

# Quality control of low-frequency variants in SARS-CoV-2 genomes

Mikhail Rayko<sup>1\*</sup>, Aleksey Komissarov<sup>2</sup>

<sup>1</sup>Center for Algorithmic Biotechnology, Institute of Translational Biomedicine, St. Petersburg State University, Saint Petersburg, Russia;

<sup>2</sup>Applied Genomics Laboratory, SCAMT Institute, ITMO University, Saint Petersburg, Russia;

\* To whom correspondence may be addressed - [m.rayko@spbu.ru](mailto:m.rayko@spbu.ru)

## Abstract

During the current outbreak of COVID-19, research labs around the globe submit sequences of the local SARS-CoV-2 genomes to the GISAID database to provide a comprehensive analysis of the variability and spread of the virus during the outbreak. We explored the variations in the submitted genomes and found a significant number of variants that can be seen only in one submission (singletons). While it is not completely clear whether these variants are erroneous or not, these variants show lower transition/transversion ratio. These singleton variants may influence the estimations of the viral mutation rate and tree topology. We suggest that genomes with multiple singletons even marked as high-covered should be considered with caution. We also provide a simple script for checking variant frequency against the database before submission.

## Introduction

Sequencing of viral genomes allowed researchers to track the distribution of the viruses on Earth, and to assess the rate of the viral evolution. This task is especially important during the active outbreaks, where arisen mutations may affect test systems and vaccines under development.

During the current pandemic of SARS-nCoV-2, the primary resource for consolidating genomic data is the GISAID database (Shu et al., 2017). As a result of the collaborative efforts of the researchers worldwide, on April 14, 2020 it contained over 8,000 SARS-nCoV-2 genomes from different countries, sequenced and assembled using various technologies and approaches.

Unfortunately, these sequences are not error-free. Different sequencing technologies are characterized by different types and frequency of errors (Ma et al., 2019). Often these sequencing errors are not random and are typical for certain sets of nucleotides such as homopolymers. At least 21% of submitted genomes on April 1, 2020 are sequenced using Oxford Nanopore technology according to GISAID (according to searching by “nanopore” or “minion” keywords in metadata). Oxford Nanopore technology is error-prone and ONT data requires careful polishing.

Another source of systematic errors can be the use of a fixed set of primers, which leads to the enrichment of some regions over others. While many assemblers imply more or

less uniform coverage, primer sets (such as described at [https://github.com/CDCgov/SARS-CoV-2\\_Sequencing](https://github.com/CDCgov/SARS-CoV-2_Sequencing)) are commonly used to enrich the sequences. The use of PCR enrichment may result in low coverage of individual genomic regions (even in case of high average coverage), which can be a serious source of errors in the downstream analysis.

In addition to sequencing errors, the variety of genome assembly methods makes it difficult to compare data, and the lack of access to raw data makes it impossible to reassemble data using a standardized approach. Prompt access to original raw sequencing data is needed to perform accurate and reproducible analysis.

GISAID database curators do a tremendous job of filtering submitted sequences, but sometimes it is difficult to distinguish real variants from errors, especially at the lack of information about coverage. Here we compared variants across the submissions and developed a pipeline to separate real variants from potential errors based on their frequency across all genomes in the database. We suppose that variations observed in a single genome from the dataset - hereinafter referred to as *singletons* - may be erroneous, and one should proceed with caution, or maybe even filter out singleton-containing genomes from downstream applications until we get additional evidence from other samples.

## Methods

### Dataset

8,053 full-length (>29,000 bp) sequences of the SARS-CoV-2 were downloaded from the GISAID database ([www.epicov.org](http://www.epicov.org)) on April 14, 2020, including 5,556 genomes marked as “high coverage”.

### Variation calling

Sequences were aligned to the reference genome (NCBI RefSeq NC\_045512.2) using minimap2 (Li, 2018). Resulting vcf files were merged using MergeVcfs tool from Picard toolkit (<http://broadinstitute.github.io/picard/>). SNVs were annotated using SnpEff 4.3t (Cingolani et al., 2012). Ts/Tv was calculated as a direct transition/transversion ratio on a filtered set of SNVs (not considering their multiplicity, and excluding indels and Ns). Scripts for data analysis and visualization are available at [https://github.com/ablab/covid19\\_variation\\_analysis](https://github.com/ablab/covid19_variation_analysis).

### Sequencing and assembly technology statistics

We used the following keywords for GISAID database to get information about sequencing methods: “Illumina”, “Nanopore”, “Ion Torrent”, “Sanger”, “dbnseq”. We used the following keywords for GISAID database to get information about assembly methods: “artic”, “phe”, “spades”, “dnbseq”, “megahit”, “clc”, “ivar”, and “seattle”. To get various assembly methods based on raw reads mapping we used the following keywords: “mpileup”, “bwa”, “bowtie”, or “mapping”.

## Results

### **Submitted sequences contain a large fraction of singleton variations**

After filtering out all variants containing Ns, there are variants in 4,562 positions (out of 29,903 bp). 3,006 of them were identified as singletons. Figure 1 illustrates quantity and distribution over the genome for SNVs of different multiplicity.

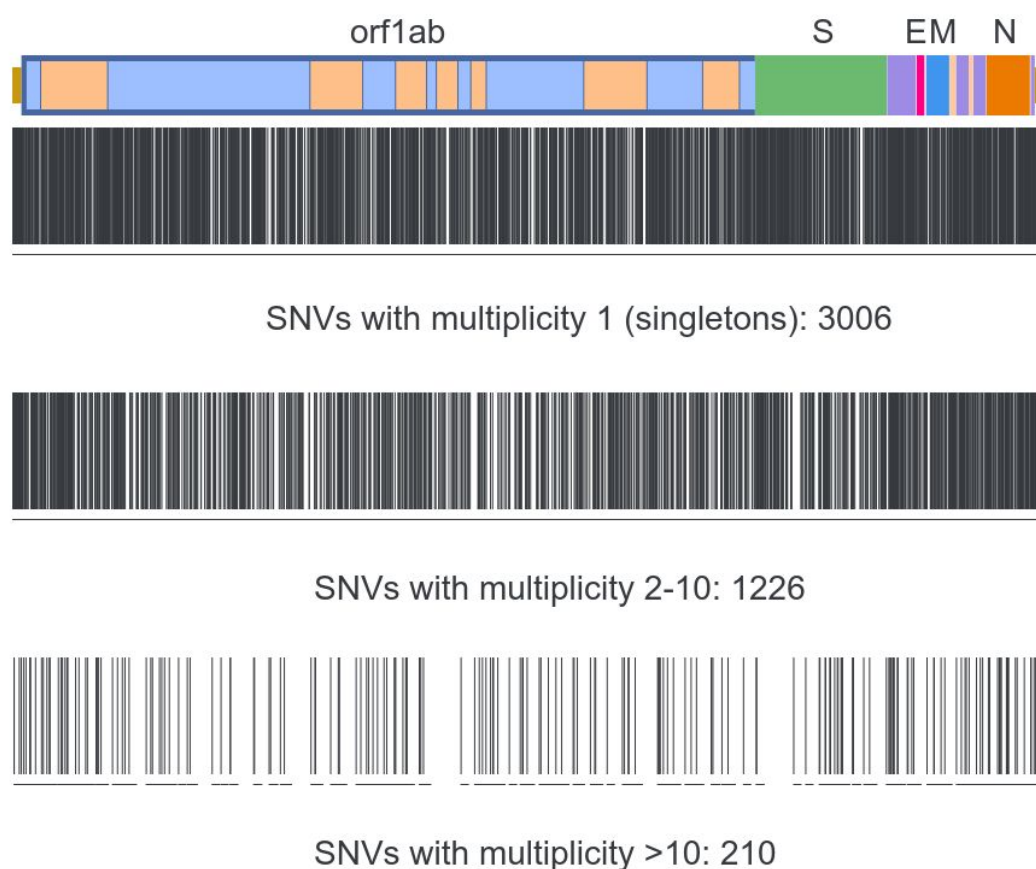


Figure 1. Visualisation of the obtained SNVs in SARS-CoV-2 genomes collected before April 14, 2020. Top to bottom: singletons, SNVs observed in 2-10 genomes, SNVs observed more than in 10 genomes.

### **Singleton variations show decreased $Ts/Tv$ ratio**

We explored transition/transversion ( $Ts/Tv$ ) ratio for the variants observed with different frequencies. For singletons this ratio is lower than for more frequent variants. (Table 1). Lower  $Ts/Tv$  ratio corresponds to false positive results (e.g. Wang et al. 2015, Guo et al 2012), and may indicate the introduced sequencing/assembly errors.

Variant Frequency	All genomes			"High coverage"		
	Ts	Tv	Ts/Tv	Ts	Tv	Ts/Tv
1 (singleton)	1,834	1,132	1.62	1,355	697	1.94
2	492	214	2.3	364	141	2.58
3	214	97	2.21	175	75	2.33
4	125	47	2.66	92	32	2.88
5	80	25	3.2	44	15	2.93
>=6	311	114	2.73	247	92	2.68

Table 1. Ts/Tv for variants that occur in SARS-nCoV-2 genomes with different frequencies.

Currently, genomes with more than 0.05% singleton mutations (i.e. more than 15 SNPs) are automatically excluded from "high-covered" in GISAID database. When comparing the fraction of genomes marked as "high-covered" among genomes that contain singletons we see that for genomes with 2 and less singletons there is no significant difference. However, the fraction of genomes with 3 or more singletons is significantly ( $p < 0.01$ , counted with  $\chi^2$  criteria) lower than in those that do not contain any singletons (see Table 2).

X (# of singletons per genome)	0	1	2	3	>3
Total genomes with X singletons	6,194	1,307	363	98	91
HC genomes (fraction) among genomes with X singletons	4,282 (0.69)	932 (0.71)	252 (0.69)	54 (0.55)	36 (0.4)

Table 2. Number of genomes with different amounts of singletons, and their fraction in genomes marked as "high-covered" in GISAID database (denoted as "HC").

### **Frameshift indels and nonsense mutations are usually correspond to singletons**

All variants in coding regions were annotated with SnpEff. Singletons showed slightly higher presence of the frameshift indels and stop-gained mutations, most probably erroneous (see Table 3). Also we see a significant difference in the percentage of synonymous variants between singleton and non-singleton SNPs. However, it is not clear whether this difference corresponds to errors or not.

	Singletons	Non-singletons	Singletons from HC genomes	Non-singletons from HC genomes

In-frame indels	42 (1.44% )	15 (0.9%)	2 (0.1%)	5 (0.3%)
Frameshift indels	101 (3.5%)	5 (0.3%)	9 (0.5%)	3 (0.2%)
Missense variants	1,838 (62.8%)	970 (59.5%)	1,064 (63.7%)	859 (58.9%)
Synonymous variants	891 (30.5%)	630 (38.6%)	575 (34.4%)	582 (39.9%)
Stop gained mutations	52 (1.8%)	11 (0.7%)	18 (1%)	9 (0.62%)
Stop lost mutations	1 (0.1%)	0	1 (0.1%)	0
Total mutations in coding regions	2,925	1,631	1,669	1,458

Table 2. Types of mutations in the coding regions.

### ***Indels should be verified prior to submission.***

Genomes marked as “high covered” in the GISAID database must not contain indels unless verified by the submitter. Thus, this is important to compare obtained indels with those already presented in the database. Out of 227 indels from samples collected and submitted to GISAID before April 14st, 2020, we observed only 33 non-singletons (see Supplemental Table 1). 30 of these non-singleton indels are observed in at least one HC genome, some of them were already described and checked (Bal et al., 2020, Su et al., 2020).

### ***Genome assembly method seems more important than sequencing technology***

We extracted information about sequencing technology by keywords in metadata, and estimated the number of genomes with singletons. We were expecting to see an elevated number of singleton-containing genomes in the Oxford Nanopore results. However, it turned out that the proportion of the singleton-containing genomes for Illumina and ONT data is almost the same (see Table 3). There is a small amount of Sanger and DNBseq data in the database at the moment, these results may change over time.

	# of genomes	% of genomes with >0 singletons	% of genomes with >1 singletons
Illumina	5602	18.58	5.23
Nanopore	1609	19.14	4.79
Ion Torrent	94	24.47	10.64
Sanger	42	26.19	11.9
DNBseq	16	25.00	6.25
Other/Not specified	690	29.06	8.82

Table 3. Percentage of singleton-containing genomes depending on sequencing technology.

Then we compared different assembly methods (see Table 4). We found the lowest number of singletons in genomes assembled by specialized virus-tailored pipelines, such as Artic Network (<https://artic.network/ncov-2019>), Phe (Public Health England), iVar (Grubaugh et

al., 2019) and Seattle flu assembly pipeline (<https://github.com/seattleflu/assembly>). De novo assembly with MEGAHIT (Li et al., 2015) shows a significant amount of singletons - one should probably interpret such results with caution. The full data shown in Supplementary table 2.

	Technology	# of genomes	# of genomes with singletons	# of genomes with >1 singletons
Phe pipeline	Illumina	1749	13.15	3.03
Artic pipeline	Illumina, ONT	825	18.42	4.97
CLC	Illumina, ONT, Ion Torrent	485	31.55	11.96
iVAR	Illumina	287	18.12	2.79
seattle	Illumina	122	11.48	2.46
SPAdes	Illumina, Ion Torrent	115	20.00	5.22
MEGAHIT	Illumina	110	47.27	22.73
Other*	-	4360	20.53	5.66

Table 4. Percentage of singleton-containing genomes depending on assembly method.

\*"Other" category includes all custom pipelines, rarely used tools and samples with incomplete or absent information about assembly methods.

### ***There is a two week delay between sample collection and genome availability***

For each day from January 1 to April 15 we computed a number of total known variants and a number of variants shared in more than one genome to this date (Figure 1). We found that the data on the new variants has a two week delay. This time delay should also be taken into account when analyzing the data, especially if one links it to other more rapidly updated data, such as infection statistics.

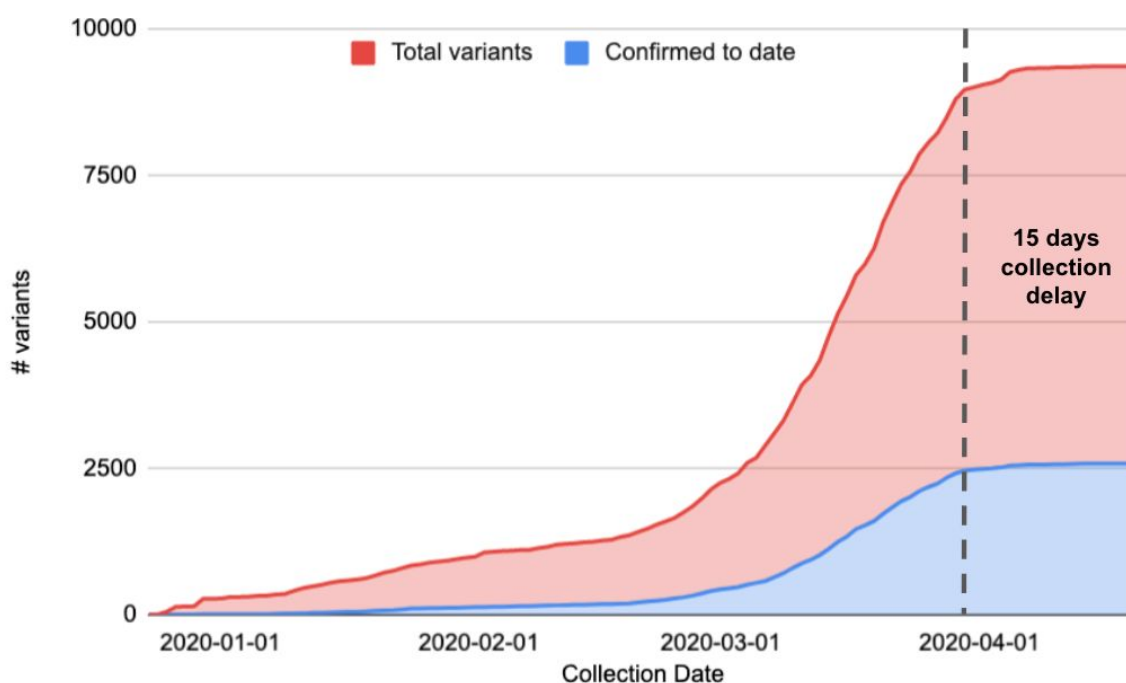


Figure 2. Fraction of confirmed variants during the four months period.

## Discussion

Based on our results, we suggest that the singletons can serve as indicators of the potential erroneous assembly. We provide a script that quickly allows to obtain sample's SNVs frequencies against the current SARS-CoV-2 samples database and to use this result as a sanity check before submission. It is definitely possible to see in a recently sequenced sample a real SNV that is not present in the database yet. However, if you see plenty of them we recommend checking these locations (i.e. in some genome visualization software like Tablet (Milne et al., 2012)).

Described artifacts may influence even the estimations of the viral mutation rate and phylogenetic tree topology. We suggest that genomes with multiple singletons even marked as high-covered should be used with caution.

Although described methods allow to notice some potential errors in the database, reliable quality control for individual samples is not possible without access to reads. We hope to extend this work when more raw sequencing data connected to GISAID genomes will become available in public databases.

## Data availability

Scripts for reproducing the analysis steps and checking variant frequency against the database before submission are available at [https://github.com/ablab/covid19\\_variation\\_analysis](https://github.com/ablab/covid19_variation_analysis).

## Acknowledgements.

Our results acknowledge, as the original source of the data, the laboratories where the clinical specimens and/or virus isolates were obtained (see full list in the [Supplementary table 3](#)). Aleksey Komissarov was financially supported by the Government of the Russian Federation through the ITMO Fellowship and Professorship Program. Mikhail Rayko was supported by St. Petersburg State University (ID 51555639).

We are grateful to Dmitry Antipov, Sonya Garushyants, Dmitry Meleshko, Anton Korobeynikov and Alla Lapidus for suggestions and comments that improved the paper.

## References

- Bal, A., Destras, G., Gaymard, A., Bouscambert-Duchamp, M., Valette, M., Escuret, V., Frobert, E., Billaud, G., Trouillet-Assant, S., Cheynet, V. and Brengel-Pesce, K., 2020. Molecular characterization of SARS-CoV-2 in the first COVID-19 cluster in France reveals an amino-acid deletion in nsp2 (Asp268Del). *bioRxiv*.
- Cingolani, P., Platts, A., Wang, L.L., Coon, M., Nguyen, T., Wang, L., Land, S.J., Lu, X. and Ruden, D.M., 2012. A program for annotating and predicting the effects of single nucleotide polymorphisms, SnpEff: SNPs in the genome of *Drosophila melanogaster* strain w1118; iso-2; iso-3. *Fly*, 6(2), pp.80-92.
- Grubaugh, N.D., Gangavarapu, K., Quick, J., Matteson, N.L., De Jesus, J.G., Main, B.J., Tan, A.L., Paul, L.M., Brackney, D.E., Grewal, S. and Gurfield, N., 2019. An amplicon-based sequencing framework for accurately measuring intrahost virus diversity using PrimalSeq and iVar. *Genome biology*, 20(1), pp.1-19.
- Guo, Y., Li, J., Li, C.I., Long, J., Samuels, D.C. and Shyr, Y., 2012. The effect of strand bias in Illumina short-read sequencing data. *BMC genomics*, 13(1), p.666.
- Ma, X., Shao, Y., Tian, L., Flasch, D.A., Mulder, H.L., Edmonson, M.N., Liu, Y., Chen, X., Newman, S., Nakitandwe, J. and Li, Y., 2019. Analysis of error profiles in deep next-generation sequencing data. *Genome biology*, 20(1), p.50.
- Li, D., Liu, C.M., Luo, R., Sadakane, K. and Lam, T.W., 2015. MEGAHIT: an ultra-fast single-node solution for large and complex metagenomics assembly via succinct de Bruijn graph. *Bioinformatics*, 31(10), pp.1674-1676.



Li, H., 2018. Minimap2: pairwise alignment for nucleotide sequences. *Bioinformatics*, 34(18), pp.3094-3100.

Milne I, Stephen G, Bayer M, Cock PJA, Pritchard L, Cardle L, Shaw PD and Marshall D. 2013. Using Tablet for visual exploration of second-generation sequencing data. *Briefings in Bioinformatics* 14(2), 193-202.

Redelings, B.D. and Suchard, M.A., 2007. Incorporating indel information into phylogeny estimation for rapidly emerging pathogens. *BMC evolutionary biology*, 7(1), p.40.

Shu, Y., & McCauley, J. (2017). GISAID: Global initiative on sharing all influenza data—from vision to reality. *Eurosurveillance*, 22(13).

Su, Y., Anderson, D., Young, B., Zhu, F., Linster, M., Kalimuddin, S., Low, J., Yan, Z., Jayakumar, J., Sun, L. and Yan, G., 2020. Discovery of a 382-nt deletion during the early evolution of SARS-CoV-2. *bioRxiv*.

Wang, J., Raskin, L., Samuels, D.C., Shyr, Y. and Guo, Y., 2015. Genome measures used for quality control are dependent on gene function and ancestry. *Bioinformatics*, 31(3), pp.318-323.