

Read Sequence from Remote Database Element

Download sequence(s) with the specified ID(s) from one of the remote databases: NCBI, Ensembl, PDB, etc.

The sequences are downloaded with the associated annotations in a file format, specific for the selected database.

The element outputs message(s) with the sequence and annotations data.

Element type: fetch-sequence

Parameters

Parameter	Description	Default value	Parameter in Workflow File	Type
Resource IDs (required)	Semicolon-separated list of resource IDs in the database.		resource-id	<i>string</i>
Database (required)	Name of the database to read from.	NCBI Genbank (DNA sequence)	database	<i>string</i> Available values are: <ul style="list-style-type: none">• ncbi-dna (NCBI GenBank (DNA sequence))• ncbi-protein (NCBI protein sequence database)• pdb (PDB)• swiss-plot (SWISS-PROT)• uniprot-swiss-prot (UniProtKB /Swiss-Prot)• uniprot-trembl (UniProtKB /TrEMBL)
Save file to directory	Directory to store a file loaded from the database.	default	save-dir	<i>string</i>
Read resource ID(s) from source	The source to read resource IDs from the list or a local file.	List of TDs	ids-source	<i>string</i>

Input/Output Ports

The element has 1 *output port*:

Name in GUI: *Sequence*

Name in Workflow File: out-sequence

Slots:

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