Software Manual





cn.mops - Mixture of Poissons for CNV detection in NGS data

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1 Introduction 3

1 Introduction

The cn.mops package is part of the Bioconductor (http://www.bioconductor.org) project. The package allows to detect copy number variations (CNVs) from next generation sequencing (NGS) data sets based on a generative model. Please visit http://www.bioinf.jku.at/software/cnmops/cnmops.html for additional information.

To avoid the false discoveries induced by read count variations along the chromosome or across samples, we propose a "Mixture Of PoissonS model for CNV detection" (cn.MOPS). The cn.MOPS model is not affected by read count variations along the chromosome, because at each DNA position a local model is constructed. Read count variations across samples are decomposed by the cn.MOPS model into integer copy numbers and noise by its mixture components and Poisson distributions, respectively. In contrast to existing methods, cn.MOPS model's posterior provides integer copy numbers together with their uncertainty. Model selection in a Bayesian framework is based on maximizing the posterior given the samples by an expectation maximization (EM) algorithm. The model incorporates the linear dependency between average read counts in a DNA segment and its copy number. Most importantly, a Dirichlet prior on the mixture components prefers constant copy number 2 for all samples. The more the data drives the posterior away from the Dirichlet prior corresponding to copy number two, the more likely the data is caused by a CNV, and, the higher is the informative/non-informative (I/NI) call. cn.MOPS detects a CNV in the DNA of an individual as a segment with high I/NI calls. I/NI call based CNV detection guarantees a low false discovery rate (FDR) because wrong detections are less likely for high I/NI calls. We assume that the genome is partitioned into segments in which reads are counted but which need not be of constant length throughout the genome. For each of such an segment we build a model. We consider the read counts x at a certain segment of the genome, for which we construct a model across samples. The model incorporates both read count variations due to technical or biological noise and variations stemming from copy number variations. For further information regarding the algorithm and its assessment see the cn.MOPS homepage at http://www.bioinf.jku.at/software/cnmops/cnmops.html.

2 Getting started and quick start

To load the package, enter the following in your R session:

```
> library(cn.mops)
```

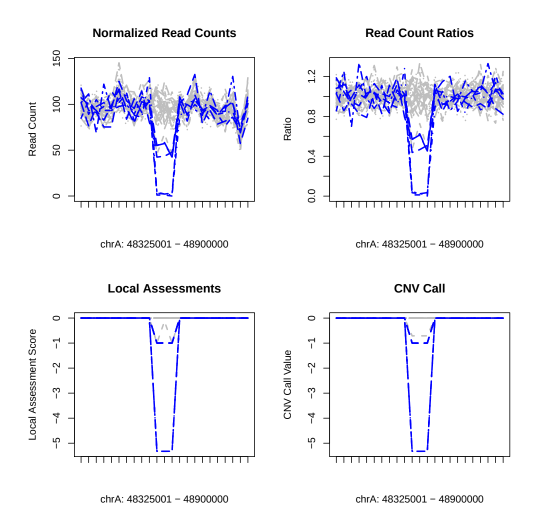
The whole pipeline will only take a few steps, if BAM files are available (for read count matrices directly go to step 2):

- 1. Getting the input data from BAM files (also see Section 3.2 and Section 3).
 - > BAMFiles <- list.files(pattern=".bam\$")</pre>
 - > bamDataRanges <- getReadCountsFromBAM(BAMFiles)
- 2. Running the algorithm (also see Section 4.2).

```
> res <- cn.mops(bamDataRanges)
```

3. Visualization of the detected CNV regions. For more information about the result objects and visualization see Section 4.2.

> plot(res, which=1)



3 Input of cn.mops: BAM files, GRanges objects, or numeric matrices

3.1 Read count matrices as input

cn.mops does not require the data samples to be of any specific kind or structure. cn.mops only requires a read count matrix, i.e., given N data samples and m genomic segments, this is an $m \times N$ real- or integer-valued matrix \mathbf{X} , in which an entry x_{ij} corresponds to the read count of sample j in the i-th segment. E.g. in the following read count matrix sample three has 17 reads in the second segment: $x_{23} = 71$.

cn.mops can handle numeric and integer matrices or GRanges objects, in which the read counts are stored as values of the object.

3.2 BAM files as input

The most widely used file format for aligned short reads is the Sequence Alignment Map (SAM) format or in the compressed form the Binary Alignment Map (BAM). We provide a simple function that makes use of the Rsamtools package to obtain the alignment positions of reads. The result object of the function can directly be used as input for cn.mops. The author can provide functions for input formats other than BAM upon request: cn.mops@bioinf.jku.at.

```
> BAMFiles <- list.files(system.file("extdata", package="cn.mops"),pattern=".bam$",
+ full.names=TRUE)
> bamDataRanges <- getReadCountsFromBAM(BAMFiles,
+ sampleNames=paste("Sample",1:3),mode="unpaired")</pre>
```

In bamDataRanges you have now stored the genomic segments (left of the |'s) and the read counts (right of the |'s):

> (bamDataRanges)

GRanges with 9 ranges and 3 metadata columns:

	seqnames		ranges	strand		Sample.1	Sample.3	Sample.2
	<rle></rle>		Ranges	<rle></rle>		<integer></integer>	<integer></integer>	<integer></integer>
[1]	20	[1,	7363000]	*	I	856	25	3
[2]	20	[7363001,	14726000]	*	I	0	0	0
[3]	20	[14726001,	22089000]	*	I	0	0	0
[4]	20	[22089001,	29452000]	*		0	0	0
[5]	20	[29452001,	36815000]	*	I	0	0	0
[6]	20	[36815001,	44178000]	*	I	0	0	0
[7]	20	[44178001,	51541000]	*	I	0	0	0
[8]	20	[51541001,	58904000]	*	I	0	0	0
[9]	20	[58904001,	63025519]	*		0	0	0

seqlengths:

20

63025520

4 Copy number estimation with cn.mops

To get a first impression, we use a data set, in which CNVs have been artificially implanted. The simulated data set was generated using distributions of read counts as they appear in real sequencing experiments. CNVs were implanted under the assumption that the expected read count is linear dependent on the copy number. For example in a certain genomic we expect λ reads for copy number 2, then we expect 2λ reads for copy number 4. The linear relationship was confirmed in different studies, like Alkan *et al.* (2009), Chiang *et al.* (2008) and Sathirapongsasuti *et al.* (2011).

4.1 Running cn. mops

The read counts are stored in the objects X and XRanges, which are the two basic input types that cn.mops allows:

```
> data(cn.mops)
> ls()

[1] "BAMFiles" "CNVRanges" "X" "XRanges"
[5] "bamDataRanges" "cn.mopsVersion" "exomeCounts" "resCNMOPS"
```

The same data is stored in a GRanges object, in which we see the genomic coordinates, as well as the read counts (values):

```
> head(XRanges[,1:3])
```

GRanges with 6 ranges and 3 metadata columns:

	seqnames	_	ranges	strand		S_1	S_2	S_3
	<rle></rle>	<	IRanges>	<rle></rle>		<integer></integer>	<integer></integer>	<integer></integer>
[1]	chrA	[1,	25000]	*		102	93	109
[2]	chrA	[25001,	50000]	*		118	99	100
[3]	chrA	[50001,	75000]	*		85	81	82
[4]	chrA	[75001,	100000]	*		87	116	106
[5]	chrA	[100001,	125000]	*	1	87	68	89
[6]	chrA	[125001,	150000]	*	1	87	91	91
seq.	lengths:							

sequenguns: chrA

NΑ

We are now ready to run cn.mops on the GRanges object:

```
> resCNMOPS <- cn.mops(XRanges)</pre>
```

To calculate integer copy number use the command calcIntegerCopyNumbers:

> resCNMOPS <- calcIntegerCopyNumbers(resCNMOPS)</pre>

Alternatively, it is possible to use an integer matrix, in which the genomic coordinates can be stored as rownames and the entries are the read counts. For example the data from above represented by an integer matrix X:

> head(X[,1:3])

	S_1	S_2	S_3
Chr_A_1_25000	102	93	109
Chr_A_25001_50000	118	99	100
Chr_A_50001_75000	85	81	82
Chr_A_75001_100000	87	116	106
Chr_A_100001_125000	87	68	89
Chr_A_125001_150000	87	91	91

We are now ready to run cn.mops on the integer matrix:

```
> resCNMOPSX <- cn.mops(X)</pre>
```

To calculate integer copy number use the command calcIntegerCopyNumbers:

```
> resCNMOPSX <- calcIntegerCopyNumbers(resCNMOPSX)</pre>
```

Note that the two results resCNMOPS and resCNMOPSRanges identify the same CNVs:

```
> all(individualCall(resCNMOPSX)==individualCall(resCNMOPS))
```

4.2 The result object

To get a summary of the CNV detection result, just enter the name of the object (which implicitly calls show):

> (resCNMOPS)

The CNVs per individual are stored in the slot cnvs:

> cnvs(resCNMOPS)[1:5]

GRanges with 5 ranges and 4 metadata columns:

median	sampleName		strand	ranges		seqnames	
<numeric></numeric>	<factor></factor>		<rle></rle>	<iranges></iranges>		<rle></rle>	
-1.0000000	S_1		*	1850000]	[1775001,	chrA	[1]
-5.3219281	S_1		*	53400000]	[53300001,	chrA	[2]
-0.9999143	S_1		*	113325000]	[113175001,	chrA	[3]
-5.3219281	S 2	1	*	486500001	[48575001,	chrA	۲4٦

```
[5]
        chrA [ 70625001, 70750000]
                                           *
                                                       S_2 -5.3219281
          mean
     <numeric> <character>
[1] -0.9999995
                        CN1
[2] -5.3219281
                        CNO
[3] -0.9997036
                        CN1
[4] -5.3219281
                        CNO
[5] -5.3219281
                        CNO
seqlengths:
chrA
   NΑ
```

Segments, in which individual CNVs accumulate, are called CNV regions and can be accessed by cnvr:

```
> cnvr(resCNMOPS)[1,1:5]
```

NA

```
GRanges with 1 range and 5 metadata columns:
      seqnames
                            ranges strand |
                                                           S_2
                                                                     S_3
                                                  S_1
         <Rle>
                         <IRanges> <Rle> | <factor> <factor> <factor>
          chrA [1775001, 1850000]
                                         *
                                                           CN2
  [1]
                                                  CN1
                                                                     CN2
           S_4
                     S_5
      <factor> <factor>
  [1]
           CN2
                    CN2
  seqlengths:
   chrA
     NA
```

We now want to check, whether cn.mops found the implanted CNVs. We have stored the implanted CNVs (see beginning of Section) in the object CNVRanges.

```
> (CNVRanges[15,1:5])
GRanges with 1 range and 5 metadata columns:
      seqnames
                            ranges strand |
                                                   S_1
                                                             S_2
                                                                        S_3
         <Rle>
                         <IRanges> <Rle> | <integer> <integer> <integer>
          chrA [1751017, 1850656]
                                        *
  [1]
                                                     1
            S_4
                      S_5
      <integer> <integer>
  [1]
              2
  seqlengths:
   chrA
```

Next we identify overlaps between CNVs that were detected by cn.mops and CNVs that were implanted. Towards this end we use the functions of the GenomicRanges package.

```
> ranges(cnvr(resCNMOPS))[1:2]
```

```
IRanges of length 2

start end width

[1] 1775001 1850000 75000

[2] 5525001 5600000 75000
```

```
> ranges(cnvr(resCNMOPS)) %over% ranges(CNVRanges)
```

The detected CNV regions all overlap with the known CNV regions contained in CNVRanges.

The function cn.mops creates an instance of the S4 class CNVDetectionResult that is defined by the present package. To get detailed information on which data are stored in such objects, enter

```
> help(CNVDetectionResult)
```

5 Visualization of the result

5.1 Chromosome plots

cn.mops allows for plotting the detected segments of an individual at one chromosome by a plot similar to the ones produced by DNAcopy:

```
> segplot(resCNMOPS, sampleIdx=13)
```

S_13 Chromosome chrA

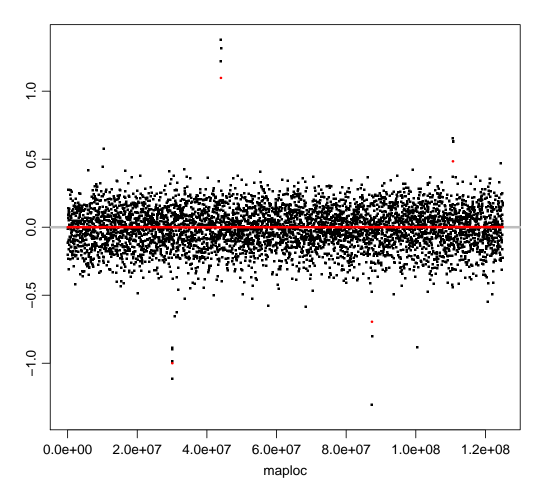


Figure 1: The x-axis represents the genomic position and on the y-axis we see the log ratio of the read counts (green) and the copy number call of each segment (red).

5.2 CNV region plots

cn.mops allows for plotting the detected CNV regions:

> plot(resCNMOPS, which=1)

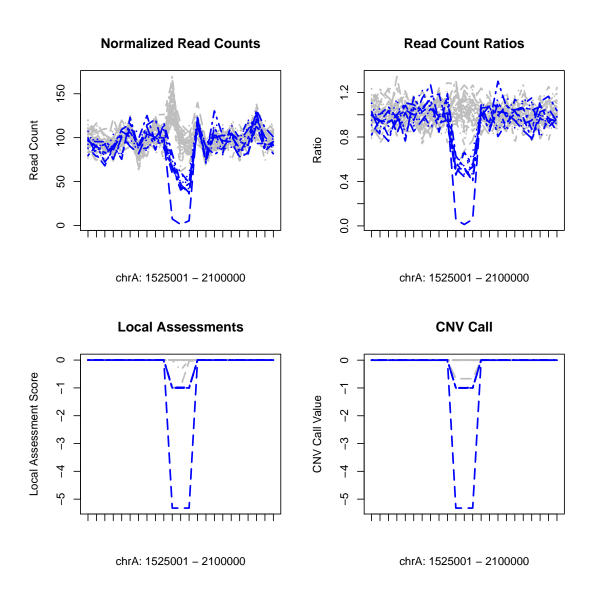


Figure 2: The x-axis represents the genomic position and on the y-axis we see the read counts (left), the call of the local model (middle) and the CNV call produced by the segmentation algorithm. Blue lines mark samples having a copy number loss.

6 Exome sequencing data

To apply cn.mops to exome sequencing data requires a different preprocessing, since constant windows spanning the whole genome are not appropriate. The initial segments in which the reads are counted should be chosen as the regions of the baits, targets or exons. The read count matrix can now be generated by using the function getSegmentReadCountsFromBAM that requires the genomic coordinates of the predefined segments as GRanges object. The resulting read count

matrix can directly be used as input for cn.mops. A possible processing script could look like the following:

```
> library(cn.mops)
> BAMFiles <- list.files(pattern=".bam$")
> segments <- read.table("targetRegions.bed",sep="\t",as.is=TRUE)
> gr <- GRanges(segments[,1],IRanges(segments[,2],segments[,3]))
> X <- getSegmentReadCountsFromBAM(BAMFiles,GR=gr,mode="unpaired")
> resCNMOPS <- exomecn.mops(X)
> resCNMOPS <- calcIntegerCopyNumbers(resCNMOPS)</pre>
```

We included an exome sequencing data set in this package. It is stored in exomeCounts.

```
> resultExomeData <- exomecn.mops(exomeCounts)
> resultExomeData <- calcIntegerCopyNumbers(resultExomeData )</pre>
```

```
> plot(resultExomeData, which=5)
```

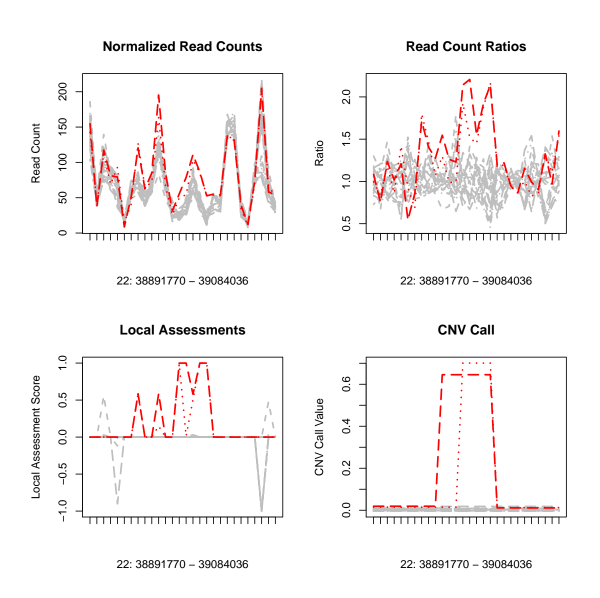


Figure 3: ExomeSeq data results.

Possible issues and notes A problem can occur, if the names of the reference sequences, e.s. chromosomes, are inconsistent between the bed file and the bam file. For example "chr1", "chr2",...,"chrX","chrY" and "1","2",...,"X","Y". This can easily be solved by replacing the sequevels of the GRanges object:

```
> #the following removes the "chr" from reference sequence names
> seqlevels(gr) <- gsub("chr","",seqlevels(gr))</pre>
```

Results can also be improved if you extend your target regions by a small amount of bases to the left and to the right (in the following case it is 30bp):

```
> gr <- GRanges(segments[,1], IRanges(segments[,2]-30, segments[,3]+30))
> gr <- reduce(gr)</pre>
```

7 Cases vs. Control or Tumor vs. Normal

For detection of CNVs in a setting in which the normal state is known, the function referencecn.mops can be applied. It implements the cn.MOPS algorithm adjusted to this setting. For tumor samples very high copy numbers can be present – the maximum copy number with the default setting is 8 – and cn.mops has to be adjusted to allow higher copy numbers.

```
> resRef <- referencecn.mops(cases=X[,1],controls=rowMeans(X),
                   classes=c("CNO", "CN1", "CN2", "CN3", "CN4", "CN5", "CN6",
                                    "CN7", "CN8", "CN16", "CN32", "CN64", "CN128"),
+
                   I = c(0.025, 0.5, 1, 1.5, 2, 2.5, 3, 3.5, 4, 8, 16, 32, 64),
+
                   segAlgorithm="DNAcopy")
Analyzing: Sample.1
> resRef <- calcIntegerCopyNumbers(resRef)</pre>
> (cnvs(resRef))
GRanges with 1 range and 4 metadata columns:
      seqnames
                      ranges strand | sampleName
                                                     median
                                                                  mean
                   <IRanges>
                              <Rle>
                                         <factor> <numeric> <numeric>
         <Rle>
  [1]
         undef [2133, 2136]
                                  *
                                           Case_1 -5.321928
                                                               -4.3686
               CN
      <character>
  [1]
              CNO
  seqlengths:
   undef
      ΝA
```

8 Heterosomes and CNVs of tumor samples

With the default settings the normalization procedure assumes that the ploidy of each sample is the same. However, it is possible to account for different karyotypes. When analyzing CNVs on the X or Y chromosome one possibility is to treat males and females separately. The second option is to provide the normalization function with the information about the gender, that is different ploidy states of the X and Y chromosome. This can be handled by the ploidy parameter of the normalization function. In the following we show the normalization for the X chromosome, if the first 10 individuals are males (ploidy set to 1) and the next 30 individuals are females (ploidy set to 2):

```
> XchrX <- normalizeChromosomes(X[1:500, ],ploidy=c(rep(1,10),rep(2,30)))
> cnvr(calcIntegerCopyNumbers(cn.mops(XchrX,norm=FALSE)))
GRanges with 1 range and 40 metadata columns:
                    ranges strand |
                                            S_1
                                                       S_2
                                                                 S_3
                                                                           S_4
       seqnames
          <Rle> <IRanges>
                             <Rle> | <factor> <factor> <factor> <factor>
  \lceil 1 \rceil
          undef
                  [1, 500]
                                  *
                                            CN1
                                                       CN1
                                                                 CN1
                                                                           CN1
            S_5
                       S_6
                                 S_7
                                           S 8
                                                      S 9
                                                               S_10
                                                                         S_11
       <factor> <factor> <factor> <factor> <factor> <factor> <factor> <factor>
            CN1
                                           CN1
                                                      CN1
                                                                CN1
  [1]
                      CN1
                                 CN1
                                                                          CN2
           S_12
                     S_13
                                S_14
                                          S_15
                                                    S_16
                                                               S_17
                                                                         S 18
       <factor> <factor> <factor> <factor> <factor> <factor> <factor> <factor>
  [1]
            CN1
                      CN2
                                 CN2
                                           CN1
                                                     CN2
                                                                CN1
                                                                          CN2
           S_19
                     S_20
                                S_21
                                          S_22
                                                    S_{23}
                                                               S_24
                                                                         S<sub>25</sub>
       <factor> <factor> <factor> <factor> <factor> <factor> <factor> <factor>
  [1]
            CN2
                      CN2
                                 CN2
                                           CN2
                                                      CN2
                                                                CN2
                                                                          CN2
           S<sub>26</sub>
                     S_27
                                S_28
                                          S_29
                                                    S_30
                                                               S_31
                                                                         S_32
       <factor> <factor> <factor> <factor> <factor> <factor> <factor>
  [1]
            CN2
                      CN1
                                 CN1
                                           CN2
                                                     CN2
                                                                CN2
                                                                          CN2
           S 33
                     S 34
                                S 35
                                          S 36
                                                    S 37
                                                               S_38
                                                                         S 39
       <factor> <factor> <factor> <factor> <factor> <factor> <factor> <factor>
  [1]
            CN2
                      CN2
                                 CN2
                                           CN2
                                                      CN2
                                                                CNO
                                                                          CN2
           S_40
       <factor>
  [1]
            CN1
  seglengths:
   undef
      NΑ
```

Karyotype information can also improve results of CNV detection in tumor samples. The best results can be reached, if for each chromosome the number of appearances in the cell is known. In this case normalization should be applied to each chromosome separately.

9 cn.mops for haploid genomes

For haploid genomes the prior assumption is that all samples have copy number 1. The function haplocn.mops implements the cn.MOPS algorithm adjusted to haploid genomes.

```
> resHaplo <- haplocn.mops(X)
> resHaplo <- calcIntegerCopyNumbers(resHaplo)</pre>
```

10 Adjusting sensitivity, specificity and resolution for specific applications

The default parameters of both the local models of cn.mops and the segmentation algorithm were optimized on a wide ranged of different data sets. However, you might want to adjust sensitivity and specificity or resolution to your specific needs.

- upperThreshold The calling threshold for copy number gains, a positive value. Lowering the threshold will increase the detections, raising will decrease the detections.
- lowerThreshold The calling threshold for copy number losses, a negative value. Raising the threshold will increase the detections, lowering will decrease the detections.
 - priorImpact This parameter should be optimized for each data set, since it is influenced by number of samples as well as noise level. The higher the value, the more samples will have copy number 2, and consequently less CNVs will be detected.
 - minWidth The minimum length of CNVs measured in number of segments. The more adjacent segments with a high or low copy number call are joined, the higher the confidence in the detections. A lower value will lead to more shorter segments, and a higher value will yield to less, but longer segments.

The length of the initial segments is also crucial. They should be chosen such that on average 50 to 100 reads lie in one segment. The WL parameter of getReadCountsFromBAM determines this resolution.

11 Overview of study designs and cn.mops functions

In Table 1 we give an overview of the functions implemented in this package and present settings for which they are appropriate. All these functions work for multiple, at least two, samples.CNV detection for single samples usually yields many false detections, because of characteristics of genomic segments that lead to a higher or lower read count (and coverage). These biases can usually not be corrected for (except for the GC content and the mappability bias). Being aware of all these problems we have implemented the function singlecn.mops for cases in which only one sample is available.

Seq. Type	Ploidy	Study	Samples	Function
WGS	2n	cohort/non-tumor/GWAS	≥5	cn.mops
WGS	2n	tumor vs. normal	≥ 2	referencecn.mops
ES	2n	cohort/non-tumor/GWAS	≥5	exomecn.mops
ES	2n	tumor vs. normal	≥ 2	referencecn.mops
WGS	1n	cohort/non-tumor/GWAS	≥5	haplocn.mops
WGS	1n	tumor vs. normal	≥ 2	not implemented
ES	1n	cohort/non-tumor/GWAS	≥5	haplocn.mops
ES	1n	tumor vs. normal	>2	not implemented

Table 1: **Seq. Type** reports the sequencing technology that was used: whole genome sequencing (WGS) or targeted/exome sequencing (ES). **Ploidy** gives the usual ploidy of the samples. In case of a tumor vs. control study the control sample is meant. **Study**: The type of study to be analyzed for CNVs: either a cohort study, such the HapMap or the 1000Genomes Project, or studies including a number of non-tumor samples or studies with both healthy and diseased individuals, i.e. genome wide association studies (GWAS). **Samples** reports the minimum number of samples needed for the analysis. **Function** gives the function of the cn.mops package that is appropriate for this setting.

18 REFERENCES

12 How to cite this package

If you use this package for research that is published later, you are kindly asked to cite it as follows: (Klambauer *et al.*, 2012).

To obtain BibTFX entries of the reference, you can enter the following into your R session:

> toBibtex(citation("cn.mops"))

References

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- Chiang, D. Y., Getz, G., Jaffe, D. B., Zhao, X., Carter, S. L., Russ, C., Nusbaum, C., Meyerson, M., and Lander, E. S. (2008). High-resolution mapping of copy-number alterations with massively parallel sequencing. *Nat. Methods*, 6, 99–103.
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- Sathirapongsasuti, F. J., Lee, H., Horst, B. A., Brunner, G., Cochran, A. J., Binder, S., Quackenbush, J., and Nelson, S. F. (2011). Exome Sequencing-Based Copy-Number Variation and Loss of Heterozygosity Detection: ExomeCNV. *Bioinformatics*, 27(19), 2648–2654.