The R package trioClasses for definition of the class SNPTrioExperiment, an extension of SummarizedExperiment, for use in trio based analyses of genetic data.

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March 11, 2013

# 1 Packages

```
> library("trioClasses")
```

## 2 Sample Data

## 2.1 Pedigree

> ped <- PedClass(ped.DF)

## 2.2 SNPTrioExperiment

```
> ste <- SNPTrioExperiment(se, pedigree = ped)
> ste

class: SNPTrioExperiment
dim: 511 968
exptData(0):
assays(1): geno
rownames(511): chr14:54383433 chr14:54383470 ...
    chr14:54444385 chr14:54444750
rowData metadata column names(0):
colnames(968): H_ME-CSc175_1-CSc175_1_a.1
    H_ME-CSc175_2-CSc175_2 ... H_ME-DS11418_2-DS11418_2
    H_ME-DS11418_3-DS11418_3
colData names(28): vcfid jb_famid ... id ped
pedigree(4139): famid id fid mid sex dx
```

## 3 Methods

### 3.1 ScanTrio

```
> ste.rare <- ste[MAF(ste) <= 0.01]</pre>
> (scan.trio <- ScanTrio(object = ste.rare, window = rowData(ste.rare[1:10]),</pre>
     block = range(rowData(ste))))
               minor.win major.win mendel.win minor.out major.out
chr14:54383433
                                  1
                                              0
                                                      142
                                                                 528
chr14:54383819
                        2
                                              0
                                                      140
                                                                 527
chr14:54384232
                        0
                                  0
                                                      142
                                                                 529
                                              1
chr14:54384251
                        0
                                  1
                                              3
                                                      142
                                                                 528
                        0
                                  0
                                              0
chr14:54384658
                                                      142
                                                                 529
chr14:54384847
                        0
                                  1
                                              0
                                                      142
                                                                 528
chr14:54384907
                        0
                                  1
                                              0
                                                      142
                                                                 528
chr14:54384924
                        0
                                  1
                                              0
                                                      142
                                                                 528
                        0
                                             0
chr14:54384970
                                  1
                                                      142
                                                                 528
                        4
                                             3
                                                                 525
chr14:54385150
                                                      138
               mendel.out
chr14:54383433
                      202
chr14:54383819
                      202
chr14:54384232
                      201
chr14:54384251
                      199
chr14:54384658
                      202
chr14:54384847
                      202
chr14:54384907
                      202
chr14:54384924
                      202
chr14:54384970
                      202
chr14:54385150
                      199
```

## 3.2 Holger-style Genotype Matrix

Or apply the aTDT method to the SNPTrioExperiment directly.

### > aTDT(ste[1:5])

#### \$stat

[1] 1.000000 0.000000 0.000000 8.727273 19.565217

## \$pval

[1] 3.173105e-01 1.000000e+00 1.000000e+00 3.134851e-03 9.722321e-06

## \$transMinor

[1] 0 9 2 108 8

### \$transMajor

[1] 1 9 2 156 38

# A Count of Transmission of Variants (TransCount())

### A.1 Window

```
> window <- GRanges(seqnames = "chr14", IRanges(start(gr[100]),</pre>
     width = 10000), strand = "*")
> window2 <- GRanges(seqnames = "chr14", IRanges(start(gr[100]),</pre>
     width = 1000), strand = "*")
> block <- range(rowData(ste))</pre>
> TransCount(ste, window)
$minor
[1] 1106
$major
[1] 1345
$mendel
[1] 393
      Not in window, but in block
> TransCount(ste, setdiff(block, window))
$minor
[1] 7727
$major
[1] 9337
$mendel
[1] 2950
     In the whole block
A.3
> TransCount(ste, block)
$minor
[1] 8833
$major
[1] 10682
$mendel
[1] 3343
```

### A.4 In both windows

> TransCount(ste, GRangesList(window, window2))

```
$minor
[1] 1106
          48
$major
[1] 1345 175
$mendel
[1] 393 58
      Not "In both windows," but in block
> TransCount(ste, GRangesList(setdiff(block, window), setdiff(block,
     window2)))
$minor
[1] 7727 8785
$major
[1] 9337 10507
$mendel
[1] 2950 3285
     Count of Transmission of Rare Variants (TransCount())
\mathbf{B}
> ste.rare <- ste[MAF(ste) <= 0.01]</pre>
      In both windows
> TransCount(ste.rare, GRangesList(window, window2))
$minor
[1] 26 3
$major
[1] 85 7
$mendel
[1] 28 5
      Not "In both windows," but in block
> TransCount(ste.rare, GRangesList(setdiff(block, window),
     setdiff(block, window2)))
$minor
[1] 116 139
$major
[1] 444 522
```

## C Classes

## C.1 SnpMatrix, DataFrame, GRanges

The four key ingredients are the SNP matrix, the pedigree information as a DataFrame, position of the SNPs given by a GRanges object, and covariate data given as a DataFrame.

```
> sm
```

A SnpMatrix with 968 rows and 511 columns
Row names: H\_ME-DS10776\_2-DS10776\_2 ... H\_ME-DS11313\_1-DS11313\_1
Col names: chr14:54383433 ... chr14:54444750

> ped.DF

DataFrame with 4139 rows and 6 columns

```
famid
                                                id
     <character>
                                      <character>
            4778
                            H_ME-4778_1-4778_1.2
1
2
            4778
                            H_ME-4778_2-4778_2.2
3
                            H_ME-4778_3-4778_3.2
            4778
4
            4783
                            H_ME-4783_1-4783_1.1
5
            4783
                            H_ME-4783_2-4783_2.1
6
            4783
                            H_ME-4783_3-4783_3.1
7
        20000492
                    H_ME-20000492_1-20000492_1.1
8
        20000492 H_ME-20000492_2-20000492_2_a.1
                    H_ME-20000492_3-20000492_3.1
9
        20000492
              . . .
. . .
4131
         DS12332
                        H_ME-DS12332_1-DS12332_1
4132
         DS12332
                        H_ME-DS12332_2-DS12332_2
                        H_ME-DS12332_3-DS12332_3
4133
         DS12332
4134
         DS99998
                      H_ME-DS10193_5-DS10193_5.2
4135
         DS99998
                        H_ME-DS10193_6-DS10193_6
4136
         DS99998
                        H_ME-DS10193_7-DS10193_7
4137
         DS99999
                      H_ME-DS10707_4-DS10707_4.2
                        H_ME-DS10707_5-DS10707_5
4138
         DS99999
4139
         DS99999
                        H_ME-DS10707_6-DS10707_6
                                 fid
                                                                mid
                         <character>
                                                        <character>
1
               H_ME-4778_2-4778_2.2
                                              H_ME-4778_3-4778_3.2
2
                                                                  NA
                                   NA
3
                                   NA
                                                                  NA
4
               H_ME-4783_2-4783_2.1
                                              H_ME-4783_3-4783_3.1
5
                                   NA
                                                                  NA
6
                                   NA
                                                                  NA
7
     H_ME-20000492_2-20000492_2_a.1 H_ME-20000492_3-20000492_3.1
8
                                   NA
```

```
9
                                 NA
                                                              NA
. . .
           H_ME-DS12332_2-DS12332_2
4131
                                        H_ME-DS12332_3-DS12332_3
4132
4133
           H_ME-DS10193_6-DS10193_6
                                        H_ME-DS10193_7-DS10193_7
4134
4135
4136
                                 NA
                                                              NA
4137
                                 NA
                                                              NA
4138
         H_ME-DS10707_4-DS10707_4.2
                                        H_ME-DS10707_6-DS10707_6
4139
                                 NA
                                                              NA
                      dx
           sex
     <numeric> <numeric>
1
            1
                       1
2
             1
                       0
3
             2
                       0
4
             2
                       1
5
                       0
             1
6
             2
                       1
7
             2
                       1
8
             1
                       0
9
             2
                       0
4131
            1
                       1
4132
            1
                       0
            2
4133
                       0
4134
            2
                       1
4135
            1
             2
                       0
4136
4137
             1
                       0
4138
                       1
             1
4139
             2
> gr
```

GRanges with 511 ranges and 0 metadata columns:

	${\tt seqnames}$		ranges	strand
	<rle></rle>		<pre><iranges></iranges></pre>	<rle></rle>
chr14:54383433	chr14	[54383433,	54383433]	*
chr14:54383470	chr14	[54383470,	54383470]	*
chr14:54383819	chr14	[54383819,	54383819]	*
chr14:54383945	chr14	[54383945,	54383945]	*
chr14:54384030	chr14	[54384030,	54384030]	*
chr14:54384232	chr14	[54384232,	54384232]	*
chr14:54384251	chr14	[54384251,	54384251]	*
chr14:54384291	chr14	[54384291,	54384291]	*
chr14:54384382	chr14	[54384382,	54384382]	*
chr14:54443495	chr14	[54443495,	54443495]	*
chr14:54443499	chr14	[54443499,	54443499]	*
chr14:54443514	chr14	[54443514,	54443514]	*
chr14:54443831	chr14	[54443831,	54443831]	*

```
chr14:54443880 chr14 [54443880, 54443880] *
chr14:54444016 chr14 [54444016, 54444016] *
chr14:54444124 chr14 [54444124, 54444124] *
chr14:54444385 chr14 [54444385, 54444385] *
chr14:54444750 chr14 [54444750, 54444750] *
---
seqlengths:
chr14
NA
```

## C.2 SummarizedExperiment

We combine three of the key ingredients when we create the SummarizedExperiment object.

## C.3 SNPTrioExperiment

Now, we include the pedigree information as an object of class PedClass. We keep PedClass independent of SNPTrioExperiment for flexibility.

```
> ped <- PedClass(ped.DF)
> ste <- SNPTrioExperiment(se, pedigree = ped)</pre>
Here is the show method.
> ste
class: SNPTrioExperiment
dim: 511 968
exptData(0):
assays(1): geno
rownames(511): chr14:54383433 chr14:54383470 ...
  chr14:54444385 chr14:54444750
rowData metadata column names(0):
colnames(968): H_ME-CSc175_1-CSc175_1_a.1
  H_ME-CSc175_2-CSc175_2 ... H_ME-DS11418_2-DS11418_2
  H_ME-DS11418_3-DS11418_3
colData names(28): vcfid jb_famid ... id ped
pedigree(4139): famid id fid mid sex dx
And now we verify that it is indeed an extension of SummarizedExperiment.
> getClass("SNPTrioExperiment")
Class "SNPTrioExperiment" [package "trioClasses"]
Slots:
                                                      exptData
Name:
                          pedigree
Class:
                          PedClass
                                                    SimpleList
```

Name: rowData colData Class: GenomicRangesORGRangesList DataFrame

Name: assays Class: Assays

Extends: "SummarizedExperiment"

#### C.4 PedClass

Now we investigate the pedigree slot of the SNPTrioExperiment object.

```
> class(pedigree(ste))
```

```
[1] "PedClass"
attr(,"package")
[1] "trioClasses"
```

> getClass("PedClass")

Class "PedClass" [package "trioClasses"]

#### Slots:

Name: rownames nrows listData Class: characterORNULL integer list

Name: elementType elementMetadata metadata Class: character DataTableORNULL list

#### Extends:

Class "DataFrame", directly

Class "DataTable", by class "DataFrame", distance 2

Class "SimpleList", by class "DataFrame", distance 2

Class "DataTableORNULL", by class "DataFrame", distance 3

Class "List", by class "DataFrame", distance 3

Class "Vector", by class "DataFrame", distance  $\boldsymbol{4}$ 

Class "Annotated", by class "DataFrame", distance 5

### > pedigree(ste)

## ${\tt PedClass\ with\ 4139\ rows\ and\ 6\ columns}$

	famid	id
	<factor></factor>	<factor></factor>
1	4778	H_ME-4778_1-4778_1.2
2	4778	H_ME-4778_2-4778_2.2
3	4778	H_ME-4778_3-4778_3.2
4	4783	H_ME-4783_1-4783_1.1
5	4783	H_ME-4783_2-4783_2.1
6	4783	H_ME-4783_3-4783_3.1
7	20000492	H_ME-20000492_1-20000492_1.1

```
8
     20000492 H_ME-20000492_2-20000492_2_a.1
9
     20000492
               H_ME-20000492_3-20000492_3.1
4131 DS12332
                    H_ME-DS12332_1-DS12332_1
4132 DS12332
                    H_ME-DS12332_2-DS12332_2
4133 DS12332
                    H_ME-DS12332_3-DS12332_3
4134 DS99998
                  H_ME-DS10193_5-DS10193_5.2
4135 DS99998
                    H_ME-DS10193_6-DS10193_6
4136 DS99998
                    H_ME-DS10193_7-DS10193_7
                  H_ME-DS10707_4-DS10707_4.2
4137 DS99999
4138 DS99999
                    H_ME-DS10707_5-DS10707_5
                    H_ME-DS10707_6-DS10707_6
4139 DS99999
                                 fid
                                                               mid
                            <factor>
                                                          <factor>
1
               H_ME-4778_2-4778_2.2
                                             H_ME-4778_3-4778_3.2
2
                                                                NA
3
                                  NA
                                                                NA
4
               H_ME-4783_2-4783_2.1
                                             H_ME-4783_3-4783_3.1
5
                                  NA
                                                                NA
6
                                  NA
7
     H_ME-20000492_2-20000492_2_a.1 H_ME-20000492_3-20000492_3.1
8
                                  NA
                                                                NA
9
                                  NA
                                                                NA
                                 . . .
. . .
                                                                . . .
4131
           H_ME-DS12332_2-DS12332_2
                                         H_ME-DS12332_3-DS12332_3
4132
                                  NA
                                                                NA
4133
                                  NA
                                                                NA
4134
           H_ME-DS10193_6-DS10193_6
                                         H_ME-DS10193_7-DS10193_7
4135
                                  NA
4136
                                  NA
                                                                NA
4137
                                  NA
                                                                NA
4138
         H_ME-DS10707_4-DS10707_4.2
                                         H_ME-DS10707_6-DS10707_6
4139
                    dx
          sex
     <factor> <factor>
            1
1
                      1
2
            1
                      0
3
            2
                      0
            2
4
                      1
5
            1
                      0
6
            2
                      1
7
            2
                      1
            1
8
                      0
            2
                      0
9
            1
4131
                      1
4132
            1
                      0
            2
                      0
4133
4134
            2
                      1
            1
4135
                      0
            2
4136
                      0
```

```
4137
     1
                    0
4138
           1
                    1
4139
           2
C.5
      geno accessor
> class(geno(ste))
[1] "SnpMatrix"
attr(,"package")
[1] "snpStats"
> getClass("SnpMatrix")
Class "SnpMatrix" [package "snpStats"]
Slots:
Name: .Data
Class: matrix
Extends:
Class "matrix", from data part
Class "array", by class "matrix", distance 2
Class "mMatrix", by class "matrix", distance 2
Class "structure", by class "matrix", distance 3
Class "vector", by class "matrix", distance 4, with explicit coerce
Known Subclasses: "XSnpMatrix"
> geno(ste)
A SnpMatrix with 968 rows and 511 columns
```

Row names: H\_ME-CSc175\_1-CSc175\_1\_a.1 ... H\_ME-DS11418\_3-DS11418\_3

#### C.6 RowData

> rowData(ste)

GRanges with 511 ranges and 0 metadata columns:

Col names: chr14:54383433 ... chr14:54444750

	seqnames		ranges	strand
	<rle></rle>		Ranges	<rle></rle>
chr14:54383433	chr14	[54383433,	54383433]	*
chr14:54383470	chr14	[54383470,	54383470]	*
chr14:54383819	chr14	[54383819,	54383819]	*
chr14:54383945	chr14	[54383945,	54383945]	*
chr14:54384030	chr14	[54384030,	54384030]	*
chr14:54384232	chr14	[54384232,	54384232]	*
chr14:54384251	chr14	[54384251,	54384251]	*
chr14:54384291	chr14	[54384291,	54384291]	*

```
chr14:54384382 chr14 [54384382, 54384382] *
... ... ... ... ... ... ...
chr14:54443495 chr14 [54443495, 54443495] *
chr14:54443499 chr14 [54443499, 54443499] *
chr14:54443514 chr14 [54443514, 54443514] *
chr14:54443831 chr14 [54443831, 54443831] *
chr14:54443880 chr14 [54443880, 54443880] *
chr14:54444016 chr14 [54444016, 54444016] *
chr14:54444124 chr14 [54444124, 54444124] *
chr14:54444385 chr14 [54444385, 54444385] *
chr14:54444750 chr14 [54444750, 54444750] *
---
seqlengths:
chr14
NA
```

## C.7 ColData

# D Validity