A User’s Guide for the

GBT Mapping Pipeline

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# Introduction

The GBT Data Processing Pipeline facilitates reducing spectroscopic data taken with the GBT. There are two primary software tools that constitute the pipeline: the first is called the “mapping pipeline” and it helps users create FITS image cubes from GBT mapping data. The second is the “spectral pipeline” and it helps users process single-pointing GBT spectra. At this time, the spectral pipeline is available only as a beta-release product. This document describes version 1.0 of the mapping pipeline.

The mapping pipeline was developed initially to support mapping with the KFPA, but it can be applied to data taken with any of the receivers that use standard position-switched or frequency-switched observing modes. Pipeline tools can also help with reducing other types of data, such as W-band mapping observations.

To use the pipeline effectively, you should follow certain guidelines and observing procedures when taking your data. The observing steps are described here, are relatively simple, and are implemented in Astrid.

While this document addresses observing strategies that use the GBT spectrometer, the methods will be nearly identical for the new VEGAS spectrometer when it is introduced for science observations.

# Getting Started

Where to Run the Pipeline: Currently, the GBT pipeline has certain dependencies and it runs only on 64-bit, RedHat 6 computers in Green Bank. The computer *arcturus.gb.nrao.edu* is reserved for pipeline use. We recommend that you use *arcturus* for all pipeline work at this time.

Disk Space: The pipeline typically requires a significant allocation of disk space. Even small data sets use several gigabytes. So generally, you must make use of the Green Bank scratch areas to use the pipeline.

We recommend /lustre/pipeline/scratch/user\_id/

In the above directory, replace “user\_id” with your linux user ID, e.g. “jbraatz”. At the command line type:

mkdir /lustre/pipeline/scratch/user\_id

It is also essential that you are a member of the ‘aipsuser’ group. You can check this by typing ‘groups’ at the command line. If ‘aipsuser’ is not listed, contact the GB helpdesk to have GB computing add you to the group. The NRAO helpdesk is accessible at: [http://help.nrao.edu](http://help.nrao.edu/)

# A First Look at the Mapping Pipeline

You can run the pipeline using the command:

% gbtpipeline

For quick usage information, type the command with no parameters.

Here is a sample pipeline execution using KFPA position-switched test data that observed NH3 toward W51:

REPLACE WITH VEGAS DATA?

% gbtpipeline -i /home/gbtpipeline/reference-data/TKFPA\_29.raw.acs.fits -m 14:24 --refscan 13 --units Jy

This command maps data from scans 14-24, using scan 13 as the reference observation. The data are calibrated to units of Jy. The program creates several products, including three maps stored as FITS files, calibrated spectra from each spectrometer sampler in separate FITS files, and logs of the pipeline execution. The output maps are:

W51\_23740\_MHz\_cont.fits – a single-channel “continuum” map

W51\_23740\_MHz\_cube.fits – a multi-channel data cube

W51\_23740\_MHz\_line.fits – a continuum-subtracted data cube

You can examine these data cubes using the casa viewer, e.g. :

% casaviewer W51\_23740\_MHz\_line.fits

Select the menu option “Tools -> Spectral Profile”, then left-click on the crosshair icon, and left-click on the map. This will display a spectral cut through the cube.

# Basic Operation of the Pipeline

The input to the pipeline is an SDFITS data file generated from a GBT mapping observation, and from that file the pipeline generates image cubes. The pipeline operates in three basic stages.

1. The first stage calibrates the spectral line data and produces FITS files compatible with the GBTIDL “keep file” format of SDFITS.
2. The second stage uses the utility program *sdfits2aips* to convert these SDFITS files into a FITS format that can be read by AIPS.
3. The third stage uses AIPS to grid these calibrated data and produce the output data cubes.

In typical use, a single call to the *gbtpipeline* command executes all three stages of operation. You can specify the range of scans to be mapped and identify the reference scans used for calibration (if the data are “position switched”), as in the NH3 mapping example, above. A standard position-switched mapping observation begins with a reference scan, executes the mapping scans, and then (optionally) finishes with another reference scan, all using a common spectrometer configuration. In the *gbtpipeline* command, you can specify a second reference scan in the parameter list, and in this case the two reference scans will be interpolated for each integration in the map, as in this example:

% gbtpipeline -i /home/gbtpipeline/reference-data/TKFPA\_29.raw.acs.fits -m 14:24 --refscans 13,26

Here are the standard options available from the *gbtpipeline* command line:

## Pipeline Options

**Input file or project directory:**

-i FILE, --infile FILE

SDFITS file name containing map scans

**data selection:**

-m MAPSCANS, --map-scans MAPSCANS

map scans, e.g. "10:20,30:40" identifies two ranges of 11 scans each

--refscan REFSCANS, --refscans REFSCANS

reference scan(s), e.g. "4,13" to identify two reference scans

-f FEED, --feed FEED

feeds, e.g. "0:6" identifies feeds 0 through 6

-p POL, --pol POL

polarizations, e.g. "0,1"

-w WINDOW, --window WINDOW

spectral windows, e.g. "0,1,2,3"

-c CHANNELS, --channels CHANNELS

channel selection, e.g. "100:200". (Passed directly to sdfits2aips)

**control:**

--imaging-off

If set, will calibrate data and write calibrated SDFITS files but will not

create image FITS files.

-a AVERAGE, --average AVERAGE

average the spectra over N channels (passed directly to sdfits2aips)

-n N, --rms-flag N flag integrations with RMS noise > N kelvin (passed directly to

sdfits2aips)

--median-baseline-subtract N

subtract median-filtered baseline, with a median window half width of

N channels (passed directly to sdfits2aips)

--display-sdfits2aips-plots

display sdfits2aips plots of calibrated spectra

**calibration:**

-u {ta,ta\*,tmb,jy}, --units {ta,ta\*,tmb,jy}

calibration units. Default: tmb

--spillover-factor SPILLOVER

rear spillover factor (eta\_l) Default: .99

--aperture-efficiency APERTURE\_EFF

aperture efficiency for spectral window 0. Other window efficiencies

are adjusted to this value. (eta\_A) Default: .71

--main-beam-efficiency MAINBEAM\_EFF

main beam efficiency for spectral window 0. Other window efficiencies

are adjusted to this value. (eta\_B) Default: .91

--beam-scaling BEAMSCALING

comma-separated beam scaling factors for each feed and polarization in

the order 0L,0R,1L,1R,etc.

-t ZENITHTAU, --zenith-opacity ZENITHTAU

zenith opacity value (tau\_z). Default: determined from GB weather

prediction tools.

**output:**

-v N, --verbose N

Set the verbosity level-- 0-1:none, 2:errors only, 3:+warnings,

4:+user info, 5:+debug (default: 0)

--clobber Overwrite existing output files.

--keep-temporary-files

Does not remove intermediate imaging files if set.

# Advanced Operation of the Pipeline

Basic usage of the *gbtpipeline* command, as described above, should meet the data reduction needs of many projects. Sometimes, however, you may need to customize the mapping results or take advantage of certain features that are not available from the standard *gbtpipeline* parameters. In this case, you can run any of the three stages of the pipeline separately, customizing the results at each stage.

### Stage 1: Calibration

The first stage of the pipeline calibrates the data and creates an SDFITS file for each spectrometer sampler. You can halt the *gbtpipeline* process after Stage 1 by specifying the command-line option “--imaging-off”.

As an alternative, you may use GBTIDL to produce the SDFITS files. GBTIDL gives you detailed control of the calibration and flagging, on an integration-by-integration basis. The SDFITS file must contain one record for each integration in the map, and each record should contain a calibrated spectrum. You can either create one SDFITS file with all the data, or multiple SDFITS files containing partial data sets (for example, separating feeds into separate SDFITS files for individual processing). Separate SDFITS files are easily combined for mapping in a later step.

The GBTIDL 'flag' procedure allows you to write NaNs for flagged channels. However, the sdfits2aips program that converts SDFITS to an AIPS-friendy format rejects any spectra that have a NaN, in any channel.

To ignore flagged data you have two options:

1. If only certain channels are predictably bad, you could use the sdfits2aips channel filtering ‘-c’ option to only select good channels for imaging.

2. You could replace bad channels with values interpolated from neighboring channels.

One option is to use the GBTIDL 'replace' procedure which replaces the values between two channels with a linear interpolation using the values of the two edge channels in the region to be replaced. i.e. if channel 100 is bad you'd do:

replace,99,101

And then channel 100 would contain the average of channels 99 and 101. Documentation can be found here: [http://www.gb.nrao.edu/GBT/DA/gbtidl/release/user/](http://www.gb.nrao.edu/GBT/DA/gbtidl/release/user/" \t "_blank)

It may be slow to do this for all spectra, but should work.

### Stage 2: File Conversion

With an SDFITS file in hand, you can convert the file to an AIPS-compatible format using the utility program *sdfits2aips*. A typical execution will look like:

% sdfits2aips –l –o myfile.aips myfile.fits

The “-l” option suppresses the creation of a plot file. The *sdfits2aips* program has a host of additional options that can be examined by typing “sdfits2aips” at the linux prompt, with no parameters.

The product of Stage 2 is one “aips” file for each SDFITS file from Stage 1.

### Stage 3: Combining Data and Imaging

Prior to running Stage 3, you should have a set of “aips” files. You can then run the third stage, with more control of the mapping procedure, manually. Here we describe that process.

**aipspy Scripts:** The pipeline includes a utility program called *aipspy* that runs ParselTongue scripts in AIPS. To use ParselTongue scripts, you need to know the AIPS ID used on your behalf by the pipeline. Get the ID from the linux prompt using:

% id –u

There are several *aipspy* scripts in the standard gbtpipeline distribution, contributed mainly by Glen Langston. Two important scripts are:

1. load.py : Combine data from multiple mapping procedures
2. image.py : Map data

So, you can execute the third stage of the pipeline, in its basic form, by executing the following two linux commands:

% aipspy /home/gbtpipeline/release/contrib/load.py <aips\_id> <file1.aips> [<file2.aips>]

% aipspy /home/gbtpipeline/release/contrib/image.py <aips\_id>

The result of these commands is three FITS files, one with the full image data cube, one with a continuum map, and one with a line map.

**load.py** The parameters to load.py are:

load.py <aipsNumber> <file1.aips> [<file2.aips>] […]

where the first parameter is the AIPS ID and subsequent parameters are the aips files to be combined in the final image. You can use one, two, or more than two files in the list. The “load” step must be executed prior to the “image” step.

**image.py** The parameters to image.py are:

image.py <aipsNumber> [<nAverage>] [<mapRaDeg>] [<mapDecDeg>] [<imageXPixels>] [<imageYPixels>] [<refFreqMHz>]

1. [<nAverage>] spectral channels to average
2. [<mapRaDeg>] RA of map center in degrees
3. [<mapDecDeg>] Dec of map center in degrees
4. [<imageXPixels>] pixel size of the map in the X direction
5. [<imageYPixels>] pixel size of the map in the Y direction
6. [<refFreqMHz>] rest frequency in MHz (used to calculate the velocity scale)

If any one of the optional parameters in the list is specified, all the preceding parameters must also be specified.

# Examples Using aipspy

Here we give examples of several common ways to use the *aipspy* pipeline tool.

### Example: Combining Data from Multiple Maps

Suppose you have several data sets from independent executions of your GBT mapping observation. You can combine and map the data using the following procedure.

First run the pipeline on each of the maps separately, specifying the “--imaging-off” option. Each execution will generate a set of SDFITS files and aips files.

gbtpipeline –i my\_sample\_data.fits –m 10:19 --refscan 9 --imaging-off

gbtpipeline –i my\_sample\_data.fits –m 21:30 --refscan 20 --imaging-off

Next create the aips files for each FITS file generated in the previous step:

sdfits2aips –l –o file1.aips file1.fits

sdfits2aips –l –o file2.aips file2.fits

Now combine the data using the *aipspy* command with load.py:

aipspy /home/gbtpipeline/release/contrib./load.py <aips\_id> <file1.aips> <file2.aips>

Finally you can map the data using the *aipspy* command with image.py:

aipspy /home/gbtpipeline/release/contrib/image.py <aips\_id>

## Example: Mapping Pre-calibrated Data

Under certain circumstances, *gbtpipeline* will be unable to calibrate GBT data. For example, the pipeline cannot currently calibrate W-band data, since observations with that receiver use a unique observing sequence for calibration. To map data such as this, you must first calibrate the data in GBTIDL and produce a “keep file” that has one record per beam, per integration in the map.

You can then grid the data with the following steps. First, convert the keep file to a FITS format (aips file) that can be read by the AIPS FITS reader:

% sdfits2aips –o my\_aips\_file.aips my\_keep\_file.fits

Next, load the data into AIPS with the *load.py* script:

% aipspy /home/gbtpipeline/release/contrib/load.py <aipsid> my\_aips\_file.aips

And finally map the data with the *image.py* script:

% aipspy /home/gbtpipeline/release/contrib/image.py <aipsid>

## Example: Line Maps and Temperature from NH3(1,1) and NH3(2,2)

Here we give an example of using *aipspy* scripts to generate line maps and a temperature map from the NH3 data. We make line maps by restricting the spectral channels used in the map to only those near the NH3 line centers. Glen Langston contributed the scripts.

To generate the maps, you can run these exact commands using the pipeline in Green Bank:

gbtpipeline -i /home/gbtpipeline/reference-data/TKFPA\_29.raw.acs.fits -m 14:24 --refscan 13 -a 3 -v 4

aipspy /home/gbtpipeline/release/contrib/sumLine.py 13482 W51-NH3-11 23694.506 60. 20.

aipspy /home/gbtpipeline/release/contrib/sumLine.py 13482 W51-NH3-22 23722.6336 60. 20.

aipspy /home/gbtpipeline/release/contrib/tempNH3\_1122.py 13482 W51 60. 10. .2

The first command is the pipeline mapping command we encountered already in our initial example. The second and third commands use the sumLine.py script to generate single-channel maps averaged through the line centers from the NH3(1,1) and NH3(2,2) lines. The parameters of the sumLine.py procedure are:

sumLine.py <aipsNumber> <fileName> <restFreq> <lineVel> <lineWid>

1. <aipsNumber> : the AIPS ID
2. <fileName> : a name for the output file
3. <restFreq> : the rest frequency of the line in MHz
4. <lineVel> : the line center velocity in km/s, and
5. <lineWid> : the line FWHM in km/s.

The fourth command in the sequence generates the temperature map from the original line cube using the temp\_NH3\_1122.py script. The parameters for this command are:

tempNH3\_1122.py <aipsNumber> <fileName> <restFreq> <lineVel> <cutOff>

1. <aipsNumber> : the AIPS ID
2. <filename> : a name for the output file
3. <lineVel> : the line center velocity in km/s
4. <lineWid> : the line width (FWHM) to average, and
5. <cutOff> : the minimum line intensity to use in the line ratio (smaller values result in blanked pixels in the temperature map)

These scripts are available in the *contrib* directory. Besides being useful in their own right, they also serve as good examples to help you develop your own scripts.

# Optimal Observing Configuration for the Mapping Pipeline

Previous examples demonstrated how you could identify specific mapping scans and reference scans for the pipeline’s calibration sequence. By specifying mapping and reference scans explicitly in this manner, you have direct control over the calibration process. An alternative is to allow the pipeline to determine mapping and reference scans automatically. If you do not specify scan ranges at the command line, the pipeline will examine the entire input data file and attempt to process each mapping observation it encounters there. To identify mapping scans and reference scans, the pipeline uses header information that must be set explicitly during the observation. The *KFPA Data Reduction Guide,* by Langston et al., describes this process. (https://safe.nrao.edu/wiki/bin/view/Kbandfpa/KfpaReduction).

The following sample astrid script demonstrates a KFPA observing sequence with keywords set so that the pipeline can identify reference scans properly.

# Example Astrid script demonstrating KFPA Mapping for use with pipeline “allmaps”

# We assume that the telescope has been configured prior to running this script

# Identify the source catalog

Catalog(“/home/userid/mycatalog.cat”)

# Define procedures with scan annotations and header values

execfile("/home/astro-util/projects/TKFPA/kfpaMapInit")

target = "W51" # define mapping target

off = "W51-Off" # define a map reference location, with no emission

Slew(target)

Balance()

# Check the levels are correct

Break("Check Balance”)

#perform a total power observation at the reference location

OffTrack( off, None, 30.0, "1")

#Tell Pipeline that mapping is starting

SetValues("ScanCoordinator",{"scanId":"Map"})

RALongMap( target,

Offset("Galactic", 0.33, 0.0), # This is a galactic coordinate map

Offset("Galactic", 0.0, 0.10),

Offset("Galactic", 0.0, 0.008),

140.0, "1")

#perform the final total power reference obs

OffTrack( off, None, 30.0, "1")

The pipeline can then process the resulting data, correctly associating the reference scans with the mapping data. The pipeline command is simply:

gbtpipeline –i my\_sample\_data.fits

# Subtracting Spectral Baselines

A typical GBT mapping observation uses short integration times per pixel (a few seconds), and the resulting spectral baselines are flat compared to the thermal noise. The pipeline therefore uses a very basic spectral baseline subtraction, implemented with the AIPS task *IMLIN* in the *image.py* script. The default behavior is to fit a DC baseline (polynomial of order 0) determined from the edges of the spectrum. Specifically, the baseline is fit to channels [0.04, 0.12]\* nchan and [0.81,0.89]\*nchan. You can modify this behavior by editing the *image.py* script directly. Follow these steps:

1. cd to the directory with your mapping data
2. copy /home/gbtpipeline/release/contrib/image.py to the current directory
3. edit image.py appropriately (search for the “imline” command)
4. re-run the pipeline with the “--imaging-off” option
5. run “aipspy load.py <aipsid> <file.aips>”
6. run “aipspy image.py <aipsid>” (note we are using the local copy of image.py)

A future upgrade to the pipeline will incorporate baseline-fitting parameters directly in the *gbtpipeline* command line.

For more detailed control over the baseline fitting and other aspects of the calibration and flagging, you can use GBTIDL to calibrate the data and use pipeline tools for “Stages” 2 and 3.

# Cleaning Up Disk Space

The pipeline creates large data files, and may completely fill the available disk space if care is not taken to clean up unused files. You should remove all intermediate files that are not needed for additional processing. This includes “aips” files generated during the calibration of data, and AIPS files used in intermediate stages of gridding the data. A utility is available to clean up old AIPS files, and this should be run after each use of the pipeline.

% aipspy /home/gbtpipeline/release/contrib/clear\_AIPS\_catalog.py <aips\_id>

Recall that you can find your AIPS ID by the linux command “id –u”.

Note: All files stored under the AIPS ID used by the pipeline will be removed with this operation, so it is important that you not store any non-pipeline AIPS files under this AIPS ID.

# Appendix 1: Calibration Equations

The pipeline uses the following set of equations for calibration:

System temperature

*Tsys = Tcal \* ref80caloff / (ref80calon – ref80caloff) + Tcal / 2*

Antenna temperature

*Ta = Trefsys \* (sig-ref)/ref*

Corrected antenna temperature

*T\*a = Ta \* eτ*0/*sin*(*el*) / *ηl*

Main beam brightness

*TMB* = *T*\**a* / (1.32 \* *ηA*)

Flux density (Jy)

*Sν = T\*a / (2.85 \* ηA)*