1 Introduction

The goal of the tutorial is to present a standard analysis workflow of 450K data with the package minfi, incorporating the functions recently added to the package. We invite you to read the software paper recently published [1] and the online package vignette on the Bioconductor project [2] for more details.

We will start from the very beginning by reading input raw data (IDAT files) from an example dataset, and ending with a list of candidate genes for differential methylation. Among others, we will cover quality control assessments, different within-array and between-array normalizations, SNPs removal, sex prediction, differentially methylated positions (DMPs) analysis and bump hunting for differentially methylated regions (DMRs).

If time permits, we will introduce a complementary interactive visualization tool package, shinyMethyl that allows interactive quality control assessment. [3] You can download the package online from gitHub [4] and try it online at http://spark.rstudio.com/jfortin/shinyMethyl/

450k Array design and terminology

In this section, we introduce briefly the 450K array as well as the terminology used throughout the minfi package. Each sample is measured on a single array, in two different color channels (red and green). As the name of the platform indicates, each array measures more than 450,000 CpG positions. For each CpG, we have two measurements: a methylated intensity and an unmethylated intensity. Depending on the probe design, the signals are reported in different colors:

For Type I design, both signals are measured in the same color: one probe for the methylated signal and one probe for the unmethylated signal.
Figure 1: **Probe design of the 450k array.** For **Type I** design, both signals are measured in the same color: one probe for the methylated signal and one probe for the unmethylated signal. For **Type II** design, only one probe is used. The **Green** intensity measures the methylated signal, and the **Red** intensity measures the unmethylated signal.

For **Type II** design, only one probe is used. The **Green** intensity measures the methylated signal, and the **Red** intensity measures the unmethylated signal.

**Some definitions**

Two commonly measures are used to report the methylation levels: Beta values and M values.

**Beta value:**

\[ \beta = \frac{M}{M + U + 100} \]

where \( M \) and \( U \) denote the methylated and unmethylated signals respectively. In minfi the constant 100 can easily be changed, if needed.

**MValue:**

\[ Mval = \log \left( \frac{M}{U} \right) \]

**DMP:** Differentially methylated position: single genomic position that has a different methylated level in two different groups of samples (or conditions)

**DMR:** Differentially methylated region: when consecutive genomic locations are differentially methylated in the same direction.

**Array:** One sample

**Slide:** Physical slide containing 12 arrays (6 × 2 grid)

**Plate:** Physical plate containing at most 8 slides (96 arrays). For this tutorial, we use **batch** and plate interchangeably.
2  Reading Data

The starting point of minfi is reading the .IDAT files with the built-in function read.450k.exp. Several options are available: the user can specify the sample filenames to be read in along with the directory path, or can specify the directory that contains the files. In the latter case, all the files with the extension .IDAT located in the directory will be loaded into R. The user can also read in a sample sheet, and then use the sample sheet to load the data into a RGChannelSet. For more information, see the minfi vignette. Here, we will load the dataset containing 6 samples from the minfiData package using the sample sheet provided within the package:

```r
> require(minfi)
> require(minfiData)
> baseDir <- system.file("extdata",package="minfiData")
> targets <- read.450k.sheet(baseDir)
```

[read.450k.sheet] Found the following CSV files:

```r
> targets
```

<table>
<thead>
<tr>
<th>Sample_Name</th>
<th>Sample_Well</th>
<th>Sample_Plate</th>
<th>Sample_Group</th>
<th>Pool_ID</th>
<th>person</th>
<th>age</th>
<th>sex</th>
</tr>
</thead>
<tbody>
<tr>
<td>GroupA_3</td>
<td>H5</td>
<td>NA</td>
<td>GroupA</td>
<td>NA</td>
<td>id3</td>
<td>83</td>
<td>M</td>
</tr>
<tr>
<td>GroupA_2</td>
<td>D5</td>
<td>NA</td>
<td>GroupA</td>
<td>NA</td>
<td>id2</td>
<td>58</td>
<td>F</td>
</tr>
<tr>
<td>GroupB_3</td>
<td>C6</td>
<td>NA</td>
<td>GroupB</td>
<td>NA</td>
<td>id3</td>
<td>83</td>
<td>M</td>
</tr>
<tr>
<td>GroupB_1</td>
<td>F7</td>
<td>NA</td>
<td>GroupB</td>
<td>NA</td>
<td>id1</td>
<td>75</td>
<td>F</td>
</tr>
<tr>
<td>GroupA_1</td>
<td>G7</td>
<td>NA</td>
<td>GroupA</td>
<td>NA</td>
<td>id1</td>
<td>75</td>
<td>F</td>
</tr>
<tr>
<td>GroupB_2</td>
<td>H7</td>
<td>NA</td>
<td>GroupB</td>
<td>NA</td>
<td>id2</td>
<td>58</td>
<td>F</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>status</th>
<th>Array</th>
<th>Slide</th>
</tr>
</thead>
<tbody>
<tr>
<td>normal</td>
<td>R02C02</td>
<td>5723646052</td>
</tr>
<tr>
<td>normal</td>
<td>R04C01</td>
<td>5723646052</td>
</tr>
<tr>
<td>cancer</td>
<td>R05C02</td>
<td>5723646052</td>
</tr>
<tr>
<td>cancer</td>
<td>R04C02</td>
<td>5723646053</td>
</tr>
<tr>
<td>normal</td>
<td>R05C02</td>
<td>5723646053</td>
</tr>
<tr>
<td>cancer</td>
<td>R06C02</td>
<td>5723646053</td>
</tr>
</tbody>
</table>

```r
> RGSet <- read.450k.exp(base = baseDir, targets = targets)
```
The class of RGSet is a \texttt{RGChannelSet} object. This is the initial object of a \texttt{minfi} analysis that contains the raw intensities in the green and red channels. Note that this object contains the intensities of the internal control probes as well. Because we read the data from a data sheet experiment, the phenotype data is also stored in the \texttt{RGChannelSet} and can be accessed via the accessor command \texttt{pData}:

```
> phenoData <- pData(RGSet)
> phenoData[,1:6]
```

```
Sample_Name Sample_Well Sample_Plate Sample_Group Pool_ID
5723646052_R02C02 GroupA_3 H5 NA GroupA NA
5723646052_R04C01 GroupA_2 D5 NA GroupA NA
5723646052_R05C02 GroupB_3 C6 NA GroupB NA
5723646053_R04C02 GroupB_1 F7 NA GroupB NA
5723646053_R05C02 GroupA_1 G7 NA GroupA NA
5723646053_R06C02 GroupB_2 H7 NA GroupB NA
```

The \texttt{RGChannelSet} stores also a manifest object that contains the probe design information of the array. This object is mainly of interest for developers; the package makes use of this behind the scenes.

```
> manifest <- getManifest(RGSet)
> manifest
```

```
IlluminaMethylationManifest object
Annotation
  array: IlluminaHumanMethylation450k
Number of type I probes: 135476
Number of type II probes: 350036
Number of control probes: 850
Number of SNP type I probes: 25
Number of SNP type II probes: 40
```

```
> head(getProbeInfo(manifest))
```

```
DataFrame with 6 rows and 8 columns
  Name AddressA AddressB Color NextBase
<character> <character> <character> <character> <DNAStringSet>
1  cg00050873 32735311 31717405 Red A
```
### 3 Class structure

minfi contains a number of classes corresponding to various transformations of the raw data. It is important to understand how these classes relate to each other. Figure 2 provides a useful overview.

#### MethylSet and RatioSet

A **MethylSet** object contains the methylated and unmethylated signals. The most basic way to construct a **MethylSet** is to use the function `preprocessRaw` which uses the array design to match up the different probes and color channels to construct the methylated and unmethylated signals. This function does not do any normalization (see later).

```r
> MSet <- preprocessRaw(RGSet)
> MSet

MethylSet (storageMode: lockedEnvironment)
assayData: 485512 features, 6 samples
```

See the `minfi` vignette for more information.
Figure 2: Flow chart of the minfi’s functions
element names: Meth, Unmeth

phenoData
  sampleNames: 5723646052_R02C02 5723646052_R04C01 ...
         5723646053_R06C02 (6 total)
  varLabels: Sample_Name Sample_Well ... filenames (13 total)
  varMetadata: labelDescription

Annotation
  array: IlluminaHumanMethylation450k
  annotation: ilmn12.hg19

Preprocessing
  Method: Raw (no normalization or bg correction)
  minfi version: 1.10.2
  Manifest version: 0.4.0

The accessors getMeth and getUnmeth can be used to get the methylated and unmethylated intensities matrices:

> head(getMeth(MSet)[,1:3])

      5723646052_R02C02 5723646052_R04C01 5723646052_R05C02
cg00050873       22041       588     20505
cg00212031        679        569       439
cg00213748       1620       421       707
cg00214611        449       614       343
cg00455876       5921       421       707
cg01707559       1238       646       637

> head(getUnmeth(MSet)[,1:3])

      5723646052_R02C02 5723646052_R04C01 5723646052_R05C02
cg00050873        1945        433      1012
cg00212031        6567        300      2689
cg00213748         384        461       295
cg00214611        4869        183      1655
cg00455876        1655        792      1060
cg01707559       12227       1009      7414

A RatioSet object is a class designed to store Beta values and/or M values instead of the methylated and unmethylated signals. An optional copy number matrix, CN, the sum of the methylated and unmethylated signals, can be also stored. Mapping a MethylSet to a RatioSet is irreversible, i.e. one cannot technically retrieve the methylated and unmethylated signals from a RatioSet. A RatioSet can be created with the function ratioConvert:

> ratioSet <- ratioConvert(MSet, what = "both", keepCN = TRUE)
> ratioSet
The functions `getBeta`, `getM` and `getCN` return respectively the Beta value matrix, M value matrix and a the Copy Number matrix.

Why do we have these two classes? This is to allow methods development where normalization is done directly on the beta and/or M-values, such as quantile normalization of the Beta matrix (which we btw. do not recommend).

**Mapping to the Genome**

The function `mapToGenome` applied to a `RatioSet` object will add genomic coordinates to each probe together with some additional annotation information. The output object is a `GenomicRatioSet` (class holding M or/and Beta values together with associated genomic coordinates). It is possible to merge the manifest object with the genomic locations by setting the option `mergeManifest` to TRUE.

```r
> gset <- mapToGenome(ratioSet)
> gset
```

```r
class: GenomicRatioSet
dim: 485512 6
exptData(0):
assays(3): Beta M CN
rownames(485512): cg13869341 cg14008030 ... cg08265308 cg14273923
rowData metadata column names(0):
colnames(6): 5723646052_R02C02 5723646052_R04C01 ... 5723646053_R05C02
5723646053_R06C02
```
colData names(13): Sample_Name Sample_Well ... Basename filenames

Annotation
   array: IlluminaHumanMethylation450k
       annotation: ilmn12.hg19

Preprocessing
   Method: Raw (no normalization or bg correction)
       minfi version: 1.10.2
       Manifest version: 0.4.0

Note that the GenomicRatioSet extends the class SummarizedExperiment. Here are the main accessors functions to access the data:

> beta <- getBeta(gset)
> m <- getM(gset)
> cn <- getCN(gset)

> sampleNames <- sampleNames(gset)
> probeNames <- featureNames(gset)
> pheno <- pData(gset)

To return the probe locations as a GenomicRanges objects, one can use the accessor granges:

> gr <- granges(gset)
> head(gr, n= 3)

GRanges with 3 ranges and 0 metadata columns:
   seqnames ranges strand
       <Rle> <IRanges> <Rle>
   cg13869341    chr1 [15865, 15865] *
   cg14008030    chr1 [18827, 18827] *
   cg12045430    chr1 [29407, 29407] *

We can similary use mapToGenome on a MethylSet to get a GenomicMethylSet.

Annotation

To access the full annotation, one can use the command getAnnotation:

> annotation <- getAnnotation(gset)
> names(annotation)
The order and content of the annotation DataFrame is in the same order as the main object (here: gset). There are a number of convenience functions to get parts of the annotation, like

```r
> islands <- getIslandStatus(gset)
> head(islands)
[1] "OpenSea" "OpenSea" "Island" "Island" "Island" "OpenSea"
> probeType <- getProbeType(gset)
> head(probeType)
[1] "I"  "II"  "I"  "I"  "I"  "II"
```
(see later regarding SNPs). Also, you can get different subsets of the annotation by using the `what` argument.

## 4 Quality control

Efficient and reliable quality control is important. Our view on this has evolved over time and currently we recommend using the qc plot described below as well as `shinyMethyl` for interactive visualization. We conclude this section with some comments on what we don’t recommend using.
**QC plot**

`minfi` provides a simple quality control plot that uses the log median intensity in both the methylated (M) and unmethylated (U) channels. When plotting these two medians against each other, it has been observed that good samples cluster together, while failed samples tend to separate and have lower median intensities [1]. In general, we advice users to make the plot and make a judgement. The line separating "bad" from "good" samples represent a useful cutoff, which may have to be adapted to a specific dataset. The functions `getQC` and `plotQC` are designed to extract and plot the quality control information from the `MethylSet`:

```r
> qc <- getQC(MSet)
> head(qc)

Dataframe with 6 rows and 2 columns

<table>
<thead>
<tr>
<th>mMed</th>
<th>uMed</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;numeric&gt;</td>
<td>&lt;numeric&gt;</td>
</tr>
<tr>
<td>5723646052_R02C02</td>
<td>11.69566</td>
</tr>
<tr>
<td>5723646052_R04C01</td>
<td>11.99046</td>
</tr>
<tr>
<td>5723646052_R05C02</td>
<td>11.55603</td>
</tr>
<tr>
<td>5723646053_R04C02</td>
<td>12.06609</td>
</tr>
<tr>
<td>5723646053_R05C02</td>
<td>12.23332</td>
</tr>
<tr>
<td>5723646053_R06C02</td>
<td>11.36851</td>
</tr>
</tbody>
</table>
```

```r
> plotQC(qc)
```

11
Moreover, the function `addQC` applied to the `MethylSet` will add the QC information to the phenotype data.

To further explore the quality of the samples, it is useful to look at the Beta value densities of the samples, with the option to color the densities by group:

```r
> densityPlot(MSet, sampGroups = phenoData$Sample_Group)
```
or density bean plots:

```r
> densityBeanPlot(MSet, sampGroups = phenoData$Sample_Group)
```
shinyMethyl [3] is particularly useful to visualize all plots at the same time in an interactive fashion.

**Control probes plot**

The 450k array contains several internal control probes that can be used to assess the quality control of different sample preparation steps (bisulfite conversion, hybridization, etc.). The values of these control probes are stored in the initial **RGChannelSet** and can be plotted by using the function **controlStripPlot** and by specifying the control probe type:

```r
> controlStripPlot(RGSet, controls="BISULFITE CONVERSION II")
```

All the plots above can be exported into a pdf file in one step using the function **qcReport**:

```r
> qcReport(RGSet, pdf= "qcReport.pdf")
```
In practice, we use the QC plot presented above as well as inspection of the bisulfite conversion probes and marginal densities to do QC. We don’t really use the \texttt{qcReport} anymore.

5 SNPs

Because the presence of SNPs inside the probe body or at the nucleotide extension can have important consequences on the downstream analysis, \texttt{minfi} offers the possibility to remove such probes. The function \texttt{getSnpInfo}, applied to a \texttt{GenomicRatioSet}, returns a data frame with 6 columns containing the SNP information of the probes:

\begin{verbatim}
> snps <- getSnpInfo(gset)
> head(snps,10)

DataFrame with 10 rows and 6 columns

<table>
<thead>
<tr>
<th>Probe_rs</th>
<th>Probe_maf</th>
<th>CpG_rs</th>
<th>CpG_maf</th>
<th>SBE_rs</th>
<th>SBE_maf</th>
</tr>
</thead>
<tbody>
<tr>
<td>cg13869341</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>cg14008030</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>cg12045430</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>cg20826792</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>cg00381604</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>cg20253340</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>cg21870274</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>cg03130891</td>
<td>rs77418980</td>
<td>0.305556</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>cg24335620</td>
<td>rs147502335</td>
<td>0.012800</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>cg16162899</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

Probe, CpG and SBE correspond the SNPs present inside the probe body, at the CpG interrogation and at the single nucleotide extension respectively. The columns with \texttt{rs} give the names of the SNPs while the columns with \texttt{maf} gives the minor allele frequency of the SNPs based on the dbSnp database. The function \texttt{addSnpInfo} will add to the \texttt{GenomicRanges} of the \texttt{GenomicRatioSet} the 6 columns:

\begin{verbatim}
> gset <- addSnpInfo(gset)
> head(granges(gset))

GRanges with 6 ranges and 6 metadata columns:

<table>
<thead>
<tr>
<th>seqnames</th>
<th>ranges</th>
<th>strand</th>
<th>Probe_rs</th>
<th>Probe_maf</th>
<th>CpG_rs</th>
</tr>
</thead>
<tbody>
<tr>
<td>chr1 [15865, 15865]</td>
<td>*</td>
<td>&lt;NA&gt;</td>
<td>&lt;NA&gt;</td>
<td>&lt;NA&gt;</td>
<td></td>
</tr>
<tr>
<td>chr1 [18827, 18827]</td>
<td>*</td>
<td>&lt;NA&gt;</td>
<td>&lt;NA&gt;</td>
<td>&lt;NA&gt;</td>
<td></td>
</tr>
<tr>
<td>chr1 [29407, 29407]</td>
<td>*</td>
<td>&lt;NA&gt;</td>
<td>&lt;NA&gt;</td>
<td>&lt;NA&gt;</td>
<td></td>
</tr>
<tr>
<td>chr1 [29425, 29425]</td>
<td>*</td>
<td>&lt;NA&gt;</td>
<td>&lt;NA&gt;</td>
<td>&lt;NA&gt;</td>
<td></td>
</tr>
<tr>
<td>chr1 [29425, 29425]</td>
<td>*</td>
<td>&lt;NA&gt;</td>
<td>&lt;NA&gt;</td>
<td>&lt;NA&gt;</td>
<td></td>
</tr>
</tbody>
</table>
We strongly recommend to drop the probes that contain either a SNP at the CpG interrogation or at the single nucleotide extension. The function `dropLociWithSnps` allows to drop the corresponding probes (introduced in minfi 1.11.9). Here is an example where we drop the probes containing a SNP at the CpG interrogation and/or at the single nucleotide extension, for any minor allele frequency:

```r
> gset <- dropLociWithSnps(gset, snps=c("SBE","CpG"), maf=0)
```

There are several options for SNP databases. These are contained inside the annotation object. A list of databases can be had by printing the annotation object. In mifni 1.11.8 we made it possible to do the following:

```r
> getAnnotationObject(gset)
```

In earlier versions of minfi, you can get the same outcome by first getting the name of the annotation object and then printing it, by

```r
> annoStr <- paste(annotation(MsetEx), collapse = "anno.")
> annoStr

[1] "IlluminaHumanMethylation450kanno.ilmn12.hg19"
> get(annoStr)
```

IlluminaMethylationAnnotation object
Annotation
  array: IlluminaHumanMethylation450k
  annotation: ilmn12
  genomeBuild: hg19
Available annotation
  Islands.UCSC
Locations
Cross-reactive probes

It has been previously reported than about 6% of the probes on the 450K array co-hybridize to alternate genomic sequences, therefore potentially generating spurious signals [5]. We are planning to include a function in minfi that drops these cross-reactive probes. The function can be either applied to a [Genomic]MethylSet or a [Genomic]RatioSet:

```r
> gset <- dropCrossReactiveProbes(gset)
```

This functionality is currently being tested.

6 Preprocessing and normalization

So far, we did not use any normalization to process the data. Different normalization procedures are available in minfi.

preprocessRaw

As seen before, it converts a RGChannelSet to a MethylSet by converting the Red and Green channels into a matrix of methylated signals and a matrix of unmethylated signals. No normalization is performed.

Input: RGChannelSet
Output: MethylSet
preprocessIllumina

Convert a RGChannelSet to a MethylSet by implementing the preprocessing choices as available in Genome Studio: background subtraction and control normalization. Both of them are optional and turning them off is equivalent to raw preprocessing (preprocessRaw):

```r
> MSet.illumina <- preprocessIllumina(RGSet, bg.correct = TRUE, +     normalize = "controls")
```

Input: RGChannelSet
Output: MethylSet

preprocessSWAN

Perform Subset-quantile within array normalization (SWAN) [6], a within-array normalization correction for the technical differences between the Type I and Type II array designs. The algorithm matches the Beta-value distributions of the Type I and Type II probes by applying a within-array quantile normalization separately for different subsets of probes (divided by CpG content). The input of SWAN is a MethylSet, and the function returns a MethylSet as well. If an RGChannelSet is provided instead, the function will first call preprocessRaw on the RGChannelSet, and then apply the SWAN normalization. We recommend setting a seed (using set.seed) before using preprocessSWAN to ensure that the normalized intensities will be reproducible.

```r
> MSet.swan <- preprocessSWAN(RGSet)
```

Input: RGChannelSet or MethylSet
Output: MethylSet

preprocessQuantile

This function implements stratified quantile normalization preprocessing. The normalization procedure is applied to the Meth and Unmeth intensities separately. The distribution of type I and type II signals is forced to be the same by first quantile normalizing the type II probes across samples and then interpolating a reference distribution to which we normalize the type I probes. Since probe types and probe regions are confounded and we know that DNAm distributions vary across regions we stratify the probes by region before applying this interpolation. Note that this algorithm relies on the assumptions necessary for quantile normalization to be applicable and thus is not recommended for cases where global changes are expected such as in cancer-normal comparisons. Note that this normalization procedure is essentially similar to one previously presented [7]. The different options can be summarized into the following list:
1) If `fixMethOutlier` is `TRUE`, the functions fixes outliers of both the methylated and unmethylated channels when small intensities are close to zero.

2) If `removeBadSamples` is `TRUE`, it removes bad samples using the QC criterion discussed previously.

3) Performs stratified subset quantile normalization if `quantileNormalize=TRUE` and `stratified=TRUE`.

4) Predicts the sex (if not provided in the `sex` argument) using the function `getSex` and normalizes males and females separately for the probes on the X and Y chromosomes.

```r
> gset.quantile <- preprocessQuantile(RGSet, fixOutliers = TRUE, removeBadSamples = TRUE, badSampleCutoff = 10.5, quantileNormalize = TRUE, stratified = TRUE, mergeManifest = FALSE, sex = NULL)
```

Input: RGChannelSet
Output: GenomicRatioSet

Note that the function returns a GenomicRatioSet object ready for downstream analysis.

**preprocessFunnorm**

The function `preprocessFunnorm` implements the functional normalization algorithm developed in [8]. Briefly, it uses the internal control probes present on the array to infer between-array technical variation. It is particularly useful for studies comparing conditions with known large-scale differences, such as cancer/normal studies, or between-tissue studies. It has been shown that for such studies, functional normalization outperforms other existing approaches [8]. By default, it uses the first two principal components of the control probes to infer the unwanted variation.

```r
> gset.funnorm <- preprocessFunnorm(RGSet)
```

Input: RGChannelSet
Output: GenomicRatioSet

As the `preprocessQuantile` function, it returns a GenomicRatioSet object.
7 dmpFinder: to find differentially methylated positions (DMPs)

While we do not encourage particularly a single position differential methylation analysis, \texttt{minfi} implements a simple algorithm called \texttt{dmpFinder} to find differentially methylated positions with respect to a phenotype covariate. The phenotype may be categorical (e.g. cancer vs. normal) or continuous (e.g. blood pressure). Below is an example of a DMP analysis for age using the \texttt{gset.funnorm} object created above:

```r
> beta <- getBeta(gset.funnorm)
> age <- pData(gset.funnorm)$age
> dmp <- dmpFinder(beta, pheno = age, type = "continuous")
> head(dmp)
```

<table>
<thead>
<tr>
<th></th>
<th>intercept</th>
<th>beta</th>
<th>t</th>
<th>pval</th>
<th>qval</th>
</tr>
</thead>
<tbody>
<tr>
<td>cg05764847</td>
<td>0.9984524</td>
<td>-0.0007670769</td>
<td>-51.86582</td>
<td>8.270863e-07</td>
<td>0.4015603</td>
</tr>
<tr>
<td>cg10467968</td>
<td>0.3434299</td>
<td>0.0061458131</td>
<td>33.35419</td>
<td>4.818942e-06</td>
<td>0.9999983</td>
</tr>
<tr>
<td>cg00135841</td>
<td>-0.6673803</td>
<td>0.0189247545</td>
<td>28.63051</td>
<td>8.857500e-06</td>
<td>0.9999983</td>
</tr>
<tr>
<td>ch.17.28266240F</td>
<td>0.1352752</td>
<td>-0.0008316485</td>
<td>-26.87454</td>
<td>1.139697e-05</td>
<td>0.9999983</td>
</tr>
<tr>
<td>cg05389038</td>
<td>0.5306783</td>
<td>-0.0056651940</td>
<td>-0.2263167</td>
<td>2.257633e-05</td>
<td>0.9999983</td>
</tr>
<tr>
<td>cg21498547</td>
<td>-1.8114330</td>
<td>0.0008316485</td>
<td>21.46843</td>
<td>2.784158e-05</td>
<td>0.9999983</td>
</tr>
</tbody>
</table>
```

8 Bumphunter: to find differentially methylated regions (DMRs)

The \texttt{bumphunter} function in \texttt{minfi} is a version of the bump hunting algorithm [9] adapted to the 450k array, relying on the \texttt{bumphunter} function implemented in the eponym package \texttt{bumphunter} [10].

Instead of looking for association between a single genomic location and a phenotype of interest, \texttt{bumphunter} looks for genomic regions that are differentially methylated between two conditions. In the context of the 450k array, the algorithm first defines \textit{clusters} of probes. Clusters are simply groups of probes such that two consecutive probe locations in the cluster are not separated by more than some distance \texttt{mapGap}. Briefly, the algorithm first computes a t-statistic at each genomic location, with optional smoothing. Then, it defines a candidate region to be a cluster of probes for which all the t-statistics exceed a predefined threshold. To test for significance of the candidate regions, the algorithm uses permutations (defined by the parameter \texttt{B}). The permutation scheme is expensive, and can take a few days when the number of candidate bumps is large. To avoid wasting time, we propose the following guideline:

- Define your phenotype of interest
> pheno <- pData(gset.funnorm)$status
> designMatrix <- model.matrix(~ pheno)

• Run the algorithm with $B = 0$ permutation on the Beta-values, with a medium difference cutoff, say 0.2 (which corresponds to 20% difference on the Beta-values):

> dmrs <- bumphunter(gset.funnorm, design = designMatrix,
+     cutoff = 0.2, B=0, type="Beta")

• If the number of candidate bumps is large, say $> 30000$, increase the cutoff to reduce the number of candidate bumps. The rationale behind this is that the most of the additional candidate regions found by lowering the cutoff will be found to be non-significant after the permutation scheme, and therefore time can be saved by being more stringent on the cutoff (high cutoff).

• Once you have decided on the cutoff, run the algorithm with a large number of permutations, say $B = 1000$:

> dmrs <- bumphunter(gset.funnorm, design = designMatrix,
+     cutoff = 0.2, B=1000, type="Beta")

Since the permutation scheme can be expensive, parallel computation is implemented in the bumphunter function. The foreach package allows different parallel “back-ends” that will distribute the computation across multiple cores in a single machine, or across machines in a cluster. For instance, if one wished to use 3 cores, the two following commands have to be run before running bumphunter:

> library(doParallel)
> registerDoParallel(cores = 3)

The results of bumphunter are stored in a data frame with the rows being the different differentially methylated regions (DMRs):

> names(dmrs)
> head(dmrs$table, n=3)

As an example, we have run the bump hunting algorithm to find DMRs between colon and kidney (20 samples each from TCGA), with $B = 1000$ permutations, and a cutoff of 0.2 on the Beta values:

<table>
<thead>
<tr>
<th>chr</th>
<th>start</th>
<th>end</th>
<th>value</th>
<th>area</th>
<th>cluster</th>
<th>indexStart</th>
<th>indexEnd</th>
</tr>
</thead>
<tbody>
<tr>
<td>chr8</td>
<td>145103393</td>
<td>145107199</td>
<td>0.3767581</td>
<td>6.404887</td>
<td>194325</td>
<td>238277</td>
<td>238293</td>
</tr>
<tr>
<td>chr13</td>
<td>113425756</td>
<td>113428172</td>
<td>0.4257673</td>
<td>5.960743</td>
<td>57562</td>
<td>337302</td>
<td>337315</td>
</tr>
<tr>
<td>chr12</td>
<td>54446019</td>
<td>54447349</td>
<td>0.3278039</td>
<td>5.900470</td>
<td>46543</td>
<td>311839</td>
<td>311856</td>
</tr>
<tr>
<td>chr10</td>
<td>11206772</td>
<td>11208339</td>
<td>-0.4148710</td>
<td>5.393322</td>
<td>21466</td>
<td>251989</td>
<td>252001</td>
</tr>
<tr>
<td>chr10</td>
<td>130844121</td>
<td>130844899</td>
<td>-0.5869309</td>
<td>5.282378</td>
<td>29724</td>
<td>269921</td>
<td>269929</td>
</tr>
<tr>
<td>chr12</td>
<td>54409207</td>
<td>54409770</td>
<td>0.4370705</td>
<td>5.244846</td>
<td>46529</td>
<td>311742</td>
<td>311753</td>
</tr>
</tbody>
</table>

L clusterL p.value fwer p.valueArea fwerArea
The start and end columns indicate the limiting genomic locations of the DMR; the value column indicates the average difference in methylation in the bump, and the area column indicates the area of the bump with respect to the 0 line. The fwer column returns the family-wise error rate (FWER) of the regions estimated by the permeation scheme. One can filter the results by picking a cutoff on the FWER.

### 9 Other important topics

**Batch effects correction with SVA**

Surrogate variable analysis (SVA) [11, 12] is a useful tool to identified surrogate variables for unwanted variation while protecting for a phenotype of interest. In our experience, running SVA after normalizing the 450K data with preprocessFunnorm or preprocessQuantile increases the statistical power of the downstream analysis. For instance, to run SVA on the M-values, protecting for case-control status, the following code can be used to estimate the surrogate variables (this can take a few hours to run):

```r
> require(sva)
> mval <- getM(gset)
> pheno <- pData(gset)
> mod <- model.matrix(~as.factor(status), data=pheno)
> mod0 <- model.matrix(~1, data=pheno)
> sva.results <- sva(mval, mod, mod0)
```

Once the surrogate variables are computed, one can include them in the downstream analysis to adjust for unknown unwanted variation. See sea package vignette for a more comprehensive use of sva.

**Cell Type Composition**

As shown in [13], biological findings in blood samples can often be confounded with cell type composition. In order to estimate the confounding levels between phenotype and cell type composition, the function `estimateCellCounts` depending on the package FlowSorted.Blood.450k [14] estimates the cell type composition of blood samples by using a mod-
ified version of the algorithm described in [15]. The function takes as input a \texttt{RGChannelSet} and returns a cell counts vector for each samples:

\begin{verbatim}
> require(FlowSorted.Blood.450k)
> cellCounts <- estimateCellCounts(RGSet)
\end{verbatim}

**Block finder**

The approximately 170,000 open sea probes on the 450K array can be used to detect long-range changes in methylation status. These large scale changes that can range up to several Mb have typically been identified only through whole-genome bisulfite sequencing. The function \texttt{blockFinder} groups the average methylation values in open-sea probe cluster (via \texttt{cpg-Collapse}) into large regions, and then run the \texttt{bumphunter} algorithm with a large (250KB+) smoothing window (see the bump hunting section for DMRs above).

**Sex prediction**

By looking at the median total intensity of the X chromosome-mapped probes, denoted $med(X)$, and the median total intensity of the Y-chromosome-mapped probes, denoted $med(Y)$, one can observe two different clusters of points corresponding to which gender the samples belong to. To predict the gender, \texttt{minfi} separate the points by using a cutoff on $\log_2 med(Y) - \log_2 med(Y)$. The default cutoff is $-2$. Since the algorithm needs to map probes to the X-chr and to the Y-chr, the input of the function \texttt{getSex()} needs to be a \texttt{GenomicMethylSet} or a \texttt{GenomicRatioSet}.

\begin{verbatim}
> predictedSex <- getSex(gset, cutoff = -2)$predictedSex
> head(predictedSex)
\end{verbatim}

[1] "M" "F" "M" "F" "F" "F"

To choose the cutoff to separate the two gender clusters, one can plot $med(Y)$ against $med(Y)$ with the function \texttt{plotSex}:

\begin{verbatim}
> plotSex(getSex(gset, cutoff = -2))
\end{verbatim}

Finally, the function \texttt{addSex} applied to the \texttt{GenomicRatioSet} will add the predicted sex to the phenotype data. \textit{Remark}: the function does not handle datasets with only females or only males.
10 Advanced functions

getSnpBeta

The array contains by design 65 probes that are not meant to interrogate methylation status, but instead are designed to interrogate SNPs. By default, minfi drops these probes. The function `getSnpBeta` developmental version allows the user to extract the Beta values for those probes from an `RGChannelSet`. The return object is a matrix with the columns being the samples and the rows being the different SNP probes:

```r
> snps <- getSnpBeta(rgset)
> head(snaps)
```

These SNP probes are intended to be used for sample tracking and sample mixups. Each SNP probe ought to have values clustered around 3 distinct values corresponding to homo-, and hetero-zygotes.

Out-of-band (or ghost) probes

The function `getOOB` applied to an `RGChannelSet` retrieves the so-called “out-of-band” (OOB) probes. These are the measurements of Type I probes in the “wrong” color channel. The function returns a list with two matrices, named `Red` and `Grn`.

```r
> oob <- getOOB(rgset)
```

11 Exercises

1) Before processing a RGChannelSet further, could you remove the probes which failed more than 50% of the samples in the example dataset?

2) For the top loci that we found differentially methylated for the predicted sex, could you tell if those loci are mostly mapped to the X and Y chromosomes?

3) It is known that the Beta-value distribution of the Type I probes is different from the Beta value distribution of the Type II probes. Can you verify this with by plotting the Beta-value distribution density for each type separately?

References


