Berkeley **H**igh **R**esolution (**BEHR**) Retrieval: Readme

Josh Laughner

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Summary:

The Berkeley High Resolution Retrieval is a high resolution NO_2 retrieval based on the NASA OMNO2 product from the Ozone Monitoring Instrument (OMI) aboard the Aura satellite. This retrieval improves the standard OMNO2 product in several ways:

- 1. A higher resolution terrain product (GLOBE Terrain Database) is used to calculate the terrain pressure of the pixels
- 2. A more frequent and finer resolution albedo is used, taken from the Moderate Resolution Imaging Spectrometer (MODIS) instruments aboard the Terra and Aqua satellites.
- 3. A MODIS cloud fraction is available to use in rejecting cloudy pixels, as the OMNO2 cloud fraction tends to overestimate cloud fractions if highly reflective surfaces are present

This document will describe the current state of BEHR, including its file structure and a changelog. As of this writing, the code is maintained in a Git repository on my machine. I will try to keep the main matter of this document up-to-date; however, if there is a discrepancy between the changelog and the main matter, defer to the changelog.

Contents

1 Authors

BEHR was initiated by Ashley Russell, who completed her Ph. D. in 2012. She showed that using high-resolution albedo and terrain data improved the OMI NASA Standard Product retrieval (see her papers in the Literature section below). Check the group website for her current contact information.

Luke Valin also completed his Ph. D. in 2012; he helped Ashley run the WRF-Chem simulations needed to get the high-resolution NO_2 profiles.

Josh Laughner took over development in 2013.

If you contribute to the development of BEHR, add your name and contribution to this list and update the change log.

- 2 Literature
- 3 Retrieving NASA data
- 3.1 Web sites
- 3.2 Group server file locations

4 Version Control

The core code for BEHR is contained in a Git repository on our group's Synology DS1813+ NAS file server. As of February 2, 2015, this server is located at IP address **128.32.208.13**. To connect to this server, your computer will need to have a UC Berkeley IP address. Connecting through an ethernet wall port is the recommended method, but you can also access it by connecting to the RC-Lab wifi network or from anywhere as long as you are connected to the Berkeley VPN. Credentials are:

Username: RCCohenLab

Password: SurfaceToStratopause

If you are not familiar with Git, I recommend reading chapters 1-4 at http://git-scm.com/doc, which includes information on installing Git on your system as well as the basic commands. If you are working on a Windows machine you make also want to look at http://guides.beanstalkapp.com/version-control/git-on-windows.html and follow their recommendations, although as I've never used Git on a Windows machine, I can't say for certain how well that works. Once you have Git installed and working on your machine, navigate to the folder that will be your working directory for the project in either Terminal (Mac) or Command Prompt (Windows) and run the command:

git clone ssh://RCCohenLab@128.32.208.13/volume1/share-sat/SAT/BEHR/BEHR_GitRepo.git

This will mirror the repository as a new folder in that directory. A second repository, at 128.32.208.13/volume1/share-sat/SAT/BEHR/MiscUtils.git contains some general utility Matlab scripts that I have found useful. Some were downloaded from the Matlab file exchange, many were functions I found myself needing rather often. A third repository is 128.32.208.13/volume1/share-sat/SAT/BEHR/AircraftProfiles.git, with functions to work with aircraft data sets. Additional repositories will be added to Table ??. I recommend that these be downloaded to folders called "BEHR", "Utils", and "NO2 Profiles" within your main Matlab directory (which you can check with userpath at the Matlab command prompt. This way, any internal links that I've set up to be robust as fullfile(userpath,x,y,z,...,file) should work smoothly on either a Windows or Mac platform.

This repository is in place now and will be kept relatively up-to-date as I progress. I will also try to remember to make a simple copy of the BEHR code to the file server which will not require the use of Git to obtain. However, there are several reasons you should consider

Address	Rec. folder	Contains
128.32.208.13/volume1/share-sat/SAT	BEHR	All the code needed to run
/BEHR/BEHR_GitRepo.git		BEHR
128.32.208.13/volume1/share-sat/SAT	Utils	Miscellaneous utilities not
/BEHR/MiscUtils.git		needed for BEHR but
		generally helpful
128.32.208.13/volume1/share-sat/SAT	NO2 Profiles	Functions to work with
/BEHR/AircraftProfiles.git		aircraft data sets, including
		code to validate satellite
		data against such data sets.
128.32.208.13/volume1/share-sat/SAT	Classes	Certain MATLAB classes I
/BEHR/BEHR_MatlabClasses_GitRepo.		wrote to help manage
git		certain tasks (e.g. error
		messages)

Table 1: Summary of the IP addresses, recommended folders within the main Matlab directory, and contents of the three Git repositories.

learning to use Git if you haven't yet. (If you have, you probably stopped reading this as soon as I told you where the repository was.)

- 1. It keeps everything in one place, and makes it easy to keep everything up to date. As long as you make changes in the directory that the Git repo consists of (and commit the changes periodically), those changes are tracked. So, you can roll back to an earlier version if something breaks, or see what code you (or I) changed.
- 2. Parallel development. If multiple people are developing BEHR at one point, each person can create their own branch and develop simultaneously, while still being able to update the project on the server, and eventually merge the development lines together.
- 3. Code sharing. Conversely, if two (or more) people are both using BEHR, this makes it easier to synchronize code when and if you want.

5 File structure

This section describes the key files of code in BEHR to make it run. Instructions for running it contained within this section are predicated on the idea that you're running BEHR on a desktop computer. For instructions on running it on a cluster, see §??.

5.1 Read data

The first step in running BEHR on new data is to read the NASA files into Matlab. This is done using the <code>read_omno2_v_aug2012.m</code> file. This is a Matlab function, but it is meant to be run from the editor without any input arguments. All of the properties that need to be set are coded into the function. However, it can accept start and end dates as inputs to the function, which can be useful to run it in batches.

The "v_aug2012" part of the name indicates that this file is intended to work with OMNO2 v. 3, which was released in Aug. 2012 (or at least the technical specs for it were). This function serves several purposes:

- 1. It reads all relevant variables from OMNO2 files into Matlab.
- 2. It averages the MYD06_L2 cloud product data to each OMI pixel.
- 3. It averages the MCD43C3 albedo product data to each OMI pixel.
- 4. It averages the GLOBE terrain data to each OMI pixel, converting from altitude to terrain pressure.

For each day that this function processes, a .mat file is saved with a single variable, Data. This variable is a data structure, in which each OMI swath for that day is stored under a different top-level index (i.e., Data(1) refers to the first swath of that day, and Data(2) the second, etc.). These data structures contain the data read from the OMNO2, MODIS, and GLOBE files as matrix fields. Each element of the matrix corresponds to an OMI pixel.

Production (i.e. not testing) files output from this script will generally have the name OMI_SP_yyyymmdd.mat. Filenames with additional information are generally testing files I have created in the course of various debugging runs, and should not be used to produce NO₂ data.

To run this file, enter the latitude and longitude boundaries for the area to retrieve (any OMI pixels outside this area will be discarded) and the date range to process. The first time you run this file, you may need to edit the paths to the various files; follow the instructions in the comments to identify which directory corresponds to which type of file. To specify a path to a folder on the lab server, Mac users should begin the path with /Volumes. PC users: the path should begin with the letter you chose when mounting the volume.

Should a new version of the OMNO2 files be released, lines 293 and 333–358 (dealing with various H5 loading functions to read OMNO2) might need updated to reflect any change in dataset names in the OMNO2 he5 files. The MYD06 and MCD43C3 files are loaded further down in the code; follow the comments.

5.2 Recalculate AMF and Tropospheric Column

This is handled by the BEHR_main.m function in BEHR/BEHR_Main/. Compared to the reading function, this is rather short, but it is the key component of BEHR. It goes through several steps:

- 1. It uses the TOMRAD look up table from NASA's OMNO2 product to generate box AMFs for each pixel, but using MODIS albedo and GLOBE terrain pressure. This is done for the clear and cloudy cases.
- 2. Reads in NO₂ profiles generated from WRF-Chem and bins them to each OMI pixel.
- 3. Calculates the full AMFs by combining the WRF profiles and box AMFs.

- 4. The new AMF is then applied to the tropospheric slant column, found by multiplying the NASA AMF with their vertical column.
- 5. Finally selected data fields are gridded onto a $0.05^{\circ} \times 0.05^{\circ}$.

The resulting outputs are saved in BEHR files that contain two variables: Data contains the original OMI pixel data, with the new BEHR AMF and NO_2 column density appended. OMI contains the variables gridded to $0.05^{\circ} \times 0.05^{\circ}$. Both retain the same structure wherein the top-level index represents a single swath. However, in OMI, every swath's grid covers the entire region of interest, and cells with no data for that swath have a fill value.

You may wonder that the grid is smaller than the regular OMI pixels; this technique is called *oversampling* and allows us to take advantage of the fact that the OMI pixels do not overlap exactly day-to-day to obtain an effective resolution greater than the pixel size if we average over longer time periods.

This function should require minimal upkeep even in the event a new NASA product is released.

5.3 Weight pixels and map

6 Running BEHR on a compute cluster

6.1 BEHR in Matlab

6.1.1 Parallelization

Starting in Jan 2015, the main BEHR code (read_omno2_v_aug2012.m and BEHR_main.m) had simple parallelization added to the code body. This is only intended to be used when the code is run on a cluster because in order to run operations in parallel, Matlab must send all relevant variables from the main instance to the parallel "workers" actually executing the code. Given the size of the variables routinely used in BEHR, this communication overhead can result in overall slower work than a serial execution if too few cores are used. As of 29 Jan 2015, I have yet to run any sort of rigorous benchmarking tests, but anecdotally, running BEHR in parallel with only 2 cores seemed to result in a slower execution than running it in serial.

There were numerous small changes to the code to enable parallelization, most importantly, the for loop over days was replaced with a parfor loop. parfor in Matlab allows multiple iterations of a for loop to run in parallel, and automatically handles the distribution of data. Compared to an spmd block, we give up control over how and when the data is distributed in exchange for Matlab handling it automatically.

Compared to previous versions of BEHR, most of the changes in the code (too numerous to list individually) were mainly to allow Matlab to "slice" variables appropriately for inclusion in a parfor loop. The change that is significant for the user is that a number of global variables were added to control both the action of the parallel loop and the paths to data.

The reason these variables were made global variables instead of inputs to the function was to allow them to be set in a run script once. This makes the run script more "bash like"

(programs compiled from a shell like bash—including GEOS-Chem and WRF-Chem—often reference environmental variables to determine how they compile).

The two variables controlling the action of the parallel loop are onCluster and numThreads. onCluster should be set to true in a run script whenever the script is being executed on a cluster. numThreads is, by default, set up in the run script template to take its value from an environmental variable, MATLAB_NUM_THREADS set in the bash shell that calls the Matlab instance running it. This was done because it is possible to set this environmental variable by referencing another that relates directly to how many CPU cores are available. See the example in §?? for an example.

The path variables are all defined in the .m files, and are set up such that if onCluster is true, the functions will look to global variables setup in the run script for those paths. (If any aren't defined, an error is thrown.) This is so that you only have to edit the run script to use BEHR on a cluster. If onCluster is false, the function looks to the paths coded into the functions.

6.1.2 Running it

The code was set up to be executing using a "runscript," which is a Matlab script file that can be called from the command line as:

```
matlab -nosplash -nodisplay -r "run('runscript.m')"
```

The argument -nosplash tells Matlab not to show the splash screen on startup, and -nodisplay tells Matlab that it shouldn't start the GUI interface. (Since we're working from the command line, we don't need it, and we don't want it to try to open it if we can't see it anyway.) -r "..." tells Matlab to execute the command given in quotation marks as soon as it starts up. In this case, that is to execute the runscript in the current directory.

A template for a BEHR runscript can be found in BEHR/Run_Scripts/. You'll notice that the runscript template does several things:

- 1. Sets the global variable onCluster to true. This lets any scripts that use that variable know that it is running on a cluster and should activate any parallel elements in the code.
- 2. Sets the variable numThreads. By default this is set to the variable of the environmental variable \$MATLAB_NUM_THREADS from the shell that executed this instance of Matlab. More on why this is preferable below.
- 3. Detects any active parallel pools and shuts them down before exiting.
- 4. Everything is wrapped in a try-catch block that will, in the event of an error, prints the error information to the console, closes any active parallel pools, and exits with status code > 0.

This is a very general template, so you can use it to call any function you parallelize using the global variables on Cluster and numThreads. For BEHR, you'll also have to set all the global path variables for read_omno2_v_aug2012.m and BEHR_main in the runscript.

6.1.3 Submitting to the cluster queue

Like any job you want to run on a computing cluster, you'll need to write a shell script that is put into the queue for the cluster. Since the Savio cluster that I use operates with the SLURM scheduler, this section will be written from the point of view of submitting to SLURM using the bash shell. (I assume that if you use tcsh or another shell that you know what you're doing well enough to make the necessary adjustments.) Below will be an example submit script, each important line will be described afterward.

```
1 #!/bin/bash
2 #
3 # Job Name:
4 #SBATCH --job-name=BEHR
5
6 # Partition:
7 #SBATCH --partition=savio
8 #
9 # Account:
10 #SBATCH --account=ac_aiolos
11 #
12 # QoS:
13 #SBATCH --qos=condo_aiolos
14 #
15 # Number of nodes and processors per node
16 #SBATCH --nodes=1
17 #SBATCH --ntasks-per-node=20
18 #
19 # Wall clock limit:
20 #SBATCH --time=24:00:00
21 #
22 ## Run command
23 export MATLAB_NUM_THREADS=$((SLURM_NTASKS_PER_NODE-1))
24 cd /global/home/users/<me>/<behr_run_dir>
25 module load matlab
26 matlab -nosplash -nodisplay -logfile "runlog.txt" -r "run('runscript_behr
27 MATLAB_EXIT=$?
28 exit $MATLAB_EXIT
```

- Line 1 Lines starting with a #! are called a *shebang* in bash-speak, this one tells the computer to run the script using the bash shell. Including this is a safety measure; it ensures the script is always run using bash if another shell interpreter is active.
- Line 2 A # indicates a comment in bash

- Line 4 The #SBATCH at the beginning of this line tells the SLURM scheduler that a setting is being passed. Bash ignores it because of the #, but SLURM does not. In this case, we're setting the name of the job that will appear in the queue.
- Line 7 Tells SLURM which partition of nodes to run on. We use "savio" unless we need lots of RAM.
- Line 10 The SLURM account that would be charged for running (I think). Ron is under the "aiolos" account; the "ac" indicates that it is the account.
- Line 13 "QoS" determines what rules the job is run under and how compute time is charged. The part after the underscore will always be the account, "aiolos." The part before the underscore determines the rules. "condo" means that we can use up to 8 nodes at a time and won't be charged for it.
- Line 16 How many nodes to to. A node is the computing unit of the cluster—each node will only run one job at a time.
- **Line 17** How many cores to use on each node. Each node has two 10-core processors for 20 cores maximum per node.
- Line 20 How long to let the job run before forcing it to quit. Here, we set it to 24 hours. It's good practice to do this to prevent a job from running indefinitely.
- Line 23 export in bash means "set this as an environmental variable" which programs executed from this shell can access. The \$(()) in bash means to evaluate an arithmetic expression and return the result. So here we're saving one less than the number of tasks per node in the MATLAB_NUM_THREADS variable. I do this to be a little bit careful to leave a core free for the main Matlab instance, although I don't know if that's strictly necessary.
- Line 24 Change to our run directory (just in case the job starts somewhere else). <me> and <bent_run_dir> are just placeholders.
- Line 25 Savio organizes applications by modules, here load the matlab module to be able to run matlab.
- Line 26 Execute matlab with command line arguments. The only new one is -logfile "runlog.txt" which saves all Matlab command window output to the file runlog.txt. One mistake to avoid is including the -nojwm flag, because (apparently) the parallel computing toolbox needs Java to work. Don't ask me why.
- Line 27 The \$? is the last given exit code. We save this to a variable...
- Line 28 ...and then exit this script with that exit code. This will let you know if the script succeeded or failed.

If this script is named e.g. matrum then typing sbatch matrum on the cluster will submit it to run. You can check the status of all jobs with squeue.

The reason that we use the variable MATLAB_NUM_THREADS to pass the number of available cores to the run script is to be a little careful about not causing Matlab to request more cores than we've set aside. By deriving MATLAB_NUM_THREADS from the SLURM variable indicating the number of cores to be used per node, we ensure that a change to the SLURM

settings is propagated through to our Matlab instance without any intervention on our part.

6.2 Resources

- 1. Matlab documentation on parfor loops: http://www.mathworks.com/help/distcomp/parallel-for-loops-parfor.html
- 2. Matlab parfor loops, classification of variables: http://www.mathworks.com/help/distcomp/classification-of-variables-in-parfor-loops.html
- 3. The High Performance Computer (HPC) User Guide: http://research-it.berkeley.edu/services/high-performance-computing/user-guide
- 4. SLURM Documentation: https://computing.llnl.gov/linux/slurm/documentation.html
- 5. List of SLURM parameters (i.e., what can be set in the bash run script on the lines beginning with #SBATCH: https://computing.llnl.gov/linux/slurm/sbatch.html

7 Additional utilities

7.1 Verification

8 Changelog

A notation of (dev) after the version number indicates that the full archive of BEHR data was never produced using that version (i.e. "development only").

Date V	Version Number	Description
2012	2.0A	Original version by A. Russell, using OMI SP 2 with MODIS cloud and albedo, GLOBE terrain database, and WRF-Chem profiles (12 km for full US).
May 2013	2.0B (dev)	Version (unvalidated at this point) updated by J. Laughner to use OMI SP 2 and MODIS cloud collection 6. Albedo, terrain products, and WRF-Chem profiles unchanged. MODIS and GLOBE data import now done directly by read_omno2_v_aug2012, "preprocessing" of these files into Matlab .mat files no longer necessary.