

Report for dataset 5prime_v4_GEX_hPBMCs_20k_Rep1-Cell6 (all samples)

Dataset a941bc6d-dc66-473a-8d34-a05a23126997

Summary

Name	5prime_v4_GEX_hPBMCs_20k_Rep1-Cell6 (all samples)
Created At	2025-04-13 21:07:17.600
Number of Records	9,408,444
Total Length	144,234,982,827
Movie Name	m84039_250412_064818_s2
ICS Version	13.3.0.253824
Well Sample	5prime_v4_GEX_hPBMCs_20k_Rep1
Biological Sample	5prime_v4_GEX_hPBMCs_20k_Rep1
Barcode Name	bcM0004--bcM0004

Barcodes

Summary

Unique Barcodes	1
Barcoded HiFi Reads	9,408,444
Unbarcoded HiFi Reads	17,508
Barcoded HiFi Reads (%)	99.81 %
Barcoded HiFi yield (Gb)	144.23 Gb
Unbarcoded HiFi yield (Gb)	0.19 Gb
Barcoded HiFi Yield (%)	99.87 %
Mean HiFi Reads per Barcode	9,408,444
Max. HiFi Reads per Barcode	9,408,444
Min. HiFi Reads per Barcode	9,408,444
Barcoded HiFi read length (mean, kb)	15.33 kb
Unbarcoded HiFi read length (mean, kb)	11.10 kb

Barcode Data

Sample Name	Barcode	Barcode Quality	HiFi Reads	HiFi Read Length (mean, bp)	HiFi Read Quality (median, QV)	HiFi Yield (bp)	Polymerase Read Length (mean, bp)
5prime_v4_GEX_hPBMCs_20k_Rep1	bcM0004--bcM0004	97.5	9408444	15330	Q29	144234982827	144351
No Name	Not Barcoded	0.0	17508	11099	Q27	194335015	114155

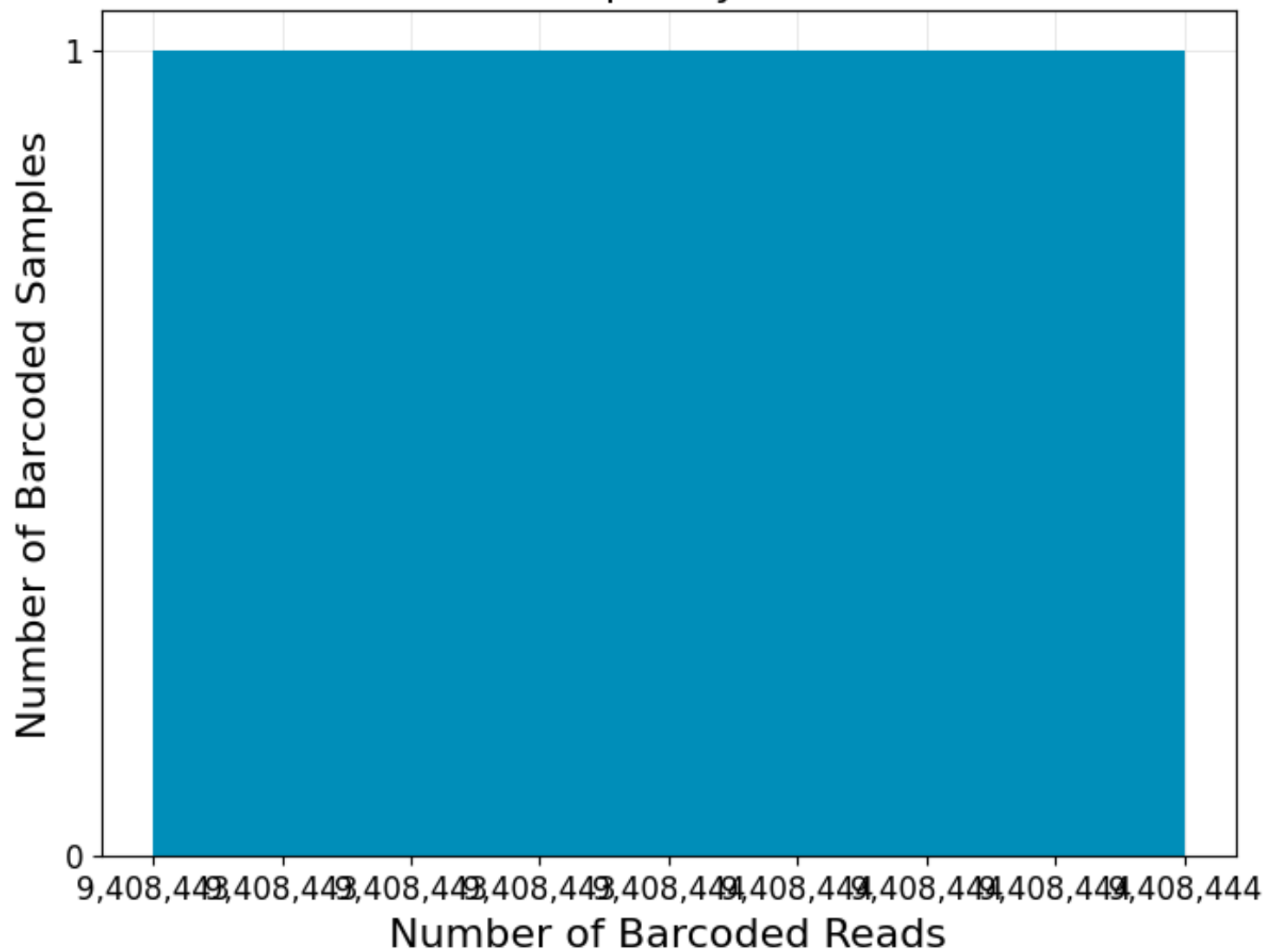
Number Of Reads Per Barcode



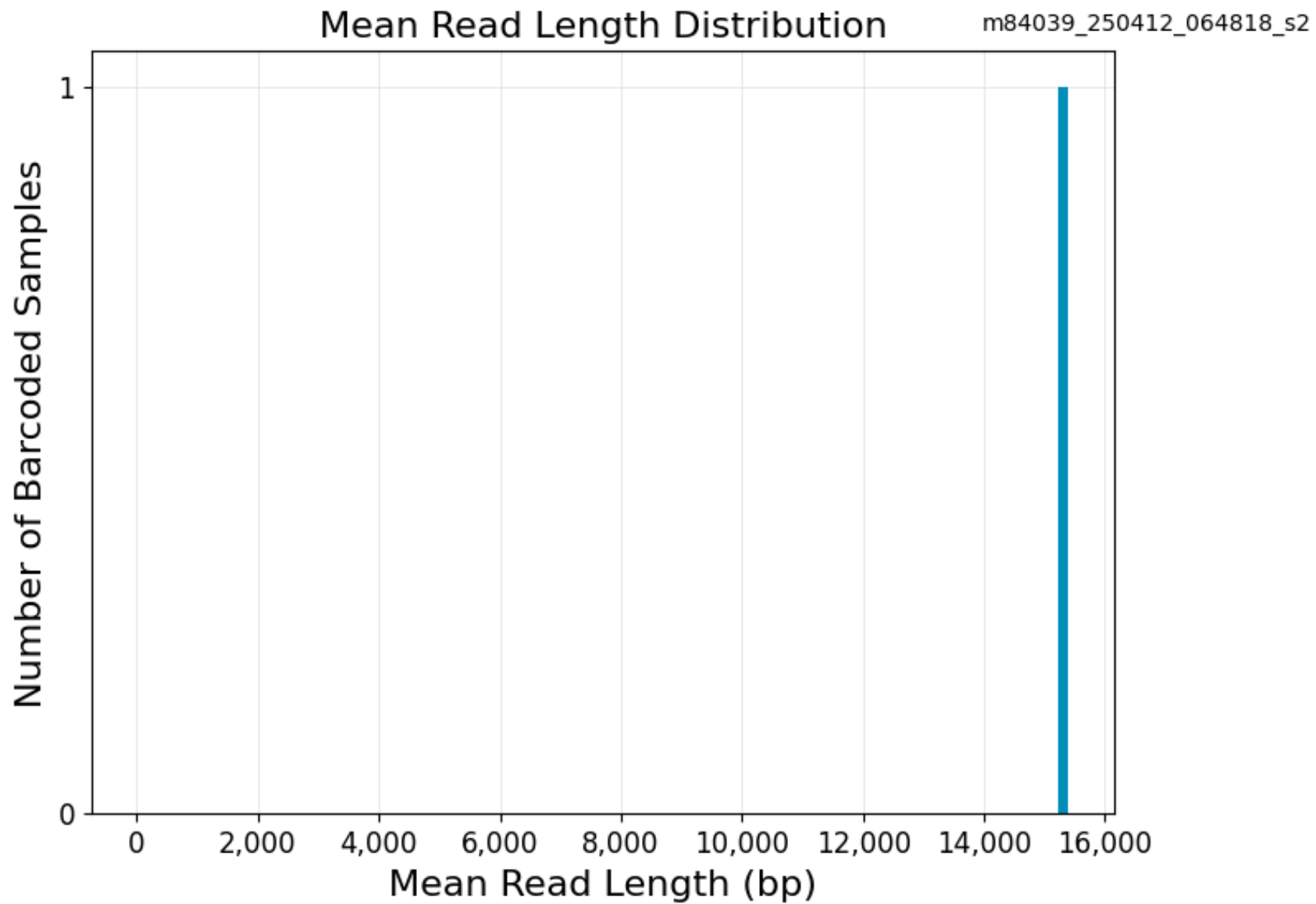
Barcode Frequency Distribution

Barcode Frequency Distribution

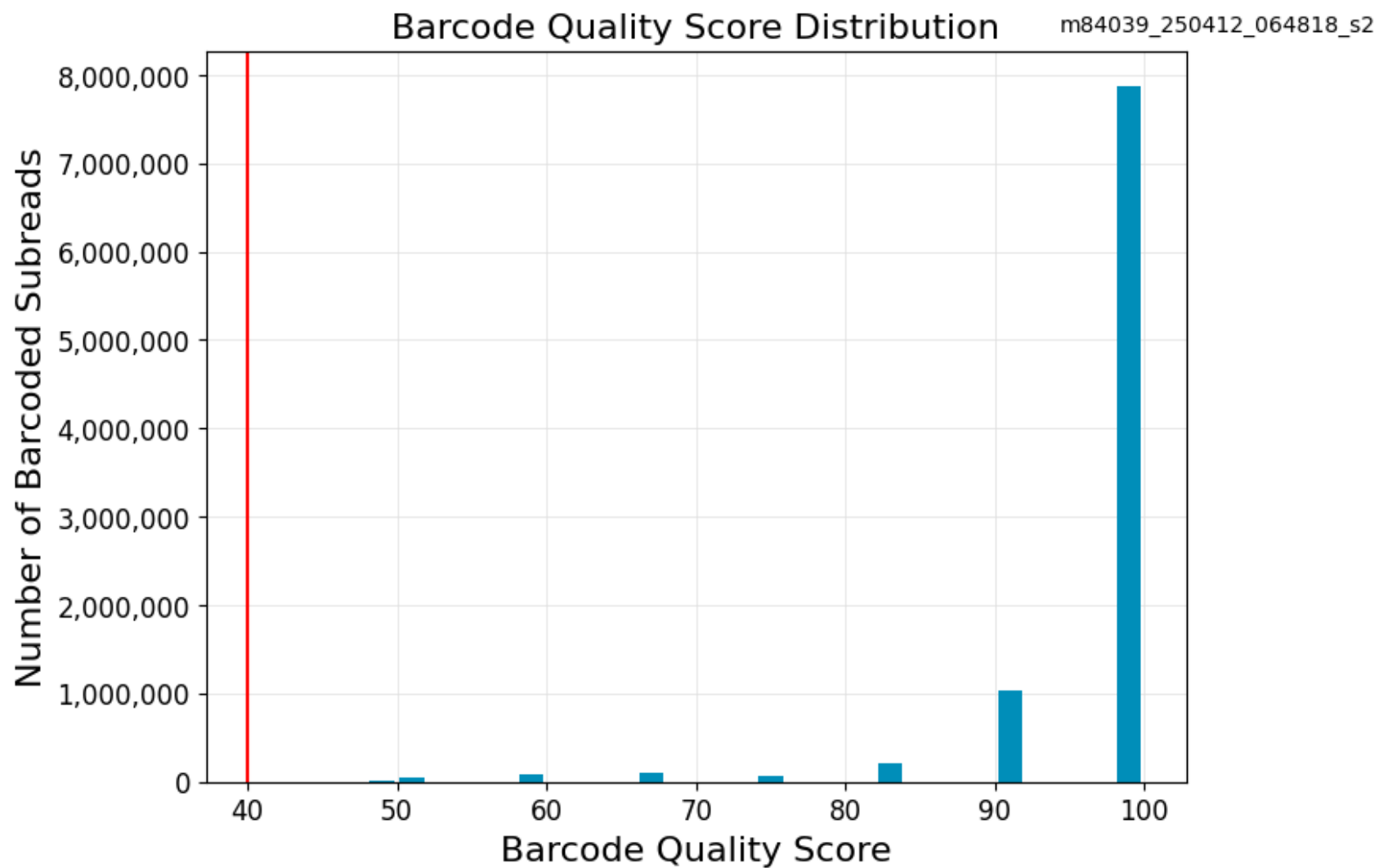
m84039_250412_064818_s2



Mean Read Length Distribution



Barcode Quality Score Distribution



CCS Analysis Report

Summary

HiFi reads	9.4 M
HiFi reads yield	144.43 Gb
HiFi reads length (mean)	15.32 kb
HiFi reads length (median, bp)	15,518
HiFi Read Length N50 (bp)	15,851
HiFi Read Quality (median)	Q29
HiFi Read Quality (median)	29
Base Quality Q30 (%)	95.59%
HiFi Number of Passes (mean)	9
Missing adapters (%)	4.65%

HiFi Read Length Summary

Read Length (kb)	Reads	Reads (%)	Yield (Gb)	Yield (%)
0	9425952	100	144.43 Gb	100
5,000	9222606	98	143.72 Gb	100
10,000	9015093	96	142.21 Gb	98
15,000	5649437	60	95.86 Gb	66
20,000	258889	3	5.46 Gb	4
25,000	2572	0	0.07 Gb	0
30,000	367	0	0.01 Gb	0
35,000	31	0	0.00 Gb	0
40,000	2	0	0.00 Gb	0

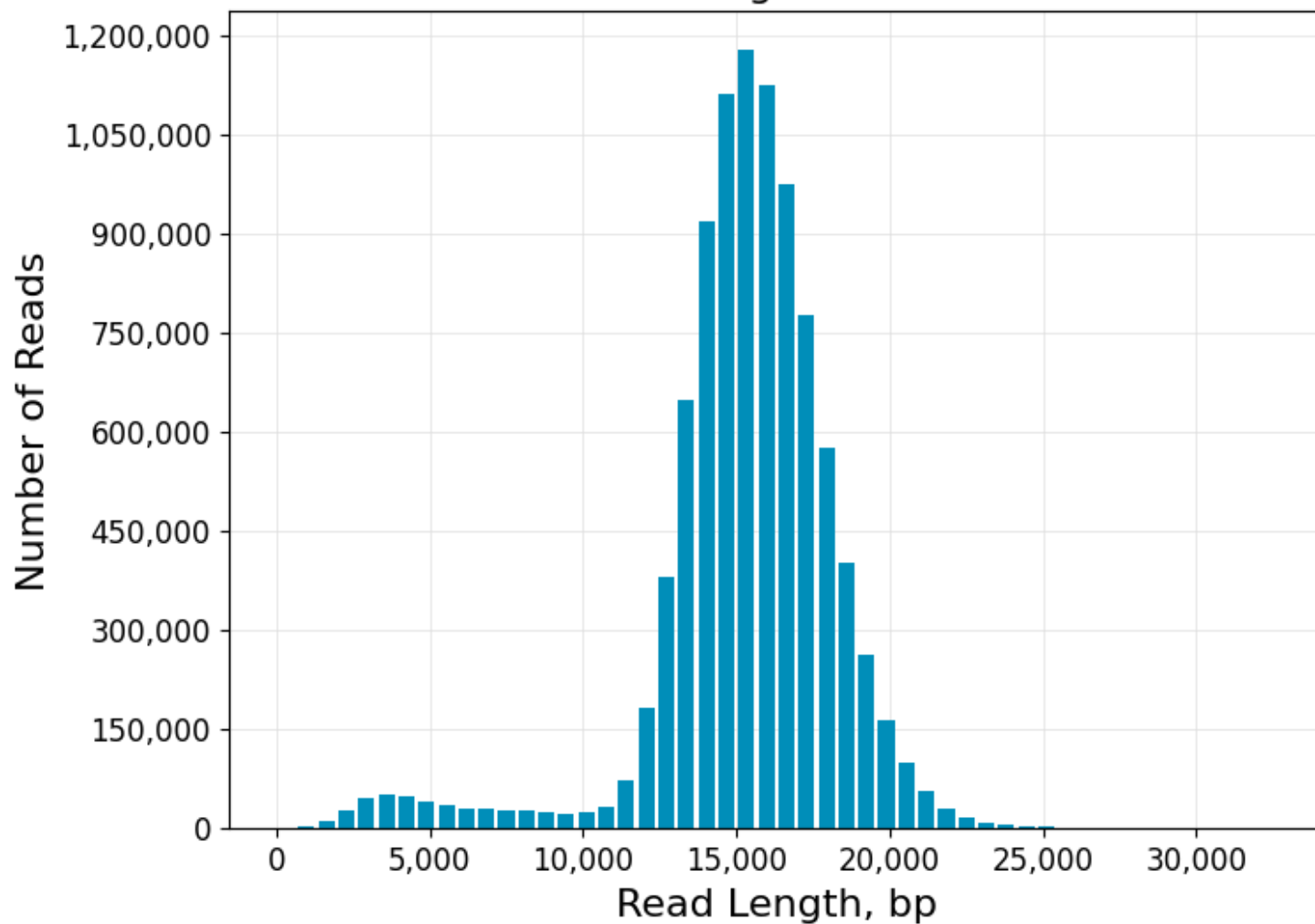
HiFi Read Quality Summary

Read Quality (Phred)	Reads	Reads (%)	Yield (Gb)	Yield (%)
Q20	9425952	100	144.43 Gb	100
Q30	3751296	40	55.80 Gb	39
Q40	129835	1	0.70 Gb	0
Q50	31778	0	0.10 Gb	0

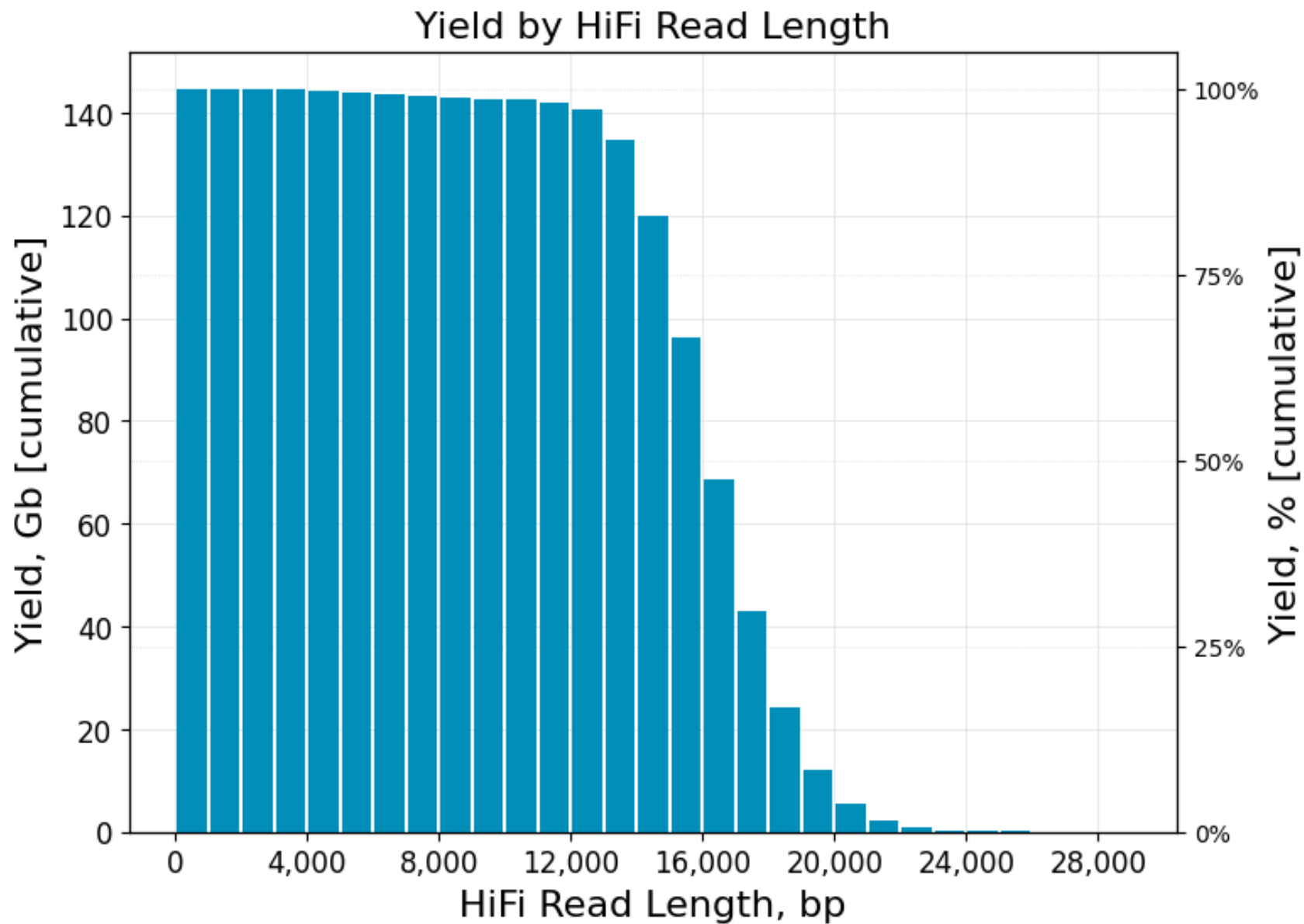
HiFi read length distribution

HiFi read length distribution

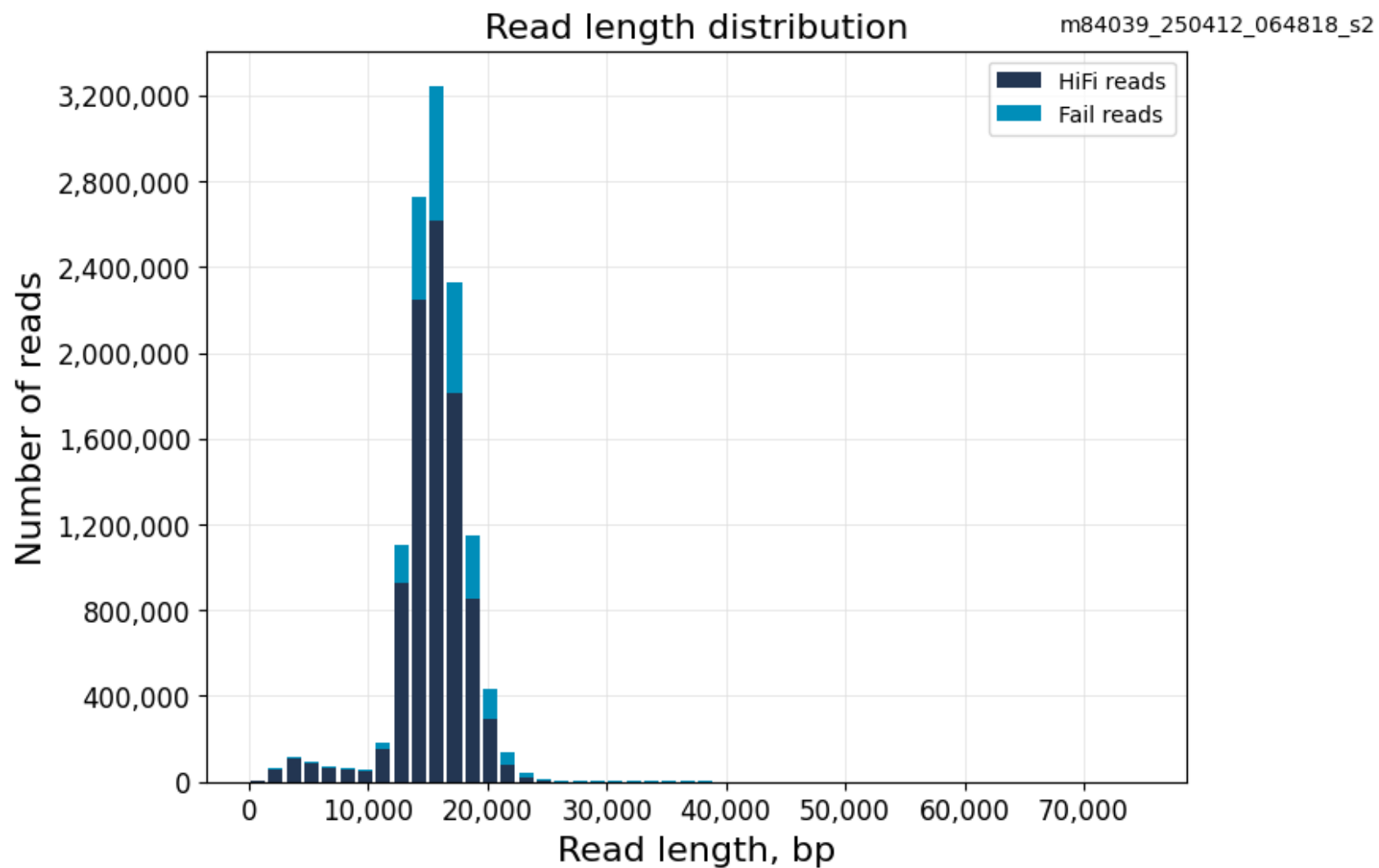
m84039_250412_064818_s2



Yield by HiFi Read Length



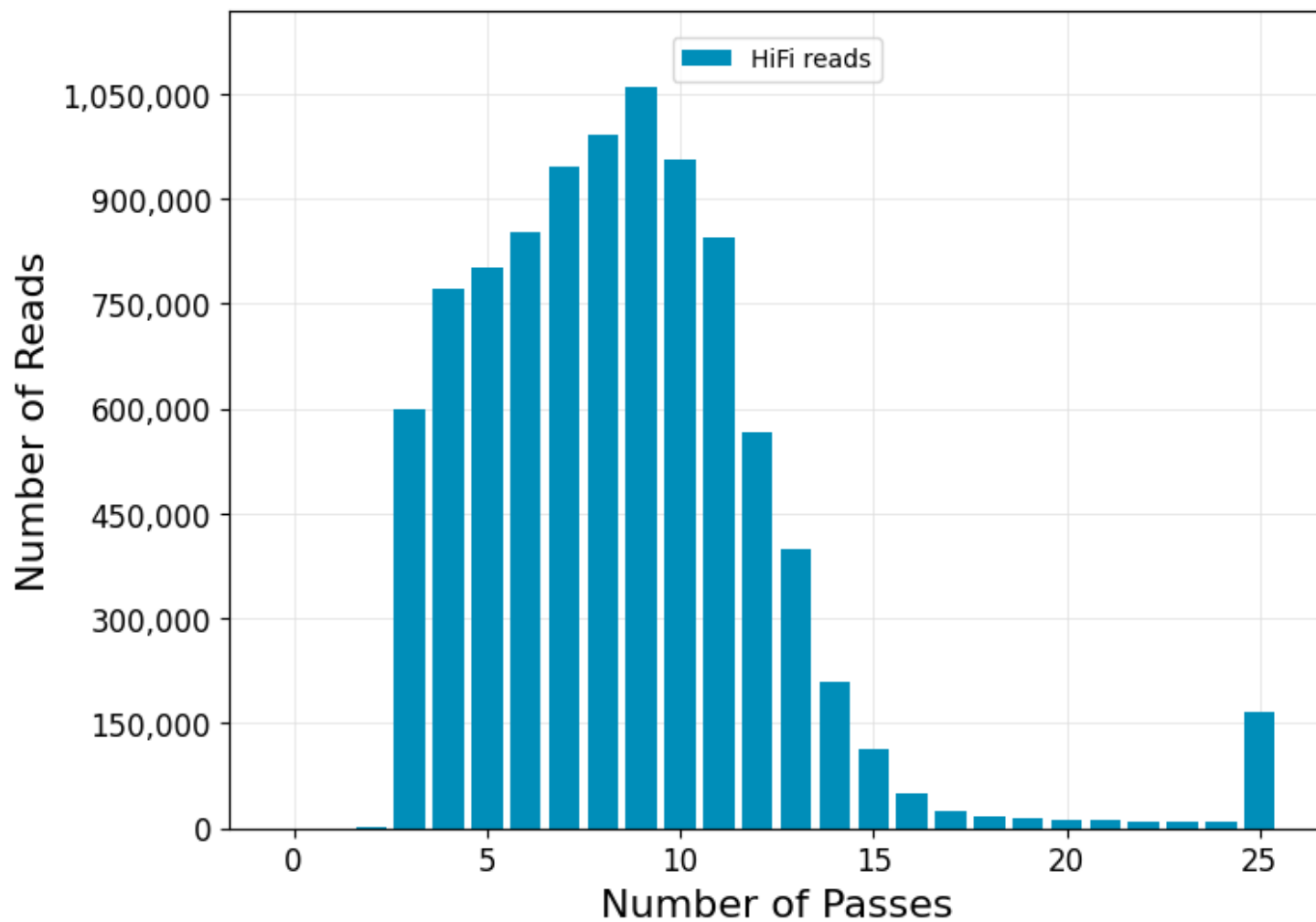
Read length distribution



Number of Passes

Number of Passes

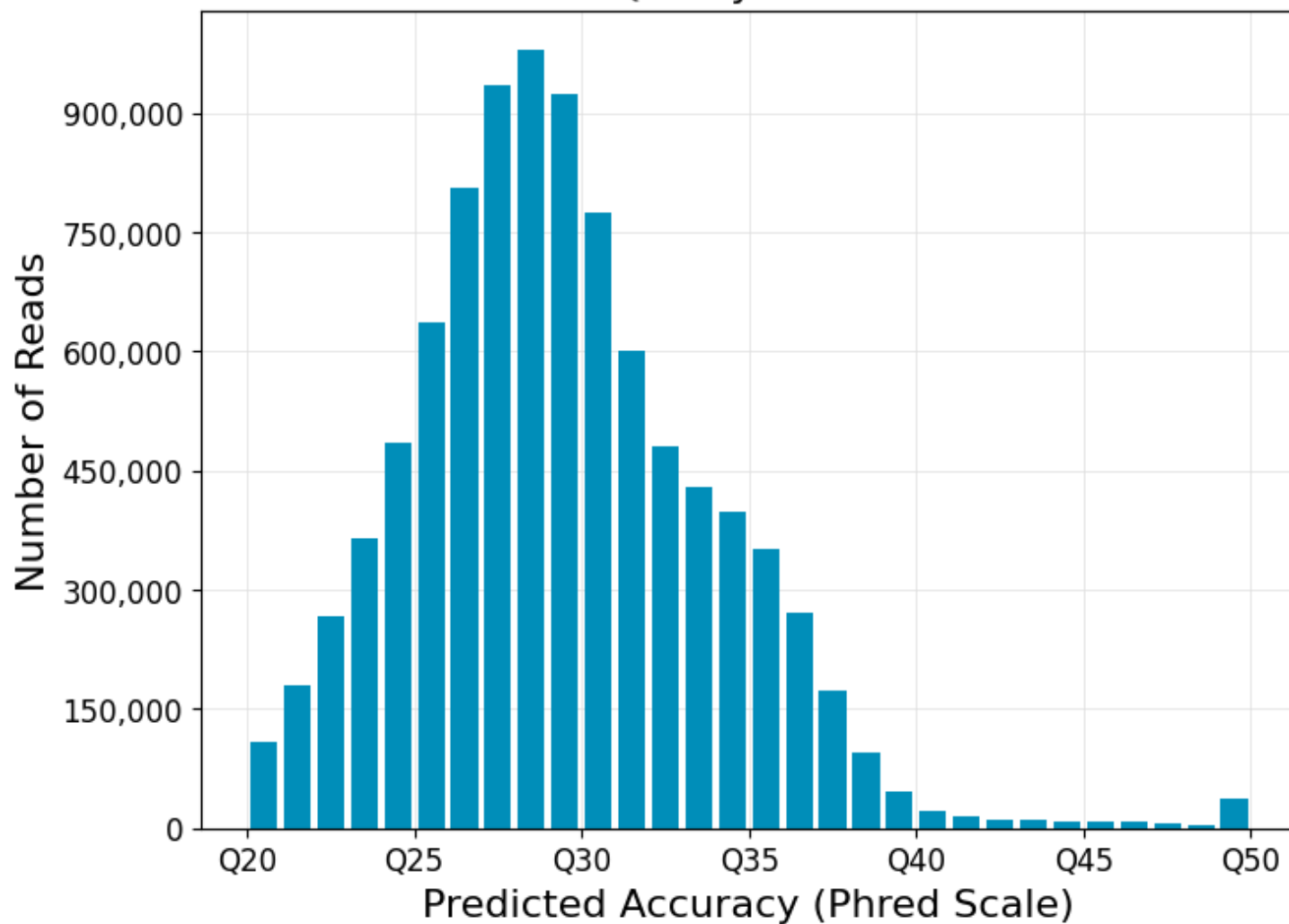
m84039_250412_064818_s2



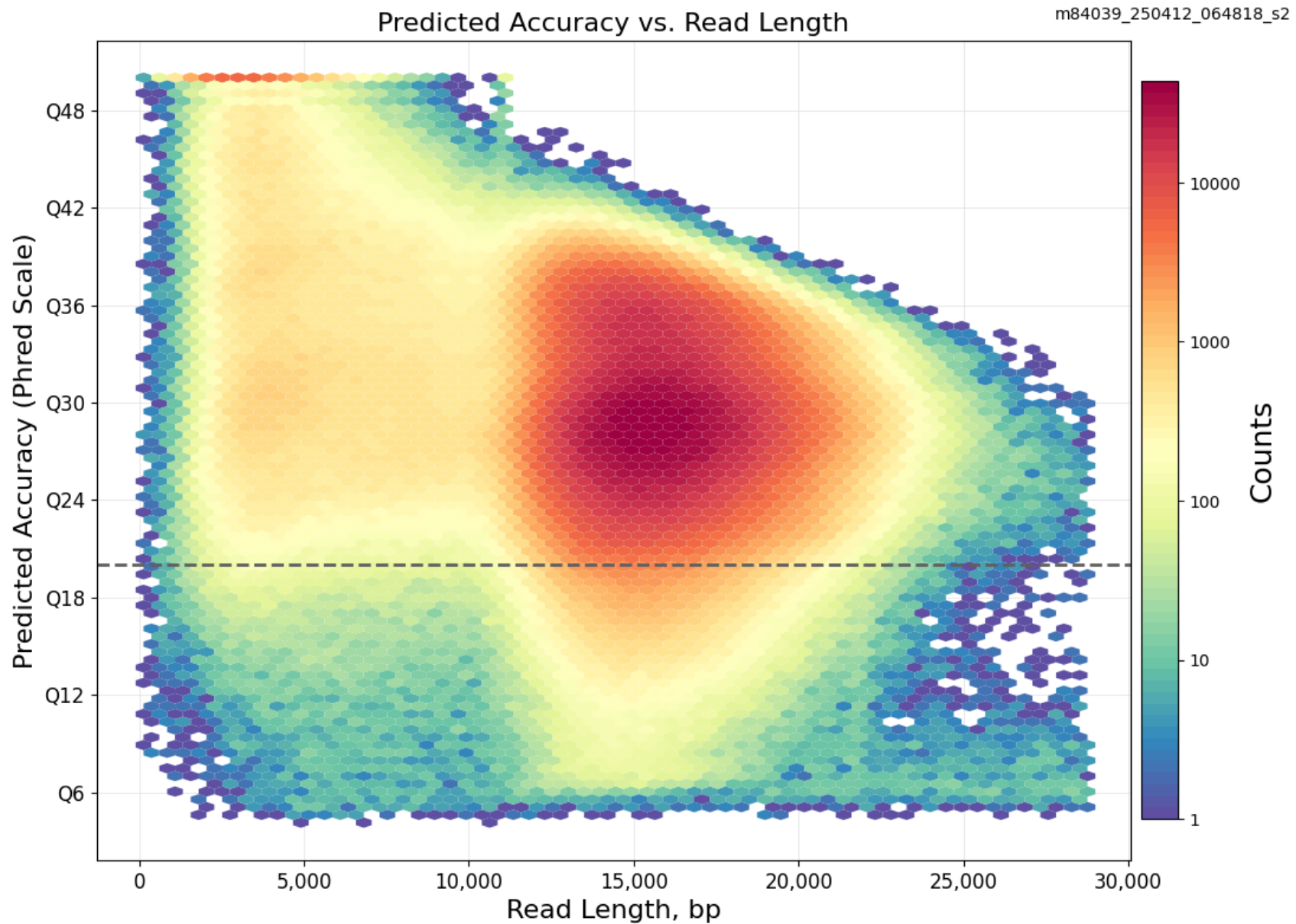
Read Quality Distribution

Read Quality Distribution

m84039_250412_064818_s2



Predicted Accuracy vs. Read Length



Loading Report

Summary

Productive ZMWs	25,165,824
Productivity 0	8,020,215
Productivity 1	17,066,377
Productivity 2	79,232

Control Report

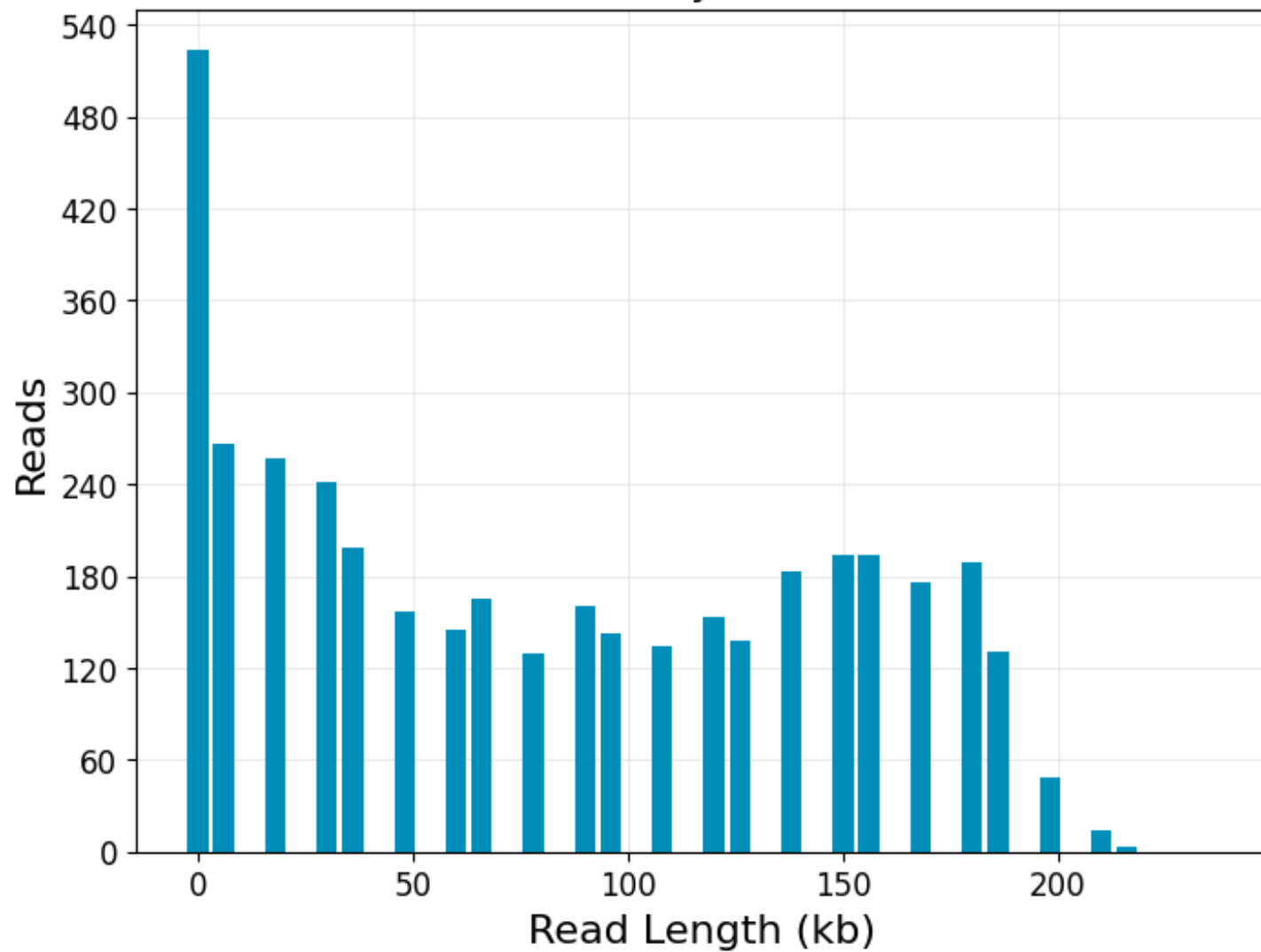
Summary

Number of Control Reads	3,943
Control Read Length Mean	83,828
Control Read Concordance Mean	0.92
Control Read Concordance Mode	0.93

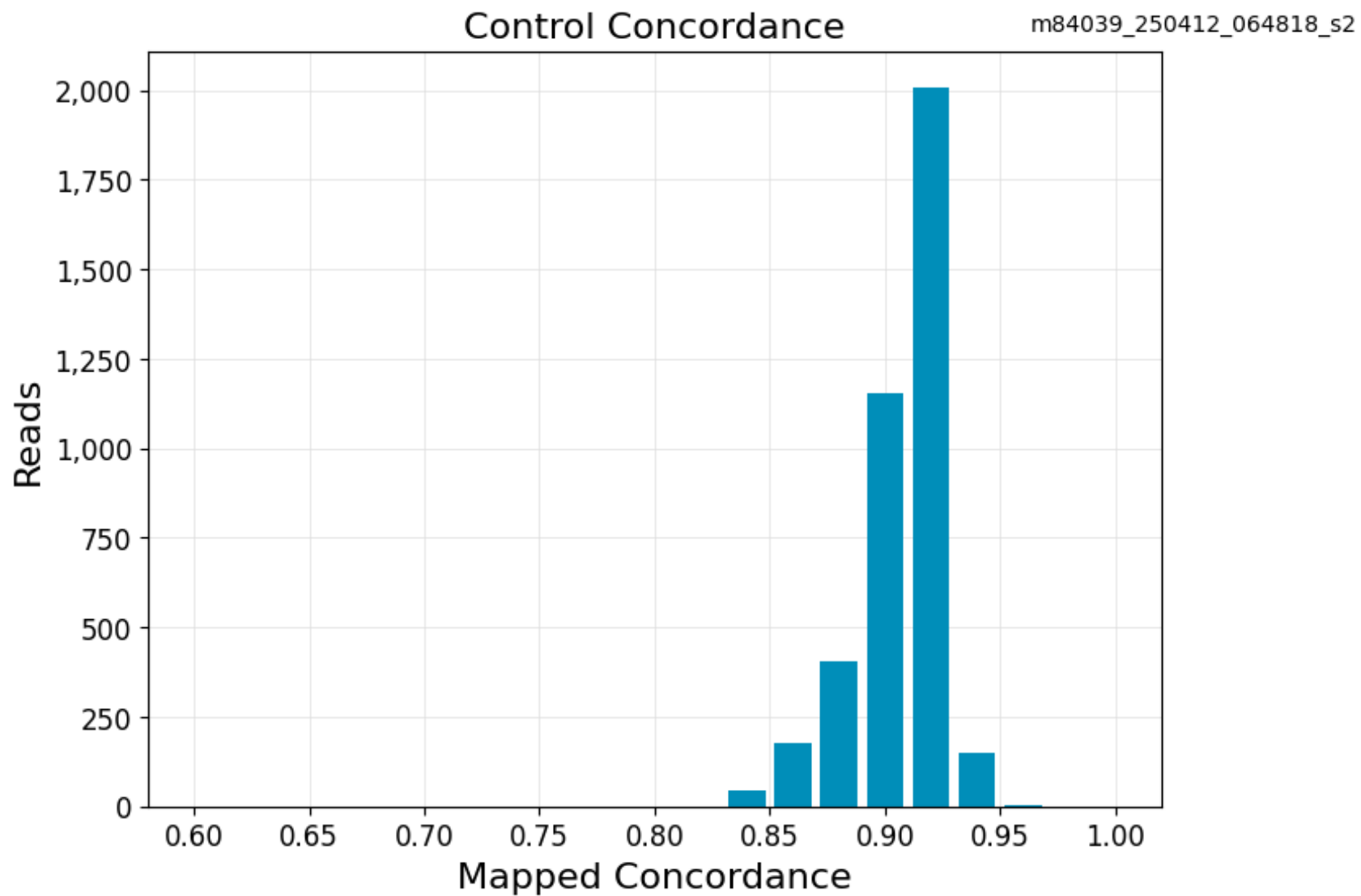
Control Polymerase RL

Control Polymerase RL

m84039_250412_064818_s2



Control Concordance



Raw Data Report

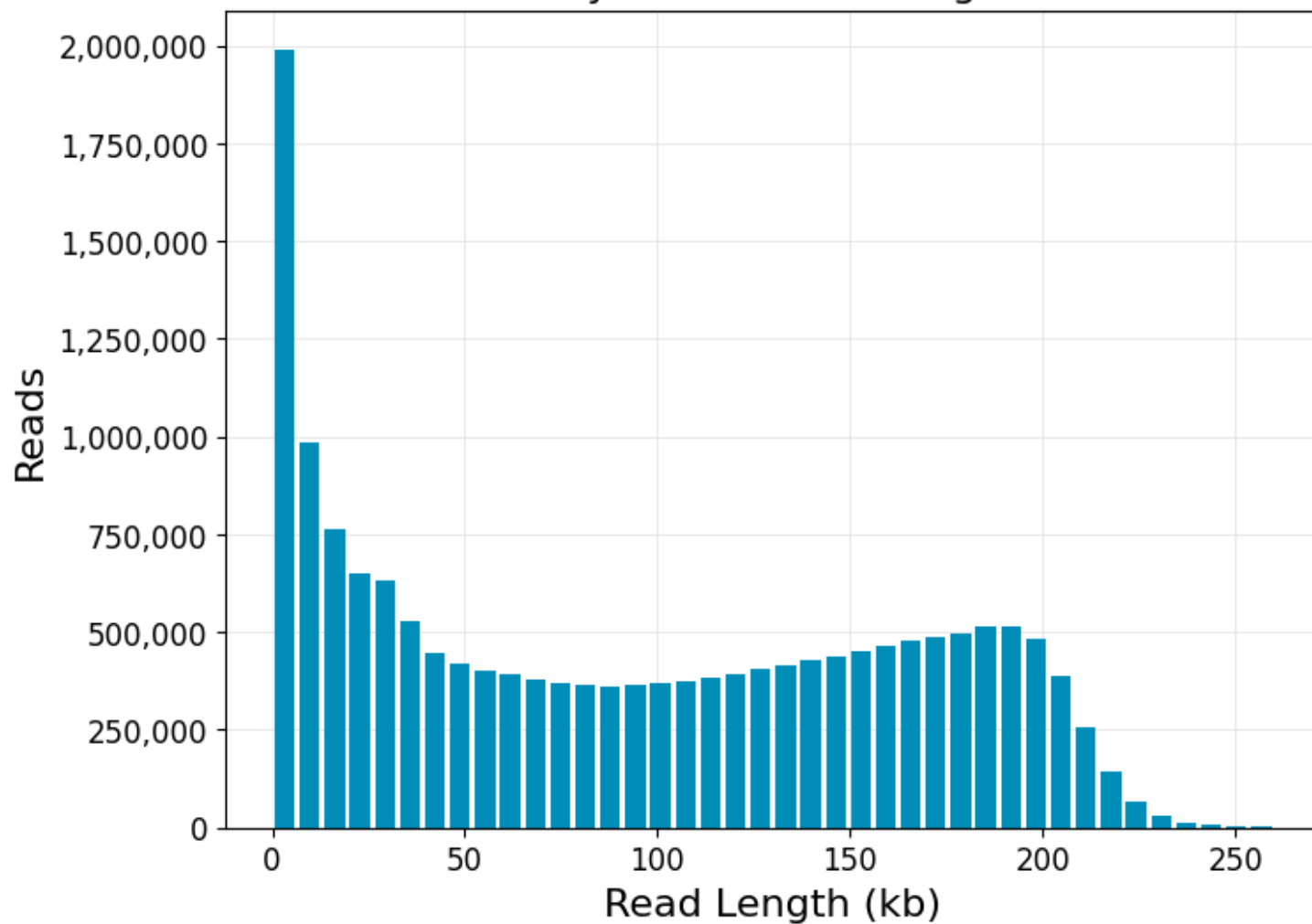
Summary

Polymerase Read Bases	1,539,197,052,410
Polymerase Reads	17,062,434
Polymerase Read Length (mean)	90.21 kb
Polymerase read length (N50)	157.25 kb
Polymerase read length longest subread length (mean)	16.03 kb
Polymerase read length longest subread length (N50)	17.25 kb
Unique Molecular Yield	261,982,945,280
Local Base Rate	2.49

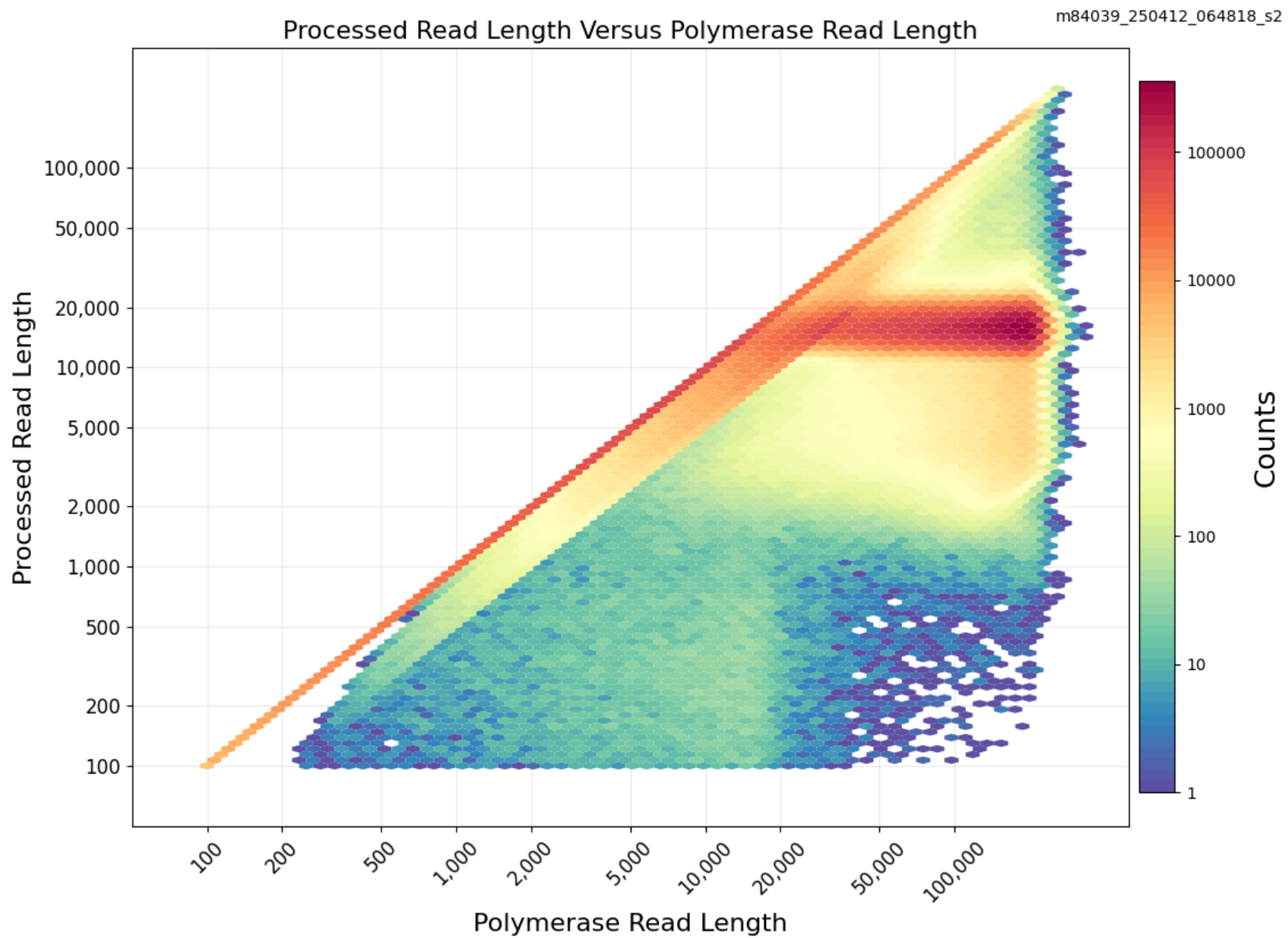
Polymerase Read Length

Polymerase Read Length

m84039_250412_064818_s2



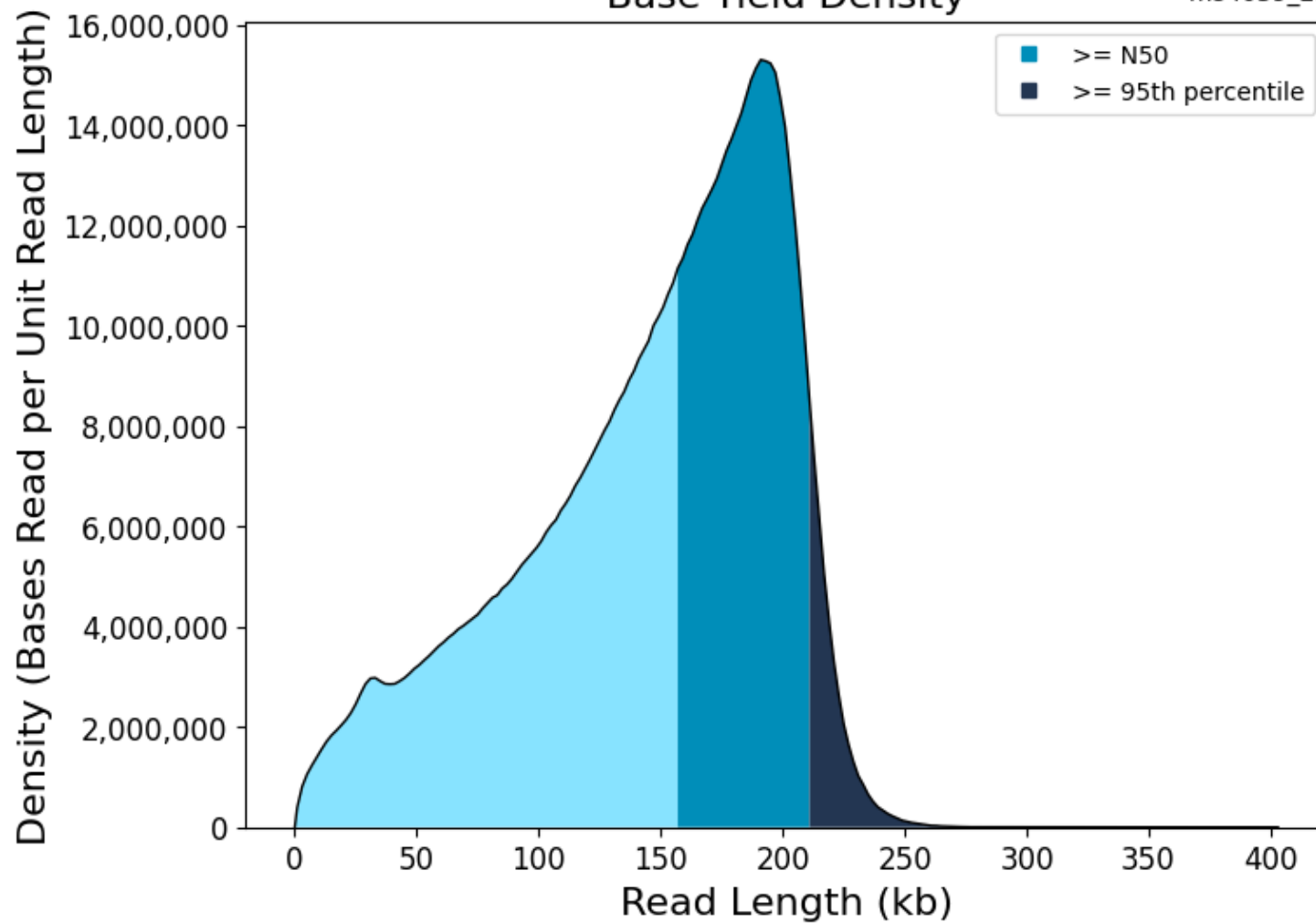
Longest Subread Length Versus Polymerase Read Length



Base Yield Density

Base Yield Density

m84039_250412_064818_s2



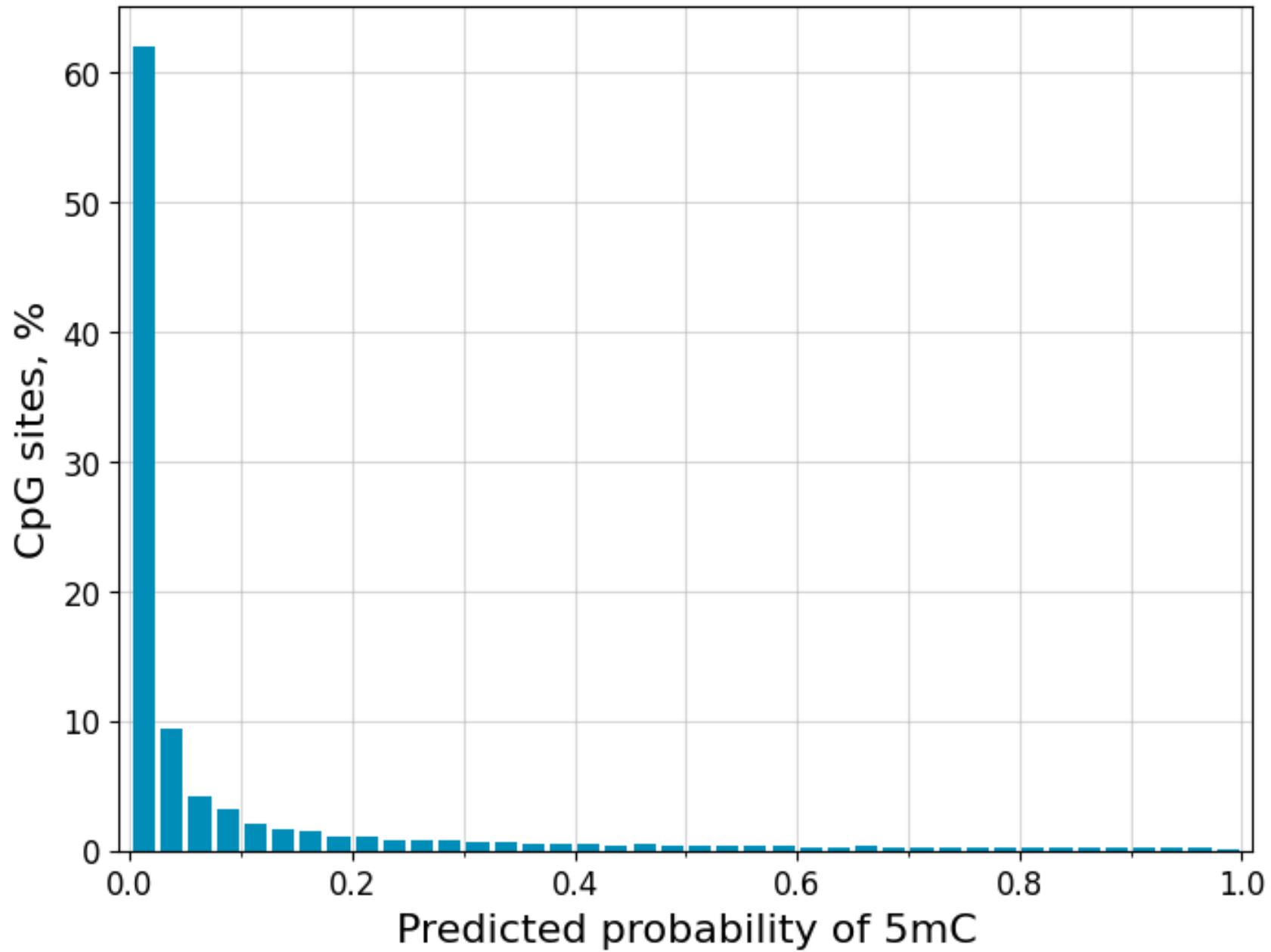
Methylation

Summary

Modification	Motif	Scored sites	Modified sites (Pr > 0.5)
5mC	CpG	98.6%	5.5%
6mA	A	0.5%	0.5%

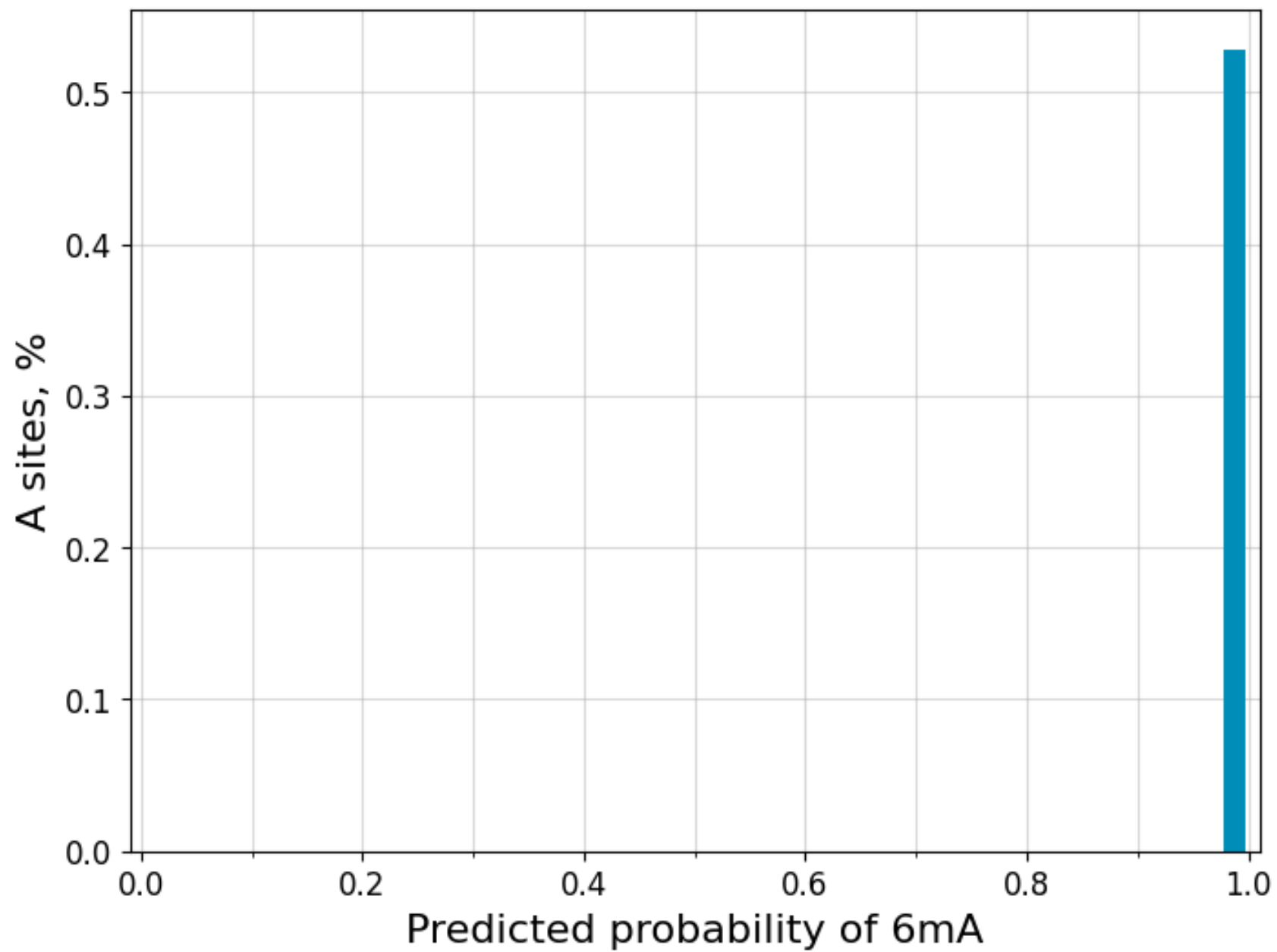
5mC methylation at CpG in reads

5mC at CpG



6mA methylation in reads

6mA



No Sample Setup found

No instrument run found

Parent jobs (1)

Job 118845

Summary

Job Type	import-dataset
Name	import-dataset
Comments	Description for job Import PacBio DataSet
Created At	2025-04-24 15:59:19.368
SMRT Link Version	25.3.0.SNAPSHOT269086

Child jobs (1)

Job 124627

Summary

Job Type analysis
Pipeline cromwell.workflows.pb_segment_reads_and_sc_isoseq
Name KinnexRelease-10x-5pGEX-20k-Revio2cell-largeMem
Comments Description for job Run Analysis Application
Created At 2025-08-02 13:32:20.076
SMRT Link Version 25.3.0.273777

Other inputs

File Type	Name	Dataset ID	Number of Records
ReferenceSet	Human Genome hg38, with Gencode v39 annotations	ba3866bf-2aba-7c99-0570-0d6709174e4a	195
BarcodeSet	MAS-Seq Adapter v1 (MAS16)	10ce52cd-91c5-4bb9-4cac-2233832ecb12	17
BarcodeSet	10X Chromium single cell 5' cDNA primers	5d9656f9-bbcc-6eca-1a46-56a2d31db49e	2
ConsensusReadSet	5prime_v4_GEX_hPBMCs_20k_Rep1-Cell7 (all samples)	7aefee39-0ce4-4640-b0fe-f554dc69a979	9,358,946

Output files

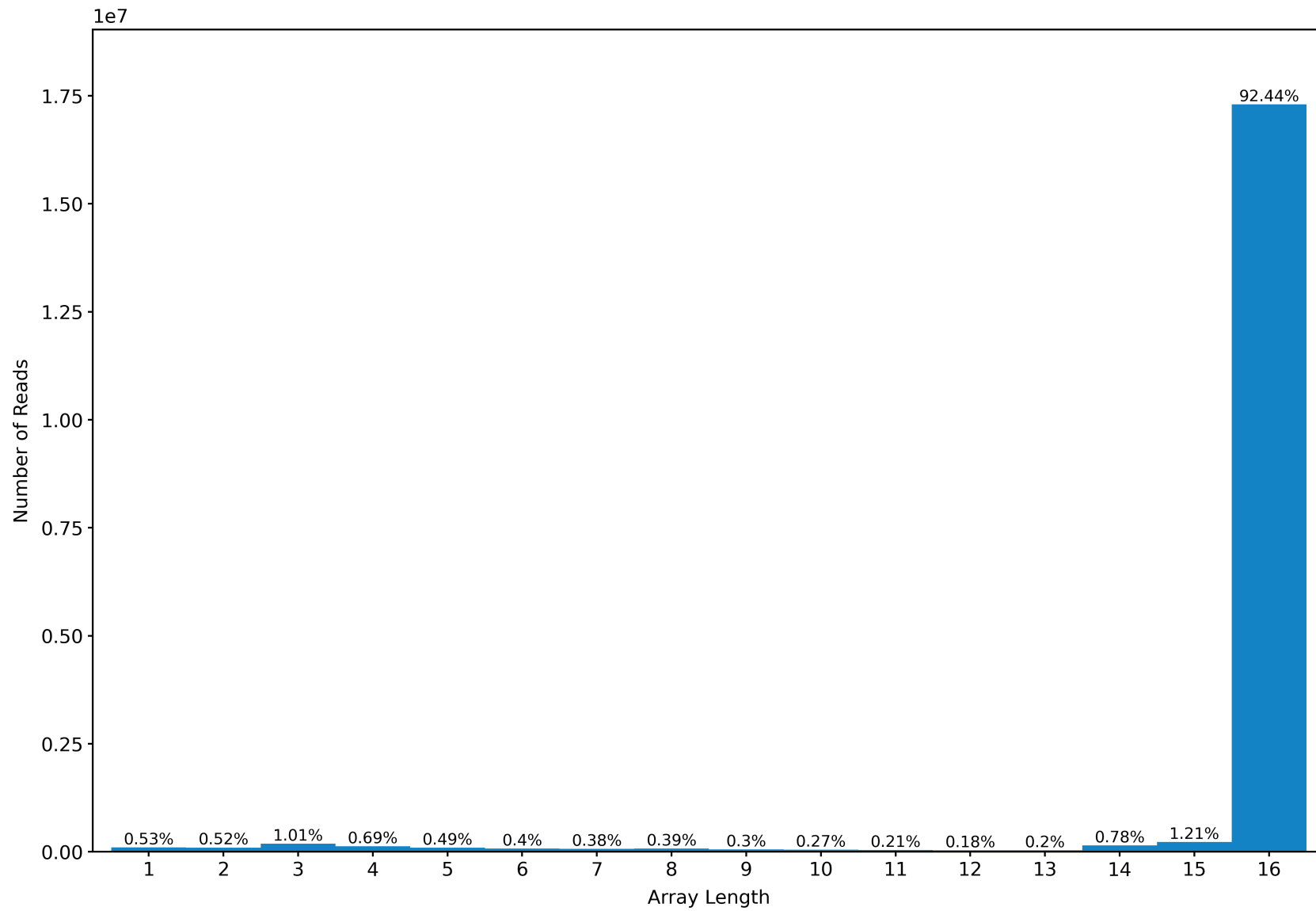
File name	Size	Type
KinnexRelease-10x-5pGEX-20k-Revio2cell-largeMem Segmented Reads	47,582	ConsensusReadSet
Deduplicated reads after cell barcode correction, unmapped, BAM	58,630,025,312	bam
Single-cell isoform and gene matrix, tar-gzipped	5,229,835,901	tgz
Segmented Reads, passing, unaligned	93,966,703,973	bam
Segmented Reads, passing, unaligned	93,478,605,474	bam
Non-passing reads, unaligned	3,598,645,913	bam
Non-passing reads, unaligned	3,359,211,767	bam
Unique mapped transcripts, GFF	2,866,215,348	gff
Unique mapped transcripts, filtered, GFF	1,241,683,689	gff
Deduplicated reads after cell barcode correction, mapped, BAM	29,712,220,750	bam
Deduplicated reads after cell barcode correction, mapped, BAM index	37,932,976	bam_bai
Unique mapped transcripts, junctions TXT	2,207,733,257	txt
Unique mapped transcripts, filtered, junctions TXT	1,173,175,043	txt
Deduplicated transcripts	42,745,109,308	Fasta
Collapsed transcript groups	2,977,338,014	txt
Unique mapped transcripts, classification TXT	1,972,267,425	txt
Unique mapped transcripts, filtered, classification TXT	704,402,006	txt
bcstats_report.tsv.gz	18,086,654	tsv
Analysis Log	101,851	log
3M_5pgex_jan_2023.txt.gz	6,945,252	txt
3M_february_2018_REVERSE_COMPLEMENTED.txt.gz	17,812,866	txt
Analysis Log	4,083	log
SMRT Link Log	20,183	log

Read Segmentation

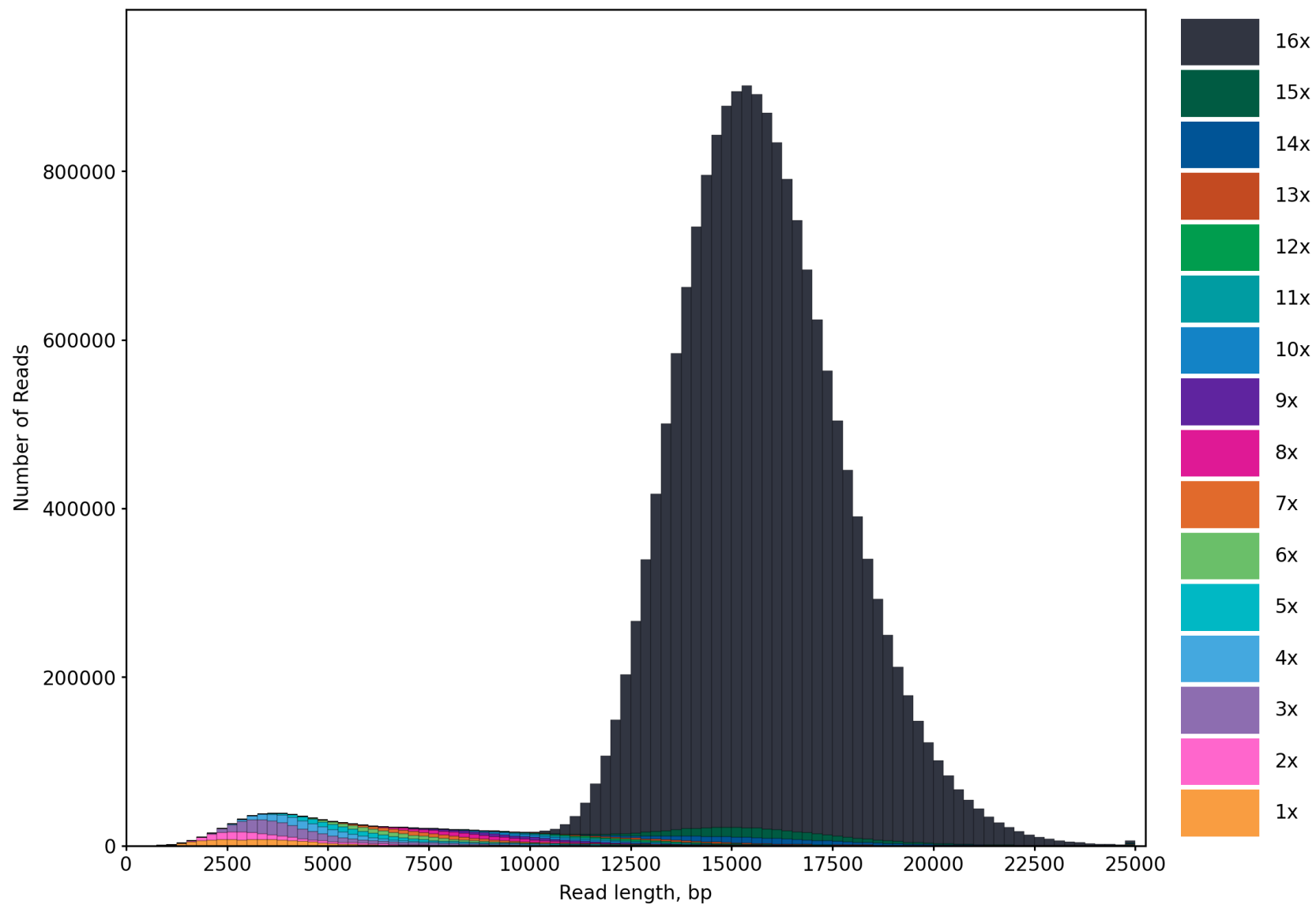
Summary

Reads	18,767,390
Segmented reads (S-reads)	288,007,196
Mean length of S-reads	970.0055384102278
Percent of reads with full arrays	92.20 %
Mean array size (concatenation factor)	15.35

Histogram of concatemers

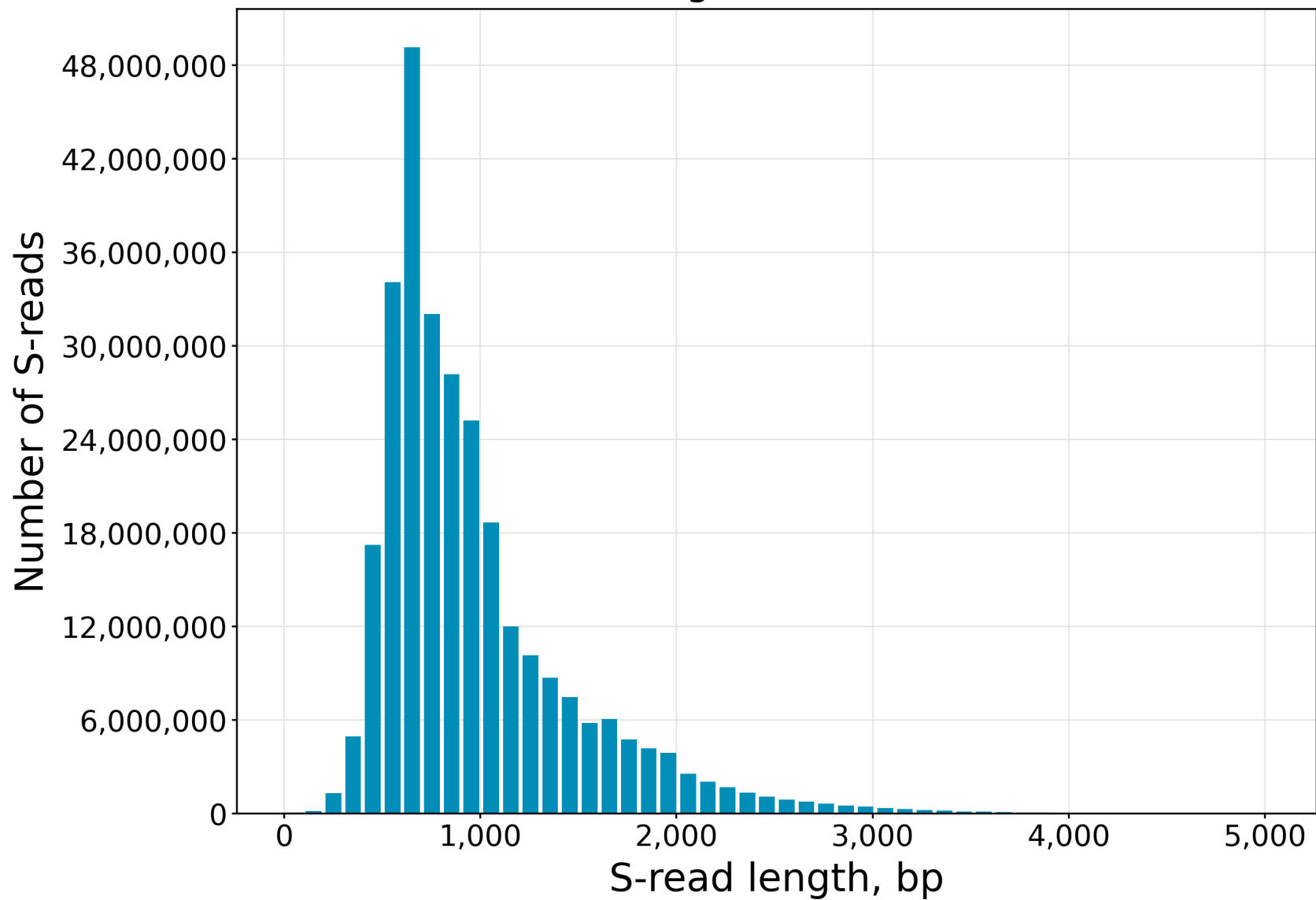


Length of Reads



Length of S-reads

Length of S-read

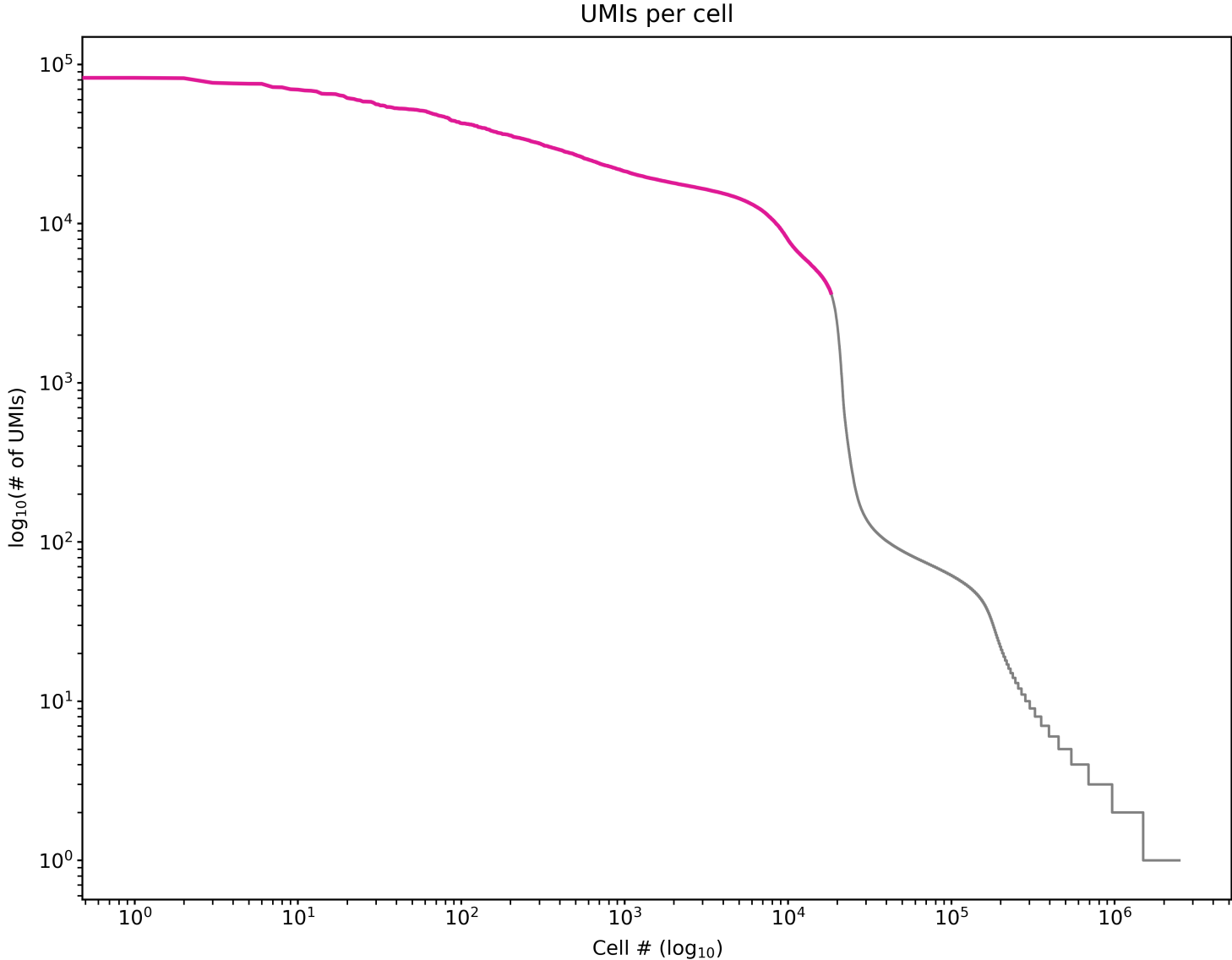


Cell Statistics

Summary

Estimated Number of Cells	17,654
Reads in Cells	87.13%
Mean Reads per Cell	14,011
Median UMIs per Cell	9,524

Barcode Rank Plot



Read Statistics

Summary

Reads	288,007,196
Read Type	SEGMENT
Reads with 5' and 3' Primers with extracted UMIs and Barcodes	285,630,076
Non-Concatamer Reads with 5' and 3' Primers and Poly-A Tail (FLNC reads)	283,891,388
FLNC Reads with Valid Barcodes	271,375,072
FLNC Reads with Valid Barcodes, corrected	280,263,363
Reads after Barcode Correction and UMI Deduplication	197,544,312

Transcript Statistics

Summary

FLNC Reads Mapped Confidently to Genome	191,189,865
FLNC Reads Mapped Confidently to Transcriptome	119,177,062
Total Unique Genes	2,160,147
Total Unique Genes, filtered	135,011
Total Unique Genes, known genes only	39,976
Total Unique Genes, filtered, known genes only	31,071
Total Unique Transcripts	5,637,804
Total Unique Transcripts, filtered	1,384,709
Total Unique Transcripts, known transcripts only	108,978
Total Unique Transcripts, filtered, known transcripts only	99,281

Transcript Summary

Value	Analysis Metric
3,154	Median Genes per Cell
2,407	Median Genes per Cell, known genes only
4,137	Median Transcripts per Cell
1,950	Median Transcripts per Cell, known transcripts only
2,160,147	Total Unique Genes
39,976	Total Unique Genes, known genes only
5,637,804	Total Unique Transcripts
108,978	Total Unique Transcripts, known transcripts only

Transcript Summary, filtered

Value	Analysis Metric
1,772	Median Genes per Cell
1,743	Median Genes per Cell, known genes only
2,255	Median Transcripts per Cell
1,670	Median Transcripts per Cell, known transcripts only
135,011	Total Unique Genes
31,071	Total Unique Genes, known genes only
1,384,709	Total Unique Transcripts
99,281	Total Unique Transcripts, known transcripts only

Transcript Classification

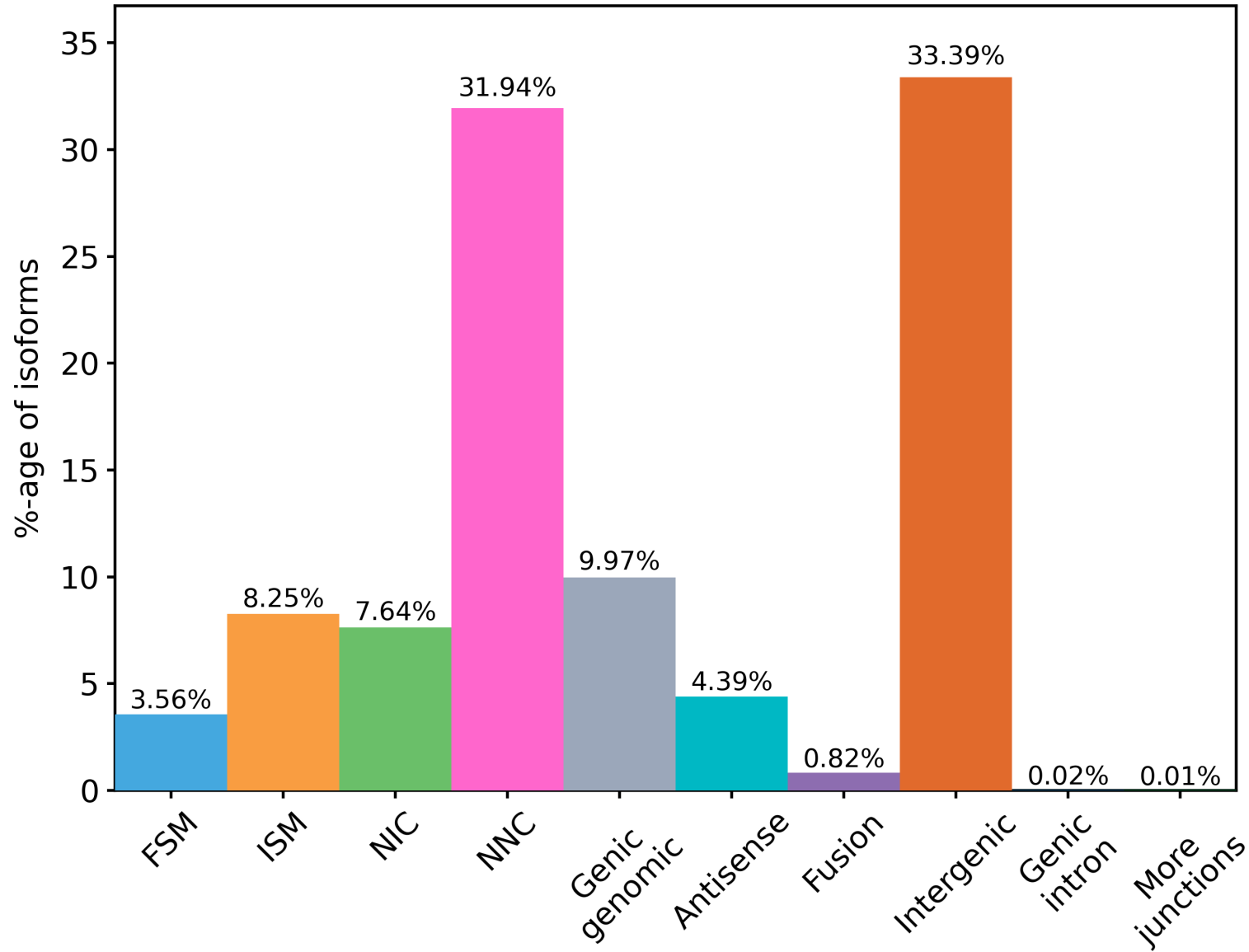
Category	Count	CAGE Detected	CAGE Detected (%)	polyA Motif Detected	polyA Motif Detected (%)
FSM	222891	128967	57.86%	67652	30.35%
ISM	517523	212697	41.10%	156434	30.23%
NIC	478879	308184	64.36%	161479	33.72%
NNC	2002321	1231439	61.50%	757232	37.82%
Antisense	275513	12153	4.41%	82855	30.07%
Fusion	51581	28536	55.32%	20474	39.69%
More junctions	857	418	48.77%	258	30.11%
Genic intron	1099	6	0.55%	362	32.94%
Genic genomic	625205	149505	23.91%	189730	30.35%
Intergenic	2093371	21491	0.999	712321	34.03%

Transcript Classification, filtered

Category	Count	CAGE Detected	CAGE Detected, (%)	polyA Detected	polyA Detected, (%)
FSM	187860	111558	59.38%	67652	36.01%
ISM	356165	161074	45.22%	138556	38.90%
NIC	375188	243279	64.84%	160170	42.69%
NNC	739730	466058	63.00%	351468	47.51%
Antisense	23799	2438	10.24%	14541	61.10%
Fusion	24723	14126	57.14%	12404	50.17%
More junctions	490	252	51.43%	218	44.49%
Genic intron	0	0	0.00%	0	0.00%
Genic genomic	24507	10487	42.79%	12053	49.18%
Intergenic	96991	2584	2.66%	70613	72.80%

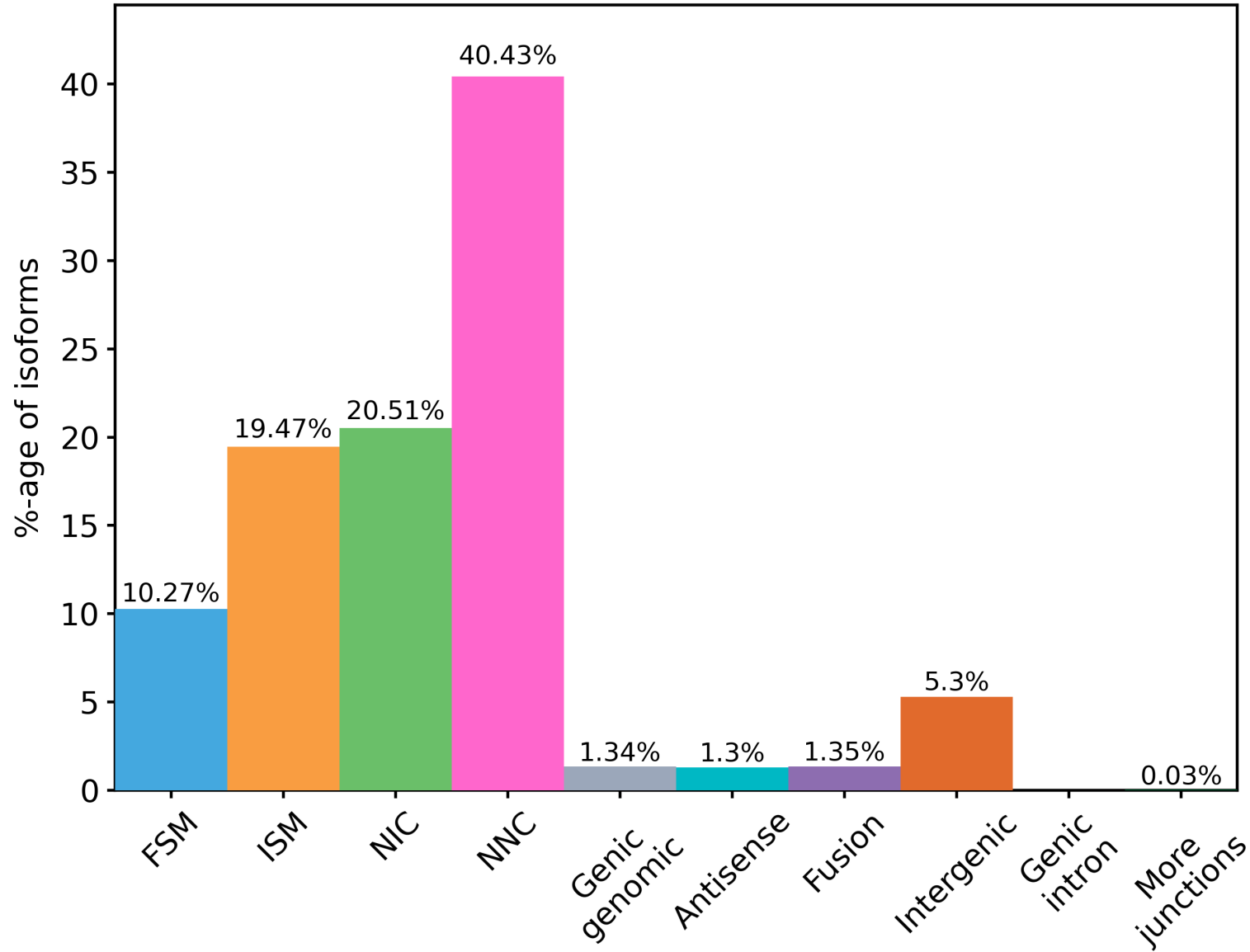
Isoform Distribution Across Categories

Isoform distribution across structural categories

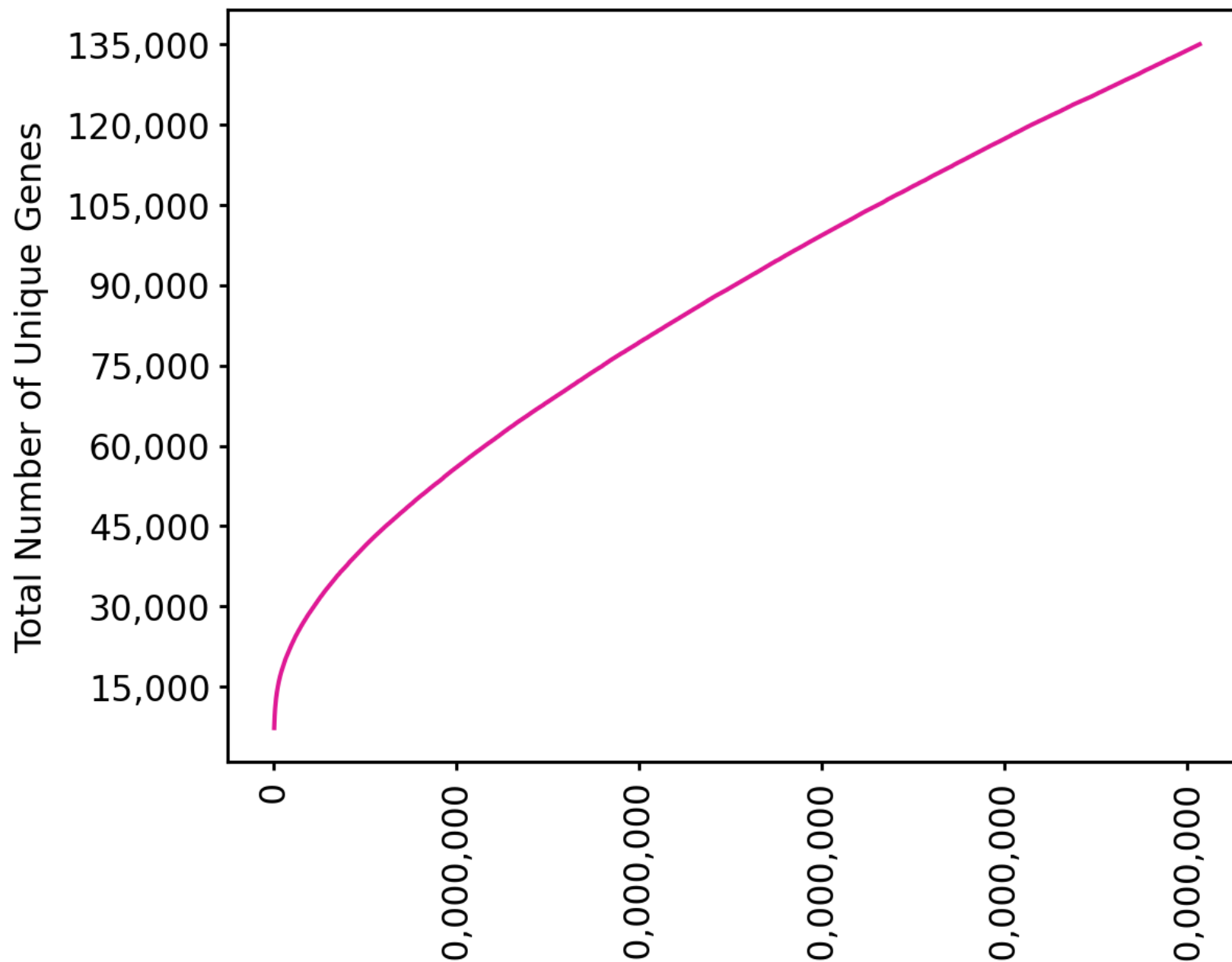


Isoform Distribution Across Categories

Isoform distribution across structural categories



Gene Saturation, all genes, filtered



2

4

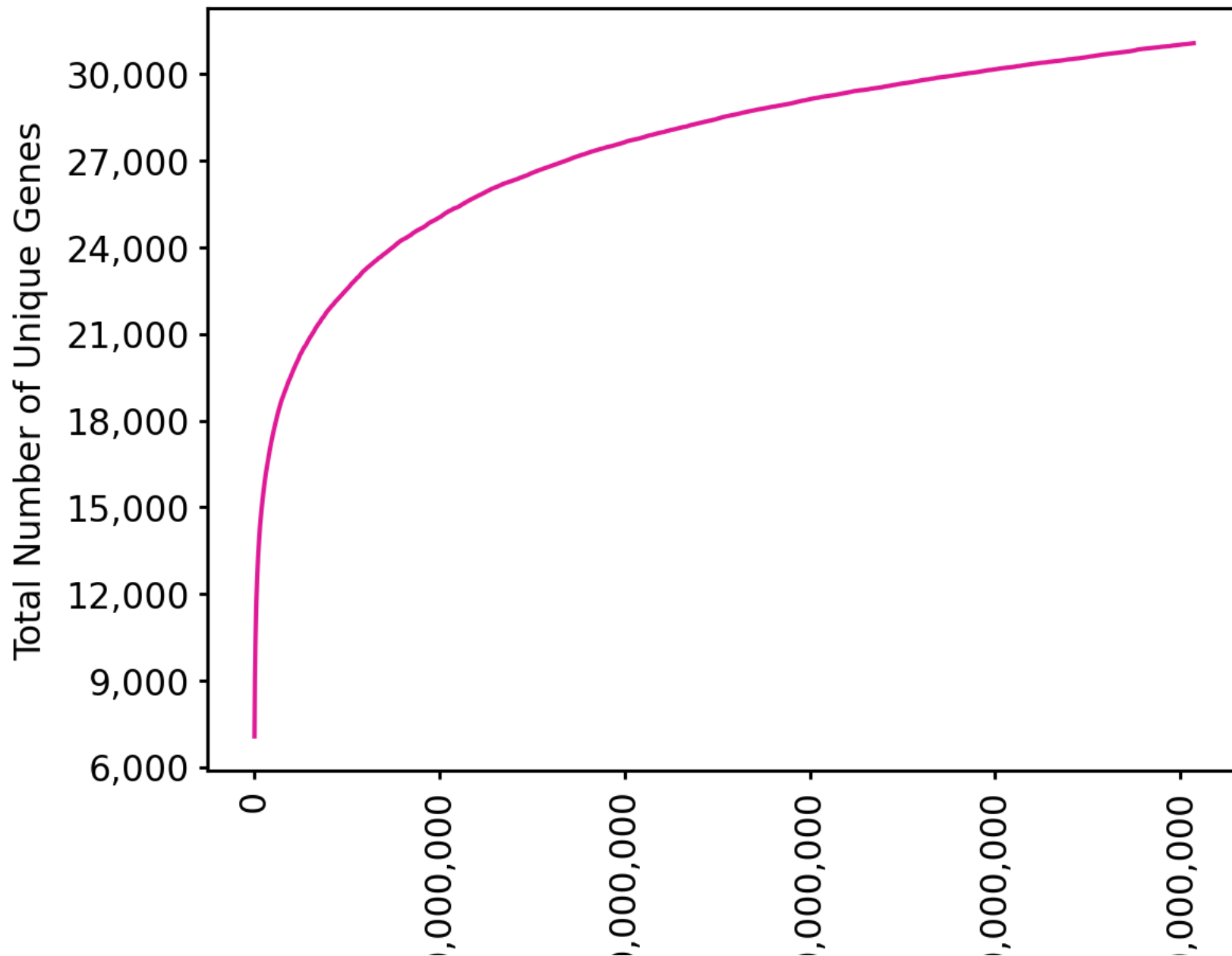
6

8

10

Total Reads

Gene Saturation, known genes only, filtered



20

40

60

80

100

Total Reads

