COVID-19 subject UPHS-0122

2021-04-01

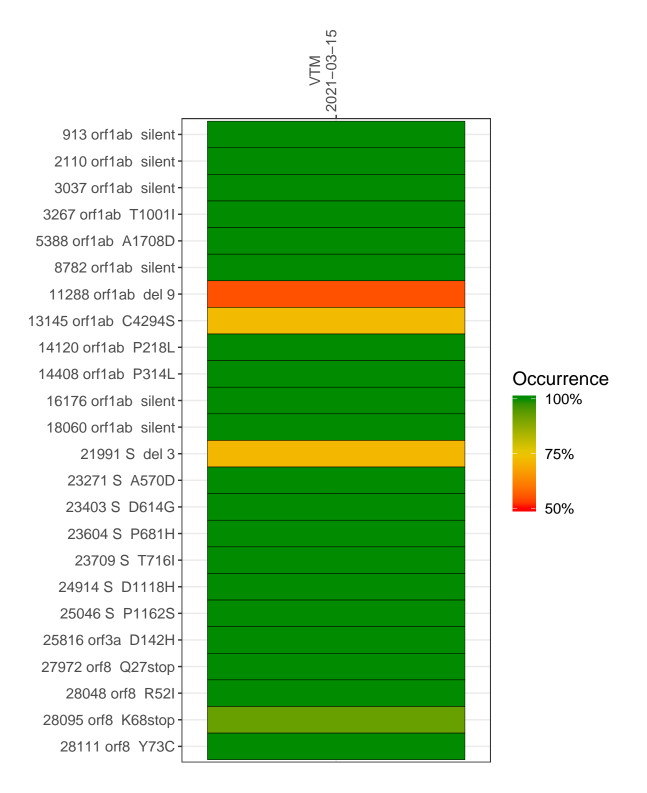
The table below provides a summary of subject samples for which sequencing data is available. The experiments column shows the number of sequencing experiments performed for each specimen. Experiment specific analyses are shown at the end of this report. Lineages are called with the Pangolin software tool (Rambaut et al 2020) for genomes with > 90% sequence coverage.

Table 1. Sample summary.

Experiment	Туре	Genomes	Sample type	Sample date	Largest contig (KD)	Lineage	Reference read coverage	Reference read coverage (>= 5 reads)
VSP1107-1	single experiment	NA	VTM	2021-03-15	17.61	B.1	99.0%	97.3%

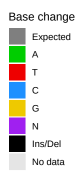
Variants shared across samples

The heat map below shows how variants (reference genome /home/everett/projects/SARS-CoV-2-Philadelphia/USA-WA1-2020) are shared across subject samples where the percent variance is colored. Variants are called if a variant position is covered by 5 or more reads, the alternative base is found in > 50% of read pairs and the variant yields a PHRED score > 20. Gray tiles denote positions where the variant was not the major variant or no variants were found. The relative base compositions of each experiment used to calculate tiles are shown in the following plot where the total number of position reads are shown atop of each plot.



VTM 2021-03-15

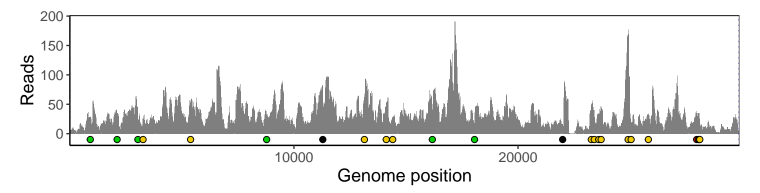
	2021-03-13
913 orf1ab silent	28
2110 orf1ab silent	32
3037 orf1ab silent	28
3267 orf1ab T1001I	26
5388 orf1ab A1708D	53
8782 orf1ab silent	29
11288 orf1ab del 9	44
13145 orf1ab C4294S	61
14120 orf1ab P218L	46
14408 orf1ab P314L	28
16176 orf1ab silent	45
18060 orf1ab silent	33
21991 S del 3	13
23271 S A570D	43
23403 S D614G	35
23604 S P681H	33
23709 S T716I	43
24914 S D1118H	177
25046 S P1162S	17
25816 orf3a D142H	31
27972 orf8 Q27stop	33
28048 orf8 R52I	32
28095 orf8 K68stop	26
28111 orf8 Y73C	14
)7–1
	P1107-1



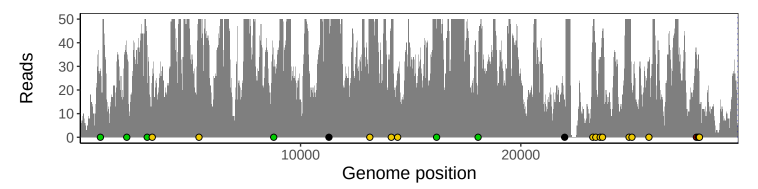
Analyses of individual experiments and composite results

$VSP1107-1 \mid 2021-03-15 \mid VTM \mid UPHS-0122 \mid genomes \mid single \; experiment$

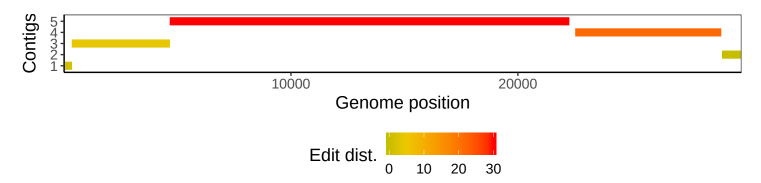
The plot below shows the number of reads covering each nucleotide position in the reference genome. Variants are shown as colored dots along the bottom of the plot and are color coded according by variant types: gray - transgenic, green - silent, gold - missense, red - nonsense, black - indel.



Excerpt from plot above focusing on reads coverage from 0 to 50 NT.



The longest five assembled contigs are shown below colored by their edit distance to the reference genome.



Software environment

Software/R package	Version
R	3.4.0
bwa	0.7.17-r1198-dirty
samtools	1.10 Using htslib 1.10
bcftools	1.10.2-34-g1a12af0-dirty Using htslib 1.10.2-57-gf58a6f3
pangolin	2.3.3
genbankr	1.4.0
optparse	1.6.0
forcats	0.3.0
stringr	1.4.0
dplyr	0.8.1
purrr	0.2.5
readr	1.1.1
tidyr	0.8.1
tibble	2.1.2
ggplot2	3.0.0
tidyverse	1.2.1
ShortRead	1.34.2
$\operatorname{GenomicAlignments}$	1.12.2
SummarizedExperiment	1.6.5
DelayedArray	0.2.7
matrixStats	0.54.0
Biobase	2.36.2
Rsamtools	1.28.0
GenomicRanges	1.28.6
$\operatorname{GenomeInfoDb}$	1.12.3
Biostrings	2.44.2
XVector	0.16.0
IRanges	2.10.5
S4Vectors	0.14.7
BiocParallel	1.10.1
BiocGenerics	0.22.1