# Package 'rEHR'

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add_to_database
-----------------

# Description

This function can be used to import a CPRD file or files into a SQLite database connection.

# Usage

```
add_to_database(db, files, table_name, dateformat = "%d/%m/%Y",
   yob_origin = 1800, practid = TRUE, filenames = FALSE)
```

# **Arguments**

db	a database connection object
files	a character vector of filenames to files to be imported
table_name	a name for the table to import to
dateformat	the format that dates are stored in the CPRD data. If this is wrong it won't break but all dates are likely to be NA
yob_origin	value to add yob values to to get actual year of birth (Generally 1800)
practid	logical should practice id variable be constructed from the patient ids?
filenames	logical should the filename be included as a variable?

#### **Details**

Will automatically unzip files before calling them in If practid is TRUE, a practid variable is constructed by converting the last 3 digits of the patient id (if supplied) to a numeric. If filenames is TRUE, source data filenames are included as a variable with the filetypes stripped away.

```
append_to_temp_table

Appends rows to a temporary table
```

# Description

This function checks if a table is a temp table and then adds to it based on the select statement The check is to maintain the integrity of the database

# Usage

```
append_to_temp_table(db, tab_name, columns, select_query)
```

# Arguments

db	a database connection
tab_name	the name of the temporary table being appended to
columns	character vector of columns in tab_name
select_query	SQL query for the selector

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build\_cohort

Converts a longitudinal data set from e.g. prev\_terms to a cohort dataset. The dataset has a single row for each patient and includes only patients in the numerator or denominator for whichever cohort type is chosen. Columns are added for start and end dates and for start and end times as integer differences from the cohort start date. A binary column is added for membership of the case group. All patients with start date > end date are removed from the dataset.

## **Description**

Converts a longitudinal data set from e.g. prev\_terms to a cohort dataset. The dataset has a single row for each patient and includes only patients in the numerator or denominator for whichever cohort type is chosen. Columns are added for start and end dates and for start and end times as integer differences from the cohort start date. A binary column is added for membership of the case group. All patients with start date > end date are removed from the dataset.

#### Usage

```
build_cohort(dat, cohort_type = c("incid", "prev"), cohort_start, cohort_end,
    diagnosis_start = "eventdate")
```

# **Arguments**

dat dataframe as output from a call to prev\_terms

cohort\_type character either 'incid' or 'prev'. This selects the numerators and denominators

for that type of cohort

cohort\_start ISO date character string for the start of the cohort

cohort\_end ISO date character string for the end of the cohort

diagnosis\_start

character string for the name of the diagnosis variable used to define the start dates, or NULL if the diagnosis date is not to be included in the definition of

start dates.

build\_date\_fn

Function to build start/enddate helper fuctions

# Description

This builds functions identical to qof\_years but they can be customised to the user's preferences

#### **Usage**

```
build_date_fn(start, end)
```

## Arguments

start list containing the offset in years, month and day as numerics for the start date end list containing the offset in years, month and day as numerics for the end date

clinical\_codes 5

#### Value

a function taking a year as an argument and returning a list of startdates and enddates

clinical\_codes

Clinical codes for 17 QOF conditions, smoking and HbA1c

#### **Description**

This data comes from http://www.clinicalcodes.org

# Usage

```
clinical_codes
```

#### **Format**

A dataframe with 1283 observations:

medcode Unique internal code for the medical term selected by the healthcare professionalreadcode Unique Read Code for the medical term selected by the healthcare professionaldesc Description of the code

**list** Name of the clinical code list this code belongs to

#### **Source**

```
http://www.clinicalcodes.org
```

compress

Compresses a dataframe to make more efficient use of resources

# **Description**

Converts date variables in a dataframe to integers Integers represent time in days from the supplied origin Converts specified numeric values to integer This function is useful for keeping file sizes down and is used by the to\_stata command to save to Stata files.

#### Usage

```
compress(dat, origin = "1970-01-01", format = "%Y-%m-%d",
  date_fields = c("eventdate", "sysdate", "lcd", "uts", "frd", "crd", "tod",
  "deathdate"), integer_fields = c("yob", "practid"))
```

# **Arguments**

```
dat a dataframe

origin ISO string representation of the dat of origin. default is UNIX start date

format character: format of the date string. Default is ISO standard

date_fields character vector of column names representing dates

integer_fields
```

character vector of column names that should be integers

#### Value

dataframe

convert\_dates

converts date fields from ISO character string format to R Date format

# **Description**

Date fields are determined by the date-fields element in the .ehr definition. Extra date fields can be added to the extras argument or by setting '.ehr\$date\_fields'.

#### Usage

```
convert_dates(dat, extras = NULL)
```

# Arguments

dat a dataframe

extras = a character vector of extra columns to convert or NULL for no extras

#### See Also

```
get_EHR_value set_EHR_value
```

```
cprd_uniform_hbalc_values
```

Transforms hba1c values to mmol/mol

# **Description**

This function converts hba1c and fructosamine values to comon mmol/mol values conversions: mmol/L, mg/dL, fructosamine Assumes doctor miscoding for values < 20 after previous transformations

#### Usage

```
cprd_uniform_hbalc_values(df)
```

# Arguments

df

a dataframe of hba1c test scores

#### **Details**

The input dataframe must include the data2, data3 and enttype columns and the enttype must equal either 275 (HbA1C) or 356 (Fructosamine)

#### Value

dataframe with all hba1c values recoded to mmol/mol

cut\_tv 7

cut\_tv

cut\_tv - Cuts a survival dataset on a time varying variable

## **Description**

Survival datasets often have time-varying covariates that need to be dealt with. For example a drug exposure may occur after the entry into the cohort and you are interested in how this might affect your outcome.

#### Usage

```
cut_tv(.data, entry, exit, cut_var, tv_name, cores = 1, id_var,
  on_existing = c("flip", "increment"))
```

## **Arguments**

.data	a dataframe
entry	name of the column in .data that defines entry time to cohort. Column must be numeric.
exit	name of the column in .data that defines exit time from cohort. Column must be numeric
cut_var	name of the column in .data that defines the time of the time-varying covariate event. Column must be numeric.
tv_name	name for the constructed time-varying covariate
cores	number of mc.cores to use.
id_var	name of the variable identifying individual cases
on_existing	see details for cutting behaviour

## **Details**

This function cuts up a dataset based on times supplied for the time-varying covariate. If there is already a variable for the time-varying covariate, you can chose to flip the existing values or increment them. This means the function can be called multiple times to, e.g. deal with drugs starting and stopping and also to deal with progression of treatment.

The function is faster than other cutting methods, does not require conversion to Lexis format, and can be parallelised for large datasets and chained with dply workflows. Arguments should not be quoted.

This function can deal with the following scenarios (see examples):

- "Binary chronic covariates" e.g. The time of diagnosis for a chronic (unresolvable) condition. This requires a single column variable of times from entry in the dataset
- "Binary covariates" e.g. times of starting and stopping medication. This requires more than one column variable in the dataset, one for each start or stop event. The state flips with each new change.
- "Incremental time-varying covariates" e.g. different stages of a condition. This requires a single column variable for each incremental stage
- "Any combination of the above"This is achieved by chaining multiple calls together

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#### **Examples**

```
# A simple example dataset to be cut
tv_test < -data.frame(id = 1:5, start = rep(0, 5), end = c(1000, 689, 1000, 874, 777),
                      event = c(0,1,0,1,1), drug_1 = c(NA, NA, NA, 340, 460),
                      drug_2 = c(NA, 234, 554, 123, NA),
                      drug_3_start = c(110, 110, 111, 109, 110),
                      drug_3\_stop = c(400, 400, 400, 400, 400),
                      stage_1 = c(300, NA, NA, NA, NA),
                      stage_2 = c(450, NA, NA, NA, NA))
# Binary chronic covariates:
tv_out1 <- cut_tv(tv_test, start, end, drug_1, id_var = id, drug_1_state)
tv_out1 <- cut_tv(tv_out1, start, end, drug_2, id_var = id, drug_2_state)
# Binary covariates:
tv_out3 <- cut_tv(tv_test, start, end, drug_3_start, id_var = id, drug_3_state)
tv_out3 <- cut_tv(tv_out3, start, end, drug_3_stop, id_var = id, drug_3_state)
# incremental covariates:
inc_1 <- cut_tv(tv_test, start, end, stage_1, id_var = id, disease_stage, on_existing = '</pre>
inc_1 <- cut_tv(inc_1, start, end, stage_2, id_var = id, disease_stage, on_existing = "ir
# Chaining combinations of the above
## Not run:
library(dplyr)
tv_all <- tv_test %>%
          cut_tv(start, end, drug_1, id_var = id, drug_1_state) %>%
          cut_tv(start, end, drug_2, id_var = id, drug_2_state) %>%
          cut_tv(start, end, drug_3_start, id_var = id, drug_3_state) %>%
          cut_tv(start, end, drug_3_stop, id_var = id, drug_3_state) %>%
          cut_tv(start, end, stage_1, id_var = id, disease_stage, on_existing = "inc") %
          cut_tv(start, end, stage_2, id_var = id, disease_stage, on_existing = "inc")
## End(Not run)
```

database

Wrapper for dbConnect

# **Description**

Connects to a SQLite database or creates one if it does not already exist

#### Usage

```
database (dbname)
```

#### **Arguments**

dbname

character name path to database file

## Details

If the '.sqlite' file extension is ommited from the dbname argument it is automatically added.

#### Value

SQLiteConnection object

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#### **Examples**

```
## Not run:
db <- database("mydb")
## End(Not run)</pre>
```

define\_EHR

Construct an EHR\_definition object.

## **Description**

This function creates the object that defines EHR simulations

# Usage

```
define_EHR(start_date = "1930-01-01", end_date = "2014-06-30",
  patient = list(num = 10000, comorbidity = list(codes = NULL, prevalence =
  NULL), sim_params = list(transfer_out_prob = 0.2, scale = 25000, weibull_shape
  = 1, censor_type = "noninformative", betas = list(conditions = NULL, gender =
  log(0.7), baseline = 0.01, transfer_out = 3.5))),
  consultation = list(per_year = 2, type = list(code = 0:1, prob = c(0.2,
  0.8))), clinical = list(mean_events = 1), referral = list(mean_events =
  0.1), therapy = list(mean_events = 1), practice = list(num = 100, regions
  = 13, imd_cats = 5, early_lcd_prob = 0.1, early_lcd_range = 2, uts_limit =
  as.Date("1998-01-01"), late_uts_prob = 0.5, late_uts_range = 3))
```

# **Arguments**

```
character date for earliest birthday of patients in EMR ("%Y-%m-%d")
end_date character date for latest data collection date ("%Y-%m-%d")

patient list of definitions for patient file (see details)

consultation list of definitions for consultation file (see details)

clinical list of definitions for clinical file (see details)

referral list of definitions for referral file (see details)

therapy list of definitions for therapy file (see details)

practice list of definitions for practice file (see details)
```

## **Details**

The arguments for this constructor function are complex:

## Patient element

This is a list containing:

- num number of patients in the EHR simulation
- Comorbidity a list containing:
  - codes a named list of data frames of clinical codes in www.clinicalcodes.org
     export format

10 definition\_search

- prevalence a named list of prevalences for the comorbidities
- sim\_params a list containing:
  - \* transfer\_out\_prob probability of a patient transferring out early
  - \* scale scaling up parameter for survival rates
  - \* weibull\_shape parameter for survival rates
  - \* censor\_type type of censoring employed by link {surv\_sims}
  - \* betas list of values for conditions, gender, baseline and transfer out hazards. The betas element is a named vector of log hazards of length == length(comorbidity\$codes)
- to do.

# **Description**

This function is used to build new definition lists based on medical definitions

# Usage

```
definition_search(def, medical_table = NULL, test_table = NULL,
   drug_table = NULL, lookup = NULL)
```

#### **Arguments**

```
def an object of class MedicalDefinition

medical_table
Dataframe lookup table of clinical codes

test_table dataframe lookup table of test codes

drug_table dataframe lookup table of medication product codes

lookup list containing elements: "codes", "terms", "tests", "drugs", "drugcodes" (see details)
```

#### **Details**

You may get an invalid multibyte string error, in which case, set fileEncoding="latin1" on read.delim when reading in the lookup tables Lookup tables are

#### **Examples**

drop\_all\_temp\_tables 11

```
drop_all_temp_tables
```

drops all temporary tables from the database

# Description

This is useful for temporary storage/memory management

# Usage

```
drop_all_temp_tables(db)
```

# **Arguments**

db

a database connection

drop\_temp\_table

Checks if a temporary table exists and then deletes if it does

# Description

Checks if a temporary table exists and then deletes if it does

# Usage

```
drop_temp_table(db, tab_name)
```

# **Arguments**

db a database connection

tab\_name character the name of the table of interest

ehr\_def

An example EHR\_definition object for defining parameters for simulating EHR data

# **Description**

An example EHR\_definition object for defining parameters for simulating EHR data

# Usage

```
ehr_def
```

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#### **Format**

object of type 'EHR\_definition' including the following elements

patient list of parameters defining the simulation of the patient table clinical list of parameters defining the simulation of the clinical table therapy list of parameters defining the simulation of the therapy table practice list of parameters defining the simulation of the patient table start\_date start\_date for patients - earliest possible birthday

end\_date Latest possible date in the ehr for data collection

entity

A sample of 6 clinical tests and meaures used in UK primary care

# Description

A sample of 6 clinical tests and meaures used in UK primary care

# Usage

entity

#### **Format**

A dataframe with 6 observations:

enttype Unique internal code for the entity term

description Description of the code

filetype The table in the EHR the entity applies to

category Category of clinical entity

data1 first data variable

data2 second data variable

data3 third data variable

data4 fourth data variable

# Source

```
http://www.clinicalcodes.org
```

expand\_string 13

expand\_string

Reads strings and expands sections wrapped in dotted parentheses

# **Description**

This is a kind of inverse of bquote

# Usage

```
expand_string(s, level = 3)
```

#### **Arguments**

```
s a string
```

level integer sets the parent frame level for evaluation

# **Examples**

```
a <- runif(10)
expand_string("The r code is .(a)")</pre>
```

```
export_definition_search
```

Exports definition searches to an excel file

#### **Description**

Exports definition searches to an excel file

# Usage

```
export_definition_search(definition_search, out_file)
```

# **Arguments**

# **Examples**

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export\_fn

Exports to a variety of formats based on the file type argument

## **Description**

Exports to a variety of formats based on the file type argument

#### Usage

```
export_fn(x, file, \dots)
```

# **Arguments**

x object to be exported

file character path to the file to be exported to

arguments to be passed to the export functions

File type is based on the file suffix and can be one of "txt", "csv", "rda", "dta". dta files use foreign::write.dta. If a match is not found, the file is written to

std.out

extract\_keywords

Function to extract rows from a lookup table based on keywords

# **Description**

This function can be used to build draft clinical code lists based on a clinical or product lookup table and a set of keywords.

#### Usage

```
extract_keywords(lookup, keywords, keyword_field = "desc")
```

# Arguments

lookup a dataframe containing a lookup table

keywords character vector containing the keyword terms to search for

keyword\_field

character identifying the field in the lookup table to be searched for keywords

#### **Details**

See www.clinicalcodes.org for clinical code lists that have been used in previous studies All keywords are collapsed together in an OR statement

#### Value

a data frame subsetted by keyword

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#### **Examples**

```
## Not run:
keywords <- c('oral ulceration', 'mouth ulceration', 'aphthous ulceration',
'oral aphthous ulceration','oral ulcer[s]?', 'mouth ulcer[s]?', 'aphthous ulcer[s]?',
'aphthous stomatitis', "stomatitis", "aphthae", 'oral aphthous stomatitis',
'oral aphthous ulcers', 'recurrent oral ulcers', 'recurrent mouth ulcers',
'recurrent oral aphthous ulcers', 'recurrent aphthous ulcers', 'recurrent aphthous stomatitis')
a <- extract_keywords(medical, keywords)

## End(Not run)</pre>
```

first\_events

Selects the earliest event grouped by patient

# Description

This function runs a select\_events() query and then groups by patient id and picks only the earliest event for each patient

#### Usage

```
first_events(db = NULL, tab, columns = "eventdate", where = NULL,
    sql_only = FALSE, group_column = "patid", date_column = "eventdate")
```

# **Arguments**

db	A database connection object
tab	the database table to extract from
columns	The other columns to be extracted
where	sting representation of the selection criteria
sql_only	logical should the function just return a string of the SQL query?
group_column	column to group by. Default is patid
date_column	the column to sort by. default is eventdate

#### Value

a dataframe or a string representing an sql query

## **Examples**

```
## Not run:
b1 <- first_events(db, tab = "Clinical", columns = c("eventdate", "medcode"),
where = "medcode %in% .(a$medcode)")
first_events(tab = "Clinical", columns = c("eventdate", "medcode"),
where = "medcode %in% c(1, 2, 3, 4)", sql_only = TRUE)
## End(Not run)</pre>
```

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flat\_files

Exports flat files from the database. One file per practice

#### **Description**

Exports flat files from the database. One file per practice

# Usage

```
flat_files(db, table = "Consultation", practice_table = "Practice", out_dir,
  file_type = c("txt", "csv", "rda", "dta"), ...)
```

# **Arguments**

```
db a database connection

table character the table to be exported

practice_table
the table that the practice definitions can be found

out_dir a directory to output to. This will be created if it does not already exist

file_type the type of file to be saved. This can be one of "txt", "csv", "rda", "dta".

... arguments to be passed to export_fn

Defaults to exporting consultation tables for use by match_on_index(). the full path to out_dir will be created if it does not already exist.
```

# See Also

```
match_on_index export_fn
```

```
get_EHR_attribute Return the value of an attribute in the .ehr environment
```

# Description

Return the value of an attribute in the .ehr environment

#### Usage

```
get_EHR_attribute(x = NULL)
```

#### **Arguments**

```
x an attribute name
```

# Examples

```
{
set_CPRD()
get_EHR_attribute()
get_EHR_attribute(patient_id)
}
```

get\_matches 17

get_matches	Find matched controls for a set of cases
-------------	--

# **Description**

This function will provide a set of matched controls for a given set of cases.

# Usage

```
get_matches(cases, control_pool, n_controls, match_vars, extra_vars,
  extra_conditions = NULL, cores = 1, track = TRUE,
  tracker = function(case_num) ".", method = c("incidence_density",
  "exact"), diagnosis_date = NULL)
```

### **Arguments**

cases	dataframe of cases	
control_pool	dataframe of potential controls to be used for matching	
n_controls	number of controls to match to each case	
match_vars	character vector of variables in the dataframes to be used to perform the matching	
extra_vars	character vector of other variables to be used in the matching to define other conditions	
extra_condit	ions	
	a character vector of length 1 defining further restrictions on matching	
cores	number of cpu cores to be used by multicore (windows users should leave set to 1)	
track	logical should a dot be printed to std.out for each case?	
tracker	function to track progress of the function (See details)	
method	The method of selection of controls (see details)	
diagnosis_date		
	character the name of the variable in the cases and control_pool datasets containing the date of diagnosis (or other event to base the IDM method on). If there is no diagnosis date for a patient, this should be represented by NA	

## **Details**

Setting method to "exact" means that the matched controls are removed from the control pool after each case has been matched. This makes this method not thread safe and so will only run on a single core (and more slowly). Setting method to "incidence\_density" is thread safe as the same controls can be used for more than one case. See Richardson (2004) Occup Environ Med 2004;61:e59 doi:10.1136/oem.2004.014472 for a description of IDS matching. Also see the introduction vignette. The tracker variable allows for different outputs to track the progress of the function. This is currently set to ouput a dot for every case matched. A function can be added to the argument For a more verbose tracking, e.g. to track number of cases, set tracker = function() paste0(case\_num, ",")

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```
head.SQLiteConnection
```

head for SQLiteConnection object

#### **Description**

If just a database connection is selected, returns a dataframe of table names If a table name is also supplied, the first n rows from this table are output

## Usage

```
## S3 method for class 'SQLiteConnection'
head(x, table = NULL, n = 6L, temp = FALSE,
...)
```

# **Arguments**

```
x A SQLiteConnection object
table character specifying a table
n integer: Number of rows to output
temp logical should the function list the temp tables
... Additional arguments
```

import\_CPRD\_data Imports all selected CPRD data into an sqlite database

# **Description**

This function can import from both cohorts downloaded via the CPRD online tool and CPRD GOLD builds

# Usage

```
import_CPRD_data(db, data_dir, filetypes = c("Additional", "Clinical",
   "Consultation", "Immunisation", "Patient", "Practice", "Referral", "Staff",
   "Test", "Therapy"), dateformat = "%d/%m/%Y", yob_origin = 1800,
   regex = "PET", recursive = TRUE, ...)
```

# Arguments

db	a database connection
data_dir	the directory containing the CPRD cohort data
filetypes	character vector of filetypes to be imported
dateformat	the format that dates are stored in the CPRD data. If this is wrong it won't break but all dates are likely to be NA
yob_origin	value to add yob values to to get actual year of birth (Generally 1800)
regex	character regular expression to identify data files in the directory. This is separated from the filetype by an underscore. e.g. 'p[0-9]3' in CPRD GOLD
recursive	logical should files be searched for recursively under the data_dir?
	arguments to be passed to add_to_database

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#### **Details**

Note that if you chose to import all the filetype, you may end up with a very large database file. You may then chose only to import the files you want to use. You can always import the rest of the files later. This function may take a long time to process because it unzips (potentially large) files, reads into R where it converts the date formats before importing to SQLite. However, this initial data preparation step will greatly accelerate downstream processing.

import\_definitions Imports definitions to be searched from a csv file into a MedicalDefinition object

# Description

Imports definitions to be searched from a csv file into a Medical Definition object

#### Usage

```
import_definitions(input_file)
```

#### **Arguments**

```
input_file character path to the input file
```

#### **Examples**

last\_events

Selects the earliest event grouped by patient

#### **Description**

This function runs a select\_events() query and then groups by patient id and picks only the latest event for each patient

#### Usage

```
last_events(db = NULL, tab, columns = "eventdate", where = NULL,
    sql_only = FALSE, group_column = "patid", date_column = "eventdate")
```

# **Arguments** db

tab	the database table to extract from
columns	The other columns to be extracted
where	sting representation of the selection criteria
sql_only	logical should the function just return a string of the SQL query?
-	1 75 6 144 441

A database connection object

group\_column to group by. Default is patid date\_column the column to sort by. default is eventdate

20 match\_case

# Value

a dataframe or a string representing an sql query

#### **Examples**

```
## Not run:
b2 <- last_events(db, tab = "Clinical", other_columns = c("eventdate", "medcode"),
where = "medcode %in% .(a$medcode)")
## End(Not run)</pre>
```

Lists all of the EHR attribute names in .ehr

list\_EHR\_attributes

# Description

Lists all of the EHR attribute names in .ehr

# Usage

```
list_EHR_attributes()
```

match\_case

Selected controls matching a list of variables from a case

# **Description**

Helper function for get\_matches() This function wil perform incidence density sampling or exact sampling up to a supplied number of matched controls

# Usage

```
match_case(matcher, control_pool, n_controls, id, replace)
```

# **Arguments**

matcher list of character strings defining the matching conditions

control\_pool dataframe of potential controls for matching

n\_controls

The number of controls for each case. If replace == FALSE this is a maximum value

id named vector of length 1 for the variable name and value of the case identifier replace logical should sampling of matched controls be with replacement (incident density sampling) or not.

## Value

A dataframe of matched controls or NULL if no controls could be found

match\_on\_index 21

match on index	Function for performing matching of controls to cases using the con-
	sultation files to generate a dummy index date for controls.

# **Description**

Controls are matched on an arbitrary number of categorical variables and on continuous variables via the extra\_conditions argument. Also the date at index\_var is matched to the eventdate in the consultation files, providing a dummy index date for controls of a consultation within +/-index\_diff\_limit days of the index date.

## Usage

```
match_on_index(cases, control_pool, index_var, match_vars,
  extra_conditions = "", index_diff_limit = 90, consult_path,
  n_controls = 5, cores = 1, import_fn = read.delim, ...)
```

## Arguments

cases	A dataframe of cases to which to match controls	
control_pool	A dataframe of possible contols to match to cases	
index_var	character string of the name of the variable containing index dates	
match_vars	$character\ vector\ detailing\ the\ common\ variables\ in\ \verb{cases}\ and\ \verb{control\_pool}$	
	to match on	
extra_condit	ions	
	character string detailing other matching constraints (see details)	
index_diff_limit		
	integer number of days before or after the case index date that dummy index dates can be picked from the consultation files	
consult_path	path to directory containing consultation files	
n_controls	integer the number of controls to attempt to match to each case	
cores	integer the number of processor cores to be used in processing	
import_fn	function name stipulating the function used to read the consultation files	
• • •	extra arguments to be passed to import_fn	

## **Details**

Note that the consultation files must be in flat-file format (i.e. not as part of the database, but as text (or other filetype, e.g stata dta) files). Set the import\_fn argument to use different file formats (e.g. foreign::read.dta or readstata13::read.dta13)

The extra\_conditions argument can add extra condtions to the matching criteria on top of the matching vars for example you could add "year > 1990". You can wrap calls to expressions in dotted brackets to automatically expand them. This is particularly useful when you want to find the value for each individual case. Each case is denoted by CASE e.g. "start\_date < .(CASE\$start\_date)" will ensure the start date for controls is prior to the start date for the matched case.

# Value

a dataframe of matched controls

22 Medical Definition

# Description

This function accepts a data frame with a column for CPRD medcodes and merges with a medical lookup table to give columns for Read/OXMIS codes and optional descriptions

# Usage

```
medcodes_to_read(medcodes_data, lookup_table, medcodes_name = "medcode",
    lookup_readcodes = "readcode", lookup_medcodes = "medcode",
    description = TRUE)
```

#### **Arguments**

```
medcodes_data
```

a dataframe with a column matching medcodes\_name

lookup\_table a dataframe with columns matching lookup\_readcodes and lookup\_medcodes medcodes\_name

character name of the CPRD medcodes column in medcodes\_data

lookup\_readcodes

character name of the Read codes column in the lookup\_table

lookup\_medcodes

character name of the CPRD medcodes column in the lookup\_table

description logical Should description and other categories from the lookup table also be included?

#### **Details**

Note that if the names of the medcodes columns are different in the data and the lookup table, the name in the data is retained To maintain sanity, a warning will be given to inform of are any name conflicts between the input data and the lookup

#### Value

a data frame matching the input medcodes\_data with the Read codes and optional description columns merged in.

```
MedicalDefinition Constructor function for MedicalDefinition class
```

## **Description**

Constructor function for MedicalDefinition class

# Usage

```
MedicalDefinition(terms = NULL, codes = NULL, tests = NULL,
   drugs = NULL, drugcodes = NULL)
```

patients\_in\_window 23

## **Arguments**

terms	list of character vectors or NULL
codes	list of character vectors or NULL
tests	list of character vectors or NULL
drugs	list of character vectors or NULL
drugcodes	list of character vectors or NULL

#### **Details**

Elements marked with a "-" are excluded. Elements marked with a "r

## **Examples**

patients\_in\_window Select patients alive and registered between certain dates

#### **Description**

This function selects patients from the patient table of a CPRD database who are alive and registered within a supplied window

# Usage

```
patients_in_window(db, startdate, enddate, qs = TRUE,
    registration_buffer = 0, patient_tablename = "Patient")
```

# **Arguments**

db a database connection

startdate character for the start of the window. format %Y-%m-%d

enddate character for the end of the window. format %Y-%m-%d

qs logical should only patients deemed to be of acceptable quality standard be selected? Most downloaded cohorts will be up to standard by default

registration\_buffer

numeric how many days must patients be registered for prior to the startdate to be included? Setting a posive value can reduce information bias

patient\_tablename

The name of the patient table. default is Patient

#### **Details**

criteria are that crd is before start of window, tod is after end of window deathdate is after end of window

24 prev\_terms

```
patients_per_medcode
```

Produce a dataset of CPRD medcodes with frequencies of patients in the clinical table

# **Description**

This function aggregates all distinct patients matching each CPRD medcode in the clinical table

# Usage

```
patients_per_medcode(db, clinical_table = "Clinical", patid = "patid",
  medcode = "medcode")
```

# Arguments

#### **Details**

Note that this does not translate to Read/OXMIS codes. This function should be fast because all of the heavy lifting happens in SQLite before the data is exported to R

## **Examples**

```
## Not run:
medcode_counts <- patients_per_medcode(db)
head(medcode_counts)
## End(Not run)</pre>
```

prev\_terms

This function adds columns enabling one to calculate numerators and denominators for prevalence and incidence.

# Description

See the vignette for more details.

# Usage

```
prev_terms(dat, event_date = "eventdate", year_fn = standard_years)
```

prev\_totals 25

# **Arguments**

dat	dataframe of longitudinal data
event_date	character name of the column used to identify clinical events
year_fn	function that determines how year start and end dates are calculated

# Value

longitudinal data frame with incidence, prevalence and followup columns

prev_totals	Calculates the prevalence totals for the output of a data frame of events/patients etc.

# Description

e.g. Run on the output of a call to prev\_terms

# Usage

```
prev_totals(dat, included_totals = c("year", .ehr$practice_id),
   time_var = "year", person_years = 100)
```

# **Arguments**

```
dat a dataframe

included_totals

character vector describing which aggregates should be included e.g. c("year",
 "practid")

time_var name of the variable determining timepoints

person_years numeric multiplier for presentation of prevalence and incidence numbers
```

# **Details**

Outputs totals by aggregated by a time variable and other variables Updated to use dplyr to do the aggregation - Now 40x faster!

#### Value

list of aggregates and/or the original data

print.MedicalDefinition

```
print.EHR_definition
```

Tools for describing EMR\_description objects.

# Description

Tools for describing EMR\_description objects.

# Usage

```
## S3 method for class 'EHR_definition'
print(x, element = NULL, level = 3, ...)
```

# Arguments

```
x A EHR_definition object
```

element an element name

level nesting level for display purposes

... Additional arguments

```
print.MedicalDefinition
```

Basic print method for medical definition classes

# Description

Basic print method for medical definition classes

# Usage

```
## S3 method for class 'MedicalDefinition' print(x, ...)
```

# **Arguments**

```
x an object of class "medical_definition"
```

... Potential further arguments (required for method/generic reasons)

product 27

product

A sample of 500 medicines used in UK primary care

# Description

A sample of 500 medicines used in UK primary care

# Usage

product

# **Format**

A dataframe with 500 observations:

prodcode Unique internal code for the entity term

**productname** Description of the code**bnfcode** BNF code for the medicine**bnfchapter** BNF chapter heading

Source

```
http://www.bnf.org/
```

qof\_15\_months

Helper function providing startdate and enddate for a given year

# **Description**

Start and end dates matching QOF year start/ends

#### Usage

```
qof_15_months(year)
```

# **Arguments**

year

integer

28 random\_dates

qof\_years

Helper function providing startdate and enddate for a given year

# Description

Start and end dates matching QOF year start/ends

# Usage

```
qof_years(year)
```

# **Arguments**

year

integer

random\_dates

Generates random dates between a start and end day.

# Description

dates are in usual R as.Date() format

# Usage

```
random_dates(n, start_day, end_day)
```

# Arguments

Number of dates to be returnedstart\_daystring representation of a start dayend\_daystring representation of a start day

# **Details**

Enter start and end dates in ISO format, e.g. "2001-02-04"

# Value

a vector of dates

read\_to\_medcodes 29

read\_to\_medcodes

Translate Read/Oxmis codes to CPRD medcodes

## **Description**

This function accepts a data frame with a column for Read/Oxmis codes and merges with a medical lookup table to give columns for CPRD medcodes and optional descriptions

#### Usage

```
read_to_medcodes(readcodes_data, lookup_table, readcodes_name = "readcode",
   lookup_readcodes = "readcode", lookup_medcodes = "medcode", description)
```

# Arguments

readcodes\_data

a dataframe with a column matching medcodes\_name

lookup\_table a dataframe with columns matching lookup\_readcodes and lookup\_medcodes readcodes name

character name of the Read codes column in readcodes\_data

lookup\_readcodes

character name of the Read codes column in the lookup\_table

lookup medcodes

character name of the CPRD medcodes column in the lookup\_table

description logical Should description and other categories from the lookup table also be included?

#### **Details**

Note that if the names of the Read/Oxmis codes columns are different in the data and the lookup table, the name in the data is retained To maintain sanity, a warning will be given to inform of are any name conflicts between the input data and the lookup

# Value

a data frame matching the input medcodes\_data with the Read codes and optional description columns merged in.

read\_zip

Reads a zipped data file to a dataframe

# Description

This function will unzip a zipped text file and read it in to an R data frame

# Usage

```
read_zip(file, ...)
```

30 repsample\_example

# **Arguments**

file character a file to read in
... extra arguments to pass to read.delim

# **Details**

Default behaviour is to read in as a standard read.delim call. extra arguments to read.delim can be passed to the function

#### Value

a dataframe

rEHR

The rEHR package.

# **Description**

The rEHR package.

repsample\_example

An example dataset to demonstrate the repsample function. 2474 theroetical UK GP Practices.

#### **Description**

An example dataset to demonstrate the repsample function. 2474 theroetical UK GP Practices.

## Usage

```
repsample_example
```

#### **Format**

A dataframe with 2474 observations:

practicecode ID for the GP practice
postcode postcode for the practice

shacode Strategic Health Authorty Codeshaname Strategic Health Authorty name

pctcode Primary care trust codepctname Primary care trust name

listsize number of patients registered at practice

ftes ftes

soaimd07 2007 IMD deprivation score

ruralvar Rurality

# Source

```
http://www.jstatsoft.org/v55/c01/paper
```

select\_by\_year 31

select_by_year Runs a series of selects over a year range and collects in a list	list of	
--	---------	--

# Description

This function applies a database select over a range of years and outputs as a list or a dataframe The function can be parallelised using parallel.

# Usage

```
select_by_year(dbname = NULL, db = NULL, tables, columns = "*", where,
  year_range, year_fn = qof_years, as_list = FALSE,
  selector_fn = select_events, cores = 1L, ...)
```

#### **Arguments**

dbname	path to the database file
db	a database connection
tables	character vector of table names
columns	character vector of columns to be selected from the tables
where	character string representation of the selection criteria
year_range	integer vector of years to be queried
year_fn	function that determines how year start and end dates are calculated
as_list	logical: Should the results be returned as a list of years? If not, the data is collapsed into a dataframe
selector_fn	function to select from the database. See notes.
cores	integer: The number of processor cores available.
	extra arguments to be passed to the selector_fn

#### **Details**

Because the same database connection cannot be used across threads, the input is a path to a database rather than a database connection itself and a new connection is made with every fork.

columns can take a character vector of arbitrary length. This means you can use it to insert SQL clauses e.g. "DISTINCT patid".

Year start and year end criteria can be added to the where argument as 'STARTDATE' and 'END-DATE'. These will get translated to the correct start and end dates specified by year\_fn

Note that if you are working with temprary tables, you need to set cores to 1 and specify the open database connection with db This is because the use of mclapply means that new database connections need to be started for each fork and temporary files can only be seen inside the same connection

The selector\_fn argument determines how the database select operates. Default is the select\_events function. Alternatives are first\_events and last\_events

32 select\_events

#### **Examples**

```
## Not run:
# Output from a single table
where_q <- "crd < STARTDATE & (is.null(tod) | tod > ENDDATE) & accept == 1"
ayears <- select_by_year(db, "Patient", columns = c("patid", "yob", "tod"),</pre>
                          where = where_q, year_range = 2000:2003)
# Output from multiple tables
load("data/medical.RData")
a <- read.csv("data/chronic-renal-disease.csv")</pre>
a <- read_to_medcodes(a, medical, "code", lookup_readcodes= "readcode",
                      lookup_medcodes="medcode", description = T)
where_q <- "eventdate >= STARTDATE & eventdate <= ENDDATE & medcode %in% .(a$medcode)"
byears <- byears <- select_by_year("~/rOpenHealth/CPRD_test/Coupland/Coupland",</pre>
                                    c("Clinical", "Referral"),
columns = c("patid", "eventdate", "medcode"),
where = where_q, year_range = 2000:2003, as_list = FALSE,
cores = 10)
## End(Not run)
```

select\_events

Extracts From the database

#### **Description**

This is a generic function for extracting EHR data from the database

## Usage

```
select_events(db = NULL, tab, columns = "*", where = NULL,
sql_only = FALSE, convert_dates = FALSE)
```

## Arguments

db a database connection

tab the database table to extract from

columns character vector of columns to extract from the table "\*" means all tables

where sting representation of the selection criteria

sql\_only logical should the function just return a string of the SQL query?

convert\_dates

logical should date fields be converted to R date format?

# **Details**

The function is the base function for a range of others It can either extract by itself or generate the SQL to make a query. In this way it can be combined to make compound queries. The where argument is equivalent to the WHERE clause in sql The elements are converted to SQL using dplyr::translate\_sql\_q If an element is wrapped in a '.()', the element is expanded. Dates should be entered as strings in ISO format (

set\_CPRD 33

#### Value

a dataframe or a string representing an sql query

# **Examples**

```
## Not run:
# medical lookup tables are provided with CPRD
load("data/medical.RData")
a <- read.csv("data/chronic-renal-disease.csv")
a <- read_to_medcodes(a, medical, "code", lookup_readcodes= "readcode",
lookup_medcodes="medcode", description = T)
b <- select_events(db, tab = "Referral", columns = c("patid", "eventdate", "medcode"),
where = "medcode %in% .(a$medcode) & eventdate < '2000-01-01'")
b1 <- select_events(db, tab = "Clinical", columns = c("patid", "eventdate", "medcode"),
where = "medcode %in% .(a$medcode) & eventdate < '2000-01-01'")
## End(Not run)</pre>
```

set\_CPRD

Sets EHR metadata to CPRD format When this is run, most functions in rEHR act as though the EHR database is CPRD

# Description

Sets EHR metadata to CPRD format When this is run, most functions in rEHR act as though the EHR database is CPRD

# Usage

```
set_CPRD()
```

# Details

Metadata on EHR type is stored in the .ehr environment. This allows the same functions to work across different data sources. The .ehr environment is not desgined to be accessible to the user, but accessor functions are provided. CPRD is the default EHR setting.

# See Also

```
get_EHR_value set_EHR_value
```

set\_EHR\_attribute Sets the value of an attribute in the .ehr environment

#### **Description**

Sets the value of an attribute in the .ehr environment

# Usage

```
set_EHR_attribute(x, value)
```

### **Arguments**

x an ehr attribute namevalue the value to set to the attribute

# **Examples**

```
set_CPRD()
set_EHR_attribute(practice_id, "pracid")
```

```
simulate_ehr_consultations
```

Generates simulated GP consultation tables.

# **Description**

This function generates simulated GP consultations based on an EHR\_definition object and a patient table, as generated by simulate\_ehr\_patients. Multicore functionality is implemented via mclapply

# Usage

```
simulate_ehr_consultations(ehr_def, patient_table, cores = 1)
```

# Arguments

#### Value

data frame of simulated GP consultations

# **Examples**

```
## Not run: patient <- simulate_ehr_patients(ehr_definition) cons <-
    simulate_ehr_consultations(ehr_def, patient_table = patient, cores = 4)
## End(Not run)</pre>
```

simulate\_ehr\_events 35

```
simulate_ehr_events
```

Generate simulated events tables

#### **Description**

This function can generate events for clinical, referral and therapy tables. These are based on the consultation tables generated by simulate\_ehr\_consultations.

# Usage

```
simulate_ehr_events(ehr_def, consultation, event_type = c("clinical",
    "referral", "therapy"), cores = 1, therapy_lookup = NULL)
```

## **Arguments**

#### **Details**

This function is relatively basic - for clinical and referral tables, it generates events according to the comorbidities defined in the <code>ehr\_def</code>, with the mean number of events for each consultation being defined in the <code>ehr\_def</code> for that event\_type. For therapy events, the function simply samples the therapy\_lookup table, with the mean number of events for each consultation being defined in the <code>ehr\_def</code> for therapy. Therefore, at the moment, the therapies bear no relationship to the conditions the patient has and are only for the purposes of explaining the functioning of the package. The random sampling is based on a poisson distribution

#### Value

dataframe

```
simulate_ehr_patients
```

Generate a dataframe of simulated patients with exit dates based on presented comorbidities.

# **Description**

The definitions of the patient file are all in the ehr\_def object

# Usage

```
simulate_ehr_patients(ehr_def)
```

36 standard\_years

#### **Arguments**

```
ehr_def an object of class EHR_definition
```

#### **Details**

Patients must have transferred out after the ealiest possible collection date ehr\_def\$practice\$uts\_limit

#### Value

a dataframe of simulated patients

```
simulate_ehr_practices
```

generates a simulated dataframe of primary care practices in the same format as is used in CPRD

# **Description**

generates a simulated dataframe of primary care practices in the same format as is used in CPRD

# Usage

```
simulate_ehr_practices(ehr_def)
```

# **Arguments**

ehr\_def an object of class ehr\_def

#### **Details**

The definitions of the practice file are all in the ehr\_def object

# Value

a dataframe of simulated practices

standard\_years

Helper function providing startdate and enddate for a given year

# Description

Standard years

# Usage

```
standard_years(year)
```

# **Arguments**

year integer

surv\_sims 37

surv\_sims

Function to simulate survival data.

# **Description**

Model: proportional hazards, h(t; cov\_mat, beta) = exp(cov\_mat indicators for Type I censoring (common censoring time 'tc').

#### Usage

```
surv_sims(cov_mat, beta, cens_type = c("typeI", "noninformative"),
  baseline_hazard, cens_hazard = 0.04, cens_prob = 0, scale = 1,
  weibull_shape = 1)
```

# **Arguments**

cov\_mat n x p matrix of cov\_matiates

beta p-vector of regression coefficients

cens\_type typeI censoring or non-informative based on exponential distribution

baseline\_hazard
for modelling death dates

cens\_hazard log(hazard) for non-informative censoring

cens\_prob expected censoring fraction (0 <= cens\_prob < 1). Used for typeI censoring

scale value to scale up the time variable by

weibull\_shape
shape parameter for the weibull distribution. 1 is the same as an exponential

# Details

Weibull\_shape is the k (shape) parameter from a weibull distribution

- A value of k < 1 indicates that the mortality rate decreases over time. This happens if there is significant infant mortality
- A value of k = 1 indicates that the mortality rate is constant over time. This might suggest random external events are causing mortality. This is the same as an exponential distribution
- A value of k > 1 indicates that the mortality rate increases with time. This happens if there is an aging process.

#### Value

Censored exponential survival times and censoring

38 temp\_table

temp\_location

Sets location of the db temporary store for temporary tables

#### **Description**

By default, sqlite stores temp tables in /tmp (Or windows equivalent). If you are building large temporary tables and don't have a large /tmp directory, you can get "database or disk is full" errors. If you have a lot of RAM you can set store to "RAM" and the temp files will be stored in RAM rather than in /tmp. This could also speed things up.

# Usage

```
temp_location(db, store = c("tmp", "RAM"))
```

# **Arguments**

db a database connection store character vector either "tmp" or "RAM"

temp\_table

Creates a temporary table in the database

# **Description**

This function is useful if most of your work is on a subset of the database

## Usage

```
temp_table(db, tab_name, select_query)
```

# Arguments

```
db a database connection object
tab_name character name for the teporary table
select_query character the query that specifies the temporary table
```

## **Details**

The table will exist for as long as the database connection is kept open The Select\_query argument will take the output from a select\_events(sql\_only = TRUE) based function

# **Examples**

to\_stata 39

to_stata	Compresses a dataframe and saves in stata format. Options to save as Stata 12 or 13.

# Description

Automatically compresses data to reduce file size

# Usage

```
to_stata(dat, fname, stata13 = FALSE, ...)
```

# Arguments

dat	dataframe
fname	character string: filepath to save to
stata13	logical Save as Stata13 compatible format?
	arguments to be passed to compress

# **Details**

Defaults to saving compressed dates to integer days from 1960-01-01 which is the standard in stata.

to_temp_table	Send a dataframe to a temporary table in the database
to_temp_table	sena a adiajrame to a temporary table in the adiabase

# Description

The table is a temporary database and is linked only to the current connection object

# Usage

```
to_temp_table(db, tab_name, dat, overwrite = FALSE)
```

# Arguments

db	a database connection
tab_name	character name for the new temporary database table
dat	dataframe to send to the temporary database table
overwrite	logical if a table already exists with the same name should it be dropped?

40 wrap\_sql\_query

wrap\_sql\_query

combines strings and vectors in a sensible way for select queries

# **Description**

This function is a variant of the sprintf function. In the query, can be placed identifier tags which are a hash character followed by a number e.g. #1 The number in the tag reflects the position of the arguments after the query The resut of evaluating that argument will then be inserted in place of the tag. If the result of evaluating the argument is a vector of length 1, it is inserted as is. If it is a vector of length > 1, it is wrapped in parentheses and comma separated.

# Usage

```
wrap_sql_query(query, ...)
```

# Arguments

```
query a character string with identifier tags (#[number]) for selecting the argument in ... optional arguments selected by the identifier tags
```

#### **Details**

Note that this function is for help in constructing raw SQL queries and should not be used as an input to the where argument in  $select\_event$  calls. This is because these calls use translate\_sql\_q to translate from R code to SQL

# **Examples**

```
medcodes1 <- 1:5
practice <- 255
wrap_sql_query("eventdate >= STARTDATE & eventdate <= ENDDATE & medcode %in% #1 &
    practice == #2", medcodes1, practice)</pre>
```

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