# Package 'mmconvert'

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<b>Date</b> 2023-04-25
Title Mouse Map Converter
<b>Description</b> Convert mouse genome positions between the build 39 physical map and the genetic map of Cox et al. (2009) <doi:10.1534 genetics.109.105486="">.</doi:10.1534>
<b>Author</b> Karl W Broman [aut, cre] ( <a href="https://orcid.org/0000-0002-4914-6671">https://orcid.org/0000-0002-4914-6671</a> )
Maintainer Karl W Broman broman@wisc.edu>
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<pre>URL https://github.com/rqtl/mmconvert</pre>
<pre>BugReports https://github.com/rqtl/mmconvert/issues</pre>
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R topics documented:
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coxmap	Mouse	genetic	map	based	on	Cox	et	al.	
	Rhrefhttps://doi.org/10.1534/genetics.109.105486doi:10.1534/								
	genetics.109.105486, revised for mouse genome build 39.								

### **Description**

A data frame with rows being markers and six columns: marker name, chromosome, sex-averaged cM position, female cM position, male cM position, and build 39 basepair position.

### **Details**

Genetic maps were re-estimated after reordering markers according to their position in mouse genome build 39. See https://github.com/kbroman/CoxMapV3. Markers were shifted so that 0 cM corresponds to 3 Mbp, using the chromosome- and sex-specific recombination rate. The maps are smoothed slightly to avoid intervals with 0 recombination.

### **Source**

```
https://github.com/kbroman/CoxMapV3
```

### **Examples**

data(coxmap)

cross2\_to\_grcm39

Convert a cross2 object to use mouse build GRCm39

### **Description**

Convert a cross2 object (with genotypes from one of the MUGA arrays) to use mouse build GRCm39 and the revised Cox map positions, revising marker order and omitting markers that are not found.

### Usage

```
cross2_to_grcm39(cross, array = c("guess", "gm", "mm", "mini", "muga"))
```

### **Arguments**

cross	Object of class "cross2", as produced by qt12::read_cross2(). Must have markers from just one of the MUGA arrays.
array	Character string indicating which of the MUGA arrays was used ("gm" for Giga-MUGA "mm" for MegaMUGA "mini" for MiniMUGA or "muga" for the orig-

MUGA, "mm" for MegaMUGA, "mini" for MiniMUGA, or "muga" for the original MUGA), or "guess" (the default) to pick the array with the most matching

marker names.

grcm39\_chrlen 3

### Value

The input cross object with markers subset to those in build GRCm39 and with pmap and gmap replaced with the GRCm39 physical map and revised Cox genetic map, respectively.

### See Also

**MUGAmaps** 

### **Examples**

grcm39\_chrlen

Mouse chromosome lengths in basepairs for build GRCm39

### **Description**

A vector of mouse chromosome lengths in basepairs for mouse genome build GRCm39.

### Source

```
https://www.ncbi.nlm.nih.gov/data-hub/genome/GCF_000001635.27/
```

### **Examples**

```
data(grcm39_chrlen)
```

mmconvert

Convert mouse genome positions

### Description

Convert mouse genome positions between the build 39 physical map and the Cox genetic map.

### Usage

```
mmconvert(
  positions,
  input_type = c("bp", "Mbp", "ave_cM", "female_cM", "male_cM")
)
```

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### **Arguments**

positions

A set of positions, in one of three possible formats

- a vector of character strings with like "chr:position", with names being marker names, e.g., c(rs13482072="14:6738536", rs13482231="14:67215850", gnf14.117.278="14:121955310").
- a list of marker positions, each list being positions on a given chromosome, e.g., list("14"=c(rs13482072=6738536, rs13482231=67215850, gnf14.117.278=121955310)
- a data frame with columns chromosome, position, and marker, e.g. data.frame(chr=c(14,14,14), pos=c(6738536, 67215850, 121955310), marker=c("rs13482072", "rs13482231", "gnf14.117.278")).

input\_type

Character string indicating the type of positions provided ("bp", "Mbp", "ave\_cM", "female\_cM", or "male\_cM")

#### **Details**

We use linear interpolation using the Cox map positions in the object coxmap. For positions outside the range of the markers on the Cox map, we extrapolate using the overall recombination rate.

### Value

A data frame with the interpolated positions, with seven columns: marker, chromosome, sex-averaged cM, female cM, male cM, basepairs, and mega-basepairs. The rows are sorted by genomic position.

### See Also

coxmap

### Examples

MUGAmaps 5

MUGAmaps	Array annotation information for the mouse MUGA arrays in mouse
	genome build 39.

### Description

A list of four data frames with annotation information for the four MUGA arrays, GigaMUGA ("gm"), MegaMUGA ("mm"), MiniMUGA ("mini") and the original MUGA ("muga"). Each has columns marker, chromosome, build 39 basepair position, and sex-averaged cM position (in Cox Map v3).

### **Details**

SNP probes for the MUGA arrays were blasted against mouse genome build GRCm39 and locations interpolated using revised Cox maps. See <a href="https://github.com/kbroman/MUGAarrays">https://github.com/kbroman/MUGAarrays</a> for the array annotations and <a href="https://github.com/kbroman/CoxMapV3">https://github.com/kbroman/CoxMapV3</a> for the genetic maps. Note that for the genetic map locations, markers were shifted so that 0 cM corresponds to 3 Mbp, using the chromosome- and sex-specific recombination rate. Moreover, the Cox map positions were smoothed slightly to avoid regions with 0 recombination.

### **Source**

https://github.com/kbroman/MUGAarrays

### **Examples**

data(MUGAmaps)

## **Index**

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