

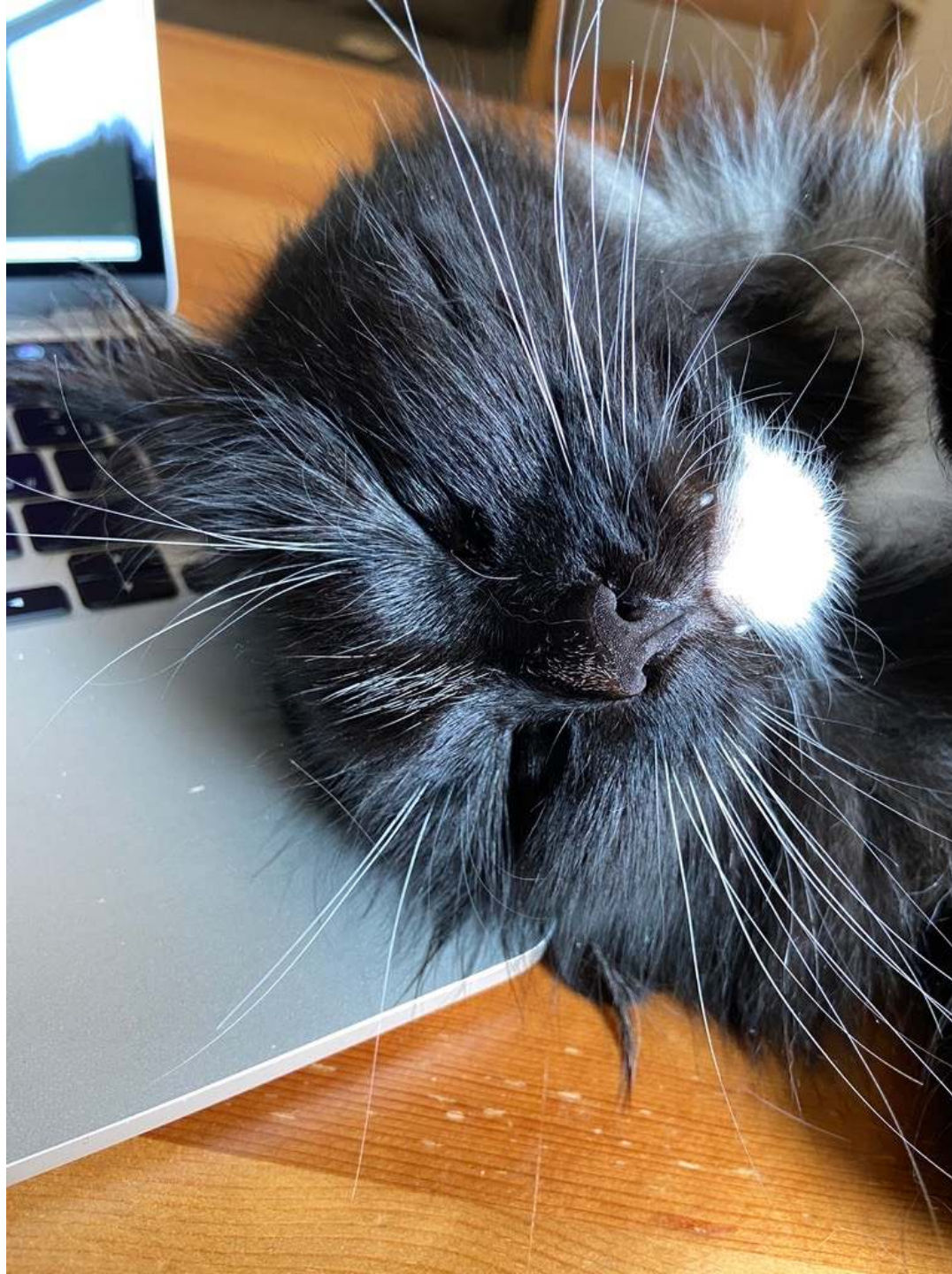
Structural variant activities!!

- 1 SV quiz
- 2 SV calling tutorial

SVs are awesome
and fun

QUIZ!

With prizes ;)



SV classification (aka SV quiz)

You can find all the files necessary to answer the questions in the folder “SV_quiz”. Within this folder you will find a subfolder corresponding to each of the questions. The tools we suggest to use are all freely available (IGV, MAFFT or LAST online) but Censor/Rebase which has a limited free use. Nonetheless, the limits of Censor should allow all of you to answer these questions. If you have any problem with Censor please ask me (Vale) or Alex :)

Visualise BAM files

IGV: Amazon instance or download on your computer ([igvteam/igv](https://github.com/IGVteam/igv))

Make dotplots

MAFFT: <https://mafft.cbrc.jp/alignment/server/index.html>

LAST: <http://lastweb.cbrc.jp>

Repeat database Censor (sequence homology to Repbase repeat database): <https://www.girinst.org/censor/index.php>

Q1: What is it?

Hint: Is there any signature of incongruent read mapping along the sequence of interest?

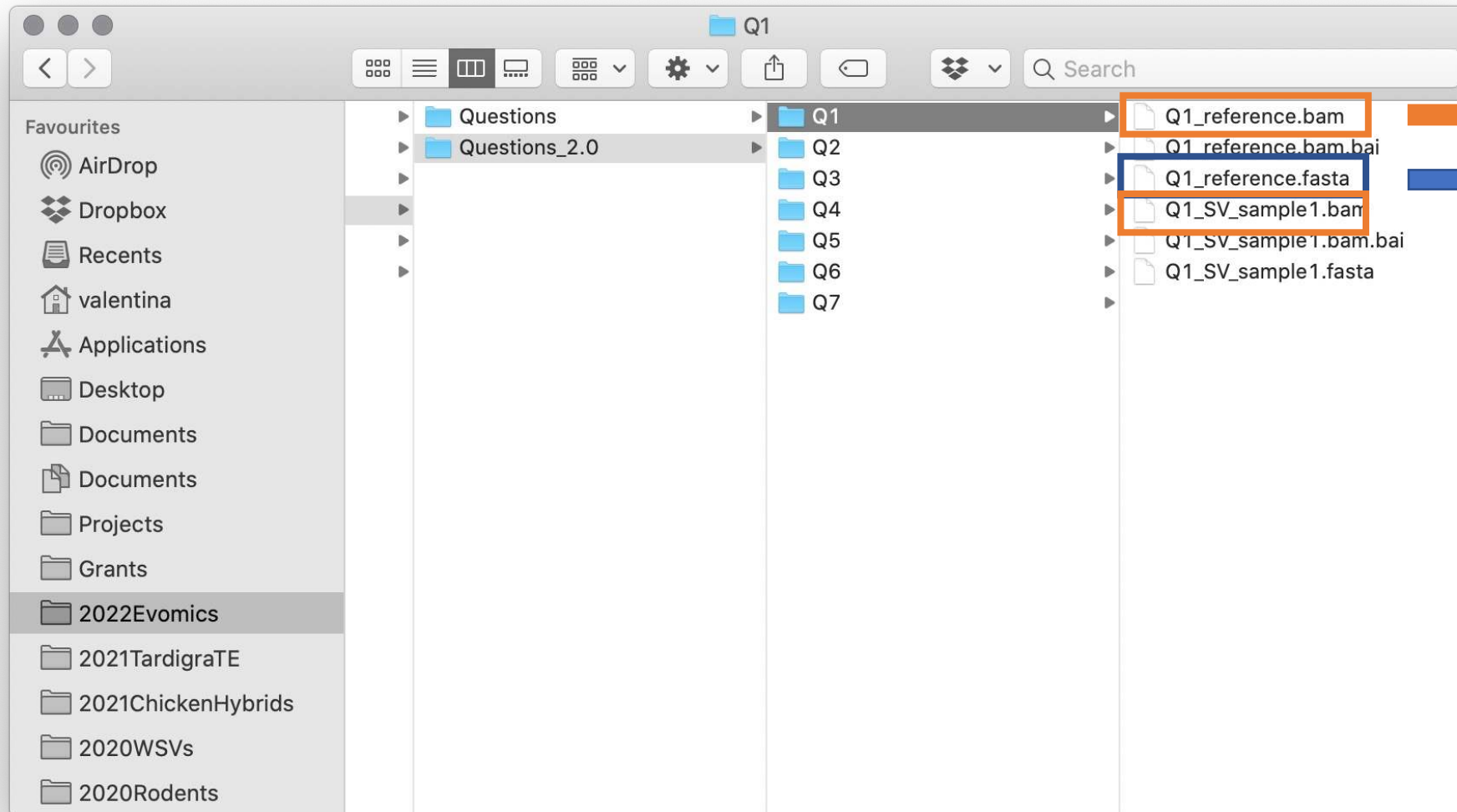
- Open the sequence and respective BAM files in IGV.

Data for the quiz

Reference fasta to be uploaded on IGV through the menu Genomes > Load Genome from File

BAM file of reads from reference mapped to reference: File > Load from File

BAM file of reads from samples mapped to reference: File > Load from File



Upload as "File"

Upload as "Genome"

Reference fasta always named in the form:

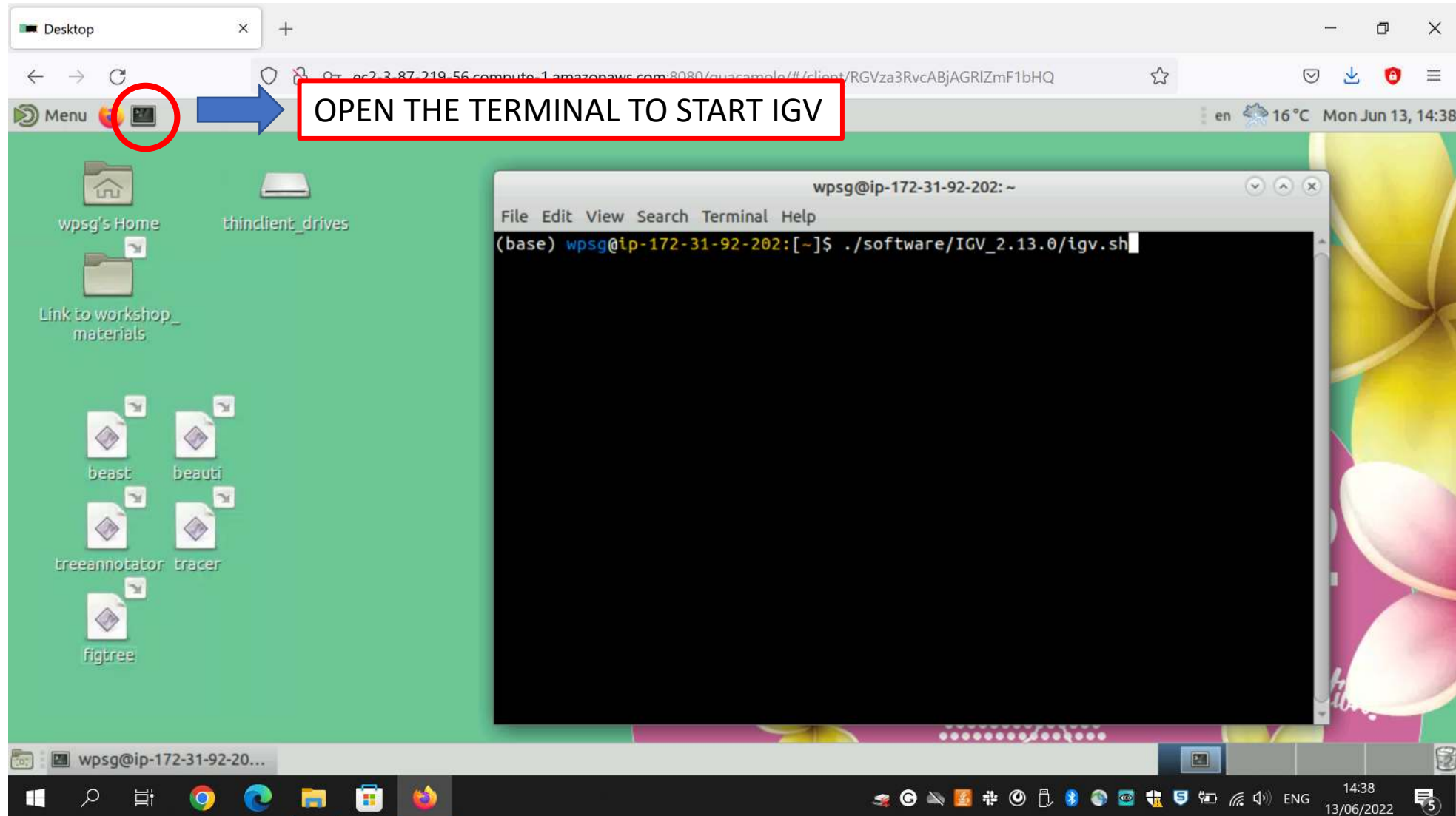
Q*_reference.fasta

BAM reference: Q*_reference.bam

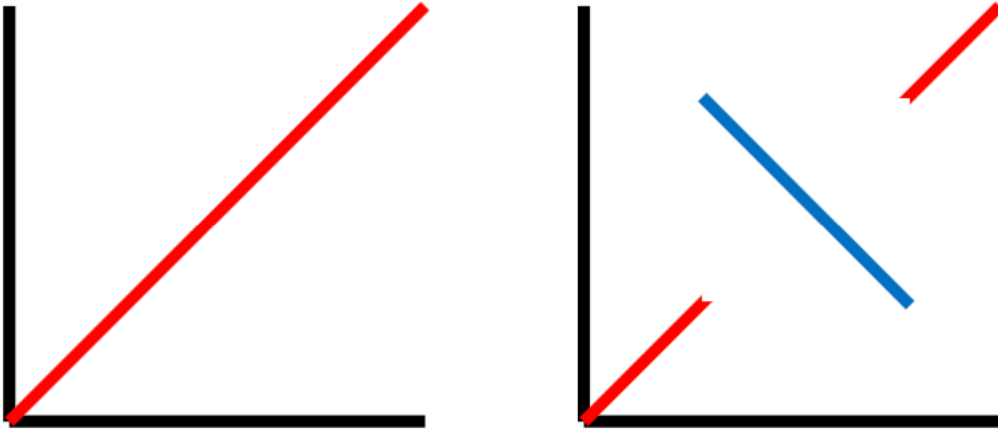
BAM sample(s): Q*_sample*.bam



Open IGV in Guacamole



Dot plot



You can get important information by aligning sequences, not only reads

Copy reference and sample sequences into LAST or MAFFT to get a dot plot

Is your SV a transposon????

Go to Rebase, click on Repeat Masking menu to run Censor on your sequences and find homologies with known transposable elements





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Submit sequence to CENSOR

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[References](#)

Submit sequence to CENSOR

CENSOR is a software tool which screens query sequences against a reference collection of repeats and "censors" (masks) homologous portions with masking symbols, as well as generating a report classifying all found repeats. If you use CENSOR as a tool in your published research, please quote:

[Kohany O](#), [Gentles AJ](#), [Hankus L](#), [Jurka J](#)
Annotation, [submission](#) and screening of repetitive elements in Repbase: RepbaseSubmitter and Censor.
BMC Bioinformatics, 2006 Oct 25;7:474

Sequence source:

All

Force translated search:

☐

Search for identity:

☐

Report simple repeats:

☐

Mask pseudogenes:

☐

Enter query file name:

(Up to 2MB; IG–Stanford, FASTA, GENBANK, EMBL formats are supported)

Choose File no file selected

Submit File

OR

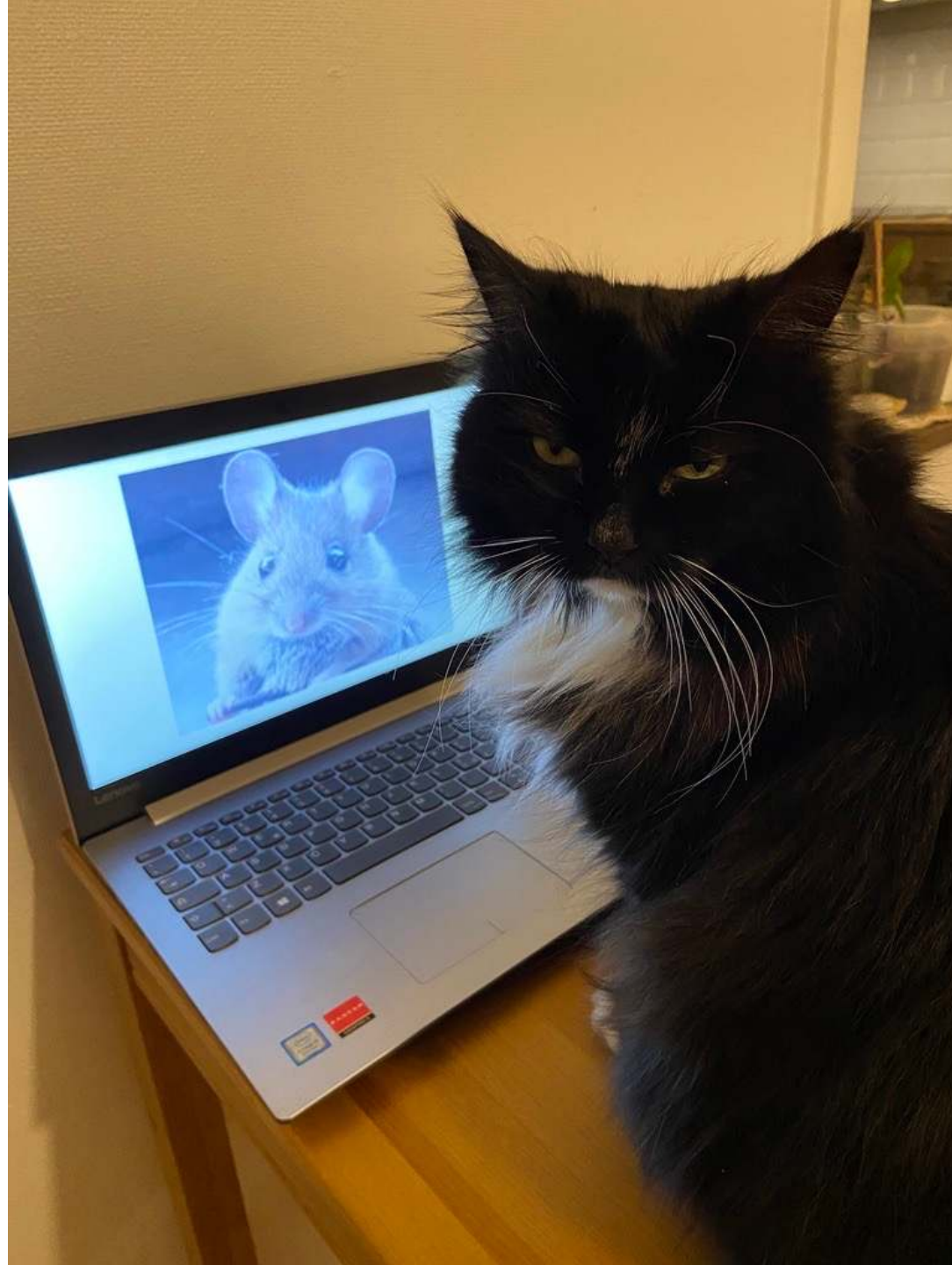
Paste query sequences here:

(Up to 2MB; IG–Stanford, FASTA, GENBANK, EMBL formats are supported)

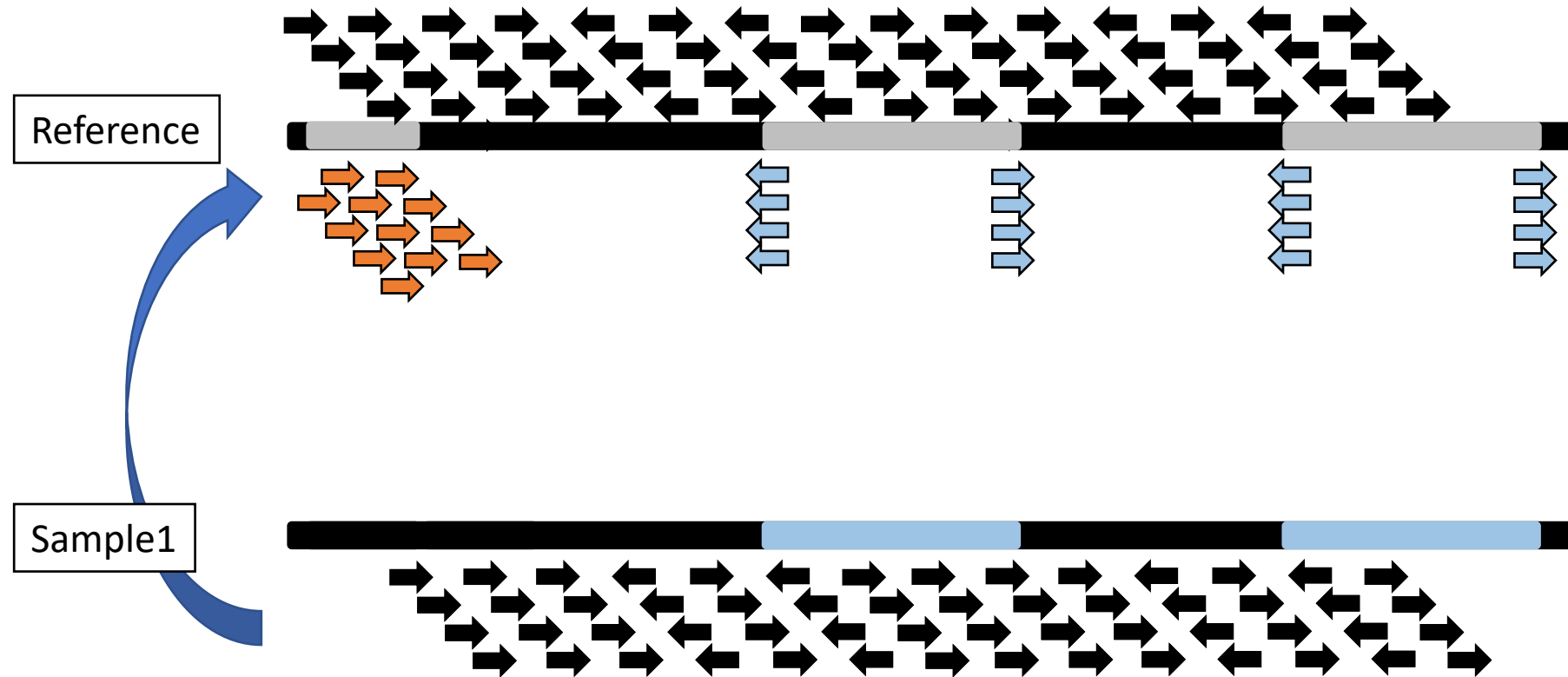
Submit Sequence

Reset

SVs are awesome
but painful



SV simulation



SV simulation

Simulated libraries 30X

Illumina paired-end reads	150 bp x 2, 500 bp insertion size
PacBio reads	6 kb
Nanopore reads	5.5 kb

Two conda envs!

One only for Manta

The second (SV_Env) for all the other analysis

1

Map reads

DO NOT RUN!

2

Call SVs

Manta for IL
Sniffles2 for PB and ONT

~20 min

3

Compare SVs

SURVIVOR merge

4

Evaluate simulation

SURVIVOR eval

WRAP-UP

2

SV calling tutorial

SV simulation

SV CALLING

SV_PB_filtered.vcf

A tibble: 5 × 8

	SVType	Number	MinLen	MaxLen	MeanLen	SDLen	NPrecise	NImprecise
	<chr>	<int>	<int>	<int>	<dbl>	<dbl>	<int>	<int>
1	DEL	10	83	534	244.	138.	10	0
2	DUP	25	1377	18975	7893.	5109.	24	1
3	INS	23	132	795	476.	251.	21	2
4	INV	10	2981	8551	5892.	2021.	10	0
5	ALL	68	83	18975	3965.	4741.	65	3

SV_ONT_filtered.vcf

A tibble: 5 × 8

	SVType	Number	MinLen	MaxLen	MeanLen	SDLen	NPrecise	NImprecise
	<chr>	<int>	<int>	<int>	<dbl>	<dbl>	<int>	<int>
1	DEL	8	83	341	228.	92.4	8	0
2	DUP	25	1377	18986	7894.	5110.	25	0
3	INS	22	118	710	437.	222.	20	2
4	INV	9	2982	8557	5663.	1993.	9	0
5	ALL	64	83	18986	4058.	4808.	62	2

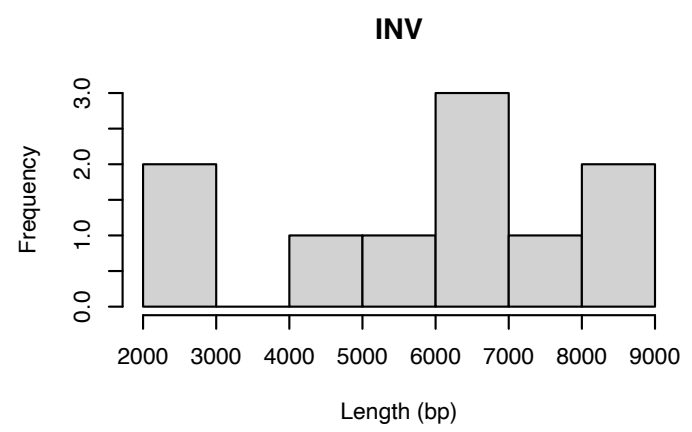
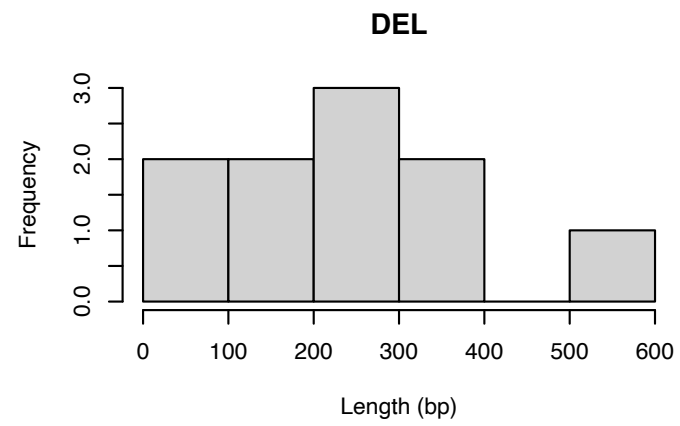
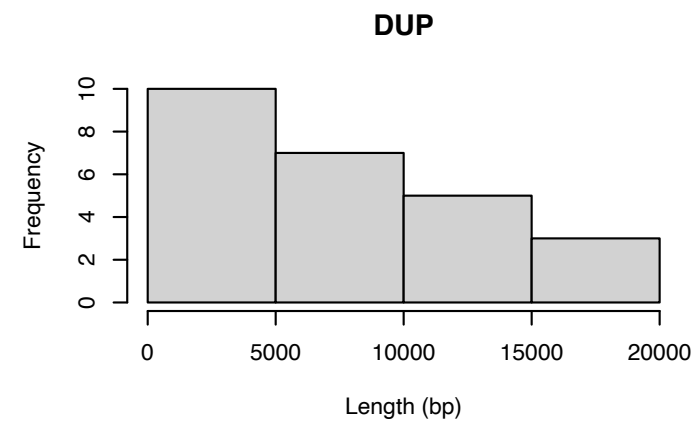
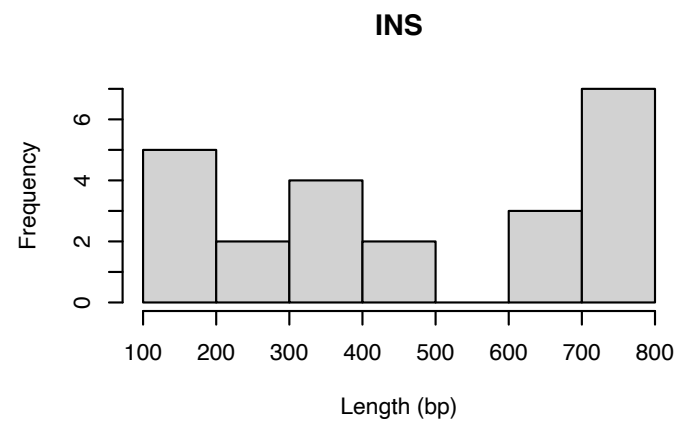
SV_IL_filtered.vcf

A tibble: 4 × 8

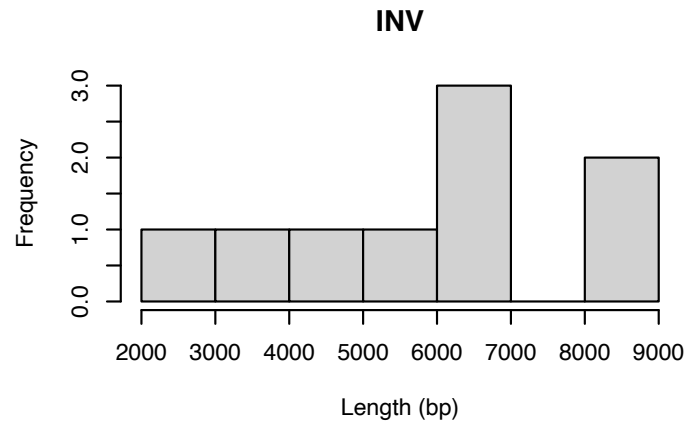
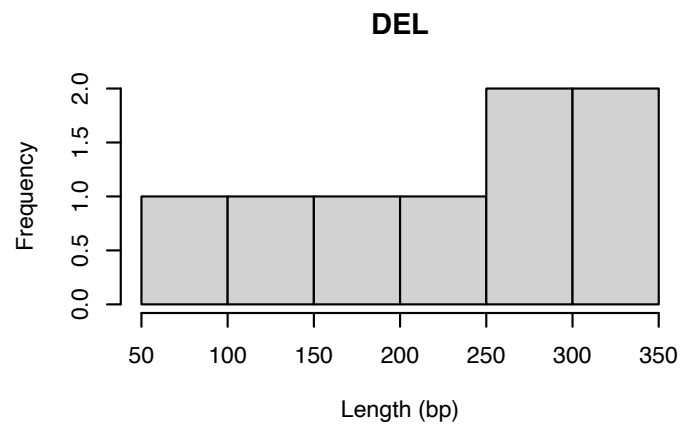
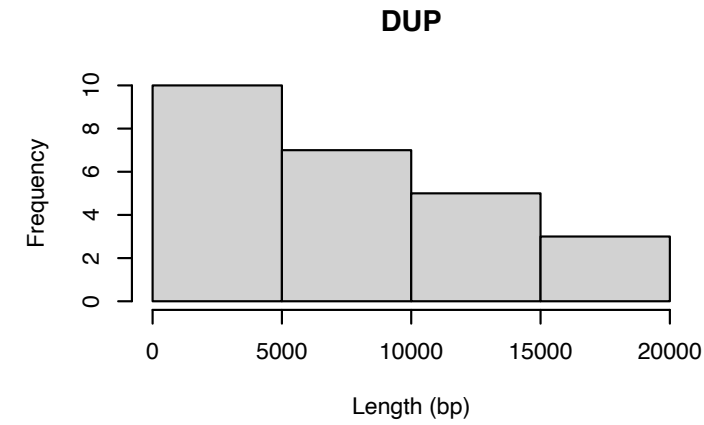
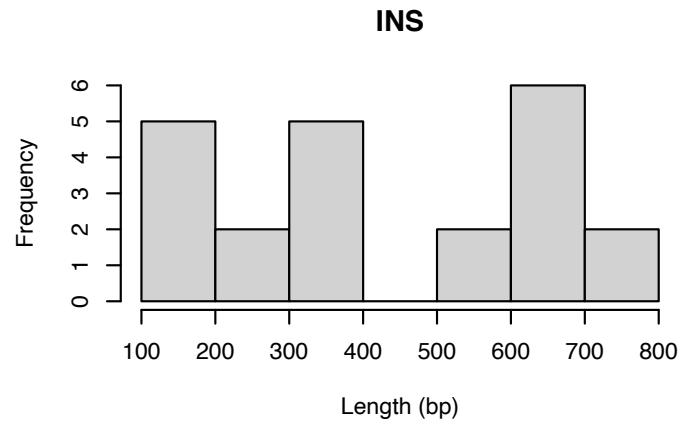
	SVType	Number	MinLen	MaxLen	MeanLen	SDLen	NPrecise	NImprecise
	<chr>	<int>	<int>	<int>	<dbl>	<dbl>	<int>	<int>
1	DEL	1	11	11	11	NA	1	0
2	DUP	14	1576	19023	7255.	5478.	0	14
3	INV	45	13	8533	2530.	2822.	1	44
4	ALL	60	11	19023	3590.	4100.	2	58

SIMULATED

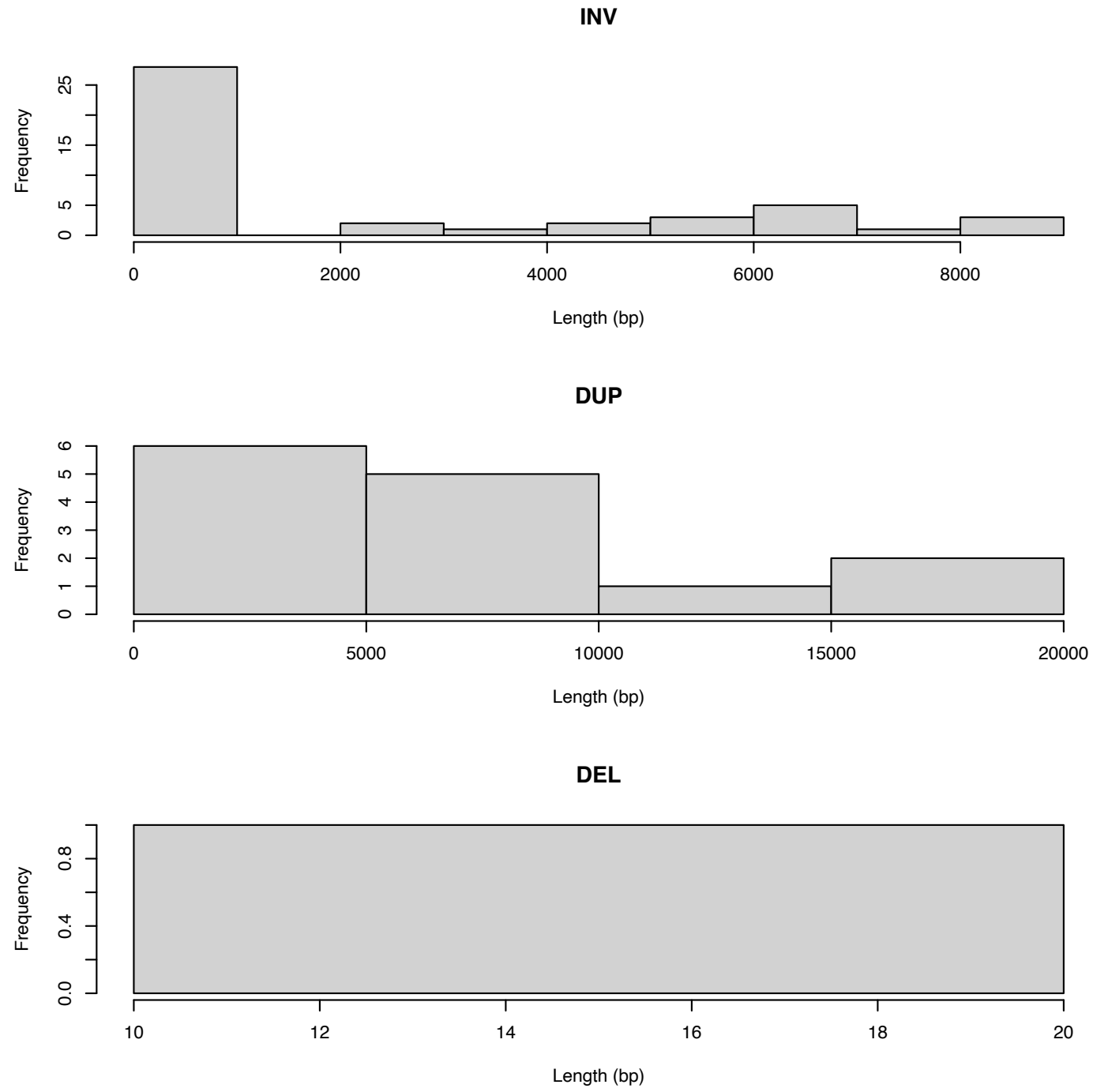
29	DEL
40	DUP
11	INS
30	INV



ONT

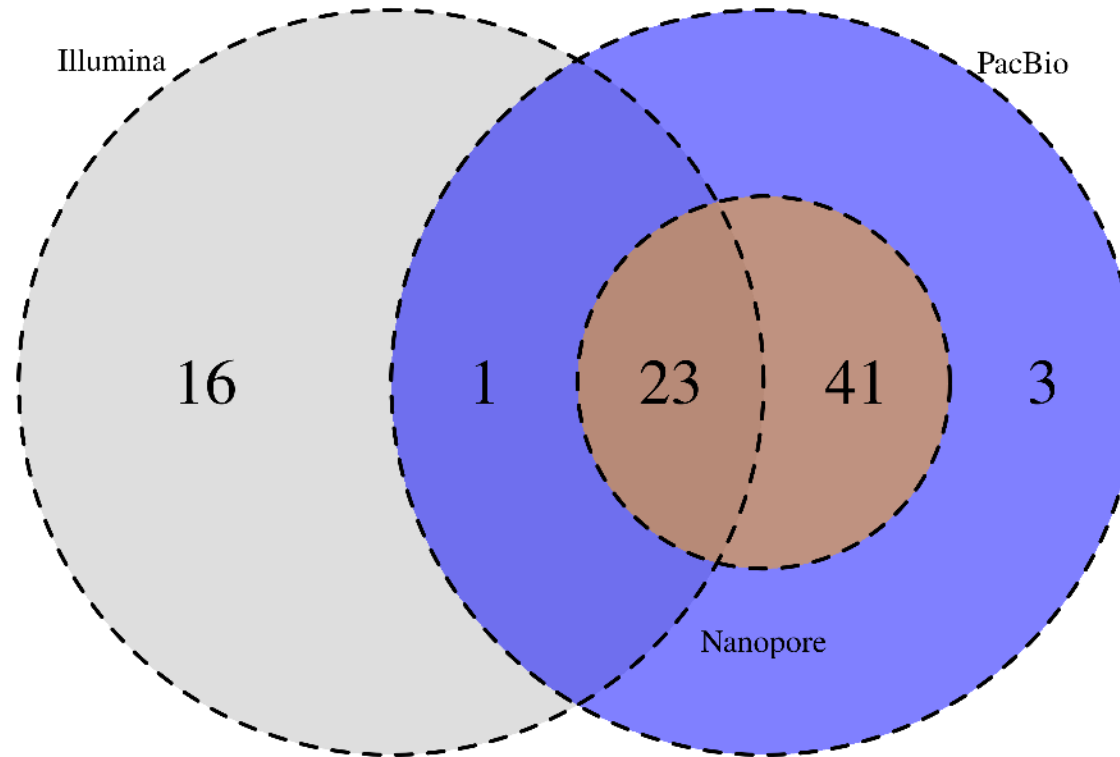


IL



SV simulation

COMPARISON



SV simulation

EVALUATION

		True positives DEL/DUP/INV/TRA/INS					False positives DEL/DUP/INV/TRA/INS					Sensitivity True positive rate		
IL	Overall:	110	0/0/0/0/0	29/40/30/0/11	1/14/45/0/0	0/1								
PB	Overall:	110	8/20/8/0/10	21/20/22/0/1	2/5/2/0/13	0.418182	0.323529							
ONT	Overall:	110	8/21/9/0/10	21/19/21/0/1	0/4/0/0/12	0.436364	0.25							
		N. SV simulated		False negatives DEL/DUP/INV/TRA/INS					FDR					

So what????

Quality control and correction of reads

Check for mappability

Test more tools

Simulate on your data and reference to find FDR

Increase in read length

Assembly-based approach

Report

Expectations and blind spots for structural variation detection from long-read assemblies and short-read genome sequencing technologies

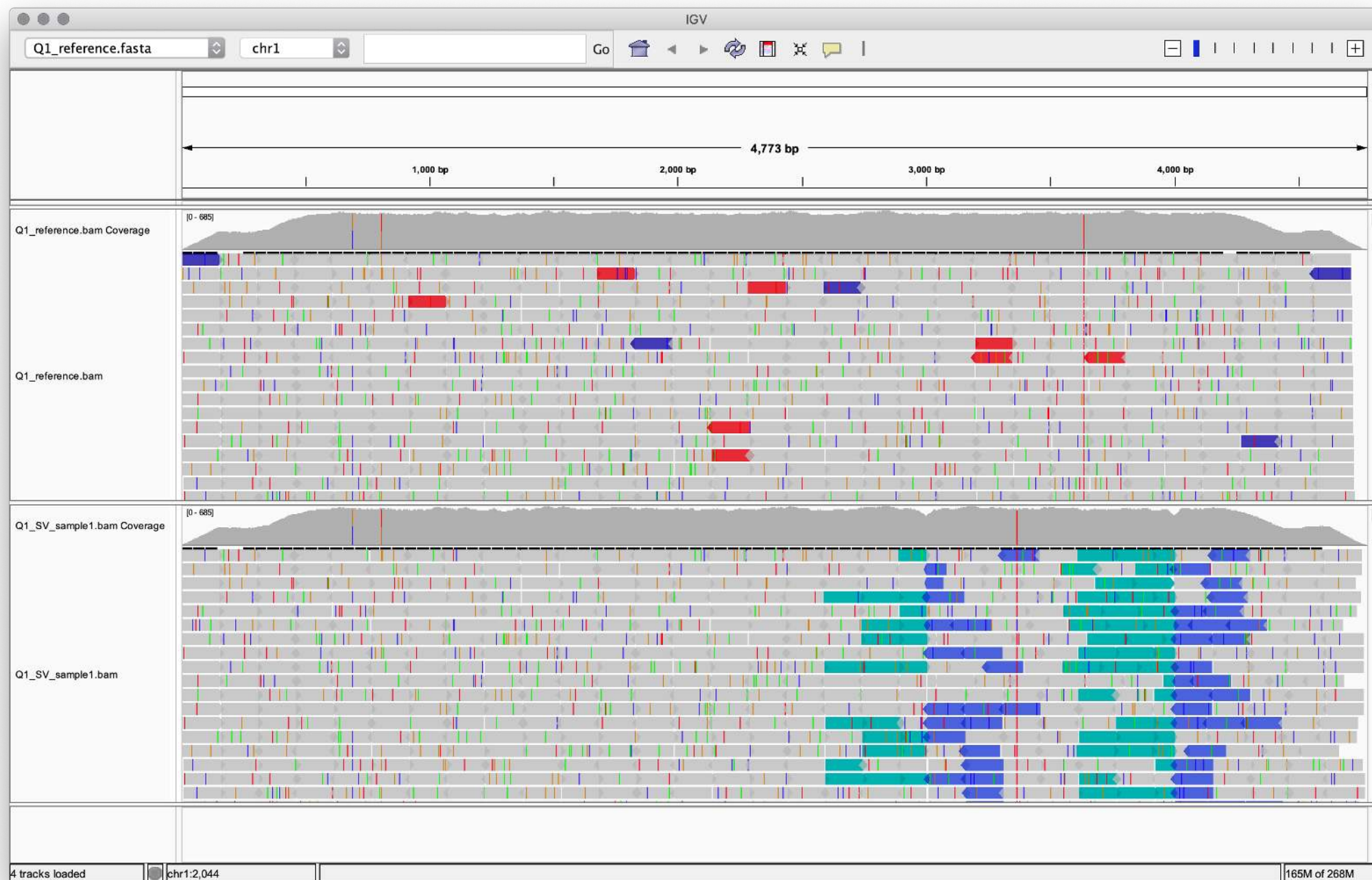
Xuefang Zhao ^{1, 2, 3}, Ryan L. Collins ^{1, 2, 4}, Wan-Ping Lee ⁵, Alexandra M. Weber ^{6, 7}, Yookyung Jun ⁵, Qihui Zhu ⁵, Ben Weisburd ², Yongqing Huang ⁸, Peter A. Audano ⁹, Harold Wang ^{1, 2}, Mark Walker ^{2, 3}, Chelsea Lowther ^{1, 2, 3}, Jack Fu ^{1, 2, 3}, Human Genome Structural Variation Consortium, Mark B. Gerstein ¹⁰, Scott E. Devine ¹¹, Tobias Marschall ¹², Jan O. Korbel ^{13, 14} ... Michael E. Talkowski ^{1, 2, 3, 4}  

Finally, we explored the concordance of SV detection for a class of SVs that is strongly enriched for pathogenic variation and appears to be a significant blind spot for long-read assembly technologies: large CNVs captured by depth-based analyses from srWGS. Our initial analyses suggested that lrWGS assembly methods failed to capture all but one of the small number of large (>5 kb) CNVs that could be detected by srWGS read-depth methods in three probands (average size = 14.7 kb). Recognizing the limitation of read-depth analyses to

1

SV quiz → ANSWERS and PRIZES

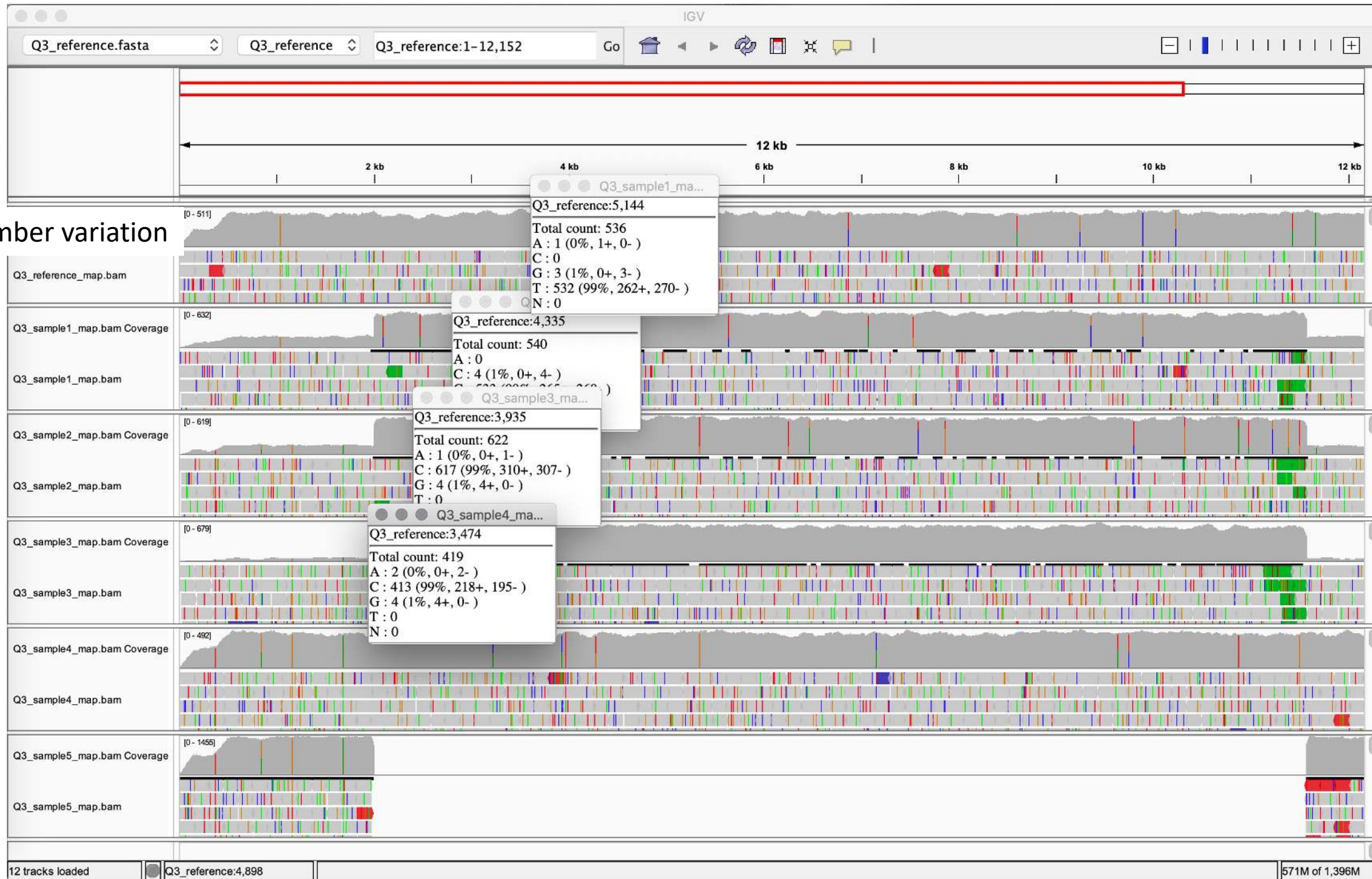
Q1: inversion



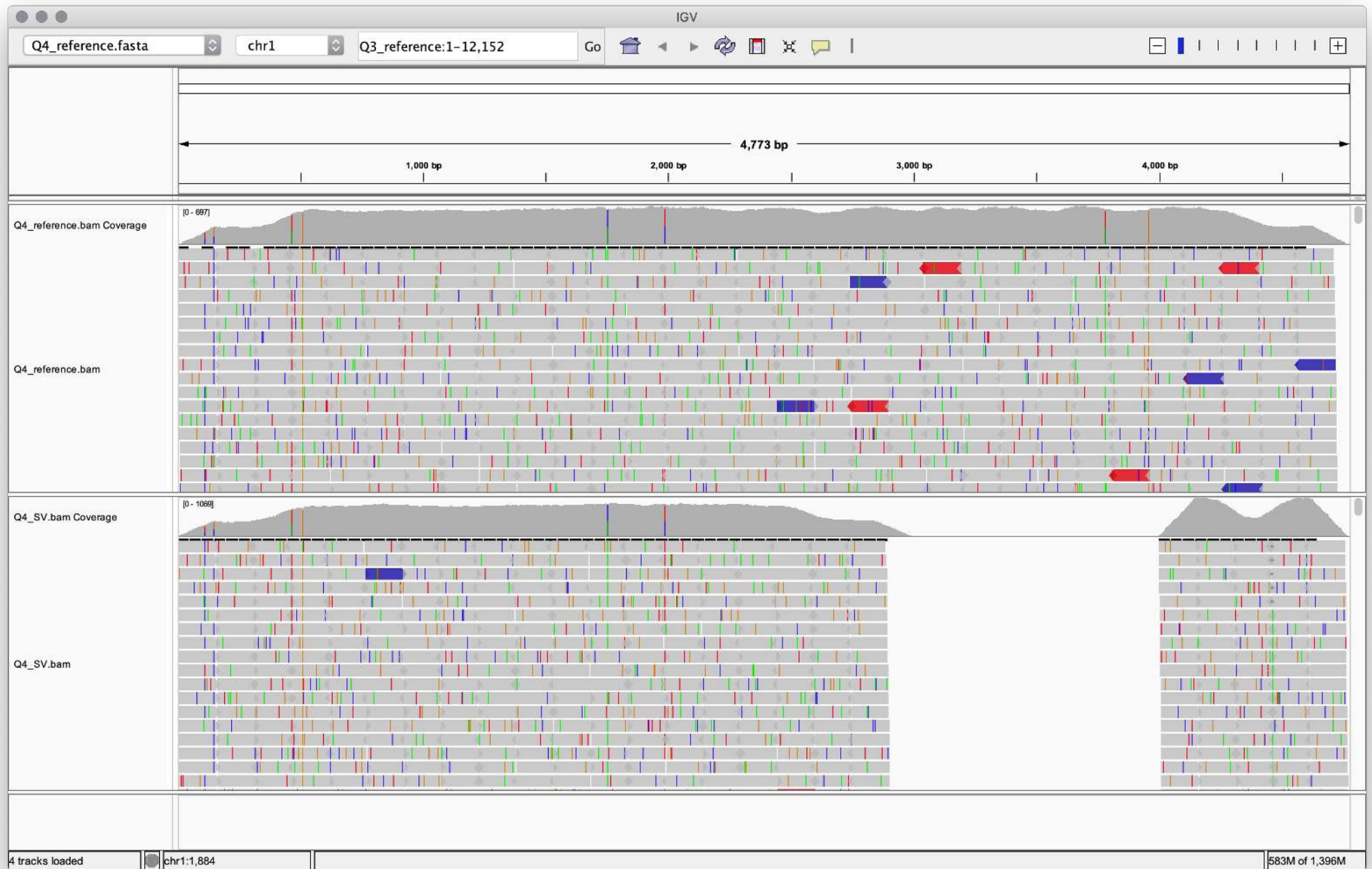
Q2: duplication



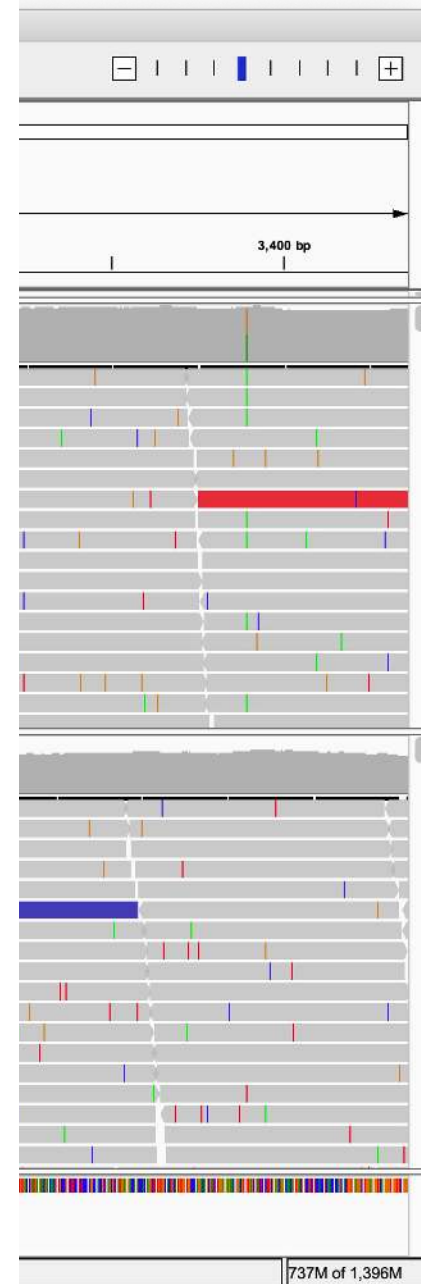
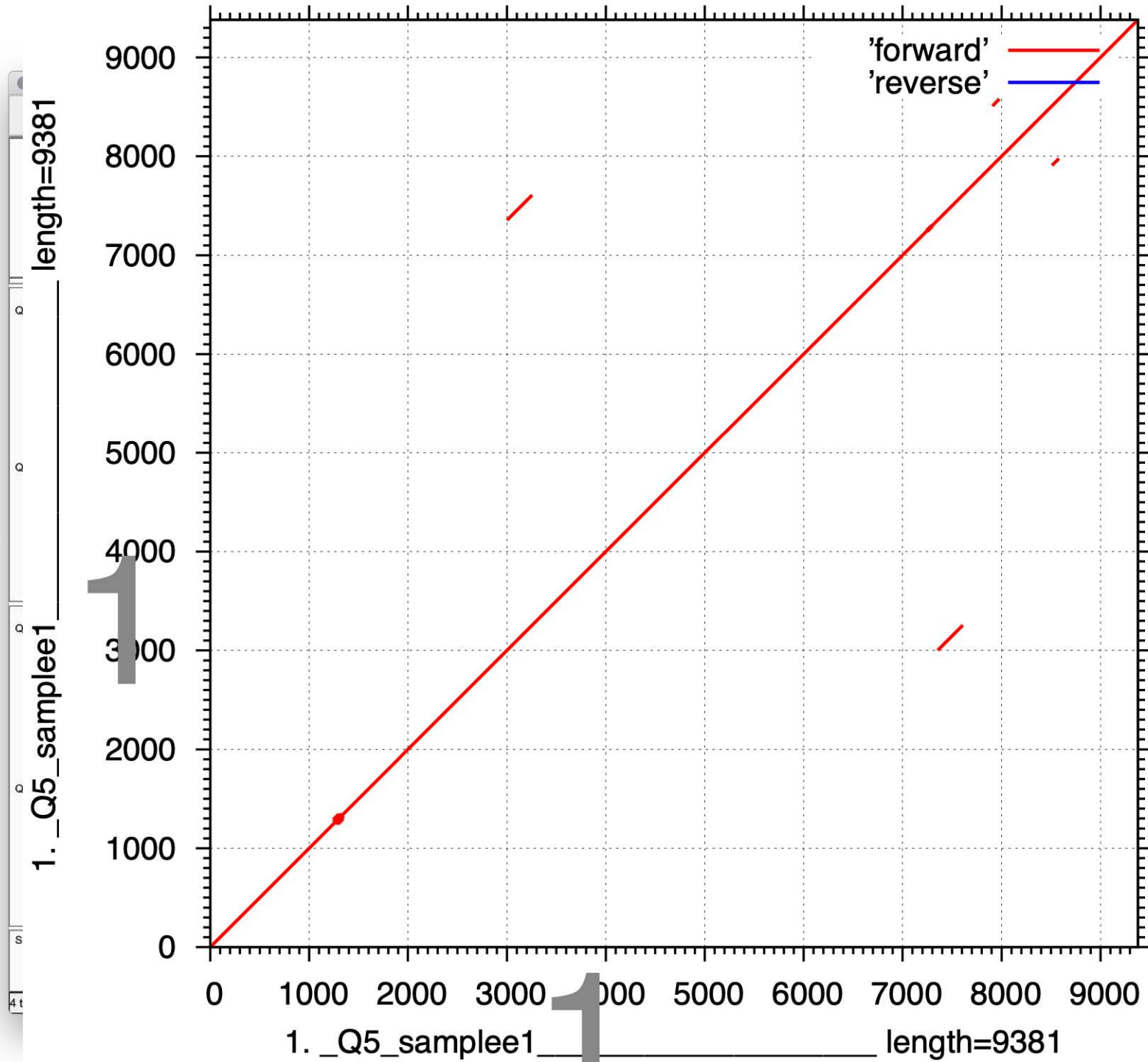
Q3: copy number variation



Q4: deletion



Q5: LTR insertion



Q6: LTR insertion
soloLTR



Q7: DNA TE

