Tidy R programming with the OMOP Common Data Model

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Welcome

Is this book for me?

We've written this book for anyone interested in working with Observational Medical Outcomes Partnership (OMOP) Common Data Model (CDM) instances using a tidyverse style approach. That is, human centered, consistent, composable, and inclusive (see Tidy design principles for more details on these principles).

New to R? We recommend you take a look at R for data science before reading this book. We assume that you have R installed together with an adequate Integrated Development Environment (IDE) such as RStudio or Positron. See this tutorial if you need guidance on how to get started. The book uses multiple packages that you will need to install. See the list in the R packages section.

New to databases? We recommend you take a look at some web tutorials on SQL, such as SQLBolt or SQLZoo to have a basic understanding of how databases work.

New to the OMOP CDM? We'd recommend you pair this book with The Book of OHDSI.

How is the book organised?

The book is divided into two parts. The first half of the book is focused on the general principles for working with databases from R. In these chapters you will see how you can use familiar tidyverse-style code to build up analytic pipelines that start with data held in a database and end with your analytic results. The second half of the book is focused on working with data in the OMOP CDM format, a widely used data format for health care data. In these chapters you will see how to work with this data format using the general principles from the first half of the book along with a set of R packages that have been built for the OMOP CDM.

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License

Code

The source code for the book can be found at this GitHub repository, please star it if you found it useful.

R Packages

This book is rendered automatically through GitHub Actions using the following version of packages:

Note: we only included the packages called explicitly in the book.

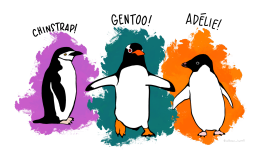
Package	Version	Link
CDMConnector	2.2.0	
CodelistGenerator	3.5.0	
CohortCharacteristics	1.0.2	
CohortConstructor	0.5.0	
DBI	1.2.3	
DatabaseConnector	7.0.0	
Lahman	13.0-0	
OmopSketch	0.5.1	
PatientProfiles	1.4.3	
RPostgres	1.4.8	
bit64	4.6.0 - 1	
cli	3.6.5	
clock	0.7.3	
dbplyr	2.5.1	
dm	1.0.12	
dplyr	1.1.4	
duckdb	1.4.1	
ggplot2	4.0.0	
nycflights13	1.0.2	
omock	0.5.0	
omopgenerics	1.3.2	
palmerpenguins	0.1.1	
purrr	1.1.0	
sloop	1.0.1	
stringr	1.5.2	
tidyr	1.3.1	

Part I Getting started with databases from R

In this first half of the book, we will explore how to work with databases from R. In the following chapters, you'll see that when working with data held in a relational database, we can leverage various open-source R packages to perform tidyverse-style data analyses.

- In Chapter 1 we will perform a simple data analysis from start to finish using a table in a database.
- In Chapter 2 we will see in more detail how familiar dplyr functions can be used to combine data spread across different tables in a database into an analytic dataset for further analysis in R.
- In Chapter 3 we will see how we can perform more complex data manipulation via translation of R code into SQL specific to the database management system being used.
- In Chapter 4 we will see how we can build data pipelines by creating a data model in R to represent the relational database we're working with and creating functions and methods to work with it.

1 A first analysis using data in a database



Artwork by @allison_horst

Before we start working with healthcare data spread across a database using the OMOP Common Data Model, let's first do a simpler analysis. In this case, we will do a quick data analysis with R using a simple dataset held in a database to understand the general approach. For this we'll use data from palmerpenguins package, which contains data on penguins collected from the Palmer Station in Antarctica.

1.1 Getting set up

```
library(dplyr)
library(dpplyr)
library(ggplot2)
library(DBI)
library(duckdb)
library(palmerpenguins)
```

1.2 Taking a peek at the data

The package palmerpenguins contains two datasets, one of them called penguins, which we will use in this chapter. We can get an overview of the data using the glimpse() command.

glimpse(penguins)

```
Rows: 344
Columns: 8
                    <fct> Adelie, Adelie, Adelie, Adelie, Adelie, Adelie, Adela-
$ species
                    <fct> Torgersen, Torgersen, Torgersen, Torgersen, Torgerse~
$ island
$ bill_length_mm
                    <dbl> 39.1, 39.5, 40.3, NA, 36.7, 39.3, 38.9, 39.2, 34.1, ~
                    <dbl> 18.7, 17.4, 18.0, NA, 19.3, 20.6, 17.8, 19.6, 18.1, ~
$ bill_depth_mm
$ flipper_length_mm <int> 181, 186, 195, NA, 193, 190, 181, 195, 193, 190, 186~
$ body_mass_g
                    <int> 3750, 3800, 3250, NA, 3450, 3650, 3625, 4675, 3475, ~
                    <fct> male, female, female, NA, female, male, female, male~
$ sex
                    <int> 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007
$ year
```

Or we could take a look at the first rows of the data using head():

```
head(penguins, 5)
```

```
# A tibble: 5 x 8
                     bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
  species island
                              <dbl>
  <fct>
          <fct>
                                             <dbl>
                                                                <int>
                                                                             <int>
1 Adelie Torgersen
                               39.1
                                              18.7
                                                                   181
                                                                              3750
2 Adelie Torgersen
                               39.5
                                              17.4
                                                                              3800
                                                                   186
3 Adelie Torgersen
                               40.3
                                              18
                                                                   195
                                                                              3250
4 Adelie Torgersen
                               NA
                                              NA
                                                                   NA
                                                                                NA
5 Adelie Torgersen
                               36.7
                                              19.3
                                                                   193
                                                                              3450
# i 2 more variables: sex <fct>, year <int>
```

1.3 Inserting data into a database

By default, the data provided by the package is local (stored in memory on your computer), so let's first put it into a DuckDB database. We need to first create the database.

```
con <- dbConnect(drv = duckdb())</pre>
```

See that now we have created an empty DuckDB database. We can easily add the penguins data to it.

```
dbWriteTable(conn = con, name = "penguins", value = penguins)
```

With the function dbListTables() we can list the tables of a database. In our case, we can see it now has one table:

```
dbListTables(conn = con)
```

[1] "penguins"

And now that the data is in a database we could use SQL to get the first rows that we saw before.

```
dbGetQuery(conn = con, statement = "SELECT * FROM penguins LIMIT 5")
```

	species	island	bill_length_mm	bill_depth_mm	flipper_length_mm	body_mass_g
1	Adelie	Torgersen	39.1	18.7	181	3750
2	Adelie	Torgersen	39.5	17.4	186	3800
3	Adelie	Torgersen	40.3	18.0	195	3250
4	Adelie	Torgersen	NA	NA	NA	NA
5	Adelie	Torgersen	36.7	19.3	193	3450
	sex v	vear				

- 1 male 2007
- 2 female 2007
- 3 female 2007
- 4 <NA> 2007
- 5 female 2007

As you can see we have the same data that we had locally but now inside the database.

? Connecting to databases from R

Database connections from R can be made using the DBI package where the back-end for DBI is facilitated by database-specific driver packages (such as RPostgres for Postgres and Amazon Redshift, and bigrquery for Google BigQuery), or the odbc R package can be used with ODBC drivers. Or instead of using ODBC drivers, JDBC drivers can be used via the DatabaseConnector R package.

In the code snippets above, we created a new, empty, in-process DuckDB via database to which we then added our dataset. But we could have instead connected to an existing duckdb database. This could, for example, look like:

```
con <- dbConnect(drv = duckdb(dbdir = "my duckdb database.ducdkb"))</pre>
```

Note that if you point to a non-existing DuckDB file, this will be created with an empty database.

In this book for simplicity we will mostly be working with in-process DuckDB databases with synthetic data. However, when analysing real patient data we will be more often working with client-server databases, where we are connecting from our computer to a central server with the database or working with data held in the cloud. The approaches shown throughout this book will work in the same way for these other types of database management systems, but the way to connect to the database will be different (although still using DBI). In general, creating connections is supported by associated back-end packages. For example a connection to a Postgres database would use the RPostgres R package and look something like:

Or if using the DatabaseConnector R package creating a connection would look something like:

1.4 Translation from R to SQL

Instead of using SQL to query our database, we might instead want to use the same R code as before. However, instead of working with the local dataset, now we will need it to query the data held in the database. To do this, first we can create a reference to the table in the database as such:

```
penguins_db <- tbl(src = con, "penguins")
penguins_db</pre>
```

```
table<penguins> [?? x 8]
# Source:
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
                     bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
   species island
   <fct>
           <fct>
                               <dbl>
                                              <dbl>
                                                                 <int>
                                                                             <int>
 1 Adelie Torgersen
                                39.1
                                               18.7
                                                                   181
                                                                              3750
2 Adelie
           Torgersen
                                39.5
                                               17.4
                                                                   186
                                                                              3800
3 Adelie Torgersen
                                40.3
                                               18
                                                                   195
                                                                              3250
4 Adelie Torgersen
                                NA
                                               NA
                                                                   NA
                                                                                NA
5 Adelie Torgersen
                                36.7
                                               19.3
                                                                   193
                                                                              3450
6 Adelie
           Torgersen
                                39.3
                                               20.6
                                                                   190
                                                                              3650
7 Adelie
           Torgersen
                                38.9
                                               17.8
                                                                   181
                                                                              3625
8 Adelie
           Torgersen
                                39.2
                                               19.6
                                                                   195
                                                                              4675
9 Adelie
           Torgersen
                                34.1
                                               18.1
                                                                   193
                                                                              3475
10 Adelie Torgersen
                                42
                                               20.2
                                                                   190
                                                                              4250
# i more rows
```

Once we have this reference, we can then use it with familiar looking R code.

i 2 more variables: sex <fct>, year <int>

```
head(penguins_db, 5)
```

```
# Source:
            SQL [?? x 8]
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
 species island
                    bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
                                            <dbl>
  <fct>
          <fct>
                              <dbl>
                                                               <int>
                                                                            <int>
1 Adelie Torgersen
                               39.1
                                              18.7
                                                                  181
                                                                             3750
2 Adelie Torgersen
                               39.5
                                             17.4
                                                                 186
                                                                             3800
                                                                 195
                                                                             3250
3 Adelie Torgersen
                               40.3
                                              18
4 Adelie Torgersen
                               NA
                                             NA
                                                                  NA
                                                                               NA
5 Adelie Torgersen
                               36.7
                                              19.3
                                                                  193
                                                                             3450
# i 2 more variables: sex <fct>, year <int>
```

The magic here is provided by the dbplyr package, which takes the R code and converts it into SQL. In this case the query looks like the SQL we wrote directly before.

```
head(penguins_db, 5) |>
show_query()
```

```
<SQL>
SELECT penguins.*
FROM penguins
LIMIT 5
```

1.5 Example analysis

More complicated SQL can also be generated by using familiar dplyr code. For example, we could get a summary of bill length by species like so:

```
penguins_db |>
  group_by(species) |>
  summarise(
    n = n(),
    min_bill_length_mm = min(bill_length_mm, na.rm = TRUE),
    mean_bill_length_mm = mean(bill_length_mm, na.rm = TRUE),
    max_bill_length_mm = max(bill_length_mm, na.rm = TRUE)
) |>
  mutate(min_max_bill_length_mm = paste0(
    min_bill_length_mm, " to ", max_bill_length_mm
)) |>
  select("species", "mean_bill_length_mm", "min_max_bill_length_mm")
```

The benefit of using dbplyr now becomes quite clear if we take a look at the corresponding SQL that is generated for us:

```
penguins_db |>
  group_by(species) |>
  summarise(
   n = n(),
   min_bill_length_mm = min(bill_length_mm, na.rm = TRUE),
  mean_bill_length_mm = mean(bill_length_mm, na.rm = TRUE),
```

```
max_bill_length_mm = max(bill_length_mm, na.rm = TRUE)
) |>
mutate(min_max_bill_length_mm = paste0(
    min_bill_length_mm, " to ", max_bill_length_mm
)) |>
select("species", "mean_bill_length_mm", "min_max_bill_length_mm") |>
show_query()

<SQL>
SELECT
```

```
SELECT
   species,
   mean_bill_length_mm,
   CONCAT_WS('', min_bill_length_mm, ' to ', max_bill_length_mm) AS min_max_bill_length_mm
FROM (
   SELECT
     species,
     COUNT(*) AS n,
     MIN(bill_length_mm) AS min_bill_length_mm,
     AVG(bill_length_mm) AS mean_bill_length_mm,
     MAX(bill_length_mm) AS max_bill_length_mm
FROM penguins
   GROUP BY species
) q01
```

Instead of having to write this somewhat complex SQL specific to DuckDB, we can use the friendlier dplyr syntax that will be more familiar if you're coming from an R programming background.

i Translation to different SQL dialects

Note this same R code will also work for other SQL dialects such as Postgres, SQL server, Snowflake and Spark. Here you can see the different generated translations:

1.5.1 Postgres

```
SELECT
  `species`,
  `mean_bill_length_mm`,
  CONCAT_WS('', `min_bill_length_mm`, ' to ', `max_bill_length_mm`) AS `min_max_bill_lengtl
FROM (
  SELECT
```

```
`species`,
    COUNT(*) AS `n`,
    MIN(`bill_length_mm`) AS `min_bill_length_mm`,
    AVG(`bill_length_mm`) AS `mean_bill_length_mm`,
    MAX(`bill_length_mm`) AS `max_bill_length_mm`
  FROM `df`
  GROUP BY `species`
) AS `q01`
1.5.2 SQL Server
<SQL>
SELECT
  `species`,
  `mean_bill_length_mm`,
  `min_bill_length_mm` + ' to ' + `max_bill_length_mm` AS `min_max_bill_length_mm`
FROM (
  SELECT
    `species`,
    COUNT_BIG(*) AS `n`,
    MIN(`bill_length_mm`) AS `min_bill_length_mm`,
    AVG(`bill_length_mm`) AS `mean_bill_length_mm`,
    MAX(`bill_length_mm`) AS `max_bill_length_mm`
  FROM `df`
  GROUP BY `species`
) AS `q01`
1.5.3 Redshift
<SQL>
SELECT
  `species`,
  `mean_bill_length_mm`,
  `min_bill_length_mm` || ' to ' || `max_bill_length_mm` AS `min_max_bill_length_mm`
FROM (
  SELECT
    `species`,
    COUNT(*) AS `n`,
    MIN(`bill_length_mm`) AS `min_bill_length_mm`,
    AVG(`bill_length_mm`) AS `mean_bill_length_mm`,
    MAX(`bill_length_mm`) AS `max_bill_length_mm`
  FROM `df`
```

```
GROUP BY `species`
) AS `q01`
1.5.4 Snowflake
<SQL>
SELECT
  `species`,
  `mean_bill_length_mm`,
  ARRAY_TO_STRING(ARRAY_CONSTRUCT_COMPACT(`min_bill_length_mm`, ' to ', `max_bill_length_mm
FROM (
  SELECT
    `species`,
    COUNT(*) AS `n`,
    MIN(`bill_length_mm`) AS `min_bill_length_mm`,
    AVG(`bill_length_mm`) AS `mean_bill_length_mm`,
    MAX(`bill_length_mm`) AS `max_bill_length_mm`
  FROM `df`
  GROUP BY `species`
) AS `q01`
1.5.5 Spark
<SQL>
SELECT
  `species`,
  `mean_bill_length_mm`,
  CONCAT_WS('', `min_bill_length_mm`, ' to ', `max_bill_length_mm`) AS `min_max_bill_lengtl
FROM (
  SELECT
    `species`,
    COUNT(*) AS `n`,
    MIN(`bill_length_mm`) AS `min_bill_length_mm`,
    AVG(`bill_length_mm`) AS `mean_bill_length_mm`,
    MAX(`bill_length_mm`) AS `max_bill_length_mm`
  FROM `df`
  GROUP BY `species`
) AS `q01`
Note that even though the different SQL statements look similar, each SQL dialect has
its own particularities. Using the dbplyr approach allows us to support multiple different
SQL dialects and back-ends by just writing R code.
```

Not having to worry about the SQL translation behind our queries allows us to query the database in a simple way even for more complex questions. For instance, suppose now that we are particularly interested in the body mass variable. We can first notice that there are a couple of missing records for this.

```
penguins_db |>
  mutate(missing_body_mass_g = if_else(is.na(body_mass_g), 1, 0)) |>
  group_by(species, missing_body_mass_g) |>
  tally()

# Source: SQL [?? x 3]

# Database: DuckDR 1 4 1 [unknown@Linux 6 11 0=1018=azure:R 4 4 1/:memory:]
```

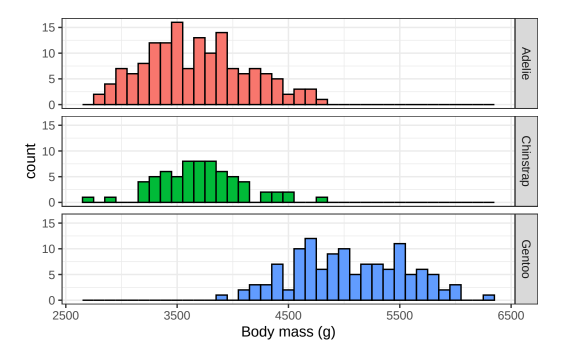
```
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
 species
            missing_body_mass_g
  <fct>
                           <dbl> <dbl>
1 Adelie
                               0
                                   151
2 Gentoo
                               0
                                   123
3 Adelie
                               1
                                      1
4 Gentoo
                                      1
                               1
                               0
5 Chinstrap
                                    68
```

We can get the mean for each of the species (dropping those two missing records).

```
penguins_db |>
  group_by(species) |>
  summarise(mean_body_mass_g = round(mean(body_mass_g, na.rm = TRUE)))
# Source: SQL [?? x 2]
```

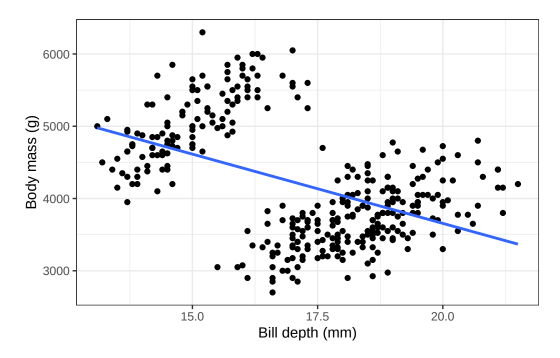
We could also make a histogram of values for each of the species using the ggplot2 package. Here we would bring our data back into R before creating our plot with the collect() function.

```
penguins_db |>
  select("species", "body_mass_g") |>
  collect() |>
  ggplot(aes(group = species, fill = species)) +
  facet_grid(species ~ .) +
  geom_histogram(aes(body_mass_g), colour = "black", binwidth = 100) +
  xlab("Body mass (g)") +
  theme_bw() +
  theme(legend.position = "none")
```



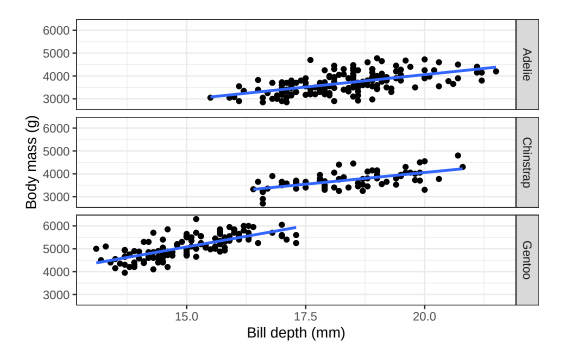
Now let's look at the relationship between body mass and bill depth.

```
penguins_db |>
  select("species", "body_mass_g", "bill_depth_mm") |>
  collect() |>
  ggplot(aes(x = bill_depth_mm, y = body_mass_g)) +
  geom_point() +
  geom_smooth(method = "lm", se = FALSE) +
  xlab("Bill_depth_(mm)") +
  ylab("Body_mass_(g)") +
  theme_bw() +
  theme(legend.position = "none")
```



Here we see a negative correlation between body mass and bill depth which seems rather unexpected. But what about if we stratify this query by species?

```
penguins_db |>
  select("species", "body_mass_g", "bill_depth_mm") |>
  collect() |>
  ggplot(aes(x = bill_depth_mm, y = body_mass_g)) +
  facet_grid(species ~ .) +
  geom_point() +
  geom_smooth(method = "lm", se = FALSE) +
  xlab("Bill depth (mm)") +
  ylab("Body mass (g)") +
  theme_bw() +
  theme(legend.position = "none")
```



As well as having an example of working with data in database from R, you also have an example of Simpson's paradox!

1.6 Disconnecting from the database

Now that we've reached the end of this example, we can close our connection to the database.

dbDisconnect(conn = con)

1.7 Further reading

- R for Data Science (Chapter 13: Relational data)
- Writing SQL with dbplyr
- Data Carpentry: SQL databases and R

2 Core verbs for analytic pipelines utilising a database

We saw in the previous chapter that we can use familiar dplyr verbs with data held in a database. In the last chapter, we were working with just a single table which we loaded into the database. When working with databases, we will typically be working with multiple tables (as we'll see later when working with data in the OMOP CDM format). For this chapter, we will see more tidyverse functionality that can be used with data in a database, this time using the nycflights13 data. As we can see, we now have a set of related tables with data on flights departing from New York City airports in 2013.

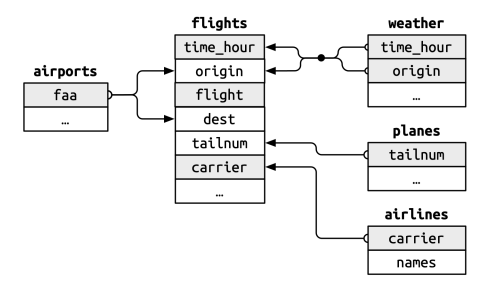


Figure 2.1: nycflights13 relational diagram from https://collinn.github.io/teaching/2023/labs/joins.html.

Let's load the required libraries, add our data to a DuckDB database, and then create references to each of these tables.

```
library(nycflights13)
library(dplyr)
library(dbplyr)
```

```
library(tidyr)
library(duckdb)
library(DBI)
# create duckdb connection
con <- dbConnect(drv = duckdb())</pre>
# copy tables in a loop
for (nm in c("airlines", "airports", "flights", "planes", "weather")) {
 dbWriteTable(conn = con, name = nm, value = get(nm))
}
airports_db <- tbl(src = con, "airports")</pre>
glimpse(airports_db)
Rows: ??
Columns: 8
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
       <chr> "04G", "06A", "06C", "06N", "09J", "0A9", "0G6", "0G7", "0P2", "~
$ faa
       <chr> "Lansdowne Airport", "Moton Field Municipal Airport", "Schaumbur~
$ name
$ lat
       <dbl> 41.13047, 32.46057, 41.98934, 41.43191, 31.07447, 36.37122, 41.4~
$ lon
       <dbl> -80.61958, -85.68003, -88.10124, -74.39156, -81.42778, -82.17342~
       <dbl> 1044, 264, 801, 523, 11, 1593, 730, 492, 1000, 108, 409, 875, 10~
$ alt
       <dbl> -5, -6, -6, -5, -5, -5, -5, -5, -5, -8, -5, -6, -5, -5, -5, -
$ tz
5, ~
       $ dst
$ tzone <chr> "America/New_York", "America/Chicago", "America/Chicago", "Ameri~
flights_db <- tbl(src = con, "flights")
glimpse(flights_db)
Rows: ??
Columns: 19
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
$ year
               <int> 2013, 2013, 2013, 2013, 2013, 2013, 2013, 2013, 2013, 2~
$ month
               $ day
               $ dep_time
               <int> 517, 533, 542, 544, 554, 554, 555, 557, 557, 558, 558, ~
$ sched_dep_time <int> 515, 529, 540, 545, 600, 558, 600, 600, 600, 600, 600, ~
               <dbl> 2, 4, 2, -1, -6, -4, -5, -3, -3, -2, -2, -2, -2, -2, -
$ dep_delay
1~
```

```
$ arr time
                <int> 830, 850, 923, 1004, 812, 740, 913, 709, 838, 753, 849,~
$ sched_arr_time <int> 819, 830, 850, 1022, 837, 728, 854, 723, 846, 745, 851,~
$ arr_delay
                <dbl> 11, 20, 33, -18, -25, 12, 19, -14, -8, 8, -2, -3, 7, -
1~
                <chr> "UA", "UA", "AA", "B6", "DL", "UA", "B6", "EV", "B6", "~
$ carrier
                <int> 1545, 1714, 1141, 725, 461, 1696, 507, 5708, 79, 301, 4~
$ flight
$ tailnum
                <chr> "N14228", "N24211", "N619AA", "N804JB", "N668DN", "N394~
$ origin
                <chr> "EWR", "LGA", "JFK", "JFK", "LGA", "EWR", "EWR", "LGA",~
$ dest
                <chr> "IAH", "IAH", "MIA", "BQN", "ATL", "ORD", "FLL", "IAD",~
$ air_time
                <dbl> 227, 227, 160, 183, 116, 150, 158, 53, 140, 138, 149, 1~
                <dbl> 1400, 1416, 1089, 1576, 762, 719, 1065, 229, 944, 733, ~
$ distance
$ hour
                <dbl> 15, 29, 40, 45, 0, 58, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0
$ minute
                <dttm> 2013-01-01 10:00:00, 2013-01-01 10:00:00, 2013-01-
$ time_hour
01 1~
weather_db <- tbl(src = con, "weather")</pre>
glimpse(weather_db)
```

```
Rows: ??
Columns: 15
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
$ origin
                               <chr> "EWR", "EW
$ year
                               <int> 2013, 2013, 2013, 2013, 2013, 2013, 2013, 2013, 2013, 2013, 2013, ~
                               $ month
                               $ day
$ hour
                               <int> 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 13, 14, 15, 16, 17, 18, ~
                               <db1> 39.02, 39.02, 39.02, 39.92, 39.02, 37.94, 39.02, 39.92, 39.~
$ temp
                               <dbl> 26.06, 26.96, 28.04, 28.04, 28.04, 28.04, 28.04, 28.04, 28.04
$ dewp
                               <dbl> 59.37, 61.63, 64.43, 62.21, 64.43, 67.21, 64.43, 62.21, 62.~
$ humid
$ wind_dir
                               <dbl> 270, 250, 240, 250, 260, 240, 240, 250, 260, 260, 260, 330,~
$ wind_speed <dbl> 10.35702, 8.05546, 11.50780, 12.65858, 12.65858, 11.50780, ~
$ wind_gust
                               $ precip
$ pressure
                               <dbl> 1012.0, 1012.3, 1012.5, 1012.2, 1011.9, 1012.4, 1012.2, 101~
                               $ visib
                               <dttm> 2013-01-01 06:00:00, 2013-01-01 07:00:00, 2013-01-01 08:00~
$ time hour
```

```
planes_db <- tbl(src = con, "planes")
glimpse(planes_db)</pre>
```

Rows: ??

```
Columns: 9
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
$ tailnum
             <chr> "N10156", "N102UW", "N103US", "N104UW", "N10575", "N105UW~
$ year
             <int> 2004, 1998, 1999, 1999, 2002, 1999, 1999, 1999, 1999, 1999
             <chr> "Fixed wing multi engine", "Fixed wing multi engine", "Fi~
$ type
$ manufacturer <chr> "EMBRAER", "AIRBUS INDUSTRIE", "AIRBUS INDUSTRIE", "AIRBU~
$ model
             <chr> "EMB-145XR", "A320-214", "A320-214", "A320-214", "EMB-
145~
$ engines
             $ seats
             <int> 55, 182, 182, 182, 55, 182, 182, 182, 182, 182, 55, 55, 5~
             $ speed
$ engine
             <chr> "Turbo-fan", "Turbo-fan", "Turbo-fan", "Turbo-fan", "Turb~
airlines_db <- tbl(src = con, "airlines")</pre>
glimpse(airlines_db)
```

2.1 Tidyverse functions

For almost all analyses, we want to go from having our starting data spread out across multiple tables in the database to a single tidy table containing all the data we need for the specific analysis. We can often get to our tidy analytic dataset using the tidyverse functions below (most of which come from dplyr, but a couple also from the tidyr package). These functions all work with data in a database by generating SQL that will have the same purpose as if these functions were being run against data in R.

Important

Until we use compute() or collect() (or printing the first few rows of the result), all we're doing is translating R code into SQL. This means no code is being executed on the database side.

- compute() will execute the query and store it in a new table in the database.
- collect() will execute the query and bring the result back to R.
- printing (e.g. glimpse() or print()) will execute the query, limiting the result to

the first set of rows, which leads to shorter computation time on the database side.

Purpose	Functions	Description
Selecting rows	filter, distinct	To select rows in a table.
Ordering rows	arrange	To order rows in a table.
Column Transformation	mutate, select, relocate, rename	To create new columns or change existing ones.
Grouping and ungrouping	group_by, rowwise, ungroup	To group data by one or more variables and to
Aggregation	count, tally, summarise	remove grouping. These functions are used for summarising data.
Data merging and joining	inner_join, left_join, right_join, full_join, anti_join, semi_join, cross_join	These functions are used to combine data from different tables based on
Data reshaping	pivot_wider, pivot_longer	common columns. These functions are used to reshape data between wide and long formats.
Data union	union_all, union	This function combines two tables.
Randomly selects rows	slice_sample	We can use this to take a random subset a table.

? Behind the scenes

By using the above functions we can use the same code regardless of whether the data is held in the database or locally in R. This is because the functions used above are generic functions which behave differently depending on the type of input they are given. Let's take <code>inner_join()</code> for example. We can see that this function is a S3 generic function (with S3 being the most common object-oriented system used in R).

```
library(sloop)
ftype(inner_join)
[1] "S3"
               "generic"
Among others, the references we create to tables in a database have tbl_lazy as a class
attribute. Meanwhile, we can see that when collected into R, the object changes to have
different attributes, one of which is data.frame:
class(flights_db)
                                                       "tbl_sql"
[1] "tbl_duckdb_connection" "tbl_dbi"
[4] "tbl_lazy"
                              "tbl"
class(flights_db |> head(1) |> collect())
[1] "tbl_df"
                  "tbl"
                                "data.frame"
We can see that inner_join() has different methods for tbl_lazy and data.frame.
s3_methods_generic("inner_join")
# A tibble: 2 x 4
 generic class
                         visible source
  <chr>
             <chr>
                         <lg1> <chr>
1 inner_join data.frame FALSE registered S3method
2 inner_join tbl_lazy
                         FALSE
                                  registered S3method
When working with references to tables in the database the tbl_lazy method will be
used.
s3_dispatch(flights_db |>
               inner_join(planes_db))
   inner_join.tbl_duckdb_connection
```

inner_join.tbl_dbi
inner_join.tbl_sql
=> inner_join.tbl_lazy
inner_join.tbl
inner_join.default

But once we bring data into R, the data.frame method will be used.

2.2 Getting to an analytic dataset

To see a little more on how we can use the above functions, let's say we want to do an analysis of late flights from JFK airport. We want to see whether there is some relationship between plane characteristics and the risk of delay.

For this, we'll first use the filter() and select() dplyr verbs to get the data from the flights table. Note, we'll rename arr_delay to just delay.

```
delayed_flights_db <- flights_db |>
  filter(!is.na(arr_delay) & origin == "JFK") |>
  select("dest", "distance", "carrier", "tailnum", "delay" = "arr_delay")
```

```
i Show query
See the resultant DuckDB query:

<SQL>
SELECT dest, distance, carrier, tailnum, arr_delay AS delay
FROM flights
WHERE (NOT((arr_delay IS NULL)) AND origin = 'JFK')
```

When executed, our results will look like the following:

```
1 MIA
             1089 AA
                           N619AA
                                       33
2 BQN
             1576 B6
                           N804JB
                                      -18
3 MCO
              944 B6
                           N593JB
                                       -8
4 PBI
             1028 B6
                           N793JB
                                       -2
                                       -3
5 TPA
             1005 B6
                           N657JB
             2475 UA
                           N29129
                                        7
6 LAX
7 BOS
              187 B6
                           N708JB
                                       -4
8 ATL
              760 DL
                           N3739P
                                       -8
9 SFO
             2586 UA
                           N532UA
                                       14
10 RSW
             1074 B6
                           N635JB
                                        4
# i more rows
```

Now we'll add plane characteristics from the planes table. We will use an inner join so that only records for which we have the plane characteristics are kept.

```
delayed_flights_db <- delayed_flights_db |>
  inner_join(
  planes_db |>
     select("tailnum", "seats"),
  by = "tailnum"
)
```

Note that our first query was not executed, as we didn't use either compute() or collect(), so we'll now have added our join to the original query.

```
i Show query

See that now the SQL code combines both queries:

<SQL>
SELECT LHS.*, seats
FROM (
    SELECT dest, distance, carrier, tailnum, arr_delay AS delay
    FROM flights
    WHERE (NOT((arr_delay IS NULL)) AND origin = 'JFK')
) LHS
INNER JOIN planes
    ON (LHS.tailnum = planes.tailnum)
```

And when executed, our results will look like the following:

delayed_flights_db

```
# Source:
             SQL [?? x 6]
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
   dest
         distance carrier tailnum delay seats
   <chr>
             <dbl> <chr>
                            <chr>
                                     <dbl> <int>
 1 MIA
              1089 AA
                            N619AA
                                        33
                                             178
 2 BQN
              1576 B6
                            N804JB
                                       -18
                                             200
3 MCO
               944 B6
                            N593JB
                                        -8
                                             200
4 PBI
              1028 B6
                                        -2
                            N793JB
                                             200
5 TPA
              1005 B6
                            N657JB
                                        -3
                                             200
 6 LAX
              2475 UA
                            N29129
                                         7
                                             178
7 BOS
               187 B6
                            N708JB
                                        -4
                                             200
8 ATL
               760 DL
                            N3739P
                                        -8
                                             189
                                         4
9 RSW
              1074 B6
                            N635JB
                                             200
10 SJU
              1598 B6
                            N794JB
                                       -21
                                             200
# i more rows
```

This tidy dataset has been created in the database via R code translated to SQL. With this, we can now collect our analytic dataset into R and proceed from there (for example, to perform statistical analyses locally that might not be possible to run in a database, such as plots, density distributions, regression, or anything beyond data manipulation).

```
delayed_flights <- delayed_flights_db |>
   collect()

glimpse(delayed_flights)
```

2.3 Disconnecting from the database

Now that we've reached the end of this example, we can close our connection to the database.

2.4 Further reading

- Wickham H, François R, Henry L, Müller K, Vaughan D (2025). dplyr: A Grammar of Data Manipulation. R package version 1.1.4, https://dplyr.tidyverse.org
- Wickham H, Vaughan D, Girlich M (2025). *tidyr: Tidy Messy Data*. R package version 1.3.1, https://tidyr.tidyverse.org.

3 Supported expressions for database queries

In the previous chapter, Chapter 2, we saw that there are a core set of tidyverse functions that can be used with databases to extract data for analysis. The SQL code used in the previous chapter is consistent across database management systems, since it only involves basic operations such as joins and variable selection.

For more complex data pipelines, we will, however, often need to incorporate additional expressions within these functions. Because of differences across database management systems, the translated SQL can vary. Moreover, some expressions are only supported for some databases.

When writing code that should work across different database management systems, we need to keep in mind which expressions are supported where. To help with this, the sections below show the available SQL translations for common expressions we might want to use.

Let's first load the packages which these expressions come from. In addition to base R types, bit64 adds support for integer64. The stringr package provides functions for working with strings, while clock has various functions for working with dates. Many other useful expressions will come from dplyr itself.

```
library(duckdb)
library(bit64)
library(dplyr)
library(dsplyr)
library(stringr)
library(clock)
```

3.1 Data types

Commonly used data types are consistently supported across database backends. We can use the base as.numeric(), as.integer(), as.character(), as.Date(), and as.POSIXct(). We can also use as.integer64() from the bit64 package to coerce to integer64, and the as_date() and as_datetime() from the clock package instead of as.Date() and as.POSIXct(), respectively.

```
    Show SQL

3.1.1 DuckDB
con <- simulate_duckdb()</pre>
translate_sql(as.numeric(var), con = con)
<SQL> CAST(`var` AS NUMERIC)
translate_sql(as.integer(var), con = con)
<SQL> CAST(`var` AS INTEGER)
translate_sql(as.integer64(var), con = con)
<SQL> CAST(`var` AS BIGINT)
translate_sql(as.character(var), con = con)
<SQL> CAST(`var` AS TEXT)
translate_sql(as.Date(var), con = con)
<SQL> CAST(`var` AS DATE)
translate_sql(as_date(var), con = con)
<SQL> CAST(`var` AS DATE)
translate_sql(as.POSIXct(var), con = con)
<SQL> CAST(`var` AS TIMESTAMP)
```

translate_sql(as_datetime(var), con = con)

translate_sql(as.logical(var), con = con)

<SQL> CAST(`var` AS TIMESTAMP)

<SQL> CAST(`var` AS BOOLEAN)

3.1.2 Postgres

```
con <- simulate_postgres()</pre>
translate_sql(as.numeric(var), con = con)
<SQL> CAST(`var` AS NUMERIC)
translate_sql(as.integer(var), con = con)
<SQL> CAST(`var` AS INTEGER)
translate_sql(as.integer64(var), con = con)
<SQL> CAST(`var` AS BIGINT)
translate_sql(as.character(var), con = con)
<SQL> CAST(`var` AS TEXT)
translate_sql(as.Date(var), con = con)
<SQL> CAST(`var` AS DATE)
translate_sql(as_date(var), con = con)
<SQL> CAST(`var` AS DATE)
translate_sql(as.POSIXct(var), con = con)
<SQL> CAST(`var` AS TIMESTAMP)
translate_sql(as_datetime(var), con = con)
<SQL> CAST(`var` AS TIMESTAMP)
translate_sql(as.logical(var), con = con)
<SQL> CAST(`var` AS BOOLEAN)
```

3.1.3 SQL Server

```
con <- simulate_mssql()</pre>
translate_sql(as.numeric(var), con = con)
<SQL> TRY_CAST(`var` AS FLOAT)
translate_sql(as.integer(var), con = con)
<SQL> TRY_CAST(TRY_CAST(`var` AS NUMERIC) AS INT)
translate_sql(as.integer64(var), con = con)
<SQL> TRY_CAST(TRY_CAST(`var` AS NUMERIC(38, 0)) AS BIGINT)
translate_sql(as.character(var), con = con)
<SQL> TRY_CAST(`var` AS VARCHAR(MAX))
translate_sql(as.Date(var), con = con)
<SQL> TRY_CAST(`var` AS DATE)
translate_sql(as_date(var), con = con)
<SQL> TRY_CAST(`var` AS DATE)
translate_sql(as.POSIXct(var), con = con)
<SQL> TRY_CAST(`var` AS DATETIME2)
translate_sql(as_datetime(var), con = con)
<SQL> TRY_CAST(`var` AS DATETIME2)
translate_sql(as.logical(var), con = con)
<SQL> TRY_CAST(`var` AS BIT)
```

3.1.4 Redshift

```
con <- simulate_redshift()</pre>
translate_sql(as.numeric(var), con = con)
<SQL> CAST(`var` AS FLOAT)
translate_sql(as.integer(var), con = con)
<SQL> CAST(`var` AS INTEGER)
translate_sql(as.integer64(var), con = con)
<SQL> CAST(`var` AS BIGINT)
translate_sql(as.character(var), con = con)
<SQL> CAST(`var` AS TEXT)
translate_sql(as.Date(var), con = con)
<SQL> CAST(`var` AS DATE)
translate_sql(as_date(var), con = con)
<SQL> CAST(`var` AS DATE)
translate_sql(as.POSIXct(var), con = con)
<SQL> CAST(`var` AS TIMESTAMP)
translate_sql(as_datetime(var), con = con)
<SQL> CAST(`var` AS TIMESTAMP)
translate_sql(as.logical(var), con = con)
<SQL> CAST(`var` AS BOOLEAN)
```

3.1.5 Snowflake

```
con <- simulate_snowflake()</pre>
translate_sql(as.numeric(var), con = con)
<SQL> CAST(`var` AS DOUBLE)
translate_sql(as.integer(var), con = con)
<SQL> CAST(`var` AS INT)
translate_sql(as.integer64(var), con = con)
<SQL> CAST(`var` AS BIGINT)
translate_sql(as.character(var), con = con)
<SQL> CAST(`var` AS STRING)
translate_sql(as.Date(var), con = con)
<SQL> CAST(`var` AS DATE)
translate_sql(as_date(var), con = con)
<SQL> CAST(`var` AS DATE)
translate_sql(as.POSIXct(var), con = con)
<SQL> CAST(`var` AS TIMESTAMP)
translate_sql(as_datetime(var), con = con)
<SQL> CAST(`var` AS TIMESTAMP)
translate_sql(as.logical(var), con = con)
<SQL> CAST(`var` AS BOOLEAN)
```

3.1.6 **Spark** con <- simulate_spark_sql()</pre> translate_sql(as.numeric(var), con = con) <SQL> CAST(`var` AS DOUBLE) translate_sql(as.integer(var), con = con) <SQL> CAST(`var` AS INT) translate_sql(as.integer64(var), con = con) <SQL> CAST(`var` AS BIGINT) translate_sql(as.character(var), con = con) <SQL> CAST(`var` AS STRING) translate_sql(as.Date(var), con = con) <SQL> CAST(`var` AS DATE) translate_sql(as_date(var), con = con) <SQL> CAST(`var` AS DATE) translate_sql(as.POSIXct(var), con = con) <SQL> CAST(`var` AS TIMESTAMP) translate_sql(as_datetime(var), con = con) <SQL> CAST(`var` AS TIMESTAMP) translate_sql(as.logical(var), con = con) <SQL> CAST(`var` AS BOOLEAN)

3.2 Comparison and logical operators

Base R comparison operators, such as <, <=, ==, >=, >=, are also well supported in all database backends. Logical operators, such as & and |, can also be used as if the data were in R.

```
Show SQL
3.2.1 DuckDB
con <- simulate_duckdb()</pre>
translate_sql(var_1 == var_2, con = con)
<SQL> `var_1` = `var_2`
translate_sql(var_1 >= var_2, con = con)
<SQL> `var_1` >= `var_2`
translate_sql(var_1 < 100, con = con)</pre>
<SQL> `var_1` < 100.0
translate_sql(var_1 %in% c("a", "b", "c"), con = con)
<SQL> `var_1` IN ('a', 'b', 'c')
translate_sql(!var_1 %in% c("a", "b", "c"), con = con)
<SQL> NOT(`var_1` IN ('a', 'b', 'c'))
translate_sql(is.na(var_1), con = con)
<SQL> (`var_1` IS NULL)
translate_sql(!is.na(var_1), con = con)
<SQL> NOT((`var_1` IS NULL))
translate_sql(var_1 >= 100 \& var_1 < 200, con = con)
<SQL> `var_1` >= 100.0 AND `var_1` < 200.0
translate_sql(var_1 >= 100 \mid var_1 < 200, con = con)
<SQL> `var_1` >= 100.0 OR `var_1` < 200.0
```

3.2.2 Postgres

```
con <- simulate_postgres()</pre>
translate_sql(var_1 == var_2, con = con)
\langle SQL \rangle  var_1 = var_2 
translate_sql(var_1 >= var_2, con = con)
<SQL> `var_1` >= `var_2`
translate_sql(var_1 < 100, con = con)</pre>
<SQL> `var_1` < 100.0
translate_sql(var_1 %in% c("a", "b", "c"), con = con)
<SQL> `var_1` IN ('a', 'b', 'c')
translate_sql(!var_1 %in% c("a", "b", "c"), con = con)
<SQL> NOT(`var_1` IN ('a', 'b', 'c'))
translate_sql(is.na(var_1), con = con)
<SQL> (`var_1` IS NULL)
translate_sql(!is.na(var_1), con = con)
<SQL> NOT(('var_1' IS NULL))
translate_sql(var_1 \ge 100 \& var_1 < 200, con = con)
<SQL> `var_1` >= 100.0 AND `var_1` < 200.0
translate_sql(var_1 >= 100 \mid var_1 < 200, con = con)
<SQL> `var_1` >= 100.0 OR `var_1` < 200.0
```

3.2.3 SQL Server

```
con <- simulate_mssql()</pre>
translate_sql(var_1 == var_2, con = con)
\langle SQL \rangle  var_1 = var_2 
translate_sql(var_1 >= var_2, con = con)
<SQL> `var_1` >= `var_2`
translate_sql(var_1 < 100, con = con)</pre>
<SQL> `var_1` < 100.0
translate_sql(var_1 %in% c("a", "b", "c"), con = con)
<SQL> `var_1` IN ('a', 'b', 'c')
translate_sql(!var_1 %in% c("a", "b", "c"), con = con)
<SQL> NOT(`var_1` IN ('a', 'b', 'c'))
translate_sql(is.na(var_1), con = con)
<SQL> (`var_1` IS NULL)
translate_sql(!is.na(var_1), con = con)
<SQL> NOT(('var_1' IS NULL))
translate_sql(var_1 \ge 100 \& var_1 < 200, con = con)
<SQL> `var_1` >= 100.0 AND `var_1` < 200.0
translate_sql(var_1 >= 100 \mid var_1 < 200, con = con)
<SQL> `var_1` >= 100.0 OR `var_1` < 200.0
```

3.2.4 Redshift

```
con <- simulate redshift()</pre>
translate_sql(var_1 == var_2, con = con)
\langle SQL \rangle  var_1 = var_2 
translate_sql(var_1 >= var_2, con = con)
<SQL> `var_1` >= `var_2`
translate_sql(var_1 < 100, con = con)</pre>
<SQL> `var_1` < 100.0
translate_sql(var_1 %in% c("a", "b", "c"), con = con)
<SQL> `var_1` IN ('a', 'b', 'c')
translate_sql(!var_1 %in% c("a", "b", "c"), con = con)
<SQL> NOT(`var_1` IN ('a', 'b', 'c'))
translate_sql(is.na(var_1), con = con)
<SQL> (`var_1` IS NULL)
translate_sql(!is.na(var_1), con = con)
<SQL> NOT(('var_1' IS NULL))
translate_sql(var_1 \ge 100 \& var_1 < 200, con = con)
<SQL> `var_1` >= 100.0 AND `var_1` < 200.0
translate_sql(var_1 >= 100 \mid var_1 < 200, con = con)
<SQL> `var_1` >= 100.0 OR `var_1` < 200.0
```

3.2.5 Snowflake

```
con <- simulate snowflake()</pre>
translate_sql(var_1 == var_2, con = con)
\langle SQL \rangle  var_1 = var_2 
translate_sql(var_1 >= var_2, con = con)
<SQL> `var_1` >= `var_2`
translate_sql(var_1 < 100, con = con)</pre>
<SQL> `var_1` < 100.0
translate_sql(var_1 %in% c("a", "b", "c"), con = con)
<SQL> `var_1` IN ('a', 'b', 'c')
translate_sql(!var_1 %in% c("a", "b", "c"), con = con)
<SQL> NOT(`var_1` IN ('a', 'b', 'c'))
translate_sql(is.na(var_1), con = con)
<SQL> (`var_1` IS NULL)
translate_sql(!is.na(var_1), con = con)
<SQL> NOT(('var_1' IS NULL))
translate_sql(var_1 \ge 100 \& var_1 < 200, con = con)
<SQL> `var_1` >= 100.0 AND `var_1` < 200.0
translate_sql(var_1 >= 100 \mid var_1 < 200, con = con)
<SQL> `var_1` >= 100.0 OR `var_1` < 200.0
```

3.2.6 **Spark** con <- simulate_spark_sql()</pre> translate_sql(var_1 == var_2, con = con) <SQL> `var_1` = `var_2` translate_sql(var_1 >= var_2, con = con) <SQL> `var_1` >= `var_2` translate_sql(var_1 < 100, con = con)</pre> <SQL> `var_1` < 100.0 translate_sql(var_1 %in% c("a", "b", "c"), con = con) <SQL> `var_1` IN ('a', 'b', 'c') translate_sql(!var_1 %in% c("a", "b", "c"), con = con) <SQL> NOT(`var_1` IN ('a', 'b', 'c')) translate_sql(is.na(var_1), con = con) <SQL> (`var_1` IS NULL) translate_sql(!is.na(var_1), con = con) <SQL> NOT((`var_1` IS NULL)) $translate_sql(var_1 >= 100 \& var_1 < 200, con = con)$ <SQL> `var_1` >= 100.0 AND `var_1` < 200.0 $translate_sql(var_1 >= 100 \mid var_1 < 200, con = con)$ <SQL> `var_1` >= 100.0 OR `var_1` < 200.0

3.3 Conditional statements

The base ifelse function, along with if_else and case_when from dplyr are translated for each database backend. As can be seen in the translations, case_when maps to the SQL CASE WHEN statement.

```
    Show SQL

3.3.1 DuckDB
con <- simulate_duckdb()</pre>
translate_sql(ifelse(var == "a", 1L, 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 WHEN NOT ('var' = 'a') THEN 2 END
translate_sql(if_else(var == "a", 1L, 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 WHEN NOT ('var' = 'a') THEN 2 END
translate_sql(case_when(var == "a" ~ 1L, .default = 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 ELSE 2 END
translate_sql(case_when(var == "a" ~ 1L, var == "b" ~ 2L, var == "c" ~ 3L, .default = NULL)
              con = con)
<SQL> CASE
WHEN ('var' = 'a') THEN 1
WHEN (`var` = 'b') THEN 2
WHEN ('var' = 'c') THEN 3
END
translate_sql(case_when(var == "a" ~ 1L, var == "b" ~ 2L, var == "c" ~ 3L, .default = "some
              con = con)
<SQL> CASE
WHEN ('var' = 'a') THEN 1
WHEN (`var` = 'b') THEN 2
WHEN ('var' = 'c') THEN 3
ELSE 'something else'
END
```

```
3.3.2 Postgres
```

```
con <- simulate_postgres()</pre>
translate_sql(ifelse(var == "a", 1L, 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 WHEN NOT ('var' = 'a') THEN 2 END
translate_sql(if_else(var == "a", 1L, 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 WHEN NOT ('var' = 'a') THEN 2 END
translate_sql(case_when(var == "a" ~ 1L, .default = 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 ELSE 2 END
translate sql(case when(var == "a" ~ 1L, var == "b" ~ 2L, var == "c" ~ 3L, .default = NULL
              con = con)
<SQL> CASE
WHEN ('var' = 'a') THEN 1
WHEN ('var' = 'b') THEN 2
WHEN ('var' = 'c') THEN 3
END
translate_sql(case_when(var == "a" ~ 1L, var == "b" ~ 2L, var == "c" ~ 3L, .default = "some
              con = con)
<SQL> CASE
WHEN ('var' = 'a') THEN 1
WHEN ('var' = 'b') THEN 2
WHEN ('var' = 'c') THEN 3
ELSE 'something else'
END
3.3.3 SQL Server
con <- simulate_mssql()</pre>
translate_sql(ifelse(var == "a", 1L, 2L), con = con)
\langle SQL \rangle IIF(`var` = 'a', 1, 2)
```

```
translate_sql(if_else(var == "a", 1L, 2L), con = con)
\langle SQL \rangle IIF(`var` = 'a', 1, 2)
translate_sql(case_when(var == "a" ~ 1L, .default = 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 ELSE 2 END
translate_sql(case_when(var == "a" ~ 1L, var == "b" ~ 2L, var == "c" ~ 3L, .default = NULL)
              con = con)
<SQL> CASE
WHEN ('var' = 'a') THEN 1
WHEN ('var' = 'b') THEN 2
WHEN ('var' = 'c') THEN 3
END
translate sql(case when(var == "a" ~ 1L, var == "b" ~ 2L, var == "c" ~ 3L, .default = "some
              con = con)
<SQL> CASE
WHEN ('var' = 'a') THEN 1
WHEN ('var' = 'b') THEN 2
WHEN ('var' = 'c') THEN 3
ELSE 'something else'
END
3.3.4 Redshift
con <- simulate_redshift()</pre>
translate_sql(ifelse(var == "a", 1L, 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 WHEN NOT ('var' = 'a') THEN 2 END
translate_sql(if_else(var == "a", 1L, 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 WHEN NOT ('var' = 'a') THEN 2 END
translate_sql(case_when(var == "a" ~ 1L, .default = 2L), con = con)
```

```
<SQL> CASE WHEN ('var' = 'a') THEN 1 ELSE 2 END
translate_sql(case_when(var == "a" ~ 1L, var == "b" ~ 2L, var == "c" ~ 3L, .default = NULL)
              con = con)
<SQL> CASE
WHEN ('var' = 'a') THEN 1
WHEN ('var' = 'b') THEN 2
WHEN ('var' = 'c') THEN 3
END
translate sql(case when(var == "a" ~ 1L, var == "b" ~ 2L, var == "c" ~ 3L, .default = "some
              con = con)
<SQL> CASE
WHEN ('var' = 'a') THEN 1
WHEN ('var' = 'b') THEN 2
WHEN ('var' = 'c') THEN 3
ELSE 'something else'
END
3.3.5 Snowflake
con <- simulate_snowflake()</pre>
translate_sql(ifelse(var == "a", 1L, 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 WHEN NOT ('var' = 'a') THEN 2 END
translate_sql(if_else(var == "a", 1L, 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 WHEN NOT ('var' = 'a') THEN 2 END
translate_sql(case_when(var == "a" ~ 1L, .default = 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 ELSE 2 END
translate sql(case when(var == "a" ~ 1L, var == "b" ~ 2L, var == "c" ~ 3L, .default = NULL
              con = con)
```

```
<SQL> CASE
WHEN ('var' = 'a') THEN 1
WHEN ('var' = 'b') THEN 2
WHEN ('var' = 'c') THEN 3
END
translate_sql(case_when(var == "a" ~ 1L, var == "b" ~ 2L, var == "c" ~ 3L, .default = "some
             con = con)
<SQL> CASE
WHEN ('var' = 'a') THEN 1
WHEN (`var` = 'b') THEN 2
WHEN ('var' = 'c') THEN 3
ELSE 'something else'
END
3.3.6 Spark
con <- simulate_spark_sql()</pre>
translate_sql(ifelse(var == "a", 1L, 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 WHEN NOT ('var' = 'a') THEN 2 END
translate_sql(if_else(var == "a", 1L, 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 WHEN NOT ('var' = 'a') THEN 2 END
translate_sql(case_when(var == "a" ~ 1L, .default = 2L), con = con)
<SQL> CASE WHEN ('var' = 'a') THEN 1 ELSE 2 END
translate_sql(case_when(var == "a" ~ 1L, var == "b" ~ 2L, var == "c" ~ 3L, .default = NULL)
              con = con)
<SQL> CASE
WHEN ('var' = 'a') THEN 1
WHEN ('var' = 'b') THEN 2
WHEN ('var' = 'c') THEN 3
END
```

3.4 Working with strings

Compared to the previous sections, there is much more variation in support of functions to work with strings across database management systems. In particular, although various useful **stringr** functions do have translations ubiquitously, it can be seen below that more translations are available for some databases compared to others.

```
Show SQL

3.4.1 DuckDB

con <- simulate_duckdb()
translate_sql(nchar(var), con = con)

<SQL> LENGTH(`var`)

translate_sql(nzchar(var), con = con)

<SQL> ((`var` IS NULL) OR `var` != '')

translate_sql(substr(var, 1, 2), con = con)

<SQL> SUBSTR(`var`, 1, 2)

translate_sql(trimws(var), con = con)

<SQL> LTRIM(RTRIM(`var`))

translate_sql(tolower(var), con = con)
```

```
<SQL> LOWER(`var`)
translate_sql(str_to_lower(var), con = con)
<SQL> LOWER(`var`)
translate_sql(toupper(var), con = con)
<SQL> UPPER(`var`)
translate_sql(str_to_upper(var), con = con)
<SQL> UPPER(`var`)
translate_sql(str_to_title(var), con = con)
<SQL> INITCAP(`var`)
translate_sql(str_trim(var), con = con)
<SQL> LTRIM(RTRIM(`var`))
translate_sql(str_squish(var), con = con)
<SQL> TRIM(REGEXP_REPLACE(`var`, '\s+', ' ', 'g'))
translate_sql(str_detect(var, "b"), con = con)
<SQL> REGEXP_MATCHES(`var`, 'b')
translate_sql(str_detect(var, "b", negate = TRUE), con = con)
<SQL> (NOT(REGEXP_MATCHES(`var`, 'b')))
translate_sql(str_detect(var, "[aeiou]"), con = con)
<SQL> REGEXP_MATCHES(`var`, '[aeiou]')
```

```
translate_sql(str_replace(var, "a", "b"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', 'b')
translate_sql(str_replace_all(var, "a", "b"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', 'b', 'g')
translate_sql(str_remove(var, "a"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', '')
translate_sql(str_remove_all(var, "a"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', '', 'g')
translate_sql(str_like(var, "a"), con = con)
<SQL> `var` LIKE 'a'
translate_sql(str_starts(var, "a"), con = con)
<SQL> REGEXP_MATCHES(`var`, '^(?:' || 'a' || ')')
translate_sql(str_ends(var, "a"), con = con)
<SQL> REGEXP_MATCHES(`var`, '(?:' || 'a' || ')$')
3.4.2 Postgres
con <- simulate_postgres()</pre>
translate_sql(nchar(var), con = con)
<SQL> LENGTH(`var`)
translate_sql(nzchar(var), con = con)
<SQL> (('var' IS NULL) OR 'var' != '')
```

```
translate_sql(substr(var, 1, 2), con = con)
<SQL> SUBSTR(`var`, 1, 2)
translate_sql(trimws(var), con = con)
<SQL> LTRIM(RTRIM(`var`))
translate_sql(tolower(var), con = con)
<SQL> LOWER(`var`)
translate_sql(str_to_lower(var), con = con)
<SQL> LOWER(`var`)
translate_sql(toupper(var), con = con)
<SQL> UPPER(`var`)
translate_sql(str_to_upper(var), con = con)
<SQL> UPPER(`var`)
translate_sql(str_to_title(var), con = con)
<SQL> INITCAP(`var`)
translate_sql(str_trim(var), con = con)
<SQL> LTRIM(RTRIM(`var`))
translate_sql(str_squish(var), con = con)
<SQL> LTRIM(RTRIM(REGEXP_REPLACE(`var`, '\s+', ' ', 'g')))
translate_sql(str_detect(var, "b"), con = con)
```

```
<SQL> `var` ~ 'b'
translate_sql(str_detect(var, "b", negate = TRUE), con = con)
<SQL> !(`var` ~ 'b')
translate_sql(str_detect(var, "[aeiou]"), con = con)
<SQL> `var` ~ '[aeiou]'
translate_sql(str_replace(var, "a", "b"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', 'b')
translate_sql(str_replace_all(var, "a", "b"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', 'b', 'g')
translate_sql(str_remove(var, "a"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', '')
translate_sql(str_remove_all(var, "a"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', '', 'g')
translate_sql(str_like(var, "a"), con = con)
<SQL> `var` ILIKE 'a'
translate_sql(str_starts(var, "a"), con = con)
Error in `str_starts()`:
! Only fixed patterns are supported on database backends.
translate_sql(str_ends(var, "a"), con = con)
Error in `str_ends()`:
! Only fixed patterns are supported on database backends.
```

3.4.3 SQL Server con <- simulate_mssql()</pre> translate_sql(nchar(var), con = con) <SQL> LEN(`var`) translate_sql(nzchar(var), con = con) <SQL> (('var' IS NULL) OR 'var' != '') translate_sql(substr(var, 1, 2), con = con) <SQL> SUBSTRING(`var`, 1, 2) translate_sql(trimws(var), con = con) <SQL> LTRIM(RTRIM(`var`)) translate_sql(tolower(var), con = con) <SQL> LOWER(`var`) translate_sql(str_to_lower(var), con = con) <SQL> LOWER(`var`) translate_sql(toupper(var), con = con) <SQL> UPPER(`var`) translate_sql(str_to_upper(var), con = con) <SQL> UPPER(`var`)

translate_sql(str_to_title(var), con = con)

! `str_to_title()` is not available in this SQL variant.

Error in `str_to_title()`:

```
translate_sql(str_trim(var), con = con)
<SQL> LTRIM(RTRIM(`var`))
translate_sql(str_squish(var), con = con)
Error in `str_squish()`:
! `str_squish()` is not available in this SQL variant.
translate_sql(str_detect(var, "b"), con = con)
Error in `str_detect()`:
! Only fixed patterns are supported on database backends.
translate_sql(str_detect(var, "b", negate = TRUE), con = con)
Error in `str_detect()`:
! Only fixed patterns are supported on database backends.
translate_sql(str_detect(var, "[aeiou]"), con = con)
Error in `str_detect()`:
! Only fixed patterns are supported on database backends.
translate_sql(str_replace(var, "a", "b"), con = con)
Error in `str_replace()`:
! `str_replace()` is not available in this SQL variant.
translate_sql(str_replace_all(var, "a", "b"), con = con)
Error in `str_replace_all() `:
! `str_replace_all()` is not available in this SQL variant.
translate_sql(str_remove(var, "a"), con = con)
Error in `str_remove()`:
! `str_remove()` is not available in this SQL variant.
```

```
translate_sql(str_remove_all(var, "a"), con = con)
Error in `str_remove_all()`:
! `str_remove_all()` is not available in this SQL variant.
translate_sql(str_like(var, "a"), con = con)
<SQL> `var` LIKE 'a'
translate_sql(str_starts(var, "a"), con = con)
Error in `str_starts()`:
! Only fixed patterns are supported on database backends.
translate_sql(str_ends(var, "a"), con = con)
Error in `str_ends()`:
! Only fixed patterns are supported on database backends.
3.4.4 Redshift
con <- simulate_redshift()</pre>
translate_sql(nchar(var), con = con)
<SQL> LENGTH(`var`)
translate_sql(nzchar(var), con = con)
<SQL> (('var' IS NULL) OR 'var' != '')
translate_sql(substr(var, 1, 2), con = con)
<SQL> SUBSTRING(`var`, 1, 2)
translate_sql(trimws(var), con = con)
<SQL> LTRIM(RTRIM(`var`))
translate_sql(tolower(var), con = con)
```

```
<SQL> LOWER(`var`)
translate_sql(str_to_lower(var), con = con)
<SQL> LOWER(`var`)
translate_sql(toupper(var), con = con)
<SQL> UPPER(`var`)
translate_sql(str_to_upper(var), con = con)
<SQL> UPPER(`var`)
translate_sql(str_to_title(var), con = con)
<SQL> INITCAP(`var`)
translate_sql(str_trim(var), con = con)
<SQL> LTRIM(RTRIM(`var`))
translate_sql(str_squish(var), con = con)
<SQL> LTRIM(RTRIM(REGEXP_REPLACE(`var`, '\s+', ' ', 'g')))
translate_sql(str_detect(var, "b"), con = con)
<SQL> `var` ~ 'b'
translate_sql(str_detect(var, "b", negate = TRUE), con = con)
<SQL> !(`var` ~ 'b')
translate_sql(str_detect(var, "[aeiou]"), con = con)
<SQL> `var` ~ '[aeiou]'
```

```
translate_sql(str_replace(var, "a", "b"), con = con)
Error in `str_replace()`:
! `str_replace()` is not available in this SQL variant.
translate_sql(str_replace_all(var, "a", "b"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', 'b')
translate_sql(str_remove(var, "a"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', '')
translate_sql(str_remove_all(var, "a"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', '', 'g')
translate_sql(str_like(var, "a"), con = con)
<SQL> `var` ILIKE 'a'
translate_sql(str_starts(var, "a"), con = con)
Error in `str_starts()`:
! Only fixed patterns are supported on database backends.
translate_sql(str_ends(var, "a"), con = con)
Error in `str_ends()`:
! Only fixed patterns are supported on database backends.
3.4.5 Snowflake
con <- simulate_snowflake()</pre>
translate_sql(nchar(var), con = con)
<SQL> LENGTH(`var`)
translate_sql(nzchar(var), con = con)
```

```
<SQL> (('var' IS NULL) OR 'var' != '')
translate_sql(substr(var, 1, 2), con = con)
<SQL> SUBSTR(`var`, 1, 2)
translate_sql(trimws(var), con = con)
<SQL> LTRIM(RTRIM(`var`))
translate_sql(tolower(var), con = con)
<SQL> LOWER('var')
translate_sql(str_to_lower(var), con = con)
<SQL> LOWER(`var`)
translate_sql(toupper(var), con = con)
<SQL> UPPER(`var`)
translate_sql(str_to_upper(var), con = con)
<SQL> UPPER(`var`)
translate_sql(str_to_title(var), con = con)
<SQL> INITCAP(`var`)
translate_sql(str_trim(var), con = con)
<SQL> TRIM(`var`)
translate_sql(str_squish(var), con = con)
<SQL> REGEXP_REPLACE(TRIM(`var`), '\\s+', ' ')
```

```
translate_sql(str_detect(var, "b"), con = con)
<SQL> REGEXP INSTR(`var`, 'b') != 0
translate_sql(str_detect(var, "b", negate = TRUE), con = con)
<SQL> REGEXP_INSTR(`var`, 'b') = 0
translate_sql(str_detect(var, "[aeiou]"), con = con)
<SQL> REGEXP_INSTR(`var`, '[aeiou]') != 0
translate_sql(str_replace(var, "a", "b"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', 'b', 1.0, 1.0)
translate_sql(str_replace_all(var, "a", "b"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', 'b')
translate_sql(str_remove(var, "a"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a', '', 1.0, 1.0)
translate_sql(str_remove_all(var, "a"), con = con)
<SQL> REGEXP_REPLACE(`var`, 'a')
translate_sql(str_like(var, "a"), con = con)
<SQL> `var` LIKE 'a'
translate_sql(str_starts(var, "a"), con = con)
<SQL> REGEXP_INSTR(`var`, 'a') = 1
translate_sql(str_ends(var, "a"), con = con)
<SQL> REGEXP_INSTR(`var`, 'a', 1, 1, 1) = (LENGTH(`var`) + 1)
```

3.4.6 **Spark** con <- simulate_spark_sql()</pre> translate_sql(nchar(var), con = con) <SQL> LENGTH(`var`) translate_sql(nzchar(var), con = con) <SQL> (('var' IS NULL) OR 'var' != '') translate_sql(substr(var, 1, 2), con = con) <SQL> SUBSTR(`var`, 1, 2) translate_sql(trimws(var), con = con) <SQL> LTRIM(RTRIM(`var`)) translate_sql(tolower(var), con = con) <SQL> LOWER(`var`) translate_sql(str_to_lower(var), con = con) <SQL> LOWER('var') translate_sql(toupper(var), con = con) <SQL> UPPER(`var`) translate_sql(str_to_upper(var), con = con) <SQL> UPPER(`var`) translate_sql(str_to_title(var), con = con) <SQL> INITCAP(`var`)

```
translate_sql(str_trim(var), con = con)
<SQL> LTRIM(RTRIM(`var`))
translate_sql(str_squish(var), con = con)
Error in `str_squish()`:
! `str_squish()` is not available in this SQL variant.
translate_sql(str_detect(var, "b"), con = con)
Error in `str_detect()`:
! Only fixed patterns are supported on database backends.
translate_sql(str_detect(var, "b", negate = TRUE), con = con)
Error in `str_detect()`:
! Only fixed patterns are supported on database backends.
translate_sql(str_detect(var, "[aeiou]"), con = con)
Error in `str_detect()`:
! Only fixed patterns are supported on database backends.
translate_sql(str_replace(var, "a", "b"), con = con)
Error in `str_replace()`:
! `str_replace()` is not available in this SQL variant.
translate_sql(str_replace_all(var, "a", "b"), con = con)
Error in `str_replace_all() `:
! `str_replace_all()` is not available in this SQL variant.
translate_sql(str_remove(var, "a"), con = con)
Error in `str_remove()`:
! `str_remove()` is not available in this SQL variant.
```

```
translate_sql(str_remove_all(var, "a"), con = con)

Error in `str_remove_all()`:
! `str_remove_all()` is not available in this SQL variant.

translate_sql(str_like(var, "a"), con = con)

<SQL> `var` LIKE 'a'

translate_sql(str_starts(var, "a"), con = con)

Error in `str_starts()`:
! Only fixed patterns are supported on database backends.

translate_sql(str_ends(var, "a"), con = con)

Error in `str_ends()`:
! Only fixed patterns are supported on database backends.
```

3.5 Working with dates

Like with strings, support for working with dates is somewhat mixed. In general, we would use functions from the clock package such as get_day(), get_month(), get_year() to extract parts from a date, add_days() to add or subtract days to a date, and date_count_between() to get the number of days between two date variables.

```
Show SQL

3.5.1 DuckDB

con <- simulate_duckdb()
translate_sql(get_day(date_1), con = con)

<SQL> DATE_PART('day', `date_1`)

translate_sql(get_month(date_1), con = con)

<SQL> DATE_PART('month', `date_1`)
```

```
translate_sql(get_year(date_1), con = con)
<SQL> DATE_PART('year', `date_1`)
translate_sql(add_days(date_1, 1), con = con)
<SQL> DATE_ADD(`date_1`, INTERVAL (1.0) day)
translate_sql(add_years(date_1, 1), con = con)
<SQL> DATE_ADD(`date_1`, INTERVAL (1.0) year)
translate sql(date_count_between(date_1, date_2, "day"), con = con)
<SQL> DATEDIFF('day', `date_1`, `date_2`)
3.5.2 Postgres
con <- simulate_postgres()</pre>
translate_sql(get_day(date_1), con = con)
<SQL> DATE_PART('day', `date_1`)
translate_sql(get_month(date_1), con = con)
<SQL> DATE_PART('month', `date_1`)
translate_sql(get_year(date_1), con = con)
<SQL> DATE_PART('year', `date_1`)
translate_sql(add_days(date_1, 1), con = con)
<SQL> (`date_1` + 1.0*INTERVAL'1 day')
translate_sql(add_years(date_1, 1), con = con)
<SQL> (`date_1` + 1.0*INTERVAL'1 year')
translate_sql(date_count_between(date_1, date_2, "day"), con = con)
<SQL> `date_2` - `date_1`
```

```
3.5.3 SQL Server
con <- simulate_mssql()</pre>
translate_sql(get_day(date_1), con = con)
<SQL> DATEPART(DAY, `date_1`)
translate_sql(get_month(date_1), con = con)
<SQL> DATEPART(MONTH, `date_1`)
translate_sql(get_year(date_1), con = con)
<SQL> DATEPART(YEAR, `date_1`)
translate_sql(add_days(date_1, 1), con = con)
<SQL> DATEADD(DAY, 1.0, `date_1`)
translate_sql(add_years(date_1, 1), con = con)
<SQL> DATEADD(YEAR, 1.0, `date_1`)
translate sql(date_count_between(date_1, date_2, "day"), con = con)
<SQL> DATEDIFF(DAY, `date_1`, `date_2`)
3.5.4 Redshift
con <- simulate redshift()</pre>
translate_sql(get_day(date_1), con = con)
<SQL> DATE_PART('day', `date_1`)
translate_sql(get_month(date_1), con = con)
<SQL> DATE_PART('month', `date_1`)
translate_sql(get_year(date_1), con = con)
```

```
<SQL> DATE_PART('year', `date_1`)
translate_sql(add_days(date_1, 1), con = con)
<SQL> DATEADD(DAY, 1.0, `date_1`)
translate_sql(add_years(date_1, 1), con = con)
<SQL> DATEADD(YEAR, 1.0, `date_1`)
translate_sql(date_count_between(date_1, date_2, "day"), con = con)
<SQL> DATEDIFF(DAY, `date_1`, `date_2`)
3.5.5 Snowflake
con <- simulate_snowflake()</pre>
translate_sql(get_day(date_1), con = con)
<SQL> DATE_PART(DAY, `date_1`)
translate_sql(get_month(date_1), con = con)
<SQL> DATE_PART(MONTH, `date_1`)
translate_sql(get_year(date_1), con = con)
<SQL> DATE_PART(YEAR, `date_1`)
translate_sql(add_days(date_1, 1), con = con)
<SQL> DATEADD(DAY, 1.0, `date_1`)
translate_sql(add_years(date_1, 1), con = con)
<SQL> DATEADD(YEAR, 1.0, `date_1`)
translate_sql(date_count_between(date_1, date_2, "day"), con = con)
<SQL> DATEDIFF(DAY, `date_1`, `date_2`)
```

```
3.5.6 Spark
con <- simulate_spark_sql()</pre>
translate_sql(get_day(date_1), con = con)
<SQL> DATE_PART('DAY', `date_1`)
translate_sql(get_month(date_1), con = con)
<SQL> DATE_PART('MONTH', `date_1`)
translate_sql(get_year(date_1), con = con)
<SQL> DATE_PART('YEAR', `date_1`)
translate_sql(add_days(date_1, 1), con = con)
<SQL> DATE_ADD(`date_1`, 1.0)
translate_sql(add_years(date_1, 1), con = con)
<SQL> ADD_MONTHS(`date_1`, 1.0 * 12.0)
translate_sql(date_count_between(date_1, date_2, "day"), con = con)
<SQL> DATEDIFF(`date_2`, `date_1`)
```

3.6 Data aggregation

Within the context of using summarise(), we can get aggregated results across entire columns using functions such as n(), n_distinct(), sum(), min(), max(), mean(), and sd(). As can be seen below, the SQL for these calculations is similar across different database management systems.

```
• Show SQL
```

3.6.1 DuckDB

```
lazy_frame(x = c(1, 2), con = simulate_duckdb()) |>
  summarise(
    n = n(),
   n_unique = n_distinct(x),
    sum = sum(x, na.rm = TRUE),
   sum_is_1 = sum(x == 1, na.rm = TRUE),
   min = min(x, na.rm = TRUE),
   mean = mean(x, na.rm = TRUE),
   max = max(x, na.rm = TRUE),
   sd = sd(x, na.rm = TRUE)
  ) |>
  show_query()
<SQL>
SELECT
  COUNT(*) AS `n`,
  COUNT(DISTINCT row(`x`)) AS `n_unique`,
  SUM('x') AS 'sum',
  SUM(x = 1.0) AS sum_is_1,
  MIN('x') AS 'min',
  AVG(`x`) AS `mean`,
  MAX('x') AS 'max',
  STDDEV(`x`) AS `sd`
FROM `df`
3.6.2 Postgres
```

```
lazy_frame(x = c(1, 2), con = simulate_postgres()) |>
  summarise(
    n = n(),
    n unique = n_distinct(x),
    sum = sum(x, na.rm = TRUE),
    sum_is_1 = sum(x == 1, na.rm = TRUE),
    min = min(x, na.rm = TRUE),
   mean = mean(x, na.rm = TRUE),
   max = max(x, na.rm = TRUE),
    sd = sd(x, na.rm = TRUE)
  ) |>
  show_query()
<SQL>
SELECT
  COUNT(*) AS `n`,
  COUNT(DISTINCT `x`) AS `n_unique`,
  SUM(`x`) AS `sum`,
  SUM(x) = 1.0 AS sum_is_1,
  MIN('x') AS 'min',
  AVG('x') AS 'mean',
  MAX('x') AS 'max',
  STDDEV_SAMP(`x`) AS `sd`
FROM `df`
3.6.3 SQL Server
lazy_frame(x = c(1, 2), con = simulate_mssql()) |>
  summarise(
    n = n(),
    n_unique = n_distinct(x),
    sum = sum(x, na.rm = TRUE),
    sum_is_1 = sum(x == 1, na.rm = TRUE),
    min = min(x, na.rm = TRUE),
   mean = mean(x, na.rm = TRUE),
   max = max(x, na.rm = TRUE),
    sd = sd(x, na.rm = TRUE)
  ) |>
  show_query()
<SQL>
```

```
SELECT
  COUNT_BIG(*) AS `n`,
  COUNT(DISTINCT `x`) AS `n_unique`,
  SUM(`x`) AS `sum`,
  SUM(CAST(IIF(`x` = 1.0, 1, 0) AS BIT)) AS `sum_is_1`,
  MIN('x') AS 'min',
  AVG('x') AS 'mean',
  MAX('x') AS 'max',
  STDEV('x') AS 'sd'
FROM `df`
3.6.4 Redshift
lazy_frame(x = c(1, 2), con = simulate_redshift()) |>
 summarise(
   n = n(),
    n_unique = n_distinct(x),
   sum = sum(x, na.rm = TRUE),
    sum_is_1 = sum(x == 1, na.rm = TRUE),
   min = min(x, na.rm = TRUE),
   mean = mean(x, na.rm = TRUE),
   max = max(x, na.rm = TRUE),
    sd = sd(x, na.rm = TRUE)
  ) |>
  show_query()
<SQL>
SELECT
  COUNT(*) AS `n`,
  COUNT(DISTINCT `x`) AS `n_unique`,
  SUM('x') AS 'sum',
  SUM(x) = 1.0) AS sum_is_1,
  MIN('x') AS 'min',
 AVG('x') AS 'mean',
  MAX(`x`) AS `max`,
  STDDEV_SAMP(`x`) AS `sd`
FROM `df`
```

3.6.5 Snowflake

```
lazy_frame(x = c(1, 2), con = simulate_snowflake()) |>
  summarise(
    n = n(),
    n unique = n_distinct(x),
    sum = sum(x, na.rm = TRUE),
    sum_is_1 = sum(x == 1, na.rm = TRUE),
    min = min(x, na.rm = TRUE),
   mean = mean(x, na.rm = TRUE),
   max = max(x, na.rm = TRUE),
    sd = sd(x, na.rm = TRUE)
  ) |>
  show_query()
<SQL>
SELECT
  COUNT(*) AS `n`,
  COUNT(DISTINCT `x`) AS `n_unique`,
  SUM(`x`) AS `sum`,
  SUM(x) = 1.0 AS sum_is_1,
  MIN('x') AS 'min',
  AVG('x') AS 'mean',
  MAX('x') AS 'max',
  STDDEV(`x`) AS `sd`
FROM `df`
3.6.6 Spark
lazy_frame(x = c(1, 2), con = simulate_spark_sql()) >
  summarise(
    n = n(),
   n_unique = n_distinct(x),
    sum = sum(x, na.rm = TRUE),
    sum_is_1 = sum(x == 1, na.rm = TRUE),
    min = min(x, na.rm = TRUE),
   mean = mean(x, na.rm = TRUE),
   max = max(x, na.rm = TRUE),
    sd = sd(x, na.rm = TRUE)
  ) |>
  show_query()
<SQL>
```

```
SELECT

COUNT(*) AS `n`,

COUNT(DISTINCT `x`) AS `n_unique`,

SUM(`x`) AS `sum`,

SUM(`x` = 1.0) AS `sum_is_1`,

MIN(`x`) AS `min`,

AVG(`x`) AS `mean`,

MAX(`x`) AS `max`,

STDDEV_SAMP(`x`) AS `sd`

FROM `df`
```

3.7 Window functions

Window functions differ from data aggregation functions in that they perform calculations across rows that are related to the current row, rather than collapsing multiple rows into a single result. For these operations, we use mutate() instead of summarise().

For instance, we can use window functions like cumsum() and cummean() to calculate running totals and averages, or lag() and lead() to help compare rows to their preceding or following rows.

Given that window functions compare rows to rows before or after them, we will often use arrange() or window_order() to specify the order of rows. This will translate into an ORDER BY clause in the SQL. In addition, we may well also want to apply window functions within some specific groupings in our data. Using group_by() would result in a PARTITION BY clause in the translated SQL so that the window function operates on each group independently.



```
con <- simulate_duckdb()</pre>
lazy_frame(x = c(10, 20, 30), z = c(1, 2, 3), con = con) |>
  window_order(z) |>
  mutate(
    sum_x = cumsum(x),
   mean_x = cummean(x),
   lag_x = lag(x),
   lead_x = lead(x)
  ) |>
  show_query()
<SQL>
SELECT
  `df`.*,
  SUM('x') OVER (ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'sum x',
  AVG('x') OVER (ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'mean_x',
  LAG('x', 1, NULL) OVER (ORDER BY 'z') AS 'lag_x',
  LEAD('x', 1, NULL) OVER (ORDER BY 'z') AS 'lead_x'
FROM `df`
lazy_frame(x = c(10, 20), y = c("a", "b"), z = c(1, 2), con = con) |>
  window_order(z) |>
  group_by(y) |>
  mutate(
   sum_x = cumsum(x),
    mean_x = cummean(x),
   lag_x = lag(x),
   lead_x = lead(x)
  ) |>
  show_query()
<SQL>
SELECT
  `df`.*,
  SUM('x') OVER (PARTITION BY 'y' ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'sum_x',
  AVG('x') OVER (PARTITION BY 'y' ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS
                                                                              `mean_x`,
  LAG('x', 1, NULL) OVER (PARTITION BY 'y' ORDER BY 'z') AS 'lag_x',
  LEAD('x', 1, NULL) OVER (PARTITION BY 'y' ORDER BY 'z') AS 'lead_x'
FROM `df`
```

3.7.2 Postgres

```
con <- simulate_postgres()</pre>
lazy_frame(x = c(10, 20, 30), z = c(1, 2, 3), con = con) |>
  window_order(z) |>
  mutate(
   sum_x = cumsum(x),
   mean_x = cummean(x),
   lag_x = lag(x),
   lead_x = lead(x)
  ) |>
  show_query()
<SQL>
SELECT
  `df`.*,
  SUM(`x`) OVER `win1` AS `sum_x`,
  AVG('x') OVER 'win1' AS 'mean_x',
  LAG('x', 1, NULL) OVER 'win2' AS 'lag_x',
  LEAD(`x`, 1, NULL) OVER `win2` AS `lead_x`
FROM `df`
WINDOW
  `win1` AS (ORDER BY `z` ROWS UNBOUNDED PRECEDING),
  `win2` AS (ORDER BY `z`)
lazy_frame(x = c(10, 20), y = c("a", "b"), z = c(1, 2), con = con) >
  window_order(z) |>
  group_by(y) |>
  mutate(
    sum_x = cumsum(x),
   mean_x = cummean(x),
   lag_x = lag(x),
   lead_x = lead(x)
  ) |>
  show_query()
<SQL>
SELECT
  `df`.*,
  SUM(`x`) OVER `win1` AS `sum_x`,
  AVG(`x`) OVER `win1` AS `mean_x`,
```

```
LAG('x', 1, NULL) OVER 'win2' AS 'lag_x',
  LEAD(`x`, 1, NULL) OVER `win2` AS `lead_x`
FROM `df`
WINDOW
  `win1` AS (PARTITION BY `y` ORDER BY `z` ROWS UNBOUNDED PRECEDING),
  `win2` AS (PARTITION BY `y` ORDER BY `z`)
3.7.3 SQL Server
con <- simulate_mssql()</pre>
lazy_frame(x = c(10, 20, 30), z = c(1, 2, 3), con = con) |>
  window_order(z) |>
  mutate(
    sum_x = cumsum(x),
   mean x = cummean(x),
   lag_x = lag(x),
   lead_x = lead(x)
  ) |>
  show_query()
<SQL>
SELECT
  `df`.*,
  SUM('x') OVER (ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'sum_x',
  AVG('x') OVER (ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'mean x',
  LAG(`x`, 1, NULL) OVER (ORDER BY `z`) AS `lag_x`,
  LEAD('x', 1, NULL) OVER (ORDER BY 'z') AS 'lead_x'
FROM `df`
lazy_frame(x = c(10, 20), y = c("a", "b"), z = c(1, 2), con = con) >
  window_order(z) |>
  group_by(y) |>
  mutate(
    sum_x = cumsum(x),
   mean_x = cummean(x),
   lag_x = lag(x),
   lead_x = lead(x)
  ) |>
  show_query()
<SQL>
```

```
SELECT
  `df`.*,
  SUM('x') OVER (PARTITION BY 'y' ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'sum_x',
  AVG('x') OVER (PARTITION BY 'y' ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'mean_x',
  LAG('x', 1, NULL) OVER (PARTITION BY 'y' ORDER BY 'z') AS 'lag_x',
  LEAD('x', 1, NULL) OVER (PARTITION BY 'y' ORDER BY 'z') AS 'lead_x'
FROM `df`
3.7.4 Redshift
con <- simulate_redshift()</pre>
lazy_frame(x = c(10, 20, 30), z = c(1, 2, 3), con = con) >
  window_order(z) |>
  mutate(
    sum_x = cumsum(x),
    mean x = cummean(x),
   lag x = lag(x),
   lead_x = lead(x)
  ) |>
  show_query()
<SQL>
SELECT
  `df`.*,
  SUM('x') OVER (ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'sum x',
  AVG('x') OVER (ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'mean_x',
  LAG('x', 1) OVER (ORDER BY 'z') AS 'lag_x',
  LEAD(`x`, 1) OVER (ORDER BY `z`) AS `lead_x`
FROM `df`
lazy_frame(x = c(10, 20), y = c("a", "b"), z = c(1, 2), con = con) >
  window_order(z) |>
  group_by(y) |>
  mutate(
    sum x = cumsum(x),
   mean x = cummean(x),
   lag_x = lag(x),
   lead_x = lead(x)
  ) |>
  show_query()
```

```
<SQL>
SELECT
  `df`.*,
  SUM('x') OVER (PARTITION BY 'y' ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'sum_x',
  AVG('x') OVER (PARTITION BY 'y' ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'mean_x',
  LAG('x', 1) OVER (PARTITION BY 'y' ORDER BY 'z') AS 'lag_x',
  LEAD(`x`, 1) OVER (PARTITION BY `y` ORDER BY `z`) AS `lead_x`
FROM `df`
3.7.5 Snowflake
con <- simulate snowflake()</pre>
lazy_frame(x = c(10, 20, 30), z = c(1, 2, 3), con = con) |>
  window_order(z) |>
  mutate(
    sum_x = cumsum(x),
   mean_x = cummean(x),
   lag_x = lag(x),
   lead_x = lead(x)
  ) |>
  show_query()
<SQL>
SELECT
  `df`.*,
  SUM('x') OVER (ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'sum_x',
  AVG('x') OVER (ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'mean x',
  LAG('x', 1, NULL) OVER (ORDER BY 'z') AS 'lag_x',
  LEAD('x', 1, NULL) OVER (ORDER BY 'z') AS 'lead x'
FROM `df`
lazy_frame(x = c(10, 20), y = c("a", "b"), z = c(1, 2), con = con) |>
  window_order(z) |>
  group_by(y) |>
  mutate(
   sum_x = cumsum(x),
    mean x = cummean(x),
   lag_x = lag(x),
   lead_x = lead(x)
  ) |>
  show_query()
```

```
<SQL>
SELECT
  `df`.*,
  SUM('x') OVER (PARTITION BY 'y' ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'sum_x',
  AVG('x') OVER (PARTITION BY 'y' ORDER BY 'z' ROWS UNBOUNDED PRECEDING) AS 'mean_x',
 LAG('x', 1, NULL) OVER (PARTITION BY 'y' ORDER BY 'z') AS 'lag_x',
  LEAD('x', 1, NULL) OVER (PARTITION BY 'y' ORDER BY 'z') AS 'lead x'
FROM `df`
3.7.6 Spark
con <- simulate_spark_sql()</pre>
lazy_frame(x = c(10, 20, 30), z = c(1, 2, 3), con = con) |>
  window_order(z) |>
 mutate(
   sum x = cumsum(x),
   mean x = cummean(x),
   lag_x = lag(x),
   lead_x = lead(x)
  ) |>
 show_query()
<SQL>
SELECT
  `df`.*,
  SUM(`x`) OVER `win1` AS `sum_x`,
  AVG(`x`) OVER `win1` AS `mean_x`,
 LAG('x', 1, NULL) OVER 'win2' AS 'lag_x',
  LEAD(`x`, 1, NULL) OVER `win2` AS `lead_x`
FROM `df`
WINDOW
  `win1` AS (ORDER BY `z` ROWS UNBOUNDED PRECEDING),
  `win2` AS (ORDER BY `z`)
```

```
lazy_frame(x = c(10, 20), y = c("a", "b"), z = c(1, 2), con = con) |>
  window_order(z) |>
  group_by(y) |>
  mutate(
    sum_x = cumsum(x),
    mean_x = cummean(x),
    lag_x = lag(x),
    lead_x = lead(x)
  ) |>
  show_query()
<SQL>
SELECT
  `df`.*,
  SUM('x') OVER 'win1' AS 'sum x',
  AVG(`x`) OVER `win1` AS `mean_x`,
  LAG('x', 1, NULL) OVER 'win2' AS 'lag_x',
  LEAD('x', 1, NULL) OVER 'win2' AS 'lead_x'
FROM `df`
WINDOW
  `win1` AS (PARTITION BY `y` ORDER BY `z` ROWS UNBOUNDED PRECEDING),
  `win2` AS (PARTITION BY `y` ORDER BY `z`)
```

TODO add note arrange vs window_order

i arrange() vs window_order()

Although arrange() and window_order() both involve ordering, they serve different purposes.

- arrange(): changes the order of the final output of a table, by ordering it as the user specified.
- window_order(): defines the order within window functions. It controlls how functions (e.g., lag(), lead(), rank(), cumsum()) are applied across rows.

In conclusion, for all dialects, arrange() changes the output row order, while window_order() changes how window functions calculate values.

3.8 Calculating quantiles, including the median

So far we've seen that we can perform various data manipulations and calculate summary statistics for different database management systems using the same R code. Although the translated SQL has been different, the databases all supported similar approaches to perform these queries.

A case where this is not true is when we are interested in summarising distributions of the data and estimating quantiles. For example, let's take estimating the median as an example. Some databases only support calculating the median as an aggregation function similar to how min, mean, and max were calculated above. However, some others only support it as a window function like lead and lag above. Unfortunately, this means that for some databases, quantiles can only be calculated using the summarise aggregation approach, while in others only the mutate window approach can be used.

```
Show SQL
3.8.1 DuckDB
con <- simulate_duckdb()</pre>
lazy_frame(x = c(1,2), con = con) >
  summarise(median = median(x, na.rm = TRUE)) |>
  show_query()
<SQL>
SELECT MEDIAN(`x`) AS `median`
FROM `df`
lazy_frame(x = c(1,2), con = con) >
  mutate(median = median(x, na.rm = TRUE)) |>
  show_query()
<SQL>
SELECT `df`.*, MEDIAN(`x`) OVER () AS `median`
FROM `df`
3.8.2 Postgres
con <- simulate_postgres()</pre>
lazy_frame(x = c(1,2), con = con) >
  summarise(median = median(x, na.rm = TRUE)) |>
  show query()
```

```
<SQL>
SELECT PERCENTILE_CONT(0.5) WITHIN GROUP (ORDER BY `x`) AS `median`
FROM `df`
lazy_frame(x = c(1,2), con = con) >
  mutate(median = median(x, na.rm = TRUE)) |>
  show_query()
Error in `median()`:
! Translation of `median()` in `mutate()` is not supported for
  PostgreSQL.
i Use a combination of `summarise()` and `left_join()` instead:
  `df %>% left_join(summarise(<col> = median(x, na.rm = TRUE)))`.
3.8.3 SQL Server
con <- simulate_mssql()</pre>
lazy_frame(x = c(1,2), con = con) >
  summarise(median = median(x, na.rm = TRUE)) |>
  show_query()
Error in `median()`:
! Translation of `median()` in `summarise()` is not supported for SQL
  Server.
i Use a combination of `distinct()` and `mutate()` for the same result:
  `mutate(<col> = median(x, na.rm = TRUE)) %>% distinct(<col>)`
lazy_frame(x = c(1,2), con = con) >
  mutate(median = median(x, na.rm = TRUE)) |>
  show_query()
<SQL>
SELECT
  `df`.*,
  PERCENTILE CONT(0.5) WITHIN GROUP (ORDER BY 'x') OVER () AS 'median'
FROM `df`
3.8.4 Redshift
con <- simulate redshift()</pre>
lazy_frame(x = c(1,2), con = con) >
  summarise(median = median(x, na.rm = TRUE)) |>
  show_query()
```

```
<SQL>
SELECT PERCENTILE_CONT(0.5) WITHIN GROUP (ORDER BY `x`) AS `median`
FROM `df`
lazy_frame(x = c(1,2), con = con) |>
  mutate(median = median(x, na.rm = TRUE)) |>
  show_query()
Error in `median()`:
! Translation of `median()` in `mutate()` is not supported for
  PostgreSQL.
i Use a combination of `summarise()` and `left_join()` instead:
  `df %>% left_join(summarise(<col> = median(x, na.rm = TRUE)))`.
3.8.5 Snowflake
con <- simulate_snowflake()</pre>
lazy_frame(x = c(1,2), con = con) >
  summarise(median = median(x, na.rm = TRUE)) |>
  show_query()
<SQL>
SELECT PERCENTILE_CONT(0.5) WITHIN GROUP (ORDER BY `x`) AS `median`
FROM `df`
lazy_frame(x = c(1,2), con = con) >
  mutate(median = median(x, na.rm = TRUE)) |>
  show_query()
<SQL>
SELECT
  `df`.*,
  PERCENTILE_CONT(0.5) WITHIN GROUP (ORDER BY `x`) OVER () AS `median`
FROM `df`
3.8.6 Spark
con <- simulate_spark_sql()</pre>
lazy_frame(x = c(1,2), con = con) >
  summarise(median = median(x, na.rm = TRUE)) |>
  show_query()
```

```
<SQL>
SELECT MEDIAN(`x`) AS `median`
FROM `df`

lazy_frame(x = c(1,2), con = con) |>
    mutate(median = median(x, na.rm = TRUE)) |>
    show_query()

<SQL>
SELECT `df`.*, MEDIAN(`x`) OVER () AS `median`
FROM `df`
```

4 Building analytic pipelines for a data model

In the previous chapters, we've seen that after connecting to a database, we can create references to the various tables we're interested in and write custom analytic code to query them. However, if we are working with the same database over and over again, we might want to build some tooling for tasks we often perform.

To see how we can develop a data model with associated methods and functions, we will use the Lahman baseball data. The data is stored across various related tables.

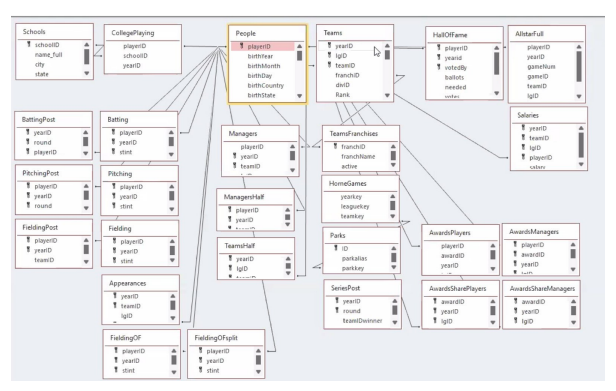


Figure 4.1: Lahman's Baseball Dabatase schema from https://cdalzell.github.io/Lahman/.

4.1 Defining a data model

```
library(duckdb)
library(dplyr)
library(tidyr)
library(purrr)
library(cli)
library(dbplyr)
library(Lahman)

con <- dbConnect(drv = duckdb())
copy_lahman(con = con)</pre>
```

i copy_lahman

The copy_lahman() function inserts all the different tables in the connection. It works in the same way as we have done before with the for loop and the dbWriteTable() function. See that there are 28 new tables inserted in our DuckDB database:

```
dbListTables(conn = con)
 [1] "AllstarFull"
                            "Appearances"
                                                   "AwardsManagers"
 [4] "AwardsPlayers"
                            "AwardsShareManagers" "AwardsSharePlayers"
 [7] "Batting"
                            "BattingPost"
                                                   "CollegePlaying"
[10] "Fielding"
                            "FieldingOF"
                                                   "FieldingOFsplit"
[13] "FieldingPost"
                            "HallOfFame"
                                                   "HomeGames"
[16] "LahmanData"
                            "Managers"
                                                   "ManagersHalf"
[19] "Parks"
                            "People"
                                                   "Pitching"
[22] "PitchingPost"
                            "Salaries"
                                                   "Schools"
                            "Teams"
                                                   "TeamsFranchises"
[25] "SeriesPost"
[28] "TeamsHalf"
```

Instead of manually creating references for each one of the tables (so we can access them easily), we will write a function to create a single reference to the Lahman data.

```
lahmanFromCon <- function(con) {
  lahmanRef <- set_names(c(
    "AllstarFull", "Appearances", "AwardsManagers", "AwardsPlayers", "AwardsManagers",
    "AwardsShareManagers", "Batting", "BattingPost", "CollegePlaying", "Fielding",
    "FieldingOF", "FieldingOFsplit", "FieldingPost", "HallOfFame", "HomeGames",
    "LahmanData", "Managers", "ManagersHalf", "Parks", "People", "Pitching",</pre>
```

```
"PitchingPost", "Salaries", "Schools", "SeriesPost", "Teams", "TeamsFranchises",
    "TeamsHalf"
))

lahmanRef <- map(lahmanRef, \(x) tbl(src = con, from = x))

class(lahmanRef) <- c("lahman_ref", class(lahmanRef))
    return(lahmanRef)
}</pre>
```

With this function we can now easily get references to all our Lahman tables in one go using our lahmanFromCon() function.

```
lahman <- lahmanFromCon(con = con)
lahman$People |>
  glimpse()
```

```
Rows: ??
Columns: 26
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
                                  <chr> "aardsda01", "aaronha01", "aaronto01", "aasedo01", "abada~
$ playerID
                                  <int> 1981, 1934, 1939, 1954, 1972, 1985, 1850, 1877, 1869, 186~
$ birthYear
$ birthMonth
                                  <int> 12, 2, 8, 9, 8, 12, 11, 4, 11, 10, 6, 9, 3, 10, 2, 8, 9, ~
$ birthDay
                                  <int> 27, 5, 5, 8, 25, 17, 4, 15, 11, 14, 1, 20, 16, 22, 16, 17~
$ birthCity
                                  <chr> "Denver", "Mobile", "Mobile", "Orange", "Palm Beach", "La~
$ birthCountry <chr> "USA", "USA", "USA", "USA", "USA", "D.R.", "USA", "USA", ~
$ birthState
                                  <chr> "CO", "AL", "AL", "CA", "FL", "La Romana", "PA", "PA", "V~
                                  <int> NA, 2021, 1984, NA, NA, NA, 1905, 1957, 1962, 1926, NA, N~
$ deathYear
$ deathMonth
                                 <int> NA, 1, 8, NA, NA, NA, 5, 1, 6, 4, NA, NA, 2, 6, NA, NA, N~
                                  <int> NA, 22, 16, NA, NA, NA, 17, 6, 11, 27, NA, NA, 13, 11, NA~
$ deathDay
$ deathCountry <chr> NA, "USA", "USA", NA, NA, NA, "USA", 
                                  <chr> NA, "GA", "GA", NA, NA, NA, "NJ", "FL", "VT", "CA", NA, N~
$ deathState
$ deathCity
                                  <chr> NA, "Atlanta", "Atlanta", NA, NA, NA, "Pemberton", "Fort ~
                                  <chr> "David", "Hank", "Tommie", "Don", "Andy", "Fernando", "Jo~
$ nameFirst
                                 <chr> "Aardsma", "Aaron", "Aaron", "Aase", "Abad", "Abad", "Aba~
$ nameLast
                                 <chr> "David Allan", "Henry Louis", "Tommie Lee", "Donald Willi~
$ nameGiven
                                  <int> 215, 180, 190, 190, 184, 235, 192, 170, 175, 169, 192, 22~
$ weight
$ height
                                  <int> 75, 72, 75, 75, 73, 74, 72, 71, 71, 68, 72, 74, 71, 70, 7~
$ bats
                                  <fct> R, R, R, R, L, L, R, R, R, L, L, R, R, R, R, R, R, L, R, L, ~
$ throws
                                  <fct> R, R, R, R, L, L, R, R, R, L, L, R, R, R, R, R, L, L, R, L, ~
```

i The dm package

In this chapter we will be creating a bespoke data model for our database. This approach can be further extended using the $\frac{dm}{dm}$ package, which also provides various helpful functions for creating a data model and working with it.

Similar to above, we can use dm() to create a single object to access our database tables.

```
library(dm)
lahman_dm <- dm(batting = tbl(con, "Batting"), people = tbl(con, "People"))</pre>
lahman_dm
-- Table source ------
src: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
-- Metadata -----
Tables: `batting`, `people`
Columns: 48
Primary keys: 0
Foreign keys: 0
Using this approach, we can make use of various utility functions. For example here we
specify primary and foreign keys and then check that the key constraints are satisfied.
lahman_dm <- lahman_dm |>
 dm_add_pk(table = "people", columns = "playerID") |>
 dm_add_fk(table = "batting", columns = "playerID", ref_table = "people")
lahman_dm
src: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
-- Metadata -----
Tables: `batting`, `people`
```

```
Columns: 48
Primary keys: 1
Foreign keys: 1

dm_examine_constraints(.dm = lahman_dm)

i All constraints satisfied.

For more information on the dm package see https://dm.cynkra.com/index.html
```

4.2 Creating functions for the data model

Given that we know the structure of the data, we can build a set of functions tailored to the Lahman data model to simplify data analyses.

Let's start by creating a simple function that returns the teams each player has played for. We can see that the code we use follows on from the last couple of chapters.

```
getTeams <- function(lahman, name = "Barry Bonds") {
   lahman$Batting |>
    inner_join(
        lahman$People |>
        mutate(full_name = pasteO(nameFirst, " ", nameLast)) |>
        filter(full_name %in% name) |>
        select("playerID"),
        by = "playerID"
        ) |>
        distinct(teamID, yearID) |>
        left_join(
        lahman$Teams,
        by = c("teamID", "yearID")) |>
        distinct(name)
}
```

Now we can easily get which teams a player has represented. We can see how changing the player name changes the SQL that is run behind the scenes.

```
getTeams(lahman = lahman, name = "Babe Ruth")
```

```
i Show query
<SQL>
SELECT DISTINCT q01.*
FROM (
  SELECT "name"
  FROM (
    SELECT DISTINCT q01.*
    FROM (
      SELECT teamID, yearID
      FROM Batting
      INNER JOIN (
        SELECT playerID
        FROM (
          SELECT People.*, CONCAT_WS('', nameFirst, '', nameLast) AS full_name
         FROM People
        ) q01
        WHERE (full_name IN ('Babe Ruth'))
        ON (Batting.playerID = RHS.playerID)
    ) q01
  ) LHS
  LEFT JOIN Teams
    ON (LHS.teamID = Teams.teamID AND LHS.yearID = Teams.yearID)
) q01
```

```
getTeams(lahman = lahman, name = "Barry Bonds")
```

```
i Show query
<SQL>
SELECT DISTINCT q01.*
FROM (
  SELECT "name"
  FROM (
    SELECT DISTINCT q01.*
    FROM (
      SELECT teamID, yearID
      FROM Batting
      INNER JOIN (
        SELECT playerID
        FROM (
          SELECT People.*, CONCAT WS('', nameFirst, '', nameLast) AS full name
          FROM People
        ) q01
        WHERE (full_name IN ('Barry Bonds'))
        ON (Batting.playerID = RHS.playerID)
    ) q01
  ) LHS
  LEFT JOIN Teams
    ON (LHS.teamID = Teams.teamID AND LHS.yearID = Teams.yearID)
) q01
```

? Choosing the right time to collect data into R

The function <code>collect()</code> brings data out of the database and into R. When working with large datasets, as is often the case when interacting with a database, we typically want to keep as much computation as possible on the database side. In the case of our <code>getTeams()</code> function, for example, everything is done on the database side. Collecting the result will bring the result of the teams the person played for out of the database. In this case, we could also use <code>pull()</code> to get our result out as a vector rather than a data frame.

```
getTeams(lahman = lahman, name = "Barry Bonds") |>
  collect()

# A tibble: 2 x 1
  name
  <chr>
```

```
1 Pittsburgh Pirates
2 San Francisco Giants

getTeams(lahman = lahman, name = "Barry Bonds") |>
   pull()

[1] "San Francisco Giants" "Pittsburgh Pirates"
```

However, in other cases we may need to collect the data to perform analyses that can not be done in SQL. This might be the case for plotting or for other analytic steps(i.e., fitting statistical models). In such cases, it is important to only bring out the data that we need (as our local computer will typically have far less memory available than the database system).

Similarly, we can make a function to add a player's year of birth to another Lahman table.

```
addBirthCountry <- function(x){
    x |>
    left_join(
        lahman$People |>
            select("playerID", "birthCountry"),
        by = "playerID"
    )
}
```

```
lahman$Batting |>
  addBirthCountry()
```

```
# Source:
             SQL [?? x 23]
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
                                                                         X2B
              yearID stint teamID lgID
                                                G
                                                      AB
                                                             R
                                                                    Η
                                                                               ХЗВ
                                                                                       HR
   <chr>
               <int> <int> <fct>
                                     <fct> <int> <int> <int> <int> <int> <int> <int> <int> <int>
                           1 SFN
 1 aardsda01
                2004
                                               11
                                                       0
 2 aardsda01
                2006
                           1 CHN
                                     NL
                                               45
                                                       2
                                                             0
                                                                    0
                                                                           0
                                                                                  0
                                                                                        0
 3 aardsda01
                2007
                          1 CHA
                                     ΑL
                                               25
                                                       0
                                                             0
                                                                    0
                                                                           0
                                                                                  0
                                                                                        0
4 aardsda01
                2008
                          1 BOS
                                               47
                                                       1
                                                             0
                                                                    0
                                                                           0
                                                                                  0
                                                                                        0
                                     AL
5 aardsda01
                2009
                          1 SEA
                                     ΑL
                                               73
                                                       0
                                                             0
                                                                    0
                                                                           0
                                                                                  0
                                                                                        0
 6 aardsda01
                2010
                                               53
                                                       0
                                                             0
                                                                    0
                                                                           0
                                                                                  0
                                                                                        0
                          1 SEA
                                     AL
7 aardsda01
                2012
                           1 NYA
                                     ΑL
                                                1
                                                       0
                                                             0
                                                                    0
                                                                           0
                                                                                  0
                                                                                        0
8 aardsda01
                2013
                           1 NYN
                                     NL
                                               43
                                                       0
                                                             0
                                                                    0
                                                                           0
                                                                                  0
                                                                                        0
9 aardsda01
                2015
                          1 ATL
                                     NL
                                               33
                                                             0
                                                                    0
                                                                           0
                                                                                  0
                                                                                        0
```

```
10 aaronha01
               1954
                       1 ML1
                                NL
                                        122
                                              468
                                                     58
                                                          131
                                                                 27
                                                                             13
# i more rows
# i 11 more variables: RBI <int>, SB <int>, CS <int>, BB <int>, SO <int>,
    IBB <int>, HBP <int>, SH <int>, SF <int>, GIDP <int>, birthCountry <chr>
 i Show query
 <SQL>
 SELECT Batting.*, birthCountry
 FROM Batting
 LEFT JOIN People
   ON (Batting.playerID = People.playerID)
lahman$Pitching |>
  addBirthCountry()
# Source:
           SQL [?? x 31]
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
   playerID yearID stint teamID lgID
                                                L
                                                      G
                                                           GS
                                                                  CG
                                                                              SV
                                          W
                                                                       SHO
              <chr>
                                                                        0
 1 aardsda01
               2004
                        1 SFN
                                NL
                                          1
                                                0
                                                      11
                                                             0
                                                                   0
                                                                              0
                        1 CHN
                                          3
 2 aardsda01
              2006
                                NL
                                                      45
                                                                         0
                                                                               0
3 aardsda01
              2007
                       1 CHA
                                AL
                                           2
                                                1
                                                      25
                                                                         0
                                                                              0
4 aardsda01
              2008
                       1 BOS
                                          4
                                                2
                                                     47
                                                                         0
                                                                              0
                                AL
                                                             0
                                                                   0
5 aardsda01
                                          3
              2009
                       1 SEA
                                AL
                                                6
                                                     73
                                                             0
                                                                   0
                                                                         0
                                                                              38
6 aardsda01
              2010
                       1 SEA
                                AL
                                          0
                                                6
                                                     53
                                                             0
                                                                   0
                                                                        0
                                                                              31
7 aardsda01
              2012
                       1 NYA
                                AL
                                          0
                                                0
                                                      1
                                                             0
                                                                   0
                                                                         0
                                                                              0
8 aardsda01
                       1 NYN
                                NL
                                          2
                                                2
                                                     43
                                                             0
                                                                   0
                                                                         0
                                                                              0
              2013
9 aardsda01
              2015
                       1 ATL
                                NL
                                           1
                                                 1
                                                      33
                                                             0
                                                                   0
                                                                         0
                                                                              0
                                                                         2
                                                2
                                                                              0
10 aasedo01
               1977
                        1 BOS
                                AL
                                           6
                                                      13
                                                            13
# i more rows
# i 19 more variables: IPouts <int>, H <int>, ER <int>, HR <int>, BB <int>,
   SO <int>, BAOpp <dbl>, ERA <dbl>, IBB <int>, WP <int>, HBP <int>, BK <int>,
#
   BFP <int>, GF <int>, R <int>, SH <int>, SF <int>, GIDP <int>,
   birthCountry <chr>
 i Show query
```

```
i Show query

<SQL>
SELECT Pitching.*, birthCountry
```

```
FROM Pitching
LEFT JOIN People
ON (Pitching.playerID = People.playerID)
```

We can then use our addBirthCountry() function as part of a larger query to summarise and plot the proportion of players from each country over time (based on their presence in the batting table).

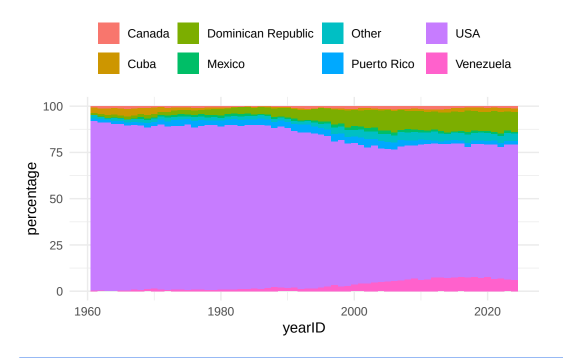
```
plot_data <- lahman$Batting |>
  select("playerID", "yearID") |>
  addBirthCountry() |>
  filter(yearID > 1960) |>
  mutate(birthCountry = case_when(
    birthCountry == "USA" ~ "USA",
    birthCountry == "D.R." ~ "Dominican Republic",
    birthCountry == "Venezuela" ~ "Venezuela",
    birthCountry == "P.R." ~ "Puerto Rico ",
    birthCountry == "Cuba" ~ "Cuba",
    birthCountry == "CAN" ~ "Canada",
    birthCountry == "Mexico" ~ "Mexico",
    .default = "Other"
  )) |>
  group_by(yearID, birthCountry) |>
  summarise(n = n(), .groups = "drop") |>
  group_by(yearID) |>
  mutate(percentage = n / sum(n) * 100) |>
  ungroup() |>
  collect()
```

```
i Show query

<SQL>
SELECT q01.*, (n / SUM(n) OVER (PARTITION BY yearID)) * 100.0 AS percentage
FROM (
    SELECT yearID, birthCountry, COUNT(*) AS n
    FROM (
    SELECT
        playerID,
        yearID,
        CASE
WHEN (birthCountry = 'USA') THEN 'USA'
WHEN (birthCountry = 'D.R.') THEN 'Dominican Republic'
```

```
WHEN (birthCountry = 'Venezuela') THEN 'Venezuela'
WHEN (birthCountry = 'P.R.') THEN 'Puerto Rico '
WHEN (birthCountry = 'Cuba') THEN 'Cuba'
WHEN (birthCountry = 'CAN') THEN 'Canada'
WHEN (birthCountry = 'Mexico') THEN 'Mexico'
ELSE 'Other'
END AS birthCountry
    FROM (
      SELECT Batting.playerID AS playerID, yearID, birthCountry
      FROM Batting
     LEFT JOIN People
        ON (Batting.playerID = People.playerID)
    ) q01
   WHERE (yearID > 1960.0)
  ) q01
  GROUP BY yearID, birthCountry
) q01
```

```
library(ggplot2)
plot_data |>
    ggplot() +
    geom_col(
        mapping = aes(yearID, percentage, fill = birthCountry),
        width = 1
    ) +
    theme_minimal() +
    theme(
        legend.title = element_blank(),
        legend.position = "top"
    )
```



i Defining methods for the data model

As part of our lahmanFromCon() function, our data model object has the class "lahman_ref". Therefore, apart from creating user-friendly functions to work with our Lahman data model, we can also define methods for this object.

class(lahman)

[1] "lahman ref" "list"

With this we can make some specific methods for a "lahman_ref" object. For example, we can define a print method like so:

```
print.lahman_ref <- function(x, ...) {
  len <- length(names(x))
  cli_h1("# Lahman reference - {len} tables")
  cli_li(paste("{.strong tables:}", paste(names(x), collapse = ", ")))
  invisible(x)
}</pre>
```

Now we can see a summary of our Lahman data model when we print the object.

lahman

```
-- # Lahman reference - 28 tables ------

* tables: AllstarFull, Appearances, AwardsManagers, AwardsPlayers,
AwardsManagers, AwardsShareManagers, Batting, BattingPost, CollegePlaying,
Fielding, FieldingOF, FieldingOFsplit, FieldingPost, HallOfFame, HomeGames,
LahmanData, Managers, ManagersHalf, Parks, People, Pitching, PitchingPost,
Salaries, Schools, SeriesPost, Teams, TeamsFranchises, TeamsHalf

And we can see that this print is being done by the method we defined.

library(sloop)
s3_dispatch(print(lahman))

=> print.lahman_ref
    print.list
    * print.default
```

4.3 Building efficient analytic pipelines

4.3.1 The risk of "clean" R code

Following on from the above approach, we might think it is a good idea to make another function addBirthYear(). We can then use it along with our addBirthCountry() to get a summarised average salary by birth country and birth year.

```
addBirthYear <- function(lahmanTbl){
  lahmanTbl |>
    left_join(
    lahman$People |>
        select("playerID", "birthYear"),
    by = "playerID"
  )
}

lahman$Salaries |>
  addBirthCountry() |>
  addBirthYear() |>
  group_by(birthCountry, birthYear) |>
  summarise(average_salary = mean(salary), .groups = "drop")
```

```
# Source:
            SQL [?? x 3]
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
   birthCountry birthYear average_salary
                     <int>
   <chr>
                                     <dbl>
1 USA
                      1966
                                 1761151.
2 Venezuela
                      1974
                                 4269365.
3 D.R.
                      1984
                                 2924854.
4 Mexico
                      1982
                                 1174912.
5 Panama
                      1981
                                  555833.
6 USA
                      1978
                                 3133596.
7 P.R.
                      1959
                                  297786.
8 USA
                      1961
                                  811250.
9 USA
                                  728740.
                      1990
10 USA
                      1950
                                  625076.
# i more rows
```

Although the R code looks fine, when we look at the SQL we can see that our query has two joins to the People table. One join gets information on the birth country and the other on the birth year.

```
SQL>
SELECT birthCountry, birthYear, AVG(salary) AS average_salary
FROM (
    SELECT
        Salaries.*,
        "People...2".birthCountry AS birthCountry,
        "People...3".birthYear AS birthYear
FROM Salaries
    LEFT JOIN People "People...2"
        ON (Salaries.playerID = "People...2".playerID)
    LEFT JOIN People "People...3"
        ON (Salaries.playerID = "People...3".playerID)
) q01
GROUP BY birthCountry, birthYear
```

To improve the performance of the code, we can build a single function to get simultaneously the birth country and birth year, so only one join is done.

```
addCharacteristics <- function(lahmanTbl){
  lahmanTbl |>
  left_join(
```

```
lahman$People |>
        select("playerID", "birthYear", "birthCountry"),
      by = "playerID"
}
lahman$Salaries |>
  addCharacteristics() |>
  group_by(birthCountry, birthYear) |>
  summarise(average_salary = mean(salary), .groups = "drop")
# Source:
            SQL [?? x 3]
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
   birthCountry birthYear average_salary
   <chr>
                    <int>
                                    <dbl>
 1 USA
                     1966
                                 1761151.
 2 Venezuela
                                 4269365.
                     1974
 3 D.R.
                     1984
                                 2924854.
 4 Mexico
                     1982
                                 1174912.
 5 Panama
                     1981
                                  555833.
 6 USA
                     1978
                                 3133596.
 7 P.R.
                                  297786.
                     1959
 8 USA
                     1961
                                  811250.
 9 USA
                                  728740.
                     1990
10 USA
                     1950
                                  625076.
# i more rows
  i Show query
  SELECT birthCountry, birthYear, AVG(salary) AS average_salary
 FROM (
    SELECT Salaries.*, birthYear, birthCountry
    FROM Salaries
    LEFT JOIN People
      ON (Salaries.playerID = People.playerID)
  GROUP BY birthCountry, birthYear
```

This query produces the same result but is simpler than the previous one, thus reducing the computational cost of the analysis. This shows the importance of being aware of the SQL code being executed when working in R with databases.

4.3.2 Piping and SQL

Piping functions has little impact on performance when using R with data in memory. However, when working with a database, the SQL generated will differ when using multiple function calls (with a separate operation specified in each) instead of multiple operations within a single function call.

For example, a single mutate function creating two new variables would generate the below SQL.

```
lahman$People |>
mutate(
   birthDatePlus1 = add_years(x = birthDate, n = 1L),
   birthDatePlus10 = add_years(x = birthDate, n = 10L)
) |>
select("playerID", "birthDatePlus1", "birthDatePlus10") |>
show_query()

<SQL>
SELECT
   playerID,
   DATE_ADD(birthDate, INTERVAL (1) year) AS birthDatePlus1,
   DATE_ADD(birthDate, INTERVAL (10) year) AS birthDatePlus10
FROM People
```

Whereas the SQL will be different if these were created using multiple mutate calls (with now one being created in a sub-query).

```
lahman$People |>
  mutate(birthDatePlus1 = add_years(x = birthDate, n = 1L)) |>
  mutate(birthDatePlus10 = add_years(x = birthDate, n = 10L)) |>
  select("playerID", "birthDatePlus1", "birthDatePlus10") |>
  show_query()
```

```
<SQL>
SELECT
  playerID,
  birthDatePlus1,
  DATE_ADD(birthDate, INTERVAL (10) year) AS birthDatePlus10
FROM (
  SELECT People.*, DATE_ADD(birthDate, INTERVAL (1) year) AS birthDatePlus1
  FROM People
) q01
```

4.3.3 Computing intermediate queries

Let's now summarise home runs (Batting table) and strike outs (Pitching table) by college player and their birth year. We can do this like so:

```
players_with_college <- lahman$People |>
  select("playerID", "birthYear") |>
  inner_join(
    lahman$CollegePlaying |>
      filter(!is.na(schoolID)) |>
      distinct(playerID, schoolID),
    by = "playerID"
  )
lahman$Batting |>
  left_join(
    players_with_college,
    by = "playerID"
  ) |>
  group_by(schoolID, birthYear) |>
  summarise(home_runs = sum(H, na.rm = TRUE), .groups = "drop") |>
  collect()
```

```
# A tibble: 6,205 x 3
  schoolID birthYear home_runs
                            <dbl>
  <chr>
                 <int>
1 rice
                  1981
2 ucsd
                 1968
                                3
                               27
3 virginia
                  1987
4 cacerri
                                3
                   1971
5 usc
                   1947
                               11
6 lsu
                   1927
                             1832
7 wake
                               72
                   1915
8 pepperdine
                   1969
                                1
9 lsu
                   1978
                                2
10 miamidade
                                0
                   1982
# i 6,195 more rows
```

```
lahman$Pitching |>
left_join(
   players_with_college,
   by = "playerID"
```

```
) |>
group_by(schoolID, birthYear) |>
summarise(strike_outs = sum(SO, na.rm = TRUE), .groups = "drop")|>
collect()
```

```
# A tibble: 3,663 x 3
   schoolID
              birthYear strike_outs
   <chr>
                  <int>
                               <dbl>
1 pennst
                    1981
                                  340
                                 327
2 cacerri
                    1971
3 usc
                                  275
                    1947
                                   4
4 pepperdine
                    1969
                                  162
5 lsu
                    1978
6 miamidade
                    1982
                                   56
7 upperiowa
                                   11
                    1918
8 jamesmad
                    1966
                                    4
9 flinternat
                                  133
                    1971
10 ucla
                                  323
                    1984
# i 3,653 more rows
```

If we look at the SQL code we will realise that there is code duplication, because as part of each full query, we have run our players_with_college query.

```
Show query
<SQL>
SELECT schoolID, birthYear, SUM(H) AS home_runs
FROM (
  SELECT Batting.*, birthYear, schoolID
  FROM Batting
  LEFT JOIN (
    SELECT People.playerID AS playerID, birthYear, schoolID
    FROM People
    INNER JOIN (
      SELECT DISTINCT playerID, schoolID
      FROM CollegePlaying
      WHERE (NOT((schoolID IS NULL)))
    ) RHS
      ON (People.playerID = RHS.playerID)
    ON (Batting.playerID = RHS.playerID)
```

```
) q01
GROUP BY schoolID, birthYear
SELECT schoolID, birthYear, SUM(SO) AS strike_outs
FROM (
  SELECT Pitching.*, birthYear, schoolID
  FROM Pitching
  LEFT JOIN (
    SELECT People.playerID AS playerID, birthYear, schoolID
    FROM People
    INNER JOIN (
      SELECT DISTINCT playerID, schoolID
      FROM CollegePlaying
      WHERE (NOT((schoolID IS NULL)))
    ) RHS
      ON (People.playerID = RHS.playerID)
    ON (Pitching.playerID = RHS.playerID)
) q01
GROUP BY schoolID, birthYear
```

To avoid this, we can make use of the **compute()** function to force the computation of the players_with_college query to a temporary table in the database.

```
players_with_college <- players_with_college |>
compute()
```

Now we have a temporary table with the result of our players_with_college query, and we can use this in both of our aggregation queries.

```
players_with_college |>
    show_query()

<SQL>
SELECT *
FROM dbplyr_qHA60HEppP

lahman$Batting |>
    left_join(players_with_college, by = "playerID") |>
```

group_by(schoolID, birthYear) |>

```
summarise(home_runs = sum(H, na.rm = TRUE), .groups = "drop") |>
collect()
```

```
# A tibble: 6,205 x 3
   schoolID
             birthYear home_runs
   <chr>
                  <int>
                             <dbl>
1 vermont
                   1869
                                38
2 michigan
                   1967
                                 2
3 nmstate
                   1968
                                 0
4 cacerri
                   1971
                                 3
5 chicago
                                 2
                   1874
                                28
6 byu
                   1961
7 pepperdine
                   1969
                                 1
8 lsu
                                 2
                   1978
9 miamidade
                   1982
                              1611
10 stanford
                   1961
# i 6,195 more rows
```

```
lahman$Pitching |>
  left_join(players_with_college, by = "playerID") |>
  group_by(schoolID, birthYear) |>
  summarise(strike_outs = sum(SO, na.rm = TRUE), .groups = "drop") |>
  collect()
```

```
# A tibble: 3,663 x 3
   schoolID birthYear strike_outs
   <chr>
                 <int>
                              <dbl>
 1 longbeach
                  1968
                                273
2 elon
                  1921
                                13
3 lehigh
                  1901
                                 1
                                275
4 usc
                  1947
5 tamukvill
                                409
                  1978
6 stanford
                  1972
                                218
7 upenn
                                14
                  1964
8 arkansas
                  1962
                                537
9 kentucky
                  1985
                                91
10 txsjjcn
                  1983
                                571
# i 3,653 more rows
```

```
i Show query
<SQL>
SELECT schoolID, birthYear, SUM(H) AS home_runs
  SELECT Batting.*, birthYear, schoolID
  FROM Batting
  LEFT JOIN dbplyr_qHA60HEppP
    ON (Batting.playerID = dbplyr_qHA60HEppP.playerID)
GROUP BY schoolID, birthYear
<SQL>
SELECT schoolID, birthYear, SUM(SO) AS strike_outs
  SELECT Pitching.*, birthYear, schoolID
  FROM Pitching
  LEFT JOIN dbplyr_qHA60HEppP
    ON (Pitching.playerID = dbplyr_qHA6OHEppP.playerID)
) q01
GROUP BY schoolID, birthYear
```

In this example, the SQL code of the intermediate table, players_with_college, was quite simple. However, in some cases, the SQL associated code can become very complicated and unmanageable, resulting with inefficient code. Therefore, although we do not want to overuse computation of intermediate queries, it is often useful when creating our analytic pipelines.

indexes

Some SQL dialects use indexes for more efficient 'joins' performance. Briefly speaking, indexes store the location of the different values of a column. Every time that you create a new table with <code>compute()</code>, the indexes will not be carried over. Hence, if you want your new table to keep some indexes, you will have to add them manually. That is why sometimes it will not be more efficient to add a <code>compute()</code> in between, because the new table generated will not have the indexes that make your query to be executed faster.

4.4 Disconnecting from the database

Now that we have reached the end of this example, we can close our connection to the database.

dbDisconnect(conn = con)

Part II Working with the OMOP CDM from R

In this second half of the book, we will focus on how we can work with data in the OMOP CDM format from R.

- In Chapter 5 we will see how to create a cdm_reference in R, a data model that contains references to the OMOP CDM tables and provides the foundation for analysis.
- The OMOP CDM is a person-centric model, and the person and observation period tables are two key tables for any analysis. In Chapter 6 we will see more on how these tables can be used as the starting point for identifying your study participants.
- In Chapter 7 we will see how to add demographics information to different tables of interest and summarise it using dplyr code. Finally, we will also see how to use tidyverse verbs to add some custom features.
- In Chapter 8 we will have a look at the cohort object, how it is defined and what are their attributes. We will also see how to create some simple base cohorts and apply some inclusion criteria to them.
- Finally, in Chapter 9 we will learn how to intersect cohorts with one another, extracting counts, presence indicators, specific dates, or time differences to obtain the information of interest for our study population.

5 Creating a CDM reference

5.1 The OMOP CDM layout

The OMOP CDM standardises the structure of healthcare data. Data is stored across a system of tables with established relationships between them. In other words, the OMOP CDM provides a relational database structure, with version 5.4 of the OMOP CDM shown below.



Figure 5.1: OMOP CDM v5.4 entity-relationship diagram retrieved from https://ohdsi.github.io/CommonDataModel/cdm54erd.html

5.2 Creating a reference to the OMOP CDM

As we saw in Chapter 4, creating a data model in R to represent the OMOP CDM can provide a basis for analytic pipelines using the data. Luckily for us, we won't have to create functions and methods for this ourselves. Instead, we will use the omopgenerics package which defines a data model for OMOP CDM data and the CDMConnector package which provides functions for connecting to OMOP CDM data held in a database.

To see how this works, we will use the <code>omock</code> package to create example data in the format of the OMOP CDM, which we will then copy to a DuckDB database created by the <code>duckdb</code> package.

```
library(duckdb)
library(dplyr)
library(omock)
library(CDMConnector)
library(palmerpenguins)
cdm local <- mockCdmReference() |>
  mockPerson(nPerson = 100) |>
  mockObservationPeriod() |>
  mockConditionOccurrence() |>
  mockDrugExposure() |>
  mockObservation() |>
  mockMeasurement() |>
  mockVisitOccurrence() |>
  mockProcedureOccurrence()
con <- dbConnect(drv = duckdb())</pre>
src <- dbSource(con = con, writeSchema = "main")</pre>
cdm <- insertCdmTo(cdm = cdm local, to = src)</pre>
```

Note that <code>insertCdmTo()</code> output is already a <code><cdm_reference></code> object. But how would we create this cdm reference from the connection? We can use the function <code>cdmFromCon()</code> from <code>CDMConnector</code> to create our cdm reference. Note that as well as specifying the schema containing our OMOP CDM tables, we will also specify a write schema where any database tables we create during our analysis will be stored. Often, our OMOP CDM tables will be in a schema that we only have read-access to, and we'll have another schema where we can have write-access and where intermediate tables can be created for a given study.

```
cdm <- cdmFromCon(con = con,</pre>
                   cdmSchema = "main",
                   writeSchema = "main",
                   cdmName = "example_data")
```

cdm

```
-- # OMOP CDM reference (duckdb) of example_data -----
```

- * omop tables: cdm_source, concept, concept_ancestor, concept_relationship, concept_synonym, condition_occurrence, drug_exposure, drug_strength, measurement, observation, observation_period, person, procedure_occurrence, visit_occurrence, vocabulary
- * cohort tables: -
- * achilles tables: -
- * other tables: -

Setting a write prefix

We can also specify a write prefix and this will be used whenever permanent tables are created in the write schema. This can be useful when we're sharing our write schema with others and want to avoid table name conflicts and easily drop tables created as part of a particular study.

```
cdm <- cdmFromCon(con = con,</pre>
                   cdmSchema = "main",
                   writeSchema = "main",
                   writePrefix = "my_study_",
                   cdmName = "example_data")
```

Note you only have to specify this writePrefix once at the connection stage, and then the cdm reference object will store that and use it every time that you create a new table.

We can see that we now have an object that contains references to all the OMOP CDM tables. We can reference specific tables using the "\$" or "[[...]]" operators.

cdm\$person

```
# Source:
            table<person> [?? x 18]
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
   person_id gender_concept_id year_of_birth month_of_birth day_of_birth
       <int>
                          <int>
                                        <int>
                                                        <int>
                                                                      <int>
           1
                           8507
                                          1978
                                                            8
                                                                         18
1
2
           2
                           8532
                                                           10
                                          1972
                                                                         11
3
           3
                           8507
                                          1996
                                                            9
                                                                         20
           4
4
                           8532
                                          2000
                                                            11
                                                                         13
5
           5
                           8507
                                          1954
                                                            7
                                                                         19
6
           6
                           8532
                                          1954
                                                           10
                                                                         11
7
           7
                           8507
                                          1998
                                                             1
                                                                         16
8
           8
                           8507
                                          1991
                                                            4
                                                                         27
9
           9
                                                            9
                           8532
                                          1966
                                                                         16
          10
                                                            9
10
                           8507
                                          1963
                                                                         26
# i more rows
# i 13 more variables: race_concept_id <int>, ethnicity_concept_id <int>,
    birth_datetime <dttm>, location_id <int>, provider_id <int>,
#
    care_site_id <int>, person_source_value <chr>, gender_source_value <chr>,
#
   gender_source_concept_id <int>, race_source_value <chr>,
```

race_source_concept_id <int>, ethnicity_source_value <chr>,

cdm[["observation_period"]]

#

Source: table<observation_period> [?? x 5]

ethnicity_source_concept_id <int>

Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
 observation_period_id person_id observation_period_s~1 observation_period_e~2

	<int></int>	<int></int>	<date></date>	<date></date>
1	1	1	1996-05-26	1996-07-23
2	2	2	1992-06-02	2015-08-23
3	3	3	2013-09-03	2016-04-27
4	4	4	2014-08-26	2016-06-25
5	5	5	1960-03-30	2007-01-19
6	6	6	1980-03-14	2003-01-11
7	7	7	2011-04-21	2014-01-04
8	8	8	2019-05-05	2019-10-12
9	9	9	1994-04-05	2012-02-19
10	10	10	2014-12-25	2018-08-11

i more rows

```
# i abbreviated names: 1: observation_period_start_date,
# 2: observation_period_end_date
# i 1 more variable: period_type_concept_id <int>
```

Note that here we have first created a local version of the cdm with all the tables of interest with <code>omock</code> (<code>cdm_local</code>), then copied it to a DuckDB database, and finally created a reference to it with <code>CDMConnector</code>, so that we can work with the final <code>cdm</code> object as we normally would for one created with our own healthcare data. In that case, we would directly use <code>cdmFromCon()</code> with our own database information. Throughout this chapter, however, we will keep working with the mock dataset.

5.3 CDM attributes

5.3.1 CDM name

Our cdm reference will be associated with a name. By default, this name will be taken from the cdm_source_name field from the cdm_source table. We will use the function cdmName() from omopgenerics to get it.

```
cdm <- cdmFromCon(con = con,</pre>
                  cdmSchema = "main",
                  writeSchema = "main")
cdm$cdm_source
            table < cdm_source > [?? x 10]
# Source:
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
  cdm_source_name cdm_source_abbreviation cdm_holder source_description
  <chr>
                   <chr>
                                            <chr>
                                                       <chr>>
1 mock
                  <NA>
                                            <NA>
                                                       <NA>
# i 6 more variables: source_documentation_reference <chr>,
    cdm_etl_reference <chr>, source_release_date <date>,
    cdm_release_date <date>, cdm_version <chr>, vocabulary_version <chr>
cdmName(cdm)
```

[1] "mock"

However, we can instead set this name to whatever else we want when creating our cdm reference.

```
cdm <- cdmFromCon(con = con,</pre>
                   cdmSchema = "main",
                   writeSchema = "main",
                   cdmName = "my_cdm")
cdmName(cdm)
```

[1] "my_cdm"

Note that we can also get our cdm name from any of the tables in our cdm reference.

cdmName(cdm\$person)

[1] "my_cdm"



Pehind the scenes

The class of the cdm reference itself is <cdm_reference>.

class(cdm)

[1] "cdm_reference"

Each of the tables has class <cdm_table>. If the table is one of the standard OMOP CDM tables, it will also have class <omop_table>. This latter class is defined so that we can allow different behavior for these core tables (person, condition_occurrence, observation_period, etc.) compared to other tables that are added to the cdm reference during the course of running a study.

class(cdm\$person)

```
[1] "omop_table"
                             "cdm_table"
                                                       "tbl_duckdb_connection"
[4] "tbl_dbi"
                              "tbl_sql"
                                                       "tbl_lazy"
[7] "tbl"
```

We can see that cdmName() is a generic function, which works for both the cdm reference as a whole and individual tables.

```
library(sloop)
s3_dispatch(cdmName(cdm))
=> cdmName.cdm_reference
 * cdmName.default
```

```
s3_dispatch(cdmName(cdm$person))

cdmName.omop_table
=> cdmName.cdm_table
    cdmName.tbl_duckdb_connection
    cdmName.tbl_dbi
    cdmName.tbl_sql
    cdmName.tbl_lazy
    cdmName.tbl
* cdmName.default
```

5.3.2 CDM version

We can also easily check the OMOP CDM version that is being used with the function cdmVersion() from omopgenerics like so:

```
cdmVersion(cdm)
```

[1] "5.3"

i cdmVersion

Note, the cdmVersion() function also works for <cdm_table> objects:

cdmVersion(cdm\$person)

[1] "5.3"

i Methods functions

Although as stated, the <code>cdmName()</code> and <code>cdmVersion()</code> functions are defined by the omopgenerics packages, these functions are re-exported in other packages and you won't need to load omopgenerics explicitly.

5.4 Including cohort tables in the cdm reference

A cohort is a fundamental piece in epidemiological studies. Later, we'll see how to create cohorts in more detail in Chapter 8. For the moment, let's just outline how we can include

the reference to an existing cohort in our cdm reference. For this, we'll use omock to add a cohort to our local cdm and upload that to a DuckDB database again.

```
cdm_local <- cdm_local |>
  mockCohort(name = "my_study_cohort")
con <- dbConnect(drv = duckdb())
src <- dbSource(con = con, writeSchema = "main")
cdm <- insertCdmTo(cdm = cdm_local, to = src)</pre>
```

Now we can specify we want to include this existing cohort table to our cdm object when creating our cdm reference.

```
cdm$my_study_cohort |>
  glimpse()
```

Tables included in the cdm reference

Note that by default the cohort table won't be included in the cdm_reference object.

```
-- # OMOP CDM reference (duckdb) of example_data ---
* omop tables: cdm source, concept, concept ancestor, concept relationship,
concept_synonym, condition_occurrence, drug_exposure, drug_strength,
measurement, observation, observation period, person, procedure occurrence,
visit_occurrence, vocabulary
* cohort tables: -
* achilles tables: -
* other tables: -
Even if the cohort exists in the database:
dbListTables(conn = con)
                                   "concept"
 [1] "cdm_source"
 [3] "concept_ancestor"
                                   "concept_relationship"
                                   "condition_occurrence"
 [5] "concept_synonym"
 [7] "drug_exposure"
                                   "drug_strength"
 [9] "measurement"
                                   "my_study_cohort"
[11] "my_study_cohort_attrition" "my_study_cohort_codelist"
[13] "my_study_cohort_set"
                                  "observation"
[15] "observation_period"
                                  "person"
[17] "procedure_occurrence"
                                  "visit_occurrence"
[19] "vocabulary"
By default, only the default omop tables omopTables() will be included (if they exist)
into the cdm reference object.
```

5.5 Including achilles tables in the cdm reference

If we have the results tables from the **Achilles** package in our database, we can also include these in our cdm reference.

Just to show how this can be done, let's upload some empty results tables in the Achilles format.

```
dbWriteTable(conn = con,
             name = "achilles_analysis",
             value = tibble(
               analysis_id = NA_integer_,
               analysis name = NA character,
               stratum_1_name = NA_character_,
               stratum_2_name = NA_character_,
               stratum_3_name = NA_character_,
               stratum_4_name = NA_character_,
               stratum_5_name = NA_character_,
               is_default = NA_character_,
               category = NA_character_))
dbWriteTable(conn = con,
             name = "achilles_results",
             value = tibble(
               analysis_id = NA_integer_,
               stratum_1 = NA_character_,
               stratum_2 = NA_character_,
               stratum_3 = NA_character_,
               stratum_4 = NA_character_,
               stratum_5 = NA_character_,
               count_value = NA_character_))
dbWriteTable(conn = con,
             name = "achilles_results_dist",
             value = tibble(
               analysis_id = NA_integer_,
               stratum_1 = NA_character_,
               stratum_2 = NA_character_,
               stratum_3 = NA_character_,
               stratum_4 = NA_character_,
               stratum_5 = NA_character_,
               count_value = NA_character_,
               min_value = NA_character_,
               max_value = NA_character_,
               avg_value = NA_character_,
               stdev_value = NA_character_,
               median_value = NA_character_,
               p10_value = NA_character_,
               p25_value = NA_character_,
               p75_value = NA_character_,
               p90_value = NA_character_))
```

We can now include these achilles tables in our cdm reference as in the previous case.

Note we specified the achillesSchema that in this case is the same as the writeSchema and cdmSchema, but each one of them can be different and point to a separate schema in our database.

5.6 Adding other tables to the cdm reference

Let's say we have some additional **local** data that we want to add to our cdm reference. We can add this both to the same source (in this case a database) and to our cdm reference using <code>insertTable()</code> from <code>omopgenerics</code> (<code>insertTable()</code> is also re-exported in <code>CDMConnector()</code>. We will show this with the dataset <code>cars</code> built-in to R.

```
cars |>
  glimpse()
```

```
Rows: 50

Columns: 2
$ speed <dbl> 4, 4, 7, 7, 8, 9, 10, 10, 10, 11, 11, 12, 12, 12, 12, 13, 13, 13~
$ dist <dbl> 2, 10, 4, 22, 16, 10, 18, 26, 34, 17, 28, 14, 20, 24, 28, 26, 34~
```

```
cdm <- insertTable(cdm = cdm, name = "cars", table = cars)</pre>
```

We can see that now this extra table has been uploaded to the database behind our cdm reference and also added to our reference.

```
\operatorname{\mathsf{cdm}}
```

```
-- # OMOP CDM reference (duckdb) of example_data -----
```

```
* omop tables: cdm_source, concept, concept_ancestor, concept_relationship, concept_synonym, condition_occurrence, drug_exposure, drug_strength, measurement, observation, observation_period, person, procedure_occurrence, visit_occurrence, vocabulary
```

```
* cohort tables: my_study_cohort
```

* achilles tables: achilles_analysis, achilles_results, achilles_results_dist

* other tables: cars

```
cdm$cars
```

```
# Source:
            table<cars> [?? x 2]
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
   speed dist
   <dbl> <dbl>
1
       4
             2
2
       4
            10
3
       7
             4
4
       7
            22
5
       8
            16
6
       9
            10
7
      10
            18
8
      10
            26
9
      10
            34
10
            17
      11
# i more rows
```

If we already had the table in the database we could have instead just assigned it to our existing cdm reference. To see this let's upload the penguins table to our DuckDB database.

```
dbWriteTable(conn = con, name = "penguins", value = penguins)
```

Once we have this table in the database, we can just read it using the readSourceTable() function.

```
cdm <- readSourceTable(cdm = cdm, name = "penguins")
cdm</pre>
```

```
-- # OMOP CDM reference (duckdb) of example data -----
```

- * omop tables: cdm_source, concept, concept_ancestor, concept_relationship, concept_synonym, condition_occurrence, drug_exposure, drug_strength, measurement, observation, observation_period, person, procedure_occurrence, visit_occurrence, vocabulary
- * cohort tables: my_study_cohort
- * achilles tables: achilles_analysis, achilles_results, achilles_results_dist
- * other tables: cars, penguins

Note that omopgenerics provides these functions readSourceTable(), listSourceTables(), and dropSourceTable() for the easier management of the tables in the writeSchema.

listSourceTables(cdm = cdm)

```
[1] "achilles_analysis"
                                  "achilles_results"
 [3] "achilles_results_dist"
                                  "cars"
 [5] "cdm_source"
                                  "concept"
 [7] "concept_ancestor"
                                  "concept_relationship"
 [9] "concept_synonym"
                                  "condition_occurrence"
[11] "drug_exposure"
                                  "drug_strength"
[13] "measurement"
                                  "my_study_cohort"
[15] "my_study_cohort_attrition" "my_study_cohort_codelist"
[17] "my_study_cohort_set"
                                  "observation"
[19] "observation_period"
                                  "penguins"
                                  "procedure_occurrence"
[21] "person"
[23] "visit_occurrence"
                                  "vocabulary"
```

```
dropSourceTable(cdm = cdm, name = "penguins")
listSourceTables(cdm = cdm)
```

```
[1] "achilles_analysis"
                                  "achilles_results"
 [3] "achilles_results_dist"
                                  "cars"
                                  "concept"
 [5] "cdm_source"
[7] "concept_ancestor"
                                  "concept_relationship"
 [9] "concept synonym"
                                  "condition occurrence"
[11] "drug exposure"
                                  "drug strength"
[13] "measurement"
                                  "my study cohort"
[15] "my_study_cohort_attrition"
                                 "my_study_cohort_codelist"
[17] "my study cohort set"
                                  "observation"
[19] "observation_period"
                                  "person"
[21] "procedure_occurrence"
                                  "visit_occurrence"
[23] "vocabulary"
```

Difference between insertTable and dbWriteTable

- dbWriteTable() is a function from the DBI package that writes a local R data frame to a database. You need to manually specify the schema and table name and it does not update the cdm reference object.
- insertTable() is a function from the omopgenerics package designed for use with cdm reference objects. It writes a local table to the database and adds it to the list of tables in the cdm reference. Internally, it uses dbWriteTable() but also handles the schema and table name automatically using the writeSchema and writePrefix from the cdm reference.

In general, for studies using OMOP CDM data, you should use <code>insertTable()</code> rather than <code>dbWriteTable()</code>. It ensures the table is both written to the correct location in the database and accessible through the cdm reference object. Only use <code>dbWriteTable()</code> if you are confident working directly with the database and understand its structure. Note <code>insertTable()</code> would also work for a local cdm reference or any other defined cdm reference source, whereas the <code>dbWriteTable()</code> is a database specific function. TODO reference to omopgenerics supported sources.

5.7 Mutability of the cdm reference

An important characteristic of our cdm reference is that we can alter the tables in R, but the OMOP CDM data will not be affected. We will therefore only be transforming the data in our cdm object but the original datasets behind it will remain intact.

For example, let's say we want to perform a study with only people born in 1970. For this we could filter our person table to only people born in this year.

```
cdm$person <- cdm$person |>
  filter(year_of_birth == 1970)

cdm$person
```

```
# Source:
            SQL [?? x 18]
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:]
 person_id gender_concept_id year_of_birth month_of_birth day_of_birth
      <int>
                        <int>
                                      <int>
                                                      <int>
1
         67
                         8532
                                       1970
                                                         10
                                                                      20
         97
                         8532
                                       1970
                                                                      19
# i 13 more variables: race_concept_id <int>, ethnicity_concept_id <int>,
   birth_datetime <dttm>, location_id <int>, provider_id <int>,
    care_site_id <int>, person_source_value <chr>, gender_source_value <chr>,
   gender_source_concept_id <int>, race_source_value <chr>,
   race source concept id <int>, ethnicity source value <chr>,
    ethnicity_source_concept_id <int>
```

From now on, when we work with our cdm reference this restriction will continue to have been applied.

The original OMOP CDM data itself however will remain unaffected. We can see that, indeed, if we create our reference again the underlying data is unchanged.

The mutability of our cdm reference is a useful feature for studies as it means we can easily tweak our OMOP CDM data if needed. Meanwhile, leaving the underlying data unchanged is essential so that other study code can run against the data, unaffected by any of our changes.

One thing we can't do, though, is alter the structure of OMOP CDM tables. For example, the following code would cause an error as the person table must always have the column person_id.

```
cdm$person <- cdm$person |>
rename("new_id" = "person_id")
```

```
Error in `newOmopTable()`:
! person_id is not present in table person
```

In such a case we would have to call the table something else first, and then run the previous code:

```
cdm$person_new <- cdm$person |>
  rename("new_id" = "person_id") |>
  compute(name = "person_new")
```

Now we would be allowed to have this new table as an additional table in our cdm reference, knowing it was not in the format of one of the core OMOP CDM tables.

```
\operatorname{cdm}
```

```
-- # OMOP CDM reference (duckdb) of example_data -----
```

^{*} omop tables: cdm_source, concept, concept_ancestor, concept_relationship, concept_synonym, condition_occurrence, drug_exposure, drug_strength, measurement, observation, observation_period, person, procedure_occurrence, visit_occurrence, vocabulary

```
* cohort tables: -
* achilles tables: -
* other tables: person_new
```

The package omopgenerics provides a comprehensive list of the required features of a valid cdm reference. You can read more about it here.

i Name consistency

Note also that there must be a name consistency between the name of the table and the assignment in the cdm reference object.

```
cdm$new_table <- cdm$person |>
   compute(name = "not_new_table")

Error in `[[<-`:
   x You can't assign a table named not_new_table to new_table.
i You can change the name using compute:
cdm[['new_table']] <- yourObject |>
   dplyr::compute(name = 'new_table')
i You can also change the name using the `name` argument in your function:
   `name = 'new_table'`.
```

5.8 Working with temporary and permanent tables

When we create new tables and our cdm reference is in a database we have a choice between using temporary or permanent tables. In most cases we can work with these interchangeably. Below we create one temporary table and one permanent table. We can see that both of these tables have been added to our cdm reference and that we can use them in the same way. Note that any new computed table will by default be temporary unless otherwise specified.

```
cdm$person_new_temp <- cdm$person |>
  head(5) |>
  compute(temporary = TRUE)
```

```
cdm$person_new_permanent <- cdm$person |>
  head(5) |>
  compute(name = "person_new_permanent", temporary = FALSE)
```

cdm

cdm\$person_new_temp

- # Source: table<og_001_1761240524> [?? x 18]
- # Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:] person_id gender_concept_id year_of_birth month_of_birth day_of_birth

	<int></int>	<int></int>	<int></int>	<int></int>	<int></int>
1	1	8507	1978	8	18
2	2	8532	1972	10	11
3	3	8507	1996	9	20
4	4	8532	2000	11	13
5	5	8507	1954	7	19

- # i 13 more variables: race_concept_id <int>, ethnicity_concept_id <int>,
- # birth_datetime <dttm>, location_id <int>, provider_id <int>,
- # care_site_id <int>, person_source_value <chr>, gender_source_value <chr>,
- # gender_source_concept_id <int>, race_source_value <chr>,
- # race_source_concept_id <int>, ethnicity_source_value <chr>,
- # ethnicity_source_concept_id <int>

cdm\$person_new_permanent

- # Source: table<person_new_permanent> [?? x 18]
- # Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1/:memory:] person_id gender_concept_id year_of_birth month_of_birth day_of_birth

-					
	<int></int>	<int></int>	<int></int>	<int></int>	<int></int>
1	1	8507	1978	8	18
2	2	8532	1972	10	11
3	3	8507	1996	9	20
4	4	8532	2000	11	13
5	5	8507	1954	7	19

- # i 13 more variables: race_concept_id <int>, ethnicity_concept_id <int>,
- # birth_datetime <dttm>, location_id <int>, provider_id <int>,
- # care_site_id <int>, person_source_value <chr>, gender_source_value <chr>,
- # gender_source_concept_id <int>, race_source_value <chr>,
- # race_source_concept_id <int>, ethnicity_source_value <chr>,
- # ethnicity_source_concept_id <int>

One benefit of working with temporary tables is that they will be automatically dropped at the end of the session, whereas the permanent tables will be left in the database until explicitly dropped. This helps maintain the original database structure tidy and free of irrelevant data. However, one disadvantage of using temporary tables is that we will generally accumulate more and more of them as we go (in a single R session), whereas we can overwrite permanent tables continuously. For example, if our study code contains a loop that requires a compute, we would either overwrite an intermediate permanent table 100 times or create 100 different temporary tables in the process. In the latter case we should be wary of consuming a lot of drive memory, which could lead to performance issues or even crashes.

i name argument in compute()

Note that in the previous examples we explicitly specified the name of the new table and whether it must be temporary or permanent (temporary = FALSE), but we do not need to populate the temporary field explicitly as if name is left as NULL (default behavior), then the table will be temporary (temporary = TRUE), and if the name argument is populated with a character (e.g., name = "my_custom_table"), then the created table will be permanent:

```
cdm$person_new_temp <- cdm$person |>
  compute()

cdm$person_new_permanent <- cdm$person |>
  compute(name = "person_new_permanent")
```

5.9 Disconnecting

Once we have finished our analysis we can close our connection to the database behind our cdm reference.

```
cdmDisconnect(cdm)
```

5.10 Further reading

- Català M, Burn E (2025). omopgenerics: Methods and Classes for the COMOP Common Data Model. R package version 1.3.1, https://darwin-eu.github.io/omopgenerics/.
- Black A, Gorbachev A, Burn E, Català M, Nika I (2025). CDMConnector: Connect to an OMOP Common Data Model. R package version 2.2.0, https://darwin-eu.github.io/CDMConnector/.
- OmopOnPostgres (in progress)
- OmopOnSpark (in progress)
- OmopOnDuckDB (in progress)

6 Exploring the OMOP CDM

For this chapter, we'll use a synthetic COVID-19 dataset ("synthea-covid19-10k") that can be downloaded with the **omock** package. A characterisation of this dataset to better understant its content can be found in the following Shiny App https://dpa-pde-oxford.shinyapps.io/OmopSketchCharacterisation/.

```
library(dplyr)
library(omock)
library(ggplot2)
library(clock)
library(omopgenerics) # TODO https://github.com/OHDSI/omock/issues/189
library(PatientProfiles)
library(OmopSketch)
```

You can download the dataset using the function downloadMockDataset():

```
downloadMockDataset(datasetName = "synthea-covid19-10k")
```

i Setup OMOP_DATA_FOLDER

The downloadMockDataset function checks if the database has already been downloaded, and if it is not, it is downloaded in a temporary directory. To avoid downloading the database every time we want to use it, we need to set up the OMOP_DATA_FOLDER. To do that, you need to create an environment variable named OMOP_DATA_FOLDER. You can add it in the R environment file (usethis::edit_r_environ()) or using Sys.setenv(OMOP_DATA_FOLDER = "..."). Either way, OMOP_DATA_FOLDER should point to a folder where the dataset will be downloaded. This way, the dataset will be stored permanently on your computer and you will not have to download it every time you want to use it.

Note that this folder is defined by omopgenerics and is also used by other packages to store OMOP-related data.

Once the dataset is downloaded you can create the cdm reference:

cdm

```
-- # OMOP CDM reference (duckdb) of synthea-covid19-10k -----
```

* omop tables: attribute_definition, care_site, cdm_source, cohort_definition, concept, concept_ancestor, concept_class, concept_relationship, concept_synonym, condition_era, condition_occurrence, cost, death, device_exposure, domain, dose_era, drug_era, drug_exposure, drug_strength, fact_relationship, location, measurement, metadata, note, note_nlp, observation, observation_period, payer_plan_period, person, procedure_occurrence, provider, relationship, source_to_concept_map, specimen, visit_detail, visit_occurrence, vocabulary

```
* cohort tables: -
```

- * achilles tables: -
- * other tables: -

i mockCdmFromDataset()

Note that if you call the function without downloading the database first you will be prompted to download it (or it will be downloaded directly if the session is not interactive).

By default, the mock cdm returns a local dataset that can be inserted in any source of interest using the insertCdmTo() function. Setting the source to source = "duckdb" will return an in-memory DuckDB cdm_reference with two schemas: cdmSchema = "main" and writeSchema = "results".

6.1 Counting people

The OMOP CDM is person-centric, with the person table containing records to uniquely identify each person in the database. As each row refers to a unique person, we can quickly get a count of the number of individuals in the database like so

```
cdm$person |>
count()
```

The person table also contains some demographic information, including a gender concept for each person. We can easily get a count grouped by this variable. As the concept id is just a number it is more useful to get the concept name, this can be done with a join to the concept table.

```
cdm$person |>
  group_by(gender_concept_id) |>
  count() |>
  left_join(cdm$concept, by = c("gender_concept_id" = "concept_id")) |>
  select("gender_concept_id", "concept_name", "n") |>
  collect()
```

• Vocabulary tables

Above we've got counts by specific concept IDs recorded in the condition occurrence table. What these IDs represent is described in the concept table. Here we have the name associated with the concept, along with other information such as its domain and vocabulary id.

```
cdm$concept |>
  glimpse()
```

Rows: ?? Columns: 10

```
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/RtmpRTbnLD/file31c052422a58.duckdb]
$ concept id
                <int> 45756805, 45756804, 45756803, 45756802, 45756801, 457~
$ concept_name
                 <chr> "Pediatric Cardiology", "Pediatric Anesthesiology", "~
                 <chr> "Provider", "Provider", "Provider", "Provider", "Prov-
$ domain id
                 <chr> "ABMS", "ABMS", "ABMS", "ABMS", "ABMS", "ABMS", "ABMS"
$ vocabulary_id
$ concept_class_id <chr> "Physician Specialty", "Physician Specialty", "Physic~
<chr> "OMOP4821938", "OMOP4821939", "OMOP4821940", "OMOP482~
$ concept_code
$ valid_start_date <date> 1970-01-01, 1970-01-01, 1970-01-01, 1970-01-
01, 1970~
$ valid_end_date <date> 2099-12-31, 2099-12-31, 2099-12-31, 2099-12-
31, 2099~
$ invalid reason
                 Other vocabulary tables capture other information about concepts, such as the direct
relationships between concepts (the concept relationship table) and hierarchical relation-
ships between (the concept ancestor table).
cdm$concept_relationship |>
 glimpse()
Rows: ??
Columns: 6
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/RtmpRTbnLD/file31c052422a58.duckdb]
                 <int> 35804314, 35804314, 35804314, 35804327, 35804327, 358~
$ concept id 1
$ concept_id_2
                 <int> 912065, 42542145, 42542145, 35803584, 42542145, 42542
$ relationship_id <chr>> "Has modality", "Has accepted use", "Is current in", ~
$ valid_start_date <date> 2021-01-26, 2019-08-29, 2019-08-29, 2019-05-
27, 2019~
$ valid_end_date
                <date> 2099-12-31, 2099-12-31, 2099-12-31, 2099-12-
31, 2099~
cdm$concept_ancestor |>
 glimpse()
Rows: ??
Columns: 4
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/RtmpRTbnLD/file31c052422a58.duckdb]
```

More information on the vocabulary tables (as well as other tables in the OMOP CDM version 5.3) can be found at https://ohdsi.github.io/CommonDataModel/cdm53.html# Vocabulary Tables.

addConceptName()

\$ race_source_concept_id

The PatientProfiles package has a utility function that helps you to add concept names to a table. By default, any column that ends in concept_id will be used to join to the concept table and add a concept_name column:

```
cdm$person |>
 addConceptName() |>
 glimpse()
Rows: ??
Columns: 24
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/RtmpRTbnLD/file31c052422a58.duckdb]
                                <int> 10752, 10754, 10746, 10753, 10745, 10~
$ person_id
$ gender_concept_id
                                <int> 8532, 8507, 8532, 8507, 8532, 8507, 8~
$ year_of_birth
                                <int> 2017, 1942, 2017, 1953, 2010, 1930, 1~
                                <int> 2, 6, 10, 5, 9, 1, 3, 7, 5, 12, 8, 10~
$ month_of_birth
$ day_of_birth
                                <int> 1, 1, 18, 9, 27, 10, 16, 28, 21, 21, ~
$ birth_datetime
                                <dttm> 2017-02-01, 1942-06-01, 2017-
10-18, ~
$ race_concept_id
                                <int> 8527, 8527, 8515, 8527, 8527, 8515, 8~
                                <int> 38003564, 38003564, 38003564, 3800356~
$ ethnicity_concept_id
$ location id
                                $ provider id
                                <int> NA, NA, NA, NA, NA, NA, NA, NA,
$ care_site_id
                                $ person_source_value
                                <chr> "ffecf9fe-26c1-605c-0ce7-
6133f75eb6dc~
$ gender_source_value
                                <chr> "F", "M", "F", "M", "F", "M", "F", "M~
$ gender_source_concept_id
                                <int> 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0~
                                <chr> "white", "white", "asian", "white", "~
$ race_source_value
```

<int> 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0~

```
$ ethnicity_source_value
                              <chr> "nonhispanic", "nonhispanic", "nonhis~
$ ethnicity_source_concept_id
                              <int> 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0,
                              <chr> "FEMALE", "MALE", "FEMALE", "MALE",
$ gender_concept_id_name
$ race_concept_id_name
                              <chr> "White", "White", "Asian", "White",
$ ethnicity_concept_id_name
                              <chr> "Not Hispanic or Latino", "Not Hispan~
$ gender_source_concept_id_name
                              <chr> "No matching concept", "No matching c~
$ race_source_concept_id_name
                              <chr> "No matching concept", "No matching c~
$ ethnicity_source_concept_id_name <chr> "No matching concept", "No matching c~
Note you can edit the arguments to only use one desired column or edit the concept name
column.
cdm$person |>
 addConceptName(column = "gender_concept_id", nameStyle = "sex") |>
 glimpse()
Rows: ??
Columns: 19
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/RtmpRTbnLD/file31c052422a58.duckdb]
                          <int> 10752, 10754, 10746, 10753, 10745, 10751, ~
$ person_id
$ gender_concept_id
                          <int> 8532, 8507, 8532, 8507, 8532, 8507, 8532, ~
                          <int> 2017, 1942, 2017, 1953, 2010, 1930, 1970, ~
$ year_of_birth
$ month_of_birth
                          <int> 2, 6, 10, 5, 9, 1, 3, 7, 5, 12, 8, 10, 5, ~
$ day_of_birth
                          <int> 1, 1, 18, 9, 27, 10, 16, 28, 21, 21, 20, 1~
$ birth_datetime
                          <dttm> 2017-02-01, 1942-06-01, 2017-10-
18, 1953-~
$ race concept id
                          <int> 8527, 8527, 8515, 8527, 8527, 8515, 8527, ~
$ ethnicity_concept_id
                          <int> 38003564, 38003564, 38003564, 38003564, 38~
                          $ location id
$ provider_id
                          $ care site id
                          $ person_source_value
                          <chr> "ffecf9fe-26c1-605c-0ce7-
6133f75eb6dc", "f~
                          <chr> "F", "M", "F", "M", "F", "M", "F", "M", "F~
$ gender_source_value
$ gender_source_concept_id
                          <chr> "white", "white", "asian", "white", "white~
$ race_source_value
$ race_source_concept_id
                          <chr> "nonhispanic", "nonhispanic", "nonhispanic~
$ ethnicity_source_value
<chr> "FEMALE", "MALE", "FEMALE", "MALE", "FEMAL~
$ sex
```

6.2 Summarising observation periods

The observation period table contains records indicating spans of time over which clinical events can be reliably observed for the people in the person table (see formal definition). Someone can potentially have multiple observation periods. So, say we wanted a count of people grouped by the year during which their first observation period started.

To do this first we would need to get the first observation period per person:

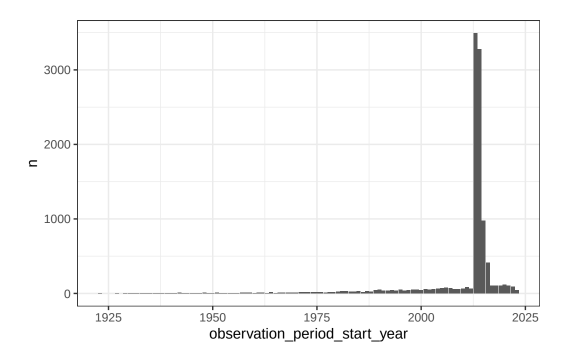
```
first_observation_period <- cdm$observation_period |>
  group_by(person_id) |>
  arrange(observation_period_start_date) |>
  filter(row_number() == 1) |>
  compute()
```

Now we can add this to the person table to make sure that all individuals defined in the observation period table are also defined the person table. Later we can extract the *observation_period_start_year* and count the number of records associated in each year:

```
first_records_per_year <- cdm$person |>
  left_join(first_observation_period, by = "person_id") |>
  mutate(observation_period_start_year = get_year(observation_period_start_date)) |>
  group_by(observation_period_start_year) |>
  count() |>
  collect()
```

Finally we can plot the counts with ggplot2.

```
ggplot(first_records_per_year) +
  geom_col(mapping = aes(x = observation_period_start_year, y = n)) +
  theme_bw()
```

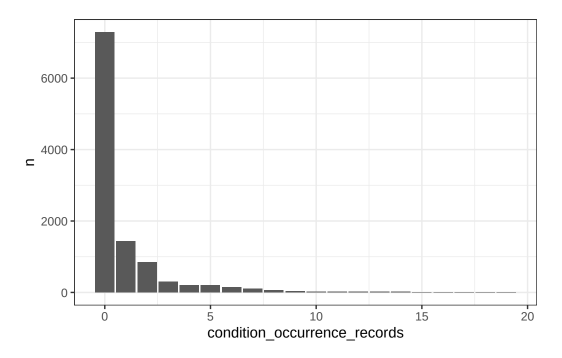


6.3 Summarising clinical records

What's the number of condition occurrence records per person in the database? We can find this out like so

```
number_condition_occurrence_records <- cdm$person |>
  left_join(
    cdm$condition_occurrence |>
        group_by(person_id) |>
        count(name = "condition_occurrence_records"),
        by="person_id"
  ) |>
  mutate(condition_occurrence_records = coalesce(condition_occurrence_records, 0)) |>
  group_by(condition_occurrence_records) |>
  count() |>
  count() |>
  collect()

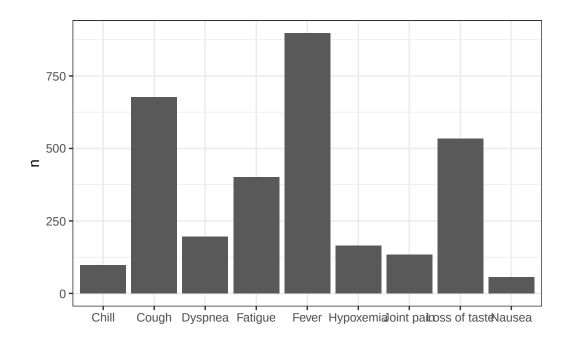
ggplot(number_condition_occurrence_records) +
  geom_col(mapping = aes(x = condition_occurrence_records, y = n)) +
  theme_bw()
```



How about we were interested in getting record counts for some specific concepts related to Covid-19 symptoms?

```
codes <- c(437663, 437390, 31967, 4289517, 4223659, 312437, 434490, 254761, 77074)
symptoms_records <- cdm$condition_occurrence |>
    filter(condition_concept_id %in% codes) |>
    group_by(condition_concept_id) |>
    count() |>
    addConceptName(column = "condition_concept_id", nameStyle = "concept_name") |>
    collect()

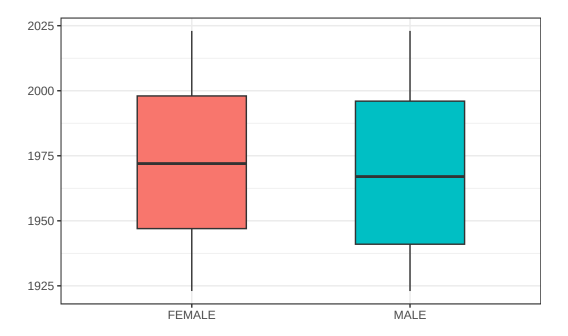
ggplot(symptoms_records) +
    geom_col(mapping = aes(x = concept_name, y = n)) +
    theme_bw()+
    xlab("")
```



We can also use summarise for various other calculations

As we've seen before, we can also quickly get results for various groupings or restrictions

```
grouped_summary <- cdm$person |>
   group_by(gender_concept_id) |>
   summarise(min_year_of_birth = min(year_of_birth, na.rm=TRUE),
             q25_year_of_birth = quantile(year_of_birth, 0.25, na.rm=TRUE),
             median_year_of_birth = median(year_of_birth, na.rm=TRUE),
             q75_year_of_birth = quantile(year_of_birth, 0.75, na.rm=TRUE),
             max_year_of_birth = max(year_of_birth, na.rm=TRUE)) |>
 left_join(cdm$concept, by = c("gender_concept_id" = "concept_id")) |>
 collect()
grouped_summary |>
 ggplot(mapping = aes(x = concept_name, group = concept_name, fill = concept_name)) +
 geom_boxplot(mapping = aes(
   lower = q25_year_of_birth,
   upper = q75_year_of_birth,
   middle = median_year_of_birth,
   ymin = min_year_of_birth,
   ymax = max_year_of_birth),
   stat = "identity", width = 0.5) +
 theme_bw()+
 theme(legend.position = "none") +
 xlab("")
```



			CDM name
Observation period ordinal	Variable name	Estimate name	synthea-covid19-10k
all	Number records	N	10,754
	Number subjects	N	10,754
	Records per person	mean (sd)	1.00(0.00)
		median [Q25 - Q75]	1 [1 - 1]
	Duration in days	mean (sd)	3,958.53 (3,430.29)
		median [Q25 - Q75]	3,326 [2,927 - 3,340]
1st	Number subjects	N	10,754
	Duration in days	mean (sd)	3,958.53 (3,430.29)
		median [Q25 - Q75]	3,326 [2,927 - 3,340]

6.4 The OmopSketch package

The OmopSketch R package aims to characterise and visualise an OMOP CDM instance to asses if it meets the necessary criteria to answer a specific clinical question and conduct a certain study. It can be very useful to conduct some of the analyses that we were conducting in the previous sections with bespoke code. For example, you can quickly summarise and visualise the observation period table as:

```
result <- summariseObservationPeriod(cdm$observation_period)
tableObservationPeriod(result = result)</pre>
```

With this table we can see that only one pbservation period is defined by person and that individuals have a median follow up of ~ 9 years.

We can also visualise how many individuals are in observation per year:

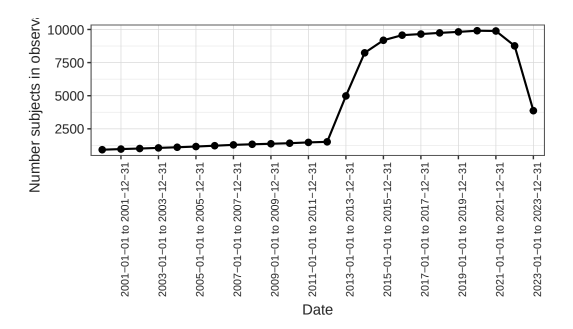
```
result <- summariseInObservation(
  cdm$observation_period,
  interval = "years",
  output = "person",
  dateRange = c("2000-01-01", "2023-12-31")
)</pre>
```

> The observation period in the cdm ends in 2023-06-14

```
plotInObservation(result = result)
```

[`]result_id` is not present in result.

`result_id` is not present in result.



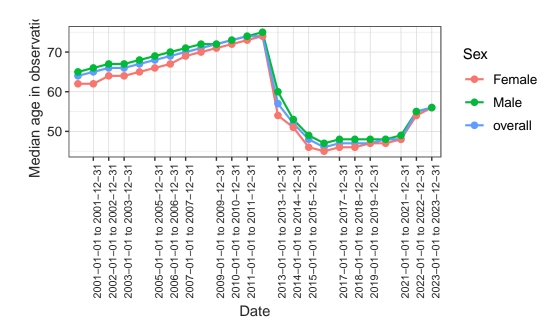
Or even the median age of the individuals in observation stratified by sex:

```
result <- summariseInObservation(
  cdm$observation_period,
  interval = "years",
  output = "age",
  sex = TRUE,
  dateRange = c("2000-01-01", "2023-12-31")
)</pre>
```

- > The observation period in the cdm ends in 2023-06-14
- i The following estimates will be computed:
- * age: median
- > Start summary of data, at 2025-10-23 17:30:23.318492
- v Summary finished, at 2025-10-23 17:30:23.482003
- i The following estimates will be computed:
- * age: median
- > Start summary of data, at 2025-10-23 17:30:25.506412

```
plotInObservation(result = result, colour = "sex")
```

```
`result_id` is not present in result.
`result_id` is not present in result.
```



The package also provides functions to characterise the clinical tables to show percentage of records in observation, domains recorded or the source vocabularies:

```
result <- summariseClinicalRecords(cdm = cdm, omopTableName = "drug_exposure")</pre>
```

- i Adding variables of interest to drug_exposure.
- i Summarising records per person in drug_exposure.

```
tableClinicalRecords(result = result)
```

			Database name
Variable name	Variable level	Estimate name	synthea-covid19-10k
drug_exposure			
Number records	-	N	337,509
Number subjects	-	N (%)	$10,754 \ (100.00\%)$
Records per person	-	Mean (SD)	31.38 (45.63)
		Median [Q25 - Q75]	22 [13 - 33]
		Range [min to max]	[1 to 1,152]
In observation	Yes	N (%)	337,509 (100.00%)
Domain	Drug	N (%)	337,509 (100.00%)
Source vocabulary	Cvx	N (%)	310,584 (92.02%)
· ·	Rxnorm	N(%)	26,925 (7.98%)
Standard concept	S	N (%)	337,509 (100.00%)
Type concept id	Pharmacy claim	N (%)	337,509 (100.00%)

Database characterisation

When you start using a database characterising it is probably a good idea to know what information is recorded in the database and explore the studies that are feasible. You can perform a database characterisation using OmopSketch as:

```
result <- databaseCharacteristics(cdm = cdm)
```

Note this process can take several hours or even days depending on the size of the database. Later you can visualise all the results in a Shiny App such as the one shown at the beginning of this chapter.

```
shinyCharacteristics(result = result, directory = getwd())
```

6.5 Disconnecting

Once we have finished our analysis we can close our connection to the database behind our cdm reference.

cdmDisconnect(cdm)

6.6 Further reading

• Alcalde-Herraiz M, Lopez-Guell K, Rowlands E, Campanile C, Burn E, Català M (2025). OmopSketch: Characterise Tables of an OMOP Common Data Model Instance. R package version 0.5.1, https://OHDSI.github.io/OmopSketch/.

7 Identifying patient characteristics

For this chapter, we'll again use our example COVID-19 dataset.

```
library(dplyr)
library(omock)
library(PatientProfiles)
library(ggplot2)
library(omopgenerics) # TODO https://github.com/OHDSI/omock/issues/189
```

```
cdm <- mockCdmFromDataset(datasetName = "synthea-covid19-10k", source = "duckdb")</pre>
```

- i Reading synthea-covid19-10k tables.
- i Adding drug_strength table.

As part of an analysis, we almost always have a need to identify certain characteristics related to the individuals in our data. These characteristics might be time-invariant (i.e. a characteristic that does not change as time passes and a person ages) or time-varying.¹

7.1 Adding specific demographics

The PatientProfiles package makes it easy for us to add demographic information to tables in the OMOP CDM. Like the CDMConnector package we've seen previously, the fact that the structure of the OMOP CDM is known allows the PatientProfiles package to abstract away some common data manipulations required to do research with patient-level data.²

Let's say we're interested in individuals' age and sex at the time of COVID-19 diagnosis. We can add these variables to the table as follows (noting that, since age is time-varying, we need to specify the date relative to which it should be calculated).

¹In some datasets, characteristics that could conceptually be considered as time-varying are encoded as time-invariant. One example of the latter is that in some cases an individual may be associated with a particular socioeconomic status or nationality that for the purposes of the data is treated as time-invariant.

²Although these manipulations can seem quite simple on the face of it, their implementation across different database platforms with different data granularity (for example, whether day of birth has been filled in for all patients or not) presents challenges that the PatientProfiles package solves for us.

```
cdm$condition_occurrence <- cdm$condition_occurrence |>
   addSex() |>
   addAge(indexDate = "condition_start_date")

cdm$condition_occurrence |>
   glimpse()
```

```
Rows: ??
Columns: 18
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1//tmp/RtmpMgnkRf/file32277317]
                           <int> 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 1~
$ condition_occurrence_id
                           <int> 2, 6, 7, 8, 8, 8, 8, 16, 16, 18, 18, 25,~
$ person_id
$ condition_concept_id
                           <int> 381316, 321042, 381316, 37311061, 437663~
$ condition_start_date
                           <date> 1986-09-08, 2021-06-23, 2021-04-
07, 202~
                           <dttm> 1986-09-08, 2021-06-23, 2021-04-
$ condition_start_datetime
07, 202~
                           <date> 1986-09-08, 2021-06-23, 2021-04-
$ condition_end_date
07, 202~
$ condition_end_datetime
                           <dttm> 1986-09-08, 2021-06-23, 2021-04-
07, 202~
$ condition_type_concept_id
                           <int> 38000175, 38000175, 38000175, 38000175, ~
$ condition_status_concept_id
                           $ stop_reason
$ provider_id
                           $ visit_occurrence_id
                           <int> 19, 55, 67, 79, 79, 79, 79, 168, 171, 19~
$ visit_detail_id
                           <int> 1000019, 1000055, 1000067, 1000079, 1000~
                           <chr> "230690007", "410429000", "230690007", "~
$ condition_source_value
                           <int> 381316, 321042, 381316, 37311061, 437663~
$ condition_source_concept_id
<chr> "Female", "Male", "Male", "Male", "Male"~
$ sex
                           <int> 57, 25, 97, 2, 2, 2, 75, 77, 57, 76, ~
$ age
```

We have now added two variables containing values for age and sex.

```
cdm$condition_occurrence |>
   glimpse()
```

Rows: ?? Columns: 18

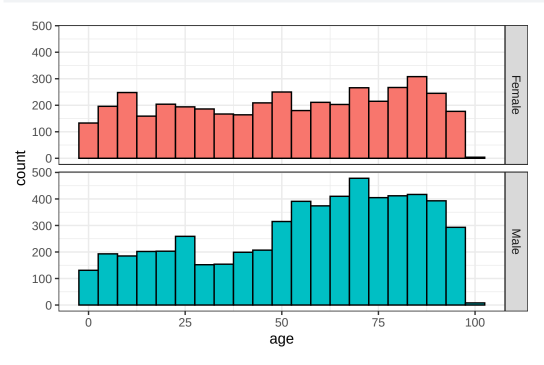
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1//tmp/RtmpMgnkRf/file32277317

```
$ condition_occurrence_id
                          <int> 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 1~
                          <int> 2, 6, 7, 8, 8, 8, 8, 16, 16, 18, 18, 25,~
$ person_id
$ condition_concept_id
                          <int> 381316, 321042, 381316, 37311061, 437663~
$ condition_start_date
                          <date> 1986-09-08, 2021-06-23, 2021-04-
07, 202~
                          <dttm> 1986-09-08, 2021-06-23, 2021-04-
$ condition_start_datetime
07, 202~
$ condition_end_date
                          <date> 1986-09-08, 2021-06-23, 2021-04-
07, 202~
                          <dttm> 1986-09-08, 2021-06-23, 2021-04-
$ condition_end_datetime
07, 202~
                          <int> 38000175, 38000175, 38000175, 38000175, ~
$ condition_type_concept_id
                          $ condition_status_concept_id
$ stop_reason
                          $ provider_id
                          $ visit_occurrence_id
                          <int> 19, 55, 67, 79, 79, 79, 79, 168, 171, 19~
$ visit_detail_id
                          <int> 1000019, 1000055, 1000067, 1000079, 1000~
                          <chr> "230690007", "410429000", "230690007", "~
$ condition_source_value
$ condition_source_concept_id
                          <int> 381316, 321042, 381316, 37311061, 437663~
<chr> "Female", "Male", "Male", "Male", "Male"~
                          <int> 57, 25, 97, 2, 2, 2, 75, 77, 57, 76, ~
$ age
```

With these now added, it is straightforward to calculate the mean age at condition start date by sex or even plot the distribution of age at diagnosis by sex.

```
cdm$condition_occurrence |>
  group_by(sex) |>
  summarise(mean_age = mean(age, na.rm=TRUE)) |>
 collect()
# A tibble: 2 x 2
  sex
         mean_age
  <chr>
            <dbl>
1 Female
             50.8
2 Male
             56.5
cdm$condition_occurrence |>
  select("person id", "age", "sex") |>
  collect() |>
 ggplot(aes(fill = sex)) +
```

```
facet_grid(sex ~ .) +
geom_histogram(aes(age), colour = "black", binwidth = 5) +
theme_bw() +
theme(legend.position = "none")
```



```
i Show query

cdm$condition_occurrence |>
   addSexQuery() |>
   show_query()

Warning: ! The following columns will be overwritten: sex

<SQL>
   SELECT
   condition_occurrence_id,
   og_002_1761240722.person_id AS person_id,
   condition_concept_id,
   condition_start_date,
   condition_start_datetime,
   condition_end_date,
   condition_end_datetime,
```

```
condition_type_concept_id,
  condition_status_concept_id,
  stop_reason,
  provider_id,
  visit occurrence id,
  visit_detail_id,
  condition_source_value,
  condition_source_concept_id,
  condition_status_source_value,
  age,
  RHS.sex AS sex
FROM og_002_1761240722
LEFT JOIN (
  SELECT
    person_id,
WHEN (gender_concept_id = 8507.0) THEN 'Male'
WHEN (gender concept id = 8532.0) THEN 'Female'
ELSE 'None'
END AS sex
  FROM person
) RHS
  ON (og 002 1761240722.person id = RHS.person id)
The difference between addSexQuery() and addSex() will be explained in the next tip
chunk.
```

7.2 Adding multiple demographics simultaneously

We've now seen individual functions from PatientProfiles that add specific patient characteristics, such as age and sex. The package also includes functions to add other characteristics, such as the number of days of prior observation in the database (rather unimaginatively named addPriorObservation()). In addition to these individual functions, the package also provides a more general function that retrieves all of these characteristics at the same time.³

```
cdm$drug_exposure <- cdm$drug_exposure |>
addDemographics(
```

³This function also provides a more time-efficient method than getting the characteristics one by one. This is because these characteristics are all derived from the OMOP CDM person and observation period tables, and so can be identified simultaneously.

```
indexDate = "drug_exposure_start_date",
   age = TRUE,
   sex = TRUE,
   priorObservation = TRUE,
   futureObservation = TRUE,
   dateOfBirth = TRUE
)

cdm$drug_exposure |>
   glimpse()
```

Rows: ?? Columns: 28 Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1//tmp/RtmpMgnkRf/file32277317 <int> 245761, 245762, 245763, 245764, 245765, 2~ \$ drug_exposure_id \$ person_id <int> 7764, 7764, 7764, 7764, 7764, 7764, 7764,~ \$ drug_concept_id <int> 40213227, 40213201, 40213198, 40213154, 4~ \$ drug_exposure_start_date <date> 2015-02-08, 2010-01-10, 2010-01-10, 2017~ \$ drug_exposure_start_datetime <dttm> 2015-02-08 22:40:04, 2010-01-10 22:40:04~ \$ drug_exposure_end_date <date> 2015-02-08, 2010-01-10, 2010-01-10, 2017~ <dttm> 2015-02-08 22:40:04, 2010-01-10 22:40:04~ \$ drug_exposure_end_datetime \$ verbatim_end_date <date> 2015-02-08, 2010-01-10, 2010-01-10, 2017~ <int> 32869, 32869, 32869, 32869, 32869, ~ \$ drug_type_concept_id \$ stop_reason \$ refills \$ quantity \$ days_supply \$ sig \$ route_concept_id \$ lot_number \$ provider_id <int> 14656, 14656, 14656, 14656, 14656, ~ \$ visit_occurrence_id <int> 80896, 80891, 80891, 80892, 80895, 80896,~ <int> 1080896, 1080891, 1080891, 1080892, 10808~ \$ visit_detail_id <chr> "113", "33", "133", "140", "140", "140", ~ \$ drug_source_value <int> 40213227, 40213201, 40213198, 40213154, 4~ \$ drug_source_concept_id \$ route_source_value \$ dose_unit_source_value <int> 71, 66, 66, 73, 72, 71, 69, 67, 65, 70, 6~ \$ age <chr> "Male", "Male", "Male", "Male", "~ \$ sex <int> 2597, 742, 742, 3339, 2968, 2597, 1855, 1~ \$ prior_observation <int> 896, 2751, 2751, 154, 525, 896, 1638, 238~ \$ future_observation <date> 1943-10-10, 1943-10-10, 1943-10-10, 1943~ \$ date_of_birth

With these characteristics added, we can now calculate mean age, prior observation (the number of days have passed since each individual's most recent observation start date), and future observation (the number of days until the individual's nearest observation end date) at drug exposure start date, stratified by sex.

• Returning a query from PatientProfiles rather than the result

In the above examples, the functions from PatientProfiles execute queries and write the results to a table in the database (either a temporary table if no name is provided when calling the function, or a permanent table). We might instead want to just get the underlying query back so that we have more control over how and when the query is executed.

```
cdm$visit_occurrence |>
   addSex() |>
   filter(sex == "Male") |>
   show_query()

<SQL>
SELECT og_004_1761240724.*
FROM og_004_1761240724
WHERE (sex = 'Male')

cdm$visit_occurrence |>
   addSex(name = "my_new_table") |>
   filter(sex == "Male") |>
   show_query()

<SQL>
```

```
SELECT test_my_new_table.*
FROM results.test_my_new_table
WHERE (sex = 'Male')
cdm$visit_occurrence |>
  addSexQuery() |>
 filter(sex == "Male") |>
  show_query()
<SQL>
SELECT q01.*
FROM (
  SELECT visit_occurrence.*, sex
  FROM visit_occurrence
  LEFT JOIN (
    SELECT
      person_id,
      CASE
WHEN (gender_concept_id = 8507.0) THEN 'Male'
WHEN (gender_concept_id = 8532.0) THEN 'Female'
ELSE 'None'
END AS sex
    FROM person
  ) RHS
    ON (visit_occurrence.person_id = RHS.person_id)
) q01
WHERE (sex = 'Male')
Query functions can be useful in some contexts where you don't want to generate multiple
```

Query functions can be useful in some contexts where you don't want to generate multiple temporary tables or do not want to lose indexes of a certain table, but they can also generate large queries that could result in low performance.

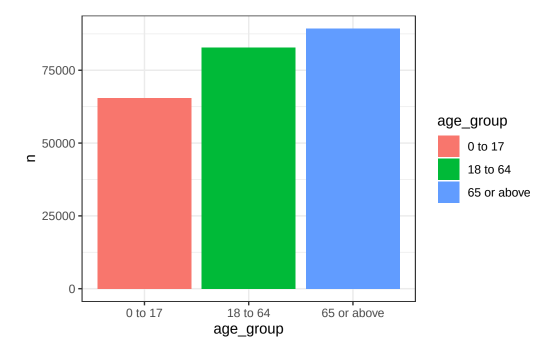
7.3 Creating categories

When adding age, either via addAge or addDemographics, we can also include an additional variable that groups individuals into age categories. These age groups must be specified in a list of vectors, each of which containing the lower and upper bounds.

```
cdm$visit_occurrence <- cdm$visit_occurrence |>
  addAge(
```

```
indexDate = "visit_start_date",
    ageGroup = list(c(0,17), c(18, 64), c(65, Inf))
)

cdm$visit_occurrence |>
    # data quality issues with our synthetic data means we have
# some negative ages so will drop these
filter(age >= 0) |>
group_by(age_group) |>
tally() |>
collect() |>
ggplot() +
geom_col(aes(x = age_group, y = n, fill = age_group)) +
theme_bw()
```

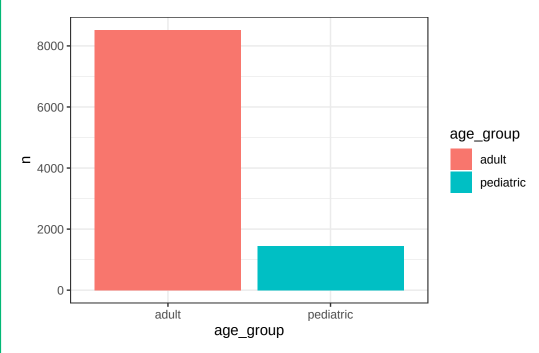


• Naming age groups

As we have seen, by default the age groups are named according to their lower and upper bounds ('0 to 17', '18 to 64', and '65 or above'). However, we can customise these labels by assigning names to the list of age groups:



Warning: ! The following columns will be overwritten: age



If you take a close look at the documentation of the function, you'll see that it also allows you to add multiple age groups and to control the name of the new column, which by default is 'age_group'.

PatientProfiles also provides a more general function for adding categories. Can you guess its name? That's right, we have addCategories() for this.

```
cdm$condition_occurrence |>
  addPriorObservation(indexDate = "condition_start_date") |>
  addCategories(
   variable = "prior_observation",
   categories = list("prior_observation_group" = list(
      c(0, 364), c(365, Inf)
   ))
  ) |>
  glimpse()
```

```
Rows: ??
Columns: 20
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1//tmp/RtmpMgnkRf/file32277317
                            <int> 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 1~
$ condition occurrence id
$ person_id
                            <int> 2, 6, 7, 8, 8, 8, 8, 16, 16, 18, 18, 25,~
$ condition concept id
                            <int> 381316, 321042, 381316, 37311061, 437663~
$ condition_start_date
                            <date> 1986-09-08, 2021-06-23, 2021-04-
07, 202~
$ condition_start_datetime
                            <dttm> 1986-09-08, 2021-06-23, 2021-04-
07, 202~
                            <date> 1986-09-08, 2021-06-23, 2021-04-
$ condition_end_date
07, 202~
$ condition_end_datetime
                            <dttm> 1986-09-08, 2021-06-23, 2021-04-
07, 202~
                            <int> 38000175, 38000175, 38000175, 38000175, ~
$ condition_type_concept_id
$ condition_status_concept_id
                            <int> 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0~
$ stop_reason
                            $ provider_id
                            <int> 19, 55, 67, 79, 79, 79, 79, 168, 171, 19~
$ visit_occurrence_id
$ visit_detail_id
                            <int> 1000019, 1000055, 1000067, 1000079, 1000~
                            <chr> "230690007", "410429000", "230690007", "~
$ condition source value
$ condition_source_concept_id
                            <int> 381316, 321042, 381316, 37311061, 437663~
<chr> "Female", "Male", "Male", "Male", "Male"~
$ sex
                            <int> 57, 25, 97, 2, 2, 2, 75, 77, 57, 76, ~
$ age
                            <int> 3437, 2898, 2842, 872, 872, 872, 872, 23~
$ prior_observation
$ prior_observation_group
                            <chr> "365 or above", "365 or above", "365 or ~
```

7.4 Adding custom variables

While PatientProfiles provides a range of functions that can help you add characteristics of interest, you may also want to add other features. Obviously, the package can't cover all the possible custom characteristics you may wish to add. However, we will see two common groups of custom variables you may want to add:

- variables derived from existing columns within the same table,
- variables obtained from other tables and joined to our table of interest.

In the first case, where we want to add a new variable derived from existing variables within our table, we'll typically use mutate() (from the dplyr package). For example, perhaps we just want to add a new variable to our observation period table that contains the year of each individual's observation period start date. This is rather straightforward.

```
cdm$observation_period <- cdm$observation_period |>
  mutate(observation_period_start_year = get_year(observation_period_start_date))
cdm$observation_period |>
  glimpse()
```

The second case is usually a more complex task, as adding a new variable involves joining to some other table following a certain logic. This table may have been created by some intermediate query that we wrote to derive the variable of interest. For example, let's say we want to add the number of condition occurrence records for each individual to the person table (remember that we saw how to calculate this in the previous chapter). To do this, we will need to perform a join between the person and condition occurrence tables (as some people might not have any records in the condition occurrence table). Here we'll create a table containing just the information we're interested in and compute it to a temporary table.

```
left_join(
    cdm$condition_occurrence |>
      group_by(person_id) |>
      count(name = "condition_occurrence_records"),
    by="person_id"
  ) |>
  select("person_id", "condition_occurrence_records") |>
  mutate(condition_occurrence_records = coalesce(condition_occurrence_records, 0)) |>
  compute()
condition_summary |>
  glimpse()
Rows: ??
Columns: 2
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1//tmp/RtmpMgnkRf/file32277317]
$ person_id
                                <int> 2, 6, 7, 8, 16, 18, 25, 36, 40, 42, 44, 4~
$ condition_occurrence_records <dbl> 1, 1, 1, 4, 2, 2, 1, 4, 1, 3, 2, 5, 1, 3,~
We can see what goes on behind the scenes by viewing the associated SQL.
cdm$person |>
  select("person_id") |>
  left_join(
    cdm$condition_occurrence |>
      group_by(person_id) |>
      count(name = "condition_occurrence_records"),
    by="person id"
  select("person_id", "condition_occurrence_records") |>
  mutate(condition_occurrence_records = coalesce(condition_occurrence_records, 0)) |>
  show_query()
<SQL>
SELECT
  person_id,
  COALESCE(condition_occurrence_records, 0.0) AS condition_occurrence_records
FROM (
  SELECT person.person_id AS person_id, condition_occurrence_records
```

condition_summary <- cdm\$person |>

select("person_id") |>

```
FROM person
LEFT JOIN (
    SELECT person_id, COUNT(*) AS condition_occurrence_records
    FROM og_002_1761240722
    GROUP BY person_id
) RHS
    ON (person.person_id = RHS.person_id)
) q01
```

Taking care with joins

When adding variables through joins we need to pay particular attention to the dimensions of the resulting table. While sometimes we may want to have additional rows added as well as new columns, this is often not desired. For example, if we have a table with one row per person, performing a left join to another table containing multiple rows per person will result in multiple rows per person in the output.

Examples where to be careful include when joining to the observation period table, as individuals can have multiple observation periods, and when working with cohorts (which are the focus of the next chapter) as individuals can also enter the same study cohort multiple times.

Just to underline how problematic joins can become if we don't take care, here we join the condition occurrence table and the drug exposure table, both of which have multiple records per person. Even with our small synthetic dataset, this produces an extremely large table. When working with real patient data, which is oftentimes much, much larger, this would be extremely problematic (and would unlikely be needed to answer any research question). In other words, don't try this at home!

```
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/RtmpMgnkRf/file32277317fee0.duckdb]
   <dbl>
1 337509
cdm$condition_occurrence |>
  select(person_id, condition_start_date) |>
  left_join(
    cdm$drug_exposure |>
      select(person_id, drug_exposure_start_date),
    by = "person_id"
  ) |>
  tally()
            SQL [?? x 1]
# Source:
# Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/RtmpMgnkRf/file32277317fee0.duckdb]
   <dbl>
1 410683
```

7.5 Disconnecting

Once we have finished our analysis we can close our connection to the database behind our cdm reference.

```
cdmDisconnect(cdm)
```

7.6 Further reading

• Català M, Guo Y, Du M, Lopez-Guell K, Burn E, Mercade-Besora N (2025). Patient-Profiles: Identify Characteristics of Patients in the OMOP Common Data Model. R package version 1.4.3, https://darwin-eu.github.io/PatientProfiles/.

8 Adding cohorts to the CDM

8.1 What is a cohort?

When conducting research using the OMOP Common Data Model (CDM), we often aim to identify groups of individuals who share specific characteristics. The inclusion criteria for these groups can range from relatively simple (e.g. people diagnosed with asthma) to highly complex (e.g. adults diagnosed with asthma who had at least one year of prior observation in the database before their diagnosis, no prior history of chronic obstructive pulmonary disease (COPD), and no history of using short-acting beta-antagonists).

The groups of individuals we identify are called cohorts. In the OMOP CDM, cohorts are represented using a specific structure: a cohort table with four required fields:

- 1) Cohort definition id a unique identifier for each cohort (multiple cohorts can be defined in the same cohort table).
- 2) Subject id a foreign key linking the subject in the cohort to the person table.
- 3) Cohort start date date indicating the beginning of the cohort record.
- 4) Cohort end date date indicating the end of the cohort record.

Individuals must be defined in the person table and must be under observation (i.e. have an ongoing record in the observation period table) to be part of a cohort. Individuals can enter a cohort multiple times, but the time periods in which they are in the cohort cannot overlap.

It is beyond the scope of this book to describe all the different ways cohorts could be created, however in this chapter we provide a summary of some of the key building blocks for cohort creation. Cohort-building pipelines can be created following these principles to create a wide range of study cohorts.

8.2 Set up

We'll use the same Covid-19 synthetic dataset that we used before for demonstrating how cohorts can be constructed.

```
library(omock)
library(CohortConstructor)
library(CohortCharacteristics)
library(dplyr)

cdm <- mockCdmFromDataset(datasetName = "synthea-covid19-10k", source = "duckdb")</pre>
```

8.3 General concept based cohort

Often study cohorts will be based around a specific clinical event identified by some set of clinical codes. Here, for example, we use the CohortConstructor package to create a cohort of people with Covid-19. For this we are identifying any clinical records with the code 37311061.

- # Source: table<results.test_covid> [?? x 4]
- # Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1//tmp/Rtmp2eznHK/file32a472/ cohort_definition_id subject_id cohort_start_date cohort_end_date

	<int></int>	<int></int>	<date></date>	<date></date>
1	1	1635	2020-07-14	2020-08-08
2	1	2000	2020-11-15	2020-12-10
3	1	4649	2021-09-08	2021-10-12
4	1	9822	2021-02-06	2021-02-25
5	1	10394	2020-11-29	2020-12-31
6	1	5670	2021-02-20	2021-03-02
7	1	5823	2021-01-11	2021-01-26
8	1	5910	2020-10-11	2020-11-15
9	1	10547	2020-07-29	2020-08-11
10	1	2386	2021-01-29	2021-02-24

i more rows

i Name consistency

Note that the name argument determines the name of the permanent table written in the database and as we have seen before, we have to be consistent assigning the tables to the cdm object. That's why we used name = "covid" and then we were able to assign it to cdm\$covid. Otherwise, see this failing example:

```
cdm$not_covid <- conceptCohort(cdm = cdm,</pre>
                                conceptSet = list("covid" = 37311061),
                                name = "covid")
Warning: ! `codelist` casted to integers.
i Subsetting table condition_occurrence using 1 concept with domain: condition.
i Combining tables.
i Creating cohort attributes.
i Applying cohort requirements.
i Merging overlapping records.
v Cohort covid created.
Error in `[[<-`:</pre>
x You can't assign a table named covid to not_covid.
i You can change the name using compute:
cdm[['not_covid']] <- yourObject |>
  dplyr::compute(name = 'not_covid')
i You can also change the name using the `name` argument in your function:
  `name = 'not_covid'`.
```

• Finding appropriate codes

In defining the cohorts above, we have needed to provide concept IDs for our outcomes of interest. But where do these come from?

We can search for codes of interest using the CodelistGenerator package. This can be done using a text search with the function getCandidateCodes(). For example, we can have found the code we used above (and many others) like so:

Time taken: 0 minutes and 1 seconds

v 37 candidate concepts identified

```
# A tibble: 37 x 6
   concept id found from
                             concept_name domain_id vocabulary_id standard_concept
        <int> <chr>
                             <chr>
                                           <chr>
                                                                     <chr>
                                                      <chr>
     37310254 From initia~ Otitis medi~ Condition SNOMED
                                                                     S
     37310287 From initia~ Myocarditis~ Condition SNOMED
                                                                     S
       703445 From initia~ Low risk ca~ Condition SNOMED
       756039 From initia~ Respiratory~ Condition OMOP Extensi~ S
       705076 From initia~ Post-acute ~ Condition OMOP Extensi~
     37311061 From initia~ COVID-19
                                           Condition SNOMED
 7
      3655975 From initia~ Sepsis due ~ Condition SNOMED
                                                                     S
      3655977 From initia~ Rhabdomyoly~ Condition SNOMED
                                                                     S
 8
      3656667 From initia~ Cardiomyopa~ Condition SNOMED
                                                                     S
      3661885 From initia~ Fever cause~ Condition SNOMED
10
                                                                     S
# i 27 more rows
We can also do automated searches that make use of the hierarchies in the vocabularies.
Here, for example, we find the code for the drug ingredient Acetaminophen and all of its
descendants.
codes <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")</pre>
codes
- 161_acetaminophen (25747 codes)
Note that in practice clinical expertise is vital in the identification of appropriate codes
so as to decide which the codes are in line with the clinical idea at hand. Additionally,
not all codes may be used in the database. To check for this, we can use the function
subsetToCodesInUse().
```

We can see that as well as having the cohort entries above, our cohort table is associated with several attributes.

cdm = cdm)

usedCodes <- subsetToCodesInUse(list("acet" = codes[[1]]),</pre>

First, we can see the settings associated with cohort.

```
settings(cdm$covid) |>
glimpse()
```

In settings, we can see the cohort name that by default is the name of the codelist used, in this case 'covid' as we used conceptSet = list(covid = 37311061). Also, the cdm and vocabulary versions are recorded in the settings by the CohortConstructor package.

Second, we can get counts of each cohort.

```
cohortCount(cdm$covid) |>
glimpse()
```

```
Rows: 1
Columns: 3
$ cohort_definition_id <int> 1
$ number_records <int> 964
$ number_subjects <int> 964
```

Where you can see the number of records and number of subjects for each cohort. In this case, there are no multiple records per subject.

Attrition can also be retrieved from any cohort.

```
attrition(cdm$covid) |>
  glimpse()
```

And finally, you can extract the codelists used to create a cohort table:

```
codelist <- cohortCodelist(cdm$covid, cohortId = 1)
codelist</pre>
```

- -- 1 codelist -----
- covid (1 codes)

codelist\$covid

[1] 37311061

Note that in this case, we had to provide the cohortId of the cohort of interest.

All these attributes can be retrieved because it is a cohort_table object, a class on top of the usual cdm_table class that we have seen before:

class(cdm\$covid)

As we will see below, these attributes of the cohorts become particularly useful as we apply further restrictions on our cohort.

Behind the scenes

All these attributes that we have seen are part of the attributes of the cohort_table object and are used by these utility functions:

names(attributes(cdm\$covid))

- [1] "names" "class" "tbl_source" "tbl_name"
- [5] "cohort_set" "cohort_attrition" "cohort_codelist" "cdm_reference"

In particular, the cohort_set (contains the settings() source), cohort_attrition

(contains the source for <code>cohortCount()</code> and <code>attrition()</code>) and <code>cohort_codelist</code> (contains the source for <code>cohortCodelist()</code>) attributes are the ones of interest. For database backends, these attributes are stored directly in the database so that they persist for when we read them again. Note that although it may appear that there is only one table—<code>cdm\$covid</code>—in fact, four tables are written to the database:

We do not have to worry about the attributes and the naming of the tables as CohortConstructor, CDMConnector and omopgenerics take care of that and if we create the cohorts with functions such as conceptCohort() then we will be able to read them back with the cohortTables argument of cdmFromCon() or the readSourceTable() function and all the attributes will be in place.

8.4 Applying inclusion criteria

8.4.1 Only include first cohort entry per person

Let's say we first want to restrict our cohort to only include the first record for each person. This can be done by using the funtion requireIsFirstEntry():

```
cdm$covid <- cdm$covid |>
    requireIsFirstEntry()
```

8.4.2 Restrict to study period

Then we are only interested in records from January 1st, 2020 onwards.

```
cdm$covid <- cdm$covid |>
    requireInDateRange(dateRange = c(as.Date("2020-01-01"), NA))
```

8.4.3 Applying demographic inclusion criteria

Finally, we want to restrict our population of interest to only adult males under 65 years old. We can do that with the requireDemographics() function.

```
cdm$covid <- cdm$covid |>
  requireDemographics(ageRange = c(18, 64), sex = "Male")
```

i Similarity of naming with PatientProfiles

Note that all these require*() functions that come from the CohortConstructor package use functionalities from PatientProfiles and the naming is consistent. For example, requireDemographics() uses addDemographics(), requirePriorObservation() uses addPriorObservation(), and so on...

8.4.4 Applying cohort-based inclusion criteria

In addition to demographic requirements, we may also want to use another cohort as part of the inclusion or exclusion criteria. For example, we might exclude anyone with a history of cardiac conditions prior to their COVID-19 cohort entry.

We can first generate a cohort table with records of myocardial infarction.

```
cdm$cardiac <- conceptCohort(
  cdm = cdm,
  conceptSet = list("myocaridal_infarction" = c(317576L, 313217L, 321042L, 4329847L)),
  name = "cardiac"
)
cdm$cardiac</pre>
```

```
# Source: table<results.test_cardiac> [?? x 4]
```

Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-azure:R 4.4.1//tmp/Rtmp2eznHK/file32a472/ cohort_definition_id subject_id cohort_start_date cohort_end_date

	<int></int>	<int></int>	<date></date>	<date></date>
1	1	873	2015-08-28	2015-08-28
2	1	1022	2012-01-11	2012-01-11
3	1	1080	1996-12-19	1996-12-19

```
1737 1961-11-26
                                                         1961-11-26
4
                       1
 5
                       1
                                1914 2007-07-03
                                                         2007-07-03
6
                       1
                                1933 2021-08-29
                                                         2021-08-29
7
                                2721 2006-09-23
                       1
                                                         2006-09-23
8
                       1
                                4602 2018-05-09
                                                         2018-05-09
9
                       1
                                5007 2013-11-16
                                                         2013-11-16
10
                       1
                                5232 1983-05-16
                                                         1983-05-16
# i more rows
```

Now we can apply the inclusion criteria that individuals have no records of myocardical infarction prior to their Covid-19 cohort entry.

Note that if we had wanted to require that individuals did have a history of a cardiac condition, we would instead have set intersections = c(1, Inf) above.

i Use requireConceptIntersect

We could have applied the exact same inclusion criteria using the requireConceptIntersect() function, this code would be equivalent:

```
cdm$covid <- cdm$covid |>
  requireConceptIntersect(
    conceptSet = list("myocaridal_infarction" = c(317576L, 313217L, 321042L, 4329847L)),
  indexDate = "cohort_start_date",
  window = c(-Inf, -1),
  intersections = 0
)
```

In fact, this approach is generally more efficient unless we plan to reuse the myocar-dial_infarction cohort for another inclusion criteria or analysis. Note, however, that intersecting with the cohort table is more flexible, as it allows for more complex inclusion and exclusion logic. However, you need to be careful with the order of criteria, for example if we restricted the myocardial_infarction cohort to a specific time period before performing the intersection, we would need to avoid applying a separate inclusion criterion based on that same time span.

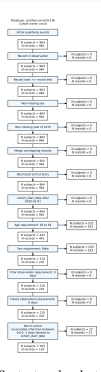
8.5 Cohort attributes

Using the require*() functions, the cohort attributes have been updated to reflect the applied inclusion criteria.

```
settings(cdm$covid) |>
 glimpse()
Rows: 1
Columns: 8
$ cohort_definition_id
                      <int> 1
$ cohort_name
                      <chr> "covid"
                      <chr> "5.3"
$ cdm_version
$ vocabulary_version
                      <chr> "v5.0 22-JUN-22"
                      <chr> "18_64"
$ age_range
                      <chr> "Male"
$ sex
$ min_prior_observation
                      <dbl> 0
$ min_future_observation <dbl> 0
cohortCount(cdm$covid) |>
 glimpse()
Rows: 1
Columns: 3
$ cohort_definition_id <int> 1
$ number_records
                    <int> 193
$ number_subjects
                    <int> 193
attrition(cdm$covid) |>
 glimpse()
Rows: 13
Columns: 7
$ cohort_definition_id <int> 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1
$ number_records
                    $ number_subjects
$ reason_id
                    <int> 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13
$ reason
                    <chr> "Initial qualifying events", "Record in observati~
                    <int> 0, 0, 0, 0, 0, 0, 0, 521, 233, 0, 0, 17
$ excluded_records
$ excluded_subjects
                    <int> 0, 0, 0, 0, 0, 0, 0, 521, 233, 0, 0, 17
```

We can visualize the attrition with the CohortCharacteristics package. We can first extract it with summariseCohortAttrition() and then plotCohortAttrition to better view the impact of applying each inclusion criteria:

```
attrition_summary <- summariseCohortAttrition(cohort = cdm$covid)
plotCohortAttrition(result = attrition_summary, type = 'png')</pre>
```



Note that the conceptCohort() first step leads to several rows in the attrition table, whereas any other require*() function always adds just one record of attrition.

- i Combining tables.
- i Creating cohort attributes.
- i Applying cohort requirements.

```
i Merging overlapping records.
v Cohort my_cohort created.
settings(cdm$my cohort)
# A tibble: 2 x 4
  cohort_definition_id cohort_name cdm_version vocabulary_version
                <int> <chr>
                             <chr> <chr>
                                 5.3
1
                    1 concept_1
                                            v5.0 22-JUN-22
2
                    2 concept_2
                                 5.3
                                             v5.0 22-JUN-22
But maybe we are interested in renaming a cohort (e.g., after applying the inclusion
criteria). We can do that with the renameCohort() utility function:
cdm$my_cohort <- cdm$my_cohort |>
 requirePriorObservation(minPriorObservation = 365, cohortId = 1) |>
 renameCohort(cohortId = 1, newCohortName = "concept_1_365obs")
settings(cdm$my_cohort)
# A tibble: 2 x 5
  <int> <chr>
                                      <chr>
                                                  <chr>
                                                  v5.0 22-JUN-22
1
                    1 concept_1_365obs 5.3
                    2 concept_2 5.3
                                                 v5.0 22-JUN-22
# i 1 more variable: min_prior_observation <dbl>
Note that for arguments such as cohortId, targetCohortId, etc., we are able to use the
name of the cohort of interest. See for example:
cdm$my cohort <- cdm$my cohort |>
 requireSex(sex = "Female", cohortId = "concept 2") |>
 renameCohort(cohortId = "concept_2", newCohortName = "concept_2_female")
settings(cdm$my_cohort)
# A tibble: 2 x 6
  cohort_definition_id cohort_name
                                      cdm_version vocabulary_version
                <int> <chr>
                                      <chr>
                                                  <chr>
1
                    1 concept_1_365obs 5.3
                                                  v5.0 22-JUN-22
2
                    2 concept_2_female 5.3
                                                 v5.0 22-JUN-22
# i 2 more variables: min_prior_observation <dbl>, sex <chr>
```

This functionality also applies to other packages, such as CohortCharacteristics, Patient-Profiles and DrugUtilisation. In some cases, it is useful to add the cohort_name as a column to not have to check manually the equivalence between *cohort definition id* and *cohort name*. This can be done using the PatientProfiles utility function addCohortName():

```
library(PatientProfiles)
cdm$my_cohort |>
 addCohortName() |>
 glimpse()
Rows: ??
Columns: 5
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/Rtmp2eznHK/file32a4729aaa6b.duckdb]
<int> 71, 82, 134, 151, 153, 183, 194, 196, 200, 215,
$ subject_id
$ cohort_start_date
                     <date> 1936-01-24, 1997-08-04, 2001-01-09, 2022-
01-13, ~
$ cohort_end_date
                     <date> 1936-01-24, 1997-08-04, 2001-01-09, 2022-
01-13, ~
                     <chr> "concept_2_female", "concept_2_female", "concept_~
$ cohort_name
Also other utility functions that can be useful are those provided by omopgenerics:
library(omopgenerics)
getCohortId(cohort = cdm$my_cohort, cohortName = "concept_2_female")
concept_2_female
              2
getCohortId(cohort = cdm$my_cohort)
concept_1_365obs concept_2_female
                              2
getCohortName(cohort = cdm$my cohort, cohortId = 1)
"concept_1_365obs"
getCohortName(cohort = cdm$my_cohort, cohortId = c(2, 1))
```

8.6 Disconnecting

Once we have finished our analysis we can close our connection to the database behind our cdm reference.

```
cdmDisconnect(cdm)
```

8.7 Further reading

- Cohort tables
- Burn E, Català M, Mercade-Besora N, Alcalde-Herraiz M, Du M, Guo Y, Chen X, Lopez-Guell K, Rowlands E (2025). *CohortConstructor: Build and Manipulate Study Cohorts Using a Common Data Model.* R package version 0.5.0, https://ohdsi.github.io/CohortConstructor/.

9 Working with cohorts

9.1 Cohort intersections

When conducting research, it is often necessary to study patients who meet multiple clinical criteria simultaneously. For example, we may be interested in analysing outcomes among patients who have both diabetes and hypertension. Using the OMOP CDM, this typically involves first creating two separate cohorts: one for patients with diabetes and another for those with hypertension. To identify patients who meet both conditions, the next step is to compute the intersection of these cohorts. This ensures that the final study population includes only individuals who satisfy all specified criteria. Hence, finding cohort intersections is a common and essential task when working with the OMOP CDM, enabling researchers to define precise target populations that align with their research objectives.

Depending on the research question, the definition of a cohort intersection may vary. For instance, you might require patients to have a diagnosis of hypertension before developing diabetes, or that both diagnoses occur within a specific time window. These additional temporal or clinical criteria can make cohort intersection more complex. The PatientProfiles R package addresses these challenges by providing a suite of flexible functions to support the calculation of cohort intersections under various scenarios.

9.2 Intersection between two cohorts

Suppose we are interested in studying patients with gastrointestinal (GI) bleeding who have also been exposed to acetaminophen. First, we would create two separate cohorts: one for patients with GI bleeding and another for patients with exposure to acetaminophen. Below is an example of the code used to create these cohorts within the GiBleed synthetic database. A characterisation of this dataset can be found here.

```
library(omock)
library(dplyr)
library(PatientProfiles)
library(CohortConstructor)
library(omopgenerics)
```

```
cdm <- mockCdmFromDataset(datasetName = "GiBleed", source = "duckdb")</pre>
# gi bleed contains all records of GiBbleed, end date is 30 days after index
# date
cdm$gi_bleed <- conceptCohort(</pre>
 cdm = cdm,
 conceptSet = list("gi_bleed" = 192671L),
 name = "gi_bleed",
 exit = "event_start_date"
) |>
 padCohortEnd(days = 30)
# drugs cohort contains records of acetaminophen using start and end dates of
# the drug records and collapsing records separated by less than 30 days
cdm$drugs <- conceptCohort(</pre>
 cdm = cdm,
 conceptSet = list("acetaminophen" = c(
    1125315L, 1127078L, 1127433L, 40229134L, 40231925L, 40162522L, 19133768L
 )),
 name = "drugs",
 exit = "event_end_date"
) |>
 collapseCohorts(gap = 30)
```

The PatientProfiles package contains functions to obtain the intersection flag, count, date, or number of days between cohorts.

9.2.1 Flag

To get a binary indicator showing the presence of an intersection between the cohorts within a given time window, we can use addCohortIntersectFlag().

```
x <- cdm$gi_bleed |>
  addCohortIntersectFlag(
    targetCohortTable = "drugs",
    window = list("prior" = c(-Inf, -1), "index" = c(0, 0), "post" = c(1, Inf))
)

x |>
  summarise(
  acetaminophen_prior = sum(acetaminophen_prior, na.rm = TRUE),
```

```
acetaminophen_index = sum(acetaminophen_index, na.rm = TRUE),
   acetaminophen_post = sum(acetaminophen_post, na.rm = TRUE)
) |>
collect()
```

i Window naming

\$ acetaminophen 1 to inf

Windows work very similarly to age groups that we have seen before. If a name is not provided, an automatic name will be obtained from the values of the window limits:

```
cdm$gi_bleed |>
 addCohortIntersectFlag(
  targetCohortTable = "drugs",
  window = list(c(-Inf, -1), c(0, 0), c(1, Inf))
 ) |>
 glimpse()
Rows: ??
Columns: 7
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/Rtmp5ribhQ/file330f6115a898.duckdb]
$ cohort_definition_id
                   $ subject_id
                   <int> 1733, 507, 2777, 4210, 2836, 3195, 3958, 149,~
$ cohort_start_date
                   <date> 2016-02-01, 1977-10-29, 1999-12-
04, 2017-02-~
$ cohort end date
                   <date> 2016-03-02, 1977-11-28, 2000-01-
03, 2017-03-~
```

Note that to avoid conflicts with column naming, all names will be lower case, spaces are not allowed, and the - symbol for negative values is replaced by m. That's why it is usually nice to provide your own custom names:

<dbl> 0, 1, 1, 0, 0, 1, 1, 1, 0, 0, 0, 0, 0, 0, 1, ~

```
cdm$gi_bleed |>
 addCohortIntersectFlag(
  targetCohortTable = "drugs",
  window = list("prior" = c(-Inf, -1), "index" = c(0, 0), "post" = c(1, Inf))
 ) |>
 glimpse()
Rows: ??
Columns: 7
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/Rtmp5ribhQ/file330f6115a898.duckdb]
$ subject id
                <int> 1733, 507, 2777, 4210, 2836, 3195, 3958, 149, 301~
$ cohort_start_date
                <date> 2016-02-01, 1977-10-29, 1999-12-04, 2017-
02-17, ~
$ cohort end date
                <date> 2016-03-02, 1977-11-28, 2000-01-03, 2017-
03-19, ~
$ acetaminophen_post
                <dbl> 0, 1, 1, 0, 0, 1, 1, 1, 0, 0, 0, 0, 0, 0, 1, 0, 0~
```

• New column naming

By default, the name of new columns is '{cohort_name}_{window_name}' as we have seen in the prior examples. In some cases, you only have one cohort or one window and you might want to rename the column as you please. In that case, you can use the nameStyle argument to change the new naming of the columns:

```
$ subject id
                     <int> 1733, 507, 2777, 4210, 2836, 3195, 3958, 149, 301~
$ cohort_start_date
                     <date> 2016-02-01, 1977-10-29, 1999-12-04, 2017-
02-17, ~
$ cohort_end_date
                     <date> 2016-03-02, 1977-11-28, 2000-01-03, 2017-
03-19, ~
$ my_column_post
                     <dbl> 0, 1, 1, 0, 0, 1, 1, 1, 0, 0, 0, 0, 0, 0, 1, 0, 0~
$ my_column_prior
                     $ my_column_index
                     If multiple windows are provided but '{window name}' is not included in nameStyle,
then an error will prompt:
cdm$gi_bleed |>
 addCohortIntersectFlag(
   targetCohortTable = "drugs",
   window = list("prior" = c(-Inf, -1), "index" = c(0, 0), "post" = c(1, Inf)),
   nameStyle = "my_new_column"
 ) |>
 glimpse()
Error in `.addIntersect()`:
! The following elements are not present in nameStyle:
* {window_name}
Many functions that create new columns (usually functions that start with add*()) have
this nameStyle functionality that allows you to control the naming of the new columns
created.
```

9.2.2 Count

To get the count of occurrences of intersection between two cohorts, we can use addCohortIntersectCount():

```
x <- cdm$gi_bleed |>
  addCohortIntersectCount(
    targetCohortTable = "drugs",
    window = list("prior" = c(-Inf, -1), "index" = c(0, 0), "post" = c(1, Inf)),
)
x |>
  summarise(
```

```
sum_prior = sum(acetaminophen_prior, na.rm = TRUE),
mean_prior = mean(acetaminophen_prior, na.rm = TRUE),
sum_index = sum(acetaminophen_index, na.rm = TRUE),
mean_index = mean(acetaminophen_index, na.rm = TRUE),
sum_post = sum(acetaminophen_post, na.rm = TRUE),
mean_post = mean(acetaminophen_post, na.rm = TRUE)
) |>
collect()
```

i Handling the observation period

Note that **only intersections in the current observation period are considered**. The count and flag new columns can also have NA values meaning that the individual was not in observation in that window of interest. If we see individual 2070, it has 3748 days of future observation:

```
cdm$gi bleed |>
  filter(subject_id == 2070) |>
  addFutureObservation() |>
  glimpse()
Rows: ??
Columns: 5
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/Rtmp5ribhQ/file330f6115a898.duckdb]
$ cohort_definition_id <int> 1
$ subject_id
                       <int> 2070
$ cohort start date
                       <date> 2008-08-15
$ cohort_end_date
                       <date> 2008-09-14
$ future observation
                       <int> 3748
Now we will perform the intersect with the following window of interest: c(2000, 3000),
c(3000, 4000), c(4000, 5000).
```

```
cdm$gi_bleed |>
  filter(subject id == 2070) |>
 addCohortIntersectCount(
    targetCohortTable = "drugs",
    window = list(c(2000, 3000), c(3000, 4000), c(4000, 5000)),
  ) |>
glimpse()
Rows: ??
Columns: 7
Database: DuckDB 1.4.1 [unknown@Linux 6.11.0-1018-
azure:R 4.4.1//tmp/Rtmp5ribhQ/file330f6115a898.duckdb]
$ cohort_definition_id
                             <int> 1
$ subject_id
                             <int> 2070
$ cohort start date
                             <date> 2008-08-15
$ cohort end date
                             <date> 2008-09-14
$ acetaminophen_2000_to_3000 <dbl> 0
$ acetaminophen_3000_to_4000 <dbl> 0
$ acetaminophen_4000_to_5000 <dbl> NA
```

See that for the window 2000 to 3000, where the individual is still in observation, a 0 is reported. The same happens for the window 3000 to 4000 even if the individual does not have complete observation in the window. But for the last window, as the individual is not in observation at any point of the window, NA is reported.

9.2.3 Date and times

To get the date of the intersection with a cohort within a given time window, we can use addCohortIntersectDate(). To get the number of days between the index date and intersection, we can use addCohortIntersectDays().

Both functions allow the order argument to specify which value to return:

- first returns the first date/days that satisfy the window
- last returns the last date/days that satisfy the window

```
x <- cdm$gi_bleed |>
  addCohortIntersectDate(
  targetCohortTable = "drugs",
  window = list("post" = c(1, Inf)),
  order = "first"
```

```
x |>
  summarise(acetaminophen_post = median(acetaminophen_post, na.rm = TRUE)) |>
  collect()
# A tibble: 1 x 1
  acetaminophen post
  <dttm>
1 2004-02-01 00:00:00
x <- cdm$gi_bleed |>
  addCohortIntersectDays(
    targetCohortTable = "drugs",
    window = list("prior" = c(-Inf, -1)),
    order = "last"
  )
x |>
  summarise(acetaminophen_prior = median(acetaminophen_prior, na.rm = TRUE)) |>
  collect()
# A tibble: 1 x 1
  acetaminophen_prior
                <dbl>
1
                -3159
```

Note that for the window in the future, we used order = "first" and for the window in the past, we used order = "last" as in both cases we wanted to get the intersection that was closer to the index date. Individuals with no intersection will have NA on the newly created columns.

9.3 Intersection between a cohort and tables with patient data

Sometimes we might want to get the intersection between a cohort and another OMOP table. PatientProfiles also includes several addTableIntersect* functions to obtain intersection flags, counts, days, or dates between a cohort and clinical tables.

For example, if we want to get the number of general practitioner (GP) visits for individuals in the cohort, we can use the visit_occurrence table:

9.4 Disconnecting

Once we have finished our analysis we can close our connection to the database behind our cdm reference.

```
cdmDisconnect(cdm)
```

9.5 Further reading

Full details on the intersection functions in PatientProfiles can be found on the package website: $\frac{https:}{darwin-eu.github.io} = \frac{PatientProfiles}{PatientProfiles}.$

Final remarks

Tidy R programming with the OMOP Common Data Model aims to (1) explain the main principles for working with databases from R and (2) how to apply these principles and use them with the OMOP CDM. Hopefully, after reading this book, you can understand how the dplyr and dbplyr packages interact with the databases, in particular with data formatted to the OMOP CDM; how the cdm_reference object can be used to extract and identify your population of interest; and add the desired features to your dataset. Note that in this book we always worked with relatively small synthetic data with unrealistic performance. Any analysis conducted with real-world data and big databases will take more time, that's why we would always recommend you test your code against synthetic data or subsets of your data to ensure good performance. Be careful, especially while writing custom code using join functions that can create some ugly SQL.

Learning more

If you find this book useful then joining the **Tidy R in OMOP** OHDSI working group will likely be of interest. Building on many of the concepts and tools seen in this book, the Tidy R in OMOP OHDSI working group aims to (1) create and promote a unified set of resources to guide Tidy R development in OMOP and support the OHDSI community, and (2) establish an overview of available packages relevant to Tidy R programming in OMOP (tidyverse style packages). If you are interested in joining the working group then please email Martí Català or Raivo Kolde.

Support us

We encourage you to support this work by either citing the book in your papers or documentation, recommending it to your colleagues, or starring the GitHub repository, or simply letting us know how it helped you. Most importantly, please **use it** in research that results in patient benefit.