

Package ‘yeastCC’

April 12, 2018

Version 1.18.0

Title Spellman et al. (1998) and Pramila/Breeden (2006) yeast cell cycle microarray data

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Description ExpressionSet for Spellman et al. (1998) yeast cell cycle microarray experiment

License Artistic-2.0

LazyData yes

Depends Biobase (>= 2.5.5)

biocViews ExperimentData, CellCulture, Saccharomyces_cerevisiae_Data, CancerData, MicroarrayData, OneChannelData, GEO

NeedsCompilation no

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breeden	<i>Breeden et al. yeast cell cycle experiment</i>
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Description

ExpressionSet instance; 50 samples from a 25-sample dye-swap of alpha-synchronized yeast cultures

Usage

data(breeden)

Format

The format is:

Formal class 'ExpressionSet' [package "Biobase"] with 7 slots

..@ assayData :<environment: 0x10221ebc8>

..@ phenoData :Formal class 'AnnotatedDataFrame' [package "Biobase"] with 4 slots

.. ..@ varMetadata :'data.frame': 37 obs. of 1 variable:

..\$ labelDescription: chr [1:37] NA NA NA NA ...

.. ..@ data :'data.frame': 50 obs. of 37 variables:

..\$ title : Factor w/ 50 levels "Yeast cell cycle-time point 0 min 2001-08-17_0000.rfm Yeast W303 cells",...: 1 29 3 15 17 19 21 23 25 27 ...

..\$ geo_accession : Factor w/ 50 levels "GSM112133","GSM112134",...: 1 2 3 4 5 6 7 8 9 10 ...

.. ..

..\$ status : Factor w/ 1 level "Public on Aug 05 2006": 1 1 1 1 1 1 1 1 1 1 ...

..\$ submission_date : Factor w/ 1 level "Jun 01 2006": 1 1 1 1 1 1 1 1 1 1 ...

..\$ last_update_date : Factor w/ 1 level "Jun 23 2006": 1 1 1 1 1 1 1 1 1 1 ...

..\$ type : Factor w/ 1 level "RNA": 1 1 1 1 1 1 1 1 1 1 ...

..\$ channel_count : Factor w/ 1 level "2": 1 1 1 1 1 1 1 1 1 1 ...

..\$ source_name_ch1 : Factor w/ 25 levels "Yeast cell cycle-time point 0 min",...: 1 15 2 8 9 ...

10 11 12 13 14 ...

..\$ organism_ch1 : Factor w/ 1 level "Saccharomyces cerevisiae": 1 1 1 1 1 1 1 1 1 1 ...

..\$ characteristics_ch1 : Factor w/ 25 levels "Yeast cell cycle-time point 0 min",...: 1 15 2 8 9 ...

10 11 12 13 14 ...

..\$ treatment_protocol_ch1 : Factor w/ 1 level "Cells were arrested with alpha factor,and released into YEPD to get a synchronized population. Cells were sampled every 5 min a"| __truncated__ : 1 1 1 1 1 1 1 1 1 1 ...

..\$ molecule_ch1 : Factor w/ 1 level "total RNA": 1 1 1 1 1 1 1 1 1 1 ...

..\$ label_ch1 : Factor w/ 2 levels "Cy3","Cy5": 1 1 1 1 1 1 1 1 1 1 ...

..\$ source_name_ch2 : Factor w/ 1 level "Yeast asynchronous culture": 1 1 1 1 1 1 1 1 1 1 ...

..\$ organism_ch2 : Factor w/ 1 level "Saccharomyces cerevisiae": 1 1 1 1 1 1 1 1 1 1 ...

..\$ characteristics_ch2 : Factor w/ 1 level "Yeast asynchronous culture": 1 1 1 1 1 1 1 1 1 1 ...

..\$ treatment_protocol_ch2 : Factor w/ 1 level "Cells were grown overnight to an OD of 0.6 in YEPD": 1 1 1 1 1 1 1 1 1 1 ...

..\$ molecule_ch2 : Factor w/ 1 level "total RNA": 1 1 1 1 1 1 1 1 1 1 ...

..\$ label_ch2 : Factor w/ 2 levels "Cy3","Cy5": 2 2 2 2 2 2 2 2 2 2 ...

..\$ description : Factor w/ 1 level "Yeast cell cycle": 1 1 1 1 1 1 1 1 1 1 ...

..\$ data_processing : Factor w/ 1 level "normalized log ratio using Rosetta Resolver": 1 1 1 1 1 1 1 1 1 1 ...

..\$ platform_id : Factor w/ 1 level "GPL1914": 1 1 1 1 1 1 1 1 1 1 ...

..\$ contact_name : Factor w/ 1 level "Tata,,Pramila": 1 1 1 1 1 1 1 1 1 1 ...

..\$ contact_email : Factor w/ 1 level "tpramila@fhcrc.org": 1 1 1 1 1 1 1 1 1 1 ...

..\$ contact_phone : Factor w/ 1 level "(206)6674483": 1 1 1 1 1 1 1 1 1 1 ...

..\$ contact_laboratory : Factor w/ 1 level "Breeden Lab": 1 1 1 1 1 1 1 1 1 1 ...

..\$ contact_department : Factor w/ 1 level "Basic Sciences": 1 1 1 1 1 1 1 1 1 1 ...

..\$ contact_institute : Factor w/ 1 level "FHCRC": 1 1 1 1 1 1 1 1 1 1 ...

..\$ contact_address : Factor w/ 1 level "1100, Fairview Avenue N": 1 1 1 1 1 1 1 1 1 1 ...

..\$ contact_city : Factor w/ 1 level "Seattle": 1 1 1 1 1 1 1 1 1 1 ...

..\$ contact_state : Factor w/ 1 level "WA": 1 1 1 1 1 1 1 1 1 1 ...

..\$ contact_zip/postal_code: Factor w/ 1 level "98109": 1 1 1 1 1 1 1 1 1 1 ...

..\$ contact_country : Factor w/ 1 level "USA": 1 1 1 1 1 1 1 1 1 1 ...

..\$ supplementary_file : Factor w/ 50 levels "ftp://ftp.ncbi.nih.gov/pub/geo/DATA/supplementary/samples/GSM112133",...: 1 2 3 4 5 6 7 8 9 10 ...

..\$ data_row_count : Factor w/ 1 level "6228": 1 1 1 1 1 1 1 1 1 1 ...

```

.. .. ..$ mins : num [1:50] 0 5 10 15 20 25 30 35 40 45 ...
.. .. ..$ sign : num [1:50] 1 1 1 1 1 1 1 1 1 1 ...
.. .. ..@ dimLabels : chr [1:2] "sampleNames" "sampleColumns"
.. .. ..@ __classVersion__:Formal class 'Versions' [package "Biobase"] with 1 slots
..@ featureData :Formal class 'AnnotatedDataFrame' [package "Biobase"] with 4 slots
.. .. ..@ varMetadata :'data.frame': 10 obs. of 3 variables:
.. .. .. ..$ Column : chr [1:10] "ID" "ORF" "SPOT_ID" "Gene" ...
.. .. .. ..$ Description : Factor w/ 2 levels "", "LINK_PRE:\http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?CMD=search&LINK_SUF:\[gene name]"": 1 2 1 1 1 1 1 1 1 1
.. .. .. ..$ labelDescription: chr [1:10] NA NA NA NA NA ...
.. .. ..@ data :'data.frame': 6228 obs. of 10 variables:
.. .. .. ..$ ID : Factor w/ 6337 levels "YPR105C","YPR106W",...: 116 6322 119 6327 124 133 6328
6329 6330 6331 ...
.. .. .. ..$ ORF : Factor w/ 6222 levels "YPR105C","YPR106W",...: 6221 6221 6221 6221 6221
6221 6221 6221 6221 6221 ...
.. .. .. ..$ SPOT_ID : chr [1:6228] "<blank>" "blank" "E. coli control" "empty" ...
.. .. .. ..$ Gene : Factor w/ 3347 levels "", "AOS1", "APG13",...: 1 1 1 1 1 1 1 1 1 1 ...
.. .. .. ..$ SGDID : Factor w/ 6205 levels "S0006309","S0006310",...: 1001 1001 1001 1001 1001
1001 1001 1001 1001 1001 ...
.. .. .. ..$ CHR : Factor w/ 17 levels "XVI","XV","XIV",...: 13 13 13 13 13 13 13 13 13 13 ...
.. .. .. ..$ ORF.Length: chr [1:6228] "" "" "" "" "" ...
.. .. .. ..$ Process : Factor w/ 488 levels "DNA repair*",...: 194 194 194 194 194 194 194 194 194
194 ...
.. .. .. ..$ Function : Factor w/ 760 levels "CDP-diacylglycerol-inositol 3-phosphatidyltransferase",...:
215 215 215 215 215 215 215 215 215 ...
.. .. .. ..$ Component : Factor w/ 211 levels "19S proteasome regulatory particle",...: 90 90 90 90 90
90 90 90 90 90 ...
.. .. ..@ dimLabels : chr [1:2] "featureNames" "featureColumns"
.. .. ..@ __classVersion__:Formal class 'Versions' [package "Biobase"] with 1 slots
.. .. .. ..@ .Data:List of 1
.. .. .. .. ..$ : int [1:3] 1 1 0
..@ experimentData :Formal class 'MIAME' [package "Biobase"] with 13 slots
.. .. ..@ name : chr [1:2] "Pramila T" "Breeden LL"
.. .. ..@ lab : chr "Fred Hutchinson Cancer Research Center, Seattle, Washington 98109, USA."
.. .. ..@ contact : chr ""
.. .. ..@ title : chr "The Forkhead transcription factor Hcm1 regulates chromosome segregation
genes and fills the S-phase gap in the transcriptional "l __truncated__
.. .. ..@ abstract : chr "Transcription patterns shift dramatically as cells transit from one phase of
the cell cycle to another. To better define this t"l __truncated__
.. .. ..@ url : chr "http://labs.fhcrc.org/breeden/cellcycle/index.html"
.. .. ..@ pubMedIds : chr "16912276"
.. .. ..@ samples : list()
.. .. ..@ hybridizations : list()
.. .. ..@ normControls : list()
.. .. ..@ preprocessing : list()
.. .. ..@ other : list()
.. .. ..@ __classVersion__:Formal class 'Versions' [package "Biobase"] with 1 slots
.. .. .. ..@ .Data:List of 1
.. .. .. .. ..$ : int [1:3] 1 0 0
..@ annotation : chr(0)
..@ protocolData :Formal class 'AnnotatedDataFrame' [package "Biobase"] with 4 slots
.. .. ..@ varMetadata :'data.frame': 0 obs. of 1 variable:

```

```

.. .. ..$ labelDescription: chr(0)
.. .. ..@ data :'data.frame': 50 obs. of 0 variables
.. .. ..@ dimLabels : chr [1:2] "sampleNames" "sampleColumns"
.. .. ..@ __classVersion__:Formal class 'Versions' [package "Biobase"] with 1 slots
.. .. .. ..@ .Data:List of 1
.. .. .. ..$ : int [1:3] 1 1 0
.. .. ..@ __classVersion__:Formal class 'Versions' [package "Biobase"] with 1 slots
.. .. ..@ .Data:List of 4
.. .. ..$ : int [1:3] 2 10 0
.. .. ..$ : int [1:3] 2 5 5
.. .. ..$ : int [1:3] 1 3 0
.. .. ..$ : int [1:3] 1 0 0

```

Details

Retrieved from GEO using `getGEO` in package `GEOquery`, August 27 2009. Variables `mins` and `sign` added to `pData` manually.

Source

PMID 16912276; see `url` slot of `experimentData` slot.

Examples

```

data(breeden)
#
# show how to use the dye-swap 'sign' variable
#
plot(exprs(breeden)["YBL002W",]~breeden$mins)
plot(I(exprs(breeden)["YBL002W",])*breeden$sign)~breeden$mins)

```

orf800

Cell cycle regulated genes from Spellman et al. (1998)

Description

Vector of ORF names for the 800 cell cycle regulated genes identified by the analysis of Spellman et al. (1998). The expression measures and sample descriptions are stored in the [ExpressionSet](#) instance `yeastCC`.

Usage

```
data(orf800)
```

Format

The format is: chr [1:800] "YAL022C" "YAL040C" "YAL053W" "YAL067C" "YAR003W" "YAR007C"
 ...

Source

The 800 ORF names were obtained from the file "CellCycle98.xls" on the Yeast Cell Cycle Analysis Project website (<http://genome-www.stanford.edu/cellcycle/>). The raw data (images, TIFF) and processed data "combine.txt" used to create the `ExpressionSet` instance `yeastCC` are also available on the website. Gene annotation information is available from the Saccharomyces Genome Database (SGD, http://genome-www.stanford.edu/Saccharomyces/gene_list.shtml). The script "createYeastCC.R" for generating the `yeastCC` package is available in `../doc`.

References

Spellman et al. (1998). Comprehensive Identification of Cell Cycle-regulated Genes of the Yeast *Saccharomyces cerevisiae* by Microarray Hybridization. *Molecular Biology of the Cell*, 9: 3273-3297 (`../doc`).

Examples

```
data(orf800)
```

```
spYCCmeta
```

```
metadata on yeast cell cycle microarray experiment
```

Description

`data.frame` instance with metadata on 800 genes

Usage

```
data(spYCCmeta)
```

Format

A data frame with 800 observations on the following 75 variables.

`Process` a factor with levels 4-nitroquinoline-N-oxide resistance ATP synthesis ...

`Function` a factor with levels (1->6)-beta-glucan synthase subunit (putative) Glc7p regulatory subunit ...

`X` a logical vector

`Peak` a factor with levels G1 G2/M M/G1 S S/G2

`Phase.Order` a numeric vector

`Cluster.Order` a numeric vector

`ORF` a factor with levels YAL022C YAL040C YAL053W YAL067C YAR003W YAR007C ...

`YPD` a factor with levels AAD10 ACE2 ADA2 ADK2 AFR1 AGA1 AGA2 AGP1 ...

`SGD` a factor with levels AAD10 ACE2 ADA2 ADK2 AFR1 AGA1 AGA2 ...

`YPD.1` a factor with levels YPD

`SGD.1` a factor with levels SGD

`MIPS` a factor with levels MIPS

`n1` a numeric vector

n2 a numeric vector
 Geomean a numeric vector
 Absolute a numeric vector
 g1 a numeric vector
 g2 a numeric vector
 Geomean.1 a numeric vector
 Absolute.1 a numeric vector
 Deletion a factor with levels irrelevant lethal undocumented viable
 Known. a factor with levels Known New New
 Description a factor with levels Inhibitor of Cdc28p/Cln1p and Cdc28p/Cln2p complexes involved in o
 mating1,3-beta-D-glucan synthase 3'-Phosphoadenylylsulfate reductase; part of the sulfate
 Aggregate.Score a numeric vector
 Phase a numeric vector
 No..Elements a numeric vector
 Most.Relevant.Promoter.Elements a factor with levels AATAGATGACCCGATTTGGAAAAAGGTAACAACAATG
 ATTTGATTGCCGAAAGAGGCAAAAC GTAAATAGTTGT C 156 TCTGCCAGCCAA C 253 AAAGCCAGCCAT
 C 256 TATGCCAGCCAA C 276 AAGGCCAGCCTC C 293 TTGACCAGCTAA ...
 X.1 a factor with levels ATATAGCGACCGAATCAGGAAAAAG GTCAACAACGAAG C 102 CGAGCCAGCATT
 C 252 AAGACCAGCATG C 301 AGTGCCAGCAAA C 496 GAAGCCAGCAC C 550 GCGCCAGCAAC
 c 106 attACGCGaaaat c 112 aaaACGCGagaaa c 121 ggaACGCGgacg ...
 X.2 a factor with levels C 125 GCAACCAGCTCT C 146 CAAGCCAGCCAT C 195 CGCACCAGCAAC C 212
 TATACCAGCGTT C 245 TAAACCAGCGCA C 402 TATGCCAGCAAA c 112 ttaACGCGatcga
 c 115 agtACGCGaaagg c 123 acaACGCGaacac c 127 gtgACGCGaaaaa ...
 X.3 a factor with levels C 307 AAGACCAGCATT c 163 ctgACGCGgaaa c 190 aatACGCGagaaa
 c 220 tagACGCGcctta c 241 cgaACGCGaaact c 275 aaaACGCGaccgt c 282 aagACGCGatttt
 c 289 attACGCGcatta c 290 aggACGCGaaact ...
 X.4 a factor with levels C 200 CAAACCAGCATC c 117 gtcACGCGaaaaa c 314 cctACGCGaaagt
 c 338 caaACGCGaaaaa c 359 acgACGCGccttc c 382 gttACGCGaaagtc c 384 tcaACGCGaattt
 c 397 aaaACGCGaaaac c 440 gtgACGCGcggtt ...
 X.5 a factor with levels C 306 GGAGCCAGCGCG c 467 accACGCGaaaag c 588 gaaACGCGccaaa
 w 266 ATAACCAGCAAA w 383 cagACGCGagaac w 478 GGAGCCAGCGCG w401 tatCGCGAAAatt
 X.6 a factor with levels C 337 AGAGCCAGCAAG C 417 TCGCCAGCAAT c 501 acaACGCGaaaaa
 w 370 gcgACGCGaaaaa w 447 AGAGCCAGCAAG
 X.7 a factor with levels C 388 GGAACCAGCAGA w 396 GGAACCAGCAGA
 Number a numeric vector
 SCB a factor with levels c 103 gacCACGAAAttt c 105 atgCACGAAAaag c 106 ctaCACGAAAcac
 c 108 tacCACGAAAgta c 110 ccaCACGAAAaga c 123 agaCACGAAAtgt c 127 acaCACGAAAacg
 c 181 cagCACGAAAtgg ...
 SCB.1 a factor with levels c178 tgaCACGAAAaac c232 gaaCACGAAAtgc c539 gtaCACGAAAttc
 w269 agcCACGAAAtgc w347 tgaCACGAAAtgt w541 agtCACGAAAacg w601 tgtCACGAAAtgt
 SCB.2 a factor with levels c330 aacCACGAAAaaa c582 agtCACGAAAacg w467 attCACGAAAtaa
 SCB.3 a factor with levels w435 atcCACGAAAatc
 X.8 a factor with levels w252 aacCACGAAAagt
 Number.1 a numeric vector

SCB_d a factor with levels c 156 gatCGCGAAAttt c 184 cgaCGCGAAAatg c 218 cagCGCGAAAagt
c 222 tatCGCGAAAaaa c 229 tgaCGCGAAAacg c 237 tatCGCGAAAacg c 238 atcCGCGAAAagga
c 283 aagCGCGAAAcaa ...

SCB_d.1 a factor with levels c 126 tttCGCGAAAactg c 415 tttCGCGAAAatct c 566 ttcCGCGAAAaaa
c 592 aggCGCGAAAatc c 633 aaaCGCGAAAatg c 242 gaaCGCGAAAactt c 297 ctcCGCGAAAaat
c 306 tcgCGCGAAAaga ...

SCB_d.2 a factor with levels c 468 ccaCGCGAAAaga c 508 tttCGCGAAAatct

SCB_d.3 a factor with levels c 502 caaCGCGAAAaat

Number .2 a numeric vector

MCB a factor with levels w 126 gcaACGCGTcgc w 187 caaACGCGTaca w 207 ctcACGCGTcgg w 209
attACGCGTtta w 226 cagACGCGTtgc w 228 acaACGCGTtct w 23 acaACGCGTgct
w 267 cccACGCGTtagg ...

MCB.1 a factor with levels w 111 gaaACGCGTtct w 124 ttgACGCGTtct w 128 gtgACGCGTtat
w 130 agaACGCGTtct w 131 gcgACGCGTaac w 138 aagACGCGTgaa w 139 attACGCGTttaw w 153 ctaACGCGTttt
...

MCB.2 a factor with levels w 374 taaACGCGTcat

MCB.3 a factor with levels w 309 aggACGCGTaaa

Number .3 a numeric vector

MCB_d a factor with levels c 106 attACGCGGaaaat c 109 acaACGCGactgg c 112 aaaACGCGagaaa
c 115 agtACGCGGaaagg c 117 gtcACGCGGaaaaa c 121 ggaACGCGGacgc c 127 gtgACGCGGaaaaa
c 129 acaACGCGGcccga ...

MCB_d.1 a factor with levels c 123 acaACGCGaacac c 136 aatACGCGattgg c 147 gcaACGCGagaga
c 158 tctACGCGggaag c 163 ctgACGCGggaac c 176 gcgACGCGgttgt c 187 agtACGCGattg
c 189 gaaACGCGggcac ...

MCB_d.2 a factor with levels c 112 ttaACGCGatcga c 220 tagACGCGcctta c 294 ttcACGCGcttaa
c 382 gttACGCGaagtc c 477 gcaACGCGcctgg c 501 acaACGCGaaaaa c 549 attACGCGcacg
c 557 tgtACGCGgcaac ...

MCB_d.3 a factor with levels c 617 gaaACGCGcagta w 50 gtaACGCGctttt

X.9 a factor with levels c 359 acgACGCGccttc

Number .4 a numeric vector

SFF a factor with levels AATAGATGACCCGATTTGGAAAAAGGTAAACAACAATG ATTTGATTGCCGAAAGAGGCAAAC GTAAATAGGTT
CAAAACAACCAATAAAGAAAATCCAAAATATAGAAC GTACTTTAACCTGTTTAGGAAAAAG GTAAACAATAACA
TCGAACAATTCTAAAAAGGTAAAT AAAAAAATGGTA ...

Number .5 a factor with levels 1 2 3 4 ATATAGCGACCGAATCAGGAAAAGGTCAACAACGAAG

Swi5 a factor with levels C 102 CGAGCCAGCATT C 156 TCTGCCAGCCAA C 200 CAAACCAGCATC
C 252 AAGACCAGCATG C 253 AAAGCCAGCCAT C 256 TATGCCAGCCAA C 276 AAGGCCAGCCTC
C 293 TTGACCAGCTAA ...

Swi5.1 a factor with levels C 125 GCAACCAGCTCT C 146 CAAGCCAGCCAT C 195 CGCACCAGCAAC
C 245 TAAACCAGCGCA C 301 AGTGCCAGCAAA C 306 GGAGCCAGCGCG C 307 AAGACCAGCATT
C 402 TATGCCAGCAAA ...

Swi5.2 a factor with levels C 212 TATACCAGCGTT C 337 AGAGCCAGCAAG c 19 AGAACCAGCTGA
c 320 ACCACCAGCTTA c 545 ACCACCAGCGTA c 569 TTCACCAGCGGC c 642 GAGACCAGCGGA
c 651 ATCACCAGCAAA ...

Swi5.3 a factor with levels C 388 GGAACCAGCAGA C 417 TCGGCCAGCAAT c 336 TTTACCAGCTCA
c 363 TGCACCAGCATT c 494 CTGGCCAGCAAG w 396 GGAACCAGCAGA

Number .6 a numeric vector

Swi5e a factor with levels c 102 CGAGCCAGCATT c 137 TAGGCCAGCAAA c 155 ACAACCAGCAGT
c 156 CTAACCAGCAAG c 16 AGAGCCAGCAGA c 174 TAAACCAGCATT c 184 ATGCCAGCATA
c 200 CAAACCAGCATC ...

Swi5e.1 a factor with levels c 222 TTGACCAGCGCC c 256 TAAACCAGCAAA c 306 GGAGCCAGCGCG
c 307 AAGACCAGCATT c 637 GGAGCCAGCGAT w 265 TAAACCAGCAAT w 266 ATAACCAGCAAA
w 467 TGAGCCAGCAAT w 478 GGAGCCAGCGCG w 536 GAAACCAGCAAC w 554 ATGCCAGCACC

Swi5e.2 a factor with levels c 337 AGAGCCAGCAAG c 417 TCGGCCAGCAAT c 642 GAGACCAGCGGA
w 447 AGAGCCAGCAAG

Swi5e.3 a factor with levels c 388 GGAACCAGCAGA w 396 GGAACCAGCAGA

Number.7 a numeric vector

ECB a factor with levels c 185 TTACCCATTTAGGAAA c 221 TTACCCAATTAGGAAA c 251 TTTCCCTTTAAGGAAA
c 258 TTTCCCAAAAAGGAAA c 387 TTTCCCTTTTAGGAAA c 394 TTACCCACTTAGGAAA w 154 TTTCCCTTTTAGGAAA
w 177 TTACCCACTTAGGAAA w 229 TTACCCAGAAAAGGAAA w 378 TTTCCCTAATAGGAAA w 453 TTTCCGTTTAGGAAA
w 595 TTTCCCACTAAGGAAA

Number.8 a numeric vector

STE12 a factor with levels c 243 CCTTTTTTCAGTTTCTATTTTTAACACTGAAACT w 112 CCCTATTTGGTTGCAATTCAATTCCGTGA
w 119 CCAATGTAGAAAAGTACATCATATGAAACA w 218 CCTAATTGGGTAAGTACATGATGAAACA
w 224 CCCAAAAGGAAATTTACATGTAAATGAAACC ...

MIG1.sites a factor with levels c 114 AATAGACTGGGG c 137 TCTATCCTGGGG c 147 TGAATGCTGGGG
c 165 AATAAAGTGGGG c 215 TATAATGCGGGG c 304 AAATCGCCGGGG c 332 AAATATCTGGGG
c 368 AATTGCGCGGGG ...

X.10 a factor with levels c 161 AGTTTGGTGGGG c 262 AAGATGGTGGGG c 498 AAAAAACCGGGG
c 499 AAAAAATGCGGGG w 296 TATTCGGCGGGG w 578 CTTTTGCCGGGG

X.11 a logical vector

Details

taken from the Spellman support web site.

Source

cellcycle-www.stanford.edu

References

PMID 9843569

Examples

```
data(spYCCmeta)
spYCCmeta[1:5,1:6]
```

`yeastCC`*Data from the Spellman et al. (1998) yeast cell cycle microarray experiment*

Description

This data package contains an `ExpressionSet` instance for the yeast cell cycle microarray experiment. The dataset contains gene expression measures (log-ratios, with Cy3-labeled common reference) for 6,178 yeast genes in 77 conditions.

Usage

```
data(yeastCC)
```

Details

There are four main timecourses: alpha (alpha factor arrest), cdc15, cdc28, and elu (elutriation), corresponding to different synchronization methods. For details on experimental procedures and analysis, refer to Spellman et al. (1998) (in `./doc`) and the Yeast Cell Cycle Analysis Project website (<http://genome-www.stanford.edu/cellcycle/>). The `ExpressionSet` instance `yeastCC` was derived from the file "combined.txt" on the website. The ORF names for the 800 cell cycle regulated genes are stored in `orf800`.

Source

The raw data (images, TIFF) and processed data "combine.txt" used to create the `ExpressionSet` instance `yeastCC` are available from the Yeast Cell Cycle Analysis Project website (<http://genome-www.stanford.edu/cellcycle/>). Gene annotation information is available from the Saccharomyces Genome Database (SGD, http://genome-www.stanford.edu/Saccharomyces/gene_list.shtml). The script "createYeastCC.R" for generating the `yeastCC` package is available in `./doc`.

Note that `spYCCES` is an `ExpressionSet` instance with the same data and slightly different phenodata annotation.

References

Spellman et al. (1998). Comprehensive Identification of Cell Cycle-regulated Genes of the Yeast *Saccharomyces cerevisiae* by Microarray Hybridization. *Molecular Biology of the Cell*, 9: 3273-3297.

Examples

```
data(yeastCC)
yeastCC
varLabels(yeastCC)
pData(yeastCC)
description(yeastCC)
abstract(yeastCC)
featureNames(yeastCC)[1:10]
dim(exprs(yeastCC))
```

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