

Slopbed Element

Increases the size of each feature in files using bedtools slop.

Parameters in GUI

| Parameter | Description | Default value |
|--------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------|
| Output directory | Select an output directory. Custom - specify the output directory in the 'Custom directory' parameter. Workflow - internal workflow directory. Input file - the directory of the input file. | Input file |
| Custom directory | Specify the output directory. | |
| Output file name | A name of an output file. If default of empty value is provided the output name is the name of the first file with additional extention. | |
| Genome | In order to prevent the extension of intervals beyond chromosome boundaries, bedtools slop requires a genome file defining the length of each chromosome or contig (-g). | human.hg18 |
| Each direction increase | Increase the BED/GFF/VCF entry by the same number base pairs in each direction. If this parameter is used -l and -l are ignored. Enter 0 to disable (-b). | 0 |
| Subtract from start | The number of base pairs to subtract from the start coordinate. Enter 0 to disable (-l). | 0 |
| Add to end | The number of base pairs to add to the end coordinate. Enter 0 to disable (-r). | 0 |
| Strand-based | Define -l and -r based on strand. For example. if used, -l 500 for a negative-stranded feature, it will add 500 bp to the end coordinate (-s). | False |
| As fraction | Define -l and -r as a fraction of the feature's length. E.g. if used on a 1000bp feature, -l 0.50, will add 500 bp upstream (-pct). | False |
| Print header | Print the header from the input file prior to results (-header). | False |

Parameters in Workflow File

Type: sloped

| Parameter | Parameter in the GUI | Type |
|-------------------|--------------------------------|----------------|
| out-mode | Output directory | <i>numeric</i> |
| custom-dir | Custom directory | <i>string</i> |
| out-name | Output file name | <i>string</i> |
| genome-id | Genome | <i>string</i> |
| b-id | Each direction increase | <i>numeric</i> |
| l-id | Subtract from start | <i>numeric</i> |
| r-id | Add to end | <i>numeric</i> |
| s-id | Strand-based | <i>boolean</i> |
| pct-id | As fraction | <i>boolean</i> |
| header-id | Print header | <i>boolean</i> |

Input/Output Ports

The element has 1 *input port*:

Name in GUI: Input File

Name in Workflow File: in-file

Slots:

| Slot In GUI | Slot in Workflow File | Type |
|-------------|-----------------------|---------------|
| Source URL | url | <i>string</i> |

And 1 *output port*:

Name in GUI: Output File

Name in Workflow File: out-file

Slots:

| Slot In GUI | Slot in Workflow File | Type |
|-------------|-----------------------|---------------|
| Source URL | url | <i>string</i> |