

# Topological Surveillance of Recurrent Mutations in SARS-CoV-2

CoVtRec report as of 17 May 2022

Michael Bleher<sup>2\*</sup>, Lukas Hahn<sup>2\*</sup>, Maximilian Neumann<sup>1\*</sup>, Samuel Braun<sup>3</sup>, Holger Obermaier<sup>3</sup>, Mehmet Soysal<sup>3</sup>, René Caspart<sup>3</sup>, Andreas Ott<sup>1,2\*</sup>

\*Correspondence: mbleher@mathi.uni-heidelberg.de (M.B.) Ihahn@mathi.uni-heidelberg.de (L.H.) maximilian.neumann@kit.edu (M.N.) andreas.ott@kit.edu (A.O.)

#### **Abstract**

The appearance of new variants of the coronavirus SARS-CoV-2 in the current COVID-19 pandemic underlines the importance of being able to quickly identify mutations that could confer some adaptive advantage to the virus, such as immune evasion or higher infectivity. Here we apply CoVtRec, a fast and scalable early warning system based on Topological Data Analysis, for the identification and surveillance of emerging potentially adaptive mutations in the ongoing evolution of SARS-CoV-2. CoVtRec is based on a new topological approach to the surveillance of recurrent mutations in large genomic datasets developed in [1].

<sup>&</sup>lt;sup>1</sup>Mathematics Department, Karlsruhe Institute of Technology, Karlsruhe, Germany

<sup>&</sup>lt;sup>2</sup>Mathematical Institute, Heidelberg University, Heidelberg, Germany

<sup>&</sup>lt;sup>3</sup>Steinbuch Centre for Computing, Karlsruhe Institute of Technology, Karlsruhe, Germany



### Results

We analyzed topological signals for the ongoing convergent evolution of the coronavirus SARS-CoV-2 on the Spike gene from 17 April 2021 until 17 May 2022. To that end, we performed a topological recurrence analysis for a curated alignment of 9,894,062 high-quality SARS-CoV-2 Spike gene sequences shared via GISAID, the global data science initiative [2, 3]. For each Spike mutation we computed its topological recurrence index (tRI) and the corresponding time series analysis chart. The topological recurrence index is a topological measure for convergence of a given mutation (see [1, 4] for details).

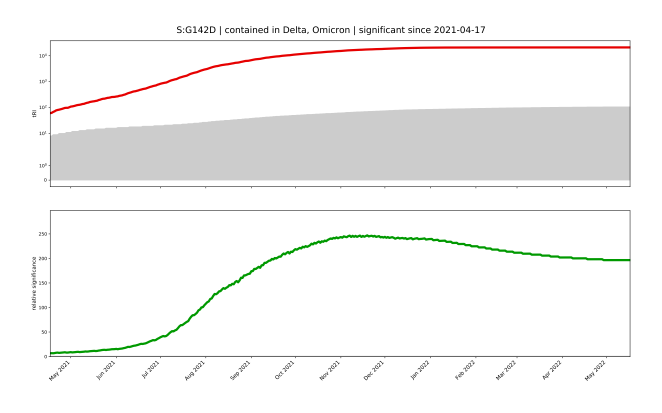
We present a list featuring the top ten amino acid variations on the Spike gene that show strongest topological signal of convergence as of 17 May 2022 (see Table 1). Here signals with  $tRI \ge 106$  are statistically significant (p < 0.05). It was demonstrated in [1] that these mutations are potentially adaptive in the current phase of the pandemic and might therefore appear in future variants. We also present time series analysis charts (see Figure 1) showing (i) the development of the topological signal as well as its significance over time, (ii) major lineages containing the mutation, and (iii) the date from which on the topological signal became significant.

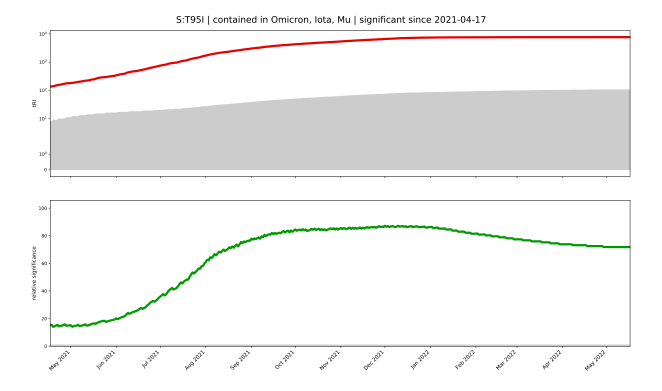
SAAV	tRI	relative significance	notable variants
G142D	21037	198.5	Delta, Omicron
T95I	7684	72.5	Omicron, Iota, Mu
L5F	5916	55.8	lota
D950N	4568	43.1	Delta, Mu
R346K	3240	30.6	Mu
A222V	3016	28.5	
V1264L	2372	22.4	
N440K	2304	21.7	Omicron
A701V	1986	18.7	Beta, Iota
S112L	1920	18.1	

**Table 1.** The top ten amino acid changes on the Spike gene showing strongest topological signal of convergence as of 17 May 2022. For a given mutation, the table displays its topological recurrence index (tRI), its relative significance, and notable variants containing the mutation.

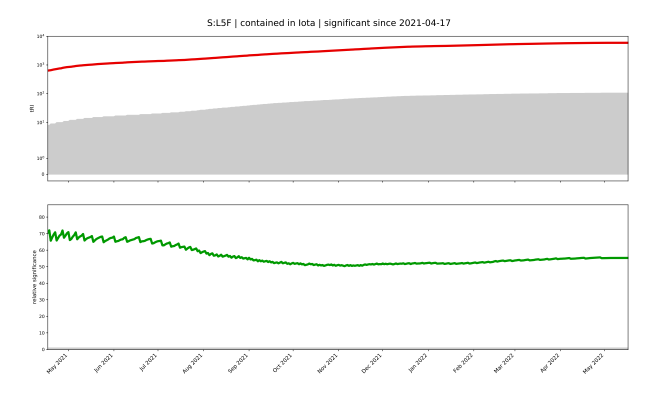


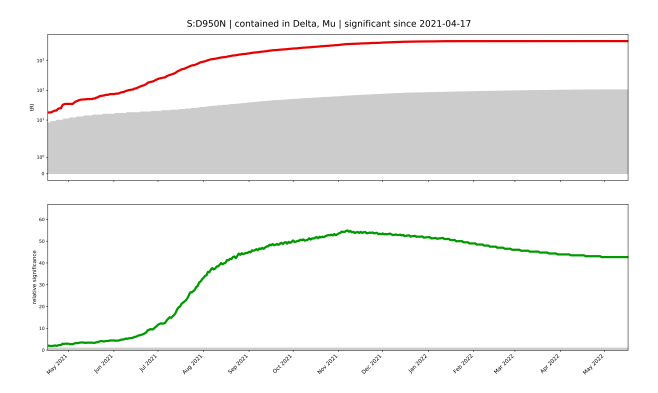
**Figure 1.** Time series analysis charts for the mutations listed in Table 1. Each chart shows the topological recurrence index (red) and its relative significance (green) from 17 April 2021 until 17 May 2022. In each chart, in the upper diagram the shaded region marks the level of significance.



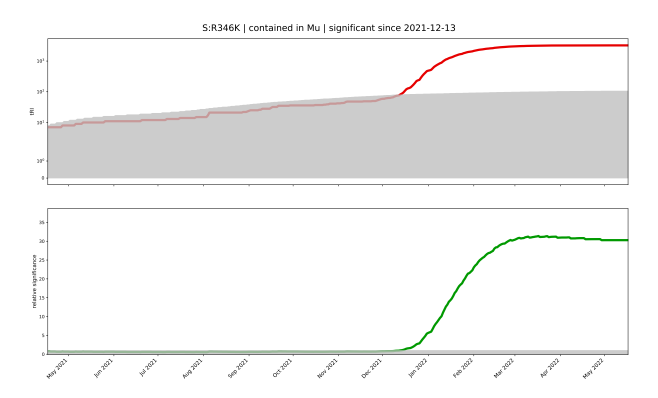


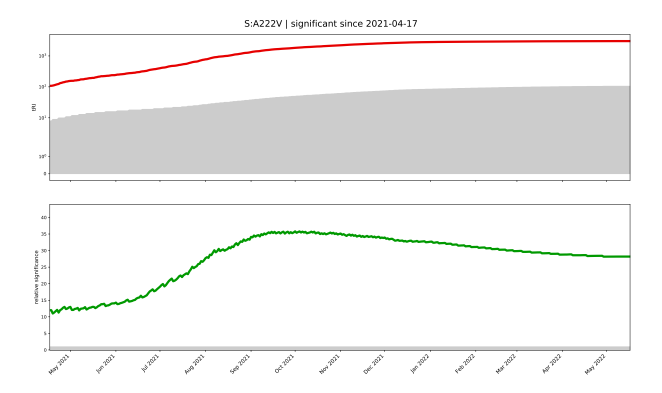




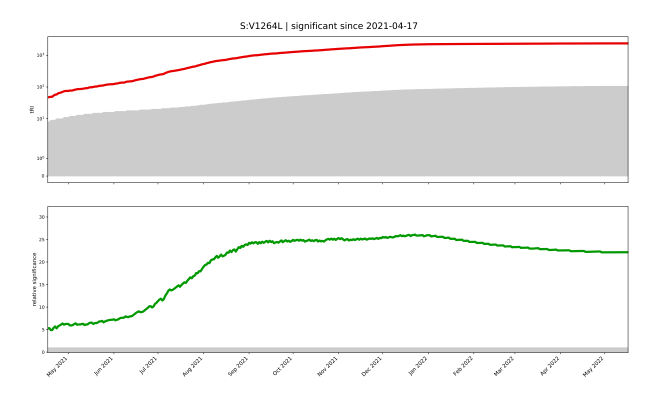


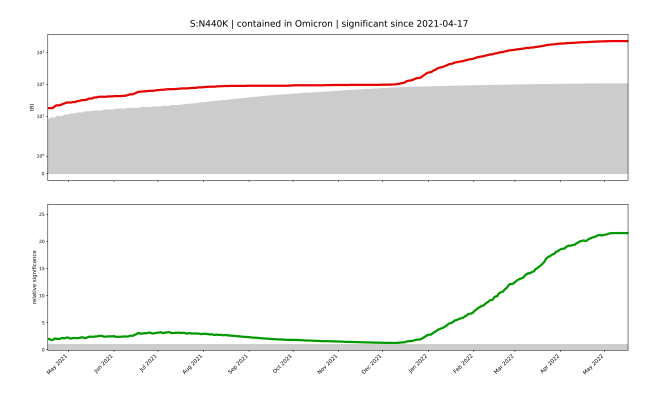




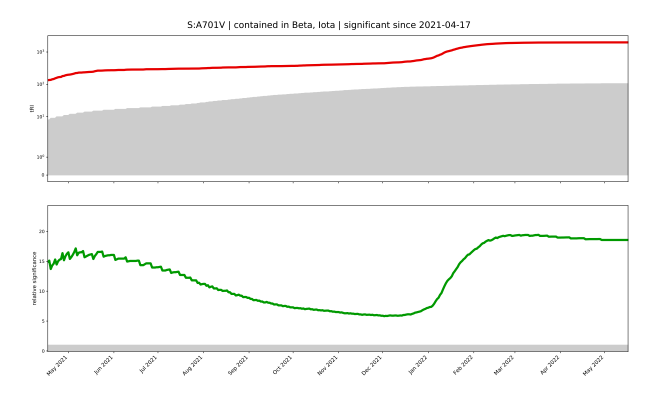


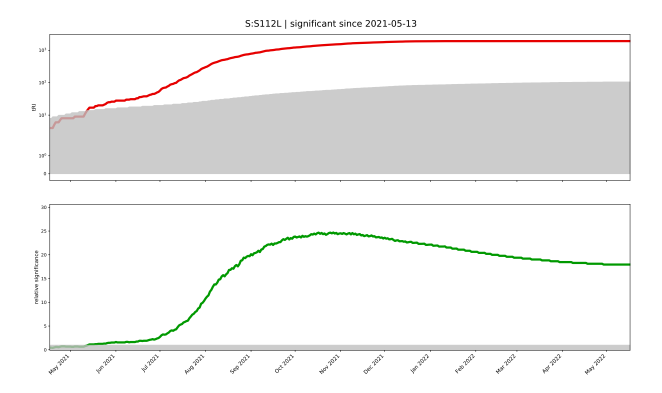














#### Methods

#### Data acquisition and data preparation

Our analysis is based on the alignment msa\_0517.fasta downloaded from the GISAID EpiCoV Database [2, 3] on 19 May 2022. This alignment comprises 9,894,062 SARS-CoV-2 whole genome sequences that have been aligned to the reference sequence Wuhan/WIV04 with GISAID accession number EPI\_ISL\_402124 using MAFFT (Version 7.497) [5]. Sequences in this alignment were truncated to the Spike gene (reference site positions 21,563 to 25,384), and subsequently sequences containing any characters other than A, C, T, G or - were removed. This resulted in an alignment comprising 5,322,559 complete SARS-CoV-2 Spike genes of length 5,850nt. A list of accession numbers of all sequences in this alignment, along with an acknowledgement of the contributions of both the submitting and the originating laboratories, can be retrieved through the Data Acknowledgement Locator at https://www.gisaid.org with ID EPI\_SET\_20220520dz.

### Topological recurrence analysis

The Spike gene alignment contains 335,598 genetically distinct sequences. We used Hammingdist (Version 0.15.0) [6] to compute the genetic distance matrix of this alignment. Subsequently we used Ripser [7] to compute the representative cycles for the persistent homology of the Vietoris-Rips filtration associated to the genetic distance matrix. The computation of persistence barcodes was restricted to small genetic distance scales (Ripser scale parameter threshold set to 2). Next a complete list of SNV cycles (topological cycles all of whose edges correspond to single nucleotide variations) in the given alignment was generated from the corresponding Ripser output. Then we used custom code implemented in Python to compute the topological recurrence index (tRI) for each such SNV. Summing over all SNVs determining an SAAV (single amino acid variation), we computed the tRI for each SAAV. Lastly, from the distribution of the tRI measurements over the whole Spike gene we inferred the level of significance for the tRI per SAAV. For each SAAV, its relative significance is then defined as the quotient of its tRI by the level of significance. Using dimensional reduction in multipersistence via deformations of distance matrices, we computed tRI time series analysis charts at daily resolution from the natural stratification by time of genomic data. For a more detailed description of the topological recurrence analysis see [1, 4].

#### Data availability

All SARS-CoV-2 genome data used in this work are available from the GISAID EpiCov Database [2, 3] at https://www.gisaid.org and can be retrieved through the Data Acknowledgement Locator with ID EPI\_SET\_20220520dz.

#### Code availability

Code used for the analyses is available at https://github.com/ssciwr/hammingdist and https://github.com/Ripser/ripser/tree/tight-representative-cycles. All other code is available from the corresponding authors upon request.

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the genetic sequence and metadata and sharing via the GISAID Initiative [2, 3], on which this research is based. Acknowledgement tables can be retrieved through the Data Acknowledgement Locator at https://www.gisaid.org with ID EPI\_SET\_20220520dz. The authors acknowledge the use of de.NBI Cloud and the support by the High Performance and Cloud Computing Group at the Zentrum für Datenverarbeitung of the University of Tübingen and the German Federal Ministry of Education and Research (BMBF) through grant no 031 A535A. They thank M. Hanussek for IT support and early access to VALET [8]. The authors further acknowledge support from the Interdisciplinary Center for Scientific Computing at Heidelberg University and the development work of the Scientific Software Center of Heidelberg University carried out by L. Keegan and D. Kempf [6]. This research was funded by the Federal Ministry of Education and Research (BMBF) and the Baden-Württemberg Ministry of Science as part of the Excellence Strategy of the German Federal and State Governments (KIT Centers, "Topological Genomics"), and by the Vector Foundation ("Topological Genomics"). M.B. was supported by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) under Germany's Excellence Strategy EXC 2181/1 - 390900948 (the Heidelberg STRUCTURES Excellence Cluster). L.H. thanks the Evangelisches Studienwerk Villigst for their support.

#### **Author contributions**

M.B., L.H., M.N., A.O. designed the study; M.B., L.H., A.O. curated data; M.B., L.H., M.N., A.O. performed computational analyses; M.B., L.H., M.N., A.O., S.B., H.O., M.S., R.C. developed and implemented software; M.B., L.H., A.O. acquired computing resources; M.B., L.H., A.O., M.N. drafted the manuscript; all authors contributed to the final version of the report.

# **Competing interests**

The authors declare no competing interests.

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