



User's Guide Version 3.7

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Foreword

This document is the 2018 Gridpoint Statistical Interpolation (GSI) User's Guide, geared particularly for beginners. It describes the fundamentals of using GSI version (v) 3.7 released in November 2018. More advanced applications of GSI as well as details of assimilating specific data types can be found in the Advanced GSI User's Guide.

This User's Guide includes six chapters and three appendices:

Chapter 1 provides a background introduction of GSI.

Chapter 2 contains basic information about how to install and compile GSI - including system requirements, required software, GSI system download, and information about compilers, libraries, and building the code.

Chapter 3 focuses on the input files needed to run GSI and how to configure and run GSI through a sample run script. It also provides an example of a successful GSI run and explanations of often-used namelist variables.

Chapter 4 includes information about diagnostics and tuning of the GSI system through GSI standard output, statistic fit files, and some diagnostic tools.

Chapter 5 illustrates the GSI applications for regional WRF-ARW cases, including the setup of different data types such as conventional, radiance, and GPSRO data, and different analysis functions available in the GSI, such as running a hybrid analysis. GSI applications for global and chemical cases are also illustrated.

Appendix A introduces community tools available for GSI users.

Appendix B describes the content of the GSI namelist section OBS_INPUT.

Appendix C contains a complete list of the GSI namelist with explanations and default values.

For the latest version of the GSI documentation and code, please visit the GSI User's Website:

<http://www.dtcenter.org/com-GSI/users/index.php>

Please send questions and comments to the GSI help desk:

gsi-help@ucar.edu

This document and the annual GSI releases are made available through a community GSI effort jointly led by the Developmental Testbed Center (DTC) and the National Centers for Environmental Prediction (NCEP) Environmental Modeling Center (EMC), in collaboration with other GSI developers. To help sustain this effort, we request that those who use the

community-released GSI, the GSI helpdesk, the GSI User's Guide, or other DTC GSI services, please refer to this community GSI effort in their work and publications.

To reference this user's guide, please use:

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For referencing the general aspect of the GSI community effort, please use:

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1

Overview

1.1 GSI History and Background

The Gridpoint Statistical Interpolation (GSI) system is a unified data assimilation (DA) system for both global and regional applications. It was initially developed by the National Centers for Environmental Prediction (NCEP) Environmental Modeling Center (EMC) as a next generation analysis system based on the then operational Spectral Statistical Interpolation (SSI) analysis system ([4]; [1]; [2]). Instead of being constructed in spectral space like the SSI, the GSI is constructed in physical space and is designed to be a flexible, state-of-art system that is efficient on available parallel computing platforms. Starting with a three-dimensional variational (3DVar) data assimilation technique, the current GSI can be run as a data assimilation system of 2DVar (for surface data analysis), 3DVar, 3D ensemble-variational (3D EnVar), 4D EnVar, 3D/4D hybrid EnVar, or 4DVar (if coupled with an adjoint model from a GSI supported forecast system).

After initial development, the GSI analysis system became operational as the core of the North American Data Assimilation System (NDAS) for the North American Mesoscale (NAM) system in June 2006 and the Global Data Assimilation System (GDAS) for the Global Forecast System (GFS) in May 2007 at National Oceanic and Atmospheric Administration (NOAA). Since then, the GSI system has been adopted in various operational systems, including the National Aeronautics and Space Administration (NASA) Goddard Earth Observing System Model (GEOS), the United States Air Force (USAF) mesoscale data assimilation system, the NOAA Real-Time Mesoscale Analysis (RTMA) system, the Hurricane Weather Research and Forecasting (WRF) model (HWRF), and the Rapid Refresh (RAP) and High Resolution Rapid Refresh (HRRR) systems. The current version of GSI is also used as the data assimilation system in coming Finite-Volume Cubed-Sphere Dynamical Core (FV3) GFS system (Fv3GFS).

1.2 GSI Becomes Community Code

In 2007, the Developmental Testbed Center (DTC) began collaborating with major GSI development groups to transform the operational GSI system into a community system and support distributed development ([3]). The DTC complements the development groups in providing GSI documentation, porting GSI to multiple platforms, and testing GSI in an independent and objective environment, while maintaining equivalent functionality to what used in the operational centers. Since 2009, due to the NOAA security constraints, the DTC has been maintaining a community GSI code repository, which mirrors the EMC operational GSI code repository and facilitates community users to develop GSI. In 2017, the DTC and EMC worked together to build a unified GSI code repository for both operational and community developers and users. The unified repository facilitates direct communication among developers and helps accelerate transitions between research and operations. Based on this unified repository, the DTC releases this version of the GSI code with updated documentation.

The first community version of the GSI system was released in 2009. This user's guide describes the release of GSI (v3.7) in November 2018. The DTC provides user support through the GSI Helpdesk (gsi-help@ucar.edu), tutorials, and workshops. More information about the GSI community services can be found at the DTC GSI webpage (<https://dtcenter.org/com-GSI/users>).

1.2.1 GSI Code Management and Review Committee

The GSI code development and maintenance are managed by the Data Assimilation Review Committee (DARC). It was originally formed as the GSI Review Committee in 2010, with the goal of incorporating all major GSI development teams in the United States within a unified community framework. In 2014, EMC and DTC decided to merge their GSI code repository with the code repository of the NOAA ensemble Kalman filter (EnKF) data assimilation system. This merge enabled coordinated development of both systems and joint community support. Following the repository merging, the GSI Review Committee was transitioned to DARC, incorporating new members representing EnKF development and applications. Currently, DARC contains members from NCEP/EMC, NASA's Goddard Global Modeling and Assimilation Office (GMAO), NOAA's Earth System Research Laboratory (ESRL), the Joint Center for Satellite Data Assimilation (JCSDA), the National Center for Atmospheric Research (NCAR) Mesoscale & Microscale Meteorology Laboratory (MMM), the National Environmental Satellite, Data, and Information Service (NESDIS), USAF, the University of Maryland, and the DTC. The DTC also releases the EnKF system annually (along with GSI). Please refer to the community EnKF user's webpage (<https://dtcenter.org/EnKF/users/>) for more information.

DARC primarily steers distributed GSI/EnKF development, community code management, and support. The responsibilities of the committee are divided into two major aspects: coordination and code review. The purpose and guiding principles of the review committee are as follows:

1. Overview

- Coordination and advisory
 - Propose and shepherd new development
 - Coordinate on-going and new development
 - Establish and manage a code review and transition process
 - Community support recommendation
- Code review
 - Establish and manage a unified coding standard followed by all GSI/EnKF developers
 - Review proposed modifications to the unified code repository
 - Make decisions on whether code change proposals are accepted or denied for inclusion in the repository
 - Manage the repository
 - Oversee the timely testing and inclusion of code into the repository

1.2.2 Community Code Contributions

GSI is a community data assimilation system, open to contributions from scientists and software engineers from both the operational and research communities. DARC oversees the code transition from prospective contributors. This committee reviews proposals for code commits to the GSI repository and ensures that coding standards and tests are being fulfilled. Once the committee approves, the contributed code will be committed to the GSI code repository and available for operational implementation and public release.

To facilitate this process, the DTC is providing code transition assistance to the general research community. Prospective code contributors should contact the DTC GSI helpdesk (gsi-help@ucar.edu) as early as possible. It is contributor's responsibility to ensure the proposed code change is correct, meeting GSI coding standards, and well documented. The DTC will help the contributor run regression tests and merge the code with the top of the repository. Prospective contributors can also apply to the DTC visitor program for their GSI research and code transition. The visitor program is open to applications year-round. Please check the visitor program webpage (<https://dtcenter.org/visitor-program>) for the latest announcement of opportunity and application procedures.

1.3 About This GSI Release

As a critical part of the GSI user support, this document is provided to assist users in applying GSI to data assimilation and analysis studies. It was composed by the DTC and reviewed by the DARC members. Please note that the major focuses of the DTC are currently on testing and evaluation of GSI for regional numerical weather prediction (NWP). GSI for global and chemical applications are briefly discussed in the document. The document is based on GSI v3.7 release. Active users can contact the DTC (gsi-help@ucar.edu) for developmental versions of GSI and access to the GSI code repository.

1.3.1 What Is New in This Release Version

The following lists some of the new functions and changes included in the v3.7 release of the GSI versus the v3.6 release:

Observational aspects:

- Further code generalization on all-sky radiance assimilation
- Added the capability of assimilating satellite hydrometeor retrievals (integrated liquid-water content (LWC) and integrated solid-water content (SWC)) into GSI
- Added assimilation of KOMPSAT5 GPSRO data
- Modifications and improvements for CrIS and CrIS-FSR analysis
- Updated CRTM library and coefficients to version 2.3

Code optimization and refactoring:

- Added wrf interface as a library (wrflib). No need to compile WRF with GSI and EnKF
- Added code to read in multiple-time level ARW ensemble forecast to get 4D perturbations

Application specific updates:

- Application for FV3
 - Added FV3GFS interface
 - Added regional GSI interface with single FV3 tile
 - Added new scripts and fix files for using global and regional FV3 applications
- RTMA
 - Updated mesonet uselist
 - Added assimilation of mesonet visibility data

Other enhancements and updates:

- Removed GOTO statements from the code.
- Added capability to read GFS ensemble member nemsio files in parallel simultaneously
- Removed srw data type option
- Clean up the output and add a "verbose" namelist flag.
- Updated surface pressure observation error
- Added Safeguard to filter out bad data in the radiance bias correction.
- Utility updates such as radar data plotting.
- Bug fixes

Besides the above-mentioned changes, this version release code have a unified cmake-based build system. This cmake build system has been tested on multiple platform by DTC and EMC and released as the only building utility for this release version of GSI.

1.3.2 Observations Used by This Version

GSI is used by various applications on multiple scales. The types of observations GSI can assimilate vary from conventional to aerosol observations. Users should use observations with caution to fit their specific applications. The GSI v3.7 can assimilate, but is not limited to, the following types of observations:

Conventional observations (including satellite retrievals):

- Radiosondes
- Pilot balloon (PIBAL) winds
- Synthetic tropical cyclone winds
- Wind profilers: USA, Jan Meteorological Agency (JMA)
- Conventional aircraft reports
- Aircraft to Satellite Data Relay (ASDAR) aircraft reports
- Meteorological Data Collection and Reporting System (MDCRS) aircraft reports
- Drosondes
- Moderate Resolution Imaging Spectroradiometer (MODIS) IR and water vapor winds
- Geostationary Meteorological Satellite (GMS), JMA, and Meteosat cloud drift IR and visible winds
- European Organization for the Exploitation of Meteorological Satellites (EUMETSAT) and GOES water vapor cloud top winds
- GEOS hourly IR and cloud top wind
- Surface land observations
- Surface ship and buoy observations
- Special Sensor Microwave Imager (SSM/I) wind speeds
- Quick Scatterometer (QuikSCAT), the Advanced Scatterometer (ASCAT) and Oceansat-2 Scatterometer (OSCAT) wind speed and direction
- RapidScat observations
- SSM/I and Tropical Rainfall Measuring Mission (TRMM) Microwave Imager (TMI) precipitation estimates
- Velocity-Azimuth Display (VAD) Next Generation Weather Radar ((NEXRAD) winds
- Global Positioning System (GPS) precipitable water estimates
- Sea surface temperatures (SSTs)
- Doppler wind Lidar
- Aviation routine weather report (METAR) cloud coverage
- Flight level and Stepped Frequency Microwave Radiometer (SFMR) High Density Observation (HDOB) from reconnaissance aircraft
- Tall tower wind

Satellite radiance/brightness temperature observations (instrument/satellite ID):

- SBUV: *NOAA-17, NOAA-18, NOAA-19*
- High Resolution Infrared Radiation Sounder (HIRS): *Meteorological Operational-A (MetOp-A), MetOp-B, NOAA-17, NOAA-19*
- GOES imager: *GOES-11, GOES-12*
- Atmospheric IR Sounder (AIRS): *aqua*

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- AMSU-A: *MetOp-A, MetOp-B, NOAA-15, NOAA-18, NOAA-19, aqua*
- AMSU-B: *MetOp-B, NOAA-17*
- Microwave Humidity Sounder (MHS): *MetOp-A, MetOp-B, NOAA-18, NOAA-19*
- SSMI: *DMSP F14, F15, F19*
- SSMI/S: *DMSP F16*
- Advanced Microwave Scanning Radiometer for Earth Observing System (AMSR-E): *aqua*
- GOES Sounder (SNDR): *GOES-11, GOES-12, GOES-13*
- Infrared Atmospheric Sounding Interferometer (IASI): *MetOp-A, MetOp-B*
- Global Ozone Monitoring Experiment (GOME): *MetOp-A, MetOp-B*
- Ozone Monitoring Instrument (OMI): *aura*
- Spinning Enhanced Visible and Infrared Imager (SEVIRI): *Meteosat-8, Meteosat-9, Meteosat-10*
- Advanced Technology Microwave Sounder (ATMS): *Suomi NPP*
- Cross-track Infrared Sounder (CrIS): *Suomi NPP*
- GCOM-W1 AMSR2
- GPM GMI
- Megha-Tropiques SAPHIR
- Himawari AHI

Others:

- GPS Radio occultation (RO) refractivity and bending angle profiles
- Solar Backscatter Ultraviolet (SBUV) ozone profiles, Microwave Limb Sounder (MLS) (including NRT) ozone, and Ozone Monitoring Instrument (OMI) total ozone
- Doppler radar radial velocities
- Radar reflectivity Mosaic
- Tail Doppler Radar (TDR) radial velocity and super-observation
- Tropical Cyclone Vitals Database (TCVital)
- Particulate matter (PM) of 10-um diameter, 2.5-um diameter or less
- MODIS AOD (when using GSI-chem package)
- Significant wave height observations from JASON-2, JASON-3, SARAL/ALTIKA and CRYOSAT-2

Please note that some of these above mentioned data are not yet fully tested and/or implemented for operations. Therefore, the current GSI code might not have an optimal setup for these data.



2

Software Installation

2.1 Introduction

The DTC GSI is a community distribution of NOAA's operational GSI. The community GSI expands the portability of the operational code by adding a flexible build system which provides support for many common platforms and compilers. GSI v3.7 builds and runs on most standard Linux platforms using Intel, PGI, or Gnu compilers.

This chapter describes how to build and install the DTC community GSI software on your local Linux computing resources.

The GSI build process consists of three general steps:

- Obtaining the source code for GSI.
- Setting the appropriate environment variables for the GSI build.
- Configuring and compiling the GSI source code using CMake.

This chapter is organized as follows: Section 2.2 describes how to obtain the source code. Section 2.3 covers the directory structure and supplemental NCEP libraries included with the distribution. Section 2.4 starts with a brief overview of CMake and continues with an example of the build process in section 2.4.1. Section 2.4.2 describes all of the current optional settings for the build. Sections 2.5 and 2.5.1 cover the system requirements (tools, libraries, and environment variable settings) and currently supported platforms in detail. Section 2.6 discusses what to do if you have problems with the build and where to get help.

Advanced topics, such as customizing the build, porting to new platforms, and debugging

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can be found in the GSI Advanced User's Guide.

2.2 Obtaining and Setting Up the Source Code

The community GSI resources, including source code, build system, utilities, practice data, and documentation, are available from the DTC community GSI users website, located at

<http://www.dtcenter.org/com-GSI/users/index.php>

The source code is available by first selecting the Download tab on the vertical menu located on the left column of the page, and then selecting the GSI/EnKF System submenu. New users must first register before downloading the source code. Returning users only need to enter their registration email address to log in. After accessing the download page, select the link to the `comGSIv3.7_EnKFv1.3.tar` tarball to download the most recent version of the source code (November 2018). Selecting the newest release of the community GSI is critical for having the most recent capabilities, versions of supplemental libraries, and bug fixes. Full support is only offered for the latest code releases.

To analyze satellite radiance observations, GSI requires the use of CRTM coefficients. It is crucial to use **only** the version of CRTM coefficients provided on the GSI website. Due to their large size, these are available as a separate tarfile. They can be downloaded by selecting the link to the tarball for the CRTM 2.3.0 Big Endian coefficients from the web page. For all compilers, use the big endian byte order coefficients found in the first CRTM link.

The download page also contains links to the fixed files necessary for running global GSI:

- Global fix files to run Global GSI

The community GSI version 3.7 comes in a tar file named `comGSIv3.7_EnKFv1.3.tar`. The tar file may be unpacked by using the UNIX commands:

```
gunzip comGSIv3.7_EnKFv1.3.tar.gz
tar -xvf comGSIv3.7_EnKFv1.3.tar
```

This creates the top level GSI directory `comGSIv3.7_EnKFv1.3/`. After downloading the source code, and prior to building, the user should check the known issues link on the download page of the DTC website to determine if any bug fixes or platform specific customizations are needed.

2.3 Directory Structure, Source Code and Supplemental Libraries

The GSI system includes the GSI source code, the build system, supplemental libraries, fixed files, and run scripts. Again, with V3.7 of GSI the directory structure has been slightly

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changed to accommodate the new CMake build system. The following table lists the system components found inside the root GSI directory.

Table 2.1: Description of top level directories

Directory Name	Content
cmake/	CMake build rules and configure files.
fix/	Fixed input files required by a GSI analysis, such as background error covariances, observation error tables; Excluding the CRTM coefficients
libsrc/	Supplemental library source code, required for the CMake build
README.comgsi	An overview on how to build/run GSI/EnKF and get helps
README.GSI_Docker	An overview on how to run GSI/EnKF in a docker container
ush/	Sample run scripts and namelists
src/	GSI source code and makefiles
src/enkf	EnKF source code and makefiles
util/	Tools for GSI diagnostics

For convenience, supplemental NCEP libraries for building GSI are included in the `libsrc/` directory. These libraries will be built prior to compiling GSI. These supplemental libraries are listed in the table below.

Table 2.2: List of NCEP libraries used by GSI

Directory Name	Content
bacio/	NCEP BACIO library
bufr/	NCEP BUFR library
crtm/	JCSDA community radiative transfer model
GSD/	GSD Cloud analysis library
ncdiag/	Misc NetCDF libraries
nemsio/	NEMS I/O library
sfcio/	NCEP GFS surface file i/o module
sigio/	NCEP GFS atmospheric file i/o module
sp/	NCEP spectral - grid transforms
w3emc/	NCEP/EMC W3 library (date/time manipulation, GRIB)
w3nco/	NCEP/NCO W3 library (date/time manipulation, GRIB)
wrfliib	WRF IO libraries

2.4 CMake Build System

A new unified build system based on CMake has been added to the GSI code. While the new build system is a significant departure from the previous DTC build system, CMake is a very powerful cross-platform open-source build system that has many capabilities to make building GSI easier.

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The CMake build infrastructure consists of a top level directory with the name `cmake/` and configuration files in each directory named (`CMakeLists.txt`). CMake relies on a two step command line process, similar to the old UNIX "configure" and "compile." Command line arguments are used to specify paths and compilers. By default CMake is configured to build the source code "out-of-place," meaning that it does not populate the GSI directory with the build. The location for the build must be specified by the user.

Once a compiler has been chosen, CMake generates local makefiles by invoking the `cmake` command with the proper arguments. One of those arguments selects that a local build of the NCEP libraries needed by GSI will be conducted prior to the source code being built. The final step is to invoke a parallel build (`make`) of the code.

A significant advantage with using CMake to build the code, is that CMake automatically generates code dependencies each time a build is invoked, allowing the use of a parallel `make` as the final step of the build. This greatly reduces the time it take to complete the build. Because of the parallel compiling, the time to complete the CMake build is typically a quarter of the time needed for the serial DTC "configure" and "compile" to complete.

Once the build is complete, the two executables `gsi.x` and `enkf_wrf.x` are placed in the `bin` directory of the build directory. Note that the name of the executables and their location differs from the traditional DTC build.

Summary of CMake build steps:

1. Set up the build environment
 - Select a specific compiler by loading modules or setting paths as needed
 - Set the environment path for the NetCDF library (The HDF5 library is also needed if your NetCDF version is 4+).
 - Set the LaPack/MKL mathematics libraries as needed.
2. Create a directory for your build and change into it.
3. Run the `cmake` command with the appropriate arguments.
4. Run `make`
5. The completed executables `gsi.x` and `enkf_wrf.x` will be located in the build directory within the `bin` directory.

2.4.1 Example build using the Intel Fortran Compiler

This section details how to build GSI on the NCAR supercomputer Cheyenne using the Intel V18.0.1 fortran compiler, the 2018 MKL libraries, NetCDF v4.6.1, and the 2018 Intel MPI libraries.

- Set up your build environment

```
module purge
module load cmake/3.9.1
module load intel/18.0.1 ncarenv/1.2 ncarcompilers/0.4.1 mkl/2018.0.1
module load netcdf/4.6.1
module load impi/2018.1.163
```

- Next, create a build directory and move into it.

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```
mkdir ./build
cd build
```

- From within the build directory, invoke the cmake command

```
cmake path_to_the_comGSI_directory
```

The makefiles customized to your platform and the environment you have set are now created.

- Run make (parallel compile) on eight cores

```
make -j8
```

Because CMake has already figured out the dependencies, the make can run in parallel.

Lets look at relevant arguments you may use for cmake when building GSI. The first is `-DBUILD_CORELIBS=ON`. This argument directs CMake to look in the `libsrc/` directory to build the NCEP libraries needed for GSI. Default is ON, so you don't need to use this argument by default.

The argument `CMAKE_C_COMPILER` specifies the C compiler to use with the build. For the Intel compiler, use `icc`. The argument `CMAKE_CXX_COMPILER` specifies the C++ compiler to use with the build. For the Intel compiler, use `icpc`. The argument `CMAKE_Fortran_COMPILER` specifies the Fortran compiler to use with the build. Because this is an parallel MPI code, on Cheyenne use `mpif90` instead of `ifort`. Typically, the only time one would explicitly specify the compilers on the command line is when you wish to override a default choice set in your dot files or if you were to create a batch script to conduct the build.

To run the cmake command with a compiler other than Intel only requires replacing the values for `CMAKE_C_COMPILER`, `CMAKE_CXX_COMPILER`, and `CMAKE_Fortran_COMPILER` that correspond to the new compiler. Table 2.3 lists these values.

Table 2.3: The compiler specific values for the cmake compiler argument

Compiler	C	C++	FC
Intel	icc	icpc	mpif90
PGI	pgcc	pgcc++	mpif90
GNU	gcc	g++	mpif90

2.4.2 Default Options

The CMake build has a number of options to when building. These are enacted by adding the option with the prefix `-D` to the command line call to cmake. For instance, in the example above, the `BUILD_CORELIBS` option is enacted by adding `-DBUILD_CORELIBS=ON` to the cmake call. The currently available command line build options are:

- `BUILD_ENKF`: builds the enkf executable (default is ON)
- `BUILD_CORELIBS`: Attempts to build the core libraries from source
- `USE_WRF`: Builds GSI with WRF dependencies (default is ON)
- `BUILD_GFS`: Builds ENKF using GFS (default is ON)

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- BUILD_WRF: Builds ENKF using WRF (default is OFF)
- BUILD_NMMB: Builds ENKF using NMMB (default is OFF)

Note that by default GSI is built with the regional WRF IO bindings, and EnKF is built with GFS. See the readme file *README.cmake* in the top directory for a current list of options; or look in the CMakeLists.txt in the top directory and search for options.

2.5 Software Requirements and Compiler Specific Notes

The source code for GSI is written in FORTRAN, FORTRAN 90, and C. In addition, the parallel executables require some flavor of MPI and OpenMP for the distributed memory parallelism. The I/O relies on the NetCDF I/O libraries. And CMake is needed to build the code.

The basic requirements for building and running the GSI system are the following:

- CMake V2.8+
- A FORTRAN compiler that supports the 2003 or newer standard
- C compiler
- MPI v1.2+
- OpenMP
- NetCDF V4.2+ and HDF5
- LAPACK and BLAS mathematics libraries, or equivalent

In addition, GSI and EnKF require linear algebra libraries such as LAPACK and BLAS. The Intel compiler usually comes with a vendor provided mathematics library known as the *Mathematics Kernel Libraries* or MKL for short. While *most* installations of the Intel compiler come with the MKL libraries installed not all do. In addition, if it is not installed correctly, the ifort compiler does not automatically load the library. It is therefore necessary to set the LAPACK_PATH variable to the location of the MKL libraries when using the Intel compiler. You may need to ask your system administrator for the correct path to these libraries.

IBM systems typically come installed with the LAPACK equivalent ESSL library that links automatically. Likewise, the PGI compiler often comes with a vendor provided version of LAPACK that links automatically with the compiler. The Gnu compiler is the only one that does not come with any installed version of the LAPACK and BLAS libraries.

Because all but the last of these tools and libraries are typically the purview of system administrators to install and maintain, they are lumped together here as part of the basic system requirements.

2.5.1 Compilers Tested for Release

Version 3.7 of the DTC community GSI system extensively tested with a variety of compilers and versions of those compilers, on standard Linux platforms. To summarize, Intel com-

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piler versions 16-18 should build and run without issue. Early version 15 has some known issues, but 15.1, 15.3, and 15.6 should work. For the Portland Group compiler, versions 16-18 should work without issue, and Gnu version 6.3, 7.1, and 7.3 are known to work. Version 18 of Gnu has issues and is not currently supported. See release notes (https://dtcenter.org/com-GSI/users/support/release_notes/index_GSIv3.7.php) for the complete list compilers and versions tested.

Unforeseen build issues may occur when using older compiler and library versions. As always, the best results will be achieved by using the most recent compiler versions.

2.6 Getting Help and Reporting Problems

Should a user experience any difficulty building GSI on their system, please first confirm that all the required software is properly installed. Should all these check out, feel free to contact the community GSI Help Desk for assistance at

`gsi-help@ucar.edu`

At a minimum, when reporting code building problems to the helpdesk, please include a copy of the build log with your e-mail.



3

Running GSI

This chapter discusses the issues of running GSI. It starts with introductions to the input data required to run GSI, then proceeds with a detailed explanation of an example GSI run script and introductions to files produced by a successful GSI run. It concludes with some frequently used options from the GSI namelist.

3.1 Input Data Required to Run GSI

In most cases, three types of input data (background, observations, and fixed files) must be available before running GSI. In some special idealized cases, such as a pseudo single observation test, GSI can be run without any observations. If running GSI with the 3D EnVar hybrid option, global or regional ensemble forecasts are also needed.

3.1.1 Background or First Guess Field

As with other data analysis systems, the background or first guess fields may come from a model forecast conducted separately or from a previous data assimilation cycle. The following is a list of the types of background files that can be used by this release version of GSI:

- a) WRF-NMM input fields in binary format
- b) WRF-NMM input fields in NetCDF format
- c) WRF-ARW input fields in binary format
- d) WRF-ARW input fields in NetCDF format

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- e) GFS input fields in binary format or through NEMS I/O
- f) NEMS-NMMB input fields
- g) RTMA input files (2-dimensional binary format)
- h) WRF-Chem GOCART input fields with NetCDF format
- i) CMAQ binary file

The Weather Research and Forecasting (WRF) community modeling system includes two dynamical cores: the Advanced Research WRF (ARW) and the Nonhydrostatic Mesoscale Model (NMM). The GFS (Global Forecast System), NEMS (National Environmental Modeling System)-NMMB (Nonhydrostatic Mesoscale Model B-Grid), and RTMA (Real-Time Mesoscale Analysis) are operational systems at the National Center for Environmental Prediction (NCEP). The DTC mainly supports GSI for regional WRF applications. Therefore, most of the multiple platform tests were conducted using WRF netcdf background files (d). The DTC also supports the GSI in global and chemical applications with limited resources. The following backgrounds have been tested for this release:

1. ARW NetCDF (d) were tested with multiple cases
2. GFS (e) was tested with multiple NCEP cases
3. WRF-Chem NetCDF (h) was tested with a single case
4. NEMS-NMMB(f) was tested with a single case

3.1.2 Observations

GSI can analyze many types of observational data, including conventional data, satellite radiance observations, GPS Radio Occultations, and radar data, among others. The default observation file names are given in the released GSI namelist, with corresponding observations included in each file. Sample BUFR files available for download from the NCEP website listed in table 3.1.

The observations are complex and many observations need format converting and quality control before being used by GSI. GSI ingests observations saved in BUFR format (with NCEP specified features). The NCEP processed PrepBUFR and BUFR files can be used directly. If users need to introduce their own data into GSI, please check the following website for the User's Guide and examples of BUFR/PreBUFR processing:

<http://www.dtcenter.org/com-GSI/BUFR/index.php>

DTC supports BUFR/PrepBUFR data processing and quality control as part of the GSI community tasks.

GSI can analyze all of the data types in table 3.1, but each GSI run (for both operation and case study purposes) only uses a subset of the data. Some data may be outdated and not available, some are in monitoring mode, and some may have quality issues during certain periods. Users are encouraged to check data quality prior to running an analysis. The following NCEP links provide resources that include data quality history:

http://www.emc.ncep.noaa.gov/mmb/data_processing/Satellite_Historical_Documentation.htm

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http://www.emc.ncep.noaa.gov/mmb/data_processing/Non-satellite_Historical_Documentation.htm

Because the current regional models do not have ozone as a prognostic variable, ozone data are not assimilated on the regional scale.

GSI can be run without any observations to see how the moisture constraint modifies the first guess (background) field. GSI can also be run in a pseudo single observation mode, which does not require any BUFR observation files. In this mode, users should specify observation information in the namelist section SINGLEOB_TEST (see Section 4.2 for details). As more data files are used, additional information will be added through the GSI analysis.

3.1.3 Fixed Files (Statistics and Control Files)

A GSI analysis also needs to read specific information from statistic files, configuration files, bias correction files, and CRTM coefficient files. We refer to these files as fixed files and they are located in a directory called `fix/` in the release package, except for CRTM coefficients.

Table 3.2 lists fixed files required for a GSI run, the content of the files, and corresponding example files from the regional and global applications:

Because most of those fixed files have hardwired names inside the GSI, a GSI run script needs to copy or link those files (right column in table 3.2) from the `./fix` directory to the GSI run directory with the file name required in GSI (left column in table 3.2). For example, if GSI runs with an ARW background, the following line should be in the run script:

```
cp ${path of the fix directory}/anavinfo_arw_netcdf anavinfo
```

Note that in this release, there is a strict rule that the numbers of vertical levels in the file `anavinfo` must match the background file (for example, `wrfinput_d01`) for the 3-dimensional variables. Otherwise GSI will fail. To identify the correct numbers of vertical levels, users can dump out (use `ncdump -h`) the dimensions from the NetCDF background file and find the number for `bottom_top` and `bottom_top_stag`. For example, if the dimensions for the background file is:

```
bottom_top = 50 ;  
bottom_top_stag = 51 ;
```

Then the corresponding `anavinfo` file should have 51 levels for `prse` (3-dimensional pressure field) and 50 levels for other three-dimensional variables such as `u`, `v`, `tv`, `q`, `oz`, `cw`, etc. For details, users can dump out the global attributes of the background file and find the number of vertical levels for each variable. The following shows part of the `anavinfo` file for the above background:

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Table 3.1: GSI observation file names, content, and examples

GSI Name	Content	Example file names
prepbuf	Conventional observations, including ps, t, q, pw, uv, spd, dw, sst	gdasl.t12z.prepbuf.nr
satwndbuf	satellite winds observations	gdasl.t12z.satwnd.tm00.buf_d
amsuabuf	AMSU-A 1b radiance (brightness temperatures) from satellites NOAA-15, 16, 17,18, 19 and METOP-A/B	gdasl.t12z.lbamua.tm00.buf_d
amsubbuf	AMSU-B 1b radiance (brightness temperatures) from satellites NOAA-15, 16,17	gdasl.t12z.lbamub.tm00.buf_d
radarbuf	Radar radial velocity Level 2.5 data	ndas.t12z.radwnd.tm12.buf_d
gpsrobuf	GPS radio occultation and bending angle observation	gdasl.t12z.gpsro.tm00.buf_d
ssmirrbuf	Precipitation rate observations from SSM/I	gdasl.t12z.spssmi.tm00.buf_d
tmirrbuf	Precipitation rate observations from TMI	gdasl.t12z.sptrmm.tm00.buf_d
sbuvbuf	SBUV/2 ozone observations from satellite NOAA-16, 17, 18, 19	gdasl.t12z.osbuv8.tm00.buf_d
hirs2buf	HIRS2 1b radiance from satellite NOAA-14	gdasl.t12z.lbhrs2.tm00.buf_d
hirs3buf	HIRS3 1b radiance observations from satellite NOAA-16, 17	gdasl.t12z.lbhrs3.tm00.buf_d
hirs4buf	HIRS4 1b radiance observation from satellite NOAA-18, 19 and METOP-A/B	gdasl.t12z.lbhrs4.tm00.buf_d
msubuf	MSU observation from satellite NOAA 14	gdasl.t12z.lbmsu.tm00.buf_d
airsbuf	AMSU-A and AIRS radiances from satellite AQUA	gdasl.t12z.airsev.tm00.buf_d
mhsbuf	Microwave Humidity Sounder observation from NOAA-18, 19 and METOP-A/B	gdasl.t12z.lbmhs.tm00.buf_d
ssmitbuf	SSMI observation from satellite f13, f14, f15	gdasl.t12z.ssmi.tm00.buf_d
amsrebuf	AMSRE radiance from satellite AQUA	gdasl.t12z.amsre.tm00.buf_d
ssmisbuf	SSMIS radiances from satellite f16	gdasl.t12z.ssmis.tm00.buf_d
gsndlbuf	GOES sounder radiance (sndrd1, sndrd2, sndrd3, sndrd4) from GOES-11, 12, 13, 14, 15.	gdasl.t12z.goesfv.tm00.buf_d
l2rwbuf	NEXRAD Level 2 radial velocity	ndas.t12z.nexrad.tm12.buf_d
gsndrbuf	GOES sounder radiance from GOES-11, 12	gdasl.t12z.goesnd.tm00.buf_d
gimgbuf	GOES imager radiance from GOE-11, 12	
omibuf	Ozone Monitoring Instrument (OMI) observation NASA Aura	gdasl.t12z.omi.tm00.buf_d
iasibuf	Infrared Atmospheric Sounding Interferometer sounder observations from METOP-A/B	gdasl.t12z.mtiasi.tm00.buf_d
gomebuf	The Global Ozone Monitoring Experiment (GOME) ozone observation from METOP-A/B	gdasl.t12z.gome.tm00.buf_d
mlsbuf	Aura MLS stratospheric ozone data from Aura	gdasl.t12z.mlsbuf.tm00.buf_d
tcvitl	Synthetic Tropic Cyclone-MSLP observation	gdasl.t12z.syndata.tcvitals.tm00
seviribuf	SEVIRI radiance from MET-08,09,10	gdasl.t12z.sevcsr.tm00.buf_d
atmsbuf	ATMS radiance from Suomi NPP	gdasl.t12z.atms.tm00.buf_d
crisbuf	CRIS radiance from Suomi NPP	gdasl.t12z.cris.tm00.buf_d
modisbuf	MODIS aerosol total column AOD observations from AQUA and TERRA	

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```
state_derivatives::
!var level src
ps 1 met_guess
u 50 met_guess
v 50 met_guess
tv 50 met_guess
q 50 met_guess
oz 50 met_guess
cw 50 met_guess
prse 51 met_guess
::
```

Table 3.2: GSI fixed files, content, and examples

GSI Name	Content	Example file names
anavinfo	Information file to set control and analysis variables	anavinfo_arw_netcdf anavinfo_ndas_netcdf global_anavinfo.l64.txt
berror_stats	background error covariance	nam_nmmstat_na.gcv nam_glb_berror.f77.gcv global_berror.l64y386.f77
errtable	Observation error table	nam_errtable.r3dv prepobs_errtable.global
<i>Observation data control file (more detailed explanation in Section 4.3)</i>		
convinfo	Conventional observation information file	global_convinfo.txt nam_regional_convinfo.txt
satinfo	satellite channel information file	global_satinfo.txt
pcpinfo	precipitation rate observation information file	global_pcpinfo.txt
ozinfo	ozone observation information file	global_ozinfo.txt
<i>Bias correction and Rejection list</i>		
satbias_angle	satellite scan angle dependent bias correction file	global_satangbias.txt
satbias_in	satellite mass bias correction coefficient file	sample.satbias
	combined satellite angle dependent and mass bias correction coefficient file	gdasl.t00z.abias.new
t_rejectlist, w_rejectlist,..	Rejection list for T, wind, et al. in RTMA	new_rtma_t_rejectlist new_rtma_w_rejectlist

Each operational system, such as GFS, NAM, RAP, and RTMA, has their own set of fixed files. For your specific GSI runs, you need to get the correct set of fixed files. Fixed files for regional applications are included in this GSI/EnKF release and put under the *fix/* directory. Fixed files for global applications are not included in this release in order to save space. Please download `comGSIv3.7_EnKFv1.3_fix_global.tar.gz` if you need to run global cases. Note that little endian background error covariance files are no longer supported.

Each release version of the GSI calls a certain version of the CRTM library and needs corresponding CRTM coefficients to do radiance data assimilation. This version of GSI uses CRTM 2.3.0. The coefficient files are listed in table 3.3.

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Table 3.3: List of radiance coefficients used by CRTM

File name used in GSI	Content	Example files
Nalli.IRwater.EmisCoeff.bin NPOESS.IRice.EmisCoeff.bin NPOESS.IRsnow.EmisCoeff.bin NPOESS.IRland.EmisCoeff.bin NPOESS.VISice.EmisCoeff.bin NPOESS.VISland.EmisCoeff.bin NPOESS.VISsnow.EmisCoeff.bin NPOESS.VISwater.EmisCoeff.bin FASTEM6.MWwater.EmisCoeff.bin	IR surface emissivity coefficients	Nalli.IRwater.EmisCoeff.bin NPOESS.IRice.EmisCoeff.bin NPOESS.IRsnow.EmisCoeff.bin NPOESS.IRland.EmisCoeff.bin NPOESS.VISice.EmisCoeff.bin NPOESS.VISland.EmisCoeff.bin NPOESS.VISsnow.EmisCoeff.bin NPOESS.VISwater.EmisCoeff.bin FASTEM6.MWwater.EmisCoeff.bin
AerosolCoeff.bin	Aerosol coefficients	AerosolCoeff.bin
CloudCoeff.bin	Cloud scattering and emission coefficients	CloudCoeff.bin
\$(satsen).SpCcoeff.bin	Sensor spectral response characteristics	\$(satsen).SpCcoeff.bin
\$(satsen).TauCoeff.bin	Transmittance coefficients	\$(satsen).TauCoeff.bin

3.2 GSI Run Script

In this release version, three sample run scripts are available for different GSI applications:

- `ush/comgsi_run_regional.ksh` for regional GSI
- `ush/comgsi_run_global.ksh` for global GSI (GFS)
- `ush/comgsi_run_chem.ksh` for chemical analysis

These scripts will be called to generate GSI namelists:

- `ush/comgsi_namelist.sh` for regional GSI
- `ush/comgsi_namelist_gfs.sh` for global GSI (GFS)
- `ush/comgsi_namelist_chem.sh` for GSI chemical analysis

We will introduce the regional run scripts (`comgsi_run_regional.ksh`) in detail in the following sections and introduce the global run script when we discuss the GSI global application in Section 6.1.

Note there is also a run script for regional EnKF (`comenkf_run_regional.ksh`), a run script for global EnKF (`comenkf_run_global.ksh`) and the EnKF namelist script (`comenkf_namelist.sh`) in the same directory, which will be introduced in the EnKF User's Guide.

3.2.1 Steps in the GSI Run Script

The GSI run script creates a run time environment necessary to run the GSI executable. A typical GSI run script includes the following steps:

1. Request computer resources to run GSI.
2. Set environmental variables for the machine architecture.
3. Set experimental variables (such as experiment name, analysis time, background, and observation).
4. Set the script that generates the GSI namelist.
5. Check the definitions of required variables.
6. Generate a run directory for GSI (sometimes called a working or temporary directory).
7. Copy the GSI executable to the run directory.
8. Copy the background file to the run directory and create an index file listing the location and name of ensemble members if running with a hybrid set up.
9. Link observations to the run directory.
10. Link fixed files (statistic, control, and coefficient files) to the run directory.
11. Generate namelist for GSI.
12. Run the GSI executable.
13. Post-process: save analysis results, generate diagnostic files, and clean the run directory.
14. Run GSI as observation operator for EnKF, only for `if_observer=Yes`.

Typically, users only need to modify specific parts of the run script (steps 1, 2, and 3) to fit their specific computer environment and point to the correct input/output files and directories. Users may also need to modify step 4 if changes are made to the namelist and it is under a different name or at a different location. The next section (3.2.2) covers each of these modifications for steps 1 to 3. Section 3.2.3 will dissect a sample regional GSI run script and introduce each piece of this sample GSI run script. Users should start with the run script provided in the same release package with the GSI executable and modify it for their own run environment and case configuration.

3.2.2 Customization of the GSI Run Script

3.2.2.1 Setting Up the Machine Environment

This section focuses on step 1 of the run script: modifying the machine specific entries. Specifically, this consists of setting Unix/Linux environment variables and selecting the correct parallel run time environment (batch system with options).

GSI can be run with the same parallel environments as other MPI programs, for example:

- IBM supercomputer using LSF (Load Sharing Facility)
- IBM supercomputer using LoadLevel
- Linux clusters using PBS (Portable Batch System)
- Linux clusters using LSF

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- Linux workstation (no batch system)
- Intel Mac Darwin workstation with PGI compiler (no batch system)

Two queuing systems are listed below as examples:

Machine & queue system	Linux Cluster with LSF	Linux Cluster with PBS	Workstation
example	<pre>#BSUB -P ????????? #BSUB -W 00:10 #BSUB -n 4 #BSUB -R "span[ptile=16] #BSUB -J gsi #BSUB -o gsi.%J.out #BSUB -e gsi.%J.err #BSUB -q small</pre>	<pre>#PBS -l procs=4 #PBS -n #PBS -o gsi.out #PBS -e gsi.err #PBS -N GSI #PBS -l walltime=00:20 #PBS -A ???????</pre>	No batch system, skip this step

In both of the examples above, environment variables are set specifying system resource management, such as the number of processors, the name/type of queue, maximum wall clock time allocated for the job, options for standard out and standard error, etc. Some platforms need additional definitions to specify Unix environment variables that further define the run environment.

These variable settings can significantly impact the GSI run efficiency and accuracy of the GSI results. Please check with your system administrator for optimal settings for your computer system. Note that while the GSI can be run with any number of processors, it will not scale well with the increase of processor numbers after a certain threshold based on the case configuration and GSI application types.

3.2.2.2 Setting up the Running Environment

There are only two options to define in this block.

```
# GSIPROC = processor number used for GSI analysis
#-----
GSIPROC=4
ARCH='LINUX_LSF'
# Supported configurations:
# IBM_LSF,
# LINUX, LINUX_LSF, LINUX_PBS,
# DARWIN_PGI
```

The option ARCH selects the machine architecture. It is a function of platform type and batch queuing system. The option GSIPROC sets the number of cores used in the run. This option also decides if the job is run as a multiple core job or as a single core run. Several choices of the option ARCH are listed in the sample run script. Please check with your system administrator about running parallel MPI jobs on your system.

3.2.2.3 Setting Up an Analysis Case

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Option ARCH	Platform	Compiler	batch queuing system
IBM_LSF	IBM AIX	xlf, xlc	LSF
LINUX	Linux workstation	Intel/PGI/GNU	mpirun if GSIPROC > 1
LINUX_LSF	Linux cluster	Intel/PGI/GNU	LSF
LINUX_PBS	Linux cluster	Intel/PGI/GNU	PBS
DARWIN_PGI	MAC DARWIN	PGI	mpirun if GSIPROC > 1

This section discusses setting up variables specific to a given case, such as analysis time, working directory, background and observation files, location of fixed files and CRTM coefficients, the GSI executable file, and the script generating GSI namelist.

```
#####
# case set up (users should change this part)
#####
#
# ANAL_TIME= analysis time (YYYYMMDDHH)
# WORK_ROOT= working directory, where GSI runs
# PREPBURF = path of PreBUFR conventional obs
# BK_FILE = path and name of background file
# OBS_ROOT = path of observations files
# FIX_ROOT = path of fix files
# GSI_EXE = path and name of the gsi executable
# ENS_ROOT = path where ensemble background files exist
ANAL_TIME=2017051318
JOB_DIR=the_job_directory
    #normally you put run scripts here and submit jobs form here,
    #require a copy of gsi.x at this directory
RUN_NAME=a_descriptive_run_name_such_as_case05_3denvar_etc
OBS_ROOT=the_directory_where_observation_files_are_located
BK_ROOT=the_directory_where_background_files_are_located
GSI_ROOT=the_comgsi_main_directory_where_src/ ush/ fix/ etc are located
CRTM_ROOT=the_CRTM_directory
ENS_ROOT=the_directory_where_ensemble_backgrounds_are_located
    #ENS_ROOT is not required if not running hybrid EnVAR
HH='echo $ANAL_TIME | cut -c9-10'
GSI_EXE=${JOB_DIR}/gsi.x #assume you have a copy of gsi.x here
WORK_ROOT=${JOB_DIR}/${RUN_NAME}
FIX_ROOT=${GSI_ROOT}/fix
GSI_NAMELIST=${GSI_ROOT}/ush/comgsi_namelist.sh
PREPBURF=${OBS_ROOT}/nam.t${HH}z.prepbufr.tm00
BK_FILE=${BK_ROOT}/wrfinput_d01.${ANAL_TIME}
```

When picking the observation BUFR files, please be aware of the following:

- GSI run will stop if the time in the background file does not match the cycle time in the observation BUFR file used for the GSI run (there is a namelist option to turn this verification step off).
- Since release version 3.2, the BUFRLIB in the GSI package compiled with PGI and Intel can automatically handle byte order issues in PrepBUFR and BUFR files. The BUFRLIB in this version GSI can works with GNU also.

The next part of this block focuses on additional options that specify important aspects of

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the GSI configuration.

```
#-----
# bk_core= which WRF core is used as background (NMM or ARW or NMMB)
# bkcvc_option= which background error covariance and parameter will be used
#                (GLOBAL or NAM)
# if_clean = clean  : delete temporal files in working directory (default)
#                no   : leave running directory as is (this is for debug only)
# if_observer = Yes  : only used as observation operator for enkf
# if_hybrid   = Yes  : Run GSI as 3D/4D EnVar
# if_4DEnVar = Yes  : Run GSI as 4D EnVar
# if_nemsio   = Yes  : The GFS background files are in NEMSIO format
# if_oneob    = Yes  : Do single observation test
if_hybrid=No    # Yes, or, No -- case sensitive !
if_4DEnVar=No   # Yes, or, No -- case sensitive (set if_hybrid=Yes first)!
if_observer=No  # Yes, or, No -- case sensitive !
if_nemsio=No    # Yes, or, No -- case sensitive !
if_oneob=No     # Yes, or, No -- case sensitive !

bk_core=ARW
bkcvc_option=NAM
if_clean=clean
#
# setup whether to do single obs test
if [ ${if_oneob} = Yes ]; then
    if_oneobtest='.true.'
else
    if_oneobtest='.false.'
fi
#
# setup for GSI 3D/4D EnVar hybrid
if [ ${if_hybrid} = Yes ] ; then
    PDYa='echo $ANAL_TIME | cut -c1-8'
    cyca='echo $ANAL_TIME | cut -c9-10'
    gdate='date -u -d "$PDYa $cyca -6 hour" +%Y%m%d%H' #guess date is 6hr ago
    gHH='echo $gdate |cut -c9-10'
    datem1='date -u -d "$PDYa $cyca -1 hour" +%Y-%m-%d_%H:%M:%S' #1hr ago
    datep1='date -u -d "$PDYa $cyca 1 hour" +%Y-%m-%d_%H:%M:%S' #1hr later
    if [ ${if_nemsio} = Yes ]; then
        if_gfs_nemsio='.true.'
        ENSEMBLE_FILE_mem=${ENS_ROOT}/gdas.t${gHH}z.atmf006s.mem
    else
        if_gfs_nemsio='.false.'
        ENSEMBLE_FILE_mem=${ENS_ROOT}/sfg_${gdate}_fhr06s_mem
    fi

    if [ ${if_4DEnVar} = Yes ] ; then
        BK_FILE_P1=${BK_ROOT}/wrfout_d01_${datep1}
        BK_FILE_M1=${BK_ROOT}/wrfout_d01_${datem1}

        if [ ${if_nemsio} = Yes ]; then
            ENSEMBLE_FILE_mem_p1=${ENS_ROOT}/gdas.t${gHH}z.atmf009s.mem
            ENSEMBLE_FILE_mem_m1=${ENS_ROOT}/gdas.t${gHH}z.atmf003s.mem
        else
            ENSEMBLE_FILE_mem_p1=${ENS_ROOT}/sfg_${gdate}_fhr09s_mem
            ENSEMBLE_FILE_mem_m1=${ENS_ROOT}/sfg_${gdate}_fhr03s_mem
        fi
    fi
fi
```

3. Running GSI

```
fi
fi

# The following two only apply when if_observer = Yes, i.e. run observation operator for EnKF
# no_member      number of ensemble members
# BK_FILE_mem    path and base for ensemble members
no_member=20
BK_FILE_mem=${BK_ROOT}/wrfarw.mem
#
```

Option `if_hybrid` controls whether to run a hybrid ensemble/variational data analysis. If `if_hybrid=Yes`, option `if_4DEnVar=Yes` indicates a hybrid 4D-EnVar analysis will be run, while `if_4DEnVar=No` indicates a hybrid 3DENVAR analysis will be run. Option `if_observer` determines whether GSI is run as an observation operator for EnKF.

Option `bk_core` indicates the specific dynamic core used to create the background files and specifies the core in the namelist. Option `bk_core` can be ARW or NMMB. Option `bkcv_option` specifies the background error covariance to be used in the case. Two regional background error covariance matrices are provided with the release, one from NCEP global data assimilation (GDAS), and one from the NAM data assimilation system (NDAS). Please check Section 4.8 for more details about GSI background error covariance. Option `if_clean` tells the script if it needs to delete temporary intermediate files in the working directory after a GSI run is completed.

In most cases, users should only make minor changes after the following:

```
#####
# Users should NOT change script after this point
#####
#
BYTE_ORDER=Big_Endian
# BYTE_ORDER=Little_Endian
```

3.2.3 Description of the Sample Regional Run Script to Run GSI

Listed below is an annotated regional run script with explanations on each function block.

For further details on the first three blocks of the script that users need to change, see sections 3.2.2.1, 3.2.2.2, and 3.2.2.3:

```
#!/bin/ksh
#####
# machine set up (users should change this part)
#####

set -x
#
# GSIPROC = processor number used for GSI analysis
#-----
```

3. Running GSI

```
GSIPROC=1
ARCH='LINUX_LSF'

# Supported configurations:
#   IBM_LSF,
#   LINUX, LINUX_LSF, LINUX_PBS,
#   DARWIN_PGI
#
#####
# case set up (users should change this part)
#####
#
# ANAL_TIME= analysis time (YYYYMMDDHH)
# WORK_ROOT= working directory, where GSI runs
# PREPBURF = path of PreBUFR conventional obs
# BK_FILE   = path and name of background file
# OBS_ROOT  = path of observations files
# FIX_ROOT  = path of fix files
# GSI_EXE   = path and name of the gsi executable
# ENS_ROOT  = path where ensemble background files exist
ANAL_TIME=2017051318
JOB_DIR=the_job_directory
#normally you put run scripts here and submit jobs form here, require a copy of gsi.x at this d
RUN_NAME=a_descriptive_run_name_such_as_case05_3denvar_etc
OBS_ROOT=the_directory_where_observation_files_are_located
BK_ROOT=the_directory_where_background_files_are_located
GSI_ROOT=the_comgsi_main_directory_where_src/ ush/ fix/ etc are located
CRTM_ROOT=the_CRTM_directory
ENS_ROOT=the_directory_where_ensemble_backgrounds_are_located
#ENS_ROOT is not required if not running hybrid EnVAR
HH='echo $ANAL_TIME | cut -c9-10'
GSI_EXE=${JOB_DIR}/gsi.x #assume you have a copy of gsi.x here
WORK_ROOT=${JOB_DIR}/${RUN_NAME}
FIX_ROOT=${GSI_ROOT}/fix
GSI_NAMELIST=${GSI_ROOT}/ush/comgsi_namelist.sh
PREPBURF=${OBS_ROOT}/nam.t${HH}z.prepbufr.tm00
BK_FILE=${BK_ROOT}/wrfinput_d01.${ANAL_TIME}
#
#-----
# bk_core= which WRF core is used as background (NMM or ARW or NMMB)
# bkcvc_option= which background error covariance and parameter will be used
#               (GLOBAL or NAM)
# if_clean = clean : delete temperal files in working directory (default)
#            no    : leave running directory as is (this is for debug only)
# if_observer = Yes : only used as observation operater for enkf
# if_hybrid   = Yes : Run GSI as 3D/4D EnVar
# if_4DEnVar  = Yes : Run GSI as 4D EnVar
# if_nemsio   = Yes : The GFS background files are in NEMSIO format
# if_oneob    = Yes : Do single observation test
if_hybrid=No # Yes, or, No -- case sensitive !
if_4DEnVar=No # Yes, or, No -- case sensitive (set if_hybrid=Yes first)!
if_observer=No # Yes, or, No -- case sensitive !
if_nemsio=No # Yes, or, No -- case sensitive !
if_oneob=No # Yes, or, No -- case sensitive !

bk_core=ARW
bkcvc_option=NAM
if_clean=clean
```


3. Running GSI

```
#
# setup whether to do single obs test
if [ ${if_oneob} = Yes ]; then
    if_oneobtest='.true.'
else
    if_oneobtest='.false.'
fi
#
# setup for GSI 3D/4D EnVar hybrid
if [ ${if_hybrid} = Yes ] ; then
    PDYa='echo $ANAL_TIME | cut -c1-8'
    cyca='echo $ANAL_TIME | cut -c9-10'
    gdate='date -u -d "$PDYa $cyca -6 hour" +%Y%m%d%H' #guess date is 6hr ago
    gHH='echo $gdate |cut -c9-10'
    datem1='date -u -d "$PDYa $cyca -1 hour" +%Y-%m-%d_%H:%M:%S' #1hr ago
    datep1='date -u -d "$PDYa $cyca 1 hour" +%Y-%m-%d_%H:%M:%S' #1hr later
    if [ ${if_nemsio} = Yes ]; then
        if_gfs_nemsio='.true.'
        ENSEMBLE_FILE_mem=${ENS_ROOT}/gdas.t${gHH}z.atmf006s.mem
    else
        if_gfs_nemsio='.false.'
        ENSEMBLE_FILE_mem=${ENS_ROOT}/sfg_${gdate}_fhr06s_mem
    fi

    if [ ${if_4DEnVar} = Yes ] ; then
        BK_FILE_P1=${BK_ROOT}/wrfout_d01_${datep1}
        BK_FILE_M1=${BK_ROOT}/wrfout_d01_${datem1}

        if [ ${if_nemsio} = Yes ]; then
            ENSEMBLE_FILE_mem_p1=${ENS_ROOT}/gdas.t${gHH}z.atmf009s.mem
            ENSEMBLE_FILE_mem_m1=${ENS_ROOT}/gdas.t${gHH}z.atmf003s.mem
        else
            ENSEMBLE_FILE_mem_p1=${ENS_ROOT}/sfg_${gdate}_fhr09s_mem
            ENSEMBLE_FILE_mem_m1=${ENS_ROOT}/sfg_${gdate}_fhr03s_mem
        fi
    fi
fi

# The following two only apply when if_observer = Yes, i.e. run observation operator for EnKF
# no_member      number of ensemble members
# BK_FILE_mem    path and base for ensemble members
no_member=20
BK_FILE_mem=${BK_ROOT}/wrfarw.mem
#
#
```

At this point, users should be able to run the GSI for simple cases without changing the scripts. However, some advanced users may need to change some of the following blocks for special applications, such as use of radiance data, cycled runs, specifying certain namelist variables, or running GSI on a platform not tested by the DTC.

```
#####
# Users should NOT change script after this point
#####
```

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The next block sets the run command for GSI on multiple platforms. The ARCH variable is set at the beginning of the script. Option BYTE_ORDER has been set as Big_Endian because GSI compiled to read a Big_Endian background error file, BUFR files, and CRTM coefficient files.

```
#####
# Users should NOT make changes after this point
#####
#
BYTE_ORDER=Big_Endian
# BYTE_ORDER=Little_Endian

case $ARCH in
  'IBM_LSF')
    ##### IBM LSF (Load Sharing Facility)
    RUN_COMMAND="mpirun.lsf " ;;

  'LINUX')
    if [ $GSIPROC = 1 ]; then
      ### Linux workstation - single processor
      RUN_COMMAND=""
    else
      ##### Linux workstation - mpi run
      RUN_COMMAND="mpirun -np ${GSIPROC} -machinefile ~/mach "
    fi ;;

  'LINUX_LSF')
    ##### LINUX LSF (Load Sharing Facility)
    RUN_COMMAND="mpirun.lsf " ;;

  'LINUX_PBS')
    ##### Linux cluster PBS (Portable Batch System)
    RUN_COMMAND="mpirun -np ${GSIPROC} " ;;

  'DARWIN_PGI')
    ### Mac - mpi run
    if [ $GSIPROC = 1 ]; then
      ### Mac workstation - single processor
      RUN_COMMAND=""
    else
      ##### Mac workstation - mpi run
      RUN_COMMAND="mpirun -np ${GSIPROC} -machinefile ~/mach "
    fi ;;

  * )
    print "error: $ARCH is not a supported platform configuration."
    exit 1 ;;
esac
```

The next block checks if all the variables needed for a GSI run are properly defined. These variables should have been defined in the first three parts of this script.

```
#####
# Check GSI needed environment variables are defined and exist
#
```

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```
# Make sure ANAL_TIME is defined and in the correct format
if [ ! "${ANAL_TIME}" ]; then
    echo "ERROR: \${ANAL_TIME} is not defined!"
    exit 1
fi

# Make sure WORK_ROOT is defined and exists
if [ ! "${WORK_ROOT}" ]; then
    echo "ERROR: \${WORK_ROOT} is not defined!"
    exit 1
fi

# Make sure the background file exists
if [ ! -r "${BK_FILE}" ]; then
    echo "ERROR: \${BK_FILE} does not exist!"
    exit 1
fi

# Make sure OBS_ROOT is defined and exists
if [ ! "${OBS_ROOT}" ]; then
    echo "ERROR: \${OBS_ROOT} is not defined!"
    exit 1
fi
if [ ! -d "${OBS_ROOT}" ]; then
    echo "ERROR: OBS_ROOT directory '\${OBS_ROOT}' does not exist!"
    exit 1
fi

# Set the path to the GSI static files
if [ ! "${FIX_ROOT}" ]; then
    echo "ERROR: \${FIX_ROOT} is not defined!"
    exit 1
fi
if [ ! -d "${FIX_ROOT}" ]; then
    echo "ERROR: fix directory '\${FIX_ROOT}' does not exist!"
    exit 1
fi

# Set the path to the CRTM coefficients
if [ ! "${CRTM_ROOT}" ]; then
    echo "ERROR: \${CRTM_ROOT} is not defined!"
    exit 1
fi
if [ ! -d "${CRTM_ROOT}" ]; then
    echo "ERROR: fix directory '\${CRTM_ROOT}' does not exist!"
    exit 1
fi

# Make sure the GSI executable exists
if [ ! -x "${GSI_EXE}" ]; then
    echo "ERROR: \${GSI_EXE} does not exist!"
    exit 1
fi

# Check to make sure the number of processors for running GSI was specified
if [ -z "${GSIPROC}" ]; then
    echo "ERROR: The variable $GSIPROC must be set to contain the number of processors to run GSI"
    exit 1
fi
```

The next block creates a working directory (workdir) in which GSI will run. The directory should have enough disk space to hold all the files needed for this run. This directory is cleaned before each run, therefore, save all the files needed from the previous run before rerunning GSI.

#####

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```
# Create the work directory and cd into it

workdir=${WORK_ROOT}
echo " Create working directory:" ${workdir}

if [ -d "${workdir}" ]; then
  rm -rf ${workdir}
fi
mkdir -p ${workdir}
cd ${workdir}

#
#####

echo " Copy GSI executable, background file, and link observation bufr to working directory"

# Save a copy of the GSI executable in the workdir
cp ${GSI_EXE} gsi.exe

# Bring over background field (it's modified by GSI so we can't link to it)
cp ${BK_FILE} ./wrf_inout
if [ ${if_4DnVar} = Yes ] ; then
  cp ${BK_FILE_P1} ./wrf_inou3
  cp ${BK_FILE_M1} ./wrf_inou1
fi
```

Note: You can link observation files to the working directory because GSI will not overwrite these files. The observations that can be analyzed in GSI are listed in the column "dfile" of the GSI namelist section OBS_INPUT, as specified in run/comgsi_namelist.sh. Most of the conventional observations are in one single file named prepbufr, while different radiance data are in separate files based on satellite instruments, such as AMSU-A or HIRS. All these observation files must be linked as GSI recognized file names in "dfile." Please check table 3.1 for a detailed explanation of links and the meanings of each file name listed below.

```
# Link to the prepbufr data
ln -s ${PREPBUFR} ./prepbufr

# ln -s ${OBS_ROOT}/gdas1.t${HH}z.sptrmm.tm00.bufr_d tmirrbufr
# Link to the radiance data
srcobsfile[1]=${OBS_ROOT}/gdas1.t${HH}z.satwnd.tm00.bufr_d
gsiobsfile[1]=satwnd
srcobsfile[2]=${OBS_ROOT}/gdas1.t${HH}z.1bamua.tm00.bufr_d
gsiobsfile[2]=amsuabufr
srcobsfile[3]=${OBS_ROOT}/gdas1.t${HH}z.1bhrs4.tm00.bufr_d
gsiobsfile[3]=hirs4bufr
srcobsfile[4]=${OBS_ROOT}/gdas1.t${HH}z.1bmhs.tm00.bufr_d
gsiobsfile[4]=mhsbufr
srcobsfile[5]=${OBS_ROOT}/gdas1.t${HH}z.1bamub.tm00.bufr_d
gsiobsfile[5]=amsubbufr
srcobsfile[6]=${OBS_ROOT}/gdas1.t${HH}z.ssmisu.tm00.bufr_d
gsiobsfile[6]=ssmirrbufr
# srcobsfile[7]=${OBS_ROOT}/gdas1.t${HH}z.airsev.tm00.bufr_d
gsiobsfile[7]=airsbufr
srcobsfile[8]=${OBS_ROOT}/gdas1.t${HH}z.sevcsr.tm00.bufr_d
gsiobsfile[8]=seviribufr
srcobsfile[9]=${OBS_ROOT}/gdas1.t${HH}z.iasidb.tm00.bufr_d
gsiobsfile[9]=iasibufr
srcobsfile[10]=${OBS_ROOT}/gdas1.t${HH}z.gpsro.tm00.bufr_d
gsiobsfile[10]=gpsrobufr
srcobsfile[11]=${OBS_ROOT}/gdas1.t${HH}z.amsr2.tm00.bufr_d
```

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```
gsiobsfile[11]=amsrebufr
srcobsfile[12]=${OBS_ROOT}/gdas1.t${HH}z.atms.tm00.bufr_d
gsiobsfile[12]=atmsbufr
srcobsfile[13]=${OBS_ROOT}/gdas1.t${HH}z.geoimr.tm00.bufr_d
gsiobsfile[13]=gimgrbufr
srcobsfile[14]=${OBS_ROOT}/gdas1.t${HH}z.gome.tm00.bufr_d
gsiobsfile[14]=gomebufr
srcobsfile[15]=${OBS_ROOT}/gdas1.t${HH}z.omi.tm00.bufr_d
gsiobsfile[15]=omibufr
srcobsfile[16]=${OBS_ROOT}/gdas1.t${HH}z.osbuv8.tm00.bufr_d
gsiobsfile[16]=sbuvbufr
srcobsfile[17]=${OBS_ROOT}/gdas1.t${HH}z.eshrs3.tm00.bufr_d
gsiobsfile[17]=hirs3bufrears
srcobsfile[18]=${OBS_ROOT}/gdas1.t${HH}z.esamua.tm00.bufr_d
gsiobsfile[18]=amsuabufrears
srcobsfile[19]=${OBS_ROOT}/gdas1.t${HH}z.esmhs.tm00.bufr_d
gsiobsfile[19]=mhsbufrears
srcobsfile[20]=${OBS_ROOT}/rap.t${HH}z.nexrad.tm00.bufr_d
gsiobsfile[20]=l2rwbufr
srcobsfile[21]=${OBS_ROOT}/rap.t${HH}z.lgyclld.tm00.bufr_d
gsiobsfile[21]=larcglb
ii=1
while [[ $ii -le 21 ]]; do
    if [ -r "${srcobsfile[$ii]}" ]; then
        ln -s ${srcobsfile[$ii]} ${gsiobsfile[$ii]}
        echo "link source obs file ${srcobsfile[$ii]}"
    fi
    (( ii = $ii + 1 ))
done
```

The following block copies constant fixed files from the fix/ directory and links CRTM coefficients. Please check Section 3.1 for the meanings of each fixed file.

```
#####
echo " Copy fixed files and link CRTM coefficient files to working directory"

# Set fixed files
# berror = forecast model background error statistics
# specoef = CRTM spectral coefficients
# trncoef = CRTM transmittance coefficients
# emiscoef = CRTM coefficients for IR sea surface emissivity model
# aerocoeff = CRTM coefficients for aerosol effects
# cldcoef = CRTM coefficients for cloud effects
# satinfo = text file with information about assimilation of brightness temperatures
# satangl = angle dependent bias correction file (fixed in time)
# pcpinfo = text file with information about assimilation of precpitation rates
# ozinfo = text file with information about assimilation of ozone data
# errtable = text file with obs error for conventional data (regional only)
# convinfo = text file with information about assimilation of conventional data
# bufrtable= text file ONLY needed for single obs test (oneobstest=.true.)
# bftab_sst= bufr table for sst ONLY needed for sst retrieval (retrieval=.true.)
```

Note: For background error covariances, observation errors, and analysis variable information, we provide two sets of fixed files. One set is based on GFS statistics and another is

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based on NAM statistics. For this release there is an additional setting in the ANAVINFO file for "bk_core" for both GFS and NAM statistics.

```
if [ ${bkcv_option} = GLOBAL ] ; then
  echo ' Use global background error covariance'
  BERROR=${FIX_ROOT}/${BYTE_ORDER}/nam_glb_berror.f77.gcv
  OBERROR=${FIX_ROOT}/prepobs_errtable.global
  if [ ${bk_core} = NMM ] ; then
    ANAVINFO=${FIX_ROOT}/anavinfo_ndas_netcdf_glbe
  fi
  if [ ${bk_core} = ARW ] ; then
    ANAVINFO=${FIX_ROOT}/anavinfo_arw_netcdf_glbe
  fi
  if [ ${bk_core} = NMMB ] ; then
    ANAVINFO=${FIX_ROOT}/anavinfo_nems_nmm_b_glb
  fi
else
  echo ' Use NAM background error covariance'
  BERROR=${FIX_ROOT}/${BYTE_ORDER}/nam_nmmstat_na.gcv
  OBERROR=${FIX_ROOT}/nam_errtable.r3dv
  if [ ${bk_core} = NMM ] ; then
    ANAVINFO=${FIX_ROOT}/anavinfo_ndas_netcdf
  fi
  if [ ${bk_core} = ARW ] ; then
    ANAVINFO=${FIX_ROOT}/anavinfo_arw_netcdf
  fi
  if [ ${bk_core} = NMMB ] ; then
    ANAVINFO=${FIX_ROOT}/anavinfo_nems_nmm_b
  fi
fi

SATINFO=${FIX_ROOT}/global_satinfo.txt
CONVINFO=${FIX_ROOT}/global_convinfo.txt
OZINFO=${FIX_ROOT}/global_ozinfo.txt
PCPINFO=${FIX_ROOT}/global_pcpinfo.txt

# copy Fixed fields to working directory
cp $ANAVINFO anavinfo
cp $BERROR berror_stats
cp $SATINFO satinfo
cp $CONVINFO convinfo
cp $OZINFO ozinfo
cp $PCPINFO pcpinfo
cp $OBERROR errtable

#
# # CRTM Spectral and Transmittance coefficients
CRTM_ROOT_ORDER=${CRTM_ROOT}/${BYTE_ORDER}
emiscoef_IRwater=${CRTM_ROOT_ORDER}/Nalli.IRwater.EmisCoeff.bin
emiscoef_IRice=${CRTM_ROOT_ORDER}/NPOESS.IRice.EmisCoeff.bin
emiscoef_IRland=${CRTM_ROOT_ORDER}/NPOESS.IRland.EmisCoeff.bin
emiscoef_IRsnow=${CRTM_ROOT_ORDER}/NPOESS.IRsnow.EmisCoeff.bin
emiscoef_VISice=${CRTM_ROOT_ORDER}/NPOESS.VISice.EmisCoeff.bin
emiscoef_VISland=${CRTM_ROOT_ORDER}/NPOESS.VISland.EmisCoeff.bin
emiscoef_VISsnow=${CRTM_ROOT_ORDER}/NPOESS.VISsnow.EmisCoeff.bin
emiscoef_VISwater=${CRTM_ROOT_ORDER}/NPOESS.VISwater.EmisCoeff.bin
emiscoef_MWwater=${CRTM_ROOT_ORDER}/FASTEM6.MWwater.EmisCoeff.bin
aercoef=${CRTM_ROOT_ORDER}/AerosolCoeff.bin
```

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```
cldcoef=${CRTM_ROOT_ORDER}/CloudCoeff.bin

ln -s $emiscoef_IRwater ./Nalli.IRwater.EmisCoeff.bin
ln -s $emiscoef_IRrice ./NPOESS.IRrice.EmisCoeff.bin
ln -s $emiscoef_IRsnow ./NPOESS.IRsnow.EmisCoeff.bin
ln -s $emiscoef_IRland ./NPOESS.IRland.EmisCoeff.bin
ln -s $emiscoef_VISice ./NPOESS.VISice.EmisCoeff.bin
ln -s $emiscoef_VISland ./NPOESS.VISland.EmisCoeff.bin
ln -s $emiscoef_VISsnow ./NPOESS.VISsnow.EmisCoeff.bin
ln -s $emiscoef_VISwater ./NPOESS.VISwater.EmisCoeff.bin
ln -s $emiscoef_MWwater ./FASTEM6.MWwater.EmisCoeff.bin
ln -s $aercoef ./AerosolCoeff.bin
ln -s $cldcoef ./CloudCoeff.bin
# Copy CRTM coefficient files based on entries in satinfo file
for file in `awk '{if($1~"!"){print $1}}' ./satinfo | sort | uniq` ;do
    ln -s ${CRTM_ROOT_ORDER}/${file}.SpcCoeff.bin ./
    ln -s ${CRTM_ROOT_ORDER}/${file}.TauCoeff.bin ./
done

# Only need this file for single obs test
bufhtable=${FIX_ROOT}/prepobs_prep.bufhtable
cp $bufhtable ./prepobs_prep.bufhtable

# for satellite bias correction
# Users may need to use their own satbias files for correct bias correction
cp ${GSI_ROOT}/fix/comgsi_satbias_in ./satbias_in
cp ${GSI_ROOT}/fix/comgsi_satbias_pc_in ./satbias_pc_in
```

Please note that in the above sample script, two files related to radiance bias correction are copied to the work directory:

```
cp ${GSI_ROOT}/fix/comgsi_satbias_in ./satbias_in
cp ${GSI_ROOT}/fix/comgsi_satbias_pc_in ./satbias_pc_in
```

There are two options on how to perform the radiance bias correction. The first method is to do the angle dependent bias correction offline and do the mass bias correction inside the GSI analysis, therefore requiring two input files: `satbias_angle`, corresponding to the angle dependent bias correction file and `satbias_in`, being the input file for mass bias correction. The second method is to combine the angle dependent and mass bias correction together and do it within the GSI analysis, requiring one combined input file: `satbias_in`. Note that the input bias correction coefficients file, `satbias_in`, is different for the two options, therefore it is important to use the appropriate input file for each method. This version GSI only provide the second option - combined angle dependent and mass bias correction. A sample file, `comgsi_satbias_in`, is provided as an example. As a starting point, users may also download a GDAS `satbias` coefficient file from the NOMADS ftp site as the input file (starting in spring 2015, the GDAS `satbias` files have adopted the following format):

```
ftp://nomads.ncdc.noaa.gov/GDAS/YYYYMM/YYYYMMDD/gdas1.tHHz.abias
```

In order to use the combined angle dependent and mass bias correction, users also need to set `adp_anglebc=.true.` in the `&SETUP` section of the GSI namelist

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(comgsi_namelist.sh). For more details about the namelist, please see Appendix C in this document.

Set up some constants used in the GSI namelist. Please note that `bkcv_option` is set for background error tuning. They should be set based on specific applications. Here we provide three sample sets of the constants for different background error covariance options, one set is used in the NAM operations, one for the GFS operations and one for the NMMB operations. In this release, the capability of NMMB application is included and therefore the namelist settings for NMMB are provided in addition to NMM and ARW applications.

```
#####
# Set some parameters for use by the GSI executable and to build the namelist
echo " Build the namelist "

# default is NAM
# as_op='1.0,1.0,0.5 ,0.7,0.7,0.5,1.0,1.0,'
vs_op='1.0,'
hzscl_op='0.373,0.746,1.50,'
if [ ${bkcv_option} = GLOBAL ] ; then
# as_op='0.6,0.6,0.75,0.75,0.75,0.75,1.0,1.0'
vs_op='0.7,'
hzscl_op='1.7,0.8,0.5,'
fi
if [ ${bk_core} = NMMB ] ; then
vs_op='0.6,'
fi

# default is NMM
bk_core_arw='.false.'
bk_core_nmm='.true.'
bk_core_nmb='.false.'
bk_if_netcdf='.true.'
if [ ${bk_core} = ARW ] ; then
bk_core_arw='.true.'
bk_core_nmm='.false.'
bk_core_nmb='.false.'
bk_if_netcdf='.true.'
fi
if [ ${bk_core} = NMMB ] ; then
bk_core_arw='.false.'
bk_core_nmm='.false.'
bk_core_nmb='.true.'
bk_if_netcdf='.false.'
fi
```

The following section specifies the number of outer loops and whether to save GSI read observations based on the setting of "if_observer".

```
if [ ${if_observer} = Yes ] ; then
nummiter=0
if_read_obs_save='.true.'
if_read_obs_skip='.false.'
```


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```
else
  nummiter=2
  if_read_obs_save='.false.'
  if_read_obs_skip='.false.'
fi
```

The following section of the script is used to generate the GSI namelist called `gsiparm.anl` in the working directory. A detailed explanation of each variable can be found in Section 3.4 and Appendix C.

```
# Build the GSI namelist on-the-fly
. $GSI_NAMELIST
```

The following block modifies the `anavinfo` file so that its vertical levels are consistent with the `wrf_inout` file for WRF ARW or NMM. Users no longer need to manually modify the `anavinfo` file.

```
# modify the anavinfo vertical levels based on wrf_inout for WRF ARW and NMM
if [ ${bk_core} = ARW ] || [ ${bk_core} = NMM ] ; then
bklevels='ncdump -h wrf_inout | grep "bottom_top =" | awk '{print $3}' '
bklevels_stag='ncdump -h wrf_inout | grep "bottom_top_stag =" | awk '{print $3}' '
anavlevels='cat anavinfo | grep ' sf ' | tail -1 | awk '{print $2}' ' # levels of sf, vp, u, v, t,
anavlevels_stag='cat anavinfo | grep ' prse ' | tail -1 | awk '{print $2}' ' # levels of prse
sed -i 's/ '$anavlevels'/ '$bklevels'/g' anavinfo
sed -i 's/ '$anavlevels_stag'/ '$bklevels_stag'/g' anavinfo
fi
```

The following block runs GSI and checks if GSI has successfully completed.

```
#####
# run GSI
#####
echo ' Run GSI with' ${bk_core} 'background'

case $ARCH in
  'IBM_LSF')
    ${RUN_COMMAND} ./gsi.exe < gsiparm.anl > stdout 2>&1 ;;

  * )
    ${RUN_COMMAND} ./gsi.exe > stdout 2>&1 ;;
esac

#####
# run time error check
#####
error=$?

if [ ${error} -ne 0 ]; then
  echo "ERROR: ${GSI} crashed Exit status=${error}"
  exit ${error}
fi
```

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The following block saves the analysis results with an understandable name and adds the analysis time to some output file names. Among them, "stdout" contains runtime output of GSI and wrf_inout is the resulting analysis file.

```
#####  
#  
#   GSI updating satbias_in  
#  
# GSI updating satbias_in (only for cycling assimilation)  
  
# Copy the output to more understandable names  
ln -s stdout      stdout.anl.${ANAL_TIME}  
ln -s wrf_inout   wrfanl.${ANAL_TIME}  
ln -s fort.201    fit_p1.${ANAL_TIME}  
ln -s fort.202    fit_w1.${ANAL_TIME}  
ln -s fort.203    fit_t1.${ANAL_TIME}  
ln -s fort.204    fit_q1.${ANAL_TIME}  
ln -s fort.207    fit_rad1.${ANAL_TIME}
```

The following block collects the diagnostic files. The diagnostic files are merged and categorized based on outer loop and data type. Setting "write_diag" to true in the namelist directs GSI to write out diagnostic information for each observation. This information is very useful to check analysis details. Please check Appendix A.2 for the tool to read and analyze these diagnostic files.

```
# Loop over first and last outer loops to generate innovation  
# diagnostic files for indicated observation types (groups)  
#  
# NOTE: Since we set miter=2 in GSI namelist SETUP, outer  
#       loop 03 will contain innovations with respect to  
#       the analysis. Creation of o-a innovation files  
#       is triggered by write_diag(3)=.true. The setting  
#       write_diag(1)=.true. turns on creation of o-g  
#       innovation files.  
#  
loops="01 03"  
for loop in $loops; do  
  
case $loop in  
  01) string=ges;;  
  03) string=anl;;  
  *) string=$loop;;  
esac  
  
# Collect diagnostic files for obs types (groups) below  
# listall="conv amsua_metop-a mhs_metop-a hirs4_metop-a hirs2_n14 msu_n14 \  
#         sndr_g08 sndr_g10 sndr_g12 sndr_g08_prep sndr_g10_prep sndr_g12_prep \  
#         sndrd1_g08 sndrd2_g08 sndrd3_g08 sndrd4_g08 sndrd1_g10 sndrd2_g10 \  
#         sndrd3_g10 sndrd4_g10 sndrd1_g12 sndrd2_g12 sndrd3_g12 sndrd4_g12 \  
#         hirs3_n15 hirs3_n16 hirs3_n17 amsua_n15 amsua_n16 amsua_n17 \  
#         amsub_n15 amsub_n16 amsub_n17 hsb_aqua airs_aqua amsua_aqua \  
#         goes_img_g08 goes_img_g10 goes_img_g11 goes_img_g12 \  
#         pcp_ssmi_dmsp pcp_tmi_trmm sbuv2_n16 sbuv2_n17 sbuv2_n18 \  
#         "
```

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```
#      omi_aura  ssmi_f13  ssmi_f14  ssmi_f15  hirs4_n18  amsua_n18  mhs_n18  \  
#      amsre_low_aqua  amsre_mid_aqua  amsre_hig_aqua  ssmis_las_f16  \  
#      ssmis_uas_f16  ssmis_img_f16  ssmis_env_f16  mhs_metop_b  \  
#      hirs4_metop_b  hirs4_n19  amusa_n19  mhs_n19"  
listall='ls pe* | cut -f2 -d"." | awk '{print substr($0, 0, length($0)-3)}' | sort | uniq'  
  
  for type in $listall; do  
    count='ls pe*${type}_${loop}* | wc -l'  
    if [[ $count -gt 0 ]]; then  
      cat pe*${type}_${loop}* > diag_${type}_${string}.${ANAL_TIME}  
    fi  
  done  
done
```

The following scripts clean the temporary intermediate files:

```
# Clean working directory to save only important files  
ls -l * > list_run_directory  
if [[ ${if_clean} = clean && ${if_observer} != Yes ]]; then  
  echo 'Clean working directory after GSI run'  
  rm -f *Coeff.bin      # all CRTM coefficient files  
  rm -f pe0*           # diag files on each processor  
  rm -f obs_input.*    # observation middle files  
  rm -f siganl sigf03  # background middle files  
  rm -f fsize_*       # delete temperal file for bufr size  
fi
```

The following block of the script runs only for `if_observer=Yes`, which runs GSI as an observation operator for EnKF and without doing minimization. The script first renames the previous diagnostics files and GSI analysis file by appending `.ensmean` to the filenames to avoid these files being overwritten by the new GSI run.

```
#####  
# start to calculate diag files for each member  
#####  
#  
if [ ${if_observer} = Yes ] ; then  
  string=ges  
  for type in $listall; do  
    count=0  
    if [[ -f diag_${type}_${string}.${ANAL_TIME} ]]; then  
      mv diag_${type}_${string}.${ANAL_TIME} diag_${type}_${string}.ensmean  
    fi  
  done  
  mv wrf_inout wrf_inout_ensmean
```

Next, the script generates the namelist for each ensemble member.

```
# Build the GSI namelist on-the-fly for each member  
nummiter=0  
if_read_obs_save='.false.'  
if_read_obs_skip='.true.'  
. $GSI_NAMELIST
```

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The rest of the script loops through the ensemble members to get the background ready, run GSI, and check the run status:

```
# Loop through each member
loop="01"
ensmem=1
while [[ $ensmem -le $no_member ]];do

    rm pe0*

    print "\$ensmem is $ensmem"
    ensmemid='printf %3.3i $ensmem'

# get new background for each member
    if [[ -f wrf_inout ]]; then
        rm wrf_inout
    fi

    BK_FILE=${BK_FILE_mem}${ensmemid}
    echo $BK_FILE
    ln -s $BK_FILE wrf_inout

# run GSI
    echo ' Run GSI with' ${bk_core} 'for member ', ${ensmemid}

    case $ARCH in
        'IBM_LSF')
            ${RUN_COMMAND} ./gsi.exe < gsiparm.anl > stdout_mem${ensmemid} 2>&1 ;;
        * )
            ${RUN_COMMAND} ./gsi.exe > stdout_mem${ensmemid} 2>&1 ;;
    esac

# run time error check and save run time file status
    error=$?

    if [ ${error} -ne 0 ]; then
        echo "ERROR: ${GSI} crashed for member ${ensmemid} Exit status=${error}"
        exit ${error}
    fi

    ls -l * > list_run_directory_mem${ensmemid}

done
```

The following lines generate the diagnostics files for each member.

```
# generate diag files

for type in $listall; do
    count='ls pe*${type}_${loop}* | wc -l'
    if [[ $count -gt 0 ]]; then
        cat pe*${type}_${loop}* > diag_${type}_${string}.mem${ensmemid}
    fi
done
```

3. Running GSI

The following section is to move on to the next ensemble member and run GSI.

```
# next member
  (( ensmem += 1 ))

done

fi
```

If this point is reached, the GSI successfully finishes and exits with status "0":

```
exit 0
```

3.3 GSI Analysis Result Files in Run Directory

Once the GSI run script is set up, it is ready to be submitted like any other batch job. When completed, GSI will create a number of files in the run directory. Below is an example of the files generated in the run directory from one of the GSI test case runs. This case was run to perform a regional GSI analysis with a WRF-ARW NetCDF background using conventional (prepbufr), radiance (AMSU-A, HIRS4, and MHS), and GPSRO data. The analysis time is 1200Z on 13 May 2017. Four processors were used. To make the run directory more readable, we turned on the clean option in the run script, which deleted all temporary intermediate files.

amsuabufr	fort.206	hirs3bufrears
amsuabufrears	fort.207	hirs4buffr
anavinfo	fort.208	l2rwbuffr
atmsbuffr	fort.209	larcglb
berror_stats	fort.210	list_run_directory
convinfo	fort.211	mhsbuffr
diag_amsua_n15_anl.2017051312	fort.212	mhsbufrears
diag_amsua_n15_ges.2017051312	fort.213	omibuffr
diag_amsua_n18_anl.2017051312	fort.214	ozinfo
diag_amsua_n18_ges.2017051312	fort.215	pcpbias_out
diag_amsua_n19_anl.2017051312	fort.217	pcpinfo
diag_amsua_n19_ges.2017051312	fort.218	prepbufr
diag_conv_anl.2017051312	fort.219	prepobs_prep.bufhtable
diag_conv_ges.2017051312	fort.220	radar_supobs_from_level2
diag_hirs4_n19_anl.2017051312	fort.221	satbias_angle
diag_hirs4_n19_ges.2017051312	fort.223	satbias_ang.out
diag_mhs_n18_anl.2017051312	fort.224	satbias_in
diag_mhs_n18_ges.2017051312	fort.225	satbias_out
diag_mhs_n19_anl.2017051312	fort.226	satbias_out.int
diag_mhs_n19_ges.2017051312	fort.227	satbias_pc_in
errtable	fort.228	satbias_pc.out
fit_p1.2017051312	fort.229	satinfo
fit_q1.2017051312	fort.230	satwnd
fit_rad1.2017051312	fort.232	sbuvbuffr
fit_t1.2017051312	fort.233	seviribuffr
fit_w1.2017051312	fort.234	ssmirrbuffr
fort.201	gimgrbuffr	stdout
fort.202	gomebuffr	stdout.anl.2017051312
fort.203	gpsrobuffr	wrfanl.2017051312
fort.204	gsi.exe	wrf_inout
fort.205	gsiparm.anl	

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It is important to know which files hold the GSI analysis results, standard output, and diagnostic information. We will introduce these files and their contents in detail in the following chapter. The following is a brief list of what these files contain:

- *stdout* or *stdout.anl.(time)*: standard text output file. *stdout.anl.(time)* is a link to *stdout* with the analysis time appended. This is the most commonly used file to check the GSI analysis processes and contains basic and important information about the analyses. We will explain the contents of the *stdout* file in Section 4.1 and users are encouraged to read this file in detail to become familiar with the order of GSI analysis processing.
- *wrf_inout* or *wrfanl.(time)*: analysis results if GSI completes successfully. It exists only if using WRF for the background. The *wrfanl.(time)* file is a link to *wrf_inout* with the analysis time appended. The format is the same as the background file.
- *diag_conv_anl.(time)*: binary diagnostic files for conventional and GPS RO observations at the final analysis step (analysis departure for each observation).
- *diag_conv_ges.(time)*: binary diagnostic files for conventional and GPS RO observations before the initial analysis step (background departure for each observation)
- *diag (instrument_satellite)_anl*: diagnostic files for satellite radiance observations at the final analysis step.
- *diag (instrument_satellite)_ges*: diagnostic files for satellite radiance observations before the initial analysis step.
- *gsiparm.anl*: GSI namelist, generated by the run script.
- *fit_(variable).(time)*: links to fort.2?? with meaningful names (variable name plus analysis time). They are statistic results of observation departures from background and analysis results according to observation variables. Please see Section 4.5 for more details.
- *fort.220*: output from the inner loop minimization (in *pcgsoi.f90*). Please see Section 4.6 for details.
- *anavinfo*: info file to set up control, state, and background variables. Please see the Advanced GSI User's Guide for details.
- **info (convinfo,satinfo, ...)*: info files that control data usage. Please see Section 4.3 for details.
- *berror_stats* and *errtable*: background error file (binary) and observation error file (text).
- **bufr*: observation BUFR files linked to the run directory. Please see Section 3.1 for details.
- *satbias_in*: the input coefficients of bias correction for satellite radiance observations.
- *satbias_out*: the output coefficients of bias correction for satellite radiance observations after the GSI run.
- *satbias_pc_in*: the input coefficients of bias correction for passive satellite radiance observations.
- *list_run_directory*: the complete list of files in the run directory before cleaning takes place. This is generated by the GSI run script.

The *diag* files, such as *diag (instrument_satellite)_anl.(time)* and *diag_conv_anl.(time)*, contain important information about the data used in the GSI, including observation departure from analysis results for each observation (O-A). Similarly, *diag_conv_ges* and *diag (instrument_satellite)_ges.(time)* include the observation innovation for each observation (O-B). These files can be very helpful in understanding the detailed impact of data on the analysis. A tool is provided to process

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these files, which is introduced in Appendix A.2.

There are many intermediate files in this directory while GSI is running or if the run crashes. The complete list of files in the directory (prior to cleaning) is saved in file `list_run_directory`. Some knowledge about the content of these files is very helpful for debugging if the GSI run crashes. Please check table 3.4 for the meaning of these files. (Note: you may not see all the files in the list because different observational data are used. Also, the fixed files prepared for a GSI run, such as CRTM coefficient files, are not included.)

Table 3.4: List of GSI intermediate files

File name	Content
<code>sigf03</code>	This is a temporary file, holding binary format background files (typically <code>sigf03</code> , <code>sigf06</code> and <code>sigf09</code> if FGAT used). When you see this file, at the minimum, a background file was successfully read in.
<code>siganl</code>	Analysis results in binary format. When this file exists, the analysis has finished.
<code>pe????.(conv or instru- ment_satellite)_(outer loop)</code>	Diagnostic files for conventional and satellite radiance observations at each outer loop and each sub-domain (????=subdomain id)i.
<code>obs_input.????</code>	Observation scratch files (each file contains observations for one observation type within the whole analysis domain and time window. ????=observation type id in namelist).
<code>pcpbias_out</code>	Output precipitation bias correction file.

3.4 Introduction to Frequently Used GSI Namelist Options

The complete namelist options and their explanations are listed in Appendix C. For most GSI analysis applications, only a few namelist variables need to be changed. Here we introduce frequently used variables for regional analyses:

3.4.1 Set Up the Number of Outer and Inner Loops

To change the number of outer loops and the number of inner iterations in each outer loop, the following three variables in the namelist need to be modified:

- `miter`: number of outer analysis loops.
- `niter(1)`: maximum iteration number of inner loop iterations for the 1st outer loop. The inner loop will stop when it reaches this maximum number, when it reaches the convergence threshold, or when it fails to converge.
- `niter(2)`: maximum iteration number of inner loop iterations for the 2nd outer loop.
- If `miter` is larger than two, repeat `niter` with larger index.

3.4.2 Set Up the Analysis Variable for Moisture

There are two moisture analysis variable options. It is based on the following namelist variable:

`qoption = 1 or 2:`

- If `qoption=1`, the moisture analysis variable is pseudo-relative humidity. The saturation specific humidity, `qsatg`, is computed from the guess and held constant during the inner loop. Thus, the relative humidity control variable can only change via changes in specific humidity, `q`.
- If `qoption=2`, the moisture analysis variable is normalized relative humidity. This formulation allows relative humidity to change in the inner loop via changes to surface pressure, temperature, or specific humidity.

3.4.3 Set Up the Background File

The following four variables define which background field will be used in the GSI analyses:

- `regional`: if true, perform a regional GSI run using either ARW or NMM inputs as the background. If false, perform a global GSI analysis. If either `wrf_nmm_regional` or `wrf_mass_regional` are true, it will be set to true.
- `wrf_nmm_regional`: if true, the background comes from WRF-NMM. When using other background fields, set it to false.
- `wrf_mass_regional`: if true, the background comes from WRF-ARW. When using other background fields, set it to false.
- `nems_nmmb_regional`: if true, the background comes from NMMB. When using other background fields, set it to false.
- `netcdf`: if true, WRF files are in NetCDF format, otherwise WRF files are in binary format. This option only works for a regional GSI analysis.

3.4.4 Set Up the Output of Diagnostic Files

The following variables tell the GSI to write out diagnostic results in certain loops:

- `write_diag(1)`: if true, write out diagnostic data in the beginning of the analysis, so that we can have information on observation – background (O-B) differences.
- `write_diag(2)`: if true, write out diagnostic data at the end of the 1st outer loop (before the 2nd outer loop starts).
- `write_diag(3)`: if true, write out diagnostic data at the end of the 2nd outer loop (after the analysis finishes if the outer loop number is two), so that we can have information on observation – analysis (O-A) differences.

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Please check appendix A.2 for the tools to read the diagnostic files.

3.4.5 Set Up the GSI Recognized Observation Files

The following sets up the GSI recognized observation files for GSI observation ingest:

```
OBS_INPUT::
! dfile          dtype      dplat      dsis          dval      dthin dsfcalc
  prepbuf      ps          null       ps            1.0       0      0
  prepbuf      t          null       t             1.0       0      0
  prepbuf      q          null       q             1.0       0      0
  prepbuf      pw          null       pw            1.0       0      0
  satwndbuf    uv          null       uv            1.0       0      0
  prepbuf      uv          null       uv            1.0       0      0
  prepbuf      spd         null       spd           1.0       0      0
  prepbuf      dw          null       dw            1.0       0      0
  radarbuf     rw          null       rw            1.0       0      0
  prepbuf      sst         null       sst           1.0       0      0
  gpsrobuf     gps_ref    null       gps           1.0       0      0
  ssmirrbufr   pcp_ssmi  dmsp      pcp_ssmi     1.0      -1      0
```

- `dfile`: GSI recognized observation file name. The observation file contains observations used for a GSI analysis. This file can include several observation variables from different observation types. The file name listed by this parameter will be read in by GSI. This name can be changed as long as the name in the link from the BUFR/Prep-BUFR file in the run scripts also changes correspondingly.
- `dtype`: analysis variable name that GSI can read in. Please note this name should be consistent with that used in the GSI code.
- `dplat`: sets up the observation platform for a certain observation, which will be read in from the file `dfile`.
- `dsis`: sets up the data name (including both data type and platform name) used inside GSI.

Please see Section 4.3 for examples and explanations of these variables.

3.4.6 Set Up Observation Time Window

In the namelist section `OBS_INPUT`, use `time_window_max` to set the maximum half time window (hours) for all data types. In the `convinfo` file, you can use the column "twindow" to set the half time window for a certain data type (hours). For conventional observations, only observations within the smaller window of these two will be kept for further processing. For others, observations within `time_window_max` will be kept for further processing.

3.4.7 Set Up Data Thinning

1) Radiance data thinning

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Radiance data thinning is controlled through two GSI namelist variables in the section &OBS_INPUT. Below is an example:

```
&OBS_INPUT
  dmesh(1)=120.0,dmesh(2)=60.0,dmesh(3)=30,time_window_max=1.5,ext_sonde=.true.,
/
OBS_INPUT::
!  dfile      dtype      dplat      dsis      dval      dthin dsfcalc
  prepbuf    ps          null       ps         1.0       0      0

  gpsrobuf   gps_ref     null       gps        1.0       0      0
  ssmirrbufr pcp_ssmi    dmsp      pcp_ssmi   1.0      -1      0
  tmirrbufr  pcp_tmi    trmm      pcp_tmi    1.0      -1      0

  hirs3buf   hirs3       n17       hirs3_n17  6.0       1      0
  hirs4buf   hirs4       metop-a   hirs4_metop-a 6.0       2      0
```

The two namelist variables that control the radiance data thinning are real array "dmesh" in the 1st line and the "dthin" values in the 6th column. The "dmesh" array sets mesh sizes for radiance thinning grids in kilometers, while "dthin" defines if the data type it represents needs to be thinned and which thinning grid (mesh size) to use. If the value of dthin is:

- an integer less than or equal to zero, no thinning is needed
- an integer larger than zero, this kind of radiance data will be thinned using the mesh size defined as dmesh (dthin).

The following section provides several thinning examples defined by the above sample &OBS_INPUT section:

- Data type ps from prepbuf: no thinning because dthin=0
- Data type gps_ref from gpsrobuf: no thinning because dthin=0
- Data type pcp_ssmi from dmsp: no thinning because dthin(01)=-1
- Data type hirs3 from NOAA-17: thinning in a 120 km grid because dthin=1 and dmesh(1)=120
- Data type hirs4 from metop-a: thinning in a 60 km grid because dthin=2 and dmesh(2)=60

2) Conventional data thinning

The conventional data can also be thinned. However, the setup of thinning is not in the namelist. To give users a complete picture of data thinning, conventional data thinning is briefly introduced here. There are three columns, ithin, rmesh, pmesh, in the convinfo file (more details on this file are in Section 4.3) to configure conventional data thinning:

- ithin: 0 = no thinning; 1 = thinning with grid mesh decided by rmesh and pmesh
- rmesh: horizontal thinning grid size in km
- pmesh: vertical thinning grid size in mb; if 0, then use background vertical grid.

3.4.8 Set Up Background Error Factor

In the namelist section BKGERR, `vs` is used to set up the scale factor for vertical correlation length and `hzscl` is defined to set up scale factors for horizontal smoothing. The scale factors for the variance of each analysis variables are set in the `anavinfo` file. The typical values used in operations for regional and global background error covariance are given and picked based on the choice of background error covariance in the run scripts and sample `anavinfo` files

3.4.9 Single Observation Test

To do a single observation test, the following namelist option has to be set to true:

```
oneobtest=.true.
```

Then go to the namelist section SINGLEOB_TEST to set up the single observation location and variable to be tested, please see Section 4.2 for an example and details on the single observation test.



4

GSI Diagnostics and Tuning

The guidance in this chapter will help users understand how and where to check output from GSI to determine whether a run was successful. Properly checking the GSI output will also provide useful information to diagnose potential errors in the system. This chapter starts with an introduction to the content and structure of the GSI standard output file: (**stdout**). It continues with the use of a single observation to check the features of the GSI analysis. Then, observation usage control, analysis domain partitioning, fit files, and the optimization process will all be presented from information within the GSI output files (including **stdout**).

This chapter follows a case example for 2014061700. This case uses a WRF-ARW NetCDF file as the background and analyzes several observations typical for operations, including most conventional observation data, several radiance data sets (AMSU-A, HIRS4, and MHS), and GPSRO data. The case was run on a Linux cluster supercomputer, using four processors. Users can execute this test to reproduce the following results by visiting:

<http://www.dtcenter.org/com-GSI/users/tutorial/index.php>

4.1 Understanding Standard Output (*stdout*)

In Section 3.3, we listed the files present in the GSI run directory following a successful GSI analysis and briefly introduced the contents of several important files. Of these, **stdout** is the most useful because critical information about the GSI analysis can be obtained from the file. From **stdout**, users can check if the GSI has successfully completed, if optimal iterations look correct, and if the background and analysis fields are reasonable. Understanding the

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content of this file can also be very helpful for users to find where and why the GSI failed if it crashes.

The structure of **stdout** follows the typical steps of a meteorological data analysis system:

1. Read in all data and prepare analysis:
 - Read in configuration (namelist)
 - Read in the background
 - Read in observations
 - Partition domain and data for parallel analysis
 - Read in constant fields (fixed files)
2. Calculate observation innovations
3. Optimal iteration (analysis)
4. Save analysis results

In this section, the detailed structure and content of **stdout** are explained using the online example case 5: Hybrid 3DEnVar with all observations. To keep the output concise and make it more readable, most repeated content was deleted (shown with a dotted line). For the same reason, the precision of some numbers has been reduced to avoid line breaks in **stdout**.

The following indicates the start of the GSI analysis. It shows the date and time that GSI started running:

```
* . . . . .
PROGRAM GSI_ANL HAS BEGUN. COMPILED 1999232.55   ORG: NP23
STARTING DATE-TIME  NOV 06,2018  11:13:35.994 310 TUE  2458429
```

The following shows the content of **anavinfo**, a list of state and control variables:

```
gsi_metguess_mod*init_: 2D-MET STATE VARIABLES:
ps
z
gsi_metguess_mod*init_: 3D-MET STATE VARIABLES:
u
v
div
vor
tv
q
oz
cw
gsi_metguess_mod*init_: ALL MET STATE VARIABLES:
u
v
div
vor
tv
q
oz
cw
ps
z
state_vectors*init_anasv: 2D-STATE VARIABLES ps          sst
state_vectors*init_anasv: 3D-STATE VARIABLES u            v
                                     tv            tseni
                                     q            oz
```

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```

                                cw                prse
state_vectors*init_anasv: ALL STATE VARIABLES u          v
                                tv                tsen
                                q                 ozi
                                cw                prsei
                                ps                sst

control_vectors*init_anacv: 2D-CONTROL VARIABLES ARE ps      sst
control_vectors*init_anacv: 3D-CONTROL VARIABLES ARE sf      vp
                                t                 q
                                oz                cw
control_vectors*init_anacv: MOTLEY CONTROL VARIABLES stl     sti
control_vectors*init_anacv: ALL CONTROL VARIABLES sf         vp
                                ps                t
                                q                 oz
                                sst              cw
                                stl              sti

```

Next is the content of all namelist variables used in this analysis. The 1st part shows 4DVAR setup information. Please note that while this version of the GSI includes a 4DVAR option, it remains untested. The general setup for the GSI analysis (3DVAR, 3D/4D EnVar) is located in the &SETUP section of the GSI namelist. Please check Appendix C for definitions and default values of each namelist variable.

```

GSI_4DVAR: nobs_bins =          3
SETUP_4DVAR: nobs_bins =    3, ntlevs_ens =    1
SETUP_4DVAR: allocate array containing time levels for ensemble
SETUP_4DVAR: timelevel =    1 , ens_fhrlevs =    6
SETUP_4DVAR: l4dvar= F
SETUP_4DVAR: l4densvar= F
SETUP_4DVAR: winlen=  2.0000000000000000
SETUP_4DVAR: winoff=  1.0000000000000000
SETUP_4DVAR: hr_obsbin=  1.0000000000000000
SETUP_4DVAR: nobs_bins=    1
SETUP_4DVAR: ntlevs_ens=    1
SETUP_4DVAR: nsubwin,nhr_subwin=    1    2
SETUP_4DVAR: lsqrtb= F
SETUP_4DVAR: lbicg= F
SETUP_4DVAR: lcongrad= F
SETUP_4DVAR: lbfgsmin= F
SETUP_4DVAR: ltlint= F
SETUP_4DVAR: ladtest,ladtest_obs,lgrtest= F F F
SETUP_4DVAR: iwrtinc=    -1
SETUP_4DVAR: lanczosave= F
SETUP_4DVAR: ltcost= F
SETUP_4DVAR: jsiga=    -1
SETUP_4DVAR: nwrvecs=    -1
SETUP_4DVAR: iorthomax=    0
SETUP_4DVAR: liauon= F
SETUP_4DVAR: ljc4tlevs= F
SETUP_4DVAR: ibin_anl=    1
in gsimod: use_gfs_stratosphere,nems_nmmb_regional,wrf_nmm_regional= F F F
GSIMOD: ***WARNING*** set l_cloud_analysis=false
INIT_OBSMOD_VARS: reset time window for one or more OBS_INPUT entries to  1.5000000000000000
INIT_OBSMOD_VARS: ndat_times,ndat_types,ndat=    1    82    82
INIT_OBSMOD_VARS: nhr_assimilation=    2
GSIMOD: ***WARNING*** reset oberrflg= T
calling gsisub with following input parameters:

```

```

&SETUP
GENCODE=  78.0000000000000000    ,
FACTQMIN=  0.0000000000000000    ,
FACTQMAX=  0.0000000000000000    ,
CLIP_SUPERSATURATION=F,
FACTV=  0.0000000000000000    ,
FACTL=  0.0000000000000000    ,

```

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```
FACTP= 0.0000000000000000 ,
FACTG= 0.0000000000000000 ,
FACTW10M= 0.0000000000000000 ,
FACTHOWV= 0.0000000000000000 ,
FACTCLDCH= 0.0000000000000000 ,
R_OPTION=F,
DELTIM= 1200.0000000000000 ,
DTPHYS= 3600.0000000000000 ,
BIASCOR= 0.97999999999999998 , 0.10000000000000001 ,
BCOPTION= 0,
DIURNALBC= 0.0000000000000000 ,
NITER= 0, 2*50 , 48*0 ,
NITER_NO_QC= 51*1000000 ,
MITER= 2,
QOPTION= 2,
CWOPTION= 0,
NHR_ASSIMILATION= 2,
MIN_OFFSET= 60,
PSEUDO_Q2=F,
IOUT_ITER= 220,
NPREDP= 6,
...
/
&GRIDOPTS
...
&BKGERR
...
&ANBKGERR
...
&JCOPTS
...
&STRONGOPTS
...
&OBSQC
...
&SUPEROB_RADAR
...
&LAG_DATA
...
&HYBRID_ENSEMBLE
...
&RAPIDREFRESH_CLDSURF
...
&CHEM
...
&NST
...
```

This version of GSI attempts to read multiple time level backgrounds for option FGAT (First Guess at Appropriate Time), however we only have provided one time level in this test case. Therefore, there is an error message while reading background that does not exist:

```
CONVERT_NETCDF_MASS: problem with flnm1 = wrf_inou1, Status = -1021
```

We can ignore errors for missing files *wrf_inou1*, ..., *wrf_inou9*, because we are only running hybrid 3D EnVAR with one background.

Next, the background fields for the analysis are read in, and the maximum, minimum, and median values of the fields at each vertical level are displayed. Here, only part of the variables ZNU and T are shown, with all other variables read by the GSI listed solely as the variable name in the NetCDF file (*rmse_var = T*). Maximum and minimum values are useful for a quick verification that the background fields have been read successfully. From this section, we also know the time (*iy,m,d,h,m,s*) and dimension (*nlon,lat,sig_regional*) of the background field.

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```

convert wrf_inout to sigf02
iy,m,d,h,m,s=      2018          8          12          12          0          0
dh1 =              2
rmse_var = SMOIS
ndim1 =            3
ordering = XYZ
staggering = N/A
start_index =      1          1          1          0
end_index =       190        114          4          0
WrfType =          104
ierr =             0
rmse_var = T ndim1 = 3 dh1 = 2
WrfType =          104 ierr = 0
ordering = XYZ staggering = N/A
start_index =      1          1          1          0 end_index = 190 114 32
nlon,lat,sig_regional= 190 114 32
rmse_var = P_TOP ndim1= 0
WrfType =          104 WRF_REAL= 104 ierr = 0
ordering = 0 staggering = N/A
start_index =      1          1          1          0 end_index = 190 114 32
p_top= 5000.00000
rmse_var = ZNU ndim1= 1
WrfType =          104 WRF_REAL= 104 ierr = 0
ordering = Z staggering = N/A
start_index =      1          1          1          0 end_index = 32 114 32
k,znu(k)=          1 0.996907353
k,znu(k)=          2 0.989882708
k,znu(k)=          3 0.980982423
...
k,znu(k)=          31 1.61046982E-02
k,znu(k)=          32 5.04574692E-03
rmse_var = ZNW ndim1= 1
...
rmse_var = RDX ndim1= 0
...
rmse_var = RDY ndim1= 0
...
rmse_var = MAPFAC_M ndim1= 2
...
rmse_var = XLAT ndim1= 2
...
rmse_var = XLONG ndim1= 2
...
rmse_var = MUB ndim1= 2
...
rmse_var = MU ndim1= 2
...
rmse_var = PHB ndim1= 3
...
rmse_var = T ndim1= 3
WrfType =          104 WRF_REAL= 104 ierr = 0
ordering = XYZ staggering = N/A
start_index =      1          1          1          0 end_index = 190 114 32
k,max,min,mid T=    1 315.226959 274.242065 295.652863
k,max,min,mid T=    2 321.905151 276.876068 299.062286
k,max,min,mid T=    3 321.910370 278.549225 301.189453
k,max,min,mid T=    4 322.678375 279.197327 302.514221
k,max,min,mid T=    5 323.098846 280.000549 303.465729
...
...
k,max,min,mid T=    30 465.381653 415.209686 435.941864
k,max,min,mid T=    31 488.598053 446.339020 463.024048
k,max,min,mid T=    32 513.816467 476.332092 490.115540
rmse_var = QVAPOR ndim1= 3
...
rmse_var = U ndim1= 3
...
rmse_var = V ndim1= 3
...
rmse_var = XLAND ndim1= 2
...
rmse_var = SEAICE ndim1= 2

```


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```
...
  rmse_var = SST ndim1=      2
...
  rmse_var = IVGTYP ndim1=   2
...
  rmse_var = ISLTYP ndim1=   2
...
  rmse_var = VEGFRA ndim1=   2
...
  rmse_var = SNOW ndim1=     2
...
  rmse_var = U10 ndim1=      2
...
  rmse_var = V10 ndim1=      2
...
  rmse_var = SMOIS ndim1=    3
...
  rmse_var = TSLB ndim1=     3
...
  rmse_var = TSK ndim1=      2
```

Again, some error information on missing background files shows up. Ignore if you are not doing FGAT.

```
CONVERT_NETCDF_MASS: problem with flnm1 = wrf_inou4, Status = -1021
```

Following this is information on the byte order of the binary background files. Since we used a NetCDF file, there is no need to be concerned with byte order. When using a binary format background, byte-order can be a problem. Beginning with the release version v3.2, GSI can automatically check the background byte-order and read it in the right order:

```
in convert_regional_guess, for wrf arw binary input, byte_swap= F
```

Information on setting the grid related variables, and the beginning and ending indices for thread one:

```
INIT_GRID_VARS: number of threads          1
INIT_GRID_VARS: for thread          1  jtstart,jtstop =          1          97
```

Information on the initial pointer location for each variable in the Jacobian for the use of satellite radiance data:

```
Vars in Rad-Jacobian (dims)
-----
sst          0
u            1
v            2
tv           3
q           35
oz           67
```

The following shows some output for debug. Those lines start with core numbers, such as [000]. We will not discuss those debug information but there are many similar lines in stdout file:

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```
[000]gsisub(): : starting ...
[000]gsisub(): init_pass = T
[003]gsisub(): iadate(5)= 0
```

Starting subroutine *gsisub* (major GSI control subroutine) and displaying the analysis and background file time (they should be the same):

```
in gesinfo: iadatemn with minutes      2018      8      12      12      0
in gesinfo: iadatemn with minutes      2018      8      12      12      0
READ_wrf_mass_FILES: analysis date,minutes      2018      8      12      12      0      21360240
READ_wrf_mass_FILES: sigma guess file, nming2      0.0000000000000000      2018      8      12
READ_wrf_mass_FILES: sigma fcst files used in analysis :      2      1.0000000000000000      1
READ_wrf_mass_FILES: surface fcst files used in analysis:      2      1.0000000000000000      1
RADAR_BUFREAD_READ_ALL: radial wind infile for superobbing =
***WARNING*** NOT USING ANY RADIAL WIND OBSERVATIONS!!!
GESINFO: Guess date is      12      8      12      2018      0.0000000000000000
GESINFO: Analysis date is      2018      8      12      12      0      2018081212      1.00
```

Read in radar location information and generate superobs for radar level-II radial velocity. This case didn't have radar level-II velocity data linked, therefore there is warning about missing observations, but this will not impact the rest of the GSI analysis.

```
***WARNING*** NOT USING ANY RADIAL WIND OBSERVATIONS!!!
```

Read in information from fix file *satbias_in* and other info files (see table 3.2). Here shows warning information on channels in *satbias_in* but are not listed in *satinfo*.

```
***WARNING file scaninfo not found, use default
***WARNING instrument/channel cris_npp      27 found in satbias_in file but not found in satinfo
***WARNING instrument/channel cris_npp      28 found in satbias_in file but not found in satinfo
***WARNING instrument/channel cris_npp      31 found in satbias_in file but not found in satinfo
...
***WARNING instrument/channel cris_npp      1298 found in satbias_in file but not found in satinfo
***WARNING instrument/channel cris_npp      1301 found in satbias_in file but not found in satinfo
INIT_PREDX: enter routine
...
radiance_obstype_init: total_rad_type=      14 types are: hirs      goes_img      airs      amsua      amsub      mhs
                      ssmi      amstre      ssmis      sndr      iasi      seviri      atms      cris
radiance_obstype_init: type=goes_img      nch=      4      lcloud_fwd= F      lallsky= F      laerosol_fwd= F      laerosol= F
radiance_obstype_init: type=airs      nch=      281      lcloud_fwd= F      lallsky= F      laerosol_fwd= F      laerosol= F
...
radiance_obstype_init: type=seviri      nch=      8      lcloud_fwd= F      lallsky= F      laerosol_fwd= F      laerosol= F
radiance_obstype_init: type=atms      nch=      22      lcloud_fwd= F      lallsky= F      laerosol_fwd= F      laerosol= F
```

Read in and show the content of the conventional observation information file (*convinfo*; see Section 4.3 for details). Here is the part of the **stdout** file showing information from *convinfo*:

```
READ_CONVINFO: tcp      112      0      1      3.00000      0      0      0      75.0000      5.00000      1.00000      75.0000      0.00000      0      0.00000
READ_CONVINFO: ps      120      0      1      3.00000      0      0      0      4.00000      3.00000      1.00000      4.00000      0.300000E-03      0      0.00000
...
READ_CONVINFO: t      120      0      1      3.00000      0      0      0      8.00000      5.60000      1.30000      8.00000      0.100000E-05      0      0.00000
READ_CONVINFO: t      126      0      -1      3.00000      0      0      0      8.00000      5.60000      1.30000      8.00000      0.100000E-02      0      0.00000
...
READ_CONVINFO: gps      4      0      1      3.00000      0      0      0      10.0000      10.0000      1.00000      10.0000      0.00000      0      0.00000
READ_CONVINFO: gps      41      0      -1      3.00000      0      0      0      10.0000      10.0000      1.00000      10.0000      0.00000      0      0.00000
...
READ_CONVINFO: gps      44      0      -1      3.00000      0      0      0      10.0000      10.0000      1.00000      10.0000      0.00000      0      0.00000
```

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Starting subroutine *glbsoi* with information on reading in background fields from the intermediate binary file *sigf02* and partitioning the whole 2D field into subdomains for parallel analysis:

```
glbsoi: starting ...
before compute nlat_ens,nlon_ens, nlat,nlon,nlat_ens,nlon_ens,r_e,eps=      114      190      0      0
dual_res,nlat,nlon,nlat_ens,nlon_ens,r_e,eps= F      114      190      114      190  1.0000000000000000
gsi_metguess_mod*create_: alloc() for met-guess done
guess_grids*create_chemges_grids: trouble getting number of chem/gases
at 0 in read_wrf_mass_guess
read_wrf_mass_guess: lm = 32
read_wrf_mass_guess: num_mass_fields= 142
read_wrf_mass_guess: nfldsig = 1
read_wrf_mass_guess: num_all_fields= 142
read_wrf_mass_guess: npe = 4
read_wrf_mass_guess: num_loc_groups= 35
read_wrf_mass_guess, num_all_pad = 144
read_wrf_mass_guess, num_loc_groups= 36
READ_WRF_MASS_GUESS: open lendian_in= 15 to file=sigf02
in read_wrf_mass_guess, min,max(soil_moi)= 5.5290825664997101E-002 1.0000000000000000
in read_wrf_mass_guess, min,max(soil_temp)= 273.16000366210938 307.12820434570312
in read_wrf_mass_guess, num_doubtful_sfct_all = 0
in read_wrf_mass_guess, num_doubtful_sfct_all = 0
```

Show observation observer as successfully initialized and inquire about the control vectors (space for analysis variables).

```
observer_init: successfully initialized
control_vectors: length= 2064450
control_vectors: currently allocated= 0
control_vectors: maximum allocated= 0
control_vectors: number of allocates= 0
control_vectors: number of deallocates= 0
control_vectors: Estimated max memory used= 0.0 Mb
```

Show the source of observation error used in the analysis (details see Section 4.7.1):

```
CONVERR: using observation errors from user provided table
```

The following information is related to the observation ingest processes, which is distributed over all the processors with each processor reading in at least one observation type. To speed up the reading process, some of the large datasets will use more than one (ntasks) processor for the ingest process.

Before reading in data from BUFR files, GSI checks the file status to insure the observation time matches the analysis time and whether the namelist option *offtime_data* is set (can be used to turn off the time consistency check between observation and analysis time). This step also checks for consistency between the satellite radiance data types in the BUFR files and the usage setups in the *satinfo* files. The following shows **stdout** information from this step:

```
read_obs_check: bufr file date is 2018081212 prepbufr t
read_obs_check: bufr file date is 2018081212 prepbufr ps
read_obs_check: bufr file uv not available satwndbufr
read_obs_check: bufr file rw not available l2rwbufr
```

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```

read_obs_check: bufr file pcp_tmi   trmm       not available tmirrbufr
read_obs_check: bufr file hirs3     n17       not available hirs3bufr
read_obs_check: bufr file hirs3     n17       not available hirs3bufrears
read_obs_check: bufr file goes_img  g11       not available gimgrbufr
read_obs_check: bufr file date is   2018081212 prepbufr q
read_obs_check: bufr file date is   2018081212 amsuabufr amsua   n18
read_obs_check: bufr file amsua     n18       not available amsuabufrears
read_obs_check: bufr file amsua     n18       not available amsuabufr_db

...
...

read_obs_check: bufr file mhs       n19       not available mhsbufr_db
data type ssmi_f13                 not used in info file -- do not read file ssmitbufr
data type ssmi_f14                 not used in info file -- do not read file ssmitbufr
read_obs_check: bufr file amsre_low aqua   not available amsrebufr
read_obs_check: bufr file ssmis_uas f16    not available ssmisbufr
read_obs_check: bufr file sndrd2   g12     not available gsnd1bufr
read_obs_check: bufr file sndrd2   g11     not available gsnd1bufr
read_obs_check: bufr file sndrd2   g13     not available gsnd1bufr
read_obs_check: bufr file sndrd2   g15     not available gsnd1bufr
read_obs_check: bufr file gome      metop-a not available gomebufr
read_obs_check: bufr file seviri    m08     not available seviribufr
read_obs_check: bufr file gome      metop-b not available gomebufr
data type cris_npp                 not used in info file -- do not read file crisbufr
data type mta_cld                  not used in info file -- do not read file prepbufr

```

The list of observation types that were read in and processors used to read them:

```

number of extra processors          1
READ_OBS: read 33 mhs               mhs_n18      using ntasks= 2 0 2123 999999 999999
READ_OBS: read 34 mhs               mhs_n19      using ntasks= 2 2 159 999999 999999
READ_OBS: read 35 mhs               mhs_metop-a  using ntasks= 2 0 590 999999 999999
READ_OBS: read 36 mhs               mhs_metop-b  using ntasks= 2 2 3 999999 999999
READ_OBS: read 1 ps                 ps           using ntasks= 1 0 1 999999 999999
READ_OBS: read 2 t                  t            using ntasks= 1 1 1 999999 999999
READ_OBS: read 3 q                  q            using ntasks= 1 2 1 999999 999999
READ_OBS: read 4 pw                 pw           using ntasks= 1 0 4218 999999 999999
READ_OBS: read 6 uv                 uv           using ntasks= 1 1 1 999999 999999
READ_OBS: read 10 sst               sst         using ntasks= 1 2 611 999999 999999
READ_OBS: read 11 gps_ref           gps          using ntasks= 1 0 1 999999 999999
READ_OBS: read 19 hirs4             hirs4_metop-a using ntasks= 2 1 292 999999 999999
READ_OBS: read 21 hirs4             hirs4_n19    using ntasks= 1 2 80 999999 999999
READ_OBS: read 22 hirs4             hirs4_metop-b using ntasks= 1 0 3 999999 999999
READ_OBS: read 26 amsua             amsua_n15    using ntasks= 1 1 72 999999 999999
READ_OBS: read 27 amsua             amsua_n18    using ntasks= 1 2 509 999999 999999
READ_OBS: read 28 amsua             amsua_n19    using ntasks= 1 0 31 999999 999999
READ_OBS: read 29 amsua             amsua_metop-a using ntasks= 1 1 159 999999 999999
READ_OBS: read 30 amsua             amsua_metop-b using ntasks= 1 2 3 999999 999999

```

Display basic statistics for full horizontal surface fields (If radiance BUFR files are not linked, this section will not be in the **stdout** file):

```

=====
Status  Var      Mean              Min              Max
sfcges2 FC10  1.000000000000E+00  1.000000000000E+00  1.000000000000E+00
sfcges2 SNOW  7.240454056477E-02  0.000000000000E+00  1.990934906006E+02
sfcges2 VFRC  2.058203305344E-01  0.000000000000E+00  8.997633457184E-01
sfcges2 SRGH  4.999980274880E-02  5.000000074506E-02  5.000000074506E-02
sfcges2 STMP  2.799090258541E+02  2.724390563965E+02  3.071282043457E+02
sfcges2 SMST  7.058461197195E-01  2.784143574536E-02  1.000000000000E+00
sfcges2 SST   2.937425438596E+02  2.656092224121E+02  3.091810302734E+02
sfcges2 VTYP  1.335590951062E+01  1.000000000000E+00  1.900000000000E+01
sfcges2 ISLI  3.752077562327E-01  0.000000000000E+00  1.000000000000E+00
sfcges2 STYP  1.072373037858E+01  1.000000000000E+00  1.600000000000E+01
=====

```

4. GSI Diagnostics and Tuning

Loop over all data files to read in observations, also read in rejection list for surface observations, if new VAD wind is used, and show GPS observations outside the time window:

```

READ_BUFERTOVS      : file=mhsbufr      type=mhs      sis=mhs_n18      nread= 99700 ithin= 2 rmesh= 60.000000 isfcalc= 0
                      nkeep= 10880 ntask= 2
READ_BUFERTOVS      : file=mhsbufr      type=mhs      sis=mhs_n19      nread= 164720 ithin= 2 rmesh= 60.000000 isfcalc= 0
nkeep= 17575 ntask= 2
READ_BUFERTOVS      : file=mhsbufr      type=mhs      sis=mhs_metop-a   nread= 1190 ithin= 2 rmesh= 60.000000 isfcalc= 0
nkeep= 225 ntask= 2
READ_BUFERTOVS      : file=mhsbufr      type=mhs      sis=mhs_metop-b   nread= 39330 ithin= 2 rmesh= 60.000000 isfcalc= 0
nkeep= 4410 ntask= 2
new vad flag:: T
new vad flag:: T
new vad flag:: T
READ_PREPBUFR: messages/reports = 4173 / 501125 nthread = 1
WARNING: GLERL program code not in this file.
mesonetuselist: listexist,nprov= F 0
w_rejectlist: wlistexist,nwrjs= F 0
t_rejectlist: tlistexist,ntrjs= F 0

...

READ_PREPBUFR      : file=prepbufr      type=q      sis=q      nread= 14530 ithin= 0 rmesh= 120.000000 isfcalc= 0
nkeep= 13316 ntask= 1
READ_PREPBUFR: clobf( 13 )
READ_PREPBUFR      : file=prepbufr      type=pw     sis=pw     nread= 817 ithin= 0 rmesh= 120.000000 isfcalc= 0
nkeep= 817 ntask= 1
READ_GPS: time outside window -3.0000000000000000 skip this report

...

READ_BUFERTOVS      : file=amsuabufr     type=amsua   sis=amsua_n18   nread= 33465 ithin= 2 rmesh= 60.000000 isfcalc= 0
nkeep= 27762 ntask= 1
READ_BUFERTOVS      : file=amsuabufr     type=amsua   sis=amsua_metop-b nread= 13335 ithin= 2 rmesh= 60.000000 isfcalc= 0
nkeep= 10112 ntask= 1
dval_use = T

```

Using the above output information, many details on the observations can be obtained. For example, the last line indicates that subroutine *READ_BUFERTOVS* was called to read in METOP-B AMSU-A (*sis=amsua_metop-b*) data from the BUFR file *amsuabufr* (*file=amsuabufr*). Furthermore, there are 13335 observations in this file (*nread=13335*) and 10112 in the analysis domain and within the time window (*nkeep=10112*). The data was thinned on a 60 km coarse grid (*rmesh=60.000000*).

The next step partitions observations into subdomains. The observation distribution is summarized below by listing the number of observations for each variable per subdomain (see Section 4.4 for more information):

OBS_PARA: ps		809	1447	2099	2982
OBS_PARA: t		2380	2792	4398	5207
OBS_PARA: q		2324	2690	4127	4653
OBS_PARA: pw		165	245	147	277
OBS_PARA: uv		2784	2683	4657	4856
OBS_PARA: sst		5	2	44	21
OBS_PARA: gps_ref		0	526	0	1112
OBS_PARA: hirs4	metop-a	0	0	0	33
OBS_PARA: hirs4	n19	1080	1921	0	299
OBS_PARA: hirs4	metop-b	0	0	58	180
OBS_PARA: amsua	n15	0	0	882	889
OBS_PARA: amsua	n18	0	0	898	971
OBS_PARA: amsua	n19	1026	1797	0	263
OBS_PARA: amsua	metop-a	0	0	0	25
OBS_PARA: amsua	metop-b	0	0	148	580
OBS_PARA: mhs	n18	0	0	1047	1148
OBS_PARA: mhs	n19	1198	2049	1	338
OBS_PARA: mhs	metop-a	0	0	0	45
OBS_PARA: mhs	metop-b	0	0	213	678

Information on ingesting background error statistics:

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```
m_berror_stats_reg::berror_read_bal_reg(PREBAL_REG): get balance variables "berror_stats". mype,nsigstat,nlatstat =
    0          60          93
m_berror_stats_reg::berror_read_wgt_reg(PREWGT_REG): read error amplitudes "berror_stats". mype,nsigstat,nlatstat =
    0          60          93
Assigned default statistics to variable oz
Assigned default statistics to variable cw
```

For hybrid analysis, the following information is about the GFS ensemble forecast, including time of the ensemble forecast, vertical coordinate, resolution of the forecast and location of the files:

```
nemsio: fhour, idate= 6.0000000000000000 2018 8 12 6
iadate(y,m,d,hr,min)= 2018 8 12 12 0
nemsio: jcap, levs= 62 64
nemsio: latb, lonb= 94 192
nemsio: idvc, nvcoord= 2 2
nemsio: idsl= 1
in get_gefs_for_regional, iadate_gefs= 2018 8 12 12 0
in get_gefs_for_regional, iadate = 2018 8 12 12 0
ak5,bk5,ck5= 0.0000000000000000 1.0000000000000000 0.0000000000000000
ak5,bk5,ck5= 0.0000000000000000 0.99467116594314575 0.0000000000000000
ak5,bk5,ck5= 5.7499998807907101E-004 0.98862659931182861 0.0000000000000000
ak5,bk5,ck5= 5.7410001754760742E-003 0.98174226284027100 0.0000000000000000
...
ak5,bk5,ck5= 0.31826599121093752 0.0000000000000000 0.0000000000000000
ak5,bk5,ck5= 0.22195799255371093 0.0000000000000000 0.0000000000000000
ak5,bk5,ck5= 0.13778999328613281 0.0000000000000000 0.0000000000000000
ak5,bk5,ck5= 6.4247001647949220E-002 0.0000000000000000 0.0000000000000000
ak5,bk5,ck5= 0.0000000000000000 0.0000000000000000 0.0000000000000000
GENERAL_READ_GFSATM_NEMS: read lonb,latb,levs= 192 94 64, scatter nlon,nlat= 192 94, hour= 6.0,
idate= 6 8 12 2018 /glade/p/ral/jntp/DATask/case_data/com_2018081212/gfsens/gdas.t06z.atmf006s.mem001.nemsio
GENERAL_READ_GFSATM_NEMS: read lonb,latb,levs= 192 94 64, scatter nlon,nlat= 192 94, hour= 6.0,
idate= 6 8 12 2018 /glade/p/ral/jntp/DATask/case_data/com_2018081212/gfsens/gdas.t06z.atmf006s.mem002.nemsio
```

Show the hybrid setups in each vertical level:

```
HYBENS_LOCALIZATION_SETUP: s_ens_hv,s_ens_vv,beta_s,beta_e
110.0 3.0 0.5000 0.5000
110.0 3.0 0.5000 0.5000
110.0 3.0 0.5000 0.5000

110.0 3.0 0.5000 0.5000
110.0 3.0 0.5000 0.5000
110.0 3.0 0.5000 0.5000
110.0 3.0 0.5000 0.5000
110.0 3.0 0.5000 0.5000
```

From this point forward in the **stdout** file, the output shows many repeated entries. This is because the information is written from inside the outer loop. Typically the outer loop is run twice.

For each outer loop, the work begins with the calculation of the observation innovation. This calculation is done by the subroutine **setuprhalls**, which sets up the right hand side (rhs) of the analysis equation. This information is contained within the **stdout** file, which is shown in the following sections:

Start the first outer analysis loop:

```
GLBSOI: jiter,jiterstart,jiterlast,jiterend= 1 1 2 1
```

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Calculate observation innovation for each data type in the first outer loop. When computing the radiance observation innovation, information on reading in CRTM coefficients follows SETUPALL information. In the **stdout** file, only information related to available radiance data are printed. The complete innovation information can be found in the diagnostic files for each observation (for details see Appendix A.2):

```
crtm_interface*init_crtm: crt_m_init() on path "./"

SpCcoeff_ReadFile(Binary)(INFORMATION) : FILE: ./hirs4_n19.SpcCoeff.bin;
SpCcoeff RELEASE.VERSION: 8.04
  N_CHANNELS=19
...
FitCoeff RELEASE.VERSION : 1.6; DIMENSIONS= 8
FitCoeff_ReadFile(INFORMATION) : FILE: ./FASTEM6.MWwater.EmisCoeff.bin;
FitCoeff RELEASE.VERSION : 1.6; DIMENSIONS= 3, 6, 2
MWwaterCoeff_ReadFile(INFORMATION) : FILE: ./FASTEM6.MWwater.EmisCoeff.bin;
MWwaterCoeff RELEASE.VERSION: 1.6
SETUPPRAD: write header record for mhs_metop-a           12           30           8           0           0
           23           4           40000 to file pe0003.mhs_metop-a_01 2018081212
```

The inner iteration of the first outer loop is discussed in the example below. In this example, the maximum number of iterations is 50.

Print cost function values for each inner iteration (see section 4.6 for more details):

```
GLBSOI: START pcgsoi jiter= 1
pcgsoi: gnorm(1:3),b= 2.729446579175403500E+15 2.621641902596249565E+03 2.621641902596249565E+03 0.0000000000000000E+00
Begin J table inner/outer loop 0 1
  J term J
surface pressure 4.7741393535600264E+03
temperature 7.8419025904486416E+03
wind 8.2939160322155458E+03
moisture 4.2982963156969481E+03
sst 4.4435897008983467E+01
gps 1.4249707290215210E+03
radiance 1.0312906716180896E+04
-----
J Global 3.6990567634132560E+04
End Jo table inner/outer loop 0 1
Initial cost function = 3.699056763413256704E+04
Initial gradient norm = 5.224410568834922463E+07
cost,grad,step,b,step? = 1 0 3.699056763413256704E+04 5.224410568834922463E+07 8.253769138443810549E-01 0.0000000000000000E+00 good
pcgsoi: gnorm(1:3),b= 1.111843080460129375E+15 3.081400755207042721E+02 3.081400755207050679E+02 1.175370576795971844E-01
cost,grad,step,b,step? = 1 1 3.482672493136255798E+04 3.334431106590941921E+07 2.966537900900262148E+00 1.175370576795971844E-01 good
pcgsoi: gnorm(1:3),b= 2.618119688568596000E+15 5.233620050663861321E+02 5.233620050663854499E+02 1.698454847789571343E+00
cost,grad,step,b,step? = 1 2 3.391261571854411886E+04 5.116756480983432382E+07 4.105823018764231058E+00 1.698454847789571343E+00 good
pcgsoi: gnorm(1:3),b= 4.809494828186123000E+15 9.711879327337466066E+02 9.711879327337500172E+02 1.855671453663429338E+00
cost,grad,step,b,step? = 1 3 3.17637839509595218E+04 6.935052147018162906E+07 2.671127324144195292E+00 1.855671453663429338E+00 good
pcgsoi: gnorm(1:3),b= 4.315369758675125500E+15 4.836127760071037756E+02 4.836127760071009334E+02 4.979600340026935079E-01
...
pcgsoi: gnorm(1:3),b= 2.588270087151630554E+11 8.107253043776205814E-01 8.107253043774353962E-01 1.278420303738640351E+00
cost,grad,step,b,step? = 1 49 1.930005078536898873E+04 5.087504385405117646E+05 4.239602407785440619E+00 1.278420303738640351E+00 good
pcgsoi: gnorm(1:3),b= 2.826303730012705078E+12 1.740492590323112099E+00 1.740492590323460043E+00 2.146833928737002850E+00
cost,grad,step,b,step? = 1 50 1.929661363241649815E+04 1.681161422949237516E+06 1.545575515630557950E+00 2.146833928737002850E+00 good
update_guess: successfully complete
```

At the end of the 1st outer loop, print some diagnostics about the analysis increments as well as information on the guess fields after adding the analysis increments to the background:

```
=====
Status Var Mean Min Max
analysis U 1.362868221082E+00 -2.561965601145E+01 7.336788270602E+01
analysis V 3.965723797005E-02 -3.324405649283E+01 3.951375834823E+01
analysis TV 2.588579356507E+02 1.905883571635E+02 3.102596713291E+02
analysis Q 4.581851101606E-03 1.000000000000E-07 2.203928944498E-02
analysis TSEN 2.580511700079E+02 1.905879091828E+02 3.074921960954E+02
analysis OZ 1.000000000002E-15 1.000000000000E-15 1.000000000000E-15
analysis CW 0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
```

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```

analysis DIV      0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
analysis VDR      0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
analysis PRSL     5.385620929685E+01  5.323168380321E+00  1.032286059067E+02
analysis PS       9.890425623867E+01  6.892640994729E+01  1.035367484906E+02
analysis SST      2.937426126383E+02  2.656092224121E+02  3.091810302734E+02
analysis radb     9.425585403234E-02  -1.655751310000E+02  2.821827560000E+02
analysis pcpb     0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
analysis aftb     0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
=====
increment u       7.848960181417E-03  -8.054188393640E+00  6.055781218838E+00
increment v       3.258610899476E-03  -5.799328704874E+00  6.419030188202E+00
increment tv      1.108815673124E-01  -2.029480417256E+00  4.015145579676E+00
increment tsen    1.015912624120E-01  -2.029503916971E+00  4.014574667498E+00
increment q       5.147359869825E-05  -3.832308425511E-03  3.683516094586E-03
increment oz      0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
increment cw      0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
increment prse    -7.866162615204E-03  -2.412199253002E-01  1.123612852372E-01
increment ps      -1.513943279830E-02  -2.412199253002E-01  1.123612852372E-01
increment sst     -3.284418147534E-02  -1.125829038410E+00  1.522857245108E+00

```

Calculate observation innovations for each data type in the second outer loop. When calculating the radiance data innovation, there is no need to read in CRTM coefficients again because they were already read in during the first outer loop:

```

in setupw, bad ikx, ikx,i,nconvtype=      0      1278      215
in setupw, num_bad_ikx ( ikx<1 or ikx>nconvtype ) =      395
      0 setuprad: passive obs      21 hrs4_n19

```

Start the second outer loop.

```

RGLBSOI:  START pcgsoi jiter=      2

```

The output from the inner iterations in the second outer loop is shown below. In this example, the maximum number of iterations is 50.

Print cost function values for each inner iteration (see section 4.6 for more details):

```

GLBSOI:  START pcgsoi jiter=      2
pcgsoi:  gnorm(1:3),b=  1.748934444754522188E+14  2.498908356599028195E+02  2.498908356599027911E+02  0.000000000000000000E+00
Begin J table inner/outer loop      0      2
      J term      J
background      4.0576634502296029E+03
surface pressure 2.0186253679757897E+03
temperature      3.7863130457920138E+03
wind             4.2221633626971798E+03
moisture         1.6848145824733588E+03
sst              4.4414461980361125E+01
gps              4.0673587211891697E+02
radiance        6.0413870599169568E+03
-----
J Global        2.2262117203184182E+04
End Jo table inner/outer loop      0      2
Initial cost function = 2.226211720318417792E+04
Initial gradient norm = 1.322472852180536091E+07
cost,grad,step,b,step? =  2  0  2.226211720318417792E+04  1.322472852180536091E+07  8.743095124801847362E-01  0.000000000000000000E+00  good
pcgsoi:  gnorm(1:3),b=  1.779987726829237500E+14  3.119955441500999171E+01  3.119955441501095450E+01  1.248527355259720639E-01
cost,grad,step,b,step? =  2  1  2.204363526848510082E+04  1.334161806839499250E+07  3.917049140953876130E+00  1.248527355259720639E-01  good
pcgsoi:  gnorm(1:3),b=  9.563111338115188750E+14  6.386441855459911210E+01  6.386441855459896288E+01  2.046965726019221776E+00
cost,grad,step,b,step? =  2  2  2.192142508066563823E+04  3.092428065148030221E+07  3.934891262992486816E+00  2.046965726019221776E+00  good
pcgsoi:  gnorm(1:3),b=  3.674803322863953125E+14  1.058836900113621198E+02  1.058836900113624182E+02  1.657944946619063664E+00
cost,grad,step,b,step? =  2  3  2.167012553807904987E+04  1.916977653198897839E+07  1.535679976735633501E+00  1.657944946619063664E+00  good
...
...
cost,grad,step,b,step? =  2  48  2.074202075612976478E+04  9.824897960038228193E+05  8.345643356966736093E+00  1.195023573152231178E+00  good
pcgsoi:  gnorm(1:3),b=  1.332191889788007812E+11  5.734034451509922858E-02  5.734034451322606724E-02  5.550851189922036966E-01
cost,grad,step,b,step? =  2  49  2.074115865039333221E+04  3.649920396101821680E+05  2.55556896059054228E+00  5.550851189922036966E-01  good
pcgsoi:  gnorm(1:3),b=  4.847178172167221680E+11  1.252577268549982525E-01  1.25257726855055134E-01  2.184460660610548555E+00
cost,grad,step,b,step? =  2  50  2.074101211388048978E+04  6.962167889506272040E+05  3.984326453749022612E+00  2.184460660610548555E+00  good
update_guess: successfully complete

```


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Diagnostics of the analysis results after adding the analysis increment to the guess, as well as diagnostics about the analysis increments:

```

=====
Status  Var          Mean          Min          Max
analysis U      1.360483466628E+00 -2.574925274989E+01  7.336415204906E+01
analysis V      4.012681261518E-02 -3.323994949780E+01  3.955809426534E+01
analysis TV     2.588857375746E+02  1.907340360305E+02  3.102363261552E+02
analysis Q      4.587484199149E-03  1.000000000000E-07  2.211380527114E-02
analysis TSEN   2.580779914425E+02  1.907335880235E+02  3.075871602128E+02
analysis OZ     1.000000000002E-15  1.000000000000E-15  1.000000000000E-15
analysis CW     0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
analysis DIV    0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
analysis VDR   0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
analysis PRSL   5.384833387035E+01  5.322556486019E+00  1.032320091505E+02
analysis PS     9.890409150001E+01  6.893130649187E+01  1.035374921768E+02
analysis SST    2.937426126383E+02  2.656092224121E+02  3.091810302734E+02
analysis radb   9.425576966354E-02  -1.655751310000E+02  2.821827560000E+02
analysis pcpb  0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
analysis aftb  0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
=====
increment u      -2.384754454102E-03  -4.515404663843E-01  1.114576021589E+00
increment v      4.695746451575E-04  -1.189114439170E+00  6.887491209179E-01
increment tv     2.780192388882E-02  -1.101279692098E+00  1.157477342725E+00
increment tsen   2.682062610000E-02  -1.100968731762E+00  1.157476000169E+00
increment q      5.632793095050E-06  -1.900471275333E-03  1.425101132072E-03
increment oz     0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
increment cw     0.000000000000E+00  0.000000000000E+00  0.000000000000E+00
increment prse  -8.559509115317E-05  -2.626616668875E-02  1.007126363345E-02
increment ps    -1.647386653147E-04  -2.626616668875E-02  1.007126363345E-02
increment sst   -4.539864282467E-03  -6.574366341706E-01  7.323275951184E-01
=====

```

Because the outer loop is set to two, the completion of the 2nd outer loop marks the end of the analysis. The next step is to save the analysis results. Again, only a portion of variable "T" is shown and all other variables are listed according to variable name in the NetCDF file (rmse_var = T). The maximum and minimum values are useful information for a quick sanity check of the analysis:

```

at 2 in wrwrfmassa
update sigf02
at 3 in wrwrfmassa
at 6 in wrwrfmassa
iy,m,d,h,m,s=      2018      8      12      12      0      0
nlon,lat,sig_regional=      190      114      32
rmse_var=P_TOP
ordering=0
WrfType,WRF_REAL=      104      104
ndim1=      0
staggering= N/A
start_index=      1      1      1      0
end_index1=      190      114      32      0
p_top= 5000.00000
rmse_var=MUB
ordering=XY
WrfType,WRF_REAL=      104      104
ndim1=      2
staggering= N/A
start_index=      1      1      1      0
end_index1=      190      114      32      0
max,min MUB= 95000.0000 61690.7969
max,min psfc= 103286.703 68803.6797
max,min MU= 3286.70312 -389.023438
rmse_var=MU
ordering=XY
WrfType,WRF_REAL=      104      104

```

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```

ndim1=          2
staggering= N/A
start_index=          1          1          1          0
end_index1=      190          114          32          0
k,max,min,mid T=          1    316.327240    274.314423    295.946381
k,max,min,mid T=          2    322.989807    276.998138    299.261627
...

k,max,min,mid T=          31    489.376556    447.367950    464.863403
k,max,min,mid T=          32    514.591003    476.494690    491.469971
rmse_var=T
...
rmse_var=QVAPOR
...
rmse_var=U
...
rmse_var=V
...
rmse_var=SEAICE
...
rmse_var=SST
...
rmse_var=TSK
...
rmse_var=Q2
...

```

Deallocate the data arrays and finalize the GSI run:

```

prad_bias: number of passive channels=          2564
gsi_metguess_mod*destroy_: dealloc() for met-guess done
observer_final: successfully finalized
glbsoi: complete
[000]gsisub(): : complete.

```

The end of the GSI analysis (reaching this point does not necessarily guarantee a successful analysis), which shows the date and time when GSI finished and some additional resource statistics:

```

          ENDING DATE-TIME    NOV 06,2018  11:15:55.503  310  TUE   2458429
          PROGRAM GSI_ANL HAS ENDED.
* . . . . .
*****RESOURCE STATISTICS*****
The total amount of wall time              = 139.509586
The total amount of time in user mode      = 135.700000
The total amount of time in sys mode      = 3.472000
The maximum resident set size (KB)        = 519420
Number of page faults without I/O activity = 1471061
Number of page faults with I/O activity   = 0
Number of times filesystem performed INPUT = 0
Number of times filesystem performed OUTPUT = 0
Number of Voluntary Context Switches      = 5723
Number of InVoluntary Context Switches    = 3159
*****END OF RESOURCE STATISTICS*****

```

Different GSI applications may write out slightly different **stdout** file information but the major flow and information are the same. A good knowledge of the **stdout** file gives users a clear picture of how GSI runs and the key information provided during a GSI run like data distribution and inner iterations.

4.2 Single Observation Test

A single observation test is a GSI where only one (pseudo) observation is assimilated from a specific time and location within the analysis domain. By examining the analysis increments from a single observation test, one can visualize the important features of the analysis, such as the ratio of background error and observation error variance and the pattern of the background error covariance. Therefore, the single observation test is the first thing that users should run after successfully installing the GSI.

4.2.1 Setup a Single Observation Test

To perform the single observation test with the GSI, the following GSI namelist variables need to be set, which should be done through editing the script *ush/comgsi_run_regional.ksh* and *ush/comgsi_namelist.sh*:

In *ush/comgsi_run_regional.ksh*, turn on the single observation test:

```
if_oneob=Yes      # Yes, or, No -- case sensitive !
```

In *ush/comgsi_namelist.sh*, set up single observation features like:

```
maginnov=1.0,  
magoberr=0.8,  
oneob_type='t',  
oblat=38.,  
oblon=262.,  
obpres=500.,  
obdattim= 2014061700,  
obhourset=0.,
```

Note:

- Please check Appendix C in the User's Guide for the explanation of each parameter. From these parameters, we can see that a useful observation in the analysis should include information like the observation type (*oneob_type*), value (*maginnov*), observation error (*magoberr*), location (*oblat*, *oblong*, *obpres*), and time (*obdattim*, *obhourset*). Users can dump out (use *ncdump*) the global attributes from the NetCDF background file and set *oblat=CEN_LAT*, *oblong=360-CEN_LON* to have the observation at the center of the domain.
- In the analysis, the GSI first generates a *prepbufr* file including only one observation based on the information given in the namelist &SINGLEOB_TEST section. To generate this *prepbufr* file, the GSI needs to read in a *PrepBUFR* table, which is not needed when running a GSI analysis with real observations. The *BUFR* table is in the *fix/* directory and needs to be copied to the run directory. We have put the following lines in the GSI run script for the single observation test:

```
bufrtable=${FIX_ROOT}/prepobs_prep.bufrtable  
cp $bufrtable ./prepobs_prep.bufrtable
```

4.2.2 Examples of Single Observation Tests for GSI

Figure 4.1 is a single observation test that has a temperature observation (`oneob_type='t'`) with a one degree innovation (`maginnov=1.0`) and a 0.8 degree observation error (`magoberr=0.8`). The background error covariance converted from global (GFS) BE was picked to provide for better illustration.

This single observation was located at the center of the domain. The results are shown with figures of the horizontal and vertical cross sections through the point of maximum analysis increment. The Figure 4.1 was generated using NCL scripts, which can be found in the `util/Analysis_Uilities/plots_ncl` directory, introduced in Section A.4.

4.3 Control Data Usage

Observation data used in the GSI analysis can be controlled through three parts of the GSI system:

1. In the GSI run script, by linking observation BUFR files to the working directory
2. In section `&OBS_INPUT` of the GSI namelist (inside `comgsi_namelist.sh`)
3. Through parameters in info files (e.g.: `convinfo`, `satinfo`, etc.)

Each part provides different levels of control for data usage in the GSI, which is introduced below:

1. Link observation BUFR files to the working directory in the GSI run script:

All BUFR/PrepBUFR observation files need to be linked to the working directory with GSI recognizable names before they can be used in a GSI analysis. The run script (`run_gsi_regional.ksh`) makes these links after locating the working directory. Turning these links on or off can control the use of all the data contained in the BUFR files. Table 4.1 provides a list of all default observation file names recognized by GSI and the corresponding examples of the observation BUFR files from NCEP. The following is the first three rows of the table as an example:

Table 4.1: List of all default observation file names recognized by GSI.

GSI Name	Content	Example file names
prepbuf	Conventional observations, including ps, t, q, pw, uv, spd, dw, sst, from observation platforms such as METAR, soundings, etc.	<i>gdas1.t12z.prepbuf</i>
satwndbuf	satellite winds	<i>gdas1.t12z.satwnd.tm00.buf_d</i>
amsuabuf	AMSU-A 1b radiance (brightness temperatures) from satellites NOAA-15, 16, 17,18, 19, and METOP-A/B	<i>gdas1.t12z.1bamua.tm00.buf_d</i>

The left column is the GSI recognized name (bold) and the right column are names of BUFR files from NCEP (italic). In the run script, the following lines are used to link the

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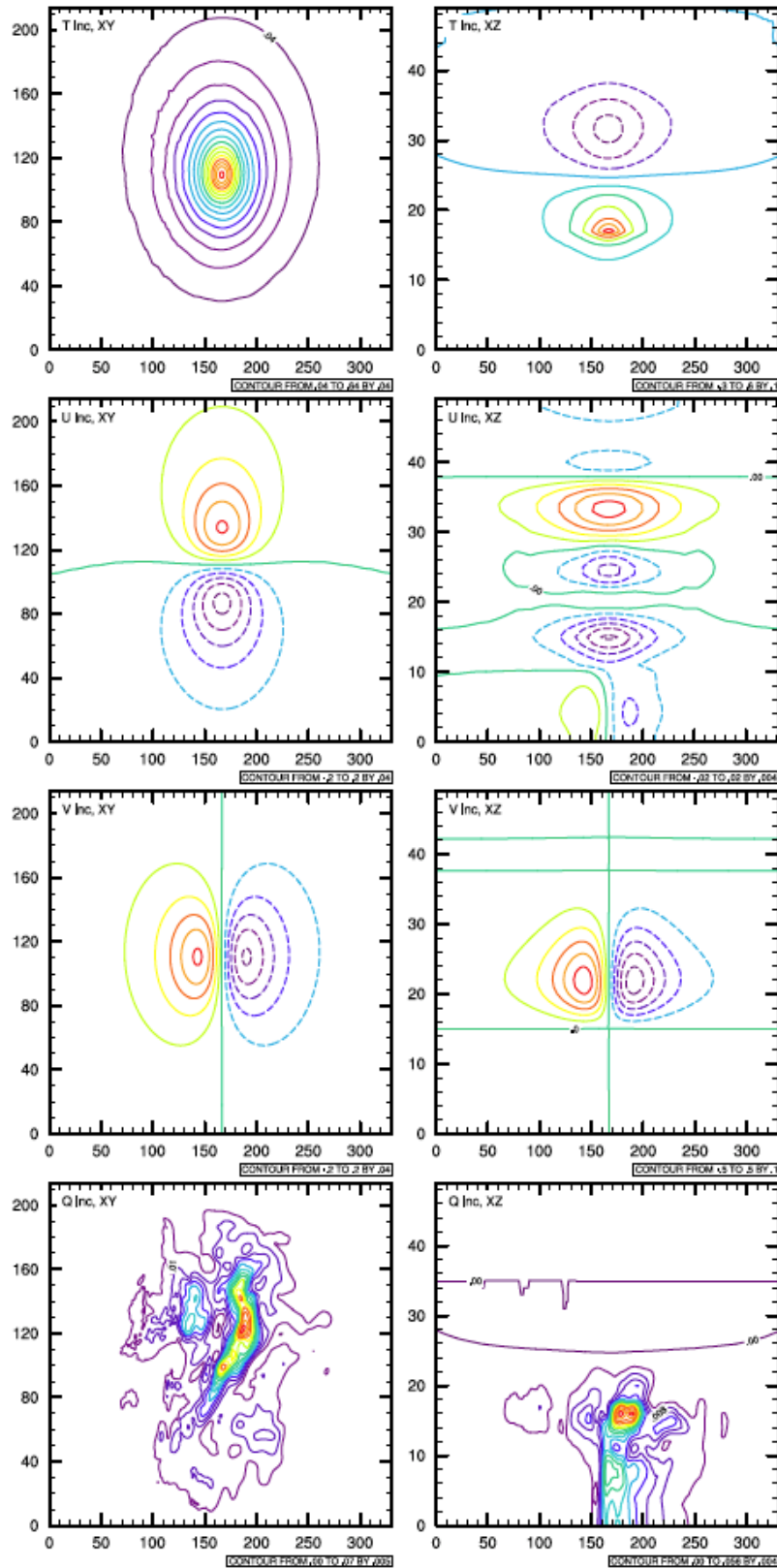


Figure 4.1: Horizontal cross sections (left column) and vertical cross sections (right column) of analysis increment of T, U, V, and Q from a single T observation

BUFR files in the right column to the working directory using the GSI recognized names

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shown in the left column:

```
# Link to the prepbuf data
ln -s ${PREPBUFR} ./prepbuf

# Link to the radiance data
ln -s ${OBS_ROOT}/gdas1.t12z.1bamua.tm00.buf_r_d amsuabuf
```

The GSI recognized default observation filenames are set up in the namelist section &OBS_INPUT, which can be changed based on application needs (see below for details).

2. In the GSI namelist (inside *comgsi_namelist.sh*), section &OBS_INPUT:

In this namelist section, observation files ("dfile" column) are tied to the observation variables used inside the GSI code ("dsis" column). For example, part of section OBS_INPUT shows:

```
&OBS_INPUT
  dmesh(1)=120.0,dmesh(2)=60.0,dmesh(3)=30,time_window_max=1.5,ext_sonde=.true. ,
/
OBS_INPUT::
!  dfile          dtype          dplat          dsis          dval          dthin dsfcalc
  prepbuf         ps             null           ps             1.0           0       0
  prepbuf         t             null           t             1.0           0       0
  prepbuf         q             null           q             1.0           0       0
  prepbuf         pw             null           pw            1.0           0       0
  satwndbuf       uv             null           uv            1.0           0       0
  prepbuf         uv             null           uv            1.0           0       0
  prepbuf         spd            null           spd            1.0           0       0
  prepbuf         dw             null           dw            1.0           0       0
  radarbuf        rw             null           rw            1.0           0       0
  prepbuf         sst            null           sst            1.0           0       0
  gpsrobuf        gps_ref        null           gps            1.0           0       0
  ssmirrbuf       pcp_ssmi      dmsp          pcp_ssmi      1.0          -1       0
  ...
  amsuabuf        amsua         n15           amsua_n15     10.0          2       0
  amsuabuf        amsua         n18           amsua_n18     10.0          2       0
  ...
```

This setup tells GSI that conventional observation variables ps, t, and q should be read in from the file prepbuf, while AMSU-A radiances from NOAA-15 and NOAA-18 satellites should be read in from the file **amsuabuf**. Deleting a particular line in &OBS_INPUT will turn off the use of the observation variable presented by the line in the GSI analysis but other variables under the same type can still be used. For example, if we delete:

```
amsuabuf        amsua         n15           amsua_n15     10.0          2       0
```

Then, the AMSU-A observation from NOAA-15 will not be used in the analysis but the AMSU-A observations from NOAA-18 will still be used.

The observation filename in "dfile" can be different from the sample script (*comgsi_namelist.ksh*). If the filename in "dfile" has been changed, the link from the BUFR files to the GSI recognized name in the run script also needs to be changed correspondingly. For example, if we change the "dfile" in **amsuabuf** for NOAA-15 to be amsuabuf_n15,

```
amsuabuf_n15    amsua         n15           amsua_n15     10.0          2       0
amsuabuf        amsua         n18           amsua_n18     10.0          2       0
```

Then a new link needs to be added in the run script:

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```
# Link to the radiance data
ln -s ${OBS_ROOT}/le_gdas1.t00z.1bamua.tm00.bufr_d amsuabufr
ln -s ${OBS_ROOT}/le_gdas1.t00z.1bamua.tm00.bufr_d amsuabufr_n15
```

The GSI will read NOAA-18 AMSU-A observations from file **amsuabufr** and NOAA-15 AMSU-A observations from file **amsuabufr_n15** based on the above changes to the run scripts and namelist. In this example, both **amsuabufr** and **amsuabufr_n15** are linked to the same BUFR file and NOAA-15 AMSU-A and NOAA-18 AMSU-A observations are still read in from the same BUFR file. If **amsuabufr** and **amsuabufr_n15** link to different BUFR files, then NOAA-15 AMSU-A and NOAA-18 AMSU-A will be read in from different BUFR files. The changeable filename in *dfile* gives GSI more flexibility to handle multiple data resources.

3. Use info files to control data usage

For each variable, observations can come from multiple platforms (data types or observation instruments). For example, surface pressure (ps) can come from METAR observation stations (data type 187) and rawinsonde (data type 120). There are several files named **info* in the GSI system (located in *./fix*) to control the usage of observations based on the observation platform. Table 4.2 is a list of info files and their function:

Table 4.2: The content of info files

File name in GSI	Function and Content
convinfo	Control the usage of conventional data, including tcp, ps, t, q, pw, sst, uv, spd, dw, radial wind (Level 2 <i>rw</i> and 2.5 <i>srw</i>), gps, and <i>pm2_5</i>
satinfo	Control the usage of satellite data. Instruments include AMSU-A/B, HIRS3/4, MHS, ssmi, ssmis, iasi, airs, sndr, cris, amsre, imgr, seviri, atms, avhrr3, etc., and satellites include NOAA 15, 17, 18, 19, aqua, GOES 11, 12, 13, METOP-A/B, NPP, DMSP 15,16,17,18,19,20, M08, M09, M10, etc.
ozinfo	Control the usage of ozone data, including sbuv6, 8 from NOAA 14, 16, 17, 18, 19. omi_aura, gome_metop-a, and mls_aura
pcpinfo	Control the usage of precipitation data, including pcp_ssmi and pcp_tmi
aeroinfo	Control the usage of aerosol data, including modis_aqua and modis_terra

The header of each info file includes an explanation of the content of the file. Here we discuss the two most commonly used info files:

- convinfo

The **convinfo** file controls the usage of conventional data. The following is part of the **convinfo** file:

```
!otype  type  sub iuse twindow numgrp ngroup nmitter gross ermax ermin var_b  var_pg ithin rmesh  pmesh  npred  pmot  ptime
tcp      112   0  1  3.0  0  0  0  75.0  5.0  1.0  75.0  0.000000  0  0.  0.  0  0.  0.
ps       120   0  1  3.0  0  0  0  4.0  3.0  1.0  4.0  0.000300  0  0.  0.  0  0.  0.
ps       132   0 -1  3.0  0  0  0  4.0  3.0  1.0  4.0  0.000300  0  0.  0.  0  0.  0.
ps       180   0  1  3.0  0  0  0  4.0  3.0  1.0  4.0  0.000300  0  0.  0.  0  0.  0.
ps       180  01  1  3.0  0  0  0  4.0  3.0  1.0  4.0  0.000300  0  0.  0.  0  0.  0.
ps       181   0  1  3.0  0  0  0  3.6  3.0  1.0  3.6  0.000300  0  0.  0.  0  0.  0.
ps       182   0  1  3.0  0  0  0  4.0  3.0  1.0  4.0  0.000300  0  0.  0.  0  0.  0.
ps       183   0 -1  3.0  0  0  0  4.0  3.0  1.0  4.0  0.000300  0  0.  0.  0  0.  0.
ps       187   0  1  3.0  0  0  0  4.0  3.0  1.0  4.0  0.000300  0  0.  0.  0  0.  0.
t        120   0  1  3.0  0  0  0  8.0  5.6  1.3  8.0  0.000001  0  0.  0.  0  0.  0.
t        126   0 -1  3.0  0  0  0  8.0  5.6  1.3  8.0  0.001000  0  0.  0.  0  0.  0.
t        130   0  1  3.0  0  0  0  7.0  5.6  1.3  7.0  0.001000  0  0.  0.  0  0.  0.
t        131   0  1  3.0  0  0  0  7.0  5.6  1.3  7.0  0.001000  0  0.  0.  0  0.  0.
```

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```

t      132  0  1  3.0  0  0  0  7.0  5.6  1.3  7.0  0.001000  0  0.  0.  0  0.  0.
t      133  0  1  3.0  0  0  0  7.0  5.6  1.3  7.0  0.004000  0  0.  0.  0  0.  0.
t      134  0 -1  3.0  0  0  0  7.0  5.6  1.3  7.0  0.004000  0  0.  0.  0  0.  0.
t      135  0 -1  3.0  0  0  0  7.0  5.6  1.3  7.0  0.004000  0  0.  0.  0  0.  0.
t      180  0  1  3.0  0  0  0  7.0  5.6  1.3  7.0  0.004000  0  0.  0.  0  0.  0.
t      180  0  1  3.0  0  0  0  7.0  5.6  1.3  7.0  0.004000  0  0.  0.  0  0.  0.
t      181  0 -1  3.0  0  0  0  7.0  5.6  1.3  7.0  0.004000  0  0.  0.  0  0.  0.
t      182  0  1  3.0  0  0  0  7.0  5.6  1.3  7.0  0.004000  0  0.  0.  0  0.  0.

```

The meaning of each column is explained in the header of the file and is listed in Table 4.3.

Table 4.3: list of the content for each convinfo column

Column Name	Content of the column
otype	observation variables (t, uv, q, etc.)
type	prepbufr observation type (if available)
sub	prepbufr subtype (not yet available)
iuse	flag if to use/not use / monitor data; = 1, use data, the data type will be read and used in the analysis after quality control; = 0, read in and process data, use for quality control, but do NOT assimilate; = -1, monitor data. This data type will be read in and monitored but not be used in the GSI analysis.
twindow	time window (+/- hours) for data used in the analysis
numgrp	cross validation parameter - number of groups
ngroup	cross validation parameter - group to remove from data use
nmitter	cross validation parameter - external iteration to introduce removed data
gross	gross error parameter - gross error
ermax	gross error parameter - maximum error
ermin	gross error parameter - minimum error
var_b	variational quality control parameter - b parameter
var_pg	variational quality control parameter - pg parameter
ithin	Flag to turn on thinning (0, no thinning, 1 - thinning)
rmesh	size of horizontal thinning mesh (in kilometers)
pmesh	size of vertical thinning mesh
npred	Number of bias correction predictors
pmot	Option to keep thinned data as monitored, 0: do not keep, other values: keep
ptime	time interval for thinning, 0, no temporal thinning, other values define time interval (less than six)

From this table, we can see that parameter "iuse" is used to control the usage of data and parameter "twindow" is used to control the time window of data usage. Parameters gross, ermax, and ermin are for gross quality control. Through these parameters, GSI can control how to use certain types of data in the analysis.

- satinfo

The *satinfo* file contains information about the channels, sensors, and satellites. It specifies observation error (cloudy or clear) for each channel, how to use the channels

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(assimilate, monitor, etc), and other useful information. The following is part of the content of *satinfo*. The meaning of each column is explained in Table 4.4.

!sensor/instr/sat	chan	iuse	error	error_cld	ermax	var_b	var_pg	cld_det
amsua_n15	1	1	3.000	20.000	4.500	10.000	0.000	-2
amsua_n15	2	1	2.200	18.000	4.500	10.000	0.000	-2
amsua_n15	3	1	2.000	12.000	4.500	10.000	0.000	-2
amsua_n15	4	1	0.600	3.000	2.500	10.000	0.000	-2
amsua_n15	5	1	0.300	0.500	2.000	10.000	0.000	-2
amsua_n15	6	-1	0.230	0.300	2.000	10.000	0.000	-2
amsua_n15	7	1	0.250	0.250	2.000	10.000	0.000	-2
amsua_n15	8	1	0.275	0.275	2.000	10.000	0.000	-2
amsua_n15	9	1	0.340	0.340	2.000	10.000	0.000	-2
amsua_n15	10	1	0.400	0.400	2.000	10.000	0.000	-2
amsua_n15	11	-1	0.600	0.600	2.500	10.000	0.000	-2
amsua_n15	12	1	1.000	1.000	3.500	10.000	0.000	-2
amsua_n15	13	1	1.500	1.500	4.500	10.000	0.000	-2
amsua_n15	14	-1	2.000	2.000	4.500	10.000	0.000	-2
amsua_n15	15	1	3.500	15.000	4.500	10.000	0.000	-2
hirs3_n17	1	-1	2.000	0.000	4.500	10.000	0.000	-1
hirs3_n17	2	-1	0.600	0.000	2.500	10.000	0.000	1
hirs3_n17	3	-1	0.530	0.000	2.500	10.000	0.000	1

Table 4.4: list of the content for each satinfo column

Column Name	Content of the column
sensor/instr/sat	Sensor, instrument, and satellite name
chan	Channel number for certain sensor
iuse	= 1, use this channel data; =-1, don't use this channel data
error	Variance for each satellite channel
error_cld	Variance for each satellite channel if cloudy
ermax	Error maximum for gross check to observations
var_b	Possible range of variable for gross errors
var_pg	Probability of gross error
icld_det	Use this channel in cloud detection if > 0

4.4 Domain Partition for Parallelization and Observation Distribution

The standard output file (*stdout*) has an information block that shows the distribution of different kinds of observations in each sub-domain. This block follows the observation input section. The following is the observation distribution from the case shown in Section 4.1. From the introduction, we know the prepbufr (conventional data), radiance BUFR files, and GPS BUFR files were used. In this list, the conventional observations (ps, t, q, pw, and uv), GPSRO (gps_ref), and radiance data (amusa, hirs4, and mhs from Metop-a, Metop-b, NOAA 15, and 18) were distributed among four sub-domains:

OBS_PARA: ps	1429	3190	4655	6774
OBS_PARA: t	2564	5200	7057	11128
OBS_PARA: q	2346	4626	6148	8128

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OBS_PARA: pw		65	80	63	49
OBS_PARA: uv		3358	6453	8091	11998
OBS_PARA: gps_ref		1799	1368	2664	3520
OBS_PARA: hirs4	metop-a	0	0	1146	1661
OBS_PARA: hirs4	n19	213	1020	0	0
OBS_PARA: hirs4	metop-b	0	0	85	555
OBS_PARA: amsua	n15	1458	2026	830	234
OBS_PARA: amsua	n18	2223	2318	108	0
OBS_PARA: amsua	n19	176	960	0	0
OBS_PARA: amsua	metop-a	0	0	1077	1559
OBS_PARA: amsua	metop-b	0	0	265	1829
OBS_PARA: mhs	n18	2550	2695	160	0
OBS_PARA: mhs	n19	246	1103	0	0
OBS_PARA: mhs	metop-a	0	0	1237	1809
OBS_PARA: mhs	metop-b	0	0	321	2128

This list is a good way to quickly check which kinds of data are used in the analysis and how they are distributed in the analysis domain.

4.5 Observation Innovation Statistics

The GSI analysis provides a set of files named *fort.2** to summarize observations fit to the current solution in each outer loop (except for *fort.220*, see explanation in the next section). The content of some of these files is listed in Table 4.5:

To help users understand the information inside these files, some examples are given in the following sub-sections with corresponding explanations.

4.5.1 Conventional observations

Example of files, including single level data (*fort.201*, *fort.205*, and *fort.213*)

```
current fit of surface pressure data, ranges in mb
-----
pressure levels (hPa)= 0.0 2000.0
it obs type stype count bias rms cpen qcpen
o-g 01 ps 120 0000 101 0.1813 0.6089 0.4711 0.4711
o-g 01 ps 180 0000 601 0.0378 0.6354 0.7171 0.7171
o-g 01 ps 180 0001 1428 0.1857 0.8555 0.7164 0.7164
o-g 01 ps 181 0000 610 0.1959 0.8991 0.7387 0.7387
o-g 01 ps 182 0000 1 0.9173 0.9173 2.8976 2.8976
o-g 01 ps 187 0000 11149 0.1999 0.7877 0.3360 0.3360
o-g 01 all 13890 0.1912 0.7931 0.4105 0.4105
o-g 01 ps rej 120 0000 1 -3.5799 3.5799 0.0000 0.0000
o-g 01 ps rej 180 0001 7 -0.3349 4.9273 0.0000 0.0000
o-g 01 ps rej 181 0000 47 1.1146 54.8539 0.0000 0.0000
o-g 01 ps rej 183 0000 3 10.4797 10.4836 0.0000 0.0000
o-g 01 ps rej 187 0000 52 4.2528 7.4112 0.0000 0.0000
o-g 01 rej all 110 2.7186 36.2804 0.0000 0.0000
o-g 01 ps mon 132 0000 1 0.9173 0.9173 0.2104 0.2104
o-g 01 ps mon 180 0000 113 0.0447 0.5158 0.8709 0.8709
o-g 01 ps mon 180 0001 24 0.2122 0.4050 0.3559 0.3559
o-g 01 ps mon 181 0000 207 0.0910 0.9492 1.2514 1.2514
o-g 01 ps mon 183 0000 1386 0.3974 1.0861 0.0000 0.0000
o-g 01 ps mon 187 0000 88 -0.0100 0.5290 0.7534 0.7534
o-g 01 mon all 1819 0.3188 1.0169 0.2378 0.2378
```

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Table 4.5: List of the content and units for each fort files

File Name	Variables in file	Ranges/units
<i>fort.201</i> <i>or</i> <i>fit_p1.analysis_time</i>	fit of surface pressure data	mb
<i>fort.202</i> <i>or</i> <i>fit_w1.analysis_time</i>	fit of u, v wind data	m/s
<i>fort.203</i> <i>or</i> <i>fit_t1.analysis_time</i>	fit of temperature data	K
<i>fort.204</i> <i>or</i> <i>fit_q1.analysis_time</i>	fit of moisture data	percent of qsaturation guess
<i>fort.205</i>	fit of precipitation water data	mm
<i>fort.206</i>	fit of ozone observations from sbuv6_n14 (, _n16, _n17, _n18), sbuv8_n16 (, _n17, _n18, _n19), omi_aura, gome_metop-a/b, and mls_aura	
<i>fort.207</i> <i>or</i> <i>fit_rad1.analysis_time</i>	fit of satellite radiance data, such as: am-sua_n15(, n16, n17, n18, metop-a, aqua, n19), amsub_n17, hirs3_n17, hirs4_n19 (, metop-a), etc.	
<i>fort.208</i>	fit of precipitation rate (pcp_ssmi and pcp_tmi)	
<i>fort.209</i>	fit of radar radial wind (rw)	
<i>fort.210</i>	fit of lidar wind (dw)	
<i>fort.211</i>	fit of radar superob wind data (srw)	
<i>fort.212</i>	fit of GPS data (refractivity or bending angle)	fractional difference
<i>fort.213</i>	fit of conventional sst data	C
<i>fort.214</i>	fit of tropical cyclone central pressure	
<i>fort.215</i>	fit of Lagrangian tracer data	
<i>fort.217</i>	fit of aerosol product (aod)	
<i>fort.218</i>	fit of wind gust	
<i>Fort.219</i>	fit of visibility	

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Example of files including multiple level data (*fort.202*, *fort.203*, and *fort.204*)

it	obs	type	styp	ptop pbot	1000.0 1200.0	900.0 1000.0	800.0 900.0	600.0 800.0	100.0 150.0	50.0 100.0	0.0 2000.0
o-g 01	uv	220	0000	count	44	223	223	478	437	519	4231
o-g 01	uv	220	0000	bias	0.26	0.51	0.04	0.35	0.99	0.97	0.59
o-g 01	uv	220	0000	rms	2.21	2.48	2.51	2.84	5.88	4.79	4.18
o-g 01	uv	220	0000	cpen	0.31	0.38	0.44	0.61	1.64	1.18	0.90
o-g 01	uv	220	0000	qcpn	0.31	0.38	0.44	0.60	1.63	1.18	0.89
o-g 01	uv	223	0000	count	0	6	16	88	32	8	331
o-g 01	uv	223	0000	bias	0.00	0.23	-0.29	0.85	-0.76	-4.89	0.27
o-g 01	uv	223	0000	rms	0.00	1.28	1.32	2.86	5.21	6.63	3.76
o-g 01	uv	223	0000	cpen	0.00	0.06	0.07	0.45	0.88	1.31	0.73
...											
o-g 01		all		count	1799	1726	2001	3050	469	527	13124
o-g 01		all		bias	0.05	0.91	0.78	0.63	0.87	0.88	0.62
o-g 01		all		rms	2.46	2.58	2.65	3.21	5.83	4.82	3.54
o-g 01		all		cpen	0.24	0.24	0.30	0.45	1.58	1.19	0.52
o-g 01		all		qcpn	0.23	0.24	0.30	0.44	1.58	1.19	0.52
o-g 01	uv rej	220	0000	count	0	0	0	0	0	0	800
o-g 01	uv rej	220	0000	bias	0.00	0.00	0.00	0.00	0.00	0.00	3.11
o-g 01	uv rej	220	0000	rms	0.00	0.00	0.00	0.00	0.00	0.00	6.41
...											
o-g 01		rej all		count	23	108	21	41	2	0	1008
o-g 01		rej all		bias	44.72	6.40	-0.30	-7.38	11.46	0.00	3.84
o-g 01		rej all		rms	56.08	16.60	6.87	10.47	43.15	0.00	12.28
o-g 01		rej all		cpen	0.00	0.00	0.00	0.00	0.00	0.00	0.00
o-g 01		rej all		qcpn	0.00	0.00	0.00	0.00	0.00	0.00	0.00
o-g 01	uv mon	220	0000	count	0	4	0	1	8	9	29
o-g 01	uv mon	220	0000	bias	0.00	1.07	0.00	6.04	-1.03	0.23	-2.21
o-g 01	uv mon	220	0000	rms	0.00	20.49	0.00	17.78	9.06	8.51	12.52
o-g 01	uv mon	220	0000	cpen	0.00	0.00	0.00	0.00	0.00	0.00	0.00
o-g 01	uv mon	220	0000	qcpn	0.00	0.00	0.00	0.00	0.00	0.00	0.00
...											
o-g 01		mon all		count	3573	7736	1004	563	8	9	13052
o-g 01		mon all		bias	-0.01	0.40	0.05	-0.30	-1.03	0.23	0.23
o-g 01		mon all		rms	2.55	3.25	4.67	5.98	9.06	8.51	3.60
o-g 01		mon all		cpen	0.81	1.21	1.67	1.28	0.00	0.00	1.12
o-g 01		mon all		qcpn	0.80	1.15	1.53	1.16	0.00	0.00	1.07
...											

Please note that five layers from 600 to 150 hPa have been deleted to make each row fit into one line. Only observation type 220 and 223 are shown as an example.

Table 4.6 lists the meaning of each item in file *fort.201-213* except file *fort.207*:

The contents of the fit files are calculated based on O-B or O-A for each observation. The detailed departure information about each observation is saved in the diagnostic files. For the content of the diagnostic files, please check the content of the array "rdiagbuf" in one of the setup subroutines for conventional data (for example, *setupt.f90*). We provide a tool in appendix A.2 to help users read in the information from the diagnostic files.

These fit files give lots of useful information on how data are analyzed by the GSI, such as how many observations are used and rejected, the bias and root mean squared (RMS) error for certain data types or for all observations, and how analysis results fit to the observation before and after analysis. Again, we use observation type 220 in *fort.202* (*fit_w1.2014061700*)

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Table 4.6: List of each item in file fort.201-213 (except fort.207).

Name	Explanation
<i>it</i>	outer loop number = 01: observation - background = 02: observation - analysis (after 1 st outer loop) = 03: observation - analysis (after 2 nd outer loop)
<i>obs</i>	observation variable type (such as uv or ps) and usage, which includes: blank: used in GSI analysis mon: monitored (read in but not assimilated by GSI) rej: rejected because of quality control in GSI
<i>type</i>	prepbufr observation type (see the BUFR User's Guide for details)
<i>styp</i>	prepbufr observation subtype (not used)
<i>ptop</i>	for multiple level data: pressure at the top of the layer
<i>pbot</i>	for multiple level data: pressure at the bottom of the layer
<i>count</i>	the number of observations summarized under observation types and vertical layers
<i>bias</i>	bias of observation departure for each outer loop (<i>it</i>)
<i>rms</i>	root mean square error of observation departure for each outer loop (<i>it</i>)
<i>cpen</i>	observation part of penalty (cost function)
<i>qcpen</i>	nonlinear qc penalty

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as an example to illustrate how to read this information. The fit information for observation type 220 (soundings) is listed below. Like the previous example, five layers from 600 to 150 hPa were deleted to make each row fit into one line. All fit information of observation type 220 is shown.

it	obs	type	styp	ptop	1000.0	900.0	800.0	600.0	100.0	50.0	0.0
				pbot	1200.0	1000.0	900.0	800.0	150.0	100.0	2000.0

o-g 01	uv	220	0000	count	44	223	223	478	437	519	4231
o-g 01	uv	220	0000	bias	0.26	0.51	0.04	0.35	0.99	0.97	0.59
o-g 01	uv	220	0000	rms	2.21	2.48	2.51	2.84	5.88	4.79	4.18
o-g 01	uv	220	0000	cpen	0.31	0.38	0.44	0.61	1.64	1.18	0.90
o-g 01	uv	220	0000	qcpen	0.31	0.38	0.44	0.60	1.63	1.18	0.89
...											
o-g 01	uv	rej	220	0000	count	0	0	0	0	0	800
o-g 01	uv	rej	220	0000	bias	0.00	0.00	0.00	0.00	0.00	3.11
o-g 01	uv	rej	220	0000	rms	0.00	0.00	0.00	0.00	0.00	6.41
o-g 01	uv	rej	220	0000	cpen	0.00	0.00	0.00	0.00	0.00	0.00
o-g 01	uv	rej	220	0000	qcpen	0.00	0.00	0.00	0.00	0.00	0.00
...											
o-g 01	uv	mon	220	0000	count	0	4	0	1	8	29
o-g 01	uv	mon	220	0000	bias	0.00	1.07	0.00	6.04	-1.03	-2.21
o-g 01	uv	mon	220	0000	rms	0.00	20.49	0.00	17.78	9.06	12.52
o-g 01	uv	mon	220	0000	cpen	0.00	0.00	0.00	0.00	0.00	0.00
o-g 01	uv	mon	220	0000	qcpen	0.00	0.00	0.00	0.00	0.00	0.00
...											
it	obs	type	styp	ptop	1000.0	900.0	800.0	600.0	100.0	50.0	0.0
				pbot	1200.0	1000.0	900.0	800.0	150.0	100.0	2000.0

o-g 02	uv	220	0000	count	44	223	223	478	437	519	4231
o-g 02	uv	220	0000	bias	0.21	0.05	-0.43	0.03	0.69	0.96	0.39
o-g 02	uv	220	0000	rms	2.13	2.26	2.29	2.56	5.19	4.55	3.73
o-g 02	uv	220	0000	cpen	0.32	0.31	0.37	0.50	1.27	1.07	0.71
o-g 02	uv	220	0000	qcpen	0.32	0.31	0.37	0.50	1.27	1.07	0.71
...											
o-g 02	uv	rej	220	0000	count	0	0	0	0	0	800
o-g 02	uv	rej	220	0000	bias	0.00	0.00	0.00	0.00	0.00	2.96
o-g 02	uv	rej	220	0000	rms	0.00	0.00	0.00	0.00	0.00	6.33
o-g 02	uv	rej	220	0000	cpen	0.00	0.00	0.00	0.00	0.00	0.00
o-g 02	uv	rej	220	0000	qcpen	0.00	0.00	0.00	0.00	0.00	0.00
...											
o-g 02	uv	mon	220	0000	count	0	4	0	1	8	29
o-g 02	uv	mon	220	0000	bias	0.00	2.16	0.00	5.80	-1.00	0.23
o-g 02	uv	mon	220	0000	rms	0.00	18.44	0.00	12.60	10.31	8.58
o-g 02	uv	mon	220	0000	cpen	0.00	0.00	0.00	0.00	0.00	0.00
o-g 02	uv	mon	220	0000	qcpen	0.00	0.00	0.00	0.00	0.00	0.00
...											
it	obs	type	styp	ptop	1000.0	900.0	800.0	600.0	100.0	50.0	0.0
				pbot	1200.0	1000.0	900.0	800.0	150.0	100.0	2000.0

o-g 03	uv	220	0000	count	44	223	223	478	437	519	4231
o-g 03	uv	220	0000	bias	0.28	0.08	-0.32	0.11	0.74	1.00	0.43
o-g 03	uv	220	0000	rms	2.11	2.17	2.20	2.38	5.00	4.46	3.60
o-g 03	uv	220	0000	cpen	0.31	0.29	0.34	0.43	1.18	1.03	0.65
o-g 03	uv	220	0000	qcpen	0.31	0.29	0.34	0.43	1.18	1.03	0.65
...											
o-g 03	uv	rej	220	0000	count	0	0	0	0	0	800

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```

o-g 03      uv rej 220 0000  bias    0.00  0.00  0.00  0.00  0.00  0.00  2.98
o-g 03      uv rej 220 0000  rms     0.00  0.00  0.00  0.00  0.00  0.00  6.35
o-g 03      uv rej 220 0000  cpen    0.00  0.00  0.00  0.00  0.00  0.00  0.00
o-g 03      uv rej 220 0000  qcpn    0.00  0.00  0.00  0.00  0.00  0.00  0.00
...
o-g 03      uv mon 220 0000 count      0      4      0      1      8      9      29
o-g 03      uv mon 220 0000 bias     0.00  1.86  0.00  6.09 -0.98  0.07 -0.94
o-g 03      uv mon 220 0000 rms     0.00 18.76  0.00 12.59 10.34  8.69 11.01
o-g 03      uv mon 220 0000 cpen    0.00  0.00  0.00  0.00  0.00  0.00  0.00
o-g 03      uv mon 220 0000 qcpn    0.00  0.00  0.00  0.00  0.00  0.00  0.00

```

In loop section o-g 01, from the count line, we can see there were 4231 sounding observations used in the analysis. Among them, 44 were within the 1000-1200 hPa layer. Also from the count lines, in the rejection and monitoring section, there were 800 observations rejected and 29 observations monitored. In the same loop section, from the bias line and rms lines, we can see the total bias and RMS error of O-B for the sounding information is 0.59 and 4.18. The bias and RMS error for each vertical layer can also be found in this file.

Next we can see bias and RMS error values from different loops, as shown with the comparison in the following three lines:

```

o-g 01      uv      220 0000  rms    2.21  2.48  2.51  2.84  5.88  4.79  4.18
o-g 02      uv      220 0000  rms    2.13  2.26  2.29  2.56  5.19  4.55  3.73
o-g 03      uv      220 0000  rms    2.11  2.17  2.20  2.38  5.00  4.46  3.60

```

These three lines show that the RMS error reduced from 4.18 (o-g 01, which is O-B) to 3.73 (o-g 02, which is O-A after the 1st outer loop) and then to 3.60 (o-g 03, which is O-A after the 2nd outer loop, which is also the final analysis result). The reduction in the RMS error shows that observation type 220 (sounding) was used in the GSI analysis to modify the background fields to fit to the observations.

4.5.2 Satellite Radiance

The file *fort.207* is the fit file for radiance data. Its content includes important information about the radiance data analysis.

The first part of the file *fort.207* lists the content that corresponds to those in the file **satinfo**, which is the info file to control radiance data usage.

```

RADINFO_READ:  jpch_rad=  2723
 1 amsua_n15      chan=   1 var=   3.000 varch_cld= 20.000 use=   1 ermax=   4.500 b_rad=   10.00 pg_rad=   0.00 icld_det=-2
 2 amsua_n15      chan=   2 var=   2.200 varch_cld= 18.000 use=   1 ermax=   4.500 b_rad=   10.00 pg_rad=   0.00 icld_det=-2
 3 amsua_n15      chan=   3 var=   2.000 varch_cld= 12.000 use=   1 ermax=   4.500 b_rad=   10.00 pg_rad=   0.00 icld_det=-2
 4 amsua_n15      chan=   4 var=   0.600 varch_cld=   3.000 use=   1 ermax=   2.500 b_rad=   10.00 pg_rad=   0.00 icld_det=-2
 5 amsua_n15      chan=   5 var=   0.300 varch_cld=   0.500 use=   1 ermax=   2.000 b_rad=   10.00 pg_rad=   0.00 icld_det=-2
...

```

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This shows there are 2723 channels listed in the *satinfo* file and the 2723 lines following this line include the details for each channel.

The second part of the file is a list of the coefficients for bias correction, after reading the *satbias_in* file:

```
RADINFO_READ: ***WARNING instrument/channel ahi_himawari8          16
not found in satbias_pc file - set to zero
RADINFO_READ: guess air mass bias correction coefficients below
1      amsua_n15  -1.965181  0.000000  39.068176  -0.923195  0.408460  0.000000  0.000000  0.001735  3.689709
          2.367472  8.862940  -1.848292
2      amsua_n15  -7.415403  0.000000  80.138019  -0.818472  -1.013216  0.000000  0.000000  0.014279  2.638804
          8.315096  22.041458  -1.564773
...
```

Each channel has 12 coefficients listed in a line. Therefore, there are 2723 lines of radiance bias correction coefficients for all channels, though some of the coefficients are zero.

The 3rd part of the *fort.207* file is similar to other fit files with content repeated in three sections, providing detailed statistics about the data in stages before the 1st outer loop, between the 1st and 2nd outer loops, and after the 2nd outer loop. The results before the 1st outer loop are used here as an example to explain the content of the results:

- Summaries for various statistics as a function of observation type

sat	type	penalty	nobs	iland	isnoice	icoast	ireduce	ivar1	nlgross
metop-a	hirs4	371.46818888	385	143	0	25	0	1452	0
	qcpenalty		qc1	qc2	qc3	qc4	qc5	qc6	qc7
		371.46818888	0	0	1651	3330	0	0	0

sat	type	penalty	nobs	iland	isnoice	icoast	ireduce	ivar1	nlgross
metop-b	hirs4	0.00000000	34	25	0	0	0	139	0
	qcpenalty		qc1	qc2	qc3	qc4	qc5	qc6	qc7
		0.00000000	0	0	97	279	0	0	0

6

```
rad total penalty_all= 13986.0933845987129
rad total qcpenalty_all= 13986.0933845987129
rad total failed nonlinqc= 0
```

...

Table 4.7 outlines the meaning of each item in the above statistics:

Note: One radiance observation may include multiple channels, and not all channels are necessarily used in the analysis.

- Summaries for various statistics as a function of channel

1	2	3	4	5	6	7	8	9	10	11
1	1	amsua_n15	915	247	3.000	0.6623748	1.2673027	0.2414967	2.4209147	2.0627099
2	2	amsua_n15	902	261	2.000	0.5209886	1.1773765	0.2626527	2.2705374	1.9414234
3	3	amsua_n15	1114	48	2.000	1.2349467	-1.5090660	0.3421312	1.9416738	1.2218089
4	4	amsua_n15	1162	0	0.600	-0.2114506	-0.3195250	0.5286580	0.6271317	0.5396275
5	5	amsua_n15	1162	3	0.300	-0.0888496	-0.1685199	0.4352839	0.2734131	0.2153040
6	6	amsua_n15	2093	3	-0.230	-1.8141034	-0.0695503	0.8415303	0.2646033	0.2552992
7	7	amsua_n15	2342	28	0.250	-0.0830293	0.0453835	0.6141132	0.2522587	0.2481427
8	8	amsua_n15	2311	59	0.275	0.0201868	0.1758359	0.7782391	0.3173862	0.2642267
9	9	amsua_n15	2124	246	0.340	0.1262987	0.5008846	1.4907547	0.5544190	0.2376868
10	10	amsua_n15	82	2288	0.400	0.6700157	0.8263504	2.0387089	0.8304076	0.0819868
15	15	amsua_n15	862	300	3.000	0.8244923	-2.2960147	0.3769625	2.7445692	1.5036544

...

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Table 4.7: Summary of the radiance observation fit file (fort.207)

Name	Explanation
<i>sat</i>	satellite name
<i>type</i>	instrument type
<i>penalty</i>	contribution to cost function from this observation type
<i>nobs</i>	number of good observations used in the assimilation
<i>iland</i>	number of observations over land
<i>isnoice</i>	number of observations over sea ice and snow
<i>icoast</i>	number of observations over coast
<i>ireduce</i>	number of observations that reduce qc bounds in tropics
<i>ivarl</i>	number of observations removed by gross check
<i>nlgross</i>	number of observation removed by nonlinear qc
<i>qcpenalty</i>	nonlinear qc penalty from this data type
<i>qc1-7</i>	number of observations whose quality control criteria has been adjusted by each qc method (1-7). For details, see the Radiance Chapter of the Advanced User's Guide
<i>rad total penalty_all</i>	summary of penalty for all radiance observation types
<i>rad total qcpenalty_all</i>	summary of qcpenalty for all radiance observation types
<i>rad total failed nonlingc</i>	summary of observations removed by nonlinear qc for all radiance observation types

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```

63 4 hirs4_metop-a 11 217 0.400 0.9717384 0.9536444 1.8583033 0.9570578 0.0807581
64 5 hirs4_metop-a 81 4 0.360 0.1655806 -0.2231640 0.3231878 0.3433454 0.2609289
65 6 hirs4_metop-a 25 27 0.460 -0.8415009 -1.1454801 2.6082742 1.1558578 0.1545404
66 7 hirs4_metop-a 20 3 0.570 -0.9563649 -1.1287970 1.4138248 1.1699278 0.3074870
67 8 hirs4_metop-a 23 0 1.000 1.3954294 0.6716651 0.1668694 0.9134028 0.6190078
...

```

Table 4.8 lists the meaning of each column in the above statistics:

Table 4.8: Content of fit statistics for each channel in the fort.207 file.

Column #	Content
1	series number of the channel in satinfo file
2	channel number for certain radiance observation type
3	radiance observation type (for example: amsua_n15)
4	number of observations (nobs) used in GSI analysis within this channel
5	number of observations (nobs) tossed by gross check within this channel
6	variance for each satellite channel
7	bias (observation-guess before bias correction)
8	bias (observation-guess after bias correction)
9	penalty contribution from this channel
10	square root of (observation-guess with bias correction)**2
11	standard deviation

- Final summary for each observation type

```

it      satellite instrument # read # keep # assim penalty qcpnlty cpen qccpen
o-g 01 rad n16 hirs3 0 0 0 0.0000 0.0000 0.0000 0.0000
o-g 01 rad n17 hirs3 0 0 0 0.0000 0.0000 0.0000 0.0000
o-g 01 rad metop-a hirs4 13832 7315 651 371.47 371.47 0.57061 0.57061
o-g 01 rad n18 hirs4 0 0 0 0.0000 0.0000 0.0000 0.0000
o-g 01 rad n19 hirs4 0 0 0 0.0000 0.0000 0.0000 0.0000
o-g 01 rad metop-b hirs4 2527 646 0 0.0000 0.0000 0.0000 0.0000
o-g 01 rad g11 goes_img 0 0 0 0.0000 0.0000 0.0000 0.0000
o-g 01 rad g12 goes_img 0 0 0 0.0000 0.0000 0.0000 0.0000
o-g 01 rad aqua airs 0 0 0 0.0000 0.0000 0.0000 0.0000
o-g 01 rad n15 amsua 43515 30810 12976 8854.4 8854.4 0.68236 0.68236
o-g 01 rad n18 amsua 30690 25575 8203 3642.7 3642.7 0.44407 0.44407
o-g 01 rad n19 amsua 0 0 0 0.0000 0.0000 0.0000 0.0000
o-g 01 rad metop-a amsua 4890 3878 1466 652.20 652.20 0.44489 0.44489
o-g 01 rad metop-b amsua 810 718 294 201.79 201.79 0.68636 0.68636
o-g 01 rad aqua amsua 0 0 0 0.0000 0.0000 0.0000 0.0000
o-g 01 rad n17 amsub 0 0 0 0.0000 0.0000 0.0000 0.0000
...

```

Table 4.9 lists the meaning of each column in the above statistics:

Similar to other fit files, a comparison between results from different outer loops can give us very useful information on how much impact each channel and data type has in the GSI.

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Table 4.9: Content of the final summary section for the fort.207 file.

Name	Explanation
<i>it</i>	stage (o-g 01 rad = before 1st outer loop for radiance data)
<i>satellite</i>	satellite name (n16=NOAA -16)
<i>instrument</i>	instrument name (HIRS-3)
<i># read</i>	number of data (channels) read in within analysis time window and domain
<i># keep</i>	number of data (channels) kept after data thinning
<i># assim</i>	number of data (channels) used in analysis (passed all qc process)
<i>penalty</i>	contribution from this observation type to cost function
<i>qcpnlty</i>	nonlinear qc penalty from this data type
<i>cpen</i>	penalty divided by (the number of data assimilated)
<i>qccpen</i>	qcpnlty divided by (the number of data assimilated)

4.6 Convergence Information

There are two ways to check the convergence information for each iteration of the GSI:

1. Standard output file (*stdout*):

The value of the cost function and norm of the gradient for each iteration are listed in the file *stdout*.

The following is an example showing the iterations from the first outer loop:

```

GLBSOI:  START pcgsoi  jiter=          1
pcgsoi:  gnorm(1:2),b=  9.869857497554413276E+05  9.869857497554413276E+05  0.000000000000000000E+00
Initial cost function =  3.915930707165839704E+04
Initial gradient norm =  9.934715646436194447E+02
cost,grad,step,b,step? =  1  0  3.915930707165839704E+04  9.934715646436194447E+02  4.821051367939106942E-03  0.000000000000000000E+00  good
pcgsoi:  gnorm(1:2),b=  4.101618851769856410E+05  4.101618851769853500E+05  4.155702200144395508E-01
cost,grad,step,b,step? =  1  1  3.440099807266351127E+04  6.404388223530688720E+02  5.837551195665082078E-03  4.155702200144395508E-01  good
pcgsoi:  gnorm(1:2),b=  4.679752995326447999E+05  4.679752995326457312E+05  1.140952673675940110E+00
cost,grad,step,b,step? =  1  2  3.200665706943234181E+04  6.840872017021256397E+02  3.727016244273905453E-03  1.140952673675940110E+00  good
pcgsoi:  gnorm(1:2),b=  2.579351145588153449E+05  2.579351145588165091E+05  5.511724973869557287E-01
...
cost,grad,step,b,step? =  1  9  2.515019683100273687E+04  2.511847728725914237E+02  5.552768045789822915E-03  9.872916643640710088E-01  good
pcgsoi:  gnorm(1:2),b=  5.868194961445817898E+04  5.868194961445837544E+04  9.300748853414525508E-01
cost,grad,step,b,step? =  1  10  2.479985164931967302E+04  2.422435749704379191E+02  5.481216572289848883E-03  9.300748853414525508E-01  good

```

The following are the iterations from the second outer loop:

```

Initial cost function =  2.792919782749931983E+04
Initial gradient norm =  4.241369976412337337E+02
cost,grad,step,b,step? =  2  0  2.792919782749931983E+04  4.241369976412337337E+02  4.301269527061492466E-03  0.000000000000000000E+00  good
pcgsoi:  gnorm(1:2),b=  1.641799966598818428E+05  1.641799966598813771E+05  9.126577097845959274E-01
cost,grad,step,b,step? =  2  1  2.715543302058953486E+04  4.051913087171068923E+02  3.965037770683364597E-03  9.126577097845959274E-01  good
pcgsoi:  gnorm(1:2),b=  7.369958699881075881E+04  7.369958699881142820E+04  4.488950450613589660E-01
cost,grad,step,b,step? =  2  2  2.650445313264244396E+04  2.714766785541821719E+02  6.065878225371862040E-03  4.488950450613589660E-01  good
pcgsoi:  gnorm(1:2),b=  5.010117245725526300E+04  5.010117245725520479E+04  6.798026216627733875E-01
...
cost,grad,step,b,step? =  2  9  2.488459155328830457E+04  1.491452772915179139E+02  6.349376870755444636E-03  9.522651754285673675E-01  good
pcgsoi:  gnorm(1:2),b=  2.350218829989638834E+04  2.350218829989681763E+04  1.056548139732611746E+00
cost,grad,step,b,step? =  2  10  2.474335402213209454E+04  1.533042344486817683E+02  5.304880218120413757E-03  1.056548139732611746E+00  good

```

Here, we can see the number of outer and inner loops (minimization iteration). The meaning of the names (**bold**) used in *stdout* are explained in the following:

- **cost**: the cost function values, (=J)

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- grad: inner product of gradients (norm of the gradient (Y*X))
- step: stepsize
- b: parameter to estimate the new search direction

As a quick check, the cost function reduced from $3.915930707165839704E+04$ to $2.479985164931967302E+04$ in the 1st outer loop and reduced from $2.792919782749931983E+04$ to $2.474335402213209454E+04$ in the 2nd outer loop.

2. Convergence information in file *fort.220*:

In file *fort.220*, users can find more detailed minimization information about each iteration. A detailed description and example are provided in the Advanced User's Guide.

To evaluate the iteration convergence, we usually make plots based on the information from *fort.220*, such as the value of the cost function and the norm of the gradient. The following are example plots showing the evolution of the cost function and the norm of the gradient in different outer loops:

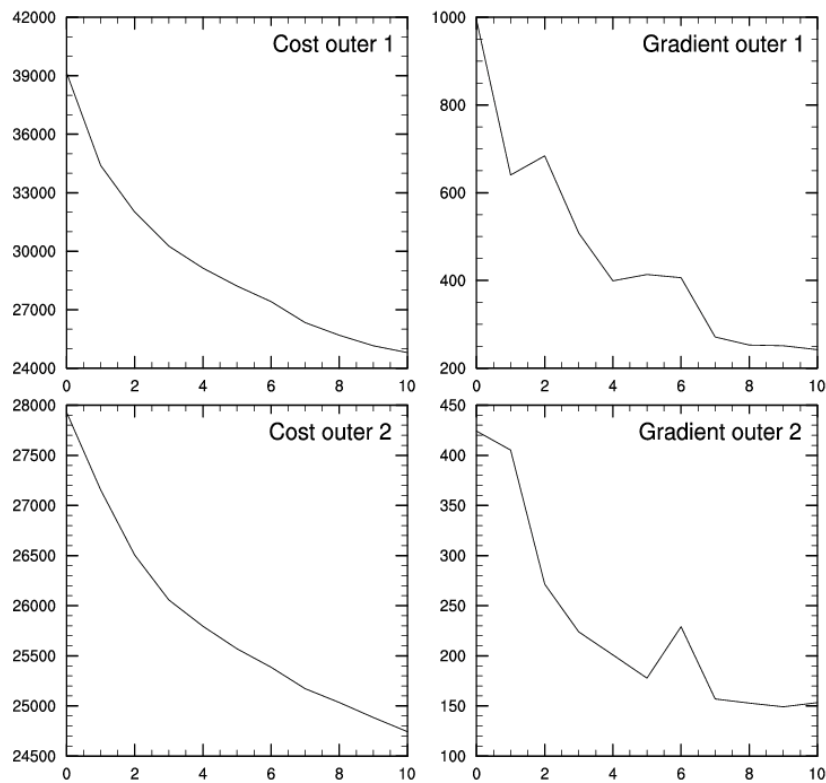


Figure 4.2: Evolution of the cost function (left column) and the norm of the gradient (right column) in the first outer loop (top row) and the second outer loop (bottom row). The Y-axis is the iteration number.

Scripts are available in the release code to read convergence information from *fort.220* and produce the above plots. Please see Section A.3 for information on where to locate and how to run these scripts.

4.7 Conventional Observation Errors

Each observation type has its own observation errors. In this section, we introduce several topics related to the conventional observation error processing in GSI. The observation error for satellite radiance and its adjustment is discussed in the Advanced User's Guide.

4.7.1 Getting Original Observation Errors

For the global GSI analysis, when `oberrflg` (a namelist option in section `&obsqc`) is true, observation errors are generated based on an external observation error table according to the observation type. Otherwise, observation errors are read in from the PrepBUFR file.

For regional analyses, GSI forces the use of an external observation error table to get observation errors no matter what the `oberrflg` is set to (`oberrflg` is forced to be true for regional runs in *gsimod.F90*).

The external observation error table file, *errtable*, includes observation errors for all types of conventional observations. It is copied from the `./fix` directory by the run script. This release package has three sample external observation error table files, *nam_errtable.r3dv*, *prepobs_errtable.global*, and *rtma/new_rtma_nam_errtable.r3dv* in the `./fix` directory. The *nam_errtable.r3dv* is used in the sample run script as a default observation error table. The observation error file is a text file that can be easily edited to tune the error values. The following shows a portion of *nam_errtable.r3dv* file for rawinsondes and its description of each column in Table 4.10:

```

Column # 1      2      3      4      5      6
120 OBSERVATION TYPE
0.11000E+04 0.12696E+01 0.60737E+00 0.10000E+10 0.76322E+00 0.10000E+10
0.10500E+04 0.13282E+01 0.66294E+00 0.10000E+10 0.76322E+00 0.10000E+10
0.10000E+04 0.13932E+01 0.74223E+00 0.10000E+10 0.76322E+00 0.10000E+10
0.95000E+03 0.14390E+01 0.83688E+00 0.10000E+10 0.79899E+00 0.10000E+10
0.90000E+03 0.14354E+01 0.94025E+00 0.10000E+10 0.83561E+00 0.10000E+10
0.85000E+03 0.13669E+01 0.10439E+01 0.10000E+10 0.87224E+00 0.10000E+10

220 OBSERVATION TYPE
0.11000E+04 0.10000E+10 0.10000E+10 0.18521E+01 0.10000E+10 0.10000E+10
0.10500E+04 0.10000E+10 0.10000E+10 0.20636E+01 0.10000E+10 0.10000E+10
0.10000E+04 0.10000E+10 0.10000E+10 0.22799E+01 0.10000E+10 0.10000E+10
0.95000E+03 0.10000E+10 0.10000E+10 0.24211E+01 0.10000E+10 0.10000E+10
0.90000E+03 0.10000E+10 0.10000E+10 0.24934E+01 0.10000E+10 0.10000E+10
0.85000E+03 0.10000E+10 0.10000E+10 0.25155E+01 0.10000E+10 0.10000E+10

```

Table 4.10: Description of each column in the observation error table file

Column #	1	2	3	4	5	6
<i>Content</i>	Pressure	T	RH	UV	Ps	Pw
<i>Unit</i>	hPa	degree C	percent/10	m/s	mb	kg/m2(or mm)

For each observation type, the error table has six columns and 33 rows (levels). The 1st column prescribes 33 pressure levels, covering 1100 hPa to 0 hPa. Columns 2-6 prescribe

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observation errors for temperature (T), relative humidity (RH), horizontal wind component (UV), surface pressure (Ps), and the total column precipitable water (Pw). The missing value is 0.10000E+10.

The observation error table for each observation type starts with the observation type number defined for the PrepBUFR files, such as:

```
120 OBSERVATION TYPE
220 OBSERVATION TYPE
```

The PrepBUFR data type numbers 100-199 are for temperature (T), moisture (q), and surface pressure (Ps) observations, while numbers 200-299 are for horizontal wind component (UV) observations. A detailed explanation of each data type number can be found in the following table on the EMC website:

http://www.emc.ncep.noaa.gov/mmb/data_processing/prepbufr.doc/table_2.htm

For more details on PrepBUFR/BUFR, please check the BUFR/PrepBUFR User's Guide, which is freely available at the DTC BUFR/PrepBUFR website:

<http://www.dtcenter.org/com-GSI/BUFR/index.php>

4.7.2 Observation Error Gross Error Check within GSI

The gross error check is an important quality control step to exclude questionable observations that degrade the analysis. Users can adjust the threshold of the gross error check for each data type within the *convinfo* file to make the gross error check more or less strict for a certain data type. For example, the following is a part of the *convinfo* file without the last five columns:

!otype	type	sub	iuse	twindow	numgrp	ngroup	nmitter	gross	ermax	ermin	var_b	var_pg	ithin
ps	183	0	-1	3.0	0	0	0	4.0	3.0	1.0	4.0	0.000300	0
ps	187	0	1	3.0	0	0	0	4.0	3.0	1.0	4.0	0.000300	0
t	120	0	1	3.0	0	0	0	8.0	5.6	1.3	8.0	0.000001	0
t	126	0	-1	3.0	0	0	0	8.0	5.6	1.3	8.0	0.001000	0

The gross check for each data type is controlled by columns "gross", "ermax", and "ermin." If an observation has observation error set to "obserror," then a gross check ratio is calculated:

$$ratio = (Observation-Background)/max(ermin, min(ermax, obserror))$$

If *ratio* > *gross*, then this observation fails the gross check and will not be used in the analysis. The unused observation is indicated as a "rejection" in the fit files.

4.8 Background Error Covariance

The GSI package has several files in *./fix* to hold the pre-computed background error statistics for different GSI applications with different grid configurations. Within the *./fix* subdirectories *./fix/Big_Endian* and *./fix/Little_Endian* contain the fix files corresponding to each endianness. Since the GSI code has a build-in mechanism to interpolate the input background error matrix to any desired analysis grid, the following two background error files can be used to specify the B matrix for any GSI regional application.

- *nam_nmmstat_na.gcv* : contains the regional background error statistics, computed using forecasts from the NCEP's NAM model covering North America. The values of this B matrix cover the northern hemisphere with 93 latitude lines, from -2.5 degrees to 89.5 degrees with 60 vertical sigma levels from 0.9975289 to 0.01364.
- *nam_glb_berror.f77.gcv* : contains the global background errors based on NCEP's GFS model, a global forecast model. The values of this B matrix cover the globe with 192 latitude lines from -90 degrees to 90 degrees and 42 vertical sigma levels from 0.99597 to 0.013831.

The background error matrix files listed above are in Big Endian binary form (therefore located in the *Big_Endian* directory). In the *Little_Endian* directory, *nam_nmmstat_na.gcv* and *nam_glb_berror.f77.gcv* are their Little Endian versions for certain computer platforms that cannot compile GSI with the Big Endian option. In this release version, GSI can be compiled with the Big Endian option with PGI and Intel.

4.8.1 Tuning Background Error Covariance through the Namelist and Anavinfo File

The final background error covariance matrix used in the GSI analysis is the content from the fixed file "berror", which is a copy of *nam_nmmstat_na.gcv* or *nam_glb_berror.f77.gcv*, multiplied by several factors set by the namelist and the **anavinfo** file.

In GSI namelist, three variables are used for tuning horizontal and vertical impact scales:

- *vs* scale factor for vertical correlation lengths for background error
- *hzsc1(3)* scale factor for three scales specified for horizontal smoothing
- *hswgt(3)* weights to apply to each horizontal scales

In the GSI anavinfo file, the column **as/tsfc_sdv** in the *control_vector* section are factors for tuning the variance of each analysis control variable.

These values can be used to tune the background error covariance used in the GSI analysis. For each background error matrix file, there are recommended values for these parameters listed in table 4.11.

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Table 4.11: Recommended tuning values for the provided B matrix.

	Global	Regional
<i>fixed B matrix</i>	nam_glb_berror.f77.gcv	nam_nmmstat_na.gcv
<i>vs</i>	0.7	1.0
<i>hzscl</i>	1.7, 0.8, 0.5	0.373,0.746,1.50
<i>hswgt</i>	0.45, 0.3,0.25	0.45, 0.3,0.25
<i>ss/tsfc_sdv</i>	<pre>control_vector:: !var as/tsfc_sdv sf 0.60 vp 0.60 ps 0.75 t 0.75 q 0.75 oz 0.75 sst 1.00 cw 1.00 stl 3.00 sti 3.00</pre>	<pre>control_vector:: !var as/tsfc_sdv sf 1.00 vp 1.00 ps 0.50 t 0.70 q 0.70 oz 0.50 sst 1.00 cw 1.00 stl 1.00 sti 1.00</pre>

4.9 Analysis Increments

Analysis increments are defined as the difference between analysis results and the background (A-B). A plot of analysis increments can help users understand how the analysis procedure modifies the background fields according to observations, background and observation error covariances, and other constraints. You can either calculate *analysis-guess* and plot the difference field or use the tools introduced in Appendix A.4 to make analysis increment figures for different analysis fields.

4.10 Running Time and Memory Usage

In addition to analysis increments, run time and memory usage are other important features of an analysis system, especially for operational code like the GSI.

The GSI standard output file (*stdout*) gives the GSI start time and end time of the analysis at the beginning and end of the file. For example:

```
* . . . . .
PROGRAM GSI_ANL HAS BEGUN. COMPILED 1999232.55      ORG: NP23
STARTING DATE-TIME  JUL 02,2016  20:36:21.760  184  SAT  2457572
```

...

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```
ENDING DATE-TIME    JUL 02,2016  20:43:40.422  184  SAT   2457572
PROGRAM GSI_ANL HAS ENDED.
* . * . * . * . * . * . * . * . * . * . * . * . * . * . * . * . * . * . * .
```

This tells us the analysis started at 20:36:21.760 and ended at 20:43:40.422, which means GSI used 7 minutes and 19 seconds to finish.

Following the ending date-time, there is a resource statistics section at the end of the *stdout* file, which gives information about run time and memory usage for the analysis:

```
*****RESOURCE STATISTICS*****
The total amount of wall time           = 438.663534
The total amount of time in user mode   = 427.578998
The total amount of time in sys mode    = 9.457562
The maximum resident set size (KB)     = 2020132
Number of page faults without I/O activity = 312762
Number of page faults with I/O activity = 0
Number of times filesystem performed INPUT = 0
Number of times filesystem performed OUTPUT = 0
Number of Voluntary Context Switches    = 7641
Number of InVoluntary Context Switches  = 851
*****END OF RESOURCE STATISTICS*****
```



5

GSI Applications

In this chapter, information from the previous chapters will be applied to the regional GSI cases with different data sources and assimilation methods, as well as the global and chemical analysis. These examples are to give users a clear idea of how to set up GSI with various configurations and properly check the run status and analysis results in order to determine if a particular GSI application was successful. Note that the regional examples here only use the WRF-ARW system - WRF-NMM runs are similar, but require different background and namelist options.

For illustrations of all the cases, it is assumed that the reader has successfully compiled GSI on a local machine.

For regional WRF-ARW case studies, users should have the following data available:

1. Background file
 - When using WRF, WPS and real.exe will be run to create a WRF input file: `wrfinput_<domain>`
 - Users can also use the forecast files from running WRF: `wrfout_<domain>_<yyyy-mm-dd_hh:mm:ss>`. In this chapter, WRF forecast files are being used as the background files.
2. Conventional data
 - Real time RAP PrepBUFR data can be obtained from the server:
`ftp://ftpprd.ncep.noaa.gov/pub/data/nccf/com/rap/prod`
Note: A RAP PrepBUFR file, prepared by the NCEP and used by the NOAA Rapid Refresh (RAP) and High Resolution Rapid Refresh (HRRR), was chosen to increase the amount of data used in the analysis (compared to a GDAS PrepBUFR file)
3. Radiance data and GPS RO data
 - Real time GDAS BUFR files can be obtained from the following server:

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`ftp://ftpprd.ncep.noaa.gov/pub/data/nccf/com/gfs/prod`

Note: GDAS data was chosen to get better coverage for radiance and GPS Radio Occultation (RO).

The background and observation files for the global and chemical analysis will be introduced in the later sections.

The following cases will give users an example of a successful GSI run with various data sources. Users are welcome to download these example data from the GSI users' webpage (online case) or create a new background and get the observational data from the above servers.

The background and observations used in the regional WRF-ARW cases are as follows:

1. Background files: `wrfout_d01_2018-08-12_12:00:00`
 - The horizontal grid spacing is 45-km with 33 vertical sigma levels

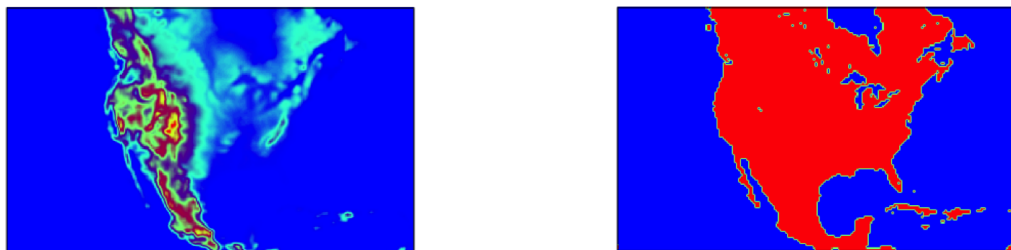


Figure 5.1: The terrain (left) and land mask (right) of the background used in this case study.

2. Conventional data: RAP PrepBUFR data from 12 UTC 12 August 2018
 - File: `rap.t12z.prepbufr.tm00`
3. Radiance and GPS RO data: GDAS BUFR data from 12 UTC 12 August 2018
 - Files: `gdas1.t12z.1bamua.tm00.bufr_d`
`gdas1.t12z.1bhrs4.tm00.bufr_d`
`gdas1.t12z.1bmhs.tm00.bufr_d`
`gdas1.t12z.gpsro.tm00.bufr_d`

This case study was run on a Linux cluster. As of version 3.2, the BUFR/PrepBUFR files do not need to be byte-swapped to little endian format. BUFRLIB can automatically handle byte order issues.

Assume the background file is located at:

`casedata/2018081212/bkg`

All the observations are located at:

`casedata/2018081212/obs`

And the GSI release version is located at:

`code/comGSI(version_number)_EnKF(version_number)`

5.1 Assimilating Conventional Observations with Regional GSI

5.1.1 Run Script

With GSI successfully compiled and background and observational data acquired, move to the `./ush` directory under `./comGSIv3.7_EnKFv1.3` to run the GSI using the sample script `comgsi_run_regional.ksh`. The `comgsi_run_regional.ksh` script must be modified in several places before running:

- Set up batch queuing system.

To run GSI with multi-processors, a job queuing section has to be added at the beginning of the `comgsi_run_regional.ksh` script. The set up of the job queue depends on the machine and the job control system. More setup examples are described in section 3.2.2. The following example is set up to run on a Linux cluster supercomputer with PBS. The job section is as follows:

```
#PBS -A ??????
#PBS -l walltime=00:20:00
#PBS -N cg02-case03-conv
#PBS -l select=1:ncpus=4:mpiprocs=4
#PBS -q premium
#PBS -o out.case03-conv
#PBS -j oe
```

In order to find out how to set up the job section, a good start is to use an existing MPI job script and copy the job section over.

- Set up the number of processors and the job queue system used. For this example, LINUX_PBS and four processors are used:

```
GSIPROC=4
ARCH='LINUX_PBS'
```

- Set up the case data, analysis time, GSI fix files, GSI executable, and CRTM coefficients:

Set up analysis time:

```
ANAL_TIME=2018081212
```

Set up a working directory, which will hold all the analysis results. This directory must have correct write permissions, as well as enough space to hold the output.

```
JOB_DIR=the_job_directory
#normally you put run scripts here and submit jobs form here, require a copy of gsi.x at this directory
RUN_NAME=a_descriptive_run_name_such_as_case05_3denvar_etc
```

Set path to the background directory and file:

```
BK_ROOT=casedata/2018081212/bkg
BK_FILE=${BK_ROOT}/wrfout_d01_2018-08-12_12:00:00
```

Set path to the observation directory and the PrepBUFR file within the observation directory. All observations to be assimilated should be in the observation directory.

```
OBS_ROOT=casedata/2018081212/obs
PREPBUFR=${OBS_ROOT}/rap.t${HH}z.prepbuf.fr.tm00
```

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Set up the GSI system used for this case, including the paths of fix files and the CRTM coefficients as well as the location of the GSI executable and the namelist file. Please be sure to modify accordingly.

```
GSI_ROOT=the_comgsi_main_directory_where_src/ush/fix/etc_are_located
CRTM_ROOT=the_CRTM_directory
ENS_ROOT=the_directory_where_ensemble_backgrounds_are_located
#ENS_ROOT is not required if not running hybrid EnVAR
HH='echo $ANAL_TIME | cut -c9-10'
GSI_EXE=${JOB_DIR}/gsi.x #assume you have a copy of gsi.x here
WORK_ROOT=${JOB_DIR}/${RUN_NAME}
FIX_ROOT=${GSI_ROOT}/fix
GSI_NAMELIST=${GSI_ROOT}/ush/comgsi_namelist.sh
```

- Set which background and background error file to use:

```
bk_core=ARW
bkcv_option=NAM
if_clean=clean
```

This example uses the ARW NetCDF background; therefore `bk_core` is set to 'ARW'. The regional background error covariance file is used in this case, as set by `bkcv_option=NAM`. Finally, the run scripts is set to clean the run directory to delete all temporary intermediate files.

5.1.2 Run GSI and Check the Run Status

Once the run script is set up properly for the case and machine, GSI can be run through the run script. On our test machine, the GSI run is submitted as follows:

```
$ qsub comgsi_run_regional.ksh
```

Next, move to the working directory and check the details. While GSI is running, the contents of this directory should include files such as:

```
imgr_g12.TauCoeff.bin      ssmi_f15.SpcCoeff.bin
imgr_g13.SpcCoeff.bin     ssmi_f15.TauCoeff.bin
imgr_g13.TauCoeff.bin     ssmis_f16.SpcCoeff.bin
```

These are CRTM coefficient files that have been linked to this run directory through the GSI run script. Additionally, many other files are linked or copied to this run directory or generated during the run, such as:

- `stdout`: standard output file
- `wrf_inout`: background file
- `gsiparm.anl`: GSI namelist
- `prepbufr`: prepBUFR file for conventional observation
- `convinfo`: data usage control for conventional data
- `berror_stats`: background error file
- `errtable`: observation error file

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The presence of these files indicates that the GSI run scripts have successfully set up a run environment for GSI and that the GSI executable is running. While GSI is still running, checking the content of the standard output file (*stdout*) can monitor the status of the GSI analysis:

```
$ tail stdout
cost_grad_step_b_step? = 1 17 1.377774626348101629E+04 7.451202547871611515E+00 4.626071752079010935E+01 6.438326819051856109E-01 good
cost_grad_step_b_step? = 1 18 1.377517784904208384E+04 5.686038514327907301E+00 4.998548357396024500E+01 5.823269047738203197E-01 good
cost_grad_step_b_step? = 1 19 1.377356176667382715E+04 4.191035073402087363E+00 5.924613648750278116E+01 5.432790981528288210E-01 good
cost_grad_step_b_step? = 1 20 1.377252112161760306E+04 3.424520237073231943E+00 5.200324046450971593E+01 6.676623448435244912E-01 good
cost_grad_step_b_step? = 1 21 1.377191126199516657E+04 2.570152810776232144E+00 5.140680102179312172E+01 5.632723291196570781E-01 good
cost_grad_step_b_step? = 1 22 1.377157168483655914E+04 2.013346124444937324E+00 5.880772879344541337E+01 6.136475366177717161E-01 good
cost_grad_step_b_step? = 1 23 1.377133330402554202E+04 1.606051135539575236E+00 4.451842783131810677E+01 6.363291982442578210E-01 good
```

The above output shows that GSI is in the inner iteration stage. It may take several minutes to finish the GSI run. Once GSI has finished running, the number of files in the directory will be greatly reduced from those during the run stage. This is because the run script was set to clean the working directory after a successful run. The important analysis result files and configuration files will remain. Please check Section 3.3 for more details on GSI run results. Upon successful completion of GSI, the run directory will look as follows:

```
anavinfo          fort.202 fort.214 fort.228          satbias_ang.out
berror_stats     fort.203 fort.215 fort.229          satbias_in
convinfo         fort.204 fort.217 fort.230          satbias_out
diag_conv_anl.2018081212 fort.205 fort.218 gsi.exe          satbias_out.int
diag_conv_ges.2018081212 fort.206 fort.219 gsiparm.anl       satbias_pc
errtable         fort.207 fort.220 12rwbufr         satbias_pc.out
fit_p1.2018081212 fort.208 fort.221 list_run_directory satinfo
fit_q1.2018081212 fort.209 fort.223 ozinfo           stdout
fit_rad1.2018081212 fort.210 fort.224 pcpbias_out      stdout.anl.2018081212
fit_t1.2018081212 fort.211 fort.225 pcpinfo          wrfanl.2018081212
fit_w1.2018081212 fort.212 fort.226 prepbufr         wrf_inout
fort.201         fort.213 fort.227 prepobs_prep.bufhtable
diag_light_anl.2018081212 diag_light_ges.2018081212
```

5.1.3 Check for Successful GSI Completion

It is important to always check for a successful completion of the GSI analysis. However, completion of the GSI run without crashing does not guarantee a successful analysis. First, it is necessary to check the *stdout* file in the run directory to make sure GSI completed each step without any obvious problems. The following are several important steps to check:

1. Read in the anavinfo and namelist files

The following lines show that GSI started normally and has read in the anavinfo and namelist files:

```
gsi_metguess_mod*init_: 2D-MET STATE VARIABLES:
ps
z
gsi_metguess_mod*init_: 3D-MET STATE VARIABLES:
u
v
... ..
state_vectors*init_anasv: 2D-STATE VARIABLES ps      sst
state_vectors*init_anasv: 3D-STATE VARIABLES u      v      tv      tsen      q      oz      cw      prse
... ..
```

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```

radiance_mode_init: icloud_fwd= F iallsky= F cw_cv= T iaerosol_fwd= F iaerosol= F
radiance_mode_init: n_actual_clouds= 1
radiance_mode_init: cloud_names=cw
... ..

GSI_4DVAR: nobs_bins = 3
SETUP_4DVAR: nobs_bins = 3, ntlevs_ens = 1
SETUP_4DVAR: allocate array containing time levels for ensemble
SETUP_4DVAR: timelevel = 1, ens_fhrlevs = 6
... ..

&SETUP
GENCODE = 78.00000000000000 ,
FACTQMIN = 0.000000000000000E+000,
FACTQMAX = 0.000000000000000E+000,
CLIP_SUPERSATURATION = F,
FACTV = 0.000000000000000 ,
FACTL = 0.000000000000000 ,
FACTP = 0.000000000000000 ,
FACTG = 0.000000000000000 ,
... ..

```

2. Read in the background field

The following lines in standard output file, immediately following the namelist section, indicate that GSI is reading the background fields.

```

... ..
CONVERT_NETCDF_MASS: problem with flnm1 = wrf_inou9, Status = -1021
READ_wrf_mass_FILES: analysis date,minutes 2018 8 12 12 0 21360240
READ_wrf_mass_FILES: sigma guess file, nming2 0.0000000000000000 2018 8 12
READ_wrf_mass_FILES: sigma fcst files used in analysis : 2 1.0000000000000000 1
READ_wrf_mass_FILES: surface fcst files used in analysis: 2 1.0000000000000000 1
GESINFO: Guess date is 12 8 12 2018 0.0000000000000000
... ..

```

3. Read in observational data

Skipping through a majority of the content towards the middle of the standard output file, the following lines will appear:

```

READ_OBS: read 1 ps ps using ntasks= 1 0 1 999999 999999
READ_OBS: read 2 t t using ntasks= 1 1 1 999999 999999
READ_OBS: read 3 q q using ntasks= 1 2 1 999999 999999
READ_OBS: read 4 pw pw using ntasks= 1 0 4218 999999 999999
READ_OBS: read 6 uv uv using ntasks= 1 1 1 999999 999999
READ_OBS: read 10 sst sst using ntasks= 1 2 611 999999 999999

```

This table is an important step to check if the observations have been read in, which types of observations have been read in. At this point, GSI has read in all the data needed for the analysis.

4. Inner iteration

The inner iteration step in the standard output file will look like this:

```

Begin J table inner/outer loop 0 1
J term J
surface pressure 4.7741393535600264E+03
temperature 7.8419025904486407E+03
wind 8.2939160322155458E+03
moisture 4.2982963156969481E+03
sst 4.4435897008983467E+01
-----
J Global 2.5252690188930144E+04
End Jo table inner/outer loop 0 1
Initial cost function = 2.525269018893014800E+04

```

5. GSI Applications

```
Initial gradient norm = 3.940145784968640328E+02
cost_grad_step_b_step? = 1 0 2.525269018893014800E+04 3.940145784968640328E+02 1.978317674865380837E+01 0.0000000000000000E+00 good
cost_grad_step_b_step? = 1 1 2.218140169269516628E+04 3.212948564609200730E+02 2.133681787464458424E+01 6.649407734248591328E-01 good
cost_grad_step_b_step? = 1 2 1.997879377333894445E+04 2.485530349560721959E+02 4.318845368915710736E+01 5.984537528616304947E-01 good
cost_grad_step_b_step? = 1 3 1.731067108515735890E+04 1.990853912639774137E+02 2.912427563070909997E+01 6.415649729561008208E-01 good
cost_grad_step_b_step? = 1 4 1.615633062397510366E+04 1.865345902820838262E+02 2.809840761991527103E+01 8.778897313990374762E-01 good
cost_grad_step_b_step? = 1 5 1.517864222133947260E+04 1.132155175882477351E+02 3.973365751208305596E+01 3.68377520644228529E-01 good
... ..
```

In this case, two outer loops with 50 inner loops in each outer loop are shown. The last iteration looks like this:

```
...
cost_grad_step_b_step? = 2 34 1.363468612559743997E+04 6.120955749102529111E-03 4.469155955050479889E+01 4.112810488623842020E-01 good
cost_grad_step_b_step? = 2 35 1.363468612392302020E+04 5.111121533712363954E-03 3.663810108770889684E+01 6.972573690510027733E-01 good
cost_grad_step_b_step? = 2 36 1.363468612296590072E+04 5.549509485982077710E-03 2.861531574243975129E+01 1.178899912735292821E+00 good
cost_grad_step_b_step? = 2 37 1.363468612208463492E+04 3.684737720368412486E-03 4.055547505420312149E+01 4.408643555780700218E-01 good
PCGSOI: WARNING *** Stopping inner iteration ***
gnorm 0.874557922763519453E-10 less than 0.100000000000000004E-09
update_guess: successfully complete
```

At the 37th iteration in the second outer loop, GSI met the stop threshold before getting to the maximum iteration number (50). As a quick check, the J value should descend with each iteration. Here, J has a value of 2.525269018893014800E+04 at the beginning and a value of 1.363468612208463492E+04 for the final iteration. Therefore, the value has reduced by about one half, which is an expected reduction.

5. Write out analysis results

The final step of the GSI analysis procedure is to write out the Mean, Min and Max values of the analysis fields and the increments:

```
... ..
Status Var          Mean              Min              Max
analysis U         1.361850805619E+00 -2.562917075965E+01 7.369990867998E+01
analysis V         3.886248550541E-02 -3.332195325872E+01 3.974182200227E+01
analysis TV        2.588919513045E+02 1.915694191183E+02 3.103375632340E+02
analysis Q         4.585064517727E-03 1.000000000000E-07 2.222948552069E-02
analysis TSEN      2.580842411077E+02 1.915687083175E+02 3.073688885641E+02
analysis OZ        1.000000000000E-15 1.000000000000E-15 1.000000000000E-15
analysis CW        0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
analysis DIV       0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
analysis VDR       0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
analysis PRSL      5.384764795104E+01 5.322522378726E+00 1.032257938844E+02
analysis PS        9.890316609835E+01 6.892487155270E+01 1.035305116598E+02
analysis SST       2.937426126383E+02 2.656092224121E+02 3.091810302734E+02
analysis radb      9.425591451060E-02 -1.655751310000E+02 2.821827560000E+02
analysis pcpb      0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
analysis aftb      0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
=====
increment u        2.300198794805E-04 -1.798607862276E-01 1.110612583584E-01
increment v        -9.481863951665E-06 -4.211866487153E-01 1.226689231956E-01
increment tv       2.141218696352E-04 -4.154807080575E-01 3.866293159332E-01
increment tsen     2.848999733065E-04 -4.700394309512E-01 3.587168154417E-01
increment q        -4.041678396967E-07 -2.283104749143E-04 6.133666653629E-04
increment oz       0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
increment cw       0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
increment prse     1.186961953910E-04 -2.742236780568E-03 7.248552921752E-03
increment ps       2.284459604308E-04 -2.742236780568E-03 7.248552921752E-03
increment sst      3.549767776646E-04 -6.460545382111E-01 6.877060492842E-01
... ..
```

6. As an indication that GSI has successfully run, several lines will appear at the bottom of the file:

```
glbsoi: complete
[000]gsisub(): : complete.

ENDING DATE-TIME  NOV 01,2018 16:34:33.026 305 THU 2458424
PROGRAM GSI_ANL HAS ENDED.
* . * . * . * . * . * . * . * . * . * . * . * . * . * . * . * .
```


5. GSI Applications

After carefully investigating each portion of the standard output file, it can be concluded that GSI successfully ran through every step and there were no run issues. A more complete description of the standard output file can be found in Section 4.1. However, it cannot be concluded that GSI successfully produced an analysis until more diagnosis has been completed.

5.1.4 Diagnose GSI Analysis Results

a Check Analysis Fit to Observations

The analysis uses observations to correct the background fields to fit to the observations under certain constraints. The easiest way to confirm the GSI analysis results fit the observations better than the background is to check a set of files with names *fort.???*, where ?? is a number from 01 to 19 or larger than 20. In the run script, several "fort" files have also been renamed as *fit_t1(q1, p1, rad1, w1).YYYYMMDDHH*. Please check Section 4.5.1 for a detailed explanation of the fit files. Here, we illustrate how to use these fit files.

- *fit_t1.2018081212 (fort.203)*

This file shows how the background and analysis fields fit to temperature observations. The contents of this file show three data types were assimilated in the analysis: 120, 130, and 180. Also included are the number of observations, bias, and RMS errors of observation minus background (o-g 01) or analysis (o-g 03) on each level for the three data types. The following is part of the file:

```

current fit of temperature data, ranges in K
o-g      ptop 1000.0 900.0 800.0 600.0 400.0 300.0 250.0 200.0 150.0 100.0 50.0 0.0
o-g it   obs use typ styp pbot 1200.0 1000.0 900.0 800.0 600.0 400.0 300.0 250.0 200.0 150.0 100.0 50.0 2000.0
-----
o-g 01   t asm 120 0000 count 123 447 448 820 1169 778 277 433 499 903 363 6260
o-g 01   t asm 120 0000 bias 0.55 0.62 0.39 0.13 0.49 0.69 0.49 0.61 0.97 0.96 1.00 0.61
o-g 01   t asm 120 0000 rms 1.46 1.79 1.62 0.99 1.05 1.27 1.11 1.75 2.46 2.56 2.37 1.73
o-g 01   t asm 120 0000 cpen 0.23 0.69 0.46 0.62 0.73 0.94 0.56 1.25 2.34 2.38 1.62 1.16
o-g 01   t asm 120 0000 qcpen 0.23 0.69 0.46 0.62 0.73 0.94 0.56 1.25 2.34 2.38 1.62 1.16
o-g 01   t asm 130 0000 count 0 0 0 0 0 0 0 1 13 5 0 0 19
o-g 01   t asm 130 0000 bias 0.00 0.00 0.00 0.00 0.00 0.00 -1.02 0.88 0.85 0.00 0.00 0.77
o-g 01   t asm 130 0000 rms 0.00 0.00 0.00 0.00 0.00 0.00 1.02 2.16 1.89 0.00 0.00 2.05
o-g 01   t asm 130 0000 cpen 0.00 0.00 0.00 0.00 0.00 0.00 0.57 2.67 2.06 0.00 0.00 2.40
o-g 01   t asm 130 0000 qcpen 0.00 0.00 0.00 0.00 0.00 0.00 0.57 2.67 2.06 0.00 0.00 2.40
o-g 01   t asm 180 0000 count 300 27 0 0 0 0 0 0 0 0 0 0 327
o-g 01   t asm 180 0000 bias 0.50 2.54 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.67
o-g 01   t asm 180 0000 rms 1.28 3.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 1.50
o-g 01   t asm 180 0000 cpen 0.45 4.13 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.75
o-g 01   t asm 180 0000 qcpen 0.45 4.13 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.75
o-g 01   t asm 180 0001 count 472 7 0 0 0 0 0 0 0 0 0 0 479
o-g 01   t asm 180 0001 bias 0.52 2.98 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.56
o-g 01   t asm 180 0001 rms 1.99 3.20 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 2.01
o-g 01   t asm 180 0001 cpen 0.58 6.02 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.66
o-g 01   t asm 180 0001 qcpen 0.58 6.02 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.66
o-g 01   asm all count 895 481 448 820 1169 778 278 446 504 903 363 7085
o-g 01   asm all bias 0.52 0.76 0.39 0.13 0.49 0.69 0.49 0.62 0.97 0.96 1.00 0.61
o-g 01   asm all rms 1.71 1.90 1.62 0.99 1.05 1.27 1.11 1.76 2.45 2.56 2.37 1.75
o-g 01   asm all cpen 0.49 0.97 0.46 0.62 0.73 0.94 0.56 1.29 2.34 2.38 1.62 1.11
o-g 01   asm all qcpen 0.49 0.97 0.46 0.62 0.73 0.94 0.56 1.29 2.34 2.38 1.62 1.11
... ..
current fit of temperature data, ranges in K
o-g      ptop 1000.0 900.0 800.0 600.0 400.0 300.0 250.0 200.0 150.0 100.0 50.0 0.0
o-g it   obs use typ styp pbot 1200.0 1000.0 900.0 800.0 600.0 400.0 300.0 250.0 200.0 150.0 100.0 2000.0
-----
o-g 03   t asm 120 0000 count 123 447 448 820 1169 778 277 433 499 903 363 6260
o-g 03   t asm 120 0000 bias 0.38 0.35 0.20 0.02 0.17 0.30 0.14 0.12 0.32 0.32 0.41 0.23
o-g 03   t asm 120 0000 rms 1.34 1.47 1.36 0.72 0.64 0.76 0.61 0.95 1.31 1.52 1.66 1.12
o-g 03   t asm 120 0000 cpen 0.18 0.41 0.25 0.32 0.27 0.35 0.18 0.36 0.65 0.85 0.81 0.44
o-g 03   t asm 120 0000 qcpen 0.18 0.41 0.25 0.32 0.27 0.35 0.18 0.36 0.65 0.85 0.81 0.44
o-g 03   t asm 130 0000 count 0 0 0 0 0 0 0 1 13 5 0 0 19
o-g 03   t asm 130 0000 bias 0.00 0.00 0.00 0.00 0.00 0.00 -0.60 0.62 0.58 0.00 0.00 0.54
o-g 03   t asm 130 0000 rms 0.00 0.00 0.00 0.00 0.00 0.00 0.60 1.52 1.50 0.00 0.00 1.48
o-g 03   t asm 130 0000 cpen 0.00 0.00 0.00 0.00 0.00 0.00 0.20 1.32 1.29 0.00 0.00 1.25
o-g 03   t asm 130 0000 qcpen 0.00 0.00 0.00 0.00 0.00 0.00 0.20 1.32 1.29 0.00 0.00 1.25
o-g 03   t asm 180 0000 count 300 27 0 0 0 0 0 0 0 0 0 0 327
o-g 03   t asm 180 0000 bias 0.23 0.77 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.28

```

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o-g 03	t asm 180 0000	rms	0.92	1.55	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.99
o-g 03	t asm 180 0000	cpen	0.14	0.77	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.19
o-g 03	t asm 180 0000	qcpn	0.14	0.77	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.19
o-g 03	t asm 180 0001	count	472	7	0	0	0	0	0	0	0	0	0	0	479
o-g 03	t asm 180 0001	bias	0.18	1.48	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.19
o-g 03	t asm 180 0001	rms	1.68	1.79	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	1.68
o-g 03	t asm 180 0001	cpen	0.22	1.60	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.24
o-g 03	t asm 180 0001	qcpn	0.22	1.60	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.24
o-g 03	asm all	count	895	481	448	820	1169	778	278	446	504	903	363	7085	
o-g 03	asm all	bias	0.22	0.39	0.20	0.02	0.17	0.30	0.13	0.14	0.32	0.32	0.41	0.23	
o-g 03	asm all	rms	1.42	1.48	1.36	0.72	0.64	0.76	0.61	0.97	1.31	1.52	1.66	1.16	
o-g 03	asm all	cpen	0.19	0.45	0.25	0.32	0.27	0.35	0.18	0.39	0.66	0.85	0.81	0.42	
o-g 03	asm all	qcpn	0.19	0.45	0.25	0.32	0.27	0.35	0.18	0.39	0.66	0.85	0.81	0.42	

For example, data type 120 has 1169 observations in layer 400.0-600.0 hPa, a bias of 0.49, and a RMS error of 1.05. The last column shows the statistics for the whole atmosphere. There are several summary lines for all data types, which is indicated by "asm all" in the data types column. For summary O-B (which is "o-g 01" in the file), there are 7085 observations in total, for a bias of 0.61, and a RMS error of 1.75.

Skipping ahead in the "fort.203" file, "o-g 03" columns (under "it") show the observation minus analysis (O-A) information. Under the summary ("all") rows, it can be seen that there were 7085 total observations, a bias of 0.23, and a RMS error of 1.16. This shows that from the background to the analysis, the bias reduced from 0.61 to 0.23, and the RMS error reduced from 1.75 to 1.16. This is about a 62% and 34% reduction respectively, which is a good reduction for the regional analysis.

- *fit_w1.2018081212 (fort.202)*

This file demonstrates how the background and analysis fields fit to wind observations. This file is formatted the same way as *fort.203*. Therefore, only the summary lines for O-B and O-A will be shown here to gain a quick view of the fit to observations:

current vfit of wind data, ranges in m/s																			
o-g	it	obs	use	typ	styp	ptop	1000.0	900.0	800.0	600.0	400.0	300.0	250.0	200.0	150.0	100.0	50.0	0.0	
o-g	it	obs	use	typ	styp	pbot	1200.0	1000.0	900.0	800.0	600.0	400.0	300.0	250.0	200.0	150.0	100.0	50.0	2000.0
o-g 01	asm all	count					830	619	615	1201	705	526	132	257	364	695	372	6316	
o-g 01	asm all	bias					-0.50	0.13	0.50	0.55	0.52	0.25	0.23	1.24	0.76	0.19	0.91	0.35	
o-g 01	asm all	rms					2.85	3.48	3.70	3.46	3.73	3.57	4.41	4.62	4.78	4.55	4.49	3.81	
o-g 03	asm all	count					826	621	616	1202	705	526	132	257	364	695	372	6316	
o-g 03	asm all	bias					0.05	0.18	0.42	0.39	0.41	0.16	0.11	0.58	0.40	0.40	0.79	0.34	
o-g 03	asm all	rms					2.16	2.36	2.08	1.92	2.20	1.87	2.55	2.85	3.41	3.81	4.00	2.61	

O-B: 6316 observations in total, bias is 0.35, and RMS error is 3.81

O-A: 6316 observations in total, bias is 0.34, and RMS error is 2.61

The total bias was not reduced much but the RMS error was reduced from 3.81 to 2.61 (31% reduction).

- *fit_q1.2018081212 (fort.204)*

This file demonstrates how the background and analysis fields fit to moisture observations (relative humidity). The summary lines for O-B and O-A are as follows:

current fit of q data, units in per-cent of guess q-sat																			
o-g	it	obs	use	typ	styp	ptop	1000.0	950.0	900.0	850.0	800.0	700.0	600.0	500.0	400.0	300.0	0.0	0.0	
o-g	it	obs	use	typ	styp	pbot	1200.0	1000.0	950.0	900.0	850.0	800.0	700.0	600.0	500.0	400.0	300.0	0.0	2000.0
o-g 01	asm all	count					497	238	227	262	186	447	373	492	677	757	0	4156	
o-g 01	asm all	bias					7.48	7.40	4.43	1.80	-0.26	-0.36	-1.76	-5.06	-7.64	-9.35	0.00	-2.08	
o-g 01	asm all	rms					11.76	12.18	10.32	12.09	14.13	12.47	14.90	18.43	22.84	24.35	0.00	17.86	
o-g 03	asm all	count					497	238	227	262	186	447	373	492	677	757	0	4156	
o-g 03	asm all	bias					1.29	0.81	1.13	0.37	0.33	-0.06	-0.41	-1.50	-2.99	-6.36	0.00	-1.57	
o-g 03	asm all	rms					5.98	5.14	4.81	7.98	9.41	8.66	10.83	13.50	17.19	19.94	0.00	13.26	

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O-B: 4156 observations in total, bias is -2.08, and RMS error is 17.86

O-A: 4156 observations in total, bias is -1.57, and RMS error is 13.26

The total bias and RMS error were both largely reduced.

- *fit_p1.2018081212 (fort.201)*

This file demonstrates how the background and analysis fields fit to surface pressure observations. Because surface pressure is a two-dimensional field, the table is formatted differently than the three-dimensional fields shown above. Once again, only the summary lines will be shown for O-B and O-A to gain a quick view of the fit to observations:

```
-----
o-g it      obs use typ styp      count      bias      rms      cpen      qcpn
o-g 01      asm all                6251     -0.2037    0.7633    0.7637    0.7637
-----
o-g 03      asm all                6251      0.0150    0.5376    0.3596    0.3596
```

O-B: 6251 observations in total, bias is -0.2037, and RMS error is 0.7633

O-A: 6251 observations in total, bias is 0.0150, and RMS error is 0.5376

Both the total bias and RMS error were reduced.

These statistics show that the analysis results fit to the observations closer than the background, which is what we would expect. How close the analysis fits to the observations is based on the ratio of background error covariance and observation error.

b Check the Minimization

In addition to the minimization information in the standard output file, GSI writes more detailed information into a file called "fort.220." The content of "fort.220" is explained in the Advanced GSI User's Guide. Below is an example of a quick check of the cost function trend and the norm of gradient. The values should get smaller with each iteration.

In the run directory, information on the cost function and norm of the gradient can be dumped into an output file by using the following command:

```
$ grep 'cost,grad,step,b' fort.220 | sed -e 's/cost,grad,step,b,step? = //g' | sed -e 's/good//g' > cost_gradient.txt
```

The output file *cost_gradient.txt* includes six columns, however only the first four columns are needed and are explained below. The first and last five lines are:

```
1  0  2.525269018893014800E+04  3.940145784968640328E+02  1.978317674865380837E+01  0.000000000000000000E+00
1  1  2.218140169269516628E+04  3.212948564609200730E+02  2.133681787464458424E+01  6.6494077342428591328E-01
1  2  1.997879377333894445E+04  2.485530349560721959E+02  4.318845368915710736E+01  5.984537528616304947E-01
1  3  1.731067108515735890E+04  1.990853912639774137E+02  2.912427563070909997E+01  6.415649729561008208E-01
1  4  1.615633062397510366E+04  1.865345902820838262E+02  2.809840761991527103E+01  8.778897313990374762E-01
...
2  33 1.363468612787015809E+04  9.544428453907339432E-03  2.494860880527297908E+01  1.064566883607217962E+00
2  34 1.363468612559743997E+04  6.120955749102529111E-03  4.469155955050479889E+01  4.112810488623842020E-01
2  35 1.363468612392302020E+04  5.111121533712363954E-03  3.663810108770889684E+01  6.972573690510027733E-01
2  36 1.363468612296590072E+04  5.549509485982077710E-03  2.861531574243975129E+01  1.178899912735292821E+00
2  37 1.363468612208463492E+04  3.684737720368412486E-03  4.055547505420312149E+01  4.408643555780700218E-01
```

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The first column is the outer loop number and the second column is the inner iteration number. The third column is the cost function, and the fourth column is the norm of the gradient. It can be seen that both the cost function and norm of the gradient are descending.

To get a complete picture of the minimization process, the cost function and norm of the gradient can be plotted using an included NCL script located here:

```
./util/Analysis_Uutilities/plot_ncl/GSI_cost_gradient.ncl
```

The plot is shown as Fig.5.2:

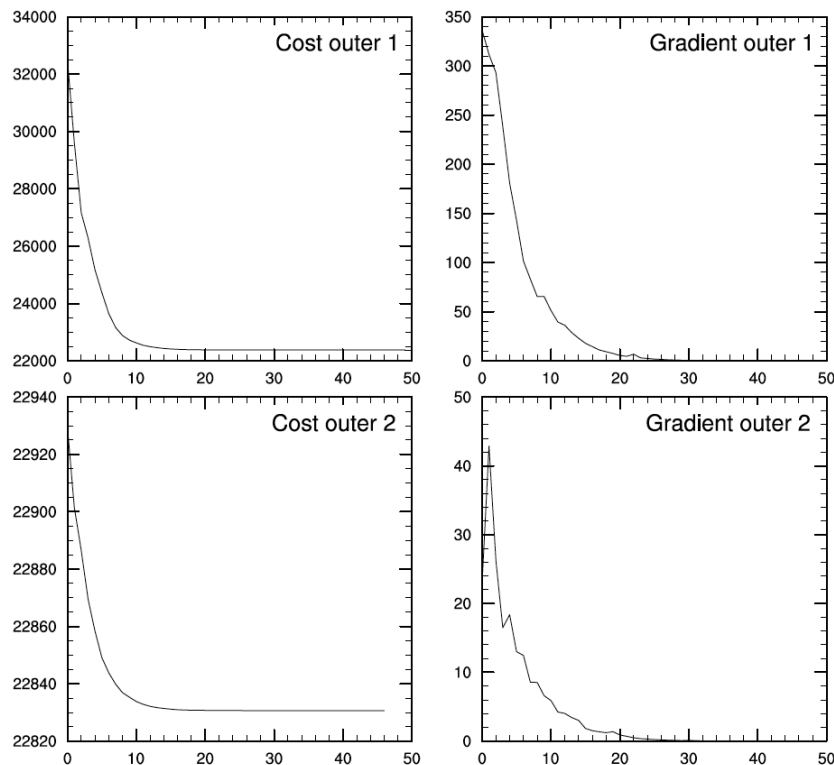


Figure 5.2: The cost function (y-axes) and norm of the gradient (y-axes) change with each iteration (x-axes).

The above plots demonstrate that both the cost function and norm of the gradient descend very fast in the first ten iterations in both outer loops and drop very slowly afterward.

c Check the Analysis Increment

The analysis increment gives us an idea of where and how much the background fields have been modified by the observations through the analysis. Another useful graphics tool that can be used to look at the analysis increment is located here:

```
./util/Analysis_Uutilities/plot_ncl/Analysis_increment.ncl
```

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The graphic below shows the analysis increment at the 2nd vertical level on the analysis grid. Notice that the scales are different for each of the plots.

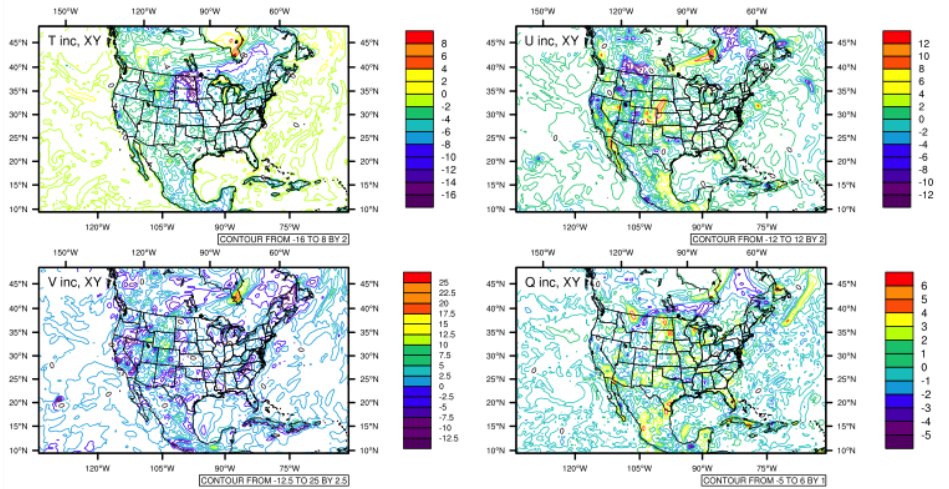


Figure 5.3: Analysis increment at the 2nd level

The analysis increment indicates that conventional observations are mostly located within the continental United States and that data availability over the ocean is very sparse.

5.2 Assimilating Radiance and GPS RO data with Regional GSI

5.2.1 Run Script

Adding radiance and GPS RO data into the GSI analysis is straightforward after having already run GSI with conventional data. The same run script from the above section can be used to run GSI with radiance and GPS RO data (with or without PrepBUFR data). The key step to adding the data is linking the radiance and GPS RO BUFR files to the GSI run directory with the names listed in the `&OBS_INPUT` section of the GSI namelist. The following example adds three radiance BUFR files and the GPS RO file:

```
AMSU-A: gdas1.t12z.1bamua.tm00.bufr_d  
HIRS4: gdas1.t12z.1bhrs4.tm00.bufr_d  
MHS: gdas1.t12z.1bmhs.tm00.bufr_d  
GPS RO: gdas1.t12z.gpsro.tm00.bufr_d
```

Please make sure these BUFR files are located as included in the variable `OBS_ROOT` and link them to the work directory in the script `comgsi_run_regional.ksh`:

```
# Link to the radiance data  
srcobsfile[1]=${OBS_ROOT}/gdas1.t${HH}z.satwnd.tm00.bufr_d  
gsiobsfile[1]=satwnd  
srcobsfile[2]=${OBS_ROOT}/gdas1.t${HH}z.1bamua.tm00.bufr_d  
gsiobsfile[2]=amsuabufr
```

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```
srcobsfile[3]=${OBS_ROOT}/gdas1.t${HH}z.1bhrs4.tm00.bufr_d
gsiobsfile[3]=hirs4bufr
srcobsfile[4]=${OBS_ROOT}/gdas1.t${HH}z.1bmhs.tm00.bufr_d
... ..
ii=1
while [[ $ii -le 21 ]]; do
  if [ -r "${srcobsfile[$ii]}" ]; then
    ln -s ${srcobsfile[$ii]} ${gsiobsfile[$ii]}
    echo "link source obs file ${srcobsfile[$ii]}"
  fi
  (( ii = $ii + 1 ))
done
```

If radiance and RO data are desired in addition to conventional prepBUFR data, the following link to the PrepBUFR data should be kept as is:

```
ln -s ${PREPBUFR} ./prepbufr
```

Alternatively, to analyze radiance and RO data without conventional PrepBUFR data, this line can be commented out in the script *comgsi_run_regional.ksh*:

```
# ln -s ${PREPBUFR} ./prepbufr
```

In the following example, conventional observations will be assimilated, in addition to the radiance and GPS RO.

In order to link the correct name for the radiance BUFR file, the namelist section `&OBS_INPUT` should be referenced. This section has a list of data types and BUFR file names that can be used in GSI. The 1st column "dfile" is the file name recognized by GSI. The 2nd column "dtype" and 3rd column "dplat" are the data type and data platform that are included in the file listed in "dfile," respectively. For example, the following lines tell us the AMSU-A observation from NOAA-15 should be read from a BUFR file named "*amsuabufr*", and the GPS RO refractivity observations "*gps_ref*" should be read from a BUFR file named "*gpsrobufr*":

!	dfile	dtype	dplat	dsis	dval	dthin	dsfcalc
	amsuabufr	amsua	n15	amsua_n15	10.0	2	0
	gpsrobufr	gps_ref	null	gps	1.0	0	0

With radiance data assimilation, data thinning and bias correction need to be checked carefully. The following is a brief description of these two:

- Radiance data thinning

Radiance data thinning is found in the namelist section `&OBS_INPUT`. The following is a part of namelist in that section:

```
dmesh(1)=120.0,dmesh(2)=60.0,dmesh(3)=30,time_window_max=1.5,ext_sonde=.true.
! dfile      dtype      dplat      dsis      dval      dthin dsfcalc
  amsuabufr  amsua      n15        amsua_n15 10.0      2      0
```

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The first line of `&OBS_INPUT` lists multiple mesh grids as elements of the array `dmesh` (three mesh grids in the above example). For the line specifying data type, the 2nd to last element of that line is used to specify the choice of `dthin`. This selects the mesh grid to be used for thinning. The data thinning option for NOAA-15 AMSU-A observations is set to 60 km because the value of `dthin` is two, corresponding to `dmesh(2)=60` km. For more information about radiance data thinning, please refer to the Advanced GSI User's Guide.

- Radiance data bias correction

Radiance data bias correction is very important for successful radiance data assimilation. In the sample run scripts, there are two files related to bias correction:

```
# for satellite bias correction
# Users may need to use their own satbias files for correct bias correction
cp ${GSI_ROOT}/fix/comgsi_satbias_in ./satbias_in
cp ${GSI_ROOT}/fix/comgsi_satbias_pc_in ./satbias_pc_in
```

For this case, the GDAS bias correction files were downloaded and saved in the `fix` directory as examples. For other cases, the run script should link to corresponding bias correction coefficient files. The first line sets the path to the bias coefficient file, and the second copies the bias correction coefficients for passive (monitored) channels into the working directory. These two coefficient files are usually calculated from within GSI in the previous cycle. Two files are provided in `./fix` as examples of the bias correction coefficients. For the best results, it is necessary for the user to generate his or her own bias files. The details of radiance data bias correction are discussed in the Advanced GSI User's Guide. Please note that GSI releases prior to v3.5 have coefficients for mass bias correction and angle bias correction calculated separately.

Once these links are set, we are ready to run GSI.

5.2.2 Run GSI and Check Run Status

The process for running GSI is the same as described in section 5.1.2. Once `comgsi_run_regional.ksh` has been submitted, move into the run directory to check the GSI analysis results. For the current case, the run directory will look almost as it did for the conventional data case, the exception being the links to the radiance and GPS RO BUFR files and new diag files for the radiance data types used. Following the same steps as in section 5.1.2, check the `stdout` file to see if GSI has run through each part of the analysis process successfully. In addition to the information outlined for the conventional run, the radiance BUFR files should have been read in, as well as the refractivity data from GPS RO:

```
READ_OBS: read 33 mhs      mhs_n18      using ntasks= 2 0 2123 999999 999999
READ_OBS: read 34 mhs      mhs_n19      using ntasks= 2 2 159 999999 999999
READ_OBS: read 35 mhs      mhs_metop-a  using ntasks= 2 0 590 999999 999999
READ_OBS: read 36 mhs      mhs_metop-b  using ntasks= 2 2 3 999999 999999
READ_OBS: read 1 ps        ps           using ntasks= 1 0 1 999999 999999
READ_OBS: read 2 t         t            using ntasks= 1 1 1 999999 999999
READ_OBS: read 3 q         q            using ntasks= 1 2 1 999999 999999
READ_OBS: read 4 pw        pw           using ntasks= 1 0 4218 999999 999999
READ_OBS: read 6 uv        uv           using ntasks= 1 1 1 999999 999999
READ_OBS: read 9 rw        rw           using ntasks= 1 2 1 999999 999999
```

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```

READ_OBS: read 10 sst      sst      using ntasks= 1 0 611 999999 999999
READ_OBS: read 11 gps_ref  gps      using ntasks= 1 1 1 999999 999999
READ_OBS: read 19 hirs4    hirs4_metop-a using ntasks= 1 2 292 999999 999999
READ_OBS: read 21 hirs4    hirs4_n19     using ntasks= 1 0 80 999999 999999
READ_OBS: read 22 hirs4    hirs4_metop-b using ntasks= 1 1 3 999999 999999
READ_OBS: read 26 amsua    amsua_n15     using ntasks= 1 2 72 999999 999999
READ_OBS: read 27 amsua    amsua_n18     using ntasks= 1 0 509 999999 999999
READ_OBS: read 28 amsua    amsua_n19     using ntasks= 1 1 31 999999 999999
READ_OBS: read 29 amsua    amsua_metop-a using ntasks= 1 2 159 999999 999999
READ_OBS: read 30 amsua    amsua_metop-b using ntasks= 1 0 3 999999 999999

```

When comparing this output to the content in step three of section 5.1.3, it can be seen that, in addition to the GPS refractivity observations (*gps_ref*), there are twelve new radiance data types that have been read in: HIRS4 from METOP-A, METOP-B and NOAA-19, AMSU-A from NOAA-15, NOAA-18, NOAA-19, METOP-A, and METOP-B, and MHS from NOAA-18, NOAA-19, METOP-A, and METOP-B.

5.2.3 Diagnose GSI Analysis Results

a Check Analysis Fit to Observations

The file *fort.207* contains the statistics for the radiance data, similar to file *fort.203* for temperature. This file contains important details about the radiance data analysis. Section 4.5.2 explains this file in detail. Below are some values from the file *fort.207* to provide a quick look at the radiance assimilation for this example.

The *fort.207* file contains the following lines:

For O-B, before the first outer loop:

```

      it  satellite instrument  # read  # keep  # assim  penalty  qcplty  cpen  qccpen
o-g 01 rad  n15    amsua    32535  22797  3193  3593.4  3593.4  1.1254  1.1254
o-g 01 rad  n18    amsua    33465  27762  9127  2973.1  2973.1  0.32575  0.32575

```

For O-A, after the second outer loop:

```

o-g 03 rad  n15    amsua    32535  22797  6006  2020.2  2020.2  0.33636  0.33636
o-g 03 rad  n18    amsua    33465  27762  10483  1386.9  1386.9  0.13230  0.13230

```

From the above information, it can be seen that AMSU-A data from NOAA-15 provides 32535 observations within the analysis time window and domain. After thinning, 22797 observations remained, and only 3193 passed the quality check and were used in the analysis. The penalty for this data decreased from 3593.4 to 2020.2 after two outer loops. It is important to note that the number of AMSU-A observations assimilated in the O-A calculation increased to 6006 from 3193 as more data passed the quality check in the 2nd outer loop.

The statistics for each channel can be viewed in the *fort.207* file as well. Here, channels from AMSU-A NOAA-15 are listed as an example:

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For O-B, before the first outer loop:

1	1	amsua_n15	190	0	3.000	2.9421171	0.4438706	0.1561571	1.5278684	1.4619715
2	2	amsua_n15	190	0	2.200	0.8933773	0.0750037	0.1639086	1.3984509	1.3964381
3	3	amsua_n15	190	0	2.000	1.2106089	-0.0473438	0.1661452	1.0258870	1.0247940
4	4	amsua_n15	190	0	0.600	-0.4461829	-0.5369729	0.7065383	0.6161111	0.3020812
5	5	amsua_n15	190	0	0.300	-0.0791235	-0.0665128	0.3446176	0.2013947	0.1900944
6	6	amsua_n15	292	342	-0.230	-2.4084362	0.5222298	4.1440417	0.5364275	0.1225986
7	7	amsua_n15	1320	435	0.250	0.0772148	0.2545220	1.1522083	0.3141868	0.1842060
8	8	amsua_n15	733	1022	0.275	0.2375745	0.4957631	2.3953990	0.5205394	0.1586827
15	15	amsua_n15	190	0	3.500	0.3668241	0.3548074	0.1291879	1.4624421	1.4187490

For O-A, after the second outer loop:

1	1	amsua_n15	361	0	3.000	2.3526577	-0.0691773	0.0570915	0.9734396	0.9709784
2	2	amsua_n15	361	0	2.200	1.0975100	0.2032144	0.0916095	1.0612608	1.0416229
3	3	amsua_n15	361	0	2.000	1.4728299	0.1679623	0.0870602	0.7443155	0.7251168
4	4	amsua_n15	361	0	0.600	-0.2468814	-0.3416600	0.3559445	0.4243844	0.2517351
5	5	amsua_n15	361	0	0.300	0.0601439	0.0652633	0.2859038	0.1824454	0.1703732
6	6	amsua_n15	571	289	-0.230	-2.4451832	0.4898659	3.7588334	0.5051936	0.1234989
7	7	amsua_n15	1560	195	0.250	-0.1917289	0.0354134	0.4463849	0.1931275	0.1898529
8	8	amsua_n15	1512	243	0.275	-0.1662265	0.0570724	0.3423785	0.1943532	0.1857846
9	9	amsua_n15	768	987	0.340	0.1799446	0.3241856	0.6087271	0.3750867	0.1886630
15	15	amsua_n15	361	0	3.500	0.3218240	0.4970433	0.0604030	1.0051883	0.8736998

The second column is the channel number for AMSU-A and the last column is the standard deviation for each channel. It can be seen that most of the channels fit better to the observations after the second outer loop.

The file *fort.212* shows the fit of the analysis/background to the GPS RO data as fractional differences. It has the same structure as the fit files for conventional data. Below is a quick look to be sure the GPS RO data were used:

```

Observation - Background (O-B)
current fit of gps data in fractional difference
o-g      ptop 1000.0 900.0 800.0 600.0 400.0 300.0 250.0 200.0 150.0 100.0 50.0 0.0
o-g it   obs use typ styp pbot 1200.0 1000.0 900.0 800.0 600.0 400.0 300.0 250.0 200.0 150.0 100.0 50.0 2000.0
-----
o-g 01   asm all count      0      0      1      33      85      79      47      49      62      89      108      553
o-g 01   asm all bias      0.00  0.00  0.35  0.33 -0.57 -0.22 -0.22 -0.05 -0.34  0.10  0.31 -0.08
o-g 01   asm all rms      0.00  0.00  0.35  1.12  1.32  0.57  0.51  0.73  0.78  0.39  0.67  0.80

Observation - Analysis (O-A)
      it   obs   type styp  ptop 1000.0 900.0 800.0 600.0 400.0 300.0 250.0 200.0 150.0 100.0 50.0 0.0
-----
o-g 03   asm all count      0      0      1      33      85      80      50      55      66      88      110      568
o-g 03   asm all bias      0.00  0.00  0.07  0.28  0.03 -0.06 -0.08  0.09 -0.10  0.01  0.05  0.01
o-g 03   asm all rms      0.00  0.00  0.07  0.77  0.50  0.30  0.24  0.32  0.36  0.23  0.28  0.37

```

It can be seen that most of the GPS RO data are located in the upper levels, with a total of 553 observations used in the analysis during the 1st outer loop, and 568 used to calculate O-A. After the analysis, the data bias reduced from -0.08 to 0.01, and the RMS error was reduced from 0.80 to 0.37. It can be concluded that the analysis with GPS RO data looks reasonable from these statistics.

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b Check the Analysis Increment

The same methods for checking the optimal minimization as demonstrated in section 5.1.4.2 can be used for this case. Similar features to the conventional assimilation should be seen with the minimization. The figures below show detailed information on how the radiance and GPS RO data impact the analysis results on top of the conventional data. Using the same NCL script as in section 5.1.4.3, analysis increment fields are plotted comparing the analysis results with radiance, GPS RO and conventional data, to the analysis results with conventional data assimilation only. Figure 5.5 is for vertical level 21 and Figure 5.4 is for vertical level 2, respectively.

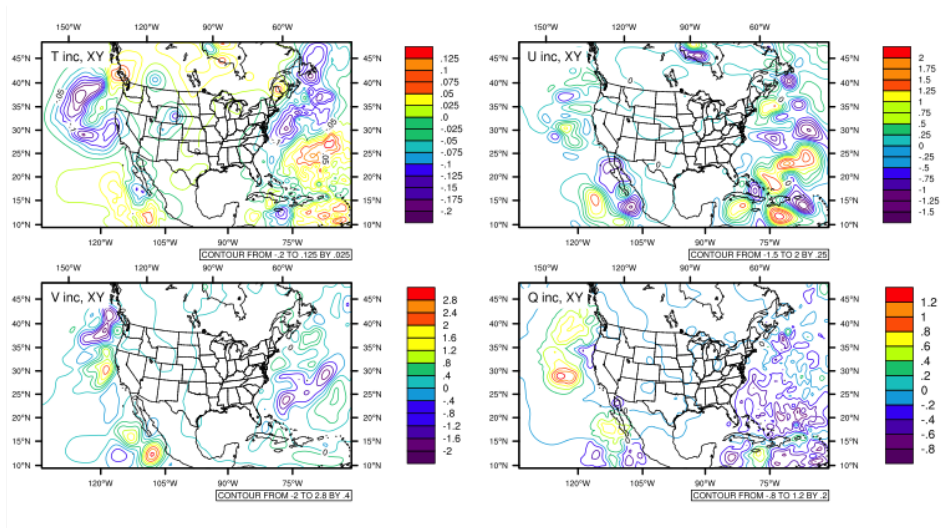


Figure 5.4: Analysis increment fields of the PrepBUFR, GPS RO and radiance data analysis compared to the analysis with prepBUFR only at vertical level 2

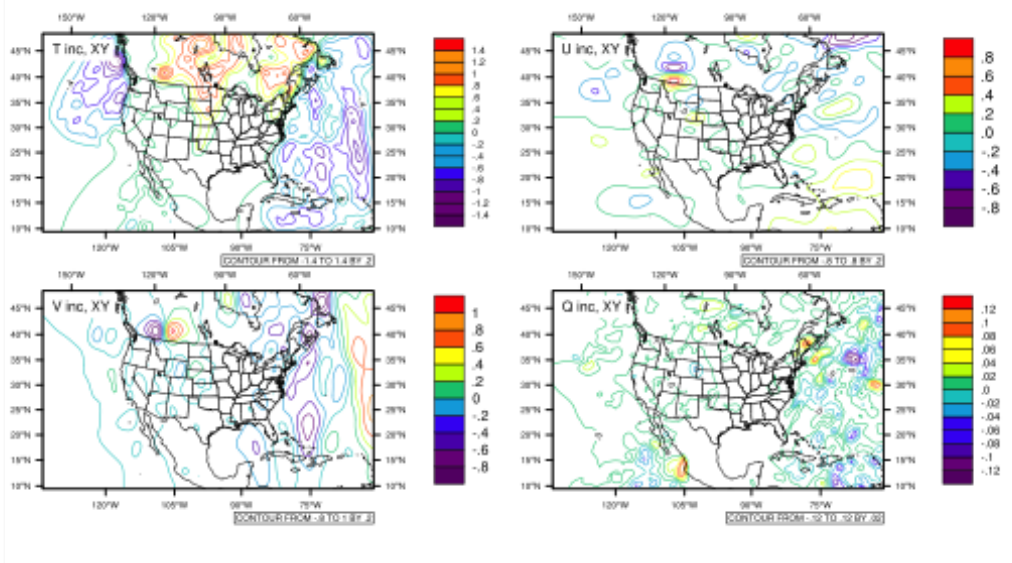


Figure 5.5: Analysis increment fields of the PrepBUFR, GPS RO and radiance data analysis compared to the analysis with prepBUFR only at vertical level 21

In order to fully understand the analysis results, the following topics should be reviewed:

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1. The weighting functions of each channel and the data coverage at the analysis time. There are several sources on the internet that show the weighting functions of the AMSU-A channels, as an example. Channel 1 is the moisture channel, while the others are mainly temperature channels (Channels 2, 3, and 15 also have large moisture signals). Because a model top of 50 mb was specified for this case study, the actual impact should come from channels with peak weighting below 50 hPa.
2. The usage of each channel is located in the file named *'satinfo'* in the run directory. The first two columns show the observation type and platform of the channels, and the third column indicates if the channel is used in the analysis. Because many amsua_n15 and amsua_n18 data were used, they should be checked in detail. In this case, Channels 6, 11, and 14 from amsua_n15 and channels 5, 8, 9 and 14 from amsua_n18 were turned off.
3. Thinning information, including a quick look at the namelist in the run directory. The file "gsparm.anl" shows that both amsua_n15 and amsu_n18 use thinning grid two, which is 60 km.
4. Bias correction: Radiance bias correction was previously discussed. It is very important for a successful radiance data analysis. The run script can only link to the GDAS bias correction coefficients that are provided as an example in *./fix*:

```
cp ${GSI_ROOT}/fix/comgsi_satbias_in ./satbias_in
cp ${GSI_ROOT}/fix/comgsi_satbias_pc_in ./satbias_pc_in
```

Users can download the operational bias correction coefficients during their experiment period as a starting point to calculate the coefficients suitable for their experiments.

Radiance bias correction for regional analyses is a difficult issue because of the limited coverage of radiance data. This topic is out of the scope of this document, but this issue should be considered and understood when using GSI with radiance applications.

5.3 GSI Hybrid EnVar Analysis

5.3.1 Introduction to GSI Hybrid 3DEnVar Analysis

The three-dimensional hybrid ensemble-variational (hybrid 3DEnVar) analysis is an important option in the GSI system that has been used operationally. It provides the ability to bring the flow dependent background error covariance into the analysis based on ensemble forecasts. If ensemble forecasts have been generated, setting up GSI to do a hybrid analysis is straightforward and only requires two changes in the run script in addition to the current regional run script `comgsi_run_regional.ksh`:

- *Change 1: Specify the location of the ensemble members*

In the run script, point to the directory that contains the ensemble files:

```
ENS_ROOT=case_data/2018081212/gfsens
```

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This step is required to link the ensemble members to the GSI run directory and assign each ensemble member a name that GSI recognizes. GSI can accept four kinds of ensemble forecasts, controlled by the namelist variable *regional_ensemble_option*. Table 5.1 provides a list of options for *regional_ensemble_option* and the naming convention for linking the ensemble files to GSI recognized names.

Table 5.1: List of ensemble forecasts that can be read by GSI

regional_ensemble_option	explanation	Function called	GSI recognized ensemble file names
1	GFS ensemble internally interpolated to hybrid grid	get_gefs_for_regional	<i>filelist</i> : a text file including the path and name of ensemble files
2	Ensemble is in WRF-NMM (HWRF) format	get_wrf_nmm_enspers	<i>d01_en001</i> , <i>d01_en002</i> , ...
3	Ensemble is in ARW netcdf format	get_wrf_mass_enspers_netcdf	<i>wrf_en001</i> , <i>wrf_en002</i> , ...
4	Ensemble is in NMMB format	get_nmmb_enspers	<i>nmmb_ens_mem001</i> , <i>nmmb_ens_mem002</i> , ...

The example script assumes that the hybrid 3DEnVar uses GFS ensemble files. If this is the case, the script will link proper files from the directory ENS_ROOT to the run directory. Please note that starting from this release version, the GFS ensemble has a new format using NEMS IO. If you are using GFS ensemble files like `gdas.t{HH}z.atmf006s.mem005.nemsio`, please set it up in the run script:

```
if_nemsio=Yes
```

If using non-GFS ensemble files, users have to change the GSI run script to add the links to the ensemble forecasts if they want to use the GSI hybrid function. Below is an example of using an ensemble in ARW netcdf format, assuming that all the ensemble members are located in a directory defined by the parameter *\$mempath* and the ensemble members have a name such as: *wrfout_d01_\${iimem}*, where *\$iimem* is an integer indicating the ensemble member ID. The following lines should be added to the run script with loop *iimem* from one to the total number of ensemble members:

```
if [ -r ${mempath}/wrfout_d01_${iimem} ]; then
  ln -sf ${mempath}/wrfout_d01_${iimem} ./wrf_en${iimem}
else
  echo "member ${mempath}/wrfout_d01_${iimem} does not exist"
fi
```

- *Change 2: Set up the namelist options in section HYBRID_ENSEMBLE*

In the run script `comgsi_run_regional`, set it up like this for the hybrid 3DEnVar analysis:

```
if_hybrid=Yes # Yes, or, No -- case sensitive !
if_4DEnVar=No # Yes, or, No -- case sensitive (set if_hybrid=Yes first)!
```

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Table 5.2: The list of namelist options for GSI hybrid

Options	explanation
<code>l_hyb_ens</code>	if true, turn on hybrid ensemble option
<code>uv_hyb_ens</code>	if true, ensemble perturbation wind variables are u and v; otherwise, ensemble perturbation wind variables are stream function and velocity potential
<code>generate_ens</code>	if true, generate an internal ensemble based on the existing background error; recommended = false
<code>n_ens</code>	number of ensemble members
<code>beta_s0</code>	value between 0 and 1, default = 1.0 relative weight given to static background B when (<code>readin_beta=.false.</code>). when (<code>readin_beta=.true.</code>), the verti- cal weighting parameters are read from a file, instead of being defined based on <code>beta_s0</code> namelist or default value. =1, ensemble information turned off =0, static background errors turned off
<code>s_ens_h</code>	homogeneous isotropic horizontal ensemble localization scale (km)
<code>s_ens_v</code>	vertical localization scale If positive, in grid units; if negative, in lnp unit
<code>regional_ensemble_</code> <code>_option</code>	integer, used to select the type of ensemble to read in for re- gional applications. Currently takes values from one to four: =1: use GEFS internally interpolated to ensemble grid; =2: ensembles are in WRF-NMM format; =3: ensembles are in ARW netcdf format; =4: ensembles are in NMMB format.
<code>grid_ratio_ens</code>	for regional runs, the ratio of ensemble to analysis grid resolu- tion. If turned on and specified with an appropriate value, this could increase the computational efficiency.
<code>l_ens_in_diff_time</code>	if use ensembles that are available at different time from analysis time. =false: only ensembles available at analysis time can be used for hybrid. (default); =true: ensembles available time can be different from analysis time in hybrid analysis.

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This will set `l_hyb_ens=.true.` to turn on the hybrid ensemble analysis. Commonly used namelist options for the hybrid analysis are listed in table 5.2:

Please note: the parameters `s_ens_h`, `s_ens_v`, and `beta1_inv` are tunable parameters. They should be tuned for best performance.

After setting up the namelist parameters and the path/name of the ensemble members, GSI can be run in the same manner as the other 3DVAR cases introduced in this chapter. The same procedures could be followed as in the previous sections to check the run status and diagnose the GSI analysis.

5.3.2 Introduction to GSI Hybrid 4DEnVar Analysis

The GSI hybrid 4DEnVar analysis is similar to the hybrid 3DEnVar except that the hybrid 4DEnVar will use multiple background files and GFS ensemble forecasts. As an example, the following shows how to conduct a hybrid 4DEnVar analysis using three time levels of background files and ensembles.

Before creating a hybrid 4DEnVar analysis, be sure to read the previous section about how to run hybrid 3DEnVar first. The following steps are additional procedures beyond hybrid 3DEnVar and assume that all hybrid 3DEnVar settings have already been set.

- (1). Set `if_4DEnVar=Yes` in `comgsi_run_regional.ksh`, in addition to `if_hybrid=Yes`.
- (2). Set the correct background files and ensemble files at different time levels in `comgsi_run_regional.ksh`. See the following example:

```
if [ ${if_4DEnVar} = Yes ] ; then
  BK_FILE_P1=${BK_ROOT}/wrfout_d01_${datep1}
  BK_FILE_M1=${BK_ROOT}/wrfout_d01_${datem1}

  if [ ${if_nemsio} = Yes ]; then
    ENSEMBLE_FILE_mem_p1=${ENS_ROOT}/gdas.t${gHH}z.atmf009s.mem
    ENSEMBLE_FILE_mem_m1=${ENS_ROOT}/gdas.t${gHH}z.atmf003s.mem
  else
    ENSEMBLE_FILE_mem_p1=${ENS_ROOT}/sfg_${gdate}_fhr09s_mem
    ENSEMBLE_FILE_mem_m1=${ENS_ROOT}/sfg_${gdate}_fhr03s_mem
  fi
fi
```

Note that the background file at the analysis time (2018081218 for the above example) is set by `BK_FILE` and the ensemble files at the analysis time are set by `ENSEMBLE_FILE_mem` as introduced in previous sections.

Now GSI can be run following the hybrid 3DEnVar case introduced in the previous section. Similar procedures can be conducted to check the GSI run status and results.

5.4 Introduction to Global GSI analysis

The *Global Forecast System (GFS)* is a global numerical weather prediction system containing a global computer model and variational analysis run by the U.S. National Weather Service (NWS). As of February 2015, the numerical model is run four times a day, and produces forecasts for up to 16 days in advance, with decreased spatial resolution after 10 days. It is a spectral model with a resolution of T1534 from 0 to 240 hours (0-10 days) and T574 from 240 to 384 hours (10-16 days). In the vertical, the model is divided into 64 layers and temporally, it produces forecast output every hour for the first 12 hours, every 3 hours out to 10 days, and every 12 hours after that. Its data assimilation system runs 6-hourly continuous cycles using the GSI-hybrid.

GSI has many functions specifically designed and tuned for GFS. Although the release version of the community GSI includes all the functions used by the operational systems, the DTC can only support the GSI regional applications because the DTC is not able to run GFS on community computers. Beginning with release version 3.2, the DTC began to introduce the use of GSI for global applications, assuming users can obtain the GFS background through the NCEP data hub or by running GFS themselves.

5.4.1 The Difference between Global and Regional GSI

As mentioned above, all of the NCEP operational systems use GSI as their analysis system. The majority of the GSI code is common to these operational systems. Very little source code is specific to a particular operational system. The main differences in the GSI operational application come from the configuration the run scripts and namelist parameters.

The different GSI applications need different backgrounds, observations, and fixed files. For the GFS system, GSI needs:

- **GFS Backgrounds:** Typically, GSI uses 6-h GFS forecasts as the background. GFS 3-h and 9-h forecasts are also needed for the FGAT function in the GSI analysis. Both surface and atmosphere forecasts are needed.
- **Observations:** NCEP has several sets of BUFR/prepBUFR observation files with global coverage for global systems. The files that start with the prefix **GDAS** are for the 6-hourly global data assimilation system. These files have more data available for the analysis, but have a longer delay for use in real-time. The files that start with **gfs** are for the GFS forecasts. Different operational systems need different observation data files because they require different kinds of observations with different coverage, cut-off times, and quality control processes. All these observation files are read in and processed in GSI by the same section of code. Therefore, there is no problem using GFS observation data files for regional GSI applications, as is described in the practice cases and the GSI User's Guide. Using regional BUFR files for global applications will cause problems since the data only cover part of the analysis domain, but GSI can still read in the observations and perform the analysis.
- **Fixed files:** Section 3.1 of the GSI User's Guide introduced the notion that different operational systems have their own fixed files. These global fixed

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files can be downloaded as a separate tar ball from the GSI user's website (<http://www.dtcenter.org/com-GSI/users/downloads/index.php>). For the GFS GSI application, the big difference is the background error covariance (BE). Different resolutions of the GFS backgrounds use their matched BE files, which are different from the BE files used by the regional GSI applications. In this release version, 14 BE files were provided for users:

1. global_berror.l63y386.f77
2. global_berror.l64y192.f77
3. global_berror.l64y290.f77
4. global_berror.l64y674.f77
5. global_berror.l64y96.f77
6. global_berror.l64y1154.f77
7. global_berror.l64y194.f77
8. global_berror.l64y386.f77
9. global_berror.l64y770.f77
10. global_berror.l64y98.f77
11. global_berror.l64y130.f77
12. global_berror.l64y258.f77
13. global_berror.l64y578.f77
14. global_berror.l64y882.f77

5.4.2 Global GFS Scripts

Starting with release version 3.3, support for running GSI global applications was added to the community release. Specifically a sample global analysis run script was added to the `./ush` directory named `comgsi_run_gfs.ksh` of the community release. This script `comgsi_run_gfs.ksh` is based on the GSI GFS regression tests. This script retains the same basic structure as the regional run script `comgsi_run_regional.ksh`, but includes additional details needed for the global analysis.

The first part of the global analysis run script, just as in the regional script, sets up the computer environment and case configuration. The primary differences between the global and regional scripts are the specification of the GFS case and the global application namelist.

```
GFSCASE=T62
GSI_NAMELIST=${GSI_ROOT}/ush/comgsi_namelist_gfs.sh
```

While the regional script simply specifies the background and BE files, the global script needs to know the background resolution by defining the following parameters:

```
# Set the JCAP resolution which you want.
# All resolutions use LEVS=64
if [[ "$GFSCASE" = "T62" ]]; then
    JCAP=62
    JCAP_B=62
elif [[ "$GFSCASE" = "T126" ]]; then
    JCAP=126
```


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```
JCAP_B=126
elif [[ "$GFSCASE" = "enkf_glb_t254" ]]; then
  JCAP=254
  JCAP_B=254
elif [[ "$GFSCASE" = "T254" ]]; then
  JCAP=254
  JCAP_B=574
elif [[ "$GFSCASE" = "T574" ]]; then
  JCAP=574
  JCAP_B=1534
else
  echo "INVALID case = $GFSCASE"
  exit
fi
LEVS=64
#
```

Just as with the regional analysis run script, the global script double checks the needed parameters, creates a run directory, and copies the background, observations, and fixed files into the run directory. It generates the namelist, and places it in the run directory as well.

1. Specify the values of LATA, LONA, DELTIME, resol based on the choice of JCAP:

```
# Given the requested resolution, set dependent resolution parameters
if [[ "$JCAP" = "382" ]]; then
  LONA=768
  LATA=384
  DELTIM=180
  resol=1
elif [[ "$JCAP" = "574" ]]; then
  LONA=1152
  LATA=576
  DELTIM=1200
  resol=2
elif [[ "$JCAP" = "254" ]]; then
  LONA=512
  LATA=256
  DELTIM=1200
  resol=2
elif [[ "$JCAP" = "126" ]]; then
  LONA=256
  LATA=128
  DELTIM=1200
  resol=2
elif [[ "$JCAP" = "62" ]]; then
  LONA=192
  LATA=94
  DELTIM=1200
  resol=2
else
  echo "INVALID JCAP = $JCAP"
  exit
fi
NLAT=' expr $LATA + 2 '
```

2. Set up CO₂, CH₄, N₂O, CO file decisions:

```
IC02=${IC02:-0}
if [ $IC02 -gt 0 ] ; then
  # Copy co2 files to $workdir
  co2dir=${FIX_ROOT}
  yyyy='echo $ANAL_TIME | cut -c1-4'
  rm ./global_co2_data.txt
  co2=$co2dir/global_co2.gcmscl_$yyyy.txt
  while [ ! -s $co2 ] ; do
    ((yyyy-=1))
    co2=$co2dir/global_co2.gcmscl_$yyyy.txt
  done
  if [ -s $co2 ] ; then
    cp $co2 ./global_co2_data.txt
```

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```
fi
if [ ! -s ./global_co2_data.txt ] ; then
    echo "\./global_co2_data.txt" not created
    exit 1
fi
#CH4 file decision
...
```

3. Set up the namelist parameters and generate the namelist:

```
# Set some parameters for use by the GSI executable and to build the namelist
echo " Build the namelist "

vs_op='0.7,'
hzscl_op='1.7,0.8,0.5,'

... ..

# Build the GSI namelist on-the-fly
. $GSI_NAMELIST
```

4. Multiple time level backgrounds are needed:

```
if [[ "$GFSCASE" = "enkf_glb_t62" ]]; then
    cp $BK_ROOT/bfg_${gdate}_fhr03_ensmean ./sfcf03
    cp $BK_ROOT/bfg_${gdate}_fhr06_ensmean ./sfcf06
    cp $BK_ROOT/bfg_${gdate}_fhr09_ensmean ./sfcf09

    cp $BK_ROOT/sfg_${gdate}_fhr03_mem001 ./sigf03
    cp $BK_ROOT/sfg_${gdate}_fhr06_mem001 ./sigf06
    cp $BK_ROOT/sfg_${gdate}_fhr09_mem001 ./sigf09
else
    cp $BK_ROOT/sfcf03 ./sfcf03
    cp $BK_ROOT/sfcf06 ./sfcf06
    cp $BK_ROOT/sfcf09 ./sfcf09

    cp $BK_ROOT/sigf03 ./sigf03
    cp $BK_ROOT/sigf06 ./sigf06
    cp $BK_ROOT/sigf09 ./sigf09
fi
```

Both surface and atmosphere files at 03, 06, and 09 hour forecasts are needed.

5. Link observations files:

```
# Link to the prepbufr data
ln -s ${PREPBUFR} ./prepbufr

# Link to the other observation data
if [ -r "${OBS_ROOT}/satwnd" ]; then
    ln -s ${OBS_ROOT}/satwnd .
fi
if [ -r "${OBS_ROOT}/gpsrobufr" ]; then
    ln -s ${OBS_ROOT}/gpsrobufr .
fi
if [ -r "${OBS_ROOT}/ssmirrbufr" ]; then
    ln -s ${OBS_ROOT}/ssmirrbufr .
fi
... ..
```

Table 5.3 lists the grid dimensions for GFS of different resolutions from *Running Global Model Parallel Experiments Version 7.0* from NCEP/EMC, which may provide users more information on the above mentioned resolution parameters.

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Table 5.3: The grid dimensions for GFS.

SPECTRAL RESOLUTION	EULERIAN		SEMI-LAGRANGIAN	
	LONB	LATB	LONB	LATB
T62	192	94	128	64
T126	384	190	256	128
T170	512	256	352	176
T190	576	288	384	192
T254	768	384	512	256
T382	1152	576	768	384
T574	1760	880	1152	576
T878	2304	1152	1760	880
T1148			2304	1152
T1534			3072	1536
T2014			4032	2016
T2046			4096	2048
T3070			6144	3072

5.4.3 Sample Results

After a successful run of the GSI GFS analysis, the contents of the run directory, with the clean option turned on, will look something like this:

```

airsbufr
amsuabufr
amsuabufrears
amsuabufrears
anavinfo
atms_beamwidth.txt
atmsbufr
berror_stats
bftab_sstphr
cloudy_radiance_info.txt
convinfo
crisbufr
diag_airs_aqua_anl.2014080400
diag_airs_aqua_ges.2014080400
diag_amsua_aqua_anl.2014080400
diag_amsua_aqua_ges.2014080400
diag_amsua_metop-a_anl.2014080400
diag_amsua_metop-a_ges.2014080400
diag_amsua_metop-b_anl.2014080400
diag_amsua_metop-b_ges.2014080400
diag_amsua_n15_anl.2014080400
diag_amsua_n15_ges.2014080400
diag_amsua_n18_anl.2014080400
diag_amsua_n18_ges.2014080400
diag_amsua_n19_anl.2014080400
diag_amsua_n19_ges.2014080400
diag_atms_npp_anl.2014080400
diag_atms_npp_ges.2014080400
diag_conv_anl.2014080400
diag_conv_ges.2014080400
diag_gome_metop-a_anl.2014080400
diag_gome_metop-a_ges.2014080400
diag_gome_metop-b_anl.2014080400
diag_gome_metop-b_ges.2014080400
diag_hirs4_metop-a_anl.2014080400
diag_hirs4_metop-b_ges.2014080400
diag_hirs4_n19_anl.2014080400
diag_hirs4_n19_ges.2014080400
diag_iasi_metop-a_anl.2014080400
diag_iasi_metop-a_ges.2014080400
diag_iasi_metop-b_anl.2014080400
diag_iasi_metop-b_ges.2014080400
diag_light_anl.2014080400
diag_light_ges.2014080400
diag_mhs_metop-a_anl.2014080400
diag_mhs_metop-a_ges.2014080400
diag_mhs_metop-b_anl.2014080400
diag_mhs_metop-b_ges.2014080400
diag_mhs_n18_anl.2014080400
diag_mhs_n18_ges.2014080400
diag_mhs_n19_anl.2014080400
diag_mhs_n19_ges.2014080400
diag_mls30_aura_anl.2014080400
diag_mls30_aura_ges.2014080400
diag_omi_aura_anl.2014080400
diag_omi_aura_ges.2014080400
diag_pcp_tmi_trmm_anl.2014080400
diag_pcp_tmi_trmm_ges.2014080400
diag_sbuv2_n19_anl.2014080400
diag_sbuv2_n19_ges.2014080400
diag_seviri_m10_anl.2014080400
diag_seviri_m10_ges.2014080400
diag_sndrd1_g13_anl.2014080400
diag_sndrd1_g13_ges.2014080400
diag_sndrd2_g13_anl.2014080400
diag_sndrd2_g13_ges.2014080400
diag_sndrd3_g13_anl.2014080400
diag_sndrd3_g13_ges.2014080400
diag_sndrd4_g13_anl.2014080400
diag_sndrd4_g13_ges.2014080400
fit_q1.2014080400
fit_rad1.2014080400
fit_t1.2014080400
fit_w1.2014080400
fort.201
fort.202
fort.203
fort.204
fort.205
fort.206
fort.207
fort.208
fort.209
fort.210
fort.211
fort.212
fort.213
fort.214
fort.215
fort.217
fort.218
fort.219
fort.220
fort.221
fort.222
fort.223
fort.224
fort.225
fort.226
fort.227
fort.228
fort.229
fort.230
fort.232
fort.233
fort.234
fort.235
gomebufr
gpsrobufr
gsiparm.anl
gsi.x
gsnd1bufr
hirs3bufrears
hirs4bufr
hybens_info
iasibufr
list_run_directory
mhsbufr
mlsbufr
omibufr
ozinfo
pcpbias_out
pcpinfo
prepbufr
prepobs_prep.bufrtable
satbias_angle
satbias_ang.out
satbias_in
satbias_out
satbias_out.int
satbias_pc.in
satbias_pc.out
satinfo
satwndbufr
sbuvbufr
scaninfo
seviribufr
sfcanl.gsi
siganl
stdout
stdout.anl.2014080400
tcviltl

```

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diag_hirs4_metop-a_ges.2014080400 errtable fort.236 tmirrbuf
diag_hirs4_metop-b_anl.2014080400 fit_p1.2014080400 fort.237

The majority of these files existed after running the GSI regional analysis examples in section 3.2.3 of the Basic User's Guide, and they provide the same information about the GSI run. Of note, the GSI global analysis run includes more radiance observations, resulting in more radiance diag files in this list. Instead of the single background file wrf_inout as seen with the regional analysis, the global analysis background is split between the two files siganl, for the atmosphere, and sfcanl.gsi for the surface. A quick check of the standard output file stdout shows information similar to that generated by the regional runs for the namelist, data ingest, and minimization, but is quite different with respect to information on the background IO.

Please visit our online tutorial for more details regarding how to conduct a global GSI run.

5.5 Introduction to Chemical Analysis

The GSI has also been developed to analyze chemical observations, such as MODIS AOD or PM2.5, to improve the pollution forecasts with chemical models. In this release, GSI can do the following chemical analyses:

Table 5.4: List of GSI chemical analyses

case	Chemical Model	background species	Observations
1	WRF-Chem	GOCART	MODIS AOD
2	WRF-Chem	GOCART	PM2.5
3	WRF-Chem	PM2.5	PM2.5
4	CMAQ	CMAQ	PM2.5

The GSI run script for a chemical analysis (`./ush/comgsi_run_chem.ksh`) and associated namelist (`./ush/comgsi_namelist_chem.sh`) are provided with this release. Sample background and observation files are provided through the on-line tutorial.

5.5.1 Setup GSI Run Scripts for Chemical Analysis

The script `comgsi_run_chem.ksh` was built based on regional GSI run scripts and has a similar structure to the regional run script `comgsi_run_regional.ksh`, but include a couple of differences.

The first part of the run script sets up the computer environment and case configuration. This is similar to the regional analysis run scripts, except for the specification of (`bk_core` and `obs_type`) for a given chemical case, and the namelist for the chemical application:

```
GSI_NAMELIST=${GSI_ROOT}/ush/comgsi_namelist_chem.sh
```

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```
#-----
# bk_core= set background (WRFCHEM_GOCART WRFCHEM_PM25 or CMAQ)
# obs_type= set observation type (MODISAOD or PM25)
  bk_core=WRFCHEM_GOCART
  obs_type=PM25
```

The choices of (`bk_core` and `obs_type`) for a chemical case need to match with the options `PREPBUFR` and `BK_FILE`, which set background and observation files. Table 5.5 shows how to set up these two options for each case:

Table 5.5: List of GSI chemical analyses.

case	background (<i>BK_FILE</i> ; <i>bk_core</i>)	Observation (<i>PREPBUFR</i> ; <i>obs_type</i>)
1	wrfinput_enkf_d01_2012-06-03_18:00:00; WRFCHEM_GOCART	Aqua_Terra_AOD_BUFR:2012-06-03_00:00:00; MODISAOD
2	wrfinput_enkf_d01_2012-06-03_18:00:00; WRFCHEM_GOCART	anow.2012060318.bufr; PM25
3	wrfinput_enkf_d01_2012-06-03_18:00:00; WRFCHEM_PM25	anow.2012060318.bufr; PM25
4	cmaq2gsi_4.7_20130621_120000.bin; CMAQ	anow.2013062112.bufr; PM25

Similar to the regional run script, this chemical run script will also double check the needed parameters. Then it creates a run directory, generates the namelist, and copies the background, observations, and fixed files into the run directory. Users who run the cases listed in table 5.4 do not need to change the rest of the run script. But users who need to build new cases may need to know the differences between chemical and regional applications, which is shown below.

1. Specify the name of the background and observations:

```
# Bring over background field (it's modified by GSI so we can't link to it)

if [ ${bk_core} = WRFCHEM_GOCART ] ; then
  cp ${BK_FILE} ./wrf_inout
fi
if [ ${bk_core} = WRFCHEM_PM25 ] ; then
  cp ${BK_FILE} ./wrf_inout
fi
if [ ${bk_core} = CMAQ ] ; then
  cp ${BK_FILE} ./cmaq_in.bin
fi

# Link to the observation data
if [ ${obs_type} = MODISAOD ] ; then
  ln -s ${PREPBUFR} ./modisbufr
fi
if [ ${obs_type} = PM25 ] ; then
  ln -s ${PREPBUFR} ./pm25bufr
fi
```

2. Specify the background error file and anavinfo:

```
if [ ${bk_core} = WRFCHEM_GOCART ] ; then
  BERROR=${FIX_ROOT}/wrf_chem_berror_little_endian
  BERROR_CHEM=${FIX_ROOT}/wrf_chem_berror_little_endian
  ANAVINFO=${FIX_ROOT}/anavinfo_wrfchem_gocart
fi
if [ ${bk_core} = WRFCHEM_PM25 ] ; then
  BERROR=${FIX_ROOT}/wrf_chem_berror_little_endian
  BERROR_CHEM=${FIX_ROOT}/wrf_chem_berror_little_endian
  ANAVINFO=${FIX_ROOT}/anavinfo_wrfchem_pm25
fi
```

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```
if [ ${bk_core} = CMAQ ] ; then
  BERROR=${FIX_ROOT}/cmaq_berror_little_endian
  BERROR_CHEM=${FIX_ROOT}/cmaq_berror_little_endian
  ANAVINFO=${FIX_ROOT}/anavinfo_cmaq_pm25
fi
```

3. Specify the options for building the namelist:

```
if [ ${bk_core} = WRFCHEM_GOCART ] ; then
  bk_core_arw='.true.'
  bk_if_netcdf='.true.'
  bk_core_cmaq='.false.'
  bk_wrf_pm2_5='.false.'
  bk_laeroana_gocart='.true.'
fi
if [ ${bk_core} = WRFCHEM_PM25 ] ; then
  bk_core_arw='.true.'
  bk_if_netcdf='.true.'
  bk_core_cmaq='.false.'
  bk_wrf_pm2_5='.true.'
  bk_laeroana_gocart='.false.'
fi
if [ ${bk_core} = CMAQ ] ; then
  bk_core_arw='.false.'
  bk_if_netcdf='.false.'
  bk_core_cmaq='.true.'
  bk_wrf_pm2_5='.false.'
  bk_laeroana_gocart='.false.'
fi
```

5.5.2 Sample Results

In this section, case two in Table 5.4 will be used as an example. After a successful run of the GSI Chem analysis, the contents of the run directory, with the clean option turned on, will look something like this:

aeroinfo	fit_rad1.2012060318	fort.210	fort.225	gsiparm.anl	satbias_out.int
anavinfo	fit_t1.2012060318	fort.212	fort.226	gsi.x	satbias_pc_in
berror_stats	fit_w1.2012060318	fort.213	fort.227	list_run_directory	satbias_pc.out
berror_stats_chem	fort.201	fort.214	fort.228	ozinfo	satinfo
convinfo	fort.202	fort.215	fort.229	pcpbias_out	stdout
diag_conv_anl.2012060318	fort.203	fort.217	fort.230	pcpinfo	stdout.anl.2012060318
diag_conv_ges.2012060318	fort.204	fort.218	fort.232	pm25buf	wrfanl.2012060318
diag_light_anl.2012060318	fort.205	fort.219	fort.233	prepobs_prep.bufhtable	wrf_inout
diag_light_ges.2012060318	fort.206	fort.220	fort.234	satbias_angle	
errtable	fort.207	fort.221	fort.235	satbias_ang.out	
fit_p1.2012060318	fort.208	fort.223	fort.236	satbias_in	
fit_q1.2012060318	fort.209	fort.224	fort.237	satbias_out	

Following instructions from the previous sections, the following steps are conducted to check the results of this GSI chemical analysis:

1. Check the standard output file:

- Read in chemical background fields:

```
Vars in Aero-Jacobian (dims)
-----
sulf                0
bc1                 40
bc2                 80
oc1                 120
oc2                 160
```

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```

dust1                200
dust2                240
dust3                280
dust4                320
dust5                360
seas1                400
seas2                440
seas3                480
seas4                520
p25                  560
READ_wrf_mass_FILES: analysis date,minutes      2012          6          3          18          0          18
READ_wrf_mass_FILES: sigma guess file, nming2   0.0000000000000000          2012          6          3
READ_wrf_mass_FILES: sigma fcst files used in analysis :          3  3.0000000000000000          1
READ_wrf_mass_FILES: surface fcst files used in analysis:          3  3.0000000000000000          1
RADAR_BUFREAD_ALL: radial wind infile for superobbing =
***WARNING*** NOT USING ANY RADIAL WIND OBSERVATIONS!!!
GESINFO: Guess date is          18          6          3          2012  0.0000000000000000
GESINFO: Analysis date is          2012          6          3          18          0  2012060318  3.000000

```

- Read in ANOW PM2.5 observations:

```

READ_OBS: read 3 pm2_5 TEOM using ntasks= 1 0 1 999999 999999
...
READ_PM2_5: AIRNOW data type, subset=NC008032
READ_PM2_5: AIRNOW data type, subset=NC008032
READ_PM2_5: AIRNOW data type, subset=NC008032
READ_PM2_5: AIRNOW data type, subset=NC008032
READ_PM2_5: AIRNOW data type, subset=NC008032
READ_PM2_5: AIRNOW data type, subset=NC008032
READ_PM2_5: AIRNOW data type, subset=NC008032
READ_PM2_5: AIRNOW data type, subset=NC008032
READ_PM2_5: AIRNOW data type, subset=NC008032
READ_ANOWBUFR : file=pm25bufr type=pm2_5 sis=TEOM nread= 3981 ithin= 0 rmesh= 120.000000 isfcalc= 0 nkeep=

```

- Minimizations:

```

Begin J table inner/outer loop          0          1
J term                                  J
in-situ pm2_5 obs          1.3542969726002650E+03
-----
J Global          1.3542969726002650E+03
End Jo table inner/outer loop          0          1
Initial cost function = 1.354296972600265008E+03
Initial gradient norm = 5.380895196513087342E+01
cost,grad,step,b,step? = 1 0 1.354296972600265008E+03 5.380895196513087342E+01 1.854543171825029049E+02
cost,grad,step,b,step? = 1 1 9.182744794443540286E+02 2.959044916403530223E+01 1.524423141344592239E+02
cost,grad,step,b,step? = 1 2 8.108389171340875237E+02 1.786383268161806726E+01 2.399303783474798877E+02
cost,grad,step,b,step? = 1 3 7.473770294407435131E+02 7.768344135228370639E+00 2.566860407896649576E+02
...
cost,grad,step,b,step? = 2 9 7.373536121943825492E+02 8.155140099219158889E-03 2.931414472056467844E+02
cost,grad,step,b,step? = 2 10 7.373533822730103111E+02 2.973194603226746082E-03 2.525124974839193328E+02
cost,grad,step,b,step? = 2 11 7.373534027955187184E+02 2.181776986223444344E-03 1.584222202549521228E+02
PCGSOI: WARNING **** Stopping inner iteration ***
Penalty increase or constant 2 11 0.544454737560102542E+00 0.544454722406477631E+00
update_guess: successfully complete
\end{scriptsize}
\end{tiny}

\item Write out the increments statistics of the chemical variables, in addition to the traditional ones:
\begin{scriptsize}
\begin{verbatim}
increment u          0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
increment v          0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
increment tv         0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
increment tsen       0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
increment q          0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
increment oz         0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
increment cw         0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
increment p3d        0.000000000000E+00 0.000000000000E+00 0.000000000000E+00
increment sulf       -1.967693793248E-03 -1.771890969044E-01 1.968484278187E-03
increment bc1        -2.096573104319E-06 -2.930508183883E-04 1.019992534548E-05
increment bc2        -1.957703007985E-06 -8.967896393568E-05 7.146442589949E-08
increment oc1        -1.717720652720E-05 -3.473656841105E-03 1.800854291887E-04
increment oc2        -1.725425185217E-04 -8.176868955994E-03 6.630504242842E-06
increment dust1      -3.422858809284E-04 -2.224762201284E-02 4.144340973942E-05
increment dust2      -1.251405434281E-03 -8.311293736162E-02 2.331265353796E-04
increment dust3      0.000000000000E+00 0.000000000000E+00 0.000000000000E+00

```

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increment dust4	0.000000000000E+00	0.000000000000E+00	0.000000000000E+00
increment dust5	0.000000000000E+00	0.000000000000E+00	0.000000000000E+00
increment seas1	-3.418585319158E-06	-3.676233967264E-04	1.031774010195E-06
increment seas2	-2.824559081996E-04	-3.184781285067E-02	9.082224602894E-05
increment seas3	0.000000000000E+00	0.000000000000E+00	0.000000000000E+00
increment seas4	0.000000000000E+00	0.000000000000E+00	0.000000000000E+00
increment p25	-4.930178884615E-04	-3.499448870273E-02	1.367146375236E-04
increment ps	0.000000000000E+00	0.000000000000E+00	0.000000000000E+00
increment sst	0.000000000000E+00	0.000000000000E+00	0.000000000000E+00

2. Analysis increments: After successfully running GSI, the analysis increments should be checked to see if data impacts are reasonable.

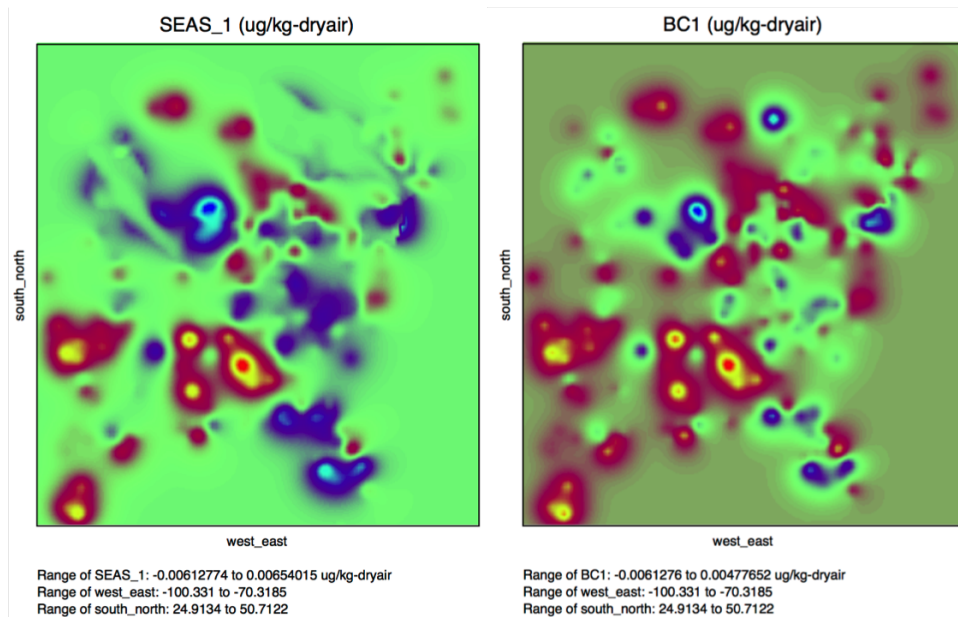


Figure 5.6: Analysis increments in the lowest level for SEAS_1 (left) and BC1 (right).

5.6 Summary

This chapter applied previous information outlined in the user's guide to demonstrate how to set up, run, and analyze GSI for various applications. It is important to always check for a successful GSI analysis, as running to completion does not always indicate a successful analysis was generated. Using the tools and methods described in this chapter, a complete picture of the GSI analysis can be obtained.



GSI Community Tools

A.1 BUFR Format and BUFR Tools

Under *./util/bufr_tools*, there are many Fortran examples to illustrate basic BUFR/PrepBUFR file process skills such as encoding, decoding, and appending. For details of these examples and the BUFR format, please see the BUFR/PrepBUFR User's Guide, which is freely available on-line

<http://www.dtcenter.org/com-GSI/BUFR/docs/index.php>

The observation BUFR files generated by NCEP (for example, PrepBUFR and BUFR files from NCEP ftp server or *gdasl.t12z.prepbufr.nr* in tutorial case) are in Big Endian binary format. For release version older than 3.2, the BUFR files have to be converted from Big Endian BUFR file to Little Endian file when used by the GSI on Linux platform. Please refer to older version User's guide on how to convert.

Since release 3.2, BUFRLIB can automatically identify and handle either byte orders. For Intel and PGI compilers on Linux, the Big Endian BUFR/PrepBUFR files can be used by GSI without byte swap.

A.2 Read GSI Diagnostic Files

Lots of useful information about how one observation was used in the analysis such as innovation, observation values, observation error and adjusted observation error, and quality control information, has been saved in diagnostic files. To generate these diagnosis files,

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namelist variable *write_diag* in namelist section *&SETUP* needs to be true. The *write_diag* variable has been introduced in Part 4 of Section 3.4. The following is an example of using the *write_diag* variable to control diagnostic files. When we set the number of outer loops to 2, and set the *write_diag* namelist variable to the following:

```
write_diag(1)=.true.,write_diag(2)=.false.,write_diag(3)=.true.,
```

GSI will write out diagnostic files before the start of the 1st outer loop start (O-B) and after the completion of the 2nd outer loop finish (O-A). We don't want GSI to write out diagnosis files after the 1st outer loop because we set *write_diag(2)=.false.*

This is what we set in our example case described in section 5.2. From this case, we can see the following diagnostic files generated from the GSI analysis:

```
diag_amsua_metop-a_anl.2014061700  diag_amsua_n18_ges.2014061700
diag_amsua_metop-a_ges.2014061700  diag_conv_anl.2014061700
diag_amsua_n15_anl.2014061700      diag_conv_ges.2014061700
diag_amsua_n15_ges.2014061700      diag_hirs4_metop-a_anl.2014061700
diag_amsua_n18_anl.2014061700      diag_hirs4_metop-a_ges.2014061700
```

All files are identified with a filename containing three elements. The first element "diag" indicates these are combined diagnostic files. The second element identifies the observation type (here, "conv" means conventional observation from prepbuf and "amsua_n15" corresponds to radiance observation AMSU-A from NOAA 15). The last element identifies which step of outer loop the files were generated for. Here, "anl" means the contents were written after the last outer loop (from *write_diag(3)=.true.*) and "ges" means the contents were written before the first output loop (from *write_diag(1)=.true.*).

To help users to read the information from these diagnostic files, we have provided two Fortran programs in the *./util/Analysis_Uilities/read_diag/* directory:

read_diag_conv.f90 : Reads the diagnostic files for conventional observations. For example:
diag_conv_anl.2014061700 and *diag_conv_ges.2014061700*

read_diag_rad.f90 : Reads the diagnostic files for radiance observation. For example:

```
diag_amsua_n15_ges.2014061700  diag_hirs4_metop-a_anl.2014061700
diag_amsua_n18_anl.2014061700  diag_hirs4_metop-a_ges.2014061700
```

To compile the programs, use the makefile provided:

```
./make
```

Note: since information in the GSI *include* directory is required, the GSI must have been compiled first.

To run *read_diag_conv.exe*, a namelist file *namelist.conv* needs to be in the directory along with the executable. The *namelist.conv* only has two parameters:

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```
&iosetup
  infilename='./diag_conv_anl',      : The path and name of GSI diagnosis file
  outfilename='./results_conv_anl', : The path and name of a text file used to
    save the content of the diagnostic file
/
```

The user can set the test case directory and file *diag_conv_anl.2014061700* from section 5.2 as the entry for *infilename* in the namelist, then run the executable

```
./read_diag_conv.exe
```

The results are placed in the file specified by the *outfilename* entry in the namelist. In this case that would be a file *results_conv_anl* located in the directory where the executable was run.

Similarly, to run *read_diag_rad.exe*, the namelist file *namelist.rad* is needed. It contains the same parameters as *namelist.conv* but it links to radiance diag files. After setting it to use the same case from section 5.2, such as:

```
&iosetup
  infilename='(test directory)/diag_amsua_n18_ges.2014061700',
  outfilename='./results_amsua_n18_ges',
/
```

Running the executable creates the text file *results_amsua_n18_ges* specified by the namelist in the directory *read_diag_rad.exe* runs.

For the conventional observations, the data is stored in two arrays: *rdiagbuf* and *cdiagbuf*. Their contents are listed as follows, for temperature (check *src/main/setupt.f90*) as an example:

```
cdiagbuf      = station id
rdiagbuf(1)   = observation type
rdiagbuf(2)   = observation subtype
rdiagbuf(3)   = observation latitude (degrees)
rdiagbuf(4)   = observation longitude (degrees)
rdiagbuf(5)   = station elevation (meters)
rdiagbuf(6)   = observation pressure (hPa)
rdiagbuf(7)   = observation height (meters)
rdiagbuf(8)   = observation time (hours relative to analysis time)
rdiagbuf(9)   = input prepbufr qc or event mark
rdiagbuf(10)  = setup qc or event mark (currently qtflg only)
rdiagbuf(11)  = read_prepbufr data usage flag
rdiagbuf(12)  = analysis usage flag (1=use, -1=not used)
rdiagbuf(13)  = nonlinear qc relative weight
rdiagbuf(14)  = prepbufr inverse obs error (K**-1)
rdiagbuf(15)  = read_prepbufr inverse obs error (K**-1)
rdiagbuf(16)  = final inverse observation error (K**-1)
rdiagbuf(17)  = observation (K)
rdiagbuf(18)  = obs-ges used in analysis (K)
rdiagbuf(19)  = obs-ges without bias correction (K)
```

For wind observations, the content after index 16 (check *src/main/setupw.f90*) is:

```
rdiagbuf(17) = earth relative u wind component observation (m/s)
```

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```

rdiagbuf(18) = earth relative u obs-ges used in analysis (m/s)
rdiagbuf(19) = earth relative u obs-ges w/o bias correction (m/s)
rdiagbuf(20) = earth relative v wind component observation (m/s)
rdiagbuf(21) = earth relative v obs-ges used in analysis (m/s)
rdiagbuf(22) = earth relative v obs-ges w/o bias correction (m/s)

```

The *read_diag_conv.exe* reads these arrays and outputs important information in the text file *results_conv_anl* specified by the user in the *&iosetup* namelist. For example:

station	obs	obs	obs	obs	obs	usag	obs	O-B		
ID	type	time	latitude	longitude	pressure		value			
ps @ 46047	: 180	-0.17	32.40	240.50	1014.30	1	1014.30	-0.09		
t @ 72293	: 120	-0.98	32.85	242.88	996.00	1	297.25	1.26		
uv @ 72293	: 220	-0.98	32.85	242.88	996.00	1	3.90	0.38	3.30	2.66

For wind, the last 4 columns are the wind components in the order of: U observation, O-B for U, V observation, O-B for V.

For radiance observations, the data is stored in two arrays: *diagbuf* and *diagbufchan*. Their contents are listed as follows (please refer to *src/main/setuprad.f90* for more details):

```

diagbuf(1) = observation latitude (degrees)
diagbuf(2) = observation longitude (degrees)
diagbuf(3) = model (guess) elevation at observation location
diagbuf(4) = observation time (hours relative to analysis time)

diagbuf(5) = sensor scan position
diagbuf(6) = satellite zenith angle (degrees)
diagbuf(7) = satellite azimuth angle (degrees)
diagbuf(8) = solar zenith angle (degrees)
diagbuf(9) = solar azimuth angle (degrees)
diagbuf(10) = sun glint angle (degrees) (sgagl)

diagbuf(11) = surface fractional coverage by water
diagbuf(12) = surface fractional coverage by land
diagbuf(13) = surface fractional coverage by ice
diagbuf(14) = surface fractional coverage by snow
if(.not. retrieval)then
diagbuf(15) = surface temperature over water (K)
diagbuf(16) = surface temperature over land (K)
diagbuf(17) = surface temperature over ice (K)
diagbuf(18) = surface temperature over snow (K)
diagbuf(19) = soil temperature (K)
diagbuf(20) = soil moisture
diagbuf(21) = surface land type
else
diagbuf(15) = SST first guess used for SST retrieval
diagbuf(16) = NCEP SST analysis at t
diagbuf(17) = Physical SST retrieval
diagbuf(18) = Navy SST retrieval
diagbuf(19) = d(ta) corresponding to sstph
diagbuf(20) = d(qa) corresponding to sstph
diagbuf(21) = data type
endif
diagbuf(22) = vegetation fraction
diagbuf(23) = snow depth
diagbuf(24) = surface wind speed (m/s)
! Note: The following quantities are not computed for all sensors
if (.not.microwave) then
diagbuf(25) = cloud fraction (%)
diagbuf(26) = cloud top pressure (hPa)
else
diagbuf(25) = cloud liquid water (kg/m**2)
diagbuf(26) = total column precip. water (km/m**2)
endif
endif

```

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```
diagbuf(27) = foundation temperature: Tr
diagbuf(28) = diurnal warming: d(Tw) at depth zob
diagbuf(29) = sub-layer cooling: d(Tc) at depth zob
diagbuf(30) = d(Tz)/d(Tr)
```

diagbufchan include loop through channel *i* from 1 to *nchanl*:

```
diagbufchan(1,i)= observed brightness temperature (K)
diagbufchan(2,i)= observed - simulated Tb with bias corrrection (K)
diagbufchan(3,i)= observed - simulated Tb with no bias correction (K)
diagbufchan(4,i)= inverse observation error
diagbufchan(5,i)= quality control mark or event indicator
diagbufchan(6,i)= surface emissivity
diagbufchan(7,i)= stability index
diagbufchan(8,i)= d(Tb)/d(Ts)
do j=1,npred+1
  diagbufchan(7+j,i)= Tb bias correction terms (K)
end do
```

In the sample output file *results_amsua_n18_ges*, only the observation location and time are written in the file. Users can write out other information based on the list.

A.3 Read and Plot Convergence Information from fort.220

In section 4.6, we introduced how to check the convergence information in the *fort.220* file. Further detail on the *fort.220* convergence information can be found in the Advanced User's Guide. Here, we provide tools to filter this file and to plot the values of the cost function and the norm of gradient during each iteration.

These tools - one ksh script and one ncl script - are in: *./util/Analysis_Uutilities/plot_cost_grad* directory:

The ksh script, *filter_fort220.ksh*, only has one line:

```
grep 'cost,grad,step,b' fort.220 | sed -e 's/cost,grad,step,b,step? = //g' | sed -e 's/good//g' > cost_gradient.txt
```

To run *filter_fort220.ksh*, the *fort.220* needs to be in the same directory. The script will filter out the values of the cost function and the norm of gradient at each iteration from *fort.220* into a text file called *cost_gradient.txt*.

Once the file *cost_gradient.txt* is ready, run ncl script *GSI_cost_gradient.ncl* to generate the plot:

```
ncl GSI_cost_gradient.ncl
```

The pdf file *GSI_cost_gradient.pdf* is created. The pdf file contains plots of the convergence of the GSI analysis like those in section 4.6.

A.4 Plot Single Observation Test Result and Analysis Increment

In Section 4.2, we introduced how to do a single observation test for GSI. Here we provide users with the ncl scripts to plot the single observation test results.

There are 5 ncl scripts in the *util/Analysis_Uutilities/plots_ncl* directory:

Table A.1: the list of ncl plotting tools

GSI_singleobs_arw.ncl	Plot single observation test with ARW NetCDF background
GSI_singleobs_nmm.ncl	Plot single observation test with NMM NetCDF background
Analysis_increment.ncl	Plot analysis increment from the case with ARW NetCDF background
Analysis_increment_nmm.ncl	Plot analysis increment from the case with NMM NetCDF background
fill_nmm_grid2.ncl	E grid to A grid convertor

The main difference between the ARW and NMM core used in GSI is that ARW is on a C grid, while NMM is on an E grid. *GSI_singleobs_nmm.ncl* calls *fill_nmm_grid2.ncl* to convert the E grid to an A grid for plotting, while *GSI_singleobs_arw.ncl* itself includes a C grid to A grid convertor.

Before running ncl scripts, users need to set up two links:

- *cdf_analysis* - Link to analysis result in NetCDF format (*wrf_inout*)
- *cdf_bk* - Link to background file in netCDF format

These scripts read in the analysis and background fields of temperature (T), U component of wind (U), V component of wind (V), and moisture (Q) and calculate the difference of the analysis field minus the background field. Then XY sections (left column) and XZ sections (right column) are plotted for T, U, V, and Q through the point that has maximum analysis increment of single observation. Here the default single observation test is T. If the user conducts other single observation tests, the corresponding changes should be made based on the current scripts.

The scripts *Analysis_increment.ncl* and *Analysis_increment_nmm.ncl* are very similar the one for the single observation but only the XY section for a certain analysis level is plotted.

For more information on how to use ncl, please check the NCL website at:

<http://www.ncl.ucar.edu/>

A.5 Generate initial regional ensembles

Under the `./util/EnKF/arw/src` directory, there are two sub-directories: `enspreproc_regional.fd/` and `initialens_regional.fd/`. The first one is to extract ensemble perturbations from GDAS 80 member ensembles and the second one is to add the extracted ensemble perturbations to a regional WRF background field (considered as the mean field) to generate initial regional ensembles.

Before using these two utilities, you should have already successfully compiled the GSI and gotten the "gsi.x" file. After that, under the build directory, type `"cmake -DBUILD_UTIL_COM path_to_comGSI"` to compile the utilities. A successful compilation should yield "enspreproc.x" and "initialens.x" in the bin/ directory.

Now, the next step is to get GDAS spectrally smoothed atmospheric ensemble forecasts. The old GDAS ensemble files are in the sigma format. You need to contact NCEP or other appropriate contacts to download these kind of ensembles. These ensemble files follow the name convention of "sfg_\$(CDATE)_fhr\$(FEs)_mem\$(MEM)". \$(CDATE) is the cycle date, such as 2017011518 which means 18z of Jan. 15th, 2017. \$(FE) is the forecast hour, for example, 06 means 6 hour of forecasts. \$(MEM) is the member number. Here is an example of GDAS ensembles: `sfg_2017011518_fhr06s_mem001`. After GFS system update in early 2017, the GDAS ensembles are in NEMSIO format. The name convention for NEMSIO format GDAS ensemble files follows "gdas.t\$(HR)z.atmf\$(FE)s.mem\$(MEM).nemsio". The FE and MEM have the same means as old ones and HR is cycling hour. For example, `gdas.t00z.atmf006s.mem001.nemsio`.

After you download the required GDAS ensembles, follow the following steps:

1. Running "enspreproc.exe", enter the `enspreproc_regional.fd/` directory:

(1). generate the file "fileslist01". This file lists the ensemble files to be used in the calculation of ensemble perturbations. For example, if it is determined to use 20 members to generate ensemble perturbations, the file "filelist01" will be as follows:

```
sfg_2017011518_fhr06s_mem001
sfg_2017011518_fhr06s_mem002
sfg_2017011518_fhr06s_mem003
...
sfg_2017011518_fhr06s_mem018
sfg_2017011518_fhr06s_mem019
sfg_2017011518_fhr06s_mem020
```

(2). Modify the file "namelist.input", change "n_ens" to the total number of ensembles to be used. For old GFS binary format ensemble files, the namelist option "use_gfs_nemsio" needs to be ".false.", while for NEMSIO format files, it needs to be ".true.".

(3). Copy the "anavinfo" file used by GSI into current directory.

(4). Copy the background WRF file, name it as "wrf_inout".

(5). Create a job description file, submit the job to get it run in parallel.

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After the successful running of "enspreproc.x", you will get ensemble perturbations as follows:

```
en_perts4ars.mem0001
en_perts4ars.mem0002
en_perts4ars.mem0003
...
en_perts4ars.mem0018
en_perts4ars.mem0019
en_perts4ars.mem0020
```

2. Running "initialens.x", enter the *initialens_regional.fd/* directory:

(1). Modify the file "namelist.input", change "n_ens" to the total number of ensembles to be used.

(2). Copy wrf_inout to current directory

(3). Copy wrf_inout to wrfinput_d01.mem\$MEM files as follows:

```
cp wrf_inout wrfinput_d01.mem0001
cp wrf_inout wrfinput_d01.mem0002
cp wrf_inout wrfinput_d01.mem0003
...
cp wrf_inout wrfinput_d01.mem0018
cp wrf_inout wrfinput_d01.mem0019
cp wrf_inout wrfinput_d01.mem0020
```

Be sure that each member has a corresponding wrfinput_d01 file. These files will be updated by "initialens.x" later.

(4). Link the ensemble perturbations generated by "enspreproc.x" to current directory. Something like this *ln -s ../enspreproc_regional.fd/en_perts4arw.mem**.

(5). Create a job description file, submit the job to get it run in parallel. Please note that only 1 processor is required to run "initialens.x" but submitting it to run on computing node is a must.

After the successful running of "initialens.x", all the *wrfinput_d01.mem\$MEM* files are updated with ensemble perturbation added to the background or "mean" state of the original wrf_inout.

Now the initial regional ensembles have been successfully generated.



Contents of Namelist Section

OBS_INPUT

```
&OBS_INPUT
  dmesh(1)=120.0,dmesh(2)=60.0,dmesh(3)=30,time_window_max=1.5,ext_sonde=.true.,
/
OBS_INPUT::
!  dfile      dtype      dplat      dsis          dval      dthin dsfcalc
  prepbufr    ps          null       ps            1.0       0       0
  prepbufr    t          null       t            1.0       0       0
  prepbufr    q          null       q            1.0       0       0
  prepbufr    pw         null       pw            1.0       0       0
  satwndbufr  uv         null       uv            1.0       0       0
  prepbufr    uv         null       uv            1.0       0       0
  prepbufr    spd        null       spd           1.0       0       0
  prepbufr    dw         null       dw            1.0       0       0
  radarbufr   rw         null       rw            1.0       0       0
  prepbufr    sst        null       sst           1.0       0       0
  gpsrobufr   gps_ref     null       gps           1.0       0       0
  ssmirrbufr  pcp_ssmi   dmstp     pcp_ssmi     1.0      -1       0
  tmirrbufr   pcp_tmi    trmm      pcp_tmi     1.0      -1       0
  sbuvbufr    sbuv2      n16       sbuv8_n16   1.0       0       0
  sbuvbufr    sbuv2      n17       sbuv8_n17   1.0       0       0
  sbuvbufr    sbuv2      n18       sbuv8_n18   1.0       0       0
  hirs3bufr   hirs3      n16       hirs3_n16   0.0       1       0
  hirs3bufr   hirs3      n17       hirs3_n17   6.0       1       0
  hirs4bufr   hirs4      metop-a   hirs4_metop-a 6.0       2       0
  hirs4bufr   hirs4      n18       hirs4_n18   0.0       1       0
  hirs4bufr   hirs4      n19       hirs4_n19   1.0       2       0
  hirs4bufr   hirs4      metop-b   hirs4_metop-b 1.0       1       0
  gimgrbufr   goes_img   g11       imgr_g11    0.0       1       0
  gimgrbufr   goes_img   g12       imgr_g12    0.0       1       0
  airsbufr    airs       aqua      airs281SUBSET_aqua 20.0      2       0
  amsuabufr   amsua     n15       amsua_n15   10.0      2       0
  amsuabufr   amsua     n18       amsua_n18   10.0      2       0
  amsuabufr   amsua     n19       amsua_n19   10.0      2       0
  amsuabufr   amsua     metop-a   amsua_metop-a 10.0      2       0
```

B. Contents of Namelist Section OBS_INPUT

amsuabufr	amsua	metop-b	amsua_metop-b	10.0	2	0
airsbufr	amsua	aqua	amsua_aqua	5.0	2	0
amsubbufr	amsub	n17	amsub_n17	1.0	1	0
mhsbufr	mhs	n18	mhs_n18	3.0	2	0
mhsbufr	mhs	n19	mhs_n19	3.0	2	0
mhsbufr	mhs	metop-a	mhs_metop-a	3.0	2	0
mhsbufr	mhs	metop-b	mhs_metop-b	3.0	2	0
ssmitbufr	ssmi	f13	ssmi_f13	0.0	2	0
ssmitbufr	ssmi	f14	ssmi_f14	0.0	2	0
ssmitbufr	ssmi	f15	ssmi_f15	0.0	2	0
amsrebufr	amsre_low	aqua	amsre_aqua	0.0	2	0
amsrebufr	amsre_mid	aqua	amsre_aqua	0.0	2	0
amsrebufr	amsre_hig	aqua	amsre_aqua	0.0	2	0
ssmisbufr	ssmis_las	f16	ssmis_f16	0.0	2	0
ssmisbufr	ssmis_uas	f16	ssmis_f16	0.0	2	0
ssmisbufr	ssmis_img	f16	ssmis_f16	0.0	2	0
ssmisbufr	ssmis_env	f16	ssmis_f16	0.0	2	0
gsnd1bufr	sndrd1	g12	sndrD1_g12	1.5	1	0
gsnd1bufr	sndrd2	g12	sndrD2_g12	1.5	1	0
gsnd1bufr	sndrd3	g12	sndrD3_g12	1.5	1	0
gsnd1bufr	sndrd4	g12	sndrD4_g12	1.5	1	0
gsnd1bufr	sndrd1	g11	sndrD1_g11	1.5	1	0
gsnd1bufr	sndrd2	g11	sndrD2_g11	1.5	1	0
gsnd1bufr	sndrd3	g11	sndrD3_g11	1.5	1	0
gsnd1bufr	sndrd4	g11	sndrD4_g11	1.5	1	0
gsnd1bufr	sndrd1	g13	sndrD1_g13	1.5	1	0
gsnd1bufr	sndrd2	g13	sndrD2_g13	1.5	1	0
gsnd1bufr	sndrd3	g13	sndrD3_g13	1.5	1	0
gsnd1bufr	sndrd4	g13	sndrD4_g13	1.5	1	0
gsnd1bufr	sndrd1	g15	sndrD1_g15	1.5	2	0
gsnd1bufr	sndrd2	g15	sndrD2_g15	1.5	2	0
gsnd1bufr	sndrd3	g15	sndrD3_g15	1.5	2	0
gsnd1bufr	sndrd4	g15	sndrD4_g15	1.5	2	0
iasibufr	iasi	metop-a	iasi616_metop-a	20.0	1	0
gomebufr	gome	metop-a	gome_metop-a	1.0	2	0
omibufr	omi	aura	omi_aura	1.0	2	0
sbuvbufr	sbuv2	n19	sbuv8_n19	1.0	0	0
tcvltl	tcp	null	tcp	1.0	0	0
seviribufr	seviri	m08	seviri_m08	1.0	1	0
seviribufr	seviri	m09	seviri_m09	1.0	1	0
seviribufr	seviri	m10	seviri_m10	1.0	1	0
iasibufr	iasi	metop-b	iasi616_metop-b	0.0	1	0
gomebufr	gome	metop-b	gome_metop-b	0.0	2	0
atmsbufr	atms	npp	atms_npp	0.0	1	0
crisbufr	cris	npp	cris_npp	0.0	1	0
mlsbufr	mls30	aura	mls30_aura	0.0	0	0
oscatbufr	uv	null	uv	0.0	0	0
prepbufr	mta_cld	null	mta_cld	1.0	0	0
prepbufr	gos_ctp	null	gos_ctp	1.0	0	0
refInGSI	rad_ref	null	rad_ref	1.0	0	0
lghtInGSI	lghtn	null	lghtn	1.0	0	0
larcInGSI	larccld	null	larccld	1.0	0	0
::						



GSI Namelist: Name, Default Value, Explanation

The following are lists and explanations of the GSI namelist variables. You can also find them in the source code **gsimod.F90**.

Variable name	Default value	Description
SETUP		General control namelist
gencode	80	source generation code
factqmin	1	weighting factor for negative moisture constraint
factqmax	1	weighting factor for supersaturated moisture constraint
clip_supersaturation	.false.	flag to remove supersaturation during each outer loop
factv	1	weighting factor for negative visibility constraint
factl		
factp		
factg		
factw10m		
fachowv		
deltim	1200	model timestep
dtphys	3600	physics timestep
biascor	-1	background error bias correction coefficient
bcoption	1	0=ibc (no bias correction to bkg); 1= sbc(original implementation)
diurnalbc	0	1= diurnal bias; 0= persistent bias

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
SETUP		General control namelist
niter(0:50)	0,...	Maximum number of inner loop iterations for each outer loop
niter_no_qc(0:50)	1000000	Inner loop iteration at which to turn on variational quality control
miter	1	number of outer loops
qoption	1	option for moisture analysis variable; 1:q/qsatg 2:normalized RH
cwoption		
pseudo_q2	.false.	breed between q1/q2 options, that is, (q1/sig(q))
nhr_assimilation	6	assimilation time interval (currently 6 hours for global, 3 hours for regional)
min_offset	3	time of analysis in assimilation window
iout_iter	220	output file number for iteration information
npredp	6	number of predictors for precipitation bias correction
retrieval	.false.	logical to turn off or on the SST physical retrieval
nst_gsi	0	indicator to control the Tr Analysis mode: 0 = no nst info ingi at all; 1 = input nst info, but used for monitoring only 2 = input nst info, and used in CRTM simulation, but no Tr analysis 3 = input nst info, and used in CRTM simulation and Tr analysis is on
nst_tzr	0	indicator to control the Tzr_QC mode: 0 = no Tz retrieval; 1 = Do Tz retrieval and applied to QC
nstinfo	0	number of nst variables
fac_dtl	0	index to apply diurnal thermocline layer or not: 0 = no; 1 = yes
fac_tsl	0	index to apply thermal skin layer or not: 0 = no; 1 = yes.
nst_tzr		
tzr_bufsave	.false.	logical to turn off or on the bufr Tz retrieval file true=on
diag_rad	.true.	logical to turn off or on the diagnostic radiance file (true=on)
diag_pcp	.true.	logical to turn off or on the diagnostic precipitation file (true=on)
diag_conv	.true.	logical to turn off or on the diagnostic conventional file (true=on)
diag_ozone	.true.	logical to turn off or on the diagnostic ozone file (true=on)
diag_aero	.false.	logical to turn off or on the diagnostic aerosol file (true=on)
diag_co	.false.	logical to turn off or on the diagnostic carbon monoxide file (true=on)
iguess	1	flag for guess solution (currently not working) -1 do not use guess file 0 write only guess file 1 read and write guess file 2 read only guess file
write_diag	.false., ...	logical to write out diagnostic files for outer iteration
reduce_diag	.false.	namelist logical to produce reduced radiance diagnostic files

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
SETUP		General control namelist
oneobtest	.false.	one observation test flag true=on
sfcmodel	.false.	if true, then use boundary layer forward model for surface temperature data.
dtbduv_on	.true.	logical for switching on (.true.) sensitivity of uv winds to microwave brightness temperatures. if true, use d(microwave brightness temperature)/d(uv wind) in inner loop
ifact10	0	flag for recomputing 10m wind factor = 1 compute using GFS surface physics = 2 compute using MM5 surface physics = 0 or any other value - DO NOT recompute - use value from guess file
l_foto	.false.	option for First-Order Time extrapolation to observation
offtime_data	.false.	if true, then allow use of obs files with ref time different from analysis time. default value = .false., in which case analysis fails if observation file reference time is different from analysis time.
npred_conv_max	0	maximum number of conventional observation bias correction coefficients
id_bias_ps	0	prepbufr id to have conv_bias added for testing
id_bias_t	0	prepbufr id to have conv_bias added for testing
id_bias_spd	120	prepbufr id to have conv_bias added for testing
conv_bias_ps	0	magnitude of ps bias(mb)
conv_bias_t	0	magnitude of t bias(deg K)
conv_bias_spd	0	magnitude of spd bias(m/sec)
id_bias_pm2_5		
conv_bias_pm2_5		
id_bias_pml0		
conv_bias_pml0		
stndev_conv_ps	1.0	
stndev_conv_t	1.0	
stndev_conv_spd	1.0	
use_pbl	.false.	Logical flag to include PBL effects in tendency model.
use_compress	.false.	option to turn on the use of compressibility factors in geopotential heights
nsig_ext	13	number of layers above the model top which are necessary to compute the bending angle for gpsro
gpstop	30.0	maximum height for gpsro data assimilation. Reject anything above this height. (km)
perturb_obs	.false.	logical flag to perturb observation (true=on)
perturb_fact	1	magnitude factor for observation perturbation
oberror_tune	.false.	logical to tune (=true) oberror
preserve_restart_date	.false.	if true, then do not update regional restart file date.
crtm_coeffs_path	./	path of directory w/ CRTM coeffs files
berror_stats	berror_stats	filename if other than "berror_stats"
newpc4pred	.false.	option for additional preconditioning for pred coeff
adp_anglebc	.false.	option to perform variational angle bias correction
angord	0	order of polynomial for variational angle bias correction

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
SETUP		General control namelist
passive_bc	.false.	option to turn on bias correction for passive (monitored) channels
use_edges	.true.	option to exclude radiance data on scan edges
biaspredvar	0.1	set background error variance for radiance bias coeffs
lobsdiagsave	.false.	write out additional observation diagnostics
l4dvar	.false.	turn 4D-Var on/off (default=off=3D-Var)
lbicg	.false.	use B-precond w/ bi-conjugate gradient for minimization
lsqrtb	.false.	Use sqrt(B) preconditioning
lcongrad	.false.	Use conjugate gradient/Lanczos minimizer
lbfgsmin	.false.	Use L-BFGS minimizer
ltilint	.false.	Use TL inner loop (ie TL intall)
nhr_obsbin	-1	length of observation bins
nhr_subwin	-1	length of weak constraint 4d-Var sub-window intervals
nwrvecs	-1	Number of precond vectors (Lanczos) or pairs of vectors (QN) being saved
iorthomax	0	max number of vectors used for orthogonalization of various CG options
ladtest	.false.	Run adjoint test
ladtest_obs	.false.	if true, doing the adjoint check for the observation operators
lgrtest	.false.	Run gradient test
lobskeep	.false.	keep obs from first outer loop for subsequent OL
lsensrecompute	.false.	does adjoint by recomputing forward solution
jsiga	-1	calculate approximate analysis errors from lanczos for jiter=jsiga
lrcost	.false.	calculate true cost when using Lanczos (this is very expensive)
lobsensfc	.false.	compute forecast sensitivity to observations
lobsensjb	.false.	compute Jb sensitivity to observations
lobsensincr	.false.	compute increment sensitivity to observations
lobsensadj	.false.	use adjoint of approx. Hessian to compute obs sensitivity
lobsensmin	.false.	use minimisation to compute obs sensitivity
iobsconv	0	compute convergence test in observation space =1 at final point, =2 at every iteration
idmodel	.false.	uses identity model when running 4D-Var (test purposes)
iwrtinc	.false.	when .t., writes out increments instead of analysis
jiterstart	1	first outloop iteration number
jiterend	1	last outloop iteration number
lobserver	.false.	when .t., calculate departure vectors only
lanczosave	.false.	save lanczos vectors for forecast sensitivity computation
llancdone	.false.	use to tell adjoint that Lanczos vecs have been pre-computed
lferrscale	.false.	Something related to forecast error
print_diag_pcg	.false.	logical turn on of printing of GMAO diagnostics in pcgsoi.f90
tsensible	.false.	option to use sensible temperature as the analysis variable. Works only for twodvar_regional=.true.
lgschmidt	.false.	option for re-biorthogonalization of the gradx and grady set from pcgsoi when twodvar_regional=.true.

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
SETUP		General control namelist
lread_obs_save	.false.	option to write out collective obs selection info
lread_obs_skip	.false.	option to read in collective obs selection info
use_gfs_ozone	.false.	option to read in gfs ozone and interpolate to regional model domain
check_gfs_ozone_date	.false.	option to date check gfs ozone before interpolating to regional model domain
regional_ozone	.false.	option to turn on ozone in regional analysis
lwrite_predterms	.false.	option to write out actual predictor terms instead of predicted bias to the radiance diagnostic files
lwrite_peakwt	.false.	option to writ out the approximate pressure of the peak of the weighting function for satellite data to the radiance diagnostic files
use_gfs_nemsio	.false.	option to use nemsio to read global model NEMS/GFS first guess
liauon	.false.	treat 4dvar CV as tendency perturbation (default=false)
use_prepb_satwnd	.false.	allow using satwnd's from prepbuf (historical) file
l4densvar	.false.	logical to turn on ensemble 4dvar
ens4d_nstarthr	3	start hour for ensemble perturbations (generally should match min_offset)
use_gfs_stratosphere		When true, a guess gfs valid at the same time as the nems-nmmb guess is used to replace the upper levels with gfs values. The purpose of this is to allow direct use of gdas derived sat radiance bias correction coefs.
pblend0	152	The nems-nmmb vertical coordinate is smoothly merged with gfs above this level. Below this level, is original nems-nmmb.
pblend1	79.0	The nems-nmmb vertical coordinate is smoothly merged with gfs below this level. Above this level, is gfs.
step_start	1.e-4	initial stepsize in minimization
diag_precon	.false.	if true do preconditioning
lrun_subdirs	.false.	logical to toggle use of subdirectires at runtime for pe specific files
emiss_bc	.false.	option to turn on emissivity bias predictor
upd_pred	1	bias update indicator for radiance bias correction; 1.0=bias correction coefficients evolve
use_reflectivity	.false.	option of using reflectivity
lnested_loops	.false.	allow for nested resolution outer/inner loops
lwrite4danl	.false.	logical to write out 4d analysis states if 4dvar or 4denvar mode
lsingleradob	.false.	logical for single radiance observation assimilation. Uses existing bufr file and rejects all radiances that don't fall within a tight threshold around oblat/oblon (SINGLEOB_TEST)
ssmis_method	1	choose method for SSMIS noise reduction 0=no smoothing 1=default
ssmis_precond	0.01	weighting factor for SSMIS preconditioning (if not using newpc4pred)
R_option	.false.	Option to use variable correlation length for lcbas based on data density - follows Hayden and Purser (1995) (twodvar_regional only)
thin4d		

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
GRIDOPTS		Grid setup variables, including regional specific variables
jcap	62	spectral resolution of the analysis
jcap_b	62	spectral resolution of background (model guess field)
nsig	42	number of sigma levels
nlat	96	number of latitudes
nlon	384	number of longitudes
hybrid	logical	hybrid data file flag true=hybrid
nlat_regional	0	Number of y grid point in whole regional domain
nlon_regional	0	Number of x grid point in whole regional domain
diagnostic_reg	.false.	logical for regional debugging
update_regsfc	.false.	logical to write out updated surface fields to the regional analysis file (default = false)
netcdf	.false.	if true, then wrf files are in netcdf format, otherwise wrf files are in binary format.
regional	.false.	logical for regional GSI run
wrf_nmm_regional	.false.	logical for input from WRF NMM
nems_nmmb_regional	.false.	logical for input from NEMS NMMB
wrf_mass_regional	.false.	logical for input from WRF MASS-CORE (ARW)
twodvar_regional	.false.	logical for regional 2d-var analysis
filled_grid	.false.	logical to fill in points on WRF-NMM E-grid
half_grid	.false.	logical to use every other row of WRF-NMM E-Grid
nvege_type	24	number of types of vegetation; old=24, IGBP=20
nlayers(100)	1	number of sub-layers to break indicated model layer into prior to calling radiative transfer model
cmaq_regional	.false.	Background input is from CMAQ model
nmmb_reference_grid	H	= 'H', then analysis grid covers H grid domain = 'V', then analysis grid covers V grid domain
grid_ratio_nmmb	sqrt(2)	ratio of analysis grid to nmmb model grid in nmmb model grid units.
grid_ratio_wrfmass	1.0	ratio of analysis grid to wrf mass grid in wrf grid units
jcap_gfs		spectral truncation used to transform high wavenumber spectral coefficients to a coarser resolution grid, when use_gfs_ozone = .true. or use_gfs_stratosphere = .true.
jcap_cut		
use_sp_eqspac	.false.	if .true., then ensemble grid is equal spaced, staggered 1/2 grid unit off poles. if .false., then gaussian grid assumed for ensemble (global only)

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
BKGERR		Background error related variables
vs	1/1.5	scale factor for vertical correlation lengths for background error
nhscrf	3	number of horizontal scales for recursive filter
hzscl(3)	1, 1, 1	scale factor for horizontal smoothing, n=1,number of scales (3 for now) specifies factor by which to reduce horizontal scales (i.e. 2 would then apply 1/2 of the horizontal scale)
hswgt(3)	1/3, 1/3, 1/3	empirical weights to apply to each horizontal scale
norh	2	order of interpolation in smoothing
ndeg	4	degree of smoothing in recursive filters
noq	3	1/4 of accuracy in compact finite differencing
bw	0	factor in background error calculation
norsp	0	order of interpolation for smooth polar cascade routine default is norsp=0, in which case norh is used with original polar cascade interpolation (global only).
fstat	.false.	logical to separate f from balance projection
pert_berr	.false.	logical to turn on random inflation/deflation of background error tuning parameters
pert_berr_fct	0	factor for increasing/decreasing berror parameters, this is multiplied by random number
bkgv_flowdep	.false.	flag to turn on flow dependence to background error variances
bkgv_rewgtfct	0	factor used to perform flow dependent reweighting of error variances
bkgv_write	.false.	flag to turn on=.true. /off=.false. generation of binary file with reweighted variances
fpsproj	.true.	controls full nsig projection to surface pressure
fut2ps		controls the projection from unbalance T to surface pressure
adjustozvar		adjusts ozone variances in the stratosphere based on guess field
cwcoveqqcov		sets cw Bcov to be the same as B-cov(q) (presently glb default)

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
ANBKGERR		Anisotropic background error related variables
anisotropic	.false.	if true, then use anisotropic background error covariance
ancovmdl	0	covariance model settings - 0: pt-based, 1: ensemble based
triad4	.true.	for 2d variables, if true, use blended triad algorithm
ifilt_ord	4	filter order for anisotropic filters
npass	1	2 \sqrt{N} npass = number of factors in background error
normal	200	number of random vectors to use for filter normalization (if < 0 then slightly slower, but results independent of number of processors)
binom	.true.	if true, weight correlation lengths of factors using binomial distribution, with shortest scales on outside, longest scales on inside. This can help to produce smoother correlations in the presence of strong anisotropy
ngauss	3	number of Gaussians to add together in each factor
rgauss	0	multipliers on reference aspect tensor for each Gaussian factor
anhswgt	1.0	empirical weights to apply to each gaussian
an_vs	1	scale factor for background error vertical scales (temporary carry over from isotropic inhomogeneous option)
grid_ratio	2.0	ratio of coarse to fine grid in fine grid units
grid_ratio_p	0	ratio of coarse to fine grid in fine grid units for polar patches
nord_f2a	4	order of interpolation for transfer operators between filter grid and analysis grid
an_flen_u	1	coupling parameter for connecting horizontal wind to background error
an_flen_t	1	coupling parameter for connecting grad(potential temperature) to background error
an_flen_z	1	coupling parameter for connecting grad(terrain) to background error
rtma_subdomain_option	.false.	if true, then call alternative code which calls recursive filter directly from subdomain mode, bypassing transition to/from horizontal slabs. This is mainly to improve efficiency for 2d rtma analysis. at the moment, this only works for twodvar_regional=.true. rtma_subdomain_option will be forced to false when twodvar_regional=.false.
lreadnorm	.false.	if true, then read normalization from fixed files
nsmooth	0	number of 1-2-1 smoothing passes before and after background error application
nsmooth_shapiro	0	number of 2nd moment preserving (shapiro) smoothing passes before and after background error application. NOTE: default for nsmooth and nsmooth_shapiro is 0. if both are > 0, then nsmooth will be forced to zero.
afact0	0.0	anistropy effect parameter, the range must be in 0.0-1.0.
covmap	.false.	if true, covariance map would be drawn

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
JCOPTS		Constraint term in cost function (Jc)
ljcdfi	.false.	if .false., uses original formulation based on wind, temp, and ps tends when .t. uses digital filter initialization of increments (4dvar)
alphajc	10.0	parameter for digital filter
switch_on_derivatives	.false., $\hat{\Delta}\epsilon$	if true, then compute horizontal derivatives of all state variables (to be used eventually for time derivatives, dynamic constraints and observation forward models that need horizontal derivatives)
tendsflag	.false.	if true, compute time tendencies
ljcpdry	.false.	when .t. uses dry pressure constraint on increment
bamp_jcpdry	0.0	parameter for pdry_jc
eps_eer	-1.0	Errico-Ehrendofer parameter for q-term in energy norm
ljc4tlevs	.false.	when true and in 4D mode, apply any weak constraints over all time levels instead of just at a single time

Variable name	Default value	Description
STRONGOPTS		Strong dynamic constraint
reg_tlnmc_type	1	=1 for 1st version of regional strong constraint =2 for 2nd version of regional strong constraint
tlnmc_option	0	integer flag for strong constraint (various capabilities for hybrid): =0: no TLNMC =1: TLNMC for 3DVAR mode =2: TLNMC on total increment for single time level only (for 3D EnVar) or if 4D EnVar mode, TLNMC applied to increment in center of window =3: TLNMC on total increment over all time levels (if in 4D EnVar mode) =4: TLNMC on static contribution to increment ONLY for any EnVar mode
nstrong	0	if > 0, then number of iterations of implicit normal mode initialization to apply for each inner loop iteration
period_max	1000000.0	cutoff period for gravity waves included in implicit normal mode initialization (units = hours)
period_width	1.0	defines width of transition zone from included to excluded gravity waves
nmodes_keep	0	number of vertical modes to use in implicit normal mode initialization
baldiag_full	.false.	flag to toggle balance diagnostics for the full fields
baldiag_inc	.false.	flag to toggle balance diagnostics for the analysis increment

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
OBSQC		Observation quality control variables Parameters used for gross error checks are set in file con- vinfo (ermin, ermax, ratio) Parameters below used for non- linear (variational) quality control
dfact	0	factor for duplicate observation at same location for conven- tional data
dfact1	3.0	time factor for duplicate observation at same location for con- ventional data
erradar_inflate	1	radar error inflation factor
tdrerr_inflate	.false.	logical for tdr obs error inflation
tdrgross_fact	1	factor applied to tdr gross error
oberrflg	.false.	logical for reading in new observation error table (if set to true)
vadfile	'none'	character(10) variable holding name of VAD wind bufr file
noiqc	.false.	logical flag to bypass OI QC (if set to true)
c_varqc	1	constant number to control variance qc turning on speed
blacklst	.false.	logical for reading in raob blacklist (if set to true)
use_poq7	.false.	Logical to toggle accept (.true.) or reject (.false.) SBUV/2 ozone observations flagged with profile ozone quality mark
hilbert_curve	.false.	option for hilbert-curve based cross-validation. works only with twodvar_regional=.true.
tcp_refps	1000.0	reference pressure for tcps oberr calculation (mb)
tcp_width	50.0	parameter for tcps oberr inflation (width, mb)
tcp_ermin	0.75	parameter for tcps oberr inflation (minimum oberr, mb)
tcp_erman	5.0	parameter for tcps oberr inflation (maximum oberr, mb)
qc_noirjaco3	.false.	controls whether to use O3 Jac from IR instruments
qc_noirjaco3_pole	.false.	controls wheter to use O3 Jac from IR instruments near poles
qc_satwnds	.true.	allow bypass sat-winds qc normally removing lots of mid-tropo obs
njqc		
vqc		
aircraft_t_bc_pof	.false.	logical for aircraft temperature bias correction, pof is used for predictor
aircraft_t_bc	.false.	logical for aircraft temperature bias correction
aircraft_t_bc_ext	.false.	logical for reading aircraft temperature bias correction from external file
buddycheck_t	.false.	When true, run buddy check algorithm on temperature obser- vations
buddydiag_save	.false.	When true, output files containing buddy check QC info for all obs run through the buddy check
biaspredt	1	berror var for temperature bias correction coefficients
upd_aircraft	.true.	indicator if update bias at 06Z & 18Z
cleanup_tail	.false.	logical to remove tail number no longer used

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
OBS_INPUT		Controls input data
dfile	' '	input observation file name
dtype	' '	observation type
dplat	' '	satellite (platform) id (for satellite data)
dsis	' '	sensor/instrument/satellite flag from satinfo files
dthin	' '	satellite group
dval	' '	relative value of each profile within group relative weight for observation = dval/sum(dval) within grid box
dmesh(max(dthin))		thinning mesh for each group mesh size (km) for radiance thinning grid (used in satthin)
dsfcalc	' '	specifies method to determine surface fields within a FOV. when equal to one, integrate model fields over FOV. when not one, bilinearly interpolate model fields to FOV center.
time_window_max	3	upper limit on time window for all input data
ext_sonde	.false.	logical for extended forward model on sonde data
l_foreaft_thin	.false.	separate TDR fore/aft scan for thinning

Variable name	Default value	Description
SINGLEOB_TEST		Single observation test case setup
maginnov	1	magnitude of innovation for one observation
magoberr	1	magnitude of observational error
oneob_type	' '	observation type (t, u, v, etc.)
oblat	0	observation latitude
oblon	0	observation longitude
obpres	1000.0	observation pressure (hPa)
obdattim	2000010100	observation date (YYYYMMDDHH)
obhourset	0	observation delta time from analysis time
pctswitch	.false.	if .true. innovation & oberr are relative (%) of background value (level ozone only)
obchan	0	if > 0, selects the channel number. If <= zero, it will use all channels that pass qc in setuprad.

Variable name	Default value	Description
SUPEROB_RADAR		Level 2 bufr file to radar wind superobs
del_azimuth	5.0	azimuth range for superob box (default 5 degrees)
del_elev	0.25	elevation angle range for superob box (default .05 degrees)
del_range	5000.0	radial range for superob box (default 5 km)
del_time	0.5	1/2 time range for superob box (default .5 hours)
elev_angle_max	5.0	max elevation angle (default of 5 deg) minnum 50 minimum number of samples needed to make a superob
range_max	100000.0	max radial range in meters to use in constructing superobs (default 100km)
l2superob_only	.false.	if true, then process level 2 data creating superobs, then quit. (added for easier retrospective testing, since level 2 bufr files are very large and hard to work with)

Variable name	Default value	Description
LAG_DATA 		Lagrangian data assimilation related variables
lag_accur	1.0e-6	Accuracy used to decide whether or not a balloon is on the grid
infile_lag	inistate_lag.dat	File containing the initial position of the balloon
lag_stepduration	900.0	Duration of one time step for the propagation model
lag_nmax_bal	1000	Maximum number of balloons at starting time
lag_vorcore_stderr_a	2.0e3	Observation error for vorcore balloon
lag_vorcore_stderr_b	0.0	error = b + a*timestep(in hours)

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
HYBRID_ENSEMBLE		Parameters for use with hybrid ensemble option
l_hyb_ens	.false.	if true, then turn on hybrid ensemble option
uv_hyb_ens	.false.	if true, then ensemble perturbation wind variables are u,v, otherwise, ensemble perturbation wind variables are stream, pot. Functions.
q_hyb_ens	.false.	if true, then use specific humidity ensemble perturbations, otherwise, use relative humidity
aniso_a_en	.false.	if true, then use anisotropic localization of hybrid ensemble control variable a_en.
generate_ens	.true.	if true, then generate internal ensemble based on existing background error
n_ens	0	number of ensemble members.
nlon_ens	0	number of longitudes on ensemble grid (may be different from analysis grid nlon)
nlat_ens	0	number of latitudes on ensemble grid (may be different from analysis grid nlat)
jcap_ens	0	for global spectral model, spectral truncation
pseudo_hyb_ens	.false.	if true, turn on pseudo ensemble hybrid for HWRF
merge_two_grid_enspers	.false.	if true, merge ensemble perturbations from two forecast domains to analysis domain (one way to deal with hybrid DA for HWRF moving nest)
regional_ensemble_option	0	integer, used to select type of ensemble to read in for regional application. Currently takes values from 1 to 4 =1: use GEFS internally interpolated to ensemble grid. =2: ensembles are WRF NMM format =3: ensembles are ARW netcdf format. =4: ensembles are NEMS NMMB format.
full_ensemble	.false.	if true, first ensemble perturbation on first guess instead of on ens mean
betaflg	.false.	if true, use vertical weighting on beta1_inv and beta2_inv, for regional
coef_bw	0.9	fraction of weight given to the vertical boundaries when betaflg is true
pwgtflg	.false.	if true, use vertical integration function on ensemble contribution of Psfc
jcap_ens_test	0	for global spectral model, test spectral truncation (to test dual resolution)
beta1_inv	1	1/beta1, the default weight given to static background error covariance if (.not. readin_beta) 0 <= beta1_inv <= 1, tuned for optimal performance =1, then ensemble information turned off =0, then static background turned off the weights are applied per vertical level such that : betas_inv(:) = beta1_inv , vertically varying weights given to static B ; betae_inv(:) = 1 - beta1_inv , vertically varying weights given ensemble derived covariance. If (readin_beta) then betas_inv and betae_inv are read from a file and beta1_inv is not used.

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
HYBRID_ENSEMBLE		Parameters for use with hybrid ensemble option
s_ens_h	2828	homogeneous isotropic horizontal ensemble localization scale (km)
s_ens_v	30	vertical localization scale (grid units for now) s_ens_h, s_ens_v, and betal_inv are tunable parameters.
use_gfs_ens	.true.	controls use of global ensemble: .t. use GFS (default); .f. uses user-defined ens
readin_localization	.false.	flag to read (.true.)external localization information file
readin_beta	.false.	flag to read (.true.) the vertically varying beta parameters betas_inv and betae_inv from a file.
eqspace_ensgrid	.false.	if .true., then ensemble grid is equal spaced, staggered 1/2 grid unit off ploe. if .false., then gaussian grid assumed for ensemble (global only)
use_localization_grid	.false.	if true, then use extra lower res gaussian grid for horizontal localization (global runs only--allows possiblity for non-gaussian ensemble grid)
grid_ratio_ens	1	for regional runs, ratio of ensemble grid resolution to analysis grid resolution default value = 1 (dual resolution off)
oz_univ_static	.false.	if true, decouple ozone from other variables and defaults to static B (ozone only)
write_ens_sprd	.false.	writing global ensemble spread in byte addressable format for plotting with grads
enspreproc	.false.	flag to read(.true.) pre-processed ensemble data already
i_en_perts_io	0	flag to read in ensemble perturbations in ensemble grid. This is to speed up RAP/HRRR hybrid runs because the same ensemble perturbations are used in 6 cycles =0: No ensemble perturbations IO (default) =2: skip get_gefs_for_regional and read in ensemble perturbations from saved files.
l_ens_in_diff_time	.false.	if use ensembles that are available at different time from analysis time. =false: only ensembles available at analysis time can be used for hybrid. (default) =true: ensembles available time can be different from analysis time in hybrid analysis
ensemble_path		

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
rapidrefresh_cldsrf		Options for cloud analysis and surface enhancement for RR application
dfi_radar_latent_heat_time_period	30.0	DFI forward integration window in minutes
metar_impact_radius	10.0	metar cloud observation impact radius in grid number
metar_impact_radius_lowCloud	4.0	impact radius for METAR cloud observation that indicate low cloud base
l_gsd_terrain_match_surfTobs	.false.	if .true., GSD terrain match for surface temperature observation
l_sfcobserror_ramp_t	.false.	namelist logical for adjusting surface temperature observation error
l_sfcobserror_ramp_q	.false.	namelist logical for adjusting surface moisture observation error
l_PBL_pseudo_SurfobsT	.false.	if .true. produce pseudo-obs in PBL layer based on surface obs T
l_PBL_pseudo_SurfobsQ	.false.	if .true. produce pseudo-obs in PBL layer based on surface obs Q
l_PBL_pseudo_SurfobsUV	.false.	if .true. produce pseudo-obs in PBL layer based on surface obs UV
pblH_ration	0.75	percent of the PBL height within which to add pseudo-obs
pps_press_incr	30hPa	pressure increase for each additional pseudo-obs on top of previous level
l_gsd_limit_ocean_q	.false.	if .true. do GSD limitation of Q over ocean
l_pw_hgt_adjust	.false.	if .true. do GSD PW adjustment for model vs. obs station height
l_limit_pw_innov	.false.	if .true. do GSD limitation of PW obs
max_innov_pct	0.1	sets limit of PW ob to a percent of the background value (0-1)
l_cleanSnow_WarmTs	.false.	if .true. do GSD limitation of using retrieved snow over warn area ($T_s > r_cleanSnow_WarmTs_threshold$)
l_conserve_thetaV	.false.	if .true. conserve thetaV during moisture adjustment in cloud analysis
r_cleanSnow_WarmTs_threshold	8.0	threshold for using retrieved snow over warn area
i_conserve_thetaV_iternum	3	iteration number for conserving thetaV during moisture adjustment
l_gsd_soilTQ_nudge	.false.	if .true. do GSD GOES cloud building
l_cld_bld	.false.	if .true. do GSD soil T and Q nudging based on the lowest t analysis increment
cld_bld_hgt	1200m	sets limit below which GOES cloud building occurs
build_cloud_frac_p	0.95	sets the threshold for building clouds from satellite
clear_cloud_frac_p	0.1	sets the threshold for clearing clouds from satellite
nesdis_npts_rad	1	NESDIS cloud product impact radiu (grid points)
iclean_hydro_withRef	1	if =1, then clean hydrometeors if the grid point has no echo and maxref=0
iclean_hydro_withRef_allcol	0	if =1, then clean whole column hydrometeors if the observed max ref =0 and satellite cloud shows clean

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
rapidrefresh_cldsrf		Options for cloud analysis and surface enhancement for RR application
l_use_2mq4b	0	background used for calculate surface moisture observation innovation =0 Use Q from the 1st model level. (default) =1 use 2m Q as part of background
i_use_2mt4b	0	background used for calculate surface temperature observation innovation =0 Use T from the 1st model level. (default) =1 use 2m T as part of background
i_gsdclanal_type	0	options for how GSD cloud analysis should be conducted =0. no cloud analysis (default) =1. cloud analysis after var analysis =5. skip cloud analysis and NETCDF file update
i_gsdsc_uselist	0	options for how to use surface observation use or rejection list =0 . EMC method (default) =1 . GSD method
i_lightpcp	0	options for how to deal with light precipitation =0 . don't add light precipitation (default) =1 . add light precipitation in warm section
i_sfct_gross	0	if use extended threshold for surface T gross check =0 use threshold from convinfo (default) =1 for cold surface, threshold for gross check is enlarged to bring more large negative innovation into analysis.

C. GSI Namelist: Name, Default Value, Explanation

Variable name	Default value	Description
CHEM		Chemistry data assimilation
berror_chem	.false.	if berror file is supplied for chemistry
oneobtest_chem	.false.	single observation test for chemistry
maginnov_chem	30.0	if oneobtest_chem=T magnitude of innovation for chemistry
magoberr_chem	2.0	if oneobtest_chem=T magnitude of observation error for chemistry
oneob_type_chem	pm2_5	if oneobtest_chem=T type of chemical observation
oblat_chem	45.0	if oneobtest_chem=T latitude of the observation
oblon_chem	270.0	if oneobtest_chem=T longitude of the observation
obpres_chem	1000.0	if oneobtest_chem=T pressure of the observation
diag_incr	.false.	if user wishes to output to a binary file increment
elev_tolerance	500.0	for surface chemical observation sometimes elevation (elev_obs) of the measurement is available (sometimes not).
tunable_error	0.5	tuning parameter to specify representativeness error for in-situ observations
in_fname	cmaq_input.bin	name of background file for cmaq
out_fname	cmaq_output.bin	name analysis file for cmaq
incr_fname	chem_increment.bin	if diag_incr=T name of the binary dump for pm2_5
laeroana_gocart	.false.	when true, do chem analysis with wrfchem and modis
l_aoderr_table		
aod_qa_limit		
luse_deepblue		
aero_ratios		
wrf_pm2_5		

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