## Package 'karlen'

April 30, 2024

```
Title Real-Time PCR Data Sets by Karlen et al. (2007)
```

Version 0.0.2

```
Description Real-time quantitative polymerase chain reaction (qPCR) data sets by Karlen et al. (2007) <doi:10.1186/1471-2105-8-131>. Provides one single tabular tidy data set in long format, encompassing 32 dilution series, for seven PCR targets and four biological samples. The targeted amplicons are within the murine genes: Cav1, Ccn2, Eln, Fn1, Rpl27, Hspg2, and Serpine1, respectively. Dilution series: scheme 1 (Cav1, Eln, Hspg2, Serpine1): 1-fold, 10-fold, 50-fold, and 100-fold; scheme 2 (Ccn2, Rpl27, Fn1): 1-fold, 10-fold, 50-fold, 100-fold and 1000-fold. For each concentration there are five replicates, except for the 1000-fold concentration, where only two replicates were performed. Each amplification curve is 40 cycles long. Original raw data file is Additional file 2 from "Statistical significance of quantitative PCR" by Y. Karlen, A. McNair, S. Perseguers, C. Mazza, and N. Mermod (2007) <a href="https://static-content.springer.com/esm/art%3A10.1186%2F1471-2105-8-131/MediaObjects/12859_2006_1503_MOESM2_ESM.ZIP>">https://static-content.springer.com/esm/art%3A10.1186%2F1471-2105-8-131/MediaObjects/12859_2006_1503_MOESM2_ESM.ZIP>">https://static-content.springer.com/esm/art%3A10.1186%2F1471-2105-8-131/MediaObjects/12859_2006_1503_MOESM2_ESM.ZIP>">https://static-content.springer.com/esm/art%3A10.1186%2F1471-2105-8-131/MediaObjects/12859_2006_1503_MOESM2_ESM.ZIP>">https://static-content.springer.com/esm/art%3A10.1186%2F1471-2105-8-131/MediaObjects/12859_2006_1503_MOESM2_ESM.ZIP>">https://static-content.springer.com/esm/art%3A10.1186%2F1471-2105-8-131/MediaObjects/12859_2006_1503_MOESM2_ESM.ZIP>">https://static-content.springer.com/esm/art%3A10.1186%2F1471-2105-8-131/MediaObjects/12859_2006_1503_MOESM2_ESM.ZIP>">https://static-content.springer.com/esm/art%3A10.1186%2F1471-2105-8-131/MediaObjects/12859_2006_1503_MOESM2_ESM.ZIP>">https://static-content.springer.com/esm/art%3A10.1186%2F1471-2105-8-131/MediaObjects/12859_2006_1503_MOESM2_ESM.ZIP>">https://static-content.springer.com/esm/art%3A10.1186%2F1471-2105-
```

License CC BY 4.0 Encoding UTF-8 RoxygenNote 7.3.1 Imports tibble Depends R (>= 2.10)

LazyData true

URL https://rmagno.eu/karlen/, https://github.com/ramiromagno/karlen

BugReports https://github.com/ramiromagno/karlen/issues

NeedsCompilation no

Author Ramiro Magno [aut, cre] (<a href="https://orcid.org/0000-0001-5226-3441">https://orcid.org/0000-0001-5226-3441</a>),
Pattern Institute [cph, fnd]

Maintainer Ramiro Magno <rmagno@pattern.institute>

Repository CRAN

**Date/Publication** 2024-04-30 08:30:07 UTC

2 karlen

## **R** topics documented:

	karlen	 				 . 2
Index						4
karle	en	qPCR d	lata sets by	Karlen et al.	(2007)	

#### **Description**

One single tabular tidy data set in long format, encompassing 32 dilution series, for seven PCR targets and four biological samples. The targeted amplicons are within the murine genes: Cav1, Ccn2, Eln, Fn1, Rpl27, Hspg2, and Serpine1, respectively. Dilution series: scheme 1 (Cav1, Eln, Hspg2, Serpine1): 1-fold, 10-fold, 50-fold, and 100-fold; scheme 2 (Ccn2, Rpl27, Fn1): 1-fold, 10-fold, 50-fold, 100-fold and 1000-fold. For each concentration there are five replicates, except for the 1000-fold concentration, where only two replicates were performed. Each amplification curve is 40 cycles long. Please read the sections *Experimental set of qPCR data* and *Quantitative PCR assays* of Karlen et al. (2007) for more details.

#### **Format**

A tibble providing amplification curve data in long format.

plate Plate identifier. This corresponds, loosely, to the name of the targets. In the original publication the amplicons are indicated by gene symbol synonyms which we do use here for naming each plate. This differs from the names used in the column target where actual murine gene symbols are used.

well Well identifier, i.e. the position within the 96-well plate.

target Target identifier: murine gene symbol where the amplicon maps to.

dye Type of fluorescence dye, in this data set it is always SYBR Green I master mix (Roche) ("SYBR").

sample Name of the biological sample.

sample\_type Sample type. Most reactions in this data set are standard curves, i.e. "std", but a few no template controls ("ntc") are also included.

replicate Replicate identifier: 1 thru 5.

copies Standard copy number.

dilution Dilution factor. Higher number means greater dilution, e.g. 10 means a 1:10 (ten-fold) dilution.

cycle PCR cycle.

fluor Raw fluorescence values.

#### Source

doi:10.1186/147121058131

karlen 3

### Examples

karlen

# **Index**

```
* datasets
karlen, 2
karlen, 2
tibble, 2
```