

# Package ‘yriMulti’

October 18, 2017

**Title** support for expression, methylation, DHS, VCF for YRI

**Version** 0.6.0

**Author** VJ Carey <stvjc@channing.harvard.edu>

**Description** expression, methylation, DHS for YRI

**Suggests** erma, BiocStyle, knitr, rmarkdown

**Depends** gQTLBase, SummarizedExperiment, GenomicRanges, Homo.sapiens, dsQTL, geuvPack, MultiAssayExperiment

**Maintainer** VJ Carey <stvjc@channing.harvard.edu>

**License** Artistic-2.0

**LazyLoad** yes

**VignetteBuilder** knitr

**NeedsCompilation** no

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yriMulti-package      *support for expression, methylation, DHS, VCF for YRI*

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## Description

expression, methylation, DHS for YRI

## Details

The DESCRIPTION file:

Package:	yriMulti
Title:	support for expression, methylation, DHS, VCF for YRI
Version:	0.6.0
Author:	VJ Carey <stvjc@channing.harvard.edu>
Description:	expression, methylation, DHS for YRI
Suggests:	erma, BiocStyle, knitr, rmarkdown
Depends:	gQTLBase, SummarizedExperiment, GenomicRanges, Homo.sapiens, dsQTL, geuvPack, MultiAssay
Maintainer:	VJ Carey <stvjc@channing.harvard.edu>
License:	Artistic-2.0
LazyLoad:	yes
VignetteBuilder:	knitr

Index of help topics:

banovichSE	Illumina 450k methylation assay applied to Yoruba cell lines
mexGR	combine expression data for a gene with nearby methylation data
pwplot	pairwise plot for components of MultiAssayExperiment
yriMulti-package	support for expression, methylation, DHS, VCF for YRI

This is a relatively experimental package that is used to exercise aspects of MultiAssayExperiment and gQTLstats. The primary innovations are illustration of VcfStack class of GenomicFiles as a distributed on-disk (or in-cloud) resource for assay elements representing genotypes, and adoption of MultiAssayExperiment to unite relatively uncommonly integrated assays such as DnaseI hypersensitivity and 450k methylation.

## Author(s)

VJ Carey <stvjc@channing.harvard.edu>  
Maintainer: VJ Carey <stvjc@channing.harvard.edu>

## Examples

```
# see the vignette
```

banovichSE

*Illumina 450k methylation assay applied to Yoruba cell lines*

## Description

Illumina 450k methylation assay applied to Yoruba cell lines

**Usage**

```
data("banovichSE")
```

**Format**

The format is:

```
Formal class 'RangedSummarizedExperiment' [package "SummarizedExperiment"] with 6 slots
..@ rowRanges :Formal class 'GRanges' [package "GenomicRanges"] with 6 slots
...@ seqnames :Formal class 'Rle' [package "S4Vectors"] with 4 slots
...@ values : Factor w/ 24 levels "chr1","chr2",..: 16 1 8 14 1 15 19 3 12 15 ...
...@ lengths : int [1:301401] 1 1 1 1 1 1 1 1 1 ...
...@ elementMetadata: NULL
...@ metadata : list()
...@ ranges :Formal class 'IRanges' [package "IRanges"] with 6 slots
...@ start : int [1:329469] 53468112 91194674 42263294 69341139 230560793 23034447
54695678 128902377 124086477 59785306 ...
...@ width : int [1:329469] 2 2 2 2 2 2 ...
...@ NAMES : chr [1:329469] "cg00000029" "cg00000165" "cg00000236" "cg00000289"
...
...@ elementType : chr "integer"
...@ elementMetadata: NULL
...@ metadata : list()
...@ strand :Formal class 'Rle' [package "S4Vectors"] with 4 slots
...@ values : Factor w/ 3 levels "+","-","*": 3
...@ lengths : int 329469
...@ elementMetadata: NULL
...@ metadata : list()
...@ elementMetadata:Formal class 'DataFrame' [package "S4Vectors"] with 6 slots
...@ rownames : NULL
...@ nrow : int 329469
...@ listData :List of 10
...$ addressA : chr [1:329469] "14782418" "12637463" "12649348" "18766346" ...
...$ addressB : chr [1:329469] "" "" "" ...
...$ channel :Formal class 'Rle' [package "S4Vectors"] with 4 slots
...@ values : Factor w/ 3 levels "Both","Grn","Red": 1 3 1 2 1 2 1 3 1 2 ...
...@ lengths : int [1:140453] 5 1 2 1 3 1 1 1 2 1 ...
...@ elementMetadata: NULL
...@ metadata : list()
...$ platform :Formal class 'Rle' [package "S4Vectors"] with 4 slots
...@ values : Factor w/ 2 levels "BOTH","HM450": 2 1 2 1 2 1 2 1 ...
...@ lengths : int [1:35429] 38 1 31 1 41 1 19 1 8 1 ...
...@ elementMetadata: NULL
...@ metadata : list()
...$ percentGC : num [1:329469] 0.46 0.48 0.5 0.44 0.42 0.8 0.6 0.58 0.78 0.54 ...
...$ sourceSeq :Formal class 'DNAStringSet' [package "Biostrings"] with 5 slots
...@ pool :Formal class 'SharedRaw_Pool' [package "XVector"] with 2 slots
...@ xp_list :List of 1
...$ <externalptr>
...@ .link_to_cached_object_list:List of 1
...$ <environment: 0x7f87a4476990>
...@ ranges :Formal class 'GroupedIRanges' [package "XVector"] with 7 slots
...@ group : int [1:329469] 1 1 1 1 1 1 1 1 ...
```

```

.....@ start : int [1:329469] 1 151 201 251 401 451 551 651 701 851 ...
.....@ width : int [1:329469] 50 50 50 50 50 50 50 50 50 50 ...
.....@ NAMES : NULL
.....@ elementType : chr "integer"
.....@ elementMetadata: NULL
.....@ metadata : list()
.....@ elementType : chr "DNAString"
.....@ elementMetadata: NULL
.....@ metadata : list()
.....$ probeType :Formal class 'Rle' [package "S4Vectors"] with 4 slots
.....@ values : Factor w/ 3 levels "cg","ch","rs": 1 2
.....@ lengths : int [1:2] 327286 2183
.....@ elementMetadata: NULL
.....@ metadata : list()
.....$ probeStart : chr [1:329469] "53468112" "91194626" "42263246" "69341139" ...
.....$ probeEnd : chr [1:329469] "53468161" "91194675" "42263295" "69341188" ...
.....$ probeTarget: num [1:329469] 5.35e+07 9.12e+07 4.23e+07 6.93e+07 2.31e+08 ...
.....@ elementType : chr "ANY"
.....@ elementMetadata: NULL
.....@ metadata : list()
.....@ seqinfo :Formal class 'Seqinfo' [package "GenomeInfoDb"] with 4 slots
.....@ seqnames : chr [1:24] "chr1" "chr2" "chr3" "chr4" ...
.....@ seqlengths : int [1:24] 249250621 243199373 198022430 191154276 180915260
171115067 159138663 146364022 141213431 135534747 ...
.....@ is_circular: logi [1:24] FALSE FALSE FALSE FALSE FALSE FALSE FALSE ...
.....@ genome : chr [1:24] "hg19" "hg19" "hg19" "hg19" ...
.....@ metadata :List of 1
.....$ :Formal class 'DataFrame' [package "S4Vectors"] with 6 slots
.....@ rownames : NULL
.....@ nrows : int 13
.....@ listData :List of 2
.....$ name : chr [1:13] "Db type" "Supporting package" "data_nrow" "Db created by" ...
.....$ value: chr [1:13] "FeatureDb" "GenomicFeatures" "487173" "GenomicFeatures
package from Bioconductor" ...
.....@ elementType : chr "ANY"
.....@ elementMetadata: NULL
.....@ metadata : list()
@ colData :Formal class 'DataFrame' [package "S4Vectors"] with 6 slots
.....@ rownames : chr [1:64] "NA18498" "NA18499" "NA18501" "NA18502" ...
.....@ nrows : int 64
.....@ listData :List of 35
.....$ title : Factor w/ 64 levels "GM18489","GM18498",...: 2 3 4 5 11 12 13 14 19 20 ...
.....- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
.....$ geo_accession : Factor w/ 64 levels "GSM1383567","GSM1383568",...: 1 2 3 4 5 6 7 8 9
10 ...
.....- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
.....$ status : Factor w/ 1 level "Public on May 09 2014": 1 1 1 1 1 1 1 1 1 1 ...
.....- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
.....$ submission_date : Factor w/ 1 level "May 08 2014": 1 1 1 1 1 1 1 1 1 ...
.....- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
.....$ last_update_date : Factor w/ 1 level "Aug 12 2014": 1 1 1 1 1 1 1 1 1 ...
.....- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...

```

```

... . . . . $ type : Factor w/ 1 level "genomic": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ channel_count : Factor w/ 1 level "1": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ source_name_ch1 : Factor w/ 1 level "HapMap_Yoruba": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ organism_ch1 : Factor w/ 1 level "Homo sapiens": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ characteristics_ch1 : Factor w/ 2 levels "gender: Female", ...: 2 1 2 1 2 1 2 1 2 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ characteristics_ch1.1 : Factor w/ 1 level "cell type: Lymphoblastoid Cell Line": 1 1 1 1 1
1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ molecule_ch1 : Factor w/ 1 level "genomic DNA": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ extract_protocol_ch1 : Factor w/ 1 level "DNA was extracted from lymphoblastoid Cell
Line": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ label_ch1 : Factor w/ 1 level "cy3 and cy5": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ label_protocol_ch1 : Factor w/ 1 level "Standard illumina protocols": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ taxid_ch1 : Factor w/ 1 level "9606": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ hyb_protocol : Factor w/ 1 level "The samples were bisulphite-converted and hybridized
to the Infinium HumanMethylation450 BeadChip at the University of Chicago" | __truncated__: 1
1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ scan_protocol : Factor w/ 1 level "not provided": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ description : Factor w/ 64 levels "Sample 1", "Sample 10", ...: 1 12 23 34 45 56 62 63 64 2
...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ data_processing : Factor w/ 1 level "To ensure high data quality, probes were mapped to a
bisulfite converted genome and only uniquely mapped probes were retained. | __truncated__: 1 1
1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ data_processing.1 : Factor w/ 1 level "Normalized data: Normalized average beta (PCs
removed)": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ data_processing.2 : Factor w/ 1 level "Un-normalized data: Unmethylated and methylated
signal intensities and detection p-value": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ platform_id : Factor w/ 1 level "GPL13534": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ contact_name : Factor w/ 1 level "Nicholas,E,Banovich": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ contact_laboratory : Factor w/ 1 level "Gilad": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ contact_department : Factor w/ 1 level "Human Genetics": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ contact_institute : Factor w/ 1 level "University of Chicago": 1 1 1 1 1 1 1 1 1 ...

```

```

... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ contact_address : Factor w/ 1 level "920 E. 58th Street, CLSC 317": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ contact_city : Factor w/ 1 level "Chicago": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ contact_state : Factor w/ 1 level "IL": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ contact_zip.postal_code: Factor w/ 1 level "60453": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ contact_country : Factor w/ 1 level "USA": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ supplementary_file : Factor w/ 1 level "NONE": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ data_row_count : Factor w/ 1 level "329469": 1 1 1 1 1 1 1 1 1 ...
... . . . .- attr(*, "names")= chr [1:64] "V2" "V3" "V4" "V5" ...
... . . . .$ naid : chr [1:64] "NA18498" "NA18499" "NA18501" "NA18502" ...
... . . . @ elementType : chr "ANY"
... . . . @ elementMetadata: NULL
... . . . @ metadata : list()
... @ assays :Reference class 'ShallowSimpleListAssays' [package "GenomicRanges"] with 1 field
... . $ data: NULL
... and 14 methods.
... @ NAMES : NULL
... @ elementMetadata:Formal class 'DataFrame' [package "S4Vectors"] with 6 slots
... . . @ rownames : NULL
... . . @ nrows : int 329469
... . . @ listData : Named list()
... . . . @ elementType : chr "ANY"
... . . . @ elementMetadata: NULL
... . . . @ metadata : list()
.. @ metadata : list()

```

## Examples

```
data(banovichSE)
```

mexGR

*combine expression data for a gene with nearby methylation data*

## Description

combine expression data for a gene with nearby methylation data,

## Usage

```
mexGR(methSE, exprSE, gradius = 0, symbol = "ORMDL3", etagname = "gene_name")
```

*pwpplot*

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### Arguments

methSE  
exprSE  
gradius  
symbol  
etagnome

### Value

GRanges instance with assay results in mcols

### Examples

```
data(geuFPKM)
data(banovichSE)
m1 = mexGR(banovichSE, geuFPKM, gradius=5000)
ii = bindelms(geuFPKM, banovichSE)
```

---

*pwpplot*

*pairwise plot for components of MultiAssayExperiment*

---

### Usage

```
pwpplot(fmla1, fmla2, mae, ytx = force, xtx = force, ...)
```

### Arguments

fmla1            a formula with one name in each of lhs and rhs. these are the names of the experiments to be extracted  
fmla2            a formula with lhs the name of a feature in rowData(lhs(fmla1)) and rhs the name of a feature in rowData(rhs(fmla1)). these are the features to be plotted  
mae              instance of MultiAssayExperiment  
ytx              function to transform lhs(fmla2)  
xtx              function to transform rhs(fmla2)  
...                passed to plot

### Value

runs plot

### Author(s)

VJ Carey <stvjc@channing.harvard.edu>

### Examples

```
data(demoMAE)
pwpplot(geuvRNaseq~yri450k, ENSG00000139618.9~cg20073910,
demoMAE, ytx=log, main="demo")
```

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