

Vignette ecophylo

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Rules for working in Rmarkdown with python.

we will need to check the setup chunk for location of python3 binary across different os (ubuntu work with this code)

Python chunks calling R obj.

R chunks works just fine as usual.

```
a <- 42 # set a in R
```

This chunk only work when the markdown is knitted. See that we can call R object when looking for them in 'r' namespace.

```
print(r.a)
```

```
## 42.0
```

```
r.a += 1 # modify R variable in python  
print(r.a)
```

```
## 43.0
```

```
b = 666 # set b in python
```

R chunks calling Python obj.

See that we can call python object when looking for them in 'py' list.

```
print(a) # call a in R after modification in python
```

```
## [1] 43
```

```
print(py$b) # call b in R
```

```
## [1] 666
```

```
# Also possible to call directly python function in r  
rand <- import('numpy')$random # import python module numpy and submod random  
rand$randint(3) # call of function randint
```

```
## [1] 0
```

You can run python code in interactive mod in the console using the following function : `repl_python(quiet = T)`. It set the console in 'python mode' so you can type in python commands. However you can't send lines from a chunk or a script.

To exit the consol in python, just type *exit* and tadaa the consol is back to R.

Using Python modules in R.

```
# Be carefull for idiotproof of ecophylo and int class !
eco <- import('ecophylo')
eco$timeframes(I=as.integer(3), T=2, a=0.3)
```

```
## [[1]]
## [1] 0.565357
##
## [[2]]
## [1] 1.226603
##
## [[3]]
## [1] 2
```

```
# make dataframes with pandas is same as R
pd <- import('pandas')
d <- list(col1 = c(1,2,3), col2 = c(4,5,6))
pd$DataFrame(d)
```

```
##   col1 col2
## 1    1    4
## 2    2    5
## 3    3    6
```

Serious Part

Introduction

Installation

You need to download the *tar.gz* file from the repository and install it with pip. You need to use python3 and install the dependencies before.

impossible at this moment as the repository is private impossible to install from popy because not pushed there at this time

```
# NOT RUN
wget -L https://github.com/thegreatlizzyator/ecophylo/blob/master/dist/ecophylo-0.0.5.tar.gz
python3 -m pip install ecophylo-0.0.5.tar.gz
```

Installation on windows (the painfull way)

For François : First create a new project in rstudio with **version control** from the github.

You will need to install {reticulate} package and python dependencies.

```
# NOT RUN
install.packages('reticulate') # answer yes
library(reticulate)
# python dependencies
conda_install('r-reticulate', c('msprime', 'ete3', 'pandas'))
```

Single simulation

```
# First we need to import the python module
eco <- import('ecophylo')
# Be carefull for idiotproof of ecophylo and int class ! we need to
eco$timeframes(I=as.integer(3), T=2, a=0.3)

## [[1]]
## [1] 0.565357
##
## [[2]]
## [1] 1.226603
##
## [[3]]
## [1] 2

sim1 <- eco$simulate(as.integer(100), as.integer(1e4), as.integer(1e-3))
sim1

##
##      /-20
## --|
##      /-20
```

Multiple simulations

```
# NOT RUN  
truc <- eco$dosimuls(nsim = 1, sample_size = 100, comprior = list(1000,10e9),  
  muprior = list(1e-3) , verbose = TRUE, seed = 42)
```

```
# NOT RUN  
eco.dosimuls(nsim = 5, sample_size = 100, comprior = [1000,10e9],  
  muprior = [1e-3] , verbose = True, seed = 42)
```