

QTLseqrRiceColdTolerance

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```
# install devtools and all other dependent packages first
#utils::install.packages("devtools")
#utils::install.packages("tidyr")
#utils::install.packages("ggplot2")
#utils::install.packages("dplyr")
#utils::install.packages("data.table")
#utils::install.packages("vcfR")
```

```
# use devtools to install QTLseqr from my "Forked" Repository at PBGL
devtools::install_github("PBGLMichaelHall/QTLseqr")
```

```
# Load them into your session so they are available for use
base::library("QTLseqr")
base::library("data.table")
base::library("dplyr")
base::library("tidyr")
base::library("vcfR")
base::library("ggplot2")
```

```
QTLseqr::importFromVCF(file="wGQ-Filt-freebayes~bwa~IRGSP-1.0~both-segregant_bulks~filtered-default.vcf",
highBulk = "ET-pool-385", lowBulk = "ES-pool-430", chromList =c("NC_029256.1", "NC_029257.1", "NC_029258.1",
"NC_029259.1", "NC_029260.1", "NC_029261.1", "NC_029262.1", "NC_029263.1", "NC_029264.1", "NC_029265.1",
"NC_029266.1", "NC_029267.1"), filename = "Hall", filter=FALSE)
```

Scanning file to determine attributes.

File attributes:

- meta lines: 126
- header_line: 127
- variant count: 1714745
- column count: 11

Meta line 126 read in.

All meta lines processed.

gt matrix initialized.

Character matrix gt created.

- Character matrix gt rows: 1714745
- Character matrix gt cols: 11
- skip: 0
- nrows: 1714745
- row_num: 0

Processed variant 1000Processed variant 2000Processed variant 3000Processed variant 4000Processed variant 5000

All variants processed

Keeping SNPs that pass all filters Either PASS or No Filter

Extracting gt element AD

Extracting gt element DP

Extracting gt element GQ

Removing the following chromosomes:

```
HighBulk <- "ET-pool-385"
```

```
LowBulk <- "ES-pool-430"
```

```
file <- "Hall.csv"
```

#Choose which chromosomes/contigs will be included in the analysis,

```
chromList <- c("NC_029256.1","NC_029257.1","NC_029258.1","NC_029259.1","NC_029260.1","NC_029261.1",  
"NC_029262.1","NC_029263.1","NC_029264.1","NC_029265.1","NC_029266.1","NC_029267.1")
```

```
df <-
```

```
  QTLseqr::importFromTable(  
    file = file,  
    highBulk = HighBulk,  
    lowBulk = LowBulk,  
    chromList = chromList,  
    sep = ",",  
  )
```

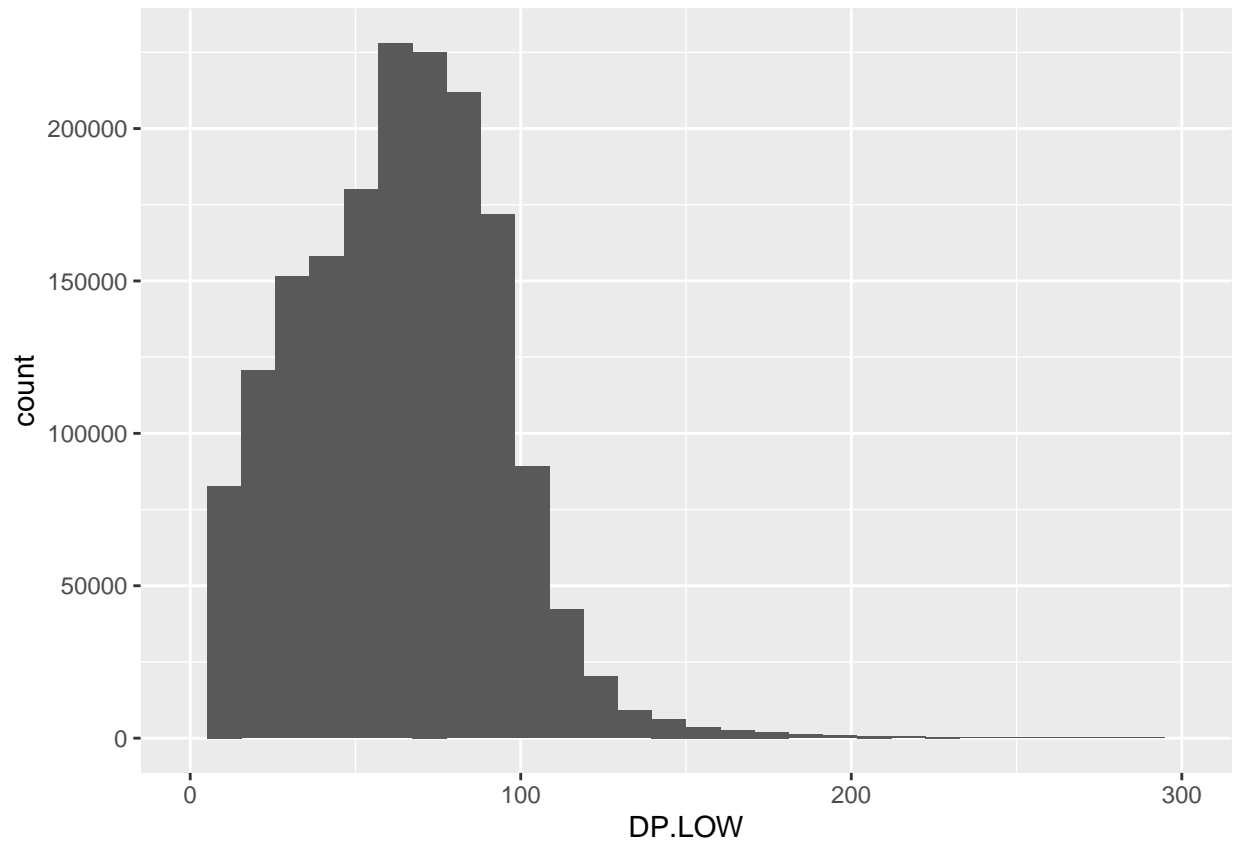
Removing the following chromosomes:

Renaming the following columns: AD_REF.ET-pool-385, AD_ALT.ET-pool-385

Renaming the following columns: AD_REF.ES-pool-430, AD_ALT.ES-pool-430

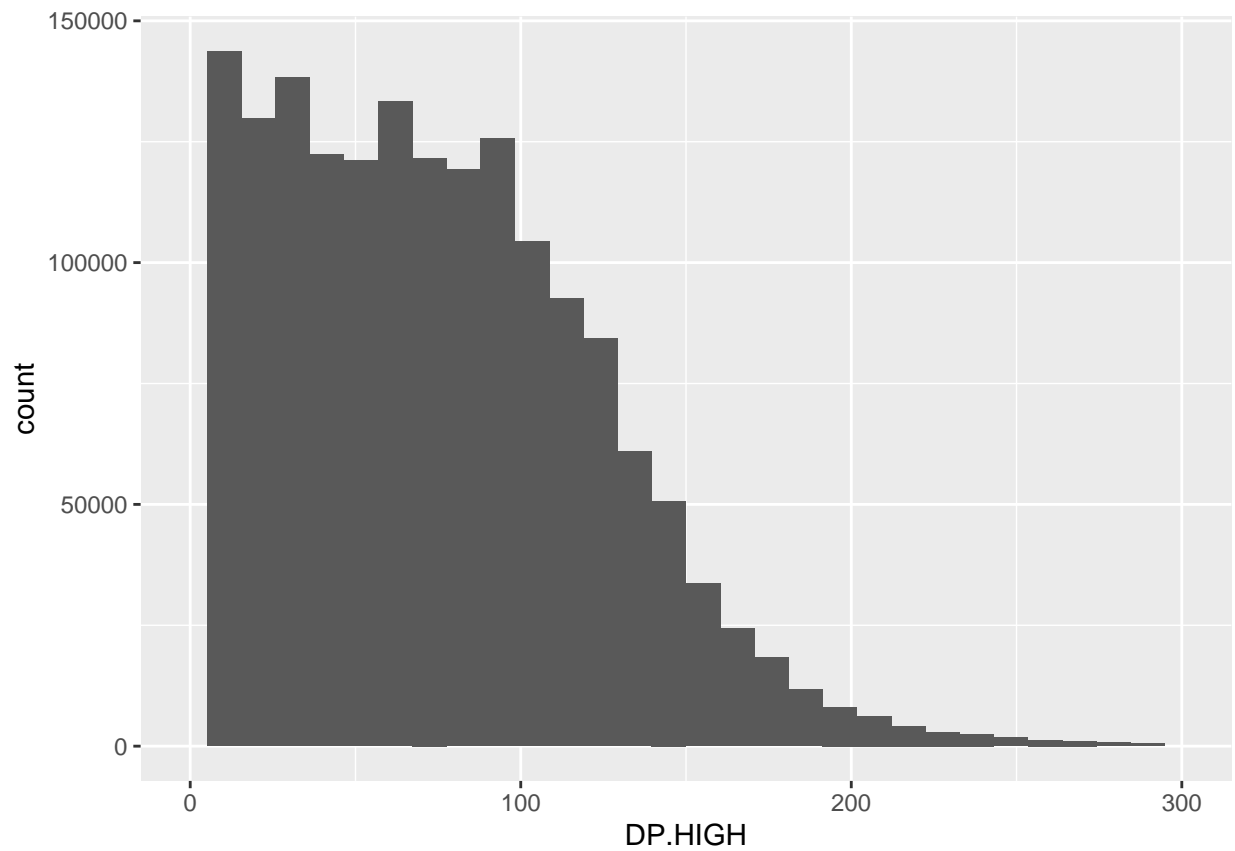
```
ggplot(data = df) + geom_histogram(aes(x = DP.LOW)) + xlim(0,300)
```

``stat_bin()` using `bins = 30`. Pick better value with `binwidth`.`



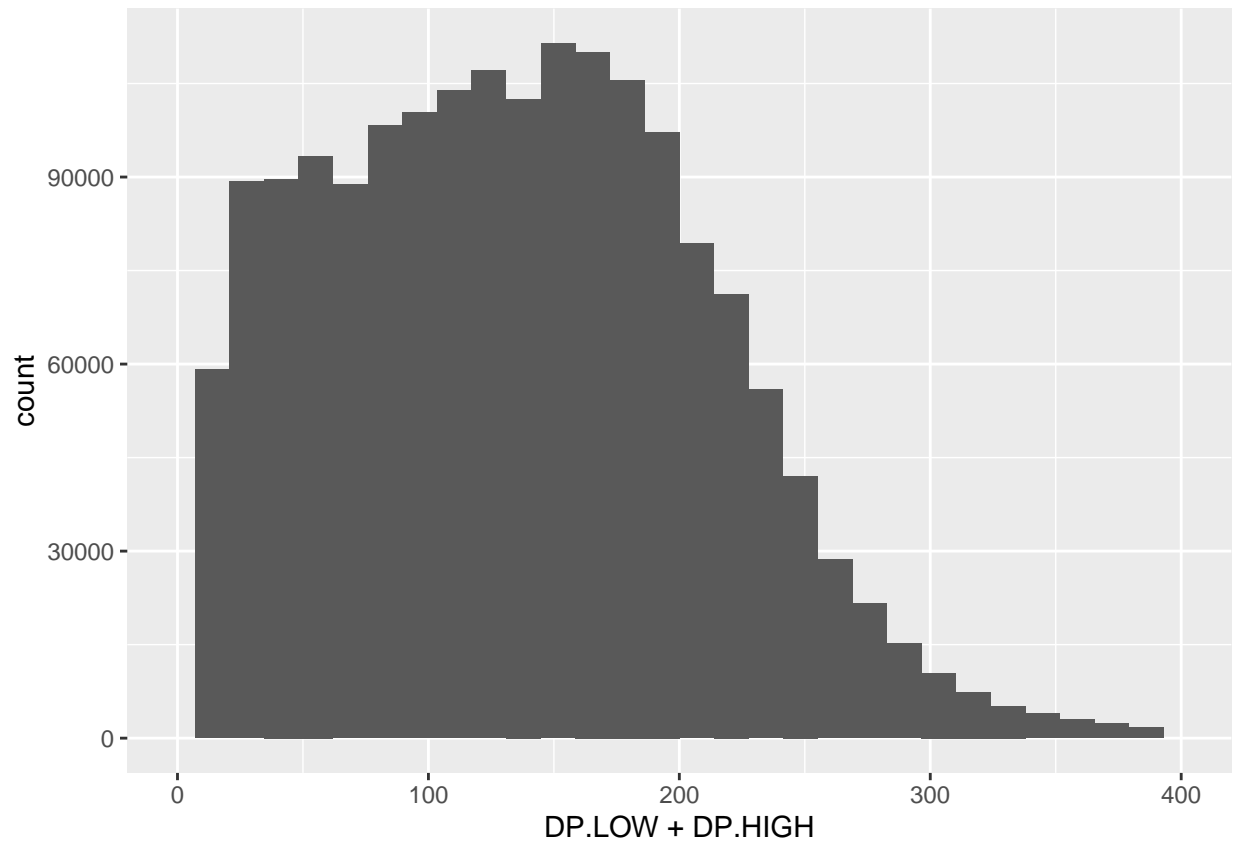
```
ggplot(data = df) + geom_histogram(aes(x = DP.HIGH)) + xlim(0,300)
```

``stat_bin()`` using ``bins = 30``. Pick better value with ``binwidth``.



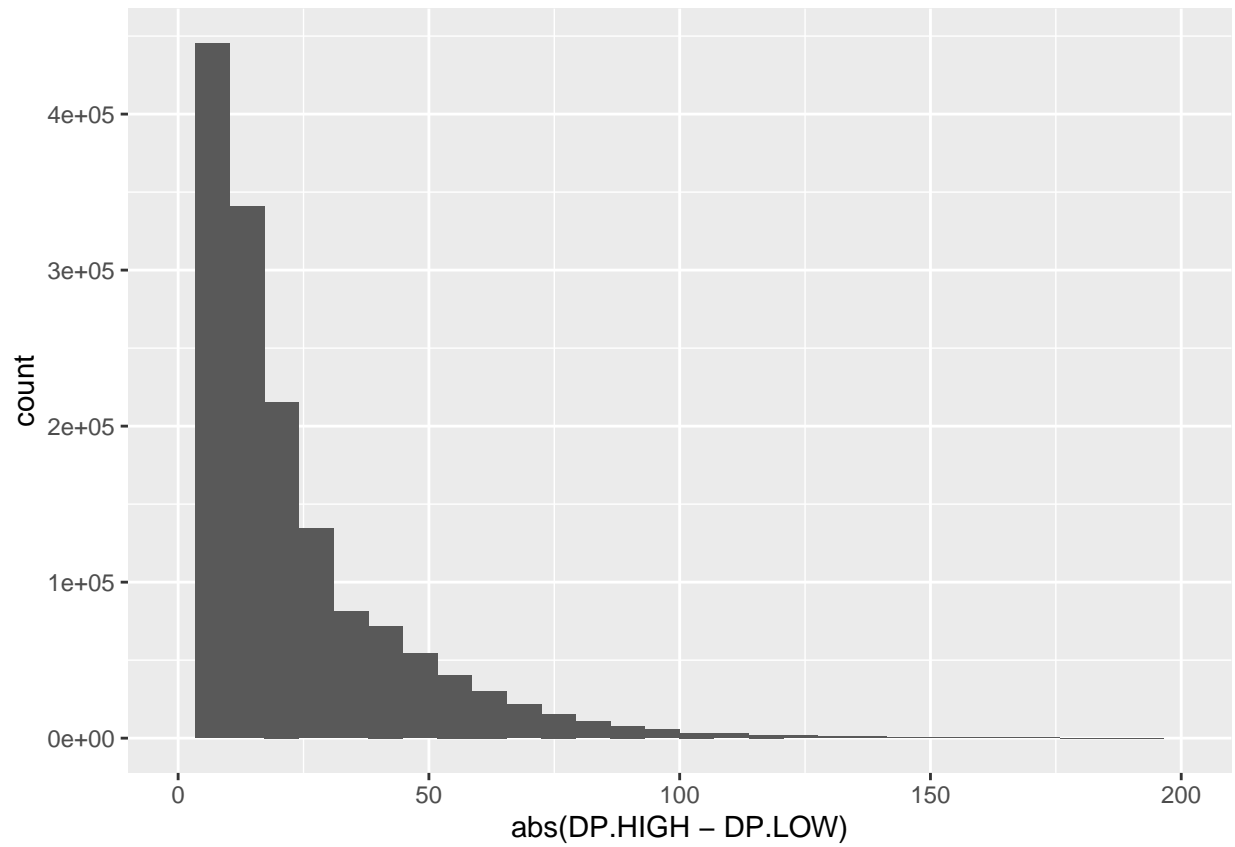
```
ggplot(data = df) + geom_histogram(aes(x = DP.LOW + DP.HIGH)) + xlim(0,400)
```

``stat_bin()`` using ``bins = 30``. Pick better value with ``binwidth``.



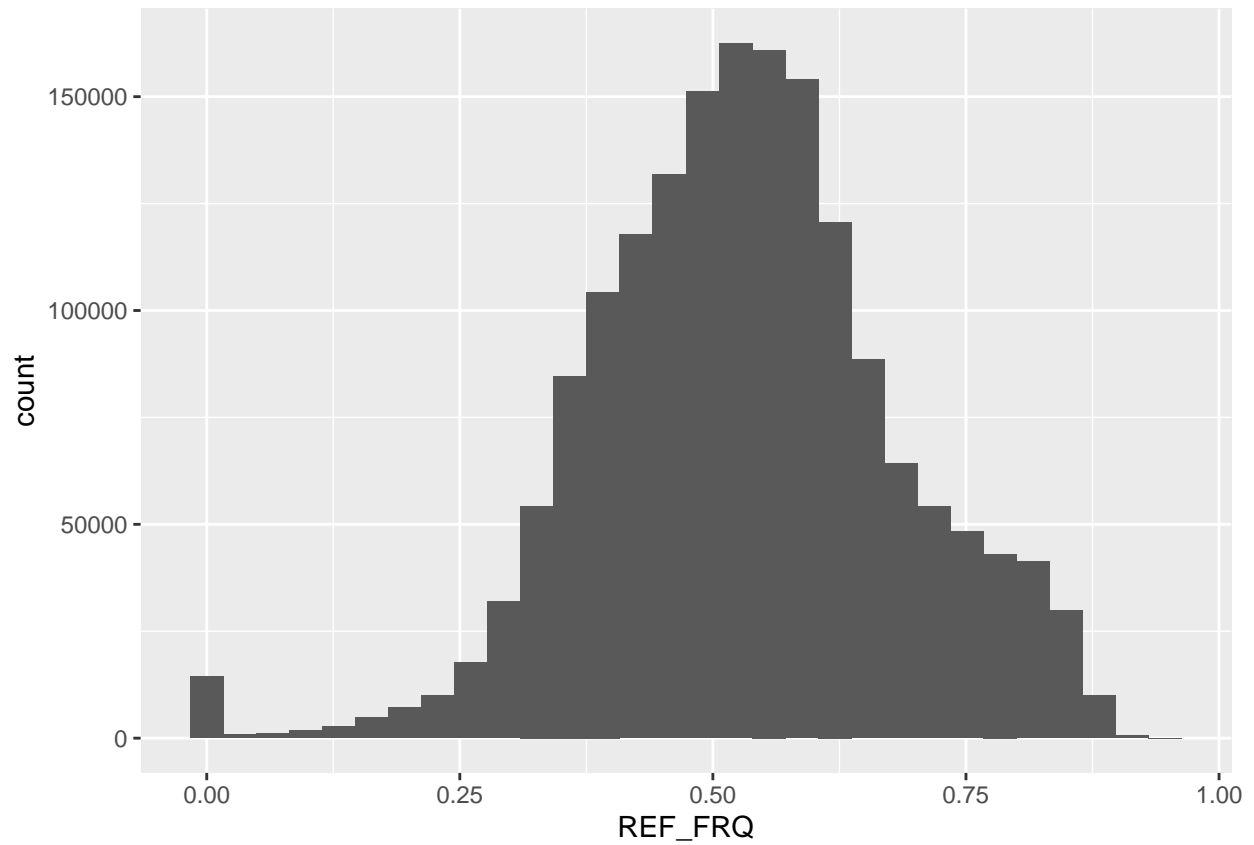
```
ggplot(data = df) + geom_histogram(aes(x = abs(DP.HIGH - DP.LOW))) + xlim(0, 200)
```

``stat_bin()`` using ``bins = 30``. Pick better value with ``binwidth``.



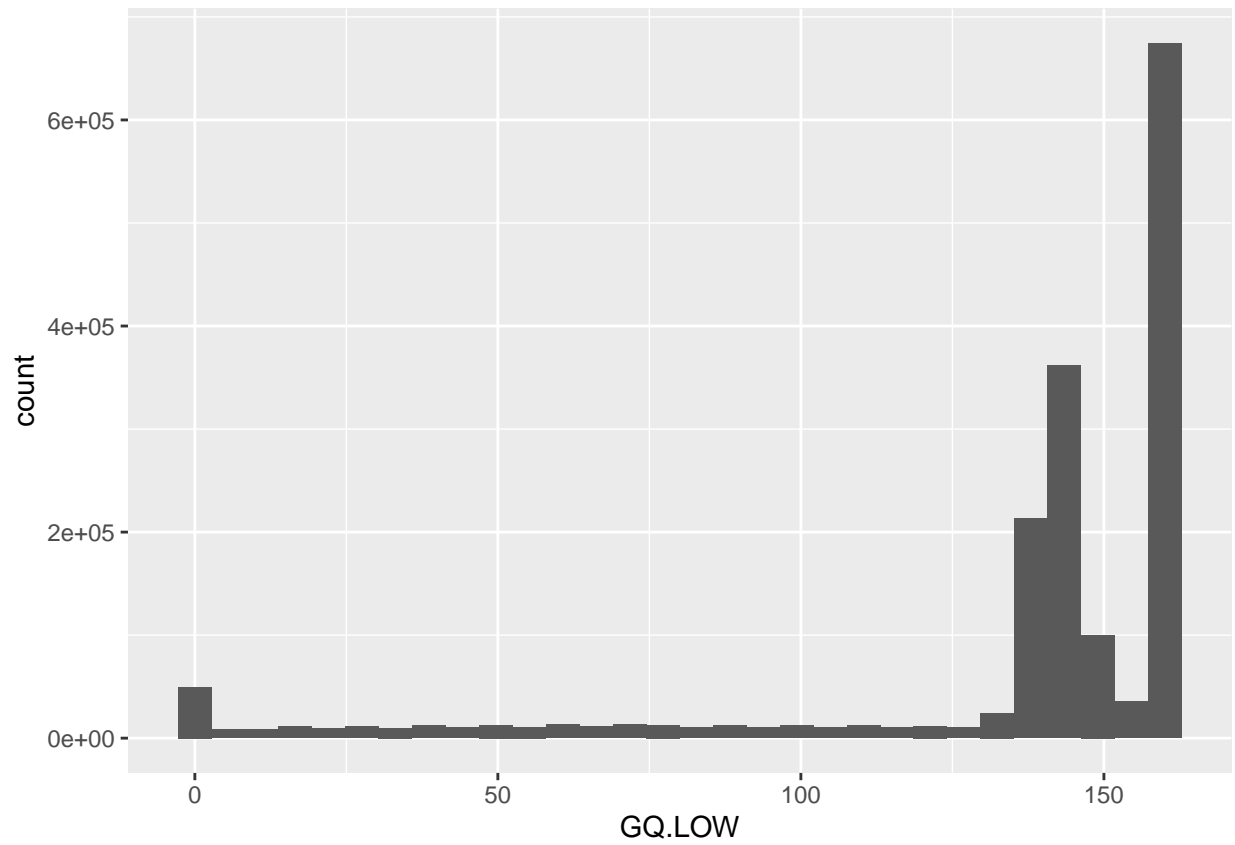
```
ggplot(data = df) + geom_histogram(aes(x = REF_FRQ))
```

``stat_bin()`` using ``bins = 30``. Pick better value with ``binwidth``.



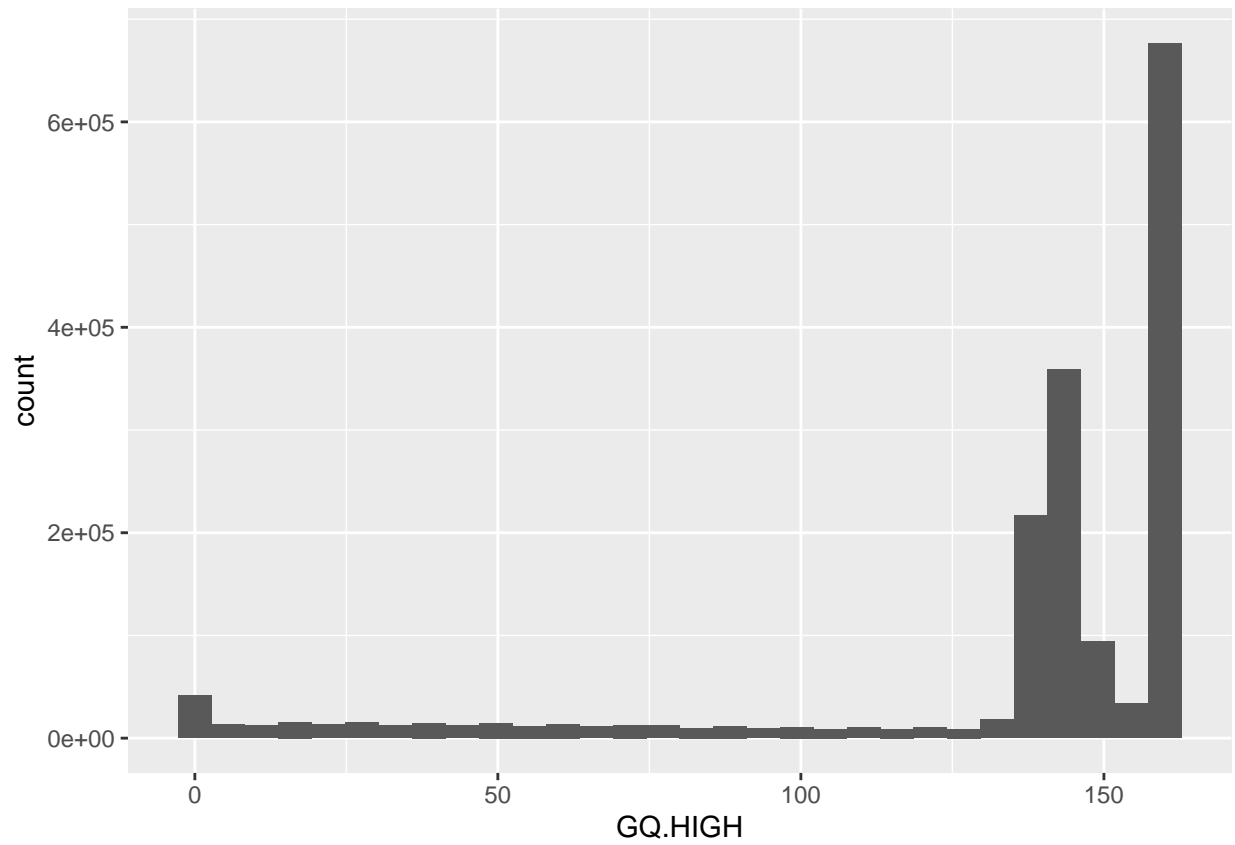
```
ggplot(data = df) + geom_histogram(aes(x = GQ.LOW))
```

``stat_bin()`` using ``bins = 30``. Pick better value with ``binwidth``.



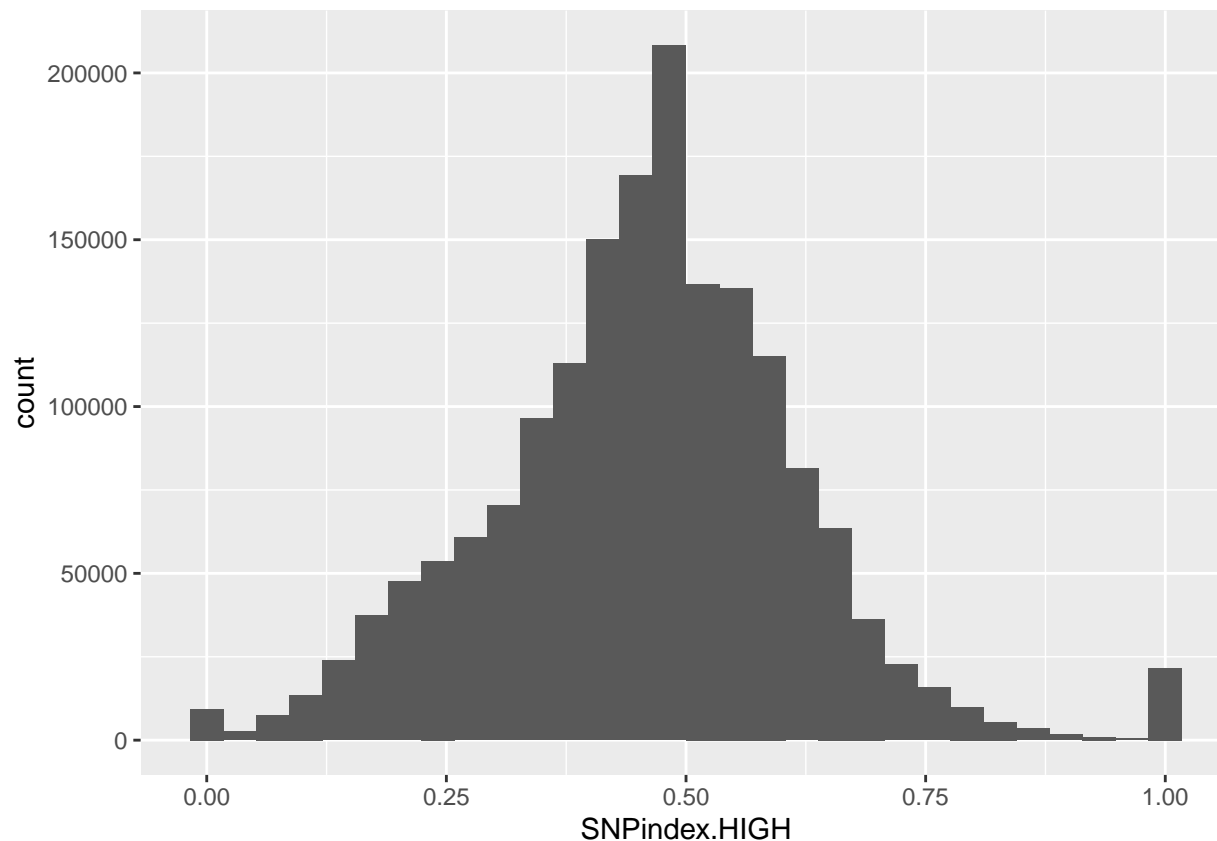
```
ggplot(data = df) + geom_histogram(aes(x = GQ.HIGH))
```

``stat_bin()`` using ``bins = 30``. Pick better value with ``binwidth``.



```
ggplot(data = df) + geom_histogram(aes(x = SNPindex.HIGH))
```

``stat_bin()`` using ``bins = 30``. Pick better value with ``binwidth``.



#Filter SNPs based on some criteria

```
df_filt <-
  QTLseqr::filterSNPs(
    SNPset = df,
    refAlleleFreq = 0.20,
    minTotalDepth = 100,
    maxTotalDepth = 400,
    depthDifference = 100,
    minSampleDepth = 40,
    minGQ = 99,
    verbose = TRUE
  )
```

Filtering by reference allele frequency: $0.2 \leq \text{REF_FRQ} \leq 0.8$

...Filtered 112443 SNPs

Filtering by total sample read depth: Total DP ≥ 100

...Filtered 537579 SNPs

Filtering by total sample read depth: Total DP ≤ 400

...Filtered 6975 SNPs

Filtering by per sample read depth: DP ≥ 40

...Filtered 8015 SNPs

Filtering by Genotype Quality: GQ ≥ 99

...Filtered 37905 SNPs

Filtering by difference between bulks ≤ 100

...Filtered 8746 SNPs

Original SNP number: 1714745, Filtered: 711663, Remaining: 1003082

#Run G' analysis

```
df_filt<-QTLseqr::runGprimeAnalysis(  
  SNPset = 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...

Calculating p-values...

#Run QTLseq analysis

```
df_filt <- QTLseqr::runQTLseqAnalysis(  
  SNPset = df_filt,  
  windowSize = 1e6,  
  popStruc = "F2",  
  bulkSize = c(385, 430),  
  replications = 10000,  
  intervals = c(95, 99)  
)
```

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: 40-199

Assuming bulks selected from F2 population, with 385 and 430 individuals per bulk.

Simulating 10000 SNPs with reads at each depth: 40-199

Keeping SNPs with ≥ 0.3 SNP-index in both simulated bulks

Joining, by = "tricubeDP"

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

Warning: Removed 2 rows containing missing values (geom_bar).

Removed 2 rows containing missing values (geom_bar).

```
QTLseqr::plotGprimeDist(SNPset = df_filt, outlierFilter = "deltaSNP", filterThreshold = 0.1)
```

Warning: Removed 2 rows containing missing values (geom_bar).

Removed 2 rows containing missing values (geom_bar).

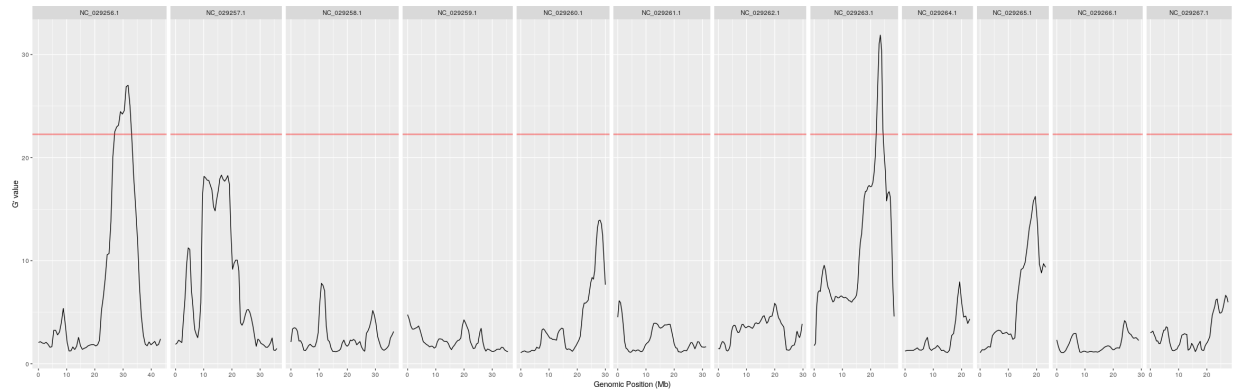
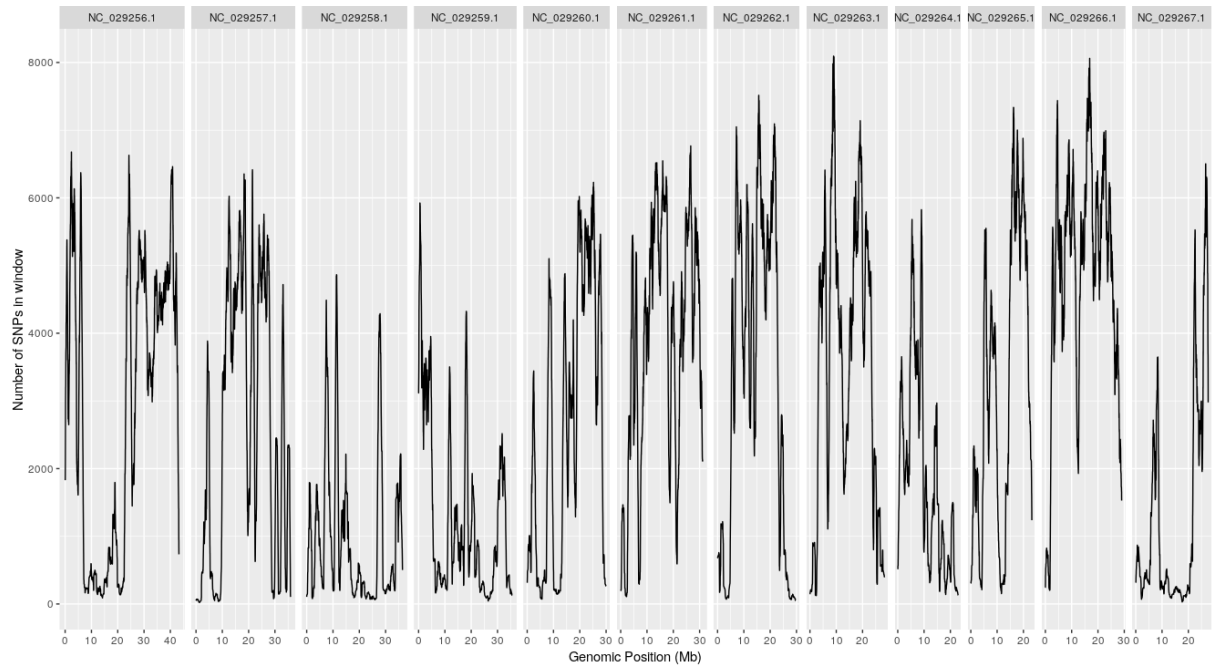
```
QTLseqr::plotQTLStats(SNPset = df_filt, var = "nSNPs")
```

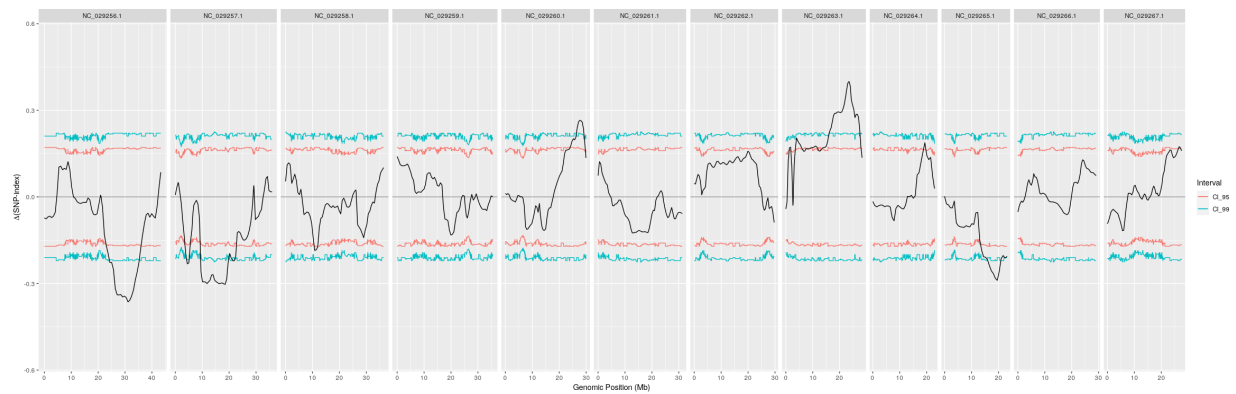
```
QTLseqr::plotQTLStats(SNPset = df_filt, var = "Gprime", plotThreshold = TRUE, q = 0.01)
```

```
QTLseqr::plotQTLStats(SNPset = df_filt, var = "deltaSNP", plotIntervals = TRUE)
```

```
QTLseqr::plotQTLStats(SNPset = df_filt, var = "negLog10Pval", plotThreshold = TRUE, q=0.01,
  subset = c("NC_029256.1", "NC_029257.1", "NC_029263.1", "NC_029265.1"))
```

```
QTLseqr::plotQTLStats(SNPset = df_filt, var = "Gprime", plotThreshold = TRUE, q=0.01,
  subset = c("NC_029256.1", "NC_029257.1", "NC_029263.1", "NC_029265.1"))
```





```
QTLseqr::getQTLTable(SNPset = df_filt, alpha = 0.01, export = TRUE, fileName = "my_BSA_QTL.csv")
```

```
##          CHROM qtl      start      end  length nSNPs avgSNPs_Mb peakDeltaSNP
## 1 NC_029256.1   1 26986793 33033424 6046631 26816      4435   -0.3636383
## 2 NC_029263.1   2 21980124 24413447 2433323  8704      3577    0.3999064
##   posPeakDeltaSNP avgDeltaSNP maxGprime posMaxGprime meanGprime sdGprime
## 1          31110319  -0.3456919  27.01544    31780847    24.38459  1.321401
## 2          23458666   0.3756486  31.87973    23458666    28.26922  3.219559
##          AUCaT      meanPval      meanQval
## 1 13606901 0.0002315438 0.009554921
## 2 14451588 0.0001277682 0.009555889
```