

# R documentation

of ‘README.Rd’

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PDACMOC

*PDACMolecularOmniClassifier*

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## Description

This package classifies tumor samples according to several molecular classifiers available using different Machine Learning (ML) approaches. It classifies both tumor and stroma fractions. To classify stroma compartment, it first makes a virtual microdissection through the **ADVOCATE** package.

Classifiers available for tumor fractions:

1. Collisson et al., 2011.
2. Moffitt et al., 2015.
3. Bailey et al., 2016.
4. Puleo et al., 2018.
5. Chan-Seng-Yue et al., 2020.
6. PDAConsensus (following the methodology of Kamoun et al., 2020).

Classifiers available for stroma fractions:

1. Moffitt et al., 2016.
2. Maurer et al., 2019.
3. PDAConsensus (following the methodology of Kamoun et al., 2020).

## Details

The package has been built for Linux (Ubuntu 22.04.3 LTS).

It contains the following files:

- inst:
  - examples: it contains a R script ('example.R') with some examples.
  - extdata: it contains the 'Certificate.pdf'.
  - gene\_signatures: it contains a file with all the final genes included.
  - graphs\_and\_tables: it contains two figures that appear in the Shiny app.
  - logos: it contains some institutional logos.
  - models: it contains ML models saved as pickle format.
  - packages: it contains a compressed file ('ADVOCATE\_0.1.0.tar.gz') used to install ADVOCATE package.
  - saved\_workspaces: it contains the workspaces generated with the example script.
  - training\_data: it contains several files used by the classifiers.
- man: it contains the R markdown files produced automatically by devtools::document() using roxygen2 comments.
- R: it contains the different functions of the package.
- releases:
  - download:
    - \* v2.4.0:
      - PDACMOC\_2.4.0.tar.gz: v2.4.0 distribution file
- DESCRIPTION: description file.
- LICENSE: license file.
- NAMESPACE: namespace file produced automatically by devtools::document() using roxygen2 comments.
- PDACMOC.Rproj: R project file.
- README.Rd: readme R markdown file.
- README.pdf: readme pdf file.
- .Rproj.user: directory generated by RStudio that stores user-specific configurations and session states.
- .Rbuildignore: file used by R to exclude specific files and directories from package building.
- .git: directory for Git control version.
- .gitattributes: file used to manage attributes for paths in the repository, such as configuration for Git LFS.
- .gitignore: file used by Git to exclude specific files and directories from tracking and version control.

## Installation

To install the **PDACMOC** package, we recommend to create a Miniconda environment following the next instructions.

```
## Download and uncompressed the distribution file
TBI

## Install Miniconda
wget https://repo.anaconda.com/miniconda/Miniconda3-latest-Linux-x86_64.sh
bash Miniconda3-latest-Linux-x86_64.sh
rm Miniconda3-latest-Linux-x86_64.sh

## Close and open terminal

## Update conda
conda update conda

## Create conda environment
conda create -n PDACMOC

## Activate conda environment
conda activate PDACMOC

## Add conda-forge and bioconda channels
conda config --env --add channels conda-forge
conda config --env --add channels bioconda

## Install Python
conda install python=3.11.4

## Install Python modules
pip3 install numpy pandas pickle-mixin scikit-learn==1.3.1

## Install R
conda install -c conda-forge r-base=4.3.2

## Install libuv, libcurl, and libxml2
conda install -c conda-forge libuv
conda install -c conda-forge libcurl
conda install -c conda-forge libxml2

## Change PKG_CONFIG_PATH
# change path to your miniconda3 environment
export PKG_CONFIG_PATH="/home/pvilloslada/Software/miniconda3/envs/PDACMOC/lib/pkgconfig/:$PKG_CONFIG_PATH"

## Open R
R
```

```
## Install R packages
options(repos = c(CRAN = 'https://cloud.r-project.org/'))

install.packages(c('BiocManager', 'doMC', 'dplyr', 'DT', 'foreach',
                  'iterators', 'mclust', 'plyr', 'reticulate', 'scales',
                  'shiny', 'shinydashboard', 'shinyjs', 'shinythemes'))
BiocManager::install(c('AnnotationDbi', 'DESeq2', 'org.Hs.eg.db', 'sva'))

## Close R
q()

## Install ADVOCATE
R CMD INSTALL ./PDACMOC/inst/packages/ADVOCATE_0.1.0.tar.gz

## Install PDACMOC
R CMD INSTALL ./PDACMOC_2.4.0.tar.gz

## Open R
R

## Check if PDACMOC can be loaded
library(PDACMOC)

## Close R
q()
```

### Shiny app instructions

To run the program online:

1. Connect to server.
2. Upload a tsv file with genes in rows and samples in columns (max. 1GB).
3. Select if you want to apply or not batch correction.
4. Select the gene ID type.
5. Select one or more tumor classifiers.
6. Select if you want to include or not stroma classification.
7. If you want to include stroma classification, select one or more stroma classifiers.
8. Press 'Run classification' and wait about 15-20 minutes.
9. If you want to upload another tsv file press 'Reset app'.

After the process you will find the following:

1. Table of tumor/stroma proportions. 'E' stands for epithelium, 'S' stands for stroma, and 'O' stands for others. 'conf' refers to 95% confidence intervals.

2. Up to 6 tables of tumor classification.
3. Up to 6 summary tables of tumor classification.
4. Up to 3 tables of stroma classification.
5. Up to 3 summary tables of stroma classification.

**Keywords**

PDAC, consensus molecular classifier, transcriptomics, Machine Learning, translational medicine, personalized medicine.

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## Examples

```
library(PDACMOC)

file1 <- system.file('examples', 'example.R', package = 'PDACMOC')
dir <- file.path(dirname(file1), '../saved_workspaces/example.RData')
#load(dir)
rm(file1)

# change path to your miniconda3 environment
```

```
reticulate::use_python("~/Software/miniconda3/envs/PDACMOC/bin/python3.11", required = TRUE)

file2 <- system.file('training_data', 'all_datasets_corrected.csv', package = 'PDACMOC')

samples <- read.csv(file2, row.names = 1, check.names = FALSE)
rm(file2)

new_samples <- PDACMOC::import.and.normalize(samples, batch = FALSE, gene_id = 'EnsemblID')

results_collisson <- PDACMOC::collisson.classify(new_samples)

results_moffitt <- PDACMOC::moffitt.classify(new_samples)

results_bailey <- PDACMOC::bailey.classify(new_samples)

results_puleo <- PDACMOC::puleo.classify(new_samples)

results_chan <- PDACMOC::chan.classify(new_samples)

results_consensus <- PDACMOC::PDAConsensus.classify(new_samples)

vm_result <- PDACMOC::virtual.microdissect(new_samples)

results_stroma_moffitt <- PDACMOC::stroma.moffitt.classify(vm_result$vm_S)

results_stroma_maurer <- PDACMOC::stroma.maurer.classify(vm_result$vm_S)

results_stroma_consensus <- PDACMOC::stroma.PDAConsensus.classify(vm_result$vm_S)

classification_tumor <- PDACMOC::omni.classify(samples, batch = FALSE, gene_id = 'EnsemblID',
  classifier = c('Collisson', 'Moffitt', 'Bailey',
    'Puleo', 'Chan-Seng-Yue', 'PDAConsensus'))

classification_all <- PDACMOC::omni.classify(samples, batch = FALSE, gene_id = 'EnsemblID',
  classifier = c('Collisson', 'Moffitt', 'Bailey',
    'Puleo', 'Chan-Seng-Yue', 'PDAConsensus'),
  stroma = TRUE,
  stroma_classifier = c('Moffitt',
    'Maurer',
    'PDAConsensus'))

# change path to your browser
options(browser = '/usr/bin/firefox')
shinyjs::useShinyjs()
runPDACMOC()

#save.image(dir)
```

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