Extending oligo with SNPchip

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Introduction

This vignette describes a pipeline for preprocessing and visualizing SNP-level summaries using the packages oligo and SNPchip. We use a set of unprocessed Affymetrix files (CEL files) available as experimental data packages on Bioconductor. A minimal set of commands to perform pre-processing with oligo are provided here, though one should consult the oligo vignette for additional information. An object of the processed data is provided with this package to reduce the time of computation – the code chunks for the preprocessing steps are not evaluated in the vignette. An example of using oligo to process a batch of 209 Affymetrix 100k CEL files and VanillaICE to identify regions of alterations are provided in the hapmap100k vignette in the directory inst/testing of the VanillaICE package. The hapmap100k vignette is not reproducible as it depends on access to the 209 CEL hapmap CEL files that are not provided with the VanillaICE package, but may be useful as a guideline when performing your own analyses. Comparable vignettes for hapmapAffy500k, hapmapAffy5.0, hapmapAffy6.0, and Illumina will also be added to VanillaICE in the near future.

1 Creating an instance of oligoSnpSet

The *oligo* vignette creates an instance of SnpCallSetPlus, crlmmOut, from the call to the function crlmm. For purposes of illustration, I subset the object to only include SNPs on chromosome 1. I also took the liberty of adding chromosome and physical position to the featureData slot. This object can be loaded by

```
> library(SNPchip)
> data(crlmmOut)
> class(crlmmOut)

[1] "SnpCallSetPlus"
attr(,"package")
[1] "oligoClasses"
```

The elements in the assayData for instances of SnpCallSetPlus is dependent on the Affymetrix platform.

```
> annotation(crlmmOut)
[1] "pd.mapping50k.xba240"
> ls(assayData(crlmmOut))
[1] "antisenseThetaA" "antisenseThetaB" "calls"
[4] "callsConfidence" "senseThetaA" "senseThetaB"
> callset <- crlmmOut</pre>
```

1.1 Estimating copy number

We expect that the intensities for the A and B alleles averaged over the sense and antisense strands will be proportional to the total copy number. Because the fluorescence at a SNP is very much SNP-dependent, we'll center each SNP at the median value and then recenter at the 'normal' copy number. Here, we define normal copy number to be two for autosomes, 1 for the male X, and 2 for the female X. Because we've noticed that homozygous genotype calls appear to have overall less fluorescence than the heterozygous genotype calls, we'll recenter the median intensities for the homozygous and heterozygous genotypes to be equivalent.

We begin by extracting an array of average values for the sense (A and B alleles) and antisense (A and B) strands.

[1] 4669

The homozygous and heterozygous genotypes are centered separately since the average intensities differ:

```
> chr.matrix <- matrix(chromosome(callset), ncol = ncol(log2cn),</pre>
      nrow = nrow(log2cn))
> median.hom <- median(log2cn[(calls(callset) ==</pre>
      1 | calls(callset) == 3) & chr.matrix != "X"],
      na.rm = TRUE)
+
> median.het <- median(log2cn[calls(callset) ==</pre>
      2 & chr.matrix != "X"], na.rm = TRUE)
 recenterByGenotype <- function(x, callset, recenter.hom,
      recenter.het) {
      calls <- as.vector(calls(callset))</pre>
+
      x[calls == 1 \mid calls == 3] \leftarrow x[calls == 1 \mid
           calls == 3] - recenter.hom
      x[calls == 2] \leftarrow x[calls == 2] - recenter.het
+
+
+ }
> for (j in 1:ncol(log2cn)) {
+
      log2cn[, j] <- recenterByGenotype(log2cn[,</pre>
+
           j], callset[, j], recenter.hom = median.hom,
+
           recenter.het = median.het)
```

Next, we sweep out a robust estimate of the median from the samples (tries to put fluorescence intensities on a similar scale for each of the samples)

```
tmp2 <- split(x, chromosome)</pre>
      if (length(tmp2) > 15) {
+
+
           idx <- order(sapply(tmp2, "median"))</pre>
           tmp2 \leftarrow tmp2[idx]
+
           tmp3 \leftarrow tmp2[-c(1:5, (length(tmp2) - 4):length(tmp2))]
+
           med <- median(unlist(tmp3))</pre>
+
      }
+
      else {
           med <- median(sapply(tmp2, "median"))</pre>
+
      return (med)
+ }
> robust.median <- apply(log2cn, 2, f, chromosome(callset))</pre>
> log2cn <- sweep(log2cn, 2, robust.median)
We then sweep out the SNP-specific median intensities and recenter to the expected copy number
(depends on chromosome):
> rowSweep <- function(callset, X, value, recenter,
      j) {
      if (length(value) == 1) {
           i <- chromosome(callset) == value</pre>
+
      }
+
      else {
           i <- chromosome(callset) %in% value
+
+
      i[is.na(i)] <- FALSE
+
      if (sum(i) > 1) {
+
           if (!missing(j)) {
               if (sum(j) < 5)
                    warning("very few samples for calculating a robust average")
               avg <- rowMedians(X[i, j], na.rm = TRUE)</pre>
+
               X[i, j] \leftarrow sweep(X[i, j], 1, avg) +
                    recenter
           }
           else {
               avg <- rowMedians(X[i, ], na.rm = TRUE)</pre>
+
               X[i, ] \leftarrow sweep(X[i, ], 1, avg) +
+
                   recenter
           }
+
+
      }
+
      Χ
+ }
> male <- callset$gender == 1
> female <- callset$gender == 2</pre>
> chromosome(callset)[is.na(chromosome(callset))] <- "NA"</pre>
> log2cn <- rowSweep(callset, log2cn, "NA", log2(2))
> log2cn <- rowSweep(callset, log2cn, "M", log2(2))</pre>
> log2cn <- rowSweep(callset, log2cn, "X", log2(1),
> log2cn <- rowSweep(callset, log2cn, "X", log2(2),</pre>
```

> f <- function(x, chromosome) {</pre>

```
+ female)
> log2cn <- rowSweep(callset, log2cn, "Y", log2(1),
+ male)
> log2cn <- rowSweep(callset, log2cn, "Y", log2(0.5),
+ female)
> log2cn <- rowSweep(callset, log2cn, as.character(1:22),
+ log2(2))
> chromosome(callset)[chromosome(callset) == "NA"] <- NA
> copyNumber <- 2^log2cn</pre>
```

The proceding codechunks may be replaced by the function calculateCopyNumber.

```
> cn <- calculateCopyNumber(callset, center.autosomes = 2,
+ center.X.male = 1, center.X.female = 2, center.Y.male = 1,
+ center.Y.female = 0.4)
> identical(cn, copyNumber)
```

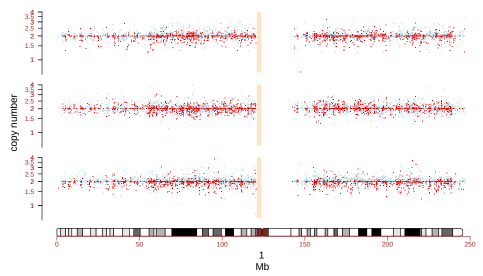
[1] TRUE

We may now create an object of class oligoSnpSet that contains assayData elements for SNP-level summaries of genotype calls and copy number. For now, we will assign a matrix of missing values for copy number confidence estimates.

Estimating copy number from a SnpCallSetPlus object and creating an instance of the class oligoS-npSet are encapsulated in the method for coercing an object between the two classes:

```
> snpset <- as(callset, "oligoSnpSet")
A plot of chromosome 1:
> gp <- new("ParSnpSet")
> gp <- getPar(gp, snpset)

[1] "one.ylim is FALSE. Calculating ylim based on the percentiles of the copy number distribution"
> gp$ylab <- "copy number"
> gp$ylim <- c(0.7, 4)
> plotSnp(gp, snpset)
NULL
```



A hidden Markov model can be used to identify chromosomal alterations using genotype and copy number estimates as described in the *VanillaICE* vignette.

2 Combining objects that use different annotation packages

Here we illustrate how one may combine two objects of class SnpCallSetPlus that use different annotation packages: e.g., pd.mapping50k.hind240 and pd.mapping50k.xba240. Following the oligo vignette, I created hind and xba instances of SnpCallSetPlus. The following code is not evaluated due to time constraints.

```
> library("oligo")
> library("hapmap100kxba")
> pathCelFiles <- system.file("celFiles", package = "hapmap100kxba")</pre>
 fullFilenames <- list.celfiles(path = pathCelFiles,
      full.names = TRUE)
  aboutSamples <- data.frame(gender = c("female",
      "female", "male"))
 rownames(aboutSamples) <- basename(fullFilenames)</pre>
 aboutVars <- data.frame(labelDescription = "male/female")</pre>
 rownames(aboutVars) <- "gender"
  pd <- new("AnnotatedDataFrame", data = aboutSamples,</pre>
      varMetadata = aboutVars)
 xba <- justCRLMM(fullFilenames, phenoData = pd,</pre>
      verbose = FALSE)
> library("hapmap100khind")
 pathCelFiles <- system.file("celFiles", package = "hapmap100khind")</pre>
 fullFilenames <- list.celfiles(path = pathCelFiles,</pre>
      full.names = TRUE)
  aboutSamples <- data.frame(gender = c("female",</pre>
      "female", "male"))
 rownames(aboutSamples) <- basename(fullFilenames)</pre>
> aboutVars <- data.frame(labelDescription = "male/female")</pre>
```

```
> rownames(aboutVars) <- "gender"
> pd <- new("AnnotatedDataFrame", data = aboutSamples,
+ varMetadata = aboutVars)
> hind <- justCRLMM(fullFilenames, phenoData = pd,
+ verbose = FALSE)

To combine into one object, simply
> callset <- combine(xba, hind)</pre>
```

3 Session Information

- R version 2.7.0 Under development (unstable) (2008-01-28 r44219), powerpc-apple-darwin8.11.0
- Locale: C
- Base packages: base, datasets, grDevices, graphics, methods, splines, stats, tools, utils
- Other packages: AnnotationDbi 0.99.23, Biobase 1.99.0, DBI 0.2-3, RSQLite 0.6-4, SNPchip 1.3.16, affxparser 1.9.5, oligo 1.3.15, oligoClasses 1.1.15, pd.mapping50k.xba240 0.3.4, preprocessCore 0.99.22