1 Introduction

Data representations for genetics of gene expression in humans must be as efficient as possible. Up to Bioconductor 2.1, integrative genomics experiments could be represented by racExSet instances as defined in package GGtools. Genotypes were represented as integers.

For Bioconductor 2.2, we have access to the snpMatrix package of D. Clayton, and we have introduced the smlSet class to represent genetics of gene expression studies with lists of snp.matrix instances, along with ordinary representations of expression data.

2 Primary class structure, and associated methods

```r
> library(GGBase)
> getClass("smlSet")
```

Class "smlSet"

Slots:

- Name: smlEnv, annotation, chromInds
  - Class: environment, character, numeric

- Name: organism, assayData, phenoData
  - Class: character, AssayData, AnnotatedDataFrame

- Name: featureData, experimentData, ._classVersion_
  - Class: AnnotatedDataFrame, MIAME, Versions

Extends:
Class "eSet", directly
Class "VersionedBiobase", by class "eSet", distance 2
Class "Versioned", by class "eSet", distance 3

> library(GGtools)
> showMethods(class = "smlSet", where = "package:GGBase")

Function: [ (package base)
  x="smlSet", i="ANY", j="ANY", drop="ANY"

Function: combine (package Biobase)
  x="smlSet", y="smlSet"

Function: exprs (package Biobase)
  object="smlSet"

Function: getAlleles (package GGBase)
  x="smlSet", rs="rsid"

Function: getSnpsLocs (package GGBase)
  x="smlSet", c="chrnum"
  x="smlSet", c="missing"

Function: plot_EvG2 (package GGBase)
  gsym="genesym", rsid1="rsid", rsid2="rsid", sms="smlSet"
  gsym="probeId", rsid1="rsid", rsid2="rsid", sms="smlSet"

Function: plot_EvG (package GGBase)
  gsym="genesym", rsid="rsid", sms="smlSet"
  gsym="probeId", rsid="rsid", sms="smlSet"

Function: show (package methods)
  object="smlSet"

Function: smlEnv (package GGBase)
  x="smlSet"

Function: smList (package GGBase)
  x="smlSet"

Function: snpNames (package GGBase)
  x="smlSet", c="chrnum"
  x="smlSet", c="missing"
> showMethods(class = "smlSet", where = "package:GGtools")

Function: [ (package base)
x="smlSet", i=ANY, j=ANY, drop=ANY"

Function: gwSnpTests (package GGtools)
sym="formula", sms="smlSet", cnum="cnumOrMissing"
sym="formula", sms="smlSet", cnum="snpdepth"

Function: hbTests (package GGtools)
fmla="genesym", sms="smlSet", cnum="chrnum", rsid=numeric", rad="numeric"

Function: residTests (package GGtools)
fit="cwSnpScreenResult", sms="smlSet", litfmla="formula", rsnum="missing"

3  Example data structure

> data(hmceuB36.2021)
> hmceuB36.2021

snp.matrix-based genotype set:
number of samples: 90
number of snp.matrix: 2
annotation: illuminaHumanv1.db
Expression data dims: 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
    ...: ...
    male: logical TRUE if male
    (7 total)

4  SNP location infrastructure

see the newSNPloc vignette

5  Conclusion

The concepts discussed above define the bulk of the tools supporting the gwSnpScreen method of GGtools. Consult the vignette of GGtools for more information.