# CNVrd2: A package for measuring gene copy number, identifying SNPs tagging copy number variants, and detecting copy number polymorphic genomic regions

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## 1 Introduction

The CNVrd2 package <sup>1</sup> (Nguyen et~al., 2013) utilizes next-generation sequencing (NGS) data to measure human-gene copy number (CN) and identify single-nucleotide polymorphisms (SNPs), and insertions and deletions (INDELs) that are in linkage disequilibrium with a gene of interest. Typically, the data being used are low- or medium-coverage whole genome sequence (WGS) data from multiple individuals in a population. Such data comprise collections of sequence reads that have been aligned (or "mapped") to an appropriate reference genome. Changes in read depth (i.e., the number of reads aligned to a specific region of the genome) can indicate changes in DNA copy number in this region (i.e., deletions or duplications of specific portions of DNA). If this region encompasses a gene, then changes in copy number may also be reflected by changes in gene activity - such changes have been shown to be associated with altered risk of disease in human populations, and altered trait distributions in agricultural settings.

To measure gene CN, CNVrd2 firstly divides a region (usually at least 1Mb) flanking a gene of interest into constant-sized windows, and counts reads mapped in these windows. Next, these read-count windows are transformed and standardized. After that, the DNAcopy package (Venkatraman and Olshen, 2007) is used to join the per-window standardized counts into regions (or "segments") of similar values. The package then refines the segmentation step and outputs segmentation results, namely segmentation scores (SS), for each sample. A function in the CNVrd2 package is then used to group SSs into copy-number groups.

To calculate linkage disquilibrium (LD) between gene CNVs and SNPs/INDELs nearby, SNPs/INDELs are coded into numeric values (0, 1, 2) and Fisher's Exact Test is used to assess associations between SNPs/INDELs and copy number. CNVrd2 is designed to identify SNPs/INDELs that can be used as a surrogate marker for CNVs, therefore multiple samples are needed to obtain reliable results. The package also uses distribution quantiles to identify highly polymorphic regions of the genome (within a collection of samples) and can identify regions with variable polymorphism between populations. The BAM format (Li  $et^*al.$ , 2009) for aligned-NGS data and VCF format (Danecek  $et^*al.$ , 2011) for structural variant information are used as the main forms of input for the package.

 $<sup>^1</sup>$ CNVrd2 is an improved version of the pipeline CNVrd used to identify tagSNPs of FCGR3A/B CNV

## 2 Getting started

First, we load the package in our R session. Note that the *rjags* package (Plummer and Plummer, 2013) requires the associated JAGS application to be installed.

```
library("CNVrd2")
```

#### Working with BAM and VCF files.

The following section describes the workflow of the CNVrd2 package in reading BAM and VCF files into R. The 58 MXL-sample BAM files (chr1:161100000-162100000) were downloaded from the 1000 Genomes Project to measure copy number counts of FCGR3B gene (chr1:161592986-161601753). Users can download the file MXLexample.zip on

http://code.google.com/p/cnvrdfortagsnps/downloads/list and unzip it into a directory.

Alternatively, to run the example without downloading the associated BAM files, users can skip to section 2.1.3 to load a pre-processed verison of the same data.

#### 2.1 Measuring FCGR3B CN

#### 2.1.1 CNVrd2 object

We need to make an object of class CNVrd2 to define a region we want to investigate (regions sized > 1Mb tend to work well - multiple genes can be included by specifying the start and end positions of each). Here, we choose 1000bp-constant windows. We also need to supply a directory that consists of BAM files including only mapped reads. Users who have **not downloaded the BAM files**, should skip to section 2.1.3

```
objectCNVrd2 <- new("CNVrd2", windows = 1000, chr = "chr1", st = 161100001,
en = 162100000, dirBamFile = "BamMXL", genes = c(161592986, 161601753),
geneNames = "3B")
```

#### 2.1.2 Count reads in windows

Use the function countReadInWindow to read the BAM files into R and count the number of reads in each of the windows.

```
readCountMatrix <- countReadInWindow(Object = objectCNVrd2, correctGC = TRUE)</pre>
```

If GC-content correction is selected (correctGC=TRUE) then a reference genome must be supplied. The default reference genome is the human reference genome (UCSC version hg19). A full list of reference genomes available through Bioconductor can be obtained from:

```
http://www.bioconductor.org/packages/release/bioc/html/BSgenome.html
```

#### 2.1.3 Segmentation

Use the function segmentSamples to segment and obtain segmentation scores for the FCGR3B gene (Figure 1):

```
## Obtain segmentation scores
resultSegment <- segmentSamples(Object = objectCNVrd2, stdCntMatrix = readCountMatrix)</pre>
```

Instead of reading BAM files directly, we can use a matrix of read counts for the function segmentSamples. Here, we obtain a read-count matrix from data in the CNVrd2 package.

```
## Load data into R
data(fcgr3bMXL)
## Reload readCountMatrix
readCountMatrix <- resultSegment$stdCntMatrix
## Take a quick look the data
readCountMatrix[1:2, 1:2]
## Make a CNVrd2 object
objectCNVrd2 <- new("CNVrd2", windows = 1000, chr = "chr1", st = 161100001,
        en = 162100000, dirBamFile = "BamMXL", genes = c(161592986, 161601753),
        geneNames = "3B")
## Obtain segmentation scores
resultSegment <- segmentSamples(Object = objectCNVrd2, stdCntMatrix = readCountMatrix)
## View these segmentation results
sS <- resultSegment$segmentationScores
hist(sS[, 1], 100, xlab = "Segmentation score", main = "")</pre>
```

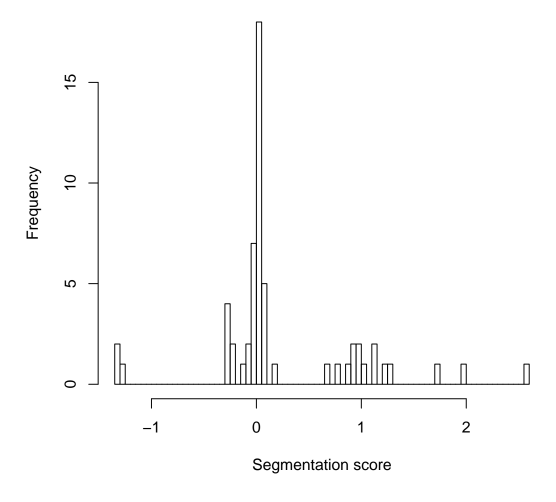


Figure 1: FCGR3B segmentation score.

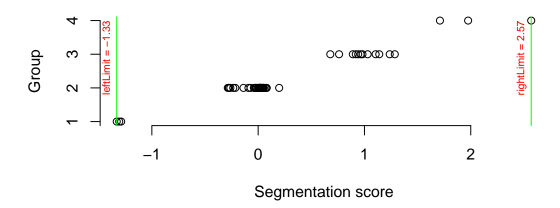
#### 2.1.4 Obtain copy-number count

The data in Figure 1 suggest four distinct groups of segmentation scores, likely related to four different copy number genotypes. The function  $group\,CNVs$  uses a normal mixture model to cluster SSs into groups. Unequal variances are assumed by default (EV = FALSE), however, if there are relatively few SS values in one group then we can set EV = TRUE (see the  $group\,CNVs$  manual page for additional details).

Clustering results are shown in Figure 2, and the group assignments for the samples are contained in the *allGroups* object. For example, the NA19648 sample is assigned to the second group because the probability associated with membership of this group is higher than that of the other groups (nearly 1).

```
copynumberGroups$allGroups[1:3, ]
##
                             Name Classification
                                                    Group1
                                                              Group2
## NA19648.MXL.bam NA19648.MXL.bam
                                               2 8.962e-19 1.000e+00
## NA19649.MXL.bam NA19649.MXL.bam
                                               2 1.985e-17 1.000e+00
## NA19651.MXL.bam NA19651.MXL.bam
                                               3 3.918e-62 8.216e-14
                     Group3
                               Group4
                                        score
## NA19648.MXL.bam 9.781e-12 5.203e-47 -0.0195
## NA19649.MXL.bam 8.309e-13 3.171e-49 -0.0710
## NA19651.MXL.bam 1.000e+00 2.959e-10 1.1393
```

If we would like to force outliers into the lowest or highest CN genotype groups (e.g., dividing



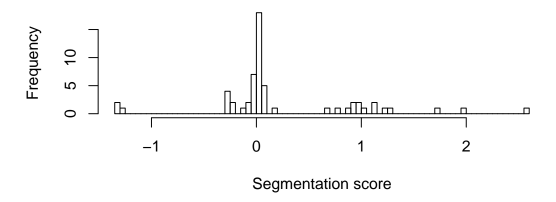
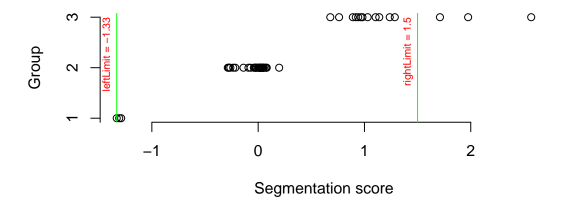


Figure 2: FCGR3B CN groups.

the data into three groups: deletions, normal CN, duplications) then we can use options rightLimit (Figure 3) or leftLimit or both.

#### 2.1.5 Plots

The function *plotCNVrd2* can plot multiple samples. Trace plots of some of the samples exhibiting duplications at the FCGR locus are shown in Figure 4. Here, based on information from the literature, we assume that a copy number of two is the most common CN genotype.



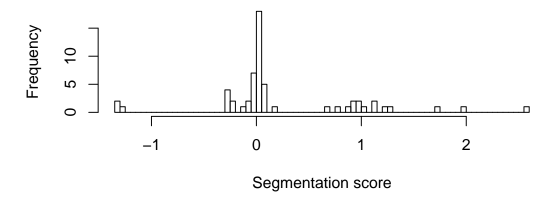


Figure 3: FCGR3B CN groups (right Limit =1.5).

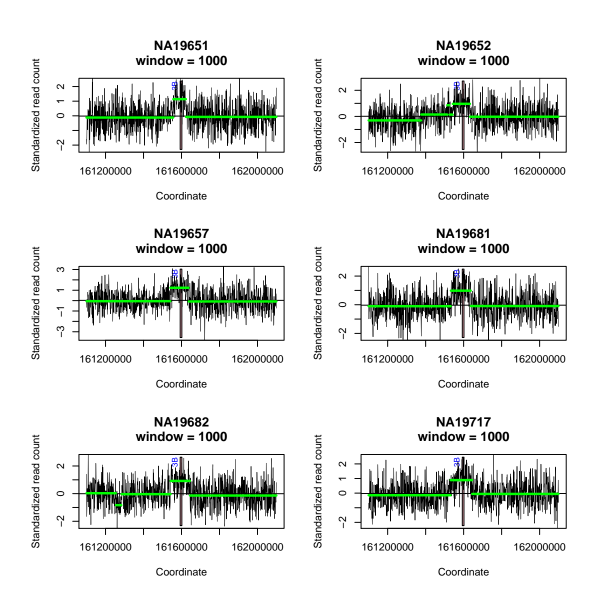


Figure 4: MXL duplicated samples.

## 2.2 Identifying tag SNPs/INDELs for FCGR3B CNVs

The function calculateLDSNP and CNV is used to calculate LD between CNVs and SNPs/INDELs. This function will read a VCF file into R and transform phased/unphased values (00, 01, 10, 11) into numeric values (0, 1, 2 or 0, 1). For a large VCF file (e.g., >= 1Mb), we generally use the option nChunkForVcf=50 to break the file into 50 chunks for reading into R.

```
## Obtain VCF-file information in CNVrd2 package
vcfFile <- system.file(package = "CNVrd2", "extdata", "chr1.161600000.161611000.vcf.gz")</pre>
## Make a data frame named sampleCNV including samples, CNs, population
sampleCNV <- data.frame(copynumberGroups$allGroups[, c(1, 2)], rep("MXL",</pre>
   dim(copynumberGroups$allGroups)[1]))
rownames(sampleCNV) <- substr(sampleCNV[, 1], 1, 7)</pre>
sampleCNV[, 1] <- rownames(sampleCNV)</pre>
## The first column must be the sample names and some samples should be in
## the vcf file
tagSNPandINDELofMXL <- calculateLDSNPandCNV(sampleCNV = sampleCNV, vcfFile = vcfFile,</pre>
    cnvColumn = 2, population = "MXL", popColumn = 3, nChunkForVcf = 5, chr = "1",
    st = 161600001, en = 161611000, codeSNP = "Three", codeCNV = "ThreeGroup")
## Reading the VCF file 1:161600001-161611000 with 5 blocks each
## VCF file: 161600001 to 161602751
## VCF file: 161602752 to 161605500
## VCF file: 161605501 to 161608250
## VCF file: 161608251 to 161611000
## Calculating p and r2 values for MXL population.
```

```
head(tagSNPandINDELofMXL)
               1CN_{n=3} 2CN_{n=36} 3CN_{n=15} p.values
## rs117435514
                                        86.67 2.1e-09 0.81
                   0.00
                              2.78
## rs185696163
                   0.00
                              2.78
                                        86.67 2.1e-09 0.81
## rs34015117
                                        80.00 2.3e-08 0.77
                   0.00
                              2.78
## rs76736176
                                        66.67 1.5e-06 0.68
                   0.00
                              2.78
## esv2661911
                                         0.00 2.1e-03 -0.39
                   66.67
                              0.00
## rs72704050
                                        93.33 4.6e-03 0.34
                   33.33
                             52.78
```

From the results of the LD analysis, rs117435514 is the best tagSNP for duplications: 0%, 2.78% and 86.7% of deleted, normal and duplicated samples have this SNP (adjusted p-value = 7.1e-08,  $r^2 = 0.66$ ).

## 3 Working with complex loci

CNVrd2 can also be used to measure multiallelic copy-number polymorphisms. For loci having high CN, users should use the function segmentSamplesUsingPopInformation to adjust the segmentation process across populations. An xample of a gene exhibiting this type of complex CN polymorphism is CCL3L1. Below we use the package to measure CCL3L1 CN and identify tag SNPs/INDELs for CCL3L1 CNVs.

The data set used here includes 1,917 samples of five large populations European, East Asian, West African, South Asian ancestry and Americas with a total of 26 small populations as in the table below:

| Large Pop    | Small Pop | Sample size |
|--------------|-----------|-------------|
| Americas     | ACB       | 74          |
| Americas     | ASW       | 50          |
| Americas     | CLM       | 65          |
| Americas     | MXL       | 59          |
| Americas     | PEL       | 60          |
| Americas     | PUR       | 74          |
| East Asian   | CDX       | 88          |
| East Asian   | СНВ       | 83          |
| East Asian   | CHS       | 104         |
| East Asian   | JPT       | 82          |
| East Asian   | KHV       | 78          |
| European     | CEU       | 96          |
| European     | FIN       | 78          |
| European     | GBR       | 77          |
| European     | IBS       | 77          |
| European     | TSI       | 100         |
| South Asian  | BEB       | 50          |
| South Asian  | GIH       | 81          |
| South Asian  | ITU       | 39          |
| South Asian  | PJL       | 37          |
| South Asian  | STU       | 49          |
| West African | ESN       | 64          |
| West African | GWD       | 105         |
| West African | LWK       | 90          |
| West African | MSL       | 68          |
| West African | YRI       | 89          |

# 3.1 Measuring CCL3L1 CN

The *ccl3l1data* data includes 1917 samples downloaded from the 1000 Genomes Project in October 2012 and March 2013, their corresponding populations, segmentation scores and CNs <sup>2</sup>. The segmentation scores were obtained by using the

<sup>&</sup>lt;sup>2</sup>Manuscript in preparation

function segmentSamplesUsingPopInformation for a 1Mb region (chr17:33670000-34670000) with 500bp-constant windows.

```
## Load data into R:

data(ccl311data)

head(ccl311data)

## Name Pop SS CN

## 1 HG00096 GBR -0.6933 1

## 2 HG00100 GBR -0.2308 2

## 3 HG00103 GBR -0.3511 2

## 4 HG00106 GBR -0.8012 1

## 5 HG00108 GBR -0.4120 2

## 6 HG00111 GBR -0.2880 2

hist(ccl311data$SS, 100)
```

As can be seen in Figure 5, the data is multimodal and there are not clear clusters on the right. Therefore, we can use a single population which has clear clusters to obtain prior information for the clustering process into CN groups. Here, we used the large European-ancestry population to obtain prior information.

```
xyEuro <- ccl3l1data[grep("CEU|TSI|IBS|GBR|FIN", ccl3l1data[, 2]), ]
yEuro <- xyEuro[, 3]
names(yEuro) <- rownames(xyEuro)
hist(yEuro, 100, xlab = "", main = "")</pre>
```

As can be seen from Figure 6, the European-ancestry data exhibit relatively clear clusters, allowing us to classify the samples into different CN groups.

**Note:** if we use the option autoDetermineGroup = TRUE in the function groupC-NVs then the Bayesian information criterion (BIC) will be used to choose a suitable number of components (See Schwarz (1978)).

```
## Clustering European segmentation scores into group: 5 groups were
## chosen
```

# Histogram of ccl3l1data\$SS

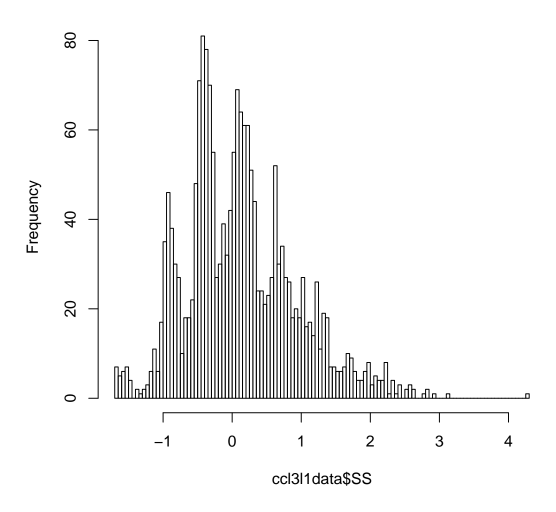
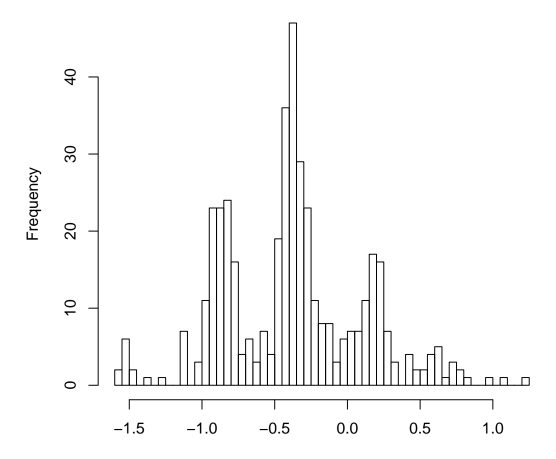


Figure 5: CCL3L1 segmentation score.



 $\label{eq:Figure 6: European-ancestry segmentation score.}$ 

Next, we use these results to infer *CCL3L1* CN in all populations. The following code collects information about the means, standard deviations and proportions of the mixture components from the European population.

```
# Means
lambda0 <- as.numeric(europeanCCL3L1Groups$m)
# SD
sdEM <- as.numeric(europeanCCL3L1Groups$sigma)
# Proportions
pEM <- as.numeric(europeanCCL3L1Groups$p)</pre>
```

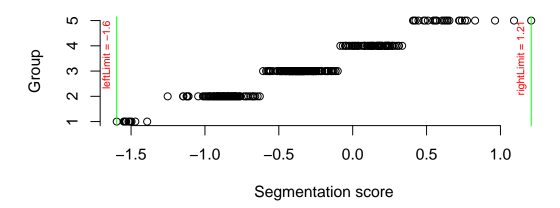
#### Take a look these results:

```
lambda0
## [1] -1.5140 -0.8719 -0.3596  0.1442  0.6111

sdEM
## [1] 0.04986  0.11320  0.10089  0.11547  0.23072

pEM
## [1] 0.02569  0.28138  0.44650  0.17878  0.06764

### Calculate the distances between groups
for (ii in 2:5) {
```



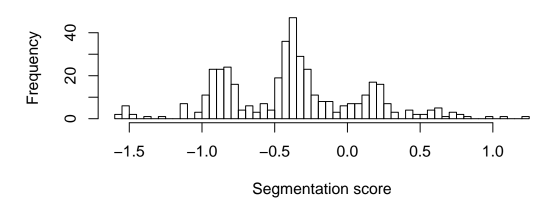


Figure 7: Clustering results of European-ancestry sample sets.

```
print(lambda0[ii] - lambda0[ii - 1])

## [1] 0.6422

## [1] 0.5123

## [1] 0.5038

## [1] 0.4668

### All segmentation scores

ccl311X <- ccl311data$SS

names(ccl311X) <- as.character(ccl311data$Name)

range(ccl311X)

## [1] -1.675 4.258</pre>
```

The information above is then used by the function groupBayesianCNVs to cluster the segmentation scores for the combined set of European-ancestry cohorts into different CN groups. There is a high value in the full SS set (Figure 5), which we eliminate in the following steps by setting rightLimit=4 so that it is automatically allocated into the highest CN group. Using the other values, combined with locus-specific information from the literature, we set the number of groups to be 10. All prior information was used in our work to obtain CN for CCL3L1 gene (the four group).

```
## Set prior information: prior for the sd of the means of groups: 5 was
## set for the third group = 2 CN

sd <- c(1, 1, 5, 1, 1)

ccl311X <- sort(ccl311X)

### Data

xData <- ccl311X

### Number of groups

nGroups <- 10

### prior for means of groups

lambda0 <- lambda0

### Prior for mixing proportions</pre>
```

```
alpha0 <- c(3, 29, 44, 18, 7, 5, rep(2, nGroups - length(pEM) - 1))

## Prior for the distances between groups

distanceBetweenGroups <- 0.485

sdEM <- sdEM
```

The final  $(5^{th})$  group of the results of the European-ancestry sample sets has a large standard deviation as a result of the scattering of values on the right (Figure 6), therefore, we can set this value to equal the standard deviation of the fourth group to avoid an overly wide mixture component.

```
## Adjust standard deviation for the fifth group
sdEM[5] <- sdEM[4]</pre>
```

Run the *groupBayesianCNVs* function to obtain CN groups.

```
set.seed(123)
groupCCL3L1allPops <- groupBayesianCNVs(xData = xData, nGroups = nGroups,
    lambda0 = lambda0, sd0 = sdEM, alpha0 = alpha0, distanceBetweenGroups = distanceBetweenGroups,
    sd0ftau = sd, rightLimit = 4)</pre>
```

These results would be similar to the results of the fourth column in the data of the package.

## 3.2 Identifying tag-SNPs for CCL3L1 CNVs

We can obtain tag-SNPs/INDELs for multiple populations simultaneously. Below, we reuse the CCL3L1 data to obtain tag-SNPs for some populations.

```
rownames(ccl3l1data) <- ccl3l1data[, 1]
```

Load VCF file into R and choose populations which we would like to find tagSNPs/IN-DELs.

```
## Obtain vcf-file information in CNVrd2
vcfFileCCL3L1 <- system.file(package = "CNVrd2", "extdata", "chr17.34800000.34830000.vcf.gz")
## Set populations we would like to identify tagSNPs
allPops <- c("TSI", "CEU", "GBR", "FIN", "IBS")</pre>
```

Take a quick look some significant results (multiple populations: the return value of calculateLDSNPandCNV is a list of populations).

```
lapply(tagSNPandINDELofCCL3L1, head)
## [[1]]
          OCN_(n=5) 1CN_(n=32) 2CN_(n=41) 3CN_(n=16) 4CN_(n=4)
             80 87.50 12.20 18.75 0
                                    6.25
## rs113877493 60 71.88 2.44
## rs11316723 80 84.38 12.20
## rs8072769 80 78.12 12.20
                                     18.75
                                     12.50
             80 87.50 17.07 18.75
## rs9911791
## rs113435750
            80 78.12 12.20 12.50
## p.values r p.valuesAdjusted r2 POP
## rs8064426 3.2e-12 -0.64 5.8e-10 0.40 TSI
## rs113877493 6.7e-12 -0.63
                           6.2e-10 0.39 TSI
## rs11316723 3.1e-11 -0.62
                           1.9e-08 0.37 TSI
## rs8072769 5.3e-10 -0.61
                          1.1e-08 0.34 TSI
3.8e-08 0.33 TSI
## rs9911791 2.5e-10 -0.58
## rs113435750 1.4e-09 -0.57
## [[2]]
          OCN_(n=2) 1CN_(n=20) 2CN_(n=39) 3CN_(n=14) 4CN_(n=3)
## rs8072769 0 75 2.56 0.00 0
## rs113435750 0
                      75 5.13 0.00
## rs138153523 0 75 5.13 0.00
## rs11316723 0 80 5.13 7.14
```

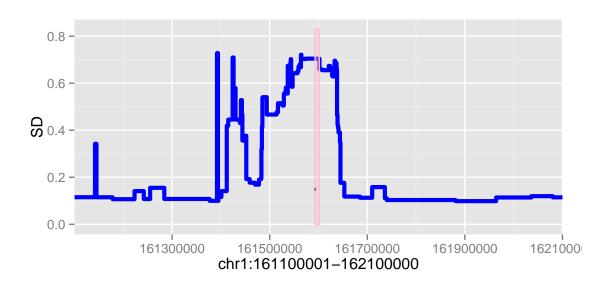
```
## rs8064426 0 80 5.13 7.14 0
## rs9911791 0 85 10.26 7.14 0
## 5CN_{n=1} p.values r p.valuesAdjusted r2 POP
## rs8072769 0 7.6e-10 -0.61 1.4e-07 0.37 CEU
               0 6.4e-09 -0.60
                                2.4e-07 0.35 CEU
              0 6.4e-09 -0.60
## rs138153523
              0 3.7e-09 -0.57 2.3e-07 0.33 CEU
## rs11316723
             ## rs8064426
               0 8.8e-09 -0.58
## [[3]]
## 1CN_(n=18) 2CN_(n=37) 3CN_(n=11) 4CN_(n=4) p.values
## rs11316723 72.22 2.70 0
                                      0 1.2e-08 -0.65
             72.22
                      2.70
## rs8064426 72.22 2.70
## rs9911791 72.22 8.11
                             0 0 5.6e-07 -0.62
## rs113877493 50.00 0.00 0 0 3.8e-06 -0.54
## rs8072769 50.00 2.70 0
## rs113435750 50.00 2.70 0
                                     0 5.1e-05 -0.52
                                       0 5.1e-05 -0.52
## p.valuesAdjusted r2 POP
## rs11316723 1.1e-06 0.43 GBR
## rs8064426
              1.1e-06 0.43 GBR
## rs9911791
                3.4e-05 0.38 GBR
## rs113877493
                1.8e-04 0.29 GBR
## rs8072769
               1.6e-03 0.27 GBR
## rs113435750
               1.6e-03 0.27 GBR
## OCN_(n=2) 1CN_(n=19) 2CN_(n=28) 3CN_(n=18) 4CN_(n=6)
## rs113877493 100 73.68 7.14 0.00 0
## rs6607368 100 84.21 21.43 5.56
                          7.14
                                   0.00
## rs8067765 100 63.16
## rs60952743 100 63.16
                            7.14
                                     0.00
## rs8070238 100 63.16 7.14 0.00 ## rs8072238 100 63.16 7.14 0.00
## 5CN_(n=1) p.values r p.valuesAdjusted r2 POP
## rs113877493 0 2.7e-09 -0.67
              0 4.7e-08 -0.66
## rs6607368
                                 2.8e-06 0.43 FIN
              0 2.3e-07 -0.61 4.2e-06 0.38 FIN
## rs60952743
              0 2.3e-07 -0.61 4.2e-06 0.38 FIN
               0 2.3e-07 -0.61
                                  4.2e-06 0.38 FIN
## rs8072238
               0 2.3e-07 -0.61
                                  4.2e-06 0.38 FIN
##
         1CN (n=2) 2CN (n=1) 3CN (n=1) 4CN (n=2) p.values
              0
                            0
                                   100 2.0e-01 0.85
## rs28856610
                     0
## rs11651338 100 0 100 0 2.0e-01 -0.70
## p.valuesAdjusted r2 POP
## rs4796217
              1e+00 0.73 IBS
## rs28856610
                 1e+00 0.73 IBS
## rs138347191
                1e+00 0.29 IBS
## rs4796216
                1e+00 0.29 IBS
## rs60520102 1e+00 0.29 IBS
```

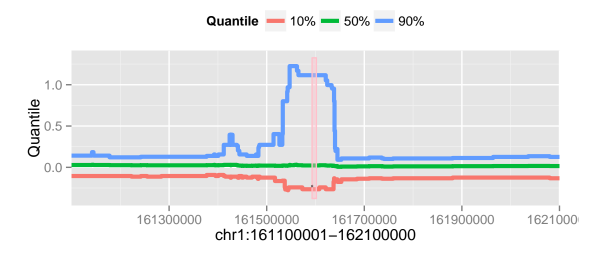
Th eoutput above provides evidence that rs113877493 may be a tagSNP for CCL3L1 deletions in the FIN (p = 1.1e-07,  $r^2$  = 0.44) and GBR (p = 2.6e-04,  $r^2$  = 0.27) populations.

## 4 Indentifying poplymorphic regions

CNVrd2 can also be used to identity CN polymorphic regions and the putative boundaries of these regions. We reuse the data from the FCGR3 locus to investigate the polymorphic region around the two genes.

```
fcgr3PolymorphicRegion <- identifyPolymorphicRegion(Object = objectCNVrd2,
    segmentObject = resultSegment, plotLegend = FALSE)
## Calculate segmentation scores for polymorphic regions</pre>
```





To plot a small region around the gene, we use the funtion plotPolymorphicRegion. Users can change typePlot to only plot SDs or percentiles.

```
## IRanges of length 5

## start end width

## [1] 161392001 161394000 2000

## [2] 161424001 161426000 2000

## [3] 161543001 161544000 1000

## [4] 161564001 161602000 38000

## [5] 161633001 161635000 2000
```

```
## IRanges of length 4

## start end width

## [1] 161392001 161394000 2000

## [2] 161424001 161426000 2000

## [3] 161564001 161565000 1000

## [4] 161573001 161602000 29000
```

Here, we are using standard deviations to identify polymorphic regions. Therefore, the putative boundaries of these regions rely on the parameter sdThreshold. Alternatively, users can use different percentiles to identify these regions. In that case, two parameters quantileValue and thresholdForPolymorphicRegions can be used to adjust the boundaries of regions. For example, here we can set small sdThreshold values to obtain only high-polymorphic regions (e.g., CCL3L1), but it can omit some medium-polymorphic regions (e.g., FCGR3A/3B). Figure 8 and 9 depicts two different thresholds resulting in different polymorphic regions.

In the function identifyPolymorphicRegion, if we would like to obtain only polymorphic regions which differentiate between populations (e.g., to detect evidence of selection) then we can use the option VstTest=TRUE. This option will calculate the Vst statistics (Redon  $et^{\sim}al.$ , 2006). Users have to supply a vector which includes population information in popName. The returned putative boundaries will be the intersection of polymorphic regions and regions having maxVst >= thresholdVST.

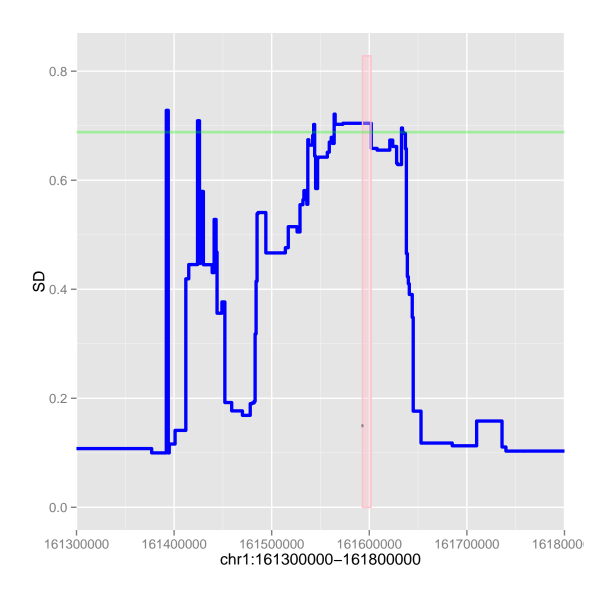


Figure 8: CN polymorphic region at FCGR3 locus, represented by quantiles of the distribution of segmentation scores across samples.

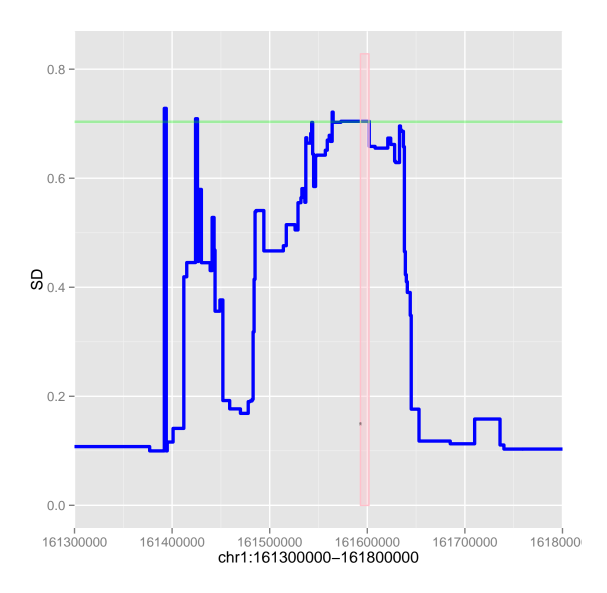


Figure 9: CN polymorphic region at FCGR3 locus, represented by quantiles of the distribution of segmentation scores across samples.

## 5 Note

If we use the option entireGene = FALSE in the step segmentation then the pipeline will not refine the segmentation results (the results will be the same as the pipeline used in Nguyen  $et^{\sim}al$ . (2013)).

## 6 Session information

```
sessionInfo()
## R version 3.0.2 (2013-09-25)
## Platform: x86_64-pc-linux-gnu (64-bit)
##
## locale:
   [1] LC_CTYPE=en_NZ.UTF-8
                                  LC_NUMERIC=C
## [3] LC_TIME=en_NZ.UTF-8
                                  LC_COLLATE=en_NZ.UTF-8
   [5] LC_MONETARY=en_NZ.UTF-8
                                  LC_MESSAGES=en_NZ.UTF-8
## [7] LC_PAPER=en_NZ.UTF-8
                                  LC_NAME=C
## [9] LC_ADDRESS=C
                                  LC_TELEPHONE=C
## [11] LC_MEASUREMENT=en_NZ.UTF-8 LC_IDENTIFICATION=C
## attached base packages:
## [1] grid
               parallel stats
                                    graphics grDevices utils
## [7] datasets methods base
##
## other attached packages:
## [1] codetools_0.2-8
                               knitr_1.5
## [3] CNVrd2_1.1.3
                               gridExtra_0.9.1
## [5] ggplot2_0.9.3.1
                               rjags_3-11
## [7] coda_0.16-1
                               lattice_0.20-24
## [9] VariantAnnotation_1.6.8 Rsamtools_1.12.4
## [11] Biostrings_2.28.0
                               GenomicRanges_1.12.5
## [13] IRanges_1.18.4
                               BiocGenerics_0.6.0
## [15] DNAcopy_1.34.0
## loaded via a namespace (and not attached):
```

```
[1] AnnotationDbi_1.22.6 Biobase_2.20.1
    [3] biomaRt_2.16.0
                               bitops_1.0-6
    [5] BSgenome_1.28.0
##
                               colorspace_1.2-4
    [7] compiler_3.0.2
                               DBI_0.2-7
    [9] dichromat_2.0-0
                               digest_0.6.3
## [11] evaluate_0.5.1
                               formatR_0.10
## [13] GenomicFeatures_1.12.4 gtable_0.1.2
## [15] highr_0.3
                               labeling_0.2
## [17] MASS_7.3-29
                               munsell_0.4.2
## [19] plyr_1.8
                               proto_0.3-10
## [21] RColorBrewer_1.0-5
                               RCurl_1.95-4.1
## [23] reshape2_1.2.2
                               RSQLite_0.11.4
## [25] rtracklayer_1.20.4
                               scales_0.2.3
## [27] stats4_3.0.2
                               stringr_0.6.2
## [29] tools_3.0.2
                               XML_3.98-1.1
## [31] zlibbioc_1.6.0
```

## References

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Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G, Durbin R, et ~al. (2009). "The sequence alignment/map format and SAMtools." Bioinformatics, 25(16), 2078–2079.

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