Introduction to the plotAlongChrom function

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1 Introduction to the example data

The purpose of this vignette is to demonstrate some of the functionalities of the plotAlongChrom in the tilingArray package. We use a small subset data from an expression profiling paper [1]; The data only include the region from 35000bp to 50000bp in yeast chromosome one. Expression profiling is done in YPE and YPD conditions, 3 replicates each. Further information about the experimental design can be found at the paper website http://steinmetzlab.embl.de/NFRsharing/.

```r
> library("grid")
> library("RColorBrewer")
> library("tilingArray")

> data("segnf")
> class(segnf)

[1] "environment"

> ls(segnf)

[1] "1.-" "1.+"

> segnf$"1.+"

Object of class 'segmentation':
Data matrix: 1775 x 6
Change point estimates for number of segments S = 1:17
Confidence intervals for 1 fits from S = 17 to 17
Selected S = 17

> head(segnf$"1.+"@y)

YPE1  YPE2  YPE3  YPD1  YPD2  YPD3
[1,] -1.55 -2.70 -2.52 -4.78 -6.66 -4.62
[2,] -1.95 -4.20 -3.55 -3.93 -4.75 -4.74
[3,] -2.24 -2.24 -1.86 -6.15 -4.74 -3.82
[4,] -2.59 -2.52 -2.39 -4.31 -4.44 -4.40
[5,] -2.62 -3.91 -4.41 -4.94 -5.19 -4.65
[6,] -4.30 -4.82 -4.61 -5.62 -5.14 -5.60

> dim(segnf$"1.+"@y)

[1] 1775 6
```
The `segnf` object is an environment which contains two objects of class `segmentation`. `segnf` is the output of the `segChrom` in the `tilingArray` package. The `segmentation` object in `segnf` stores the probe expression information in the slot `y`. As can be seen, it contains 1775 probes and 6 array hybes in two conditions. The genomic coordinates where the probes aligned to is stored in the slot `x`. The order of the slot `x` is the same as the probe row order in slot `y`. The segment boundary information is stored in the slot `breakpoints` which is a list that contains all the optimal placement of 1 segment to the designate number (here in this data set is 17) of segments for this data. A log likelihood score for each placement is stored in slot `logLik` from which the best one is choosen and stored in the slot `nrSegments`. Further information about how the segmentation algorithm works, please read the vignette segmentation demo.

The `gffSub` object is a data frame that contains the SGD annotated features of the region 35000bp-50000bp for yeast chromosome one.
2 Visualizing the expression profiling with the plotAlongChrom function

The function `plotAlongChrom` accepts an environment as its first argument, which is expected to contain objects of class `segmentation` with names given by `paste(chr, c("+", "-"), sep=".")`, where `chr` is the chromosome identifier.

The following code generates Figure 1, a dot plot that averaged across all hybes.

```r
> grid.newpage()
> plotAlongChrom(segnf, chr=1, coord=c(35000,50000), what="dots", gff=gffSub)
```

![Figure 1: Along-chromosome dot plot of the averaged value across all hybes.](image)

We could also make separate dot plot for different hybes by setting the parameter `sepPlot` as TRUE. The following code generates Figure 2 that plots the expression separately for the two conditions.

```r
> segObj = new.env(parent = baseenv())
> nmLabel = colnames(segnf"1.+.y)
> lab = gsub("\\d","",nmLabel)
> for(nm in paste(1,c("+","-"),sep=".")){
+   s = get(nm,env = segnf)
+   rpY = tapply(1:length(lab),lab,function(i)rowMeans(s@y[,i]))
+   s@y = do.call(cbind,rpY)
+   assign(nm,s,segObj)
+ }
> grid.newpage()
> plotAlongChrom(segObj, chr=1, coord=c(35000,50000), what="dots", gff=gffSub, sepPlot = T)
```

However, with the number of hybes increases, it is very hard to see the difference in dot plots in a normal screen. Thus, if the number of hybes is more than 4, the function will force to take the average. A better alternative of displaying multiple hybes is to use the heatmap. The following code generates Figure 3 that makes the heatmap plot.

```r
> grid.newpage()
> plotAlongChrom(segnf, chr=1, coord=c(35000,50000), what="heatmap", gff=gffSub, sepPlot = T)
```

![Figure 3: Heatmap plot of the expression across all hybes.](image)
Figure 2: Along-chromosome dot plot of the averaged value among different replicates for YPD and YPE condition.

Figure 3: Along-chromosome heatmap plot of all the replicates in YPD and YPE condition.
Start with R 2.11.0, the \texttt{grid} package introduced the raster array image function \texttt{grid.raster} which is a faster and efficient way of generating heatmap images. From R 2.11.0, The \texttt{herringArray} package will use the \texttt{grid.raster} function as default to make heatmap images replacing the previous \texttt{grid.rect} function. The choice between the two drawing functions can be changed by the parameter \texttt{makeRasterImage}. The following code generates Figure 4 that makes the raster heatmap plot.

```r
> grid.newpage()
> plotAlongChrom(segnf,chr=1, coord=c(35000,50000), what="heatmap", gff=gffSub, +   rowNamesHeatmap=nmLabel,makeRasterImage=TRUE)
```

![Figure 4: Along-chromosome raster heatmap plot of all the replicates in YPD and YPE condition.](image)

The color gradient of the heatmap could be changed by the parameter \texttt{colHeatmap}. The following code generates Figure 5 that makes the raster heatmap plot using a blue color gradient.

```r
> grid.newpage()
> plotAlongChrom(segnf,chr=1, coord=c(35000,50000), what="heatmap", gff=gffSub, +   rowNamesHeatmap=nmLabel,makeRasterImage=TRUE, +   colHeatmap = colorRamp(brewer.pal(9, "Blues")))
```

References

Figure 5: Along-chromosome raster heatmap plot of all the replicates in YPD and YPE condition with a blue color gradient.