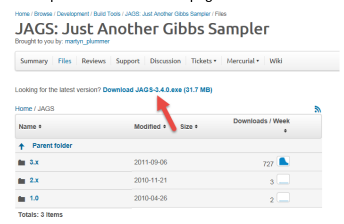
How to install the R-package "crmPack" for model-based dose escalation in R

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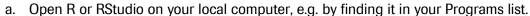
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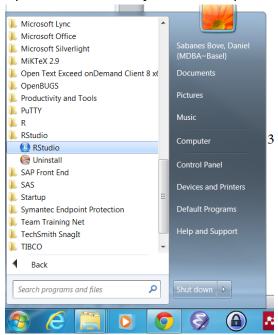
Following this installation guide, you will install the R-package "crmPack" on your local computer with your local R installation. Please note that it has nothing to do with the remote RStudio that you can access on https://r.roche.com. Your local R and the remote RStudio on the server are completely independent. We assume here that you have already installed R or RStudio on your computer.

- 1. First, you have to install JAGS ("Just Another Gibbs Sampler") on your computer. Note that this requires you to have administrative rights on your computer.
 - Click on the following link to visit the JAGS webpage: http://sourceforge.net/projects/mcmc-jags/files/JAGS/
 - b. Download the JAGS installation program suitable for your computer by clicking on the top link in the JAGS webpage:



- c. Run the JAGS installation program by executing the downloaded file, and clicking through and confirming all installation setup steps (including warnings).
- 2. Now we will install the latest version of the R-package "crmPack" to your computer.





b. Please copy the following lines into your R console and press Enter:

```
options(repos=structure(c(CRAN="http://stat.ethz.ch/CRAN/")))
install.packages("httr")
library("httr")
url <-
  paste("https://stash.intranet.roche.com/stash/plugins/servlet/archive/projects/RSTAT",
         "/repos/crmpack?at=refs%2Fheads%2Fmaster",
         sep="")
tmp <- tempdir()
target <- file.path(tmp, "crmpack.zip")
x <- GET(url, config = list(ssl.verifypeer = FALSE))
bin <- content(x, "raw")
writeBin(bin, target)
dir <- file.path(tmp, "crmpack")
unzip(zipfile=target, exdir=dir)
install.packages(c("Rcpp", "RcppArmadillo", "rjags", "ggplot2", "gridExtra",
            "GenSA", "BayesLogit", "mvtnorm"), dependencies=TRUE)
install.packages(dir, repos=NULL, type="source")
```

Please be patient as this may take a minute.

c. The successful installation looks like this:

```
Console -/ Console -/
```

3. Before you can use the R-package "crmPack", you have to load it into your R session, by typing the command

library("crmPack")

into your console.

Note that you have to run library("crmPack") again for each new R session!

On the other hand, you only have to do the *installation* process (step 2) *once* on your computer. When you have successfully completed step 2, then only in case you would like to update your "crmPack" installation with a new version, you would have to run completely through step 2 of this installation guide again.

4. In order to read an introductory example to the package, type the following into your console: crmPackExample()

In order to get help on specific functions or classes, get a list of all help pages by typing crmPackHelp()

Thank you very much for installing crmPack.

If you have any questions or feedback, please post on the Hive page:

https://roche.jiveon.com/groups/open-source-framework/projects/crmpack

Thank you!