

# Overview of GGtools for genetics of gene expression in Bioconductor for Bioc 2.2, 2008

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## 1 Introduction

The *GGtools* package contains infrastructure and demonstration data for joint analysis of transcriptome and genome through combination of DNA expression microarray and high-density SNP genotyping data. For Bioconductor 2.2 we adopted a representation of genotypes due to Clayton (in package *snpMatrix*) allowing reasonably convenient storage and manipulation of 4 megaSNP phase II HapMap genotypes on all the CEPH CEU samples. This contrasts with the previous version of *GGtools* which was limited to 550 kiloSNP and 58 CEU founders.

To give an immediate taste of the capabilities, we attach the package and load some test data.

```
> library(GGtools)
> data(hmceuB36.2021)
> hmceuB36.2021
```

```
snp.matrix-based genotype set:
number of samples: 90
```

```

number of snp.matrix: 2
annotation:
  exprs: illuminaHumanv1.db
  snps: snp locs package: GGBase ; SQLite ref: hmceuAmbB36_23a_dbconn
Expression data: 47293 x 90
Phenodata: An object of class "AnnotatedDataFrame"
  sampleNames: NA06985, NA06991, ..., NA12892 (90 total)
  varLabels and varMetadata description:
    famid: hapmap family id
    persid: hapmap person id
    ....: ...
    isAdad: logical TRUE if person is a father
    (9 total)

```

Expression data are recoverable in a familiar way:

```
> exprs(hmceuB36.2021)[1:5, 1:5]
```

	NA06985	NA06991	NA06993	NA06994	NA07000
GI_10047089-S	5.983962	5.939529	5.912270	5.891347	5.906675
GI_10047091-S	6.544493	6.286516	6.244446	6.277397	6.330893
GI_10047093-S	9.905235	10.353804	10.380972	9.889223	10.155686
GI_10047099-S	7.993935	7.593970	8.261215	6.598430	6.728085
GI_10047103-S	11.882265	12.204753	12.249708	11.798415	12.015252

Genotype data have more complex representation.

```
> smList(hmceuB36.2021)
```

```
$`20`
```

```

A snp.matrix with 90 rows and 119921 columns
Row names: NA06985 ... NA12892
Col names: rs4814683 ... rs6090120

```

```
$`21`
```

```

A snp.matrix with 90 rows and 50165 columns
Row names: NA06985 ... NA12892
Col names: rs885550 ... rs10483083

```

This shows that we use a named list to hold items of the *snp.matrix* class from *snpMatrix*. We can dig in with a different accessor:

```
> rawSNP(hmceuB36.2021, 20)[1:5, 1:5]
```

	rs4814683	rs6076506	rs6139074	rs1418258	rs7274499
NA06985	03	03	03	03	03
NA06991	02	03	02	02	03
NA06993	01	03	01	01	03
NA06994	01	03	01	01	03
NA07000	03	03	03	03	03

and now we see that an unusual data representation is in use: the leading zeroes indicate that a raw byte representation is being shown.

We can coerce from this representation to allele tokens:

```
> as(smList(hmceuB36.2021)[["20"]][1:5, 1:5], "character")
```

```
      [,1] [,2] [,3] [,4] [,5]
[1,] "B/B" "B/B" "B/B" "B/B" "B/B"
[2,] "A/B" "B/B" "A/B" "A/B" "B/B"
[3,] "A/A" "B/B" "A/A" "A/A" "B/B"
[4,] "A/A" "B/B" "A/A" "A/A" "B/B"
[5,] "B/B" "B/B" "B/B" "B/B" "B/B"
```

## 2 Chromosome-wide SNP screens

The demonstration object `hmceuB36.2021` has only chromosomes 20 and 21. An object in the *GGdata* package called `hmceuB36` has all 24 chromosomes. The *GGBase* package has the SNP locations for `hmceuB36` in an external SQLite store that is exported dynamically upon attaching *GGBase*, on which *GGtools* depends.

It is fairly easy to test for eQTL for a given gene on a specified chromosome:

```
> g1 = gwSnpScreen(genesym("CPNE1"), hmceuB36.2021, chrnum(20))
> class(g1)
```

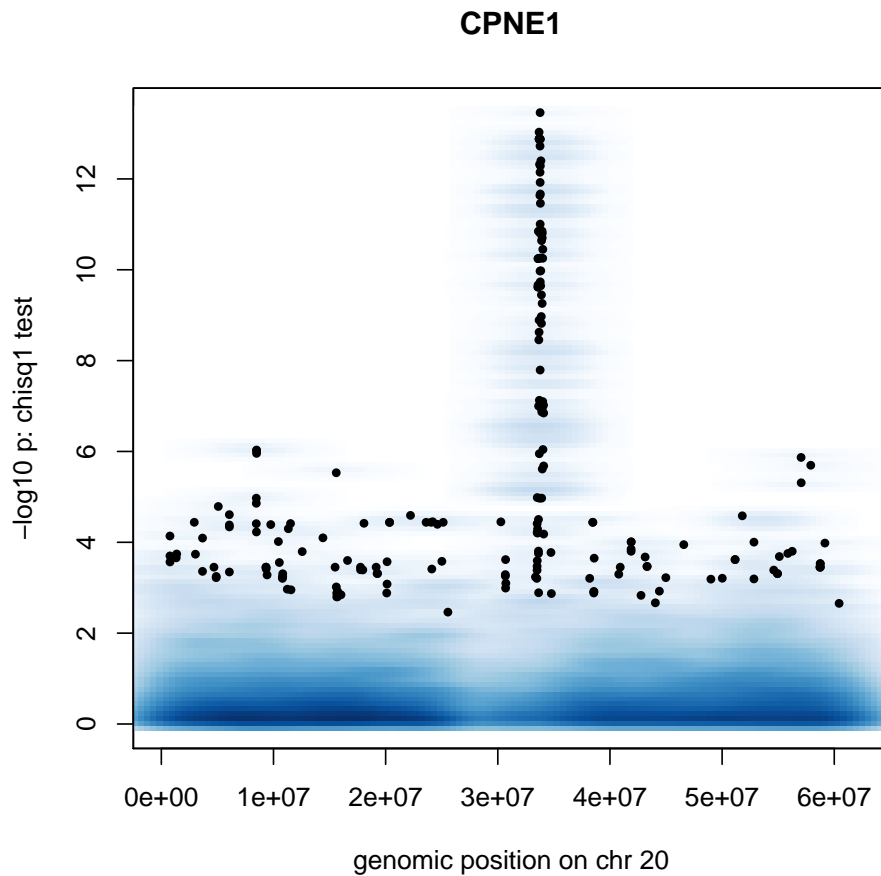
```
[1] "cwSnpScreenResult"
attr(,"package")
[1] "GGBase"
```

```
> g1
```

```
gwSnpScreenResult with 1 inference data.frames
gene used: CPNE1 ; expression platform: illuminaHumanv1.db
```

We can visualize the results over the chromosome:

```
> plot(g1)
```



and we can get a

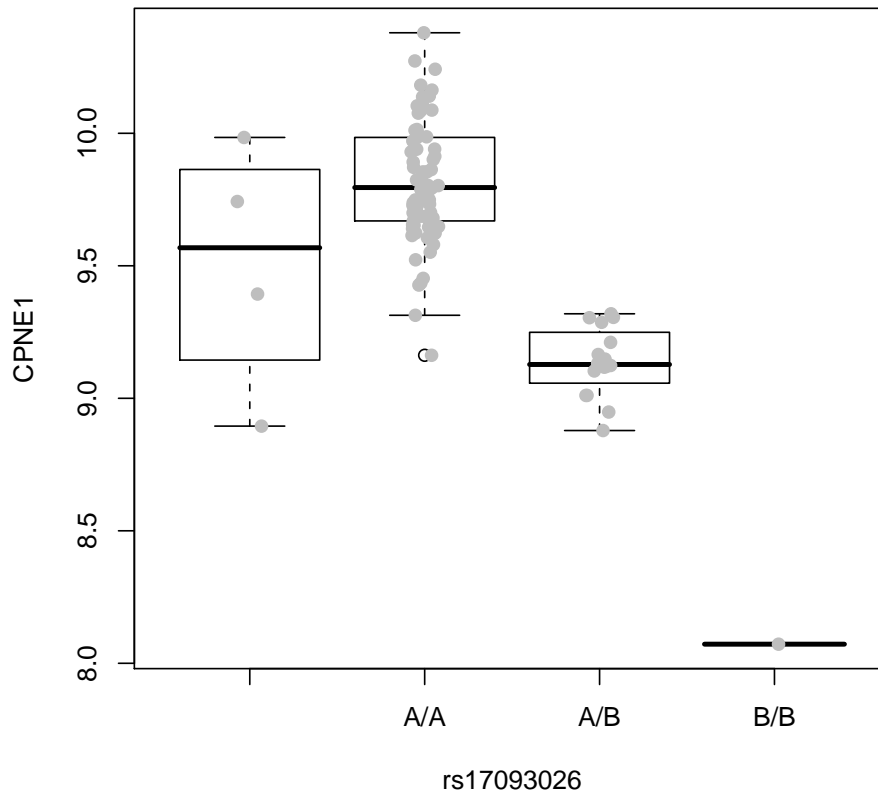
report on the most highly associated SNPs:

```
> tt = topSnps(g1)
> tt
```

	p.1df
rs17093026	3.464560e-14
rs1118233	9.322326e-14
rs7273815	1.304436e-13
rs12480408	1.330170e-13
rs6060535	1.330170e-13
rs11696527	1.330170e-13
rs6058303	1.330170e-13
rs2425078	1.330170e-13
rs6060578	1.330170e-13
rs1970357	1.330170e-13

To see the relationship between expression values and genotype, the `plot_EvG` method can be used.

```
> plot_EvG(genesym("CPNE1"), rsNum(rownames(tt)[1]), hmceuB36.2021)
```



### 3 Genome-wide SNP screens

The *GGdata* package contains a representation of all chromosomes.

```
> library(GGdata)
> if (!exists("hmceuB36")) data(hmceuB36)
```

We can use `gwSnpScreen` to find eQTL, if we have sufficient memory. On windows, we skip this step.

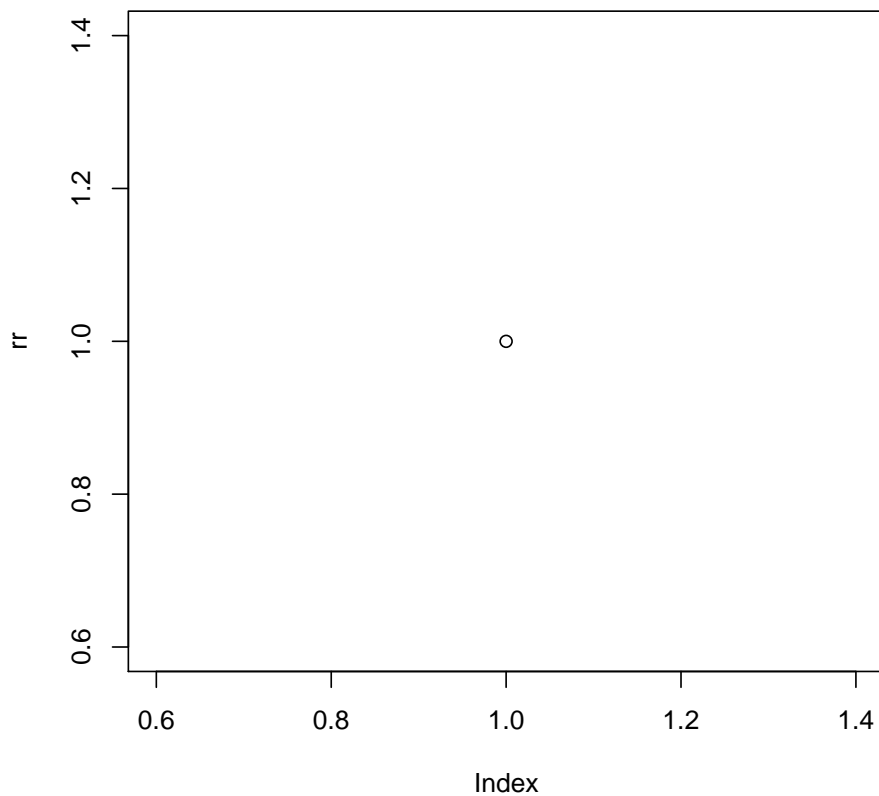
```
> ONWIN = FALSE
> ONWIN = (Sys.info()["sysname"] == "Windows")
> if (!ONWIN) {
+   rr = gwSnpScreen(genesym("RPS26"), hmceuB36)
+   rrt = topSnps(rr)
```

```
+   rrt[[12]]  
+ }
```

Note that `topSnps`, when applied to a genome-wide screen output, will return a list of chromosome-specific results.

The whole-genome plot is automatic but somewhat slow; on windows, this is a placeholder. Please get the genuine vignette from the web site.

```
> if (ONWIN) {  
+   rr = 1  
+ }  
> plot(rr)
```



## 4 Phenotype data available on CEPH samples

```
> names(pData(hmceuB36))
```

```
[1] "famid"      "persid"    "mothid"    "fathid"    "sampid"    "isFounder"
[7] "male"      "isAmom"    "isAdad"
```

## 5 Session information

```
> sessionInfo()
```

```
R version 2.7.0 (2008-04-22)
i386-pc-mingw32
```

```
locale:
```

```
LC_COLLATE=English_United States.1252;LC_CTYPE=English_United States.1252;LC_MONETARY=E
```

```
attached base packages:
```

```
[1] grid      splines  tools      stats      graphics  grDevices  utils
[8] datasets  methods  base
```

```
other attached packages:
```

```
[1] GGdata_0.1.1          illuminaHumanv1.db_1.0.0 GGtools_2.0.2
[4] Biostrings_2.8.6      RColorBrewer_1.0-2      mgu74av2.db_2.2.0
[7] geneplotter_1.18.0    hgfocus.db_2.2.0        GGBase_2.0.3
[10] snpMatrix_1.4.0       hexbin_1.14.0           lattice_0.17-8
[13] survival_2.34-1       GSEABase_1.2.1          annotate_1.18.0
[16] xtable_1.5-2          AnnotationDbi_1.2.0      RSQLite_0.6-8
[19] DBI_0.2-4             Biobase_2.0.1
```

```
loaded via a namespace (and not attached):
```

```
[1] cluster_1.11.10      graph_1.18.1           KernSmooth_2.22-22 Ruuid_1.18.0
[5] XML_1.95-2
```