cdfenvs and oligonucleotide arrays

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Introduction

This document describes briefly how the package gets (or tries to get) the needed cdfenvs. Various issues like security and configuring the options for the package affy are outlined. To some extend, what is developed here could¹ be applied to the package *oligochips*.

As usual, loading the package in your R session is required.

R> library(affy) ##load the affy package

Why and how

The *cdfenvs* are associative data structures to map efficiently a *probeset id* with indices for the corresponding probes. These indices are used to subset a matrix storing probe level intensities. Technically, a *cdfenv* is a **R** environment. The functions get and multiget (in *Biobase*) are a convenient way to access what is in an environment. Expert users only will consider modifying what is those environment.

In the case of Affymetrix data, the cdfenvs are built from the .CDF files. The package makecdfenvs is dedicated to the building of packages with cdfenvs. The simplest way to proceed is to have the package needed for the analysis installed. A number of cdfenv packages for Affymetrix chips are available for download on the Bioconductor data packages repository. This release (1.2.x) of the affy package has an option to allow automatic downloading and install of a cdfenv that would be found missing during an analysis. However, if you are using an unconventional chip, it is possible that Bioconductor has not created the appropriate package for your .CDF file. We recommend that you use the package makecdfenvs to create the appropriate source code for the cdfenv package you need. For Microsoft Windows binaries you need to do more, see http://www.stats.ox.ac.uk/pub/R/rw-FAQ.html. If you contribute the package to

¹Currently this would require a bit of effort, but it should become very easy very soon

Bioconductor a windows binary will be made for you (and everybody else). The last section outlines briefly how to change the associated *cdfenv*.

The complete structure of the options for the package is not completely described, but one can refer the source code for the function .setAffyOptions(). The entry *probesloc* in the options details the path used to look for the corresponding *cdfenv*. We introduce a simple function to display the content of the options for the obtention of the information about probe locations:

```
> print.probesloc.opt <- function(affy.opt, fields) {</pre>
      all.fields <- c("what", "where", "autoload", "repository",
+
           "installdir")
+
      if (sum(!(fields %in% all.fields)) > 0)
+
           stop(paste("'fields' can only contain elements of:",
+
               paste(all.fields, collapse = " ")))
+
      1 <- lapply(affy.opt$probesloc, function(x) x[fields])</pre>
+
      1 <- lapply(l, function(x) {</pre>
+
           unk <- is.na(names(x))</pre>
+
           x[unk] <- rep(list(unk = NA), sum(unk))</pre>
+
           x <- lapply(x, function(y) if (is.null(y))</pre>
+
+
                "NULL"
           else y)
+
           return(x)
+
      })
+
      ul <- as.character(unlist(1))</pre>
+
      m <- t(matrix(ul, nr = length(fields)))</pre>
+
      colnames(m) <- fields</pre>
+
+
      print(m)
+ }
```

The default search path for *cdfenvs* will be:

```
> affy.opt <- getOption("BioC")$affy
> print.probesloc.opt(affy.opt, c("what", "where", "autoload"))
    what where autoload
[1,] "environment" "<environment>" "NA"
[2,] "libPath" "NULL" "NA"
[3,] "data" "affy" "NA"
[4,] "bioC" "/tmp/Rinst212077129" "NA"
```

The option autoload is only relevant where what is equal to *package*. Having it set to TRUE means that an attempt will be made to download the package if it is not found in .libPaths() or in the search path shown above.

The following function returns a set of options with any existing automatic download deactivated :

```
> deactivate.autoload <- function(affy.opt) {</pre>
       1 <- lapply(affy.opt$probesloc, function(x) {</pre>
+
           i <- names(x) == "autoload"</pre>
+
           x[i] <- list(FALSE)</pre>
+
+
           return(x)
+
       7)
+
      affy.opt$probesloc <- 1
      return(affy.opt)
+
+ }
```

It can be used to deactivate any automatic download:

```
> affy.opt <- getOption("BioC")$affy
> affy.opt.noauto <- deactivate.autoload(affy.opt)
> .setAffyOptions(affy.opt.noauto)
```

Security

The autoload mecanism can be perceived as a security breach. It is the case, but not more than any \mathbf{R} package you might install (unless you inspect carefully the source of every single package you install).

By default the *cdfenv* packages are downloaded from the bioconductor repository. This can be changed through the options for the package. One might want to check what it is like on his/her local installation:

```
> print.probesloc.opt(affy.opt, c("what", "autoload", "repository"))
```

	what	autoload	repository
[1,]	"environment"	NA	NA
[2,]	"libPath"	NA	NA
[3,]	"data"	NA	NA
[4,]	"bioC"	NA	NA

If you do not have install permissions, you might consider installing the *cdfenvs* packages to a particular place. For that you need to change the *installdir* (passed to the function install.packages).

```
> my.installdir <- "mydir/is/here"
> has.installdir <- unlist(lapply(affy.opt$probesloc, function(x) if ("installdir" %i
+ names(x)) grep("installdir", names(x)) else numeric(0)))
> l <- lapply(affy.opt$probesloc, function(x) {
+ if ("installdir" %in% names(x)) {
+ x$installdir <- my.installdir</pre>
```

```
+ }
+ return(x)
+ })
> affy.opt$probesloc <- l
> .setAffyOptions(affy.opt)
```

Note that a given *cdfenv* package is searched as specified in the argument *where*. Depending on your settings, you might prefer updating the .libPaths() of your **R** session, or the field *where* for the probelocs).

Changing the associated *cdfenv*

If you do not care about mapping *probeset ids* with indices and only want to deal with probe intensities, or if you would like to make a custom *cdfenv* from scratch, you will have to modify the association between your AffyBatch object(s) and *cdfenvs*.

If you want to experiment with this feature of the package, we would recommend to create a dummy *cdfenv* and associate your AffyBatch to it:

```
> data(affybatch.example)
```

```
> dummymap.name <- "dummymap"</pre>
```

```
> assign(dummymap.name, new.env())
```

```
> affybatch.example@cdfName <- paste(dummymap.name, "cdf", sep = "")</pre>
```

As explained in the first part, the *affy* package will then find a matching environment in the current environment (.GlobaEnv) and will not try to find a package. Note: the environment should still be in the section *probelocs* of the package options.