# Reading PSI-25 XML file from IntAct with the *Rintact* package

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#### Abstract

This document serves as a user's vignette to the R package *Rintact*. We present examples of how to use the two main function's of *Rintact*, and also how to take the output data from these two functions and create the input data for statistical methods in proteomic analysis provided by other Bioconductor packages.

### 1 Introduction

Rintact is an R package mainly used to parse the PSI-25 files generated by the IntAct data repository which collects, curates and stores thousands of protein interactions. Currently, there are two main functions within Rintact:

- 1. psi25interaction
- 2. psi25complex

The first function, psi25interaction, takes either a PSI-25 XML file from IntAct or an URL containing the web address of where such an XML file can be obtained. The XML file must contain binary protein protein interaction data. Example for such data are direct physical interactions, complex co-membership, synthetic genetic interactions. The second function, psi25complex, also takes a PSI-25 XML file or URL as an input parameter, but the file must contain protein complex membership information. In principle, these two functions can take any XML file which adheres to the PSI-25 standards. We have constructed these functions, however, to work primarily with the IntAct PSI-25 XML files, as there are subtle implementation differences between repositories such as IntAct and DIP, although both use the PSI-25 standards. In this vignette, we shall demonstrate the use of these functions on the data generated by [Ewing et al.(2007)] and the manually curated protein complexes derived by the IntAct curators.

## 1.1 Loading R Libraries

We begin by loading the various R libraries with which we shall use. Our primary focus will be with the *Rintact* package, but we will also examine and exploit statistical methods found in various Bioconductor packages for the analysis of the interaction data obtain from *IntAct*.

```
> library("Rintact")
> library("graph")
> library("Rgraphviz")
> #library("ppiStats")
> library("RBGL")
> library("apComplex")
> library("xtable")
```

# 2 Obtaining the Interaction Information

#### 2.1 psi25interaction

We first demonstrate the use of the function psi25interaction. We can either download the *IntAct* PSI-25 XML file onto a local directory or we can simply use the URL (of where the file can be obtained) as the input parameter. We have chosen the latter:

```
> url <- system.file("PSI25XML", "interactionSample.xml", package="Rintact")
> ewing <- psi25interaction(url)</pre>
```

Once the XML file has been parsed by psi25interaction, we can look at its overall structure.

```
> class(ewing)
[1] "interactionEntry"
attr(,"package")
[1] "Rintact"
```

We can see that the output of psi25interaction is an instance of the class *interactionEntry*. This class has 5 slots:

```
> slotNames(ewing)
[1] "organismName" "taxId" "releaseDate" "interactors" "interactions"
```

Three of them contain simple character vectors:

```
> ewing@organismName
[1] "Homo sapiens" "Human adenovirus E"
[3] "Human papillomavirus type 1a"
> ewing@taxId
[1] "9606" "130308" "10583"
> ewing@releaseDate
[1] "2007-04-27"
```

organismName records all the organisms for which interactions were assayed. For each organism, we have also included its taxonomy identification code. Because IntAct does not currently version its weekly release, we have added the releaseDate as a time stamp to act as a surrogate for the version number.

Let us investigate the structure of the *interactions* slot. This slot contains a list which holds all the binary interactions given within the XML file (along with information about each particular interaction). Each element of the list is an instance of the class *intactInteraction* class. This class has 9 slots:

> slotNames(interactions(ewing)[[1]])

```
[1] "intact" "interactionType" "expPubMed" "expIntAct"
[5] "confidenceValue" "bait" "prey" "inhibitor"
```

[9] "neutralComponent"

The various slots contain information which is relevant for each individual interaction. The *interactionType* slot details what manner of interaction was found between the bait protein and the prey protein, which are specified in the *bait* and *prey* slots. Another important attribute is the experimental confidence value given in the *confidenceValue* slot. This confidence value is reported by the experimenters; it does not report scores derived by third parties.

We can extract the names of the bait and prey proteins for all of the interactions in the ewing dataset:

```
> ewbait <- sapply(interactions(ewing), bait)
> ewprey <- sapply(interactions(ewing), prey)</pre>
```

We now have two character vectors, ewbait and ewprey, that are aligned to each other: the  $i^{th}$  protein in ewprey is found by the  $i^{th}$  protein in ewbait.

#### > ewbait

```
intactId intactId intactId intactId intactId
"EBI-491274" "EBI-491274" "EBI-963841" "EBI-491274" "EBI-963841"
```

# > ewprey

```
intactId intactId intactId intactId intactId
"EBI-448924" "EBI-448924" "EBI-491274" "EBI-448924" "EBI-765551"
```

The IntAct accession codes are useful as unique and uniform identifiers in the IntAct repository, but we will usually want to translate them to other identifier schemes such as HUGO gene name Ensembl gene identifier. The PSI-25 XML files from IntAct contain a look-up table for this purpose. This look-up table is stored in the interactors slot of the interactionEntry object ewing, in the form of a character matrix. Its rows are indexed by the IntAct accession numbers of the molecules in the data structure, and its has 7 columns.

#### > interactors(ewing)

```
uniprotId geneName fullName locusName
EBI-491274 "P06400" "RB1" "Retinoblastoma-associated protein" NA
EBI-987160 "Q6H1D8" "E1A" "E1A" NA
EBI-448924 "Q01094" "E2F1" "Transcription factor E2F1" NA
```

```
"E7"
                               "Protein E7"
EBI-963841 "P06465"
                                                                      NA
EBI-765551 "000716"
                      "E2F3"
                               "Transcription factor E2F3"
                                                                      NA
           orfName organismName
                                                     taxId
                                                     "9606"
EBI-491274 NA
                    "Homo sapiens"
EBI-987160 NA
                    "Human adenovirus E"
                                                     "130308"
EBI-448924 NA
                    "Homo sapiens"
                                                     "9606"
EBI-963841 NA
                    "Human papillomavirus type 1a" "10583"
                    "Homo sapiens"
EBI-765551 NA
                                                     "9606"
```

The IntAct accession codes can be translated into any of the associated identifier schemes. Two further properties are given for each molecule: the organism in which the molecule is native and the corresponding taxonomy ID. Most of the interactions found in IntAct will be protein-protein interactions; other types of interactions, however, are also stored such as small molecule to protein interactions as well as gene-gene interactions. As a result, there will be times when a molecule cannot be mapped to a locus name or an ORF etc. We also remark that interactions have been tested between proteins of different organisms (i.e human protein against mice). Thus the organism attribute is vital to keep such interactions in the proper context.

Using the look-up table is quick and efficient because of the subsetting functionality of R. For instance, say we would like to translate the following IntAct accession codes

# 3 Obtaining Protein Complex Composition Information

Now we will demonstrate the parser function psi25complex. We remark here that the protein complexes which this function obtains have been manually curated from literature sources by *IntAct* curators.

The parameters of psi25complex are identical to those of psi25interaction, while its output only contains 3 slots:

```
> url2 <- system.file("PSI25XML/complexSample.xml", package="Rintact")
> comps <- psi25complex(url2)
> slotNames(comps)
[1] "releaseDate" "interactors" "complexes"
```

Again the releaseDate slot serves as a surrogate version number. The interactors slot again holds a look-up table that can be used to translate the IntAct accession codes. The complexes slot is a list of intactComplex objects. Each list entry is an instance of the class intactComplex, which itself has 7 slots.

```
> length(complexes(comps))
[1] 174
> class(complexes(comps)[[1]])
[1] "intactComplex"
attr(,"package")
[1] "Rintact"
> slotNames(complexes(comps)[[1]])
[1] "intactId" "shortLabel" "fullName" "organismName" "taxId"
[6] "members" "attributes"
```

These slots describe the multi-protein complex. The three most important ones are full-Name, attributes and members slots. The fullName slot gives the exact name of the multi-protein complex while the attributes slots gives a short description as to the known functionality of the complex. The interactors slot gives the members of the complex and their multiplicity.

```
> complexes(comps)[[1]]
complex ( EBI-706546 )
 _____
 [ short label ]: bcl2_bcl2_human
 [full name]: BCL-2 homodimer
 [ organism ]: Homo sapiens
 [ taxonomy ID ]:
                 9606
 [ attributes ]:
curated-complex: Role of homodimer is unclear, may act as reservoir of protein f
 or heterodimer formation.
complex-synonym: Bcl2 homodimer; Bcl-2:Bcl-2; Bcl2:Bcl2;
kd: 0.0
 [ members ]:
         intActId multiplicity
intactId EBI-77694
```

#### > toLatex(sessionInfo())

- R version 2.9.0 (2009-04-17), x86\_64-unknown-linux-gnu
- Locale: LC\_CTYPE=en\_US; LC\_NUMERIC=C; LC\_TIME=en\_US; LC\_COLLATE=en\_US; LC\_MONETARY=C; LC\_MESS
- Base packages: base, datasets, graphics, grDevices, grid, methods, stats, tools, utils
- Other packages: apComplex 2.10.0, DBI 0.2-4, graph 1.22.0, RBGL 1.20.0, Rgraphviz 1.22.0, Rintact 1.6.0, RSQLite 0.7-1, XML 2.3-0, xtable 1.5-5
- Loaded via a namespace (and not attached): AnnotationDbi 1.6.0, Biobase 2.4.0, cluster 1.11.13, hypergraph 1.16.0, org.Sc.sgd.db 2.2.11

Table 1: The output of sessionInfo on the build system after running this vignette.

# References

[Gavin et al.(2007)] Gavin AC, Aloy P, Grandi P, Krause R, Boesche M, Marzioch M, Rau C, Jensen LJ, Bastuck S, Dümpelfeld B, et al. Proteome survey reveals modularity of the yeast cell machinery. *Nature* 2006, 440:631–636.

[Ewing et al.(2007)] Ewing EM et al. Large-scale Mapping of Protein-Protein Interactions by Mass Spectrometry. *Molecular Systems Biology* 2007, 3.