

# Package ‘svisits’

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**Type** Package

**Title** Shared Clinic Visit Dates Identify Steady HIV Positive Partnerships for Research

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**Description** This package helps to find some stable HIV-infected partnerships in cohort studies based on the assumption that some patients frequently attend the clinical visits together. These pairs are useful for biological and epidemiological research.

**License** GPL ( $\geq 2$ )

**Depends** R ( $\geq 2.10$ ), Rcpp ( $\geq 0.12.5$ ), compiler, parallel, chron, data.table

**Imports** Rcpp ( $\geq 0.12.5$ ), compiler, parallel, chron, data.table

**LinkingTo** Rcpp

**Suggests** knitr, rmarkdown

**VignetteBuilder** knitr

**NeedsCompilation** yes

## R topics documented:

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svisits-package	<i>svisits: finding HIV transmission and serosorting pairs using shared clinic visit dates</i>
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### Description

This package helps to find some stable HIV-infected partnerships in cohort studies based on the assumption that some patients frequently attend the clinical visits together. These pairs are useful for biological and epidemiological research.

### Author(s)

Alex Marzel, Teja Turk, Roger Kouyos

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adjust_Rcpp_min	<i>Adjustment function</i>
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### Description

This is the C++ adjustment function that is called by adjust\_visits. It adjusts the numbers of shared visits for each pair.

### Usage

```
adjust_Rcpp_min(mat, prob)
```

### Arguments

mat	matrix with 3 columns (unadjusted number of shared visits, number of visits of the first member, number of visits of the second member) to adjust
prob	probability of sharing a single visit, default 1/75

### Details

written in C++

### Value

a vector of adjusted number of shared visits

### Examples

```
visits <- data.frame(Freq = c(1,1,1,2,3),
                     N_visits.x = c(46,39,10,40,55),
                     N_visits.y = c(68,68,68,24,24))
mat <- as.matrix(visits)
adj_mat <- adjust_Rcpp_min(mat, prob=1/75)
adj_mat
```

---

adjust\_R\_Rcpp\_short\_binomial

*Probabilites of sharing a given number of visits*


---

### Description

This is the C++ function that is called by Bonferroni\_m.

### Usage

```
adjust_R_Rcpp_short_binomial(mat, prob)
```

### Arguments

mat	matrix with 3 columns (unadjusted number of shared visits, number of visits of the first member, number of visits of the second member)
prob	probability of sharing a single visit, default 1/75

### Details

written in C++

### Value

a vector of probabilities to share the given number of visits for each pair

### Examples

```
visits <- data.frame(Freq = c(1,1,1,2,3),
                     N_visits.x = c(46,39,10,40,55),
                     N_visits.y = c(68,68,68,24,24))
mat <- as.matrix(visits)
Bonferroni_mat <- adjust_R_Rcpp_short_binomial(mat, prob=1/75)
Bonferroni_mat
```

---

adjust\_visits

*Adjust the number of shared visits*


---

### Description

The function adjusts the total number of shared visits per pair.

### Usage

```
adjust_visits(unadjusted_pairs, ids, prob = 1/75)
```

**Arguments**

unadjusted_pairs	a "table" object containing the pairs and the number of shared visits
ids	a "data.table" object containing columns "ids" and "N_visits". The "ids" column represents patient identifiers and is the reference (see setkey from package data.table for more details). The "N_visits" contains the number of visits for each patient.
prob	probability of sharing a single visit, depending on the frequency of visits. Default 1/75~visit every three months.

**Value**

a data frame with the columns:

"allPairs" pair identifiers  
 "Freq" unadjusted number of shared visits  
 "id\_1" first member of the pair's id  
 "id\_2" second member of the pair's id  
 "N\_visits.x" first member of the pair's number of visits  
 "N\_visits.y" second member of the pair's number of visits  
 "Corrected" the adjusted number of shared visits per pair

**Examples**

```
# load data
data("simulated_data")
db_dates <- prepare_db(your_database = simulated_data,
                      ids_column = "subject",
                      dates_column = "sim_dates")

# first get unadjusted pairs
unadjusted_observed_pairs <- get_observed_pairs(db_dates)
ids <- data.table(ids = as.character(names(table(db_dates$subject))),
                 N_visits = as.character(as.numeric(table(db_dates$subject))))
setkey(ids, "ids")

# now adjust the pairs
adjusted_observed <- adjust_visits(unadjusted_pairs = unadjusted_observed_pairs,
                                ids = ids,
                                prob = 1/75)

head(adjusted_observed)
```

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barcode

*Unique pair identifier*


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**Description**

It makes a unique identifier for each pair. Each member's id should be numeric! For example for id\_1=6758 and id\_2=3456 the barcode will be "3456\_6578".

**Usage**

```
barcode(raw_for_eval)
```

**Arguments**

`raw_for_eval` a vector with two ids

**Value**

string of a form id1\_id2

**Examples**

```
barcode(c(id_2=6578,id_1=3456))
barcode(c(3456, 6578))
```

---

Bonferroni\_m

*Detecting the pairs using Bonferroni correction*

---

**Description**

This method is much faster than the shuffling method, but less precise.

**Usage**

```
Bonferroni_m(unadjusted_pairs,
             ids=ids,
             prob = 1/75,
             alpha = 0.01,
             only_significant = TRUE)
```

**Arguments**

`unadjusted_pairs` a "table" object containing the pairs and the number of shared visits. See [get\\_observed\\_pairs](#).

`ids` a "data.table" object with columns "ids" and "N\_visits". The "ids" column represents patient identifiers and is the reference (see `setkey` from package `data.table` for more details). The "N\_visits" contains the number of visits for each patient.

`prob` probability of sharing a single visit. Default 1/75.

`alpha` type I error

`only_significant` returns only significant pairs above the threshold (default) or all the pairs

**Value**

a data frame with

"allPairs" pair identifier

"Freq" the unadjusted number of shared visits

"id\_1" identifier of the first pair member

"id\_2" identifier of the second pair member

"N\_visits.x" total number of visits of the first pair member

"N\_visits.y" total number of visits of the second pair member

"Prob\_for\_Bonferr" probability of sharing the given number of shared visits

"BP" row number

"ltp" the log-transformed probability of sharing the given number shared visits

**Examples**

```
# load data
data("simulated_data")
db_dates <- prepare_db(your_database = simulated_data,
                      ids_column = "subject",
                      dates_column = "sim_dates")

# first get unadjusted pairs
unadjusted_observed_pairs <- get_observed_pairs(db_dates)
# prepare ids
ids <- data.table(ids = as.character(names(table(db_dates$subject))),
                  N_visits = as.character(as.numeric(table(db_dates$subject))))
setkey(ids, "ids")

# run
Bonferroni_m_output <- Bonferroni_m(unadjusted_observed_pairs,
                                   ids = ids, prob = 1/75, alpha = 0.01)

# number of significant pairs
nrow(Bonferroni_m_output)
```

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chooseC

*Binomial coefficient*

---

**Description**

It calculates the binomial coefficient.

**Usage**

```
chooseC(n, k)
```

**Arguments**

n	set size
k	number of unordered outcomes

**Details**

implemented in C++

**Value**

numeric

**Examples**

```
chooseC(n=6, k=2)
```

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find_pairs	<i>Find significant pairs</i>
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**Description**

Find significant pairs using the Bonferroni method and (optionally) with the shuffling algorithm to avoid False-Positives. This is the wrap-up of the following functions:

- [prepare\\_db](#)
- [get\\_observed\\_pairs](#)
- [Bonferroni\\_m](#)
- [adjust\\_visits](#) (only if the shuffling algorithm is performed)
- [Shuffling\\_simulation](#) (optional)

**Usage**

```
find_pairs(your_database,
           ids_column,
           dates_column,
           prob = 1/75,
           alpha = 0.01,
           only_significant = TRUE,
           shuffling = FALSE,
           n_shuffl = 100,
           period_months = 3,
           FP_threshold = max,
           max_threshold = NULL)
```

**Arguments**

your_database	a data frame with the clinic visit dates in a long format.
ids_column	character, column name with the person identifiers
dates_column	character, column name with the visit dates
prob	probability of sharing a single visit, default 1/75
alpha	type I error
only_significant	returns only the significant pairs above the threshold (default) or all the pairs with the corresponding p-values

shuffling	logical indicating if the shuffling should be performed. By default (FALSE) only the Bonferroni method is used.
n_shuffl	number of repetitions of shuffling algorithm (default 100)
period_months	the length of the follow-up period in months (by default 3)
FP_threshold	the function used to get the False-Positive fraction from the shuffling repetitions at each position. By default the max is called at each threshold. Another possibilities are mean or median.
max_threshold	maximum false-positive threshold to be inspected. If it is NULL, the 99.995%-quantile of the distribution of the unadjusted number of shared visits is used.

### Value

A list containing

"unadjusted\_pairs" pairs with unadjusted number of shared visits

"Bonferroni\_pairs" pairs obtained with the Bonferroni correction. If significant\_only==TRUE only those above the threshold are returned.

"adjusted\_pairs" pairs with adjusted number of shared visits returned only if shuffling==TRUE.

"Shuffled\_pairs" pairs obtained by the shuffling algorithm (returned only if shuffling==TRUE).

"shuffling\_threshold" threshold determined by the shuffling algorithm

"shuffling\_simulation\_output" output from the shuffling function if shuffling==TRUE

### See Also

[prepare\\_db](#), [get\\_observed\\_pairs](#), [Bonferroni\\_m](#), [adjust\\_visits](#), [Shuffling\\_simulation](#)

### Examples

```
data("simulated_data")
pairs <- find_pairs(simulated_data,
                    ids_column = "subject",
                    dates_column = "sim_dates",
                    shuffling = TRUE, n_shuffl = 2)
```

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getPairs

*Pairs with shared visit on a certain date*

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### Description

From a given list of patient with the clinic visit on a certain date the function returns all the pairs.

### Usage

```
getPairs(ids)
```

### Arguments

ids                      a vector of patient identifiers



**Details**

This function is called by [get\\_observed\\_pairs](#) which date-wise finds all the pairs with at least one shared visit (for all the dates that appear in your database).

**Value**

a character vector containing the [barcodes](#) of the found pairs on a certain date

**Examples**

```
dates_long <- data.frame(subject=c(12,15,2,202,31,3),
                        dates=as.Date(c(rep("2001-03-16", 5), "2011-03-17"))))
# subjects with a visit on "2001-03-16"
id_s <- dates_long$subject[which(dates_long$dates==as.Date("2001-03-16"))]
getPairs(id_s)
```

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get_observed_pairs	<i>Pairs with at least one shared visit</i>
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**Description**

Find all the pairs that shared at least a single visit.

**Usage**

```
get_observed_pairs(tested_db)
```

**Arguments**

tested_db	visits in long format with each row representing a visit date. See <a href="#">simulated_data</a> for an example. It should contain columns "subject" with (numeric) patient identifiers and "dates" with the dates of clinic visits.
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**Value**

a table object containing the number of shared visits for all pairs with at least one shared visit

**Examples**

```
# a short example (with artificial data)
dates_long <- data.frame(subject=c(12,15,2,202,31,15, 202),
                        dates=as.factor(c(rep("2001-03-16",5),rep("2011-06-20",2))))
get_observed_pairs(dates_long)

# example with the simulated dates based on the real distribution
# load data
data("simulated_data")
db_dates <- prepare_db(your_database = simulated_data,
                      ids_column = "subject",
                      dates_column = "sim_dates")

# get unadjusted pairs
unadjusted_observed_pairs <- get_observed_pairs(db_dates)
```

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prepare_db	<i>Prepare the database</i>
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### Description

Transform the database to the appropriate format for finding pairs.

### Usage

```
prepare_db(your_database, ids_column, dates_column)
```

### Arguments

your_database	the data frame to be analyzed
ids_column	character, column name with the person identifiers
dates_column	character, column name with the visit dates

### Details

The dates are ordered in an increasing order within each subject.

### Value

a data frame with two columns:

"subject" person identifiers

"dates" dates of the clinic visits

### Examples

```
data("simulated_data")
db_dates<-prepare_db(your_database = simulated_data,
                     ids_column = "subject",
                     dates_column = "sim_dates")
```

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Shuffling_simulation	<i>Shuffling simulation to determine shared visits false-positive threshold</i>
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### Description

Shuffling simulation to determine shared visits false-positive threshold. It performs only one run. Please use "replicate" for several (see Example).

### Usage

```
Shuffling_simulation(db_dates, ids, adjusted_observed, max_threshold=NULL)
```

**Arguments**

<code>db_dates</code>	a data frame with visits to be shuffled (in long format) which column "fupdate-Period" containing the time period which each clinic visit belongs to.
<code>ids</code>	a "data.table" object with columns "ids" and "N_visits". The "ids" column represents patient identifiers and it is the reference (see <code>setkey</code> from package <code>data.table</code> for more details). The "N_visits" contains the number of visits for each patient.
<code>adjusted_observed</code>	a data frame with adjusted observed number of visits.
<code>max_threshold</code>	a maximum false-positive threshold to be inspected. If it is NULL, the 99.995%-quantile of the distribution of the unadjusted number of shared visits is used.

**Details**

Due to the multiple comparisons problem and strongly simplifying assumptions about the uniform distribution of visits, the corrected number of shared visits is not an optimal test statistic. Instead, the visits are first shuffled within each period (such that the original distribution of the number of visits per individual is preserved) and the number of randomly collided shared visits per pair is counted and penalized using "adjust\_visits". In other words, the observed visit dates from a given period (by default a quarter) are randomly re-assigned between the patients that attended the clinic during this period.

Define the number of desired shuffling simulations (ideally > 100, but it will take some time) and run.

**Value**

a numeric vector of size `max_threshold`. Each value represents the false positive fraction for a cutoff of adjusted shared visits that corresponds to the position (index) of the value in the vector.

**Examples**

```
# load data
data("simulated_data")
db_dates <- prepare_db(your_database = simulated_data,
                      ids_column = "subject",
                      dates_column = "sim_dates")

# prepare 3 months periods for later shuffling
db_dates <- cbind(db_dates, month.day.year(db_dates$dates))
periodMo <- 3
db_dates <- cbind(db_dates,
                  fupdatePeriod=with(db_dates, year+round(floor((month-1)/periodMo)
                                                         *periodMo/12,
                                                         digits=2)))

# first get unadjusted pairs
unadjusted_observed_pairs <- get_observed_pairs(db_dates)
ids <- data.table(ids=as.character(names(table(db_dates$subject))),
                  N_visits=as.character(as.numeric(table(db_dates$subject))))
setkey(ids, "ids")

# now adjust the pairs
adjusted_observed <- adjust_visits(unadjusted_pairs = unadjusted_observed_pairs,
```

```

                                ids = ids,
                                prob = 1/75)
# number of simulations
n_shuffl <- 2 # two
Shuffling_simulation_output <- replicate(n_shuffl,
                                         Shuffling_simulation(db_dates,
                                                             ids=ids,
                                                             adjusted_observed),
                                         simplify=FALSE)
head(Shuffling_simulation_output)

```

---

simulated\_data

*Simulated dataset of visit dates based on the SHCS*


---

### Description

It contains 20 pseudo transmission pairs that should be detected with the method.

### Usage

```
data("simulated_data")
```

### Format

A data frame with 90266 observations on the following 3 variables:

X row's indexing  
 subject a numeric vector with patient identifiers  
 sim\_dates a factor with the visit dates

### Examples

```
data("simulated_data")
head(simulated_data)
```

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