Review Article

miRNAs: Key Molecules in the Immunopathogenesis of Betacoronaviruses

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Abstract

A series of patients hospitalized with acute respiratory disease was reported in Wuhan, Hubei Province, China, in December 2019. Many patients have had direct or indirect links with the Huanan Seafood Wholesale Market, Wuhan. Millions of people worldwide have been impacted by the 2019 coronavirus disease (COVID-19) in numerous nations. The pandemic has once again drawn public attention to the coronaviruses that developed epidemics in China (2002) and Saudi Arabia (2012). Given the structural and phylogenetic similarity of the 2019 novel coronavirus (2019-nCoV) with the severe acute respiratory syndrome coronavirus (SARS-CoV) and the Middle East respiratory syndrome coronavirus (MERS-CoV), the results of recent studies have been combined with new findings to complete one of the strangest pneumonia puzzles in human history. Coronaviruses establish extremely complex interactions with the immune system, especially in order to evade immune responses. Undoubtedly, increasing our knowledge of the immunopathogenesis of diseases caused by these viruses will eventually lead to more effective treatment and diagnosis. Non-coding RNAs (ncRNAs) are among the leading immune response regulators. MicroRNAs (miRNAs) play an important role in the expression and regulation of both innate and adaptive immune responses and in many immune disorders from autoimmunity to cancer and allergies. Our understanding of the functions of human and viral miRNAs in the pathogenesis of many viruses has increased in recent years. Accordingly, the present review article aims to review studies evaluating the role of miRNAs in the pathogenesis of other betacoronaviruses. The results of these studies, given the similarity of viruses within the family Coronaviridae, could be helpful for future research on SARS-CoV2.

Keywords: COVID19; SARS-CoV2; miRNA; SARS-CoV; MERS-CoV

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Introduction

The appearance of a solar corona, with its clubshaped spikes on its surface, inspired the name coronavirus. These viruses are potential pathogens for humans and vertebrates. They can infect the respiratory system, the gastrointestinal tract, the liver, and the central nervous system of humans, livestock, birds, bats, mice, and many other wild animals (1). Alpha- and Betacoronaviruses commonly infect mammals. In contrast, gam-

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This work is licensed under a Creative Commons Attribution-NonCommercial 4.0 International license (https://creativecommons.org/ licenses/by-nc/4.0/). Non-commercial uses of the work are permitted, provided the original work is properly cited. ma- and delta-coronaviruses infect birds and fish. called severe acute respiratory syndrome corona-These viruses can impose a huge economic burden because of their ability to infect livestock and pets (1). A bat coronavirus known as HKU2 caused the Swine Acute Diarrhea Syndrome coronavirus (SADS-CoV) in 2016, causing a large-scale naviruses is capable of causing severe respiratory outbreak and a deadly disease in pigs in southern China that eventually resulted in the loss of more tem (6). Most patients deceased from COVID-19 than 24,000 pigs (2). Before 2019, there were only in China were over 60 years old with a history six coronavirus strains known to infect humans and cause respiratory diseases. HCoV-229E, HCoV-OC43, HCoV-NL63, and HKU1 only tion, and cardiovascular disease (7). Clinically, cause mild upper respiratory disease, and some of them can rarely cause severe infection in infants, children, and adults (3). In addition, SARS-CoV and MERS-CoV can infect the lower respiratory tract and develop severe acute respiratory syndrome in humans (4). A series of acute respiratory diseases was reported in Wuhan, Hubei Province, China, in December 2019. Many patients have had direct or indirect links with the Huanan Seafood Wholesale Market, Wuhan, which is believed to be the primary source for the outbreak of the 2019 novel coronavirus (2019-nCoV) (5). The novel coronavirus was identified by the Chinese Center for Disease Control and Prevention (China CDC) as the cause of the disease, initially

virus 2 (SARS-CoV2). The World Health Organization (WHO) has now named the disease as Coronavirus Disease 2019 (COVID-19) (6). The novel Coronavirus 2019 belonging to Betacorosyndrome by engaging the lower respiratory sysof underlying diseases such as abdominal mass, chronic liver disease, myocarditis, renal dysfuncthe disease is accompanied by symptoms such as fever, cough, myalgia, fatigue, diarrhea pneumonia, and even death in severe cases (8-10). According to the phylogenetic tree, the SARS-CoV2 is closer to bat coronaviruses such as CoV ZC45 and SL-CoV ZXC21 and was more distant from SARS-CoV (11). The novel coronavirus sequencing shows that the SARS-CoV2 genome sequence is 96% homogeneous with the bat coronavirus. Sequence analysis indicates that the novel coronavirus has the typical genomic structure of coronaviruses and belongs to Betacoronaviruses, including the bat corona viruses of SL-ZC45, Bat-SL ZXC21, SARS-CoV and MERS-CoV (11). (Figure 1)

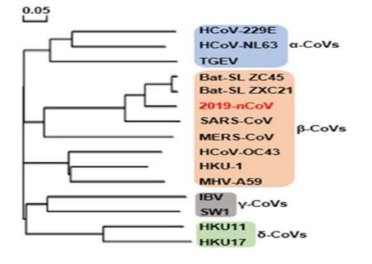


Figure 1. 2019-nCoV is highlighted in red. Genomic sequence analysis indicates that the novel coronavirus has the typical genomic structure of coronaviruses and belongs to Betacoronaviruses, including the bat coronaviruses of SL-ZC45, Bat-SL ZXC21, SARS-CoV, and MERS-CoV. According to the phylogenetic tree, 2019-nCov is closer to bat coronaviruses such as SL-ZC45 and SL-ZXC21 (with structural similarity up to 89%) and is more distant from SARS-CoV. Adapted from Chen, et al. Journal of Medical Virology 2020. (12).

Genomic structure and proliferation of corona- ous virus, 2019-nCov utilizes a unique furin-deviruses

The genome of coronaviruses contains a posi-Ebola viruses, which could be a justification for much higher infectivity (15, 16). tive-sense single-stranded RNA (+ssRNA) with ~30 kb in length and a 5' cap structure along with a 3' poly (A) tail. The viral genome has at least six Immunopathogenesis of Betacoronaviruses ORFs. The first ORF (ORF1a / b) comprises two-Studies have shown that important betacoronaviruses, such as MERS-CoV, SARS-CoV, and thirds of the viral genome encodes 16 non-structural proteins of the virus. Other ORFs are near SARS-CoV2 have extensive but complex interthe 3' ends of the virus genome encoding spike actions with the immune system (17-19). Studies (S), membrane (M) envelope (E), and nucleocaphave shown that the betacoronaviruses are able to sid (N) proteins. In addition to the four major disrupt IFNI-dependent antiviral responses (20). structural proteins, various coronaviruses encode These viruses prevent IFNI production by blockspecific structural and additional proteins such as ing the identification of their PAMPs by innate HE protein, 3a / b protein, and 4a / b protein. In immune-specific PRRs such as RIG1 and MDA5, the genome of various coronaviruses, the regions on the other hand, and by utilizing their NSPs to block the signaling pathways of these cytokines encoding non-structural proteins (NSPs) have 58% overlap, and the genomic regions encoding from IFNIR receptor to downstream transcripstructural proteins have up to 43% overlap. This tion factors of IRF and STAT1 (20). In addition, suggests that NSPs are more conserved, whereas betacoronaviruses have been shown to increase structural proteins have more diversity to adapt cell resistance to IFNI (20). Examination of specto novel hosts. Since the mutation rate in RNA imens from patients with SARS-CoV and MERSviruses is much higher than that of DNA virus-CoV reveals an accumulation of macrophages es, the genomes of RNA viruses are usually less in the lung tissue (9). These viruses, by infecting than 10 kb in length but coronaviruses are more macrophages on the one hand and stimulating than 30 kb in length. Thus, the coronavirus is the inflammasome and other inflammatory response largest RNA virus ever known. Coronaviruses are pathways on the other hand, impair the function different from all RNA viruses because of their of macrophages and enhance the production of so-called proof-reading properties, which is due proinflammatory cytokines such as TNF, IL6, to the ability of the 3'-5 'exoribonucleases of these and IL1, as well as overexpress the inflammatory chemokines such as CXCL10, CCXL10 and, viruses (11). CCL5 and CCL8 on their surface (21, 22). Studies The penetration of the virus into host cells have shown that T lymphocytes are significantly Penetration into host cells is an essential step in reduced in the peripheral blood of patients inthe transmission of various strains of coronavirus, fected with SARS-CoV2 and Findings reported especially for betacoronaviruses. All coronavirussevere lymphopenia in patients with COVID-19. es encode the spike (S) glycoprotein. Part of the The lymphopenia has been reported to be much S protein that interacts with the host cell surface more severe in the ICU patients with severe forms of the infection. In contrast, it has been found protein is called the receptor-binding domain. Upon receptor binding, the adjacent host cell prothat the T-cell counts increase and return to their tease cleaves the viral S protein, which releases normal levels in patients who are in the recovery the S fusion peptide and facilitates virus penetraphase (7, 8).

tion (13). Receptors of betacoronaviruses known SARS-CoV directly infects T lymphocytes so far include Angiotensin-converting enzyme-2 and causes lymphopenia, splenic atrophy, and lymph node atrophy. It has also been shown that (ACE2) for the SARS virus and dipeptidyl peptidase-4 or CD26 for MERS virus, which binds to MERS-CoV is capable of activating intrinsic and host cell receptors and mediates viral penetration. extrinsic apoptotic pathways in T lymphocytes. The 2019-nCov receptor is also ACE2 (13, 14). The virus appears to not only develop lympho-The affinity of 2019-nCov is higher than that of penia but also impair the function of normal T SARS-CoVs. On the other hand, unlike the previlymphocytes. A study reported that the T lym-

pendent mechanism previously seen for HIV and

phocytes in the patients with COVID-19 were The miRNA and immune system over-controlled and over-activated, confirmed by the high proportion of lymphocytes expressing the HLA-DR (CD4 3.47) and CD38 (CD8 39.4) markers (23). This over-activation process appears to result in T-lymphocyte exhaustion in the patients, as a study examined 522 Chinese patients in Wuhan for the expression of T-cell exhaustion markers such as PD-1 and Tim-3 using range of bioactivities in the body. The synthesis of flow cytometry (24). The results showed that their expression was significantly increased in patients with COVID-19 compared to controls. The increased expression of these markers is associated with the shift of disease to symptomatic stages. Even if no antigen is present, the memory CD4+ and CD8+ T cells can persist for four years in a number of subjects recovered from SARS-CoV infection and be able to perform T cell proliferation, DTH response, and IFN-γ production. Six years after SARS-CoV infection, the SARS-CoV S peptide-specific T-cell responses were detected in 14 of 23 patients recovered from SARS (25). In summary, recent studies have shown that SARS-CoV2 reduces, over-activates, and eventually exhausts the T lymphocytes; however, memory lymphocytes also seem to remain in the patient's investigating changes in miRNAs in the pathobody for years.

Finally, the cytokine storm is one of the reported causes of mortality in patients with COVID-19. Therefore, the use of the anti-rheumatoid drug Actemra in China has already shown promising results in the treatment of patients with COVID-19. MERS-CoV, SARS-CoV, and 2019-nCOV are able to widely modify the process of developing immune responses, which is undoubtedly caused by changes in gene expression profiles or, in other words, changes in gene in SARS-CoV and MERS-CoV has been investiexpression regulation. In recent years, more than ever it has been shown that epigenetic modifications play an undeniable role in the regulation The Coronavirus N protein is one of the structural and homeostasis of immune responses (26). The ncRNAs also possess a prominent function (27).

is being seriously studied in a variety of immune disorders such as autoimmune diseases, cancers, autoinflammatory syndromes, allergies, and infectious diseases to understand more precisely mechanisms of pathogenesis and ultimately their more effective diagnosis and treatment (28-30).

The ncRNAs act as regulatory molecules and do not encode a protein. Studies revealed that only 2% of mammalian genomes contain protein-coding genes and a large percentage of genomes account for the ncRNAs (31). The miR-NAs are a major class of ncRNAs that are short in length (19-23 nucleotides) and regulate a wide miRNAs is a complex and multistep process (32). Reportedly, the miRNAs play a central role in immune cell plasticity, especially the differentiation of T helper cells, such as TH17 and TH1, TH2, TH9, TH35, and Treg (33). The regulation of producing cytokines and their signaling pathways, including IFNI and proinflammatory cytokines that play important roles in the pathogenesis of coronaviruses, are all regulated by epigenetic modifications such as miRNAs (34). Studies have shown well that the differentiation and function of macrophages M1, M2, and Mreg depend on alterations in miRNA expression (35). Given the widespread interaction of MERS-CoV, SARS-CoV, and 2019-nCov with the immune system, this review article aimed to investigate the studies genesis of these diseases. Given the similarity of viruses in the family Coronaviridae, the results of these studies could be helpful for future research into the 2019-nCoV.

The miRNAs in coronavirus infections

There have been few in vivo studies on the role of miRNAs in coronavirus infections, but this issue has been investigated in HCoV-OC43 in the in vitro condition. In addition, the role of miRNAs gated using bioinformatics tools. The OC43 virus causes the common cold around the world (36). proteins of the virus that binds to genomic RNA to facilitate virus proliferation as well as to form a helical capsid. It acts as a potent stimulus by bind-Thus, the function of these types of ncRNAs ing to the negative inhibitor of NF-κB, miR-9. NF-kB is one of the most important transcription factors in the immune system. It is not yet clear whether NF-kB activation is beneficial or harmful to the virus, whether it is directly beneficial for viral proliferation, and whether it is a random effect that ultimately limits the viral pathogenicity. However, the low pathogenicity of OC43 can ment of patients with COVID-19. Roche Compabe investigated in other respects. Accordingly, the ny is working with the FDA to start a randomreduction in OC43 pathogenicity, which is assoized, double-blind, placebo-controlled clinical trial to study more closely the efficacy of Actemra ciated with more limited clinical symptoms, may result in increased contact between infected and in hospitalized patients with COVID-19. Therefore, can intervention of the virus mechanism in non-infected individuals, resulting in the spread of the virus in a population when compared with the case of miR9 slightly reduce the intensity of the cytokine storm? Does SARS-CoV2 also target coronaviruses with higher pathogenicity (37). miR9? Bioinformatics and empirical studies can This finding is important because the binding of virus proteins to host miRNAs and altering gene answer all of these questions. expression may be a clever strategy for coronaviruses to evade immune responses more effective-Effect of SARS-CoV on the expression of miRly (38). Importantly, understanding the mecha-NA in Bronchoalveolar stem cells (BASCs) nisms of escape from immune responses can also Bronchoalveolar stem cells (BASCs, including help to treat COVID-19 more effectively. Previ-Sca-1+ CD34+ CD45- Pecam-) are among the ous studies in patients with COVID-19 show well cells targeted and infected with SARS-CoV. The the high ability of SARS-CoV2 to inhibit antiviral viruses can downregulate miR-223 and miR-98 in responses and avoid their damage. Coronaviruses BASCs, thereby controlling several different stages of their differentiation and production of their take advantage of the reduced rate of proliferation within cells infected with the virus at the onset of anti-inflammatory cytokines. Viral nucleocapsid and spike proteins appear to downregulate miRinfection as a strategy to evade immune responses so that they can infect more cells. In the cells 223 and miR-98 in the BASCs, simultaneously, affected by the coronavirus, the involvement of to control different stages of BASC differentia-PRRs such as RIG1 and MDA5 provokes intration, activation of inflammatory chemokines, and cellular signaling cascades, thereby enhancing the ACE2 inhibition. These expression changes in the expression of NFKB1 and miR-9. It should be notmiRNAs by the virus actually play a dual role beed that the miR-9 targets the NFKB1 mRNA and cause, on the one hand, these changes facilitate thus inhibits the translation of NFKB; however, easier viral cell-to-cell transmission, and on the other hand, disrupt the BASCs to drastically rethis outcome is inhibited by the activity of OC43, duce the ability of affected lung tissue regenerawhich binds to miR-9 and consequently increastion. Taken together, this study demonstrates yet es the translation of NFKB1. The translation of another clever strategy of the virus in how it uti-NFKB1 itself further leads to higher expression of lizes host cell miRNAs for its advantage (39). proinflammatory cytokines (37), whose over-expression is one of the major contributing factors to the immunopathogenesis of ARDS in patients. SARS-CoV and the expression of svRNA It should be noted that the occurrence of the cy-Three svRNAs including nsp3 (svRNA-nsp3.1 tokine storm is one of the major pathological and and -nsp3.2) and N (svRNA-N) were obtained from the genomic regions of SARS-CoV. The biodetrimental events in SARS-CoV and 2019-nCoV. genesis of CoV svRNAs is independent of RNase High levels of proinflammatory and suppressive III, cell type and host species, but is dependent cytokines such as IL-2, IL-7, G-CSF, IP-10, MCP-1, MIP-1A, and TNFa have been reported in the on the rate of viral proliferation. The inhibition acute cases of the disease, which is termed the of svRNA-N using Antagomir tools significantcytokine storm. In the patients with COVID-19 ly correlated with a decrease in lung pathology induced by SARS-CoV2, a study published in the and proinflammatory cytokine expression. Takauthoritative journal of Lancet reported that the en together, these data suggest that the svRNAs deaths of 6 out of 41 patients were due to cytokine contribute to the pathogenesis of SARS-CoV and storm, subsequently the ARDS and ultimately the therefore their inhibition by Antagomir could be dysfunction of several vital organs of these pa- a potential therapeutic approach in the infection tients. Accordingly, the use of Actemra in China with these viruses (40). As previously described, has already shown promising results in the treat- SARS-CoV2 is phylogenetically and structurally

more than 80% similar to SARS-CoV and to a quent decrease in the expression of CASK-inlesser extent similar to MERS-CoV. Therefore, it is not out of the question that multiple svRNAs could be used as effective factors in the pathogenesis and invasion of SARS-CoV2 and ultimately as cell proliferation, migration, invasion and mein the worsening status of COVID19 patients. Currently, more than 110 clinical trials in China and other countries are evaluating the efficacy of a ies and can be used as a target for the treatment variety of drugs, monoclonal antibodies, vaccines of viral diseases. Overexpression of miR-21a-5p and immune cell therapy in COVID19, so why or deletion of Caskin1 in the host significantly should we not test our chances of evaluating the inhibition of svRNAs in SARS-CoV2 infection.

The miRNAs in porcine hemagglutinating encephalomyelitis virus (PHEV) infection

The porcine hemagglutinating encephalomyelitis virus (PHEV) is another Betacoronaviruse that causes neurological and / or digestive disease in pigs. The PHEV was the first coronavirus identified in and isolated from pigs and is the only neurotropic virus ever identified in pigs so far. The first epidemic caused by the virus was reported in 1957 in Ontario, Canada. Studies have shown that the miRNAs also play a prominent role in the pathogenesis of the virus (Table 1) (41). A study found that miR-10a-5p suppresses The miR-221-5p in porcine epidemic diarrhea the downstream Syndecan 1 gene and acts as an antiviral mechanism in PHEV-induced disease. The Syndecan 1 is a cell surface proteoglycan that interacts with extracellular matrix molecules and growth factors to maintain epithelial cell morphology, anchor protein-dependent growth, and invasive inhibition in cell culture (42). Another study reported a significant increase in the expression level of miR-21a-5p in the rat brain as well as PHEV-infected N2a cells and a conse-

teracting protein 1 (Caskin1). It should be noted that the miR-21 regulates the expression of target genes involved in several cellular processes such tastasis. The role of miR-21 in the viral infection process has been confirmed in a number of studcontributes to PHEV proliferation. In contrast, the miR-21a-5p silencing by miR-21a 5p inhibitors results in viral suppression. Altogether, the results of this study indicate that the Caskin1 gene is a direct target of miR-21a-5p, and helps to increase virus proliferation by suppressing the Caskin expression. This illustrates well how a high-virulence beta-coronavirus is using miRNA to overwhelm the host immune system, and this raises the possibility of replicating such a strategy in the pathogenicity of other coronaviruses, especially SARS-CoV2, which we are now seeing as its pandemic, and these findings may help to develop strategies for therapeutic applications (43).

virus (PEDV) infection

The porcine epidemic diarrhea virus (PEDV), the causative agent of porcine epidemic diarrhea, causes significant economic burden to the pig industry around the world. The researchers found that overexpression of miR-221-5p inhibits the dose-dependent PEDV proliferation and interestingly reduced expression of miR-221-5p increases PEDV proliferation. It was subsequently found that the miR-221-5p directly targets the

Coronavirus	miRNA	Target genes	Functional activity	Ref
PHEV	miR-142a-3p	miR-142a-3p bind directly bound to the 3'UTR of Rab3a	miR-142a-3p promotes PHEV proliferation by directly targeting Rab3a mRNA.	(41)
PHEV	miR-10a-5p	Syndecan 1, a cell surface proteoglycan	miR-10a-5p leads to downstream suppression of Syndecan 1, and it functions as an antiviral mechanism in the PHEV-induced disease.	(42)
PHEV	miR-21a-5p	Caskin1	Over-expression of miR-21a-5p or Caskin1 knockdown in the host significantly contributes to PHEV proliferation.	(43)
PEDV	miRNA-221-5p	NF-kappaB-inhibitor alpha and suppressor of cytokine signaling 1.	miR-221-5p directly targets the 3' UTR of PEDV genomic RNA to inhibit PEDV proliferation.	(47)

Table 1. Deregulation of miRNA expression in PHEV and PEDV

Betacoronaviruses have posed serious chalgenomic RNA of PEDV and in turn activates the NFKB signaling pathway (44). The results of this lenges to the healthcare systems of the internastudy are certainly amazing. The host miRNA tional community in recent years. Previously, SARS-CoV and MERS-CoV have caused widedirectly targets the betacronavirus RNA and simultaneously alters one of the major intracellular spread epidemics that have imposed heavy fisignaling pathways involved in the exacerbation nancial burdens on national healthcare systems, of inflammation. These findings raise some key but SARS-CoV2 is undoubtedly one of the most questions: Are miRNAs similarly able to target complex viruses that has ever challenged mandirectly the genomic RNA of SARS-CoV2 in the kind. Although it is only about three months human body? And prevent its proliferation? And since the birth of SARS-CoV2, it has caused a at the same time affect key signaling pathways pandemic and involved more than 150 countries, in the emergence of immune responses? What infecting more than 250,000 people and killing changes does the virus make in the expression tens of thousands of people. The promising point level of this miRNA? Ultimately, the existence of is that SARS-CoV2 is structurally similar to its such a strategy can lead to the development of exother counterparts in the family Coronaviridae. tremely effective treatment strategies. The 96% similarity with bat Betacoronaviruses and the 80% similarity with SARS-CoV has led Bioinformatics studies of miRNAs in coronavimany of the drugs previously evaluated in acute respiratory diseases developed by MERS and Bioinformatics studies on the interaction of vi-SARS to be rapidly evaluated today to control SARS-CoV2 in over 105 clinical trials. Approxral mRNA and miRNA have suggested the idea that SARS-CoV may evade immune responses by imately 40 large and small pharmaceutical companies are currently trying to evaluate a variety increasing the expression of miR-17, miR-574of therapies, from immune cell therapy to the use 5p, and miR-214 and subsequently reducing their of nucleoside analogues and protease inhibitors, proliferation at the onset of infection. It should be noted that these host miRNAs target all four and monoclonal antibodies. Although the gestructural proteins of the virus called spike (S), nome sequence of the novel coronavirus has been nucleocapsid (N), envelope (E), and matrix (M) rapidly determined, the mechanism of its pathoproteins (45). These results demonstrate how genesis remains unknown. Undoubtedly, a better understanding of immunopathogenesis and the SARS-CoV may alter host miRNA expression interaction between the immune system and the profiles. In another study, bioinformatics studies have shown that some miRNAs such as miR-628virus can be extremely useful in this regard. Un-5p, miR-6804-3p, miR-4289, miR-208a-3p, miRderstanding the immune responses in patients 510-3p, miR-18a-3p miR-329-3p, miR-548ax, with severe forms of COVID-19 may lead to sigmiR-3934-5p, miR-4474-5p, miR-7974, miRnificant progress in the development of effective 6865-5p, and miR-342-3p were strongly similar therapies. The miRNAs are the regulators of imin hairpin structure to the MERS-CoV genome. mune responses. The most prominent feature of Therefore, all of these miRNAs may possibly help the human immune system is plasticity and spethe virus to evade the immune response and evencialized function, and this can only be achieved tually infect more cells by reducing virus proliferby rigorous regulation while also being flexible ation within the infected cell (46). In addition, in in gene expression through a variety of epigenetic mechanisms, especially miRNAs. These small vitro studies should be performed to evaluate the inhibitory effect on viral proliferation through but extremely effective non-coding RNAs play an the effect of specific human miRNAs. Let us just important role in the interaction between the imnot forget that it is only three months since the mune system and infectious agents. There is cur-SARS-CoV2 came out. Therefore, the role of host rently no disease in which the key role of miRNAs miRNAs as well as the virus and the complex inhas not been investigated and detected. Therefore, investigating the role of host miRNAs as well as teraction between them and the immune system must be carefully evaluated, which will definitely svRNAs that are specifically involved in regulathelp to better understand the immunopathogening immune responses should be prioritized for a

ruses

infection. Can one imagine that evading the virus from immune responses, such as disruption mune response can help accelerate access to novel of the antiviral system of type I interferon, initiation of destructive cytokine storm, induction of severe lymphopenia, and T-lymphocyte exhaustion through increased expression of Tim3 and LAG3 on them, recruitment of macrophages and of interest. dendritic cells and impaired lymphocyte differentiation, could be practically without extensive References alterations of miRNAs? Therefore, the current review article presents the role of miRNAs in the pathogenesis and immunopathogenesis of other pathogenic betacoronaviruses in humans and other species, and how the virus uses miRNAs to overcome immune system and exacerbate its infectivity, as well as how alterations in miRNA expression can alter virus infection. Therefore, in situations where excessive costs are spent on clinical trials to find effective treatments and vaccines for SARS-CoV2, research on the role of miRNAs in pathogenicity, and in particular virus-host interactions, could lead to effective treatment strategies or adjunctive therapies in patients with COVID-19. Studies in recent years have focused on identifying miRNAs as therapeutic targets for viral diseases, and there are promising clinical trials, including antagomirs that target host miR-NAs such as miR-122 in hepatitis C. However, it should be kept in mind that changes in the level of the host cell miRNA can cause multiple abnormalities in different biological pathways due to the multiple regulatory roles of the miRNA. However, small RNAs have been identified in the genomes of coronaviruses such as SARS-CoV2, which are able to target important human genes and can be a therapeutic target in these viruses.

Conclusion

The inhibitors of viral miRNAs can be considered as effective antiviral therapies. Targeting viral miRNAs with Antagomirs specifically in the 9. infected cell can be a therapeutic candidate in this field. In spite of all the promising evidence, in vivo, in vitro, and ex vivo studies are necessary to evaluate the efficacy of miRNAs as targeted therapy in coronaviruses. Increasing levels of viral miRNAs in the host may alter conditions favoring the immune system for virus control. Undoubt-

deep and accurate understanding of SARS-CoV2 edly increasing our understanding of the molecular details governing the viral life cycle and imantiviral therapies.

Conflict of Interest

The authors declare that they have no conflict

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