

# Representative sequences of SARS-CoV-2 variants that are considered VOC<sup>™</sup>, VOI, or VUM by the World Health Organization.

Enabled by Data from



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## I. Descriptions of WHO variants

Designations: WHO Tracking SARS-CoV-2 variants
VOC, Variant of Concern
VOI, Variant of Interest
VUM, Variant Under Monitoring
VOC-LUM, VOC Lineages Under Monitoring

#### **Spike Variant Data Table**

The accompanying spreadsheet lists a natural form of a SARS-CoV-2 sequence that was found to be most representative of each of the WHO designated VOC, VOI, or VUM (See "WHO Tracking SARS-CoV-2 variants": WHO Tracking SARS-CoV-2 variants). The spreadsheet provides details of mutational patterns and of sampling frequencies of these common forms both 60 days prior to this report and historically within entire collection at GISAID.

The accompanying fasta files include (i) a full-length genome codon aligned set of the each of the representative forms, and (ii) an alignment of Spike reference forms. We emphasize Spike to facilitate monitoring and reagent design for immune response and testing of enhanced infectivity. The most commonly sampled natural version of each variant listed as a "baseline form"; this form generally corresponds to an early consensus form of the lineage, and to the sequence of founder of the lineage. Mutations in other regions of the genome that are most typical of particular forms are included in the full-length genome alignments.

Variants are continuously evolving and so the most representative forms may change over time.

Note that the WHO reference sequences set is a subset of a larger set of VOI's we have defined based on our own selection strategies at Los Alamos; the larger full set is also available at GISAID. Our analyses are generally based on actual Spike sequences, not Pango lineages designations, as some very distinctive forms of Spike are repeated in many different Pango lineages, and conversely, some Pango lineages contain very distinctive forms of Spike. Also, Pango lineage designations often change over time. Current information about the designated Pango lineages most associated with each variant form is provided, to enable the variants to be recognized in the context of the current designations in the literature and appropriately reflect the World Health Organization SARS-CoV-2 variant nomenclature.

## II. Reference Alignments for WHO GISAID Sequences

#### WHO-GISAID\_RepresentativeSpikeSequences\_2022-05-31

The team at Los Alamos National Lab (cov.lanl.gov) builds an alignment of all intact SARS-CoV-2 Spike sequences available in GISAID, and the most common form of the spike protein for each WHO variant is resolved. We then identify the most common nucleotide sequence encoding that spike variant, and the natural sequence with the lowest EPI\_ISL number that perfectly matches that gene sequence is selected to represent that particular form of Spike.

#### WHO-GISAID RepresentativeGenomes CodonAligned 2022-05-31.fasta

We next build an alignment based on all full-length (ORF-complete) GISAID sequences that encodes the exact representative Spike form of interest. The consensus of the rest of the genome among that set is determined. The earliest-sampled intact natural sequence in GISAID EpiCoV that completely matches the full genome consensus is selected to be the representative sequence of that variant. If no intact natural sequence completely matching the consensus is available, the natural sequence that is closest to the consensus is selected (i.e. with the lowest Hamming distance); in practice, these have differed from the consensus by only a few bases.

We the build a codon-aligned genome alignment of these most representative sequences spanning all open reading frames, so the reading frame is preserved throughout the alignment of the full genome. At the top of the full genome alignment is the GISAID SARS-CoV-2 reference strain, immediately followed by a separate aligned sequence specifying each open reading frame/gene in isolation; this is intended as a guide to enable one to easily resolve where mutations are occurring throughout the genome.

## III. Key to the Variants Table

**Key to Variants Table.** This spreadsheet/table contains the most common forms of Spike among WHO designated VOI/VOC/VUMs sampled in GISAID EpiCoV (<a href="www.gisaid.org">www.gisaid.org</a>). Table counts were updated based on GISAID data from May 25, 2022; the alignment data upload precedes the report date (May 31, 2022) because it requires several days to complete the analyses.

#### Column headers with notes, Variants Table:

- A. WHO Classification group of Variant
- B. Dates of WHO classifications
- **C. WHO designation:** The WHO Greek letter variant designation from WHO Tracking SARS-CoV-2 variants; \*indicates continued monitoring.
- **D.** Pango lineage most often associated with a particular WHO variant. Pango lineage descriptions can be found at <u>Pango Lineages</u>.
- **E. GISAID reference sequence accession number** for the full-length genome most representative variant. This is the first sampled sequence with the most common variant form that available in GISAID.
- F. GISAID reference sequence virus name.
- **G.** WHO variant linked to the most common Pango lineage in which it is found. A short list of mutations relative to the most appropriate Omicron baseline variant is provided.
- **H. Most common Spike backbones.** Mutation list relative to the Wuhan reference strain for the most common natural form of each Spike variant lineage.

Note that "-" indicates a deletion at a site (e.g. Y144-, the Y at position 144 in the reference Spike is deleted), "+" indicates an insertion following the specified site (e.g. +143T indicates a T was inserted after position 143 in the reference Spike), and "\_" indicates the ancestral value (e.g. D614\_ indicates the ancestral D at site 614; it is equivalent to D614D).

Colors highlight mutation in regions of interest:

Blue: Addition of positive charge near the furin cleavage site: 675, 677, 681 are positively charged, or the H655Y substitution

Green: NTD supersite: 13-20, 140-158, 242-264

Magenta: RBD: 330-521

Red: D614\_, The ancestral Spike D614 amino acid is the dominant form in this lineage, the underscore indicates ancestral.

Turquoise: The Heptad Repeat 1 region, HR1: 908-985

- I. Is the representative sequence available in the current Spike fasta file (X == yes)
- J. Is the representative sequence available in the Reference Genome fasta file (X== yes)
- **K.** Number of sequences that exactly match this pattern in Spike, full data. This tally includes all data in our quality filtered GISAID data set starting in December 2019.
- L. Number of sequences that contain this pattern, full data. The most common form of Spike representing a lineage is always part of an evolving lineage. This tally represents the number of variants that contain the full specified set of mutations (the "sequence backbone"), but that may also contain additional mutations. As lineages spread over time, they diversify, and the most common form becomes a smaller percentage of the total. As new variants of variants become more prevalent, we identify them as distinctive common forms, and add the most interesting of these to the "variants-of-variants" listing.
- M. Number of sequences that exactly match this pattern, last 60 days. This tally is a rough indication of whether a particular form of Spike is still present in a contemporary GISAID global sampling, is declining and being replaced by other variants, or is no longer sampled.
- N. Number of sequences that contain this pattern, last 60 days. See above.

- O. All Pango Lineages that contain sequences that exactly match this pattern in Spike (with counts). Quite distinctive Spikes are often assigned to an array of Pango lineages. Some are closely related, but some not obviously phylogenetically related; such exceptions can arise due to recombination, or mis-classification; they also can change over time as Pango lineage designations can be reassigned. The current counts are based on Pango lineage assignments in the 2022-05-31 GISAID data, after being filtered for quality control (QC) at cov.lanl.gov.
- P. All Pango Lineages that are associated with sequences that contain this pattern in Spike, including Spikes that also contain additional mutations (with counts).
- Q. Total count of sequences with the Pango lineage designation in which the Spike variant is most commonly found. These counts are based on our QC filtered set used at cov.lanl.gov.
- R. Number of Spikes in the Pango lineage that exactly match this pattern.
- S. Fraction of the Pango lineage that exactly match this pattern (column R/Q).
- T. Number in the Pango lineage that contain this pattern, but that also contain additional mutations.
- U. Fraction of the Pango lineage sequences that contain this pattern (column T/Q).
- V. The three countries where the exact Spike variant is most commonly sampled historically (with counts).
- W. The three countries where Spikes that include the variant mutations are most commonly sampled historically (with counts).
- X. The three countries where the exact Spike variant is most commonly sampled in the last 60 days (with counts).
- Y. The three countries where Spikes that include the variant mutations are most commonly sampled in the last 60 days (with counts).
- Z. The lowest (first) EPI\_ISL accession number among the set of sequences that exactly matches the pattern of Spike mutations listed in column H, used in the Spike fasta file.
- AA. The full sequence name used in the full length genome fasta file.
- BB. Brief notes regarding the lineage.

# **Citation for these reference alignments:**

The tools we use for tracking SARS-CoV-2 variants were first described in Korber et al. Cell. 2020 Aug 20;182(4):812-827.e19. doi: 10.1016/j.cell.2020.06.043.

Further information can be found at cov.lanl.gov.