(PART) Demos and Vignettes

BioCro Run Using the Virtual Machine

- 1. Download the VirtualBox application from here and open the application.
- 2. Download the PEcAn **virtual machine** .ova file from here and open it from within VirtualBox by going to "File" -> "Import appliance" and selecting the .ova. Start the virtual machine by clicking on the green arrow in VirtualBox. This will bring up a new window, and the virtual machine is ready when it displays pecan login:
- 3. Open up the **RStudio** interface to the VM by going to localhost:6480/rstudio/ in your browser.
- 4. Use ssh to open up the **command line** for the VM. Do this by opening up a command line program on your computer and execute ssh -p 6422 carya@localhost. You will be prompted for a password with carya@localhost's password:. The password is illinois, and the letters will not appear when you type them in.
- 5. Create a new folder for the run results using mkdir biocro_results. Edit the XML to include this file path by doing nano pecan/tests/pecan64.biocro.xml and changing the third line from <outdir>pecan/outdir> to <outdir>biocro_results/outdir> and saving these changes. Do the model run by putting the BioCro XML file through PEcAn by executing pecan/web/workflow.R --settings pecan/tests/pecan64.biocro.xml.
- 6. Go to the RStudio instance that is open in your browser. In the file directory, there should be a new biocro_results folder, and you can see all the files by clicking on it. The .nc data files that are in the out subfolder can be plot in an R script, as is shown for leaf area index.

PEcAn.visualization::plot.netcdf("biocro_results/out/SA-salix-chi_leaf-0.159/2004.nc", "LAI")

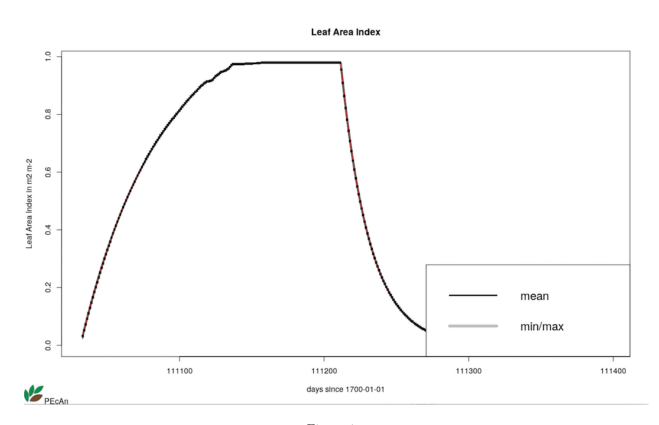


Figure 1: