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Microbial co-infections in COVID-19: Associated microbiota and underlying mechanisms of pathogenesis

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\begin{abstract}

The novel coronavirus infectious disease-2019 (COVID-19), caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has traumatized the whole world with the ongoing devastating pandemic. A plethora of microbial domains including viruses (other than SARS-CoV-2), bacteria, archaea and fungi have evolved together, and interact in complex molecular pathogenesis along with SARS-CoV-2. However, the involvement of other microbial co-pathogens and underlying molecular mechanisms leading to extortionate ailment in critically ill COVID-19 patients has yet not been extensively reviewed. Although, the incidence of co-infections could be up to 94.2% in laboratory-confirmed COVID-19 cases, the fate of co-infections among SARS-CoV-2 infected hosts often depends on the balance between the host’s protective immunity and immunopathology. Predominantly identified co-pathogens of SARS-CoV-2 are bacteria such as \textit{Streptococcus pneumoniae}, \textit{Staphylococcus aureus}, \textit{Klebsiella pneumoniae}, \textit{Haemophilus influenzae}, \textit{Mycoplasma pneumoniae}, \textit{Actinobacter baumannii}, \textit{Legionella pneumophila} and \textit{Clamydia pneumoniae} followed by viruses including influenza, coronavirus, rhinovirus/enterovirus, parainfluenza, metapneumovirus, influenza B virus, and human immunodeficiency virus. The cross-talk between co-pathogens (especially lung microbiomes), SARS-CoV-2 and host is an important factor that ultimately increases the difficulty of diagnosis, treatment, and prognosis of COVID-19. Simultaneously, co-infecting microbiotas may use new strategies to escape host defense mechanisms by altering both innate and adaptive immune responses to further aggravate SARS-CoV-2 pathogenesis. Better understanding of co-infections in COVID-19 is critical for the effective patient management, treatment and containment of SARS-CoV-2. This review therefore necessitates the comprehensive investigation of commonly reported microbial co-pathogens amid COVID-19, their transmission pattern along with the possible mechanism of co-infections and outcomes. Thus, identifying the possible co-pathogens and their underlying molecular mechanisms during SARS-CoV-2 pathogenesis may shed light in developing diagnostics, appropriate curative and preventive interventions for suspected SARS-CoV-2 respiratory infections in the current pandemic.

\end{abstract}

\section{Introduction}

The novel coronavirus infectious disease-2019 (COVID-19) is a rapidly transmissible pneumonia-like disease caused by the SARS-CoV-2 which emerged in Wuhan, China in December 2019, and is currently circulating throughout the world \cite{1,2}. The positive-sense enveloped RNA virus (SARS-CoV-2) is genetically different from the previously known coronaviruses such as SARS-CoV-1 and Middle East respiratory syndrome coronavirus (MERS-CoV) \cite{1–3}. Immediately after its first outbreak in China, this fearsome virus has emerged as one of the

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deadliest human pathogens in the last hundred years after the Spanish Flu in 1918–1920 [4,5]. The SARS-CoV-2 infection has become a public health challenge worldwide, and thus, the World Health Organization (WHO) has declared this disease as a public health emergency of international concern [1,2]. The deadly outbreaks of SARS in 2003 and MERS in 2012, with case fatality rate of 9.6% and 34.4%, respectively were successfully contained within six months. The COVID-19 disease affected total 217 countries and territories until April 7, 2021, and more than 133,688,126 cases have been confirmed globally with 2,901,038 deaths (https://www.worldometers.info/coronavirus/?utm_campaign=homeAdvegas17). Therefore, this quickly spreading COVID-19 pandemic highlights the critical need for rapid development of vaccines and antiviral treatments to reduce the number of hospitalizations and deaths by this worrisome pathogen [6].

Co-infections and superinfections are common in many respiratory viral infectious diseases [6,7]. Bacterial co-infections can significantly increase the mortality rate in patients infected with any viral infections [6,8,9]. Previously, bacterial co-infections were also reported in MERS-CoV patients requiring intensive care [10]. The co-infection of the SARS-CoV-2 with other microorganisms is a very important factor in COVID-19 pathogenesis that may complicate the accurate diagnosis, treatment, prognosis of COVID-19, and even increase the mortality rates [11]. Clinical trials and metagenomic investigations indicated the co-presence of other viruses, bacteria, archaea and fungi with SARS-CoV-2 in COVID-19 patients [11–13]. About 50% of the patients who died of COVID-19 had secondary bacterial infections [12,14] which further intensifies the pathophysiological progression of COVID-19. Better understanding of co-infections in COVID-19 is critical for the effective patient management, treatment and containment of SARS-CoV-2. It is therefore, necessary to strengthen the investigation of the co-infection in COVID-19 patients. Moreover, regarding COVID-19 several issues such as useful strategies to prevent disease spread, collection of appropriate clinical specimens, transmission route, viral dynamics and effective drug treatments are still largely unknown. However, the possibility of co-infections with other respiratory pathogens including bacteria, archaea, viruses (other than betacoronavirus) and fungi are not yet clearly understood. The association of these secondary pathogens to causing co-infections should be an important concern for the clinician in the management of COVID-19 cases. The emerging infectious diseases section of the Centers for Disease Control and Prevention (CDC, USA) endorsed testing for other respiratory pathogens, suggesting that evidence of another infection could aid the evaluation of patients with potential COVID-19 in the absence of widely available rapid testing for SARS-CoV-2 [15].

The two earlier coronaviruses (SARS-CoV-1 and MERS-CoV), influenza virus, and SARS-CoV-2 show highly similar respiratory symptoms, including high fever, cough, headache and even pneumonia [16–19]. Recent clinical and in silico studies showed that viral co-infections mainly includes respiratory viruses such as enterovirus/rhinovirus (hRV), human metapneumovirus (hMPV), Respiratory Syncytial Virus (RSV), Siphovirus, Alphapapillomavirus, Myovirus, Tumbavirus, Victoriaivirus, Parvivirus, Chrysoivirus, Totivirus, and other coronaviruses (non-SARS-CoV-2) [11,20]. Concurrent co-infections in COVID-19 can also change the respiratory microbiome homeostasis, and thus triggers the infection and stimulates immune cells to produce more severe inflammation [11,20]. The gut bacterial diversity of the COVID-19 patients is also reduced with the increased relative abundance of opportunistic pathogens, and it can affect the relative bacteria and alters the beneficial symbionts [21]. Recent metagenomic studies reveal the concurrent association of bacteria, archaea and non-COVID viruses in nasal swabs of COVID-19 patient [11]. During the earlier epidemics of SARS-CoV-1 and MERS-CoV, patients receiving invasive mechanical ventilation were easily developed bacterial co-infections, and had higher mortality rates [16,18]. Therefore, bacterial co-infections might be a key element that promotes severities of the COVID-19 and mortality rates [22]. Recent study on deceased patients showed that sepsis (100%) acted as one of the main complications [23], indicating that co-infection is of great importance to prognosis and subsequent treatment of COVID-19 patients. Furthermore, co-infection has been associated with more severe outcomes in pandemic and seasonal influenza [24]. It has been suggested that influenza-related bacterial infections overall may account for up to 30% of community-acquired pneumonia (CAP) cases [25]. Several studies of hospitalized patients with COVID-19 noted the empiric use of antibiotics in majority of patients [24,26,27]. However, there is an evidence of increased inflammatory serological markers associated with bacterial infections including procalcitonin and C-reactive protein in patients with COVID-19 without a corresponding bacterial co-infections [24,28]. Meanwhile, several descriptive studies showed that the ecosystem of commensal microbiota can both regulate and be regulated by invading viruses, facilitating either stimulatory or suppressive effects [23,29,30]. More importantly, the coinfected microorganisms may also be a new strategy for the development of new treatment of SARS-CoV-2 infection. Despite increasing evidence for its salience to COVID-19 outcomes, the effect of co-infection clearance on SARS-CoV-2 load has not yet been systematically reviewed or critically discussed. This systematic review updates our knowledge on the microbial co-infections associated with SARS-CoV-2 pandemic, and the possible molecular mechanisms of co-infections in COVID-19 to emphasize that microbial co-infections.

2. Rationale and review methodology

To date, thousands of reports on genomics, origin, genome evolution, molecular diagnosis and vaccine and/or therapeutics of SARS-CoV-2 have been published. However, a comprehensive review on microbial (virus, bacteria, fungus, archaea) co-infections associated with COVID-19 and the impact of co-infections on COVID-19 patients, characterization of co-pathogens, and their underlying molecular mechanisms in COVID-19 patients are lacking. Therefore, we conducted a rigorous literature survey on the co-infections, identifying of co-infecting microorganisms and their pathogenesis. The concept and evidence of co-infection with COVID-19 disease and a rationale of this comprehensive review are described in the introduction section. Later sections of this review were arranged coherently from the literature available in the PubMed central, Google Scholar, ResearchGate, bioRxiv, medRxiv, Preprints archives, World Health Organization (WHO) COVID-19 blog, National Institute of Health (NIH), Centers for Disease Control and Prevention (CDC, USA), Clinical Trials Registry databases, and COVID-19 vaccine and therapeutics tracker (https://biorender.com/covid-vaccine-tracker). The original research articles that discussed the evidence and significance of co-infections amid COVID-19, detection and possible molecular mechanisms of co-infections were considered for the content of this review. This literature survey also included case studies, case series and observational studies published from the very beginning of COVID-19 outbreak in Wuhan, China in late December 2019 to April 7, 2021. The literature search was done through screening of titles, abstracts and full articles for eligibility. Proposed molecular mechanisms of co-infections concurrent in SARS-CoV-2 infections have been represented in Fig. 1.

3. Microbial co-infections in hospitalized patients with COVID-19

Co-infection refers to the concurrent infection of a cell or host by two or multiple pathogen species and/or strains, whereas, superinfection is a scenario where one pathogen infects the host some time before infection by the second pathogen [31]. For both of these cases, the fate of the infected host often depends on a balance between the host’s protective immunity and immunopathology [32]. The universal pervasiveness or incidence of co-infection among humans is unknown, but it is thought to be commonplace, sometimes more common than single infection [33]. Co-infecting pathogens can alter the population of the primary
human diseases [19]. The COVID-19 causing SARS-CoV-2 strains show pathogen, as for example, Van der Hoek et al. (2004) reported that in respiratory co-infections, the human coronavirus (hCoV) load was much lower than for a single infection [34]. Co-infections may occur by multiple infectious agents of viral, bacterial, archaeal and fungal origin (Fig. 1), and appear to occur simultaneously with the initial onset of illness [35]. Co-infection morbidity has previously been studied within certain cohorts (e.g., age and sex), and is often reported to be worse than single infections [36]. Recently, several observational and cohort studies reported that pulmonary complications occurred in 51-2% COVID-19 patients, of which 82.6% accounted for deaths, and independent risk factors for mortality were male sex, age 65 years or older [37]. However, the occurrence of co-infection in death across age and sex cohorts of COVID-19 patients has not been studied yet. Co-infections in COVID-19 may not accounted for by age group, and unexplained heterogeneity in COVID-19 severity may be due to differences between studies in disease severity, patient comorbidities, treatment differences (such as corticosteroid administration), use of antibiotics prior to and during hospitalization, or other unidentified covariates. However, underestimation of secondary bacterial or fungal infections developing later in the course of COVID-19 is likely.

3.1. Viral co-infections in hospitalized patients with COVID-19

Co-infections with other viruses are very common in the viral infections of respiratory diseases. The prevalence of respiratory viral co-infection varies from 3.0% to 68.0% [20,38]. Several clinical studies indicated that viral co-infections of SARS-CoV-2 occurred with other virus from different countries [23,26]. Lin et al. (2020) reported that in Shenzhen Third People’s Hospital, 3.2% SARS-CoV-2 patients suffered from viral co-infections [20]. Association of other viruses, bacteria, fungi, with SARS-CoV-2 infection has been reported [13]. Bacteriophages are naturally occurring viruses that use bacteria as hosts, and play an extremely important part in allowing relatively harmless bacteria to become pathogens [19,39]. The bacteriophages are overlooked human pathogens that imply in triggering and worsening of a number of human diseases [19]. The COVID-19 causing SARS-CoV-2 strains show neighboring relationship to human classic coronavirus, the SARS coronavirus isolate Tor2 (SARS-CoV Tor2) corroborating with the recent findings [40]. In a recent metagenomic study, Hoque et al. (2020) reported that COVID-19 samples have sole association with 16 viral genera (other than betacoronavirus), and of them, Tombusvirus, Victorivirus, Partitivirus, Chrysovirus and Totivirus were the most abundant genera associated with SARS-CoV-2 co-infections [11].

3.1.1. SARS-CoV-2 and influenza: the disarray

Influenza is an acute and highly contagious respiratory disease that is responsible for significant morbidity and mortality worldwide. Annually, influenza can affect approximately 9% of the world’s population, with up to 1 billion infections, 3 to 5 million severe cases, and 0.3 to 0.65 million deaths [41]. The current COVID-19 pandemic caused by SARS-CoV-2 demonstrates similar symptoms of influenza such as fever, headache, sore throat and so on. Although, at the early stage of COVID-19 pandemic, the co-infection of SARS-CoV-2 and influenza did not draw any significant attention, the first report of influenza and SARS-CoV-2 co-infection [42] indicated that only 0.4% influenza positive patients were infected with SARS-CoV-2 in January 2020. The incidence of co-infection by influenza viruses in COVID-19 patients have been reported frequently from different countries other than China [27]. A case report of a 78-year old woman in Japan, showed the presence of Influenza A co-infection with SARS-CoV-2 [43] which intensified the necessity for specific and accurate detection of etiologic agents considering the travel history and medical condition of the patients. Another study showed 4.35% presence of influenza co-infection in hospitalized COVID-19 positive patients [44]. Although no anomalies in hematology screening were reported, all coinfected patients were clinically cured after treatment with oxygen inhalation, oseltamivir and antimicrobial agents without any invasive ventilator, intensive care unit (ICU) care and extracorporeal membrane oxygenation treatment [44]. In another retrospective study in Jiangsu Province in China demonstrated that 94.2% of laboratory-confirmed COVID-19 patients had co-infection, specifically 31% of them had viral co-infections with influenza virus [45]. In a recent retrospective cohort study on COVID-19 patients in a
hospital in Barcelona showed that only 0.4% patients were co-infected with community-acquired influenza A virus, whereas 3.1% patients infected with community-acquired bacterial infections, 4.7% of which were hospital-acquired superinfections [46]. However, the concomitant outbreak of influenza and COVID-19 in the winter season confirmed the presence of co-infection in multiple cases and inevitably emphasized the simultaneous laboratory diagnosis facilities for both SARS-CoV-2 and influenza viruses. The aphasymic symptoms of both COVID-19 and typical influenza have necessitated the co-diagnosis of influenza in COVID-positive patients, specifically with recent travel in influenza-endemic areas.

3.1.2. Hepatitis B co-infections in COVID-19 patients
Hepatic complications associated with COVID-19 are particularly concerning among people living with hepatitis B virus (HBV) co-infection with pre-existing liver complications (e.g., cirrhosis, liver failure, hepatocellular carcinoma) [47,48]. Co-infections among COVID-19 patients with hepatitis symptoms before developing the respiratory syndromes have been documented in several studies [49,50]. While our understanding of SARS-CoV-2’s pathogenesis continues to grow, initial studies suggest that the virus could lead to liver injury mainly by binding to angiotensin-converting enzyme 2 (ACE2) receptors on hepatocytes or causing an immune-mediated hepatic injury through activation of cytokine storm [51]. It is now clear that COVID-19 could lead to liver injuries and elevate alanine aminotransferase (ALT), aspartate aminotransferase (AST), and total bilirubin, particularly among the severe COVID-19 cases who need ICU [52–54]. Abnormal liver functions in COVID-19 patients have also been associated with increased disease severity and risk of mortality. Interestingly, viral co-infections caused by non-respiratory infectious agents have been reported from the Wuhan city in China, which stated the co-incidence of hepatitis B virus infections in 12.2% patients with acute COVID-19 [23]. The significant rate of liver cirrhosis and abnormally higher liver functions in severe COVID-19 patients corroborated with the findngs of co-infection with hepatitis B including elevated ALT, AST, gamma-glutamyl transferase (GGT) levels accompanied by moderately elevated prothrombin time (PT), and total bilirubin (TB) levels in COVID-19 patients [55,56]. The possibilities of liver impairment in severe COVID-19 patients have been suggested due to viral tropism to hepatic tissues, drug toxicity and systemic inflammation [54]. Therefore, extensive screening for hepatitis B infections in critical COVID-19 patients can be more useful for disease progression analysis, and effective treatment plan application in patients with remarkable diagnostic indications of abnormalities in liver functions.

3.1.3. SARS-CoV-2 and dengue: the deadly duo
The spread of SARS-CoV-2 in temperate countries such as Switzerland and France, where arboviral dengue fever endemic, has been described with a contemporary travel history [57]. In France, the first diagnostic test of SARS-CoV-2 RT-PCR revealed negative with a flu-like syndrome in patients, which became positive a week later with severe clinical onset of COVID-19 symptoms like fever, fatigue, loss of appetite and diarrhea [57]. The appearance of diffuse maculopapular exanthema made the clinicians to screen for Leptospira spp., Rift Valley fever virus, dengue virus and Chikungunya virus infections which indicated that RT-PCR for type-1 dengue virus was positive. Another report of two patients from Singapore revealed that rapid serological test for dengue can generate false positive results accelerating the respiratory complications in patients who later became positive for SARS-CoV-2 [58]. The initially worsening fever, increasing thrombocytopenia and sero-positivity for dengue misguided the clinicians who lately found the absence of dengue specific immunoglobulins, but with positive results from RT-PCR test of nasopharyngeal swab for SARS-CoV-2. In several countries of South America, a significant numbers of dengue cases were reported along with a gradual increase of COVID-19 cases. A study from Brazil depicted the possibility of under-reporting of dengue cases due to extensive mobilization of epidemiological sero-surveillance response team for COVID-19 emergency response, which may indirectly affected the reporting and treatment of dengue during COVID-19 outbreak [59]. That study urged the robust integrated national strategy for combined surveillance, treatment and prevention plan for dengue and COVID-19 management in Brazil. Another study from Columbia analyzed the dual epidemiological features of dengue and SARS-CoV-2 in the first 20 weeks of COVID-19 pandemic [60]. The viral interference resulting in blocking entry and replication of dengue during SARS-CoV-2 infection may also have contributed to the decreased onset of clinical dengue in subclinically or, mildly COVID-19 infected population [61]. Contemporary reports from Ecuador revealed that the eco-epidemiological dynamics and high endemic-epidemic transmission of dengue in large coastal areas can drastically affect the mitigation campaign and COVID-19 containment measures through extensive loads of patients on public health facilities for diagnosis, treatment and preventive actions [62]. These reports indicated the urgency of sero-surveillance for dengue virus infections in dwellers and travelers from temperate endemic areas. It has also been stated that the physical distancing and reduced public mobility may contribute to the containment of dengue infections amid COVID-19 [63].

3.1.4. COVID-19 patients with chronic viral diseases: HIV and HCV
The triple burden of COVID-19, human immunodeficiency virus (HIV) and hepatitis C virus (HCV) is one of the major and persistent global health challenges of the twenty-first century. The HIV, HCV and newly emerging infectious diseases such as coronavirus epidemics are expected to overlap in high HIV and HCV burden countries [64]. How COVID-19 will manifest itself in persons co-infected with HIV/HCV is still unclear [64]. Populations infected with HIV and HCV may be at elevated risk for severe responses if they are infected with COVID-19. In the future, lung lesions associated with COVID-19 may increase the risk of HIV or HCV, which induces a truly vicious circle of HIV-HCV-COVID-19 co-infections [64,65]. While COVID-19 continues to spread across the world, many areas face the risk of infection with SARS-CoV-2 and the obstacles and challenges to sustaining the continuum of HIV and HCV treatment in high-burden HIV/HCV countries are increasing [64]. In fact, the pathogenicity of COVID-19 could be accelerated in people living with HIV, who have compromised immunity [65]. Recent evidence has indicated a substantial association between coronavirus-related lower respiratory tract infections (LRTIs) and increased risk of death in immuno-compromised individuals [64]. Co-infections of HIV and SARS-CoV-2 in five individuals—three male and two transgender patients in Spain has been reported [32]. Interestingly, the HIV/SARS-CoV-2 patients had similar clinical, laboratory and radiographical features to the HIV-negative patients infected with SARS-CoV-2. During the current SARS-CoV-2 pandemic, this lack of information is a concern in countries with high HIV cases, especially in Sub-Saharan Africa, where 70% of people living with HIV infection [32]. As the patient had the history of co-infection with HIV-1 and HCV before 4 years, the follow-up study of anti-SARS-CoV-2 immune response revealed the delayed antibody response but with repeatedly negative RT-PCR test for SARS-CoV-2 RNA [66]. However, the compromised immune status of the patient caused the delayed humoral response development against the SARS-CoV-2, but the anti-HIV therapeutics and elevated level of activated IFN-γ due to anti-HIV agents may be suppressing SARS-CoV-2 infection leading to persistently positive SARS-CoV-2 RNA in RT-PCR tests [66]. Therefore, history of viral co-infection with immuno-compromised status and antiviral therapies may lead to delayed antibody response along with indistinct COVID-19 diagnosis.

3.1.5. SARS-CoV-2 with hCoV-HKU1
The endemic human coronaviruses (hCoVs) have been known to cause co-infections, sequential infections or can be co-detected with each other or with other respiratory viruses, including influenza A/B,
Respiratory Syncytial Virus (RSV), metapneumovirus, enterovirus, and adenovirus [67]. A critical case of co-infection by human coronavirus HKU1 in a COVID-19 patient, reported from Indonesia, indicated the sequential infections by hCoV-HKU1 and SARS-CoV-2, which was confirmed by a FilmArray Respiratory Panel (RP) test [67]. Thus, clinicians need to be aware of hCoV co-infections among COVID-19 patients. A high degree of suspicion in this rapidly evolving outbreak is required to make the diagnosis, and thereby, to contain and control the spread of the COVID-19.

3.2. Bacterial, archaeal and fungal co-infections in hospitalized patients with COVID-19

Like other well studied respiratory viral infections including the 1918 influenza outbreak [1], and 2009 H1N1 pandemic [68], the current pandemic outbreak of SARS-CoV-2 is also reported to associate with secondary microbial infections [11,12,69]. Bacterial co-infections were previously reported for respiratory diseases including SARS-CoV, MERS-CoV and influenza patients receiving intensive care [10,70]. Several retrospective studies showed that during the 1918 Spanish flu pandemic, bacterial pneumonia was a major cause of morbidity and mortality [71]. In a previous study, the prevalence of bacterial co-infections during the pandemic of influenza A (H1N1) between 2009 and 2012 was 23.0% [68]. Recent study reported that about 65% of laboratory-confirmed cases of influenza infection are known to be complicated by bacterial co-infections [72].

Bacterial co-infections develop in patients amid or after the primary infection initiated by an infectious agent. Bacterial co-infections also play a significant role during COVID-19 and are associated with an increasing rate of disease severity and case fatality [35]. Nevertheless until now, the incidence of bacterial co-infections in patients admitted to the ICU for acute respiratory failure associated with SARS-CoV-2 pneumonia is poorly studied [73,74]. SARS-CoV-2 primarily enters human body through nasopharyngeal tract and then gradually move to lung to initiate infection. The pathophysiology of COVID-19 can be attributed to aberrant immune responses in clearing the virus [75,76].

The most unequivocal association between viral and bacterial co-infection and respiratory disease severity, there is a pressing need to better understand how interactions of the SARS-CoV-2 with the host microbiome in the respiratory tracts correlate with viral infections that facilitate opportunistic co-infections [77]. However, little is known the outcome of the interactions of SARS-CoV-2 with nasal commensal bacteria which is thought to be critical for ultimate severity of COVID-19 disease development. The prevalence rate of bacterial co-infections in critically ill hospitalized COVID-19 patients was around 14% revealed by a meta-analysis [78]. A recent North American study reported 41% prevalence of bacterial co-infections among 17 COVID-19 patients admitted to ICU [79]. The incidence of co-infection associated with bacterial pneumonia ranged between 11% and 35% among the patients who had been infected with respiratory viruses [72]. Recently, Fu et al. (2020) reported that among ICU admitted COVID-19 patients, 13.9% were suffering from bacterial co-infections. Despite having a varying rate of bacterial co-infections among COVID-19 patients, the rate of prevalence could be as high as 50% among the non-survivors [80]. A series of retrospective case studies on SARS-CoV-2 confirmed that severely and non-severely ill patients had 7.7% and 3.2%, bacterial and fungal co-infections, respectively [54]. In Italy, a study conducted among 16,654 patients with critical condition, who died of SARS-CoV-2 infection depicted that 11% of those cases were associated with bacterial and fungal co-infections [81].

Co-infections with Streptococcus pneumoniae, Staphylococcus aureus or other colonizing bacteria during the pathophysiology of COVID-19 impairs both innate and adaptive antibacterial host defenses and temporarily compromise the physical and immunological barrier to cause secondary bacterial pneumonia, leading to severe and deadly disease in people with pre-existing comorbidities and previously healthy people [82]. Data regarding bacterial co-infections in COVID-19 pneumonia are still emerging, but an association has been made between the detection of bacterial pathogens in samples with disease severity in COVID-19 patients. The most commonly identified coinfected bacterial pathogens include, Acinetobacter baumannii, Klebsiella pneumoniae, Mycoplasma pneumoniae, Legionella pneumophila, Streptococcus pneumoniae, and Clamydia pneumoniae [11,70,83,84], while Aspergillus flavus, Candida glabrata, and Candida albicans are the most common coinfected fungi [78]. In addition, bacterial pathogens such as Staphylococcus aureus, Haemophilus influenzae, Klebsiella pneumoniae, Streptococcus pneumoniae, Neisseria meningitides [83,84] as well as some genera of Proteus, Enterobacter, and Citrobacter species have also been reported in hospitalized COVID-19 patients. In a recent microbiome study, Hoque et al. reported 527 and 306 bacterial genera in COVID-19 patients of Bangladesh and China, respectively [11]. Moreover, Pseudomonas aeruginosa and E. coli are the most frequently isolated multi-drug resistant (MDR) pathogens to be associated with hospital acquired superinfections [46]. Remarkably, SARS-CoV-2 RNA has also been detected in fecal samples of COVID-19 patients. It raises the question of gastrointestinal infection of SARS-CoV-2 and a possible fecal-oral route of disease transmission [85,86]. Moreover, high expression levels of ACE2 mRNA in the gastrointestinal system revealed a strong interaction of SARS-CoV-2 with the gastrointestinal system that has high microbiome diversity and possible chances of immune suppression and bacterial co-infections [85,86]. The role of SARS-CoV-2 in modulation of microbiome diversity in the gastrointestinal tracts is an important question for further research. The infection of SARS-CoV-2 of intestinal cells can lead to the change of intestinal microbiota. Gu et al. reported that the gut bacterial community of COVID-19 patients significantly reduced compared to those of healthy humans [87]. This dysbiosis (loss of beneficial commensals) was reflected by the significantly higher abundance of opportunistic pathogens such as Streptococcus, Rothia, Veillonella and Actinomyces in COVID-19 patients, while the relative of beneficial symbionsts abundance, such as Blautia, Romboutsia, Collinsella, and Bifidobacterium remained much lower [87].

A study on hospitalized COVID-19 patients with oropharyngeal candidiasis (OPC) showed that C. albicans was found to be the most prevalent pathogen, which was counted for 70.7%, followed by other fungi including C. glabrata (10.7%), C. dubliniensis (9.2%), C. tropicalis (3%), and C. krusei (1.5%) [88]. On the other hand, Chen et al. reported 5% prevalence of fungal co-infections in 99 COVID-19 patients in China, including one case of Aspergillus flavus, one case of Candida glabrata and three cases of C. albicans [23]. Yang et al. found that 5.8% (3/52) of the critical COVID-19 patients had fungal co-infections with A. fuscus, A. fumigatus and C. albicans [89]. In addition, 8–15% incidence of non-specific co-infections among COVID-19 patients were reported in different studies from China, but it is not clear whether it is bacterial or fungal infections [52,90]. A significant percentage of the SARS-CoV-2 infected patients developed co-infections associated with MDR typically from nosocomial pathogens [91]. An appreciable minority cases of superinfection, most commonly pneumonia and bacteremia can be developed due to MDR bacterial pathogens and fungus specially Aspergillus spp. Fungal originated co-infections including pulmonary aspergillosis and candidiasis were reported to complicate SARS-CoV-2 infection [46]. To date COVID-19-associated pulmonary aspergillosis (CAPA) has been documented in >30% of the cases [78].

Until now, most of the reported respiratory tract co-infections are limited to viral, bacterial, and fungal pathogens [57,92], while a plethora of other concomitant microbial components including archaea could also be found [11,93]. Unlike bacteria, the incidence, diversity and composition of these co-pathogens always remain much lower compared to the infectious agent of COVID-19. Metagenomic investigations confirmed presence of Methanosarcina, Methanocaldolectococcus, Thermococcus, Methanothermobacter, Haloarcula, Staphylothermus, Natronomonas, Ferrolobus, Caldivirga, Halobacterium, Natrilalba, Methanosphaerula and Picrophilus as the archaeal genera in samples of...
COVID-19 patients [11,93].

4. Molecular mechanism of co-infection in COVID-19

Co-infections can augment the pathogenesis, morbidity and mortality in most of the respiratory viral diseases [35]. Table 1 discusses the commonly reported microbial co-pathogens amid COVID-19, their transmission pattern, possible mechanism of co-infections and outcomes. Co-infections in COVID-19 patients may also complicate the clinical outcomes of the disease. The SARS-CoV-2 enters human cells by binding to the ACE2 protein of the cells lining the upper and lower airways. Recently, Lee et al. reported that the ACE2 receptor protein robustly localizes within the motile cilia of airway epithelial cells, which likely represents the initial or early subcellular site of SARS-CoV-2 viral entry during host respiratory transmission [94]. However, the ciliary ACE2 expression in the upper airway is influenced by patient demographics (age, sex and smoking), clinical characteristics, comorbidities/co-infections or medication use [94]. Remarkably, specific molecular kinetics of these additional infections in COVID-19 patients are still remained unclear although a few studies proposed some models for the co/superinfections in COVID-19 patients in different countries [26]. Based on available literature, we propose a plausible mechanism of co-infection in COVID-19 patients (Fig. 1, Table 1).

Respiratory viruses are frequently collaborated by secondary bacterial infections due to the outgrowth of opportunistic bacterial pathogens. Although the specific molecular mechanisms of co-infections in COVID-19 patients remain unclear, it may include virus-induced airway damage, cell loss, goblet cell hyperplasia, altered mucus secretion, reduced ciliary beat frequency, function and clearance, reduced oxygen exchange, and damage to the immune system [27,45,95]. Co-infection increases the levels of C-reactive protein (CRP) and procalcitonin (PCT) [5]. Viral infections damage the respiratory airway both histologically and functionally [69,96]. The co-infection mechanisms include virus-induced airway damage, cell loss, goblet cell hyperplasia, altered mucus secretion, reduced ciliary beat frequency, reduced mucociliary clearance, dis-coordinated mucociliary functions, reduced oxygen exchange, and damage to the immune system [69,96,97]. Viral co-infection can also facilitate bacterial adhesion, disrupt the tight junction and epithelial barrier integrity favoring paracellular transmigration of bacteria, and alter both innate and adaptive immune responses that render the lung more vulnerable to SARS-CoV-2 infections [20,38]. Many viruses can destroy the airway epithelium which facilitates other viral co-infection [98]. The COVID-19 patient having co-infected with HIV had a longer progression of the disease and slower generation of specific antibody because of the collapse of immune system [99]. SARS-CoV-2 infection may cause liver damage [5], and thus, drug-induced liver injury (DILI) is more likely to occur in patients who already have certain viral infections including HCV and HIV [23]. Therefore, the development and outcome of SARS-CoV-2 associated co-infections with other viruses are highly dependent on the host immune response, especially in the elderly [23]. The co-infection of viruses is associated with different molecular mechanisms by which the pre-disposition of the virus occurs in the respiratory tract that promotes simultaneous bacterial infection. The epithelial cells of the respiratory tract help bacterial adherence using different mechanisms during viral infection, while the disease severity varies upon virus, bacterial strain, other co-pathogens and hosts immunity. Respiratory viruses can up-regulate the expression of host cell membrane protein to facilitate their binding [69]. Respiratory Syncytial Virus (RSV) reported to bind directly with Haemophilus influenzae and Staphylococcus pneumoniae, and thus, favoring bacterial proximity to the epithelial monolayer and supplementing attachment to the host cell receptors. Moreover, the expression and localization of the RSV glycoprotein on the host cell membrane during infection can further act as bacterial receptors for pneumococcal binding [100]. Previous studies demonstrated that influenza virus can make mice susceptible to pneumonia caused by S. aureus where both virus and bacterial load increased during co-infection [100,101]. Several respiratory viruses such as RSV, parainfluenzavirus-3, and influenza viruses reported to increase the bacterial adherence upon infection, in both primary and immortalized epithelial cells [69]. The surface glycoprotein adhesion molecule-1 (ICAM-1) expression is upregulated during RSV and adenovirus infection, and thereby, increases adherence for S. pneumoniae in human nasopharyngeal cells (HeP-2) and pneumocyte type II cells (A549). The enhanced pneumococcal adherence in epithelial cells results in bacterial accumulation which may facilitate other bacterial co-infections [69,102]. Methanogenic archaea coexist and interact closely with anaerobic bacteria [39]. Methanogenic archaea utilize low molecular weight compounds, such as H2 + CO2, formic acid, or acetate, and therefore, have symbiotic relationships with the producers of these substrates. It is reasonable to assume that the presence or increase in level of methanogenic archaea modulates the composition of the polymicrobial community and changes the virulence property of the microflora [39]. Although, several earlier studies stated that archaeal co-infection is frequently detected in viral and bacterial hosts [92,105], the systematic tests of the factors explaining variation in viral co-infection across different taxa and environments are still lacking.

Recent studies suggested that COVID-19 patients having microbial co-infections are characterized by lymphopenia and enhanced levels of proinflammatory cytokines including interleukin-6 (IL-6) and IL-1β as well as MCP-1, IP-10, and granulocyte colony-stimulating factor (G-CSF) in the plasma. It has been proposed that high levels of proinflammatory cytokines might lead to shock as well as respiratory failure or multiple organ failure, and several trials to assess inflammatory mediators are under way [104]. Cytokine storm, or hypercytokinemia, describes hyperactivation of the immune system that may be provoked or worsened by co-infections. This can lead to devastating and irreversible destruction of lung tissue as proinflammatory cytokines damage the alveoli, tiny sacs in the lungs responsible for gas exchange and oxygenation [51]. Damage to lung tissues caused by SARS-CoV-2 may explore the receptors for other pathogenic or opportunistic microorganisms facilitating secondary infections. Specific domains in which viruses play such facilitating role including enhancement of bacterial adhesion by unmasking cryptic receptors and upregulation of adhesion proteins, disruption of tight junction integrity favoring paracellular transmission of bacteria and loss of epithelial barrier integrity, increased availability of nutrients, such as mucins and iron, alteration of innate and adaptive immune responses, and disabling defense against bacteria, and lastly, changes in airway microbiome that render the lungs more vulnerable to pathogens [105]. Moreover, SARS-CoV-2 infection can damage lymphocytes, especially B cells, T cells, and NK cells, which will lead to the immune system’s impairment during the period of disease [26]. The decrease of lymphocytes and host immune function may be the main reason for further super- or secondary infection [106]. The mortality is more significant in severe cases compared with the non-severe group [107] due to the higher co-infection rate in severe patients [106,107]. The mechanistic evidence of influenza and pneumococcal co-infections showed that influenza virus causes a depletion of alveolar macrophage which allows a smaller inoculum of bacteria to establish productive infection [106], and consequent severity of the bacterial co-infection is exacerbated by preexisting host factors such as obesity [109]. For more severely ill patients, they are more likely to receive treatment with invasive catheters, resulting in increased sensitivity of co-infections with multidrug-resistant pathogens such as Acinetobacter baumannii, Escherichia coli, Pseudomonas aeruginosa, and Enterococcus species [110]. In addition, during the pathophysiology of COVID-19, SARS-CoV-2 viruses can interact with a large number of cellular proteins (virus-host interactome) and protein-protein interactions between unrelated viruses, bacteria, archaea and fungi are also possible [110]. Co-infections may result in genetic exchange among heterologous viruses, and/or agents [111] leading to the generation of recombinant or chimeric pathogens, and this recombination effects can
### Table 1
Commonly reported microbial co-pathogens amid COVID-19, their transmission pattern along with the possible mechanism of co-infections and outcomes.

| Type of co-infection | Co-pathogens | Route of transmission | Person to person transmission | Possible mechanism of co-infection and pathogenesis | Possible outcomes |
|----------------------|--------------|-----------------------|-----------------------------|---------------------------------------------------|-------------------|
| **Viral**            | Influenza    | Respiratory           | Yes                         | IFN induced overexpression of ACE2 triggered by influenza virus aids SARS-CoV-2 infection [112]. | Severe respiratory distress followed by pleural effusion and necrotizing pneumonia [128], higher mortality rate [129]. |
|                      | HBV          | Body fluid            | Yes                         | Increased liver tissue damage and inflammatory responses due to COVID-19 may aid HBV co-infection by overexpressing host cell receptors [114]. It may also fuel the reactivation of pre-existing chronic HBV [115]. | HBV patients under ART exhibits mild COVID-19 symptoms. But ART-naive patients show acute COVID-19 clinical presentation [121]. Higher maximum body temperatures, longer duration of fever and longer improvement time of chest CT image was reported due to co-infection [122]. |
|                      | Dengue       | Mosquito bite         | No                          | NR | Increase the severity of symptoms [118]. Decrease in white blood cell, neutrophils, lymphocytes and platelets count and eventual higher mortality rate [119]. |
|                      | HIV          | Body fluid            | Yes                         | Suppression of T lymphocyte mediated immunity (as observed in HIV patients) leads to the prognosis of increased disease severity and higher mortality rate during COVID-19 co-infection [126]. | The actual outcome is not reported till date. It has been speculated that some investigational COVID-19 drugs may adversely affect the HCV-related decompensated cirrhosis patients [124]. |
|                      | HCV          | Body fluid            | Yes                         | Both SARS-CoV-2 E and HCV p7 proteins can form similar ion channels which ensure their success in attacking their host and effective replication during co-infection [123]. | One case has been reported in a young patient expressing critical illness as the outcome of co-infection [126]. |
|                      | Rhinovirus   | Respiratory           | Yes                         | Major disease-causing rhinovirus serotype HRV-A16 infection upregulates ACE2 and TMPRSS2 expression in epithelial cells by inducing IFNb1. This event facilitates SARS-CoV-2 transmission and further disease severity [125]. | Unfavorable prognostic outcome including ARDS [127]. |
|                      | Adenovirus   | Respiratory           | Yes                         | Similar ion channel forming capability of SARS-CoV-2 E and Adenovirus 6K proteins facilitates co-infection [123] | |
| **Bacterial**        | Streptococcus pneumoniae | Respiratory | Yes | Opportunistic normal flora of human upper respiratory tract | Severe respiratory distress followed by pleural effusion and necrotizing pneumonia [128], higher mortality rate [129]. |
|                      | Staphylococcus aureus | Respiratory/Digestive/Contact | Yes | Opportunistic normal flora of human upper respiratory tract, gut mucosa and skin | Necrotizing pneumonia [139]. |
|                      | Pseudomonas aeruginosa | Contact | Yes | Opportunistic pathogen causing HAI mostly related with poor hygiene, mechanical ventilation and urinary catheterization. | Bacteremia and higher mortality [131]. |
|                      | Acinetobacter baumannii | Contact | Yes | NR | |
|                      | Klebsiella pneumoniae | Respiratory/Contact | Yes | Opportunistic normal flora of human mouth, skin, and intestines | Fatal sepsis [132]. |
|                      | Mycoplasma pneumoniae | Respiratory/Contact | Yes | NR | Severe pneumonia [133]. Increased morbidity, mortality and disease severity [134] |
|                      | Clamydia pneumoniae | Respiratory/Contact | Yes | NR | Severe pneumonia [135]. |
|                      | Legionella pneumophila | Digestive/Respiratory | Yes | NR | Elevated aspartate aminotransferase, blood urea nitrogen, creatinine, lactate dehydrogenase and C-reactive protein [135]. |
|                      | Haemophilus influenzae | Respiratory/contact | Yes | Opportunistic normal flora of human upper respiratory tract | |
|                      | Neisseria meningitides | Respiratory/contact | Yes | NR | Convulsion [136], elevated C-reactive protein, headache, neck stiffness, rigor, confusion, and a new purpuric rash over hands and feet [137] |
|                      | Mycobacterium tuberculosis | Respiratory | Yes | Cytokine storm produced by COVID-19 may reactivate latent TB or boost the development of active TB. Lung damages caused by TB may also escalate the disease severity caused by SARS-CoV-2 [138]. | Co-infection is associated with disease severity and disease progression rate [139]. 2.17 times higher risk-of-death and 26% lower risk-of-recovery was reported. Also shorter time-to-death and longer time-to-recovery was found [140]. |
|                      | Aspergillus spp. | Respiratory | No | Pro-inflammatory cytokines (especially IL-6 and IL-10) released during COVID-19 can result in tissue necrosis and ARDS, which eventually makes patient more vulnerable to Aspergillosis [141]. | Invasive pulmonary aspergillosis, higher case fatality rate (64.7% reported) [141] |
|                      | Candida spp. | Perinatal/Contact | No | Opportunistic pathogen found in human skin. | Candidemia and increased mortality rate [142]. |

IFN: Interferon; ACE2: Angiotensin-converting enzyme 2; SARS-CoV-2: Severe Acute Respiratory Syndrome Coronavirus 2; COVID-19: Coronavirus disease 2019; HBV: Hepatitis B Virus; HIV: Human Immunodeficiency Virus; HCV: Hepatitis C Virus; ALT: Alanine transaminase; AST: Aspartate transaminase; TBIL: Total bilirubin; ALP: Alkaline phosphatase; γ-GT: Gamma-glutamyl transferase; ART: Antiretroviral therapy; CT: Computed Tomography; HRV-A16: Human rhinovirus A16; TMPRSS2: Transmembrane protease, serine 2; IFNb1: Interferon Beta 1; ARDS: Acute respiratory distress syndrome; HAI: Hospital Acquired Infections; TB: Tuberculosis; IL-6: Interleukin 6; IL-10: Interleukin 10; NR: Not Reported.
influence viral evolution, disease dynamics, sensitivity to antiviral therapy, and eventually the fate of the host [110] Though not studied yet, similar mechanistic events may be found in SARS-CoV-2 and associated co-infections aggravating the pathophysiology of COVID-19. In addition, four factors including host ecology, host taxonomy or phylogeny, host defense mechanisms, and the interactions of co-pathogens with SARS-CoV-2 are likely to play vital role in the pathophysiology and severity of COVID-19 disease. The relevance and importance of these molecular events of pathogenesis are likely to vary for cross-infectivity, culture co-infection, and single-cell co-infection [92].

Overall, due to some risk factors including the epithelial lung damage, immune system dysregulation, prolonged period of hospitalization etc., the possible development of superinfections is somewhat expected in severely ill COVID-19 patients [78]. However, the actual scenario of the possible development of superinfections is somewhat expected in COVID-19 is a crucial step in the design of rational therapeutic strategies. Therefore, the effect of SARS-CoV-2 replication and induction of innate immune response on the composition of the human or animal upper respiratory tract (URT) microbiome remains to be elucidated and analyzed in depth on a community wide scale. Further extensive investigation is warranted for a better understanding and evaluating the risk factors associated and the disease spectrum of co-secondary and superinfection in critically ill patients suffering from SARS-CoV-2 infection.

5. Conclusion and perspectives

Co-infections with various microorganisms are commonly found in SARS-CoV-2 infected patients that significantly influences the severity and mortality rates of COVID-19. However, our understanding about co-infecting organisms, their cross-talks and ultimate interactions with the hosts are poor. The COVID-19 co-infections are associated with multiple domains of microorganisms including viruses, bacteria, fungi and archaea. Although the specific molecular events of co-pathogenesis in SARS-CoV-2 pathophysiology is yet unknown, the co-infecting pathogens may participate to damage the respiratory airway, cell loss, goblet cell hyperplasia, alter mucus secretion, reduced ciliary beat frequency, function and clearance, reduced oxygen exchange, and damage the immune system. Furthermore, viral co-infection facilitates bacterial adhesion, disrupt the tight junction and epithelial barrier integrity favoring paracellular transmigration of bacteria, and alter both innate and adaptive immune responses that render the lung more vulnerable to SARS-CoV-2 infections. Although, this review provides a comprehensive scenario of co-infection in COVID-19 patients, further studies are needed to focus the epidemiology, clinical and laboratory characteristics of co-infecting pathogens among diverse group of COVID-19 patients from different geo-climatic conditions, and assess the effect of co-infecting microorganisms on the outcome of COVID-19 patients.

Declaration of competing interest

The authors declare no conflict of interests.

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