

# *ontoCAT*: package for ontology parsing

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20 September, 2010

Version 0.0.1. released 20 September, 2010.

## 1 Introduction

The *ontoCAT* package provides a simple interface to the Experimental Factor Ontology (<http://www.ebi.ac.uk/efo>) and to any other ontology described in OWL or OBO format.

Package can load the ontology from a local file or on the fly from a URL and internally create the inferred ontology view. Experimental Factor Ontology (EFO) is the default ontology, loaded from: [http://efo.svn.sourceforge.net/viewvc/efo/trunk/src/efoinowl/InferredEFOOWLview/EFO\\_inferred.owl](http://efo.svn.sourceforge.net/viewvc/efo/trunk/src/efoinowl/InferredEFOOWLview/EFO_inferred.owl). The package's methods allow to parse an ontology, search terms in it, find out term parents and children. The package is based on the Ontology Common API Tasks Java library (<http://www.ontocat.org>) as well as various other utilites methods and depends on *rJava* R package.

There are two classes in the *ontoCAT* package:

- *OntologyParser* for parsing and representation of the ontology hierarchy.
- *OntologyTerm* for external view of an ontology term in the hierarchy.

## 2 Functions

The *ontoCAT* package provides a number of functions to work with ontologies in R.

### 2.1 Create Ontology Parser

To create an object of *OntologyParser* you can use one of the two methods:

- `getEFOParser()` loads the latest EFO version on the fly from the EFO SVN repository and creates *OntologyParser* object.
- `getOntologyParser("pathToOntology")` loads the ontology described in OWL or OBO format from a local file or a URL and creates *OntologyParser* object.

```
> library(ontoCAT)
> efo <- getEFOParser()
> biotop <- getOntologyParser("http://purl.org/biotop/biotop.owl#Disposition")
```

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## 2.2 Ontology Parsing

- To find all ontology terms two functions can be used: `getAllTerms` - returns a list of *OntologyTerm* objects. In turn, `getAllTermIds` - returns a list of term accessions.

```
> getAllTerms(biotop)
> getAllTermIds(efo)
```

- Function `getTermById` returns the accession number of the term. In turn, `getTermNameById` returns the name of the term.

```
> term <- getTermById(efo, "EFO_0000322")
> term_biotop <- getTermById(biotop, "DeadBody")
> getTermNameById(efo, "EFO_0000311")
> getTermNameById(biotop, "EmbryonicStructure")
```

- To find out all term parents or children the following functions can be used.

```
> getAllTermParents(efo, "EFO_0000322")
> getAllTermChildren(efo, "EFO_0000322")
> getAllTermParents(biotop, "DeadBody")
> getAllTermChildren(biotop, "DeadBody")
```

Arguments: appropriate `OntologyParser` and the term accession.

- To find out only direct parents or children of the term functions `getTermParents` or `getTermChildren` can be used.

```
> getTermParents(efo, "EFO_0000322")
> getTermChildren(efo, "EFO_0000322")
> getTermParents(biotop, "DeadBody")
> getTermChildren(biotop, "DeadBody")
```

Arguments: appropriate `OntologyParser` and the term accession.

- One more function to get term children together with the queried term accession is `getTermAndAllChildrenIds`.

```
> getTermAndAllChildrenIds(efo, "EFO_0000322")
> getTermAndAllChildrenIds(biotop, "DeadBody")
```

Arguments: appropriate `OntologyParser` and the term accession.

- To create a flat subtree representation of the ontology "opened" down to the specified term function `getTreeDownTo` can be used. All possible paths from the root will be returned.

```
> getTreeDownTo(efo, "EFO_0000322")
```

Arguments: appropriate `OntologyParser` and the term accession.

- A few simple functions allow to get term definition and synonyms:

```
> getDefinitions(efo, "EFO_0000322")
> getSynonyms(efo, "EFO_0000322")
```

Arguments: appropriate `OntologyParser` and the term accession.

- A few simple functions allow to get some metadata about the used ontology:

```
> getOntologyAccession(efo)
> getOntologyDescription(efo)
```

Arguments: appropriate `OntologyParser`.

- To check if the term is present in the ontology function `hasTerm` can be used.

```
> hasTerm(efo, "CL000023")
> hasTerm(efo, "EFO_0000322")
```

Arguments: appropriate `OntologyParser` and the term accession.

- The following functions can be used to search terms in the ontology:

```
> searchTerm(efo, "thymus")
> searchTermPrefix(efo, "thym")
```

Arguments: appropriate `OntologyParser` and stringstring prefix to search for.

- The following functions can be used to investigate the ontology hierarchy:

```
> isRoot(efo, "EFO_0000322")
> getRoots(efo)
> getRootIds(efo)
```

For some ontologies these functions might fail when the ontology used was not design to have root classes.

## 2.3 Functions Specific for EFO

There are few functions specific for EFO class hierarchy to work with EFO branch roots.

```
> getEFOBranchRootIds(efo)
> isEFOBranchRoot(efo, "EFO_0000322")
```

## 2.4 Ontology Term

There are only three functions for the `OntologyTerm` class.

- `getLabel` to get term name;
- `getAccession` to get term accession;
- `show` to view the term.

```
> term <- getTermById(efo, "EFO_0000322")
> getLabel(term)
> getAccession(term)
> term
```

## References

1. Adamusiak T, Burdett T, van der Velde K J, Abeygunawardena N, Antonakaki D, Parkinson H and Swertz M: OntoCAT – a simpler way to access ontology resources. *Available from Nature Precedings* <http://dx.doi.org/10.1038/npre.2010.4666.1> (2010)
2. Malone J, Holloway E, Adamusiak T, Kapushesky M, Zheng J, Kolesnikov N, Zhukova A, Brazma A, Parkinson H: Modeling Sample Variables with an Experimental Factor Ontology. *Bioinformatics* 2010, **26**(8):1112–1118
3. Experimental Factor Ontology <http://www.ebi.ac.uk/efo>
4. Ontology Common API Tasks java library <http://www.ontocat.org>
5. Java sources and javadocs: <http://sourceforge.net/projects/ontocat/files/>