

Run Info

Host Name GXB01190 (localhost)

Experiment Name
EIMock_6kbp_Sdysgalactiae_04012021
Sample ID
EIMock_6kbp_Sdysgalactiae_04012021
Run ID
484966ff-9253-4fd7-85df-c7d02096e2bf

Flow Cell Id FAO54786
Start Time January 4, 18:13

Run Length 5h 25m

Run Summary

Reads Generated487.74 KPassed Bases220.81 MbFailed Bases32.03 MbEstimated Bases262.99 Mb

Run Parameters

Flow Cell Type FLO-MIN106 SQK-LSK109 Kit -180 mV Initial Bias Voltage FAST5 Output **Enabled FASTQ Output Enabled BAM Output Enabled** Active Channel Selection **Enabled** Basecalling on Specified Run Length 72 hours

reference_files=

Read Until ["/data/references/S_dysgalactiae_ref.fasta"],filter_type=enrich,first_

 $channel = 1, last_channel = 512$

FAST5 Reads per File 4000

FAST5 Output Options **zlib_compress,fastq,raw**

FASTQ Reads per File 4000

Mux Scan Period 1 hour 30 minutes

Reserved Pores 0 %

Basecall Model High-accuracy basecalling

Alignment reference_files=["/data/references/the7references.fasta"]

Read Filtering min_qscore=7

Versions

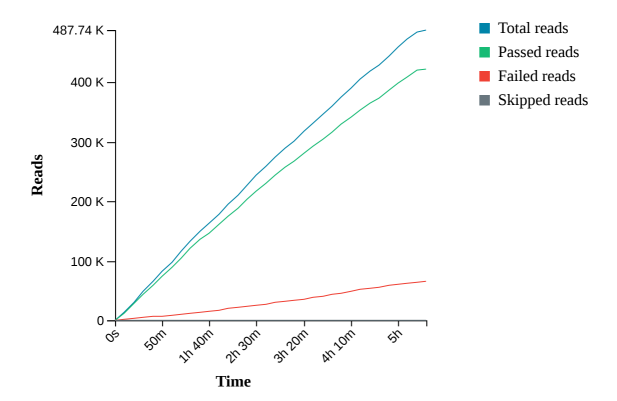
 MinKNOW
 20.10.6

 MinKNOW Core
 4.1.2

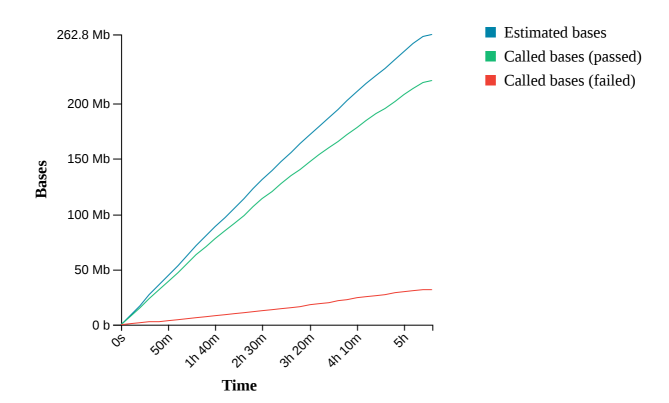
 Bream
 6.1.4

 Guppy
 4.2.3

Cumulative Output Reads

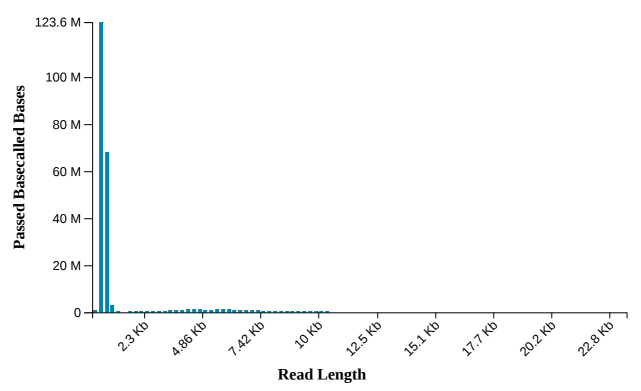


Cumulative Output Bases



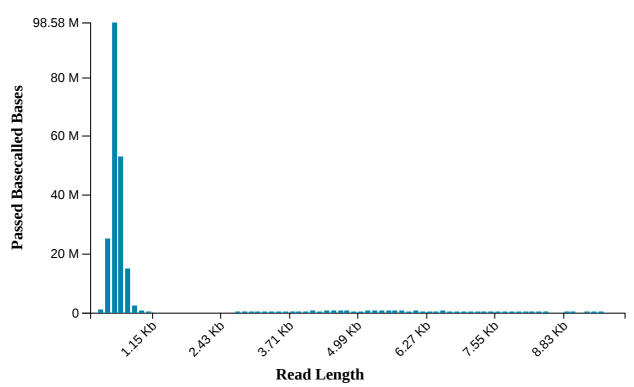
Read Length Histogram Estimated Bases - Outliers Discarded

Estimated N50: 494



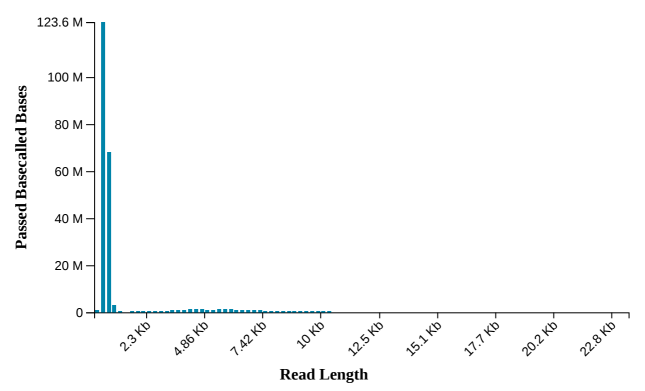
Read Length Histogram Basecalled Bases - Outliers Discarded

Estimated N50: 493



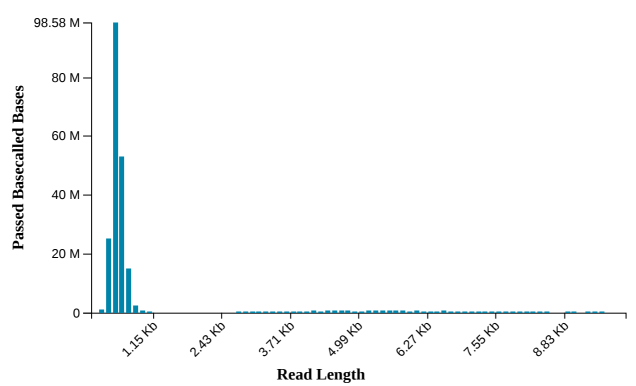
Read Length Histogram Estimated Bases

Estimated N50: 494

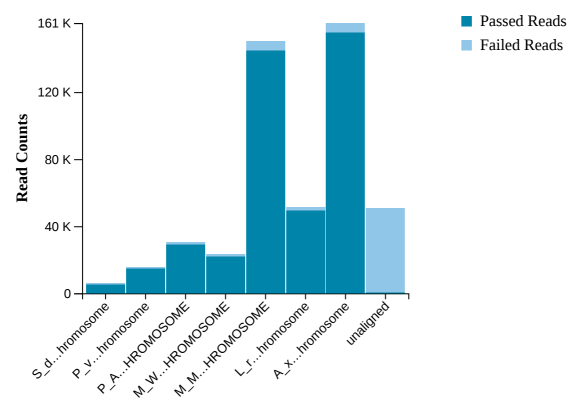


Read Length Histogram Basecalled Bases

Estimated N50: 493

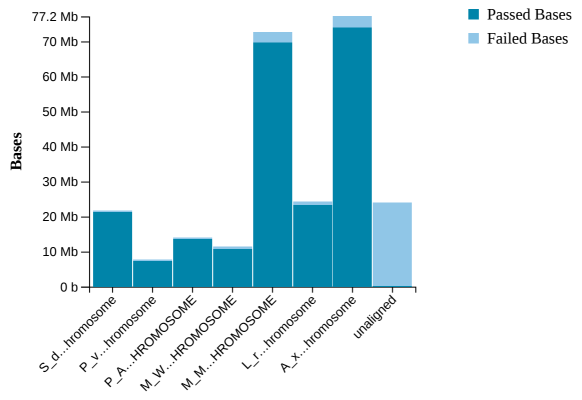


Alignment Target Hits (reads)



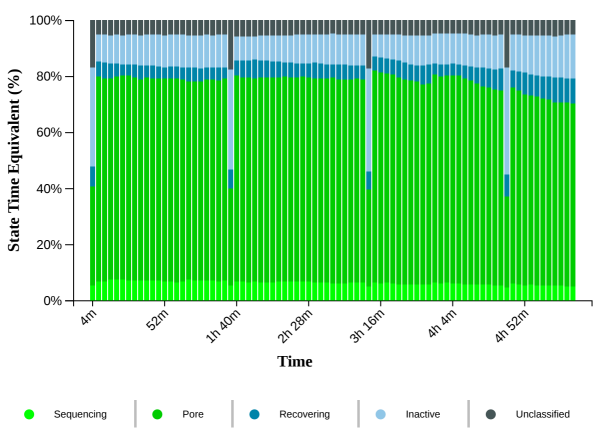
Alignment Target

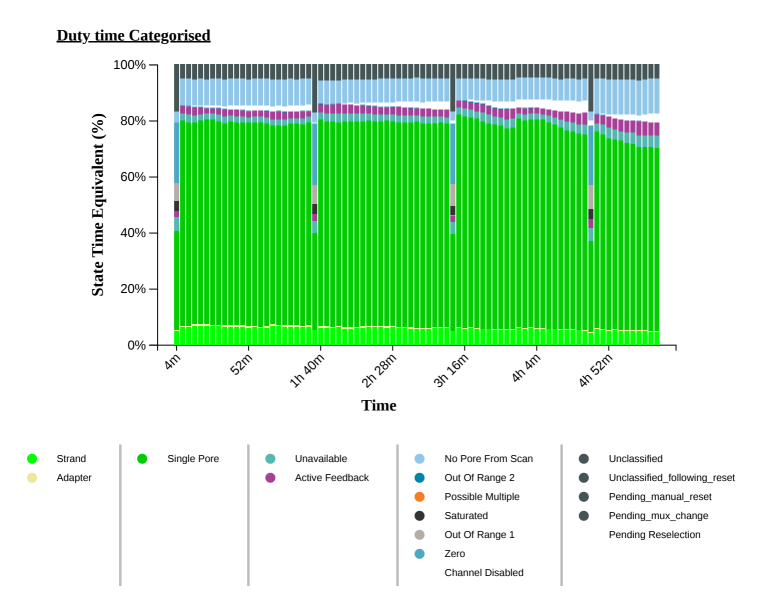
Alignment Target Hits (bases)



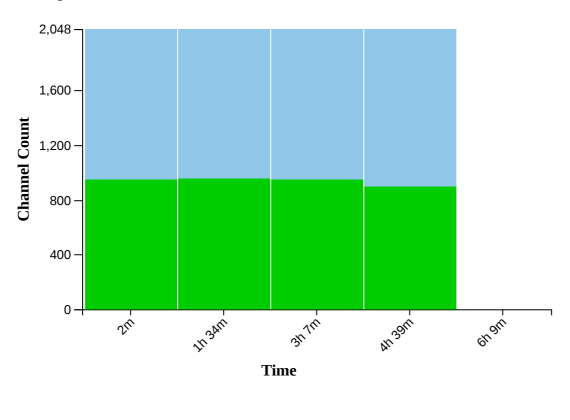
Alignment Target

Duty Time Grouped



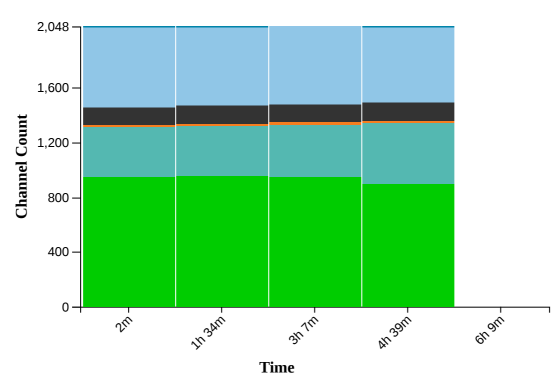


Mux Scan Grouped



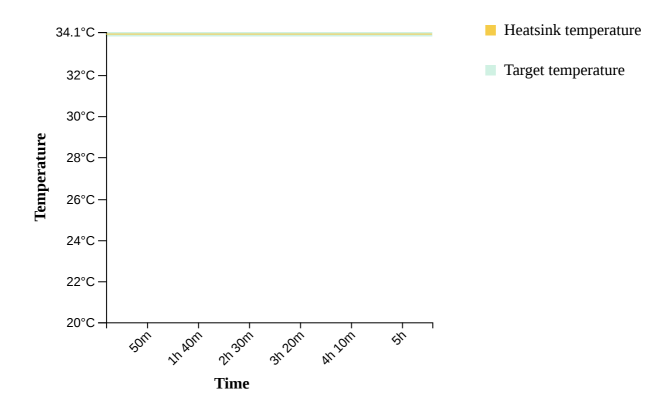
ActiveInactive

Mux Scan Categorised

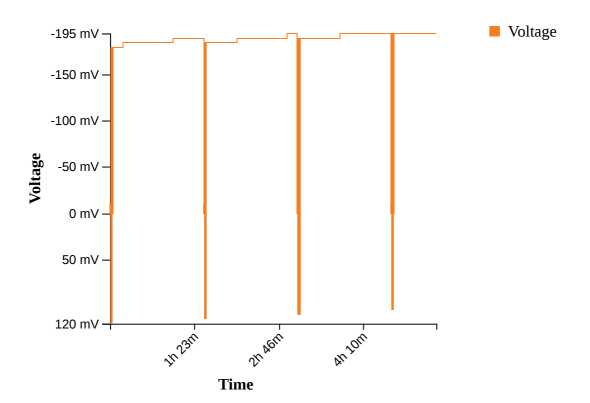


Single Pore
 Reserved Pore
 Unavailable
 Multiple
 Saturated
 Zero
 Other

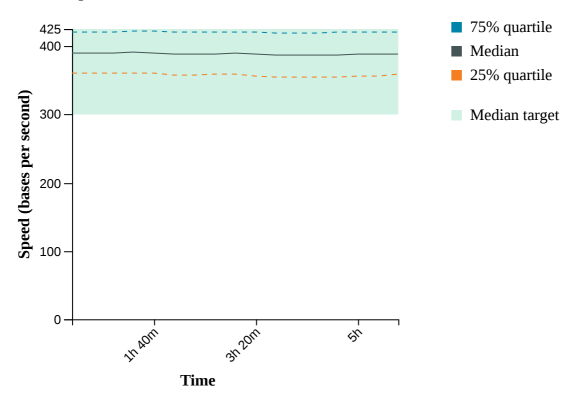
Temperature History



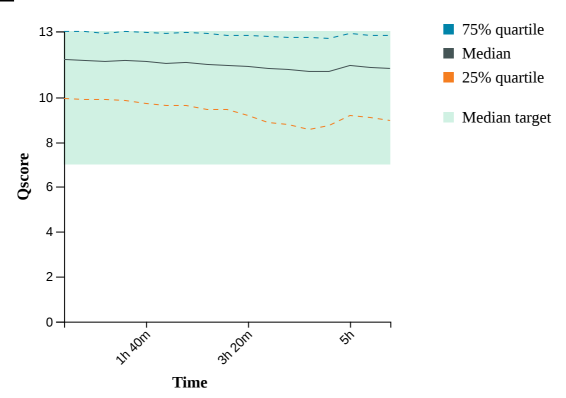
<u>Bias Voltage History</u>



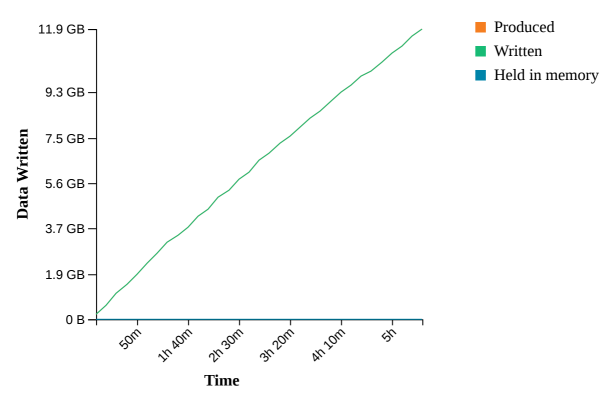
Translocation Speed



QScore



Disk Write Performance



Run Debug Messages

- Mux scan for flow cell FAO54786 has found a total of 901 pores. 448 pores available for immediate sequencing January 4, 22:56
- Performing Mux Scan January 4, 22:54
- Mux scan for flow cell FAO54786 has found a total of 953 pores. 472 pores available for immediate sequencing January 4, 21:24
- Performing Mux Scan January 4, 21:21
- Mux scan for flow cell FAO54786 has found a total of 961 pores. 469 pores available for immediate sequencing January 4, 19:51
- Performing Mux Scan January 4, 19:49
- Mux scan for flow cell FAO54786 has found a total of 952 pores. 463 pores available for immediate sequencing January 4, 18:19
- Performing Mux Scan January 4, 18:16
- Starting sequencing procedure January 4, 18:16
- Waiting up to 300 seconds for temperature to stabilise at 34.0°C January 4, 18:13