Influenza-like illness, pneumonia, cough.</td>
<td>Viral culture; IFA; DNA-microarray based, sequencingbased tests, ELISA, PCR.</td>
<td>[14-17]</td>
</tr>
<tr>
<td>Adenovirus</td>
<td>Adenoviridae family, Atadenovirus, Avladenovirus, Mastadenovirus and Siadenovirus genus</td>
<td>Major symptoms: Pneumonia.</td>
<td>Viral culture; Indirect ELISA; IFA; LAT; EIA; Real-time PCR based</td>
<td>[18-21]</td>
</tr>
<tr>
<td>Coronavirus</td>
<td>Coronavirus family &amp; coronavirus (alpha, beta, delta and gamma genus)</td>
<td>Minor symptoms: Influenza-like illness, common cold, bronchiolitis.</td>
<td>RT-PCR; rRT-PCR; RT-LAMP</td>
<td>[22-25]</td>
</tr>
<tr>
<td>Rhinovirus</td>
<td>Picornaviridae family &amp; Enterovirus genus</td>
<td>Minor symptoms: Common cold.</td>
<td>CFT; HI; IFA; ELISA; Semi-nested RT-PCR assay; One step Panenterhino /Ge/08 real-time RTRCR assay;</td>
<td>[26-28]</td>
</tr>
<tr>
<td>Human RSV</td>
<td>Paramyxoviridae family &amp; Pneumovirus genus</td>
<td>Minor symptoms: Pneumonia.</td>
<td>RT – PCR</td>
<td>[29-31]</td>
</tr>
</tbody>
</table>

*Abbreviations:* CFT: Complement Fixation Test; DFA: Direct Fluorescent Assay; EIA: Enzyme-Immuoassay; ELISA: Enzymelinked Immunosorbent Assay; HI: Hemagglutination inhibition; IFA: Immunofluorescence assay; LAMP: Loop-mediated isothermal amplification; LAT: Lateral-Flow Test; LFIA: Lateral-Flow Immunochromatographic Assay; PCR: polymerase Chain Reaction; RT-PCR: Reverse Transcription-Polymerase Chain Reaction; rRT-PCR: Real-Time Reverse Transcription-polymerase chain reaction; RT-LAMP: Reverse Transcription Loop-Mediated Isothermal Amplification.
protease cleaved the S2 cleavage site and initiate the critical process of viral infections [47-49]. In recent research, it has been established that the ectodomain of the S glycoprotein of SARS-CoV-2 binds with the PD of ACE2 with a dissociation constant (Kd) of ~ 15nM [50]. ACE2 in human physiology is a type 1 membrane protein expressed in the lungs, heart, kidneys, and intestine which helps in the maturation of peptide hormone angiotensin that controls the vasoconstriction and blood pressure [51,52]. The reported cryo-electron microscopy (cryo-EM) structures suggest that two S proteins of SARS-CoV-2 trimers can simultaneously bind to an ACE2 homodimer [53]. This binding of S protein of SARS-CoV-2 with ACE2 is 10- to 20-fold greater in attraction than with S protein of SARS-CoV virus, which may be a reason behind the rapid human to human transmission worldwide [54,55]. In this context, it may be mentioned that this ACE2 expression rate is higher in less effected X-heterozygous females compared to males, as the ACE2 gene lays on the X-chromosome. Further, several genes involved with inflammation and immune response (ABO locus, SRY, SOX3, ADAM17) are also located on the X-chromosomes and directly or indirectly plays an impact on the ACE1/ACE2 balance. Rebalancing of this ACE1/ACE2 ratio or higher level of ACE2 might reduce the inflammation, thrombosis, and death related to nCOVID-19 [56]. The monoclonal antibodies (mAbs) S230, m396, and 80R showing cross-reactivity with SARS-CoV-2 [57]. The reported cryoelectron microscopy (cryo-EM) structures suggest that two S proteins of SARS-CoV-2 have no possible binding affinity towards SARS-CoV-2 RBD despite the structural homology between SARS-CoV and SARS-CoV-2 [50]. It may further suggest that SARS-directed mAbs will not necessarily cross-reactive with SARS-CoV-2 and this information will help in antibody isolation and therapeutic design for SARS-CoV-2 [50]. It is also evident that SARS-CoV-2 employs transmembrane protease serine 2 (TMPRSS2) for S protein priming. S protein priming by TMPRSS2 is found to be essential for the entry of the virus into the target cells as well as the spreading of infections [57-59]. Like many other infectious diseases such as influenza A viruses and coronaviruses, this TMPRSS2 is the critical factor for spreading of SARS-CoV-2 and thus a convenient drug target to combat SARS-CoV-2 [57-62]. In this context, it may be mentioned that camostat mesylate a serine protease inhibitor has blocking activity against TMPRSS2 [57,58].

Structural elucidation of the receptor-binding domain of SARS-CoV-2

The X-ray crystallographic study of SARS-CoV-2 RBD as demonstrated by Lan, et al. (2020) has shown loops forming core along with short connecting helices with twisted five-stranded antiparallel β sheet (β1, β2, β3, β4, and β7) [60-63]. An extended insertion formed with short β5 and β6 strands, α4 and α5 helices and loops forms receptor-binding motif (RBM) is the main contacting residues for ACE2 binding. The β sheet structure is stabilized by three pairs of a cysteine residue. This extended concave outer surface in RBM of SARS-CoV-2 RBD gets an attachment to the bottom side of the N-terminal helix ACE2 small lobe Figure 2.

**Immunopathology of nCOVID-19**

The SARS-CoV-2–induced immune responses are occurred in two phases [64]. During the first phase of incubation, the specific adaptive immune response gets activated to eliminate the virus and to mitigate the disease progression [64]. The development of this endogenous protective immune response during the early phase solely depends on the health of the patients and the immune-boosting approaches are helpful in this phase only. In this context, it may be mentioned that genetic differences are strong contributors to the variation in the immune response to pathogens in individuals [64]. A recent investigation of Ellinghaus, et al., (2020) have shown a 3p21.31 gene cluster as a genetic susceptibility locus for the nCOVID-19 patients as a reason for respiratory failure. The ABO blood group system has a potential involvement in this risk as locus 9q34.2 coincided with the ABO blood group locus. The study further revealed that blood group A has a higher and blood group O has lower chances of respiratory failure during nCOVID-19 infection in comparison to the other groups [65]. Impairment of immune response initiates the viral attack to the high ACE2 expressed organs such as intestine and kidney. In the second phase of the disease, cytokine release syndrome (CRS) or increased level of pro-inflammatory macrophages and granulocytes are observed followed by inflammation of lungs, respiratory distress, and associated disorders in severe stage [66]. Laguna-Rangel and Chávez-Valencia (2020) have reported that severe nCOVID-19 patients have higher IL-6/IFN-γ ratio than moderate nCOVID-19 patients which could be the possible reason behind the cytokine storm followed by lung damage [67]. Cytokine storm syndrome is a resultant effect of excessive release of pro-inflammatory stimuli and/ or deregulation of inflammation due to host-derived or environmental causes and associated with severe systemic inflammation, hemodynamic instability, multiple organ failure, and finally death [68]. These proinflammatory stimuli are superantigens (causing massive activation of T-cell receptors), antigens, toll-like receptor (TLR) ligands, allergens, or proinflammatory cytokines [68]. The lack of capacity to induce an immune response in the patient of SARS-CoV-2 may be the reason behind the remaining or returning of viruses after the complete cure of the patients [67]. In severe nCOVID-19 patients, lymphocytopenia is also observed with a significant reduction in B cells, natural killer (NK) cells, CD4+ T cells, and CD8+ T cells, monocytes, eosinophils and basophils along with an increase in neutrophil count [66,69,70]. The upregulation of exhaustion markers such as NKG2A on cytotoxic lymphocytes (such as NK, CD8+ T cells) is observed in nCOVID-19 positive
patients [71–74]. SARS-CoV-2-specific antibodies have been detected from the plasma of convalescent patients and used to treat SARS-CoV-2 positive with acute respiratory distress syndrome (ARDS) patients [75]. Another pathological condition of nCOVID-19 is thrombocytopenia which has been observed in 36.2% of cases of severe nCOVID-19 positive cases and also carrying the risk of developing disseminated intravascular coagulation (DIC) [76–78]. It has been reported as per the International Society on Thrombosis and Haemostasis diagnostic criteria that DIC has been observed in 15 out of 21 non–survivors of the nCOVID-19 patients [79]. Some other reports have shown by a meta–analysis that the presence of thrombocytopenia increases the chance of severe diseases up to five folds whereas in severe disease the platelet counts decrease (mean difference: -31×10^9/L, 95% CI: -35 to -29×10^9/L) [80]. The occurrence of thrombocytopenia and elevated D–dimer are probably due to the over activation of coagulation cascade and platelets as the procoagulant and anticoagulant homeostatic mechanisms get disrupted during viral infections [81]. This phenomenon also involves endothelial dysfunction, Toll–like receptor activation, von Willebrand factor elevation, and tissue–factor pathway activation [80,81]. It is also known that platelets, the key mediator of inflammatory pathways, are responsible for white blood cell activation and clot formation after interacting with pathogen pattern recognition receptors or immunoglobulin Fc receptors and complement receptors (pathogen recognition) [82]. Disruption of this mechanism has a major role in viral infections associated procoagulant effect [82,83]. One rare pathological condition of nCOVID-19 is viral encephalitis which has been found in one nCOVID-19 infected patient in Beijing, China. The study with cerebrospinal fluid (CSF) of nCOVID-19 infected patients has revealed the presence of SARS-CoV-2 in the central nervous system (CNS) and showing the symptoms of encephalitis [84].

Symptoms of nCOVID-19

The nCOVID-19 is mainly transmitted through the respiratory droplets from coughing and sneezing and the common symptoms are fever and respiratory symptoms like cough and dyspnoea, and also there is evidence of bilateral atypical pneumonia during infections. It has also been observed that aerosolized SARS–CoV–2 particles remain for hours in the air and are able to transport over a usual distance (6 feet) like outside of rooms and intra building [85]. WHO has also reported the existence of this virus in stools of the patient in 30% cases which persist up to 4–5 weeks, although not known whether it signifies the presence of an infectious virus or not [2,86,87]. It has been found that nCOVID-19 also present in the glandular cells of the rectum along with interstitial edema and lymphoblasts–cytosis, although their association with pathological evidence such as diarrhea during nCOVID-19 infection has not been clear yet [2,88]. The increased level of proinflammatory cytokines (IL–1β, TNF–alpha) has an association with inflammatory bowel diseases (IBD), characterized by fever, abdominal pain, and diarrhea, during relapse and remission phases of nCOVID-19 [86,89]. Zhang, et al. (2020) have investigated by using single–cell transcriptomic analysis that the digestive system is also a potential route of SARS–CoV–2 infections along with the respiratory tract [90]. Zhang, et al. have also suggested that this may be a possible reason behind symptomatic diarrhea as an enteric symptom of nCOVID–19 as the sites of enterocytes are highly expressed with ACE2 and TMPRSS2 [89]. The rare case of nCOVID19 associated encephalitis has shown meningeal irritation signs in the patients along with low WBC count (3.3 × 10^9/L) and lymphopenia (0.8 × 10^9/L) [91].

Diagnosis of nCOVID-19

Globally, RT–PCR is recognized as an only validated tool for the confirmation of nCOVID19 cases to verify the asymptomatic cases (those who have been in close contacts of nCOVID–19 positive cases) and also to differentiate nCOVID–19 from the other respiratory infections [92–94]. As suggested by WHO, various RT–PCR protocols are examining the amplification of nCOVID–19 specific genes i.e. (a) envelope protein (E); (b) RNA–dependent RNA polymerase (RdRPR); (c) open reading frame 1a and b 226 (Orf1a,b) ; (d) nucleocapsid protein; (N) genes of SARS–CoV–2; or (e) combination of those genes (two or more), for screening and identification of nCOVID–19 positive cases. At least one of these genes should be detected by RT–PCR for interpreting the result as positive as per the guideline of WHO [94–98]. At present, the knowledge regarding nCOVID–19 is not fully proved, so it has been recommended to collect the specimen from the upper respiratory tract (Nasopharyngeal swab, oropharyngeal swab, nasopharyngeal aspirates, etc.) and also from the lower respiratory tract (Sputum, bronchial washing, tracheal aspirates, transtracheal aspirates, bronchoalveolar lavage, tranchonbral lung biopsy, etc.) within seven days of symptoms onset [99]. Reports are available regarding the possibilities of transmission of nCOVID–19 via the fecal–oral route [100–102]. SARS–CoV–2 viral nucleotide detection has been found to be positive in anal swabs but negative in nasopharyngeal swabs even after 42 days of infections suggesting the requirement of resting the fecal specimen of the nCOVID–19 patients also [103].

Due to the shortage of laboratory–based molecular testing capacity and reagents, diagnostic test manufacturers are now more focused on manufacturing easy to use devices to expedite the testing process in less time [104]. The test kits are mainly able to detect (a) viral protein fraction from the human respiratory sample or (b) human antibodies generated in response to infection. For the detection of the target antigen, the testing kit usually contains specific antibodies fixed paper strip or plastic casing that can give visual signal upon antigen–antibody reactions [104]. The testing process usually takes 30 minutes of time and the accuracy rates varying from 38% to 80% in different kits [104,105]. The testing is only able to detect in acute or early infection time when the target antigens are present in sufficient concentration [104]. The limitation of such a test is the specificity of antibodies towards nCOVID–19 antigen as antibodies on the test strip may also be able to recognize the human coronavirus that causes the common cold. In that case, the test should be performed only in those patients whose symptoms are very likely to nCOVID–19. Due to this reason WHO has not recommended these types of antigen–detecting rapid diagnostic tests for patient care. The other
kind of readily available test kit can detect the virus–specific antibody in the blood or plasma of the patient [106,107]. Several studies have suggested that antibody response develops only in the second week of infection, which means detection of the infection by antibody response is possible in the recovery phase and may not be effective enough to control the transmission or implementing clinical interventions for therapy [106,108–110]. Chances of cross–reaction are still possible and may produce false–positive results that is why WHO has not recommended this type of test for patient care so far [104,111,112].

**Preventive measures for nCOVID-19**

1. National Health Commission of the People’s Republic of China has recommended the following precautionary measures for the infected patients according to the different phase of nCOVID-19 infections [113,114].

   (a) During the initial days of infections, suspected patients may be considered for home isolation with routine health check-ups if the symptoms of the patient are mild. These classes of patients should be treated with antiviral or other recommended drugs for symptomatic relief in the initial stage to prevent further deterioration of health. [114,115].

   (b) Patients with worst symptoms such as aged patients and those with other medical issues should be treated with more care and repetitive assessment based on routine chest CT scan of the infected individuals [113].

   (c) If in some patients, after 10 days, any expansion of lung damage or acute cardiovascular diseases persists with an increase in inflammatory responses, then the necessary treatments are required to control immune modulation [113].

   (d) More progression in infection would become more difficult for critically ill patients to tolerate and in that phase, the mortality rates become high/maximum [113].

2. Guidelines, as mentioned below, are needed to be followed as precautionary/ preventive measures by the patients and the persons who are in close contact with the patient (Persons who are suspected of nCOVID-19 exposure) as recommended by several authorities:

   (a) Advised to take 14 days of self–quarantine from the day of exposure [115,116].

   (b) Medical attention to be taken, if any kind of fever or coughing, shortness of breath, or diarrhea observed during this period [115].

   (c) Surveillance is needed for the persons who have been exposed at a low–level to the suspected or nCOVID–19 positive patients [116].

3. Preventive measures recommended for travelers:

   (a) Routine precautions should be taken before entering and leaving the nCOVID–19 affected zones [117].

### Table 2: Home remedies and precautions for early prevention of nCOVID–19.

1. Regular drinking of warm water.
2. Minimum 30 minutes practice of yoga and meditation.
3. Daily administration of natural immune boosters like turmeric, cumin, coriander, and garlic, etc.
4. Drinking of herbal tea with basil, cinnamon, black pepper, dry ginger, raisin once or twice a day.
5. Consumption of milk with turmeric powder.
6. Application of sesame oil, coconut oil, or ghee in both the nostrils twice a day (in morning and evening).
7. Inhalation of steam with mint leaves or caraway seeds.
8. Application of clove powder with honey in a sore throat or cough.
9. Practicing the oil pulling therapy by swishing 1 tablespoon sesame or coconut oil in the mouth for 2 to 3 minutes and then rinsing the mouth with warm water for once or twice a
10. Naturopathic nutrition (balanced diet with fruits and vegetables) is important for improving innate immunity, especially in comorbid patients.
11. Intermittent fasting or lemon juice fasting helped in boosting the immune system; reduce oxidative stress as well as may improve autophagy in comorbid patients.

The above information are taken from Ayush, Indian Council of Medical Research, Government of India [118].

(b) Frequently washing the hands especially after being in contact with affected persons [117].

(c) Avoidance of contact with dead farming animals or wild animals [117].

Recently the Government of India has published a list of home remedies and precautions for early prevention of nCOVID–19 which is expected to be effective as precautionary measures against the viral contaminations (Table 2) [118]. The government of India has also recommended “Dinacharya” –daily regimes and “Ritucharya”–seasonal regimes for immune–boosting based on Ayurveda’s extensive knowledge in relation to prevention, care and to maintain a healthy life [118]. However, the application of “Dinacharya and Ritucharya” with the precautions of nCOVID–19 needs further investigation [118].

United nation and WHO both recommended yoga/meditation to reduce disease–related anxiety or stress [119]. Yoga is also recommended as a preventive measure for infection control as it has been well proved that yoga can reduce the IL–6 and TNF–alpha as well as decrease the IL–1beta showing an effective impact on populations at risk or already suffering from the diseases associated with inflammations, although the consistent and regular practice is essential for achieving effective results [120].

Another possible aspect is the homeopathic treatment for the prevention of nCOVID–19 infection. The Cuban government has already promoted proactively the use of homeopathic medicine as protection for elderly persons against nCOVID–19 infection [121]. Cuba successfully prevented the outbreak of Leptospirosis epidemic by using homeoprophylactic.
approaches (oral administration of highly diluted pathogens) in the year 2007 [122,123]. In January 2020, Govt. of India has recommended Arsenicum album 30C as a preventive measure [123]. Recently in Naiminath Homeopathic Medical College, hospital and Research Centre, Agra, India, trials of homeopathic medicines on 100 nCOVID-19 patients have been initiated by providing a combination of Arsenicum Album 30, Bryonia Alba 30, Gelsemium, Antimonia Tartaricum and Crotalus Horridus, to the patient by Dr. Surabh Kumar with an expectation of preventing the transfer of asymptomatic symptoms of nCOVID-19 to the symptomatic condition [124].

Administration of micronutrients like vitamins, minerals, etc. can protect the respiratory tract as antioxidants are important factors to boost immune function during the nCOVID-19 pandemic [125]. Vitamins like A, C or E are a good source of anti-oxidants which helps to increase (a) T-cell subsets, (b) lymphocyte response to mitogen, (c) interleukin-2 production, (d) potential of the activity of the natural killer cell, and (d) response to influenza virus vaccine [126]. Vitamin D, which is likely to synthesized less in the body during quarantine due to lack of exposure in sunlight, able to protect the respiratory tract by preserving tight junctions and also helps to decrease the level of pro-inflammatory cytokines followed by reducing the risk of cytokine storm leading to pneumonia [127]. It is also evident that zinc has the potential to inhibit severe acute respiratory syndrome (SARS) coronavirus RNA-dependent RNA polymerase (RdRp) template binding [128].

Therapeutic interventions of nCOVID-19

The in-vitro activity and clinical applications of some drugs applied so far for the treatment of nCOVID-19 are summarized below although none of them has been claimed to mitigate this infectious disease:

**Chloroquine and hydroxychloroquine:** Chloroquine and hydroxychloroquine, the anti-malaria and anti-inflammatory drugs, are now in limelight as a potential therapy of nCOVID-19 [129]. The mechanism of action of chloroquine and hydroxychloroquine is the inhibition of glycosylation of host receptors, proteolytic processing, and endosomal acidification followed by blocking the viral entry inside the host body [130,131]. Two research groups have observed the antiviral efficacy of chloroquine and hydroxychloroquine against SARS–CoV in a high dose regimen showing a reduction in the exacerbation of pneumonia and reducing the disease course. The hydroxychloroquine, a derivative of chloroquine, has EC50 14μM against SARS-CoV-2 which is much lower than the EC50 of chloroquine, i.e. 23.90 μM [132]. Long-term administration in high dose (400mg/day orally for 5 days in case of hydroxychloroquine or 500mg/ day orally for chloroquine) of these drugs are associated with several side–effects (in less than 10% cases) such as myopathy, arrhythmias such as ventricular tachycardia and torsades de pointes [133–135]. China has reported the successful application of chloroquine in the treatment of more than 100 nCOVID-19 patients with improved radiological findings but the report is not yet published in any peer-reviewed journals [129]. Also, a nonrandomized French trial has published the report of 20 in the hydroxychloroquine group (receiving 200mg hydroxychloroquine by mouth in every 8 hrs) and 16 in the control group and found positive results in treated group with virologic clearance at day 6 and even better result with concomitant use of azithromycin [136]. But the associated cytotoxicity of drugs cannot be avoided as very recently, Borba, et al. (2020) have mentioned that this effect of chloroquine or hydroxychloroquine is lethal for aged nCOVID–19 infected patients especially those who have the previous report of cardiac diseases and application of such drugs need randomized clinical trials before further applications [137]. The mixed report as obtained for chloroquine and hydroxychloroquine is not able to state the exact mechanism of action of these drugs on nCOVID–19 as it showed effectively in some patients and thus need further exploration of the scientific background of the drugs before drawing any positive or negative conclusion.

**Antiretrovirals:** Anti–HIV antiviral such as Lopinavir/ritonavir is also active against other coronaviruses and act by inhibiting 3–chymotryptase–like protease although no published literature is available regarding their activity against SARS–CoV–2 [138–140]. The report available so far are retrospective, nonrandomized cohort studies, and difficult to conclude [141,142]. The dosing regimen for nCOVID–19 for lopinavir/ritonavir as per the report of the National Health Commission and State Administration of Traditional Chinese Medicine (March 2020) and Chao, et al. (2020) is 400mg/100mg twice daily for up to 14 days [143]. The drugs have significant drug–drug interactions and adverse effects i.e. gastrointestinal distress such as nausea and diarrhea, hepatotoxicity, etc which creates limitations for their applications [144,145]. Similarly, other antiretrovirals like darunavir have shown anti–SARS–CoV–2 activities in-vitro but their effects on humans are still under investigation [140,144].

**Antivirals other than antiretrovirals:** Ribavirin is an RNA polymerase inhibitor and effective against other coronaviruses [129]. However, information related to its activity against SAR–CoV–2 is limited and so far only extrapolated from results against other coronaviruses [129]. A high dose of ribavirin or in combination therapies has shown in vitro activity against SARS–CoV but that is also associated with several side effects and toxicities such as hematologic and liver toxicity [146]. This substantial teratogenicity and toxicity is the limitation of Ribavirin as a potential therapeutics of nCOVID–19 [147]. Influenza–like symptoms have been treated with Oseltamivir like neuraminidase inhibitor but it has no in–vivio activity against SARS–CoV–2 [148]. So far none of the clinical trials have included oseltamivir as proposed therapeutic interventions [149]. Recently, China and Russia have approved umifenovir as a potential antiviral against SARS as this drug can target S protein/ACE2 interaction and inhibit the membrane fusion of the viral envelope [150,151]. A nonrandomized clinical data has recently established that the use of umifenovir for a median duration of 9 days in nCOVID–19 patients shows a lower mortality rate and higher compared to those who not receiving this drug [152]. Anti Ebola virus drug remdesivir or GS-5734, is a monophosphate pro–drug which converted
into active C-adenosine nucleoside triphosphate analog after metabolism, is now in limelight as a potential candidate for nCOVID-19 therapy due to its anti–RNA virus activities against coronaviridae and flaviviridae [153]. Similarly, another anti–Ebola analog favipiravir or T705, a purine nucleotide which after activation inhibits the RNA polymerase and halt the viral replication, has shown a broad activity against RNA viruses [154]. EC50 of favipiravir against SARS–CoV–2 is 61.88 μM/L in Vero E6 cells [155]. For the treatment of the SARS–CoV–2 higher range of dosing has been proposed such that the recommended loading dose is 24,000mg to 30000mg every 12 hours/day followed by a maintenance dose of 1200mg to 1800mg every 12 hours per day considering its mild side-effects in comparison to other antiviral drugs [156]. Recently, India–based Glenmark Pharmaceuticals has got the regulatory approval in India for the manufacturing and marketing of favipiravir under the brand name FabiFlu for the treatment of mild to moderate nCOVID–19 infections. As per the data obtained by their phase III clinical trial, the company has claimed that favipiravir has shown promising results in mild to moderate nCOVID–19 in age groups of 20 to more than 90 years and provide rapid decrease in viral load within four days, along with quicker symptomatic relief [157].

Vaccines: Vaccines are a fundamental requirement for long term protection against infection outbreaks [158]. Various approaches have been taken for the development of vaccines for nCOVID–19 worldwide. The S glycoprotein is mainly responsible for the receptor binding of the host cell, so it is a good target for the development of host Recombinant Subunit Vaccine [159]. Recently, Clover Biopharmaceuticals, Chengdu, China, collaborated with GSK, Middlesex, UK, and have announced the pre–clinical testing of a recombinant subunit vaccine for nCOVID–19. The vaccine has been generated with native–like trimeric viral spike, similar to trimeric S protein (S–Trimer), by using a mammalian cell culture–based expression system [160,161]. The immune response of the vaccine has been improved by using GSK’s adjuvant system to S Trimer and finally, the detection of antigen–specific neutralizing antibodies has been done in the sera of fully–recovered nCOVID–19 patients [160,161]. Subunit vaccine is also under development by the University of Queensland, Australia, as they are using a transformative technology “molecular lamp” which is a polypeptide that helps to stabilize a surface protein and improves antigen recognition proving a stronger immune response [162]. Other than the subunit vaccine, DNA vaccines can also be served by direct injection of plasmids encoding the antigens followed by a wide range of immune responses [163]. The DNA vaccine development is focused on the improvement of the efficacy by using electroporation to deliver plasmids [164]. Inovio Pharmaceuticals, Plymouth Meeting, PA, and Beijing Advaccine Biotechnology, Beijing, China are conducting pre–clinical trials for DNA vaccine (INO–4800) against nCOVID–19 [165]. INO–4800 activates T cells by delivering DNA plasmids with the spike protein of SARS–CoV–2 [166]. Intradermal application is also under investigation to produce therapeutic antibodies and activate immune cells. The phase I trial is initiated by Inovio Pharmaceuticals in the USA and China with the help of the Coalition for Epidemic Preparedness Innovations (CEPI) body [167]. On the other side, Moderna, Inc., Cambridge, MA, in collaboration with the National Institute of Allergy and Infectious Diseases (NIAID), USA has started the phase I clinical trials for an in silico mRNA vaccine (mRNA–1273) encoded with viral Spike(S) protein [168]. This mRNA–based vaccine contains mRNAs encoding the antigens having advantages of absence of genome integration causing improved immune responses followed by the rapid development of multimeric antigens [158,169]. Genexine Inc., Gyeonggi–do, Republic of Korea, is also developing an nCOVID–19 vaccine using Hyleukin–7 platform technology by fusion of interleukin–7(IL–7) to hyFc [170]. This fusion is designed to hybridize the flexible hinge structured IgD and an unexposed junction site IgG4 which increases the sustained action of Fc fusion proteins and prevents the antibody–dependent cellular cytotoxicity (ADCC) and complement–dependent cytotoxicity (CDC) [170,171]. Very recently AstraZeneca and the University of Oxford have conducted a phase III clinical trial (ISRCTN89951424) in Brazil to determine the safety, efficacy and immunogenicity of the non–replicating ChAdOx1 nCoV–19 vaccine. The single intervention of this vaccine will be followed by 1g paracetamol every 6 h for a day. It is expected to get the trial outcome by July 2021 [172].

External respiratory support: Studies suggest that respiratory failure is the main reason behind nCOVID–19 death [173,174]. The patients with respiratory distress and/or hypoxemia transnasal high–flow oxygen, endotracheal intubation, and invasive mechanical ventilation should use the external respiratory support as and when necessary if standard oxygen therapy does not function properly. In severe cases, a fiberoptic bronchoscope may be required for high–density sputum suction or bronchoalveolar lavage for their relief [175].

Myocardial protection: Vasoaducting drugs like norepinephrine or dopamine, dobutamine for increasing the systemic functions, creatine sodium phosphate, vitamin C, coenzyme Q for avoiding comorbid myocardial injury during severe nCOVID–19 infections, conservative fluid treatment for fluid resuscitation and to improve oxygen supply to tissues are also sometimes needed to the patients in severe conditions [176,177].

Artificial liver support therapy: Treatments like plasma replacement, blood adsorption, perfusion, artificial liver blood purification system should be carried out in the severe nCOVID–19 patients with excessive inflammatory responses and failure of the liver. It may help to reduce the level of excess cytokines [175].

Steroids: Low doses of glucocorticoids for the short term could be a possible therapy for excessive inflammatory responses, progressive worsening of oxygenation markers, and rapid progressive imaging. Glucocorticoids can be considered in severe nCOVID–19 cases to prevent the progression of ARDS [175].

Specialized pro–resolving lipid autacoid mediators (SPMs): Resolvins and other SPMs, derived from omega–3 fatty acids have a stimulatory effect on the macrophage–mediated clearance
of debris and also found to counteract the pro-inflammatory cytokine production by the process, inflammation resolution [178]. Thus, SPMs affect viruses in a very low dose (in nanogram level) without being an immunosuppressive. SPMs also promote various anti-viral B cell antibodies and lymphocytes, making it a potential candidate for the treatment of nCOVID-19 [178].

Nutritional supplements for gut microbiota: The intestinal barrier can be disrupted by the high inflammatory responses during nCOVID-19 infection–causing bacterial translocation and secondary infections [179]. This phenomenon further increases the lipopolysaccharides influx which takes part in the release of TNF-α, IL1β, and IL-6, causing further exacerbation of systemic inflammation [180]. Gut and respiratory tract flora interact and it has been found that gut microbiota can reduce enteritis and ventilator–associated pneumonia [179,181]. Administration of probiotics, energy source(s), amino acid, and trace elements are thus essential for the nCOVID-19 patients for maintaining the equilibrium of the gut microbiota [175].

Vitamin C infusion: Administration of Vitamin C (15g/ day, I.V. for 4 days) to the nCOVID-19 patients with sepsis–related ARDS may have reduced the rate of mortality as observed in 167 ICU patients in the USA [182]. Very recently, a clinical trial for the investigation of the effect of Vitamin C infusion to treat the nCOVID-19 infected pneumonia has been initiated in Wuhan, China [183]. A total of 140 patients are taken under the clinical trial and infusion of 24 g of vitamin C per day for 7 days has been applied for trial [183]. The study is most likely to be ended by the month of September 2020. The patients are assessed based on the requirement of ventilation, organ failure score, length of stay in ICU, and also the rate of 28 days mortality [183].

Treatment with the plasma of recovered patients: From the experience of SARS and MERS, it is evident that plasma of recovered patients can be effective as a treatment of nCOVID-19 [184,185]. This is currently under clinical trial but it is expected to be effective on severe nCOVID-19 cases before applying any non-specific antiviral drugs although a very dynamic evaluation is needed [186,187].

nCOVID-19 and global mental health

Although the prevalence of nCOVID-19 is uncertain in the community there is no doubt that older persons are at much higher risk than the younger generation especially those with comorbidity [188,189]. The primary prevention for the aged generation is social isolation which is highly associated with depression, cognitive damage, cardiovascular, and other aging–related diseases ultimately leading to an increased rate of mortality. It is a well-known fact that people with mental illness have a lower expectancy of life than the general population [190]. A large group of populations is psychologically vulnerable due to poor physical and mental health, impaired access to services, and losing control over day-to-day normal lives. A greater focus is needed for the mental health of society during this lockdown phase [191]. Otherwise, anxiety, panic-like mental issues will worsen the outbreak of nCOVID-19 among the public. Retrospective case studies have shown that severe nCOVID-19 patients are showing neurological disorders like cerebrovascular diseases, consciousness impairment, and skeletal muscle symptoms [192]. The uncertainty of infection, their growth rate, and the corresponding mortality, etc. create a psychological disturbance in the population [193]. Meng and his team have conducted a survey in china with a total of 1556 samples and have observed that 37.1% of the seniors are experiencing depression and anxiety due to nCOVID-19 [194]. Qiu and others have recently shown that aged individuals are more prompt to emotional reactions [192]. They have also found that aged women are experiencing more anxiety and depression than aged men. However, no significant difference between age segments (60–64 years, 65–69 years, 70–74 years, 75–79 years, and above 80 years) are observed by them. A good diet, exercise, and sleep are most important during the phase of heightened emotion along with that reaching out to others for mutual support where ever required are also needed. Ironically the nCOVID-19 infection control needs public health measures that harm mental health such as social isolation causing fear of infection, frustration, boredom, and anxiety, etc [195]. The India based survey on the younger generation has shown that 80% of the educated young generation is preoccupied with the thought of the pandemic situation and 12.5% are facing sleep difficulties, 37.8% are going through paranoia and 36.4% of the participants are in distressful condition [8]. Even the academicians and researchers are also going through the same stressful situation due to lack of access to the university and their respective professional fields of activities [196]. The only option is to withdraw the social isolation as soon as the infection is under control and enough treatment is available in the market.

Conclusion

The global threat of nCOVID-19 has declared as pandemic and the world is still fighting with a crisis and millions of lives are still under threat. The occurrence of infection is based on age and sex following a ratio of 1:0.3:1 in the chances of infection between males and females. The diverse single-stranded RNA virus SARS-CoV–2, showing severe respiratory and bowel syndromes, is responsible for impairment of immune response followed by cytokine release syndrome, and carrying the risk of disseminated intravascular coagulation. Prevention of the infections by social distancing depending on the concept of aerosolized/droplet transmission of virus and maintaining personal hygiene are most important to break the viral chain as there are no fully proven remedies available for this virus so far. Some remedies like ayurvedic, homeopathic, micronutrients such as vitamins and minerals along with regular physical exercises like yoga and meditation are also important for disease prevention. The treatments such as antimalarial chloroquine and hydroxychloroquine, antivirals, plasma therapy, steroids, omega–3-fatty acid derivatives, etc. have been reported so far as a possible therapeutic regimen. So far no full proved antiviral treatment or vaccine is available for the treatment leaving control and prevention as the only option to deal with this deadly infection [197]. Not only heath,
but this global pandemic also has a huge impact over, social, economic, cultural as well as psychological damages [198]. One thing that cannot be ignored is the chances of the reoccurrence of this disease condition soon after nCOVID-19 with the same or some other infections and therefore, control is needed from every part of the society. It is important to understand and reorganized global healthcare policies and management. Clinicians also need to be trained with updated interim guidance on the effective, safe, efficient and prompt supportive management of the nCOVID-19 patients especially those who are admitted to the hospital with severe respiratory illness. The hospital authority should ensure that the clinical management of the patient and patient care should be started from the entry point of the hospital to reduce the risk of the contamination. It is important to implement the application of telemedicine and video consultations for regular health checkups and non-emergency cases to avoid the unnecessary contamination of the patients as well as health care professionals [199]. On the other hand, governments are responsible for social management like the implementation of strategies, rules, guidelines for food habits to prevent the recurrence of this type of disease in the future. The rules regarding the prohibition on consumption of fast foods, raw uncooked food eating should be taken into strict considerations, as food consumption is the probable reason behind the initiation of nCOVID-19 infections. While the sudden socio-economic turmoil has been taken care of by the government, personal safety, wellbeing, health, and hygiene is the responsibility of each one of the citizens. However, long term protection against nCOVID-19 is only possible when immune boosting can be achieved scientifically as well as by taking up certain changes in habits in daily lives such as consumption of fruits, antioxidants, exercise, yoga, proper sleep, and a routine healthy lifestyle. Finally, it may be stated that this is wise to pay proper scientific attention to the current situation and take it as a lesson for the future in all possible aspects of life.

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