Annotations with NetAffx

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Affymetrix provides annotations for all arrays they produce. The annotations are made available in Bioconductor with the NetAffxResource class in the AffyCompatible package; additional packages complement Affymetrix annotation information with data collected from additional public repositories. This document outlines a simple workflow to retrieve annotations available through NetAffx.

> library(AffyCompatible)

To use these facilities, one must be a registered Affymetrix user; see the Affymetrix user registration site for details.

The first step is to create an instance of the NetAffxResource class. Do this using the NetAffxResource function. Important arguments are user and password length 1 character vectors containing the registered user name and password. The password is printed, saved, and transmitted in clear text, and so is not secure. An additional argument is directory, which is the location where the NetAffx data base and downloaded files are stored. directory defaults to a session-specific temporary directory, meaning that if it is not supplied the data base and any downloaded annotations are removed when the R session ends. To create the NetAffxResource instance, evaluate a command like

```
> rsrc <- NetAffxResource(user = "mtmorgan@fhcrc.org", password = password)
> rsrc
directory: /tmp/RtmpIbme3d
annotationsFile: NetAffxAnnotFileList.xml
affxUrl: https://www.affymetrix.com/analysis/downloads/netaffxapi/GetFileList.jsp
affxLicence: ********
user: mtmorgan@fhcrc.org
password: ********
```

This creates the resource, but does not validate the user name and password (the user name and password are verified when the NetAffx resource is first retrieved from Affymetrix, typically the first time the code in the following paragraph is evaluated).

A typical workflow involves querying rsrc for the names of available arrays, and the descriptions of annotations available for an array of interest:

"Probe Sequences, FASTA format"

[9] "Probe Sequences, tabular format"

[7] "PSI Library File"

Annotations usually include a comma-separated value (CSV) file that can be represented in R as a data.frame. The data frame usually includes a probe identifier column, and columns of additional information Affymetrix has collated from a variety of sources, as described on the NetAffx site. Additional annotation files usually include a (much larger physically, but containing comparable information) MAGE-ML representation of the CSV file, channel description files (CDF), other files describing probes present on chips, probe sequences in FASTA format, and possibly other files specific to the chip platform.

An R representation of the annotations of a particular array can be created with

```
> annos <- rsrc[["Porcine"]]
> annos

affxName: Porcine
affxAnnotation: AffxAnnotation(8)
```

A particular annotation can be selected from this using R commands to navigate the implied class structure:

```
> sapply(affxAnnotation(annos), force)[1:5]
[[1]]
```

affxType: Annot CSV
affxDescription: Annotations, CSV format

affxFile: AffxFile(1)

[[2]]
affxType: Annot MAGE-ML

affxDescription: Annotations, MAGE-ML XML Format

affxFile: AffxFile(1)

[[3]]

affxType: BLASTp CSV

affxDescription: BLASTP annotations, CSV Format

```
affxFile: AffxFile(1)

[[4]]
affxType: CDF
affxDescription: CDF Library File
affxFile: AffxFile(1)

[[5]]
affxType: CIF
affxDescription: CIF Library File
affxFile: AffxFile(1)

> anno <- affxAnnotation(annos)[[3]]
> anno

affxType: BLASTp CSV
affxDescription: BLASTP annotations, CSV Format
affxFile: AffxFile(1)
```

(The Porcine BLASTP Annotation file is chosen because it is small). The annotation file may also be obtained by subsetting the reseource with a second argument corresponding to the annotation description or index

```
> anno <- rsrc[["Porcine", "BLASTP annotations, CSV Format"]]
> anno <- rsrc[["Porcine", 3]]</pre>
```

Annotation files can be retrieved with

```
> df <- readAnnotation(rsrc, annotation = anno)</pre>
```

This checks to see if the relevant annotation file is in the directory specified in the rsrc object. If the annotation file is not present, it is retrieved from the Affymetrix site. The argument update=TRUE forces retrieval. readAnnotation will read files with known type (e.g., CSV) into appropriate R objects (e.g., data frames), and return these from readAnnotation. Some file types (e.g., CDF) are not meant for representation as R objects, and for these readAnnotation returns the (local) path to the relevant file. For all annotations, the argument content=FALSE returns the local file path, without loading the content of the file into R.

Affymetrix does not specify the format of all files, so some files might reasonably be read into R but the readAnnotation code is not able to identify the appropriate format. The user is free to explore these annotation files using standard R commands, e.g.,

```
> anno <- rsrc[["Porcine", "PSI Library File"]]
> fl <- readAnnotation(rsrc, annotation=anno, content=FALSE)
> fl
```

^{[1] &}quot;/tmp/RtmpIbme3d/Porcine.psi.zip"

- > ## a zip file, containing 'Porcine.psi'
- > conn <- unz(f1, "Porcine.psi")</pre>
- > readLines(conn, n=6)
- [1] "#Probe Sets: 24123" "1\tAFFX-BioB-5_at\t20" "2\tAFFX-BioB-M_at\t20"
- [4] "3\tAFFX-BioB-3_at\t20" "4\tAFFX-BioC-5_at\t20" "5\tAFFX-BioC-3_at\t20" $^{"}$
- > read.table(conn, header=FALSE, skip=1, sep="\t", nrows=5)
 - V1 V2 V3
- 1 1 AFFX-BioB-5_at 20
- 2 2 AFFX-BioB-M_at 20
- 3 3 AFFX-BioB-3_at 20
- 4 4 AFFX-BioC-5_at 20
- 5 5 AFFX-BioC-3_at 20