

The signature of SARS-CoV-2 evolution reflects selective pressures within human guts

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Abstract

In somatic cells, microRNAs (miRNAs) bind to the genomes of RNA viruses and influence their translation and replication. Here we demonstrate that a significant number of miRNA binding sites locate in the NSP4 region of the SARS-CoV-2 genome, and the intestinal human miRNAs exert evolutionary pressure on this region. Notably, in infected cells, NSP4 promotes the formation of double-membrane vesicles, which serve as the scaffolds for replication-transcriptional complexes and protect viral RNA from intracellular destruction. In three years of selection, the loss of many miRNA binding sites, in particular, those within the NSP4, has shaped the SARS-CoV-2 genomes to promote the descendants of the BA.2 variants as the dominant strains that define current momentum of the pandemics. Findings highlight the possibility that intestinal tissue may significantly impact evolution of the SARS-CoV-2 genome and may play a pivotal role in the long COVID.

The signature of SARS-CoV-2 evolution reflects selective pressures within human guts

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Abstract

In somatic cells, microRNAs (miRNAs) bind to the genomes of RNA viruses and influence their translation and replication. Here we demonstrate that a significant number of miRNA binding sites locate in the NSP4 region of the SARS-CoV-2 genome, and the intestinal human miRNAs exert evolutionary pressure on this region. Notably, in infected cells, NSP4 promotes the formation of double-membrane vesicles, which serve as the scaffolds for replication-transcriptional complexes and protect viral RNA from intracellular destruction. In three years of selection, the loss of many miRNA binding sites, in particular, those within the NSP4, has shaped the SARS-CoV-2 genomes to promote the descendants of the BA.2 variants as the dominant strains that define current momentum of the pandemics. Findings highlight the possibility that intestinal tissue may significantly impact evolution of the SARS-CoV-2 genome and may play a pivotal role in the long COVID.

1 Introduction

The (+)RNA genome of the pandemic SARS-CoV-2 virus is about 30,000 nucleotides long [1]. Since December 2019, when the original strain (WuhanHu-1) has emerged, the SARS-CoV-2 acquired more than 10,000 recorded nucleotide mutations [2], with an estimated mutation rate between 0.0004 and 0.002 mutations per nucleotide per year [3]. The current release of the GISAID database contains more than 14 million SARS-CoV-2 sequences [4] and allows the tracking of the evolution of the virus with unprecedented precision. As widely discussed, SARS-CoV-2 is being shaped by the pressure exerted by neutralizing antibodies and applied upon the receptor-binding domain of the Spike (S) protein [5]. It should be, however, noted that antibodies are far from being the only adversarial factor encountered by the virus upon infection. In particular, the binding of host miRNA to other (+)RNA viruses provides a substantial evolutionary pressure [6] as these molecules suppress both the translation and the replication of virus along with altering some aspects of their pathogenesis [7–9].

Several works have predicted interactions of the SARS-CoV-2 single-stranded

(+)RNA virus with multiple human miRNAs [10–13]. A majority of studies have not, however, taken into account the ability of the SARS-CoV-2 virus to evolve, and, accordingly, to selectively alter its miRNA binding regions, especially these enriched in binding seeds. Only a few published works utilized the sequences from various Variants of Concern (VOCs) [14]. Human genomes encode many miRNAs, a majority of which display differential expression in human tissues. It is obvious that only miRNA species expressed in replication-permissive host cells will be relevant to SARS-CoV-2 evolution. In humans, the SARS-CoV-2 virus predominantly replicates in the type 2 alveocytes of the lung. In addition, significant clinical data indicate the presence of the virus in the intestinal epithelium, which expresses both ACE2 and TMPRSS2 [15]. Moreover, intestinal viral persistence is commonly detected. It is now recognized that long-term residence of SARS-CoV-2 in the human gut supports low-intensity inflammation, which contributes to the complications of COVID-19, collectively known as post-COVID or long COVID [16]. For three years of the pandemic, the SARS-CoV-2 virus has significantly mutated. Due to the degeneracy of the genetic code, many mutations have not led to a change in the sequence of viral proteins and, therefore, are classified as silent [17]. Nevertheless, silent nucleotide changes have managed to get fixed in the virus genome. In evolution, the fixation of the mutation may take place either by chance, or under the influence of the selective forces acting upon RNA itself rather than on the encoded proteins. The binding of the cellular miRNAs to the viral genome may be capable of exerting such pressure. The hypothesis that the evolutionary pressure exerted by binding of miRNA species expressed in human lung and intestinal tissue has shaped up the genome of the current variants of SARS-CoV-2 virus dominating the globe was tested.

2 Materials and methods

2.1 SARS-CoV-2 variants

A total of 14,472,180 SARS-CoV-2 RNA sequences were downloaded from the GISAID database [18] on January 7th 2023 and annotated with Pangolin [19] (program’s version 4.2, Pangolin’s data version 1.17). Among 28,420 variants associated with the City of Berlin (Federal Republic of Germany), 3,755 ones satisfying the following conditions were selected: viral RNA does not contain unrecognized nucleotides; the difference between viral RNA length and the “Wuhan” reference variant (GISAID: EPI_ISL_402125) is less than 5%. All selected RNA sequences were collected between the 14th January 2021 and 12th December 2022. All sequences were divided into two groups, “Ancestral variants” and “Omicrons”, according to their GISAID descriptions. “Ancestral variants” included variants of “Alpha”, “Beta”, “Delta”, and others (Table 1 of Supplementary materials) and collected before the 30th January 2022, while all “Omicrons” species were sequenced after 1st December 2021 (Figure 1).

2.2 Highly expressed miRNAs

miRNA-seq read count tables were downloaded from the GDC portal at <https://portal.gdc.cancer.gov/projects/> for colon (COAD) and lung (LUAD) tissues. Following the corresponding clinical description, the dataset samples were referred to as “Normal” (“Solid Tissue Normal”) and “Tumor” (“Primary Tumor”) groups. Only “Normal” samples were used to assess miRNA expression in the analysis: 8 entries in COAD dataset and 46 in the LUAD project. Highly expressed miRNAs were selected independently for each tissue by the following procedure: read counts associated with a particular miRNA by the samples were summed up; then, miRNAs were sorted by the total number of reads; finally, miRNAs accounting for 95% of all reads were selected; As a final expression of the miRNAs, corresponding CPM expressions were averaged by the samples.

2.3 Binding model

To model the binding interaction between miRNA and viral mRNAs, the regions of viral RNA that were a reverse complement to the seed region of miRNA (nucleotides from 2 to 7 on the 5'-end of mature miRNA) were mapped to SARSCoV-2 genomes. Following the common classification [20], such interactions are called “6mer”; they lead to translational inactivation of target mRNAs [21]. In addition to the reverse complementarity of the seed region, miRNA target prediction tools may rely on additional features. For example, TargetScan [22] utilizes context++ score based on site type, 3⁰-supplementary pairing, local AU content and others, MirTarget [23] considers GC content of target site, and UTR length, while RNA22 [24] predicts target sites by placing “Target Islands” within 3'-UTRs, 5'-UTRs or CDS.

Commonly, the overlap of the targets by different software is weak at best [25,

26]. While RNA22 predicts more individual miRNA-mRNA interactions than TargetScan, it misses the majority of classical seed region binding sites in 3'-UTRs. Moreover, the prediction results heavily depend on the tool-specific internal thresholds, with sensitivity and specificity expected to vary depending on particular thresholds. To overcome these limitations, a bottom-up approach based on the mapping of SARS-CoV-2 regions with reverse complement “6mers” matching the seed regions within tissue-specific and relevant host miRNAs was utilized. All SARS-CoV-2 RNA sequences were aligned to the reference “Wuhan” variant using MAFFT tool [27]. After the alignment, we transformed viral RNAs coordinates on the “Wuhan” variant and annotated protein-coding regions using

“Wuhan” annotation (<https://www.ncbi.nlm.nih.gov/nuccore/1798174254>).

2.4 Statistical analysis

In all comparisons, the significances of the findings were evaluated in the nonparametric Mann–Whitney U tests [28]. In addition, the standard Student’s t-test procedure [29] was used to test the hypothesis that the Spearman correlation [30] was zero. The Monte-Carlo sampling procedure was used to show that the decrease in the occurrence of miRNA binding motifs in the “Omicron” sequence group was specific to the highly expressed miRNAs. More precisely, the expression profile of the highly expressed miRNAs was fixed and the miRNA sequences were randomly replaced with other miRNAs expressed in a tissue. After that, the weighted number of binding motifs in “Omicrons” and “Ancestral” groups were calculated: the number of binding motifs within a particular miRNA was multiplied by the proportion of its read counts, summed up and normalized by the total lengths of the compared viral genomes. Then, for “Ancestral variants” and “Omicrons” groups, the computed numbers were compared using the MannWhitney U test. The described procedure was repeated 1,000 times. Thus, the distribution of statistics characterizing the change in the number of binding motifs between “Ancestral variants” and “Omicrons” was obtained. Finally, using this distribution, the probability that the decrease in the number of binding motifs for highly expressed miRNAs was greater than for randomly selected ones was calculated.

2.5 Other technical details

The statistical analysis (hypothesis testing, correlation computation) was performed with SciPy library [31]. All vector computations were implemented using NumPy library [32] and parallelized using Parallel tool [33]. Miscellaneous computations with tables were produced with Pandas [34]. Finally, figures were made using Matplotlib [35] and Seaborn [36] libraries. The source code of the analysis can be found at <https://github.com/zhiyanov/covid-miRNA-evolution>.

3 Results

In this study, both the tissue-specificity and expression volume restrictions were applied to facilitate identification of the most prominent interactions between host miRNAs and the viral genomes. Firstly, the scope of the study was limited to the lungs and colon tissues as the two main sites of SARS-CoV-2 replication in the human body [15]. Secondly, only those miRNA species were analyzed that represent the top 95% of the miRNAome for their tissue type [37], with the following rationale. In COVID-19, each infected cell expresses from 1,500 to 15,000 copies of the virus [38] and from 100,000 to 200,000 miRNA molecules [39]. To visualize the kinetics of their interactions, one should take into account that each miRNA species, in addition to the virus, binds to 25-50 host mRNAs species, each expressed as 50 to 100 copies per cell, thus, and therefore, being supplied with more than 1,250 endogenous target molecules [21]. Because of that, only highly represented miRNA species are likely to exert a significant impact on the evolution of the viral genomes. For lung tissue, there were 32 miRNAs (Figure 2 of Supplementary materials), and for the intestines, 40 miRNAs (Figure 3) that fulfilled the abovementioned criteria. Twenty-one of these miRNAs were found in both of these tissues, and 19 were expressed in either one or another tissue. In lung tissue, lung-specific miRNAs accounted for 81% of all miRNAs, while in the colon only 56% of expressed miRNAs were colon-specific. In lungs, the most prevalent miRNA was miR-143-3p, followed by miR-21-5p, miR-22-3p and miR-30a-5p. In colon, miRNA expression profile had a predominance of let-7b-5p, miR-92a-3p and miR-200c-3p. From the GISAD database, a total of 3,755 SARS-CoV-2 sequences were retrieved. All of them were tagged by location within the same metropolitan area (Berlin, Germany) and a time stamp between the 1st January 2020 and the 20th December 2022 (Figure 1), see “On choosing a city of the study” section of Supplementary materials for more details). All selected sequences were divided into two groups:

“Ancestral variants”, which included “Alpha” (B.1.1.7-like, 759 variants), “Beta” (B.1.351-like, 4 variants), Delta (B.1.617.2-like, 841 variants; AY.4.2like, 12 variants) as well as 7 other strains (Table 1 of Supplementary materials). All of these sequences were collected in January 2022 and earlier; “Ancestral variants” included “Alpha” (B.1.1.7-like, 759 variants), “Beta” (B.1.351-like, 4 variants), Delta (B.1.617.2-like, 841 variants; AY.4.2like, 12 variants) and 7 other strains (Table 1 of Supplementary materials) and belonged to the period until January 2022;

“Omicrons”, which included 8 varieties of “Omicron” strain. All of these were collected in December 2021 or later.

Figure 1: SARS-CoV-2 variants obtained from GISAID portal according to Berlin, Germany geotag.

To assess the evolutionary pressure exerted by colon and lung-specific miR-

NAs, the expression-normalized amounts of binding motifs numbers within 29,903 nucleotides of viral SARS-CoV-2 RNA were analyzed (see “Materials and methods” for more details). In this way, the regulatory contribution of each miRNA was taken separately into account. As compared to “Ancestral variants”, Omicron variants contain strikingly lesser amounts of binding sites for either colon miRNAs (Mann-Whitney U test, p [?] $4.44 \cdot 10^{-254}$, Figure 2, A) or lung miRNAs (Mann-Whitney U test, p [?] $1.07 \cdot 10^{-146}$, Figure 1 of Supplementary materials). By reshuffling the list of miRNAs with a predefined expression repertoire specific to the highly expressed miRNAs, the proportion of random shuffles for which the drop in the number of binding motifs was greater than the drop detected for the highly expressed miRNAs was calculated (the

detailed description of the procedure can be found in “Materials and Methods” section). This proportion turned to be lower than 5% for colon miRNAs (p [?] $2.50 * 10^{-2}$), but not for lung miRNAs (p [?] $7.79 * 10^{-2}$). When the significance of the miRNA influence was tested for other tissue compartments, including the prostate, the breast, the bladder, and the liver, the decrease in the number of observed motifs was not significant for either of these tissues (Table 2 of Supplementary materials). Further study of the colon-specific dropdown showed that it was mainly observed within Omicron brunch (BA.2-like) (see Figure 2, A). However, even after exclusion of Omicron (BA.2-like) from “Omicrons” group, the indicated trend remained strong (Mann-Whitney U test, p [?] $2.57 * 10^{-69}$). Moreover, the weighted number of miRNA binding motifs decreased steadily according to time stamps (Spearman correlation [?] -0.34 , p [?] $7.37 * 10^{-80}$, Figure 2, B), thus, pointing at continuous evolution of “Omicron” sequences within the intestinal niche.

The difference was calculated between the number of binding motifs in the

“Wuhan” and the subsequent “Omicrons” (Figure 3). It turned out that the loss of the binding sites was primarily due to diminished binding to hsa-let-7b-5p, hsa-let7a-5p and hsa-miR-93-5p. Notably, for hsa-miR-194-5p and hsa-miR-29a-3p, the amounts of binding sites in the “Omicrons” have increased. The pairwise alignment of SARS-CoV-2 RNAs with reference “Wuhan” variant allowed us to compare the distributions of the miRNA seed regions within each variant of the virus. Further, the binding distributions for individual miRNA were calculated. It was found that the contribution of each miRNA in those distributions was proportional to its expression in colon tissues. Finally, the distributions were averaged over “Ancestral variants” and different groups of “Omicrons”, either including or excluding Omicron (BA.2-like) variants, and analyzed. As seen from Figure 4, the decrease in amounts of the complementary seeds observed in Omicron BA.2-like variants was primarily due to a loss of hsa-let-7b5p binding motifs located within NSP4. Indeed, C9861T nucleotide mutations occurred mostly in Omicron BA.2-like sequences (872 variants of 887), while the 9861st position in all “Ancestral variants” sequences retained the cytosine. Interestingly, that transition was silent as it has not lead to amino acid mutation: both CTC and CTT triplets encode for lysine.

Notably, a comparison of “Ancestral variants” and other “Omicrons” (Figure

3 of Supplementary materials) showed that the overall dropdown of miRNA binding sites was not due to any specific mutation within a particular miRNA binding motif. On the contrary, a gradual decrease in the number of binding motifs was observed over time (Figure 2, B). To access the contribution of virus RNA coding regions to the observed dropdown, the weighted (by miRNA expression) amounts of miRNA binding motifs per coding region were calculated for “Ancestral variants” and for “Omicrons” with or without Omicron (BA.2-like) sequences separately. By comparing these amounts within “Ancestral variants” with “Omicrons” using Mann-Whitney U test (Table 1), it turned out that Spike and NSP15 regions were changed the most. In addition, the sequences classified as Omicron (BA.2-like) displayed an additional region with accumulated changes, the one encoding for NSP4, while in other “Omicrons” the third most divergent location was the region of NSP12. Figure 2: Weighted number (by miRNA expression in colon tissues) of binding motifs per 100 miRNA and 29,903 nucleotides **A**: grouped by “Ancestral variants” and “Omicrons”, **B**: distributed over time. Spearman correlation between the weighted number and number of days since the start of pandemics is [?] -0.34 , (p [?] $7.37 * 10^{-80}$). In this analysis, Omicron (BA.2 like) variants were excluded. Figure 3: Amounts of the binding motifs in analyzed SARS-CoV-2 variants and reference “Wuhan” variant differ. Here, miRNAs are sorted by the levels of their expression in the intestine. The share of miRNome represented by particular miRNA could be found adjacent to miRNA ID. Figure 4: Distributions of miRNA binding motifs along to “Wuhan” RNA sequences and averaged by **A**: “Ancestral variants”, **B**: difference between “Ancestral variants” and Omicron (BA.2-like) variants, and **C**: Omicron (BA.2like) variants.

4 Discussion

In this study, the hypothesis that host cell miRNAs may have influenced the evolution of SARS-CoV-2 was tested. Two sets of miRNAs expressed in the main sites of virus reproduction, the lungs and the intestine, were considered. It was found that those sets significantly differed in the representation of individual miRNAs. On one hand, hsa-miR-143-3p, hsa-miR-21-5p, hsa-miR-22-3p and hsa-miR-30a-5p were specific for the lungs, with large predominance of miR-1433p expression (41.5%). On the other hand, the intestine was characterized by a more uniform profile with the presence of tissue-specific let-7b-5p, miR-92a-3p and miR-200c-3p. During pandemics, SARS-CoV-2 genomes lost many miRNA binding sites specific for the lung and intestinal tissue expressed miRNAs. Importantly, MonteCarlo simulations showed that observed changes were significant only for the set

Region	Length	Ancestral variants	Omicron BA.2-like	Omicron BA.2-like	Other Omicrons	Other Omicrons
			Diff	FDR	Diff	FDR
NSP4	1500	962	464	< 0.0001	1	0.1127
Spike	3822	709	21	< 0.0001	20	< 0.0001
NSP15	1038	376	21	< 0.0001	11	< 0.0001
NSP9	339	1518	6	< 0.0001	7	< 0.0001
NSP3	5835	467	4	< 0.0001	0	0.7554
NSP6	870	1162	3	< 0.0001	-3	< 0.0001
NSP12	2769	356	1	0.1789	2	< 0.0001
NSP14	1581	1781	1	0.8104	2	0.0129
N	1260	683	1	0.4540	-1	0.0123
NS7a	366	223	0	0.5782	0	0.0458
NSP13	1803	1659	0	< 0.0001	0	< 0.0001
NS9b	294	444	0	0.9704	-4	0.0040
E	228	2338	-2	0.1030	-2	0.0486
NSP2	1914	455	-5	0.3511	3	0.0015
NS9c	222	2474	-5	0.1000	-7	0.0166
NS3	828	243	-5	< 0.0001	0	0.0002
NSP1	540	498	-7	< 0.0001	-4	0.0001
3P	370	188	-9	< 0.0001	0	0.9165
RNA	29903	744	27	< 0.0001	3	< 0.0001

Table 1: The differences in the weighted number of the binding motifs for colon-specific miRNA within various coding regions of SARS-CoV-2 RNA genome. All weighted numbers are averaged by variants of the groups and normalized by “Wuhan” RNA length. “Diff” columns contain differences between the averaged values of the corresponding groups. “FDR” columns contain adjusted p-values of Mann-Whitney U tests comparing the weighted numbers of virus variants of the corresponding groups. of miRNAs expressed in the intestinal tissue (p [?] $2.50 \cdot 10^{-2}$). Coronaviruses are generally perceived as Acute Respiratory Infections (ARIs), and the viral production in the lungs provides for a reservoir underlining the virus spread. In addition to the lungs, SARS-CoV-2 is capable of replication in many other organs and tissues [40], the main of which is an intestine, where the virus may reside for up to three month [41]. Prolonged period of intestinal replication provides sufficient time for evolutionary forces to work on the population of the viruses within the same host, with the host factors serving as the drivers for such evolution. Arguably, binding of host miRNAs represents such a force. Resultant viral populations may be returned to circulating pool of the viruses either through the fecal-oral route [42] or during secondary viremia in the immunocompromised hosts capable of supporting internal reinfection of the lungs [43,44], and subsequent airborne spread. It has been shown that the initial version of Omicron variant shared a high similarity with the variants of the

SARS-CoV-2 virus found in rodents, particularly in mice [45]. Therefore, it has been hypothesized that one of the ancestral variants of SARS-CoV-2, from approximately mid-2020 [46], has entered the population of rodents and has evolved within these animals for quite some time till reemerging in humans [47,48]. An alternative theory suggests a longterm virus evolution within an immunocompromised patient, predominantly in the intestines [49]. An observation described in this study highlights the significance of the intestinal site for the pathogenesis of COVID-19. Although pneumonia and respiratory complications account for the most of the morbidity and mortality of COVID-19, extrapulmonary manifestations of COVID-19 infection include prominently diarrhea, as well as nausea, emesis, anorexia, abdominal pain, and heartburn [41]. These complications are observed in 10%-20% of patients with COVID-19 [50], which does not exclude the possibility of asymptomatic gastrointestinal infections with SARS-CoV-2. Moreover, the latent persistence of SARS-CoV-2 in the intestine may be one of the contributors to the long COVID [51]. A dramatic decrease in miRNA binding sites was especially pronounced in BA.2-like SARS-CoV-2 variants. To a large extent, this drop was due to the mutations within the region encoding the NSP4 protein, which promotes the formation of double-membrane vesicles serving as a scaffold for replication/transcriptional complexes [52] of the virus. Additionally, attachment to NSP4 protects viral RNA from intracellular immunity [53]. Along with dramatic changes in S-protein, the miRNA-driven alterations of NSP4, acquired during prolonged replication with the intestine, may have contributed to the current global dominance of the descendants of the BA.2 variant [54]. Findings highlight the possibility that intestinal tissue may significantly impact evolution of the SARS-CoV-2 genome and may play a pivotal role in the long COVID.

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6 Conflict of interest

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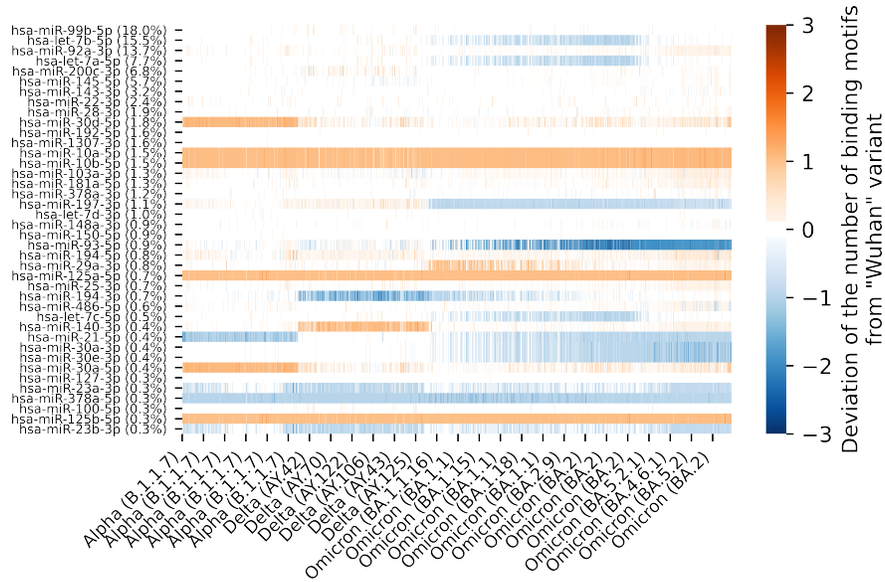
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