

**Using the WRF-ARW on the UWM *mortimer* HPC**  
**Guide for WRF-ARW Version 3.9.1.1**  
**May 23, 2018**

***Introduction***

This guide is designed to facilitate basic compilation and execution of the WRF-ARW v3.9.1.1 model core using OpenMPI on the UWM *mortimer* HPC. I cannot cover every option available to you when installing or running the model; for all of the gory details, see the WRF-ARW User's Guide, available online at:

[http://www2.mmm.ucar.edu/wrf/users/docs/user\\_guide\\_V3.9/contents.html](http://www2.mmm.ucar.edu/wrf/users/docs/user_guide_V3.9/contents.html)

I also recommend reading over and familiarizing yourself with the information found in the UWM HPC User's Guide, available online (from on-campus machines only) at:

<http://www.peregrine.hpc.uwm.edu/Webdocs/uwm-pc-user-guide.pdf>

Note that the steps outlined in this User's Guide are generally applicable to both earlier and newer versions of the WRF-ARW model; however, there may be subtle differences in compilation options between each version of the model. Always read the installation information from the appropriate version of the WRF-ARW User's Guide first!

***What You Need to Know First***

*Compile Environment:* We will use the Intel compilers to build WRF on the UWM HPC. The Intel compilers are available through a module on the UWM HPC that will set all of the necessary path, library, and execution variables for you automatically. To load this module, run:

```
module use /sharedapps/ICC/modulefiles/  
module load icc/18.0.1
```

*MPI Libraries:* You will be using OpenMPI to run parallelized (multiple CPU) jobs on the cluster. To correctly configure OpenMPI, you need to load the appropriate module. This is done by issuing the following command *after* you issue the `module load icc/18.0.1` command listed above:

```
module load openmpi/3.0.0_v2
```

*NetCDF Libraries:* Before compilation and execution of the model code, you'll need to point to a netCDF installation on the server. This is also done using modules:

```
module use /bigraid-02/LS/evans36/modules/  
module load netcdf-4.6.1
```

If you wish to enable netCDF4-formatted output, set the appropriate environment variable:

```
export NETCDF4=1
```

Note that the WRF-ARW namelist.input variable `use_netcdf_classic` (in `&time_control`) must also be set to `.false.` in order to output in netCDF4 format.

*GRIB2 Libraries:* To use GRIB2-formatted data as inputs to WRF-ARW, you'll need to point to the GRIB2-related libraries on the server. To do so, load the following module:

```
module load grib2-libs-2018q2
```

I recommend adding all of the above commands to your `~/.bashrc` file so that they are loaded automatically each time you log in. To do so, issue the following command:

```
nano ~/.bashrc
```

In the nano text editor, simply add the commands from above to the end of the file, save the file (Ctrl-O), exit nano (Ctrl-X), and then log out and log back in to the supercomputer. There are other variables that you may need to set along the way during WRF or WPS compilation and/or run-time; I will alert you to those where appropriate.

### ***Mortimer Access Considerations***

*Remote Access:* mortimer is accessible *only* from machines with `uwm.edu` IP addresses. Thus, if you wish to connect to mortimer from off-campus, you will first need to connect to the UWM VPN (Virtual Private Network). Three steps are involved. The first is to visit <https://vpn.uwm.edu/>, login with your ePanther credentials, and download and install the appropriate Pulse Secure VPN client. The second is to e-mail Dan Siercks and request that your ePanther ID be given VPN access credentials to mortimer. The third, completed each time you wish to access mortimer, is to run the VPN client and login to it with your ePanther credentials. Once you are connected to the VPN, you should be able to access mortimer as if you were on campus.

*Data Transfer:* To transfer data from mortimer to your laptop or desktop using `sftp` or `scp`, you must specify the remote port that you wish to access. For your data, which resides on `/bigraid-02`, this is port 22102. This port is specified with the `-P #####` flag, i.e.,

```
sftp -P 22102 <user>@login.mortimer.hpc.uwm.edu
```

*Visualization:* By default, only command-line resources are available on mortimer. Graphical, or XWindows, resources are available only on a dedicated visualization node. This node is accessed using port 22201, i.e.,

```
ssh -Y -p 22201 <user>@login.mortimer.hpc.uwm.edu
```

Alternatively, if you are connected to mortimer (on the login node rather than a computational node), you can issue the following command to access the visualization node:

```
ssh -Y vis-1
```

Note that you must be on a Linux/Unix machine, one of the Macs in EMS W434, or using the MobaXterm package for Windows to be able to use graphical resources via the visualization node.

If you wish to use the ImageMagick (display, convert, etc.) programs on the visualization node, you must first issue the following command (all on one line):

```
module load /sharedapps/pkg-  
2017Q2/etc/modulefiles/pkgsrc/2017Q2-non-exclusive
```

---

## ***Part I: Obtaining & Compiling the Model***

### **Connect to a Compute Node on mortimer**

To begin, we wish to connect to a compute node on mortimer. Because the WRF model requires a lot of memory in order to successfully compile, the typical “slurm-shell” command will not work for this task. Instead, we manually connect from the mortimer login node using srun:

```
srun --mem-per-cpu=8gb --ntasks=1 --pty --preserve-env @$ $SHELL -l
```

Subsequently, your command prompt will change slightly, reflecting that you are now on a compute node (compute-#-##) rather than the login node.

### **Obtaining the WRF Model Code**

You can download the latest version of the WRF model code from:

[http://www2.mmm.ucar.edu/wrf/users/download/get\\_sources.html](http://www2.mmm.ucar.edu/wrf/users/download/get_sources.html)

From this website, you need to download the “WRF model tar file” and “WPS code tar file.” Download these tar files to a working directory within your “data” directory and untar them using the “tar -zxvf” command for each file. I recommend doing this directly on mortimer using the wget command. You should also review the “Known Problems” pages for both WRF and WPS and download any fixed code that is made available for each. We will first install the WRF v3.9.1.1 model code itself, followed by the pre-processor.

### **Installing the WRF v3.9.1.1 Model Code**

Once you have unzipped the WRF tar file, you need to switch into the newly created WRFV3 directory. First, open the arch/Config\_new.pl script with a text editor. Change line 250 (\$I\_really\_want\_to\_output\_grib2\_from\_WRF) from FALSE to TRUE, then save the file and exit the text editor. Next, issue “./configure” (without the quotes) to the command line to start model configuration. A list of configuration options will appear. You will want to choose option 15, “15. (dmpar) INTEL (ifort/icc)”. It will then ask you for a nesting option. Unless you plan on using advanced nesting options with the model, I recommend using option #1, the

default, for basic nesting capabilities.

Once configuration is complete, open the `configure.wrf` file that is created with a text editor. Change what comes after the equal sign for both `DM_FC` and `FC` – lines 155 and 157 – to read `mpif90`. Save this file, then exit out of the text editor.

If you wish to use this WRF installation with the Data Assimilation Research Testbed software, a couple of additional edits to the WRF Registry are needed before compiling the model. First, open `Registry/Registry.EM_COMMON` with a text editor. Search for `h_diabatic`. Change `rdu` to read `i012rhdu`. Next, search for `refl_10cm`. Change `hdu` to read `i012rhdu`. Save this file, exit the text editor, and return to the main `WRFV3` directory.

Finally, compile the model by issuing the following command:

```
./compile em_real
```

In all, compilation will take approximately 60 minutes to complete. The job may appear to hang when compiling several portions of the model code; this is okay. A successful compilation will produce “`ndown.exe`,” “`real.exe`,” “`tc.exe`,” and “`wrf.exe`” files in `WRFV3/main`. After this has completed, return to the login node by typing `exit` and pressing `enter`.

### Installing the WRF Pre-Processor Code

Installing the WPS code is similar to installing the WRF model code itself. Once you have unzipped the WPS tar file, switch into the newly created “WPS” directory. Ensure that this directory is on the same level as the `WRFV3` directory but is not *in* the `WRFV3` directory.

To enable GRIB2 support for WPS, ensure you have loaded the `grib2-libs` module introduced in the outset of this document. Next, move to a compute node, just as was done above for WRF. After having done so, issue a “`./configure`” command to the command line to start model configuration. A list of configuration options will appear. Choose the option that reads most similar to, “`Linux x86_64, Intel compiler (dmpar)`” (currently option 19).

After configuration is complete, open up the `configure.wps` file with a text editor. Edit the entry for `NCARG_LIBS` to read like the following:

```
NCARG_LIBS = -L/bigraid-05/LS/evans36/software/ncl/lib -lncarg -lncarg_gks -lncarg_c \  
             -L/usr/lib64 -lX11 -lXext -lpng -lz -lcairo -lfontconfig -lpixman-1 \  
             -lfreetype -lexpat -lpthread -lbz2 -lXrender -lgfortran -lgcc
```

This change is necessary to ensure that utilities requiring the NCAR Graphics or NCL libraries are built. After making these changes, save `configure.wps` and exit back to the command line. Once this is done, compile the code (`./compile`). Compilation may take 1-3 minutes. You can safely ignore any warnings to the effect of “ignoring unknown option ‘-f90=ifort.’” Once compilation has completed, look for “`geogrid.exe`,” “`ungrib.exe`,” and “`metgrid.exe`” in the current directory; this is your sign of success. If one or more of these programs are missing, check your `configure.wps` file to ensure that the library paths are all set correctly (as above), then run `./clean -a` and start the

process anew. Once done, type exit and press enter to return to the login node.

## ***Part II: WRF Pre-Processor (WPS)***

### **What does the WRF Pre-Processor do?**

The WPS has three tasks: defining model domains, extracting initialization data for the model simulation from GRIB files, and horizontally interpolating that data to the model domain and boundaries. All of this is accomplished through the use of a namelist file and a few command line options. If you are working through this tutorial and using the WRF-ARW model for the first time, I recommend that you do so alongside the related material within the “Basics for Running the Model” section of the WRF-ARW Online Tutorial, available at:

<http://www2.mmm.ucar.edu/wrf/OnLineTutorial/Basics/index.html>

### **Step 1: Defining a Domain**

Defining a domain is done through the `geogrid.exe` program of the WPS. Options for the domain are set in the `namelist.wps` file. Open this file in some text editor. The first two sections, `&share` and `&geogrid`, are the only two sections of this file to worry about now.

In `&share`, assuming you are not creating a nested model run, change `max_dom` in 1. Change `start_date` and `end_date` to the appropriate dates and times. These take the form of 'YYYY-MM-DD\_HH:MM:SS'. The interval between input times of your model data is specified by `interval_seconds`; for three hourly data, this will be 10,800. Note that unless you are doing a nested run, only the first option in each list matters. Information on nested runs can be found at the end of this document.

In `&geogrid`, `e_we` and `e_sn` define the size of your domain in gridpoints, with `e_we` defining the east-west size and `e_sn` defining the north-south size. Change these to appropriate values. `geog_data_res` defines the horizontal resolution of the geographic data files that you wish to use to setup your domain and has four options: 30s, 10m, 5m, and 2m. Generally 5m is fine for a grid spacing of about 20 km or larger; switch down to 2m or 30s for lower grid spacing values. If you want to include inland lake information with the 30s data set, you can use “30s\_with\_lakes” in lieu of 30s. There are other options available, and I recommend that you review the WRF-ARW User’s Guide for details.

`dx` and `dy` control your grid spacing; generally they should be equal to one another and are given in meters (default = 30000 = 30 km). `map_proj` deals with the desired map projection of your run; `lambert`, `mercator`, or `lat-lon` are suggested for most tasks, with the Lambert projection favored for mid-latitude simulations and the Mercator projection favored for simulations at tropical latitudes. `ref_lat` and `ref_lon` are the center point of your domain in latitude and longitude, respectively. Note that for west longitudes, `ref_lon` should be negative. The remaining fields vary depending upon which map projection you use. For instance, the Lambert projection requires three additional fields: `truelat1`, `truelat2`, and `stand_lon`. `truelat1` and `truelat2` define the “true” latitudes for the Lambert map projection; unless moving to the southern hemisphere, the default values should be

fine. `stand_lon` specifies the longitude parallel to the x-axis for conic and azimuthal projections; this value should generally be close to that of `ref_lon`. The Mercator projection requires just one additional field, `truelat1`. The Latitude-Longitude projection requires three additional fields: `pole_lat`, `pole_lon`, and `stand_lon`. This projection is recommended only for global runs and for such runs should not require any changes to `pole_lat` and `pole_lon`.

Finally, `geog_data_path` defines the path to where the geographic data resides on the server. Set this to `'/bigraid-05/LS/evans36/WRF/geog'`. Once these variables are set, save the `namelist.wps` file and return to the command line.

To perform a visual check on your domains' extent to ensure they meet your requirements, `ncl` and the `util/plotgrids_new.ncl` script may be used. First, log on to the visualization node following the instructions earlier in this document, then change into your WPS directory. Load the `ncl` module:

```
module use /bigraid-02/LS/evans36/modules/  
module load ncl-6.4
```

Finally, run the script:

```
ncl util/plotgrids_new.ncl
```

This will produce a display containing your domain(s) with the relevant underlying geography. To exit from this window, simply click anywhere inside of it. If you are satisfied with the domain(s), proceed to run `geogrid.exe` as below; if not, edit the domain particulars in `namelist.wps` and use `ncl` and the `plotgrids_new.ncl` script to re-visualize the domain(s) until you are satisfied.

Once ready, you can run `geogrid.exe` using a `slurm-shell`, i.e.,

```
slurm-shell  
./geogrid.exe
```

Once it is done (and it should give you a success message if it successfully completed), check that you have a `geo_em.d01.nc` file in the current directory; this ensures that it worked correctly.

## Step 2: Getting Model Data

Extracting model data from GRIB files is accomplished through the `ungrib.exe` program. There are two steps you do need to do before running the program: linking the appropriate variable table (`Vtable`) and linking the appropriate GRIB data.

Residing in the `WPS/ungrib/Variable_Tables/` directory are a series of `Vtable.xxx` files, where the `xxx` denotes a model name. These files tell the `ungrib` program about the format of the data files to be degribbed. If you are using the GFS model, for instance, you'll note a `Vtable.GFS` file in that directory. In the main WPS directory, issue the following command to link this `Vtable` file:

```
ln -s ungrib/Variable_Tables/Vtable.GFS Vtable
```

where you would substitute for GFS as appropriate for your initialization data set. For data sets that do not have a corresponding Vtable file in the default WRF installation, you will either need to create your own using the GRIB packing information from your data set or find one that someone else has already made for that data set.

Next, you need to link your model data GRIB files to the WPS directory. You can obtain GRIB data used to initialize the model from NCEP's NOMADS data server and/or UCAR's Research Data Archive for most data sources. If you need help finding data, please ask! Download these files to the supercomputer, identify where you have placed these files on the supercomputer, and issue the following command:

```
./link_grib.csh /path/to/model/data/model_data*
```

where you will replace /path/to/model/data with the appropriate path and model\_data.t00z\* with the appropriate file name format of the data files that you wish to link. This will create a series of GRIBFILE.xxx files in the WPS directory.

Before running the ungrid program, clear out all old GRIBFILE., FILE:, and PFILE: files that may exist to avoid inadvertent errors when running the model. Finally, run ungrid.exe in a similar fashion to geogrid.exe. If all goes well, you'll see a success message on screen and multiple files of the format FILE:YYYY-MM-DD\_HH will be present in the WPS directory.

### Step 3: Interpolating Model Data

Finally, to horizontally interpolate the model data (obtained in Step 2) to the domain (obtained in Step 1), the metgrid.exe program is used. At this point, except in rare circumstances, you will not need to change any of the variables in the namelist.wps file.

Instead of running metgrid.exe directly, as with geogrid.exe and ungrid.exe, we use a job submission script to run metgrid.exe on multiple CPUs. A job submission script specific to our model configuration takes the form:

```
#!/bin/sh
#SBATCH -n ##
#SBATCH --mem-per-cpu=2gb
module use /sharedapps/ICC/modulefiles/
module load icc/18.0.1
module load openmpi/3.0.0_v2
module use /bigraid-02/LS7/evans36/modules/
module load netcdf-4.6.1
module load grib2-libs-2018q2

mpirun ./metgrid.exe
```

The second and third line tell the SLURM scheduler, which is used by mortimer to schedule jobs on multiple CPUs, how many processors to use (replace ## with some number greater than one; for metgrid, I recommend 16) and to use 2 GB of RAM for each processor. We saw an instance of the mem-per-cpu directive in the third line when we used `srun` to connect to a compute node with 8 GB of memory to compile WRF/WPS; in general, however, `metgrid.exe` (and `real.exe` and `wrf.exe` to come) only require <2 GB of RAM per processor. The last line uses the `mpirun` program to run `metgrid.exe` on the specified number of processors.

Place the above lines in a new text file named `metgrid.sbatch` (or similar), change ## accordingly, and then save the file. To run metgrid, you submit this script to the job scheduler:

```
sbatch metgrid.sbatch
```

Assuming that the necessary resources exist to start your job immediately, metgrid will take 1-30 minutes to complete, with larger amounts of time necessary for longer-duration simulations and/or larger numbers of simulation domain grid points. You may check on its progress by examining the `metgrid.log.0000` file in the WPS directory, or by using `squeue` to look for your `metgrid.sbatch` job. Once metgrid has finished, ensure that you have a series of `met_em.d01.YYYY-MM-DD_HH:00:00` files in the WPS directory. If so, you're done with the WPS and can skip ahead to Part III of this document. If not, check the `metgrid.log.0000` file for possible insight into any errors that may have occurred at this step.

### **Advanced Uses: Multiple Domains**

If you want to set up for a run using multiple domains, it is fairly simple to do so. When editing the `namelist.wps` file, the following things will be different than as presented in Step 1 above:

- Under `&share`, set `max_dom` to 2 (or how many domains you wish to have).
- Edit the second listing in the `start_date` and `end_date` options.
- Under `&geogrid`, change the second listing in `parent_grid_ratio` to whatever downscaling factor you wish to have for the inner domain. The default of 3 is fine for most circumstances (e.g. will take a 30 km outer domain and create a 10 km grid spacing inner domain).
- Change the second listings of `i_parent_start` and `j_parent_start` to where in the outer domain you wish the lower left of the inner domain to begin.
- Change the second listings `e_we` and `e_sn` to the desired size values of the inner domain. (Note: the values for these must be some integer multiple of `parent_grid_ratio` plus 1.)
- Change `geog_data_res` as needed.
- You will not need to change `parent_id` from 1 unless you wish to create further inner domains that are not based off of the outer domain.

Note that if you have more than two domains, simply add a comma at the end of the second listing under the options listed above and manually type in your third (and beyond) values.

### **Advanced Uses: “Constant” Input Data Sources**

If you want to use a data set as a “constant” value, such as SST data, simply follow steps 1 and 2

above only for the GRIB files containing this constant data, noting that you will be doing this for just one time. Then, in `namelist.wps` under the `&metgrid` section, add a line called `constants_name`, e.g.

```
constants_name = 'SST_FILE:YYYY-MM-DD_HH'
```

where the file name is whatever the output file from the `ungrib.exe` program is named. In the example above, it is an explicitly named (using the prefix option in `&ungrib` in `namelist.wps`) SST data file. If you are using multiple data sets, make sure they have different prefix names so as to not overwrite one data set with the other inadvertently! To do this, edit the `prefix` listing under `&ungrib` in the `namelist.wps` file to reflect the desired prefix name (often for the constant data set), then change it back when re-running it for the actual model input.

### **Advanced Uses: Time-Varying SST, Sea Ice, Albedo, and Vegetation Data**

For long-duration simulations, or shorter-duration simulations of phenomena such as tropical cyclones that significantly influence fields such as sea surface temperature, you may wish to use time-varying inputs for surface fields that are typically held constant throughout the duration of a WRF-ARW model simulation. While WPS geographic data can describe the climatological evolution of albedo and vegetation fraction through time, external time-varying input data are needed in order for sea surface temperature and/or sea ice to be updated throughout the duration of your model simulation.

Sometimes, these data may be provided in the same input GRIB files you use to provide initial and lateral boundary conditions for your model simulations. In this case, no additional steps are necessary at the WPS stage. Other times, however, you may wish to use external data in lieu of those provided by your atmospheric model data set. In such cases, first use `ungrib` to process your atmospheric data. Next, link the other data sources' GRIB files to the WPS directory, link the appropriate `Vtable` to the WPS directory, edit `namelist.wps` to change the prefix entry under `&ungrib` to some new descriptive name, then run `ungrib` for these data. Finally, add the prefix for these data to the `fg_name` entry under `&metgrid` in `namelist.wps` (e.g., `fg_name = 'FILE','SST'` if you used SST as your new prefix), then run `metgrid.exe`.

I recommend reading through the WPS advanced tutorial information, particularly that related to `METGRID.TBL`, if you desire to make use of this option for SST data. Care must be taken to ensure that the datasets are blended appropriately.

[http://www2.mmm.ucar.edu/wrf/users/tutorial/201801/wps\\_advanced.pdf](http://www2.mmm.ucar.edu/wrf/users/tutorial/201801/wps_advanced.pdf)

There are some additions to `namelist.input` for WRF-ARW when using time-varying surface data sets. Please see the companion section to this one in Part III below for the relevant information. More advanced uses of multiple input data sources (e.g. multiple time-varying data sets, ocean mixed layer depth information, etc.) are detailed within the WRF-ARW User's Guide.

### ***Part III: Configuring and Running the WRF Model***

Except in the case of a nested domain or idealized simulation, there are two programs that will be used to setup and run the WRF model: `real.exe` and `wrf.exe`. Both of these programs are housed in the `WRFV3/run/` directory; change over to that directory now. We'll first use `real.exe` to take the data from the WPS and get it ready for use in the model, then use `wrf.exe` to actually run the model. All of this is accomplished on the command line with no GUI options available. For more information, please refer to Chapter 5 of the WRF-ARW User's Guide.

### Step 1: Real-Data Initialization

Before editing any of the files necessary for this step, first link the `met_em.d01.*` files from the WPS to the current working directory (`WRFV3/run/`) by issuing the following command:

```
ln -s ../../WPS/met_em.d01.* .
```

From here, we can move on to editing the `namelist.input` file with the necessary parameters. Many of the parameters in the first few sections of `namelist.input` will be the same as those in the `namelist.wps` file from the WPS program, so it might be useful to have those parameters handy at this time.

The `namelist.input` file has several parts, each with multiple variables and multiple options for each of those variables. In particular, you will see sections headed by `&time_control`, `&domains`, `&physics`, `&fdda`, `&dynamics`, `&bdy_control`, `&grib2`, and `&namelist_quilt`; some of these will be edited, others will not. Note that this is not intended to be an end-all listing of the options available to you here, particularly in terms of physics packages. Refer to the section of Chapter 5 of the WRF-ARW User's Guide entitled "Description of Namelist Variables" for more information on all of these options. The meanings of many of these variables are readily apparent, so I will only cover those that are not. As noted before, many of these values are the same as those input in the `namelist.wps` file from the WPS section. Only edit values in the first column (if there are multiple columns for a given variable) for now.

The `&time_control` section of the `namelist.input` file is where you will input the basic model timing parameters. Change `history_interval` to the time (in minutes) between output times you wish for model output. Otherwise, simply change all values above `history_interval` (except for `input_from_file`) to the appropriate values and leave all values below `history_interval` alone. If you want output in netCDF4-format, add a `use_netcdf_classic` entry and set it to `.false,`.

The `&domains` section of the `namelist.input` file is where you will be inputting information about your model's domain. Your first option relates to the model's time step. If you wish to use a fixed time step, set the `time_step` variable to a value (in seconds) that is approximately 6 times as large as your model grid spacing in kilometers. For example, for a 15 km model simulation, set this value to 90. It is helpful, but not necessary, if this `time_step` value is evenly divisible into 3600, the number of seconds in an hour. (If this is desired, you may modify `time_step` from `6*grid spacing` to some other similar value.)

Set the values from `max_dom` to `e_sn` to their appropriate values from `namelist.wps`. Set `e_vert` to the desired number of vertical levels. Slightly more complicated is `num_metgrid_levels`. For this

value, open a terminal window to the WRFV3/run directory and issue the following command:

```
ncdump -h met_em.d01.YYYY-MM-DD_HH:00:00 | grep
num_metgrid_levels
```

where you put in the desired time of one of the met\_em files. In the output from ncdump, look for the num\_metgrid\_levels toward the top of the screen, then cancel out using Control-C. Edit the namelist.input file variable to match this. For NAM input data, this value will be 40; for recent GFS input data, this value will be 32; and for older GFS input data, this value will be 27. Next, set dx and dy to the appropriate values from namelist.wps. Ignore the rest of the options for now; these are generally only relevant to nested runs.

The &physics section is where you will choose what physics packages you wish to include in your model. Refer to the User's Guide for what numeric values you need to select for each of these parameters. Since v3.9, WRF has handled this via physics suites; however, you can override any of the parameterizations in a given suite by manually adding the relevant entries to the namelist. For example, the mp\_physics variable defines what microphysics package you wish to use. Longwave and shortwave physics packages are defined in ra\_lw\_physics and ra\_sw\_physics. radt is a time step increment and should be set to the same as dx in kilometers (e.g. set this to 18 for dx = 18 km). Surface physics packages are handled with sf\_sfclay\_physics (surface layer) and sf\_surface\_physics (land-surface model). Boundary layer parameterizations are given in bl\_pbl\_physics. bldt is a time step increment; setting this equal to zero will match the model time step. Cumulus parameterization is handled by cu\_physics, with the time step to calls to that package given by cutd. Setting cutd to 0 works the same as for bldt. Set ifsnow to 1. The value for num\_soil\_layers will depend on the land-surface model chosen; for the NOAH and NOAH-MP land-surface models, this should be set to 4. The urban canopy model may be enabled by setting sf\_urban\_physics to 1. If modeling a tropical cyclone, the following line may be added to enable the usage of a modified surface flux formulation appropriate for tropical cyclones:

```
isftcflx = 2,
```

You may also want to consider employing the 1-dimensional oceanic mixed layer model for such simulations. See the WRF-ARW User's Guide for more information on this option.

Ignore the &fdda section. This handles four-dimensional data assimilation options and will not be used (or maybe even present in the namelist) unless specifically performing data assimilation. In general, you will not need to edit anything in &dynamics either; however, the diff\_opt and km\_opt variables may be tweaked to modify how the model handles diffusion and eddy coefficients. Refer to the User's Guide for more if you choose to modify those variables. I do recommend changing gwd\_opt from 1 to 0, however, to turn off the gravity wave drag option. Otherwise, you should not need to edit any other data in namelist.input.

## Step 2: Running the Model

The WRF model utilizes two programs, real.exe and wrf.exe, to prepare to and to actually run the model respectively. These programs are both run in parallel (e.g. on multiple machines) using OpenMPI on the cluster. To do so, you will need an appropriate job submission script. An example

is given below for real.exe; you may use the same one for wrf.exe if you change all instances of “real” within the text to “wrf.”

```
#!/bin/sh
#SBATCH -n ##
#SBATCH --mem-per-cpu=2gb
module use /sharedapps/ICC/modulefiles/
module load icc/18.0.1
module load openmpi/3.0.0_v2
module use /bigraid-02/LS/evans36/modules/
module load netcdf-4.6.1
module load grib2-libs-2018q2

mpirun ./real.exe
```

You will need to replace ## with the number of processors on which to run real.exe (or wrf.exe). This number should ideally be some multiple of 8 up to and including 96. The usage of more than 96 CPUs is generally discouraged to avoid excessive use of limited cluster resources. You will want to make sure, however, that you use an identical number of processors for real.exe and wrf.exe as well as for any and all simulation(s) you may conduct for a given research project.

Please also note that not every simulation will require 2 GB of memory per processor (the --mem-per-cpu=2gb flag); some will require less, while some will require more. You are encouraged to check the memory usage of each job using:

```
ssh -t compute-#-## top
```

where #-## is replaced with a compute node on which your job runs. This can be identified using the `squeue` command and looking for the entry corresponding to your running job, then identifying the nodes listed in the last column of that entry. You’ll want to look for the value in the “VIRT” column for one of your processes (e.g., real.exe, wrf.exe, or metgrid.exe) and set the value of --mem-per-cpu to a value somewhat higher than the actual usage.

Once the job submission script is ready, submit it to the cluster using sbatch, e.g.,

```
sbatch (name-of-submission-script)
```

After running sbatch, you will want to exit from the slurm-shell prompt by hitting Ctrl-D on the keyboard (or, alternatively, type exit and hit enter). You can monitor the progress of real.exe by looking at the rsl.error.0000 file in the WRFV3/run directory or by running `squeue -j #`, where # = the job number returned to you upon running sbatch. Once real.exe has finished, you run wrf.exe in a similar fashion utilizing a nearly identical job submission script. Model progress can be examined in the same manner as for real.exe. Once the model has completed, you are ready for post-processing.

### **Common Errors: SIGTERM Statements in the rsl.\* Files**

If you are getting SIGTERM errors in your rsl.\* files that keep the model from successfully completing, check to ensure that all relevant parameters between your namelist.wps and namelist.input files are identical. If these are identical, then you may be running into an issue where your simulation requires more memory than is available on the selected nodes (i.e., you've already requested the maximum available per node). To increase the amount of available memory, simply add additional processors in multiples of 8 to the real.exe and wrf.exe job submission scripts.

### **Advanced Uses: Adaptive Time Steps**

With the adaptive time step option, the WRF model will modify the time step up and down as the model integrates in order to find the most efficient yet computationally stable time step. Oftentimes, this speeds up model integration by 25-60%. Note that this option is discouraged for research applications. To use this option, simply add the following line (including the ending comma) immediately below the max\_dom option within the &domains section of the namelist:

```
use_adaptive_time_step           = .true.,
```

### **Advanced Uses: Two-Way Nesting**

Most options to get a two-way nest going are handled with one single run of the WRF model and through the namelist.input file. When editing this file, you will note multiple column listings for some of the variables; these extra columns handle information for the inner nest(s). Edit these variables to match the desired values for the inner nest, using the values for the outer nest as a guide. Variables that you did not edit for the single domain run but will need to be edited for a nested run include input\_from\_file, fine\_input\_stream, max\_dom (the total number of nests), grid\_id (1, 2, 3, etc.), parent\_id (generally one less than the grid\_id), i/j\_parent\_start (where in the outer domain you want the inner grid lower left hand corner to be), parent\_grid\_ratio (generally 3 is a good number), parent\_time\_step\_ratio (generally at or near the parent\_grid\_ratio), and feedback (1 is yes, where the inner grid writes back to the outer one; requires an odd value for parent\_grid\_ratio). Also, num\_metgrid\_levels needs to be changed for the nests as well to the number of WRF model levels; see the procedure above to see how to check this.

Notice that I did not discuss input\_from\_file and fine\_input\_stream in the previous paragraph. There are many interrelated options to consider for these two variables. The first option is to have all of the fields interpolated from the coarse domain rather than created on their own. This is probably the fastest method, but also may not lead to as accurate of results as otherwise expected. In this case, input\_from\_file would be .false. and you don't need to worry about fine\_input\_stream. The second option is to have separate input files from each domain, with input\_from\_file set to .true.; unfortunately, this means that the nest has to start at the same time as the outer domain. The final option also has input\_from\_file set to .true., but requires you add a new line after input\_from\_file for fine\_input\_stream and set it to a value of 2 for all domains. This allows you to start the nest at a different time than the initial time.

For a nested run, you run real.exe and wrf.exe as before. Make sure you link over any necessary met\_em.d02 (or .d03, etc.) files to the working directory before running real.exe.

## Advanced Uses: Time-Varying SST, Sea Ice, Albedo, and Vegetation Data

If you desire to use time-varying surface datasets, new entries must be added to `namelist.input` prior to running `real.exe` and `wrf.exe`. The first three of these new entries go at the bottom of the `&time_control` section (i.e., before the ending `/`) and take the form:

```
io_form_auxinput4      = 2
auxinput4_inname       = "wrflowinp_d<domain>"
auxinput4_interval     = 360, 360, 360,
```

Note that `auxinput4_inname` should appear exactly as it does above. If the input data are available at a different frequency than every 6 h, change `auxinput4_interval` to match this frequency (where `auxinput4_interval` is given in minutes).

A new entry must also be added to the `&physics` section of `namelist.input`. This takes the form:

```
sst_update = 1
```

Note that `sf_ocean_physics` should be set to 0 if this option is activated.

## Other Advanced Uses

Recent versions of the WRF-ARW model have added a number of additional options to the model, including the ability to use a digital filter to aid in model initialization; apply nudging (to some specified data set) over some specified time interval; use a 1-D or 3-D ocean model primarily for tropical cyclone simulations; and many, many others. In addition, a number of new physical parameterization packages have been added, many of which have their own additional `namelist` parameters that can be tweaked to (theoretically) improve model performance for specific forecasting applications. A complete listing of all that is possible with the WRF-ARW model is available within Chapter 5 of the WRF-ARW User's Guide. If you are in doubt as to whether a particular option should be utilized for your simulations, or what particular selections following from using such an option should be for your simulations, please ask.

---

## *Part IV: Post-Processing*

### Post-Processing Using ARWpost and GrADS

WRF-ARW may be used with the ARWpost post-processor. ARWpost converts the WRF model output data into a GrADS-readable format. In the same level directory as the WRFV3 and WPS directories, `untar` (using a `tar -xzvf ARWpost.tar.gz` command) the post-processor code obtained from the WRF-ARW User's Site. This will create a new ARWpost directory.

Now, switch into the ARWpost directory. Next, run the configuration program by typing `./configure`. Choose option 2 for Linux using the Intel compiler. Once configuration is complete, open `configure.arwp` with a text editor. Change the entry for `LDFLAGS` in line 35 so

that it reads:

```
LDFFLAGS = -L/bigraid-05/LS/evans36/software/netcdf-4.6.1/lib -lnetcdf -lnetcdff
```

Save the file and exit the text editor. Next, compile the post-processor by opening up a slurm-shell and then running `./compile` (assuming that all modules have been loaded and environment variables set prior to doing so). If everything goes according to plan, you'll get an `ARWpost.exe` file in the current directory. At this point, you may reset all of your environment variables to their defaults (such as by logging out and back in again).

Options for the `ARWpost` program are handled in the `namelist.ARWpost` file. First, a note about the `namelist.ARWpost` file. Options that are active in the `namelist` are found between the `&_____` label and the ending `/` for that section; options that are inactive (i.e. commented out) are founded after the `.`. Make sure that you have the right options set in your active options sections when editing this `namelist` file. With that said, let's explore the `namelist.ARWpost` file section by section. In the `&datetime` section, set the appropriate values for `start_date` and `end_date`. The value for `interval_seconds` will be equal to that from the `history_interval` variable in the `namelist.input` file from running the model, except here in seconds instead of minutes. In the `&io` section, `input_root_name` refers to where your `wrfout_d01*` files are located. Set this equal to the full path there plus `wrfout_d01` and it will read in the correct file. The `output_root_name` variable refers to what you want to call the GrADS output control and data files; set these accordingly.

What is output is controlled in the rest of the `&io` section of the `namelist.ARWpost` file. There are several options for the variables `plot` and `fields`. Generic options are listed by default; other options for `plot` and `fields` are given below the `mercator_defs` variable line. In general, it is probably best to output all available variables, meaning to change `plot` to `'all_list'` and add in all of the variables in the `fields` definition after `mercator_defs` to the `fields` definition immediately below `plot`. Note that the lines past `mercator_defs` before `&interp` are only comment lines; they will not modify your output! I do, however, recommend adding the following line immediately after `mercator_defs` (but before the `/`) to split post-processed output into individual files:

```
split_output = .true.
```

There are other options available; see Chapter 9 of the `WRF-ARW User's Guide` or the `README` file in the `ARWpost` directory for more information.

Finally, under `&interp`, you have the choice of how you want the data output, as controlled by `interp_method`. The default value of 1 is for pressure level data; 0 is for sigma levels while -1 is for user-defined height levels. The levels to interpolate to, no matter what you choose for `interp_method`, are given by `interp_levels`. The default pressure levels are good for most uses; however, you may desire to add additional levels. Examples for pressure and height levels are given in the last three lines of the file; I don't believe you can control specific sigma levels. Finally, if you wish to interpolate data below ground, you can set the `extrapolate` variable in this section to `.true.`.

Once the `namelist` file has been edited to your liking, simply run `./ARWpost.exe` on the command line and wait for data to come out. Once it is done and gives you a success message, you

can move on to GrADS to view your output. A working installation of GrADS v2.1.1.b0 can be enabled by loading the appropriate module:

```
module use /bigraid-02/LS/evans36/modules/  
module load grads-2.1.1.b0
```

You may wish to add this to your ~/.bashrc file so that it loads automatically on login. These statements tell the supercomputer where the GrADS executables and supplementary data sets (e.g., maps, fonts) may be located. A working installation of GrADS v2.2.1 is also available by request, but its use is discouraged unless you have a specific reason to require it over v2.1.1.b0.

### **Post-Processing Using NCL**

It is also possible to post-process WRF-ARW data using utilities developed around the NCAR Graphics framework, specifically using NCL. A working version of NCL 6.4 is available on mortimer as a module:

```
module use /bigraid-02/LS/evans36/modules/  
module load ncl-6.4
```

You will also want to add the following entry to your ~/.bashrc file:

```
export NCARG_ROOT=/bigraid-05/LS/evans36/software/ncl
```

The primary benefit of using NCL rather than ARWpost for WRF-ARW output and visualization is that no post-processing step is necessary; the necessary libraries for reading and manipulating WRF data are built into NCL. The learning curve is somewhat steep, however, particularly if you do not have extensive experience with modern, object-based programming languages. I highly recommend reading the NCL User's Guide (<http://www.ncl.ucar.edu>) and working through a number of examples to become familiar with NCL before relying on it exclusively for WRF visualization. Also recommended are the WRF Tutorial slides on NCL, available at <http://www2.mmm.ucar.edu/wrf/users/tutorial/201801/ncl.pdf>.

### **Post-Processing Using the Unified Post-Processor (UPP) and GrADS**

WRF-ARW model output may also be post-processed using the Unified Post-Processor, or UPP, initially developed by NCEP for use with the GFS and NAM models. It produces output in GRIB format, which can then be read in (after some manipulation) by either GrADS or GEMPAK. The primary advantage to using UPP is the extensive number of derived quantities provided by UPP. The primary disadvantage of using UPP lies in its complexity, particularly with regards to compiling and executing the UPP code.

This guide focuses upon compiling and executing version 3.2 of the UPP code. It assumes that you have added all of the modules loaded to compile and run WRF to your ~/.bashrc file. If you have not done so, please run each module load statement first before continuing.

In the directory in which your WRFV3 and WPS directories reside, issue the following command

to download the UPP v3.2 source code:

```
wget https://dtcenter.org/upp/users/downloads/UPP_releases/DTC_upp_v3.2.tar.gz
```

Next, unzip and untar this file, (using `tar -zxvf`). After the file has been untarred, change into the UPPV3.2 directory and run `./configure`. Of the eight selections, you will choose option #4, "Linux x86\_64, Intel compiler (dm\_par)." Once this option has been given to the configure script, you are ready to compile. Open a slurm-shell and type `./compile`, again assuming that all of the necessary module files were loaded and environment variables set.

Compilation will take a few minutes. To ensure that the compilation completed successfully, check the output log (`UPPcompile.log`) for any errors. You also will want to check to ensure that a total of sixteen files are present within two directories:

```
UPPV3.2/lib/: libbacio.a, libCRTM.a, libg2.a, libg2tmpl.a,  
libgfsio.a, libip.a, libnemsio.a, ibsfcio.a, libsigio.a, libsp.a,  
libw3emc.a, libw3nco.a, libxmlparse.a
```

```
UPPV3.2/bin/: unipost.exe, ndate.exe, copygb.exe
```

Presuming that compilation completed successfully, you are ready to read and edit the UPP namelist file, `UPPV3.2/parm/wrf_cntrl.parm`. This file controls the fields that UPP will post-process and the vertical (isobaric) levels on which post-processed data will be available. The first three lines of `wrf_cntrl.parm` can safely be ignored. The remaining lines, two for each output field, control what will be processed and on what vertical levels it will be processed. The first line for each record looks like this:

```
(DWPT TEMP ON P SFCS ) SCAL=( 4.0)
```

The field controlled by that record is listed within the first set of parentheses. In this example, dewpoint (DWPT) temperature (TEMP) on pressure (P) surfaces (SFCS) is the output field. A full listing of output files is available on pages 18 through 33 of the [UPP User's Guide](#). The remainder of this line, the `SCAL=( 4.0)`, controls the accuracy of floating-point output. In this example, `SCAL=( 4.0)` states that the output data will be accurate to within 1/16 (i.e.,  $1/2^4$ ). Usually, it is fine to leave this number as-is; however, if you wish for there to be greater precision to a given field, set this value higher (e.g., 5.0 or 6.0).

The second line for each record looks like this:

```
L=(00000 01001 01010 10101 01010 10101 01010 11111 11111 10000  
00000 00000 00000 00000)
```

This line controls the vertical levels on which post-processed data will be available. A value of 0 indicates NO output at a given level; a value of 1 indicates that output WILL be provided at a given level. For surface-level output, only the first digit is of interest. For pressure-level output, the first 47 digits are of interest. In order, these levels are 2, 5, 7, 10, 20, 30, 50, 70, 75, 100, 125, 150, 175,

200, 225, 250, 275, 300, 325, 350, 375, 400, 425, 450, 475, 500, 525, 550, 575, 600, 625, 650, 675, 700, 725, 750, 775, 800, 825, 850, 875, 900, 925, 950, 975, and 1000 hPa. The isobaric levels on which you choose to output data, and the fields that you desire to output, will vary depending upon your specific application.

For model-level output, the first N digits are of interest, where N is the number of vertical levels used within the model simulation (i.e., `e_vert` in your `WRFV3/run/namelist.input` file). Other variables for specific applications (e.g., soil data, PBL averages, etc.) have slightly different specifications; please see pages 10 and 11 of the [UPP User's Guide](#) for more details.

Decide which model fields you desire to output, modify `wrf_cntrl.parm` appropriately, and then save the file. Once this is done, you are ready to post-process the data. To do so, move into the `UPPV3.2/scripts` directory and open the `run_unipost` script with your favorite text editor. A number of environment and script variables must be edited, including:

- `export TOP_DIR` (line 75) – the full directory path to your UPPV3.2 directory.
- `export DOMAINPATH` (line 76) – the full directory path you some working directory where, ideally, you want your post-processed data to reside. You'll want to make sure that this folder exists.
- `export WRFPATH` (line 77) – the full directory path to your WRF installation (the WRFV3 directory).
- `export modelDataPath` (line 81) – change this to the full path to your `wrfout_d01` file(s).
- `export startdate` (line 93) – change this to the start time of your WRF-ARW model forecast, in `YYYYMMDDHH` format.
- `export fhr` (line 94) – usually can keep this at 00 unless you wish to start post-processing at some time after model initialization.
- `export lastfhr` (line 95) – change this to the length (in hours) of your WRF-ARW model forecast or the last desired forecast output time.
- `export incrementthr` (line 96) – change this to match how frequently (in hours) you wish to have post-processed model output.

You will also want make sure that lines 107 and 108 are commented out (`#` in front) and that line 104 is uncommented (no `#` in front). We will run this job on one processor on a compute node, and have compiled it with distributed memory access. But, for the job submission to work correctly, we submit it like a serial job.

Once this is done, we are ready to post-process the data. First, create a job submission script (perhaps named `process.lsf`) in your `UPPV3.2/scripts/` directory that looks like this:

```
#!/bin/bash
#SBATCH -J process
#SBATCH -o process.log
#SBATCH -n 1
module use /sharedapps/ICC/modulefiles/
```

```
module load icc/18.0.1
module load openmpi/3.0.0_v2
module use /bigraid-02/LS/evans36/modules/
module load netcdf-4.6.1
module load grib2-libs-2018q2
mpirun ./real.exe
rm process.log
cd /path/to/Uppv3.2/scripts
mpirun ./run_unipost
```

Make sure that you modify the `cd` statement above with the appropriate directory path.

Despite whatever you may set for your `DOMAINPATH` variable above, output will likely appear in your `Uppv3.2/scripts/` directory as `WRFPRS_d01.##`, where `##` = forecast hour. If all goes well, you'll get output files for each forecast time requested. This may take a few minutes to an hour or more, particularly for lengthy, high-resolution model simulations. Afterward, you may safely move the `WRFPRS_d01*` files to the directory that you entered as the `DOMAINPATH` variable; in fact, I recommend doing in order to facilitate analysis of multiple cases.

Once this has completed, we need to create GrADS control files for these data. First, load the GrADS module (see the ARWpost section above). Next, issue one of the following commands from your data/working directory:

```
./grib2ctl.pl WRFPRS_d01.%f2 > (casename).ctl (if max fcst hour < 100)
./grib2ctl.pl WRFPRS_d01.%f3 > (casename).ctl (if max fcst hour > 100)
```

where you will want to replace `(casename)` with some descriptive name for the dataset. Once this has completed, the `(casename).ctl` file will contain a nearly-complete GrADS control file. Before reading it in, however, we need to edit one line within this file. Open it up with your favorite text editor, then go to the line that starts with `'ddef 1'`. Change the `1` to the number of forecast times (i.e., the number of `WRFPRS_d01*` files). Change the `'1mo'` at the end of this line to the time increment between files (e.g., `1hr`, `3hr`, etc.).

Next, we need to create a "map" for the output data file so that GrADS can read it in. To do so, issue the following command from this same directory:

```
gribmap -e -i (casename).ctl
```

where you replace `(casename)` with whatever you used as the prefix for your `.ctl` file. Once this has completed, you are ready to read in the data using GrADS.

### Other Post-Processing Systems

Other post-processing systems for WRF-ARW output do exist. VAPOR is a self-contained tool developed by UCAR for graphics-intensive multi-dimensional visualizations; in many ways, it is the natural successor to the deprecated Vis5d package. Information on VAPOR may be found in Chapter 9 of the WRF-ARW User's Guide. Newer utilities such as `wrf-python` exist to visualize

and process WRF-ARW output using Python are also available. WRF-ARW model output may also be used with a number of other utilities, such as the HYSPLIT trajectory and dispersion model and LAGRANTO trajectory tool. If you are interested in using WRF-ARW model output with such a utility, I encourage you to consult the documentation for that utility to understand what is necessary to be able to do so.

---

### *Conclusions*

If you have any questions with WRF model installation, setup, or debugging, please feel free to ask me via e-mail (UWM affiliates only; otherwise, please contact [wrfhelp@ucar.edu](mailto:wrfhelp@ucar.edu)).