

Chapter 6: WRF Data Assimilation (WRFDA)

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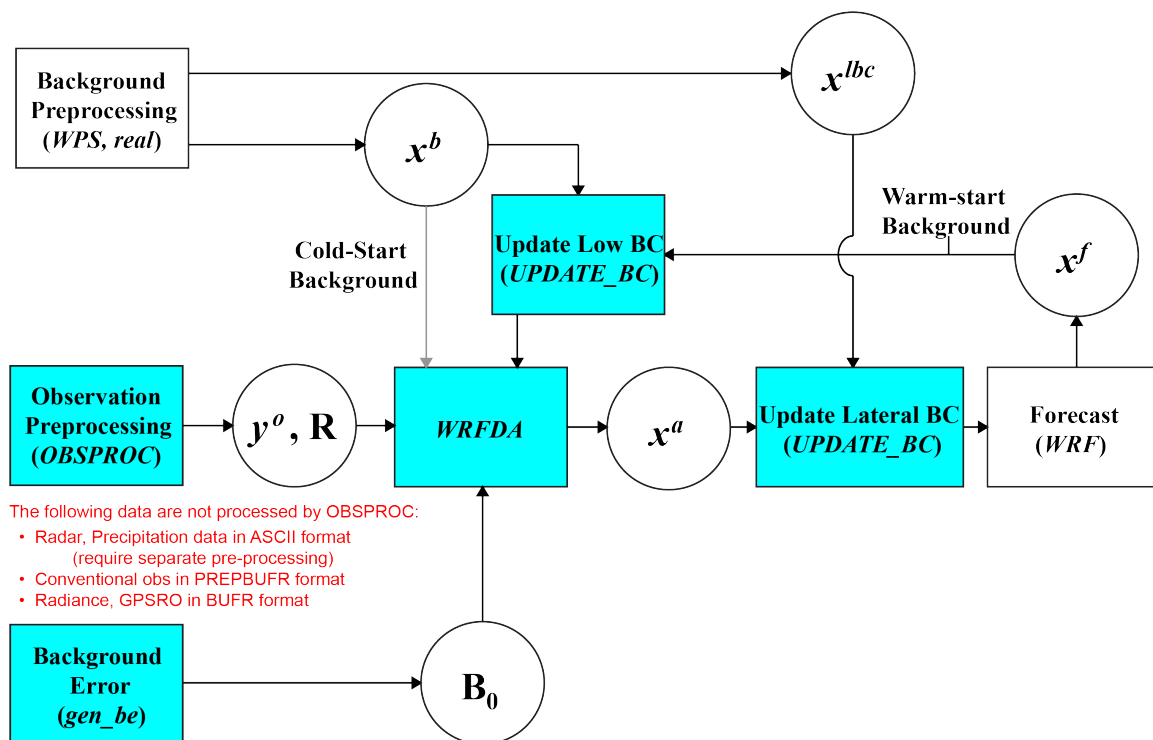
Introduction

Data assimilation is the technique by which **observations** are combined with an NWP product (the **first guess** or background forecast) and their respective error statistics to provide an improved estimate (the **analysis**) of the atmospheric (or oceanic, Jovian, etc.) state. Variational (Var) data assimilation achieves this through the iterative minimization of a prescribed cost (or penalty) function. Differences between the analysis and observations/first guess are penalized (damped) according to their perceived error. The difference between three-dimensional (3D-Var) and four-dimensional (4D-Var) data assimilation is the use of a numerical forecast model in the latter.

The MMM Laboratory of NCAR supports a unified (global/regional, multi-model, 3/4D-Var) model-space data assimilation system (WRFDA) for use by the NCAR staff and collaborators, and is also freely available to the general community, together with further documentation, test results, plans etc., from the WRFDA web-page (<http://www2.mmm.ucar.edu/wrf/users/wrfda/index.html>).

Various components of the WRFDA system are shown in blue in the sketch below, together with their relationship with the rest of the WRF system.

WRFDA in the WRF Modeling System



x^b	first guess, either from a previous WRF forecast or from WPS/real.exe output.
x^{lbc}	lateral boundary from WPS/real.exe output.
x^a	analysis from the WRFDA data assimilation system.
x^f	WRF forecast output.
y^o	observations processed by OBSPROC. (note: PREPBUFR input, radar, radiance, and rainfall data do not go through OBSPROC)
B_0	background error statistics from generic BE data (CV3) or gen_be.
R	observational and representative error statistics.

In this chapter, you will learn how to install and run the various components of the WRFDA system. For training purposes, you are supplied with a test case, including the following input data:

- observation files,
- a netCDF background file (WPS/real.exe output, the first guess of the analysis)
- background error statistics (estimate of errors in the background file).
- This tutorial dataset can be downloaded from the WRFDA Users Page (<http://www2.mmm.ucar.edu/wrf/users/wrfda/download/testdata.html>), and will be described later in more detail. In your own work, however, you will have to create all these input files yourself. See the section “[Running Observation Preprocessor](#)” for creating your observation files. See the section “[Background Error and running GEN_BE](#)” for generating your background error statistics file, if you want to use cv_options=5, 6, or 7.

Before using your own data, we suggest you start by running through the WRFDA- related programs using the supplied test case. This serves two purposes: First, you can learn how to run the programs with data we have tested ourselves, and second you can test whether your computer is capable of running the entire data assimilation system. After you have completed the tutorial, you can try running other, more computationally-intensive case studies, and experimenting with some of the many namelist variables.

WARNING: It is impossible to test every permutation of computer, compiler, number of processors, case, namelist option, etc. for every WRFDA release. The namelist options that are supported are indicated in the “WRFDA/var/README.namelist”, and these are the default options.

Hopefully, our test cases will prepare you for the variety of ways in which you may wish to run your own WRFDA experiments. Please inform us about your experiences through the [WRF and MPAS-A Support Forum](https://forum.mmm.ucar.edu) (<https://forum.mmm.ucar.edu>).

As a professional courtesy, we request that you include the following references in any publication that uses any component of the community WRFDA system:

Barker, D.M., W. Huang, Y.R. Guo, and Q.N. Xiao., 2004: A Three-Dimensional (3DVAR) Data Assimilation System For Use With MM5: Implementation and Initial Results. *Mon. Wea. Rev.*, **132**, 897-914.

Huang, X.Y., Q. Xiao, D.M. Barker, X. Zhang, J. Michalakes, W. Huang, T. Henderson, J. Bray, Y. Chen, Z. Ma, J. Dudhia, Y. Guo, X. Zhang, D.J. Won, H.C. Lin, and Y.H. Kuo, 2009: Four-Dimensional Variational Data Assimilation for WRF: Formulation and Preliminary Results. *Mon. Wea. Rev.*, 137, 299–314.

Barker, D., X.-Y. Huang, Z. Liu, T. Auligné, X. Zhang, S. Rugg, R. Ajjaji, A. Bourgeois, J. Bray, Y. Chen, M. Demirtas, Y.-R. Guo, T. Henderson, W. Huang, H.-C. Lin, J. Michalakes, S. Rizvi, and X. Zhang, 2012: The Weather Research and Forecasting Model's Community Variational/Ensemble Data Assimilation System: WRFDA. *Bull. Amer. Meteor. Soc.*, 93, 831–843.

Liu, Z., J. Ban, J.-S. Hong, and Y.-H. Kuo, 2020: Multi-resolution incremental 4D-Var for WRF: Implementation and application at convective scale, *Q. J. R. Meteorol. Soc.*, 146, 3661-3674.

Sun, W., Liu, Z., Chen, D., Zhao, P., and Chen, M., 2020: Development and application of the WRFDA-Chem three-dimensional variational (3DVAR) system: aiming to improve air quality forecasting and diagnose model deficiencies, *Atmos. Chem. Phys.*, 20, 9311-9329.

Running WRFDA requires a Fortran 90 compiler. The WRFDA system can be compiled on the following platforms: Linux (ifort, gfortran, pgf90), Macintosh (gfortran, ifort), IBM (xlf), and SGI Altix (ifort). Please let us know if this does not meet your requirements, and we will attempt to add other machines to our list of supported architectures, as resources allow. Although we are interested in hearing about your experiences in modifying compiler options, we do not recommend making changes to the configure file used to compile WRFDA.

Installing WRFDA for 3DVAR Run

a. Obtaining WRFDA Source Code

Users can download the WRFDA source code from http://www2.mmm.ucar.edu/wrf/users/wrfda/download/get_source.html.

Note: The source code for WRFDA and WRFPLUS are contained in the Basic WRF tar file. Although the WRF and WRFDA source code is packaged together, they can not be built together. To build basic WRF, one would need to download the tar file again and build it separately (in a separate directory). For instructions to build WRFDA versions prior to V4.0, please see: http://www2.mmm.ucar.edu/wrf/users/docs/user_guide_V3.9/users_guide_chap6.htm#_Installing_WRFDA_for_1.

After the tar file is unzipped (`gunzip WRFV4.1.TAR.gz`) and untarred (`tar -xvf WRFV4.1.TAR`), the directory WRF should be created. This directory contains the WRFDA

source, the WRFPLUS source (now fully integrated into WRF and TLM/ADJ code, located under the ‘wrftladj’ directory), external libraries, and fixed files. The following is a list of the system components and content for each subdirectory:

Directory Name	Content
var/da	WRFDA source code
var/run	Fixed input files required by WRFDA, such as background error covariance, radiance-related files, CRTM coefficients and VARBC.in
var/external	Libraries needed by WRFDA, includes CRTM, BUFR, LAPACK, BLAS
var/obsproc	OBSPROC source code, namelist, and observation error files
var/gen_be	Source code of gen_be, the utility to create background error statistics files
var/build	Builds all .exe files.

b. Compile WRFDA and Libraries

Some external libraries (e.g., LAPACK, BLAS, and NCEP BUFR) are included in the WRF tar file. To compile the WRFDA code, the only mandatory library is the netCDF library. You should set an environment variable NETCDF to point to the directory where your netCDF library is installed

```
> setenv NETCDF path-to-your-netcdf-library/netcdf
```

The source code for BUFRLIB 10.2.3 (with minor modifications) is included in the WRFDA tar file, and is compiled automatically. This library is used for assimilating files in PREPBUFR and NCEP BUFR format.

AMSR2 data can be assimilated in HDF5 format, which requires the use of HDF5 libraries. If you wish to make use of this capability, you must ensure that HDF5 libraries are installed on your system (or download and install them yourself; the source code is available from <https://www.hdfgroup.org/HDF5/>). To use HDF5 in WRFDA, you should set the environment variable “HDF5” to the parent path of your HDF5 build:

```
> setenv HDF5 your_hdf5_path
```

The HDF5 path should contain the directories “include” and “lib”.

For some platforms, you may have to also add the HDF5 “lib” directory to your environ-

ment variable LD_LIBRARY_PATH:

```
> setenv LD_LIBRARY_PATH ${LD_LIBRARY_PATH}:your_hdf5_path/lib
```

If satellite radiance data are to be used, a Radiative Transfer Model (RTM) is required. The current RTM versions that WRFDA supports are CRTM V2.3.0 and RTTOV V12.1.

The CRTM V2.3.0 source code is included in the WRFDA tar file, and is compiled automatically. No action is needed from the user. CRTM Coefficients can be downloaded here: http://www2.mmm.ucar.edu/wrf/users/wrfda/download/crtm_coeffs.html.

If you wish to use RTTOV, download and install the RTTOV v12 library before compiling WRFDA. This library can be downloaded from <https://nwp-saf.eumetsat.int/site/software/rttov/rttov-v12/>. **The RTTOV libraries must be compiled with the “emis_atlas” option in order to work with WRFDA; see the RTTOV “re-adme.txt” for instructions on how to do this.** After compiling RTTOV (see the RTTOV documentation for detailed instructions), set the “RTTOV” environment variable to the path where the `lib` directory resides. For example, if the library files can be found in `/usr/local/rttov11/gfortran/lib/librttov11.*.a`, you should set RTTOV as:

```
> setenv RTTOV /usr/local/rttov11/gfortran
```

Note: Make sure the required libraries were all compiled using the same compiler that will be used to build WRFDA, since the libraries produced by one compiler may not be compatible with code compiled with another.

Assuming all required libraries are available and the WRFDA source code is ready, you can start to build WRFDA using the following steps:

For organization and consistency, rename the WRF directory:

```
> mv WRF WRFDA
```

Enter the WRFDA directory and run the configure script:

```
> cd WRFDA
> ./configure wrfda
```

A list of configuration options should appear. Each option combines an operating system, a compiler type, and a parallelism option. Since the configuration script doesn’t check which compilers are *actually* installed on your system, be sure to select only among the options that you have available to you. The available parallelism options are single-processor (serial), shared-memory parallel (smpar), distributed-memory parallel (dmpar), and distributed-memory with shared-memory parallel (sm+dm). **However, shared-memory (smpar and sm+dm) options are not supported as of WRFDA Version 4.1, so we do not recommend selecting any of these options.**

For example, on a Linux machine such as NCAR’s Cheyenne, the above steps will look similar to the following:

```
checking for perl5... no
checking for perl... found /usr/bin/perl (perl)
Will use NETCDF in dir: /glade/apps/opt/netcdf/4.3.0/gnu/4.8.2/
Will use HDF5 in dir: /glade/u/apps/opt/hdf5/1.8.12/gnu/4.8.2/
PHDF5 not set in environment. Will configure WRF for use without.
Will use 'time' to report timing information
$JASPERLIB or $JASPERINC not found in environment, configuring to build without grib2
I/O...
-----
Please select from among the following Linux x86_64 options:

    1. (serial)   2. (smpar)   3. (dmpar)   4. (dm+sm)   PGI (pgf90/gcc)
    5. (serial)   6. (smpar)   7. (dmpar)   8. (dm+sm)   PGI (pgf90/pgcc): SGI MPT
    9. (serial)  10. (smpar)  11. (dmpar)  12. (dm+sm)   PGI (pgf90/gcc): PGI accelerator
   13. (serial)  14. (smpar)  15. (dmpar)  16. (dm+sm)   INTEL (ifort/icc)
                                           17. (dm+sm)   INTEL (ifort/icc): Xeon Phi (MIC
architecture)
   18. (serial)  19. (smpar)  20. (dmpar)  21. (dm+sm)   INTEL (ifort/icc): Xeon (SNB with
AVX mods)
   22. (serial)  23. (smpar)  24. (dmpar)  25. (dm+sm)   INTEL (ifort/icc): SGI MPT
   26. (serial)  27. (smpar)  28. (dmpar)  29. (dm+sm)   INTEL (ifort/icc): IBM POE
   30. (serial)                               31. (dmpar)   PATHSCALE (pathf90/pathcc)
   32. (serial)  33. (smpar)  34. (dmpar)  35. (dm+sm)   GNU (gfortran/gcc)
   36. (serial)  37. (smpar)  38. (dmpar)  39. (dm+sm)   IBM (xlf90_r/cc_r)
   40. (serial)  41. (smpar)  42. (dmpar)  43. (dm+sm)   PGI (ftn/gcc): Cray XC CLE
   44. (serial)  45. (smpar)  46. (dmpar)  47. (dm+sm)   CRAY CCE (ftn/cc): Cray XE and XC
   48. (serial)  49. (smpar)  50. (dmpar)  51. (dm+sm)   INTEL (ftn/icc): Cray XC
   52. (serial)  53. (smpar)  54. (dmpar)  55. (dm+sm)   PGI (pgf90/pgcc)
   56. (serial)  57. (smpar)  58. (dmpar)  59. (dm+sm)   PGI (pgf90/gcc): -f90=pgf90
   60. (serial)  61. (smpar)  62. (dmpar)  63. (dm+sm)   PGI (pgf90/pgcc): -f90=pgf90
   64. (serial)  65. (smpar)  66. (dmpar)  67. (dm+sm)   INTEL (ifort/icc): HSW/BDW
   68. (serial)  69. (smpar)  70. (dmpar)  71. (dm+sm)   INTEL (ifort/icc): KNL MIC

Enter selection [1-71] : 34
-----
Configuration successful!
-----
... ..
```

After entering the option that corresponds to your machine/compiler combination, the configure script should print the message “Configuration successful!” followed by a large amount of configuration information. Depending on your system, you may see a warning message mentioning that some Fortran 2003 features have been removed: this message is normal and can be ignored. However, if you see a message “One of compilers testing failed! Please check your compiler”, configuration has probably failed, and you should make sure you have selected the correct option.

After running the configuration script and choosing a compilation option, a `configure.wrf` file will be created. Because of the variety of ways that a computer can be configured, if the WRFDA build ultimately fails, there is a chance that minor modifications to the `configure.wrf` file may be needed.

To compile WRFDA, type

```
> ./compile all_wrfvar >& compile.out
```


Successful compilation will produce 44 executables: 43 of which are in the `var/build` directory and linked in the `var/da` directory, with the 44th, `obsproc.exe`, found in the `var/obsproc/src` directory. You can list these executables by issuing the command:

```
>ls -l var/build/*exe var/obsproc/src/obsproc.exe
-rwxr-xr-x 1 user 885143 Apr 4 17:22 var/build/da_advance_time.exe
-rwxr-xr-x 1 user 1162003 Apr 4 17:24 var/build/da_bias_airmass.exe
-rwxr-xr-x 1 user 1143027 Apr 4 17:23 var/build/da_bias_scan.exe
-rwxr-xr-x 1 user 1116933 Apr 4 17:23 var/build/da_bias_sele.exe
-rwxr-xr-x 1 user 1126173 Apr 4 17:23 var/build/da_bias_verif.exe
-rwxr-xr-x 1 user 1407973 Apr 4 17:23 var/build/da_rad_diags.exe
-rwxr-xr-x 1 user 1249431 Apr 4 17:22 var/build/da_tune_obs_desroziers.exe
-rwxr-xr-x 1 user 1186368 Apr 4 17:24 var/build/da_tune_obs_hollingsworth1.exe
-rwxr-xr-x 1 user 1083862 Apr 4 17:24 var/build/da_tune_obs_hollingsworth2.exe
-rwxr-xr-x 1 user 1193390 Apr 4 17:24 var/build/da_update_bc_ad.exe
-rwxr-xr-x 1 user 1245842 Apr 4 17:23 var/build/da_update_bc.exe
-rwxr-xr-x 1 user 1492394 Apr 4 17:24 var/build/da_verif_grid.exe
-rwxr-xr-x 1 user 1327002 Apr 4 17:24 var/build/da_verif_obs.exe
-rwxr-xr-x 1 user 26031927 Apr 4 17:31 var/build/da_wrfvar.exe
-rwxr-xr-x 1 user 1933571 Apr 4 17:23 var/build/gen_be_addmean.exe
-rwxr-xr-x 1 user 1944047 Apr 4 17:24 var/build/gen_be_cov2d3d_contrib.exe
-rwxr-xr-x 1 user 1927988 Apr 4 17:24 var/build/gen_be_cov2d.exe
-rwxr-xr-x 1 user 1945213 Apr 4 17:24 var/build/gen_be_cov3d2d_contrib.exe
-rwxr-xr-x 1 user 1941439 Apr 4 17:24 var/build/gen_be_cov3d3d_bin3d_contrib.exe
-rwxr-xr-x 1 user 1947331 Apr 4 17:24 var/build/gen_be_cov3d3d_contrib.exe
-rwxr-xr-x 1 user 1931820 Apr 4 17:24 var/build/gen_be_cov3d.exe
-rwxr-xr-x 1 user 1915177 Apr 4 17:24 var/build/gen_be_diags.exe
-rwxr-xr-x 1 user 1947942 Apr 4 17:24 var/build/gen_be_diags_read.exe
-rwxr-xr-x 1 user 1930465 Apr 4 17:24 var/build/gen_be_ensmean.exe
-rwxr-xr-x 1 user 1951511 Apr 4 17:24 var/build/gen_be_ensrf.exe
-rwxr-xr-x 1 user 1994167 Apr 4 17:24 var/build/gen_be_epl.exe
-rwxr-xr-x 1 user 1996438 Apr 4 17:24 var/build/gen_be_ep2.exe
-rwxr-xr-x 1 user 2001400 Apr 4 17:24 var/build/gen_be_etkf.exe
-rwxr-xr-x 1 user 1942988 Apr 4 17:24 var/build/gen_be_hist.exe
-rwxr-xr-x 1 user 2021659 Apr 4 17:24 var/build/gen_be_stage0_gsi.exe
-rwxr-xr-x 1 user 2012035 Apr 4 17:24 var/build/gen_be_stage0_wrf.exe
-rwxr-xr-x 1 user 1973193 Apr 4 17:24 var/build/gen_be_stage1_ldvar.exe
-rwxr-xr-x 1 user 1956835 Apr 4 17:24 var/build/gen_be_stage1.exe
-rwxr-xr-x 1 user 1963314 Apr 4 17:24 var/build/gen_be_stage1_gsi.exe
-rwxr-xr-x 1 user 1975042 Apr 4 17:24 var/build/gen_be_stage2_ldvar.exe
-rwxr-xr-x 1 user 1938468 Apr 4 17:24 var/build/gen_be_stage2a.exe
-rwxr-xr-x 1 user 1952538 Apr 4 17:24 var/build/gen_be_stage2.exe
-rwxr-xr-x 1 user 1202392 Apr 4 17:22 var/build/gen_be_stage2_gsi.exe
-rwxr-xr-x 1 user 1947836 Apr 4 17:24 var/build/gen_be_stage3.exe
-rwxr-xr-x 1 user 1928353 Apr 4 17:24 var/build/gen_be_stage4_global.exe
-rwxr-xr-x 1 user 1955622 Apr 4 17:24 var/build/gen_be_stage4_regional.exe
-rwxr-xr-x 1 user 1924416 Apr 4 17:24 var/build/gen_be_vertloc.exe
-rwxr-xr-x 1 user 2057673 Apr 4 17:24 var/build/gen_mbe_stage2.exe
-rwxr-xr-x 1 user 2110993 Apr 4 17:32 var/obsproc/src/obsproc.exe
```

The main executable for running WRFDA is `da_wrfvar.exe`. Make sure it has been created after the compilation: it is fairly common that all the executables will be successfully compiled except this main executable. If this occurs, please check the compilation log file carefully for any errors.

The basic `gen_be` utility for the regional model consists of `gen_be_stage0_wrf.exe`, `gen_be_stage1.exe`, `gen_be_stage2.exe`, `gen_be_stage2a.exe`, `gen_be_stage3.exe`, `gen_be_stage4_regional.exe`, and `gen_be_diags.exe`.

`da_update_bc.exe` is used for updating the WRF lower and lateral boundary conditions before and after a new WRFDA analysis is generated. This is detailed in the section on [Updating WRF Boundary Conditions](#).

`da_advance_time.exe` is a very handy and useful tool for date/time manipulation. Type `$WRFDA_DIR/var/build/da_advance_time.exe` to see its usage instructions.

`obsproc.exe` is the executable for preparing conventional observations for assimilation by WRFDA. Its use is detailed in the section on [Running Observation Preprocessor](#).

If you plan on using CRTM for radiance assimilation, check `$WRFDA_DIR/var/external/crtm_2.3.0/libsrc` to ensure that `libCRTM.a` was generated.

c. Clean old compilation

To remove all object files and executables, type:

```
./clean
```

To remove all build files, including `configure.wrf`, type:

```
./clean -a
```

The `clean -a` command is recommended if your compilation fails, or if the configuration file has been changed and you wish to restore the default settings.

Installing WRFPLUS and WRFDA for 4DVAR Run

If you intend to run WRFDA 4DVAR, it is necessary to have WRFPLUS installed first. WRFPLUS contains the adjoint and tangent linear models based on a simplified WRF model, which includes a few simplified physics packages, such as surface drag, large scale condensation and precipitation, and cumulus parameterization.

Note: if you intend to run both 3DVAR and 4DVAR experiments, it is not necessary to compile the code twice. The `da_wrfvar.exe` executable compiled for 4DVAR can be used for both 3DVAR and 4DVAR assimilation.

To install WRFPLUS (for versions prior to V4.0, see:

http://www2.mmm.ucar.edu/wrf/users/docs/user_guide_V3.9/users_guide_chap6.htm#_Installing_WRFPLUS_and_2)

- The WRFPLUS code is fully integrated into WRF and TLM/ADJ code, located under the ‘`wrftladj`’ directory. To obtain the WRF packaged file, see: http://www2.mmm.ucar.edu/wrf/users/download/get_source.html.
- Unzip and untar the WRF file, rename it appropriately to WRFPLUS, and then

run the `configure` script

```
> gunzip WRFV4.1.tar.gz
> tar -xf WRFV4.1.tar
> mv WRF WRFPLUS
> cd WRFPLUS
> ./configure wrfplus
```

As with 3D-Var, “serial” means single-processor, and “dmpar” means Distributed Memory Parallel (MPI). Be sure to select the same option for WRFPLUS as you will use for WRFDA.

- Compile WRFPLUS

```
> ./compile wrfplus >& compile.out
> ls -ls main/*.exe
```

If compilation was successful, you should see the WRFPLUS executable (named `wrfplus.exe`):

```
53292 -rwxr-xr-x 1 user man 54513254 Apr  6 22:43 main/wrfplus.exe
```

Finally, set the environment variable `WRFPLUS_DIR` to the appropriate directory:

```
>setenv WRFPLUS_DIR ${your_source_code_dir}/WRFPLUS
```

To install WRFDA for the 4D-Var run:

- If you intend to use RTTOV to assimilate radiance data, you will need to set the appropriate environment variable at compile time. See [the previous 3DVAR section](#) for instructions.

```
>./configure 4dvar

>./compile all_wrfvar >& compile.out

>ls -ls var/build/*.exe var/obsproc/*.exe
```

You should see the same 44 executables as are listed in [the above 3DVAR section](#), including `da_wrfvar.exe`

Running Observation Preprocessor (OBSPROC)

The OBSPROC program reads observations in LITTLE_R format (a text-based format, in use since the MM5 era). We have provided observations for the tutorial case, but for your

own applications, you will have to prepare your own observation files. Please see http://www2.mmm.ucar.edu/wrf/users/wrfda/download/free_data.html for the sources of some freely-available observations. Because the raw observation data files have many possible formats, such as ASCII, BUFR, PREPBUFR, MADIS (*note*: a converter for MADIS data to LITTLE_R is available on the WRFDA website:

<http://www2.mmm.ucar.edu/wrf/users/wrfda/download/madis.html>), and HDF, the free data site also contains instructions for converting the observations to LITTLE_R format. To make the WRFDA system as general as possible, the LITTLE_R format was adopted as an intermediate observation data format for the WRFDA system, however, *the conversion of the user-specific source data to LITTLE_R format is the user's task*. A more complete description of the LITTLE_R format, as well as conventional observation data sources for WRFDA, can be found by reading [The LITTLE_R for WRFDA help page](#), the “Observation Pre-processing” tutorial found at http://www2.mmm.ucar.edu/wrf/users/wrfda/Tutorials/2016_Aug/tutorial_presentations_summer_2016.html, or by referencing [Chapter 7 of this User's Guide](#).

The purpose of OBSPROC is to:

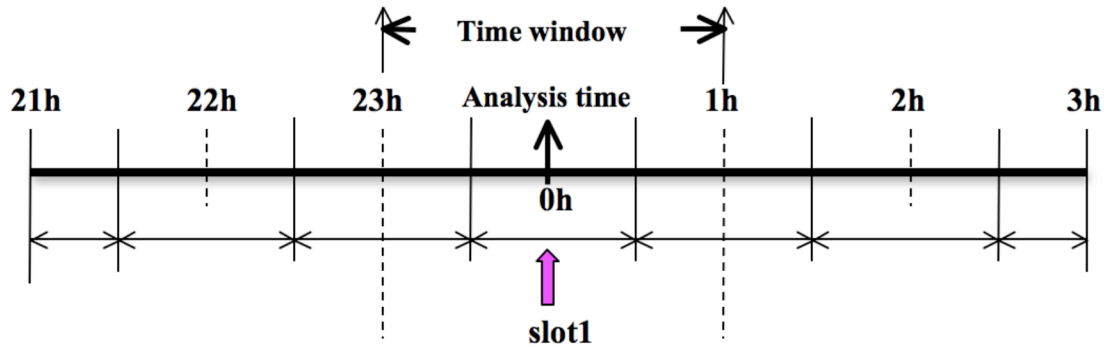
- Remove observations outside the specified temporal and spatial domains
- Re-order and merge duplicate (in time and location) data reports
- Retrieve pressure or height based on observed information using the hydrostatic assumption
- Check multi-level observations for vertical consistency and superadiabatic conditions
- Assign observation errors based on a pre-specified error file
- Write out the observation file to be used by WRFDA in ASCII or BUFR format

The OBSPROC program (`obsproc.exe`) should be found under the directory `$WRFDA_DIR/var/obsproc/src` if “`compile all_wrfvar`” completed successfully.

If you haven't already, you should download the tutorial case, which contains example files for all the exercises in this User's Guide. The case can be found at the WRFDA website (<http://www2.mmm.ucar.edu/wrf/users/wrfda/download/testdata.html>).

a. OBSPROC for 3DVAR

As an example, to prepare the observation file at the analysis time, all the observations in the range ± 1 h will be processed, which means that (in the example case) the observations between 23h and 1h are treated as the observations at 0h. This is illustrated in the following figure:



OBSPROC requires at least 3 files to run successfully:

- A namelist file (`namelist.obsproc`)
- An observation error file (`obserr.txt`)
- One or more observation files
- Optionally, a table for specifying the elevation information for marine observations over the US Great Lakes (`msfc.tbl`)

The files `obserr.txt` and `msfc.tbl` are included in the source code under `var/obsproc`. To create the required namelist file, we have provided an example file (`namelist_obsproc.3dvar.wrfvar-tut`) in the `var/obsproc` directory. Thus, proceed as follows.

```
> cd $WRFDA_DIR/var/obsproc
> cp namelist_obsproc.3dvar.wrfvar-tut namelist.obsproc
```

Next, edit the namelist file, `namelist.obsproc`, to accommodate your experiments. You will likely only need to change variables listed under records 1, 2, 6, 7, and 8. See `$WRFDA_DIR/var/obsproc/README.namelist`, or the section [OBSPROC namelist variables](#) for details. You should pay special attention to the record 7 and record 8 variables: these will determine the domain for which observations will be written to the output observation file. Alternatively, if you do not wish to filter the observations spatially, you can set `domain_check_h = .false.` under `&record4`.

If you are running the tutorial case, you should copy or link the sample observation file (`ob/2008020512/obs.2008020512`) to the `obsproc` directory. Alternatively, you can edit the namelist variable `obs_gts_filename` to point to the observation file's full path.

To run OBSPROC, type

```
> ./obsproc.exe >& obsproc.out
```

Once `obsproc.exe` has completed successfully, you will see an observation data file,

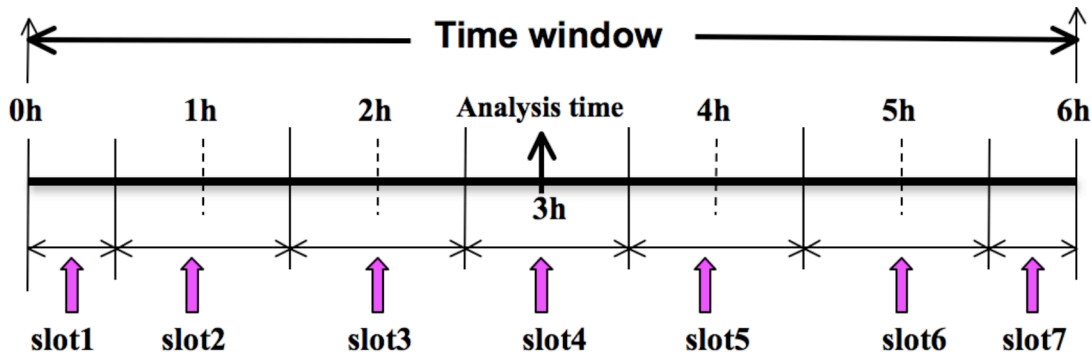
with the name formatted `obs_gts_YYYY-MM-DD_HH:NN:SS.3DVAR`, in the `obsproc` directory. For the tutorial case, this will be `obs_gts_2008-02-05_12:00:00.3DVAR`. This is the input observation file to WRFDA. It is an ASCII file that contains a header section (listed below) followed by observations. The meanings and format of observations in the file are described in the last six lines of the header section.

```
TOTAL = 9066, MISS. = -888888.,
SYNOP = 757, METAR = 2416, SHIP = 145, BUOY = 250, BOGUS = 0, TEMP =
86,
AMDAR = 19, AIREP = 205, TAMDAR = 0, PILOT = 85, SATEM = 106, SATOB =
2556,
GPSPW = 187, GPSZD = 0, GPSRF = 3, GPSEP = 0, SSMT1 = 0, SSMT2 =
0,
TOVS = 0, QSCAT = 2190, PROFL = 61, AIRSR = 0, OTHER = 0,
PHIC = 40.00, XLONC = -95.00, TRUE1 = 30.00, TRUE2 = 60.00, XIM11 = 1.00, XJM11 =
1.00,
base_temp= 290.00, base_lapse= 50.00, PTOP = 1000., base_pres=100000.,
base_tropo_pres= 20000., base_strat_temp= 215.,
IXC = 60, JXC = 90, IPROJ = 1, IDD = 1, MAXNES= 1,
NESTIX= 60,
NESTJX= 90,
NUMC = 1,
DIS = 60.00,
NESTI = 1,
NESTJ = 1,
INFO = PLATFORM, DATE, NAME, LEVELS, LATITUDE, LONGITUDE, ELEVATION, ID.
SRFC = SLP, PW (DATA,QC,ERROR).
EACH = PRES, SPEED, DIR, HEIGHT, TEMP, DEW PT, HUMID (DATA,QC,ERROR)*LEVELS.
INFO_FMT = (A12,1X,A19,1X,A40,1X,I6,3(F12.3,11X),6X,A40)
SRFC_FMT = (F12.3,I4,F7.2,F12.3,I4,F7.3)
EACH_FMT = (3(F12.3,I4,F7.2),11X,3(F12.3,I4,F7.2),11X,3(F12.3,I4,F7.2))
#-----#
..... observations .....
```

Before running WRFDA, you may find it useful to learn more about various types of data that will be processed (e.g., their geographical distribution). This file is in ASCII format and so you can easily view it. For a graphical view of the file's content, there are NCL scripts available which can display the distribution and type of observations. In the WRFDA Tools package (can be downloaded at <http://www2.mmm.ucar.edu/wrf/users/wrfda/download/tools.html>), the relevant script is located at `$TOOLS_DIR/var/graphics/ncl/plot_ob_ascii_loc.ncl`. You will need to have NCL installed in your system to use this script; for more information on NCL, the NCAR Command Language, see <http://www.ncl.ucar.edu/>.

b. OBSPROC for 4DVAR

To prepare the observation file, for example, at the analysis time 0h for 4D-Var, all observations from 0h to 6h will be processed and grouped in 7 sub-windows (slot1 through slot7) as illustrated in the following figure:



NOTE: The “Analysis time” in the above figure is not the actual analysis time (0h). It indicates the `time_analysis` setting in the namelist file, which in this example is three hours later than the actual analysis time. The actual analysis time is still 0h.

An example namelist (`namelist_obsproc.4dvar.wrfvar-tut`) has already been provided in the `var/obsproc` directory. Thus, proceed as follows:

```
> cd $WRFDA_DIR/var/obsproc
> cp namelist_obsproc.4dvar.wrfvar-tut namelist_obsproc
```

In the namelist file, you need to change the following variables to accommodate your experiments. In this tutorial case, the actual analysis time is `2008-02-05_12:00:00`, **but in the namelist, `time_analysis` should be set to 3 hours later**. The different values of `time_analysis`, `num_slots_past`, and `time_slots_ahead` contribute to the actual times analyzed. For example, if you set `time_analysis = 2008-02-05_16:00:00`, and set the `num_slots_past = 4` and `time_slots_ahead=2`, the final results will be the same as before.

Edit all the domain settings according to your own experiment; a full list of namelist options and descriptions can be found in the section [Description of Namelist Variables](#). You should pay special attention to the record 7 and record 8 variables: these will determine the domain for which observations will be written to the output observation file. Alternatively, if you do not wish to filter the observations spatially, you can set `do_main_check_h = .false.` under `&record4`.

If you are running the tutorial case, you should copy or link the sample observation file (`ob/2008020512/obs.2008020512`) to the `obsproc` directory. Alternatively, you can edit the namelist variable `obs_gts_filename` to point to the observation file’s full path.

To run OBSPROC, type

```
> obsproc.exe >& obsproc.out
```

Once `obsproc.exe` has completed successfully, you will see 7 observation data files, which for the tutorial case are named

```

obs_gts_2008-02-05_12:00:00.4DVAR
obs_gts_2008-02-05_13:00:00.4DVAR
obs_gts_2008-02-05_14:00:00.4DVAR
obs_gts_2008-02-05_15:00:00.4DVAR
obs_gts_2008-02-05_16:00:00.4DVAR
obs_gts_2008-02-05_17:00:00.4DVAR
obs_gts_2008-02-05_18:00:00.4DVAR

```

They are the input observation files to WRF 4D-Var.

Running WRFDA

a. Download Test Data

The WRFDA system requires three input files to run:

- WRF *first guess* file, output from either WPS/real.exe (wrfinput) or a WRF forecast (wrfout)
- Observations (in ASCII format, PREPBUFR or BUFR for radiance)
- A background error statistics file (containing background error covariance)

The following table summarizes the above info:

<i>Input Data</i>	<i>Format</i>	<i>Created By</i>
First Guess	NETCDF	WRF Preprocessing System (WPS) and real.exe or WRF
Observations	ASCII (PREPBUFR also possible)	Observation Preprocessor (OBSPROC)
Background Error Statistics	Binary	WRFDA gen_be utility or generic CV3

In the test case, you will store data in a directory defined by the environment variable \$DAT_DIR. This directory can be in any location, and it should have read access. Type

```
> setenv DAT_DIR your_choice_of_dat_dir
```

Here, your_choice_of_dat_dir is the directory where the WRFDA input data is stored.

If you have not already done so, download the sample data for the tutorial case, valid at 12 UTC 5th February 2008, from

<http://www2.mmm.ucar.edu/wrf/users/wrfda/download/testdata.html>

Once you have downloaded the `WRFDAV4.0-testdata.tar.gz` file to `$DAT_DIR`, extract it by typing

```
> gunzip WRFDAV4.0-testdata.tar.gz
> tar -xvf WRFDAV4.0-testdata.tar
```

Now you should find the following four files under “`$DAT_DIR`”

```
ob/2008020512/ob.2008020512      # Observation data in “little_r” format
rc/2008020512/wrfinput_d01       # First guess file
rc/2008020512/wrfbdy_d01        # lateral boundary file
be/be.dat                       # Background error file
.....
```

At this point you should have three of the input files (first guess, observations from OB-SPROC, and background error statistics files in the directory `$DAT_DIR`) required to run WRFDA, and have successfully downloaded and compiled the WRFDA code. If this is correct, you are ready to run WRFDA.

b. Run 3DVAR test case

The data for the tutorial case is valid at 12 UTC 5 February 2008. The first guess comes from the NCEP FNL (Final) Operational Global Analysis data, passed through the WRF-WPS and `real.exe` programs.

To run WRF 3D-Var, first create and enter into a working directory (for example, `$WRFDA_DIR/workdir`), and set the environment variable `WORK_DIR` to this directory (e.g., `setenv WORK_DIR $WRFDA_DIR/workdir`). Then follow the steps below:

```
> cd $WORK_DIR
> cp $DAT_DIR/namelist.input.3dvar namelist.input
> ln -sf $WRFDA_DIR/run/LANDUSE.TBL .
> ln -sf $DAT_DIR/rc/2008020512/wrfinput_d01 ./fg
> ln -sf $DAT_DIR/ob/2008020512/obs_gts_2008-02-05_12:00:00.3DVAR
./ob.ascii (note the different name!)
> ln -sf $DAT_DIR/be/be.dat .
> ln -sf $WRFDA_DIR/var/da/da_wrfvar.exe .
```

Now edit the file `namelist.input`, which is a very basic namelist for the tutorial test case, and is shown below.

```
&wrfvar1
var4d=false,
print_detail_grad=false,
/
&wrfvar2
/
&wrfvar3
ob_format=2,
/
&wrfvar4
/
&wrfvar5
/
```

```
&wrfvar6
max_ext_its=1,
ntmax=50,
orthonorm_gradient=true,
/
&wrfvar7
cv_options=5,
/
&wrfvar8
/
&wrfvar9
/
&wrfvar10
test_transforms=false,
test_gradient=false,
/
&wrfvar11
/
&wrfvar12
/
&wrfvar13
/
&wrfvar14
/
&wrfvar15
/
&wrfvar16
/
&wrfvar17
/
&wrfvar18
analysis_date="2008-02-05_12:00:00.0000",
/
&wrfvar19
/
&wrfvar20
/
&wrfvar21
time_window_min="2008-02-05_11:00:00.0000",
/
&wrfvar22
time_window_max="2008-02-05_13:00:00.0000",
/
&time_control
start_year=2008,
start_month=02,
start_day=05,
start_hour=12,
end_year=2008,
end_month=02,
end_day=05,
end_hour=12,
/
&fdda
/
&domains
e_we=90,
e_sn=60,
e_vert=41,
dx=60000,
dy=60000,
/
&dft_control
/
&tc
/
&physics
mp_physics=3,
ra_lw_physics=1,
ra_sw_physics=1,
radt=60,
sf_sfclay_physics=1,
```

```

sf_surface_physics=1,
bl_pbl_physics=1,
cu_physics=1,
cudt=5,
num_soil_layers=5,
mp_zero_out=2,
co2tf=0,
/
&scm
/
&dynamics
/
&bdy_control
/
&grib2
/
&fire
/
&namelist_quilt
/
&perturbation
/

```

No edits should be needed if you are running the tutorial case without radiance data. If you plan to use the PREPBUFR-format data, change the `ob_format=1` in `&wrfvar3` in `namelist.input` and link the data as `ob.bufr`,

```
> ln -fs $DAT_DIR/ob/2008020512/gdas1.t12z.prepbufr.nr ob.bufr
```

Once you have changed any other necessary namelist variables, run WRFDA 3D-Var:

```
> da_wrfvar.exe >& wrfda.log
```

The file `wrfda.log` (or `rs1.out.0000`, if run in distributed-memory mode) contains important WRFDA runtime log information. Always check the log after a WRFDA run:

```

*** VARIATIONAL ANALYSIS ***
WRFDA V4.1

DYNAMICS OPTION: Eulerian Mass Coordinate
  alloc_space_field: domain      1 ,          499448408 bytes allocated
  hybrid_opt = 0

  Tile Strategy is not specified. Assuming 1D-Y
WRF TILE   1 IS      1 IE      89 JS      1 JE      59
WRF NUMBER OF TILES = 1
Domain mapping info:
map_proj = 1
cen_lat  = 0.400000E+02
cen_lon  = -0.950000E+02
truelat1 = 0.300000E+02
truelat2 = 0.600000E+02
start_lat = 0.207406E+02
start_lon = -0.119501E+03
pole_lat  = 0.900000E+02
dsm       = 0.600000E+02

Set up observations (ob)

Using ASCII format observation input

Observation summary
  ob time 1
    sound                86 global,      86 local

```

```

synop          750 global,    750 local
pilot          85 global,    85 local
satem         105 global,   105 local
geoamv       2499 global,  2499 local
airep        221 global,   221 local
gpspw        187 global,   187 local
gpsrf         3 global,     3 local
metar       2408 global,  2408 local
ships        140 global,   140 local
qscat       2126 global,  2126 local
profiler      61 global,    61 local
buoy        247 global,   247 local
sonde_sfc     86 global,    86 local

```

Set up background errors for regional application for cv_options = 5

Using the averaged regression coefficients for unbalanced part

```

cloud_cv_options = 0
WRFDA dry control variables are: psi, chi_u, t_u and ps_u
WRFDA Humidity control variable is rh

```

Vertical truncation for psi = 15(99.00%)

Vertical truncation for chi_u = 20(99.00%)

Vertical truncation for t_u = 29(99.00%)

Vertical truncation for rh = 22(99.00%)

```

Scaling: var, len, ds: 0.100000E+01 0.100000E+01 0.600000E+05
Scaling: var, len, ds: 0.100000E+01 0.100000E+01 0.600000E+05
Scaling: var, len, ds: 0.100000E+01 0.100000E+01 0.600000E+05
Scaling: var, len, ds: 0.100000E+01 0.100000E+01 0.600000E+05
Scaling: var, len, ds: 0.100000E+01 0.100000E+01 0.600000E+05

```

Calculate innovation vector(iv)

Minimize cost function using CG method

```

Starting outer iteration : 1
Starting cost function: 5.366048535905661D+04, Gradient= 8.134548965597619D+02
For this outer iteration gradient target is: 8.134548965597618D+00

```

```

-----
      Loop Iter Cost Function      Gradient      Step
minimize_cg 1 0 5.366048535905661D+04 8.134548965597619D+02 0.000000000000000D+00
minimize_cg 1 1 4.160895118661885D+04 4.087246011996226D+02 3.642548782952763D-02
minimize_cg 1 2 3.684015367850271D+04 3.371798989064126D+02 5.709227119079368D-02
minimize_cg 1 3 3.374263324064368D+04 2.579817263649239D+02 5.449050389717097D-02
minimize_cg 1 4 3.163760162325043D+04 1.959172950097825D+02 6.325731144809946D-02
minimize_cg 1 5 3.069560214937115D+04 1.426201418835412D+02 4.908345259881822D-02
minimize_cg 1 6 2.994553670826729D+04 1.016946487735333D+02 7.375091679036799D-02
minimize_cg 1 7 2.963703986213524D+04 8.614527420925806D+01 5.966017496500481D-02
minimize_cg 1 8 2.944840596364260D+04 5.586889886231801D+01 5.083780846639206D-02
minimize_cg 1 9 2.932974897428447D+04 4.810455663750825D+01 7.602966851781012D-02
minimize_cg 1 10 2.925909806811914D+04 3.154639503336832D+01 6.106260102681394D-02
minimize_cg 1 11 2.922595474924949D+04 2.511864257306522D+01 6.660801879172493D-02
minimize_cg 1 12 2.920702117122884D+04 1.953264766296704D+01 6.001645744010778D-02
minimize_cg 1 13 2.919536455912784D+04 1.421581370380062D+01 6.110547268188220D-02
minimize_cg 1 14 2.918931706353119D+04 1.000448731094664D+01 5.984971814842433D-02
minimize_cg 1 15 2.918568177329574D+04 7.457040724273403D+00 7.264059789587740D-02
-----

```

Inner iteration stopped after 15 iterations

Final: 15 iter, J= 2.918568177329574D+04, g= 7.457040724273385D+00

```

Diagnostics
  Final cost function J      =      29185.68

  Total number of obs.      =      37570
  Final value of J          =      29185.68177
  Final value of Jo         =      23086.99122
  Final value of Jd         =           0.00000
  Final value of Jb         =      6098.69056
  Final value of Jc         =           0.00000
  Final value of Je         =           0.00000
  Final value of Jp         =           0.00000
  Final value of Jl         =           0.00000
  Final J / total num_obs   =      0.77683
  Jb factor used(1)         =      1.00000
  Jb factor used(2)         =      1.00000
  Jb factor used(3)         =      1.00000
  Jb factor used(4)         =      1.00000
  Jb factor used(5)         =      1.00000
  Jb factor used            =      1.00000
  Je factor used            =      1.00000
  VarBC factor used        =     10.00000

```

```
*** WRF-Var completed successfully ***
```

The file `namelist.output.da` (which contains the complete namelist settings) will be generated after a successful run of `da_wrfvar.exe`. The settings appearing in `namelist.output.da`, but not specified in your `namelist.input`, are the default values from `$WRFDA_DIR/Registry/registry.var`.

After successful completion, `wrfvar_output` (the WRFDA analysis file, i.e. the new initial condition for WRF) should appear in the working directory along with a number of diagnostic files. Text files containing various diagnostics will be explained in the [WRFDA Diagnostics](#) section.

To understand the role of various important WRFDA options, try re-running WRFDA by changing different namelist options. Some examples are listed below:

1. Response of convergence criteria:

Run the tutorial case with

```

&wrfvar6
  eps = 0.0001,
/

```

You may wish to compare various diagnostics with an earlier run.

2. Response of outer loop on minimization:

Run the tutorial case with

```

&wrfvar6
  max_ext_its = 2,
/

```

With this setting, the “outer loop” for the minimization procedure will be activated. You may wish to compare various diagnostics with an earlier run.

Note that when running multiple outer loops with the CV3 background error option, you must specify the scaling factors which are called as1, as2, as3, as4, and as5. More details can be found in the section [“Modifying CV3 length scales and variance”](#).

3. Response of suppressing particular types of data in WRFDA:

The types of observations that WRFDA gets to use actually depend on what is included in the observation file and the WRFDA namelist settings. For example, if you have SYNOP data in the observation file, you can suppress its usage in WRFDA by setting `use_synopobs=false` in record `&wrfvar4` of `namelist.input`. It is OK if there are no SYNOP data in the observation file and `use_synopobs=true`.

Turning on and off certain types of observations is widely used for assessing the impact of observations on data assimilations.

Note: It is important to go through the default “use_*” settings in record `&wrfvar4` in `WRFDA/Registry/registry.var` to know what observations are activated in default.

For example, try making the WRFDA convergence criterion more stringent. This is achieved by reducing the value of “EPS” to e.g. 0.0001 by adding “EPS=0.0001” in the `namelist.input` record `&wrfvar6`. See the section [Additional Background Error options](#) for more namelist options.

c. Run 4DVAR test case

To run WRF 4D-Var, first create and enter a working directory, such as `$WRFDA_DIR/workdir`. Set the `WORK_DIR` environment variable (e.g. `setenv WORK_DIR $WRFDA_DIR/workdir`)

For the tutorial case, the analysis date is 2008020512 and the test data directories are:

```
> setenv DAT_DIR {directory where data is stored}
> ls -lr $DAT_DIR
ob/2008020512
ob/2008020513
ob/2008020514
ob/2008020515
ob/2008020516
ob/2008020517
ob/2008020518
rc/2008020512
be
```

Note: WRFDA 4D-Var is able to assimilate conventional observational data, satellite radiance BUFR data, and precipitation data. The input data format can be PREPBUFR format data or ASCII observation data, processed by OBSPROC.

Now follow the steps below:

1) Link the executable file

```
> cd $WORK_DIR
> ln -fs $WRFDA_DIR/var/da/da_wrfvar.exe .
```

2) Link the observational data, first guess, BE and LANDUSE.TBL, etc.

```
> ln -fs $DAT_DIR/ob/2008020512/ob01.ascii ob01.ascii
> ln -fs $DAT_DIR/ob/2008020513/ob02.ascii ob02.ascii
> ln -fs $DAT_DIR/ob/2008020514/ob03.ascii ob03.ascii
> ln -fs $DAT_DIR/ob/2008020515/ob04.ascii ob04.ascii
> ln -fs $DAT_DIR/ob/2008020516/ob05.ascii ob05.ascii
> ln -fs $DAT_DIR/ob/2008020517/ob06.ascii ob06.ascii
> ln -fs $DAT_DIR/ob/2008020518/ob07.ascii ob07.ascii

> ln -fs $DAT_DIR/rc/2008020512/wrfinput_d01 .
> ln -fs $DAT_DIR/rc/2008020512/wrfbdy_d01 .
> ln -fs wrfinput_d01 fg

> ln -fs $DAT_DIR/be/be.dat .
> ln -fs $WRFDA_DIR/run/LANDUSE.TBL .
> ln -fs $WRFDA_DIR/run/GENPARM.TBL .
> ln -fs $WRFDA_DIR/run/SOILPARM.TBL .
> ln -fs $WRFDA_DIR/run/VEGPARM.TBL .
> ln -fs $WRFDA_DIR/run/RRTM_DATA_DBL RRTM_DATA
```

3) Copy the sample namelist

```
> cp $DAT_DIR/namelist.input.4dvar namelist.input
```

4) Edit necessary namelist variables, link optional files

WRFDA 4D-Var has the capability to consider lateral boundary conditions as control variables as well during minimization. The namelist variable `var4d_lbc=true` turns on this capability. To enable this option, WRF 4D-Var needs not only the first guess at the beginning of the time window, but also the first guess at the end of the time window.

```
> ln -fs $DAT_DIR/rc/2008020518/wrfinput_d01 fg02
```

Please note: WRFDA beginners should not use this option until you have a good understanding of the 4D-Var lateral boundary conditions control. To disable this feature, make sure `var4d_lbc` in `namelist.input` is set to `false`.

If you use PREPBUFR format data, set `ob_format=1` in `&wrfvar3` in `namelist.input`. Because 12UTC PREPBUFR data only includes the data from 9UTC to 15UTC, for 4D-Var you should include 18UTC PREPBUFR data as well:

```
> ln -fs $DAT_DIR/ob/2008020512/gdas1.t12z.prepbuftr.nr ob01.bufr
> ln -fs $DAT_DIR/ob/2008020518/gdas1.t18z.prepbuftr.nr ob02.bufr
```

Edit `namelist.input` to match your experiment settings. The most important namelist variables related to 4D-Var are listed below. Please refer to `README.namelist` under the `$WRFDA_DIR/var` directory. A common mistake users make is in the time information

settings. The rules are: `analysis_date`, `time_window_min` and `start_xxx` in `&time_control` should always be equal to each other; `time_window_max` and `end_xxx` should always be equal to each other; and `run_hours` is the difference between `start_xxx` and `end_xxx`, which is the length of the 4D-Var time window.

```
&wrfvar1
var4d=true,
var4d_lbc=false,
var4d_bin=3600,
.....
/
.....
&wrfvar18
analysis_date="2008-02-05_12:00:00.0000",
/
.....
&wrfvar21
time_window_min="2008-02-05_12:00:00.0000",
/
.....
&wrfvar22
time_window_max="2008-02-05_18:00:00.0000",
/
.....
&time_control
run_hours=6,
start_year=2008,
start_month=02,
start_day=05,
start_hour=12,
end_year=2008,
end_month=02,
end_day=05,
end_hour=18,
interval_seconds=21600,
debug_level=0,
/
.....
```

5) Run WRF 4D-Var

```
> cd $WORK_DIR
> ./da_wrfvar.exe >& wrfda.log
```

4DVAR is much more computationally expensive than 3DVAR, so running may take a while; you can set `ntmax` to a lower value so that WRFDA uses fewer minimization steps. You can also MPI with multiple processors to speed up the execution:

```
> mpirun -np 4 ./da_wrfvar.exe >& wrfda.log &
```

The “`mpirun`” command may be different depending on your machine. The output logs will be found in files named `rs1.out.####` and `rs1.error.####` for MPI runs.

Please note: If you utilize the lateral boundary conditions option (`var4d_lbc=true`), in addition to the analysis at the beginning of the time window (`wrfvar_output`), the analysis at the end of the time window will also be generated as `ana02`, which will be used in subsequent updating of boundary conditions before the forecast.

Radiance Data Assimilation in WRFDA

This section gives a brief description for various aspects related to radiance assimilation in WRFDA. Each aspect is described mainly from the viewpoint of usage, rather than more technical and scientific details, which will appear in a separate technical report and scientific paper. Namelist parameters controlling different aspects of radiance assimilation will be detailed in the following sections. It should be noted that this section does not cover general aspects of the assimilation process with WRFDA; these can be found in other sections of chapter 6 of this user's guide, or other WRFDA documentation.

a. Running WRFDA with radiances

In addition to the basic input files (LANDUSE.TBL, fg, ob.ascii, be.dat) mentioned in the “[Running WRFDA](#)” section, the following additional files are required for radiances: radiance data (typically in NCEP BUFR format), `radiance_info` files, `VARBC.in` (if you plan on using variational bias correction VARBC, as described in [the section on bias correction](#)), and RTM (CRTM or RTTOV) coefficient files. Note that a subset of binary CRTM coefficient files is not included in the WRFDA package, and the users need to download it from http://www2.mmm.ucar.edu/wrf/users/wrfda/download/crtm_coefs.html.

Edit `namelist.input` (Pay special attention to `&wrfvar4`, `&wrfvar14`, `&wrfvar21`, and `&wrfvar22` for radiance-related options. A very basic `namelist.input` for running the radiance test case is provided in `WRFDA/var/test/radiance/namelist.input`)

```
> ln -sf $DAT_DIR/gdas1.t00z.1bamua.tm00.bufr_d ./amsua.bufr
> ln -sf $DAT_DIR/gdas1.t00z.1bamub.tm00.bufr_d ./amsub.bufr
> ln -sf $WRFDA_DIR/var/run/radiance_info ./radiance_info # (radi-
ance_info is a directory)
> ln -sf $WRFDA_DIR/var/run/VARBC.in ./VARBC.in
(CRTM only) > ln -sf directory_of_crtm_coefs ./crtm_coefs
#(crtm_coefs is a directory)
(RTTOV only) > ln -sf your_RTTOV_path/rtcoef_rttov11/rttov7pred54L
./rttov_coefs # (rttov_coefs is a directory)
(HDF5 only) > ln -sf $WRFDA_DIR/var/run/leapsec.dat .
```

See the following sections for more details on each aspect of radiance assimilation.

Note: You can also specify the path of the “`crtm_coefs`” directory via the namelist; see the following section for more details

b. Reading radiance data in WRFDA

Currently, the ingest interface for NCEP BUFR radiance data is implemented in WRFDA. The radiance data are available through NCEP's public ftp server ([ftp://ftp.ncep.noaa.gov/pub/data/nccf/com/gfs/prod/gdas.\\${yyyymmddhh}](ftp://ftp.ncep.noaa.gov/pub/data/nccf/com/gfs/prod/gdas.${yyyymmddhh})) in near real-time (with a 6-hour delay) and can meet requirements for both research purposes and some real-time applications.

WRFDA can read data from NOAA ATOVS instruments (HIRS, AMSU-A, AMSU-B and MHS), EOS Aqua instruments (AIRS, AMSU-A), DMSP instruments (SSMIS), METOP instruments (HIRS, AMSU-A, MHS, IASI), Meteosat instruments (SEVIRI), and JAXA GCOM-W1 instruments (AMSR2). Note that NCEP radiance BUFR files are separated by instrument names (i.e., one file for each type of instrument), and each file contains global radiance (generally converted to brightness temperature) within a 6-hour assimilation window, from multi-platforms. For running WRFDA, users need to rename NCEP corresponding BUFR files (table 1) to `hirs3.bufr` (including HIRS data from NOAA-15/16/17), `hirs4.bufr` (including HIRS data from NOAA-18/19, METOP-2), `amsua.bufr` (including AMSU-A data from NOAA-15/16/18/19, METOP-1 and -2), `amsub.bufr` (including AMSU-B data from NOAA-15/16/17), `mhs.bufr` (including MHS data from NOAA-18/19 and METOP-1 and -2), `airs.bufr` (including AIRS and AMSU-A data from EOS-AQUA), `ssmis.bufr` (SSMIS data from DMSP-16, AFWA provided), `iasi.bufr` (IASI data from METOP-1 and -2) and `seviri.bufr` (SEVIRI data from Meteosat 8-10) for WRFDA filename convention. Note that the `airs.bufr` file contains not only AIRS data but also AMSU-A, which is collocated with AIRS pixels (1 AMSU-A pixel collocated with 9 AIRS pixels). Users must place these files in the working directory where the WRFDA executable is run. It should also be mentioned that WRFDA reads these BUFR radiance files directly without the use of any separate pre-processing program. All processing of radiance data, such as quality control, thinning, bias correction, etc., is carried out within WRFDA. This is different from conventional observation assimilation, which requires a pre-processing package (OBSPROC) to generate WRFDA readable ASCII files. For reading the radiance BUFR files, WRFDA must be compiled with the NCEP BUFR library (see <http://www.nco.ncep.noaa.gov/sib/decoders/BUFRLIB/>).

Table 1: NCEP and WRFDA radiance BUFR file naming convention

NCEP BUFR file names	WRFDA naming convention
gdas1.t00z.airsev.tm00.bufr_d	airs.bufr
gdas1.t00z.1bamua.tm00.bufr_d	amsua.bufr
gdas1.t00z.1bamub.tm00.bufr_d	amsub.bufr
gdas1.t00z.atms.tm00.bufr_d	atms.bufr
gdas1.t00z.1bhirs3.tm00.bufr_d	hirs3.bufr
gdas1.t00z.1bhirs4.tm00.bufr_d	hirs4.bufr
gdas1.t00z.mtiiasi.tm00.bufr_d	iasi.bufr
gdas1.t00z.1bmhs.tm00.bufr_d	mhs.bufr
gdas1.t00z.sevcsr.tm00.bufr_d	seviri.bufr

Namelist parameters are used to control the reading of corresponding BUFR files into WRFDA. For instance, `USE_AMSUAOBS`, `USE_AMSUBOBS`, `USE_HIRS3OBS`,

`USE_HIRS4OBS`, `USE_MHSOBS`, `USE_AIRSOBS`, `USE_EOS_AMSUAOBS`, `USE_SSMISOBS`, `USE_ATMSOBS`, `USE_IASIOBS`, and `USE_SEVIRIOBS` control whether or not the respective file is read. These are logical parameters that are assigned to `.FALSE.` by default; therefore they must be set to `.TRUE.` to read the respective observation file. Also note that these parameters only control whether the data is read, not whether the data included in the files is to be assimilated. This is controlled by other namelist parameters explained in the next section.

Sources for downloading these and other data can be found on the WRFDA website: http://www2.mmm.ucar.edu/wrf/users/wrfda/download/free_data.html.

Other data formats

Most of the above paragraphs describe NCEP BUFR data, but some of the satellite data supported by WRFDA are in alternate formats. Level-1R AMSR2 data from the JAXA GCOM-W1 satellite are available in HDF5 format, which requires compiling WRFDA with HDF5 libraries, as described in the “[Compile WRFDA and Libraries](#)” section.

HDF5 file naming conventions are different than those for BUFR files. For AMSR2 data, WRFDA will look for two data files: `L1SGRTBR.h5` (brightness temperature) and `L2SGCLWLD.h5` (cloud liquid water). Only the brightness temperature file is mandatory (you will also need to copy or link the “`leapsec.dat`” file from `WRFDA/var/run`). If you have multiple data files for your assimilation window, you should name them `L1SGRTBR-01.h5`, `L1SGRTBR-02.h5`, etc. and `L2SGCLWLD-01.h5`, `L2SGCLWLD-02.h5`, etc.

GOES-Imager GVAR radiance data are in NETCDF format with file names like `goes-13-imager-01.nc`, `goes-13-imager-02.nc`, `goes-13-imager-03`, and `goes-13-imager-04.nc` for channel 2, 3, 4, and 6, respectively.

Newly-introduced Himawari-8-AHI radiance data in V4.1 are in NETCDF format as well with file names ‘`L1AHITBR`’ (Level-1 brightness temperature) and ‘`L2AHICLP`’ (level-2 cloud product) generated by GEOCAT software. Notice that the file ‘`~var/run/ahi_info`’ defines a sub-area of full-disk data to read.

c. Radiative Transfer Models

The core component for direct radiance assimilation is to incorporate a radiative transfer model (RTM) into the WRFDA system as one part of observation operators. Two widely used RTMs in the NWP community, RTTOV (developed by ECMWF and UKMET in Europe), and CRTM (developed by the Joint Center for Satellite Data Assimilation (JCSDA) in US), are already implemented in the WRFDA system with a flexible and consistent user interface. WRFDA is designed to be able to compile with or without RTTOV by the definition of the “RTTOV” environment variable at compile time (see the “[Compile WRFDA and Libraries](#)” section). At runtime the user must select which RTM they intend to use via the namelist parameter `RTM_OPTION` (1 for RTTOV, the default,

and 2 for CRTM).

Both RTMs can calculate radiances for almost all available instruments aboard the various satellite platforms in orbit. An important feature of the WRFDA design is that all data structures related to radiance assimilation are dynamically allocated during running time, according to a simple namelist setup. The instruments to be assimilated are controlled at run-time by four integer namelist parameters: **RTMINIT_NSENSOR** (the total number of sensors to be assimilated), **RTMINIT_PLATFORM** (the platforms IDs array to be assimilated with dimension **RTMINIT_NSENSOR**, e.g., 1 for NOAA, 9 for EOS, 10 for METOP and 2 for DMSP), **RTMINIT_SATID** (satellite IDs array) and **RTMINIT_SENSOR** (sensor IDs array, e.g., 0 for HIRS, 3 for AMSU-A, 4 for AMSU-B, etc.). The full list of instrument triplets can be found in the table below:

Instrument	Satellite	Format	(PLATFORM, SATID, SENSOR)
AIRS	EOS-Aqua	BUFR	(9,2,11)
AMSR2	GCOM-W1	HDF5	(29,1,63)
AMSU-A	EOS-Aqua	BUFR	(9,2,3)
AMSU-A	METOP-A	BUFR	(10,2,3)
AMSU-A	NOAA 15–19	BUFR	(1,15–19,3)
AMSU-B	NOAA 15–17	BUFR	(1,15–17,4)
ATMS	Suomi-NPP	BUFR	(17,0,19)
HIRS-3	NOAA 15–17	BUFR	(1,15–17,0)
HIRS-4	METOP-A	BUFR	(10,2,0)
HIRS-4	NOAA 18–19	BUFR	(1,18–19,0)
IASI	METOP-A	BUFR	(10,2,16)
IMAGER	GOES 13-15	NETCDF	(4,13-15,22)
MHS	METOP-A	BUFR	(10,2,15)
MHS	NOAA 18–19	BUFR	(1,18–19,15)
MWHS	FY-3A–FY-3B	Binary	(23,1–2,41)
MWTS	FY-3A–FY-3B	Binary	(23,1–2,40)
SEVIRI	Meteosat 8–10	BUFR	(12,1–3,21)
SSMIS	DMSP 16–18	BUFR	(2,16–18,10)
AHI	Himawari 8	NETCDF	(31, 8, 56)

Here’s an example of this section of the namelist for a user assimilating IASI observations from METOP-A, utilizing RTTOV as their RTM:

```
&wrfvar14
  rtminit_nsensor = 1
  rtminit_platform = 10,
  rtminit_satid = 2,
  rtminit_sensor = 16,
  rtm_option = 1,
/
```

Here’s another example of this section of the namelist, this time for a user assimilating AMSU-A from NOAA 18–19 and EOS-Aqua, MHS from NOAA 18–19, and AIRS from EOS-Aqua, utilizing CRTM as their RTM:

```
&wrfvar14
  rtminit_nsensor = 6
  rtminit_platform = 1, 1, 9, 1, 1, 9
  rtminit_satid = 18, 19, 2, 18, 19, 2
  rtminit_sensor = 3, 3, 3, 15, 15, 11
  rtm_option = 2,
/
```

The instrument triplets (platform, satellite, and sensor ID) in the namelist can be ranked in any order. More detail about the convention of instrument triples can be found in tables 2 and 3 in the RTTOV v11 User's Guide

(https://www.nwpsaf.eu/site/download/documentation/rtm/docs_rtov12/users_guide_rtov12_v1.3.pdf)

CRTM uses a different instrument-naming method, however, a conversion routine inside WRFDA is implemented such that the user interface remains the same for RTTOV and CRTM, using the same instrument triplet for both.

When running WRFDA with radiance assimilation switched on, a set of RTM coefficient files need to be loaded. For the RTTOV option, RTTOV coefficient files are to be copied or linked to a sub-directory `rttov_coeffs/` under the working directory. For the CRTM option, CRTM coefficient files are to be copied or linked to a sub-directory `crtm_coeffs/` under the working directory, or the location of this directory can be specified in the namelist:

```
&wrfvar14
  crt_coef_path = WRFDA/var/run/crtm_coeffs (Can be a relative or absolute path)
/
```

Only coefficients for instruments listed in the namelist are needed. Potentially WRFDA can assimilate all sensors as long as the corresponding coefficient files are provided. In addition, necessary developments on the corresponding data interface, quality control, and bias correction are important to make radiance data assimilate properly; however, a modular design of radiance relevant routines already facilitates the addition of more instruments in WRFDA.

The RTTOV package is not distributed with WRFDA, due to licensing restrictions. Users need to follow the instructions at <http://nwpsaf.eu/site/software/rttov/> to download the RTTOV source code and supplement coefficient files and the emissivity atlas dataset. Only RTTOV v11 (11.1—11.3) can be used since WRFDA version 3.9, so if you have an older version of RTTOV you must upgrade. Only RTTOV v12 is supported in version 4.1.

As mentioned in a previous paragraph, the CRTM package is distributed with WRFDA, and is located in `$WRFDA_DIR/var/external/crtm_2.3.0`. The CRTM code in WRFDA is the same as the source code that users can download from <ftp://ftp.emc.ncep.noaa.gov/jcsda/CRTM>, with only minor modifications (mainly for ease of compilation).

To use one or both of the above radiative transfer models, you will have to set the appropriate environment variable(s) at compile time. See the section “[Compile WRFDA and Libraries](#)” for details.

d. Channel Selection

Channel selection in WRFDA is controlled by radiance ‘info’ files, located in the sub-directory `radiance_info`, under the working directory. These files are separated by satellites and sensors; e.g., `noaa-15-amsua.info`, `noaa-16-amsub.info`, `dmsp-16-ssmis.info` and so on. An example of 5 channels from `noaa-15-amsub.info` is shown below. The fourth column is used by WRFDA to control when to use a corresponding channel. Channels with the value “-1” in the fourth column indicate that the channel is “not assimilated,” while the value “1” means “assimilated.” The sixth column is used by WRFDA to set the observation error for each channel. Other columns are not used by WRFDA. It should be mentioned that these error values might not necessarily be optimal for your applications. It is the user’s responsibility to obtain the optimal error statistics for his/her own applications.

Sensor	channel	IR/MW	use	idum	varch	polarization(0:vertical;1:horizontal)
415	1	1	-1	0	0.5500000000E+01	0.0000000000E+00
415	2	1	-1	0	0.3750000000E+01	0.0000000000E+00
415	3	1	1	0	0.3500000000E+01	0.0000000000E+00
415	4	1	-1	0	0.3200000000E+01	0.0000000000E+00
415	5	1	1	0	0.2500000000E+01	0.0000000000E+00

e. Bias Correction

Satellite radiance is generally considered to be biased with respect to a reference (e.g., background or analysis field in NWP assimilation) due to systematic error of the observation itself, the reference field, and RTM. Bias correction is a necessary step prior to assimilating radiance data. There are two ways of performing bias correction in WRFDA. One is based on the Harris and Kelly (2001) method, and is carried out using a set of coefficient files pre-calculated with an off-line statistics package, which was applied to a training dataset for a month-long period. The other is Variational Bias Correction (VarBC). Only VarBC is introduced here, and recommended for users because of its relative simplicity in usage.

Variational Bias Correction

To use VarBC, set the namelist option **USE_VARBC** to TRUE and have the `VARBC.in` file in the working directory. `VARBC.in` is a VarBC setup file in ASCII format. A template is provided with the WRFDA package (`$WRFDA_DIR/var/run/VARBC.in`).

All VarBC input is passed through a single ASCII file called `VARBC.in`. Once WRFDA has run with the VarBC option switched on, it will produce a `VARBC.out` file in a similar ASCII format. This output file will then be used as the input file for the next assimilation

cycle.

VarBC Coldstart

Coldstarting means starting the VarBC from scratch; i.e. when you do not know the values of the bias parameters.

The coldstart is a routine in WRFDA. The bias predictor statistics (mean and standard deviation) are computed automatically and will be used to normalize the bias parameters. All coldstart bias parameters are set to zero, except the first bias parameter (= simple offset), which is set to the mode (=peak) of the distribution of the (uncorrected) innovations for the given channel.

A threshold of a number of observations can be set through the namelist option **VARBC_NOBSMIN** (default = 10), under which it is considered that not enough observations are present to keep the coldstart values (i.e. bias predictor statistics and bias parameter values) for the next cycle. In this case, the next cycle will do another coldstart.

Background constraint for bias parameters

The background constraint controls the inertia you want to impose on the predictors (i.e. the smoothing in the predictor time series). It corresponds to an extra term in the WRFDA cost function.

It is defined in the namelist via the option **VARBC_NBGERR**; the default value is 5000. This number is related to a number of observations; the bigger the number, the more inertia constraint. If these numbers are set to zero, the predictors can evolve without any constraint.

Scaling factor

The VarBC uses a specific preconditioning, which can be scaled through the namelist option **VARBC_FACTOR** (default = 1.0).

Offline bias correction

The analysis of the VarBC parameters can be performed "offline" ; i.e. independently from the main WRFDA analysis. No extra code is needed. Just set the following **MAX_VERT_VAR*** namelist variables to be 0, which will disable the standard control variable and only keep the VarBC control variable.

```
MAX_VERT_VAR1=0.0
MAX_VERT_VAR2=0.0
MAX_VERT_VAR3=0.0
MAX_VERT_VAR4=0.0
MAX_VERT_VAR5=0.0
```

Freeze VarBC

In certain circumstances, you might want to keep the VarBC bias parameters constant in time ("frozen"). In this case, the bias correction is read and applied to the innovations, but it is not updated during the minimization. This can easily be achieved by setting the namelist options:

```
USE_VARBC=false  
FREEZE_VARBC=true
```

Passive observations

Some observations are useful for preprocessing (e.g. Quality Control, Cloud detection) but you might not want to assimilate them. If you still need to estimate their bias correction, these observations need to go through the VarBC code in the minimization. For this purpose, the VarBC uses a separate threshold on the QC values, called "qc_varbc_bad". This threshold is currently set to the same value as "qc_bad", but can easily be changed to any ad hoc value.

f. Other radiance assimilation options

RAD_MONITORING (30)

Integer array of dimension RTMINIT_NSENSOR, 0 for assimilating mode, 1 for monitoring mode (only calculates innovation).

THINNING

Logical, TRUE will perform thinning on radiance data.

THINNING_MESH (30)

Real array with dimension RTMINIT_NSENSOR, values indicate thinning mesh (in km) for different sensors.

QC_RAD

Logical, controls if quality control is performed, always set to TRUE.

WRITE_IV_RAD_ASCII

Logical, controls whether to output observation-minus-background (O-B) files, which are in ASCII format, and separated by sensors and processors.

WRITE_OA_RAD_ASCII

Logical, controls whether to output observation-minus-analysis (O-A) files (including also O-B information), which are in ASCII format, and separated by sensors and processors.

USE_ERROR_FACTOR_RAD

Logical, controls use of a radiance error tuning factor file

(`radiance_error.factor`) which is created with empirical values, or generated using a variational tuning method (Desroziers and Ivanov, 2001).

ONLY_SEA_RAD

Logical, controls whether only assimilating radiance over water.

TIME_WINDOW_MIN

String, e.g., "2007-08-15_03:00:00.0000", start time of assimilation time window

TIME_WINDOW_MAX

String, e.g., "2007-08-15_09:00:00.0000", end time of assimilation time window

USE_ANTCORR (30)

Logical array with dimension RTMINIT_NSENSOR, controls if performing Antenna Correction in CRTM.

USE_CLDDET

Integer, controls cloud detection scheme for infrared radiance.

0: no cloud detection

1: MMR scheme

2: Particle filter scheme (default)

3: ECMWF scheme

AIRS_WARMEST_FOV

Logical, controls whether using the observation brightness temperature for AIRS Window channel #914 as criterion for GSI thinning.

USE_CRTM_KMATRIX

Logical, controls whether using the CRTM K matrix rather than calling CRTM TL and AD routines for gradient calculation.

CRTM_CLOUD

Logical, include cloud effects in CRTM calculations.

Further information on this capability can be found in the following publication:

Chun Yang, Zhiquan Liu, Jamie Bresch, Syed R. H. Rizvi, Xiang-Yu Huang and Jinzhong Min, 2016: [AMSR2 all-sky radiance assimilation and its impact on the analysis and forecast of Hurricane Sandy with a limited-area data assimilation system](#). *Tellus A*, **68**, 30917, doi:10.3402/tellusa.v68.30917.

USE_RTTOV_KMATRIX

Logical, controls whether using the RTTOV K matrix rather than calling RTTOV TL and AD routines for gradient calculation.

RTTOV_EMIS_ATLAS_IR

Integer, controls the use of the IR emissivity atlas.

Emissivity atlas data (should be downloaded separately from the RTTOV web

site) need to be copied or linked under a sub-directory of the working directory (`emis_data`) if `RTTOV_EMIS_ATLAS_IR` is set to 1.

RTTOV_EMIS_ATLAS_MW

Integer, controls the use of the MW emissivity atlas.

Emissivity atlas data (should be downloaded separately from the RTTOV web site) need to be copied or linked under a sub-directory of the working directory (`emis_data`) if `RTTOV_EMIS_ATLAS_MW` is set to 1 or 2.

g. Diagnostics and Monitoring

Monitoring capability within WRFDA

Run WRFDA with the `rad_monitoring` namelist parameter in record `wrfvar14` in `namelist.input`.

0 means assimilating mode. Innovations (O minus B) are calculated and data are used in minimization.

1 means monitoring mode: innovations are calculated for diagnostics and monitoring. Data are not used in minimization.

The value of `rad_monitoring` should correspond to the value of `rtminit_nsensor`. If `rad_monitoring` is not set, then the default value of 0 will be used for all sensors.

Outputting radiance diagnostics from WRFDA

Run WRFDA with the following namelist options in record `wrfvar14` in `namelist.input`.

write_iv_rad_ascii

Logical. TRUE to write out (observation-background, etc.) diagnostics information in plain-text files with the prefix 'inv,' followed by the instrument name and the processor id. For example, `01_inv_noaa-17-amsub.0000` (01 is outerloop index, 0000 is processor index)

write_oa_rad_ascii

Logical. TRUE to write out (observation-background, observation-analysis, etc.) diagnostics information in plain-text files with the prefix 'oma,' followed by the instrument name and the processor id. For example, `01_oma_noaa-18-mhs.0001`

Each processor writes out the information for one instrument in one file in the

WRFDA working directory.

Radiance diagnostics data processing

One of the 44 executables compiled as part of the WRFDA system is the file `da_rad_diags.exe`. This program can be used to collect the `01_inv*` or `01_oma*` files and write them out in netCDF format (one instrument in one file with prefix `diags` followed by the instrument name, analysis date, and the suffix `.nc`) for easier data viewing, handling and plotting with netCDF utilities and NCL scripts. See `WRFDA/var/da/da_monitor/README` for information on how to use this program.

Radiance diagnostics plotting

Two NCL scripts (available as part of the WRFDA Tools package, which can be downloaded at <http://www2.mmm.ucar.edu/wrf/users/wrfda/download/tools.html>) are used for plotting: `$TOOLS_DIR/var/graphics/ncl/plot_rad_diags.ncl` and `$TOOLS_DIR/var/graphics/ncl/advance_cymdh.ncl`. The NCL scripts can be run from a shell script, or run alone with an interactive `ncl` command (the NCL script and set the plot options must be edited, and the path of `advance_cymdh.ncl`, a date-advancing script loaded in the main NCL plotting script, may need to be modified).

Steps (3) and (4) can be done by running a single ksh script (also in the WRFDA Tools package: `$TOOLS_DIR/var/scripts/da_rad_diags.ksh`) with proper settings. In addition to the settings of directories and what instruments to plot, there are some useful plotting options, explained below.

<code>setenv OUT_TYPE=ncgm</code>	ncgm or pdf pdf will be much slower than ncgm and generate huge output if plots are not split. But pdf has higher resolution than ncgm.
<code>setenv PLOT_STATS_ONLY=false</code>	true or false true: only statistics of OMB/OMA vs channels and OMB/OMA vs dates will be plotted. false: data coverage, scatter plots (before and after bias correction), histograms (before and after bias correction), and statistics will be plotted.
<code>setenv PLOT_OPT=sea_only</code>	all, sea_only, land_only
<code>setenv PLOT_QCED=false</code>	true or false true: plot only quality-controlled data false: plot all data
<code>setenv PLOT_HISTO=false</code>	true or false: switch for histogram plots
<code>setenv PLOT_SCATT=true</code>	true or false: switch for scatter plots
<code>setenv PLOT_EMISS=false</code>	true or false: switch for emissivity plots
<code>setenv PLOT_SPLIT=false</code>	true or false true: one frame in each file false: all frames in one file

<code>setenv PLOT_CLOUDY=false</code>	true or false true: plot cloudy data. Cloudy data to be plotted are defined by PLOT_CLOUDY_OPT (si or clwp), CLWP_VALUE, SI_VALUE settings.
<code>setenv PLOT_CLOUDY_OPT=si</code>	si or clwp clwp: cloud liquid water path from model si: scatter index from obs, for amsua, amsub and mhs only
<code>setenv CLWP_VALUE=0.2</code>	only plot points with clwp >= clwp_value (when clwp_value > 0) clwp > clwp_value (when clwp_value = 0)
<code>setenv SI_VALUE=3.0</code>	

Evolution of VarBC parameters

NCL scripts (also in the WRFDA Tools package: \$TOOLS_DIR/var/graphics/ncl/plot_rad_varbc_param.ncl and \$TOOLS_DIR/var/graphics/ncl/advance_cymdh.ncl) are used for plotting the evolution of VarBC parameters.

Radar Data Assimilation in WRFDA

WRFDA has the ability to assimilate Doppler radar data, either for 3DVAR or 4DVAR assimilation. Both Doppler velocity and reflectivity can be assimilated, and there are several different reflectivity operator options available.

a. Preparing radar observations

Radar observations are read by WRFDA in a text-based format. This format is described in the radar tutorial presentation available on the WRFDA website (http://www2.mmm.ucar.edu/wrf/users/wrfda/Tutorials/2015_Aug/docs/WRFDA_Radar.pdf). Because radar data comes in a variety of different formats, it is the user's responsibility to convert their data into this format. For 3DVAR, these observations should be placed in a file named `ob.radar`. For 4DVAR, they should be placed in files named `ob01.radar`, `ob02.radar`, etc., with one observation file per time slot, as described in the earlier [4DVAR section](#).

b. Running WRFDA for radar assimilation

Once your observations are prepared, you can run WRFDA the same as you would normally (see the previous sections on how to run either 3DVAR or 4DVAR). For guidance, there is an example 3DVAR case available for download at http://www2.mmm.ucar.edu/wrf/users/wrfda/download/V38/wrfda_radar_testdata.tar.gz.

Edit `namelist.input` and pay special attention to the radar options listed below. Further details on some of these options can be found in the following sections

```

&wrfvar4
use_radarobs      true: radar observation files will be read by WRFDA
use_radar_rv      true: Assimilate radar velocity observations
use_radar_rf      true: Assimilate radar reflectivity using total mixing ratio
radar_rf_opt      1 or 2: Choose radar reflectivity option
use_radar_rhv     true: Assimilate retrieved hydrometeors (qr, qs, qg) from radar reflectivity
use_radar_rqv     true: Assimilate estimated humidity (qv) from radar reflectivity
/

...
&wrfvar7
cloud_cv_options  0 (default): no hydrometeor control variables
                  1: use total water (water vapor+cloud liquid water+rain water) control variable
                  2: use individual hydrometeor control variables with statistical error covariances
                     in be.dat file produced using gen_be_v3 program available in WRFDA package
                     or https://github.com/wrf-model/GENBE 2.0
                  3: use individual hydrometeor control variables with hard-coded error
                     covariances
use_cv_w          false (default): use  $\omega$  (vertical velocity with respect to pressure) control variable
                  true: use W (vertical velocity with respect to height) control variable.
                  For cloud_cv_options = 3 only
/

...
&radar_da
radar_non_precip_opt  0 (default): no null-echo assimilation
                     1: KNU null-echo scheme
radar_non_precip_rf   Reflectivity flag value (dBz) in observation file indicating non-
                     precipitation echoes (default: -999.99)

The following options apply for use_radar_rqv only:
radar_non_precip_rh_w RH (%) with respect to water for non-precip retrieved Q_vapor (rqv) (de-
                     fault: 95)
radar_non_precip_rh_i RH (%) with respect to ice for non-precip retrieved rqv (default: 85)
cloudbase_calc_opt    Option for calculating cloud-base height: below this height retrieved hu-
                     midity will not be assimilated for the use_radar_rqv option
                     0: fixed value of 1500 meters
                     1 (default): KNU scheme
                     2: NCAR scheme
radar_saturated_rf    rf value (dBz) used to indicate precipitation for rqv (default 25.0)
radar_rqv_thresh1     rf value (dBz) used to scale down retrieved rqv (default 40.0)
radar_rqv_thresh2     rf value (dBz) used to scale down retrieved rqv (default 50.0)
radar_rqv_rh1         RH (%) for radar_saturated_rf < rf < radar_rqv_thresh1 (default 85)
radar_rqv_rh2         RH (%) for radar_rqv_thresh1 < rf < radar_rqv_thresh2 (default 95)
radar_rqv_h_lbound    height (meters) lower bound for assimilating rqv (default -999.0)
radar_rqv_h_ubound    height (meters) upper bound for assimilating rqv (default -999.0)
NOTE: both namelist settings radar_rqv_h_lbound and ra-
dar_rqv_h_ubound must be set and greater than zero for either to
have an impact

```

c. Reflectivity assimilation options

There are three different options for assimilating radar reflectivity data. The first directly assimilates the observed reflectivity using a reflectivity operator to convert the model rainwater mixing ratio into reflectivity and the total mixing ratio as the control variable, as described in Xiao and Sun, 2007

(<http://journals.ametsoc.org/doi/full/10.1175/MWR3471.1>). For this option, the hydrometeors are partitioned using a warm rain scheme described in the above reference. Namelist setting for this option is given below.

```
radar_rf_opt = 1,
use_radar_rf = T,
use_radar_rhv = F,
use_radar_rqv = F,
cloud_cv_options = 1,
```

The second (`use_radar_rhv`) is a scheme described in Wang et al, 2013 (<http://journals.ametsoc.org/doi/full/10.1175/JAMC-D-12-0120.1>), which assimilates rainwater mixing ratio that is estimated from radar reflectivity, described as an “indirect method” in the paper. This second option also includes an option (`use_radar_rqv`) that allows the assimilation of in-cloud humidity estimated from reflectivity using a method described in Wang et al, 2013. It also includes the assimilation of snow and graupel converted from reflectivity using formulas as described in Gao and Stensrud, 2012

(<http://journals.ametsoc.org/doi/abs/10.1175/JAS-D-11-0162.1>). Namelist setting for this (default) option is given below.

```
radar_rf_opt = 1, ! no effect for this option
use_radar_rf = F,
use_radar_rhv = T,
use_radar_rqv = T, ! or false
cloud_cv_options = 2, ! or 3
```

There are many tunable parameters that go along with the `use_radar_rqv` option, which retrieves a value of cloud humidity for assimilation. There are three possible options for cloud base height (below which cloud humidity will not be assimilated) as specified by `cloudbase_calc_opt` (0 is the previous default behavior and is not recommended). There are also a few thresholds for scaling the calculated cloud humidity by certain amounts, as well as an upper and lower altitude bound for assimilating cloud humidity.

The third option newly available in version 4.2 directly assimilates reflectivity with a new observation operator and its TL/AD taking into account snow and graupel (Wang and Liu, 2019, <https://www.geosci-model-dev.net/12/4031/2019/>). Namelist setting for this (default) option is given below.

```
radar_rf_opt = 2,
use_radar_rf = T,
use_radar_rhv = F,
use_radar_rqv = F,
cloud_cv_options = 2, ! or 3
```


d. Null-echo assimilation options

WRFDA includes the capability to assimilate null-echo observations (`radar_non_precip_opt=1`): reflectivity values with a set flag value (`radar_non_precip_rf`) will be assimilated as non-precipitation points. This can be an important radar assimilation technique, as normally you can not remove precipitation from your analysis without using the retrieved water vapor option (`use_radar_rqv`). This capability was developed by Ki-Hong Min from Kyungpook National University, South Korea (WRF Workshop abstract:

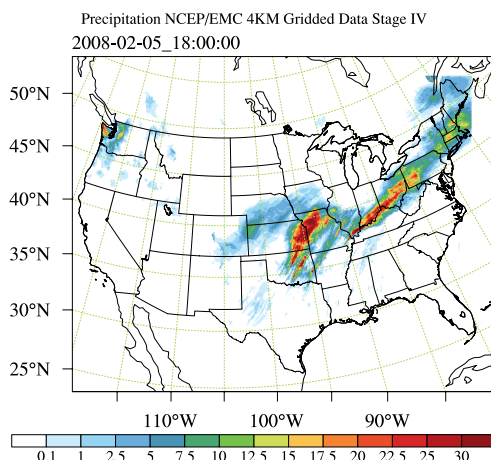
http://www2.mmm.ucar.edu/wrf/users/workshops/WS2016/short_abstracts/P78.pdf).

Precipitation Data Assimilation in WRFDA 4DVAR

The assimilation of precipitation observations in WRFDA 4D-Var is described in this section. Currently, WRFPLUS has already included the adjoint and tangent linear codes of large-scale condensation and cumulus scheme, therefore precipitation data can be assimilated directly in 4D-Var. Users who are interested in the scientific detail of 4D-Var assimilation of precipitation should refer to related scientific papers, as this section is only a basic guide to running WRFDA Precipitation Assimilation. This section instructs users on data processing, namelist variable settings, and how to run WRFDA 4D-Var with precipitation observations.

a. Preparing precipitation observations

WRFDA 4D-Var can assimilate NCEP Stage IV radar and gauge precipitation data. NCEP Stage IV archived data are available on the NCAR CODIAC web page at: <http://data.eol.ucar.edu/codiac/dss/id=21.093> (for more information, please see the NCEP Stage IV Q&A Web page at <http://www.emc.ncep.noaa.gov/mmb/ylin/pcpanl/QandA/>). The original precipitation data are at 4-km resolution on a polar-stereographic grid. Hourly, 6-hourly and 24-hourly analyses are available. The following image shows the accumulated 6-h precipitation for the tutorial case.



It should be mentioned that the NCEP Stage IV archived data is in GRIB1 format and it cannot be ingested into the WRFDA directly. A tool “precip_converter” is provided to reformat GRIB1 observations into the WRFDA-readable ASCII format. It can be downloaded from the WRFDA users page at

http://www2.mmm.ucar.edu/wrf/users/wrfda/download/precip_converter.tar.gz. The NCEP GRIB libraries, w3 and g2 are required to compile the precip_converter utility. These libraries are available for download from NCEP at <http://www.nco.ncep.noaa.gov/pmb/codes/GRIB2/>. The output file to the precip_converter utility is named in the format `ob.rain.yyyymmddhh.xxh`; The 'yyymmddhh' in the file name is the ending hour of the accumulation period, and 'xx' (=01,06 or 24) is the accumulating time period.

For users wishing to use their own observations instead of NCEP Stage IV, it is the user's responsibility to write a Fortran main program and call subroutine `writerainobs` (in `write_rainobs.f90`) to generate their own precipitation data. For more information please refer to the README file in the `precip_converter` directory.

b. Running WRFDA with precipitation observations

WRFDA 4D-Var is able to assimilate hourly, 3-hourly and 6-hourly precipitation data. According to experiments and related scientific papers, 6-hour precipitation accumulations are the ideal observations to be assimilated, as this leads to better results than directly assimilating hourly data.

The tutorial example is for assimilating 6-hour accumulated precipitation. In your working directory, link all the necessary files as follows,

```
> ln -fs $WRFDA_DIR/var/da/da_wrfvar.exe .
> ln -fs $DAT_DIR/rc/2008020512/wrfinput_d01 .
> ln -fs $DAT_DIR/rc/2008020512/wrfbdy_d01 .
```

```

> ln -fs $DAT_DIR/rc/2008020518/wrfinput_d01 fg02 (only necessary
for var4d_lbc=true)
> ln -fs wrfinput_d01 fg
> ln -fs $DAT_DIR/be/be.dat .
> ln -fs $WRFDA_DIR/run/LANDUSE.TBL .
> ln -fs $WRFDA_DIR/run/RRTM_DATA_DBL ./RRTM_DATA
> ln -fs $DAT_DIR/ob/2008020518/ob.rain.2008020518.06h ob07.rain

```

Note: The reason why the observation ob.rain.2008020518.06h is linked as ob07.rain will be explained in section c.

Edit namelist.input (you can start with the same namelist as for the 4dvar tutorial case) and pay special attention to &wrfvar1 and &wrfvar4 for precipitation-related options.

```

&wrfvar1
var4d          true: Run WRFDA for 4DVAR. This is the only supported option for precipitation
               assimilation (default value is false)
var4d_bin_rain length (seconds) of the precipitation assimilation window (default 3600). This can be
               different from var4d_bin, which controls the assimilation window for all other ob-
               servation types
/
...
&wrfvar4
use_rainobs    true (default) : read precipitation data
thin_rainobs   true (default): thin precipitation observations
thin_mesh_conv Size of thinning mesh (in km) for non-radiance observations, including precipitation
               observations (default value 20.0)
/

```

Then, run 4D-Var in serial or parallel mode,

```
> ./da_wrfvar.exe >& wrfda.log
```

c. Properly linking observation files

In section b, ob.rain.2008020518.06h is linked as ob07.rain. The number 07 is assigned according to the following rule:

$$x = i * (\text{var4d_bin_rain} / \text{var4d_bin}) + 1,$$

Here, i is the sequence number of the observation.

for $x < 10$, the observation file should be renamed as ob0x.rain;

for $x \geq 10$, it should be renamed as obx.rain

In the example above, 6-hour accumulated precipitation data is assimilated in 6-hour time window. In the namelist, values should be set at var4d_bin=3600 and var4d_bin_rain=21600, and there is one observation file (i.e., $i=1$) in the time window, Thus the value of x is 7. The file ob.rain.2008020518.06h should be renamed as ob07.rain.

Let us take another example for how to rename observation files for 3-hourly precipitation data in 6-hour time window. The sample namelist is as follows,

```
&wrfvar1
var4d=true,
var4d_lbc=true,
var4d_bin=3600,
var4d_bin_rain=10800,
.....
/
```

There are two observation files, `ob.rain.2008020515.03h` and `ob.rain.2008020518.03h`. For the first file (`i=1`) `ob.rain.2008020515.03h`, it should be renamed as `ob04.rain`, and the second file (`i=2`) renamed as `ob07.rain`.

Updating WRF Boundary Conditions

The ultimate goal of WRFDA is to combine a WRF file (`wrfinput` or `wrfout`) with observations and error information, in order to produce a “best guess” of the atmospheric state for your domain. While this “best guess” can be useful on its own for research purposes, it is often more useful as the initial conditions to a WRF forecast, so that the better initial conditions will ultimately provide a better forecast.

A common use of WRF/WRFDA for research or realtime forecast purposes is by following these steps:

1. Generate initial conditions for WRF (`wrfinput`) via WPS (as described in [Chapter 3 of this users guide](#)) and `real.exe` (as described in [Chapter 5 of this users guide](#))
2. Run WRFDA on this `wrfinput` to assimilate observations and produce a `wrfvar_output` file (a new, improved `wrfinput`)
3. **Run `da_update_bc.exe` to update the WRF lateral boundary conditions file created in step 1 (`wrfbdy_d01`) to be consistent with the new `wrfinput_d01` file**
4. Run `wrf.exe` to produce a WRF forecast

The highlighted step 3 will be described in the following section.

a. Lateral boundary conditions

When using a WRFDA analysis (`wrfvar_output`) to run a WRF forecast, it is essential that you update the WRF lateral boundary conditions (contained in the file `wrfbdy_d01`, created by `real.exe`) to match your new analysis. Domain-1 (`wrfbdy_d01`) must be updated to be consistent with the new WRFDA initial condition (analysis). This is absolutely essential for domain 1; otherwise your forecast will have discontinuities and poor results at the boundary. For nested domains, domain-2, domain-

3, etc., the lateral boundary conditions are provided by their parent domains, so no lateral boundary update is needed for these domains. The update procedure is performed by the WRFDA utility called `da_update_bc.exe`, and after compilation can be found in `$WRFDA_DIR/var/build`.

`da_update_bc.exe` requires three input files: the WRFDA analysis (`wrfvar_output`), the `wrfbdy` file from `real.exe`, and a namelist file: `parame.in`. To run `da_update_bc.exe` to update lateral boundary conditions, follow the steps below:

```
> cd $WRFDA_DIR/var/test/update_bc
> cp -p $DAT_DIR/rc/2008020512/wrfbdy_d01 .
    (IMPORTANT: make a copy of wrfbdy_d01, as the wrf_bdy_file will be over-
    written by da_update_bc.exe)
> vi parame.in
&control_param
da_file           = '../tutorial/wrfvar_output'
wrf_bdy_file      = './wrfbdy_d01'
domain_id         = 1
debug             = .true.
update_lateral_bdy = .true.
update_low_bdy    = .false.
update_lsm        = .false.
iswater           = 16
var4d_lbc         = .false.
/

> ln -sf $WRFDA_DIR/var/da/da_update_bc.exe .
> ./da_update_bc.exe
```

At this stage, you should have the files `wrfvar_output` and `wrfbdy_d01` in your WRFDA working directory. They are the WRFDA updated initial and boundary condition files for any subsequent WRF model runs. To use, link a copy of `wrfvar_output` and `wrfbdy_d01` to `wrfinput_d01` and `wrfbdy_d01`, respectively, in your WRF working directory.

You should also see two additional output files: `fort.11` and `fort.12`. These contain information about the changes made to `wrfbdy_d01`.

Note: The above instructions for updating lateral boundary conditions do not apply for child domains (`wrfinput_d02`, `wrfinput_d03`, etc.). This is because the lateral boundary conditions for these domains come from the respective parent domains, so `update_bc` is not necessary after running WRFDA when a child domain is used as the first guess.

b. Cycling with WRF and WRFDA

While the above procedure is useful, often for realtime applications it is better to run a so-called “cycling” forecast. In a WRF/WRFDA cycling system, rather than using WPS to generate the initial conditions for your assimilation/forecast, the output from a previ-

ous forecast is used. In this way, information from previous observations can be used to improve the current “first guess” of the atmosphere, ultimately resulting in an even better analysis and forecast. The procedure for cycling is as follows:

1. For your initial forecast time (T1), generate initial and boundary conditions for WRF (`wrfinput` and `wrfbdy_d01`) via WPS (as described in [Chapter 3 of this users guide](#)) and `real.exe` (as described in [Chapter 5 of this users guide](#))
2. Run WRFDA on this `wrfinput` to assimilate observations to produce a `wrfvar_output` file (a new, improved `wrfinput`)
3. Run `da_update_bc.exe` to update the WRF lateral boundary conditions file created in step 1 (`wrfbdy_d01`) to be consistent with the new `wrfinput` file
4. Run `wrf.exe` to produce a WRF forecast (`wrfout`) for the next forecast time (T2)
5. Repeat step 1 for the next forecast time (T2) to produce initial and boundary conditions for WRF (`wrfinput` and `wrfbdy_d01`) via WPS and `real.exe`
6. **Run `da_update_bc.exe` to update the lower boundary conditions of the WRF forecast file (`wrfout`) with data from the `wrfinput` file generated in step 5.**
7. Run WRFDA on this `wrfout` file to assimilate observations and produce a `wrfvar_output` file (a new, improved `wrfinput` for the next WRF forecast)
8. Run `da_update_bc.exe` *again* to update the WRF boundary conditions file created in step 5 (`wrfbdy_d01`) to be consistent with the new `wrfinput` file
9. Run `wrf.exe` to produce a WRF forecast (`wrfout`) for the next forecast time (T2)
10. Repeat steps 5-9 for the next forecast time(s) ad infinitum (T3, T4, T5...)

In cycling mode, as you can see above, the program `da_update_bc.exe` is used for two distinct purposes: prior to running WRFDA it updates the *lower* boundary conditions of the WRF forecast file that is used as the first guess for WRFDA, then after running WRFDA it updates the *lateral* boundary conditions file (`wrfbdy_d01`) to be consistent with the WRFDA output (a new, improved `wrfinput` for the next WRF forecast). The use of `da_update_bc.exe` to update the lateral boundary conditions was covered in the previous section, and this section will cover the second use: using `da_update_bc.exe` to update the *lower* boundary conditions.

The reason that this additional step is necessary for cycling forecasts: While a WRF forecast integrates atmospheric variables forward in time, it does not update certain lower boundary conditions, such as vegetation fraction, sea ice, snow cover, etc, which are important for both forecasts and data assimilation. For short periods of time, this is not a problem, as these fields do not tend to evolve much over the course of a few days. How-

ever, for a cycling forecast that runs for weeks, months, or even years, it is essential to update these fields regularly from the initial condition files through WPS.

To do this, prior to the assimilation process, the first guess file needs to be updated based on the information from the `wrfinput` file, generated by WPS/real.exe at analysis time. You should run `da_update_bc.exe` with the following namelist options:

```
da_file           = './fg'
wrf_input         = './wrfinput_d01'
update_lateral_bdy = .false.
update_low_bdy    = .true.
iswater          = 16
```

Note: “iswater” (water point index) is 16 for USGS LANDUSE and 17 for MODIS LANDUSE.

This creates a lower-boundary updated first guess (`da_file` will be overwritten by `da_update_bc` with updated lower boundary conditions from `wrf_input`). Then, after WRFDA has finished, run `da_update_bc.exe` again with the following namelist options:

```
da_file           = './wrfvar_output'
wrf_bdy_file      = './wrfbdy_d01'
update_lateral_bdy = .true.
update_low_bdy    = .false.
```

This updates the lateral boundary conditions (`wrf_bdy_file` will be overwritten by `da_update_bc` with lateral boundary conditions from `da_file`).

As mentioned previously, lateral boundary conditions for child domains (`wrfinput_d02`, `wrfinput_d03`, etc.) come from the respective parent domains, so `update_bc` is not necessary after running WRFDA. However, in a cycling procedure, the lower boundaries in each of the nested domains’ WRFDA analysis files still need to be updated. In these cases, you must set the namelist variable, `domain_id > 1` (default is 1 for domain 1) and provide the appropriate `wrfinput` file (`wrf_input = './wrfinput_d02'` for domain 2, for instance).

c. WRFDA 4DVAR with lateral boundary conditions as control variables

If you activate the `var4d_lbc` option in a WRF 4D-Var run, in addition to the above-mentioned files you will also need the `ana02` file from the WRFDA working directory. In `parame.in`, set `var4d_lbc` to TRUE and use “`da_file_02`” to point to the location of the `ana02` file.

```
da_file_02        = './ana02'
var4d_lbc         = .true.
```

Background Error and running GEN_BE

Quick-start guide: running WRFDA with different BE options

To run WRFDA with the generic CV3 option, simply link the provided be.dat file in the var/run directory:

```
> cp -p $WRFDA_DIR/var/run/be.dat.cv3 $WORK_DIR/be.dat
```

To run WRFDA with any other option, you will have to run GEN_BE first. GEN_BE takes a series of forecasts initialized at different times, and compares the forecasts that are valid at the same time (e.g., compare a 24-hour forecast initialized at 00Z with a 12-hour forecast initialized at 12Z) to get an estimate of the background error statistics.

You will use the wrapper script `gen_be_wrapper.ksh` to run GEN_BE. For instructions on how to set up your experiment to run GEN_BE, reference the test case as described in the section “[Domain-specific background error options: Running GEN_BE](#)” below.

Background error options in WRFDA

Users have four choices to define the background error covariance (BE). We call them CV3, CV5, CV6, and CV7. Each of these has different properties, which are outlined in the table below:

<i>CV option</i>	<i>Data source</i>	<i>Control variables</i>	<i>cv_options =</i>
CV3	Provided be.dat file	$\psi, \chi_u, T_u, q, P_{s,u}$	3
CV5	GEN_BE	$\psi, \chi_u, T_u, RH_s, P_{s,u}$	5
CV6	GEN_BE	$\psi, \chi_u, T_u, RH_{s,u}, P_{s,u}$	6
CV7	GEN_BE	u, v, T, RH_s, P_s	7

With CV3, the control variables are in physical space while with CV5, CV6, and CV7, the control variables are in eigenvector space. The major difference between these two kinds of BE is the vertical covariance; CV3 uses the vertical recursive filter to model the vertical covariance but the others use an empirical orthogonal function (EOF) to represent the vertical covariance. The recursive filters to model the horizontal covariance are also different with these BEs. We have not conducted the systematic comparison of the analyses based on these BEs. However, CV3 (a BE file provided with our WRFDA system) is a global BE and can be used for any regional domain, while CV5, CV6, and CV7 BE's are domain-dependent, and so should be generated based on forecast or ensemble data from the same domain.

As summarized in the above table, CV5, CV6, and CV7 differ in the control variables they use. CV5 utilizes streamfunction (ψ), unbalanced velocity potential (χ_u), unbalanced

temperature (T_u), pseudo relative humidity (RH_s), and unbalanced surface pressure ($P_{s,u}$). The pseudo relative humidity is defined as $Q/Q_{b,s}$, where $Q_{b,s}$ is the saturated specific humidity from the background field. For CV6 the moisture control variable is the unbalanced portion of the pseudo relative humidity ($RH_{s,u}$). Additionally, CV6 introduces six additional correlation coefficients in the definition of the balanced part of analysis control variables. See the section [GEN_BE for CV6](#) for more details on this option. Finally, CV7 uses a different set of control variables: u , v , temperature, pseudo relative humidity (RH_s), and surface pressure (P_s).

Generic BE option: CV3

CV3 is the NCEP background error covariance. It is estimated in grid space by what has become known as the NMC method (Parrish and Derber 1992). The statistics are estimated with the differences of 24 and 48-hour GFS forecasts with T170 resolution, valid at the same time for 357 cases, distributed over a period of one year. Both the amplitudes and the scales of the background error have to be tuned to represent the forecast error in the estimated fields. The statistics that project multivariate relations among variables are also derived from the NMC method.

The variance of each variable, and the variance of its second derivative, are used to estimate its horizontal scales. For example, the horizontal scales of the stream function can be estimated from the variance of the vorticity and stream function.

The vertical scales are estimated with the vertical correlation of each variable. A table is built to cover the range of vertical scales for the variables. The table is then used to find the scales in vertical grid units. The filter profile and the vertical correlation are fitted locally. The scale of the best fit from the table is assigned as the scale of the variable at that vertical level for each latitude. Note that the vertical scales are locally defined so that the negative correlation further away, in the vertical direction, is not included.

Theoretically, CV3 BE is a generic background error statistics file which can be used for any case. It is quite straightforward to use CV3 in your own case. To use the CV3 BE file in your case, set `cv_options=3` in `&wrfvar7` in `namelist.input` in your working directory, and use the `be.dat` is located in `WRFDA/var/run/be.dat.cv3`.

Modifying CV3 length scales and variance

Because CV3 is a generic background error option, it is necessary to tune the default background error scale length and variance values for each experiment. These can be controlled at run time with a series of namelist variables described below.

The scaling factors for CV3 are stored as an array of values for each individual control variable:

```
as1: stream function
as2: unbalanced velocity potential
```

```
as3: unbalanced temperature
as4: pseudo relative humidity
as5: unbalanced surface pressure
```

These variables are all 3-element vectors. The first element is the variance scaling factor. The second is the horizontal length scale factor. The third is the vertical length scale factor. So setting the default values in your namelist would look like this:

```
&wrfvar7
  cv_options = 3,
  as1       = 0.25,   1.00,   1.50,
  as2       = 0.25,   1.00,   1.50,
  as3       = 0.25,   1.00,   1.50,
  as4       = 0.25,   1.00,   1.50,
  as5       = 0.25,   1.00,   1.50,
/
```

The first column is the variance, the second is the horizontal length scale factor, and the third is the vertical length scale factor.

For multiple outer loops, the next 3 elements of each vector must be filled in. So for 2 outer loops (`max_ext_its=2`), to use the default values, the namelist should look like this:

```
&wrfvar7
  cv_options = 3,
  as1       = 0.25,   1.00,   1.50,   0.25,   1.00,   1.50,
  as2       = 0.25,   1.00,   1.50,   0.25,   1.00,   1.50,
  as3       = 0.25,   1.00,   1.50,   0.25,   1.00,   1.50,
  as4       = 0.25,   1.00,   1.50,   0.25,   1.00,   1.50,
  as5       = 0.25,   1.00,   1.50,   0.25,   1.00,   1.50,
/
```

Again, the first column is the variance, the second is the horizontal length scale factor, and the third is the vertical length scale factor for the first outer loop. The fourth column is the variance, the fifth is the horizontal length scale factor, and the sixth is the vertical length scale factor for the second outer loop.

Continue in this manner for more outer loops. The values listed above are the default values, but can be adjusted for each individual experiment.

Domain-specific background error options: Running GEN_BE

To use CV5, CV6 or CV7 background error covariance, it is necessary to generate your domain-specific background error statistics with the `gen_be` utility. The default CV3 background error statistics file, supplied with the WRFDA source code, can NOT be used with these control variable options.

The main source code for the various `gen_be` stages can be found in `WRFDA/var/gen_be`. The executables of `gen_be` should have been created when you compiled the WRFDA

code (as described earlier). The scripts to run these codes are in `WRFDA/var/scripts/gen_be`. The user can run `gen_be` using the wrapper script `WRFDA/var/scripts/gen_be/gen_be_wrapper.ksh`.

The input data for `gen_be` are WRF forecasts, which are used to generate model perturbations, used as a proxy for estimates of forecast error. For the NMC-method, the model perturbations are differences between forecasts (e.g. T+24 minus T+12 is typical for regional applications, T+48 minus T+24 for global) valid at the same time. Climatological estimates of background error may then be obtained by averaging these forecast differences over a period of time (e.g. one month). Given input from an ensemble prediction system (EPS), the inputs are the ensemble forecasts, and the model perturbations created are the transformed ensemble perturbations. The `gen_be` code has been designed to work with either forecast difference or ensemble-based perturbations. The former is illustrated in this tutorial example.

It is important to include forecast differences valid at different parts of the day (for example, forecasts valid at 00Z and 12Z through the forecast period) to remove contributions from the diurnal cycle (i.e. do not run `gen_be` using model perturbations valid for a single time each day).

The inputs to `gen_be` are netCDF WRF forecast output ("wrfout") files at specified forecast ranges. To avoid unnecessary large single data files, it is assumed that all forecast ranges are output to separate files. For example, if we wish to calculate BE statistics using the NMC-method with (T+24)-(T+12) forecast differences (default for regional) then by setting the WRF `namelist.input` options `history_interval=720`, and `frames_per_outfile=1` we get the necessary output datasets. Then the forecast output files should be arranged as follows: directory name is the forecast initial time, time info in the file name is the forecast valid time. `2008020512/wrfout_d01_2008-02-06_00:00:00` means a 12-hour forecast valid at `2008020600`, initialized at `2008020512`.

Example dataset for a test case (90 x 60 x 41 gridpoints) can be downloaded from <http://www2.mmm.ucar.edu/wrf/users/wrfda/download/testdata.html>. Untar the `gen_be_forecasts_20080205.tar.gz` file. You will have:

```
>ls $FC_DIR
-rw-r--r-- 1 users 11556492 2008020512/wrfout_d01_2008-02-06_00:00:00
-rw-r--r-- 1 users 11556492 2008020512/wrfout_d01_2008-02-06_12:00:00
-rw-r--r-- 1 users 11556492 2008020600/wrfout_d01_2008-02-06_12:00:00
-rw-r--r-- 1 users 11556492 2008020600/wrfout_d01_2008-02-07_00:00:00
-rw-r--r-- 1 users 11556492 2008020612/wrfout_d01_2008-02-07_00:00:00
-rw-r--r-- 1 users 11556492 2008020612/wrfout_d01_2008-02-07_12:00:00
```

In the above example, only 1 day (12Z 05 Feb to 12Z 06 Feb. 2008) of forecasts, every 12 hours is supplied to `gen_be_wrapper` to estimate forecast error covariance. It is only for demonstration. The minimum number of forecasts required depends on the application, number of grid points, etc. Month-long (or longer) datasets are typical for the NMC-method. Generally, at least a 1-month dataset should be used.

Under WRFDA/var/scripts/gen_be, `gen_be_wrapper.ksh` is used to generate the BE data. The following variables need to be set to fit your case:

```
export WRFVAR_DIR=/glade/work/wrfhelp/PRE_COMPILED_CODE/WRFDA
export NL_CV_OPTIONS=5      # 5 for CV5, 7 for CV7
export START_DATE=2008020612 # the first perturbation valid date
export END_DATE=2008020700  # the last perturbation valid date
export NUM_LEVELS=40       # e_vert - 1
export BIN_TYPE=5          # How data is binned for calculating statistics
export FC_DIR=/glade/work/wrfhelp/WRFDA_DATA/fc  # where wrf forecasts are
export RUN_DIR=`pwd`/gen_be # Where GEN_BE will run and output files
```

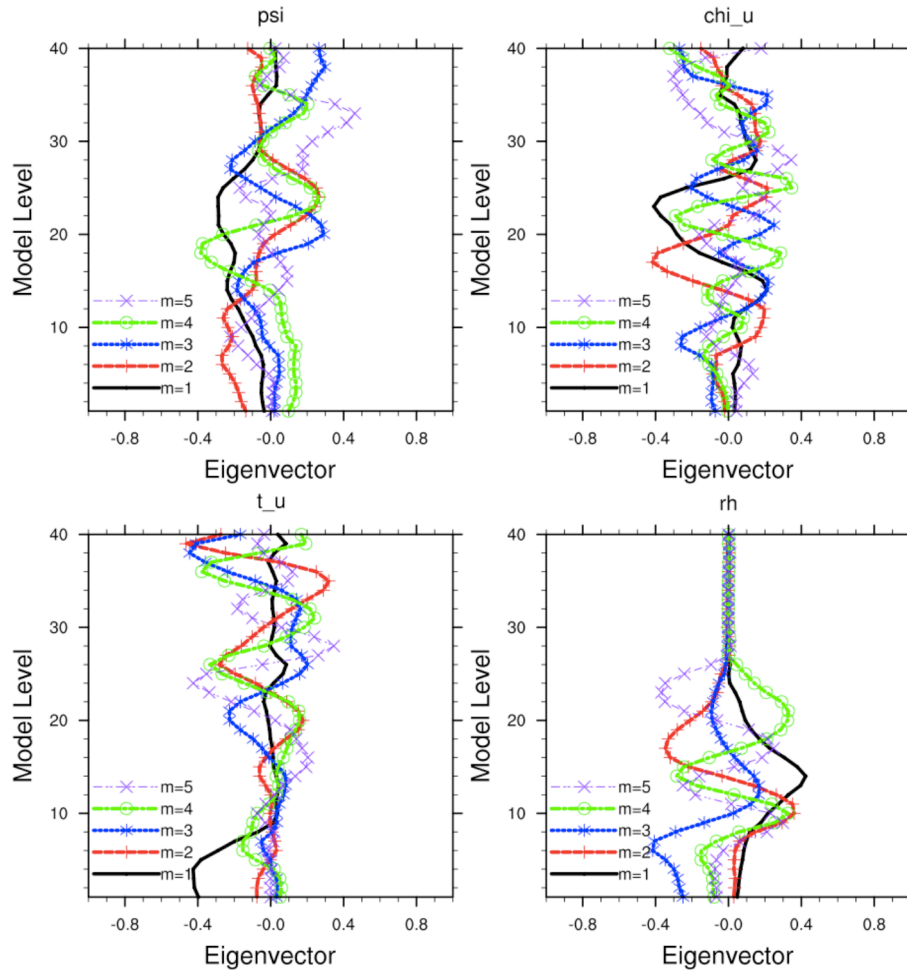
Note: The `START_DATE` and `END_DATE` are perturbation valid dates. As shown in the forecast list above, when you have 24-hour and 12-hour forecasts initialized at 2008020512, through 2008020612, the first and final forecast difference valid dates are 2008020612 and 2008020700, respectively.

Note: The forecast dataset should be located in `$FC_DIR`. Then type:

```
> gen_be_wrapper.ksh
```

Once the `gen_be_wrapper.ksh` run is completed, the `be.dat` can be found under the `$RUN_DIR` directory.

To get a clear idea about what is included in `be.dat`, the script `gen_be_plot_wrapper.ksh` may be used. This plots various data in `be.dat`; for example:



GEN_BE for CV6

CV6 is a multivariate background error statistics option in WRFDA. It may be activated by setting the namelist variable `cv_options=6`. This option introduces six additional correlation coefficients in the definition of the balanced part of analysis control variables. Thus with this implementation, moisture analysis is multivariate, in the sense that temperature and wind may lead to moisture increments, and vice-versa. Further details may be seen at: http://www2.mmm.ucar.edu/wrf/users/wrfda/Docs/WRFDA_updated_for_cv6.pdf

How to generate CV6 background error statistics for WRFDA

CV6 background error statistics for WRFDA are generated by executing a top-level script, `gen_be/wrapper_gen_mbe.ksh`, residing under `SCRIPTS_DIR` via a suitable wrapper script. The rest of the procedure remains the same as with normal running of the `gen_be` utility. A successful run will create a `be.dat` file in the `RUN_DIR` directory.

How to run WRFDA with CV6 background error statistics

After successfully generating the CV6 background error statistics file `be.dat`, the procedure for running WRFDA is straight-forward: Include `cv_options=6` in the `namelist.input` file under the `&wrfvar7` list of `namelist` options.

How to tune CV6 background error statistics

Below is a list of nine tuning parameters available in WRFDA; these can be specified under `&wrfvar7` in the `namelist`. Default values for these variables are set as “1.0”. Setting corresponding values > 1.0 (< 1.0) will increase (decrease) the corresponding contributions:

Variable name	Description
<code>psi_chi_factor</code>	Parameter to control contribution of stream function in defining balanced part of velocity potential
<code>psi_t_factor</code>	Parameter to control contribution of stream function in defining balanced part of temperature
<code>psi_ps_factor</code>	Parameter to control contribution of stream function in defining balanced part of surface pressure
<code>psi_rh_factor</code>	Parameter to control contribution of stream function in defining balanced part of moisture
<code>chi_u_t_factor</code>	Parameter to control contribution of unbalanced part of velocity potential in defining balanced part of temperature
<code>chi_u_ps_factor</code>	Parameter to control contribution of unbalanced part of velocity potential in defining balanced part of surface pressure
<code>chi_u_rh_factor</code>	Parameter to control contribution of unbalanced part of velocity potential in defining balanced part of moisture
<code>t_u_rh_factor</code>	Parameter to control contribution of unbalanced part of temperature in defining balanced part of moisture
<code>ps_u_rh_factor</code>	Parameter to control contribution of unbalanced part of surface pressure in defining balanced part of moisture

Additional Background Error options

a. Single Observation response in WRFDA

With the single observation test, you may get some ideas of how the background and observation error statistics work in the model variable space. A single observation test is done in WRFDA by setting `num_pseudo=1`, along with other pre-specified values in record `&wrfvar15` and `&wrfvar19` of `namelist.input`.

With the settings shown below, WRFDA generates a single observation with a pre-

specified *innovation* (Observation – First Guess) value at the desired location; e.g. at (in terms of grid coordinate) 23x23, level 14 for “U” observation with error characteristics 1 m/s, and innovation size = 1.0 m/s.

```
&wrfvar15
num_pseudo = 1,
pseudo_x = 23.0,
pseudo_y = 23.0,
pseudo_z = 14.0,
pseudo_err = 1.0,
pseudo_val = 1.0,
/
&wrfvar19
pseudo_var = 'u',
/
```

You may wish to repeat this exercise for other observation types. “pseudo_var” can take any of the following values:

Variable name	Description	Units
u	East-west wind	m/s
v	North-south wind	m/s
t	Temperature	K
p	Pressure	Pa
q	Water vapor mixing ratio	<i>unitless</i> (kg/kg)
qcw	Water vapor mixing ratio	<i>unitless</i> (kg/kg)
qrn	Rain water mixing ratio	<i>unitless</i> (kg/kg)
qci	Cloud ice mixing ratio	<i>unitless</i> (kg/kg)
qsn	Snow mixing ratio	<i>unitless</i> (kg/kg)
qgr	Graupel mixing ratio	<i>unitless</i> (kg/kg)
tpw	Total precipitable water	cm
ztd	GPS zenith total delay	cm
ref	GPS Refractivity	<i>Unitless</i>

b. Response of BE length scaling parameter

Run the single observation test with the following additional parameters in record &wrfvar7 of namelist.input.

```
&wrfvar7
len_scaling1 = 0.5, # reduce psi length scale by 50%
len_scaling2 = 0.5, # reduce chi_u length scale by 50%
len_scaling3 = 0.5, # reduce T length scale by 50%
len_scaling4 = 0.5, # reduce q length scale by 50%
len_scaling5 = 0.5, # reduce Ps length scale by 50%
/
```

Note: You may wish to try the response of an individual variable by setting one parameter at a time. Note the spread of analysis increment.

c. Response of changing BE variance

Run the single observation test with the following additional parameters in record `&wrfvar7` of `namelist.input`.

```
&wrfvar7
var_scaling1 = 0.25,    # reduce psi variance by 75%
var_scaling2 = 0.25,    # reduce chi_u variance by 75%
var_scaling3 = 0.25,    # reduce T variance by 75%
var_scaling4 = 0.25,    # reduce q variance by 75%
var_scaling5 = 0.25,    # reduce Ps variance by 75%
/
```

Note: You may wish to try the response of individual variable by setting one parameter at a time. Note the magnitude of analysis increments.

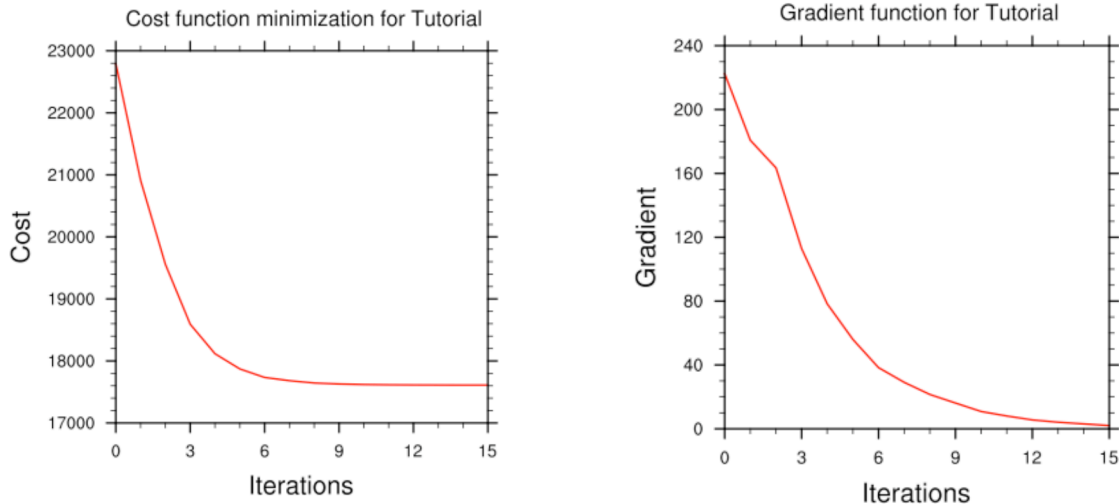
WRFDA Diagnostics

WRFDA produces a number of diagnostic files that contain useful information on how the data assimilation has performed. This section will introduce you to some of these files, and what to look for.

After running WRFDA, it is important to check a number of output files to see if the assimilation appears sensible. The WRFDA package, which includes several useful scripts, may be downloaded from <http://www2.mmm.ucar.edu/wrf/users/wrfda/download/tools.html>

The content of some useful diagnostic files are as follows:

`cost_fn` and `grad_fn`: These files hold (in ASCII format) WRFDA cost and gradient function values, respectively, for the first and last iterations. If you run with `calculate_cg_cost_fn=true` and `write_detail_grad_fn=true`, however, these values will be listed for each iteration; this can be helpful for visualization purposes. The NCL script in the WRFDA TOOLS package located at `var/graphics/ncl/plot_cost_grad_fn.ncl` may be used to plot the content of `cost_fn` and `grad_fn`, if these files are generated with `calculate_cg_cost_fn=true` and `write_detail_grad_fn=true`.



Note: Make sure that you remove the first two lines (header) in `cost_fn` and `grad_fn` before you plot. You also need to specify the directory name for these two files.

`gts_omb_oma_01`: It contains (in ASCII format) information on all of the observations used by the WRFDA run. Each observation has its observed value, quality flag, observation error, observation minus background (OMB), and observation minus analysis (OMA). This information is very useful for both analysis and forecast verification purposes.

`namelist.input`: This is the WRFDA input namelist file, which contains all the user-defined non-default options. Any namelist-defined options that do not appear in this file should have their names checked against the values in `$WRFDA_DIR/Registry/registry.var`.

`namelist.output.da`: A list of all the namelist options used. If an option was not specified in `namelist.input`, the default listed in the registry value will be used.

`rs1*`: Files containing information for standard WRFDA output from individual processors when multiple processors are used. It contains a host of information on a number of observations, minimization, timings, etc. Additional diagnostics may be printed in these files by including various “print” WRFDA namelist options. To learn more about these additional “print” options, search for the “print_” string in `$WRFDA_DIR/Registry/registry.var`.

`statistics`: Text file containing OMB (OI) and OMA (OA) statistics (minimum, maximum, mean and standard deviation) for each observation type and variable. This information is very useful in diagnosing how WRFDA has used different components of the observing system. Also contained are the analysis minus background (A-B) statistics, i.e. statistics of the analysis increments for each model variable at each model level. This information is very useful in checking the range of analysis increment values found in the analysis, and where they are in the WRF-model grid space.

The WRFDA analysis file is `wrfvar_output`. It is in WRF (netCDF) format. It will become the input file `wrfinput_d01` of any subsequent WRF run after lateral boundary and/or lower boundary conditions are updated by another WRFDA utility (See the section [Updating WRF boundary conditions](#)).

An NCL script, `$TOOLS_DIR/var/graphics/ncl/WRF-Var_plot.ncl`, is provided in the tools package for plotting. You need to specify the `analysis_file` name, its full path, etc. Please see the in-line comments in the script for details.

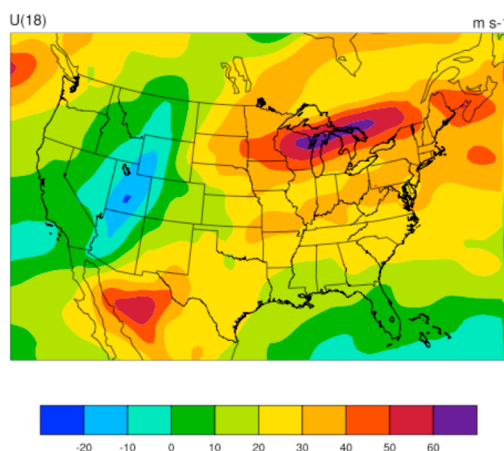
As an example, if you are aiming to display the U-component of the analysis at level 18, use the script `WRF-Var_plot.ncl`, and make sure the following pieces of codes are uncommented:

```
var = "U"
fg = first_guess->U
an = analysis->U
plot_data = an
```

When you execute the following command from `$WRFDA_DIR/var/graphics/ncl`.

```
> ncl WRF-Var_plot.ncl
```

The plot should look like:



You may change the variable name, level, etc. in this script to display the variable of your choice at the desired eta level.

Take time to look through the text output files to ensure you understand how WRFDA works. For example:

How closely has WRFDA fit individual observation types? Look at the `statistics` file to compare the O-B and O-A statistics.

How big are the analysis increments? Again, look in the `statistics` file to see

minimum/maximum values of A-B for each variable at various levels. It will give you a feel for the impact of the input observation data you assimilated via WRFDA by modifying the input analysis first guess.

How long did WRFDA take to converge? Does it really converge? You will get the answers of all these questions by looking into the `rs1.*` -files, as it indicates the number of iterations taken by WRFDA to converge. If this is the same as the maximum number of iterations specified in the namelist (`NTMAX`), or its default value (=200) set in `$WRFDA_DIR/Registry/registry.var`, then it means that the analysis solution did not converge. If this is the case, you may need to increase the value of “`NTMAX`” and rerun your case to ensure that the convergence is achieved. On the other hand, a normal WRFDA run should usually converge within 100 iterations. If it still doesn’t converge in 200 iterations, that means there may be a problem in the observations or first guess.

A good way to visualize the impact of assimilation of observations is to plot the analysis increments (i.e. analysis minus the first guess difference). Many different graphics packages (e.g. RIP4, NCL, GRADS etc) can do this.

You need to modify this script to fix the full path for `first_guess` and `analysis` files. You may also use it to modify the display level by setting `kl` and the name of the variable to display by setting `var`. Further details are given in this script.

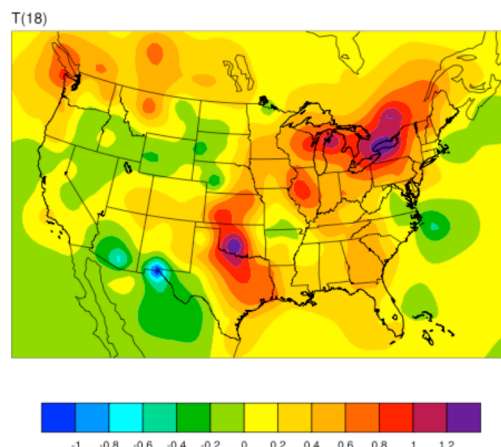
If you are aiming to display the increment of potential temperature at level 18, after modifying `$WRFDA_DIR/var/graphics/ncl/WRF-Var_plot.ncl`, make sure the following pieces of code are uncommented:

```
var = "T"
fg = first_guess->T ;Theta- 300
an = analysis->T    ;Theta- 300
plot_data = an - fg
```

When you execute the following command from `WRFDA_DIR/var/graphics/ncl`.

```
> ncl WRF-Var_plot.ncl
```

The plot created will look as follows:



Note: Larger analysis increments indicate a larger data impact in the corresponding region of the domain.

Generating ensembles with RANDOMCV

In addition to the variational methods previously mentioned, the WRFDA system supports both ensemble and hybrid ensemble/variational assimilation methods. To utilize these methods, having an ensemble of forecasts is necessary. WRFDA has a built-in method for generating ensemble initial conditions known as RANDOMCV. RANDOMCV works by adding random noise to the analysis in control variable space.

a. Running WRFDA for RANDOMCV

RANDOMCV is a capability of the main WRFDA executable, so you will run it by setting the following variables in `namelist.input` and then running `da_wrfvar.exe` as you would with any data assimilation run:

```
&wrfvar5
put_rand_seed  true:  enter your own seed numbers to generate random background perturbations.
                  The advantage of this setting is that the same seed numbers will always produce the
                  same perturbation
                  false: (default behavior) perturbations will be generated "randomly" and give new
                  perturbations each time

&wrfvar11
seed_array1    First integer for seeding the random function (default: 1)
seed_array2    Second integer for seeding the random function (default: 1) It is not necessary to
                  change both seeds to get different perturbations

&wrfvar17
analysis_type  Set this to 'RANDOMCV' to use the RANDOMCV capability
n_randomcv     1 (default), number of ensemble to be generated in one
                  run. New in version 4.2.
```

When setting your own random seeds, a common good practice is to set the first seed as the experiment date/time in integer form, and the second seed as the ensemble member number. *The seeds should not be set to zero.*

Because the perturbations are made in control variable space, the general pattern of perturbations will depend on your background error. You should be able to use any background error option with RANDOMCV (CV3, CV5, CV6, or CV7). Additionally, this means you can control their magnitude and lengthscales by modifying the background error variance and length scaling variables respectively:

For CV5, CV6, or CV7

```
&wrfvar7
var_scaling1 = 0.25, # reduce psi perturbation magnitude by 75%
var_scaling2 = 0.25, # reduce chi_u perturbation magnitude by 75%
var_scaling3 = 0.0,  # reduce T perturbation by 100% (there will be no
                    # T perturbation!)
var_scaling4 = 2.0,  # increase q perturbation by 100%
var_scaling5 = 1.0,  # Keep Ps perturbation magnitude the same
len_scaling1 = 0.5,  # reduce psi perturbation length scale by 50%
len_scaling2 = 0.5,  # reduce chi_u perturbation length scale by 50%
len_scaling3 = 1.0,  # Keep T perturbation length scale the same
len_scaling4 = 2.0,  # increase q perturbation length scale by 100%
len_scaling5 = 1.5,  # increase Ps perturbation length scale by 50%
/
```

For CV3, see the section “[Modifying CV3 length scales and variance](#)”.

Hybrid Data Assimilation in WRFDA

The WRFDA system also includes hybrid data assimilation techniques—both 3DVar and 4DVar, which is based on the previously-described variational capability.

The difference between hybrid and variational schemes is that WRFDA 3DVAR and 4DVAR rely solely on a static covariance model to specify the background errors, while the hybrid system uses a combination of static error covariances and ensemble-estimated error covariances to incorporate a flow-dependent estimate of the background error statistics. The following sections will give a brief introduction to using the hybrid system. Please refer to these papers for a detailed description of the methodology used in the WRFDA hybrid system:

Xuguang Wang, Dale M. Barker, Chris Snyder, and Thomas M. Hamill, 2008: [A hybrid ETKF-3DVAR data assimilation scheme for the WRF model. Part I: Observing system simulation experiment](#). *Mon. Wea. Rev.*, **136**, 5116–5131.

Xuguang Wang, Dale M. Barker, Chris Snyder, and Thomas M. Hamill, 2008: [A Hybrid ETKF–3DVAR Data Assimilation Scheme for the WRF Model. Part II: Real Observation Experiments](#). *Mon. Wea. Rev.*, **136**, 5132–5147.

Four executables are used in the hybrid system. If you have successfully compiled the WRFDA system, the following executables will exist in the WRFDA/var/build directory:

```
WRFDA/var/build/gen_be_ensmean.exe
WRFDA/var/build/gen_be_ep2.exe
WRFDA/var/build/da_wrfvar.exe
WRFDA/var/build/gen_be_vertloc.exe
```

`gen_be_ensmean.exe` is used to calculate the ensemble mean, while `gen_be_ep2.exe` is used to calculate the ensemble perturbations. `gen_be_vertloc.exe` is used for vertical localization. As with 3DVAR/4DVAR, `da_wrfvar.exe` is the main WRFDA program. However, in this case, `da_wrfvar.exe` will run in hybrid mode.

a. Running the hybrid system for 3DEnVar

The procedure is the same as running 3DVAR, with the exception of some extra input files and namelist settings. The basic input files for WRFDA are `LANDUSE.TBL`, `ob.ascii` or `ob.bufr` (depending on which observation format you use), and `be.dat` (static background errors). Additional input files required for 3DEnVar are a single ensemble mean file (used as the `fg` for the hybrid application) and a set of ensemble perturbation files (used to represent flow-dependent background errors).

A set of initial ensemble members must be prepared before the hybrid application can be started. The ensemble can be obtained from other ensemble model outputs, or you can generate them yourself. This can be done, for example, adding random noise to the initial conditions at a previous time and integrating each member to the desired time. A tutorial case with a test ensemble can be found at http://www2.mmm.ucar.edu/wrf/users/wrfda/download/V38/wrfda_hybrid_etkf_testdata.tar.gz. In this example, the ensemble forecasts were initialized at 2015102612 and valid 2015102712. A hybrid analysis at 2015102712 will be performed using the ensemble valid 2015102712 as input. Once you have the initial ensemble, the ensemble mean and perturbations can be calculated following the steps below:

- 1) Set an environment variable for your working directory and your data directory

```
> setenv WORK_DIR your_hybrid_path
> setenv DAT_DIR your_data_path
> cd $WORK_DIR
```

- 2) Calculate the ensemble mean

- a) From your working directory, copy or link the ensemble forecasts to your work-

ing directory. The ensemble members are identified by the characters “.e###” at the end of the file name, where ### represents three-digit numbers following the valid time.

```
> ln -sf $DAT_DIR/Hybrid/fc/2015102612/wrfout_d01_2015-10-27_12:00:00.e* .
```

- b) Provide two template files (ensemble mean and variance files) in your working directory. These files will be overwritten with the ensemble mean and variance as discussed below.

```
> cp $DAT_DIR/Hybrid/fc/2015102612/wrfout_d01_2015-10-27_12:00:00.e001 ./wrfout_d01_2015-10-27_12:00:00.mean
> cp $DAT_DIR/Hybrid/fc/2015102612/wrfout_d01_2015-10-27_12:00:00.e001 ./wrfout_d01_2015-10-27_12:00:00.vari
```

- c) Copy `gen_be_ensmean_nl.nl` (`cp $DAT_DIR/Hybrid/gen_be_ensmean_nl.nl .`) You will need to set the information in this script as follows:

```
&gen_be_ensmean_nl
directory = '.'
filename = 'wrfout_d01_2015-10-27_12:00:00'
num_members = 10
nv = 7
cv = 'U', 'V', 'W', 'PH', 'T', 'MU', 'QVAPOR'
/
```

where `directory` is the folder containing the ensemble members and template files, `filename` is the name of the files before their suffixes (e.g., `.mean`, `.vari`, etc), `num_members` is the number of ensemble members you are using, `nv` is the number of variables, and `cv` is the name of variables used in the hybrid system. Be sure `nv` and `cv` are consistent!

- d) Link `gen_be_ensmean.exe` to your working directory and run it.

```
> ln -sf $WRFDA_DIR/var/build/gen_be_ensmean.exe .
> ./gen_be_ensmean.exe
```

Check the output files. `wrfout_d01_2015-10-27_12:00:00.mean` is the ensemble mean; `wrfout_d01_2015-10-27_12:00:00.vari` is the ensemble variance

3) Calculate ensemble perturbations

- a) Create a sub-directory in which you will be working to create ensemble perturbations.

```
> mkdir -p ./ep
> cd ./ep
```

- b) Run `gen_be_ep2.exe`. The executable requires four command-line arguments

(DATE, NUM_MEMBER, DIRECTORY, and FILENAME) as shown below for the tutorial example:

```
> ln -sf $WRFDA_DIR/var/build/gen_be_ep2.exe .
> ./gen_be_ep2.exe 2015102712 10 . ../wrfout_d01_2015-10-27_12:00:00
```

- c) Check the output files. A list of binary files should now exist. Among them, `tmp.e*` are temporary scratch files that can be removed.

- 4) Back in the working directory, create the input file for vertical localization. This program requires one command-line argument: the number of vertical levels of the model configuration (same value as `e_vert` in the namelist; for the tutorial example, this should be 42).

```
> cd $WORK_DIR
> ln -sf $WRFDA_DIR/var/build/gen_be_vertloc.exe .
> ./gen_be_vertloc.exe 42
```

The output is `./be.vertloc.dat` in your working directory.

- 5) Run WRFDA in hybrid mode

- a) In your hybrid working directory, link all the necessary files and directories as follows:

```
> ln -fs ../wrfout_d01_2015-10-27_12:00:00.mean ./fg (first
guess is the ensemble mean for this test case)
> ln -fs $WRFDA_DIR/run/LANDUSE.TBL .
> ln -fs $DAT_DIR/Hybrid/ob/2015102712/ob.ascii ./ob.ascii
> ln -fs $DAT_DIR/Hybrid/be/be.dat ./be.dat
> ln -fs $WRFDA_DIR/var/build/da_wrfvar.exe .
> cp $DAT_DIR/Hybrid/namelist.input .
```

- b) Edit `namelist.input`, paying special attention to the following hybrid-related settings:

```
&wrfvar7
je_factor = 2.0
/
&wrfvar16
ensdim_alpha = 10
alphacv_method = 2
alpha_corr_type=3
alpha_corr_scale = 500.0
alpha_std_dev=1.000
alpha_vertloc_opt = 2
ep_format = 1
/
```

- c) Finally, execute the WRFDA file, running in hybrid mode

```
> ./da_wrfvar.exe >& wrfda.log
```

Check the output files; the output file lists are the same as when you run WRF 3D-Var.

b. Running the hybrid system for 4DEnVar

The procedure for 4DEnVar is very similar to the procedure for 3DEnVar. Unlike 4DVAR, 4DEnVar does not require WRFPLUS, the tangent linear/adjoint model. It is more analogous to the First Guess at Appropriate Time (FGAT) method than full 4DVAR. Like FGAT, 4DEnVar requires multiple first guess files, one for each assimilation time window. In addition, you must have ensemble output for each of the first guess times, and generate ensemble perturbations from each of these sets of ensemble forecasts.

To activate the 4DEnVar option, set the namelist variable `use_4denvar=.true.` under `&wrfvar16`. A tutorial case with a test ensemble can be found at http://www2.mmm.ucar.edu/wrf/users/wrfda/download/V39/WRFDA_4denvar_test_data.tar.gz. In this example, the ensemble forecasts were initialized at 2005071512 and valid for 2005071521—2005071603. The 4DEnVar hybrid analysis (`wrfvar_output`) will be valid for 2005071600.

c. Dual-resolution hybrid

WRFDA has an option for dual-resolution hybrid data assimilation, where a high-resolution background can make use of a lower-resolution ensemble for extracting the flow-dependent contribution to the background error. The lower-resolution ensemble should be the parent domain of the higher-resolution child domain that the analysis is performed on.

Performing dual-resolution hybrid assimilation is similar to the process for regular assimilation described above. The main difference is that you must include some settings in the `&domains` section of the namelist in a different way: The setting `max_dom=2` must be used, and for each column of settings, **the low-resolution domain settings must be listed first**, even though the second column will be the domain on which assimilation is being performed. An example of the `&domains` namelist section is shown below; in this case, the low-resolution ensemble domain is 222x128 grid points at 45 km resolution, and the high-resolution analysis domain is 184x196 grid points at 15 km resolution:

```
&domains
  time_step      = 90,
  max_dom        = 2,
  s_we           = 1,  1,
  e_we           = 222, 184,
  s_sn           = 1,  1,
  e_sn           = 128, 196,
  s_vert         = 1,  1,
  e_vert         = 45,  45,
  dx             = 45000, 15000,
  dy             = 45000, 15000,
  grid_id        = 1,  2,
  parent_id      = 0,  1,
```

```

i_parent_start      = 0, 89,
j_parent_start      = 0, 22,
parent_grid_ratio    = 1, 3,
/

```

For further details about any of the above settings, see [Chapter 5 of this User's Guide](#).

In addition to the above, you must set the namelist variable **hybrid_dual_res=true**, as well as providing a file named “fg_ens” in the working directory. This file can be any WRF input or wrfout file that has the same domain as the low-resolution ensemble; it is merely used for reading in mapping parameters for the low-resolution ensemble domain.

The dual-resolution hybrid capability is described in more detail in the following publication:

Schwartz, C. S., Z. Liu, X.-Y. Huang, 2015: [Sensitivity of Limited-Area Hybrid Variational-Ensemble Analyses and Forecasts to Ensemble Perturbation Resolution](#). *Mon. Wea. Rev.*, **143**, 3454-3477.

d. Hybrid namelist options

&wrfvar7

je_factor

ensemble covariance weighting factor. This factor controls the weighting component of ensemble and static covariances. The corresponding **jb_factor** = $je_factor/(je_factor - 1)$.

&wrfvar16

use_4denvar

.true. will activate 4DEnVar

hybrid_dual_res

.true. will activate dual-resolution mode

ep_para_read

Method for reading ensemble perturbation files

0 (default): Serial read

1: Parallel read.

rden_bin

bins for parallel reading of ensemble perturbation files. Default is 1. Lower numbers use more memory, but are faster. If memory use becomes too large, increase this value.

ensdim_alpha

the number of ensemble members. Hybrid mode is activated when **ensdim_alpha** is larger than zero

alphacv_method

2 (default): perturbations in model space (“u”, “v”, “t”, “q”, “ps”). Option 2 is extensively tested and recommended to use.

1: perturbations in control variable space (“psi”, “chi_u”, “t_u”, “rh”, “ps_u”)

alpha_corr_type

correlation function. 1=Exponential; 2=SOAR; 3=Gaussian.

alpha_corr_scale

hybrid covariance localization scale in km unit. Default value is 200.

alpha_std_dev

alpha standard deviation. Default value is 1.0

alpha_vertloc_opt (new in version 4.2)

0: no vertical localization

1: old option prior to version 4.2 (obsolete).

2: default, recommended, let WRFDA internally calculate logP-based vertical localization. `be.vertloc.dat` will be written out.

ep_format (new in version 4.2)

1: (default) original format, double precision, each ep file is for one variable and one member, as the output from the current `gen_be_ep2`.

11: same as `ep_format=1` except data are in single precision

2: single precision, each ep file is for one variable and all members.

3: single precision, each ep file is for one variable and all members but on decomposed patch domain. This is recommended for large domain.

Different format of ep files can be generated using `gen_be_v3` program (new in version 4.2). See `var/gen_be_v3/README.gen_be_v3` for more information.

ETKF Data Assimilation

The WRFDA system also includes a ETKF assimilation technique. The ETKF system updates the ensemble perturbations. Please refer to Bishop et al. (2001) and Wang et al. (2003) for a detailed description of the methodology. The following section will give a brief introduction of some aspects of using the ETKF system.

a. Source Code

Three executables are used in the ETKF system. If you have successfully compiled the WRFDA system, you will see the following:

```
WRFDA/var/build/gen_be_etkf.exe
```

```
WRFDA/var/build/gen_be_addmean.exe
```

```
WRFDA/var/build/da_wrfvar.exe
```

The file `gen_be_etkf.exe` is used to update the ensemble perturbations, while `gen_be_addmean.exe` is used to combine the ensemble mean and the ensemble perturbations. As with 3D-Var/4D-Var, `da_wrfvar.exe` is the main WRFDA program. However, in this case, `da_wrfvar.exe` will create filtered observations and prepare formatted omb files for ETKF.

b. Running the ETKF System

The first procedure is to update the ensemble perturbations. A set of initial ensemble members must be prepared before the ETKF application can be started. The ensemble can be obtained from a previous ensemble forecast. A tutorial case with a test ensemble can be found at

http://www2.mmm.ucar.edu/wrf/users/wrfda/download/V38/wrfda_hybrid_etkf_testdata.tar.gz. In this example, the ensemble forecasts were initialized at 2015102612 and valid 2015102712. ETKF will be performed using the ensemble valid 2015102712 as input. Once you have the initial ensemble, the ensemble perturbations can be updated by following the steps below:

1) Set environment variables for convenience

```
> setenv WORK_DIR_ETKF your_etkf_path
> setenv DAT_DIR your_data_path
> setenv WRFDA_DIR your_WRFDA_path
> cd $WORK_DIR_ETKF
```

2) Prepare the filtered observations

- a) In your ETKF working directory, make a subdirectory to prepare the filtered observations and link all the necessary files and directories as follows:

```
> mkdir obs_filter
> cd obs_filter
> ln -fs $DAT_DIR/Hybrid/fc/2015102612/wrfout_d01_2015-10-
27_12:00:00.mean ./fg (first guess is the ensemble mean)
> ln -fs $WRFDA_DIR/run/LANDUSE.TBL .
> ln -fs $DAT_DIR/Hybrid/ob/2015102712/ob.ascii ./ob.ascii
> ln -fs $DAT_DIR/Hybrid/be/be.dat ./be.dat
> ln -fs $WRFDA_DIR/var/build/da_wrfvar.exe .
> cp $DAT_DIR/ETKF/namelist.input .
```

- b) Edit namelist.input, paying special attention to the following 'QC-OBS'-related settings:

```
&wrfvar17
analysis_type = 'QC-OBS',
/
```

- c) Execute the WRFDA file, running in QC-OBS mode

```
> ./da_wrfvar.exe >& wrfda.log
```

Check the output files; you should see a 'filtered_obs_01' file which contains the filtered observations.

3) Prepare omb files for ETKF

a) In your ETKF working directory, make a subdirectory to prepare the omb files for each ensemble member and link all the necessary files and directories as follows:

```
> cd $WORK_DIR_ETKF
> mkdir -p omb/working.e001
> cd omb/working.e001
> ln -fs $DAT_DIR/Hybrid/fc/2015102612/wrfout_d01_2015-10-
27_12:00:00.e001 ./fg (first guess is the ensemble member)
> ln -fs $WRFDA_DIR/run/LANDUSE.TBL .
> ln -fs $WORK_DIR_ETKF/obs_filter/filtered_obs_01 ./ob.ascii
> ln -fs $DAT_DIR/Hybrid/be/be.dat ./be.dat
> ln -fs $WRFDA_DIR/var/build/da_wrfvar.exe .
> cp $DAT_DIR/ETKF/namelist.input .
```

b) Edit namelist.input, paying special attention to the following 'VERIFY'-related settings:

```
&wrfvar17
analysis_type                = 'VERIFY',
/
```

c) Execute the WRFDA file, running in VERIFY mode

```
> ./da_wrfvar.exe >& wrfda.log
```

Check the output files. The output files are the same as when you run WRF 3D-Var (except wrfvar_output will NOT be created), and the 'ob.etkf.0*' files are omb files.

d) Combine the ob.etkf.0* files and add the observation number in the head of ob.etkf.e0*

```
> cat ob.etkf.0* > ob.all
> wc -l ob.all > ob.etkf.e001
> cat ob.all >> ob.etkf.e001
```

e) Likewise, prepare ob.etkf.e0* files for other ensemble members

4) Run ETKF

a) Copy or link the ensemble mean and forecasts and ob.etkf.e0* files to your working directory and make a parameter directory to save the parameter files.

```
> cd $WORK_DIR_ETKF
> setenv PAR_DIR_ETKF $WORK_DIR_ETKF/param
> mkdir $PAR_DIR_ETKF
> ln -sf $DAT_DIR/Hybrid/fc/2015102612/wrfout_d01_2015-10-
27_12:00:00.mean ./etkf_input
> ln -sf $DAT_DIR/Hybrid/fc/2015102612/wrfout_d01_2015-10-
27_12:00:00.e001 ./etkf_input.e001
...
> ln -sf $DAT_DIR/Hybrid/fc/2015102712/wrfout_d01_2015-10-
```

```
27_12:00:00.e010 ./etkf_input.e010
```

```
> ln -sf omb/working.e001/ob.etkf.e001 .
...
> ln -sf omb/working.e010/ob.etkf.e010 .
```

b) Provide template files. These files will be overwritten with the ensemble perturbations.

```
> cp $DAT_DIR/Hybrid/fc/2015102612/wrfout_d01_2015-10-
27_12:00:00.e001 ./etkf_output.e001
...
> cp $DAT_DIR/Hybrid/fc/2015102612/wrfout_d01_2015-10-
27_12:00:00.e010 ./etkf_output.e010
```

c) Copy `gen_be_etkf_nl.nl` (`cp $DAT_DIR/ETKF/gen_be_etkf_nl.nl .`)
You will need to set the information in this script as follows:

```
&gen_be_etkf_nl
  num_members = 10,
  nv = 7,
  cv = 'U', 'V', 'W', 'PH', 'T', 'QVAPOR', 'MU',
  naccumt1 = 20,
  naccumt2 = 20,
  nstartaccum1 = 1,
  nstartaccum2 = 1,
  nout = 1,
  tainflatinput = 1,
  rhoinput = 1,
  infl_fac_file = '$PAR_DIR_ETKF/inflation_factor.dat',
  infl_let_file = '$PAR_DIR_ETKF/inflation_letkf.dat',
  eigen_val_file = '$PAR_DIR_ETKF/eigen_value.dat',
  inno2_val_file = '$PAR_DIR_ETKF/innovation_value.dat',
  proj2_val_file = '$PAR_DIR_ETKF/projection_value.dat',
  infl_fac_TRNK = .false.,
  infl_fac_WG03 = .false.,
  infl_fac_WG07 = .true.,
  infl_fac_BOWL = .false.,
  letkf_flg=.false.,
  rand_filt = .false.,
  rnd_seed = 2015102712,
  rnd_nobs = 5000
  etkf_erro_max = 20.
  etkf_erro_min = .00001
  etkf_inno_max = 20.
  etkf_inno_min = .00001
  etkf_erro_flg = .true.
  etkf_inno_flg = .true.
  etkf_wrfda = .false.
/
```

Important note: since environment variables are not parsed when reading namelists, you MUST manually change `$PAR_DIR_ETKF` to its actual value in the namelist

Where the various namelist parameters are as follows:

- `num_members` is the ensemble members size
- `nv` is the number of variables
- `cv` the name of variables
- `naccumt1` and `naccumt2` are number of previous cycles used to accumulate for inflation and rho factor
- `nstartaccumt1` and `nstartaccumt2` are not used for ordinary ETKF
- `nout` is the cycle index
- `tainflatinput` and `rhoinput` are prescribed factors for inflation and rho factor
- `infl_fac_file`, `eigen_val_file`, `inno2_val_file` and `proj2_val_file` are files to save template parameters
- `infl_fac_TRNK`, `infl_fac_WG03`, `infl_fac_WG07`, and `infl_fac_BOWL` are options for different adaptive inflation schemes
- `rand_filt`, `rnd_seed` and `rnd_nobs` are options for using filtered observation and random observations
- `etkf_erro_max`, `etkf_erro_min`, `etkf_inno_max`, `etkf_inno_min`, `etkf_erro_flg`, `etkf_inno_flg`, and `etkf_wrfda` are options to conduct further observation filtering.

d) Link `gen_be_etkf.exe` to your working directory and run it.

```
> ln -sf $WRFDA_DIR/var/build/gen_be_etkf.exe .  
> ./gen_be_etkf.exe
```

Check the output files. `etkf_output.*` files are updated ensemble perturbations.

5) Add updated ensemble perturbations to the ensemble mean to get new ensemble members

```
> cd $WORK_DIR_ETKF
```

a) Copy `add_mean_nl.nl` (`cp $DAT_DIR/ETKF/add_mean_nl.nl .`)

You will need to set the information in this script as follows for each member:

```
&add_mean_nl  
num_members = 10  
cv           = 'U', 'V', 'W', 'PH', 'T', 'QVAPOR', 'MU'  
nv           = 7  
path         = '$WORK_DIR_ETKF'  
file_mean    = 'etkf_input'  
file_pert    = 'etkf_output.e001' (for each member,  
etkf_output.e0*...)  
/  

```

Again, be sure to substitute the actual path in the place of `$WORK_DIR_ETKF`

b) Run `gen_be_addmean.exe`.

```
> ln -sf $WRFDA_DIR/var/build/gen_be_addmean.exe .
> ./gen_be_addmean.exe
```

Check the output files. `etkf_output.e0*` files are the new ensemble members.

Multi-Resolution Incremental 4DVAR

Prior to version 4.3, WRFDA's 4DVAR analysis can only be run at the same horizontal resolution as that of running the WRF model forecast. This limits 4DVAR's application at high resolution due to its high computational demand. From version 4.3, an option of running incremental 4DVAR analysis (i.e., iterative minimization process) at a lower resolution than that for the model forecast is available to speed up 4DVAR and the stand-alone interpolation utility is also provided to obtain high-resolution analysis from the low-resolution analysis. With multiple outer loops, incremental 4DVAR analysis can have gradually increased minimization resolutions for different outer loops. This new 4DVAR capability is referred to as the Multi-Resolution Incremental 4DVAR (MRI-4DVAR) and its implementation details are documented by Liu et al. (2020).

Liu, Z., J. Ban, J.-S. Hong, and Y.-H. Kuo, 2020: Multi-resolution incremental 4D-Var for WRF: Implementation and application at convective scale, *Q. J. R. Meteorol. Soc.*, 146, 3661-3674.

The standard 4DVAR with multiple outer loops (i.e., outer loop and inner loop are at the same resolution) is a non-stop (i.e., executable runs in memory without disk I/O) procedure even with multiple outer loops. However, the technical implementation of MRI-4DVAR is a three-stage procedure for each outer loop and the three stages are linked together through stand-alone programs and scripts. The three-stage procedure is described by Liu et al. (2020) and is also given below for users' quick reference.

Stage-1: Observer. This stage runs WRFDA to call WRF nonlinear model (NLM) to produce the model trajectory (initialized from the first guess x_g) at the full model resolution of running WRF forecast, compute the model trajectory departure to observations within the time window, and perform the quality control. At the end of this stage, the departure information along with QC flags is written out to separate files for different observation types and different time slots. Note that this stage is always run at the model forecast resolution (i.e., outer-loop resolution) for every outer loop and the appropriate high-resolution terrain field shall be used. Namelist setting for running WRFDA in observer mode is given below.

```
&wrfvar1
var4d    = true,
var4d_bin = 3600,
multi_inc = 1,
/
```


The key parameter is ‘multi_inc’, which is zero by default (i.e., turning off multi-resolution option) for a standard 4DVAR run. Setting multi_inc = 1 means that running WRFDA in observer mode (i.e., stage-1) for the multi-resolution option. Note that the setting above is for MRI-4DVAR, but the multi-resolution capability is also working for 3DVAR (i.e., MRI-3DVAR). However, usually there is no need to run 3DVAR in this multi-resolution setting because of low computational demand of 3DVAR. Also note that in observer mode, minimization will not be triggered even if namelist parameter ‘ntmax’ is set to >0.

Stage-2: Minimizer. This stage runs WRFDA for 4DVAR inner-loop minimization, which can be at the same or (more typically) at a lower resolution than that for observer stage. The observation-space departure files (output from Stage-1) will be read in. The model trajectory at inner-loop resolution is obtained by integrating WRF NLM from the first guess at the corresponding resolution and serves as the base state for the integration of WRF TLM/ADM during each inner iteration. Also be cautious that the corresponding inner-loop resolution model terrain should be used for the integration of TLM and ADM in this stage. At the end of stage-2 after minimization, WRFDA will write out the model space analysis (the normal WRFDA analysis file) and the “control variable space” analysis increment at the current inner-loop resolution. In this stage, ‘multi_inc’ needs to be set to 2 (i.e., stage-2) and a new namelist parameter **use_inverse_squarerootb** needs to be set to true.

```
&wrfvar1
var4d      = true,
var4d_bin  = 3600,
multi_inc  = 2,
/
&wrfvar6
max_ext_its = 1,
ntmax       = 20,
eps         = 0.01,
use_inverse_squarerootb = true,
/
```

Note that ‘use_inverse_squarerootb = true’ works only with cv_options=5, 6, 7, but not for cv_options=3.

Stage-3: Regridder. This stage prepares the model space and control variable space first guess for the next outer loop, which can use a higher inner-loop resolution. Two updated model space first-guess files are needed with one at the outer loop resolution and another at the inner-loop resolution. The former (the latter) will be used in Stage-1 (Stage-2) of the next outer loop. Both first-guess files are produced using an offline program **da_bilin.f90** that interpolates the model space analysis increment (from low to high resolution) and then adds the increment to the first-guess files from the previous outer loop. The control variable space first guess needs also to be interpolated (using another offline program **da_vp_bilin.f90**) to a higher inner-loop resolution, which will be read in by WRFDA in Stage-2 of the next outer loop. It is worth mentioning that some or all of three

regriddings (for two model space first guess and one control variable space first guess) can be skipped if there is no resolution difference between the inner loops or between inner loop and outer loop.

Stages-1/2/3 can be repeated for multiple outer loops with different inner-loop resolutions. A sample script file **run_mri3d4dvar.csh_pbs** under `~var/mri4dvar` is provided to integrate these 3-stage procedures. However, the script is provided only for helping the understanding of procedures described above. With the complexity of the procedures, it is not possible for us to provide the assistance for users' own applications of MRI-4DVAR without funding support. A readme file **README.MRI-4DVar** in the same directory provides some more information about compiling stand-alone utilities and the requirement for the grid size of high and low resolutions. Note that only a ratio 3 configuration between low and high resolution has been tested and other configurations of resolution ratio might not work properly without code modification.

Aerosol/Chemical Data Assimilation

Aerosol/Chemical data assimilation capability is added in version 4.3. Currently it allows the assimilation of 6 types of surface measurements (PM2.5, PM10, O3, CO, NO2, SO2) with 3DVAR, which can provide the analyzed initial conditions for WRF-Chem. This new capability is documented by Sun et al. (2020).

Sun, W., Liu, Z., Chen, D., Zhao, P., and Chen, M., 2020: Development and application of the WRFDA-Chem three-dimensional variational (3DVAR) system: aiming to improve air quality forecasting and diagnose model deficiencies, *Atmos. Chem. Phys.*, 20, 9311-9329.

Here we give some information about how to compile and configure WRFDA for this capability. The four gas-phase analysis variables include O3, CO, NO2 and SO2. For aerosols, it works with either GOCART or MOSAIC (4 bin) aerosol scheme in WRF-Chem, consisting of either 15 or 32 species, respectively.

To compile 3DVAR with aerosol/chemical DA capability, following the commands below:

```
setenv WRF_CHEM 1
./configure wrfda
./compile all_wrfvar
```

The executable is still `da_wrfvar.exe`, but now it can take WRF-Chem's initial condition file as the background (i. e., the 'fg' file) and perform the aerosol/chemical analysis in addition to the usual function for meteorological analysis. All related settings are in a new namelist section named **wrfvarchem**. Each namelist parameter is explained below.

&wrfvarchem

chem_cv_options= 10/20, control variables are 4 gas + GOCART or MOSAIC aerosol variables.
Default is 0, no aerosol/chemical DA.

chemicda_opt=1/2/3/4/5, assimilate pm2.5/pm10/pm2.5+pm10/co+o3+so2+no2/6-species.
Default is 1.

var_scaling12=1.0, 1.0, ..., array with a dimension of 36*max_outer_iterations, inflation factors of chemical background error variances of aerosol/gas analysis variables at different outer loops. Default values are 1 (no inflation).

len_scaling12=1.0, 1.0, ..., array with a dimension of 36*max_outer_iterations, inflation factors of chemical background error length scales. Default values is 1.

use_chemic_surfbobs = .true., Default is true, read in **ob_chemsfc.ascii** file that includes 6 types of observations at analysis time. Please refer to var/da/da_obs_io/da_read_obs_chem_sfc.inc for format.
/

The standard GEN_BE package included in WRFDA cannot obtain the background error statistics for aerosol/chemical variables, users will need to use https://github.com/wrf-model/GENBE_2.0 to generate a be.dat file that contains the background errors of chemical variables. Please refer to the online commit message for its use.

A testcase is provided in https://www2.mmm.ucar.edu/wrf/users/wrfda/chemda_testcase/. We have no resource to provide the assistance of this capability for users' own applications without funding support.

Additional WRFDA Options

1. Wind speed/direction assimilation

If observations containing wind speed/direction information are provided to WRFDA, you can assimilate these observations directly, rather than converting the wind to its u- and v-components prior to assimilation.

Wind speed/direction assimilation is controlled by the following namelist options:

```
&wrfvar2
wind_sd           true: all wind values which are reported as speed/direction will be assimilated as
                  such
                  false: (default behavior) all wind obs are converted to u/v prior to assimilation
wind_stats_sd     Assimilate wind in u/v format, but output speed/direction statistics
```

The following settings only matter if `check_max_iv=true` (if innovation is greater than observation error times the error factor listed below, the observation will be rejected):

```
&wrfvar2
qc_rej_both       true: if either u or v (spd or dir) do not pass quality control, both obs are rejected
                  false: (default behavior) qc on wind obs is handled individually
```

```

&wrfvar5
max_omb_spd      Max absolute value of innovation for wind speed obs in m/s; greater than this and the
                  innovation will be set to zero (default: 100.0)
max_omb_dir      Max absolute value of innovation for wind direction obs in degrees; greater than this
                  and the innovation will be set to zero (default: 1000.0)
max_error_spd    Speed error factor (default: 5.0)
max_error_dir    Direction error factor (default: 5.0)

```

The assimilation of wind speed/direction can also be controlled by observation type, using the following variables (note: setting `wind_sd = .true.` as above will override these individual settings):

```

&wrfvar2
wind_sd_airep    Aircraft reports
wind_sd_buoy     Buoy reports
wind_sd_geoamv   Geostationary satellite atmospheric motion vectors
wind_sd_metar    METAR reports
wind_sd_mtgirs   Meteosat Third Generation
wind_sd_pilot    Pilot reports
wind_sd_polaramv Polar satellite atmospheric motion vectors
wind_sd_profiler Wind profiler reports
wind_sd_qscat    QuikScat reports
wind_sd_ships    Ship reports
wind_sd_sound    Sounding reports
wind_sd_synop    Synoptic reports
wind_sd_tamdar   TAMDAR reports

```

true: wind values which are reported as speed/direction will be assimilated as such
 false: (default behavior) all wind obs are converted to u/v prior to assimilation

Further details about this method can be found in the following publications:

Huang, X.-Y., F. Gao, N. A. Jacobs, and H. Wang, 2013: [Assimilation of wind speed and direction observations: a new formulation and results from idealised experiments](#). *Tellus A*, **65**, 19936, doi:10.3402/tellusa.v65i0.19936.

Gao, Feng, Xiang-Yu Huang, Neil A. Jacobs, and Hongli Wang, 2015: [Assimilation of wind speed and direction observations: results from real observation experiments](#). *Tellus A*, **67**, 27132, doi:10.3402/tellusa.v67.27132.

2. The Weak Penalty Constraint option

A “weak penalty constraint” (WPEC) option is included in WRFDA, which aims to enforce quasi-gradient balance on a WRFDA analysis. It was designed with the specific aim of improving assimilation of radar data within tropical cyclones, but may be useful for other weather phenomena of similar scales. It can be used with 3DVAR or hybrid 3DVAR (4DVAR is not compatible with this new capability).

Further details about this method can be found in the following publication:

Li, X., J. Ming, M. Xue, Y. Wang, and K. Zhao, 2015: [Implementation of a dynamic equation constraint based on the steady state momentum equations within the WRF hybrid ensemble-3DVar data assimilation system and test with radar T-TREC wind assimilation for tropical Cyclone Chanthu \(2010\)](#). *J. Geophys. Res. Atmos.*, 120, 4017–4039, doi: 10.1002/2014JD022706.

This new option is controlled by the following set of namelist options:

```
&wrfvar12
use_wpec           true: enables the constraint term
                   false: (default behavior) disables the constraint term
wpec_factor        The constraint's weighting factor (1/Γ) as described in the paper
balance_type       1 = geostrophic term
                   2 = cyclostrophic term
                   3 = geostrophic + cyclostrophic terms (default; recommended)
```

3. Options for improving surface data assimilation

There are a number of options in WRFDA that are specifically for surface observations (e.g. METAR, SYNOP). Surface observations should be handled especially cautiously, as their impact can vary widely based on vertical and horizontal resolution, as well as other factors. Adjusting the options listed below can help investigate the assimilation of surface observations, especially in mountainous terrain.

```
&wrfvar11
These two options work for BUOY, METAR, SHIP, and SYNOP observations, as well as surface-
level sounding and TAMDAR observations.
sfc_assi_options    1 (default): surface observations will be assimilated based on the lowest
                   model level first guess. Observations are not used when the elevation dif-
                   ference between the observing site and the lowest model level is larger
                   than max_sheight_diff
                   2: surface observations will be assimilated based on surface similarity
                   theory in PBL. Innovations are computed based on 10-m wind, 2-m tem-
                   perature and 2-m moisture.
max_sheight_diff    Stations whose model-interpolated elevation is different from the actual
                   observation elevation by greater than this value (default: 100.0) in meters
                   will be rejected.
```

The following options apply only for SYNOP observations

```
sfc_hori_intp_options  Specifies the method of interpolating the background to observation space
                   1 (default): 4-point interpolation
                   2: Chooses values from neighboring model gridpoint with smallest height
                   difference (among land points: grid points over water will not be used).
obs_err_inflate       false (default): Observation error will be used as specified from observa-
                   tion files
                   true: Inflate observation error values by a factor of
                    $e^{(|Z_{diff}|/stn\_ht\_diff\_scale)}$ 
stn_ht_diff_scale      If obs_err_inflate=true, observation error will be inflated by a
                   factor of  $e^{(|Z_{diff}|/stn\_ht\_diff\_scale)}$ . Default is 200.0
```

4. Divergence constraint (DIVC)

A constraint term DIVC was added to model the correlation between u and v . The DIVC was implemented by adding a term in the cost function that constrains the horizontal divergence. For the detailed formulation of the new term, the user is referred to Tong et al. (2016) (<https://journals.ametsoc.org/doi/pdf/10.1175/WAF-D-16-0041.1>). The DIVC is evoked by setting `use_divc = .true.`. The effect of the DIVC is controlled by the value of the parameter `divc_factor`. The greater this value is, the smaller effect the DIVC has. Our recent experience showed that a value between 5000 and 10000 produced appropriate results, which did not significantly alter the pattern of precipitation while noise was eliminated.

5. Large Scale Analysis Constraint (LSAC)

The purpose of LSAC is to ensure that the convective-scale analysis does not distort the underlining large-scale balance and to eliminate possible large-scale bias in WRF background. The global analysis or forecast, such as that from GFS or FNL, is treated as bogus observations and assimilated via WRF 3DVAR. The user can choose the scale of the large-scale to be enforced by skipping grid points both in the horizontal and vertical directions (see below). For more detailed description of the LSAC, the user is referred to Vendrasco et al. (2016) (<https://journals.ametsoc.org/doi/pdf/10.1175/JAMC-D-15-0010.1>). The input data for LSAC is prepared using WPS, the same as preparing WRF input data by downscaling. The options related to LSAC are explained below:

<code>use_lsac</code>	True: use the large scale analysis constraint
<code>lsac_nh_step</code>	Number of increment steps in grid points in the horizontal direction
<code>lsac_nv_step</code>	Number of increment steps in grid points in the vertical direction
<code>lsac_nv_start</code>	Vertical level index to start LSAC (skip the levels below)
<code>lsac_use_u</code>	True: use data from large scale u analysis
<code>lsac_use_v</code>	True: use data from large scale v analysis
<code>lsac_use_t</code>	True: use data from large scale t analysis
<code>lsac_use_q</code>	True: use data from large scale qv analysis
<code>lsac_u_error</code>	Error in m/s of large-scale u analysis
<code>lsac_v_error</code>	Error in m/s of large-scale v analysis
<code>lsac_t_error</code>	Error in degree C of large-scale T analysis
<code>lsac_q_error</code>	Error in kg/kg of large-scale qv analysis
<code>lsac_print_details</code>	Switch for printout

Description of Namelist Variables

a. WRFDA namelist variables

&wrfvar1

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
write_increments	false	.true.: write out a binary analysis increment file
var4d	false	.true.: 4D-Var mode
var4d_lbc	true	.true.: on/off for lateral boundary control in 4D-Var
var4d_bin	3600	seconds, observation sub-window length for 4D-Var
var4d_bin_rain	3600	seconds, precipitation observation sub-window length for 4D-Var
multi_inc	0	> 0: multi-incremental run
print_detail_radar	false	print_detail_xxx: output extra (sometimes can be too many) diagnostics for debugging; not recommended to turn these on for production runs
print_detail_xa	false	
print_detail_xb	false	
print_detail_obs	false	
print_detail_grad	false	.true.: to print out a detailed gradient of each observation type at each iteration
check_max_iv_print	true	obsolete (used only by Radar)
update_sfc_diags	false	.true.: update T2/Q2/U10/V10/TH2 with WRFDA re-diagnosed values. Use only with sf_sfclay_physics=91 in WRF

&wrfvar2

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
analysis_accu	900	in seconds: if the time difference between the namelist date (analysis_date) and date info read-in from the first guess is larger than analysis_accu, WRFDA will abort.
calc_w_increment	false	.true.: the increment of the vertical velocity, W, will be diagnosed based on the increments of other fields. .false.: the increment of the vertical velocity W is zero if no W information is assimilated. If there is information on the W from observations assimilated, such as radar radial velocity, the W increments are always computed, whether calc_w_increment=true. or .false.
wind_sd	false	true: wind values which are reported as speed/direction will be assimilated as such false: (default behavior) all wind obs are converted to u/v prior to assimilation
qc_rej_both	false	true: if either u or v (spd or dir) do not pass quality

control, both obs are rejected
false: (default behavior) qc on wind obs is handled individually

&wrfvar3

Variable Name

Default Value

Description

fg_format

1

1: fg_format_wrf_arw_regional (default)
3: fg_format_wrf_arw_global
4: fg_format_kma_global
Options 3 and 4 are untested; use with caution!

ob_format

2

1: read in NCEP PREPBUFR data from ob.bufr
2: read in data from ob.ascii (default)

ob_format_gpsro

2

1: read in GPSRO data from gpsro.bufr
2: read in GPSRO data from ob.ascii (default)

num_fgat_time

1

1: 3DVar
> 1: number of time slots for FGAT and 4DVAR

&wrfvar4

Variable Name

Default Value

Description

thin_conv

true

Turns on observation thinning for ob_format=1 (NCEP PREPBUFR) only. thin_conv can be set to .false., but this is not recommended.

thin_conv_ascii

false

Turns on observation thinning for ob_format=2 (ASCII from OBSPROC) only.

thin_mesh_conv

20.

km, each observation type can set its thinning mesh (max_instrument and the index/order follows the definition in ts)

use_synopobs

true

WRFDA/var/da/da_control/da_control.f90

use_XXXobs - .true.: assimilate XXX obs if available
.false.: do not assimilate XXX obs even available

use_shipsobs

true

use_metarobs

true

use_soundobs

true

use_pilotobs

true

use_airepobs

true

use_geoamvobs

true

use_polaramvobs

true

use_bogusobs

true

use_buoyobs

true

use_profilerobs

true

use_satemobs

true

use_gpsspobs

true

use_gpsztdobs

false

Note: unlike most use_*obs variables, the default for use_gpsztdobs is false. This is because PW and ZTD observations can not be assimilated simultaneously, so one of them must be false.

use_gpsrefobs	true	
use_qscatobs	true	
use_radarobs	false	.true.: Assimilate radar data
use_radar_rv	false	Assimilate radar velocity observations
use_radar_rf	false	Assimilate radar reflectivity using original reflectivity operator (total mixing ratio)
use_radar_rhv	false	Assimilate retrieved hydrometeors (qr, qs, qg) from radar reflectivity
use_radar_rqv	false	Assimilate estimated humidity (qv) from radar reflectivity
use_rainobs	false	.true.: Assimilate precipitation data
thin_rainobs	true	.true.: perform thinning on precipitation data
use_airsretobs	true	
; use_hirs2obs, use_hirs3obs, use_hirs4obs, use_mhsobs, use_msuobs,		
; use_amsuaobs, use_amsubobs, use_airsobs, use_eos_amsuaobs, use_ssmisobs are		
; radiance-related variables that only control if corresponding BUFR files are read		
; into WRFDA or not, but do not control if the data is assimilated or not. Additional		
; variables have to be set in &wrfvar14 in order to assimilate radiance data.		
use_hirs2obs	false	.true.: read in data from hirs2.bufr
use_hirs3obs	false	.true.: read in data from hirs3.bufr
use_hirs4obs	false	.true.: read in data from hirs4.bufr
use_mhsobs	false	.true.: read in data from mhs.bufr
use_msuobs	false	.true.: read in data from msu.bufr
use_amsuaobs	false	.true.: read in data from amsua.bufr
use_amsubobs	false	.true.: read in data from amsub.bufr
use_airsobs	false	.true.: read in data from airs.bufr
use_eos_amsuaobs	false	.true.: read in data EOS-AMSUA data from airs.bufr
use_ssmisobs	false	.true.: to read in data from ssmis.bufr
use_atmsobs	false	.true.: to read in data from atms.bufr
use_iasioobs	false	.true.: to read in data from iasi.bufr
use_seviriobs	false	.true.: to read in data from seviri.bufr
use_amsr2obs	false	.true.: to read in data from AMSR2 files (see section “Other data formats” for file names)
use_obs_errfac	false	.true.: apply obs error tuning factors if errfac.dat is available for conventional data only

&wrfvar5

Variable Name

Default Value

Description

check_max_iv	true	.true.: reject the observations whose innovations (O-B) are larger than a maximum value defined as a multiple of the observation error for each observation. i.e., $inv > (obs_error * factor)$ --> fails_error_max; the default maximum value is 5 times the observation error ; the factor of 5 can be changed through max_error_* settings.
max_error_t	5.0	maximum check_max_iv error check factor for t

max_error_uv	5.0	maximum check_max_iv error check factor for u and v
max_error_pw	5.0	maximum check_max_iv error check factor for precipitable water
max_error_ref	5.0	maximum check_max_iv error check factor for gps refractivity
max_error_q	5.0	maximum check_max_iv error check factor for specific humidity
max_error_p	5.0	maximum check_max_iv error check factor for pressure
max_error_thickness	5.0	maximum check_max_iv error check factor for thickness
max_error_rv	5.0	maximum check_max_iv error check factor for radar radial velocity
max_error_rf	5.0	maximum check_max_iv error check factor for radar reflectivity
max_error_rain	5.0	maximum check_max_iv error check factor for precipitation
max_error_spd	5.0	maximum check_max_iv error check factor for wind speed (wind_sd=.true. only)
max_error_dir	5.0	maximum check_max_iv error check factor for wind direction (wind_sd=.true. only)
put_rand_seed	false	For RANDOMCV: setting to “true” allows you to enter your own seed numbers (see &wrfvar11) to generate random background perturbations.

&wrfvar6 (for minimization options)

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
max_ext_its	1	number of outer loops
ntmax	75	maximum number of iterations in an inner loop criterion (uses dimension: max_ext_its)
eps	0.01	minimization convergence criterion (uses dimension: max_ext_its); minimization stops when the norm of the gradient of the cost function gradient is reduced by a factor of eps. inner minimization stops either when the criterion is met or when inner iterations reach ntmax.
orthonorm_gradient	false	.true.: the gradient vectors are stored during the Conjugate Gradient for each iteration and used to re-orthogonalize the new gradient. This requires extra storage of large vectors (each one being the size of the control variable) but results in a better convergence of the Conjugate Gradient after around 20 iterations.

&wrfvar7

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
cv_options	5	3: NCEP Background Error model

		5: NCAR Background Error model (default)
		6: Use of moisture-multivariate background error statistics (CV6)
		7: New NCAR Background Error model (CV7)
cloud_cv_options	0	0: no hydrometeor/cloud control variables
		1: Q_total control variable
		3: Use individual cloud/hydrometeor control variables: Q_cloud, Q_rain, Q_ice, Q_snow, Q_graupel
use_cv_w	false	true: turns on W (vertical velocity) as a control variable. Works for cloud_cv_options=3 only
as1 (3)	0.25, 1.0, 1.5	tuning factors for variance, horizontal and vertical scales for control variable 1 = stream function. For cv_options=3 only.
as2 (3)	0.25, 1.0, 1.5	tuning factors for variance, horizontal and vertical scales for control variable 2 - unbalanced potential velocity. For cv_options=3 only.
as3 (3)	0.25, 1.0, 1.5	tuning factors for variance, horizontal and vertical scales for control variable 3 - unbalanced temperature. For cv_options=3 only.
as4 (3)	0.25, 1.0, 1.5	tuning factors for variance, horizontal and vertical scales for control variable 4 - pseudo relative humidity. For cv_options=3 only.
as5 (3)	0.25, 1.0, 1.5	tuning factors for variance, horizontal and vertical scales for control variable 5 - unbalanced surface pressure. For cv_options=3 only.
rf_passes	6	number of passes of recursive filter.
var_scaling1	1.0	tuning factor of background error covariance for control variable 1 - stream function. For cv_options=5, 6, and 7 only.
var_scaling2	1.0	tuning factor of background error covariance for control variable 2 - unbalanced velocity potential. For cv_options=5, 6, and 7 only.
var_scaling3	1.0	tuning factor of background error covariance for control variable 3 - unbalanced temperature. For cv_options=5, 6, and 7 only.
var_scaling4	1.0	tuning factor of background error covariance for control variable 4 - pseudo relative humidity. For cv_options=5, 6, and 7 only.
var_scaling5	1.0	tuning factor of background error covariance for control variable 5 - unbalanced surface pressure. For cv_options=5, 6, and 7 only.
len_scaling1	1.0	tuning factor of scale-length for stream function. For cv_options=5, 6, and 7 only.
len_scaling2	1.0	tuning factor of scale-length for unbalanced velocity potential. For cv_options=5, 6, and 7 only.
len_scaling3	1.0	tuning factor of scale-length for unbalanced temper-

len_scaling4	1.0	ature. For cv_options=5, 6, and 7 only.
len_scaling5	1.0	tuning factor of scale-length for pseudo relative humidity. For cv_options=5, 6, and 7 only.
je_factor	1.0	tuning factor of scale-length for unbalanced surface pressure. For cv_options=5, 6, and 7 only.
		ensemble covariance weighting factor

&wrfvar8 not used

&wrfvar9 (for program tracing)

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
stdout	6	unit number for standard output
stderr	0	unit number for error output
trace_unit	7	Unit number for tracing output. Note that units 10 and 9 are reserved for reading namelist.input and writing namelist.output respectively.
trace_pe	0	Currently, statistics are always calculated for all processors, and output by processor 0.
trace_repeat_head	10	the number of times any trace statement will produce output for any particular routine. This stops overwhelming trace output when a routine is called multiple times. Once this limit is reached a 'going quiet' message is written to the trace file, and no more output is produced from the routine, though statistics are still gathered.
trace_repeat_body	10	see trace_repeat_head description
trace_max_depth	30	define the deepest level to which tracing writes output
trace_use	false	.true.: activate tracing. Tracing gives additional performance diagnostics (calling tree, local routine timings, overall routine timings, & memory usage). It does not change results, but does add runtime overhead.
trace_use_frequent	false	.true.: activate tracing for all subroutines, even frequently called ones. Adds significant runtime overhead
trace_use_dull	false	
trace_memory	true	.true.: calculate allocated memory using a mallinfo call. On some platforms (Cray and Mac), mallinfo is not available and no memory monitoring can be done.
trace_all_pes	false	.true.: tracing is output for all pes. As stated in trace_pe, this does not change processor statistics.
trace_csv	true	.true.: tracing statistics are written to a xxxx.csv file in CSV format
use_html	true	.true.: tracing and error reporting routines will include HTML tags.

warnings_are_fatal	false	.true.: warning messages that would normally allow the program to continue are treated as fatal errors.
--------------------	-------	---

&wrfvar10 (for code developers)

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
test_transforms	false	.true.: perform adjoint tests
test_gradient	false	.true.: perform gradient test

&wrfvar11

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
check_rh	0	0 --> No supersaturation check after minimization. 1 --> supersaturation (rh> 100%) and minimum rh (rh<10%) check, and make the local adjustment of q. 2 --> supersaturation (rh> 95%) and minimum rh (rh<11%) check and make the multi-level q adjustment under the constraint of conserved column integrated water vapor
sfc_assi_options	1	1 --> surface observations will be assimilated based on the lowest model level first guess. Observations are not used when the elevation difference between the observing site and the lowest model level is larger than max_sheight_diff. 2 --> surface observations will be assimilated based on surface similarity theory in PBL. Innovations are computed based on 10-m wind, 2-m temperature and 2-m moisture.
max_sheight_diff	100.0	Height difference in meters. Stations whose model-interpolated height is different from the actual observation station height by more than this value will be rejected.
sfc_hori_intp_options	1	(SYNOP only) Specifies the method of interpolating the background to observation space 1: 4-point interpolation 2: Chooses values from neighboring model gridpoint with smallest height difference (among land points: grid points over water will not be used).
q_error_options	1	(SYNOP only) Method for calculating Q error values from RH error values 1 (default): original method 2: new method
obs_err_inflate	false	(SYNOP only) Observation error will be used as specified from observation files true: Inflate observation error values by a factor of $e^{(Zdiff /stn_ht_diff_scale)}$
stn_ht_diff_scale	200.0	(SYNOP only)

		If <code>obs_err_inflate=true</code> , observation error will be inflated by a factor of $e^{(Z_{diff} /stn_ht_diff_scale)}$. Default is 200.0
<code>psfc_from_slp</code>	false	.true.: when <code>sfc_assi_options=1</code> , re-calculates Psfc from SLP when the observation elevation is below the lowest model level height. This is not recommended.
<code>calculate_cg_cost_fn</code>	false	conjugate gradient algorithm does not require the computation of cost function at every iteration during minimization. .true.: Compute and write out cost function for each iteration into file <code>cost_fn</code> for diagnostic purposes false.: Only the initial and final cost functions are computed and output.
<code>write_detail_grad_fn</code>	false	.true.: Write out gradient for each iteration into file <code>grad_fn</code> for diagnostic purposes
<code>seed_array1</code>	1	For RANDOMCV when <code>put_rand_seed=true</code> , first integer for seeding the random function
<code>seed_array2</code>	1	For RANDOMCV when <code>put_rand_seed=true</code> , second integer for seeding the random function

&wrfvar12

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
<code>use_wpec</code>	false	true: enables the WPEC dynamic constraint term
<code>wpec_factor</code>	0.001	WPEC dynamic constraint weighting factor
<code>balance_type</code>	3	1 = geostrophic term only 2 = cyclostrophic term only 3 = geostrophic + cyclostrophic terms

&wrfvar13

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
<code>max_vert_var1</code>	99.0	specify the maximum truncation value (percentage) to explain the variance of stream function in eigenvector decomposition
<code>max_vert_var2</code>	99.0	specify the maximum truncation value (percentage) to explain the variance of unbalanced potential velocity in eigenvector decomposition
<code>max_vert_var3</code>	99.0	specify the maximum truncation value (percentage) to explain the variance of the unbalanced temperature in eigenvector decomposition
<code>max_vert_var4</code>	99.0	specify the maximum truncation value (percentage) to explain the variance of pseudo relative humidity in eigenvector decomposition
<code>max_vert_var5</code>	99.0	for unbalanced surface pressure, it should be a non-zero positive number. set <code>max_vert_var5=0.0</code> only for offline VarBC applications.

psi_chi_factor	1.0	Contribution of stream function in defining balanced part of velocity potential. For cv_options=6 only.
psi_t_factor	1.0	Contribution of stream function in defining balanced part of temperature. For cv_options=6 only.
psi_ps_factor	1.0	Contribution of stream function in defining balanced part of surface pressure. For cv_options=6 only.
psi_rh_factor	1.0	Contribution of stream function in defining balanced part of moisture. For cv_options=6 only.
chi_u_t_factor	1.0	Contribution of the unbalanced part of velocity potential in defining balanced part of temperature. For cv_options=6 only.
chi_u_ps_factor	1.0	Contribution of the unbalanced part of velocity potential in defining balanced part of surface pressure. For cv_options=6 only.
chi_u_rh_factor	1.0	Contribution of the unbalanced part of velocity potential in defining balanced part of moisture. For cv_options=6 only.
t_u_rh_factor	1.0	Contribution of the unbalanced part of temperature in defining balanced part of moisture. For cv_options=6 only.
ps_u_rh_factor	1.0	Contribution of the unbalanced part of surface pressure in defining balanced part of moisture. For cv_options=6 only.

&wrfvar14 (radiance options)

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
rtinit_nsensord	1	total number of sensors to be assimilated
rtinit_platform	-1 (max_instruments)	platforms IDs array (used dimension: rtinit_nsensord); e.g., 1 for NOAA, 9 for EOS, 10 for METOP and 2 for DMSP
rtinit_satid	-1.0 (max_instruments)	satellite IDs array (used dimension: rtinit_nsensord)
rtinit_sensor	-1.0 (max_instruments)	sensor IDs array (used dimension: rtinit_nsensord); e.g., 0 for HIRS, 3 for AMSU-A, 4 for AMSU-B, 15 for MHS, 10 for SSMIS, 11 for AIRS
rad_monitoring	0 (max_instruments)	integer array (used dimension: rtinit_nsensord); 0: assimilating mode; 1: monitoring mode (only calculate innovations)
thinning_mesh	60.0 (max_instruments)	real array (used dimension: rtinit_nsensord); specify thinning mesh size (in km) for different sensors.
thinning	false	.true.: perform thinning on radiance data
qc_rad	true	.true.: perform quality control. Do not change.
write_iv_rad_ascii	false	.true.: output radiance Observation minus Background files, which are in ASCII format and separated by sensor and processor.

<code>write_oa_rad_ascii</code>	<code>true</code>	.true.: output radiance Observation minus Analysis files (Observation minus Background information is also included), which are in ASCII format and separated by sensor and processor.
<code>use_error_factor_rad</code>	<code>false</code>	.true.: use a radiance error tuning factor file <code>radiance_error.factor</code> , which can be created with empirical values or generated using variational tuning method (Desroziers and Ivanov, 2001)
<code>use_antcorr</code>	<code>false</code>	.true.: perform Antenna Correction in CRTM
<code>rtm_option</code>	(<code>max_instruments</code>) <code>1</code>	which RTM (Radiative Transfer Model) to use (To use RTTOV, WRFDA must be compiled to include RTTOV libraries; see first section for details) 1: RTTOV 2: CRTM
<code>only_sea_rad</code>	<code>false</code>	.true.: assimilate radiance over water only
<code>use_varbc</code>	<code>true</code>	.true.: perform Variational Bias Correction. A parameter file in ASCII format called <code>VARBC.in</code> (a template is provided with the source code tar ball) is required.
<code>freeze_varbc</code>	<code>false</code>	.true: together with <code>use_varbc=.false.</code> , keep the VarBC bias parameters constant in time. In this case, the bias correction is read and applied to the innovations, but it is not updated during the minimization.
<code>varbc_factor</code>	<code>1.0</code>	for scaling the VarBC preconditioning
<code>varbc_nobsmin</code>	<code>10</code>	defines the minimum number of observations required for the computation of the predictor statistics during the first assimilation cycle. If there are not enough data (according to "VARBC_NOBSMIN") on the first cycle, the next cycle will perform a coldstart again.
<code>use_clddet_mmr</code>	<code>false</code>	.true. :use the MMR scheme to conduct cloud detection for infrared radiance
<code>use_clddet_ecmwf</code>	<code>false</code>	.true. :use the ECMWF operational scheme to conduct cloud detection for infrared radiance.
<code>airs_warmest_fov</code>	<code>false</code>	.true.: uses the observation brightness temperature for AIRS Window channel #914 as criterion for GSI thinning (with a higher amplitude than the distance from the observation location to the nearest grid point).
<code>use_crtm_kmatrix</code>	<code>true</code>	.true. use CRTM K matrix rather than calling CRTM TL and AD routines for gradient calculation, which reduces runtime noticeably.

crtm_cloud	false	.true. include cloud effects in CRTM calculations (AMSR2 instrument only) .false. do not include cloud effects (non-clear-sky pixels will be rejected)
use_rtov_kmatrix	false	.true. use RTTOV K matrix rather than calling RTTOV TL and AD routines for gradient calculation, which reduces runtime noticeably.
rtov_emis_atlas_ir	0	0: do not use IR emissivity atlas 1: use IR emissivity atlas (recommended)
rtov_emis_atlas_mw	0	0: do not use MW emissivity atlas 1: use TELSEM MW emissivity atlas (recommended) 2: use CNRM MW emissivity atlas
use_blacklist_rad	true	true.: switch off the assimilation of known problematic channels (up to year 2012) that are hard-coded in var/da/da_radiance/da_blacklist_rad.inc. false.: users need to specify proper channel selections in the radiance_info files.

&wrfvar15 (needs to be set together with &wrfvar19)

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
num_pseudo	0	Set the number of pseudo observations, either 0 or 1 (single ob)
pseudo_x	1.0	Set the x-position (I) of the OBS in unit of grid-point.
pseudo_y	1.0	Set the y-position (J) of the OBS in unit of grid-point.
pseudo_z	1.0	Set the z-position (K) of OBS with the vertical level index, in bottom-up order.
pseudo_val	1.0	Set the innovation of the ob; wind in m/s, pressure in Pa, temperature in K, specific humidity in kg/kg
pseudo_err	1.0	set the error of the pseudo ob. Unit the same as pseudo_val.; if pseudo_var="q", pseudo_err=0.001 is more reasonable.

&wrfvar16 (hybrid DA options)

<u>Variable Name</u>	<u>Default Value</u>	<u>Description</u>
use_4denvar	.false.	.true.: activate 4DEnVar capability .false.: No 4DEnVar
hybrid_dual_res	.false.	.true.: activate dual-resolution hybrid capability .false.: No dual-resolution hybrid
ensdim_alpha	0	ensemble size
alphacv_method	2	1: ensemble perturbations in control variable space 2: ensemble perturbations in model variable space
alpha_corr_type	3	1: alpha_corr_type_exp 2: alpha_corr_type_soar

alpha_corr_scale	200.0	3: alpha_corr_type_gaussian (default)
alpha_std_dev	1.0	Hybrid covariance localization (km)
alpha_vertloc_opt	2	Alpha standard deviation let WRFDA internally calculate logP-based vertical localization. be.vertloc.dat will be written out.

&wrfvar17**Variable Name**

analysis_type

Default Value

"3D-VAR"

Description

"3D-VAR": 3D-VAR mode (default);
 "QC-OBS": 3D-VAR mode plus extra filtered_obs
 output;
 "VERIFY": verification mode. WRFDA resets
 check_max_iv=.false. and ntmx=0;
 "RANDOMCV": for creating ensemble perturba-
 tions
 .true.: write out gradient of Jo for adjoint sensitivity

adj_sens

false

&wrfvar18**Variable Name**

analysis_date

Default Value

"2002-08-
 03_00:00:00.00
 00"

Description

specify the analysis time. It should be consistent
 with the first guess time; if time difference between
 analysis_date and date info read in from first guess
 is larger than the &wrfvar2 setting "analysis_accu",
 WRFDA will abort.

&wrfvar19 (needs to be set together with &wrfvar15)**Variable Name**

pseudo_var

Default Value

"t"

Description

Set the name of the OBS variable:
 'u' = X-direction component of wind,
 'v' = Y-direction component of wind,
 't' = Temperature,
 'p' = Pressure,
 'q' = Specific humidity
 "tpw": total precipitable water
 "ref": refractivity
 "ztd": zenith total delay

&wrfvar20

documentation_url

"http://www.m
 mm.ucar.edu/pe
 ople/wrfhelp/wr
 fvar/code/trunk"

&wrfvar21

time_window_min

"2002-08-
 02_21:00:00.00
 00"

start time of assimilation time window used for
 ob_format=1 and radiances to select observations
 inside the defined time_window. Note: this variable

is also used for ob_format=2 to double-check if the obs are within the specified time window.

&wrfvar22

time_window_max	"2002-08-03_03:00:00.0000"	end time of assimilation time window used for ob_format=1 and radiances to select observations inside the defined time_window. Note: this variable is also used for ob_format=2 to double-check if the obs are within the specified time window.
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&perturbation (settings related to 4DVAR options)

jcdfi_use	false	.true.: Include JcDF term in cost function. .false.: Ignore JcDF term in cost function.
jcdfi_diag	1	0: Doesn't print out the value of Jc. 1: Print out the value of Jc.
jcdfi_penalty	10	The weight to Jc term.
enable_identity	.false.	.true.: use identity adjoint and tangent linear model in 4D-Var. .false.: use full adjoint and tangent linear model in 4D-Var.
trajectory_io	.true.	.true.: use memory I/O in 4D-Var for data exchange NOTE: trajectory_io=false is deprecated and should not be used. .false.: use disk I/O in 4D-Var for data exchange
var4d_detail_out	false	.true.: output extra diagnostics for debugging 4D-Var

&radar_da (settings related to radar options)

radar_non_precip_opt	0	0 (default): no null-echo assimilation 1: KNU null-echo scheme
radar_non_precip_rf	-999.99	Reflectivity flag value (dBz) in observation file indicating non-precipitation echoes
radar_non_precip_rh_w	95	RH (%) with respect to water for non-precip retrieved Q_vapor (rqv)
radar_non_precip_rh_i	85	RH (%) with respect to water for non-precip rqv
cloudbase_calc_opt	1	Option for calculating cloud-base height: below this height retrieved humidity will not be assimilated for the use_radar_rqv option 0 (not recommended): fixed value of 1500 meters 1 (default): KNU scheme 2: NCAR scheme
radar_saturated_rf	25.0	rf value (dBz) used to indicate precipitation for rqv
radar_rqv_thresh1	40.0	rf value (dBz) used to scale down retrieved rqv
radar_rqv_thresh2	50.0	rf value (dBz) used to scale down retrieved rqv
radar_rqv_rh1	85	RH (%) for radar_saturated_rf < rf < radar_rqv_thresh1
radar_rqv_rh2	95	RH (%) for radar_rqv_thresh1 < rf < radar_rqv_thresh2

radar_rqv_h_lbound	-999.0	height (meters) lower bound for assimilating rqv
radar_rqv_h_ubound	-999.0	height (meters) upper bound for assimilating rqv

b. OBSPROC namelist variables

Variable Names	Description
&record1	
obs_gts_filename	name and path of decoded observation file
fg_format	'MM5' for MM5 application, 'WRF' for WRF application
obserr.txt	name and path of observational error file
first_guess_file	name and path of the first guess file
&record2	
time_window_min	The earliest time edge as ccyy-mm-dd_hh:mn:ss
time_analysis	The analysis time as ccyy-mm-dd_hh:mn:ss
time_window_max	The latest time edge as ccyy-mm-dd_hh:mn:ss
	** Note : Only observations between [time_window_min, time_window_max] will kept.
&record3	
max_number_of_obs	Maximum number of observations to be loaded, i.e. in domain and time window, this is independent of the number of obs actually read.
fatal_if_exceed_max_obs	.TRUE.: will stop when more than max_number_of_obs are loaded .FALSE.: will process the first max_number_of_obs loaded observations.
&record4	
qc_test_vert_consistency	.TRUE. will perform a vertical consistency quality control check on sounding
qc_test_convective_adjust	.TRUE. will perform a convective adjustment quality control check on sounding
qc_test_above_lid	.TRUE. will flag the observation above model lid
remove_above_lid	.TRUE. will remove the observation above model lid
domain_check_h	.TRUE. will discard the observations outside the domain
Thining_SATOB	.FALSE.: no thinning for SATOB data. .TRUE.: thinning procedure applied to SATOB data.
Thining_SSMI	.FALSE.: no thinning for SSMI data. .TRUE.: thinning procedure applied to SSMI data.
Thining_QSCAT	.FALSE.: no thinning for SATOB data. .TRUE.: thinning procedure applied to SSMI data.
&record5	
print_gts_read	TRUE. will write diagnostic on the decoded obs reading in file obs_gts_read.diag
print_gpspw_read	.TRUE. will write diagnostic on the gpsppw obs reading in file obs_gpspw_read.diag

print_recoverp	.TRUE. will write diagnostic on the obs pressure recovery in file obs_recover_pressure.diag
print_duplicate_loc	.TRUE. will write diagnostic on space duplicate removal in file obs_duplicate_loc.diag
print_duplicate_time	.TRUE. will write diagnostic on time duplicate removal in file obs_duplicate_time.diag
print_recoverh	.TRUE. will write diagnostic on the obs height recovery in file obs_recover_height.diag
print_qc_vert	.TRUE. will write diagnostic on the vertical consistency check in file obs_qc1.diag
print_qc_conv	.TRUE. will write diagnostic on the convective adjustment check in file obs_qc1.diag
print_qc_lid	.TRUE. will write diagnostic on the above model lid height check in file obs_qc2.diag
print_uncomplete	.TRUE. will write diagnostic on the uncompleted obs removal in file obs_uncomplete.diag
user_defined_area	.TRUE.: read in the record6: x_left, x_right, y_top, y_bottom, .FALSE.: not read in the record6.

&record6

x_left	West border of sub-domain, not used
x_right	East border of sub-domain, not used
y_bottom	South border of sub-domain, not used
y_top	North border of sub-domain, not used
ptop	Reference pressure at model top
ps0	Reference sea level pressure
base_pres	Same as ps0. User must set either ps0 or base_pres.
ts0	Mean sea level temperature
base_temp	Same as ts0. User must set either ts0 or base_temp.
tlp	Temperature lapse rate
base_lapse	Same as tlp. User must set either tlp or base_lapse.
pis0	Tropopause pressure, the default = 20000.0 Pa
base_tropo_pres	Same as pis0. User must set either pis0 or base_tropo_pres
tis0	Isothermal temperature above tropopause (K), the default = 215 K.
base_start_temp	Same as tis0. User must set either tis0 or base_start_temp.

&record7

IPROJ	Map projection (0 = Cylindrical Equidistance, 1 = Lambert Conformal, 2 = Polar stereographic, 3 = Mercator)
PHIC	Central latitude of the domain
XLONC	Central longitude of the domain
TRUELAT1	True latitude 1
TRUELAT2	True latitude 2
MOAD_CEN_LAT	The central latitude for the Mother Of All Domains
STANDARD_LON	The standard longitude (Y-direction) of the working domain.

&record8

IDD	Domain ID (1=< ID =< MAXNES), Only the observations geo-
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	graphically located on that domain will be processed. For WRF application with XLONC /= STANDARD_LON, set IDD=2, otherwise set 1.
MAXNES	Maximum number of domains as needed.
NESTIX	The I(y)-direction dimension for each of the domains
NESTJX	The J(x)-direction dimension for each of the domains
DIS	The resolution (in kilometers) for each of the domains. For WRF application, always set NESTIX(1),NESTJX(1), and DIS(1) based on the information in wrfinput.
NUMC	The mother domain ID number for each of the domains
NESTI	The I location in its mother domain of the nest domain's low left corner -- point (1,1)
NESTJ	The J location in its mother domain of the nest domain's low left corner -- point (1,1). For WRF application, NUMC(1), NESTI(1), and NESTJ(1) are always set to be 1.
&record9	
prep- bufr_output_filename	Name of the PREPBUFR OBS file.
prep- bufr_table_filename	'prepbufr_table_filename' ; do not change
output_ob_format	output 1, PREPBUFR OBS file only; 2, ASCII OBS file only; 3, Both PREPBUFR and ASCII OBS files.
use_for	'3DVAR' obs file, same as before, default 'FGAT ' obs files for FGAT '4DVAR' obs files for 4DVAR
num_slots_past	the number of time slots before time_analysis
num_slots_ahead	the number of time slots after time_analysis
write_synop	If keep synop obs in obs_gts (ASCII) files.
write_ship	If keep ship obs in obs_gts (ASCII) files.
write_metar	If keep metar obs in obs_gts (ASCII) files.
write_buoy	If keep buoy obs in obs_gts (ASCII) files.
write_pilot	If keep pilot obs in obs_gts (ASCII) files.
write_sound	If keep sound obs in obs_gts (ASCII) files.
write_amdar	If keep amdar obs in obs_gts (ASCII) files.
write_satem	If keep satem obs in obs_gts (ASCII) files.
write_satob	If keep satob obs in obs_gts (ASCII) files.
write_airep	If keep airep obs in obs_gts (ASCII) files.
write_gpspw	If keep gpspw obs in obs_gts (ASCII) files.
write_gpsztd	If keep gpsztd obs in obs_gts (ASCII) files.
write_gpsref	If keep gpsref obs in obs_gts (ASCII) files.
write_gpseph	If keep gpseph obs in obs_gts (ASCII) files.
write_ssmt1	If keep ssmt1 obs in obs_gts (ASCII) files.
write_ssmt2	If keep ssmt2 obs in obs_gts (ASCII) files.
write_ssmi	If keep ssmi obs in obs_gts (ASCII) files.
write_tovs	If keep tovs obs in obs_gts (ASCII) files.
write_qscat	If keep qscat obs in obs_gts (ASCII) files.

<code>write_profl</code>	If keep profile obs in obs_gts (ASCII) files.
<code>write_bogus</code>	If keep bogus obs in obs_gts (ASCII) files.
<code>write_airs</code>	If keep airs obs in obs_gts (ASCII) files.