2.2 - Forest water/energy balance (practice)

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Outline

- 1. Water balance input object
 - 2. Basic water balance
- 3. Evaluating model performance
- 4. Advanced water/energy balance
 - 5. Modifying model inputs



Creating the water balance input object

We assume we have an appropriate forest object and species parameter data frame:

```
data(exampleforestMED)
data(SpParamsMED)
```



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a soil object:

```
examplesoil <- soil(defaultSoilParams(4))</pre>
```



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data(exampleforestMED)
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a soil object:
    examplesoil <- soil(defaultSoilParams(4))

and a simulation control list:
    control <- defaultControl("Granier")</pre>
```



Creating the water balance input object

We assume we have an appropriate forest object and species parameter data frame:

x <- forest2spwbInput(exampleforestMED, examplesoil, SpParamsMED, control)</pre>

```
data(exampleforestMED)
data(SpParamsMED)

a soil object:
    examplesoil <- soil(defaultSoilParams(4))

and a simulation control list:
    control <- defaultControl("Granier")

With these four elements we can build our input object for function spwb():</pre>
```



Structure of the water balance input object (1)

The water balance input object is a list with several elements:



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Elements soil and control contain copies of the parameters used in the call to forest2spwbInput().



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Elements soil and control contain copies of the parameters used in the call to forest2spwbInput().

Element cohorts contains the species identity of each cohort:

```
x$cohorts

## SP Name
## T1_148 148 Pinus halepensis
## T2_168 168 Quercus ilex
## S1_165 165 Quercus coccifera
```



Structure of the water balance input object (2)

Element above contains above-ground description of vegetation:



Structure of the water balance input object (2)

Element above contains above-ground description of vegetation:

Element below contains below-ground description of vegetation:

```
x$below

## Z50 Z95 fineRootBiomass coarseRootSoilVolume
## T1_148 100 600 1574.98321 4.381122
## T2_168 300 1000 667.64217 7.526346
## S1_165 200 1000 13.57757 5.544086
```



Structure of the water balance input object (3)

Elements params * contain cohort-level parameters, for example...

```
x$paramsTranspiration
              Gswmin Tmax_LAI Tmax_LAIsq Psi_Extract Psi_Critic
##
                                                                        WUE
                                                                              WUE_par
                                                                                          WUE_co2
## T1_148 0.003086667 0.1756248 -0.007863797
                                              -2.303000
                                                          -5.14000 8.543861 0.6826373 0.002517464
## T2_168 0.004473333 0.1121914 -0.005023494
                                              -2.840000
                                                         -6.96500 8.605536 0.2572564 0.002718680
## S1_165 0.010455247 0.1340000 -0.006000000
                                                         -6.95092 7.900000 0.3643000 0.002757000
                                              -2.121073
##
            WUE_vpd
## T1_148 -0.3032638
## T2_168 -0.5780737
## S1_165 -0.4636000
```



Structure of the water balance input object (3)

Elements params * contain cohort-level parameters, for example...

```
x$paramsTranspiration
              Gswmin Tmax_LAI
                                Tmax_LAIsq Psi_Extract Psi_Critic
                                                                       WUE
                                                                             WUE_par
                                                                                        WUE_co2
## T1_148 0.003086667 0.1756248 -0.007863797
                                             -2.303000
                                                         -5.14000 8.543861 0.6826373 0.002517464
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                                             -2.840000
                                                        -6.96500 8.605536 0.2572564 0.002718680
## S1_165 0.010455247 0.1340000 -0.006000000
                                             -2.121073 -6.95092 7.900000 0.3643000 0.002757000
##
            WUE_vpd
## T1_148 -0.3032638
## T2_168 -0.5780737
## S1_165 -0.4636000
or ...
x$paramsAnatomy
                             SLA LeafDensity WoodDensity FineRootDensity
##
            Al2As Ar2Al
                                                                             SRL RLD
                                                                                        r635
## T1_148 1317.523
                                   0.2982842 0.6077016
                      1 5.140523
                                                              0.2982842 3172.572 10 1.964226
## T2_168 3908.823
                      1 6.340000
                                   0.4893392 0.9008264
                                                              0.4893392 4398.812 10 1.805872
## S1_165 4189.325
                      1 4.980084
                                   0.3709679 0.4389106
                                                              0.3709679 4398.812 10 2.289452
```



Structure of the water balance input object (4)

Elements internal* contain cohort-level state variables, for example:

```
x$internalPhenology
         gdd sen budFormation leafUnfolding leafSenescence leafDormancy phi
##
## T1_148
                         FALSE
                                       FALSE
                                                      FALSE
                                                                   FALSE
## T2_168 0
                         FALSE
                                       FALSE
                                                                   FALSE
                0
                                                      FALSE
## S1_165 0
                         FALSE
                                       FALSE
                                                      FALSE
                                                                   FALSE
                0
```



Structure of the water balance input object (4)

0

Elements internal* contain cohort-level state variables, for example:

```
x$internalPhenology
         gdd sen budFormation leafUnfolding leafSenescence leafDormancy phi
##
## T1_148
                         FALSE
                                       FALSE
                                                      FALSE
                                                                   FALSE
## T2_168 0
                         FALSE
                                       FALSE
               0
                                                      FALSE
                                                                   FALSE
## S1_165 0 0
                         FALSE
                                       FALSE
                                                      FALSE
                                                                   FALSE
or...
x$internalWater
```

S1_165 -0.033



Water balance run

Let us assume we have an appropriate weather data frame:

data(examplemeteo)



Water balance run

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```
data(examplemeteo)
```

The call to function spwb() needs the water balance input object, the weather data frame, latitude and elevation:

```
S <- spwb(x, examplemeteo, latitude = 41.82592, elevation = 100)
```

```
## Initial soil water content (mm): 291.257
## Initial snowpack content (mm): 0
## Performing daily simulations
##
##
    [Year 2001]:......
##
## Final soil water content (mm): 265.781
## Final snowpack content (mm): 0
## Change in soil water content (mm): -25.4761
## Soil water balance result (mm): -25.4761
## Change in snowpack water content (mm): 0
## Snowpack water balance result (mm): 7.10543e-15
## Water balance components:
    Precipitation (mm) 513
##
    Rain (mm) 462 Snow (mm) 51
    Interception (mm) 96 Net rainfall (mm) 366
##
    Infiltration (mm) 408 Runoff (mm) 9 Deep drainage (mm) 66
##
    Soil evaporation (mm) 17 Transpiration (mm) 350
##
    Plant extraction from soil (mm) 350 Plant water balance (mm) -0 Hydraulic redistribution (mm) 10
##
```



Water balance output object (1)

Function spwb() returns an object of class with the same name, actually a list:

```
class(S)
## [1] "spwb" "list"
```



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It is interesting to inspect the list element names:



Water balance output object (1)

Function spwb() returns an object of class with the same name, actually a list:

```
class(S)
## [1] "spwb" "list"
```

It is interesting to inspect the list element names:

```
names(S)

## [1] "latitude" "topography" "weather" "spwbInput" "spwbOutput" "WaterBalance"
## [7] "Soil" "Stand" "Plants"
```

Elements	Information
latitude, topography, weather, spwbInput	Copies of the information used in the call to spwb()
spwb0utput	State variables at the end of the simulation (can be used as input to a subsequent one)
WaterBalance, Soil, Stand, Plants	Daily outputs (days as rows, variables as columns)
subdaily	Sub-daily outputs (not relevant here)



Water balance output object (2)

Actually, Plants is itself a list with several data frames of results by cohort (days as rows, cohorts as columns):



Summaries

The package provides a summary() function for objects of class spwb. It can be used to extract/summarize the model's output at different temporal steps (i.e. weekly, annual, ...).

For example, to obtain the average soil moisture and water potentials by months one can use:

```
summary(S, freq="months", FUN=mean, output="Soil")
```



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Parameter output indicates the element of the spwb object for which we desire a summary. Similarly, it is possible to calculate the average stress of plant cohorts by months:

```
summary(S, freq="months",FUN=mean, output="PlantStress")
```



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```
summary(S, freq="months",FUN=mean, output="PlantStress")
```

The summary function can be also used to aggregate the output by species. In this case, the values of plant cohorts belonging to the same species will be averaged using LAI values as weights:

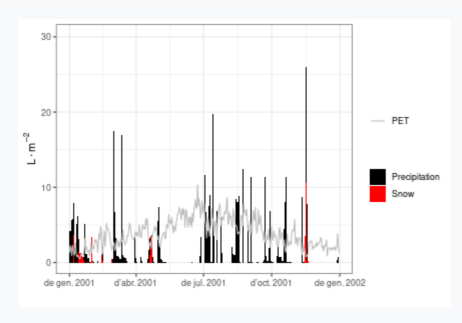
```
summary(S, freq="months", output="PlantStress", bySpecies = TRUE)
```



Plots

The package provides a plot() function for objects of class spwb. It can be used to show weather inputs and different components of the water balance, for example:

```
plot(S, type = "PET_Precipitation")
```

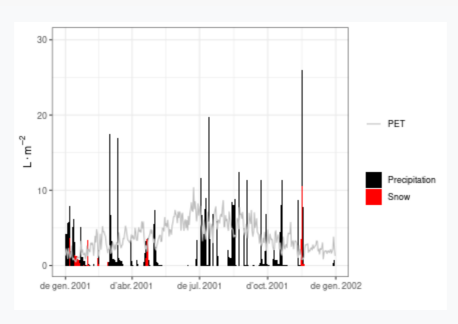




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The help page of ?plot.spwb lists all the possible plots...



Interactive plots

... but instead of typing all plots, we can call the interactive plot function and explore them all:

shinyplot(S)



Post-processing functions

The package provides some functions to extract or transform specific outputs from spwb() simulations.



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Function droughtStress() allows calculating several plant stress indices, such as the maximum drought stress value per month:

```
DS <- droughtStress(S, index = "MDS", freq = "months", draw=FALSE)
head(DS)

## T1_148 T2_168 S1_165
## 2001-01-01 3.461212e-06 1.575275e-06 4.102597e-06
## 2001-02-01 1.898577e-05 6.896248e-06 2.038892e-05
## 2001-03-01 2.016387e-05 7.291361e-06 2.161446e-05
## 2001-04-01 6.543452e-05 2.241727e-05 6.863687e-05
## 2001-05-01 1.909799e-04 6.520781e-05 2.000474e-04
## 2001-06-01 5.836492e-03 1.978796e-03 6.085632e-03
```



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## 2001-05-01 1.909799e-04 6.520781e-05 2.000474e-04
## 2001-06-01 5.836492e-03 1.978796e-03 6.085632e-03
```

Other similar post-processing functions are waterUseEfficiency() or fireHazard().

They (should) also work on the output of functions growth() and fordyn().



Evaluation metrics

The package provides functions to compare predictions with observations (use ?evaluation for details on how observations should be arranged).



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For example, a single evaluation metric can be calculated:

```
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## [1] 0.008684958
```



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For example, a single evaluation metric can be calculated:

```
evaluation_metric(S, exampleobs, type = "SWC", metric = "MAE")
## [1] 0.008684958
```

or many of them:

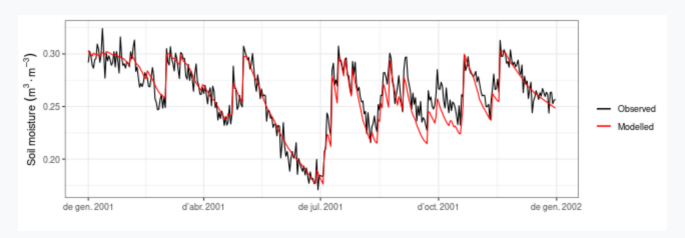
```
evaluation_stats(S, exampleobs, type = "SWC")
##
                          Bias
                                    Bias.rel
                                                       MAE
                                                                 MAE.rel
                                                                                                  NSF
               n
## 365.000000000
                 -0.003369394 -1.285026012
                                               0.008684958
                                                             3.312285595
                                                                           0.943031030
                                                                                         0.871877761
        NSF.abs
    0.634006817
##
```



Evaluation plots and interactive evaluation

Evaluation functions also allow visualizing the comparison as time series or scatter plots:

```
evaluation_plot(S, exampleobs, type = "SWC", plotType = "dynamics")
```

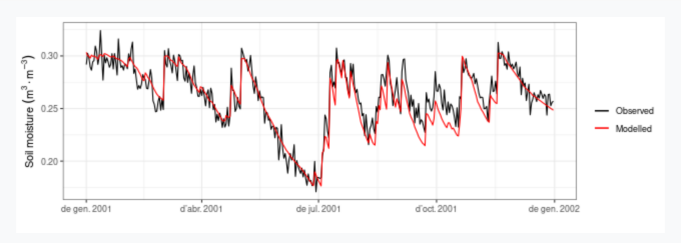




Evaluation plots and interactive evaluation

Evaluation functions also allow visualizing the comparison as time series or scatter plots:

```
evaluation_plot(S, exampleobs, type = "SWC", plotType = "dynamics")
```



Alternatively, the observed data can be supplied as an additional parameter to shinyplot() for interactive graphics including model evaluation:

```
shinyplot(S, exampleobs)
```



4. Advanced water/energy balance

Creating an input object for the advanced model

The most important step to run the advanced model is to specify the appropriate transpiration mode in the control parameters:

```
control <- defaultControl("Sperry")</pre>
```



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control <- defaultControl("Sperry")</pre>
```

We can build our input object for function spwb() using the same function as before:

```
x_adv <- forest2spwbInput(exampleforestMED, examplesoil, SpParamsMED, control)</pre>
```



Creating an input object for the advanced model

The most important step to run the advanced model is to specify the appropriate transpiration mode in the control parameters:

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control <- defaultControl("Sperry")</pre>
```

We can build our input object for function spwb() using the same function as before:

```
x_adv <- forest2spwbInput(exampleforestMED, examplesoil, SpParamsMED, control)</pre>
```

The water balance input object contains the same elements...



Creating an input object for the advanced model

... but the main difference with the basic model is in the number of parameters, e.g.:

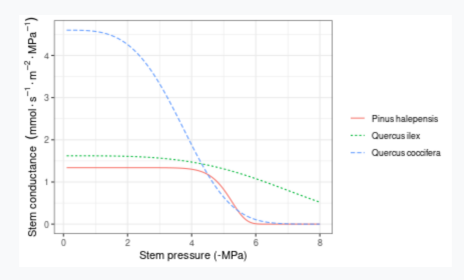
```
x_adv$paramsTranspiration
           Gswmin
                   Gswmax Vmax298 Jmax298 Kmax_stemxylem Kmax_rootxylem VCleaf_kmax VCleaf_c
##
## T1 148 0.003086667 0.2850000 72.19617 124.1687
                                              0.15
                                                          0.60
                                                                4.000000 11.137051
## T2_168 0.004473333 0.2007222 68.51600 118.7863
                                              0.40
                                                          1.60
                                                                4.000000 1.339370
## S1_165 0.010455247 0.2830167 62.78100 118.4486
                                              0.29
                                                          1.16
                                                                9.579077 1.844224
##
        ## T1_148 -2.380849
                 1.339563 12.710000 -5.290000
                                          1.505250 11.137051 -3.065569
                                                                    29078608
92609924
## S1_165 -3.030130
                 4.599269 3.537784 -4.126512
                                          4.660990 1.760307 -2.797092
                                                                   473105480
       Plant_kmax
##
## T1_148 0.6021000
## T2_168 0.6920994
## S1_165 1.8643961
```



Vulnerability curves

We can inspect *hydraulic vulnerability curves* (i.e. how hydraulic conductance of a given segment changes with the water potential) for each plant cohort and each of the different segments of the soil-plant hydraulic network:

```
hydraulics_vulnerabilityCurvePlot(x_adv, type="stem", speciesNames = TRUE)
```

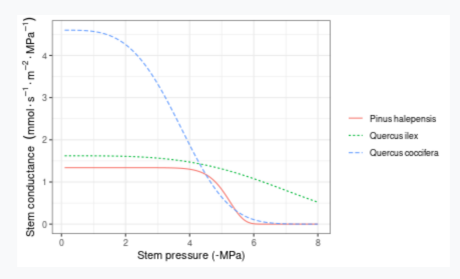




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```



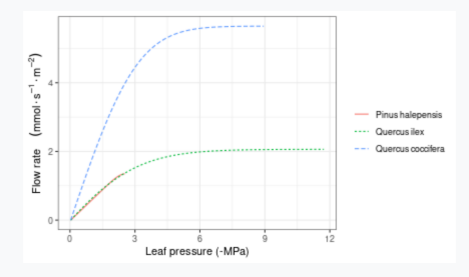
The maximum values and shape of vulnerability curves for leaves and stems are regulated by parameters in paramsTranspiration.



Supply functions

The vulnerability curves conforming the hydraulic network are used in the model to build the **supply function**, which relates water flow (i.e. transpiration) with the drop of water potential along the whole hydraulic pathway.

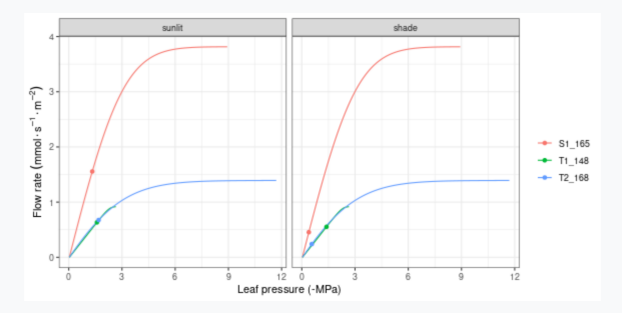
```
hydraulics_supplyFunctionPlot(x_adv, type="E", speciesNames = TRUE)
```





Stomatal regulation

Stomatal conductance is determined after building a photosynthesis function corresponding to the supply function and finding the value of stomatal conductance that maximizes carbon revenue while avoiding hydraulic damage: the *profit-maximization* approach.





Water/energy balance run for a single day

Since the model operates at a daily and sub-daily temporal scales, it is possible to perform soil water balance for one day only, by using function spwb_day():



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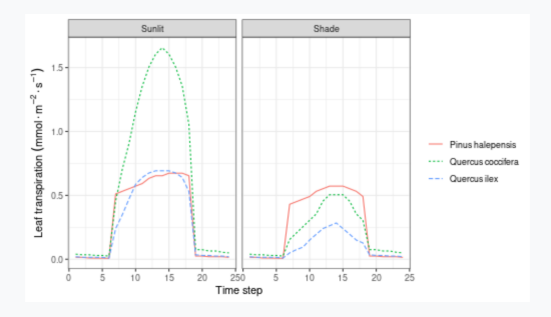
The output of spwb_day() is a list with several elements:

```
names(sd1)
                            "topography"
                                               "weather"
                                                                                       "EnergyBalance"
    [1] "cohorts"
                                                                   "WaterBalance"
                                                                                       "SunlitLeaves"
                            "Stand"
                                               "Plants"
                                                                   "RhizoPsi"
## [6] "Soil"
## [11] "ShadeLeaves"
                            "ExtractionInst"
                                               "PlantsInst"
                                                                   "SunlitLeavesInst" "ShadeLeavesInst"
## [16] "LightExtinction"
                            "LWRExtinction"
                                               "CanopyTurbulence"
```



Plotting single-day results

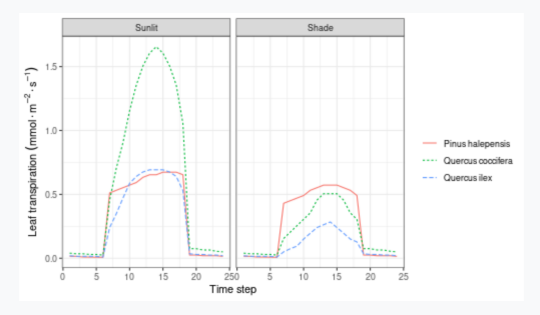
```
plot(sd1, type = "LeafTranspiration", bySpecies = TRUE)
```





Plotting single-day results

```
plot(sd1, type = "LeafTranspiration", bySpecies = TRUE)
```



More conveniently, you can examine multiple plots interactively:

```
shinyplot(sd1)
```



Resetting the input object

However, running spwb_day() modified the input object. In particular, the soil moisture at the end of the simulation was:

```
x_adv$soil$W
## [1] 0.9926237 0.9982876 0.9997178 0.9998916
```

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```
x_adv$soil$W
## [1] 0.9926237 0.9982876 0.9997178 0.9998916
```

We simply use function resetInputs() to reset state variables to their default values, so that the new simulation is not affected by the end state of the previous simulation:

```
resetInputs(x_adv)
x_adv$soil$W
## [1] 1 1 1 1
```



Water/energy balance run for multiple days

We can now run the advanced water balance model (which takes 1 min aprox.)

```
S_adv \leftarrow spwb(x_adv, examplemeteo, latitude = 41.82592, elevation = 100)
```



Water/energy balance run for multiple days

We can now run the advanced water balance model (which takes 1 min aprox.)

```
S_adv <- spwb(x_adv, examplemeteo, latitude = 41.82592, elevation = 100)
```

Function spwb() returns a list of class spwb, like the basic water balance model, but which contains more information:



Summaries, plots and interactive plots

Summaries and plots can be obtained from simulation results, using functions summary():

```
summary(S_adv, freq="months", output="PlantStress", bySpecies = TRUE)
```



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```
summary(S_adv, freq="months", output="PlantStress", bySpecies = TRUE)
and plot():
plot(S_adv, type="LeafPsiMin", bySpecies = TRUE)
```



Summaries, plots and interactive plots

Summaries and plots can be obtained from simulation results, using functions summary():

```
summary(S_adv, freq="months", output="PlantStress", bySpecies = TRUE)
and plot():
plot(S_adv, type="LeafPsiMin", bySpecies = TRUE)
```

Alternatively, one can interactively create plots using function shinyplot(), e.g.:

```
shinyplot(S_adv)
```



5. Modifying model inputs

Let's imagine one is not happy with a particular cohort parameter. For example, LAI estimates produced by forest2spwbInput() do not match known values:



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We should not manually modify x_{adv} because some parameters are related and we may break their relationship.



5. Modifying model inputs

Let's imagine one is not happy with a particular cohort parameter. For example, LAI estimates produced by forest2spwbInput() do not match known values:

We should not manually modify x_{adv} because some parameters are related and we may break their relationship.

Instead, we should use function modifyInputParams():

```
x_mod <- modifyInputParams(x_adv, c("T2_168/LAI_live" = 1.1))</pre>
```

which will display messages describing the parameters that are modified.

M.C. Escher - Night and day, 1938

