

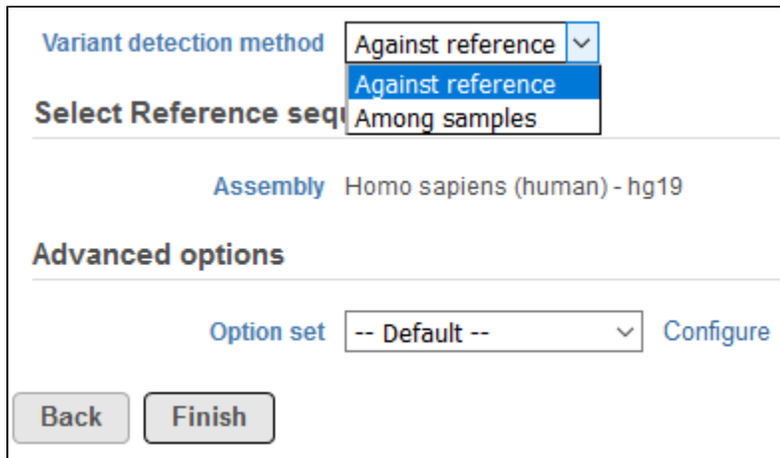
SAMtools

SAMtools¹ (version 1.2) utilizes the mpileup command to look at observed bases in the reads covering every genomic position represented in the aligned sequence data and calculate the likelihood of every possible genotype at a locus. Subsequently, bcftools applies the prior probability and uses Bayesian inference to call actual genotypes, outputting variant information in Variant Call Format (vcf). This method can identify both single nucleotide variants and insertions/deletion events. General information about the underlying algorithm utilized by SAMtools is detailed by Li.^{2,3}

SAMtools dialog

Selecting SAMtools from the context sensitive menu will bring up the SAMtools task dialog, which contains three default sections: *Variant detection method*, *Select Reference sequence*, and *Advanced options*.

In the *Variant detection method* drop-down list, *Against reference* will compare base composition for each sample against the reference sequence assembly, independently (Figure 1).



The image shows a screenshot of the SAMtools task dialog. The 'Variant detection method' dropdown menu is open, showing two options: 'Against reference' (which is highlighted in blue) and 'Among samples'. Below this, the 'Select Reference sequence' section shows 'Assembly Homo sapiens (human) - hg19'. The 'Advanced options' section has an 'Option set' dropdown menu set to '-- Default --' and a 'Configure' button. At the bottom, there are 'Back' and 'Finish' buttons.

Figure 1. Selecting a variant detection method in the Samtools dialog

Selection of *Among samples* (in the absence of sample attributes or when *Paired analysis* is not selected) utilizes information from all samples in the project in a joint variant detection. *Against reference* typically performs well when samples have good coverage and may contain unique SNPs. In instances where there is low coverage, *Among samples* may perform well for identifying SNPs shared between samples (but not for unique SNPs).

In the event paired samples exist within the project, detection *Among samples* can be utilized to identify loci with differing genotypes between the pair once each sample has been compared to the reference sequence assembly. In instances where there is limited information to accurately determine genotypes in one or both of the samples, the same genotype may be called for case and control if it differs from the reference. The [Filter variants](#) task can be used to exclude these spurious loci. To perform this analysis, sample attributes must be added in the [Data tab](#) of the project (Figure 2). Specifically, an attribute must be added for sample ID (shared between the paired samples) and an attribute must also be added for sample type that differentiates the paired samples. Examples of the latter can include case and control or tumor and normal. If these attributes are present, a section for *Analysis options* will be displayed below the Variant detection method (Figure 3). To utilize this feature, select *Paired analysis*. *Match ID* must then be specified and should correspond to the attribute that references the sample ID shared between the pair. Selecting *Case/control* will allow for discriminating genotypes between paired samples in downstream tasks. *Attribute* should correspond to the attribute that defines type within sample pairs, and *Control* can be specified for whatever category relates to the reference sample.

	Sample name	Attributes	
		Sample	Type
1	29a	29	Normal
2	29b	29	Tumor
3	128a	128	Normal
4	128b	128	Tumor
5	129a	129	Normal
6	129b	129	Tumor
7	287a	287	Normal
8	287b	287	Tumor

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Figure 2. Example of attributes required for paired variant analysis

Variant detection method Among samples

Analysis options

Paired analysis

Match ID Sample

Case/control

Attribute Type

Control Normal

Select Reference sequence

Assembly Homo sapiens (human) - hg19

Advanced options

Option set -- Default -- [Configure](#)

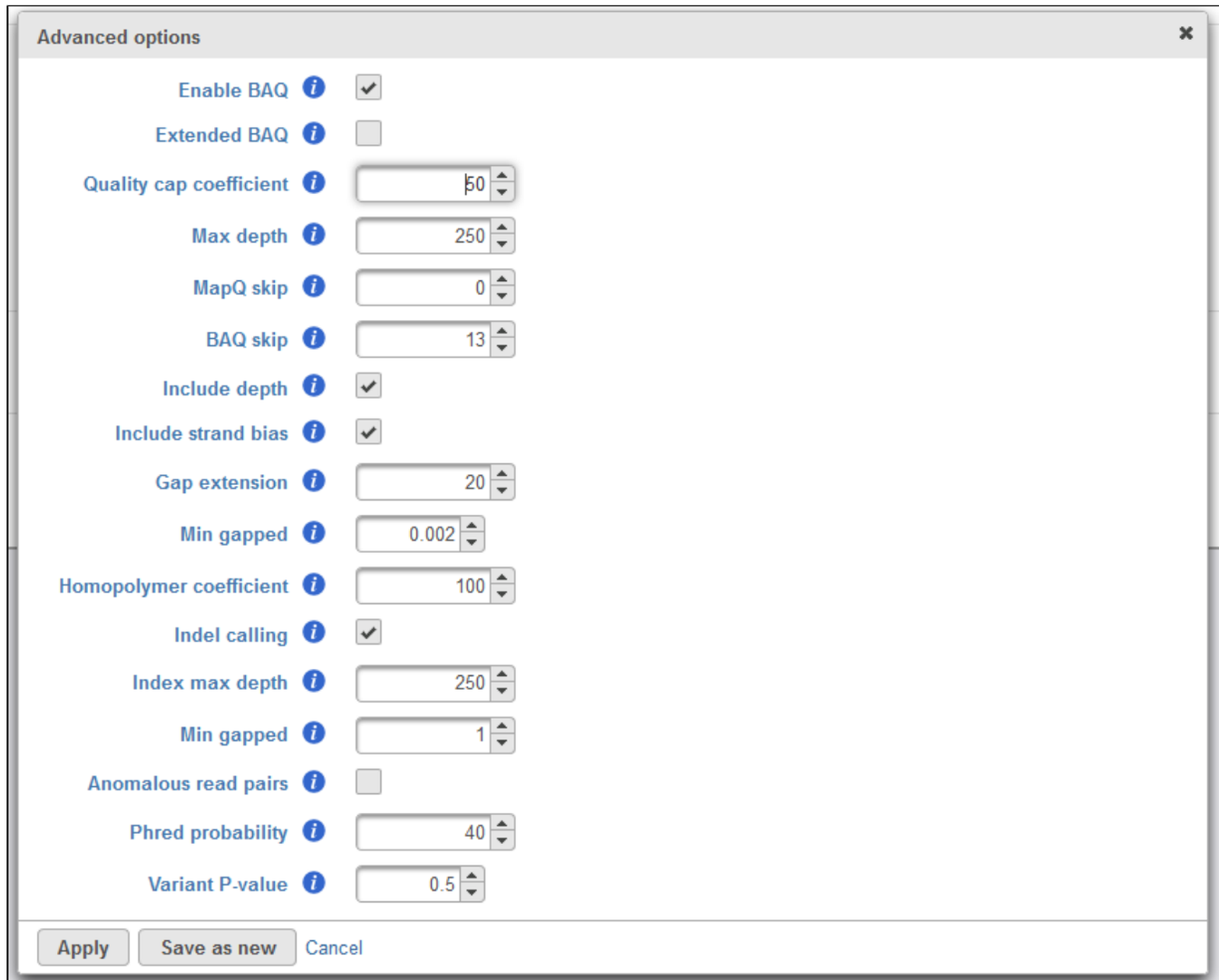
Back
Finish

Figure 3. Specifying options for paired variant detection

Select Reference sequence will specify the reference assembly to utilize for variant detection. If the alignment was generated in Partek® Flow®, the *Assembly* will be displayed as text in the section, and you do not have the option to change the reference. In the event that alignment was performed outside of Partek Flow, you will need to select the appropriate *Assembly* utilized for alignment in the drop-down list. Assemblies previously added to library files (see [Library File Management](#)) will be available for selection or *New assembly...* can be utilized to import the reference sequence to library files from within the task.

Advanced options provides a means to tune parameters in the variant detection for optimal performance. Upon invoking the task dialog, *Option set* is set to *Default*, and these parameters are provided by the SAMtools developers. Clicking *Configure* will open a window to tune advanced options (Figure 3).

Moving the mouse cursor over the info button  will provide details for each parameter. Please refer to the [SAMtools documentation](#) for further details on any of these parameters.



Parameter	Value
Enable BAQ	<input checked="" type="checkbox"/>
Extended BAQ	<input type="checkbox"/>
Quality cap coefficient	50
Max depth	250
MapQ skip	0
BAQ skip	13
Include depth	<input checked="" type="checkbox"/>
Include strand bias	<input checked="" type="checkbox"/>
Gap extension	20
Min gapped	0.002
Homopolymer coefficient	100
Indel calling	<input checked="" type="checkbox"/>
Index max depth	250
Min gapped	1
Anomalous read pairs	<input type="checkbox"/>
Phred probability	40
Variant P-value	0.5

Figure 4. Configuring advanced Samtools options

References

1. Li H, Handsaker B, Wysoker A, et al. The Sequence Alignment/Map format and SAMtools. *Bioinforma Oxf Engl*. 2009;25(16):2078-2079.
2. Li H. A statistical framework for SNP calling, mutation discovery, association mapping and population genetical parameter estimation from sequencing data. *Bioinformatics*. 2011;27(21):2987-2993.
3. Li H. Improving SNP discovery by base alignment quality. *Bioinformatics*. 2011;27(8):1157-1158.

Additional Assistance

If you need additional assistance, please visit [our support page](#) to submit a help ticket or find phone numbers for regional support.



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