# AS23063\_49284006\_lon\_V2\_RBC2\_BC2 5\_rawlib.basecaller

Overall QC Status: PASS Sample QC Status: PASS Fusion QC Status: PASS Variations QC Status: PASS Job: 20231221 AS23063 [3519]

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:** 21-Dec-2023 4:34 Report Creator: mpvghtpe@gmail.com

Report Date: 21-Dec-2023 18:54



#### **Statistics**

#### **Molecular Barcode Statistics**

Total Fragments	Fragments with Complete Adapter	Number of Reads After Trimming Adapters
2,771,479	2,595,319	2,436,193

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#### **Read Statistics**

Туре	Total Fragments (# / %)	Mapped (# / %)	Pass Alignment Filter (%)	On Target (%)
All Fragments	2,425,213 / 100.0	2,425,213 / 100.0	100.0	99.1
Unique Fragments	634,570 / 26.2	634,570 / 100.0	100.0	99.1

### **DNA/RNA Statistics**

Туре	DNA Reads (# / %)	RNA Reads (# / %)	Ambiguous Reads (# / %)
All Fragments	207,284.0 / 8.6	1,774,244.0 / 73.9	420,729.0 / 17.5
Molecular Bins	66,613.0 / 10.6	410,878.0 / 65.4	151,065.0 / 24.0
Average Molecular Bins per GSP2	101.08	623.49	229.23
Unique Start Sites	24,880.0 / 31.9	47,585.0 / 61.1	18,800.0 / 24.1
Average Unique Start Sites per GSP2	39.69	102.05	33.74
Average Unique Start Sites per GSP2 Control	23.58	148.92	27.67

### **QC Statistics**

Avg. Unique DNA And Ambiguous Start Sites Per GSP2	Avg. Unique RNA Start Sites Per GSP2 Control	
72.62	148.92	

### **Miscellaneous Statistics**

On Target Deduplication Ratio
3.82:1

## **DNA/RNA Fragment Lengths**

DNA Median Fragment Length	DNA Mean Fragment Length	RNA Median Fragment Length	RNA Mean Fragment Length
139.0	148.8	137.0	143.6

# **Reportable Variants**

None Found

## **Reportable Isoforms**

☑ Passed all strong-evidence filters

Likely off-target mispriming event

Exact breakpoint known

☎ Cross contamination

✗ User-annotated false positive

■ Known fusion partners in Archer Quiver™

₹ Percent GSP2 reads below threshold

 $\Delta \hat{}$  Fusion expression imbalance

1 Low confidence

 $\triangle$  User-annotated true positive

% Intronic fusion

↓ Not enough unique start sites

O Transcriptional readthrough event

☐ Known ensembl paralogue

Fusion: SS18 → SSX1			
Filters: ☑ ⊚	Reads: 1873 (46.55%)	<u>Segments</u>	
GSP2: SS18_chr18_23612369_25A1_GSP2  Mutation Classification: Undefined	Start Sites: 214	chr18:23612496→23612363 SS18(-) NM_005637.3, exon:9 chrX:48123217→48123352	
Is Artifact: no		SSX1(+) NM_005635.3, exon:6	