CPRDTools: a CPRD GOLD data wrangling toolkit

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Abstract

The analysis of large scale data, and moreover the analysis of large scale electronic health records data has become more commonplace. The ease and ability of modern data generation and capture means there is the potential across almost all industries to capture more data now than ever before, and that equates to large data resources, with the Clinical Practice Research Datalink (CPRD) no exception. These large data resources, as attractive and appealing as they may be to researchers, pose a significant problem how to manage all that data? CPRDTools is a collection of wrapper R functions intended to simplify the loading, extraction and management of CPRD GOLD specific and associated electronic health records data. Allowing for the loading of CPRD and non-CPRD data into a SQLite database, providing an efficient, secure and updatable repository for the data, keeing the origional source files intact. Through data queries, user-defined data are drawn allowing for subsetting, joining and filtering in a single step, creating alaysis ready data.

Introduction

The CPRD is one of the largest longitudinal medical records databases in the world, supported by the National Institute of Health (NIHR) and the Medicines and Healthcare products Regulatory Agency (MHRA). It was first established in 1987 as the Value Added Medical Products (VAMP) dataset, this grew into the General Practice Research Database (GPRD) in 1993, before its final transition into CPRD in 2012 (Herrett et al. 2015).

CPRD GOLD data are comprised of ten separate datasets: patient, practice, staff, consultation, clinical, additional clinical details, referral, immunisation, test and therapy (Padmanabhan 2017). These datasets contain their specific data and are linkable through a unique linkage key field, where key does not imply importance but a unique variable contained in two datasets allowing them to be joined, such as the CPRD-assigned and anonymised unique patient identifier patid.

Due to the size of CPRD, data extracted for research are spread over multiple text (.txt) files within each dataset to enable file transfer. This means that for CPRD clinical data, for instance, a researcher may receive their requested clinical data broken up over numerous individual text files. This is done to aid with file completeness and reduce turn-around times if errors are found. If an error occurred during the transfer of data between the data owner and the researcher or in the extraction of the requested data by the data owner, the error can potentially be limited to only select files, requiring only their replacement with the corrected/error-free versions.

Often these text files are additionally zipped, or compressed, requiring that these files first be uncompressed or unzipped. These multiple files from each dataset (clinical, referral etc.) then need to be grouped together and amalgamated into a single table. Tables are a collection of data of the same shape, from the same dataset. In CPRD, each separate dataset (patient, practice, clinical etc.) forms a table. These tables are then stored in the SQLite database.

SQLite is an open source, SQL based database engine (SQLite 2017). The use of a SQLite database provides an efficient storage solution, allowing for the loading, updating and maintenance of the database, all while retaining the original raw data files unaltered. A SQLite database permits for rapid data extraction through the use of data queries, drawing the required data from the database, allowing filtering, sub-setting, limiting and the joining of data in a single execution step. This document aims to provide a simple and introductory overview of CPRDTools and its application to arbitrary (fictional) data. This data though are provided in the manner in which many CPRD data extract are received, where data are spread over multiple files and often located in sub-folders.

CPRDTools is loaded using:

```
library(devtools)
install_github("JamesCFSchmidt/CPRDTools")
library(CPRDTools)
```

CPRDTools overview

The functions within 'CPRDTools' broadly fall into three groups, categorised by their general application area: (1) - loading, (2) - maintenance and other tasks and (3) - extraction. **Loading** encompasses all functions used in the reading, converting and writing of data into the database, including a function used to list all available files and all available CPRD files in a specified location. Functions used in database maintenance, query speed improvements and date conversion fall under **maintenance and other tasks**, and finally, functions used to, and in the process of, drawing and retrieving data from the database fall within **extraction**.

The data

The example data supplied with this package resembles CPRD GOLD data in layout and in file naming convention. Additionally included are example secondary files from 'Hospital Episode Statistics' (HES) patient information and 'Office for National Statistics' (ONS) actuarial life tables. The location of the file and the location where the database will be build must be provided

```
#database location
DB.path = "/home/j/jcfs2/Documents/Test"
#location of example data
FILE.path = dirname(system.file("extdata", "hes patient 20336R.txt", package = "CPRDTools"))
```

Loading

Before loading any data into the database, it is best practice to understand the layout of the *raw* data. This can be achieved by either navigating to the location where the data are stored or by employing the list_files and list_cprd functions.

Listing available data

The list_files function provides a list of all files of a specified type in a specified location. To view all text (.txt) files in the data location

To view all compressed (.zip) files in the data location

```
list_files(file_location = FILE.path,
           file_type = ".zip")
## $file_location
## [1] "/lustre/ahome3/j/jcfs2/R/x86_64-pc-linux-gnu-library/4.2/CPRDTools/extdata"
##
## $files
##
                                   files
## 1 /1234.aa1 Extract Practice 001.zip
## 2 /1234.aa2_Extract_Clinical_001.zip
## 3 /1234.aa2_Extract_Clinical_002.zip
## 4 /1234.aa2_Extract_Clinical_003.zip
## 5 /1234.aa2_Extract_Clinical_004.zip
## 6 /1234.aa2_Extract_Practice_001.zip
And to view all files in the data location, excluding files in sub-folders
list_files(file_location = FILE.path,
           file_type = "all")
## $file_location
## [1] "/lustre/ahome3/j/jcfs2/R/x86_64-pc-linux-gnu-library/4.2/CPRDTools/extdata"
##
## $files
##
                                   files
## 1 /1234.aa1_Extract_Practice_001.zip
## 2 /1234.aa2_Extract_Clinical_001.zip
## 3 /1234.aa2_Extract_Clinical_002.zip
## 4 /1234.aa2 Extract Clinical 003.zip
## 5 /1234.aa2_Extract_Clinical_004.zip
## 6 /1234.aa2_Extract_Practice_001.zip
## 7
                /hes_patient_20336R.txt
## 8
                             /ons_lt.csv
## 9
                                /patient
```

From the above it can be seen that there are six zipped text files (.zip), one excel file (.xlsx), one standard text file (.txt) and a folder, patient. In order to view the CPRD GOLD specific files, corresponding to named CPRD GOLD datasets (patient, practice, staff, consultation, clinical, additional clinical details, referral, immunisation, test and therapy), the list_cprd function is used. This function provides the core functionality to the loading of CPRD GOLD data, generating a list of all the CPRD GOLD specific files in the specified data location.

```
## 1 /1234.aa1_Extract_Practice_001.zip Practice
## 6 /1234.aa2_Extract_Practice_001.zip Practice
##
## $tables
## table file_count
## 1 Clinical 4
## 2 Practice 2
```

Here the folders input is defined as , showing only the .zip files data contained in the data location. The .txt file is ignored as this is not a CPRD GOLD dataset. When the folder input is changed to , the available data found in the patient folder is displayed.

```
## $file_location
## [1] "/lustre/ahome3/j/jcfs2/R/x86_64-pc-linux-gnu-library/4.2/CPRDTools/extdata"
##
## $all_files_tables
## files table
## 1 /patient/aa1_Extract_Patient_001.zip Patient
## 2 /patient/aa2_Extract_Patient_001.zip Patient
## ## $tables
## table file_count
## 1 Patient 2
```

In the **\$table** output, the **list_cprd** function defines the CPRD GOLD tables and the count of *raw* files associated with that CPRD GOLD table in the data location. This outputted list is core to the automated loading of CPRD GOLD data, using the names CPRD GOLD tables and files to load all or a selection of specified CPRD GOLD tables

Loading all available data

The load_cprd function (relying internally on the list_cprd function for the list of files and table in the data location) is able to automatically load in all or a selection available CPRD GOLD files in the specified data location. Importing, uncompressing (unzipping) and appending all files from a CPRD GOLD-specific table into a database.

To load the patient data into a new database, the load_cprd function is defined as

```
## Rows: 12 Columns: 20
## -- Column specification ------
## Delimiter: "\t"
## chr (4): frd, crd, tod, deathdate
## dbl (14): patid, vmid, gender, yob, marital, famnum, CHSreg, prescr, capsup,...
## lgl (2): mob, CHSdate
```

```
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
## -----LOAD OF TABLE Patient, FILE No. 1 SUCCESSFUL-----
##
## Rows: 12 Columns: 20-- Column specification ------
## Delimiter: "\t"
## chr (4): frd, crd, tod, deathdate
## dbl (14): patid, vmid, gender, yob, marital, famnum, CHSreg, prescr, capsup,...
## lgl (2): mob, CHSdate
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
## -----LOAD OF TABLE Patient, FILE No. 2 SUCCESSFUL-----
##
## $database_location
## [1] "/home/j/jcfs2/Documents/Test"
##
## $files_in_location
##
                                   files
                                           table
## 1 /patient/aa1_Extract_Patient_001.zip Patient
## 2 /patient/aa2 Extract Patient 001.zip Patient
##
## $tables in location
      table file_count
##
## 1 Patient
##
## $loaded tables
## [1] "Patient"
## $load_report
      table load_total file_total size_Mb size_Gb
                                    0.001
## 1 Patient
##
## $database_tables
## [1] "Patient"
                     "lookup_table"
##
## $load_start
## [1] "2022-10-04 15:34:07 BST"
##
## $load end
## [1] "2022-10-04 15:34:08 BST"
##
## $load_time
## Time difference of 1.283845 secs
```

The patient data are found in a sub-folder within the data location () and the data are compressed (). As this is a new load of the database, no overwriting is required () and no CPRD specific mapping is needed ().

Alternatively to specifying each individual table to load (additional, clinical, consultation, immunisation, patient, practice, referral, staff, test, therapy), the specification of the loading of all tables can be performed.

```
tables_to_load = "all",
folder = F,
zip = T,
load_mapping = F,
overwrite = F)
```

This returns an error ## Error in load_cprd(db_path = DB.path, file_location = FILE.path, tables_to_load = "all", : tables_to The error is due to the patient table already being contained in the database. The should only be applied on new databases or in conjunction with . The remainder of the CPRD GOLD data are loaded with load_cprd(db_path = DB.path, file_location = FILE.path, tables_to_load = c("Clinical", "Practice"), folder = F, zip = T, load_mapping = F, overwrite = F) ## Rows: 12 Columns: 10 ## -- Column specification -----## Delimiter: "\t" ## chr (2): eventdate, sysdate ## dbl (8): patid, constype, consid, medcode, staffid, episode, enttype, adid ## i Use 'spec()' to retrieve the full column specification for this data. ## i Specify the column types or set 'show_col_types = FALSE' to quiet this message. ## -----LOAD OF TABLE Clinical, FILE No. 1 SUCCESSFUL-----## ## Rows: 12 Columns: 10-- Column specification -----## Delimiter: "\t" ## chr (2): eventdate, sysdate ## dbl (8): patid, constype, consid, medcode, staffid, episode, enttype, adid ## i Use 'spec()' to retrieve the full column specification for this data. ## i Specify the column types or set 'show_col_types = FALSE' to quiet this message. ## -----LOAD OF TABLE Clinical, FILE No. 2 SUCCESSFUL------## Rows: 12 Columns: 10-- Column specification ------## Delimiter: "\t" ## chr (2): eventdate, sysdate ## dbl (8): patid, constype, consid, medcode, staffid, episode, enttype, adid ## i Use 'spec()' to retrieve the full column specification for this data. ## i Specify the column types or set 'show_col_types = FALSE' to quiet this message. ## -----LOAD OF TABLE Clinical, FILE No. 3 SUCCESSFUL-----## Rows: 12 Columns: 10-- Column specification ------## Delimiter: "\t" ## chr (2): eventdate, sysdate ## dbl (8): patid, constype, consid, medcode, staffid, episode, enttype, adid

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i Specify the column types or set 'show_col_types = FALSE' to quiet this message.

i Use 'spec()' to retrieve the full column specification for this data.

```
## -----LOAD OF TABLE Clinical, FILE No. 4 SUCCESSFUL-----
##
## Rows: 5 Columns: 4-- Column specification ------
## Delimiter: "\t"
## chr (2): lcd, uts
## dbl (2): pracid, region
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
## -----LOAD OF TABLE Practice, FILE No. 1 SUCCESSFUL-----
## Rows: 5 Columns: 4-- Column specification -----
## Delimiter: "\t"
## chr (2): lcd, uts
## dbl (2): pracid, region
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
## -----LOAD OF TABLE Practice, FILE No. 2 SUCCESSFUL------
## $database_location
## [1] "/home/j/jcfs2/Documents/Test"
## $files_in_location
                                files
## 1 /1234.aa1_Extract_Practice_001.zip Practice
## 2 /1234.aa2_Extract_Clinical_001.zip Clinical
## 3 /1234.aa2_Extract_Clinical_002.zip Clinical
## 4 /1234.aa2_Extract_Clinical_003.zip Clinical
## 5 /1234.aa2_Extract_Clinical_004.zip Clinical
## 6 /1234.aa2_Extract_Practice_001.zip Practice
##
## $tables_in_location
      table file_count
## 1 Clinical
## 2 Practice
##
## $loaded_tables
## [1] "Clinical" "Practice"
## $load_report
       table load_total file_total size_Mb size_Gb
## 1 Clinical
                          4 2e-03
                     4
## 2 Practice
                                   5e-04
##
## $database_tables
## [1] "Clinical"
                    "Patient"
                                   "Practice"
                                                 "lookup_table"
##
## $load_start
## [1] "2022-10-04 15:34:08 BST"
## $load_end
## [1] "2022-10-04 15:34:09 BST"
```

```
##
## $load_time
## Time difference of 0.5281887 secs
```

The \$loaded_tables output shows the current tables loaded and available in the database '

Loading a single CPRD table

The patient table previously could have been loaded singularly using the load_table function. Similar to the load_cprd function, this will load CPRD-specific data tables, using the compiled list of available files and their respective tables generated in the list_cprd function. This function imports, unzips and appends the files specific to a CPRD GOLD table but performs the task on a single table at a time.

This may be of use when the data are stored in separate file locations, not within sub-folders in the same data location. The loading of the practice information is performed using

```
load_table(db_path = DB.path,
         file location = FILE.path,
         table name = "Practice",
         zip = T,
         overwrite = T)
## -----Overwrite = TRUE, DELETION OF Practice COMPLETE-----
##
## Rows: 5 Columns: 4
## Delimiter: "\t"
## chr (2): lcd, uts
## dbl (2): pracid, region
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
## -----LOAD OF TABLE Practice, FILE No. 1 SUCCESSFUL-----
## Rows: 5 Columns: 4-- Column specification ------
## Delimiter: "\t"
## chr (2): lcd, uts
## dbl (2): pracid, region
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
## -----LOAD OF TABLE Practice, FILE No. 2 SUCCESSFUL------
##
## $database_location
## [1] "/home/j/jcfs2/Documents/Test"
## $files_in_location
##
                              files
                                      table
## 1 /1234.aa2_Extract_Clinical_001.zip Clinical
## 2 /1234.aa2_Extract_Clinical_002.zip Clinical
## 3 /1234.aa2_Extract_Clinical_003.zip Clinical
## 4 /1234.aa2 Extract Clinical 004.zip Clinical
```

5 /1234.aa1_Extract_Practice_001.zip Practice

```
## 6 /1234.aa2_Extract_Practice_001.zip Practice
##
## $tables in location
##
        table file_count
## 1 Clinical
## 2 Practice
                        2
##
## $loaded_table
## [1] "Practice"
##
## $load_report
##
        table load_total file_total size_Mb size_Gb
## 1 Clinical
                       2
                                   2
                                       5e-04
## 2 Practice
                       2
                                   2
                                       5e-04
                                                    0
##
## $database_tables
## [1] "Clinical"
                       "Patient"
                                      "Practice"
                                                      "lookup_table"
##
## $load_start
## [1] "2022-10-04 15:34:09 BST"
##
## $load end
## [1] "2022-10-04 15:34:09 BST"
##
## $load_time
## Time difference of 0.1625137 secs
```

Here as the practice table was previously loaded.

Updating a CPRD GOLD table

If you were to receive a new month or year of patient data, this would be append to the data already contained in the database using

Note that capitalisation of the table name for CPRD-specific tables in unimportant. In general, SQL and therefore SQLite is not case sensitive.

Updating an additional file

For peripheral or additional data, whether CPRD or non-CPRD specific, the use of the load_additional function is employed. This function loads a singular file into a table (new or existing).

```
## Rows: 24 Columns: 5
## -- Column specification -----
## Delimiter: "\t"
## dbl (5): patid, pracid, gen_hesid, n_patid_hes, match_rank
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
## -----LOAD OF TABLE hes patient SUCCESSFUL-----
##
## $database_location
## [1] "/home/j/jcfs2/Documents/Test"
##
## $loaded table
## [1] "hes_patient"
##
## $load_report
          table size_Mb size_Gb
## 1 hes_patient
                  6e-04
## $database_tables
                                                    "hes_patient"
## [1] "Clinical"
                      "Patient"
                                     "Practice"
                                                                   "lookup_table"
##
## $load_start
## [1] "2022-10-04 15:34:09 BST"
## $load_end
## [1] "2022-10-04 15:34:09 BST"
##
## $load_time
## Time difference of 0.07197189 secs
```

table_name = "hes_patient",

overwrite = F)

The accepted files types are .txt, .csv, .dta, .rds, .excel, .xl, .xls and .xlsx with no additional input for file importing.

Updating an additional file

If sub-setting, manipulation or sheet selection, for instance, is required prior to the loading of a file into a table (new or existing) is required, it is first recommended to import the file in the usual manner into R. Once the data are in the format and layout that is required, the <code>load_global</code> function can be employed

```
## -----LOAD OF TABLE Life_table SUCCESSFUL-----
##
## $database_location
## [1] "/home/j/jcfs2/Documents/Test"
##
## $loaded table
## [1] "Life_table"
##
## $load_report
          table size_Mb size_Gb
## 1 Life_table 0.1722
##
## $database_tables
## [1] "Clinical"
                      "Life_table"
                                     "Patient"
                                                    "Practice"
                                                                    "hes_patient"
## [6] "lookup_table"
##
## $load start
## [1] "2022-10-04 15:34:09 BST"
## $load_end
## [1] "2022-10-04 15:34:09 BST"
##
## $load time
## Time difference of 0.02614522 secs
```

Extracting

With the database successfully loaded with our CPRD GOLD and associated data, we are now able to perform data extracts from the database. Before executing data extracts, it is *best practice* to first interrogate the databases for loaded table (ignoring that this is provided after each load iteration previously) and the for the structure and layout of a selection of tables.

List tables, fields and first rows

table_name = "PaTient")

In order to list all loaded tables in the database, similar to the output in the \$loaded_tables, returned when loading data via the data-loading function previously, specify

```
db_tables(db_path = DB.path)

## $database_location
## [1] "/home/j/jcfs2/Documents/Test"
##

## $tables
## [1] "Clinical" "Life_table" "Patient" "Practice" "hes_patient"
## [6] "lookup_table"

loaded are the Clinical, Patient, Practice and hes_patient tables.

To view the fields within the Patient table

table_fields(db_path = DB.path,
```

```
## $database_location
## [1] "/home/j/jcfs2/Documents/Test"
##
## $fields
##
   [1] "patid"
                     "vmid"
                                  "gender"
                                              "yob"
                                                           "mob"
                                                                        "marital"
                     "CHSreg"
                                  "CHSdate"
                                              "prescr"
                                                                        "frd"
##
  [7] "famnum"
                                                           "capsup"
## [13] "crd"
                     "regstat"
                                  "reggap"
                                              "internal"
                                                           "tod"
                                                                        "toreason"
## [19] "deathdate" "accept"
                                  "pracid"
To view the first 5 row of the hes_patient table
first_n(db_path = DB.path,
        table_name = "hes_patient",
        n = 5)
## $database_location
## [1] "/home/j/jcfs2/Documents/Test"
##
## $query
## [1] "SELECT * FROM hes_patient LIMIT 5;"
##
## $first_n_rows
##
       patid pracid gen_hesid n_patid_hes match_rank
## 1 1000001
                       1236275
                  1
                                          1
                                                      1
## 2 1001001
                       1235450
                                          1
                                                      1
                   1
## 3 1002003
                  3
                      1235459
                                          1
                                                      1
## 4 1003004
                   4
                      1236872
                                          1
                                                      1
## 5 1004005
                   5
                       1235080
                                                      1
```

Extracting data

Data extraction can be performed via two functions (1) the user-input defined query_builder function and (2) query_direct, the more customisable and preferred method of data extraction.

The basic structure of the query_bilder function is

This function allows for only a simple join between two tables, the main table (from_table) and secondary table (join_table). Fields can be extracted from both tables, including the field used to join the tables together (join_on), known as a key field. Additional filtering, sub-setting and restrictions of data can be done using where, order and limit inputs.

A simple extract is defined as

```
from_table = "Patient",
              join_table = "hes_patient",
              join_fields = c("gen_hesid", "match_rank"),
              join_on = "patid",
              limit to = 5)
## $database location
## [1] "/home/j/jcfs2/Documents/Test"
##
## $database_tables
## [1] "Clinical"
                      "Life_table"
                                      "Patient"
                                                     "Practice"
                                                                     "hes_patient"
## [6] "lookup_table"
##
## $query
## [1] "SELECT DISTINCT Patient.patid, Patient.gender, Patient.yob, Patient.deathdate, hes_patient.ge
##
## $query_data
       patid gender yob deathdate gen_hesid match_rank
##
## 1 1000001
                  2 1999 2020-01-08
                                      1236275
## 2 1001001
                  2 1989
                               <NA>
                                       1235450
                                                        1
## 3 1002003
                  1 1993
                               <NA>
                                       1235459
                                                        1
```

1

1

field_list = c("patid", "gender", "yob", "deathdate"),

query_builder(db_path = DB.path,

4 1003004

5 1004005

\$query_time

##

unique_obs = T,

2 1984

1 1982

Time difference of 0.005812407 secs

Here, a unique extract of patid, gender, yob and deathdate from the Patient table and gen_hesid and match_rank from the hes_patient table are extracted, joined together via the patid field and limited to the first 10 rows of outputted data. This data can be returned directly into a dataset using

1236872

1235080

<NA>

<NA>

Building on the previous extract, now filtered for only <code>gender==1</code> and ordered by <code>patid</code> in ascending order is defined as

```
order_field = "PATID",
              order_type = "aSC",
              limit to = 5)
## $database location
## [1] "/home/j/jcfs2/Documents/Test"
##
## $database_tables
## [1] "Clinical"
                      "Life table"
                                      "Patient"
                                                      "Practice"
                                                                     "hes_patient"
## [6] "lookup table"
##
## $query
## [1] "SELECT DISTINCT Patient.paTid, Patient.gender, Patient.yob, Patient.deathdate, hes_patient.ge
## $query_data
       patid gender yob deathdate gen_hesid match_rank
## 1 1002003
                  1 1993
                               <NA>
                                      1235459
## 2 1004005
                  1 1982
                               <NA>
                                      1235080
                                                        1
## 3 1005006
                  1 1989
                               <NA>
                                      1235878
                                                        1
## 4 1007001
                  1 1987
                               <NA>
                                      1236339
                                                        1
## 5 1009010
                  1 1986
                               <NA>
                                      1235558
                                                        1
```

Note that SQLite is not case sensitive.

Time difference of 0.006338835 secs

##

\$query_time

where_filter = "geNder==1",

The preferred method for data extraction is through the direct execution of SQL code defined by the user. This allows for the most customisable, comprehensive and completed data extraction, providing all the functionality found in a SQLite SELECT statement.

A introductory overview of the SELECT statement can be found at https://www.sqlitetutorial.net/sqlite-select/2, with the basic syntax defined as

```
SELECT DISTINCT column_list
FROM table_list
    JOIN table ON join_condition
WHERE row_filter
ORDER BY column
LIMIT count OFFSET offset
GROUP BY column
HAVING group_filter;
```

To replicate the query executed using the query_builder function previously, the following is provided

```
## $database location
## [1] "/home/j/jcfs2/Documents/Test"
##
## $database_tables
## [1] "Clinical"
                       "Life table"
                                       "Patient"
                                                      "Practice"
                                                                      "hes_patient"
## [6] "lookup table"
## $query
## [1] "SELECT DISTINCT Patient.patid, \n
                                                                                   Patient.gender, \n
##
## $query_data
##
       patid gender yob deathdate gen_hesid match_rank
## 1 1000001
                  2 1999 2020-01-08
                                        1236275
## 2 1001001
                                        1235450
                  2 1989
                                < NA >
                                                         1
## 3 1002003
                   1 1993
                                <NA>
                                        1235459
                                                         1
## 4 1003004
                   2 1984
                                <NA>
                                        1236872
                                                         1
## 5 1004005
                   1 1982
                                <NA>
                                        1235080
                                                         1
##
## $query_time
## Time difference of 0.001212597 secs
```

The query here is identical to the query returned in the \$query output in the query_builder function. The table from which a field should be extracted is provided before the field such as patient.yob, extracting yob from Patient.

This query can be made more aesthetically pleasing by using an alias for the Pateint and hes_patient tables

```
##
       patid sex bith_date
                                  death
                                          hesid match_rank
## 1 1000001
                       1999 2020-01-08 1236275
               2
## 2 1001001
               2
                       1989
                                   <NA> 1235450
                                                          1
## 3 1002003
               1
                       1993
                                   <NA> 1235459
                                                          1
## 4 1003004
               2
                       1984
                                   <NA> 1236872
                                                          1
## 5 1004005
                       1982
                                   <NA> 1235080
                                                          1
```

To view the procedure in which a query will be executed, use EXPLAIN QUERY PLAN. This will provide an overview of the strategy or plan SQLite will use to execute a SQL query, providing

```
FROM Patient pat

LEFT JOIN hes_patient hes ON hes.patid = pat.patid

LIMIT 5 ;")$query_data
```

```
id parent notused
## 1 5
             0
## 2 22
             0
                      0
## 3 53
             0
                      0
##
                                                               detail
                                                            SCAN pat
## 1
## 2 SEARCH hes USING AUTOMATIC COVERING INDEX (patid=?) LEFT-JOIN
                                       USE TEMP B-TREE FOR DISTINCT
## 3
```

Shows that first a full-table scan is used on the Patient table, followed by a subset search of the hes_patient table followed by a temporary b-tree structure used to defined the unique (DISTINCT) output data. A more complete overview of the EXPLAIN QUERY PLAN command can be found at https://www.sqlite.org/eqp.html.

Maintenace and other tasks

The removal and remaining of tables and the deletion of the databases, as well as query speed improvement function make up the final section of the CPRDTools package.

Speeding up a query

A method to improve the execution times of a query is indexing. Indexing a table or dataset creates a secondary data table containing the indexing variable and the row number (rowid) associated with that number. The index is initially search, returning the row numbers for the searched fields, before using these row numbers to return the required data from.

Consider the following query, returning data from the Patient and Clinical tables, where the {sql} medcodes IN (1004,1046,1076,1146,1157,1314,1425,1454,1508,1842), defined as

```
## $database_location
## [1] "/home/j/jcfs2/Documents/Test"
##
## $database_tables
## [1] "Clinical" "Life_table" "Patient" "Practice" "hes_patient"
## [6] "lookup_table"
##
## $query
```

```
## [1] "EXPLAIN QUERY PLAN\n
                                          SELECT pat.patid,\n
                                                                                   pat.yob,\n
##
## $query_data
     id parent notused
                                                                      detail
##
## 1 3
             0
                                                                   SCAN clin
## 2 58
             Λ
                     O SEARCH pat USING AUTOMATIC COVERING INDEX (patid=?)
##
## $query_time
## Time difference of 0.0007603168 secs
Executing the EXPLAIN QUERY PLAN command provides
     id parent notused
                                                                       detail
## 1
     3
             0
                                                                   SCAN clin
## 2 58
                     O SEARCH pat USING AUTOMATIC COVERING INDEX (patid=?)
             0
```

Using a full-table scan of the clinical table for the desired medcodes before using an automatically generated, temporary index on patid.

An index can be defined on the clinical table using the medcodes as

2 60

```
add_index(db_path = DB.path,
          index_name = "clinical_index",
          index table = "clinical",
          index_field = "medcode")
## $database_location
## [1] "/home/j/jcfs2/Documents/Test"
##
## $index
## [1] "clinical index"
##
## $index_statement
## [1] "CREATE INDEX clinical_index ON clinical (medcode);"
## $query time
## Time difference of 0.002881765 secs
Now the query plan shows
     id parent notused
                                                                      detail
## 1 4
                       SEARCH clin USING INDEX clinical_index (medcode=?)
             0
```

First using the clinical_index to retrieve the desired medcodes before completing the rest of the query. The speed improvements are dependent on the complexity of the data, the query and the size of the data. There is a once-off execution time cost in indexing data but for large, complex datasets where queries rely on searching lists of medical codes, for instance, this may be beneficial to the user.

O SEARCH pat USING AUTOMATIC COVERING INDEX (patid=?)

If a SQL statement must be passed directly to the database, specifically for maintenance or speed improvements, the **statement_direct** function is used. If the index created previously were to be deleted, the following is used

```
## $database_tables
## [1] "Clinical" "Life_table" "Patient" "Practice" "hes_patient"
## [6] "lookup_table"
##
## $statement
## [1] "DROP INDEX clinical_index;"
##
## $query_time
## Time difference of 0.002105236 secs
```

statement_direct provides no output data and so should only be used for maintenance and speed improvement tasks such as UPDATE, DELETE, INSERT INTO, DROP TABLE etc.

Convert fields to dates

Often date fields are provided in datasets in the format of a date ($^{12}/03/1989$) but read as a character variable into the database. To convert a field to a date field use

```
data <- query_direct(db_path = DB.path,</pre>
                     query = "SELECT* FROM Practice;")$query_data
str(data)
## 'data.frame':
                    10 obs. of 4 variables:
  $ pracid: num 1 2 3 4 5 6 7 8 9 10
## $ region: num 8 8 7 8 7 5 5 1 1 2
            : chr "2015-11-21" "2016-05-09" "2015-11-20" "2016-08-26" ...
## $ 1cd
            : chr "1999-09-11" "2001-10-01" "1989-11-17" "2001-02-23" ...
## $ uts
convert_data <- convert_dates(data = data,</pre>
                              date_fields = c("lcd","uts"),
                              format = "%Y-%m-%d")$convert_data
str(convert_data)
## 'data.frame':
                    10 obs. of 4 variables:
## $ pracid: num 1 2 3 4 5 6 7 8 9 10
   $ region: num 8 8 7 8 7 5 5 1 1 2
## $ lcd
            : Date, format: "2015-11-21" "2016-05-09" ...
            : Date, format: "1999-09-11" "2001-10-01" ...
```

Here the format provided is the format the fields are prior to conversion and not the desired output required, with the default set to %d/%m/%Y.

Maintenance

in order to rename a table in the database, use

```
##
## $old_table_name
  [1] "practice"
##
## $new_table_name
## [1] "gp_pracs"
##
## $after_rename
## [1] "Clinical"
                       "Life_table"
                                       "Patient"
                                                                        "hes_patient"
                                                        "gp_pracs"
## [6] "lookup_table"
While to remove an entire table from the database, the following is used
delete_table(db_path = DB.path,
           remove tables = "gp pracs")
## $database_location
## [1] "/home/j/jcfs2/Documents/Test"
##
## $before_drop
## [1] "clinical"
                       "life_table"
                                       "patient"
                                                        "gp_pracs"
                                                                       "hes_patient"
   [6] "lookup_table"
##
## $dropped_table
## [1] "gp_pracs"
##
## $after_drop
## [1] "Clinical"
                       "Life_table"
                                       "Patient"
                                                       "hes patient"
                                                                       "lookup table"
For the complete deletion of a database, use with caution
delete_db(db_path = DB.path)
This will require the inputting of confirmation of the deletion into the Console
## -----WARNING, DELETION IN PROGRESS-----
##
## Press y+[enter] to proceed, n+[enter] to stop:
##
```

Conclusion

CPRDTools provides a comprehensive set of function to aid the general loading, maintenance and extraction of CPRD GOLD and its associated and peripheral data. These functions should provide a *toolkit* to any researcher, aiding in there ability to self-manage their data using an efficient and reliable SQLite data repository, providing tools to view, interrogate, load, extract, maintain, convert and delete in a single package.

References

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