Pipeline Report

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April 29, 2022

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1 Metadata information

1.1 Constants

The following information is not automatically updated in the report: Build: b38~(hg38)

1.2 Dynamic

The following information is gathered dynamically when a report is generated:

Library	Sample	Preparation	Gbp
HALB3002753	NA12878	PCR-free-illumina	181.14
HALB3002754	HG002	PCR-free-illumina	145.87
HALB3002755	HG003	PCR-free-illumina	149.77
HALB3002756	HG004	PCR-free-illumina	138.30
HALB3002757	HG005	PCR-free-illumina	159.85
HALB3003200	NA12878	PCR-free-illumina	167.53
HALB3003201	HG002	PCR-free-illumina	129.81
HALB3003222	HG003	PCR-free-illumina	156.20
HALB3010451	NA12878	PacBio-CCS-high	77.46
HALB3010452	HG002	PacBio-CCS-high	166.22
HALB3010453	HG003	PacBio-CCS-high	129.52
HALB3010454	HG004	PacBio-CCS-high	123.81
HALB3010455	HG005	PacBio-CCS-high	145.34
HALB3010456	HG006	PacBio-CCS-high	122.17
HALB3010457	HG007	PacBio-CCS-high	113.30

2 Summarized Small Variant Comparison

This section contains the summarized results of the small variant analysis. For each aligner-caller pair, we ran RTG on the Genome in a Bottle (GIAB) high confidence truth set ($\approx 3.6 \mathrm{M}$ variants). The following results are gathered from the RTG 'summary.txt' file using the final row (i.e. "None" indicating no variants were filtered) and with the --all-records option enabled.

2.1 Preparation: "PCR-free-illumina"

Reference	Aligner	Caller	N	Precision	Sensitivity	F-measure
clinical	sentieon-201808.07	strelka-2.9.10	8	0.9923 + -0.0014	0.9920 + -0.0008	0.9922+-0.0010
hg38_T2T_masked	dragmap-1.2.1	clair3-0.1-r11	8	0.9976 + -0.0003	0.9920 + -0.0006	0.9948 + -0.0003
hg38_T2T_masked	dragmap-1.2.1	dnascope-1.0-202112.01-PO	8	0.9984 + -0.0003	0.9942 + -0.0008	0.9963 + -0.0005
hg38_T2T_masked	dragmap-1.2.1	octopus-0.7.4	8	0.9962 + -0.0003	0.9899 + -0.0008	0.9931 + -0.0005
hg38_T2T_masked	dragmap-1.2.1	strelka-2.9.10	8	0.9893 + -0.0039	0.9922 + -0.0009	0.9907 + -0.0023
hg38_T2T_masked	sentieon-202112.01	clair3-0.1-r11	8	0.9974 + -0.0003	0.9927 + -0.0009	0.9950 + -0.0005
hg38_T2T_masked	sentieon-202112.01	dnascope-1.0-202112.01-PO	8	0.9983 + -0.0003	0.9944 + -0.0007	0.9963 + -0.0005
hg38_T2T_masked	sentieon-202112.01	octopus-0.7.4	8	0.9946 + -0.0003	0.9903 + -0.0007	0.9924 + -0.0005
hg38_T2T_masked	sentieon-202112.01	strelka-2.9.10	8	0.9869 + -0.0039	0.9932 + -0.0007	0.9900 + -0.0022
hg38_T2T_masked	sentieon-202112.01-recal	clair3-0.1-r11	8	0.9974 + -0.0003	0.9918+-0.0010	0.9945 + -0.0006
hg38_T2T_masked	sentieon-202112.01-recal	dnascope-1.0-202112.01-PO	8	0.9986 + -0.0002	0.9932 + -0.0009	0.9959 + -0.0005
hg38_T2T_masked	sentieon-202112.01-recal	octopus-0.7.4	8	0.9947 + -0.0004	0.9903 + -0.0008	0.9925 + -0.0005
hg38_T2T_masked	sentieon-202112.01-recal	strelka-2.9.10	8	0.9905 + -0.0016	0.9930 + -0.0007	0.9918 + -0.0011
hg38_T2T_masked	snap-2.0.0	clair3-0.1-r11	8	0.9966 + -0.0003	0.9921 + -0.0011	0.9943 + -0.0007
hg38_T2T_masked	snap-2.0.0	dnascope-1.0-202112.01-PO	8	0.9976 + -0.0003	0.9943 + -0.0006	0.9959 + -0.0004
hg38_T2T_masked	snap-2.0.0	octopus-0.7.4	8	0.9938 + -0.0003	0.9907 + -0.0007	0.9922 + -0.0004
hg38_T2T_masked	snap-2.0.0	strelka-2.9.10	8	0.9861 + -0.0042	0.9934 + -0.0007	0.9898 + -0.0024
hg38_asm5_alt	dragmap-1.2.1	clair3-0.1-r11	8	0.9975 + -0.0002	0.9695 + -0.0012	0.9833 + -0.0007
hg38_asm5_alt	dragmap-1.2.1	dnascope-1.0-202112.01-PO	8	0.9978 + -0.0002	0.9735 + -0.0010	0.9855 + -0.0006
hg38_asm5_alt	dragmap-1.2.1	octopus-0.7.4	8	0.9969 + -0.0002	0.9681 + -0.0011	0.9823 + -0.0006
hg38_asm5_alt	dragmap-1.2.1	strelka-2.9.10	8	0.9902 + -0.0038	0.9699 + -0.0011	0.9800 + -0.0023
hg38_asm5_alt	sentieon-202112.01	clair3-0.1-r11	8	0.9977 + -0.0002	0.9922 + -0.0009	0.9949 + -0.0005
hg38_asm5_alt	sentieon-202112.01	dnascope-1.0-202112.01-PO	8	0.9983 + -0.0002	0.9938 + -0.0007	0.9961 + -0.0004
hg38_asm5_alt	sentieon-202112.01	octopus-0.7.4	8	0.9958 + -0.0002	0.9893 + -0.0008	0.9926 + -0.0004
hg38_asm5_alt	sentieon-202112.01	strelka-2.9.10	8	0.9890 + -0.0038	0.9921 + -0.0007	0.9906+-0.0022

Reference	Aligner	Caller	N	Precision	Sensitivity	F-measure
hg38_asm5_alt	sentieon-202112.01-recal	clair3-0.1-r11	8	0.9978 + -0.0002	0.9911 + -0.0008	0.9944 + -0.0005
hg38_asm5_alt	sentieon-202112.01-recal	dnascope-1.0-202112.01-PO	8	0.9986 + -0.0002	0.9926 + -0.0008	0.9956 + -0.0005
hg38_asm5_alt	sentieon-202112.01-recal	octopus-0.7.4	8	0.9960 + -0.0002	0.9893 + -0.0008	0.9926 + -0.0004
hg38_asm5_alt	sentieon-202112.01-recal	strelka-2.9.10	8	0.9923 + -0.0014	0.9920 + -0.0008	0.9922 + -0.0010
hg38_asm5_alt	snap-2.0.0	clair3-0.1-r11	8	0.9967 + -0.0003	0.9920 + -0.0008	0.9944 + -0.0005
hg38_asm5_alt	snap-2.0.0	deepvariant-1.2.0	8	0.9984 + -0.0002	0.9936 + -0.0005	0.9960 + -0.0003
hg38_asm5_alt	snap-2.0.0	dnascope-1.0-202112.01-PO	8	0.9976 + -0.0003	0.9941 + -0.0006	0.9959 + -0.0004
hg38_asm5_alt	snap-2.0.0	octopus-0.7.4	8	0.9944+-0.0003	0.9905 + -0.0007	0.9925 + -0.0004
hg38_asm5_alt	snap-2.0.0	strelka-2.9.10	8	0.9868 + -0.0042	0.9932 + -0.0007	0.9900+-0.0024

${\bf 2.2}\quad {\bf Preparation:~"PacBio-CCS-high"}$

Reference	Aligner	Caller	N	Precision	Sensitivity	F-measure
hg38_T2T_masked	minimap2-2.23	clair3-0.1-r11	7	0.9980 + -0.0013	0.9987 + -0.0006	0.9984 + -0.0010
hg38_T2T_masked	minimap2-2.23	dnascope_hifi-0.4-202112.01	7	0.9986 + -0.0008	0.9986 + -0.0006	0.9986 + -0.0007
hg38_T2T_masked	minimap2-2.23	peppermargindv-r0.8	7	0.9983 + -0.0009	0.9985 + -0.0005	0.9984 + -0.0007
hg38_T2T_masked	pbmm2-1.7.0	clair3-0.1-r11	7	0.9984 + -0.0013	0.9987 + -0.0006	0.9985 + -0.0009
hg38_T2T_masked	pbmm2-1.7.0	dnascope_hifi-0.4-202112.01	7	0.9987 + -0.0008	0.9987 + -0.0005	0.9987 + -0.0007
hg38_T2T_masked	pbmm2-1.7.0	peppermargindv-r0.8	7	0.9985 + -0.0009	0.9984 + -0.0005	0.9985 + -0.0007
hg38_T2T_masked	sentieon_mm2-202112.01	clair3-0.1-r11	7	0.9980 + -0.0013	0.9987 + -0.0006	0.9983+-0.0010
hg38_T2T_masked	sentieon_mm2-202112.01	dnascope_hifi-0.4-202112.01	7	0.9986 + -0.0008	0.9986 + -0.0006	0.9986 + -0.0007
hg38_T2T_masked	sentieon_mm2-202112.01	peppermargindv-r0.8	7	0.9983 + -0.0009	0.9985 + -0.0005	0.9984+-0.0007

3 Summarized SNP/Indel Comparison

This section contains the summarized results of the small variant analysis when stratified by SNP/Indel. Files were filtered using bcftools filter to pair down to SNPs or Indels only. For each aligner-caller pair, we ran RTG on the Genome in a Bottle (GIAB) high confidence truth set (≈3.6M variants). The following results are gathered from the RTG 'summary.txt' file using the final row (i.e. "None" indicating no variants were filtered) and with the --all-records option enabled.

3.1 Preparation: "PCR-free-illumina"

Reference	Aligner	Caller	Type	N	Precision	Sensitivity	F-measure
			SNP	8	0.9912+-0.0016	0.9922+-0.0007	0.9917+-0.0010
clinical	sentieon-201808.07	strelka-2.9.10	Indel	8	0.9885 + -0.0009	0.9839 + -0.0018	0.9862 + -0.0013
1 00 TOTT 1 1	1 101	1 . 0 0 1 11	SNP	8	0.9976 + -0.0003	0.9920 + -0.0006	0.9948+-0.0003
hg38_T2T_masked	dragmap-1.2.1	clair3-0.1-r11	Indel	8	0.9949 + -0.0006	0.9890+-0.0010	0.9920+-0.0008
l20 TOTll	d 1 0 1	-l	SNP	8	0.9986 + -0.0002	0.9940 + -0.0008	0.9963 + -0.0005
hg38_T2T_masked	dragmap-1.2.1	dnascope-1.0-202112.01-PO	Indel	8	0.9968 + -0.0005	0.9946 + -0.0008	0.9957 + -0.0006
ha20 TOT modered	dragman 1 2 1	actorus 0.7.4	SNP	8	0.9961 + -0.0003	0.9851 + -0.0012	0.9906 + -0.0006
hg38_T2T_masked	38_T2T_masked dragmap-1.2.1	octopus-0.7.4	Indel	8	0.9476 + -0.0026	0.9612 + -0.0024	0.9544+-0.0024
hg38_T2T_masked	dragmap-1.2.1	strelka-2.9.10	SNP	8	0.9878 + -0.0045	0.9924+-0.0009	0.9900 + -0.0025
lig56_121_illasked	dragmap-1.2.1	streika-2.9.10	Indel	8	0.9883 + -0.0010	0.9841 + -0.0019	0.9862 + -0.0014
hg38_T2T_masked	sentieon-202112.01	clair3-0.1-r11	SNP	8	0.9973 + -0.0003	0.9929 + -0.0009	0.9951 + -0.0005
lig56_121_illasked	sentieon-202112.01	Claif 5-0.1-111	Indel	8	0.9948 + -0.0006	0.9889 + -0.0015	0.9918+-0.0010
hg38_T2T_masked	sentieon-202112.01	dnascope-1.0-202112.01-PO	SNP	8	0.9984 + -0.0002	0.9942 + -0.0007	0.9963 + -0.0004
ngoo_121_masked	Sentieon-202112.01	dnascope-1:0-202112:01-1 O	Indel	8	0.9968 + -0.0005	0.9946 + -0.0008	0.9957 + -0.0006
hg38_T2T_masked	sentieon-202112.01	octopus-0.7.4	SNP	8	0.9944 + -0.0004	0.9854 + -0.0011	0.9899 + -0.0006
ngoo_121_masked	Sentieon-202112.01	octopus-0.7.4	Indel	8	0.9469 + -0.0026	0.9612 + -0.0023	0.9540 + -0.0023
hg38_T2T_masked	sentieon-202112.01	strelka-2.9.10	SNP	8	0.9852 + -0.0045	0.9935 + -0.0006	0.9893 + -0.0025
lig50_121_lilasked	Sentieon-202112.01	Streika-2.9.10	Indel	8	0.9870 + -0.0010	0.9844 + -0.0018	0.9857 + -0.0014
hg38_T2T_masked	sentieon-202112.01-recal	clair3-0.1-r11	SNP	8	0.9974 + -0.0003	0.9919 + -0.0009	0.9947 + -0.0005
iig56_121_iiia5keu	Sentieon-202112.01-recar	Claif 5-0.1-111	Indel	8	0.9942 + -0.0008	0.9881 + -0.0016	0.9911+-0.0011
hg38_T2T_masked	sentieon-202112.01-recal	dnascope-1.0-202112.01-PO	SNP	8	0.9987 + -0.0002	0.9931 + -0.0009	0.9959 + -0.0005
ngoo_121_maskeu senueon-202112.01-recar	dnascope-1.0-202112.01-PO	Indel	8	0.9967 + -0.0006	0.9927 + -0.0014	0.9947 + -0.0010	
hg38_T2T_masked	sentieon-202112.01-recal	octopus-0.7.4	SNP	8	0.9944 + -0.0004	0.9854 + -0.0011	0.9899+-0.0006
11500_121_111a5kCd	Scholodi-202112.01-10cai	- 00:0pub 0.1.4	Indel	8	0.9476 + -0.0025	0.9613 + -0.0023	0.9544 + -0.0023

Reference	Aligner	Caller	Type	N	Precision	Sensitivity	F-measure
l20 TOT ll	sentieon-202112.01-recal	strelka-2.9.10	SNP	8	0.9893+-0.0018	0.9933 + -0.0006	0.9913+-0.0012
hg38_T2T_masked	sentieon-202112.01-recai	streika-2.9.10	Indel	8	0.9870 + -0.0010	0.9844 + -0.0018	0.9857+-0.0014
hg38_T2T_masked	snap-2.0.0	clair3-0.1-r11	SNP	8	0.9964 + -0.0003	0.9921 + -0.0011	0.9942 + -0.0006
ngoo_121_masked	Shap-2.0.0	Ciaii 5-0.1-111	Indel	8	0.9943 + -0.0006	0.9895 + -0.0018	0.9919 + -0.0012
hg38_T2T_masked	snap-2.0.0	dnascope-1.0-202112.01-PO	SNP	8	0.9977 + -0.0002	0.9941 + -0.0006	0.9959 + -0.0004
ng36_121_masked	snap-2.0.0	dilascope-1.0-202112.01-1 O	Indel	8	0.9962 + -0.0004	0.9945 + -0.0007	0.9953 + -0.0005
hg38_T2T_masked	snap-2.0.0	octopus-0.7.4	SNP	8	0.9936+-0.0003	0.9858 + -0.0011	0.9897 + -0.0006
ngoo_121_masked	Shap-2.0.0	octopus-0.1.4	Indel	8	0.9463 + -0.0025	0.9609 + -0.0023	0.9535 + -0.0023
hg38_T2T_masked	snap-2.0.0	strelka-2.9.10	SNP	8	0.9845 + -0.0048	0.9937 + -0.0006	0.9891 + -0.0026
ngoo_121_masked	Shap-2.0.0	Strcika-2.3.10	Indel	8	0.9853 + -0.0009	0.9849 + -0.0018	0.9851 + -0.0014
hg38_asm5_alt	dragmap-1.2.1	clair3-0.1-r11	SNP	8	0.9975 + -0.0002	0.9691 + -0.0012	0.9831 + -0.0006
11500_a51119_a10	dragmap-1.2.1	Clairy 0.1 111	Indel	8	0.9948 + -0.0006	0.9703 + -0.0019	0.9824 + -0.0012
hg38_asm5_alt	dragmap-1.2.1	dnascope-1.0-202112.01-PO	SNP	8	0.9980 + -0.0002	0.9728 + -0.0010	0.9853 + -0.0006
11500_a51119_a10	dragmap-1.2.1	dhascope 1.0 202112.01-1 0	Indel	8	0.9966 + -0.0006	0.9776 + -0.0010	0.9870 + -0.0007
hg38_asm5_alt	dragmap-1.2.1	octopus-0.7.4	SNP	8	0.9968 + -0.0002	0.9628 + -0.0016	0.9796 + -0.0008
11890-2001119-2010	dragmap 1.2.1	octopus o.r.1	Indel	8	0.9482 + -0.0027	0.9436 + -0.0024	0.9459 + -0.0024
$hg38_asm5_alt$	dragmap-1.2.1	strelka-2.9.10	SNP	8	0.9886 + -0.0044	0.9696 + -0.0011	0.9790 + -0.0026
11890-201119-2010	dragmap 1.2.1	5010Hat 2.5.10	Indel	8	0.9894+-0.0009	0.9656 + -0.0021	0.9773 + -0.0015
hg38_asm5_alt	sentieon-202112.01	clair3-0.1-r11	SNP	8	0.9976 + -0.0002	0.9923 + -0.0009	0.9950 + -0.0005
11890260511102611	5011010011 202112.01	010110 0.1 111	Indel	8	0.9950 + -0.0005	0.9890 + -0.0012	0.9920+-0.0008
hg38_asm5_alt	sentieon-202112.01	dnascope-1.0-202112.01-PO	SNP	8	0.9984 + -0.0002	0.9936 + -0.0007	0.9960 + -0.0004
11890241011102411	5011010011 202112.01	diageope 1.0 202112.01 1 0	Indel	8	0.9970 + -0.0004	0.9943 + -0.0007	0.9957 + -0.0006
hg38_asm5_alt	sentieon-202112.01	octopus-0.7.4	SNP	8	0.9957 + -0.0002	0.9844 + -0.0012	0.9900+-0.0006
11890-001110-011	5011010011 202112101	occopias cr.vi	Indel	8	0.9478 + -0.0025	0.9608 + -0.0023	0.9543 + -0.0023
hg38_asm5_alt	sentieon-202112.01	strelka-2.9.10	SNP	8	0.9874 + -0.0043	0.9924 + -0.0007	0.9899 + -0.0024
	5011010011 202112101	20120	Indel	8	0.9884 + -0.0008	0.9839 + -0.0018	0.9861+-0.0013
$hg38_asm5_alt$	sentieon-202112.01-recal	clair3-0.1-r11	SNP	8	0.9978 + -0.0002	0.9912 + -0.0008	0.9945 + -0.0004
	20212101 10001	010110 011 111	Indel	8	0.9945 + -0.0008	0.9880+-0.0011	0.9912+-0.0009
$hg38_asm5_alt$	sentieon-202112.01-recal	dnascope-1.0-202112.01-PO	SNP	8	0.9988+-0.0001	0.9925+-0.0008	0.9956+-0.0004
0			Indel	8	0.9968+-0.0006	0.9923+-0.0014	0.9946+-0.0010
$hg38_asm5_alt$	sentieon-202112.01-recal	octopus-0.7.4	SNP	8	0.9958+-0.0002	0.9844+-0.0012	0.9901+-0.0006
3	- 7701-	•	Indel	8	0.9486+-0.0024	0.9609+-0.0023	0.9547+-0.0022
$hg38_asm5_alt$	sentieon-202112.01-recal	strelka-2.9.10	SNP	8	0.9912+-0.0016	0.9922+-0.0007	0.9917+-0.0010

Reference	Aligner	Caller	Type	N	Precision	Sensitivity	F-measure
			Indel	8	0.9885 + -0.0009	0.9839 + -0.0018	0.9862 + -0.0013
hg38_asm5_alt	sm5_alt snap-2.0.0 clair3-0.1-1	aloir2 0.1 r11	SNP	8	0.9966 + -0.0003	0.9920 + -0.0008	0.9943 + -0.0005
ligoo_asino_an		Claff 5-0.1-111	Indel	8	0.9944 + -0.0006	0.9895 + -0.0015	0.9919 + -0.0010
hg38_asm5_alt	snap-2.0.0	deepvariant-1.2.0	SNP	8	0.9984 + -0.0002	0.9936 + -0.0005	0.9960 + -0.0003
ngoo_asmo_an	Shap-2.0.0		Indel	8	0.9961 + -0.0004	0.9920 + -0.0009	0.9940 + -0.0006
hg38_asm5_alt	snap-2.0.0	dnascope-1.0-202112.01-PO	SNP	8	0.9977 + -0.0002	0.9939 + -0.0006	0.9958 + -0.0004
ngoo_asmo_an	Shap-2.0.0	dnascope-1.0-202112.01-FO	Indel	8	0.9963 + -0.0004	0.9944 + -0.0007	0.9953 + -0.0005
hg38_asm5_alt	gnan 2.0.0	octopus-0.7.4	SNP	8	0.9942 + -0.0003	0.9856 + -0.0011	0.9899 + -0.0006
ngoo_asmo_an	snap-2.0.0	octopus-0.7.4	Indel	8	0.9467 + -0.0026	0.9608 + -0.0023	0.9537 + -0.0023
har 20 sam 5 alt	$sm5$ _alt $snap-2.0.0$ $strelka-2.9$	stroller 2.0.10	SNP	8	0.9852 + -0.0048	0.9935 + -0.0006	0.9894 + -0.0026
ngoo_asmo_an		streika-2.9.10	Indel	8	0.9860 + -0.0009	0.9848+-0.0018	0.9854+-0.0013

${\bf 3.2}\quad {\bf Preparation:~"PacBio-CCS-high"}$

Reference	Aligner	Caller	Туре	N	Precision	Sensitivity	F-measure
hg38_T2T_masked	minimap2-2.23	clair3-0.1-r11	SNP	7	0.9979 + -0.0003	0.9994 + -0.0001	0.9986+-0.0001
118002121211111111111111111111111111111		0.0000	Indel	7	0.9880 + -0.0089	0.9859 + -0.0064	0.9869 + -0.0076
hg38_T2T_masked	minimap2-2.23	dnascope_hifi-0.4-202112.01	SNP	7	0.9993 + -0.0002	0.9993 + -0.0001	0.9993 + -0.0001
iig50_121_iiia5kcd	IIIIIIIIapz-2.23	dnascope_mii-0.4-202112.01	Indel	7	0.9935 + -0.0049	0.9932 + -0.0050	0.9934 + -0.0049
hg38_T2T_masked	minimap2-2.23	peppermargindv-r0.8	SNP	7	0.9983 + -0.0003	0.9990 + -0.0001	0.9986 + -0.0001
lig56_121_lilasked	mmmap2-2.25		Indel	7	0.9886 + -0.0056	0.9858 + -0.0055	0.9872 + -0.0055
hg38_T2T_masked	pbmm2-1.7.0	clair3-0.1-r11	SNP	7	0.9987 + -0.0002	0.9993 + -0.0002	0.9990 + -0.0001
lig56_121_masked	pomm2-1.7.0	Clair 5-0.1-111	Indel	7	0.9903 + -0.0087	0.9913 + -0.0059	0.9908 + -0.0073
hg38_T2T_masked	pbmm2-1.7.0	dnascope_hifi-0.4-202112.01	SNP	7	0.9993 + -0.0003	0.9993 + -0.0001	0.9994 + -0.0001
lig56_121_lilasked	poinin2-1.7.0	dnascope_mii-0.4-202112.01	Indel	7	0.9940 + -0.0046	0.9940 + -0.0045	0.9940 + -0.0046
hg38_T2T_masked	pbmm2-1.7.0	peppermargindv-r0.8	SNP	7	0.9989 + -0.0002	0.9989 + -0.0002	0.9989 + -0.0001
ngoo_121_masked	pomm2-1.7.0	peppermargmuv-10.8	Indel	7	0.9908 + -0.0055	0.9914 + -0.0049	0.9911 + -0.0052
hg38_T2T_masked	sentieon_mm2-202112.01	clair3-0.1-r11	SNP	7	0.9979 + -0.0003	0.9994 + -0.0001	0.9986 + -0.0001
ngoo_121_masked	Sentieon_mm2-202112.01		Indel	7	0.9880 + -0.0089	0.9859 + -0.0064	0.9869 + -0.0076

Reference	Aligner	Caller	Туре	N	Precision	Sensitivity	F-measure
hg38_T2T_masked	sentieon_mm2-202112.01	dnascope_hifi-0.4-202112.01	SNP	7	0.9993 + -0.0002	0.9993 + -0.0001	0.9993+-0.0001
ligoo_121_lilasked	Schricon_mm2-202112.01	dnascope_im-0.4-202112.01	Indel	7	0.9935 + -0.0049	0.9932 + -0.0050	0.9934 + -0.0049
hg38_T2T_masked	sentieon_mm2-202112.01	n ann anns angin des no 0	SNP	7	0.9983 + -0.0003	0.9990 + -0.0001	0.9986 + -0.0001
lig56_121_lilasked	sentieon_mmz-zuz11z.u1	peppermargindv-r0.8	Indel	7	0.9886 + -0.0056	0.9858 + -0.0055	0.9872 + -0.0055

4 CMRG Comparison

GIAB recently released a pre-print describing Challenging Medically-Relevant Autosomal Genes (CMRG) benchmark. These regions are harder to capture than the typical GIAB benchmarks. This section contains the summarized results of the small variant analysis, specifically on the CMRG regions. For each aligner-caller pair, we ran RTG on the CMRG high confidence truth set (currently only for HG002 samples). The following results are gathered from the RTG 'summary.txt' file using the final row (i.e. "None" indicating no variants were filtered) and with the --all-records option enabled.

4.1 Preparation: "PCR-free-illumina"

				Precision	Sensitivity	-measure
Reference	Aligner	Caller	N			<u> </u>
clinical	sentieon-201808.07	strelka-2.9.10	2	0.9542 + -0.0031	0.9512 + -0.0007	0.9527+-0.0019
hg38_T2T_masked	dragmap-1.2.1	clair3-0.1-r11	2	0.9708 + -0.0001	0.9592 + -0.0002	0.9650 + -0.0000
hg38_T2T_masked	dragmap-1.2.1	dnascope-1.0-202112.01-PO	2	0.9753 + -0.0001	0.9668 + -0.0007	0.9710 + -0.0004
hg38_T2T_masked	dragmap-1.2.1	octopus- $0.7.4$	2	0.9804 + -0.0007	0.9634 + -0.0009	0.9718 + -0.0008
hg38_T2T_masked	dragmap-1.2.1	strelka-2.9.10	2	0.9405 + -0.0033	0.9562 + -0.0008	0.9482 + -0.0021
hg38_T2T_masked	sentieon-202112.01	clair3-0.1-r11	2	0.9696 + -0.0004	0.9594 + -0.0010	0.9645 + -0.0004
hg38_T2T_masked	sentieon-202112.01	dnascope-1.0-202112.01-PO	2	0.9750 + -0.0009	0.9661 + -0.0015	0.9706 + -0.0011
hg38_T2T_masked	sentieon-202112.01	octopus-0.7.4	2	0.9734 + -0.0016	0.9640 + -0.0015	0.9687 + -0.0016
hg38_T2T_masked	sentieon-202112.01	strelka-2.9.10	2	0.9357 + -0.0045	0.9608 + -0.0006	0.9481 + -0.0025
hg38_T2T_masked	sentieon-202112.01-recal	clair3-0.1-r11	2	0.9707 + -0.0005	0.9564 + -0.0010	0.9634+-0.0003
hg38_T2T_masked	sentieon-202112.01-recal	dnascope-1.0-202112.01-PO	2	0.9767 + -0.0015	0.9626 + -0.0015	0.9695 + -0.0015
hg38_T2T_masked	sentieon-202112.01-recal	octopus-0.7.4	2	0.9738 + -0.0001	0.9635 + -0.0007	0.9687 + -0.0005
hg38_T2T_masked	sentieon-202112.01-recal	strelka-2.9.10	2	0.9449 + -0.0026	0.9604 + -0.0006	0.9526 + -0.0017
hg38_T2T_masked	snap-2.0.0	clair3-0.1-r11	2	0.9644 + -0.0016	0.9586 + -0.0011	0.9614+-0.0014
hg38_T2T_masked	snap-2.0.0	dnascope-1.0-202112.01-PO	2	0.9704 + -0.0012	0.9654 + -0.0006	0.9679 + -0.0009
hg38_T2T_masked	snap-2.0.0	octopus-0.7.4	2	0.9670 + -0.0018	0.9652 + -0.0007	0.9661 + -0.0005
hg38_T2T_masked	snap-2.0.0	strelka-2.9.10	2	0.9269 + -0.0055	0.9606 + -0.0006	0.9435 + -0.0031
hg38_asm5_alt	dragmap-1.2.1	clair3-0.1-r11	2	0.9750 + -0.0001	0.8495 + -0.0001	0.9080+-0.0001
hg38_asm5_alt	dragmap-1.2.1	deepvariant-1.2.0	2	0.9854 + -0.0003	0.8513 + -0.0006	0.9134+-0.0002
hg38_asm5_alt	dragmap-1.2.1	dnascope-1.0-202112.01-PO	2	0.9780 + -0.0001	0.8656 + -0.0008	0.9183+-0.0004
hg38_asm5_alt	dragmap-1.2.1	octopus-0.7.4	2	0.9747 + -0.0000	0.8539 + -0.0012	0.9103+-0.0007
hg38_asm5_alt	dragmap-1.2.1	strelka-2.9.10	2	0.9430+-0.0029	0.8447+-0.0010	0.8911+-0.0019
hg38_asm5_alt	sentieon-202112.01	clair3-0.1-r11	2	0.9777 + -0.0004	0.9517 + -0.0000	0.9646+-0.0002

Reference	Aligner	Caller	N	Precision	Sensitivity	F-measure
$hg38_asm5_alt$	sentieon-202112.01	dnascope-1.0-202112.01-PO	2	0.9831 + -0.0005	0.9622 + -0.0012	0.9726 + -0.0009
$hg38_asm5_alt$	sentieon-202112.01	octopus- $0.7.4$	2	0.9738 + -0.0010	0.9551 + -0.0014	0.9644+-0.0013
hg38_asm5_alt	sentieon-202112.01	strelka-2.9.10	2	0.9449 + -0.0050	0.9516 + -0.0007	0.9483 + -0.0028
hg38_asm5_alt	sentieon-202112.01-recal	clair3-0.1-r11	2	0.9789 + -0.0002	0.9486 + -0.0001	0.9635 + -0.0002
hg38_asm5_alt	sentieon-202112.01-recal	dnascope-1.0-202112.01-PO	2	0.9847 + -0.0012	0.9589 + -0.0010	0.9717 + -0.0011
hg38_asm5_alt	sentieon-202112.01-recal	octopus- $0.7.4$	2	0.9748 + -0.0000	0.9550 + -0.0013	0.9648 + -0.0007
hg38_asm5_alt	sentieon-202112.01-recal	strelka-2.9.10	2	0.9542 + -0.0031	0.9512 + -0.0007	0.9527 + -0.0019
hg38_asm5_alt	snap-2.0.0	clair3-0.1-r11	2	0.9724 + -0.0010	0.9505 + -0.0005	0.9612 + -0.0007
hg38_asm5_alt	snap-2.0.0	deepvariant-1.2.0	2	0.9869 + -0.0007	0.9533 + -0.0004	0.9698 + -0.0001
hg38_asm5_alt	snap-2.0.0	dnascope-1.0-202112.01-PO	2	0.9802 + -0.0010	0.9583 + -0.0003	0.9691 + -0.0006
hg38_asm5_alt	snap-2.0.0	octopus-0.7.4	2	0.9666 + -0.0018	0.9575 + -0.0011	0.9620 + -0.0003
hg38_asm5_alt	snap-2.0.0	strelka-2.9.10	2	0.9335 + -0.0053	0.9525 + -0.0008	0.9429+-0.0031

4.2 Preparation: "PacBio-CCS-high"

Reference	Aligner	Caller	N	Precision	Sensitivity	F-measure
hg38_T2T_masked	minimap2-2.23	clair3-0.1-r11	1	0.9661 + -0.0000	0.9927 + -0.0000	0.9792 + -0.0000
hg38_T2T_masked	minimap2-2.23	dnascope_hifi-0.4-202112.01	1	0.9876 + -0.0000	0.9894 + -0.0000	0.9885 + -0.0000
hg38_T2T_masked	minimap2-2.23	peppermargindv-r0.8	1	0.9862 + -0.0000	0.9908 + -0.0000	0.9885 + -0.0000
hg38_T2T_masked	pbmm2-1.7.0	clair3-0.1-r11	1	0.9681 + -0.0000	0.9902 + -0.0000	0.9790 + -0.0000
hg38_T2T_masked	pbmm2-1.7.0	dnascope_hifi-0.4-202112.01	1	0.9881+-0.0000	0.9889 + -0.0000	0.9885 + -0.0000
hg38_T2T_masked	pbmm2-1.7.0	peppermargindv-r0.8	1	0.9806 + -0.0000	0.9888+-0.0000	0.9847 + -0.0000
hg38_T2T_masked	sentieon_mm2-202112.01	clair3-0.1-r11	1	0.9661 + -0.0000	0.9928 + -0.0000	0.9792 + -0.0000
hg38_T2T_masked	sentieon_mm2-202112.01	dnascope_hifi-0.4-202112.01	1	0.9876 + -0.0000	0.9894+-0.0000	0.9885 + -0.0000
hg38_T2T_masked	sentieon_mm2-202112.01	peppermargindv-r0.8	1	0.9860 + -0.0000	0.9908+-0.0000	0.9884 + -0.0000

5 Summarized Haplotyper Comparison

This section contains the summarized results of the haplotype analysis. For each aligner-haplotyper pair, we attempted to run the haplotyper and gather the resulting prediction. "Output" is the percentage that generated an output file. This is most commonly below 100% when a tool crashes while trying to run. "Computed" is the percentage that generated a prediction in the output file. This is most commonly below 100% when a tool successfully runs, but it does not generate a prediction in the output file (e.g. "None"). "Matching" is the percentage that generated the correct haplotype prediction based on our lookup table. These haplotyping tools often assume a particular data type, and are not designed to work under all of the conditions tested by this benchmark.

5.1 Preparation: "PCR-free-illumina"

Reference	Aligner	Haplotyper	N	Output (%)	Computed (%)	Matching (%)
hg38_T2T_masked	dragmap-1.2.1	cyrius-1.1.1	8	100.0	75.0	75.0
hg38_T2T_masked	sentieon-202112.01	cyrius-1.1.1	8	100.0	87.5	87.5
hg38_T2T_masked	sentieon-202112.01-recal	cyrius-1.1.1	8	100.0	87.5	87.5
hg38_T2T_masked	snap-2.0.0	cyrius-1.1.1	8	100.0	87.5	87.5
hg38_asm5_alt	dragmap-1.2.1	cyrius-1.1.1	8	25.0	0.0	0.0
hg38_asm5_alt	sentieon-202112.01	cyrius-1.1.1	8	100.0	100.0	100.0
hg38_asm5_alt	sentieon-202112.01-recal	cyrius-1.1.1	8	100.0	100.0	100.0
hg38_asm5_alt	snap-2.0.0	cyrius-1.1.1	8	100.0	87.5	87.5

5.2 Preparation: "PacBio-CCS-high"

Reference	Aligner	Haplotyper	N	Output (%)	Computed (%)	Matching (%)
hg38_T2T_masked	minimap2-2.23	cyrius-1.1.1	5	100.0	20.0	20.0
hg38_T2T_masked	pbmm2-1.7.0	cyrius-1.1.1	5	100.0	40.0	40.0

Reference	Aligner	Haplotyper	N	Output (%)	Computed (%)	Matching (%)
$hg38_T2T_masked$	sentieon_mm2-202112.01	cyrius-1.1.1	5	100.0	20.0	20.0

6 Summarized Deletion Comparisons

This section contains the summarized results of the deletion variant analysis. For each aligner-caller pair, we ran Truvari on the corresponding truth set with these options: "--pctsim 0.0 --sizemax 1000000000". The following results are gathered from the Truvari "summary.txt" file.

6.1 Restricted calls

These evaluations are **RESTRICTED**, meaning that a BED file was used to limit the analysis regions, typically to "high-confidence" regions associated with the benchmark. This corresponds to the "--includebed" option in Truvari. As a result, benchmark sets that were not released with a corresponding BED file are excluded from this analysis. At this time, only the HG002 GIAB v0.6 benchmark is included in this analysis.

6.1.1 Preparation: "PCR-free-illumina"

				ecision	call	
Reference	Aligner	Caller	N	pr	re	fJ
hg38_T2T_masked	dragmap-1.2.1	dysgu-1.3.10-PO	2	0.9016 + -0.0233	0.7096 + -0.0038	0.7940 + -0.0066
hg38_T2T_masked	dragmap-1.2.1	dysgu-1.3.4-PO	2	0.9105 + -0.0212	0.7073 + -0.0035	0.7960 + -0.0059
hg38_T2T_masked	dragmap-1.2.1	manta-1.6.0	2	0.9561 + -0.0011	0.6677 + -0.0188	0.7862 + -0.0127
hg38_T2T_masked	sentieon-202112.01	dysgu-1.3.10-PO	2	0.9019 + -0.0201	0.6656 + -0.0023	0.7658 + -0.0057
hg38_T2T_masked	sentieon-202112.01	dysgu-1.3.4-PO	2	0.9025 + -0.0211	0.6508 + -0.0014	0.7562 + -0.0064
hg38_T2T_masked	sentieon-202112.01	manta-1.6.0	2	0.9529 + -0.0007	0.6909 + -0.0153	0.8009 + -0.0101
hg38_T2T_masked	sentieon-202112.01-recal	dysgu-1.3.10-PO	2	0.8991+-0.0211	0.6657 + -0.0024	0.7649 + -0.0060
hg38_T2T_masked	sentieon-202112.01-recal	dysgu-1.3.4-PO	2	0.9031 + -0.0172	0.6516 + -0.0060	0.7568 + -0.0020
hg38_T2T_masked	sentieon-202112.01-recal	manta-1.6.0	2	0.9518+-0.0009	0.6909 + -0.0149	0.8005 + -0.0097
hg38_T2T_masked	snap-2.0.0	dysgu-1.3.10-PO	2	0.9401+-0.0050	0.6418+-0.0104	0.7628 + -0.0057
hg38_T2T_masked	snap-2.0.0	dysgu-1.3.4-PO	2	0.9420+-0.0040	0.6266 + -0.0043	0.7526 + -0.0018
hg38_T2T_masked	snap-2.0.0	manta-1.6.0	2	0.9490+-0.0008	0.6805 + -0.0128	0.7925 + -0.0084
hg38_asm5_alt	dragmap-1.2.1	dysgu-1.3.10-PO	2	0.9052 + -0.0229	0.6944 + -0.0032	0.7857 + -0.0066
hg38_asm5_alt	dragmap-1.2.1	dysgu-1.3.4-PO	2	0.9146 + -0.0191	0.6927 + -0.0040	0.7882 + -0.0045
hg38_asm5_alt	dragmap-1.2.1	manta-1.6.0	2	0.9616 + -0.0007	0.6550 + -0.0169	0.7791 + -0.0117
hg38_asm5_alt	sentieon-202112.01	dysgu-1.3.10-PO	2	0.9013 + -0.0195	0.6655 + -0.0022	0.7655 + -0.0056
hg38_asm5_alt	sentieon-202112.01	dysgu-1.3.4-PO	2	0.9061 + -0.0184	0.6463 + -0.0002	0.7544 + -0.0062
hg38_asm5_alt	sentieon-202112.01	manta-1.6.0	2	0.9563 + -0.0018	0.6899 + -0.0149	0.8015+-0.0094
hg38_asm5_alt	sentieon-202112.01-recal	dysgu-1.3.10-PO	2	0.8991 + -0.0206	0.6651 + -0.0028	0.7645 + -0.0056
hg38_asm5_alt	sentieon-202112.01-recal	dysgu-1.3.4-PO	2	0.9059 + -0.0176	0.6464 + -0.0042	0.7543 + -0.0032
hg38_asm5_alt	sentieon-202112.01-recal	manta-1.6.0	2	0.9547 + -0.0018	0.6896 + -0.0140	0.8007+-0.0088

Reference	Aligner	Caller	N	precision	recall	11
hg38_asm5_alt	snap-2.0.0	dysgu-1.3.10-PO	2	0.9411+-0.0039	0.6406 + -0.0104	0.7623 + -0.0061
hg38_asm5_alt	snap-2.0.0	dysgu-1.3.4-PO	2	0.9430 + -0.0048	0.6225 + -0.0043	0.7500 + -0.0016
hg38_asm5_alt	snap-2.0.0	manta-1.6.0	2	0.9509 + -0.0022	0.6791 + -0.0127	0.7923 + -0.0079

6.1.2 Preparation: "PacBio-CCS-high"

Reference	Aligner	Caller	N	precision	recall	fl
hg38_T2T_masked	minimap2-2.23	dysgu-1.3.10-PO	1	0.9463 + -0.0000	0.9729 + -0.0000	0.9594+-0.0000
hg38_T2T_masked	minimap2-2.23	dysgu-1.3.4-PO	1	0.9415 + -0.0000	0.9731 + -0.0000	0.9571 + -0.0000
hg38_T2T_masked	minimap2-2.23	pbsv-2.8.0	1	0.9463 + -0.0000	0.9847 + -0.0000	0.9651 + -0.0000
hg38_T2T_masked	minimap2-2.23	sniffles-2.0.2	1	0.9494+-0.0000	0.9808 + -0.0000	0.9649 + -0.0000
hg38_T2T_masked	pbmm2-1.7.0	dysgu-1.3.10-PO	1	0.9372 + -0.0000	0.9729 + -0.0000	0.9547 + -0.0000
hg38_T2T_masked	pbmm2-1.7.0	dysgu-1.3.4-PO	1	0.9554 + -0.0000	0.1849 + -0.0000	0.3098+-0.0000
hg38_T2T_masked	pbmm2-1.7.0	pbsv-2.8.0	1	0.9420 + -0.0000	0.9859 + -0.0000	0.9634 + -0.0000
hg38_T2T_masked	pbmm2-1.7.0	sniffles-2.0.2	1	0.9450 + -0.0000	0.9811 + -0.0000	0.9627 + -0.0000
hg38_T2T_masked	sentieon_mm2-202112.01	dysgu-1.3.10-PO	1	0.9468+-0.0000	0.9736 + -0.0000	0.9600 + -0.0000
hg38_T2T_masked	sentieon_mm2-202112.01	dysgu-1.3.4-PO	1	0.9415 + -0.0000	0.9727 + -0.0000	0.9568 + -0.0000
hg38_T2T_masked	sentieon_mm2-202112.01	pbsv-2.8.0	1	0.9459 + -0.0000	0.9847 + -0.0000	0.9649 + -0.0000
hg38_T2T_masked	sentieon_mm2-202112.01	sniffles-2.0.2	1	0.9492 + -0.0000	0.9808 + -0.0000	0.9647+-0.0000

6.2 Unrestricted calls

These evaluations are **UNRESTRICTED**, meaning that there is no BED file used to limit the analysis to a subset of the genome (e.g. no high-confidence region applied). As a result, precision results are skewed downwards and should be considered a lower-bound on precision with respect to the full benchmark dataset. However, this allows us to add benchmarks that were not released with a BED file, specifically HG001/NA12878's Personalis benchmark set.

6.2.1 Preparation: "PCR-free-illumina"

				l d		
				precision		
				cik	all:	
Reference	Aligner	Caller	N	pre	recall	ĮĮ.
hg38_T2T_masked	dragmap-1.2.1	dysgu-1.3.10-PO	4	0.5638 + -0.0896	0.7866 + -0.1637	0.6354 + -0.0077
hg38_T2T_masked	dragmap-1.2.1	dysgu-1.3.4-PO	4	0.5682 + -0.0907	0.7865 + -0.1645	0.6380 + -0.0072
hg38_T2T_masked	dragmap-1.2.1	manta-1.6.0	4	0.5691 + -0.0968	0.7417 + -0.1500	0.6221 + -0.0078
hg38_T2T_masked	sentieon-202112.01	dysgu-1.3.10-PO	4	0.5717 + -0.0914	0.7482 + -0.1599	0.6260 + -0.0068
hg38_T2T_masked	sentieon-202112.01	dysgu-1.3.4-PO	4	0.5693 + -0.0916	0.7369 + -0.1598	0.6199 + -0.0078
hg38_T2T_masked	sentieon-202112.01	manta-1.6.0	4	0.5404+-0.0990	0.7589 + -0.1444	0.6098 + -0.0175
hg38_T2T_masked	sentieon-202112.01-recal	dysgu-1.3.10-PO	4	0.5714 + -0.0919	0.7485 + -0.1599	0.6258 + -0.0064
hg38_T2T_masked	sentieon-202112.01-recal	dysgu-1.3.4-PO	4	0.5692 + -0.0924	0.7371 + -0.1589	0.6198 + -0.0064
hg38_T2T_masked	sentieon-202112.01-recal	manta-1.6.0	4	0.5392 + -0.0979	0.7588 + -0.1451	0.6092 + -0.0167
hg38_T2T_masked	snap-2.0.0	dysgu-1.3.10-PO	4	0.6112 + -0.0957	0.7082 + -0.1522	0.6336 + -0.0120
hg38_T2T_masked	snap-2.0.0	dysgu-1.3.4-PO	4	0.6057 + -0.0977	0.6900 + -0.1455	0.6229 + -0.0111
hg38_T2T_masked	snap-2.0.0	manta-1.6.0	4	0.4960 + -0.0947	0.7504 + -0.1473	0.5758 + -0.0217
hg38_asm5_alt	dragmap-1.2.1	dysgu-1.3.10-PO	4	0.5743 + -0.0933	0.7661 + -0.1580	0.6348 + -0.0078
hg38_asm5_alt	dragmap-1.2.1	dysgu-1.3.4-PO	4	0.5745 + -0.0922	0.7648 + -0.1575	0.6346 + -0.0062
hg38_asm5_alt	dragmap-1.2.1	manta-1.6.0	4	0.5851 + -0.1027	0.7229 + -0.1443	0.6242 + -0.0073
hg38_asm5_alt	sentieon-202112.01	dysgu-1.3.10-PO	4	0.5686 + -0.0904	0.7475 + -0.1594	0.6241 + -0.0072
hg38_asm5_alt	sentieon-202112.01	dysgu-1.3.4-PO	4	0.5629 + -0.0909	0.7279 + -0.1555	0.6130 + -0.0078
hg38_asm5_alt	sentieon-202112.01	manta-1.6.0	4	0.5502 + -0.1032	0.7558 + -0.1430	0.6148 + -0.0182
hg38_asm5_alt	sentieon-202112.01-recal	dysgu-1.3.10-PO	4	0.5677 + -0.0898	0.7477 + -0.1600	0.6235 + -0.0067
hg38_asm5_alt	sentieon-202112.01-recal	dysgu-1.3.4-PO	4	0.5628 + -0.0919	0.7274 + -0.1549	0.6126 + -0.0057
hg38_asm5_alt	sentieon-202112.01-recal	manta-1.6.0	4	0.5492 + -0.1020	0.7553 + -0.1435	0.6141 + -0.0173
hg38_asm5_alt	snap-2.0.0	dysgu-1.3.10-PO	4	0.6121 + -0.0976	0.7053 + -0.1509	0.6327 + -0.0108
hg38_asm5_alt	snap-2.0.0	dysgu-1.3.4-PO	4	0.6027 + -0.0977	0.6880 + -0.1475	0.6199 + -0.0122
hg38_asm5_alt	snap-2.0.0	manta-1.6.0	4	0.4999 + -0.0970	0.7479 + -0.1467	0.5774 + -0.0223

6.2.2 Preparation: "PacBio-CCS-high"

Reference	Aligner	Caller	N	precision	recall	fì
hg38_T2T_masked	minimap2-2.23	dysgu-1.3.10-PO	2	0.4273 + -0.1364	0.9429+-0.0214	0.5732 + -0.1263
hg38_T2T_masked	minimap2-2.23	dysgu-1.3.4-PO	2	0.4148 + -0.1336	0.9438+-0.0201	0.5618 + -0.1264

Reference	Aligner	Caller	N	precision	recall	17
hg38_T2T_masked	minimap2-2.23	pbsv-2.8.0	2	0.3681+-0.1158	0.9564 + -0.0071	0.5204+-0.1206
hg38_T2T_masked	minimap2-2.23	sniffles-2.0.2	2	0.3764 + -0.1182	0.9473 + -0.0192	0.5265 + -0.1189
hg38_T2T_masked	pbmm2-1.7.0	dysgu-1.3.10-PO	2	0.4160 + -0.1336	0.9440 + -0.0172	0.5632 + -0.1267
hg38_T2T_masked	pbmm2-1.7.0	dysgu-1.3.4-PO	2	0.6125 + -0.0607	0.4066 + -0.2475	0.4280 + -0.1706
hg38_T2T_masked	pbmm2-1.7.0	pbsv-2.8.0	2	0.3813+-0.1210	0.9593 + -0.0087	0.5338+-0.1234
hg38_T2T_masked	pbmm2-1.7.0	sniffles-2.0.2	2	0.3836 + -0.1205	0.9486 + -0.0172	0.5339 + -0.1203
hg38_T2T_masked	sentieon_mm2-202112.01	dysgu-1.3.10-PO	2	0.4275 + -0.1363	0.9434+-0.0216	0.5736 + -0.1261
hg38_T2T_masked	sentieon_mm2-202112.01	dysgu-1.3.4-PO	2	0.4148+-0.1338	0.9434+-0.0201	0.5617 + -0.1265
hg38_T2T_masked	sentieon_mm2-202112.01	pbsv-2.8.0	2	0.3672 + -0.1156	0.9561 + -0.0070	0.5195 + -0.1206
hg38_T2T_masked	sentieon_mm2-202112.01	sniffles-2.0.2	2	0.3761 + -0.1180	0.9473 + -0.0192	0.5262 + -0.1187