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***In silico* Analysis of Human miRNAs in SARS-CoV-2 Genome**

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Abstract

In December 2019, a new coronavirus (SARS-CoV-2) was discovered in Wuhan (China) that was rapidly transmitted to many other countries. Henceforth, the World Health Organization (WHO) Emergency Committee declared a global health emergency on January 30, 2020. Statistics depicted the fatality rate as about 1.4%. In this study, a potential antiviral treatment for the SARS-CoV-2 virus using host miRNAs was explored which may slow down the expression of viral genes to suppress viral replication. The miRNAs from genome (coronavirus / SARS-CoV-2) were analyzed using various computational approaches. The complete genome sequence was retrieved from NCBI. The result of our study highlighted that hsa-miR-3675-3p (MD19), hsa-miR-363-5p (MD220), hsa-miR-325 (MD306), hsa-miR-2114-5p (MD306), hsa-miR-744-3p (MR186) and hsa-miR-448 (MR186) can be used as an antiviral treatment to quell the replication of SARS-CoV-2 virus in human beings. The findings and observations of our study opened new possibilities to explore both the pathogenesis function of miRNAs and for the development of novel antiviral drugs.

1. Introduction

In December 2019, a new coronavirus (SARS-CoV-2) emerged in Wuhan (China) and rapidly spread to many other countries [1–3]. The World Health Organization (WHO) Emergency Committee declared a global health emergency on Jan. 30, 2020, based on growing case rates. As of 24 April 2020, 177,108,695 individuals were infected by SARS-CoV-2 worldwide and 3,840,223 people died because of it. As of 19 June 2021, a total of 2,412,226,768 vaccine doses have been administered, worldwide [4]. SARS-CoV-2 is listed as a top category pathogen by several organizations including WHO, CDC and NIH because its fatality rate is up to 1.4%

[5–7]. Clinical signs of SARS-CoV-2 closely resemble those seen in MERS and SARS infections [8, 9]. A recent report [10] indicated that the potential source(s) has not been identified yet which caused the transmission of the virus to human beings.

Coronaviruses have a monopartite plus-strand RNA genome and belong to the *Coronaviridae*, often pleiomorphic virions, with a diameter of approximately 80 to 120 nm [5]. Coronaviruses contain a positive, capped and polyadenylated RNA strand with the largest genomic RNA (approximately 27 – 32 kb) in size that causes respiratory, gastrointestinal, hepatic, and neurologic diseases in human beings and animals [5, 11, 12]. The “N”

protein was found to be bound to viral RNA and packaged into ribonucleoprotein complexes, which are located at the viral membrane's internal face [13]. Coronaviruses have at least three viral proteins in their membrane including a) Spike (S), which reveals the structure of the virus as a crown; b) membrane protein (M), coated three times and provided with a small N-terminal ecto-domain; and c) a hydrophobic protein, that is, a cytoplasmic tail and a small membrane protein (E) [14, 15]. So far, the presence of coronaviruses has been identified in mice, rats, pigs, cats, rabbits, horses and livestock, causing a number of serious diseases including gastroenteritis and respiratory tract problems [16].

MicroRNAs (miRNAs) are single stranded RNAs (ssRNAs), around 18 – 25 nucleotides long that modulate protein-coding genes [17, 18]. Introns of protein coding genes, UTR of protein coding genes, exons of non-coding genes, and introns of non-coding genes are all sites where miRNAs can be found [19, 20]. It is well documented that miRNAs perform different biological or physiological functions including apoptosis, development, tumorigenesis, stress response, proliferation and fat metabolism [21, 22]. RNA polymerase II are generally used to make miRNAs [23]. Main Drosha converts the main transcript into a hairpin pre-miRNA using RNase III enzyme along with the dsRNA binding protein [24, 25], exporting 5/Ran-GTP transport to the nuclear pre-miRNA, which is then cleaved by the cytoplasmic RNase III Dicer to create an incomplete 21 – 25 nucleotide dsRNA [26, 27]. In the RNA Induced Silencing Complex (RISC), a strand known as the mature miRNA strand is loaded and RISC is guided to target it

where it hybridizes with complementary sequences, causing cleavage or translational inhibition. The presence of viral miRNA is associated with the role of virus infection, as indicated by numerous researches. Additionally, emerging evidence has confirmed the connection of viral miRNAs with human diseases [28, 29].

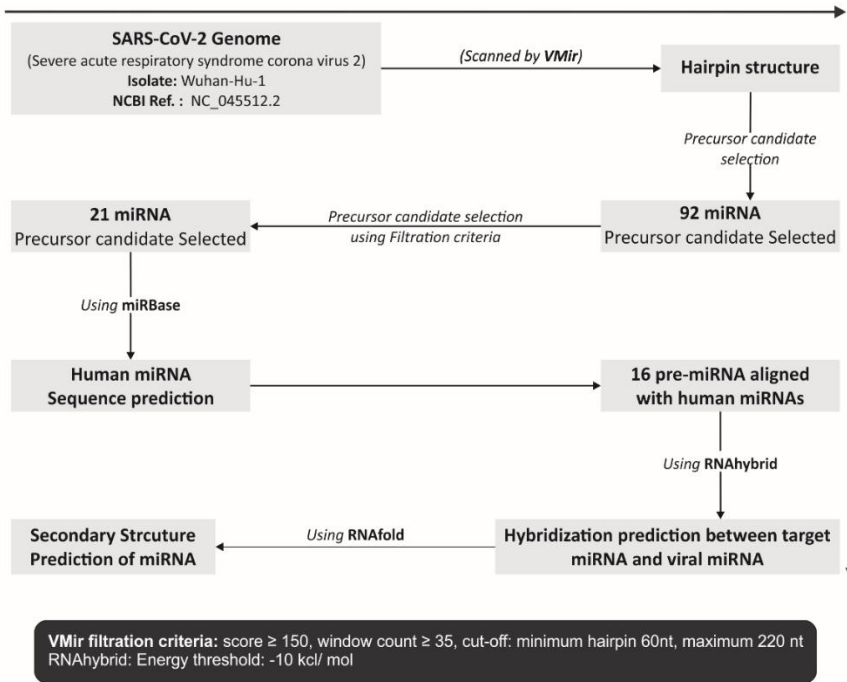
Viral miRNAs were found to alter the life cycle of a virus and also affect its survival in hosts [30, 31]. Significantly, viral miRNAs can target not only the virus but also the host's miRNA regulation. Identifying viral miRNAs using bioinformatics technologies and techniques is, therefore, an evolving approach to explore the mechanisms of virus-host interaction [30, 32].

In this study, we utilized various computational methods and techniques including the RNA-hybridization technique to identify the potential targets of human microRNAs of the SARS-CoV-2 genome. This study aids in enhancing the understanding of host-pathogen interactions as well as the development of new antiviral therapies for all SARS-CoV-2 strains.

2. Materials and Methods

2.1. Data Retrieval

The complete SARS-CoV-2 genome sequence was obtained from the National Center for Biotechnology Information (NCBI) (Isolate: Wuhan-Hu-1, NCBI Reference Sequence: NC_045512.2) and used for miRNA prediction. Figure 1 shows the overall workflow used in this study. Complete genome isolate from Wuhan-Hu-1, revised by authors on 30 March 2020, contains 29903 bp ss-RNA.



Workflow: *In-silico* approach for anticipating role of Human miRNAs

Figure 1. Human miRNA prediction of SARS-CoV-2 workflow (Isolate Wuhan-Hu-1)

2.2. Hairpin-structured miRNA Precursors Prediction – Pre-miRNA Extraction

A flowchart (Figure 1) describes the computational prediction of miRNA precursors. In our study, VMir Analyzer tool was used to search the genome for experimentally confirmed hairpin-structure miRNAs precursor [33-35]. Predictions for VMir were performed using default parameters. For further investigation, pre-miRNAs having a VMir score of less than and equal to 150 (Window Count, WC = 35) were chosen. VMir Viewer was used for the visualization of scanned hairpins [35, 36, 37].

2.3. Human miRNAs Sequence Prediction

The miRbase database contains the sequences of human miRNAs (<http://www.mirbase.org/search.shtml>) [38]. The genome nucleotide segment under analysis was scanned using the

VMir tool and each segment’s input and nucleotide similarity to all human microRNAs was extensively analyzed using blast program in miRbase search tool.

2.4. Hybridization Prediction between Target miRNA and Viral miRNA

RNAhybrid predicts miRNA based on the minimum free energy and site complementarity. RNAhybrid is also used in viral genome to locate the exact match for miRNA target [39]. RNAhybrid (<https://bibiserv.cebitec.uni-bielefeld.de/rnahybrid>) was utilized to evaluate miRNAs against SARS-CoV-2 genome attachment at an energy threshold of -10 kcl/ mol and other filters were set to default parameters. The tool identified some minimum free energy miRNA precursors that deviated from the threshold values, so these were removed from the final list. RNAhybrid’s result was

categorized in terms of pairing energy and pattern hybridization.

2.5. Secondary Structure Prediction of miRNA

The structure of pre-miRNAs was predicted using the online server “RNAfold” with default parameters (<http://rna.tbi.univie.ac.at/cgi-bin/RNAWebSuite/RNAfold.cgi>) [40].

3. Results

3.1. miRNA Precursor (Pre-miRNA) Hairpins Prediction

The viral genome was screened and visualized using VMIR Analyzer and VMIR Viewer, respectively. VMIR

Viewer displays the entire output graphically, including the sequence score and length. Figure 2 illustrates the visual representation of miRNA hairpin precursor of SARS-CoV-2 genome. We filtered 92 candidate hairpins through the default parameters of VMir Analyzer tool as shown in Figure 2(a). A filter using specific parameters and custom configuration, that is, for minimum hairpin score of 150, minimum window count of 35, a minimum cut-off value of 60 nt for hairpin size, and a maximum hairpin size of 220 nt was applied to avoid the bona fide candidate hairpin. Finally, for further study, 21 pre-miRNA hairpins were chosen as candidate hairpins (Figure 2).

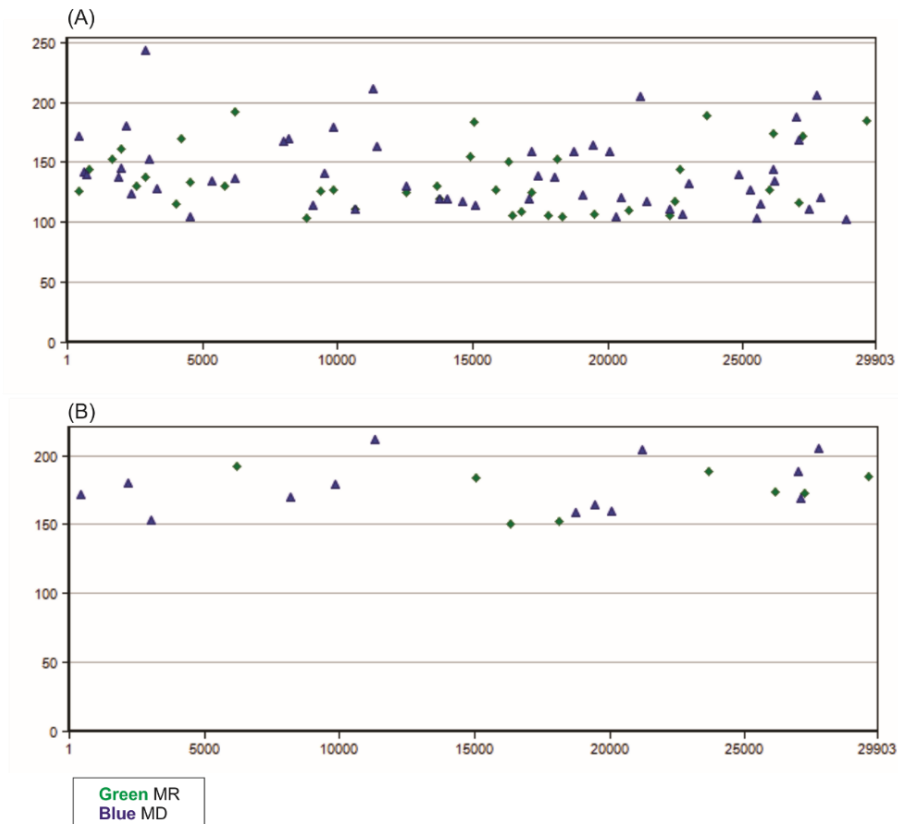


Figure 2. VMir analysis of SARS-CoV-2 genome is represented graphically (A) default setting was used to display all pre-miRNA hairpins (B) representation of predicated pre-miRNA after filtration

3.2. Human miRNAs Prediction from miRNAs Precursor (Hairpin)

The nucleotide similarity of 21 candidate miRNAs, precursors to human miRNAs, was searched using miRBase database (<http://www.mirbase.org/search.shtml>) (Table 1). Based on the considerable sequence similarities with human miRNAs, 16 precursors were selected as probable miRNA precursor candidates. Human miRNAs were identified as the primary target miRNAs with a similarity of at least 10 bp sequence with candidate miRNA precursors. Then, in the candidate miRNA precursors' 3' untranslated region, near or almost perfect alignment of all those miRNAs' seed areas (2 – 8) were possible miRNA targets. Perfect complementary matching between 3' untranslated region (UTR) of miRNA and

the seed region of miRNA is essential for the fruitful cleavage of miRNA or translational repression. Viral precursors miRNA hairpins MD3, MD19, MD29, MD134, MD220, MD228, MD240, MD306, MD307, MD311, MR155, MR165, MR186, MR243, MR274, MR304 showed a significant similarity with hsa-miR-4471, hsa-miR-3675-3p, (hsa-miR-383-5p and hsa-miR-5197-3p), hsa-miR-190b-5p, hsa-miR-363-5p, hsa-miR-4802-3p, hsa-miR-23b-5p, (hsa-miR-325 and hsa-miR-2114-5p), (hsa-miR-215-3p, hsa-miR-548y, hsa-miR-338-3p and hsa-miR-3065-5p), hsa-miR-4699-3p and hsa-miR-6739-3p), hsa-miR-363-5p, hsa-miR-153-5p, (hsa-miR-744-3p, hsa-miR-4420 and hsa-miR-448), hsa-miR-4796-5p, hsa-miR-6867-5p, (hsa-miR-3064-5p and hsa-miR-411-5p), respectively.

Table 1. miRNAs Hairpin Precursor Sequence and Human miRNAs

S. No	Hairpin	Score	Alignment (SARS-CoV-2 and Human microRNA)
1	MD3	62	UserSeq 24 cuuaguagauguugaa 39
			hsa-miR-4471 7 cuuaguagaguuuaa 22
2	MD19	62	UserSeq 77 gaguuucuuagagacg 92
			hsa-miR-3675-3p 16 gaguuccuuagagaug 1
3	MD29	70	UserSeq 81 agaaggugauugug 94
		70	hsa-miR-383-5p 6 agaaggugauugug 19 UserSeq 98 aagaagaguugagccaaucaacu 120
4	MD134	69	hsa-miR-5197-3p 1 aagaagagacugagucaucgaa 23 UserSeq 34 gauaugguugauacuaguuug 54
			hsa-miR-190b-5p 2 gauauguuugauauuggguug 22 UserSeq 30 cguguauaacacguugcaauuu 51
5	MD220	65	hsa-miR-363-5p 1 cggguggaucacgaugcaauuu 22 UserSeq 62 uaaaagguuuacaaccau 79
			hsa-miR-4802-3p 21 uugaagguuuccauccau 4 UserSeq 50 uucuuggaaucgugaucu 67
7	MD240	63	hsa-miR-23b-5p 5 uuccuggaucgugauuu 22



S. No	Hairpin	Score	Alignment (SARS-CoV-2 and Human microRNA)
8	MD306	68	UserSeq 60 uugcuggacaccaucuagg 78
			hsa-miR-325 19 uuacuggacaccuacuagg 1
			UserSeq 1 accgcuucuagaaguga 18
		63	 hsa-miR-2114-5p 21 accgcuucaaggaaggga 4
		UserSeq 20 gccuaaagaaucacug 36	
		67	 hsa-miR-215-3p 18 gccuaaagaaugacag 2
		UserSeq 24 aaagaaucacuguu 38	
9	MD307	66	 hsa-miR-548y 2 aaaguaaucacuguu 16
			UserSeq 24 aaagaaucacugugcu 41
		63	 hsa-miR-338-3p 21 aacaaaucacugaugcu 4
			UserSeq 24 aaagaaucacugugcu 41
		63	 hsa-miR-3065-5p 3 aacaaaucacugaugcu 20
		UserSeq 13 acuuuacucuccaauuuu 30	
10	MD311	63	 hsa-miR-4699-3p 1 aauuuacucugcaauuu 18
			UserSeq 80 agaaagacagaaugau 95
		62	 hsa-miR-6739-3p 16 agaaagacagaacaau 1
11	MR155	68	 hsa-miR-363-5p 22 aaauugcaucugauccac 4
		UserSeq 37 uauuuuugugauguug 53	
12	MR165	67	 hsa-miR-153-5p 1 ucauuuuugugauguug 17
			UserSeq 50 uugaauuuagugucaaca 67
		63	 hsa-miR-744-3p 19 uugagguuaguggcaaca 2
13	MR186	63	 hsa-miR-4420 3 cacugaugucuguagcug 20
			UserSeq 82 ugccuguguaggau 97
		62	 hsa-miR-448 2 ugcauauugaugu 17
14	MR243	68	 hsa-miR-4796-5p 19 aagucagagauagaca 1
			UserSeq 47 uguagaugaagaaggua 63
15	MR274	67	 hsa-miR-6867-5p 7 uguagaggaagaaggga 23

S. No	Hairpin	Score	Alignment (SARS-CoV-2 and Human microRNA)
16	MR304	64	UserSeq 68 ucugccuuguguggucugca 87
			hsa-miR-3064-5p 1 ucuggcuguuguggugugca 20
			UserSeq 19 aguagacuaauauaucgua 36
		63	hsa-miR-411-5p 2 aguagaccguauagcgua 19

3.3. Hybridization between Viral Precursor miRNAs and Human miRNAs

RNAhybrid (https://bibiserv.cebitec.uni-bielefeld.de/rnahybrid?id=rnahybrid-view_submission), was utilized for the successful and positive hybridization of the human target miRNAs and miRNA

precursors of SARS-CoV-2 measurement. Pairing energy revealed hybridization stability (or minimum free energy). A cut-off score was used to pick the potential miRNAs at -10 kcal/mol. In a large viral RNA hairpin, this software discovered the most energetically favorable miRNA hybridization sites. Effective hybridizations are shown in Table 2.

Table 2. MicroRNA and Viral RNA Hybridization using RNAhybrid Program

S. No.	Target and miRNA	RNAhybrid Result	*mfe (kcal/mol)
1	MD3 & hsa-miR-4471	Hit not found	
2	MD19 & hsa-miR-3675-3p	target 5' A U A G 3' GUUUCU AG GAC UAGAGA UC UUG miRNA 3' G U C AG 5'	-13.7
3	MD29 & hsa-miR-383-5p MD29 & hsa-miR-5197-3p	Hit not found target 5' A GCCA A 3' GUUUGA UCA UAAGCU AGU miRNA 3' ACUG CAGAGAAGAA 5'	-11.6
4	MD134 & hsa-miR-190b-5p	Hit not found	
5	MD 220 & hsa-miR-363-5p	target 5' UAUAA G 3' CGUG CAC GCAC GUG miRNA 3' UUUAACGUA UAG GGC 5'	-12.7
6	MD228 & hsa-miR-4802-3p	target 5' A UUUAC A 3' GG AACC CC UUGG miRNA 3' UA UACCU AAGUU 5'	-11.0
7	MD240 & hsa-miR-23b-5p	target 5' C A G 3' UUGG AUGCU AGUC UACGG miRNA 3' UUU G UCCUU 5'	-12.3
8	MD306 & hsa-miR-325	target 5' G ACACCA G 3' CUGG UCUAG GAUC AGGUC miRNA 3' G AUCCAC AUU 5'	-12.7

S. No.	Target and miRNA	RNAhybrid Result	*mfe (kcal/mol)
	MD306 & hsa-miR-2114-5p	target 5' A G AGA A 3' CC CUUCU AAGUG GG GAAGG UUCGC miRNA 3' A AAC CA 5'	-16.8
9	MD307 & hsa-miR-215-3p	Hit not found	
	MD307 & hsa-miR-548y	Hit not found	
	MD307 & hsa-miR-338-3p	Hit not found	
	MD307 & hsa-miR-3065-5p	Hit not found	
10	MD311 & hsa-miR-4699-3p	Hit not found	
	MD311 & hsa-miR-6739-3p	Hit not found	
11	MR155 & hsa-miR-363-5p	target 5' A C UU C 3' GUG AUC GAU CAC UAG CUA miRNA 3' C UG CGUAAA 5'	-11.8
12	MR165 & hsa-miR-153-5p	Hit not found	
13	MR186 & hsa-miR-744-3p	target 5' AAU UG C 3' UUG UUAG UCAA AAC GAUU AGUU miRNA 3' AC GGU GG 5'	-10.3
	MR186 & hsa-miR-4420	target 5' C U G GU A 3' AC GA GU GU UG CU UA CA miRNA 3' GUCGA U G GU C 5'	-13.1
	MR186 & hsa-miR-448	target 5' C G 3' UGC UGUGUA AUG AUACGU miRNA 3' UGUAGG U 5	-12.0
14	MR243 & hsa-miR-4796-5p	Hit not found	
15	MR274 & hsa-miR-6867-5p	Hit not found	
16	MR304 & hsa-miR-3064-5p	target 5' G GU 3' UGUG CUGCA ACGU GGUGU miRNA 3' GU UGUCGGUCU 5'	-11.7
	MR304 & hsa-miR-411-5p	target 5' G UA C A 3' AC UAUAU GU UG AUAUG CA miRNA 3' A CG C GAUGA 5'	-11.7

3.4. Secondary Structure miRNA Precursor

The pre-miRNA secondary structure was predicated using the online web-server tool RNAfold (<http://rna.tbi.univie.ac.at/cgi-bin/RNAWebSuite/RNAfold.cgi>) (Figures 4). The RNAfold results were

used to predict SARS-CoV-2 hairpin sequences with the most stable secondary structures. Pre-miRNA, around 200bp from the precursor's end, was included in the sequence used for prediction analysis. Folding structures with a centroid were presented in every case.

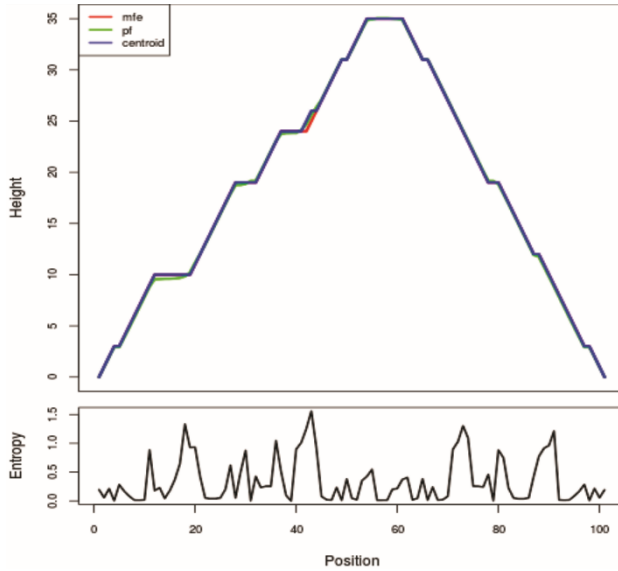


Figure 3. Predicated secondary structure of precursor miRNA hairpin – mounting plot

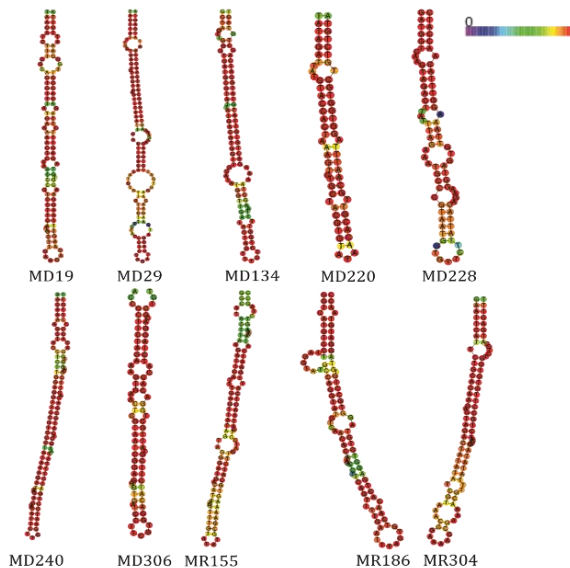


Figure 4. Structure (*Secondary*) of potential hairpin candidates of SARS-CoV-2

4. Discussion

Over the last few decades, miRNA research has been accelerated to explore the pathogenesis function and its role in the development of novel antiviral therapy [41]. MicroRNAs are ~ 21-nt non-coding RNAs derived from large primary

miRNAs (pre-miRNAs) by binding to the 3' UTR of the target miRNAs, which slices gene expression post-transcriptionally and they are well conserved between different organisms [42, 43]. Each miRNA possesses hundreds of target genes and a single gene can be targeted by several miRNAs [44]. There is

increasing evidence suggesting that miRNAs use partial nucleotide sequence complementarity to suppress the expression of protein-coding genes and many biological processes, such as development, proliferation, differentiation (cellular), and pathophysiology are dependent on them [45].

Since the discovery of the first miRNAs, over 2,500 human and a total of 28,645 miRNA sequences have been stored in the miRbase [38, 46-48]. Since miRNAs are essential post-transcriptional regulators of both viral and host gene expression, so they play a significant role in viral pathogenesis. In target selection, the ideal binding position between 3' UTR of the miRNA and the seed region (2 to 7 or 2 to 8 of the 5' ends of the miRNA) is essential, it should be sufficient for effective cleavage [48]. Due to a highly conserved nucleotide position upstream, the minimum pairing requirement is 5 – 6 nucleotide match [49, 50].

The SARS-CoV-2 genome was investigated using different bioinformatics methods, resulting in the identification of 16 potential miRNA precursors. Among those based on bioinformatics analysis were effective hybridization, hybridization pattern and pairing energy. We identified considerable sequence similarity with the SARS-CoV-2 genome where the seed region is concerned and it showed an ideal identity with 3' UTR of viral miRNA. So, we propose that hsa-miR-3675-3p (MD19), hsa-miR-325 (MD306), hsa-miR-2114-5p (MD306), hsa-miR-744-3p (MR186) and hsa-miR-448 (MR186) would be the best potential cellular target miRNAs to develop a post-exposure therapy.

5. Conclusion

In our current investigation, we identified miRNAs for SARS-CoV-2 in human beings using computational tools. This study was based on an interesting

hypothesis of the utilization of host miRNA as a potential post-exposure therapy because the current evidence suggests that host miRNAs may down-regulate the viral gene expression.

Although most of the predicated human miRNAs of SARS-CoV-2 genome functions are yet to be discovered, still we hypothesize that those miRNAs may down-regulate viral gene expression to block its replication [51, 52]. However, further *in vitro* research is needed to determine the effect of chosen miRNAs on viral replication inhibition.

Conflict of Interest

The authors declare no conflict of interest.

Funding

None

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