RpepXML: an R interface to the pepXML format for peptide identification.

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Abstract

This vignette describes the data structures and associated methods and functions to import and inspect peptide identification data in the pepXML format.

Keywords: mass spectrometry, proteomics, MSMS, identification, XML.

1. Introduction

From the pepXML webpage¹:

pepXML is an open data format developed at the SPC/Institute for Systems biology² for the storage, exchange, and processing of peptide sequence assignments of MS/MS scans. pepXML is intended to provide a common data output format for many different MS/MS search engines and subsequent peptide-level analyses. Several search engines already have native support for outputting pepXML and converters are available to transform output files to pepXML.

Note that the HUPO Proteomics Standards Initiative³ (PSI) has also develop an exchange standard for database search results, called mzIdentXML⁴. It is not yet as widely used as pepXML but should supersede the latter. A RmzIdentML library will be developed at a later stage.

Currently, only the generic MSMS identification results of the msms_pipeline_analysis is implemented. This package has currently been tested and developed around Mascot⁵ search results.

2. Getting data into R

The functionality of RpepXML will be illustrated with a dummy example file that is provided with the package. It is parsed with the parseMSMSpepXML function, that takes a file name as first parameter. The second one defined whether a progress bar should be displayed. This function returns an object of class MSMSpepXML.

```
> library("RpepXML")
```

```
> pepxml <- system.file("extdata", "tinySearch.pepXML", package = "RpepXML")</pre>
```

```
> ms <- parseMSMSpepXML(pepfile = pepxml, verbose = FALSE)</pre>
```

> class(ms)

²http://tools.proteomecenter.org/wiki/index.php?title=Main_Page

```
<sup>3</sup>http://www.psidev.info/
```

```
<sup>4</sup>http://www.psidev.info/index.php?q=node/319
```

```
<sup>5</sup>http://www.matrixscience.com/
```

¹http://tools.proteomecenter.org/wiki/index.php?title=Formats:pepXML

[1] "MSMSpepXML" attr(,"package") [1] "RpepXML"

3. Data structures and manipulation

The classes implemented mimic a simplified version of the XML structure for the $msms_pipeline_analysis$ element (see the (older) docs⁶, for a browsable description). The structure of the 4 classes is described below, starting with the main, high-level, data structure. Each class and slots are described in the respective on-line manuals.

The classes are versioned individually and the class version can be retrieved with classVersion().

3.1. MSMSpepXML

> getClass("MSMSpepXML")

```
Class "MSMSpepXML" [package "RpepXML"]
```

Slots:

Name:	pepFile	searchEngine	sampleEnzyme	searchDatabase
Class:	character	character	character	character
Name: Class:	spectrumQueries list	classVersion Versions		

Extends: "Versioned"

MSMSpepXML is the main class that holds general meta-data about the search like the search engine, the sample enzyme and the original database used to query the MSMS spectra. The name of the pepXML file also recorded. All the spectrum queries are stored as a list.

> ms

```
Object of class MSMSpepXML
File: tinySearch.pepXML
Database: erwinia_pro.fasta
Engine: MASCOT
Enzyme: Trypsin
4 spectrum queries
4 peptides identified
```

> pepFile(ms)

[1] "/tmp/RtmpD62wid/Rinstf968a17/RpepXML/extdata/tinySearch.pepXML"

```
> searchEngine(ms)
```

```
[1] "MASCOT"
```

⁶http://sashimi.sourceforge.net/schema_revision/pepXML/Docs/pepXML_v18.html

```
> sampleEnzyme(ms)
```

[1] "Trypsin"

> searchDatabase(ms)

[1] "erwinia_pro.fasta"

The spectrum queries can be accessed using the **spectrumQueries** method or the [operator. The former returns a list that can be subsetted as usual. The latter return a list when multiple spectra are selected or and object of class **SpectrumQuery** if one query is provided.

```
> class(spectrumQueries(ms))
```

[1] "list"

```
> sq <- ms[3]
> class(sq)
```

[1] "SpectrumQuery" attr(,"package") [1] "RpepXML"

3.2. SpectrumQuery

```
> getClass("SpectrumQuery")
```

Class "SpectrumQuery" [package "RpepXML"]

Slots:

Name:	searchResults	spectrumId	startScan	endScan
Class:	list	character	integer	integer
			0	0
Name:	precNeutralMass	assumedCharge	queryIndex	classVersion
Class:	numeric	numeric	integer	Versions
			9	

Extends: "Versioned"

SpectrumQuery objects hold the information about specific spectra that have been queried. Note that all pieces of information may not be provided in the pepXML file.

> sq

```
Object of class SpectrumQuery
Spectrum id: 186_c660.3871_u662.3625_r58.85
Start - end scans: 0 - 0
Precursor neutral mass: 659.3798
Assumed charge: 1
Query index: 3
2 search result(s)
3 search hits(s)
6 proteins(s)
```

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```
> precNeutralMass(sq)
```

[1] 659.3798

```
> assumedCharge(sq)
```

[1] 1

4

```
> spectrumId(sq)
```

[1] "186_c660.3871_u662.3625_r58.85"

If any, the search results can be accessed with the **searchResuts** or [operator, that works as described above.

> sr <- sq[2]

3.3. SearchResult

```
> getClass("SearchResult")
```

Class "SearchResult" [package "RpepXML"]

Slots:

Name:	searchHits	searchId	classVersion
Class:	list	integer	Versions

Extends: "Versioned"

The SearchResult class holds the information about a specific search result and the hits that have been identified are available as a list of SearchHit objects. The latter can be accessed by the searchHits method or the [operator.

> sr

```
Object of class SearchResult
Search id: 2
2 search hit(s)
```

> sh <- sr[1]

3.4. SearchHit

> getClass("SearchHit")
Class "SearchHit" [package "RpepXML"]
Slots:
Name: hitRank proteins

ns pepSequence

modifications

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Class:	integer	character	character	data.frame
Name:	scores0	lassVersion		
Class:	numeric	Versions		

```
Extends: "Versioned"
```

The actual results are stored as individual SearchHit instances. The class records the rank of the hit, the peptide sequence with the previous and next amino acids, possible modifications, associated proteins and scores associated to the hit. The scores are returned as a named vector to retrieve individual scores. The ionScore method allows to extract specifically the ion score for Mascot results.

> sh

```
Object of class SearchHit
 Rank: 1
 Sequence: K.TPVSEK.V
 Scores: 34.67 27 22 0.0089
 1 proteins: BSA
 0 modifications
> hitRank(sh)
[1] 1
> pepSequence(sh)
[1] "K.TPVSEK.V"
> scores(sh)
     ionscore identityscore homologyscore
                                                   expect
      34.6700
                    27.0000
                                   22.0000
                                                   0.0089
> ionScore(sh)
ionscore
  34.67
> proteins(sh)
[1] "BSA"
```

4. Getting a general overview of the data

The number of spectrum queries and search results or hits can be extracted with the nSpectrum-Queries, nSearchResults and nSearchhits methods. Several methods can also be applied to parent classes to those that actually have the slot to retrieve the slot values from the whole set of objects.

> nSearchResults(sq)

```
[1] 2
```

6

proteinSummary and proteinSummary can be called on any of the classes to retrieve the number of inferred proteins and a table summary.

The filterHits method can filter any of the classes based on the ion score and the rank of a search hit. Except when filtering a SearchHit, where TRUE or FALSE is returned, depending if the hit passes the filtering criteria, filterHits returns an object of the same class.

```
> msf <- filterHits(ms, rank = 2, ionscore = 20)
> class(msf)
[1] "MSMSpepXML"
attr(,"package")
[1] "RpepXML"
> proteins(ms[3])
[1] "ECA0851" "ECA3748" "BSA" "ECA0851" "ECA3174" "ECA3748"
> proteins(msf[3])
```

[1] "BSA"

An MSMSpepXML object can also be converted to a matrix with as.matrix. Only first rank hits and the first search results are considered and partial information is returned. The all parameter defined whether all spectrum queries should be returned or only those with search results.

```
> as.matrix(msf, all = TRUE)
```

	QueryIndex	${\tt SpectrumId}$			ionscore	identityscore
1	"1"	"679_c403.2	119_u403.2178	_r63.12"		
2	"2"	"511_c402.24	474_u402.2695	_r41.13"		
3	"3"	"186_c660.38	.3625.3625	_r58.85"	"34.67"	"27"
4	"4"	"40_c402.76	15_u402.7805_1	r41.44"		
	homologysco	ore expect	Sequence	NumProte	ins Prote	eins
1				"0"		
2				"0"		
3	"22"	"0.0089"	"K.TPVSEK.V"	"1"	"BSA'	ı
4				"0"		

5. More details

A full list of functions and methods can be obtained with help(package="RpepXML"). Methods, functions and classes are documented individually.

6. Session information

- R version 2.13.0 Under development (unstable) (2010-10-09 r53266), x86_64-unknown-linux-gnu
- Locale: LC_CTYPE=en_GB.utf8, LC_NUMERIC=C, LC_TIME=en_GB.utf8, LC_COLLATE=C, LC_MONETARY=C, LC_MESSAGES=en_GB.utf8, LC_PAPER=en_GB.utf8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_GB.utf8, LC_IDENTIFICATION=C
- Base packages: base, datasets, grDevices, graphics, methods, stats, utils
- Other packages: Biobase².9.2, RpepXML^{0.3.3}, XML^{3.2-0}, plyr^{1.2.1}
- Loaded via a namespace (and not attached): tools².13.0