# Next-generation sequencing bulk segregant analysis with QTLseqr

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#### Contents

rent version: Development - 0.6.3
dard workflow
uick Start
put data
aporting SNPs from GATK
iltering SNPs
unning the analysis
lotting the data $\dots$ 1
xtracting QTL data
ımmary

### Current version: Development - 0.6.3

#### Standard workflow

#### If you use QTLseqr in published research, please cite:

Mansfeld B.N. and Grumet R, QTLseqr: An R package for bulk segregant analysis with next-generation sequencing *The Plant Genome* doi:10.3835/plantgenome2018.01.0006

We also recommend citing the paper for the corresponding method you work with.

#### QTL-seq method:

Takagi, H., Abe, A., Yoshida, K., Kosugi, S., Natsume, S., Mitsuoka, C., Uemura, A., Utsushi, H., Tamiru, M., Takuno, S., Innan, H., Cano, L. M., Kamoun, S. and Terauchi, R. (2013), QTL-seq: rapid mapping of quantitative trait loci in rice by whole genome resequencing of DNA from two bulked populations. *Plant J*, 74: 174–183. doi:10.1111/tpj.12105

#### G prime method:

Magwene PM, Willis JH, Kelly JK (2011) The Statistics of Bulk Segregant Analysis Using Next Generation Sequencing. *PLOS Computational Biology* 7(11): e1002255. https://doi.org/10.1371/journal.pcbi.1002255

#### **Quick Start**

Here are the basic steps required to run and plot QTLseq and G' analysis

```
#load the package
library("QTLseqr")
```

```
#Set sample and file names
HighBulk <- "SRR834931"
LowBulk <- "SRR834927"
file <- "SNPs_from_GATK.table"</pre>
#Choose which chromosomes will be included in the analysis (i.e. exclude smaller contigs)
Chroms <- paste0(rep("Chr", 12), 1:12)</pre>
#Import SNP data from file
df <-
    importFromGATK(
        file = file,
        highBulk = HighBulk,
        lowBulk = LowBulk,
        chromList = Chroms
    )
#Filter SNPs based on some criteria
df_filt <-
    filterSNPs(
        SNPset = df,
        refAlleleFreq = 0.20,
        minTotalDepth = 100,
        maxTotalDepth = 400,
        minSampleDepth = 40,
        minGQ = 99
    )
#Run G' analysis
df_filt <- runGprimeAnalysis(SNPset = df_filt,</pre>
                              windowSize = 1e6,
                              outlierFilter = "deltaSNP")
#Run QTLseq analysis
df_filt <- runQTLseqAnalysis(</pre>
    SNPset = df_filt,
    windowSize = 1e6,
    popStruc = "F2",
    replications = 10000,
    intervals = c(95, 99)
)
#Plot
plotQTLStats(
    SNPset = df_filt,
    var = "Gprime",
    plotThreshold = TRUE,
    q = 0.01
)
plotQTLStats(
    SNPset = df_filt,
    var = "deltaSNP",
```

```
plotIntervals = TRUE)

#export summary CSV
getQTLTable(
    SNPset = df_filt,
    alpha = 0.01,
    export = TRUE,
    fileName = "my_BSA_QTL.csv"
)
```

#### Input data

QTLseqr currently only supports table format SNP data exported from the VariantsToTable function built in to GATK. We hope to support import from any VCF file soon.

#### Importing SNPs from GATK

Working directly with the GATK best practices guide for whole genome sequence should result in a VCF that is compatible with QTLseqr. In general the workflow suggested by GATK is per-sample variant calling followed by joint genotyping across samples. This will produce a VCF file that includes **BOTH** bulks, each with a different sample name (here SRR834927 and SRR834931), one SNP for example:

CHROM	POS	ID	REF	ALT	QUAL	FILTER	INFO	FORMAT	SRR834927	SRR834931
Chr1	31071		A	G	1390.44	PASS	\*	GT:AD:DP:GQ:PL	0/1:34,36:70:99:897,0,855	0/1:26,22:48:99:522,0,698

<sup>\*</sup>info column removed for brevity

GATK have provided a fast VCF parser, the VariantsToTable tool, that extracts the necessary fields for easy use in downstream analysis.

We highly recommend reading What is a VCF and how should I interpret it? for more information on GATK VCF Fields and Genotype Fields

Though the use of GATK's VariantsToTable function is out of the scope of this vignette, the syntax for use with QTLseqr should look something like this:

```
java -jar GenomeAnalysisTK.jar \
-T VariantsToTable \
-R ${REF} \
-V ${NAME} \
-F CHROM -F POS -F REF -F ALT \
-GF AD -GF DP -GF GQ -GF PL \
-o ${NAME}.table
```

Where \${REF} is the reference genome file and \${NAME} is VCF file you wish to parse.

To run QTLseqr successfully, the required VCF fields (-F) are CHROM (Chromosome) and POS (Position). the required Genotype fields (-GF) are AD (Allele Depth), DP (Depth). Recommended fields are REF (Reference allele) and ALT (Alternative allele) Recommended Genotype fields are PL (Phred-scaled likelihoods) and GQ (Genotype Quality).

#### Import function

Let's install and load the QTLseqr package:

```
#Install step if you have not done so yet:
#devtools::install_github("bmansfeld/QTLseqr")
library("QTLseqr")
```

The importFromGATK function imports SNP data from the output of the VariantsToTable function in GATK. After importing the data, the function then calculates total reference allele frequency for both bulks together, the SNP index for each bulk, and the  $\Delta(SNP-index)$ .

To demonstrate the use of this function we will load the Yang et al. (2013) data file. We first need to download the package that contains the data from github.

```
#download and load data package (~50Mb)
devtools::install_github("bmansfeld/Yang2013data")
library("Yang2013data")

#Import the data
rawData <- system.file(
    "extdata",
    "Yang_et_al_2013.table",
    package = "Yang2013data",
    mustWork = TRUE)</pre>
```

If you have your own data you can simply refer to it directly:

```
rawData <- "C:/PATH/TO/MY/DIR/My_BSA_data.table"
```

We define the sample name for each of the bulks. This should correspond to the sample names in the VCF returned by GATK. We also define a vector of the chromosomes to be included in the analysis (i.e. exclude smaller contigs), In this case, Chr1, Chr2... Chr12.

```
HighBulk <- "SRR834931"
LowBulk <- "SRR834927"
Chroms <- paste0(rep("Chr", 12), 1:12)</pre>
```

We then use the importFromGATK function to import the raw data. After importing the data, the function then calculates total reference allele frequency for both bulks together, the SNP-index for each SNP in each bulk and the  $\Delta(SNP$ -index) and returns a data frame.

```
Reference \ allele \ frequency = \frac{Ref \ allele \ depth_{HighBulk} + Ref \ allele \ depth_{LowBulk}}{Total \ read \ depth \ for \ both \ bulks} SNP\text{-}index_{per \ bulk} = \frac{Alternate \ allele \ depth}{Total \ read \ depth} \Delta(SNP\text{-}index) = SNP\text{-}index_{HighBulk} - SNP\text{-}index_{LowBulk}
```

Let's import

```
#import data
df <-
   importFromGATK(
      file = rawData,
      highBulk = HighBulk,
      lowBulk = LowBulk,</pre>
```

```
chromList = Chroms
)
```

#### Loaded data frame

The loaded data frame should look like this:

#### head(df)

```
POS REF ALT AD_REF.LOW AD_ALT.LOW DP.LOW GQ.LOW
##
     CHROM
                                                                       PL.LOW
## 1
      Chr1 31071
                         G
                                    34
                                                36
                                                        70
                                                                    897,0,855
                         Τ
## 2
      Chr1 31478
                    C
                                                52
                                                        86
                                                               99 1363,0,844
                                    34
                         G
## 3
      Chr1 33667
                    Α
                                    20
                                                48
                                                        68
                                                               99
                                                                  1331,0,438
                         Т
## 4
      Chr1 34057
                    С
                                    38
                                                40
                                                        78
                                                               99
                                                                  1059,0,996
                         C
## 5
      Chr1 35239
                    Α
                                    25
                                                36
                                                        61
                                                               99
                                                                    987,0,630
                    Т
                         C
                                                42
                                                        78
                                                               99 1066,0,906
##
      Chr1 38389
                                    36
##
     SNPindex.LOW AD_REF.HIGH AD_ALT.HIGH DP.HIGH
                                                      GQ.HIGH
                                                                    PL.HIGH
## 1
        0.5142857
                             26
                                          22
                                                   48
                                                            99
                                                                  522,0,698
## 2
        0.6046512
                             40
                                          34
                                                   74
                                                            99
                                                                848,0,1099
## 3
        0.7058824
                             24
                                          29
                                                   53
                                                            99
                                                                  765,0,599
## 4
        0.5128205
                             29
                                          26
                                                   55
                                                            99
                                                                  673,0,772
## 5
        0.5901639
                             40
                                          60
                                                  100
                                                            99 1632,0,1015
##
                             42
                                          40
  6
        0.5384615
                                                   82
                                                                984,0,1105
                                                            99
##
     SNPindex.HIGH
                       REF_FRQ
                                    deltaSNP
## 1
         0.4583333 0.5084746 -0.055952381
         0.4594595 0.4625000 -0.145191703
## 2
         0.5471698 0.3636364 -0.158712542
## 3
## 4
         0.4727273 0.5037594 -0.040093240
## 5
         0.6000000 0.4037267
                                0.009836066
         0.4878049 0.4875000 -0.050656660
```

Let's review the column headers:

- CHROM The chromosome this SNP is in
- POS The position on the chromosome in nt
- REF The reference allele at that position
- ALT The alternate allele
- DP.HIGH The read depth at that position in the high bulk
- AD\_REF.HIGH The allele depth of the reference allele in the high bulk
- AD\_ALT.HIGH The alternate allele depth in the the high bulk
- GQ.HIGH The genotype quality score, (how confident we are in the genotyping)
- SNPindex.HIGH The calculated SNP-index for the high bulk
- Same as above for the low bulk
- REF FRQ The reference allele frequency as defined above
- deltaSNP The  $\Delta(SNP-index)$  as defined above

#### Filtering SNPs

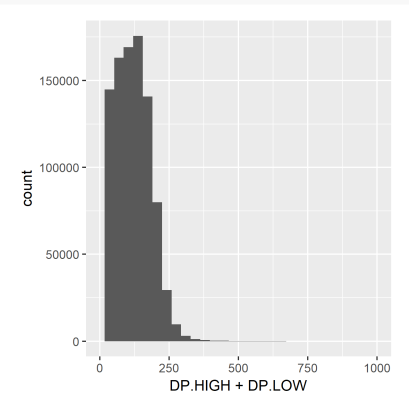
Now that we have loaded the data into R we can start cleaning it up by filtering some of the low confidence SNPs. While GATK has its own filtering tools, QTLseqr offers some options for filtering that may help reduce noise and improve results. Filtering is mainly based on read depth for each SNP, such that we can try to eliminate SNPs with low confidence, due to low coverage, and SNPs that may be in repetitive regions and thus have inflated read depth.

#### Read depth histograms

One way to assess filtering thresholds is by plotting histograms of the read depths. We can get an idea of where to draw our thresholds. We'll use the ggplot2 package for this purpose, but you could use base R to plot as well.

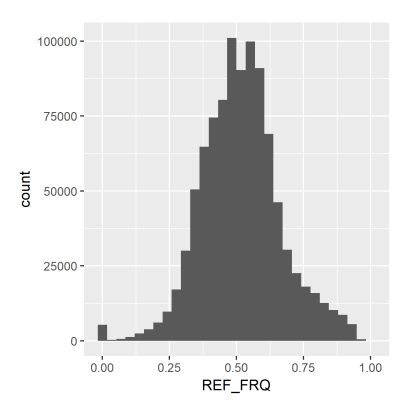
Lets look at total read depth for example:

```
library("ggplot2")
ggplot(data = df) +
    geom_histogram(aes(x = DP.HIGH + DP.LOW)) +
    xlim(0,1000)
```



... or look at total reference allele frequency:

```
ggplot(data = df) +
  geom_histogram(aes(x = REF_FRQ))
```



#### Using the filterSNPs function

## ...Filtered 4729 SNPs

Now that we have an idea about our read depth distribution we can filter out low confidence SNPS. In general we recommend filtering extremely low and high coverage SNPs, either in both bulks (minTotalDepth/maxTotalDepth) and/or in each bulk separately (minSampleDepth). We have the option to filter based on reference allele frequency (refAlleleFreq), this removes SNPs that for some reason are overor under-represented in BOTH bulks. We can also use the GATK GQ score (Genotype Quality) to filter out low confidence SNPs. If the verbose parameter is set to TRUE (default) the function will report the numbers of SNPs filtered in each step.

```
df_filt <-
   filterSNPs(
        SNPset = df,
        refAlleleFreq = 0.20,
        minTotalDepth = 100,
        maxTotalDepth = 400,
        minSampleDepth = 40,
        minGQ = 99,
        verbose = TRUE
## Filtering by reference allele frequency: 0.2 <= REF_FRQ <= 0.8
## ...Filtered 59754 SNPs
## Filtering by total sample read depth: Total DP >= 100
## ...Filtered 378814 SNPs
## Filtering by total sample read depth: Total DP <= 400
## ...Filtered 744 SNPs
## Filtering by per sample read depth: DP >= 40
```

```
## Filtering by Genotype Quality: GQ >= 99
## ...Filtered 6677 SNPs
## Original SNP number: 969487, Filtered: 450718, Remaining: 518769
```

This step is quick and we can go back and plot some histograms to see if we are happy with the results, and we can quickly re-run the filtering step if not.

#### Running the analysis

The analysis in QTLseqr is an implementation of both pipelines for bulk segregant analysis, G' and  $\Delta(SNP-index)$ , described by Magwene et al. (2011) and Takagi et al. (2013), respectively. We recommend reading both papers to fully understand the considerations and math behind the analysis.

There are two main analysis functions: 1. runGprimeAnalysis - performs Magwene et al type G' analysis 1. runQTLseqAnalysis - performs Takagi et al type QTLseq analysis

#### QTLseq analysis

Takagi et al. (2013) developed the method for QTLseq type NGS-BSA. The analysis is based on calculating the allele frequency differences, or  $\Delta(SNP\text{-}index)$ , from the allele depths at each SNP. To determine regions of the genome that significantly differ from the expected  $\Delta(SNP\text{-}index)$  of 0, a simulation approach is used. Briefly, at each read depth, simulated SNP frequencies are bootstrapped, and the extreme quantiles are used as simulated confidence intervals. The true data are averaged over a sliding window and regions that surpass the CI are putative QTL.

When the analysis is run the following steps are performed:

- 1. First the number of SNPs within the sliding window are counted.
- 2. A tricube-smoothed  $\Delta(SNP-index)$  is calculated within the set window size.
- 3. The minimum read depth at each position is calculated and the tricube-smoothed depth is calculated for the window.
- 4. The simulation is performed for data derived read depths (can be set by the user): Alternate allele frequency is calculated per bulk based on the population type and size (F2 or RIL)  $\Delta(SNP\text{-}index)$  is simulated over several replications (default = 10000) for each bulk. The quantiles from the simulations are used to estimate the confidence intervals. Say for example the 99th quantile of 10000  $\Delta(SNP\text{-}index)$  simulations represents the 99% confidence interval for the true data.
- 5. Confidence intervals are matched with the relevant window depth at each SNP.

Here is an example for running the analysis for an F2 population, with a bulk size of 300 individuals in each bulk. The simulation is bootstrapped 10000 times and the two-sided 95 and 99% confidence intervals are calculated:

```
df_filt <- runQTLseqAnalysis(df_filt,
    windowSize = 1e6,
    popStruc = "F2",
    bulkSize = 300,
    replications = 10000,
    intervals = c(95, 99)
    )</pre>
```

```
## Counting SNPs in each window...
## Calculating tricube smoothed delta SNP index...
## Returning the following two sided confidence intervals: 95, 99
## Variable 'depth' not defined, using min and max depth from data: 40-197
```

```
## Assuming bulks selected from F2 population, with 300 individuals per bulk.
## Simulating 10000 SNPs with reads at each depth: 40-197
## Keeping SNPs with >= 0.3 SNP-index in both simulated bulks
## Joining, by = "tricubeDP"
```

#### G' analysis

An alternate approach to determine statistical significance of QTL from NGS-BSA was proposed by Magwene et al. (2011) – calculating a modified G statistic for each SNP based on the observed and expected allele depths and smoothing this value using a tricube smoothing kernel. Using the smoothed G statistic, or G', Magwene et al. allow for noise reduction while also addressing linkage disequilibrium between SNPs. Furthermore, as G' is close to being log normally distributed, p-values can be estimated for each SNP using non-parametric estimation of the null distribution of G'. This provides a clear and easy-to-interpret result as well as the option for multiple testing corrections.

Here, we will briefly summarize the steps performed by the main analysis function, runGprimeAnalysis.

The following steps are performed:

- 1. First the number of SNPs within the sliding window are counted.
- 2. A tricube-smoothed  $\Delta(SNP\text{-}index)$  is calculated within the set window size.
- 3. Genome-wide G statistics are calculated by getG. G is defined by the equation:

$$G = 2 * \sum n_i * ln(\frac{obs(n_i)}{exp(n_i)})$$

Where for each SNP,  $n_i$  from i = 1 to 4 corresponds to the reference and alternate allele depths for each bulk, as described in the following table:

Allele	High Bulk	Low Bulk
Reference Alternate	$n_1$ $n_3$	$n_2$ $n_4$

... and  $obs(n_i)$  are the observed allele depths as described in the data frame. getG calculates the G statistic using expected values assuming read depth is equal for all alleles in both bulks:

$$exp(n_1) = \frac{(n_1 + n_2) * (n_1 + n_3)}{(n_1 + n_2 + n_3 + n_4)}$$

$$exp(n_2) = \frac{(n_2 + n_1) * (n_2 + n_4)}{(n_1 + n_2 + n_3 + n_4)}$$

$$exp(n_3) = \frac{(n_3 + n_1) * (n_3 + n_4)}{(n_1 + n_2 + n_3 + n_4)}$$

$$exp(n_4) = \frac{(n_4 + n_2) * (n_4 + n_3)}{(n_1 + n_2 + n_3 + n_4)}$$

4. G' - A tricube-smoothed G statistic is predicted by constant local regression within each chromosome using the tricubeStat function. This works as a weighted average across neighboring SNPs that accounts for Linkage disequilibrium (LD) while minimizing noise attributed to SNP calling errors. G values for neighboring SNPs within the window are weighted by physical distance from the focal SNP.

- 5. P-values are estimated based using the non-parametric method described by Magwene et al. 2011 with the function getPvals. Briefly, using the natural log of G' a median absolute deviation (MAD) is calculated. The G' set is trimmed to exclude outlier regions (i.e. QTL) based on Hampel's rule. An alternate method for filtering out QTL that we propose is using absolute  $\Delta(SNP\text{-}index)$  values greater than a set threshold (default = 0.1) to filter out potential QTL. An estimation of the mode of the trimmed set is calculated using the mlv function from the package modeest. Finally, the mean and variance of the set are estimated using the median and mode and p-values are estimated from a log normal distribution.
- 6. Negative Log10- and Benjamini-Hochberg adjusted p-values are calculated using p.adjust.

Let's run the function:

## Calculating p-values...

```
df_filt <- runGprimeAnalysis(df_filt,
    windowSize = 1e6,
    outlierFilter = "deltaSNP",
    filterThreshold = 0.1)

## Counting SNPs in each window...
## Calculating tricube smoothed delta SNP index...
## Calculating G and G' statistics...
## Using deltaSNP-index to filter outlier regions with a threshold of 0.1
## Estimating the mode of a trimmed G prime set using the 'modeest' package...</pre>
```

As this is window is using a tricube-smoothing kernel the window size can be much larger than you might expect. We however choose a window size of 1Mb for the sliding window analysis, for a discussion about window size we recommend reading Magwene et al. (2011). In general larger windows will produced smoother data. The functions making these calculations are rather fast, so we recommend testing several window sizes for your data, and deciding on the optimal size.

Some additional columns are added to the filtered data frame:

#### head(df filt)

```
POS REF ALT AD_REF.LOW AD_ALT.LOW DP.LOW GQ.LOW
##
     CHROM
                                                                      PL.LOW
                                                                   897,0,855
## 1
      Chr1 31071
                    Α
                         G
                                    34
                                                36
                                                       70
                                                               99
                         Т
## 2
      Chr1 31478
                    С
                                    34
                                                52
                                                       86
                                                               99 1363,0,844
## 3
      Chr1 33667
                         G
                                                48
                                                               99 1331,0,438
                    A
                                    20
                                                       68
## 4
      Chr1 34057
                    C
                         Т
                                    38
                                                40
                                                       78
                                                               99 1059,0,996
                         С
                                    25
                                                36
                                                       61
## 5
      Chr1 35239
                    Α
                                                               99
                                                                   987,0,630
  6
      Chr1 38389
                    Τ
                         C
                                    36
                                                42
                                                       78
                                                               99 1066,0,906
                                                      GQ.HIGH
##
     SNPindex.LOW
                   AD_REF.HIGH AD_ALT.HIGH DP.HIGH
                                                                   PL.HIGH
## 1
        0.5142857
                             26
                                          22
                                                   48
                                                            99
                                                                 522,0,698
                                                   74
## 2
        0.6046512
                             40
                                          34
                                                            99
                                                                848,0,1099
## 3
        0.7058824
                             24
                                          29
                                                   53
                                                            99
                                                                 765,0,599
                                          26
                                                   55
## 4
        0.5128205
                             29
                                                            99
                                                                 673,0,772
## 5
        0.5901639
                             40
                                          60
                                                  100
                                                            99 1632,0,1015
##
  6
        0.5384615
                             42
                                          40
                                                   82
                                                            99
                                                                984,0,1105
                                    deltaSNP
##
     SNPindex.HIGH
                       REF_FRQ
                                             nSNPs tricubeDeltaSNP minDP
## 1
         0.4583333 0.5084746 -0.055952381
                                                881
                                                        -0.07365553
                                                                         48
## 2
         0.4594595 0.4625000 -0.145191703
                                                881
                                                        -0.07365807
                                                                         74
## 3
         0.5471698 0.3636364 -0.158712542
                                                883
                                                        -0.07367173
                                                                         53
## 4
         0.4727273 0.5037594 -0.040093240
                                                883
                                                        -0.07367416
                                                                         55
## 5
         0.6000000 0.4037267
                               0.009836066
                                                883
                                                        -0.07368154
                                                                         61
## 6
         0.4878049 0.4875000 -0.050656660
                                                883
                                                        -0.07370120
                                                                         78
     tricubeDP
##
                     CI 95
                                  CI 99
                                                  G
                                                      Gprime
                                                                 pvalue
## 1
             70 -0.1714286 -0.2285714 0.35697092 1.919586 0.4670188
```

```
## 2
            70 -0.1714286 -0.2285714 3.38161778 1.919650 0.4669994
## 3
            70 -0.1714286 -0.2285714 3.23692853 1.919994 0.4668952
## 4
            70 -0.1714286 -0.2285714 0.20748641 1.920056 0.4668766
            70 -0.1714286 -0.2285714 0.01521766 1.920242 0.4668204
## 5
## 6
            70 -0.1714286 -0.2285714 0.41076961 1.920737 0.4666705
     negLog10Pval
##
                     qvalue
        0.3306657 0.6683592
## 1
        0.3306837 0.6683425
## 2
## 3
        0.3307806 0.6682994
## 4
        0.3307979 0.6682904
## 5
        0.3308502 0.6682467
## 6
        0.3309896 0.6681446
```

- nSNPs the number of SNPs bracketing the focal SNP within the set sliding window
- tricubeDeltaSNP the tricube-smoothed  $\Delta(SNP-index)$
- G the G value for the SNP
- Gprime the tricube-smoothed G value
- pvalue the p-value calculated by non-parametric estimation
- negLog10Pval the  $-log_{10}(p\text{-}value)$
- qvalue Benjamini-Hochberg adjusted p-values

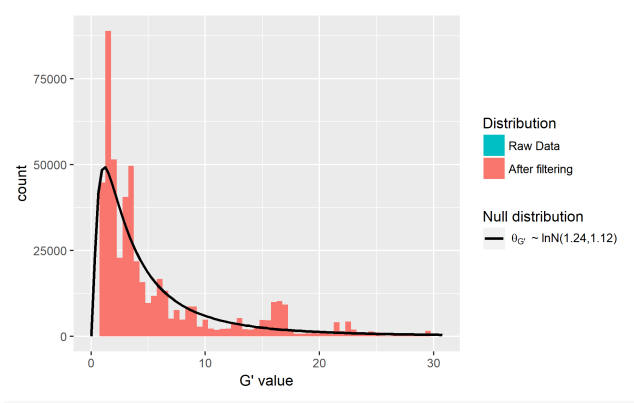
#### Plotting the data

QTLseqr offers two main plotting functions to check the validity of the G' analysis and to plot genome-wide or chromosome specific QTL analysis plots.

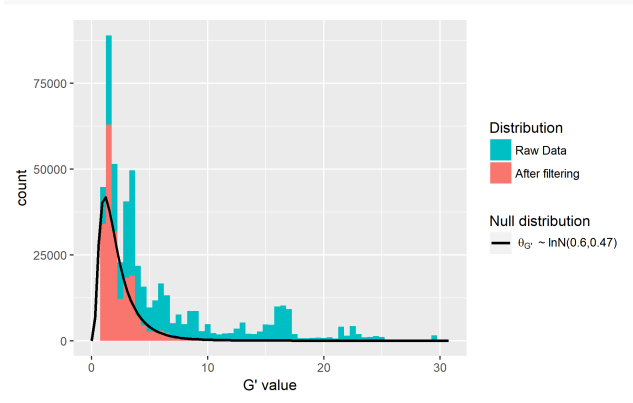
#### G' distribution plots

Due to the fact that p-values are estimated from the null distribution of G', an important check is to see if the null distribution of G' values is close to log normally distributed. For this purpose we use the plotGprimeDist function, which plots the G' histograms of both raw and filtered G' sets (see P-value calculation above) alongside the log-normal null distribution (which is reported in the legend). We can also use this to test which filtering method (Hampel or DeltaSNP) estimates a more accurate null distribution. If you use the "deltaSNP" method plotting G' distributions with different filter thresholds might also help reveal a better G' null distribution.

```
plotGprimeDist(SNPset = df_filt, outlierFilter = "Hampel")
```



plotGprimeDist(SNPset =df\_filt, outlierFilter = "deltaSNP", filterThreshold = 0.1)

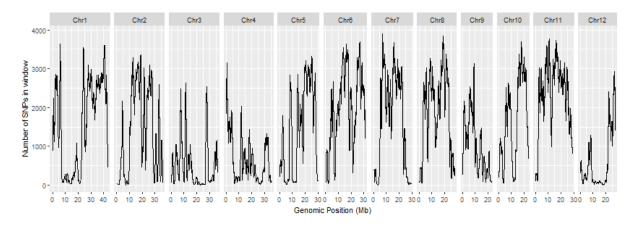


#### QTL analysis plots

Now that we are happy with our filtered data and it seems that the G' distribution is close to log-normal, we can finally plot some genome-wide figures and try to identify QTL.

Let's start by plotting the SNP/window distribution:

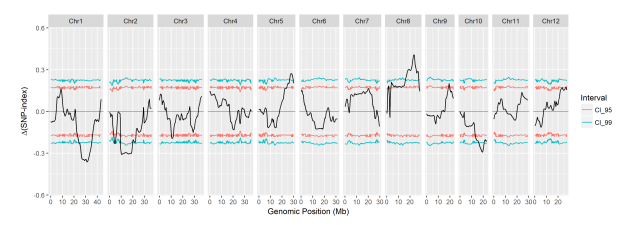
```
p1 <- plotQTLStats(SNPset = df_filt, var = "nSNPs")
p1</pre>
```



This is informative as we can assess if there are regions with extremely low SNP density.

More importantly lets identify some QTL by plotting the smoothed  $\Delta(SNP\text{-}index)$  and G' values. If we've performed QTLseq analysis we can also set plotIntervals to TRUE and plot the confidence intervals to identify QTL using that method.

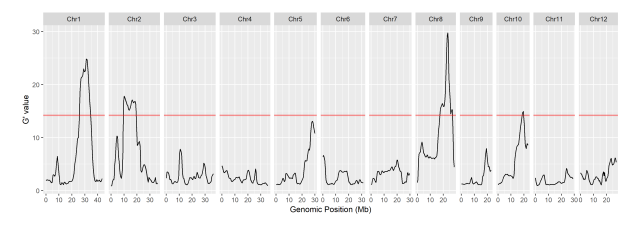
```
p2 <- plotQTLStats(SNPset = df_filt, var = "deltaSNP", plotIntervals = TRUE)
p2</pre>
```



We can see that there are some regions that have  $\Delta(SNP\text{-}index)$  that pass the confidence interval thresholds, and are putative QTL. The directionality of the  $\Delta(SNP\text{-}index)$  is also important for G' analysis. If the allele contributing to the trait is from the reference parent the  $\Delta(SNP\text{-}index)$  should be less than 0. However, if the  $\Delta(SNP\text{-}index) > 0$  then the contributing parent is the one with the alternate alleles.

Let's look at the G' values to see if these regions are significant and pass the FDR (q) of 0.01.

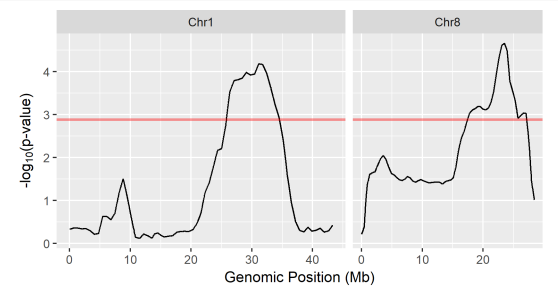
```
p3 <- plotQTLStats(SNPset = df_filt, var = "Gprime", plotThreshold = TRUE, q = 0.01)
p3
```



Great! It looks like there are QTL identified on Chromosomes 1, 2, 5, 8 and 10. Based on the  $\Delta(SNP\text{-}index)$  and G' plots the QTL from Chr1 originates from the reference parent (Nipponbare rice, in this case) and the QTL on Chr8 was contributed by the other parent, for example.

We can also use the plotQTLStats function to the  $-log_{10}(p\text{-}value)$ . While this number is a direct derivative of G' it can be more self explanatory for some. We can use the subset parameter to plot one or a few of the chromosomes, say for a close up figure of a QTL of interest. Here we look at the  $-log_{10}(p\text{-}value)$  plots of Chromosomes 1 and 8:

```
QTLplots <- plotQTLStats(
    SNPset = df_filt,
    var = "negLog10Pval",
    plotThreshold = TRUE,
    q = 0.01,
    subset = c("Chr1", "Chr8")
    )
QTLplots</pre>
```



#### Extracting QTL data

Now that we've plotted and identified some putative QTL we can extract the data using two functions getSigRegions and getQTLTable.

#### Extracting significant regions

The getSigRegions function will produce a list in which each element represents a QTL region. The elements are subsets from the original data frame you supplied. Any contiguous region above with an adjusted p-value above the set alpha will be returned. If there is a dip below the alpha this region will be split to two elements.

Let's examine the head of the first QTL:

```
QTL <- getSigRegions(SNPset = df_filt, alpha = 0.01)
head(QTL[[1]])</pre>
```

```
ALT AD_REF.LOW AD_ALT.LOW DP.LOW GQ.LOW
##
     qtl CHROM
                     POS REF
          Chr1 25842439
## 1
                           Т
                                С
                                          21
                                                      59
                                                              80
                                                                     99
       1
## 2
          Chr1 25844940
                                G
                                          20
                                                      59
                                                              79
                                                                     99
       1
                           Α
## 3
       1
          Chr1 25846683
                           Α
                                G
                                          16
                                                      47
                                                              63
                                                                     99
          Chr1 25847043
## 4
       1
                           G
                                Α
                                          26
                                                      61
                                                              87
                                                                     99
## 5
       1
          Chr1 25849237
                           G
                                Α
                                          12
                                                      46
                                                              58
                                                                     99
##
  6
          Chr1 25851646
                           Α
                                Т
                                          12
                                                              53
                                                                     99
##
         PL.LOW SNPindex.LOW
                              AD_REF.HIGH AD_ALT.HIGH DP.HIGH
                                                                GQ.HIGH
## 1 1651,0,439
                    0.7375000
                                        58
                                                     47
                                                             105
                                                                      99
## 2 1675,0,399
                                        74
                                                     52
                                                             126
                                                                      99
                    0.7468354
## 3 1342,0,326
                                        48
                    0.7460317
                                                     30
                                                              78
                                                                      99
## 4 1679,0,563
                                                     90
                                                             180
                                                                      99
                    0.7011494
                                        90
                    0.7931034
## 5 1311,0,222
                                        30
                                                     20
                                                              50
                                                                      99
## 6 1150,0,227
                                                     26
                                                                      99
                    0.7735849
                                        35
                                                              61
##
         PL.HIGH SNPindex.HIGH
                                   REF_FRQ
                                             deltaSNP nSNPs tricubeDeltaSNP
                      0.4476190 0.4270270 -0.2898810
## 1 1189,0,1546
                                                        1068
                                                                   -0.2762683
## 2 1270,0,1984
                      0.4126984 0.4585366 -0.3341370
                                                        1065
                                                                   -0.2764951
## 3
                                                        1055
      768,0,1329
                      0.3846154 0.4539007 -0.3614164
                                                                   -0.2766532
## 4 2369,0,2351
                      0.5000000 0.4344569 -0.2011494
                                                        1055
                                                                   -0.2766858
## 5
       499,0,826
                      0.4000000 0.3888889 -0.3931034
                                                        1054
                                                                   -0.2768848
## 6
       650,0,915
                      0.4262295 0.4122807 -0.3473554
                                                        1052
                                                                   -0.2771032
##
     minDP tricubeDP
                           CI 95
                                       CI 99
                                                      G
                                                          Gprime
                                                                       pvalue
                   71 -0.1690141 -0.2253521 15.998496 14.52061 0.001185737
## 1
        80
## 2
        79
                   71 -0.1690141 -0.2253521 22.572856 14.54205 0.001177278
## 3
        63
                   71 -0.1690141 -0.2253521 18.929521 14.55700 0.001171423
## 4
        87
                   71 -0.1690141 -0.2253521 9.885982 14.56008 0.001170218
## 5
                   71 -0.1690141 -0.2253521 17.901883 14.57889 0.001162903
        50
##
  6
        53
                   71 -0.1690141 -0.2253521 14.579060 14.59955 0.001154930
##
     negLog10Pval
                        qvalue
         2.926012 0.009145189
## 1
## 2
         2.929121 0.009096584
## 3
         2.931286 0.009060519
         2.931733 0.009052934
## 4
## 5
         2.934457 0.009008602
## 6
         2.937444 0.008962478
```

#### Output QTL summary

While getSigRegions is useful for examining every SNP within each QTL and perhaps for some downstream analysis, the getQTLTable will summarize those results and can output a CSV by setting export = TRUE and fileName = "MyQTLsummary.csv". We can set method as either "Gprime" or "QTLseq" depending on the type of analysis; "Gprime" will use alpha as FDR threshold and "QTLseq" will use the interval parameter, which should match one of the intervals calculated above.

Here is the summary for significant regions with a FDR of 0.01:

```
results <- getQTLTable(SNPset = df_filt, method = "Gprime",alpha = 0.01, export = FALSE)
results</pre>
```

```
length nSNPs avgSNPs_Mb peakDeltaSNP
##
     CHROM qtl
                  start
                              end
             1 25842439 34556923 8714484 20263
## 1
      Chr1
                                                      2325
                                                              -0.3631140
                9575918 19411590 9835672 24267
                                                              -0.3081058
## 2
      Chr2
                                                      2467
             1 17427057 27196250 9769193 20529
## 3
      Chr8
                                                      2101
                                                               0.4082090
## 4 Chr10
             1 18619373 19792840 1173467 3590
                                                      3059
                                                              -0.2922478
     avgDeltaSNP maxGprime meanGprime sdGprime
                                                      AUCaT
                                                                 meanPval
## 1
     -0.3348544
                  24.83453
                             21.29574 2.6768852 60704185.7 0.0002362947
## 2
     -0.3007383
                  17.82461
                             16.44684 0.6348595 22256829.6 0.0006525797
## 3
                             18.89438 4.6368929 46533829.4 0.0005309317
       0.3215896
                  29.75614
## 4
     -0.2888448
                  14.95028
                             14.68470 0.2098474
                                                   577793.3 0.0011258224
##
        meanQval
## 1 0.005050808
## 2 0.007346565
## 3 0.006606070
## 4 0.008877309
```

#### The columns are:

- chromosome The chromosome on which the region was identified
- qtl the QTL identification number in this chromosome
- start the start position on that chromosome, i.e. the position of the first SNP that passes the FDR threshold
- end the end position
- length the length in base pairs from start to end of the region
- nSNPs the number of SNPs in the region
- avgSNPs Mb the average number of SNPs/Mb within that region
- peakDeltaSNP the  $\Delta(SNP\text{-}index)$  value at the peak summit
- maxGprime the max G' score in the region
- mean Gprime - the average G' score of that region
- sdGprime the standard deviation of G' within the region
- AUCaT the **A**rea **U**nder the **C**urve but **a**bove the **T**hreshold line, an indicator of how significant or wide the peak is
- meanPval the average p-value in the region
- meanQval the average adjusted p-value in the region

#### Summary

We've reviewed how to load SNP data from GATK and filter the data to contain high confidence SNPs. We then performed  $\Delta(SNP\text{-}index)$  and G' analysis and calculate p-values and q-values based on the tricube-smoothed G' values. The QTL regions that pass our defined threshold can be stored as a list for further analysis or summarized as a table for publication.