CGHnormaliter Package (Version 1.20.0)

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1 Introduction

This package contains an implementation of the CGHnormaliter strategy for normalization of two-channel array Comparative Genomic Hybridization (aCGH) data displaying many copy number imbalances. The key idea of our method is that temporary exclusion of aberrations from the aCGH data allows for a more appropriate calculation of the LOWESS regression curve. As a result, after normalization, the log₂ intensity ratios of the normals will generally be closer to zero and better reflect the biological reality. We coined this normalization strategy 'local-LOWESS' since only a subset of the log₂ ratios is considered in the LOWESS regression.

The strategy can be summarized as follows (see Figure 1). Initially the log₂ intensity ratios are segmented using DNAcopy [5]. The segmented data are then given as input to a calling tool named CGHcall [2] to discriminate the normals from gains and losses. These normals are subsequently used for normalization based on LOWESS. These steps are then iterated to refine the normalization. For more detailed information we refer to the publications of the method [4, 3].

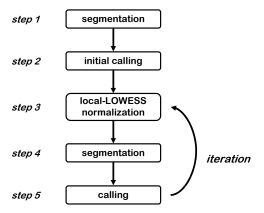


Figure 1: Overview of the CGHnormaliter method.

2 Data format

The input should be either a data.frame or the file name of a tabseparated text file (text files must contain a header). The first four columns should describe the clone and its position on the genome:

- 1. ID: The unique identifiers of array elements.
- 2. Chromosome: Chromosome number of each array element.
- 3. Start: Chromosomal start position in bp of each array element.
- 4. End: Chromosomal end position in bp of each array element.

The start and end positions must be numeric. The next columns hold the actual data. For each sample in the experiment, there must be two adjacent columns with the *test* and *reference* intensities, respectively. All entries must be delimited by tabs, and missing entries must be denoted with NA or by an empty value. Below, an example is given of a correctly formatted data file or data frame containing measurements on 7 clones in 2 samples.

ID	Chromosome	Start	End	Case1.test	Case1.ref	Case2.test	Case2.ref
RP11-34P13	1	1	254479	279	294	NA	NA
RP11-379K15	1	95421	244136	1815	2269	2793	3996
RP11-776018	1	357737	465038	387	349	429	362
RP11-45C18	1	579118	696613	786	734	900	735
RP11-242B5	1	606617	711982	2955	4158	4478	5229
RP13-586C17	1	619355	783174	NA	NA	823	841
RP11-414L23	1	658751	846904	630	937	959	744

3 Example

First, we load the example acute lymphoblastic leukemia dataset [1] which comes with the CGHnormaliter package:

- > library(CGHnormaliter)
- > data(Leukemia)

Next, we run the CGHnormaliter routine on the first four chromosomes of the Leukemia data:

```
> result <- CGHnormaliter(Leukemia, nchrom=4, cellularity=0.9)
```

CGHnormaliter -- Running an initial segmentation and calling Start data segmentation .. Analyzing: Sample.1 $\,$

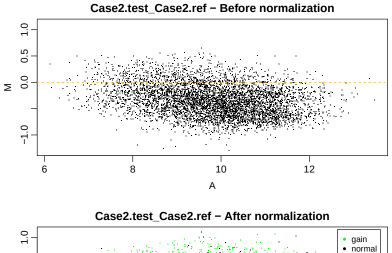
Analyzing: Sample.1 Analyzing: Sample.2 Analyzing: Sample.3 Start data calling ...

```
CGHnormaliter -- Iteration # 1
Mean normalization shift per sample:
   Case1.test_Case1.ref : 0.1053031
   Case2.test_Case2.ref : 0.07000189
   Case3.test_Case3.ref : 0.128865
Start data segmentation ...
Analyzing: Sample.1
Analyzing: Sample.2
Analyzing: Sample.3
Start data calling ...
CGHnormaliter -- Iteration # 2
Mean normalization shift per sample:
   Case1.test_Case1.ref : 0.00856964
   Case2.test_Case2.ref : 0.0162889
   Case3.test_Case3.ref : 0.01226434
Start data segmentation ...
Analyzing: Sample.1
Analyzing: Sample.2
Analyzing: Sample.3
Start data calling ...
CGHnormaliter -- Iteration # 3
Mean normalization shift per sample:
   Case1.test_Case1.ref : 0.001326081
   Case2.test_Case2.ref : 0.002476787
   Case3.test_Case3.ref : 0.001817676
CGHnormaliter -- Reached convergence. Running a final segmentation and calling...
Start data segmentation ...
Analyzing: Sample.1
Analyzing: Sample.2
Analyzing: Sample.3
Start data calling ...
Writing MA-plots to file: MAplot.pdf
CGHnormaliter -- FINISHED
```

To enable a visual assessment of the bias reduction, MA-plots are (by default) automatically generated before and after normalization of each sample. These plots are stored into a PDF, usually named MAplot.pdf (the exact file name is supplied at the end of each CGHnormaliter run). See Figure 2 for such MA-plots of the second Leukemia sample.

Now, several fields of the result object can be accessed, for example:

```
> normalized.data <- copynumber(result) # log2 ratios
> segmented.data <- segmented(result)
> called.data <- calls(result)</pre>
```



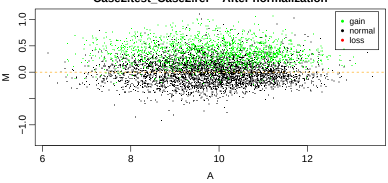


Figure 2: MA-plot for the second leukemia sample before and after normalization. Note that the normalization is based on the normals only, represented by the black dots.

Plotting the normalized log₂ ratios in a density plot provides another means (besides MA plots) to inspect whether or not the centralization has been successful. Figure 3 shows such a density plot for sample 2:

The results, including segments and calls, can be visualized using the plot function. In Figure 4 the results of sample 2 are plotted in full resolution:

```
> plot(result[,2], ylimit=c(-2,2), dotres=1)
```

Plotting sample Case2.test_Case2.ref

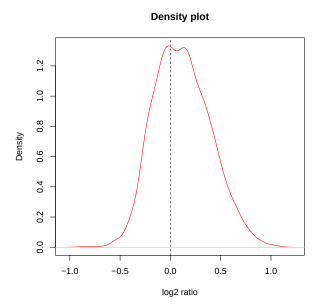


Figure 3: Density plot after CGHnormaliter normalization for the second leukemia sample. The data are adequately centralized around peak at the left, which corresponds to the normals. The peak at the right corresponds to the gains.

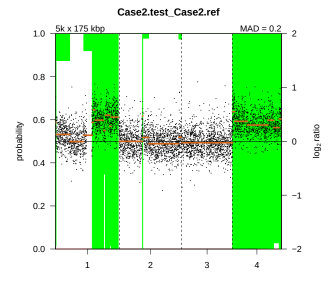


Figure 4: Results of the CGHnormal mediation for the second leukemia sample.

Finally, the package provides the function CGHnormaliter.write.table to save the normalized data into a tabdelimited plain text file:

> CGHnormaliter.write.table(result)

Saving normalized log2 ratios to file: normalized.txt

The segmented and called data from the **result** object can be saved to file as well using this function:

> CGHnormaliter.write.table(result, data.type="segmented")

Saving segmented log2 ratios to file: segmented.txt

> CGHnormaliter.write.table(result, data.type="called")

Saving calls to file: called.txt

References

- [1] K. Paulsson, M. Heidenblad, H. Mörse, Å. Borg, T. Fioretos, and B. Johansson. Identification of cryptic aberrations and characterization of translocation breakpoints using array CGH in high hyperdiploid childhood acute lymphoblastic leukemia. *Leukemia*, 20:2002–2007, 2006.
- [2] M.A. van de Wiel, K.I. Kim, S.J. Vosse, W.N. van Wieringen, S.M. Wilting, and B. Ylstra. CGHcall: calling aberrations for array CGH tumor profiles. *Bioinformatics*, 23:892–894, 2007.
- [3] B.P.P. van Houte, T.W. Binsl, H. Hettling, and J. Heringa. CGHnormaliter: a bioconductor package for normalization of array CGH data with many CNAs. *Bioinformatics*, 26(10):1366–1367, 2010.
- [4] B.P.P. van Houte, T.W. Binsl, H. Hettling, W. Pirovano, and J. Heringa. CGHnormaliter: an iterative strategy to enhance normalization of array CGH data with imbalanced aberrations. *BMC Genomics*, 10:401, 2009.
- [5] E.S. Venkatraman and A.B. Olshen. A faster circular binary segmentation algorithm for the analysis of array CGH data. *Bioinformatics*, 23(6):657–663, 2007.