AS22011_lon_V2_RBC1_BC22_rawlib.basecaller

Overall QC Status: PASS Sample QC Status: PASS Fusion QC Status: PASS Variations QC Status: PASS

Job: 20220408_AS22010_11 [4725]

Type: RNA FusionRNA SNP/InDel Targeted Mutations: Archer Comprehensive Targets NIH v1.3.1 2 Include Non-Targeted

Variants: No

Software Version: Suite_Analysis_v6.2.7

Analysis Date: 08-Apr-2022 9:26

Report Creator: mpvghtpe@gmail.com

Report Date: 19-Jun-2023 3:27



Statistics

Molecular Barcode Statistics

Total Fragments	Fragments with Complete Adapter	Number of Reads After Trimming Adapters
3,071,260	2,838,546	2,677,019

Read Statistics

Туре	Total Fragments (# / %)	Mapped (# / %)	Pass Alignment Filter (%)	On Target (%)
All Fragments	2,662,542 / 100.0	2,662,542 / 100.0	100.0	97.8
Unique Fragments	421,214 / 15.8	421,214 / 100.0	100.0	98.2

DNA/RNA Statistics

Туре	DNA Reads (# / %)	RNA Reads (# / %)	Ambiguous Reads (# / %)
All Fragments	236,848.0 / 9.1	2,080,627.0 / 79.9	287,449.0 / 11.0
Molecular Bins	46,675.0 / 11.3	295,342.0 / 71.4	71,627.0 / 17.3
Average Molecular Bins per GSP2	79.51	503.14	122.02
Unique Start Sites	19,938.0 / 31.3	38,204.0 / 59.9	14,651.0 / 23.0
Average Unique Start Sites per GSP2	35.23	90.19	28.34
Average Unique Start Sites per GSP2 Control	35.67	184.25	30.0

QC Statistics

Avg. Unique DNA And Ambiguous Start Sites Per GSP2	Avg. Unique RNA Start Sites Per GSP2 Control	
63.18	184.25	

Miscellaneous Statistics

On Target Deduplic	tion Ratio
6.30:1	

DNA/RNA Fragment Lengths

DNA Median Fragment Length	DNA Mean Fragment Length	RNA Median Fragment Length	RNA Mean Fragment Length
138.0	147.6	132.0	140.6

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

△ Fusion expression imbalance

1 Low confidence

 \triangle User-annotated true positive

% Intronic fusion

1. Not enough unique start sites

O Transcriptional readthrough event

☐ Known ensembl paralogue

Fusion: FUS → DDIT3			
Filters: ☑ ⊚	Reads: 220 (19.21%)	<u>Segments</u>	
GSP2: FUS_chr16_31195655_27_+_A1_GSP2 Mutation Classification: Undefined	Start Sites: 71	chr16:31195530→31195717 FUS(+) NM_004960.3, exon:5 chr12:57911536→57911489 DDIT3(-) NM_004083.5, exon:2	
Is Artifact: no		223()30 1000.0, 0.012	