# RHAPSODY WP3 Pre-Diabetes Analysis Plan

(Version 1.0.2)

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## Files description

At this stage, you were provided with the following files from the (hidden) Gist (don't share outside RHAPSODY):

- README.Rmd/README.pdf, the file you are actually reading;
- RHAPSODY\_WP3\_PreDiab.Rmd, a Rmarkdown script which perform:
  - opal node access,
  - phenotype QC,
  - preliminary modelling,
  - VCFs formatting/filtering,
  - variants analysis;
- opal\_credentials.txt, the server, login and password informations to access phenotype data on your local node;

Utils files, can be downloaded from the following (hidden) Gist:

- ./utils/RHAPSODY\_WP3\_PreDiab\_DEBUG.R, a small part of the main Rmarkdown script, which includes only node access and phenotype QC;
- ./utils/RunAnalysis.sh, a example of shell script to run the whole analysis in bash (parameters have to be filled properly according to your informations);
- ./utils/handleVCF.sh, the shell script used to format/filter VCFs files prior to the variant analysis. This shell script is also included within the main script, but you can use it directly as a standalone, if so (or if your already done formatting and don't want to do it again), parameter format\_vcfs must be set to FALSE).
- ./utils/Install\_Rpackages.R, a R script which install a predefined version of all R packages loaded in the Rmarkdown script, only if needed (also avaiable within the main script).

## Run the analysis

## R Setup

Please use R version 3.5.1 (2018-07-02) (if not possible, newer version "3.4.\*" can be used). If VCFtools is not available on your machine, please install it: VCFtools.

Be sure the needed packages are installed (this is also done within the Rmarkdown script) or copy/paste the following command. session\_info.csv provided should be used, by default in the working directory.

```
if (!'tidyverse'%in%installed.packages()[, 'Package']) {
  install.package(pkgs = 'tidyverse', repos = 'http://cran.us.r-project.org')
}
if (!'devtools'%in%installed.packages()[, 'Package']) {
  install.package(pkgs = 'devtools', repos = 'http://cran.us.r-project.org')
library(tidyverse)
check_packages_version <- function(</pre>
  list_packages,
  session_info_csv = 'session_info.csv'
) {
  check_packages <- function(package) {</pre>
    if (!package%in%installed.packages()[, 'Package']) {
      install.packages(
       pkgs = package, repos = 'http://cran.us.r-project.org', dependencies = TRUE
      )
   }
   library(
      package = package, character.only = TRUE, quietly = TRUE, warn.conflicts = FALSE
   )
  }
  read csv(file = session info csv) %>%
   dplyr::mutate(
      local_version = purrr::map(.x = package, .f = function(x) {
        if (x%in%utils::installed.packages()[, 'Package']) {
          utils::packageDescription(pkg = x, fields = 'Version')
       }
      })
   ) %>%
   filter(local_version!=version) %>%
   dplyr::arrange(package) %>%
   dplyr::mutate(
      install = purrr::map2(.x = package, .y = version, .f = function(x, y) {
        devtools::install_version(
          package = x, version = as.character(y),
          repos = c(
            'https://rhap-fdb01.vital-it.ch/repo/',
            'http://cran.us.r-project.org'
        )
      })
  invisible(sapply(list_packages, check_packages))
  tidyverse_conflicts()
}
```

```
list_packages <- c(
   'devtools', 'parallel', 'grid', 'scales', 'broom', 'viridis', 'readxl', 'writexl',
   'cowplot', 'knitr', 'kableExtra', 'lme4', 'lmerTest', 'Hmisc', 'data.tree', 'opal'
)
check_packages_version(
   list_packages = list_packages,
   session_info_csv = 'session_info.csv'
)</pre>
```

## Using Docker

If you are familiar with Docker, you can find an image which include everything you need to run the analyses.

#### Run the docker container with the latest image

To run the image, you can use the following command (need docker to be installed):

```
docker run \
   --name rhapsody \
   --hostname rhapsody \
   --detach \
   --volume /your_local_volume:/media/remote_volume \
   --publish 8787:8787 \
   --publish 22:22 \
   biostatgood/rapsody:latest
```

#### Connect to the docker container

Login and password have been set respectively to rhapsody and wp3 (rhapsody is root via sudo). You can connect through SSH using standard IP address: ssh rhapsody@localhost (equivalent to ssh rhapsody@localhost().

You can also connect through Rstudio using a web browser at http://localhost:8787/ (equivalent to http://127.0.0.1:8787/).

Once you are connected directly by SSH or with Rstudio, you can go to the bash terminal and change the password using passwd (just follow the instructions).

#### Docker configuration

The (hidden) Gist s cloned in /home/rhapsody/WP3/scripts.

Make sure after starting the Docker image to update the scripts, if necessary.

```
cd /home/rhapsody/WP3/scripts
git pull origin master
```

VCFtools is also installed in its latest version using github repository.

vcftoolsbinary is located in the default directory: /usr/local/bin (default path in the Rmarkdown scripts).

<sup>&</sup>quot;/your\_local\_volume" is the path to the volume (where your data are located) on the machine when you run the docker software.

<sup>&</sup>quot;/media/remote\_volume" will be the path within the docker container (keep /media/ as root directory).

## Run the Rmarkdown script

- 1. First, create (or change) the opal\_credentials.txt with your informations.
- 2. Fill in the proper informations.
- 3. Set the analysis\_step:
  - 1. Node access
  - 2. Phenotype QC
  - 3. Run Mixed Model without variants
  - 4. Check VCFs files
  - 5. Format VCFs files
  - 6. Run Mixed Model with (imputed) variants
  - 7. Check output files
- 4. Run the R code below within interactive R session.
- 5. a/ If everything worked, you should have a HTML report. Increment analysis\_step and repeat steps 3 to 5.
  - b/ If something went wrong, you should have an error message, for example:

```
label: list_vcf (with options)
List of 2
$ eval : language params_steps$step_5
$ engine: chr "sh"

Quitting from lines 1597-1598 (RHAPSODY_WP3_PreDiab.Rmd)
Error in if (options$eval) { : argument is of length zero}
```

Either try to solve the issue yourself and/or send the informations to mickael.canouil@cnrs.fr

```
# set the output directory or leave as is;
# output will be generated where the Rmarkdown file is
working directory <- './'
analysis step <- 7
cohort_name <- 'Cohort_Name'</pre>
author_name <- 'Firstname LASTNAME'</pre>
opal_credentials <- 'opal_credentials.txt'</pre>
vcf directory <- '/media/.../vcf'</pre>
imputation_quality_tag <- 'INFO' # To be set according to VCF (could also be 'R2')
# Run the analysis
dir.create(path = working_directory, showWarnings = FALSE, mode = '0777')
rmarkdown::render(
  input = '/home/rhapsody/WP3/scripts/RHAPSODY_WP3_PreDiab.Rmd',
  output_format = 'html_document',
  output_file = paste0(
    'RHAPSODY_WP3_PreDiab_',
    cohort_name, '_step',
    analysis_step, '.html'
  ),
  output_dir = working_directory,
  params = list(
    cohort_name = cohort_name,
    author name = author name,
    opal_credentials = opal_credentials,
    vcf_input_directory = vcf_directory,
    imputation_quality_tag = imputation_quality_tag,
    vcftools_binary_path = '/usr/local/bin',
    output_directory = working_directory,
    analysis_step = analysis_step,
    format_vcfs = TRUE,
    variants_analysis = TRUE,
    chunk_size = 10000,
    n_{cpu} = 2,
    echo = FALSE, # Should R code be printed in the report
    warning = FALSE, # Should warnings be printed in the report
    message = FALSE # Should messages be printed in the report
  ),
  encoding = 'UTF-8'
)
```