

Monday morning 2:22:00

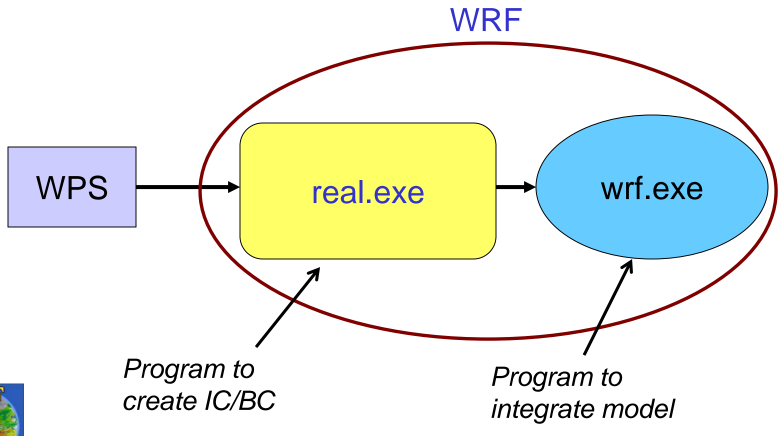
# Set Up and Run WRF

(real data)

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# WRF System Flowchart





### **Outline**

- Running WRF code
  - Things to check before you run..
  - Running real-data case
- Basic runtime options for a single domain run (namelist)
- Check output
- Simple trouble shooting



## Before You Run ..

- Top directory is now wrf/
- Make sure appropriate executables are created in WRF/main/ directory:
  - real.exe executable to create IC/BC
  - wrf.exe executable for model integration
  - ndown.exe utility
  - tc.exe utility routine for TC bogusing

 If you are working with real data, be sure that files for <u>a few time periods</u> from WPS are correctly generated:



- met\_em.d01.\*

## WRF test case directories

You have these choices in **WRF/test/** 

(choices made at compile time. E.g. compile em real):

em\_real



## Steps to Run

- 1. cd to *run*/ or the *test/em\_real*/ directory
- 2. Move or link WPS output files to the directory for real-data cases
- 3. Edit *namelist.input* file for the appropriate grid dimensions and times of the case
- 4. Run a initialization program (real.exe)
- 5. Run model executable, wrf.exe



# WRF/run directory

```
README.namelist
LANDUSE. TBL
GENPARM. TBL
SOILPARM. TBL
VEGPARM. TBL
URBPARM, TBL
RRTM DATA
RRTMG SW DATA
RRTMG LW DATA
CAM ABS DATA
CAM AEROPT DATA
ozone.formatted
ozone lat.formatted
ozone plev.formatted
aerosol.formatted
aerosol lat.formatted
aerosol lon.formatted
aerosol plev.formatted
gribmap.txt
grib2map.tbl
.... (a total of 60 files)
```

description of namelists

These are model physics data files: they are used to either initialize physics variables, or make physics computation faster

\* Some of these files are text files,

for grib 10

hence editable



# WRF/run directory after compile

```
LANDUSE . TBL
SOILPARM. TBL
VEGPARM. TBL
GENPARM, TBL
URBPARM, TBL
RRTM DATA
                             An example after
RRTMG SW DATA
                             em real case
RRTMG LW DATA
                             compile
ozone, formatted
ozone lat.formatted
ozone plev.formatted
namelist.input - copied from ../test/em real/namelist.input
real.exe -> ../main/real.exe
wrf.exe -> ../main/wrf.exe
ndown.exe -> ../main/ndown.exe
.... (a few more)
```





 If you have compiled the em\_real case, you should have:

```
real.exe - real data initialization program
wrf.exe - model executable
ndown.exe - program for doing one-way nesting
tc.exe - program for TC bogusing
```

These executables are linked to:

```
WRF/run
and
WRF/test/em_real
```



One can go to either directory to run.

## WRF/test/em\_*real* directory

```
LANDUSE.TBL -> ../../run/LANDUSE.TBL
GENPARM.TBL -> ../../run/GENPARM.TBL
SOILPARM.TBL -> ../../run/SOILPARM.TBL
VEGPARM.TBL -> ../../run/VEGPARM.TBL
URBPARM.TBL -> ../../run/URBPARM.TBL
RRTM DATA -> ../../run/RRTM DATA
RRTMG SW DATA -> ../../run/RRTMG SW DATA
RRTMG LW DATA -> ../../run/RRTMG LW DATA
ozone.formatted -> ../../run/ozone.formatted
ozone lat.formatted -> ../../run/ozone lat.formatted
ozone_plev.formatted -> ../../run/ozone_plev.formatted

    editing required

namelist.input
real.exe -> ../../main/real.exe
wrf.exe -> ../../main/wrf.exe
ndown.exe -> ../../main/ndown.exe
.... (many more)
```



 One must successfully run WPS to prepare data required, and create met\_em.\* files for multiple time periods for initial and boundary conditions

 Move or link WPS/metgrid output files to the run directory:

```
cd test/em_real
ln -s ../../WPS/met_em.d01.*
```



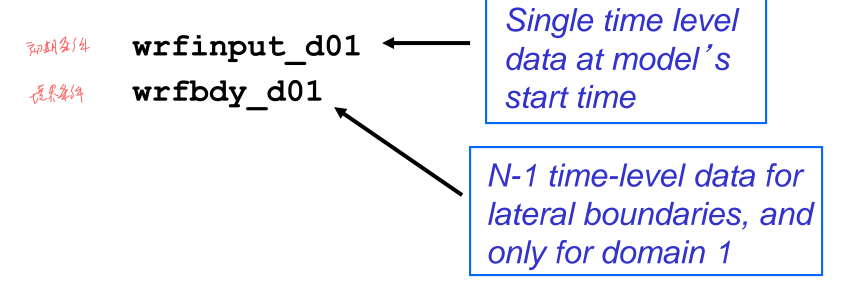
- Edit namelist.input file for runtime options (at mininum, one must edit &time\_control for start, end and integration times, and &domains for grid dimensions)
- Run the real-data initialization program:

```
./real.exe if compiled serially / SMP, or 和斯维·德特特 a Met CDF 生版的 serial ですか
```

mpirun -np N./real.exe for a MPI job where N is the number of processors requested.



 Successfully running real.exe will create model initial and boundary files:



N: the number of time periods processed





- Typing 'ncdump -v Times wrfbdy\_d01' will give you, for a 24 hour period, 6 hourly data interval:
  - .. a bunch of prints and at the end:

#### data:

```
Times =

"2005-08-28_00:00:00",

"2005-08-28_06:00:00",

"2005-08-28_12:00:00",

"2005-08-28_18:00:00";
```

\* BC data consists of values at the start of the time interval and rate of change in the time interval.



Run the model executable by typing:

```
./wrf.exe >& wrf.out &
or
mpirun -np N ./wrf.exe &
```

 Successfully running the model will a create model <u>history</u> file:

```
wrfout_d01_2005-08-28_00:00:00
```

Based on start date set in namelist

and a <u>restart</u> file if **restart\_interval** is set to a time within the range of the forecast time:



Exact time at a restart

```
wrfout_d01_2005-08-28_00:00:00
```

#### Based on start date set in namelist

```
start_year
start_month
start_day
start_hour
start_minute
start_second
end_year
end_month
end_day
end_hour
end_minute
end_second
```

```
2008, 2008, 2008,
= 08,
      08,
            08,
= 28, 28, 28,
= 00, 00, 00,
= 00, /00, 00,
→ 00, / 00,
            00,
= 2008, 2008, 2008,
= 08,
      08,
            08,
= 29, 29, 29,
= 00, 00, 00,
= 00, 00, 00,
= 00, 00,
            00,
```



# **Basic namelist Options**



## What is a namelist?

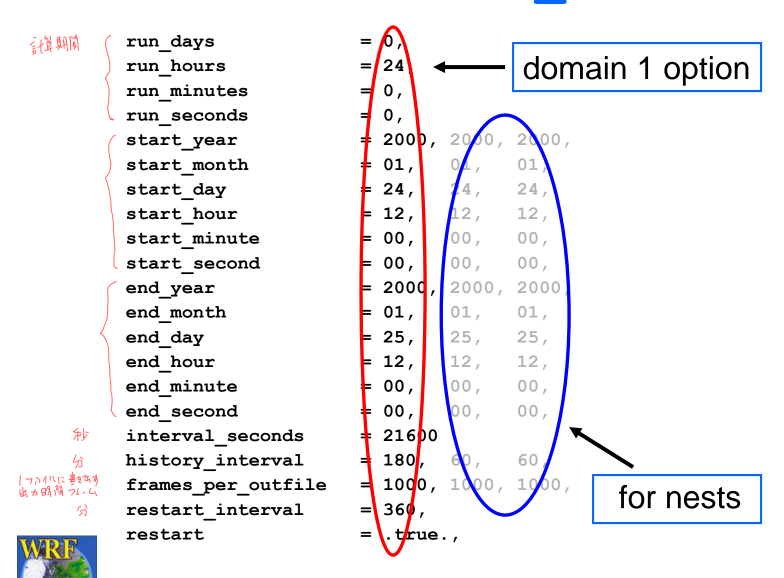
- A Fortran namelist contains a list of runtime options for the code to read in during its execution. Use of a namelist allows one to change runtime configuration without the need to recompile the source code.
- Fortran 90 namelist has very specific format, so edit with care:

```
&namelist-record - start
/ - end
```

- As a general rule:
  - Multiple columns: domain dependent
  - Single column: value valid for all domains



## namelist record &time\_control



# Notes on &time\_control

- run\_\* time variables:
  - Model simulation length: wrf.exe and domain 1 only
- start\_\* and end\_\* time variables:
  - Program real will use WPS output between these times to produce lateral (and lower) boundary file
  - They can also be used to specify the start and end of simulation times for the coarse grid if run\_\*
     variables are not set (or set to 0)



# Notes on &time\_control

- interval\_seconds:
  - Time interval between WPS output times, and lateral BC (and lower BC) update frequency
- history\_interval:
  - Time interval in <u>minutes</u> when a history output is written (<u>note</u> output is instantaneous)
  - If the time\_step cannot be evenly divided by history\_interval, then nearest time step output is used
  - The time stamp in a history file name is the time when the history file is first written, and multiple time periods may be written in one file. e.g. a history file for domain 1 that is first written for 1200 UTC Jan 24 2000 is

wrfout\_d01\_2000-01-24\_12:00:00



# Notes on &time\_control

- frames\_per\_outfile:
  - Number of history times written to one file
- restart\_interval
  - Time interval in minutes when a restart file is written
  - By default, restart file is not written at hour 0
  - A restart file contains only one time level data, and its <u>valid time</u> is in its file name, e.g. a restart file for domain 1 valid for 0000 UTC Jan 25 2000 is

```
wrfrst_d01_2000-01-25_00:00:00
```

restart:



whether this is a restart run

# Notes on &time control

#### Example 1: all output times are in a single file

```
history_interval = 180, 60, 60, frames_per_outfile = 1000, 1000, 1000, wrfout_d01_2000-01-24_12:00:00
```

#### Example 2: each output file only contains a single time

```
history_interval = 180, 60, 60, frames_per_outfile = 1, 1, 1, wrfout_d01_2000-01-24_12:00:00 wrfout_d01_2000-01-24_15:00:00 wrfout_d01_2000-01-24_18:00:00
```



## Notes on restart

- What is a restart run?
  - A restart run is a continuation of a model run
- How to do a restart run:
  - In the first run, set <u>restart\_interval</u> to a value that is within the model integration time
  - A restart file will be created. e.g. wrfrst\_d01\_2000-01-25\_00:00:00
- When doing a restart run:
  - Set restart = .true.,
  - Set start time to restart time
  - Set run\_\* to be the hours remaining in the run



## &time control

```
io_form_history = 2,
io_form_restart = 2,
io_form_input = 2,
io_form_boundary = 2,
```

#### For large files:

io\_form\_restart = 102 :
write output in patch
sizes: fast for large grid
and useful for restart file

#### IO format options:

- = 1, binary
- = 2, netcdf (most common)
- = 4, PHDF5
- = 5, Grib 1
- =10, Grib 2
- =11, pnetCDF



#### namelist record &domains

```
time step
                        = 180
time step fract num
                        = 0,
time_step_fract_den = 1,
                        = 1,
max dom
                        = 74,
e we
                         = 61,
                                     , nest<sup>1</sup>,
e sn
                        = 33,
e vert
                        = 32,
num metgrid levels
num_metgrid_soil_levels
                         = 30000, 10000, 333
dx
                        = 30000, 1000, 3
dy
                        = 1.0, 0.996, 0.99, 0.98, ... 0.0
eta levels
                        = 5000,
p top requested
```



## Notes on &domains

- time\_step\_fract\_num, time\_step\_frac\_den:
  - Time step for model integration in seconds
  - Fractional time step specified in separate integers of numerator and denominator
  - Typically 5 to 6xDX (DX is grid distance in km) 30km to 30x 6= 180(5) が 造れなる
- e\_we, e\_sn, e\_vert.
  - Model grid dimensions (staggered) in X, Y and Z directions, need to match those defined in *geogrid* program
- num\_metgrid\_levels:
  - Number of metgrid (input) data levels
- num\_metgrid\_soil\_levels:
  - Number of soil data levels in the input data
- ⇒ Both can be found by typing ncdump -h met\_em.d01.<date> | more
- *dx, dy*.
- WRIF
- grid distance: in meters

## Notes on &domains

- p\_top\_requested:
  - Pressure value at the model top
  - Constrained by the available data from WPS
  - Default is 5000 Pa (recommended as lowest model top)
- eta\_levels:
  - Specify your own model levels from 1.0 to 0.0
  - If not specified, program real will calculate a set of levels
    - V4 has a new and better way to compute the levels
  - Use a minimum of 33 or more levels with 5000 Pa model top to limit vertical grid distance < 1 km. Use more vertical levels when decreasing horizontal grid sizes.



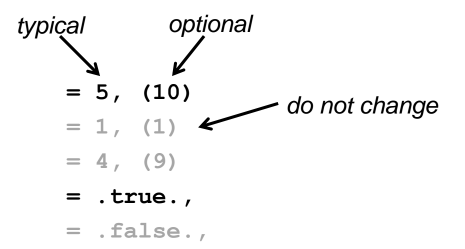
## namelist record <a href="mailto:bdy\_control">bdy\_control</a>

spec bdy width

spec\_zone
relax\_zone

specified

nested



May change relax\_zone
and spec\_bdy\_width
(spec\_zone + relax\_zone
= spec\_bdy\_width)

\* Wider boundary zone may work better for coarser driving data



### Other namelists

#### &physics

Model physics options

#### &dynamics

- Damping, diffusion options
- Advection options
- In 4.0, the hybrid vertical coordinate option is the default.
   Turn it off by setting the following for *real* and *wrf*:

$$hybrid_opt = 0$$



## Where do I start?

- Always start with a namelist template provided in a test case directory, whether it is an ideal case, or a real data case.
  - A number of namelist templates are provided in test/test\_<case>/ directories

```
For example: in test/em_real/, there are namelist.input.4km ~ 4 km grid size namelist.input.jun01 ~ 10 km grid size namelist.input.jan00 ~ 30 km grid size
```



## Where do I start?

- For different applications, please refer to p5-38 to 5-40 of the ARW User's Guide:
  - 2 or 4 km microphysics-only runs
  - -20 30 km, 2 3 day runs
  - Antarctic region
  - Tropical storm forecasting
  - Regional climate
  - Try physics suites (since V3.9)



## Where do I start?

- Use document to guide the modification of the namelist values:
  - run/README.namelist
  - test/em\_real/examples.namelist
  - User's Guide, Chapter 5 (online version has the latest)
  - Full list of namelists and their default values can be found in Registry files: Registry.EM\_COMMON, registry.io\_boilerplate (for IO options) and other registry files - look for character string 'namelist'



# To run a job in a different directory...

- Directories run/ and test\_<case>/ are convenient places to run, but it does not have to be.
- Copy or link the content of these directories to another directory, including physics data files, wrf input and boundary files, wrf namelist and executables, and you should be able to run a job anywhere on your system.



# **Check Output**



# Output After a Model Run

Standard out/error files:

```
wrf.out, or rsl.* files
```

Model history file(s):

```
wrfout_d01_<date>
```

Model restart file(s), optional



## Output from a multi-processor run

The standard out and error will go to the following files for a MPI run:

```
mpirun -np 4 ./wrf.exe →
```

```
rsl.out.0000 rsl.error.0000 rsl.out.0001 rsl.error.0001 rsl.error.0002 rsl.error.0002 rsl.error.0003
```

There is one pair of files for each processor requested

## What to Look for in a standard out File?

Check run log file by typing

```
tail wrf.out, or tail rsl.out.0000
```

You should see the following if the job is successfully completed:

wrf: SUCCESS COMPLETE WRF



# How to Check Model History File?

• Use ncdump:

```
ncdump -v Times wrfout_d01_<date>
to check output times. Or
  ncdump -v U wrfout_d01_<date>
to check a particular variable (U)
```

- Use ncview (great tool!)
- Use post-processing tools (see talks later)



## What is in a *wrf.out* or *rsl* file?

Model version, decomposition info:

```
Ntasks in X 2, ntasks in Y 4
WRF V4.0 MODEL
```

Time taken to compute one model step:

```
Timing for main: time 2000-01-24_20:03:00 on domain 1: 0.89475 elapsed seconds Timing for main: time 2000-01-24_20:06:00 on domain 1: 0.09011 elapsed seconds Timing for main: time 2000-01-24_20:09:00 on domain 1: 0.08634 elapsed seconds Timing for main: time 2000-01-24_20:12:00 on domain 1: 0.09004 elapsed seconds
```

Time taken to write history and restart file:

```
Timing for Writing wrfout_d01_2000-01-25_00:00:00 for domain 1: 0.07091 elapsed seconds
```

Any model error prints:

```
エラーの 可必性
5 points exceeded cfl=2 in domain 1 at time 4.200000 MAX AT i,j,k: 123 48 3 cfl,w,d(eta)= 4.165821
```

→ An indication the model has become numerically unstable

# Simple Trouble Shooting



## Often-seen runtime problems

- module\_configure: initial\_config: error reading namelist: &dynamics
  - > Typos or erroneous namelist variables exist in namelist record & dynamics in namelist.input file
- input\_wrf.F: SIZE MISMATCH: namelist
  ide,jde,num\_metgrid\_levels= 70 61 27; input
  data ide,jde,num\_metgrid\_levels= 74 61 27
  - > Grid dimensions in error



# Often-seen runtime problems

- Segmentation fault (core dumped)
  - > Often typing 'unlimit' or 'ulimit -s
    unlimited' or equivalent can help when this
    happens quickly in a run, and on a small computer
- If you do: grep cfl rsl.error.\* and see
  121 points exceeded cfl=2 in domain 1 at time
  4.200000 MAX AT i,j,k: 123 48 3 cfl,w,d(eta)=
  4.165821
  - Model becomes unstable due to various reasons. If it happens soon after the start time, check input data, and/or reduce time step.



### References

 Information on compiling and running WRF, and a more extensive list of namelist options and their definition / explanations can be found in the User's Guide, Chapter 5

