

1 ***coronapp: A Web Application to Annotate and Monitor***
2 ***SARS-CoV-2 Mutations***

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1 **Abstract**

2 The avalanche of genomic data generated from the SARS-CoV-2 virus requires the
3 development of tools to detect and monitor its mutations across the world. Here, we
4 present a webtool, *coronapp*, dedicated to easily processing user-provided
5 SARS-CoV-2 genomic sequences and visualizing current worldwide status of
6 SARS-CoV-2 mutations.

7 The webtool allows users to highlight mutations and categorize them by frequency,
8 country, genomic location and effect on protein sequences, and to monitor their
9 presence in the population over time.

10 The tool is available at <http://giorgilab.unibo.it/coronapp/> for the worldwide dataset
11 and at <http://giorgilab.unibo.it/coronannotator/> for the annotation of user-provided
12 sequences. The full code is freely shared at
13 <https://github.com/federicogiorgi/giorgilab/tree/master/coronapp>

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15

16 **KEYWORDS:** COVID-19; SARS-CoV-2; mutations; genetics; web application

17

18 **Data Availability Statement**

19 The data that support the findings of this study derive from the GISAID consortium
20 and are openly available in Github , in Rdata format for the R environment, in files
21 results.rda and metadata.rda, at the following link:

22 <https://github.com/federicogiorgi/giorgilab/tree/master/coronapp/data>

23

1 **Introduction**

2 SARS-CoV-2 is a novel pathogenic enveloped RNA beta-coronavirus causing a
3 severe illness in human hosts known as coronavirus disease-2019 (COVID-19). The
4 predominant COVID-19 illness is a viral pneumonia, often requiring hospitalization
5 and in some cases intensive care [1]. With almost 27.5 million laboratory-confirmed
6 positive cases worldwide as of 9 September 2020 and an estimated case fatality rate
7 across 204 countries of 5.2%, COVID-19 has become a global health challenge in
8 only a few months [2]. SARS-CoV-2 infection depends on the recognition of host
9 angiotensin converting enzyme 2 (ACE2), exposed on the cell surface in human lung
10 tissues [3,4]. SARS-CoV-2 spike glycoprotein binds ACE2, mediating membrane
11 fusion and cell entry [5]. Upon cell entry, the virus subverts host cell molecular
12 processes, inducing interferon responses and eventually apoptosis [6].

13 To date, much effort has been made to develop therapeutic strategies to limit
14 SARS-CoV-2 transmission and replication, but no treatment or vaccine has proven
15 effective against the virus, and repurposing of approved therapeutic agents has been
16 the main practical approach to manage the emergency so far [7]. As viruses mutate
17 during replication, the emergence of SARS-CoV-2 sub-strains and the challenge of a
18 probable antigenic drift require attention, especially for vaccine development [8].

19 Although sequence analyses of SARS-CoV-2 have shown that genomic variability
20 is very low [9], new SARS-CoV-2 mutation hotspots are emerging due to the high
21 number of infected individuals across countries and to viral replication rates [10].
22 Three major SARS-CoV-2 clades known as clade G, V, and S have emerged, showing
23 a different geographical prevalence [10]. The most frequent mutation detected so far
24 defines the G clade and causes an aminoacidic change, aspartate (D) or glycine (G), at
25 position 614 (D614G) of the viral Spike protein [11].

26 Continual genomic surveillance should be considered to monitor the possible
27 appearance of viral subtypes characterized by altered tropism, or causing more
28 aggressive symptoms. Constant and widespread monitoring of mutations is also a
29 powerful means of informing drug development and global or local pandemic

1 management. The Global Initiative on Sharing All Influenza Data (GISAID) has
2 collected to date (9 September 2020) over 90,000 publicly accessible SARS-CoV-2
3 sequences. The GISAID effort has made it possible to compare genomes on a
4 geographical and temporal scale and an increasing number of laboratories have started
5 to sequence COVID-19 patient samples worldwide [12,13]. Several online tools have
6 been developed to monitor the evolution of the virus from a phylogenetic perspective,
7 such as Nextstrain [14], or to visualize epidemiological data such as number of cases
8 and deaths [15]. However, no online tool currently exists to annotate user-provided
9 SARS-CoV-2 genomic sequences, which may derive from specific GISAID subsets
10 or from sequencing efforts of individual laboratories. Neither does any tool
11 specifically monitor the prevalence of specific SARS-CoV-2 mutations associated to
12 particular geographic regions or protein locations, nor their frequency in the
13 population over time.

14 To overcome these limitations, we have developed *coronapp*, a web application
15 with two purposes: real-time tracking of SARS-CoV-2 mutational status and
16 annotation of user-provided viral genomic sequences. Our tool enables users to easily
17 perform genomic comparisons and provides an instrument to monitor SARS-CoV-2
18 genomic variance, both worldwide and by uploading custom and locally produced
19 genomic sequences. The webtool is available at <http://giorgilab.dyndns.org/coronapp/>
20 and the full source code is shared on Github
21 <https://github.com/federicogiorgi/giorgilab/tree/master/coronapp>

22

23 **Results**

24 The webtool *coronapp* is available at the website
25 <http://giorgilab.dyndns.org/coronapp/> and it automatically provides the user with the
26 current status of SARS-CoV-2 mutations worldwide. The app also allows users to
27 annotate user-provided sequences (Figure 1 A). There are multiple functionalities of
28 *coronapp*, described in the following paragraphs.

29

1 **Current Status of SARS-CoV-2 mutational data**

2 A worldwide analysis is shown, generated using data from GISAID. Specifically, we
3 processed all SARS-CoV-2 complete (>29,000 sequenced nucleotides) genomic
4 sequences, excluding low-quality sequences (>5% undefined nucleotide “N”) and
5 viruses extracted from non-human hosts.

6 The underlying database is updated weekly, and we provide the date of the last
7 version as a reference for studies based on the data provided. We indicate the number
8 of samples processed and the total number of mutational events detected (Figure 1 A).
9 We also show the number of distinct mutated loci. Currently, this number is slightly
10 below 20,000, meaning that two thirds of the original Wuhan SARS-CoV-2 genome
11 has been affected by mutations and/or sequencing errors (the full length of the
12 reference genome is 29,903 nucleotides, based on sequence id NC_045512.2).

13

14 **Mutation frequency in SARS-CoV-2 proteins**

15 We show the frequency of mutations along the length of every SARS-CoV-2 protein,
16 reporting in the X-axis the amino acid position and on the Y-axis its frequency, either
17 as number of observed samples carrying the mutation, the base 10 logarithm of that
18 number, or the percentage over all sequenced samples. In the example in Figure 1 B,
19 we show the most frequent mutations affecting the viral Spike protein S,
20 distinguishing silent mutations and amino acid-changing mutations (including the
21 introduction of STOP codons). For Spike, the mutations appear to be evenly
22 distributed in frequency along the protein length, with the most frequent mutation
23 being the aforementioned D614G. Mouse-over functionality is provided to allow the
24 user to identify the selected mutation (e.g. N439K in Figure 1 B).

25

26 **The SARS-CoV-2 mutation table**

27 The user can visualize or download the full table of mutations on which the webtool
28 operates (Figure 2 A). This table is frequently updated and allows the user to specify a
29 worldwide or a country-specific dataset. The table also provides a Search function to

1 look for specific variants or sample ids, and it can be viewed online or downloaded in
2 full as a Comma-Separated Values (CSV) file.

3 The table shows every mutation in a specific geographical area, reporting:

- 4 • the GISAID sample ID (useful for cross-reference with the GISAID database
5 and other analyses based on it, e.g. Nexstrain).
- 6 • The country where the sample was collected.
- 7 • The position of the mutation, on the reference genome (refpos) and on the
8 sample (qpos).
- 9 • The sequence at the mutation site, on the reference genome (refvar) and on the
10 sample (qvar).
- 11 • The length of the sample genome (qlength); the reference genome is 29,903
12 nucleotides long.
- 13 • The protein affected by the mutation or, if the mutation is extragenic, the
14 denomination of the untranslated region (UTR), e.g. 5'UTR or 3'UTR.
- 15 • The effect of the mutation on the amino acid sequence of the protein (variant).
16 This uses the canonical mutational standard, indicating the original amino
17 acid(s), the position on the protein, and the mutated amino acid(s). An asterisk
18 (*) indicates a STOP codon, while the letters indicate amino acids in IUPAC
19 code. E.g. a mutation P315L indicates a leucine mutation (L) on the amino
20 acid location 315, normally occupied by a proline (P). Nucleotide mutations
21 can be silent, i.e. not yielding any aminoacidic change, e.g. the mutation
22 F106F, where the codon of phenylalanine 106 is affected but without changing
23 the corresponding amino acid. As in the previous column, mutations affecting
24 UTR regions are simply reported as the location of the nucleotide affected.
- 25 • The class of the mutation, of which there are currently 10 types:
 - 26 ○ SNP: a change of one or more nucleotides, determining a change in
27 amino acid sequence.
 - 28 ○ SNP_stop: a change of one or more nucleotides, yielding the generation
29 of one or more STOP codons.

-
- 1 ○ SNP_silent: a change of one or more nucleotides with no effect in
2 protein sequence.
- 3 ○ Insertion: the insertion of 3 (or multiples of 3) nucleotides, causing the
4 addition of 1 or more amino acids to the protein sequence.
- 5 ○ Insertion_stop: the insertion of 3 (or multiples of 3) nucleotides, causing
6 the generation of a novel STOP codon.
- 7 ○ Insertion_frameshift: the insertion of nucleotides not as multiples of 3,
8 causing a frameshift mutation.
- 9 ○ Deletion: the deletion of 3 (or multiples of 3) nucleotides, causing the
10 removal of 1 or more amino acids to the protein sequence.
- 11 ○ Deletion_stop: the removal of 3 (or multiples of 3) nucleotides, causing
12 the generation of a novel STOP codon.
- 13 ○ Deletion_frameshift: the deletion of nucleotides not as multiples of 3,
14 causing a frameshift mutation.
- 15 ○ Extragenic: a mutation affecting intergenic or UTR regions.
- 16 • The extended annotation of the protein region affected by the mutation (e.g.
17 “Spike” for “S” or “Predicted phosphoesterase, papain-like proteinase” for
18 NSP3, the Non-Structural Protein 3).
- 19 • The full name of the variant (varname), in the format
20 proteinName:AApositionAA, to allow for unique denomination of viral
21 proteome variants.
- 22

23 **Mutational overview**

24 The user is also provided with a general overview of the mutational status of the
25 selected country or the entire world (Figure 2 B). Six bar plots provide a summary and
26 highlights of the dataset, specifically:

- 27 • The most mutated samples, indicating which samples (in GISAID IDs) carry
28 the highest number of mutations

-
- 1 • The overall mutations per sample, indicating the distributions of mutations per
2 sample. It has been previously reported [10] that the current mode for
3 mutation number compared to the reference NC_045512.2 genome is 7.5.
- 4 • The most frequent events per class. Classes are the same as reported in the
5 mutation table and are described in the previous paragraph.
- 6 • The most frequent events per type. Individual mutation types are shown as
7 specific nucleotides events, e.g. cytosine to thymidine transitions (C>T),
8 guanosine to thymidine transversion (G>T) or even multinucleotide mutations
9 (e.g. GGG>AAC, observed in the Nucleocapsid protein). As reported before,
10 nucleotide transitions seem to be the most abundant SARS-CoV-2 type of
11 mutational event worldwide [11].
- 12 • The most frequent events, either in nucleotide coordinates or in aminoacidic
13 coordinates. Currently, the most frequent events are four mutations affecting
14 SARS-CoV-2 genomes belonging to clade G, which is the most sequenced
15 worldwide and predominant in Europe. These mutations are A23403G
16 (associated to the already mentioned D614G mutation in the Spike protein),
17 C3037T, C14408T and C241T.

18

19 **Analysis of mutations over time**

20 The *coronapp* webtool allows users to monitor the abundance and frequency of any
21 SARS-CoV-2 mutation in any country specified (Figure 3 A). Both plots in this
22 section report continuous dates on the X-axis, starting on the day of the first collected
23 SARS-CoV-2 genome available on GISAID: December 24, 2019.

24 The “abundance” plot reports on the Y-axis the number of samples carrying a
25 selected mutation in a particular day, in the specified country or worldwide. Since the
26 date reported is the collection date (not the submission date to the GISAID database),
27 there is usually a drop towards the right part of the plot, as there are fewer sequences
28 collected approaching the day of the analysis. The “frequency” plot on the other hand
29 normalizes the abundance of mutations by the total number of sequences generated on

1 each day. The plot currently shows a sharp increase in clade G-associated mutations
2 (e.g. S:D614G), as these mutations are most frequent in countries where sequencing is
3 more pervasive (e.g. United Kingdom).

4

5 **Annotation of user-provided SARS-CoV-2 genomic sequence.**

6 *coronapp* provides the user with an optional tool, *coronannotor*, providing the
7 optional possibility of uploading one or more SARS-CoV-2 genomic sequences,
8 which can be complete or partial. The format of the sequences is standard FASTA,
9 and an example input FASTA containing 12 sequences is provided (Figure 3 B). The
10 analysis is almost instantaneous and shows an overall breakdown of the most mutated
11 samples and most frequent mutations in the dataset. Moreover, a full table of all
12 detected mutations is provided: this can be visualized and searched on the web
13 browser or downloaded as a standard CSV file. Finally, a mutation frequency plot is
14 provided, allowing the user to visualize mutation frequency in selected proteins.

15 The user can easily return to the worldwide status of the app by refreshing or
16 reopening the page.

17

18 **Discussion**

19 Our webtool *coronapp* provides a fast, simple tool to annotate user-provided
20 SARS-CoV-2 genomes and visualize all mutations currently present in viral
21 sequences collected worldwide. The results provided by this instrument can have
22 several applications. The main purpose of *coronapp* is to help medical laboratories at
23 the front lines of COVID-19 fight with the opportunity to quickly define the
24 mutational status of their sequences, even without dedicated bioinformaticians.

25 Additionally, it enables scientists to perform mutational co-variance analyses and
26 to identify present and future significant functional interactions between viral
27 mutations, as previously attempted for the influenza virus and the human
28 immunodeficiency virus (HIV) [16]. Another application is the identification of the
29 most frequent mutations in specific protein regions: for example, our tool can quickly

1 identify that the most frequent mutation in the Spike protein, D614G, lies outside the
2 known interaction domain with the human protein ACE2, which spans roughly
3 between Spike amino acids 330 and 530 [17].

4 A recently published structural model simulating the effect of the D614G mutation
5 on the 3D structure of the spike protein has suggested that this mutation may result in
6 a viral particle which binds ACE2 receptors less efficiently, due to the masking of the
7 host receptor binding site on viral spikes [18]. The same researchers have reported a
8 possible correlation of the D614G form with increased case fatality rates,
9 hypothesizing that this mutation may lead to a viral form which is better suited to
10 escape immunologic surveillance by eliciting a lower immunologic response [18].
11 The *coronapp* analysis highlighted in Figure 1 B shows that a mutation located within
12 the Spike/ACE2 interaction domain is the change of Asparagine (N) to a Lysine (K)
13 in position 439 of the Spike sequence; this mutation could affect the protein folding or
14 its affinity with ACE2, as Asparagine is less charged than the basic amino acid
15 Lysine.

16 One of *coronapp*'s key strengths is to help prioritize scientific efforts on specific
17 aminoacidic variations that could affect the efficacy of anti-viral strategies or the
18 development of a vaccine by tracking the most frequent mutations in the population.
19 A further novelty of *coronapp* is that it provides a mean to assess the growth or
20 decline of specific mutations over time, in order to identify possible viral adaptation
21 mechanisms.

22 We provide not only the webtool, but also all the underlying code for the
23 annotation and visualization steps on a public Github repository, in order to help other
24 computational scientists in the ongoing battle against COVID-19. Furthermore, the
25 *coronapp* structure and concept could be expanded to other current and future
26 pathogens as well (e.g. the seasonal influenza or HIV), in order to monitor the
27 mutational status across proteins, countries and time.

28

29 **Materials and methods**

1 The webtool *coronapp* has been developed using the programming language R and is
2 based on a Shiny server (current version 1.4.0.2) running on R version 3.6.1. The app
3 is based on two distinct files, *server.R* and *ui.R*, managing the server functionalities
4 and the browser visualization processes, respectively. The results visualization utilizes
5 both basic R functions and Shiny functionalities; for tooltip functionality, *coronapp*
6 uses the R package *googleVis* v0.6.4, which provides an interface between R and the
7 Google visualization API [19].

8 The core of the annotation of the user-provided sequences rests in the NUCMER
9 (Nucleotide Mummer) alignment tool, version 3.1 [20]. Nucmer output is processed
10 by UNIX and R scripts provided in Github within the *server.R* file.

11

12

1 **Authors' contributions**

2 DM drafted the manuscript and performed the mutational analysis and literature
3 search. LT developed the user interface code and drafted the methodological parts of
4 the manuscript. EF worked on graphical interface of the webtool. FR wrote the
5 manuscript and performed literature search. FMG designed the study, developed the
6 server code, finalized the manuscript and provided financial support. All authors
7 tested the webtool and provided original contributions to its development. All authors
8 read and approve the final manuscript.

9

10 **Competing interests**

11 The authors have declared no competing interests.

12

13 **Acknowledgements**

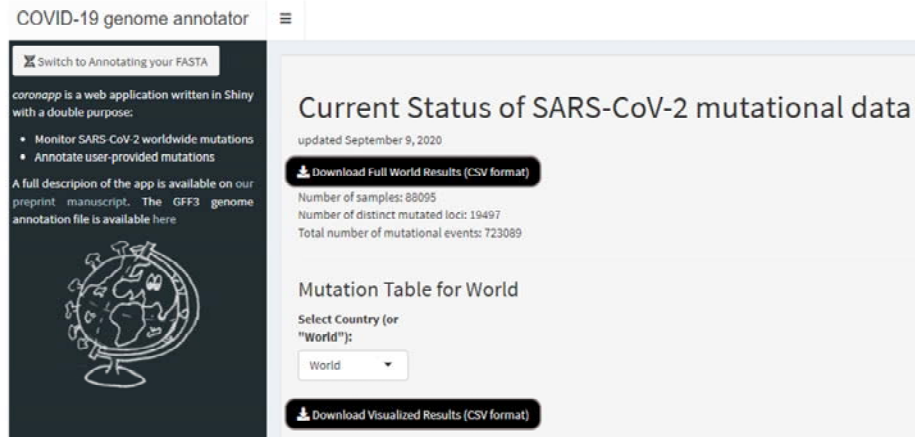
14 We thank the Italian Ministry of University and Research for their support, under the
15 Montalcini Grant 2016.

16

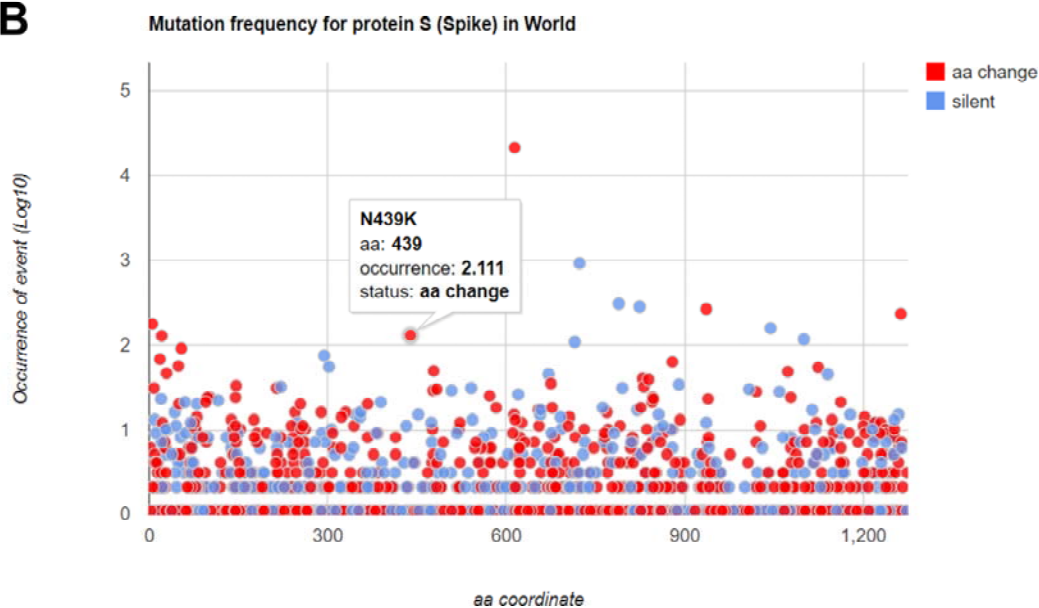
17

1 Figures and Legends

A



B



2

3 Figure 1 Overview of *coronapp*

4 A. Screenshot of the entry page of *coronapp* showing the basic tool description, the
5 interface to upload user-provided sequences and the overall summary of the mutations
6 detected worldwide. B. Common interface showing mutation frequency in
7 SARS-CoV-2 proteins, with occurrence of the mutation on the Y-axis and protein
8 coordinate on the X-axis. Red dots indicate amino acid (aa)-changing mutations, and
9 blue dots indicate silent mutations. Tooltip functionality is also provided to identify
10 and quantify each mutation on mouse-over.

11

A

Showing results for World

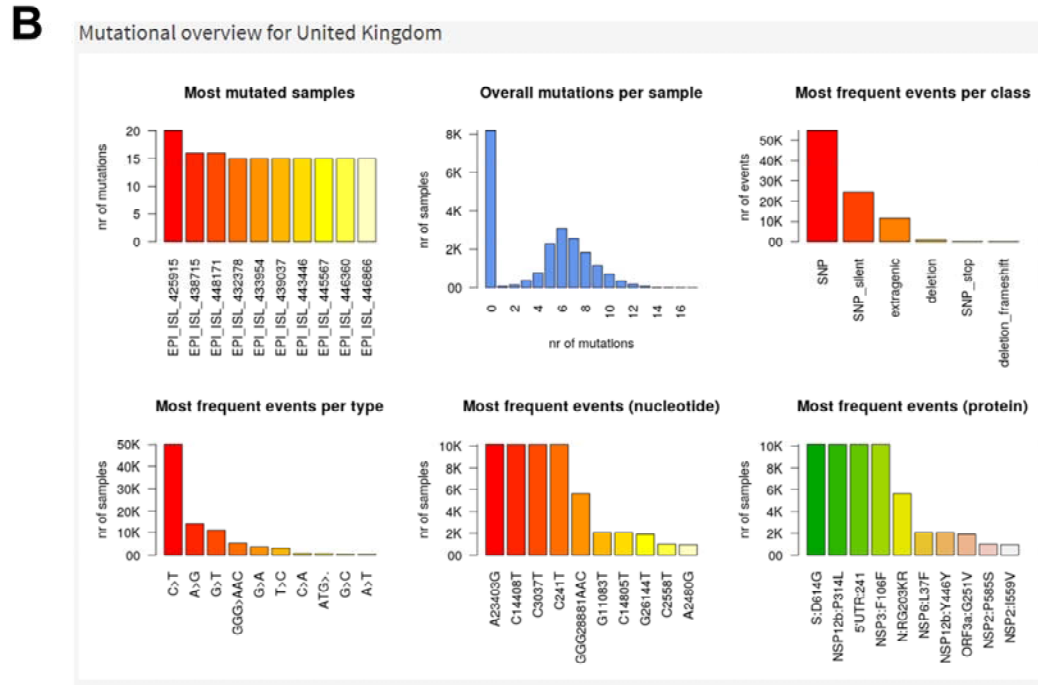
Download Full table (CSV format)

Show 10 entries Search:

sample	country	refpos	refvar	qvar	qpos	qlength	protein	variant	varclass	annotation	varname
EPI_ISL_415708	Switzerland	4	A	T	4	29903	S'UTR	4	extragenic		S'UTR:4
EPI_ISL_415708	Switzerland	241	C	T	241	29903	S'UTR	241	extragenic		S'UTR:241
EPI_ISL_415706	Switzerland	3037	C	T	3037	29903	NSP3	F106F	SNP_silent	Predicted phosphoesterase, papain-like proteinase	NSP3:F106F
EPI_ISL_415706	Switzerland	14408	C	T	14408	29903	NSP12b	P314L	SNP	RNA-dependent RNA polymerase, post-ribosomal frameshift	NSP12b:P314L
EPI_ISL_415706	Switzerland	15324	C	T	15324	29903	NSP12b	N619N	SNP_silent	RNA-dependent RNA polymerase, post-ribosomal frameshift	NSP12b:N619N
EPI_ISL_415706	Switzerland	23403	A	G	23403	29903	S	D614G	SNP	Spike	S:D614G
EPI_ISL_416497	France	4	A	T	4	29862	S'UTR	4	extragenic		S'UTR:4
EPI_ISL_416497	France	241	C	T	241	29862	S'UTR	241	extragenic		S'UTR:241
EPI_ISL_416497	France	2416	C	T	2416	29862	NSP2	Y537Y	SNP_silent	Non-Structural protein 2	NSP2:Y537Y
EPI_ISL_416497	France	3037	C	T	3037	29862	NSP3	F106F	SNP_silent	Predicted phosphoesterase, papain-like proteinase	NSP3:F106F

Showing 1 to 10 of 192,208 entries

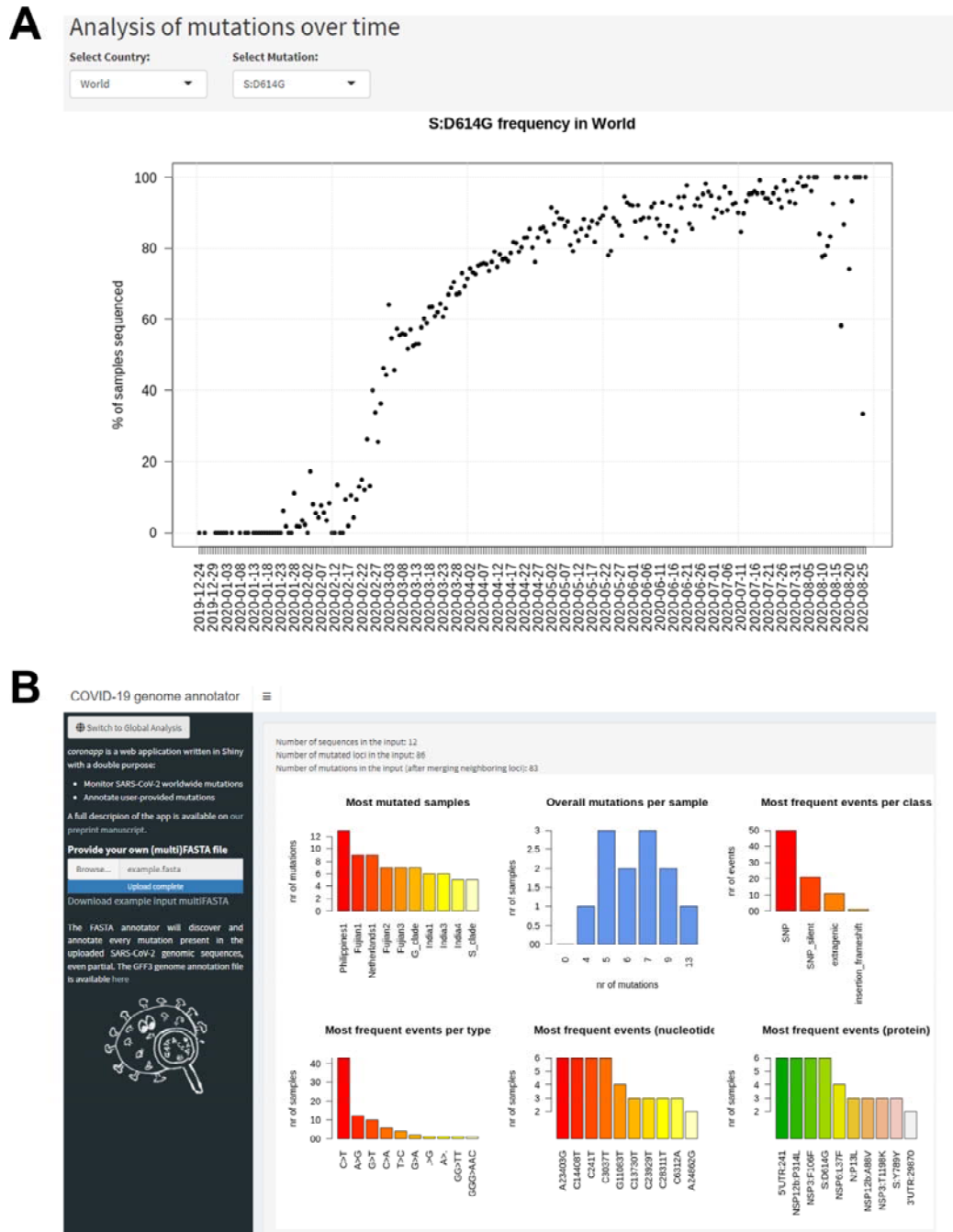
Previous 1 2 3 4 5 ... 19221 Next



1

2 **Figure 2 Mutation table and overview in *coronapp***

3 **A.** Result table of *coronapp*, available both for worldwide-precomputed and
 4 user-input analyses. A “download full table” button is provided to allow the user to
 5 perform larger-scale analyses autonomously. **B.** Barplots showing the most mutated
 6 samples, overall sample mutations and most frequent mutation events, classes and
 7 types. This analysis is also available both for worldwide-precomputed and user-input
 8 analyses.



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