Tutorial #2

LMDZ team

December 6th, 2016

Contents

\mathbf{E}_{2}	kperime	
2.1	l Tracei	rs
	2.1.1	Radon and lead
	2.1.2	Inserting new tracers
2.2	2 Switch	hing physics parameterizations
2.3	3 Sensit	ivity to the soil scheme
	2.3.1	ORCHIDEE 2-layers
	2.3.2	ORCHIDEE 11-layers
	2.3.3	Bucket
	2.3.4	Bucket scheme with imposed soil water content

This tutorial makes use of a zoomed configuration of LMDZ. Section 1 is an initialization phase. In section 2, you can either take a closer look at tracers, or the Orchidee model or switch some parameterizations of the physics part of the program. The three experiments in section 2 are independent. In section 3, you will nudge the model toward analyzed wind fields.

Start by installing the model with the script install_lmdz.sh, if you have not already done so in tutorial #1 by using:

```
wget http://www.lmd.jussieu.fr/~lmdz/Distrib/install_lmdz.sh
chmod +x install_lmdz.sh
./install_lmdz.sh -d 48x36x39
```

1 Setting up a case with a zoomed grid

• Go to the directory LMDZtesting/modipsl/modeles/LMDZ5, which contains files makelmdz_fcm, libf...In this directory, download the following tar file:

```
wget http://www.lmd.jussieu.fr/~lmdz/Distrib/tuto.tar
tar -xf tuto.tar
cd TUTO
```

- In the directory TUTO, take a look at the extracted files and at the DEF directory. In this directory, you should edit the file called gcm.def if you want to place the center of the zoom at your preferred location. For that you just have to change the longitude and latitude of the zoom center, clon and clat.
- If you are interested in coupling LMDZ with the surface scheme Orchidee, edit the init.sh file and set option veget=1. If not (veget=0), the model will be run with a simplified scheme for surface hydrology: the "bucket" scheme.
- Increase the stack memory you can use by typing the following command:

```
ulimit -Ss unlimited
```

or, even better, add this command in you shell start-up file (if you use Bash, your start-up file can be .profile or .bash_profile).

• Run the script :

./init.sh

The script first compiles the model (gcm.e) again and also compiles the program ce01.e, which creates initial state and boundary conditions. init.sh then downloads new NetCDF files which contain the surface orography, sea-surface temperature, and so on, as well as 3D meteorological files taken from ECMWF analyses at a particular date. init.sh then runs ce01.e which creates files start.nc, startphy.nc and limit.nc in a directory called INITIAL. These files are then used to initialize a new simulation in a directory called SIMU1. You can check that these 3 files have actually been created in the directory SIMU1. If not, please ask for our help. If you set veget=1, you also need to run a first simulation to create the ORCHIDEE initial file. To do so:

cd SIMU0
./gcm.e > listing0

This simulation should create a file called sechiba_rest_out.nc. The ORCHIDEE initial file of your simulation, sechiba_rest_in.nc, should point to this file (this is done automatically for the SIMU1 experiment when veget=1).

• If you have the files start.nc, startphy.nc and limit.nc (and optionally sechiba_rest_in.nc for veget=1) in your SIMU1 directory, you are almost done. In the file config.def, add the line:

flag_pres__00003 =

Thus, the NetCDF variable pres, which contains the pressure at model layers, will be included in the file #3, histhf.nc.

• You can now run the model by executing the command ./gcm.e. The simulation should end with this message "Everything is cool" and the output files histday.nc, histmth.nc and histhf.nc should be present.

2 Experiments

The three subsections 2.1, 2.2 and 2.3 are independent.

2.1 Tracers

2.1.1 Radon and lead

In the file traceur.def, you can see that two trace species, named RN, for radon, and PB, for lead, are already defined in addition to vapor and condensed water. Visualize these two tracers in the file histhf.nc. The NetCDF variables RN and PB are numbers of atoms per unit mass, in kg⁻¹. Usually, activities are preferred for comparison with observations. The activities per unit mass, in Bq kg⁻¹, are:

$$A_{\rm Rn} = \frac{\rm RN}{4,765 \cdot 10^5}$$

$$A_{\rm Pb} = \frac{\rm PB}{1,028 \cdot 10^9}$$

(The numerical values that appear in these fractions are the lifetimes of radon and lead, in s.) You can take a look at LMDZ info number 7, figure 19, or the plot in the General introduction of the LMDZ model (§ 4 "Operating modes") to check that you have sensible results. A recent work on the modeling of tracers with LMDZ is Pilon et al. (2015, QJRMS). If you want to see activities per unit volume, you should divide by the mass density, using NetCDF variables pres, temp and ovap in the file histhf.nc. (The Ferret color palette in figure 19 of LMDZ info number 7 is saz2.)

2

2.1.2 Inserting new tracers

We will now explain how to modify the code in order to add idealized tracers. As an example, we will add two tracers, and call them NewTr1 and NewTr2. We will define a domain of the horizontal grid in which the two tracers will initially have the same constant value. The first tracer will be transported by the boundary layer and convective sub-grid-scale motion, in addition to large-scale advection. The second tracer will only be transported by large-scale advection.

We will have to modify the Fortran program but let us first consider the run-time parameters that we have to modify. At run-time, we decide to include the tracers NewTr1 and NewTr2 in the simulation by changing the file traceur.def. Change the number of tracers at the first line of traceur.def and append one line for each tracer:

```
7
14 14 H2Ov
10 10 H2Ol
10 10 H2Oi
10 10 RN
10 10 PB
10 10 NewTr1
10 10 NewTr2
```

Now let us turn to the Fortran program. The only file we need to modify is

LMDZtesting/modipsl/modeles/LMDZ5/libf/phylmd/traclmdz_mod.F90

Here are the changes you should make in that file:

- Declare two new module variables, id_NewTr1 and id_NewTr2, with type integer. These are the identifying numbers of the tracers in the program. You can take a previous declaration (id_pcsat, id_pcocsat...) as a template. (Do not forget the OpenMP directives.)
- All remaining changes will be made in the procedure traclmdz_init, which is inside the module traclmdz_mod. In order to define id_NewTr1 and id_NewTr2, the program will scan the file traceur.def, looking for NewTr1 and NewTr2. So you should initialize id_NewTr1 and id_NewTr2 to 0 before the loop beginning at line 175. There is a comment just above, saying "Recherche des traceurs connus", which means "looking for known tracers".
- In the body of this loop, set id_NewTr1 to the value of index it if tname(iiq) equals NewTr1. You can add the test near line 265, for instance, after the test for pcq0. You can take another tracer as a template. Do the same for id NewTr2.
- For NewTr2, just after setting id_NewTr2, deactivate convective and boundary layer transport by setting conv_flg(it) and pbl_flg(it) to 0.
- Finally, we will initialize the tracers. There is a loop on tracers, beginning at line 294, which tests whether the initial tracer field, read from the file start.nc, is zero everywhere. At this point, the tracer field would also be zero if it was not found in start.nc. There is a comment just above the test which says "Initalize tracer that was not found in restart file" (the typo "Initalize" is in the code!). In the body of the test, for our two tracers, change the value at the surface in some horizontal domain. The value of tracers is in the variable tr_seri. The first dimension of tr_seri is for the horizontal position, the second dimension is for the vertical level and the third dimension identifies the tracer. The index of the surface in the vertical dimension is 1. Use variables xlat (latitudes) and xlon (longitudes) to choose the horizontal domain.

In summary, after making those changes, svn diff should give you something like this:

```
+ INTEGER, SAVE:: id_newtr1, id_newtr2
 !$OMP THREADPRIVATE(id newtr1, id newtr2)
 CONTAINS
@@ -172,6 +174,8 @@
     id_rn=0; id_pb=0; id_aga=0; id_be=0; id_o3=0
     id_pcsat=0; id_pcocsat=0; id_pcq=0; id_pcs0=0; id_pcos0=0; id_pcq0=0
     id newtr1 = 0
     id_newtr2 = 0
     DO it=1,nbtr
        iiq=niadv(it+2)
        IF ( tname(iiq) == "RN" ) THEN
@@ -262,6 +266,12 @@
        ELSE IF ( tname(iiq) == "pcq0" .OR. tname(iiq) == "Pcq0" ) THEN
           id_pcq0=it
           conv_flg(it)=0 ! No transport by convection for this tracer
        else if (tname(iiq) == "NewTr1") then
           id newtr1 = it
        else if (tname(iiq) == "NewTr2") then
           id_newtr2 = it
           conv_flg(it) = 0
           pbl_flg(it) = 0
        ELSE
           WRITE(lunout,*) 'This is an unknown tracer in LMDZ : ', trim(tname(iiq))
        END IF
@@ -325,6 +335,9 @@
                    tr_seri(i,:,it) = 100.
                 END IF
              END DO
           else if (it == id_newtr1 .or. it == id_newtr2) then
              where (xlat >= 40. .and. xlat <= 45. .and. xlon >= 0. &
                   .and. xlon \le 5.) tr_seri(:, 1, it) = 1.
           ELSE
              ! No specific initialization exist for this tracer
              tr_seri(:,:,it) = 0.
   Re-compile the program:
cd some_path/LMDZtesting/modipsl/modeles/LMDZ5/TUTO/SIMU1
rm gcm.e
cd ../..
If you ran init.sh with veget=0:
./makelmdz -d 48x36x39 gcm
If you set veget=1 in init.sh:
./makelmdz -d 48x36x39 -v orchidee2.0 -cpp ORCHIDEE_NOZOH gcm
Rename restart files:
cd TUTO/SIMU1
mv restart.nc start.nc
mv restartphy.nc startphy.nc
mv sechiba_rest_out.nc sechiba_rest_in.nc
Run the model:
../../gcm.e > listing
(could take about 15 mn). Visualize the two new tracers in histhf.nc and the difference between them.
```

2.2 Switching physics parameterizations

You can change the parameters of the model that are set up in the .def files and run new simulations without having the compile the model again. This is very convenient to test different parameterizations.

- Prepare for five more days of simulation: rename files restart.nc and restartphy.nc (created at the end of the first simulation) to start.nc and startphy.nc.
- Create a directory where you will run the model with modified .def files. In this new directory, you can create symbolic links to the files start.nc, startphy.nc and limit.nc which are in TUTO/SIMU1, and you need to copy the *.def files from TUTO/SIMU1, except those beginning with used_. If you used veget=1, don't forget to create a link called sechiba_rest_in.nc pointing to the ../SIMU0/sechiba_rest_out.nc file. Here are the commands you will need:

```
mkdir SIMU1_test1
cd SIMU1_test1
ln -s ../SIMU1/start.nc .
ln -s ../SIMU1/startphy.nc .
ln -s ../SIMU1/limit.nc .
ln -s ../SIMU0/sechiba_rest_out.nc sechiba_rest_in.nc
cp ../SIMU1/*.def .
rm -f used_*.def
```

In case you re-run the simulation, make sure to remove the sechiba_rest_out.nc file in the directory, or the model will stop with an error (STOP 1).

• In your new directory, in file physiq.def, change the boundary layer parameterization by deactivating the Mellor and Yamada scheme for the boundary layer, based on a prognostic equation for the Turbulent Kinetic Energy:

```
iflag_pbl = 1
```

instead of the nominal value 11 (which corresponds to the variant for very stable planetary boundary layer and exact dissipation). Thus, you will switch to the "Standard Physics" (SP, LMDZ5A).

• Run the model in the two directories and compare the results. You can use a convenient command called ncdiff to directly compute the difference between the two NetCDF files in a new file:

```
ncdiff ../SIMU1/histday.nc histday.nc histday_diff.nc
```

• Another test you can try is to deactivate the thermal plume model:

```
iflag_thermals=0
```

instead of the nominal value 18.

Remember that you can modify the content or time frequency of the output files if you want to focus on a particular question.

2.3 Sensitivity to the soil scheme

2.3.1 Running with ORCHIDEE 2-layers

Execute init.sh with veget=1. As explained in section 1, the model is run automatically for one day and creates a start file named sechiba_rest_in.nc (sechiba is the hydrological part of the soil/vegetation model Orchidee).

Then you can run gcm.e to do a simulation with Orchidee activated. The number of days, set in run.def, is nday=5. To get results faster, decrease nday to 3, and change 5day in 3day in config.def, in the line 'phys_out_filetimesteps' (otherwise your output file histmth.nc will be empty).

You can play with the sechiba output frequency by changing in orchidee.def the variable WRITE_STEP (in seconds; default: 86400 for daily output); 0 means no sechiba output; -1 means monthly output; N*86400 means output written every N days). A second output file sechiba_out_2.nc is for high-frequency output, modulated by WRITE_STEP2 (default: 10800, for 3 hours)

You can change the complexity level of outputs by playing with the SECHIBA_HISTLEVEL variable: higher SECHIBA_HISTLEVEL means more variables in output. The variables corresponding to the various output levels are coded in

modipsl/modeles/ORCHIDEE/src_sechiba/intersurf.f90

2.3.2 Running with ORCHIDEE 11-layers

Set HYDROL_CWRR to y in orchidee.def (the default is n, meaning that ORCHIDEE 2-layers is used).

If it's the first time that you do a run with ORCHIDEE in your current working directory (which is the case if you don't have a sechiba_rest_in.nc file), the procedure is the same as described in § 2.3.1 for ORCHIDEE 2-layers.

If you have already done a run with ORCHIDEE -2 layers, you need to re-create an initial state file adapted to the multi-layer hydrology:

- 1. remove the file sechiba_rest_in.nc in your working directory,
- 2. set SECHIBA_restart_in.nc to NONE in orchidee.def, and
- 3. relaunch the gcm.

2.3.3 Running with the simple bucket scheme

If VEGET=n (meaning that the vegetation is not activated) instead of y in file config.def, the soil scheme is a simple bucket (even if you compiled with makegcm -v true as done by install.sh when ran with veget=1).

2.3.4 Running with bucket scheme with imposed soil water content

Same as in § 2.3.3, you should run with VEGET=n in config.def. Evaporation is computed as the potential evaporation multiplied by the aridity coefficient vbeta, which is a function of the soil water content qsol0:

```
vbeta(i) = MIN(2.0*qsol/mx_eau_sol, 1.0)
```

(here mx_eau_sol=150mm). So, if qsol0 is constant, vbeta is constant as well. You can fix qsol0 to a chosen value qsol0_val (in mm), by adding in physiq.def the line qsol0_val; try for example qsol0_val=5 or 10, that result in vbeta values typical of summertime.

You can compare the turbulent fluxes for the austral summer (variables flat and sens in the LMZ output files) computed using the different options.

3 Nudging

- The program ce01.e (which was run by init.sh in the first part of this tutorial) created the file grilles_gcm.nc in the INITIAL directory. grilles_gcm.nc contains the longitudes and latitudes of the model grids (staggered grids for zonal wind, meridional wind and temperature). You can visualize the orography map as seen by the zoomed grid, by opening this file with ferret, ncview or grads and plotting the variable phis (which, despite its name, actually contains the surface altitude, in m). You can also easily plot the horizontal resolution of the model as the square root of the grid mesh area (variable aire, in m²).
- We will nudge the model with reanalysis fields. The script <code>get_era.sh</code>, in the directory <code>TUTO</code>, gets the reanalysis files. Run the script. It will also interpolate the winds on the model grid (by reading the model grid from file <code>grilles_gcm.nc</code>). You should end up with a directory called <code>GUIDAGE</code> (nudging in french) that contains two files, <code>u.nc</code> and <code>v.nc</code>. Note that for this tutorial we have given open access to a subset of the ERA-interim wind fields. ERA-interim files are stored at IDRIS, CCRT and Climserv, with restricted access. To access these files at IDRIS or on Climserv, you should contact Sophie Bouffies-Cloché (IPSL). For access at CCRT, contact Anne Cozic (LSCE). <code>get_era.sh</code> is a very simplified script for the tutorial, but more general scripts are available on the Subversion server of IGCMG:http://forge.ipsl.jussieu.fr/igcmg/browser/CONFIG/LMDZOR/branches/LMDZOR_v4/CREATE/SCRIPT. The file called <code>era2gcm.ksh</code> interpolates the ERA data on the GCM grid. This script can be called by another script (see <code>interp_from_era.ksh</code> for example) on multiple months, if you want to run long simulations.
- As in section 2.2, create a new directory, called for example SIMU1_nudged, which contains all the files you need to run a new simulation:

```
mkdir SIMU1_nudged
cd SIMU1_nudged
ln -s ../SIMU1/start.nc .
```

```
ln -s ../SIMU1/startphy.nc .
ln -s ../SIMU1/limit.nc .
ln -s ../SIMU0/sechiba_rest_out.nc sechiba_rest_in.nc
cp ../SIMU1/*.def .
rm -f used_*.def

Don't forget to also point to the nudging files:
ln -s ../GUIDAGE/u.nc .
ln -s ../GUIDAGE/v.nc .
```

- Take a look at the file guide.def. Nudging is activated for variables u and v only (as is often the case). The relaxation time is set to 3 hours inside the zoomed area (tau_max=0.125 days) and half an hour outside (tau_min=0.0208333 days). The smaller the relaxation time, the stronger the nudging. You can change the parameters of this file if you want.
- Add the line:

```
INCLUDEDEF=guide.def
in run.def.
```

• Run the model again with nudging:

```
../gcm.e > listing
```

- Make sure that the nudged simulation worked by comparing the winds of the outputs with the winds contained in the u.nc and v.nc files. You should see the same patterns, for the same time and altitude of course (for example look at the first output of the histhf.nc file at the 500 hPa level and compare with the first timestep of the u.nc and v.nc files).
- Compare the results of the simulations with and without nudging.