# Package 'iMKT'

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Title McDonald and Kreitman Test and its extensions calculation
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<b>Description</b> McDonald and Kreitman Test and its extensions.
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### **Description**

asymptoticMK developed in "Haller BC, Messer PW. asymptoticMK: A Web-Based Tool for the Asymptotic McDonald-Kreitman Test. G3 (Bethesda). 2017 May 5;7(5):1569-1575". Adapted from: http://github.com/MesserLab/asymptoticMK

### Usage

```
asymptoticMK(daf, divergence, xlow, xhigh, seed)
```

### **Arguments**

daf	data frame containing DAF, Pi and P0 values
divergence	data frame containing divergent and analyzed sites for selected (i) and neutral (0) classes
xlow	lower limit for asymptotic alpha fit
xhigh	higher limit for asymptotic alpha fit
seed	seed value (optional). No seed by default

### **Details**

The standard McDonald and Kreitman test (MKT) is used to detect the signature of selection at the molecular level. The MKT compares the amount of variation within a species (polymorphism, P) to the divergence (D) between species at two types of sites, one of which is putatively netral and used as the reference to detect selection at the other type of site. In the standard MKT, these sites are synonymous (putatively neutral, 0) and non-synonymous sites (selected sites, i) in a coding region. Under strict neutrality, the ratio of the number of selected and neutral polymorphic sites (Pi/P0) is equal to the ratio of the number of selected and neutral divergence sites (Di/D0).

### Value

Estimation of alpha asymptotic value and details of the model fit

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### **Examples**

```
asymptoticMK(mydafdata, mydivergencedata, 0, 0.9)
```

check\_input check\_input

### **Description**

Check input data and return detailed errors when it is malformed.

### Usage

```
check_input(daf, divergence, xlow, xhigh)
```

### **Arguments**

daf data frame containing DAF, Pi and P0 values

divergence data frame containing divergent and analyzed sites for selected (i) and neutral

(0) classes

xlow lower limit for asymptotic alpha fit xhigh higher limit for asymptotic alpha fit

### **Details**

This function checks input data used in most package's functions (arguments daf, divergence, xlow and xhigh) and returns a brief description of the error(s) found. If data does not pass check\_input() the requested analysis is not performed.

#### Value

None

completeMKT completeMKT

### Description

```
completeMKT() put details here
```

```
completeMKT(daf, divergence, xlow, xhigh, seed)
```

DGRP

### Arguments

daf data frame containing DAF, Pi and P0 values

divergence data frame containing divergent and analyzed sites for selected (i) and neutral

(0) classes

xlow lower limit for asymptotic alpha fit
xhigh higher limit for asymptotic alpha fit

seed seed value (optional). No seed by default

#### **Details**

put description here

### Value

Execute all the MKT extensions

### **Examples**

```
completeMKT(mydafdata, mydivergencedata, 0, 0.9)
```

DGRP DGRP

### Description

DGRP() Perform MKT corrected with DGRP method

### Usage

```
DGRP(daf, divergence, list_cutoffs = c(0, 0.05, 0.2), plot = FALSE)
```

### **Arguments**

daf data frame containing DAF, Pi and P0 values

divergence data frame containing divergent and analyzed sites for selected (i) and neutral

(0) classes

list\_cutoffs list of cutoffs to use (optional). Default cutoffs are: 0, 0.05, 0.2

plot report plot (optional). Default is FALSE

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#### **Details**

The standard McDonald and Kreitman test (MKT) is used to detect the signature of selection at the molecular level. The MKT compares the amount of variation within a species (polymorphism, P) to the divergence (D) between species at two types of sites, one of which is putatively netral and used as the reference to detect selection at the other type of site. In the standard MKT, these sites are synonymous (putatively neutral, 0) and non-synonymous sites (selected sites, i) in a coding region. Under strict neutrality, the ratio of the number of selected and neutral polymorphic sites (Pi/P0) is equal to the ratio of the number of selected and neutral divergence sites (Di/D0). The null hypothesis of neutrality is rejected in a MKT when Di/D0 > Pi/P0. The excess of divergence relative to polymorphism for class i, is interpreted as adaptive selection for a subset of sites i. The fraction of adaptive fixations, alpha.symbol, is estimated from 1-(Pi/P0)(D0/Di). The significance of the test can be assessed with a Fisher exact test. The estimate of alpha.symbol can be easily biased by the segregation of slightly deleterious non-synonymous substitutions. Specifically, slightly deleterious mutations tend to contribute more to polymorphism than to divergence, and thus, lead to an underestimation of alpha. Because adaptive mutations and weakly deleterious selection sct in opposite directions on the MKT, alpha and the fraction of substitutions that are sligholty deleterious, b, will be both underestimated when the two selection regimes occur. To take adaptive and slighlty deleterious mutations mutually into account, Pi, the count off segregatning sites in class i, should be seaprated into the number of neutral variants and the number of weakly deleterious variants, Pi = Pineutral + Pi weak del. Alpha is then estimated as 1-(Pineutral/P0)(D0/Di)

#### Value

MKT corrected by the DGRP method. List with alpha results, graph, divergence metrics, MKT tables and negative selection fractions

### **Examples**

```
## Using default cutoffs
DGRP(mydafdata, mydivergencedata)
## Using custom cutoffs
DGRP(mydafdata, mydivergencedata, c(0.05, 0.1, 0.15))
```

FWW FWW

### **Description**

FWW() Perform MKT corrected with FWW method

```
FWW(daf, divergence, list_cutoffs = c(0, 0.05, 0.1), plot = FALSE)
```

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### Arguments

daf data frame containing DAF, Pi and P0 values

divergence data frame containing divergent and analyzed sites for selected (i) and neutral

(0) classes

list\_cutoffs list of cutoffs to use (optional). Default cutoffs are: 0, 0.05, 0.2

plot report plot (optional). Default is FALSE

#### **Details**

The standard McDonald and Kreitman test (MKT) is used to detect the signature of selection at the molecular level. The MKT compares the amount of variation within a species (polymorphism, P) to the divergence (D) between species at two types of sites, one of which is putatively netral and used as the reference to detect selection at the other type of site. In the standard MKT, these sites are synonymous (putatively neutral, 0) and non-synonymous sites (selected sites, i) in a coding region. Under strict neutrality, the ratio of the number of selected and neutral polymorphic sites (Pi/P0) is equal to the ratio of the number of selected and neutral divergence sites (Di/D0). The null hypothesis of neutrality is rejected in a MKT when Di/D0 > Pi/P0. The excess of divergence relative to polymorphism for class i, is interpreted as adaptive selection for a subset of sites i. The fraction of adaptive fixations, alpha.symbol, is estimated from 1-(Pi/P0)(Ds/Dn). The significance of the test can be assesed with a Fisher exact test. The estimate of alpha. symbol can be easily biased by the segregation of slightly deleterious non-synonymous substitutions. Specifically, slightly deleterious mutations tend to contribute more to polymorphism than to divergence, and thus, lead to an underestimation of alpha. Bevause they tend to segregate at lower frequencies than do neutral mutations, they can be apartially controlled for by removing low frequency polymorphisms from the analysis (Fay et al. 2001). This is known as the FWW method.

#### Value

MKT corrected by the FWW method

#### **Examples**

```
## Using default cutoffs
FWW(mydafdata, mydivergencedata)
## Using custom cutoffs and rendering plot
FWW(mydafdata, mydivergencedata, c(0.05, 0.1, 0.15), plot=TRUE)
```

iMK iMK

#### **Description**

iMK: alpha asymptotic + negative selection (d,b,f)

#details details here

loadPopFly 7

### **Usage**

```
iMK(daf, divergence, xlow, xhigh, seed, plot = FALSE)
```

### **Arguments**

daf data frame containing DAF, Pi and P0 values

divergence data frame containing divergent and analyzed sites for selected (i) and neutral

(0) classes

xlow lower limit for asymptotic alpha fit
xhigh higher limit for asymptotic alpha fit
seed seed value (optional). No seed by default

plot report plots of daf, alpha and negative selection fractions (optional). Default is

**FALSE** 

### Value

iMK

### **Examples**

```
## Without plot
iMK(mydafdata, mydivergencedata, 0, 0.9)
## With plot
iMK(mydafdata, mydivergencedata, 0, 0.9, plot=TRUE)
```

loadPopFly loadPopFly

### Description

Load PopFly dataset

### Usage

loadPopFly()

### **Details**

This function loads PopFly data (http://popfly.uab.cat/) into the current workspace. Data is stored in a dataframe named PopFlyData.

### Value

None

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### **Examples**

```
# loadPopFly()
```

loadPopHuman

loadPopHuman

### **Description**

Load PopHuman dataset

### Usage

loadPopHuman()

### **Details**

This function loads PopHuman data (http://pophuman.uab.cat/) into the current workspace. Data is stored in a dataframe named PopFlyData.

### Value

None

### Examples

# loadPopHuman()

multipleDatasets

multipleDatasets

### Description

Perform any MK test using all files (or a subset of them) in a given directory.

```
multipleDatasets(directory = directory, test = c("standard", "DGRP", "FWW",
    "asymptotic", "iMK"), xlow = 0, xhigh = 1, fullAnalysis = TRUE/FALSE,
    idList = "NA")
```

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### **Arguments**

directory directory (path/to/files/) where daf and divergence files are stored in your local

machine

test which test to perform. Options include: standard (default), DGRP, FWW, asymp-

totic, iMK

xlow lower limit for asymptotic alpha fit (default=0)
xhigh higher limit for asymptotic alpha fit (default=1)

fullAnalysis decide whether to analyze all files in directory or not (default=TRUE)

idList used when fullAnalysis = F, list of IDs to analyze

### **Details**

Files in directory must be named: file1\*daf\*, file1\*divergence\*, file2\*daf\*, file2\*divergence\*, ...

#### Value

None

### **Examples**

##example here

mydafdata

Example data frames

### **Description**

Containing basic information to execute the funcionts

- price. price in US dollars (\\$326-\\$18,823)
- carat. weight of the diamond (0.2–5.01)
- cut. quality of the cut (Fair, Good, Very Good, Premium, Ideal)
- colour. diamond colour, from J (worst) to D (best)
- clarity. a measurement of how clear the diamond is (I1 (worst), SI1, SI2, VS1, VS2, VVS1, VVS2, IF (best))
- x. length in mm (0–10.74)
- y. width in mm (0–58.9)
- z. depth in mm (0–31.8)
- depth. total depth percentage = z / mean(x, y) = 2 \* z / (x + y) (43-79)
- table. width of top of diamond relative to widest point (43–95)

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### **Usage**

mydafdata

### **Format**

A data frame with x rows and y variables

mydivergencedata

Example data frames

### **Description**

Containing basic information to execute the funcionts

- price. price in US dollars (\\$326-\\$18,823)
- carat. weight of the diamond (0.2–5.01)
- cut. quality of the cut (Fair, Good, Very Good, Premium, Ideal)
- colour. diamond colour, from J (worst) to D (best)
- clarity. a measurement of how clear the diamond is (I1 (worst), SI1, SI2, VS1, VS2, VVS1, VVS2, IF (best))
- x. length in mm (0–10.74)
- y. width in mm (0–58.9)
- z. depth in mm (0–31.8)
- depth. total depth percentage = z / mean(x, y) = 2 \* z / (x + y) (43-79)
- table. width of top of diamond relative to widest point (43–95)

### Usage

mydivergencedata

### **Format**

A data frame with x rows and y variables

PopFlyAnalysis 11

|--|

### **Description**

Perform any MK test using a subset of PopFly data defined by custom genes and populations lists

### Usage

```
PopFlyAnalysis(genes = c("gene1", "gene2", "..."), pops = c("pop1", "pop2", "..."), recomb = TRUE/FALSE, bins = 0, test = c("standard", "DGRP", "FWW", "asymptotic", "iMK"), xlow = 0, xhigh = 1)
```

### Arguments

genes	list of genes
pops	list of populations
recomb	group genes according to recombination values (must specify number of bins). TRUE/FALSE. Recomb values (cM/Mb) from Comeron et al. 2012.
bins	number of recombination bins to compute (mandatory if recomb = TRUE)
test	which test to perform. Options include: standard (default), DGRP, FWW, asymptotic, iMK $$
xlow	lower limit for asymptotic alpha fit (default=0)
xhigh	higher limit for asymptotic alpha fit (default=1)

### **Details**

Recombination values (recomb=T) from Comeron et al. 2012 (reference!)

#### Value

None

### **Examples**

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standard

mkt\_standard

### **Description**

```
mkt_standard() MKT standard formula
```

### Usage

```
standard(daf, divergence)
```

### **Arguments**

daf data frame containing DAF, Pi and P0 values

divergence data frame containing divergent and analyzed sites for selected (i) and neutral

(0) classes

#### **Details**

The standard McDonald and Kreitman test (MKT) is used to detect the signature of selection at the molecular level. The MKT compares the amount of variation within a species (polymorphism, P) to the divergence (D) between species at two types of sites, one of which is putatively netral and used as the reference to detect selection at the other type of site. In the standard MKT, these sites are synonymous (putatively neutral, 0) and non-synonymous sites (selected sites, i) in a coding region. Under strict neutrality, the ratio of the number of selected and neutral polymorphic sites (Pi/P0) is equal to the ratio of the number of selected and neutral divergence sites (Di/D0).

### Value

Standard McDonald and Kreitman Test

### **Examples**

standard(mydafdata, mydivergencedata)

subsetPopData

subsetPopData

### **Description**

Perform iMK using a subset of PopFly or PopHuman data defined by custom genes and populations lists

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### Usage

```
subsetPopData(data = c("PopFly", "PopHuman"), genes = c("gene1", "gene2",
   "..."), pops = c("pop1", "pop2", "..."), recomb = TRUE/FALSE, bins = 0,
   test = c("standard", "DGRP", "FWW", "asymptotic", "iMK"), xlow = 0,
   xhigh = 1)
```

### Arguments

input PopFly or PopHuman data data list of genes genes list of populations pops recomb group genes according to recombination values (must specify number of bins). TRUE/FALSE bins number of recombination bins to compute (mandatory if recomb = TRUE) which test to perform. Options include: standard (default), DGRP, FWW, iMK, test asymptotic, none. lower limit for asymptotic alpha fit (default=0) xlow

xlow lower limit for asymptotic alpha fit (default=0) xhigh higher limit for asymptotic alpha fit (default=1)

#### **Details**

put details here

#### Value

None

### **Examples**

```
## Load PopFly data
# loadPopFly()
## Perform analysis
# mygenes <- c("FBgn0053196", "FBgn0000008")
# subsetPopData("PopFly", mygenes , c("RAL","ZI"), recomb=F)</pre>
```

theme\_Publication

ggplot theme for publication ready Plots

### **Description**

```
Date = 04/07/2015 Author = Koundinya Desiraju
```

```
theme_Publication(base_size = 14, base_family = "sans")
```

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### Arguments

base\_size base size required from theme\_Publication

base\_family font to load in theme\_Publication

### **Details**

Default theme used for plot images. From http://rpubs.com/Koundy/71792

### Value

plot theme

### Examples

```
# theme_Publication(14, "sans")
```

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