Pathogen Genomics Center of Excellence Situation Report

2024-04-26

# Key Findings

* Current data reflects a mixture of JN.1 descendents as the likely near term variants.
* Globally no other variants with unusual characteristics have been identified as having unusual growth.
* Some other point

# Situation Update Details

* Based on what - XYZ(?), JN.1 and descendents continue to dominate. Some recombinations from JN.1 and other BA.5 variants are being monitoredtracked, but have yet to show significant growth relative other variants.
* Together this diversity suggests steady evolution against general population immunity with no indications of a variant driven wave of COVID-19 infections.
* As of MM/DD/YYYY, there were X samples from MM/DD/YYYY - MM/DD/YYYY, some comment on trend
* Some text here about image one. There is this variant that’s here
* Some text about image two
* Image 3 has this
* Findings from a site’s analysis of national data

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| Figure 1: Proportion of variants by year. |

Source: [NorthWest Genomics Center of Excellence](https://coe-test-org.github.io/sitrep-demo/notebooks\nwcoe-preview.html#cell-fig-countprop)

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| Figure 2: From the New York Times: A couple of observations are obvious. First when cases start to rise, deaths follow with a lag. Second, we have had three spikes in cases so far and in each successive instance the mortality has risen by a smaller amount. This suggests that, thankfully, we are getting better at treating this disease. It is NOT a function of increased testing because positivity rates have not been falling. |

Source: [New England Genomics Center of Excellence](https://coe-test-org.github.io/sitrep-demo/notebooks\necoe-preview.html#cell-fig-state-analysis)

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| Figure 3: COVID-19 fatalities, outputs from New York Times modeling. |

Source: [Virginia Genomics Center of Excellence](https://coe-test-org.github.io/sitrep-demo/notebooks\vacoe-preview.html#cell-fig-fatality-plot)

# Site Summaries

Source: [Article Notebook](https://coe-test-org.github.io/sitrep-demo/index.qmd.html)

* Washington State Department of Health - highest variant proportion is 50%
* Georgia Department of Public Health probablity of detection: 60 and the consensus genomes are uploaded to public repositories like GISAID and GenBank.
* Massachusetts Department of Health prop - 50
* Virginia Deparment of Health - 60%

## Section

This is a simple placeholder for the manuscript’s main document (Knuth 1984).

* To monitor SARS-CoV-2 in Washington state, Washington state Department of Health (WA DOH) implemented a Sentinel Surveillance system, a type of genomic surveillance that tracks SARS-CoV-2 variants across the state.
* Laboratories across the state, including the Washington state Public Health Laboratories (PHL) will sequence SARS-CoV-2 from collected specimens. Raw sequencing data is assembled into consensus genomes using publicly available bioinformatics pipeline, and the consensus genomes are uploaded to public repositories like GISAID and GenBank. This report demonstrates how the NW PGCoE utilizes SARS-CoV-2 sequencing data to monitor emerging variants biweekly, forecast emerging SARS-CoV-2 variants, and infers relative abundance estimates of SARS-CoV-2 variants in the wastewater. Previous work that looked at the disease severity of SARS-CoV-2 variants is currently being implemented to analyze the disease severity of current variants utilizing hospitalization data. This work is ongoing and will be presented at a later time.

Knuth, Donald E. 1984. “Literate Programming.” *Comput. J.* 27 (2): 97–111. <https://doi.org/10.1093/comjnl/27.2.97>.