

## Comparative Genomics: SARS-CoV-2 vs SARS-CoV Tor2

### Sequences:

- SARS-CoV-2 (MN908947.3)
- SARS-CoV Tor2 (AY274119.3)

### Alignment Method:

- Full genome alignment using MAFFT (FFT-NS-2 strategy).
- Alignment length: 30,016 bases.

### Results:

- Mutations detected: 6,319.
- Genomic divergence: 21.05%.

### Interpretation:

- SARS-CoV-2 diverges significantly from SARS-CoV (2003).
- Early genome region shows substitution and indel hotspots.
- Divergence suggests changes in regulatory RNA regions and spike-mediated host interaction.

### Tools Used:

- Ubuntu (WSL)
- MAFFT v7.505
- Python (Biopython)
- Custom mutation counting script

### Conclusion:

The alignment confirms substantial sequence evolution between the two viral genomes and highlights regions potentially associated with biochemical/functional differences.