# The bumphunter user's guide

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## Introduction

This package implements the statistical procedure described in [1] (with some small modifications). Notably, batch effect removal needs additional code.

For any given type of data, it is usually necessary to make a number of choices and/or transformations before the bump hunting methodology is ready to be applied. Typically, these modifications resides in other packages. Examples are **charm** (for CHARM-like methylation microarrays), **bsseq** (for whole-genome bisulfite sequencing data) and **minfi** (for Illumina 450k methylation arrays). In some cases (specifically **bsseq**) only parts of the methodology as implemented in the **bumphunter** package is applied, although the conceptual approach is still build on bump hunting.

In other words, this package is mostly intended for developers wishing to adapt the general methodology to their specific applications.

The core of the package is encapsulated in the bumphunter method which uses the underlying bumphunterEngine to do the heavy lifting. However, bumphunterEngine consists of a number of useful functions that does much of the specific tasks involved in bump hunting. This document attempts to describe the overall workflow as well as the specific functions. The relevant functions are clusterMaker, getSegments, findRegions.

> library(bumphunter)

Note that this package is written with genomic data as an illustrative example but most of it is easily generalizable to other data types.

#### Other functions

Most of the **bumphunter** package is code for bump hunting. But we also include a number of convenience functions we have found useful, which are not necessarily part of the bump hunting exercise. At the time of writing, this include **annotateNearest**.

### The Statistical Model

The bump hunter methodology is meant to work on data with several biological replicates, similar to the lmFit function in limma. While the package is written using genomic data as an illustrative example, most of it is generalizable to other data types (with some one-dimensional location information).

We assume we have data  $Y_{ij}$  where *i* represents (biological) replicate and  $l_j$  represents genomic location. The use of *j* and  $l_j$  is a convenience notation, allowing us to discuss the "next" observation j + 1 which may be some distance  $lj + 1 - l_j$  away. Note that we assume in this notation that all replicates have been observed at the same set of genomic locations.

The basic statistical model is the following:

$$Y_{ij} = \beta_0(l_j) + \beta_1(l_j)X_j + \varepsilon_{ij}$$

with *i* representing subject,  $l_j$  representing the *j*th location,  $X_j$  is the covariate of interest (for example  $X_j = 1$  for cases and  $X_j = 0$  otherwise),  $\varepsilon_{ij}$  is measurement error,  $\beta_0(l)$  is a baseline function, and  $\beta_1(l)$  is the parameter of interest, which is a function of location. We assume that  $\beta_1(l)$  will be equal to zero over most of the genome, and we want to identify stretched where  $\beta_1(l) \neq 0$ , which we call *bumps*.

We want to share information between nearby locations, typically through some form of smoothing.

### **Creating clusters**

For many genomic applications the locations are clustered. Each cluster is a distinct unit where the model fitting will be done separately, and each cluster does not depend on the data, only on the locations  $l_j$ . Typically there is some maximal distance, and we do not want to smooth between observations separated by more than this distance. The choice of distance is very application dependent.

"Clusters" are simply groups of locations such that two consecutive locations in the cluster are separated by less than some distance maxGap. For genomic applications, the biggest possible clusters are chromosomes.

The function clusterMaker defines such grouping locations.

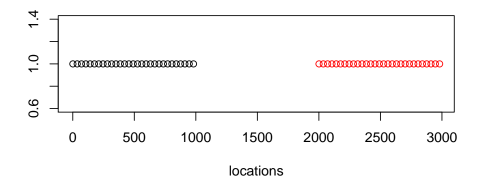
Example: We first generate an example of typical genomic locations

Now we run the function to obtain the three clusters from the positions. We use the default gap of 300 base pairs (bps), i.e. any two points more than 300 bps away are put in a new cluster. Also, locations from different chromosomes are separated.

```
> cl <- clusterMaker(chr, pos, maxGap = 300)
> table(cl)
cl
    1   2   3
29   29   20
```

The output is an indexing variable telling us which cluster each location belongs to. Locations on different chromosomes are always on different clusters.

Note that data from the first chromosome has been split into two clusters:



### Breaking into segments

The function getSegments is used to find segments that are positive, near zero, and negative. Specifically we have a vector of numbers  $\theta_j$  with each number associated with a genomic location  $l_j$  (thinks either test statistics or estimates of  $\beta_i(l)$ ). A segment is a list of consecutive locations such that all  $\theta_l$  in the segment are either "positive", "near zero" or "negative". In order to define "positive" etc we need a cutoff which is one number L (in which case "near zero" is [-L, L]) or two numbers L, U (in which case "near zero" is [L; U]).

Example: we are going to create a simulated  $\beta_1(l)$  with a couple of real bumps.

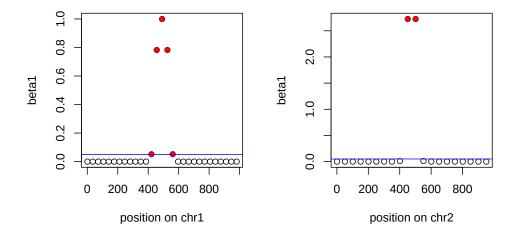
```
> Indexes <- split(seq_along(cl), cl)
> beta1 <- rep(0, length(pos))
> for(i in seq(along=Indexes)){
+     ind <- Indexes[[i]]
+     x <- pos[ind]
+     z <- scale(x, median(x), max(x)/12)
+     beta1[ind] <- i*(-1)^(i+1)*pmax(1-abs(z)^3,0)^3 ##multiply by i to vary size
+ }
```

We now find bumps of this functions by

> segs <- getSegments(beta1, cl, cutoff=0.05)</pre>

Now we can make, for example, a plot of all the positive bumps

```
> par(mfrow=c(1,2))
> for(ind in segs$upIndex){
+     index <- which(cl==cl[ind[1]])
+     plot(pos[index], beta1[index],
+          xlab=paste("position on", chr[ind[1]]),
+         ylab="beta1")
+     points(pos[ind], beta1[ind], pch=16, col=2)
+     abline(h = 0.05, col = "blue")
+ }</pre>
```



This function is used by regionFinder which is described next.

## regionFinder

This function packages up the results of getSegments into a table of regions with the location and characteristics of bumps.

```
> tab <- regionFinder(beta1, chr, pos, cl, cutoff=0.05)
> tab
```

```
area cluster indexStart indexEnd L
   chr start end
                       value
                                             2
3 chr1
      2281 2701 -1.2636037 16.426848
                                                       38
                                                                50 13
2 chr2
        451
             501 2.7262463 5.452493
                                             3
                                                       68
                                                                69
                                                                    2
1 chr1
        421
             561 0.5336474 2.668237
                                             1
                                                       13
                                                                17 5
 clusterL
3
       29
2
       20
1
       29
```

In the plot in the preceding section we show two of these regions in red.

Note that regionFinder and getSegments do not really contain any statistical model. All it does is finding regions based on segmenting a vector  $\theta_j$  associated with genomic locations  $l_j$ .

#### Bumphunting

Bumphunter is a more complicated function. In addition to regionFinder and clusterMaker it also implements a statistical model as well as permutation testing to assess uncertainty.

We start by creating a simulated data set of 10 cases and 10 controls (recall that **beta1** was defined above).

```
> beta0 <- 3*sin(2*pi*pos/720)
> X <- cbind(rep(1,20), rep(c(0,1), each=10))
> error <- matrix(rnorm(20*length(beta1), 0, 1), ncol=20)
> y <- t(X[,1])%x%beta0 + t(X[,2])%x%beta1 + error</pre>
```

Now we can run bumphunter

```
> tab <- bumphunter(y, X, chr, pos, cl, cutoff=.5)
> names(tab)
[1] "table" "fitted" "pvaluesMarginal"
[4] "null"
```

```
> tab$table
   chr start
              end
                        value
                                   area cluster indexStart indexEnd L
1 chr1
        2176 2701 -0.7715593 12.34495
                                              2
                                                         35
                                                                  50 16
  clusterL p.value fwer p.valueArea fwerArea
1
        29
                 0
                       0
                                    0
                                             0
```

Briefly, the **bumphunter** function fits a linear model for each location (like **lmFit** from the **limma** package), focusing on one specific column (coefficient) of the design matrix. This coefficient of interest is optionally smoothed. Subsequently, a permutation test is formed for this specific coefficient.

#### Faster bumphunting with multiple cores

bumphunter can be speeded up by using multiple cores. We use the foreach package which allows different parallel "back-ends" that will distribute the computation across multiple cores in a single machine, or across multiple machines in a cluster. The most straightforward usage, illustrated below, involves multiple cores on a single machine. See the foreach documentation for more complex use cases, as well as the packages doParallel and doSNOW (among others). Finally, we use doRNG to ensure reproducibility of setting the seed within the parallel computations.

In order to use the **foreach** package we need to register a backend, in this case a multicore machine with 2 cores.

```
> library(doParallel)
> registerDoParallel(cores = 2)
```

bumphunter will now automatically use this backend

```
> tab <- bumphunter(y, X, chr, pos, cl, cutoff=.5, B=250, verbose = TRUE)</pre>
```

### References

[1] Andrew E Jaffe, Peter Murakami, Hwajin Lee, Jeffrey T Leek, M Daniele Fallin, Andrew P Feinberg, and Rafael A Irizarry. Bump hunting to identify differentially methylated regions in epigenetic epidemiology studies. International Journal of Epidemiology, 41(1):200–209, 2012.

### Cleanup

This is a cleanup step for the vignette on Windows; typically not needed for users.

```
> bumphunter:::closeSockets()
```

### SessionInfo

- R version 3.0.1 (2013-05-16), x86\_64-unknown-linux-gnu
- Locale: LC\_CTYPE=en\_US.UTF-8, LC\_NUMERIC=C, LC\_TIME=en\_US.UTF-8, LC\_COLLATE=C, LC\_MONETARY=en\_US.UTF-8, LC\_MESSAGES=en\_US.UTF-8, LC\_PAPER=C, LC\_NAME=C, LC\_ADDRESS=C, LC\_TELEPHONE=C, LC\_MEASUREMENT=en\_US.UTF-8, LC\_IDENTIFICATION=C
- Base packages: base, datasets, grDevices, graphics, methods, parallel, stats, utils
- Other packages: BiocGenerics 0.6.0, GenomicRanges 1.12.4, IRanges 1.18.2, bumphunter 1.0.2, doParallel 1.0.3, doRNG 1.5.3, foreach 1.4.1, iterators 1.0.6, pkgmaker 0.16, registry 0.2, rngtools 1.2.1
- Loaded via a namespace (and not attached): R.methodsS3 1.4.4, codetools 0.2-8, compiler 3.0.1, digest 0.6.3, itertools 0.1-1, limma 3.16.7, matrixStats 0.8.5, stats4 3.0.1, stringr 0.6.2, tools 3.0.1, xtable 1.7-1