# Package 'cubar'

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Title Codon Usage Bias Analysis

Version 1.0.0

**Description** A suite of functions for rapid and flexible analysis of codon usage bias. It provides in-depth analysis at the codon level, including relative synonymous codon usage (RSCU), tRNA weight calculations, machine learning predictions for optimal or preferred codons, and visualization of codon-anticodon pairing. Additionally, it can calculate various genespecific codon indices such as codon adaptation index (CAI), effective number of codons (ENC), fraction of optimal codons (Fop), tRNA adaptation index (tAI), mean codon stabilization coefficients (CSCg), and GC contents (GC/GC3s/GC4d). It also supports both standard and non-standard genetic code tables found in NCBI, as well as custom genetic code tables.

```
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```

```
URL https://github.com/mt1022/cubar, https://mt1022.github.io/cubar/
```

BugReports https://github.com/mt1022/cubar/issues

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aa2codon

amino acids to codons

# Description

A data.frame of mapping from amino acids to codons

# Usage

aa2codon

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#### **Format**

```
a data.frame with two columns: amino_acid, and codon.

amino_acid amino acid corresponding to the codon

codon codon identity
```

#### **Source**

It is actually the standard genetic code.

# **Examples**

aa2codon

check\_cds

Quality control of CDS

## **Description**

check\_cds performs quality control of CDS sequences by filtering some peculiar sequences and optionally remove start or stop codons.

#### Usage

```
check_cds(
    seqs,
    codon_table = get_codon_table(),
    min_len = 6,
    check_len = TRUE,
    check_start = TRUE,
    check_stop = TRUE,
    check_istop = TRUE,
    rm_start = TRUE,
    rm_stop = TRUE,
    start_codons = c("ATG")
)
```

## Arguments

```
seqs input CDS sequences

codon_table codon table matching the genetic code of seqs

min_len minimum CDS length in nt

check_len check whether CDS length is divisible by 3

check_start check whether CDSs have start codons

check_stop check whether CDSs have stop codons
```

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check\_istop check internal stop codons

rm\_start whether to remove start codons

rm\_stop whether to remove stop codons

start\_codons vector of start codons

#### Value

DNAStringSet of filtered (and trimmed) CDS sequences

## **Examples**

```
# CDS sequence QC for a sample of yeast genes
s <- head(yeast_cds, 10)
print(s)
check_cds(s)</pre>
```

codon\_diff

Differential codon usage analysis

#### **Description**

codon\_diff takes two set of coding sequences and perform differential codon usage analysis.

#### Usage

```
codon_diff(seqs1, seqs2, codon_table = get_codon_table())
```

#### **Arguments**

seqs1 DNAStringSet, or an object that can be coerced to a DNAStringSet seqs2 DNAStringSet, or an object that can be coerced to a DNAStringSet

codon\_table a table of genetic code derived from 'get\_codon\_table' or 'create\_codon\_table'.

#### Value

a data.table of the differential codon usage analysis. Global tests examine whether a codon is used differently relative to all the other codons. Family tests examine whether a codon is used differently relative to other codons that encode the same amino acid. Subfamily tests examine whether a codon is used differently relative to other synonymous codons that share the same first two nucleotides. Odds ratio > 1 suggests a codon is used at higher frequency in seqs1 than in seqs2.

```
yeast_exp_sorted <- yeast_exp[order(yeast_exp$fpkm),]
seqs1 <- yeast_cds[names(yeast_cds) %in% head(yeast_exp_sorted$gene_id, 1000)]
seqs2 <- yeast_cds[names(yeast_cds) %in% tail(yeast_exp_sorted$gene_id, 1000)]
cudiff <- codon_diff(seqs1, seqs2)</pre>
```

codon\_optimize 5

codon\_optimize

Optimize codons

## **Description**

codon\_optimize takes a coding sequence (without stop codon) and replace each codon to the corresponding synonymous optimal codon.

## Usage

```
codon_optimize(
  seq,
  optimal_codons,
  codon_table = get_codon_table(),
  level = "subfam"
)
```

## **Arguments**

seq DNAString, or an object that can be coerced to a DNAString.

optimal\_codons table optimze codons as generated by est\_optimal\_codons.

codon\_table a table of genetic code derived from 'get\_codon\_table' or 'create\_codon\_table'.

level "subfam" (default) or "amino\_acid". Optimize codon usage at which level.

#### Value

a DNAString of the optimized coding sequence.

## **Examples**

```
cf_all <- count_codons(yeast_cds)
optimal_codons <- est_optimal_codons(cf_all)
seq <- 'ATGCTACGA'
codon_optimize(seq, optimal_codons)</pre>
```

count\_codons

Count occurrences of different codons

## **Description**

count\_codons tabulates the occurrences of all the 64 codons in input CDSs

#### Usage

```
count_codons(seqs, ...)
```

6 create\_codon\_table

#### **Arguments**

```
seqs CDS sequences, DNAStringSet.
... additional arguments passed to 'Biostrings::trinucleotideFrequency'.
```

#### Value

```
matrix of codon (column) frequencies of each CDS (row).
```

## **Examples**

```
# count codon occurrences
cf_all <- count_codons(yeast_cds)
dim(cf_all)
cf_all[1:5, 1:5]
count_codons(yeast_cds[1])</pre>
```

create\_codon\_table

create custom codon table from a data frame

# Description

create\_codon\_table creates codon table from data frame of aa to codon mapping.

#### Usage

```
create_codon_table(aa2codon)
```

## Arguments

aa2codon

a data frame with two columns: amino\_acid (Ala, Arg, etc.) and codon.

#### Value

```
a 'data.table' with four columns: aa_code, amino_acid, codon, and subfam.
```

```
head(aa2codon)
create_codon_table(aa2codon = aa2codon)
```

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est\_csc

Estimate Codon Stabilization Coefficient

## **Description**

get\_csc calculate codon occurrence to mRNA stability correlation coefficients (Default to Pearson's).

# Usage

```
est_csc(
  seqs,
  half_life,
  codon_table = get_codon_table(),
  cor_method = "pearson"
)
```

## **Arguments**

seqs CDS sequences of all protein-coding genes. One for each gene.

half\_life data.frame of mRNA half life (gene\_id & half\_life are column names).

codon\_table a table of genetic code derived from 'get\_codon\_table' or 'create\_codon\_table'.

cor\_method method name passed to 'cor.test' used for calculating correlation coefficients.

## Value

data.table of optimal codons.

#### References

Presnyak V, Alhusaini N, Chen YH, Martin S, Morris N, Kline N, Olson S, Weinberg D, Baker KE, Graveley BR, et al. 2015. Codon optimality is a major determinant of mRNA stability. Cell 160:1111-1124.

```
# estimate yeast mRNA CSC
est_csc(yeast_cds, yeast_half_life)
```

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est\_optimal\_codons

Estimate optimal codons

## **Description**

est\_toptimal\_codons determine optimal codon of each codon family with binomial regression. Usage of optimal codons should correlate negatively with enc.

#### Usage

```
est_optimal_codons(
  cf,
  codon_table = get_codon_table(),
  level = "subfam",
  gene_score = NULL,
  fdr = 0.001
)
```

#### **Arguments**

cf matrix of codon frequencies as calculated by 'count\_codons()'.

codon\_table a table of genetic code derived from 'get\_codon\_table' or 'create\_codon\_table'.

level "subfam" (default) or "amino\_acid". For which level to determine optimal codons.

gene\_score a numeric vector of scores for genes. The order of values should match with

gene orders in the codon frequency matrix. The length of the vector should be equal to the number of rows in the matrix. The scores could be gene expression levels (RPKM or TPM) that are optionally log-transformed (for example, with 'log1p'). The opposite of ENC will be used by default if 'gene\_score' is not

provided.

fdr false discovery rate used to determine optimal codons.

## Value

data.table of optimal codons.

```
# perform binomial regression for optimal codon estimation
cf_all <- count_codons(yeast_cds)
codons_opt <- est_optimal_codons(cf_all)
codons_opt <- codons_opt[optimal == TRUE]
codons_opt</pre>
```

est\_rscu 9

## **Description**

est\_rscu returns the RSCU value of codons

## Usage

```
est_rscu(cf, weight = 1, pseudo_cnt = 1, codon_table = get_codon_table())
```

## **Arguments**

cf	matrix of codon frequencies as calculated by 'count_codons()'.
weight	a vector of the same length as 'seqs' that gives different weights to CDSs when count codons. for example, it could be gene expression levels.
pseudo_cnt	pseudo count to avoid dividing by zero. This may occur when only a few sequences are available for RSCU calculation.
codon_table	a table of genetic code derived from 'get_codon_table' or 'create_codon_table'.

#### Value

a data.table of codon info. RSCU values are reported in the last column.

#### References

Sharp PM, Tuohy TM, Mosurski KR. 1986. Codon usage in yeast: cluster analysis clearly differentiates highly and lowly expressed genes. Nucleic Acids Res 14:5125-5143.

```
# compute RSCU of all yeast genes
cf_all <- count_codons(yeast_cds)
est_rscu(cf_all)

# compute RSCU of highly expressed (top 500) yeast genes
heg <- head(yeast_exp[order(-yeast_exp$fpkm), ], n = 500)
cf_heg <- count_codons(yeast_cds[heg$gene_id])
est_rscu(cf_heg)</pre>
```

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	4	weight
A C T	Trna	WAIGHT

Estimate tRNA weight w

## **Description**

 ${\tt est\_trna\_weight}$  compute the tRNA weight per codon for TAI calculation. This weight reflects relative tRNA availability for each codon.

## Usage

```
est_trna_weight(
  trna_level,
  codon_table = get_codon_table(),
  s = list(WC = 0, IU = 0, IC = 0.4659, IA = 0.9075, GU = 0.7861, UG = 0.6295)
)
```

## **Arguments**

trna_level	named vector of tRNA level (or gene copy numbers), one value for each anticodon. vector names are anticodons.
codon_table	a table of genetic code derived from 'get_codon_table' or 'create_codon_table'.
S	list of non-Waston-Crick pairing panelty.

#### Value

data.table of tRNA expression information.

#### References

dos Reis M, Savva R, Wernisch L. 2004. Solving the riddle of codon usage preferences: a test for translational selection. Nucleic Acids Res 32:5036-5044.

```
# estimate codon tRNA weight for yeasts
est_trna_weight(yeast_trna_gcn)
```

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get\_cai

Calculate CAI

## **Description**

```
get_cai calculates Codon Adaptation Index (CAI) of each input CDS
```

## Usage

```
get_cai(cf, rscu)
```

## Arguments

cf matrix of codon frequencies as calculated by 'count\_codons()'.

rscu rscu table containing CAI weight for each codon. This table could be generated

with 'est\_rscu' or prepared manually.

#### Value

a named vector of CAI values

#### References

Sharp PM, Li WH. 1987. The codon Adaptation Index—a measure of directional synonymous codon usage bias, and its potential applications. Nucleic Acids Res 15:1281-1295.

```
# estimate CAI of yeast genes based on RSCU of highly expressed genes
heg <- head(yeast_exp[order(-yeast_exp$fpkm), ], n = 500)
cf_all <- count_codons(yeast_cds)
cf_heg <- cf_all[heg$gene_id, ]
rscu_heg <- est_rscu(cf_heg)
cai <- get_cai(cf_all, rscu_heg)
head(cai)
hist(cai)</pre>
```

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get\_codon\_table

get codon table by NCBI gene code ID

# Description

get\_codon\_table creates a codon table based on the given id of genetic code in NCBI.

## Usage

```
get_codon_table(gcid = "1")
```

## **Arguments**

gcid

a string of genetic code id. run 'show\_codon\_tables()' to see available codon tables.

#### Value

a 'data.table' with four columns: aa\_code, amino\_acid, codon, and subfam.

# **Examples**

```
# Standard genetic code
get_codon_table()

# Vertebrate Mitochondrial genetic code
get_codon_table(gcid = '2')
```

get\_cscg

Mean Codon Stabilization Coefficients

# Description

get\_cscg calculates Mean Codon Stabilization Coefficients of each CDS.

## Usage

```
get_cscg(cf, csc)
```

## **Arguments**

cf matrix of codon frequencies as calculated by 'count\_codons()'.
csc table of Codon Stabilization Coefficients as calculated by 'est\_csc()'.

## Value

a named vector of cscg values.

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#### References

Presnyak V, Alhusaini N, Chen YH, Martin S, Morris N, Kline N, Olson S, Weinberg D, Baker KE, Graveley BR, et al. 2015. Codon optimality is a major determinant of mRNA stability. Cell 160:1111-1124.

# Examples

```
# estimate CSCg of yeast genes
yeast_csc <- est_csc(yeast_cds, yeast_half_life)
cf_all <- count_codons(yeast_cds)
cscg <- get_cscg(cf_all, csc = yeast_csc)
head(cscg)
hist(cscg)</pre>
```

get\_dp

Deviaiton from Proportionality

#### **Description**

get\_dp calculates Deviation from Proportionality of each CDS.

#### Usage

```
get_dp(cf, host_weights, codon_table = get_codon_table())
```

## **Arguments**

cf matrix of codon frequencies as calculated by 'count\_codons()'.

host\_weights a named vector of tRNA weights for each codon that reflects the relative avail-

ability of tRNAs in the host organism.

codon\_table a table of genetic code derived from 'get\_codon\_table' or 'create\_codon\_table'.

#### Value

a named vector of dp values.

#### References

Chen F, Wu P, Deng S, Zhang H, Hou Y, Hu Z, Zhang J, Chen X, Yang JR. 2020. Dissimilation of synonymous codon usage bias in virus-host coevolution due to translational selection. Nat Ecol Evol 4:589-600.

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#### **Examples**

```
# estimate DP of yeast genes
cf_all <- count_codons(yeast_cds)
trna_weight <- est_trna_weight(yeast_trna_gcn)
trna_weight <- setNames(trna_weight$w, trna_weight$codon)
dp <- get_dp(cf_all, host_weights = trna_weight)
head(dp)
hist(dp)</pre>
```

get\_enc

Calculate ENC

## **Description**

```
get_enc computes ENC of each CDS
```

## Usage

```
get_enc(cf, codon_table = get_codon_table())
```

## **Arguments**

cf matrix of codon frequencies as calculated by 'count\_codons()'.

codon\_table codon\_table a table of genetic code derived from 'get\_codon\_table' or 'cre-

ate\_codon\_table'.

#### Value

vector of ENC values, sequence names are used as vector names

#### References

- Wright F. 1990. The 'effective number of codons' used in a gene. Gene 87:23-29. - Sun X, Yang Q, Xia X. 2013. An improved implementation of effective number of codons (NC). Mol Biol Evol 30:191-196.

```
# estimate ENC of yeast genes
cf_all <- count_codons(yeast_cds)
enc <- get_enc(cf_all)
head(enc)
hist(enc)</pre>
```

get\_fop 15

|--|

# Description

get\_fop calculates the fraction of optimal codons (Fop) of each CDS.

# Usage

```
get_fop(cf, op = NULL, codon_table = get_codon_table(), ...)
```

# Arguments

cf	matrix of codon frequencies as calculated by 'count_codons()'.
ор	a character vector of optimal codons. Can be determined automatically by running 'est_optimal_codons'.
codon_table	a table of genetic code derived from 'get_codon_table' or 'create_codon_table'.
• • •	other arguments passed to 'est_optimal_codons'.

# Value

a named vector of fop values.

#### References

Ikemura T. 1981. Correlation between the abundance of Escherichia coli transfer RNAs and the occurrence of the respective codons in its protein genes: a proposal for a synonymous codon choice that is optimal for the E. coli translational system. J Mol Biol 151:389-409.

```
# estimate Fop of yeast genes
cf_all <- count_codons(yeast_cds)
fop <- get_fop(cf_all)
head(fop)
hist(fop)</pre>
```

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get\_gc

GC contents

# Description

Calculate GC content of the whole sequences.

## Usage

```
get_gc(cf)
```

## **Arguments**

cf

matrix of codon frequencies as calculated by 'count\_codons()'.

## Value

a named vector of GC contents.

## **Examples**

```
# estimate GC content of yeast genes
cf_all <- count_codons(yeast_cds)
gc <- get_gc(cf_all)
head(gc)
hist(gc)</pre>
```

get\_gc3s

GC contents at synonymous 3rd codon positions

# Description

Calculate GC content at synonymous 3rd codon positions.

# Usage

```
get_gc3s(cf, codon_table = get_codon_table())
```

# **Arguments**

```
cf matrix of codon frequencies as calculated by 'count_codons()'.
codon_table a table of genetic code derived from 'get_codon_table' or 'create_codon_table'.
```

## Value

a named vector of GC3s values.

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#### References

Peden JF. 2000. Analysis of codon usage.

## **Examples**

```
# estimate GC3s of yeast genes
cf_all <- count_codons(yeast_cds)
gc3s <- get_gc3s(cf_all)
head(gc3s)
hist(gc3s)</pre>
```

get\_gc4d

GC contents at 4-fold degenerate sites

# Description

Calculate GC content at synonymous position of codons (using four-fold degenerate sites only).

## Usage

```
get_gc4d(cf, codon_table = get_codon_table())
```

## **Arguments**

```
cf matrix of codon frequencies as calculated by 'count_codons()'.

codon_table a table of genetic code derived from 'get_codon_table' or 'create_codon_table'.
```

#### Value

a named vector of GC4d values.

```
# estimate GC4d of yeast genes
cf_all <- count_codons(yeast_cds)
gc4d <- get_gc4d(cf_all)
head(gc4d)
hist(gc4d)</pre>
```

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get\_tai

Calculate TAI

## **Description**

```
get_tai calculates tRNA Adaptation Index (TAI) of each CDS
```

## Usage

```
get_tai(cf, trna_w)
```

## **Arguments**

cf matrix of codon frequencies as calculated by 'count\_codons()'.

trna\_w tRNA weight for each codon, can be generated with 'est\_trna\_weight()'.

#### Value

a named vector of TAI values

#### References

dos Reis M, Savva R, Wernisch L. 2004. Solving the riddle of codon usage preferences: a test for translational selection. Nucleic Acids Res 32:5036-5044.

## **Examples**

```
# calculate TAI of yeast genes based on genomic tRNA copy numbers
w <- est_trna_weight(yeast_trna_gcn)
cf_all <- count_codons(yeast_cds)
tai <- get_tai(cf_all, w)
head(tai)
hist(tai)</pre>
```

human\_mt

human mitochondrial CDS sequences

## **Description**

CDSs of 13 protein-coding genes in the human mitochondrial genome extracted from ENSEMBL Biomart

#### Usage

human\_mt

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## **Format**

```
a DNAStringSet of 13 sequences
```

# Source

<a href="https://www.ensembl.org/index.html">https://www.ensembl.org/index.html</a>

# **Examples**

```
head(human_mt)
```

plot\_ca\_pairing

Plot codon-anticodon pairing relationship

## **Description**

```
plot_ca_pairing show possible codon-anticodons pairings
```

# Usage

```
plot_ca_pairing(codon_table = get_codon_table(), plot = TRUE)
```

# Arguments

```
codon_table a table of genetic code derived from 'get_codon_table' or 'create_codon_table'.

plot whether to plot the pairing relationship
```

# Value

a data.table of codon info and RSCU values

```
ctab <- get_codon_table(gcid = '2')
pairing <- plot_ca_pairing(ctab)
head(pairing)</pre>
```

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rev\_comp

Reverse complement

# Description

rev\_comp creates reverse complemented version of the input sequence

## Usage

```
rev_comp(seqs)
```

# Arguments

seqs

input sequences, DNAStringSet or named vector of sequences

#### Value

reverse complemented input sequences as a DNAStringSet.

## **Examples**

```
# reverse complement of codons
rev_comp(Biostrings::DNAStringSet(c('TAA', 'TAG')))
```

seq\_to\_codons

Convert CDS to codons

# Description

seq\_to\_codons converts a coding sequence to a vector of codons

# Usage

```
seq_to_codons(seq)
```

## **Arguments**

seq

DNAString, or an object that can be coerced to a DNAString

#### Value

a character vector of codons

show\_codon\_tables 21

#### **Examples**

```
# convert a CDS sequence to a sequence of codons
seq_to_codons('ATGTGGTAG')
seq_to_codons(yeast_cds[[1]])
```

show\_codon\_tables

show available codon tables

## **Description**

show\_codon\_tables print a table of available genetic code from NCBI through 'Biostrings::GENETIC\_CODE\_TABLE'.

## Usage

```
show_codon_tables()
```

#### Value

No return value (NULL). Available codon tables will be printed out directly.

## **Examples**

```
# print available NCBI codon table IDs and descriptions.
show_codon_tables()
```

slide

slide window interval generator

## **Description**

slide generates a data.table with start, center, and end columns for a sliding window analysis.

#### Usage

```
slide(from, to, step = 1, before = 0, after = 0)
```

# **Arguments**

from integer, the start of the sequence to integer, the end of the sequence

step integer, the step size

before integer, the number of values before the center of a window after integer, the number of values after the center of a window

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## Value

data.table with start, center, and end columns

# Examples

```
slide(1, 10, step = 2, before = 1, after = 1)
```

slide\_apply

apply a cub index to a sliding window

# Description

slide\_apply applies a function to a sliding window of codons.

# Usage

```
slide_apply(seq, .f, step = 1, before = 0, after = 0, ...)
```

# Arguments

seq	DNAString, the sequence
. f	function, the codon index calculation function to apply, for example, 'get_enc'.
step	integer, the step size in number of codons
before	integer, the number of codons before the center of a window
after	integer, the number of codons after the center of a window
	additional arguments to pass to the function '.f'

## Value

data.table with start, center, end, and codon usage index columns

```
slide_apply(yeast_cds[[1]], get_enc, step = 1, before = 10, after = 10)
```

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slide_codon	sliding window of codons

## **Description**

slide\_codon generates a data.table with start, center, and end columns for a sliding window analysis of codons.

## Usage

```
slide_codon(seq, step = 1, before = 0, after = 0)
```

## **Arguments**

seq	DNAString, the sequence
step	integer, the step size
before	integer, the number of codons before the center of a window
after	integer, the number of codons after the center of a window

#### Value

data.table with start, center, and end columns

## **Examples**

```
x \leftarrow Biostrings::DNAString('ATCTACATAGCTACGTAGCTCGATGCTAGCATCGTACGATCGTAGC') slide_codon(x, step = 3, before = 1, after = 1)
```

|--|--|--|

# Description

slide\_plot visualizes codon usage in sliding window.

## Usage

```
slide_plot(windt, index_name = "Index")
```

## Arguments

windt data.table, the sliding window codon usage generated by 'slide\_apply'.

index\_name character, the name of the index to display.

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## Value

```
ggplot2 plot.
```

#### **Examples**

```
sw <- slide_apply(yeast_cds[[1]], get_enc, step = 1, before = 10, after = 10)
slide_plot(sw)</pre>
```

yeast\_cds

yeast CDS sequences

# Description

CDSs of all protein-coding genes in Saccharomyces\_cerevisiae

#### Usage

yeast\_cds

#### **Format**

a DNAStringSet of 6600 sequences

#### Source

<a href="https://ftp.ensembl.org/pub/release-107/fasta/saccharomyces\_cerevisiae/cds/Saccharomyces\_cerevisiae.R64-1-1.cds.all.fa.gz">https://ftp.ensembl.org/pub/release-107/fasta/saccharomyces\_cerevisiae/cds/Saccharomyces\_cerevisiae.R64-1-1.cds.all.fa.gz</a>

## **Examples**

```
head(yeast_cds)
```

yeast\_exp

yeast mRNA expression levels

## **Description**

Yeast mRNA FPKM determined from rRNA-depleted (RiboZero) total RNA-Seq libraries. RUN1 $_0$ WT and RUN2 $_0$ WT (0 min after RNA Pol II repression) were averaged and used here.

## Usage

```
yeast_exp
```

yeast\_half\_life 25

#### **Format**

a data.frame with 6717 rows and three columns:

```
gene_id gene ID
gene_name gene name
fpkm mRNA expression level in Fragments per kilobase per million reads
```

#### **Source**

<a href="https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE57385">https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE57385></a>

#### References

Presnyak V, Alhusaini N, Chen YH, Martin S, Morris N, Kline N, Olson S, Weinberg D, Baker KE, Graveley BR, et al. 2015. Codon optimality is a major determinant of mRNA stability. Cell 160:1111-1124.

## **Examples**

```
head(yeast_exp)
```

yeast\_half\_life

Half life of yeast mRNAs

## **Description**

Half life of yeast mRNAs in Saccharomyces\_cerevisiae calculated from rRNA-deleted total RNAs by Presnyak et al.

## Usage

```
yeast_half_life
```

#### **Format**

```
a data.frame with 3888 rows and three columns:
```

```
gene_id gene id
gene_name gene name
half_life mRNA half life in minutes
```

#### Source

```
<a href="https://doi.org/10.1016/j.cell.2015.02.029">https://doi.org/10.1016/j.cell.2015.02.029</a>
```

26 yeast\_trna

#### References

Presnyak V, Alhusaini N, Chen YH, Martin S, Morris N, Kline N, Olson S, Weinberg D, Baker KE, Graveley BR, et al. 2015. Codon optimality is a major determinant of mRNA stability. Cell 160:1111-1124.

## **Examples**

```
head(yeast_half_life)
```

yeast\_trna

yeast tRNA sequences

# Description

Yeast tRNA sequences obtained from gtRNAdb.

## Usage

```
yeast_trna
```

#### **Format**

a RNAStringSet with a length of 275.

#### Source

<a href="http://gtrnadb.ucsc.edu/genomes/eukaryota/Scere3/sacCer3-mature-tRNAs.fa">http://gtrnadb.ucsc.edu/genomes/eukaryota/Scere3/sacCer3-mature-tRNAs.fa</a>

#### References

Chan PP, Lowe TM. 2016. GtRNAdb 2.0: an expanded database of transfer RNA genes identified in complete and draft genomes. Nucleic Acids Res 44:D184-189.

```
yeast_trna
```

yeast\_trna\_gcn 27

yeast\_trna\_gcn

yeast tRNA gene copy numbers (GCN)

# Description

Yeast tRNA gene copy numbers (GCN) by anticodon obtained from gtRNAdb.

# Usage

```
yeast_trna_gcn
```

#### **Format**

a named vector with a length of 41. Value names are anticodons.

# Source

<a href="http://gtrnadb.ucsc.edu/genomes/eukaryota/Scere3/sacCer3-mature-tRNAs.fa">http://gtrnadb.ucsc.edu/genomes/eukaryota/Scere3/sacCer3-mature-tRNAs.fa</a>

## References

Chan PP, Lowe TM. 2016. GtRNAdb 2.0: an expanded database of transfer RNA genes identified in complete and draft genomes. Nucleic Acids Res 44:D184-189.

```
yeast_trna_gcn
```

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