# AS21001\_lon\_V2\_RBC1\_BC10\_rawlib.basecaller

Overall QC Status: PASS Sample QC Status: PASS Fusion QC Status: PASS Variations QC Status: PASS Job: 20210826 AS21001 [3437]

**Type:** RNA FusionRNA SNP/InDel Targeted Mutations: Archer Comprehensive Targets NIH v1.3.1 Include Non-Targeted

Variants: No

Software Version: Suite\_Analysis\_v6.2.7

Analysis Date: 26-Aug-2021 5:52 Report Creator: mpvghtpe@gmail.com

Report Date: 19-Jun-2023 2:30



#### **Statistics**

#### **Molecular Barcode Statistics**

Total Fragments	Fragments with Complete Adapter	Number of Reads After Trimming Adapters
3,500,000	3,236,877	2,959,596

#### **Read Statistics**

Туре	Total Fragments (# / %)	Mapped (# / %)	Pass Alignment Filter (%)	On Target (%)
All Fragments	2,945,379 / 100.0	2,945,379 / 100.0	100.0	98.2
Unique Fragments	488,690 / 16.6	488,690 / 100.0	100.0	98.6

### **DNA/RNA Statistics**

Туре	DNA Reads (# / %)	RNA Reads (# / %)	Ambiguous Reads (# / %)
All Fragments	215,313.0 / 7.4	2,206,098.0 / 76.2	471,896.0 / 16.3
Molecular Bins	45,277.0 / 9.4	325,582.0 / 67.6	111,109.0 / 23.1
Average Molecular Bins per GSP2	77.13	554.65	189.28
Unique Start Sites	14,987.0 / 28.7	32,321.0 / 61.9	12,739.0 / 24.4
Average Unique Start Sites per GSP2	26.29	74.54	25.23
Average Unique Start Sites per GSP2 Control	57.42	222.33	48.33

### **QC Statistics**

Avg. Unique DNA And Ambiguous Start Sites Per GSP2	Avg. Unique RNA Start Sites Per GSP2 Control	
51.07	222.33	

### **Miscellaneous Statistics**

On Targ	get Deduplication Ratio
	6.00:1

## **DNA/RNA Fragment Lengths**

DNA Median Fragment Length	DNA Mean Fragment Length	RNA Median Fragment Length	RNA Mean Fragment Length
111.0	124.7	119.0	125.0

# **Reportable Variants**

None Found

# **Reportable Isoforms**

None Found