

# 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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**Description** CNV analysis in groups of tumor samples.

**License** LGPL (>= 3)

**Imports** BiocGenerics (>= 0.31.6), BiocStyle, Biobase, ggplot2,  
WriteXLS, gtools, RColorBrewer, limma, grid, gtable, gridExtra,  
biovizBase, e1071, methods, grDevices, stats, utils

**Depends** R (>= 3.5), IRanges, GenomicRanges, plyr, dplyr, magrittr

**Suggests** knitr, rmarkdown

**biocViews** CopyNumberVariation, Software, Sequencing, Coverage

**VignetteBuilder** knitr

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## Contents

|                               |    |
|-------------------------------|----|
| all.somatic.lst . . . . .     | 2  |
| allCall.lst . . . . .         | 3  |
| allCNV.lst . . . . .          | 3  |
| allHetero.lst . . . . .       | 4  |
| allRBD.lst . . . . .          | 4  |
| annoByGenesAndCyto . . . . .  | 4  |
| Annotate . . . . .            | 5  |
| bafTrack . . . . .            | 6  |
| btcompare . . . . .           | 7  |
| btpredict . . . . .           | 7  |
| BTreePlotter . . . . .        | 8  |
| BTreePredictor . . . . .      | 9  |
| cancer.genes.minus2 . . . . . | 9  |
| centromere.dat . . . . .      | 9  |
| cnv.gr . . . . .              | 10 |
| cyto.gr . . . . .             | 10 |
| drawBTree . . . . .           | 10 |
| drawBubbles . . . . .         | 11 |
| drawFeatures . . . . .        | 12 |
| gene.uni.clean.gr . . . . .   | 13 |
| getTracks . . . . .           | 14 |
| heteroLociTrack . . . . .     | 15 |
| hg19.seqinfo . . . . .        | 16 |
| info . . . . .                | 16 |
| loadRBD . . . . .             | 17 |
| makeRBD . . . . .             | 18 |
| mergeSnpCnv . . . . .         | 20 |
| RBD . . . . .                 | 21 |
| RscoreTrack . . . . .         | 21 |
| saveXLS . . . . .             | 22 |
| snp.gr . . . . .              | 23 |
| trackBTree . . . . .          | 23 |
| TrackPlotter . . . . .        | 24 |
| vol.genes . . . . .           | 25 |
| xyTrack . . . . .             | 25 |

|              |           |
|--------------|-----------|
| <b>Index</b> | <b>27</b> |
|--------------|-----------|

---

|                 |                        |
|-----------------|------------------------|
| all.somatic.lst | <i>all.somatic.lst</i> |
|-----------------|------------------------|

---

## 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 | <i>allHetero.lst</i> |
|---------------|----------------------|

---

**Description**

S4 GRanges dataset containing pre-calculated heterozygosity data.

**Format**

S4

**Source**

internal

---

|            |                   |
|------------|-------------------|
| allRBD.lst | <i>allRBD.lst</i> |
|------------|-------------------|

---

**Description**

A dataset containing precalculated data from CNV segment analysis.

**Format**

S4 object with rbd, rbd.adj

**Source**

internal

---

|                    |                           |
|--------------------|---------------------------|
| annoByGenesAndCyto | <i>annoByGenesAndCyto</i> |
|--------------------|---------------------------|

---

**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.1st.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.1st[[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

*bafTrack*


---

**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**

|            |                              |
|------------|------------------------------|
| .Object    | the object                   |
| result.dat | the result dataframe         |
| gr2        | the gr2 object               |
| somatic.gr | somatic gr object annotation |
| min.prev   | previous min                 |
| cex        | 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 | <i>btcompare</i> |
|-----------|------------------|

---

**Description**

btcompare

**Usage**

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

**Arguments**

|      |                       |
|------|-----------------------|
| set1 | first set             |
| set2 | 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 | <i>btpredict</i> |
|-----------|------------------|

---

**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

*BTreePlotter*


---

**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

---

|                     |               |
|---------------------|---------------|
| <code>cnv.gr</code> | <i>cnv.gr</i> |
|---------------------|---------------|

---

**Description**

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

**Format**

S4

**Source**

internal

---

|                      |                |
|----------------------|----------------|
| <code>cyto.gr</code> | <i>cyto.gr</i> |
|----------------------|----------------|

---

**Description**

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

**Format**

S4

**Source**

internal

---

|                        |                  |
|------------------------|------------------|
| <code>drawBTree</code> | <i>drawBTree</i> |
|------------------------|------------------|

---

**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.1st.RData", package="BubbleTree"))

btreeplotter <- new("BTreePlotter", max.ploidy=5, max.size=10)
nn <- "sam2"
rbd1 <- allCall.1st[[nn]]@rbd
rbd2 <- allCall.1st[[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.1st.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

*gene.uni.clean.gr*


---

## Description

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

## Format

S4

## Source

internal

---

getTracks

*getTracks*


---

## 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 | <i>heteroLociTrack</i> |
|-----------------|------------------------|

---

## 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"))

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

nn <- "sam6"

rbd <- allRBD.lst[[nn]]
btreepredictor@config$high.ploidy <- high.ploidy[nn]
btreepredictor@config$high.purity <- high.purity[nn]
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")

```

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"))

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

nn <- "sam6"

rbd <- allRBD.lst[[nn]]
btreepredictor@config$high.ploidy <- high.ploidy[nn]
btreepredictor@config$high.purity <- high.purity[nn]
btreepredictor <- loadRBD(btreepredictor, 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
btrepplotter <- new("BTreePlotter", max.ploidy=5, max.size=10)
btree <- drawBTree(btrepplotter, pred@rbd)
print(btree)

# create rbd.adj plot
btrepplotter <- new("BTreePlotter", branch.col="gray50")
btree <- drawBTree(btrepplotter, 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
btrepplotter <- new("BTreePlotter", max.ploidy=5, max.size=10)
arrows <- trackBTree(btrepplotter,
                     pred@rbd,
                     pred@rbd.adj,
                     min.srcSize=0.01,
                     min.trtSize=0.01)

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

# create a plot with overlays of significant genes
```

```

btreeplotter <- 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.1st.RData", package="BubbleTree"))
load(system.file("data", "centromere.dat.rda", package="BubbleTree"))
load(system.file("data", "allCNV.1st.RData", package="BubbleTree"))
load(system.file("data", "hg19.seqinfo.rda", package="BubbleTree"))

gr2 = centromere.dat
trackplotter <- new("TrackPlotter")
nn <- "sam2"
z <- RscoreTrack(trackplotter, allCall.1st[[nn]]@result, gr2, allCNV.1st[[nn]])
```

---

saveXLS

*saveXLS*


---

**Description**

saveXLS

**Usage**

```
saveXLS(dat.1st, xls.fn, row.names = FALSE, ...)
```

**Arguments**

|           |           |
|-----------|-----------|
| dat.1st   | dataframe |
| xls.fn    | filename  |
| row.names | row names |
| ...       | misc      |

**Value**

new Excel file

**Examples**

```
load(system.file("data", "allCall.1st.RData", package="BubbleTree"))

all.summary <- plyr::ldply(allCall.1st, 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 | <i>snp.gr</i> |
|--------|---------------|

---

**Description**

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

**Format**

S4

**Source**

internal

---

|            |                   |
|------------|-------------------|
| trackBTree | <i>trackBTree</i> |
|------------|-------------------|

---

**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

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

---

**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)
```

# Index

## \* datasets

- [all.somatic.lst](#), [2](#)
  - [allCall.lst](#), [3](#)
  - [allCNV.lst](#), [3](#)
  - [allHetero.lst](#), [4](#)
  - [allRBD.lst](#), [4](#)
  - [cancer.genes.minus2](#), [9](#)
  - [centromere.dat](#), [9](#)
  - [cnv.gr](#), [10](#)
  - [cyto.gr](#), [10](#)
  - [gene.uni.clean.gr](#), [13](#)
  - [hg19.seqinfo](#), [16](#)
  - [snp.gr](#), [23](#)
  - [vol.genes](#), [25](#)
- [all.somatic.lst](#), [2](#)
- [allCall.lst](#), [3](#)
- [allCNV.lst](#), [3](#)
- [allHetero.lst](#), [4](#)
- [allRBD.lst](#), [4](#)
- [annoByGenesAndCyto](#), [4](#)
- [annoByGenesAndCyto](#), [Annotate-method \(annoByGenesAndCyto\)](#), [4](#)
- [Annotate](#), [5](#)
- [Annotate-package \(Annotate\)](#), [5](#)
- [bafTrack](#), [6](#)
- [bafTrack](#), [TrackPlotter-method \(bafTrack\)](#), [6](#)
- [btcompare](#), [7](#)
- [btpredict](#), [7](#)
- [btpredict](#), [BTreePredictor-method \(btpredict\)](#), [7](#)
- [BTreePlotter](#), [8](#)
- [BTreePlotter-package \(BTreePlotter\)](#), [8](#)
- [BTreePredictor](#), [9](#)
- [BTreePredictor-package \(BTreePredictor\)](#), [9](#)
- [cancer.genes.minus2](#), [9](#)
- [centromere.dat](#), [9](#)
- [cnv.gr](#), [10](#)
- [cyto.gr](#), [10](#)
- [drawBTree](#), [10](#)
- [drawBTree](#), [BTreePlotter-method \(drawBTree\)](#), [10](#)
- [drawBubbles](#), [11](#)
- [drawBubbles](#), [BTreePlotter-method \(drawBubbles\)](#), [11](#)
- [drawFeatures](#), [12](#)
- [drawFeatures](#), [BTreePlotter-method \(drawFeatures\)](#), [12](#)
- [gene.uni.clean.gr](#), [13](#)
- [getTracks](#), [14](#)
- [heteroLociTrack](#), [15](#)
- [heteroLociTrack](#), [TrackPlotter-method \(heteroLociTrack\)](#), [15](#)
- [hg19.seqinfo](#), [16](#)
- [info](#), [16](#)
- [info](#), [BTreePredictor-method \(info\)](#), [16](#)
- [loadRBD](#), [17](#)
- [loadRBD](#), [BTreePredictor-method \(loadRBD\)](#), [17](#)
- [makeRBD](#), [18](#)
- [makeRBD](#), [RBD-method \(makeRBD\)](#), [18](#)
- [mergeSnpCnv](#), [20](#)
- [mergeSnpCnv](#), [RBD-method \(mergeSnpCnv\)](#), [20](#)
- [RBD](#), [21](#)
- [RBD-package \(RBD\)](#), [21](#)
- [RscoreTrack](#), [21](#)
- [RscoreTrack](#), [TrackPlotter-method \(RscoreTrack\)](#), [21](#)
- [saveXLS](#), [22](#)

snp.gr, [23](#)

trackBTree, [23](#)

trackBTree, BTreePlotter-method  
(trackBTree), [23](#)

TrackPlotter, [24](#)

TrackPlotter-package (TrackPlotter), [24](#)

vol.genes, [25](#)

xyTrack, [25](#)

xyTrack, TrackPlotter-method (xyTrack),  
[25](#)