Genomic and Viral Structure of SARS-CoV-2 in the Context of Pathogenesis, Symptomology, and Transmission

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## Abstract

The novel coronavirus SARS-CoV-2, which emerged in late 2019, has been a major force shaping the year 2020 as it spread around the world infecting tens of millions of people with coronavirus disease 2019 (COVID-19). While the viral species was unknown prior to January 2020, its similarity to other coronaviruses that infect humans has allowed for rapid insight into the mechanisms that it uses to infect humans, as well as the ways in which the human immune system can respond. Here, we contextualize SARS-CoV-2 among other coronaviruses and identify what is known and what can be inferred about its behavior once inside a human host. Because the genomic content of coronaviruses, which specifies the virus’s structure, is highly conserved, early genomic analysis provided a significant head start in predicting viral pathogenesis. The pathogenesis of the virus offers insights into symptomology, transmission, and individual susceptibility. Additionally, prior research into interactions between the human immune system and coronaviruses identified how these viruses can evade the immune system’s protective mechanisms. We also explore systems-level research into the regulatory and proteomic effects of SARS-CoV-2 infection and the immune response. Understanding the structure and behavior of the virus serves to contextualize the many facets of the COVID-19 pandemic and can influence efforts to control the virus and treat the disease.

## Importance

COVID-19 involves a number of organ systems and can present with a wide range of symptoms. However, understanding how the virus infects epithelial cells can contextualize how these systems connect. Similarly, the modes of viral transmission have been under debate throughout much of 2020, yet the available research suggests that these patterns are very similar to those observed in closely related viruses like SARS-CoV-1 and possibly MERS-CoV. Exploring the structure, phylogeny, and pathogenesis of the virus therefore helps to guide interpretation of the broader impacts of the virus on the human body and on human populations. For this reason, an in-depth exploration of viral mechanisms is critical to a robust understanding of the COVID-19 pandemic.

## Introduction

The current coronavirus disease 2019 (COVID-19) pandemic, caused by the *Severe acute respiratory syndrome-related coronavirus 2* (SARS-CoV-2) virus, represents an acute global health crisis. Symptoms of the disease can range from mild to severe to fatal [[1](#ref-gHWlMufv)] and can affect a variety of organs and systems [[2](#ref-DkrkT4Mb)]. Outcomes of the disease can include acute respiratory distress (ARDS) and acute lung injury, as well as damage to other organ systems [[2](#ref-DkrkT4Mb),[3](#ref-1H0HgI6iy)]. Understanding the progression of the disease, including these diverse symptoms, depends on understanding how the virus interacts with the host. Additionally, the fundamental biology of the virus can provide insights into how it is transmitted among people, which can, in turn, inform efforts to control its spread. As a result, a thorough understanding of the pathogenesis of SARS-CoV-2 is a critical foundation on which to build an understanding of COVID-19 and the pandemic as a whole.

The rapid identification and release of the genomic sequence of the virus in January 2020 [[4](#ref-Bp847Lfa)] provided early insight into the virus in a comparative genomic context. The viral genomic sequence clusters with known coronaviruses (order *Nidovirales*, family *Coronaviridae*, subfamily *Orthocoronavirinae*). Phylogenetic analysis of the coronaviruses reveals four major subclades, each corresponding to a genus: the alpha, beta, delta, and gamma coronaviruses. Among them, alpha- and betacoronaviruses infect mammalian species, gammacoronaviruses infect avian species, and deltacoronaviruses infect both mammalian and avian species [[5](#ref-17DSmRo9H)]. The novel virus now known as SARS-CoV-2 was identified as betacoronavirus belonging to the B lineage based on phylogenetic analysis of a polymerase chain reaction (PCR) amplicon fragment from five patients along with the full genomic sequence [[6](#ref-IFLf8rsY)]. This lineage also includes the *Severe acute respiratory syndrome-related coronavirus* (SARS-CoV-1) that caused the 2002-2003 outbreak of Severe Acute Respiratory Syndrome (SARS) in humans [[6](#ref-IFLf8rsY)]. Because viral structure and mechanisms of pathogenicity are highly conserved within the order, this phylogenetic analysis provides a basis for forming hypotheses about how the virus interacts with hosts, including which tissues, organs, and systems would be most susceptible to SARS-CoV-2 infection. Coronaviruses that infect humans (HCoV) are not common, but prior research into other HCoV such as SARS-CoV-1 and *Middle East respiratory syndrome-related coronavirus* (MERS-CoV), as well as other viruses infecting humans such as a variety of influenza species, established a strong foundation that accelerated the pace of SARS-CoV-2 research.

Coronaviruses are large viruses that can be identified by their distinctive “crown-like” shape (Figure [1](#fig:genome-structure)). Their spherical virions are made from lipid envelopes ranging from 100 to 160 nanometers (nm) in which peplomers of two to three spike (S) glycoproteins are anchored, creating the crown [[7](#ref-RGAUx68v),[8](#ref-DlBAegL1)]. These spikes, which are critical to both viral pathogenesis and to the response by the host immune response, have been visualized using cryo-electron microscopy [[9](#ref-CqQVkaqj)]. Because they induce the human immune response, they are also the target of many proposed therapeutic agents. Viral pathogenesis is typically broken down into three major components: entry, replication, and spread [[10](#ref-jsMmdH4D)]. However, in order to draw a more complete picture of pathogenesis, it is also necessary to examine how infection manifests clinically, identify systems-level interactions between the virus and the human body, and consider the possible effects of variation or evolutionary change on pathogenesis and virulence. In this way, both biomedicine and genomics are important pieces of the puzzle of SARS-CoV-2 presentation and pathogenesis.

## Coronavirus Structure and Pathogenesis

### Structure of Coronaviruses

Genome structure is highly conserved among coronaviruses, meaning that the relationship between the SARS-CoV-2 genome and its pathogenesis can be inferred from prior research in related viral species. The genomes of viruses in the *Nidovirales* order share several fundamental characteristics. They are non-segmented, which means the viral genome is contained in a single capsid, and are enveloped, which means that the genome and capsid are encased by a lipid bilayer. Coronaviruses have large positive-sense RNA (ssRNA+) genomes ranging from 27 to 32 kilobases in length [[11](#ref-47hnjvDb),[12](#ref-1Fqilxaum)]. The SARS-CoV-2 genome lies in the middle of this range at 29,903 bp [[12](#ref-1Fqilxaum)]. Genome organization is highly conserved within the order [[11](#ref-47hnjvDb)]. There are three major regions: one containing the replicase gene and one containing the genes encoding structural proteins [[11](#ref-47hnjvDb)] (Figure [1](#fig:genome-structure)). The replicase gene comprises about two-thirds of the genome and consists of two open reading frames that are translated with ribosomal frameshifting [[11](#ref-47hnjvDb)]. This polypeptide is then translated into 16 non-structural proteins (nsp), except in Gammacoronaviruses where nsp1 is absent, that form replication machinery used to synthesize viral RNA [[13](#ref-18GflyMj)]. The remaining third of the genome encodes structural proteins, including the spike (S), membrane (M), envelope (E), and nucleocapsid (N) proteins. Additional accessory genes are sometimes present between these two regions, depending on the species or strain. Much attention has been focused on the S protein, which is a critical structure involved in cell entry.



Figure 1: **Structure of SARS-CoV-2 capsid and genome.** A) The genomic structure of coronaviruses is highly conserved, including the order and organization of genes such as the spike (S), envelope (E), membrane (M) and nucleocapsid (N) proteins. B) The physical structure of the coronavirus virion, including the components encoded by conserved genes S, E, M and N.

### Pathogenic Mechanisms of Coronaviruses

While, like most viruses, it is possible that SARS-CoV-1 and SARS-CoV-2 can enter cells through endocytosis, coronaviruses are able to target cells for entry through fusion with the plasma membrane [[14](#ref-FFwktXgx),[15](#ref-FNetQkRI)]. This process is conserved among coronaviruses and is closely associated with the content of their genomes. Cell entry proceeds in three steps: binding, cleavage, and fusion. First, the viral spike protein binds to a host cell via a recognized receptor. Coronaviruses can bind to a range of host receptors [[16](#ref-13wWdgODZ),[17](#ref-OVsxrEuX)], with binding conserved only at the genus level [[5](#ref-17DSmRo9H)]. Viruses in the betacoronavirus genus, to which SARS-CoV-2 belongs, are known to bind to the CEACAM1 protein, 5-N-acetyl-9-O-acetyl neuraminic acid (Neu 5,9 Ac2), and to the angiotensin-converting enzyme 2 (ACE2) [[16](#ref-13wWdgODZ)]. SARS-CoV-2 has a high affinity for the human ACE2 receptor, which is expressed in the vascular epithelium, other epithelial cells, and cardiovascular and renal tissues [[18](#ref-So6ceUwB),[19](#ref-NodtSApl)]. The binding process is guided by the molecular structure of the spike protein, which is structured in three segments: an ectodomain, a transmembrane anchor, and an intracellular tail [[20](#ref-skHKTFMe)]. The ectodomain forms the crown-like structures on the viral membrane and contains two subdomains known as the S1 and S2 subunits [[21](#ref-ueMPuUMe)]. The S1 (N-terminal) domain forms the head of the crown and contains the receptor binding motif, and the S2 (C-terminal) domain forms the stalk that supports the head [[21](#ref-ueMPuUMe)]. The S1 subunit guides the binding of the virus to a host cell receptor, and the S2 subunit guides the fusion process [[20](#ref-skHKTFMe)].

After the binding of the S1 subunit to a receptor, the spike protein is often cleaved at the S1-S2 boundary by a host protease [[22](#ref-qcVbT0w4),[23](#ref-kCVy8vjj),[24](#ref-YeKaMzdJ)]. Similar to SARS-CoV, SARS-CoV-2 exhibits redundancy in which host proteases can cleave the S protein [[25](#ref-15l3di3Wj)]. Specifically, both transmembrane protease serine protease-2 (TMPRSS2) and cathepsins B/L have been shown to mediate SARS-CoV-2 S protein proteolytic priming, and small molecule inhibition of these enzymes fully inhibited viral entry *in vitro* [[25](#ref-15l3di3Wj),[26](#ref-JOJ2n3gC)]. Proteolytic priming prepares the S protein for fusion [[23](#ref-kCVy8vjj),[24](#ref-YeKaMzdJ)]. The two subunits remain bound by van der Waals forces, with the S1 subunit stabilizing the S2 subunit during the membrane fusion process [[22](#ref-qcVbT0w4)]. Electron microscopy suggests that in some coronaviruses, including SARS-CoV-1 and MERS-CoV, a six-helix bundle separates the two subunits in the postfusion conformation, and the unusual length of this bundle facilitates membrane fusion through the release of additional energy [[5](#ref-17DSmRo9H)]. Cleavage at a second site within S2 by these same proteases activates *S* for fusion by inducing conformational changes [[22](#ref-qcVbT0w4)]. The viral membrane can then fuse with the endosomal membrane to release the viral genome into the host cytoplasm. Once the virus enters a host cell, the replicase gene is translated and assembled into the viral replicase complex. This complex then synthesizes the double-stranded RNA (dsRNA) genome from the genomic ssRNA(+). The dsRNA genome is transcribed and replicated to create viral mRNAs and new ssRNA(+) genomes [[11](#ref-47hnjvDb),[27](#ref-ZUuGRxk9)]. From there, the virus can spread into other cells. In this way, the genome of SARS-CoV-2 provides insight into the pathogenic behavior of the virus.

### Immune Evasion Strategies

Research in other HCoV provides some indication of how SARS-CoV-2 infection proceeds in spite of the human immune response. By infecting the epithelium, viruses such as SARS-CoV-1 are known to bypass the physical barriers such as skin and mucus that comprise the immune system’s first line of defense [[28](#ref-1XbF4GLn)]. Once the virus infiltrates host cells, it is adept at evading detection. CD163+ and CD68+ macrophage cells are especially crucial for the establishment of SARS-CoV-1 in the body [[28](#ref-1XbF4GLn)]. These cells most likely serve as viral reservoirs that help shield SARS-CoV-1 from the innate immune response. According to a study on the viral dissemination of SARS-CoV-1 in Chinese macaques, viral RNA could be detected in some monocytes throughout the process of differentiation into dendritic cells [[28](#ref-1XbF4GLn)]. Thus lack of active viral replication allows SARS-CoV-1 to escape innate immunity because reduced levels of detectable viral RNA allow the virus to avoid both natural killer (NK) cells and Toll-like receptors [[28](#ref-1XbF4GLn)]. Even during replication, SARS-CoV-1 is able to mask its dsRNA from detection by the immune system. Although dsRNA is a pathogen-associated molecular pattern (PAMP) that would typically initiate a response from the innate immune system [[29](#ref-159GOVLV1)], *in vitro* analysis of nidoviruses including SARS-CoV-1 suggests that these viruses can induce the development of double-membrane vesicles that protect the dsRNA signature from being detected by the host immune system [[30](#ref-yQO4HFTZ)]. This protective envelope can therefore insulate these coronaviruses from the innate immune response’s detection mechanism [[31](#ref-OlEbK4fc)].

HCoV are also known to interfere with the host immune response, rather than just evade it. For example, the virulence of SARS-CoV is increased by nsp1, which can suppress host gene expression by stalling mRNA translation and inducing endonucleolytic cleavage and mRNA degradation [[32](#ref-Q5Q9k6Nc)]. SARS-CoV-1 also evades the immune response by interfering with type I IFN induction signaling, which is a mechanism that leads to cellular resistance to viral infections. SARS-CoV-1 employs methods such as ubiquitination and degradation of RNA sensor adaptor molecules MAVS and TRAF3/6 [[33](#ref-G6A0wWdn)]. Also, MERS-CoV downregulates antigen presentation via MHC class I and MHC class II, which leads to a reduction in T cell activation [[33](#ref-G6A0wWdn)]. These evasion mechanisms, in turn, can lead to systemic infection. Coronaviruses such as SARS-CoV-1 are also able to evade the humoral immune response through other mechanisms, such as inhibiting certain cytokine pathways or down-regulating antigen presentation by the cells [[30](#ref-yQO4HFTZ)].

### Host Cell Susceptibility

ACE2 and TMPRSS2 have been identified as the primary entry portal and as a critical protease, respectively, in facilitating the entry of SARS-CoV-1 and SARS-CoV-2 into a target cell [[9](#ref-CqQVkaqj),[25](#ref-15l3di3Wj),[34](#ref-2QTH37Xi),[35](#ref-MXWaV7SO),[36](#ref-JjrO6YbD)]. This finding has led to a hypothesized role for ACE2 and TMPRSS2 expression in determining which cells, tissues, and organs are most likely to be infected by SARS-CoV-2. The ACE2 receptor is expressed in numerous organs, such as the heart, kidney, and intestine, but it is most prominently expressed in alveolar epithelial cells; this pattern of expression is expected to contribute to the virus’ association with lung pathology [[18](#ref-So6ceUwB),[37](#ref-UwVweB2M)]. Clinical investigations of COVID-19 patients have detected SARS-CoV-2 transcripts in bronchoalveolar lavage fluid (BALF) (93% of specimens), sputum (72%), nasal swabs (63%), fibrobronchoscopy brush biopsies (46%), pharyngeal swabs (32%), feces (29%) and blood (1%) [[38](#ref-vr7AH83b)]. Two studies reported that SARS-CoV-2 could not be detected in the urine specimens [[38](#ref-vr7AH83b),[39](#ref-azgJqujy)]; however, a third study identified four urine samples (out of 58) that were positive for SARS-CoV-2 nucleic acids [[40](#ref-F7nSMvZk)]. Although respiratory failure remains the leading cause of death for COVID-19 patients [[41](#ref-1AJFJxzmJ)], SARS-CoV-2 infection can damage many other organ systems including the heart [[42](#ref-5gnCuPzp)], kidneys [[43](#ref-5ET1D3cK),[44](#ref-w5B6qRKv)], liver [[45](#ref-TTSFlLVC)], and gastrointestinal tract [[46](#ref-PmE9xedP),[47](#ref-3Ak4Mata)]. As it becomes clear that SARS-CoV-2 infection can damage multiple organs, the scientific community is pursuing multiple avenues of investigation in order to build a consensus about how the virus affects the human body.

## Clinical Presentation of COVID-19

Pathogenesis is closely linked with the clinical presentation of the disease caused by the virus. Reports have described diverse symptom profiles associated with COVID-19. Differences in the frequency of symptoms are found when comparing both between institutions in similar locations and between different regions. A large study from Wuhan, China conducted early in the pandemic identified fever and cough as the two most common symptoms that patients reported at hospital admission [[48](#ref-15IDA5kX6)], while a retrospective study in China described the clinical presentations of patients infected with SARS-CoV-2 as including lower respiratory tract infection with fever, dry cough, and dyspnea [[49](#ref-10THxyeCg)]. This study [[49](#ref-10THxyeCg)] noted that upper respiratory tract symptoms were less common, suggesting that the virus preferentially targets cells located in the lower respiratory tract. However, data from the New York City region [[50](#ref-19ytsiSpq),[51](#ref-1H8gSsSkn)] revealed variable rates of fever as a presenting symptom, suggesting that symptoms may not be consistent across samples. For example, even within New York City, one study [[50](#ref-19ytsiSpq)] identified low oxygen saturation (<90% without the use of supplemental oxygen or ventilation support) in a significant percentage of patients upon presentation, while another study [[51](#ref-1H8gSsSkn)] reported cough, fever, and dyspnea as the most common presenting symptoms. The variability of both which symptoms present and their severity has presented challenges for public health agencies to provide clear recommendations for citizens regarding what symptoms indicate SARS-CoV-2 infection and should prompt isolation. Patients may also experience loss of smell, myalgias (muscle aches), fatigue, or headache. Gastrointestinal symptoms can also present [[52](#ref-18tT1tLJI)], and the CDC includes nausea and vomiting, as well congestion and runny nose, on its list of symptoms consistent with COVID-19 [[1](#ref-gHWlMufv)]. A recent preprint using data from an app-based survey of 500,000 individuals in the US found that among those tested for SARS-CoV-2, a loss of taste or smell, fever, and a cough were significant predictors of a positive test result [[53](#ref-19IBzQBhB)]. It is important to note that in this study, the predictive value of symptoms may be underestimated if they are not specific to COVID-19 because the outcome measured was a positive, as opposed to a negative, COVID-19 test result. At the time the surveys were conducted, due to limits in US testing infrastructure, respondents typically needed to have some symptoms known to be specific to COVID-19 in order to qualify for testing in the first place. Widespread testing of asymptomatic individuals may therefore provide additional insight into the range of symptoms associated with COVID-19.

Consistent with the wide range of symptoms observed, COVID-19 can affect diverse body systems in addition to causing respiratory problems [[54](#ref-wK2afyL8)]. For example, COVID-19 can lead to acute kidney injury, especially in patients with severe respiratory symptoms or certain preexisting conditions [[55](#ref-17hdoiBo9)]. It can also cause neurological complications [[56](#ref-1CqoZGSKK),[57](#ref-1EF9EpbZ5)], potentially including stroke, seizures or meningitis [[58](#ref-nBGGUV9s),[59](#ref-LLm9d62d)]. COVID-19 has also been associated with an increased incidence of large vessel stroke, particularly in patients under the age of 40 [[60](#ref-aRNqjkMg)], and other thrombotic events including pulmonary embolism and deep vein thrombosis [[61](#ref-3UBbDrG2)]. The mechanism behind these complications has been suggested to be related to coagulopathy, with reports indicating the presence of antiphospholipid antibodies [[62](#ref-8OnbWuhF)] and elevated levels of d-dimer and fibrinogen degradation products in deceased patients [[63](#ref-dUfws1q0)]. Other viral infections have been associated with coagulation defects and changes to the coagulation cascade; notably SARS was also found to lead to disseminated intravascular coagulation and was associated with both pulmonary embolism and deep vein thrombosis [[64](#ref-p5EhN9Qd)]. The mechanism behind these insults has been suggested to be related to inflammation-induced increases in the von Willebrand factor clotting protein, leading to a pro-coagulative state [[64](#ref-p5EhN9Qd)]. Abnormal clotting (thromboinflammation or coagulopathy) has been increasingly discussed recently as a possible key mechanism in many cases of severe COVID-19, and may be associated with the high d-dimer levels often observed in severe cases [[65](#ref-v1EIzwfx),[66](#ref-18AiyvhO8),[67](#ref-PyKMLraw)]. This excessive clotting in lung capillaries has been suggested to be related to a dysregulated activation of the complement system, part of the innate immune system [[68](#ref-14dyYQY7s),[69](#ref-17y6YeJ6R)].

### Cytokine Release Syndrome

Symptoms of a disease can be caused by a pathogen, but they can also be caused by the immune system’s reaction to the pathogen. A dysregulated immune response can cause significant damage to the host [[70](#ref-o6BQnEt7),[71](#ref-b1QVdemU),[72](#ref-1GnFL9zeN)]. The inflammatory response has received particular attention for its role in both a healthy response to infection and a pathogenic one. Inflammation is one of the most visible components of the immune response, as it is responsible for the hallmarks of injury, such as pain, heat, and swelling [[73](#ref-sXusUkLI)]. In response to injury or to signaling by pattern recognition receptors (PRRs) indicating the detection of a molecular pattern associated with a pathogen or foreign body, the immune system stimulates leukocytes that travel to the site of the threat, where they then produce cytokines [[73](#ref-sXusUkLI)]. Cytokines are a diverse group of small proteins that play an important role in intercellular signaling [[74](#ref-bZMKqj6e)]. Cytokines can be both pro- and anti-inflammatory, which means they can either stimulate or inhibit the production of additional cytokines [[74](#ref-bZMKqj6e),[75](#ref-DYbswZ6D)]. Some notable pro-inflammatory cytokines include the interleukins: IL-1β and IL-6 and tumor necrosis factor α (TNF-α) [[75](#ref-DYbswZ6D)]. Anti-inflammatory cytokines play an immunoregulatory role complementary to the cascading effect of pro-inflammatory cytokines [[74](#ref-bZMKqj6e),[75](#ref-DYbswZ6D)]. A number of interleukins and interferons play anti-inflammatory roles, and receptors or receptor antagonists for inflammatory cytokines are also important for regulating inflammation [[75](#ref-DYbswZ6D)]. IL-10 is an anti-inflammatory cytokine of particular note because it regulates the expression of TNF-α, IL-1, and IL-6 [[75](#ref-DYbswZ6D)]. When the pro- and anti-inflammatory responses are both commensurate with the threat posed, the immune system drives a shift back to homeostasis [[76](#ref-16CYY7vzG)]. However, when the responses are disproportionate, the cytokine response can become dysregulated. Too low of an inflammatory response will not eliminate the immune threat [[76](#ref-16CYY7vzG)]. In contrast, if the response is dysregulated towards excessive pro-inflammatory cytokine activity, inflammation can cascade [[77](#ref-xuMYmc7W)] and cause cell damage, among other problems [[73](#ref-sXusUkLI)]. Elevated levels of inflammation over the long-term are associated with many chronic health conditions, including type 2 diabetes, dementia and Alzheimer’s, and arthritis, among others [[78](#ref-OvhepUrA)]. On a shorter timescale, dysregulated systemic inflammation can cause sepsis, which can lead to multi-organ failure and death [[74](#ref-bZMKqj6e),[79](#ref-6eWgCrWj)].

Cytokines have been investigated for their role in the immune response to lung infections long before the COVID-19 pandemic. Dysregulation of the inflammatory response, including elevated levels of pro-inflammatory cytokines, is found in patients with ARDS, which is a severe condition that can arise from pneumonia, SARS, and COVID-19 [[77](#ref-xuMYmc7W)]. One study of patients with and at risk for ARDS, specifically those who were intubated for medical ventilation, found that shortly after the onset of ARDS, anti-inflammatory cytokine concentration in BALF increased relative to the concentration of pro-inflammatory cytokines [[80](#ref-f61jsRKY)]. The results suggest that an increase in pro-inflammatory cytokines such as IL-6 may signal the onset of ARDS, but recovery depends on an increased anti-inflammatory response [[80](#ref-f61jsRKY)]. However, patients with severe ARDS were excluded from this study. Acute phase response to an infection can also cause damage to the capillary endothelium, allowing leaks that disrupt the balance between pro-inflammatory cytokines and their regulators [[80](#ref-f61jsRKY)]. Hyperactivity of the pro-inflammatory response due to lung infection is commonly associated with acute lung injury and more rarely with the more severe manifestation, ARDS [[74](#ref-bZMKqj6e)]. The heightened inflammatory response in the lungs can also serve as a source for systemic inflammation, or sepsis, which can lead to multi-organ failure [[74](#ref-bZMKqj6e)]. The shift from local to systemic inflammation is a phenomenon often referred to broadly as a cytokine storm [[74](#ref-bZMKqj6e)] or, more precisely, as cytokine release syndrome (CRS) [[81](#ref-NITOJ0Ka)]. Sepsis is a known possible complication of pneumonia, and in an analysis of over 1,400 US pneumonia patients, IL-6, tumor necrosis factor (TNF), and IL-10 were found to be elevated at intake in patients who developed severe sepsis and/or ultimately deceased [[82](#ref-oSOG8N0f)]. IL-6 and TNF are pro-inflammatory cytokines, while IL-10; is anti-inflammatory [[82](#ref-oSOG8N0f)]. However, this study reported that unbalanced pro-/anti-inflammatory cytokine profiles were rare, although they measured only the three cytokines listed above. Prior work therefore made it clear that pulmonary infection and injury were associated with systemic inflammation through sepsis. While IL-6 is a biomarker sometimes used to assess cytokine storm activity in sepsis [[74](#ref-bZMKqj6e)], the relationship between cytokine profiles and the risks associated with sepsis may be more complex. IL-6 is a pleiotropic cytokine that plays an integral role in both the inflammatory and anti-inflammatory responses and is associated with both healthy and pathological responses to viral threat [[83](#ref-dBAe8aYi)].

The inflammatory response was identified early on as a potential driver of COVID-19 outcomes due to existing research in SARS and emerging research in COVID-19. In addition to the known role of cytokines in ARDS and lung infection more broadly, immunohistological analysis at autopsy of patients who deceased from SARS revealed that ACE2-expressing cells that were infected by SARS-CoV-1 showed elevated expression of IL-6, IL-1β, and TNF-α [[84](#ref-NNFGje9g)]. Similarly, the introduction of the S protein from SARS-CoV-1 to mouse macrophages was found to increase production of IL-6 and TNF-α [[85](#ref-QkTGQUcj)]. For SARS-CoV-2 infection leading to COVID-19, early reports described a cytokine storm syndrome-like response in patients with particularly severe infections [[37](#ref-UwVweB2M),[86](#ref-3HdlV9Vf),[87](#ref-cyUG1Zi2)]. Among patients hospitalized with COVID-19 in Wuhan, China, 112 out of 191 (59%) developed sepsis, including all 54 of the non-survivors [[49](#ref-10THxyeCg)]. However, the argument has been made that while the cytokine levels observed in COVID-19 patients fall outside of the normal range, they are not as high as typically found in patients with ARDS [[88](#ref-Xcrxlxnp)]. Regardless, inflammation has received significant interest both in regards to the pathology of COVID-19 as well as potential avenues for treatment, as the similarities between the cytokine storm and the pathophysiology of COVID-19 has led to the suggestion that a number of immunomodulatory pharmaceutical interventions could hold therapeutic value for the treatment of COVID-19 [[89](#ref-RIMWgsWt)].

### Pediatric Presentation

The presentation of COVID-19 infection can vary greatly among pediatric patients and, in some cases, manifests in distinct ways from COVID-19 in adults. Evidence suggests that while children and adolescents tend to have mostly asymptomatic infections, those that are symptomatic exhibit a mild illness [[90](#ref-g435TGYc),[91](#ref-SHtceaPy),[92](#ref-8sVs0pB8),[93](#ref-L81Cb2ZZ)]. A review examined symptoms reported in 17 studies of children infected with COVID-19 during the early months of the COVID-19 epidemic in China and one study from Singapore [[94](#ref-1BhN2uq3f)]. Of the more than a thousand cases described, the most common reports were for mild symptoms such as fever, dry cough, fatigue, nasal congestion and/or runny nose, while three children were reported to be asymptomatic. Severe lower respiratory infection was described in only one of the pediatric cases reviewed. Gastrointestinal symptoms such as vomiting or diarrhea were occasionally reported. Radiologic findings were not always reported in the case studies reviewed, but when they were mentioned they included bronchial thickening, ground-glass opacities, and/or inflammatory lesions [[94](#ref-1BhN2uq3f)]. Neurological symptoms have also been reported [[95](#ref-12lVfu2Qe)].

These analyses indicate that most pediatric cases of COVID-19 are not severe. Indeed, it is estimated that less than 1% of pediatric cases result in critical illness [[92](#ref-8sVs0pB8),[96](#ref-ShMJKFvG)]. However, serious complications and, in rare cases, deaths have occurred [[97](#ref-MCVe25tf)]. Of particular interest, children have occasionally experienced a serious inflammatory syndrome, multisystem inflammatory syndrome in children (MIS-C), following COVID-19 infection. This syndrome is similar in some respects to Kawasaki disease or to Kawasaki disease shock syndrome [[98](#ref-fZwrv8M),[99](#ref-AmvwCWm3),[100](#ref-68JorBeU)] and is thought to be a distinct clinical manifestation of SARS-CoV-2 due to its distinct cytokine profile and the presence of burr cells in peripheral blood smears [[101](#ref-10gSXfOKm),[102](#ref-10QhaDu6N)]. MIS-C has been associated with heart failure in some cases [[103](#ref-FYgN6gUL)]. One case study [[104](#ref-71To4laE)] described an adult who appeared to show symptoms similar to MIS-C after exposure to COVID-19, but cautioned against broad conclusions; a second possible adult case has also been reported [[105](#ref-UBDAxWWv)]. The presentation of SARS-CoV-2 infection is therefore likely to be largely distinct between adult and pediatric populations.

## Systems-Level Effects

Systems biology provides a cross-disciplinary analytical paradigm through which the host response to an infection can be analyzed. This field integrates the “omics” (genomics, transcriptomics, proteomics, metabolomics, etc.) with bioinformatics and other computational approaches. These cutting-edge research paradigms hold enormous potential for the study of the complexity of biological systems and human diseases [[106](#ref-5MhhHQO0)]. Over the last decade, systems biology approaches have been used widely to study the pathogenesis of diverse types of life-threatening acute and chronic infectious diseases [[107](#ref-8esIIWt9)]. Omics-based studies have also provided meaningful information regarding host immune responses and surrogate protein markers in several viral, bacterial and protozoan infections [[108](#ref-8vUHj9cm)]. Though the complex pathogenesis and clinical manifestations of SARS-CoV-2 infection are not yet fully understood, omics technologies offer the opportunity for discovery-driven analysis of biological changes associated with SARS-CoV-2 infection. For example, previous studies suggest that infection by coronaviruses, such as SARS-CoV-1 and MERS-CoV, and other viruses is associated with the upregulation of ACE2. In several preliminary assays and an analysis of previous microarray data, ACE2 expression was reported to be significantly upregulated following infection of human embryonic kidney cells and human airway epithelial cells [[37](#ref-UwVweB2M)]. This study also reported that direct stimulation with inflammatory cytokines such as type I interferons (e.g., IFNβ) resulted in the upregulation of ACE2 in human bronchial epithelial cells, with treated groups showing four-fold higher ACE2 expression than control groups at 18 hours post-treatment [[37](#ref-UwVweB2M)]. Whether SARS-CoV-2 facilitates the positive regulation of its own transmission between host cells is still unclear, the host immune response itself likely plays a key role in mediating infection-associated pathologies. A systems-biology approach allows for analyses such as these to identify possible phenotypic and endophenotypic responses to SARS-CoV-2 infection and to develop new hypotheses about how pathogenesis proceeds.

### Transcriptomics

In addition to the study described above, two other studies have profiled expression following SARS-CoV-2 infection using human cell lines. The first study [[109](#ref-pHrstqMQ)] compared transcriptional responses to SARS-CoV-2 and to other respiratory viruses, including MERS-CoV, SARS-CoV, human parainfluenza virus 3 (HPIV3), respiratory syncytial virus (RSV), and influenza A virus (IAV). The authors analyzed the responses of three human cell lines: A549 (adenocarcinomic human alveolar basal epithelial cells), Calu-3 (human airway epithelial cells derived from human bronchial submucosal glands), and MRC-5 (human fetal lung fibroblast cells). As the viral receptor ACE2 has low expression in A549 cells, they supplemented the A549 cells with adenovirus (AdV)-based vectors expressing either mCherry (a fluorescent protein used as a control) or ACE2 (A549-ACE2). The authors also measured host transcriptional responses to SARS-CoV-2 in primary normal human bronchial epithelial cells (HBEC or NHBE cells), nasal washes from an animal model (ferret), and lung samples from two COVID-19 patients. Differential expression (DE) analysis was then carried out to compare infected cells with control cells that underwent only a mock treatment. In the hosts where SARS-CoV-2 was able to replicate efficiently, DE analysis revealed that the transcriptional response was significantly different from the response to all of the other viruses tested. A unique proinflammatory cytokine signature associated with SARS-CoV-2 was present in cells exposed to both high and low doses of the virus, with the cytokines IL-6 and IL1RA uniquely elevated in response to SARS-CoV-2 relative to other viruses. However, the A549-ACE2 cells showed significant IFN-I or IFN-III expression when exposed to high, but not low, doses of SARS-CoV-2. This finding suggests that IFN induction is dependent on the extent of exposure. Similarly, in cells from the NHBE line, ferrets, and COVID-19 patients, chemokine signaling was significantly enriched, but there was no significant induction of IFN-I or IFN-III. Together, these results suggest that SARS-CoV-2 induces a limited antiviral state with low IFN-I or IFN-III expression and a moderate IFN-stimulated gene response, in contrast to other viruses. However, in ACE2-expressing A549 cells, this state could be overcome by using a 10-fold increase in SARS-CoV-2 exposure. This finding suggests that the SARS-CoV-2 interferon antagonist is insufficient for large doses of the virus [[109](#ref-pHrstqMQ)]. This hypothesis was further supported by a recent study [[110](#ref-PJgYPcKM)] that showed that the SARS-CoV-2 *ORF3b* gene suppresses IFNB1 promoter activity (IFN-I induction) more efficiently than the SARS-CoV-1 *ORF3b* gene. Taken together, these findings suggest that a unique cytokine profile is associated with the response to the SARS-CoV-2 virus and that this response differs depending on the extent of exposure.

Another study [[111](#ref-Gj8vlc0W)] analyzed cells’ transcriptional response to SARS-CoV-2 and SARS-CoV-1 over time. They characterized the response of three human cell lines, H1299 (human non-small cell lung carcinoma cell line), Calu-3, and Caco-2 (human epithelial colorectal adenocarcinoma cell line), at 4 to 36 hours post infection (hpi). Using poly(A) bulk RNA-seq, the authors found negligible susceptibility of H1299 cells (< 0.08 viral read percentage of total reads) compared to Caco-2 and Calu-3 cells (>10% of viral reads). This finding suggests that the risk of infection varies among cell types, and that cell type could influence which hosts are more or less susceptible. Based on visual inspection of microscopy images alongside transcriptional profiling, the authors also showed distinct responses among the host cell lines evaluated. In contrast to Caco-2, Calu-3 cells infected with SARS-CoV-2 showed signs of impaired growth and cell death at 24 hpi, as well as moderate IFN induction with a strong up-regulation of IFN-stimulated genes. Interestingly, the results were similar to those reported in Calu-3 cells exposed to much higher levels of SARS-CoV-2 [[109](#ref-pHrstqMQ)], as described above. This finding suggests that IFN induction in Calu-3 cells is not dependent on the level of exposure, in contrast to A549-ACE2 cells. The discrepancy could be explained by the observations that Calu-3 cells are highly susceptible to SARS-CoV-2 and show rapid viral replication [[26](#ref-JOJ2n3gC)], whereas A549 cells are incompatible with SARS-CoV-2 infection [[112](#ref-16EFyBURq)]. This discrepancy raises the concern that *in vitro* models may vary in their similarity to the human response, underscoring the importance of follow-up studies in additional models.

### Proteomics

One early proteomics study investigated changes associated with *in vitro* SARS-CoV-2 infection using Caco-2 cells [[113](#ref-11xZWeHN3)]. This study reported that SARS-CoV-2 induced alterations in multiple vital physiological pathways, including translation, splicing, carbon metabolism and nucleic acid metabolism in the host cells. Another area of interest is whether SARS-CoV-2 is likely to induce similar changes to other HCoV. For example, because of the high level of sequence homology between SARS-CoV-2 and SARS-CoV-1, it has been hypothesized that sera from convalescent SARS-CoV-1 patients might show some efficacy in cross-neutralizing SARS-CoV-2-S-driven entry [[25](#ref-15l3di3Wj)]. However, despite the high level of sequence homology, certain protein structures might be immunologically distinct, which would be likely to prohibit effective cross-neutralization across different SARS species [[114](#ref-GhJYjnft)]. Consequently, proteomic analyses of SARS-CoV-1 might also provide some essential information regarding the new pathogen [[115](#ref-GpnngtWK),[116](#ref-JLTf2Fwb)].

Considering the paucity of omics-level big data sets for SARS-CoV-2 currently available, existing data hubs that contain information for other coronaviruses such as UniProt, NCBI Genome Database, The Immune Epitope Database and Analysis Resource (IEDB), and The Virus Pathogen Resource (ViPR) will serve as useful resources for computational and bioinformatics research on SARS-CoV-2. Using such databases, the systems-level reconstruction of the protein-protein interaction (PPI) will enable the generation of hypotheses about the mechanism of action of SARS-CoV-2 and suggest potential drug targets. In an initial study [[117](#ref-phJM8g2Y)], 26 of the 29 SARS-CoV-2 proteins were cloned and expressed in HEK293T kidney cells, allowing for the identification of 332 high-confidence human proteins interacting with them. Notably, this study suggested that SARS-CoV-2 interacts with innate immunity pathways. Ranking pathogens by the similarity between their interactomes and that of SARS-CoV-2 suggested *West Nile virus*, *Mycobacterium tuberculosis*, and *human papillomavirus* infections as the top three hits. Therefore, given the lung symptoms associated with COVID-19, the *Mycobacterium tuberculosis* host-pathogen interactome in particular might provide new insights to the mechanism of SARS-CoV-2 infection. Additionally, it was suggested that the envelope protein, E, could disrupt host bromodomain-containing proteins, i.e., BRD2 and BRD4, that bind to histones, and the spike protein could likely intervene in viral fusion by modulating the GOLGA7-ZDHHC5 acyl-transferase complex to increase palmitoylation, which is a post-translational modification that affects how proteins interact with membranes [[118](#ref-mXUCjmCh)].

Another study [[119](#ref-NLoN4aYj)] used patient-derived peripheral blood mononuclear cells (PBMCs) to identify 251 host proteins targeted by SARS-CoV-2. This study also reported that more than 200 host proteins were disrupted following infection. In particular, a network analysis showed that nsp9 and nsp10 interacted with NF-Kappa-B-Repressing Factor (NKRF), which encodes a transcriptional repressor that mediates repression of genes responsive to Nuclear Factor kappa-light-chain-enhancer of activated B-cells (NF-kB). These genes are important to pro-, and potentially also anti-, inflammatory signaling [[120](#ref-KAqOiTeZ)]. This finding could explain the exacerbation of the immune response that shapes the pathology and the high cytokine levels characteristic of COVID-19, possibly due to the chemotaxis of neutrophils mediated by IL-8 and IL-6. Finally, it was suggested [[121](#ref-1ccnm0N9)] that the E protein of both SARS-CoV-1 and SARS-CoV-2 has a conserved Bcl-2 Homology 3 (BH3)-like motif, which could inhibit anti-apoptosis proteins, e.g., BCL2, and trigger the apoptosis of T cells. Several compounds are known to disrupt the host-pathogen protein interactome, largely through the inhibition of host proteins. Therefore, preliminary research in the proteomics of SARS-CoV-2 infection suggests that drugs modulating the protein-level interactions between virus and host might be worth investigating.

## Viral Evolution and Virulence

Like that of SARS-CoV-1, the entry of SARS-CoV-2 into host cells is mediated by interactions between the viral spike glycoprotein, S, and human ACE2 (hACE2) [[22](#ref-qcVbT0w4),[25](#ref-15l3di3Wj),[122](#ref-15EIBRhef),[123](#ref-12TZ7hPMA),[124](#ref-wCbhn23d),[125](#ref-VX8OWaGj),[126](#ref-15Q2XgkK7),[127](#ref-t1e4CW9A)]. Differences in how the S proteins of the two viruses interact with the hACE2 receptor could also partially account for the increased transmissibility of SARS-CoV-2. Recent studies have reported conflicting binding constants for the S-hACE2 interaction, though they have agreed that the SARS-CoV-2 S protein binds with equal, if not greater, affinity than the SARS-CoV-1 S protein does [[9](#ref-CqQVkaqj),[22](#ref-qcVbT0w4),[125](#ref-VX8OWaGj)]. The C-terminal domain of the SARS-CoV-2 S protein in particular was identified as the key region of the virus that interacts with hACE2, and the crystal structure of the C-terminal domain of the SARS-CoV-2 S protein in complex with hACE2 reveals stronger interaction and a higher affinity for receptor binding than that of SARS-CoV-1 [[126](#ref-15Q2XgkK7)]. Among the 14 key binding residues identified in the SARS-CoV-1 S protein, eight are conserved in SARS-CoV-2, and the remaining six are semi-conservatively substituted, potentially explaining variation in binding affinity [[22](#ref-qcVbT0w4),[125](#ref-VX8OWaGj)]. Recent crystal structures have shown that the receptor-binding domain (RBD) of the SARS-CoV-2 S protein, like that of other coronaviruses, undergoes stochastic hinge-like movement that flips it from a “closed” conformation, in which key binding residues are hidden at the interface between protomers, to an “open” one [[9](#ref-CqQVkaqj),[22](#ref-qcVbT0w4)]. Because the RBD plays such a critical role in viral entry, blocking its interaction with ACE2 could represent a promising therapeutic approach. Nevertheless, despite the high structural homology between the SARS-CoV-2 RBD and that of SARS-CoV-1, monoclonal antibodies targeting SARS-CoV-1-RBD failed to bind to SARS-CoV-2-RBD [[9](#ref-CqQVkaqj)]. Promisingly, though, sera from convalescent SARS patients inhibited SARS-CoV-2 viral entry *in vitro*, albeit with lower efficiency than it inhibited SARS-CoV-1 [[25](#ref-15l3di3Wj)].

Comparative genomic analysis reveals that several regions of the coronavirus genome are likely critical to virulence. The S1 domain of the spike protein, which contains the receptor binding motif, evolves more rapidly than *S*’s S2 domain [[16](#ref-13wWdgODZ),[17](#ref-OVsxrEuX)]. However, even within the S1 domain, some regions are more conserved than others, with the receptors in S1’s N-terminal domain (S1-NTD) evolving more rapidly than those in its C-terminal domain (S1-CTD) [[17](#ref-OVsxrEuX)]. Both S1-NTD and S1-CTD are involved in receptor binding and can function as RBDs to bind proteins and sugars [[16](#ref-13wWdgODZ)], but RBDs in the S1-NTD typically bind to sugars, while those in the S1-CTD recognize protein receptors [[5](#ref-17DSmRo9H)]. Viral receptors show higher affinity with protein receptors than sugar receptors [[5](#ref-17DSmRo9H)], which suggests that positive selection on or relaxed conservation of the S1-NTD might reduce the risk of a deleterious mutation that would prevent binding. The SARS-CoV-2 S protein also contains an RRAR furin recognition site at the S1/S2 junction [[9](#ref-CqQVkaqj),[22](#ref-qcVbT0w4)], setting it apart from both bat coronavirus RaTG13, with which it shares 96% genome sequence identity, and SARS-CoV-1 [[128](#ref-VSkK7CeP)]. Such furin cleavage sites are commonly found in highly virulent influenza viruses, and as such may contribute to the heightened pathogenicity of SARS-CoV-2 [[129](#ref-NsORsLig),[130](#ref-vhHB3yyS)]. Effective cell entry is a critical component to pathogenesis and therefore an important process to understand when examining possible therapeutics.

## Mechanism of Transmission

Once a human host is infected with a virus, person-to-person viral transmission of a virus can occur through several possible mechanisms. The primary mechanisms associated with respiratory viruses are contact, droplet, and aerosol transmission [[131](#ref-CJbohYmK)]. Contact transmission can occur through either direct contact with a contagious person or indirect contact with active viral particles on a contaminated surface [[132](#ref-KMb6knfE)]. This latter mode of transmission is also called fomite transmission [[133](#ref-1FLLN4PSl)]. Viral particles can enter the body if they then come in contact with the oral, nasal, eye, or other mucus membranes [[132](#ref-KMb6knfE)]. Droplet transmission occurs when a contagious individual sneezes, coughs, or exhales and produces respiratory droplets that can contain a large number of viral particles [[132](#ref-KMb6knfE)]. Contact with these droplets can occur either through direct exposure to the droplets, such as breathing in droplets produced by a sneeze, or exposure to particles that have settled on a surface [[132](#ref-KMb6knfE)]. Aerosol transmission refers to much smaller particles (less than 5 micrometers) that are also produced by sneezing, coughing, or exhaling [[131](#ref-CJbohYmK),[132](#ref-KMb6knfE)]. The small size of these particles allows them to remain suspended over a longer period of time and potentially to be moved by air currents [[132](#ref-KMb6knfE)]. Additionally, viral particles deposited on surfaces via large respiratory droplets can also later be aerosolized [[132](#ref-KMb6knfE)]. Droplet and/or contact transmission are both well-accepted modes of transmission for many viruses associated with common human illnesses, including influenza and rhinovirus [[132](#ref-KMb6knfE)]. The extent to which aerosol transmission contributes to the spread of respiratory viruses is less clear. In influenza A, for example, viral particles can be detected in aerosols produced by infected individuals, but the extent to which these particles drive the spread of influenza A infection remains under debate [[131](#ref-CJbohYmK),[132](#ref-KMb6knfE),[134](#ref-AC4okoVf),[135](#ref-Hct9jRcb),[136](#ref-i6tTpqwA)]. Regardless of its role in the spread of influenza A, however, aerosol transmission likely played a role in outbreaks such as the 1918 Spanish Influenza (H1N1) and 2009 “swine flu” (pH1N1) [[136](#ref-i6tTpqwA)]. Contact, droplet, and aerosol transmission are therefore all worth evaluating when considering possible modes of transmission for a respiratory virus like SARS-CoV-2.

### Transmission of HCoV

All three of these mechanisms have been identified as contributors to the transmission of HCoV [[132](#ref-KMb6knfE)], including the highly pathogenic coronaviruses SARS-CoV-1 and MERS-CoV [[137](#ref-rjVw7V94),[138](#ref-G5NJrE75)]. Transmission of SARS-CoV-1 is thought to proceed primarily through droplet transmission, but aerosol transmission is also considered possible [[132](#ref-KMb6knfE)], and fomite transmission may have also played an important role in some outbreaks [[139](#ref-yR57NFIB)]. Similarly, the primary mechanism of MERS transmission is thought to be droplets because inter-individual transmission appears to be associated with close interpersonal contact (e.g., household or healthcare settings), but aerosolized particles of the MERS virus have been reported to persist much more robustly than influenza A under a range of environmental conditions [[140](#ref-s3oVNbGV),[141](#ref-1FjDLPCye)]. While droplet-based and contact transmission were initially considered to be the primary modes by which SARS-CoV-2 spread [[142](#ref-vP6yTZ0y)], as additional information has emerged, the possibility of aerosol transmission has also been raised [[143](#ref-16SDeiudC),[144](#ref-yXKcviw8),[145](#ref-9ouDX5IN)]. For example, the detection of SARS-CoV-2 viral particles in air samples taken from hospitals treating COVID-19 patients led to the concern that the virus could be spreading via aerosols [[146](#ref-bQLMgMGC)]. The stability of the virus both in aerosols and on a variety of surfaces appeared similar to that of SARS-CoV-1 [[144](#ref-yXKcviw8)], and fomite transmission could also play a role in transmission (e.g., [[147](#ref-1HOZHff6)]). However, while the possibility of aerosol transmission seems plausible, the evidence suggests that droplet and contact transmission are the dominant mechanisms driving the spread of the virus [[148](#ref-82XnTbtX)], and the risk of fomite transmission under real-world conditions is likely to be substantially lower than the conditions used for experimental analyses [[149](#ref-cw5j7x80)]. These mechanisms may differ in their relevance to different types of transmission events, such as transmission within households, nosocomial transmissions, and transmission in indoor versus outdoor spaces.

### Symptoms and Viral Spread

Other aspects of pathogenesis are also important to understanding how the virus spreads, especially the relationship between symptoms, viral shedding, and contagiousness. Symptoms associated with reported cases of COVID-19 range from mild to severe [[1](#ref-gHWlMufv)], but some individuals who contract COVID-19 remain asymptomatic throughout the duration of the illness [[150](#ref-wxVni9Hz)]. The incubation period, or time period between exposure and the onset of symptoms, has been estimated at five to eight days (4.91 and 7.54 in two different cities) [[151](#ref-NIxttl2v),[152](#ref-QFVSrboR)], and estimates suggest that viral shedding may begin long before the onset of symptoms (12.3 days with a 95% CI of 5.9 - 17.0) and peak around the onset of symptoms [[153](#ref-1654TaAK4)]. As these trends became apparent, concerns arose due to the potential for individuals who did not or did not yet show symptoms to transmit the virus [[154](#ref-Oyy1OUTT)]. Recovered individuals may also be able to transmit the virus after their symptoms cease. Estimates of the communicable period based on twenty-four individuals who tested positive for SARS-CoV-2 prior to or without developing symptoms estimated that individuals may be contagious for one to twenty-one days, but they note that this estimate may be low [[150](#ref-wxVni9Hz)]. Initially, viral nucleic acids were reported to remain at observable levels in the respiratory specimens of recovering hospitalized COVID-19 patients for a median of 20 days and with a maximum observed duration through 37 days, when data collection for the study ceased [[49](#ref-10THxyeCg)]. As more estimates of the duration of viral shedding are released, they are beginning to converge around approximately three weeks from first positive PCR test and/or onset of symptoms (which, if present, are usually identified within three days of the initial PCR test). For example, viral shedding was reported for up to 28 days following symptom onset by a second study [[155](#ref-Xp5HT2SF)] and for one to 24 days from first positive PCR test with a median of 12 days [[39](#ref-azgJqujy)]. On the other hand, almost 70% of patients were reported to still have symptoms at the time that viral shedding ceased, although all symptoms reduced in prevalence between onset and cessation of viral shedding (CVS) [[156](#ref-14fWuiUKS)]. They also reported that the median time that elapsed between the onset of symptoms and CVS was 23 days and between first positive PCR test and CVS was 17 days [[156](#ref-14fWuiUKS)]. The fact that this study reported symptom onset to predate the first positive PCR test by an average of three days, however, suggests that there may be some methodological differences between it and related studies. Furthermore, an analysis of residents of a nursing home with a known SARS-CoV-2 case measured similar viral load in residents who were asymptomatic regardless of whether they later developed symptoms, and the load in the asymptomatic residents was comparable to that of residents who displayed either typical of atypical symptoms [[157](#ref-ac2kt3rh)]. Taken together, these results suggest that the presence or absence of symptoms are not reliable predictors of viral shedding or of SARS-CoV-2 status (e.g, [[158](#ref-lVDiHYp)]). However, viral shedding is not necessarily indicative of contagiousness. The risk of spreading the infection was low after ten days from the onset of symptoms, as viral load in sputum was found to be unlikely to pose a significant risk based on their efforts to culture samples *in vitro* [[155](#ref-Xp5HT2SF)].

The extent to which asymptomatic (or presymptomatic) individuals are able to transmit SARS-CoV-2 has been a question of both scientific and community interest. Early reports (February and March 2020) described transmission from presymptomatic SARS-CoV-2-positive individuals to close family contacts [[159](#ref-E6Qv6YQ0),[160](#ref-18Orbxz8t)]. One of these reports [[160](#ref-18Orbxz8t)] also included a description of an individual who tested positive for SARS-CoV-2 but never developed symptoms. Later analyses also sought to estimate the proportion of infections that could be traced back to a presymptomatic or asymptomatic individual (e.g., [[161](#ref-4M3GM1sg)]). Estimates of the proportion of individuals with asymptomatic infections have varied widely. The proportion of asymptomatic individuals on board the Diamond Princess cruise ship, which was the site of an early COVID-19 outbreak, was estimated at 17.9% [[162](#ref-6K5lsF5i)]. In contrast, a model using the prevalence of antibodies among residents of Wuhan, China estimated a much higher of asymptomatic cases, at approximately 7 in 8, or 87.5% [[163](#ref-sk1NbA7K)]. An estimate of the population of care homes in London found that, among the residents (median age 85), the rate of asymptomatic infection was 43.8%, and among the caretakers (median age 47), the rate was 49.1% [[164](#ref-kIumgXPI)]. The duration of viral shedding may also be longer in individuals with asymptomatic cases of COVID-19 compared to those who do show symptoms [[165](#ref-10OGkFiGJ)]. As a result, the potential for individuals who do not know they have COVID-19 to spread the virus raises significant concerns. In Singapore and Tianjin, two cities studied to estimate incubation period, an estimated 40-50% and 60-80% of cases, respectively, were estimated to be caused by contact with an asymptomatic individuals [[151](#ref-NIxttl2v)]. An analysis of viral spread in the Italian town of Vo’, which was the site of an early COVID-19 outbreak, revealed that 42.5% of cases were asymptomatic and that the rate was similar across age groups [[166](#ref-dMjSbAQV)]. They argued that the town’s lockdown was imperative for controlling the spread of COVID-19 because it isolated the asymptomatic individuals. While more models are likely to emerge to better explore the effect of asymptomatic individuals on SARS-CoV-2 transmission, these results suggest that strategies for identifying and containing asymptomatic, but contagious individuals are important for managing community spread.

### Estimating the Fatality Rate

Estimating the occurrence of asymptomatic and mild COVID-19 cases is important to identifying the mortality rate associated with COVID-19. The mortality rate of greatest interest would be the total number of fatalities as a fraction of the total number of people infected. One metric reported is often case fatality rate (CFR), which simply compares the number of COVID-19 related deaths to the number of confirmed or suspected cases. However, in locations without universal testing protocols, it is impossible to identify all exposed or all infected individuals because so many asymptomatic or mild cases go undetected. Therefore, a more informative metric is the infection fatality rate (IFR), which compares the known deaths to the estimated number of cases. Meta-analyses have produced estimates of global IFR ranging from as low as 0.1% to as high as 1.04% [[167](#ref-BENQIwOM),[167](#ref-BENQIwOM),[168](#ref-AavOV1He),[169](#ref-12CiweatZ)], and this estimate was also supported by a repeated cross-sectional serosurvey conducted in New York City that revealed an estimated IFR of 0.97% [[170](#ref-tIjAygts)]. All of these estimates note that IFR varies widely around the world. Estimates of infection rates are becoming more feasible as more data becomes available for modeling and will be bolstered as serological testing becomes more common and available.

## Dynamics of Transmission

Disease spread dynamics can be estimated using R0, the basic reproduction number, and Rt, the effective reproduction number. Thus, accurate estimates are crucial to understanding the dynamics of infection and to predicting the effects of different interventions. R0 and the timescale of infection (measured by the infectious period and the exposed period) govern population-level epidemic dynamics, with R0 being one of the most important epidemiological parameters [[171](#ref-14l4fvU71)]. R0 is the average number of new (secondary) infections caused by one infected person, assuming a wholly susceptible population [[172](#ref-16CD4voW1)]. A simple mechanistic model used to describe infectious disease dynamics is a susceptible-infected-recovered (SIR) compartmental model. In this formulation individuals move through three states: susceptible, infected, and recovered; two parameters, and , specify the rate at which the infectious recover, and the infection transmission rate, respectively. In this simple formulation, R0 is estimated as the ratio of and .[[171](#ref-14l4fvU71),[173](#ref-YubbhU7G)]. A pathogen can invade a susceptible population only if R0 > 1 [[171](#ref-14l4fvU71),[174](#ref-JRaVgcNd)]. The spread of an infectious disease at a particular time t can be quantified by Rt, the effective reproduction number, which assumes that part of the population has already recovered (and thus gained immunity to reinfection) or that mitigating interventions were put into place. For example, if only a fraction St of the population is still susceptible, Rt = St x R0. When Rt is greater than 1, an epidemic grows (i.e., the proportion of the population that is infectious increases); when Rt is less than 1, the proportion of the population that is infectious decreases. R0 and Rt can be estimated directly from epidemiological data or inferred using mathematical modeling. Modeling approaches are typically based upon a classic epidemiological model structure: the SIR model and its extensions [[175](#ref-1Aozf7KBW),[176](#ref-SPInPew0)]. In the context of SARS-CoV-2, more complex modified susceptible-exposed-infectious-recovered (SEIR) models are commonly used.

Estimates of R0 for COVID-19 lie in the range R0=1.4-6.5 [[177](#ref-1E0r4uZy9),[178](#ref-LHtuVmaq),[179](#ref-mHYmt0mv)]. Variation in R0 is expected between different populations, and the estimated values of R0 discussed below are for specific populations in specific environments. The different estimates of R0 should not necessarily be interpreted as a range of estimates of the same underlying parameter. In one study of international cases, the predicted value was R0=1.7 [[180](#ref-vITui6ac)]. In China (both Hubei province and nationwide), the value was predicted to lie in the range R0=2.0-3.6 [[177](#ref-1E0r4uZy9),[181](#ref-nxM0rP5R),[182](#ref-WLc2UMgQ)]. Another estimate based on a cruise ship where an outbreak occurred predicted R0=2.28 [[183](#ref-10bBqMHH7)]. SEIR model-derived estimates of R0 range from 2.0 - 6.5 in China [[184](#ref-ITh0Anof),[185](#ref-1BMU7sKbs),[186](#ref-Yj8Xh4Wz),[187](#ref-ZzrrVDoE)] to R0=4.8 in France [[188](#ref-itj26agd)]. Using the same model as for the French population, a study estimated R0=2.6 in South Korea [[188](#ref-itj26agd)], which is consistent with other studies [[189](#ref-YfOcGRRa)]. From a meta-analysis of studies estimating R0, [[178](#ref-LHtuVmaq)] predict the median as R0=2.79.

Inference of the effective reproduction number can provide insight into how populations respond to an infection and the effectiveness of interventions. In China, Rt was predicted to lie in the range 1.6-2.6 in January 2020, before travel restrictions [[190](#ref-tiRfUgvs)]. Rt decreased from 2.35 one week before travel restrictions were imposed (January 23, 2020), to 1.05 one week after. Using their model, the authors also estimated the probability of new outbreaks occurring: the probability of a single individual exporting the virus and causing a large outbreak is 17-25%, assuming MERS-like or SARS-like transmission, and the probability of a large outbreak occurring after ≥4 infections exist at a new location is greater than 50%. An independent study came to similar conclusions, finding Rt=2.38 in the two-week period before January 23 with a decrease to Rt = 1.34 (using data from January 24 to February 3) or Rt=0.98 (using data from January 24 to February 8) [[179](#ref-mHYmt0mv)]. In South Korea, Rt was inferred for February through March 2020 in two cities, Daegu (the center of the outbreak) and Seoul [[189](#ref-YfOcGRRa)]. Metro data was also analyzed to estimate the effects of social distancing measures. Rt decreased in Daegu from around 3 to <1 over the period that social distancing measures were introduced. In Seoul, Rt decreased slightly, but remained close to 1 (and larger than Rt in Daegu). These findings indicate that social distancing measures appeared to be effective in containing the infection in Daegu, but in Seoul, Rt remained above 1, meaning secondary outbreaks remained possible. The study also shows the importance of region-specific analysis: the large decline in case load nationwide was mainly due to the Daegu region and could mask persistence of the epidemic in other regions, such as Seoul and Gyeonggi-do. In Iran, estimates of Rt declined from 4.86 in the first week to 2.1 by the fourth week after the first cases were reported [[191](#ref-TH4ymqvJ)]. In Europe, analysis of 11 countries inferred the dynamics of Rt over a time range from the beginning of the outbreak until March 28, 2020, by which point most countries had implemented major interventions (such as school closures, public gathering bans, and stay-at-home orders) [[192](#ref-Frya4XA4)]. Across all countries, the mean Rt before interventions began was estimated as 3.87; Rt varied considerably, from below 3 in Norway to above 4.5 in Spain. After interventions, Rt decreased by an average of 64% across all countries, with mean Rt=1.43. The lowest predicted value was 0.97 for Norway and the highest was 2.64 for Sweden, which may be in part because Sweden did not implement social distancing measures on the same scale as other countries. The study concludes that while large changes in Rt are observed, it is too early to tell whether the interventions put into place are sufficient to decrease Rt below 1.

More generally, population-level epidemic dynamics can be both observed and modelled [[173](#ref-YubbhU7G)]. Data and empirically determined biological mechanisms inform models, while models can be used to try to understand data and systems of interest or to make predictions about possible future dynamics, such as the estimation of capacity needs [[193](#ref-NCcssjLV)] or the comparison of predicted outcomes among prevention and control strategies [[194](#ref-11CWFYFDL),[195](#ref-J4RCpntH)]. Many current efforts to model Rt have led to tools that assist the visualization of estimates in real time or over recent intervals [[196](#ref-WA1R5R95),[197](#ref-17CQnstjh)]. While these may be valuable resources, it is important to note that the estimates arise from models containing many assumptions and are dependent on the quality of the data they use, which varies widely by region.

## Molecular Signatures and Transmission

Genetic variation in SARS-CoV-2 has been used to elucidate patterns over time and space. Mutations observed in individual SARS-CoV-2 genome sequences can be used to trace transmission patterns and have provided insights during outbreak investigations [[198](#ref-p9LUiyCN),[199](#ref-UUAeVUaR),[200](#ref-2w7lNKxQ)]. Similar mutations observed in several patients may indicate that the patients belong to the same transmission group. The tracking of SARS-CoV-2 mutations is recognized as an essential tool for controlling outbreaks that may facilitate tracing the paths of SARS-CoV-2’s spread [[201](#ref-jVbH3kJR)]. Several studies used phylogenetic analysis to determine the source of local COVID-19 outbreaks in Connecticut (USA), [[202](#ref-W5Z7Ztlg)], the New York City area (USA) [[203](#ref-FVmbE2oW)], and Iceland [[204](#ref-WJrnUyZf)]. There is an ongoing effort to collect SARS-CoV-2 genomes throughout the COVID-19 outbreak, and as of January 18, 2021 more than 381,000 genome sequences have been collected from patients. The sequencing data can be found at GISAID [[205](#ref-qIXPib3m)], NCBI [[206](#ref-cyzrC7qd)], and COVID-19 data portal [[207](#ref-BjPoOYSA)].

## Conclusions

The novel coronavirus SARS-CoV-2 is the third HCoV to emerge in the 21st century, and research into previous HCoVs has provided a strong foundation for characterizing the pathogenesis and transmission of SARS-CoV-2. Critical insights into how the virus interacts with human cells have been gained from previous research in HCoV and other viral infections. As with other HCoV, the immune response to SARS-CoV-2 is likely driven by detection of its spike protein, which allows it to enter cells through the ACE2 receptor. Epithelial cells have also emerged as the major cellular target of the virus, contextualizing the respiratory and gastrointestinal symptoms that are frequently observed in COVID-19. Many of the mechanisms that facilitate the pathogenesis of SARS-CoV-2 are currently under consideration as possible targets for the treatment or prevention of COVID-19. Research in other viruses also provides a foundation for understanding the transmission of SARS-CoV-2 among people and can therefore inform efforts to control the virus’s spread. Though it remains a question whether aerosol and fomite transmission contribute to the spread of SARS-CoV-2 in real-world settings, in general, much like SARS-CoV-1 and MERS-CoV, this virus seems to be spread primarily by droplet transmission. Asymptomatic transmission was also a concern in the SARS outbreak of 2002-03 and, as in current pandemic, presented challenges for estimating rates of infection [[208](#ref-QGjpUct1)]. However, in 2021, we are fortunate to be able to build on top of 18 years of SARS-CoV research in order to rapidly ascertain the identity and behavior of the virus.

COVID-19 is a complex disease with many possible presentations that appear to vary across the lifespan. Variability in presentation, including cases with no respiratory symptoms or with no symptoms altogether, were also reported during the SARS epidemic at the beginning of the 21st century [[208](#ref-QGjpUct1)]. The variability of both which symptoms present and their severity have presented challenges for public health agencies seeking to provide clear recommendations regarding which symptoms indicate SARS-CoV-2 infection and should prompt isolation. Additionally, asymptomatic cases add complexity both to efforts to estimate critical statistics such as R0 and Rt, which are critical to understanding the transmission of the virus, and IFR, which is an important component of understanding its impact on a given population. The development of diagnostic technologies over the course of the pandemic have facilitated more accurate identification, including of asymptomatic cases. As more cases have been diagnosed, the health conditions and patient characteristics associated with more severe infection have also become more clear, although there are likely to be significant sociocultural elements that also influence these outcomes. While many efforts have focused on adults, and especially older adults because of the susceptibility of this demographic, additional research is needed to understand the presentation of COVID-19 and MIS-C in pediatric patients. As more information is uncovered about the pathogenesis of HCoV and SARS-CoV-2 specifically, the diverse symptomology of COVID-19 has and likely will continue to conform with the ever-broadening understanding of how SARS-CoV-2 functions within a human host.

While the SARS-CoV-2 virus is very similar to other HCoV in several ways, including in its genomic structure and the structure of the virus itself, there are also some differences that may account for differences in the COVID-19 pandemic compared to the SARS and MERS of the past two decades. The R0 of SARS-CoV-2 has been estimated to be similar to SARS-CoV-1 but much higher than that of MERS-CoV [[209](#ref-sP4wQEiM),[209](#ref-sP4wQEiM)]. While the structures of the viruses are very similar, evolution among these species may account for differences in their transmissibility and virulence. For example, the acquisition of a furin cleavage site the S1/S2 boundary within the SARS-CoV-2 S protein may be associated with increased virulence. Additionally, concerns have been raised about the accumulation of mutations within the SARS-CoV-2 species itself, and whether these could influence virulence. The coming of age of genomic technologies has made these types of analyses feasible, and genomics research characterizing changes in SARS-CoV-2 along with temporal and spatial movement is likely to provide additional insights into whether within-species evolution influences the effect of the virus on the human host.

Additionally, omics technologies have made it feasible to characterize the host response to the virus. Although at present, much of the omics research has utilized *in vitro* models, such systems-level approaches represent a promising opportunity to characterize the host response to the SARS-CoV-2 virus. For example, analysis of PPI using publicly available data identified some similarities between COVID-19 and tuberculosis infection [[117](#ref-phJM8g2Y)]. This finding suggests that insights into COVID-19 may be gained by systems-level analysis of a wide array of infections, even those that are not causes by viruses, as is the case with the bacterium *M. tuberculosis*. As more data is collected, large-scale, omics-based analyses will become more feasible.

Thus, though the COVID-19 crisis is still evolving, the insights acquired over the past 20 years of HCoV research have provided a solid foundation for understanding the SARS-CoV-2 virus and the disease it causes. As the scientific community continues to respond to COVID-19 and to elucidate more of the relationships between viral pathogenesis, transmission, and symptomology, and as more data about the regulatory shifts associated with COVID-19 becomes available, this understanding will no doubt continue to develop to reveal additional interdisciplinary connections among virology, pathogenesis, and health. As additional information becomes available, this review will be updated to reflect the changing state of research in this area. At present, understanding the SARS-CoV-2 virus and its pathogenesis is critical to a holistic understanding of the COVID-19 pandemic.

# Additional Items

## Competing Interests

|  |  |  |
| --- | --- | --- |
| Author | Competing Interests | Last Reviewed |
| Adam L. MacLean | None | 2020-11-11 |
| Alexandra J. Lee | None | 2020-11-09 |
| Anthony Gitter | Filed a patent application with the Wisconsin Alumni Research Foundation related to classifying activated T cells | 2020-11-10 |
| Ashwin N. Skelly | None | 2020-11-11 |
| Casey S. Greene | None | 2020-11-07 |
| Christian Brueffer | Employee and shareholder of SAGA Diagnostics AB. | 2020-11-11 |
| David Mai | None | 2021-01-08 |
| Elizabeth Sell | None | 2020-11-11 |
| Gregory L Szeto | None | 2020-11-16 |
| Halie M. Rando | None | 2020-03-22 |
| James Brian Byrd | Funded by FastGrants to conduct a COVID-19-related clinical trial | 2020-11-12 |
| Jinhui Wang | None | 2020-04-13 |
| John J. Dziak | None | 2020-11-11 |
| John P. Barton | None | 2020-11-11 |
| Lamonica Shinholster | None | 2020-11-11 |
| Lucy D'Agostino McGowan | Received consulting fees from Acelity and Sanofi in the past five years | 2020-11-10 |
| Marouen Ben Guebila | None | 2020-11-11 |
| Nils Wellhausen | None | 2020-11-03 |
| Ronan Lordan | None | 2020-11-03 |
| Ryan Velazquez | None | 2020-11-10 |
| Sandipan Ray | None | 2020-11-11 |
| Sergey Knyazev | None | 2020-11-11 |
| Serghei Mangul | None | 2020-11-11 |
| Simina M. Boca | None | 2020-11-07 |
| Tiago Lubiana | None | 2020-11-11 |
| Vikas Bansal | None | 2020-05-26 |
| YoSon Park | None | 2020-11-11 |
| COVID-19 Review Consortium | None | 2021-01-16 |

## Author Contributions

|  |  |
| --- | --- |
| Author | Contributions |
| Adam L. MacLean | **MISSING** |
| Alexandra J. Lee | **MISSING** |
| Anthony Gitter | **MISSING** |
| Ashwin N. Skelly | **MISSING** |
| Casey S. Greene | X, Y, Z |
| Christian Brueffer | **MISSING** |
| David Mai | **MISSING** |
| Elizabeth Sell | **MISSING** |
| Gregory L Szeto | **MISSING** |
| Halie M. Rando | A, B, C, D |
| James Brian Byrd | **MISSING** |
| Jinhui Wang | **MISSING** |
| John J. Dziak | **MISSING** |
| John P. Barton | **MISSING** |
| Lamonica Shinholster | **MISSING** |
| Lucy D'Agostino McGowan | **MISSING** |
| Marouen Ben Guebila | **MISSING** |
| Nils Wellhausen | **MISSING** |
| Ronan Lordan | **MISSING** |
| Ryan Velazquez | **MISSING** |
| Sandipan Ray | **MISSING** |
| Sergey Knyazev | **MISSING** |
| Serghei Mangul | **MISSING** |
| Simina M. Boca | **MISSING** |
| Tiago Lubiana | **MISSING** |
| Vikas Bansal | **MISSING** |
| YoSon Park | **MISSING** |
| COVID-19 Review Consortium | Project Administration |

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DOI: [10.1146/annurev-virology-110615-042301](https://doi.org/10.1146/annurev-virology-110615-042301) · PMID: [27578435](https://www.ncbi.nlm.nih.gov/pubmed/27578435) · PMCID: [PMC5457962](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5457962)

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