

# Package: qtl2pattern (via r-universe)

August 26, 2024

**Type** Package

**Title** Pattern Support for 'qtl2' Package

**Version** 1.2.2

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**Description** Routines in 'qtl2' to study allele patterns in quantitative trait loci (QTL) mapping over a chromosome. Useful in crosses with more than two alleles to identify how sets of alleles, genetically different strands at the same locus, have different response levels. Plots show profiles over a chromosome. Can handle multiple traits together. See <https://github.com/byandell/qtl2pattern>.

**Depends** R (>= 3.1.0)

**Imports** dplyr, tidyr, stringr, ggplot2, assertthat, qtl2, qtl2fst, fst, rlang, stats, graphics

**Suggests** knitr, rmarkdown, qtl2ggplot

**VignetteBuilder** knitr

**License** GPL-3

**URL** <https://github.com/byandell/qtl2pattern>

**Encoding** UTF-8

**RoxygenNote** 7.2.3

**Repository** <https://byandell.r-universe.dev>

**RemoteUrl** <https://github.com/byandell/qtl2pattern>

**RemoteRef** HEAD

**RemoteSha** 44e129ddc5bcd2b075a90bf01b883ed96200fea6

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---

allele1

*Allele plot for SNPs, alleles and allele pairs*

---

## Description

Create table of alleles for various model fits.

Plot alleles for haplotype, diplotype and top patterns and genome position.

## Usage

```
allele1(
  probD,
  phe_df = NULL,
  cov_mx = NULL,
  map = NULL,
  K_chr = NULL,
  patterns = NULL,
  alt = NULL,
  blups = FALSE,
  ...
)

ggplot_allele1(
  object,
  scan1_object = NULL,
  map = NULL,
  pos = NULL,
```

```

    trim = TRUE,
    legend.position = "none",
    ...
)

## S3 method for class 'allele1'
autoplot(object, ...)

```

### Arguments

|                 |   |
|-----------------|---|
| probD           | object of class <code>calc_genoprob</code>                      |
| phe_df          | data frame with one phenotype                                   |
| cov_mx          | covariate matrix  |
| map             | Genome map (required if <code>scan1_object</code> present).     |
| K_chr           | kinship matrix  |
| patterns        | data frame of pattern information                               |
| alt             | Haplotype allele letter(s) for alternative to reference.        |
| blups           | Create BLUPs if TRUE  |
| ...             | Other parameters ignored.                                       |
| object          | Object of class <code>allele1</code> .                          |
| scan1_object    | Optional object of class <code>scan1</code> to find peak.       |
| pos             | Genome position in Mbp (supercedes <code>scan1_object</code> ). |
| trim            | If TRUE, trim extreme alleles.                                  |
| legend.position | Legend position (default is "none").                            |

### Value

Table with allele effects across sources.  
 object of class `ggplot`

---

```
create_probs_query_func
```

*Create a function to query genotype probabilities*

---

### Description

Create a function that will connect to a database of genotype probability information and return a list with ‘probs’ object and a ‘map’ object.

### Usage

```
create_probs_query_func(dbfile, method_val = "fst", probdir_val = "genoprob")
```

**Arguments**

|             |   |
|-------------|---|
| dbfile      | Name of database file                                     |
| method_val  | either "fst" or "calc" for type of genotype probabilities |
| probdir_val | name of probability directory (default "genoprob")        |

**Details**

Note that this function assumes that `probdir_val` has a file with the physical map with positions in Mbp and other files with genotype probabilities. See [read\\_probs](#) for details on how probabilities are read. See [create\\_variant\\_query\\_func](#) for original idea.

**Value**

Function with six arguments, 'chr', 'start', 'end', 'allele', 'method' and 'probdir'. It returns a list with 'probs' and 'map' objects spanning the region specified by the first three arguments. The 'probs' element should be either a 'calc\_genoprob' or 'fst\_genoprob' object (see [fst\\_genoprob](#)).

**Examples**

```
dirpath <- "https://raw.githubusercontent.com/rqt1/rtl2data/master/DOex"

create_qv <- function(dirpath) {
  # Download SNP info for DOex from web via RDS.
  # snpinfo is referenced internally in the created function.

  tmpfile <- tempfile()
  download.file(file.path(dirpath, "c2_snpinfo.rds"), tmpfile, quiet=TRUE)
  snpinfo <- readRDS(tmpfile)
  unlink(tmpfile)
  snpinfo <- dplyr::rename(snpinfo, pos = pos_Mbp)

  function(chr, start, end) {
    if(chr != "2") return(NULL)
    if(start < 96.5) start <- 96.5
    if(end > 98.5) end <- 98.5
    if(start >= end) return(NULL)
    dplyr::filter(snpinfo, .data$pos >= start, .data$pos <= end)
  }
}

query_variants <- create_qv(dirpath)

create_qg <- function(dirpath) {
  # Download Gene info for DOex from web via RDS
  # gene_tbl is referenced internally in the created function.

  tmpfile <- tempfile()
  download.file(file.path(dirpath, "c2_genes.rds"), tmpfile, quiet=TRUE)
  gene_tbl <- readRDS(tmpfile)
  unlink(tmpfile)
}
```

```

function(chr, start, end) {
  if(chr != "2") return(NULL)
  if(start < 96.5) start <- 96.5
  if(end > 98.5) end <- 98.5
  if(start >= end) return(NULL)
  dplyr::filter(gene_tbl, .data$end >= start, .data$start <= end)
}
}

query_genes <- create_qg(dirpath)

# Examples for probs require either FST or RDS storage of data.

```

---

|           |                                   |
|-----------|-----------------------------------|
| gene_exon | <i>Get exons for set of genes</i> |
|-----------|-----------------------------------|

---

### Description

Match up exon start,stop,strand with genes. Use `query_genes` to find features; see [create\\_gene\\_query\\_func](#).

Returns table of gene and its exons.

Uses [gene\\_exon](#) to plot genes, exons, mRNA with SNPs.

### Usage

```

gene_exon(
  top_snps_tbl,
  feature_tbl = query_genes(chr_id, range_Mbp[1], range_Mbp[2])
)

## S3 method for class 'gene_exon'
summary(object, gene_name = NULL, top_snps_tbl = NULL, extra = 0.005, ...)

## S3 method for class 'gene_exon'
subset(x, gene_val, ...)

ggplot_gene_exon(
  object,
  top_snps_tbl = NULL,
  plot_now = TRUE,
  genes = unique(object$gene),
  ...
)

## S3 method for class 'gene_exon'
autoplot(object, ...)

```

**Arguments**

|              |  |
|--------------|--|
| top_snps_tbl | table from <a href="#">top_snps</a>  |
| feature_tbl  | table of features from query_genes; see <a href="#">create_gene_query_func</a> |
| object       | Object of class gene_exon.   |
| gene_name    | name of gene as character string   |
| extra        | extra region beyond gene for SNPs (in Mbp)                                     |
| ...          | arguments passed along to <a href="#">gene_exon</a>                            |
| x            | Object of class gene_exon.   |
| gene_val     | Name of gene from object x.  |
| plot_now     | plot now if TRUE   |
| genes        | Names of genes in object   |

**Value**

tbl of exon and gene features  
tbl of summary  
list of ggplots (see [gene\\_exon](#))

**Author(s)**

Brian S Yandell, <brian.yandell@wisc.edu>

**Examples**

```
dirpath <- "https://raw.githubusercontent.com/rqtl/ql2data/master/D0ex"

# Read D0ex example cross from 'ql2data'
D0ex <- subset(ql2::read_cross2(file.path(dirpath, "D0ex.zip")), chr = "2")

# Download genotype probabilities
tmpfile <- tempfile()
download.file(file.path(dirpath, "D0ex_genoprobs_2.rds"), tmpfile, quiet=TRUE)
pr <- readRDS(tmpfile)
unlink(tmpfile)

# Download SNP info for D0ex from web and read as RDS.
tmpfile <- tempfile()
download.file(file.path(dirpath, "c2_snpinfo.rds"), tmpfile, quiet=TRUE)
snpinfo <- readRDS(tmpfile)
unlink(tmpfile)
snpinfo <- dplyr::rename(snpinfo, pos = pos_Mbp)

# Convert to SNP probabilities
snpinfo <- ql2::index_snps(D0ex$pmap, snpinfo)
snppr <- ql2::genoprob_to_snpprob(pr, snpinfo)
```

```
# Scan SNPs.
scan_snppr <- qt12::scan1(snppr, D0ex$pheno)

# Collect top SNPs
top_snps_tbl <- top_snps_pattern(scan_snppr, snpinfo)

# Download Gene info for D0ex from web via RDS
tmpfile <- tempfile()
download.file(file.path(dirpath, "c2_genes.rds"), tmpfile, quiet=TRUE)
gene_tbl <- readRDS(tmpfile)
unlink(tmpfile)

# Get Gene exon information.
out <- gene_exon(top_snps_tbl, gene_tbl)
summary(out, gene = out$gene[1])
```

---

genoprob\_to\_patternprob

*Collapse genoprob according to pattern*

---

## Description

Collapse genoprob according to pattern

## Usage

```
genoprob_to_patternprob(probs1, sdp, alleles = FALSE)
```

## Arguments

|         |   |
|---------|---|
| probs1  | object of class <a href="#">calc_genoprob</a> |
| sdp     | SNP distribution pattern                      |
| alleles | use allele string if TRUE                     |

## Value

object of class [calc\\_genoprob](#)

## Author(s)

Brian S Yandell, <brian.yandell@wisc.edu>

**Examples**

```

dirpath <- "https://raw.githubusercontent.com/rqtl/ql2data/master/D0ex"

# Read D0ex example cross from 'ql2data'
D0ex <- subset(ql2::read_cross2(file.path(dirpath, "D0ex.zip")), chr = "2")

# Download genotype probabilities
tmpfile <- tempfile()
download.file(file.path(dirpath, "D0ex_genoprobs_2.rds"), tmpfile, quiet=TRUE)
pr <- readRDS(tmpfile)
unlink(tmpfile)

# Convert genotype probabilities to pattern probabilities for pattern 1.
pattern_pr <- genoprob_to_patternprob(pr, 7, TRUE)

str(pr)
str(pattern_pr)

```

---

get.gene.locations      *Helper function to set gene locations on plot.*

---

**Description**

Figure out gene locations to make room for gene names. Written original by Dan Gatti 2013-02-13

**Usage**

```

get.gene.locations(
  locs,
  xlim,
  text_size = 3,
  str_rect = c("iW", "i"),
  n_rows = 10,
  plot_width = 6,
  ...
)

```

**Arguments**

|           |  |
|-----------|--|
| locs      | tbl of gene information  |
| xlim      | X axis limits  |
| text_size | size of text (default 3)   |
| str_rect  | character spacing on left and right of rectangles (default c("iW", "i")) |
| n_rows    | desired number of rows (default 10)                                      |



plot\_width      width of default plot window (in inches)  
...              additional parameters (not used)

**Value**

list object used by [ggplot\\_feature\\_tbl](#)

**Author(s)**

Brian S Yandell, <brian.yandell@wisc.edu> Daniel Gatti, <Dan.Gatti@jax.org>

**References**

<https://github.com/dmgatti/DOQTL/blob/master/R/gene.plot.R>

---

get\_feature\_snp      *Match features with SNPs*

---

**Description**

Find features that overlap with SNPs

**Usage**

```
get_feature_snp(snp_tbl, feature_tbl, extend = 0.005)
```

**Arguments**

snp\_tbl              tbl of SNPs from `assoc.map`  
feature\_tbl          tbl of feature information from [create\\_gene\\_query\\_func](#)  
extend                extend region for SNPs in Mbp (default 0.005)

**Value**

tbl of features covering SNPs

**Author(s)**

Brian S Yandell, <brian.yandell@wisc.edu>

---

|              |                              |
|--------------|------------------------------|
| get_gene_snp | <i>Match genes with SNPs</i> |
|--------------|------------------------------|

---

**Description**

Internal routine to find features that overlap with SNPs

**Usage**

```
get_gene_snp(  
  snp_tbl,  
  feature_tbl,  
  feature_snp = get_feature_snp(snp_tbl, feature_tbl, 0)  
)
```

**Arguments**

|             |   |
|-------------|---|
| snp_tbl     | tbl of SNPs from query_variants; see package <a href="#">create_variant_query_func</a>          |
| feature_tbl | tbl of feature information from query_genes; see package <a href="#">create_gene_query_func</a> |
| feature_snp | tbl of feature information from <a href="#">get_feature_snp</a>                                 |

**Value**

tbl of genes covering SNPs

**Author(s)**

Brian S Yandell, <brian.yandell@wisc.edu>

---

|                      |                                     |
|----------------------|-------------------------------------|
| ggplot_merge_feature | <i>Plot of merge_feature object</i> |
|----------------------|-------------------------------------|

---

**Description**

Merge all SNPs in small region with LOD peaks across multiple phenotype.

**Usage**

```
ggplot_merge_feature(object, pheno, plot_by = c("pattern", "consequence"), ...)  
  
## S3 method for class 'merge_feature'  
autoplot(object, ...)  
  
merge_feature(  
  top_snps_tbl,
```

```

    snpinfo,
    out_lmm_snps,
    drop = 1.5,
    dropchar = 0,
    exons = gene_exon(top_snps_tbl)
  )

## S3 method for class 'merge_feature'
summary(object, sum_type = c("SNP type", "pattern"), ...)

```

### Arguments

|              |   |
|--------------|---|
| object       | of class <code>merge_feature</code>                                   |
| pheno        | name of phenotype to be plotted                                       |
| plot_by      | element to plot by (one of <code>c("pattern", "consequence")</code> ) |
| ...          | other arguments not used  |
| top_snps_tbl | tbl from <a href="#">top_snps_pattern</a> or <a href="#">top_snps</a> |
| snpinfo      | SNP information table   |
| out_lmm_snps | tbl from <a href="#">scan1</a> on SNPs                                |
| drop         | include LOD scores within drop of max for each phenotype              |
| dropchar     | number of characters to drop on phenames                              |
| exons        | table from <a href="#">gene_exon</a>                                  |
| sum_type     | one of <code>c("SNP type", "pattern")</code>                          |

### Value

ggplot2 object  
 tbl with added information on genes and exons  
 table summary

### Author(s)

Brian S Yandell, <brian.yandell@wisc.edu>

### Examples

```

dirpath <- "https://raw.githubusercontent.com/rqtl/ql2data/master/D0ex"

# Read D0ex example cross from 'ql2data'
D0ex <- subset(ql2::read_cross2(file.path(dirpath, "D0ex.zip")), chr = "2")

# Download genotype probabilities
tmpfile <- tempfile()
download.file(file.path(dirpath, "D0ex_genoprobs_2.rds"), tmpfile, quiet=TRUE)
pr <- readRDS(tmpfile)
unlink(tmpfile)

```

```

# Download SNP info for D0ex from web and read as RDS.
tmpfile <- tempfile()
download.file(file.path(dirpath, "c2_snpinfo.rds"), tmpfile, quiet=TRUE)
snpinfo <- readRDS(tmpfile)
unlink(tmpfile)
snpinfo <- dplyr::rename(snpinfo, pos = pos_Mbp)

# Convert to SNP probabilities
snpinfo <- qt12::index_snps(D0ex$pmap, snpinfo)
snppr <- qt12::genoprob_to_snpprob(pr, snpinfo)

# Scan SNPs.
scan_snppr <- qt12::scan1(snppr, D0ex$pheno)

# Collect top SNPs
top_snps_tbl <- top_snps_pattern(scan_snppr, snpinfo)
summary(top_snps_tbl)

# Download Gene info for D0ex from web via RDS
tmpfile <- tempfile()
download.file(file.path(dirpath, "c2_genes.rds"), tmpfile, quiet=TRUE)
gene_tbl <- readRDS(tmpfile)
unlink(tmpfile)

out <- merge_feature(top_snps_tbl, snpinfo, scan_snppr, exons = gene_tbl)
summary(out, "pattern")

```

---

ggplot\_scan1pattern *Plot scan pattern using ggplot2*

---

### Description

Plot scan pattern using ggplot2

Genome scan by pattern set

### Usage

```

ggplot_scan1pattern(
  object,
  map,
  plot_type = c("lod", "coef", "coef_and_lod"),
  patterns = object$patterns$founders,
  columns = 1:3,
  min_lod = 3,
  lodcolumn = seq_along(patterns),
  facet = "pheno",

```

```

    ...
  )

## S3 method for class 'scan1pattern'
autoplot(object, ...)

scan1pattern(
  probs1,
  phe,
  K = NULL,
  covar = NULL,
  map,
  patterns,
  condense_patterns = TRUE,
  blups = FALSE,
  do_scans = TRUE
)

## S3 method for class 'scan1pattern'
summary(object, map, ...)

```

### Arguments

|                   |   |
|-------------------|---|
| object            | object of class <a href="#">scan1pattern</a>                                  |
| map               | genome map  |
| plot_type         | type of plot from <code>c("lod", "coef")</code>                               |
| patterns          | data frame of pattern information   |
| columns           | columns used for coef plot  |
| min_lod           | minimum LOD peak for contrast to be retained                                  |
| lodcolumn         | columns used for scan1 plot (default all patterns)                            |
| facet             | Plot facets if multiple phenotypes and patterns provided (default = "pheno"). |
| ...               | additional parameters passed on to other methods                              |
| probs1            | object of class <a href="#">calc_genoprob</a>                                 |
| phe               | data frame with one phenotype   |
| K                 | kinship matrix  |
| covar             | covariate matrix  |
| condense_patterns | remove snp_action from contrasts if TRUE                                      |
| blups             | Create BLUPs if TRUE  |
| do_scans          | Do scans if TRUE.   |

### Value

object of class [ggplot](#)

List containing:

- patterns Data frame of summary for top patterns (column founders has pattern)
- dip\_set Diplotype sets for contrasts
- group Group for each founder pattern
- scan Object of class `scan1`.
- coef Object of class `listof_scan1coef`. See package `'qtl2ggplot'`.

### Author(s)

Brian S Yandell, <brian.yandell@wisc.edu>

### Examples

```
dirpath <- "https://raw.githubusercontent.com/rqtl/qtl2data/master/D0ex"

# Read D0ex example cross from 'qtl2data'
D0ex <- subset(qtl2::read_cross2(file.path(dirpath, "D0ex.zip")), chr = "2")

# Download genotype probabilities
tmpfile <- tempfile()
download.file(file.path(dirpath, "D0ex_genoprobs_2.rds"), tmpfile, quiet=TRUE)
pr <- readRDS(tmpfile)
unlink(tmpfile)

# Download SNP info for D0ex from web and read as RDS.
tmpfile <- tempfile()
download.file(file.path(dirpath, "c2_snpinfo.rds"), tmpfile, quiet=TRUE)
snpinfo <- readRDS(tmpfile)
unlink(tmpfile)
snpinfo <- dplyr::rename(snpinfo, pos = pos_Mbp)

# Convert to SNP probabilities
snpinfo <- qtl2::index_snps(D0ex$pmap, snpinfo)
snppr <- qtl2::genoprob_to_snpprob(pr, snpinfo)

# Scan SNPs
scan_snppr <- qtl2::scan1(snppr, D0ex$pheno)
top_snps_tbl <- top_snps_pattern(scan_snppr, snpinfo)

# Summarize to find top patterns
patterns <- dplyr::arrange(summary(top_snps_tbl), dplyr::desc(max_lod))

# Scan using patterns.
scan_pat <- scan1pattern(pr, D0ex$pheno, map = D0ex$pmap, patterns = patterns)

# Summary of scan1pattern.
summary(scan_pat, D0ex$pmap)
```

---

|                |                                      |
|----------------|--------------------------------------|
| pattern_diplos | <i>Extract pattern of diplotypes</i> |
|----------------|--------------------------------------|

---

**Description**

Extract pattern of diplotypes

Extract pattern of haplotypes

**Usage**

```
pattern_diplos(sdp, haplos, diplos, cont = NULL)
```

```
pattern_haplos(sdp, haplos)
```

**Arguments**

sdp                vector of sdp from [top\\_snps\\_pattern](#)

haplos            vector of haplotype names

diplos            vector of diplotype names

cont              vector of types of contrasts (NULL or from `c("add", "dom", "b6r", "b6d")`)

**Value**

matrix of diplotype patterns

matrix of haplotype patterns

**Author(s)**

Brian S Yandell, <[brian.yandell@wisc.edu](mailto:brian.yandell@wisc.edu)>

---

|               |  |
|---------------|--|
| pattern_label | <i>Turn genotype probabilities into labels</i> |
|---------------|--|

---

**Description**

Turn genotype probabilities into labels

**Usage**

```
pattern_label(genos, allele = TRUE)
```

```
pattern_sdp(label, sdp = NULL, geno_names = sort(unique(label)))
```

**Arguments**

|            |   |
|------------|---|
| genos      | matrix of genotype probabilities at locus           |
| allele     | Driver has alleles if TRUE, otherwise allele pairs. |
| label      | character string from <a href="#">pattern_label</a> |
| sdp        | SNP distribution pattern for plot colors            |
| geno_names | unique genotype names (alleles or allele pairs)     |

**Value**

character vector of genotype names.

---

|           |  |
|-----------|--|
| read_fast | <i>Read fast database with possible rownames</i> |
|-----------|--|

---

**Description**

Read fast database with format fst. Use first column of database (must be named 'ind') as rownames if desired. R/qtl2 routines assume data frames have rownames to use to align individuals.

**Usage**

```
read_fast(datapath, columns = NULL, rownames = TRUE)
```

**Arguments**

|          |   |
|----------|---|
| datapath | character string path to database                               |
| columns  | names or indexes for columns to be extracted                    |
| rownames | use first column of rownames if TRUE (can supply column number) |

**Value**

extracted data frame with appropriate rows and columns.

**See Also**

[read\\_fst](#)



---

|            |   |
|------------|---|
| read_probs | <i>Read genotype probability object from file</i> |
|------------|---|

---

**Description**

Read object from file stored according to method.

**Usage**

```
read_probs(  
  chr = NULL,  
  start_val = NULL,  
  end_val = NULL,  
  datapath,  
  allele = TRUE,  
  method,  
  probdir = "genoprob"  
)
```

**Arguments**

|                    |   |
|--------------------|---|
| chr                | vector of chromosome identifiers  |
| start_val, end_val | start and end values in Mbp   |
| datapath           | name of folder with Derived Data  |
| allele             | read haplotype allele probabilities (if TRUE) or diplotype allele-pair probabilities (if FALSE) |
| method             | method of genoprob storage  |
| probdir            | genotype probability directory (default "genoprob")   |

**Value**

list with probs = large object of class `calc_genoprob` and map = physical map for selected chr

**Author(s)**

Brian S Yandell, <brian.yandell@wisc.edu>

---

|                |                               |
|----------------|-------------------------------|
| sdp_to_pattern | <i>Convert sdp to pattern</i> |
|----------------|-------------------------------|

---

**Description**

Convert strain distribution pattern (sdp) to letter pattern.

**Usage**

```
sdp_to_pattern(sdp, haplos, symmetric = TRUE)
```

```
sdp_to_logical(sdp, haplos, symmetric = TRUE)
```

**Arguments**

|           |  |
|-----------|--|
| sdp       | vector of sdp values                   |
| haplos    | letter codes for haplotypes (required) |
| symmetric | make patterns symmetric if TRUE        |

**Value**

vector of letter patterns

**Author(s)**

Brian S Yandell, <brian.yandell@wisc.edu>

**Examples**

```
dirpath <- "https://raw.githubusercontent.com/rqt1/qt12data/master/D0ex"

# Download SNP info for D0ex from web and read as RDS.
tmpfile <- tempfile()
download.file(file.path(dirpath, "c2_snpinfo.rds"), tmpfile, quiet=TRUE)
snpinfo <- readRDS(tmpfile)
unlink(tmpfile)
snpinfo <- dplyr::rename(snpinfo, pos = pos_Mbp)

# Extract strain distribution pattern.
sdp <- snpinfo$sdp
# Find out how many alleles.
nallele <- ceiling(log2(max(sdp)))
out <- sdp_to_pattern(sdp, LETTERS[seq_len(nallele)])
# Show most frequent patterns.
head(rev(sort(c(table(out)))))
```

---

|                |                                |
|----------------|--------------------------------|
| snpinfo_to_map | <i>Convert SNP info to map</i> |
|----------------|--------------------------------|

---

**Description**

Convert SNP info to map

**Usage**

```
snpinfo_to_map(snpinfo)
```

**Arguments**

|         |  |
|---------|--|
| snpinfo | Data frame with SNP information with the following columns (the last three are generally derived from with <a href="#">index_snps</a> ): <ul style="list-style-type: none"> <li>• chr - Character string or factor with chromosome</li> <li>• pos - Position (in same units as in the "map" attribute in genoprobs.</li> <li>• sdp - Strain distribution pattern: an integer, between 1 and <math>2^n - 2</math> where <math>n</math> is the number of strains, whose binary encoding indicates the founder genotypes</li> <li>• snp - Character string with SNP identifier (if missing, the rownames are used).</li> <li>• index - Indices that indicate equivalent groups of SNPs.</li> <li>• intervals - Indexes that indicate which marker intervals the SNPs reside.</li> <li>• on_map - Indicate whether SNP coincides with a marker in the genoprobs</li> </ul> |
|---------|--|

**Value**

map as list of vectors of marker positions.

---

|                  |  |
|------------------|--|
| snpprob_collapse | <i>Collapse genoprobs according to pattern</i> |
|------------------|--|

---

**Description**

Collapse genoprobs according to pattern

**Usage**

```
snpprob_collapse(
  snpprobs,
  action = c("additive", "add+dom", "non-add", "recessive", "dominant", "basic")
)
```

**Arguments**

snpprobs        object of class `calc_genoprob`  
action         SNP gene action type

**Value**

object of class `calc_genoprob`

**Author(s)**

Brian S Yandell, <brian.yandell@wisc.edu>

**Examples**

```
dirpath <- "https://raw.githubusercontent.com/rqtl/ql2data/master/D0ex"

# Read D0ex example cross from 'ql2data'
D0ex <- subset(ql2::read_cross2(file.path(dirpath, "D0ex.zip")), chr = "2")

# Download genotype probabilities
tmpfile <- tempfile()
download.file(file.path(dirpath, "D0ex_genoprob_2.rds"), tmpfile, quiet=TRUE)
pr <- readRDS(tmpfile)
unlink(tmpfile)

# Download SNP info for D0ex from web and read as RDS.
tmpfile <- tempfile()
download.file(file.path(dirpath, "c2_snpinfo.rds"), tmpfile, quiet=TRUE)
snpinfo <- readRDS(tmpfile)
unlink(tmpfile)
snpinfo <- dplyr::rename(snpinfo, pos = pos_Mbp)

# Convert to snp probabilities
snpinfo <- ql2::index_snps(D0ex$pmap, snpinfo)
snppr <- ql2::genoprob_to_snpprob(pr, snpinfo)

dim(snppr[[1]])
dim(snpprob_collapse(snppr, "additive")[[1]])
```

---

summary.feature\_snp    *Summary of features with SNP information*

---

**Description**

Summary of features with SNP information

**Usage**

```
## S3 method for class 'feature_snp'
summary(object, ...)
```

**Arguments**

```
object      tbl of feature information from get\_feature\_snp
...         additional parameters ignored
```

**Value**

tbl of feature summaries by type

**Author(s)**

Brian S Yandell, <brian.yandell@wisc.edu>

---

summary.feature\_tbl    *Summary of features*

---

**Description**

Show count min and max of features by type

Plot genes as rectangles followed by names. Stagger genes for easy reading. Written original by Dan Gatti 2013-02-13

**Usage**

```
## S3 method for class 'feature_tbl'
summary(object, major = TRUE, ...)
```

```
## S3 method for class 'feature_tbl'
subset(x, start_val = 0, stop_val = max(x$stop), ...)
```

```
ggplot_feature_tbl(
  object,
  rect_col = "grey70",
  strand_col = c(`-` = "#1b9e77", `+` = "#d95f02"),
  type_col = c(gene = "black", pseudogene = "#1b9e77", other = "#d95f02"),
  text_size = 3,
  xlim = NULL,
  snp_pos = top_snps_tbl$pos,
  snp_lod = top_snps_tbl$lod,
  top_snps_tbl = NULL,
  snp_col = "grey70",
  extend = 0.005,
```

```

    ...
  )

  ## S3 method for class 'feature_tbl'
  autoplot(object, ...)

```

### Arguments

|                     |  |
|---------------------|--|
| object              | tbl of gene information from query_variants; see <a href="#">create_variant_query_func</a> |
| major               | if TRUE (default), only summarize genes and exons  |
| ...                 | additional arguments (not used)  |
| x                   | tbl of feature information from <a href="#">get_feature_snp</a>                            |
| start_val, stop_val | start and stop positions for subset  |
| rect_col            | fill color of rectangle (default "grey70")   |
| strand_col          | edge color of rectangle by strand from object (default -="blue", += "red"; none if NULL)   |
| type_col            | color of type from object (default "black" for gene, "blue" for pseudogene; none if NULL)  |
| text_size           | size of text (default 3)   |
| xlim                | horizontal axis limits (default is range of features)                                      |
| snp_pos             | position of SNPs in bp if used (default NULL)  |
| snp_lod             | LOD of SNPs (for color plotting)   |
| top_snps_tbl        | table from <a href="#">top_snps</a>  |
| snp_col             | color of SNP vertical lines (default "grey70")   |
| extend              | extend region for SNPs in bp (default 0.005)   |

### Value

tbl of feature summaries by type  
tbl of feature summaries by type  
data frame of gene information (invisible)

### Author(s)

Brian S Yandell, <brian.yandell@wisc.edu>  
Brian S Yandell, <brian.yandell@wisc.edu> Daniel Gatti, <Dan.Gatti@jax.org>

### References

<https://github.com/dmgatti/DOQTL/blob/master/R/gene.plot.R>

---

|                  |  |
|------------------|--|
| summary.gene_snp | <i>Summary of genes overlapping SNPs</i> |
|------------------|--|

---

**Description**

Summary of genes overlapping SNPs

**Usage**

```
## S3 method for class 'gene_snp'
summary(object, ...)
```

**Arguments**

|        |   |
|--------|---|
| object | tbl of feature information from <a href="#">get_feature_snp</a> |
| ...    | additional parameters ignored                                   |

**Value**

tbl of feature summaries by type

**Author(s)**

Brian S Yandell, <brian.yandell@wisc.edu>

---

|                  |   |
|------------------|---|
| top_snps_pattern | <i>Top SNPs organized by allele pattern</i> |
|------------------|---|

---

**Description**

Separate fine mapping scans by allele pattern.

**Usage**

```
top_snps_pattern(
  scan1_output,
  snpinfo,
  drop = 1.5,
  show_all_snps = TRUE,
  haplos
)

## S3 method for class 'top_snps_pattern'
summary(object, sum_type = c("range", "best", "peak"), ...)

## S3 method for class 'top_snps_pattern'
subset(x, start_val = 0, end_val = max(x$pos), pheno = NULL, ...)
```

**Arguments**

|                    |   |
|--------------------|---|
| scan1_output       | output of linear mixed model for phename (see <a href="#">scan1</a> )   |
| snpinfo            | Data frame with SNP information with the following columns (the last three are generally derived from with <a href="#">index_snps</a> ): <ul style="list-style-type: none"> <li>• chr - Character string or factor with chromosome</li> <li>• pos - Position (in same units as in the "map" attribute in genoprobs.</li> <li>• sdp - Strain distribution pattern: an integer, between 1 and <math>2^n - 2</math> where <math>n</math> is the number of strains, whose binary encoding indicates the founder genotypes</li> <li>• snp_id - Character string with SNP identifier (if missing, the rownames are used).</li> <li>• index - Indices that indicate equivalent groups of SNPs.</li> <li>• intervals - Indexes that indicate which marker intervals the SNPs reside.</li> <li>• on_map - Indicate whether SNP coincides with a marker in the genoprobs</li> </ul> |
| drop               | include all SNPs within drop of max LOD (default 1.5)   |
| show_all_snps      | show all SNPs if TRUE   |
| haplos             | optional argument identify codes for haplotypes   |
| object             | object of class top_snps_tbl  |
| sum_type           | type of summary (one of "range","best")   |
| ...                | additional parameters ignored   |
| x                  | tbl of feature information from <a href="#">get_feature_snp</a>   |
| start_val, end_val | start and end positions for subset  |
| pheno              | phenotype name(s) for subset  |

**Value**

table of top\_snps at maximum lod for pattern  
table summary  
subset of x

**Author(s)**

Brian S Yandell, <brian.yandell@wisc.edu>

**Examples**

```
dirpath <- "https://raw.githubusercontent.com/rqt1/qt12data/master/DOex"

# Read DOex example cross from 'qt12data'
DOex <- subset(qt12::read_cross2(file.path(dirpath, "DOex.zip")), chr = "2")

# Download genotype probabilities
```



```
tmpfile <- tempfile()
download.file(file.path(dirpath, "D0ex_genoprobs_2.rds"), tmpfile, quiet=TRUE)
pr <- readRDS(tmpfile)
unlink(tmpfile)

# Download SNP info for D0ex from web and read as RDS.
tmpfile <- tempfile()
download.file(file.path(dirpath, "c2_snpinfo.rds"), tmpfile, quiet=TRUE)
snpinfo <- readRDS(tmpfile)
unlink(tmpfile)
snpinfo <- dplyr::rename(snpinfo, pos = pos_Mbp)

# Convert to SNP probabilities
snpinfo <- qtl2::index_snps(D0ex$pmap, snpinfo)
snppr <- qtl2::genoprob_to_snpprob(pr, snpinfo)

# Scan SNPs.
scan_snppr <- qtl2::scan1(snppr, D0ex$pheno)

# Collect top SNPs
top_snps_tbl <- top_snps_pattern(scan_snppr, snpinfo)
summary(top_snps_tbl)
```

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