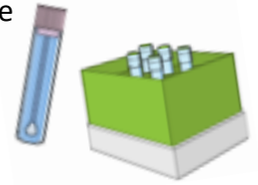


# SARS-CoV-2 Sequencing Timeline

## DAY 0: Specimens arrive at Alaska State Virology Laboratory

SARS-CoV-2 specimens submitted for sequencing are accessioned into the sequencing queue upon arrival at ASVL. Specimens submitted for diagnostic or confirmatory SARS-CoV-2 PCR testing will reflex to sequencing after a positive PCR result is obtained.



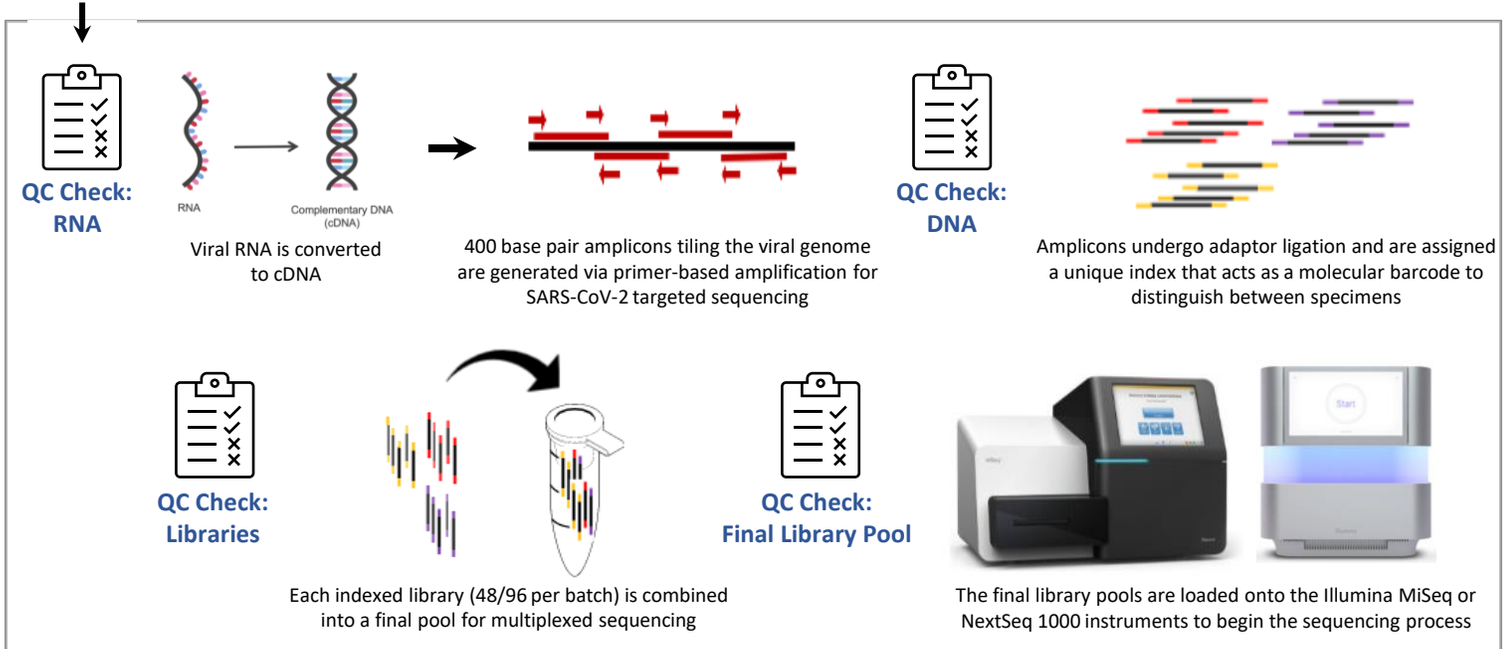
## DAY 1: Specimens are selected and batched for sequencing

ASVL sequences specimens on two Illumina sequencing platforms- the MiSeq and the NextSeq1000. Depending on the sequencing instrument used, sequencing batches consists of 46 to 94 patient specimens and 2 controls. During periods of high positive specimen volumes, specimen selection is based on a priority scheduling algorithm where samples associated with travel, rural areas, known outbreaks, vaccine breakthroughs, and reinfections are batched first. **As of May 2021, all positive SARS-CoV-2 specimens received at ASVL are being sequenced.**

## DAYS 2 & 3: Library Preparation

Sequencing libraries are prepared from extracted RNA in a 16-hour process comprised of a series of enzymatic reactions that begins with the conversion of viral RNA to cDNA and ends with the pooling of 48/96 "libraries" of barcoded DNA fragments generated from each specimen and control in the batch.

Extracted RNA



## DAYS 3 & 4: Sequencing



Prepared libraries are loaded on a flow cell where adaptors facilitate attachment of DNA fragments to the flow cell surface and clonal amplification begins. The resulting cluster of template DNA undergoes multiple rounds of sequencing to generate millions of reads. A unique fluorescent signal is emitted for each of the four bases (A,G,T,C) which allows the instrument to issue base calls for nucleotides at each location in the sequence.

## DAY 5: Genome Assembly and Data Analysis

Fragmented DNA reads are then assembled by aligning sequenced fragments to a reference genome (SARS-CoV-2 isolate Wuhan-Hu-1). Assembled genomes are assessed for quality before variant calls are issued based on nucleotide differences between the sequenced virus and reference.

