

#### **Run Info**

Host Name GXB03020 (localhost)

 Experiment Name
 ReadUntil\_38kbp\_Mmenrich\_con\_15042021

 Sample ID
 ReadUntil\_38kbp\_Mmenrich\_con\_15042021

 Run ID
 143d42a6-087f-4a6c-878f-6bbc737df140

Flow Cell Id FAP21384
Start Time April 15, 10:57

Run Length 1h 0m

#### **Run Summary**

Reads Generated34.47 KPassed Bases368.61 MbFailed Bases16.58 MbEstimated Bases382.29 Mb

#### **Run Parameters**

Flow Cell Type FLO-MIN106 SQK-LSK109 Kit Initial Bias Voltage -180 mV FAST5 Output **Enabled FASTQ Output Enabled BAM Output Enabled** Active Channel Selection **Enabled** Basecalling on Specified Run Length 72 hours 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/the7references.fasta"]

Read Filtering min\_qscore=7

#### **Versions**

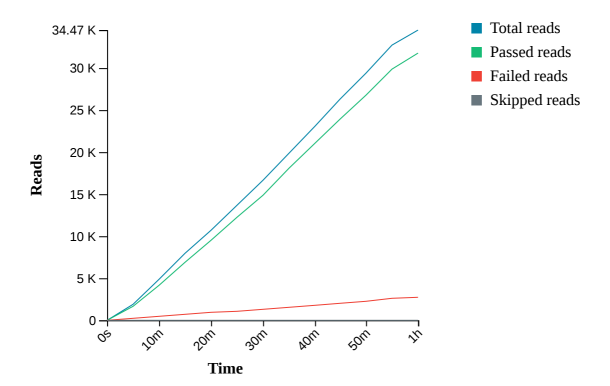
 MinKNOW
 21.02.5

 MinKNOW Core
 4.2.5

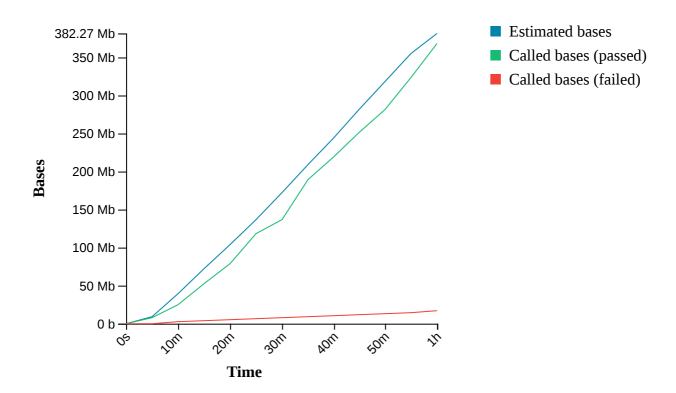
 Bream
 6.1.10

 Guppy
 4.3.4

### **Cumulative Output Reads**

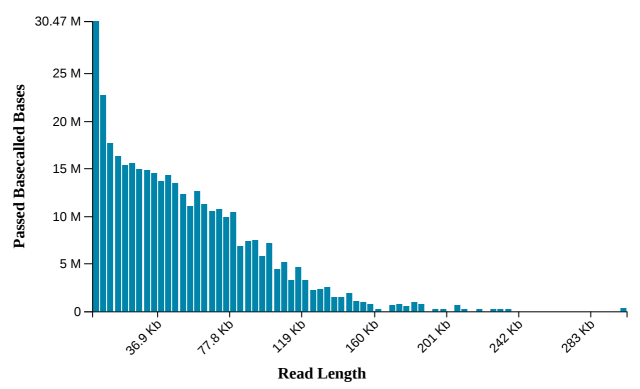


# **Cumulative Output Bases**



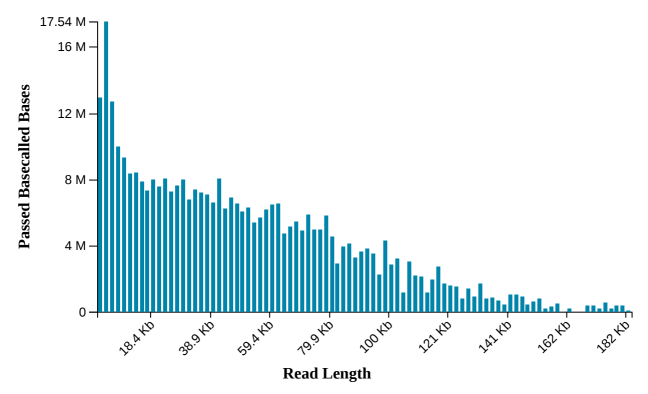
# Read Length Histogram Estimated Bases - Outliers Discarded

Estimated N50: 43.13 K



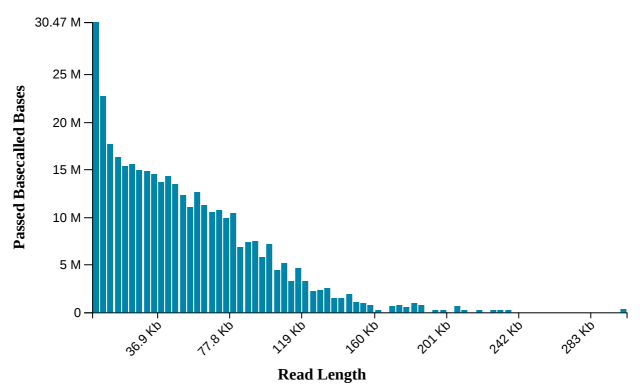
# Read Length Histogram Basecalled Bases - Outliers Discarded

Estimated N50: 42.57 K



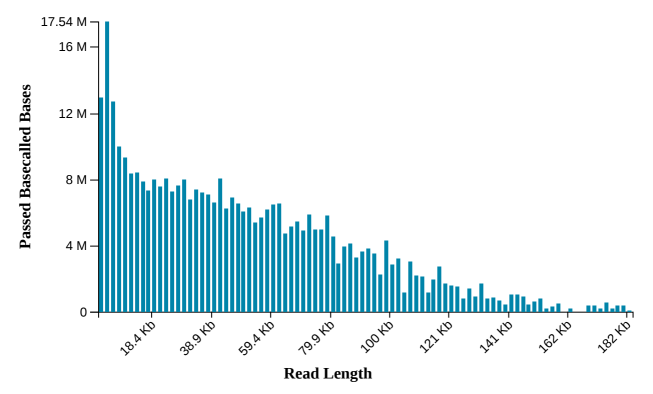
#### **Read Length Histogram Estimated Bases**

Estimated N50: 43.13 K

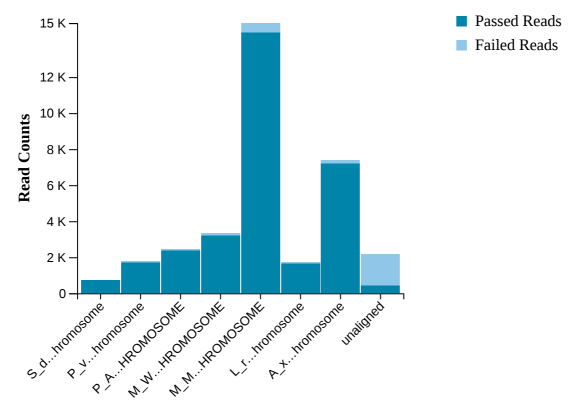


# **Read Length Histogram Basecalled Bases**

Estimated N50: 42.57 K

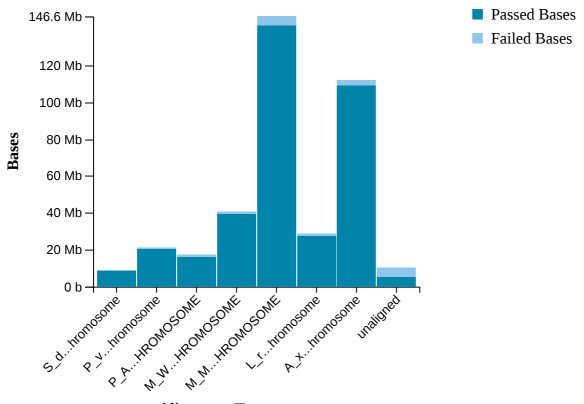


#### Alignment Target Hits (reads)



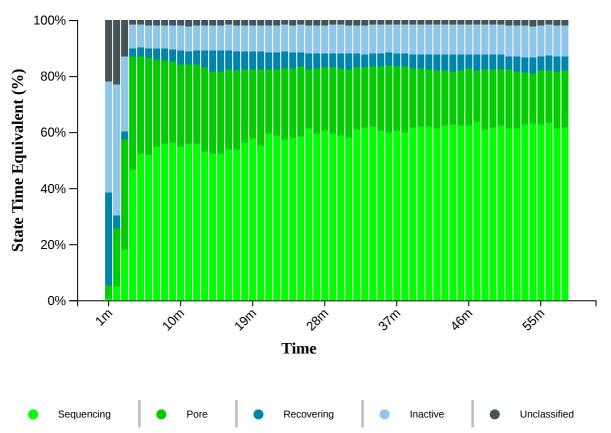
**Alignment Target** 

### **Alignment Target Hits (bases)**

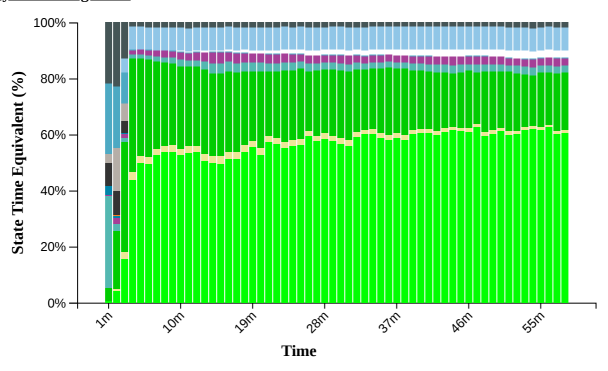


**Alignment Target** 

#### **Duty Time Grouped**

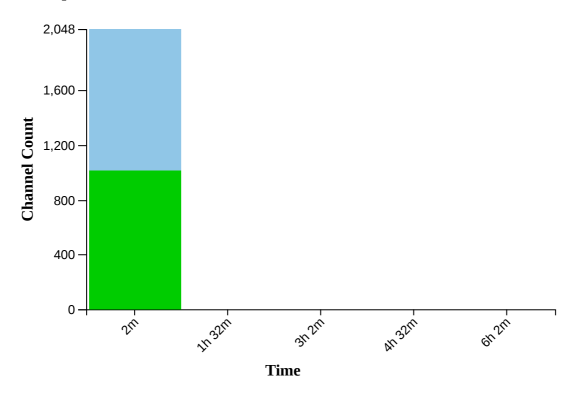


### **Duty time Categorised**

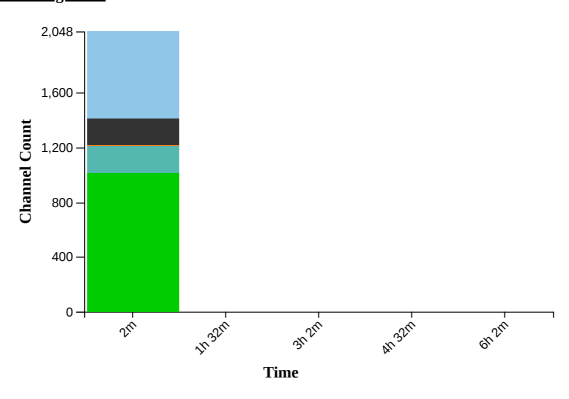




### **Mux Scan Grouped**





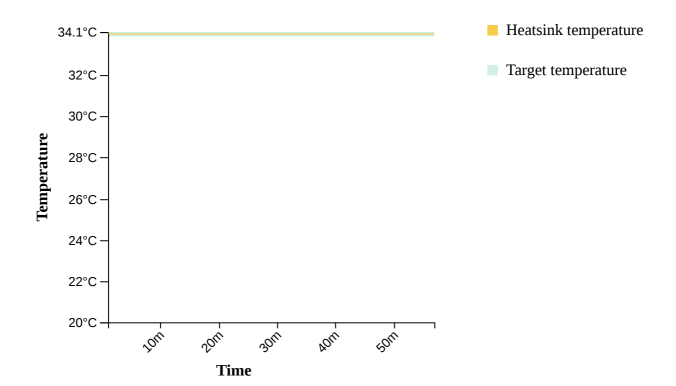


Inactive

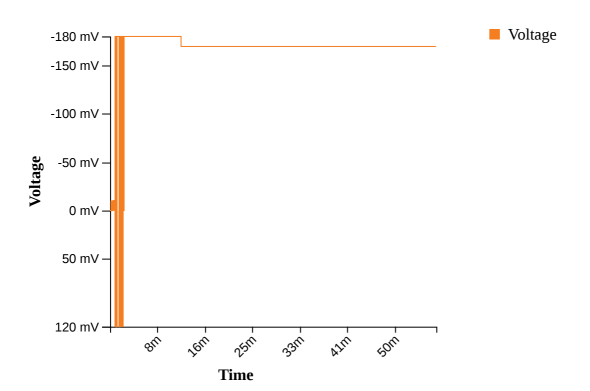
Active

Single Pore
 Reserved Pore
 Unavailable
 Multiple
 Saturated
 Zero
 Other

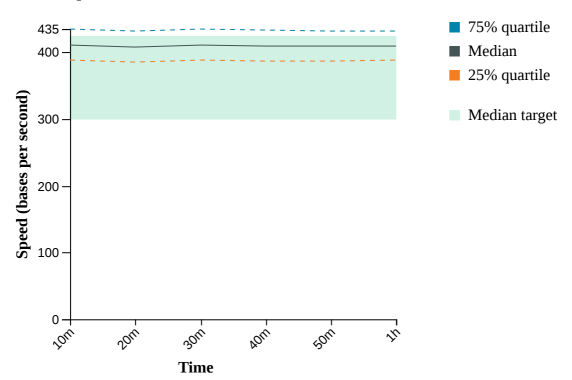
# **Temperature History**



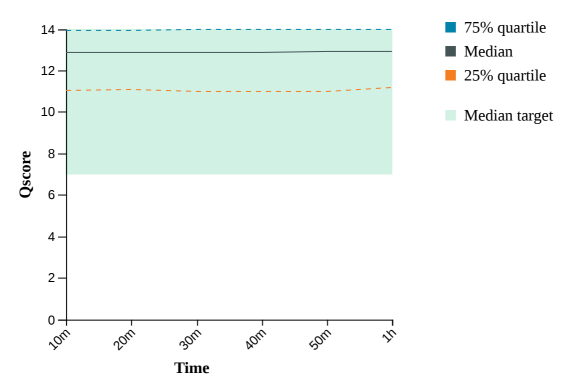
# <u>Bias Voltage History</u>



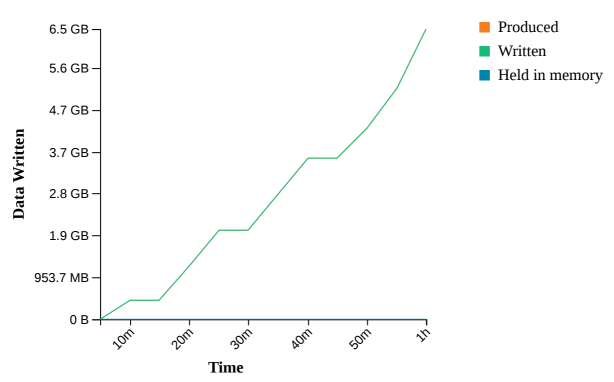
# **Translocation Speed**



# **QScore**



# **Disk Write Performance**



#### **Run Debug Messages**

- Mux scan for flow cell FAP21384 has found a total of 1019 pores. 470 pores available for immediate sequencing April 15, 11:03
- Performing Mux Scan April 15, 11:01
- Starting sequencing procedure April 15, 11:01
- Waiting up to 300 seconds for temperature to stabilise at 34.0°C April 15, 10:58