

Package ‘BubbleTree’

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Type Package

Title BubbleTree: an intuitive visualization to elucidate tumoral aneuploidy and clonality in somatic mosaicism using next generation sequencing data

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License LGPL (>= 3)

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all.somatic.lst	<i>all.somatic.lst</i>
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Description

A dataset containing pre-calculated BAF scores for annotated SNVs.

Format

S4 object with seqnames, genomic ranges, strand, BAF score

Source

internal

`allCall.lst`*allCall.lst*

Description

A dataset containing precalculated data from CNV segment analysis.

Format

S4 object with rbd, rbd.adj, results

Source

internal

`allCNV.lst`*allCNV.lst*

Description

A dataset containing pre-calculated segment calls.

Format

S4 object with seqnames, genomic ranges, num.mark, score

Source

internal

`allHetero.lst`*allHetero.lst*

Description

S4 GRanges dataset containing pre-calculated heterozygosity data.

Format

S4

Source

internal

allRBD.lst

allRBD.lst

Description

A dataset containing precalculated data from CNV segment analysis.

Format

S4 object with rbd, rbd.adj

Source

internal

annoByGenesAndCyto

annoByGenesAndCyto

Description

get annotation for genes and cytobands

Usage

```
annoByGenesAndCyto(.Object, chr, beg, end, critical.genes, gene.uni.clean.gr,
  cyto.gr)
```

```
## S4 method for signature 'Annotate'
annoByGenesAndCyto(.Object, chr, beg, end, critical.genes,
  gene.uni.clean.gr, cyto.gr)
```

Arguments

.Object	the objet
chr	the chromosome
beg	genomic start coord
end	genomic end coord
critical.genes	set of critical genes
gene.uni.clean.gr	gr object of genes
cyto.gr	gr object of cyto positions

Value

list of annotation for genes and cytobands

Examples

```
load(system.file("data", "allCall.lst.RData", package="BubbleTree"))
load(system.file("data", "cancer.genes.minus2.rda", package="BubbleTree"))
load(system.file("data", "vol.genes.rda", package="BubbleTree"))
load(system.file("data", "gene.uni.clean.gr.rda", package="BubbleTree"))
load(system.file("data", "cyto.gr.rda", package="BubbleTree"))

comm <- btcompare(vol.genes, cancer.genes.minus2)
btreeplotter <- new("BTreePlotter", branch.col="gray50")
annotator <-new("Annotate")
nn <- "sam2"
cc <- allCall.lst[[nn]]
z <- drawBTree(btreeplotter, cc@rbd.adj) +
  ggplot2::labs(title=sprintf("%s (%s)", nn, info(cc)))
out <- cc@result$dist %>%
  filter(seg.size >= 0.1 ) %>%
  arrange(gtools::mixedorder(as.character(seqnames)), start)

ann <- annoByGenesAndCyto(annotator,
  as.character(out$seqnames),
  as.numeric(out$start),
  as.numeric(out$end),
  comm$comm,
  gene.uni.clean.gr=gene.uni.clean.gr,
  cyto.gr=cyto.gr)
```

Annotate	<i>Annotate</i>
----------	-----------------

Description

Annotate

Examples

```
annotate <- new("Annotate")
```

bafTrack	<i>bafTrack</i>
----------	-----------------

Description

get the BAF track

Usage

```
bafTrack(.Object, result.dat, gr2, somatic.gr = NULL, min.prev = 0.15,
  cex = 1.2)

## S4 method for signature 'TrackPlotter'
bafTrack(.Object, result.dat, gr2, somatic.gr = NULL,
  min.prev = 0.15, cex = 1.2)
```

Arguments

<code>.Object</code>	the object
<code>result.dat</code>	the result dataframe
<code>gr2</code>	the gr2 object
<code>somatic.gr</code>	somatic gr object annotation
<code>min.prev</code>	previous min
<code>cex</code>	the cex

Value

the highlighted BAF track

Examples

```
load(system.file("data", "allCall.lst.RData", package="BubbleTree"))
load(system.file("data", "centromere.dat.rda", package="BubbleTree"))
load(system.file("data", "all.somatic.lst.RData", package="BubbleTree"))
load(system.file("data", "hg19.seqinfo.rda", package="BubbleTree"))

trackplotter <- new("TrackPlotter")
gr2 = centromere.dat
nn <- "sam2"
p2 <- bafTrack(trackplotter,
               result.dat=allCall.lst[[nn]]@result,
               gr2=gr2,
               somatic.gr=all.somatic.lst[[nn]])
```

btcompare

btcompare

Description

btcompare

Usage

```
btcompare(set1, set2)
```

Arguments

<code>set1</code>	first set
<code>set2</code>	second set to compare

Value

combined, unique list of genes

Examples

```
load(system.file("data", "cancer.genes.minus2.rda", package="BubbleTree"))
load(system.file("data", "vol.genes.rda", package="BubbleTree"))

# 77 common cancer genes
comm <- btcompare(vol.genes, cancer.genes.minus2)
```

btpredict

*btpredict***Description**

btpredict

Usage

```
btpredict(.Object)

## S4 method for signature 'BTreePredictor'
btpredict(.Object)
```

Arguments

.Object the object

Value

.Object populated with the predictions

Examples

```
load(system.file("data", "allRBD.lst.RData", package="BubbleTree"))

btreepredictor <- new("BTreePredictor")
btreepredictor@config$cutree.h <- 0.15
high.ploidy <- rep(TRUE, length(allRBD.lst))
high.purity <- rep(TRUE, length(allRBD.lst))

high.ploidy[c("sam6",
              "ovary.wgs",
              "ovary.wes",
              "TCGA-06-0145-01A-01W-0224-08",
              "TCGA-13-1500-01A-01D-0472-01",
              "TCGA-A0-A0JJ-01A-11W-A071-09")] <- FALSE

high.purity[c("sam6", "ovary.wgs", "ovary.wes")] <- FALSE

rbd <- allRBD.lst[["sam6"]]
btreepredictor@config$high.ploidy <- high.ploidy["sam6"]
btreepredictor@config$high.purity <- high.purity["sam6"]
btreepredictor <- loadRBD(btreepredictor, rbd)
btreepredictor@config$min.segSize <- ifelse(max(btreepredictor@rbd$seg.size,
                                                na.rm=TRUE) < 0.4, 0.1, 0.4)

btreepredictor <- btpredict(btreepredictor)
cat(info(btreepredictor), "\n")
```

BTreePlotter	<i>BTreePlotter</i>
Description BTreePlotter	
Examples btreeplotter <- new("BTreePlotter")	
BTreePredictor	<i>BTreePredictor</i>
Description BTreePredictor	
Examples btreepredictor <- new("BTreePredictor")	
cancer.genes.minus2	<i>cancer.genes.minus2.rda</i>
Description A dataset containing a list of known cancer genes.	
Format list	
Source internal	
centromere.dat	<i>centromere.dat</i>
Description A dataset containing an annotated list of centromere locations.	
Format list	
Source internal	

`cnv.gr`*cnv.gr*

Description

S4 GRanges object containing data on chromosomal locations with seqnames, genomic range, strand, name

Format

S4

Source

internal

`cyto.gr`*cyto.gr*

Description

S4 GRanges object containing data on chromosomal locations with seqnames, genomic range, strand, name, gieStain.

Format

S4

Source

internal

`drawBTree`*drawBTree*

Description

draw the BTree track

Usage

```
drawBTree(.Object, rbd, size = 1)
```

```
## S4 method for signature 'BTreePlotter'  
drawBTree(.Object, rbd, size = 1)
```

Arguments

.Object	the object
rbd	the rbd object
size	the size

Value

draw the BTree track

Examples

```
load(system.file("data", "allCall.lst.RData", package="BubbleTree"))
load(system.file("data", "cancer.genes.minus2.rda", package="BubbleTree"))
load(system.file("data", "vol.genes.rda", package="BubbleTree"))
load(system.file("data", "gene.uni.clean.gr.rda", package="BubbleTree"))
load(system.file("data", "cyto.gr.rda", package="BubbleTree"))

# 77 common cancer genes
comm <- btcompare(vol.genes, cancer.genes.minus2)

btreeplotter <- new("BTreePlotter", branch.col="gray50")
annotator <-new("Annotate")
cc <- allCall.lst[["sam2"]]
z <- drawBTree(btreeplotter, cc@rbd.adj) +
  ggplot2::labs(title=sprintf("%s (%s)", "sam2", info(cc)))
```

drawBubbles

drawBubbles

Description

draw the Bubbles

Usage

```
drawBubbles(.Object, rbd, col = NULL)

## S4 method for signature 'BTreePlotter'
drawBubbles(.Object, rbd, col = "gray80")
```

Arguments

.Object	the object
rbd	the rbd object
col	the col value

Value

draw the bubbles on the track

Examples

```
load(system.file("data", "allCall.lst.RData", package="BubbleTree"))

btreeplotter <- new("BTreePlotter", max.ploidy=5, max.size=10)
nn <- "sam2"
rbd1 <- allCall.lst[[nn]]@rbd
rbd2 <- allCall.lst[[nn]]@rbd.adj
arrows <- trackBTree(btreeplotter, rbd1, rbd2, min.srcSize=0.01,
                     min.trtSize=0.01)
btree <- drawBTree(btreeplotter, rbd1) +
  drawBubbles(btreeplotter, rbd2, "gray80") + arrows
```

drawFeatures

*drawFeatures***Description**

draw the features

Usage

```
drawFeatures(.Object, rbd, col = NULL)

## S4 method for signature 'BTreePlotter'
drawFeatures(.Object, rbd, col = "black")
```

Arguments

.Object	the object
rbd	the rbd object
col	the col value

Value

draw the annotation on the track

Examples

```
load(system.file("data", "allCall.lst.RData", package="BubbleTree"))
load(system.file("data", "cancer.genes.minus2.rda", package="BubbleTree"))
load(system.file("data", "vol.genes.rda", package="BubbleTree"))
load(system.file("data", "gene.uni.clean.gr.rda", package="BubbleTree"))
load(system.file("data", "cyto.gr.rda", package="BubbleTree"))

# 77 common cancer genes merged from 2 sets
comm <- btcompare(vol.genes, cancer.genes.minus2)

btreeplotter <- new("BTreePlotter", branch.col="gray50")
annotator <- new("Annotate")

nn <- "sam12"
cc <- allCall.lst[[nn]]
z <- drawBTree(btreeplotter, cc@rbd.adj) +
```

```
ggplot2::labs(title=sprintf("%s (%s)", nn, info(cc)))
out <- cc@result$dist %>% filter(seg.size >= 0.1 ) %>%
  arrange(gtools::mixedorder(as.character(seqnames)), start)

ann <- with(out, {
  annoByGenesAndCyto(annotator,
    as.character(out$seqnames),
    as.numeric(out$start),
    as.numeric(out$end),
    comm$comm,
    gene.uni.clean.gr=gene.uni.clean.gr,
    cyto.gr=cyto.gr)
})

out$cyto <- ann$cyto
out$genes <- ann$ann
v <- z + drawFeatures(btreeplotter, out)
print(v)
```

gene.uni.clean.gr	<i>gene.uni.clean.gr</i>
-------------------	--------------------------

Description

S4 GRanges object containing human gene annotation with seqnames, genomic coordinates, stand, gene.sybmol.

Format

S4

Source

internal

getTracks	<i>getTracks</i>
-----------	------------------

Description

get all tracks

Usage

```
getTracks(p1, p2, title = "")
```

Arguments

p1	set 1
p2	set 2
title	the title

Value

all of the requested tracks

Examples

```
load(system.file("data", "allCall.lst.RData", package="BubbleTree"))
load(system.file("data", "centromere.dat.rda", package="BubbleTree"))
load(system.file("data", "all.somatic.lst.RData", package="BubbleTree"))
load(system.file("data", "hg19.seqinfo.rda", package="BubbleTree"))

trackplotter <- new("TrackPlotter")
gr2 = centromere.dat
nn <- "sam2"
ymax <- ifelse(nn %in% c("lung.wgs", "lung.wes"), 9, 4.3)
p1 <- xyTrack(trackplotter,
              result.dat=allCall.lst[[nn]]@result,
              gr2=gr2,
              ymax=ymax) + ggplot2::labs(title=nn)

p2 <- bafTrack(trackplotter,
              result.dat=allCall.lst[[nn]]@result,
              gr2=gr2,
              somatic.gr=all.somatic.lst[[nn]])

t1 <- getTracks(p1, p2)
```

heteroLociTrack

heteroLociTrack

Description

get the heteroLoci track

Usage

```
heteroLociTrack(.Object, result.dat, gr2, hetero.gr = NULL, min.prev = 0.15,
               ymax = 4.3, cex = 0.5)
```

```
## S4 method for signature 'TrackPlotter'
```

```
heteroLociTrack(.Object, result.dat, gr2,
               hetero.gr = NULL, min.prev = 0.15, ymax = 4.3, cex = 0.5)
```

Arguments

.Object	the object
result.dat	the results
gr2	the gr2 object
hetero.gr	hetero annotation
min.prev	previous min
ymax	max y
cex	the cex

Value

the highlightted heterozygosity track

Examples

```
load(system.file("data", "allCall.lst.RData", package="BubbleTree"))
load(system.file("data", "centromere.dat.rda", package="BubbleTree"))
load(system.file("data", "allHetero.lst.RData", package="BubbleTree"))
load(system.file("data", "hg19.seqinfo.rda", package="BubbleTree"))

trackplotter <- new("TrackPlotter")
gr2 = centromere.dat
nn <- "sam2"
z1 <- heteroLociTrack(trackplotter, allCall.lst[[nn]]@result,
                      gr2, allHetero.lst[[nn]])
```

hg19.seqinfo	<i>hg19.seqinfo.Rd</i>
--------------	------------------------

Description

Seqinfo object containing names and lengths of each chromosome of the human genome.

Format

Seqinfo

Source

internal

info	<i>info</i>
------	-------------

Description

info

Usage

```
info(.Object)

## S4 method for signature 'BTreePredictor'
info(.Object)
```

Arguments

.Object the object

Value

print out info of prediction data

Examples

```
load(system.file("data", "allRBD.lst.RData", package="BubbleTree"))

btrepredictor <- new("BTreePredictor")
btrepredictor@config$cutree.h <- 0.15

high.ploidy <- rep(TRUE, length(allRBD.lst))
high.purity <- rep(TRUE, length(allRBD.lst))

high.ploidy[c("sam6",
              "ovary.wgs",
              "ovary.wes",
              "TCGA-06-0145-01A-01W-0224-08",
              "TCGA-13-1500-01A-01D-0472-01",
              "TCGA-AO-A0JJ-01A-11W-A071-09")] <- FALSE

high.purity[c("sam6", "ovary.wgs", "ovary.wes")] <- FALSE

nn <- "sam6"

rbd <- allRBD.lst[[nn]]
btrepredictor@config$high.ploidy <- high.ploidy[nn]
btrepredictor@config$high.purity <- high.purity[nn]
btrepredictor <- loadRBD(btrepredictor, rbd)
btrepredictor@config$min.segSize <- ifelse(max(btrepredictor@rbd$seg.size,
                                              na.rm=TRUE) < 0.4, 0.1, 0.4)

btrepredictor <- btpredict(btrepredictor)
cat(info(btrepredictor), "\n")
```

loadRBD

*loadRBD***Description**

load the RBD data

Usage

```
loadRBD(.Object, rbd, total.mark = NA)

## S4 method for signature 'BTreePredictor'
loadRBD(.Object, rbd, total.mark = NA)
```

Arguments

.Object	the object
rbd	rbd object
total.mark	total mark

Value

.Object populated with the RBD list with updated segment size

Examples

```
load(system.file("data", "allRBD.lst.RData", package="BubbleTree"))

btrepredictor <- new("BTreePredictor")
btrepredictor@config$cutree.h <- 0.15

high.ploidy <- rep(TRUE, length(allRBD.lst))
high.purity <- rep(TRUE, length(allRBD.lst))

high.ploidy[c("sam6",
              "ovary.wgs",
              "ovary.wes",
              "TCGA-06-0145-01A-01W-0224-08",
              "TCGA-13-1500-01A-01D-0472-01",
              "TCGA-AO-A0JJ-01A-11W-A071-09")] <- FALSE

high.purity[c("sam6", "ovary.wgs", "ovary.wes")] <- FALSE

nn <- "sam6"

rbd <- allRBD.lst[[nn]]
btrepredictor@config$high.ploidy <- high.ploidy[nn]
btrepredictor@config$high.purity <- high.purity[nn]
btrepredictor <- loadRBD(btrepredictor, rbd)
```

makeRBD

makeRBD

Description

make the RBD object

Usage

```
makeRBD(.Object, ...)

## S4 method for signature 'RBD'
makeRBD(.Object, snp.gr, cnv.gr, unimodal.kurtosis = -0.1)
```

Arguments

.Object	the object
...	other input (not needed)
snp.gr	SNP GenomicRanges object
cnv.gr	CNV GenomicRanges object
unimodal.kurtosis	kurtosis

Value

RBD object

Examples

```
# load sample files
load(system.file("data", "cnv.gr.rda", package="BubbleTree"))
load(system.file("data", "snp.gr.rda", package="BubbleTree"))

# load annotations
load(system.file("data", "centromere.dat.rda", package="BubbleTree"))
load(system.file("data", "cyto.gr.rda", package="BubbleTree"))
load(system.file("data", "cancer.genes.minus2.rda", package="BubbleTree"))
load(system.file("data", "vol.genes.rda", package="BubbleTree"))
load(system.file("data", "gene.uni.clean.gr.rda", package="BubbleTree"))

# initialize RBD object
r <- new("RBD", unimodal.kurtosis=-0.1)

# create new RBD object with GenomicRanges objects for SNPs and CNVs
rbd <- makeRBD(r, snp.gr, cnv.gr)
head(rbd)

# create a new prediction
btrepredictor <- new("BTreePredictor", rbd=rbd, max.ploidy=6, prev.grid=seq(0.2,1, by=0.01))
pred <- btpredict(btrepredictor)

# create rbd plot
btreetplotter <- new("BTreePlotter", max.ploidy=5, max.size=10)
btree <- drawBTree(btreetplotter, pred@rbd)
print(btree)

# create rbd.adj plot
btreetplotter <- new("BTreePlotter", branch.col="gray50")
btree <- drawBTree(btreetplotter, pred@rbd.adj)
print(btree)

# create a combined plot with rbd and rbd.adj that shows the arrows indicating change
# THIS IS VERY MESSY WITH CURRENT DATA from Dong
btreetplotter <- new("BTreePlotter", max.ploidy=5, max.size=10)
arrows <- trackBTree(btreetplotter,
                    pred@rbd,
                    pred@rbd.adj,
                    min.srcSize=0.01,
                    min.trtSize=0.01)

btree <- drawBTree(btreetplotter, pred@rbd) + arrows
print(btree)

# create a plot with overlays of significant genes
btreetplotter <- new("BTreePlotter", branch.col="gray50")
annotator <- new("Annotate")

comm <- btcompare(vol.genes, cancer.genes.minus2)
```

```

sample.name <- "22_cnv_snv"

btree <- drawBTree(btreeplotter, pred@rbd.adj) +
  ggplot2::labs(title=sprintf("%s (%s)", sample.name, info(pred)))

out <- pred@result$dist %>%
  filter(seg.size >= 0.1 ) %>%
  arrange(gtools::mixedorder(as.character(seqnames)), start)

ann <- with(out, {
  annoByGenesAndCyto(annotator,
    as.character(out$seqnames),
    as.numeric(out$start),
    as.numeric(out$end),
    comm$comm,
    gene.uni.clean.gr=gene.uni.clean.gr,
    cyto.gr=cyto.gr)
})

out$cyto <- ann$cyto
out$genes <- ann$ann

btree <- btree + drawFeatures(btreeplotter, out)
print(btree)

# print out purity and ploidy values
info <- info(pred)
cat("\nPurity/Ploidy: ", info, "\n")

```

mergeSnpCnv

mergeSnpCnv

Description

merge snp and cnv data

Usage

```
mergeSnpCnv(.Object, snp.gr, cnv.gr)
```

```
## S4 method for signature 'RBD'
```

```
mergeSnpCnv(.Object, snp.gr, cnv.gr)
```

Arguments

.Object	the object
snp.gr	SNP GenomicRanges object
cnv.gr	CNV GenomicRanges object

Value

combined, unique list of genes

RBD	<i>RBD</i>
-----	------------

Description

RBD

Examples

```
rbd <- new("RBD")
```

RscoreTrack	<i>RscoreTrack</i>
-------------	--------------------

Description

get the RScore track

Usage

```
RscoreTrack(.Object, result.dat, gr2, cnv.gr = NULL, min.prev = 0.15,  
            ymax = 3, cex = 1.5)
```

```
## S4 method for signature 'TrackPlotter'  
RscoreTrack(.Object, result.dat, gr2, cnv.gr = NULL,  
            min.prev = 0.15, ymax = 3, cex = 1.5)
```

Arguments

.Object	the object
result.dat	the results
gr2	the gr2 object
cnv.gr	cnv annotation
min.prev	previous min
ymax	max y
cex	the cex

Value

the highlighted RScore track

Examples

```
load(system.file("data", "allCall.lst.RData", package="BubbleTree"))
load(system.file("data", "centromere.dat.rda", package="BubbleTree"))
load(system.file("data", "allCNV.lst.RData", package="BubbleTree"))
load(system.file("data", "hg19.seqinfo.rda", package="BubbleTree"))

gr2 = centromere.dat
trackplotter <- new("TrackPlotter")
nn <- "sam2"
z <- RscoreTrack(trackplotter, allCall.lst[[nn]]@result, gr2, allCNV.lst[[nn]])
```

saveXLS

*saveXLS***Description**

saveXLS

Usage

```
saveXLS(dat.lst, xls.fn, row.names = FALSE, ...)
```

Arguments

dat.lst	dataframe
xls.fn	filename
row.names	row names
...	misc

Value

new Excel file

Examples

```
load(system.file("data", "allCall.lst.RData", package="BubbleTree"))

all.summary <- plyr::ldply(allCall.lst, function(.Object) {
  purity <- .Object@result$prev[1]
  adj <- .Object@result$ploidy.adj["adj"]
  # when purity is low the calculation result is not reliable
  ploidy <- (2*adj -2)/purity + 2

  with(.Object@result,
    return(c(Purity=round(purity,3),
              Prevalences=paste(round(prev,3), collapse=", "),
              "Tumor ploidy"=round(ploidy,1))))
}) %>% plyr::rename(c(".id"="Sample"))

xls.filename <- paste("all_summary", "xlsx", sep=".")
saveXLS(list(Summary=all.summary), xls.filename)
```

`snp.gr`*snp.gr*

Description

S4 GRanges object containing data on chromosomal locations with seqnames, genomic position, strand, name

Format

S4

Source

internal

`trackBTree`*trackBTree*

Description

get the geom_segment location of the BTree track

Usage

```
trackBTree(.Object, rbd1, rbd2, is.matched = FALSE, min.srcSize = 0.5,  
           min.trtSize = 0.1, min.overlap = 1e+05)
```

```
## S4 method for signature 'BTreePlotter'  
trackBTree(.Object, rbd1, rbd2, is.matched = FALSE,  
           min.srcSize = 0.5, min.trtSize = 0.1, min.overlap = 1e+05)
```

Arguments

<code>.Object</code>	the object
<code>rbd1</code>	rbd one
<code>rbd2</code>	rbd two
<code>is.matched</code>	is it matched
<code>min.srcSize</code>	min src size
<code>min.trtSize</code>	min trt size
<code>min.overlap</code>	min overlap

Value

geom_segment location of BTree track

Examples

```
load(system.file("data", "allCall.lst.RData", package="BubbleTree"))

btreeplotter <- new("BTreePlotter", max.ploidy=5, max.size=10)
nn <- "sam2"
rbd1 <- allCall.lst[[nn]]@rbd
rbd2 <- allCall.lst[[nn]]@rbd.adj
arrows <- trackBTree(btreeplotter, rbd1, rbd2, min.srcSize=0.01,
                     min.trtSize=0.01)
btree <- drawBTree(btreeplotter, rbd1) +
         drawBubbles(btreeplotter, rbd2, "gray80") + arrows
```

TrackPlotter	<i>TrackPlotter</i>
--------------	---------------------

Description

TrackPlotter

Examples

```
trackplotter <- new("TrackPlotter")
```

vol.genes	<i>vol.genes</i>
-----------	------------------

Description

A dataset containing a list of known cancer genes.

Format

list

Source

internal

xyTrack

*xyTrack***Description**

get the xy track

Usage

```
xyTrack(.Object, result.dat, gr2, min.prev = 0.15, ymax = 4.3)

## S4 method for signature 'TrackPlotter'
xyTrack(.Object, result.dat, gr2, min.prev = 0.15,
        ymax = 4.3)
```

Arguments

.Object	the object
result.dat	result dataframe
gr2	gr2 object
min.prev	previous min
ymax	the max y

Value

the highlighted xy track

Examples

```
load(system.file("data", "allCall.lst.RData", package="BubbleTree"))
load(system.file("data", "centromere.dat.rda", package="BubbleTree"))
load(system.file("data", "hg19.seqinfo.rda", package="BubbleTree"))

trackplotter <- new("TrackPlotter")
gr2 = centromere.dat
nn <- "sam2"
ymax <- ifelse(nn %in% c("lung.wgs", "lung.wes"), 9, 4.3)
p1 <- xyTrack(trackplotter,
              result.dat=allCall.lst[[nn]]@result,
              gr2=gr2,
              ymax=ymax) + ggplot2::labs(title=nn)
```

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