## Wikiprint Book

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## Simulation setup

This chapter describes how to setup your simulation.

## 1. Create submission directory

The configuration directory contains tools to compile (Makefile and AA\_make) and tools to run a simulation, i.e.

- two directories (EXPERIMENTS and GENERAL) allow you to create submission directories for your model configuration;
- if one or several submission directories (e.g. EXP00, OOL\_SEC\_STO, historical, EXP\_AER, etc...) have already been created, you can directly go to the <u>next step</u>.

In the EXPERIMENTS directory you will find subdirectories for each model configuration (included the one you work with). For example:

- IPSLCM5\_v5 includes LMDZOR, LMDZ
- IPSLCM5CHT\_v5 includes IPSLCM5, LMDZOR, LMDZORINCA
- LMDZOR\_v5 includes LMDZ
- LMDZORINCA\_v5 includes LMDZOR, LMDZ

Each of these subdirectories may contain a reference experiment (e.g. clim and amip for LMDZOR, NMHC\_AER, AER and GES for LMDZORINCA, piControl, historical and cie for IPSLCM5\_v5) and the file config.card which will be your simulation's initial setup.

Before preparing the working directory you must know which kind of simulation you want to perform. Then, you must copy the config.card file at the same level as the main Makefile.

For example, to perform a clim experiment with LMDZOR\_v5:

```
cd modipsl/config/LMDZOR_v5
cp EXPERIMENTS/LMDZOR/clim/config.card .
```

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The header of config.card contains the **JobName** field for which you must specify your simulation's name. Then run the ins\_job script that will in first time ask you, if you are working on TGCC, your id group, and then create a directory for your experiment. If you are working on IDRIS it will directly create a directory for your experiment.

In the previous example, a simulation called *DIADEME* is created:

```
cd modipsl/config/LMDZOR_v5
cp EXPERIMENTS/LMDZOR/clim/config.card .
ls
AA_Make Makefile EXPERIMENTS GENERAL config.card
vi config.card  # Change JobName=DIADEME
../../util/ins_job
ls
AA_Make Makefile EXPERIMENTS GENERAL DIADEME
```

The config.card file is deleted and a directory called DIADEME is created.

## 2. Contents of the submission directory

The contents of the new directory are described below.

```
cd DIADEME
ls
config.card COMP/ PARAM/ POST/ DRIVER/
```

## 2.1. config.card

The config.card file contains the settings of your simulation configuration. The file contains several sections with the simulation settings (e.g. name, duration, processors' number, post processing, initial state). Below is a list of the file sections:

#### 2.1.1. The [UserChoices] section

- JobName --> simulation name
- ExperimentName --> experiment name (following the CMIP5 nomenclature for the IPCC simulations)
- SpaceName --> variable indicating the type of a simulation. Choose between PROD, DEVT and TEST. SpaceName=TEST is a special case deactivating pack and storage.
- LongName --> description of your simulation
- TagName --> do not change this field; describes to which configuration family your experiment belongs
- ExpType --> do not change this field; allows you to find the EXPERIMENTS directory in which you are working
- DateBegin --> simulation start date (yyyy-mm-dd)
- DateEnd --> simulation end date. It must be the last day "included" in your simulation
- PeriodLength --> frequency of the executable run. This parameter can be 1M, 1Y or 10Y
- JobNumProcTot --> number of processors required by your simulation. This parameter depends on the configurations and on the resolutions. For example, a configuration containing LMDZ in 96x95 can use 32 processors at the most.

The parameters ExperimentName and SpaceName are optional. They impact on the path to the storage directory for the simulation output. SpaceName=TEST is a specific case which deactivate pack and storage at archive directory which means that the output will be stored only at SCRATCHDIR(curie) or WORKDIR(ada).

Example 1: The output directory for the following case will be IGCM\_OUT/LMDZOR/TEST/REINE/DIADEME

JobName=DIADEME ExperimentName=REINE SpaceName=TEST TagName=LMDZOR

The output directory will be IGCM\_OUT/LMDZOR/TEST/REINE/DIADEME

Example 2: without ExperimentName and SpaceName

JobName=DIADEME TagName=LMDZOR

The output directory will be IGCM\_OUT/LMDZOR/DIADEME

The character "\_" is not allowed in the variables JobName, ExperimentName and SpaceName

if SpaceName=TEST all output will be store on scratchdir (on curie) or workdir (on ada)

## 2.1.1.1. PeriodLength

The parameter PeriodLength allows you to determine the continuous length of a simulation period for your configuration, i.e. the frequency at which restart files are created. **The maximum PeriodLength for any LMDZ model is 1M**.

## 2.1.2. The [Restarts] section

The Restarts section allow to start from an existing simulation. This simulation can be found at the archive machine or at local scratch- or workdir. Activate by setting **OverRule=y**. All components(e.g. ATM, SRF, etc) will then use the same simulation as restart state.

| day of the experience used as restart for all component |
|---|
| ne restart simulation name for all components           |
|   |

RestartPath=\${ARCHIVE}/IGCM\_OUT/IPSLCM5A/DEVT/pdControl # Path Server Group Login

The root path for the RestartPath depend on the computing center. They are:

| \${ARCHIVE}                    | # The storage machine of the computing center    |
|--------------------------------|--|
|                                | # (CCCSTOREDIR or ERGON). This space can contain |
|                                | # tar of restarts or                             |
|                                | # usual restarts files                           |
| /ccc/store/cont003/dsm/login   | # TGCC   |
| /u/rech/ces/login              | # IDRIS  |
|                                |  |
| \${SCRATCHDIR}                 | # The large TGCC workspace (no backup)           |
|                                |  |
| /ccc/scratch/cont003/dsm/login | # This kind of space can contain                 |
|                                | # usual restarts files                           |
|                                |  |
| \${WORKDIR}                    | # The large IDRIS workspace (no backup)          |
|                                |  |
| /workgpfs/rech/ces/login       | # This kind of space can contain                 |
|                                | # usual restarts files                           |
|                                |  |

libIGCM manages the difference in treatment between a path pointing to restart files that are directly accessible (without pack) and a path pointing to restart files that are in tar format (after pack). The management is made according to the path you provided.

#### 2.1.3. The [ATM], ..., sections of the model components

This section for each of the model components allows you to:

- define the output frequency;
- define whether this component is installed which will only be considered if you specified OverRule=n in the [Restarts] section.

The possible settings for the RestartPath options are the same as for the [Restarts] section.

The possible settings for the WriteFrequency options are:

- 1M (monthly)
- 5D (5-day)
- 1D (daily)

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- HF (6-hour high frequency)
- HF3h (real-time 3-hour frequency specific to LMDZ)
- HF3hm (3-hour averaged high frequency specific to LMDZ)
- STN (instantaneous output only for the CFMIP stations specific to LMDZ).

|     | [ATM]   |  |
|-----|---|--|
|     | WriteFrequency="1M 1D"  | # Activate the writing frequency of this component                     |
|     | Restart=y   | <pre># If config_Restarts_OverRule == 'n' next 4 params are read</pre> |
|     | RestartDate=1999-12-31  | # Last day of the experience used as restart for this component if     |
|     | RestartJobName=piControl25  | # Define restart simulation name for this component                    |
|     | <pre>RestartPath=\${ARCHIVE}/IGCM_OUT/IPSLCM5A/PROD/piControl</pre> | # Path Server Group Login  |
| - 1 |   |  |

## WriteFrequency specific to the model components

• LMDZ : ([ATM]) Each of the frequencies settings 1M, 1D, HF, HF3h, HF3hm, and STN correspond to a given output file. For example, if you specify 1*M*, a histmth.nc file will be created. If you want to change the output frequency in the histmth file you must change the corresponding Imdz parameter file. See <u>here</u>.

### ORCHIDEE

• [SRF] : The first frequency corresponds to the output frequency for the sechiba\_history.nc file. The available frequencies are: xY, xM, 5D, 1D and xs, where x is an integer and s means seconds. This file is required. If you add HF, a second sechiba\_out\_2.nc file will be written with the 3H frequency.

- [SBG]: Only one frequency (xY, xM, 5D, 1D or xs) can be specified. It applies to the stomate\_history.nc file. The stomate\_history\_ipcc.nc file always contains daily outputs.
- INCA : the section WriteFrequency does not work. Click here to learn more about how to change the writing frequency.

#### 2.1.4. The section [Executable]

This section contains one line for each model component giving the executable's name in the bin/ directory and the executable's name copied to the working directory. You should only change this section if your executable is running in parallel using MPI and OpenMP or if you have changed the executable's name.

```
[Executable]
#D- For each component, Real name of executable, Name of executable for oasis, optional : number of MPI task, number of Op
ATM= (gcm.e, lmdz.x)
SRF= ("", "")
...
```

Note : (",") indicates that this component has no executable. It is defined in a library linked to another executable (e.g. Orchidee in LMDZOR or Inca in LMDZINCA).

# Example for coupled configuration : Atmosphere on 27 MPI process and 4 OMP threads per process, Ocean on 5 MPI process, IO Server on 1 MPI process.

```
[Executable]
#D- For each component, Real name of executable, Name of executable in RUN_DIR directory, Number of MPI process, Number of
ATM= (gcm.e, lmdz.x, 27MPI, 40MP)
SRF= ("" ,"" )
SBG= ("" ,"" )
OCE= (opa, opa.xx, 5MPI)
ICE= ("" ,"" )
MBG= ("" ,"" )
CPL= ("", "" )
IOS= (xios_server.exe, xios.x, 1MPI)
```

### 2.1.5. The [Post] section

The options of the [Post] section will allow you to set or disable the frequencies for submitting <u>post processing jobs</u> by changing the 5 following options (see the diagram below).

If you do not wish to run post processing jobs, you must specify NONE for both TimeSeriesFrequency and SeasonalFrequency.

RebuildFrequency and PackFrequency should not be disabled except in the case of running in expert mode.

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| RebuildFrequency=1Y       | <pre># Frequency of rebuild submission (use NONE for DRYRUN=3)</pre> |
|---------------------------|--|
| PackFrequency=1Y          | # If absent default to RebuildFrequency.                             |
| TimeSeriesFrequency=1Y    | # Frequency of post-processing submission (NONE if you don't want)   |
| SeasonalFrequency=2Y      | # Seasonal average period (NONE if you don't want,                   |
|                           | # 2Y at least, 10Y by default)                                       |
| SeasonalFrequencyOffset=0 | # Offset for seasonal average first start dates ;                    |
|                           | # same unit as SeasonalFrequency                                     |
|                           |  |

## 2.2. COMP directory

This directory contains the architecture (or map) of each model component. Each map specifies inputs and outputs required by a component.

Input files of each component are organized into different sections.

- [UserChoices] contains specific options. --> used by the component's drivers (e.g.: Imdz.driver)
- [InitialStateFiles] Initial conditions files such as vegetation maps, topography,... --> retrieved by the IGCM\_comp\_GetInputInitialStateFiles function
- [BoundaryFiles] Boundary conditions files such as forcings or a LAI --> retrieved by the IGCM\_comp\_GetInputBoundaryFiles function

- [SmoothFiles] Time-varying boundary conditions files such as aerosols --> retrieved by the IGCM\_comp\_GetInputSmoothFiles function
- [ParametersFiles] Parameters files such as namelist or the run.def file --> retrieved by the IGCM\_comp\_GetInputParametersFiles function
- [RestartFiles] Restart files --> retrieved by the IGCM\_comp\_GetInputRestartFiles function

#### 2.2.1. The [UserChoices] section

Contains several options which change the simulation setup by drivers files of the components (Imdz.driver, opa9.driver, ...). For example :

```
[UserChoices]
# Physics package to use :
# LMDZ_Physics=AP for standard/old physics(default), can be used with LMDZ4_AR5 or LMDZ5/trunk sources
# LMDZ_Physics=NPv3.1 for new physics, to be used with LMDZ5/trunk revision 1554 or later
LMDZ_Physics=AP
```

See the description for LMDZ here.

#### 2.2.2. The [InitialStateFiles] section

Files needed to create initial files. This section is not activated if you chose to start or continue from an existing simulation (Section [Restart] in config.card). The files in this list will be only copied at the startup of your simulation.

#### 2.2.3. The [BoundaryFiles] section

The files containing the boundary conditions are copied to the working directory.

The files in the **List** list will be copied at each integration period (one 1-month integration per period in general). A job can consist of several periods (PeriodNb).

The files in the ListNonDel list will only be copied for the first period of each job. These files will be accessible but will not change during the simulation.

Be very careful : if there is any space at the end of a line, libIGCM will not taking in account the next line in the list

#### 2.2.4. The [SmoothFiles] section

These are also files containing boundary conditions but their retrieval is only done at specific time integrals and it is not systematic. **1:12:** means that the file will be copied to the working directory at the first integration step and then every 12 iterations until the simulation is finished.

## 2.2.5. The [ParametersFiles] section

The parameter files of the component (namelist, run.def,...)

## 2.2.6. The [RestartFiles] section

The files providing the restart data. You must not change this section it is needed to link the jobs.

#### 2.2.7. The [OutputText] section

This section contains text files which will be produced during the simulation and model input parameter files. You might want to save these files.

```
[OutputText]
List= (NAME OF TEXT1 FILE, NAME OF TEXT2 FILE ....)
```

This files will be saved in tar stored in the output directory

• TGCC :

\$CCCSTOREDIR/IGCM\_OUT/TagName/[SpaceName]/[ExperimentName]/JobName/DEBUG

· IDRIS :

\$ARCHIVE/IGCM\_OUT/TagName/[SpaceName]/[ExperimentName]/JobName/DEBUG

## 2.2.8. The [OutputFiles] section

The netcdf files produced by the simulation are listed in this paragraph. This paragraph is associated with the [Post\_\*] sections.

```
[OutputFiles]
List = (OUTPUT_FILE_NAME, SAVE_PATH, POSSIBLE ASSOCIATED POST PROCESSING)
```

Refer to this chapter to learn everything about this section.

## 2.3. DRIVER directory

This directory contains the different *drivers* (predefined libIGCM functions for the component) of the different configuration's components. These drivers modify the parameter files of each component (\*.def, namelist, ...) setting the integration times, the outputs, and the forcing files.

Note : If this directory does not exist the *driver* files are located in the COMP directory.

## 2.4. PARAM directory

This directory contains input text files for the configuration's components.

## 2.5. POST directory

This directory contains configuration files for additional diagnostic output. Click here for more details.

## 3. Set up initial state for the simulation

When you setup a simulation make sure that the list of input files in each card file of the model components and the selected options correspond to your experiment.

There are three different ways to define your simulation's initial conditions:

- Start using restart files from an existing simulation by setting OverRule=y in the [Restart] section of the config.card file
- Start using different restart files from different simulations for each model component by setting Restart=y in each associated part of the config.card file. OverRule=n must be set in config.card.
- Use the default section InitialStateFiles in the comp.card file of the model components in config.card you must have OverRule=n and and Restart=n for this case.

## 3.1. Example for different restart

#### 3.1.1. Example with OverRule=y

If you wish to use the start state of a given simulation, set in config.card:

For the same case but if the simulation was performed by someone else, you must give the complete path of the directory, for example:

RestartPath=/u/rech/lab/plabxxx/IGCM\_OUT/IPSLCM5A/DEVT/pdControl
# or RestartPath=/dmnfs/contxxx/login/IGCM\_OUT/IPSLCM5A/DEVT/pdControl

#### 3.1.2. Example with OverRule=n and [COMP]/Restart=y

You can also distinguish the setup parameters for each model components. Set OverRule=n and use the Restart, RestartDate, RestartJobName and RestartPath parameters for each model component (section). For example, use restart files for the atmosphere but not for the surface component. For the surface component the InitialStateFiles will then be used :

```
#D-- ATM -
[ATM]
#
WriteFrequency="1M 1D HF"
# If config_Restarts_OverRule == 'n' all params are read
Restart= y
# Last day of the experience used as restart for this component
RestartDate=1999-12-30
# Define restart simulation name
RestartJobName=2L18
RestartPath=${ARCHIVE}/IGCM_OUT/IPSLCM5A/DEVT/pdControl
#
#D-- SRF -
[SRF]
#
```

```
WriteFrequency="1M"
# If config_Restarts_OverRule == 'n' all params are read
Restart= n
# Last day of the experience used as restart for this component
RestartDate=1999-12-30
# Define restart simulation name
RestartJobName=2L18
RestartPath=${ARCHIVE}/IGCM_OUT/IPSLCM5A/DEVT/pdControl
```

## 3.2. Note for LMDZ

To obtain exactly the same outputs in different simulations, you must choose the same LMDZ Bands files. This is explained in COMP/lmdz.card with the LMDZ\_NbPeriod\_adjust and LMDZ\_Bands\_file\_name parameters.

```
LMDZ_NbPeriod_adjust=0
# To force the use of this Bands file, set LMDZ_NbPeriod_adjust=0 and replace XXXXXXX by Restart Job Name
LMDZ_Bands_file_name=${ARCHIVE}/IGCM_OUT/IPSLCM5/CEPR00/ATM/Debug/CEPR00_Bands_96x95x39_3prc.dat_3
```

Click here for more details.

## 4. Main job of the simulation

The main job contains scripts that will be executed by the system. With libIGCM, this job is unique (in the beginning AA\_job and later Job\_MYJOBNAME) for all type of configurations. It contains all scripts to initialize a simulation, to summarize the chosen model configuration and to run identical experiments for all model components. It resubmits itself in order to continue the simulation if needed.

The job header depends on the machine type. It contains the job name and the parameters. **Real-times** must be chosen to match the specific classes for the computing machine and according to the simulation length (test or production).

At TGCC you must specify the project number: <u>#MSUB -A MY\_PROJECT</u>.

You should change the PeriodNb parameter in the job to change the number of runs in one job (see the example of computation in the next section) :

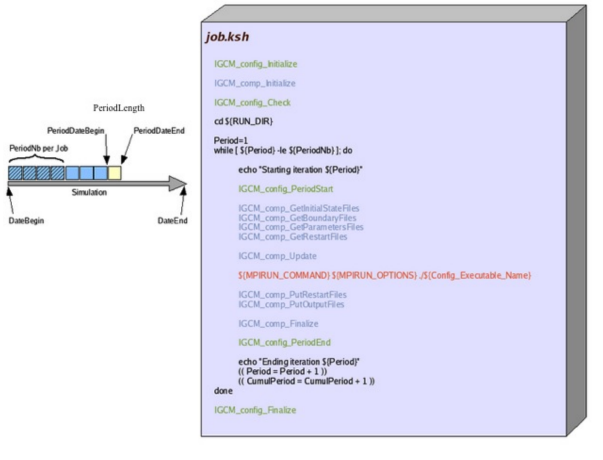
```
#D- Number of execution in one job
PeriodNb=1
```

In some cases, you must set a variable RUN\_DIR\_PATH in order to avoid deleting the working directory.

```
#D- Define running directory
#D- Default=${TMPDIR} ie temporary batch directory
#RUN_DIR_PATH=/workdir/or/scratchdir/of/this/machine
```

Here is the diagram of the steps in AA\_job :

# Script de référence : AA\_Job



## 4.1. Choosing PeriodNb

To avoid starting a lot of short jobs which might be queued, the production job starts n integrations (PeriodNb), whose length are PeriodLength.

These are calculated as followed: Time limit = PeriodNb \* max(Real time of a PeriodLength)

where Time limit is the requested time in the job header.

At the end of a simulation, the run.card file returns the used CPU time for each simulation step. This will allow you to perform this computation. It is therefore important, for each simulation with a new configuration, to perform a 1-3 month test to estimate beforehand the CPU time.

## 5. Prepare a new experiment

There are two ways to prepare a new working directory for your model configuration:

- 1. Start again from the first step described above by copying the desired config.card file to your configuration directory using a new JobName.
- 2. Copy an existing submission directory, delete the files created by the simulation, and change JobName in config.card.

#### For example:

```
cd modipsl/config/LMDZOR_v5
cp -r DIADEME CHOUCROUTE
cd CHOUCROUTE
```

```
rm -f Job_DIADEME run.card Script_Output_DIADEME.000001
vi config.card
    JobName=CHOUCROUTE
    ../../util/ins_job
```

The ins\_job script allows you to create a submission directory from a config.card file or if the directory already exists it allows you to only create the job corresponding to config.card. ins\_job will not overwrite a directory or an existing job.

## 5.1. Post-processing jobs

Jobs headers for post-processing have to be carefully checked, especially elapsed time limits. They are in libIGCM directory (xxx.job) and are adapted for IPSLCM5A with 1Y for RebuildFrequency and PackFrequency. Change time limits if you use larger frequencies.

## 6. Prepare ensembles with ins\_job -e

#### Remark ! What follows applies to IPSLCM5\_v5.

When IPSLCM5\_v5 is downloaded with ./model IPSLCM5\_v5 it will offer the possibility to launch experiments of the decadal type. To prepare an ensemble of simulations copy the config.card and ensemble.card files from the directory:

modipsl/config/IPSLCM5\_v5/EXPERIMENTS/IPSLCM5/decadal/

into the directory:

```
\texttt{modipsl/config/IPSLCM5\_v5}
```

Several types of ensemble simulations can be prepared by filling config.card and more importantly ensemble.card.

## 6.1. Config.card

The file config.card is filled as a regular config.card (ins\_job without the -e option).

The important lines for the ensemble set up are in the [UserChoices] section. Make sure that JobName and ExperimentName are filled with proper values.

The variable CalendarType should be consistent with the simulations from which you are initialising or else the restart file with the proper date may not exist.

The variables DateBegin and DateEnd will be overriden by variables present in ensemble.card.

```
#D-- UserChoices -
[UserChoices]
JobName=v3h4testB
#----- Short Name of Experiment
ExperimentName=v3h4testB
#---- DEVT TEST PROD
SpaceName=DEVT
LongName="IPSLCM5A CMIP5 DEVT phase decadal example with limited outputs."
TagName=IPSLCM5A
#D- Choice of experiment in EXPERIEMENTS directory
ExpType=IPSLCM5/decadal
#-- leap, noleap, 360d
CalendarType=noleap
#-- Experiment dates : Beginning and ending
#-- "YYYY-MM-DD"
DateBegin=2013-01-01
DateEnd=2022-12-31
#-- 1Y, 1M, 5D, 1D Period Length of one trunk of simulation
```

A section [Ensemble] should also be present. It contains the information that we want to prepare an ensemble simulation with variable EnsembleRun set to y and three unset fields to be filled in the config.card of each member after ins\_job -e has run.

```
[Ensemble]
#D- Ensemble run ? 'y' or 'n'
#D- If 'y', fill in ensemble.card !!
EnsembleRun=y
EnsembleName=
EnsembleDate=
EnsembleType=
```

#### 6.2. Ensemble.card

There are several sections in ensemble.card: [Ens\_PARAMETRIC], [Ens\_DATE] and [Ens\_PERTURB].

The choice of ensemble types is done by setting the variable active to y or n.

```
[Ens_PERTURB]
# active=y to use this ensemble type
active=y
```

We cover here the third section which allows to generate members from an initial condition which is perturbed by different means.

There are two ways to perturb the initial condition:

- · apply some random white noise of defined amplitude to the temperature field of the coupler component (CPL) restart file
- apply some previously generated 3D temperature perturbation map to the temperature field of the ocean component (OCE) restart file

Each method apply only to the relevant type of ensemble generation available inside [Ens\_PERTURB] as will be explained later.

Before detailing the different functionalities available in [Ens\_PERTURB] let us discuss the NAME variable. This variable should be set identical to the JobName variable, otherwise the script will fail to generate the proper files.

```
\ensuremath{\texttt{\#}} ensemble name (must be equal to JobName in config.card) NAME=v3h4testB
```

#### Periodic start dates

For this type of perturbed ensembles the following variables are left empty:

```
# member list (apply list of pattern to initial state)
MEMBER_LIST=()
# member list of names corresponding to each member
MEMBER_NAMESLIST=()
# member pattern global name
MEMBER_INITFROM=
# member pattern global directory for name
MEMBER_INITPATH=
....
# start dates list
NONPERIODIC=()
```

```
# length list for non periodic simulation (NOTE: use length above if not fill)
LENGTH_NONPERIODIC=()
...
# Path of Mask file
MASKPATH=
```

In ensemble.card, it is possible to specify a periodic list of start dates. Restart files will be generated for each member at each date starting from BEGIN\_INIT to END\_INIT with a periodicity of PERIODICITY. The variable MEMBER sets the number of members for each start date.

The following part of ensemble.card sets 10 members from 19900101 to 20000101 every 2 years each lasting 10 years:

# member nb (i.e nb of perturb initial restart for each date)
MEMBER=10
....
# periodic and member list simulations length
LENGTH=10Y
# start date of the first ensemble
BEGIN\_INIT=19900101
# start date of the last ensemble
END\_INIT=20000101
# timestep between each periodic simulation
PERIODICITY=2Y

This will produce 10 members starting at the dates : 19900101, 19920101, 19940101, 19960101, 19980101, 20000101. (PERIODICITY can be given in months for shorter periods)

Each time the restart file to be perturbed in order to produce each member is taken from the previous day of the start date : 19893112, 19913112, etc...

The directory in which the start date is retrieved is given by INITPATH and INITFROM. To restart from experiment v3h4BTxx in directory /ccc/store/cont003/gen2211/nguyens/IGCM\_OUT/IPSLCM5A/PROD/historical fill:

# Restart name INITFROM=v3h4BTxx # Restart directory INITPATH=/ccc/store/cont003/gen2211/nguyens/IGCM\_OUT/IPSLCM5A/PROD/historical

The way the perturbed member is generated depends on PERTURB\_BIN array. The first two elements are the most important. The first one is the executable to be used to produce the members, the second one is the component from which the restart is perturbed.

In the Periodic Case it is only possible to build the members by applying a randomly generated temperature pattern on the restart file of the coupler. PERTURB\_BIN should look like this:

PERTURB\_BIN=(AddNoise, CPL, sstoc, O\_SSTSST, 0.1)

The list is interpretted as follows:

- the used executable is AddNoise,
- the component is the coupler (CPL),
- · the restart file to perturb contains sstoc in its name,
- the variable to perturb in the restart file is O\_SSTSST,
- the randomly generated pertubation is in [-.05;+0.05] degrees

For each member (in our example we have ten) a new restart file for the coupler will be generated using the executable addnoise to add some randomly generated temperature perturbation.

For the year 1990, the corresponding restart file of member 1 will be stored in

\$WORKDIR/IGCM\_IN/v3h4testB190/v3h4testB190A/CPL/Restart/

Non-Periodic start dates

For this type of perturbed ensembles the following variables are left empty:

```
# member list (apply list of pattern to initial state)
MEMBER_LIST=( )
# member list of names corresponding to each member
MEMBER_NAMESLIST=()
# member pattern global name
MEMBER_INITFROM=
# member pattern global directory for name
MEMBER_INITPATH=
. . .
# start dates list
NONPERIODIC=()
# length list for non periodic simulation (NOTE: use length above if not fill)
LENGTH_NONPERIODIC=()
. . .
# start date of the first ensemble
BEGIN INIT=
# start date of the last ensemble
END INIT=
. . .
# Path of Mask file
MASKPATH=
```

The variable LENGTH must be set to something but is not used, PERIODICITY must be set to NONE:

```
# periodic and member list simulations length
LENGTH=10Y
...
# timestep between each periodic simulation (NONE for nonperiodic)
PERIODICITY=NONE
```

To set 10 members for the starting dates 1990 and 1992 for a duration of 10 years, set MEMBER, NONPERIODIC and LENGTH\_NONPERIODIC as follows:

```
# member nb (i.e nb of perturb initial restart for each date)
MEMBER=10
...
# start dates list
NONPERIODIC=(19900101 19920101)
# length list for non periodic simulation (NOTE: use length above if not fill)
LENGTH_NONPERIODIC=(10Y 10Y)
```

This results in 20 simulations in total.

The restart files to be perturbed to produce each member are sought in directory INITFROM which PATH is INITPATH.

```
# Restart name
INITFROM=v3h4BT00
# Restart directory
INITPATH=/ccc/store/cont003/gen0826/labetoul/dmf_import/IGCM_OUT/IPSLCM5A/PROD/historical
```

This will result in using restarts from experiment v3h4BT00 located in directory /ccc/store/cont003/gen0826/labetoul/dmf\_import/IGCM\_OUT/IPSLCM5A/PROD/historical.

The perturbation executable must be AddNoise.

PERTURB\_BIN=(AddNoise, CPL, sstoc, O\_SSTSST, 0.1)

#### List of members for a single start date

For this type of perturbed ensembles the following variables are left empty:

```
# member nb (i.e nb of perturb initial restart for each date)
MEMBER=
# timestep between each periodic simulation (NONE for nonperiodic)
PERIODICITY=NONE
# start dates list
NONPERIODIC=()
# length list for non periodic simulation (NOTE: use length above if not fill)
LENGTH_NONPERIODIC=()
```

It is important to leave PERIODICTY set to NONE and LENGTH\_NONPERIODIC as an empty list: the list of member method only works for a single start date and neither with periodic start dates nor with non periodic start dates.

The variables BEGIN\_INIT and END\_INIT are set to the same date, only BEGIN\_INIT will be used to provide the start date of the simulation for each member.

# start date of the first ensemble
BEGIN\_INIT=20560101
# start date of the last ensemble
END\_INIT=20560101

The variable LENGTH is the computation time which is the same for all members.

```
# periodic and member list simulations length
LENGTH=10Y
```

MEMBER\_NAMESLIST is the list of names given to each member. It gives the names of the subdirectories from which the Job is submitted for each member as well as the subdirectories in which the results are stored for each member.

MEMBER\_LIST is the list of perturbation maps files names prefix to apply to the restart file. It is implied that the files are named prefix.nc.

MEMBER\_INITFROM is the directory in which the perturbations maps are stored.

MEMBER\_INITPATH is the path to this directory.

```
# member list (apply list of pattern to initial state)
MEMBER_LIST=(OWN3DT_A, OWN3DT_B, OWN3DT_C, OWN3DT_D)
# member list of names corresponding to each member
MEMBER_NAMESLIST=(OWN3DTA, OWN3DTB, OWN3DTC, OWN3DTD)
# member pattern global directory name
MEMBER_INITFROM=OWN3DTpf
# member pattern global directory for name
MEMBER_INITPATH=/ccc/work/cont003/gen2211/nguyens/PERTU/VECTORS
```

The variables INITFROM and INITPATH are still used to point to the directory where the restart files including the one to be perturbed are available.

# Restart name
INITFROM=piControl2
# Restart directory
INITPATH=/ccc/store/cont003/dsm/p86caub/dmf\_import/IGCM\_OUT/IPSLCM5A/PROD/piControl

For the member list perturbation type we use the executable AddPertu3DOCE and set PERTURB\_BIN this way:

# perturbation type
PERTURB\_BIN=(AddPertu3DOCE, OCE, restart, tn, ORCA2\_mesh\_mask.nc)

The elements of the list mean:

- the executable to be called to generate the perturbation is AddPertu3DOCE
- the component is the Ocean (OCE)
- the restart file to perturb is \*restart\*.nc
- the field to perturb in the restart file is tn
- the meshmask file to tell if the gridcell is land or sea is ORCA2\_mesh\_mask.nc

The path to the mesh mask file is given in MASKPATH.

# Path of Mask file MASKPATH=/ccc/cont003/home/gen2211/nguyens/addpertu

Once config.card and ensemble.card properly filled the directories containing the jobs to launch the simulations are created by issuing the command:

ins\_job -e