

# Fetch Sequences by ID From Annotation Element

Parses annotations to find any IDs and fetches corresponding sequences.

## Parameters in GUI

| Parameter              | Description  | Default value |
|------------------------|--|---------------|
| Save file to directory | The directory to store sequence files loaded from a database.  | default       |
| NCBI database          | The database to read from.<br><br>Available values are: <ul style="list-style-type: none"><li>• nucleotide</li><li>• protein</li></ul> | nucleotide    |

## Parameters in Workflow File

Type: fetch-sequence

| Parameter | Parameter in the GUI   | Type   |
|-----------|------------------------|--------|
| save-dir  | Save file to directory | string |
| database  | NCBI database          | string |

The element has 1 *input port*:

Name in GUI: *Input annotations*

Name in Workflow File: in-annotations

Slots:

| Slot In GUI        | Slot in Workflow File | Type             |
|--------------------|-----------------------|------------------|
| Set of annotations | annotations           | annotation-table |

And 1 *output port*:

Name in GUI: *Sequence*

Name in Workflow File: out-sequence

Slots:

| Slot In GUI        | Slot in Workflow File | Type             |
|--------------------|-----------------------|------------------|
| Set of annotations | annotations           | annotation-table |
| Sequence           | sequence              | sequence         |