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Situation Report

21st June 2022

The National Institute of Health Doutor Ricardo Jorge, I.P. (INSA) has analysed 37248 SARS-CoV-2 genome sequences so far, obtained from positive samples collected in more than 100 laboratories/hospitals/institutions, across 304 municipalities.

The genetic diversity of SARS-CoV-2 has been monitored based on an average of 511 sequences/week, since the beginning of June, 2021. These sequences have been obtained from positive samples collected randomly throughout the 18 Districts of Portugal Mainland and the Autonomous Regions of the Azores and Madeira, covering an average of 139 municipalities per week (Figures 1 e 2).



Figure 1. Geographic coverage (at the municipality level) of the last weekly nationwide sequencing survey, which resulted in 283 new SARS-CoV-2 genome sequences from 114 municipalities.

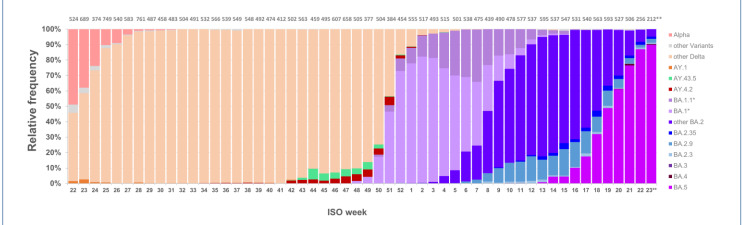


Figure 2. Evolution of the weekly relative frequency of the SARS-CoV-2 (sub-)lineages circulating in Portugal between ISO weeks 22 (31st May – 6th June, 2021) and 23 (6th – 12th June, 2022). The values on top of each bar indicate the number of sequences analysed per week in the context of the weekly random nationwide surveys. The plot highlights: i) BA.1 (and sub-lineage BA.1.1), BA.2 sub-lineages with relative frequency ≥1% in week 22 (sampling closed), BA.3, BA.4 and BA.5 lineages (all classified as *Omicron* by the WHO); ii) *Delta* sublineages of interest (AY.1, AY.4.2 e AY.43.5); and iii) the Alpha variant. *BA.1 = BA.1 and sub-lineages (with exception of BA.1.1 and sub-lineages). **The frequencies presented for the last week under analysis (ISO week 23) might change in the next report, given that some data from that period is still being processed. BA.1.1 = BA.1.1 and sub-lineages.



Main highlights:

The Variant of Concern (VOC) *Omicron*, according to the <u>WHO</u> classification, currently includes several (sub)lineages identified by the "BA" prefix. The nomenclature of the various sub-lineages is under constant review and refinement (https://www.pango.network/). The sequences identified in Portugal are reclassified weekly and the result are available on our website https://insaflu.insa.pt/covid19/. Whenever relevant, some of these sub-lineages will be highlighted in this report.

Lineage BA.5

Lineage BA.5 presents several genetic traits of interest, such as the presence of the mutations **L452R** and **F486V in Spike** (both affect Spike regions that interact with the human cells and mediate adherence/immune evasion). BA.5 also presents the deletion del69-70 in Spike, which is responsible for the SGTF profile, so this laboratory indicator has also been used as a proxy to estimate its frequency. However, due to the recent decrease in the number of tests performed with the *TaqPath – ThermoFisher* methodology (allowing the monitoring of the proportion of samples with "SGTF" profile) by the different laboratories, SGTF data now lack robustness and, as such, are no longer used for BA.5 lineage monitoring.

The nationwide weekly surveys by sequencing show that, since its first detection in week 13, the **lineage BA.5** has been markedly increasing in relative frequency, being clearly dominant in Portugal since week 19 (9th - 15th May, 2022) and presented a relative frequency of 88% in the last national sequencing survey in week 23 (6th-12th June, 2022) (Figure 2).

Lineage BA.2

- o In Portugal, BA.2 has been firstly detected in random nationwide sequencing surveys in ISO week 52 (27th December 2021 2nd January, 2022) (Figure 2), becoming dominant on week 8 (21st–27th February, 2022). After reaching a maximum relative frequency of 95% on week 13 (28th May 3rd Abril 2022), this lineage presents a continuous descrease in relative frequency, having ceased to be dominant since week 19 (9th–15th May, 2022), point from which lineage BA.5 became dominant in Portugal (Figures 2 and 3).
- The sublineage BA.2.12.1, detected in all 7 Regions (Figure 3) with 1.3% relative frequency peak at week 19 (9th-15th May, 2022), has since shown a decreasing trend. This lineage has raised particular interest internationally due to its additional mutation L452Q in Spike (affecting the same protein site described above) as well as due to its increasing circulation in some countries, namely in the United States of America.
- Another BA.2 sublineage (BA.2.35), with the additional mutation L452R in Spike protein, has been recently detected and is being monitored. The L452R mutation is associated with resistance to neutralizing antibodies and is also known as a marker mutation of other variants of interest/concern, such as *Delta*, *Kappa* and *Omicron* BA.4/BA.5. Since its first detection on 1st March, 2022 (ISO week 9), BA.2.35 relative frequency has shown a decreasing trend since week 18 (where it reached 3.6%), presenting a relative frequency of 1.4% on week 23 (analysis ongoing).



• Lineage BA. 1

BA.1 has been firstly identified in Portugal by mid November, 2021, and was dominant between ISO weeks 51, 2021 (20th-26th November) and 7, 2022 (14th-20th February), having reached its maximum circulation in week 2 (95.6%, 10th-16th January, 2022) (Figure 2). The current circulation of BA.1 in Portugal is estimated to be residual (<1% since week 16, 18th-24th April, 2022).

Lineages BA.4 and BA.3

Since week 19 (9th-15th May, 2022), 8 sequences of the BA.4 lineage of Omicron variant were identified in Portugal, associated with confirmed cases in 5 out of the 7 Regions (Figure 3). Similarly to BA.5, with which it shares multiples mutations of interest, BA.4 lineage (also classified as VOC) has a relevant circulation in some countries, in particular in South Africa. In Portugal, no BA.3 case has been detected since week 11 (14th-20th March, 2022; see Report from 29th March, 2022).

Recombinants

- The co-circulation of several lineages/variants in the community increases the possibility of the occurrence of mixed infections, i.e., the same individual being simultaneously infected by more than one of them. In this context, mixture of their genetic material might occur, resulting in a hybrid genetic profile, commonly designated as "recombinant".
- Currently, several SARS-CoV-2 recombinants have been described at a global scale (e.g., Delta+Omicron BA.1 or BA.1+BA.2), with novel designations being assigned to the recombinants with epidemiological/functional relevance. In Portugal, the few recombinant viruses identified so far were detected in sporadic cases in the weekly random surveys. Among these, we highlight cases associated with the recombinants internationally designated as XM, XN, XE, XH and XQ, being that all are characterized by a hybrid genetic profile where an initial portion of the genome corresponds to lineage BA.1 and the remaining portion to lineage BA.2. Similarly to other recombinant SARS-CoV-2 detected so far in Portugal, there is no evidence that these recombinants might present functional differences (e.g., differences in transmissibility and immune evasion) from the parental lineages BA.1, BA.2 and BA.5.
- Dynamic tables summarizing the frequency and geotemporal spread of the variants/lineages identified so far as well as the mutations of interest in protein Spike in each of them are available in the <u>website</u>.



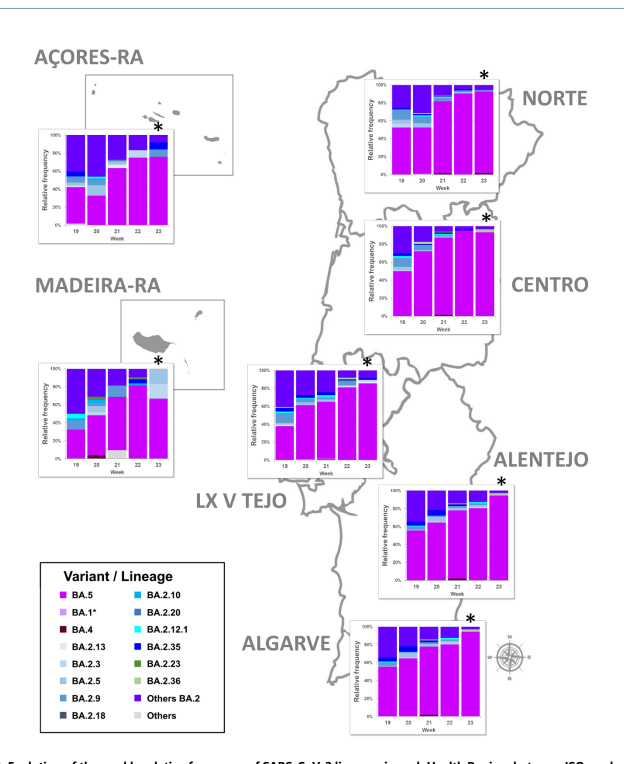


Figure 3. Evolution of the weekly relative frequency of SARS-CoV-2 lineages in each Health Region, between ISO weeks 19 (9th – 15th May) and 23 (6th – 12th June), 2022. The plots highlight the BA.1 (including its sublineage BA.1.1), BA.2 sub-lineages with relative frequency \geq 1% in at least one Region in week 22 (sampling closed), BA.4 and BA.5 (all classified as *Omicron* by WHO). BA.1 = BA.1 and sub-lineages (with exception of BA.1.1 and descendent sub-lineages).

*It is expected that the frequencies presented for the last week under analysis (ISO week 23) might change in the next report, given that some data from that period is still being processed.