oneChannelGUI Package Installation

Raffaele A Calogero, Francesca Cordero, Remo Sanges ${\rm May}\ 12\ 2011$

1 Installation

It is important to install additional software other than R and Bioconductor libraries, for the complete functionality of oneChannelGUI.

The external tools and data needed to run exon array analysis in oneChannelGUI are shown in figure 1.

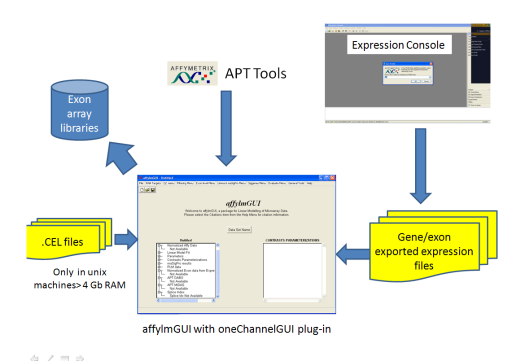


Figure 1: External tools needed for exon array analysis when oneChannelGUI add-on is loaded in affylmGUI.

1.1 Windows users

The lastest version of R need to be installed. It can be downloaded at: http://www.r-project.org/oneChannelGUI needs the installation of the following packages:

- > packageDescription("oneChannelGUI") \$Depends
- [1] "Biobase, affylmGUI, tkrplot, tkWidgets, IRanges, Rsamtools (>= 1.13.1),\nBiostrir oneChannelGUI suggests the installation of the following packages:
- > packageDescription("oneChannelGUI")\$Suggests
- [1] "annotate, genefilter, maSigPro, pamr, pdmclass, ChIPpeakAnno,\nchipseq, BSgenome,

In order to install all the Bioconductor packages needed to run oneChannelGUI a specific R function can be downloaded from http://www.bioinformatica.unito.it/oneChannelGUI/ To use it user needs to open an R session on a computer connected to internet and copy and paste the following code:

oneChannelGUI inherits the core graphical interface of affylmGUI. This interface need, under windows, the installation of TCL/TK libraries that can be download from: http://www.activestate.com/Products/ActiveTcl/

To load Hs/Mm/Rn exon 1.0 ST Affymtrix arrays users need to install APT tools 1.12.0. The windows binaries can be downloaded from: http://www.affymetrix.com/support/developer/powertools/index.affx In addition library files for gene/exon arrays are directly downloaded and installed locally from oneChannelGUI interface, the first time an Exon 1.0 ST or Gene 1.0 ST analysis is started. Or using the function oneChannelGUI: Set library folder and install Affy gene/Exon library files present in the General tools Menu.

IMPORTANT APT TOOLS AND EXON LIBRARY FILES NEED TO BE LOCATED IN A FOLDER CHARACTERIZED BY A PATH WITHOUT SPACES C:/apt.1.8.0 THIS IS OK

C:/PROGRAM FILES/APT.1.8.0 STARTING APT TOOLS FROM oneChannelGUI will produce an er

If you are interested to run RNA-seq secondary analysis it is also required the installation of perl. The easy way is to install the active perl: http://www.activestate.com/activeperl

1.2 Unix users

The packages to be downloaded for unix users are those indicated for windows users unless for the TCL/TK libraries.

To use Tcl/Tk then the interface package tcltk must be installed into R. If an R package uses Tcl/Tk then Tcl/Tk must be installed on the OS before the R package can be installed into R. Furthermore, these two Tcl/Tk packages are needed by oneChanneGUI, affylmGUI and limmaGUI for their functionality:

- 1. BWidget, this is a package written entirely in the tcl scripting language and hence is not compiled. Consequently it is the same on any platform and should be in the Tcl/Tk path.
- 2. Tktable, this package has some code written in C and hence is a compiled package. It therefore is different for each type of CPU and OS.

It is possible to encounter problems in installing those libraries on x64 machines. For this reason I found quite useful the suggestions of Keith Satterley (24 Apr 2007) subject: Re: [Bioc-devel] Tcl/TK pckage "Bwidget" available on the Bioconductor mailing list.