

# Package ‘msqc1’

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**Title** Sigma mix MSQC1 data

**Version** 1.10.0

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**Depends** R (>= 3.2), lattice, stats, utils

**Suggests** BiocStyle, knitr, testthat, specL

**Description** The data set contains an eight technical replicate data set and a three replicate dilution series of the MS Qual/Quant QC Mix standard sample (Sigma-Aldrich, Buchs, Switzerland) measured on five different mass spec platforms at the Functional Genomics Center Zurich.

**License** GPL

**LazyData** true

**VignetteBuilder** knitr

**NeedsCompilation** no

**biocViews** ExperimentData, MassSpectrometryData, ReproducibleResearch

**URL** <https://panoramaweb.org/labkey/MSQC1.url>,  
<http://fgcz-bfabric.uzh.ch/bfabric/project.html?projectId=1959>

**git\_url** <https://git.bioconductor.org/packages/msqc1>

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## Description

The data set contains an eight technical replicate data set and a three replicate dilution series of the MS Qual/Quant QC Mix standard sample (Sigma-Aldrich, Buchs, Switzerland) measured on five different mass spec platforms at the Functional Genomics Center Zurich.

## Details

To obtain the MSQC1 stock solution, MS Qual/Quant QC Mix (Catalog #: MSQC1, Lot # 081M6281) was purchased from SigmaAldrich (Buchs, Switzerland).

The first sample `msqc1_8rep` is based on eight technical replicates. To derive a complex sample matrix, 10 μl of MSQC1 working solution was mixed with 10μl (approx 10 μg) tryptic yeast digest (*Saccharomyces cerevisiae*), 1μl iRT peptide stock solution (Biognosys, Schlieren, Switzerland), and 19 μl 0.1% FA. This sample is denoted as standard sample.

The `msqc1_dil` sample was derived from a six point dilution series containing relative MSQC1 amounts of 0.025, 0.05, 0.2, 1, 2, and 5 with respect to the standard sample. The amount of yeast digest and iRT peptides was kept constant across all dilution steps. The reference L:H ratio vs. the on column amount of SIL peptide can be found in the data set `peptides`.

Both samples were measured on five mass spectrometers. The mass spectrometers and operation modes are:

- QTRAP (ABSciex, Concord, Canada) - SRM mode
- TSQvantage (ThermoScientific, Bremen, Germany) - SRM mode
- Qexactive (ThermoScientific, Bremen, Germany) - PRM mode
- QExactiveHF (ThermoScientific, Bremen, Germany) - DIA mode
- TRIPLETOF 5600 (ABSciex, Concord, Canada) - SWATH and MS1 mode

All Raw LC-MS data from all platforms were imported into Skyline 3.1 (doi: [10.1101/bioinformatics/btq054](https://doi.org/10.1101/bioinformatics/btq054)). A single expert user adjusted the automatic peak group selection and integration border positioning (skyline legacy peak scoring).

The data were exported from skyline as csv files and included in this package as `data.frame` in RData files.

## Author(s)

Tobias Kockmann, Christian Trachsel, Christian Panse

Maintainer: Christian Panse <[cp@fgcz.ethz.ch](mailto:cp@fgcz.ethz.ch)>

## References

- [www.sigmaaldrich.com](http://www.sigmaaldrich.com)
- skyline software - doi: [10.1101/bioinformatics/btq054](https://doi.org/10.1101/bioinformatics/btq054)
- peptide mix <http://www.sigmaaldrich.com/life-science/proteomics/mass-spectrometry/ms-qual-quant-qc-mix.html>, 2014

- Functional Genomic Center Zurich data repository (internal project p1959) <http://fgcz-bfabric.uzh.ch/bfabric/userlab/show-project.html?projectId=1959> - contains all mass spectrometer generated raw files. access for registered users.
- <https://panoramaweb.org/labkey/project/PanoramaPublic/> search for msqc1 (not published yet) - contains all skyline files and the export template

## Examples

```
# Have Fun!
browseVignettes('msqc1')
```

msqc1\_8rep

*MSQC1 8 technical replicates*

## Description

The data set contains a 8 technical replicates measured on five mass spec devices.

## Usage

```
data("msqc1_8rep")
```

## Format

A data frame with 6272 observations on the following 16 variables.

Replicate.Name	a factor with levels 04_MSQC1_in_yd_incl_iRT_1in40_rep1 05_MSQC1_in_yd_incl_iRT_1in40_rep2 06_MSQC1_in_yd_incl_iRT_1in40_rep3 07_MSQC1_in_yd_incl_iRT_1in40_rep4 08_MSQC1_in_yd_incl_iRT_1in40_rep5 09_MSQC1_in_yd_incl_iRT_1in40_rep6 10_MSQC1_in_yd_incl_iRT_1in40_rep7 11_MSQC1_in_yd_incl_iRT_1in40_rep8 20131204_001_HS_yeast 20131204_002_HS_yeast 20131204_003_HS_yeast 20131204_003_HS_yeast_01 20131204_004_HS_yeast 20131204_004_HS_yeast_02 20131204_005_HS_yeast 20131204_005_HS_yeast_03 20131204_006_HS_yeast 20131204_006_HS_yeast_04 20131204_007_HS_yeast 20131204_008_HS_yeast 20131204_008_HS_yeast_05 20131204_009_HS_yeast_06 20131204_010_HS_yeast_07 20131204_011_HS_yeast_08 20140212_006_MSQC1_QTrap_01 20140212_007_MSQC1_QTrap_02 20140212_008_MSQC1_QTrap_03 20140212_009_MSQC1_QTrap_04 20140212_011_MSQC1_QTrap_05 20140212_012_MSQC1_QTrap_06 20140212_013_MSQC1_QTrap_07 20140212_014_MSQC1_QTrap_08 20140326_01_MSQC1_Yeast_iRT_12MS2_1e5_70k 20140326_02_MSQC1_Yeast_iRT_12MS2_1e5_70k 20140326_03_MSQC1_Yeast_iRT_12MS2_1e5_70k 20140326_04_MSQC1_Yeast_iRT_12MS2_1e5_70k 20140326_05_MSQC1_Yeast_iRT_12MS2_1e5_70k 20140326_06_MSQC1_Yeast_iRT_12MS2_1e5_70k 20140326_07_MSQC1_Yeast_iRT_12MS2_1e5_70k 20140326_08_MSQC1_Yeast_iRT_12MS2_1e5_70k
File.Name	a factor with levels 20131204_001_HS_yeast.wiff 20131204_002_HS_yeast.wiff 20131204_003_HS_yeast_01.raw 20131204_003_HS_yeast.wiff 20131204_004_HS_yeast_02.raw 20131204_004_HS_yeast.wiff 20131204_005_HS_yeast_03.raw 20131204_005_HS_yeast.wiff 20131204_006_HS_yeast_04.raw 20131204_006_HS_yeast.wiff 20131204_007_HS_yeast.wiff 20131204_008_HS_yeast_05.raw 20131204_008_HS_yeast.wiff 20131204_009_HS_yeast_06.raw 20131204_010_HS_yeast_07.raw 20131204_011_HS_yeast_08.raw 20140212_006_MSQC1_QTrap_01.wiff 20140212_007_MSQC1_QTrap_02.wiff 20140212_008_MSQC1_QTrap_03.wiff 20140212_009_MSQC1_QTrap_04.wiff 20140212_011_MSQC1_QTrap_05.wiff 20140212_012_MSQC1_QTrap_06.wiff 20140212_013_MSQC1_QTrap_07.wiff 20140326_01_MSQC1_Yeast_iRT_12MS2_1e5_70k.raw 20140326_02_MSQC1_Yeast_iRT_12MS2_1e5_70k.raw 20140326_03_MSQC1_Yeast_iRT_12MS2_1e5_70k.raw 20140326_04_MSQC1_Yeast_iRT_12MS2_1e5_70k.raw 20140326_05_MSQC1_Yeast_iRT_12MS2_1e5_70k.raw

```

20140326_06_MSQC1_Yeast_iRT_12MS2_1e5_70k.raw 20140326_07_MSQC1_Yeast_iRT_12MS2_1e5_70k.raw
20140326_08_MSQC1_Yeast_iRT_12MS2_1e5_70k.raw 20150526_04_MSQC1_in_yd_incl_iRT_1in40_rep1.raw
20150526_05_MSQC1_in_yd_incl_iRT_1in40_rep2.raw 20150526_06_MSQC1_in_yd_incl_iRT_1in40_rep3.
20150526_07_MSQC1_in_yd_incl_iRT_1in40_rep4.raw 20150526_08_MSQC1_in_yd_incl_iRT_1in40_rep5.
20150526_09_MSQC1_in_yd_incl_iRT_1in40_rep6.raw 20150526_10_MSQC1_in_yd_incl_iRT_1in40_rep7.
20150526_11_MSQC1_in_yd_incl_iRT_1in40_rep8.raw

Protein.Name a factor with levels CAH1/P00915 CAH2/P00918 Catalase/P04040 CRP/P02741
DT-Diaphorase(NQ01)/P15559 iRT-C18 Standard Peptides PPIA/Q13427

Peptide.Sequence a factor with levels ADVTPADFSEWSK ALIVLAHSER AVQQPDGLAVLGIFLK DGLDAASYYAPVR
EGHLSPDIVAEQK ESDTSYVSLK FEDENFILK FSTVAGESGSADTVR GAGAFGYFEVTHDITK GAGSSEPVGTLAK
GGPFSDSYR GTFIIDPGGVIR GYSIFSYATK LFLQFGAQGSPFLK LGGNEQVTR NLSVEDAAR
SADFTNFDPR TAENFR TPVISGGPYEYR TPVITGAPYEYR VEATFGVDESNAK VLDALQAIK VSFELFADK
YILAGVENSK

Isotope.Label.Type a factor with levels heavy light
Precursor.Mz a numeric vector
Precursor.Charge a numeric vector
Product.Mz a numeric vector
Product.Charge a numeric vector
Fragment.Ion a factor with levels b8 precursor precursor [M+1] precursor [M+2] y10 y11
y12 y4 y5 y6 y7 y8 y9
Retention.Time a numeric vector
Area a numeric vector
Background a numeric vector
relative.amount a numeric vector
instrument a factor with levels QExactive QExactiveHF QTRAP TRIPLETOF TSQVantage
File.Name.Id a numeric vector

```

### **Value**

A `data.frame` and `msqc1` S3 object.

### **Examples**

```

data(msqc1_8rep)
## maybe str(msqc1_8rep) ; plot(msqc1_8rep) ...

```

**msqc1\_8rep\_QEXACTIVEHF\_assayDevel**  
*QEXACTIVEHF peptide signal response.*

### **Description**

This data set contains peptide signal response for the QEXACTIVEHF mass spec device. The peptide level signals as measured by DIA.

### **Usage**

```
data("msqc1_8rep_QEXACTIVEHF_assayDevel")
```

## Format

A data frame with 1864 observations on the following 16 variables.

```
Replicate.Name a factor with levels 04_MSQC1_in_yd_incl_iRT_1in40_rep1 05_MSQC1_in_yd_incl_iRT_1in40_r
  06_MSQC1_in_yd_incl_iRT_1in40_rep3 07_MSQC1_in_yd_incl_iRT_1in40_rep4 08_MSQC1_in_yd_incl_iRT_1in40_
  09_MSQC1_in_yd_incl_iRT_1in40_rep6 10_MSQC1_in_yd_incl_iRT_1in40_rep7 11_MSQC1_in_yd_incl_iRT_1in40_r
File.Name a factor with levels 20150526_04_MSQC1_in_yd_incl_iRT_1in40_rep1.raw 20150526_05_MSQC1_in_yd_
  20150526_06_MSQC1_in_yd_incl_iRT_1in40_rep3.raw 20150526_07_MSQC1_in_yd_incl_iRT_1in40_rep4.
  20150526_08_MSQC1_in_yd_incl_iRT_1in40_rep5.raw 20150526_09_MSQC1_in_yd_incl_iRT_1in40_rep6.
  20150526_10_MSQC1_in_yd_incl_iRT_1in40_rep7.raw 20150526_11_MSQC1_in_yd_incl_iRT_1in40_rep8.
Protein.Name a factor with levels CAH1/P00915 CAH2/P00918 Catalase/P04040 CRP/P02741
  DT-Diaphorase(NQ01)/P15559 iRT-C18 Standard Peptides P00915MSQC1|CAH1_HUMAN_MSQC1
  P00918MSQC1|CAH2_HUMAN_MSQC1 P02741MSQC1|CRP_HUMAN_MSQC1 P04040MSQC1|CATA_HUMAN_MSQC1
  P15559MSQC1|NQ01_HUMAN_MSQC1 P62937MSQC1|PPIA_HUMAN_MSQC1 PPIA/Q13427
Peptide.Sequence a factor with levels ADVLTGAGNPVGDK ADVTPADFSEWSK AFYVNVLNEEQR ALIVLAHSER
  APLTKPLK AVQQPDGLAVLGIFLK DAQIFIQK DGLDAASYYAPVR DLFNAIATGK DPANFQYPAESVLAYK
  DPILFPSFIHSQK DYPLIPVGK EGHLSPDIVAEQK EIINVGHFSHVNFEDNDR EPISVSSEQVLK ESDTSYVSLK
  ESISVSSEQLAQFR FEDENFILK FGLSGVHHLGK FNTANDDNVTQVR FSTVAGESGSADTVR GAGAFGYFEVTHDITK
  GAGSSEPVTLDAK GGPFSDSYR GGPLDGTYR GTFIIDPAAVIR GTFIIDPGGVIR GYSIFSYATK HDTSLKPISVSYNPATAK
  HNGPEHWHK ILNNNGHAFNVEFDDSQDK IQILEGWK LCENIAGHLK LFAYPDTHR LFLQFGAQGSPFLK LFQFHFWGSTNEHGSEHTVD
  LGGNEQVTR LGPNYLHIPVNCPYR LIQFHFHWGSLDGQGSEHTVDK LNVTVGPR LSQEDPDYGIR LVNANGEAVYCK
  LYPIANGNNQSPVDIK NAIHTFVQSGSHLAAR NGPEQWSK NLSVEDAAR QDNEILIFWSK QSPVIDTHTAK
  SADFTNFDPR SIPTDNQIK TEWLDGK TPVISGGPYEYR TPVITGAPYEYR VEATFGVDESNAK VFEHIGK
  VGSAKPGLQK VLDAHQAIK VSFELFADK VVDVLDSIK YAAELHLVHWNTK YDPSLKPLSVSYDQATSLR YEVQGEVFTKPKQLWP
  YILAGVENSK YNAEKPK YSAELHVAHWNSAK YSSLAEAASK
Isotope.Label.Type a factor with levels light
Precursor.Mz a numeric vector
Precursor.Charge a numeric vector
Product.Mz a numeric vector
Product.Charge a numeric vector
Fragment.Ion a factor with levels b3 b4 b5 b6 b7 b8 y10 y11 y12 y3 y4 y5 y6 y7 y8 y9
Retention.Time a numeric vector
Area a numeric vector
Background a numeric vector
relative.amount a numeric vector
QEXACTIVEHF a factor with levels QEXACTIVEHF
File.Name.Id a numeric vector
```

## Value

A `data.frame` and `msqc1` S3 object.

## Examples

```
data(msqc1_8rep_QEXACTIVEHF_assayDevel)
## maybe str(msqc1_8rep_QEXACTIVEHF_assayDevel) ; plot(msqc1_8rep_QEXACTIVEHF_assayDevel) ...
```

**msqc1\_8rep\_tripleTOF5600\_assayDevel**  
*tripleTOF5600 peptide signal response*

## Description

This data set contains peptide signal response for the tripleTOF5600 mass spec device. The peptide level signals were measured in SWATH-MS mode.

## Usage

```
data("msqc1_8rep_tripleTOF5600_assayDevel")
```

## Format

A data frame with 1616 observations on the following 17 variables.

**Replicate.Name** a factor with levels 20131204\_001\_HS\_yeast 20131204\_002\_HS\_yeast 20131204\_003\_HS\_yeast  
 20131204\_004\_HS\_yeast 20131204\_005\_HS\_yeast 20131204\_006\_HS\_yeast 20131204\_007\_HS\_yeast  
 20131204\_008\_HS\_yeast

**File.Name** a factor with levels 20131204\_001\_HS\_yeast.wiff 20131204\_002\_HS\_yeast.wiff  
 20131204\_003\_HS\_yeast.wiff 20131204\_004\_HS\_yeast.wiff 20131204\_005\_HS\_yeast.wiff  
 20131204\_006\_HS\_yeast.wiff 20131204\_007\_HS\_yeast.wiff 20131204\_008\_HS\_yeast.wiff

**Protein.Name** a factor with levels CAH1/P00915 CAH2/P00918 Catalase/P04040 CRP/P02741  
 DT-Diaphorase(NQ01)/P15559 iRT-C18 Standard Peptides P00915MSQC1|CAH1\_HUMAN\_MSQC1  
 P00918MSQC1|CAH2\_HUMAN\_MSQC1 P02741MSQC1|CRP\_HUMAN\_MSQC1 P04040MSQC1|CATA\_HUMAN\_MSQC1  
 P15559MSQC1|NQ01\_HUMAN\_MSQC1 P62937MSQC1|PPIA\_HUMAN\_MSQC1 PPIA/Q13427

**Peptide.Sequence** a factor with levels ADGLAVIGVLMK ADVLTTGAGNPVGDK ADVTPADFSEWSK AFYVNVLNEEQR  
 ALIVLAHSER APLTKPLK AVQQPDGLAVLGIFLK DAQIFIQK DGLDAASYYAPVR DLFNAIATGK DPILFPSFIHSQK  
 DYPLIPVGK EGHLSPDIVAEQK EGMNIVEAMER EPISVSSEQVLK ESDTSYVSLK ESISVSSEQLAQFR FEDENFILK  
 FGLSVGHHLGK FNTANDDNVTQVR FSTVAGESGSADTVR GAGAFGYFEVTHDITK GAGSSEPVTLDAK GGPFSDSYR  
 GPLDGTYR GTFIIDPAAVIR GTFIIDPGGVIR GYSIFSYATK HDTSLKPISVSYNPATAK ILNNNGHAFNVVFDDSQDK  
 IQILEGWK LFAYPDTHR LFLQFGAQGSPFLK LGGNEQVTR LNVITVGPR LSQEDPDYGYR LYPIANGNNQSPVDIK  
 NAIHTFVQSGSHLAAR NGPEQWSK NLSVEDAAR QSPVDIDTHTAK SADFTNFDPR TEWLDGK TPVISGGPYEYR  
 TPVITGAPYEYR TSFNYAMK VEATFGVDESNAK VGSAKPGLQK VLDALQAIK VSFELFADK VVDVLDTSIK  
 YAAELHLVHWNTK YDPSLKPLSVSYDQATSLR YILAGVENS K YSAELHVAHWSAK YSSLAEAASK

**Isotope.Label.Type** a factor with levels light

**Precursor.Mz** a numeric vector

**Precursor.Charge** a numeric vector

**Product.Mz** a numeric vector

**Product.Charge** a numeric vector

**Fragment.Ion** a factor with levels b2 b3 b7 b8 precursor precursor [M+1] precursor [M+2]  
 y10 y11 y12 y13 y14 y2 y3 y4 y5 y6 y7 y8 y9

**Retention.Time** a numeric vector

**Area** a numeric vector

**Background** a numeric vector

**relative.amount** a numeric vector

```
flag a factor with levels False True
instrument a factor with levels TRIPLETOF
File.Name.Id a numeric vector
```

## Details

The signal response of the peptide \*TAENF[R]\* was not measured due to the lower m/z limit used by the DIA/SWATH-MS methods.

## Value

A `data.frame` and `msqc1` S3 object.

## Examples

```
data(msqc1_8rep_tripleTOF5600_assayDevel)

#filter <- !(msqc1_8rep_tripleTOF5600_assayDevel$Protein.Name %in% peptides$Protein.Name)

#msqc1:::.signal_response_ident_plot(msqc1_8rep_tripleTOF5600_assayDevel[filter, ],
#  instrument="tripleTOF560",
#  prot="P02741")
```

`msqc1_dil`

*msqc1 dilution series*

## Description

MSQC1 3 technical replicates dilution series measured on five mass spec devices.

## Usage

```
data("msqc1_dil")
```

## Format

A data frame with 15300 observations on the following 15 variables.

```
Replicate.Name a factor with levels 20140807_01_MSQC1_1_40dil_1_140808162146 20140807_02_MSQC1_1_40dil_
20140807_02_MSQC1_1_40dil_2_140808175142 20140807_03_MSQC1_1_40dil_3_140808192115
20140807_04_MSQC1_1_20dil_1_140808214050 20140807_05_MSQC1_1_20dil_2_140808231024
20140807_06_MSQC1_1_20dil_3_140809004002 20140807_07_MSQC1_1_5dil_1_140809025941
20140807_08_MSQC1_1_5dil_2_140809051908 20140807_09_MSQC1_1_5dil_3_20140807_10_MSQC1_nodil_1
20140807_11_MSQC1_nodil_2_20140807_12_MSQC1_nodil_3_20140807_13_MSQC1_x2_1_20140807_14_MSQC1_
20140807_15_MSQC1_x2_3_20140807_16_MSQC1_x5_1_20140807_17_MSQC1_x5_2_20140807_18_MSQC1_x5_3
20140808_02_MSQC1_1_40dil_2_140808175141 20140812_004_QC_dil1_40_1_20140812_005_QC_dil1_40_2
20140812_006_QC_dil1_40_3_20140812_008_QC_dil1_20_1_20140812_009_QC_dil1_20_2
20140812_010_QC_dil1_20_3_20140812_012_QC_dil1_5_1_re_20140812_013_QC_dil1_5_2_re
20140812_014_QC_dil1_5_3_20140812_016_QC_nodil_1_20140812_017_QC_nodil_2_20140812_018_QC_nodil_
20140812_020_QC_conc2_1_20140812_021_QC_conc2_2_20140812_022_QC_conc_2_3_20140812_024_QC_conc_
20140812_025_QC_conc5_2_20140812_026_QC_conc5_3_20140818_003_MSQC1_1_40dil_1
```

*msqc1\_dil*

20140818\_004\_MSQC1\_1\_40dil\_1 20140818\_005\_MSQC1\_1\_40dil\_3 20140818\_007\_MSQC1\_1\_20dil\_1  
 20140818\_008\_MSQC1\_1\_20dil\_2 20140818\_009\_MSQC1\_1\_20dil\_3 20140818\_011\_MSQC1\_NoDil\_1  
 20140818\_013\_MSQC1\_NoDil\_2 20140818\_015\_MSQC1\_NoDil\_3 20140818\_017\_MSQC1\_x2\_1  
 20140818\_019\_MSQC1\_x2\_2 20140818\_021\_MSQC1\_x2\_3 20140818\_023\_MSQC1\_x5\_1 20140818\_025\_MSQC1\_x5\_2  
 20140818\_027\_MSQC1\_x5\_3 20140818\_030\_MSQC1\_1\_5dil\_1 20140818\_032\_MSQC1\_1\_5dil\_2  
 20140818\_034\_MSQC1\_1\_5dil\_3 20150526\_16\_MSQC1\_40xdil\_rep1 20150526\_17\_MSQC1\_40xdil\_rep2  
 20150526\_18\_MSQC1\_40xdil\_rep3 20150526\_20\_MSQC1\_20xdil\_rep1 20150526\_21\_MSQC1\_20xdil\_rep2  
 20150526\_22\_MSQC1\_20xdil\_rep3 20150526\_24\_MSQC1\_5xdil\_rep1 20150526\_25\_MSQC1\_5xdil\_rep2  
 20150526\_26\_MSQC1\_5xdil\_rep3 20150526\_28\_MSQC1\_nodil\_rep1 20150526\_29\_MSQC1\_nodil\_rep2  
 20150526\_30\_MSQC1\_nodil\_rep3 20150526\_32\_MSQC1\_2x\_rep1 20150526\_33\_MSQC1\_2x\_rep2  
 20150526\_34\_MSQC1\_2x\_rep3 20150526\_36\_MSQC1\_5x\_rep1 20150526\_37\_MSQC1\_5x\_rep2  
 20150526\_38\_MSQC1\_5x\_rep3 20150601\_001\_QCdil\_40dil\_1 20150601\_002\_QCdil\_40dil\_2  
 20150601\_003\_QCdil\_40dil\_3 20150601\_004\_QCdil\_20dil\_1 20150601\_005\_QCdil\_20dil\_2  
 20150601\_006\_QCdil\_20dil\_3 20150601\_007\_QCdil\_5dil\_1 20150601\_008\_QCdil\_5dil\_2  
 20150601\_009\_QCdil\_5dil\_3 20150601\_010\_nodil\_1 20150601\_011\_nodil\_2 20150601\_012\_nodil\_3  
 20150601\_013\_2conc\_1\_re 20150601\_014\_2conc\_2\_re 20150601\_015\_2conc\_3 20150601\_016\_5conc\_1  
 20150601\_017\_5conc\_2 20150601\_018\_5conc\_3

**File.Name** a factor with levels 20140807\_01\_MSQC1\_1\_40dil\_1\_140808162146.raw 20140807\_02\_MSQC1\_1\_40dil\_1\_140808192115.raw 20140807\_03\_MSQC1\_1\_40dil\_1\_140808231024.raw 20140807\_04\_MSQC1\_1\_20dil\_1\_140808214050.raw 20140807\_05\_MSQC1\_1\_20dil\_2\_140808231024.raw 20140807\_06\_MSQC1\_1\_20dil\_3\_140809004002.raw 20140807\_07\_MSQC1\_1\_5dil\_1\_140809025941.raw 20140807\_08\_MSQC1\_1\_5dil\_2\_140809051908.raw 20140807\_09\_MSQC1\_1\_5dil\_3.raw 20140807\_10\_MSQC1\_nodil\_1.raw 20140807\_11\_MSQC1\_nodil\_2.raw 20140807\_12\_MSQC1\_nodil\_3.raw 20140807\_13\_MSQC1\_x2\_1.raw 20140807\_14\_MSQC1\_x2\_2.raw 20140807\_15\_MSQC1\_x2\_3.raw 20140807\_16\_MSQC1\_x5\_1.raw 20140807\_17\_MSQC1\_x5\_2.raw 20140807\_18\_MSQC1\_x5\_3.raw 20140812\_004\_QC\_dil1\_40\_1.raw 20140812\_005\_QC\_dil1\_40\_2.raw 20140812\_006\_QC\_dil1\_40\_3.raw 20140812\_008\_QC\_dil1\_20\_1.raw 20140812\_009\_QC\_dil1\_20\_2.raw 20140812\_010\_QC\_dil1\_20\_3.raw 20140812\_012\_QC\_dil1\_5\_1\_re.raw 20140812\_013\_QC\_dil1\_5\_2\_re.raw 20140812\_014\_QC\_dil1\_5\_3.raw 20140812\_016\_QC\_nodil\_1.raw 20140812\_017\_QC\_nodil\_2.raw 20140812\_018\_QC\_nodil\_3.raw 20140812\_020\_QC\_conc2\_1.raw 20140812\_021\_QC\_conc2\_2.raw 20140812\_022\_QC\_conc2\_3.raw 20140812\_024\_QC\_conc5\_1.raw 20140812\_025\_QC\_conc5\_2.raw 20140812\_026\_QC\_conc5\_3.raw 20140818\_003\_MSQC1\_1\_40dil\_1.wiff 20140818\_004\_MSQC1\_1\_40dil\_1.wiff 20140818\_005\_MSQC1\_1\_40dil\_3.wiff 20140818\_007\_MSQC1\_1\_20dil\_1.wiff 20140818\_008\_MSQC1\_1\_20dil\_3.wiff 20140818\_009\_MSQC1\_1\_20dil\_3.wiff 20140818\_011\_MSQC1\_NoDil\_1.wiff 20140818\_013\_MSQC1\_NoDil\_2.wiff 20140818\_015\_MSQC1\_NoDil\_3.wiff 20140818\_017\_MSQC1\_x2\_1.wiff 20140818\_019\_MSQC1\_x2\_2.wiff 20140818\_021\_MSQC1\_x2\_3.wiff 20140818\_023\_MSQC1\_x5\_1.wiff 20140818\_025\_MSQC1\_x5\_2.wiff 20140818\_027\_MSQC1\_x5\_3.wiff 20140818\_030\_MSQC1\_1\_5dil\_1.wiff 20140818\_032\_MSQC1\_1\_5dil\_2.wiff 20140818\_034\_MSQC1\_1\_5dil\_3.wiff 20150526\_16\_MSQC1\_40xdil\_rep1.raw 20150526\_17\_MSQC1\_40xdil\_rep2.raw 20150526\_18\_MSQC1\_40xdil\_rep3.raw 20150526\_20\_MSQC1\_20xdil\_rep1.raw 20150526\_21\_MSQC1\_20xdil\_rep2.raw 20150526\_22\_MSQC1\_20xdil\_rep3.raw 20150526\_24\_MSQC1\_5xdil\_rep1.raw 20150526\_25\_MSQC1\_5xdil\_rep2.raw 20150526\_26\_MSQC1\_5xdil\_rep3.raw 20150526\_28\_MSQC1\_nodil\_rep1.raw 20150526\_29\_MSQC1\_nodil\_rep2.raw 20150526\_30\_MSQC1\_nodil\_rep3.raw 20150526\_32\_MSQC1\_2x\_rep1.raw 20150526\_33\_MSQC1\_2x\_rep2.raw 20150526\_34\_MSQC1\_2x\_rep3.raw 20150526\_36\_MSQC1\_5x\_rep1.raw 20150526\_37\_MSQC1\_5x\_rep2.raw 20150526\_38\_MSQC1\_5x\_rep3.raw 20150601\_001\_QCdil\_40dil\_1.wiff 20150601\_002\_QCdil\_40dil\_2.wiff 20150601\_003\_QCdil\_40dil\_3.wiff 20150601\_004\_QCdil\_20dil\_1.wiff 20150601\_005\_QCdil\_20dil\_2.wiff 20150601\_006\_QCdil\_20dil\_3.wiff 20150601\_007\_QCdil\_5dil\_1.wiff 20150601\_008\_QCdil\_5dil\_2.wiff 20150601\_009\_QCdil\_5dil\_3.wiff 20150601\_010\_nodil\_1.wiff 20150601\_011\_nodil\_2.wiff 20150601\_012\_nodil\_3.wiff 20150601\_013\_2conc\_1\_re.wiff 20150601\_014\_2conc\_2\_re.wiff 20150601\_015\_2conc\_3.wiff 20150601\_016\_5conc\_1.wiff 20150601\_017\_5conc\_2.wiff 20150601\_018\_5conc\_3.wiff

**Protein.Name** a factor with levels CAH1/P00915 CAH2/P00918 Catalase/P04040 CRP/P02741

DT-Diaphorase(NQ01)/P15559 iRT-C18 Standard Peptides PPIA/Q13427

**Peptide.Sequence** a factor with levels ADVTPADFSEWSK ALIVLAHSER AVQQPDGLAVLGIFLK DGLDAASYYAPVR

```

EGHLSPDIVAEQK ESDTSYVSLK FEDENFILK FSTVAGESGSADTVR GAGAFGYFEVTHDITK GAGSSEPVTLDAK
GGPFSDSYR GTFIIDPAAVIR GTFIIDPGGVIR GYSIFSYATK LFLQFGAQGSPFLK LGGNEQVTR NLSVEDAAR
SADFTNFDPR TAENFR TPVISGGPYEYR TPVITGAPYEYR VEATFGVDESNAK VLDALQAIK VSFELFADK
YILAGVENSK

Isotope.Label.Type a factor with levels heavy light
Precursor.Mz a numeric vector
Precursor.Charge a numeric vector
Product.Mz a numeric vector
Product.Charge a numeric vector
Fragment.Ion a factor with levels b8 precursor precursor [M+1] precursor [M+2] y10 y11
y12 y4 y5 y6 y7 y8 y9
Retention.Time a numeric vector
Area a numeric vector
Background a numeric vector
relative.amount a numeric vector
instrument a factor with levels QExactive QExactiveHF QTRAP TRIPLETOF TSQVantage

```

**Value**

A `data.frame` and `msqc1` S3 object.

**Examples**

```

data(msqc1_dil)
## maybe str(msqc1_dil) ; plot(msqc1_dil) ...

```

msqc1\_userstudy

*msqc1 user study***Description**

contains a user study

**Usage**

```
data("msqc1_userstudy")
```

**Format**

A data frame with 21204 observations on the following 17 variables.

```

Replicate.Name a factor with levels 01_QCdil_40dil_1 02_QCdil_40dil_2 03_MSQC1_1_40dil_1
03_QCdil_40dil_3 04_MSQC1_1_40dil_1 04_QCdil_20dil_1 05_MSQC1_1_40dil_3 05_QCdil_20dil_2
06_QCdil_20dil_3 07_MSQC1_1_20dil_1 07_QCdil_5dil_1 08_MSQC1_1_20dil_2 08_QCdil_5dil_2
09_MSQC1_1_20dil_3 09_QCdil_5dil_3 10_nodil_1 11_MSQC1_NoDil_1 11_nodil_2 12_nodil_3
13_2conc_1_re 13_MSQC1_NoDil_2 14_2conc_2_re 15_2conc_3 15_MSQC1_NoDil_3 16_5conc_1
17_5conc_2 17_MSQC1_x2_1 18_5conc_3 19_MSQC1_x2_2 20140818_003_MSQC1_1_40dil_1
20140818_004_MSQC1_1_40dil_1 20140818_005_MSQC1_1_40dil_3 20140818_007_MSQC1_1_20dil_1
20140818_008_MSQC1_1_20dil_2 20140818_009_MSQC1_1_20dil_3 20140818_011_MSQC1_NoDil_1

```

20140818\_013\_MSQC1\_NoDil\_2 20140818\_015\_MSQC1\_NoDil\_3 20140818\_017\_MSQC1\_x2\_1  
 20140818\_019\_MSQC1\_x2\_2 20140818\_021\_MSQC1\_x2\_3 20140818\_023\_MSQC1\_x5\_1 20140818\_025\_MSQC1\_x5\_2  
 20140818\_027\_MSQC1\_x5\_3 20140818\_030\_MSQC1\_1\_5dil\_1 20140818\_032\_MSQC1\_1\_5dil\_2  
 20140818\_034\_MSQC1\_1\_5dil\_3 20150601\_001\_QCdil\_40dil\_1 20150601\_002\_QCdil\_40dil\_2  
 20150601\_003\_QCdil\_40dil\_3 20150601\_004\_QCdil\_20dil\_1 20150601\_005\_QCdil\_20dil\_2  
 20150601\_006\_QCdil\_20dil\_3 20150601\_007\_QCdil\_5dil\_1 20150601\_008\_QCdil\_5dil\_2  
 20150601\_009\_QCdil\_5dil\_3 20150601\_010\_nodil\_1 20150601\_011\_nodil\_2 20150601\_012\_nodil\_3  
 20150601\_013\_2conc\_1\_re 20150601\_014\_2conc\_2\_re 20150601\_015\_2conc\_3 20150601\_016\_5conc\_1  
 20150601\_017\_5conc\_2 20150601\_018\_5conc\_3 21\_MSQC1\_x2\_3 23\_MSQC1\_x5\_1 25\_MSQC1\_x5\_2  
 27\_MSQC1\_x5\_3 30\_MSQC1\_1\_5dil\_1 32\_MSQC1\_1\_5dil\_2 34\_MSQC1\_1\_5dil\_3

File.Name a factor with levels 20140818\_003\_MSQC1\_1\_40dil\_1.wiff 20140818\_004\_MSQC1\_1\_40dil\_1.wiff  
 20140818\_005\_MSQC1\_1\_40dil\_3.wiff 20140818\_007\_MSQC1\_1\_20dil\_1.wiff 20140818\_008\_MSQC1\_1\_20dil\_3.wiff  
 20140818\_009\_MSQC1\_1\_20dil\_3.wiff 20140818\_011\_MSQC1\_NoDil\_1.wiff 20140818\_013\_MSQC1\_NoDil\_2.wiff  
 20140818\_015\_MSQC1\_NoDil\_3.wiff 20140818\_017\_MSQC1\_x2\_1.wiff 20140818\_019\_MSQC1\_x2\_2.wiff  
 20140818\_021\_MSQC1\_x2\_3.wiff 20140818\_023\_MSQC1\_x5\_1.wiff 20140818\_025\_MSQC1\_x5\_2.wiff  
 20140818\_027\_MSQC1\_x5\_3.wiff 20140818\_030\_MSQC1\_1\_5dil\_1.wiff 20140818\_032\_MSQC1\_1\_5dil\_2.wiff  
 20140818\_034\_MSQC1\_1\_5dil\_3.wiff 20150601\_001\_QCdil\_40dil\_1.wiff 20150601\_002\_QCdil\_40dil\_2.wiff  
 20150601\_003\_QCdil\_40dil\_3.wiff 20150601\_004\_QCdil\_20dil\_1.wiff 20150601\_005\_QCdil\_20dil\_2.wiff  
 20150601\_006\_QCdil\_20dil\_3.wiff 20150601\_007\_QCdil\_5dil\_1.wiff 20150601\_008\_QCdil\_5dil\_2.wiff  
 20150601\_009\_QCdil\_5dil\_3.wiff 20150601\_010\_nodil\_1.wiff 20150601\_011\_nodil\_2.wiff  
 20150601\_012\_nodil\_3.wiff 20150601\_013\_2conc\_1\_re.wiff 20150601\_014\_2conc\_2\_re.wiff  
 20150601\_015\_2conc\_3.wiff 20150601\_016\_5conc\_1.wiff 20150601\_017\_5conc\_2.wiff  
 20150601\_018\_5conc\_3.wiff

Protein.Name a factor with levels CAH1/P00915 CAH2/P00918 Catalase/P04040 CRP/P02741  
 DT-Diaphorase(NQ01)/P15559 iRT-C18 Standard Peptides PPIA/Q13427

Peptide.Sequence a factor with levels ADVTPADFSEWSK ALIVLAHSER AVQQPDGLAVLGIFLK DGLDAASYYAPVR  
 EGHLSPDIVAEQK ESDTSYVSLK FEDENFILK FSTVAGESGSADTVR GAGAFGYFEVTHDITK GAGSSEPVTLDAK  
 GGPFSDSYR GTFIIDPAAVIR GTFIIDPGGVIR GYSIFSYATK LFLQFGAQGSPFLK LGGNEQVTR NLSVEDAAR  
 SADFTNFDPR TAENFR TPVISGGPYEYR TPVITGAPYEYR VEATFGVDES#N/AK VL DALQAIK VSFELFADK  
 YILAGVENSK

Isotope.Label.Type a factor with levels heavy light

Precursor.Mz a numeric vector

Precursor.Charge a numeric vector

Product.Mz a numeric vector

Product.Charge a numeric vector

Fragment.Ion a factor with levels b8 y10 y11 y12 y4 y5 y6 y7 y8 y9

Retention.Time a numeric vector

Area a numeric vector

Background a numeric vector

relative.amount a numeric vector

instrument a factor with levels QTRAP TRIPLETOF

user a factor with levels algorithm\_1 algorithm\_2 algorithm\_3 user\_1 user\_2 user\_3 user\_4  
 user\_5 user\_6

attempt a factor with levels beginner expert legacy mProphet second\_best

## Details

The data were derived by a user study. Therfore each individual was given a skyline document containing the dilution series data recorded on the QTRAP and TripleTOF 5600. Starting from skyline legacy results, users curated the data.

In addition, two mProphet doi: [10.1038/nmeth.1584](https://doi.org/10.1038/nmeth.1584) models (as implemented in skyline) were trained on the 8 replicate data from the same instrument.

## Value

A `data.frame` and `msqc1` S3 object.

## Examples

```
data(msqc1_userstudy)
## maybe str(msqc1_userstudy) ; plot(msqc1_userstudy) ...
```

---

peptides	<i>sigma mix peptides</i>
----------	---------------------------

---

## Description

This table contains `msqc1` peptide properties.

## Usage

```
data("peptides")
```

## Format

A data frame with 14 observations on the following 5 variables.

```
Peptide.Sequence a factor with levels ALIVLAHSER AVQQPDGLAVLGIFLK EGHLSPDIVAEQK ESDTSYVSLK
  FEDENFILK FSTVAGESGSADTVR GAGAFGYFEVTHDITK GGPFSDSYR GYSIFSYATK NLSVEDAAR SADFTNFDPR
  TAENFR VLDALQAIK VSFELFADK
SIL.peptide.per.vial a numeric vector
LH.ratio a numeric vector
actual.LH.ratio a numeric vector
Protein.Name a factor with levels CAH1/P00915 CAH2/P00918 Catalase/P04040 CRP/P02741
  DT-Diaphorase(NQ01)/P15559 PPIA/Q13427
```

## Value

A `data.frame`.

## Source

<http://www.sigmaldrich.com/life-science/proteomics/mass-spectrometry/ms-qual-quant-qc-mix.html>, 2014

## Examples

```
data(peptides)
## maybe str(peptides) ; plot(peptides) ...
```

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