# Correcting parameters

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#### I. Introduction

#### I.a) Set up

```
wd = "D:/Internship 3A/GitHub/Project-Klamath-2021/Model Parameterization"
files_dir = "E:/FIF/Stage 3A/Results single cell/CC single cell/"

setwd(wd)
library(magrittr)
library(cowplot)
library(ggplot2)
library(raster)
library(knitr)
```

### I.b) Presentation

When looking at the results after a first run of the scenario with only the climate change (CC only), it was observed that the ecosystem was starved in nitrogen. The more likely explanation is a too important stock of nitrogen in one (or several) of the soils pools of carbon and nitrogen.

These pools simulates the different soil stocks of nitrogen and carbon that interveine in the biochemical cycles. As decay progresses, carbon and nitrogen are transferred to and among the fast, slow, and passive soil organic matter (SOM) pools, with some loss to the atmosphere and through leaching. The picture below is a schematic diagram of NECN functionning, based on the CENTURY modelling.

The process described concern only the bottom panel, showing the 3 pools SOM1, SOM2 and SOM3. NECN allows to parameter a forth pool, simulating the soil litter.

Thus, it was decided to change the decay rate of these pools. To do so and not spend too many time on computing, these tests where done using the single cell landscape. The method used to build this test landscape is detailed on the pdf document "Test\_Landscape".

# I.c) Scenario

The scenario used to test the parameters correspond to a single cell landscape with only climate change i.e. without disturbancies (insect breakout, fire or harvest). A summary of the settings used in the different subscenarios of the signle cell run are presented on the following table. They correspond to the decay rate of the four soil layers.

#### Cohort Net Primary Productivity H2O, T, LAI, available N, maximum monthly NPP by species, maximum capacity by species L = leaf W W W W W one to many W = woodFR = fine root cohorts ... FR CR FR CR FRICR FRICR FR CR CR = coarse root Surface Residue Soil Residue Detrital Dead Dead Wood Structural Metabolic Metabolic Coarse Roots Structural Pools M---->CO<sub>2</sub>, Available N Non-Lignin Lignin SOM 1 SOM 2 Fast pool Slow pool Legend SOM 3 Passive pool ► Cand N flow ---▶ Decomposition CO2, Available N

Figure 1: Capture of figure NECN publication

Table 1: Decay rates used in the sub-scenarios

	V1	V2	V3	V4	V5
DecayRateSurf	0.8200	0.9900	0.9900	0.9900	0.9900
DecayRateSOM1	0.5300	0.7600	0.9000	0.9900	0.9900
${\bf DecayRateSOM2}$	0.0200	0.0400	0.0500	0.9900	0.0400
DecayRateSOM3	0.0003	0.0003	0.0003	0.0003	0.0003

# II. Carbon and nitrogen dynamics

The succetion log files of the NECN extention are imported for all the sub-scenarios. Only the variables of interest are selected.

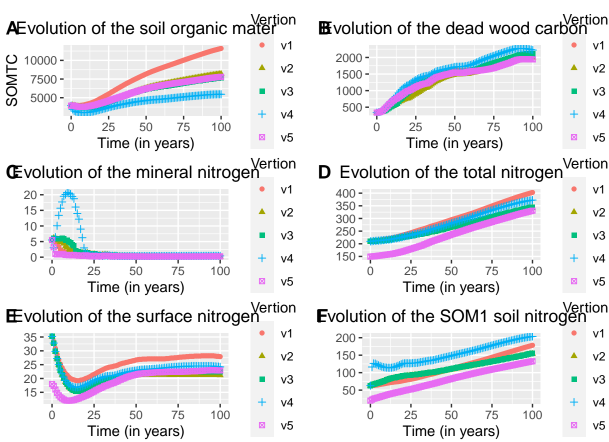
```
NECN_V1_long = pasteO(files_dir,"NECN_V1.csv") %>% read.csv()
NECN_V1 = aggregate(data=NECN_V1_long, cbind(SOMTC, MineralN, TotalN, C_DeadWood,
                                             N_SOM1surf, N_SOM1soil, N_SOM2) ~ Time, mean)
NECN_V1$Vertion = "v1"
NECN_V2_long = pasteO(files_dir,"NECN_V2.csv") %>% read.csv()
NECN_V2 = aggregate(data=NECN_V2_long, cbind(SOMTC, MineralN, TotalN, C_DeadWood,
                                             N SOM1surf, N SOM1soil, N SOM2) ~ Time, mean)
NECN V2$Vertion = "v2"
NECN_V3_long = pasteO(files_dir,"NECN_V3.csv") %>% read.csv()
NECN_V3 = aggregate(data=NECN_V3_long, cbind(SOMTC, MineralN, TotalN, C_DeadWood,
                                             N_SOM1surf, N_SOM1soil, N_SOM2) ~ Time, mean)
NECN_V3$Vertion = "v3"
NECN_V4_long = pasteO(files_dir,"NECN_V4.csv") %>% read.csv()
NECN_V4 = aggregate(data=NECN_V4_long, cbind(SOMTC, MineralN, TotalN, C_DeadWood,
                                             N_SOM1surf, N_SOM1soil, N_SOM2) ~ Time, mean)
NECN V4$Vertion = "v4"
```

The carbon and nitrogen dynamics are then plotted against time for all the sub-scenario.

```
SOMTp = ggplot(NECN, aes(x=Time, y=SOMTC, col=Vertion, shape = Vertion)) +
  geom_point() +
  scale_x_continuous(name= "Time (in years)") +
  scale_y_continuous(name = "SOMTC") +
  ggtitle("Evolution of the soil organic mater") +
  theme(plot.title = element_text(hjust = 0.5))
CDWp = ggplot(NECN, aes(x=Time, y=C_DeadWood, col=Vertion, shape = Vertion)) +
  geom point() +
  scale_x_continuous(name= "Time (in years)") +
  scale y continuous(name = "") +
  ggtitle("Evolution of the dead wood carbon") +
  theme(plot.title = element text(hjust = 0.5))
MNp = ggplot(NECN, aes(x=Time, y=MineralN, col=Vertion, shape = Vertion)) +
  geom point() +
  scale_x_continuous(name= "Time (in years)") +
  scale_y_continuous(name = "") +
  ggtitle("Evolution of the mineral nitrogen") +
  theme(plot.title = element_text(hjust = 0.5))
TNp = ggplot(NECN, aes(x=Time, y=TotalN, col=Vertion, shape = Vertion)) +
  geom_point() +
  scale_x_continuous(name= "Time (in years)") +
  scale_y_continuous(name = "") +
  ggtitle("Evolution of the total nitrogen") +
  theme(plot.title = element_text(hjust = 0.5))
SNp = ggplot(NECN, aes(x=Time, y=N_SOM1surf, col=Vertion, shape = Vertion)) +
  geom point() +
  scale_x_continuous(name= "Time (in years)") +
  scale y continuous(name = "") +
  ggtitle("Evolution of the surface nitrogen") +
  theme(plot.title = element_text(hjust = 0.5))
N1p = ggplot(NECN, aes(x=Time, y=N_SOM1soil, col=Vertion, shape = Vertion)) +
  geom_point() +
  scale_x_continuous(name= "Time (in years)") +
  scale_y_continuous(name = "") +
  ggtitle("Evolution of the SOM1 soil nitrogen") +
  theme(plot.title = element_text(hjust = 0.5))
```

```
N2p = ggplot(NECN, aes(x=Time, y=N_SOM2, col=Vertion, shape = Vertion)) +
  geom_point() +
  scale_x_continuous(name= "Time (in years)") +
  scale_y_continuous(name = "") +
  ggtitle("Evolution of the SOM2 nitrogen") +
  theme(plot.title = element_text(hjust = 0.5))

plot_grid(SOMTp, CDWp, MNp, TNp, SNp, N1p, N2p, labels="AUTO", ncol = 2, nrow = 3)
```



V4 is choosen

# III. Initial quantity of soil nitrogen

#### III.a) Modification of initial nitrogen cocentration for a single cell raster

It is observed

```
Nsoil1 = paste(files_dir,"som1nsoil2.tif",sep="") %>% raster()
Nsurf1 = paste(files_dir,"som1nsurf4.tif",sep="") %>% raster()
a = freq(Nsoil1, merge=T) %>% as.data.frame()
b = freq(Nsurf1, merge=T) %>% as.data.frame()
```

Table 2: Initial and modified concentration of nitrogen on the SOM1 layers

	value	count
Initial N on SOM1 soil layer	63	1
Initial N on SOM1 surface layer	35	1
New N on SOM1 soil layer	20	1
New N on SOM1 surface layer	18	1

## Warning in .gd\_SetProject(object, ...): NOT UPDATED FOR PROJ >= 6

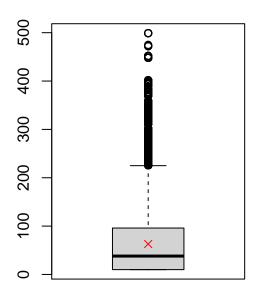
# III.b) Modification of initial nitrogen cocentration for the initial raster

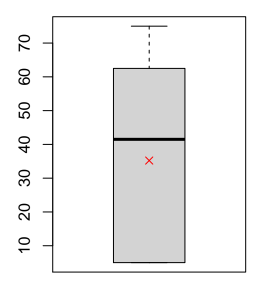
```
Nsoil1_Tot = paste(files_dir, "Tot/som1nsoil2.tif", sep="") %>% raster()
Nsurf1_Tot = paste(files_dir, "Tot/som1nsurf4.tif", sep="") %>% raster()

SoilMean = cellStats(Nsoil1_Tot, stat='mean', na.rm=TRUE, asSample=TRUE)
SurfMean = cellStats(Nsurf1_Tot, stat='mean', na.rm=TRUE, asSample=TRUE)

par(mfrow=c(1,2))
boxplot(Nsoil1_Tot, maxpixels=1000000, xlab = "SOM1 Soil N")
cellStats(Nsoil1_Tot, stat='mean', na.rm=TRUE, asSample=TRUE) %>% points(pch=4,col="red")

boxplot(Nsurf1_Tot, maxpixels=1000000, xlab = "SOM1 surface N")
cellStats(Nsurf1_Tot, stat='mean', na.rm=TRUE, asSample=TRUE) %>% points(pch=4,col="red")
```





### SOM1 Soil N

#### SOM1 surface N

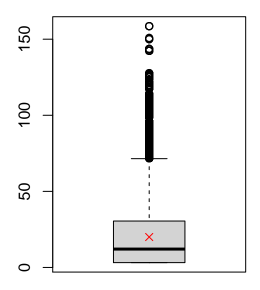
```
print(paste("Mean of SOM1 soil raster:", SoilMean, "Mean of SOM1 surface raster:", SurfMean, sep=" "))
## [1] "Mean of SOM1 soil raster: 62.9439680398293 Mean of SOM1 surface raster: 35.2173852118269"

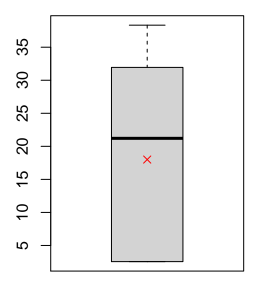
Nsoil1_New = Nsoil1_Tot*(20/SoilMean)

Nsurf1_New = Nsurf1_Tot*(18/SurfMean)

par(mfrow=c(1,2))
boxplot(Nsoil1_New, maxpixels=1000000, xlab = "SOM1 Soil N")
cellStats(Nsoil1_New, stat='mean', na.rm=TRUE, asSample=TRUE) %>% points(,pch=4,col="red")

boxplot(Nsurf1_New, maxpixels=1000000, xlab = "SOM1 surface N")
cellStats(Nsurf1_New, stat='mean', na.rm=TRUE, asSample=TRUE) %>% points(,pch=4,col="red")
```





### SOM1 Soil N

SOM1 surface N