

# Package ‘qtl2convert’

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**Title** Convert Data among QTL Mapping Packages

**Description** Functions to convert data structures among the 'qtl2', 'qtl', and 'DOQTL' packages for mapping quantitative trait loci (QTL).

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stats

**Suggests** testthat, devtools, roxygen2

**License** GPL-3

**URL** <https://kbroman.org/qtl2/>, <https://github.com/rqtl/qtl2convert>

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## Contents

|                                |   |
|--------------------------------|---|
| cbind_smother . . . . .        | 2 |
| count_unique_geno . . . . .    | 3 |
| cross2_do_to_genail8 . . . . . | 3 |
| cross2_ril_to_genril . . . . . | 4 |

|                               |           |
|-------------------------------|-----------|
| encode_geno . . . . .         | 5         |
| find_consensus_geno . . . . . | 6         |
| find_unique_geno . . . . .    | 6         |
| map_df_to_list . . . . .      | 7         |
| map_list_to_df . . . . .      | 8         |
| probs_doqtl_to_qtl2 . . . . . | 9         |
| probs_qtl2_to_array . . . . . | 10        |
| probs_qtl2_to_doqtl . . . . . | 10        |
| probs_qtl_to_qtl2 . . . . .   | 11        |
| scan_qtl2_to_qtl . . . . .    | 12        |
| scan_qtl_to_qtl2 . . . . .    | 13        |
| write2csv . . . . .           | 13        |
| <b>Index</b>                  | <b>16</b> |

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|               |                                                                                    |
|---------------|------------------------------------------------------------------------------------|
| cbind_smother | <i>Combine matrices by columns, replacing matching ones and adding unique ones</i> |
|---------------|------------------------------------------------------------------------------------|

---

**Description**

This is like `base::cbind()` but if a column in the second matrix has the same name as a column in the first matrix, the column in the first matrix is deleted and that in the second matrix is used in its place.

**Usage**

```
cbind_smother(mat1, mat2)
```

**Arguments**

- mat1            A matrix
- mat2            Another matrix, with the same number of rows as mat.

**Value**

The two matrices combined by columns, but columns in the first matrix that also appear in the second matrix are deleted and replaced by those in the second matrix. Uses the row names to align the rows in the two matrices, and to expand them as needed.

**Examples**

```
df1 <- data.frame(x=c(1,2,3,NA,4), y=c(5,8,9,10,11), row.names=c("A", "B", "C", "D", "E"))
df2 <- data.frame(z=c(7,8,0,9,10), y=c(6,NA,NA,9,10), row.names=c("A", "B", "F", "C", "D"))
df1n2 <- cbind_smother(df1, df2)
```

---

|                   |                                                                     |
|-------------------|---------------------------------------------------------------------|
| count_unique_geno | <i>Count the unique genotypes for each row of a genotype matrix</i> |
|-------------------|---------------------------------------------------------------------|

---

**Description**

For genotype data (markers x individuals) on a set of individuals, count the unique genotypes for each marker

**Usage**

```
count_unique_geno(genotypes, na.strings = c("N", "H", "NA", ""), cores = 1)
```

**Arguments**

|            |                                                                                                                                                                                                                                        |
|------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| genotypes  | Matrix of genotypes (markers x individuals)                                                                                                                                                                                            |
| na.strings | Genotypes to be considered as missing values.                                                                                                                                                                                          |
| cores      | Number of CPU cores to use, for parallel calculations. (If 0, use <a href="#">parallel::detectCores()</a> .)<br>Alternatively, this can be links to a set of cluster sockets, as produced by <a href="#">parallel::makeCluster()</a> . |

**Value**

Vector of counts of unique genotypes.

**See Also**

[find\\_unique\\_geno\(\)](#)

**Examples**

```
g <- rbind(c("NA", "A", "A", "A", "T"),
           c("NA", "NA", "NA", "A", "A"),
           c("A", "A", "T", "G", "G"),
           c("C", "C", "G", "G", "NA"))
counts <- count_unique_geno(g)
```

---

|                      |                                                 |
|----------------------|-------------------------------------------------|
| cross2_do_to_genail8 | <i>Convert cross2 object from do to genail8</i> |
|----------------------|-------------------------------------------------|

---

**Description**

Convert a cross2 object from cross type "do" to cross type "genail8".

**Usage**

```
cross2_do_to_genail8(cross)
```

```
cross2_do_to_genail(cross)
```

**Arguments**

cross                      Object of class "cross2", as produced by `qt12::read_cross2()`.

**Value**

The input object cross with cross type changed to class "genril8" and the cross information revised to match.

**Examples**

```
## Not run:
file <- paste0("https://raw.githubusercontent.com/rqt1/",
               "qt12data/master/D0ex/D0ex.zip")
D0ex <- read_cross2(file)

D0ex_genril <- cross2_do_to_genril8(D0ex)

## End(Not run)
```

---

cross2\_ril\_to\_genril    *Convert cross2 object from ril to genril*

---

**Description**

Convert a cross2 object from cross type "risibn" to cross type "genriln".

**Usage**

```
cross2_ril_to_genril(cross)
```

**Arguments**

cross                      Object of class "cross2", as produced by `qt12::read_cross2()`.

**Value**

The input object cross with cross type changed to class "genriln" for some value of n, and the cross information revised to match.

**Examples**

```
## Not run:
file <- paste0("https://raw.githubusercontent.com/rqt1/",
               "qt12data/master/CC/cc.zip")
cc <- read_cross2(file)

cc_genril <- cross2_ril_to_genril(cc)

## End(Not run)
```

```
## Not run:
file <- paste0("https://raw.githubusercontent.com/rqtl/",
               "qtl2data/master/ArabMAGIC/arabmagic_tair9.zip")
arab <- read_cross2(file)

arab_genril <- cross2_ril_to_genril(arab)

## End(Not run)
```

---

encode\_geno

---

*Encode a matrix of genotypes using a set of allele codes*


---

## Description

Encode a matrix of genotypes using a set of allele codes.

## Usage

```
encode_geno(
  geno,
  allele_codes,
  output_codes = c("-", "A", "H", "B"),
  cores = 1
)
```

## Arguments

|              |                                                                                                                                                                                                                                        |
|--------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| geno         | Character matrix of genotypes (rows as markers, columns as individuals)                                                                                                                                                                |
| allele_codes | Two-column matrix of alleles (rows as markers)                                                                                                                                                                                         |
| output_codes | Vector of length four, with missing, AA, AB, BB codes                                                                                                                                                                                  |
| cores        | Number of CPU cores to use, for parallel calculations. (If 0, use <a href="#">parallel::detectCores()</a> .)<br>Alternatively, this can be links to a set of cluster sockets, as produced by <a href="#">parallel::makeCluster()</a> . |

## Value

Matrix of same dimensions as geno, but with values in output\_codes.

## See Also

[find\\_consensus\\_geno\(\)](#), [find\\_unique\\_geno\(\)](#)

## Examples

```
geno <- rbind(c("C", "G", "C", "GG", "CG"),
             c("A", "A", "AT", "TA", "TT"),
             c("T", "G", NA, "GT", "TT"))
codes <- rbind(c("C", "G"), c("A", "T"), c("T", "G"))
geno_encoded <- encode_geno(geno, codes)
```

---

|                     |                                                                      |
|---------------------|----------------------------------------------------------------------|
| find_consensus_geno | <i>Find the consensus genotype for each row of a genotype matrix</i> |
|---------------------|----------------------------------------------------------------------|

---

### Description

For genotype data (markers x individuals) on a set of individuals from a single inbred line, find the consensus genotype at each marker.

### Usage

```
find_consensus_geno(genotypes, na.strings = c("N", "H", "NA", ""), cores = 1)
```

### Arguments

|            |                                                                                                                                                                                                                                        |
|------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| genotypes  | Matrix of genotypes (markers x individuals)                                                                                                                                                                                            |
| na.strings | Genotypes to be considered as missing values.                                                                                                                                                                                          |
| cores      | Number of CPU cores to use, for parallel calculations. (If 0, use <a href="#">parallel::detectCores()</a> .)<br>Alternatively, this can be links to a set of cluster sockets, as produced by <a href="#">parallel::makeCluster()</a> . |

### Value

Vector of consensus genotypes, one value per row of genotypes

### See Also

[find\\_unique\\_geno\(\)](#), [encode\\_geno\(\)](#)

### Examples

```
g <- rbind(c("NA", "N", "A", "A", "T", "G", NA, "H"),
           c("C", "C", "G", "G", "A", NA, NA, NA),
           rep(NA, 8),
           c("C", "C", "G", "G", "G", "C", "G", "G"))
consensus <- find_consensus_geno(g)
```

---

|                  |                                                                    |
|------------------|--------------------------------------------------------------------|
| find_unique_geno | <i>Find the unique genotypes for each row of a genotype matrix</i> |
|------------------|--------------------------------------------------------------------|

---

### Description

For genotype data (markers x individuals) on a set of individuals, find the unique genotypes for each marker, provided that there are exactly two. (If more than two or fewer than two, return NAs.)

### Usage

```
find_unique_geno(genotypes, na.strings = c("N", "H", "NA", ""), cores = 1)
```

**Arguments**

|            |                                                                                                                                                                                                                                  |
|------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| genotypes  | Matrix of genotypes (markers x individuals)                                                                                                                                                                                      |
| na.strings | Genotypes to be considered as missing values.                                                                                                                                                                                    |
| cores      | Number of CPU cores to use, for parallel calculations. (If 0, use <code>parallel::detectCores()</code> .)<br>Alternatively, this can be links to a set of cluster sockets, as produced by <code>parallel::makeCluster()</code> . |

**Value**

Matrix with two columns. Each row corresponds to a marker, and has the two unique genotypes, or NAs (if >2 or <2 unique genotypes).

**See Also**

`count_unique_geno()`, `encode_geno()`

**Examples**

```
g <- rbind(c("NA", "A", "A", "A", "T"),
           c("NA", "NA", "NA", "A", "A"),
           c("A", "A", "T", "G", "G"),
           c("C", "C", "G", "G", "NA"))
ug <- find_unique_geno(g)
```

---

|                |                                      |
|----------------|--------------------------------------|
| map_df_to_list | <i>Marker map data frame to list</i> |
|----------------|--------------------------------------|

---

**Description**

Convert a marker map organized as data frame to a list

**Usage**

```
map_df_to_list(
  map,
  chr_column = "chr",
  pos_column = "cM",
  marker_column = "marker",
  Xchr = c("X", "X")
)
```

**Arguments**

|            |                                                               |
|------------|---------------------------------------------------------------|
| map        | Data frame with marker map                                    |
| chr_column | Name of the column in map that contains the chromosome IDs.   |
| pos_column | Name of the column in map that contains the marker positions. |

marker\_column    Name of the column in map that contains the marker names. If NULL, use the row names.

Xchr             Vector of character strings indicating the name or names of the X chromosome. If NULL, assume there's no X chromosome.

**Value**

A list of vectors of marker positions, one component per chromosome

**See Also**

[map\\_list\\_to\\_df\(\)](#)

**Examples**

```
map <- data.frame(chr=c(1,1,1, 2,2,2, "X","X"),
                  pos=c(0,5,10, 0,8,16, 5,20),
                  marker=c("D1M1","D1M2","D1M3", "D2M1","D2M2","D2M3", "DXM1","DXM2"))
map_list <- map_df_to_list(map, pos_column="pos")
```

---

|                |                                      |
|----------------|--------------------------------------|
| map_list_to_df | <i>Marker map list to data frame</i> |
|----------------|--------------------------------------|

---

**Description**

Convert a marker map organized as a list to a data frame

**Usage**

```
map_list_to_df(
  map_list,
  chr_column = "chr",
  pos_column = "pos",
  marker_column = "marker"
)
```

**Arguments**

map\_list            List of vectors containing marker positions

chr\_column          Name of the chromosome column in the output

pos\_column          Name of the position column in the output

marker\_column      Name of the marker column in the output. If NULL, just put them as row names.

**Value**

A data frame with the marker positions.

**See Also**[map\\_df\\_to\\_list\(\)](#)**Examples**

```
library(qtl2)
iron <- read_cross2(system.file("extdata", "iron.zip", package="qtl2"))
iron_map <- map_list_to_df(iron$gmap)
```

---

probs\_doqtl\_to\_qtl2     *Convert DOQTL genotype probabilities to R/qtl2 format*

---

**Description**

Convert DOQTL genotype probabilities to R/qtl2 format

**Usage**

```
probs_doqtl_to_qtl2(
  probs,
  map,
  is_female = NULL,
  chr_column = "chr",
  pos_column = "cM",
  marker_column = "marker"
)
```

**Arguments**

|               |                                                                                                                                                    |
|---------------|----------------------------------------------------------------------------------------------------------------------------------------------------|
| probs         | 3d array of genotype probabilities as calculated from DOQTL (individuals x genotypes x positions)                                                  |
| map           | Data frame with marker map                                                                                                                         |
| is_female     | Optional logical vector indicating which individuals are female. Names should contain the individual identifiers, matching the row names in probs. |
| chr_column    | Name of the column in map that contains the chromosome IDs.                                                                                        |
| pos_column    | Name of the column in map that contains the marker positions.                                                                                      |
| marker_column | Name of the column in map that contains the marker names. If NULL, use the row names.                                                              |

**Details**

We assume that the X chromosome is labeled "X" (must be upper-case) and that any other chromosomes are autosomes. We assume that the genotypes are labeled using the 8 letters A-H.

If the probabilities are for the full 36 states and the X chromosome is included but `is_female` is not provided, we'll guess which individuals are females based on their genotype probabilities. (If the average, across loci, of the sum of the heterozygote probabilities is small, we'll assume it's a female.)

**Value**

An object of the form produced by `qtl2::calc_genoprob()`.

---

|                     |                                                            |
|---------------------|------------------------------------------------------------|
| probs_qtl2_to_array | <i>Convert R/qtl2 genotype probabilities to a 3d array</i> |
|---------------------|------------------------------------------------------------|

---

**Description**

Convert R/qtl2 genotype probabilities to a 3d array

**Usage**

```
probs_qtl2_to_array(probs)
```

**Arguments**

|       |                                                                                                                                 |
|-------|---------------------------------------------------------------------------------------------------------------------------------|
| probs | A "calc_genoprob" object (a list of 3d arrays of genotype probabilities), as calculated by <code>qtl2::calc_genoprob()</code> . |
|-------|---------------------------------------------------------------------------------------------------------------------------------|

**Details**

We convert just the autosomal genotype probabilities, because they should all have the same number of genotypes (columns). The main application of this is for identifying possible sample mix-ups among batches of genotype probabilities (e.g., using the [R/lineup2](#) package), and for this the autosomal genotype probabilities should be sufficient.

**Value**

A single three-dimensional array, with just the autosomal genotype probabilities.

---

|                     |                                                              |
|---------------------|--------------------------------------------------------------|
| probs_qtl2_to_doqtl | <i>Convert R/qtl2 genotype probabilities to DOQTL format</i> |
|---------------------|--------------------------------------------------------------|

---

**Description**

Convert R/qtl2 genotype probabilities to DOQTL format

**Usage**

```
probs_qtl2_to_doqtl(probs)
```

**Arguments**

|       |                                                                                                                                 |
|-------|---------------------------------------------------------------------------------------------------------------------------------|
| probs | A "calc_genoprob" object (a list of 3d arrays of genotype probabilities), as calculated by <code>qtl2::calc_genoprob()</code> . |
|-------|---------------------------------------------------------------------------------------------------------------------------------|

## Details

If the arrays in probs all have 8 columns, they're assumed to be allele dosages and we paste them all together into one big array.

Otherwise, it should be that the autosomes all have 36 columns the X chromosome has 44. In this case, the male hemizygotes on the X are placed where the female homozygotes are, and then we reorder the genotypes into alphabetical order.

## Value

A single three-dimensional array, for use with **DOQTL**.

---

|                   |                                                              |
|-------------------|--------------------------------------------------------------|
| probs_qtl_to_qtl2 | <i>Convert R/qtl genotype probabilities to R/qtl2 format</i> |
|-------------------|--------------------------------------------------------------|

---

## Description

Convert R/qtl genotype probabilities to R/qtl2 format

## Usage

```
probs_qtl_to_qtl2(cross)
```

## Arguments

|       |                                                                                                                                                                          |
|-------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| cross | An R/qtl "cross" object (see <a href="#">qtl::read.cross()</a> for details.) Must contain genotype probabilities as calculated by <a href="#">qtl::calc.genoprob()</a> . |
|-------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

## Value

A list with two components:

- "probs" - the genotype probabilities in the form produced by [qtl2::calc\\_genoprob\(\)](#)
- "map" - Map of marker/pseudomarker positions (a list of vectors of positions)

## Examples

```
library(qtl)
data(hyper)
hyper <- calc.genoprob(hyper, step=1, error.prob=0.002)
result <- probs_qtl_to_qtl2(hyper)
pr <- result$probs
map <- result$map
```

---

|                  |                                                    |
|------------------|----------------------------------------------------|
| scan_qtl2_to_qtl | <i>Convert scan1 results to the scanone format</i> |
|------------------|----------------------------------------------------|

---

## Description

Convert the results of `qtl2::scan1()` to the form used by the R/qtl function `qtl::scanone()`.

## Usage

```
scan_qtl2_to_qtl(scan1_output, map)
```

## Arguments

|              |                                                                     |
|--------------|---------------------------------------------------------------------|
| scan1_output | Matrix of LOD scores, as calculated by <code>qtl2::scan1()</code> . |
| map          | Map of markers/pseudomarkers (as a list of vectors).                |

## Value

A data frame with class "scanone", containing chromosome and position columns followed by the LOD scores in scan1\_output.

## See Also

`scan_qtl2_to_qtl()`

## Examples

```
library(qtl2)
iron <- read_cross2(system.file("extdata", "iron.zip", package="qtl2"))
map <- insert_pseudomarkers(iron$gmap, step=1)
probs <- calc_genoprob(iron, map, error_prob=0.002)
pheno <- iron$pheno
covar <- match(iron$covar$sex, c("f", "m")) # make numeric
names(covar) <- rownames(iron$covar)
Xcovar <- get_x_covar(iron)
out <- scan1(probs, pheno, addcovar=covar, Xcovar=Xcovar)

out_rev <- scan_qtl2_to_qtl(out, map)
```

---

|                  |                                                             |
|------------------|-------------------------------------------------------------|
| scan_qtl_to_qtl2 | <i>Convert R/qtl scanone results to R/qtl2 scan1 format</i> |
|------------------|-------------------------------------------------------------|

---

**Description**

Convert the results of R/qtl `qtl::scanone()` to the form used by the R/qtl2 `qtl2::scan1()`.

**Usage**

```
scan_qtl_to_qtl2(scanone_output)
```

**Arguments**

scanone\_output Data frame as output by the R/qtl function `qtl::scanone()`.

**Value**

List with two objects: the LOD scores in `qtl2::scan1()` format, and the map (as a list of marker/pseudomarker positions).

**See Also**

`scan_qtl_to_qtl2()`

**Examples**

```
library(qtl)
data(hyper)
hyper <- calc.genoprob(hyper, step=1, error.prob=0.002)
out <- scanone(hyper)
out2 <- scan_qtl_to_qtl2(out)
```

---

|           |                                         |
|-----------|-----------------------------------------|
| write2csv | <i>Write a data frame to a CSV file</i> |
|-----------|-----------------------------------------|

---

**Description**

Write a data frame to a CSV file in a special form, with info about the number of rows and columns.

**Usage**

```
write2csv(
  df,
  filename,
  comment = "",
  sep = ",",
  comment.char = "#",
  row.names = NULL,
  overwrite = FALSE
)
```

**Arguments**

|                           |                                                                                                                                                                                                        |
|---------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>df</code>           | A data frame (or matrix)                                                                                                                                                                               |
| <code>filename</code>     | File name to write                                                                                                                                                                                     |
| <code>comment</code>      | Comment to place on the first line                                                                                                                                                                     |
| <code>sep</code>          | Field separator                                                                                                                                                                                        |
| <code>comment.char</code> | Character to use to initiate the comment lines                                                                                                                                                         |
| <code>row.names</code>    | If NA or NULL (the default), row names are not included in the output file. Otherwise, the row names are included as the first column of the output, and this is taken to be the name for that column. |
| <code>overwrite</code>    | If TRUE, overwrite file if it exists. If FALSE (the default) and the file exists, stop with an error.                                                                                                  |

**Details**

If the file already exists, the function will refuse to write over it.

The file will include comments at the top, using # as a comment character, including the number of rows (not including the header) and the number of columns.

By default, row names are not included. But with the option `row.names` provided as a character string, they are added as an initial column, with the value of this argument defining the name for that column. If a column with that name already exists, the function halts with an error.

**Value**

None.

**Examples**

```
nr <- 10
nc <- 5
x <- data.frame(id=paste0("ind", 1:nr),
                matrix(rnorm(nr*nc), ncol=nc))
colnames(x)[1:nc + 1] <- paste0("col", 1:nc)

testfile <- file.path(tempdir(), "tmpfile.csv")
write2csv(x, testfile, "A file created by write2csv")
```

```
# Remove the file, to clean up temporary directory  
unlink(testfile)
```

# Index

`base::cbind()`, [2](#)

`cbind_smother`, [2](#)

`count_unique_geno`, [3](#)

`count_unique_geno()`, [7](#)

`cross2_do_to_genail`  
    (`cross2_do_to_genail8`), [3](#)

`cross2_do_to_genail8`, [3](#)

`cross2_ril_to_genril`, [4](#)

`encode_geno`, [5](#)

`encode_geno()`, [6](#), [7](#)

`find_consensus_geno`, [6](#)

`find_consensus_geno()`, [5](#)

`find_unique_geno`, [6](#)

`find_unique_geno()`, [3](#), [5](#), [6](#)

`map_df_to_list`, [7](#)

`map_df_to_list()`, [9](#)

`map_list_to_df`, [8](#)

`map_list_to_df()`, [8](#)

`parallel::detectCores()`, [3](#), [5–7](#)

`parallel::makeCluster()`, [3](#), [5–7](#)

`probs_doqtl_to_qtl2`, [9](#)

`probs_qtl2_to_array`, [10](#)

`probs_qtl2_to_doqtl`, [10](#)

`probs_qtl_to_qtl2`, [11](#)

`qtl2::calc_genoprob()`, [10](#), [11](#)

`qtl2::read_cross2()`, [4](#)

`qtl2::scan1()`, [12](#), [13](#)

`qtl::calc.genoprob()`, [11](#)

`qtl::read.cross()`, [11](#)

`qtl::scanone()`, [12](#), [13](#)

`scan_qtl2_to_qtl`, [12](#)

`scan_qtl2_to_qtl()`, [12](#)

`scan_qtl_to_qtl2`, [13](#)

`scan_qtl_to_qtl2()`, [13](#)

`write2csv`, [13](#)