

NLMJournalFetch

Overview

This tool is used to ingest data from the [National Library of Medicine's catalog](#) using their [EUtils web service](#). PubMed allows queries to be designed to return very specific record sets based on a range of different attributes such as date added, date modified, number range, affiliation, etc. See their [help for using the NLM Catalog search tool](#) and also the [advanced search](#) tool, which helps you build your query.

The catalog records that match the query given to the tool will be fetched as XML data and loaded into the output [RecordHandler](#).

Setup

Command Line Arguments

NLMJournalFetch extends [NIHFetch](#), so it uses the same arguments.

Short Option	Long Option	Parameter Value Map	Description	Required
b	batchSize	NUMBER	number of records to fetch per batch	true
m	email	EMAIL_ADDRESS	your contact email address	true
n	numRecords	NUMBER	maximum records to return	true
o	output	CONFIG_FILE	RecordHandler config file path	true
O	outputOverride	VALUE	override the RH_PARAM of output recordhandler using VALUE	false
t	termSearch	SEARCH_STRING	term to search against nlmcatalog	true

Configuration File

As with all the [Harvester command-line tools](#), you can provide all the arguments as parameters in a configuration file (`XML-config`). Here is a sample configuration for NLMJournalFetch.

```
<?xml version="1.0" encoding="UTF-8"?>
<Task>
    <Param name="email">sample.email@mydomain.tld</Param>
    <Param name="termSearch">sample AND edu[ad]</Param>
    <!-- these are set inside the example scripts, so are not needed
    <Param name="output">config/recordhandlers/tfrh.xml</Param>
    <Param name="outputOverride">fileDir=harvested-data/examples/nlmcatalog</Param>
    -->
    <Param name="numRecords">ALL</Param>
    <Param name="batchSize">1000</Param>
</Task>
```

Execution

To execute the NLMJournalFetch tool from the commandline, there is a convenient environment config file that, when loaded in a bash shell, will allow you to execute NLMJournalFetch with a simple `$NLMJournalFetch [args]`.

Or you can simply call (paths relative to base harvester folder):

```
java -cp bin/harvester-<version>.jar:bin/dependency/* -Dprocess-task=PubmedFetch org.vivoweb.harvester.fetch.
NLMJournalFetch
```

Design

See [Design of NLMJournalFetch](#) and its [javadoc page](#)