

Get Sequences by Annotations Element

Extracts annotated regions from the input sequence.

Element type: extract-annotated-sequence

Parameters

Parameter	Description	Default value	Parameter in Workflow File	Type
Translate	Translates the annotated regions if the corresponding annotation marks a protein subsequence.	False	translate	<i>boolean</i>
Complement	Complements the annotated regions if the corresponding annotation is located on the complement strand.	False	complement	<i>boolean</i>
Split joined	Split joined annotations to single region annotations.	False	split-joined-annotations	<i>boolean</i>
Extend left	Extends the resulted regions to left.	0	extend-left	<i>numeric</i>
Extend right	Extends the resulted regions to right.	0	extend-right	<i>numeric</i>
Gap length	Inserts a gap of a specified length between the merged locations of the annotation.	0	merge-gap-length	<i>numeric</i>

Input/Output Ports

The element has 1 *input port*:

Name in GUI: *Input sequence*

Name in Workflow File: in-sequence

Slots:

Slot In GUI	Slot in Workflow File	Type
Sequence	sequence	<i>sequence</i>
Set of annotations	annotations	<i>annotation-table</i>

And 1 *output port*:

Name in GUI: *Annotated regions*

Name in Workflow File: out-sequence

Slots:

Slot In GUI	Slot in Workflow File	Type
Sequence	sequence	<i>sequence</i>