

WRF test case directories

You have these choices in WRF/test/ (made at compile time):

	em_real	}	3-d	imensional	l r	eal-data – real.exe			
	em_quarter_ss		٦						
	em_b_wave								
	em_les		}	3d ideal	~				
	em_tropical_cyc	lor							
	em_heldsuarez		J						
VRF	em_hill2d_x		ſ						
	em_squall2d_x					≻ ideal.exe			
	em_squall2d_y		ļ			ideal.exe			
	em_grav2d_x		Í	2d ideal					
	em_seabreeze2d_	x	J						
	em_scm_xy		}	1d ideal	,				
E	Mesoscale & Microscale Meteorological Laboratory / NCAR 5								

WRF/run directory





Running a Real-Data Case • If you have compiled the em real case, you should have: **real.exe** - real data initialization program wrf.exe - model executable Running a Real-Data Case 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. 9 Mesoscale & Microscale Meteorological Laboratory / NCAR Mesoscale & Microscale Meteorological Laboratory / NCAR 10 WRF/test/em real directory Running a Real-data Case LANDUSE.TBL -> ../../run/LANDUSE.TBL GENPARM.TBL -> ../../run/GENPARM.TBL One must successfully run WPS to prepare SOILPARM.TBL -> ../../run/SOILPARM.TBL data required, and create met em. * files for VEGPARM.TBL -> ../../run/VEGPARM.TBL URBPARM.TBL -> ../../run/URBPARM.TBL multiple time periods for initial and boundary RRTM DATA -> ../../run/RRTM DATA RRTMG SW DATA -> ../../run/RRTMG SW DATA conditions 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 Move or link WPS/metgrid output files to the namelist.input editing required run directory:

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



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ln -s ../../../WPS/met em.d01.* .

cd test/em real

Running a Real-data Case

- 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

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



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Running a Real-data Case

- 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.



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Running a Real-data Case

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



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Running a Real-data Case

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:

wrfrst_d01_2005-08-28_12:00:00



Exact time at a restart



WRF

➔ One can go to either directory to run.

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Running an *Idealized* Case

Then run the ideal initialization program: ./ideal.exe

The input to this program is typically a sounding file (file named *input_sounding*), or a pre-defined 2D input (e.g. *input_jet* in em_b_wave case).

Running ideal.exe only creates WRF initial condition file: wrfinput_d01



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Running an Idealized Case

• To run the model interactively, type

./wrf.exe>& wrf.out &

for single processor (serial) or SMP run. Or

mpirun -np N ./wrf.exe &

```
for a MPI run (3D cases only)
```

 Successful running of the model executable will create a model history file called wrfout_d01_<date>
 e.g. wrfout d01 0001-01-01 00:00:00



Based on start date set in namelist (dates are important for radiation physics)

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Running an Idealized Case

Note that wrfbdy file is not needed for idealized cases.

Instead, the boundary condition options are set in the namelist.input file. For example, these are for options in east-west, or x direction:

periodic_x	= .true.,
symmetric_xs	= .false.,
symmetric_xe	= .false.,
open_xs	= .false.,
open_xe	= .false.,



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Running an Idealized Case

wrfout_d01_0001-01-01_00:00:00

Based on start date set in namelist

	1	\sim		
start_year	F	0001,	0001,	0001,
start_month	=	01,	01,	01,
start_day	=	01,	01,	01,
start_hour	=	00,	00,	00,
start_minute	=	00,	00,	00,
start_second	7	00,	00,	00,
end_year	=	0001,	0001,	0001,
end_month	=	01,	01,	01,
end_day	=	01,	01,	01,
end_hour	=	00,	00,	00,
end_minute	=	120,	120,	120,
end_second	=	00,	00,	00,

Running an Idealized Case

- Edit namelist.input file to change options.
- For your own case, you may provide a different sounding.
- You may also edit <u>dyn_em/module_initialize_<case>.F</u> to change other aspects of the initialization (more on Thur.)

Note:

- For 2D cases and baroclinic wave case, ideal.exe must be run serially
- For all 2D cases, wrf.exe must be run serially or
 with SMP



For the 1D case, compile and run serially

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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
```

- As a general rule:
 - Multiple columns: domain dependent
 - Single column: value valid for all domains
- A namelist file may contain a number of records

- end



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Basic namelist Options

namelist record &time control



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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)



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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

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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



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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







Where do I start?

- For different applications, please refer to p5-36 to 5-38 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)



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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.

WRF

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'



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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 wrfrst d01 <date>



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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



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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.out.0002	rsl.error.0002
rsl.out.0003	rsl.error.0003

There is one pair of files for each processor requested



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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)





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
- Also see nesting talk and demonstration tomorrow.

