RImmPort: Quick Start Guide

Ravi Shankar 2016-01-27

1 Introduction

ImmPort study data is available for download in two formats currently: MySQL and TSV (Tab) formats. The RImmPort workflow is as follows: 1) MySQL formatted study data: User downloads one or more studies in MySQL zip files. Unzips the files. Loads local database instance. Connects to the database. Sets the ImmPort data source to the connection handle. Invokes RImmPort functions. 2) Tab: User downloads one or more studies in Tab format. Passes the folder where the zip files are located to an RImmPort function that builds SQLite database. Connects to the database. Sets the ImmPort data source to the connection handle. Invokes RImmPort functions.

User downloads study data of interest from the ImmPort website (http://www.immport.org) **. Depending on the file format MySQL or Tab the data is loaded into a local MySQL and SQLite database respectively. The user installs the RImmPort package, loads the RImmPort library, connects to the ImmPort database, and calls RImmPort methods to load study data from the database into R. Please refer to RImmPort_Article.pdf for a detailed discussion on RImmPort.

** User need to regsiter to the ImmPort website for downloading the datasets.

2 Initial Steps

- Download MySQL or Tab formatted data of studies of interest from the ImmPort website
- If working with MySQL-format, load the data in to a local MySQL database
- Install and load RImmPort package, and other required packages.

3 Load the RImmPort library

```
library(RImmPort)
library(DBI)
library(sqldf)

## Loading required package: gsubfn

## Loading required package: proto

## Loading required package: RSQLite

library(plyr)
```

- 4 Setup ImmPort data source that all RImmPort functions will use
- 4.1 Option 1: ImmPort MySQL database
- 4.1.1 Download zip files of ImmPort study data in MySQL format. e.g. SDY139' and 'SDY208'
- 4.1.2 Load the data into a local MySQL database
- 4.1.3 Connect to the ImmPort MySQL database.

4.1.4 Set the data source as the ImmPort MySQL database.

```
setImmPortDataSource(mysql_conn)
```

- 4.2 Option 2: ImmPort SQLite database
- 4.2.1 Download zip files of ImmPort data, in Tab format. e.g. SDY139' and SDY208'

```
# get the directory where ImmPort sample data is stored in the directory structure of RImmPort package
studies_dir <- system.file("extdata", "ImmPortStudies", package = "RImmPort")

# set tab_dir to the folder where the zip files are located
tab_dir <- file.path(studies_dir, "Tab")
list.files(tab_dir)</pre>
```

[1] "SDY139-DR16_Tab.zip" "SDY208-DR16_Tab.zip"

[1] "ImmPort.sqlite"

4.2.2 Build a local SQLite ImmPort database instance.

```
# set db_dir to the folder where the database file 'ImmPort.sqlite' should be stored
db_dir <- file.path(studies_dir, "Db")

# build a new ImmPort SQLite database with the data in the downloaded zip files
buildNewSqliteDb(tab_dir, db_dir)

list.files(db_dir)</pre>
```

4.2.3 Connect to the ImmPort SQLite database

```
# connect to the private instance of the ImmPort database
sqlite_conn <- dbConnect(SQLite(), dbname=file.path(db_dir, "ImmPort.sqlite"))</pre>
```

4.2.4 Set the data source to the ImmPort SQLite DB

```
setImmPortDataSource(sqlite_conn)
```

- 5 NOTE: In rest of the script, all RImmPort functions will use the SQLite ImmPort database as the data source.
- 6 Get all study ids

```
getListOfStudies()
## [1] "SDY139" "SDY208"
```

7 Get all data of a specific study

The getStudyFromDatabase queries the ImmPort database for the entire dataset of a specific study, and instantiates the Study reference class with that data.

```
?Study
# load all the data of study: `SDY139`
study_id <- 'SDY139'
sdy139 <- getStudy(study_id)</pre>
## loading Study ID = SDY139
## loading Demographics data....done
## loading Subject Visits data....done
## loading Concomitant Medications data....done
## loading Exposure data....done
## loading Substance Use data....done
## loading Adverse Events data....done
## loading Protocol Deviations data....done
## loading Medical History data....done
## loading Associated Persons Medical History data....done
## loading Laboratory Test Results data....done
## loading Physical Examination data....done
## loading Vital Signs data....done
## loading Questionnaires data....done
## loading Findings About data....done
```

```
## loading Skin Response data....done
## loading Genetics Findings data....loading HLA Typing Results data....done
## loading Array Results data....done
## done
## loading Protein Quantification data....loading ELISA Results data....done
## loading MBAA Results data....done
## loading Cellular Quantification data....loading FCS Results data....done
## loading ELISPOT Results data....done
## loading Nucleic Acid Quantification data....loading PCR Results data....done
## loading Titer Assay Results data....loading HAI Assay Results data....done
## loading Neut. Ab Titer Results data....done
## loading TrialArms data....done
## loading TrialInclusionExclusionCriteria data....done
## loading TrialSummary data.... SDY139 done
## done loading Study ID = SDY139
# access Demographics data of SDY139
dm_df <- sdy139$special_purpose$dm_l$dm_df</pre>
head(dm df)
                                                      SPECIES STRAIN SBSTRAIN
##
     STUDYID DOMAIN
                      USUBJID AGE
                                    AGEU
                                             SEX
                DM SUB118053 2 Months Unknown Mus musculus BALB/c
## 1 SDY139
## 2 SDY139
                DM SUB118054 2 Months Unknown Mus musculus BALB/c
                                                                           NA
## 3 SDY139
                DM SUB118055 2 Months Unknown Mus musculus BALB/c
                                                                           NA
## 4 SDY139
                DM SUB118056 2 Months Unknown Mus musculus BALB/c
                                                                           NA
## 5 SDY139
                DM SUB118057 2 Months Unknown Mus musculus BALB/c
                                                                           NA
                DM SUB118058 2 Months Unknown Mus musculus BALB/c
## 6 SDY139
                                                                           NA
      ARMCD
##
              ARM
## 1 ARM678 BALB/c
## 2 ARM678 BALB/c
## 3 ARM678 BALB/c
## 4 ARM678 BALB/c
## 5 ARM678 BALB/c
## 6 ARM678 BALB/c
# access Concomitant Medications data of SDY139
cm_df <- sdy139$interventions$cm_1$cm_df</pre>
head(cm_df)
## NULL
# get Trial Title from Trial Summary
ts_df <- sdy139$trial_design$ts_l$ts_df
title <- ts_df$TSVAL[ts_df$TSPARMCD== "TITLE"]</pre>
title
```

[1] "The peptide specificity of the endogenous T follicular helper cell repertoire generated after p

8 Get the list of Domain names.

Note that some RImmPort functions take a domain name as input.

```
# get the list of names of all supported Domains
getListOfDomains()
```

```
##
                              Domain Name Domain Code
## 1
                           Adverse Events
                  Concomitant Medications
                                                    CM
## 3
                                                    DM
                             Demographics
## 4
                                                    EX
                                 Exposure
## 5
                          Medical History
                                                    MH
      Associated Persons Medical History
## 6
                                                  APMH
## 7
                 Laboratory Test Results
                                                    LB
## 8
                     Physical Examination
                                                    PΕ
## 9
                      Protocol Deviations
                                                    DV
## 10
                               Trial Arms
                                                    TA
                                                    ΤI
      Trial Inclusion Exclusion Criteria
## 12
                            Trial Summary
                                                    TS
## 13
                           Subject Visits
                                                    SV
## 14
                            Substance Use
                                                    SU
## 15
                              Vital Signs
                                                    ٧S
                           Questionnaires
## 16
                                                    QS
## 17
                           Findings About
                                                    FA
## 18
                            Skin Response
                                                    SR
## 19
                        Genetics Findings
                                                    PF
## 20
                  Protein Quantification
                                                    ZA
## 21
                  Cellular Quantification
                                                    ZB
## 22
                                                    ZC
             Nucleic Acid Quantification
                      Titer Assay Results
## 23
                                                    ZD
```

?"Demographics Domain"

9 Get list of studies with specifc domain data

The Domain name should be exact to what is found in the list of Domain names.

```
# get list of studies with Cellular Quantification data
domain_name <- "Cellular Quantification"
study_ids_l <- getStudiesWithSpecificDomainData(domain_name)
study_ids_l</pre>
```

[1] "SDY139" "SDY208"

10 Get specifc domain data of one or more studies

The Domain name should be exact to what is found in the list of Domain names.

```
# get Cellular Quantification data of studies `SDY139` and `SDY208
# get domain code of Cellular Quantification domain
domain_name <- "Cellular Quantification"</pre>
getDomainCode(domain_name)
## [1] "ZB"
study_ids <- c("SDY139", "SDY208")
domain name <- "Cellular Quantification"
zb 1 <- getDomainDataOfStudies(domain name, study ids)</pre>
## loading Cellular Quantification data....loading FCS Results data....done
## loading ELISPOT Results data....done
## done
## loading Cellular Quantification data....loading FCS Results data....done
## loading ELISPOT Results data....done
## done
if (length(zb_l) > 0)
  names(zb_1)
## [1] "zb df"
                    "suppzb_df"
head(zb 1$zb df)
##
     STUDYID DOMAIN
                       USUBJID ZBSEQ
                                            ZBTEST
                                                                 ZBCAT ZBMETHOD
## 1 SDY139
                 ZB SUB118106
                                   1 Figure-1_FCM Cellular_Phenotype
                                                                            FCM
## 2 SDY139
                 ZB SUB118106
                                   2 Figure-1_FCM Cellular_Phenotype
                                                                            FCM
## 3 SDY139
                 ZB SUB118106
                                   3 Figure-1_FCM Cellular_Phenotype
                                                                            FCM
## 4 SDY139
                 ZB SUB118106
                                   4 Figure-1_FCM Cellular_Phenotype
                                                                            FCM
## 5 SDY139
                                                                            FCM
                 ZB SUB118106
                                   5 Figure-1_FCM Cellular_Phenotype
## 6
     SDY139
                 ZB SUB118106
                                   6 Figure-1 FCM Cellular Phenotype
                                                                            FCM
##
                                                                          ZBPOPDEF
                                  B220- live, doublet excluded, CD4+ CD44low, CCR7+
## 1
## 2
                B220- live, doublet excluded, CD4+ CD44high CXCR5high PD1high, CCR7+
## 3
      B220- live, doublet excluded, CD4+ CD44high, either CXCR5dim or PD1dim, CCR7+
## 4
               B220- live, doublet excluded, CD4+ CD44highCXCR5highPD1high, CD62L+
## 5
                                 B220- live, doublet excluded, CD4+ CD44low, CD62L+
## 6 B220- live, doublet excluded, CD4+ CD44high, either CXCR5dim or PD1dim, CD62L+
##
               ZBPOPNAM ZBORRES
                                                      ZBORRESU ZBBASPOP ZBSPEC
## 1
      CCR7+ CD4+CD44low
                          192974 events per experiment sample
                                                                    <NA>
                                                                           Cell
## 2
              CCR7+ Tfh
                                                                           Cell
                            1701 events per experiment sample
                                                                    <NA>
## 3
          CCR7+ Non-Tfh
                           43396 events per experiment sample
                                                                    <NA>
                                                                           Cell
## 4
             CD62L+ Tfh
                            1386 events per experiment sample
                                                                    <NA>
                                                                           Cell
## 5 CD62L+ CD4+CD44low 166232 events per experiment sample
                                                                    <NA>
                                                                           Cell
## 6
         CD62L+ Non-Tfh
                           38427 events per experiment sample
                                                                    < NA >
                                                                           Cell
##
     VISITNUM
                                                         VISIT ZBELTM
## 1
            3 Day 8 Sample collection, SL_Sant_Plos1_2012_d8
                                                                   P8D
                                                                   P8D
            3 Day 8 Sample collection, SL Sant Plos1 2012 d8
            3 Day 8 Sample collection, SL_Sant_Plos1_2012_d8
                                                                   P8D
## 3
```

```
3 Day 8 Sample collection, SL_Sant_Plos1_2012_d8
## 5
            3 Day 8 Sample collection, SL_Sant_Plos1_2012_d8
                                                                P8D
## 6
           3 Day 8 Sample collection, SL_Sant_Plos1_2012_d8
                                                                P8D
##
                                   ZBTPTREF ZBREFID
                                                                      ZBXFN
## 1 Time of initial vaccine administration BS645819 CCR7 PAN_Tube_009.fcs
## 2 Time of initial vaccine administration BS645819 CCR7 PAN Tube 009.fcs
## 3 Time of initial vaccine administration BS645819 CCR7 PAN_Tube_009.fcs
## 4 Time of initial vaccine administration BS645819 CD62L Pan_Tube_009.fcs
## 5 Time of initial vaccine administration BS645819 CD62L Pan_Tube_009.fcs
## 6 Time of initial vaccine administration BS645819 CD62L Pan_Tube_009.fcs
```

Get the list of assay types from ImmPort studies 11

```
getListOfAssayTypes()
## [1] "ELISA"
                        "ELISPOT"
                                                          "PCR"
                                         "Array"
## [5] "HLA Typing"
                        "MBAA"
                                         "HAI"
                                                          "Neut Ab Titer"
## [9] "Flow"
```

12 Get specific assay data of one or more Immport studies

The assay type should be exact to what is found in the list of supported assay types.

```
# get 'ELISPOT' data of study `SDY139`
assay_type <- "ELISPOT"
study_id = "SDY139"
elispot_l <- getAssayDataOfStudies(study_id, assay_type)</pre>
## loading Protein Quantification data....done
## loading Cellular Quantification data....loading ELISPOT Results data....done
## loading Nucleic Acid Quantification data....done
## loading Titer Assay Results data....done
## loading Genetics Findings data....done
if (length(elispot_l) > 0)
 names(elispot_l)
## [1] "zb_df"
                   "suppzb_df"
head(elispot_l$zb_df)
    STUDYID DOMAIN
                      USUBJID ZBSEQ
                                              ZBTEST
## 1 SDY139
                ZB SUB118053 8657 Figure-4_ELISPOT Cellular_Quantification
## 2 SDY139
                ZB SUB118053 8658 Figure-4_ELISPOT Cellular_Quantification
```

3 SDY139

4 SDY139

ZB SUB118053 8659 Figure-4_ELISPOT Cellular_Quantification

ZB SUB118053 8660 Figure-4_ELISPOT Cellular_Quantification

```
## 5 SDY139
                ZB SUB118053 8661 Figure-4_ELISPOT Cellular_Quantification
## 6 SDY139
                ZB SUB118053 8662 Figure-4_ELISPOT Cellular_Quantification
    ZBMETHOD ZBPOPDEF ZBPOPNAM ZBORRES
                                                     ZBORRESU
                                             1000000 TfH cells
## 1 ELISPOT
                IL-21
                         IL-21 5312.50
                                                                   TfH cells
## 2 ELISPOT
                IL-21
                         IL-21 8337.50
                                             1000000 TfH cells
                                                                   TfH cells
## 3 ELISPOT
                IL-21
                         IL-21 11775.00
                                            1000000 TfH cells
                                                                   TfH cells
## 4 ELISPOT
                                            1000000 TfH cells
                IL-21
                         IL-21 3925.00
                                                                   TfH cells
                                             1000000 TfH cells
                                                                   TfH cells
## 5 ELISPOT
                IL-21
                         IL-21 1931.25
## 6 ELISPOT
                IL-21
                         IL-21
                                  600.00 1000000 Non-TfH cells Non-TfH cells
    ZBSPEC VISITNUM
##
                                                              VISIT ZBELTM
## 1
      Cell
                  3 Day 8 Sample collection, SL_Sant_Plos1_2012_d8
                   3 Day 8 Sample collection, SL_Sant_Plos1_2012_d8
## 2
                                                                       P8D
      Cell
                   3 Day 8 Sample collection, SL_Sant_Plos1_2012_d8
## 3
      Cell
                                                                       P8D
                  3 Day 8 Sample collection, SL_Sant_Plos1_2012_d8
                                                                       P8D
## 4
      Cell
## 5
      Cell
                  3 Day 8 Sample collection, SL_Sant_Plos1_2012_d8
                                                                       P8D
## 6
      Cell
                   3 Day 8 Sample collection, SL_Sant_Plos1_2012_d8
                                                                       P8D
##
                                   ZBTPTREF ZBREFID
                                                                   ZBXFN
## 1 Time of initial vaccine administration BS645680 ELISPOT Results.txt
## 2 Time of initial vaccine administration BS645680 ELISPOT_Results.txt
## 3 Time of initial vaccine administration BS645680 ELISPOT_Results.txt
## 4 Time of initial vaccine administration BS645680 ELISPOT_Results.txt
## 5 Time of initial vaccine administration BS645680 ELISPOT_Results.txt
## 6 Time of initial vaccine administration BS645679 ELISPOT_Results.txt
```

13 Serialize RImmPort-formatted study data as .rds files

```
# serialize all of the data of studies `SDY139` and `SDY208'
study_ids <- c('SDY139', 'SDY208')

# the folder where the .rds files will be stored
rds_dir <- file.path(studies_dir, "Rds")

serialzeStudyData(study_ids, rds_dir)
list.files(rds_dir)</pre>
```

14 Load the serialzed data (.rds) files of a specific domain of a study from the directory where the files are located

```
# get the directory where ImmPort sample data is stored in the directory structure of RImmPort package
studies_dir <- system.file("extdata", "ImmPortStudies", package = "RImmPort")

# the folder where the .rds files will be stored
rds_dir <- file.path(studies_dir, "Rds")

# list the studies that have been serialized
list.files(rds_dir)</pre>
```

```
# load the serialized data of study `SDY208`
study_id <- 'SDY208'
dm_1 <- loadSerializedStudyData(rds_dir, study_id, "Demographics")</pre>
##
## domain_file_path = /Users/user/Projects/RImmPort/inst/extdata/ImmPortStudies/Rds/SDY208/dm.rds
## suppdomain_file_path = /Users/user/Projects/RImmPort/inst/extdata/ImmPortStudies/Rds/SDY208/suppdm.
head(dm_1[[1]])
    STUDYID DOMAIN
                   USUBJID AGE AGEU
                                       SEX
                                               SPECIES STRAIN SBSTRAIN
## 1 SDY208
               DM SUB120516 6 Weeks Female Mus musculus
                                                        <NA>
                                                                  NA
## 2 SDY208
               DM SUB120517 6 Weeks Female Mus musculus
                                                        <NA>
                                                                  NA
               DM SUB120518 6 Weeks Female Mus musculus
## 3 SDY208
                                                        <NA>
                                                                  NA
               DM SUB120519 6 Weeks Female Mus musculus
## 4 SDY208
                                                        <NA>
                                                                  NA
               DM SUB120520 6 Weeks Female Mus musculus
                                                                  NA
## 5 SDY208
                                                        <NA>
               DM SUB120521 6 Weeks Female Mus musculus
## 6 SDY208
                                                        <NA>
                                                                  NA
##
     ARMCD
## 1 ARM881
## 2 ARM882
## 3 ARM883
## 4 ARM884
## 5 ARM885
## 6 ARM886
##
## 1
                                                Microneedle vaccination- 5 ug inactivated A/Califo
## 2
                                              Subcutaneous vaccination- 5 ug inactivated A/Califo
## 3
                                                                      Uncoated microneedle vacci:
## 4 Microneedle vaccination- 5 ug inactivated A/California/04/09 virus, Challenged: 10x LD50 A/Califo
```

Uncoated microneedle vaccination- Placebo, Challenged: 10x LD50 A/Califo

6