# DBEN\_testing\_all\_vars

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## **Summary:**

Some models already have biomass at disturbance year, while others have 0 values. We consider this a feature of the models and accept that some models therefore are on year 'ahead' in the dynamics.

For some models, some variables which have a value of 0 at disturbance year, but then have non-zero values for other variables. Please double-check. i.e. WBgrowth shows 0, but height is already at >0 values? That's of course plausible if height is initialised, but please clarify.

BiomeEP and BiomeES sometimes contain grasses in output that should be explicitly for wood. I can post-process the grasses 'away', but am wondering whether we in this case received output we asked for (i.e. WBgrowth). Please double-check.

Some of the jitteryness comes from the climate forcing. We may create a 30-year running mean to smoothen this in the paper. Other jitteryness I would like to dicuss for individual outputs below (see my comments below).

From Amsterdam: -still not really resolved for me: How dependent is the disturbance recovery on how "disturbance/bare ground" was set up?

NA = no output provided. I mention in the individual model TODOs sent via mail, whether I an output is needed. For ther rest, if you find you can easily add it, please go ahead for the next submission.

[TODOs for ANNEMARIE] JULES-RED sizeclass <1 currently set to 0, but is actually "NA". Right now seems like there are 0 trees, instead of no output. if(model name =="JULES-RED"){ df[,4] <- NULL

```
#knitr::opts_chunk$set(echo = TRUE)
library(ncdf4)
library(DGVMTools)
library(dplyr)
library(maps)
library(maptools)
library(gridExtra)
#source the format metadata for DBEN project to integrate with DGVMTools:
#source('Format-DBEN_paper1.R')
source("/Users/annemarie/Documents/1_TreeMort/2_Analysis/3_analysis_demographic_model_intercomparison/P
```

```
## Layer: BINE (Boreal/Temperate Shade-Intolerant Needleleaved Evergreen Tree), plot colour = dodgerblu
## With properties: =, growth.form=Tree, leaf.form=Needleleaved, phenology=Evergreen, shade.tolerance
## Layer: NE (Boreal/Temperate Needleleaved Evergreen Tree), plot colour = darkblue
## With properties: =, growth.form=Tree, leaf.form=Needleleaved, phenology=Evergreen, shade.tolerance
## Layer: IBS (Shade-intolerant Broadleaved Summergreen Tree), plot colour = chartreuse
## With properties: =, growth.form=Tree, leaf.form=Broadleaved, phenology=Summergreen, shade.toleranc
## Layer: TeBS ((intermediate) shade tolerant broadleaf deciduous), plot colour = purple
```

```
With properties: =, growth.form=Tree, leaf.form=Broadleaved, phenology=Summergreen, climate.zone=T
## Layer: TrIBE (Tropical Shade-intolerant Broadleaved Evergreen Tree), plot colour = orchid
     With properties: =, growth.form=Tree, leaf.form=Broadleaved, phenology=Evergreen, climate.zone=Tro
## Layer: TrBE (Tropical Broadleaved Evergreen Tree), plot colour = orchid4
     With properties: =, growth.form=Tree, leaf.form=Broadleaved, phenology=Evergreen, climate.zone=Tro
## Layer: TrBR (Tropical Broadleaved Raingreen Tree), plot colour = palevioletred
     With properties: =, growth.form=Tree, leaf.form=Broadleaved, phenology=Raingreen, climate.zone=Tro
## Layer: Grasses (C3 and/or C4 grasses), plot colour = dark grey
     With properties: =, growth.form=Grass, shade.tolerance=None, land.cover=Natural,
## Layer: Total (Total), plot colour = black
     With properties: type=Sum, land.cover=All,
## Layer: <1 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #00FF00
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <5 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #0AF102
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <10 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #14E405
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <15 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #1ED707
    With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <20 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #29C90A
    With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <30 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #33BCOD
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
##
## Layer: <40 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #3DAF0F
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <50 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #48A112
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <60 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #529415
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <70 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #5C8717
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <80 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #67791A
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <90 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #716C1C
    With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <100 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #7B5F1F
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <150 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #865122
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <200 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #904424
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <250 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #9A3727
    With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: 300+ (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #A52A2A
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: 1 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = light green
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: 2 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = dark green
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: 3 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = brown
    With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <1 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #00FF00
    With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
```

## Layer: <5 (sizeclass\_by\_diameter\_at\_breastheight (cm)), plot colour = #0AF102

```
With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <10 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #14E405
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <20 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #1ED707
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <30 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #29C90A
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <40 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #33BCOD
##
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <50 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #3DAF0F
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <60 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #48A112
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <70 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #529415</pre>
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <80 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #5C8717
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
##
## Layer: <90 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #67791A
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <100 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #716C1C
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <150 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #7B5F1F
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
##
## Layer: <200 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #865122
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <250 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #904424
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <300 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #9A3727</pre>
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: 300+ (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #A52A2A
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <1 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #00FF00
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <5 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #0BF002
    With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <10 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #16E205
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <15 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #21D408
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <20 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #2CC60B
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <30 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #37B80E
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <40 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #42A910
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <50 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #4D9B13
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <60 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #588D16
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <70 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #637F19
    With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <80 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #6E711C</pre>
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
```

## Layer: <90 (sizeclass\_by\_diameter\_at\_breastheight (cm)), plot colour = #79621E

```
With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <100 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #845421
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <150 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #8F4624
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: <200 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #9A3827
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
## Layer: >=200 (sizeclass_by_diameter_at_breastheight (cm)), plot colour = #A52A2A
     With properties: =, growth.form=Tree, leaf.form=Any, phenology=Any, climate.zone=Any, shade.tolera
                    cveg (Vegetation Carbon Mass by PFT): Units=kgC m-2, Defined for format: DBEN, CF
## Quantity:
## Quantity:
                    albedo (surface reflectiveness of PFTs): Units=-, Defined for format: DBEN, CF stan-
                    z0 (roughness length (momentum)): Units=m, Defined for format: DBEN, CF standard_na
## Quantity:
                    cwood (Wood Carbon Mass by PFT): Units=kgC m-2, Defined for format: DBEN, CF standa
## Quantity:
## Quantity:
                    AGcwood (Aboveground woody biomass): Units=kgC m-2, Defined for format: DBEN, CF st
## Quantity:
                    AGB (Aboveground total biomass): Units=kgC m-2, Defined for format: DBEN, CF standa
## Quantity:
                    cwood_size (Wood Carbon Mass by size class): Units=kgC m-2, Defined for format: DBE
                    nstem_size (Stem number by size class): Units=count ha-1, Defined for format: DBEN,
## Quantity:
## Quantity:
                    stemmort rate (Stem mortality rate by size class): Units=% yr-1, Defined for format
                    lai (Leaf area index): Units=m2 m-2, Defined for format: DBEN, CF standard_name=lea
## Quantity:
## Quantity:
                    CA (Crown area): Units=m2 ha-1, Defined for format: DBEN, CF standard_name=crown_ar
## Quantity:
                    BA (Basal area): Units=m2 ha-1, Defined for format: DBEN, CF standard_name=basal_ar
                    height (Mean PFT height in the 95% height percentile): Units=m, Defined for format:
## Quantity:
                    WBgrowth (Woody biomass growth): Units=kgC m-2 yr-1, Defined for format: DBEN, CF s
## Quantity:
                    BAgrowth (Basal area growth): Units=m2 ha-1 yr-1, Defined for format: DBEN, CF stan-
## Quantity:
## Quantity:
                    cmort (Wood Carbon Mass flux lost by size class): Units=kgC m-2 s-1, Defined for for
## Quantity:
                    cmort_size (Wood Carbon Mass flux lost by size class): Units=kgC m-2 s-1, Defined f
                    cmort_rate (% Wood Carbon Mass lost due to mortality, by size class): Units=% yr -1
## Quantity:
                    stemmort (Stem number flux lost by size class): Units=Count ha-1 yr-1, Defined for
## Quantity:
                    stemmort_size (Stem number flux lost by size class): Units=Count ha-1 yr-1, Defined
## Quantity:
                    gpp (gpp): Units=kgC m-2 yr-1, Defined for format: DBEN, CF standard_name=Gross_pri
## Quantity:
## Quantity:
                    npp (npp): Units=kgC m-2 yr-1, Defined for format: DBEN, CF standard_name=Net_prima
## Quantity:
                    nbp (Net Biospheric Production): Units=kgC m-2 yr-1, Defined for format: DBEN, CF s
## Quantity:
                    cveg_dyn (Total Carbon mass Flux by Pft): Units=kgC m-2 yr-1, Defined for format: Di
                    BA_size (BA by sizeclass): Units=m2 ha-1, Defined for format: DBEN, CF standard_nam
## Quantity:
## Quantity:
                    stemmort_freezing (Stems lost due to freezing mortality): Units=count ha-1, Defined
                    stemmort_freezing_rate (% stems lost due to freezing mortality): Units=% yr-1, Defi
## Quantity:
                    stemmort_cstarv (Stems lost due to carbon starvation mortality): Units=count ha-1,
## Quantity:
## Quantity:
                    stemmort_cstarv_rate (% Stems lost due to carbon starvation mortality): Units=% yr-
                    stemmort_background (Stems lost due to background mortality): Units=count ha-1, Def
## Quantity:
                    stemmort_background_rate ( %Stems lost due to background mortality): Units=% yr-1,
## Quantity:
                    stemmort impact (Stems lost duee to impact mortality): Units=count ha-1, Defined fo
## Quantity:
                    stemmort_impact_rate (#Stems lost duee to impact mortality): Units=% yr-1, Defined
## Quantity:
                    stemmort_termination (Stems lost due to termination mortality): Units=count ha-1, D
## Quantity:
## Quantity:
                    stemmort_termination_rate (% Stems lost due to termination mortality): Units=% yr-1
                    stemmort_hydro (Stems lost due to hydraulic failure(?) mortality): Units=count ha-1
## Quantity:
                    stemmort_hydro_rate (% Stems lost due to hydraulic failure(?) mortality): Units=% y
## Quantity:
                    cmort_cstarv (Wood Carbon Mass flux lost through carbon starvation mortality): Unit
## Quantity:
   carbon starvation mortality
## Quantity:
                    cmort_cstarv (%Wood Carbon Mass flux lost through carbon starvation mortality): Uni
   carbon starvation mortality
                    cmort_cstarv_rate ( % Wood Carbon Mass lost through carbon starvation mortality): U
## Quantity:
## carbon starvation mortality
## Quantity:
                    cmort_hydro (Wood Carbon Mass flux lost through hydarulic failur mort.): Units=m2 h
```

## hydraulic failure mortality

```
## Quantity:
                    cmort_hydro_rate (% Wood Carbon Mass flux lost through hydarulic failur mort.): Uni
## hydraulic failure mortality
                    cmort res (Wood Carbon Mass flux lost due to Resource mortality): Units=kg C m-2 s-
## Quantity:
                    cmort_res_rate (% Wood Carbon Mass lost, Resource mortality): Units=% yr-1, Defined
## Quantity:
## Quantity:
                    cmort_crowd (Wood Carbon Mass flux lost due to crowding mortality): Units=kg C m-2
                    cmort_crowd_rate ( % Wood Carbon Mass lost due to crowding mortality): Units=% yr
## Quantity:
## Quantity:
                    cmort dist (Wood Carbon Mass flux lost due to disturbance mortality): Units=kg C m-
## Quantity:
                    cmort_dist_rate (% Wood Carbon Mass lost, disturbance mortality): Units=% yr-1, Def
## Quantity:
                    cmort_age (Wood Carbon Mass flux lost through old age mortality): Units=kg C m-2 s-
                    cmort_age_rate (% Wood Carbon Mass lost, old age mortality): Units=% s-1, Defined f
## Quantity:
## Quantity:
                    cmort_dist (Wood Carbon Mass flux lost through disturbance mortality): Units=kg C m
                    cmort_dist_rate (% Wood Carbon Mass lost through disturbance mortality): Units=% s-
## Quantity:
                    cmort_fire (Wood Carbon Mass flux lost through fire mortality): Units=kg C m-2 s-1,
## Quantity:
## Quantity:
                    cmort_fire_rate (% Wood Carbon Mass lost, fire mortality): Units=% s-1, Defined for
## Quantity:
                    cmort_greff (Wood Carbon Mass flux lost, low growth efficiency mortality): Units=kg
## Quantity:
                    cmort_greff_rate (% Wood Carbon Mass lost, growth efficiency mortality): Units=% s-
                    cmort_other (Wood Carbon Mass flux lost, other mortality mechs.): Units=kg C m-2 s-
## Quantity:
                    cmort_res (Wood Carbon Mass flux lost, other mortality mechs.): Units=kg C m-2 s-1,
## Quantity:
## Quantity:
                    cmort_other_rate ( % Wood Carbon Mass lost, other mortality mechs.): Units=kg C m-2
                    stemmort_fire (Number of stems lost through fire mortality): Units=count ha-1, Defi
## Quantity:
## Quantity:
                    stemmort_fire_rate (fire mortality rate): Units=% yr-1, Defined for format: DBEN, C
## Quantity:
                    stemmort_greff (Number of stems lost through low growth efficiency mortality): Unit
                    stemmort_greff_rate (low growth efficiency mortality rate): Units=% yr-1, Defined f
## Quantity:
## Quantity:
                    stemmort_other (Number of stems lost through other mortality mechs.): Units=count h
                    stemmort_other_rate (mortality rate, other mechs.): Units=% yr-1, Defined for forma
## Quantity:
## Quantity:
                    stemmort_age (Number of stems lost through age related mortality): Units=count ha-1
## Quantity:
                    stemmort_age_rate (old age related mortality rate): Units=% yr-1, Defined for forma
                    stemmort_dist (Number of stems lost through disturbance mortality): Units=count ha-
## Quantity:
                    stemmort_dist_rate ( disturbance mortality rate): Units=% yr-1, Defined for format:
## Quantity:
#conversion helpers:
year_{to}_{seconds} = 60*60*24*365
seconds_to_year = 1/year_to_seconds
create_figs = TRUE
Figs dir <- "/Users/annemarie/Documents/1 TreeMort/2 Analysis/3 analysis demographic model intercompari
## set all file directories here:
file.dir.fates
                  <- "/Users/annemarie/Documents/1_TreeMort/2_Analysis/3_analysis_demographic_model_int</pre>
file.dir.lpjguess <- "/Users/annemarie/Documents/1_TreeMort/2_Analysis/3_analysis_demographic_model_int
file.dir.orchidee <- "/Users/annemarie/Documents/1_TreeMort/2_Analysis/3_analysis_demographic_model_int
file.dir.cablepop <- "/Users/annemarie/Documents/1_TreeMort/2_Analysis/3_analysis_demographic_model_int
file.dir.julesred <- "/Users/annemarie/Documents/1_TreeMort/2_Analysis/3_analysis_demographic_model_int
file.dir.biomeEP <- "/Users/annemarie/Documents/1_TreeMort/2_Analysis/3_analysis_demographic_model_inte
file.dir.biomeE_Standalone <- "/Users/annemarie/Documents/1_TreeMort/2_Analysis/3_analysis_demographic_
#load observations
obs.path <- "/Users/annemarie/Documents/1_TreeMort/2_Analysis/3_analysis_demographic_model_intercompari
get_output_LPJGUESS <- function(site,run,var,file.dir,co2_levels = "PS_412ppm/"){</pre>
  model_name = "LPJGUESS"
  if(co2 levels == "PS 562ppm/"){
   file.dir. = pasteO(file.dir, "PS_562ppm/")
```

```
}else{
    file.dir = paste0(file.dir, "PS_412ppm/")
     # set metadata
      source.in <- defineSource(id = site,
                            dir = file.dir ,
                            format = DBEN,
                            name = paste(model_name, site,'-', run),
                            forcing.data = "cru_jra2.2")
      var.to.plot <- getField_DBEN(source.in,</pre>
                                 quant = get quantity(var),
                                 file.name = pasteO(file.dir,
                                  model_name,"_",var,"_",run,"_",site,".nc"),
                                     model_name = model_name)
      # change metadata
      var.to.plot@first.year <- 1 # first year of simulation</pre>
      var.to.plot@last.year <- dim(var.to.plot@data)[1]</pre>
       # change real data:
      var.to.plot@data$Year <- seq(1,dim(var.to.plot@data)[1])</pre>
            #[TODO] probably change unit later in actual model output and for dben standard..
       if(var == "cmort" | var == "cmort age" |
           var == "cmort_other" | var == "cmort_greff" | var == "nbp") {
         #convert to -yr:
          end <- dim(var.to.plot@data)[2]</pre>
          var.to.plot@data[,4:end] <- var.to.plot@data[,4:end] * 1/year_to_seconds</pre>
          var.to.plot@quant@units <- "kgC m-2 yr-1"</pre>
       }
     if(var =="gpp" ){
        var.to.plot <- change_unit_stoy(var.to.plot)</pre>
      return(var.to.plot)
}
get_output_BiomeEP <- function(site,run,var,file.dir,co2_levels){</pre>
 model_name = "BiomeEP"
  #site = c("Fi1")
  \#run = c("p0")
     # set metadata
      source.in <- defineSource(id = site,</pre>
                            dir = file.dir ,
                            format = DBEN,
                            name = paste(model_name, site,'-', run),
                            forcing.data = "cru_jra2.2")
      #adress the fact that for filename with var = "cmort_size", the variable name within = "cmort"
      if(var== "cmort_size"){
         var.to.plot <- getField_DBEN(source.in,</pre>
```

```
quant = get_quantity("cmort"),
                                 file.name = pasteO(file.dir,co2_levels,"/",site,"/",
                                 model_name,"_",var,"_",run,"_",site,"_",co2_levels,".nc"),
                                 model name = model_name)
      }else if(var == "stemmort_size"){ # see above.
        var.to.plot <- getField_DBEN(source.in,</pre>
                                 quant = get_quantity("stemmort"),
                                 file.name = pasteO(file.dir,co2_levels,"/",site,"/",
                                 model_name,"_",var,"_",run,"_",site,"_",co2_levels,".nc"),
                                 model name = model name)
      }else{ # normal procedure:
         var.to.plot <- getField_DBEN(source.in,</pre>
                                 quant = get_quantity(var),
                                 file.name = pasteO(file.dir,co2_levels,"/",site,"/",
                                 model_name,"_",var,"_",run,"_",site,"_",co2_levels,".nc"),
                                 model_name = model_name)
      }
            #[TODO] probably change unit later...
       if(var == "cmort" | var == "cmort_age" |
           var == "cmort_other" | var == "cmort_greff") {
         #convert to -yr:
          end <- dim(var.to.plot@data)[2]</pre>
          var.to.plot@data[,4:end] <- var.to.plot@data[,4:end] * 1/year_to_seconds</pre>
          var.to.plot@quant@units <- "kgC m-2 yr-1"</pre>
       }
     # if(var =="npp"){
      # var.to.plot <- change_npp_unit(var.to.plot)</pre>
      #}
      return(var.to.plot)
}
get_output_BiomeE_standalone <- function(site,run,var,file.dir,co2_levels){</pre>
  model_name = "BiomeE"
  if(co2_levels=="562ppm"){
    co2_levels ="eCO2"
  if(co2_levels =="412ppm"){
     co2_levels ="aCO2"
  \#run = c("p0")
     # set metadata
      source.in <- defineSource(id = site,</pre>
                            dir = file.dir.biomeE Standalone ,
                            format = DBEN,
                            name = paste(model_name, site,'-', run),
                            forcing.data = "cru_jra2.2")
        var.to.plot <- getField_DBEN(source.in,</pre>
                                        quant = get_quantity(var),
                                        file.name = pasteO(file.dir,model_name,"_PS_",
                                                            site,"_",co2_levels,
```

```
"_",run,"_",var,".nc") ,
                                       model_name = model_name)
      #pasteO(file.dir,model_name,"_",run,"_",site,"_",co2_levels,"_00_",var,".nc")
   # /Users/annemarie/Documents/1_TreeMort/2_Analysis/3_analysis_demographic_model_intercomparison/Pape
            #[TODO] probably change unit later...
       if(var == "cmort" | var == "cmort_age" |
           var == "cmort_other" | var == "cmort_greff") {
         #convert to -yr:
          end <- dim(var.to.plot@data)[2]</pre>
          var.to.plot@data[,4:end] <- var.to.plot@data[,4:end] * 1/year_to_seconds</pre>
          var.to.plot@quant@units <- "kgC m-2 yr-1"</pre>
       }
      return(var.to.plot)
get_output_CABLEPOP <- function(site,run,var,file.dir,co2_levels){</pre>
 model name="CABLE-POP"
  # flexibly account for file naming:
  if(run == "0"){
    #benchmark run
    file.name = pasteO(file.dir,co2_levels,"/",model_name,"_",var,"_PO_",site,".nc")
  }else{ # sensitivity runs:
    file.name = paste0(file.dir,co2_levels,"/",model_name,"_",var,"_PS_",site,"_",run,".nc")
  }
     # set metadata
      source.in <- defineSource(id = site,</pre>
                            dir = file.dir ,
                            format = DBEN,
                            name = paste(model_name, site,'-', run),
                            forcing.data = "cru_jra2.2")
      var.to.plot <- getField_DBEN(source.in,</pre>
                                 quant = get_quantity(var),
                                 file.name = file.name,
                                 model_name="CABLE-POP")
      # CABLE-POP_AGcwood_PS_BCI_1.nc
      # split up size classes graph to make more readable.
      # create shared maximum y axis to make graphs more interpretable. maybe change back later
      ymax <- max(var.to.plot@data$Total)</pre>
     # print(plotTemporal(var.to.plot))#y.scale_log = TRUE))
     # p <- plotTemporal(var.to.plot)</pre>
      # [TODO] tidy...
    c("cmort_crowd", "cmort_dist", "cmort_res")
       #[TODO] probably change unit later...
```

```
if(var == "cmort" | var == "cmort_crowd" | var == "cmort_dist" | var == "cmort_res"){
          var.to.plot <- change_unit_stoy(var.to.plot)</pre>
      if(var =="npp"|var =="gpp" |var =="nbp"){
        var.to.plot <- change_unit_stoy(var.to.plot)</pre>
return(var.to.plot)
get_output_JULESRED <- function(site,run,var,file.dir,co2_levels){</pre>
     model name = "JULES-RED"
     # set metadata
      source.in <- defineSource(id = site,</pre>
                            dir = file.dir ,
                            format = DBEN,
                            name = paste(model_name, site,'-', run),
                            forcing.data = "cru_jra2.2")
      var.to.plot <- getField_DBEN(source.in,</pre>
                                 quant = get_quantity(var),
                                 file.name = pasteO(file.dir,
                                  model_name,"_",var,"_PS_",co2_levels,"_",site,"_",run,".nc"),
                                 model_name = model_name)
      # split up size classes graph to make more readable.
      # create shared maximum y axis to make graphs more interpretable. maybe change back later
      ymax <- max(var.to.plot@data$Total)</pre>
      #print(plotTemporal(var.to.plot))#y.scale_log = TRUE))
      p <- plotTemporal(var.to.plot)</pre>
            #[TODO] probably change unit later...
       if(var == "cmort" | var == "cmort_age" |
           var == "cmort_other" | var == "cmort_greff") {
         #convert to -yr:
          end <- dim(var.to.plot@data)[2]</pre>
          var.to.plot@data[,4:end] \leftarrow var.to.plot@data[,4:end] * 1/year_to_seconds
          var.to.plot@quant@units <- "kgC m-2 yr-1"</pre>
       }
       if(var =="npp"|var =="gpp" |var =="nbp"){
        var.to.plot <- change_unit_stoy(var.to.plot)</pre>
       }
return(var.to.plot)
get_output_FATES <- function(site,run,var,file.dir,model_name){</pre>
model name ="FATES"
if(site == "FIN"){siteFates="fi"}
if(site == "BIA"){siteFates ="bia"}
if(site == "BCI"){siteFates ="bci"}
```

```
if(run == "P0"){runFates ="p0"}
     # set metadata
      source.in <- defineSource(id = site,</pre>
                            dir = file.dir ,
                            format = DBEN,
                            name = paste(model_name, site,'-', run),
                            forcing.data = "cru_jra2.2")
# obtain total stemmort from under and overstory:
        if(var == "stemmort"){ #
             var.to.plot_under <- getField_DBEN(source.in,</pre>
                                 quant = get_quantity(var),
                                  file.name = pasteO(file.dir,site,"/",runFates,"/",
                                                    model_name,"_",var,"_",
                                                    runFates,"_",siteFates,"_understsory.nc"),
                                 model_name = model_name)
              var.to.plot_over <- getField_DBEN(source.in,</pre>
                                 quant = get_quantity(var),
                               file.name = pasteO(file.dir,site,"/",runFates,"/",
                                                   model_name,"_",var,"_",
                                                   runFates,"_",siteFates,"_overstsory.nc"),
                               model_name = model_name)
              var.to.plot <- var.to.plot_under # "initalise" to keep metadata</pre>
              var.to.plot@data[,4:20] <- var.to.plot_under@data[,4:20] + var.to.plot_over@data[,4:20]</pre>
              #print(plotTemporal(var.to.plot))
                if(create_figs == TRUE){
                   jpeg(filename = paste0(Figs_dir,paste(var, run, site, model_name,sep="_"),".jpeg") )
                   print(plotTemporal(var.to.plot))
                   dev.off()
                }
        }else{
          var.to.plot <- getField_DBEN(source.in,</pre>
                                 quant = get_quantity(var),
                                 file.name = pasteO(file.dir,site,"/",runFates,"/",
                                 model_name,"_",var,"_",runFates,"_",siteFates,".nc"),
                                 model_name = model_name)
          #[TODO] probably change unit later...
      if(var =="npp"|var =="gpp"|var =="nbp"){
        var.to.plot <- change_unit_stoy(var.to.plot)</pre>
        } # done reading in variables
        return(var.to.plot)
}
#temporary function, to change units for some fluxes where needed
change_unit_stoy <- function(var_to_plot){</pre>
          year_to_seconds = 31536000
```

```
#convert to -yr:
          end <- dim(var_to_plot@data)[2]</pre>
          var_to_plot@data[,4:end] <- var_to_plot@data[,4:end] * year_to_seconds</pre>
          var_to_plot@quant@units <- "kgC m-2 yr-1"</pre>
          return(var_to_plot)
         }
#aesthetics for consistent plotting
#x allowed to be biome name or site name, for more convenient use
get_biome_colour <- function(x){</pre>
  #preparation:
  biomes <- c("Boreal", "Temperate", "Tropics")</pre>
  sites <- c("FIN", "BIA", "BCI")</pre>
  cols <- c(1,1,1)#c("dark green","light green","brown")</pre>
  #automatic colour coding for biome based on site name:
  if(x %in% sites){
  #retrieval:
  ret col <- cols[sites==x]</pre>
  }
  if(x %in% c("Boreal", "Temperate", "Tropics")){
  #retrieval:
  ret_col <- cols[biomes==x]</pre>
  }
 return(ret_col)
}
#turn into object with function that returns colour when giving it model-name later..
get_model_colour <- function(model){</pre>
  # PREPARATION:
 models <- c("FATES", "JULES-RED", "ORCHIDEE", "LPJ-GUESS", "CABLE-POP", "BiomeEP", "BiomeE-Standalone")
 model_cols <- c("orange","red","blue","light blue","grey","purple","magenta")</pre>
 plot_mod_cols <- data.frame(models,model_cols)</pre>
  #RETRIEVAL:
 ret_col <- plot_mod_cols[which(plot_mod_cols$models == model), ]$model_cols</pre>
  return(ret_col)
  }
#stand-structure related shared objects:
sc <- c( "<1" , "<5" , "<10", "<15", "<20" , "<30", "<40" , "<50" , "<60" , "<70" , "<80" ,
#to deal with different file naming conventions:
runs = c("P1","P2","P3","P4","P5","P6","P7")
```

```
runs_biomeEP <- c("P0","PS1","PS2","PS3","PS4","PS5","PS6")</pre>
#Disturbance: stand-replacing (resetting to initial conditions) and stochastic with mean frequency
#of:0.01, 0.02, 0.04, 0.08, 0.20, 0.40 (corresponding to the file names: _01, _02, _04, _08, _20, _40)
runs_biomeES <- c("00","01", "02", "04", "08", "20", "40")
runs_julesred = c("1","2","3","4","5","6","7")
runs_cablepop = c("1","2","3","4","5","6","7")
#[TODO] re-naming of biomeEP scenarios once PS1 is run.
#biome-specific points on graphs:
Tropics_pch = 16
Temperate_pch = 16
Boreal_pch = 16
# function to plot a model's stand stucture output alongside observations.
# nstem_size can be plotted on log yaxis (log = TRUE) or not
# ylim_ext can override automatically generated ylim values.
# model name must match the model name used for qet_model_colour
# var the DBEN variable that should be plotted.
stand_structure_benchmarks <- function(model_name = "LPJ-GUESS", site_in = site, log =TRUE, var_in = var,
 if(model_name == "LPJ-GUESS"){
 if(model name == "BiomeEP"){
   model_out <- get_output_BiomeEP(site = site_in,run = "PO",var = var_in,file.dir = file.dir.biomeEP,</pre>
 if(var_in =="nstem_size"){
    ##plot models - nstem_size
   dbh_classes_plot <- stand_structure_obs[which(stand_structure_obs$site == site),]$dbh_classes_num
    #subset of dbh- classes for plotting, because in the observations there are no more than that.
   #[TODO]would ideally subset with this vector of column names, but it somehow doesnt, work. doing th
   dbh_classes_sel \leftarrow c( "<1" ,  "<5" ,  "<10" ,  "<15" ,  "<20" ,  "<30" ,  "<40" ,  "<50" ,  "<60" ,
   #make ylims flexible, or prescribe from external:
   max_model_output <- max(model_out@data[,</pre>
                                           c( "<1" , "<5" , "<10", "<15", "<20" , "<30".
                                                                                                "<40
                                              "<90" , "<100", "<150" , "<200", ">=200")])
   min_model_output <- max(model_out@data[,</pre>
                                           c("<1", "<5", "<10", "<15", "<20", "<30",
                                                                                                "<40
                                              "<90", "<100", "<150", "<200", ">=200")])
   max_obs <- max( na.omit(stand_structure_obs[,c("AGB_size_kgCm.2","AGB_size_upper_kgCm.2","AGB_size_</pre>
   min_obs <- min( na.omit(stand_structure_obs[,c("AGB_size_kgCm.2","AGB_size_upper_kgCm.2","AGB_size_
    #check if external ylims are present or not:
   if(is.null(ylim_ext)){ # not present, obtain ylim_set from within
    ylim_set = c(min(min_obs,min_model_output),max(max_obs,max_model_output))
```

```
}else{
ylim_set = ylim_ext # present. pass external ylims to ylim_set
  #if nstem should be logged:
if(log ==TRUE){
 if(is.null(ylim ext)){ # not present, obtain ylim set from within
 ylim_set = log(ylim_set)
 ylim_set[ylim_set ==-Inf] <- -2</pre>
 plot(stand_structure_obs[which(stand_structure_obs$site == site),]$dbh_classes_num,
    log(model_out@data[model_out@data$Year == max(model_out@data$Year),
                      ylim = ylim_set ,col= get_model_colour(model_name),pch=16,cex=1.5,ylab="")
  points(stand_structure_obs[which(stand_structure_obs$site == site),]$dbh_classes_num,
      log(stand_structure_obs[which(stand_structure_obs$site == site),]$nstem_size_ha.1),
      col= get_biome_colour(site))
  arrows(stand structure obs$dbh classes num,
      log(stand_structure_obs[which(stand_structure_obs$site == site),]$nstem_size_lower_ha.1),
      stand_structure_obs$dbh_classes_num,
      log(stand_structure_obs[which(stand_structure_obs$site == site),]$nstem_size_upper_ha.1),
       length = 0.00, angle = 90, code = 3,
       col= get_biome_colour(site) )
  mtext(site,adj=0.95,side=3,line=-1.3)
  mtext(side=2,"log(nstem_size) (nstem)",line=2,outer=FALSE)
}else { # nstem not logged
  plot(stand_structure_obs[which(stand_structure_obs$site == site),]$dbh_classes_num,
   "<10". "<1
                                         "<90", "<100", "<150", "<200", ">=200")],
    ylim = ylim_set ,col= get_model_colour(model_name),pch=16,cex=1.5)
  points(stand_structure_obs[which(stand_structure_obs$site == site),]$dbh_classes_num,
      stand_structure_obs[which(stand_structure_obs$site == site),]$nstem_size_ha.1,
      col= get biome colour(site))
  arrows(stand structure obs$dbh classes num,
      stand_structure_obs[which(stand_structure_obs$site == site),]$nstem_size_lower_ha.1,
      stand_structure_obs$dbh_classes_num,
      stand_structure_obs[which(stand_structure_obs$site == site),]$nstem_size_upper_ha.1,
       length = 0.00, angle = 90, code = 3,
       col= get_biome_colour(site) )
  mtext(site,adj=0.95,side=3,line=-1.3)
mtext(side=2, "nstem_size (nstem)", line=2, outer=FALSE)
}
```

```
}
   if(var in =="cwood size"){
       # cwood plots
      "<30", "<40", "<50", "<60", "<70", "<80", "<90",
   max obs <- max( na.omit(stand structure obs[,c("AGB size kgCm.2","AGB size upper kgCm.2","AGB size
    #check if external ylims are present or not:
   if(is.null(ylim_ext)){ # not present, obtain ylim_set from within
     ylim_set = c(0,max(max_obs,max_model_output) + max(max_obs,max_model_output)*0.1)
   }else{
     ylim_set = ylim_ext # present. pass external ylims to ylim_set
   plot(stand_structure_obs[which(stand_structure_obs$site == site),]$dbh_classes_num,
       "<30",
                                                                         "<40" , "<50" ,
        vlim = ylim set ,col= get model colour(model name),pch=16,cex=1.5,ylab="")
   points(stand_structure_obs[which(stand_structure_obs$site == site),]$dbh_classes_num,
         stand_structure_obs[which(stand_structure_obs$site == site),]$AGB_size_kgCm.2, col= get_bios
   arrows(stand_structure_obs$dbh_classes_num,
         stand_structure_obs[which(stand_structure_obs\site == site),]\sharpartage AGB_size_lower_kgCm.2,
         stand_structure_obs$dbh_classes_num,
         stand_structure_obs[which(stand_structure_obs$site == site),]$AGB_size_upper_kgCm.2,
          length = 0.00, angle = 90, code = 3,
         col= get_biome_colour(site) )
   mtext(site,adj=0.95,side=3,line=-1.3)
   mtext(side=2, "AGcwood (kgC m-2)", line=2, outer=FALSE)
   }
}
#convenience function for grid.arrange
#passes back ggplot object for each model_output (input)
create_gobj <-function(model_out, y.lim = NULL){</pre>
 p_out <- plotTemporal(model_out, legend.position = "none", title = model_out@source@name, y.lim )</pre>
return(p_out)
```

## Individual variables

Plotted for all model variables, in order of occurrence in the protocol table. Sometimes I have questions to the model outputs. Please, everyone if you come across something fishy, point it out, and the people who know the models, please feel responsible in answering the questions.

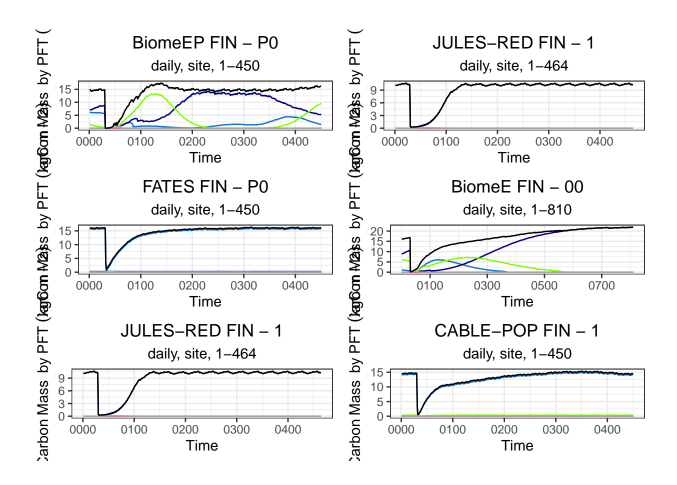
cwood\_size and nstem\_size are not plotted here.

#### cveg

```
Model Comments

BiomeEP
LPJ-GUESS
FATES AHES: Why is eveg not 0 at disturbance year?
BiomeES AHES: Why is eveg not 0 at disturbance year?AHES: -> Grasses?
JULES-RED AHES: Why is eveg not 0 at disturbance year?
```

```
var = "cveg"
site ="FIN"
run ="P0"
BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
pBiomeEP <- create_gobj(BiomeEP)</pre>
LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create gobj(LPJGUESS)</pre>
FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name ="FATES")
pFATES <- create_gobj(FATES)</pre>
BiomeES <- get_output_BiomeE_standalone(site, run = "00", var = var, file.dir.biomeE_Standalone, co2_level
## [1] "vobjtovarid4: **** WARNING **** I was asked to get a varid for dimension named PFT BUT this dim
pBiomeES <- create_gobj(BiomeES)</pre>
JULESRED <- get_output_JULESRED(site,run ="1",var = var,file.dir.julesred,co2_levels = "412ppm")
pJULESRED <- create_gobj(JULESRED)</pre>
CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)</pre>
grid.arrange(pBiomeEP,pJULESRED,pFATES,pBiomeES,pJULESRED,pCABLEPOP)
```



### **AG**cwood

Model	Comments
BiomeEP LPJ-GUESS FATES	
BiomeES	AHES: no AGcwood variable. Right now we are not comparing like-for-like here. What do we do for the paper: argue with low leaf carbon content ("The biomass of leaves is low, 0.2~0.4 kg C m-2.") or remove leaf fraction?
JULES-RED	AHES: Why does this not go to 0 at distyear?

```
var = "AGcwood"
site ="FIN"
run ="P0"
BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
pBiomeEP <- create_gobj(BiomeEP)

LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)

FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name ="FATES")</pre>
```

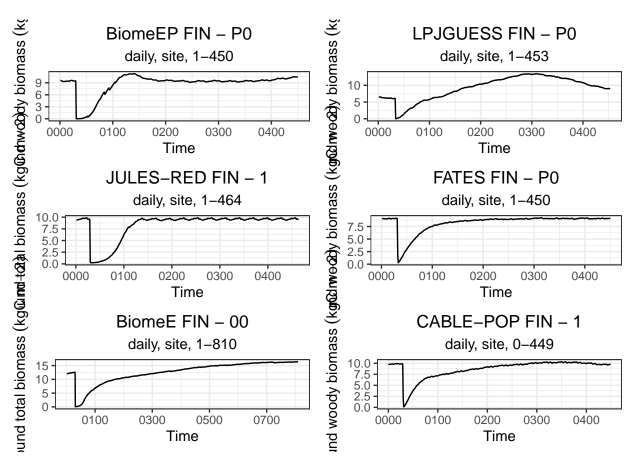
```
pFATES <- create_gobj(FATES)

#AHES: AGB not AGcwood
BiomeES <- get_output_BiomeE_standalone(site,run = "00",var = "AGB", file.dir.biomeE_Standalone,co2_lev
pBiomeES <- create_gobj(BiomeES)

#AHES: AGB here AGcwood - note that the automated plotting will provide the wrong metadata for this plo
JULESRED <- get_output_JULESRED(site,run ="1",var = "AGB",file.dir.julesred,co2_levels = "412ppm")
pJULESRED <- create_gobj(JULESRED)

CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)

grid.arrange(pBiomeEP,pLPJGUESS,pJULESRED,pFATES,pBiomeES,pCABLEPOP)</pre>
```



### cwood

Model Comments

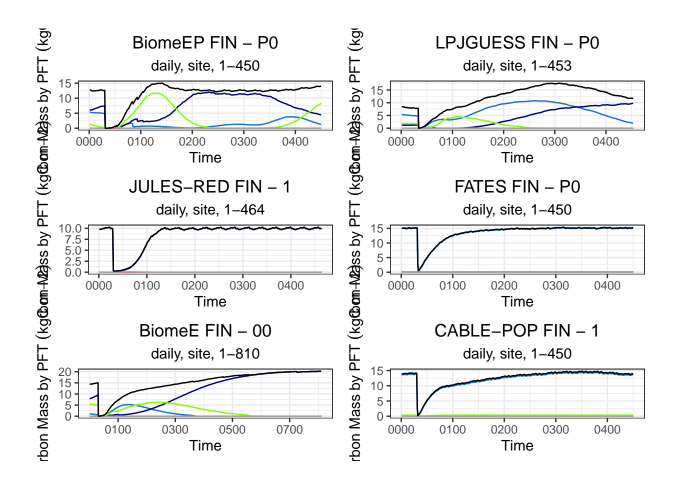
BiomeEP LPJ-GUESS FATES

AHES: Why does this not go to 0 at distyear?

Model	Comments
BiomeES	AHES: cwood goes to 0, so in other plots i.e. eveg
	the non-0 amount is indeed cause by the grass
	fraction ( makes sense to me)
JULES-RED	AHES: Why does this not go to 0 at distyear?

```
var = "cwood"
site ="FIN"
run ="P0"
BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
pBiomeEP <- create_gobj(BiomeEP)</pre>
LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)</pre>
FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name ="FATES")</pre>
pFATES <- create_gobj(FATES)</pre>
BiomeES <- get_output_BiomeE_standalone(site, run = "00", var = var, file.dir.biomeE_Standalone, co2_level
## [1] "vobjtovarid4: **** WARNING **** I was asked to get a varid for dimension named PFT BUT this dim
pBiomeES <- create_gobj(BiomeES)</pre>
JULESRED <- get_output_JULESRED(site, run ="1", var = var, file.dir.julesred, co2_levels = "412ppm")
pJULESRED <- create_gobj(JULESRED)</pre>
CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)</pre>
```

grid.arrange(pBiomeEP,pLPJGUESS,pJULESRED,pFATES,pBiomeES,pCABLEPOP)



lai

Model	Comments
BiomeEP	
LPJ-GUESS	
FATES	Why is LAI never 0?
BiomeES	Why is LAI never 0? -> Grass fraction?
JULES-RED	Why is LAI never 0?
CABLE-POP	Why is LAI never 0?

```
var = "lai"
site ="FIN"
run ="PO"
BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
pBiomeEP <- create_gobj(BiomeEP)

LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)

FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name ="FATES")
pFATES <- create_gobj(FATES)

BiomeES <- get_output_BiomeE_standalone(site,run = "00",var = var, file.dir.biomeE_Standalone,co2_level</pre>
```

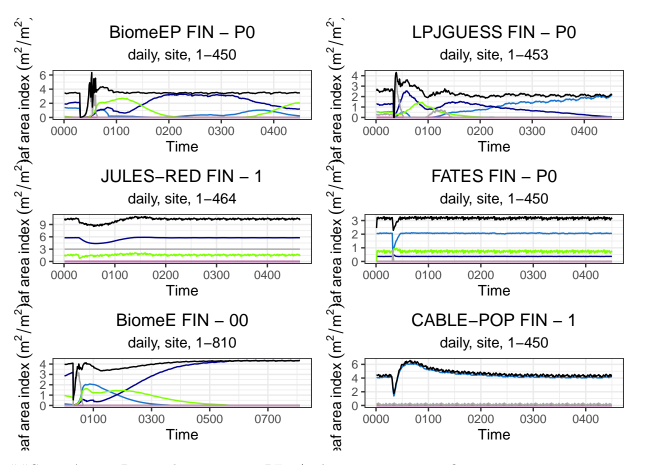
## [1] "vobjtovarid4: \*\*\*\* WARNING \*\*\*\* I was asked to get a varid for dimension named PFT BUT this dim

```
pBiomeES <- create_gobj(BiomeES)

JULESRED <- get_output_JULESRED(site,rum ="1",var = var,file.dir.julesred,co2_levels = "412ppm")
pJULESRED <- create_gobj(JULESRED)

CABLEPOP <- get_output_CABLEPOP(site,rum = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)

grid.arrange(pBiomeEP,pLPJGUESS,pJULESRED,pFATES,pBiomeES,pCABLEPOP)</pre>
```



##Crown Area — Projected crown area or PFT /multi canopy crown area?

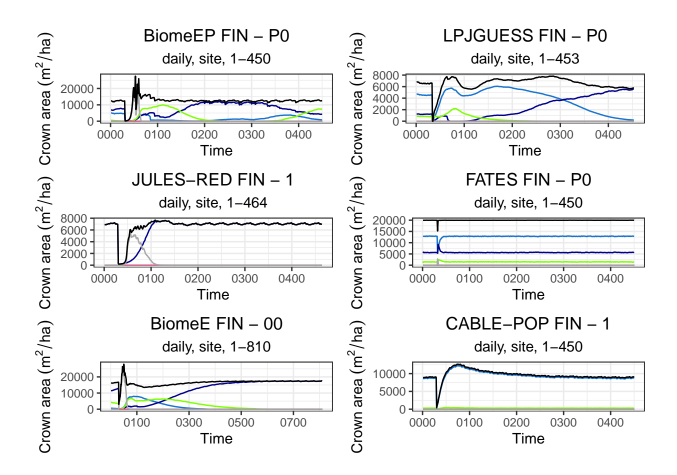
Issue with Crown area variable definition

How to standardise amongst models? What do we want to compare against? I would say satellite observations? Is Crownarea otherwise comparable against any other observations? Maybe some LIDAR observations?

Model	Comments
JULES-RED	projected crown area?
LPJ-GUESS	projected crown area?

Model	Comments
BiomeEP	multi-canopy structure /multi LAI based?
BiomeES	multi-canopy structure /multi LAI based?
FATES	multi-canopy structure /multi LAI based?
CABLE-POP	projected crown area?

```
var = "CA"
site ="FIN"
run ="P0"
BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")</pre>
pBiomeEP <- create_gobj(BiomeEP)</pre>
LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)</pre>
FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name ="FATES")
pFATES <- create_gobj(FATES)</pre>
BiomeES <- get_output_BiomeE_standalone(site, run = "00", var = var, file.dir.biomeE_Standalone, co2_level
## [1] "vobjtovarid4: **** WARNING **** I was asked to get a varid for dimension named PFT BUT this dim
pBiomeES <- create_gobj(BiomeES)</pre>
JULESRED <- get_output_JULESRED(site,run ="1",var = var,file.dir.julesred,co2_levels = "412ppm")
pJULESRED <- create_gobj(JULESRED)</pre>
CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)</pre>
grid.arrange(pBiomeEP,pLPJGUESS,pJULESRED,pFATES,pBiomeES,pCABLEPOP)
```



### BA

Model	Comments
BiomeEP LPJ-GUESS	AHES: Grasses shouldn't have basal area output.
FATES BiomeES JULES-RED CABLE-POP	AHES: BA output missing

```
var = "BA"
site ="FIN"
run ="P0"
BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
pBiomeEP <- create_gobj(BiomeEP)

LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)

#FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name ="FATES")
#plotTemporal(FATES)

BiomeES <- get_output_BiomeE_standalone(site,run = "00",var = var, file.dir.biomeE_Standalone,co2_level</pre>
```

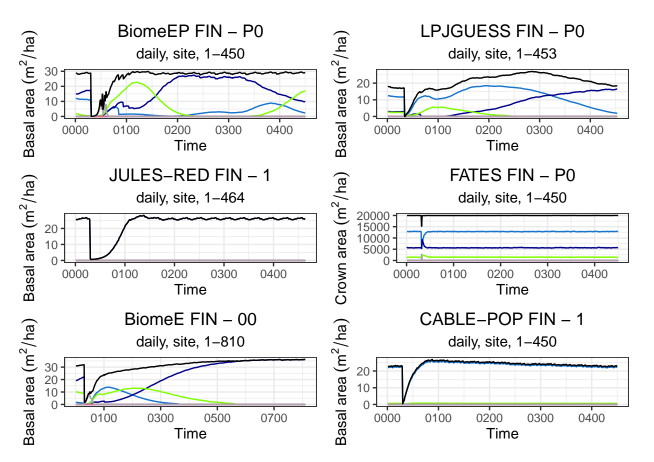
## [1] "vobjtovarid4: \*\*\*\* WARNING \*\*\*\* I was asked to get a varid for dimension named PFT BUT this dim

```
pBiomeES <- create_gobj(BiomeES)

JULESRED <- get_output_JULESRED(site,run ="1",var = var,file.dir.julesred,co2_levels = "412ppm")
pJULESRED <- create_gobj(JULESRED)

CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)

grid.arrange(pBiomeEP,pLPJGUESS,pJULESRED,pFATES,pBiomeES,pCABLEPOP)</pre>
```



## height

[TODO] for height - remove "Total" column, instead plot max, but dotted Height output ambiguous between models. So the y-axis here is misleading and I cannot fully interpret it. Reported in Protocol: "95 th percentile of tree height"

Model	Comments
BiomeEP	AHES: height never goes to 0.
LPJ-GUESS	

```
Model Comments

AHES: no height output. Jessica "We don't have 95th percentile of height in FATES. I can do crown area weighted height if useful?"

BiomeES AHES: v. jittery. Why?AHES: height never goes to 0.

JULES-RED AHES: height never goes to 0.

CABLE-POP AHES: height never goes to 0.
```

```
var = "height"
site = "FIN"
run = "PO"
BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
pBiomeEP <- create_gobj(BiomeEP)
min(BiomeEP@data$Total)</pre>
```

#### ## [1] 20.28285

```
LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")

pLPJGUESS <- create_gobj(LPJGUESS)

# height output confirmed: mean PFT height in 95th percentile.

#FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name = "FATES")

#plotTemporal(FATES)

BiomeES <- get_output_BiomeE_standalone(site,run = "00",var = var, file.dir.biomeE_Standalone,co2_level
```

## [1] "vobjtovarid4: \*\*\*\* WARNING \*\*\*\* I was asked to get a varid for dimension named PFT BUT this dim

```
pBiomeES <- create_gobj(BiomeES)
min(BiomeES@data$Total)</pre>
```

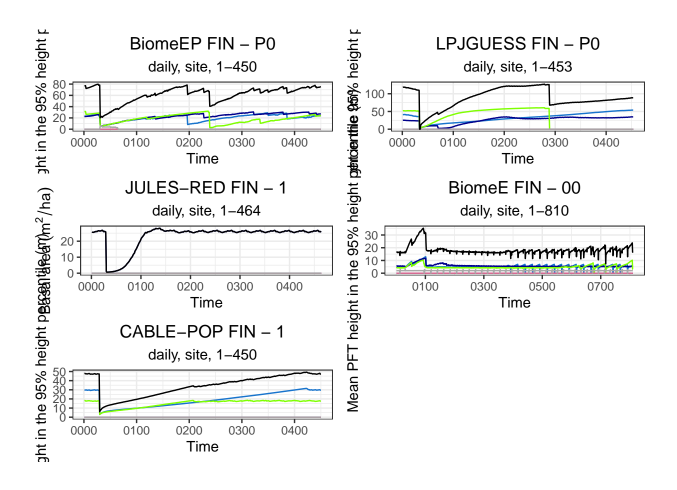
### ## [1] 7.416

```
# height confirmed: "95 th percentile of tree height"

JULESRED <- get_output_JULESRED(site,run ="1",var = var,file.dir.julesred,co2_levels = "412ppm")
pCABLEPOP <- create_gobj(JULESRED)

CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)

grid.arrange(pBiomeEP,pLPJGUESS,pJULESRED,pBiomeES,pCABLEPOP)</pre>
```



## roughness - optional output

The two models which provide this output currently have two orders of magnitude difference in their values.

Model	Comments
BiomeEP	NA
LPJ-GUESS	NA
FATES	AHES: Does show any dynamics in keeping with regrowth dynamics. why?
BiomeES	NA
JULES-RED	NI A
CABLE-POP	NA

```
var = "z0"
site ="FIN"
run ="P0"
#BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
#plotTemporal(BiomeEP)

#LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
#plotTemporal(LPJGUESS)

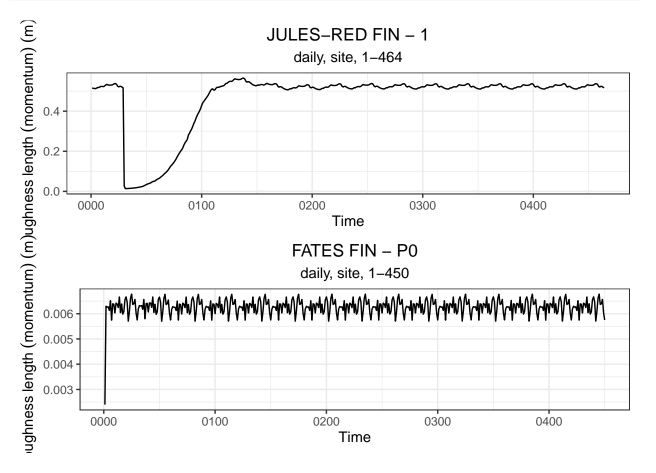
FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name ="FATES")</pre>
```

```
pFATES <- create_gobj(FATES)

#BiomeES <- get_output_BiomeE_standalone(site,run = "00",var = var, file.dir.biomeE_Standalone,co2_leve
#plotTemporal(BiomeES)

JULESRED <- get_output_JULESRED(site,run ="1",var = var,file.dir.julesred,co2_levels = "412ppm")
pJULESRED <- create_gobj(JULESRED)

#CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
#pCABLEPOP <- create_gobj(CABLEPOP)</pre>
grid.arrange(pJULESRED,pFATES)
```

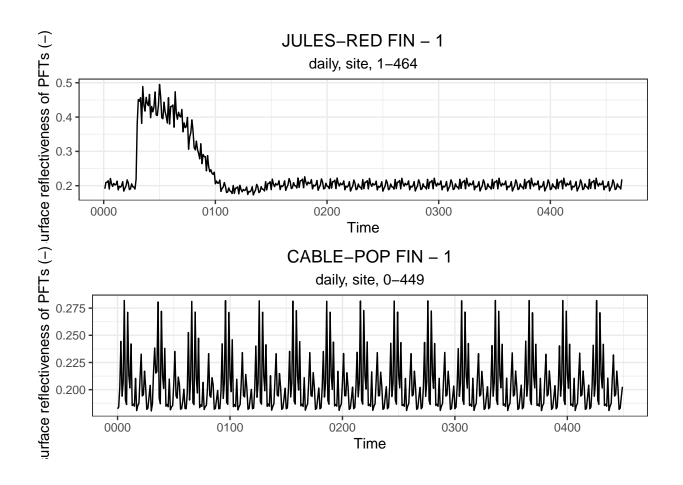


## albedo - optional output

Model	Comments
BiomeEP	NA
LPJ-GUESS	AHES: NA[TODO] get this from post processing still
FATES	AHES: Does show any dynamics in keeping with regrowth dynamics. why?
BiomeES	NA

Model	Comments
JULES-RED	
CABLE-POP	AHES: no trace of regrowth, but in the same
	bulpark as JULES-RED when in equilibrium.
	Juergen Knauer in email: drop this variable. AHES:
	but maybe easy to fix?

```
var = "albedo"
site ="FIN"
run ="P0"
#BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
#plotTemporal(BiomeEP)
#LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
#plotTemporal(LPJGUESS)
#FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name ="FATES")
#plotTemporal(FATES)
#BiomeES <- get_output_BiomeE_standalone(site,run = "00",var = var, file.dir.biomeE_Standalone,co2_leve
#plotTemporal(BiomeES)
JULESRED <- get_output_JULESRED(site,run ="1",var = var,file.dir.julesred,co2_levels = "412ppm")</pre>
pJULESRED <- create_gobj(JULESRED)</pre>
CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
pCABLEPOP <- create gobj(CABLEPOP)</pre>
grid.arrange(pJULESRED,pCABLEPOP)
```



# WBgrowth

Model	Comments
BiomeEP	AHES: remove grasses. double-check the output is
	indeed woody biomass only, not something like eveg
	or another carbon pool.
LPJ-GUESS	AHES: not gross growth?!
FATES	AHES: never goes to $0$ .
BiomeES	AHES: remove grasses, double-check the output is
	indeed woody biomass only, not something like eveg
	or another carbon pool.
JULES-RED	AHES: remove grasses. double-check the output is
	indeed woody biomass only, not something like eveg
	or another carbon pool.
AHES: why no BINE PFT in FIN?	-
CABLE-POP	AHES: WBgrowth rate not much influenced by
	regrowth-dynamics.AHES: add PFT dimension
	easily possible? (not to worry for this analysis?)

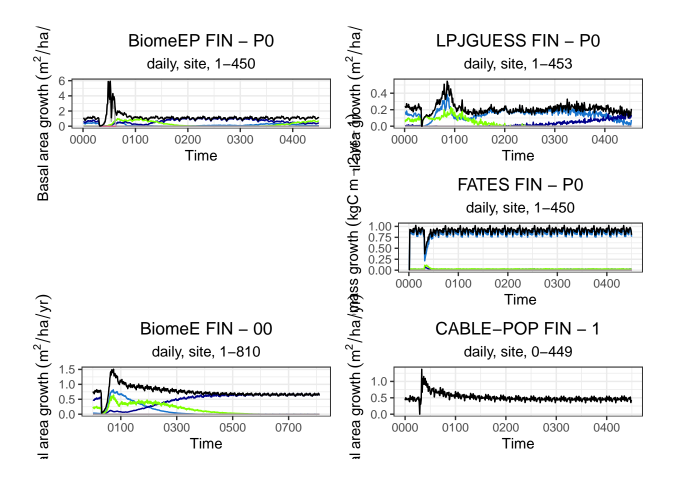
```
var = "WBgrowth"
site ="FIN"
run ="P0"
```

```
BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
pBiomeEP <- create_gobj(BiomeEP)</pre>
LPJGUESS <- get output LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2 levels = "PS 412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)</pre>
FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name = "FATES")
pFATES <- create gobj(FATES)</pre>
BiomeES <- get_output_BiomeE_standalone(site, run = "00", var = var, file.dir.biomeE_Standalone, co2_level
## [1] "vobjtovarid4: **** WARNING **** I was asked to get a varid for dimension named PFT BUT this dim
pBiomeES <- create_gobj(BiomeES)</pre>
JULESRED <- get_output_JULESRED(site,run = "1",var = var,file.dir.julesred,co2_levels = "412ppm")</pre>
pJULESRED <- create_gobj(JULESRED)</pre>
CABLEPOP <- get_output_CABLEPOP(site, run = "1", var = var, file.dir.cablepop, co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)</pre>
grid.arrange(pBiomeEP,pLPJGUESS,pJULESRED,pFATES,pBiomeES,pCABLEPOP)
 mass growth (kgC m +ragss-growth (kgC m +ragss-growth (kgC m
                 BiomeEP FIN - P0
                                                    mass growth (kgC m +ragss-grbwth (kgC m +ragss-grbwth (kgC m
                                                                   LPJGUESS FIN - P0
                    daily, site, 1-450
                                                                       daily, site, 1-453
     0.6
     0.4
                                                                                    0300
                0100
                         0200
                                 0300
                                                           0000
                                                                   0100
                                                                            0200
                                                                                            0400
        0000
                                         0400
                          Time
                                                                              Time
                JULES-RED FIN - 1
                                                                       FATES FIN - PO
                     daily, site, 1-464
                                                                        daily, site, 1-450
     0.75
     0.50
     0.00
         0000
                 0100
                         0200
                                 0300
                                         0400
                                                            0000
                                                                    0100
                                                                             0200
                                                                                     0300
                                                                                             0400
                           Time
                                                                              Time
                  BiomeE FIN - 00
                                                                   CABLE-POP FIN - 1
                    daily, site, 1-810
                                                                        daily, site, 0-449
             0100
                      0300
                               0500
                                        0700
                                                            0000
                                                                    0100
                                                                             0200
                                                                                     0300
                                                                                             0400
                          Time
                                                                              Time
```

# **BAgrowth**

Model	Comments
BiomeEP	AHES: remove grasses. double-check the output is
	BAgrowth, not something like eveg or another
	carbon pool. Units are also v.high.
LPJ-GUESS	AHES: maybe not gross growth?
FATES	NA
BiomeES	AHES: remove grasses. double-check the output is
	indeed woody biomass only, not something like eveg
	or another carbon pool.
JULES-RED	AHES: negative basal area growth.
CABLE-POP	AHES: add PFT dimension easily possible? (not to
	worry for this analysis?)

```
var = "BAgrowth"
site ="FIN"
run ="P0"
BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
pBiomeEP <- create_gobj(BiomeEP)</pre>
LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)</pre>
#FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name = "FATES")
#plotTemporal(FATES)
BiomeES <- get_output_BiomeE_standalone(site, run = "00", var = var, file.dir.biomeE_Standalone, co2_level
## [1] "vobjtovarid4: **** WARNING **** I was asked to get a varid for dimension named PFT BUT this dim
pBiomeES <- create_gobj(BiomeES)</pre>
JULESRED <- get_output_JULESRED(site,run = "1",var = var,file.dir.julesred,co2_levels = "412ppm")</pre>
pJULESRED <- create_gobj(JULESRED,y.lim = c(-6,2))</pre>
CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)</pre>
grid.arrange(pBiomeEP,pLPJGUESS,pJULESRED,pFATES,pBiomeES,pCABLEPOP)
```



## cmort

BiomeEP, BiomeE and Jules-RED seem to reset to bare ground as a mechanism to introduce disturbance. The rest of the models kills all trees. That way, the mortality flux at the disturbance event is recorded in the output.

Model	Comments
BiomeEP	
LPJ-GUESS	
FATES	
BiomeES	
JULES-RED	
CABLE-POP	Added all mortalities together. according to JK,
	have to potentially remove cmort_res. This should
	be done once I have patch-scale data.

```
var = "cmort"
site ="FIN"
run ="PO"

BiomeEP <- get_output_BiomeEP(site,run,var = "cmort_size",file.dir = file.dir.biomeEP,co2_levels = "412]
pBiomeEP <- create_gobj(BiomeEP)</pre>
```

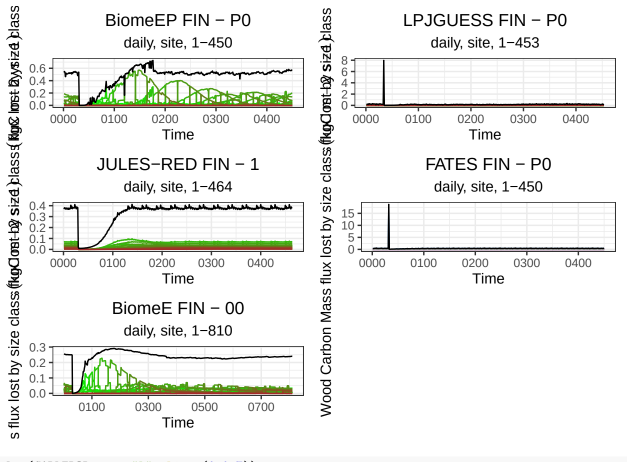
```
LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)

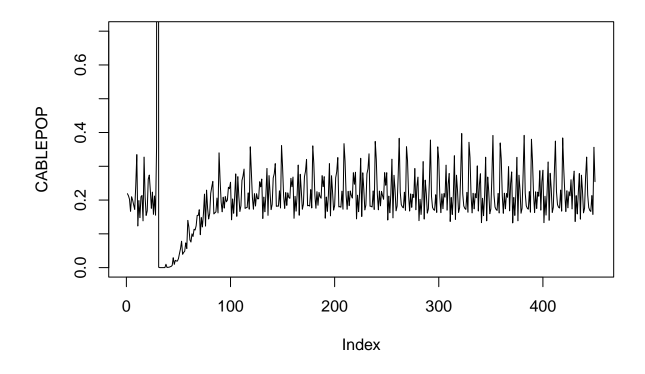
FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name = "FATES")
pFATES <- create_gobj(FATES)

BiomeES <- get_output_BiomeE_standalone(site,run = "00",var = "cmort_size", file.dir.biomeE_Standalone,
pBiomeES <- create_gobj(BiomeES)

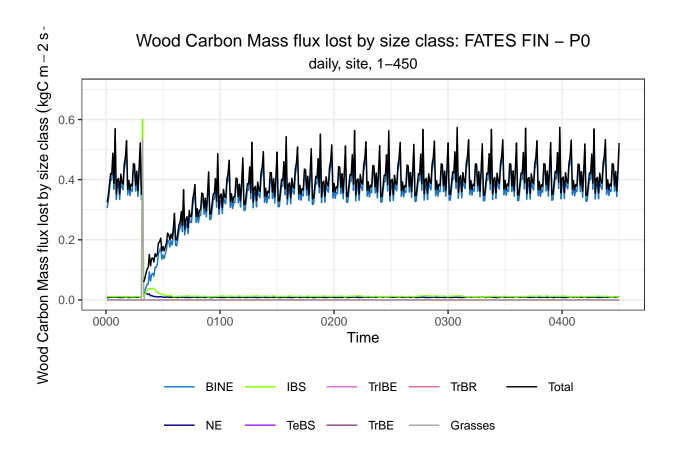
JULESRED <- get_output_JULESRED(site,run = "1",var = var,file.dir.julesred,co2_levels = "412ppm")
pJULESRED <- create_gobj(JULESRED)

#sum up all cmort mechanisms:
CABLEPOP_ccrowd <- get_output_CABLEPOP(site,run = "0",var = "cmort_crowd",file.dir.cablepop,co2_levels = CABLEPOP_cdist <- get_output_CABLEPOP(site,run = "0",var = "cmort_dist",file.dir.cablepop,co2_levels = CABLEPOP_cres <- get_output_CABLEPOP(site,run = "0",var = "cmort_res",file.dir.cablepop,co2_levels = "PCABLEPOP <- get_output_CABLEPOP(site,run = "0",var = "cmort_res",file.dir.cablepop,co2_levels = "PCABLEPOP <- GABLEPOP COMMON COMMON
```

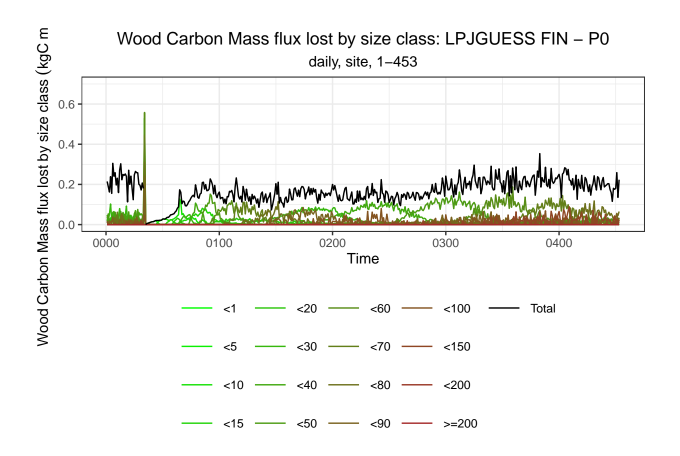




plotTemporal(FATES,y.lim=c(0,0.7))



plotTemporal(LPJGUESS,y.lim=c(0,0.7))



# is the budget closed?

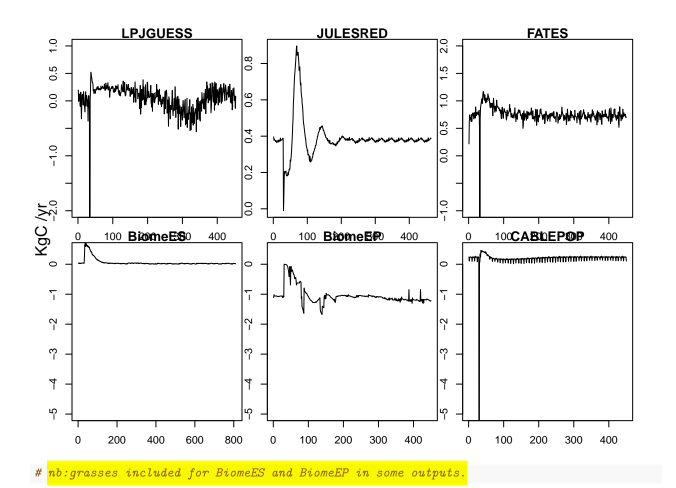
woody growth\_rate - mort\_rate = 0 ( in equilibrium)

Model	Comments
BiomeEP	AHES: too low. One possibility - are the grasses
	included in the cmort output?
LPJ-GUESS	AHES: show age mortality (=decline to
	negative), "Equilibrium" probably maybe more
	dynamic than other models. OK confirmed with
	TOM.
FATES	AHES: too high
BiomeES	
JULES-RED	TOO high in equilibrium
CABLE-POP	AHES: With cmort_res included, too low, when
	cmort res excluded too high.
Or is this "OK" for CABLE-POP? To me there seems to be a bias, that needs to be adressed.	· ·

```
par(mfrow =c(2,3),oma=c(1,2,1,1),mar=c(1,1,1,1))

var = "cmort"
site = "BCI"
```

```
run ="P0"
BiomeEP cmort <- get output BiomeEP(site,run,var = "cmort size",file.dir = file.dir.biomeEP,co2 levels
LPJGUESS_cmort <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm
FATES_cmort <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name = "FATES")
BiomeES_cmort <- get_output_BiomeE_standalone(site, run = "00", var = "cmort_size", file.dir.biomeE_Stand
JULESRED_cmort <- get_output_JULESRED(site, run = "1", var = var, file.dir.julesred, co2_levels = "412ppm")
#sum up all cmort mechanisms:
c("cmort_crowd", "cmort_dist", "cmort_res")
## [1] "cmort_crowd" "cmort_dist" "cmort_res"
CABLEPOP_ccrowd <- get_output_CABLEPOP(site,run = "0",var = "cmort_crowd",file.dir.cablepop,co2_levels
CABLEPOP_cdist <- get_output_CABLEPOP(site,run = "0",var = "cmort_dist",file.dir.cablepop,co2_levels =
CABLEPOP_cres <- get_output_CABLEPOP(site,run = "0",var = "cmort_res",file.dir.cablepop,co2_levels = "Particle of the content 
CABLEPOP <-CABLEPOP_ccrowd@data$Total +CABLEPOP_cdist@data$Total +CABLEPOP_cres@data$Total
CABLEPOP <-CABLEPOP ccrowd@data$Total +CABLEPOP cdist@data$Total
#########################
##wbgrowth
var = "WBgrowth"
BiomeEP_WBgrowth <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412"
#plotTemporal(BiomeEP_WBgrowth)
LPJGUESS_WBgrowth <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412"
FATES_WBgrowth <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name = "FATES")
BiomeES_WBgrowth <- get_output_BiomeE_standalone(site, run = "00", var = var, file.dir.biomeE_Standalone,
## [1] "vobjtovarid4: **** WARNING **** I was asked to get a varid for dimension named PFT BUT this dim
JULESRED_WBgrowth <- get_output_JULESRED(site,run = "1",var = var,file.dir.julesred,co2_levels = "412pp:
CABLEPOP_WBgrowth <- get_output_CABLEPOP(site,run = "0",var = var,file.dir.cablepop,co2_levels = "PS_41"
#####################################
##AGcwood
var ="AGcwood"
BiomeEP_AGcwood <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412p"
LPJGUESS_AGcwood <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412p
#plotTemporal(LPJGUESS_AGcwood)
FATES_AGcwood <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name = "FATES")
```



## stemmort

Model	Comments
BiomeEP	AHES: Unit low: should be numer of stems /year. *nstems[y-1]?
LPJ-GUESS	
BiomeES	AHES: Unit low: should be numer of stems /year.
	*nstems[y-1] ?Why so jittery? Biology model
	structure or post-processing?
JULES-RED	AHES: what causes the small oscillations on top of
	the general number flux dynamics?
CABLE-POP	NA

FATES- AHES: output year 1: stem number flux almost 0 AHES: strange oscillations in  $\_$ overstory.nc, due to PFT2 (e.g. simyear 32)

Year	Lat	Lon	<1	<5	<10	<15	<20
32	62.25	23.25	0	1678.8756	572.7487876	164.84464856	88.34444358

these numbers are also present in the .netcdf file, so not a read-in error. or is this just a quick transition of trees from under- to over-story after the disturbance? But I what is this oscillation exactly?

```
var = "stemmort"
site = "FIN"
run = "PO"

BiomeEP <- get_output_BiomeEP(site,run,var = "stemmort_size",file.dir = file.dir.biomeEP,co2_levels = "pBiomeEP <- create_gobj(BiomeEP)

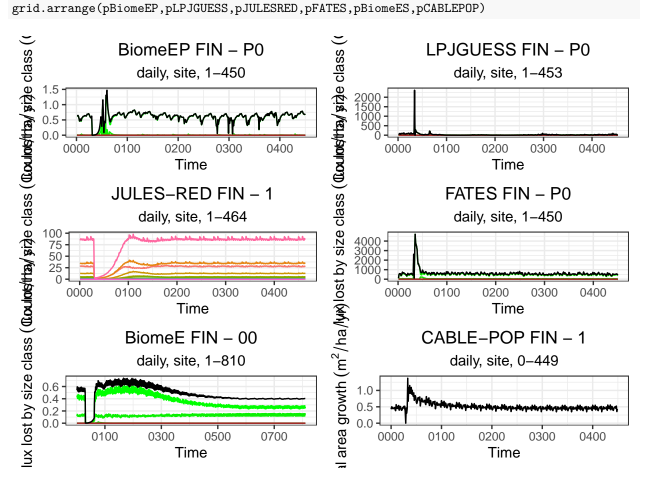
LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)

FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name = "FATES")
pFATES <- create_gobj(FATES)

BiomeES <- get_output_BiomeE_standalone(site,run = "00",var = "stemmort_size", file.dir.biomeE_Standalone)
BiomeES <- create_gobj(BiomeES)

JULESRED <- get_output_JULESRED(site,run = "1",var = var,file.dir.julesred,co2_levels = "412ppm")
pJULESRED <- create_gobj(JULESRED)

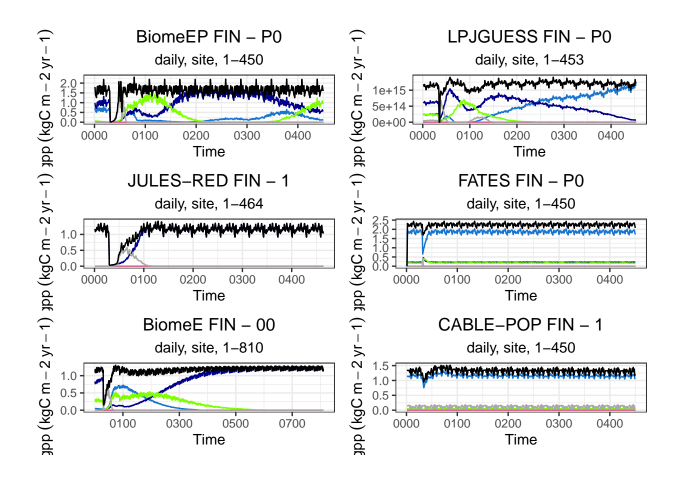
#CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
#plotTemporal(CABLEPOP)</pre>
```



### gpp

Model	Comments
BiomeEP	AHES: netcdf only 0 values for gpp for BCI -double check output
LPJ-GUESS	one of a dapare
FATES	
BiomeES	
JULES-RED	AHES:FIN: only PFT2 really active. PFT1 never present?
CABLE-POP	•

```
var = "gpp"
site ="FIN"
run ="P0"
BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
pBiomeEP <- create_gobj(BiomeEP)</pre>
LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)</pre>
FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name = "FATES")</pre>
pFATES <- create_gobj(FATES)</pre>
BiomeES <- get_output_BiomeE_standalone(site, run = "00", var = var, file.dir.biomeE_Standalone, co2_level
## [1] "vobjtovarid4: **** WARNING **** I was asked to get a varid for dimension named PFT BUT this dim
pBiomeES <- create_gobj(BiomeES)</pre>
JULESRED <- get_output_JULESRED(site,rum = "1",var = var,file.dir.julesred,co2_levels = "412ppm")</pre>
pJULESRED <- create_gobj(JULESRED)</pre>
CABLEPOP <- get_output_CABLEPOP(site, run = "1", var = var, file.dir.cablepop, co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)</pre>
grid.arrange(pBiomeEP,pLPJGUESS,pJULESRED,pFATES,pBiomeES,pCABLEPOP)
```



### npp

Model	Comments
BiomeEP LPJ-GUESS	
FATES BiomeES	AHES: simulation output year 1 negative
· · · · · ·	AHES:FIN: only PFT2 really active. PFT1 never present?

#### **CABLE-POP**

```
var = "npp"
site ="FIN"
run ="P0"

BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
pBiomeEP <- create_gobj(BiomeEP)

LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)

FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name = "FATES")</pre>
```

```
pFATES <- create_gobj(FATES)</pre>
BiomeES <- get_output_BiomeE_standalone(site, run = "00", var = var, file.dir.biomeE_Standalone, co2_level
## [1] "vobjtovarid4: **** WARNING **** I was asked to get a varid for dimension named PFT BUT this dim
pBiomeES <- create_gobj(BiomeES)</pre>
JULESRED <- get_output_JULESRED(site,run = "1",var = var,file.dir.julesred,co2_levels = "412ppm")
pJULESRED <- create_gobj(JULESRED)</pre>
CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)</pre>
grid.arrange(pBiomeEP,pLPJGUESS,pJULESRED,pFATES,pBiomeES,pCABLEPOP)
                 BiomeEP FIN - P0
                                                                   LPJGUESS FIN - P0
 (\log (\log m - 2 \text{ yr} - 1) \log (\log m - 2 \text{ yr} - 1) \log (\log m - 2 \text{ yr} - 1)
                    daily, site, 1-450
                                                                        daily, site, 1-453
                                                        0.6
     1.0
    0.5
     0.0
                         0200
                0100
                                 0300
                                          0400
                                                                    0100
                                                                            0200
                                                                                     0300
                                                                                             0400
        0000
                                                           0000
                          Time
                                                                              Time
                                                                      FATES FIN - P0
                JULES-RED FIN - 1
                                                                        daily, site, 1-450
                    daily, site, 1-464
    0.6
                                                        1.0
                                                       0.5
                                                        0.0
        0000
                0100
                        0200
                                0300
                                         0400
                                                           0000
                                                                    0100
                                                                            0200
                                                                                     0300
                                                                                             0400
                          Time
                                                                              Time
                  BiomeE FIN - 00
                                                                   CABLE-POP FIN - 1
                    daily, site, 1-810
                                                                        daily, site, 1-450
    0.6
```

0000

0100

0200

Time

0300

0400

### nbp

0100

0300

From the protocol: "Positive flux is into the land." post-disturbance, there is a less positive flux into the land for FATES

0500

Time

0700

Model	Comments
BiomeEP	
LPJ-GUESS	AHES: dynamics post disturbance look a bit odd double-check output; does soil C increase?
FATES	AHES: Seems to me in terms of NBP, FATES is not yet(?) in equilibrium. Seems to always take up more than it releases. different to other models or overaccounting of output? First output value needs double-checking.
BiomeES JULES-RED CABLE-POP	NA

```
var = "nbp"
site ="FIN"
run ="P0"
BiomeEP <- get_output_BiomeEP(site,run,var = var,file.dir = file.dir.biomeEP,co2_levels = "412ppm")
pBiomeEP <- create_gobj(BiomeEP)</pre>
LPJGUESS <- get_output_LPJGUESS(site,run,var,file.dir = file.dir.lpjguess,co2_levels = "PS_412ppm/")
pLPJGUESS <- create_gobj(LPJGUESS)</pre>
FATES <- get_output_FATES(site,run,var,file.dir=file.dir.fates,model_name = "FATES")
pFATES <- create_gobj(FATES)</pre>
BiomeES <- get_output_BiomeE_standalone(site, run = "00", var = var, file.dir.biomeE_Standalone, co2_level
pBiomeES <- create_gobj(BiomeES)</pre>
#JULESRED <- get_output_JULESRED(site,run = "1",var = var,file.dir.julesred,co2_levels = "412ppm")
#plotTemporal(JULESRED)
CABLEPOP <- get_output_CABLEPOP(site,run = "1",var = var,file.dir.cablepop,co2_levels = "PS_412ppm")
pCABLEPOP <- create_gobj(CABLEPOP)</pre>
grid.arrange(pBiomeEP,pLPJGUESS,pJULESRED,pFATES,pBiomeES,pCABLEPOP)
```

