GRANAR User Guide

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What is GRANAR?

GRANAR stands for Generator of Root Anatomy in R. The GRANAR model is able to generate complete cell networks of root cross-sections using a small set of root anatomical parameters These root anatomical features can typically be gathered using software image analysis such as ImageJ (Schneider, Rasband, and Eliceiri 2012). Once the root anatomy is created, the cell network can be saved as an eXtended Markup Language (XML) file. The structure of the XML files created by GRANAR are identical to the ones created by CellSeT (Pound et al. 2012) (a software which digitize root anatomical network).

How to use GRANAR?

GRANAR is available as an R package. The 'granar' package can be found on GitHub. Alternatively, GRANAR can be installed locally by simply executing the following line in the R environment:

```
install.packages("devtools")
library(devtools)
install_github("granar/granar")
library(granar)
```

Granar was built upon the following dependencies dependencies:

```
# Load the libraries
library(xm12)
library(tidyverse)
library(plyr)
library(deldir)
library(alphahull)
library(viridis)
library(sp)
library(cowplot)
library(retistruct)
library(Hmisc)
# Working with Python in R
library(reticulate)
```

You do not need your own data to try it out, some examples of input parameters of GRANAR can be found in a Repository. The following table shows an example of all input variables inside one of the .XML file used to run GRANAR:

```
##
            name
                           type
                                 value
## 1
       planttype
                                  1.000
                          param
## 2 randomness
                                  3.000
                          param
## 3
                 cell diameter
           stele
                                  0.011
## 4
           stele
                       n layers
                                 1.000
## 5
           stele layer_diameter 0.210
```

```
## 6
          stele
                          order
                                1.000
## 7
      pericycle cell_diameter
                                0.011
                      n layers
## 8
      pericycle
                                1.000
## 9
      pericycle
                          order
                                2.000
## 10 endodermis cell diameter
                                0.025
## 11 endodermis
                      n layers 1.000
## 12 endodermis
                         order 3.000
## 13
          cortex cell diameter 0.030
## 14
          cortex
                      n_layers
                                6.000
## 15
          cortex
                          order
                                4.000
## 16
      exodermis cell_diameter 0.030
## 17
      exodermis
                      n_layers
                                1.000
## 18
      exodermis
                         order
                                5.000
      epidermis cell_diameter 0.010
## 19
## 20
      epidermis
                      n_layers
                                1.000
## 21
      epidermis
                         order
                                6.000
## 22
                      max_size 0.050
          xylem
## 23
          xvlem
                       n files
                                5.000
## 24
          xylem
                         order 1.500
## 25
          xylem
                         ratio
                                1.860
## 26
         phloem
                      max_size 0.010
## 27
         phloem
                       n files 3.000
## 28 aerenchyma
                    proportion 0.200
## 29 aerenchyma
                       n files 10.000
```

Importing the .XML parameter file into R can be done with the function read_param_xml(). Example:

```
# Read the parameters
params <- read_param_xml("granar_examples/modelparam/Zea_mays_Heymans_2019.xml")</pre>
```

It is possible to change directly some of the parameters one-by-one in R.

```
# Copy and modify the parameters
params_no_aerenchyma <- params
params_no_aerenchyma$value[params$name == "aerenchyma" & params$type == "proportion"] <- 0</pre>
```

Once, the input file is loaded, the function $create_anatomy()$ can be called to generate the corresponding cross-section.

```
# Run the model with the loaded parameters
sim <- create_anatomy(parameters = params, verbatim=F)
## Time difference of 7.892196 secs</pre>
```

```
# Run the model with the modified parameters
sim_no_aer <- create_anatomy(parameters = params_no_aerenchyma, verbatim=F)
```

```
## Time difference of 1.045678 secs
```

The funtion plot_anatomy() can be used to vizualize the generated anatomy. Different informations can be displayed such as the cell type (fig. 1) or the cell size of the cross section (fig. 2)

```
# Create figure of the generated anatomies.
plot_anatomy(sim, col = "type")
plot_anatomy(sim_no_aer, col = "area")
```

When the anatomy is completed it can be saved using the write anatomy xml() function.

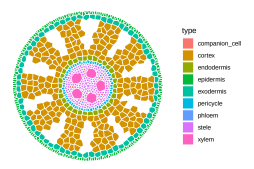


Figure 1: Maize cross-section anatomy made by GRANAR.

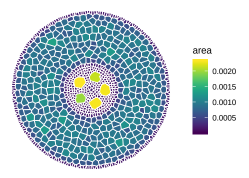


Figure 2: Maize cross-section anatomy made by GRANAR.

Loading the nodes data of a cross-section can be achieved with get_root_section(). Below, we load a fully digitilized cross section done by CellSeT cross-section of a maize (Zea mays. B73 line) crown root grown aeroponically (fig. 3). It was collected 5 cm above the root tip. And the initial image cross section come from Hachez et al. (2006).

Next, to compare visually the root anatomical network obtain with CellSeT with the one of GRANAR, we generated the coresponding root cross section with GRANAR thanks to the cell data inside the CellSeT file.

```
# It is possible to load cross-section xml under cellset format
Cross_section <- list()
Cross_section$nodes <- get_root_section(
   path="granar_examples/cellsetdata/Maize_pip_cross8_phloem.xml")</pre>
```

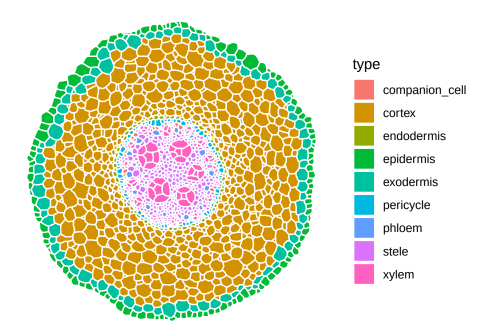


Figure 3: Cross-section anatomy made by CellSeT. Maize (Zea mays. B73 line) crown root grown aeroponically. It was collected 5 cm above the root tip. And the initial image cross section come from Hachez et al. (2006). The digitilization with CellSeT was performed to run MECHA (Couvreur et al. 2018) on it.

```
node$area[node$id_cell == i] <- pol@area</pre>
}
# Get information per cell type.
condens <- node%>%
  dplyr::group_by(type)%>%
  dplyr::summarise(area_type = sum(area),
                   cell type = mean(area),
                    cell_size = 2*sqrt(cell_type/pi),
                    n = n()
                    r_{type} = (max(x) - min(x) + max(y) - min(y))/4)%
  dplyr::ungroup()
# Replace input parameter of GRANAR with the cell data gathered from CellSeT.
# Cell data are turned from ?m to mm.
params$value[params$name == "aerenchyma" &
             params$type == "proportion"] <- 0</pre>
params$value[params$name == "randomness" &
             params$type == "param"] <- 4</pre>
params$value[params$name == "stele" &
             params$type == "cell_diameter"] <- condens$cell_size[</pre>
               condens$type == "stele"]/1000
params$value[params$name == "stele" &
             params$type == "layer_diameter"] <- condens$r_type[</pre>
               condens$type == "stele"]*2/1000
params$value[params$name == "pericycle" &
             params$type == "cell_diameter"] <- condens$cell_size[</pre>
               condens$type == "pericycle"]/1000
params$value[params$name == "endodermis" &
             params$type == "cell_diameter"] <- condens$cell_size[</pre>
               condens$type == "endodermis"]/1000
params$value[params$name == "cortex" &
             params$type == "cell_diameter"] <- condens$cell_size[</pre>
               condens$type == "cortex"]*0.9/1000
params$value[params$name == "cortex" &
             params$type == "n_layers"] <- round((condens$r_type[</pre>
               condens$type == "cortex"] - condens$r_type[
               condens$type == "endodermis"])/condens$cell_size[
               condens$type == "cortex"])
params$value[params$name == "exodermis" &
             params$type == "cell_diameter"] <- condens$cell_size[</pre>
               condens$type == "exodermis"]/1000
params$value[params$name == "epidermis" &
             params$type == "cell_diameter"] <- condens$cell_size[</pre>
               condens$type == "epidermis"]/1000
params$value[params$name == "xylem" &
             params$type == "max_size"] <- condens$r_type[</pre>
               condens$type == "stele"]*0.425/1000
params$value[params$name == "xylem" &
             params$type == "ratio"] <- 3</pre>
# Run GRANAR
sim <- create_anatomy(parameters = params)</pre>
```

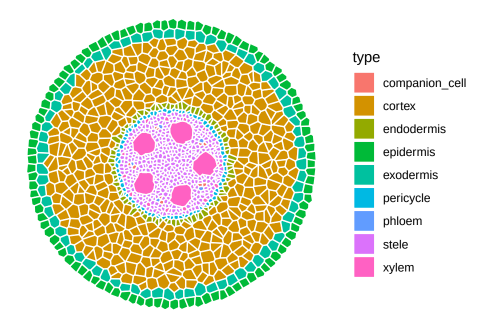


Figure 4: Maize cross-section anatomy made by GRANAR.

type	id_group		##
exodermis	1	1	##
epidermis	2	2	##
endodermis	3	3	##
cortex	4	4	##
stele	5	5	##
xylem	13	6	##
pericycle	16	7	##
companion_cell	12	8	##
phloem	11	9	##

Coupling GRANAR with MECHA

MECHA

MECHA is an explicit cross-section model of the root hydraulic anatomy which connects hydraulic concepts across scales (Couvreur et al. 2018).

The model computes horizontal water flow at the level of individual cells, quantifies the contribution of water composite pathways, and predicts root radial permeability (k_r) , using detailed anatomical descriptions and experimental data on the permeability of cell walls (k_w) , membranes (L_p) and the conductance of individual plasmodesmata (K_{PD}) .

Install MECHA

Here, we installed Canopy to run MECHA which was coded in Python. Canopy offer the possibilty to install easily the dependencies of MECHA:

- numpy
- scipy
- networkx (v.1.9.1)

Time difference of 1.290753 mins

• lxml

How to calculate hydraulic properties on newly aquired root cross section

```
# The user is invited to select the pathway to Python
use_python("../../AppData/Local/Enthought/Canopy/edm/envs/User")
os <- import("os")</pre>
# If os produce error: verify your python directory
# use Canopy package manager to install
# numpy, scipy, networkx, lxml, matplotlib and pyqt
# For windows user, pygt allows matplotlib to run without error
numpy <- import("numpy")</pre>
scipy <- import("scipy")</pre>
networkx <- import("networkx")</pre>
lxml <- import("lxml")</pre>
matplotlib <- import("matplotlib")</pre>
# The following function call MECHA and return the radial conductivity
source("granar examples/MECHA R.R")
Kr_Granar <- MECHA_R(path = "granar_examples/MECHAv4_light.py")</pre>
## Time difference of 46.40314 secs
fc <- file.copy(from = "granar_examples/cellsetdata/Maize_pip_cross8_phloem.xml",</pre>
                 to = paste0("granar_examples/cellsetdata/current_root.xml"),
                 overwrite = T)
Kr_CellSet <- MECHA_R(path = "granar_examples/MECHAv4_light_1.py")</pre>
```

- The $Kr\theta$: The Radial conductivity with only a endodermal casparian strip (cm hPa-1 d-1).
- The Kr1: The Radial conductivity with suberized endodermis (cm hPa-1 d-1).
- The Kr2: The Radial conductivity with suberized endodermis and exodermal Casparian strip (cm hPa-1 d-1).

References

Couvreur, Valentin, Marc Faget, Guillaume Lobet, Mathieu Javaux, François Chaumont, and Xavier Draye. 2018. "Going with the Flow: Multiscale Insights into the Composite Nature of Water Transport in Roots." *Plant Physiol.* 178 (4). Am Soc Plant Biol: 1689–1703.

Hachez, Charles, Menachem Moshelion, Enric Zelazny, Damien Cavez, and François Chaumont. 2006. "Localization and Quantification of Plasma Membrane Aquaporin Expression in Maize Primary Root: A Clue to Understanding Their Role as Cellular Plumbers." *Plant Molecular Biology*.

Pound, Michael P, Andrew P French, Darren M Wells, Malcolm J Bennett, and Tony P Pridmore. 2012. "CellSeT: Novel Software to Extract and Analyze Structured Networks of Plant Cells from Confocal Images." *Plant Cell* 24 (4): 1353–61.

Schneider, Caroline A, Wayne S Rasband, and Kevin W Eliceiri. 2012. "NIH Image to ImageJ: 25 Years of Image Analysis." *Nat. Methods* 9 (7). nature.com: 671–75.