Map Reads with Bowtie2 Element

Performs alignment of short reads with Bowtie2.

Element type: align-reads-with-bowtie2

Parameters in GUI

Parameter	Description	Default value	Parameter in Workflow File	Туре
Output directory	Directory to save Bowtie2 output files.		output-dir	string
Reference genome	Path to an indexed reference genome.		reference	string
Output file name	Base name of the output file. 'out.sam' by default.	out.sam	outname	string
Library	Is this library mate-paired?	single-end	library	string
Mode	When the -n option is specified (which is the default), bowtie determines which alignments are valid according to the following policy, which is similar to Maq's default policy. In -v mode, alignments may have no more than V mismatches, where V may be a number from 0 through 3 set using the -v option. Quality values are ignored. The -v option is mutually exclusive with the -n option.	end-to- end	mode	string
Number of mismatches	Sets the number of mismatches to allowed in a seed alignment. Can be set to 0 or 1. Setting this higher makes alignment slower (often much slower) but increases sensitivity.	0	mismatches_ number	numerio
Seed length (L)	Sets the length of the seed substrings to align. Smaller values make alignment slower but more senstive.	20	seed_len	numerio
Add columns to allow gaps (dpad)	"Pads" dynamic programming problems by the specified number of columns on either side to allow gaps.	15	dpad	numerio
Disallow gaps (gbar)	Disallow gaps within a specified number of positions of the beginning or end of the read.	4	gbar	numerio
Seed (seed)	Use as the seed for pseudo-random number generator.	0	seed	numerio
Threads	Launch specified number of parallel search threads. Threads will run on separate processors/cores and synchronize when parsing reads and outputting alignments. Searching for alignments is highly parallel, and speedup is close to linear.	1	threads	numerio
No unpaired alignments (no-mixed)	If Bowtie2 cannot find a paired-end alignment for a pair, by default it will go on to look for unpaired alignments for the constituent mates. This is called "mixed mode." To disable mixed mode, set this option. Bowtie2 runs a little faster in the mixed mode, but will only consider the alignment status of pairs per se, not individual mates.	False	nomixed	boolear
No discordant alignments (no- discordant)	By default, Bowtie2 looks for discordant alignments if it cannot find any concordant alignments. A discordant alignment is an alignment where both mates align uniquely, but that does not satisfy the paired-end constraints. This option disables that behavior.	False	nodiscordant	boolear
No forward orientation (nofw)	Ifnofw is specified, bowtie will not attempt to align against the forward reference strand.	False	nofw	boolear
No reverse- complement orientation (norc)	Ifnorc is specified, bowtie will not attempt to align against the reverse-complement reference strand.	False	norc	boolear
No overlapping mates (no- overlap)	If one mate alignment overlaps the other at all, consider that to be non-concordant. Default: mates can overlap in a concordant alignment.	False	nooverlap	boolear
No mates containing one another (no-contain)	If one mate alignment contains the other, consider that to be non-concordant. Default: a mate can contain the other in a concordant alignment.	False	nocontain	boolear

Input/Output Ports

The element has 1 input port.

Name in GUI: Bowtie2 data

Name in Workflow File: in-data

Slots:

Slot In GUI	Slot in Workflow File	Туре
URL of a file with mate reads	readsurl	string
URL of a file with reads	readspairedurl	string

And 1 output port:

Name in GUI: Bowtie2 output data

Name in Workflow File: out-data

Slots:

Slot In GUI	Slot in Workflow File	Туре
Assembly URL	assembly-out	string