Exercise/Assignment

Case study: There are 7 reported positive cases of SARS-CoV-2 in location **X** which needed urgent sequencing to ascertain the potential threat. You have been provided with the raw sequencing data for all 7 samples, generated via Illumina NextSeq using the ARTIC protocol (FastQ Format). For all 7 samples you need to:

- 1. Assemble the full genome using Galaxy and Exatype
- 2. Assess the quality of each sample and flag those that might be poor quality
- 3. Assign lineage (pangolin) and clades (nextclade) for each sample and also assess quality based on pangolin and nextclade outputs
- 4. Compare the quality assignments between Galaxy and Exatype
- 5. Compare the lineage and clade assignments between Galaxy and Exatype
- 6. Identify any mutations or variants of concern/interest
- 7. Create a report that outlines this information and your recommendations to your Department of Health.

Find raw sequencing data and additional data files here:

Jottacloud – [9_10E] Galaxy pipeline