

Detection of de novo copy number alterations in case-parent trios using the R package **MinimumDistance**

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Abstract

In studies involving case-parent trios assayed on high-throughput genotyping arrays, a common goal is to identify regions of de novo copy number alterations. This package defines a statistic, referred to as the minimum distance, that can be useful for identifying de novo copy number alterations in the offspring. We smooth the minimum distance using the circular binary segmentation algorithm implemented in the Bioconductor package **DNAcopy**. Trio copy number states are inferred from the maximum a posteriori probability of the segmented data, incorporating information from the log R ratios and B allele frequencies. As both log R ratios and B allele frequencies can be estimated from Illumina and Affymetrix arrays, this package supports de novo copy number inference in both platforms.

1 Introduction

There are numerous R packages available from Bioconductor for smoothing copy number alterations. For example, the biocview **CopyNumberVariants** in the 2.9 release of Bioconductor lists 27 packages. For the analysis of germline diseases, hidden Markov models have emerged as a popular tool as inference regarding the latent copy number state incorporates information from both the estimates of copy number (or relative copy number) and the allelic frequencies, such as the B allele frequency [4] or genotype calls [1, 7, 5]. For the analysis of somatic cell diseases such as cancer, algorithms that segment the genome into regions of constant copy number (referred to here as segmentation algorithms) may be more preferable for the detection of copy number aberrations (CNA) as a mixture of cells with different copy numbers give rise to mosaic (non-integer) copy numbers. Examples include circular binary segmentation implemented in the R package **DNAcopy** and the **GLAD**, both of which were originally developed for array CGH platforms [3, 2, 6]. One disadvantage of segmentation algorithms is that inference regarding duplication and deletions is not directly available.

More recently, HMMs and segmentation algorithms have been developed for inferring common regions of copy number alterations in multiple samples. However, relatively few algorithms are available for inferring copy number alterations, especially de novo copy number alterations, in family-based study designs involving case-parent trios. Instead, a common strategy has been a two-step approach of identifying copy number alterations in the individual samples and then comparing the results across samples to infer whether an alteration observed in the offspring is de novo. A disadvantage of the two-step approach is that technical sources of variation, such as genomic waves that can effect parents and offspring differentially, often contribute to false positives. To our knowledge, the joint HMM implemented in the **PennCNV** software is the only software to date that provides direct inference regarding de novo alterations in case parent studies.

This package develops an alternative framework for inferring regions of de novo copy number alterations in case-parent trios. Like **PennCNV**, inference regarding de novo alterations integrates information from both the log R ratios and B allele frequencies. Differences in the two approaches are most apparent in non-HapMap experimental datasets in which technical and experimental sources of variation appear to contribute to a large number of false positives. Section 2 describes the data structures required for using this package. Section 3 outlines the steps for computing the minimum distance, segmenting the minimum distance, and computing maximum a posteriori probabilities to infer regions of de novo copy number alterations. We provide convenient tools for visualizing the low-level statistical summaries in the context of the segmented (*ranged* data) based on the **lattice** graphics. The workflow is illustrated using publicly available HapMap

trios assayed on the Illumina 610quad array. Where possible, we have adopted standard data structures for encapsulating the low-level data (**eSet** extensions) and the segmented data (**RangedData** extensions).

2 Required data structures

We begin by describing how to instantiate the data structures required for inferring changes in the de novo copy number. A key design consideration is that the 'unit' of our analysis is a trio, the sample size is the number of trios, and a common query will be to access the low level statistical summaries (log R ratios and B allele frequencies) for a set of markers for all members in a trio. For this particular use-case, it is much more convenient to store statistical summaries in a **feature x trios x family member** array rather than a **feature x samples** matrix. As the number of trios in a typical genome-wide association study may exceed 1000 (corresponding to several thousand samples), it may be impractical to store the low-level summaries in a single array. To make the size of the arrays more manageable, we organize the low-level summaries by chromosome in an object termed a **TrioSet**. We define a formal S4 class for a list of **TrioSet** objects as a **TrioSetList**. In addition to containing a **TrioSet** for each chromosome, the **TrioSetList** class has a slot for pedigree information defining the family relationships and a slot for the **sampleSheet** containing metadata on the experiment. We begin by constructing object of class **Pedigree** and **SampleSheet** that contain this information.

A constructor for the **SampleSheet** class is the **SampleSheet** function. The **SampleSheet** has 3 arguments (filename, id, and plate), but additional metadata on the samples can be supplied. For example, below we also include information in the well of the chemistry plate, the gender, and the SentrrixPosition for the Illumina array. The id can either be the same as the filename, or a more convenient sample identifier. For example, we use the NA identifiers instead of the lengthy sample names in the example below.

```
R> library(MinimumDistance)
R> library(human610quadv1bCr1mm)
R> path <- system.file("extdata", package="MinimumDistance")
R> load(file.path(path, "sample.sheet.rda"))
R> samplesheetExample <- SampleSheet(filename=sample.sheet$Sample.Name,
                                     id=paste("NA",substr(sample.sheet$Sample.Name, 6, 10),sep=""),
                                     plate=sample.sheet$Plate,
                                     well=sample.sheet$Well,
                                     gender=sample.sheet$Gender,
                                     SentrrixPosition=sample.sheet$SentrrixPosition)
```

An object of class *Pedigree* is constructed in the following codechunk:

```
R> load(file.path(path, "pedigreeInfo.rda"))
R> indId <- unlist(pedigreeInfo)
R> trioFactor <- matrix(seq_len(nrow(pedigreeInfo)), nrow(pedigreeInfo), 3, byrow=FALSE)
R> memberId <- substr(names(indId), 1, 1)
R> ## which row in the pedigree file
R> trio.index <- as.integer(trioFactor) ## note this gives the index in the pedigree
R> pedigreeIndex <- data.frame(individualId=indId,
                               memberId=memberId,
                               index.in.pedigree=trio.index,
                               stringsAsFactors=FALSE)
R> rownames(pedigreeIndex) <- NULL
R> pedigreeExample <- new("Pedigree", trios=pedigreeInfo,
                          trioIndex=pedigreeIndex)
R> allNames(pedigreeExample)

[1] "NA06993" "NA11881" "NA06985" "NA11882" "NA06991" "NA10859"

R> show(pedigreeExample)
```

```

trios:
      F      M      O
1 NA06993 NA06985 NA06991
2 NA11881 NA11882 NA10859

```

```

pedigreeIndex:
  individualId memberId index.in.pedigree
1      NA06993      F      1
2      NA11881      F      2
3      NA06985      M      1
4      NA11882      M      2
5      NA06991      O      1
6      NA10859      O      2

```

Next, we load matrices of the \log_2 R ratios and B allele frequencies extracted from the BeadStudio output. These matrices contain 25 markers for each chromosome as the intent is to illustrate the steps required for initializing a data structure of the `TrioSetList` class. (In order to illustrate some of the smoothing functions downstream, we will use the `trioSetListExample` data provided with the package. The `trioSetListExample` contains many more markers, but only for two autosomes). A *TrioSetList* object can be created using the helper function `TrioSetList`.

```

R> load(file.path(path, "logRratio.rda"))
R> load(file.path(path, "baf.rda"))
R> trioSetList <- TrioSetList(lrr=logRratio,
                             baf=baf,
                             pedigree=pedigreeExample,
                             sampleSheet=samplesheetExample,
                             cdfname="human610quadv1bCr1mm")

```

Note that the log R ratio and B allele frequency matrices have been transformed into arrays using the information stored in the `pedigree` object.

```

R> trioSetList

```

TrioSetList of length 22

```

R> dims(trioSetList)

```

	chr 1	chr 2	chr 3	chr 4	chr 5	chr 6	chr 7	chr 8	chr 9
Features	25	25	25	25	25	25	25	25	25
Trios	2	2	2	2	2	2	2	2	2
F, M, O	3	3	3	3	3	3	3	3	3

	chr 10	chr 11	chr 12	chr 13	chr 14	chr 15	chr 16	chr 17
Features	25	25	25	25	25	25	25	25
Trios	2	2	2	2	2	2	2	2
F, M, O	3	3	3	3	3	3	3	3

	chr 18	chr 19	chr 20	chr 21	chr 22
Features	25	25	25	25	25
Trios	2	2	2	2	2
F, M, O	3	3	3	3	3

```

R> sampleSheet(trioSetList)

```

	filename	id	plate	well	gender
1	CIDR_06993@0072958140	NA06993	WG1001552-DNA	C12	Male
4	CIDR_11881@0072958128	NA11881	WG1001557-DNA	E06	Male

```

2 CIDR_06985@0072958138 NA06985 WG1001574-DNA F12 Female
5 CIDR_11882@0072958127 NA11882 WG1001558-DNA G09 Female
3 CIDR_06991@1007845680 NA06991 WG1001289-DNA D10 Female
6 CIDR_10859@0072958146 NA10859 WG1001558-DNA E03 Female

```

```

  SentrrixPosition
1 4256206347_R01C02
4 4256206437_R01C01
2 4256206089_R02C01
5 4256206428_R01C02
3 4256193742_R01C02
6 4256193768_R01C01

```

```
R> pedigree(trioSetList)
```

```
trios:
```

```

      F      M      O
1 NA06993 NA06985 NA06991
2 NA11881 NA11882 NA10859

```

```
pedigreeIndex:
```

	individualId	memberId	index.in.pedigree
1	NA06993	F	1
2	NA11881	F	2
3	NA06985	M	1
4	NA11882	M	2
5	NA06991	O	1
6	NA10859	O	2

```
R> trios(trioSetList)
```

```

      F      M      O
1 NA06993 NA06985 NA06991
2 NA11881 NA11882 NA10859

```

```
R> trioSetList[1:5]
```

```
TrioSetList of length 5
```

```
R> trioSetList[[1]]
```

```

TrioSet (storageMode: lockedEnvironment)
assayData: 25 features, 2 samples
element names: BAF, logRRatio

```

As the offspring name uniquely identifies a trio, `sampleNames` and `offspringNames` are interchangeable.

```
R> ## Note
```

```
R> identical(sampleNames(trioSetList), offspringNames(trioSetList))
```

```
[1] TRUE
```

Feature annotation such as chromosome and position for each SNP is added by the function `featureDataFrom` defined in the `oligoClasses` package. Alternatively, users may supply their own feature annotation through the optional `featureData` argument to the R function `TrioSetList`. Platforms supported by the function `featureDataFrom` are listed below.

```
R> annotationPackages()[grep("Crlmm", annotationPackages())]
```

```
[1] "genomewidesnp6Crlmm" "genomewidesnp5Crlmm"
[3] "human370v1cCrlmm"    "human370quadv3cCrlmm"
[5] "human550v3bCrlmm"    "human650v3aCrlmm"
[7] "human610quadv1bCrlmm" "human660quadv1aCrlmm"
[9] "human1mduov3bCrlmm"  "humanomni1quadv1bCrlmm"
```

3 Detection of de novo copy number

In order to make the size of this package manageable, the `trioSetList` object created in the previous section contained only 25 markers for each of the 22 autosomes. While useful for illustrating the steps for creating the required data structures, it is not useful for detecting de novo copy number alterations. To illustrate the smoothing steps, we load an object that contains all the markers for 2 chromosomes (7 and 22) for 2 HapMap trios. A few summary statistics for this object are displayed below.

```
R> data(trioSetListExample)
R> show(trioSetList)

TrioSetList of length 2

R> dims(trioSetList)

      chr 7 chr 22
Features 34080  9284
Trios      2      2
F, M, O    3      3

R> sampleNames(trioSetList)

[1] "NA12878" "NA12864"

R> allNames(trioSetList)

[1] "NA12891" "NA12872" "NA12892" "NA12873" "NA12878" "NA12864"

R> annotation(trioSetList)

[1] "human610quadv1bCrlmm"
```

For a given trio, the signed minimum absolute difference of the offspring and parental \log_2 R ratios (\mathbf{r}) is defined as

$$\mathbf{d} \equiv (\mathbf{r}_O - \mathbf{r}_M) \times \mathbb{I}_{[|\mathbf{r}_O - \mathbf{r}_F| > |\mathbf{r}_O - \mathbf{r}_M|]} + (\mathbf{r}_O - \mathbf{r}_F) \times \mathbb{I}_{[|\mathbf{r}_O - \mathbf{r}_F| \leq |\mathbf{r}_O - \mathbf{r}_M|]}. \quad (1)$$

The above calculation is vectorized in R. The following codechunk calculates the minimum distance for each chromosome in the `trioSetList` object, returning a list of matrices for each chromosome. We store the minimum distances in the *TrioSet* objects using the replacement method `mindist`.

```
R> md <- calculateMindist(trioSetList)

|
|                                     | 0%
|
|=====| 50%
|
|=====| 100%
```

```

|
|
|
|=====| 50%
|
|=====| 100%

```

```
R> mindist(trioSetList) <- md
```

We calculate the median absolute deviation (MAD) of the minimum distance and the log R ratios. For the log R ratios, we calculate the MAD across markers and across independent offspring.

```

R> ## calculate the mad of the minimum distance
R> mads.md <- mad2(mindist(trioSetList), byrow=FALSE)
R> mad.mindist(trioSetList) <- mads.md
R> mads.lrr.sample <- mad2(lrr(trioSetList), byrow=FALSE)
R> mads.lrr.marker <- mad2(lrr(trioSetList), byrow=TRUE)
R> mad.sample(trioSetList) <- mads.lrr.sample
R> mad(trioSetList)

```

```

          F          M          O
NA12878 0.1291345 0.1125293 0.1083781
NA12864 0.1377335 0.2068227 0.1654582

```

```

R> mad.marker(trioSetList) <- mads.lrr.marker
R> fvarLabels(trioSetList[[1]])

```

```

[1] "chromosome"      "position"         "isSnp"
[4] "marker.mad"      "marker.mad.father" "marker.mad.mother"

```

The circular binary segmentation algorithm implemented in the R package DNAcopy is used to smooth the minimum distance estimates. The method `segment2` is a wrapper to the `CNA.smooth` and `segment` functions defined in DNAcopy. Arguments to the `segment` function can be passed through the `...` argument in `segment2`, allowing users to modify the segmentation from the default values in DNAcopy.

```

R> md.segs <- segment2(object=mindist(trioSetList),
                      pos=position(trioSetList),
                      chrom=chromosome(trioSetList), verbose=1)

```

The object returned by the `segment2` method is an object of class `RangedDataCBS`. Currently, I do not use `space` in `RangedData` objects.

```
R> md.segs
```

RangedDataCBS with 33 rows and 6 value columns across 1 space

	space	ranges	chrom	num.mark
	<factor>	<IRanges>	<integer>	<numeric>
1	1	[37124, 8197116]	7	1936
2	1	[8200167, 8200467]	7	2
3	1	[8200896, 49923859]	7	11315
4	1	[37124, 49923859]	7	13212
5	1	[50003183, 57941963]	7	1636
6	1	[50003183, 57941963]	7	1632
7	1	[61060840, 62120420]	7	101
8	1	[62128107, 72534486]	7	1829
9	1	[61060840, 61668154]	7	79

```

...      ...      ...      ...      ...
25      1 [14456412, 17074487] |      22      515
26      1 [17092563, 17118296] |      22      2
27      1 [17135059, 20780261] |      22      825
28      1 [20784580, 21554058] |      22      290
29      1 [21579040, 39642296] |      22      4621
30      1 [39719455, 44690948] |      22      1439
31      1 [39719455, 44690948] |      22      1432
32      1 [44801602, 49562479] |      22      1559
33      1 [44801602, 49562479] |      22      1552

```

```

      id start.index end.index seg.mean
<character> <integer> <integer> <numeric>
1      NA12878      1      1936      0.0003
2      NA12878     1937      1938     -0.6592
3      NA12878     1939     13254     -0.0028
4      NA12864      1     13254     -0.0159
5      NA12878     13255     14890     -0.0092
6      NA12864     13255     14890     -0.0274
7      NA12878     14891     14991      0.0784
8      NA12878     14992     16820     -0.0080
9      NA12864     14891     14969     -0.0421
...      ...      ...      ...
25     NA12864      1      516     -0.0199
26     NA12864     517      518     -0.7901
27     NA12864     519     1348     -0.0250
28     NA12864     1349     1640     -0.3484
29     NA12864     1641     6286     -0.0180
30     NA12878     6287     7725     -0.0029
31     NA12864     6287     7725     -0.0194
32     NA12878     7726     9284     -0.0026
33     NA12864     7726     9284     -0.0131

```

Accessors for objects of this class include

```
R> sampleNames(md.segs)
```

```

[1] "NA12878" "NA12878" "NA12878" "NA12864" "NA12878" "NA12864"
[7] "NA12878" "NA12878" "NA12864" "NA12864" "NA12864" "NA12878"
[13] "NA12864" "NA12878" "NA12864" "NA12878" "NA12864" "NA12878"
[19] "NA12878" "NA12878" "NA12878" "NA12878" "NA12878" "NA12878"
[25] "NA12864" "NA12864" "NA12864" "NA12864" "NA12864" "NA12878"
[31] "NA12864" "NA12878" "NA12864"

```

```
R> coverage2(md.segs)
```

```

[1] 1936      2 11315 13212 1636 1632 101 1829      79      64
[11] 1782 3746 3730 12176 12143 1334 1330 445      2 1181
[21] 10 604      70 3974 515      2 825 290 4621 1439
[31] 1432 1559 1552

```

```
R> chromosome(md.segs)
```

```

[1] 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 22 22 22 22
[22] 22 22 22 22 22 22 22 22 22 22 22 22 22

```

```
R> mean(md.segs)
```

```
[1] 0.0003 -0.6592 -0.0028 -0.0159 -0.0092 -0.0274 0.0784 -0.0080
[9] -0.0421 0.2125 -0.0190 -0.0028 -0.0215 -0.0059 -0.0190 -0.0009
[17] -0.0215 0.0063 0.5924 0.0017 -0.3244 -0.0080 -0.2317 -0.0064
[25] -0.0199 -0.7901 -0.0250 -0.3484 -0.0180 -0.0029 -0.0194 -0.0026
[33] -0.0131
```

Having partitioned the genome by the segmentation of the minimum distance, we can call the trio copy number state for each genomic interval. Before we do so, consider that minimum distance estimates near zero are consistent with normal copy number in all members of the trio and a copy number alteration in the offspring inherited from either parent. Or that minimum distance estimates near -1 are consistent with a denovo hemizygous deletion in which both parents are normal and a denovo homozygous deletion in the offspring when both parents are hemizygous. While we can distinguish between these possibilities by examining the distribution of log R ratios and B allele frequencies in the family members, an automatic means to correctly classify the trio copy number state becomes more difficult if a single segment contains a mixture of these states. We therefore edit minimum distance ranges that are nonzero, where nonzero is defined by the number of MADs the mean is from zero. The editing is achieved by segmenting the log R ratios of the offspring. If a breakpoint is identified within a nonzero minimum distance segment, the breakpoints of the minimum distance segment are edited. The following diagram using vertical dashes (|) to denote breakpoints illustrates the basic principle.

```
--|-----|--      ## minimum distance segment (before editing)
---|-----|-----  ## segmenation of log R ratios for offspring

->

--|-|-----|---|--  ## after editing
```

In the following codechunk, we segment the log R ratios for all chromosomes in the `trioSetList` object for the father, mother, and offspring. While only the segmentation of the offspring log R ratios are required to edit the minimum distance breakpoints, the segmentation of the parental log R ratios are useful for the visualization methods discussed in Section 3.1.

```
R> lrr.segs <- segment2(object=lrr(trioSetList),
                        pos=position(trioSetList),
                        chrom=chromosome(trioSetList),
                        id=trios(trioSetList), ## id is required
                        verbose=TRUE)
```

Note that the object `lrr.segs` is also a `RangedDataCBS` object. Minimum distance segments with a mean absolute value greater than 0.9 MADs from zero are edited using the `narrow`:

```
R> md.segs2 <- narrow(md.segs, lrr.segs, 0.9, mad.minimumdistance=mads.md)
```

Note that the number of segments after editing is typically greater than the original number of segments:

```
R> nrow(md.segs2) > nrow(md.segs)
```

```
[1] FALSE
```

Finally, we use the ranged data in the `md.segs2` object to call the trio copy number state for each interval. Conditional on the underlying trio copy number state, we calculate the probability of the observed log R ratios and the probability of the observed B allele frequencies. Details regarding the mixture model used to estimate the emission probabilities are discussed elsewhere. The copy number state with the maximum posterior probability is assigned to each genomic range. We assign to each range both the maximum posterior probability and the posterior probability of the normal range. The ratio of posterior probabilities provides a useful statistic to rank the genomic ranges by the evidence of de novo copy number alteration. Consecutive

genomic ranges in a sample that are assigned the same copy number state are collapsed into a single range. As a consequence, the number of ranges in the object returned by `computeBayesFactor` is typically smaller than the number of ranges passed to the `ranges` argument. We continue the previous example to illustrate how ranges are subject to pruning following the posterior classification of the copy number states:

```
--|-|-----|---|--      ## after editing
--|-----|---|--      ## result 1: all segments have the same trio state call
--|-----|---|--      ## result 2: the first two segments have the same call
```

Note that 'result 1' recovers the original segmentation of the minimum distance. Pragmatically, the above steps are implemented in the function `computeBayesFactor`:

```
R> map.segs <- computeBayesFactor(object=trioSetList, ranges=md.segs2)
R> map.segs <- map.segs[order(sampleNames(map.segs), chromosome(map.segs), start(map.segs)), ]
R> map.segs
```

```
RangedDataCBS with 16 rows and 11 value columns across 1 space
```

	space	ranges	id	chrom	num.mark
	<factor>	<IRanges>	<character>	<integer>	<integer>
1	1	[37124, 61668154]	NA12864	7	14969
2	1	[61673005, 62336389]	NA12864	7	64
3	1	[62342453, 158812247]	NA12864	7	19047
4	1	[14456412, 20780261]	NA12864	22	1348
5	1	[20784580, 21554058]	NA12864	22	292
6	1	[21579040, 49562479]	NA12864	22	7644
7	1	[37124, 8197116]	NA12878	7	1936
8	1	[8200167, 8200467]	NA12878	7	2
9	1	[8200896, 158812247]	NA12878	7	32142
10	1	[14456412, 16871770]	NA12878	22	445
11	1	[16872032, 16872749]	NA12878	22	2
12	1	[16873980, 21419339]	NA12878	22	1181
13	1	[21441199, 21538519]	NA12878	22	10
14	1	[21540389, 23970628]	NA12878	22	604
15	1	[23980406, 24244593]	NA12878	22	70
16	1	[24254444, 49562479]	NA12878	22	6972

	seg.mean	start.index	end.index	mindist.mad	lik.state
	<numeric>	<numeric>	<numeric>	<numeric>	<numeric>
1	-0.017295137	1	14969	0.11682888	4.657932e+04
2	0.212500000	14970	15033	0.11682888	1.001422e+02
3	-0.019666772	15034	34080	0.11682888	6.021705e+04
4	-0.024182938	1	1348	0.11682888	4.531105e+03
5	-0.348400000	1349	1640	0.11682888	7.875919e+02
6	-0.017264194	1641	9284	0.11682888	2.689665e+04
7	0.000300000	1	1936	0.08880774	7.132457e+03
8	-0.659200000	1937	1938	0.08880774	-9.202714e-01
9	-0.004262367	1939	34080	0.08880774	1.163936e+05
10	0.006300000	1	445	0.08880774	1.571798e+03
11	0.592400000	446	447	0.08880774	-4.211252e+00
12	0.001700000	448	1628	0.08880774	4.166802e+03
13	-0.324400000	1629	1638	0.08880774	1.974743e+01
14	-0.008000000	1639	2242	0.08880774	2.178886e+03
15	-0.231700000	2243	2312	0.08880774	2.068662e+02
16	-0.004827897	2313	9284	0.08880774	2.660287e+04

	lik.norm	argmax	state
	<numeric>	<integer>	<character>

copy number	symbol for copy number state
0	1
1	2
2	3
3	4
4	5

Table 1: We use the notation adopted by PennCNV for representing the copy number states. The trio copy number state is indicated by the triplet xyz where x is the symbol for the copy number state of the father, y is the symbol for the copy number state of the mother, and z is the symbol for the offspring copy number state. For example, the trio copy number state '332' indicates a de novo hemizygous deletion in the offspring as both parents have *normal* copy number (copy number 2) and the offspring has a single copy.

1	41730.970277	61	333
2	45.951226	85	334
3	5691.972685	61	333
4	1670.035038	61	333
5	470.318982	37	332
6	16416.834538	61	333
7	7132.456873	61	333
8	-1.013983	37	332
9	40615.749758	61	333
10	1571.798135	61	333
11	-11.899091	55	223
12	4166.802470	61	333
13	16.348665	37	332
14	2178.885858	61	333
15	119.096697	37	332
16	15420.296813	61	333

The integer code for the copy number states are provided in the Table 1 and can be accessed by the function `state`. For example, here we tabulate the frequency of the trio copy number states by chromosome from the `map.segs` object.

```
R> table(state(map.segs), chromosome(map.segs))
```

```

      7 22
223 0  1
332 1  3
333 4  6
334 1  0

```

3.1 Visualizations

The R package `VanillaICE` contains methods for visualizing *RangedData* objects that build on the infrastructure in the `lattice` package. In the following codechunk, we construct a trellis object for the log R ratios, minimum distance, and B allele frequencies.

```

R> trioSet <- stack(trioSetList)
R> rd.denovoDel <- map.segs[state(map.segs) == 332, ]
R> library("VanillaICE")
R> ##md.segs$seg.mean <- -1*md.segs$seg.mean
R> lrrfig <- xyplot(lrr ~ x | id,
                   data=trioSet,

```

```

range=rd.denovoDel[1, ],
lrr.segs=lrr.segs,
md.segs=md.segs,
frame=500e3,
panel=xypanelMD,
xlab="physical position (Mb)",
cex=0.2,
pch=21,
border="orange",
scales=list(cex=0.5),
par.strip.text=list(lines=0.8, cex=0.6),
col.np="royalblue", fill.np="grey",
index.cond=list(4:1),
return.data.frame=FALSE,
cex.state=0.5)
R> baffig <- xyplot(baf ~ x | id,
data=trioSet,
range=rd.denovoDel[1, ],
xlab="",
lrr.segs=lrr.segs,
md.segs=md.segs,
frame=500e3,
panel=VanillaICE::xypanel,
cex=0.2,
pch=21,
border="orange",
scales=list(cex=0.6),
par.strip.text=list(lines=0.8, cex=0.6),
col.np="royalblue", fill.np="grey",
index.cond=list(3:1), cex.state=0.5)

```

The final graphic is displayed in Figure 1.

```
R> print(VanillaICE:::arrangeSideBySide(lrrfig, baffig))
```

4 Session information

```
R> toLatex(sessionInfo())
```

- R Under development (unstable) (2011-11-03 r57560), x86_64-unknown-linux-gnu
- Locale: LC_CTYPE=en_US.iso885915, LC_NUMERIC=C, LC_TIME=en_US.iso885915, LC_COLLATE=en_US.iso885915, LC_MONETARY=en_US.iso885915, LC_MESSAGES=en_US.iso885915, LC_PAPER=C, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.iso885915, LC_IDENTIFICATION=C
- Base packages: base, datasets, graphics, grDevices, grid, methods, stats, utils
- Other packages: Biobase 2.15.0, BiocInstaller 1.3.1, DNACopy 1.29.0, human610quadv1bCrlmm 1.0.2, IRanges 1.13.1, MinimumDistance 0.1.47, oligoClasses 1.17.1, VanillaICE 1.17.2
- Loaded via a namespace (and not attached): affyio 1.23.0, Biostings 2.23.0, bit 1.1-7, ff 2.2-3, lattice 0.20-0, msm 1.1, mvtnorm 0.9-9991, SNPchip 1.19.0, splines 2.15.0, survival 2.36-10, tools 2.15.0, zlibbioc 1.1.0

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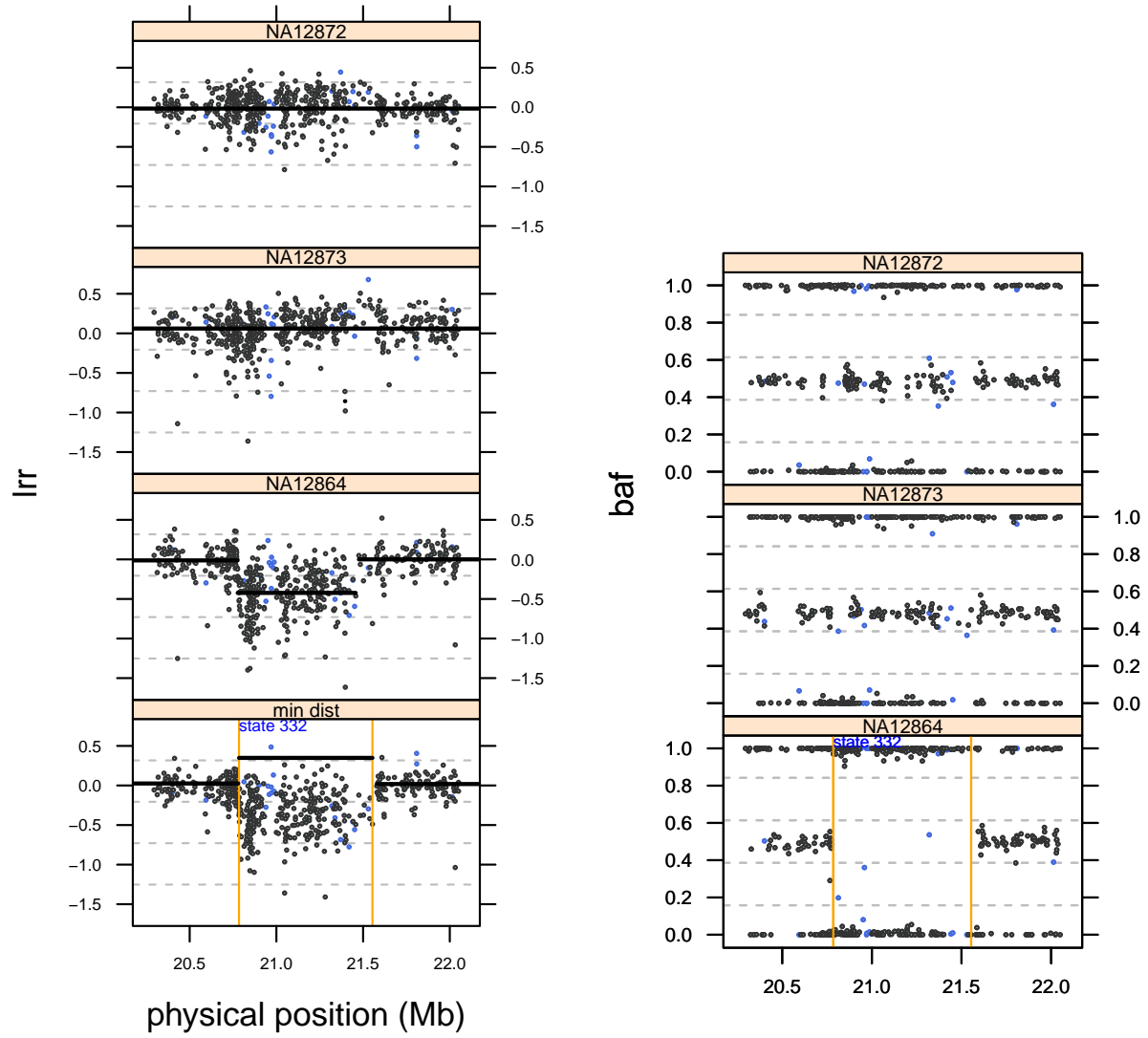


Figure 1: The left panels display the log R ratios for the father (top left), mother, and offspring, as well as the minimum distance (bottom left). The right panels display the B allele frequencies for the father, mother, and offspring. The orange vertical lines indicate the start and stop positions of an inferred de novo hemizygous deletion in the offspring (state '332').