

From the Genepix data files to RGList to NChannelSet

Audrey Kauffmann, Wolfgang Huber

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Load the required packages

```
> library("Biobase")
> library("limma")
> library("CCl4")
```

Read the data and convert them into an RGList

The Genepix (.gpr) data files are in the `extdata` directory of the `CCl4` package. If you have the package installed, we can locate them on your filesystem with the function `system.file`. If the files are somewhere else, please adapt the below assignment to `datapath`.

```
> datapath = system.file("extdata", package="CCl4")

> p = read.AnnotatedDataFrame("samplesInfo.txt", path=datapath)
> CCl4_RGList = read.maimages(files=sampleNames(p),
+     path = datapath,
+     source = "genepix",
+     columns = list(R = "F635 Median", Rb = "B635 Median",
+                     G = "F532 Median", Gb = "B532 Median"))
```

If this code is run in the `inst/doc` directory of the `CCl4` (source) package, the output data files will be written directly into the `data` directory of the package. Otherwise, just write into a temporary directory.

```
> outdir = file.path("../", "..", "data")
> if(!isTRUE(file.info(outdir)$isdir))
+   outdir = tempdir()
> save(CCl4_RGList, file = file.path(outdir, "CCl4_RGList.RData"))
```

The function `read.maimages` from the `limma` package reads the .gpr files and builds an `RGList` object from it.

The output is written to

```
> outdir
[1] "../data"
```

Build an NChannelSet from the RGList

Once the `RGList` object has been created, we can build an `NChannelSet`.

```

> sessionInfo()
R version 2.10.0 Under development (unstable) (2009-09-03 r49541)
x86_64-unknown-linux-gnu

locale:
[1] LC_CTYPE=en_US.UTF-8      LC_NUMERIC=C
[3] LC_TIME=en_US.UTF-8       LC_COLLATE=en_US.UTF-8
[5] LC_MONETARY=C            LC_MESSAGES=en_US.UTF-8
[7] LC_PAPER=en_US.UTF-8     LC_NAME=C
[9] LC_ADDRESS=C              LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

attached base packages:
[1] tools      stats      graphics   grDevices  utils      datasets   methods
[8] base

other attached packages:
[1] CC14_1.0.9    limma_2.19.4  Biobase_2.5.6

```

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

```

> featureData = new("AnnotatedDataFrame", data = CC14_RGList$genes)
> assayData = with(CC14_RGList, assayDataNew(R=R, G=G, Rb=Rb, Gb=Gb))
> varMetadata(p)$channel=factor(c("G", "R", "G", "R"),
+                                levels=c(ls(assayData), "_ALL_"))
> CC14 <- new("NChannelSet",
+             assayData = assayData,
+             featureData = featureData,
+             phenoData = p)
> save(CC14, file = file.path(outdir, "CC14.RData"))

```