# LPJmL Runner

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LPJmL Runner is a lpjmlkit module of functions that have the goal to simplify the execution of simulations with LPJmL and further to execute complex, nested and multiple simulation sequences fast and less error prone without having a big (bash) script overhead.

# Setup

Please make sure to have set the working environment for LPJmL correctly if you are not working on the PIK cluster (with Slurm Workload Manager). On the PIK cluster please load the lpjml module (below) or add it to your ".profile".

```
# load lpjml module to use LPJmL internal functions
module load lpjml
```

# Overview

The LPJmL Runner generally requires 3 to 4 working steps: Define a modified parameter table (1), create the corresponding configuration files (2), check if the these are valid for LPJmL (3 - optional) and run or submit LPJmL with each configurations (4).

### 1. Define a table of modified configuration parameters

Define what LPJmL parameters/settings (here all referred to as parameters) to be changed. They can be changed directly in the corresponding "js" file or in a tibble table (see example). For a single simulation this is a matter of personal routine, but when it comes to multiple runs where these parameters differ from each other, they have to be specified in a tibble.

?write\_config for more information.

```
my_params <- tibble(
    sim_name = c("scenario1", "scenario2"),
    random_seed = c(42, 404),
    param.k_temp = c(NA, 0.03),
    new_phenology = c(TRUE, FALSE)
)</pre>
```

# 2. Create corresponding Configuration files

Now the central function is write\_config, create and write LPJmL Configuration (Config) file(s) "config\_\*.json" from a table (tibble) with the parameters of a base "lpjml.js" file to be changed. ?write\_config for more information.

```
config_details <- write_config(my_params, model_path, output_path)</pre>
```

## 3. Check validity of Configurations

Check whether your Config(s) are valid for LPJmL by passing the returned tibble to check\_lpjml. It won't raise an error (dependencies might not be satisfied yet) but will print/return the information of lpjcheck.

```
lpjml_check(config_details, model_path, output_path)
```

#### 4. Run or submit LPJmL

Execute LPJmL for each Configuration via submit\_lpjml or run it interactively via run\_lpjml. run\_lpjml can also be utilized within slurm jobs to execute multiple single cell runs. ?submit\_lpjml or ?run\_lpjml for more information.

```
# submit to Slurm
run_details <- submit_lpjml(config_details, model_path, output_path)
# OR run interactively
run_details <- run_lpjml(config_details, model_path, output_path)</pre>
```

### miscellaneous

More helpful functions that come with LPJmL Runner are:

- read\_config to read a "config\_\*.json" file as a nested R list object
- use the R internal View function for a tree visualization of a "config\_\*.json" file
- make\_lpjml function for compiling LPJmL.

# Usage

```
library(lpjmlkit)
# why tibble? -> https://r4ds.had.co.nz/tibbles.html
library(tibble)

model_path <- "./LPJmL_internal"
output_path <- "./my_runs"</pre>
```

# Single cell simulations

Single cell (or short number of multiple cells) simulations can be executed locally or on a login node. This mode is especially useful when it comes to testing or comparing local data.

# Example Potential natural vegetation and land-use run

```
# create parameter tibble
params <- tibble(</pre>
  sim_name = c("spinup", "lu", "pnv"),
  landuse = c("no", "yes", "no"),
  # only for demonstration
  nspinup = c(1000, NA, NA),
  reservoir = c(FALSE, TRUE, FALSE),
  startgrid = c(27410, 27410, 27410),
  river_routing = c(FALSE, FALSE, FALSE),
  wateruse = c("no", "yes", "no"),
  const_deposition = c(FALSE, FALSE, TRUE),
  # run parameter: order defines the execution order
     (spinup is always 1 -> no -DFROM_RESTART),
     could be followed by a historic (order=2) and a future run (order=3)
     can be extended as far as required (order=n)
  order = c(1, 2, 2),
  # run parameter: dependency sets the restart paths to the corresponding
  # restart_filename
  dependency = c(
    NA, "spinup", "spinup"
# write confiq files
config_details <- write_config(</pre>
 params = params, # pass the defined parameter tibble
 model_path = model_path,
 output_path = output_path,
  js_filename = "lpjml.js" # (default) the base js file
# read and view config
config_lu <- read_config(</pre>
  filename = pasteO(output_path,"/configurations/config_lu.json")
View(config_lu)
# check confiq & LPJmL
check_config(
 x = config_details, # can be filename (vector) or tibble
 model_path = model_path,
  output_path = output_path
# execute runs sequentially
run details <- run lpjml(</pre>
  config_details,
```

```
model_path = model_path,
output_path = output_path)
```

# Example Old vs. new phenology and old land-use vs. input toolbox

```
# create parameter tibble
params <- tibble(</pre>
  sim_name = c("spinup_oldphen",
               "spinup_newphen",
               "oldphen",
               "old_lu",
               "lu toolbox"),
  # object oriented like syntax to access nested json elements
  input.landuse.name = c(
   NA,
   NA,
   NA,
   NA.
    "input_toolbox_30arcmin/cftfrac_1500-2017_64bands_f2o.clm"
  ),
  nspinup = c(1000, 1000, NA, NA, NA),
 new_phenology = c(FALSE, TRUE, FALSE, TRUE, TRUE),
 startgrid = c(27410, 27410, 27410, 27410, 27410),
 river_routing = c(FALSE, FALSE, FALSE, FALSE),
 order = c(1, 1, 2, 2, 2),
 dependency = c(NA, NA, "spinup_oldphen", "spinup_newphen", "spinup_newphen")
# write config files
config_details <- write_config(params, model_path, output_path)</pre>
# check config & LPJmL
check_config(config_details, model_path, output_path)
# execute runs sequentially
run_details <- run_lpjml(config_details, model_path, output_path)</pre>
```

#### Global simulations on the PIK cluster

Global simulations are simulations on all available cells with a coherent water cycle. It requires more computational ressources which is why they have to be run at dedicated compute nodes, at PIK Cluster only accessible via SLURM Job scheduler. Therefore LPJmL has to be "submitted".

# Example Compare old vs new land use (lpjml input toolbox)

```
# create parameter tibble
params <- tibble(</pre>
  sim_name = c("spinup",
               "old lu",
               "lu_toolbox"),
  input.landuse.name = c(
    NA,
    NA,
    "input_toolbox_30arcmin/cftfrac_1500-2017_64bands_f2o.clm"
  ),
  order = c(1, 2, 2),
  dependency = c(NA, "spinup", "spinup"),
  # slurm option wtime: analogous to sbatch -wtime defines slurm option
      individually per config, overwrites submit_lpjml argument
      (same for sclass, ntasks, blocking)
  wtime = c("15:00:00", "3:00:00", "3:00:00")
# write config files
config_details <- write_config(</pre>
 params = params,
 model_path = model_path,
 output path = output_path,
  output_list = c("vegc", "soilc", "cftfrac", "pft_harvestc", "irrig"),
  output_list_timestep = c("annual", "annual", "annual", "annual", "monthly"),
  # output_list_timestep = "annual",
 output_format = "clm"
# check confiq & LPJmL
check_config(config_details, model_path, output_path)
# submit runs to slurm
run_details <- submit_lpjml(</pre>
 x = config_details,
 model_path = model_path,
 output_path = output_path,
 group = "open")
```

# Notes & tips

1. You can save the generated config tibble by applying saveRDS to it to reuse for a rerun or resubmission next time . . .

2. Also if you want do not want to submit all runs you can ...

```
# use a subset for the rows - in this example you may only want to resubmit the
# transient runs
run_details <- submit_lpjml(
    x = config_details[2:3, ],
    model_path = model_path,
    output_path = output_path,
    group = "open")</pre>
```

3. a bit dirty though If you want to reuse an old spinup simulation, you can copy the file or create a symlink of the file to "<output\_path>/restart/<spinup\_sim\_name>/restart.lpj". Make sure the file/symlink is named "restart.lpj"